Vilse, a conserved Rac/Cdc42 GAP mediating Robo repulsion in tracheal cells and axons

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Slit proteins steer the migration of many cell types through their binding to Robo receptors, but how Robo controls cell motility is not clear. We describe the functional analysis of vilse, a Drosophila gene required for Robo repulsion in epithelial cells and axons. Vilse defines a conserved family of RhoGAPs (Rho GTPase-activating proteins), with representatives in flies and vertebrates. The phenotypes of vilse mutants resemble the tracheal and axonal phenotypes of Slit and Robo mutants at the CNS midline. Dosage-sensitive genetic interactions between vilse, slit, and robo mutants suggest that vilse is a component of robo signaling. Moreover, overexpression of Vilse in the trachea of robo mutants ameliorates the phenotypes of robo, indicating that Vilse acts downstream of Robo to mediate midline repulsion. Vilse and its human homolog bind directly to the intracellular domains of the corresponding Robo receptors and promote the hydrolysis of RacGTP and, less efficiently, of Cdc42GTP. These results together with genetic interaction experiments with robo, vilse, and rac mutants suggest a mechanism whereby Robo repulsion is mediated by the localized inactivation of Rac through Vilse.

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Directed cell migration is a striking feature of the development and physiology of virtually all animals. A particularly dramatic example of directed cell migration takes place in the developing nervous system, as the axons and dendrites of differentiating neurons are guided toward their appropriate target regions by attractive and repulsive cues provided in the extracellular environment (Tessier-Lavigne and Goodman 1996). Several conserved families of axon guidance molecules and their receptors have recently been identified (Dickson 2002). Perhaps not surprisingly, many of these molecules act not only on neuronal growth cones, but can also direct the migration of entire cells, including both neurons and nonneuronal cells. One such guidance cue is Slit, which acts through receptors of the Robo family to direct axon and cell migration in a variety of systems and species (Kidd et al. 1998, 1999; Zallen et al. 1998; Nguyen Ba-Charvet et al. 1999; Kramer et al. 2001). In the Drosophila ventral nerve cord, some axons but not others grow across the midline. Those axons that do cross, called commissural axons, cross the midline only once. This choice between a crossing (commissural) and noncrossing (longitudinal) pathway is controlled by Slit and Robo. Slit is expressed on midline cells, and repels axons expressing its receptor Robo. Commissural axons express only very low levels of Robo, whereas longitudinal axons express high levels of Robo. Accordingly, only the former can cross (Kidd et al. 1998, 1999).

Guided cell migration is also a central process in the development of the tracheal (respiratory) network in Drosophila, which extends its branches into most tissues of the animal. Air enters the system through the spiracles and is delivered directly to internal organs through unicellular thin capillaries, which are made by specialized terminal cells that target and invade the different tissues (Uv et al. 2003). The embryonic nerve cord receives 20 ganglionic branches (GBs), which are initially guided toward the ventral side of the embryo by the expression of the Drosophila FGFR homolog branchless (Sutherland et al. 1996). The tip cell of the ganglionic

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branch (GB1) leads the way toward the ventral nerve cord (VNC) and during embryogenesis navigates a tortuous but invariable path of 50 µm tracking along distinct nerves and glia [Englund et al. 1999]. Inside the VNC, GB1 becomes exposed to a highly diverse array of positional signals provided by its surrounding cells and, with its relatively large size and distinct lineage, provides an advantageous single-cell model for the genetic dissection of the signaling events that steer its migration. GB1 is guided in part by Slit acting through the two receptors Robo and Robo2. Slit appears to act as a repellent through Robo to help prevent GB1 from crossing the midline, and as an attractant through Robo2 to facilitate GB1’s initial extension toward the midline [Englund et al. 2002].

Guidance receptors such as the Robo proteins are thought to direct cell or axon migration by inducing dynamic and spatially coordinated changes in the actin and microtubule network in the growth cone or migrating cell. There is compelling evidence that the Rho family of small GTPases play a critical role in signaling between guidance receptors and the cytoskeleton [Luo 2002]. In particular, Rac (and possibly also Cdc42) plays a critical role downstream of Robo in midline repulsion of CNS axons by Slit and Robo. This was first suggested by the observation that some longitudinal axons aberrantly cross the midline in various mutant combinations for the three Drosophila Rac genes [Hakeda-Suzuki et al. 2002]. This finding was recently extended by Bashaw and colleagues [Fan et al. 2003], who reported genetic interactions between the Rac genes and both slit and robo, as well as biochemical evidence that stimulation of Robo receptors with Slit can lead to an increase in the amount of Rac in its active GTP-bound state. These genetic and biochemical data imply that one or more of the Racs play a positive role in transducing the Robo repulsive signal in axon guidance.

