miRNA–target chimeras reveal miRNA 3′-end pairing as a major determinant of Argonaute target specificity

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microRNAs (miRNAs) act as sequence-specific guides for Argonaute (AGO) proteins, which mediate posttranscriptional silencing of target messenger RNAs. Despite their importance in many biological processes, rules governing AGO–miRNA targeting are only partially understood. Here we report a modified AGO HITS-CLIP strategy termed CLEAR (covalent ligation of endogenous Argonaute-bound RNAs)-CLIP, which enriches miRNAs ligated to their endogenous mRNA targets. CLEAR-CLIP mapped ~130,000 endogenous miRNA–target interactions in mouse brain and ~40,000 in human hepatoma cells. Motif and structural analysis define expanded pairing rules for over 200 mammalian miRNAs. Most interactions combine seed-based pairing with distinct, miRNA-specific patterns of auxiliary pairing. At some regulatory sites, this specificity confers distinct silencing functions to miRNA family members with shared seed sequences but divergent 3′-ends. This work provides a means for explicit biochemical identification of miRNA sites in vivo, leading to the discovery that miRNA 3′-end pairing is a general determinant of AGO binding specificity.
miRNAs (miRNAs) are small, non-coding RNAs that mediate posttranscriptional RNA silencing by sequence-specific targeting of Argonaute (AGO) proteins to miRNAs. miRNAs regulate the development, homeostasis and pathologies of virtually all vertebrate tissues. Many miRNAs have specific or enriched expression in the central nervous system, regulating such diverse processes as neuronal differentiation, excitation, synaptogenesis and plasticity. Accordingly, miRNA dysregulation is implicated in neurological disorders and many cancers including glioma and liver cancer. However, miRNA function in these contexts remains unclear, as in vivo miRNA targets are unknown.

Accurate miRNA target identification remains a formidable challenge. Canonical miRNA binding involves base pairing of the miRNA seed region (nucleotides 2–8) to complementary target sites. Such short motifs occur frequently in the transcriptome and are not sufficient to predict miRNA binding, leading to high false discovery rates for purely bioinformatic predictions. To mitigate this limitation, evolutionary conservation and local AU sequence content are employed as screens for site functionality and accessibility, respectively. However, the importance of non-conserved miRNA regulation, especially in the brain, and limitations of context predictions without empirical binding information are well established. Moreover, the assumption of uniform rules for all miRNAs ignores non-canonical miRNA binding, increasingly recognized as widespread. Rules beyond seed-based pairing such as in vivo-end auxiliary pairing have been described but are generally considered rare. Other non-canonical binding modes include 3′-end centric ‘seedless’ pairing, centred miRNA pairing and nucleation bulges in the seed region.

Empirical mapping of miRNA target sites in vivo was first achieved with ultraviolet cross-linking and immunoprecipitation with high-throughput sequencing (HITS-CLIP) of AGO proteins. AGO HITS-CLIP generates two data sets—a transcriptome-wide target binding map and an empirical catalogue of expressed miRNAs—that empower accurate identification of functional miRNA-binding sites. However, the inability to link miRNA and target unambiguously remains a limitation. Two groups reported experimental strategies to ligate miRNA to target RNA in purified AGO complexes. CLASH (cross-linking and sequencing of hybrids) identified thousands of miRNA–target chimeras using dual-tagged AGO1 in HEK-293T cells, revealing frequent seed-independent miRNA binding. Soon after, modified photoactivatable ribonucleoside-enhanced CLIP identified ~3,600 unambiguous events in Caenorhabditis elegans. Although identifying thousands of novel interactions, the reliance of these studies on exogenous AGO expression excludes them from analysis of human tissues and, currently, in vivo mouse models, and raises concerns about the stoichiometry of RNA-binding events.

We have developed modifications of AGO HITS-CLIP, termed CLEAR (covalent ligation of endogenous Argonaute-bound RNAs)-CLIP, permitting isolation of miRNA–target chimeras from endogenous AGO–miRNA–mRNA complexes. CLEAR-CLIP identifies tens of thousands of miRNA target sites in mouse brain including novel targets for many neuron-specific miRNAs. In mouse brain and human liver cells, we defined expanded pairing rules for over 200 mammalian miRNAs illustrating widespread use of miRNA 3′-end auxiliary pairing in vivo and tolerance of diverse, although constrained, pairing patterns for many miRNAs. Integrated with HITS-CLIP binding information, CLEAR-CLIP provides an improved empirical basis for identification of physiologic canonical and non-canonical miRNA regulation.

Results

CLEAR-CLIP defines miRNA–target interactions in vivo. We modified AGO HITS-CLIP to facilitate direct ligation of miRNA and target RNA. Endogenous AGO–RNA complexes were purified from ultraviolet-irradiated mouse brain neocortex using monoclonal anti-AGO and were washed in stringent conditions that disrupt native AGO–mRNA interactions. Complexes were treated with dilute RNAseA to generate footprint-sized fragments. To test whether T4 RNA ligase I treatment could join free RNA ends, AGO–RNA was radiolabelled with polynucleotide kinase (PNK) and 32P-γ-ATP, then treated with RNA ligase. Complexes were treated with alkaline phosphatase and visualized by SDS–polyacrylamide gel electrophoresis (PAGE) and autoradiography to assess dephosphorylation. Compared with untreated samples, ligase-treated complexes were ‘protected’ from dephosphorylation, indicating ligation of RNA ends.

Figure 1 | CLEAR-CLIP unambiguously identifies endogenous in vivo miRNA–target interactions. (a) In CLEAR-CLIP, AGO–target contacts are cross-linked in vivo by ultraviolet irradiation. Endogenous AGO is immunopurified from tissue lysates and washed under stringent conditions that disrupt the interaction of AGO–miRNA with non-cross-linked target RNAs. Target regions cannot be cloned from no-ultraviolet controls, indicating that cross-linking of AGO to target mRNA (shown as ‘X’) is required. Cross-linking of the microRNA may not be necessary, because the AGO–miRNA interaction is uniquely strong and survives stringent washing. After washing, RNA ends are modified to facilitate miRNA–target ligation and joined with T4 RNA Ligase I treatment, yielding miRNA–target chimeric RNAs in two orientations at the indicated frequencies. All depicted post-IP manipulations up to SDS–PAGE occur on beads. Correlation plots of miRNA abundance of all miR-first (b) and miR-last (c) chimeras versus small RNA sequencing data in the brain. Pearson’s correlation coefficients (r) are shown. CDF plots of cognate miRNA seed matches in target regions relative to ligation site for all miR-first chimeras in plus-ligase (d) and no-ligase (e) samples, and for all miR-last chimeras in plus-ligase (f) and no-ligase (g) samples. (h) Distribution of standard AGO CLIP and miRNA-target chimeras in transcript regions. (i) CLEAR-CLIP confirmed known miRNA regulation, here exemplified by miR-124 regulation of the Ptxp1 3′-UTR. Other examples are shown in Supplementary Fig. 3c.
not be cloned from no-ultraviolet controls, indicating that in vivo AGO–mRNA ultraviolet cross-linking was strictly required for CLEAR-CLIP.

miRNA frequency in miR-first chimeras correlated with brain miRNA abundance (Fig. 1b and Supplementary Fig. 1b–d). miR-first chimeras were dominated by a small number brain-abundant miRNAs (Supplementary Fig. 1e). In contrast, miR-last chimeras did not correlate to miRNA abundance and were dominated by dubiously annotated miRNAs (Fig. 1c and Supplementary Fig. 1d). Target regions in miR-first chimeras

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**Diagram Description:**

- **Panel a:** UV irradiation of live brain tissue leading to tissue lysis and DNAse treatment.
- **Panel b:** Canonical seed matches in miR-first, plus-ligase.
- **Panel c:** Canonical seed matches in miR-last, no-ligase.
- **Panel d:** Canonical seed matches in miR-first, plus-ligase.
- **Panel e:** Canonical seed matches in miR-last, no-ligase.
- **Panel f:** Canonical seed matches in miR-first, plus-ligase.
- **Panel g:** Canonical seed matches in miR-last, no-ligase.
- **Panel h:** Canonical seed matches in miR-first, plus-ligase.
- **Panel i:** Canonical seed matches in miR-last, no-ligase.

