

Published in final edited form as:

*Dev Biol.* 2014 August 15; 392(2): 141–152. doi:10.1016/j.ydbio.2014.06.009.

## The Fat-like Cadherin CDH-4 Acts Cell-Non-Autonomously in Anterior-Posterior Neuroblast Migration

Lakshmi Sundararajan, Megan L. Norris, Sebastian Schöneich, Brian D. Ackley, and Erik A. Lundquist\*

Programs in Genetics and Molecular, Cellular, and Developmental Biology, Department of Molecular Biosciences, The University of Kansas, 1200 Sunnyside Avenue, Lawrence, KS, 66045

### Abstract

Directed migration of neurons is critical in the normal and pathological development of the brain and central nervous system. In *C. elegans*, the bilateral Q neuroblasts, QR on the right and QL on the left, migrate anteriorly and posteriorly, respectively. Initial protrusion and migration of the Q neuroblasts is autonomously controlled by the transmembrane proteins UNC-40/DCC, PTP-3/LAR, and MIG-21. As QL migrates posteriorly, it encounters and EGL-20/Wnt signal that induces MAB-5/Hox expression that drives QL descendant posterior migration. QR migrates anteriorly away from EGL-20/Wnt and does not activate MAB-5/Hox, resulting in anterior QR descendant migration. A forward genetic screen for new mutations affecting initial Q migrations identified alleles of *cdh-4*, which caused defects in both QL and QR directional migration similar to *unc-40*, *ptp-3*, and *mig-21*. Previous studies showed that in QL, PTP-3/LAR and MIG-21 act in a pathway in parallel to UNC-40/DCC to drive posterior QL migration. Here we show genetic evidence that CDH-4 acts in the PTP-3/MIG-21 pathway in parallel to UNC-40/DCC to direct posterior QL migration. In QR, the PTP-3/MIG-21 and UNC-40/DCC pathways mutually inhibit each other, allowing anterior QR migration. We report here that CDH-4 acts in both the PTP-3/MIG-21 and UNC-40/DCC pathways in mutual inhibition in QR, and that CDH-4 acts cell-non-autonomously. Interaction of CDH-4 with UNC-40/DCC in QR but not QL represents an inherent left-right asymmetry in the Q cells, the nature of which is not understood. We conclude that CDH-4 might act as a permissive signal for each Q neuroblast to respond differently to anterior-posterior guidance information based upon inherent left-right asymmetries in the Q neuroblasts.

### Introduction

Directed neuronal migration is of fundamental importance in nervous system development. The Q neuroblasts of *Caenorhabditis elegans* are an established and useful model for understanding the molecular mechanisms of directed neuronal migration. The Q neuroblasts are born embryonically in the posterior lateral region of the worm on the right (QR) and left

© 2014 Elsevier Inc. All rights reserved.

\*Corresponding author (erikl@ku.edu).

**Publisher's Disclaimer:** This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

side (QL), and are the sister cells of the V5 hypodermal seam cells (Chalfie and Sulston, 1981; Sulston and Horvitz, 1977). QR and its descendants migrate anteriorly, and QL and descendants migrate posteriorly. Q migration can be divided into two steps, the first of which involves the protrusion and migration of the Q cells, QR to the anterior over the V4 hypodermal seam cell and QL to the posterior over V5 (Chapman et al., 2008; Honigberg and Kenyon, 2000). The Q cells then undergo their first division. The second stage of migration is regulated by EGL-20/Wnt signaling and the Hox molecule MAB-5 (Chalfie et al., 1983; Eisenmann, 2005; Harris et al., 1996; Herman, 2003; Kenyon, 1986; Korswagen et al., 2000; Salser and Kenyon, 1992; Whangbo and Kenyon, 1999). As QL migrates to the posterior, it encounters an EGL-20/Wnt signal from posterior cells, which through canonical Wnt signaling, activates expression of MAB-5/Hox in QL. QR migrates anteriorly away from the EGL-20/Wnt signal and does not activate MAB-5/Hox. MAB-5/Hox is a determinant for further posterior migration of QL descendants. QR continues anterior migration because it does not express MAB-5/Hox.

Initial Q neuroblast protrusion and migration resembles neuroblast migration in the developing vertebrate central nervous system, which extend leading processes followed by nuclear translocation in a saltatory fashion (reviewed in (Solecki et al., 2006)). At 1–2.5 h after hatching to L1 larvae, QR extends a protrusion anteriorly over V4, and QL posteriorly over V5. At 3–3.5 h post-hatching, the cell bodies follow the protrusions and migrate over the respective seam cells. At 4–4.5 h post hatching, the Q cells divide.

Clues about the molecules that control initial Q neuroblast directed protrusion and migration were first provided by (Honigberg and Kenyon, 2000), who showed that the Immunoglobulin-superfamily receptor UNC-40, similar to vertebrate Deleted in Colorectal Cancer, DCC, was required for directed protrusion and migration. Subsequent work delineated a group of transmembrane molecules that interact genetically in regulating Q directional migration, including UNC-40/DCC, the LAR receptor protein tyrosine phosphatase PTP-3, and the small thrombospondin type I-repeat containing protein MIG-21 (Honigberg and Kenyon, 2000; Middelkoop et al., 2012; Sundararajan and Lundquist, 2012). Mutations in all three genes cause misdirected QL and QR migrations. In QL, UNC-40/DCC acts redundantly in parallel to MIG-21 and PTP-3 in posterior QL migration (Middelkoop et al., 2012; Sundararajan and Lundquist, 2012). These molecules interact distinctly in QR, as genetic analysis indicates that UNC-40 and PTP-3/MIG-21 mutually inhibit each other's roles in posterior migration, allowing anterior migration of QR (Sundararajan and Lundquist, 2012). Finally, cell autonomy experiments indicate that UNC-40/DCC, PTP-3/LAR, and MIG-21 act autonomously in the Q cells (Sundararajan and Lundquist, 2012), possibly as receptors for extracellular guidance information. Other molecules have been identified that act in cytoplasmic signaling involving Q cell migrations, including the DPY-19 C-mannosyltransferase that glycosylates thrombospondin repeat proteins including MIG-21 (Buettner et al., 2013; Honigberg and Kenyon, 2000), the MIG-15 NIK-family kinase (Chapman et al., 2008), the Rac GTPases CED-10 and MIG-2, and the GTP exchange factors UNC-73/Trio and PIX-1/PIX (Dyer et al., 2010). These molecules might act downstream of receptor signals to regulate cellular and or cytoskeletal polarity in initial Q migrations.

To identify genes that interact with *unc-40*, *ptp-3* and *mig-21* in QR and QL migration, we performed a forward genetic screen for mutations that disrupted both QR and QL directional migration. We isolated three novel mutations in *cdh-4* gene, which encodes a cadherin repeat-containing transmembrane protein most similar to the Fat family of cadherins (Ackley, 2013; Najarro et al., 2012; Schmitz et al., 2008). In *C. elegans*, CDH-4 has been implicated in axon development and cell migration, including migration of Q cell descendants (Najarro et al., 2012; Schmitz et al., 2008). In *Drosophila*, Fat is thought to coordinate cellular asymmetry generated by the core planar cell polarity pathway components with tissue and organismal polarity in a non-autonomous fashion, and to be distributed asymmetrically across imaginal disc primordia in response to Wnt signaling (Cho and Irvine, 2004; Loveless and Hardin, 2012; Ma et al., 2003; Matakatsu and Blair, 2004; Rawls et al., 2002; Strutt and Strutt, 2002; Tanoue and Takeichi, 2004; Thomas and Strutt, 2012; Yang et al., 2002).

In *C. elegans*, initial Q cell migration is independent of EGL-20/Wnt and MAB-5/Hox (Chapman et al., 2008), although the initial migration can influence the expression of MAB-5 in the Q descendants (Middelkoop et al., 2012). Furthermore, CDH-4 acts independently of MAB-5/Hox in Q descendant migration (Schmitz et al., 2008). Consistent with Wnt-independence of initial Q migrations and of CDH-4 function, we found that the triple Wnt mutant *egl-20 cwn-2; cwn-1* displayed severe Q descendant migration defects but no initial directional migration defects. To determine if CDH-4 acts with the previously-defined signaling system of UNC-40/DCC, PTP-3/LAR and MIG-21 in directing early Q neuroblast migration, we analyzed double and triple mutants. In QL migration, CDH-4 acted genetically in the PTP-3/LAR pathway in parallel to UNC-40/DCC in posterior migration. In QR migration, CDH-4 acted genetically in both the PTP-3/LAR and UNC-40/DCC pathways in mutual inhibition of posterior migration. Mosaic analysis indicated that CDH-4 acts cell-non-autonomously to control QR and QL migration, possibly providing extracellular information to the Q cells about migration. Our results suggest a differential genetic interaction of CDH-4 with UNC-40/DCC in QL versus QR, acting in parallel in QL and acting with both PTP-3/LAR and UNC-40/DCC in QR in mutual inhibition. The nature of this inherent left-right asymmetry in the Q cells is unknown, but QL and QR display other left-right asymmetries, such as sensitivity to the EGL-20/Wnt signal and to MAB-5/Hox (Middelkoop et al., 2012; Whangbo and Kenyon, 1999). Possibly, an as-yet unidentified molecule that is expressed in QR but not QL mediates CDH-4 interaction with UNC-40/DCC in QR but not QL, thus establishing the mutual inhibition mechanism between PTP-3/LAR and UNC-40/DCC that allows anterior migration of QR.

## Results

### ***cdh-4* mutations were identified in a screen for neuronal migration defects**

Our earlier work showed that the transmembrane molecules UNC-40/DCC, PTP-3/LAR and MIG-21 act cell-autonomously in a genetic pathway directing anterior-posterior Q neuroblast migrations (Sundararajan and Lundquist, 2012). To identify new genes that could function with *unc-40*, *ptp-3* and *mig-21*, we performed an ethyl methansulfonate forward genetic screen for new mutations affecting Q migrations (see Materials and Methods), using

the positions of Q descendant neurons AQR and PQR. AQR and PQR are descendants of QR and QL, respectively, and make the longest anterior and posterior migrations of Q descendants (Sulston and Horvitz, 1977; White et al., 1986). AQR migrates anteriorly to a position near the anterior deirid just behind the posterior pharyngeal bulb, and PQR migrates into the phasmid ganglion posterior to the anus. Directional migrations of AQR and PQR are disrupted when earlier Q migrations are defective, and defects in AQR and PQR migrations serve as a proxy for earlier defects in Q neuroblast migrations (Chapman et al., 2008). We focused on new mutations that affected both AQR and PQR directional migration, as do *unc-40*, *ptp-3*, and *mig-21*. The new mutations *lq56*, *lq83*, and *lq97* were identified (Table 1). *lq56* and *lq83* were mapped to linkage group III by single nucleotide polymorphism (snp) mapping against the CB4856 Hawaiian background (Davis et al., 2005). The strains were then subject to whole genome sequencing (see Methods) to identify potential mutations. *lq97* was mapped to LGIII and sequenced using the CloudMap strategy (see Materials and Methods) (Minevich et al., 2012). Each of the three strains carried a novel premature stop codon in the *cdh-4* gene on LGIII (Figure 1A). The lesions were confirmed by polymerase chain reaction of the genomic region and Sanger sequencing.

