The parallel growth of motoneuron axons with the dorsal aorta depends on Vegfc/Vegfr3 signaling in zebrafish

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SUMMARY
Blood vessels and neurons grow often side by side. However, the molecular and cellular mechanisms underlying their parallel development remain unclear. Here, we report that a subpopulation of secondary motoneurons extends axons ventrally outside of the neural tubes and rostrocaudally as a fascicle beneath the dorsal aorta (DA) in zebrafish. We tried to clarify the mechanism by which these motoneuron axons grow beneath the DA and found that Vegfc in the DA and Vegfr3 in the motoneurons were essential for the axon growth. Forced expression of either Vegfc in arteries or Vegfr3 in motoneurons resulted in enhanced axon growth of motoneurons over the DA. Both vegfr3 morphants and vegfc morphants lost the alignment of motoneuron axons with DA. In addition, forced expression of two mutant forms of Vegfr3 in motoneurons, potentially trapping endogenous Vegfc, resulted in failure of growth of motoneuron axons beneath the DA. Finally, a vegfr3 mutant fish lacked the motoneuron axons beneath the DA. Collectively, Vegfc from the preformed DA guides the axon growth of secondary motoneurons.

KEY WORDS: Vegfc, Vegfr3, Motoneuron, Wiring, Guidance

INTRODUCTION
Blood vessels exhibit hierarchical branching from the central vessel (aorta) to the periphery (capillaries in the organs). Similarly, the neural network is formed according to the hierarchical connection originating from the central nervous system to the peripheral nervous system. Blood vessels and neurons are found side by side in the vertebrates (Bearden and Segal, 2005; Martin and Lewis, 1989), suggesting that both mutually affect the growth direction or that both develop following the guidance cues from the tissue surrounding them. Indeed, neurons extend or make contact according to repulsive or attractive cues, including Ephrin/Eph, Netrin/Unc5, Semaphorin/Plexin and Neuropilin (NRP), and Slit/Robo signaling, to form the network in the whole body (Carmeliet and Tessier-Lavigne, 2005; Guan and Rao, 2003). During blood vessel formation, migration and retraction of endothelial cells also follow these guidance cues (Adams and Eichmann, 2010; Eichmann et al., 2005; Larrivée et al., 2009).

There are three possible patterns for parallel growth of neurons and blood vessels: simultaneous growth, one preceding the other and common precursor vessels expressing Vegfr2 (Kdrl) in the main trunk (Lawson et al., 2002). Endothelin 3 released from smooth muscle cells of the carotid artery contributes to the alignment of sympathetic neurons with the carotid artery (Makita et al., 2008). The essential role of vascular endothelial growth factor A (VEGFA) from the preformed sensory neurons for vascular patterning has been reported in mice; loss of nerve-derived VEGFA results in impaired arterial differentiation in the skin of mice (Mukouyama et al., 2002). The congruence of neurons and blood vessels is also found in adult mouse skin, suggesting that VEGFA might not only regulate branching pattern of blood vessels but might also promote co-alignment of both nerves and vessels (James and Mukouyama, 2011). Recently, peripheral nerve-derived VEGFA and CXCL12 have been reported to regulate the patterning of arteries (Li et al., 2013). In addition, VEGFA not only functions as an axonal guidance cue but also regulates neuronal migration and plays a trophic factor for neurons (Mackenzie and Ruhrberg, 2012).

The VEGF family consists of VEGFA, VEGFB, VEGFC, VEGFD (FGF) and PLGF (PGF). Whereas VEGFA mainly activates VEGFR1 (FLT1) and VEGFR2 (FLK1; KDR) expressed on the vascular endothelial cells (Olsson et al., 2006), VEGFB and PLGF activate VEGFR1. VEGFC and VEGFD activate VEGFR3 (FLT4), which is mainly expressed on lymphatic endothelial cells (Karkkainen et al., 2004). During developmental angiogenesis of zebrafish, Vegfa from somites determines the fate of angioblasts and common precursor vessels expressing Vegfr2 (Kdr) in the main trunk (Lawson et al., 2002). Posterior cardinal vein (PCV) segregated from the dorsal aorta (DA) expresses Vegfr3 (Flt4), although the DA does not express Vegfr3 but rather Vegfc (Covassin et al., 2006; Lawson et al., 2001). In the central nervous system, VEGFR3 is expressed in the neural progenitor cells in Xenopus laevis and mouse embryos (Le Bras et al., 2006). VEGFC is able to stimulate VEGFR3-expressing neural stem cells in mice (Calvo et al., 2011). The proliferation of neural progenitor cells depends on the VEGFC/VEGFR3-mediated signal. In addition, VEGFC acts as

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a neurotransmitter for dopamine neurons (Piltonen et al., 2011). These reports indicate that the signal mediated by VEGFC/VEGFR3 is not restricted to within the mesoderm-derived cells but is also used outside of mesodermal tissues. Consistent with this, in zebrafish, Vegfc is required for coalescence of endodermal cells in the anterior midline and for the initial formation of dorsal endoderm (Ober et al., 2004).

