**Summary**

Long-term memory and synaptic plasticity are thought to require the synthesis of new proteins at activated synapses. The CPEB family of RNA binding proteins, including *Drosophila* Orb2, has been implicated in this process. The precise mechanism by which these molecules regulate memory formation is however poorly understood. We used gene targeting and site-specific transgenesis to specifically modify the endogenous *orb2* gene in order to investigate its role in long-term memory formation. We show that the Orb2A and Orb2B isoforms, while both essential, have distinct functions in memory formation. These two isoforms have common glutamine-rich and RNA-binding domains, yet Orb2A uniquely requires the former and Orb2B the latter. We further show that Orb2A induces Orb2 complexes in a manner dependent upon both its glutamine-rich region and neuronal activity. We propose that Orb2B acts as a conventional CPEB to regulate transport and/or translation of specific mRNAs, whereas Orb2A acts in an unconventional manner to form stable Orb2 complexes that are essential for memory to persist.

**Introduction**

Most behaviors can be modified through the process of learning and memory, allowing the individual to adapt its innate behavioral repertoire to the specific contingencies of the local environment. Depending on the duration, intensity and salience of the learning experience, memories can be either short or long lasting. These behavioral modifications are thought to reflect anatomical and functional changes at specific synapses. Long-term synaptic plasticity requires new protein synthesis both at the soma and locally at the synapse (Sutton and Schuman, 2006). To ensure that local protein synthesis is restricted to the relevant synapses, either through the local capture or translation of mRNAs only in specific synapses, a "synaptic tag" has been postulated (Frey and Morris, 1997; Martin et al., 1997). Candidates for such a local protein synthesis regulator are members of the cytoplasmic polyadenylation element binding (CPEB) family. The founding members of this family mediate local protein synthesis in early development (Mendez and Richter, 2001), but some CPEB proteins are also thought to mediate protein synthesis in neurons (Alarcon et al., 2004; Atkins et al., 2004; Huang et al., 2002, 2003, 2006; Liu and Schwartz, 2003; Si et al., 2003a; Wells et al., 2001; Wu et al., 1998; Zearfoss et al., 2008; Miniaci et al., 2008; Si et al., 2003a).

CPEB proteins can be divided into two subfamilies. The CPEB-I subfamily includes the Xenopus CPEB1 and its *Drosophila* ortholog Orb1, both of which regulate mRNA translation during oogenesis (Mendez and Richter, 2001). CPEB1 and Orb1 bind cytoplasmic polyadenylation elements (CPEs) in the 3'UTR of dormant mRNAs, triggering their polyadenylation and translation (Fox et al., 1989; Hake et al., 1998). Members of the CPEB-II subfamily, including *Drosophila* Orb2, have been found to function in synaptic plasticity (mCPEB2–4) (Richter, 2001) or long-term memory formation (*Drosophila* Orb2) (Keleman et al., 2007; Majumdar et al., 2012). The mechanism by which these proteins might regulate protein synthesis is still unclear. Indeed, it has been suggested that neither polyadenylation nor CPEs are involved in translational regulation by CPEB-II proteins (Huang et al., 2006).

Almost all CPEBs exist in multiple variants generated by alternative mRNA splicing (Theis et al., 2003; Wang and Cooper, 2009). The *orb2* locus potentially generates six distinct proteins, only two of which contain the well-conserved RNA-binding domain (RBD) in the C terminus that is characteristic of CPEB proteins. These two isoforms, Orb2A and Orb2B, also share a glutamine-rich domain (Q domain) in the N terminus similar to that found in some but not all CPEB proteins in other species (Hafer et al., 2011; Si et al., 2003a). Orb2A and Orb2B differ only in their N termini, which do not contain any conserved domains. In *Drosophila*, long-term memory mediated by Orb2 is critically dependent on the Q domain (Keleman et al., 2007). The corresponding Q domain in *Aplysia* CPEB is thought to maintain long-term synaptic facilitation, possibly due to its putative prion-like properties (Heinrich and Lindquist, 2011; Si et al., 2010; Si et al., 2003b).

In order to further understand the cellular and molecular contributions of Orb2 to learning and memory in *Drosophila*, we have conducted detailed genetic and biochemical analyses of the endogenous Orb2 protein. To ensure that the modified proteins are expressed at the appropriate level and in the appropriate spatial and temporal pattern, we have made all
neuron locus either wild-type sequences (orb2 locus. This new allele, orb2 modified alleles were generated by insertion into the orb2 locus. An allele that allows rapid modification of the endogenous modifications directly in the orb2 locus. Our genetic and biochemical data support a model in which Orb2B acts as a conventional CPEB molecule by a mechanism dependent on its RBD. Orb2A appears to function in an unconventional mechanism that requires the Q domain but is independent of its RBD, possibly by seeding the formation of Orb2A:Orb2B complexes upon neuronal stimulation. We propose that these complexes mediate changes in mRNA translation at activated synapses, contributing to experience-dependent changes in synaptic function and animal behavior.

