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Developmental inhibition of *miR-iab8-3p* disrupts mushroom body neuron structure and adult learning ability

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Abstract

MicroRNAs are small non-coding RNAs that inhibit protein expression post-transcriptionally. They have been implicated in many different physiological processes, but little is known about their individual involvement in learning and memory. We recently identified several miRNAs that either increased or decreased intermediate-term memory when inhibited in the central nervous system, including *miR-iab8-3p*. We report here a new developmental role for this miRNA. Blocking the expression of *miR-iab8-3p* during the development of the organism leads to hypertrophy of individual mushroom body neuron soma, a reduction in the field size occupied by axonal projections, and adult intellectual disability. We further identified four potential mRNA targets of *miR-iab8-3p* whose inhibition modulates intermediate-term memory including ceramide phosphoethanolamine synthase, which may account for the behavioral effects produced by *miR-iab8-3p* inhibition. Our results offer important new information on a microRNA required for normal neurodevelopment and the capacity to learn and remember normally.

Keywords

MicroRNA; *Drosophila*; Learning; Memory; Neurodevelopment; Mushroom bodies

1. Introduction

MicroRNAs (miRNAs) are ~21 nucleotide (nt)-long RNAs and part of a family of small non-coding RNAs implicated in many different broad processes extending from normal development, tumorigenesis, to human disease (Bartel, 2004; Krützfeldt and Stofell, 2006; Bushati and Cohen, 2007; Chang and Mendell, 2007; Kloosterman and Plasterk, 2006; Adams et al., 2014). They have spurred much interest due to their unique biological function

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Competing interests

No competing interests declared.

Author contributions

GUB, TGO and RLD designed the project, analyzed the data and wrote the manuscript. GUB performed the experiments. MC performed immunohistochemistry and brain dissociations.

and as possible blood biomarkers for disease progression and therapeutic targets (Hunsberger et al., 2009; Im and Kenny, 2012; Rao et al., 2013).

MiRNAs repress protein expression post-transcriptionally by binding to complementary sequences in the 3' untranslated region (3' UTR) of target mRNAs (Bartel, 2009). About two-thirds of miRNAs found in mammalian species are transcribed from individual genes by RNA polymerase II, but others are processed from intronic sequences in primary transcripts and are called 'miRtrons' (Bartel, 2004; Rodriguez et al., 2004; Filipowicz et al., 2008; Krol et al., 2010). MiRNA gene expression is regulated in ways analogous to protein-coding genes (Krol et al., 2010; Aksoy-Aksel et al., 2014). The primary miRNA transcript (pri-miRNA) is processed in the nucleus by the microprocessor protein complex, Drosha/Pasha, to produce a ~85 nt precursor hairpin miRNA (pre-miRNA). The pre-miRNAs are shuttled by Exportin-5 to the cytoplasm where they are further processed by the Dicer/Argonaute proteins to yield to a mature miRNA duplex. The guide strand of the miRNA duplex is integrated into a ribonucleoprotein complex named the miRNA-induced silencing complex (miRISC) while the passenger strand is, in some cases, degraded (Bartel, 2009). The target sequence in the mRNA, termed the miRNA recognition element (MRE), is usually recognized by a 'seed' sequence located between nt 2–8 in the 5'-end of the miRNA (Bartel, 2009). Depending on the extent of nucleotide complementarity, miRNAs inhibit translation and/or induce mRNA degradation (Bartel, 2009; Jonas and Izaurralde, 2015).

Many studies have shown that miRNAs are involved in neural plasticity and memory formation (Bredy et al., 2011; Aksoy-Aksel et al., 2014) and that miRNA dysregulation is part of the pathophysiology of neurological diseases and neurodevelopmental disorders (Hunsberger et al., 2009; Xu et al., 2010; Wang et al., 2012; Maciotta et al., 2013; Tan et al., 2015). Neurodevelopmental disorders, which account for more than 10% of the burden of disease globally (Durkin et al., 2006; Hunsberger et al., 2009), are characterized by a complex set of phenotypic traits, including learning disabilities and comorbidity with other diseases. A unique feature of miRNAs is that they target multiple mRNAs that could account for such pleiotropy (Hunsberger et al., 2009; Xu et al., 2010; Maciotta et al., 2013). MiRNA dysregulation has been associated with Down's syndrome, Fragile X syndrome, schizophrenia, bipolar disorder, DiGeorge syndrome, Tourette's syndrome, autism spectrum disorders and Rett syndrome (Xu et al., 2010; Feng and Feng, 2011; Olde Loohuis et al., 2012; Nowak and Michlewski, 2013; Saab and Mansuy, 2014; Sun and Shi, 2015). Obtaining insights into how individual miRNAs mediate the development of the central nervous system in ways that impact learning and memory processes may help disentangle the mechanisms underlying these disorders and offer new ways for the management of such disorders.

Two hundred and fifty six miRNAs sequences have been described so far in *Drosophila melanogaster* including 150 with a high level of confidence (www.mirbase.org, Griffiths-Jones, 2004). MiRNA function in memory formation was first deduced from studies showing miRISC pathway involvement in long-term memory formation (Ashraf et al., 2006). Until recently, however, only one specific miRNA, *miR-276a*, was identified as important for the physiology underlying memory formation (Li et al., 2013).

In *Drosophila*, olfactory classical conditioning is a robust type of learning and memory assay (Busto et al., 2010; Kahsai and Zars, 2011; Davis, 2011). In this paradigm, flies learn to associate the noxious stimulus of electric shock, the unconditioned stimulus (US), with an odor, the conditioned stimulus (CS) (Tully and Quinn, 1985; Beck et al., 2000). Although a growing number of neurons and circuits are involved in olfactory memory (Güven-Ozkan and Davis, 2014), a large part of the association between the CS and the US occurs in the mushroom body neurons (MBn; Tomchik and Davis, 2009). To obtain a comprehensive view of the roles for individual miRNAs in memory formation, we recently completed a large screen of ~140 individual miRNAs to identify those that are functionally involved (Busto et al., 2015). We employed the miRNA sponge strategy (Loya et al., 2009; Ebert and Sharp, 2010; Fulga et al., 2015) to selectively inhibit individual miRNAs in a CNS-wide fashion. From these studies, we identified five miRNAs whose inhibition reproducibly modulated intermediate-term memory (Busto et al., 2015). A subsequent secondary screen (unpublished) using two genomic copies of the miRNA sponge to increase potency allowed us to identify one additional miRNA candidate, *miR-iab8-3p*.

Here, we provide data showing that *miR-iab8-3p* function in the developing MBn is required for the adult fly to learn about olfactory cues. This cognitive deficit is associated with a requirement during development for normal *miR-iab8-3p* function. The cellular consequence of inhibiting *miR-iab8-3p* function is that individual MBn exhibit an increase in cell soma size and a decrease in the neuropil volume occupied by their axons. We further identify four potential *miR-iab8-3p* mRNA-targets that are required for normal olfactory memory. We provide evidence and arguments supporting the model that ceramide phosphoethanolamine synthase is a likely effector target for both the normal structural development of the MBn as well as adult learning capacity.

2. Results

2.1. MiR-iab8-3p inhibition in MBn impairs memory formation and expression

We used inducible complementary transcripts, called miRNA-sponges (*uas-miR-SP*) (Fulga et al., 2015), to decrease the expression of *miR-iab8-3p* in the CNS using the *elav^{c155}-gal4* driver (*c155-gal4*). Memory tested at 3 h after conditioning, which is regarded as intermediate-term memory, was impaired when compared to two control genotypes, one carrying the *elav^{c155}-gal4* driver crossed with *miR-scr* and the other carrying only the *uas-miR-SP* transgene (Fig. 1A). The *miR-scr* line carries a scrambled sequence instead of an authentic *miR-SP* inserted into the same vector and at the same defined genomic docking sites (*attP40/attP2*) as the *miR-SP* transgenes. The intermediate-term memory phenotype of the experimental flies (*elav^{c155}-gal4 > uas-miR-SP*) was reproduced in multiple experiments, including that shown in Fig. 1D.

We subsequently mapped the impairing effect of the *miR-iab8-3p-SP* in the CNS by expressing it with a panel of *gal4* drivers with some focus on the MB. Expression of the disrupting sponge sequences with a very specific MB *gal4* driver, *R13F02*, or a *gal4* driver with preferential expression in the MB, *OK107*, reproduced the 3 h memory impairment whereas *gal4* elements that drive expression in other neurons of the olfactory nervous system were ineffective (Fig. 1B–D). One exception to this was the *GAD-gal4* driver, which also

produced modest memory impairment (Fig. 1D). The memory impairment was observed with *R13F02* using flies that contain two copies of the *miR-iab8-3p-SP* or when these were separated into single copies at either the *attP40* or *attP2* integration sites (Fig. 1C), despite the marked difference in performance between the control *miR-scr* inserted at the two different docking sites. We and others have recently shown that luciferase reporters and sponge transgenes can exhibit different expression levels or behavioral performance depending on their *attP* insertion sites (Markstein et al., 2008; Busto et al., 2015). In addition, there exists substantial variation in memory expression for the various *gal4 > miR-scr* controls, presumably due to genetic background effects (Fig. 1D). This restricts the possible comparisons to be between each *gal4 > miR-scr* control and its corresponding *gal4 > miR-SP* experimental genotype.

