PCP Signaling between Migrating Neurons and their Planar-Polarized Neuroepithelial Environment Controls Filopodial Dynamics and Directional Migration

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Abstract

The planar cell polarity (PCP) pathway is a cell-contact mediated mechanism for transmitting polarity information between neighboring cells. PCP “core components” (Vangl, Fz, Pk, Dsh, and Celsr) are essential for a number of cell migratory events including the posterior migration of facial branchiomotor neurons (FBMNs) in the plane of the hindbrain neuroepithelium in zebrafish and mice. While the mechanism by which PCP signaling polarizes static epithelial cells is well understood, how PCP signaling controls highly dynamic processes like neuronal migration remains an important outstanding question given that PCP components have been implicated in a range of directed cell movements, particularly during vertebrate development. Here, by systematically disrupting PCP signaling in a rhombo-mere-restricted manner we show that PCP signaling is required both within FBMNs and the hindbrain rhombomere 4 environment at the time when they initiate their migration. Correspondingly, we demonstrate planar polarized localization of PCP core components Vangl2 and Fzd3a in the hindbrain neuroepithelium, and transient localization of Vangl2 at the tips of retracting FBMN filopodia. Using high-resolution timelapse imaging of FBMNs in genetic chimeras we uncover opposing cell-autonomous and non-cell-autonomous functions for Fzd3a and Vangl2 in regulating FBMN protrusive activity. Within FBMNs, Fzd3a is required to stabilize filopodia while Vangl2 has an antagonistic, destabilizing role. However, in the migratory environment Fzd3a acts to destabilize FBMN filopodia while Vangl2 has a stabilizing role. Together, our findings suggest a model in which PCP signaling between the planar polarized neuroepithelial environment and FBMNs directs migration by the selective stabilization of FBMN filopodia.
Author Summary

Planar cell polarity (PCP) is a common feature of many animal tissues. This type of polarity is most obvious in cells that are organized into epithelial sheets, where PCP signaling components act to orient cells in the plane of the tissue. Although, PCP is best understood for its function in polarizing stable epithelia, PCP is also required for the dynamic process of cell migration in animal development and disease. The goal of this study was to determine how PCP functions to control cell migration. We used the migration of facial branchiomotor neurons in the zebrafish hindbrain, which requires almost the entire suite of PCP core components, to address this question. We present evidence that PCP signaling within migrating neurons, and between migrating neurons and cells of their migratory environment promote migration by regulating filopodial dynamics. Our results suggest that broadly conserved interactions between PCP components control the cytoskeleton in motile cells and non-motile epithelia alike.

Introduction

The Planar Cell Polarity (PCP) signaling pathway is best understood as a cell contact dependent mechanism for generating and maintaining polarity in the plane of an epithelium [1, 2]. Its function was first described in the static epithelial cells of the fly where the molecular asymmetry of “core” PCP proteins results in the morphological asymmetry of a single actin-rich hair at the distal side of each wing cell [3–5]. Subsequently, planar polarity established by the core pathway has been shown to be a characteristic of many epithelial tissues in vertebrates and invertebrates alike [6–10]. The core PCP pathway is comprised of two protein complexes that localize to distinct cell membranes. In the fly wing, the transmembrane protein Frizzled (Fz) is confined to distal apical cell junctions along with the cytosolic proteins Disheveled (Dsh) and Diego (Dgo), while the transmembrane protein Van Gogh (Vang) (Strabismus(Stbm)) and the cytosolic protein Prickle (Pk) are proximally localized. This molecular asymmetry of PCP promotes actin polymerization at the distal side of the cell, downstream of Fz and Dsh [11–13]. While the factors that initially polarize PCP components are context dependent [14], the asymmetric localization of PCP proteins is maintained within polarized cells via intracellular destabilizing interactions between the Vang complex and the Fz complex [15, 16]. This polarization of PCP proteins is coordinated between cells by the formation of intercellular stabilizing interactions between Vang and Fz complexes across cell junctions [17–21]. In spite of the antagonistic roles of Vang and Fz complexes, loss of function of any core PCP component results in a loss of polarity.

While PCP is well known for its role in stable epithelia [22–24], core PCP components have also been implicated in dynamic cellular processes such as cell migration. How PCP controls directed cell movements is best, though incompletely, understood in coherently migrating cells such as those undergoing convergent extension [25–37]. However, independently migrating cells also require PCP [38–44]. Here, as our model we use the stereotyped and conserved migration of cranial motor neurons in the vertebrate hindbrain [45–47]. This enabled us to study in vivo how PCP can regulate the migration of non-coherent cells and to determine how PCP signaling between different cell types, the migrating neurons and the cells through which they migrate, can modulate migratory cell behaviors.

The PCP pathway drives the stereotyped tangential migration of facial branchiomotor neurons (FBMNs) in the vertebrate hindbrain. FBMNs are a subset of cranial branchiomotor neurons that originate ventrally in rhombomere (r)4 and undergo a posterior migration to r6.
where they form the facial motor nucleus, whose axons exit the hindbrain in r4 and innervate muscles derived from the second branchial arch [45, 47]. Forward genetic screens in the zebrafish have identified multiple core PCP components (Vangl2, Pk1b, Fzd3a, Celsr2 and Scribble) as being required for FBMN migration [31, 48–51]; this PCP requirement has also been shown for mouse FBMN migration [52–54]. Unlike the cell migrations mentioned above, screens have failed to identify a role for Wnts or other chemotactic cues. Although it is clear that many components of the PCP pathway are required for tangential FBMN migration, how these components regulate this highly dynamic process is unknown.

As a first step in answering this question we defined the cell types participating in PCP signaling during FBMN migration, as previous studies using a range of approaches have yielded conflicting results [31, 48, 49, 51, 55]. Using the Gal4/UAS system to systematically disrupt PCP in a cell-type and rhombomere-specific manner, we demonstrate the dual requirement for PCP within FBMNs and the planar-polarized r4 neuroepithelial environment in which they arise, and identify reciprocal PCP-dependent interactions between FBMNs and the planar-polarized floorplate as being sufficient, though not required, to promote migration. Since cell migration results from the contact-dependent stabilization of cellular protrusions and PCP signaling is known to regulate actin dynamics, we examined the protrusive activity of single FBMNs using high-resolution single-cell time-lapse microscopy in chimeric embryos and demonstrate opposing functions for the PCP core components Fzd3a and Vangl2 in regulating FBMN filopodial protrusive activity in vivo. Within FBMNs we show that Fzd3a is required to stabilize filopodia while Vangl2 has an antagonistic, destabilizing role. However, in the migratory environment we show that Fzd3a is required to destabilize filopodia while Vangl2 has a stabilizing role. In spite of having antagonistic roles at the cellular level, Vangl2 and Fzd3a mutants have the same FBMN migration phenotype. These findings are thus reminiscent of the intracellular antagonistic versus intercellular stabilizing roles that core PCP proteins perform in stably polarized epithelia. Consistent with a role for Vangl2 in regulating filopodial dynamics, we show that Vangl2 localizes transiently to the tips of retracting FBMN filopodia; consistent with a role for Vangl2 and Fzd3a in the microenvironment, we show planar polarized localization of these proteins in the adjacent floorplate. Together, our findings support a model in which canonical interactions between PCP components within FBMNs and between the FBMNs and their planar polarized neuroepithelial environment promotes migration via the selective stabilization of FBMN filopodia.

**Results**

**PCP signaling within FBMNs is required for their migration**

Initial chimeric analyses suggested that the PCP components Vangl2, Fzd3a, Celsr2 and Scrib primarily act non-cell-autonomously to regulate FBMN migration [31, 48, 49]. An additional cell-autonomous role for Vangl2 and Scrib in FBMN migration has been demonstrated [51], but refuted by others [55]. To determine whether PCP signaling is required cell-autonomously within FBMNs for their migration, we expressed a dominant negative (DN) form of the PCP core component Dvl specifically in branchiomotor neurons using the islet-1 (isl1) CREST enhancer (Fig 1A, 1B and 1C) [56]. Dvl is the branching point between multiple Wnt signaling pathways, and the overexpression of its individual domains exert pathway-specific DN properties [57]. Work in multiple vertebrate systems has demonstrated that Xdd1 and Dvl-DEP, two truncated forms of Dvl, act as PCP-specific DNs [25–27].

In previous studies mRNA injection of these DNs failed to disrupt zebrafish FBMN migration [31, 54]. We reasoned that this could be due to decreased DN mRNA levels or activity by the time of FBMN migration. To stably express DN forms of Dvl in FBMNs we raised stable Tg
Fig 1. PCP signaling is required within FBMNs and in their r4 environment. (A) Schematic showing a dorsal view of a 48 hours post fertilization (hpf) zebrafish hindbrain with anterior to the top. Facial Branchiomotor neurons (FBMNs) (green) migrate posteriorly from rhombomere (r) 4 to r6, leaving a trailing axon that exits from r4. The enhancer element *islet-1* (*isl1*) CREST drives expression in branchiomotor neurons (green); the *hoxb1a* element drives expression in r4 (light blue); *egr2b* drives expression in r3 and r5 (yellow) and *shh* drives expression in the floorplate (purple). (B,C, G-L) Live or (H) fixed confocal images showing dorsal views of the hindbrain of 48 hpf zebrafish embryos with anterior to the top. Brackets mark rhombomere (r) position. Scale
Bar: 50μm (B) Tg isl1:GFP) expression in a wild type embryo at 48 hpf. (C) Tg isl1:Dvl-DEP-GFP) embryo with unmigrated Dvl-DEP-GFP-expressing FBMNs in r4. (D,E) Dorsal view of E12.5 mouse hindbrains with FBMNs (magenta) labeled with anti-Isl1 antibody. Dotted lines indicate the length of facial motor nucleus. Scale Bar: 100μm (D) Migrating FBMNs in Tg isl1:GFP; isl1 Cre control embryos. N = 6 embryos. (E) Blocked FBMNs in Tg isl1:GFP; isl1 Cre embryos. N = 9 embryos. (F) Quantitation of FBMN migration stream length in Tg isl1:GFP; isl1 Cre control embryos and Tg isl1:GFP; isl1 Cre embryos. ***p = 0.0003. Significance was determined using an unpaired, two-tail t-test. (G-L) FBMNs (magenta) are either expressing Tg isl1:mRFP) (G-L) or are stained with anti-Isl1 (H). (G,H) Tg isl1:GFP; isl1 Cre-driven expression of Tg isl1:GFP) (H) and Tg isl1:Cre-mRFP) (H), throughout r3 and r5 does not block FBMN migration. (I,J) Tg rl1b:Gal4)-driven expression of Tg isl1:GFP) (I) and Tg isl1:Cre-mRFP) (J), throughout r4 blocks FBMN migration out of r4. (K,L) Chimeric embryos with transplant conditions indicated as donor-host. Cascade blue-dextran marks all donor-derived cells (blue) and Tg isl1:GFP) marks all donor-derived FBMNs (magenta). (K) Wild type donor-derived FBMNs migrate normally in a non-transgenic control host. N = 37 embryos, 378 FBMNs. (L) Wild type donor-derived FBMNs fail to migrate out of r4 that is expressing Tg isl1:GFP). N = 26 embryos, 190 FBMNs. Inset: same image without the magenta channel showing that donor-derived FBMNs (blue, circled) are not themselves expressing isl1-GFP (green). (M) Histograms indicate the percent of donor-derived FBMNs at 48 hpf that failed to migrate (r4), migrated partially (r5) or migrated fully (r6). Each histogram corresponds to the chimeric condition in the image to its left and numbers indicate the number of FBMNs represented in each bar.

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FBMN migration requires PCP signaling non-autonomously in r4

While these data support a cell-autonomous requirement for PCP signaling in FBMN migration, PCP signaling in FBMNs is not sufficient for their migration. Indeed, a non-autonomous requirement for PCP signaling in FBMN migration has been well established in chimeras in which wild type FBMNs are unable to migrate in FBMN, fzd3a, celsr2 or scrib mutant hosts [31, 48, 49, 51]. Since PCP is a cell-contact mediated signaling pathway in which the same trans-membrane protein components are required in both contacting cells [2], an attractive
hypothesis is that FBMNs receive PCP cues from cells in their environment that promote or
direct their migration. Thus we sought to determine where PCP signaling is required in the
FBMN migratory path for migration.

