# Identification of *Drosophila* MicroRNA Targets

Alexander Stark®, Julius Brennecke®, Robert B. Russell, Stephen M. Cohen\*

European Molecular Biology Laboratory, Heidelberg, Germany

MicroRNAs (miRNAs) are short RNA molecules that regulate gene expression by binding to target messenger RNAs and by controlling protein production or causing RNA cleavage. To date, functions have been assigned to only a few of the hundreds of identified miRNAs, in part because of the difficulty in identifying their targets. The short length of miRNAs and the fact that their complementarity to target sequences is imperfect mean that target identification in animal genomes is not possible by standard sequence comparison methods. Here we screen conserved 3' UTR sequences from the Drosophila melanogaster genome for potential miRNA targets. The screening procedure combines a sequence search with an evaluation of the predicted miRNA-target heteroduplex structures and energies. We show that this approach successfully identifies the five previously validated let-7, lin-4, and bantam targets from a large database and predict new targets for Drosophila miRNAs. Our target predictions reveal striking clusters of functionally related targets among the top predictions for specific miRNAs. These include Notch target genes for miR-7, proapoptotic genes for the miR-2 family, and enzymes from a metabolic pathway for miR-277. We experimentally verified three predicted targets each for miR-7 and the miR-2 family, doubling the number of validated targets for animal miRNAs. Statistical analysis indicates that the best single predicted target sites are at the border of significance; thus, target predictions should be considered as tentative until experimentally validated. We identify features shared by all validated targets that can be used to evaluate target predictions for animal miRNAs. Our initial evaluation and experimental validation of target predictions suggest functions for two miRNAs. For others, the screen suggests plausible functions, such as a role for miR-277 as a metabolic switch controlling amino acid catabolism. Cross-genome comparison proved essential, as it allows reduction of the sequence search space. Improvements in genome annotation and increased availability of cDNA sequences from other genomes will allow more sensitive screens. An increase in the number of confirmed targets is expected to reveal general structural features that can be used to improve their detection. While the screen is likely to miss some targets, our study shows that valid targets can be identified from sequence alone.

#### Introduction

MicroRNAs (miRNAs) are small RNAs, typically of approximately 21-23 nt, that direct posttranscriptional regulation of gene expression by binding to messenger RNAs (mRNAs). Many endogenously encoded miRNAs have been cloned from plants and animals (Lagos-Quintana et al. 2001, 2002; Lau et al. 2001; Lee and Ambros 2001; Mourelatos et al. 2002; Reinhart et al. 2002; Ambros et al. 2003; Aravin et al. 2003; Lim et al. 2003). Combining these data with computational cross-genome comparison predicts 100-120 miRNA-encoding genes in Caenorhabditis and Drosophila and approximately 250 in mouse and human (Ambros et al. 2003; Grad et al. 2003; Lai et al. 2003; Lim et al. 2003a, 2003b). However, functions have been assigned to only four animal miRNAs (Reinhart et al. 2000; Brennecke et al. 2003; Lee et al. 1993; Wightman et al. 1993; Xu et al. 2003), in part owing to the difficulty in identifying mutations in such small genes. A method to identify the target genes that are regulated by miRNAs would greatly help the study of miRNA function in animals (Ambros 2001).

Two modes of miRNA-directed target inhibition have been demonstrated. The same small RNA can cause degradation of its target mRNA or block its translation depending on the degree of miRNA-target sequence complementary (Hutvagner and Zamore 2002; Doench et al. 2003). Target RNAs containing sequences with perfect complements of the miRNA (or exogenously supplied short interfering RNA [siRNA]) are cleaved by the RNA-induced silencing complex (RISC) ribonuclease (Hutvagner and Zamore 2002; Martinez et al. 2002; Zeng et al. 2002). Endogenous plant miRNAs have been shown to regulate target RNAs by RNA interference

(RNAi) involving perfect or near-perfect target site complementarity (Llave et al. 2002b; Kasschau et al. 2003; Palatnik et al. 2003; Tang et al. 2003; Xie et al. 2003). Targets for plant miRNAs have been identified on a genome-wide scale by searches that require a high degree of sequence complementarity (Rhoades et al. 2002). However, this approach did not find targets for known animal miRNAs. The animal miRNAs tested until now pair imperfectly with their targets and act to control translation. Indeed, systematic analysis of the complete *C. elegans* miRNA complement has confirmed the absence of targets with perfect or near-perfect sequence complementarity (Ambros et al. 2003).

To date targets have been experimentally validated for just three animal miRNAs: the *lin-4* targets *lin-14* and *lin-28* (Wightman et al. 1993; Ha et al. 1996; Moss et al. 1997; Olsen

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**Abbreviations:** bHLH, basic helix-loop-helix; CDS, coding sequence; domT, domain bit score threshold; EGFP, enhanced green fluorescent protein; *E(spl), Enhancer of split* gene; GFP, green fluorescent protein; kb, kilobase; miRNA, microRNA; mRNA, messenger RNA; nt, nucleotides; ORF, open reading frame; ptc–Gal4, yeast GAL4 transcriptional activator expressed under control of the *Drosophila patched* gene promoter; RISC, RNA-induced silencing complex; RNAi, RNA interference; SD, standard deviation; siRNA, short interfering RNA; 3' UTR, untranslated region of a messenger RNA following the coding sequence

Academic Editor: Ronald H. A. Plasterk, Utrecht University

\*To whom correspondence should be addressed. E-mail: cohen@embl.de  $\,$ 

These authors contributed equally to this work



and Ambros 1999; Seggerson et al. 2002), the let-7 targets lin-41 and lin-57/hbl-1 (Reinhart et al. 2000; Slack et al. 2000; Abrahante et al. 2003; Lin et al. 2003), and the bantam target hid (Brennecke et al. 2003). These miRNA-target duplexes contain mismatches, gaps, and G:U basepairs at different positions. Even allowing for G:U basepairs, the longest contiguous alignments in these examples range from 8 to 10 nt. Such limited information content makes it difficult to identify targets within whole-genome or transcriptome databases, since standard alignment methods produce many false positives with such short variable sequences. Furthermore, the small number of validated examples makes the development and benchmarking of a generally applicable computational method problematic at present. Here we present a screen for miRNA targets in Drosophila that identifies all of the previously known miRNA targets and we demonstrate that it successfully predicts new targets that we validate experimentally.

#### Results

## Database Design

For each of the validated miRNA-target pairs, functional target sites are located in the 3' untranslated region (UTR) of the mRNA and are conserved in the 3' UTRs of the homologous genes from related species (Wightman et al. 1993; Moss et al. 1997; Pasquinelli et al. 2000; Brennecke et al. 2003). We used pairwise comparison of the 3' UTRs of orthologous genes in related genomes to identify conserved 3' UTR sequences. Figure 1A shows the resulting pattern of 3' UTR conservation for the known targets in worms and flies. The vast majority of miRNA target sites (red bars in Figure 1A) are located in blocks of conserved sequence (white blocks in Figure 1A). Figure 1B shows cross-genome conservation of these miRNA target sites. A striking pattern of uninterrupted conservation emerges at the end of the target sequences that pair with the 5' end of the miRNAs.