We tested for normal avoidance behavior to the olfactory cues and shock used during conditioning. Odor and shock avoidances were not significantly different between flies expressing *miR-iab8-3p-SP* or the *miR-scr* control in the MBn, indicating that the impairment in memory expression was not attributable to a defect in odor or shock perception (Table S1).

We next asked whether the 3 h memory impairment was due to a deficit in learning – the acquisition of the odor/shock association – or a deficit in the stability of memory of the association. We submitted flies to an increasing number of electric shocks along with the paired odor during training and measured the performance immediately (3 min) after such conditioning (Fig. 1E). Memory expression was drastically reduced for 1, 2, 3, 6, or 12 shock pairings. These data show that flies expressing *miR-iab8-3p-SP* were able to acquire the task but were limited in their learning abilities, with expression plateauing at a Performance Index (PI) of ~0.4 compared to ~0.8 for control flies. We also tested memory expression at different time points (1, 3 and 6 h) following 12 shock conditioning (Fig. 1F). Memory was reduced for flies expressing *miR-iab8-3p-SP* relative to the *miR-scr* control at 1 and 3 h after conditioning with no memory expression detectable in the experimental group at 6 h. Memory retention was also tested with another odorant combination (BEN/MCH instead of BEN/OCT) and in an alternative genetic background (*w¹¹¹⁸*) with similar results (Fig. S1A,B). We measured the memory retention relative to the 3 min memory score following 12 shock training to discriminate between acquisition and memory stability impairment. Memory expression when transformed in this way was identical between the experimental and control genotypes (Fig. S1C), indicating that acquisition is the primary process impaired by *miR-iab8-3p* inhibition in MBn.

2.2. MiR-iab8-3p inhibition in all classes of MBn reduces three-hour memory

Three major classes of MBn have been identified from their birth during development and according to the axonal projection patterns in the MB lobes (Kunz et al., 2012). The γ neurons are born first in early larval stages followed by the $\alpha'\beta'$ MBn in late larval stages and finally by the $\alpha\beta$ MBn during the pupal stage. The axons of the γ , β' and β MBn extend into the horizontal lobes of the MB while the α' and α MBn extend collateral projections into the vertical lobes. In addition, evidence has accumulated that these MBn subtypes are differentially involved in the sequential phases of memory formation (Davis, 2011). The γ

MBn are most strongly implicated in short-term memory while the $\alpha'\beta'$ MBn are implicated in intermediate-term memory and consolidation. The $\alpha\beta$ MBn are important for long-term memory and for the expression of all temporal phases of memory (Yu et al., 2006; Krashes et al., 2007; Akalal et al., 2011; Cervantes-Sandoval et al., 2013).

To determine whether *miR-iab8-3p* was required in a specific sub-population of MBn for normal adult memory, we expressed *miR-iab8-3p-SP* using a battery of Gal4 drivers expressing in the γ , $\alpha'\beta'$ or $\alpha\beta$ MBn, sampling at least two different Gal4 drivers for each class of MBn (Fig. 2). Although there exists substantial variation in memory expression across the controls for the various Gal4 drivers (see above), the *miR-iab8-3p-SP* expression in all three classes of MBn significantly decreased 3 h memory relative to their genotype-matched *miR-scr* control. These data indicate that normal levels of *miR-iab8-3p* expression are required in all classes of MBn for normal adult memory expression.

2.3. MiR-iab8-3p-SP expression limits acquisition and subsequent memory retention

We focused our subsequent studies on the roles for *miR-iab8-3p* in memory processes such as acquisition and memory retention. We initially tested memory acquisition after expressing *miR-iab8-3p-SP* in the $\alpha\beta$ MBn. Memory expression in the experimental flies was tested immediately (3 min) after conditioning with 1, 2, 3, 6, or 12 electric shock pulses, with the sponge-expressing flies showing poor performance after 3, 6 and 12 shocks conditioning (Fig. 3A). The ceiling level of performance for flies expressing *miR-iab8-3p-SP* was ~ 0.6 while the control group reached a ceiling of ~ 0.75 . These data strongly suggest a deficit in acquisition processes in the experimental group. Similar behavioral results were obtained upon expressing *miR-iab8-3p-SP* in the $\alpha'\beta'$ or γ MBn (Fig. S1A, S3A). This limitation was not associated with a deficit in odor or shock perception as reflected by avoidance behavior (Table S2). Note that the degree of memory impairment with disruption in individual classes of MBn (Fig. 3A, S2A, S3A) was milder than that observed with pan-MB inhibition (Fig. 1E), consistent with a requirement in all classes of MBn. Therefore, these data argue that normal expression of *miR-iab8-3p* in all three classes of MBn is required for normal acquisition.

In addition to our tests of memory retention at 3 h after conditioning (Fig. 2), we tested memory retention at 1 and 6 h after conditioning. The performance of flies expressing *miR-iab8-3p-SP* in the $\alpha\beta$ MBn was reduced at the 1 h time point, probably reflecting the deficit in acquisition (Fig. 3B). This was also true for flies that express *miR-iab8-3p-SP* in the $\alpha'\beta'$ or γ MBn (Fig. S2B, S3B).

Memory expression within a few hours after conditioning has been dissociated in two major components including anesthesia-resistant memory (ARM) and anesthesia-sensitive memory (ASM), named for their sensitivity to cold anesthesia (Isabel et al., 2004). We exposed control and experimental groups of flies to a cold-shock after conditioning in order to remove the ASM component and measure the remaining ARM (Fig. 3C). We observed a similar level of performance following cold-shock in flies expressing *miR-iab8-3p-SP* or *miR-scr*, indicating that ARM was preserved. We conclude from this result that a specific deficit in ASM was produced by *miR-iab8-3p* inhibition (Fig. 3C). Similar conclusions were obtained with *miR-iab8-3p* inhibition in $\alpha'\beta'$ and γ MBn (Fig. S2C, S3C).

We next determined the temporal requirement for *miR-iab8-3p* expression using the TARGET system (Fig. 3D, left panel, McGuire et al., 2003). Raising experimental genotype *Drosophila* throughout development and adulthood at 18 °C (negative control) until conditioning failed to produce a significant memory phenotype (Fig. 3D, right panel). In contrast, culturing the organism throughout development and adulthood at 30 °C induced *miR-iab8-3p-SP* expression (positive control) and impaired memory as previously observed. Culturing the organism at 18 °C across embryonic, larval and pupal stages and imposing a temperature shift to 30 °C upon eclosion to inhibit *miR-iab8-3p* in adult flies failed to produce a phenotype. However, culturing the organism at 30 °C until eclosion, with a transfer to 18 °C for adulthood, reduced memory expression in a quantitative way similar to the positive control. These results indicate that developmental knockdown in the *R28H05-gal4* expressing neurons results in an adult memory deficit. A similar experiment was performed with disruption of expression in the $\alpha'\beta'$ MBn with nearly identical results (Fig. S2D).

2.4. MiR-iab8 has a bi-functional role on memory

The results above showing that diminishing *miR-iab8-3p* expression in the MBn impairs memory beg the question of what occurs when *miR-iab8* is overexpressed. We used the MBn driver *R13F02* to overexpress this *miRNA* and found no change in memory expression when tested at 3 h (Fig. 4A). For this experiment, in which the *uas-miR-iab8* transgene was inserted in the *attP-86Fb* landing site, we compared the performance of the bi-genic progeny to that of progeny carrying only the *gal4* driver or only the *uas-miR-iab8* transgene. We obtained the same conclusion of no phenotype using the *elav^{c155}-gal4* driver (data not shown). However, the *miR-iab8* gene produces two different mature *miRNAs*, including *miR-iab8-3p* and *miR-iab8-5p* (Fig. 4C; <http://www.mirbase.org/>). To determine whether this alternative *miR-iab8-5p* is involved in memory expression, we blocked its expression in the MBn using *miR-iab8-5p-SP* at the same characterized *attP* docking sites used for *miR-iab8-3p-SP* and compared memory expression relative to the *miR-scr* control. Surprisingly, 3 h memory expression was increased by the inhibition of *miR-iab8-5p* in the MBn (Fig. 4B). This result indicates that each strand of *miR-iab8* participates in memory processes with opposite effects, such that over-expression of the complete transgene nullifies the effect of the two encoded *miRNAs* (Fig. 5C).

2.5. MiR-iab8-3p inhibition alters $\alpha\beta$ MBn projections, distribution and cell body size

The precise development of neural circuitry that mediates cognitive abilities is critical for normal cognition during adulthood (Xu et al., 2010; Feng and Feng, 2011; Olde Loohuis et al., 2012; Nowak and Michlewski, 2013; Sun and Shi, 2015). Given that normal *miR-iab8-3p* expression was necessary in $\alpha\beta$ MBn during development for normal adult memory, we asked whether the structure of $\alpha\beta$ MBn in adults was perturbed by *miR-iab8-3p* inhibition during development (Fig. 5).

