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BDNF mediates non cell-autonomous regulation of sensory neuron position and identity

Melissa A. Wright^{1,2,3} and Angeles B. Ribera^{1,2,3}

¹Department of Physiology and Biophysics, University of Colorado at Anschutz Medical Center, Aurora, CO 80045

²Neuroscience Graduate Program, University of Colorado at Anschutz Medical Center, Aurora, CO 80045

³Medical Scientist Training Program, University of Colorado at Anschutz Medical Center, Aurora, CO 80045

Abstract

During development, neurons migrate considerable distances to reside in locations that enable their individual functional roles. Whereas migration mechanisms have been extensively studied, much less is known about how neurons remain in their ideal locations. We sought to identify factors that maintain the position of post-migratory dorsal root ganglion neurons, neural crest derivatives for which migration and final position play an important developmental role. We found that an early developing population of sensory neurons maintains the position of later born dorsal root ganglia neurons in an activity-dependent manner. Further, inhibiting or increasing the function of brain-derived neurotrophic factor induces or prevents, respectively, migration of dorsal root ganglia neurons out of the ganglion to locations where they acquire a new identity. Overall, the results demonstrate that neurotrophins mediate non cell-autonomous maintenance of position and thereby identity of differentiated neurons.

Keywords

dorsal root ganglion; sodium channel; nav1.6; non cell-autonomous; BDNF; sensory deprivation

Introduction

In both the peripheral and central nervous systems, migration results in neurons acquiring positions that differ from their sites of genesis. Whereas numerous studies have investigated migration mechanisms, much less is known about factors that lead to maintenance of neuronal position after migration has occurred. Neuronal activity plays an important role in regulating many differentiated properties of neurons. On this basis, we were interested in whether activity also plays a role in maintaining the final position of neurons after migration has occurred.

Clues to mechanisms by which activity might regulate a neuron's final position might come from activity's role in neuronal migration. Several recent studies point to electrical activity as an important regulator of migration of embryonic neurons (Bolteus and Bordey, 2004; Heng et al., 2007; Tam et al., 2000; Tong et al., 2009). In some instances, electrical activity

regulates cell migration through modulation of calcium signaling cascades (Tam et al., 2000; Tong et al., 2009). In addition, electrical activity has also been shown to regulate expression and localization of cell adhesion molecules (Pierre et al., 2001), which are important regulators of cell migration and positioning. These studies point to cell autonomous mechanisms through which electrical activity regulates position of developing neurons.

However, electrical activity also exerts non cell autonomous control over neuronal migration. In both the enteric nervous system (Hao et al., 2010) and the embryonic cortex (Behar et al., 2000; Bolteus and Bordey, 2004), activation of neurotransmitter receptors activate signaling cascades that regulate migration of post-mitotic neurons or neuronal precursors. For these cells electrical activity regulates migration non cell autonomously, through its effects on neurotransmitter release. Further, many neurotrophins, which guide migration, differentiation, and maintenance of neurons, are secreted in an activity-dependent manner (Bruno and Cuello, 2006; Kuczewski et al., 2009; Wu et al., 2004), suggesting another mechanism for non cell autonomous control of position of developing neurons.

Recent work points to an important role for the voltage-gated sodium channel *nav1.6a* in maintenance of position of post-migratory dorsal root ganglion (DRG) neurons (Wright et al., 2010). However, DRG as well as many other neurons express the gene (*scn8aa*) encoding *nav1.6a* (Novak et al., 2006), indicating that *nav1.6a* could act either cell autonomously or non cell autonomously to regulate DRG position. Here, using mosaic knock-down of *nav1.6a*, we determined that *nav1.6a* exerts non cell autonomous control over DRG position, through channel activity in early-developing Rohon-Beard (RB) sensory neurons. Further, brain-derived neurotrophic factor (BDNF) rescues the DRG migration phenotype in embryos with reduced *nav1.6a*, whereas BDNF function-blocking antibodies induce migration of differentiated DRG neurons in control embryos. Consistent with previous work, once outside the ganglion, dorsal root ganglion neurons transdifferentiate into sympathetic neurons, indicating that position plays a key role in maintaining neuronal identity. Our present results indicate that by preventing migration of DRG neurons, BDNF maintains neuronal identity. Moreover, when embryos are raised under conditions that minimize stimulation of RB neurons, migration of post-migratory DRG neurons increases, revealing that maintenance of the DRG sensory system requires early tactile input.

Materials and Methods

Animal care

Adult zebrafish (*Danio rerio*) were maintained at 28.5°C on a 10 hour dark/14 hour light cycle in the Center for Comparative Medicine (CCM) at the University of Colorado Denver and bred according to established procedures (Westerfield, 1995). The University of Colorado Committees on Use and Care of Animals approved all animal protocols. The transgenic zebrafish line Tg(-3.4neurog1:GFP) was kindly provided by Uwe Strähle at the University of Heidelberg (Blader et al., 2003). The *macho* mutant line (*mao1*^{tt261/+}) was obtained from the Tübingen Stock Center (Granato et al., 1996).

Tetrodotoxin injections

Tg(-3.4neurog1:GFP) embryos were raised in Embryo Media (EM) until they reached 16-19 hpf. Embryos were then manually dechorionated and injected with either dH₂O or tetrodotoxin (TTX; EMD Chemicals, Gibbstown, NJ). We used 0-500 μM for dose-response experiments and 250 μM for all other experiments. We included 1% FastGreen (Sigma-Aldrich Corp., St. Louis, MO) in the injection solution to allow visual tracking and confirmation of transfer of the injected solution from the yolk into the animal blastomeres.

DRG position was analyzed in live embryos at 48, 72, and 96 hpf. Embryos were fixed at 96 hpf and processed for GFP and HuA immunoreactivities.

Morpholino injections

Morpholino antisense oligonucleotides (morpholinos) were synthesized by Gene Tools (Philomath, Oregon) and used to knock down *nav1.6a* protein, as previously reported (Pineda et al., 2005; Pineda et al., 2006). One targeted the predicted translation start methionine of *nav1.6a* and had 25 residues with the following sequence: 5'GGGTGCAGCCATGTTTTCATCCTGC-3'. The second morpholino (5'-ATGTGGTTTGGATCAATA-CTTACTC-3') targeted the splice junction between exons 3 and 4. Similar results were obtained with the two different morpholinos and we refer to them collectively as 1.6mo. Embryos injected with 1.6mo are referred to as morphants throughout this paper. Two control morpholinos (1.6ctl), differing from the translation blocking or splice blocking morpholino by five base mismatches, were synthesized. Embryos injected with either 1.6ctl are referred to as controls. Morpholinos (3-4 ng/nl) were injected into 1- to 2-cell stage embryos.

