Vangl2 deficient zebrafish exhibit hallmarks of neural tube closure defects

Jacalyn MacGowan^{1,2}, Mara Cardenas^{2,3,4}, and Margot Kossmann Williams^{1,2*}

¹Center for Precision Environmental Health, Baylor College of Medicine, Houston, TX

- ² Department of Molecular and Cellular Biology, Baylor College of Medicine, Houston, TX
- ³ Center for Cell and Gene Therapy, Baylor College of Medicine, Houston, TX
- ⁴ Stem Cells and Regenerative Medicine Center, Baylor College of Medicine, Houston, TX

*Author for correspondence: margot.williams@bcm.edu

Running title

Zebrafish neural tube closure

15 **Keywords** 16

- Neurulation
- 17 Neural tube defects 18
- Planar cell polarity 19
- Zebrafish 20
- 21 22

23

26 27

28

1 2

3 4

5

6

7

8

9 10 11

12

13 14

Summary statement

The anterior neural tube of zebrafish exhibits fold-and-fuse neurulation which is disrupted upon loss of Vangl2. 24 highlighting conservation of vertebrate neurulation and the potential to model neural tube defects in zebrafish. 25

Abstract

29 Neural tube defects (NTDs) are among the most devastating and common congenital anomalies worldwide. 30 and the ability to model these conditions in vivo is essential for identifying causative genetic and environmental 31 factors. Although zebrafish are ideal for rapid candidate testing, their neural tubes develop primarily via a solid 32 neural keel rather that the fold-and-fuse method employed by mammals, raising questions about their 33 suitability as an NTD model. Here, we demonstrate that despite outward differences, zebrafish anterior 34 neurulation closely resembles that of mammals. For the first time, we directly observe fusion of the bilateral 35 neural folds to enclose a lumen in zebrafish embryos. The neural folds fuse by zippering between multiple 36 distinct but contiguous closure sites. Embryos lacking vangl2, a core planar cell polarity and NTD risk gene, 37 exhibit delayed neural fold fusion and abnormal neural groove formation, yielding distinct openings and midline 38 bifurcations in the developing neural tube. These data provide direct evidence for fold-and-fuse neurulation in 39 zebrafish and its disruption upon loss of an NTD risk gene, highlighting conservation of vertebrate neurulation 40 and the utility of zebrafish for modeling NTDs. 41

Introduction 44

42 43

45

Neural tube defects (NTDs) such as spina bifida and anencephaly are among the most common and 46 devastating congenital anomalies, affecting approximately 1 in 1,000 births in the United States (1) and even 47 more worldwide (2). These conditions result from incomplete closure of the neural tube during embryogenesis, 48 often leaving the neural tube lumen open to the outside of the body. While folate supplementation has greatly 49

reduced the incidence of NTDs, as many as 70% of cases are folate-resistant (3). Mutations in a number of 50 genes are associated with an increased likelihood of NTDs (4), but are not the sole risk factors for NTD (5, 6). 51 Additional potential risk factors include mutations in multiple risk genes (7), certain teratogenic drugs (8, 9), 52 and exposure to environmental toxicants (10-16), implicating complex gene-environment interactions in their 53 etiology. Indeed, the stalled decline in NTDs after folate fortification in the U.S. (3) demonstrates that additional 54 risk factors remain to be discovered. One major limitation is the inability to draw causal relationships between 55 reported risk factors and NTD occurrence. Some genetic and/or environmental risk factors identified from 56 patient cohorts have been shown to cause NTDs in mouse models (17-19). However, the small litter sizes, high 57 husbandry costs, and large space requirements of mice are prohibitive of large-scale genetic and/or chemical 58 screening that could convert associations into causal relationships, highlighting the need for additional animal 59 models of NTDs. 60

61

Due to their relatively low husbandry costs, rapid external development, large clutch sizes, and amenability to 62 genetic manipulation, zebrafish are an ideal vertebrate model for genetic and chemical screens to identify 63 causes of NTDs. However, the morphology of the developing neural tube differs substantially between 64 zebrafish and amniote species. Primary neurulation in chick, mouse, and human embryos is driven by 65 convergent extension (CE) of the developing neural plate followed by formation of hinge points (20-27) that 66 elevate the bilateral neural folds and bend them toward each other (26, 28, 29). The neural folds then meet at 67 the dorsal midline and fuse by zippering between discrete closure points (30, 31), completing the "fold-and-68 fuse" process that encloses the neural tube lumen. By contrast, the zebrafish spinal cord develops from the 69 neural keel, a solid structure that later undergoes cavitation to form a central lumen (32, 33). The site of this 70 lumen is established through a series of midline-crossing mitoses termed "C-divisions" which distribute one 71 daughter cell of each side of the neural keel midline (32, 34-39). For this reason, primary neurulation in 72 zebrafish has been likened to secondary neurulation in amniote embryos (33, 40, 41). Zebrafish were therefore 73 viewed not only as questionable for NTD modeling, but also as fundamentally different from other vertebrates 74 in their mechanism of neurulation. 75

76

However, more recent findings reveal that despite these outward differences, several hallmarks of primary 77 neurulation are conserved in zebrafish. For example, CE morphogenesis narrows the neural plate (38, 42), and 78 apical constriction at the midline forms a medial hinge point-like structure (43, 44). The neural folds were also 79 shown to zipper closed in the forebrain region of zebrafish (44) in a fashion strikingly similar to mice (30, 31). 80 These conserved neurulation mechanisms open the possibility of modeling NTDs, or aspects thereof, in the 81 experimentally tractable zebrafish model. Indeed, previous studies have proposed bifurcation of pineal gland 82 precursors and/or the dorsal roof plate as proxies for NTDs in zebrafish (45-48). These phenotypes are 83 suggestive of reduced neural fold convergence, but it is unclear whether they exhibit other hallmarks of NTDs, 84 such as lumens that remain open to the outside. Furthermore, bifurcated pineal and roof plate domains 85 resulted from reduced Nodal signaling and N-cadherin function (45-48), but mutations in Nodal signaling 86

components are associated with holoprosencephaly in human patients (*49-51*) rather than NTDs. Whether
 these or other NTD-like phenotypes are induced by loss of human NTD risk genes in zebrafish was unknown.

Among mutations known or suspected to cause NTDs in mice and humans, respectively, many affect 90 components of planar cell polarity (PCP) signaling, a highly conserved regulator of vertebrate morphogenesis. 91 Loss of PCP genes like Vangl2 disrupts both CE of the neural plate and hinge point formation, preventing 92 anteroposterior axis elongation and neural tube closure in mouse, chick, and Xenopus (20, 21, 52-65). 93 Mutations in PCP genes (including VANGL2) are also associated with NTDs in several patient cohorts (7, 66-94 71). Loss of PCP signaling similarly disrupts CE and neural tube development in zebrafish (38, 42, 72-81), but 95 these phenotypes differ from amniote NTDs due to differences in neurulation. For example, C-divisions (which 96 determine the site of lumen formation) are disorganized In PCP mutant vangl2 / trilobite (tri)-/- zebrafish, giving 97 rise to ectopic neuroectodermal structures and neural tube lumens (38, 39, 72). Because these phenotypes do 98 not outwardly resemble NTDs in other vertebrate species or humans (i.e. no lumens that remain open to the 99 outside), it was broadly assumed that zebrafish neurulate via fundamentally different mechanisms than other 100 vertebrates and are poor models for NTD research. 101

102

Here, we reevaluate the neural tube phenotypes of vangl2 deficient zebrafish embryos with a focus on the 103 brain region at early stages of neuralation. By examining neural tube development at the time of neural fold 104 zippering, we found that many vangl2-deficient embryos exhibit a Distinct Opening of the Neural Tube 105 (DONuT), a 3-dimensional pit-shaped structure in the forebrain region. DONuT formation correlates with the 106 severity of axis extension defects and delayed midline convergence of pineal precursors, linking it to reduced 107 CE morphogenesis. Live time-lapse imaging of neural tube closure revealed that, in addition to the previously 108 described zippering of the anterior neural folds, wild-type (WT) zebrafish exhibit a distinct but contiguous 109 posterior site of neural fold fusion. These two sites zipper in opposite directions from a central point of contact 110 to close the anterior neural tube. Using optical transverse sectioning of live embryos, we further showed that 111 the bilateral neural folds fuse to enclose a lumen in a process strikingly similar to amniote neurulation. 112 Moreover, neural fold fusion is delayed in *vangl2* deficient embryos, reflecting reduced CE cell movements and 113 abnormal neural groove formation. Together, these data provide direct evidence for fold-and-fuse neurulation in 114 zebrafish and show that this process requires Vangl2. This demonstrates the deep conservation of neurulation 115 mechanisms among vertebrates and highlights the potential utility of zebrafish for modeling the morphogenetic 116 processes underlying neural tube closure and NTDs. 117

Results

118 119

120 121

124

Bifurcated pineal precursors and roof plates reflect abnormal anterior neural tube morphology in Vangl2 deficient zebrafish embryos.