To permit genome-wide searches for targets of Drosophila miRNAs, a conserved 3' UTR database was prepared by comparison of Drosophila melanogaster and Drosophila pseudoobscura 3' UTRs. As very few 3' UTRs are defined by cDNA sequence data in D. pseudoobscura, we used genomic sequence following the last exon of the D. pseudoobscura gene as the orthologous UTR (see Materials and Methods). Last exons were reliably detected in D. pseudoobscura for approximately two-thirds of *D. melanogaster* genes. On average, 22% of the *D*. *melanogaster* 3' UTR sequence is conserved in the predicted *D*. pseudoobscura 3' UTR. Much of this reflects isolated blocks of very high conservation interspersed among less-conserved sequence. Use of conserved 3' UTRs reduces the expected number of sequence matches that would occur at random by 4- to 5-fold in relation to full-length 3' UTRs, and severalfold further compared to the full transcriptome. We considered using the Anopheles gambiae genome to extend the crossspecies' comparison. Although genome annotation identifies orthologs for two-thirds of D. melanogaster genes (Zdobnov et al. 2002), we were unable to identify the last D. melanogaster exon for approximately half of these. We therefore chose not to require conservation in Anopheles, but use it as an additional level of validation for predicted Drosophila targets, where possible.

## Screening Strategy

We have adopted a two-step approach to target identification that combines a sensitive sequence database search with an RNA folding algorithm to evaluate the quality of the RNA duplex formed between the miRNA and its predicted targets. We examined the known target sites for lin-4, let-7, and bantam for common features. All of these sites showed better complementarity to the 5' end of the miRNA, with no obvious common features elsewhere (Figure 2A and 2B). There were few sequence mismatches or G:U basepairs in the alignment of the first eight residues at the 5' end of the miRNA. We used the alignment tool HMMer (Eddy 1996) to search for sequences complementary to the first eight residues of the miRNA, allowing for G:U mismatches. Where possible, the corresponding sites were also identified in the D. pseudoobscura 3' UTR, and the sites from both genomes were considered, since the regions outside of the sequence match can vary between the two organisms, leading to differences in subsequent steps (see below).

The identified sequences were extended to the length of the miRNA plus five residues to allow for bulges and were evaluated for their ability to form energetically favorable RNA–RNA duplexes with the miRNA using Mfold, which combines knowledge of known RNA structures with thermodynamic parameters, such as those involved in basepairing to evaluate the free energy of folding ( $\Delta G$ ) (Mathews et al. 1999; Zuker et al. 1999). Mfold requires a single linear sequence as input, so each predicted target was linked to the miRNA using a standard hairpin-forming linker sequence (GCGGGGAC-GC). An example of the Mfold output is shown in Figure 2C for the top-scoring bantam miRNA target site that we had previously identified in the 3' UTR of hid (Brennecke et al. 2003).

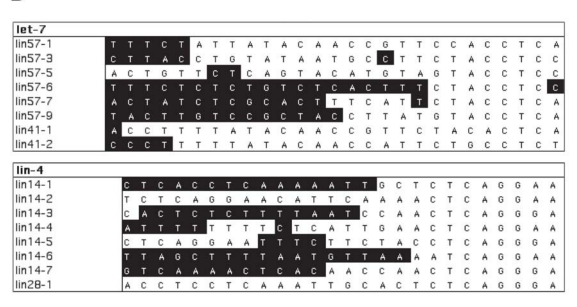
The Mfold free energy of folding ( $\Delta G$ ) was determined for each predicted target, which allows predicted sites to be ranked according to  $\Delta G$ . However,  $\Delta G$  depends on miRNA length and GC content, so it is not possible to distinguish systematically real targets from random matches using the raw  $\Delta G$  score or to compare different miRNAs. Instead, we calculated Z-values as a measure of nonrandomness, with an average random site scoring Z=0 (Z= standard deviations [SD] above the mean of background matches). Figure 2D shows the distribution of folding energies for predicted targets of the *bantam* miRNA compared to 10,000 randomly selected target sequences.

Most of the previously validated targets have more than one predicted miRNA-binding site in their 3' UTRs. Use of the Z-value allows us to add the scores of several sites within one UTR by selecting only those scores that are different from background matches. This is not possible with  $\Delta G$  alone because even average random matches have favorable energy values (Figure 2D) and the sum of several average random matches in a UTR could score better than a single true site. We have selected  $Z \ge 3$  as a cutoff value, as folding energies of more than 3 SD above the mean ( $Z \ge 3$ ) are expected to occur for only 0.3% of random matches. Use of a higher Z-value increases the likelihood that predictions are correct, but also increases the risk of missing out contributions from real sites of lower folding energy. For example, only three of the five conserved bantam sites previously identified in the hid 3' UTR score  $Z \ge 3$  (with the best site at Z = 7.4). We evaluated our





## В



bantam																												
hid-1	G	Т	Т	С	Α	Т	С	Α	Т	С	A	Т	Α	Т	Т	С	Α	Α	Α	Т	Т	G	G	Т	С	Т	С	Α
hid-2	Т	Т	Т	Т	Т	Т	G	G	Α	A	Т	G	С	Α	С	Α	Т	Т	A	Α	Т	G	A	Т	С	Т	C	Т
hid-3	С	С	Α	Α	Т	Т	С	С	С	Α	Α	Α	Α	Α	Т	С	G	С	Α	Т	Т	G	Α	Т	С	Т	C	Α
hid-4	Т	Т	G	С	Т	Α	Α	Т	Т	Α	G	Т	Т	Т	Т	С	Α	C	Α	Α	Т	G	Α	Т	С	Т	С	G
hid-5	A	Т	Α	Т	A	С	Α	Т	Α	Α	Α	Т	Α	Т	С	Α	Т	Т	A	Т	Т	G	A	Т	С	Т	С	Α

Figure 1. Features of Known miRNA Target Sequences

(A) Comparison of sequence conservation in the 3' UTRs of miRNA target genes. For lin-14, lin-28, lin-41, and lin-57, comparison was between C. elegans and C. briggsae. For hid, comparison was between D. melanogaster and D. pseudoobscura. White regions indicate conservation, and black regions are not conserved under the conditions used for producing the 3' UTR database (see Materials and Methods for details). The positions of predicted miRNA target sites from the literature are shown in red. Most of these UTRs contain multiple predicted target sequences, and while regulation of the UTR has been experimentally validated in each case, most individual sites have not been tested for function.

(B) Detailed comparison of the pattern of sequence conservation in the conserved sites. Target site length is miRNA length plus 5 nt. For *lin-57* we excluded three of the eight previously predicted sites that were not located in conserved sequence blocks and included a newly identified ninth site that is conserved. White type on black indicates residues that are not identical in the target sites in the two genomes. Black type on white indicates identity. All residues basepairing with positions 2–8 of the miRNA are identical in the conserved sites in both genomes. DOI: 10.1371/journal.pbio.0000060.g001



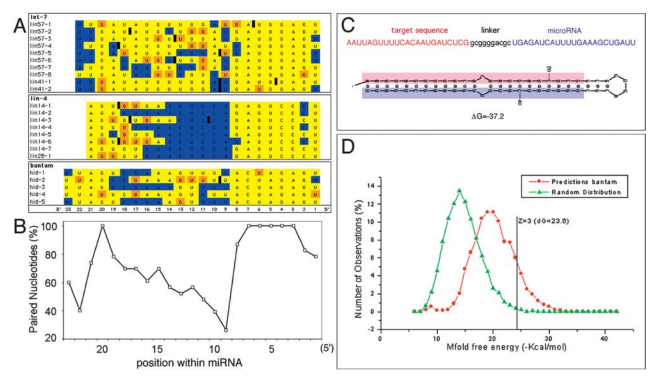


Figure 2. miRNA Target Prediction Strategy

(A) let-7, lin-4, and bantam miRNA sequences showing the pattern of basepairing to their known targets. Yellow indicates a conventional basepair. Orange indicates a G:U basepair. Blue indicates a mismatch. The black bars indicate the positions of loops in the target sequence. Note that the extra bases that form the loops in the target sequences are not shown. Sequences are shown at the length of the miRNA.