To block PCP signaling in distinct compartments of the hindbrain, we used the Gal4/UAS
system to drive rhombomere-restricted expression of Tg(UAS:Xdd1-GFP) as well as a C-termi-
nally truncated Fzd3a Tg(UAS:Fzd3aΔC-GFP) that lacks its cytoplasmic region, which has been
shown to function as a potent PCP DN tool in zebrafish [49]. We used Tg(egr2b:KalTA4) to
drive expression throughout r3 and r5 starting at 12 hpf [63] and Tg(hoxb1a:Gal4) [64] to
drive expression throughout r4 starting at 10 hpf (Fig 1A). Expression of Xdd1-GFP or
Fzd3aΔC-GFP along the migration path in the r5 neuroepithelium does not affect migration
(Fig 1G and 1H). In contrast, r4-restricted expression of Xdd1-GFP or FzdΔC-GFP completely
blocks FBMN migration (Fig 1I and 1J). This suggests that PCP signaling is required at the
onset, but not throughout the course of FBMN migration. However we note that r5 expression
of Xdd1-GFP or Fzd3aΔC-GFP using egr2b:KalTA4 comes on slightly later than r4 expression
using hoxb1a:Gal4 (12 hpf compared to 10 hpf), so the caveat remains that PCP signaling is
not fully disrupted in r5 at the time of migration with the available tools.

It was not surprising that FBMNs fail to migrate in Tg(hoxb1a:Gal4); Tg(UAS:DN-GFP)
embryos given that FBMNs arise in r4, and thus express hoxb1a throughout their early devel-
opment, and we had already shown a cell-autonomous requirement for PCP signaling within
FBMNs. To assess whether PCP signaling plays a role in the r4 neuroepithelium outside of
FBMNs, we transplanted wild type Tg(isl1:mRFP) donor FBMNs into the presumptive ventral
hindbrain of Tg(hoxb1a:Gal4); Tg(UAS:Xdd1-GFP) embryos and assessed the positions of
donor-derived FBMNs at 48 hpf. In control hosts, 87% (328/378) of wild type donor-derived
FBMNs migrated out of r4. In contrast, in hosts expressing Xdd1-GFP in r4, only 17% (33/190)
of donor-derived wild type FBMNs migrate out of r4 (Fig 1K, 1L and 1M). Thus, expression of
Xdd1-GFP throughout r4 significantly hinders wild type FBMNs from initiating migration
(p<0.0001, χ² = 207.8) (Fig 1M). This demonstrates that there is a non-autonomous require-
ment for PCP signaling for FBMN migration in r4.

Environmental PCP signaling is required for the migration of post-mitotic
FBMNs

Having established that PCP signaling is required both within FBMNs and their r4 neuroe-
phithelial environment for migration to occur; we asked when this signaling is required. PCP sig-
naling polarizes neuroepithelial progenitors before FBMNs differentiate [7, 65, 66]. It is
possible that this early neuroepithelial polarity is maintained in FBMNs to orient their initial
migration. Alternatively, PCP signaling active at the time of migration initiation may promote
migration. We reasoned that in the former case a planar polarized environment would not be
required for migration after FBMNs had differentiated while in the latter case PCP function in
the r4 environment would continue to be essential for migration.

To determine when PCP signaling is required for FBMN migration, we transplanted a small
number (1–5) of pre-migratory but post-mitotic FBMNs directly from r4 of a Tg(isl1:GFP)
donor into r4 of a stage-matched wild type or vangl2m209 mutant Tg(isl1:mRFP) host (Fig 2A).
During this extraction procedure, transplanted cells round up and become separated in the
transplant pipette and are unlikely to retain polarity information. Nevertheless, 28% (48/174)
of surviving post-mitotic FBMNs transplanted into a wild type host r4 migrated to r6 (Fig 2B
and 2D). To rule out the possibility that the transplanted FBMNs are simply being carried by
migrating host FBMNs, we transplanted post-mitotic FBMNs into a pklb mutant host, which
has normal neuroepithelial PCP but no host FBMN migration; in this environment 11%, (17/
of transplanted FBMNs migrated (S2 Fig). This suggests that host neurons do contribute to migration [51, 58], but that post-mitotic transplanted FBMNs can migrate without contribution from migrating host neurons. Importantly, 0% (0/73) of FBMNs migrated to r6 after being transplanted into a \textit{vangl2} mutant host (Fig 2C and 2D). Together, these results suggest that post-mitotic FBMNs engage PCP signaling as they initiate their migration out of r4.

**Floorplate PCP is not required for FBMN migration**

FBMNs migrate in the ventral neural tube adjacent to the floorplate (Fig 1A, [49, 67]) making the floorplate a potential source of PCP signaling for FBMN migration. A recent report found that floorplate expression of Vangl2 is both necessary and sufficient for FBMN migration [55]. Here, to investigate whether PCP signaling in the floorplate is required for FBMN migration, we generated a \textit{Tg(shh:Gal4)} line (see Methods) to drive Xdd1-GFP or Fzd3a\Delta C-GFP expression in the notochord and floorplate (S3A, S3B and S3C Fig). In order to determine if dominant negative expression does indeed disrupt floorplate planar polarity, we quantified the anterior-posterior position of the basal body in single floorplate cells as the ratio of its distance from the anterior membrane to the full anterior-posterior cell length (S3G Fig). Basal bodies in wild type floorplate cells are planar polarized to the posterior membrane (average position = 78% of cell length, S3D and S3H Fig, [6]). Conversely, basal body planar polarization is significantly
disrupted in floorplate cells expressing Xdd1-GFP or Fzd3ΔC-GFP (average position = 63% and 59% of cell length respectively; S3E, S3F and S3H Fig). By comparison, floorplate cells in vangl2 mutants display a complete loss of basal body planar polarity (average position = 47% of cell length, [51]) (S3H Fig). With the caveat that this effect on floorplate planar polarity was scored after FBMN migration was complete (48 hpf) rather than at the onset of migration (18–24 hpf), DN expression in the floorplate in the floorplate had no effect on FBMN migration (Fig 3A and 3B). To confirm this, we specifically knocked Vangl2 out in the mouse floorplate using the floxed Vangl2 allele described above [61] and Shh-driven Cre recombinase [68]. We found that Cre-mediated deletion of Vangl2 in the mouse floorplate does not disrupt FBMN migration (S4 Fig). These results suggest that PCP signaling in the floorplate is not required for FBMN migration, and point to the possibility that loss of PCP in the floorplate can be compensated for by other planar polarized cells in the r4 neuroepithelial environment.

Floorplate PCP could nevertheless be sufficient to rescue FBMN migration as has been suggested [55]. We tested the sufficiency of Vangl2 in the floorplate for FBMN migration in two ways. We expressed a GFP-Vangl2 fusion protein specifically in the floorplate of vangl2 mutants and wild type siblings using stable Tg(shh:Gal4) driver and Tg(UAS:GFP-Vangl2) transgenic lines (vangl2m209/m209; Tg(shh:Gal4); Tg(UAS:GFP-Vangl2)). Although GFP-Vangl2 was expressed broadly in the floorplate in these otherwise mutant embryos starting at 14 hpf, and exhibits planar-polarized localization (S3A Fig and see below), it neither disrupted FBMN migration in a wild type embryo nor rescued migration in a vangl2 mutant embryo (Fig 3C and 3D). Since a caveat of this experiment is that Vangl2 over-expression can itself disrupt planar polarity, we used targeted transplantation of wild type cells into the floorplate of vangl2 mutants to test whether floorplate Vangl2 is sufficient to rescue FBMN migration. We never observed rescue of host FBMN migration in vangl2 mutant Tg(isl1:mRFP) hosts with wild type donor-derived cells in the hindbrain floorplate (Fig 3E). This includes 9 cases with 10 or more wild type floorplate cells in rhombomere 4. This is contrary to the findings of Sittarmane et al. (2013) [55] who found that a single wild type floorplate cell in r4 of a vangl2 mutant could rescue FBMN migration. Together, our findings suggest that Vangl2 function in the floorplate is not sufficient for FBMN migration.

In these transplant experiments we noted that FBMNs as well as floorplate cells differentiate from donor-derived cells. This is not unexpected, given the close proximity of floorplate and branchiomotor neuron progenitors in the early embryo [70]. Interestingly, we observed that unlike the mutant host FBMNs, wild type donor-derived FBMNs sometimes migrate (Fig 3F), and their ability to do so correlates with the number of wild type cells in the hindbrain floor plate (R² = 0.244; p = 0.005). We conclude that Vangl2 function in the floorplate is not sufficient for FBMN migration, but that Vangl2 function in the floor plate can support the migration of vangl2+ FBMNs in an otherwise vangl2 mutant neuroepithelium. Taken together, we conclude that the floorplate can serve as a source of PCP signals for FBMN migration, but other cells in the r4 environment, which are also planar polarized (see below) can compensate for the loss of normal floorplate PCP signaling.

### Vangl2 localization in the migratory environment

Thus far, we have shown that PCP signaling in FBMNs and their immediate neuroepithelial/floorplate r4 environment can drive migration. The localization of core PCP components is known to be crucial for many PCP mediated processes [2, 22]. Therefore, to better understand how PCP signaling might be used in neuronal migration we asked where PCP proteins localize within FBMNs and in their neuroepithelial microenvironment. Using a polyclonal antibody against zebrafish Vangl2, we observed localization of Vangl2 to cell membranes throughout the...
Fig 3. Floorplate PCP is neither required nor sufficient for FBMN migration but can support the migration of WT FBMNs. Confocal images showing dorsal views of 48 hpf hindbrains with anterior to the top. (A,B) Tg(shh:Gal4)-driven expression of Tg(UAS:Xdd1-GFP) (A) and Tg(UAS:Fzd3aΔC-GFP) (B) does not disrupt FBMN (magenta) migration. N = 13 Xdd1-GFP expressing embryos and 26 Fzd3aΔC-GFP expressing embryos. (C,D) Tg(shh:Gal4)-driven floorplate expression of GFP-Vangl2 (blue) in the floorplate of a wild type sibling does not disrupt FBMN migration (magenta) (C) and does not rescue migration in a vangl2 mutant (D). N = 24 vangl2 mutants with GFP-Vangl2 expression in the r4 floorplate, 14 with 5 or more expressing floorplate cells in r4. (C',D') Boxed regions from panels C and D respectively, showing a single z-plane where GFP-Vangl2 (blue) is expressed broadly in floorplate cells whose membranes are marked with the Zn5 antibody (yellow) [69]. (E,F) Genetic chimeras. Cascade blue-dextran marks all donor-derived cells (blue), Tg(isl1:mRFP) marks host FBMNs (magenta in E) and Tg(isl1:GFP) marks wild type donor-derived FBMNs (green in F). (E) The presence of wild type floorplate cells (blue) in a vangl2 mutant host embryo does not rescue the migration of host FBMNs. N = 16 embryos with extensive contribution of WT cells to the floorplate. (F) The presence of wild type floorplate cells (blue) in a vangl2 mutant can, however, support the migration of co-transplanted wild type donor derived FBMNs (green, arrows). N = 8 embryos with migrated donor-derived FBMNs/22 embryos with donor-derived FBMNs; N = 76 migrated FBMNs/383 total donor-derived FBMNs. (E',F') Single Z-planes of the boxed regions from panels E and F respectively, show that donor-derived cells (blue) are in the Zn-5-positive floorplate (yellow). Scale bars: 50 μm, 5μm in the insets.