Bifurcation of pineal gland precursors and the dorsal roof plate around 24 hours post fertilization (hpf) were previously suggested as proxies for NTDs in zebrafish embryos with disrupted Nodal signaling or mutations in

cdh2 (encoding N-cadherin) (45-48). To determine whether such phenotypes were present in zebrafish 127 embryos lacking the PCP signaling component (and ortholog of a human NTD risk gene (70, 71)) vangl2, we 128 examined the morphology of these structures in *vangl2/trilobite (tri)* homozygous mutants, *vangl2* morphants, 129 and sibling controls at approximately 28 hpf. Using both a transgenic flh:kaede (82) line in which pineal 130 precursors fluoresce green (Fig. 1A-B) and whole mount *in situ* hybridization (WISH) for the pineal marker otx5 131 (Fig. 1D), we found that embryos injected with 2 ng vangl2 morpholino oligonucleotide (MO) (83) exhibited 132 significantly wider pineal domains than control siblings when measured from the lateral-most edges (Fig. 1C) 133 and were significantly more likely to have split or elongated oval-shaped pineal domains than controls (Fig. 134 1E). Notably, homozygous vangl2/trilobite (tri)-/- embryos did not exhibit increased pineal width at this stage 135 (Fig. 1C). This is likely due to maternally deposited vangl2 that is eliminated by the MO (42, 84), consistent 136 with more severe phenotypes in maternal zygotic (MZ)vangl2-/- than zygotic (Z)vangl2-/- embryos (72). WISH 137 for the dorsal neural tube marker wnt1 also revealed bifurcated roof plates in a subset vanal2 morphants (Fig. 138 1F-I). Histological analysis of 28 hpf vangl2 morphant anterior neural tubes (fore- through hind-brain) with split 139 or oval-shaped pineal domains revealed ectopic tissue and supernumerary midline structures (Fig. 1J-M, 140 yellow arrows) similar to those reported in the spinal cord region of vangl2 deficient embryos (38, 39, 72). Eight 141 of the ten embryos with split pineal domains contained ectopic midline tissue at all rostrocaudal positions 142 examined, including at the level of the pineal (Fig. 1K). Among the embryos examined with oval-shaped 143 pineals, only one contained ectopic midline tissue at the level of the pineal (Fig. 1L) while the remaining four 144 contained a single midline in the rostral-most sections but ectopic midlines further rostrally (Fig. 1M). This 145 suggests that split pineal domains are associated with ectopic midline tissues resulting from abnormal C-146 divisions, but that the anterior neural tube is less sensitive to loss of vangl2 than more posterior regions. 147

We compared these phenotypes to those of Nodal signaling-deficient embryos, in which split pineal 148 domains were hypothesized to represent open neural tubes (45-47). As previously reported, embryos 149 completely lacking Nodal signaling through maternal and zygotic loss of the Nodal coreceptor oep/tdgf1 150 (MZoep-/-) or treated with the Nodal inhibitor SB505124 at either sphere or dome stage (4-4.3 hpf) exhibited 151 substantially wider and split pineal domains (Supp. Fig. 1A-E). Later SB505124 treatment at 30% epiboly (4.7 152 hpf) yielded embryos with a range of pineal phenotypes from split to round, as seen previously (46). 153 Histological analysis of SB505124-treated embryos at 28 hpf revealed a striking Swiss cheese-like pattern of 154 multiple small holes in the neural tube of every embryo examined, regardless of whether their pineal domains 155 were split or closed (Supp. Fig. 1F-H). This internal anatomy of the neural tube is consistent with previous 156 reports of multiple lumen-like structures in MZoep-/- embryos (85), and is distinct from the ectopic bilateral 157 midlines observed in vangl2 morphants (Fig.1 J-M) despite similar pineal phenotypes. This demonstrates that 158 bifurcation of the pineal precursors and roof plate externally can be underlain by multiple distinct internal 159 phenotypes, which do not appear to share features with amniote NTDs. 160

bioRxiv preprint doi: https://doi.org/10.1101/2023.11.09.566412; this version posted November 9, 2023. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.



Figure 1. Loss of vangl2 disrupts pineal and neural roof plate morphology.

A-B) Example images of pineal precursor morphology in Tg[*flh:kaede*] control (A) or *vangl2* MO-injected (B) embryos at 28 hpf, viewed from the dorsal side. C) Width of pineal precursor domains in embryos of the conditions indicated, as measured from *otx5* whole mount in situ hybridization (WISH) at 28 hpf (D, blue lines). Each dot represents a single embryo, black bars are median values. ****p<0.0001, ns p=0.10, Mann-Whitney tests. E) Classification of pineal shape in 28 hpf control and *vangl2* MO-injected embryos expressing *flh:kaede* or WISH stained for *otx5*. n values indicate the number of embryos of each condition measured from at least 3 independent trials. ****p<0.0001, Chi-square test. Scale bars = 50 μ m. F-I) WISH for roof plate marker *wnt1* at 28 hpf in control (F-G) and *vangl2* MO-injected (H-I) embryos. Yellow arrowheads indicate the midline roof plate, cyan arrowheads indicate the epithalamus, and magenta arrowheads indicate the mid-hindbrain boundary (MHB). J-M) Transverse histological sections through the anterior neural tube at the level of the epithalamus (top panels), midbrain (middle panels), and MHB (bottom panels) in 28 hpf embryos of the conditions indicated. Cyan arrowheads indicate pineal precursors stained by *otx5* WISH. Yellow arrowheads indicate the neural tube midline(s)/lumen(s). Fractions indicate the number of embryos with the depicted phenotype over the total number of embryos examined for each condition. Anterior is up in (A,B,D,G,I), dorsal is up in (F,H,J-M).



Supplemental Figure 1. Distinct neural tube morphologies underlie split pineal phenotypes in Nodal deficient embryos.

A-D) Representative images of the anterior neural tube in DMSO- treated WT (A), MZoep-/-, or SB505124treated (C-D) embryos at 28 hpf WISH stained for *otx5* and *her5*, viewed dorsally. Cyan arrowheads indicate pineal precursors, magenta arrowheads indicate the MHB. **E**) Width of pineal precursor domains in embryos of the conditions indicated, measured from *otx5* WISH at 28 hpf (as shown in A-D). Each dot represents a single embryo, black bars are median values. **F-H**) Transverse histological sections through the anterior neural tube at the level of the epithalamus (top panels), midbrain (middle panels), and MHB (bottom panels) in 28 hpf embryos of the conditions indicated. Cyan arrowheads indicate pineal precursors stained by *otx5* WISH. Yellow arrowheads indicate ectopic lumens in a Swiss cheese-like pattern. Fractions indicate the number of embryos with the depicted phenotype over the total number of embryos examined for each condition. Anterior is up in (A-D), dorsal is up in (F-H).

Vangl2 deficient embryos exhibit delayed pineal precursor convergence and distinct openings of the neural tube.

Previous analysis of the spinal cord region in *vangl2* deficient embryos determined that ectopic tissue at the 165 neural tube midline results from abnormal C-divisions (38, 39, 72) that normally distribute pairs of neural 166 progenitor cells on either side of the neural midline (32, 34, 37) but occur ectopically in the absence of vanal2. 167 We frequently observed a tissue mass between the split pineal domains of *vangl2* morphants (Fig. 1K), raising 168 the possibility that ectopic midline tissue prevents pineal fusion. However, WT pineal fusion and anterior neural 169 tube closure (at the 7-somite stage (44, 45)) precede C-divisions (at approximately 10-somite stage (34, 35, 170 41)), and are therefore unlikely to be caused by abnormal cell divisions. To identify the developmental origins 171 of pineal phenotypes in embryos lacking vanal2, we examined neural development at the time of neural fold 172 zippering (44) from 3-10 somite stages. In control embryos, pineal precursors marked by flh/noto expression 173 began as bilateral domains that converged at the midline and fused at approximately 7 somite stage (Fig. 2A. 174 D), as previously described (45). In vangl2 morphant (and to a lesser degree, tri-/- mutant) embryos, we found 175 that this convergence was delayed, as reflected by the increased width of *flh*+ domains across several time 176 points (Fig. 2B, D; Supp. Fig. 2A). 177

However, we observed an even more striking phenotype in *vangl2* morphant embryos during this analysis: 178 a distinct, horseshoe-shaped pit in the neural plate between the *flh*+ pineal domains (Fig. 2B-C). This structure 179 was present in most vanal2 morphant embryos from 3-6 somite stages, and although its incidence decreased 180 as development proceeded, it persisted in approximately 20% of vangl2 morphants at the 8-somite stage (Fig. 181 2D). This pit-like structure was not common to all embryos whose pineal precursors had not yet fused, as only 182 one control embryo (of 163 examined) possessed such a structure even at pre-pineal fusion stages. Although 183 the pineal domains of vanal2/ tri-/- embryos were wider than their WT and heterozygous controls, none 184 exhibited a pit in their forebrain region (Supp. Fig. 2A-B), consistent with more severe phenotypes in vangl2 185 morphants than mutants. 3-dimensional reconstructions of these structures from confocal Z-stacks of fixed 186 vangl2 morphant and control embryos stained with phalloidin (Fig. 2F-G) corroborated our WISH images. At 3-187 5 somite stages, this structure resembled a pit that was open to the exterior of the embryo (Fig. 2G, yellow 188 arrowheads). Many vangl2 morphants also possessed a second, more posteriorly positioned pit structure (Fig. 189 **2G**, cyan arrowheads). By contrast, the anterior neural plate of control embryos was covered by periderm 190 (which develops from and is sometimes referred to as the enveloping layer (EVL)) and appeared smooth (Fig. 191 2F). This smooth appearance persisted to the 7-somite stage, whereas vangl2 morphant embryos exhibited 192 openings from which rounded cells protruded to the embryo's exterior. Because these finding provide direct 193 evidence for open neural tubes in vangl2 morphant embryos, we termed this pit-shaped structure the Distinct 194 Opening of the Neural Tube (DONuT). 195

196 197



Figure 2. *vangl2* deficient embryos exhibit delayed pineal convergence and a Distinct Opening of the Neural Tube (DONuT).

