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Vascular endothelial growth factor controls neuronal migration and cooperates with Sema3A to pattern distinct compartments of the facial nerve

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Developing neurons accurately position their somata within the neural tube to make contact with appropriate neighbors and project axons to their preferred targets. Taking advantage of a collection of genetically engineered mouse mutants, we now demonstrate that the behavior of somata and axons of the facial nerve is regulated independently by two secreted ligands for the transmembrane receptor neuropilin 1 (Nrp1), the semaphorin Sema3A and the VEGF164 isoform of Vascular Endothelial Growth Factor. Although Sema3A is known to control the guidance of facial nerve axons, we now show that it is not required for the pathfinding of their somata. Vice versa, we find that VEGF164 is not required for axon guidance of facial motor neurons, but is essential for the correct migration of their somata. These observations demonstrate, for the first time, that VEGF contributes to neuronal patterning in vivo, and that different compartments of one cell can be co-ordinately patterned by structurally distinct ligands for a shared receptor.

[Keywords: Neuronal migration, axon guidance, neuropilin, VEGF, semaphorins]

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are less well defined, however, much progress has recently been made through the analysis of hereditary neuropathologies in humans and mice (for review, see Rice and Curran 2001). The perhaps best-studied genetic pathway controlling neuronal migration is centered around Reelin, a secreted protein that binds to the very-low-density lipoprotein receptor (Vldlr) and the apolipoprotein E receptor 2 (ApoER2) to activate the intracellular adaptor disabled (Dab1). The reelin pathway cooperates with a signaling cascade involving Cdk5, a major neuronal kinase implicated in the control of cytoskeletal dynamics. Cdk5 can phosphorylate neurofilaments and several proteins affected in human neurological disorders, most notably doublecortin and the Lissencephaly gene LIS1 (for reviews, see Feng and Walsh 2001; Ohshima and Mikoshiba 2002). LIS1, in turn, can interact with dynein motors, possibly to regulate the transport of the nucleus and other cytoplasmic contents during neuronal migration. Consistent with a role for dynein motors in neuronal migration, we have recently shown that point mutations in cytoplasmic dynein perturb neuronal migration of facial branchiomotor neurons [Hafezparast et al. 2003].

Other pathways implicated in the control of neuronal migration contain the secreted netrin or slit proteins and their receptors. For example, loss of function for netrin impairs the migration of neurons of the inferior olive and the pontine nuclei [Bloch-Gallego et al. 1999; Yee et al. 1999]. In addition to their roles in axon guidance, Nrp1 and Nrp2 have been implicated in the control of neuronal migration. Both proteins are required for the correct pathfinding of GABA-containing interneurons in the telencephalon, and the expression patterns of their semaphorin ligands Sema3A and Sema3F are consistent with a role in guiding this migration [Marin et al. 2001; Tamamaki et al. 2003]. However, it is not known if Nrp1 contributes to both soma and axon guidance of the same neuron. To address this question, we examined the development of facial branchiomotor neurons, as they express Nrp1 to control their axon behavior [Kawakami et al. 1996; Kitsukawa et al. 1997] and undergo an extensive migration from their birthplace to the sites of facial motor nucleus assembly (e.g., see Altman and Bayer 1982; Auclair et al. 1996; Gavalas et al. 2003).

Facial branchiomotor neurons are born in a ventral position in a hindbrain segment termed rhombomere [r] 4. Concomitant with the extension of axons into the second branchial arch, they move their somata caudally to travel through r5 and into r6, where they move to the opposite side of the basal plate. This migration spans several days, covers many cell diameters, and culminates in the formation of the paired motor nuclei of the VIIth cranial (facial) nerve, which controls movement of the facial musculature. The soma migration of facial branchiomotor neurons can be divided into two distinct phases, a tangential and a radial migration (Fig. 1A). The movement from the ventricular side to the pial side of the hindbrain follows the more common pattern of radial migration, where neurons migrate from the ventricular zone toward the outer surface of the brain in close association with radial glia; in contrast, the earlier movement from r4 through r5 and into r6 is a tangential migration, as it occurs perpendicular to the radial glia scaffold (for review, see Marin and Rubenstein 2003).

We found that loss of Nrp1 compromised the soma migration of facial branchiomotor neurons during the tangential phase and resulted in the formation of misshapen and malpositioned facial motor nuclei. Two semaphorins with a high affinity for Nrp1, Sema3A and Sema3C, were expressed in the hindbrain in a spatiotemporal pattern consistent with a role in the migration of facial branchiomotor somata. However, to our surprise, neither Nrp1 ligand was essential for this process, nor was the ability of Nrp1 to bind semaphorins. Rather, soma migration of facial branchiomotor neurons relied on the presence of a structurally unrelated Nrp1 ligand that is better known for its effect on the growth, differentiation, and patterning of blood vessels, an isoform of Vascular Endothelial Growth Factor (VEGF) termed VEGF164 [Soker et al. 1998; for review, see Ruhrberg 2003].

Our observations suggest that the dual affinity of Nrp1 for VEGF and semaphorin ligands allows facial branchiomotor neurons to coordinate the migratory behavior of their axons and somata: Whereas Sema3A is required for their axon guidance [Taniguchi et al. 1997], VEGF164 supports the correct positioning of their somata within the neural tube. These findings establish, for the first time, that VEGF is required to control the complex behavior of differentiated neurons in vivo, and they have widespread implications for our understanding of signaling pathways implicated in human neurological disorders.

Results

Nrp1 controls the migration of facial branchiomotor neurons

Using a previously characterized probe for in situ hybridization [He and Tessier-Lavigne 1997], we confirmed that Nrp1 was expressed by facial branchiomotor neurons; moreover, we found that it was expressed by these neurons both during the time of soma migration and after the soma had condensed into nuclei [Supplementary Fig. S1]. To examine if Nrp1 signaling was required for the pathfinding of facial branchiomotor somata, we followed their journey in Nrp1-deficient mouse embryos with a marker for postmitotic motor neurons, an in situ probe for the LIM homeodomain protein Isl1 [Ericson et al. 1992].