CDH-4 is a member of the Cadherin superfamily most similar to Fat, a transmembrane molecule with multiple extracellular cadherin repeats, EGF-like repeats, and a laminin G domain (Figure 1B) (Schmitz et al., 2008). *lq83* was in the second exon, *lq56* was in the eighth exon that codes for the sixth cadherin repeat, and *lq97* was in penultimate exon eighteen, introducing a premature stop thirty codons upstream of the region coding for the transmembrane domain (Figure 1A, B). *lq56* and *lq83* failed to complement each other and the previously identified *cdh-4(hd40)* mutation for AQR and PQR migration (data not shown), and a fosmid transgene harboring a wild-type *cdh-4(+)* gene rescued AQR and PQR migration defects of *lq56* animals (Table 1).

Previous studies showed that CDH-4 is required for migration of the Q descendants AVM/PVM and SDQL/R (Schmitz et al., 2008). We found that AQR often migrated posteriorly and PQR often migrated anteriorly in *cdh-4* mutants, including in the previously-characterized alleles *rh310*, *ok1323*, and *hd40* (Schmitz et al., 2008), and the new alleles *lq56*, *lq83*, and *lq97* (Table 1). The AQR migration defects of *lq97* were significantly less severe than *lq56*, *lq83*, *rh310*, *ok1323*, and *hd40* (Table 1), indicating that the *lq97* allele might retain some *cdh-4* function. That the *lq97* premature stop is in the second-to-the-last exon, it is possible that some *lq97* transcripts escape nonsense mediated mRNA decay (Popp and Maquat, 2013) and produce a truncated CDH-4 protein with a signal sequence but no transmembrane domain or cytoplasmic region. This putative truncated protein would be expected to be secreted, and might retain some function, resulting in the less-severe *lq97* phenotype. Double mutants of *lq97* with *smg-1(e1228)* (Grimson et al., 2004) and *smg-2(e2008)* (Johns et al., 2007) mRNA surveillance mutants were lethal, suggesting that in the absence of nonsense-mediated decay, a truncated CDH-4 is produced that results in lethality. *smg-1* feeding RNAi slightly but significantly suppressed PQR migration defects of *lq97* (Table 1). Together, these results suggest that *cdh-4(lq97)* might produce a truncated, secreted version of CDH-4 that is functional.

## CDH-4 is required for anterior-posterior Q neuroblast migrations

We showed previously that defects in Q descendent AQR and PQR migrations can be due to defects in initial QR and QL protrusions and migrations (Chapman et al., 2008; Dyer et al., 2010; Sundararajan and Lundquist, 2012). We next analyzed the early Q neuroblast migrations in the putative null *cdh-4(lq56)* and *cdh-4(lq83)* using the *scm::GFP::CAAX* reporter to observe QR and QL migration as described previously (Chapman et al., 2008). In *wild-type*, at 1–2.5 h post hatching, QR and QL extend protrusions anteriorly and posteriorly over seam cells V4 and V5, respectively (Figure 2A, E). At 3–3.5 h post hatching, the cell bodies follow the protrusions and migrate over the seam cells. At 4–4.5 h post hatching, the Q cells divide on top of their respective seam cells to produce two daughter cells (Figure 2B, F) (Chapman et al., 2008; Dyer et al., 2010; Ou and Vale, 2009; Sundararajan and Lundquist, 2012). In *cdh-4* mutants, we observed directional migration defects and failures to migrate of both QR and QL. Figure 2C shows a QR that has migrated posteriorly over V5 at 3–3.5h, and figure 2D shows a QR that has divided over V5 at 4–4.5h. Figure 2G shows a QL that protruded anteriorly over V4 at 1–2.5 h, and Figure 2H a QL that has divided between V4 and V5, but in contact with the anterior V4.

We quantified the extent of QR and QL migration defects by determining the cell position upon division at 4–4.5 h (Figures 3 and 4). Defects observed at this stage were similar in percentage to defects at the protrusion and migration stages (data not shown) (Sundararajan and Lundquist, 2012). In wild type, no migration defects were observed in QR and QL (Figure 3A and 4A). In putative null *cdh-4(lq56)* and *lq83* mutants, 42% and 27% of QLs migrated anteriorly and divided over V4 (Figure 3A), and 72% and 73% of QRs migrated posteriorly and divided over V5 (Figure 4A). These results suggest that CDH-4 is required for directional migration of both QR to the anterior and QL to the posterior. QR posterior migration was greater than 50% in each mutant, suggesting that QR migration was not randomized. Rather, this result suggests that CDH-4 might alter QR's response to A-P guidance information.

*cdh-4(lq97)* showed significantly reduced AQR migration defects compared to *lq56* and *lq83* (Table 1). Early QL and QR migrations were both significantly less severe in *lq97* compared to *lq56* and *lq83* (Figure 3A and 4A), further indicating that *lq97* might retain function, possibly via the production of a truncated, secreted form of CDH-4.

Fat-like cadherins have been implicated in acting in parallel to Wnt signaling in planar cell polarity, (Cho and Irvine, 2004; Loveless and Hardin, 2012; Matakatsu and Blair, 2004; Rawls et al., 2002; Strutt and Strutt, 2002; Tanoue and Takeichi, 2004; Yang et al., 2002). EGL-20/Wnt, which controls Q descendant migrations including PQR to the posterior, showed no early Q migration defects (Chapman et al., 2008). The five *C. elegans* Wnts act in a complex and redundant fashion in guiding Q descendants (Zinovyeva et al., 2008). The triple mutant *cwn-1; egl-20 cwn-2* displayed extensive defects in the migrations of AQR and PQR (Table 1) but showed no apparent defects in early QL or QR migrations (n = 10; data not shown). While we cannot rule out a role of Wnt in early Q protrusion and migration, these results are consistent with a Wnt-independent role of CDH-4 in early Q protrusion and migration.



## CDH-4 acts genetically in the PTP-3/LAR pathway, in parallel to UNC-40/DCC, in posterior QL migration

Previous studies revealed that UNC-40/DCC, PTP-3/LAR, and MIG-21 each stimulated posterior migration of QL and QR, and that distinct interactions between these molecules in QL versus QR governed posterior versus anterior migration (Sundararajan and Lundquist, 2012). In QL, UNC-40 acts redundantly with PTP-3 and MIG-21 to control posterior QL migration, and in QR, UNC-40 and PTP-3/MIG-21 mutually repress each other's posterior migration function to allow anterior migration. We sought to determine how CDH-4 acts with these molecules in Q cell migration.

We first focused on interactions of *cdh-4* with *unc-40*, *ptp-3* and *mig-21* in QL posterior migration. Double mutants of *unc-40* with *ptp-3* or *mig-21* showed synergistic defects in QL posterior migration (Sundararajan and Lundquist, 2012). *unc-40(n324)* mutants alone displayed 4% QL migration defects wherein QL migrated anteriorly and divided over V4 (Fig 3A). *unc-40; cdh-4* double mutants were lethal, so we reduced *unc-40* function using expression of *unc-40* RNAi from the *scm* promoter expressed in the seam cells and Q cells as described previously (Sundararajan and Lundquist, 2012). *unc-40(RNAi)* showed no defects alone, but *cdh-4; unc-40(RNAi)* double mutants displayed a significant increase in the percentage of anterior QL compared to *cdh-4* alone (Figure 3A). This result indicates that CDH-4 acts in parallel to UNC-40/DCC in posterior QL migration, similar to PTP-3 and MIG-21 (Sundararajan and Lundquist, 2012).

We used *ptp-3(RNAi)* to reduce *ptp-3* gene function, as double mutants of *cdh-4* with the null *ptp-3(mu245)* allele were inviable. The previously-described *scm::ptp-3(RNAi)* transgene was used in these studies, which enhanced QL defects of *unc-40* mutants (Sundararajan and Lundquist, 2012). We found that QL defects in *cdh-4; ptp-3(RNAi)* double mutants did not differ significantly from *cdh-4* single mutants (Figure 3B). The double mutant of *cdh-4(lq56)* with the hypomorphic *ptp-3(mu256)* allele (Sundararajan and Lundquist, 2012) was also no more severe than *cdh-4(lq56)* alone. These data suggest that PTP-3 and CDH-4 might act in the same pathway in parallel to UNC-40 to direct posterior QL migration. Consistent with this idea, the triple mutant *cdh-4(feeding RNAi); unc-40(RNAi); ptp-3(mu256)* showed no enhancement relative to *unc-40; ptp-3* and *cdh-4; unc-40* double mutants (Figure 3A). While *cdh-4* RNAi by feeding alone had no effect, it did significantly enhance PQR migration defects of *unc-40(n324)* ( $p < 0.0001$ ) (Table 1), suggesting that *cdh-4* gene function is reduced by feeding RNAi.

The small transmembrane thrombospondin repeat containing protein MIG-21 acts in the PTP-3 pathway in parallel to UNC-40 in posterior QL migration (Middelkoop et al., 2012; Sundararajan and Lundquist, 2012). Due to close proximity on LGIII and similarity in Q cell phenotype, we were unable to construct *mig-21; cdh-4* double mutants. *cdh-4* RNAi by feeding did not modify the *mig-21(u787)* QL phenotype (Figure 3B). The consistent lack of enhancement between *cdh-4* and *ptp-3/mig-21* support the idea that CDH-4 acts in the PTP-3/MIG-21 pathway, in parallel to UNC-40, in posterior QL migration. While our double and triple mutant analysis using RNAi was consistent with CDH-4 acting in the PTP-3 pathway in parallel to UNC-40, it is also possible that RNAi did not effectively

reduce gene function in *ptp-3* double mutants and that CDH-4 defines a third pathway in parallel to both UNC-40 and PTP-3.

### CDH-4 acts genetically in both the UNC-40/DCC and PTP-3/LAR pathways in QR anterior migration

*unc-40*, *ptp-3* and *mig-21* mutations each displayed QRs that migrated to the posterior over V5 (Sundararajan and Lundquist, 2012). In *unc-40; ptp-3* and *unc-40; mig-21* double mutants, this aberrant posterior migration was reduced, suggesting a mutual inhibition between UNC-40 and PTP-3/MIG-21 in QR that allows anterior migration (Sundararajan and Lundquist, 2012). In other words, in the absence of PTP-3, UNC-40 is free to drive posterior migration, and *vice versa*.

*cdh-4* mutants each displayed over 70% of QRs migrating posteriorly and dividing over V5, significantly more severe than *unc-40*, *ptp-3*, and *mig-21* mutants (Figure 4A).