(B) Quantitation of the data from (A). This comparison shows that the 5' ends of the miRNA are always involved in good pairing with target sequences and suggests that searches for complementarity to the first eight residues of the miRNA would select all known targets.

(C) Graphic representation of the Mfold output for the bantam miRNA and a target site from the 3' UTR of the hid gene. To use Mfold, it is necessary to join the predicted target site (red) and the miRNA (blue) into a single sequence using a hairpin-forming linker sequence. In this example, the target sequence and the miRNA are the same length, so the additional 5 nt in the tail of the predicted target sequence are not shown. (D) Plot of the Mfold free energy distribution for 10,000 random sequences (green) and for predicted targets of the bantam miRNA (red). X-axis:  $\Delta G$  calculated for each site by Mfold.

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predictions by the best single site in the 3' UTR ( $Z_{max}$ ) and by the sum of sites with  $Z \ge 3$  ( $Z_{UTR}$ ).

#### Tests with Previously Validated Targets

Table 1 summarizes the performance of the method in predicting the known targets of the C. elegans miRNAs lin-4 and let-7 and the Drosophila miRNA bantam. The Drosophila hid gene ranked first of all predicted bantam targets sorted by the single best site  $(Z_{\text{max}})$  or by the sum of sites  $(Z_{\text{UTR}})$ . All of the known targets of lin-4 and let-7 were found when their 3' UTRs were added to the *Drosophila* 3' UTR database. Like *hid*, the let-7 target lin-57 ranked near the top of the list by both measures. With several sites predicted of  $Z \ge 3$ , lin-57 ranked first by  $Z_{\text{UTR}}$ . Its best single site ranked in position 2 (Z = 6.8). C. elegans lin-14 was predicted to contain multiple lin-4 sites (Wightman et al. 1993). Three of these scored  $Z \ge 3$ . lin-14 ranked first when the list of predicted lin-4 targets was sorted for Z<sub>UTR</sub>, although the best single site in lin-14 placed it in position 20 (Z = 4.3). The *lin-4* target *lin-28* and the *let-7* target lin-41 ranked highly when the lists were sorted for the best single site, but ranked lower when multiple sites were summed because they had few high-scoring sites. The Drosophila homolog of lin-41, dpld, also ranked high among let-7 targets (Z = 5.6; see below). We compared our results with previous target predictions from the literature that have not been experimentally validated (Table 1). Our screen supports some of them (e.g., let-7 regulating lin-14), but we consider others unlikely because they rank very low on their lists or have no sites of  $Z \geq 3$  (e.g., let-7 and lin-28 or miR-4 and m4). The predicted miR-14 target Drice (Xu et al. 2003) is unlikely to be valid because the site is not conserved in the predicted Drice 3' UTR from D. pseudoobscura.

This analysis shows that all known targets were detected and ranked among the top-scoring predictions in genome-wide searches. This suggests that other valid targets should rank among the small number of best predictions that can be tested experimentally. Of particular interest were three miRNAs for which we predicted clusters of functionally related targets: miR-7, the miR-2 family, and miR-277 (Table 2; Table 3). Clustering of top-scoring sites in a group of related genes is likely to be significant when it arises from an unbiased genome-wide analysis. miR-7 and miR-2 were selected for target validation.

## miR-7 Regulates Notch Targets

Among the top 10 predictions for *miR*-7, we found *Enhancer* of split (*E*(spl)) and *Bearded* (*Brd*) complex genes (Figure 3A). *HLHm3* encodes a basic helix-loop-helix (bHLH) transcrip-



Table 1. Assessment of Predictions for Known and Predicted miRNA Targets

miRNA-Target Pair	$\Delta G$	$Z_{ m Max}$	Rank $Z_{\text{Max}}$	$\#Z \geq 3$	Rank Z <sub>UTR</sub>	References
Confirmed Dain						
Confirmed Pairs						
lin-4/lin-14	-29.9	4.3	20	3	1	Wightman et al. (1993); Ha et al. (1996)
lin-4/lin-28	-30.9	4.6	8	1	15	Moss et al. (1997)
let-7/lin-41	-32.3	6.4	3	2	20	Reinhart et al. (2000);
						Slack et al. (2000)
let-7/lin-57 (hbl-1)	-33.4	6.8	2	14	1	Abrahante et al. (2003);
						Lin et al. (2003)
bantam/hid	-37.4	7.4	1	3	1	Brennecke et al. (2003)
Predicted Pairs						
lin-4/lin-41	-28.9	4.0	32	1	36	Slack et al. (2000)
lin-4/lin-57	-21.6	1.7	361	0	_	Abrahante et al. (2003);
						Lin et al. (2003)
let-7/lin-14	-35.1	7.2	1	13	2	Reinhart et al. (2000)
let-7/lin-28	-20.6	2.8	861	0	_	Moss and Tang (2003)
miR-13a/hb	_	_	_	0	_	Abrahante et al. (2003)
miR-4/hb	_	_	_	0	_	Abrahante et al. (2003)
miR-3/hb	_	_	_	0	_	Abrahante et al. (2003)
miR-11/HLHm8	-29.4	4.7	27	1	46 (predicted UTR)	Lai (2002)
miR-4/m4	-21.5	2.1	272	0	<u> </u>	Lai (2002)
miR-7/HLHm3	-37.3	7.0	2	1	16	Lai (2002)
miR-7/Tom	-34.5	6.1	5	2	1	Lai (2002)
miR-14/Drice	_	_	_	0	(Site not conserved)	Xu et al. (2003)

Confirmed pairs indicates experimentally validated target 3' UTRs.  $\Delta G$ ,  $Z_{Max}$ , and  $Z_{UTR}$  are defined in the text. Predicted pairs indicates examples predicted in the literature for which there was no experimental validation. The let-7/lin-14 pair ranks very high on the list of let-7 predictions and is likely to be a functional target. The lin-4/lin-41 pair requires experimental validation. The other C, elegans predictions cannot be distinguished from random matches. The 5' end of the K box shows sequence complementarity to the miR-2/miR-13 family and to miR-6 and miR-11 (Lai 2002). The prediction of HLHm8 as a target for miR-11 seems plausible (using predicted UTR), as do the two miR-7 GY box-based predictions. None of the conserved sites predicted for Drosophila hunchback (Abrahante et al. 2003) were on our lists because of interrupted 5' alignments. DOI: 10.1371/journal.pbio.000060.t001

tional repressor; *Tom* and *m4* encode Brd family proteins. The bHLH repressor *hairy* was also among the top 10. These sites were conserved in the orthologous genes from *Anopheles*, when those could be identified. This prompted us to examine all the genes in *E(spl)* and *Brd* complexes for *miR-7* sites. We found possible target sites in many of them. Alignment of these sites showed a pattern of 5' end conservation quite similar to that for validated targets, with no mismatches and few G:U basepairs for about half of these genes (Figure 3B).

To assess the validity of some of the predicted targets, transgenic flies expressing the miR-7 miRNA and several sensor transgenes were prepared. A genomic fragment containing the miR-7 hairpin was cloned into the 3' UTR of a UAS-DSRed2 plasmid to allow GAL4-dependent expression of miR-7. The 3' UTRs of HLHm3, m4, and hairy were cloned into a tubulin promoter-EGFP (enhanced green fluorescent protein) reporter plasmid. As a control, a specific miR-7 sensor transgene was produced by cloning two copies of a perfect complement of the *miR*-7 miRNA sequence into the 3' UTR of the tubulin promoter-EGFP reporter. The miR-7 sensor was expressed uniformly in the wing imaginal disc. GAL4-dependent expression of miR-7 miRNA reduced expression of miR-7 green fluorescent protein (GFP) sensor transgene (Figure 3C). As the target sites in the sensor construct are perfect complements of the miR-7 miRNA, this is expected to be due to RNAi. GAL4-dependent expression of *miR*-7 also reduced expression of the *m4* 3' UTR sensor transgene (Figure 3D). The *miR*-7 site in *m4* is identical in *D. pseudoobscura* and conserved in *Anopheles*.

