

Development/Plasticity/Repair

Neogenin and RGMA Control Neural Tube Closure and Neuroepithelial Morphology by Regulating Cell Polarity

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In humans, neural tube closure defects occur in 1:1000 pregnancies. The design of new strategies for the prevention of such common defects would benefit from an improved understanding of the molecular events underlying neurulation. Neural fold elevation is a key morphological process that acts during neurulation to drive neural tube closure. However, to date, the molecular pathways underpinning neural fold elevation have not been elucidated. Here, we use morpholino knock-down technology to demonstrate that Repulsive Guidance Molecule (RGMA)-Neogenin interactions are essential for effective neural fold elevation during *Xenopus* neurulation and that loss of these molecules results in disrupted neural tube closure. We demonstrate that Neogenin and RGMA are required for establishing the morphology of deep layer cells in the neural plate throughout neurulation. We also show that loss of Neogenin severely disrupts the microtubule network within the deep layer cells suggesting that Neogenin-dependent microtubule organization within the deep cells is essential for radial intercalation with the overlying superficial cell layer, thereby driving neural fold elevation. In addition, we show that sustained Neogenin activity is also necessary for the establishment of the apicobasally polarized pseudostratified neuroepithelium of the neural tube. Therefore, our study identifies a novel signaling pathway essential for radial intercalation and epithelialization during neural fold elevation and neural tube morphogenesis.

Key words: Neogenin; RGMA; cell polarity; neuroepithelium; neural tube; neurulation

Introduction

The earliest neural structure in the vertebrate is the neural tube comprising a layer of pseudostratified epithelium surrounding a central lumen. The neural tube derives from a flat sheet of neuroectoderm (neural plate) arising on the dorsal surface of the blastoderm at the end of gastrulation (Lowery and Sive, 2004; Copp, 2005; Wallingford, 2005). At the onset of neurulation, the lateral regions of the neural plate elevate producing the neural folds (NFs). As the folds continue to elevate and migrate toward the dorsal midline they bend around hinge-points forming the neural groove. Finally, the dorsal tips of the NFs fuse over the midline, generating a closed neural tube. Overall, progression through neurulation requires tight spatiotemporal regulation of cell polarity, shape, and motility as well as cell–cell contact.

Much of our current understanding of neural tube formation comes from the study of *Xenopus* neurulation. Throughout posterior (hindbrain, spinal cord) neurulation in the *Xenopus* embryo, convergent extension (the process by which intercalation of the neuroectodermal and underlying mesenchymal cells narrows

the embryo along the mediolateral axis while lengthening the anteroposterior axis) reduces the distance between the NFs allowing apposition at the midline (Keller et al., 1992; Wallingford and Harland, 2002). Failure of this process results in posterior neural tube closure defects. However, convergent extension contributes little to anterior neural tube closure (Keller et al., 1992; Copp, 2005; Wallingford, 2005). Instead, NF elevation and neural plate bending are the primary forces driving anterior neurulation. Neural plate bending is generated by the constriction of actin filaments under the apical surface of the neuroectodermal cells (apical constriction) leading to the formation of hinge-points on either side of the dorsal midline (Haigo et al., 2003; Lee et al., 2007). Inability to undergo apical constriction results in failure of anterior neural tube closure (anencephaly) (Hildebrand and Soriano, 1999; Haigo et al., 2003). To date, the molecular mechanisms promoting NF elevation have not been identified. Here we demonstrate that NF elevation is dependent on the axon guidance receptor, Neogenin (Vielmetter et al., 1994; Keeling et al., 1997; De Vries and Cooper, 2008).

Neogenin has been shown to act as a chemoattractive axon guidance receptor upon Netrin-1 binding, and a chemorepulsive receptor when binding Repulsive Guidance Molecule (RGMA) (Rajagopalan et al., 2004; Wilson and Key, 2006). Neogenin is also required for adhesion between multipotent progenitor cells and the adjacent epithelial layer of the developing mammary gland (Srinivasan et al., 2003). In addition, Netrin-1-Neogenin interactions promote adhesion and migration of vascular smooth muscle cells, and induce myotube formation *in vitro* (Kang et al., 2004; Park et al., 2004). Evidence that Neogenin is required for successful neurulation comes from our previous study in the ze-

Received Sept. 5, 2008; revised Oct. 7, 2008; accepted Oct. 15, 2008.

This work was supported by the National Health and Medical Research Council of Australia. N.W., M.D.V., and D.B. were supported by Australian Postgraduate Awards. The E7 and Xen1 antibodies, developed by M. Klymkowsky and Ruiz i Altaba, respectively, were obtained from the Developmental Studies Hybridoma Bank developed under the auspices of the National Institute of Child Health and Human Development and maintained by The University of Iowa (Iowa City, IA). We thank Rowan Tweedale for her critical reading of this manuscript and Grace J. Shin Lah for assistance with the generation of embryos.

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DOI:10.1523/JNEUROSCI.4265-08.2008

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brafish where loss of Neogenin resulted in a severe disruption of neural tube morphology (Mawdsley et al., 2004). Here, we demonstrate that RGMa and Neogenin are essential for neural tube formation as they establish apicobasal polarity during NF elevation and also in the neuroepithelium after neural tube closure.

Materials and Methods

In situ hybridization

Xenopus laevis borealis embryos were obtained and reared as previously described (Connor and Key, 2002). Embryos were staged according to Nieuwkoop and Faber (1994). The use of animals as described here was approved by the Animal Ethics Committee of the University of Queensland in accordance with the guidelines stipulated by the National Health and Medical Research Council of Australia. A 3.1kb 5' *Xneog1a* cDNA fragment (Wilson and Key, 2006) was used as the template for the synthesis of the riboprobes. It should be noted that two distinct *Xneog* genes (*Xneog1a* and *Xneog1b*) have been identified in the *Xenopus* genome (N. H. Wilson and B. Key, unpublished observations). The riboprobe used in this analysis does not discriminate between the two. The DIG RNA Labeling Kit (Roche Diagnostics) was used to synthesize the digoxigenin (DIG)-labeled riboprobes. Embryos were fixed with 4% paraformaldehyde (PFA) and *in situ* hybridization was performed as described in the DIG Application Manual for Nonradioactive *In situ* Hybridization (Roche). Bound riboprobe was visualized using an anti-DIG-AP Fab fragment (Roche) followed by the BM Purple chromogenic reaction (Roche).

Morpholinos and cRNA

The target sequences of all *Xenopus* morpholinos (MOs) are listed in supplemental Figure 1, available at www.jneurosci.org as supplemental material. Additional specificity controls included XplexinA and Xneuropilin1 MOs (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Zebrafish morpholinos are described by Mawdsley et al. (2004). MOs were synthesized by GeneTools. The standard control MO (ContMO) from GeneTools was used as a nonspecific control. Capped RNA (cRNA) encoding full-length *Xneog1b* was synthesized using the mMessage mMachine cRNA Transcription Kit (Ambion).

Blastomere injections and phenotype analysis

Xenopus. Fertilized eggs were collected and two-cell stage blastomeres injected as previously described (Connor and Key, 2002). The neurulation delay in the morphants was quantitated as follows: the number of morphants at each embryonic stage was determined when 90% of ContMO embryos had reached stage 18 (st18). The mean number of delayed embryos over a minimum of three independent experiments was then expressed as a percentage of the total number of embryos injected with that MO (\pm SEM). The delay phenotype was analyzed using ANOVA followed by Bonferroni's or Dunnett's *post hoc* test, or by the Kruskal–Wallis test followed by the Mann–Whitney test.

Zebrafish. Zebrafish embryos were injected with 2 ng of zfccont or zfneogATG MO as previously described (Mawdsley et al., 2004).