We first used *mCD8::GFP* expression in the $\alpha\beta$ MBn to gain insight into the global structure of these neurons in adults (Fig. 5A–F). We imaged the MB lobes and calyx (Fig. 5A,B) using confocal microscopy and quantified the GFP signal intensity in sub-stacks for *miR-iab8-3p-SP* or the *miR-scr* expressing flies. We uncovered a significant decrease in GFP

intensity in the lobes and calyces of flies expressing *miR-iab8-3p-SP* (Fig. 5C,E). In addition, the volume occupied by the GFP signal was reduced in flies expressing *miR-iab8-3p-SP* compared to the control (Fig. 5D,F). We also examined the GFP signal intensity across the dorsal/ventral, medial/lateral, and anterior/posterior axes of the MB lobes using a sliding window and found that the distribution intensity was altered slightly in the *miR-iab8-3p-SP* expressing flies across the medial/lateral axis of the vertical lobes (Fig. S4A,B) and across the dorsal/ventral axis of the horizontal lobes (Fig. S4C,D). These results together suggest that *miR-iab8-3p-SP* expression decreases the number and spatial distribution of neurites extending from $\alpha\beta$ MBn.

A decreased number of axonal or dendritic fibers might be due to fewer MB cells. We estimated the number of $\alpha\beta$ MBn cell bodies from image stacks of adult brains. To achieve this goal, we co-expressed *miR-iab8-3p-SP* or *miR-scr* together with a nuclear-localized GFP (*uas-GFP.nls*). We processed images of confocal stacks to obtain an estimate of the number of $\alpha\beta$ MBn nuclei. Each brain (both hemispheres) contained ~2300 $\alpha\beta$ MBn counted in this way, but there was no significant difference between genotypes (Fig. 5G). Surprisingly, we identified a significant increase in the size of the nuclear signal in the *miR-iab8-3p-SP* expressing flies (Fig. 5H). This conclusion was confirmed by measuring the size of $\alpha\beta$ MBn nuclei in dissociated neurons (Fig. S4E,F). We conclude from these structural analyses that MBn cell body size is increased in the *miR-iab8-3p-SP* expressing flies, using the proxy of nuclear area, and that there is a decrease in the neuropil volume occupied by the MBn axonal and dendritic fibers.

To gain a more precise understanding of the effect of *miR-iab8-3p-SP* on the structure of individual $\alpha\beta$ MBn, we activated C3PA-GFP (Patterson and Lippincott-Schwartz, 2002; Ruta et al., 2010; Caron et al., 2013) in the cell body of MBn to visualize the neuronal projections in the $\alpha\beta$ MB lobes (Fig. 5I). The C3PA-GFP was expressed together with *miR-iab8-3p-SP* or *miR-scr* in the MBn. After photoactivation and diffusion of the C3PA-GFP, the projections were manually traced using NeuroLucida and analyzed with NeuroLucida explorer. The total length (cumulative length of all segments) and the number of bifurcations were not affected (Fig. 5J, K). But the global volume occupied by the axonal projections was significantly decreased (Fig. 5L) in *miR-iab8-3p-SP* expressing flies. This result confirmed the decreased cumulative area of the mCD8::GFP signal observed in flies expressing *miR-iab8-3p-SP* (Fig. 5D). We also compared the distribution of axonal segment sizes and found a significant change, with a decrease in the small and large size axonal segments in flies expressing *miR-iab8-3p-SP* (Fig. 5M). We asked whether this difference was due to α or β primary (1°) segments (Fig. 5N) and found that the difference was restricted to the α neurite. Together, these results show that the α lobe neurite is shortened with the expression of *miR-iab8-3p-SP*.

2.6. Potential targets of MiR-iab8-3p

We next sought to obtain preliminary information about the mRNA targets for *miR-iab-3p* that might influence memory formation (Fig. 6A). We used the bio-informatic tool microRNA.org to predict potential mRNA targets based on multiple parameters: sequence complementarity at the seed region and 3' end of the mRNA, A/U composition near the

target site, secondary structure accessibility, length of the UTR, relative position of the target site in the UTR and conservation score (Betel et al., 2010). The micro-RNA.org algorithm predicted 1786 potential mRNA targets (Fig. 6). We selected the top ~5% (105) of these based on mirSVR scores (-1.2) and tested for the involvement of these candidates in memory formation using RNAi interference (Walkinshaw et al., 2015), expressing RNAi transgene for each in the CNS using the pan-CNS, *n-syb-gal4* driver, and subsequently in the MBn (Table S3). Thirty-four genes modulated 3 h memory when inhibited in the CNS, and 4 of these reproducibly influenced memory scores when subsequently tested in the MBn and relative to the 60100-control line, host genotype for producing the RNAi transgenics (Fig. 6C,D and Table S3).

We focused our interest on *CG12229* and *CG4585* because the RNAi knockdowns increased the memory performance, the prediction as an authentic target based on the opposite effect produced by the *miR-iab8-3p-SP* sponge (Fig. 6A). *CG12229* is predicted to code for a pyruvate kinase and *CG4585* an ethanolamine phosphotransferase. Since expression of *miR-iab8-3p-SP* decreases memory when expressed in any of the three major classes of MBn, we tested RNAi knockdowns for *CG12229* and *CG4585* in these same three classes of neurons. *CG12229* increased memory only when knocked down in the $\alpha\beta$ MBn, while *CG4585* increased memory when inhibited in all three classes of MBn (Fig. 6E). These results identify ethanolamine phosphotransferase as an attractive target of *miR-iab8-3p* for mediating perhaps some of the structural and behavioral effects of *miR-iab8-3p* inhibition.

3. Discussion

In the present report, we offer compelling results indicating that *miR-iab8-3p* exerts an important and specific role during development of MBn, a cell population that is critical for memory formation in adult flies (Busto et al., 2010; Davis, 2011). We found that developmental inhibition of *miR-iab8-3p* perturbs the structure of adult $\alpha\beta$ neurons, including an increased cell body size and a decreased volume of neuropil occupancy by MBn axons. The latter cellular phenotype seems primarily attributable to a decrease, specifically in the α lobe, in the length of the very long, and short axonal segments, as we found no difference in branching. It seems probable that the limitation in territory occupied by the MBn axons affects their connectivity with either extrinsic input and/or output neurons, leading to poor learning of olfactory cues with this alteration in circuit arrangement (Perisse et al., 2013).

The *miR-iab8* hairpin (including *miR-iab8-5p* and *-3p*), which is also called *miR-iab4AS*, was first described as a Hox *miRNA* gene, and a member of the bithorax complex (BX-C) necessary for fertility (Bender, 2008; Stark et al., 2008; Tyler et al., 2008). It resides between the *abd-A* and *Abd-B* genes as part of a noncoding RNA (ncRNA) on the antisense strand of the *miR-iab4* locus (Bender, 2008; Stark et al., 2008; Tyler et al., 2008). The *miR-iab8* hairpin decreases Ubx and abd-A protein expression and induces a haltere-to-wing homeotic transformation when ectopically expressed (Stark et al., 2008; Tyler et al., 2008). The primary transcript for *miR-iab8* hairpin is detected in embryos, larvae, pupae and adult male and females (Stark et al., 2008). Deep sequencing results indicate that the *miR-iab8-3p*

strand is expressed at all developmental phases at low levels with some preference in embryos (Ruby et al., 2007; Chung et al., 2008).

Our approach and results highlight the advantages of using *miR-SP* for dissecting *miRNA* function. Although they probably offer a hypomorphic insult, they have the great advantage of targeting individual mature *miRNAs*, while clean genomic knockouts may be difficult when multiple *miRNA* genes are clustered (Ebert and Sharp, 2010). In our specific case, two different mature *miRNAs*, *miR-iab8-3p* and *miR-iab8-5p*, are encoded from the same RNA hairpin embedded in the larger *iab8* noncoding RNA (Bender, 2008; Stark et al., 2008; Tyler et al., 2008). Effects of a genomic deletion would consequently be difficult to interpret due to the loss of multiple *miRNAs*. The deletion of this locus (called *mir*) affects fertility, segment identity (Bender, 2008; Stark et al., 2008; Tyler et al., 2008) and self-righting behavior through the inhibition of *miR-iab4*, the *miRNA* hairpin encoded by the complementary DNA strand of *miR-iab8* (Picao-Osorio et al., 2015). Remarkably, we found that each strand of the *miR-iab8* hairpin, when inhibited, produced an opposite effect on memory performance. This indicates a dual role for this *miRNA* on memory processes and stresses the unique advantage of *miR-SP* to properly dissect the individual role of each strand of the *miRNA* hairpin. Both strands of the bifunctional *miR-9* have been shown to regulate two molecular components (REST and CoREST) of the same repressor complex (Packer et al., 2008). This precedent offers the attractive hypothesis that *miR-iab8* is similarly bifunctional and with both strands acting in opposite ways during the development of the MBn. Although intriguing, we currently have no data speaking to the mechanism whereby *miR-iab8-5p* disruption produces memory enhancement.

The effect of *miR-iab8-3p* inhibition on memory was primarily restricted to the MBn, although we observed a slight effect with inhibition in GABAergic neurons as well. Thus, the effects are largely localized to the MBn population. This is important as it shows that *miR-iab8-3p* has cell-type specific roles during development rather than exerting an effect on target genes in all types of brain neurons. However, we did find a role for this *miRNA* in all three major classes of MBn, suggesting that *miR-iab8-3p* function is engaged when neurons are differentiated into MBn and before they become further differentiated into one of the three individual classes of MBn.