RESULTS

Generation and Validation of orb2attp

We generated by homologous recombination (Gong and Golic, 2003) an allele that allows rapid modification of the endogenous orb2 locus. This new allele, orb2attp, replaces most of the orb2 open reading frame (including sequences encoding the RBD and Q domains) with an attp recognition site. This attp site can be targeted by the site-specific recombinase phiC31 to insert any desired sequences directly into the orb2 locus (Bischof et al., 2007; Groth et al., 2004; Figure 1A).

To validate our approach we first reintroduced into the orb2attp locus either wild-type sequences (orb2GFP) or a modification designed to delete the Q domain (orb2QGFP). In both cases, as in most of the modifications reported here, the targeted orb2 allele additionally carried sequences encoding a C-terminal GFP tag. The structure of these modified orb2 loci were confirmed by Southern blots, PCR and transheterozygote (orb2QGFP/ orb2QGFP), and transheterozygote (orb2GFP/ orb2QGFP) were tested. p values are for H0: LI = 0, **p < 0.01, ***p < 0.001 (permutation test). See also Table S1 for LTM and Table S2 for STM.

Figure 1. Generation of the orb2attp Allele

(A) Most of the orb2 open reading frame was replaced with the acceptor site, attp, for the phi-C31 recombinase by ends-out homologous recombination. Dashed lines indicate deleted region. orb2 modified alleles were generated by insertion into the orb2attp allele of the relevant donor construct bearing the donor site, attB, and the modified genomic orb2 fragment, by phi-C31 mediated transgenesis.

(B) orb2 modified alleles were verified molecularly by Southern blots (SB), PCR, and RT-PCR. SB (left panel) was performed using probe a indicated in (A) and EcoRI. SpeI restriction digest. PCR (middle panel) amplification was performed using primers b and c indicated in (A). Obtained products were of the predicted size. RT-PCR (right panel) fragments were amplified using primers e and f indicated in (A). Product sizes were as predicted and were verified by DNA sequencing. RpS8 was used as an internal control. orb2GFP/ orb2QGFP heterozygotes were used for SB and PCR. Adult rare homozygous escapers orb2attp were used for RT-PCR.

(C) LIs (green bars) of males carrying the indicated orb2 alleles (on the left) with the corresponding Or2B protein organization (in the middle), tested in single-pair assays with mated females as trainers and testers for long-term memory (LTM) and short-term memory (STM). Control Canton-S flies, wild-type rescue allele (on the left) with the corresponding Or2B protein organization (in the middle), tested in single-pair assays with mated females as trainers and testers for long-term memory (LTM) and short-term memory (STM). Control Canton-S flies, wild-type rescue allele (orb2GFP), Q domain-deleted allele (orb2QGFP), and transheterozygote (orb2GFP/ orb2QGFP) were tested. p values are for H0: LI = 0, **p < 0.01, ***p < 0.001 (permutation test). See also Table S1 for LTM and Table S2 for STM.
C-terminal GFP tag does not impair Orb2 function. Accordingly, we also introduced the GFP tag for other modifications to the orb2 locus reported below, although for simplicity it is only indicated in allele or protein names when it is exploited in immunolabeling or biochemistry experiments.

Orb2A and Orb2B Have Distinct Expression Patterns and Developmental Roles
We used antibodies against the GFP tag on the endogenous Orb2 protein encoded by orb2\(^{AGFP}\) to determine its expression pattern and subcellular localization. At the level of light microscopy, Orb2 appeared to be broadly expressed throughout the nervous system of embryo, larvae, and adult, including the ventral nerve cord (VNC) and the brain. In the adult brain Orb2 appeared to be widely expressed throughout various regions including the lobes, calyces, and soma of the mushroom bodies (MB), a center for olfactory memory formation in insect brains (Heisenberg, 2003; Figure 2A).

