Ribonomic Analysis of Human Pum1 Reveals cis-trans Conservation across Species despite Evolution of Diverse mRNA Target Sets

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Received 29 January 2008/Returned for modification 20 March 2008/Accepted 1 April 2008

PUF family proteins are among the best-characterized regulatory RNA-binding proteins in nonmammalian species, but relatively little is known about mRNA targets or functions of mammalian PUF proteins. In this study, we used ribonomic analysis to identify and analyze mRNAs associated with ribonucleoproteins containing an endogenous human PUF protein, Pum1. Pum1-associated mRNAs were highly enriched for genes encoding proteins that function in transcriptional regulation and cell cycle/proliferation, results consistent with the posttranscriptional RNA regulon model and the proposed ancestral functions of PUF proteins in stem cell biology. Analysis of 3′ untranslated region sequences of Pum1-associated mRNAs revealed a core Pum1 consensus sequence, UGUAIUAU. Pum1 knockdown demonstrated that Pum1 enhances decay of associated mRNAs, and relocalization of Pum1 to stress granules suggested that Pum1 functions in repression of translation. This study is the first in vivo genome-wide mRNA target identification of a mammalian PUF protein and provides direct evidence that human PUF proteins regulate stability of associated mRNAs. Comparison of Pum1-associated mRNAs to mRNA targets of PUF proteins from Saccharomyces cerevisiae and Drosophila melanogaster demonstrates how a well-conserved RNA-binding domain and cognate binding sequence have been evolutionarily rewired to regulate the collective expression of different sets of functionally related genes.

PUF family RNA-binding proteins (RBPs) are among the best-characterized regulators of posttranscriptional gene expression in nonmammalian eukaryotes. Named for the founding members Pumilio (32) and FBF (61), PUF proteins are represented throughout the eukaryotic lineage (47). The common feature of PUF proteins is the PUF homology domain (PUF-HD), an RNA-binding domain typically consisting of eight imperfect repeats of a 32-amino-acid sequence (60). The overall sequences of PUF proteins from different species are not highly similar outside of the PUF-HD, although the PUF-HD is incredibly well-conserved (46). This extreme conservation of the PUF-HD suggests that posttranscriptional regulation of gene expression by PUF proteins has remained important throughout evolution. Although there is a growing body of knowledge concerning PUF proteins in nonmammalian model organisms, relatively little is known about the functions or mRNA targets of PUF proteins in mammals.

Genetic analyses have revealed diverse functions of PUF proteins, such as embryo patterning in Drosophila melanogaster (32), germ line establishment and maintenance in Drosophila (14, 33, 42) and Caenorhabditis elegans (4, 10, 30, 53, 61), and mitochondrial function in Saccharomyces cerevisiae (16). PUF proteins have been found to function as repressors of gene expression through both repression of translation and enhancement of decay of target mRNAs (13, 19, 20, 41, 55). Crystal structure analysis of the PUF-HD from Drosophila Pumilio revealed that it forms a crescent shape (12), with protein-RNA interactions occurring on the inner concave surface and protein-protein interactions on the outer surface.

 Genome-wide target identification of five S. cerevisiae PUF proteins, Puf1 to -5 (17), and the single Drosophila PUF protein, Pumilio (18), demonstrated that each protein binds to a specific group of functionally related mRNAs distinct from those mRNAs bound by any other PUF protein (except for Puf1 and Puf2, which bind overlapping sets of mRNAs). Puf3 was found to bind almost exclusively to mRNAs of nuclear-encoded mitochondrial proteins, and this binding was later demonstrated to have functional consequences when it was shown that Puf3 regulates stability of target messages in a condition-specific manner (13) and regulates mitochondrial biogenesis and motility in S. cerevisiae (16). These experiments represent a compelling example of the posttranscriptional RNA operon/regulon model (26, 28), demonstrating a mechanism through which expression of functionally related genes can be coordinately regulated at the level of the mRNA. Targets of Drosophila Pumilio also contained potential RNA regulons, most notably the vacuolar-type ATPase and the embryopatterning cascade, which Pumilio mutants are known to disrupt (18). These studies also demonstrated for the first time that while the cis-trans interactions between PUF proteins and target mRNAs are similar, target messages of PUF proteins are not conserved through evolution, at least from S. cerevisiae to Drosophila (18).

Mammalian genomes contain two PUF genes, Pum1 and Pum2 (46), both of which have been studied to a limited extent. The Pum2 gene has been knocked out in the mouse, but the only obvious phenotype was smaller testis size with no effect on fertility (58). Several potential human Pum2 mRNA targets have been discovered by various groups (15, 31, 48, 56); how-
ever, the in vivo genome-wide targets of the protein have not been identified. Expression of reporter constructs containing Pum1 target 3′ untranslated regions (UTRs) was shown to be repressed by Pum1 overexpression, but the mechanism of repression was not determined (31). Rat Pum2 was found to localize to stress granules in hippocampal neurons, with Pum2 knockdown interfering with stress granule formation and Pum2 overexpression inducing aggregates that co-stained with stress granule markers (52). Even less is known about Pum1. The human Pum1 RNA-binding domain can bind to the Nanos response element, an mRNA sequence bound by Drosophila Pumilio, and has been found by crystal structure analysis to interact with RNA in a very modular fashion, with each of the eight PUF repeats directly contacting a single RNA base (54). Recombinant Pum1 was shown in vitro to interact with the CNOT8 protein, a member of the CCR4-NOT deadenylase complex (20), suggesting that enhancement of target mRNA deadenylation and decay may be a conserved mechanism of PUF protein function (57).

