Regulation of *Drosophila* Brain Wiring by Neuropil Interactions via a Slit-Robo-RPTP Signaling Complex

**Highlights**

- In the *Drosophila* brain, mushroom bodies are a source of the Slit guidance cue

- Slit regulates axon growth in the vicinity of mushroom bodies via Robo receptors

- The phosphatase RPTP69D regulates Robo signaling in the brain

- RPTP69D regulates Robo3 membrane presentation independent of its enzymatic activity

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**In Brief**

Slit/Robo signaling is a conserved pathway regulating axon guidance. In the *Drosophila* ventral nerve cord, Robo is regulated by the protein Commissureless, but this mechanism is not conserved. Oliva, Soldano et al. show that in the *Drosophila* brain, the phosphatase RPTP69D regulates Robo membrane levels, independently of its enzymatic activity.
Regulation of Drosophila Brain Wiring by Neuropol Interactions via a Slit-Robo-RPTP Signaling Complex

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SUMMARY

The axonal wiring molecule Slit and its Round-About (Robo) receptors are conserved regulators of nerve cord patterning. Robo receptors also contribute to wiring brain circuits. Whether molecular mechanisms regulating these signals are modified to fit more complex brain wiring processes is unclear. We investigated the role of Slit and Robo receptors in wiring Drosophila higher-order brain circuits and identified differences in the cellular and molecular mechanisms of Robo/Slit function. First, we find that signaling by Robo receptors in the brain is regulated by the Receptor Protein Tyrosine Phosphatase RPTP69d. RPTP69d increases membrane availability of Robo3 without affecting its phosphorylation state. Second, we detect no midline localization of Slit during brain development. Instead, Slit is enriched in the mushroom body, a neuronal structure covering large areas of the brain. Thus, a divergent molecular mechanism regulates neuronal circuit wiring in the Drosophila brain, partly in response to signals from the mushroom body.

INTRODUCTION

During nervous system development, proper axon guidance is achieved through the interaction between neuronal cell surface receptors and their chemo-attractive or repulsive ligands present in the environment (Chilton, 2006; Dickson and Gilestro, 2006; Lowery and Van Vactor, 2009). The Slit/Robo signaling pathway (Dickson and Gilestro, 2006) plays essential functions during axon pathfinding in many neural populations, and special attention has been given to its role in commissural axon development (Dickson and Gilestro, 2006). Upon Slit binding to its receptor Robo, axons are repelled from the Slit source in most systems (Dickson and Gilestro, 2006). The Drosophila embryonic ventral nerve cord (VNC) has served as a powerful model system for the study of axon guidance by the Slit/Robo pathway (Dickson and Gilestro, 2006) whereby glial cells along the midline express Slit, which acts as a repulsive cue to guide neuronal axons toward or away from the midline in function of their repertoire of Robo receptors. This is similar to mammalian spinal cord where a specialized midline structure called the floor plate acts as a major source of guidance cues (Chedotal, 2011). In Drosophila there are three Robo receptors (Robo1, Robo2, and Robo3) and one Slit ligand. Robo1 and Robo2 are involved in commissure formation in the embryo while Robo 2 and Robo 3 regulate the formation of ipsilateral pathways (Rajagopalan et al., 2000). In contrast to the VNC, it is less clear how axon guidance is organized in higher-order brain centers. Roles for Slit and/or Robo receptors have been established in guiding peripheral axons to the brain (Jhaveri et al., 2004; Pappu et al., 2011) and Robo loss-of-function mutants, or pan-neuronal downregulation, causes broad defects (Nicolas and Preat, 2005; Tayler et al., 2004), suggesting a potentially important role for this ligand–receptor pair in adult brain connectivity. In the Drosophila VNC a major mechanism of regulating Robo activity is via the protein Commissureless (Comm) (Keleman et al., 2002, 2005; Tear et al., 1996), which binds Robo receptors and negatively regulates their activity. Comm has not been identified in other taxa, however, suggesting that this mechanism is unlikely to be conserved.

Receptor protein tyrosine phosphatases (RPTP) belong to a family of transmembrane proteins that are characterized by three extracellular immunoglobulin (Ig) domains and usually four to eight FNIII repeats, resembling adhesion molecules such as N-CAM (Chagnon et al., 2004; Siu et al., 2007), and two tandem
Figure 1. Slit Expression Pattern and Phenotypes during Brain Development