Previous studies in other species have variously placed CPEBs at either pre- or postsynaptic sites. Mouse CPEB3, for example, was reported to be present in postsynaptic densities, whereas Aplysia CPEB was shown to localize in presynaptic compartments (Huang et al., 2003, 2006; Liu and Schwartz, 2003; Wu et al., 1998). To examine the subcellular localization of Drosophila Orb2, we examined the calyx (input) region of the MB (see Experimental Procedures for details). Using immunoelectron microscopy, we detected Orb2 both in the presynaptic compartment of the extrinsic MB neurons, characterized by the presence of presynaptic specializations such as electron-dense active zones, synaptic vesicles and occasionally T bars, and the postsynaptic compartment likely to be the termini of the Kenyon cells (KCs) in the calyx, characterized by the presence of close membrane alignments with the presumptive presynaptic region (Figure 2B). Furthermore, consistent with the reported role for Orb2 during development (Hafer et al., 2011; Keleman et al., 2007), we observed morphological defects in the brains of rare adult escapers of orb2\(^{AGFP}\) null mutants (Figure S1).

To distinguish the expression of the Orb2A and Orb2B isoforms, we next generated alleles designed to tag one isoform while eliminating the other (Figure 2C). By inserting a single nucleotide in the exon specific to orb2A, we disrupted the Orb2A reading frame while leaving the GFP-tagged Orb2B reading frame intact. In a second allele, we additionally removed a single nucleotide in the first common exon, thereby restoring the reading frame of Orb2A, including the GFP tag, while now disrupting that of Orb2B. We refer to these two alleles as orb2\(^{AGFP}\) and orb2\(^{ABGFP}\), respectively. Homozygous orb2\(^{AGFP}\) mutants were lethal, whereas orb2\(^{AGFP}\) flies were viable and healthy, indicating that Orb2B but not Orb2A has an essential role in development. To examine the respective distributions of the GFP-tagged Orb2B and Orb2A proteins we used homozygous orb2\(^{AGFP}\) and adult escaper orb2\(^{ABGFP}\) animals. The distribution of Orb2B was grossly similar to that observed for Orb2, but Orb2A was undetectable in our experiments (Figure 2D).

The Q Domain in Orb2A Is Both Required and Sufficient for Long-Term Memory
We used orb2 isof orm-specific alleles to test the function of Orb2A and Orb2B in long-term memory. Viable orb2\(^{AB}\) mutant males, expressing only the B isoform, were tested as homozygous. These mutants had a normal short-term memory (Table S5D) and a strong detriment in long-term memory in comparison to the wild-type flies (3, orb2\(^{AB}\), LI = 12.69; 1, orb2\(^{AB}\), LI = 30.31), almost as severe as mutants lacking the Q domain in both isoforms (2, orb2\(^{AQ}\), LI = 2.15). suggesting that Orb2A function is critically required for long-term memory (Figure 3; Table S4). However, these mutant flies were able to form residual but statistically significant memory likely to be mediated by Orb2B.

To assess the role of the Q domain in Orb2 isoforms, we generated a specific deletion of this domain by reinserting into orb2\(^{amp}\) a genomic fragment in which disruption of either Orb2A or Orb2B was combined with the deletion of the Q domain. orb2\(^{AQ\Delta}\) males, expressing only Orb2B lacking its Q domain, had a normal short-term memory (Table S5D) but, like orb2\(^{AQ}\) mutants, almost no long-term memory (4, orb2\(^{AQ\Delta}\), LI = 5.16; 2, orb2\(^{AQ}\), LI = 2.15) (Figure 3; Table S4), suggesting that the residual memory of the orb2\(^{AQ\Delta}\) mutants might be mediated by the Q domain of Orb2B.

Since the orb2\(^{AQ\Delta}\) mutation was lethal when homozygous, we tested this allele in combination with the viable orb2\(^{AB}\) allele. These flies, which lack the Q domain specifically in Orb2A, had a normal short-term memory (Table S5D) but no long-term memory (5, orb2\(^{AQ\Delta}/\)orb2\(^{AB}\), LI = 2.86) (Figure 3; Table S4). This lack of memory shows that the Q domain in Orb2A is essential, and that of Orb2B insufficient, for long-term memory.

To test for the sufficiency of the Q domain in Orb2A, we tested the memory of the transheterozygotes in which the Q domain is present only in Orb2A. The learning index of these mutants was indistinguishable from control flies in which both isoforms are intact (6, orb2\(^{AB}/\)orb2\(^{AQ\Delta}\), LI = 16.97; 7, orb2\(^{AB}/\)orb2\(^{AQ\Delta}\), LI = 20.83) (Figure 3; Table S4). These results indicate that Orb2A has a specific role in long-term memory that requires the Q domain, which in Orb2B is both dispensable and insufficient.

