The Role of \textit{inab} in Axon Morphology of an Identified Zebrafish Motoneuron

Liesl Van Ryswyk*, Levi Simonson, Judith S. Eisen*

Institute of Neuroscience, Eugene, Oregon, United States of America

Abstract
The ability of an animal to move and to interact with its environment requires that motoneurons correctly innervate specific muscles. Although many genes that regulate motoneuron development have been identified, our understanding of motor axon branching remains incomplete. We used transcriptional expression profiling to identify potential candidate genes involved in development of zebrafish identified motoneurons. Here we focus on \textit{inab}, an intermediate filament encoding gene dynamically expressed in a subset of motoneurons as well as in an identified interneuron. We show that \textit{inab} is necessary for proper axon morphology of a specific motoneuron subtype.

Introduction
For correct locomotor circuitry to form in a developing embryo, motoneuron axons must contact appropriate muscle targets. Motoneurons are classified into subtypes that are based on their axon projections and morphologies, a hallmark of appropriate motoneuron subtype specification. Correct motoneuron development is mediated by expression of genes that allow motoneurons to acquire subtype-specific characteristics, such as axon morphology. Although a number of genes have been identified with necessary roles in motoneuron development, we still have an incomplete picture of the genes required for motoneuron differentiation and axon morphology.

The zebrafish spinal cord is an ideal model in which to study questions of early neuronal development, because there are a small number of individually identifiable neurons that can be observed in live animals over the course of development [1]. The earliest-developing motoneurons in the zebrafish spinal cord are referred to as primary motoneurons (PMNs); there are also later-developing motoneurons referred to as secondary motoneurons [2]. PMNs are especially amenable to study, because they have three distinct subtypes. Each of these subtypes projects an axon to a subtype-specific region in the overlying muscle [2,3] and expresses a number of genes differentially [4,5,6]. Not only can the mechanisms of PMN subtype specification be addressed genetically in zebrafish, but the genes known to be involved in zebrafish motoneuron specification are conserved across vertebrates.

To uncover additional genes involved in motoneuron development, including both differentiation and acquisition of axon morphology, we performed a microarray screen. By comparing the transcriptome of embryonic zebrafish spinal cords manipulated to have supernumerary motoneurons to that of spinal cords manipulated to have fewer motoneurons, we were able to select a number of candidate genes. These candidates were up-regulated or down-regulated in the same direction in each condition as genes already known to be expressed in zebrafish motoneurons. One of these candidate genes is \textit{interneuron neuronal intermediate filament alpha b} (\textit{inab}) that encodes a neuronal intermediate filament. We chose to focus on this gene because of our interest in genes that potentially regulate motoneuron differentiation and axon morphology.

Neuronal intermediate filaments are a family of proteins thought to be involved in neuronal differentiation as well as acquisition of axonal morphology [7]. Upregulation of a number of neuronal intermediate filament proteins has been correlated with alterations in cellular morphology that accompany differentiation [8,9]. The genes that encode some intermediate filament proteins, such as \textit{nestin} and \textit{ina}, are expressed in mammalian neuroblasts at the end of migration and beginning of differentiation [10,11,12,13]. \textit{nestin} is a well-characterized marker of neural progenitors [10,14] and both \textit{nestin} and \textit{ina} are thought to be involved in differentiation of neural progenitor cells and the accompanying acquisition of proper axonal morphology [7,9,15,16,17].

The goldfish and frog \textit{ina} homologues are required for axon outgrowth and are upregulated in retinal ganglion cell axons after optic nerve crush [18,19,20]. Interestingly, different intermediate filament genes are expressed in different subpopulations of neurons as they acquire distinct morphologies [9], suggesting that intermediate filaments could be important for neuronal subtype specification.

In addition to its well-characterized role in the optic nerve, \textit{ina} is expressed in spinal motoneurons of goldfish and frog, although its role in cells other than retinal ganglion cells has not yet been elucidated [18,21]. Zebrafish have two homologs of \textit{ina} – \textit{inab} and \textit{inab} – of which \textit{inab} has been shown to be expressed in
motoneurons [19,22]. Its presence there raises the possibility that it could be involved in aspects of motoneuron development.