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**Key Points:**

- miRNA frequency in miR-first chimeras correlated with brain miRNA abundance.
- miR-first chimeras were dominated by a small number brain-abundant miRNAs.
- miR-last chimeras did not correlate to miRNA abundance.

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**Figure Captions:**

- **Figure 1b:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1c:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1d:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1e:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1f:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1g:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1h:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
- **Figure 1i:** Chimeras abundance log₂ (counts) vs. miRNA abundance log₂ (miRNA-seq).
were also strongly enriched for canonical seed matches to their cognate miRNAs (Fig. 1d). Seed enrichment occurred within ~75 nt of the miRNA ligation junction in the expected downstream (3′) region, but not the upstream region (5′) (Fig. 1d). Consistent with prior findings, chimeras were present at low levels in no-ligase samples26, although with reduced seed enrichments (Fig. 1e). For miR-last chimeras, the reversed pattern of seed distribution around the ligation junction was expected; however, this pattern was weak in ligase-treated samples and was absent in no-ligase samples (Fig. 1f,g). As they better reflected miRNA abundance and known miRNA targeting features, we focused exclusively on miR-first chimeras (henceforth ‘chimeras’).

Notably, many CLEAR-CLIP target regions lacked canonical seed matches (Fig. 1d), consistent with similar analyses19,26. We took two approaches to assess miRNA ligation to non-cross-linked targets, which could falsely identify non-physiologic interactions. First, we tested chimera ligation after denaturing AGO complexes in 6M guanidine hydrochloride, as in CLASH19. Interactions from denatured samples were similar to other samples based on miRNA seed match frequency, indicating bona fide interactions. However, compared with other samples, the yield of chimeric and non-chimeric CLIP reads was low (Supplementary Table 1) and skewed to non-genic sites (Supplementary Fig. 1f); thus, we pursued it no further.

Second, we performed mixing experiments to assess miRNA ligation to non-target sequences after postlysis re-association. CLEAR-CLIP was done on lysates from cross-linked mouse cortex mixed with *Escherichia coli* total RNA, which contains thousands of potential miRNA sites by random chance at a per-nucleotide frequency comparable to mouse. For two replicates each, equal mass amounts of mouse and *E. coli* RNA or a large excess of *E. coli* RNA (sixfold) were mixed. We confirmed that *E. coli* RNA was not degraded in brain lysates (Supplementary Fig. 2). Across four mouse-only control samples, 1% of chimeric CLIP reads mapped to the *E. coli* genome, establishing the ‘background’ from cross-mapped reads and minute RNA contaminants from commercial enzymes29 (Supplementary Table 2). Average *E. coli* mapping rates were 1.9% in equal-mixture samples and 5.2% in excess-mixture samples. To examine a more complex competitor RNA pool, we performed CLEAR-CLIP on mixed lysates from ultraviolet-irradiated mouse brain and non-cross-linked *Drosophila* S2 cells containing equal amounts of RNA. Here, 0.7% of mouse-only chimeric sequences mapped to the *Drosophila* genome compared with 2.9% of mixed mouse/*E. coli* samples (Supplementary Table 2). Collectively, these experiments indicate low (<5%) false discovery comparable to related methods19.

**CLEAR-CLIP enhances the brain miRNA regulatory map.** Chimeras with the same miRNA and overlapping genomic coordinates were clustered to yield 130,120 brain miRNA–target interactions (Fig. 1a and Supplementary Data 1). Seventy-nine per cent (102,882) of interactions were also supported by non-chimeric AGO CLIP reads. We combined chimeric CLEAR-CLIP reads with conventional CLIP reads from 15 total biological replicates, to generate an enhanced brain miRNA regulatory map. We identified 96,685 AGO peaks supported in at least 5 mice, defined as biological complexity (BC) ≥5 (Supplementary Data 2)22. Twenty-seven per cent of BC ≥5 peaks (26,304) had chimera support unambiguously identifying the miRNA(s) and this proportion increased substantially for peaks with greater BC (Supplementary Fig. 1g). Consistent with our prior studies, ~20% of brain AGO peaks were ‘orphans’ lacking 6mer seed matches for the 35 most abundant miRNA families22. Chimera data linked miRNAs to 6,136 (~28%) orphan peaks, disambiguating thousands of biologically robust non-canonical miRNA-binding sites.

Chimera-defined interactions and non-chimeric AGO CLIP reads were similarly distributed in the transcriptome (Fig. 1h). In addition to 3′-untranslated region (UTR) and coding DNA sequence (CDS) sites, chimeras identified many intronic sites with miRNA-dependent AGO binding (Supplementary Fig. 3a)30–32. Intronic interactions were not previously reported for CLASH in 293T cells, because reads were only aligned against mature transcripts19. Our alignment of raw CLASH data against a genomic reference recovered many intronic (~15%) and other non-3′-UTR sites (~60%), independently confirming such binding. To examine whether annotated intronic interactions in the brain fall in mis-annotated exons, we examined polyA+ RNA sequencing from age-matched mouse cortex33. As polyA selection strongly enriches mature transcripts, introns show much lower coverage than coding or 3′-UTR exons. Accordingly, chimera-identified intronic sites showed low RNA sequencing coverage relative to exonic sites (Supplementary Fig. 3b). For comparison, binding sites for NOVA and RBFOX in the brain, which also bind intronic and exonic sequences, showed similar patterns34,35.

CLEAR-CLIP retrieved known miRNA regulatory sites (Fig. 1i and Supplementary Fig. 3c) and functions for well-characterized neuronal miRNAs, such as miR-124 and miR-9, in neuron development, synapse formation and axon guidance (Supplementary Data 3)22,36,37. Gene Ontology analysis indicated neuronal regulatory functions for less-characterized brain miRNAs, including miR-26 (for example, axon development and locomotion), miR-138 (neurotransmitter transport and secretion, and calcium transport) and miR-9* (cell migration and motility; Supplementary Data 3). In addition, Kyoto Encyclopedia of Genes and Genomes (KEGG) database analysis recovered known associations of miR-124, miR-9 and miR-26 with glioma, including known and many novel targets (Supplementary Fig. 4).