Immunohistochemistry

The mouse anti- β -tubulin antibody (E7) (1:150) and the neural specific Xen1 antibody (1:500) were obtained from the Developmental Studies Hybridoma Bank (University of Iowa, Iowa City, IA). *Xenopus* embryos were fixed in 80% methanol, 20% DMSO and incubated for 2 h in block-

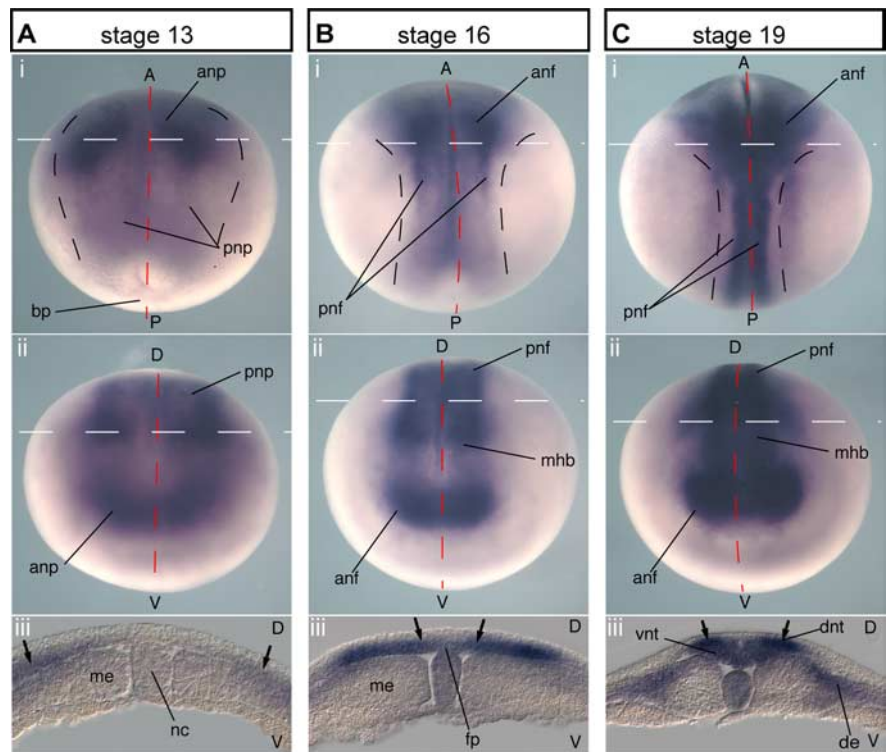


Figure 1. *Xneog* is expressed in the deep layer of the lateral and intermediate neural plate during neurulation. *Xneog* is expressed (A) in the deep layer of the lateral regions of the st13 posterior neural plate (pnp) (Ai, Aiii, arrows) and the anterior neural plate (anp) (Aii), (B) in the st16 posterior NFs (pnf) (Bi, Biii, arrows) and anterior NFs (anf) (Bii) with expression expanding into the medial neural plate and floor plate (fp) (Biii, arrows), and (C) in the apposed dorsal NFs (Ci, ii, iii, arrows). (i), Dorsal view; (ii), frontal view; (iii), transverse view at the level of the white dashed line in above panels. Dashed red lines indicate midline. A, Anterior; bp, blastopore; de, dermamyotome; D, dorsal; dnt, dorsal neural tube; me, mesoderm; mhb, midbrain–hindbrain junction; nc, notocord; P, posterior; V, ventral; vnt, ventral neural tube.

ing solution (2% fetal calf serum, 2% goat serum, 0.2% Triton X-100 in PBS), then overnight at 4°C with primary antibodies in blocking solution. Mouse anti-ZO-1 antibody (Zymed Laboratories) was used at 1:70. Zebrafish embryos were fixed for 12 h in 4% PFA at 4°C, washed in PBS, incubated in 30% sucrose for 2 h and then embedded in Tissue-Tek OCT medium. Sections (20 μ m) were rehydrated in PBS, permeabilized in PBD (PBS containing 0.1% Tween 20 and 0.5% Triton X-100), and blocked for 1 h with 20% goat serum and 1% BSA in PBD. Sections were incubated with primary antibodies in blocking solution overnight at 4°C. Bound antibody was visualized with goat anti-mouse IgG-AlexaFluor 488 (1:1000; Invitrogen). DiI (1, 1'-diiodo-3, 3', 3'-tetramethylindocarbocyanine perchlorate; Invitrogen) was used to label the plasma membrane of neuroepithelial cells in zebrafish embryos. Frozen sections were incubated in 100 μ M DiI for 2 h and then washed extensively with PBD after application of the secondary antibody. Images were acquired on an Olympus IX81 microscope using AnalySIS software or on a Zeiss Axioplan 2 microscope using Axiovision software. Higher resolution images were obtained using a Leica LSM laser scanning confocal microscope or Zeiss LSM 510 Meta confocal microscope.

Results

Loss of *Xneog* results in failure of neural tube closure

We initially determined the expression pattern of *Xenopus Neogenin* (*Xneog*) mRNA from st11 through to st19 of development in the *Xenopus* embryo by *in situ* hybridization. The *Xenopus* neural plate comprises a superficial layer of flat, neuroectodermal cells and a deep layer of densely packed mesenchymal-like cells which have lengthened in the apicobasal dimension (Keller, 2002; Wallingford, 2005). Intense *Xneog* expression was first detected at the beginning of neurulation (st13), with the highest expression localized to the deep layer of the lateral neural plate at the level of

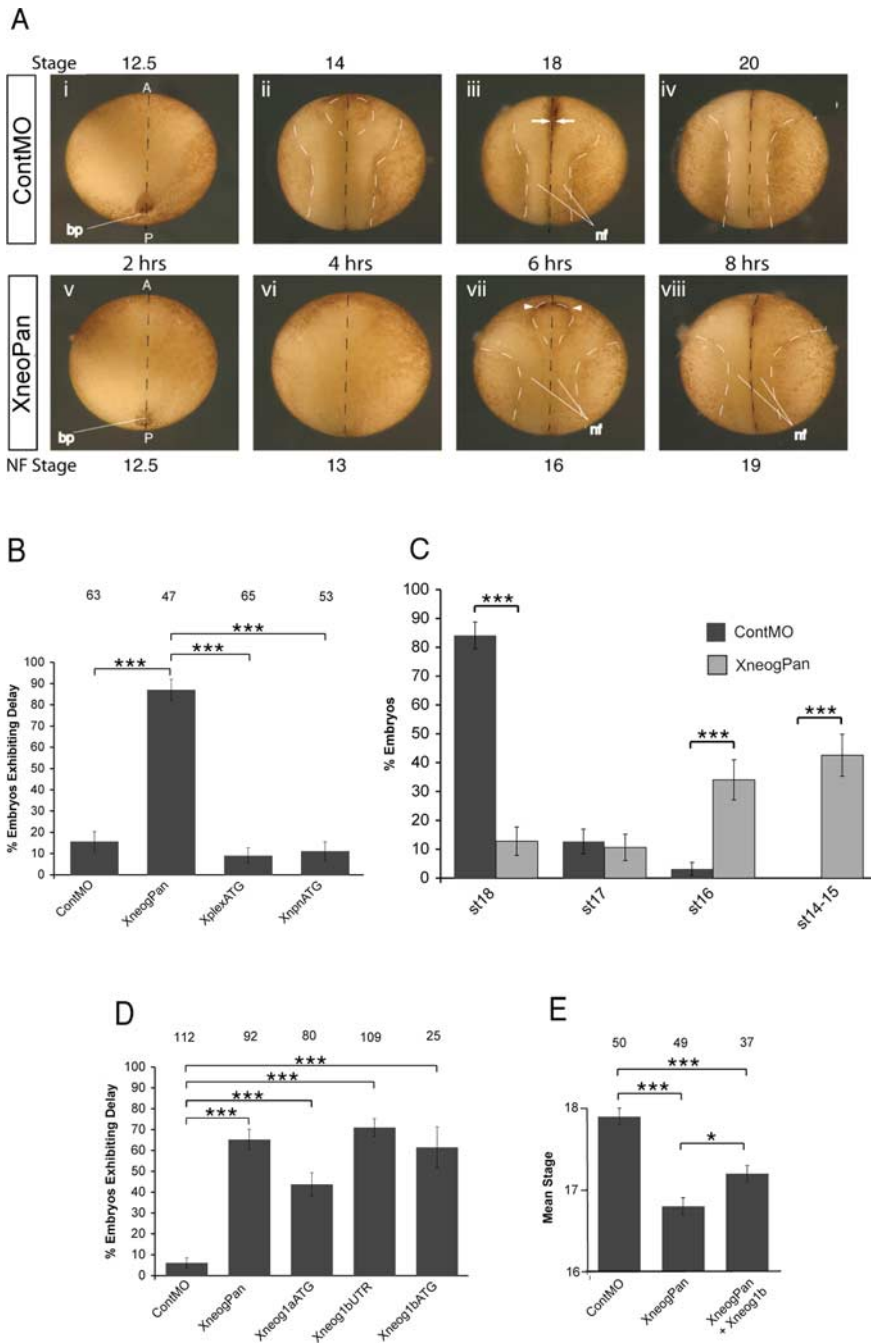


Figure 2. Loss of Xneog results in neural tube closure defects. **A**, Injection of 16 ng of XneogPan MO but not ContMO results in delayed neural tube closure (dorsal view). Hours indicate time since embryos were at st11. Dashed white lines indicate the elevating NFs. Dashed black lines indicate midline. **Aiii**, Arrows: NF apposition. **Avii**, Arrowheads: NFs have failed to reach apposition. **B**, Quantification of the number of embryos exhibiting a neurulation delay when injected with ContMO, XneogPan MO, XplexATG MO or XnpnATG MO. **C**, Percentage of XneogPan-injected embryos at each stage of neurulation when 90% of ContMO-injected embryos have reached st18. **D**, Percentage of embryos exhibiting a neurulation delay when injected with 5' UTR and ATG MOs specific for *Xneog1a* and *Xneog1b*. **E**, Coinjection of XneogPan with full-length Xneog1b cRNA increase the mean neurulation stage compared with injection of XneogPan alone. The number of embryos in each condition is given at the top of each graph. Data are represented as mean \pm SEM. *** $p < 0.001$; * $p < 0.05$.

the presumptive forebrain and hindbrain (Fig. 1A). During NF elevation (st13–16), *Xneog* expression intensified in the deep layer cells of the rising NFs (Fig. 1B). Again *Xneog* was most intense in the anterior neural tube. In addition, the *Xneog* expression domain had expanded into the midline, with expression appearing in the medial neural plate and presumptive floor plate (Fig. 1Biii, arrows). In the late neurula, when NF apposition is

occurring (st19), *Xneog* expression was most intense in the dorsal NFs along the entire neuraxis (Fig. 1C, arrows) while lower expression was observed in the ventral regions. In summary, *Xneog* is expressed in the deep layer of the neural plate throughout neurulation. It is most prominent in the lateral neural plate which gives rise to the NFs and remains high in the folds as they elevate and migrate toward the midline.