*unc-40(RNAi)* significantly reduced posterior QR migration of *cdh-4(lq56 and lq83)* (Figure 4A), as it did with *ptp-3* (Sundararajan and Lundquist, 2012). Double mutant combinations with *ptp-3* and *cdh-4* showed reduced posterior QR compared to *cdh-4* alone, but only one was significant at the  $p < 0.05$  level (the *ptp-3(RNAi); cdh-4(lq83)* double) (Figure 4B). *cdh-4(RNAi)* did not significantly modify the QR defects of *mig-21(u787)* (Figure 4B). CDH-4 was clearly involved in inhibition of UNC-40 in QR, as *unc-40(RNAi)* suppressed posterior QR migration in *cdh-4* mutants. CDH-4 might also act in inhibition of PTP-3, as *ptp-3; cdh-4* doubles showed consistent reductions in posterior QR, only one of which was significant at  $p < 0.05$ . These data are consistent with CDH-4 acting in both the PTP-3/LAR and UNC-40/DCC pathways to facilitate a mutual inhibition between them to allow anterior QR migration. The triple *unc-40(RNAi); ptp-3(mu256); cdh-4(feeding RNAi)* showed reduced posterior QR, but was not significantly different compared to *unc-40(RNAi) ptp-3(mu256)* alone. It is possible that posterior QR defects in *unc-40(RNAi) ptp-3(mu256)* are already too low to detect any significant enhancement. However, this result is consistent with CDH-4 acting in both the UNC-40/DCC and PTP-3/LAR pathways in QR. This model accounts for why the QR defects of *cdh-4* mutants are significantly stronger than *unc-40* and *ptp-3*, as in *cdh-4* mutants, both UNC-40 and PTP-3 would be free to drive posterior migration. *cdh-4; unc-40* and *cdh-4; ptp-3* double mutants have significantly more posterior QR than the *unc-40; ptp-3* double mutant, suggesting that CDH-4 is not the only molecule that acts with UNC-40 and PTP-3 in mutual inhibition.

### *cdh-4::gfp* is expressed in Q cells and surrounding cells

A recombiner version of a fosmid clone containing the full-length *cdh-4* genomic locus and surrounding region with *gfp* fused in frame at the *cdh-4* 3' end was obtained from the TransgeneOme project (Sarov et al., 2006). This transgene is predicted to drive expression of CDH-4::GFP under its endogenous promoter and regulatory regions. Expression of *cdh-4::gfp* was observed in many cells at the time of Q cell polarization and migration in early L1 larvae (1–4.5h post hatching), including hypodermal seam cells and P cells (Figure 5). *cdh-4::gfp* expression was also observed in the Q cells, as noted by overlapping expression with *egl-17::mCherry*, expression of which is limited to the Q cells in the posterior of early L1 animals (Branda and Stern, 2000; Cordes et al., 2006; Sundararajan and

Lundquist, 2012). Thus *cdh-4::gfp* is expressed in the Q cells but also in neighboring hypodermal cells. CDH-4::GFP accumulated throughout the cell bodies and margins, including the nucleus. This accumulation could reflect endogenous localization of CDH-4, or could be an artifact of transgenic expression, possibly overexpression. However, *cdh-4::gfp* rescued the Q migration defects of *cdh-4* mutants, indicating that some functional CDH-4::GFP is expressed from the transgene.

### CDH-4 acts non-cell autonomously in Q migrations

To test the functional requirement of *cdh-4* in the Q cells, we performed a mosaic analysis using the *cdh-4::gfp* fosmid transgene and a strategy used previously to demonstrate autonomy of function of *mig-15/NIK* kinase in the Q cells (Chapman et al., 2008) (see Materials and Methods). A stably-integrated *gcy-32::cfp* transgene was used to score the positions of AQR and PQR, and an unstable extrachromosomal array containing rescuing *cdh-4::gfp* and *gcy-32::yfp* was used to generate genetic mosaics of *cdh-4(+)* activity in the *cdh-4(lq56)* mutant (Figure 6A). The *cdh-4::gfp* extrachromosomal array rescued AQR and PQR defects in *cdh-4(lq56)* mutants, indicating that it supplies *cdh-4(+)* activity (Figure 7). We next screened for mosaic animals in which the *cdh-4::gfp* extrachromosomal array was specifically lost in either AQR or PQR but retained in the other and in the URX neurons using *gcy-32::cfp* expression (Figure 6). The positions of AQR or PQR that lost the array were then determined by *gcy-32::yfp*. In such mosaic animals, AQR and PQR migration defects were not significantly different from non-mosaic, rescued animals (Figure 7). These results indicate that *cdh-4* is not required in AQR and PQR for their proper migration. A similar strategy with *mig-15/NIK* showed opposite results, that loss of the rescuing array in AQR and PQR led to migration defects, indicating autonomy of function (Chapman et al., 2008). These results suggest that CDH-4 acts non-autonomously in directing AQR and PQR migrations. These results do not indicate which tissues require *cdh-4(+)* for AQR and PQR migration, but *cdh-4::gfp* is expressed broadly, including in hypodermal cells neighboring the Q cells (Figure 5).

## Materials and Methods

### C. *elegans* genetics and microscopy

Experiments were performed at 20° C using standard *C. elegans* techniques (Sulston and Brenner, 1974). The following mutations and transgenics were used: LGI *unc-40(n324)*, *smg-1(e1228)*, *smg-2(e2008)*; LGII *ptp-3(mu256)*, *cwn-1(ok546)*; LGIII *mig-21(u787)*, *cdh-4(lq56)*, *cdh-4(lq83)*, *cdh-4(lq97)*, *cdh-4(rh310)*, *cdh-4(ok1323)*, *cdh-4(hd40)*; LGIV *egl-20(n585)*, *cwn-2(ok895)*, *lqls80[Pscm::gfp::caax]*; LGV *lqls58[Pgcy-32::cfp]*. Chromosomal locations for the following transgenes were not assigned: *lqls146[Pscm::unc-40(RNAi)]*, *lqls166[Pscm::ptp-3(RNAi)]*, *lqls249[Pegl-17::mCherry]*. Extrachromosomal arrays used were *lhEx397[unc-119; cdh-4(+):gfp fosmid]*, and *lqEx755[Pgcy-32::yfp; cdh-4::gfp fosmid]*. The extrachromosomal arrays were built using standard gonadal microinjection techniques and were integrated into the genome using UV-TMP techniques (Mello and Fire, 1995). *cdh-4* and *smg-1* feeding RNAi was conducted using the standard RNAi by feeding protocol with bacterial stocks from the “Ahringer”



library (Source Bioscience, UK) (Kamath and Ahringer, 2003). The *cdh-4::gfp* fosmid clone 77167333417072 A06 was obtained from the TransgeneOme project (Sarov et al., 2012).

Micrographs were obtained using a Leica DM550 compound microscope with filters for GFP, YFP, CFP, and mCherry and a Qimaging Retiga camera. The micrographs in Figure 5 were obtained using a Nikon laser scanning confocal microscope.

### Imaging and scoring Q neuroblast defects

Previously described larval synchronization techniques were used (Chapman et al., 2008; Sundararajan and Lundquist, 2012). Adult and larval animals were washed from plates using M9 buffer. Eggs remained adhered to the agar surface of the plates. Newly-hatched L1 larvae were collected every 30 minutes by washing the plates with M9 and placing the larvae onto fresh NGM plates. These larvae were then visualized at 2, 3 and 4 hours post hatching using a the *Pscm::gfp::caax* transgene *lqIs80*. All the wild-type larvae visualized at 2 hours post synchronization (2–2.5 h timepoint) displayed defined anterior QR protrusions over the seam cell V4 and defined posterior QL protrusions over the seam cell V5. At 3 hours post synchronization (the 3–3.5 h timepoint), QR had migrated anteriorly on V4 and QL posteriorly on V5. At 4 hours post synchronization (4–4.5 h timepoint) QR had divided on V4 and QL on V5. Defects in mutant backgrounds were scored for direction of protrusion, migration and division. QR that protruded, migrated and divided posteriorly on V5 were scored as defective. QL that protruded, migrated and divided anteriorly on V4 were scored as defective. Cells also failed to migrate in *cdh-4* and other mutants, but only directional migration defects were scored and included in Figures 3 and 4. At least 25 cells were scored for all stages, but only the 4–4.5 h division stage is included in Figures 3 and 4, as the position of division correlates with defects in protrusion and migration (Sundararajan and Lundquist, 2012). Statistical differences between genotypes were determined by Fisher's exact test.

### Scoring and analysis of AQR and PQR defects in Table 1 and Figure 7

Previously described quantification methods were used (Chapman et al., 2008; Sundararajan and Lundquist, 2012). The *gcy-32::cfp* transgene *lqIs58* was used to visualize the positions of AQR and PQR in L4 larval animals, well after AQR and PQR undergo their migrations in L1 larvae. The final positions of AQR and PQR were binned into five positions along the body of the worm. Position 1 is the wild-type position of AQR in the anterior deirid ganglion just posterior to the pharyngeal bulb. Position 2 represents the region posterior to the pharyngeal bulb and just anterior to the vulva. Position 3 represents the region around the vulva. Position 4 represents the region posterior to the vulva and anterior to posterior deirid, where the Q cells are born and begin their migrations. Position 5 is the wild-type position of PQR, behind the anus in the phasmid ganglion. To best represent the defects in direction of migration, only the percentage of PQR in position 1 and the percentage of AQR in position 5 were included in Figure 7, but Table 1 shows each position. At least 100 worms were scored for each mutant background and all comparisons were done using Fisher's exact test.

## Mosaic analysis

The *cdh-4(+):gfp* fosmid transgene *lqEx755* rescued the defects in AQR and PQR migration observed in a *cdh-4(lq56)* background (Table 1 and Figure 7). A *Pgcy-32::yfp* construct was included with the *cdh-4(+)* extra chromosomal array to visualize the presence and absence of *cdh-4(+)* in AQR, PQR, and URX L/R. During mitotic divisions, extrachromosomal arrays are spontaneously lost from some daughter cells, resulting in genetically mosaic animals. Animals with losses in AQR and PQR were identified by lack of *Pgcy-32::yfp* expression in these cells (Figure 6). The strain also carried the stably-integrated *Pgcy-32::cfp* transgene *lqIs58* to mark the positions of AQR in all animals, including those that lost the rescuing *cdh-4(+)* array in AQR and PQR. Animals that lost the rescuing *cdh-4(+)* array in AQR, but retained it in PQR and the URX neurons were scored for AQR position; and those that lost the rescuing *cdh-4(+)* array in PQR, but retained it in AQR and the URX neurons were scored for PQR position (Figure 7).

## Discussion

Fat-like cadherins have been implicated extensively in cooperating with Wnt signaling and PCP core pathway components to generate tissue polarity (reviewed in (Thomas and Strutt, 2012)). Here we show that the Fat-like cadherin CDH-4 in *C. elegans* is required for anterior-posterior protrusion and migration of the Q neuroblasts. In *cdh-4* mutants, QR often migrated posteriorly, and QL anteriorly. Greater than 50% of QR cells migrated in the wrong direction posteriorly in *cdh-4* mutants, indicating that migration direction is not randomized in *cdh-4* mutants, but rather that CDH-4 might change the direction of migration of Q cells in response to similar guidance information. Wnts control the anterior-posterior migration of Q cell descendants. EGL-20/Wnt activates expression of MAB-5/Hox in QL and descendants to cause posterior migration (Chapman et al., 2008; Eisenmann, 2005; Harris et al., 1996; Harterink et al., 2011; Herman, 2001; Kenyon, 1986; Maloof et al., 1999; Salsler and Kenyon, 1992), and the five *C. elegans* Wnts display complex interactions in anterior-posterior migration of Q descendants (Zinovyeva et al., 2008). However, EGL-20/Wnt is not required for Q neuroblast initial migrations (Chapman et al., 2008), and despite having severe defects in AQR and PQR migration, the Wnt triple mutant *egl-20 cwn-2; cwn-1* had no defects in initial Q migrations. These results are consistent with early Q cell migration and thus CDH-4 function in early Q migration being Wnt-independent.