**CLEAR-CLIP-identified sites are functional.** Chimera-identified sites from the brain are functional in global analyses of miRNA perturbation. For brain polyribosome-associated miRNAs from miR-128 knockout (KO) and wild-type (WT) mice, the presence of miR-128 chimeras in transcript 3′-UTRs correlated with enhanced polyosome association in miR-128 KO brain (Fig. 2a)2. Sites with canonical seed matches and non-canonical sites predicted significant de-repression (Fig. 2b).

More detailed analysis was possible for miR-124 due to the large number of identified sites. In CAD neuroblastoma cells transfected with miR-124 mimic, the presence of miR-124 chimeras in 3′-UTRs in mouse brain correlated with repressed transcript levels compared with control cells (Fig. 2c)38. Chimera sites identified once (cluster size, N = 1) predicted significant regulation and sites identified multiple times (N > 1) or overlapping AGO CLIP peaks conferred stronger repression.

Consistent with our prior studies, AGO peaks encompassing miR-124 seed matches predicted significant transcript repression in miR-124-transfected cells (Fig. 2d)22. Critically, when such peaks overlapped miR-124 chimeras, repression was significantly greater. Thus, chimera information improved identification of functional miRNA sites in vivo. To examine different types of miR-124 sites, we defined mutually exclusive sets of transcripts possessing only chimera-defined canonical miR-124 sites or only non-canonical sites. Canonical sites correlated with significant transcript repression (Fig. 2e). Non-canonical sites predicted only a small shift in RNA levels (Fig. 2f) due largely to bulged 8mer miR-124 sites, the only non-canonical group predicting
significant transcript repression in this data set. These analyses show that AGO HITS-CLIP maps supplemented with chimera data improved identification of functional miRNA target sites, including specific non-canonical sites.

Diverse miRNA–mRNA pairing patterns. In addition to canonical sites, motif searches allowing expanded seed match variants revealed a high proportion of single mismatch and bulged sites (>30% together), and many (~20%) lacking appreciable seed homology (Fig. 3a). These patterns were similar across different transcript regions, showing that CDS and intronic AGO targeting follows similar rules to 3′-UTR binding. For chimera clusters of increasing sizes (N) and chimeras overlapping AGO peaks, canonical sites were slightly enriched (Fig. 3b). Similar canonical motifs were used by all miRNAs but relative frequencies varied (Fig. 3c).

We determined overlap of chimera-defined sites with TargetScan predictions, a purely bioinformatic approach, for six abundant brain miRNA families’. Chimera-identified sites in 3′-UTRs for a given miRNA were much more likely to overlap TargetScan-predicted sites for that miRNA than random control sites (Fig. 3d). Nonetheless, TargetScan supported only a minority of chimera-defined sites and concordance varied for different miRNAs. A major source of discrepancy was the preponderance of 6mer and imperfect seed match variants in chimera-identified binding sites, functional categories not present in TargetScan. Detailed analysis of imperfect seed sites confirmed established patterns, such as the miR-124 target G bulge between miRNA positions 5 and 6 (Fig. 3e and Supplementary Fig. 5a,b). Notably, 22 of the top 25 brain miRNAs disallowed bulging at one or more sites, most often position 5 (16/25). These preferences identify specific single-nucleotide target deletions that, presumably by forcing unfavourable miRNA bulges, should effectively abolish AGO binding and regulation. Compared with bulged motifs, seed mismatches were more evenly distributed and showed less miRNA-specific variation (Fig. 3e and Supplementary Fig. 5c). An exception was G−U wobble interactions, which showed strong preferences such as miR-30 position 3 (Supplementary Fig. 3d).

Unbiased de novo motif analysis of chimera target regions identified strong enrichment of seed-complementary motifs (Fig. 3f)39. miRNAs without significant seed binding were mostly low-abundance, often passenger-strand isoforms, which could be affected by sampling error. In addition, many miRNA targets had

Figure 2 | miRNA–target chimeras identify functional interactions. (a) Polyribosoma association in miR-128 KO versus WT mouse brain plotted as a CDF for 3′-UTR sites identified with miR-128 chimeras (red) and non-miR-128 3′-UTR chimeras (black). (b) CDF as in a shown for canonical (blue) and non-canonical (orange) miR-128 sites. (c) Fold change in mRNA levels in CAD cells transfected with miR-124 mimics versus control38 are plotted as a CDF for 3′-UTR sites identified with miR-124 chimeras in the brain. miR-124 sites identified once (red), multiple times (blue) or overlapping AGO CLIP peaks (magenta) are shown compared with non-miR-124 sites (black). (d) CDF of 3′-UTR miR-124 sites as in c, showing miR-124 sites identified with chimeras (violet), peaks overlapping miR-124 seed matches (cyan) or peaks overlapping both seeds and miR-124 chimeras (orange). (e) CDF plots for transcripts with only chimera-defined canonical 3′-UTR miR-124 sites, broken down by site type. (f) CDF as in e for all 3′-UTR non-canonical sites (green) and bulged 8mer sites (cyan). All panels, P-values from Kolmogorov-Smirnov testing comparing coloured subsets with control (black) sites are shown, along with the number of sites (n) in each set.
strong enrichments for motifs complementary to miRNA 3′-end sequences. Several auxiliary motifs included the classic complimentary pairing region from nucleotides 13–16, but many different regions of auxiliary binding were evident.17

Expanded miRNA–target pairing rules in the brain. Motif analysis revealed extensive seed-based and auxiliary miRNA targeting in vivo. For resolution of individual events, we performed duplex structure predictions for target regions and their

Figure 3 | Motif analysis reveals miRNA binding dependent on seed and auxiliary pairing. The proportion of chimera-defined target regions with the indicated seed variants is plotted, broken down (a) by transcript region, (b) by the number of times interactions were identified with chimeras (N) or whether chimeras overlapped AGO CLIP peaks and (c) for the most abundant miRNA families in mouse brain, ranked from the top by decreasing abundance. (d) Overlap of 3′-UTR chimera-identified sites in the brain with TargetScan predicted sites for the same miRNA (red) or three equally sized random control sets of TargetScan sites (black). Control sets were restricted to the top 20 brain miRNAs. Only target sites in miRNAs with detectable expression in the cortex were considered. (e) The distributions of mismatched and bulged nucleotides for chimera-identified sites with imperfect seed motifs are plotted for the top 25 mouse brain miRNAs (black), miR-124 (red) and miR-9 (blue). Error bars show the s.d. at each position for the top 25 miRNAs in the brain. miRNA seed sequences for miR-124 and miR-9 are shown below mismatch and miRNA bulge plots. Below the target bulge plot, the most frequently bulged target nucleotide at the indicated position is shown when strong preferences (>50% of sites) were apparent. Sites from all transcript regions were included in this analysis. (f) De novo analysis of cognate miRNA-complementary-enriched 7mer motifs in all chimera target regions plotted as a heat map across the miRNA. Each line represents one miRNA and colour intensity scales with abundance in target sequences. miRNAs are ordered by hierarchical clustering.
k-means clustering of structures revealed six major modes of miRNA–target binding, with five dominated by seed-site pairing combined with various auxiliary binding patterns (Fig. 4a,b). Four clusters (k = 1–4) closely mirrored similar analyses of 293T CLASH sites, including a seed-independent class (k = 4)\(^1\). A fifth group identified by CLASH, encompassing ~20% of interactions and lacking significant miRNA–target pairing, was not identified here. We also observed novel classes with seed pairing coupled with bipartite or tripartite auxiliary pairing patterns. These cognate miRNAs using RNAhybrid (Supplementary Data 1)\(^4\).
clusters, including the distinctive patterns of auxiliary binding, were not observed when target regions and miRNAs were shuffled by randomly re-assigning each chimeric target region the miRNA from a different chimera. Shuffled interactions showed significantly lower duplex hybridization energies than true ones, consistent with the discovery of real binding events (Fig. 4c).