Individual *miRNAs* serve to orchestrate the expression level of many target genes in tissue and developmental time-specific ways. This makes the task of identifying the specific effectors of any *miRNA*-associated phenotype a long and difficult task, especially given the possibility that a given phenotype may emerge from changing the expression level of multiple target effectors. But as an initial step in this direction, we chose to provide a preliminary identification of *miR-iab8-3p* targets using a behavioral approach rather than through mRNA quantification since some *miRNAs* decrease protein expression without changing mRNA expression (Griggs et al., 2013). In addition, the approach interrogates the important endpoint of function, rather than simple changes in abundance that may or may not relate to function.

We identified four genes whose inhibition in MBn modulated memory formation. These include *Efa.1*, *MadM*, *CG12229* and *CG4585*. We focused on the two genes whose

inhibition increased memory as predicted by our model (Fig. 6A). We provide three arguments for favoring *CG4585* as a major effector of the *miR-iab8-3p* role in learning. First, *CG4585* improved memory when inhibited. Second, the effect was obtained with inhibition in all three classes of MBn. Third, the function of *CG4585* aligns with the phenotypic observations.

Drosophila sphingolipids are critical for normal development with metabolic intermediates being involved in signal transduction cascades and progressive neurodegenerative disorders (Kraut, 2011). *Drosophila* lacks sphingomyelin (SM) but instead synthesizes the SM analog ceramide phosphoethanolamine (CPE) from ceramide (Vacaru et al., 2013). In general, dysfunction of sphingolipid biosynthesis leads to the accumulation of lipid substrates leading to cellular enlargement, dementia, and intellectual disability (Mencarelli and Martinez-Martinez, 2013). Interestingly, *CG4585* has recently been shown to possess a CPE synthase activity from expression experiments in cultured S2 insect cells (Vacaru et al., 2013). *MiR-iab8-3p* inhibition should increase *CG4585* expression leading to an excess of CPE production, analogous to the accumulation of SM observed in Niemann Pick's disease (Mencarelli and Martinez-Martinez, 2013). These observations are generally consistent with the phenotypes of neuronal enlargement and behavioral alterations observed with *miR-iab8-3p* inhibition. An immediate goal from these results will be to further characterize the link between *miR-iab8-3p* and *CG4585*.

Our results begin to fill an important gap in knowledge about *Drosophila* learning and memory – the identification and characterization of genetic factors that influence the development of the nervous system important for adult cognitive functions. Past research on memory formation in *Drosophila* has emphasized biological functions that influence the adult physiology of MBn critical for memory formation (Davis, 2005, 2011). The critical experimental distinction in showing developmental vs adult physiological requirement has been temporal knockdown or rescue experiments, typified by adult-specific rescue of *dunce* and *rutabaga* functions (Dauwalder and Davis, 1995; McGuire et al., 2003; Mao et al., 2004). Understanding developmental genetic functions is critically important to elucidate the molecular logic underlying the development of adult brain structures important for cognition, such as MBn here for olfactory learning and help gain potential insights into the myriad of human neurodevelopmental disorders such as intellectual disability and autism.

4. Materials and methods

4.1. Fly lines

Drosophila was raised using standard food at room temperature. Fly crosses were performed at 25 °C with ~70% relative humidity across a 12 h light/12 h dark cycle. The *uas-miR-iab8-3p-SP*, *uas-miR-iab8-5p-SP* and *uas-miR-scr* fly lines were previously described (Busto et al., 2015; Fulga et al., 2015). These first two *uas* lines were crossed to various Gal4 lines to establish the experimental genotype. The control genotype, in general, was made by crossing the same Gal4 line to *uas-miR-scr*, a transgenic line containing scrambled sequences in the same vector and inserted at the same docking site (*attP40/attP2*) in the genome as the *uas-miR-SP* lines. The *uas-miR-iab8* over-expression line was backcrossed in the *w^{CS10}* background for six generations. The other lines used in this study include:

elav^{c155}-gal4 (Lin and Goodman, 1994), *R13F02-gal4* (Jenett et al., 2012), *uas-mCD8::GFP* (Lee and Luo, 1999), *Or83b(orco)-gal4* (Wang et al., 2003), *GHI46-gal4* (Stocker et al., 1997), *C316-gal4* (Waddell et al., 2000), *TH-gal4* (Friggi-Grelin et al., 2003), *Gad-gal4* (Ng et al., 2002), *OK107-gal4* (Connolly et al., 1996), *MZ604-gal4* (Suster et al., 2003), *NP2492-gal4* (Tanaka et al., 2008), *R11D09-gal4* (Jenett et al., 2012), *R84G09-gal4* (Jenett et al., 2012), *c305a-gal4* (Armstrong et al., 2006), *R35B12-gal4* (Jenett et al., 2012), *R28H05-gal4* (Jenett et al., 2012), *c739-gal4* (O'Dell et al., 1995), *tub-Gal80ts* (McGuire et al., 2003), *uas-GFP.nls* (Shiga et al., 1996), *n-syb-gal4* (Pauli et al., 2008), *uas-dicer2* (Dietzl et al., 2007) and *uas-PA-GFP* (Patterson and Lippincott-Schwartz, 2002; Ruta et al., 2010; Caron et al., 2013). RNA interference (*uas-RNAi*) lines, and their control line (*60100*, host genotype used to construct the *uas-RNAi* transgenics), were obtained from the KK library of the Vienna Drosophila RNAi Center (VDRC) (Dietzl et al., 2007).

4.2. Behavioral analyses

One-to-four day old flies were collected for behavioral experiments ~16-to-24 h prior to experiments. Thirty minutes before conditioning, flies were transferred in fresh food vials and put in the behavioral test room to adapt to the conditions (dim red light, 25 °C, ~80% relative humidity). Groups of ~65 flies were trained using the standard aversive olfactory conditioning test already described (Beck et al., 2000). Flies were subjected to a succession of two odor stimuli lasting 1 min each and separated by 30 s. The first odor stimulus (CS+) was associated with 12 repeats of a 90-V electric shock (US). The second odor stimulus was not paired with electric shocks and thus constituted a non-conditioned stimulus (CS-). The odorants used were benzaldehyde (BEN), 3-octanol (OCT) or 4-methylcyclohexanol (MCH) when specified. They were diluted in mineral oil at concentrations of ~0.05, ~0.2% and ~0.1%, respectively. Memory retention was tested by allowing flies to choose for 2 min between the sides of a T-maze containing the CS+ odor in one arm and CS- odor in the alternative arm.

Memory acquisition was tested giving an increasing number (from 1 to 12) of electric shocks during the CS+ stimulation (Fig. 1E, bottom). Memory was tested immediately following training. Anesthesia-sensitive (ASM) and anesthesia-resistant (ARM) memories were tested 3 h following conditioning (Fig. 3C, bottom). These two constituents of memory were distinguished by giving a brief cold-shock induced anesthesia (2 min, 4 °C) 2 h following conditioning to erase ASM. After that period, flies were transferred back in their food vial and kept unperturbed for an additional hour before the memory test. For the TARGET experiments (Fig. 3D, left panel), three-to-five day old flies were used; conditioning and memory tests were performed at 25 °C (McGuire et al., 2003).

To test odor avoidance, naive flies were allowed to distribute for 2 min between the two arms of the T-maze with an odor stream on one side and a non-odorized air stream on the other. To test shock avoidance, naive flies were allowed to distribute for 2 min between the two arms of a T-maze containing copper grids with only one side being electrified. An avoidance index was computed with the following formula: [(number of flies in neutral arm) – (number of flies in odor/shocked arm)]/(total number of flies in both arms)].

4.3. Immunohistochemistry

We followed the protocol described by the Fly Light Project (Jenett et al., 2012) using 1–6 day old female fly brains. The primary antibodies used were rabbit anti-GFP (1:1000, Invitrogen) and mouse anti-nc82 (1:50; DSHB). The secondary antibodies included Alexa 633 anti-mouse (1:400) and Alexa 488 anti-rabbit (1:800). Images were obtained using Leica TCS SP5 and SP8 confocal microscopes using the exact same set-up for both experimental groups.

4.4. Image analysis

Analyses of GFP immunohistochemistry on confocal images were performed using ImageJ (Abramoff et al., 2004). For each brain, the cumulative GFP signal intensity was established by calculating the average intensity of the GFP signal for each confocal plane using the ‘mean gray value’ function. The average intensity values for each confocal plane were then summed for all the brains to obtain a quantitative value of GFP intensity representing the GFP signal throughout the brain. To assess the area spanned by the GFP signal, we used the same intensity threshold for all brains using the ‘threshold’ function of ImageJ. The area covered by the GFP signal was calculated for each confocal plan and then summed for the all brain. To gain access to the distribution of the GFP signal in the three dimensions of the MB lobes, we made a maximal projection of the confocal stack of the lobes. We used the ‘plot profile’ function to obtain an average of the GFP signal every 0.36 μm (0.49 μm for the antero-posterior axis) along the axis of our choice. Samples were aligned using the section of the lobe with the highest GFP intensity value. Only regions represented with $n = 7$ were considered for statistical analysis. Average intensity was plotted relative to the highest intensity region.