The RBD Is Essential for Function of Orb2B but Not of Orb2A in Long-Term Memory
To assess the role of the RBD in long-term memory, as a first step we chose to replace the Orb2 RBD with the RBDs of other protein names when it is exploited in immuno
Figure 2. Orb2 Is Enriched in the Nervous System and Localizes to Pre- and Postsynaptic Compartments

(A) Confocal projections of the Drosophila embryo, larvae, and adult brain of orb2\(^{GFP}\) animals (upper panels) stained with antibody to GFP (green) and counterstained with the general neuropil marker, antibody to either FasII or nc82 (red). Scale bar is 50 \(\mu\)m. Confocal projections of the adult Drosophila ventral nerve cord (VNC), mushroom body calyx (ca), lobes (\(\alpha, \gamma\)), and cell bodies of the Kenyon cells (KC) (lower panels) of orb2\(^{GFP}\) animals stained with antibody to GFP.
orb2 specific targets involved in long-term memory formation. When individually expressed, Orb2A and Orb2B have distinct localizations. While both isoforms are localized to the cytoplasm, Orb2A has a granular appearance which to test for aggregation and the potential role of the Orb2 Q domain in this process. When individually expressed, Orb2A and Orb2B form heteromeric complexes mediated by the Q domain. One possible explanation for this interallelic complementation between orb2A and orb2B alleles could be that the proteins they encode form a functional complex. We examined this possibility by light microscopy and biochemistry. Due to the small size of Drosophila neurons, we used expression studies in the Drosophila S2 cell line. S2 cells do not express Orb2 (our unpublished deep seq. data); therefore, we had a clean background in which to test for aggregation and the potential role of the Orb2 Q domain in this process. When individually expressed, Orb2A and Orb2B, have distinct localizations. While both isoforms are localized to the cytoplasm, Orb2A has a granular appearance whereas Orb2B is diffuse. Interestingly, loss of the Q domain in
Orb2A (Orb2AΔQGFP) led to a loss of the granular appearance whereas deletion of this domain in Orb2B (Orb2BΔQGFP) did not cause any detectable change (Figure 5A). This observation was extended by IP experiments. In immunoprecipitates from S2 cells transfected with both Orb2AGFP and Orb2BGFP, we observed large Orb2 complexes ranging between 100–400 kDa. Deletion of the Q domain in Orb2A eliminated higher molecular weight multimers above 250 kDa, and deletion of the Q domain both in A and B isoforms eliminated them completely, leaving only the Orb2 monomer band at ~100 kDa (Figure 5B).

These experiments suggest that Orb2 can form multimers in S2 cells that are dependent on the Q domain of both isoforms.

To examine multimerization of both isoforms in vivo, we analyzed immunoprecipitates from fly brains. In orb2AΔGFP brains, we found Orb2 present both in monomers and oligomers (~100 and 200 kDa). In immunoprecipitates from orb2BΔAGFP brains we found Orb2B mostly in a lower molecular weight band of ~100 kDa. Since deletion of Orb2B is lethal, to analyze multimerization properties of Orb2A we immunoprecipitated Orb2A from the brains of heterozygous animals (orb2BΔBGFP/+).

We observed Orb2A almost exclusively in a high molecular weight band of ~200 kDa. Consistent with Orb2B being expressed at higher levels than Orb2A, Orb2A could not be detected from the same amount of input material as for Orb2B.
In summary, Orb2A preferentially exists in multimeric complexes, whereas Orb2B has a lower propensity to aggregate but may be induced to aggregate in the presence of Orb2A. In order to test whether Orb2A and Orb2B are present in the same complex, we turned to mass spectrometry (MS), which can readily distinguish between the two isoforms. As we were unable to detect the 9 amino acids specific to Orb2A, we looked for Orb2B-specific peptides when Orb2A was immunoprecipitated. The presence of Orb2B in such immunoprecipitates would indicate that Orb2A is able to pull down Orb2B, and that these two proteins are present in one complex. We precipitated Orb2A from \textit{orb2}D\text{BGFP}/+ and \textit{orb2}D\text{BQGFP}/+ brains (Figure 5D). These results show that both Orb2 isoforms are present in the Drosophila brain in one complex, provided Orb2A has an intact Q domain.

Neuronal Activity Induces Orb2 Multimers through the Q Domain

Both dopamine and octopamine have been shown to mediate memory formation in olfactory and courtship learning paradigms (Keleman et al., 2012; Schwaezrel et al., 2003; Tempel et al., 1984). We fed adult flies carrying wild-type \textit{orb2}\text{AGFP} with either dopamine or tyramine (a neurotransmitter and precursor of octopamine) to stimulate broadly neuromodulatory pathways in the brain, and monitored Orb2 multimers at specific time points postfeeding. Orb2 in brain extracts from flies fed with either tyramine or dopamine exists both as monomers (\textasciitilde 100 kDa) and oligomers (\textasciitilde 200 kDa). The oligomer band appears between 4–6 hr postfeeding and lasts for at least 20 more hours (Figures 6A and 6B). This result parallels our previous finding that memory in \textit{orb2}\text{DQ} mutants does not last beyond 6 hr (Keleman et al., 2007). In control animals fed with sucrose only (point 0), the oligomer band was absent. The amount of extract we used for these experiments should only monitor the Orb2B isoform. Therefore, we interpret our results as demonstrating that Orb2B the mono- to oligomeric state upon neuronal stimulation.