Here we investigate which zebrafish PMN subtypes express inab. Zebrafish have three PMN subtypes, all derived from the progenitor of motoneuron (pMN) domain: CaP, VaP, MiP, and RoP [23]. CaP and VaP are initially equivalent MNs, referred to as CaP/VaP, that go on to acquire distinct CaP and VaP fates [24,25,26,27]. CaP is present in all spinal hemisegments, and can be identified by a long, ventrally-extending axon. VaP is present in only approximately half of the spinal hemisegments, projects a short, ventrally-extending axon, and typically dies during embryonic development [24]. Both MiP and RoP are present in all spinal hemisegments. MiP can be identified by a dorsally-projecting axon, whereas RoP has a ventrally-projecting axon that arborizes more laterally than the CaP axon [2].

We also focus on an identified interneuron that arises from the pMN domain and can be a sibling to motoneurons, VeLD [23]. The VeLD cell body is located just dorsal to the PMNs [28]. VeLD can be identified by its ventral axon that descends ipsilaterally and projects caudally for many segment lengths [29,30].

We demonstrate that inab is expressed only in the CaP/VaP MN subtype. Surprisingly, VeLD interneurons also express inab. VeLDs and PMNs are derived from the same progenitor domain, can be siblings, and have been shown to express some of the same genes [6,23,31]. inab appears unnecessary for motoneuron or interneuron specification, but the axon morphology of VaP MNs is disrupted when inab is misspliced – VaP axons have an increased number of processes – however the number of processes on other PMN axons is not influenced by missplicing of inab. This work suggests that neuronal intermediate filament proteins are involved in acquisition of axon morphology and are often differentially expressed in neuronal subtypes.

Materials and Methods

Zebrafish

Wild-type (AB), smu^641 [32], Tg(nrp1a:GFP)^212 [33], Tg(nsn1:GFP)^212 [34], Tg(vsx1:GFP) [35], and Tg(mnx1:GAL4VP16)^125 [36]; Tg(UAS;cd8Tomato;ze1:EGFP)^6 [36] zebrafish were maintained in a laboratory breeding colony according to established protocols [36]. Embryos collected from natural crosses were allowed to develop at 28.5°C and staged according to morphological criteria and hours postfertilization (hpf) [37].

Ethics statement

This study was carried out in strict accordance with the recommendations from the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the University of Oregon Institutional Animal Care and Use Committee (11-20). All embryonic dissociations were performed under MS-222 anesthesia, and every effort was made to minimize suffering.

RNA synthesis and injections

Capped dominant negative ribpa (formerly Suppressor of Hairless) RNA was synthesized and injected as previously described [38]. This was sufficient to generate ectopic motoneurons as assayed by in situ hybridization with islet2a probe [31].

Full-length inab RNA was transcribed using the mMessage Machine kit (Ambion/Life Technologies, Grand Island, NY, USA) according to instructions. One-cell stage embryos were injected with 400 pg inab RNA for rescue experiments.

Spinal cord dissociations

Our embryonic dissociation protocol was adapted from one previously used in the lab [39]. Embryos were grown to 20 hpf and anesthetized using Finquel (MS-222) (Argent Laboratories, Redmond, WA, USA). To obtain isolated spinal cords, embryos were incubated in 7.5 × pancreatin (MP Biomedicals, Irvine, CA, USA) in zebrafish physiological saline [36] until tissues began to separate (about 1 minute). Embryos were then triturated with Pasteur pipettes of decreasing size. Isolated spinal cords were transferred to Leibowitz’s L15 medium (Gibco/Life Technologies, Grand Island, NY, USA) containing 10% fetal bovine serum for 1–2 minutes to inactivate the pancreatin, after which they were stored in TriReagent (Molecular Research Center, Inc., Cincinnati, OH, USA) at −80°C.

Microarray sample preparation

Isolated spinal cords in TriReagent were pooled and homogenized. RNA extraction was performed according to standard protocols [40]. The time investment required for tissue collection prohibited analysis of biological or technical replicates. However, all genes of interest were validated as described in the Results. Extracted RNA was amplified using a MessageAmp II aRNA kit (Ambion/Life Technologies, Grand Island, NY, USA) according to supplied instructions. Amplified RNA was shipped to NimbleGen (Roche NimbleGen Inc., Madison, WI, USA) where cDNA probes were synthesized and hybridized to Zebrafish Gene Expression 385K Arrays.

Microarray data analysis

Data analysis was carried out using ArrayStar software (DNASTAR Inc., Madison, WI, USA). Candidate genes were selected by relative transcript abundance across conditions and gene expression pattern criteria, and verified by RT-PCR and in situ hybridization.