Remarkably, of 212 miRNAs with > 50 identified target sites in the brain, 196 (~90%) showed significant enrichment or depletion in one or more k-mers binding class (Fig. 4d and Supplementary Table 3). For example, miR-124 was strongly enriched in groups 1 \((P = 1.6 \times 10^{-245})\), Fisher’s exact test) and 5 \((P = 1.0 \times 10^{-174})\), and marginally in group 2 \((P = 2.1 \times 10^{-3})\). In contrast, miR-124 was strongly depleted in groups 3 \((P < 1.6 \times 10^{-245})\), 4 \((P = 3.7 \times 10^{-140})\) and 6 \((P = 1.1 \times 10^{-174})\). This pattern confirmed strong seed dependence for miR-124 binding and revealed distinct patterns of favoured auxiliary binding (Fig. 4b,d). Motif analysis also supported auxiliary binding, showing an enriched 7mer motif complementary to miR-124 positions 14 to 20 (Fig. 3). Structural inference revealed distinct binding patterns contributing to this motif conservation.

Some miRNAs tolerated striking diversity in pairing interactions. miR-9 was enriched in group 3 \((P = 3.3 \times 10^{-138})\), Fisher’s exact test), characterized by strong seed dependence and frequent auxiliary pairing from positions 14 to 22, and group 6 \((P = 3.9 \times 10^{-117})\), characterized by a tripartite auxiliary pattern (Fig. 4b). miR-9 was also enriched for seedless binding \((k = 4, P = 2.2 \times 10^{-9})\). Similarly, miR-181 family members were enriched in both seed-dependent and -independent classes. Globally, interactions with more predicted seed pairing exhibited fewer predicted auxiliary base pairs and vice versa (Fig. 4e). Canonical sites with less seed pairing (6mer and 7mer-A1) had slightly more predicted auxiliary pairing than stronger seed sites (8mer and 7mer-m8), consistent with supplementary 3’-pairing (Supplementary Fig. 6a)\(^17\). A stronger effect was evident for bulged or mismatched 8mer and 7mer motifs, which had more auxiliary pairing than their perfect match counterparts, indicating complementary pairing to offset imperfect seed matches (Supplementary Fig. 6b-d)\(^18\).

Specific classes of CLEAR-CLIP-defined sites are preferentially conserved in mammals, consistent with functional significance\(^4,42\). In both CDS and 3’-UTRs, groups 1, 2 and 3 were modestly more conserved than groups 4, 5 and 6, with seedless interactions \((k = 4)\) showing lowest overall conservation (Supplementary Fig. 7a,b). The 3’-UTR sites with canonical seed matches and certain bulged or mismatched motifs were more conserved than sites lacking seed homology (Supplementary Fig. 7c). CDS sites showed a similar pattern, except for mismatched sites (Supplementary Fig. 7d). To compare conservation of seed and auxiliary pairing regions, we calculated conservation scores in the seed and auxiliary portions of 3’-UTR target sites. For 8mer and 7mer-m8 sites, target seed regions were modestly more conserved than the auxiliary region \((P < 0.05,\) one-tailed \(t\)-test). For other sites, seed and auxiliary regions were similarly conserved (Supplementary Fig. 7e), implying evolutionary pressure to maintain the whole miRNA binding site.

We confirmed chimera-identified regulation by transfecting miRNA mimics into mouse neuroblastoma (N2A) cells and measuring endogenous target mRNA levels by quantitative reverse transcriptase–PCR (qRT–PCR). miRNA mimics repressed most miR-9 (6/7) and miR-181a (5/6) targets examined, including all with canonical seeds and several with seedless interactions and no canonical seed matches in their 3’-UTRs (Fig. 4f). These experiments support prior findings that seed-independent miRNA targeting is functional but weaker than seed-dependent regulation\(^14,19\).

Endogenous miRNA–target chimeras in human hepatoma cells.

To independently assess miRNA–target pairing patterns, we searched for miRNA–target chimeras in standard HITS-CLIP libraries from human hepatoma (Huh-7.5) cells. miR-first chimeras were present at ~0.5% of unique reads, suggesting that on-bead RNA ligase I treatment for 3’-linker addition in the standard protocol can form chimeras (Supplementary Table 4 and Supplementary Data 4). As in the brain, miR-first chimera target regions were strongly enriched for cognate miRNA seed matches, whereas miR-last were less so (Fig. 5a,b). In total, 34,986 miRNA–target interactions were identified in Huh-7.5 cells (Supplementary Fig. 8a)\(^43\), confirming that standard HITS-CLIP libraries contain miRNA–target chimeras, albeit at reduced frequency\(^26\).

To further test the functionality of chimera-identified sites, we examined data from Huh-7.5 cells treated with locked nucleic acid (LNA) against miR-122 or miraviren, a clinical miR-122 inhibitor\(^44\). AGO binding to 3’-UTR regions with miR-122 7mer or 8mer seed matches was specifically reduced in miR-122 LNA versus control cells (Fig. 5c). This effect was stronger for sites overlapping miR-122 chimeras and even stronger when both predictors were combined. When regions outside 3’-UTRs were included, a significant effect was only observed when miR-122 chimeras were present (Fig. 5d). These results indicate that chimeras enhanced prediction of 3’-UTR and non-3’-UTR sites. For miraviren treatment, miR-122 seed presence alone was predictive in all cases, but miR-122 chimeras enhanced these predictions (Supplementary Fig. 8b,c). This analysis provided further evidence that miRNA chimeras improve identification of miRNA regulatory sites.

miRNA–target chimeras in the absence of exogenous ligase.

Chimeras independent of exogenous ligase were present in small numbers in mouse brain and were reported in \(C.\) \(e\)\(g\)\(a\)\(s\)\(^26\). These interactions showed significant seed enrichment, suggesting many are real (Fig. 1e). We used CLEAR-CLIP in Huh-7.5 cells to investigate mammalian transfer RNA ligase HSPC117 as a potential source of these chimeras and a means to enhance chimeria ligation\(^45\). As in mouse brain, Huh-7.5 CLEAR-CLIP yielded chimeras at ~2% of mapped reads. Ligase-treated samples showed a ~10-fold enrichment for miR-first chimeras and a smaller enrichment for miR-last (Fig. 5e). CLEAR-CLIP without ligase addition was also done on Huh-7.5 cells with induced overexpression of HSPC117 or efficient depletion by RNA interference (Supplementary Fig. 8d). In both conditions, chimeria frequencies were not significantly different from controls with endogenous HSPC117 levels (Fig. 5e). We also searched for chimeras containing truncated miRNAs, in case RNAs cleavage was a prerequisite for HSPC117-mediated ligation\(^26\), yielding the same result (Fig. 5f). Interestingly, truncated chimeras in Huh-7.5 cells comprised an additional ~1% of mapped reads, far more than in the brain, with most truncated one nucleotide (Supplementary Fig. 8e). This analysis ruled out HSPC117 as a major endogenous source of chimeras.