(A) Slit RNA expression in the larval brain using in situ hybridization. KC, Kenyon cells; OL, optic lobe; CB, central brain.
(B) slit enhancer (GMR31A10-Gal4) driving GFP shows expression in MB. Cx, mushroom body calyx.
(C) Slit expression in L1 stage; larval CNS was stained with Slit (gray). Note that Slit is not expressed in any structure in the brain at this stage while strong expression is seen in the ventral midline glia (MG). VNC, ventral nerve cord; OL, optic lobe; CB, central brain.
(D–H) Slit expression from L2 until adult stage. Starting at L2 Slit is enriched in the mushroom bodies (MB). Note that the expression in the VNC disappears at 75 hr after puparium formation but Slit continues to be expressed until adult stage in MB.
(I) The wild-type pattern of the neuropil in the adult brain revealed with anti-N-cadherin. MBVL, mushroom body vertical lobe; SMP, superior medial protocerebrum; MBML, mushroom body medial lobe; AL, antennal lobe.
(J) Pattern of the neuropil in the adult brain revealed with anti-N-cadherin in slit mutants presents widespread and strong defects in neuropil architecture (arrowheads).
(K–N) Mushroom body and ellipsoid body architectures revealed with anti-Fas2 in wild-type animals (K, M) and slit mutants (L, N). Arrowheads indicate specific brain structures (mushroom body α-lobes in K and L, ellipsoid body in M and N).
(O–R) Pattern of Robo2- and Robo3-expressing axons in L3 stage in normal and slit mutant animals. Note that in wild-type brains Robo2- and Robo3-positive commissures are present (O and Q, see arrowheads). In slit mutant brains the integrity of these commissures is disturbed (P and R, see arrowheads).

(legend continued on next page)
intracellular catalytic domains (D1 and D2) with putative phosphatase activity. RPTPs have been shown to play an important role in nervous system development. In both flies and vertebrates, RPTPs play a role in guidance of motor axons (Stepanek et al., 2005). In the fly embryonic nervous system, RPTP69d and RPTP10d have been shown to regulate commissure development and to genetically interact with the Slit/Robo pathway (Sun et al., 2000). Based on the presence of phosphatase domains in these proteins, it has been speculated that RPTP69d and RPTP10d activate Robo by dephosphorylating it. However, this assumption has not been experimentally tested in any model.

Here we show that the higher-order Drosophila brain region, known as the protocerebrum, does not contain midline sources for the major axon growth and guidance cue Slit. Instead, the mushroom body (MB) is the major source of Slit in the developing protocerebrum. The MB is a large, highly conserved, insect neuropil composed of the axons and dendrites of approximately 2,500 neurons called the Kenyon cells. The MB is required for associative learning and memory, as well as a host of innate and learned behaviors (Heisenberg et al., 1985; Krashes et al., 2007; Pitman et al., 2006). We show that Slit expression within the MB is essential for the correct patterning of neighboring higher-order neural circuits. Specifically, the interaction between Robo receptors and RPTP69d is necessary and sufficient for repulsive axonal responses to Slit from the MB. While RPTP69d co-expression enhances the effect of both Robo receptors, it has no repulsive or Slit binding activity on its own. Surprisingly, the RPTP69d phosphatase domain is dispensable for both Robo receptor binding and axon repulsion. Instead, we find that RPTP69d enhances the cell surface presentation of Robo receptors. We propose that the MB acts as a spatially distributed neuronal source of Slit for Drosophila brain connectivity, which could contribute to the higher level of complexity observed in the brain compared with the VNC. Furthermore, we identify a different Slit-Robo-RPTP signaling mechanism acting in the brain.

RESULTS

Slit Is Expressed in the Mushroom Body and Is Required for Central Brain Connectivity

To explore how central brain connectivity is organized, we examined Slit and Robo expression during brain development. Visualization of Slit RNA by in situ hybridization reveals its known pattern in the VNC and optic lobe (Figure 1A). Moreover, signal is observed in two bilateral clusters consistent with the localization of MB Kenyon cells (Figure 1A). Furthermore, an enhancer fragment from the Slit locus reports strong GFP expression in the MB (Figure 1B). Next, we explored the expression pattern of Slit protein. We find that starting at the second larval instar and throughout pupal brain development, Slit protein is strongly expressed in the MB (Figures 1C–1H). Slit is expressed in both the axons and dendrites of MB neurons themselves and not in the glial scaffold that surrounds them (Figure S1). Protein localization in MB is confirmed by MB-specific knockdown of Slit showing a decreased signal in MB, in contrast to glial-specific knockdown (Figure S1). The MB is a large and spatially distributed neuropil structure, whose axons and dendrites span a significant expanse of the developing brain in all three axes (Figure 1S). All three Robo receptors are expressed in the brain in many different axonal tracts, but are specifically absent from the MB at the third instar larval stage (Figures S1L–S1W). While slit null mutant animals die at embryonic or early larval stages, we were able to examine the neuropil structure of viable slit mutants bearing a combination (slitfl/slitfas) of a null allele (slitfl) and a hypomorphic allele (slitfas) (Taylor et al., 2004) exhibiting a strong reduction of Slit in the larval and pupal nervous system (Dasceno et al., 2015). We found major defects in neuropil organization in several central brain areas, including MB lobes, the central complex, antennal lobes, and Robo2/Robo3-expressing axons in the developing brain (Figures 1I–1R). In summary, reduction of Slit activity in the brain causes widespread disturbances in brain neuropil architecture. Whereas in the VNC midline glia acts as a point source of a Slit gradient, in the protocerebrum the axons and dendrites of bilateral MBs are a neuronal source of broadly distributed Slit during brain development (Figure 1S).