RNA probe generation

Full-length cDNA sequences containing the genes of interest were obtained from Open Biosystems (Thermo Fischer Scientific, Lafayette, CO, USA). ccdh55al cDNA (Clone ID: 7035729) was amplified using the following primers: forward 5’-TGTACG-GAAGTTGTTACCTTCGGTC-3’ and reverse 5’-GAGTACCTATACCGAAGGCGCCGACCTGACGTCG-3’. inaa cDNA (Clone ID: 6968318) was amplified using the following primers: forward 5’-ACAGTCTATCGAGATTAGCAGTCG-3’ and reverse 5’-GTAAACCGGACGGCCGATG-3’. inaa cDNA was amplified using the following primers: forward 5’-CAGGTCTCACTGTCTGTGG-3’ and reverse 5’-TGACAATCGTCCTCTCCGC-3’. nr2f1b cDNA (Clone ID: 7149507) was amplified using the following primers: forward 5’-AACAGTCATGACATGATACACGCT-3’ and reverse 5’-GTTAAACCGGACGCGATTG-3’. inab cDNA was amplified using the following primers: forward 5’-CAGGTCTCACTGTCTGTGG-3’ and reverse 5’-TGACAATCGTCCTCTCCGC-3’. For all genes, T3 RNA Polymerase (Roche Applied Sciences, Indianapolis, IN, USA) was used to make probe for RNA in situ hybridization according to standard protocols [41].

Fluorescent RNA in situ hybridization

RNA in situ hybridization was performed according to standard protocols [41], with the following modifications: For 2-color fluorescent in situ hybridization, anti-sense probes were labeled with digoxigenin-UTP (Roche Applied Sciences, Indianapolis, IN, USA) and dinitrophenol-UTP (Perkin-Elmer, Waltham, MA, USA). Following overnight hybridization, unbound probe was removed with three 30-minute washes at 67°C in 50% formamide.
5× SSC, and 0.1% SDS, followed by stringent washes in 50% formamide, 2× SSC, and 0.1% Tween-20. Labeled probes were detected with HRP-conjugated anti-DIG (1:1000; Jackson ImmunoResearch, West Grove, PA, USA) or HRP-conjugated anti-DIG (1:1000; Perkin-Elmer), and stained with fluorescein or Cy-3-tyramide (1:50; Perkin-Elmer) for 1-10 minutes.

Probes used include islet1 and islet2a [31]; gad1b and gad2 (collectively referred to as gad, skl17a6a, skl17a6b, and skl17a7 (collectively referred to as glda, slc6a9 and slc6a5 (collectively referred to as gfs) [42]; vsx2 [43]; and ced65al, w2f1b, inn, and inab.

Immunohistochemistry

Embryos were fixed overnight in 4% paraformaldehyde and 1x Fix Buffer at 4°C. Embryos were then block in 5% normal goat serum, 2.5% DMSO, and 0.1% Tween-20 in 1× PBS before overnight incubation in diluted primary antibody at 4°C. Unbound primary antibodies were removed by washing for 2 hours in 1× PBS plus 0.1% Tween-20, followed by overnight incubation in diluted secondary antibody at 4°C.

Antibodies used include monoclonal mouse anti-GFP 3E6 (1:1000; A11120, Molecular Probes/Life Technologies, Eugene, OR, USA); rabbit polyclonal anti-Mnx1 and anti-Mnx2b [6]; rabbit polyclonal anti-Mnx1 (1:1000), anti-Mnx2a (1:1000), and anti-Mnx2b (1:1000; AnaSpec, Fremont, CA, USA); rabbit polyclonal anti-Mnx1 and anti-Mnx2b [6]; and Alexa-Fluor dye-labeled secondary antibodies (Molecular Probes/Life Technologies).

Morpholino injections

Approximately 2 nL of a combination of 300 μM SB2: 5’-GGAATCTAGTAGCTGT-GATAATTCC-3’ and 300 μM SB3: 5’-CAGTGATGTTTAT-TACCTGTAAGC-3’ were injected into 1 to 2-cell stage embryos. This was sufficient to cause complete missplicing as assayed by primers designed to flank the splicing sites (forward: 5’- CCTGGAGAAAAAGTGCTGATCC-3’ and reverse: 5’- GCTATTTTCTATGTCAAGCGCC-3’ (Figure 1B-C). Using these primers, the wild-type band is 584 bp and the SB MO band is 459 bp, indicating a complete removal of the 125 bp second exon. An additional splice-blocking MO against inab, SBI (5’-ATGAAAACCTGAAAACCAACTGTGTT-3’) was tested, although it did not cause complete missplicing at any concentration and thus was not used in this study.