Due to the lack of knowledge regarding both targeting and function of mammalian PUF proteins, this study was undertaken to identify genome-wide mRNAs associated with Pum1 protein, along with mechanisms by which Pum1 regulates these mRNAs. By determining the genome-wide targets of a PUF protein from a third species, humans, we gained insight as to how the use of a highly conserved RNA-binding domain and cognate binding sequence has changed throughout evolution by regulating different sets of functionally related mRNAs.

MATERIALS AND METHODS

Cell culture. HeLa S3 cells, used for all experiments except immunofluorescence (IF) assays, were grown in Ham’s F-12 supplemented with 10% fetal bovine serum. HeLa CCL2 cells were used for IF and were grown in Dulbecco’s modified Eagle’s medium supplemented with nonessential amino acids and 10% fetal bovine serum. All cells were grown in humidified incubators at 37°C and 5% CO2.

Immunoprecipitations. Immunoprecipitation (IP) reactions were performed as described previously (27, 50). Briefly, 10 μg of anti-Pum1 antibody (goat polyclonal, Bethyl Labs) was incubated overnight with protein G-agarose beads. The beads were washed, then buffer and cell lysate were added, and the reactions were performed as described. Western blotting or 1 ml of Trizol was added and RNA extracted for microarray and reverse transcription-PCR (RT-PCR) experiments. Identical IPs were performed using the CNOT8 protein, a member of the CCR4-NOT deadenylase complex (20), suggesting that enhancement of target mRNA deadenylation and decay may be a conserved mechanism of PUF protein function (57).

Due to the lack of knowledge regarding both targeting and function of mammalian PUF proteins, this study was undertaken to identify genome-wide mRNAs associated with Pum1 protein, along with mechanisms by which Pum1 regulates these mRNAs. By determining the genome-wide targets of a PUF protein from a third species, humans, we gained insight as to how the use of a highly conserved RNA-binding domain and cognate binding sequence has changed throughout evolution by regulating different sets of functionally related mRNAs.

Microarray analysis. Arrays were printed at the Duke Microarray Facility using the Genomics Solutions OmniGrid 300 arrayer. The arrays contained the following microarray oligonucleotide sets: the Human Operon v3.0.2 oligonucleotide set (Oligo Source), which consists of 34,602 unique optimized 70-mers. RNA quality was ascertained using an Agilent 2100 bioanalyzer (Agilent Technologies). All arrays were subject to Loess normalization within each array and scale normalization across all arrays using the Agilent Feature Extraction software (Agilent Technologies). The analysis method, using either the Mixtools package in R (43, 59), each iteration of mixture modeling was initialized at different values in the distribution, and the parameters from the model with the highest likelihood were used to create log of odds (LOD) scores of Pum1 ribonucleoprotein (RNP) association by comparing the weighted probability density functions of the Pum1-associated versus background distributions. LOD scores greater than 0 have a higher probability of being in the Pum1-RNP-associated distribution compared to background distribution; therefore, a LOD score of 0 was used as a cutoff for determining Pum1-associated probes.

mRNA decay rates. mRNA levels of target messages were determined by RT-qPCR at hourly time points after addition of 5 μM actinomycin D. The nontarget message GAPDH was used for normalization. Exponential decay curves were fit to points representing the means of three biological replicates, and half-lives were calculated based on the equations describing these best-fit curves.

Immunofluorescence. IF was performed essentially as described elsewhere (25). Briefly, cells were grown on glass coverslips precoated with 0.1% gelatin and then fixed in 4% paraformaldehyde followed by 100% methanol. Coverslips were then blocked, incubated with appropriate antibodies, and mounted to slides. Pum1 and Ddx6 antibodies and Pum1 blocking peptide were from Bethyl Labs, and HuR antibody was supernatant from hybridoma 3A2 cells. Images were captured using a Zeiss Axioscap microscope with a 63x objective and Axiovision 4.6.3 SP2 software. Images were then combined and minimally processed using Adobe Photoshop, with all images processed identically. For cell stress, sodium arsenite was added to growth medium at a concentration of 0.5 mM, and cells were fixed as described after 45 min of stress. Orthologues of Puf3 and Pumilino target genes were determined using the online database mining tool Biomart (www.biomart.org).