We found that loss of Nrp1 did not impair the generation of facial branchiomotor neurons in r4, but that it disrupted the tangential migration of their somata in 13/13 cases examined (Fig. 1B–K). More specifically, loss of Nrp1 delayed soma migration and resulted in the separation of the migratory stream into several distinct streams on the ventricular side of the hindbrain [Fig. 1C,G, arrowheads]. Consequently, many somata initi-
ated their radial migration toward the pial side of the hindbrain in ectopic anterior locations (Fig. 1E,II) and contributed to the formation of elongated or dumbbell-shaped nuclei, rather than the characteristic rounded nuclei seen in wild-type littermates [Fig. 1, cf. H and I, J and K; data not shown]. In severe cases, the entire nucleus was shifted anteriorly (Fig. 1, cf. the relative position of facial and trigeminal nuclei in D and E).

The abnormal soma migration of facial branchiomotor neurons is not explained by vascular defects in the hindbrain or axon defects in the periphery

Loss of Nrp1 alters the migratory behavior of facial nerve axons, as they defasciculate once they reach their target area, the second branchial arch (Kitsukawa et al. 1997). Because facial branchiomotor neurons translocate their somata at a time when their axons navigate in the periphery, we asked if the abnormal soma behavior was secondary to the disturbance of axonal patterning. To answer this question, we cultured 11.5-dpc (days post-coitum) hindbrains that had been axotomized and were free of branchial arch tissue under serum-free conditions. We found that removing distal axons or peripheral targets did not compromise soma migration in any of 15 wild-type hindbrain explants (Fig. 2A–D). In contrast, after 2 d in culture, 14/14 explants lacking Nrp1 contained facial branchiomotor somata in an anterior position (cf. Fig. 2I–L). Taken together, the observations from explant culture experiments suggest that soma migration in the neural tube and axon guidance in the periphery are regulated independently.

The radial glia scaffold assists the radial migration of neurons in the central nervous system, but it is not known if specific structural features support the tangential migration of neurons. Because mutants lacking Nrp1 are unable to form a subventricular vascular plexus [Kawasaki et al. 1999; Q. Schwarz and C. Ruhrberg, unpubl.], and the tangential migration of facial branchiomotor somata occurs in close proximity to the subventricular vascular network [see Fig. 2E; Q. Schwarz and C. Ruhrberg, unpubl.], we asked if their abnormal migration was caused indirectly by the vascular defect. This was not the case, as the soma migration of facial branchiomotor neurons proceeded normally in hindbrain explant cultures, in which vessel networks rapidly degenerated (Fig. 2E–H). Moreover, the somata of facial branchiomotor neurons migrated normally in 2/2 hindbrains of mutants that specifically lack Nrp1 in vessel endothelium (Fig. 2M,N) and that have vascular defects similar to the full Nrp1 knockout (Gu et al. 2003). We therefore conclude that normal vessel networks are not essential for the guidance of migrating facial branchiomotor somata.

Figure 1. Nrp1 is required for the correct pathfinding of facial branchiomotor somata. (A) Schematic representation of the spatial relationship of trigeminal, facial, abducens, and glossopharyngeal motor neurons in a flat-mounted mouse hindbrain. The axon exit points for trigeminal and facial branchiomotor neurons [hatched circles] and the floorplate (gray) are indicated. The cross-sections on the right-hand side show the position of facial branchiomotor somata on the ventricular (v) side in r4 (top section) and on the pial (p) side in r6 [bottom section]. (B,C) The migration of ISL1-positive facial branchiomotor somata (VIIm) from r4 to r6 was observed on the ventricular side in the presence (B) or absence (C) of Nrp1 at 12.5 dpc. Wild-type somata stayed loosely associated to form one continuous stream, but mutant somata separated into several distinct streams (arrows and arrowheads indicate separation in r4 and r5/r6, respectively). (D) In the presence of Nrp1, many facial somata have reached the pial side of the hindbrain by 12.5 dpc and formed facial motor nuclei (VIIm). (E) In the absence of Nrp1, many mutant somata emerged on the pial side in an ectopic anterior location [star] instead of their normal location [A]. Panel D shows one side of a wild-type hindbrain, and panel E shows the opposite side of a stage-matched mutant hindbrain to highlight differences in the position of facial branchiomotor neurons relative to the trigeminal nucleus [Vn]. (F) Few facial branchiomotor somata were normally present on the ventricular side at 13.5 dpc [bracket]. (G) In contrast, many mutant somata were still migrating on the ventricular side at 13.5 dpc (arrowheads). (H) On the pial side, some mutant somata had contributed to normally positioned nuclei (VIIm), while others had formed an ectopic nucleus (star). One side of a wild-type hindbrain (F,G) and the opposite side of a stage-matched mutant hindbrain (H,I) are shown. (J,K) By 14.5 dpc, all facial branchiomotor somata had integrated into nuclei on the pial side, which in wild types appeared round [J], but in mutant littermates were usually dumbbell-shaped, with an ectopic anterior component [star; K]. Bars, 250 µm (one for B,C; one for D,E; one for F–I; and one for J,K). The midline is indicated with an asterisk.
VEGF controls neuronal migration

As the abnormal soma migration of facial branchiomotor neurons in the absence of Nrp1 was not likely secondary to axon defects or disrupted vessel networks, we asked if it instead was caused by an inability of the somata to respond to a secreted Nrp1 ligand. Nrp1 binds with high affinity to several class 3 semaphorins, but also to an isoform of VEGF, termed VEGF165 in humans and VEGF164 in the mouse (Fig. 2O; He and Tessier-Lavigne 1997; Kolodkin et al. 1997; Soker et al. 1998).

