FGF and retinoic acid activity gradients control the timing of neural crest cell emigration in the trunk

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Introduction

The neural crest is formed by a transient population of multipotent cells that arises from the dorsal neural tube. Once specified, neural crest cells (NCCs) undergo a process of epithelial–mesenchymal transition (EMT) that confers the ability to delaminate and migrate away from the dorsal neural tube. The NCCs migrate along characteristic pathways to differentiate into a wide variety of derivatives according to their rostro-caudal (R-C) position in the neural tube and to the order of emigration (Krispin et al., 2010). NCC derivatives include craniofacial skeleton, sensory neurons and glia, sympathetic neurons, and melanocytes, among others (Le Douarin and Kalcheim, 1999).

The process of neural crest formation involves the orchestration of a complex gene regulatory network (Morales et al., 2005; Sauka-Spengler and Bronner-Fraser, 2008). It involves signaling pathways and transcription factors that are responsible for the sequence of early induction of the NCC during gastrulation (Wnt, bone morphogenetic proteins [BMPs], FGF, and retinoic acid [RA]; Liem et al., 1995; Saint-Jeannet et al., 1997; Villanueva et al., 2002; Monsoro-Burq et al., 2003), the specification of the neural plate border (Msx1/2, Pax3, Pax7, and Zic1/3; Liem et al., 1995; Nakata et al., 1997; Brewster et al., 1998), the expression of bona fide NCC transcription factors (AP2, Snail2, FoxD3, Sox9, and Sox10; Nieto et al., 1994; Dottori et al., 2001; Kos et al., 2001; Paratore et al., 2001; Cheung and Briscoe, 2003; Barrallo-Gimeno et al., 2004; Perez-Alcala et al., 2004), and the regulation of numerous downstream effectors involved in cell adhesion and cell cycle control, among others (Sauka-Spengler and Bronner-Fraser, 2008).

Several neural crest specifier genes are coexpressed with neural plate border specifiers at early stages during gastrulation, suggesting possible early regulatory relationships (Khudyakov and Bronner-Fraser, 2009). However, this early population of neural crest progenitors mostly represents the cephalic NCCs, whereas the trunk neural crest progenitors will progressively be specified as the trunk neural tube elongates (Le Douarin and Kalcheim, 1999).
In that sense, the role of BMP, FGF, and Wnt signaling in the early NCC induction has been derived from work in *Xenopus laevis* at cephalic NCC territory. Thus, the signal-promoting trunk neural crest specification has not been fully elucidated.

The development of trunk NCCs is highly coordinated with the development of functionally related adjacent territories. In particular, flanking the neural tube, the paraxial or presomitic mesoderm (PSM) gets progressively segmented into smaller units, somites, and this metamORIZATION imposes a segmented organization to the trunk NCCs. First, opposite the PSM, NCCs are confined to the dorsal neural tube, whereas NCC emigration begins facing epithelial somites (Teillet et al., 1987; Sela-Donenfeld and Kalcheim, 1999). Moreover, the first indication of peripheral nervous system segmentation is the patterned ventral migration of NCCs through the anterior part of each somitic sclerotome (Tosney, 1978; Rickmann et al., 1985; Bronner-Fraser, 1986; Loring and Erickson, 1987).

There is little knowledge about the integration of signals involved in trunk NCC emigration. At trunk level, it has been shown that a BMP-Wnt1 signaling cascade controls NCC emigration (Sela-Donenfeld and Kalcheim, 1999; Burstyn-Cohen et al., 2004). The regulation of that cascade is exerted through the caudal (high)-rostral (low) gradient of the BMP inhibitor noggin, which in turn is controlled by undetermined signals coming from somites (Sela-Donenfeld and Kalcheim, 2000).

In the paraxial mesoderm, the FGF and RA pathways operate as a signaling switch that controls the R-C sequence of mesodermal and neural development. Opposite gradients of declining caudal FGF and rostral RA signaling in the extending body axis create a wavefront, which determines the onset of neuronal differentiation and patterning in newly generated spinal cord (Diez del Corral et al., 2002, 2003) and also positions somite boundaries (Dubrulle et al., 2001; Diez del Corral et al., 2003). The trunk neural tube is generated during a long period of time as the main body axis extends caudally, and trunk neural crest is also generated after this R-C sequence. Thus, FGF and RA signaling pathways are candidates to coordinate NCC development with somite and neuron formation as the embryo elongates.

In this work, using both gain- and loss-of-function approaches, we establish that FGF signaling through the MAPK pathway prevents the initiation of trunk NCC specification and, consequently, NCC EMT. In contrast, RA signaling triggers the EMT of specified NCCs. Furthermore, we show that FGF and RA signaling control the timing of EMT and emigration in part through modulation of elements of the BMP and Wnt signaling pathways. Thus, opposition of FGF and RA signaling orchestrates the onset of trunk NCC specification and emigration, coordinating them with somite formation and neurogenesis in the extending vertebrate body axis.

**Results**

**Onset of NCC marker expression coincides with FGF8 decline**

To investigate whether the paraxial mesoderm has a general influence on NCC specification and migration in the spinal cord, we compared the expression patterns of a cohort of genes involved in dorsal patterning and NCC specification with respect to the formation of somites in chicken embryos at Hamburger and Hamilton (HH) stages 10–14 (Fig. 1, A–G). At the HH10 stage, the caudal open neural plate expressed dorsal patterning genes such as *Pax7* (Fig. 1 A; Kawakami et al., 1997), *Pax3* (not depicted; Goulding et al., 1991), *RhoB* (not depicted; Liu and Jessell, 1998), and *Msx1* (not depicted; Liem et al., 1995). This expression is maintained along the neural tube as the body axis extends (Fig. 1 A). More rostrally, when the neural folds move toward the midline to fuse, the expression of the NCC marker *Snail2* (Nieto et al., 1994) was found in the dorsal neural folds (Fig. 1 B) adjacent to the mid-PSM level. Subsequently, expression of *Sox* family genes involved in NCC formation could be observed in the dorsal neural tube. First, *Sox9* (Fig. 1 C; Cheung and Briscoe, 2003) appeared rostral to the initiation of *Snail2* and then *Sox5* (Fig. 1 D; Perez-Alcala et al., 2004). Finally, adjacent to forming somites, *Sox10* (Fig. 1 E; Paratore et al., 2001) was observed in premigratory NCCs about to emigrate.

This order in the expression of the genes described is maintained throughout stages HH12–14 (unpublished data). However, at HH10, *FoxD3*, an NCC specifier (Stewart et al., 2006), was not expressed dorsally in the neural folds (Fig. 1 F). Only by HH12 could *FoxD3* mRNA be observed at the rostral PSM level (Fig. 1 G).

The initiation of the expression of NCC markers seems to coincide with the transition zone where neural progenitors stop expressing caudal neural markers and initiate ventral patterning gene expression. This transition is governed by the opposing activities of the FGF and RA signaling gradients (Diez del Corral et al., 2003). In fact, by situ hybridization of half embryos, we showed that the onset of *Snail2* expression coincided with the decline of the caudal neural gene *Sox1* expression (Fig. 1 H; Bertrand et al., 2000) and the appearance of the ventral patterning gene *Sax6* (Fig. 1 I; Bertrand et al., 2000). Moreover, the initiation of *Snail2* expression is close to the decay of FGF8 transcripts in the neural tube (Fig. 1 K), and high levels of *Snail2* mRNA coincided with the onset of the RA-synthesizing enzyme *Raldh2* expression in the adjacent mesoderm (Fig. 1 J).

In summary, NCC markers are progressively transcribed in a precise temporal–spatial order in the presumptive region of premigratory NCCs. Moreover, these data suggest that the onset of specification and EMT of NCCs could be controlled by FGF and RA signaling gradients (Fig. 1 L).

**FGF signaling controls the onset of trunk NCC specification through the MAPK pathway**

FGF signaling is involved in the maintenance of the neural caudal stem zone, and it also represses neuronal differentiation and ventral patterning (Bertrand et al., 2000; Diez del Corral et al., 2003). To determine whether the onset of NCC specification was regulated by the decline of the caudal gradient of FGF8 signaling, we used the FGF receptor (FGFR) type 1 (FGFR1) antagonist SU5402 (Mohammadi et al., 1997). Cultured HH11 embryos exposed to this drug for 4 h exhibited a severe down-regulation of the well-known FGF signaling target *Sprouty2*.
This did not alter the onset of Snail2 expression ($n = 12/15$; Fig. S2) in relation to control DMSO-treated embryos ($n = 13/15$; Fig. S2). However, FGFR1 inhibition either with PD173074 or SU5402 did not alter the expression of FoxD3, another NCC specifier ($n = 4$ and $7$, respectively; Fig. S1 and not depicted), with respect to control embryos ($n = 9$).