RESULTS

Identification of Pum1-associated mRNAs. In this study, we used the RNA-binding protein immunoprecipitation-chip (RIP-chip) method to identify genome-wide mRNAs associated with RNPs containing Pum1 (21, 27, 50). All experiments were performed in human HeLa cells, and all Pum1 IPs were performed with a goat polyclonal antibody raised against a small peptide region specific to the Pum1 protein. Negative control IPs were performed with preimmune goat serum. Be-
fore performing RIP-chip, we first confirmed specific IP of Pum1 protein from cytoplasmic lysate by IP-Western blotting, using identical conditions except that in the final step protein was extracted from the beads and used for sodium dodecyl sulfate-polyacrylamide gel electrophoresis and immunoblotting. We were able to efficiently recover Pum1 protein in our IP reactions and correspondingly deplete a substantial amount from the original starting material (Fig. 1A). After confirmation of efficient Pum1 protein recovery, we next sought to determine whether specific mRNAs associated with Pum1 were enriched in the Pum1 IP pellet. We chose cyclin B1 as a potential target due to its conservation as a PUF target in other species (3, 40, 45, 47). RT-PCR analysis confirmed specific enrichment of cyclin B1 mRNA in the Pum1 IP compared to the negative control (Fig. 1B). After confirming that specific mRNAs associated with the Pum1 protein could be identified using these procedures, we next performed RIP-chip in biological triplicates using custom spotted cDNA microarrays. Each biological replicate consisted of a Pum1 IP sample, a negative control IP sample, and a total RNA sample, which were each hybridized to a separate microarray along with a common reference sample. Total RNA microarrays were used to identify the transcriptome of the cells from which IPs were performed, providing an accurate background for subsequent analyses. A probe was considered present in the transcriptome if the signal from the spot was at least twofold greater than local background in all three total RNA or Pum1 IP microarrays. T scores, based on the t statistic, for Pum1 IP versus the negative control IP were calculated for all probes on the microarrays (49). A visual inspection of the t score values of the probes (Fig. 2, histogram) suggested two distributions: a background distribution and a Pum1-associated distribution.

In order to objectively define Pum1 target mRNAs based on a distribution of t scores, we employed Gaussian mixture modeling (GMM). GMM uses probabilistic modeling to identify single Gaussian distributions in a population consisting of a mixture of multiple Gaussian distributions (43). GMM uses expectation maximization modeling, which is prone to converging on a local optimum; therefore, several iterations of mixture modeling were performed which initialized at different values in the distribution, and the model with the greatest probability was then used to define Pum1-associated mRNAs. Both Pum1 and background distributions were defined as an equation relating t scores to probability (Fig. 2, curves), and using these equations we calculated the LOD for each probe appearing in the Pum1 distribution versus the background distribution. Those probes with a greater likelihood of being in the Pum1 distribution (LOD > 0) were considered Pum1-associated mRNAs (targets). While one might expect that this cutoff would result in a high false-positive rate, downstream analysis proved that a LOD cutoff of >0 was appropriate. A summary of probes, t scores, and LOD scores is available in Table S1 of the supplemental material. Use of GMM and creation of LOD scores allowed us to objectively define probabilities of probes being in the Pum1 distribution versus the background distribution and thus allowed for a less arbitrary determination of probes that could be considered Pum1 associated. Probes were collapsed into unique genes, and of the 6,539 unique genes represented in total, 726 (11.1%) were considered Pum1 targets on this basis.

We used RT-qPCR to confirm select targets by measuring their enrichment in the Pum1 IP versus either the negative IP or total RNA, using the nontarget message β2-microglobulin for normalization and GAPDH mRNA as a negative control. qPCR analysis confirmed levels of enrichment up to 240-fold for target messages in the Pum1 IP, with similar levels of enrichment versus either the negative IP or total RNA (Fig. 3). Target messages confirmed by RT-qPCR represented a range of LOD scores greater than zero, thus confirming these targets serve as partial validation of the LOD cutoff of >0 when defining Pum1-associated mRNAs, as noted above.

**Pum1 target mRNA analysis.** After defining mRNAs associated with the Pum1 RNP, we proceeded to determine if the proteins encoded by Pum1 target mRNAs were functionally related, as is predicted by the RNA regulon model (26, 28). We analyzed Pum1 target genes using two Web-based programs, Panther (37) and WebGestalt (62), which search gene lists for significant enrichment in gene ontology (GO) categories and other functional groupings. We also analyzed Pum1 targets using GSEA, which compares a rank-ordered gene list of interest to other gene sets in the GSEA Molecular Signatures database to discover significant correlations between sets of genes (49). Results from WebGestalt and Panther, as seen in Table 1, were very similar. The Panther program searches for both positively and negatively enriched categories and applies

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**FIG. 1.** A. IP-Western blot of Pum1, probed with anti-Pum1-specific antibody. In, input; Sup, supernatant. (Only −20% of the IP sample was loaded.) B. RT-PCR analysis of RNA recovered from Pum1 IP, Pum1 IP; Neg, negative IP; Tot, total RNA.