The Mushroom Body Regulates Axon Growth and Guidance via Slit/Robo Signaling

To uncover the specific mechanisms of Slit function in the MB, we chose two neuronal populations which express or lack the Robo receptors, respectively. The circadian clock neurons, called small lateral neurons (sLNv) (Helfrich-Forster et al., 2007), express all three Robo receptors and localize them to MB, whereas the dorsal cluster neurons (DCNs), which are higher-order contralateral projecting neurons with axons that innervate the optic lobes (Langen et al., 2013; Zschatsch et al., 2014) do not express Robo1–3 (Figures S2F–S2H). The four to five sLNv axons project dorsally and then turn medially close to the MB dendritic tree known as the calyx (Figures 2A and 2B). In slit mutants, the medial projections extend significantly further than in control brains (Figures 2C–2E), suggesting that normally Slit limits the growth of sLNv axons. To test whether this effect is dependent on Robo receptors, we used the Gal4/UAS system to inhibit the activity of all Robo receptors specifically in the sLNv using the LNV-specific pdf-Gal4 driver and a dominant negative Robo2 transgene, which lacks the intracellular domain (Robo2ΔC) known to inhibit all three receptors (Bashaw et al., 2000; Godenschwege et al., 2002; Kraut and Zinn, 2004). This results in a significant increase in sLNv axon length (Figures 2F–2H). Similar results were obtained with a dominant negative Robo1 transgene, which also inhibits all three receptors (data not shown). Single-cell visualization, using flip-out clones, confirmed that sLNv individual axons overshoot their targets (Figures S2J and S2K). RNAi knockdown (S) Schematic representation of Slit responses in the VNC compared with the brain. In the VNC Slit spreads from a point source, the midline glia, whereas in the brain expression of Slit in MB generates a distributed source reaching large aspects of the developing and adult brain. Blue indicates the presence and green the absence of Robo. Gray axons indicate unknown Robo expression status, and question marks indicate lack of knowledge of how Robo-Slit signaling regulates brain wiring.

Scale bar in (B) represents 50 µm; all other scale bars represent 60 µm. See also Figure S1.
of each single Robo receptor alone, or the combination of Robo1/2 knockdown, did not significantly alter the length of sLNv axons, although knockdown of Robo3 alone did show a moderate tendency toward longer axons (Figures S2O–S2S). Knockdown of the Robo2/3 combination significantly increased sLNv axonal length (Figures 2I–2K). Conversely, overexpression of either Robo2 (Rajagopalan et al., 2000) or Robo3-GFP fusion protein, which has been shown to express in a manner comparable with endogenous Robo3 (Katsuki et al., 2009; Trunova et al., 2011), was sufficient to significantly decrease sLNv axonal...
growth toward the MB calyx (Figures 2L–2N). Single-cell clones show that sLNv terminal branches arrest and do not grow like wild-type axons (Figures S2J and S2L). Therefore, Robo receptor signaling regulates sLNv axonal growth and guidance and Robo3 appears to exert the strongest effect on sLNv axonal repulsion. The pdf-Gal4 driver also labels the ILNv neurons, which however project axons far from the MB in a more ventral aspect of the brain (Figure S2M). ILNv axons are unaffected by the Robo1–3 manipulations described above, indicating that Robo receptor levels and distance from the MB together determine axonal responses to Slit (Figure S2N). To test this idea, we overexpressed Robo2 in the LNV in a slf mutant background. Reduction in Slit function resulted in a full suppression of the Robo2 gain-of-function phenotype in sLNv axonal growth (Figures 2O–2Q). Finally, RNAi knockdown of Slit specifically in the MB in slf heterozygous animals (Figures 2R–2T) was sufficient to induce increased sLNv axon growth similar to that seen in slf mutants. To reveal whether these defects are due to overgrowth or lack of retraction during development, we performed a developmental analysis of sLNvs in wild-type, Robo2AC, and Robo2 overexpressing conditions (Figures 2U–2W”). We observe that axons do not overshoot and then retract during development in wild-type animals, indicating that sLNv axons grow until they reach their final target. When Robo receptor activity is inhibited, sLNv axons overgrow their normal target area. In contrast, these axons arrest early during development under Robo receptor gain-of-function conditions.