Together, these results indicate that FGF caudal gradient prevents the premature expression of the early NCC specifier Snail2 in the caudal neural tube.

Activation of the tyrosine kinase FGFR can initiate transduction via several downstream pathways including MAPK and phosphatidylinositol 3-kinase (PI3K; Schönwasser et al., 1998). Both pathways have been involved in the transmission of FGF signaling during early stages of neural development (Ribisi et al., 2000; Stavridis et al., 2007), and dual phosphorylated MAPK has been specifically detected in the caudal FGF signaling domain (Lunn et al., 2007). We investigated the potential role of the MAPK kinase (MEK)–extracellular signal-regulated kinase (ERK) regulatory pathway in the control of NCC development ($n = 13/13$; Fig. 2 B; Minowada et al., 1999) in comparison with control DMSO-treated embryos ($n = 0/12$; Fig. 2 A). After a 4-h culture with SU5402, Snail2 expression was enhanced in the dorsal part of the neural tube and was prematurely initiated at the level of Hensen’s node ($n = 11/12$; Fig. 2 F) in comparison with control embryos ($n = 4/18$; Fig. 2 E). A similar premature Snail2 up-regulation was observed using the specific FGFR1 inhibitor PD173074 ($n = 7/7$; Fig. S1; Skaper et al., 2000) and was quantified using quantitative RT-PCR as a twofold increase in relative Snail2 mRNA levels in PD173074 with respect to DMSO-treated samples in the neural tube at the level of the caudal PSM (2.06 [1.77–2.39]; Fig. S1). As a control, a similar quantitative RT-PCR analysis showed a threefold decrease in Sprouty relative mRNA levels in PD173074 with respect to DMSO-treated samples (0.31 ± 0.16; Fig. S1). PD173074 and SU5402 not only inhibit FGFR signaling but also target VEGF receptors (VEGFRs; Mohammadi et al., 1997, 1998).

To determine whether these effects are specifically mediated by FGFR signaling, we used a specific VEGFR inhibitor, KRN633 (Nakamura et al., 2004). This did not alter the onset of Snail2 expression ($n = 12/15$; Fig. S2) in relation to control DMSO-treated embryos ($n = 13/15$; Fig. S2). However, FGFR1 inhibition with PD173074 or SU5402 did not alter the expression of FoxD3, another NCC specifier ($n = 4$ and $7$, respectively; Fig. S1 and not depicted), with respect to control embryos ($n = 9$). Together, these results indicate that FGF caudal gradient prevents the premature expression of the early NCC specifier Snail2 in the caudal neural tube.
specification by selectively suppressing activation of MEK1 with the drug PD184352. In HH11 and 12 embryos cultured for 4 h, this drug effectively down-regulated the expression of a direct target of FGF signaling, Sprouty2 (n = 7/7; Fig. 2 C), and reduced ERK1/2 phosphorylation (92% decreased in phospho-ERK/relative to total ERK; n = 3; Fig. 2 I) in comparison with DMSO-treated embryos (n = 6; Fig. 2, A and I). As with inhibition of FGFR1, blockage of the MAPK pathway leads to a premature and enhanced caudal expression of the NCC specifier Snail2 (n = 4/4; Fig. 2 G). On the other hand, blockage of the PI3K pathway using the LY294002 treatment, which effectively reduces AKT phosphorylation (73% decreased in phospho-AKT/relative to total AKT; n = 3; Fig. 2 I), had very little effect on Sprouty2 (n = 1/7; Fig. 2 D; Echevarria et al., 2005) and on Snail2 mRNA expression (n = 0/4; Fig. 2 H). In conclusion, FGF signaling acting through the MAPK but not the PI3K pathway is required to prevent premature Snail2 expression in NCCs and, consequently, to delay NCC specification.

The FGF signaling pathway controls the initiation of trunk NCC emigration

Premature specification of NCCs could lead to premature emigration at axial levels previous to somite formation. In the aforementioned experiments, NCCs from embryos incubated with FGFR1 inhibitors for 4 h that prematurely expressed Snail2 were not able to emigrate, as assessed by the expression of Sox10 (n = 9; Fig. S1 and not depicted). To better test the consequence on NCC emigration, we examined the effect of blocking FGF signaling for longer periods of time using a dominant-negative FGFR1-EYFP protein (dn-FGFR1-EYFP; Yang et al., 2002) introduced into the right caudal hemitube at HH10 and 11. 18–20 h after electroporation, this construct effectively blocked the expression of FGF signaling targets such as Sprouty2 (n = 7/9; Fig. S3; Minowada et al., 1999) and induced Pax6 expression (n = 5/6; Fig. S3; Bertrand et al., 2000) in the neural tube in comparison with control pCIG embryos (n = 6 and 4/5, respectively; Fig. S3). Remarkably, dorsal neural progenitors expressing the dn-FGFR1-EYFP protein prematurely activated Snail2 (n = 4/4; not depicted) and Sox10 expression and exited the neural tube (n = 9/10; Fig. 3 B, arrowhead) in comparison with the control side or with cells electroporated with a control EGFP (n = 7; Fig. 3 A). Moreover, the progressive disruption of the basal lamina, identified by Laminin expression (Martins-Green and Erickson, 1987), was also more advanced in the side where FGF signaling had been blocked (n = 5/7; Fig. 3 F) with respect to the control side or control embryos (n = 7; Fig. 3 C).

Figure 2. FGF signaling controls the onset of trunk NCC specification through the MAPK pathway. (A–H) Stage HH11 and 12 embryos cultured for 4 h in the presence of control (DMSO) or inhibitors for FGFR1 (SU5402), MEK (PD184352), or PI3K (LY294002) and analyzed for Sprouty2 expression (A–D) and for Snail2 (E–H). Black arrowheads point to the last formed somite. (E′–H′) Transverse sections at the level are indicated by horizontal lines in the corresponding figures. (I) Western blot analysis from embryos treated with DMSO or the different inhibitors to detect levels of phosphorylated ERK (p-ERK) in comparison with total ERK and levels of phosphorylated AKT (p-AKT) in comparison with total AKT. Results shown are representative of at least three experiments. Bars: (A, for A–H) 100 µm; (E′, for E′–H′) 50 µm.
significantly increased in FGFRdn embryos with respect to control pCIG (68 ± 28% in FGFRdn and 42 ± 15% in pCIG; Fig. S4 H). Interestingly, the relative number of GFP–Sox5+ cells was also significantly increased in FGFRdn embryos with respect to control pCIG (192 ± 61% in FGFRdn and 65 ± 7% in pCIG; Fig. S4 H). All these data suggest that the premature migration of the FGFRdn-GFP–electroporated cells was also promoting the exit of adjacent non-GFP NCCs, and, thus, the effect on FGFR blockage would be in part non–cell autonomous on NCCs.

Moreover, although NCC markers such as Sox10, Snail2, and Sox5 were prematurely expressed (Fig. 3 B and not depicted) and more ventrally expressed (Sox10; Fig. 3 B), expression of dorsal markers such as Pax7 and Msx1 was not affected in the neural tube at epithelial somite levels (n = 6 and 10, respectively; Fig. 3, G and H). In conclusion, blocking FGF signaling in the neural tube adjacent to PSM promotes premature NCC specification and emigration from the neural tube.