To determine whether Xneog plays a role in *Xenopus* neurulation we knocked down *Xneog* expression using antisense MOs. Initially, both cells in two-cell stage *Xenopus* embryos were injected with 8 ng (16 ng total) of XneogPan MO or ContMO (supplemental Fig. 1, available at www.jneurosci.org as supplemental material) and development was monitored throughout neurulation. Both ContMO- and XneogPan MO-injected embryos developed normally until late gastrulation (Fig. 2A) (st12.5). No defects were observed in blastopore formation or involution in embryos injected with either the ContMO or XneogPan MO. However, from st12.5 onwards, neurulation proceeded at a markedly slower rate in embryos injected with XneogPan MO (morphants) (Fig. 2A). In ContMO embryos, NF apposition (st18) occurred 6 h after reaching mid-gastrulation (st11). By the end of this 6 h period ~90% of ContMO embryos had reached st18. In contrast, within the same time period, only 10–15% of morphants exhibited a closed neural tube (Fig. 2A–C). We observed that the severity of the delay phenotype varied within each experiment. On average, 6 h after reaching st11, 11% of morphants were at st17, 34% were at st16, and 43% were still at st14–15 (Fig. 2C). While the progression through neurulation in the most severely affected morphants was delayed by >2 h, NF fusion did eventually occur and the embryos survived past st35 (see Fig. 7).

We also observed that anterior neural tube closure appeared to be affected to a greater extent than posterior closure in the morphants (Fig. 2Avii). To further test this hypothesis the amount of injected MO was increased to 24 ng (12 ng per cell of two-cell stage embryos) and anterior neural tube closure was scored when >80% of ContMO-injected embryos had reached st21–22 (2 h after NF fusion). In the XneogPan MO-injected embryos, an open anterior neural tube was observed in 100% of morphants compared with 20% in control embryos ($p < 0.001$, Student's *t* test). However, in all cases, the posterior neural tube had fused. Therefore, the anterior neural tube is more severely affected by knock-down of Xneog when compared with the posterior neural tube.

Table 1. Mean neurulation stage in XneogPan MO- versus control MO-injected embryos

Morpholino ^a	Mean stage ^b ± SEM	No. of embryos
ContMO	17.81 ± 0.06 <i>p</i> < 0.001 ^c	63
XneogPan	15.89 ± 0.20	47
XplexATG	17.91 ± 0.04 <i>p</i> < 0.001	65
XnnpnATG	17.89 ± 0.04 <i>p</i> < 0.001	53

^aA total of 16 ng of each MO was injected per embryo.

^bMean neurulation stage for embryo cohorts injected with specified MOs. Embryos are staged according to Nieuwkoop and Faber (1994), when 90% of ContMO-injected embryos exhibit neural fold apposition (st18).

^cNeurulation stage was analyzed for each condition relative to XneogPan using the Mann–Whitney test.

Table 2. Mean neurulation stage in Xneog1a MO- and 1b MO-injected embryos

Morpholino ^a	Mean stage ^b ± SEM	No. of embryos
ContMO	17.94 ± 0.02	112
XneogPan	17.09 ± 0.08 <i>p</i> < 0.001 ^c	92
Xneog1aATG	17.46 ± 0.09 <i>p</i> < 0.001	80
Xneog1bUTR	16.99 ± 0.07 <i>p</i> < 0.001	109
Xneog1bATG	17.04 ± 0.20 <i>p</i> < 0.001	25

^aA total of 16 ng of each MO was injected per embryo.

^bMean neurulation stage for embryo cohorts injected with specified MOs. Embryos are staged according to Nieuwkoop and Faber (1994), when 90% of ContMO-injected embryos exhibit neural fold apposition (st18).

^cNeurulation stage was analyzed for each condition relative to ContMO using ANOVA and Dunnett's *T3 post hoc* test.

To demonstrate that defective neural tube closure did not result from a generalized, nonspecific disruption of neurulation resulting from our experimental procedure we injected MOs specific for the semaphorin guidance receptors *XplexinA* and *Xneuropilin1* (Fig. 2*B*). While 87% of XneogPan-injected embryos exhibited a delay in neural tube closure, only 10% of embryos injected with the XplexinA (Xplex) MO or Xneuropilin1 (Xnnpn) MO were delayed. We also calculated the mean neurulation stage for each embryo cohort when 90% of control embryos had reached st18 (Table 1) and demonstrated that the cohort injected with XneogPan MO (mean stage: st15.9 ± 0.2, *p* < 0.001) was significantly delayed when compared with the cohort injected with ContMO (st17.8 ± 0.1), Xplex MO (st17.9 ± 0.0), or Xnnpn MO (st17.9 ± 0.0).

There are two distinct *Xneog* genes (*Xneog1a* and *Xneog1b*) in the *Xenopus* genome (Wilson and Key, unpublished observations) (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Significant delays in NF apposition were also observed when the Xneog1aATG, Xneog1bATG or Xneog1bUTR MOs were used (Fig. 2*D*). In all instances, when control embryos displayed NF apposition (st18), the NFs of Xneog morphants had not yet reached the midline, and were between one to three neurulation stages behind (i.e., st15–17). While the efficiency of each Xneog MO varied, we observed a significant delay (*p* < 0.001) in mean neurulation stage for the cohorts injected with these MOs compared with ContMO-injected cohort (Table 2). The observed differences in the mean neurulation stage resulting from injection of each Xneog MO reflects the severity of the delay. These data indicate that Xneog1aATG is less effective in silencing Xneog function than the other MOs tested. In summary, both Xneog1a and Xneog1b are required for efficient neurulation in the *Xenopus* embryo.

The sequence of the 5' UTR target region of the XneogPan MO is derived from *Xneog1a* and differs from the corresponding sequence in *Xneog1b* by only one nucleotide (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Therefore, the XneogPan MO is likely to prevent translation from both the *Xneog1a* and *1b* transcripts. To further demonstrate specificity we performed rescue experiments in which the XneogPan MO

was coinjected with a cRNA encoding full-length Xneog1b. We were not able to perform a similar rescue experiment with *Xneog1a* as we do not have a full-length *Xneog1a* cDNA available to us. Nonetheless, coinjection of wild-type *Xneog1b* decreased to a small but significant extent the severity of the neurulation delay when compared with that resulting from the injection of the XneogPan MO alone. Figure 2*E* shows that coinjection of *Xneog1b* cRNA with XneogPan MO increased the mean neurulation stage from st16.8 ± 0.1 (XneogPan MO alone) to st17.2 ± 0.1 (*p* < 0.05). That both Xneog1a and 1b MOs cause delays in neural tube closure indicates that both Xneog1a and 1b are required for efficient neurulation. This observation provides an explanation for the small degree of rescue observed with the *Xneog1b* cRNA alone.

Xrgma is a ligand for Xneog during neurulation

In situ hybridization analysis demonstrated strong expression of *Xrgma* mRNA adjacent to the midline along the entire anteroposterior axis during neurulation (st13–19) where it was localized to the deep cells of the neural plate (Fig. 3). At st13 it was found only in the medial half of the neural plate and was therefore expressed directly adjacent to the domain of *Xneog* expression in the lateral neural plate (Fig. 3*Aiv*). During NF elevation (st13–16), the *Xrgma* expression domain contracted toward the midline (Fig. 3*B*). At this stage, *Xneog* expression had expanded toward the midline and therefore overlapped that of *Xrgma* (Fig. 3*Biii, iv*). At NF apposition (st19), *Xrgma* expression became prominent in the ventral neural tube (Fig. 3*C*). However, little *Xrgma* was observed within the strong *Xneog* expression domain in the dorsal neural tube (Fig. 3*Ciii, iv*). Therefore, *Xrgma* is expressed adjacent to the midline and partially overlaps the *Xneog* domain during neurulation.