In contrast to the sLNv, the DCN axons do project in the vicinity of the MB and appear not to be repelled by Slit (Figures 3A and 3B). However, overexpression of Robo2 in the DCNs does result in the failure of contralateral projections in 75% of the brains examined, showing that expression of Robo(s) in DCNs is sufficient to make them respond to Slit. DCN axons arrest precisely at the level of the MB in larval brains (Figures 3B and 3C) and eventually turn around and innervate the ipsilateral optic lobes in adult brains (Figures 3D–3G), although some axons stay in the vicinity of the MB, indicating that a few of them may be attracted. This phenotype is rescued in the slt mutant background (Figures S3A–S3C”). This result is consistent with the lack of Robo1–3 expression observed in DCNs (Figures S2F–S2H”). Together, these data suggest that axons respond to Slit in the MB in function of (1) their distance from the MB and (2) the composition of Robo1–3 expression. Next, we analyzed the architecture of sLNv axons in robo1, 2, and 3 mutants (Figures S3D–S3H). Robo1 mutants show normal axon lengths, while robo2 mutants display a variety of early sLNv axonal defects with variable penetrance (Figures S3I–S3L). However, the few robo2 mutant axons that do reach the calyx display normal length of the terminal branches (Figure S3F). Only robo3 mutants phenocopy the slt mutant phenotypes (Figure S3G). Altogether, these data indicate that in the sLNv all three Robo receptors play some role in regulating axonal growth, with Robo3 being the main receptor normally necessary for regulating the length and guidance of the terminal axonal arbors.

**Slit Ligand and Robo Receptors Interact Genetically with RPTP69d**

We sought to gain mechanistic insight into the regulation of Robo activity in central brain development. A major mechanism by which Slit/Robo signaling is regulated in the *Drosophila* VNC is negative regulation of Robo(s) by the Comm protein (Keleman et al., 2002, 2005). However, neither overexpression nor knockdown (using four different RNAi lines) of Comm in sLNv had any negative effects on their axonal projection (Figures S4A–S4E). Another mechanism involves the inhibition of Robo by phosphorylation of a highly conserved intracellular tyrosine (Bashaw et al., 2000). Furthermore, Robo1 was shown to genetically interact
with four RPTPs (Sun et al., 2000), although no direct Robo dephosphorylation has been demonstrated. To test whether RPTPs play a role in MB-mediated Slit/Robo signaling in the brain, we overexpressed three RPTPs previously shown to regulate axon guidance (RPTP10d, RPTP69d, and Lar [Clandinin et al., 2001; Sun et al., 2000]) in sLNv. We find that overexpression of RPTP69d, but not Lar or RPTP10d, phenocopies the gain of function of Robo receptors (Figures S4 F–S4J), causing premature sLNv axon arrest. In contrast, Lar leads to a moderate increase in the length of the axons, indicating that different RPTPs can have even opposite functions in axon growth.

Based on this, we studied the role of RPTP69d in sLNv axon growth and its potential interactions with Slit and Robo. RPTP69d is widely expressed in the larval brain including sLNv and DCNs (Figures S2E–S2I). LNv-specific knockdown of RPTP69d by RNAi significantly increases the length of sLNv axons (Figures 4A–4C). Next, we tested whether RPTP69d interacts with the Slit/Robo pathway. Co-expression of Robo2 or Robo3-GFP with RPTP69d-RNAi partially suppresses Robo2/3 gain of function (Figures 4D–4F). Furthermore, we find that Slit and RPTP69d act synergistically as brain double heterozygous for slit (slit1) and RPTP69d (ptp69d1) show a significant increase in sLNv axon length compared with heterozygous controls (Figures 4 G–4K). These data suggest that RPTP69d acts in the Slit/Robo pathway to regulate sLNv axonal growth.

RPTP69d gain of function stunts sLNv axonal growth and the sLNv express and require all three Robo receptors. We wanted to ascertain whether RPTP69d can induce axonal repulsion in the absence of Robo receptors. To this end, we tested RPTP69d overexpression in DCN axons, which do not express any of the three Robo receptors. RPTP69d overexpression alone in the DCNs has no effect on axonal projection, nor does RPTP69d knockdown (data not shown). In contrast, the expression of Robo2 alone causes disruption of the DCN axon repulsion at the level of the MB axonal lobes in 75% of the brains examined. When Robo2 and RPTP69d are co-expressed, the penetrance of DCN axon repulsion loss increases from 75% to 100% (Figures 4L–4O). This suggests that RPTP69d acts synergistically with Robo receptors to enhance axonal responses to Slit.

