# Genome-wide identification of Ago2 binding sites from mouse embryonic stem cells with and without mature microRNAs

Anthony K L Leung<sup>1,5</sup>, Amanda G Young<sup>1,2,4,5</sup>, Arjun Bhutkar<sup>1</sup>, Grace X Zheng<sup>1,3,4</sup>, Andrew D Bosson<sup>1,2</sup>, Cydney B Nielsen<sup>2,4</sup> & Phillip A Sharp<sup>1,2</sup>

MicroRNAs (miRNAs) are 19–22-nucleotide noncoding RNAs that post-transcriptionally regulate mRNA targets. We have identified endogenous miRNA binding sites in mouse embryonic stem cells (mESCs), by performing photo-cross-linking immunoprecipitation using antibodies to Argonaute (Ago2) followed by deep sequencing of RNAs (CLIP-seq). We also performed CLIP-seq in Dicer<sup>-/-</sup> mESCs that lack mature miRNAs, allowing us to define whether the association of Ago2 with the identified sites was miRNA dependent. A significantly enriched motif, GCACUU, was identified only in wild-type mESCs in 3' untranslated and coding regions. This motif matches the seed of a miRNA family that constitutes ~68% of the mESC miRNA population. Unexpectedly, a G-rich motif was enriched in sequences cross-linked to Ago2 in both the presence and absence of miRNAs. Expression analysis and reporter assays confirmed that the seed-related motif confers miRNA-directed regulation on host mRNAs and that the G-rich motif can modulate this regulation.

miRNAs are key regulators of gene expression in fundamental processes including cell proliferation, cell death, cell differentiation and cellular responses to the environment<sup>1–3</sup>. These short non-coding RNAs guide a ribonucleoprotein complex, containing a member of the conserved Argonaute (Ago) protein family, to sites predominantly in the 3' UTRs of their target mRNAs, resulting in the destabilization of the message and/or inhibition of translation<sup>4,5</sup>. Biochemical and computational studies have shown that base pairing between the 'seed' (second to seventh nucleotide) at the 5' end of the miRNA and mRNA target is important for this regulation in animals<sup>6-10</sup>. Comparative genomic analysis for miRNA seed sites in 3' UTRs suggests that miRNAs regulate ~60% of all mammalian mRNAs<sup>11</sup>. Moreover, both comparative genomic analysis and emerging data from a handful of genes suggest that miRNAs also target coding sequences<sup>8,12,13</sup>, but the prevalence of this interaction is unclear. Therefore, recent efforts<sup>14-16</sup> have aimed at identifying bona fide miRNA binding sites on a genome-wide scale in samples from whole mouse brain and whole-animal nematode preparations. However, one challenge of these studies is to deconvolute the miRNA-target relationships in the mixed cell types from these samples<sup>15,16</sup>. In this study, we dissect the miRNA-target relationship in a homogeneous cell population-mouse embryonic stem cells (mESCs)—with defined miRNA characteristics<sup>17–19</sup>. We isolated RNAs photo-cross-linked to Ago2, of which hundreds contained a seed match to the mESC-enriched miRNA cluster miR-290-295, and, unexpectedly, the majority also contained a G-rich motif.

#### RESULTS

Ago2-CLIP-seq identified miRNA-dependent and -independent sites RNA tags photo-cross-linked to Ago2 in mESCs were isolated by immunoprecipitation and subjected to deep sequencing (CLIPseq)<sup>15,16,20,21</sup>. Notably, no RNA species were detectable by autoradiography in Ago2 immunoprecipitates without cross-linking, suggesting that cloned RNA tags require cross-linking and thus are in direct association with Ago2 (Fig. 1a and Supplementary Fig. 1a). In addition, we performed a parallel analysis in derivative mESCs that lacks Dicer and hence mature miRNAs<sup>22</sup>. Unexpectedly, we identified specific RNAs cross-linked to Ago2 in Dicer<sup>-/-</sup> cells (Fig. 1a), indicating that Ago2 can associate with RNAs in a miRNA-independent manner<sup>23,24</sup>.

Approximately 24.5 million sequenced RNA tags from three wildtype mESC libraries representing two biological replicates (WT1A, WT1B, WT2) and 10.6 million tags from two Dicer<sup>-/-</sup> libraries (KO1, KO2) were processed and mapped to the mouse genome (Supplementary Methods and Supplementary Fig. 1b-d). Across all libraries, 79% reads uniquely matched to the genome, 21% mapped to nonunique locations and 0.05% could not be aligned.

miRNAs cross-linked to Ago2 in mESCs were identified by screening reads with unique and repeat matches to the genome against noncoding RNA databases (Supplementary Fig. 1e and Supplementary Methods). Mature miRNAs are highly enriched, as expected<sup>17,18</sup>, in Ago2-cross-linked samples from wild-type cells compared with

Received 5 April 2010; accepted 29 November 2010; published online 23 January 2011; corrected after print 12 May 2011; doi:10.1038/nsmb.1991

<sup>&</sup>lt;sup>1</sup>The Koch Institute for Integrative Cancer Research, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA. <sup>2</sup>Department of Biology, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA. <sup>3</sup>Computational and Systems Biology, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA. <sup>4</sup>Present addresses: Salk Institute for Biological Studies, La Jolla, California, USA (A.G.Y.); Howard Hughes Medical Institute and Program in Epithelial Biology, Stanford University School of Medicine, Stanford, California, USA (G.X.Z.); Michael Smith Genome Sciences Centre, Vancouver, British Columbia, Canada (C.B.N.). <sup>5</sup>These authors contributed equally to this work. Correspondence should be addressed to P.A.S. (sharppa@mit.edu).

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Figure 1 Identification of miRNA-dependent and -independent motifs associated with Ago2. (a) Autoradiograph of <sup>32</sup>P-labeled RNA from Ago2 complexes immunoprecipitated from Dicer-I- mESCs, WT mESCs and WT mESCs without UV cross-linking. Bracketed area indicates the region of the blot that was excised and analyzed; arrow indicates where endogenous Ago2 migrates. (b) Log plot of mature miRNA cloning frequency in the WT1A Ago2-CLIP library versus whole-cell miRNA from wild-type mESCs (Pearson coefficient r =0.63; Spearman coefficient  $\rho = 0.86$ ). (c) Data processing pipeline. After preprocessing for linker matching and stripping, and subsequently mapping to the mouse genome, Ago2-CLIP sequence reads were subjected to the filtering steps as described in the text (Supplementary Methods and Supplementary Table 1 for more details). WT reads were subjected to all four filters, KO reads were subjected to first three filters, Clustering, Normalization, and Multi-Library (K01,K02 overlaps).