Among the primary motoneurons of zebrafish [rostro primary (RoP), middle primary (MiP) and caudal primary (CaP) motoneurons] and CaP-like secondary motoneurons, RoP, CaP and CaP-like motoneurons exit the neural tube and extend their axons ventrally towards the axial vessels (Lewis and Eisen, 2003). In addition to these motoneurons, dorsallyventrally projecting secondary motoneurons, ventrally projecting secondary motoneurons and intermyotomal secondary motoneurons extend axons ventrally (Asakawa et al., 2013; Menelaou and McLean, 2012). In contrast to the initial neural axon growth of these motoneurons, intersegmental vessels sprout from the DA and extend dorsally towards the neural tube (Isogai et al., 2001). However, once the former and the latter reach the ventral-most and dorsal-most points, respectively, both extend rostrally and caudally along the anterior-posterior axis. These neural and vascular networks during embryogenesis can be spatiotemporally monitored in transgenic fish in which fluorescence proteins are produced under the control of neuron-specific or endothelial cell-specific promoters.

Here, we demonstrate the growth of secondary motoneuron axons descending ventrally and extending both rostrally and caudally as a fascicle beneath the DA using transgenic fish expressing fluorescent proteins: monomeric Cherry (mCherry) in endothelial cells and green fluorescent protein (GFP) in motoneurons. We show that the parallel growth of secondary motoneuron axons with the preformed fascicle beneath the DA using transgenic fish expressing fluorescent proteins, (z)Vegfc, (z)Vegfr3 tagged with Flag followed by 2A peptide and mCherry were subcloned into pcdNA3.1 (Invitrogen) and named as follows: pcdNA3.1 [(z)Vegfr3-f2amcherry] and pTol mnx2b [(z)Vegfr3-f2amcherry] (Asakawa et al., 2013). The pDNA encoding partial extracellular domain of human VEGFR3 fused with human Ig Fc fragment was subcloned into pcdNA3.1 and pTol mnx2 plasmids and named as follows: pcdNA3.1 (h)VEGFR3-Fc and pTol mnx2b (h)VEGFR3-Fc. The DNA encoding (z)Vegfr3 was subcloned into pCS2+ and pTol flt1 plasmids and named as follows: pCS2+ (z)Vegfc and pTol flt1 (z)Vegfr3. mCherry cDNA was inserted into pTol flt1 (pTol flt1:mcherry). pTol usGal4-specific recognition sequence; for upstream activating sequence; egfp was used to identify motoneurons by being injected into one-cell-stage Tg(mnx2b:gff) embryos.

MATERIALS AND METHODS
Zebrafish and transgenesis

The experiments using zebrafish were approved by the institutional animal committee of National Cerebral and Cardiovascular Center and performed according to the guidelines of the Institute. Zebrafish (Danio rerio) embryos were obtained from natural spawning of laboratory lines. Tg(fli1a:egfp)y1/1 fish were kindly provided by Nathan Lawson (University of Massachusetts Medical School, MA, USA). Tg(mnx1:gfp)m22 fish were obtained from the Zebrafish International Resource Center (University of Oregon, OR, USA). Tg(mnx2b:gfpy1) fish in which Gal4FF was used under the BAC-derived mnx2b promoter were established (Asakawa et al., 2008). Mutant vegfr3<sup>96hpf<sup>2</sup></sup> (expando) in double-transgenic Tg(fli1a:egfp)y1:(kdrl:mcherry) was previously reported (Hogan et al., 2009). Zebrafish were raised, injected and maintained under standard laboratory conditions (Westerfield, 2000). We used wild-type (AB), Tg(fli1a:myr-mcherry), Tg(buc:gfp), Tg(mnx1:gfp)y22, Tg(mnx2b:gfpy1) and Tg(fli1a:egfp)y1 embryos of either sex. Tg(fli1a:myr-mcherry) fish were developed by injecting the Tol2-based plasmid containing fli1a promoter followed by cDNA coding myristoylated (Myr) mCherry (pTol fli1a:myr-mcherry; 25 ng) with Tol2 transposase mRNA (25 ng) into one-cell-stage embryos of AB fish. Embryos were selected at 2 days post-fertilization (dpf) for high expression and grown to adults, among which germline founders were identified by specific expression of Myr-mCherry in the blood vessels.