How might Robo regulate the activity of Rac and other Rho family GTPases? Like other small GTPases, these proteins are regulated by the combined activity of guanine nucleotide exchange factors (GEFs), which stimulate the release of GDP and uptake of GTP, and GTPase-activating proteins (GAPs), which stimulate the hydrolysis of the bound GTP. Because the small GTPases are only active when bound to GTP, they become activated by the GEFs and inactivated by the GAPs. In vertebrates, stimulation of Robo1 by Slit has been shown to result in the recruitment and activation of members of the srGAP family of RhoGAPs, which appear to have some specificity for Cdc42 [Wong et al. 2001]. These srGAPs could thus locally inactivate Cdc42 in response to Slit stimulation of the Robo receptor. However, srGAPs neither bind to nor regulate Rac, and so are unlikely to provide a direct link between Robo and Rac [Wong et al. 2001]. Moreover, the srGAPs are not found in flies and worms, where the guidance functions of Slit and Robo proteins were first defined and are best understood. Thus, additional regulatory proteins linking Robo and Rho GTPases are predicted to exist, in both invertebrates and vertebrates.

Here we report the identification of a conserved family of RhoGAPs, the Vilse proteins, with clear representatives in flies and vertebrates. We show that Drosophila Vilse is required for midline repulsion of both CNS axons and tracheal GBs. Genetic and biochemical data support a model in which Vilse provides a direct link between Robo and Rac, and perhaps also Cdc42. Somewhat surprisingly, Vilse appears to have a positive role in Robo repulsion, despite its function as a negative regulator of Rac. This mirrors the presumed action of srGAPs in signaling from Robo to Cdc42, but is perplexing in light of the genetic and biochemical data suggesting a positive role for Rac. Our data are consistent with a more complex model in which both positive and negative factors cooperate to ensure that Rac activity is regulated in the precise temporal and spatial manner required for directed cell or growth cone migration.

Results

vilse is required for tracheal and axonal pathfinding at the VNC midline

The vilse locus was identified in a P-element screen for genes with pathfinding defects in the tracheal GB [J. Hemphälä and C. Samakovlis, unpubl.]. The vilse lacZ strain contained a single P[w + act = lacW] transposon in chromosomal position 93B10-11, which caused misroutings in GB outgrowth and, at lower frequency, crossing of the VNC midline; we therefore named the gene “vilse” [which means “lost” in Swedish]. We generated additional mutants and revertants of this phenotype by excision of the P-element, and chose to study the allele vilse1 because the analysis of the genomic region in this mutant and in situ hybridization indicated that it represents the zygotic null condition for the gene.

In wild-type midstage-16 embryos, the GB1 cell has reached the ventral side of the neuropil, and it turns posteriorly as it migrates in the proximity of the ventral longitudinal glia. Then, just before it reaches the midline, it abruptly turns to migrate dorsally to reach its final target on the dorsal side of the neuropil [Fig. 1A; Englund et al. 1999]. In vilse1 mutants, 20% of the GBs (n = 154) migrated normally to the midline, but stalled once they reached it. An additional 14% of GBs failed to turn posteriorly, instead they extended straight toward the ventral midline, where most of them stalled [Fig. 1B, arrowhead] or, occasionally, continued to migrate across the midline [Fig. 1B, arrow]. In wild-type embryos, <1% of the GBs (n = 140) migrated straight toward the midline, and none of them crossed it. Despite the low penetrance, this misguidance phenotype was interesting because it was similar to the tracheal phenotype seen in robo mutants, where GBs also migrate straight toward the midline but instead of halting there they often cross it [Fig. 1C, arrows; Englund et al. 2002].

Given these similarities between vilse and robo mutants in GB1 guidance, we wondered whether vilse mutants might also show defects in CNS axon guidance similar to those in robo mutants. In wild-type embryos,
Vilse encodes a conserved protein with WW, MyTH4, and RhoGAP domains

To further study the function of the affected gene in tracheal and axonal pathfinding, we cloned the genomic region surrounding the transposon insertion. Database searches with the sequence from genomic DNA flanking the vilse lacZ P-element showed that it was inserted in the 5’-untranslated region of the predicted gene CG3421 [RhoGap93B in GadFly]. The search also identified several cDNAs deriving from this gene, and we sequenced the longest available clone, LD10379 [BDGP]. The predicted Vilse protein contains a number of conserved domains: two N-terminal WW domains [residues 6–36 and 45–75], a more C-terminal myosin tail homology 4 [MyTH4] domain [amino acids 997–1124], a RhoGAP domain [amino acids 1154–1303], and a Pfam-B 53745 domain [amino acids 1304–1330] (Fig. 2A). A predicted human protein, KIAA1688 [Nagase et al. 2000] has an identical domain structure and overall 29% identity and 51% similarity to Drosophila Vilse [Fig. 2], which is in turn the closest match to KIAA1688 in Drosophila. We therefore refer to KIAA1688 as the human Vilse protein. In addition, the human genome encodes a second Vilse homolog with an additional extensin-2 domain (48% similarity, 26% identity, GenBank accession no. gi|37574693|). Vilse homologs can also be found in the mosquito (66% similarity, 29% identity for gi|30176853|) and mouse (50% similarity, 28% identity, gi|28380066|).