Other previously validated morpholinos used in this study include: 2.5 nL of 100 μM mnx1 (5’-ACCCTCACAACAAGAT-TAACCAGCTCG-3’), mnx2a (5’-ACCCCAACAACAGAT-TAACGCTCGTCCG-3’), and mnx2b (5’-GACCTTTTCGTTGCAA-CACCTTTTGTGTT-3’) to knock down mnx genes [6]. Morpholino effectiveness was verified by whole-mount immunohistochemistry using anti-Mnx1, anti-Mnx2a, and anti-Mnx2b antibodies (see Immunohistochemistry).

DNA Injections

To visualize individual neurons in the trunk, approximately 3 nL of 10 ng/μL UAS:EGFP 434×193 plasmid (prepared using multisite Gateway technology [44]) was injected into 1 to 2-cell stage Tg(mnx1:GAL4) embryos. We selected embryos with GFP-expressing cells for immunohistochemistry.

Image acquisition

All images were acquired on a Zeiss Pascal confocal microscope (Carl Zeiss Microscopy, LLC, Thornwood, New York, USA) using either a 20× objective or a 40× water-immersion objective. The brightness and contrast of images was adjusted using Photoshop CS5 (Version 12.0, Adobe Systems, Inc., San Jose, CA, USA).

Image Quantification

All observations of PMNs were made in mid-trunk spinal cord adjacent to somites 8–12.

Image analysis was performed using a previously described pipeline [45]. First, we used the NeuroJ] plugin [46] [NeuroJ: http://www.imagescience.org/meijering/software/neuroj/] for ImageJ [NIH, Bethesda, MD; ImageJ: http://rsb.info.nih.gov/ij/] to define the positions of neurites within raw confocal images. We used this neurite location data and the neuronStudio software [47] (NeuronStudio: http://research.mssm.edu/cnic/tools-ns.html) to construct a connectivity diagram of each neuron, such that each branching neurite may only give rise to two daughter neurites. We examined these data, corresponding to the complete two-dimensional arbor of each cell, using the MATLAB-based automated Sholl analysis program - Bonfire [45]. Sholl analysis produces a matrix of values corresponding to neurite length, number of branch points, and number of terminal points, as a function of radial distance from the soma. Images were scored blind, and resulting data were transferred to Excel for statistical analysis. Significance was determined using a two-tailed Student’s T-test, with a p-value of less than 0.05 being considered significant. We also performed a noncentral t-distribution cumulative distribution function test which provides the minimum total sample size necessary to determine whether two samples are different from one another [48]. According to this test, a sample size of 10 provides a reliable measure for our data. We examined 14 VaPs, 11 CaPs with VaPs, 11 CaPs without VaPs, and 10 MiPs.

![Figure 1. Splice-blocking MOs effectively disrupt inab. (A) Schematic of the inab gene. inab has three exons and two introns. Splice-blocking (SB) MOs target various splice sites in the gene. SB2 targets the intron1/exon2 boundary, and SB3 targets the exon2/intron2 boundary. Forward (Fw) and reverse (Rv) primers sit in the first and third exons, respectively. (B) PCR results confirming that uninjected embryos have wild-type inab (584 bp band), embryos injected with 2 nL of a combination of 31 μM SB2 and 31 μM SB3 MOs, or 2 nL of a combination of 125 μM SB2 and 125 μM SB3 MOs have both wild-type and misspliced inab, and embryos injected with 2 nL of a combination of 500 μM SB2 and 500 μM SB3 MOs have only misspliced inab (asterisk, 459 bp band). (C) PCR results confirming that uninjected embryos have wild-type inab (584 bp band), and embryos injected with 2 nL of a combination of 300 μM SB2 and 300 μM SB3 also have only misspliced inab (asterisk, 459 bp band). doi:10.1371/journal.pone.0088631.g001]
Results

Expression profiling of zebrafish spinal cord using NimbleGen microarrays

We performed a microarray screen to uncover additional genes that are expressed in zebrafish PMNs and thus potentially involved in motoneuron development. For the screen, we compared spinal cords of embryonic zebrafish with supernumerary motoneurons to spinal cords with fewer motoneurons. To create embryos with more motoneurons than normal, we injected wild-type fish with mRNA encoding a dominant negative Rbpja protein (dnrbpja). This dominant negative protein interferes with Delta/Notch signaling in the embryo, thus creating supernumerary early-born neurons and increasing the number of PMNs [38]. On average, the number of islet2 PMNs per hemisegment increases from 1–2 in control embryos to 3–6 in dnrbpja RNA-injected embryos [49]. To create embryos with fewer than normal motoneurons, we injected fish of the smoothened mutant line, smub641. The Smoothened protein plays a crucial role in Hedgehog signaling, which is required for normal motoneuron development [50]. Homozygous smub641 mutants have disrupted Hedgehog signaling and consequently almost completely lack motoneurons, having on average 18 islet2 PMNs per embryo [32,50].