A-B) Representative images of pineal precursors (*flh* WISH) and somites/adaxial cells (*myoD* WISH) in control (A) and *vangl2* MO-injected (B) embryos at the stages indicated, viewed dorsally. Blue and burgundy lines indicate pineal precursor width, yellow arrowheads indicate the presence of a DONuT. **C**) Enlargements of the DONuTs from images in (B). **D**) Width of pineal precursor domains (as shown in A-B) in control (blue) and *vangl2* morphant (burgundy) embryos at the stages indicated. Each dot represents a single embryo, black bars are median values. n values indicate the number of embryos of each stage/condition measured from 3 independent trials, **p=0.003, p=0.008, p=0.007, ****p<0.0001, T-test. **E**) Percentage of embryos (as shown in A-B) of the stage/condition indicated with (black) or without (gray) a DONuT. **F-G**) 3-dimensional reconstructions of confocal Z-stacks through the anterior neural plate of fixed and phalloidin-stained embryos of the stages and conditions indicated. Yellow arrowheads indicate a DONuT, cyan arrowheads indicate openings in more posterior regions. n values indicate the number of embryos examined for each condition from 3 independent trials. Anterior is up in all images, scale bars = 100 µm.



Supplemental Figure 2. Trilobite mutants do not exhibit DONuTs.

A) Width of pineal precursor domains (as shown in A-B) in WT (blue), *tri* het (indigo), and *tri* homozygous (purple) embryos at the stages indicated. Each dot represents a single embryo, black bars are median values. n values indicate the number of embryos of each stage/condition measured from 3 independent trials. **B**) Percentage of embryos from *tri* heterozygote incrosses of the stage/genotype indicated with (black) or without (gray) a DONuT.

Anterior neural tube openings correlate with severity of convergent extension defects.

199 Vangl2 deficient embryos have well described defects in CE morphogenesis during gastrulation (42, 78, 84), 200 and their neural tube phenotypes at later stages are thought to be secondary to reduced CE (38). We 201 hypothesized that defective CE also underlies the observed delay in convergence of the bilateral pineal 202 precursors, which in-turn manifests as a DONuT. Consistent with this hypothesis, we found that both the *flh*+ 203 pineal domains and myoD+ somites were significantly wider (consistent with reduced CE) in 3-7 somite staged 204 vangl2 morphants with a DONuT than those without a DONuT (Fig. 3A-C, E-F). Notably, the width of pineal 205 domains and somites of vangl2/ tri-/- embryos did not differ from morphants without DONuTs but were 206 narrower than morphants with DONuTs (Fig. 3D, E-F). This is consistent with our failure to observe DONuTs in 207 mutant embryos, and with previous findings that Zvangl2-/- CE phenotypes are less severe than those of 208 MZvangl2-/- and vangl2 morphants (72). Together, these findings indicate that embryos with more severe CE 209 defects are more likely to exhibit openings in the neural tube, strongly implicating reduced CE during 210 astrulation in neural tube closure. We further speculate that some vanal2 deficient embryos can never 211 overcome this delay to successfully fuse the anterior neural folds, and that any embryos with a persistent 212 DONuT and/or whose pineal domains have not yet fused by the onset of C-divisions around 10 somite stage 213 go on to become the minority of embryos with split pineal precursors and/or roof plates at 28 hpf (Fig. 1). 214 215



Figure 3. DONuT formation is correlated with severity of convergent extension defects.

A-D') Representative images of pineal precursors (*flh* WISH) and somites/adaxial cells (*myoD* WISH) in the same control (A), *vangl2* MO-injected (B-C), and *vangl2/tri-/*- (D) embryos at the 7-somite stage, viewed dorsally. Double arrows indicate pineal precursor width (A-D) or somite width (A'-D'). **E-F**) Width of pineal precursor domains (E, as shown in A-D) and somites (F, as shown in A'-D') in control (blue), *tri-/*- (purple), and *vangl2* morphant embryos with (burgundy) or without (pink) a DONuT at 3-7 somite stages. Each dot represents a single embryo, black/gray bars are median values. ****p<0.0001, **p<0.001, **p<0.01, *p<0.05, Kruskal-Wallis test with multiple comparisons. n values indicate the number of embryos of each stage/condition measured from 3 independent trials. Anterior is up in all images.

Zebrafish neural folds fuse by zippering of distinct anterior and posterior openings.

A recent study used live time-lapse imaging to directly observe closure of the forebrain neural tube in zebrafish. 219 revealing bidirectional zippering of an eye-shaped opening between the fusing neural folds beginning at 220 approximately 6-7 somite stage (44). To characterize neural tube closure more fully in WT embryos, we 221 performed confocal time-lapse imaging of the anterior neural plate beginning at the 4-somite stage. In each of 222 the 23 control embryos imaged, we observed the presence and zippering of an eye-shaped opening in the 223 forebrain region as previously reported (44). However, examining the neural plate at earlier stages revealed a 224 sequence of preceding morphogenetic changes (Fig. 4). At around the 5-somite stage, we observed a 225 continuous keyhole-shaped groove in the neural plate midline with the round portion positioned anteriorly (Fig. 226 **4A-B**', green shading). The bilateral neural folds then elevated on either side of this groove, resembling the 227 two sides of a hot dog bun (Supp. videos 2, 5), and came together near the center of the keyhole to "pinch off" 228 the anterior and posterior portions (Fig. 4B and Supp. Fig. 3, white arrows, and Supp. videos 1-5). Elevation 229 of the neural folds was accompanied by apical constriction of midline neuroectoderm cells (see Supp. video 230 **2**), consistent with observations in the future forebrain and hindbrain (43, 44). 231

The posterior opening began zippering closed first, beginning at the "pinch point" and continuing 232 posteriorly (Fig. 4A-B', blue shading and arrows, Supp. videos 1, 3,), leaving the anterior portion to form the 233 eye-shaped opening later. In some control embryos, a small opening at the posterior end of this zipper could 234 later be seen completing closure (Fig. 4A-A'). Once the posterior opening had zippered (mostly) closed, the 235 neural folds of the anterior portion approached one another at the midline and made contact to generate the 236 posterior closure point of the eve-shaped opening (Fig. 4A and Supp. Fig. 3, vellow shading and arrows), as 237 previously described (44). The anterior closure point of the eye-shaped opening arose from the anterior-most 238 edge of the initial keyhole-shaped groove (Fig. 4 and Supp. Fig. 3, yellow arrows), and the opening zippered 239 closed predominantly from anterior to posterior (Supp. videos 4, 6). As mentioned above, the posterior-most 240 end of the posterior opening also completed its zippering at this stage. Notably, this live imaging also enabled 241 examination of the relationship between midline-crossing C-divisions and neural tube closure. In embryos in 242 which the left and right sides of the developing neural keel exhibited distinct levels of fluorescent protein 243 expression (Supp. videos 1, 3, 4, 5), cells were only seen crossing the midline after the anterior eye-shaped 244 opening had closed. This provides further evidence that neural fold fusion precedes C-divisions. 245

Together, these observations delineate a complex series of morphogenetic events that close the 246 anterior zebrafish neural tube. First, apical constriction of midline neuroectoderm cells creates a medial hinge 247 point and elevates the bilateral neural folds, producing a shallow groove along the dorsal midline. The neural 248 folds come together near the center of this groove, pinching it into anterior and posterior segments. The neural 249 folds zipper together posteriorly from the pinch point while the neural folds continue toward the midline in the 250 anterior portion of the groove, creating the previously described eye-shaped opening that then zippers shut 251 between two closure points. Around the time the anterior eye-shaped opening closes, the caudal-most end of 252 the posterior portion completes its zipper closure (see model in Fig. 7). 253

254



Figure 4. Neural fold fusion proceeds bidirectionally from a central "pinch point".

A-B') Still frames from time-lapse series of anterior neural tube development in WT or *tri* sibling embryos expressing membrane GFP or mCherry beginning at the 5-somite stage, viewed dorsally from more anterior (A-A') or posterior (B-B') positions. **A-B**) Yellow arrowheads indicate the anterior edge of the neural groove and eventually the eye-shaped opening. White arrowheads indicate the pinch-point at which the bilateral neural folds make contact. Cyan arrowheads indicate the posterior opening that zippers closed in the posterior direction from the pinch-point. **A'-B'**) Pseudo-colored versions of the images shown in (A-B). Green indicates the early neural groove before formation of the pinch-point. Thereafter, yellow and blue indicate the anterior and posterior openings, respectively. Each image series is a single Z plane from a confocal stack and is representative of 23 individual WT and sibling embryos imaged in 6 independent trials. Additional examples are shown in Supp. Fig. 3 and Supp. videos 1-5. Anterior is up in all images, scale bar = 50 µm.



Supplemental Figure 3. Live imaging reveals neural fold fusion dynamics in live WT embryos.

A-C) Still frames from time-lapse series of anterior neural tube development in WT or *tri* sibling embryos expressing membrane GFP or mCherry beginning at the 5-somite stage, viewed dorsally from more anterior (A-B) or posterior (C) positions. Yellow arrowheads indicate the anterior edge of the neural groove and eventually the eye-shaped opening. White arrowheads indicate the pinch-point at which the bilateral neural folds make contact. Cyan arrowheads indicate the posterior opening that zippers closed in the posterior direction from the pinch-point. Anterior is up in all images, scale bar = 50 μ m.

Neural fold fusion is delayed in Vangl2 deficient embryos.