Figure 1. Ganglionic branch (GB) and axonal pathfinding defects in vilse1 and robo mutants. Late-stage-16 embryos stained to reveal tracheal lumen [mAb2A12, A–C] and longitudinal fascicles [mAb1D4, D–F]. All panels show ventral views, anterior to the left. In wild-type embryos, GBs [A] and longitudinal fascicles [D] never cross the midline. [B] In vilse1, a few GBs cross the midline [arrow], and several arrest upon reaching it [arrowhead]. [E] Rare midline crosses are observed in vilse longitudinal fascicles. [C] GB1 midline crossing phenotypes in robo [arrows]. [F] Axonal roundabouts at the midline [C–F, arrows]. Bar, 20µm.

Fasciclin II-positive axons project along specific pathways in the longitudinal connectives; they never cross the CNS midline [Fig. 1D]. In robo mutants, many of these axons project along or across the midline [Fig. 1F; Kidd et al. 1998]. A similar phenotype is also seen in vilse mutants [Fig. 1E], albeit at much lower frequency [1 in 30 embryos]. This phenotype too suggested a link between vilse and the slit signaling pathway.
Vilse acts downstream of robo

To further test the idea that Robo and Vilse might act in a common signaling pathway, we examined dosage-sensitive genetic interactions between vilse and slit, robo, or robo2 mutants (Fig. 3). Specifically, we asked whether mutations in these genes would act as dominant enhancers of the vilse phenotype.

First, we analyzed vilse mutant embryos lacking one copy of robo, or of both slit and robo, and we found that the frequency of GBs crossing the midline was increased in both genotypes. In vilse homozygous mutant embryos, only 1% of GBs (n = 154) crossed the midline. In robo+/+,

| Genotype | GB midline cross (%) | Axonal midline cross (cross/embryo) |
|----------|----------------------|-------------------------------------|
| wild type | 0 (n=280)            | 0 (n=0)                             |
| vilse/+ | 1 (n=154)           | 0.03 (n=30)                         |
| robo/+ | 0.5 (n=210)         | 0 (n=10)                            |
| robo/+; vilse/+ | 9 (n=140) | 1.2 (n=19)                         |
| slit;robo/+ | 2 (n=196)       | 3.3 (n=10)                         |
| slit;robo/+; vilse/+ | 6 (n=154) | 12 (n=10)                          |

Figure 3. Genetic interactions of vilse with slit and robo. Ventral views of late-stage-16 embryos showing longitudinal fascicles stained by anti-FasII (A–D) and GBs (mAb2A12, E,F) of different mutant combinations. (A) In heterozygous robo embryos, longitudinal fascicles never cross the midline. [anti-FasII] (B) Midline crossing is evident in all robo/+; vilse embryos. (C) Embryos lacking one copy of slit, robo exhibit three to four midline crosses per embryo. (D) This phenotype is also enhanced in slit, robo/+; vilse embryos. (E,F) Ventral views of robo and robo;BtlGAL4/UASvilse embryos. (F) Expression of UASvilse in all tracheal cells suppresses the ganglionic branch midline crossing phenotype of robo mutants. Overexpression of Vilse also causes GB premature turns (arrowhead) and stalling outside the VNC (arrow). The table shows the quantitation of the phenotypes.

genomes. We did not identify a protein with the same modular structure as Vilse in the Caenorhabditis elegans genome, the closest relative in worms (33% similarity, 13% identity) is encoded by C38D4.5 CE and contains a WW, a PH, and a RhoGAP domain.

In situ hybridization revealed that vilse transcript is ubiquitous during the first stages of development, suggesting a robust maternal contribution (data not shown). Zygotic transcripts were prominent at stage 15 in the tip cells of all tracheal branches, the muscles, and midline cells of the VNC [Fig. 2D, data not shown]. This pattern was the same as the β-gal marker expression in the enhancer trap strain (Fig. 2C, and was absent in vilse mutants (data not shown), indicating that vilse is a strong loss-of-function mutant in RhoGap93B. To analyze the expression of Vilse protein, we raised anti-Vilse antisera. Immunostainings of whole-mount wild-type embryos detected Vilse protein expression in a pattern that mirrored the pattern of the vilse transcript and that of β-gal expression in the vilse LacZ enhancer trap (data not shown). In addition, Vilse antisera stained the epidermis, the peripheral nervous system (PNS) segmental and intersegmental nerves, and the CNS longitudinal connectives and commissures [Fig. 2E, data not shown]. This expression was not detected in the enhancer trap or by in situ hybridization, and may in part reflect the maternal protein. The antisem is specific for Vilse, as the staining was much reduced in vilse mutants [Fig. 2F]. Vilse staining showed a subcellular distribution consistent with a cytoplasmic localization of the protein [Fig. 2G,H, arrow].