To determine whether ectopically maintaining the FGF could prevent NCCs from delaminating, we placed FGF8/4-soaked beads in contact with the HH11 and 12 dorsal neural tube at a level in which NCCs have already been specified but before they start Sox10 expression. As expected, 18 h later, FGF4/8 promoted expression of its direct target Sprouty2 in the adjacent neuroepithelial cells in HH15–17 embryos (FGF8 n = 3

Migratory NCCs are a mixture of different population subtypes identified by differential gene expression (Del Barrio and Nieto, 2004; Krispin et al., 2010). To determine whether the NCCs delaminating prematurely upon FGF signaling blockage were specifically those expressing Sox10, we examined the effects on a whole cohort of NCC markers. After 18–20 h of dn-FGFR1 expression using a nuclear EGFP (pCIG-FGFRdn) as a reporter, there was an increase in the migratory NCC subpopulations expressing Pax7 (n = 10/11; 170 ± 9% with respect to the control side; Figs. 3 G and S4 G), Sox5 (n = 13/13; 159 ± 24% increase; Fig. S4, D and G), AP2 (n = 7/7; 149 ± 13% increase; Fig. S4, E and G), Pax3, Snail2, and FoxD3 (n = 9/9, 4/4, and 5/6, respectively; not depicted) transcription factors in comparison with the control side and with pCIG-electroporated embryos (n = 8, 8, 6, 3, and 8, respectively; Figs. 3 D and S4, A–C).

Remarkably, not all the prematurely migrating NCCs were electroporated GFP+ cells (Fig. S4 D). To determine whether there was a non–cell-autonomous effect upon FGFR blockage, we analyzed the number of Sox5+ migratory cells that were either GFP+ or GFP− in relation to the number of migratory Sox5+ cells in the control nonelectroporated side (as a way of normalization to reduce axial level variations; Fig. S4 H). The relative number of double GFP+ Sox5+ migratory cells with respect to the total Sox5+ migratory cells in the control side was significantly increased in FGFRdn embryos with respect to control pCIG (68 ± 28% in FGFRdn and 42 ± 15% in pCIG; Fig. S4 H).

Interestingly, the relative number of GFP–Sox5+ cells was also significantly increased in FGFRdn embryos with respect to control pCIG (192 ± 61% in FGFRdn and 65 ± 7% in pCIG; Fig. S4 H). All these data suggest that the premature migration of the FGFRdn–GF–electroporated cells was also promoting the exit of adjacent non–GF NCCs, and, thus, the effect on FGFR blockage would be in part non–cell autonomous on NCCs. Moreover, although NCC markers such as Sox10, Snail2, and Sox5 were prematurely expressed (Fig. 3 B and not depicted) and more ventrally expressed (Sox10; Fig. 3 B), expression of dorsal markers such as Pax7 and Msx1 was not affected in the neural tube at epithelial somite levels (n = 6 and 10, respectively; Fig. 3, G and H). In conclusion, blocking FGF signaling in the neural tube adjacent to PSM promotes premature NCC specification and emigration from the neural tube.
EGFP-glycosylphosphatidylinositol (GPI; cell membrane)–expressing vectors into stage HH10 chick embryos to follow individual cells (Fig. 5, A and C). Using MetaMorph software, we first compared the behavior of the migratory NCCs during the initiation or first wave of migration at the level of dissociating somites (somites V–XII) in control EGFP-GPI embryos (Fig. 5 [A, B, and G] and Video 1) in relation to dn-FGFR1-EYFP–electroporated NCCs that started prematurely migrating adjacent to epithelial somites (somites I–IV; Fig. 5 J and Video 3). We observed a similar speed of migration (16.2 ± 1.8 µm/h with respect to 12.4 ± 1 µm/h; Fig. 5 E) and a similar migration straightness (distance covered/length of the trajectory; 0.28 ± 0.07 with respect to 0.15 ± 0.03; Fig. 5 F) of the two spatially different located populations. These results indicate that the prematurely migrating NCCs, upon FGF signaling blockage, at epithelial somite levels have a similar behavior to the first wave of migrating control NCCs facing dissociating somites.

Conversely, we next analyzed the behavior of dn-FGFR1-EYFP–expressing NCCs (Fig. 5 [C, D, and H] and Video 2) that were migrating next to dissociating somites in comparison with the first migrating NCCs expressing EGFP-GPI (Fig. 5 A). The mean speed of the migratory NCCs upon a 12-h period was higher in NCCs expressing dn-FGFR1-EYFP (20.2 ± 1.9 µm/h; Fig. 5 E) with respect to control NCCs (12.4 ± 1 µm/h; Fig. 5, B and E). Moreover, tracking NCC trajectories revealed that most dn-FGFR1–expressing NCCs showed a straight trajectory (Fig. 5 H) contrary to control NCCs that exhibited tortuous trajectories and forward and backward movements (straightness factor of 0.39 ± 0.1 with respect to 0.15 ± 0.03; Fig. 5, F and G).

Figure 4. Ectopic FGF inhibits NCC specification and emigration. (A–F) Stage HH11 and 12 embryos were exposed to heparin-coated beads either with PBS (A, C, and E) or with FGF4 (B, D, and F) after 16–18 h in culture. In situ hybridization analysis of Sprouty, Sox10, and Snail2 expression. Black arrowheads point to the last formed somite. [A–F] Transversal sections through the region where the bead was located, indicated by horizontal lines in the corresponding figures. Black arrows point to ectopic expression in B’ and to down-regulation of expression in D’ and F’. Bars: [A, for A–F] 100 µm; [A’, for A’–F‘] 40 µm.

The timing of NCC emigration confers different migratory behavior

In the trunk region of avian embryos, NCCs migrate in a stereotypical manner, leading to a general ventral to dorsal order of derivative colonization beginning with sympathetic ganglia and, finally, melanocytes (Le Douarin and Kalcheim, 1999). The final localization of crest cells can be predicted from their relative ventrodorsal position within the premigratory domain or by their time of emigration (Henion and Weston, 1997; Krispin et al., 2010). However, it is also possible that distinct properties of the differentiated mesodermal cells that the NCCs encounter during migration influence their trajectories (Harris and Erickson, 2007).

To determine whether the NCCs prematurely forced to delaminate upon blockage of FGF signaling had alterations in their migratory behavior, we followed early NCC migration using wide-field real-time microscopy of trunk dorsal neural tubes in slice culture (Wilcock et al., 2007; Ahlstrom and Erickson, 2009). We used electroporation of H2B-RFP (nucleus)– and EGFP-glycosylphosphatidylinositol (GPI; cell membrane)–expressing vectors into stage HH10 chick embryos to follow individual cells (Fig. 5, A and C). Using MetaMorph software, we first compared the behavior of the migratory NCCs during the initiation or first wave of migration at the level of dissociating somites (somites V–XII) in control EGFP-GPI embryos (Fig. 5 [A, B, and G] and Video 1) in relation to dn-FGFR1-EYFP–electroporated NCCs that started prematurely migrating adjacent to epithelial somites (somites I–IV; Fig. 5 J and Video 3). We observed a similar speed of migration (16.2 ± 1.8 µm/h with respect to 12.4 ± 1 µm/h; Fig. 5 E) and a similar migration straightness (distance covered/length of the trajectory; 0.28 ± 0.07 with respect to 0.15 ± 0.03; Fig. 5 F) of the two spatially different located populations. These results indicate that the prematurely migrating NCCs, upon FGF signaling blockage, at epithelial somite levels have a similar behavior to the first wave of migrating control NCCs facing dissociating somites.

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form for the three RARs (RARdn–internal ribosome entry site (IRES)–GFP; Novitch et al., 2003). As expected, blocking RAR signaling at HH12 and 13 decreased the expression of Pax6 in neural precursors 24 h after electroporation (Fig. S5; Novitch et al., 2003). In HH12 and 13 embryos in which RA signaling was blocked where NCC emigration was starting, the number of cells expressing \(\text{Sox10}^+\) was reduced, and there was a dramatic delay in the emigration of \(\text{Sox10}^+\)NCCs (\(n = 5/5\); Fig. 6 C) in comparison with the pCIG-electroporated cells (\(n = 5/5\); Fig. 6 A). However, more caudally, the initiation of the expression of \(\text{Snail2}\) and \(\text{FoxD3}\), early markers of NCC specification, was not affected (\(n = 5\) and 6, respectively; Figs. 6 F and S5) in comparison with control pCIG-electroporated embryos (\(n = 3\) and 8, respectively; Fig. 6 E), indicating that RA is required to establish the onset of NCC emigration independently of NCC specification.

Nevertheless, in HH11–13 embryos electroporated at the level of ongoing NCC migration (somites V–X), RA signaling blockage did not affect the EMT of \(\text{Sox10}^+\)-expressing NCCs (\(n = 4\); Fig. 6 B) compared with the control (\(n = 6\); Fig. 6 A). Similarly, blocking RA signaling did not alter the EMT of AP2\(^+\) NCCs in comparison with control (\(n = 6\); Fig. S5). These data indicate that RA signaling is required to establish the onset but not the maintenance of NCC emigration.