Expression of *miR-7* also caused a clear downregulation of the *hairy* 3' UTR sensor transgene, although its overall level of expression was lower in the wing disc (Figure 3E). The *hairy* gene has been cloned and cDNAs sequenced from two additional insect genomes: the flour beetle *Tribolium castanaeum* and *Drosophila simulans*. The predicted *miR-7*-binding site is conserved in these genomes, as well as in *Anopheles*, and shows striking conservation of alignment at the 5' and 3' ends of the predicted miRNA-binding site (Figure 3F). The level of expression of the *HLHm3* 3' UTR sensor was too low to be reliably studied, but also showed regulation by *miR-7*. Again, the *miR-7* site in *HLHm3* is identical in *D. pseudoobscura*. These observations validate the utility of the screen in predicting new miRNA targets.

To assess *miR*-7 function in vivo, we examined wings in which *miR*-7 was overexpressed under ptc-Gal4 control. Notching of the wing margin was observed (Figure 3G), which is characteristic of reduced *Notch* signaling (de Celis and Garcia Bellido 1994; Diaz-Benjumea and Cohen 1995; Rulifson and Blair 1995; Micchelli et al. 1997). The *Notch* target *cut* was expressed at reduced levels in the *miR*-7-



Table 2. Top Ten Predictions for miR-7 and miR-2a

niR-7	$\Delta G$	$Z_{\rm max}$	# Z ≥ 3	$Z_{\rm UTR}$	Gene	Alignment	Flags	$Ag~Z \geq$
1	-38.7	7.47	1	7.47	CG14989-RB	ACAGCAGAAUCACGC-AGGG-CUUCCA UGUUGUUUUAGUGAUCAGAAGGU ***+**	_	+
2	-37.3	7.03	1	7.03	HLHm3	GCAACAAGAUCCGUUGUCUUCCA GUUGUUUUAGUGAUCAGAAGGU +************************************	-	NF
3	-35.3	6.39	1	6.39	CG17657-RA	ACAACGGUUAAG CGCUGCGUCUUCCA UGUUGUU UUAGUGAU - CAGAAGGU	-	NF
4	-35.0	6.29	1	6.29	hairy	*****++ ACAGCAAAUCAGCAAAAGUCUUCCA UGUUGUUUUAGU-GAUCAGAAGGU	-	+
5	-34.5	6.13	2	11.78	Tom	***+**** -UAGCC - GAAUCAUU - GUCUUCCA UGUUG - UUUUAGUGAUCAGAAGGU	-	+
6	-33.9	5.94	1	5.94	hep	-+*+*+*****+*-**********************	5' cons	NF
0	-33.9	3.54	,	3.54	Пер	UGUUGUU - UUAGUGAUCAGAAGGU +**+***-*+***	CDS+	IVI
7	-33.8	5.91	1	5.91	CG8944-RA	ACGACAAGAUCAAGCGCUACGUCUG-CCA UGUUGUUUUAGUGAU-CAGA-AGGU **+***+***+***	5' helix CDS+	NF
8	-33.1	5.68	1	5.68	CG10540-RA	- CGACAAAGCG GCCCAAUAGUCUUCCA UGUUGUUUU AGUG AUCAGAAGGU - * + * * * * * + + * * * * *	_	_
9	-31.8	5.27	1	5.27	CG10444-RA	GCGACC - AAAA - CAG AGUCUUCCA		NF
9	-31.6	3.27	'	3.27	CG10444-NA	UGUUG-UUUU-AGU-GAUCAGAAGGU +*+********	_	INF
10	-31.5	5.17	1	5.17	m4	- CAGCUUU AAUCAAC GUCUUCCG UGUUG UUUUAGU GAUCAGAAGGU - * * + * * * * * * * * * * *	-	+
niR-2a	$\Delta G$	$Z_{\rm max}$	$\#Z \geq 3$	$Z_{\rm UTR}$	Gene	Alignment	Flags	$Ag Z \ge 1$
1	-39.0	6.78	2	11.85	CG1969-RB	GCUGGCUGGC-GGUG CGAGUAGUUUCGACCGACACUAU	5' cons 5' helix mispairing	NF
2	-38.6	6.66	1	6.66	CG4269-RA	GCUCCUG CAU - GGAUUGGCUGUGAUA CGAG UAGU - UUC - GACCGACACUAU **** ** +* - +*************	-	-
3	-38.0	6.49	1	6.49	reaper	- CUCAUCAAAGCGA UUGUGAUA CGAGUAGUUUCG ACCGACACUAU -************	-	NF
4	-34.3	5.42	1	5.42	GLaz	GCUUUGAUGAGCGCUGUGAUA CGAGUAGUUUCGACCGACACUAU ***+*****************************	mispairing	-
5	-33.5	5.19	1	5.19	BG:DS05899.3	GUUCAUCCCUUGGCGUUG-GGCUGUGU-UA CGAGUAGUUUCGACCGACAC-UAU *+*****	5' helix	NF
6	-33.2	5.1	1	5.1	Scr	GCUCGGUG - GGAGUG - GGUG - GUGGUG CGAGU - A - GUUUCG - ACCG - ACACUAU ***+- * ++**+ - ***+*+	-	_
7	-33.0	5.04	1	5.04	hbs	CAUGCGGC-GCUCGAAGGCUGUGAUA CGAGUA-GUU-UCGACCGACACUAU	-	-
8	-32.8	4.99	1	4.99	amon	GUUCAA - UAAAAGUGCUGGCUGUG CGAGU - AGUUU CGACCGACACUAU	5' helix	-
9	-32.5	4.9	1	4.9	grim	GCUCAAUCAAAGCGCA UUGUGAU -	_	NF
-	-210				<b>3</b> ·····	CGAGU - AGUUUCG ACCGACACUAU		
10	-32.1	4.78	2	8.01	CG7187-RA	GCUUUGAU GAGC GCUGUGGUG	5' cons	NF

 $\Delta$ G,  $Z_{\text{max}}$  and  $Z_{\text{UTR}}$  are explained in the text. Alignment: The target site is shown on top. Conventional basepairs are indicated by asterisks and G:U basepairs by plus signs; mismatches and gaps are indicated by dashes. Flags: The "5' conservation" (5' cons) flag identifies sites that differ in the two genomes at any residue complementary to positions 2–7 of the miRNA. The "5' helix" flag identifies sites that do not have at least six contiguous basepairs in positions 1–8. The "CDS+" flag indicates that the predicted site overlaps coding sequence on the same strand; "CDS-" indicates that the overlap is on the opposite strand. In some cases, Mfold structures include basepairs that are not between the miRNA and its target. "Mispairing" flags sites with artificially high folding energies. Ag ( $Z \ge 3$ ): Anopheles genes that cannot be reliably identified by our criteria are indicated as "NF." For the cases in which the orthologous Anopheles gene was found, the presence of a target site with a Z score  $\ge 3$  is indicated by a plus sign. Absence of a site or a Z score  $\ge 3$  is indicated by a minus sign. Heavy outlining indicates those loci that would pass stringent filtering of the lists using the flags and requiring lack of a conserved target in an Anopheles ortholog.