The robust maternal contribution of vilse is a probable explanation for the relatively weak phenotype of the vilse mutant embryos, whose phenotypes are likely to be hypomorphic, given the long-lasting presence of maternal product. To analyze the function of maternal vilse and make embryos that lack Vilse completely, we generated vilse germ-line clones using the FLP/FRT technique (Chou and Perrimon 1996). vilse mosaic females produced only a few embryos, all of which were arrested early in embryogenesis, indicating that Vilse is also required for oogenesis (data not shown).

Next, we attempted to rescue the vilse GB phenotype by tracheal-directed expression of a UAS-LD10379 transgene. For this, we used the SRFGAL4 driver which selectively expresses GAL4 in the tracheal terminal cells [Jarecki et al. 1999]. Only 3% of the GBs (n = 154) stalled or migrated straight to the midline in vilse embryos expressing LD10379 in the terminal cells [Fig. 2B], as opposed to 34% in vilse (see above). This rescue experiment confirms our identification of the vilse gene as RhoGAP93B, and also shows that the vilse tracheal phenotype is not a secondary consequence of the axonal defects.

We also noticed a mild gain-of-function phenotype in these experiments, in that 15% of the GBs stalled or turned away from the midline before even reaching the positions of ventral longitudinal glia [Fig. 2B, asterisk]. This phenotype is even stronger in a wild-type background [25%, n = 168], indicating that it is likely due to increased levels of vilse expression. These opposing loss- and gain-of-function phenotypes resemble those for loss- and gain-of-function conditions of Robo [Englund et al. 2002], suggesting that Vilse may be a critical regulatory factor in Robo signaling during GB guidance.
vilse embryos, 9% of GBs (n = 140) crossed the midline. Reduction of both slit and robo in vilse embryos also increased midline crossing (Fig. 3), but the tracheal phenotypes of robo2+/+; vilse embryos were indistinguishable from those vilse embryos. Thus, there is a selective enhancement of the vilse tracheal midline crossing phenotype when the dose of robo or slit and robo is reduced.

In parallel we analyzed the potential functional interaction between vilse, robo, and slit by looking at the longitudinal nerve fascicles. In vilse mutants, there was about one crossover of Fasciclin II positive axons per 30 embryos [0.03 crossovers/embryo, Fig. 3]. This phenotype was enhanced when robo or both slit and robo function was reduced. In robo+/+; vilse mutants, it increased to 1.2 crossovers per embryo [n = 19] and in slit, robo+/+; vilse embryos, this phenotype was enhanced to 12 crossovers per embryo [n = 10; Fig. 3B,D]. Lack of one robo copy alone caused only very weak if any defect, whereas in slit, robo heterozygous embryos only two to four longitudinal axon bundles per embryo crossed the midline [Fig. 3A,C; Kidd et al. 1999].

The genetic interaction between vilse and mutations in robo and slit suggested a function of Vilse in repulsive Slit signaling. If vilse acts downstream of robo in a common pathway, we would predict that overexpression of vilse might reduce the number of tracheal branches that aberrantly cross the midline in robo mutants. To test this, we expressed UAS-vilse in the trachea of homozygous robo embryos, using the btl-GAL4 driver. UAS-vilse expression in robo mutants reduced the GB crossing of the midline phenotype [Fig. 3E,F] by more than half [11%, n = 154 vs. 29% in robo mutants, n = 168]. In addition, an increased number of GBs failed to enter the CNS in these embryos, 21% compared with 1% in wild-type (n = 200) and 6% in robo mutants. This failure of GBs to enter the CNS was also evident when UAS-vilse was overexpressed in wild-type embryos using the btl-GAL4 driver (35%, n = 200) and suggests that vilse overexpression is sufficient to halt the early GB migration toward the CNS.

The WW domain of Vilse binds to the intracellular CC2 Robo domain

We asked whether Vilse might be a direct transducer of Slit signaling by first assessing the ability of glutathione S-transferase (GST)-Vilse-fusion proteins immobilized on glutathione beads to bind different fragments of the Robo intracellular domain, which were translated in vitro and 35S-labeled. Full-length GST-Vilse efficiently interacted with the intracellular part of the Robo receptor and a fragment containing the conserved domains CC0, CC1, and CC2. Constructs containing the CC0 and CC1 or only the CC3 domains did not bind to GST-Vilse, and none of the Robo fragments bound to beads loaded with GST alone [Fig. 4A]. To further test the direct binding of Vilse to Robo and to identify the binding domains...
of the two proteins, we used a yeast two-hybrid assay. Variants of Vilse were expressed as LexA fusions whereas Robo deletions were expressed as fusions to the B42 activation domain. After cotransformation into yeast, only clones containing interacting proteins grow on media lacking leucine.