(d) Sequence logos and statistics of the top



three significantly enriched motifs derived from motif analysis of Ago2-CLIP 3' UTR-mapping clusters using the motif tool MEME (Supplementary Methods for details). The number of sites containing each motif out of the number of clusters examined is as indicated. In the heat map, red indicates significant enrichment ( $E < 1 \times 10^{-5}$  cutoff) and blue indicates absence of significant enrichment in either the WT or KO libraries. The G-rich motif (far left) was highly enriched in all libraries analyzed, and the representative consensus sequence was taken from MEME analysis (width 4–8) on Normalized KO1 clusters that had overlaps with clusters from any Normalized WT library. The GCACU[UG] (middle) and CC[AG]GCC (far right) motifs were significantly enriched by MEME analysis (width 6) in only the 430 clusters that passed all four filters in the combined wild-type libraries and not in the Ago2-CLIP 3' UTR clusters from any Normalized KO library. There were 48 instances of GCACUU and 31 instances of GCACUG represented in the GCACU[UG] motif. (e) Hexamer enrichment analysis by statistical overrepresentation within individual libraries, using an enumerative approach (Supplementary Methods). From left to right: Normalized and Multi-library filtered Ago2-CLIP WT1A (WT), Normalized and Multi-library filtered Ago2-CLIP KO2 (KO), and Normalized, Multi-library and Knockout filtered Ago2-CLIP WT1A (WT-KO). Refer to Supplementary Figure 2 for hexamer enrichment analyses in other libraries. Two measures of significance are plotted: the x axis shows the z score, which is a measure of the number of s.d. the observed frequency of a hexamer exceeds its expectation; the y axis is the negative log<sub>10</sub> P value (two-sided Fisher's exact test) of the hexamer enrichment above background. All hexamers with a false discovery rate (FDR)  $\geq 0.5\%$  are in gray and not considered as significantly enriched. Significantly enriched hexamers (FDR < 0.5%) are classified into two types: G-rich (≥3 Gs, red) and non–G rich (blue). Those hexamers that match the consensus G-rich motif derived from MEME analysis are circled. Hexamers only significantly enriched in WT are GCACUU (black dots) and CCAGCC (light blue dot). Note the change in scale for the far right plot.

*Dicer*<sup>-/-</sup> cells. The *miR-290–295* cluster, *miR-467* family and *miR-302~367* cluster (most members of which share the AAGUGC seed) represent the largest fraction (~68%) of the Ago2–cross-linked mature miRNA population<sup>17–19,25</sup> (**Fig. 1b** and **Supplementary Fig. 1e**), and the Ago2–CLIP and whole-cell miRNA populations were positively correlated (**Fig. 1b**). Although the WT2 library had more reads mapping to ncRNA and repetitive regions than WT1 libraries, the distribution of cross-linked miRNAs was similar between the libraries (**Supplementary Fig. 1e**). The specificity of CLIP-seq method is shown by the absence of Ago2 cross-linking to the highly abundant rRNAs (~0.2%) and tRNAs (~0.2%).

For each library, the remaining tags that mapped uniquely to 3' UTRs were subjected to a data-processing pipeline that consists of four filtering steps (**Fig. 1c** and **Supplementary Table 1**). First, identical reads were collapsed as a single read to eliminate potential PCR bias, and overlapping reads were then clustered (*Clustering filter*, **Fig. 1c**). Second, 25-nt flanking regions were added to either side of the clusters in case an RNase cleavage separated the locations where the miRNA bound and Ago2 cross-linked. Clusters were further considered only if they were significantly enriched over background levels (*P* < 0.01, *Normalization filter*, **Fig. 1c** and **Supplementary Methods**). Third, to select for a reproducible signal, only the clusters that overlapped with at least one other cluster from a *Normalized* biological replicate library were considered (*Multi-Library filter*, **Fig. 1c**). Fourth, the remaining WT clusters that had overlaps with clusters

from either *Normalized Dicer<sup>-/-</sup>* library (Knockout, KO) were removed (*Knockout filter*, **Fig. 1c**). Finally, after removal of duplicates from technical replicates WT1A and 1B, 430 clusters in the combined WT libraries (244 in WT1[A+B] and 186 in WT2), of average length 81 nt, passed all four filters. Various sets of clusters from different filtering steps were then subjected to motif enrichment analysis using two independent approaches.

First, significantly enriched motifs were identified in 3' UTRmapped clusters from WT and KO sets independently (Fig. 1d and Supplementary Table 2 for all motif analyses). The motif discovery tool MEME<sup>26</sup> was used to search for significantly enriched motifs of variable lengths over background (Supplementary Table 2a and Supplementary Methods). We found significant enrichment for G-rich motifs in clusters from both WT and KO libraries, suggesting that Ago2 may be associated with G-rich sequences independently of miRNAs. The G-rich motifs identified independently in WT and KO libraries have an average Pearson correlation of ~0.80, suggesting a high degree of similarity between the motifs (Supplementary Table 3a). Therefore, we defined a consensus G-rich motif by performing MEME analysis on Ago2-cross-linked clusters that overlapped between WT and KO libraries. This motif was highly statistically enriched (*E* value =  $2.9 \times 10^{-386}$ ) and present in 87% of the common clusters between Normalized WT and KO libraries. Examination of individual libraries showed that this consensus G-rich motif was present at approximately equal frequency in sequences cross-linked to Ago2 from wild-type and *Dicer<sup>-/-</sup>* mESCs, again suggesting that its association with Ago2 is miRNA independent (**Supplementary Table 3b**).

One of the two significantly enriched miRNA-dependent motifs identified in 3' UTRs was GCACU[UG] (*E* value <  $1 \times 10^{-5}$  79 instances from 430 WT clusters). GCACUU (48/79 GCACU[UG] motifs) is complementary to the seed AAGUGC of several highly expressed miRNA families in mESCs. The only other statistically enriched miRNA-dependent motif in the selected clusters was CCAGCC (51 instances). Unlike GCACUU, however, this motif is not complementary to any miRNA sequence with appreciable expression in mESCs.