To determine whether RA signaling was not only necessary but sufficient to control NCC emigration, we used a higher speed and straightness of the more advanced delaminating NCCs expressing dn-FGFR1-EYFP could be a property of a more mature late-migrating NCC population in spite of both control and experimental cells facing the same somitic context. In summary, this analysis reveals distinct migratory properties according to the time of exit from the neural tube.

**RA signaling is required to control the timing of NCC emigration**

It has been shown that somites are required for the control of the onset of NCC emigration (Sela-Donenfeld and Kalcheim, 2000). However, the molecular nature of those signals emanating from the somites remains unknown. RA signaling is highly active in the neural tube at somitic levels, as revealed by in vivo reporter activity driven by RA response elements (RAREs) in mouse embryos (Rossant et al., 1991) and by the expression of a readout of RA signaling, \(\text{RAR-}\beta\) (the RARE promoter elements of which were used in the RARE-LacZ mouse) in chicken embryos (Olivera-Martinez and Storey, 2007). Moreover, the opposing gradients of FGF and RA pathways operate in concert as a switch that controls progressive neural and limb development (Diez del Corral and Storey, 2004).

To test whether RA could be involved in the control of NCC emigration in concert with FGF signaling, we performed neural tube electroporation of a truncated version of the human \(\alpha\)-RA receptor (RAR [RAR-\(\alpha\)]) that acts as a dominant-negative form for the three RARs (RARdn–internal ribosome entry site (IRES)–GFP; Novitch et al., 2003). As expected, blocking RA signaling at HH12 and 13 decreased the expression of Pax6 in neural precursors 24 h after electroporation (Fig. S5; Novitch et al., 2003). In HH12 and 13 embryos in which RA signaling was blocked where NCC emigration was starting, the number of cells expressing Sox10 was reduced, and there was a dramatic delay in the emigration of Sox10\(^+\) NCCs (\(n = 5/5\); Fig. 6 C) in comparison with the pCIG-electroporated cells (\(n = 5/5\); Fig. 6 A). However, more caudally, the initiation of the expression of Snail2 and FoxD3, early markers of NCC specification, was not affected (\(n = 5\) and 6, respectively; Figs. 6 F and S5) in comparison with control pCIG-electroporated embryos (\(n = 3\) and 8, respectively; Fig. 6 E), indicating that RA is required to establish the onset of NCC emigration independently of NCC specification.

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To determine whether RA signaling was not only necessary but sufficient to control NCC emigration, we used a
at the rostral PSM level (Fig. 7 D) in comparison with normal embryos (Fig. 7 C). To determine how early RA signaling is required for NCC specification, \( \textit{Pax7} \) expression, a key mediator of neural plate border specification (Basch et al., 2006), was examined. In VAD embryos, \( \textit{Pax7} \) is absent at the neural plate border (Fig. 7 F) in comparison with normal quails (Fig. 7 E). As the VAD quails represent an early RA-deficient state, these findings suggest that RA signaling is required for the earliest steps in dorsal and NCC specification during gastrulation before the RA activity is restricted to the neural tube adjacent to rostral PSM and is not required for NCC specification during neural tube elongation.

FGF maintains Noggin gradient in the caudal neural tube

It has been reported that an R-C gradient of BMP signaling in the dorsal neural tube, generated by the graded expression of the BMP antagonist Noggin, is involved in determining the onset of NCC emigration (Sela-Donenfeld and Kalcheim, 1999). As the Noggin gradient is controlled by unknown signals coming from the mesoderm (Sela-Donenfeld and Kalcheim, 2000), we tested whether FGF and/or RA could be those signals. First, we determined whether FGF signaling was required for Noggin expression, as both coincide in the caudal neural tube (Figs. 1 K and 8 A). In cultured embryos at stages HH10 and 11 briefly exposed (4 h) to the FGFR1 inhibitor PD173074, Noggin expression in the neural tube

Retinoid signaling is required for very early neural crest specification

Our results have shown that RA signaling is not required for NCC specification during neural tube elongation. However, in \( \textit{Raldh2} \) mutants, the expression of early NCC markers is severely down-regulated (Ribes et al., 2009). To determine whether that was exclusive of mammalian embryos, we analyzed expression of neural crest markers in vitamin A–deficient (VAD) quails, which represent a retinoid-deficient state (Maden et al., 1996). In normal quails, \( \textit{Snail2} \) expression starts in the dorsal neural tube next to mid-PSM (Fig. 7 A), whereas it is strongly reduced in VAD NCCs (Fig. 7 B). Similarly, in RA-deficient conditions, \( \textit{Sox9} \) expression is clearly absent from the dorsal neural tube at the rostral PSM level (Fig. 7 D) in comparison with normal embryos (Fig. 7 C). To determine how early RA signaling is required for NCC specification, \( \textit{Pax7} \) expression, a key mediator of neural plate border specification (Basch et al., 2006), was examined. In VAD embryos, \( \textit{Pax7} \) is absent at the neural plate border (Fig. 7 F) in comparison with normal quails (Fig. 7 E). As the VAD quails represent an early RA-deficient state, these findings suggest that RA signaling is required for the earliest steps in dorsal and NCC specification during gastrulation before the RA activity is restricted to the neural tube adjacent to rostral PSM and is not required for NCC specification during neural tube elongation.

Constitutively active form of RAR-\( \alpha \) fused to the transcriptional activator domain of VP16 (VP16-RAR; Novitch et al., 2003). This construct activates RA target genes in a ligand-independent manner (Castro et al., 1999). VP16-RAR–electroporated neuroepithelial cells at the rostral PSM level delaminated prematurely from the dorsal neural tube in comparison with control embryos, as assessed by \( \textit{Sox10} \) expression (Fig. 6 D). Moreover, the initiation of \( \textit{Snail2} \) expression in the caudal neural tube was not affected (Fig. 6 D). In summary, RA signaling is both necessary and sufficient to set the right timing for NCC emigration.
In summary, these results indicate that RA does not control NCC emigration through the down-regulation of Noggin gradient. Moreover, FGF signaling is sufficient but not necessary for Noggin expression, suggesting that the decrease in FGF signaling is required for Noggin down-regulation at somitic levels.

FGF and RA regulate NCC emigration in part through the control of Wnt1 expression

It has been reported that Wnt ligands are necessary to control Snail2 expression in NCCs in vivo and are sufficient to induce NCCs by in vitro assays (García-Castro et al., 2002). Moreover, blocking canonical Wnt signaling prevents NCC emigration (Burstyn-Cohen et al., 2004). However, apart from the control by BMP signaling (Burstyn-Cohen et al., 2004), the regulation of the Wnt pathway has not been extensively analyzed in the NCC during axial elongation. We looked to determine whether at NCC emigration FGF and RA signaling could be acting through the modulation of Wnt1.

In cultured embryos at stages HH12–14 briefly treated (4 h) with the FGFR1 inhibitor PD173074, the R-C gradient of Wnt1 expression was prematurely initiated at more caudal levels (n = 4/7; Fig. 9 B) in comparison with control DMSO-treated embryos (n = 10; Fig. 9 A). Conversely, exposing neural tube explants from rostral PSM levels to FGF4 for 4 h in culture inhibited Wnt1 expression (FGF4 n = 6; control n = 8; Fig. 9, C and D; described for longer incubation times in Olivera-Martínez et al. 2002).

Snail2

Sox9

Pax7

was not affected in comparison with control DMSO-treated embryos (n = 8 and 8, respectively; Fig. 8, A and B). Similarly, after 18–20 h of dn-FGFR1 electroporation, Noggin and BMP4 expression were not altered in relation to control GFP cells (n = 7 and 4, respectively; Fig. 8 [C and D] and not depicted). However, exposing neural tube explants at the rostral PSM level to high doses of FGF4 promoted Noggin expression after just 4 h in culture (n = 10; Fig. 8 L) relative to BSA control explants (n = 9; Fig. 8 K), whereas Snail2 expression (control n = 4; FGF4 n = 3; Fig. 8, G and H) was down-regulated, as previously observed with FGF4/8 beads (Fig. 4 F). Sox10 expression was also down-regulated after 8 h, the time required for Sox10 mRNA to be detected in control explants (control n = 12; FGF4 n = 9; Fig. 8, I and J). As a control, Pax6 was down-regulated by FGF4 in neural tube explants at the somitic level (FGF4 n = 16; control n = 15; Fig. 8, E and F; Bertrand et al., 2000). Thus, the caudo-rostral FGF signaling gradient is sufficient but not necessary for Noggin expression.