Sema3A and Sema3C expression patterns are consistent with a role in the development of facial branchiomotor neurons

Nrp1 serves as a high affinity receptor for several class 3 semaphorins, most notably Sema3A and Sema3C, and facial motor axons respond to Sema3A in culture models (Varela-Echavarria et al. 1997; for review, see Raper 2000). We found that Sema3A and Sema3C, but not Sema3B or Sema3E, were expressed strongly in the hindbrain (Fig. 3; data not shown). Whereas Sema3A was expressed more strongly in the environment (Fig. 3C, arrow), Sema3C was present at higher levels in the migrating neurons (Fig. 3E, arrowhead). At 12.5 dpc, Sema3A expression extended in a stripe throughout the length of the hindbrain, which coincided with the position where facial branchiomotor somata begin their descent through the basal plate (Fig. 3C, arrow). This observation raises the possibility that Sema3A provides a repulsive boundary that forces these neurons onto a new course. Interestingly, Sema3C expression was down-regulated once the facial branchiomotor somata had reached the pial surface (Fig. 3F, Δ), suggesting that Sema3C promoted their migration, but did not play a role in motor nucleus assembly.

Sema3A and Sema3C are not essential to guide facial motor neuron migration

Although Sema3A was expressed in a pattern consistent with a role in neuronal guidance in the hindbrain, it was not required for the pathfinding of facial branchiomotor somata. Isl1-staining showed that the somata migrated normally on the ventricular side and arrived on the pial side in their proper location and at the appropriate time in the absence of Sema3A in all four cases examined (Fig. 3, cf. G and H; data not shown). To address whether Sema3C was required for some migration, we examined four mutants lacking this protein (Feiner et al. 2001), but found that Sema3C was also dispensable for the pathfinding of facial branchiomotor somata to their destination on the pial side of the hindbrain (Fig. 3, cf. G and I).

The lack of a phenotype in Sema3A or Sema3C knockouts may be explained by their functional redundancy during soma migration. To address this possibility, we examined compound mutants lacking both Sema3A and Sema3C and found that the soma migration of facial branchiomotor neurons occurred normally in all four
Sema3C (of asterisk. were located within their nuclei (VIIn) on the pial side (cf. the relative positions of VIIn and Vn). Bars: and in neighboring regions (arrows in Sema3A to condense into the paired facial motor nuclei (VIIn) on the pial side in the area where facial somata migrate (arrowheads in Sema3A knockouts or Nrp1 was not required for this process. Moreover, the suggest that signaling of class 3 semaphorins through occurred normally in all five cases examined (Fig. 4G–L), as mentioned above.

Figure 3. Sema3A and Sema3C are expressed in the hindbrain, but are not required for the pathfinding of facial branchiomotor somata. (A,B) In situs hybridization (ISH) shows that at 12.5 dpc, the Isl1-positive somata of facial branchiomotor neurons were migrating caudally (VIIm) on the ventricular side [A] and began to condense into the paired facial motor nuclei (VIln) on the pial side [B] of the hindbrain. (C–F) In stage-matched hindbrains, Sema3A [C] and Sema3C [E] were expressed on the ventricular side in the area where facial somata migrate (arrowheads in C,E) and in neighboring regions (arrows in C,E). Sema3A [D] and Sema3C [F] were down-regulated on the pial side (cf. positions of Δ in D,F and VIln in B). By 13.5 dpc, most facial somata were located within their nuclei (VIln) on the pial side (cf. the size of facial motor nuclei in B and C). Loss of Sema3A [H] or Sema3C [I] did not impair the formation or positioning of facial motor nuclei (cf. the relative positions of VIln and Vn). Bars: A–F, 250 µm; G–I, 200 µm. The midline is indicated with an asterisk.

Between 10.5 and 12.5 dpc, VEGF is expressed more abundantly on the ventricular relative to the pial side of the hindbrain to guide angiogenic sprouting from the perineural plexus toward the subventricular zone (Breier et al. 1992; Miquerol et al. 1999). To relate the expression pattern of VEGF to the development of facial branchiomotor neurons, we performed whole-mount in situ hybridization at 12.5 dpc and found that the expression of the VEGF gene [VEGF-A] was up-regulated in the paired regions where facial motor nuclei formed (Fig. 5, cf. A and B). However, VEGF mRNA is unstable under normoxic conditions (e.g., Levy et al. 1995), and for this reason, its expression pattern is difficult to follow with the whole-mount in situ hybridization technique. To increase sensitivity and spatial resolution, we therefore monitored VEGF expression in subsequent experiments with a previously characterized β-galactosidase reporter, which carries a LacZ transgene in the 3′-UTR of VEGF-A (Miquerol et al. 1999).

As described before, we found that the LacZ reporter for VEGF-A was initially expressed throughout the ventricular aspect of the hindbrain, but became down-regulated by 11.5 dpc, correlating with the formation of an extensive vascular plexus (Ruhrberg et al. 2002). However, expression in the midline area remained high at 12.5 and 13.5 dpc (Fig. 5D, arrow; data not shown). In addition, VEGF-A expression at 12.5 dpc was elevated in paired dorso-medial stripes that extended throughout the length of the hindbrain (these paired stripes appear laterally in flat-mounted hindbrain preparations and are indicated by a curved arrow in Fig. 5D). The position of both these domains is consistent with a role as stepping stones for the somata along their journey: The somata first travel caudally in close proximity to the VEGF-rich midline and then change direction to travel toward the dorso-medial VEGF domains. However, neither the ventral nor the dorso-medial stripes of VEGF are restricted to the rhombomeres that contain the migrating somata of facial branchiomotor neurons. We therefore have to postulate the existence of cooperating cues, which prevent entry into anterior territories and help the migrating somata to change course once they arrive in r6 territory, where they dive through the basal plate toward the site of facial nucleus assembly (see above). Importantly, the LacZ reporter confirmed that VEGF-A was expressed at 12.5 dpc in the region where the facial nuclei formed (Fig. 5C,E), consistent with a role for VEGF in guiding the movement of the somata through the basal plate. At 13.5 dpc, VEGF-A expression on the pial side resolved into a domain corresponding to the facial nucleus assembly site and an adjacent stripe (Fig. 5, cf. E and F).