To determine if the DONuT is a consequence of delayed, abnormal, or failed neural fold zippering, we 258 performed confocal time-lapse imaging of the anterior neural plate in vangl2 deficient embryos beginning at the 259 4-somite stage (Fig. 5. Supp. video 7). We observed that the bilateral neural folds of vanal2 morphant and 260 vangl2/tri-/- embryos began much farther apart than control siblings, which led to a delay in formation of the 261 anterior eye-shaped opening. Indeed, while control embryos had formed the eye-shaped opening by 6-7 262 somite stage (Fig. 5A), the neural folds of stage-matched vang/2 mutants and morphants had not yet made 263 contact, leaving wide gaps between them that were open to the posterior (Fig. 5B-D). This is reflected in 264 guantitative measurements of the distance between the neural folds over time, beginning at 6-somite stage 265 when the eve-shaped opening had formed in control embryos. A simple linear regression revealed that the 266 distance between the neural folds of both vangl2 morphants and mutants started nearly three times larger (Y 267 intercepts of 167.6 and 132 μm, respectively) than sibling control embryos (Y intercepts of 57.8 and 48.2) (Fig. 268 5E). Interestingly, the rate of neural fold convergence was significantly higher in vangl2 morphants and mutants 269 (with slopes of -0.76 and -0.59, respectively) compared with their sibling controls (slopes of -0.31 and -0.25) 270 (Fig. 5E). This accelerated closure could not fully compensate for the increased width of their neural folds, 271 however, and closure of the anterior opening was significantly delayed in vangl2 morphants and mutants (with 272 X intercepts at 220.7 and 224.9 minutes, respectively) with respect to sibling controls (X intercepts of 188.2 273 and 192.6 minutes) (Fig. 5E). 274

Additional differences in neural fold fusion were apparent from these time-lapse series. First, the 275 anterior eye-shaped opening of control embryos closed predominantly from anterior to posterior (Supp. videos 276 **4. 6**), whereas *vangl2* morphant and mutant neural folds zippered closed from posterior to anterior (**Supp**. 277 video 7). The continuous nature of the anterior and posterior openings in the neural plate was also more 278 apparent in vangl2 deficient embryos, where "pinching" at the center of the keyhole-shaped groove was more 279 dramatic (Fig. 5B, white arrows). Closure of the posterior opening was also substantially delayed and 280 sometimes blocked in vanal2 mutants and morphants. While the posterior opening was only briefly visible in 281 control embryos after 6-somite stage and zippered closed rapidly, a large and persistent opening could be 282 seen in the posterior region of essentially all vangl2 morphants and mutants examined and often had not 283 closed by the end of the imaging period at 10-somite stage (Fig. 5B-D, blue arrows). Finally, rounded cells 284 were seen protruding from the neural groove during closure (Fig. 5B-D, orange arrows), as observed in 3D 285 reconstructions of fixed vanal2 morphants (Fig. 2G). These results highlight severe and regionally distinct 286 defects in neural fold fusion in the absence of Vangl2. Because the zippering process itself was not disrupted 287 in morphant and mutant embryos, we suspect that delayed neural fold fusion is largely the consequence of 288 increased width of the neural plate that ultimately results from reduced CE (Fig. 3). 289

290 291

255



Figure 5. Neural fold fusion is delayed in vangl2 deficient embryos.

A-D) Still frames from time-lapse series of anterior neural tube development in WT (A), *tri-/-* mutant (B), and *vangl2* morphant (C) embryos expressing membrane GFP or mCherry beginning at 6-7 somite stage, viewed dorsally. Yellow arrowheads indicate the anterior edge of the eye-shaped opening, white arrowheads indicate the pinch-point, cyan arrowheads indicate the posterior opening, and orange arrowheads indicate rounded cells protruding from the neural groove of *vangl2* deficient embryos. Each image series is a single Z plane from a confocal stack and is representative of multiple embryos of that condition (see n values for each condition in E). **E**) Distance between the bilateral neural folds over time in embryos of the conditions indicated, beginning when the eye-shaped opening forms around the 6-somite stage. Symbols are mean + SEM, lines are simple linear regressions, for which slopes and intercepts are provided below. n values indicate the number of embryos measured of each condition from 4 independent *vangl2* MO and 2 independent *tri* mutant trials. Anterior is up in all images, scale bar = 50 μ m. See also Supp. videos 6-7.

293 The Forebrain neural folds fuse to enclose a lumen.

292

294

315

Neural tube closure in zebrafish and amniote embryos involves not only CE and zippering, but also formation 295 of medial and dorsolateral hinge points within the neural plate (25, 26, 29, 44, 86). Using time-lapse confocal 296 microscopy to collect optical transverse sections through the developing forebrain region, we observed hinge 297 point formation and neural fold elevation in WT embryos beginning at the 3-4 somite stage. The neural plate 298 began largely flat across the apical surface but developed a prominent medial hinge point by the 5-somite 299 stage (Fig. 6A-A')(as described in (44)). In the anterior region of the forebrain, cells lining the V-shaped neural 300 groove sealed up progressively from ventral to dorsal until the neural tube was closed and smooth across its 301 outer surface (Fig. 6A). This is apparent from measurements of medial hinge point angle, which became more 302 acute as the neural folds elevated and then widened again as the folds sealed up (Fig. 6E). Optical sections 303 through a more posterior region of the forebrain, however, showed the bilateral neural folds elevating around a 304 larger U-shaped groove and then fused at the dorsal side to enclose a lumen (Fig. 6A', Supp. video 8). This is 305 strikingly similar to the mechanisms of primary neurulation in amniote embryos and distinct from those of spinal 306 cord development in zebrafish, in which the neural tube lumen is formed through cavitation of the solid neural 307 rod (33, 40). We also observed that at this more posterior position, the periderm separated slightly from the 308 underlying ectoderm and bridged the gap between the bilateral neural folds until they fused dorsally (Fig. 6A', 309 orange arrow), which can also be observed in time-lapse series from a previous study (44). These data directly 310 demonstrate that neural folds within the forebrain region of zebrafish embryos elevate and fuse to enclose a 311 lumen, as in many other vertebrate species. 312

Neural groove formation is abnormal in *vangl2* deficient embryos.

Our live confocal imaging revealed significant delays in neural fold fusion in *vangl2* deficient embryos, but it was unclear whether this delay alone underlies DONuT formation. First, control embryos almost never exhibited a DONuT even prior to fusion of the pineal precursors and neural folds. Second, some developmentally advanced (8-somite stage) *vangl2* deficient embryos exhibited bifurcated pineal precursors in the absence of a DONuT (**Fig. 2**). Together, this implies that a DONuT is not simply the consequence of unfused pineal domains but likely reflects additional developmental abnormalities. We therefore collected transverse optical sections through the developing brain of *vangl2* mutant and morphant embryos.

The anterior forebrain regions of vangl2/ tri-/- embryos (Fig. 6B) were wider than their siblings 323 throughout neural tube development (Fig. 6D) but exhibited formation of a V-shaped groove that sealed from 324 ventral to dorsal, similar to sibling controls (Fig. 6E). The posterior forebrain also exhibited a U-shaped groove 325 with bilateral neural folds that fused dorsally, although no open lumen was apparent upon fusion (Fig. 6B'). 326 This groove was substantially larger by cross-sectional area than sibling controls (Fig. 6F), likely due to 327 increased width of the neural plate. Consistent with all other data in this study, vangl2 morphants presented 328 with a more severe phenotype than mutants. The anterior forebrain regions of morphants were even wider than 329 mutants (Fig. 6D) and exhibited neither hinge points nor V-shaped grooves, instead resembling a solid mass of 330 cells at this level (Fig. 6C. E). A large U-shaped groove was apparent in a more posterior region, but fusion of 331

these folds was significantly delayed and sometimes blocked (Fig. 6C'), as evidenced by the enlarged cross-332 sectional area of the neural groove over time (Fig. 6F). Notably, cross-sectional area of the neural groove was 333 larger in vangl2 mutants than morphants at early stages (Fig. 6F), which likely reflects the combination of 334 widened neural plates and robust hinge point formation in mutants. In both vangl2 mutants and morphants, the 335 periderm spanned the gap between neural folds in the posterior region as in control embryos, although this cell 336 layer separated from the underlying neural plate earlier and by a larger distance than in controls. These cells 337 were also highly rounded and protruded outward from the neural groove (Fig. 6B'-C', orange arrows), 338 indicating that cells observed protruding from the neural tube of fixed and live vangl2 deficient embryos (Figs. 339 2 and 5) were almost certainly periderm. These results reveal that a combination of increased neural plate 340 width and decreased bending at hinge points produce larger neural grooves in the forebrain region of vangl2 341 morphant embryos, manifesting as a DONuT. 342

343 344



Figure 6. The bilateral neural folds fuse dorsally to enclose a lumen in WT, but not *vangl2* deficient embryos.

A-C') Still frames from time-lapse series of neural fold fusion in WT (A), *tri-/-* mutant (B), and *vangl2* morphant (C) embryos expressing membrane GFP or mCherry beginning at the 4-somite stage, viewed in transverse optical section through anterior (A-C) or posterior (A'-C') regions of the brain. Orange arrowheads indicate periderm cells spanning the neural groove. Each image series is a single representative Z plane from a confocal stack. **D-F**) Measurements of neural plate width (D) and neural groove angle (E) within the anterior brain region and cross-sectional area of the neural groove (F) within the posterior brain region in embryos of the conditions indicated, beginning at 4-somite stage. Symbols are mean + SEM, n values indicate the number of embryos measured of each condition from 2 independent *vangl2* MO and 2 independent *tri* mutant trials. Images to the right are illustrative of the measurements made. Dorsal is up in all images, scale bar = 50 µm. See also Supp. videos 8-9.