Mosaic embryos

Mosaic embryos were created by injecting single cells of 64-128 cell stage wild type or Tg(-3.4neurog1:GFP) embryos with 0.3 mM 1.6mo or 1.6ctl and lineage tracer (3% lysinated-fluorescein or rhodamine-labeled dextran; 10,000 MW; Invitrogen/Molecular Probes, Carlsbad, CA). Embryos were fixed at 72 hpf and processed for the dextran label and either GFP or HuA immunoreactivities, as described below.

Generation of transient transgenics

The 5' upstream region of the *scn8aa* gene contained within the DKEY-9P24 BAC was cloned in front of mCherry by the Transgenic and Gene Targeting Core of the Rocky Mountain Neurological Disorders Center Core at UCD AMC (P30 NS048154). Transient *scn8aa*:mCherry transgenics were generated by injection of 100 ng/ μ l of DNA along with 1.6ctl or 1.6mo and 1% FastGreen directly into the raised cell of 1-cell Tg(-3.4neurog1:GFP) embryos.

Neurotrophin and BDNF antibody injections

NGF (N-130, Alomone Labs, Israel), NT3 (N-260, Alomone Labs, Israel), or BDNF (B-250, Alomone Labs, Israel) was injected into 1.6 morphant or control Tg(-3.4neurog1:GFP) embryos at 16, 24, or 48 hpf at a concentration of 50 ng/mL. For dose-response experiments, BDNF (1 - 500 ng/ml) was injected into 48 hpf Tg(-3.4neurog1:GFP) 1.6 morphants. Chicken anti-BDNF (100- 1000 μ g/mL; AF248, R&D Systems, Minneapolis, MN) was injected into 48 hpf Tg(-3.4neurog1:GFP) embryos. DRG position was analyzed in live 96 hpf larvae.

Touch assays

The touch response was assessed in embryos that swam spontaneously with a normal pattern to ensure that motor deficits were not present. For each trial, the embryo was touched on the trunk with a tungsten needle and assigned a score. A score of 1 indicated a normal response (full swimming); a score of 0.5 indicated an abnormal response (twitching, but not swimming); a score of 0 indicated no response. We performed 10 trials per embryo and summed scores for each trial, resulting in final scores of 0-10.

Sensory Manipulation

Embryos raised under conditions of increased tactile stimulation were raised under normal conditions except for manual dechoriation at 16 hpf and stimulation with a soft poker eight times a day between 16 and 96 hpf. Embryos raised under conditions of decreased tactile stimulation were raised from 16-96 hpf at half density (40-50 versus 80-100 per dish), not moved, and kept in a separate incubator that was not opened. For all groups, media was changed at 16 hpf but not again until 96 hpf.

Immunocytochemistry

Whole-mount immunocytochemistry was performed as previously described with minor modifications (Svoboda et al., 2001). Briefly, embryos were fixed overnight (4°C) with 4% paraformaldehyde in PBS solution containing 0.1% Tween-20 (PBST). After washing with PBST, embryos were permeabilized by incubation in distilled H₂O, followed by exposure to 100% acetone at -20°C and a final collagenase (1 mg/ml) treatment. Embryos were blocked in 10% Heat Inactivated Goat Serum/PBST and incubated overnight (4°C) in 10% serum/PBST containing primary antibody. After extensive rinsing with PBST secondary antibody was applied. Embryos were then washed in PBST, post-fixed in 4% paraformaldehyde PBS solution, and rinsed again in PBST. Anti-HuC/D (mouse antibody, Invitrogen/Molecular Probes) was used at 1:1000; anti-fluorescein or rhodamine (rabbit antibodies, Invitrogen/Molecular Probes) were used at 1:200; goat-anti-mouse-Alexa568 and goat-anti-rabbit-Alexa568 secondary antibodies (Invitrogen/Molecular Probes) were used at 1:1000. For detection of GFP, the rabbit-anti-GFP-488 antibody (Invitrogen/Molecular Probes) was used at 1:400. For double immunocytochemistry, when used with mouse antibodies, the anti-GFP antibody was added during incubation with the secondary antibody; when used with rabbit antibodies, the anti-GFP antibody was added following extensive washing of the anti-rabbit secondary antibody.

Confocal imaging and analysis of DRG position

Confocal z-stacks were acquired using a Zeiss LSM5 Pascal Confocal Upright Microscope (Carl Zeiss Inc., Thornwood, NY) equipped with either a 10× or a 40× water immersion lens and the pinhole set to one Airy unit. For analysis of DRG position in live Tg(-3.4neurog1:GFP) embryos, specimens were mounted laterally in a drop of 0.5% Low Melting Point agarose containing 0.0002% tricaine (Sigma-Aldrich). DRG position was analyzed throughout the length of the embryo. GFP⁺ DRG cells were considered to be ectopic if they resided more than one cell body ventral to the spinal cord-notochord boundary.

Statistical analysis

All statistical analyses were performed using InStat3 software (GraphPad Software Inc., La Jolla, CA). Contingency tables were analyzed using Fisher's exact test to determine two-sided p values. The specific tests used for other analyses depended upon whether the data were normally distributed (e.g., Student t-test or ANOVA parametric tests) or not (e.g., nonparametric Mann-Whitney and Kruskal-Wallis tests) and are indicated in the text. For data analyzed for statistical significance with non-parametric tests, the graphs present the median with the inner quartiles, 5th and 95th percentiles, and any outliers (filled circles).

Results

Sodium current is required for maintenance of DRG position

Knock-down of the nav1.6a sodium channel protein increases the number of DRG neurons that migrate out of the ganglia to more ventral positions (Wright et al., 2010). To test

whether loss of sodium channel activity, rather than other effects of sodium channel protein, induces DRG migration, we blocked sodium channels pharmacologically with the sodium channel inhibitor tetrodotoxin (TTX). We used a transgenic line expressing GFP under the control of a portion of the *neurogenin 1* (*neurog1*) promoter, because neural crest expression of *neurog1* leads to a neuronal fate (McGraw et al., 2008). In control Tg(-3.4neurog1:GFP) embryos, the majority of GFP⁺ DRG neurons showed stereotypic positions at the spinal cord-notochord boundary, with only 1-2 ectopic DRG cells per embryo (Figs. 1A and C). Upon injection of TTX, the number of ectopic GFP⁺ DRG cells increased in a dose-dependent manner, reaching maximal effect at 250 μ M (Figs. 1B-C). These results indicate that loss of sodium channel activity promotes migration of cells away from the dorsal root ganglia.