Figure 4. RPTP69d Regulates Robo Function during Axonal Growth

(A–C) Effect of knocking down RPTP69d in the sLNv projection using pdf-Gal4 and UAS-RPTP69d-RNAi (B). UAS-Luc-RNAi is used as a control (A). Note that overgrowth of sLNv projections resembles Robo loss-of-function phenotypes. (C) Quantification of RPTP69d-RNAi effect on the projection length (**p < 0.001, two-tailed t test).

(D–F) Downregulation of RPTP69d partially rescue Robo2 and Robo3 overexpression phenotype. Control flies (D) with pdf-Gal4, UAS-GFP; UAS-Robo3-GFP/+; UAS-Luc-RNAi/+ and (E) flies with reduced RPTP69d in the Robo3-GFP background (pdf-Gal4, UAS-GFP; UAS-Robo3-GFP/+; UAS-RPTP69d-RNAi/+). (F) Quantification of the effect of RPTP69d knockdown in the Robo2 and Robo3 gain-of-function phenotype (**p < 0.001, two-tailed t test).

(G–K) Genetic interactions between the ptp69d and slit genes. Heterozygous flies (pdf-Gal4, UAS-GFP;pdf-Gal4, UAS-GFP; ptp69d1/+, UAS-GFP;pdf-Gal4, UAS-GFP; ptp69d1/+, UAS-GFP;pdf-Gal4, UAS-GFP) (G and H) and double heterozygotes (pdf-Gal4, UAS-GFP; pdf-Gal4, UAS-GFP; ptp69d1/+, UAS-GFP;pdf-Gal4, UAS-GFP; ptp69d1/+, UAS-GFP) (I and J). Note that double heterozygotes show axonal overgrowth while relative to heterozygote controls. (K) Quantification of multiple animals (**p < 0.001, one-way ANOVA).

(L–O) RPTP69d cooperates with Robo2. (L) Wild-type, (M) weak commissure phenotype, and (N) severe commissure phenotype. (O) Quantification of the phenotypes of the different groups (**p < 0.001, Fisher’s exact test); number of samples is indicated. Note that although expression of RPTP69d by itself does not produce defects, it enhances the Robo2 phenotype when they are expressed together. In all column scatter plots, data are presented as mean + SEM. A.U, arbitrary units.

Scale bars represent 20 μm in (A) (applies to A–J) and 60 μm in (L) (applies to L–N). See also Figure S4.
RPTP69d Regulates Robo3 Independently of Its Phosphatase Activity

Next, we examined the molecular nature of the interaction between RPTP69d and Robo receptors. We focused on Robo3 because it showed the strongest loss-of-function phenotype in sLNv, and little is known about the molecular regulation of its activity. Work in the embryonic midline suggests that Robo1 activity is regulated by dephosphorylation of a conserved tyrosine. We started by asking whether Robo3 is tyrosine phosphorylated. A GFP-tagged Robo3 was expressed in Drosophila S2 cells, immunoprecipitated, and probed for tyrosine phosphorylation. We find that Robo3, which bears many tyrosine residues, is indeed tyrosine phosphorylated, as confirmed by \( \lambda \)-phosphatase treatment (Figure 5A). Next, we asked whether this phosphorylation is RPTP69d dependent. Surprisingly, RPTP69d co-expression did not reduce tyrosine phosphorylation of Robo3 (Figure 5B) even in the presence of Slit (Figure S5), suggesting that RPTP69d may not be a Robo3 phosphatase, and may regulate Robo3 activity by a different mechanism. To further examine this issue, we asked which domains of RPTP69d are important for its function in vivo. We expressed four mutant forms of RPTP69d in the sLNv: a phosphatase domain mutant (RPTP69d-DA3); a C-terminal deletion mutant (RPTP69d-DC), removing both phosphatase domains; an extracellular domain (N-terminal) deletion mutant (RPTP69d-DN) removing the Ig and fibronectin III (FNIII) domains, but maintaining the signal peptide plus juxtamembrane and transmembrane regions; and a construct including only the extracellular domain (RPTP69d-extra). We tested all these forms in the sLNv axonal repulsion assay. Both RPTP69d-DA3 and RPTP69d-DC mutants caused a significant decrease in sLNv axonal growth indistinguishable from the wild-type RPTP69d (Figures 5D–5G and 5J). Therefore, phosphatase activity is dispensable for RPTP69d function in axonal Robo3-dependent axonal repulsion. In contrast, the RPTP69d-DN and RPTP69d-extra mutant fails to decrease axonal length (Figures 5H–5J), indicating a requirement for the membrane-tethered extracellular domains of RPTP69d in regulating Robo3-dependent axonal growth in vivo.