We found that *cdh-4; unc-40* and *cdh-4; ptp-3* double mutants were inviable. *cdh-4* mutants show some embryonic lethality coupled with variable failures in elongation and hypodermal organization (Schmitz et al., 2008). *ptp-3* mutants when combined with *vab-1/EphR* display defects in embryonic morphogenesis (Harrington et al., 2002), and *ptp-3; unc-40* double mutants are lethal (Sundararajan and Lundquist, 2012). Possibly, UNC-40/DCC, PTP-3/LAR, and CDH-4 all act in parallel to control aspects of embryonic morphogenesis and gastrulation.

### CDH-4 acts in parallel to UNC-40, in the PTP-3/LAR pathway, in posterior QL migration

Previous work showed that UNC-40/DCC, PTP-3/LAR and MIG-21 are each required for posterior migration (Honigberg and Kenyon, 2000; Middelkoop et al., 2012; Sundararajan

and Lundquist, 2012; Williams, 2003). In QL, MIG-21 and PTP-3 function in the same genetic pathway redundantly with UNC-40 to direct posterior migration (Sundararajan and Lundquist, 2012). In QL, *cdh-4* mutations enhanced *unc-40/DCC* but not *ptp-3/LAR* or *mig-21*, suggesting that CDH-4 acts in the PTP-3/LAR pathway (Figure 3). CDH-4 acted non-autonomously (Figures 6 and 7), while UNC-40/DCC and PTP-3/LAR were required autonomously (Sundararajan and Lundquist, 2012). One model is that CDH-4 acts as an extracellular cue that directs posterior QL migration via PTP-3/LAR (Figure 8). This model implies that an unidentified, possibly redundant, cue would act through UNC-40/DCC. This cue is unlikely to be the canonical UNC-40/DCC ligand UNC-6/Netrin, as *unc-6* mutants have no effect on anterior-posterior Q migrations (Honigberg and Kenyon, 2000). It is also possible that RNAi reduction of gene function was ineffective in *cdh-4(RNAi)*; *ptp-3* double mutants, and that CDH-4 defines a third pathway in parallel to both UNC-40 and PTP-3.

### **CDH-4 acts in both the UNC-40/DCC and PTP-3/LAR pathways in QR, and possibly facilitates mutual inhibition**

Previous results indicate that UNC-40/DCC and PTP-3 interact differently in QR versus QL. In QR, UNC-40/DCC and PTP-3/LAR mutually inhibit each others' roles in posterior migration, resulting in anterior migration (Sundararajan and Lundquist, 2012). *cdh-4* mutants displayed greater than 70% posterior QR migration, a significantly higher defect than *unc-40* and *ptp-3* (Figure 4). *unc-40(RNAi)* reduced posterior QR migration in *cdh-4* (Figure 4), suggesting that when CDH-4 is absent, UNC-40 drives posterior QR migration and that CDH-4 normally inhibits UNC-40. *ptp-3* loss also reduced posterior QR migration in *cdh-4* mutants. This effect was not as dramatic as with *unc-40*, but three different double mutant combinations showed reduced posterior QR, one of which was significant. Thus, CDH-4 might also normally inhibit PTP-3/LAR. The triple *unc-40(RNAi)*; *ptp-3(mu256)*; *cdh-4(feeding RNAi)* showed reduced posterior QR, but was not significantly different compared to *unc-40(RNAi) ptp-3(mu256)* alone. It is possible that posterior QR defects in *unc-40(RNAi) ptp-3(mu256)* are already too low to detect any significant enhancement. However, this result is consistent with CDH-4 acting in both the UNC-40/DCC and PTP-3/LAR pathways in QR, explaining why QR defects are so much more severe in *cdh-4* mutants versus *unc-40* and *ptp-3* (i.e. both UNC-40 and PTP-3 drive posterior QR in the absence of CDH-4) (Figure 8).

Our results are consistent with a model in which CDH-4 facilitates the mutual inhibition of UNC-40/DCC and PTP/LAR in QR by interacting with both pathways in a non-autonomous manner (Figure 8). However, CDH-4 does not act in the UNC-40/DCC pathway in QL. This model requires that a QR-specific function, such as a co-receptor or a cytoplasmic molecule, mediates interaction of CDH-4 with the UNC-40 pathway in QR and not QL, and thus facilitates mutual inhibition in QR but not QL. QL and QR have inherent differences, such as response to EGL-20/Wnt and MAB-5/Hox. QL is more sensitive to both than QR (Middelkoop et al., 2012; Whangbo and Kenyon, 1999). The conserved transmembrane protein MIG-13 specifically affects QR descendant migrations (Sym et al., 1999; Wang et al., 2013). However, MIG-13 does not affect initial QR migration (Sym et al., 1999). Instead, MIG-13 expression is activated specifically in QR descendants by the Hox transcription factors LIN-39 and MAB-5 after initial QR migration (Wang et al., 2013). A

molecule that mediates CDH-4/UNC-40 interaction in QR and not QL to mediate early migration has not yet been identified.

### CDH-4 acts non-autonomously

UNC-40/DCC and PTP-3/LAR are required in the Q cells to guide their migrations (Sundararajan and Lundquist, 2012). In contrast, mosaic analysis of *cdh-4* suggests that it acts non-autonomously to guide Q migrations. Loss of *cdh-4(+)* activity in either AQR or PQR had no effect on their migrations (Figures 6 and 7), suggesting that *cdh-4* is required in other cells that retained the *cdh-4(+)* activity in the mosaic animals. Our experiments could not determine which cells these were, but *cdh-4(+)::gfp* is expressed in cells neighboring the Q cells, such as the hypodermal seam cells and the hypodermal P cells (Figure 5). In QL, our data support a model in which CDH-4 might be a cue that directs posterior QL migrations via PTP-3/LAR (Figure 8). In QR, our data suggest CDH-4 that facilitates mutual inhibition of UNC-40/DCC and PTP-3/LAR because of an inherent left-right asymmetry in QR versus QL (Figure 8). This inherent asymmetry would allow CDH-4 to interact with the UNC-40/DCC pathway in QR but not QL, thus inhibiting both UNC-40/DCC and PTP-3/LAR and allowing anterior migration. This is similar to the role of Fat in planar cell polarity, which acts non-autonomously to orient cells polarized by the core PCP pathway to the tissue or organismal axis (Yang et al., 2002; Zeidler et al., 1999).

Despite non-autonomous function in determining direction of migration in QR and QL, *cdh-4::gfp* is expressed in the Q cells. Possibly, CDH-4 has a role in the Q cells that does not involve direction of migration, such as cell adhesion. Indeed, *cdh-4* mutants display defects in attachment of the pharynx to the hypodermis, suggestive of a defect in cell adhesion (Schmitz et al., 2008).

A non-autonomous role of CDH-4 is supported by the *lq97* mutation, which is in the second to the last exon but is still 894 bp upstream of the endogenous stop codon and is thus likely subject to nonsense-mediated mRNA decay. It is possible that a fraction of the mRNA escapes nonsense mediated decay and produces a CDH-4 molecule that lacks a transmembrane domain and is potentially secreted. AQR migration defects of *cdh-4(lq97)* were significantly less severe than other *cdh-4* alleles (Table 1), and *smg-1* RNAi slightly suppressed PQR migration defects, indicating that a potentially secreted form of CDH-4 in *cdh-4(lq97)* mutants retains some function. That this secreted version retains function in AQR migration supports a non-autonomous role.

CDH-4::GFP showed no apparent asymmetric distribution in the anterior-posterior axis, suggesting that it might act as a permissive cue for migration, rather than a directional cue. Possibly, CDH-4 modifies the response of the Q cells to the same anterior-posterior guidance information. That QR migration is not randomized in *cdh-4* mutants but rather skewed toward the posterior is consistent with a permissive role of CDH-4 in distinguishing the responses of QR and QL to similar anterior-posterior guidance information. The *Fat* gene in humans has been associated with neuropsychiatric disorders such as bipolar disorder and schizophrenia (Abou Jamra et al., 2008; Jung and Jun, 2013; Light et al., 2007; Redies et al., 2012). Mechanisms described here might be relevant in vertebrate central nervous system development and neuropsychiatric disorders.

## Acknowledgments

We thank E. Struckhoff for technical assistance, the Lundquist and Ackley labs for discussion and critical feedback, and O. Hobert, A. Boyanov, M. Doitsidu, and G. Minevich for assistance with next generation sequencing and CloudMap analysis. Some strains were provided by the CGC, which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). This work was supported by NIH grants R01NS040945 and R21NS070417 to E.A.L. and NIH grant P20GM103418, the Kansas Infrastructure Network of Biomedical Research Excellence, on which M.L.N. was an Undergraduate Scholar. M.L.N. and S.S. were University of Kansas Undergraduate Research Award Scholars. Some next generation sequencing was provided by the University of Kansas Genome Sequencing Core Laboratory, funded by the NIH infrastructure grant P20GM103638.