346 **Discussion**

Closure of the neural tube is essential for proper development of the central nervous system, and its failure 348 leads to deadly and debilitating congenital anomalies. Primary neurulation is well described in vertebrate 349 models including mouse, chick, and Xenopus, which share a core set of cellular behaviors including 350 convergent extension of the neural plate, apical constriction at hinge points, and dorsal fusion of the bilateral 351 neural folds. Although neurulation in zebrafish embryos differs outwardly from these other species, it is 352 increasingly clear that many aspects of primary neurulation are conserved, including apical constriction of 353 neural midline cells and zippering of the neural folds (40, 43, 44). In this study, we have further characterized 354 cell and tissue behaviors driving neural tube closure in the brain region of zebrafish embryos and how these 355 behaviors are affected by loss of the NTD risk gene vangl2. 356

357

358

347

Conservation of primary neurulation

It has long been appreciated that the zebrafish trunk neural tube forms through in-folding, by which the 359 lateral edges of the neural plate come together at the dorsal surface (34, 87). This is facilitated by an 360 enrichment of myosin contractility and subsequent apical constriction of midline cells, which drives their 361 internalization (43). Although these features are common to other vertebrate embryos, they differ in that the 362 neural tube lumen forms in mice and chick when the bilateral neural fold enclose an empty space upon dorsal 363 fusion, whereas the zebrafish lumen (at the level of the trunk) forms later by cavitation of a solid rod (33). 364 Hypotheses for evolutionary drivers of this unique method of neurulation include reducing exposure of the 365 neural tube lumen to the outside environment (88) and overcoming the high mechanical stress imposed by 366 axial curvature of the embryo that could otherwise prevent elevation of the neural folds (89). However, our 367 findings provide direct evidence for neural fold elevation and enclosure of a lumen in zebrafish. Our time-lapse 368 imaging directly demonstrates that, within the future forebrain, the bilateral neural folds elevate around a 369 midline groove then fuse at the dorsal surface, leaving a hollow lumen inside (Fig. 6, Fig. 7C). This mechanism 370 is apparently unique to a portion of the forebrain region, as it was not observed in transverse images through 371 more anterior (Fig. 6, Fig. 7B) or posterior regions (33, 43, 87, 90) of the neural keel, which is likely why it was 372 not described previously. An elegant live imaging study did capture formation of hinge points and elevation of 373 neural folds in the forebrain region (44), but did not describe lumen enclosure. 374

Our findings also expand our understanding of neural fold fusion within zebrafish. The aforementioned 375 live imaging study of zebrafish forebrain (44) directly demonstrated neural fold fusion by bidirectional 376 "zippering" of an eve-shaped opening. By imaging an earlier stage of neural development, the current study 377 captures the events preceding formation of this closure point (Fig. 7). We first observe elevation of bilateral 378 neural folds to create a keyhole-shaped neural groove, which then pinches together in the center to create 379 anterior and posterior opening that zipper closed away from the pinch-point (Fig. 4). The anterior portion of the 380 groove goes on to form the previously described eye-shaped opening (44). The posterior opening has not (to 381 our knowledge) been described before, but a previous time-lapse imaging study of midbrain-hindbrain 382 boundary formation captured zippering of an opening in the hindbrain (91) that we speculate is the same 383

posterior closure point. We observed that closure at each point occurs in a reproducible order - first the central 384 pinch-point, then the anterior followed by posterior ends of the eye-shaped opening (Fig. 7) - raising the 385 possibility that they are analogous to the multiple discreet closure points where neural fold zippering initiates in 386 the mouse (30, 31). It was further shown in mice that this closure is facilitated by contact between the non-387 neural ectoderm (NNE) (31, 92), which extends protrusions that meet across the neural groove to "button up" 388 the neural folds (93). Our study did not directly address the behavior of NNE and provides no evidence for 389 such protrusions, but neuroectoderm cells were previously shown to extend filopodia and make contact with 390 cells on the contralateral side (44). We did observe, however, that the periderm overlying the developing neural 391 tube spanned the neural groove as the neural folds elevated and fused (Fig. 6). Whether this thin epithelial 392 sheet contributes to neural tube closure, similarly to the NNE of mouse embryos, will require additional 393 investigation. 394

395

396 Effect of vangl2 deficiency on neural development

Loss of the planar cell polarity protein and NTD risk gene vangl2 has dramatic effects on vertebrate 397 neural tube development. While disruption of this gene prevents neural tube closure in mice and frogs (20, 21, 398 56, 94), in zebrafish it produces double neural lumens divided by a mass of neuroectoderm cells that result 399 from abnormal midline C-divisions (38, 72). By examining a more anterior region of the developing neural tube 400 in vang/2 deficient zebrafish embryos, we have identified additional phenotypes that more closely resemble 401 those of other vertebrate species. During early neurulation, we observe pit-shaped openings in the future 402 forebrain of morphant embryos (Fig. 2) that likely reflect the combination of delayed neural fold fusion and 403 abnormal neural groove formation observed during live imaging (Figs. 5-6). We note that neural fold fusion 404 itself is not disrupted by loss of vangl2 (unlike mice (95)), and that zippering is actually accelerated in these 405 embryos (Fig. 5). Instead, we find that this delay is due to increased width of the neural plate at the time of 406 fusion and therefore, is likely a consequence of reduced CE of neuroectoderm (38, 42) (Fig. 3). Because 407 anterior neural fold fusion is complete (even in most vangl2 deficient embryos) by the onset of C-divisions (34, 408 35, 41), it is unlikely that abnormal C-divisions contribute to defects in forebrain closure. Live imaging also 409 revealed abnormal morphology of the periderm overlying the neural plate in vangl2 deficient embryos. Although 410 the periderm was also seen spanning the neural folds in WT embryos (Fig. 6), it remained relatively close to 411 the neural plate surface and its constituent cells were flat. By contrast, vangl2 deficient periderm cells were 412 rounded and protruded dramatically from the neural groove (Figs. 5, 6). Whether abnormal periderm cells are 413 contributors to, a consequence of, or unrelated to delayed neural fold fusion in these embryos is unknown. 414

- 415
- 416 417
- 418



Figure 7. Model for anterior neural tube closure in zebrafish embryos.

A) Diagram of the anterior (brain region) neural plate in WT zebrafish embryos from approximately 4-10 somite stage, viewed from the dorsal surface with anterior to the top. A shallow neural groove (dark blue) forms at the dorsal midline between the bilateral neural folds (light blue). The neural folds come together at a central "pinch point" (white arrows), creating anterior and posterior openings. The posterior opening zippers closed caudally from the pinch point (cyan arrows) and the anterior opening goes on to form an eye-shaped opening in the forebrain region. The anterior edge of the eye-shaped opening regions at the posterior opening continues to zipper toward the hindbrain until the neural folds in the entire brain region have fused. Dashed lines represent the positions of the cross-sectional views shown in (B-C). **B-C**) Cross-sectional views of anterior (B) and posterior (C) forebrain regions of the neural plate at the positions of the dashed lines in (A), dorsal is up. The anterior forebrain (B) forms a V-shaped neural groove that seals up from ventral to dorsal as the neural folds fuse. The posterior forebrain (C) forms a U-shaped neural groove between the bilateral neural folds, which approach the midline and then fuse dorsally to enclose a hollow lumen.

419 Implications for NTD modeling in zebrafish

Zebrafish embryos are highly amenable to genetic and chemical screening techniques (96-98) that 420 could enable identification of causative genetic variants and gene-environmental interactions, but their utility in 421 NTD modeling has been limited by their apparent poor resemblance to mammalian neurulation. Researchers 422 have suggested bifurcated pineal precursors as a proxy for NTDs in zebrafish embryos and have even 423 identified gene-environment interactions that exacerbate pineal defects (45, 47). However, it was not clear to 424 what extent these phenotypes resemble NTDs because A) they were not examined histologically and B) the 425 mutations that induce them (in genes encoding Nodal signaling components and N-cadherin) are not 426 associated with human NTDs. Here, we show that loss of an NTD risk gene does indeed cause widened and 427 sometimes split pineal domains in the forebrain (Fig. 1). However, we find that this external phenotype can 428 manifest with a variety of internal phenotypes, some of which show little resemblance to amniote NTDs (Supp. 429 Fig. 1). By instead examining neural development at the time of neural tube closure, we avoid the confounding 430 effects of ectopic lumen formation seen at later stages upon loss of *vangl2* or Nodal signaling (Supp. Fig. 1). 431 Indeed, the delay in neural fold fusion in vangl2 morphants is readily observed as a DONuT within fixed 432 embryos at peri-closure stages, providing an easily screen-able phenotype. Whether the DONuT occurs in 433 upon loss of other NTD risk genes, and/or whether it is common to other planar cell polarity mutants, remains 434 to be tested. Given that open neural tubes are only apparent in the brain region of zebrafish embryos, it is also 435 not clear whether this is a fitting model for only anterior NTDs (like anencephaly and craniorachischisis) or 436 whether mutations causing posterior NTDs like spina bifida would yield similar phenotypes. Taken together, this 437 study provides direct evidence for conservation of fold-and-fuse neural tube closure within zebrafish, 438 highlighting their potential for NTD modeling. 439

440 441

442 443

Materials and Methods

444 Zebrafish

Adult zebrafish were maintained through established protocols (*99*) in compliance with the Baylor College of
 Medicine Institutional Animal Care and Use Committee. Embryos were obtained through natural mating and
 staging was based on established morphology (*100*). Studies were conducted using AB WT,
 tdgf1/oep^{tz257}(101), *vangl2/ trilobite^{vu67} (102)*, and TgBAC[*flh:flh-kaede*] (*82*) embryos. Fish were crossed from

- their home tank at random and embryos were chosen for injection and inclusion in experiments at random.
- 450

451 Microinjection of synthetic mRNA and morpholino oligonucleotides

Single-celled embryos were placed in agarose molds (Adaptive Science tools I-34) and injected with 0.5-2 nL
volumes using pulled glass needles (Fisher Sci #50-821-984). mRNAs were transcribed using the SP6
mMessage mMachine kit (Fisher Sci #AM1340) and purified using Biorad Microbiospin columns (Biorad
#7326250). Each embryo was injected with 100 pg memGFP or 200 pg mCherry mRNA, and/or 2 ng *vangl2*MO-4 ((*83*) sequence: 5' - AGTTCCACCTTACTCCTGAGAGAAT - 3').