An antisense MO knock-down strategy was then used to determine whether Xrgma was required for neural tube closure. Figure 4*A* demonstrates that injection of the XrgmaATG or XrgmaUTR MOs (16 ng total) (supplemental Fig. 1*B*, available at www.jneurosci.org as supplemental material) at the two-cell stage resulted in a significant delay in neural tube closure (*p* < 0.001, *p* < 0.05, respectively). The severity of delay induced by the XrgmaATG MO was equivalent to that seen with the XneogPan MO (Fig. 2*A*), while injection of the XrgmaUTR MO produced a less severe phenotype. However, we observed a significant delay in mean neurulation stage for the embryo cohort injected with either Xrgma MO compared with the ContMO-injected cohort (Table 3). To further demonstrate that Xrgma functions as a ligand for Xneog during neurulation, we coinjected a hypomorphic dose of XneogPan MO with XrgmaATG MO or ContMO (total of 8 ng of each MO) (Fig. 4*B*). While coinjection of XneogPan MO (+ ContMO) resulted in delayed NF fusion in 30% of embryos, no significant delay was observed upon coinjection of XrgmaATG MO (+ ContMO). However, coinjection of XneogPan MO and XrgmaATG MO resulted in a significant increase in the number of delayed embryos (71%) above that seen when either MO was injected with the control MO (*p* < 0.001). No additional phenotypic abnormalities were observed upon coinjection of the receptor and ligand MOs. The increase in phenotypic severity could not be attributed to nonspecific interactions as coinjection of XneogPan MO and the Xnnpn MO did not enhance the neurulation delay (data not shown).

To further confirm that knock-down of Xneog and Xrgma perturbs neurulation in an equivalent manner we plotted the time taken (post st12.5) for embryos to reach key neurulation stages (Fig. 4*C*). All embryos injected with either the XrgmaATG

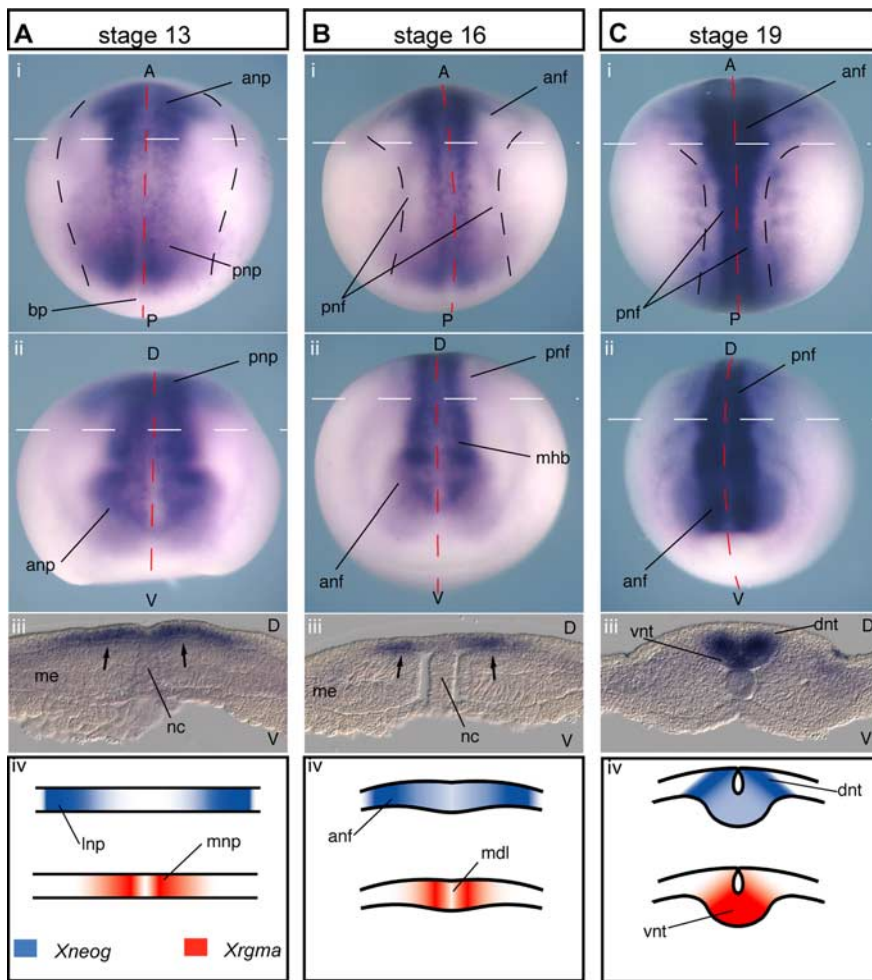


Figure 3. *Xrgma* expression in the deep layer of the medial neural plate. *Xrgma* is expressed (A) in the deep layer of the st13 medial posterior neural plate (Ai, Aiii, arrows) and the medial anterior neural plate (Aii), (B) between the midline and the posterior NFs (Bi, Biii, arrows), as well as between the midline and the anterior NFs (Bii) of the st16 neurula, and (C) in the st19 ventral neural tube (Ci, Cii, Ciii). (i), Dorsal view; (ii), frontal view; (iii), transverse view at the level of the dashed white line in above panels; (iv), schematic of complementary *Xneog* and *Xrgma* expression domains. lnp, Lateral neural plate; mdl, midline; mnp, medial neural plate.

or *Xneog*Pan MO displayed a wild-type phenotype before st12.5, and all *Xneog* and *Xrgma* morphants reached st12.5 at the same time as control embryos. However from st12.5 onwards, the progression through neurulation was markedly slower for morphants compared with control embryos. Moreover, the rate of neurulation in both *Xneog* and *Xrgma* morphants was equivalent. Therefore, we conclude that *Xrgma* interacts with *Xneog* during neurulation to promote neural tube closure.

Xneog and *Xrgma* are required for NF elevation

To identify the morphological defects underlying failure of neural tube closure, we injected 12 ng of *Xneog*Pan MO or ContMO unilaterally into one cell of two-cell stage embryos and immunolabeled at st17 with an antibody specific for β -tubulin. The temporal progression of NF elevation, neural plate bending and NF migration in embryos unilaterally injected with the ContMO was equivalent on the ContMO (Fig. 5A, right, arrow) and wild-type side (Fig. 5A, left). In addition, no disruption of neural plate morphology was observed on the ContMO-injected side. In contrast, the unilateral injection of *Xneog*Pan MO resulted in a obvious perturbation of NF elevation at both the anterior and posterior levels (Fig. 5B, I, right, arrows) when compared with the

uninjected, wild-type side (Fig. 5B, I, left). Figure 5B clearly demonstrates that while the anterior NF on the wild-type side had undergone appropriate elevation, the height of the fold on the morphant side was markedly reduced, leading to a broadening of the fold in the lateral direction (Fig. 5B, arrow). Similarly, failure of NF elevation was observed on the morphant side of the posterior neural plate (Fig. 5I, arrow). Therefore, loss of *Xneog* results in inefficient NF elevation along the entire neuraxis. Unilateral injections of the *Xrgma*ATG MO produced a similar, albeit less severe, NF elevation phenotype (data not shown), confirming that *Xrgma* is the Neogenin ligand promoting NF elevation. It is important to note that, despite the suppression of NF elevation in the absence of *Xneog* or *Xrgma*, the hinge-points formed correctly (Fig. 5B, arrowheads) and the distance between the midline and hinge-point was equivalent to that on the wild-type side, suggesting that both apical constriction and convergent extension were still operative.

Loss of *Xneog* perturbs deep layer cell polarity by disrupting the microtubule cytoskeleton

From the start of neurulation (st13) until after neural tube closure (st22) cells of the deep layer are elongated in an apicobasal direction (initially the apicobasal plane equates to the dorsoventral plane of the neural plate) and are tightly apposed in wild-type embryos. Close examination of the unilaterally injected st17 embryos immunolabeled with the anti- β -tubulin antibody revealed that while the deep cells in the wild-type anterior NF were elongated

in the apicobasal direction (Fig. 5C), those on the *Xneog*Pan-injected side had lost their apicobasal polarity and displayed a rounded morphology (Fig. 5D, arrowheads). Higher magnification revealed that the microtubule (MT) cytoskeleton in these cells was disrupted. In the presence of the *Xneog*Pan MO, β -tubulin immunoreactivity was found in punctate aggregates or dispersed throughout the cytoplasm of the deep cells, indicating depolymerization and fragmentation of the MT filaments (Fig. 5D, F). In contrast, MT filaments in wild-type deep cells formed a tight mesh of fibers adjacent to the plasma membrane (Fig. 5C, E). Disruption of the MT cytoskeleton was also observed when the *Xrgma*ATG MO was unilaterally injected (data not shown). This phenotype is highly consistent, as it was also observed in embryos when lower doses of *Xneog*Pan or *Xrgma*ATG MOs were bilaterally injected (8 ng into each cell of two-cell stage embryos). Together, these observations suggest that *Xneog*-*Xrgma* interactions establish apicobasal polarity in the deep layer cells by regulating the MT network.