## Bibliography

- Abou Jamra R, Becker T, Georgi A, Feulner T, Schumacher J, Stromaier J, Schirmbeck F, Schulze TG, Propping P, Rietschel M, Nothen MM, Cichon S. Genetic variation of the FAT gene at 4q35 is associated with bipolar affective disorder. *Mol Psychiatry*. 2008; 13:277–284. [PubMed: 17938632]
- Ackley BD. Wnt-signaling and planar cell polarity genes regulate axon guidance along the anteroposterior axis in *C. elegans*. *Dev Neurobiol*. 2013
- Branda CS, Stern MJ. Mechanisms controlling sex myoblast migration in *Caenorhabditis elegans* hermaphrodites. *Dev Biol*. 2000; 226:137–151. [PubMed: 10993679]
- Buettner FF, Ashikov A, Tiemann B, Lehle L, Bakker H. C. *elegans* DPY-19 is a C-mannosyltransferase glycosylating thrombospondin repeats. *Mol Cell*. 2013; 50:295–302. [PubMed: 23562325]
- Chalfie M, Sulston J. Developmental genetics of the mechanosensory neurons of *Caenorhabditis elegans*. *Dev Biol*. 1981; 82:358–370. [PubMed: 7227647]
- Chalfie M, Thomson JN, Sulston JE. Induction of neuronal branching in *Caenorhabditis elegans*. *Science*. 1983; 221:61–63. [PubMed: 6857263]
- Chapman JO, Li H, Lundquist EA. The MIG-15 NIK kinase acts cell-autonomously in neuroblast polarization and migration in *C. elegans*. *Dev Biol*. 2008; 324:245–257. [PubMed: 18840424]
- Cho E, Irvine KD. Action of fat, four-jointed, dachsous and dachs in distal-to-proximal wing signaling. *Development*. 2004; 131:4489–4500. [PubMed: 15342474]
- Cordes S, Frank CA, Garriga G. The *C. elegans* MELK ortholog PIG-1 regulates cell size asymmetry and daughter cell fate in asymmetric neuroblast divisions. *Development*. 2006; 133:2747–2756. [PubMed: 16774992]
- Davis MW, Hammarlund M, Harrach T, Hullett P, Olsen S, Jorgensen EM. Rapid single nucleotide polymorphism mapping in *C. elegans*. *BMC genomics*. 2005; 6:118. [PubMed: 16156901]
- Dyer JO, Demarco RS, Lundquist EA. Distinct roles of Rac GTPases and the UNC-73/Trio and PIX-1 Rac GTP exchange factors in neuroblast protrusion and migration in *C. elegans*. *Small GTPases*. 2010; 1:44–61. [PubMed: 21686119]
- Eisenmann DM. Wnt signaling. *WormBook*. 2005:1–17. [PubMed: 18050402]
- Grimson A, O'Connor S, Newman CL, Anderson P. SMG-1 is a phosphatidylinositol kinase-related protein kinase required for nonsense-mediated mRNA Decay in *Caenorhabditis elegans*. *Mol Cell Biol*. 2004; 24:7483–7490. [PubMed: 15314158]
- Harrington RJ, Gutch MJ, Hengartner MO, Tonks NK, Chisholm AD. The *C. elegans* LAR-like receptor tyrosine phosphatase PTP-3 and the VAB-1 Eph receptor tyrosine kinase have partly redundant functions in morphogenesis. *Development*. 2002; 129:2141–2153. [PubMed: 11959824]
- Harris J, Honigberg L, Robinson N, Kenyon C. Neuronal cell migration in *C. elegans*: regulation of Hox gene expression and cell position. *Development*. 1996; 122:3117–3131. [PubMed: 8898225]
- Harterink M, Kim DH, Middelkoop TC, Doan TD, van Oudenaarden A, Korswagen HC. Neuroblast migration along the anteroposterior axis of *C. elegans* is controlled by opposing gradients of Wnts and a secreted Frizzled-related protein. *Development*. 2011; 138:2915–2924. [PubMed: 21653614]
- Herman M. *C. elegans* POP-1/TCF functions in a canonical Wnt pathway that controls cell migration and in a noncanonical Wnt pathway that controls cell polarity. *Development*. 2001; 128:581–590. [PubMed: 11171341]

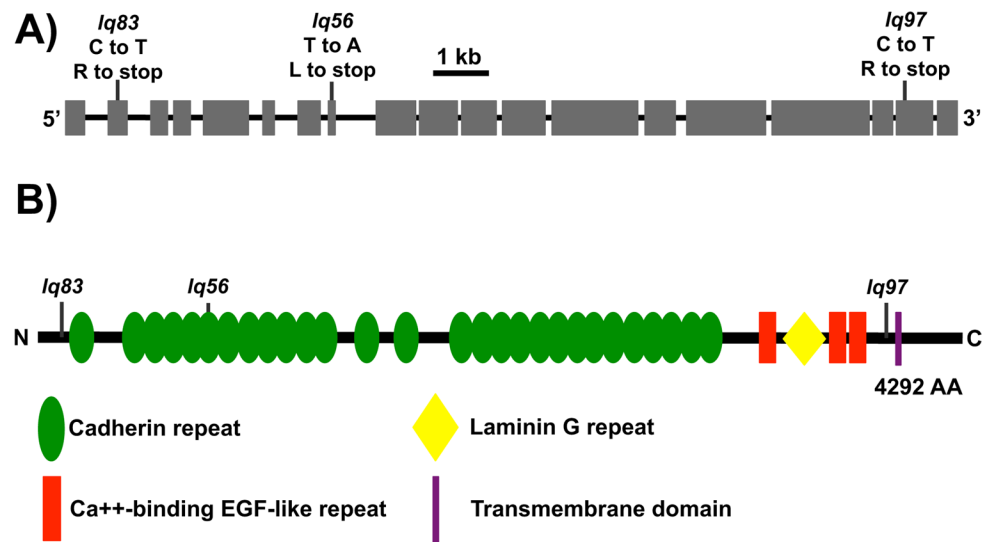


- Herman, MA. Wnt signaling in *C. elegans*. In: Kühl, M., editor. Wnt signaling in Development. Landes Biosciences; Georgetown, TX: 2003. p. 187-212.
- Honigberg L, Kenyon C. Establishment of left/right asymmetry in neuroblast migration by UNC-40/DCC, UNC-73/Trio and DPY-19 proteins in *C. elegans*. *Development*. 2000; 127:4655–4668. [PubMed: 11023868]
- Johns L, Grimson A, Kuchma SL, Newman CL, Anderson P. *Caenorhabditis elegans* SMG-2 selectively marks mRNAs containing premature translation termination codons. *Mol Cell Biol*. 2007; 27:5630–5638. [PubMed: 17562857]
- Jung YE, Jun TY. Association between FAT Gene and Schizophrenia in the Korean Population. *Clin Psychopharmacol Neurosci*. 2013; 11:67–71. [PubMed: 24023550]
- Kamath RS, Ahringer J. Genome-wide RNAi screening in *Caenorhabditis elegans*. *Methods*. 2003; 30:313–321. [PubMed: 12828945]
- Kenyon C. A gene involved in the development of the posterior body region of *C. elegans*. *Cell*. 1986; 46:477–487. [PubMed: 3731276]
- Korswagen HC, Herman MA, Clevers HC. Distinct beta-catenins mediate adhesion and signalling functions in *C. elegans*. *Nature*. 2000; 406:527–532. [PubMed: 10952315]
- Light KJ, Miller AL, Doughty CJ, Joyce PR, Olds RJ, Kennedy MA. FAT and bipolar affective disorder. *Mol Psychiatry*. 2007; 12:899–900. [PubMed: 17895925]
- Loveless T, Hardin J. Cadherin complexity: recent insights into cadherin superfamily function in *C. elegans*. *Curr Opin Cell Biol*. 2012; 24:695–701. [PubMed: 22819515]
- Ma D, Yang CH, McNeill H, Simon MA, Axelrod JD. Fidelity in planar cell polarity signalling. *Nature*. 2003; 421:543–547. [PubMed: 12540853]
- Malool JN, Whangbo J, Harris JM, Jongeward GD, Kenyon C. A Wnt signaling pathway controls hox gene expression and neuroblast migration in *C. elegans*. *Development*. 1999; 126:37–49. [PubMed: 9834184]
- Matakatsu H, Blair SS. Interactions between Fat and Dachshous and the regulation of planar cell polarity in the *Drosophila* wing. *Development*. 2004; 131:3785–3794. [PubMed: 15240556]
- Mello C, Fire A. DNA transformation. *Methods Cell Biol*. 1995; 48:451–482. [PubMed: 8531738]
- Middelkoop TC, Williams L, Yang PT, Luchtenberg J, Betist MC, Ji N, van Oudenaarden A, Kenyon C, Korswagen HC. The thrombospondin repeat containing protein MIG-21 controls a left-right asymmetric Wnt signaling response in migrating *C. elegans* neuroblasts. *Dev Biol*. 2012; 361:338–348. [PubMed: 22074987]
- Minevich G, Park DS, Blankenberg D, Poole RJ, Hobert O. CloudMap: a cloud-based pipeline for analysis of mutant genome sequences. *Genetics*. 2012; 192:1249–1269. [PubMed: 23051646]
- Najarro EH, Wong L, Zhen M, Carpio EP, Goncharov A, Garriga G, Lundquist EA, Jin Y, Ackley BD. *Caenorhabditis elegans* flamingo cadherin fmi-1 regulates GABAergic neuronal development. *J Neurosci*. 2012; 32:4196–4211. [PubMed: 22442082]
- Ou G, Vale RD. Molecular signatures of cell migration in *C. elegans* Q neuroblasts. *J Cell Biol*. 2009; 185:77–85. [PubMed: 19349580]
- Popp MW, Maquat LE. Organizing principles of mammalian nonsense-mediated mRNA decay. *Annual review of genetics*. 2013; 47:139–165.
- Rawls AS, Guinto JB, Wolff T. The cadherins fat and dachshous regulate dorsal/ventral signaling in the *Drosophila* eye. *Curr Biol*. 2002; 12:1021–1026. [PubMed: 12123577]
- Redies C, Hertel N, Hubner CA. Cadherins and neuropsychiatric disorders. *Brain Res*. 2012; 1470:130–144. [PubMed: 22765916]
- Salser SJ, Kenyon C. Activation of a *C. elegans* Antennapedia homologue in migrating cells controls their direction of migration. *Nature*. 1992; 355:255–258. [PubMed: 1346230]
- Sarov M, Murray JI, Schanze K, Pozniakovski A, Niu W, Angermann K, Hasse S, Rupprecht M, Vinis E, Tinney M, Preston E, Zinke A, Enst S, Teichgraber T, Janette J, Reis K, Janosch S, Schloissnig S, Ejsmont RK, Slightam C, Xu X, Kim SK, Reinke V, Stewart AF, Snyder M, Waterston RH, Hyman AA. A genome-scale resource for in vivo tag-based protein function exploration in *C. elegans*. *Cell*. 2012; 150:855–866. [PubMed: 22901814]