To test whether RA was the signal responsible for repression of Noggin transcription, RARdn was electroporated in NCCs adjacent to rostral PSM. RARdn-electroporated NCCs exhibited similar levels of Noggin expression (n = 4; Fig. 8 D) to that of control (n = 3; Fig. 8 C). Similarly, in trunk explants at the caudal PSM level exposed for 4 h to the RAR agonist TTNPB, which induced high levels of Pax6 (TTNPB n = 10; control n = 9; Fig. 8, M and N; Novitch et al., 2003), Noggin expression was similar to that in control DMSO-treated explants 4 and 8 h after culture (control, 4 h, n = 10; TTNPB, 4 h, n = 12; control, 8 h, n = 4; TTNPB, 8 h, n = 4; Fig. 8, O and P).

In summary, these results indicate that RA does not control NCC emigration through the down-regulation of Noggin gradient. Moreover, FGF signaling is sufficient but not necessary for Noggin expression, suggesting that the decrease in FGF signaling is required for Noggin down-regulation at somitic levels.
In summary, FGF signaling prevents the premature expression of \textit{Wnt1}, whereas RA signaling triggers the initiation of \textit{Wnt1} expression in the dorsal neural tube only at levels in which the NCCs are already specified. Given the short time required for its action in culture, FGF and RA could control the timing of NCC emigration through direct changes in \textit{Wnt1} expression.

\section*{Discussion}

During embryonic development, it is essential to coordinate the formation of adjacent tissues that maintain close functional relationships. In this work, we have established the mechanism that ensures the correct coordination between the formation of the central and peripheral nervous system in the trunk during the extension of the body axis. We have demonstrated that FGF and RA signaling set the correct timing for the crucial step of NCC EMT and emigration, which is necessary for the distribution of peripheral nervous system progenitors along the body axis.

Our work reinforces the idea that dorsal neuroepithelial progenitors in the caudal neural tube are maintained in an uncommitted state as a result of the presence of a strong FGF–MAPK signaling pathway. Therefore, the caudo-rostral gradient of FGF and Storey, 2007). In conclusion, FGF signaling levels control the initiation of \textit{Wnt1} expression in the dorsal neural tube.

RA signaling activation in the neural tube adjacent to the somitic level coincides with the initiation of \textit{Wnt1} expression (Figs. 1 K and 9 A). To determine whether RA signaling was required for the control of \textit{Wnt1}, we examined HH10 and 11 VAD quail embryos (Fig. 9, G and H). Strikingly, in VAD embryos, expression of \textit{Wnt1} and \textit{Wnt3a} (Fig. 9, I and J; at older stages in Wilson et al., 2004) was strongly reduced, particularly at the spinal cord level in comparison with normal quail embryos. As this could be caused by the early NCC specification effect described in Fig. 7, we performed trunk explant experiments in chick embryos, where the NCCs have already been specified. Thus, in trunk explants at the rostral half PSM level, \textit{Wnt1} expression was enhanced and caudally expanded after 4 h of exposure to the RAR agonist TTNPB in comparison with DMSO-treated explants (\(n = 4\) and 4, respectively; Fig. 9, E and F). However, this effect on \textit{Wnt1} expression was not observed in trunk explants at the caudal PSM level exposed to TTNPB for 4 or 8 h in culture (DMSO, 4 h, \(n = 6\); TTNPB, 4 h, \(n = 6\); DMSO, 8 h, \(n = 4\); TTNPB, 8 h, \(n = 4\); unpublished data), where NCCs do not express specifiers like \textit{Snail2} at the time of excision.

In summary, FGF signaling prevents the premature expression of \textit{Wnt1}, whereas RA signaling triggers the initiation of \textit{Wnt1} expression in the dorsal neural tube only at levels in which the NCCs are already specified. Given the short time required for its action in culture, FGF and RA could control the timing of NCC emigration through direct changes in \textit{Wnt1} expression.
Xenopus embryos concern the cephalic NCCs (Monsoro-Burq et al., 2003; Hong and Saint-Jeannet, 2007), our work shows that the FGF pathway is, so far, the only signaling pathway clearly controlling the onset of NCC specification at the trunk level. This FGF function could be another example of a general FGF role in controlling the onset of differentiation of all cell types, as they are generated at the tail end.

Our study also points out the relevance of controlling Snail2 expression to initiate NCC specification. Not only is Snail2 expression ectopically activated just after 4 h of FGF signaling blockage in comparison with FoxD3, which remained unchanged, but Snail2 is also one of the earliest NCC specifier genes being expressed during trunk NCC formation (Fig. 1; this study) and also during the caudal-most NCC development (caudal vertebrae level) in which Snail2 expression also precedes that of Sox9 and FoxD3 (Osório et al., 2009). The regulation of Snail gene expression by FGF seems to be context dependent during development. Although Snail1 is induced by FGF8 in the lateral mesoderm (Boettger et al., 1999) and primitive streak (Ciruna and Rossant, 2001) and by FGF3 in bone development (de Frutos et al., 2007), it is repressed by FGF in the somites (this study). Furthermore, controlling the right levels of FGF signaling is essential to modulate Snail gene expression. In Xenopus embryos, FGF signaling is involved in cephalic NCC induction (Monsoro-Burq et al., 2003). However, an excess of FGF signaling inhibits Snail2 expression and NCC formation (Hong and Saint-Jeannet, 2007).
In contrast to FGF signaling acting at the caudal level, RA signaling is operating at somitic levels. Our data demonstrate that, at least during trunk elongation, RA signaling is not required for the control of NCC specification but regulates the timing of NCC emigration. Thus, the onset of Snail2 expression is not affected by changes in RA signaling, but the onset of emigration of Sox10+ NCC cells at the midrostral PSM level can be modulated by changing RA signaling.

BMP activity regulates emigration of already specified NCCs (Sela-Donenfeld and Kalcheim, 1999) through the control of Wnt1 expression (Burstyn-Cohen et al., 2004). RA activity controls emigration modulating, together with FGF signaling, the onset of Wnt1 expression in the dorsal neural tube at rostral PSM levels (Fig. 10). Thus, our experiments have established that both FGF and RA signaling act as upstream pathways to the BMP/Noggin-Wnt1 signaling cascade to set the right timing for NCC emigration to keep it in register with somite formation. However, the effect of RA on emigration only occurs at rostral PSM and the last-formed somite level. More rostrally, inhibiting RA signaling does not affect ongoing emigration probably because BMP signaling is sufficient to maintain the process (Burstyn-Cohen et al., 2004). Conversely, at most, caudal neural tube level changes in FGF or RA signaling are not sufficient to induce premature Wnt1 expression (Fig. 9), and, consequently, NCCs cannot delaminate from the most caudal part of the neural tube. This time, the absence of BMP signaling at caudal levels could be responsible for the impossibility of emigration.

The consequences of altering the timing of migration could be crucial for the development of the NCC derivatives. It has recently been shown that the final localization of crest cells can be predicted by their time of emigration (Krispin et al., 2010). Here, we have demonstrated that the different NCC populations do differ in their migratory behavior, as the first migratory cells (including those prematurely forced to migrate in our experiments; Fig. 5) move at lower speed and follow more tortuous paths than later-migrating NCCs. It remains to be seen whether changes in migratory behavior adopted in an inappropriate context (nondifferentiated somites) will have consequences in the final differentiation of the NCCs.

In summary, our data show that there is a limited time window during which the onset of the NCC emigration can be modulated once those cells have acquired the expression of the essential gene network of the NCC specification program. That window coincides with the region where FGF and RA gradients collide, and it is required to keep the emergence of peripheral nervous system progenitors in register with the progressive programs of spinal cord neurogenesis and somite development during trunk axial elongation.