**FIG. 2.** A. Distribution of Pum1 IP versus negative IP t scores for three biological replicates of RIP-chip. Pum1-associated (solid black curve), background (dashed black curve), and the sum of Pum1-associated and background (gray curve) probability distributions are shown, as defined by Gaussian mixture modeling. B. Cyclin B1 and β-Actin RT-qPCR analysis.
a strict Bonferroni correction for multiple testing when calculating significance (37); thus, fewer categories were found to be enriched in the Panther analysis, and the statistical significance of that enrichment was lower. WebGestalt makes no correction for multiple testing; thus, the results obtained by this method, while having lower P values, may be less biologically significant. However, results obtained using either WebGestalt or Panther were in agreement. One of the more striking results of both the WebGestalt and Panther analyses was the large number of target genes and extreme significance of enrichment of GO categories involved in transcription and regulation of transcription. Another noteworthy result was the enrichment of mRNAs representing genes involved in regulation of the cell cycle and cell proliferation and differentiation, a result consistent with proposed ancestral functions of PUF proteins in stem cell biology (57). GSEA of the Pum1 IP rank-ordered gene list created from LOD scores revealed a high degree of correlation with various gene sets, including gene sets whose mRNA levels were found to decrease after UVC and UVB exposure, whose mRNAs increased after cytomegalovirus (CMV) infections, a gene set consisting of HOX genes, and gene sets related to the cell cycle (Table 1). This result provides further support for a role for the Pum1 protein in stem cell function, as well as a role for Pum1 in response to stress (see below).

Within the set of Pum1 target genes are several specific functional relationships that represent putative RNA operons/regulons, such as that between cyclin B1, Cdc2, p21, and Wee1. The cyclin B1-Cdc2 complex is a key regulator of the G2/M transition of the mitotic cell cycle, with p21 and Wee1 acting as negative regulators of Cdc2 (www.biocarta.com). Although it may seem counterintuitive that Pum1 would regulate expression of both one protein and a second protein that negatively regulates the first protein, this situation has been seen in C. elegans, where a PUF protein represses a mitogen-activated protein (MAP) kinase and a gene that inactivates the same MAP kinase, thereby ensuring continued repression of the MAP kinase gene after PUF-mediated repression of both proteins is relieved (31). PCNA, GSK3β, p21, and p27 form another cell cycle-related functional group, as all of these proteins function as inhibitors of cyclin D (www.biocarta.com).

One of the most striking potential Pum1 regulons is that of the E2F transcription factors: four of the five E2Fs that were represented in the total mRNA population were found to be Pum1 targets (E2F3 to E2F6 are targets, and E2F1 is not a target), showing an overlap of two highly enriched categories in Pum1 targets: cell cycle/proliferation and transcription (11). A large number of RNA processing and RNA-binding protein genes were also found to be Pum1 targets, among them the histone stem-loop-binding protein, DICER, Pum1 itself, and the other human PUF protein, Pum2.

**TABLE 1. Genes represented by Pum1-associated mRNA in various GO categories**

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<thead>
<tr>
<th>Analysis method and GO category (positively or negatively enriched)</th>
<th>No. of genes</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Panther Positively enriched mRNA transcription</td>
<td>100</td>
<td>9.66E-04</td>
</tr>
<tr>
<td>Panther Positively enriched mRNA transcription regulation</td>
<td>71</td>
<td>9.40E-03</td>
</tr>
<tr>
<td>Panther Positively enriched Transcription factor</td>
<td>103</td>
<td>3.02E-05</td>
</tr>
<tr>
<td>Panther Positively enriched Other transcription factor</td>
<td>32</td>
<td>4.42E-02</td>
</tr>
<tr>
<td>Panther Positively enriched Nucleoside, nucleotide, and nucleic acid metabolism</td>
<td>173</td>
<td>2.57E-02</td>
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<tr>
<td>Panther Positively enriched Cell proliferation and differentiation</td>
<td>51</td>
<td>1.76E-02</td>
</tr>
<tr>
<td>Panther Negatively enriched Oxidoreductase</td>
<td>5</td>
<td>9.90E-07</td>
</tr>
<tr>
<td>Panther Negatively enriched Reductase</td>
<td>0</td>
<td>1.15E-02</td>
</tr>
<tr>
<td>Panther Negatively enriched Dehydrogenase</td>
<td>2</td>
<td>2.85E-02</td>
</tr>
<tr>
<td>Panther Negatively enriched Electron transport</td>
<td>2</td>
<td>1.32E-02</td>
</tr>
<tr>
<td>WebGestalt Positively enriched Regulation of biological processes</td>
<td>220</td>
<td>1.17E-09</td>
</tr>
<tr>
<td>WebGestalt Positively enriched Regulation of transcription</td>
<td>138</td>
<td>5.48E-10</td>
</tr>
<tr>
<td>WebGestalt Positively enriched Transcription factor activity</td>
<td>67</td>
<td>1.10E-09</td>
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<tr>
<td>WebGestalt Positively enriched Regulation of cell cycle</td>
<td>42</td>
<td>2.71E-03</td>
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<tr>
<td>WebGestalt Positively enriched Ubiquitin cycle</td>
<td>30</td>
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<td>WebGestalt Positively enriched Wnt receptor signaling pathway</td>
<td>11</td>
<td>1.95E-03</td>
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<tr>
<td>WebGestalt Positively enriched Small GTPase-mediated signal transduction</td>
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<td>3.75E-04</td>
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<tr>
<td>WebGestalt Positively enriched Cell communication</td>
<td>128</td>
<td>2.13E-04</td>
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</table>