- Sarov M, Schneider S, Pozniakovski A, Roguev A, Ernst S, Zhang Y, Hyman AA, Stewart AF. A recombineering pipeline for functional genomics applied to *Caenorhabditis elegans*. *Nat Methods*. 2006; 3:839–844. [PubMed: 16990816]
- Schmitz C, Wacker I, Hutter H. The Fat-like cadherin CDH-4 controls axon fasciculation, cell migration and hypodermis and pharynx development in *Caenorhabditis elegans*. *Dev Biol*. 2008; 316:249–259. [PubMed: 18328472]
- Solecki DJ, Govek EE, Tomoda T, Hatten ME. Neuronal polarity in CNS development. *Genes Dev*. 2006; 20:2639–2647. [PubMed: 17015428]
- Strutt H, Strutt D. Nonautonomous planar polarity patterning in *Drosophila*: dishevelled-independent functions of frizzled. *Dev Cell*. 2002; 3:851–863. [PubMed: 12479810]
- Sulston JE, Brenner S. The DNA of *Caenorhabditis elegans*. *Genetics*. 1974; 77:95–104. [PubMed: 4858229]
- Sulston JE, Horvitz HR. Post-embryonic cell lineages of the nematode, *Caenorhabditis elegans*. *Dev Biol*. 1977; 56:110–156. [PubMed: 838129]
- Sundararajan L, Lundquist EA. Transmembrane Proteins UNC-40/DCC, PTP-3/LAR, and MIG-21 Control Anterior-Posterior Neuroblast Migration with Left-Right Functional Asymmetry in *Caenorhabditis elegans*. *Genetics*. 2012; 192:1373–1388. [PubMed: 23051647]
- Sym M, Robinson N, Kenyon C. MIG-13 positions migrating cells along the anteroposterior body axis of *C. elegans*. *Cell*. 1999; 98:25–36. [PubMed: 10412978]
- Tanoue T, Takeichi M. Mammalian Fat1 cadherin regulates actin dynamics and cell-cell contact. *J Cell Biol*. 2004; 165:517–528. [PubMed: 15148305]
- Thomas C, Strutt D. The roles of the cadherins Fat and Dachshous in planar polarity specification in *Drosophila*. *Dev Dyn*. 2012; 241:27–39. [PubMed: 21919123]
- Wang X, Zhou F, Lv S, Yi P, Zhu Z, Yang Y, Feng G, Li W, Ou G. Transmembrane protein MIG-13 links the Wnt signaling and Hox genes to the cell polarity in neuronal migration. *Proc Natl Acad Sci U S A*. 2013; 110:11175–11180. [PubMed: 23784779]
- Whangbo J, Kenyon C. A Wnt signaling system that specifies two patterns of cell migration in *C. elegans*. *Mol Cell*. 1999; 4:851–858. [PubMed: 10619031]
- White JG, Southgate E, Thomson JN, Brenner S. The structure of the nervous system of the nematode *Caenorhabditis elegans*. *Philos Trans R Soc Lond*. 1986; 314:1–340. [PubMed: 22462104]
- Williams, L. PhD thesis. University of California; San Francisco: 2003. A genetic analysis of the left-right asymmetric polarizations and migrations of the Q neuroblasts in *C. elegans*.
- Yang CH, Axelrod JD, Simon MA. Regulation of Frizzled by fat-like cadherins during planar polarity signaling in the *Drosophila* compound eye. *Cell*. 2002; 108:675–688. [PubMed: 11893338]
- Zeidler MP, Perrimon N, Strutt DI. The four-jointed gene is required in the *Drosophila* eye for ommatidial polarity specification. *Curr Biol*. 1999; 9:1363–1372. [PubMed: 10607560]
- Zinovyeva AY, Yamamoto Y, Sawa H, Forrester WC. Complex Network of Wnt Signaling Regulates Neuronal Migrations During *Caenorhabditis elegans* Development. *Genetics*. 2008; 179:1357–1371. [PubMed: 18622031]

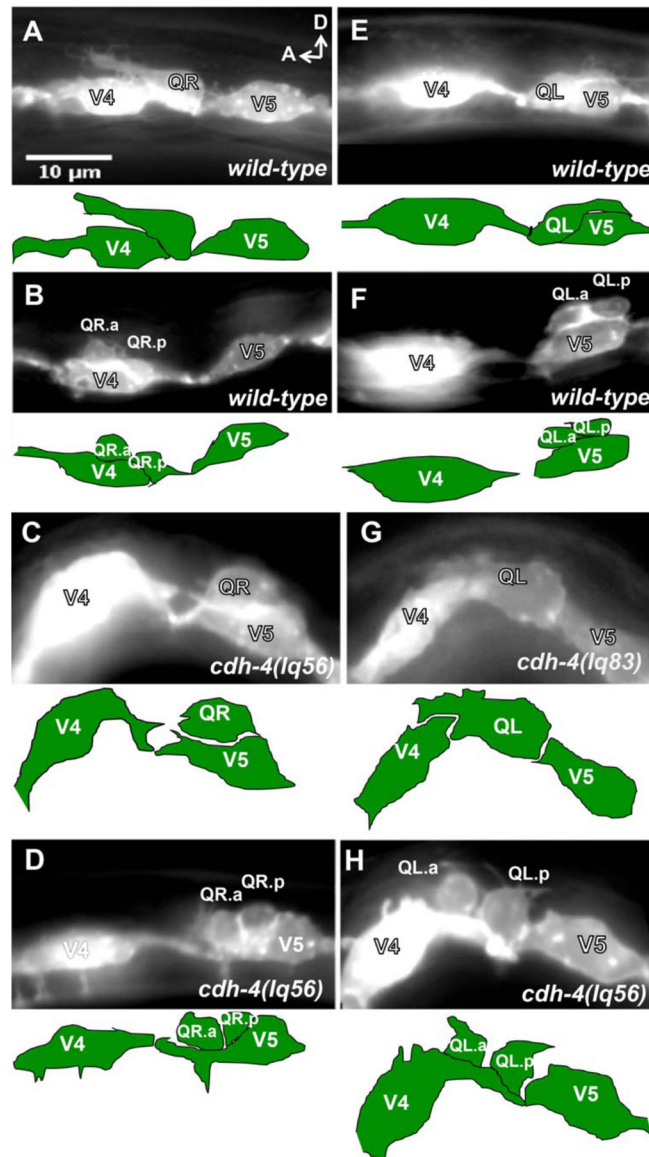
### Highlights

1. The Fat-like cadherin CDH-4 controls QR and QL neuroblast direction of protrusion and migration.
2. CDH-4 acts with the LAR receptor tyrosine phosphatase PTP-3.
3. CDH-4 acts in parallel to the Ig-family receptor UNC-40, related to DCC.
4. CDH-4 displays left-right asymmetry of function in QL (left) versus QR (right).
5. CDH-4 acts cell-non-autonomously in Q migrations.



**Figure 1. CDH-4 is a Fat-like cadherin**

A) A diagram representing the genomic region of the *cdh-4A* isoform. Exons are indicated by boxes, and introns by lines. Positions of the lesions are marked and the corresponding nucleotide changes are noted. *lq56* was a T to A transversion resulting in a leucine to stop (TTG to TAG) at LGII position 4525658 in Wormbase WS240. *lq83* was a C to T transition resulting in an arginine to stop (CGA to TGA) at position 4521154. *lq97* was C to T transition resulting in an arginine to stop (CGA to TGA) at position 4535721. B) A diagram of the 4292 residue CDH-4A polypeptide. The position of the stop codons introduced by *lq56*, *lq83*, and *lq97* are indicated.

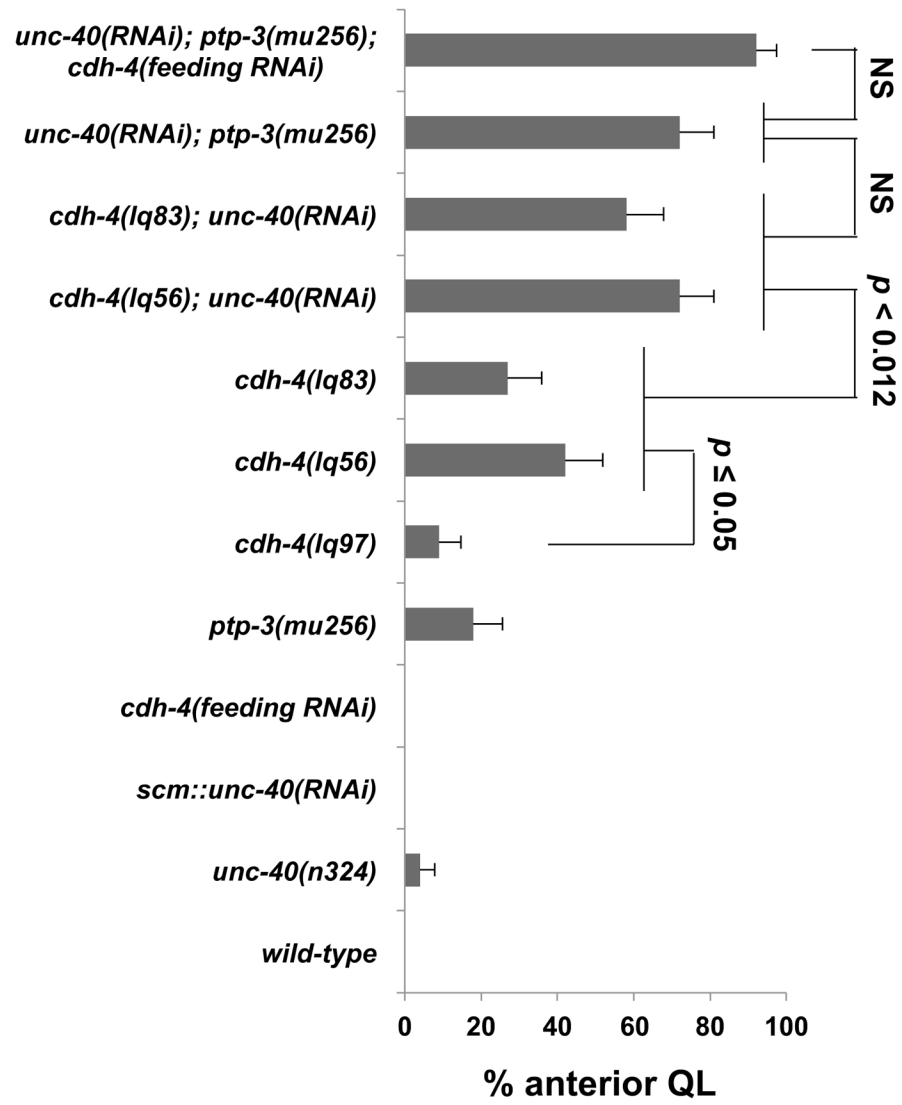


### Figure 2. Q neuroblast migration defects in *cdh-4* mutants

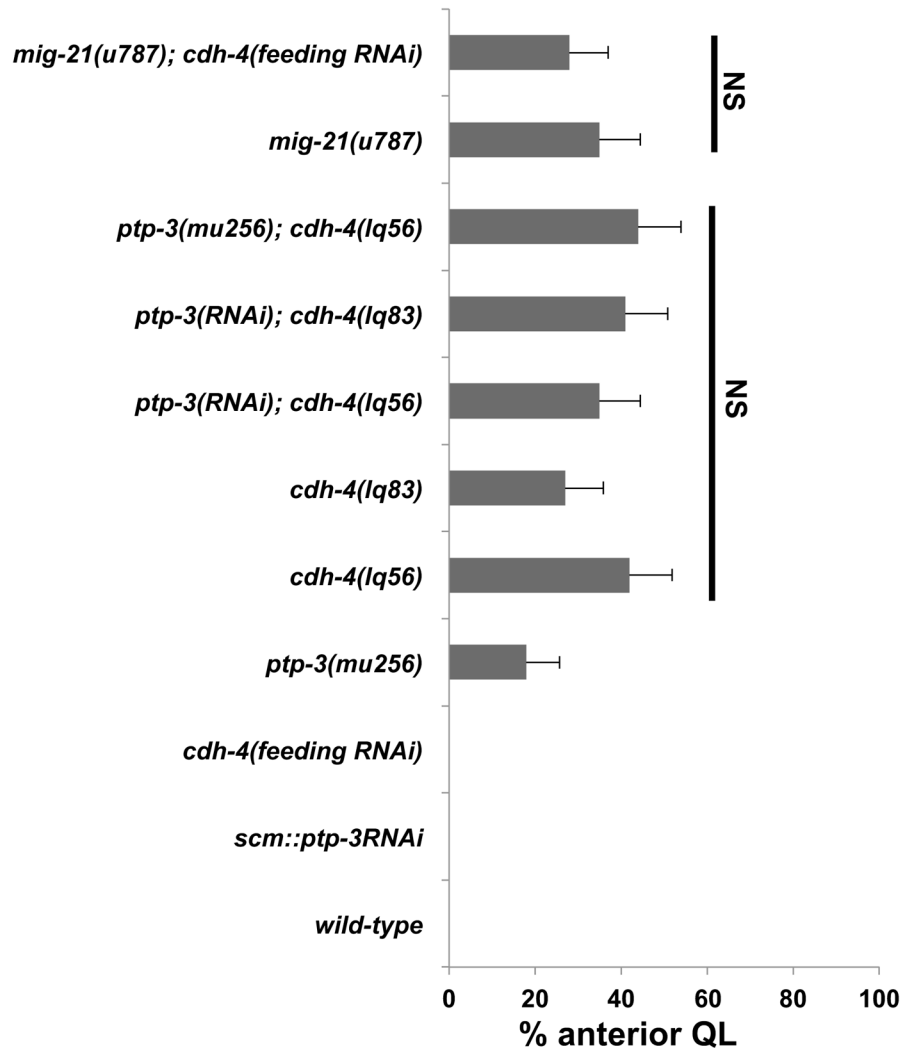
Micrographs of animals with *scm::myr::gfp* expression in the Q cells and seam cells.