Plasmids

pTol fli1a vector was constructed by modifying pTol2 vector and inserting the fli1a promoter as a driver of expression of the target molecule (Kawakami et al., 2004; Lawson and Weinstein, 2002). pTol mnx2b vector was similarly constructed by inserting the mnx2b promoter (Asakawa et al., 2012). The pTol flt1 vector was constructed by inserting the flt1 (vegfr1) promoter/enhancer corresponding to conserved non-coding elements (9a and 11) and −181 bp of the flt1 gene (Bussmann et al., 2010). An oligonucleotide encoding the myristoylation (Myr) signal derived from Lyn kinase was subcloned into pmCherry-N1 vectors (Takara) to construct the plasmid expressing Myr signal-tagged mCherry. pTol fli1a:myr-mcherry was constructed by inserting Myr-mCherry cDNA into pTol fli1a vector.

The DNA encoding zebrafish (z)Vegfr3 tagged with Flag followed by 2A peptide and mCherry was subcloned into pcdNA3.1 (Invitrogen), pPBbsr2 (provided by Michiyuki Matsuda, Kyoto University, Kyoto, Japan) for pcdNA3.1[(z)Vegfr3-f2amcherry] and pTol mnx2b [(z)Vegfr3-f2amcherry]. The DNA encoding (z)Vegfr3 lacking tyrosine kinase domain and tagged with Flage followed by 2A peptide and mCherry were inserted into pcdNA3.1 (Asakawa et al., 2013) and pTol mnx2 plasmids and named as follows: pcdNA3.1 (h)VEGFR3-Fc and pTol mnx2b (h)VEGFR3-Fc. The DNA encoding (z)Vegfc was subcloned into pCS2+ and pTol flt1 plasmids and named as follows: pCS2+ (z)Vegfc and pTol flt1 (z)Vegfr3. mCherry cDNA was inserted into pTol flt1 (pTol flt1:mcherry). pTol usGal4-specific recognition sequence; for upstream activating sequence; egfp was used to identify motoneurons by being injected into one-cell-stage Tg(mnx2b:gff) embryos.

FACS, RT-PCR and quantitative RT-PCR

Tg(fli1a:myr-mcherry);(mnx1:gfp)m22 embryos at 72 hours post-fertilization (hp) were digested with 5 mg/ml trypsin in PBS. The separated cells were sorted by a FACS Aria III Cell Sorter (BD Biosciences) according to GFP fluorescence and mCherry fluorescence. The cells sorted by FACS were suspended in 0.1% fetal bovine serum (FBS) in PBS. RNAs were isolated using the RNeasy Micro Kit (Qiagen). For RT-PCR, RNAs were reverse transcribed by random hexamer primers using SuperScript III (Invitrogen) according to the manufacturer’s instructions. PCR was performed using SpeedSTAR HS DNA polymerase (Takara). Real-time quantitative (q) RT-PCR was performed using the QuantFast SYBR RT-PCR Kit (Qiagen). Primer sequences for the target mRNAs used for PCR and qRT-PCR amplification are described in supplementary material Table S1 and Fig. S3.

Microinjections of plasmids, morpholinos and quantum dot

Transient expression of fluorescent proteins, (z)Vegfc, (z)Vegfr3 tagged with Flag followed by 2A peptide and mCherry [(z)Vegfr3-F2AmCherry], (h)VEGFR3-Fc, and (z)Vegfr3deltaRTK tagged with Flag followed by 2A peptide and mCherry [(z)Vegfr3deltaRTK2AmCherry] was carried out using the Tol2 system (Kawakami et al., 2004). Capped Tol2 transposase mRNA (25 pg) and the Tol2 plasmids (25 ng) containing either the artery-specific flt1 promoter or the motoneuron-specific mnx2b promoter were co-injected into one-cell-stage transgenic embryos. Quantum (Q) dot 655 (Molecular Probes) was injected into the axial blood vessels at 4 dpf. Three nanograms of morpholino (MO) (Gene Tools) was injected into yolk of one-to-two-cell-stage embryos of transgenic fish. Details of Tol2 plasmids, MOs and primers for the validation of inhibition of splicing by MOs are described in supplementary material Table S1 and Fig. S3.

Ex vivo assay

Tg(mnx1:gfp)m22 embryos at 36 hpf were manually dechorionated with fine forceps and were rinsed in sterile 0.5× E3 medium. Embryos were cut at the dorsoventral line between yolk and caudally extended yolk using operating scissors to expose the neural tissues of the trunk. Embryos, from which heads had been removed, were embedded in matrigel (BD Biosciences) with or without recombinant human (h)VEGF (R&D Systems). Embedded embryos were immersed in culture media consisting of CO2-independent media (Gibco) supplemented with 1× penicillin/streptomycin (Gibco) in 5% CO2 at 28°C for 12 hours.