To estimate the cell body number, each original image was first processed using the ‘unsharp mask’ function of ImageJ to highlight the nuclei. A ‘threshold’ was then imposed to isolate the nuclei from the background. The ‘despeckle’ function was then used to remove the isolated pixels. The ‘watershed’ function was used to isolate individual objects. Eventually, we used the ‘analyze particles’ tools to count objects. Values were summed for the all stacks.

4.5. Brain dissociation

One-to-six day old individual female fly brains were dissected in insect’s Ringer solution, the optic lobes removed, and brains washed once with dissecting solution. Brains were then incubated for 30 min in activated papain solution then replaced with insect’s Ringer solution and triturated in a microcentrifuge tube first with a 200 μl and then a 20 μl pipet tip. Cells were ultimately transferred in 384 well plate and allowed to settle at the bottom of a well for 3 h and then centrifuged at 12.000 RPM for 5 min. Neurons were imaged for GFP with the GE Incell 6000 high-content screening microscope at 10 \times . Cells were counted and measured using custom segmentation algorithms in GE Developer.

4.6. Photo-activation experiments

One-to-six day old individual female fly brains were dissected in ice-cold saline (in mM NaCl 124, KCl 3, MOPS 20, CaCl₂ 1.5, MgCl₂(H₂O)₆ 4, NaHCO₃ 5, NaH₂PO₄(H₂O) 1, trehalose 10, sucrose 7, glucose 10). Brains were subsequently transferred and stuck at the

bottom of a petri dish filled with saline. Individual MBn cell body was identified using the 488 nm laser of a confocal microscope (Leica). A circular 1-by-1 μm region of interest was defined at the center of a MBn and C3PA-GFP activated using a multi-photon laser (Coherent Inc.) set at 710 nm. The photoactivation stimuli consisted of three stimuli (0.325 ms each) separated by 2 min and for 1 h (90 stimuli total) with a 40x objective. The photoactivation laser power was typically between 4 and 40 mW at the objective. Brains were later flipped and axonal MBn projections imaged using the 63 \times objective with the confocal set at 488 nm.

Neuronal projections were manually traced using the NeuroLucida software (MBF Bioscience). Total projections length, bifurcations, neuronal projection volume and length of the segments were computed using the 'Convex Hull Analysis' and 'Branched Structure Analysis' functions of the NeuroLucida explorer software (MBF bioscience). α' , β' and γ neurons were discarded as the neurons with no C3PA-GFP diffusion in the calyx or in the lobes or with major gaps or unresolvable structures.

4.7. Statistics

Statistical analyses were performed using XLSTAT[®], Prism[®] and R (R Core Team, 2008). Memory Performance Indices are known to follow a normal distribution (Tully et al., 1994). Normality of samples was additionally assessed using the Kolmogorov-Smirnov test or the D'Agostino & Pearson test when possible. When average PIs of two groups were compared we used two-sample, bilateral Student *t*-tests if samples followed a normal distribution. In cases where they did not, we used Mann-Whitney tests for non-parametric data. In case where the variances were not the same between samples we used Welch's correction for inequality of variances. To compare the average performance of a group with a theoretical value, we used one-sample bilateral Student *t*-tests. For multiple group comparisons, we used ANOVA followed by Bonferroni's or two-way ANOVA followed by Tukey's post hoc tests corrected for multiple comparisons. To compare distribution and proportions we used χ -square tests. For confocal image analysis, one outlier was removed using the Grubb's test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at <http://dx.doi.org/10.1016/j.ydbio.2016.09.010>.

Summary statement

MiR-iab8-3p is necessary during development for normal olfactory learning. Inhibition of this miRNA alters the size of mushroom body neurons and their axonal projections, which impacts learning abilities in the adult.

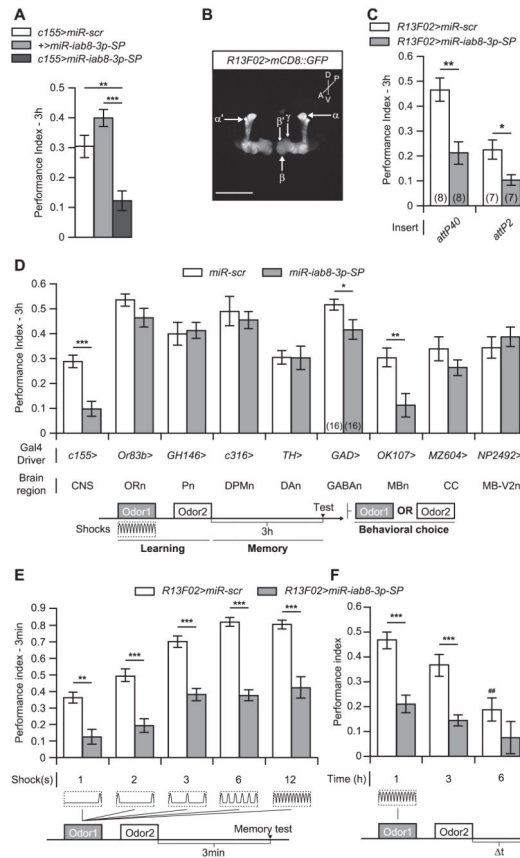


Fig. 1. *MiR-iab8-3p* inhibition in MBn impairs memory formation

(A) Three hour memory was impaired by *miR-iab8-3p-SP* expression in the CNS driven by the *elav^{c155}-gal4* driver. Memory expression was significantly different from flies carrying only the *uas-miR-iab8-3p-SP* transgene or the *gal4* driver crossed with the *miR-scr* control. Results are presented as the mean \pm s.e.m. with $n = 6$. ANOVA was used to measure the effect of genotype on PIs followed by a Bonferroni's post hoc test to compare performances between groups. ***: $p < 0.001$, **: $p < 0.01$. (B) The *R13F02-gal4* element drives expression of a *uas-mCD8::GFP* reporter specifically in the MBn. MBn are classified as three major types based on their axonal projection pattern: $\alpha\beta$, $\alpha'\beta'$ and γ MBn. The $\alpha\beta$ and $\alpha'\beta'$ MBn extend their axonal projections into the vertical and the horizontal lobes while the γ MBn extend axons into the horizontal lobe. Scale bar: 100 μ m. (C) The *uas-miR-iab8-3p-SP* transgenes, either on the 2nd chromosome at the *attP40* docking site or on the 3rd chromosome at the *attP2* docking site, reduced 3 h memory when expressed in the MBn. For each case, the *miR-scr* control transgene was inserted at the same *attP* docking site. Results are presented as the mean \pm s.e.m. with $n = 7-8$. Two-sample, bilateral Student *t*-tests were used to compare PIs between the two genotypes. **: $p < 0.01$, *: $p < 0.05$. (D) *MiR-iab8-3p-SP* impaired 3 h memory when expressed in the MBn and weakly in GABAergic neurons. *MiR-iab8-3p-SP* was expressed in specific types of neurons and in different brain regions and parts of the olfactory nervous system using different *gal4* driver lines. Expression domains for the *gal4* drivers: CNS, central nervous system; ORn, olfactory receptor neurons; Pn, projection neurons; DPMn, dorsal paired medial neurons; DAN, dopaminergic neurons;

GABAergic neurons; MBn, mushroom body neurons; CC, central complex; MB-V2n, mushroom body extrinsic neurons V2. Three hour memory performance for flies carrying a *gal4* transgene and *uas-miR-iab8-3p-SP* was compared to flies carrying *miR-scr* and the same *gal4* element. Results are presented as the mean±s.e.m. with n = 12–16. Two-sample, bilateral Student *t*-tests were used to assess an effect of the genotype on PIs. ***: p<0.001, **: p<0.01, *: p<0.05. (E) *MiR-iab8-3p-SP* dramatically impaired memory acquisition when expressed in MBn. Memory acquisition was assessed by testing immediate performance after an increasing number of electric shocks (from 1 to 12, bottom) paired with odor during training. Although flies expressing *miR-iab8-3p-SP* learned the task, their performance plateaued at PI~0.4, while control flies expressing *miR-scr* plateaued at ~0.8. For all training protocols used, immediate performance was reduced with *miR-iab8-3p-SP* expression in the MBn. Results are presented as the mean±s.e. m. with n = 8. Two-sample, bilateral Student *t*-tests were used to assess the effect of genotype on PIs. ***: p<0.001, **: p<0.01. (F) *MiR-iab8-3p-SP* impaired memory retention when expressed in the MBn. Memory retention was measured at 1, 3 and 6 h following a 12 shock conditioning protocol. Memory expression was reduced to less than half of the control value at all time points tested. The PI at 6 h for the experimental group was not significantly different from zero. Results are presented as the mean±s.e.m. with n = 8. Two-sample, bilateral Student *t*-tests were used to compare PIs between the two genotypes while a one-sample, bilateral Student *t*-test was used to compare PIs with the null value. ***: p<0.001, **: p<0.01 for two-samples. #: p<0.01 for one-sample.