Robo3 and RPTP69d Can Form Receptor Complexes

Whereas physical interactions between RPTPs, including RPTP69d, and their substrates are known to be very transient and difficult to detect by co-immunoprecipitation (Dascenco et al., 2015; Flint et al., 1997) we were able to readily co-immunoprecipitate Robo3 and RPTP69d from S2 cells (Figures 6A–6C). RPTP69d-DC and RPTP69d-extra were able to bind Robo3, indicating that the interaction may require the transmembrane and/or the juxtamembrane domain of RPTP69d. Together, these data indicate that RPTP69d binds Robo3 but that binding and functional interactions do not require enzymatic activity of RPTP69d.

RPTP69d Increases Surface Presentation of Robo3

Since RPTP69d does not regulate Robo3 through its phosphatase activity, we decided to examine alternatives. One possibility is that RPTP69d directly binds to Slit and in this way increases
Slit at the membrane, where it can bind to Robo and activate the pathway. However, although Robo3 co-immunoprecipitated RPTP69d and Slit, RPTP69d did not bind Slit in the absence of Robo3 (Figures 7A and 7B). Another possibility is that RPTP69d regulates Robo3 shedding, a recently proposed mechanism of Robo signaling activation (Coleman et al., 2010). Although we observed shedding of Robo3 in cell extracts, the level of processing was not significantly changed in the presence of RPTP69d (Figures S6A–S6E). We attempted to create an uncleavable form of Robo3, replacing the FNIII domains in the extracellular domain by the first FNIII domains of Frazzled, the same modification used for generating an uncleavable Robo (Coleman et al., 2010). However, we found no difference in the cleavage pattern between mutant (Robo3-[fraFN]) and wild-type Robo3. Interestingly, however, this form failed to co-immunoprecipitate RPTP69d (Figure S6F).

Finally, we considered the possibility that RPTP69d regulates Robo3 cell surface presentation. The molecules regulating Robo presentation outside the Drosophila VNC are largely unknown. To test whether RPTP69d might be such a regulator, we measured the amount of Robo3 on the axonal surface of primary Drosophila neurons (Figures 7C–7G). We expressed the same Robo3-GFP used in the in vivo experiments (intracellular, C-terminal tag) alone or in combination with wild-type RPTP69d, RPTP69d-ΔC, or RPTP69d-ΔN in primary Drosophila embryonic neurons. We used antibodies against the extracellular domain of Robo3 in the absence of detergents to measure Robo3 levels at the axonal surface, and used GFP fluorescence to determine the total levels of Robo3-GFP. We confirmed that intracellular antibodies do not stain in this condition (Figures S6G–S6H). Next, we calculated the ratio of surface-to-total Robo3 alone or in the presence of various forms of RPTP69d at the growth cone. We find that wild-type RPTP69d and RPTP69d-ΔC, but not RPTP69d-ΔN, result in a significant increase of the extracellular Robo3 signal, indicating an increased abundance in Robo3 on the axonal surface (Figures 7C–7G; insets show extracellular Robo3 alone). This suggests that RPTP69d increases Robo3 levels on the cell surface, consistent with a role in enhancing Robo function. To provide direct biochemical evidence for this interaction, we used a cell surface biotinylation assay to measure the amount of Robo on the cell surface (Figures 7H and 7I). We observe that indeed co-expression of Robo3 with RPTP69d increases Robo3 levels on the cell surface (immunoprecipitate/input ratio). Since the mutant form of Robo3 (Robo3-[fraFN]) did not bind RPTP69d, we used it as negative control in these assays. We find that RPTP69d has no effect on the cell surface fraction of Robo3-[fraFN] (Figures 7H and 7I), showing that RPTP69d binding is essential for increased cell surface levels. Finally, we measured the amount of Robo3 in the membrane of culture cells using an antibody feeding assay (Figures 7J–7L). Cells were transfected with Robo3-GFP alone or with RPTP69d-V5 and treated with anti-Robo-extracellular and anti-GFP to compare the levels of Robo3 on the surface with total levels of Robo3 (see Experimental Procedures for details). We quantified the levels of Robo on the cell surface when expressed alone or together with RPTP69d, and observed a significant increase at the membrane in the presence of RPTP69d (Figures S6I–S6P, for viewing separate channels). Therefore, in three independent assays we find that the presence of RPTP69d significantly increases levels of Robo3 receptors at the cell surface, including that of axonal growth cones. Together, our data indicate that RPTP69d enhances Robo(s) function by binding to it (Figure 6) and increasing its cell surface availability (Figure 7).

**DISCUSSION**

This work reveals a previously unknown function of MB in the Drosophila brain whereby neuropile-neuropile interactions guide axonal growth. Interestingly, a single subtype of neurons, the MB
Kenyon cells, acts as a major source of Slit in the post-embryonic central brain. The reach of the axonal and dendritic arbors of these neurons covers significant areas of both developing brain hemispheres. This allows Kenyon cells to exert profound effects on neighboring neuronal circuits. Consistent with this, we observe significant defects in various brain fibers and neuropils close to the MB. We also uncover a Slit/Robo signaling mechanism that relies on an RPTP, specifically RPTP69d, as a coreceptor necessary to stabilize Robo cell surface levels and enhance its signaling activity. Interestingly, RPTP69d binds to Robo3 but does not require its phosphatase domain to mediate its positive effects on Robo3 function as a repulsive receptor.