458 Whole mount in situ hybridization

Antisense riboprobes were transcribed using NEB T7 or T3 RNA polymerase (NEB #M0251s and Fisher 459 #501047499) and labeled with digoxygenin (DIG) NTPs (Sigma/Millipore #11277073910). Whole mount in situ 460 hybridization (WISH) was performed according to (103) with minor modifications. Embryos were fixed as 461 described above, washed in PBS + 0.1% Tween-20 (PBT), gradually dehydrated, and stored in methanol at -462 20C. Immediately prior to staining, embryos were rehydrated into PBT and hybridized overnight with antisense 463 probes within the wells of a 24-well plate. Embryos were gradually washed into SSC buffer and then into PBT 464 before overnight incubation with an anti-DIG primary antibody at 1:5000 (Roche #11093274910). Embryos 465 were washed in PBT and then staining buffer before developing in BM Purple staining solution (Roche 466 #11442074001). Embryos were washed and stored in stop buffer (10 mM EDTA in PBT) until imaging. 467

468

469 Histology

After whole mount in situ hybridization, the head regions of 28 hpf embryos were isolated, mounted in Tissue
Tek O.C.T. medium (VWR #25608-930) within plastic base molds, and snap frozen in liquid nitrogen. 14 μm
serial sections were cut and collected by the Baylor College of Medicine RNA In Situ Hybridization Core.
Sections were mounted under coverslips and imaged using a Nikon Fi3 color camera on the Nikon ECLIPSE
Ti2 microscope described below.

475

476 Inhibitor treatments

Nodal inhibitor SB505124 (VWR #103540-834) was stored as a 10 mM stock in DMSO at 4 C. Embryos were
dechorionated prior to treatment with 50 μM SB505124 in 0.3x Danieau's solution at the stages indicated.
Embryos were incubated at 28.5 C within the agarose-coated wells of a 6-well plate until 28 hpf, at which time
they were fixed as described above and processed for WISH.

481

482 Phalloidin staining

Embryos were fixed as described above, rinsed in PBT, and either stained directly or dehydrated and stored at
-20 C in ethanol for later use. Embryos were rehydrated in PBS + 0.1% Triton-X (PBTr) immediately before
staining and incubated with Alexa Fluor 546 Phalloidin (ThermoFisher A22283) in PBTr for several hours.
Embryos were rinsed in PBTr and mounted for confocal imaging as described below.

487

488 Microscopy

Fixed phalloidin stained embryos were mounted in 3% methylcellulose, and live embryos were mounted in
0.35% low-melt agarose (ThermoFisher #16520100) in glass bottomed 35 mm petri dishes (Fisher Sci
#FB0875711YZ) prior to imaging. Confocal Z-stacks were collected using a Nikon ECLIPSE Ti2 confocal
microscope equipped with a Yokogawa W1 spinning disk unit, PFS4 camera, and 405/488/561nm lasers
(emission filters: 455/50, 525/36, 605/52). Confocal Z-stacks were obtained with a 1 μm (fixed embryos) or 2
μm (live embryos) step size using a Plan Apo Lambda 20X lens. For time-lapse series, embryos were
maintained at 28.5 C in a Tokai Hit STX stage top incubator and Z-stacks were collected at 5-minute intervals.

Images of WISH-stained embryos were taken with a Nikon Fi3 color camera on a Nikon SMZ745T
 stereoscope.

498

499 Image analysis

ImageJ/FIJI was used to visualize and measure all microscopy data sets. Researchers were blinded to the conditions of all image data using the *blind_renamer* Perl script prior to analysis. Measurements of embryonic structures from fixed embryos, including pineal precursors and somites, were made by drawing a line from one side of the structure to the other at its widest point. Distance between neural folds, neural plate width, and neural grove angle and cross-sectional area were measured similarly from images of live embryos. 3D projections of confocal z-stacks were made using the '3D project' plugin.

506

507 Statistical Analysis

Graphpad Prism 10 software was used to perform statistical analyses and to generate graphs for the data
 collected during image analysis. Datasets were tested for normality prior to analysis and statistical tests were
 chosen accordingly. The statistical tests used for each data set are noted in figure legends.

511

512

513 Acknowledgements

We thank Dr. Patrick Blader for generously sharing the Tg[*flh:kaede*] fish line and Dr. Lila Solnica-Krezel for
sharing additional fish lines and plasmids. Histology services were provided by Dr. Cecilia Ljungberg and Rong
Jyh Kao of the BCM RNA In Situ Hybridization Core and animal care was provided by the BCM Center for
Comparative Medicine. We thank Drs. Rachel Brewster, Dan Gorelick, and Ryan Gray for helpful discussions
and comments on the manuscript. Thanks to Williams lab members for technical assistance, support, and
feedback on this project.

522 Competing Interests

523 The authors declare no competing interests.

524

520 521

525

526 Funding

This work was supported by National Institutes of Health R00HD091386 and R01HD104784 to MLKW, and a P30ES030285 (PI: Dr. Cheryl Walker) pilot grant to MLKW. The project was supported in part by the RNA In Situ Hybridization Core facility at Baylor College of Medicine, which is supported by a shared Instrumentation grant from the National Institutes of Health (1S10OD016167).

- 531
- 532
- 534

535 Supplemental video descriptions

536

540

544

548

556

560

564

568

572 573 574

575 576

Supp. video 1: Anterior view of neural fold fusion in the forebrain region of a wild-type zebrafish embryo
 labeled with membrane-GFP. Movie begins at the 4-5 somite stage and each frame = 5 minutes. Video shows
 a single Z plane of a 3D confocal time series.

Supp. video 2: Posterior view of neural fold fusion in the forebrain region of a wild-type zebrafish embryo
 labeled with membrane-GFP. Movie begins at the 4-5 somite stage and each frame = 5 minutes. Video shows
 a single Z plane of a 3D confocal time series.

Supp. video 3: Anterior view of neural fold fusion in the forebrain region of a wild-type zebrafish embryo
 labeled with membrane-GFP. Movie begins at the 4-5 somite stage and each frame = 5 minutes. Video shows
 a single Z plane of a 3D confocal time series.

Supp. video 4: Anterior view of neural fold fusion in the forebrain region of a wild-type zebrafish embryo
 labeled with membrane-GFP. Movie begins at the 4-5 somite stage and each frame = 5 minutes. Video shows
 a single Z plane of a 3D confocal time series.

Supp. video 5: Posterior view of neural fold fusion in the forebrain region of a wild-type zebrafish embryo
 labeled with membrane-GFP. Movie begins at the 4-5 somite stage and each frame = 5 minutes. Video shows
 a single Z plane of a 3D confocal time series.

557 **Supp. video 6**: Neural fold fusion in the forebrain region of a wild-type zebrafish embryo labeled with 558 membrane-GFP. Movie begins at the 6-7 somite stage and each frame = 5 minutes. Video shows a single Z 559 plane of a 3D confocal time series.

Supp. video 7: Neural fold fusion in the forebrain region of a *vangl2* morphant zebrafish embryo labeled with membrane-GFP. Movie begins at the 6-7 somite stage and each frame = 5 minutes. Video shows a single Z plane of a 3D confocal time series.

Supp. video 8: Optical transverse section through the posterior forebrain region of a wild-type zebrafish embryo labeled with membrane-GFP. Movie begins at the 4-somite stage and each frame = 5 minutes. Video shows a single Z plane of a 3D confocal time series.

Supp. video 9: Optical transverse section through the posterior forebrain region of a *vangl2* morphant zebrafish embryo labeled with membrane-GFP. Movie begins at the 4-somite stage and each frame = 5 minutes. Video shows a single Z plane of a 3D confocal time series.

References

- 577 1. S. E. Parker *et al.*, Updated National Birth Prevalence estimates for selected birth defects in the United 578 States, 2004-2006. *Birth Defects Res A Clin Mol Teratol* **88**, 1008-1016 (2010).
- 579 2. I. Zaganjor *et al.*, Describing the Prevalence of Neural Tube Defects Worldwide: A Systematic Literature 580 Review. *PLoS One* **11**, e0151586 (2016).
- 3. J. Williams *et al.*, Updated estimates of neural tube defects prevented by mandatory folic Acid fortification - United States, 1995-2011. *MMWR Morb Mortal Wkly Rep* **64**, 1-5 (2015).
- 4. M. E. Ross, C. E. Mason, R. H. Finnell, Genomic approaches to the assessment of human spina bifida risk. *Birth Defects Res* **109**, 120-128 (2017).
- 585 5. J. G. Joó *et al.*, Neural tube defects in the sample of genetic counselling. *Prenat Diagn* **27**, 912-921 (2007).
- 5876.K. L. Deak *et al.*, Further evidence for a maternal genetic effect and a sex-influenced effect contributing588to risk for human neural tube defects. *Birth Defects Res A Clin Mol Teratol* **82**, 662-669 (2008).
- 589 7. L. Wang *et al.*, Digenic variants of planar cell polarity genes in human neural tube defect patients. *Mol* 590 *Genet Metab* **124**, 94-100 (2018).