To evaluate the impact of the Q domain deletion on oligomerization we fed \textit{orb2}\text{DAGFP} mutant flies with tyramine and followed Orb2 protein as above. In \textit{orb2}\text{DQGFP} mutant brains, although Orb2 protein was expressed at the same level as in the wild-type \textit{orb2}\text{AGFP} animals, only an Orb2 monomer was observed (Figure 6A), implying an acute role for the Q domain in Orb2 oligomerization. This result parallels a complete lack of long-term memory in \textit{orb2}\text{DQ} mutant flies.

To investigate if Orb2A regulates oligomerization of Orb2B, we fed animals lacking the Orb2A isoform with tyramine. As above, we did not detect Orb2B oligomers, suggesting that Orb2A is crucial for oligomerization (Figure 6A). Finally, to test the role of Orb2A’s Q domain in Orb2 oligomer formation, we analyzed transheterozygous animals in which the Q domain present only in Orb2A and RBD only in Orb2B, able to form long-term memory (3, \textit{orb2}\text{DAGFP}/\textit{orb2}\text{RD}B\text{DAGFP}, LI = 0.68; 1,
As predicted, in brain extracts from these animals, Orb2 multimers were detected as in the wild-type brains. In contrast, in brain extracts of animals in which the Q domain was lacking specifically in Orb2A and present only in Orb2B, which are unable to form long-term memory, Orb2 oligomers were not detected (Table S4; Figure 6C). We conclude that Orb2 oligomers are induced by neuronal activity in Orb2A-dependent manner. The Q domain of Orb2A is both essential and sufficient, whereas that of Orb2B is dispensable and insufficient, for Orb2 oligomers formation. These results suggest that Orb2 complexes are essential for memory persistence.

**DISCUSSION**

Local translation of mRNAs in both pre- and postsynaptic compartments is thought to be important for the synaptic modifications that underlie long-lasting memories (Frey and Morris, 1997; Kang and Schuman, 1996; Martin et al., 1997). The CPEB family of proteins regulate local translation (Alarcon et al., 2004; Huang et al., 2006; Si et al., 2003a; Wells et al., 2001; Wu et al., 1998; Zearfoss et al., 2008), and the Drosophila CPEB protein Orb2 is acutely required for long-term memory (Keleman et al., 2007; Majumdar et al., 2012). However, the detailed molecular mechanism of CPEB function in synaptic plasticity and memory formation remains elusive.

We have shown here that the two Orb2 isoforms, Orb2A and Orb2B, both contribute to long-term memory formation, albeit by distinct mechanisms. The two isoforms share the same RNA-binding and Q domains, yet each uniquely requires only one of these domains for its function in long-term memory formation. Specifically, the Q domain is essential in Orb2A but not Orb2B, whereas the RNA-binding domain is required in Orb2B but not Orb2A. Moreover, we found that Orb2A lacking its RNA-binding domain is able to fully complement Orb2B lacking its Q domain. Such interallelic complementation often...
refers to the formation of the heteromeric complexes between the
encoded proteins (Garen and Garen, 1963; Zhang et al., 2006),
and indeed we observed that Orb2A and Orb2B are present in
the same protein complexes in vitro and in vivo, and that forma-
tion of these heteromeric Orb2A:Orb2B complexes acutely
depends on Orb2A and its Q domain. Moreover, these
complexes are induced within 6 hr after feeding with biogenic
amines (thought to provide learning signals relevant for memory
formation; Schwarzeil et al., 2003), corresponding to the time
course of memory decay in Orb2ΔΔ mutants (Keleman et al.,
2007). We therefore propose that Orb2A:Orb2B heteromeric
complexes are induced at specific synapses by the relevant
learning signals and required for memory persistence beyond
6 hr.