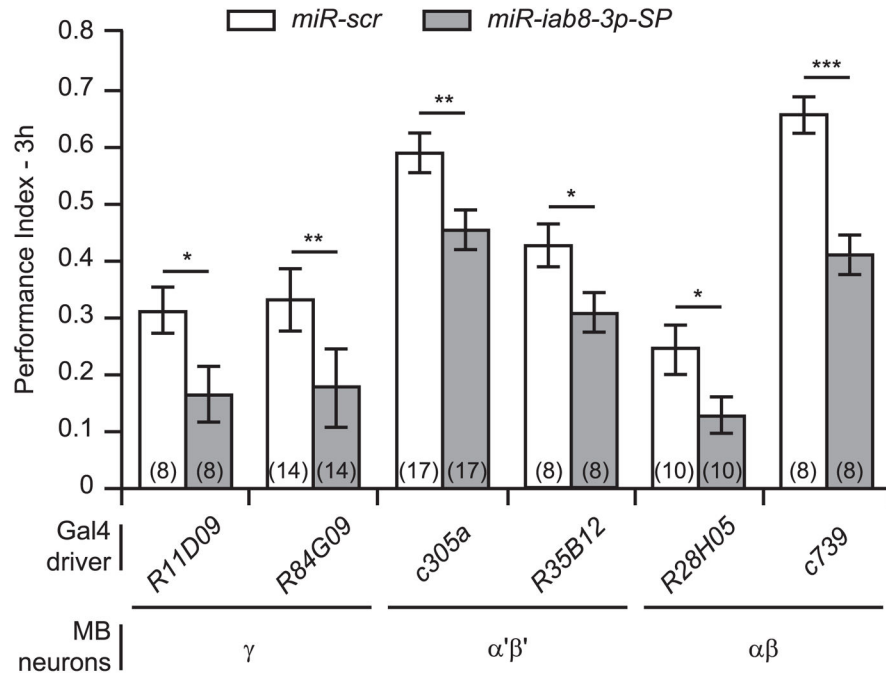


Fig. 2. *MiR-iab8-3p* inhibition in all three classes of MBn impairs 3 h memory expression
MiR-iab8-3p inhibition specifically in the γ neurons using the *R11D09*- and *R84G09-gal4* driver lines reduced 3 h memory. *MiR-iab8-3p* inhibition in $\alpha'\beta'$ neurons using *c305a*- and *R35B12-gal4* drivers also reduced 3 h memory. In addition, *miR-iab8-3p* inhibition in $\alpha\beta$ MBn using the *c739*- and *R28H05-gal4* drivers decreased 3 h memory. Results are presented as the mean \pm s.e.m. with $n = 8$ – 17 . Two-sample, bilateral Student *t*-tests were used to assess significant PI differences between the two genotypes. ***: $p < 0.001$, **: $p < 0.01$, *: $p < 0.05$.

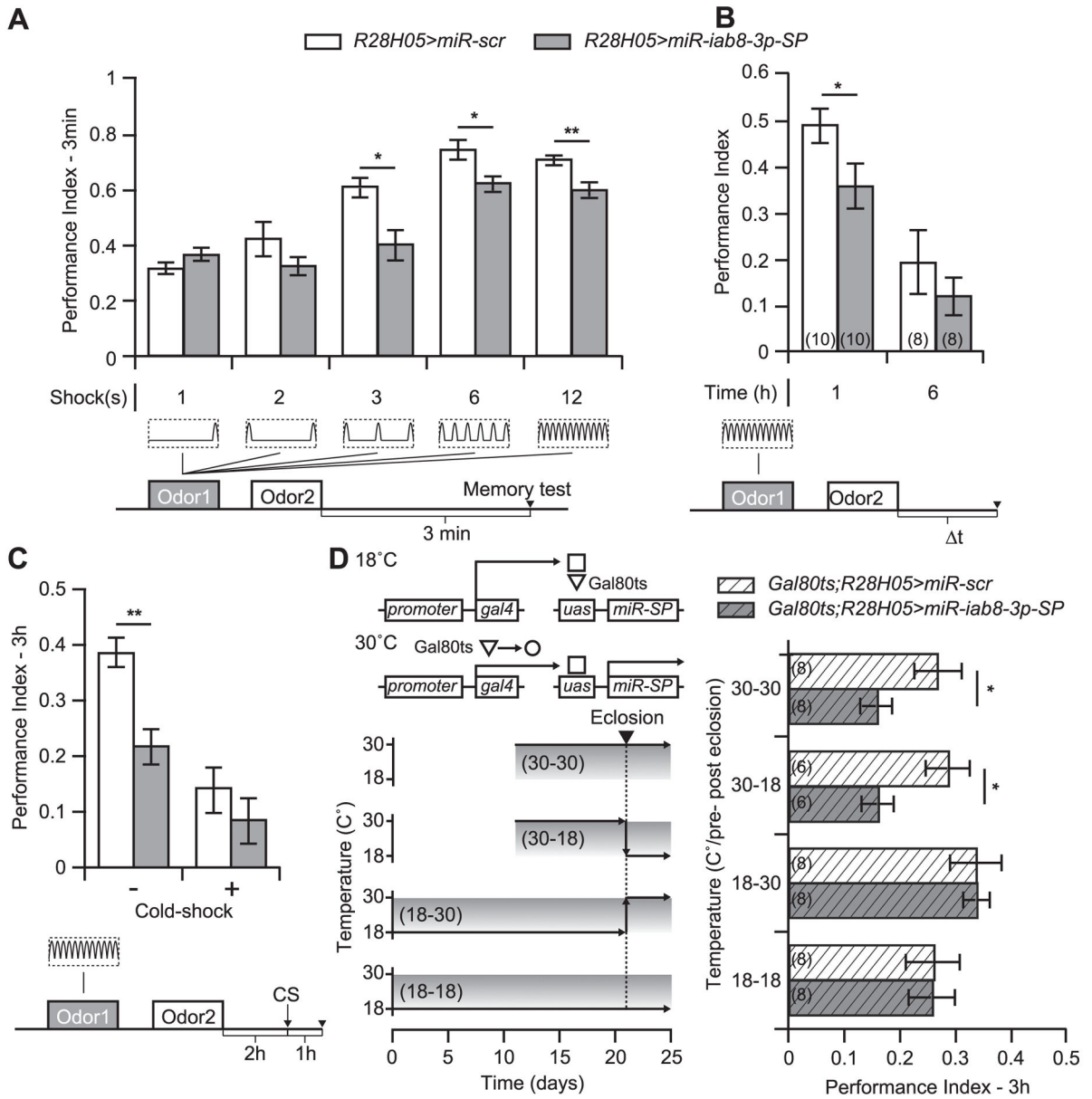


Fig. 3. Inhibition of *miR-iab8-3p* in $\alpha\beta$ MBN impairs memory in adults

(A) Memory acquisition was constrained by *miR-iab8-3p* inhibition in $\alpha\beta$ MBN compared to the *miR-scr* control group. Three-minute memory was assessed after conditioning flies with an increasing number of electric shocks (1-12, bottom panel). After 1 and 2 shocks, both genotypes exhibited similar performance. For increasing shock numbers, flies expressing the *miR-iab8-3p-SP* exhibited reduced memory performance. Results are presented as the mean \pm s.e.m. with $n = 6$. Two-sample, bilateral Student *t*-tests were used to test the effect of the genotype on PIs. **: $p < 0.01$; *: $p < 0.05$. (B) Memory retention was reduced by *miR-iab8-3p-SP* expression in $\alpha\beta$ MBN. Memory retention was measured at 1 and 6 h following a 12 shocks conditioning protocol. One hour after training flies expressing *miR-iab8-3p-SP* exhibited a significant decrease in memory performance relative to flies expressing *miR-scr*. After 6 h, memory performance was nearly zero and indistinguishable between genotypes.

Results are presented as the mean±s.e.m. with n = 8–10. Two-sample, bilateral Student *t*-tests were used to test the effect of the genotype on PIs. **: p<0.01; *: p<0.05. (C) Anesthesia-Resistant Memory (ARM) was not affected by *miR-iab8-3p-SP* expression. A cold-shock (CS, bottom panel) was delivered 2 h after training to remove ASM. The residual memory (ARM) was tested 1 h later. *MiR-iab8-3p-SP* expression had no effect on ARM but significantly reduced ASM. Results are presented as the mean±s.e.m. with n = 6. Two-way ANOVA was used to assess the effects of the genotype and cold-shock on PIs. Multiple comparisons between groups were performed using Tukey's post hoc tests. **: p<0.01. (D) *MiR-iab8-3p* is required in αβ MBn during development. The Gal80^{ts} protein (TARGET system) allows temporal control of *miR-iab8-3p-SP* expression through control of environmental temperature across the life cycle starting at mating (Time 0). At 18 °C, Gal80^{ts} and Gal4 proteins interact together preventing Gal4 binding to the *uas* sequences and consequently preventing *miR-SP* expression. When the organism is cultured at 30 °C, Gal80^{ts} undergoes a conformational change releasing Gal4 from inhibition. Gal4 is thus free to bind to the *uas* sequences and promote *miR-SP* expression (top of the left panel). Temperature shifts between 18 °C and 30 °C were performed at eclosion (dotted vertical line). For flies grown at 18 °C across the life cycle (18-18, bottom), *miR-iab8-3p-SP* expression is inhibited and memory performance is similar to the *miR-scr* control. When flies are cultured at 30 °C across the life cycle (30-30), *miR-iab8-3p-SP* is expressed causing a reduction in memory performance relative to *miR-scr*. To determine whether *miR-iab8-3p* is necessary before or after eclosion, two groups were generated and cultured at 30 °C prior to eclosion (30-18), or after eclosion (18–30). *MiR-iab8-3p* inhibition restricted to developmental periods prior to eclosion reduced memory and phenocopied the flies kept at 30 °C across the life cycle. *MiR-iab8-3p-SP* expression restricted to periods after eclosion had no effect on 3 h memory. Results are presented as the mean±s.e.m. with n = 6–8. Two-sample, bilateral Student *t*-tests were used to test the effect of the genotype on PIs. *: p<0.05.