The number of ectopic DRG cells in TTX-injected embryos increased after 3 days post-fertilization (dpf; data not shown), as would be expected if DRG neurons migrate after differentiating. To test directly whether the ectopic DRG cells that develop in TTX-injected larvae are differentiated neurons, we examined expression of the neuronal differentiation marker, HuA. Ectopic GFP⁺ cells in both control and TTX-injected larvae expressed HuA (Figs. 1D-E). Taken together, these data indicate that sodium channel inactivity increases the number of migratory differentiated DRG neurons in zebrafish larvae.

RB sodium current prevents ventral migration of DRG neurons

TTX injection affects sodium channels throughout the embryo (Fig. 1), preventing determination of the location and identity of the relevant sodium channels. However, previous work implicated the nav1.6a sodium channel encoded by the *scn8aa* gene (Wright et al., 2010). Nonetheless, developing DRG as well as other neurons express *scn8aa*, leaving the location of the relevant sodium channels an open question. To identify the cell type expressing the relevant sodium channels, we generated embryos that had mosaic knock-down of nav1.6a by injecting well-characterized antisense morpholinos targeting *scn8aa* (1.6mo; Pineda et al., 2005; Pineda et al., 2006) into single cells of 64-cell stage embryos (Figs. 2B-C, E-F). At 72 hpf, we immunostained embryos for the co-injected lineage tracer rhodamine dextran (RhDx) and HuA to determine the relationship between DRG position and morpholino distribution (Figs. 2 and 3).

Embryos mosaic for 1.6mo (Figs. 2B-C, E-F) showed a small increase in the number of migratory DRG neurons compared to 1.6ctl mosaic embryos (Figs. 2A and D; 1.56 versus 0.58, 1.6mo versus 1.6ctl; $p < 0.0001$, Mann-Whitney U-test). This finding demonstrates that mosaic knock-down of nav1.6a suffices to induce ventral migration of DRG neurons (Figs. 2E-F). However, less than 1% of ectopic DRGs in 1.6mo mosaic embryos were positive for the fluorescent tracer. In contrast, 96% of DRGs that contained the morpholino were normally-positioned (Figs. 2B-C; $p < 0.0001$; Fisher's exact test). Taken together, these data do not support the DRG as the location of the relevant sodium channels, making a cell autonomous requirement for nav1.6a in DRG neurons an unlikely possibility.

We next analyzed DRG position and the tracer distribution in 1.6mo mosaic embryos. DRG position was not affected by the presence of 1.6mo in motor neurons, intermediate interneurons, or dorsal interneurons (data not shown). However, embryos that contained fluorescent tracer-positive RB sensory neurons showed a significant increase in the number of migratory DRG neurons compared to embryos having fluorescent tracer-negative RB sensory neurons (2.7 versus 1.5, respectively; $p = 0.02$, Mann-Whitney U-test; Figs. 3A-B). These data indicate that nav1.6a maintains DRG position non cell autonomously via activity in RB cells.

The above results predict that other manipulations that reduce sodium current in RB cells will also increase the number of migratory DRG neurons. The previously characterized *macho* (*mao*) mutant has reduced sodium current density in RB cells (Ribera and Nüsslein-Volhard, 1998). As predicted, 4 dpf *mao* mutants have an increased number of migratory DRG neurons compared to siblings (6.9 versus 3.6, respectively; $p < 0.0001$, Mann-Whitney U-test; Fig. 3C). These findings further support the hypothesis that maintenance of DRG position within the ganglia requires sodium current in RB neurons.

RB and DRG peripheral arbors overlap

Although RB and DRG somata reside in different locations, both neurons innervate the skin, implicating their cutaneous peripheral arbors as a possible site of interaction. RB neurons extend processes that innervate the skin around 1 dpf but the majority subsequently retract their processes and die before 5 dpf (Reyes et al., 2004; Williams et al., 2000). The timing of axogenesis varies for individual DRG neurons but begins by 2 dpf (An et al., 2002; Williams et al., 2000). Further, DRG neurons do not migrate ventrally in embryos injected with TTX or in 1.6 morphants until 2-3 dpf, stages when their axons have reached the periphery (Wright et al., 2010; data not shown). These considerations support the possibility of an activity-dependent interaction between DRG and RB peripheral processes. However, it is not known whether RB and DRG peripheral arbors overlap before the RB processes retract.

We determined whether DRG and RB peripheral processes overlap by expressing two different fluorescent proteins in their peripheral arbors. In *Tg(-3.4neurog1:GFP)* larvae, both RB and DRG processes express GFP, making it difficult to distinguish one from the other in the periphery. To reliably identify RB processes, we generated a transgenic construct that drives expression of another fluorescent protein (mCherry) in RB cells. We isolated a portion of the 5' upstream region of the *scn8aa* gene and used it to transgenically drive expression of mCherry (supplemental Fig. 1), because RB cells show particularly robust *scn8aa* expression (Novak et al., 2006). We identified 1.6 control and morphant *Tg(-3.4neurog1:GFP)* embryos expressing GFP in both RB and DRG neurons but mCherry in RB, but not DRG, neurons. For both 1.6 control and morphant embryos, tracings of individual DRG neuron processes indicated that many DRG processes reach the skin by 3 dpf (Figs. 4B and D). Further, in both controls and morphants, DRG processes in the skin overlapped with mCherry-expressing RB processes. These results further support the peripheral arbors as a potential site for activity-dependent interaction between RB and DRG neurons.

BDNF prevents ventral migration of DRG neurons

The finding that sodium channel activity as well as protein is required for maintenance of DRG position raises the possibility that the relevant signal is released from RB cells in an activity-dependent manner. Neurotrophins, which mediate nervous system development and function, are secreted in an activity-dependent manner and function in the skin (Wu et al., 2004; for reviews, see Bothwell, 1997; Shinder and Poo, 2000). Further, various aspects of RB and DRG development require NGF, NT-3, or BDNF (Hapner et al., 2006; Heinrich and Pagtakhan, 2004; Patel et al., 2000; Valdes-Sanchez et al.; Williams et al., 2000). Accordingly, we tested the ability of these neurotrophins to prevent migration of DRG neurons away from the ganglia in 1.6 morphants (Fig. 5). We injected NGF, NT-3 or BDNF into 1.6 morphant *Tg(-3.4neurog1:GFP)* embryos at either 16 hpf—when RB neurons are just starting to elaborate arbors, 28 hpf—when RB neurons have established peripheral arbors but DRG neurons have not yet differentiated, or 48 hpf—when DRG neurons are growing axons.