Also in this assay, Vilse interacted with the intracellular part of the Robo receptor (Fig. 4B,C). The WW domains of Vilse were necessary and sufficient to mediate binding to Robo. In contrast, Vilse did not interact with the intracellular part of the Robo2 receptor. To pinpoint the Robo domain that confers Vilse binding, different deletions of Robo were tested. All Robo variants containing the CC2 motif could bind to the WW domain of Vilse, whereas variants lacking CC2 showed no binding [Fig. 4E]. Moreover, a small 25-amino acid residue fragment of Robo containing CC2 was sufficient to bind to the WW domain of Vilse. This domain is absent in Robo2, providing an explanation for the specificity of Vilse for Robo.

Given the extensive homology between the fly and human Vilse, we tested whether the human proteins could also bind to the human receptor Robo1. Human Vilse and Robo1 variants were tested in the yeast two-hybrid system. Full-length Vilse and the WW domain alone both showed binding to the intracellular domain of human Robo1 [Fig. 5]. Again, the domain in human Robo1 involved in the binding to Vilse contained the proline-rich CC2-motif [Fig. 5], suggesting that the molecular interaction between Robo and Vilse is conserved.

**Vilse is a Rac/cdc42 GAP**

The predicted domain structure of Vilse suggested that it may regulate the activity of RhoGTPases. To examine the ability of Vilse to stimulate GTP hydrolysis, GST-fusion proteins of *Drosophila* Vilse [amino acids 923–1330] and the corresponding part of the human homolog were incubated with Rac1, Cdc42, or RhoA preloaded with \([\gamma-S]GTP\). At regular intervals, the \([\gamma-S]GTP\) that remained bound to the GTPases was measured. The RhoGAP domains of both *Drosophila* Vilse and human Vilse effectively stimulated GTP hydrolysis of Rac1. In addition, human and *Drosophila* Vilse also stimulated the GTP hydrolysis of Cdc42, but less efficiently. None of the Vilse RhoGAP domains was able to stimulate the GTPase activity of RhoA. In contrast, p50RhoGAP was effective against all three GTPases and provided a positive control for the reactions [Fig. 6A].

**Vilse down-regulates Rac activity at the midline**

To identify the GTPase substrate of Vilse during GB1 pathfinding, we first analyzed the tracheal phenotypes of *Drosophila* zygotic mutants in *cdc42* and *rac1 rac2* [Fe-hon et al. 1997; Nq et al. 2002]. Removal of the maternal contribution for *cdc42* and *rac1 rac2* results in defects in oogenesis and early embryonic phenotypes in epithelial morphogenesis and axonal growth, respectively [Genova et al. 2000; Hakeda-Suzuki et al. 2002]. We therefore analyzed embryos lacking the zygotic products of these genes. Those embryos are likely to exhibit reduced levels of *cdc42* or *rac1* and *rac2* at late stages of embryogenesis, resulting in a sensitized background for genetic interactions. GB phenotypes of *rac* and *cdc42* zygotic mutants were heterogeneous, perhaps not surprisingly given the large number of cellular processes mediated by these GTPases. The migration of most GBs was affected in both *rac* and *cdc42*, resulting mainly in arrested branches [34% in *rac1*, *rac2*, *n* = 220; 69% in *cdc42*, *n* = 180], but no midline crosses were observed. Interestingly, both *rac* and *cdc42* mutants also showed a GB phenotype reminiscent of Robo overexpression: A fraction of GBs turned posteriorly prematurely, before reaching the CNS midline (10% in *rac1*, *rac2*, *n* = 220; 7% in *cdc42*, *n* = 180; Fig. 6B,C). This phenotype is also similar to that obtained by overactivation of Robo in GBs either by the ectopic expression of Slit in longitudinal glia or in *commissureless* mutants [Englund et al. 2002]. To test whether reduction of Robo could suppress the premature GB turns, we analyzed *rac1*, *rac2* embryos lacking one copy of functional *robo*: In these mutants the number of GBs turning early before reaching the midline was reduced to half (5%, *n* = 300; Fig. 6D), suggesting that reduction of Rac can be overcome by a decrease in Robo signaling.

The GB phenotype of *rac1*, *rac2* and *cdc42* mutants is also similar to the one generated by Vilse overexpression in the trachea, which caused 35% of the branches to stall and 8% [*n* = 168] to turn early before reaching the midline. To test whether the premature-turn phenotype
caused by the reduction in Cdc42 or Rac may be ameliorated by the removal of zygotic Vilse, we analyzed rac1, rac2, vilse and cdc42; vilse embryos. Reduction of Vilse in rac mutants effectively suppressed the early-turn phenotype [3%, n = 160; Fig. 6D]. This indicates that removal of Vilse, like reduction of Robo, counteracts the effects of reduced Rac activity on migrating GBs. Decrease of Vilse had little or no effect on cdc42 early turns [6%, n = 200]. Nevertheless, Vilse reduction in cdc42 mutants resulted in an overall improvement of the fraction of branches that migrated normally [1%–6%, n = 200]. This implies that both Rac and Cdc42 are Vilse targets during GB migration, but that Vilse mediates Robo repulsion at the midline mainly through the inactivation of Rac.