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hindbrain neuroepithelium (S5 Fig). In the r4 floorplate, we noted a 1.6-fold enrichment of Vangl2 protein at anterior/posterior membranes of floorplate cells compared to their lateral membranes (Fig 4A and 4B). Co-staining with ZO1 shows that this staining is sub-apical, at the level of the tight junctions (Fig 4A'). In order to distinguish anterior from posterior membrane localization we mosaically expressed GFP-Vangl2 in the floorplate so we could visualize Vangl2 localization in isolated floorplate cells. This revealed that Vangl2 is specifically enriched at the anterior subapical membrane (Fig 4C and 4E). The normalized mean fluorescent intensity ratio of GFP-Vangl2 at the anterior membrane versus the posterior membrane in expressing floorplate cells is 2.2 (std. deviation 0.9; N = 29 cells in 8 embryos). Conversely, Fzd3a-GFP is enriched at the posterior membrane (Fig 4D). These findings for PCP protein localization in the floorplate are consistent both with the requirement for PCP core components in the posterior localization of the floor plate primary cilium [6], and with a conserved deployment of PCP core components in vertebrate and invertebrate epithelia.

While the regular organization of floorplate cells makes it easy to visualize their planar polarization, our findings suggest that the primary source of environmental PCP signaling in FBMN migration comes from neuroepithelial progenitor cells outside of the floorplate. Previous studies demonstrated a planar polarization of GFP-Pk and GFP-Vangl2 in neuroepithelial progenitor cells during zebrafish neurulation [7, 65], and of endogenous Vangl2 in the Xenopus neural plate [66]. We asked whether neuroepithelial progenitor cells display planar polarization of Vangl2 in r4 at the time of FBMN migration. Using the Tg(hoxb1a:Gal4) line we mosaically expressed GFP-Vangl2 and observed a subtle but significant asymmetry of GFP-Vangl2 to the anterior sub-apical side of r4 neural progenitors. While GFP-Vangl2 polarization is subtle and not detectable in all expressing neuroepithelial progenitors, in blinded experiments we were able to correctly guess the A-P orientation of embryos based exclusively on GFP-Vangl2 localization in r4 progenitors in 18/23 mosaically expressing embryos (p = 0.004 that 18/23 correct guesses were due to chance alone). The normalized mean fluorescent intensity ratio of GFP-Vangl2 at the anterior membrane versus the posterior membrane in cells where asymmetry is detectable is 1.82 (std. deviation 0.47, N = 17 embryos, 23 cells.) (Fig 4F). Thus both the r4 neuroepithelium and floorplate exhibit planar polarized Vangl2 localization at the time of FBMN migration.

Vangl2 is enriched at the tips of retracting FBMN filopodia

We next sought to determine where Vangl2 localizes in migrating FBMNs. Endogenous Vangl2 in FBMN membranes and the membranes of surrounding cells could not be resolved using the anti-Vangl2 antibody and, unlike static floorplate cells and neuroepithelial progenitors, FBMNS are highly dynamic, extending primarily filopodia-like protrusions as they migrate [51, 71]. Reasoning that Vangl2 localization would be similarly dynamic, we mosaically expressed GFP-Vangl2 in FBMNs and visualized localization using spinning disc time-lapse imaging. We found that GFP-Vangl2 localizes throughout the membrane as well as in putative cytoplasmic vesicles, as is predicted for a transmembrane protein (Fig 5A). However, in addition to its membrane localization, we observe transient enrichment of GFP-Vangl2 at the tips of a subset filopodia immediately preceding filopodia retraction (Fig 5A’–5C and S1 Movie). Before enrichment the mean fluorescent intensity ratio of GFP at the filopodia tip versus the filopodia base is approximately 1 (0.99 ± 0.01), as is the case for mRFP (background membrane marker) (0.92 ± 0.02). During the enrichment event, this ratio for GFP-Vangl2 increased to 1.31 ± 0.05 while the ratio for mRFP remained close to 1 (0.97 ± 0.02) (Fig 5B). Since the ratio for mRFP remained close to 1, this suggests that the enrichment of GFP-Vangl2 correlates with increased Vangl2 protein levels at filopodia tips and not simply condensation of the membrane
Fig 4. Polarization of PCP protein localization in the migratory environment. (A–A") Dorsal view with anterior to the top of a 24 hpf wild type floorplate at the level of r4 co-immunostained with anti-Vangl2 (green) and anti-ZO-1 (magenta), a marker of apical tight junctions. The boxed regions in A are examples of anterior-posterior membranes (AP) (61–90° from AP axis), intermediate membranes (I) (31–60° from AP axis) and lateral membranes (L) (0–30° from AP axis). Arrows in A" indicate enrichment of anti-Vangl2 labeling at AP membranes. (B) Quantitation of fluorescent intensity of anti-Vangl2 labeling for AP, I and L membranes. N = 5 embryos, 192 membranes (57 L, 47 I, 88 AP). Graph represents data as mean ± SEM. *p = 0.018, ****p < 0.0001; Significance was determined using a paired two-tail t-test with Welch’s correction. (C–D) Live lateral views of 48 hpf wild type floorplate cells at the level of the spinal cord with mosaic expression of GFP-Vangl2 (C) and Fzd3a-GFP (D). Anterior is to the left and dorsal/apical is up; white dots indicate the center of each expressing
due to retraction. Furthermore, as described below, addition of exogenous GFP-Vangl2 in FBMs results in a reduced filopodial lifetime which is opposite to the effect of loss of Vangl2 in FBMs. This suggests that exogenous GFP-Vangl2 is functioning in FBMs and that this observed localization of Vangl2 at the tips of filopodia is correlated with retraction. This enrichment of GFP-Vangl2 in filopodia never lasted for more than one time-point (images were taken at 30–45 second intervals) and was only detected in a subset of filopodia (N = 11/84 filopodia on 8 neurons in 7 embryos); it is likely that due to the transient nature of enrichment events and the constraints of our imaging rate we failed to observe many enrichment events. Importantly, however, the enrichment events we captured invariably preceded filopodial

Fig 5. GFP-Vangl2 is enriched to the tip of retracting FBMN filopodia. (A) Live confocal image of a single GFP-Vangl2 expressing FBMN (green) in a Tg isl1:mRFP (magenta) 24 hpf embryo. Scale bar: 10 μm. (A’,A”) Magnified views of the boxed region in A of the individual channels, GFP-Vangl2 and Tg isl1:mRFP respectively, at the time points indicated. The arrow in E’ indicates enrichment of GFP-Vangl2 at the filopodial tip. Scale bar: 10 μm. (B) Quantitation of filopodia tip/base mean fluorescent intensity ratio for mRFP and GFP-Vangl2 at the time-point before and during GFP-Vangl2 enrichment. Before enrichment the mean fluorescent intensity ratio of GFP and mRFP at the filopodia tip versus the filopodia base is approximately 1 (N = 9 filopodia). During the enrichment event this ratio for GFP-Vangl2 is 1.31 while the ratio remains close to 1 for mRFP (N = 12 filopodia). (C) Plot showing the change in filopodial length over time for 10 filopodia. The stars indicate the time-point that GFP-Vangl2 is enriched at each filopodium tip. The black trace corresponds to the filopodium in A’,A”. Graph represents data as mean ± SEM. ***p<0.001; Significance was determined using an unpaired, two-tail t-test with Welch’s correction.
retraction; filopodia never extended further after an enrichment event (Fig 5C). Consequently, we infer that Vangl2 may function in FB MN filopodia to signal retraction events.

**Vangl2 and Fzd3a function cell-autonomously to regulate FB MN filopodial activity in an antagonistic manner**

Our findings that PCP signaling is required within FB MNs for migration, and that Vangl2 localizes transiently to the tips of retracting filopodia, suggested the possibility that PCP signaling influences filopodial dynamics in migrating neurons in vivo. In order to determine the cellular basis of FB MN migration defects in PCP mutants, we imaged the protrusive dynamics of single mutant FB MNs at high resolution in vivo. Previous studies have described membrane protrusions in fixed or live embryos expressing cytoplasmic GFP or membrane-RFP in bulk FB MNs at low time resolution, however the overlap between FB MNs allows only a subset of protrusions to be visualized and their dynamics could only be inferred from distant time points [31, 55, 58, 71]. To visualize the protrusive activity of single FB MNs at high time resolution, we utilized cell transplantation to generate embryos in which one or a few FB MNs express membrane-localized teal fluorescent protein (Tg isl1:mTFP), and imaged protrusion dynamics of single FB MNs at 30-second intervals, the shortest interval at which we could acquire comprehensive z-stacks on our instruments. We focused on the function of Vangl2 and Fzd3a, the mutually antagonistic transmembrane core components, whose localized activity is both the hallmark and the driver of classical epithelial planar polarity [2].

Time-lapse imaging of FB MN membranes and f-actin dynamics revealed that filopodia are the prevalent protrusion type in FB MNs (Fig 6A–6E and S2 Movie). To determine whether PCP affects the polarized orientation of filopodia on migrating cells, we quantified the positions of filopodia on single isl1:mTFP-expressing FB MNs by measuring the angle from the anterior–posterior axis of a vector from the center of the cell to the base of each filopodium (S6A Fig). When wild type FB MNs are in r4, protrusive activity is largely radial 46.4% (13/28) of filopodia are located in the anterior half of the neuron, while 53.6% (15/28) of filopodia were on the posterior side (S6A and S6B Fig). Once neurons are migrating through r5 and r6, membrane protrusive activity becomes highly enriched posteriorly, in the direction of migration with 84.6% (44/52) of filopodia on the posterior side of the cell (S6C and S6D Fig). However, filopodia on vangl2 mutant FB MNs fail to polarize. Time-lapse images were collected at a developmental time-point at which wild type neurons would have already migrated out of r4. Similar to wild type neurons in r4, filopodia in vangl2 mutant FB MNs are fairly evenly distributed along the anterior–posterior axis with 47.5% (29/61) of filopodia located anteriorly and 52.5% (32/61) located posteriorly (S6E and S6F Fig). These findings suggest that PCP signaling through Vangl2 is required to properly localize cytoskeletal dynamics in FB MNs.

To characterize protrusive membrane dynamics we quantified filopodial lifetime (number of seconds each filopodium is present during a 15 minute time-lapse period) and filopodial maximum length (the greatest length during the lifetime of filopodia lasting 90 seconds or longer) of Tg isl1:mTFP) FB MNs. Wild type FB MNs generate filopodia with an average lifetime of 224.5 ± 18.66 (SEM) seconds and an average maximum length of 3.6 ± 0.3 μm (Fig 6A, 6F and 6G and S3 Movie). When compared to wild type, FB MNs in vangl2 mutant embryos have much more stable filopodia with a longer average lifetime of 537.3 ± 81.78 seconds (Fig 6F, S4 Movie; p = 0.0059). Filopodia of these vangl2 mutant FB MNs also reach a greater average maximum length of 6.4 ± 1.1 μm (Fig 6G; p = 0.0406). We saw a similar trend when we used microinjection rather than transplantation to mosaically express
Fig 6. Vangl2 and Fzd3a have opposing cell-autonomous and non-cell-autonomous roles in modulating filopodial dynamics. (A-E) Time-lapse spinning-disc confocal series of donor-derived FBMNs in chimeric embryos at 24–30 hours post-fertilization (hpf). Transplant conditions are indicated on the left as donor—host. Colored arrows indicate individual filopodia at different time-points. Anterior is to the top and medial is to the right. Scale bar: 5 μm. (F) Quantitation of filopodial lifetime for donor-derived FBMNs. Each data point is an average of all the filopodial lifetimes for one FBMN. (G) Quantitation of the maximum filopodial length for donor-derived FBMNs. Each data point is the average maximum length for all the filopodia of one FBMN. WT—WT: N = 6
mTFP in FBMs in wild type and vangl2 mutant embryos. These results suggest that Vangl2 is required to destabilize FBMN membrane protrusions. Previous studies have demonstrated that FBMs in vangl2 mutants move more slowly than wild type FBMs and in random directions, which is consistent with Vangl2 being required to destabilize protrusions [31].