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Table 3. miR-277 Targets

Rank	Gene	Function	Enzyme	$\#Z \geq 3$	$Z_{ m UTR}$
1	CG31651	Protein GalNAc transferase	EC:2.4.1.41	2	7.31
2	CG5599	Val Leu Ile degradation	EC:2.3.1-	2	6.53
3	CG1673	Val Leu Ile degradation	EC:2.6.1.42	1	5.75
4	fz	Cell polarity		1	4.89
5	CG8199	Val Leu Ile degradation	EC:1.2.4.4	1	4.53
6	CG18549	_		1	4.23
7	CG1140	Val Leu Ile degradation	EC:2.8.3.5	1	3.9
8	scu	Val Leu Ile degradation	EC:1.1.1.35	1	3.81
9	CG15093	Val Leu Ile degradation	EC:1.1.1.31	1	3.79
10	CG7740	Membrane protein		1	3.64
11	CG17896	Val Leu lle degradation	EC:1.2.1.27	1	3.61

Required:  $Z \ge 3$  for *D. melanogaster, D. pseudoobscura, Anopheles.* DOI: 10.1371/journal.pbio.000060.t003

expressing cells at the wing margin (Figure 3E); wingless expression was also reduced (data not shown). Although bHLH transcription factors and Brd-like genes of the E(spl) complex are not strictly required for all aspects of Notch activity at the wing margin, they are required for cut expression (Ligoxygakis et al. 1999). miR-7 expression might provide a means to simultaneously downregulate these and other proteins, which might otherwise function redundantly to mediate Notch activity in the wing margin. We also noted reduced spacing of veins 3 and 4 in these wings, which may also reflect reduced Notch activity in controlling tissue growth (Baonza and Garcia-Bellido 1999). E(spl) genes are also expressed in proneural clusters, where they are required for sense organ determination and bristle patterning. Clones of cells lacking multiple genes of the E(spl) complex form extra bristles (de Celis et al. 1996). Consistent with this, we observed that expression of miR-7 under ptc-Gal4 control causes ectopic bristles and bristle duplication in the scutellum (data not shown). Taken together, these findings support the prediction that miR-7 miRNA regulates expression of bHLH and Brd-like proteins encoded by hairy and the E(spl) and Brd complex genes and implicates miR-7 as a possible regulator of Notch target gene expression. A more detailed analysis of miR-7 function will require isolation of lack of function mutations in the miR-7 gene.

Lai (2002) has reported complementarity between some miRNAs and sequence elements known as K boxes, Brd boxes, and GY boxes in the 3' UTRs of *E(spl)* and *Brd* complex genes. K boxes and Brd boxes have been implicated in posttranscriptional regulation, though no function was assigned to the *miR-7* complementary GY boxes (Lai and Posakony 1997; Lai et al. 1998). The presence of GY boxes in several *E(spl)* and *Brd* complex genes, as well as in *hairy* and *extramachrochatae* has been reported (Lai and Posokony 1998). Based on the presence of these boxes, Lai (2002) predicted *miR-7* target sites in *HLHm3* and in *Tom*. We extend these predictions to a much larger gene family and provide experimental validation for some of them, indicating that GY boxes participate in the regulation of *Notch* target genes.

#### miR-2 Regulates Proapoptotic Genes

The proapoptotic genes *reaper* and *grim* were among the top predictions for miR-2a and miR-2b (see Table 2). reaper, grim, and the third proapoptotic gene sickle are clustered in the genome and show blocks of high conservation in their 3' UTRs, which include the miR-2 family target sites (underlined in Figure 4A). Alignment of these sites shows a very similar pattern of predicted miRNA binding (Figure 4B). The corresponding sites are highly similar in D. pseudoobscura, but the orthologous genes cannot be identified in Anopheles. To evaluate these predictions, we made 3' UTR sensor transgenes for reaper, grim, and sickle. The expression level of the reaper 3' UTR sensor transgene was too low to be reliably studied in transgenic flies, so we used an in vitro assay to assess its function. Drosophila Schneider S2 cells express the miR-2 family of miRNAs (Lagos-Quintana et al. 2001). S2 cells were transfected with the reaper 3' UTR construct or with a version of the construct in which the miR-2-binding site was mutated (the residues shown in Figure 4B were replaced by a NotI site). A low level of GFP expression was detected in immunoblots of cells transfected with the reaper 3' UTR construct (Figure 4C, lane 2). The level of GFP expression was much higher in cells transfected with the mutated UTR construct, suggesting that the endogenous miR-2 family miRNAs in S2 cells can repress expression of a reporter construct via the reaper 3' UTR. The grim and sickle 3' UTR sensor transgenes were expressed at detectable levels in transgenic flies and were both downregulated by expression of miR-2b in the wing disc (Figure 4D and 4E). The miR-13 family is similar in sequence to the miR-2 family. Experimental validation will be needed to determine whether reaper, grim, and sickle are also regulated by miR-13. Identification of reaper, grim, and sickle as targets suggests that miR-2 family miRNAs may be involved in control of apoptosis.

## Statistical Evaluation of Target Predictions

Although a number of the top-ranking sites identified in our screen have been experimentally validated, we wanted to assess the likelihood that sites with equivalent scores can be found by chance. To do so, we calculated *E*-values for the



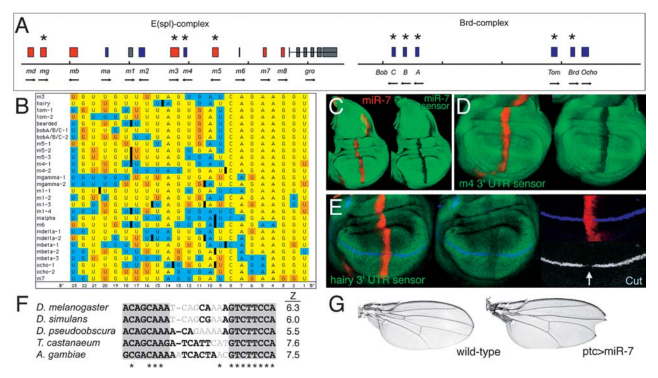


Figure 3. Experimental Validation of miR-7 Targets

(A) Schematic representation of the E(spl) and Brd gene complexes, which contain multiple predicted miR-7 target genes. bHLH-type transcriptional repressors are shown in red. Brd-type proteins are shown in blue. Other transcripts in the E(spl) cluster are in gray. Black asterisks indicate sites with no mismatch in the first eight residues (likely to be valid sites).

(B) miR-7 miRNA sequence showing the pattern of basepairing with target sites in *E(spl)* and *Brd* complex genes sorted in order of predicted folding energy. Yellow indicates a conventional basepair. Orange indicates a G:U basepair. Blue indicates a mismatch. The black bars indicate the position of loops in the target sites.

(C) Expression of the miR-7 sensor transgene is shown in green. Expression of the red fluorescent protein miR-7 miRNA under ptc–Gal4 control is shown in red. The right panel shows the miR-7 sensor alone.

(D and E) Expression of the m4 3' UTR and hairy 3' UTR sensor transgenes (green) were downregulated by miR-7 (red). Expression of the hairy 3' UTR sensor was much lower than the m4 3' UTR sensor overall. Cut protein, shown in blue, was downregulated in miR-7 expressing cells. The right panel shows a second example of Cut repression. The lower panel shows Cut channel alone.

(F) ClustalW alignment of miR-7 target sites in the 3' UTRs of hairy from several species. Asterisks indicate sequence identity. Black type indicates basepairs by Mfold (including G:U basepairs). Gray shading highlights the conserved miRNA-target binding region in all five species.