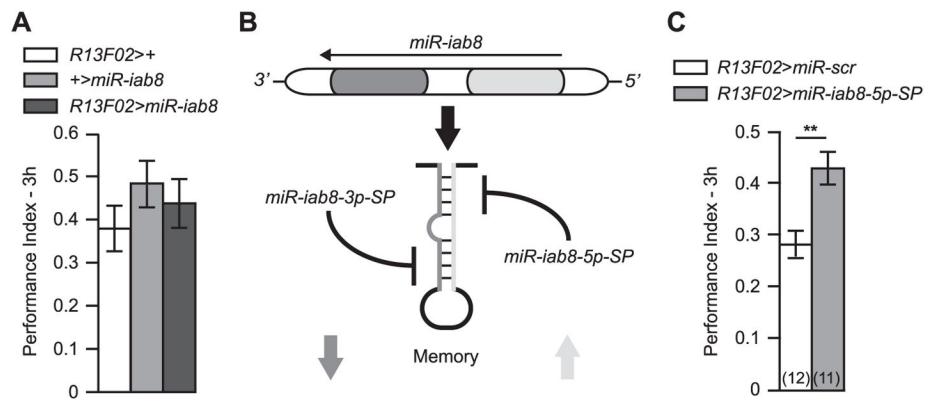


Fig. 4. *MiR-iab8* has a bi-functional effect on memory

(A) Overexpression of *miR-iab8* in the MBn had no effect on 3 h memory expression relative to control genotypes carrying only the *R13F02-gal4* driver or the *miR-iab8* transgene. Results are presented as the mean \pm s.e.m. with $n = 6$. ANOVA was used to test the effect of genotype on PI. (B) Each *miRNA* hairpin is composed of two *miRNAs*, one called the guide strand and the other called the passenger strand. In most cases, the guide strand is integrated in the miRISC while the passenger strand is degraded. In some cases, both strands can be integrated into the miRISC to regulate protein expression. (C) *MiR-iab8-5p* inhibition in MBn increases 3 h memory. *MiR-iab8-5p-SP* expression in MBn using the *R13F02-gal4* driver increased 3 h memory relative to the expression of the *miR-scr* control. Results are presented as the mean \pm s.e.m. with $n = 11-12$. Two-sample, bilateral Student *t*-test was used to compare the two genotypes. **: $p < 0.01$.

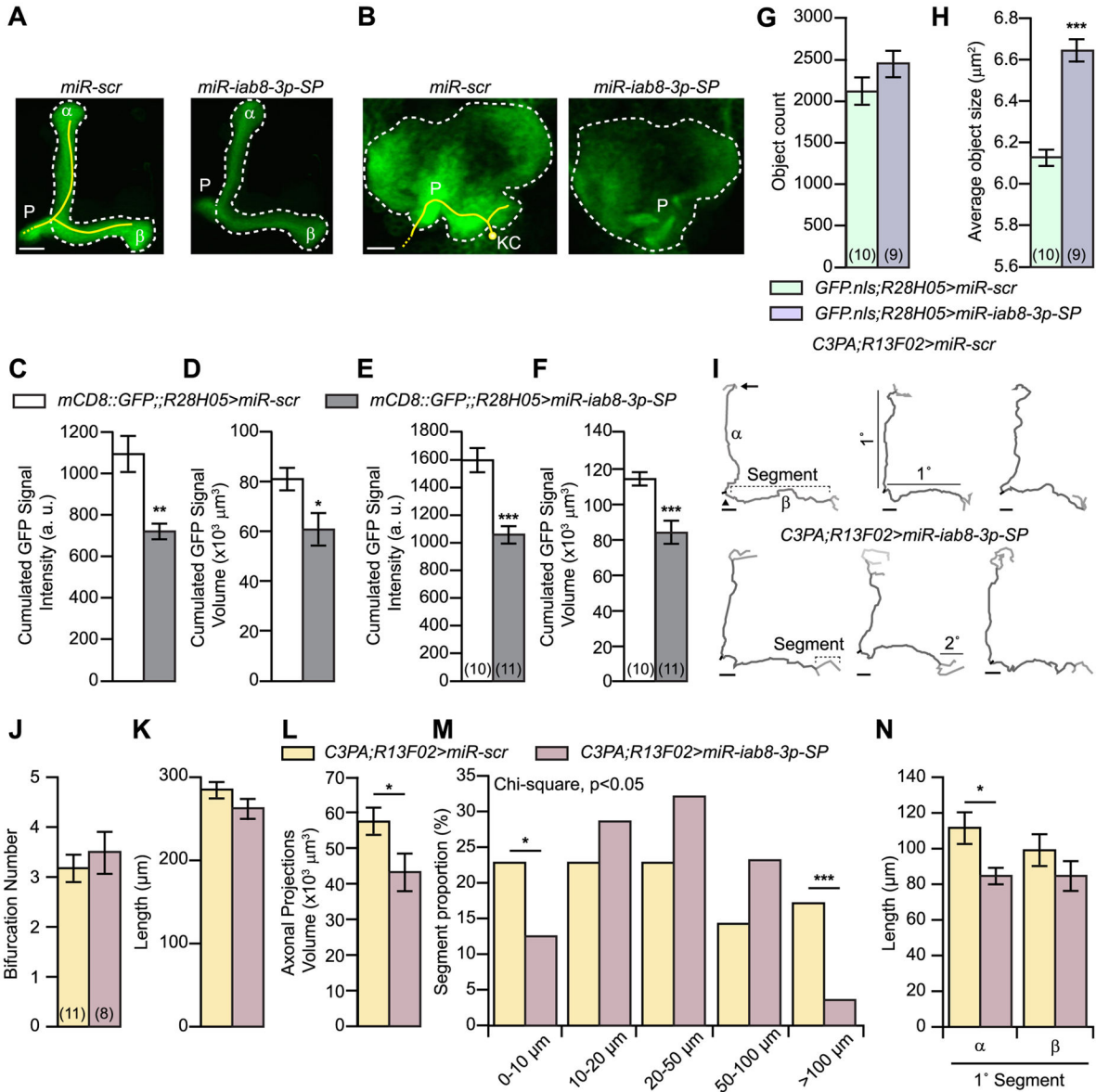


Fig. 5. *MiR-iab8-3p* inhibition alters the structure of $\alpha\beta$ MBn

(A) Representative maximal projection image of a confocal stack of the right $\alpha\beta$ MB lobes of flies co-expressing *mCD8::GFP* with *miR-scr* (left panel) or *miR-iab8-3p-SP* (right panel) and driven by the *R28H05-gal4* driver. The white dashed line was added to help visualize the contour of the lobes and a yellow line to illustrate the projections from a single MBn neuron. α : alpha lobe, β : beta lobe, P: peduncle. Scale bar: 10 μm . (B) Representative maximal projection image of a confocal stack of the right $\alpha\beta$ MB calyx from flies co-expressing *mCD8::GFP* with *miR-scr* (left panel) or *miR-iab8-3p-SP* (right panel) and driven by *R28H05-gal4* driver. A yellow line was added to help visualize the projection in the MB calyx from a single neuron and a white dashed line to identify the boundaries for the calyx. KC: Kenyon Cell, P: peduncle. Scale bar: 10 μm . (C) *MiR-iab8-3p-SP* expression decreases *mCD8::GFP* intensity in the $\alpha\beta$ MB lobes. The *mCD8::GFP* average intensity was