Vegfr inhibitors

Tg(fli1a:myr-mcherry);(mnx1:gfp)m22 embryos were incubated from 60 to 96 hpf in E3 medium containing 25 μM max51 (Merck) or 0.5 μM ki8751
Cell culture and preparation of conditioned medium
Human umbilical vein endothelial cells (HUVECs) were cultured in EBM2 medium with essential growth media (Lonza). HEK293 cells and 293T cells were cultured in DMEM (Invitrogen) containing 10% FBS supplemented with 1× penicillin/streptomycin ( Gibco). Cells were transfected using Lipofectamine2000 according to the manufacturer’s instruction (Invitrogen). Conditioned media (CM) containing (z)Vegfc and (h)VEGFR3-Fc were prepared by the culture media of the 293T cells, transfected with pCS2 (z)vegf and pcDNA3.1 (h)VEGFR3-Fc plasmids, respectively. Media were replaced with FBS-free medium 24 hours after transfection. FBS-free media from transfected cells were collected after 24 hours incubation and concentrated through an Amicon Ultra-4 filter device (10 kDa) (Millipore). The HEK293 cells stably expressing (z)Vegfr3 were established by transfecting the cells with pPBbsr2(z)vegf3-f2amcherry and pCMV-mPBase (neo–) by using Lipofectamine2000 (Invitrogen) according to the piggyBac system (Komatsu et al., 2011; Yusa et al., 2009). The cells were cultured for 2 weeks in DMEM media containing blasticidin (6 µg/ml). Those surviving under the medium containing blasticidin were used as the cells stably expressing (z)Vegfr3. Expression of (z)Vegfr3 was confirmed by immunoblot analysis using anti-Flag antibody (Sigma-Aldrich).

Immunoblot analyses and precipitation
Cells were lysed in a lysis buffer containing 10 mM HEPES (pH 7.9), 400 mM NaCl, 0.1 mM EDTA, 5% glycerol, 1 mM Dl-dithiothreitol (DTT) and a protease inhibitor cocktail (Roche). Whole-cell lysates were resolved by SDS-PAGE followed by the electrophoretic transfer onto a polyvinyl difluoride membrane (Millipore). The antibodies in this study for immunoblot analyses were: anti-phospho Erk1/2 (Santa Cruz), anti-Erk1/2 (Cell Signaling Technology), anti-Flag, anti-β-actin (Sigma-Aldrich), horseradish peroxidase-coupled goat ant-mouse IgG, horseradish peroxidase-coupled goat anti-rabbit IgG (GE Healthcare) and horseradish peroxidase-coupled goat anti-human IgG. (h)VEGFR3-Fc was precipitated on Protein G-agarose (GE Healthcare).

Image acquisition, processing and quantitative analyses of neurite outgrowth and of parallel growth
Pigmentation of embryos was inhibited by 1-phenyl-2-thiourea (Sigma). The embryos were treated with 100 µg/ml tricaine (Sigma), mounted in a drop of 1.5% low melting agarose in the E3 medium, and placed onto a glass-based dish. Fluorescence images were obtained using a FluoView FV1000 confocal microscope (Olympus) with either a ×40 or a ×60 water objective lens (LUMPlanFL N, Olympus). z-stack images were 3D volume rendered with fluorescence mode using Velocity 3D Image Analysis Software (PerkinElmer). In the ex vivo assay, the number of GFP-positive outgrown neurites from the embryos was counted after 12 hours incubation at 28°C.

Impairment of parallel growth was defined as shortened or absent axons (defined as the length of axons becoming >90% shorter than those of either untreated or uninjected controls). For quantitative analyses of the parallel growth of motoneuron axons with the DA, ‘complete’ indicates the no defects in the motoneuron axons beneath the DA above yolk tubes. For that of increased parallel growth, ‘increase’ denotes the branching over the DA in addition to the parallel growth. The number of the embryos showing ‘impaired’ parallel growth divided by the total number of the embryos we observed is indicated as the percentage of the impaired parallel growth of the embryos.

Statistical analysis
Data are expressed as mean±s.d., as indicated in figure legends. Statistical significance for paired samples and for multiple comparisons was determined by Student’s t-test and by one-way analysis of variance with Tukey’s test, respectively. Data were considered statistically significant if P value was less than 0.05.

RESULTS
Parallel growth of motoneuron axons and dorsal aorta
To examine the alignment of neurons and blood vessels during zebrafish embryogenesis, we monitored the development of neurons and blood vessels simultaneously using transgenic (Tg) fish expressing Myr-mCherry in blood vessels under the fli1a promoter and GFP in neurons under the hue promoter Tg(fli1a:myr-mcherry);(hue:gfp) (Park et al., 2000). We noticed GFP-positive axons just beneath the DA at 4 dpf (Fig. 1A). The axons beneath the DA seemed to be a single axon fascicle as confirmed by a single scan image of the lateral view and by a cross-section of confocal stack images of the lateral views (Fig. 1B). Therefore, to observe how and when the axon started to grow, we time-lapse imaged the extension of axons and found that the axons grew rostrally and caudally along the pre-existing DA from 3 dpf (Fig. 1C,D; supplementary material Movie 1).