NGF and NT-3 did not significantly affect DRG migration, regardless of the time of injection (Fig. 5). In contrast, BDNF rescued the DRG phenotype of 1.6 morphants, consistent with RB cells signaling to DRG neurons through activity-dependent release of BDNF (Fig. 5A). Further, if BDNF were required prior to 48 hpf, the results from embryos injected at 16 or 28 hpf would have differed from those injected at 48 hpf. However, the effect of BDNF on DRG position was not significantly different between embryos injected at 16, 24, or 48 hpf. This finding suggests that DRGs require BDNF after 48 hpf. At 48 hpf, *nav1.6a* already functions in RB cells (Pineda et al., 2005) but DRG peripheral processes first reach the skin.

Neurotrophins also modulate and possibly activate ion channels (Lesser et al., 1997; Blum et al., 2002; Ahn et al., 2007; for review, see Luther and Birren, 2009), raising the alternative possibility that the effects of neurotrophin injection may be due to potentiation or activation of RB sodium current in morphants. However, this possibility is unlikely given that none of the neurotrophins tested rescued the touch response phenotype in 1.6 morphants, suggesting that *nav1.6a* sodium current remained impaired after neurotrophin treatment (supplemental Fig. 2A).

If DRG migration in 1.6 morphants resulted from decreased release of BDNF by RB cells, inhibiting BDNF actions by other means would be expected to induce migration of DRG neurons away from the ganglia in wildtype zebrafish. To test this, we injected 48 hpf wildtype Tg(-3.4neurog1:GFP) embryos with a function-blocking anti-BDNF antibody. The anti-BDNF antibody increased migration of DRG neurons in a dose-dependent manner (Fig. 5B). Similar to neurotrophin injection, the anti-BDNF antibody did not affect the touch response of 60 hpf embryos (supplemental Fig. 2B) making it unlikely that the anti-BDNF antibody affected DRG position by reducing sodium current in RB cells. Taken together, these data suggest that DRG neurons migrate away from the ganglia in 1.6 morphants due to a reduction in activity-dependent release of BDNF by RB neurons.

Decreased tactile stimulation increases migratory DRG neurons in wildtype zebrafish

Our studies indicate that genetic or pharmacologic reduction of sodium current in RB cells increases the frequency of migration of DRG neurons away from the ganglia. Such reductions in sodium current, however, were produced by experimental perturbations not likely to be experienced by wildtype embryos. RB cells are mechanosensory neurons that are electrically activated in response to tactile stimulation during early development (Clarke et al., 1984). To determine whether environmentally-induced variations in RB activity also suffice to modulate migration of DRG neurons, we raised wildtype embryos to 4 dpf under conditions of varying tactile stimulation.

Embryos raised under conditions of increased tactile stimulation did not vary from controls with respect to either touch response or the number of migratory DRG neurons (Figs. 6A-B). In sharp contrast, embryos raised under conditions that minimize tactile stimulation showed a variety of phenotypes. Approximately 20% of sensory-deprived embryos failed to hatch by 4 dpf, whereas wildtype embryos typically do by 2 dpf (data not shown). Further, embryos that hatched displayed a statistically significant reduction in touch sensitivity, compared to embryos that received tactile stimulation (Fig. 6A; average touch response score 6.4 versus 8.7, respectively; $p < 0.0001$, Mann-Whitney U-test). Moreover, the number of migratory DRGs increased in sensory-deprived embryos compared to embryos raised under control conditions (Fig. 6B; 4.0 versus 2.5 migratory DRGs, respectively; $p = 0.02$, Mann-Whitney U-test).

In *nav1.6* morphant or mutant embryos, migratory DRG neurons transdifferentiate into sympathetic neurons (Wright et al., 2010). To test whether migratory DRG neurons in

sensory-deprived larvae also transdifferentiate, we allowed them to develop to 11 dpf, a time when the sympathetic nervous system has differentiated. We then examined expression of HuA (a marker of both DRG and sympathetic neurons) and tyrosine hydroxylase (TH), which labels sympathetic neurons but not DRG neurons at the stages examined in this study (Wright et al., 2010). TH-expressing migratory DRG neurons were present in larvae raised under conditions of decreased tactile stimulation (Fig. 6C). These data suggest that maintenance of DRG position and identity requires early tactile stimulation.

Discussion

In many systems, target-derived neurotrophins serve to match the level of innervation with the size or need of the target tissue (reviewed in, Purves et al., 1988; for review, see Davies 1996). Our results suggest that neurotrophins also match the size of the sensory neuron pool to the level of early sensory stimulation. Under control conditions, activity-dependent release of BDNF from RB cells transduces the early sensory requirements of the embryo into a signal that maintains DRG neurons within the ganglia. In this way, BDNF regulates the number of sensory neurons that respond to touch.

We propose that sensory deprivation reduces activity-dependent release of BDNF by RB cells (Figs. 6, 7). We did not detect any obvious effects of BDNF on channel function as noted by others (e.g., supplemental Fig. 2A; Lesser et al., 1997; Blum et al., 2002; Ahn et al., 2007; for review, see Luther and Birren, 2009). In the absence of RB activity-dependent release of BDNF, differentiated DRG neurons migrate out of the ganglion to locations where they can transdifferentiate into, e.g., sympathetic ganglia neurons (Wright et al., 2010). Further, sensory deprivation not only increases DRG migration but also significantly reduces the ability to respond to touch at later times (Fig. 4), indicating an essential requirement for activation of early sensory neurons for later sensory function.

Although activity perturbations had statistically significant effects on the number of migratory DRG neurons present in larvae, the total increase in the number of such cells per embryo was not large (e.g., a few per embryo). This finding suggests that only a subset of DRG neurons respond to sensory deprivation/BDNF loss by migrating out of the ganglion. One possible explanation is that not all DRG neurons express a receptor for BDNF such as *trkB*. The zebrafish genome contains two different *trkB* genes (Martin et al., 1995). Towards the end of gastrulation, the zebrafish embryo begins to express mRNA encoding a full-length *trkB* isoform (Lum et al., 2001). Expression of *trkB* by zebrafish DRG neurons has not yet been examined. However, in mammals, a specific DRG subpopulation expresses mRNA for *trkB* (Wright and Snider, 1995). In addition, this subpopulation co-expresses mRNA for an alternate, albeit less selective, BDNF receptor, *p75* (Wright and Snider, 1995). Many other DRG neurons also express *p75*, indicating that *trkB*, in contrast to *p75*, expression identifies a smaller subpopulation of DRG neurons.