**GSEA**

<table>
<thead>
<tr>
<th>Analysis method and GO category (positively enriched)</th>
<th>No. of genes</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>GSEA Positively enriched UVC high all DN</td>
<td>117</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>GSEA Positively enriched UBV NHEK1 DN</td>
<td>66</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>GSEA Positively enriched CMV/HCMV time course all up</td>
<td>76</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>GSEA Positively enriched HOX genes</td>
<td>10</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>GSEA Positively enriched Cell cycle</td>
<td>28</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*For GSEA results, the numbers of genes reported represent core enrichment. The GO category names for GSEA are those used in the analysis software.
Identification of the Pum1 USER. Previous analyses of PUF protein target mRNAs revealed a consensus sequence present in the 3’UTRs of target messages (17, 18). We used the motif finding program MEME (5) to search for a consensus sequence in the 3’UTRs of Pum1 targets. The 3’UTR sequences of the top 100 Pum1-associated mRNAs, by LOD score, were used as a training set for MEME analysis, resulting in discovery of the consensus sequence shown in the inset in Fig. 4. Contained in the Pum1 target consensus sequence is the 8-nucleotide core sequence UGUAHAUA, which has been shown by X-ray crystallographic analysis to be directly bound by the Pum1 PUF-HD (54). We searched for the occurrence of this core sequence in the 3’UTRs of the Pum1 targets not used in the MEME training set and found it occurred at least once in 46.5% of Pum1 target 3’UTRs but only in 13.5% of total mRNA 3’UTRs. To determine the likelihood of this enrichment occurring by chance, we created 50,000 sets of random 3’UTRs and determined the frequency of 3’UTRs in each set.

FIG. 4. A. Calculated frequency of 3’UTRs containing the Pum1 USER, UGUAHAUA, among Pum1-associated mRNAs compared with 50,000 randomly chosen sets of mRNAs. P values derived from this sampling and Fisher’s exact test are shown. Random sets of mRNAs were derived from mRNAs present in the HeLa cell transcriptome, and each set contained the same number of mRNAs as the Pum1-associated set. Inset shows the consensus Pum1 target sequence, including the 8-nucleotide Pum1 USER, as determined by MEME analysis. B. GSEA of enrichment of the Pum1 USER based on LOD scores. Enrichment scores and the family-wise error rate (FWER) P value are shown. The vertical red line in the graphs represents LOD of 0.
that contained the core consensus sequence at least once. The random sets of 3'/UTRs were chosen from the mRNAs expressed in HeLa cells, and each set contained the same number of 3'/UTRs as the Pum1-associated set. The occurrence of the Pum1 core consensus sequence was determined for each of these sets, resulting in a distribution with a mean of 13.5% and a maximum of 20.1% (Fig. 4). This represents an extremely significant enrichment of the core Pum1 consensus sequence in Pum1 targets \( (P < 2 \times 10^{-5}) \) even after excluding those 3'/UTRs used as the training set. We also used Fisher’s exact test to determine the significance of enrichment of this sequence in Pum1 targets and calculated \( P \) to be 1.99E-60. As elements of this 8-nucleotide core sequence have been shown to be important for target mRNA regulation by PUF proteins \((6, 9, 19, 24, 31, 39)\), we will henceforth refer to this sequence as the Pum1 USER (untranslated sequence element for regulation) \((28)\).

Although it may be unexpected to find that only about half of Pum1 target messages contained the Pum1 USER, there are a number of likely explanations for this outcome. The RIP-chip method is optimized to isolate entire RNPs \((27, 50)\), and thus some of those messages associated with Pum1 RNPs are not expected be directly bound by Pum1. The search for the Pum1 USER was also based on a simple string \((UGUAHAUA)\) rather than a more descriptive and flexible position-specific weight matrix. Because the position-specific weight matrix more accurately describes flexibility in the consensus sequence, it could also identify sequences in 3'/UTRs to which Pum1 binds with almost as high an affinity as the consensus sequence which were not recognized by a simple string search. Finally, only the 3'/UTRs of mRNAs were searched for the Pum1 USER; thus, any USERs present in the 5'/UTRs or coding sequences would not be identified.

In order to more thoroughly explore enrichment of the Pum1 USER in Pum1-associated messages, we also used GSEA \((49)\). This analysis used all genes with the Pum1 USER in their 3'/UTR as a gene set and calculated the running enrichment of genes containing the Pum1 USER in the Pum1 RIP-chip data ordered by LOD score. As can be seen in Fig. 4B, the peak of the running enrichment score occurs after LOD of 0, demonstrating that our earlier LOD cutoff of \(> 0\) is valid for determining targets and is likely conservative. The normalized enrichment score of 14.23 and family-wise error rate \(P\) value of \(< 0.001\) for this analysis again demonstrate the extreme enrichment of this sequence in the 3'/UTRs of Pum1-associated mRNAs.