Expanded miRNA–target pairing rules in human cells.

Motif and structural analysis revealed global miRNA–target pairing patterns in Huh-7.5 cells. As in mouse brain, seed-complementary motifs were identified for most miRNAs, in addition to many 3’-auxiliary motifs (Fig. 6a). For structure clustering, informative binding classes in Huh-7.5 cells were most evident with seven \(k\)-groups, as opposed to six in mouse brain (Fig. 6b). Two Huh-7.5 groups (5A and 5B), similar to group 5 from mouse brain, showed bipartite auxiliary pairing but at distinct sites. The other clusters closely resembled corresponding groups in mouse
**Figure 5 | miRNA-target chimeras identify functional interactions in Huh-7.5 cells.** CDF seed-enrichment plots as in Fig. 1d–g for miR-first (a) and miR-last (b) chimera target regions from Huh-7.5 HITS-CLIP. CDF plots of LNA-122 induced changes in AGO binding across 3'-UTRs (c) or all regions (d) for sites with miR-122 7-8mer seeds (magenta), miR-122 chimeras (red) or the combination of both (blue). P-values are shown for Kolmogorov-Smirnov tests comparing indicated subsets to control (black) sets. (e) miR-first and miR-last chimeras in CLEAR-CLIP on Huh-7.5 cells as a percentage of total unique reads, varying the presence of exogenous T4 RNA ligase and RNAse. For HSPC117 ligase, (+) represents endogenous levels, (−) represents siRNA knockdown and (±) represents overexpression as shown in Supplementary Fig. 8d. (f) Analysis of CLEAR-CLIP derived miR-first chimera truncation and the effects of HSPC117 manipulation. Percentage of chimeras harbouring full-length miRNAs was compared with chimeras with the indicated 3'-truncations, or with all putative chimeric reads with at least 12 nts miRNA sequence starting at the 5'-miRNA end. In e,f, the mean values of two biological replicates is shown for each sample, with error bars indicating s.d.

**Auxiliary pairing regulates miRNA-target specificity in vivo.** As a striking indication that auxiliary pairing regulates miRNA-target specificity, duplex structure analysis revealed distinct binding patterns for members of miRNA seed families (for example, let-7, miR-30, miR-181 and miR-125) (Fig. 4d). As CLEAR-CLIP does not yet provide comprehensive coverage of all miRNA-binding sites, it was not possible to compare the overlap of different miRNA paralogues by occupancy analysis. Instead, we used *de novo* motif analysis to search for distinguishing features of the target regions of individual paralogues. For most miRNA family members, motifs complementary to divergent 3'-sequences were highly enriched in cognate target regions but not their paralogues (Fig. 7a,b, below charts). Next, we reasoned that if inter-family preferences existed, family members should form more stable duplex structures with their own identified target regions than other paralogues. We calculated duplex energies for CLEAR-CLIP target regions of each abundant let-7 family member in the brain with each let-7 miRNA in a four-way pair-
wise comparison (Fig. 7c). In all cases, let-7 family miRNAs formed more stable structures with their cognate target regions than other paralogues. This observation is striking in that some paralogues (for example, let-7b and let-7c) have higher GC content and thus intrinsic potential for more stable structures. Shuffling analysis of miR-30 family members revealed similar specificity, although certain preferences were more significant than others (Fig. 7d). Specifically, miR-30b and miR-30c showed more significant differences from miR-30a, miR-30d and miR-30e than from each other and vice versa. Analysis of miR-125 and miR-181 families revealed additional intra-family target preferences (Supplementary Fig. 9a–d). Thus, motif and structure information indicate distinct targeting preferences for miRNA paralogues controlled by differential miRNA 3′-end pairing.

We validated functional specificity of miRNA family members using fluorescence reporters with paralogue-specific target sites in their 3′-UTRs (Fig. 8a)46. We examined miR-30a, miR-30c and miR-125a targets sites predicted to form more stable pairing with a specific paralogue and which were ligated to only that paralogue in at least two CLEAR-CLIP experiments. Reporters were co-transfected into N2A cells with plasmids expressing miRNA family members or a control C. elegans miRNA. miRNA expression was confirmed by northern blotting (Supplementary Fig. 10a) and silencing activity was confirmed using reporters with perfect complementary sites (Supplementary Fig. 10b,c). For
**Figure 7** | CLEAR-CLIP reveals target specificity among miRNA family members. Base pairing profiles from duplex structure maps for let-7 (a) and miR-30 (b) family members are shown. For each miRNA, the fraction of interactions with base pairing at each miRNA position is plotted. miRNA sequences are shown below with coloured bases indicating divergent nucleotides. De novo motif analysis of target regions for indicated miRNAs revealed family-member-specific motifs complementary to divergent parts of the miRNAs. For easier interpretation, the target motifs were reverse complemented to match the miRNA sequences. P-values for enrichment over background (AGO-binding regions in brain) from HOMER are indicated. No unique auxiliary motif was found for let-7a, the only such case. (** and the indicated let-7 (c) or miR-30 (d) family members is shown. Targets paired with their chimera-identified, cognate let-7 family member are shaded darker. Interactions from all transcript regions were included in these analyses. Box plots depict interquartile (25–75) values (* and **** for one paralogue was significantly more stable (Fig. 8l,m) compared to that of a perfect 8mer site (Fig. 8b), consistent with our finding that G–U pairing is well-tolerated at specific seed positions (Supplementary Fig. 5d). Conversely, more subtle differences in predicted pairing (2.8 kcal mol$^{-1}$) enhanced miR-30c activity at a 6mer site with predicted supplementary 3′-pairing (Fig. 8f). This complexity underscores the need for empirical binding maps to supplement structure- and sequence-based predictions. More broadly, these results illustrate paralogue-specific miRNA activity and diverse functional classes of non-canonical sites.

**Discussion**

CLEAR-CLIP gains its power from the formation of sequential covalent bonds that reflect *in vivo* interactions. The utility of...
AGO-binding sites were cloned into the 3′-UTR of the tagRFP cassette, with tagBFP used for internal normalization. Plasmids co-expressing miRNAs and GFP were co-transfected and measurements were taken 48 h later. In mixing experiments, CLEAR-CLIP thus provides a snapshot of true, physiologic miRNA–target interactions and is uniquely applicable to all mammalian model systems and human samples. In contrast to approaches without relying on specialized tagging strategies, CLEAR-CLIP thus provides a snapshot of true, physiologic miRNA–target interactions and is uniquely applicable to all mammalian model systems and human samples.