Second, to independently investigate enrichment of motifs from clusters within each individual CLIP library (Fig. 1e, Supplementary Fig. 2a and Supplementary Table 2b for top 20 motifs), we used an enumerative approach that guarantees global optimality by statistical over-representation and avoids the problem of becoming trapped at local optima inherent in most general motif-finding algorithms<sup>27</sup>. Briefly, we measured the statistical significance of the occurrence of all possible n-mer sequences within each library compared to their occurrence in sequences drawn randomly given a background distribution. This independent analysis confirmed the significant enrichment of G-rich hexamers (containing three or more Gs; red dots, Fig. 1e and Supplementary Fig. 2a) out of all possible hexamers in WT and KO libraries, as demonstrated by their P values (for example,  $5.59 \times 10^{-11} < P < 2.09 \times 10^{-2}$  for WT1A) and z scores at a false discovery rate (FDR) < 0.5% (Supplementary Methods for derivations). The three hexamers encompassed in the consensus G-rich motif are among the top seven significantly enriched hexamers in the WT and KO libraries (black circle, Fig. 1e). Enrichment was observed exclusively in WT libraries for several non-G rich hexamers (blue dots, Fig. 1e), including GCACUU (black dot) and CCAGCC (light-blue dot). 29 non-G rich hexamers matched to miRNA seeds, but these miRNAs are associated with Ago2 at a median frequency of 0.003% (*P* < 0.05; Supplementary Fig. 1f and Supplementary Table 2b,c). Several other miRNA seed-matching hexamers occurred with high frequency, but these were not observed significantly more than expected by chance and thus were not further considered (Supplementary Table 2c). The miRNA-dependent motif GCACUU is one of the top significantly enriched non-G rich hexamers in all WT libraries, including WT2, which had a lower proportion of 3' UTR-mapping clusters. The enrichment of GCACUU is particularly apparent after applying the Knockout filter, where common clusters between WT and KO libraries, many of which contain G-rich hexamers (red dots), are removed from the WT set. In effect, the Knockout filter reveals GCACUU as the most significantly enriched non-G rich hexamer in mESCs expressing miRNAs (black dot, leftmost versus rightmost panels in Fig. 1e). We also observed enrichment of heptamers and octamers containing GCACUU, which matches the extended seed region<sup>6,8–10</sup> of the AAGUGC-seed family (Supplementary Table 2d).

Sequences mapping to coding sequences (CDS) were also subjected to the same data-processing pipeline, resulting in a set of 197 clusters (106 in WT1[A+B], 91 in WT2). As in the case of 3' UTR clusters, G-rich motifs were highly significantly enriched by MEME analysis in CDS clusters from both WT and KO libraries (P < 0.01, constituting ~25% and ~30% of clusters, respectively; data not shown). Moreover, the GCACUU hexamer was observed in the CDS clusters from wild-type libraries (22 instances in 197 clusters; **Supplementary Table 2a**) but not KO libraries. Similarly, in the enumerative analysis of individual libraries, G-rich hexamers were highly enriched in both WT and KO libraries, and GCACUU was enriched only in WT libraries (**Supplementary Fig. 2b** and

**Supplementary Tables 2e**, **3c**). Unlike in 3' UTR–mapped clusters, both MEME and enumerative analyses indicated no enrichment for CCAGCC in the CDS-mapped clusters.

#### Ago2-CLIP genes exhibit miRNA-dependent gene expression

mRNAs targeted by miRNAs are often destabilized, resulting in a lower abundance of targeted transcripts in wild-type cells as compared to *Dicer<sup>-/-</sup>* cells<sup>4,28,29</sup>. We used mRNA expression of two sets of Ago2-CLIP 3' UTR GCACUU transcripts in wild-type and Dicer<sup>-/-</sup> mESCs to determine whether their stability is miRNA regulated. These two sets included the high-confidence "Overlap" set, comprising 43 genes that passed the Normalization and Multi-Library filters, and a more inclusive "All" set, comprising 201 genes that passed the Normalization filter for any WT library. The log<sub>2</sub> fold expression change (LFC) between wild-type and Dicer<sup>-/-</sup> mESCs was compared to the LFC of a control set of genes that lacked the GCACUU motif. The Ago2-CLIP 3' UTR GCACUU-motif genes from both "Overlap" and "All" sets showed significantly more downregulation  $(P = 4.4 \times 10^{-6}, P = 7.73 \times 10^{-16}, \text{respectively})$  in wild-type mESCs relative to *Dicer<sup>-/-</sup>* mESCs, as compared to the control gene set (Fig. 2a–d; Supplementary Table 4 for statistics and Supplementary Table 5 for gene lists). These results independently support the hypothesis that these mRNAs physically bound to Ago2 are in vivo miRNA targets in mESCs.

Given that miRNA-dependent changes in mRNA expression have previously been shown for high-confidence predicted targets on the basis of computational analysis of conservation and context around the seed site (TargetScan 5.1; refs. 8,11,30), the properties of these predicted targets of the AAGUGC seed-related family were compared with those of the mRNAs identified by Ago2-CLIP. Comparison of expression levels in wild-type mESCs of Ago2-CLIP genes and predicted targets showed that the Ago2-CLIP 3' UTR GCACUU-motif genes tend to be more highly expressed (**Supplementary Fig. 3**). This is not surprising, as biochemical enrichment protocols tend to more effectively sample highly expressed genes.

To further compare properties of the predicted targets and CLIPidentified mRNAs other than expression level, two expression-matched and 3' UTR length-matched sets of predicted targets for the AAGUGCseed family were generated. The first set, "All predicted targets," contains 799 TargetScan GCACUU-containing predicted targets. Compared with this predicted set, both Ago2-CLIP "Overlap" and "All" gene sets showed significantly greater miRNA-dependent changes in expression (**Fig. 2a,c**; Overlap  $P = 4.00 \times 10^{-6}$ , All  $P = 1.12 \times 10^{-7}$ ), suggesting that the CLIP-identified mRNAs possess features in addition to the miRNA seed match requirement.

To assess the importance of conservation and context around the seed site, we created two gene sets ("Conserved predicted targets") containing the highest-confidence bioinformatically predicted targets, which are ranked first by branch length (that is, conservation) and then by context score (scored combinatorially by its site type, 3' pairing, local AU content and position within the 3' UTR<sup>8,11</sup>) (Fig. 2b,d). These two sets were comparable to the corresponding Ago2-CLIP "Overlap" and "All" sets in terms of gene number, expression level and 3' UTR length. No statistically significant difference in miRNA-dependent gene expression change was observed between the Ago2-CLIP "Overlap" gene set and the "All" gene set and their corresponding "Conserved predicted targets" sets (Fig. 2a,c). Yet the CLIP-identified GCACUU sites from the "Overlap" and "All" sets are generally less conserved and surrounded by a relatively less favorable sequence context than the "Conserved predicted targets" (Fig. 2b,d and Supplementary Table 4d-f). Taken together, our results suggest







(conservation) and then by context score (a, top-ranked 43 genes; c, top-ranked 201 genes); "All predicted" expression-matched TargetScan GCACUU targets (light pink, 1,469 genes); and "Control" mRNAs (gray). (e) Gene sets include Ago2-CLIP CDS GCACUU-containing mRNAs that passed the Normalization filter in any WT library ("CLIP", blue); "Predicted" GCACUU-containing CDS mRNAs (pink) that were expression-matched to the "CLIP" set but lacked GCACUU in their 3' UTRs; and matched "control" mRNAs (gray). (f) Gene sets include Ago2-CLIP 3' UTR GCACUU-containing (blue) and G motif-containing mRNAs (red) that passed the Normalization and Multi-Library filters in WT libraries; Ago2-CLIP 3' UTR CCAGCC-containing mRNAs (orange) that passed the Normalization filter in at least one WT library; and corresponding matched controls (gray). Any mRNAs with GCACUU in the 3' UTR or CDS were removed from the "G motif" and "CCAGCC" sets. P < 0.01 for all significant differences. Error bar, s.d.

that the "All" set and the smaller "Overlap" gene sets represent highconfidence sets of miRNA-regulated mRNAs and that there are other factors beside conservation and context around the GCACUU seed motif that govern which sites are targeted by miRNAs and/or are bound by Ago2 in mESCs.