(G) Cuticle preparations of a wild-type adult wing and a wing expressing miR-7 under ptc–Gal4 control in the region between veins 3 and 4. Note the notching of the wing and the reduction of the region between veins 3 and 4, leading to partial fusion proximally. The size of the posterior compartment was increased apparently to compensate for reduction of the vein 3–4 region. DOI: 10.1371/journal.pbio.0000060.g003

bantam miRNA based on the tail of the cumulative distribution of  $\Delta G$  values for 10,000 random matches. An E-value predicts the number of background matches with a similar or better score (E-values scale with database size and are applicable to any distribution profile). Values greater than 1 are not significant, while those close to 0 are very significant. The results are presented on a logarithmic scale for UTRs containing one, two, or four sites of a given  $\Delta G$  value (Figure 5). The best single bantam site in the hid UTR had an E-value of 1.3. This means that background matches reach RNA-duplex energies similar to the best sites, even in the smaller conserved 3' UTR database. Indeed, target sites predicted using shuffled bantam miRNA sequences give folding energy distributions very similar to the native sequence (data not shown). Although single sites are not statistically significant, the presence of multiple sites within a single UTR can greatly increase the significance of the prediction. Combining the three bantam sites (Z > 3) predicted in the hid 3' UTR gives an *E*-value of  $1.8 \times 10^{-5}$ . Some single sites are sufficient to mediate regulation by a miRNA; however, we emphasize that

the lack of statistical significance for even the best single site means that they require experimental validation.

## Additional Validation by Cross-Genome Comparison

One means to improve the significance of the predictions would be to require conservation in a third genome. The two Drosophila species are separated by an estimated 30 million years. The mosquito A. gambiae is separated from Drosophila by 250 million years. Orthologous mosquito genes have been defined for approximately two-thirds of Drosophila genes; however, systematic comparison showed great differences in length between orthologous gene pairs (Zdobnov et al. 2002). Indeed, we were able to identify orthologous last exons with confidence for only half of these pairs, or one-third of D. melanogaster genes. We have therefore chosen to use conservation in Anopheles to provide more stringent evaluation of target site conservation, instead of requiring it generally. The presence of a conserved site with a high Z score across all three genomes increases the confidence that the site is functional. To illustrate the utility and limitations of this, we examined the top 100 predictions for miR-7 and miR-2. The



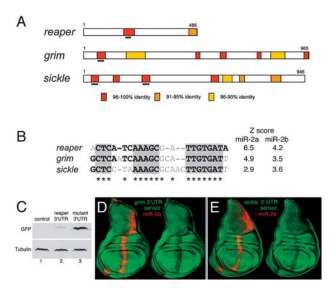


Figure 4. Experimental Validation of miR-2 Targets

- (A) Conservation of sequences in the 3' UTRs of reaper, grim, and sickle genes of *D. melanogaster* and *D. pseudoobscura*. Blocks of high sequence similarity are color-coded. The location of predicted miR-2/miR-13 family target sites are underlined in black.
- (B) ClustalW alignment of predicted miR-2/miR-13 family target sites in the reaper, grim, and sickle 3' UTRs. Z scores for miR-2a and miR-2b are shown for each site. The first bases of the grim and sickle sites do not pair with the miRNAs. Because we use a hairpin-forming linker sequence, this causes a penalty in Mfold, which gives these sites lower Z scores than they should otherwise have.
- (C) Immunoblot of S2 cells transfected to express a tubulin promoter–EGFP–reaper 3' UTR construct (lane 2) or a comparable construct from which the *miR-2/miR-13*-binding site in the UTR was deleted (lane 3). Control cells were transfected with empty vector (lane 1). The blot was probed first with antibody to GFP and then reprobed with anti-tubulin as a loading control.
- (D and E) Expression of the *grim* and *sickle* 3' UTR sensor transgenes (green) was downregulated by *miR-2b* expressed under ptc-Gal4 control (red).

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Anopheles orthologs were identified for 52 of the top 100 predicted miR-7 targets. Of these, 11 had conserved target sites ( $Z \geq 3$ ), including four of the top ten predictions: hairy, Tom, m4, and CG14989 (see Table 2). For miR-2a, forty of the top hundred predictions had a detectable ortholog in Anopheles. Of these sites, five were conserved in Anopheles ( $Z \geq 3$ ), and none of these were among the top ten predictions. Conservation in Anopheles can be used to enrich for sites with a higher probability of being valid, but increases the risk of missing real targets. It is only useful in cases where the orthologous UTR region can be identified, which, for example, is not the case for the validated miR-2a targets grim, reaper, and sichle.

#### miR-277: A Metabolic Switch?

Table 3 shows predicted miR-277 targets that are conserved  $(Z \ge 3)$  in Anopheles. Of the top 11, seven are enzymes involved in branched chain amino acid degradation (Figure 6). At more relaxed stringency  $(Z \ge 2)$ , two additional enzymes were identified (Figure 6) along with a number of unrelated loci. This striking clustering of functionally related enzymes suggests that miR-277 regulates the pathway for valine, leucine, and isoleucine degradation by downregulating many of its enzymes and thus acts as a metabolic switch. The degradation of these essential amino acids is presumably

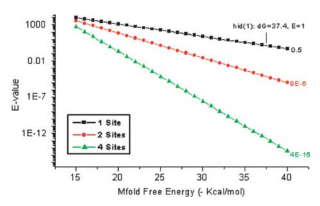


Figure 5. Statistical Evaluation of Predicted Targets

Plot of *E*-values as a function of free energy of folding. Y-axis: logarithmic scale of *E*-values. X-axis: free energy of folding calculated by Mfold. Calculations for one, two, and four sites are shown separately. The position of the best *bantam* site in *hid* is shown for reference.

DOI: 10.1371/journal.pbio.0000060.g005

regulated under conditions of starvation or excess dietary intake. *miR-277* expression has so far only been detected in adult flies (Aravin et al. 2003; Lai et al. 2003), suggesting a role in regulating metabolic responses to environmental conditions. Interestingly, the human homolog of *CG8199* is mutated in maple syrup urine disease. It remains to be determined whether these enzymes are regulated by miRNAs in vertebrates.

## Features Shared by Validated Targets

Comparison of the five previously validated targets and the six new targets validated here revealed three features shared by all sites. First, cross-genome comparison showed perfect sequence identity in the target site residues that basepair with residues 2-8 of the miRNA (see Figure 1B). This was also true for the newly validated target sites (data not shown). Second, the pattern of basepairing between the miRNAs and their targets, shown in Figure 2A, suggested that a continuous helix of at least six of the first eight basepairs might be required (allowing G:U basepairs). This was also true for the newly validated target sites (see Figures 3B and 4B). Third, many transcripts in the D. melanogaster genome overlap other transcripts on the same strand or on the opposite strand of the DNA. There are many examples of alternate splicing that produce alternate 3' UTRs so that one UTR variant may include coding exons from another variant. In such cases, the basis for the sequence conservation between genomes is unclear. None of the validated sites from Drosophila overlaps coding sequence (CDS) on either strand (this feature was not examined for the C. elegans sites).