Figure 8 | miRNA family member specificity confirmed by single cell measurements. (a) A system for single cell measurements of miRNA-mediated repression, adapted from ref. 46. tagRFP and tagBFP are expressed from the doxycycline-inducible bidirectional pTRE-3G-BI promoter. CLEAR-CLIP-defined AGO-binding sites were cloned into the 3′-UTR of the tagRFP cassette, with tagBFP used for internal normalization. Plasmids co-expressing miRNAs and GFP were co-transfected and measurements were taken 48 h later. (b–m) Log-transformed plots of tagRFP versus tagBFP fluorescence, with minimum free energies (MFEs) (ΔG) for predicted base pairing between duplex structures for indicated paralogues. A description of the site type is shown above each plot, with bold labelling denoting successful validation of paralogue specificity. Evaluation of miR-30a (red), miR-30c (blue) and negative control miRNA (black) overexpression on (b–i) sites with predicted miR-30a preference; and (j–m) sites with predicted miR-30c preference. Representative plots from at least two independent experiments for each construct are shown.
CLASH, CLEAR-CLIP does not require fully denaturing AGO and involves a single purification step. Our experiments with denatured AGO and analyses of published CLASH data showed low yield of standard non-chimeric CLIP reads compared with standard AGO HITS-CLIP, hindering robust AGO-binding peak identification. With straightforward modifications of HITS-CLIP, CLEAR-CLIP simultaneously generates chimera information and high-quality, transcriptome-wide AGO HITS-CLIP maps. These dual data sets improved identification of functional miRNA target sites compared with HITS-CLIP or chimeras alone (Figs 2d and 5c,d), a key advantage, as miRNA–target ligation remains limiting. Optimized ligation conditions yielded at least tenfold enrichment in ligase-treated versus non-ligase samples, a substantial improvement over prior methods16, but insufficient for comprehensive coverage. A key future goal is further improvement of this efficiency to reduce false negatives and achieve the global coverage of HITS-CLIP maps.

CLEAR-CLIP yielded insights into pairing rules for over 200 mammalian miRNAs. Enriched target motifs revealed seed-dependence in most miRNAs, with widespread bulged or mismatched pairing, and extensive 3′-auxiliary interactions (Figs 3 and 6). miRNA–target duplex structure prediction clarified that most interactions employed seed and auxiliary pairing in combination (Figs 4 and 6). Most miRNAs were significantly enriched or depleted in one or more binding class, with many favouring two or more categories. This tolerance for significant enrichment or depletion in one or more binding class would be expected to promote diverse pairing rules for other distinct but constrained pairing structures was most apparent for many favouring two or more categories. This tolerance for mismatched pairing, and extensive 3′-auxiliary interactions (Figs 3 and 6). miRNA–target duplex structure prediction clarified that most interactions employed seed and auxiliary pairing in combination (Figs 4 and 6).

Motifs and structure inference showed extensive pairing of miRNA 3′-ends with targets. Such auxiliary interactions can stabilize or enhance miRNA–target pairing, in particular together with imperfect seed pairing18. Global analysis of bulged and mismatched seed interactions from CLEAR-CLIP shows this phenomenon is common (Supplementary Figs 5 and 6). The importance of 3′-auxiliary binding is still debated, with some reports demonstrating significant effects18,49 and others concluding limited ones5. Analyses of miRNA mimic transfections found that supplementary pairing of miRNA bases 12–17 marginally enhanced target repression in rare instances17,50. However, the sensitivity of such analyses may be limited by stringent requirements for continuous spans of auxiliary binding7. CLEAR-CLIP revealed diverse, often discontinuous auxiliary pairing that could hinder the detection of motif presence or conservation above background (Figs 4a and 6b). A second consideration is the heavy reliance of prior conclusions on acute overexpression of miRNAs, which may perturb endogenous AGO–miRNA–target stoichiometry or interrogate different target repertoires than are available in vivo. Recent evidence for co-evolution of miRNAs and targets, in particular in neurons, underscores the importance of examining physiologic interactions31. The use of transcript destabilization in vitro as a sole functional readout may also overlook other AGO functions, including translational control, targeting to non-3′-UTR regions and interactions with other RNA-binding proteins42.

As a striking indication that auxiliary interactions regulate miRNA target specificity, we observed specificity among paralogues in miRNA seed families (Fig. 7). Such specificity was previously illustrated for two let-7 family targets in Drosophila and has been speculated elsewhere18. Functional single-cell assays confirmed paralogue specificity for several sites from brain CLEAR-CLIP (Fig. 8). Other sites were similarly regulated by different paralogues, indicating miRNA family members are functionality redundant at certain sites and specific at others. Indeed, the strict conservation of miRNA families and their unique expression patterns in vivo, including across brain regions, supports specific functions52,53.

The predominance of canonical seed pairing in mediating miRNA target level repression is supported by CLEAR-CLIP-defined sites (Fig. 2). In addition, CLEAR-CLIP data demonstrated widespread, functional non-canonical miRNA targeting and substantial diversity in canonical and non-canonical interactions among different miRNAs. CLEAR-CLIP identified functional, non-canonical regulation globally for miR-128 and miR-124 (Fig. 2), and for individual miR-9, miR-181, miR-30 and miR-125 targets (Fig. 4f and Fig. 8b–m). Non-canonical sites included diverse seed mismatch and bulged variants, and seedless interactions in both mouse brain and HuH-7.5 cells. Interestingly, a number of miRNAs enriched for seedless interactions (for example, miR-9, miR-181, miR-30 and miR-186) have AU-rich seed sites, indicating that weak seed-pairing stability may favour seedless non-canonical interactions. Our results support growing evidence of widespread non-canonical miRNA regulation that is likely to have a large collective impact13–15,17,19,20,21. We expect CLEAR-CLIP and similar methods will facilitate discovery on these sites and refine in vivo miRNA regulatory maps in future studies.

Methods

Mice. All mouse experiments were approved by The Rockefeller University Institutional Animal Care and Use Committee regulations. P13-aged C57BL6/J mice were used for all experiments, except for BR21, BR22 and BR23 (Drosophila mixing), which used 6-week-old mice.

CLEAR-CLIP. Tissue cross-linking and lysis. Neocortex was dissected and cross-linked as described and snap frozen14. Frozen pellets were re-suspended in threefold volume (w/v) lysis buffer (1 × PBS/1% Igepal/0.5% sodium deoxycholate/0.1% SDS) containing Complete protease inhibitors (Roche). Lysates were treated with 30 U RNase A (Promega) at 37 °C for 5 min with shaking.

Pre-immunoprecipitation RNase treatment. For samples BR1, BR2, BR4, BR13, BR14, BR15, BR16, BR17, BR18, BR19, BR20, BR21, BR22 and BR23, RNase A (USB Products) was added to lysates at 0.0001 U ml⁻¹ and incubated at 37 °C for 5 min. RNAsin Plus (Promega) was added at 0.5 U ml⁻¹ and lysates were cleared by ultracentrifugation (50 000g). For remaining samples, RNase treatment was done after immunoprecipitation (see below).