The Mushroom Body Is a Source of Axonal Cues

In the embryo, midline glia expresses not only Slit but also the attractive cue Netrin (Brankatschk and Dickson, 2006; Kennedy et al., 1994; Kolodziej et al., 1996; Serafini et al., 1994). Thus, the midline constitutes an organizing center for VNC connectivity. The MB also expresses Netrin, and we find that its receptor Frazered is required for growth of the sLNv axons (Figures S6Q–S6U). This suggests that the MB acts as source of both attractive and repulsive factors. Previous work (Nicolas and Preat, 2005) showed defects in the MB and the central complex, a structure close to the MB, in robo2 and robo3 mutants. The fact that the MB is a major source of Slit in the central brain explains these observations. We find that the MB itself does not express any of the Robo proteins at the larval stage, and previously it was shown that Robo2 and Robo3 are absent from MB in pupae (Nicolas and Preat, 2005), yet it has been reported to show defects in robo2/3 mutants (Nicolas and Preat, 2005). This suggests a feedback mechanism between the MB and surrounding neuropils downstream of Slit/Robo signaling. MB neuroblasts are the first to be activated when the larval brain begins its expansion to form the adult brain (Ito and Hotta, 1992; Prokop and Technau, 1994), and the MB grows during development as the brain incorporates more neurons and circuit. The finding that the MB is a major source of Slit in the central brain may have implications for the interpretation of studies using MB structural or developmental mutants to study behavior.

Differential Regulation of Brain Wiring

Our observations show a different behavior of brain axons in response to Slit/Robo signaling. In the VNC, axons use Slit to...
choose whether to cross or not to cross the midline and to select the appropriate longitudinal pathway among the three available tracts. In the central brain, sLNv axons do not change their trajectory in response to the increase or decrease of Slit; instead they change the length of the projection which is perpendicular to the Slit source (the calyx). This may be the consequence of the interaction between several guidance cues and cell-cell adhesives interactions. Furthermore, although most DCN axons are repelled from MB upon Robo2 ectopic expression, some of them appear to be attracted. This can be the results of Slit itself acting as an attractive cue for these axons or in combination with other guidance cues. Emergent properties of combinations of guidance cues have been already described in other systems. For instance, in the guidance of thalamocortical axons (TCA) Slit enables Netrin to attract TCA axons (Leyva-Diaz et al., 2014). For full understanding of the complex wiring of the brain it will eventually be necessary to characterize the expression patterns of guidance cues and their receptors, and to study their effects in combinations using live-imaging techniques.

A Distinct Mechanism of Robo Regulation in the Brain

During axon guidance at the midline, temporal regulation of the surface presentation of Robo receptors seems to be an important aspect. In Drosophila embryonic commissural axons, this is achieved by the Comm protein. However, in mammals, and even other insects, there are no Comm orthologs, indicating that Robo regulation by Comm is a non-conserved mechanism. Thus, other mechanisms of regulating Robo activity and surface availability remain to be discovered. Here we report that RPTP69d can form a complex with Robo3, thereby increasing its cell surface availability. These data suggest that direct regulation of Robo surface availability by RPTPs during axon growth represents an alternative mechanism to Comm-mediated regulation of Robo activity. RPTPs are conserved through evolution and several studies from invertebrate and vertebrate models have provided evidence of important roles of RPTPs in regulating axon guidance (Gatto et al., 2013; Stepanek et al., 2005), although the respective molecular pathways remain to be characterized. We speculate that the role of RPTPs in the Slit/Robo pathway might be an important mechanism for regulating Slit/Robo signaling in most systems.

It is interesting to note that RPTP69d positively regulates Robo receptor function independent of its enzymatic activity, in contrast to its function in guiding peripheral retinal axons (Garrity et al., 1999) and its phosphatase-dependent ability to regulate signaling of other co-receptors (Dascenzo et al., 2015). Drosophila Lar has also been suggested to function independently of enzymatic activity, but the mechanism remains unknown (Hofmeyer and Treisman, 2009). This may point to RPTPs as dual-activity molecules acting as phosphatases for some guidance receptors, and as co-receptors or chaperones for others. In vertebrates, a major control mechanism for Slit/Robo-dependent repulsion in the spinal cord and the brain appears to be via the divergent ROBO3 receptor, which is mutated in horizontal gaze palsy with progressive scoliosis (Jen et al., 2004). ROBO3 antagonizes the repulsive effects of ROBO1 and ROBO2 (Marillat et al., 2004; Sabatier et al., 2004), yet the molecular underpinnings of this antagonism are unclear. It has also been suggested that signaling and surface presentation of ROBO1/2 can be promoted by co-overexpression of RabGDI (Philipp et al., 2012), but in the absence of mutant analysis it is unclear whether, how, and under what conditions this might occur in vivo. It would therefore be interesting to examine whether ROBO3 and or RabGDI—perhaps through interactions with RPTPs—regulates surface presentation of ROBO1/2.