VEGF is made in several isoforms, but VEGF164 is the only Nrpl-binding isoform expressed in the hindbrain during the time of facial branchiomotor neuron migration: VEGF188 is not expressed at significant levels at 12.5 dpc (Ruhrberg et al. 2002), and VEGF120 (Soker et
Ventricular side (cf. A and D). At 12.5 dpc (cf. B and E) or
13.5 dpc (cf. C and F), [G–L] Mutation of the
Sema-binding domain of Nrp1 did not impair pathfinding of
Isl1-positive facial somata on the ventricular side (cf. G and F),
or the positioning of facial motor nuclei on the pial side at 12.5
dpc (cf. H and K) or 13.5 dpc (cf. I and L). Bars, 250 µm (one for
A, B, D, E, one for G, H, J, K), 200 µm (one for C, F, one for I, L). The
midline is indicated with an asterisk.

Figure 4. Semaphorin signaling through Nrp1 is not required
for the migration of facial branchiomotor somata. [A–F] Loss of
both Sema3A and Sema3C did not impair the pathfinding of
Isl1-positive facial somata on the ventricular side (cf. A and D),
nor the positioning of facial motor nuclei on the pial side at 12.5
Sema-binding domain of Nrp1 did not impair pathfinding of
Isl1-positive facial somata on the ventricular side (cf. G and F),
or the positioning of facial motor nuclei on the pial side at 12.5
dpc (cf. H and K) or 13.5 dpc (cf. I and L). Bars, 250 µm (one for
A, B, D, E, one for G, H, J, K), 200 µm (one for C, F, one for I, L). The
midline is indicated with an asterisk.

Defective soma migration in the absence of Nrp1 and VEGF164 is not caused
by abnormal hindbrain segmentation

In several mouse mutants, the irregular soma migration of
facial branchiomotor neurons can be attributed to de-
fects in hindbrain segmentation (for review, see Cordes
2001). To examine if migration defects in Nrp1−/− and
VEGF164+/− mutants were an indirect consequence of an
abnormal hindbrain structure, we used a marker that
detects trigeminal and facial branchiomotor neurons as
well as segment-specific neurogenesis patterns, the
paired homeobox gene Phox2b (Davenne et al. 1999; Pat-
tyn et al. 2000; Gavalas et al. 2003). At 12.5 dpc, Phox2b
is expressed in the wild-type hindbrain in both migrating
facial branchiomotor somata and in three neuron col-
umns that reflect the segmental organization of the
hindbrain [Supplementary Fig. S2A, B, J, Gavalas et al.
2003]. Using this marker, we confirmed the abnormal
soma migration of facial branchiomotor neurons in 4/4
Nrp1−/− and 2/2 VEGF164+/− mutants, but found that
other Phox2b-positive neurons were organized into their
appropriate columns [Supplementary Fig. S2C–F]. These
observations suggest that the abnormal soma migration
of facial branchiomotor neurons in the absence of Nrp1 or
VEGF164 is not due to a defect in hindbrain segmentation.

Defective soma migration of facial branchiomotor
neurons in the absence of VEGF164 is not
accompanied by axon guidance defects
in the periphery

Loss of Nrp1 or Sema3A results in the defasciculation
of several cranial nerves, including the trigeminal and fa-
al. 2002] showed no defects in the pathfinding of migrat-
ing facial somata and therefore assembled normal facial
motor nuclei (Fig. 5O, P; data not shown). We therefore
conclude that VEGF164 is the VEGF isoform that controls
the soma migration of facial branchiomotor neurons.

To examine if VEGF164 controlled the migration of
facial branchiomotor somata in the absence of functional
axons, we added neutralizing antibodies specific for
VEGF to the culture medium of hindbrain explants at
11.25 dpc (Fig. 5Q–S). We found that treatment with
VEGF-neutralizing antibodies, but not treatment with
control antibodies, caused the somata to prematurely
turn dorsally in r4 or r5 in 5/6 wild-type explants and in
5/5 explants expressing VEGF164 only (Fig. 5R, S; data
not shown).