Since Vangl2 is required within FBMs and their r4 microenvironment for migration, we sought to determine where Vangl2 functions to regulate filopodial dynamics. To determine the cell-autonomous function of Vangl2, we transplanted vangl2 mutant FBMs into a wild type host. Donor embryos carried the Tg(isl1:mTFP) transgene to visualize FBMs and contained rhodamine dextran to track other donor-derived cells so we could ensure that donor-derived FBMs were in fact in a genetically chimeric environment (S7 Fig). We found that vangl2 mutant FBMs in a wild type environment have longer, more stable filopodia with a mean lifetime of 432.0 ± 55.65 seconds and a maximum length of 6.8 ± 0.8 μm, similar to vangl2 mutant FBMs in a vangl2 mutant host (Fig 6B, 6F and 6G and S5 Movie; p = 0.005 and p = 0.0078 respectively). To further test if Vangl2 functions cell-autonomously to control FBMN protrusive dynamics, we mosaically expressed GFP-Vangl2 in wild type embryos. FBMs expressing GFP-Vangl2 in wild type embryos have less stable filopodia compared to wild type FBMs, with an average lifetime of 123.4 ± 14.27 seconds (S1 Movie; N = 6 embryos, 7 neurons, 42 filopodia, p = 0.0013). Together, these loss- and gain-of-function findings suggest that Vangl2 functions within FBMs to destabilize filopodia, since filopodia are affected in vangl2 mutant and GFP-Vangl2-expressing FBMs regardless of the genotype of cells in their microenvironment.

Fzd3a, like Vangl2, is required cell-autonomously and cell non-autonomously for FBMN migration (S1 Fig,[ 49]). To determine whether Fzd3a has a cell-autonomous role in FBMN protrusive activity, we transplanted fzd3a mutant FBMs into a wild type host. We found that filopodia of fzd3a mutant FBMs are significantly less stable than filopodia of wild type neurons, with a mean lifetime of 163.3 ± 8.006 seconds (Fig 6C and 6F and S6 movie; p = 0.005 and p = 0.0092). However, mean maximum filopodia length (3.4 ± 0.4 μm) was not significantly different than that of wild type FBMs (Fig 6G). This suggests that Fzd3a normally functions within FBMs to stabilize filopodia protrusions, consistent with a conserved role for Fzd in actin polymerization [11–13]. Taken together, our results suggest that Vangl2 and Fzd3a function antagonistically within FBMs to regulate filopodial stability.

Given that we observed this cell-autonomous function for Vangl2 and Fzd3a in regulating FBMN membrane protrusions, we asked whether these proteins regulate FBMN protrusive dynamics independently of cells in the migratory environment. To address this question, we analyzed the protrusive dynamics of isolated FBMs in primary culture. We found that cultured FBMs display altered filopodial dynamics compared to FBMs in vivo. Cultured wild type FBMs have a mean lifetime of 537.5 ± 32.28 seconds (during a 600 second time-lapse) and an average maximum length of 6.0 ± 0.5 μm (S9 Fig). Furthermore, there is no difference in filopodial dynamics between cultured wild type and cultured vangl2 mutant FBMs (S9 Fig). This suggested to us that the cell-autonomous functions we observe for Vangl2 and Fzd3a in vivo depend on interactions with cells in the migratory environment.
Vangl2 and Fzd3a have opposing cell non-autonomous functions in regulating FBMN filopodial activity

Since Fzd3a and Vangl2 are also required non-autonomously for FBMN migration and since FBMN protrusive dynamics depend on cells in the migratory environment, we asked whether cells in the FBMN environment influence FBMN protrusive activity in a PCP-dependent manner. In order to assess the role of Vangl2 in the environment, we imaged protrusion dynamics of wild type FBMNs in a vangl2 mutant environment. Interestingly we found that wild type FBMNs have less stable filopodia in a vangl2 mutant environment compared to a wild type environment, with a mean lifetime of 143.3 ± 18.28 seconds (Fig 6D and 6F and S7 Movie; p = 0.0056). The decrease in the average filopodia lifetime of wild type FBMNs in a vangl2 mutant environment is largely due to these neurons having a larger proportion of filopodia present for only one (30 seconds) or two (60 seconds) time-points (S8 Fig). The mean average length however was not different between wild-type FBMNs in a wild type environment and wild type FBMNs in a vangl2 mutant environment (Fig 6G) (3.1 ± 0.2 μm). Removing Fzd3a from the migratory environment had the opposite effect on FBMN filopodia. Wild type neurons in a fz3a mutant environment generate dramatically more stable filopodia compared to those in a wild type environment, with a mean lifetime of 363.8 ± 51.12 seconds (Fig 6E and 6F and S8 Movie; p = 0.0273). Together our results suggest that the core PCP components Vangl2 and Fzd3a antagonize each other’s activity to control filopodial dynamics during neuronal migration in vivo and they do so by functioning both within FBMNs and in cells in their migratory environment.

Discussion

Based on our findings, we propose a model for the role of PCP signaling in FBMN migration in which canonical interactions between the transmembrane PCP core components Vangl2 and Fzd3a regulate filopodial dynamics, thereby signaling and/or regulating adhesion for directional migration (Fig 7). We note that this model for filopodial dynamics is based on genetics and that our work does not elucidate the molecular nature of these interactions, which remain controversial even in the context of epithelial polarity [17, 76]. This model is consistent with 1) a dual cell-autonomous and cell-non-autonomous requirement for PCP core components, specifically for the transmembrane components Vangl2 and Fzd3a, in FBMNs and their rhombomere 4 environment for migration; (this work, [31, 49, 51]); 2) the cytoskeletal and conserved molecular planar polarization of the r4 neuroepithelial environment including the floorplate (this work, [6]); 3) the ability of the planar polarized floorplate to promote the migration of wild type but not mutant FBMNs; 4) the localization of Vangl2 to retracting FBMN filopodial tips; 5) the antagonistic cell-autonomous roles of Fzd3a and Vangl2 in FBMN filopodial stability and 6) the opposite roles of Fzd3a and Vangl2 in the FBMN environment on FBMN filopodial stability.

Mutual antagonism of Vangl2 and Fzd3a

Our in vivo observations of filopodial dynamics in genetic chimeras demonstrate an antagonistic intracellular relationship between Vangl2 and Fzd3a in migrating FBMNs that regulates the stability of filopodium-like protrusions. While occurring in the context of a highly dynamic structure, this antagonistic relationship of Vangl2 and Fzd3a is reminiscent of the situation in stably polarized epithelia, where mutual intracellular antagonism between Fzd and Vangl complexes sets up polarized actin dynamics within the cell, with Fzd activating actin polymerization distally and Vangl suppressing it proximally [11–13, 77, 78]. This conserved interaction
between Fzd promoting and Vangl suppressing actin growth may be common to other migratory cells. In metastatic breast cancer cells induced by stromal Wnt11-containing exosomes, Fzd6 and Vangl1 exhibit mutually exclusive localizations, with Fzd6 on the leading edge of cell protrusions and Vangl1 on non-protrusive cell surfaces, and knock-down of either protein decreases cell motility [44]. Similarly in migrating leukemia cells, Dvl-3 (part of the Fzd complex) localizes to the leading edge while Vangl2 localizes to the trailing edge [79]. During mesodermal and neuroectodermal convergence, mediolaterally oriented cell surfaces exhibit increased actomyosin contractility [33, 80] that correlates with the asymmetric localization of PCP components Dvl and Pk (part of the Vangl complex) [7, 81], suggestive of a conserved intracellular antagonism of these complexes mediating actin dynamics. In contrast, in commissural growth cones, Vangl2 promotes Fzd3-dependent outgrowth induced by diffusible Wnt5a by antagonizing a non-canonical inhibitory interaction between Dvl1 and Fzd3 identified in that context [41]. These examples show that core PCP components localize to discrete domains of moving cells and we have shown in vivo for the first time that this results in opposing effects on filopodial stability.

The role of filopodia in FBMN cell migration

Filopodia are commonly associated with the promotion of directed cell migration, although in some instances, axons and cells can achieve proper targeting and guidance without filopodia [82–84]. Due to their dynamics and long thin architecture, filopodia are capable of probing a
wide area around cells, and they can contain receptors for diverse diffusible and membrane-bound signals and extracellular matrix molecules [85]. Thus, it is thought that the primary function of filopodia is as “antennae” that cells use to sense their microenvironment to orient directed cell migration [86]. Indeed it has been demonstrated that elimination of filopodia in axon growth cones does not impair axon outgrowth, but instead impairs growth cone turning in response to environmental cues [87–89]. This sensing role for filopodia has also been demonstrated in cell migration [84, 90]. In addition to a sensing role, filopodia are thought to contribute directly to cell motility, as cells lacking filopodia tend to migrate more slowly due to the absence of filopodial adhesion molecules which could induce traction and also through force generated by actin streaming in filopodia [82, 90–93]. In the context of FBMN migration, filopodia extend in all directions from neurons when they initiate their migration, and we see no bias in the orientation of the filopodia that are affected in PCP mutants. We hypothesize that filopodia act as sensors of asymmetrically localized cell-surface PCP components on the neuroepithelial cells through which they are migrating and that this sensing fine tunes filopodium dynamics such that these filopodia can promote migration by acting as force generators or appropriately sensing other, as-yet unidentified environmental cues. In other migrating cells, several effectors have been identified as possible links between PCP signaling and cytoskeletal regulation [33, 94, 95]. While our work does not elucidate how those signals are transduced to the filopodial actin cytoskeleton in FBMNs, our previous work identified the WAVE-homology domain containing actin regulator Nhsl1 as localizing to FBMN filopodia and being required cell-autonomously for FBMN migration [51, 96]; we hypothesize that PCP signals may be transduced to the actin cytoskeleton in FBMNs via Nhsl1.

**Fzd3a and Vangl2 function in the migratory environment**

A more surprising finding than opposing cell autonomous roles for Fzd3a and Vangl2 in FBMN filopodial dynamics and migration is that the same PCP components function in the FBMN environment to influence filopodial dynamics but in the opposite way: Fzd3a in the environment destabilizes filopodia while Vangl2 in the environment stabilizes them. These non-autonomous functions for Fzd3a and Vangl2 in filopodial dynamics correlate with their non-autonomous roles in FBMN migration [31, 49]. Again, this is reminiscent of classical planar polarity, where localized Fzd activity depends on the presence of Vangl in adjacent cells in the epithelium and vice versa; this is the mechanism by which PCP is coordinated across an epithelium [17–21]. We hypothesize that the cell-autonomous activities of Fzd3a and Vangl2 are activated in different filopodia when they contact Vangl2 and Fzd3a domains of neuroepithelial cells in the r4 environment (Fig 7), with consequences on the actin dynamics regulating filopodial stability, leading to changes in signaling and/or adhesion. We have shown here that Vangl2 and Fzd3a exhibit planar polarized localization in the r4 neuroepithelium and floorplate at the time of FBMN migration. In PCP mutants, this polarized information is absent and/or cannot be correctly interpreted by filopodia resulting in a failure of directional cell migration. We note that the cell-autonomous filopodial phenotypes appear to be dominant, since in constitutive mutants filopodia have the cell-autonomous phenotype (long and stable in vangl2 mutants; unstable in fzd3a mutants). Together our findings suggest that conserved intracellular and intercellular interactions between PCP core components can have divergent effects on actin dynamics and consequently on cell behaviors.

While the similar effects on filopodial dynamics when Vangl2 is depleted from FBMNs and when Fzd3a is depleted from their environment suggest that the two proteins are working together, environmental PCP may also influence filopodia dynamics of FBMNs through an indirect mechanism. For instance, core PCP proteins have been implicated in the trafficking
and regulation of membrane levels of cadherins in fly and in vertebrate epithelial cells [97–99]. Therefore, Vangl2 and Fzd3a in the migratory environment may modulate FBMN filopodia dynamics by regulating N-cadherin levels at the surface of neuroepithelial cells. Another potential mechanism by which PCP in the migratory environment may regulate FBMN filopodial dynamics is through regulation of membrane type-1 matrix metalloproteinase (MMP14), which are known to degrade extracellular matrix proteins. During zebrafish gastrulation, an increase in Mmp14 activity was observed in vangl2 mutant embryos [100]. Thus, the decreased FBMN filopodial stability observed when Vangl2 is absent in the migratory environment could be due to decreased extracellular matrix.