To cut down on background from other tissues, we separated spinal cords from the rest of the tissues in the 20 hpf embryonic fish using our previous dissociation procedure [39]. After collecting a sufficient number of spinal cords (~200 per condition), we extracted RNA from each sample and amplified it. To confirm that spinal cords were present in each case, we used tissue-specific primers to verify presence of spinal cord genes and absence of contaminating hindbrain and muscle genes (Figure S1). The amplified RNA was delivered to NimbleGen, where it was hybridized to microarrays.

We compared 10 genes known to be involved in motoneuron development – islet1, nx6.1, nx6.2, mnx1, mnx2b, met, scd1a, dla, neurog1, and nfasca – and found that they were approximately 1.5 times upregulated in the supernumerary motoneuron condition as compared to uninjected, wild-type control embryos, as well as approximately 1.5 times downregulated in the decreased motoneuron condition as compared to wild-type control embryos. We then selected genes that were at least 1.5 times upregulated in the supernumerary motoneuron condition and 1.5 times downregulated in the decreased motoneuron condition, giving a list of 740 candidates.

We narrowed the list down by excluding “housekeeping” and hypothetical genes, giving a list of 101 candidate genes (Table S1). After excluding those genes expressed outside of the central nervous system and those initially expressed earlier than the birth of MNs or significantly after their maturation, we ended up with a list of 11 candidate genes (Table S1). To verify and further investigate expression patterns of the candidate genes, we obtained a clone of each gene from Open Biosystems and synthesized RNA probes for each of them. Here we provide information on three candidate genes that are expressed in ventral spinal cord.

One of the candidate genes, cede85al, is predicted to encode a transcription factor, and although it is expressed in the spinal cord, it is not expressed in PMNs and was therefore excluded from our analysis (Figure S2). Another candidate, m2f1b, is a transcription factor-encoding gene that is expressed in MNs, although it is also broadly expressed throughout the spinal cord (Figure S2). Because its expression pattern is so broad, we excluded m2f1b from our analysis as well. Here we describe expression and functional analysis of the third candidate gene, the intermediate filament-encoding gene inab, which we found to be expressed specifically in PMNs.

inab is dynamically expressed in a subset of primary motoneurons and in VeLD interneurons

We characterized expression of inab in the zebrafish spinal cord using RNA in situ hybridization. To determine whether PMNs express inab, we labeled embryos for islet1 mRNA, which all PMNs express before 14 hpf [31], and inab. In embryos older than 14 hpf, we labeled with probes for both islet1 mRNA (which after 14 hpf is expressed in PMN subtypes MiP and RoP) and inab, or islet2a mRNA (which after 14 hpf is expressed in PMN subtype CaP/VaP) [31] and inab (Figure 2A–H). inab mRNA was not detected before 14 hpf. inab is expressed in CaP and VaP between 16 and 24 hpf (Figure 2F–H) and is subsequently downregulated in these cells. We never saw coexpression of inab and islet1, indicating that inab is not expressed in either MiP or RoP.

In addition to its expression in a subset of PMNs, inab is also expressed in cells that are more dorsal than the PMNs. To determine the identity of these cells, we labeled transgenic mnx1:GFP embryos with inab riboprobe. The mnx1:GFP transgene is expressed in all PMN subtypes and in VeLD interneurons [INs] before 24 hpf [6]. VeLD can also be identified by expression of the neurotransmitter GABA [6,51,52]. The inab IN expresses both the mnx1 transgene and is labeled by an mRNA probe against gads, that encode the GABA synthetic enzymes, leading us to conclude that it is VeLD (Figure 3F–J). inab expression in VeLD is initiated at 14 hpf and persists through 24 hpf.

inab is also expressed in a more dorsal, glutamatergic population of interneurons, with expression initiating around 18 hpf (Figure 2K). Because we are interested in genes regulating motoneuron development, we excluded these neurons from our analysis.