Discussion

Vilse and its targets

Vilse promotes the hydrolysis of RacGTP and to a lesser extent that of Cdc42GTP. It is thus expected to antagonize the activity of these GTPases on their known effectors. Increasing experimental evidence indicates that Rac and Cdc42 regulate a multitude of cellular responses ranging from establishment of epithelial polarity and integrity to membrane trafficking and the control of planar polarity, in addition to their well known function in modulating the actin cytoskeleton (Van Aelst and Symons 2002). The tracheal phenotypes of vilse embryos are remarkably specific; all branches form, fuse, and grow toward their targets without any apparent defects on epithelial polarity, integrity, or shape. In addition, the rest of the terminal cells that target other internal organs concurrently with GB1 migrate correctly and associate with their targets. We therefore conclude that the primary function of Vilse in the trachea is in the guidance of GB1 migration.

How then does Vilse fulfill its role in cell navigation?

Both of its target GTPases are key regulators of the actin cytoskeleton in several cell types. In fibroblasts, GTP-bound Cdc42 generates actin bundles and characteristic filopodial extensions, possibly through its association with the WASP protein and subsequent stimulation of the actin polymerizing activity of the Arp2/3 complex (Rohatgi et al. 1999; Pollard and Borisy 2003). RacGTP, on the other hand, generates distinct cytoskeletal attributes, membrane ruffling, and lamellipodial protrusions (Ridley et al. 1992). Rac controls actin polymerization through the intermediary protein IRSp53, which associates to the SCAR/WAVE regulator of Arp2/3 activity (Miki et al. 2000). Cdc42 also binds to the IRSp53 adaptor, suggesting that both GTPases regulate actin polymerization through SCAR/WAVE (Govind et al. 2001).

An additional regulatory role for Rac and Cdc42 in cytoskeletal dynamics is exerted through their activating role on PAK [p21 activated kinase; Manser et al. 1994]. PAK in turn activates the LIM kinase, which can phosphorylate the actin depolymerization factor (ADF/cofilin; Arber et al. 1998). Cofilin mediates depolymerization of actin filaments and can also function as a filament-severing factor. Its phosphorylation by LIM-kinase down-regulates its activity and inhibits F-actin depolymerization [Yang et al. 1998]. Thus, the two GTPases in their active form promote the growth of actin filaments by both enhancing polymerization through the Arp2/3 complex and inhibiting severing and depolymerization at the minus end of the filaments. The phenotypic analysis of vilse and GTPase mutants leads us to propose that Vilse antagonizes the function of Rac in promoting actin polymerization locally at the migrating tip of GB1.
tebrates (Wu et al. 2001), epithelial sheets and muscle cells in *Drosophila* (Schimmelpfeng et al. 2001), and in the branching of the lung epithelium in mice (Xian et al. 2001), and more recently it was found to control movements of endothelial cells during angiogenesis (Wang et al. 2003). Slit signal interpretation and the cellular responses it elicits depend on the intracellular domains of Robo receptors (Bashaw and Goodman 1999). The best characterized examples of Robo signal transduction derive from studies of migrating neurons (Giger and Kolodkin 2001; Luo 2002; Schmucker 2003). These studies highlight two basic mechanisms for Slit signaling through Robo. In the first paradigm, derived from studies of early-stage-22 *Xenopus* spinal neurons, Robo silences the attractive Netrin receptor (Bashaw et al. 2000). More recently, Abl was found to collaborate with the cyclase-associated protein CAP, (Bashaw et al. 2000). These studies reveal two basic mechanisms for Slit signaling through Robo. In the first paradigm, derived from studies of migrating neurons (Giger and Kolodkin 2001; Luo 2002; Schmucker 2003). This Netrin-silencing function of Robo is different from the repulsive response to Slit, which is acquired by the same neurons only later, at stage 28 (Stein and Tessier-Lavigne 2001).

Slit-mediated repulsive responses involve the regulation of cytoskeletal organization in the growth cone. In *Drosophila*, the Abelson kinase [Abl] binds to CC3 and phosphorylates a tyrosine in CC1, thereby modulating Robo activity. On the other hand, the Abl substrate Enabled (Ena), a member of the profilin-binding family of proteins, associates with CC2 and mediates the repulsive role of Robo through an unknown mechanism that may involve control of cytoskeletal organization (Bashaw et al. 2000). More recently, Abl was found to collaborate with the cyclase-associated protein CAP, this time to mediate Robo repulsion (Wills et al. 2002). srGAPs bind to the CC3 domain of Robo in response to Slit and aid Cdc42GTP hydrolysis to directly mediate the repulsive response to Slit in cultured anterior subventricular rat neurons. This Cdc42GTP hydrolysis at the site of Robo activation would then result in actin filament depolymerization and severing, thus promoting the turn of the growth cone to the opposite direction. The functional analysis of Vilse identifies a direct transducer of the Slit signal to the inactivation of Rac. *vilse*, *robo*, and *slit* mutants show qualitatively the same phenotypes of midline crossings of tracheal cells and axons. The effect of *vilse* overexpression on the *robo* tracheal phenotypes and the dose-dependent interaction between *slit*, *robo*, and *vilse*, combined with the biochemical analysis indicate that Vilse acts downstream of Robo. Hence, Vilse may play an analogous role to srGAP in locally down-regulating actin polymerization through the hydrolysis of RacGTP and facilitating turning away from the midline.