Because VEGF is a chemoattractant for various cell
types, including neuronal precursors [Zhang et al. 2003],
it may also act as an attractant for migrating somata
when presented in a localized fashion. To address this
possibility, we implanted VEGF164-coated heparin
beads into normal hindbrain tissue, and found that they
promoted the migration of facial somata toward the
VEGF164 source in 5/6 cases [Fig. 5U]. In contrast, control
beads without VEGF did not attract the migrating
somata [4/4 cases] [Fig. 5T].
Figure 5. VEGF controls the pathfinding of facial branchiomotor somata. [A,B] Whole-mount in situ hybridization at 12.5 dpc shows expression of the VEGF-A gene (B) in the area where the Isl1-positive somata of facial branchiomotor neurons assemble into facial motor nuclei (VIIln) on the pial side of the hindbrain [A]. (C–F) Expression of a VEGF-A LacZ reporter at 12.5 (C–E) and 13.5 dpc (F). (F) At 12.5 dpc on the pial side, VEGF-A LacZ expression was prominent in the area of facial motor nucleus assembly [VIIln]. [D] At 12.5 dpc on the ventricular side, VEGF-A LacZ expression was prominent near the midline (arrow), in a more dorsally located path of stripes (wavy arrow), and in the area where the hypoglossal nuclei form (XIIln). (E) A higher magnification of the boxed area in C. (F) At 13.5 dpc, expression was elevated in the area of facial motor nucleus assembly and in an adjacent stripe (open arrowhead). (G–N) The migration of the Isl1-positive somata of facial branchiomotor neurons [arrowheads] was observed in the presence [wt/wt or wt/120] or absence [120/120] of VEGF164. In the absence of VEGF164, facial branchiomotor neurons formed nuclei in anterior positions, which appeared elongated or dumbbell shaped [stars, L, N]. One side of a control hindbrain and the opposite side of a stage-matched mutant hindbrain are shown next to each in G–L to highlight the position of the facial branchiomotor neurons relative to the hypoglossal [XIIln, G, H] or trigeminal [Vn, I–L] nuclei (cf. the length of the square brackets in G–I controls and mutants). In some mutants, somata emerged in an abnormally large area on the pial side and could be identified as scattered Isl1-positive cells (cf. boxed areas in K and L). (O, P) A comparison of the facial motor nuclei [VIIln] formed in littermate hindbrains capable of expressing all VEGF isoforms [wt/wt, O] or VEGF164 only [164/164, P] showed that VEGF164 was sufficient to drive normal soma migration. (Q–U) Hindbrain tissue (11.25 dpc) expressing VEGF164 only was fixed [0 h, Q] or cultured for 2 d in the presence of function-blocking antibodies for VEGF (αVEGF, R, S) to ablate VEGF164 function. When VEGF164 function was blocked, somata continued to migrate caudally out of r4 on the ventricular side, and a proportion of somata dived through the basal plate in the expected position [open arrowhead, R] to form facial motor nuclei on the pial side [VIIln, S]. However, ectopically migrating somata were seen on the ventricular side in the r5 territory [arrowheads in R]. (T, U) When heparin beads (b) were implanted into normal hindbrain tissue, somata did not respond to control beads (T), but moved toward beads coated with VEGF164 [U]; accordingly, somata were positioned more posteriorly on the side containing VEGF beads relative to the untreated hindbrain side [double bracket in U]. Bars, 250 μm (one for A–D; one for E, F one for G–I; one for K, L; on for M–P; one for Q–U). The midline is indicated with an asterisk; in K and L the midline is located outside the panel, but its position is indicated with an open arrowhead.

Discussion

VEGF164 and Nrp1 guide soma migration in the hindbrain

To address if Nrp1 controlled soma and axon migration of the same neurons, we focused on facial branchiomotor neurons; firstly, because their axons require Nrp1 for their guidance, and secondly, because their somata undergo an extensive migration through several hindbrain segments that is readily monitored (Fig. 1A; Supplementary Fig. S1). Using two markers for developing facial branchiomotor neurons, Isl1 and Phox2b, we found that loss of Nrp1 impaired soma pathfinding during their tangential, but not their radial migration phase, and therefore caused the formation of misshapen or ectopic facial [VIIth] motor nuclei (Fig. 1; Supplementary Fig. S2).
Figure 6. VEGF164 is not required for the target finding of facial nerve axons. (A–D) Neurofilament staining of cranial nerve axons in 11.5-dpc embryos. Facial nerve axons extend normally into the branchial arches in the presence [A, C] or absence [D] of VEGF164, but they defasciculate in mutants lacking Nrp1 (B). The three major branches of the lower facial nerve (VII) are indicated with arrows. The position of the maxillary (Vmx), mandibular (Vmd), and ophthalmic (Vop) branches of the trigeminal nerve, the vestibulocochlear nerve (VIII) and the trigeminal (Vg) and facial (VIIg) ganglia are indicated in A. In B, defasciculation of the facial nerve is highlighted with a circle, and an abnormally positioned branch of the facial nerve is indicated with an open arrowhead. The trigeminal nerve branches are also defasciculated. Bar, 250 µm. [E] Working model for the control of axon and soma guidance in facial branchiomotor neurons by Nrp1 ligands. Sema3A binds to Nrp1-containing receptors in the growth cone to control axon guidance in the periphery, while VEGF164 binds to Nrp1-containing receptors on the cell body in the hindbrain to control soma migration. The most prominent site of VEGF164 expression on the pial hindbrain side is shown in green to indicate its likely role as an attractive signal for migrating somata. Sema3A is shown in red to indicate its role as an axon repellent in the periphery. Nrp1 associates with plexins to transmit semaphorin signals in the growth cone, but the signaling coreceptor in the soma is not known.

As loss of Nrp1 affects axon pathfinding in the periphery and compromises vascularization of the neural tube, we first examined if the defective soma migration of facial branchiomotor neurons was secondary to axonal or vascular patterning defects. However, there was no relationship between the control of soma and axon pathfinding, and proper vessel patterning was not a prerequisite for normal soma migration [Figs. 2–4]. These observations suggested that Nrp1 was required cell-autonomously to mediate the response of migrating somata to a secreted Nrp1 ligand.

To determine which Nrp1 ligands control the behavior of facial branchiomotor somata, we analyzed the expression patterns and functional requirements for class 3 semaphorins and VEGF. Based on their expression patterns and functions in vitro, Sema3A and Sema3F have both been implicated in the Nrp1/Nrp2-mediated control of interneuron migration in the telencephalon (Marin et al. 2001; Tamamaki et al. 2003). The observation that neither Sema3A nor Sema3C, nor any other Nrp1-binding semaphorin, was required for the Nrp1-mediated pathfinding of facial branchiomotor neurons was therefore unexpected [Figs. 3, 4]. Even more surprising was the finding that soma behavior of these neurons instead relied on the alternative Nrp1 ligand VEGF164, which is better known for its role in supporting the migration of cultured endothelial cells (Soker et al. 1998).