There is a second zebrafish gene, called inaa, that is closely related to inab (http://zfin.org/action/Marker/view/ZDB-GENE-060531-65). In the zebrafish genome, inaa is located on chromosome Drc13, whereas inab is found on chromosome Drc1. Teleost orthologs of inaa and inab are found on chromosomes Ola15 and Olal in medaka, on Tetraodon chromosomes Tnl17 and Tnl18, and in stickleback on chromosomes GacVI and GacIX, respectively. All these pairs of chromosomes contain chromosomal blocks duplicated during the course of the teleost genome duplication (TGD) derived from ancestral pre-TGC chromosome α. This strongly suggests that inaa and inab are paralogs of each other generated during the TGD as well [53,54]. We characterized expression of inaa using RNA in situ hybridization. We found that inaa is not expressed in the spinal cord between 14–24 hpf (Figure 3A–D), and instead is expressed in only the diencephalon or adjacent to the diencephalic ventricle at 24 hpf (Figure 3E). We therefore excluded inaa from our analysis.

inab is expressed in hybrid cells generated by mnx gene knockdown

We recently showed that in the absence of the three zebrafish Mnx homologs (Mnx1, Mnx2a, and Mnx2b), some MiP motoneurons become MiP-CaP hybrids that develop CaP-like axons [6]. MiPs do not normally express inab (Figure 2), but we hypothesized that some MiPs would express inab in the absence of Mnx proteins. To test this hypothesis, we injected previously-validated [6], translation-blocking MOs against all three mnx genes into embryos of the Tg(mnp1a:GFP) line, in which MiPs weakly express GFP after 20 hpf. In MO-injected embryos, 32% of GFP+ MiPs expressed inab (n = 16/50 MiPs, 10 embryos; Figure 4B), in contrast to 6% of GFP+ MiPs in control embryos (n = 3/50 MiPs, 10 embryos; Figure 4A). This indicates that inab expression can be
acquired by MiP-CaP hybrids that are generated in the absence of Mnx proteins, providing another molecular marker of CaP fate.

**inab** is required for proper morphology of VaP axons

To test the function of **inab** in PMN development, we used splice-blocking (SB) MOs to prevent proper splicing of **inab** mRNA. We designed MOs to target the exon 1/intron 1 boundary (SB1), the intron 1/exon 2 boundary (SB2), and the exon 2/intron 2 boundary (SB3) (Figure 1A). Injection of up to 2 nL of either 900 mM SB1, 900 mM SB2, or 900 mM SB3 alone decreased the amount of correctly-spliced **inab** mRNA (data not shown). Only co-injection of 2 nL of 300 mM SB2 and 300 mM SB3 resulted in missplicing of all **inab** transcript (Figure 1B), although we do not know whether missplicing creates a non-functional Inab protein. What follows is an analysis of the result of missplicing of **inab** caused by co-injection of 2 nL of 300 mM SB2 and SB3 MOs.

**inab** is expressed in CaP, VaP and VeLD, so we hypothesized that its knockdown would have an effect on the axons of these particular cells. To assay any changes in axon morphology, we co-injected SB2 and SB3 MOs into **nrp1a**:GFP transgenic embryos, in which GFP is expressed exclusively in CaP/VaPs beginning at 18 hpf [33], and **mnx1**:GFP transgenic embryos, in which GFP is expressed in all PMNs and VeLD from 14 hpf on [6]. CaP/VaPs begin projecting axons ventrally around 16–17 hpf [2,3]. These axons are obvious by 20 hpf in both CaP and VaP MNs and in VeLD INs. After 24 hpf, **inab** expression in CaP and VaP is downregulated, although it persists in VeLD and an additional, dorsally-located, glutamatergic IN. Scale bar, 5 μm in A–J; 10 μm in K.

**Figure 2. inab** is dynamically expressed in PMNs and INs. (A–K) Single confocal slices of embryos labeled with **inab**, **islet**, and neurotransmitter riboprobes (**gads** for GABA and **vglut** for glutamate). Diagonal lines represent somite boundaries. At 14 hpf, **inab** is coexpressed with neither **islet1** (A) nor **islet2a** (E). Between 16 hpf and 24 hpf, **inab** is expressed in **islet2a** PMNs (circled cells in F–H) but not **islet1** PMNs (arrowheads in A–C). **inab** is expressed in the VeLD IN, as determined by coexpression with both GFP in **mnx1**:GFP transgenic embryos (arrowheads, I) and **gad** mRNA (arrowhead, J). At 24 hpf, **inab** is coexpressed with **vglut** (arrowheads, K) in a cell dorsal to the VeLD IN (circle, K). (L) Schematic of **inab** mRNA dynamics during early development. Between 16–24 hpf, **inab** is expressed in both CaP and VaP MNs and in VeLD INs. After 24 hpf, **inab** expression in CaP and VaP is downregulated, although it persists in VeLD and an additional, dorsally-located, glutamatergic IN. Scale bar, 5 μm in A–J; 10 μm in K.