Anterior is left, and dorsal is up. The tracing below each micrograph indicates the position of the Q cells and seam cells. (A–D) QR migration in wild-type and *cdh-4*. A) A QR of a wild-type L1 animal at 2–2.5h post-hatch protruded anteriorly over V4. B) A wild-type QR at 4–4.5 h post-hatch divided over V4. C) A *cdh-4(lq56)* mutant QR migrated posteriorly over V5 at 3–3.5 h post-hatch. D) A *cdh-4(lq56)* mutant QR divided over V5 at 4–4.5 h post-hatch. (E–H) QL migration in wild-type and *cdh-4*. E) A wild-type QL at 2–1.5 h post-hatch protruded posteriorly over V5. F) A wild-type QL at 4–4.5 h post-hatch divided over V5. G) A *cdh-4(lq83)* QL at 2–2.5 h post-hatch protruded anteriorly over V4. H) A *cdh-4(lq56)* mutant QL divided over V4 at 4–4.5 h post-hatch.





**Figure 3A**



### Figure 3B

#### Figure 3. Quantification of QL migration in *cdh-4* mutants

Graphs represent the division stage of early QL neuroblast migrations at 4–4.5 hours post hatching. Genotype is the Y-axis, and the percentage of defective anterior QL migration and division is the X-axis. Migration was scored as defective if QL migration was reversed and it divided while in contact with the anterior V4 seam cell. Failure to migrate, resulting in division between the seam cells, was also observed in *cdh-4* mutants but is not included in the graphs. *unc-40(RNAi)* and *ptp-3(RNAi)* represent the *Pscm::unc-40(RNAi)* and *Pscm::ptp-3(RNAi)* transgenes (Sundararajan *et al.*, 2012), and *cdh-4* RNAi by feeding is indicated (see Materials and Methods). The error bars represent two times the standard error of proportion, and the statistical difference between the genotypes were determined by

Fisher's Exact test. Twenty-five animals or more were scored for each genotype. A) *cdh-4* double mutants with *unc-40*. B) *cdh-4* double mutants with *ptp-3* and *mig-21*.

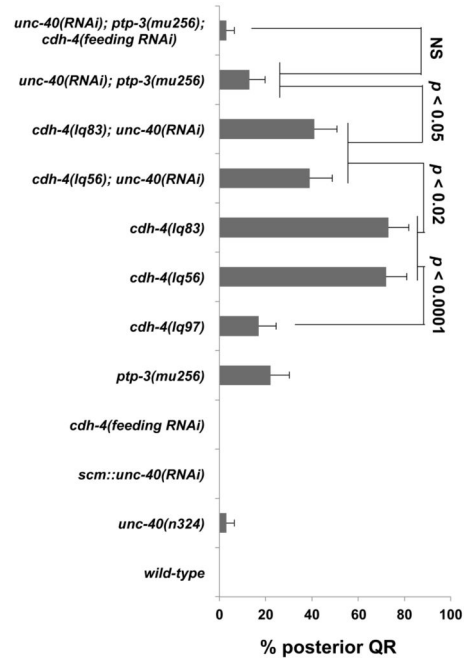
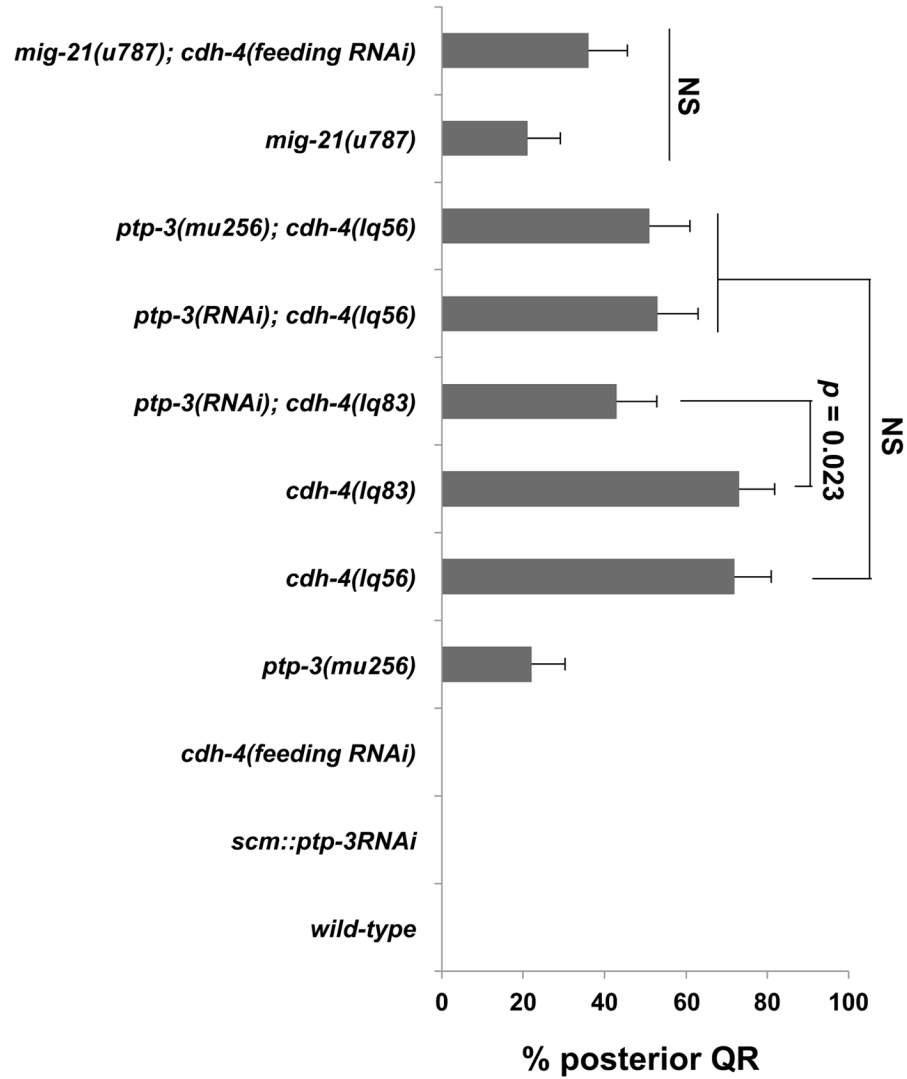


Figure 4A



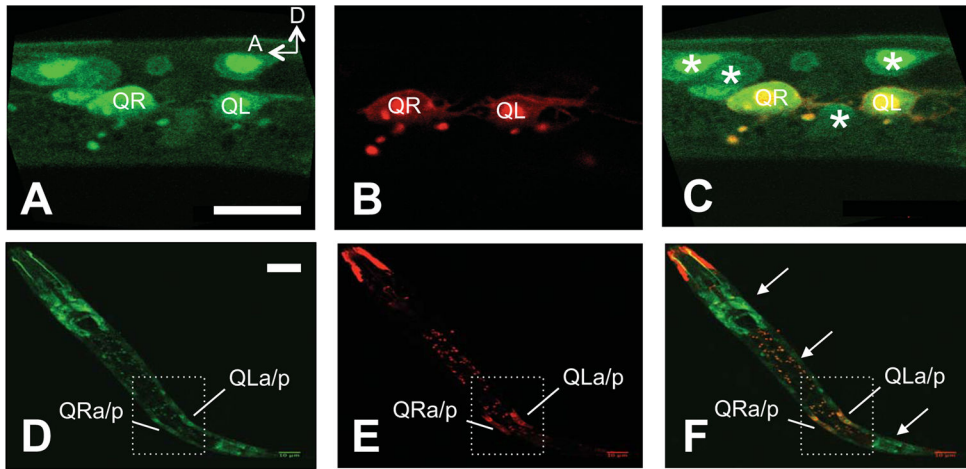
## Figure 4B

### Figure 4. Quantification of QR migration in *cdh-4* mutants

Graphs represent the division stage of early QR neuroblast migrations at 4–4.5 hours post hatching. Genotype is the Y-axis, and the percentage of defective posterior QR migration and division is the X-axis. Migration was scored as defective if QR migration was reversed and it divided while in contact with the posterior V5 seam cell. Failure to migrate, resulting in division between the seam cells, was also observed in *cdh-4* mutants but is not included in the graphs. *unc-40RNAi* and *ptp-3(RNAi)* represent the *Pscm::unc-40(RNAi)* and *Pscm::ptp-3(RNAi)* transgenes (Sundararajan *et al.*, 2012), and *cdh-4* RNAi by feeding is indicated (see Materials and Methods). The error bars represent two times the standard error of proportion, and the statistical difference between the genotypes were determined by