The lack of a soma migration defect in mutants lacking semaphorin signaling though Nrp1 may be due to compensation by Nrp2-based semaphorin signaling, as Nrp2 is expressed in migrating facial somata [Q. Schwarz and C. Ruhrberg, unpubl.]. However, we believe that this possibility is unlikely, because loss of Nrp2 or its ligands Sema3C or Sema3F did not obviously impair the soma migration of facial branchiomotor neurons [Fig. 3; Q. Schwarz, V. Vivancos, M. Studer, S. Guthrie, C. Gu, A. Sahay, A. Kolodkin, D. Ginzy, and C. Ruhrberg, unpubl.]. We therefore conclude that semaphorins at best provide a backup system for VEGF164 signals. Such a function may only be revealed in compound mutants, for example, those carrying Sema3A knockout alleles on a VEGF120 background. Alternatively, the striking expression patterns of Sema3A and Sema3C in the hindbrain may not indicate a role in soma guidance, but instead reflect other functions during hindbrain patterning. For example, Sema3A may provide repulsive boundaries for axon guidance within the hindbrain, and Sema3C may be expressed by migrating neurons to repel Nrp1/Nrp2-bearing vascular endothelial cells in the neuronal migration route.

Whereas semaphorin signaling was dispensable, VEGF164 was necessary both in vitro and in vivo for the correct pathfinding of facial branchiomotor somata [cf. Figs. 4 and 5]. The soma migration defect was fully penetrant in mutants lacking VEGF164, and they displayed a similar spectrum of pathfinding errors to Nrp1 mutants, in agreement with biochemical data that demonstrate the ability of VEGF164 to bind to Nrp1 (Soker et al. 1998). The combined analysis of hindbrains lacking VEGF164 [VEGF<sup>120/120</sup>] or expressing VEGF164 at the expense of all other isoforms [VEGF<sup>164/164</sup>] shows that VEGF164 is the only VEGF isoform that is both necessary and sufficient for the soma migration of facial branchiomotor neurons in vivo [Fig. 5]. As VEGF188 is not expressed at significant levels in the hindbrain during the time of facial branchiomotor neuron migration (Ruhrberg et al. 2002), it cannot normally contribute to facial soma pathfinding. However, VEGF188 does contain the domain that mediates Nrp1 binding in VEGF164, and it will therefore be interesting to determine if the ectopic expression of VEGF188 in the developing hindbrain can rescue the soma migration defects caused by a lack of VEGF164.
The ability of VEGF164-coated heparin beads to attract migrating facial branchiomotor somata in explant culture suggests that VEGF is a chemoattractant for Nrp1-expressing neurons [Fig. 5]. In this assay, VEGF is presented in a localized form in conjunction with heparin, an important cofactor for Nrp1 binding [Gluzman-Poltorak et al. 2000]. In contrast to heparin-rich VEGF beads, the addition of VEGF164 to the culture medium did not perturb soma migration (data not shown). This may be explained by a failure of exogenously added VEGF to be absorbed into the heparin sulfate proteoglycan (HSPG)-rich hindbrain matrix; for example, VEGF-binding sites may not be accessible or already saturated. Alternatively, VEGF may only act as a chemoattractant for migrating neurons when presented in a steep gradient, as observed in the case of migrating endothelial cells in the embryonic hindbrain and retina [Ruhrberg et al. 2002; Gerhardt et al. 2003].

Despite the striking similarity in pathfinding errors, there were some differences in the migratory behavior of facial branchiomotor somata in Nrp1−/− and VEGF120/120 mutants. For example, facial somata split into several distinct streams in Nrp1−/− mutants, but they spread out more in VEGF120/120 mutants [cf. Figs. 1 and 5]. This difference may relate to Nrp1’s ability to interact with partners other than VEGF164, for example, the cell adhesion molecule L1 [Castellani and Rougon 2002] or the Nrp2 receptor [Gluzman-Poltorak et al. 2000]. A second reason for the differences between the soma migration defects of Nrp1−/− and VEGF120/120 mutants may relate to VEGF164’s ability to participate both in isoformspecific signaling and in the formation of matrix-associated VEGF concentration gradients. VEGF120/120 mutants produce excess amounts of VEGF120 at the expense of VEGF164, and this increase in soluble heparin-binding isoforms promotes the dispersion of VEGF in the hindbrain, and therefore in the environment of migrating somata [Ruhrberg et al. 2002]. As developing motor neurons express Flt1 [Q. Schwarz, M. Golding, and C. Ruhrberg, unpubl.], they may have the capacity to detect ectopic VEGF deposits composed solely of VEGF120. A third possibility for the differences between the soma migration defects of Nrp1−/− and VEGF120/120 mutants may relate to the observation that VEGF164 and VEGF144 can both bind to Nrp2 [Gluzman-Poltorak et al. 2000]. Because Nrp2 is expressed by the migrating facial somata [see above], it is conceivable that VEGF144/164 signaling through Nrp2 could partially compensate for loss of VEGF164 signaling in Nrp1−/− mutants. In contrast, VEGF120/120 mutants lack both VEGF144 and VEGF164, and therefore loss of VEGF164 signaling through Nrp1 could not be rescued by VEGF144/164 signaling through Nrp2. As full Nrp1/Nrp2 double knockouts are lethal early during embryonic development [Takashima et al. 2002], it would be necessary to generate conditional double knockouts to test the hypothesis that Nrp2 can compensate for Nrp1 signaling in facial branchiomotor neurons.

Independent control of soma migration and axon guidance through Nrp1 ligands: cooperation, not competition of VEGF164 and Sema3A

It has previously been suggested that axon guidance can occur independently of signals from the soma, because axons of retinal ganglion cells continue to navigate after removal of their somata [Harris et al. 1987]. We now show that the reverse is also true, as soma pathfinding of facial branchiomotor neurons continues normally after axons have been removed in explant culture [Fig. 2]. The analysis of several different mouse mutants also supports the idea that soma migration and axon guidance of the developing facial nerve are regulated independently. Firstly, loss of HoxB1 or the zebrafish trilobite mutation, like loss of VEGF164, disrupt the soma migration of facial branchiomotor neurons without affecting the target finding of facial nerve axons [Fig. 6; Studer et al. 1996; Bingham et al. 2002; Gavalas et al. 2003]. Secondly, loss of Sema3A disturbs the behavior of facial nerve axons [Taniguchi et al. 1997], but does not perturb the soma migration of facial branchiomotor neurons [Fig. 3].