calculated for each slice of a stack and then summed across all slices. Results are presented as the mean \pm s.e.m. with $n = 11$. A two-sample, bilateral Student t -test was used to compare the two genotypes with Welch's correction to account for the inequality of the variances. ***: $p < 0.01$. (D) *MiR-iab8-3p-SP* expression decreased the signal area identifying the $\alpha\beta$ MB lobes. The area covered by the mCD8::GFP signal was computed for each slice of the stack and summed across slices. The signal was thresholded using the same cut-off intensity value. *MiR-iab8-3p* inhibition reduced the volume occupied by the mCD8::GFP signal in the $\alpha\beta$ MB lobes. Results are presented as the mean \pm s.e.m. with $n = 11$. A two-sample, bilateral Student t -test was used to compare the two genotypes. *: $p < 0.05$. (E) *MiR-iab8-3p-SP* expression decreases mCD8::GFP intensity in the calyx. The mCD8::GFP average intensity was calculated for each slice of a stack and then summed across all slices. Results are presented as the mean \pm s.e.m. with $n = 10-11$. A two-sample, bilateral Student t -test was used to compare the two genotypes. ***: $p < 0.001$. (F) *MiR-iab8-3p-SP* expression decreased the signal area identifying the calyx of $\alpha\beta$ MBn. The area covered by the mCD8::GFP signal was computed for each slice of the stack and summed across slices. Results are presented as the mean \pm s.e.m. with $n = 10-11$. A two-sample, bilateral Student t -test was used to compare the two genotypes. ***: $p < 0.001$. (G) *MiR-iab8-3p-SP* expression had no significant effect on the total number of $\alpha\beta$ MBn. The number of neurons estimated in flies expressing *miR-iab8-3p-SP* was not significantly different from flies expressing the *miR-scr* control. Results are presented as the mean \pm s.e.m. with $n = 9-10$. (H) *MiR-iab8-3p-SP* expression increased the size of $\alpha\beta$ MBn nuclei as estimated from the GFP.nls signal. The area occupied by the nuclei of the $\alpha\beta$ MBn was increased in flies expressing *miR-iab8-3p-SP* relative to flies expressing the *miR-scr* control. Results are presented as the mean \pm s.e.m. with $n = 9-10$. A two-sample, bilateral Student t -test was used to compare the two genotypes. ***: $p < 0.001$. (I) Representative traces of axonal projections of the $\alpha\beta$ neurons obtained after photo-conversion of C3PA-GFP in the cell body. The top panel shows three traces from flies expressing *miR-scr* together with *uas-C3PA-GFP* in MBn. The lower panels show ones obtained from flies with *miR-iab8-3p-SP* expressed in the MBn. The arrowhead identifies the initial bifurcation of the axon entering the lobes from the peduncle. The arrow identifies a bifurcation of the axon in the lobe. A segment is defined as the three dimensional distance of the axon located between two successive bifurcations or between a bifurcation and the end of a projection. Segments lengths were located between 1.4 and 161.8 μm and were ranked accordingly by class. α : alpha lobe, β : beta lobe. 1 $^\circ$: primary segments emerging from the initial bifurcation. Scale bars: 10 μm . (J) *MiR-iab8-3p-SP* expression had no effect on the number of bifurcations of $\alpha\beta$ MBn in the $\alpha\beta$ lobes. This number includes the initial bifurcation of the peduncular axon into the vertical and horizontal projections. The number of bifurcations was assessed using the 'Branched Structure Analysis' of the NeuroLucida explorer software after tracing the projections. Results are presented as the mean \pm s.e. m. with $n = 8-11$. (K) *MiR-iab8-3p-SP* expression has no effect on the total length of the $\alpha\beta$ MBn axonal projections in the $\alpha\beta$ lobes. The length was assessed using the 'Branched Structure Analysis' of the NeuroLucida explorer software after tracing the projections. Results are presented as the mean \pm s.e.m. with $n = 8-11$. (L) *MiR-iab8-3p-SP* expression decreased the neuropil volume occupied by the axonal projections in the $\alpha\beta$ lobes. The volume was computed using the 'Complex Hull Analysis' of the NeuroLucida explorer software after tracing the projections. Results are presented as

the mean \pm s.e.m. with $n = 8-11$. A two-sample, bilateral Student t -test was used to compare the two genotypes. *: $p < 0.05$. (M) *MiR-iab8-3p-SP* expression altered the length distribution of axonal segments. The definition of segment is provided in panel I. All segments were ranked according to their length and separated by genotype. *MiR-iab8-3p-SP* expression significantly reduced the fraction of small ($< 10 \mu\text{m}$) and long axonal segments ($> 100 \mu\text{m}$). Results are presented as the proportion of segments assigned to a distribution of size classes and relative to the total number of branches with $n = 56-70$. Chi-square tests were used to compare the distributions and proportions. *: $p < 0.05$, ***: $p < 0.001$. (N) *MiR-iab8-3p-SP* expression reduces the length of the primary segment of the $\alpha\beta$ MBn in the α lobe but not in the β lobe. The average length of the primary segment was computed for the α and β lobes independently and according to the genotype. Results are presented as the mean \pm s.e.m. with $n = 8-11$. A bilateral Mann-Whitney test was used to compare genotypes. *: $p < 0.05$.

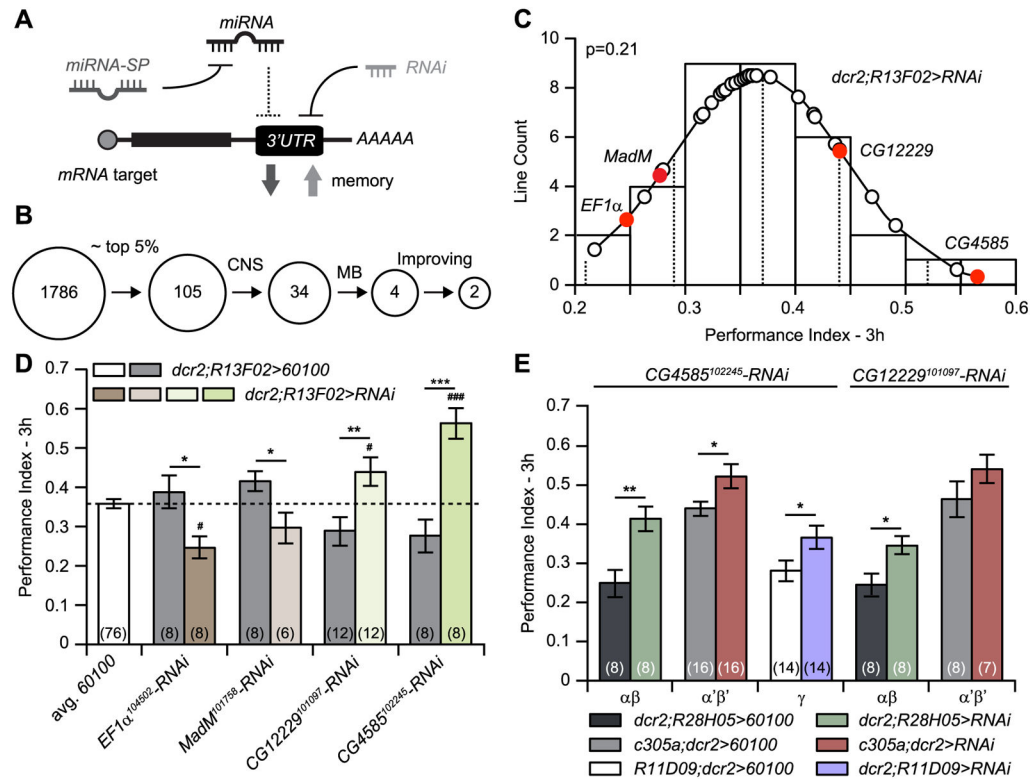


Fig. 6. *CG12229* and *CG4585* are potential mRNA targets for *miR-iab8-3p*

(A) Rational for the *miR-iab8-3p* mRNA-target screen. *miR-iab8-3p-SP* binds to *miR-iab8-3p*, preventing its interaction with the 3'UTR of mRNA targets and consequently increasing protein expression. Since *miR-iab8-3p-SP* expression impairs memory formation, we adopted the working hypothesis that inhibiting its targets using an RNAi approach would decrease protein expression and increase memory formation. (B) Schematic representation of the workflow used to identify the potential mRNA-targets. A computational approach (microRNA.org) identified 1786 potential targets. We selected the most likely targets picking lines with a mirSVR score ≥ -1.2 and obtained a primary list of 105 candidate genes. Three-hour memory was tested for flies expressing RNAi (*uas-RNAi*) to each of these candidates using the CNS-wide, *n-syb-gal4* driver. This primary screen reduced the list to 34 RNAi candidates that influenced memory formation. These RNAi's were subsequently expressed in the MBn using *R13F02-gal4*. We identified 4 genes in this manner that reproducibly modulated 3 h memory. Two of these increased memory formation. (C) PI distribution of *miR-iab8-3p* potential targets when inhibited in MBn. Three-hour memory was measured in RNAi-expressing flies using the *R13F02-gal4* driver. The average PI for 3 h memory was 0.37 ± 0.01 (mean \pm s.e.m.); the PIs followed a Normal distribution ($p = 0.21$). PIs obtained for genes ultimately identified are highlighted in red. PIs for genes represented with open circles were discarded since not confirmed by the following experimental validation. (D) Validation for the four potential targets. The memory performance for flies carrying each individual *uas-RNAi* and the *R13F02-gal4* driver was compared to *R13F02-gal4 > 60100* flies, host genotype used to produce the *uas-RNAi* lines. For each of the four RNAis, the PIs were significantly different from the paired *60100* control flies, and to the

average PI for *R13F02-gal4* > *60100* flies across the screen. Results are presented as the mean±s.e.m. with n = 6–12. Two-sample, bilateral Student *t*-tests were used to compare the two genotypes. ***: p<0.001, **: p<0.01, *: p<0.05. For comparisons to the average *60100* control (n = 76): ###: p<0.001, #: p<0.05. (E) *CG12229* and *CG4585* inhibition in subclasses of MBn increased memory expression. Each *uas-RNAi* transgene was combined with either *R11D09*, *c305a* or *R28H05-gal4* drivers to obtain expression in the, $\alpha'\beta'$ or $\alpha\beta$ neurons, respectively. *CG12229* inhibition in $\alpha\beta$ neurons increased memory expression relative to the *60100* control. *CG4585* inhibition in, $\alpha'\beta'$ and $\alpha\beta$ neurons increased memory expression. Results are presented as the mean±s.e.m. with n = 8–16. Two-sample, bilateral Student *t*-tests were used to compare the two genotypes. **: p<0.01, *: p<0.05.