Paradoxically, both activation and inactivation of Rac appear to interfere with midline crossing and Slit signaling. Expression of constitutively activated Rac causes longitudinal axons to cross the midline, and reduction of Robo signaling enhances this phenotype (Fritz and Van-Berkum 2002). On the other hand, *rac* mutants show strong phenotypes in axonal growth and guidance including midline crossings (Hakeda-Suzuki et al. 2002; Ng et al. 2002), and the latter phenotype becomes more prominent by reduction of Slit (Fan et al. 2003). One possible explanation is that Rac might be involved in multiple cellular processes affecting different aspects of the Slit/Robo pathway. For example Rac might mediate Slit secretion by midline cells or intracellular trafficking of Robo in the axons, in addition to its effect on cytoskeletal dynamics downstream of Robo. The protein adaptor Dock has also been implicated in midline repulsion downstream of Robo. In response to Slit, Dock’s binding to the intracellular domain of Robo is enhanced, leading to the recruitment of the Rac effector kinase Pak. This chain of events has been proposed to bring activated Rac to Robo in response to Slit (Fan et al. 2003). Yet, it is not clear how Rac becomes activated in response to Slit, or how the recruitment of active Rac and Pak might translate in the cellular events that lead to repulsion from Slit. The contradicting models of the function of Rac downstream of robo may be reconciled by considering a sequential interaction of the effectors with the receptor. For example, Vilse may be required initially for severing of actin filaments at the cell extensions that first encounter Slit. The inductive recruitment of Pak to Robo might occur subsequently, perhaps in response to higher concentrations of Slit, promoting cytoskeletal reorganizations that lead to a sustained turning response. This involves a new function of Rac in the context of repulsion from the signal source (Fan et al. 2003). The genetic analysis of midline repulsion reveals that Slit signaling relies on the dynamic and spatially coordinated control of Rac activity. Vilse provides both the first direct link from Robo to the inactivation of Rac, and a molecular handle to address the complex interactions that control repulsion during cell migration.

Materials and methods

**Fly strains**

The 2240, *vilse* enhancer trap line is from the third chromosome collection of lethals from the Department of Genetics, University of Szeged, Hungary (Deak et al. 1997). Excision alleles were generated as described [Robertson et al. 1988]. Excision alleles (211) were balanced over *TM3UbxlacZ* and screened for lethality and embryonic tracheal phenotype. The screen yielded several homozygous viable revertants and the lethal *vilse* (see below). This excision allele failed to complement *Df(3R)N12* (breakpoints 9B1-2;D6-7:*vilse*)); *vilse* mutants die as third instar larvae. Germ-line clones were generated as described (Chou and Perrimon 1996). The tracheal cell marker strain used was *trachealless-lacZ* (1 eve-1; Per-rimon et al. 1991). The following null or strong loss-of-function alleles were used: *robo* <sup>z70</sup> (Kidd et al. 1998), *slit-robo* (Kidd et al. 1999), *cdc42-3* (Fehon et al. 1997), *tac*<sup>100</sup>*rac2* and *tac*<sup>101</sup>*rac2* (Hakeda-Suzuki et al. 2002). For the genetic interaction experiments we constructed the following strains: *robo*<sup>z70</sup>*CyOftzlacZ*, *vilse*<sup>TM3UbxlacZ*, *slitrobo*<sup>CyOftzlacZ*, *vilse*<sup>TM3UbxlacZ*, *rac1<sup>100</sup>*<sup>rac2</sup> *vilse*<sup>/TM3UbxlacZ*, *cdc42-3*<sup>F7lzlacZ*, *vilse*<sup>/TM3UbxlacZ*, *robo*<sup>z70</sup>*CyOftzlacZ*, *rac1<sup>100</sup>*<sup>rac2</sup> *TM3UbxlacZ*, *cdc42-3*<sup>F7lzlacZ*, *robo*<sup>z70</sup>*CyOftzlacZ*. These were crossed to wild-type Oregon-R or to *vilse*<sup>/TM3UbxlacZ*. Antibody staining against β-gal allowed the identification of genotypes in embryos.
The following GAL4 and UAS strains were used: SRF-GAL4 (Jarecki et al. 1999), btl-GAL4 (Shiga et al. 1996).