Previous studies have revealed the importance of BDNF for DRG neurons. For example, several studies point to a key role for BDNF as a target-derived survival factor for sensory neurons (e.g., Liebl et al., 1997; Valdes-Sanchez et al., 2010; but see Huber et al., 2000; for review, see Barde, 1994). Further, LeMaster et al. (1999) tested the effects of BDNF overexpression in the skin; cutaneous overexpression of BDNF regulated sensory innervation of the skin, revealing a role for this neurotrophin in morphological development. In addition, Carroll et al., (1998) found that slowly adapting mechanoreceptors require BDNF for proper development of receptive properties. Adding to this list, our studies reveal a new role for BDNF as a regulator of DRG neuron position and identity.

We found that RB cell activity plays a critical early role in establishment of the DRG sensory system. RB cells are an enigmatic population of sensory neurons that develops during the earliest stages of neurogenesis (Lamborghini, 1980; Metcalfe et al., 1990). For the most part, RB cells die during larval stages (Coen et al., 2001; Humphrey, 1950; Kollros and Bovbjerg, 1997; Lamborghini, 1987; Reyes et al., 2004; Svoboda et al., 2001; Williams et al., 2000). In this way, RB cells are reminiscent of another transient population, Cajal-Retzius cells, that play a critical role in layer formation in the cerebral cortex. In amphibia and teleosts, RB cells form the initial sensory system of the developing embryo and are responsible for touch-mediated reflexes (Clarke et al., 1984; Douglass et al., 2008). Although it is typically thought that mammals lack RB cells (but see, Humphrey, 1950), such a conclusion requires further study of embryonic stages when these cells typically are present in other vertebrates.

However, if mammals lack RBs, would the mechanism we report here be relevant to humans? The diverse DRG population in mammals arises during different waves of neurogenesis (Lawson and Biscoe, 1979) allowing the possibility of earlier born sensory neurons to regulate the fate of later-born ones (e.g., Ma et al., 1999). Further, studies in rats indicate that proper functioning of the adult nociceptive circuit requires tactile stimulation during the neonatal period (Fitzgerald, 2005; Petersson et al., 2004). Similarly, adult rats that did not receive sensory inputs from their whiskers during neonatal development perform poorly on tactile discrimination tasks (Carvell and Simons, 1996). Particularly compelling are studies indicating that children who lived in institutionalized environments with minimal sensory stimulation as infants show persisting defects in sensory discrimination and processing (Cermak and Daunhauer, 1997; Lin et al., 2005).

In our studies DRG neurons migrate away from the ganglion between 2- 4 dpf, following normal hatching stages. This finding supports the idea of a sensitive period during which differentiated DRG neurons are responsive to environmental cues that regulate DRG neuron number by maintenance or loss of position within the ganglion. It is currently unknown whether other sensitive periods exist when DRG neurons could be induced to migrate and transdifferentiate. However, several findings suggest the possibility of a sensitive period for DRG neuron maintenance following nerve injury. DRG neurons acquire BDNF dependency (Geremia et al.; Li et al., 2009; Sebert and Shooter, 1993) and upregulate the BDNF-responsive trkB receptor following peripheral nerve injury (Sebert and Shooter, 1993); BDNF upregulation within the DRG appears to be crucial for sensory neuron regeneration (Geremia et al.; Li et al., 2009). To date, it has not been tested whether DRG neurons that fail to receive trophic support following nerve injury migrate away from the ganglion and transdifferentiate into sympathetic neurons. However, novel sprouting of sympathetic fibers within the DRG occurs after nerve injury; the nerve sprouting is correlated with the generation of neuropathic pain (Chien et al., 2005; Chung and Chung, 2001; Kim et al., 1999; Shinder et al., 1999; Zhang and Strong, 2008). In zebrafish, migratory DRG neurons retain their dorsal axons upon transdifferentiating into sympathetic neurons (Wright et al., 2010), suggesting a possible source for sympathetic sprouting within the DRG.

The present work identifies BDNF as a factor that maintains the position and identity of differentiated DRG neurons. BDNF mediates the effects of activity and environmental conditions, allowing early perturbations to sculpt the performance of the adult sensory system. These findings emphasize the importance of sensory input during developmental stages.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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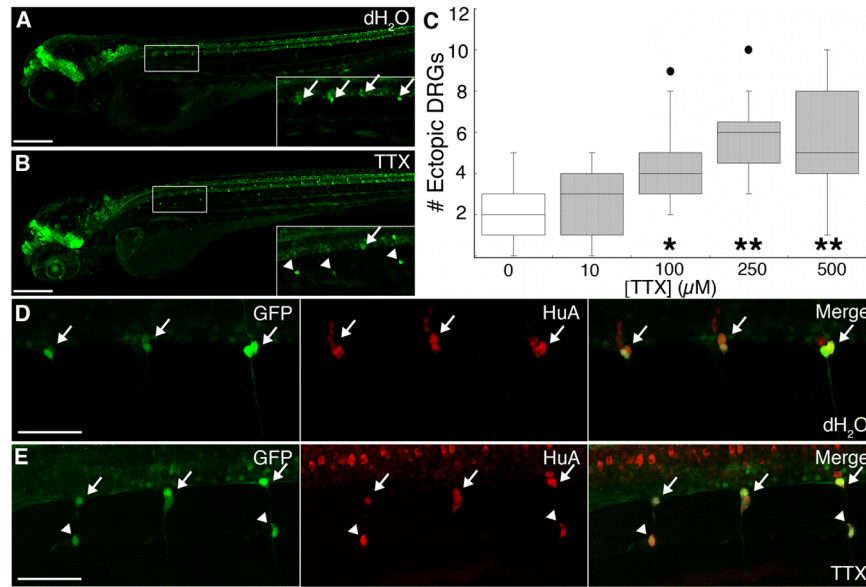


Figure 1. Maintenance of DRG position requires sodium current. **A**, Whereas DRG neurons (arrows) are regularly arranged along the ventral boundary of the spinal cord in 4 dpf Tg(-3.4neurog1:GFP) dH₂O-injected embryos. **B**, DRG cells are absent from this location in several segments and GFP-expressing cells are instead located in ectopic ventral positions (arrowheads) in sibling embryos injected with 250 μM TTX. Insets correspond to the boxed regions in panels **A** and **B**. **C**, The median number of ectopic DRG neurons per embryo at 4 dpf increases in a dose-dependent manner with TTX concentration (*, $p < 0.05$ versus 100; **, $p < 0.001$ versus 250, and 500 μM TTX; nonparametric Kruskal-Wallis test; sample sizes ranged between 15-22 embryos). The graph presents the inner quartiles as a box with an internal line indicating the median; whiskers extend to the 5th and 95th percentiles; filled circles indicate outliers. **D**, **E**, In 4 dpf Tg(-3.4neurog1:GFP) embryos, both normally-positioned as well as ectopic GFP-expressing DRG cells (arrows and arrowheads, respectively) express HuA (red), a marker of neuronal differentiation. Scale bars, 200 μm , **A** and **B**; 50 μm , **D** and **E**.