Immunoprecipitation and washing. Cleared lysates were rocked with Dynal Protein A beads (Life Technologies) prepared with 2A8 anti-AGO27,55 for 90 min at 4 °C, then washed:

- Three times lysis buffer containing 5 × Denhardt’s solution
- Twice high-detergent buffer (1 × PBS/1% Igepal/1% sodium deoxycholate/0.2% SDS)
- Three times low-salt buffer (15 mM Tris pH 7.5, 5 mM EDTA)
- Twice high-salt buffer (1 × PBS/1% Igepal/0.5% sodium deoxycholate/1% SDS, 1 M NaCl (final, including PBS))
- Twice PKN wash buffer (50 mM Tris pH 7.5, 10 mM MgCl2, 0.5% Igepal)

On-bead RNase treatment. For samples BR1, BR5, BR6, BR7, BR8, BR9, BR10, BR11 and BR12, beads were re-suspended in 0.5 ml lysis buffer containing 2 mg ml⁻¹ BSA and RNase A at 0.0002 U ml⁻¹. Samples were treated at 37 °C for 5 min with shaking, transferred to ice and supplemented with 0.5 U ml⁻¹ RNasin Plus. Beads were rocked for 20 min at 4 °C, to recover any dissociated antigen, then washed:

- Three times high-detergent buffer
- Three times low-salt buffer
- Once high-salt buffer
- Twice PKN buffer

5′-End phosphorylation and chimera ligation. Beads were treated with PKN (3′-phosphatase minus) (NEB) and 1 mM ATP to phosphorylate cleaved miRNA 5′-ends. Beads were washed three times in PKN buffer, then chimera ligation was
performed overnight at 16°C with 0.625 µM -1 T4 RNA Ligase I, 1 mM ATP and 0.1 mg mL -1 RSA in a 100 µl total volume. The following morning, fresh RNA Ligase I (25 U) and ATP (1 mM) were added to each sample and incubation was continued 4-6h. For minus-ligase controls (BR and BRS), RNA ligase was omitted. Beads were washed:

- Twice lysis buffer
- Once PKEDTA/E GTA buffer (50 mM Tris pH 7.5, 10 EDTA,10 mM E GTA, 0.5% Igepal)
- Twice PKEDNA phosphate treatment and 3’- linker ligation. Alkaline phosphate treatment was performed to remove 3’- phosphate groups 27. Pre-adenylated 3’- linker (5’-R-AppGTGCTAGCTACCTCAGCGG-3’) was added using truncated RNA Ligase 2 (NEB), with 2.5 µM 20 µM linker and 1 U enzyme per 40 µl reaction (16°C overnight).

Radiolabelling of AGO–RNA complexes. AGO–RNA complexes were radiolabelled directly with PKNDT treatment in the presence of [γ-32P]-ATP, followed by cold chase, exactly as described 27.

SDS–PAGE and amplification of RNA footprints. SDS–PAGE, nucleaseolus transfer, extraction of AGO-bound RNA, 5’- linker ligation and RT–PCR steps were performed exactly as described 27.

Addition of high-throughput sequencing adapters. Adapters for high-throughput sequencing were added to libraries with additional PCR cycles. PCR conditions were exactly as described, but indexed primers specified in Supplementary Table 7 allowed sample multiplexing. Libraries were sequenced on the Illumina Hiseq 2500 platform with 100-nucleotide single-end reads or on the Illumina Miseq with 75-nucleotide single-end reads.

CLEAR-CLIP with AGO denaturation. AGO–RNA complexes were purified as described up through PKNDT treatment, then eluted from beads with denaturation buffer (50 mM Tris pH 7.5, 0.1% Igepal, 6 µg guanidine HCl, 300 mM NaCl). Samples were diluted fivefold in 1 x PBS/0.1% Igepal and run over a buffer exchange column (Pierce) equilibrated with lysis buffer. AGO–RNA complexes were re-captured on fresh beads conjugated to 2A8 antibody, which was confirmed by western blotting. Subsequent steps were performed as described above.

CLEAR-CLIP mixing experiments. Total E. coli RNA was isolated with the RNAsnap method 56. Either equal amounts or a sixfold excess of E. coli RNA (by mass) was equilibrated in lysis buffer and added to brain lysates. CLEAR-CLIP was then performed exactly as described, starting with DNase treatment. For analyses in Supplementary Fig. 2, RNA was extracted after DNase treatment (with or without RNase) with Trizol and analysed by Bioanalyzer (Agilent) and qRT–PCR. For Dro sophila mixing experiments, lysates from non-cross-linked S2 cells and cross-linked mouse brain containing equal mass amounts RNA were combined immediately post lysis and CLEAR-CLIP was performed starting at DNase treatment.

CLEAR-CLIP in Huh7.5 cells. Huh7.5 CLEAR-CLIP was done as above with the following modifications. Cells (2 x 10^5) growing in 150 mm plates were irradiated once for 400 mJ cm -2 using a Spectrolinker XL-1500 (Spectronics Corporation). Cells were trypsinized, pelleted and stored at -80°C. Lysis was done in 1 ml lysis buffer, RNAse A (0.0004-0.00004 U µl -1; see Supplementary Table 4) or 0.1 U µl -1 RNAse T1 (Ambion) was used for RNase treatment.

AGO HITS-CLIP in Huh7.5 cells. Standard AGO CLIP was done as per the previously published protocol 27, except for multiplexing modifications described above.

Plasmids. pBRET-O-TRE3G-MSPC117 plasmid was constructed by inserting the HSPC117 (c22or28) sequence from pLX304-c22or28EFG (ref. 45) into the doxycycline-inducible retroviral vector pBRET-O-TRE3G (Clontech). The dual-colour reporter vector was described elsewhere 57. Inserts corresponding to CLEAR-CLIP-defined binding sites were synthesized as gblocks (IDT) (Supplementary Table 6) and cloned into the 3’-UTR of tagRFP by Gibson Assembly (NEB) using EcorV-linearized vector and inserts at a 1:5 molar ratio. Transformed clones were grown as maxi-preps at 30°C and confirmed by restriction digests and sequencing.

Mouse miR-125a construct was purchased from SBI (MMIR-125a-PA-1). Genomic fragments for miR-125b, miR-30a and miR-30c containing ~200 nucleotides upstream and downstream of primary hairpins were synthesized as gblocks (IDT) and inserted into the SBI vector between EcoRI and BamHI. Constructs expressing miR-30a from the miR-30c locus and miR-125b from the miR-125a locus were also made, in an effort to control for processing efficiency. However, this expression strategy required a non-sense locus (Supplementary Fig. 10). Therefore, endogenous fragments were used in all reporter experiments. The cel-mir-67 hairpin was cloned into the miR-30c genomic locus. Efficient expression of cel-mir-67 was confirmed by qRT–PCR using the miScript system (not shown).

Cell culture and transfections. N2A mouse neuroblastoma (ATCC) and Huh7.5 human hepatoma cells 58 were maintained in standard conditions.

N2A miRNA mimic ‘reverse’ transfections were done with Dharmafect reagent and mirIDIAN mouse miRNA mimics or negative control mimic #1 (Dharmacon). Complexes were pre-formed in 24-well dishes, according to manufacturer’s instructions, and 120,000 cells per well were added giving a final mimic concentration of 25 nM.

To generate N2A cells stably expressing the Tet-3G activator construct (Clontech), N2A cells were transfected with Xiremeogene 9 (6:1 reagent:plasmid ratio, 375 ng plasmid per 24-well) and split at varying dilutions into G418 media 48 h later. Functional clones were identified by transfecting pTRE-BI-RFP construct and screening for doxycycline-inducible red fluorescent protein (RFP) expression.

For inducible expression of HSPC117, Huh7.5 cells expressing Tet-3G activator (kind gift from C. Takacs) were transfected with pBRET-O-TRE3G-HSPC117. HSPC117 expression was induced by 3 µg mL -1 doxycycline.