Antibodies, embryo staining, and whole-mount in situ hybridizations

Embryo fixation, antibody staining, and light and confocal fluorescence microscopy were performed as described (Samakovlis et al. 1996). Primary antisera were mAb2A12 against tracheal lumen (diluted 1:3), mAb2-161 against DSRF [1:2000, M. Gilman, Ariad Corp., Boston, MA], anti-β-galactosidase [1:1500, Cappel], mAb1D4 against Fasciclin II [1:5, from C. Goodman; Van Vactor et al. 1993], mAb BP102 that labels all CNS axons [1:50, from B. Dickson, Institute of Molecular Biotechnology of the Austrian Academy of Sciences, Vienna, Austria]. Biotin-, Cy2-, and Cy3-conjugated (Jackson Laboratories) and Alexa Fluor-568- and Alexa Fluor-488-conjugated (Molecular Probes) secondary antibodies were used at 1:300, 1:500, and 1:400, respectively. Signal was developed using the Vectastain Elite ABC Kit (Vector Laboratories) or Cy3 Tyramide Signal Amplification Kit (TSA, NEN Life Science Products). Embryos were visualized with a Leica TCS SP2 confocal microscope.

The following GAL4 and UAS strains were used: SRF-GAL4 (Jarecki et al. 1999), btl-GAL4 (Shiga et al. 1996).

Glutathione S-transferase (GST) pull-down experiments

Vilse sequences encoding amino acid residues 1–1330, 105–1330, 923–1330, 1131–1330, 1–922, 1–104, and 105–922 into the Apal–Notl sites of the phHybLex/Zeo vector to generate fusion proteins with the LexA DNA-binding domain. PCR was also used to clone robo sequences encoding amino acid residues 941–1395, 941–1256, 1257–1395, 941–1033, 941–1097, 1098–1256, and 1098–1122 and a robo2 sequence encoding residues 1013–1463 into the EcoRI–Notl sites of the pYESTrp2 vector to generate fusion proteins with the B42 activation domain. In a similar manner, we cloned human vilse sequences encoding amino acid residues 1–1094 (starting at the first amino acid in the largest open reading frame), 1–135, 139–697, and 701–1094 into the EcoRI–Xhol sites of phHybLex/Zeo. Human robo sequences encoding amino acid residues 921–1651, 921–1154, 1149–1651, and 1180–1213 were cloned into the EcoRI–Notl sites of pYESTrp2. Interactions were detected on plates lacking leucine.
sample buffer and analyzed by 15% SDS polyacrylamide gel electrophoresis and autoradiography.

**GTPase activation assays**

GST-fusion proteins of Rac1, Cdc42 [brain isoform], and RhoA were expressed in *E. coli*, purified on glutathione-Sepharose beads (Amersham Biosciences) and isolated from GST-fusion proteins by thrombin cleavage as described (Richnau and Aspenstrom 2001). GST-fusion proteins of the RhoGAP domains of *Drosophila melanogaster* Vilse [amino acid residues 923–1330], the corresponding human Vilse residues, and p50RhoGAP [amino acid residues 230–439] were expressed in *E. coli* as described (Richnau and Aspenstrom 2001). Briefly, the bacteria were lysed in a buffer containing 50 mM Tris-HCl at pH 7.5, 5 mM MgCl2, 50 mM NaCl, 10% glycerol, 0.1% Triton X-100, 1 mM PMSF, 1% aprotinin [TrasyloL, Bayer], and 1 mM dithiothreitol (DTT). The proteins were thereafter eluted from the glutathione-Sepharose beads with 5 mM reduced glutathione, desalted on PD10 prepacked chromatography columns [Amersham Pharmacia Biotech], equilibrated in 20 mM Tris-HCl at pH 7.5, 10% glycerol and 1 mM DTT, and thereafter concentrated using Centricon-10 (Millipore). Protein concentrations were determined with the Bradford method. The GTPase assay was described as Richnau and Aspenstrom (2001). Briefly, 0.1 µg of recombinant wild-type Rac1, RhoA, or Cdc42 was preloaded with 10 µCi [γ-32P]GTP (Amersham Pharmacia Biotech) in 20 µL of 20 mM Tris-HCl at pH 7.5, 25 mM NaCl, 5 mM EDTA, and 0.1 mM DTT. The mixture was incubated for 10 min at 30°C, after which the reaction was terminated by adding 5 µL 1 M MgCl2 and the resulting [γ-32P]GTP-loaded GTPase solutions were stored on ice. For the GAP assays, equimolar amounts of the GTPases and the GTPase-domains GAPs were used. Three µL of the [γ-32P]GTP-loaded GTPase was added to a 30-µL mixture of 20 mM Tris-HCl at pH 7.5, 1 mM nonradioactive GTP, 0.87 mg/mL bovine serum albumin, 0.1 mM DTT, with either GST-RhoGAP domains of *Drosophila* melanogaster or GST-fusion proteins of Rac1, Cdc42 (brain isoform), and RhoA as described (Fan, Labrador, Hing, and Bashaw, 2003). Slit stimulation recruits Dock and Pak to the roundabout receptor and increases Rac activity to regulate axon repulsion at the CNS midline. Neuron 40: 113–127.

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Vilse, a conserved Rac/Cdc42 GAP mediating Robo repulsion in tracheal cells and axons

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