To ascertain whether axons beneath the DA developed from motoneurons or other neurons, we observed embryos expressing GFP under the mnx1 promoter, which is activated in motoneurons, crossed with fish expressing mCherry under the control of the fli1a promoter, Tg(fli1a:myr-mcherry);(mnx1:gfp) (Flanagan-Stee et al., 2005). We noticed that the axons derived from motoneurons marked by GFP located beneath the DA at 4 dpf (Fig. 2A). Therefore, we time-lapse imaged the embryos and found that the axons of motoneurons started developing caudally and rostrally as a single fascicle beneath the DA from 3 dpf (Fig. 2B-D; supplementary material Fig. S1A and Movie 2). To examine which motoneuron extends axons beneath the DA, we tried to label single cells by injecting pTol uas:egfp into Tg(fli1a:myr-mcherry);(mnx2b:gfp) embryos. The motoneurons marked by EGFP in the neural tube extended axons towards the ventromedial area and beneath the dorsal aorta. In addition, when we performed the single-cell labeling experiment, those motoneurons seemed to exist in every somite and extend axons outside of the neural tube bilaterally (Fig. 2E; supplementary material Fig. S1B-D and Movie 3). Therefore, the subpopulation of the motoneurons we identified in the present study corresponds to the intermyotomal secondary motoneurons or the ventromedial secondary motoneurons, as previously reported (Asakawa et al., 2013; Menelaou and McLean, 2012).

Motoneurons express Vegfr3
The thoracic duct (TD), a major lymphatic vessel, is known to develop between the DA and the PCV (Yaniv et al., 2006). Thus, we tried to examine the exact location of the TD and motoneuron axons beneath the DA. Motoneuron axons were located between the DA and TD. The TD formed at 5 dpf, whereas extension of motoneuron axons was completed by 3.5 dpf. Therefore, the extension of axons preceded the formation of the TD (Fig. 3A). In addition, when we looked closely at TD formation, it developed along the preformed axons of motoneurons in Tg(fli1a:egfp) embryos transiently expressing mCherry by Tol2-mediated gene transfer using mnx2b promoter, which is activated in motoneurons (Asakawa et al., 2012; Kawakami et al., 2004) (supplementary material Fig. S2A). These results prompted us to assume that the alignment of the motoneuron axons and the DA might depend on the molecular mechanism by which the TD develops along the DA, which is probably dependent on Vegfc-mediated signaling, because lymphatic vessel development depends on Vegfc/Vegfr3 signaling in zebrafish (Küchler et al., 2006).

Vegfc is essential for development of the TD and is expressed in the DA (Covassin et al., 2006; Küchler et al., 2006; Villefranc et al.,
We confirmed the expression of vegfc mRNA in the DA of embryos at 24, 48 and 72 hpf by in situ hybridization (Fig. 3B; supplementary material Fig. S2B,C). Then, we hypothesized that motoneurons might express Vegfr3 similar to lymphatic endothelial cells and extend axons in response to Vegfc from the DA. To test this hypothesis, we examined the expression of vegfr3 mRNA by in situ hybridization. We could detect it in the PCV and in the intersomitic vessels (ISVs) but not in the neurons at 24 or 48 hpf (supplementary material Fig. S2D). At 72 hpf, even its expression in the PCV was not clear. vegfr3 mRNA expression was found in the neural tube when the reaction was extended (supplementary material Fig. S2E). Therefore, to analyze vegfr3 mRNA expression in embryos at 72 hpf, we performed RT-PCR. Motoneurons and endothelial cells were collected by FACS from Tg(fli1a:myr-mcherry);(huc:gfp) embryos. RNAs from the collected cells were first analyzed by RT-PCR. Motoneurons and endothelial cells expressed vegfr3 mRNA (Fig. 3C). The purity of RNA for RT-PCR was confirmed by the observation that mnx1 mRNA was detected exclusively in motoneurons and that tie1 mRNA, tie2 (Tek) mRNA and fli1a mRNA were detected only in endothelial cells (Fig. 3C). To examine quantitatively the expression of vegfr3 mRNA, we performed qRT-PCR and found that vegfr3 mRNA was detected in motoneurons although its expression was less than that in the endothelial cells (Fig. 3D). Motoneuron expressing Vegfr3 responds to Vegfc