We also sought to determine whether the GCACUU motifs identified in CDS were associated with a miRNA-dependent gene expression signature. To this end, expression of Normalization filtered Ago2-CLIP CDS GCACUU-motif genes from all WT libraries (excluding those with GCACUU in the 3' UTR) was compared to the expression of a set of controls that lack GCACUU in the CDS. The 80 Ago2-CLIP CDS GCACUU-motif genes showed miRNA-dependent downregulation in mRNA expression as compared with the control set (Fig. 2e). Notably, other expression-matched CDS GCACUU-motif genes ("Predicted set," Fig. 2e) showed a profile similar to that of the Ago2-CLIP identified set and a significant downregulation ( $P = 1.10 \times 10^{-3}$ ) compared with the control. This indicates that the presence of the GCACUU motif in CDS, as in the case of the 3' UTR, is associated with a miRNA-dependent gene expression signature<sup>8,12</sup>.

The expression profile difference between wild-type and Dicer-/mESCs was further examined for genes with the G-rich motif, whose association with Ago2 appears to be miRNA independent, and with the CCAGCC motif, whose association with Ago2 might be miRNA dependent. Neither the G-rich motif nor the CCAGCC motif is complementary to any miRNA sequence with appreciable expression in mESCs. We compared those CCAGCCcontaining genes that passed the Normalization filter (excluding those containing GCACUU) with expression-matched sets of all mouse genes that do not contain CCAGCC (control) in the 3' UTR (Fig. 2f). Unexpectedly, we observed a significant downregulation ( $P = 2.60 \times 10^{-5}$ ) of gene expression for the CCAGCC-containing genes in wild-type mESCs relative to Dicer<sup>-/-</sup> mESCs. This expression difference seems to be specific to those Ago2-CLIP CCAGCCcontaining mRNAs, as other mRNAs containing CCAGCC did not

show a similar change ("Predicted set" in Supplementary Fig. 4). For the G-rich motif from Figure 1d, we compared the expression change for Ago2-CLIP genes that contain matches to this motif, but lack GCACUU in their 3' UTRs, and that passed the Normalization and Multi-library filters (Fig. 2f) with a set of 3' UTRs randomly chosen from the mouse genome that was matched for expression level, dinucleotide CG composition and 3' UTR length. As is the case for GCACUU- and CCAGCC-containing genes, a significant increase  $(P = 2.20 \times 10^{-16})$  in gene expression was observed upon deletion of Dicer for these G motif-containing genes identified by Ago2-CLIP. Such observed change could be due to the presence of other miRNA seed matches in the 3' UTRs of Ago2-CLIP G-rich motif genes. However, excluding those G-rich motif genes harboring seed matches to abundant mESC miRNAs did not affect the aggregate gene expression change of the G-rich motif gene set (Supplementary Fig. 4).

Notably, the degrees of change in mRNA expression observed for G motif- or CCAGCC-containing genes were not significantly different from those observed for the Ago2-CLIP GCACUU-motif genes (Fig. 2f) and their expression-matched predicted GCACUU set (compare Fig. 2a). Correlated with this, previous data suggests that the effect of miRNAs can be mimicked by miRNA-independent tethering of Argonaute proteins to reporter mRNAs<sup>31,32</sup>. Thus, the observed miRNA-dependent expression changes for Ago2-CLIP genes could be due to the close proximity between Ago2 and the cross-linked mRNA targets.

GCACUU-containing clusters confer miRNA-mediated repression Next, we sought to determine whether the GCACUU-containing regions that cross-linked to Ago2 are sufficient to confer miRNAdependent repression on luciferase reporter transgenes in the presence of endogenous levels of the corresponding miRNAs. Because only four genes<sup>33-36</sup> have been validated as GCACUU seed match targets in mESCs, it was difficult to evaluate our dataset with the existing literature. Instead, we inserted the ~80-nt Ago2-CLIP cluster sequence into the 3' UTR of luciferase and compared the expression of this construct **Figure 3** Ago2-CLIP-identified GCACUU motif-containing cluster is sufficient to confer miR-295-mediated repression. (**a**-**c**) Box plots of relative activity of *Renilla* luciferase transgenes bearing CLIP clusters plus 25-nt flanking regions in the 3' UTR. All raw *Renilla* luciferase values were first normalized to values from firefly luciferase transfection control. Error bars are s.d. (**a**) Exemplary plot of a luciferase reporter assay testing an Ago2-CLIP 3' UTR cluster, *Slc31a1* plus 25-nt flanking regions at both ends. Shown are box plots for the ratio of the expression level of a luciferase reporter containing a wild-type *Slc31a1* CLIP cluster in



its 3' UTR to an identical luciferase reporter in which the GCACUU seed was mutated to CCUCAU. This ratio is defined as "WT/seed mutant activity" and calculated in three different cellular states: *Dicer<sup>-/-</sup>* mESCs (black), wild-type mESCs (white) and *Dicer<sup>-/-</sup>* mESCs transfected with miR-295 mimic (gray). For panels **b**,**c**, "Relative luciferase activity" is defined as "WT/seed mutant activity" in specific cellular state normalized to the corresponding value in *Dicer<sup>-/-</sup>* mESCs. Genes that passed both the *Normalization* and *Multi-library filters* (that is, the "Overlap" set) are marked with an asterisk. All remaining genes passed the *Normalization filter* in one WT library. All box plots are from at least three independent experiments. (**b**) Relative luciferase activity of CLIP-identified 3' UTR clusters plus 25-nt flanking regions in wild-type mESCs (gray) or *Dicer<sup>-/-</sup>* mESCs transfected with miR-295 mimic (white). Relative luciferase activity is significantly <1 in all genes examined (*P* < 0.05, two-tailed paired *t*-test). (**c**) Relative luciferase activity of CLIP-identified CDS clusters plus 25-nt flanking regions in wild-type mESCs (gray) or *Dicer<sup>-/-</sup>* mESCs transfected with miR-295 mimic (white). *Herc1* and *Eif2c2* each have two GCACUU motifs. Relative luciferase activity is significantly <1 for all genes examined except *Tnrc6a* (*P* < 0.05, two-tailed paired *t*-test).