EXPERIMENTAL PROCEDURES

Fly Culture

Flies were cultures on standard fly food. All experiments were performed under temperature-controlled conditions at 25°C or 28°C (RNAi experiments). Flip-out clones were generated by a 1-hr heat shock at 37°C for 2 days during pupal development.

Cloning

Standard molecular biology techniques were used to make the different RPTP69d constructs and the Robo3-fraFN-GFP construct.

Antibody Staining of Drosophila Brains

For adult and larval brain staining, animals were dissected in PBS, fixed, and stained using standard procedures. The following antibodies were obtained from the Developmental Studies Hybridoma Bank: mouse anti-Slit (1:20), mouse anti-Robo (1:50), mouse anti-Robo3 extracellular (1:50), mouse anti-Robo1 (1:50), mouse anti-Fasciclin2 (1:50), rat anti N-cadherin (1:10). Other antibodies used were: rabbit anti-NetA and rabbit anti-Robo2 (gifts from Barry Dickson; 1:1,000 and 1:500, respectively), mouse anti-GFP 3E6 (Invitrogen, catalog #A11120, 1:25), rabbit anti-GFP (Invitrogen, #A11212, 1:500) rabbit anti-Robo2 (1:1,000), rabbit anti-DsRed (Clontech, #632496; 1:500), and anti-HRP (Cy5-conjugated, Jackson Immunoresearch, 1:50). Secondary antibodies conjugated with Alexa 488, Alexa 555, and Alexa 647 were obtained from Invitrogen and used at 1:500.

Cell Culture

For analysis of Robo3 localization, Drosophila embryonic primary neurons were used. HEK293T cells were used for antibody feeding experiments.

Immunoprecipitation

S2 cells were cultured in 6-well plates at 25°C in SF900II medium. Electroporation was carried out using the Amaxa V kit (Lonza). For Slit treatment, Slit was obtained from the supernatant of S2 stable-expressing cells and was added 3 days after transfection. Six hours later the cells were extracted and the pellet frozen.

Cells were lysed using 400 μL of RIPA buffer and fresh added protease inhibitors cocktail 100× (Amresco). NaF and Na3VO4 phosphatase inhibitors were added if detection of tyrosine phosphorylation was required. Immunoprecipitation was carried out using anti-GFP conjugated beads (Chromotek).

Imaging

Imaging was performed using Leica SP5 and SP6 confocal microscopes (Wetzlar). Images were processed using ImageJ software (NIH). Figures were prepared using Adobe Photoshop (Adobe).

Immunoprecipitation

S2 cells were cultured in 6-well plates at 25°C in SF900II medium. Transfections were performed with indicated constructs. Cells were lysed using RIPA buffer including NaF and Na3VO4, phosphatase inhibitors. anti-GFP conjugated beads (Chromotek) were used for immunoprecipitation.

Western Blotting

SDS-PAGE was performed using 4%–12% gradient polyacrylamide gels and then transferred to nitrocellulose membranes, according to standard protocols.

Statistics

Statistical analysis was performed using Prism software (GraphPad). For sLNv measurements, results are presented as a.u. representing the fraction
between the lengths of the dorsal projections divided by the distance between cell bodies in the two brain hemispheres. A two-tailed t test was used for analysis of two-group comparisons and ANOVA was used for multiple comparisons. For primary neuronal culture experiments the Mann-Whitney test was employed. For analysis of Robo phenotypes in DCN neurons, Fisher’s exact test was performed.

Additional protocols and details are described in Supplemental Experimental Procedures.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and six figures and can be found with this article online at http://dx.doi.org/10.1016/j.devcel.2016.09.028.

AUTHOR CONTRIBUTIONS

Conceptualization, C.O., A.S., N.S.-S., and B.A.H.; Investigation, C.O., A.S., N.M., A.R., and N.S.-S.; Resources, N.D.G., M.-L.E., J.S., D.D., and D.S.; Software, R.K.E.; Writing – Original Manuscript, C.O. and B.A.H.; Writing – Review & Editing, C.O., A.S., B.A.H., D.S., and N.M.; Supervision, B.A.H.; Funding Acquisition, B.A.H., N.S.-S., and D.S., and C.O.

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