To test whether motoneurons respond to Vegfc, we examined neurite sprouts from motoneurons in an ex vivo model. The Tg(mnx1:gfp)embryos, from which heads and yolks were dissected from the trunks at 36 hpf, were incubated in matrigel containing recombinant human (h)VEGFC (Fig. 4A). We measured the number of sprouting neurites marked by GFP from embryos in five groups: those incubated without recombinant (h)VEGFC; those treated with control DMSO and incubated with (h)VEGFC; those treated with vegfr3 morpholino (MO) and incubated with (h)VEGFC; those treated with (h)VEGFC plus the VEGFR3 inhibitor maz51 (Ny et al., 2008); and those treated with (h)VEGFC plus the VEGFR2 inhibitor ki8751 (Kubo et al., 2005) (Fig. 4B,C; supplementary material Fig. S3A). (h)VEGFC-induced sprouting of neurites was inhibited by either knockdown or inhibition of Vegfr3 but not by inhibition of Vegfr2, suggesting that neurite outgrowth from motoneurons depends on Vegfc/Vegfr3 signaling.
We then verified whether (z)Vegfr3 can respond to (h)VEGFC in cultured cells. Cells expressing (z)Vegfr3 exhibited phosphorylation of extracellular-regulated kinase (Erk) when stimulated with (h)VEGFC (Fig. 4D), confirming that motoneurons extend neurites in response to (h)VEGFC.

**Vegfc/Vegfr3 signaling augments axon growth of motoneurons**

To test whether motoneurons respond to Vegfc in vivo, we examined the effects of overexpression of (z)Vegfc in arteries or (z)Vegfr3 in motoneurons on the axonal growth of motoneurons. We observed the growth of axons in Tg(mnx1:gfpl)ml2 embryos injected with QDot 655 into the blood vessels at 4 dpf to visualize both motoneurons and blood vessels. We used the zebrafish fltl promoter/enhancer and mnx2b promoter for transient overexpression of (z)Vegfc and (z)Vegfr3 in arteries and motoneurons, respectively (Asakawa et al., 2012; Bussmann et al., 2010; Ny et al., 2008). Artery-specific expression mediated by the fltl promoter/enhancer and motoneuron-specific expression by mnx2b promoter was confirmed by transient expression of mCherry in Tg(fli1a:egfp)vl1 and Tg(mnx1:gfpl)ml2.
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Fig. 3. Motoneurons express Vegfr3. (A) 3D-rendered confocal stack of fluorescence images (lateral view) of a Tg(fli1a:myr-mcherry);(mnx1:gfp)m12 embryo at the times indicated in the top panels. Top panels, blood vessels and lymphatic vessels marked by mCherry; middle panels, merged images of GFP and mcherry; bottom panels, motoneurons marked by GFP. Arrows indicate the TD. Dashed lines in the middle panel indicate the outline of the TD. Note that the axon extension preceded TD development and that the axons were formed between DA and TD. Scale bar: 25 μm. (B) In situ hybridization analyses of vegfc mRNA in embryos at the indicated stages. Arrows and arrowheads indicate vegfc mRNA detected in the DA in the lateral views (left) and cross-section views (right), respectively. (C) RT-PCR analyses using the primers indicated at the left and RNAs prepared from the Tg(fli1a:myr-mcherry);(mnx1:gfp)m12 embryos by FACS at 72 hpf. (D) Real-time quantitative RT-PCR analyses using RNAs prepared from embryos at 72 hpf by the primers as indicated at the bottom. †P<0.05. Error bars indicate s.d. DA, dorsal aorta; EC, endothelial cells; MN, motoneurons; N, notochord; NT, neural tube; PCV, posterior cardinal vein; TD, thoracic duct.

Vegfc/Vegfr3 signaling is essential for axon growth of motoneurons beneath the dorsal aorta

To examine whether Vegfc/Vegfr3 signaling between the DA and motoneurons is essential for the axon extension of motoneurons beneath the DA, we investigated the effect of inhibition of Vegfc/Vegfr3 signaling or depletion of Vegfc or Vegfr3 on the alignment of motoneurons beneath DA in Tg(fli1a:myr-mcherry);(mnx1:gfp)m12 embryos treated with max51 and kis751 to inhibit Vegfr3 and Vegfr2, respectively. Inhibition of Vegfr3 but not Vegfr2 clearly reduced the diameter of axons and the number of branches, indicating an important role for Vegfr3-mediated signaling in the growth of axons beneath the DA (Fig. 6A). We then injected either vegfc MO or vegfr3 MO into one-cell-stage Tg(fli1a:myr-mcherry);(mnx1:gfp)m12 embryos (Fig. 6B; supplementary material Fig. S3B,C). Depletion of either Vegfc or Vegfr3 resulted in less axon extension of motoneuron beneath the DA (Fig. 6B,C). Depletion of Vegfc did not affect the ventral axon growth of motoneurons (supplementary material Fig S3D). To examine the requirement of Vegfr3 for the growth of motoneuron axon beneath the DA, we examined the presence of motoneuron axons beneath the DA by immunohistochemistry of Tg(fli1a:eGFP)Y1 and Tg(fli1a:eGFP)Y1;flt4mut662 (expando mutant;Vegfr3 I1034F) using znpl antibody against synaptotagmin 2. Whereas motoneuron axons were detectable beneath the DA in the control embryos, those in expando mutants (Vegfr3 mutants) were hardly detectable (Fig. 6D). These data indicate the essential role for Vegfc/Vegfr3 signaling in the parallel growth of the motoneuron axons and DA.