Role of Pum1 in decay of target messages. We next sought to determine whether Pum1 enhances decay of associated mRNAs, as has been shown for other PUF proteins \((19, 41)\). Thus, we knocked down Pum1 protein using siRNA and then assayed decay rates of target messages. Extent of Pum1 protein depletion was determined to be approximately 70 to 95% by Western blotting for all experiments. To determine mRNA decay rates, actinomycin D was used to inhibit transcription and qPCR was performed to determine the percentages of transcripts remaining at multiple time points after treatment, as normalized to GAPDH and averaged across three biological replicates. An exponential decay curve was fit to the mean of these measurements, and half-lives of messages were determined based on the equation describing this curve (Fig. 5). Control experiments were performed with an siRNA not known to target any mRNAs. All but one of the Pum1 target mRNAs assayed showed increased stability during Pum1 knockdown, although there was a range in the degree of increased stability. As Pum1 protein has been shown to interact with CNOT8 protein \((20)\), a member of the CCR4-NOT deadenylase complex, it will be interesting to determine in future studies whether Pum1’s effect on stability of associated mRNAs is at least partially mediated by deadenylation via this complex.

Subcellular localization of Pum1 protein following oxidative stress. To further study the function of Pum1, we used immunofluorescence analysis to determine its subcellular localization during cell growth and induced stress. Growing cells displayed a diffuse, granular cytoplasmic Pum1 staining pattern, and upon arsenite-induced oxidative stress Pum1 protein relocalized to large cytoplasmic granules \((Fig. 6, left panels)\). A blocking peptide was used to confirm specificity of the Pum1 antibody \((not shown)\). Stress granules and processing bodies \((p\)-bodies) are both candidate large cytoplasmic granules known to be induced by oxidative stress \((2)\); thus, to determine if these granules were stress granules, \(p\)-bodies, or both, we co-stained with a stress granule marker, HuR \((Fig. 6A, middle)\) or a \(p\)-body marker, Ddx6 \((Fig. 6B, middle panels)\). Co-staining confirmed that Pum1 protein was present in the same aggregates as HuR after arsenite treatment \((Fig. 6A, middle panel and zoom image)\), indicating that Pum1 protein is a component of stress granules. Pum1 also colocalized with exogenously transfected constructs expressing the recombinant fluorescent proteins G3BP-green fluorescent protein, TIA1-yellow fluorescent protein \((YFP)\), and FAST-YFP \((not shown)\), all of which are known components of stress granules \((25)\). Ddx6-containing \(p\)-bodies were sometimes juxtaposed but did not significantly overlap with Pum1 granules after induction of stress \((Fig. 6B, middle panel and zoom image)\), indicating that Pum1 is not a component of \(p\)-bodies and providing further evidence that Pum1 is a component of stress granules. Cells that had been transfected with a plasmid expressing Dcp1a-YFP protein, another \(p\)-body marker \((25)\), were also stained for Pum1, and again Pum1 granules appeared juxtaposed but distinct from Dcp1a-containing \(p\)-bodies \((not shown)\). Cells were also stained for Pum1 and HuR or Ddx6 after heat shock, with results again indicating that Pum1 is a component of stress granules but not \(p\)-bodies \((data not shown)\). The observation that Pum1 is a stress granule component suggests that it may have a role in regulating translation of target messages. Unlike the related protein Pum2 \((52)\), Pum1 knockdown did not interfere with stress granule formation after arsenite treatment, indicating that Pum1 is not necessary to form these structures in HeLa cells \((data not shown)\).

Conservation of PUF protein target mRNAs and potential RNA regulons. One of the main goals of this study was to compare Pum1 target genes to target genes of PUF proteins in other species in order to observe how the posttranscriptional regulatory adaptors of the PUF-HD and cognate binding sequence have been rewired throughout evolution. The four amino acids that directly contact RNA in each of the eight repeats of the PUF-HD \((54)\) are completely conserved between S. cerevisiae Puf3, Drosophila Pumilio, and human Pum1, as is the 8-nucleotide sequence to which each PUF-HD likely
FIG. 5. Decay rates of Pum1 target mRNAs, normalized to GAPDH, as determined by RT-qPCR after Pum1 knockdown followed by treatment with actinomycin D (ActD) to inhibit transcription. Black boxes represent Pum1 knockdown, and gray diamonds represent the control. Error bars represent standard errors of the means of three biological replicates.
binds (17, 18). Neither the RNA-contacting amino acids nor consensus binding sequence of any other yeast Puf protein show this level of conservation; thus, only Puf3 was considered in this analysis. mRNA targets of Puf3 and Pumilio have been previously determined using RIP-chip, and it was found that there was little conservation of target genes between them (17, 18). In order to determine if there is conservation of targets between Pum1 and Puf3 or Pumilio, we first determined how many Puf3 or Pumilio target genes have human orthologues that are expressed in HeLa cells and then determined how many of those genes with human orthologues are also Pum1 targets (Fig. 7A). Puf3 target genes have 89 human orthologues, and only 7 of these are also Pum1 target genes, representing no significant enrichment for Pum1 targets among Puf3 targets. Pumilio targets in the adult *Drosophila* ovary have 502 human orthologues, with 73 also being Pum1 target genes. This represents a slightly significant enrichment of Pum1 targets among Pumilio targets, with a sampling P value of 0.036. Statistical significance via sampling was determined by creating 10,000 sets of random genes containing the same number of genes as the Puf3 or Pumilio target genes with orthologues and then determining the proportion of those genes that were Pum1 targets. A possible caveat to this analysis is that each RIP-chip experiment was performed in a different cell type and thus each PUF protein had the potential to associate with different mRNAs in vivo. To address this issue we limited the universe of this analysis to only genes that were expressed in HeLa cells as defined by our study.