- 8. I. Matok *et al.*, Exposure to folic acid antagonists during the first trimester of pregnancy and the risk of major malformations. *Br J Clin Pharmacol* **68**, 956-962 (2009).
- M. Artama, A. Auvinen, T. Raudaskoski, I. Isojärvi, J. Isojärvi, Antiepileptic drug use of women with
 epilepsy and congenital malformations in offspring. *Neurology* 64, 1874-1878 (2005).
- J. Gelineau-van Waes *et al.*, Maternal fumonisin exposure and risk for neural tube defects:
 mechanisms in an in vivo mouse model. *Birth Defects Res A Clin Mol Teratol* **73**, 487-497 (2005).

591

- 597 11. E. Gamero-Estevez, A. I. Baumholtz, A. K. Ryan, Developing a link between toxicants, claudins and 598 neural tube defects. *Reprod Toxicol* **81**, 155-167 (2018).
- R. P. Rull, B. Ritz, G. M. Shaw, Neural tube defects and maternal residential proximity to agricultural pesticide applications. *Am J Epidemiol* **163**, 743-753 (2006).
- 60113.Ş. Özel *et al.*, Maternal second trimester blood levels of selected heavy metals in pregnancies602complicated with neural tube defects. J Matern Fetal Neonatal Med **32**, 2547-2553 (2019).
- N. Demir *et al.*, The relationship between mother and infant plasma trace element and heavy metal
 levels and the risk of neural tube defect in infants. *J Matern Fetal Neonatal Med* 32, 1433-1440 (2019).
- S. Kalra *et al.*, Organochlorine pesticide exposure in mothers and neural tube defects in offsprings.
 Reprod Toxicol 66, 56-60 (2016).
- 16. C. Kalliora *et al.*, Association of pesticide exposure with human congenital abnormalities. *Toxicol Appl Pharmacol* **346**, 58-75 (2018).
- M. J. Harris, D. M. Juriloff, Mouse mutants with neural tube closure defects and their role in
 understanding human neural tube defects. *Birth Defects Res A Clin Mol Teratol* **79**, 187-210 (2007).
- 18. I. E. Zohn, A. A. Sarkar, Modeling neural tube defects in the mouse. *Curr Top Dev Biol* 84, 1-35 (2008).
- I. E. Zohn, Mouse as a model for multifactorial inheritance of neural tube defects. *Birth Defects Res C Embryo Today* 96, 193-205 (2012).
- 614 20. M. Williams, W. Yen, X. Lu, A. Sutherland, Distinct apical and basolateral mechanisms drive planar cell 615 polarity-dependent convergent extension of the mouse neural plate. *Dev Cell* **29**, 34-46 (2014).
- P. Ybot-Gonzalez *et al.*, Convergent extension, planar-cell-polarity signalling and initiation of mouse
 neural tube closure. *Development* **134**, 789-799 (2007).
- R. Massarwa, H. J. Ray, L. Niswander, Morphogenetic movements in the neural plate and neural tube:
 mouse. *Wiley Interdiscip Rev Dev Biol* 3, 59-68 (2014).
- I. E. Zohn, C. R. Chesnutt, L. Niswander, Cell polarity pathways converge and extend to regulate neural
 tube closure. *Trends Cell Biol* 13, 451-454 (2003).
- 622 24. G. C. Schoenwolf, Cell movements driving neurulation in avian embryos. *Development* **Suppl 2**, 157-623 168 (1991).
- A. S. Shum, A. J. Copp, Regional differences in morphogenesis of the neuroepithelium suggest multiple
 mechanisms of spinal neurulation in the mouse. *Anat Embryol (Berl)* **194**, 65-73 (1996).
- 26. J. L. Smith, G. C. Schoenwolf, J. Quan, Quantitative analyses of neuroepithelial cell shapes during bending of the mouse neural plate. *J Comp Neurol* **342**, 144-151 (1994).
- L. A. Davidson, R. E. Keller, Neural tube closure in Xenopus laevis involves medial migration, directed protrusive activity, cell intercalation and convergent extension. *Development* **126**, 4547-4556 (1999).
- S. L. Haigo, J. D. Hildebrand, R. M. Harland, J. B. Wallingford, Shroom induces apical constriction and
 is required for hingepoint formation during neural tube closure. *Curr Biol* **13**, 2125-2137 (2003).
- S. G. McShane *et al.*, Cellular basis of neuroepithelial bending during mouse spinal neural tube closure.
 Dev Biol **404**, 113-124 (2015).
- R. Massarwa, L. Niswander, In toto live imaging of mouse morphogenesis and new insights into neural
 tube closure. *Development* 140, 226-236 (2013).
- G36 31. C. Pyrgaki, P. Trainor, A. K. Hadjantonakis, L. Niswander, Dynamic imaging of mammalian neural tube
 G37 closure. *Dev Biol* 344, 941-947 (2010).
- M. L. Concha, R. J. Adams, Oriented cell divisions and cellular morphogenesis in the zebrafish gastrula and neurula: a time-lapse analysis. *Development* **125**, 983-994 (1998).
- B. Schmitz, C. Papan, J. A. Campos-Ortega, Neurulation in the anterior trunk region of the zebrafish
 Brachydanio rerio. *Roux Arch Dev Biol* 202, 250-259 (1993).
- B. Geldmacher-Voss, A. M. Reugels, S. Pauls, J. A. Campos-Ortega, A 90-degree rotation of the mitotic spindle changes the orientation of mitoses of zebrafish neuroepithelial cells. *Development* 130, 3767-3780 (2003).
- G45 35. C. E. Buckley *et al.*, Mirror-symmetric microtubule assembly and cell interactions drive lumen formation
 in the zebrafish neural rod. *EMBO J* 32, 30-44 (2013).

36. C. Buckley, J. Clarke, Establishing the plane of symmetry for lumen formation and bilateral brain formation in the zebrafish neural rod. *Semin Cell Dev Biol* **31**, 100-105 (2014).

647

- 37. J. Clarke, Role of polarized cell divisions in zebrafish neural tube formation. *Curr Opin Neurobiol* **19**, 134-138 (2009).
- 38. M. Tawk *et al.*, A mirror-symmetric cell division that orchestrates neuroepithelial morphogenesis. *Nature* 446, 797-800 (2007).
- 653 **39**. E. Quesada-Hernández *et al.*, Stereotypical cell division orientation controls neural rod midline 654 formation in zebrafish. *Curr Biol* **20**, 1966-1972 (2010).
- 40. L. A. Lowery, H. Sive, Strategies of vertebrate neurulation and a re-evaluation of teleost neural tube formation. *Mech Dev* **121**, 1189-1197 (2004).
- 41. C. B. Kimmel, R. M. Warga, D. A. Kane, Cell cycles and clonal strings during formation of the zebrafish central nervous system. *Development* **120**, 265-276 (1994).
- 42. J. R. Jessen *et al.*, Zebrafish trilobite identifies new roles for Strabismus in gastrulation and neuronal movements. *Nat Cell Biol* **4**, 610-615 (2002).
- 43. C. Araya *et al.*, Cdh2 coordinates Myosin-II dependent internalisation of the zebrafish neural plate. *Sci Rep* **9**, 1835 (2019).
- 44. J. M. Werner *et al.*, Hallmarks of primary neurulation are conserved in the zebrafish forebrain. *Commun* Biol **4**, 147 (2021).
- 45. A. Aquilina-Beck, K. Ilagan, Q. Liu, J. O. Liang, Nodal signaling is required for closure of the anterior neural tube in zebrafish. *BMC Dev Biol* **7**, 126 (2007).
- 46. N. Gonsar *et al.*, Temporal and spatial requirements for Nodal-induced anterior mesendoderm and mesoderm in anterior neurulation. *Genesis* **54**, 3-18 (2016).
- P. Ma, M. R. Swartz, L. M. Kindt, A. M. Kangas, J. O. Liang, Temperature Sensitivity of Neural Tube
 Defects in Zoep Mutants. *Zebrafish* 12, 448-456 (2015).
- 48. Z. Lele *et al.*, parachute/n-cadherin is required for morphogenesis and maintained integrity of the zebrafish neural tube. *Development* **129**, 3281-3294 (2002).
- 49. J. M. de la Cruz *et al.*, A loss-of-function mutation in the CFC domain of TDGF1 is associated with human forebrain defects. *Hum Genet* **110**, 422-428 (2002).
- 50. K. W. Gripp *et al.*, Mutations in TGIF cause holoprosencephaly and link NODAL signalling to human neural axis determination. *Nat Genet* **25**, 205-208 (2000).
- E. Roessler *et al.*, Reduced NODAL signaling strength via mutation of several pathway members
 including FOXH1 is linked to human heart defects and holoprosencephaly. *Am J Hum Genet* 83, 18-29
 (2008).
- 680 52. W. W. Yen *et al.*, PTK7 is essential for polarized cell motility and convergent extension during mouse 681 gastrulation. *Development*, (2009).
- 53. X. Lu *et al.*, PTK7/CCK-4 is a novel regulator of planar cell polarity in vertebrates. *Nature* **430**, 93-98 (2004).
- M. Montcouquiol *et al.*, Identification of Vangl2 and Scrb1 as planar polarity genes in mammals. *Nature* 423, 173-177 (2003).
- 55. Y. Wang, N. Guo, J. Nathans, The role of Frizzled3 and Frizzled6 in neural tube closure and in the planar polarity of inner-ear sensory hair cells. *J Neurosci* **26**, 2147-2156 (2006).
- 56. N. D. Greene, D. Gerrelli, H. W. Van Straaten, A. J. Copp, Abnormalities of floor plate, notochord and
 somite differentiation in the loop-tail (Lp) mouse: a model of severe neural tube defects. *Mech Dev* 73,
 59-72 (1998).
- 57. Z. Kibar *et al.*, Ltap, a mammalian homolog of Drosophila Strabismus/Van Gogh, is altered in the mouse neural tube mutant Loop-tail. *Nat Genet* **28**, 251-255 (2001).
- 58. J. N. Murdoch *et al.*, Disruption of scribble (Scrb1) causes severe neural tube defects in the circletail mouse. *Hum Mol Genet* **12**, 87-98 (2003).
- 59. T. Goto, L. Davidson, M. Asashima, R. Keller, Planar cell polarity genes regulate polarized extracellular matrix deposition during frog gastrulation. *Curr Biol* **15**, 787-793 (2005).
- 697 60. J. B. Wallingford, R. M. Harland, Neural tube closure requires Dishevelled-dependent convergent 698 extension of the midline. *Development* **129**, 5815-5825 (2002).
- 699 61. J. Wang *et al.*, Dishevelled genes mediate a conserved mammalian PCP pathway to regulate 700 convergent extension during neurulation. *Development* **133**, 1767-1778 (2006).
- 62. O. Ossipova, K. Kim, S. Y. Sokol, Planar polarization of Vangl2 in the vertebrate neural plate is controlled by Wnt and Myosin II signaling. *Biol Open* 4, 722-730 (2015).