Dopamine is thought to provide a reinforcement signal in
Drosophila courtship learning (Tempel et al., 1984). For short-
term memory this dopamine signal is provided by neurons that
innervate the gamma lobe of the mushroom body (Keleman et al.,
2012), and for long-term memory Orb2 is required in
intrinsic gamma lobe neurons (Kenyon cells; Keleman et al.,
2007). Gamma lobe synapses are thus a likely site of Orb2
coregulation and the structural and functional modifications
that underlie courtship learning in Drosophila. Orb2 also
functions in long-term memory in an appetitive learning para-
digm (Majumdar et al., 2012), which likely maps to a distinct
class of mushroom body neuron (Waddell, 2010). Indeed,
specific long-term memories may be stored at various sites in
the fly brain, extending even beyond the mushroom body
(Chen et al., 2012; Davis, 2011). The broad distribution of Orb2
throughout the nervous system suggests that it may contribute
generally to long-term synaptic plasticity and memory formation,
regardless of where these memories are stored.

Why might Orb2A have such a critical role in Orb2 complex
formation and long-term memory, when most of its residues are
shared with the evidently more abundant Orb2B, including the
Q and RNA-binding domains? The efficacy of complex
formation of proteins containing Q domains is thought to be
determined by the length of the preceding N-terminal sequences
(Shorter and Lindquist, 2004). In this regard it is interesting to
note that Orb2A has an N-terminal extension of 9 amino acids,
comparable to the 162 N-terminal residues of Orb2B. Additionally,
a single point mutation in the unique Orb2A N-terminal extension
decreases Orb2 multimer formation in the Drosophila brain and
impaired long-term memory retention beyond 48 hr (Majumdar
et al., 2012). Thus, both the size and sequence of Orb2A’s unique
N-terminal extension might endow it with a greater propensity to
aggregate than Orb2B, and thereby nucleate heteromeric Orb2
complexes through the Q domain of Orb2A.

It has been suggested that the activation of Orb2 and other
CPEB proteins occurs via the prion-like properties of their Q
domains (Heinrich and Lindquist, 2011; Krishnan and Lindquist,
2005; Majumdar et al., 2012; Si et al., 2003a, 2010). Such Q
domains occur however in a wide range of proteins with diverse
biochemical functions, in which they are generally thought to
mediate homo- and heterotypic interactions (Michelitsch and
Weissman, 2000). In some of these proteins, for example, the
Q domains serve as polar zippers in the assembly of large multi-
meric complexes (Perutz et al., 1994). Whatever the means by
which the Q domain of Orb2 contributes to complex formation,
our data suggest that this is restricted to Orb2A, as Orb2B
does not require its Q domain to interact with Orb2A and function
in long-term memory formation. This mechanism is likely to be
conserved among CPEB proteins, as the Q domain of Orb2A
can be replaced with the analogous domain from CPEBs of
Aplysia and mouse, but not with the prion domain of ScUre2
(Figure S4).

Our data support and extend a model (Majumdar et al., 2012)
in which the two Orb2 isoforms form heteromeric complexes that
are essential for long-term memory formation. We further
propose that, upon neuronal stimulation, Orb2A, which may be
present in more limiting amounts, restricted locations, or under
specific circumstances, provides the spatial and temporal spec-
ificity for heteromeric complex formation and, we infer, synaptic
plasticity. Orb2B, in contrast, appears to be more broadly and
highly expressed and may mediate a more general function of
Orb2 in development (Cziko et al., 2009; Hafer et al., 2011;
Richter, 2007; Shieh and Bonini, 2011). Orb2 has been reported to
be present in the messenger RNPs, as we have also observed
here specifically for Orb2B (Figure S3), and is thought to control
mRNA transport and translational repression (Cziko et al., 2009;
Mendrez and Richter, 2001). During learning, Orb2A might
interact with Orb2B-containing RNPs at the relevant synapses,
releasing the associated mRNAs from translational repression
or possibly even converting Orb2B from a translation repressor
to an activator.

CPEB molecules are conserved across a wide range of
species and most of them exist in multiple isoforms generated
through alternative splicing, often varying only in their N terminus
(Theis et al., 2003; Wang and Cooper, 2009). This is the case
for mCPEB3, for example, the Q domain of which is able to
substitute for the Q domain of Orb2A both biochemically and
behaviorally (Figure S4). It is tempting to speculate that the
model we propose here is not unique for Drosophila Orb2 but
might also extend to other members of the CPEB family. More-
over, because Orb2A functions in long-term memory without
its RNA-binding domain, it is possible that proteins lacking an
RNA-binding domain, and hence not even recognized as canonical
CPEB molecules, might function in a fashion analogous to
Orb2A in Drosophila and other species.