Materials and methods

In situ hybridization

Chicken eggs were incubated at 38°C in an atmosphere of 70% humidity. Embryos were staged according to HH. Embryos were fixed overnight at 4°C with 4% PFA in PBS, rinsed, and processed for whole-mount in situ hybridization. In brief, embryos were treated with 1% H2O2 and 10 µg/ml proteinase K, rinsed in 4% PFA, and then hybridized overnight at 57°C in 5x SSC buffer containing 50% formamide with digoxigenin- or fluorescein-labeled riboprobes as previously described (Martinez-Morales et al., 2010). The chick Pax7, Snail2, Sox5, Sox10, FoxD3, Sox9, Pax6, Radh2, FGF8, Sprouty2, Noggin, and Wnt1 riboprobes have been previously described (Cheung and Briscoe, 2003; Diez del Corral et al., 2003; Burstyn-Cohen et al., 2004; Perez-Alcalá et al., 2004). Chicken Pax3 and Pax7 cDNAs were provided by H. Nakamura (Tokohu University, Sendai, Japan). Chicken VE-cadherin was cloned using primers 5’-TCCTTACATTCTCTCA-TCCTGTA-3’ (forward) and 5’-TGCCGCTCCAACACCTTACCTC-3’ (reverse) following recommendations from C.A. Roselló (Centro Nacional de Investigaciones Cardiovasculares, Madrid, Spain). 4–15 embryos were analyzed for each experimental condition. The probe hybridization was detected with AP coupled with either antidigoxigenin Fab fragments (Roche) or antifluorescein Fab fragment (Roche) and developed with nitro blue tetrazolium chloride/5-bromo-4-chloro-3-indolylphosphate-p-toluidine salt (BCIP) or 2-[4-iodophenyl]-2-[4-nitrophenyl]-5-[4-tetrazolium]-1,2-phenylenediamine chloride/BCIP. Hybridized embryos were postfixed in 4% PFA and vibratome sectioned, and immunostaining was performed as described in the Immunochemistry section to visualize GFP+-electroporated cells.

Embryo and explant culture

Chicken embryos at stages HH11 and 12 were cultured onto collagen beds [prepared as a mixture containing 2.9 mg/ml collagen rat tail (BD), 1x L15 (Invitrogen), and 0.6% NaHCO3 to achieve proper polymerization]
covered with Opti-MEM medium (Invitrogen) supplemented with 5% FBS, 2 mM glutamine, and 50 µg/ml gentamycin. Inhibitors were added to the culture medium (100 µM SU5402, 100 µM PD184352, 10 µM PD173074, 100 µM LY294002, 10 µM KRN633, or DMSO vehicle). Embryos were cultured for 4 h at 37°C and 5% CO2 before fixation in 4% PD173074, 100 µM LY294002, 10 µM KRN633, or DMSO vehicle).

serum, 2 mM glutamine, and 50 µg/ml gentamycin. Inhibitors were added (Casanova et al., 2011). After incubation at 38°C for 18–20 h, embryos were either processed for whole-mount in situ hybridization or dissected to obtain explants for quantitative RTPCR. Re-inhibitor-deficient quail embryos were a gift from E. Gole and M. Maden (Medical Research Council Centre for Developmental Neurobiology, King’s College London, London, England, UK), and normal and deficient quail embryos were fixed and processed together for in situ hybridization (Diez del Corral et al., 2003).

Explants were dissected from HH9–13 embryos treated with trypsin to separate the neural tissue from the mesoderm when necessary (Diez del Corral et al., 2002). Explants were cultured in collagen beds and cultured covered in Opti-MEM medium as described above containing either BSA or 360 ng/ml PDGF (both supplemented with 0.1% FBS or PDGF) or containing 10 µM KRN633 (EMD), 0.1% DMSO, or 10 µM T1NP8 (EMD).

Western blot analysis

Western blot analysis was performed by an established method. 20 µg protein samples of each total cell extract were separated by 12% SDS-PAGE, transferred to a polyvinylidene fluoride Immobilon-P membrane (Millipore), and probed with antibodies anti–phospho-AKT (rabbit IgG; Cell Signaling Technology), anti–total AKT1/2 (H136; Santa Cruz Biotechnology, Inc.), anti–phospho-ERK1/2 (Thr202/Tyr204; Cell Signaling Technology), and anti–total Erk1/2 (Invitrogen). Signals were detected with HRP-conjugated secondary antibodies (Jackson ImmunoResearch Laboratories, Inc.) using an ECL Advance Western Blotting Detection kit (GE Healthcare). Quantitative analysis was obtained with a densitometer (GS-800, Bio-Rad Laboratories). Results shown are representative of three or more experiments.

Quantitative real-time PCR

Explants of neural tube together with notochord at caudal PSM levels were dissected from embryos cultured for 4 h in the presence of DMSO or PD173074, and total RNA was isolated using the QuickGene RNA tissue kit SF (FujiFilm). cDNA was synthesized with SuperScript III DNA polymerase (Invitrogen) and random primers (Invitrogen). Real-time PCR was performed in a 7500 PCR System using Power SYBR green Master Mix (Applied Biosystems) and sequence-specific primers (Sigma-Aldrich). Sequences of Snail2 primers were 5′-AGCCAAAATCAAGCGAAGCTC-3′ (forward) and 5′-TGATAGGGCATTGGTAGCTTC-3′ (reverse), and sequences for Sprouty2 were 5′-ATGCATCGGCTGCGCTGCTTC-3′ (forward) and 5′-TGCTCCAAATCTTTCGCTTC-3′ (reverse). 18S ribosomal RNA was used as a reference gene. Sprouty2 levels were determined using the standard curve method, whereas Snail2 levels were determined by the comparative Ct method following recommendations by Applied Biosystems based on validation experiments assessing whether efficiencies of target and reference are approximately equal. Three lots of 4–12 explants generated in three independent experiments were used. Each of these samples was retrotranscribed, the product was used as a template for each pair of primers in triplicate-well PCR reactions, and the PCR procedure was performed twice.

Constructs

Truncated chick FGFR1c (aa 1–425) fused to EYFP in a Clontech vector (pEYFP-N1) to generate dn-FGFR1-EYFP was provided by C. Weijer (University of Dundee, Dundee, Scotland, UK; Yang et al., 2002) and was also inserted into pCAGGS-ires–nuclear EGFP (pCIG; Niwa et al., 1991) to generate pCIG-FGRFdn. The pCAGGS vector containing a membrane-tagged EGFP-GPI was used as a control. cDNAs encoding human RARα truncation mutant 403 (RARαR403; Damm et al., 1993) and a V166–RARα fusion (Castra et al., 1999) cloned into pCIG were provided by B. Novitch (University of California, Los Angeles, CA; Novitch et al., 2003). In this case, pCIG was used as a control vector.

Chick in ovo electroporation

Chick embryos were electroporated with purified plasmid DNA (Qiagen) at 1–2 µg/ml in PBS with 50 ng/ml Fast green, as previously described (Martin-Morales et al., 2010). In brief, plasmid DNA was injected into the lumen of HH10 and 12 neural tubes, electrodes were placed on either side of the neural tube, and a train of electric pulses (five pulses at 14 V for 50 ms) was applied using an electroporator (TSS20; Intracel). Eggs were further incubated for 24–48 h and were assayed for EGFP expression in the neural tube. Subsequently, the embryos were fixed and processed for immunohistochemistry or in situ hybridization.

Immunohistochemistry

Embryos were fixed for 2–4 h at 4°C with 4% PFA in PBS, and they were immersed in 30% sucrose solution, embedded in optimal cutting temperature medium, and sectioned on a cryostat (CM1900; Leica). Alternatively, embryos were embedded in 5% agarose/10% sacarose and were sectioned at 30 µm in a vibratome (VT1000S; Leica). For immunohistochemistry, 10-µm cryostat or 40-µm vibratome agarose sections were permeabalized with 0.5% Triton X-100, blocked with 10% FBS, and incubated overnight at 4°C with the primary antibody. After washing, the cryostat sections were incubated for 1 h with secondary antibodies. Primary antibodies against the following proteins were used: Sox5 (Perez-Alcala et al., 2004), GFP (Invitrogen), and Laminin (Sigma-Aldrich). Monoclonal antibodies against Pax7, Mx1 (4G1), Pax6, AP2 (3B5), Pax3, and Snail (62.1E6) were all obtained from the Developmental Studies Hybridoma Bank (developed under the auspices of the National Institute of Child Health and Development and maintained by the University of Iowa). Alexa Fluor 488– and Cy3-conjugated anti–mouse- or rabbit secondary antibodies (Invitrogen) were used for detection, and, after staining, the sections were mounted in Citiflour mounting medium plus bisbenzimide and photographed using a true confocal scanning microscope (SP5; Leica).