Motoneuron-specific inhibition of Vegfr3 reduces the parallel growth of motoneuron axons and dorsal aorta

Because the effect of MO is not restricted in either DA or motoneurons, we examined the effect of Vegfr3 inhibition specifically in motoneurons on axon extension beneath the DA. To inhibit Vegfr3, we overexpressed a mutant form of (h)VEGFR3 fused with human IgG Fc, (h)VEGFR3-Fc (Mäkinen et al., 2001; Ober et al., 2004). We first confirmed that this mutant could inhibit both (h)VEGFC- or (z)Vegfc-triggered signaling. HUVECs were stimulated with recombinant (h)VEGFC in the presence or absence of (z)Vegfc in arteries resulted in an increase in branching of motoneuron axons over the DA, whereas that of control (mCherry) did not (Fig. 5A,B,E). Similarly, overexpression of (z)Vegfr3 in motoneurons induced branching of motoneuron axons over the DA, whereas that of control (mCherry) did not alter any parallel growth of motoneuron axons with the DA (Fig. 5C-E). Overexpression of Flag-tagged (z)Vegfr3 was confirmed by expression of mCherry in the motoneurons in the neural tube (supplementary material Fig. S4C). These results indicate that gain of Vegfc/Vegfr3 signaling between the DA and motoneuron axons enhances their congruency.
of (h)VEGFR3-Fc prepared by the conditioned medium of cells transfected with the plasmid expressing (h)VEGFR3-Fc. (h)VEGFC-dependent phosphorylation of Erk was reduced in the cells treated with (h)VEGFR3-Fc compared with those without (h)VEGFR3-Fc (supplementary material Fig. S5A). We also tested whether this human VEGFR3 mutant could inhibit zebrafish Vegfr3/Vegfc signaling by treating HEK293 cells stably expressing (z)Vegfr3 with (z)Vegfc in the presence or absence of (h)VEGFR3-Fc. (z)Vegfc-dependent phosphorylation of Erk via (z)Vegfr3 was inhibited by (h)VEGFR3-Fc (supplementary material Fig. S5B). Therefore, we examined the effect of overexpression of (h)VEGFR3-Fc exclusively in motoneurons on the parallel growth of motoneurons along the DA. We transiently and specifically expressed (h)VEGFR3-Fc using the mnx2b promoter in the motoneurons of Tg(fli1a:myr-mcherry);(mnx1:gfp)ml2 embryos. The embryos injected with the plasmid expressing (h)VEGFR3-Fc exhibited less axon extension of motoneurons than did the control embryos (Fig. 7A). The impairment of parallel growth of the motoneuron axons and DA was quantitatively analyzed (Fig. 7B). The number of the embryos showing the increased, normal or decreased parallel growth (PG) divided by the total number (n) of embryos counted (indicated at the top) is expressed as percentage of embryos.
we transiently expressed this (z)Vegfr3deltaRTK in motoneurons of Tg(mnx1:gfp)ml2 embryos and examined the alignment of motoneuron and DA by injecting Qdot 655 to visualize blood vessels. Overexpression of (z)Vegfr3deltaRTK resulted in reduced extension of motoneuron axons even though the (z)Vegfc was overexpressed in the arteries (supplementary material Fig. S5E). Collectively, these data indicate that Vegfr3 expression on motoneurons is essential for the extension of axons in response to Vegfc from the DA.

**DISCUSSION**

In the present study, we provide evidence that Vegfc functions as a guidance molecule for the axon growth of motoneurons. We found that Vegfr3 was expressed in motoneurons and that forced expression of the Vegfc-trapping (h)VRGFR3 mutant resulted in the inhibition of parallel growth of motoneuron axons beneath the DA expressing Vegfc. The TD, a main lymphatic vessel formed between the DA and PCV, needs Vegfc to grow in zebrafish (Küchler et al., 2006; Villefranc et al., 2013; Yaniv et al., 2006). We found that the formation of motoneuron axons beneath the DA preceded that of the TD. These spatiotemporal data suggest that axon development employs the same mechanism underlying the development of the TD, because both lymphatic vessel and motoneurons express Vegfr3. We cannot completely exclude the involvement of Vegfr2 because Vegfc activates the heterodimer complex of Vegfr2-Vegfr3 (Herbert and Stainier, 2011). We examined the effect of a Vegfr2 inhibitor (ki8751) on the parallel growth and noticed that theaxon beneath the DA seemed to be slightly affected, although 0.5 μM ki8751 did not affect the secondary sprouting from the PCV that is promoted by Vegfc. Therefore, we can conclude at least that Vegfc/Vegfr3 signaling is required for the alignment of the axons and DA.