Because VEGF164 controls the soma positioning of facial branchiomotor neurons, but is not required for their axon guidance, and, conversely, Sema3A is essential for their axon guidance, but not the pathfinding of their somata, we conclude that Nrp1 supports soma and axon migration independently through the binding of two different ligands [Fig. 6E, model]. Therefore, Sema3A and VEGF do not appear to compete with each other, but instead cooperate by patterning different compartments of the same cell.

The concept that Nrp1 ligands cooperate to pattern neurons is novel and differs from an earlier model of Nrp1 function in neurons, which suggests that the competition of Sema3A and VEGF164 for binding to Nrp1 on the axonal growth cone is critical to steer migrating axons [Carmeliet 2003]. The competitive model for neuronal patterning by Nrp1 ligands was derived from work in several cell culture systems, where Sema3A modulates the activity of VEGF on vascular endothelial cells [Miao et al. 1999], while VEGF counters Sema3A-mediates responses of neural progenitor cells [Bagnard et al. 2001]. Both models may not be mutually exclusive, but could operate in distinct situations. For example, the balance of VEGF164 and Sema3A may influence the decision of axons to grow in tight fascicles or defasciculate, or influence the patterning of growing blood vessel networks.

The ability of Nrp1 to participate in two separate patterning events of the same neuron through its alternative ligands raises the possibility that it exists in different conformational states [1] to influence Nrp1’s preference for its structurally distinct ligands VEGF164 and Sema3A, and [2] to stimulate distinct intracellular signaling events in the soma and the axon. For example, Nrp1 may associate with a different signaling coreceptor in soma and axon, where Flk1 or Flt1 may mediate a VEGF response, while plexins would transduce a semaphorin signal. We were not able to detect Flk1 in develop-
opposing motor neurons between 10.5 and 14.5 dpc, consistent with the previous observation that Flk1 expression in neurons is not obviously required for brain development [Haigh et al. 2003]. In contrast, we found expression of Flt1 and three A-type plexins in facial branchiomotor neurons by in situ hybridization (Q. Schwarz and C. Ruhrberg, unpubl.). One attractive hypothesis may therefore be that VEGF recruits Flt1 to Nrp1 to mediate an attractive response to VEGF in the soma, whereas Sema3A/Nrp1 recruits a plexin family member to induce repulsion of axons. Alternatively, VEGF164 and Sema3A may recruit different plexin family members to Nrp1. The hypothesis that Nrp1 requires different coreceptors to signal VEGF164 and Sema3A responses might be tested through a comparative analysis of soma and axon behavior in mouse mutants carrying conditional Flk1 or Flt1 knockout alleles or plexin loss-of-function mutations.

**Relationship of VEGF 164/Nrp1 to other proteins implicated in neuronal migration**

In several mouse mutants, defective facial nerve development can be attributed to errors in the segmental patterning of the hindbrain [for review, see Cordes 2001]. In some of these mutants, the somata of facial branchiomotor neurons encounter an abnormal environment during their migration, because transcription factors that control hindbrain segmentation are lost (e.g., kreisler). In another type of hindbrain patterning mutant, facial branchiomotor neurons undergo a switch in molecular identity together with their r4 environment, and their somata therefore assume a migratory path characteristic of trigeminal branchiomotor neurons [HoxB1 knockouts]. However, the soma migration defects in Nrp1−/− and VEGF164−/− mutants are not caused by errors in the segmental patterning of the hindbrain [Supplementary Fig. S2]. In this respect, both mouse mutants resemble the zebrafish mutant trilobite, in which soma migration, but not hindbrain patterning or neuronal differentiation, is affected [Bingham et al. 2002]. In these mutants without segmentation defects, abnormal neuronal migration is likely due to cell-autonomous defects, in which neurons fail to respond to environmental cues.

We have been able to demonstrate a role for VEGF164 as a guidance cue for Nrp1-expressing motor neurons using a combination of mouse genetics and explant cultures [Fig. 5]. Moreover, we have preliminary evidence to suggest that VEGF164 and Nrp1 also contribute to the guidance of migrating inferior olive neurons (Q. Schwarz and C. Ruhrberg, unpubl.). However, just as Sema3A is only one of a large collection of axon guidance cues that act synergistically to control the pathfinding of peripheral nerves [for review, see Dickson 2002], VEGF164 is likely to act in concert with other guidance cues. In agreement with this idea, the secreted proteins reelin and netrin also play essential roles during the migration of facial branchiomotor and inferior olive neurons, but loss of either protein does not abolish soma pathfinding, as observed for loss of VEGF164 [Bloch-Gallego et al. 1999; Ohshima et al. 2002]. We therefore anticipate that VEGF synergises with other guidance pathways to control the migration of several different neuron classes in the brain.

Similar to the loss of an environmental guidance cue, the mutation of genes that control cellular responses to such extracellular stimuli can cause neuronal migration defects. For example, the mutation of the intracellular proteins Dab1 or Cdk5 or the Cdk5-associated regulatory subunits p35 and p38 disrupts the soma migration of facial branchiomotor neurons [Ohshima et al. 2002]. The VEGF/Nrp1 complex may operate in the same genetic pathway that contains Cdk5 and cytoplasmic dynein, because (1) Nrp1-containing receptor complexes can recruit Cdk5 [Sasaki et al. 2002], (2) Cdk5 can phosphorylate proteins that interact with dynein and have been implicated in nuclear translocation, and (3) loss of VEGF164/Nrp1 signaling and a dynein point mutation cause similar defects during facial branchiomotor soma migration [cf. Figs. 1, 5 shown here and Fig. 3 in Hafezparast et al. 2003]. We therefore speculate that binding of VEGF164 to Nrp1 activates a signaling pathway, which modifies Cdk5 activity to control microtubule dynamics and thereby promotes nuclear and organelle translocation during soma migration.