In addition to identities of genes encoded by target mRNAs, we also considered whether functional relationships among target genes, or potential RNA regulons (26, 28), were conserved (Fig. 7A). Target genes of Puf3 are almost entirely nuclear-encoded mitochondrial proteins (17), while those of Pumilio and Pum1 are not enriched for mitochondrial functions (18). The potential RNA regulon of cell cycle regulation is not conserved between Pumilio and Pum1, although there are many cyclin mRNAs among the Pumilio targets (18). Conversely, Pum1 targets are not enriched for components of the V-type ATPase or the embryo patterning cascade, functional groupings found to be enriched among Pumilio targets (8, 18). Pum1 targets and Pumilio targets are both enriched for the GO categories of transcription factor and membrane-bound organelle (18), showing that while most of the individual genes regulated by the proteins have changed, the types of genes regulated are similar. Combined with data regarding conservation of individual targets, this observation indicates that neither targets nor functions of PUF proteins have been conserved between *S. cerevisiae* and humans, yet some targets and functions have been conserved between *Drosophila* and humans. Thus, by keeping the PUF-HD and cognate binding sequence as static modules but changing the sets of mRNAs that are regulated, evolution has been able to rewire posttranscriptional regulation of gene expression networks, as previously suggested through experimental and computational analysis of evolutionary conservation of potential posttranscriptional regulatory elements (8, 18, 21, 26, 36).

**DISCUSSION**

While various genetic and biochemical studies have revealed mRNA targets and functions of PUF proteins in nonmamma-
lian model organisms, little is currently known about the interactions or functions of PUF proteins in mammals. This study represents both the first in vivo genome-wide mRNA target identification of a mammalian PUF protein and the first direct demonstration of a function of a mammalian PUF protein in regulating its target mRNAs. By comparing targets of PUF proteins in three organisms, we were also able to observe how mRNA targets of these proteins have changed through evolution, despite conserved cis-trans interactions between the PUF-HD and cognate binding sequence.

**Advantages of the RIP-chip method.** The RIP-chip method used in this study is a biochemical convention that allows the recovery of endogenous RNP complexes, which are more physiologically relevant than a single RBP bound to a single RNA. Isolation of entire RNPs also allows the identification of other components of these RNPs, including microRNAs (27). In addition, the RIP-chip method allows the study of RNPs during dynamic situations where RNP constituents may be changing, possibly resulting in different functional outcomes. RIP-chip is a relatively simple method with optimized conditions that provide an environment where all of the components necessary for an RBP’s binding and function are present.

Although the RIP-chip procedure used in this study was similar to that used in previous studies, some differences are noteworthy. After determining t scores of Pum1 IP versus negative IP for all of the probes on the microarrays, we used Gaussian mixture modeling to objectively determine the two distributions representing the Pum1-associated and background messages. GMM allowed us to make a more sophisticated, probabilistic determination of which probes were Pum1 associated and which represented background. By calculating probabilities of association, this method allows for more advanced downstream analysis of target mRNAs, as probabilities of association can be used rather than a simple discrete definition of which mRNAs are targets. For example, GSEA would not have been possible by simply defining which mRNAs were targets, because it depended on the continuous metric of a rank-ordered list of mRNAs created from LOD scores.

**Pum1 and the RNA regulon model.** The observation that many of the mRNAs associated with Pum1 belong to a relatively small number of functional groupings is consistent with the RNA operon/regulon model (22, 26, 28). This model describes how multiple genes with related functions can be coordinately regulated at the level of the mRNA. Indeed, many aspects of the RNA regulon model are reflected in the results from this study. For example, one key aspect is that posttranscriptional regulation at the level of the mRNA accommodates the multifunctionality of eukaryotic proteins by allowing a single gene to participate in multiple regulons (cis-combinatorial). Thus, although Pum1 targets are enriched for regulators of progression through the cell cycle yet we observed no function for Pum1 in cell cycle progression (data not shown), it is likely that these genes also function in processes that are not related to the normal progression of the cell cycle, and these alternate processes may in fact be affected by Pum1 perturbation. It will be interesting to determine whether Pum1 functions in the meiotic cell cycle, especially since PUF proteins are known to function in the germ line of various organisms, including mouse (33, 58, 61). The cell proliferation- and differentiation-related targets of Pum1 may reflect a role for Pum1 in stem cell biology, a proposed ancestral function of PUF proteins (57). Another key aspect of the RNA regulon model is that regulation of mRNAs can be trans-combinatorial, with multiple trans factors specifying the fate of a single mRNA.