Real-time imaging

Embryos at stages HH10 and 11 were electroporated with a concentration of 0.2 µg/µl H2B-EGFP and either 1 µg/µl GFP-GPI or dn-FGFR1-EYFP. Embryos were incubated at 38°C for 18 h (for imaging at stages HH15 and 16). Slices of ~150 µm were taken from the trunk on stage HH16–18 chicken embryos (Wilcock et al., 2007) at the axial level of caudal, epithelial (somite I–IV), or rostral dissociated (somite V–XII) somites. Explants were cultured as described by Wilcock et al. (2007). In brief, slices of ~150 µm were taken with a microknife, embedded in rat tail collagen type I in coverslip-based Petri dishes (Wilco glass-bottom dish [Intracel] and GW43522-coated with poly-lysine [Sigma-Aldrich]), and cultured in Neurobasal medium without phenol red (Invitrogen) and supplemented with B27 to a final 1× concentration with glucose and gentamycin maintained at 37°C with 5% CO2/air for ~4 h before imaging. Slices were imaged on a microscope workstation (Delta Vision Core; Applied Precision; Wilcock et al., 2007). The images were taken at least 20 µm past the cut surface of the slice and only where the ectoderm was intact. Images were captured using a 40× 1.30 NA objective lens with a xenon lamp (Applied Precision). Optical sections (exposure time = 50 ms, 512 × 512 pixels; bin = 2 × 2) were spaced by 2 µm and imaged at 7-min intervals for up to 24 h. The images obtained were deconvolved and projected using Delta Vision software (softWoRx, Applied Precision). Meta Morph software (Molecular Devices) was used to analyze the neural crest trajectory and the speed of migration. For each experimental condition, 10 cells were analyzed during a period of ~4–12 h from at least four embryos.

Image acquisition and processing

Transmitted light images were acquired in an upright microscope (Eclipse B01; Nikon) with Digital Sight DS-U1 or DXM1200F cameras using ACT-1 or ACT-2u software [Nikon]. Whole embryos and explants were mounted in 50% PBS/50% glycerol and photographed with 10x/0.3 Plan Fluor, 20x/0.75 Plan Apo, or 20x/0.5 Plan Fluor objectives. Vibratome sections were mounted in 50% PBS/50% glycerol and photographed with a 40x/0.75 Plan Fluor objective. Fluorescence-labeled sections were mounted in Citiflour mounting medium plus bisbenzimide, and confocal images were taken using a true confocal scanner system (SP5; Leica) and a 40x/1.40 oil UV-corrected objective. Fluorochromes used were bisbenzimide (Invitrogen), Alexa Fluor 488 (Invitrogen), and Cy3 (Jackson ImmunoResearch Laboratories, Inc.). Images were assembled and corrected in contrast and brightness using Photoshop (CS3; Adobe).

Online supplemental material

Fig. S1 shows an analysis of NCC specification and emigration upon FGFR blockage using the PD173074 inhibitor. Fig. S2 shows that VEGFR activity inhibition does not alter Snail2 expression in the neural tube. Fig. S3 shows that blockage of the FGFR signaling pathway promotes premature Pax6 expression and prevents Sprouty2 expression in the neural tube.
Fig. S4 shows that blockade of the FGF signaling pathway promotes premature NCC emigration of several populations. Fig. S5 shows that RA signaling is not required to maintain NCC emigration and does not control the onset of FoxD3 expression in the neural tube. Video 1 shows the migration of NCCs electroporated with H2B:RFP and with control EGFP:GPI at the level of differentiating somites. Videos 2 and 3 show the migration of NCCs electroporated with H2B:RFP and with dn-FGFR1: EFYP at the level of differentiating (Video 2) and epithelial (Video 3) somites. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201011077/DC1.

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References

Ahstrom, J.D., and C.A. Erickson. 2009. The neural crest epithelial-mesenchymal transition in 4D: a ‘tail’ of multiple non-obligatory cellular mechanisms. Development. 136:1801–1812. doi:10.1242/dev.035783

Barrallo-Gimeno, A., J. Holzschuh, W. Driever, and E.W. Knapik. 2004. Neural crest survival and differentiation in zebrafish depends on mont blanc/tfap2a gene function. Development. 131:1463–1477. doi:10.1242/dev.01033

Basch, M.L., M. Bronner-Fraser, and M.I. García-Castro. 2006. Specification of the neural crest occurs during gastrulation and requires Pax7. Nature. 441:218–222. doi:10.1038/nature4684

Bertrand, N., F. Médevielle, and P. Piteau. 2000. FGF signaling controls the timing of Pax9 activation in the neural tube. Development. 127:4837–4843.

Boettger, T., L. Wittler, and M. Kessel. 1999. FGF8 functions in the specification of the right body side of the chick. Curr. Biol. 9:277–280. doi:10.1016/S0960-9822(99)80119-5

Brewster, R., J. Lee, and A. Ruiz i Altaba. 1998. Gli/Zic factors pattern the neural plate by defining domains of cell differentiation. Nature. 393:579–583. doi:10.1038/31242

Bronner-Fraser, M. 1986. Guidance of neural crest migration. Lateral beads as probes of surface-substratum interactions. Dev. Biol. (NY). 3:301–337.

Bursten-Cohen, T., J. Stanleight, D. Sela-Donenfeld, and C. Kalcheim. 2004. Canonical Wnt activity regulates trunk neural crest delamination linking BMP/ noggin signaling with G1/S transition. Development. 131:5322–5339. doi:10.1242/dev.01424

Casanova, J.C., V. Uribe, C. Badia-Careaga, G. Giovinazzo, M. Torres, and J.J. Sanz-Ezquerro. 2011. Apical ectodermal ridge morphogenesis in limb extending vertebrate body axis. Dev. Dyn. 236:136–139. doi:10.1002/dvdy.20919

Diez del Corral, R., and K.G. Storey. 2004. Opposing FGF and retinoid pathways: a signalling switch that controls differentiation and patterning onset in the extending vertebrate body axis. Bioessays. 26:857–869. doi:10.1002/bies.20080

Diez del Corral, R., D.N. Breitkreuz, and K.G. Storey. 2002. Onset of neuronal differentiation is regulated by paraxial mesoderm and requires antagonism of FGF signalling. Development. 129:1681–1691.

Diez del Corral, R., I. Olivera-Martínez, A. Goriely, E. Gale, M. Maden, and K. Storey. 2003. Opposing FGF and retinoid pathways control ventral neural pattern, neuronal differentiation, and segmentation during body axis extension. Neuron. 40:65–79. doi:10.1016/S0896-6273(03)00565-8

Dottori, M., M.K. Gross, P. Labosky, and M. Goulding. 2001. The winged-helix transcription factor FoxD3 suppresses interneuron differentiation and promotes neural crest cell fate. Development. 128:4127–4138.

Dubrulle, B., M.J. McGrew, and O. Pourquié. 2001. FGF signaling controls somite boundary position and regulates segmentation clock control of spatiotemporal Hox gene activation. Cell. 106:219–232. doi:10.1016/S0092-8674(01)00437-8

Echevarría, D., S. Martínez, S. Marques, V. Lucas-Teixeira, and J.A. Belo. 2005. Mkp3 is a negative feedback modulator of Fgf8 signaling in the mammalian isomitic organism. Dev. Biol. 277:114–128. doi:10.1016/j.ydbio.2004.09.011

García-Castro, M.I., C. Marcella, and M. Bronner-Fraser. 2002. Ectodermal Wnt function as a neural crest inducer. Science. 297:848–851.

Goulding, M.D., G. Chalepakis, U. Deutsch, J.R. Erselius, and P. Gruss. 1991. Pax-3, a novel murine DNA binding protein expressed during early neural development. EMBO J. 10:1135–1147.

Harris, M.L., and C.A. Erickson. 2007. Lineage specification in neural crest cell pathfinding. Dev. Dyn. 236:1–19. doi:10.1002/dvdy.20919

Henion, P.D., and J.A. Weston. 1997. Timing and pattern of cell fate restrictions in the neural crest lineage. Development. 124:4351–4359.

Hong, C.S., and J.P. Saint-Jeannet. 2007. The activity of Pax3 and Zic1 regulates three distinct cell fates at the neural plate border. Mol. Biol. Cell. 18:2192–2202. doi:10.1091/mbc.E06-11-1047

Kawakami, A., M. Kimura-Kawakami, T. Nomura, and H. Fujisawa. 1997. Distributions of PAX6 and PAX7 proteins suggest their involvement in three distinct fate fields at the neural plate border. Dev. Dyn. 218:1479–1479.