to that of an equivalent construct with the GCACUU motif mutated to CCUCAU. The ratio of wild-type to mutant construct expression was evaluated in three cellular states: (i) wild type (endogenous miRNA levels), (ii) Dicer<sup>-/-</sup> mESCs (no mature miRNAs) and (iii) Dicer<sup>-/-</sup> mESCs transfected with a miR-295 mimic (Fig. 3a). In each cellular state, the relative repression was calculated by normalizing to the ratio in *Dicer<sup>-/-</sup>* cells. We found that eight out of eight Ago2-CLIP 3' UTR GCACUU motifs showed significant miRNA-dependent repression (P < 0.05) in wild-type cells but not in *Dicer<sup>-/-</sup>* cells (Fig. 3b). However, the repression in Dicer-/- cells was restored by addition of a miR-295 mimic, suggesting that a member from this mESC-specific miRNA cluster (with AAGUGC seed) is sufficient to provide the specificity for such regulation. Additionally, Ago2-CLIP-identified binding sites were present in three genes that have previously been shown to be regulated by the AAGUGC-related miRNA family (E2f1<sup>37</sup>, Pten<sup>38</sup> and Cdkn1a<sup>33</sup>). These data show that the Ago2-CLIP 3' UTR-bearing GCACUU motif sites are indeed endogenous targets for direct regulation by miRNAs in mESCs, and the short fragment of ~80 nt containing such sites is sufficient to confer mESC-specific miRNA-mediated repression through miR-290-295.

To determine whether the Ago2-CLIP CDS GCACUU-motif sites are sufficient for miRNA regulation, we inserted the CDS cluster sequence (~80 nt), or a seed mutant equivalent, in the 3' UTR of luciferase. Seven out of eight clusters containing CDS GCACUU motifs conferred downregulation on the luciferase reporter (**Fig. 3c**), suggesting that these sequences are recognized by the endogenous miRNA machinery even in the heterologous context of the 3' UTR.

#### miRNA regulation can be modulated by G-rich motifs

The enrichment of the G-rich motif in Ago2-CLIP sequences from both wild-type and *Dicer<sup>-/-</sup>* mESCs (**Fig. 1d**) suggests that the motif is likely a miRNA-independent binding site for Ago2. This binding preference has not previously been described for Ago2, so we used an independent method to confirm the miRNA-independent association of Ago2 with the set of the G-rich motif–containing mRNAs. We transfected *Dicer<sup>-/-</sup>* mESCs with a hemagglutinin (HA)-tagged Ago2 construct, immunoprecipitated Ago2 with anti-HA antibodies, isolated the bound mRNA and hybridized it to Affymetrix microarrays. We also performed microarray analysis on total RNA from *Dicer<sup>-/-</sup>* mESCs. The enrichment of mRNAs in the Ago2 immunoprecipitation (Ago2-IP) was determined by comparing expression values between Ago2-IP and total RNA. We then determined whether the set of genes enriched in the Ago2-IP from *Dicer<sup>-/-</sup>* mESCs significantly overlapped with the sets of Ago2-CLIP G motif-containing genes. We found that 1.6–2.1-fold more genes overlapped between the Ago2-IP set and the Ago2-CLIP G-motif set than expected by chance (**Supplementary Table 6a**). These data support the observation that the G motif-containing genes identified by Ago2-CLIP are likely bound to Ago2 or its associated protein complex in a miRNAindependent manner in mESCs.

We previously determined that the CLIP-identified 3' UTR GCACUU mRNAs have a miRNA-dependent expression change comparable to those of the high-confidence predicted targets, despite being less conserved and in a less favorable sequence context (**Fig. 2a-d**). To explore whether the G motif is a feature in these sequences that can contribute to miRNA-dependent regulation, we focused on a new miRNA target, *Txnip*, identified in this study. We validated this target by demonstrating that its endogenous mRNA and protein levels are both regulated by *Dicer* in mESCs (**Fig. 4a**), similarly to previously validated *miR-290-295* target, *Cdkn1a*<sup>33</sup>. One of the Ago2-CLIP clusters identified in *Txnip* was among the most repressed in our 3' UTR luciferase assay (**Fig. 3b**). This provides a good range of sensitivity to test whether neighboring G-rich motifs affect the miRNA-dependent activity of GCACUU seed sites (**Fig. 4b**).

The relationship between this G-rich motif and the seed motif was investigated as in **Figure 3** using the following luciferase constructs: (i) the WT cluster, (ii) the GCACUU seed motif mutated to CCUCAU, (iii) a mutant G-rich motif where all Gs are mutated to Cs (so as not to alter AU content<sup>8,10</sup>) or (iv) both these motifs mutated. For the Txnip "A" cluster, repression was the strongest with the wild-type GCACUU seed motif and G motif (**Fig. 4c**). The repression was relieved by 50% when the G motif was mutated; however, in the absence of GCACUU, the presence of the G motif alone did not confer repression (**Fig. 4c**). We extended this analysis by investigating the contributions of G-rich motifs to miRNA-dependent repression in another cluster from *Txnip*, Txnip "B" (**Fig. 4d**), and a cluster from *Ei24* (**Fig. 4e**). These clusters each have multiple G-rich motifs; mutation of each G-rich motif individually had varying impact on repression by the GCACUU seed site, with deletion of all G-rich motifs having the

## RESOURCE

Figure 4 G-rich motif modulates miRNAmediated repression. (a) Western blot of two GCACUU-containing targets, Txnip and Cdkn1a, in wild-type mESCs (WT), Dicer-I- mESCs (KO) (left) and Dicer-I- mESCs transfected with miR-295 mimic at different concentrations (nM): 0 (control), 10, 100 (right). Tubulin loading control and *Dicer* genotype control are also shown. (c) Mutation of the Txnip "A" G-rich motif reduces GCACUU seed-mediated repression. Luciferase reporter constructs were created as in Figure 3, using an Ago2-CLIP cluster, plus 25-nt flanking regions, from the Txnip 3' UTR. Along with the GCACUU seed mutant construct, a G-motif mutant construct with all the Gs in the G motif changed to Cs as well as a construct with both the seed and G motif mutated were created. The presence or absence of an intact GCACUU seed match or G motif in the construct is indicated by + (WT) or - (mutated). Relative luciferase activity was calculated as in Figure 3 as the expression level of each indicated construct relative to the seed- G motif- double mutant construct in either WT (left, blue) or KO mESCs transfected with miR-295 mimic (right, green), normalized to the same ratio in KO mESCs. Values shown are averages of three independent experiments, and P values < 0.05 were deemed significant and marked with asterisks. (b,d,e) Genomic schematic of Ago2-CLIP 3' UTR mapping clusters. The following features are depicted:



the Ago2-CLIP cluster sequence (thick black bar) with flanking regions (thin black bar), octamer G-rich motif(s) in the cluster (green bar and sequence in green to the right; Supplementary Methods for details), predicted miR-290-295-cluster GCACUU-binding site (purple) using multiple algorithms including TargetScan 5.1 (ref. 11), PITA<sup>54</sup> and the UCSC genome browser 30-Way Mammalian Multiz alignment and conservation<sup>39,55</sup> (blue; heights indicate degree of conservation at aligned position). Clusters depicted are Txnip "A" (b) and "B" (d) clusters from Txnip 3' UTR, and Ei24 cluster (e) from Ei24 3' UTR. All G-rich motifs shown match a consensus G-rich motif P < 0.05; see Supplementary Methods for details. (f) Mutation of G-rich motifs in Ago2-CLIP clusters reduces repression conferred by three independent miR-290 family seed sites. Similar data as in c but plotting the amount of seed-mediated repression on the luciferase reporter in the presence (+) or absence (-) of the G motif in either WT (blue) or KO mESCs transfected with miR-295 mimic (green), normalized to the same ratio in KO mESCs. 100% full seed repression was set as 1 minus the ratio of the expression levels of the WT construct to the seed mutant construct (G motif +). The % full seed repression in the absence of the G motif was calculated as 1 minus the ratio of the expression levels of the G-motif mutant construct to seed mutant, G-motif mutant construct (G motif-) as a percent of 100% full seed repression (as described above). For Txnip "B" and Ei24 clusters, the G motifs "+" or "-" correspond to same motifs shown left to right in the schematics (d,e). Values shown are averages of three independent experiments, and P values <0.05 deemed significant are marked wth asterisks. (g) Average phastCons score was determined for each G motif (match score > 0, data for Normalized Ago2-WT1A shown in black) along with a constrained background set of random octamers drawn from annotated mouse 3' UTRs (gray) (see Supplementary Methods for explanation of background derivation). On the x axis are bins for phastCons score cutoffs,  $\geq 0.25$ ,  $\geq 0.50$ ,  $\geq 0.75$  and  $\geq 0.90$ . On the y axis is the percentage of total G motifs or random octamers that are in each phastCons score cutoff bin. Ago2-CLIP G motifs are on average more conserved than random octamers (P values as indicated, Fisher's exact test). Error bar, s.d.

greatest effect (**Fig. 4f**). Similar loss of miRNA-dependent repression was also observed when the G-rich motif of the Txnip "A" cluster and Ei24 cluster was deleted (**Supplementary Fig. 5a**). Taken together, these data suggest that the G-rich motif is important for the full activity of the miRNA seed site but does not contribute activity in the absence of the miRNA seed site.

Given that the CLIP-identified G-rich motif modulated the miRNAmediated repression in the three clusters examined, we further investigated the general features of this motif, including its composition, conservation and location within the Ago2-associated sequence. We searched for enrichment of shorter motifs in the 3' UTR clusters and found that the original octamer G motif is composed of enriched G-rich tetramers and pentamers (**Supplementary Fig. 5b**). Next, we analyzed the conservation of the octamer G motif. We determined an average conservation score for all G motifs based on the phastCons conservation<sup>39</sup> of each nucleotide within the motifs and compared with a background set of octamers (**Supplementary Methods**). We found that G motifs are generally more conserved than random sequences  $(P < 10^{-6})$  (**Fig. 4g**). We also analyzed nucleotide positional conservation of an alignment of G motifs with 10-nucleotide (nt) flanks at either end. The level of conservation decreased immediately after the 3' end of the motif, whereas the higher level of conservation persisted in the 10 nt 5' of the motif (**Supplementary Fig. 5c**). Notably, further MEME analyses suggest that the octamer G motif is likely to be embedded in an extended G motif (**Supplementary Fig. 5b**). The excess conservation observed for G motifs was true for all 3' UTR clusters, including those lacking the GCACUU motif (**Supplementary Fig. 5d**). Thus, the excess conservation of the G motif is not a bystander effect from being near this particular miRNA seed match; rather, the G motif has attributes of a functional regulatory element<sup>40</sup>.

Another common feature of this G motif is that it tends to be present in the 5' half of the sequence that is cross-linked to Ago2 (**Supplementary Fig. 5e,f**). In contrast, there is no positional bias for the GCACUU motif (**Supplementary Fig. 5f**). In cases where both motifs are present in the Ago2-CLIP sequences, there are no biases as to whether the G motif is 5' or 3' of the GCACUU motif (data not shown). The activity of the examined G motifs is independent of its location relative to GCACUU (compare **Fig. 4b,d,e**), suggesting that the vicinity, rather than the directionality, is important for modulating miRNA repression.

#### DISCUSSION

Photo-cross-linking followed by Ago2 immunoprecipitation, Ago2-CLIP, was used to identify miRNA binding sites in mESCs. We found significantly enriched motifs in 3' UTRs and CDS that correspond to miRNA seed matches, representing 201 and 103 potential mESC miRNA targets in 3' UTRs and CDS, respectively. In regard to the latter point, this study is in agreement with other studies indicating that the presence of miRNA binding sites in CDS is more widespread than has been previously considered and nearly as prevalent as in 3' UTRs<sup>14–16</sup>. Here we provided gene expression data suggesting that these CDS sites regulate mRNA stability much like 3' UTR sites. Moreover, these sites can be recognized by miRNAs at endogenous expression levels and confer repression in a heterologous 3' UTR<sup>8,13,41,42</sup>.

Two other Ago-CLIP studies have identified potential miRNA targets in mammalian cells and tissue. Our study differed from those in that we analyzed mRNAs associated with endogenous Ago2 in a mostly homogenous cell population of mESCs, whereas Chi *et al.*<sup>15</sup> performed CLIP on brain extracts using endogenous Ago antibodies, and Hafner *et al.*<sup>14</sup> performed CLIP in 293 cells using HA-tagged Ago1–Ago4 and cross-linking by a photoactivatable nucleotide. Independently of the variations in the CLIP technique and data analysis, these studies, as well as ours, identified similar numbers of targets for each miRNA seed family (several hundred), which is comparable to the number of moderately conserved targets predicted for each miRNA seed family by TargetScan<sup>11</sup> (see **Supplementary Notes** for cross-comparison with other CLIP datasets).

There are several previously published reports of miRNA-regulated mRNAs in mESCs that we could compare to the Ago2-CLIP 3' UTR GCACUU-containing genes<sup>35,43</sup>. Only miR-294 (a member of the AAGUGC seed family)-regulated mRNAs described by Melton *et al.*<sup>43</sup> showed significant overlap with the Ago2-CLIP 3' UTR GCACUU-containing mRNAs (see **Supplementary Table 6b** for all comparisons).

Unlike other cell types, including those used in other Ago2-CLIP studies<sup>14,44</sup>, mESCs appear to be dominated by a single miRNA seed family that is probably responsible for most of the miRNA regulation in this cell type. Essentially all of the GCACUU motif–containing CLIP 3' UTR clusters conferred miRNA-dependent regulation when tested in luciferase reporter assays in the presence of endogenous levels of miRNAs. This suggests that the stringency of our filtering criteria resulted in selection of a high-confidence set of GCACUU-containing mRNAs that most likely are bona fide miRNA targets in mESCs. Previous studies have already shown that this miRNA family has important roles in mESCs, including maintaining pluripotency, self-renewal and cell cycle control<sup>33–35,43,45</sup>. But few targets have been identified and validated. This study identifying a few hundred new miRNA targets through Ago2-CLIP is a significant step in the exploration of this biology.