Which cells in the environment of FBMNs are the source of PCP cues for filopodial dynamics and migration? We have shown that disruption of PCP signaling in the r4 environment prevents FBMN migration, demonstrating that PCP signaling is required to initiate directional migration. It was recently reported that Vangl2 expression even in a single cell in the r4 floorplate is sufficient to rescue FBMN migration in a vangl2 mutant [55]. In contrast, our results show that floorplate Vangl2 is neither required nor sufficient for FBMN migration. Neither the widespread presence of GFP-Vangl2 expressing cells or of wild type donor-derived cells in the floorplate of vangl2 mutants rescued the migration of vang2 mutant FBMNs. The rescue of FBMN migration observed by Sittaramane et al. (2013) may have been due to undetected early expression of Vangl2 outside of the floorplate driven by the Gal4 genetrap line used in their experiments [101, 102].

We did, however, note that the presence of wild type cells in the floorplate could partially rescue the migration of wild type FBMNs in an otherwise vangl2 mutant embryo. This suggests that bidirectional PCP signaling between the planar-polarized floorplate and the FBMNs can promote migration. However this rescue was incomplete, indicating that other planar polarized cells in the r4 environment normally contribute to the pro-migratory environment. Consistent with this hypothesis, we found that disrupting the planar polarization of the floorplate alone in both fish and mouse embryos was insufficient to prevent FBMN migration (Fig 3 and S4 Fig). We conclude that the planar polarization of the entire r4 environment surrounding the FBMNs is required to effectively initiate migration. We were unable to confirm this by rescuing FBMN migration in a vangl2 mutant with r4-restricted expression of GFP-Vangl2, likely because the over-expression of PCP components disrupts planar polarity as efficiently as their loss [28, 31, 103, 104].

Our study provides new insights into the role of the planar cell polarity pathway in neuronal migration by identifying when and where PCP signaling is required and how it affects the dynamic cell behaviors of migrating neurons in vivo. Our data suggests that a planar polarized hindbrain rhombomere 4 neuroepithelium serves to promote FBMN migration through the selective stabilization and destabilization of FBMN filopodia using conserved intra- and intercellular interactions between the PCP components Vangl2 and Fzd3a. Whether neuroepithelial planar polarity directs posterior migration or simply enables it, and through what effectors PCP signaling regulates filopodial dynamics in vivo are important questions to be answered in future work.

Methods

Ethics statement

Experiments using animals were performed under the Fred Hutchinson Cancer Research Center Institutional Animal Care and Use Committee protocols #1392 (zebrafish, approved on 3/31/2015) and #50857 (mice, approved 4/1/2015). The Fred Hutchinson Cancer Research Center Institutional Animal Care and Use Committee (IACUC) follow the guidelines of the Office
of Laboratory Animal Welfare and set its policies according to The Guide for the Care and Use of Laboratory Animals. Fred Hutchinson Cancer Research Center maintains full accreditation from the Association for Assessment and Accreditation of Laboratory Animal Care (AAALAC) and has letters of assurance on file with OLAW. The IACUC routinely evaluates the Fred Hutchinson animal facilities and programs to assure compliance with federal, state, local, and institution laws, regulations, and policies. The OLAW Assurance number is A3226-01.

Zebrafish lines and maintenance

Zebrafish (Danio rerio) were raised at the Fred Hutchinson Cancer Research Center, and animal care and experiments were approved by the Institutional Animal Care and Use Committee. All animals were maintained according to standard procedures [105] and staged as previously described [106]. All mutant lines used were previously described and are registered at The Zebrafish International Resource Center (ZIRC): fzd3a<sup>wel689</sup> (olt<sup>wel689</sup>) [49], prickle1B<sup>fh122</sup> [50], and vangl2<sup>m209</sup> (tri<sup>m209</sup>) [31]. Previously described transgenic lines used were as follows: Tg(isl1:GFPrw0) [56], Tg(isl1CREST-hsp70:mRFP)fh1 [67], TgBAC(hoxb1a:RFP)fh3 [67], Tg(egr2b:KalTA4) [63] and Tg(hoxb1a(β-globin):Gal4VP16)um60 [64].

Cloning and transgenic line generation

The following transgenic lines were generated for this study: Tg(shh:Gal4VP16)fh445, Tg(isl1:Gal4VP16)fh452, Tg(isl1-hsp70:mTFP)fh350, Tg(isl1-hsp70:del-DEP-GFP)fh444, Tg(10XUAS:xdd1-GFP)fh446, Tg(10XUAS:fxdAC-GFP)fh447 and Tg(10XUAS:GFP-vangl2)fh453. The Gal4VP16 sequence was obtained from the Nonet Lab (http://pcg.wustl.edu/nonetlab/ResourcesF/Zebrafish.html) and the 10XUAS plasmid was obtained from the Tol2 kit (http://tol2kit.genetics.utah.edu/index.php/List_of_entry_and_destination_vectors) [107]. The mTFP construct was obtained from Alleleustrious, Inc (Cat# ABP-FP-TFA1000).

To generate Tg(shh:Gal4VP16)fh445, the ar-B enhancer element of zebrafish sonic hedgehog (shh) [108, 109] was amplified from a plasmid (gift from Uwe Strähle). For the Gal4 lines, the shh and isl1 enhancers were inserted upstream of the gata2 minimal promoter element [110]. The Xdd1 and full-length Xenopus Dvl are described in Sokol et al. (1996) [25]. Transgenic elements were cloned using the Gateway (Life Technologies) system using the primer sequences listed in S1 Table. Final DNA constructs were assembled in the pDESTpBHR4R3 plasmid (gift from the Brockerhoff Lab) or the CG5 Tol2 expression vector [107]. Transgenic embryos were generated by Tol2 transposase RNA co-injection with each plasmid at the single cell stage [111].

Mouse lines and husbandry

All mice were maintained at Fred Hutchinson Cancer Research Center under Institutional Animal Care and Use Committee approved guidelines. For general colony maintenance, all mouse lines were crossed into the C57BL/6J background (The Jackson Laboratory strain 00064). The Vangl2<sup>LoxP</sup> and Vangl2<sup>XTM</sup> lines were a gift from the Deans laboratory [61], the Isl1Cre (Ist<sup>tm1(cre)Sev</sup>) line was a gift from the Evans laboratory [62] and the Shh:gfpc-cre (Shht<sup>tm1(BGFP/cre)Cjt</sup>) line was purchased from The Jackson Laboratory (strain 005622).

Cell transplantation

Chimeric embryos were generated by transplantation at the blastula or gastrula stage as previously described [51, 112]. To track transplanted cells, donor embryos carrying the Tg(isl1:GFP) rw0, Tg(isl1:mRFP)fh1 or Tg(isl1:mTFP)fh350 transgene were injected with 1% cascade blue-
dextran or rhodamine-dextran (for live imaging) and 1% biotin-dextran (for imaging after fixation) (10,000 mw, Life Technologies). Host embryos were then processed and imaged for all donor-derived cells, donor-derived FBMs or floorplate cells, and host FBMs. Host and donor embryo genotypes were identified either by observing body axis elongation defects (for vangl2 mutant hosts), by examining FBMN location at 48 hpf or by genotyping (for fz3a mutant hosts).

For transplantation of post-mitotic FBMs, cascade blue-dextran labeled donor embryos and unlabeled host embryos were mounted in agar on coverslips at the 15-somite stage. The head of each animal was exposed by careful removal of agar with insect pins, and a hole was cut in the skin overlying the forebrain to enable entry of a thin (10 μm diameter) transplant pipette. Pre-migratory FBMs (visualized by isl1:GFP or isl1:mRFP expression) were removed from r4 of a donor embryo and transplanted into r4 of a host embryo using a Zeiss AxioSkop fixed-stage microscope fitted with a 40X long working-distance water-immersion (“dipping”) lens. During this process some non-isl1:GFP/mRFP-expressing neuroepithelial progenitor cells were inevitably co-transplanted but these usually died shortly after transplantation; any surviving donor-derived cells that were not FBMs were detected by the presence of the cascade blue dye. Due to the disruptive approach, which removes nascent axons and other processes, even wild type FBMs transplanted into a wild type environment do not migrate as well as FBMs transplanted at earlier stages.

Whole-mount immunohistochemistry

Anesthetized zebrafish embryos were fixed in 2% trichloroacetic (TCA) acid for 3 hours or 4% paraformaldehyde (PFA)/4% sucrose in PBS for 1 hour at room temperature. Dissected mouse hindbrains were fixed in 4% paraformaldehyde (PFA)/4% sucrose in PBS overnight at 4°C and permeabilized in PBS + 1% TritonX100. Fixed tissue was washed in PBS + 0.5% TritonX100 followed by standard blocking and antibody incubations. Following staining, brain tissue was dissected, cleared step-wise in a 25%, 50%, 75% glycerol series and mounted for confocal imaging. The following antibodies were used: rabbit anti-zebrafish Vangl2 (1:250, Anaspec Cat# AS-55659), mouse anti-islet1 39.4D5 (1:10 for zebrafish tissue and 1:100 for mouse tissue, Developmental Studies Hybridoma Bank); chicken anti-GFP (1:500, Abcam Cat# ab13970); rabbit anti-ZO-1 (1:1000, Zymed Cat# 61-7300); mouse anti-Cc2d2a (1:100, [113]); rabbit anti-RFP (1:1000, Abcam Cat# ab62341). For analysis of chimeric embryos after fixation, host embryos were additionally stained with a fluorescently conjugated streptavidin (Life Technologies Cat# S32351) to enhance the detection of Biotin-Dextran-containing donor-derived cells.

Primary cell culture

Primary cultures of FBMs were prepared from 24 hour post fertilization Tg(isl1:mTFP); Tg(hoxb1a:RFP) embryos. The hindbrains of embryos were micro-dissected and dissociated as previously reported [114]. Cells were plated on a chambered coverglass (Sigma Z734756) coated with 5μg/mL poly-D-lysine (Sigma L8021) and 5μg/mL laminin (Sigma L2020) at a density of 4–5 hindbrains per 1.7 cm². FBMs were distinguished from other Tg(isl1:mTFP)-expressing hindbrain motor neurons by virtue of Tg(hoxb1a:RFP) expression, which is restricted to hindbrain r4 and r4-derived neurons. Live imaging of explanted neurons was performed 5 hours after plating.

Imaging and data analysis

Imaging was performed using a Zeiss 700 confocal microscope or a Zeiss spinning disc microscope with a QuantEM EMCCD camera for live time-lapse imaging. For timelapse imaging, Z-
stack images at 1µm steps were captured every 30 seconds for 15 minutes for in vivo time-lapse images and every 5 seconds for 10 minutes for cultured neurons. Filopodia were defined as long thin protrusions, less than 0.2 µm in diameter and more than 0.75 µm in length, measured from the cell body margin to the protrusion tip. In vivo filopodia lengths, lifetimes and fluorescence intensities of mRFP and GFP-Vangl2 were quantified using Zeiss Zen 2012 software. For cultured neurons, filopodium quantification was performed semi-automatically using Imaris FilamentTracer software (http://www.bitplane.com/imaris/filamenttracer). Mean anti-Vangl2 fluorescent intensity for all cell membranes were measured in user-drawn regions of interest using Zeiss Zen 2011 software or ImageJ’s “Plot Profile” tool. Anterior/posterior GFP-Vangl2 fluorescent intensity ratios for each cell were normalized by dividing this value by the anterior/posterior ZO-1 fluorescent intensity ratio. Graphs were generated and statistics were computed using GraphPad Prism software. All statistical analyses were performed using a 95% confidence interval. In most cases significance was determined using an unpaired, two-tail t-test with Welch’s correction. For the anti-Vangl2 staining quantification significance was determined using a paired two-tail t-test with Welch’s correction. Differences in FBMN distributions were analyzed using a Chi-square test where the distribution of FBMNs in control animals served as the expected frequencies or null hypothesis to determine if the observed frequencies were significantly different. Circular plots were generated using Oriana 4 software. Figure images were created using Adobe Photoshop and Adobe Illustrator.