doi:10.1371/journal.pone.0088631.g002

**Figure 3. inaa** is expressed in the zebrafish head and not in PMNs. (A–E) Z-projections of confocal stacks of embryos labeled with **inaa** and **islet2a** riboprobes. Diagonal lines represent somite boundaries. Between 14 hpf and 24 hpf, **inaa** is never coexpressed with **islet2a** in the trunk (A–D). At 24 hpf, **inaa** expression can be seen in a region of the head that corresponds to the approximate location of the diencephalic ventricle (E). Boundaries of the fish head and eyes are marked with a dotted line. Scale bar 10 μm in A–D; 50 μm in E.

doi:10.1371/journal.pone.0088631.g003
we injected UAS:EGFP MO-injected embryos (Figure S3A–B), suggesting a role for have more small processes than ventral motor nerves in control embryos (Figure S3A–B). However, the ventral motor nerve, which contains axons of both CaP and VaP MNs appeared to be present and have a similar overall morphology to those of control embryos, CaP, VaP and VeLD axons are also present at 22 hpf. In control and Mnx MO-injected embryos, CaP, VaP and VeLD axons are present at 22 hpf. In control embryos, CaP, VaP and VeLD axons are present at 22 hpf. In MO-injected embryos, CaP, VaP and VeLD axons are also present and have a similar overall morphology to those of control embryos (Figure S3A–B). However, the ventral motor nerve, which contains axons of both CaP and VaP MNs appeared to have more small processes than ventral motor nerves in control embryos. The Bonfire program uses spatial information about the position of neurites to quantify characteristics such as the number of branch points along the length of an axon, or the relative connectivity of the neurites [45]. We found no significant difference in the number of processes of CaP axons between control and MO-injected embryos, irrespective of whether CaP was adjacent to VaP (Figure 5A–B, 5G). We also saw no difference in the number of processes along VeLD axons (data not shown). Another PMN, MiP does not express inab mRNA (n = 13, p = 0.14). We also examined the localization of these supernumerary processes and determined that they were significantly different along the VaP axon shaft (p = 0.017), whereas there was not a significant difference along the VaP growth cone (p = 0.20). The small but significant difference in the number of processes along the VaP axon shaft indicates that while the overall morphology of ventral motor axons is unchanged by inab knockdown, process formation along the length of the VaP axon is increased. Surprisingly, injection of inab MOs has little other effect on VaP cell fate — VaPs in MO-injected embryos still typically died by 36 hpf as shown in previous studies of control VaPs [24,25] (data not shown).

These results demonstrate that the phenotype of inab knockdown is surprisingly specific to a single inab-positive primary motoneuron in the ventral spinal cord.

Discussion

Microarray candidate genes are expressed in the zebrafish spinal cord

We used microarrays to identify genes that are expressed in zebrafish motoneurons and therefore potentially involved in motoneuron development. In our microarrays, we compared transcripts from spinal cords of embryos lacking Sonic hedgehog signaling to spinal cords from embryos lacking Notch signaling. We have previously shown that both of these signaling pathways

![Figure 4](image)

**Figure 4. Inab is expressed in MIP-CaP hybrids generated by Mnx protein knockdown.** (A–B) Z-projections of confocal stacks of control and Mnx MO-injected npt1a-GFP embryos labeled with inab riboprobe and anti-GFP antibody. At 26 hpf, inab is not expressed in GFP+ MIP MNs in control embryos (arrowheads, A–A’), but is expressed in a small percentage of GFP+ MIP MNs in MO-injected embryos (arrowhead, B–B’). Scale bar, 10 μm.

doi:10.1371/journal.pone.0088631.g004

![Figure 5](image)