Within polarized neuroepithelium, tight junctions are found adjacent to the apical surface of the cell. Therefore, localization of junctional proteins (e.g., ZO-1) to the tight junction indicates that correct apicobasal polarity has been established (Nelson,

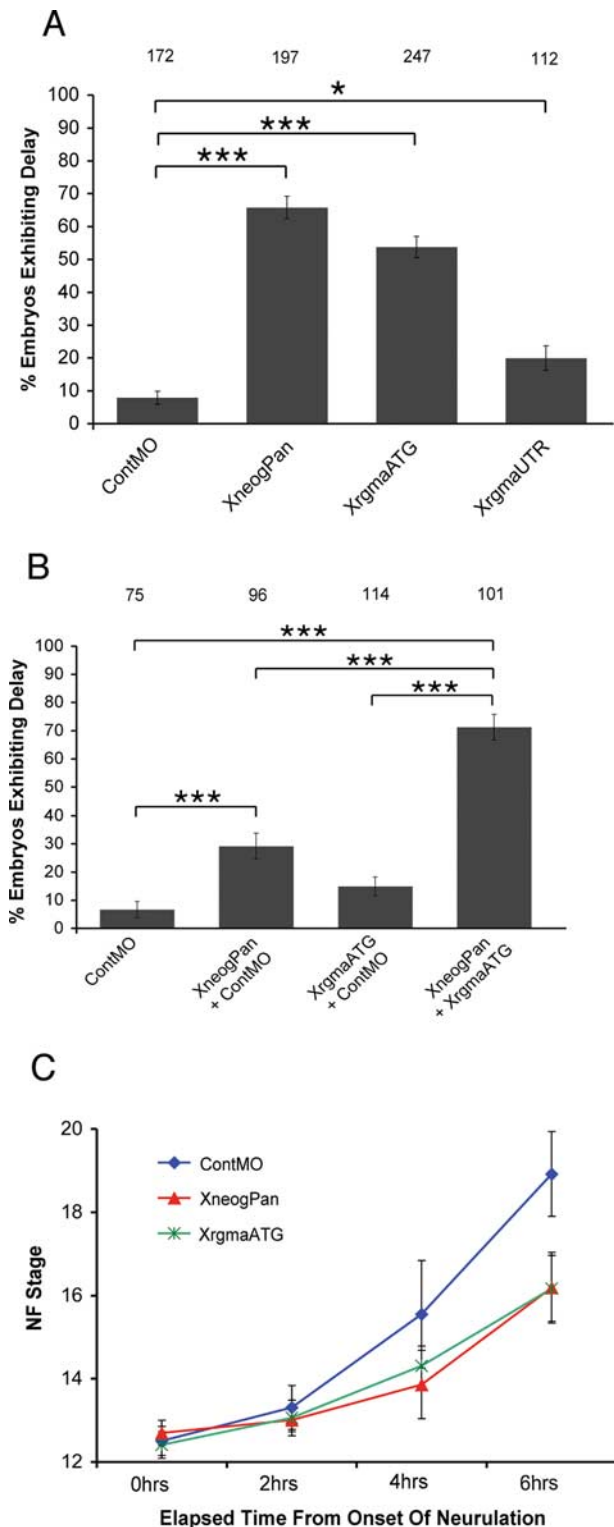


Figure 4. Loss of Xrgma results in neural tube closure defects. **A**, Injection of 16 ng of XrgmaATG MO or XrgmaUTR MO produces a neurulation delay phenotype. **B**, Coinjection of 8 ng of XneogPan and XrgmaATG MOs enhances the delay in neurulation. Data are represented as mean \pm SEM. **C**, XneogPan MO and XrgmaATG MO generate equivalent decreases in the rate of progression through neurulation. Data are represented as mean \pm SD.

2003). We wished to confirm the requirement for Neogenin in the establishment of apicobasal polarity during neurulation by investigating the distribution of ZO-1 in our *Xenopus* morphants. However, we were not able to identify an antibody that recog-

Table 3. Mean neurulation stage in Xrgma MO-injected embryos

Morpholino ^a	Mean stage ^b \pm SEM	No. of embryos
ContMO	17.92 \pm 0.02	172
XneogPan	16.81 \pm 0.08 $p < 0.001^c$	197
XrgmaATG	17.11 \pm 0.06 $p < 0.001$	247
XrgmaUTR	17.76 \pm 0.05 $p = 0.002$	112

^aA total of 16 ng of each MO was injected per embryo.

^bMean neurulation stage for embryo cohorts injected with specified MOs. Embryos are staged according to Nieuwkoop and Faber (1994), when 90% of ContMO-injected embryos exhibit neural fold apposition (st18).

^cNeurulation stage was analyzed for each condition relative to ContMO using the Mann–Whitney test.

nized the ZO-1 orthologue in *Xenopus laevis borealis* embryos. Instead, we examined ZO-1 distribution in zebrafish embryos, since the disrupted morphology of the zebrafish neural tube in the absence of Neogenin (Mawdsley et al., 2004) exactly phenocopies that observed in the *Xenopus* morphants. Like most vertebrates, the zebrafish neural plate comprises a single layer of columnar epithelium (Papan and Campos-Ortega, 1994; Lowery and Sive, 2004). During neurulation, the zebrafish neural plate rolls up and condenses into a solid mass of cells (the neural keel) which then sinks into the underlying mesoderm to form a solid rod of neuroepithelium (the neural rod) (Geldmacher-Voss et al., 2003). Lumination then progresses in a ventral to dorsal direction (Schmitz et al., 1993) in a similar manner to *Xenopus* trunk relumination (Davidson and Keller, 1999). Despite the morphological differences between *Xenopus* and zebrafish neurulation, neural keel and neural rod formation are also reliant on the establishment of apicobasal polarity and radial intercalation (for review, see Lowery and Sive, 2004).

To examine the effect of loss of Neogenin on ZO-1 distribution in the neuroepithelium of zebrafish morphants we injected a control MO (zfcontMO) or a MO directed to the ATG sequence of zebrafish *Neogenin* (zfneogATG) into the yolk sac of two-cell stage embryos. We then visualized the morphology of the neuroepithelium within the neural rod by staining the plasma membranes with the lipophilic dye, DiI (Fig. 6). In zfcontMO-injected embryos ZO-1 protein was localized to the tight junctions lying immediately below the apical surface of the neuroepithelial cells at the midline (Fig. 6A). In striking contrast, the morphology of the neural rod was severely disrupted in zfneogATG MO-injected embryos. No obvious midline could be visualized in the morphants and neuroepithelial cells were rounded and no longer aligned in an apicobasal orientation (Fig. 6B, C). Moreover, ZO-1 immunoreactivity was no longer restricted to one region of the plasma membrane, but was found distributed around the entire cell surface (Fig. 6B, C). Therefore, in the absence of Neogenin the distribution of ZO-1 was not apically polarized, demonstrating that the neuroepithelial cells had lost their apicobasal polarity.

Loss of Xneog leads to failure of radial intercalation and inefficient superficial cell wedging

In the *Xenopus* morphants (Fig. 5) we further observed that the inability of deep layer cells to maintain an apicobasally orientated elongated shape resulted in the failure of radial intercalation with the overlying superficial cell layer in the anterior neural tube. Figure 5 shows that, around the hinge-point of the wild-type side, the apical membranes of the deep cells (Fig. 5G, arrowheads) had intercalated with the superficial layer forming close contacts with adjacent wedge-shaped superficial cells (Fig. 5G, asterisks). In striking contrast, the rounded deep cells (Fig. 5H, arrowheads) on the morphant side had not interdigitated with the superficial cells (Fig. 5H, asterisks), leaving both layers only loosely associ-