Supporting Information

S1 Fig. Fzd3a has a cell-autonomous function in FBMN migration. (A-B) Live confocal images of 48 hpf chimeric embryos with anterior to the top. Transplant conditions are indicated as donor—host. Pk1bMO host embryos were used because they have normal neuroepithelial planar polarity but unmigrated FBMNs; this prevents donor-derived FBMNs from being carried to r6 by migrating host neurons in a PCP-independent manner. Cascade blue-dextran marks all donor-derived cells (blue), Tg(isl1:mRFP) marks host FBMNs (magenta) and Tg(isl1:GFP) marks donor-derived FBMNs (green). Histograms on the right indicate the percent of donor-derived FBMNs at 48 hpf that failed to migrate (rhombomere (r)4), partially migrated (r5) or fully migrated (r6) and numbers indicate the number of FBMNs represented in each bar. N indicates the number of chimeric embryos and n indicates the number of FBMNs scored in each condition. Brackets indicate rhombomere positon. Scale bar: 50 µm. (TIF)

S2 Fig. Post-mitotic FBMNs require PCP signaling for migration. (A) Live confocal image showing the dorsal view of a pk1b mutant embryo hindbrain at 48 hpf after transplantation of post-mitotic FBMNs from a wild type donor. Cascade blue-dextran marks all donor-derived cells (blue), Tg(isl1:GFP) marks host FBMNs (green) and Tg(isl1:mRFP) marks donor-derived FBMNs (magenta). (B) Histogram indicates the percent of donor-derived FBMNs at 48 hpf that failed to migrate, (rhombomere (r)4), partially migrated (r5) or fully migrated (r6) and numbers indicate the number of FBMNs represented in each bar. White arrows indicate migrated donor derived FBMNs. While post-mitotic FBMNs in general migrate poorly after being transplanted, they do sometimes migrate in WT and pk1b mutant hosts but never in vangl2 mutant hosts (see Fig 2). Brackets indicate rhombomere positon. Scale bar: 50µm. (TIF)

S3 Fig. PCP-DN expression in the floorplate disrupts planar polarity. (A-C) Tg(shh:Gal4) driven expression of Tg(UAS:Kaede) in the notochord and floorplate of a 14 hpf (A) 24 hpf embryo (B) and a 48 hpf embryo (C). Anterior is to the left. Images are live lateral views in A-C.
and live dorsal views at the hindbrain level, A’,B’. (D-F) Confocal images showing floorplate planar polarity of the anterior spinal cord in 48 hpf zebrafish embryos. Anterior is to the top. Anti-ZO-1 marks subapical tight junctions (white), anti-Cc2d2a marks the basal bodies of the primary cilia (magenta, arrows), and anti-GFP indicates dominant negative protein expression (green). Scale bar: 10μm. Whereas basal bodies are localized toward the posterior membrane in wild type embryos (D), this polarity is disrupted in floorplate cells expressing Xdd1-GFP (E) or Fzd3aΔC-GFP (F) (arrows in E’ and F’). (G) Schematic of the method used to quantify floorplate planar polarity. Total cell length (x) is measured as the distance between the anterior and posterior membranes (white) at the level of the basal body (magenta). Basal body position (y) is measured as the distance between the anterior membrane and the basal body. Cellular planar polarity is quantified as the ratio of x/y. (H) Quantitation of average basal body position in the floor plate of 48 hpf embryos. Each data point represents the mean basal body position for all cells quantitated in a single embryo. WT: N = 34 embryos, 411 cells; Xdd1-GFP: N = 14 embryos, 207 expressing cells; FzdAC-GFP: N = 29 embryos, 484 expressing cells; vangl2−/−: N = 10 embryos, 96 cells. Quantitation of floorplate polarity in vangl2−/− embryos is included for comparison. Graph represents data as mean ± SD. “p<0.001 compared to wild-type control.

(TIF)

S4 Fig. Vangl2 is not required in the mouse floorplate for FBMN migration. (A-B) Dorsal view of E13.5 mouse hindbrains with FBMNs (magenta) labeled with anti-Isl1 staining. Dotted lines indicate length of facial motor nucleus. To improve the chances that a Cre-expressing cell will have a biallelic deletion of Vangl2, in these experiments we used the Vangl2ΔTM null allele, which we discovered belatedly to cause a mild FBMN migration defect in compound heterozygotes with the floxed Vangl2loxP allele. Nevertheless, deleting the floxed allele with ShhCre did not enhance the partial migration defect in Vangl2ATM/ΔTM controls. For the experiments using Isl1Cre shown in Fig 1 we did not use the Vangl2ATM allele. (A) FBMNs in a Vangl2ATM/ΔTM control embryo. N = 6 embryos. (B) FBMNs in Vangl2loxP/ATM;ShhCre embryo. Addition of ShhCre does not further disrupt FBMN migration. N = 4 embryos. (C) Quantitation of FBMN migration stream length in Vangl2loxP/ATM control embryos and Vangl2loxP/ATM;ShhCre embryos. Scale bar: 100μm

(TIF)

S5 Fig. Specificity of the anti-Vangl2 antibody. (A-B) Dorsal view of wild type (A) and vangl2 mutant (B) 24 hpf neural tubes immunostained with anti-Vangl2-NT (green). The neuroepithelial membrane staining visible in wild type is absent in the mutant. (C) Western blot analysis of whole embryo lysates with anti-Vangl2 antibody. Anti-alpha-tubulin was used as a loading control. Zebrafish Vangl2 is expected to run at approximately 60kDa. For the anti-Vangl2 blot there is a band that is present in the wild type and absent in the vangl2 mutant, see asterisk.

(TIF)

S6 Fig. Migrating FBMNs display polarized protrusions that fail to polarize in vangl2 mutants. (A,C,E) Representative frames of mTFP expressing FBMNs from time-lapse images taken at 24 hpf to 32 hpf. (B,D,F) Each raw data point for protrusion angle is plotted on the circular graph below. Each division is 10 degrees. A, anterior. P, posterior. M, medial. L, lateral. Filopodia are radial in wild type FBMNs prior to exiting r4 (A,B, N = 3 embryos, 5 neurons, 28 filopodia) and become polarized to the posterior side of the cell during migration (C,D, N = 8 embryos, 10 neurons (6 in r5 and 4 in r6), 52 filopodia). FBMN protrusions fail to polarize in vangl2 mutants (E,F, N = 5 embryos, 7 neurons, 61 protrusions). Scale bar: 16μm.

(TIF)
S7 Fig. Donor-derived FBMNs used to quantitate filopodial dynamics were in a genetically 
chimeric environment. (A–E) Live confocal images of donor-derived FBMNs (green) and all 
other nearby donor-derived cells (magenta). Transplant conditions are indicated on as 
donor—host as in Fig 5. Rhodamine dextran marks all donor-derived cells (magenta), Tg isl1: 
mTFP marks donor-derived FBMNs (green). Anterior is to the top and medial is to the right. 
Scale bar: 5 μm. (TIF)

S8 Fig. Raw filopodial quantitation data. (A) Quantitation of filopodial lifetime for donor-
derived FBMNs. Each data point represents one filopodium. The maximum filopodial lifetime 
(900 seconds) corresponds to the full length of the time-lapse. (B) Quantitation of maximum 
filopodial length for filopodia lasting longer than 90 seconds on donor-derived FBMNs. Each 
data point represents one filopodium. (TIF)

S9 Fig. The effect of PCP on protrusion dynamics is dependent on the migratory environ-
ment. (A) Method used to isolate and identify FBMNs in primary culture. Embryos used were 
Tg isl1: mTFP; Tg hoxb1a: RFP allowing for the differentiation between FBMNs and other 
branchiomotor neurons labeled by Tg isl1: mTFP. (B,C) Cultured Tg isl1: mTFP; Tg hoxb1a: 
RFP FBMNs from a wild type (B) and a vangl2 mutant embryo (C). (B’,C’) Time-lapse spinning-disc 
confocal series of boxed region from B and C. (D) Quantitation of filopodial lifetime for cultured 
FBMNs. Each timelapse was 600 seconds total. p = 0.9044, n.s. (E) Quantitation of the 
maximum filopodial length for cultured FBMNs. p = 0.0856, n.s. Wild type: N = 8 neurons, 
64 filopodia. vangl2−/−: N = 8 neurons, 61 filopodia. Graphs represent data as mean ± SEM. 
Each data point is the average lifetime (D) or maximum length (E) for all the filopodia of one 
FBMN. Significance was determined using an unpaired, two-tail t-test with Welch’s correction. 
(TIF)

S1 Movie. Time-lapse of a GFP-Vangl2; Tg isl1: mRFP expressing FBMN. Note 
GFP-Vangl2 enrichment events preceding filopodia retraction. Movie is 4 frames per second 
(fps) with a 45 second time interval. (AVI)

S2 Movie. Time-lapse of LifeAct-GFP expressing FBMN. Movie is 4fps with a 13 second time 
interval. (AVI)

S3 Movie. Time-lapse of a Tg isl1: mTFP wild type FBMN in a wild type host. This and all 
subsequent movies are 4fps with 30 second time intervals. (AVI)

S4 Movie. Time-lapse of Tg isl1: mTFP vangl2 mutant FBMNs in a vangl2 mutant host. 
(AVI)

S5 Movie. Time-lapse of a Tg isl1: mTFP vangl2 mutant FBMN in a wild type host. 
(AVI)

S6 Movie. Time-lapse of a Tg isl1: mTFP fzd3a mutant FBMN in a wild type host. 
(AVI)

S7 Movie. Time-lapse of a Tg isl1: mTFP wild type FBMN in a vangl2 mutant host. 
(AVI)
S8 Movie. Time-lapse of a Tg(isl1:mTFP) wild type FBMNs in a fzd3a mutant host.

(AVI)

S1 Table. Primers used in the creation of transgenic constructs.

(PDF)

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Author Contributions

Conceived and designed the experiments: CFD AWM CBM. Performed the experiments: CFD AWM CBM. Analyzed the data: CFD AWM CBM. Contributed reagents/materials/analysis tools: CFD AWM CBM. Wrote the paper: CFD AWM CBM.