**Potential roles for VEGF/Nrp1 signaling in neuronal diseases**

Because VEGF164 and Nrp1 play essential roles during neuronal migration in the central nervous system, defective VEGF signaling might contribute to human disease. Moebius syndrome [MBS, OMIM 157900] is an inborn human condition that affects the VIIth cranial nerve, and three chromosomal loci have been associated with familial Moebius syndrome. MBS1 maps to chromosome 13q12.2–q13, which harbors the Flt1 gene (OMIM 165070); MBS2 maps to 3q21–q22, which contains PlexinA1 [OMIM 601055] and two other genes whose products have the potential to interact with Nrp1, PlexinD1 [OMIM 604282], and Sema5B [LOCUS LINK ID 54437]. It will therefore be interesting to determine if any of these genes are linked to the disease locus. Transient ischemia during hindbrain development may also contribute to Moebius syndrome [e.g., Verzijl et al. 1999]. As VEGF-A gene expression is regulated by hypoxia, defective VEGF signaling might contribute to the aetiology of sporadic Moebius cases by simultaneously disturbing hindbrain vascularisation and facial nerve development.

Low levels of VEGF contribute to postnatal motor neuron degeneration in mice, in which motor neuron cell death has been attributed to decreased neuroprotection by VEGF or reduced vascular perfusion of motor nerves [Oosthuysse et al. 2001]. In addition to this mechanism, our findings raise the possibility that motor nerves forming in the presence of low VEGF levels may already be compromised at birth and therefore predisposed to succumb to further insults during postnatal life. An association between abnormal motor neuron development
and postnatal motor neuron degeneration has previously been demonstrated in the Loa mouse, which carries a missense mutation in the heavy chain of cytoplasmic dynein [Hafezparast et al. 2003]. Moreover, low VEGF levels due to an inborn promoter polymorphism predispose humans to Amyotrophic Lateral Sclerosis, a fatal motor neuron disease [Lambrechts et al. 2003].

Conclusion

VEGF controls the migratory behavior of facial branchiomotor neurons in the developing brain in a mechanism that relies on the VEGF164 isoform and its receptor Nrp1. VEGF164 cooperates with the alternative Nrp1 ligand Sema3A to pattern different compartments of the same nerve, where VEGF164 is required only for soma guidance and Sema3A only for axon guidance. Altered VEGF expression during embryogenesis may therefore lead to neurological defects in addition to vascular deficiency.

Materials and methods

Animals

To obtain embryos of defined gestational ages, animals were mated in the evening, and the morning of vaginal plug formation was counted as 0.5 dpc. To stage-match embryos within a litter, or between litters from different matings, we compared somite numbers [10.5 dpc] or facial and limb development [11.5–14.5 dpc]. Nrp1−/− mutants have been described previously [Kitsukawa et al. 1997]. To obtain 14.5-dpc Nrp1−/− mutants, heterozygous females in a CD1 background were mated with heterozygous males in the JF1 background (kindly provided by J. Raper [University of Pennsylvania, Philadelphia, PA]). VEGF−/− or VEGF120/120 and VEGF164/164 mutants have been described previously [Carmeliet et al. 1999; Stalmans et al. 2002]. Male mice containing the LacZ reporter in the 3′-UTR of one VEGF-A allele [Misquero et al. 1999] were bred with wild-type CD1 females to produce heterozygous embryos for visualization of β-galactosidase activity.

In situ hybridization, immunohistochemistry, and visualization of β-galactosidase activity

Immunohistochemistry and visualization of β-galactosidase activity were performed as described previously [Ruhrberg et al. 2002; Hafezparast et al. 2003], with the exception that nerves were visualized with rabbit anti-neurofilament antibodies (Chemicon) followed by horseradish-peroxidase conjugated anti-rabbit antibodies (Dako) as described in Maina et al. [1997]. For in situ hybridization, tissues were hybridized with digoxigenin-labeled probes specific for the following genes: Is1 [Ericson et al. 1992], Nrp1 [He and Tessier-Lavigne 1997], Sema3A and Sema3C [gifts of M. Tessier-Lavigne, Stanford University, Stanford, CA; Phox2b [Pattyn et al. 2000], and VEGF164 [Breier et al. 1992]. In situ hybridization was performed as described in Hafezparast et al. [2003], with the exception that VEGF-A was detected using the method described in Riddle et al. [1993]. Images were recorded using the MZ16 microscope [Leica] equipped with a ProgRes C14 digital camera [Jenoptik] and processed using Openlab 2.2 software [Improvision Ltd.] and Adobe Photoshop 7.0 [Adobe Systems, Inc.].

Explant cultures

For explant cultures, hindbrains were dissected in cold L15 [In-vitrogen] free from branchial arch tissue, axons, sensory ganglia, and pial membranes. Hindbrain tissue was cultured for 1 or 2 d on Costar culture inserts with an 8-micrometer pore size [Corning, Inc.] in Neurobasal medium containing B27 supplement [Invitrogen]. In some experiments, explants were cultured in the presence of a function-blocking antiserum for VEGF [affinity-purified rabbit anti-mouse VEGF; Research Diagnostics, Inc.] or normal rabbit IgG [R&D Systems]. Affi-Gel heparin beads [Bio-Rad Laboratories] were washed several times in PBS and soaked in VEGF164 [Preprotech Ltd.] overnight at 4°C prior to implantation into explants. Explants were processed for in situ hybridization or immunohistochemistry as described above.

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