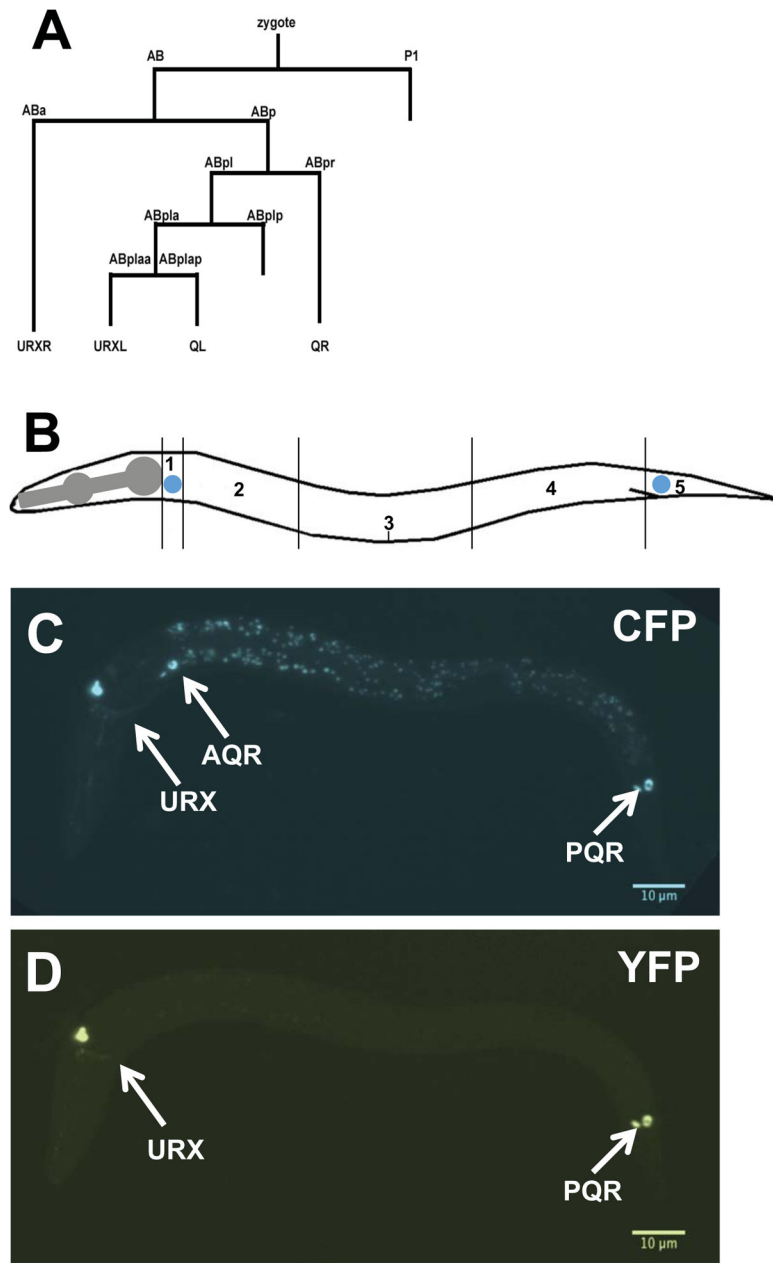


Fisher's Exact test. Twenty-five animals or more were scored for each genotype. A) *cdh-4* double mutants with *unc-40*. B) *cdh-4* double mutants with *ptp-3* and *mig-21*.



**Figure 5. Expression of *cdh-4::gfp* in the Q cells and neighboring cells**

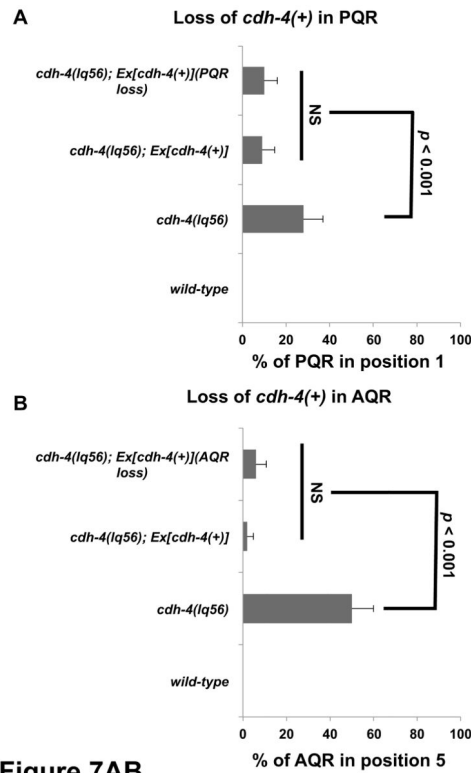
Images are micrographs of L1 animals with transgenic expression of the *cdh-4::gfp* recombinereered fosmid (green) and the *Pegl-17::mCherry* transgene expressed in the Q neuroblasts (red). The scale bar in A represents 103m for AC, and in C for C–E. A) *cdh-4::gfp* expression in the Q cells an L1 animal 3–3.5 h post-hatch. Neighboring cells expressing *cdh-4::gfp* can also be seen (asterisks in C). B) The same animal in A showing *Pegl-17::mCherry* expression in the Q cells. C) A merge of images in A and B. (C–E) A lower magnification image of expression in an L1 animal at 4–4.5 h post-hatch with the same parameters as A–C. *cdh-4::gfp* is expressed along the length of the animal (arrows in F), including in the Q cells, which have divided.



**Figure 6. Mosaic analysis of *cdh-4* function**

A) A simplified representation of the *C. elegans* cell lineage showing the origins of all *Pgcy-32::gfp*-expressing cells (AQR, PQR, URX L/R). Not all cell divisions are shown. B) A diagram of an animal subdivided into five segments along its body is shown. The wild-type positions of AQR (QR descendent) and PQR (QL descendent) are indicated in the diagram in positions 1 and 5 respectively. C–D) Micrographs of an early L2 larval animal with stably-integrated *Pgcy-32::cfp* expression to mark the positions of AQR and PQR, as well as an unstable extrachromosomal array with rescuing *cdh-4::gfp* and *Pgcy-32::yfp* to identify genetic mosaics of this array in AQR and PQR. C) *Pgcy-32::cfp* expression shows that AQR is in its normal position 1, and PQR is in its normal position 5. D) *Pgcy-32::yfp*

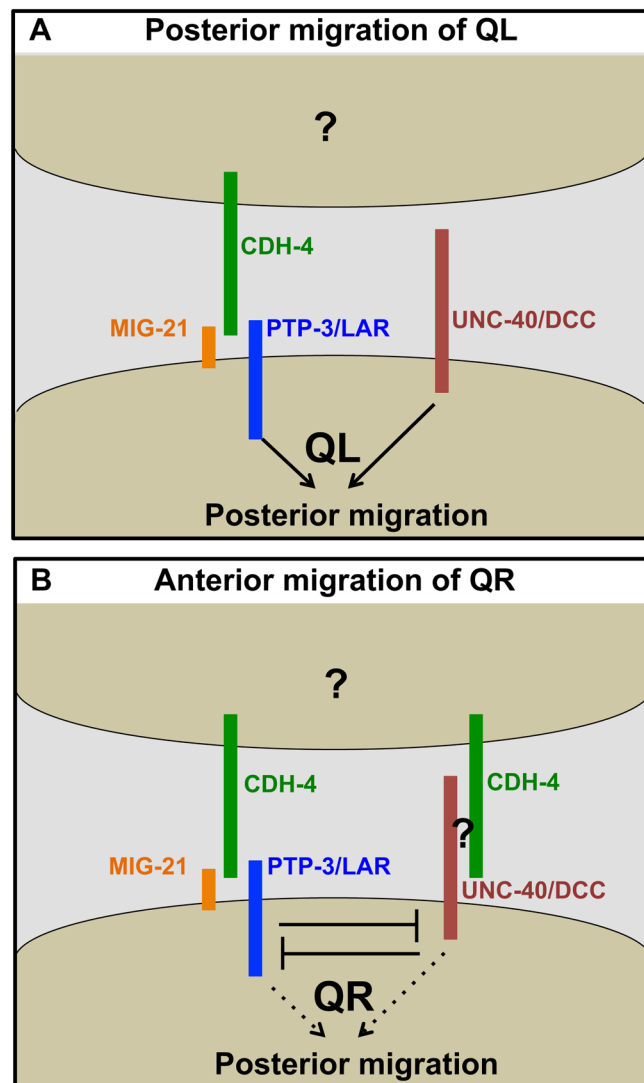
expression shows that the rescuing *cdh-4::gfp* array has been lost in AQR, but retained in PQR and the URX neurons. Despite loss of the array in AQR, it has migrated normally as judged by *Pgcy-32::cfp* expression in C.



**Figure 7AB**

**Figure 7. Quantification of AQR and PQR migration in *cdh-4* genetic mosaics**

Graphs plot the percentage of AQR or PQR migration defects (Y-axis) in different genotypes (X-axis). Error bars represent 2x SEP, and at least 100 animals of each genotype were scored. Statistical significance was determined by Fisher's exact test. A) PQR migration defects in animals that lost the rescuing *cdh-4::gfp* array in PQR (PQR loss) but retained the array in AQR and URX L/R. Animals with and without the array in PQR showed reduced defects compared to *cdh-4(lq56)* alone that were not significantly different from one another. B) AQR migration defects in animals that lost the array in AQR (AQR loss) but retained it in PQR and URX L/R. Animals with and without the array in AQR showed reduced defects compared to *cdh-4(lq56)* alone that were not significantly different from one another.

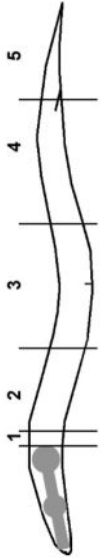


**Figure 8. A model of CDH-4 genetic interaction with UNC-40/DCC, PTP-3/LAR, and MIG-21 that is consistent with data described in this work**

A) CDH-4 acts non-autonomously with PTP-3/LAR and MIG-21 in parallel to UNC-40/DCC in directing posterior QL migration. Grouping of molecules does not imply physical interaction in this model, rather action in the same genetic pathway. The non-autonomous cellular source of CDH-4 is unknown (?). B) CDH-4 acts in both the PTP-3/LAR and UNC-40/DCC pathways in QR in mutual inhibition, allowing anterior QR migration. The QR-specific factor that mediates UNC-40/DCC function with PTP-3/LAR in QR but not QL is unknown (?).

Table 1

The Fat-like cadherin CDH-4 controls AQR and PQR migration



Genotype	AQR position (%)					PQR position (%)					
	1	2	3	4	5	1	2	3	4	5	N
<i>wild-type</i>	100	0	0	0	0	100	0	0	0	0	100
<i>cdh-4(lq56)</i>	39	3	2	7	49	100	28	3	4	4	60
<i>cdh-4(lq56); Ex[cdh-4(+)]</i>	90*	0	1	0	9	100	2*	0	1	1	96
<i>cdh-4(lq83)</i>	48	2	0	3	47	100	23	0	0	2	75
<i>cdh-4(lq97)</i>	80**	3	1	0	16	205	19	2	1	1	77
<i>cdh-4(lq97); smg-1(RNAi)</i>	82	3	1	0	14	200	13	1	1	1	84***
<i>cdh-4(rh310)</i>	50	1	1	2	46	100	19	0	0	2	79
<i>cdh-4(ok1323)</i>	45	0	2	1	52	100	24	0	1	0	75
<i>cdh-4(hd40)</i>	42	1	1	2	54	100	20	0	0	1	80
<i>cdh-4(RNAi)</i>	100	0	0	0	0	100	0	0	0	0	100
<i>unc-40(n324)</i>	96	1	0	0	3	76	32	3	8	6	51
<i>unc-40(n324); cdh-4(RNAi)</i>	99	0	0	1	0	100	76	4	2	5	13****
<i>egl-20(n585)</i>											
<i>cwn-2(ok895); cwn-1(ok546)</i>	39	49	10	2	0	100	3	16	19	25	37

\*  $p < 0.0001$  compared to other *cdh-4(lq56)*.\*\*  $p < 0.001$  compared to other *cdh-4* alleles *lq56*, *lq83*, *rh310*, *ok1323*, and *hd40*.\*\*\*  $p = 0.0416$  compared to *cdh-4(lq97)* alone.\*\*\*\*  $p < 0.0001$  compared to *unc-40(n324)* alone.