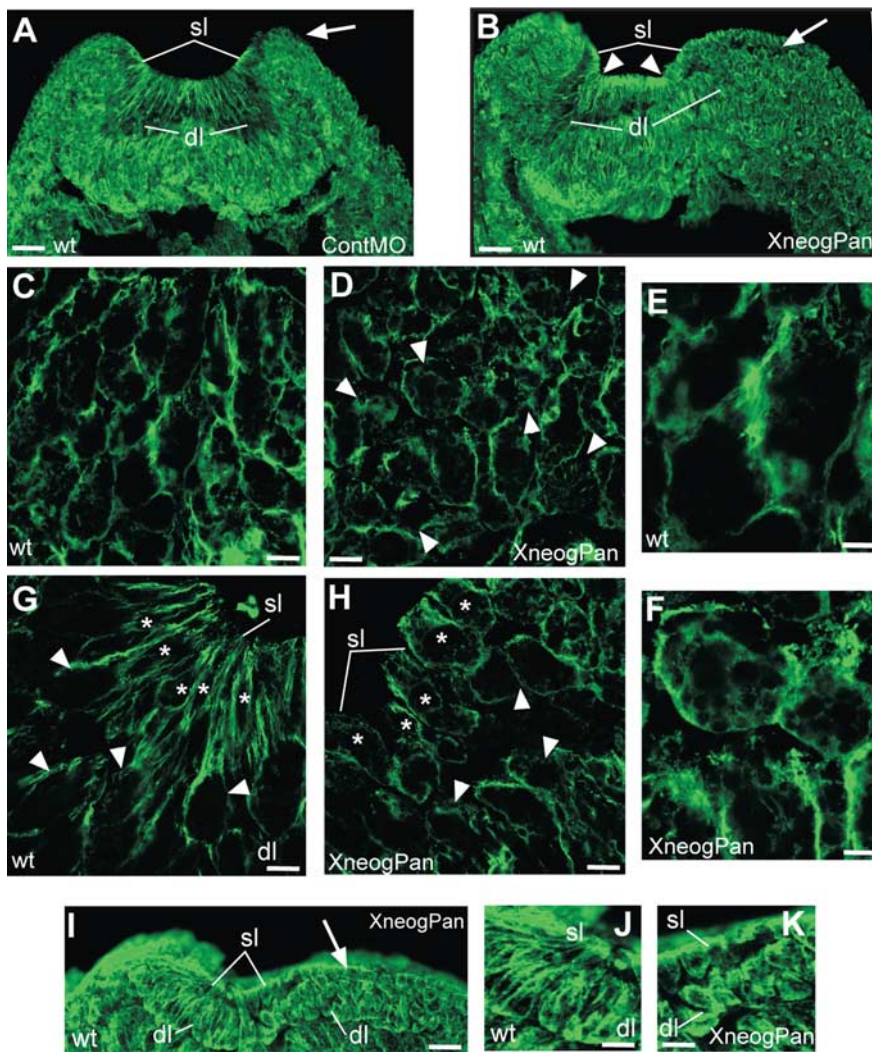


Figure 5. Loss of Xneog perturbs NF elevation and cell polarity. Transverse sections of st17 embryos immunostained with the anti- β -tubulin antibody, E7. **A**, Injection of ContMO into one cell of two-cell stage embryos did not affect anterior NF elevation on the ContMO (right side, arrow) or the wild-type (wt) side (left side) of the embryo. **B**, Injection of XneogPan MO disrupts anterior NF elevation on the morphant side of the embryo (right side, arrow). Hinge-points are indicated by arrowheads. **C–F**, Higher magnification images of the deep layer (dl) in **B** on the wild-type side (**C**, **E**) and XneogPan MO side (**D**, **F**) where deep cells have lost their apicobasal polarity (arrowheads). **G–H**, Higher magnification images of the deep (arrowheads) and superficial layers (sl, asterisks) in **B** at the hinge-point on the wild-type side (**G**) and the XneogPan MO side (**H**). **I–K**, Posterior NF elevation is inhibited on the XneogPan MO side (**J**, arrow; **K**) but not the wild-type side (**I**, left side; **J**). Scale bars: **A**, **B**, **I**, 39 μ m; **C**, **D**, **G**, **H**, 2 μ m; **E**, **F**, 1 μ m; **J**, **K**, 16 μ m.

ated. Moreover, while some degree of apical constriction had occurred in these superficial cells, they were significantly less wedge-shaped and less elongated than those on the wild-type side. Also, while β -tubulin had accumulated under the apical surface in the morphant superficial cells, long apicobasally directed MT filaments originating from the apical region (Karfunkel, 1971; Burnside, 1973; Lee et al., 2007) were only seen in wild-type cells (Fig. 5, compare *G,H*). Knock-down of Xneog in the posterior neurula also resulted in loss of the apicobasally orientated elongated shape of the deep layer cells, failure of radial intercalation, and loss of apical constriction in superficial cells (Fig. 5, compare *J,K*). Therefore, our data indicate that efficient apical constriction and elongation of superficial cells is reliant on appropriate interaction with the intercalating deep cells. Since Xneog and Xrgma are not expressed in the superficial layer, the disruption to this layer is a non-cell-autonomous effect, secondary to the disruption of deep layer cell polarity and intercalation.

These observations indicate that apicobasal polarization of the deep layer cells is essential for radial intercalation. Moreover, our data suggest that superficial cell wedging is potentiated by intimate contact with appropriately polarized deep cells.

Xneog and Xrgma are required for apicobasal polarity of the pseudostratified epithelium in the neural tube

While knock-down of Xneog or Xrgma using a total of 16 ng of MO per embryo resulted in a significant delay in NF elevation, the NFs did eventually meet and fuse at the midline to form a neural tube. To determine whether loss of Xneog and Xrgma continued to perturb cell polarity within the pseudostratified epithelium of the neural tube after closure (st35) we used the Xen1 antibody which recognizes an unknown epitope on the cell surface of *Xenopus* neuroepithelial cells (Ruiz i Altaba, 1994). While the pseudostratified epithelium of the ContMO neural tube developed normally (Fig. 7*A*), the Xneog and Xrgma morphant neural tubes displayed a severely perturbed neuroepithelium with a smaller, misshapen lumen (Fig. 7*B,C*, respectively). Within the morphant neural tube, the neuroepithelial cells had detached from the apical and basal surfaces, become rounded, and were distributed in a disorganized manner throughout the structure (Fig. 7*Bii,iii, Cii,iii*). Counterstaining with DAPI revealed many pyknotic nuclei within the neuroepithelia and luminal spaces of the Xneog and Xrgma morphants, indicating that large numbers of cells were undergoing cell death (Fig. 7*Bi, Ci*). No pyknotic cells were observed in the control neural tube. These neural tube defects were never observed in embryos injected with the XplexinA or Xneuropilin1 MOs (data not shown).

To confirm specificity, we performed a rescue experiment in which the XneogPan MO was coinjected with a cRNA encoding full-length Xneog1b. When carrying out similar experiments to rescue the neurulation delay phenotype (Fig. 2*E*), we found that coinjection of Xneog1b cRNA only rescued this phenotype to a small extent. To examine the ability of Xneog1b to rescue the aberrant morphology of the st35 neural tube we examined the neural tube of embryos that exhibited a neurulation delay in the presence of the coinjected XneogPan MO and the Xneog1b cRNA (Fig. 8*C*). That is, embryos coinjected with XneogPan MO + Xneog1b cRNA that were found to be a st13–st16 when 90% of ContMO injected embryos had reached st18 were examined. Figure 8*C* shows that Xneog1b cRNA was able to almost fully rescue the morphology of the st35 neuroepithelium after the neural tube had eventually closed. This suggests that the maintenance of neuroepithelial morphology in the st35 neural tube is predominantly dependent on Xneog1b.

In summary, we have demonstrated that Xrgma-Xneog inter-

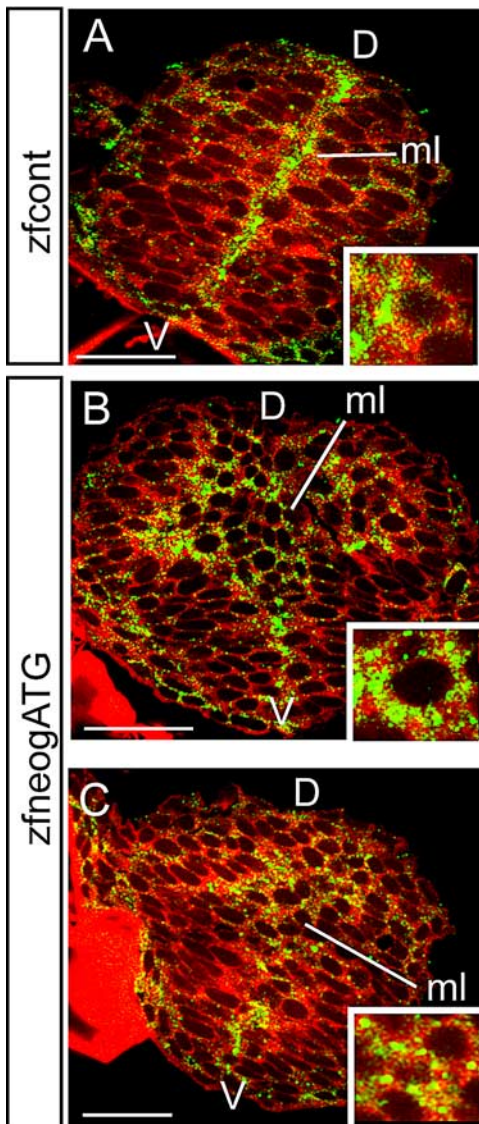


Figure 6. Loss of Neogenin disrupts localization of ZO-1 to the apical domain of neuroepithelial cells in the zebrafish neural rod. **A**, ZO-1 protein (green) is restricted to a domain adjacent to the apical surface of neuroepithelial cells at the midline (ml) of the neural rod of embryos injected with zfcont MO. **B, C**, Injection of the zfneogATG MO resulted in loss of an obvious midline. Neuroepithelial cells were rounded and no longer aligned in an apicobasal orientation. In addition, ZO-1 immunoreactivity was distributed around the entire cell surface. Insets are high-magnification images of cells exhibiting a polarized distribution of ZO-1 in control embryos (**A**) or a nonpolarized distribution of ZO-1 in morphant cells (**B, C**). The lipophilic dye, Dil (red) was used to visualize the cell shape of the neuroepithelium. Scale bars: **A–C**, 20 μ m.