- 70363.T. Nishimura, H. Honda, M. Takeichi, Planar cell polarity links axes of spatial dynamics in neural-tube704closure. Cell 149, 1084-1097 (2012).
- ⁷⁰⁵ 64. J. A. Curtin *et al.*, Mutation of Celsr1 disrupts planar polarity of inner ear hair cells and causes severe ⁷⁰⁶ neural tube defects in the mouse. *Curr Biol* **13**, 1129-1133 (2003).
- R. S. Darken *et al.*, The planar polarity gene strabismus regulates convergent extension movements in Xenopus. *EMBO J* 21, 976-985 (2002).
- 709 66. Z. Chen *et al.*, Genetic analysis of Wnt/PCP genes in neural tube defects. *BMC Med Genomics* **11**, 38
 (2018).
- 711 67. Y. Lei *et al.*, Mutations in planar cell polarity gene SCRIB are associated with spina bifida. *PLoS One* **8**, e69262 (2013).
- 713 68. Y. Lei *et al.*, Identification of novel CELSR1 mutations in spina bifida. *PLoS One* **9**, e92207 (2014).
- 71469.T. Tian *et al.*, Somatic mutations in planar cell polarity genes in neural tissue from human fetuses with715neural tube defects. *Hum Genet* **139**, 1299-1314 (2020).
- 716 **70.** T. Tian *et al.*, Rare copy number variations of planar cell polarity genes are associated with human neural tube defects. *Neurogenetics* **21**, 217-225 (2020).
- 718 **71**. Z. Kibar *et al.*, Contribution of VANGL2 mutations to isolated neural tube defects. *Clin Genet* **80**, 76-82 (2011).
- 720 72. B. Ciruna, A. Jenny, D. Lee, M. Mlodzik, A. F. Schier, Planar cell polarity signalling couples cell division 721 and morphogenesis during neurulation. *Nature* **439**, 220-224 (2006).
- 722 73. F. Carreira-Barbosa *et al.*, Prickle 1 regulates cell movements during gastrulation and neuronal 723 migration in zebrafish. *Development* **130**, 4037-4046 (2003).
- 724 **74**. C. P. Heisenberg *et al.*, Silberblick/Wnt11 mediates convergent extension movements during zebrafish gastrulation. *Nature* **405**, 76-81 (2000).
- 726 **75**. B. Kilian *et al.*, The role of Ppt/Wnt5 in regulating cell shape and movement during zebrafish gastrulation. *Mech Dev* **120**, 467-476 (2003).
- 728 76. F. Ulrich *et al.*, Slb/Wnt11 controls hypoblast cell migration and morphogenesis at the onset of zebrafish 729 gastrulation. *Development* **130**, 5375-5384 (2003).
- 730 **77.** D. Čapek *et al.*, Light-activated Frizzled7 reveals a permissive role of non-canonical wnt signaling in 731 mesendoderm cell migration. *Elife* **8**, (2019).
- 732 78. L. Solnica-Krezel *et al.*, Mutations affecting cell fates and cellular rearrangements during gastrulation in 733 zebrafish. *Development* **123**, 67-80 (1996).
- 734 **79**. J. Topczewski *et al.*, The zebrafish glypican knypek controls cell polarity during gastrulation movements 735 of convergent extension. *Dev Cell* **1**, 251-264 (2001).
- 73680.F. Carreira-Barbosa *et al.*, Flamingo regulates epiboly and convergence/extension movements through737cell cohesive and signalling functions during zebrafish gastrulation. Development **136**, 383-392 (2009).
- 73881.Y. Y. Xing *et al.*, Mutational analysis of dishevelled genes in zebrafish reveals distinct functions in739embryonic patterning and gastrulation cell movements. *PLoS Genet* **14**, e1007551 (2018).
- J. A. Clanton, K. D. Hope, J. T. Gamse, Fgf signaling governs cell fate in the zebrafish pineal complex.
 Development 140, 323-332 (2013).
- B. B. Williams *et al.*, VANGL2 regulates membrane trafficking of MMP14 to control cell polarity and migration. *J Cell Sci* **125**, 2141-2147 (2012).
- 74484.M. Park, R. T. Moon, The planar cell-polarity gene stbm regulates cell behaviour and cell fate in
vertebrate embryos. *Nat Cell Biol* **4**, 20-25 (2002).
- Resource and the second second
- 74886.J. L. Smith, G. C. Schoenwolf, Cell cycle and neuroepithelial cell shape during bending of the chick749neural plate. Anat Rec 218, 196-206 (1987).
- 75087.C. Papan, J. A. Campos-Ortega, On the formation of the neural keel and neural tube in the
zebrafishDanio (Brachydanio) rerio. *Rouxs Arch Dev Biol* **203**, 178-186 (1994).
- M. J. Harrington, E. Hong, R. Brewster, Comparative analysis of neurulation: first impressions do not count. *Mol Reprod Dev* **76**, 954-965 (2009).
- Revealed a constraint of the second se
- 75690.M. J. Harrington, K. Chalasani, R. Brewster, Cellular mechanisms of posterior neural tube757morphogenesis in the zebrafish. Dev Dyn 239, 747-762 (2010).

- 91. G. Kesavan, A. Machate, S. Hans, M. Brand, Cell-fate plasticity, adhesion and cell sorting
 complementarily establish a sharp midbrain-hindbrain boundary. *Development* 147, (2020).
- P2. E. Nikolopoulou *et al.*, Spinal neural tube closure depends on regulation of surface ectoderm identity and biomechanics by Grhl2. *Nat Commun* **10**, 2487 (2019).
- H. J. Ray, L. A. Niswander, Dynamic behaviors of the non-neural ectoderm during mammalian cranial
 neural tube closure. *Dev Biol* 416, 279-285 (2016).
- 764 94. T. Goto, R. Keller, The planar cell polarity gene strabismus regulates convergence and extension and 765 neural fold closure in Xenopus. *Dev Biol* **247**, 165-181 (2002).
- 95. D. Gerrelli, A. J. Copp, Failure of neural tube closure in the loop-tail (Lp) mutant mouse: analysis of the embryonic mechanism. *Brain Res Dev Brain Res* **102**, 217-224 (1997).
- A. Burger *et al.*, Maximizing mutagenesis with solubilized CRISPR-Cas9 ribonucleoprotein complexes.
 Development 143, 2025-2037 (2016).
- 770 97. R. S. Wu *et al.*, A Rapid Method for Directed Gene Knockout for Screening in G0 Zebrafish. *Dev Cell* 771 46, 112-125.e114 (2018).
- 772 98. K. Bambino, J. Chu, Zebrafish in Toxicology and Environmental Health. *Curr Top Dev Biol* **124**, 331-367
 (2017).
- 99. W. M. (University of Oregon Press, 1993).
- 100. C. B. Kimmel, W. W. Ballard, S. R. Kimmel, B. Ullmann, T. F. Schilling, Stages of embryonic development of the zebrafish. *Dev Dyn* **203**, 253-310 (1995).
- 101. M. Hammerschmidt *et al.*, Mutations affecting morphogenesis during gastrulation and tail formation in the zebrafish, Danio rerio. *Development* **123**, 143-151 (1996).
- 779102.X. Li *et al.*, Gpr125 modulates Dishevelled distribution and planar cell polarity signaling. *Development*780**140**, 3028-3039 (2013).
- 103. C. Thisse, B. Thisse, High-resolution in situ hybridization to whole-mount zebrafish embryos. *Nat Protoc* 3, 59-69 (2008).
- 783