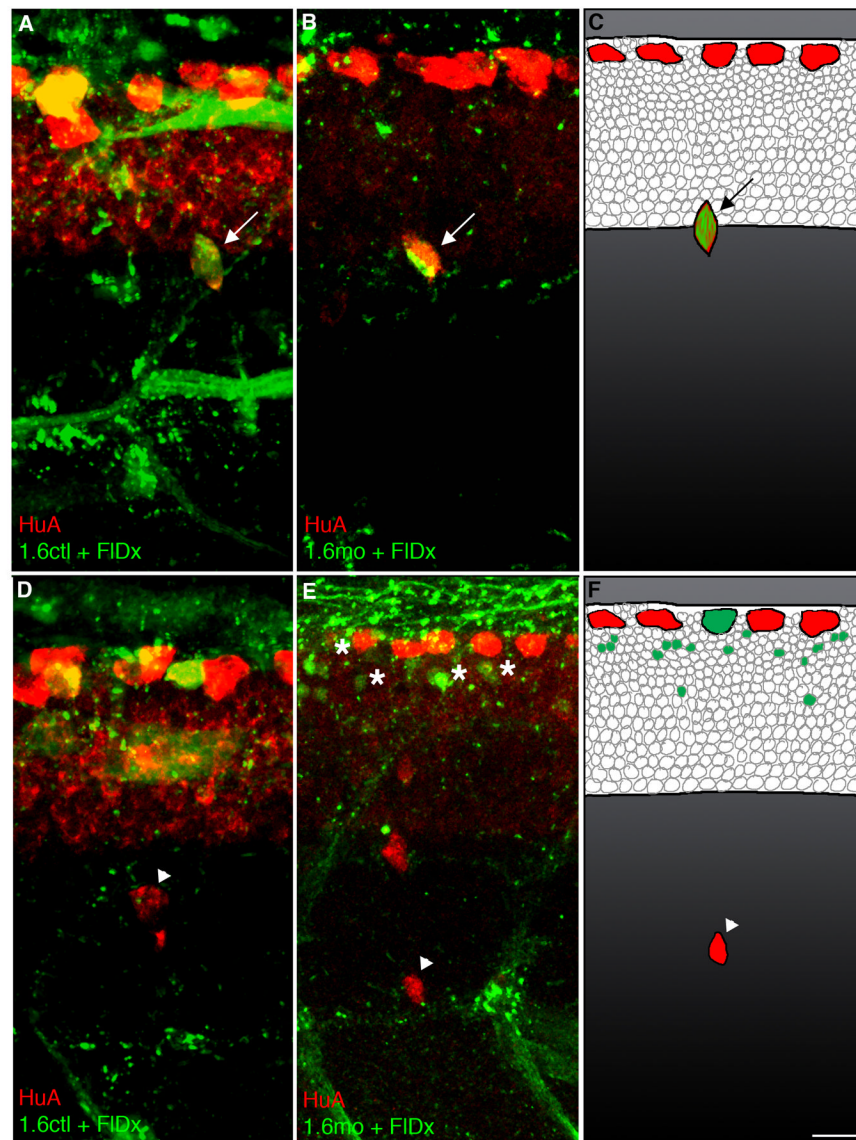


Figure 2. *nav1.6a* does not cell autonomously regulate maintenance of DRG position. **A,B,C**, In 3 dpf embryos, the presence of 1.6ctl (**A**) or 1.6mo (**B,C**) in DRG neurons does not affect DRG position; HuA immunoreactivity identified DRG neurons (red, arrows). FIDx (green) was co-injected with morpholino to serve as a lineage tracer. **D**, The rare ectopic DRG neuron (arrowhead) present in mosaic 1.6ctl embryos does not contain the FIDx. **E,F**, Similarly, even though there is an increase in the number of ectopic DRG neurons in mosaic 1.6mo embryos, the mo and tracer (green) are not found within the ectopic DRGs (arrowheads). Instead, the tracer is present in other spinal cord cells (asterisks). HuA immunoreactivity is normally present in dorsal RB sensory neurons, as shown in all panels. Scale bar, 10 μ m.

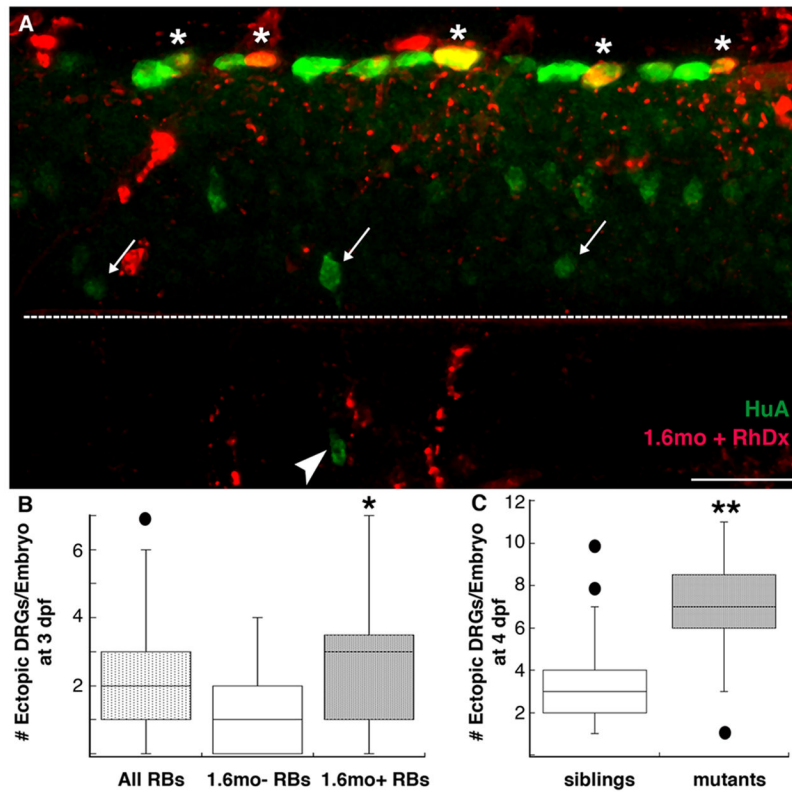


Figure 3.