**Fig. 6. Requirement of Vegfc and Vegfr3 for alignment of motoneuron axons and dorsal aorta.** (A) 3D-rendered confocal stack images (lateral view) of Tg(fli1a:myr-mcherry);(mnx1:gfp)ml2 embryos treated with control DMSO (Ctrl, upper panels), with maz51 (middle panels) or with ki8751 (lower panels) at 4 dpf. Left panels, merged images of mCherry and GFP; right panels, GFP images. White arrows indicate the motoneurons beneath the DA. Arrowheads indicate the reduced growth of motoneuron axons beneath the DA. Yellow arrows indicate the slight reduction of growth of motoneuron axon beneath the DA. (B) 3D-rendered confocal stack images (lateral view) of Tg(fli1a:myr-mcherry);(mnx1:gfp)ml2 embryos treated with control MO (Ctrl, top panels), with vegfc MO (middle panels) or with vegfr3 MO (bottom panels). Left panels, merged images of mCherry and GFP; right panels, GFP images. Arrows indicate the motoneuron axons beneath the DA. Arrowheads indicate the reduced interaction of motoneurons and dorsal aorta. (C) Quantitative analyses of impairment of parallel growth (PG) of the embryos grouped as in B. The number (n) of the embryos examined for quantitative analyses is indicated at the top. (D) Immunohistochemical study of Tg(fli1a:egfp)y1 embryos (top) and vegfr3hu4602(expando mutant);Tg(fli1a:egfp)y1 embryos using anti-znp1 antibody. Arrowheads and arrows indicate the presence or absence of axons beneath the dorsal aorta, respectively. Note that the TD is absent in the expando mutant. Scale bar: 25 μm. DA, dorsal aorta; TD, thoracic duct.

Why and how do the axons extend ventrally and grow beneath the DA? Although Vegfc is expressed in the DA at 48 hpf, it is expressed in the hypochord at an early stage (18-somite stage) (Covassin et al., 2006). We assumed that Vegfc from the hypochord might be essential for the ventral growth of axons and Vegfc from the DA for parallel growth of the axons beneath the DA. Knockdown of Vegfc did not affect the ventral growth but did affect the parallel growth with the DA, suggesting that the ventral axon growth depends on the other guidance cues. PlexinA3 expressed on motoneurons and sema3A2 (sema3ab) in somites, dcc on motoneurons and netrin in myoseptum have been reported to be involved in ventral growth of motoneuron axons from the cell body in the neural tube (Feldner et al., 2007; Lim et al., 2011). The axons of motoneurons grew more ventrally beyond the DA and grew beneath the DA. It is probable that some molecules from the PCV might attract the axons ventrally to the DA. Whereas Cxcl12a is expressed in the PCV at 72 hpf (Cha et al., 2012), Cxcr4, a receptor for Cxcl12, is expressed on neuronal cells (Tiveron and Cremer, 2008). Cxcl12a from the PCV might attract the axons ventrally to the DA. In addition, some repulsive cues from the DA might prevent the axons from touching the DA. During the pathfinding of axon growth, neurons utilize Ephrin/Eph, Netrin/Unc5, Semaphorin/Plexin and NP signaling and the Slit/Robo system (Carmeliet and Tessier-Lavigne, 2005). Because the DA expresses...
Fig. 7. Inhibition of Vegfr3 in the motoneuron results in impairment of alignment of the motoneuron axons and dorsal aorta. (A) 3D-rendered confocal stack images (lateral view) of Tg(fli1a:myr-mcherry);Tg(mnx1:gfp) embryos (upper panel) and those transiently expressing (h)VEGFR3-Fc in the motoneurons under the control of the mmx2b promoter by Tol2-mediated gene transfer (lower panels). Left panels, merged images of mCherry and GFP; right panels, GFP images. Arrows indicate the impairement of parallel growth of motoneuron axons beneath the dorsal aorta (DA). Scale bar: 25 μm. (B) Quantitative analyses of the impaired parallel growth (PG) of the embryos grouped as in A. The number (n) of the embryos observed indicated at the top.

Vegf for guiding axons

VEGF A is a bipotential, angiogenic and neurotrophic molecule. Vegf found as an angiogenic growth factor is capable of activating VEGFR2 expressed on neurons (Bearden and Segal, 2004; Jin et al., 2002). By contrast, VEGF A from neurons guides the congruence of blood vessels and neurons, involvement of other VEGFs, including VEGFC, in both are expressed in neural tissues, the expression of Np-2 in the PCV in zebrafish (Martyn and Schulte-Merker, 2004). Although the data; and K. Alitalo and N.M. wrote the paper.

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Competing interests statement

The authors declare no competing financial interests.

Author contributions

H.-B.K., K. Asakawa, K. Ando and T.K. performed the experiments; S.F., K.K., M.H., Y.-G.K., K.-W.K. and N.M. designed the research; S.F. and K.K. analyzed the data; and K. Alitalo and N.M. wrote the paper.

Supplementary material

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