actions maintain deep cell morphology within the neural plate during NF elevation and are therefore required for successful neural tube closure along the entire neuraxis, with the anterior neural tube being most sensitive to loss of this receptor–ligand pair. We also show that RGMa and Neogenin are essential for the establishment of apicobasal polarity of the pseudostratified neuroepithelium within the neural tube. Moreover, our data suggests that correct neurulation requires both Xneog1a and Xneog1b, while Xneog1b is the predominant form in the neuroepithelium after neural tube closure.

Discussion

Over the past decade, the molecular pathways regulating convergent extension (the noncanonical Wnt/PCP pathway) and neural

plate bending (Shroom3, Sonic hedgehog, BMP-2 pathways) have been identified (Ybot-Gonzalez et al., 2002, 2007; Wallingford, 2005; Kibar et al., 2007). However, as yet, no signaling pathway has been linked to NF elevation. Here we show that Xneog–Xrgma interactions are essential for effective NF elevation. We provide evidence that Xneog is required to maintain the integrity of the MT network within the deep layer cells of the neural plate. We also demonstrated that Xneog-dependent regulation of MTs within the deep cells is essential for their radial intercalation with the overlying superficial cells which is a prerequisite for NF elevation. A requirement for Xneog and Xrgma in the establishment of apicobasal polarity and therefore epithelialization of the pseudostratified neuroepithelium within the neural tube was also demonstrated.

Neogenin controls the MT cytoskeleton

At the onset of *Xenopus* neurulation (st13) the deep layer cells elongate such that they are orientated in the dorsoventral plane of the neural plate (this will become the apicobasal plane during neural groove formation) (Schroeder, 1970; Karfunkel, 1971; Keller et al., 1992; Davidson and Keller, 1999). At st13 the MTs are randomly distributed throughout the deep cell cytoplasm (Karfunkel, 1971). However, as NF elevation proceeds, the MT filaments align along the apicobasal axis of the deep cells as they undergo radial intercalation. Therefore, active reorientation of the MT network in the direction of migration accompanies the initiation of deep cell intercalation. Examination of the MT network of deep cells in the Xneog or Xrgma morphants using an antibody to β -tubulin revealed that, in the absence of Neogenin activity, tubulin had dispersed throughout the cytoplasm, indicating depolymerization of the MTs. These observations suggest that Xneog–Xrgma interactions establish the apicobasally orientated shape of the deep layer cells by regulating the MT network. The hypothesis that Neogenin is important in the establishment of apicobasal polarity is supported by the perturbation in apical localization of the polarity marker, ZO-1, in the neuroepithelium of zebrafish injected with the zfneogATG MO (Fig. 6B, C).

That the integrity of an apicobasally orientated MT network is essential for radial intercalation and NF elevation is supported by an early study on the role of MTs in *Xenopus* neurulation (Karfunkel, 1971). Strikingly, the phenotype observed after exposure of embryos throughout neurulation to the tubulin polymerization inhibitor, vinblastine, exactly phenocopies the NF and neural tube disruptions observed in our Xneog and Xrgma morphants. Therefore, disruption of the MT network by drug treatment or by silencing of Neogenin activity results in equivalent perturbations in deep cell morphology, loss of NF elevation and neural tube morphology, supporting the hypothesis that Neogenin orientates the MT network of intercalating deep layers, thereby establishing correct apicobasal polarity. At present, we do not know how Neogenin regulates the MT network. However, it is known that the neural-specific MT associated protein, MAP1B, lies downstream of the Netrin receptor DCC, and is phosphorylated by Netrin-dependent activation of GSK3 and CDK5 (Del Río et al., 2004). Thus, Neogenin may also control MT activity via a similar pathway.

Neogenin-dependent radial intercalation drives NF elevation

Our study has revealed the dependence of NF elevation on Neogenin-driven radial intercalation. We observed that the aberrant morphology of the deep layer cells prohibited intercalation with the overlying superficial layer. In the *Xenopus* neurula, it is the lateral and intermediate regions of the neural plate that un-

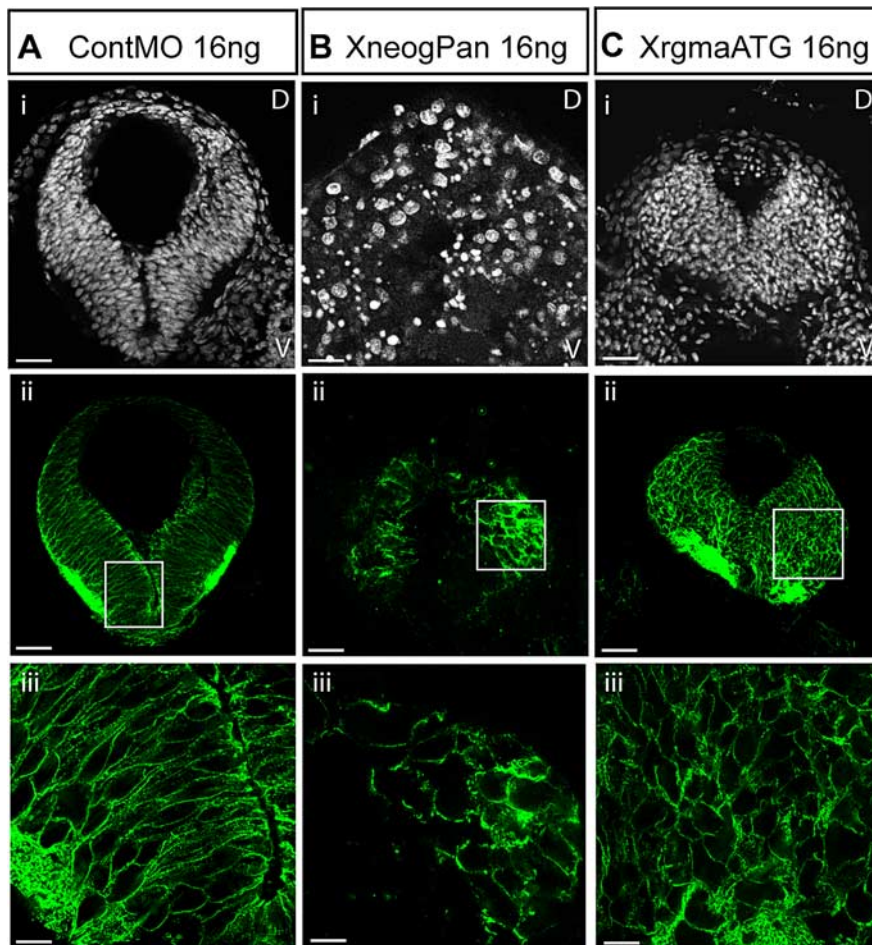


Figure 7. Xneog and Xrgma are required for the establishment of the neuroepithelium of the neural tube. Transverse sections of the st35 anterior neural tube labeled with DAPI (**Ai, Bi, Ci**) and the Xen1 antibody (**Aii, Bii, Cii, Aiii, Biii, Ciii**). Injection of XneogPan MO (**B**) or XrgmaATG MO (**C**) but not ContMO (**A**) disrupts the neuroepithelium of the neural tube. **Aiii, Biii** and **Ciii** are higher magnification of images of boxed areas in **Aii, Bii** and **Cii**, respectively. Scale bars: **Ai–Cii**, 29 μm ; **Aiii–Ciii**, 8 μm .

dergo elevation to form the NFs (Schroeder, 1970, 1971; Davidson and Keller, 1999). Intercalation of deep and superficial cells proceeds from lateral to medial, with interdigitation in the lateral and intermediate regions of the neural plate increasing threefold as the NFs elevate and migrate toward the midline (Hartenstein, 1989). We have shown that intense *Xneog* mRNA expression is initially localized to the lateral deep layer of the neural plate and then expands into the intermediate deep layer as NF elevation proceeds. Therefore, Neogenin is present in those cells actively participating in radial intercalation and NF elevation. Together these observations indicate that Neogenin activity in the deep cell layer promotes radial intercalation by regulating apicobasal polarity through the MT network.