Maintenance of DRG position/identity requires *nav1.6a* in RB cells. **A,B**, Ectopic DRG neurons (arrowhead), identified by HuA immunoreactivity (green), were present in mosaic embryos that contained RB cells (asterisks) with 1.6mo+ RB cells (red; yellow when co-present with HuA green signal). Arrows denote normally-positioned DRG neurons; the dashed white line indicates the spinal cord-notochord boundary. **B**, The number of ectopic DRG neurons increased in mosaic embryos that contained 1.6mo in RB cells compared to mosaic embryos that did not contain 1.6mo in RB cells (* $p=0.02$, Mann-Whitney U-test; $n=44$ embryos containing 1.6mo in RB cells, 22 embryos not containing 1.6mo in RB cells). **C**, *mao* mutants, characterized by reduced sodium current densities in RB cells (Ribera and Nüsslein-Volhard, 1998), have an increased number of ectopic DRG neurons versus sibling control embryos at 4 dpf (** $p<0.0001$ versus *mao* siblings, Mann-Whitney U-test; $n=43$ *mao* mutant and 99 sibling embryos). Scale bar, 25 μm .

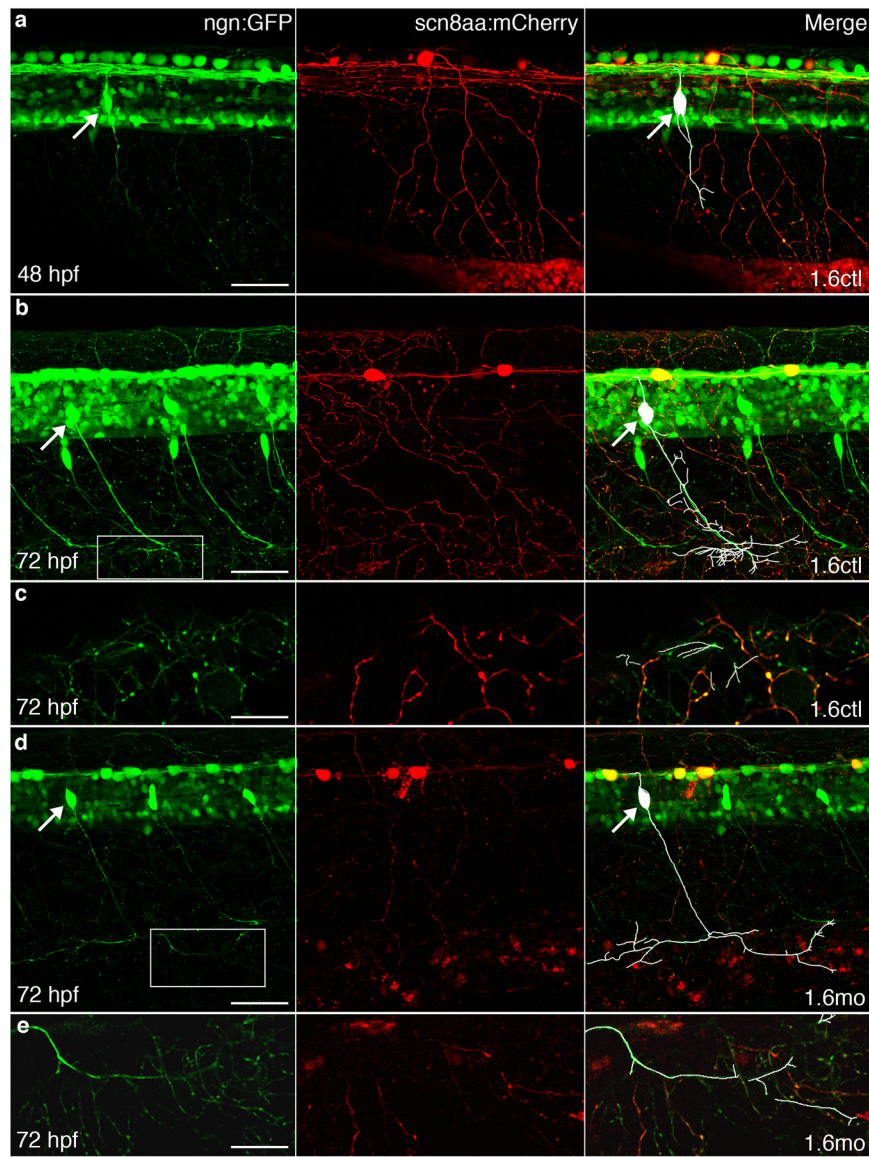


Figure 4. RB and DRG peripheral arbors overlap by 72 hpf in 1.6 control and morphant embryos. In all panels, GFP (green) in $Tg(-3.4neurog1:GFP)$ embryos labels both RB and DRG axons and cell bodies, whereas stochastic labeling with mCherry allows visualization of peripheral arbors (red) of individual RB cells. In the left merged panels, white traces the processes of the DRG neuron. White arrows indicate cell bodies of DRG neurons. **A**, In 48 hpf embryos injected with 1.6ctl, DRG axons that run next to the notochord have not yet reached the periphery and do not overlap with cutaneous RB peripheral processes. **B,C,D,E**, By 72 hpf, in embryos injected with either 1.6ctl (**B,C**) or 1.6mo (**D,E**), DRG neurons extend processes into the skin where they overlap with RB cutaneous processes. Panels **C** and **E** show 8 μ m confocal slices of the regions boxed in **B** and **D**, respectively. Scale bars, 50 μ m: **A**, **B** and **D**; 20 μ m: **C**, **E**.

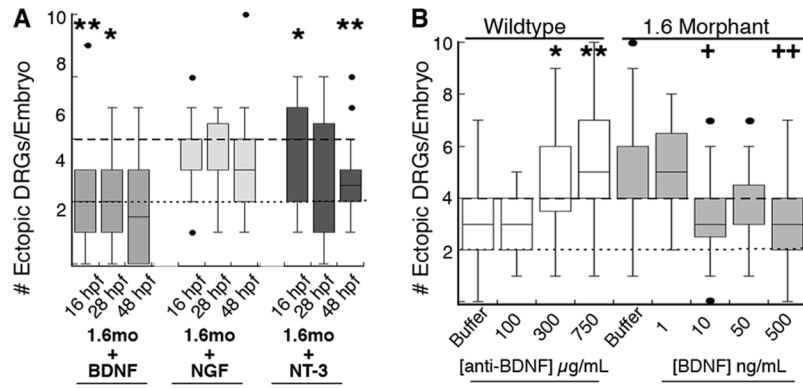


Figure 5.