EXPERIMENTAL PROCEDURES

Generation and Verification of the orb2ΔΔP and Subsequent Modified
orb2 Alleles

orb2ΔΔP was generated by ends-out homologous recombination (Gong and
Golic, 2003), using homology arms of 3.3 kb and 3.7 kb flanking the A isoform
specific and common exons of Orb2 (CG5735). In the initial recombinant,
this region was replaced with an attP followed by a white marker flanked by
mFRT11 recognition sites for the mFLPs recombinase (Hadjieconomou
et al., 2011). Removal of the white marker using hs-mFLPs generated the final
orb2ΔΔP allele, in which the A and common exons were replaced by an attP site
(Groth et al., 2004) and a single mFRT11 site. The targeted allele was verified
by genomic PCR and DNA sequencing across the entire homology region.
Southern blot and RT-PCR confirmed the intended modifications (Figure 1B).
Modified orb2 alleles were generated by cloning of the genomic fragments
with the relevant modification first into vector containing donor attB site for subse-
quent reinsertion by phiC31 (Bischof et al., 2007) mediated transgenesis into
orb2ΔΔP allele (details in the Supplemental Experimental Procedures). The

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intended modifications were verified by PCR amplification and DNA sequencing across the modified region.

**Generation of the Fly Strains**

To generate fly strains carrying modified orb2 alleles, donor constructs containing genomic fragments with the specific modification were injected into the embryos from a cross between orb2<sup>attP</sup> flies and phiC31 integrase-expressing flies, ZH11 (Bisch et al., 2007). DNA injection resulted in a site directed integration of the attB containing constructs into orb2<sup>attP</sup>, phiSFLP<sub>j</sub> (Hadjeconomou et al., 2011) was used to excise the w<sup>+</sup> marker.

**Southern Blot**

A probe (a) as indicated in Figure 1A, was generated by PCR using the primer SB1 and SB2 (Supplemental Experimental Procedures). Fifteen micromers genomic DNA was digested using EcoRl/SpeI. DNA was run on a 0.5% agarose gel at 60V at 4° C over night. The gel was blotted in 20x SSC over night. After cross-linking, the membrane was incubated in hybridization solution (ULTR Alyb Ultrasensitive Hybridization Buffer, Ambion, AM8670) before incubation with the labeled probe (Prime-It Random Primer Labeling Kit, Stratagene, 300385) for 16 hr.

**RT-PCR**

Total RNA was extracted using Trizol and reverse transcribed using primers e and f as indicated in Figure 1A (Supplemental Experimental Procedures). RpS8, was amplified with primers HH142 and HH143 (Supplemental Experimental Procedures) and used as an internal control.

**Behavioral Tests**

All orb2 alleles were backcrossed for five generations into a Canton-S background before being used in behavioral assays. Flies were raised on semi

immunohistochemistry for adult brains, VNC, and larval CNS was performed as described (Yu et al., 2010). Fly brain and VNC were dissected (between 5 to 8 days after eclosion) in PBS and fixed using 4% paraformaldehyde in PBST (PBS with 0.3% Triton X-100) for 20 min at 24° C. After washing in PBST, the tissue was blocked in 5% normal goat serum in PBST for at least 2 hr. The primary antibody and secondary antibody were incubated for 48 hr at 4° C. The brains were washed with PBST 3 x 10 min and then overnight at 4° C between the primary and secondary antibody incubations. After the secondary antibody incubation, samples were washed 3 x 10 min and overnight at 4° C before mounting in Vectashield (VectorLabs).

Antibodies used: rabbit polyclonal anti-GFP (1:5,000, Torri Pines); mouse nc82 (1:50, Hybridioma Bank); mouse anti-DAC2-3 (1:200, Hybridioma Bank); rabbit anti-eIF4e (Nakamura et al., 2004) (1:5,000); rabbit anti-Trailer-hitch (Tra) (Boag et al., 2005) (1:5,000); secondary Alexa-488, -568 antibodies (1:1,000, Invitrogen).

**Embryos**

Immunochemistry for embryos was as described (Patel et al., 1987). Embryos were collected and incubated in 50% bleach for 3 min and rinsed into a sieve using tap water. Next, they were transferred to the eppendorf tubes containing 500 μl heptane and 450 μl PBS. For fixation 50 μ1 formaldehyde was added for 20 min at RT. Lower phase was removed first, and the heptane was replaced by fresh heptane and ice-cold methanol. Then embryos were agitated strongly for 1 min to remove their vitelline membrane. After that, 3 x 5 min washes in methanol were performed followed by three washes in PBST to remove residual methanol. Next, the embryos were blocked for 1 hr in 5% normal goat serum prior to antibody incubation. Antibody incubation was done either for 1hr at RT or O/N at 4° C. Antibodies used: rabbit polyclonal anti-GFP (1:5,000, Torri Pines), mouse anti-Fasll (1:50, Hybridioma Bank, 104), secondary Alexa-488, -568 antibodies (1:1,000, Invitrogen).