References

1. Gray R.S., Roszko I., and Solnica-Krezel L., Planar cell polarity: coordinating morphogenetic cell behaviors with embryonic polarity. Dev Cell, 2011. 21(1): p. 120–33. doi:10.1016/j.devcel.2011.06.011 PMID: 21763613

2. Goodrich L.V. and Strutt D., Principles of planar polarity in animal development. Development, 2011. 138(10): p. 1877–92. doi: 10.1242/dev.054080 PMID: 21521735

3. Gubb D. and Garcia-Bellido A., A genetic analysis of the determination of cuticular polarity during development in Drosophila melanogaster. J Embryol Exp Morphol, 1982. 68: p. 37–57. PMID: 6809878

4. Wong L.L. and Adler P.N., Tissue polarity genes of Drosophila regulate the subcellular location for prehair initiation in pupal wing cells. J Cell Biol, 1993. 123(1): p. 209–21. PMID: 1365886

5. Adler P.N., The genetic control of tissue polarity in Drosophila. Bioessays, 1992. 14(11): p. 735–41. PMID: 1365886

6. Borovina A., et al., Vangl2 directs the posterior tilting and asymmetric localization of motile primary cilia. Nat Cell Biol, 2010. 12(4): p. 407–12. doi: 10.1038/ncb2042 PMID: 20305649

7. Ciruna B., et al., Planar cell polarity signalling couples cell division and morphogenesis during neurulation. Nature, 2006. 439(7073): p. 220–4. PMID: 16407953

8. Wang Y., Guo N., and Nathans J., The role of Frizzled3 and Frizzled6 in neural tube closure and in the planar polarity of inner-ear sensory hair cells. J Neurosci, 2006. 26(8): p. 2147–56. PMID: 16495441

9. Wang J., et al., Regulation of polarized extension and planar cell polarity in the cochlea by the vertebrate PCP pathway. Nat Genet, 2005. 37(9): p. 980–5. PMID: 16116426

10. Montcuquiol M., et al., Identification of Vangl2 and Scrb1 as planar polarity genes in mammals. Nature, 2003. 423(6936): p. 173–7. PMID: 12724779

11. Strutt D.I., Weber U., and Mlodzik M., The role of RhoA in tissue polarity and Frizzled signalling. Nature, 1997. 387(6630): p. 292–5. PMID: 9153394

12. Winter C.G., et al., Drosophila Rho-associated kinase (Drok) links Frizzled-mediated planar cell polarity signaling to the actin cytoskeleton. Cell, 2001. 105(1): p. 81–91. PMID: 11301004

13. Fantin M., et al., Nuclear signaling by Rac and Rho GTPases is required in the establishment of epithelial planar polarity in the Drosophila eye. Curr Biol, 2000. 10(16): p. 979–88. PMID: 10985385

14. Carvajal-Gonzalez J.M. and Mlodzik M., Mechanisms of planar cell polarity establishment in Drosophila. F1000Prime Rep, 2014. 6: p. 98. doi: 10.12703/P6-98 PMID: 25580232

15. Jenny A., et al., Diego and Prickle regulate Frizzled planar cell polarity signalling by competing for Dishevelled binding. Nat Cell Biol, 2003. 7(7): p. 691–7. PMID: 15937478
16. Tree D.R., Ma D., and Axelrod J.D., A three-tiered mechanism for regulation of planar cell polarity. Semin Cell Dev Biol, 2002. 13(3): p. 217–24. PMID: 12137730

17. Wu J. and Mlodzik M., The frizzled extracellular domain is a ligand for Van Gogh/Stbm during nonautonomous planar cell polarity signaling. Dev Cell, 2008. 15(3): p. 462–9. doi: 10.1016/j.devcel.2008.08.004 PMID: 18804440

18. Strutt D. and Strutt H., Differential activities of the core planar polarity proteins during Drosophila wing patterning. Dev Biol, 2007. 302(1): p. 181–94. PMID: 17045581

19. Chin M.L. and Mlodzik M., The Drosophila selectin furrowed mediates intercellular planar cell polarity interactions via frizzled stabilization. Dev Cell, 2013. 26(5): p. 455–68. doi:10.1016/j.devcel.2013.07.006 PMID: 23973164

20. Taylor J., et al., Van Gogh: a new Drosophila tissue polarity gene. Genetics, 1998. 150(1): p. 199–210. PMID: 9725839

21. Vinson C.R. and Adler P.N., Directional non-cell autonomy and the transmission of polarity information by the frizzled gene of Drosophila. Nature, 1987. 329(6139): p. 549–51. PMID: 3116434

22. Wallingford J.B., Planar cell polarity and the developmental control of cell behavior in vertebrate embryos. Annu Rev Cell Dev Biol, 2012. 28: p. 627–53. doi:10.1146/annurev-cellbio-092910-154206 PMID: 22905955

23. Tissir F. and Goffinet A.M., Shaping the nervous system: role of the core planar cell polarity genes. Nat Rev Neurosci, 2013. 14(8): p. 525–35. doi:10.1038/nrn3525 PMID: 23839596

24. Tada M. and Kai M., Planar cell polarity in coordinated and directed movements. Curr Top Dev Biol, 2000. 21(5): p. 976–85. PMID: 11867525

25. Goto T. and Keller R., The planar cell polarity gene strabismus regulates convergence and extension and neural fold closure in Xenopus. Dev Biol, 2002. 247(1): p. 165–81. PMID: 12074560

26. Djiane A., et al., Role of frizzled 7 in the regulation of convergent extension movements during gastrulation in Xenopus laevis. Development, 2000. 127(10): p. 2227–38. PMID: 10769246

27. Wallingford J.B., et al., Dishevelled controls cell polarity during Xenopus gastrulation. Nature, 2000. 405(6782): p. 81–5. PMID: 10811222

28. Darken R.S., et al., The planar polarity gene strabismus regulates convergent extension movements in Xenopus. EMBO J, 2002. 21(5): p. 976–85. PMID: 11867525

29. Goto T. and Keller R., The planar cell polarity gene strabismus regulates convergence and extension and neural fold closure in Xenopus. Dev Biol, 2002. 247(1): p. 165–81. PMID: 12074560

30. Djiane A., et al., Role of frizzled 7 in the regulation of convergent extension movements during gastrulation in Xenopus laevis. Development, 2000. 127(10): p. 2227–38. PMID: 11493574

31. Jessen J.R., et al., Zebrafish trilobite identifies new roles for Strabismus in gastrulation and neuronal movements. Nat Cell Biol, 2002. 4(8): p. 610–5. PMID: 12105418

32. Wallingford J.B. and Harland R.M., Xenopus Dishevelled signaling regulates both neural and mesodermal convergent extension: parallel forces elongating the body axis. Development, 2001. 128(13): p. 2581–92. PMID: 11493574

33. Shindo A. and Wallingford J.B., PCP and septins compartmentalize cortical actomyosin to direct collective cell movement. Science, 2014. 343(6171): p. 649–52. doi:10.1126/science.1243126 PMID: 24503851

34. Wallingford J.B. and Harland R.M., Neural tube closure requires Dishevelled-dependent convergent extension of the midline. Development, 2002. 129(24): p. 5815–25. PMID: 12421719

35. Wang J., et al., Dishevelled genes mediate a conserved mammalian PCP pathway to regulate convergent extension during neurotubulation. Development, 2006. 133(9): p. 1767–78. PMID: 16571627

36. Ycht-Gonzalez P., et al., Convergent extension, planar-cell-polarity signalling and initiation of mouse neural tube closure. Development, 2007. 134(4): p. 789–99. PMID: 17229766

37. Williams M., et al., Distinct apical and basolateral mechanisms drive planar cell polarity-dependent convergent extension of the mouse neural plate. Dev Cell, 2014. 29(1): p. 34–46. doi:10.1016/j.devcel.2014.02.007 PMID: 24703875

38. Lyuksyutova A.I., et al., Anterior-posterior guidance of commissural axons by Wnt-frizzled signaling. Science, 2003. 302(5652): p. 1984–8. PMID: 14671310

39. Tissir F., et al., Protocadherin Cells3 is crucial in axonal tract development. Nat Neurosci, 2005. 8(4): p. 451–7. PMID: 15778712

40. Wang Y., et al., Frizzled-3 is required for the development of major fiber tracts in the rostral CNS. J Neurosci, 2002. 22(19): p. 8563–73. PMID: 12351730
41. Shafer B., et al., Vangl2 promotes Wnt/planar cell polarity-like signaling by antagonizing Dvl1-mediated feedback inhibition in growth cone guidance. Dev Cell, 2011. 20(2): p. 177–91. doi: 10.1016/j.devcel.2011.01.002 PMID: 21316586
42. Fenstermaker A.G., et al., Wnt/planar cell polarity signaling controls the anterior-posterior organization of monoaminergic axons in the brainstem. J Neurosci, 2010. 30(47): p. 16053–64. doi: 10.1523/JNEUROSCI.4508-10.2010 PMID: 21106844
43. Cantrell V.A. and Jessen J.R., The planar cell polarity protein Van Gogh-Like 2 regulates tumor cell migration and matrix metalloproteinase-dependent invasion. Cancer Lett, 2010. 287(1): p. 54–61. doi: 10.1016/j.canlet.2009.05.041 PMID: 19577357
44. Luga V., et al., Exosomes mediate stromal mobilization of autocrine Wnt-PCP signaling in breast cancer cell migration. Cell, 2012. 151(7): p. 1542–56. doi: 10.1016/j.cell.2012.11.024 PMID: 23260141
45. Chandrasekhar A., Turning heads: development of vertebrate branchiomotor neurons. Dev Dyn, 2004. 229(1): p. 143–61. PMID: 14699587
46. Wanner S.J., et al., Facial motor neuron migration advances. Curr Opin Neurobiol, 2013. 23(6): p. 943–50. doi: 10.1016/j.conb.2013.09.001 PMID: 24090878
47. Guthrie S., Patterning and axon guidance of cranial motor neurons. Nat Rev Neurosci, 2007. 8(11): p. 859–71. PMID: 17948031
48. Wada H., et al., Dual roles of zygotic and maternal Scribble1 in neural migration and convergent extension movements in zebrafish embryos. Development, 2005. 132(10): p. 2273–85. PMID: 15829519
49. Wada H., et al., Frizzled3a and Celsr2 function in the neuroepithelium to regulate migration of facial motor neurons in the developing zebrafish hindbrain. Development, 2006. 133(23): p. 4749–59. PMID: 17079269
50. Mapp O.M., et al., Zebrafish Prickle1b mediates facial branchiomotor neuron migration via a farnesylation-dependent nuclear activity. Development, 2011. 138(10): p. 2121–32. doi: 10.1242/dev.060442 PMID: 21521740
51. Walsh G.S., et al., Planar polarity pathway and Nance-Horan syndrome-like 1b have essential cell-autonomous functions in neuronal migration. Development, 2011. 138(14): p. 3033–42. doi: 10.1242/dev.063842 PMID: 21988578
52. Vivancos V., et al., Wnt activity guides facial branchiomotor neuron migration, and involves the PCP pathway and JNK and ROCK kinases. Neural Dev, 2009. 4: p. 7. doi: 10.1186/1749-8104-4-7 PMID: 19210786
53. Qu Y., et al., Atypical cadherins Celsr1-3 differentially regulate migration of facial branchiomotor neurons in mice. J Neurosci, 2010. 30(28): p. 9392–401. doi: 10.1523/JNEUROSCI.0124-10.2010 PMID: 20631168
54. Giasco D.M., et al., The mouse Wnt/PCP protein Vangl2 is necessary for migration of facial branchiomotor neurons, and functions independently of Dishevelled. Dev Biol, 2012. 369(2): p. 211–22. doi: 10.1016/j.ydbio.2012.06.021 PMID: 22771245
55. Sittaramane V., et al., The PCP protein Vangl2 regulates migration of hindbrain motor neurons by acting in floor plate cells, and independently of cilia function. Dev Biol, 2013. 382(2): p. 400–12. doi: 10.1016/j.ydbio.2013.08.017 PMID: 23988578
56. Higashijima S., Hotta Y., and Okamoto H., Visualization of cranial motor neurons in live transgenic zebrafish expressing green fluorescent protein under the control of the islet-1 promoter/enhancer. J Neurosci, 2000. 20(1): p. 206–18. PMID: 10627598
57. Boutros M., et al., Dishevelled activates JNK and discriminates between JNK pathways in planar polarity and wingless signaling. Cell, 1998. 94(1): p. 109–18. PMID: 9674432
58. Wanner S.J. and Prince V.E., Axon tracts guide zebrafish facial branchiomotor neuron migration through the hindbrain. Development, 2013. 140(4): p. 906–15. doi: 10.1242/dev.087148 PMID: 23325758
59. Rohrschneider M.R., Elsen G.E., and Prince V.E., Zebrafish Hoxb1a regulates multiple downstream genes including prickle1b. Dev Biol, 2007. 309(2): p. 358–72. PMID: 17651720
60. Qu Y., et al., Genetic evidence that Celsr3 and Celsr2, together with Fzd3, regulate forebrain wiring in a Vangl-independent manner. Proc Natl Acad Sci U S A, 2014. 111(29): p. E2996–3004. doi: 10.1073/pnas.140210511 PMID: 25002511
61. Copley C.O., et al., Postnatal refinement of auditory hair cell planar polarity deficits occurs in the absence of Vangl2. J Neurosci, 2013. 33(35): p. 14001–16. doi: 10.1523/JNEUROSCI.1037-13.2013 PMID: 23986237
62. Yang L., et al., Isl1Cre reveals a common Bmp pathway in heart and limb development. Development, 2006. 133(8): p. 1575–85. PMID: 16556916
63. Distel M., Wullimann M.F., and Koster R.W., Optimized Gal4 genetics for permanent gene expression mapping in zebrafish. Proc Natl Acad Sci U S A, 2009. 106(32): p. 13365–70. doi: 10.1073/pnas.0903060106 PMID: 19628697