There is precedent for this combinatorial trans-regulation in yeast, where several mRNAs have been shown to be regulated by multiple PUF proteins (23, 51). This combinatorial regulation provides an explanation for the range of effects of Pum1 knockdown on target message stability, as each target message is likely regulated by various other RBPs and microRNAs. Although the level of Pum2 protein in HeLa cells is relatively low compared to some other commonly used cell lines (not shown), it is possible that Pum2 may act redundantly on some or all of the Pum1 target mRNAs, thus dampening the effect on stability seen during Pum1 knockdown. It is also likely that not every copy of a Pum1 target mRNA will be bound by a Pum1 protein, especially since Pum1 associates with mRNAs representing over 10% of expressed genes. Thus, only a portion of each Pum1 target will be stabilized by Pum1 knockdown, since not every copy would normally have been regulated by Pum1. The RNA regulon model also states that RNPs may be dynamic with changing cellular conditions, and thus it will be of interest to determine how stability and translation of Pum1 target messages change following cell stress, especially since we have observed a change in the localization of Pum1 protein upon both oxidative stress and heat shock. There is precedent for this condition-specific regulation in yeast, where Puf3 differentially regulates stability of target messages when the yeast are grown on different carbon sources (13). It will also be important to determine how the targets of human Pum1 and Pum2 change during development and stem cell maturation, processes known to involve PUF proteins in *Drosophila* (32, 33) and *C. elegans* (29).

**Evolutionary rewiring of PUF-HD and PUF-USER.** Previous studies of RBPs that bind AU-rich elements were unable to draw strong conclusions regarding conservation of RNA-protein interactions among species for two main reasons. First, the RBPs in question bind RNA through the widely represented RRM motif, and thus true orthologues between species were difficult to discern (1). Second, the sequences to which AU-rich element RBPs bind are not generally unique and involve elements that use both sequence and structure (34). Neither of these problems presents itself when considering PUF proteins. The PUF-HD is extremely well-conserved across species, yet each species has few PUF genes; mammalian genomes contain only two (47). The sequence to which the PUF-HD binds is also very well-conserved, with a UGUAR motif identified in most PUF-binding sites and the 8-nucleotide core motif, UGUARAUAA, almost identical between Puf3, Pumilio, and Pum1 (17, 18, 54). Comparison of target genes of these three proteins revealed that they likely regulate different processes; Puf3 binds messages of genes with mitochondrial function (17), but there is no enrichment for genes with mitochondrial functions among targets of Pumilio (18) or Pum1. Targets of Pumilio and Pum1 do not share specific functional relationships, and although there is some conservation of target genes, most of the Pum1 targets and Pumilio targets are different (Fig. 7). These observations show that the modules of the PUF-HD and cognate binding sequence have remained fixed through evolution, while the iden-
ties of target messages have changed phylogenetically. This study provides experimental evidence of a rewiring process that was predicted through analysis of conservation of potential postranscriptional regulatory elements, showing how a conserved cis-trans interaction can be evolutionarily rewired to coordinate the expression of different subsets of genes in different species (8, 18, 21, 26, 36).

**Pum1 as a regulator of regulators.** Many of the genes whose mRNAs are bound by Pum1 are known regulators of gene expression and cellular processes, and thus Pum1 could be described as a regulator of regulators (26, 36, 44). The GO category containing the most Pum1 targets is “regulation of biological processes,” and target genes of Pum1 contain genes that regulate gene expression at the transcriptional, postranscriptional, and posttranslational levels. GO analysis revealed that transcription factor genes and ubiquitin cycle genes are enriched in Pum1 targets, and many of the target genes involved in nucleic acid metabolism encode proteins that bind and process RNA. Pum1 targets are also enriched for GTPase-mediated signal transduction genes and other genes involved in signaling pathways that themselves could be described as regulatory, as their activation or repression typically results in changes in gene expression. Thus, Pum1 could be described as a regulator of regulators, because it associates with genes that regulate multiple levels of gene expression, as well as genes encoding members of signaling pathways that trigger changes in gene expression.

Pum1’s function as a regulator of regulators is also evident when observing genes that are not represented in Pum1 targets. The GO categories of electron transport and oxidoreductase were significantly depleted among Pum1 targets, indicating that Pum1 does not regulate mRNAs involved in processes that are not regulatory. Pum1 targets did not show significant enrichment of any GO categories involving metabolism (other than nucleic acid metabolism, which is related to transcription and RNA processing), again demonstrating the lack of a role for Pum1 in nonregulatory processes and supporting its role as a regulator of regulators.

**Importance of Pum1 target mRNA identification for study of gene expression.** Posttranscriptional regulation of gene expression has been increasingly studied in recent years, although global aspects have not been widely explored. Study of PUF family RNA-binding proteins has shown how these proteins can regulate a variety of subsets of mRNAs in a variety of organisms, with results typically supporting the proposed ancestral role of PUF proteins in stem cell biology (57). Due to the likely functions of PUF proteins in human stem cells, it is of interest to determine the role of human PUF proteins in regulating gene expression. This work serves to vastly expand the body of knowledge concerning PUF protein mRNA targeting and function in mammals, which will aid future studies in elucidating how PUF proteins act to regulate gene expression at the postranscriptional level, particularly in stem cell biology, growth, and differentiation.

**ACKNOWLEDGMENTS**

We thank Nancy Kedersha for generous contribution of constructs encoding stress granule and p-body markers and intellectual input, Blanche Capel and her group for assistance with immunofluorescence, Uwe Ohler and William Majors for access to the 3’UTR database, Sayan Mukherjee for valuable advice on statistical analyses, and Shelton Bradrick and Keene lab members for helpful comments regarding the manuscript.

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