**Confocal Microscopy**

Tissues were scanned using a Zeiss LSM 510 with a Zeiss Multi Immersion Plan Neofluar 25 x/0.8 objective (as described; Yu et al., 2010). On average 8 brains or 5 VNCs were imaged for each genotype. Scanning parameters were set to image the central brain or the entire ventral nerve cord within 30 min. Images were taken at 512 x 512 pixels and 180 slices at 1.2 μm interval. A macro plug-in was used to automate the scanning process. Images were processed in ImageJ (NIH) to obtain maximum intensity Z projections.

**Immuno-EM on Adult Brains**

The heads of 3 days old orb2<sup>attP</sup> and Canton-S male flies were fixed in 4% paraformaldehyde, 0.1% glutaraldehyde, 0.07 M phosphate buffer (pH 7.3) for 3 hr at 4° C. Frontal vibratome sections (80 μm) were collected from each head from the anterior to the posterior region and the last two sections were
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processed for immuno-EM. Fifteen heads were used per genotype. Sections were incubated with rabbit anti-GFP (Molecular Probes, dilution 1:200) for 44 hr at 4°C and avidin-biotinylated-peroxidase complexes (Vectastain Elite Kit Vector, Burlingame, CA) were formed as described (Yasuyama et al., 2002). Sections were post fixed in 0.1% osmium tetroxide in 0.1M cacodylate buffer (pH 7.4) in 0.08% potassium ferrocyanide for 1 hr at 4°C, dehydrated in ethanol and embedded in LX112 resin (Ladd Research, Williston, VT) following standard procedures. Semithin sections were collected (0.8 µm thickness) using a Leica EM UC6 microtome (Leica, Vienna, Austria), stained with toluidine blue and examined under the light microscope to determine the location in the brain. Ultrathin serial sections (110 nm) were collected on EM specimen grids when the region of the mushroom body calyx were visible, post-stained with uranyl acetate and lead citrate and imaged with a FEI Tecnai 12 transmission electron microscope, operated at 120 kV and equipped with an Eagle 4kx4k camera (FEI, Eindhoven, The Netherlands). At least three brains were analyzed per genotype.

Mass Spectrometry

Tryptic Digestion

For reduction 2 µl DTT of 1mg/ml (dissolved in 100 mM ammonium bicarbonate) stock solution were added to the samples. Reduction was performed for 30 min at 56°C. Alkylation was performed with 2 µl of 40 mM (in 100 mM ammonium bicarbonate) stock solution for 30 min at room temperature in the dark. Afterward the samples were digested with 400 ng of trypsin (Gold, ammonium bicarbonate) stock solution for 30 min at room temperature in the dark. The digestion was stopped with 10 µl of 10% TFA.

NanoLC-MS

The nano HPLC system used in all experiments was an UltiMate 3000 Dual Gradient HPLC system (Dionex, Amsterdam, The Netherlands), equipped with a Proxeon nanospray source (Proxeon, Odense, Denmark), coupled to an LTQ Velos Orbitrap mass spectrometer (Thermo Fisher Scientific). Instrument was operated in data-dependent mode using a full scan in the ICR cell followed by MS/MS scans of the twelve most abundant ions in the linear ion trap. MS/MS spectra were acquired in the multistage activation mode. Precursor ions selected for fragmentation were put on a dynamic exclusion list for 90 s. Monoisotopic precursor selection was enabled.

Analysis of MS Data

For peptide identification, all MS/MS spectra were searched using Mascot 2.2.04 (Matrix Science, London, UK) against the flybase database (49,832 sequences; 31,566,328 residues). The following search parameters were used: beta-methylthiolation on cysteine was set as a fixed modification; oxidation on methionine was set as variable modification. Monoisotopic masses were searched within unrestricted protein masses for tryptic peptides. The peptide mass tolerance was set to ± 0.5 Da. The maximal number of missed cleavages was set to ± 0.5 Da. The maximal number of missed cleavages was set to ± 0.5 Da. For transfection, confluent S2 cell cultures were split 1:3 to a fresh medium and grown overnight. Cultures were washed with 1 x PBS and resuspended to 2 million cells/ml in fresh medium before being transfected with 800 ng appropriate plasmid using Effectene Transfection Reagent (QIAGEN, 301425) according to manufacturer’s protocol. Transfected cells were subsequently grown for 72 hr and then changed to the selection medium containing hygromycin B by spinning 5 min at 1,000 g and resuspending the cell pellet in a fresh culture medium containing the antibiotic. For expression of Orb2 protein S2 cells were induced O/N with 0.25 mM CuSO4 (final concentration).

SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, six tables, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2012.08.028.

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