Maintenance of DRG identity requires BDNF. **A**, Injection of NGF or (50 ng/mL) does not prevent the increase in migratory DRG neurons present in nav1.6 morphants (dashed grey line). In contrast, injection of BDNF or NT-3 reduces the number of migratory DRG neurons in nav1.6 morphants to levels found in wild type embryos (dashed black line). * and **, $p < 0.05$ and $p < 0.01$, respectively, versus nav1.6 morphants without neurotrophin injection; Kruskal-Wallis test; sample size ranged between 28 and 63 embryos. **B**, Anti-BDNF injected at 48 hpf (light blue bars) increases the number of migratory DRG neurons in wildtype embryos. Conversely, injection of 10-500 ng/mL BDNF reduces the number of migratory DRG neurons in nav1.6 morphants. * and **, $p < 0.05$ and $p < 0.01$, respectively, versus wildtype embryos injected with buffer only; + and **, $p < 0.05$ and $p < 0.01$, respectively, versus nav1.6 morphants injected with buffer only; non-parametric Kruskal-Wallis test; sample size ranged between 13 and 34 embryos.

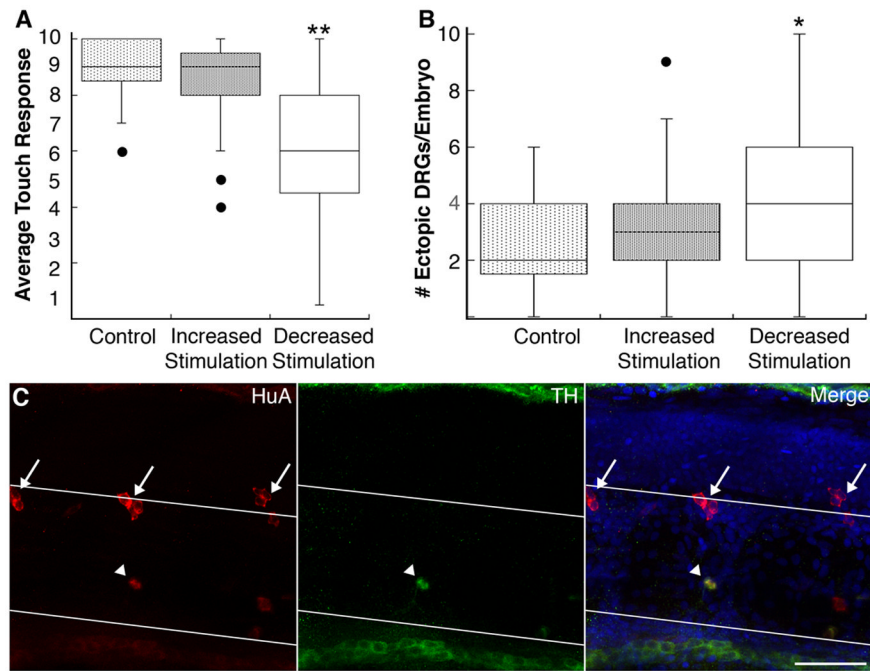


Figure 6.

Decreased tactile stimulation leads to ventral migration and transdifferentiation of DRG neurons. **A**, 4 dpf embryos raised under conditions that decrease tactile stimulation show a reduced touch response compared to embryos raised under control or increased stimulation conditions (** $p < 0.001$ versus control and increased stimulation, Kruskal-Wallis; sample size ranged between 10 and 65 embryos). **B**, Decreased tactile stimulation increases the number of ectopic DRG neurons at 4 dpf ($*p = 0.02$ versus control, Mann-Whitney U-test; sample size ranged between 20 and 50 embryos). **C**, At 11 dpf, a subset of migratory DRG neurons (arrowhead) expresses the sympathetic marker TH (green). HuA immunoreactivity is shown in red; white lines demarcate the dorsal and ventral boundaries of the notochord; arrows indicate normally-positioned DRG neurons. The fraction of migratory DRG neurons expressing TH does not differ significantly between control (9/35; 14 embryos), increased stimulation (17/58; 16 embryos) and sensory deprived embryos (16/47; 13 embryos). Scale bar, 40 μm .

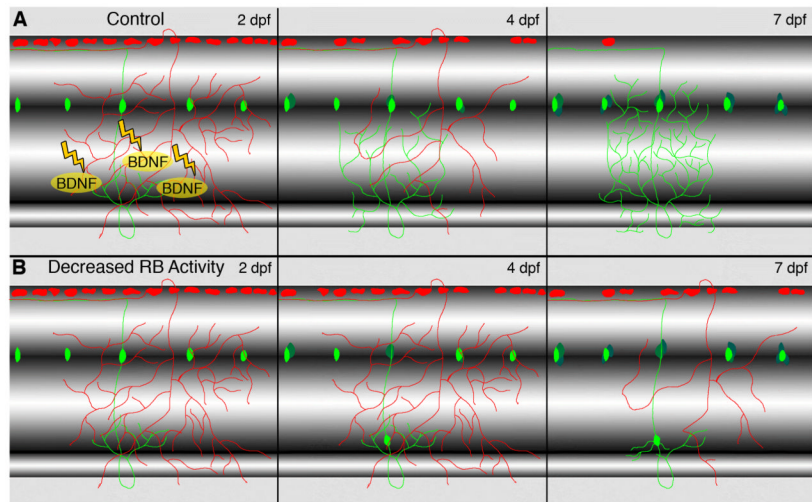


Figure 7.

Model for non cell-autonomous and activity-dependent regulation of DRG position and identity. **A**, At 2-3 dpf, RB (red) and DRG (green) peripheral processes overlap in the skin. Electrical stimulation in RB cells increases release of BDNF, which promotes maintenance of DRG neurons. Between 2 and 7 dpf, RB cells undergo cell death, while the number of DRG neurons increases and DRG peripheral processes elaborate. **B**, When RB activity is reduced genetically, pharmacologically, or environmentally, BDNF release decreases and RB cells survive to later stages. The absence of BDNF triggers migration of differentiated DRG neurons between 2 and 4 dpf. At later stages (7 dpf), migratory DRG neurons adopt sympathetic neuron characteristics.