Krispin, S., E. Nitzan, Y. Kassem, and C. Kalcheim. 2010. Evidence for a dynamic spatiotemporal fate map and early fate restrictions of premigratory avian neural crest. Development. 137:585–595. doi:10.1242/dev.041509

Le Douarin, N.M., and C. Kalcheim. 1999. The Neural Crest. Second edition. Cambridge University Press, Cambridge, UK. 445 pp.

Liem, K.F. Jr., G. Tremml, H. Roefink, and T.M. Jessell. 1995. Dorsal differentiation of neural plate cells induced by BMP-mediated signals from epidermal ectoderm. Cell. 82:969–979. doi:10.1016/0092-8674(95)90276-7

Liu, J.P., and T.M. Jessell. 1998. A role for rhb in the delamination of neural crest cells from the dorsal neural tube. Development. 125:5055–5067.

Loring, J.F., and C.A. Erickson. 1987. Neural crest cell migratory pathways in the trunk of the chick embryo. Dev. Biol. 121:220–236. doi:10.1016/0012-1606(87)90154-0

Lunn, J.S., J.K. Fishwick, P.A. Halley, and K.G. Storey. 2007. A spatial and temporal map of FGF/Erk1/2 activity and response repertoires in the early chick embryo. Dev. Biol. 302:536–552. doi:10.1016/j.ydbio.2006.10.014

Maden, M., E. Gale, I. Kostetskii, and M. Zile. 1996. Vitamin A-deficient quail embryos have half a hindbrain and other neural defects. Curr. Biol. 6:1090–1092. doi:10.1016/0960-9822(96)00126-8

Martinez-Morales, P.L., A.C. Quinrega, J.A. Barbas, and A.V. Morales. 2010. SOX5 controls cell cycle progression in neural progenitors by interfering with the WNT-beta-catenin pathway. EMBO Rep. 11:466–472. doi:10.1038 embr.2010.61

Martins-Green, M., and C.A. Erickson. 1987. Basal lamina is not a barrier to neural crest cell emigration: documentation by TEM and by immuno-fluorescent and immunogold labelling. Development. 101:517–533.
Minowa, G., L.A. Jarvis, C.L. Chi, A. Neubüser, X. Sun, N. Hacohen, M.A. Krasnow, and G.R. Martin. 1999. Vertebrate Sprouty genes are induced by FGF signaling and can cause chordodysplasia when overexpressed. Development. 126:4465–4475.

Mohammadi, M., G. McMahon, A. Monsoro-Burq, R.B. Fletcher, and R.M. Harland. 2003. Neural crest in Xenopus embryos requires FGF signals. Development. 130:3111–3124. doi:10.1242/dev.00551.

Moraes, A.V., J.A. Barbosa, and M.A. Nieto. 2005. How to become neural crest: from segregation to delamination. Semin. Cell Dev. Biol. 16:655–662. doi:10.1016/j.semcdb.2005.06.003.

Nakamura, K., A. Yamamoto, M. Kamishohara, K. Takahashi, E. Taguchi, T. Miura, K. Kubo, M. Shibuya, and T. Isoe. 2004.KRNN633: a selective inhibitor of vascular endothelial growth factor receptor-2 tyrosine kinase that suppresses tumor angiogenesis and growth. Mol. Cancer Ther. 3:1639–1649.

Nakata, K., T. Nagai, J. Aruga, and K. Mikoshiba. 1997. Xenopus Zac3, a primary regulator both in neural and neural crest development. Proc. Natl. Acad. Sci. USA. 94:11980–11985. doi:10.1073/pnas.94.22.11980.

Nieto, M.A., M.G. Sargent, D.G. Wilkinson, and J. Cooke. 1994. Control of cell behavior during vertebrate development by Slug, a zinc finger gene. Science. 264:835–839. doi:10.1126/science.7513443.

Niwa, H., K. Yamamura, and J. Miyazaki. 1991. Efficient selection for high-expression transfectants with a novel eukaryotic vector. Gene. 108:193–199. doi:10.1016/0378-1119(91)90434-D.

Novitch, B.G., H. Wichterle, T.M. Jessell, and S. Sockanathan. 2003. A requirement for retinoic acid-mediated transcriptional activation in ventral neural patterning and motor neuron specification. Neuron. 40:81–95. doi:10.1016/j.neuron.2003.08.006.

Olivera-Martinez, I., and K.G. Storey. 2007. Wnt signals provide a timing mechanism for FGF-induced Etk1/2 signalling is required for vertebrate neural specification. Development. 134:2889–2894. doi:10.1242/dev002858.

Paratore, C., D.E. Goerich, U. Suter, M. Wegner, and L. Sommer. 2001. Survival signalling pathways of neural crest progenitor cells. Development. 128:3949–3961.

Park, K.S., and B.M. Gumbiner. 2010. Cadherin 6B induces BMP signaling and negative chemotaxis mediated by FGF4 and FGF8. Dev. Cell. 3:425–437. doi:10.1016/j.devcel.2004.01.034.

Rickmann, M., J.W. Fawcett, and R.J. Keynes. 1985. The migration of neural crest cells and the growth of motor axons through the rostral half of the chick somite. J. Embryol. Exp. Morphol. 90:437–455.

Rossant, J., R. Zärbigl, D. Cado, M. Shago, and V. Giguère. 1991. Expression of a retinoic acid response element-hoplacZ transgene defines specific domains of transcriptional activity during mouse embryogenesis. Genes Dev. 5:1333–1344. doi:10.1101/gad.5.8.1333.

Saint-Jeannet, J.P., X. He, H.E. Varmus, and I.B. Dawid. 1997. Regulation of dorsal fate in the neuraxis by Wnt-1 and Wnt-3a. Proc. Natl. Acad. Sci. USA. 94:13713–13718. doi:10.1073/pnas.94.25.13713.

Sela-Donenfeld, D., and C. Kalcheim. 1999. Regulation of the onset of neural crest migration by coordinated activity of BMP4 and Noggin in the dorsal neural tube. Development. 126:4749–4762.

Sela-Donenfeld, D., and C. Kalcheim. 2000. Inhibition of noggin expression in the dorsal neural tube by somitogenesis: a mechanism for coordinating the timing of neural crest emigration. Development. 127:4845–4854.

Skaper, S.D., W.J. Kee, L. Facci, G. Macdonald, P. Doherty, and F.S. Walsh. 2000. The FGFR1 inhibitor PD 173074 selectively and potently antagonizes FGF-2 neurotrophic and neurotropic effects. J. Neurochem. 75:1520–1527. doi:10.1046/j.1471-4159.2000.0751520.x.

Stavridis, M.P., J.S. Lunn, B.J. Collins, and K.G. Storey. 2007. A discrete period of FGF-induced Etk1/2 signalling is required for neural crest specification, migration and survival. Dev. Biol. 292:174–188. doi:10.1016/j.ydbio.2005.12.035.

Teillet, M.A., C. Kalcheim, and N.M. Le Douarin. 1987. Formation of the dorsal root ganglia in the avian embryo: segmental origin and migratory behavior of neural crest progenitor cells. Dev. Biol. 120:329–347. doi:10.1016/0012-1606(87)90219-1.

Villanueva, S., A. Glavic, P. Ruiz, and R. Mayor. 2002. Posteriorization by FGF, Wnt, and retinoic acid is required for neural crest induction. Dev. Biol. 241:289–301. doi:10.1006/dbio.2001.0485.

Wilcock, A.C., J.R. Swedlow, and K.G. Storey. 2007. Mitotic spindle orientation distinguishes stem cell and terminal modes of neuron production in the early spinal cord. Development. 134:1943–1954. doi:10.1242/dev002519.

Wilson, L., E. Gale, D. Chambers, and M. Maden. 2004. Retinoic acid and the control of dorsoventral patterning in the avian spinal cord. Dev. Biol. 269:433–446. doi:10.1016/j.ydbio.2004.01.034.

Yang, X., D. Dormann, A.E. Münsterberg, and C.J. Weijer. 2002. Cell movement patterns during gastrulation in the chick are controlled by positive and negative chemotaxis mediated by FGF4 and FGF8. Dev. Cell. 3:425–437. doi:10.1016/S1534-5807(02)00256-3.