Unexpectedly, loss of Neogenin also disrupted the integrity of the superficial layer. These cells had undergone some degree of apical constriction as a high concentration of tubulin was localized under the apical membrane (Karfunkel, 1971; Burnside, 1973; Lee et al., 2007) and obvious hinge-points were generated in the mutants, indicating that apical constriction had occurred, albeit less efficiently. Since *Xneog* and *Xrgma* are not expressed in the superficial layer, this disruption is non-cell-autonomous and secondary to the disruption of the deep cell layer. Recently, the cytoskeletal organizing protein, Shroom3, has been shown to be necessary and sufficient to trigger apical constriction in superficial cells. Shroom3 activity is strictly dependent upon the prior

establishment of apicobasal polarity in epithelial cells (Haigo et al., 2003; Lee et al., 2007). However, the mechanism by which apicobasal polarity is initially established in superficial cells has not been elucidated. Our data indicate that constriction of superficial cells is potentiated by contact with the intercalating deep cells. Therefore, we propose that Neogenin activity, via its promotion of deep cell radial intercalation, also regulates superficial cell polarity and, consequently, apical constriction.

Neogenin and RGMA are essential for epithelialization

As the deep cells integrate into the superficial layer, they concomitantly undergo a mesenchymal to epithelial transition (Keller et al., 1992; Davidson and Keller, 1999). A fundamental requirement for epithelialization is the establishment of apicobasal polarity (Nelson, 2003; Pilot and Lecuit, 2005; Lecuit and Lenne, 2007). Therefore, Neogenin-dependent organization of the MT network within the intercalating deep cells appears to be an essential prerequisite for deep cell epithelialization. This hypothesis is supported by the severe perturbation and loss of apicobasal polarity of the pseudostratified neuroepithelium of the neural tube in our *Xenopus* and zebrafish morphants.

In *Xneog* and *Xrgma* morphants NFs fuse, although the neuroepithelium of the neural tube is highly disordered. The epithelial character of the cells is lost as they no longer contact the apical and basal surfaces or each other. At NF apposition, *Xneog* mRNA is present in a ventral-low, dorsal-high gradient throughout the early neural tube and is therefore expressed by deep cells actively undergoing radial interaction and epithelialization. From the early phase of NF elevation through to NF fusion, loss of Neogenin in deep cells of the lateral and intermediate neural plate disrupts MT integrity and subsequently intercalation with the superficial layer. Thus the st35 neural tube phenotype seen in our morphants provides strong evidence that sustained RGMA–Neogenin activity is also essential for deep cell radial interaction and epithelialization during, and subsequent to, NF fusion. Further support for this hypothesis comes from our zebrafish study where knock-down of Neogenin produced severe neural tube defects and loss of neuroepithelial morphology identical to that seen in our *Xenopus* morphants (Fig. 6) (Mawdsley et al., 2004). As for *Xenopus*, zebrafish neurulation is reliant on the establishment of apicobasal polarity and radial intercalation (Lowery and Sive, 2004; Hong and Brewster, 2006).

RGMA is the Neogenin ligand during neurulation

RGMA is linked to the plasma membrane by a glycosylphosphatidylinositol moiety and can be cleaved to produce a soluble isoform (Matsunaga et al., 2004; Niederkofler et al., 2004). Therefore, it can act as either a short- or long-range morphogenic factor. Throughout neurulation the highest level of *Xneog* expres-

sion is found in the lateral deep cell layer underlying the rising NFs, while the most intense *Xrgma* expression domain is restricted to the more medial deep cells (Fig. 3). This complementary expression pattern suggests that at the onset of neurulation, the relevant form of RGMA may be the cleaved isoform which is capable of establishing a morphogenic gradient. As the NFs elevate, the *Xneog* expression domains expand medially resulting in overlap between the *Xneog* and *Xrgma* domains. This raises the possibility that as the Neogenin-positive deep cells intercalate, the membrane-bound RGMA isoform on the adjacent cell may act to potentiate cell–cell contact. To date, RGMA has been shown to act as a chemorepulsive axon guidance cue (Rajagopalan et al., 2004; Wilson and Key, 2006). In contrast, our data suggest that the outcome of RGMA–Neogenin interactions in the context of neurulation is the establishment or maintenance of cell polarity and epithelial integrity. A similar situation occurs during mouse mammary gland development, where Netrin-1–Neogenin interactions are required to stabilize the structure of the developing mammary epithelium (Srinivasan et al., 2003; De Vries and Cooper, 2008). That RGMA is important for neural tube closure is further supported by the occurrence of exencephaly in *RGMA* loss-of-function mouse embryos (Niederkofler et al., 2004).

In summary, we show that RGMA and Neogenin are essential for efficient NF elevation and establishment of the pseudostratified epithelium of the neural tube via their ability to regulate apicobasal polarity and radial intercalation.

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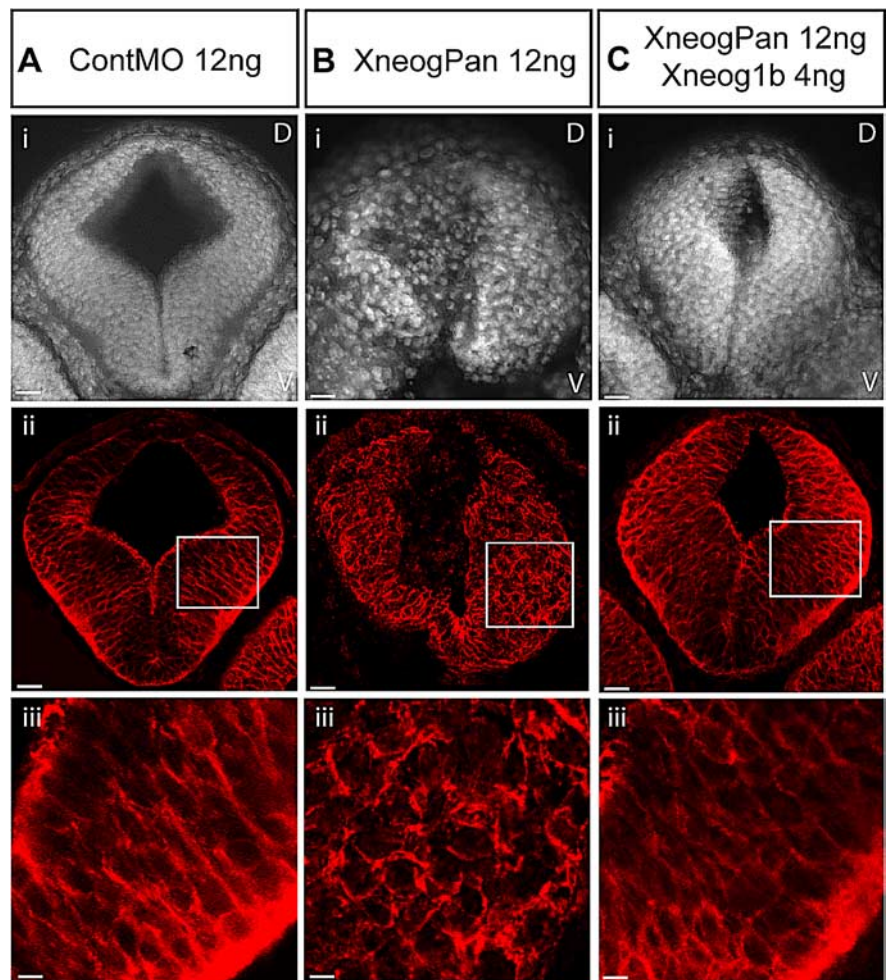


Figure 8. Coinjection of *Xneog1b* cRNA with XneogPan MO rescues the formation of the neuroepithelium in the neural tube. Transverse sections of the st35 anterior neural tube labeled with DAPI (**Ai, Bi, Ci**) and the Xen1 antibody (**Aii, iii, Bii, iii, Cii, iii**). Injection of XneogPan MO (**B**) but not ContMO (**A**) perturbs the neuroepithelium of the neural tube. **C**, Coinjection of *Xneog1b* cRNA with XneogPan MO rescues neural tube morphology. **Aiii, Biii** and **Ciii** are higher magnification of images of boxed areas in **Aii, Bii** and **Cii**, respectively. Scale bars: **Ai–Cii**, 39 μm ; **Aiii–Ciii**, 17 μm .

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