Target sites that do not share these features are indicated in Table 2. These features can be used to increase the stringency of the screen, by discounting sites that differ from validated targets. For *miR-7* this would eliminate two of the top ten predictions so that the validated targets would constitute three of the remaining top eight predictions. For *miR-2a* this would eliminate four of the top ten predictions, so that the validated targets *reaper* and *grim* would rank in positions 2 and 6. We have chosen not to implement the flags as filters to exclude predictions because they are based on a relatively small set of 11 validated targets, although we note



## VALINE, LEUCINE AND ISOLEUCINE DEGRADATION

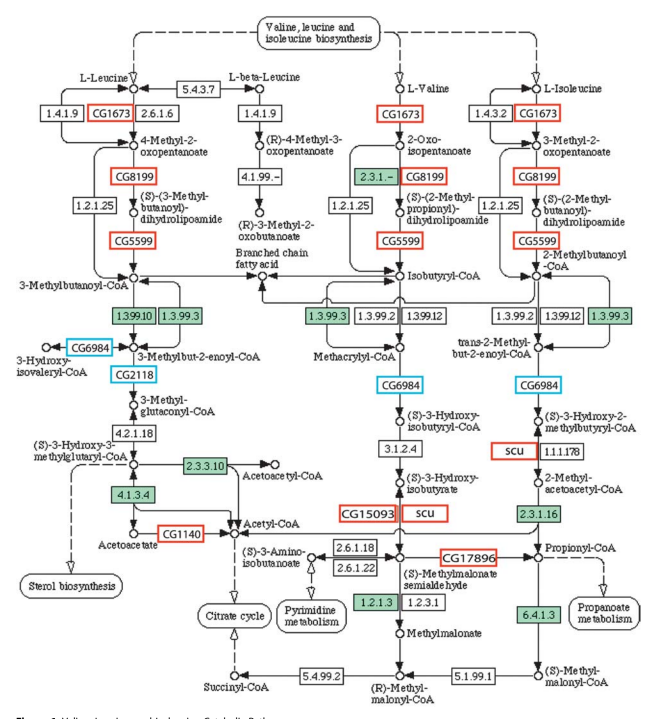


Figure 6. Valine, Leucine, and Isoleucine Catabolic Pathway

Enzymes identified as miR-277 targets are boxed and identified by CG number. Red boxes required Z > 3 in Anopheles. Blue boxes required Z > 2 in Anopheles. In addition to the predicted targets, the other enzymes for which the gene has been identified in Drosophila are shaded in green. The metabolic pathway chart is from www.genome.ad.jp/kegg/pathway/map/map00280.html. DOI: 10.1371/journal.pbio.0000060.g006



that all nine predicted *miR-277* targets would pass such a filter. When more targets are validated, we will learn whether these features have a general predictive value.

#### Discussion

One of the major limitations in studying animal miRNA function is the difficulty in identifying miRNA targets. Our screening strategy has proven to be useful for predicting new miRNA targets. Three new targets have been experimentally validated for miR-7 and for miR-2, bringing the total number of validated targets of animal miRNAs to 11. In addition, we predict a number of miRNA-target pairs or target families that seem likely to be valid, but require experimental validation. Our study depended on the high-quality annotation of the D. melanogaster genome and the availability of the D. pseudoobscura genome sequence. Where possible, we have extended the analysis to include evaluation of predicted sites in the A. gambiae genome. More complete annotations of the fly and mosquito genomes, aided by cDNA sequencing projects, will increase the number of genes for which orthologous UTR sequences can be defined. This will permit more sensitive and more extensive cross-genome comparison. We also expect improvements to come from further knowledge of the structural requirements of miRNA-target pairing.

## **Evaluation of Target Predictions**

In designing the screening strategy, we considered the balance between sensitivity and specificity. We chose a search strategy that was based on the known examples, but generalized to allow detection of similar targets. By doing so, we risk missing fewer valid targets at the expense of including more false positives, as indicated by our statistical analysis (see Figure 5). To help distinguish false positives from potentially valid targets, we identify features shared by valid targets and, where possible, test predictions for conservation in a third, more distantly related, genome. Both positive and negative results in tests of new predictions will provide a better understanding of how miRNAs bind their targets, perhaps highlighting positions that are particularly critical. This may permit us to achieve both high sensitivity and high specificity in target prediction.

Complete tables of target site predictions are available as Dataset S1 and at www.miRNA.embl.de. These tables report Z scores and sequences for the D. melanogaster and D. pseudoobscura target sites and, where possible, for the Anopheles target site. The tables contain flags to identify sites that share the features described above. We recommend using these flags to filter the predictions, but note that this may exclude valid targets. (Filtered tables containing the top 100 predictions are available as Dataset S2 and at www.miRNA.embl.de) We recommend making use of the Anopheles data to discount predictions where the othologous gene is identified and the site is absent or has a low Z score (Z < 2). We emphasize that the absence of an identified orthologous 3' UTR in the mosquito should not be taken as evidence that a target prediction is not valid.

The presence of a conserved site in all three genomes increases the confidence that a predicted site is valid, as in the case of the *miR-7* sites in *hairy* and *Tom.* Also, *dpld*, the *Drosophila* homolog of the *let-7* target *lin-41*, ranks second among *Drosophila let-7* targets when conservation in *Anopheles* 

was required. A number of other target predictions that meet these requirements look quite promising. We have high confidence that the cluster of enzymes in the branched chain amino-acid degradation pathway will prove to be valid miR-277 targets. Another promising candidate is the predicted miR-9a target Lyra. Lyra contains two predicted miR-9a sites. The best Lyra site ranks first among all predicted miR-9a targets that are conserved in Anopheles. Intriguingly, mutations affecting the Lyra 3' UTR lead to a dominant phenotype and to increased Lyra protein levels, an observation that strongly suggests that Lyra is subject to translational regulation. miR-9 is an excellent candidate to mediate this regulation. Many other miRNA-target pairs are identified with sites of a similar quality to those mentioned here (examples include four conserved sites for miR-309 in Ets65a at  $Z \geq 2$ ). As a cautionary note, we wish to emphasize that conservation of target sites in Anopheles, while compelling, should not be taken as sufficient evidence of function without experimental validation.

Although it is more difficult to distinguish functional sites from false positives in the cases where only two genomes are compared, we have made use of the clustering of related genes to identify real targets. reaper, grim, and sickle have been validated as miR-2 targets. We note that the Netrin receptor unc-5 and Netrin-A rank second and fourth among predicted miR-288 targets. We also noted an abundance of transcription factors among the predicted targets of miR-9, miR-279, and miR-286 for which orthologous UTRs were not identified in Anopheles. These predictions merit further study.

#### Single versus Multiple Sites

Our statistical analysis shows that the very best single predicted target sites are not statistically significant, even though we have used a reduced database consisting of conserved 3' UTR sequences. This means that prediction of any single target site cannot be taken as evidence for regulation of a transcript by a miRNA without experimental validation. Sites that are not statistically significant alone can be significant when combined. For example, although none of the bantam sites are significant individually, their combined scores are highly significant and supported by experimental validation. 3' UTRs with multiple predicted target sites are likely to be valid targets for regulation by the miRNA, particularly if their best single sites also rank high in the lists of predicted targets.

Despite the advantages conferred by multiple sites, single miRNA target sites can mediate regulation in vivo. The C. elegans lin-4 miRNA appears to regulate its target lin-28 through a single site (Moss et al. 1997). We have presented evidence that miR-2 family miRNAs can regulate expression of transgenes containing the 3' UTRs of reaper and grim, which have one predicted target site, as well as the sickle 3' UTR, which has two predicted sites. Similarly, miR-7 can regulate expression of transgenes containing the HLHm3, m4, and hairy 3' UTRs, which have one predicted target site. Further work will be needed to gain insight into what makes some single sites functional and others not. One possibility is that a single site for one miRNA might function in conjunction with independent target sites for other miRNAs in the same UTR. Indeed, a survey of our lists of target predictions indicates that many 3' UTRs are predicted to contain binding sites for more than one miRNA.