64. Choe S.K., et al., A Gal4/UAS system for conditional transgene expression in rhombomere 4 of the zebrafish hindbrain. Dev Dyn, 2012. 241(6): p. 1125–32. doi: 10.1002/dvdy.23794 PMID: 22499412

65. Roszko I., et al., A dynamic intracellular distribution of Vangl2 accompanies cell polarization during zebrafish gastrulation. Development, 2015. 142(14): p. 2508–20. doi: 10.1242/dev.119032 PMID: 26062934

66. Ossipova O., Kim K., and Sokol S.Y., Planar polarization of Vangl2 in the vertebrate neural plate is controlled by Wnt and Myosin II signaling. Biol Open, 2015. 4(6): p. 722–30. doi: 10.1242/bio.201511676 PMID: 25910938

67. Grant P.K. and Moens C.B., The neuroepithelial basement membrane serves as a boundary and a substrate for neuron migration in the zebrafish hindbrain. Neural Dev, 2010. 5: p. 9. doi: 10.1186/1749-8104-5-9 PMID: 20350296

68. Harfe B.D., et al., Evidence for an expansion-based temporal Shh gradient in specifying vertebrate digit identities. Cell, 2004. 118(4): p. 517–28. PMID:15315763

69. Trevarrow B., Marks D.L., and Kimmel C.B., Organization of hindbrain segments in the zebrafish embryo. Neuron, 1990. 4(5): p. 669–79. PMID:2344406

70. Xiong F., et al., Specified neural progenitors sort to form sharp domains after noisy Shh signaling. Cell, 2013. 153(3): p. 550–61. doi: 10.1016/j.cell.2013.03.023 PMID: 23622440

71. Mapp O.M., et al., Prickle1b mediates interpretation of migratory cues during zebrafish facial branchiomotor neuron migration. Dev Dyn, 2010. 239(6): p. 1596–608. doi: 10.1002/dvdy.22283 PMID: 20503357

72. Portera-Cailliau C., Pan D.T., and Yuste R., Activity-regulated dynamic behavior of early dendritic protrusions: evidence for different types of dendritic filopodia. J Neurosci, 2008. 23(18): p. 7129–42. PMID: 18555784

73. Lim K.B., et al., The Cdc42 effector IRSp53 generates filopodia by coupling membrane protrusion with actin dynamics. J Biol Chem, 2008. 283(29): p. 20454–72. doi: 10.1074/jbc.M710185200 PMID: 18448434

74. Ahmed S., Goh W.I., and Bu W., I-BAR domains, IRSp53 and filopodium formation. Semin Cell Dev Biol, 2010. 21(4): p. 350–6. doi: 10.1016/j.semcdb.2009.11.008 PMID: 19913105

75. Villefranc J.A., et al., A truncation allele in vascular endothelial growth factor c reveals distinct modes of signaling during lymphatic and vascular development. Development, 2013. 140(7): p. 1497–506. doi: 10.1242/dev.084152 PMID: 23462469

76. Chen W.S., et al., Asymmetric homotypic interactions of the atypical cadherin flamingo mediate intercellular polarity signaling. Cell, 2008. 133(6): p. 1093–105. doi: 10.1016/j.cell.2008.04.048 PMID: 18555784

77. Strutt D. and Warrington S.J., Planar polarity genes in the Drosophila wing regulate the localisation of the FH3-domain protein Multiple Wing Hairs to control the site of hair production. Development, 2008. 135(18): p. 3103–11. doi: 10.1242/dev.052520 PMID: 18701542

78. Yan J., et al., The multiple-wing-hairs gene encodes a novel GBD-FH3 domain-containing protein that functions both prior to and after wing hair initiation. Genetics, 2008. 180(1): p. 219–28. doi: 10.1534/genetics.108.091314 PMID: 18723886

79. Kaucka M., et al., Asymmetry of VANGL2 in migrating lymphocytes as a tool to monitor activity of the mammalian WNT/planar cell polarity pathway. Cell Commun Signal, 2015. 13: p. 2. doi: 10.1186/s12964-014-0079-1 PMID: 25627785

80. Nishimura T., Honda H., and Takeichi M., Planar polarity links axes of spatial dynamics in neural tube closure. Cell, 2012. 149(5): p. 1084–97. doi: 10.1016/j.cell.2012.04.021 PMID: 22632972

81. Yin C., et al., Cooperation of polarized cell intercalations drives convergence and expansion of presomitic mesoderm during zebrafish gastrulation. J Cell Biol, 2008. 180(1): p. 221–32. doi: 10.1083/jcb.200704150 PMID: 18195109

82. Phng L.K., Stanchi F., and Gerhardt H., Filopodia are dispensable for endothelial tip cell guidance. Development, 2013. 140(19): p. 4031–40. doi: 10.1242/dev.097352 PMID: 24046319

83. Dwivedy A., et al., Ena/VASP function in retinal axons is required for terminal arborization but not pathway navigation. Development, 2007. 134(11): p. 2137–46. PMID: 17507414

84. Boer E.F., et al., Fascin1-dependent Filopodia are required for directional migration of a subset of neural crest cells. PLoS Genet, 2015. 11(1): p. e1004946. doi: 10.1371/journal.pgen.1004946 PMID: 25607881
85. Mattila P.K. and Lappalainen P., Filopodia: molecular architecture and cellular functions. Nat Rev Mol Cell Biol, 2008. 9(6): p. 446–54. doi: 10.1038/nrm2406 PMID: 18464790
86. Heckman C.A. and Plummer H.K. 3rd, Filopodia as sensors. Cell Signal, 2013. 25(11): p. 2298–311. doi: 10.1016/j.cellsig.2013.07.006 PMID: 23876793
87. Bentley D. and Torian-Raymond A., Disoriented pathfinding by pioneer neurone growth cones deprived of filopodia by cytochalasin treatment. Nature, 1986. 323(6090): p. 712–5. PMID: 3773996
88. Chien C.B., et al., Navigational errors made by growth cones without filopodia in the embryonic Xenopus brain. Neuron, 1993. 11(2): p. 237–51. PMID: 8352941
89. Zheng J.Q., Wan J.J., and Poo M.M., Essential role of filopodia in chemotropic turning of nerve growth cone induced by a glutamate gradient. J Neurosci, 1996. 16(3): p. 1140–9. PMID: 8558243
90. Meyen D., et al., Dynamic filopodia are required for chemokine-dependent intracellular polarization during guided cell migration in vivo. Elife, 2015. 4.
91. Galbraith C.G., Yamada K.M., and Galbraith J.A., Polymerizing actin fibers position integrins primed to probe for adhesion sites. Science, 2007. 315(5814): p. 992–5. PMID: 17303755
92. Steketele M.B. and Tosney K.W., Three functionally distinct adhesions in filopodia: shaft adhesions control lamellar extension. J Neurosci, 2002. 22(18): p. 8071–83. PMID: 12223561
93. Johnson H.E., et al., F-actin bundles direct the initiation and orientation of lamellipodia through adhesion-based signaling. J Cell Biol, 2015. 208(4): p. 443–55. doi: 10.1083/jcb.201406102 PMID: 25666809
94. Habas R., Kato Y., and He X., Wnt/Frizzled activation of Rho regulates vertebrate gastrulation and requires a novel Formin homology protein Daam1. Cell, 2001. 107(7): p. 843–54. PMID: 11779461
95. Ulmer B., et al., Calponin 2 acts as an effector of noncanonical Wnt-mediated cell polarization during neural crest cell migration. Cell Rep, 2013. 3(3): p. 615–21. doi: 10.1016/j.celrep.2013.02.015 PMID: 23499442
96. Brooks S.P., et al., The Nance-Horan syndrome protein encodes a functional WAVE homology domain (WHD) and is important for co-ordinating actin remodelling and maintaining cell morphology. Hum Mol Genet, 2010. 19(12): p. 2421–32. doi: 10.1093/hmg/ddq125 PMID: 20332100
97. Classen A.K., et al., Hexagonal packing of Drosophila wing epithelial cells by the planar cell polarity pathway. Dev Cell, 2005. 9(6): p. 805–17. PMID: 16326392
98. Warrington S.J., Strutt H., and Strutt D., The Frizzled-dependent planar polarity pathway locally promotes E-cadherin turnover via recruitment of RhoGEF2. Development, 2013. 140(5): p. 1045–54. doi: 10.1242/dev.088724 PMID: 23364328
99. Nagaoka T., et al., Vangl2 regulates E-cadherin in epithelial cells. Sci Rep, 2014. 4: p. 6940. doi: 10.1038/srep06940 PMID: 25373475
100. Williams B.B., et al., VANGL2 regulates membrane trafficking of MMP14 to control cell polarity and migration. J Cell Sci, 2012. 125(Pt 9): p. 2141–7. doi: 10.1242/jcs.079764 PMID: 22357946
101. Asakawa K., et al., Genetic dissection of neural circuits by Tol2 transposon-mediated Gal4 gene and enhancer trapping in zebrafish. Proc Natl Acad Sci U S A, 2008. 105(4): p. 1255–60. doi: 10.1073/pnas.0704963105 PMID: 18202183
102. Koide T., et al., Olfactory neural circuitry for attraction to amino acids revealed by transposon-mediated gene trap approach in zebrafish. Proc Natl Acad Sci U S A, 2009. 106(24): p. 9884–9. doi: 10.1073/pnas.0900470106 PMID: 19497864
103. Tree D.R., et al., Prickle mediates feedback amplification to generate asymmetric planar cell polarity signaling. Cell, 2002. 109(3): p. 371–81. PMID: 12015986
104. Heisenberg C.P., et al., Silberblick/Wnt11 mediates convergent extension movements during zebrafish gastrulation. Nature, 2000. 405(6782): p. 76–81. PMID: 10811221
105. Westerfield M., The Zebrafish Book. University of Oregon Press, Eugene, OR. 1993.
106. Kimmel C.B., et al., Stages of embryonic development of the zebrafish. Dev Dyn, 1995. 203(3): p. 253–310. PMID: 8599427
107. Kwan K.M., et al., The Tol2kit: a multisite gateway-based construction kit for Tol2 transposon transgenesis constructs. Dev Dyn, 2007. 236(11): p. 3088–99. PMID: 17937935
108. Ertzer R., et al., Cooperation of sonic hedgehog enhancers in midline expression. Dev Biol, 2007. 301(2): p. 578–89. PMID: 17157288
109. Muller F., et al., Intronic enhancers control expression of zebrafish sonic hedgehog in floor plate and notochord. Development, 1999. 126(10): p. 2103–16. PMID: 10207136
110. Meng A., et al., Promoter analysis in living zebrafish embryos identifies a cis-acting motif required for neuronal expression of GATA-2. Proc Natl Acad Sci U S A, 1997. 94(12): p. 6267–72. PMID: 9177206

111. Kawakami K., Shima A., and Kawakami N., Identification of a functional transposase of the Tol2 element, an Ac-like element from the Japanese medaka fish, and its transposition in the zebrafish germ lineage. Proc Natl Acad Sci U S A, 2000. 97(21): p. 11403–8. PMID: 11027340

112. Carmany-Rampey A. and Moens C.B., Modern mosaic analysis in the zebrafish. Methods, 2006. 39(3): p. 228–38. PMID: 16829130

113. Bachmann-Gagescu R., et al., The ciliopathy gene cc2d2a controls zebrafish photoreceptor outer segment development through a role in Rab8-dependent vesicle trafficking. Hum Mol Genet, 2011. 20(20): p. 4041–55. doi: 10.1093/hmg/ddr332 PMID: 21816947

114. Andersen S.S., Preparation of dissociated zebrafish spinal neuron cultures. Methods Cell Sci, 2001. 23(4): p. 205–9. PMID: 12486331