For Huh7.5 cell miRNA inhibitor experiments, cells were seeded the day before and transfected with LNA-122 or miravirsen/SPC3649 (5’-C-AAttGtcGcA-3’; LNA in upper case and DNA in lower case). Expression was at 30 nM using RNaseMAX (Life Technologies). No significant cytotoxicity was observed from the applied concentrations of LNA and miravirsen/SPC3649, as determined using CellTiter-Glo (Promega).

qRT–PCR analysis. For miRNA mimic experiments, RNA was extracted from N2A cells 24 h post transfection with Trizol (Ambion). RNA was further purified with DNase treatment on High Pure RNA Isolation columns (Roche). Total RNA (0.5 µg) was reverse transcribed with the iScript kit (Bio Rad). Gene-specific primers (Supplementary Table 7) were designed with Primer3 and tested to confirm efficient amplification of single products 59. The following programme was carried to 40 cycles: 30 s 95°C (denaturation); 30 s 58°C (annealing); and 20 s 72°C (extension). Results were analysed by ΔΔCt, using RPL104a mRNA, an abundant transcript with negligible AGO binding in its 3’-UTR in brain, for normalization.

For E. coli/mouse mixing experiments in Supplementary Fig. 2, RNA was extracted with Trizol (Ambion). Equal volumes re-suspended RNA were reverse transcribed with the iScript kit and analysed by qPCR as above.

Western blotting. For western blotting, 10 µg protein from cleared Huh-7.5 lysates were run per lane of a 4-12% NuPage gel (Life Technologies) and blotted onto polyvinylidene difluoride membrane. HSPC117 was detected using Anti-C22or28 antibody (Abcam, ab98231, 1 µg ml -1) and Goat-anti-Rabbit-HRP (Pierce 31462, 1:50,000).

Flow cytometry. N2A-Tet3G cells were co-transfected with miRNA (250 ng) and reporter (125 ng) plasmids in media with 1 µg ml -1 doxycycline (Sigma). At 24 h media was refreshed and at 48 h cells were trypsinized, harvested and fixed with Cytofix/Cytoperm buffer (BD Biosciences). Cells were analysed on the MACS-Quant cytometer (Miltenyi Biotec). Data were processed as described 60. Briefly, side scatter vs forward scatter were gated in FlowJo software and fluorescence values were exported for analysis with custom R scripts. Cells were binned on the basis of tagRFP fluorescence and mean tagRFP fluorescence was calculated for each bin. Binned tagRFP means were plotted against binned tagBFP means.

Northern blotting. RNA was extracted from transfected N2A cells or brain with Trizol. Thirty micrograms of RNA per sample were run on 15% urea PAGE gels and then transferred to nylon membranes (Pernik Elmer). Hybridization of 32P-labelled DNA oligonucleotide probes (Supplementary Table 7) was done at 37°C in Ultrahyb-Oligo buffer (Ambion) overnight. Membranes were washed four times with 2 x SSC/0.1% SDS and exposed to film.

Bioinformatic analysis. Initial bioinformatic processing was performed exactly as described 27. An additional de-multiplexing step was added after 3’-adapter removal using a simple script for sample-specific indices (Supplementary Table 1). Peak calling for brain AGO HTS-CLIP was done as described, using pooled reads from ten biological samples in the present study and five from a prior one 23.

Identification of miRNA–mRNA chimeras. Reads containing miRNA sequences were identified by ‘reverse’ mapping mature miRNA sequences against sample libraries using Bowtie 60. Changes to default parameters were as follows: maximum mismatches allowed in the seed (n = 1), seed length (l = 8), maximum total of quality values at mismatched read positions (e = 35) and maximum reported alignments (k = 1). Reads mapped to more than one miRNA, usually members of the same miRNA family, were collapsed to a single, randomly chosen hit for initial analyses. Chimeric sequences upstream (5’) and/or downstream (3’) of miRNAs were extracted, filtered for a minimum length of 18 nt and mapped against the appropriate reference genome (mouse mm9, human hg18, Drosophila melanogaster and Danio rerio with Bowtie) at 30 nM using RNaseMAX and LNA. IUPM and LNA conserved hits were allowed and PCR duplicates were consolidated as described 27. Fragments mapping to miRNA genes were removed.
miR-first chimeras in the brain were present in ~14-fold excess of miR-last (Supplementary Table 1). This result differs from reported CLASH results, where miR-first and miR-last species were present at comparable levels. This difference may reflect an idiosyncrasy of AGO1, the only AGO paralogue analysed by CLASH, or denaturation of AGO in the CLASH protocol, which may expose the buried miRNA 5’-end. CLEAR-CLIP, miR-last chimeras frequently involved dubious canonical miRNAs, did not reflect endogenous miRNA abundance and were not formed by exogenous ligase. They were therefore excluded from subsequent analyses. Unique miR-first chimeric reads linked to same miRNA and with overlapping genomic coordinates were clustered together, using the GenomicRanges package in R.61

Analysis of chimeric targets in miRNA perturbation experiments. Normalized microarray values for polyribosome profiles in miR-128 KO and WT mouse brains were obtained from GEO. Genes with contradictory probe information (different symbols) were filtered and probe log2 fold-change (log2FC) values for remaining genes were averaged. For cumulative distribution function (CDF) analysis (Fig. 2a,b), log2FC ratios (KO/WT) in transcript polyosome association were plotted for miR-128 3’-UTR chimera sites. Non-miR-128 3’-UTR chimeras were plotted as controls.

Normalized microarray values for CAD neoblastoma cells transfected with miR-124 or control mimics were obtained from GEO and processed as for miR-128 profiles.88 In Fig. 2c, transcripts were divided into mutually exclusive sets based on the number of times (N = 1 or N > 1, where N is the number of times an interaction was identified by CLEAR-CLIP) the most frequently identified chimera site in their 3’-UTRs occurred. Log2FC ratios (miR-124/Control) were plotted as CDFs. miR-124 sites overlapping AGO-binding peaks, regardless of cluster size (N), were also plotted. To control for non-miR-124, miR-124 (N ≥ 2) and non-miR-124 (N ≤ 1) clusters were tested. The first miRNA nucleotide was assigned a score of 1 and unpaired sites a score of 0.

For LNA-122 or miravirsen-treated HuH.7 cells, standard AGO CLIP data from four biological replicates of each mock, LNA-122 and miravirsen were analysed, with alignment and peak calling as described above. Clusters were normalized to the read depth of their respective libraries after adding a pseudo-count of 1. Canonical miRNA seed searchers were mutually exclusive sets based on the presence of canonical miR-124 chimera sites (with only mismatches or indels up to 75 nt). Canonical miRNAs with the same seed site.

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miR-124 promotes neuronal differentiation by triggering brain-specific regulation.

\[ \text{miR-124} \rightarrow \text{neuronal differentiation} \]

miR-124 regulates adult miRNA interactor by CLASH reveals frequent noncanonical binding.

\[ \text{miR-124} \rightarrow \text{miRNA interactor} \]

miRNA interactome by CLASH reveals frequent noncanonical binding.

\[ \text{miRNA interactome} \rightarrow \text{noncanonical binding} \]

miRNA-mRNA interaction maps.

\[ \text{miRNA} \rightarrow \text{mRNA} \rightarrow \text{interaction maps} \]

miRNA targets using evolutionary conservation and pathway analysis.

\[ \text{miRNA} \rightarrow \text{evolutionary conservation} \rightarrow \text{pathway analysis} \]

DNA binding protein and miRNA target sites by PAR-CLIP.

\[ \text{DNA binding protein} \rightarrow \text{miRNA target sites} \rightarrow \text{PAR-CLIP} \]

mRNA binding network linked to brain development and autism.

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