#### **Materials and Methods**

Conserved 3' UTR database. D. melanogaster 3' UTRs were obtained from the Berkeley Drosophila Genome Project (www.fruitfly.org/ annot/release3.html) and those of >50 nt were selected. Duplicate UTRs from different splice variants of the same transcript were removed. For each of the resulting 10,196 nonredundant 3' UTRs, we mapped the last 50 amino acids of the corresponding open reading frame to the D. pseudoobscura genome sequence with TBLASTN (E s 10<sup>-5</sup>; hgsc.bcm.tmc.edu/projects/Drosophila). We selected UTR matches that included the last 10 residues and had a sequence identity  $\geq 80\%$  or  $E \leq 10^{-10}$  and compared these UTRs to the 3,000 nt downstream of the putative D. pseudoobscura ortholog with BLASTN (word size = 7; E < 10,000, assuming a database the size of the whole D. pseudoobscura genome). Nonconserved nucleotides or those outside the matched regions were replaced by Ns in the D. melanogaster 3' UTR database to produce the conserved 3' UTR database. The D. pseudoobscura genome has not been fully assembled. This means that some D. pseudoobscura genes are located close enough to the end of a contig that the UTR sequences may be missed. 386 D. melanogaster genes mapped to the D. pseudoobscura genome less than 1 kb from a contig end; 189 mapped less than 500 nt from a contig end. UTR conservation may be underestimated for these genes. For 3,564 genes, we did not detect a suitable ortholog using this protocol. Of these, 571 are known genes; the others are predicted genes about which little is known. For the 4,662 D. melanogaster genes lacking annotated UTRs, we assumed 3' UTRs of 2 kb after of the stop codon and built a separate database of predicted UTRs. The search for Anopheles orthologs was done using TBLASTN for the last 50 amino acids of each D. melanogaster ORF. Owing to the more extensive sequence divergence, a lower cutoff threshold was allowed (E < 0.05) if the last exon of the predicted ORF mapped to the same location (±1 kb) in the annotated genome as the orthologous gene (Zdobnov et al. 2002). If not, the cutoff was  $E \leq 10^{-5}$ , as for *D. pseudoobscura*. The second, more stringent step of comparing the last 10 amino acids was omitted.

**Sequence search.** HMMer (Eddy 1996) profiles were constructed for each of two alignments per miRNA containing copies of the reverse complement of the first (5') 8 nt of the miRNA. The first alignment contained five copies of the exact complement; the second had an additional five copies with C replaced by T and A replaced by G to allow for G:U mismatches. We searched the conserved 3' UTR database with both profiles and lenient domain bit score threshold (domT  $\geq$  3) and combined the results. Sequence matches were extended to miRNA length plus 5 nt, the hairpin loop and miRNA sequence were added, and the sequence was evaluated using Mfold. For Anopheles, predicted UTRs were searched for the presence of residues 2–7 of the predicted target site. The target sequences were extended and evaluated using Mfold. Only the best site in the Anopheles UTR was reported.

**Statistics.** For each miRNA, we calculated the mean and SD of a background distribution, i.e., the Mfold free energy  $\Delta G$  of 10,000 randomly selected sequences from the conserved UTR database with lengths of miRNA plus 5 nt. For each prediction, we calculated the Z error as the number of SDs above the mean. To compute E-values, we fit an exponential function to the cumulative background distributions extrapolated it to give a value for any observed energy and database size. E-values are not restricted to normal distributions and scale with database size, so different searches can be compared.

**Constructs.** The hairy, HLHm3, reaper, grim, and sickle 3' UTRs were amplified by polymerase chain reaction from genomic DNA and cloned into tubulin–EGFP as described (Brennecke et al. 2003). m4 lacks an annotation for its 3' UTR. A predicted UTR consisting of 900 nt was used. The miR-7 hairpin and the miR-2b hairpin from the spitz intron were cloned downstream of DSred2 (Clontech, Palo Alto, California, United States) in pUAST.

Antibodies. Rabbit anti-GFP (TP401) was purchased from Torrey Pines Biolabs (Houston, Texas, United States). Mouse anti-tubulin (T-9026) was purchased from Sigma (St. Louis, Missouri, United States). Mouse anti-Cut is described in Blochlinger et al. (1993).

## **Supporting Information**

Dataset S1. Complete Lists

These contain all predicted targets for each miRNA. There is one file per miRNA. No filtering has been done. From left, the columns are as follows:

gene: FlyBase identifier.

name: if there is one, in addition to the identifier.

GO term: gene ontology information about the gene product.

Z (me): score for the D. melanogaster site.

dG(me): folding energy for the D. melanogaster site.

Alignment (me): of the miRNA to the target site; asterisk, conventional basepair; plus sign, G:U basepair; minus sign, mismatch.

start and stop: position of the site in the 3' UTR.

*mfold (me)*: the target site, linker, and miRNA sequence as submitted to Mfold.

#sites: total number of sites for the miRNA found in that UTR.

Z(UTR): sum of Z for all sites  $Z \ge 3$  in that UTR.

Z(max): the best Z score for the UTR.

#Z > 3: total number of sites of Z > 3 found in that UTR.

*UTR*: whether the site is an experimentally defined or predicted UTR. 5' conservation, 5' helix, CDS+, CDS-, and mispairing flags are described in text and in Table 2.

*Z(ps)*: *Z* score for the corresponding site from *D. pseudoobscura*. It is not always possible to unambiguously identify the corresponding site from *D. pseudoobscura*, even though the sequence is in the database (because the genome is not assembled and some regions appear two or more times). We chose to omit ambiguous cases. This problem will disappear when the *D. pseudoobscura* genome sequence is assembled so that unambiguous gene assignment is possible.

dG(ps): folding energy for the site in the *D. pseudoobscura* UTR. The folding energy can be used to determine a *Z* score on the same scale as the site from *D. melanogaster*. This allows a direct comparison of site quality.

Alignment (ps): as above, for the D. pseudoobscura site.

Mfold (ps): as above, for the D. pseudoobscura site.

Ano-tblastn: TBLASTN score for the Anopheles ortholog (explained in the text).

Z(ano): as above, for Anopheles. dG(ano): as above, for Anopheles. Aln(ano): as above, for Anopheles. Mfold(ano): as above, for Anopheles.

DOI: 10.1371/journal.pbio.0000060.sd001

#### Dataset S2. Top 100 List

This dataset presents our recommended filtering of the lists. Only sites in experimentally validated UTRs were included. The 5' conservation, 5' helix, CDS+, and CDS- flags were used to discard sites that we consider less likely to be valid. This allows use of a lower Z score cutoff ( $Z \ge 2$ ). Sites of Z < 2 were removed and  $Z_{\rm UTR}$  includes all sites  $Z \ge 2$ . Columns are as above, except the flags have been removed.

DOI: 10.1371/journal.pbio.0000060.sd002

#### **Accession Numbers**

The accession numbers for the miRNAs discussed in this paper are bantam (AJ550546, Rfam MI0000387), let-7 (NR\_000938), lin-4 (NR\_000799), miR-2a (RF00047, AJ421757), miR-4 (AJ421762), miR-7 (AJ421767), miR-9 (AJ421769), miR-11 (AJ421771), miR-13a (AJ421773), miR-14 (AJ42177), and miR-277 (Rfam MI0000360).

The accession numbers for the target genes discussed in this paper are CG1140 (NM\_167928), CG1673 (NM\_132656), CG5599 (NM\_132772), CG8199 (NM\_141648), CG15093 (NM\_166306), CG17896 (NM\_130489), dpld (NM\_080033), Drice (NM\_079827), grim (NM\_079413), hairy (NM\_079253), hairy (D. simulans) (AY055843), hairy (T. castaneum) (AJ457831), hid (NM\_079412), HLHm3 (NM\_079785), lin-14 (NM\_077516), lin-28 (NM\_059880), lin-41 (NM\_060087), lin-57 (NM\_076575), Lyra (NM\_080079), m4 (NM\_079786), reaper (NM\_079414), scu (NM\_078672), sickle (AF460844), and Tom (NM\_079349).

## **Acknowledgments**

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**Conflicts of Interest.** The authors have declared that no conflicts of interest exist.

**Author Contributions.** AS, JB, RBR, and SMC conceived and designed the experiments. AS and JB performed the experiments. AS, JB, RBR, and SMC analyzed the data. AS, JB, RBR, and SMC wrote the paper.



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