To understand the extent of miRNA-regulated pathways represented by the Ago2-CLIP 3' UTR GCACUU-motif genes ("All" set, 201 genes), we performed pathway enrichment analysis (**Supplementary Methods**) and compared this set with the top 201 "Conserved predicted targets" and all mRNAs expressed in mESCs that contain GCACUU hexamer in the 3' UTR ("All predicted targets," 2969 genes). We found 37 and 11 pathways to be significantly enriched in the CLIP and "Conserved predicted targets" sets, respectively (**Supplementary Fig. 6a** and **Supplementary Table 7**). The pathways significantly enriched in CLIP included "Early S-phase" (four genes), a pathway in which *miR-290–295* has been previously implicated<sup>33</sup>, and "TGF-beta receptor signaling" (five genes), a pathway in which *miR-290–295* has not been implicated.

The genes identified by Ago2-CLIP in the "TGF-beta receptor signaling" pathway (P = 0.013) include two intracellular pathway inhibitors, the cytoplasm-localized Smad7 and the nucleus-localized Skil, and an extracellular inhibitor, Lefty1. Our reporter assay confirmed that these three genes are indeed targeted by miR-295 (Supplementary Fig. 6b). We extended this analysis to Lefty2, a gene that was not identified in the CLIP results but that is homologous to Lefty1 and contains the GCACUU hexamer, and showed that it is also targeted by miR-295 (Supplementary Fig. 6b). Correlated with this, miR-302 and miR-430, which are related in miRNA seed to miR-290-295, have been shown, respectively, in human ESCs<sup>46</sup> and zebrafish embryos<sup>47</sup> to regulate differentiation through targeting Lefty homologs. Here, using a genome-wide approach, we found that the miR-290-295 regulates not only the extracellular Lefty homologs but also additional inhibitory nodes of the TGFβ pathway localized in different cellular compartments (Supplementary Fig. 6c). This coordinate inhibition, as observed for other miRNAs<sup>48–50</sup>, might confer robustness in this signaling network.

We unexpectedly identified a G-rich motif in most of the sequences associated with Ago2 regardless of the miRNA status in the cell. We believe this is a true biological association, rather than a technical artifact, on the basis of the following observations. First, this motif is conserved above the general 3' UTR background even when matched for sequence content. Second, the genes containing this G-rich motif have significant overlap with the set of genes enriched in HA-tagged Ago2 immunoprecipitates from *Dicer*<sup>-/-</sup> mESCs. Third, we observe G bias only in genic sequences and not in miRNA or intergenic sequences cross-linked to Ago2. Lastly, the enrichment of G residues is not likely to be due to CLIP itself as there are no described G biases in the literature for any of the steps involved (**Supplementary Notes** for further discussion).

Yet it remains unclear whether cross-linking to this G-rich sequence is due to Ago2 itself or to a binding partner of Ago2. Given that UV cross-linking forms covalent bonds between protein and RNA molecules that are within angstroms of each other, a potential binding partner would have to be in close proximity to Ago2 and the mRNA target. Indeed, several proteins that co-immunoprecipitate with Ago2 have binding preference for G-rich sequences, including HNRNP-H and FMRP<sup>51-53</sup>, although we observed only one Ago2-dependent RNA-protein complex close to the molecular weight of native Ago2 in the CLIP procedure. Alternatively, Ago2 itself could have a previously unidentified preference for binding G-rich sequences. In either case, when a G-rich sequence occurs near a miRNA binding site, it could give the Ago2-miRNA complexes a higher affinity for this region and thus lead to an increased probability that the mRNA would be targeted for degradation and/or inhibition of translation. In three cases examined, this G-rich motif modulates the level of miRNA-dependent regulation by the miR-290-295-related seed motif but imparts no regulation by itself. Therefore, identification of this association indicates the value of Ago2-CLIP data from Dicer<sup>-/-</sup> mESCs as an invaluable background in delineating bona fide microRNA targets.

#### METHODS

Methods and any associated references are available in the online version of the paper at http://www.nature.com/nsmb/.

Accession codes. Microarray and short RNA sequence files have been deposited at the Gene Expression Omnibus database repository under accession number GSE25310. BED files for clusters in all libraries are available for download from http://rowley.mit.edu/pubs/Ago2\_CLIP/Ago2\_CLIP.html.

*Note: Supplementary information is available on the Nature Structural & Molecular Biology website.* 

#### ACKNOWLEDGMENTS

We thank C. Burge, J. Wilusz, A. Ravi and members of the Sharp laboratory for critical comments, M. Lindstrom for illustration, T. Cybulski for technical help, A. Leshinsky and R. Cook for running the Solexa sequencing samples in the KI Biopolymer & Proteomics Core Facility, M. Luo and L. Smeester for microarray technical assistance in the MIT Department of Biology Biomicrocenter and C. Whittaker for bioinformatics support in the Bioinformatics & Computing Core Facility at the Koch Institute. A.K.L.L. was supported by a special fellowship from the Leukemia and Lymphoma Society. A.G.Y. was partially supported by a David H. Koch graduate fellowship. This work was supported by United States Public Health Service grants R01-GM34277 and R01-CA133404 from the US National Institutes of Health, P01-CA42063 from the National Cancer Institute to P.A.S. and partially by Cancer Center Support (core) grant P30-CA14051 from the National Cancer Institute.

#### AUTHOR CONTRIBUTIONS

A.K.L.L. and A.G.Y. designed and performed the experiments; A.K.L.L., A.G.Y. and P.A.S. wrote the paper; A.D.B. performed experiments; A.B., C.B.N. and G.X.Z. performed the bioinformatics analyses. All authors reviewed and approved the manuscript.

#### COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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#### **ONLINE METHODS**

**Cross-linking and immunoprecipitation followed by sequencing.** We performed two independent experiments from WT and *Dicer<sup>-/-</sup>* mESCs<sup>18</sup>. 10E7 cells were plated and 48 h later irradiated once with 4,000  $\mu$ J cm<sup>-2</sup>. For immunoprecipitation, anti-mouse Ago2 monoclonal antibody (lot no. PEM0820, Wako; cat. no. 018-22021) was used. CLIP was performed as described<sup>56</sup> except that the 3' linker was not ligated 'on bead'. Another short RNA library was prepared from WT mESC total RNA. Total RNA and CLIP libraries were cloned and sequenced using the Illumina method as described previously<sup>57</sup>. Libraries "WT1A" and "WT1B" are separate PCR amplifications and sequencing runs from the same CLIP RNA sample, and "WT2" is an independent biological replicate. For further details, see **Supplementary Methods**.

Sequence processing and cluster generation. Reads were processed for linkermatching criteria and mapped to the mouse genome (mm9), miRbase (release 14) and ncRNA annotations (fRNAdb ver 3.4, Genomic tRNA Database, Rfam version 9.1). For unique RNA reads, not mapping to miRNA or noncoding RNA, identical reads were collapsed to a single observation, overlapping reads were aggregated into clusters and 25 nt of flanking sequence was added to either side of the cluster. Clusters were subjected to a series of filters: *Normalization filter*, similar to process of ref. 15; *Multi-library filter*, clusters from biological replicates with overlapping genomic coordinates were selected; and *Knockout filter*, those WT clusters that had overlapping genomic coordinates with a cluster from a *Normalized* KO library were excluded. For further details, see **Supplementary Methods**.

**Motif analysis.** Two motif-finding approaches were used: (i) MEME<sup>26</sup> for motif discovery in the aggregate set of 3' UTR– and CDS-mapping WT and KO library clusters, and (ii) an enumerative approach for analysis of significantly enriched hexamers within each individual library. We measured the statistical significance of the occurrence of *n*-mer sequences within each library compared to their occurrence in sequences drawn randomly given a background distribution. Significance was determined by *P* value, to denote enrichment over background, and *z* score, to quantify the magnitude of the enrichment. Plots made with Numbers and Photoshop. For further details, see **Supplementary Methods**.

**Gene expression analysis.** We prepared three independent sets of RNA from WT and *Dicer<sup>-/-</sup>* mESCs, labeled as per manufacturer's instructions (Affymetrix) and hybridized to Affymetrix Mouse Exon 1.0 ST arrays. Differential expression was determined as previously described<sup>10</sup>. CLIP gene sets were derived for different motif and filter combinations. 3' UTR Predicted targets for GCACUU mRNAs were derived from TargetScan 5.1 using branch length and context scores and expression-matched to CLIP set using WT microarray data as indicated. CDS GCACUU and CCAGCC predicted targets were selected from all mouse mRNAs containing the indicated hexamers and expression matched to CLIP. Controls and statistics were generated similarly to ref. 10. Plots were made in R (http://www. r-project.org/). For further details, see **Supplementary Methods**.

Luciferase assays. 3' UTR and CDS clusters plus 25-nt flanking regions were PCR-amplified from mouse genomic DNA and cloned into the 3' UTR of a cytomegalovirus promoter (CMV)-*Renilla* luciferase vector. Seed mutant versions were created by mutating GCACUU to CCUCAU. G-motif mutants were created by mutating Gs to Cs or by deleting the motif as indicated. Constructs were transfected into WT mESCs, *Dicer*<sup>-/-</sup> mESCs and *Dicer*<sup>-/-</sup> mESCs with 100 nM miR-295 mimic (Dharmacon) and luciferase assays were performed similarly to

those in reference 58. Statistical significance was determined using two-tailed paired *t*-test (P < 0.05). Plots made with Numbers and KaleidaGraph. For further details, see **Supplementary Methods**.

Western blots. 2E6 WT mESCs were plated in 10-cm dishes 48 h before lysis with 300  $\mu$ l of modified RIPA buffer (see **Supplementary Methods** for recipe). 24 h before transfection of 10 nM or 100 nM miR-295 mimics using Lipofectamine 2000 (Invitrogen), 2E6 *Dicer<sup>-/-</sup>* mESCs were seeded in a T-25 flask. 24 h after transfection, the cells were replated to 10-cm dishes; a further 24 h later, cells were lysed with 300  $\mu$ l of modified RIPA buffer. 30  $\mu$ g of lysates were subjected to SDS-PAGE and proteins transferred to nitrocellulose. Primary antibodies were to Txnip (MBL), tubulin (Abcam) and Dicer (Bethyl); appropriate horseradish peroxidase–conjugated anti-mouse or anti-rabbit secondary antibodies were from GE; and western blots were developed using Western Lightning ECL plus (Perkin Elmer).

**G-motif analysis.** The G motif was defined as the octamer G motif reported by  $MEME^{26}$  ( $E < 10^{-385}$ ) using *Normalized* KO1 clusters that overlapped with at least one WT library. MAST<sup>59</sup> was used to identify G motifs in Ago2-CLIP 3' UTR mapping clusters with the cutoff score as indicated. Conservation was determined using phastCons scores for the mouse genome (mm9) from the UCSC genome alignments (http://genome.ucsc.edu)<sup>39</sup>. Motif hits where fewer than two nucleotides had a nonzero positional alignment or score were removed. Background conservation of 3' UTR octamers was estimated by randomly selected octamers, filtered by G content, from UCSC mouse 3' UTR annotations, and the scores from three sets of 2,500 octamers were averaged and analyzed similarly to the G-motif octamer hits. Plots were made with Numbers and Photoshop. For further details, see **Supplementary Methods**.

**Microarray analysis of mRNAs in Ago2 immunoprecitates from** *Dicer<sup>-/-</sup>* **mESCs.** *Dicer<sup>-/-</sup>* mESCs were transfected with pCAGGS-FLAG/HA-hAgo2 using Lipofectamine 2000 (Invitrogen). 48 h after transfection, cells were lysed, and some of the lysate was used for total RNA extraction by Trizol (Invitrogen) following manufacturer's protocol. The remaining lysate was incubated with anti-HA beads (Sigma) for 2 h at 4 °C, then beads were washed and the associated (IP) RNA was extracted by Trizol (Invitrogen) following manufacturer's protocol. Two biological replicates of Ago2 IP and total RNA were prepared for and hybridized to Affymetrix Mouse 430A 2.0 microarrays according to manufacturer's instructions. Differential expression between IP and total RNA populations was determined similarly to previously described<sup>10</sup>.

**Pathway analysis.** For pathway analysis, we used MetaCore from GeneGO Inc. Each of the gene sets was analyzed individually for significantly enriched pathways represented in the set using an FDR < 0.57 cutoff. All annotated mouse genes are used as background for "All TargetScan GCACUU predicted targets" set. "All GCACUU predicted targets" set was used as background for Ago2-CLIP GCACUU-motif "All set" and the set of TargetScan top 201 genes (ranked by branch length, and context score) with expression on microarray > 4. For further details, see **Supplementary Methods**.

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## Corrigendum: Genome-wide identification of Ago2 binding sites from mouse embryonic stem cells with and without mature microRNAs

Anthony K L Leung, Amanda G Young, Arjun Bhutkar, Grace X Zheng, Andrew D Bosson, Cydney B Nielsen & Phillip A Sharp Nat. Struct. Mol. Biol. 18, 237-244 (2011); published online 23 January 2011; corrected after print 12 May 2011

In the version of this article initially published, the blue curve in Figure 2c was mistakenly replaced with a duplicate of that in Figure 2a. The error has been corrected in the HTML and PDF versions of the article.