**Figure 5. Inab is required for morphology of VaP axons.** (A–F) Z-projections of confocal images of control and inab MO-injected embryos. At 26 hpf, UAS:EGFP plasmid injection into mnx1:GAL4;UAS:td-Tomato transgenic embryos reveals similar overall morphology of CaP neurons in both control (A) and MO-injected embryos (B). VaP axons in MO-injected embryos (D–F) have more processes than VaP axons in control embryos (C). (G) Graph representing average number of branch points per axon in individually-labeled cells in control and MO-injected embryos. No significant difference between conditions for CaP neurons alone (n = 11, p = 0.51), CaP MNs that are located next to VaP MNs (n = 11, p = 0.38), or MiP MNs (n = 10, p = 0.38). Only the increase in VaP axon processes is significant (n = 14, p = 0.02). Scale bar is 10 μm in A–F.

doi:10.1371/journal.pone.0088631.g005

---

PLOS ONE | www.plosone.org 6 February 2014 | Volume 9 | Issue 2 | e88631
affect PMNs – excess PMNs are present in *dab7* injected embryos [38] and PMNs are decreased in *inaa* mutant embryos [50]. However, both signaling pathways also affect other cell types in the ventral spinal cord [1]. Thus, although we specifically focused our attention on PMNs, some of the genes identified in our microarray may be expressed, and regulated similarly to PMN-specific genes, but in other ventral spinal cord cell types.

We identified three candidate genes that are expressed in the zebrafish ventral spinal cord. Here we show that one of these candidates, the predicted transcription factor-encoding gene *ccdc85al*, is expressed in a population of cells just dorsal to the PMNs. Although it is not expressed in motoneurons, it may be expressed in closely-related interneurons that are generated from the same progenitor domain as motoneurons in zebrafish [23]. Investigation of *ccdc85al* could potentially be interesting, because much remains to be learned about interneurons that are generated from the zebrafish pMN domain.

We also show that another candidate gene from the microarray, the transcription factor-encoding gene *m22f1b*, is expressed broadly throughout the zebrafish spinal cord, including expression in PMNs. Based on its expression pattern, this gene may be less informative about mechanisms of motoneuron development than a gene with a cell type-specific expression pattern. The presence of *m22f1b* in the spinal cord, however, merits further investigation.

**Zebrasfish have two ina paralogs**

We show that the neuronal intermediate filament-encoding gene *ina* is expressed in a subset of zebrafish PMNs. In addition to *ina*, zebrafish also have another paralog of *ina*, named *inaa*, which we investigated alongside *ina*. We found that *inaa* is dynamically expressed in embryonic zebrafish, although not in PMNs during early development. This suggests that *ina* and *inaa* have diverged in function, leaving only *ina* with a role in PMN development.

**inab is dynamically expressed in identified zebrafish spinal neurons**

We found that *ina* is expressed in a subset of zebrafish PMNs – CaP and VaP – and a closely-related interneuron – VeLD. Previous studies of genes involved in PMN specification have underscored a close relationship between specific PMN subtypes and corresponding classes of interneurons. For instance, MiP motoneurons seem to be closely-related to V2a interneurons; in the absence of either Nks6 or Mnx proteins, MiPs form hybrids that have molecular and morphological characteristics of V2a interneurons [5,6]. On the other hand, CaP motoneurons and VeLD interneurons seem to be closely-related. In the absence of either Islet or Met proteins, CaPs form hybrids with molecular and morphological characteristics of VeLD interneurons [4,52]. Additionally, CaP and VeLD also arise from the same progenitor domain, and can even be siblings [23]. Therefore, the initial expression of *ina* in CaP/VaP and VeLD underscores the close relationship between these neurons.

We also demonstrated that *ina* is expressed in hybrid cells that have characteristics of both MiP and CaP motoneurons. We previously showed that in the combined absence of Mnx1, Mnx2a, and Mnx2b proteins, MiPs can acquire characteristics of both CaP motoneurons and V2a interneurons [6]. We were previously able to verify this MiP-CaP hybrid fate by only observing axon morphology, because there are few molecular markers that distinguish PMN subtypes. However, because we now know that *ina* is normally expressed in CaPs but not MiPs, we were able to use this marker to confirm that MiP-CaP hybrids co-express CaP markers as well as CaP morphology.

**inab is required for correct VaP axon branching**

In addition to being expressed in such a specific subset of cells, the timing of *ina* expression also suggested that this gene may be involved in directed axon outgrowth or the acquisition of axon morphology. CaP, VaP and VeLD neurons begin extending axons around 16–17 hpf [2,3,28]. *ina* is first expressed in these cells at 16 hpf, around the time of axogenesis. *ina* expression is also maintained in CaP and VaP through 24 hpf, by which time the CaP axon has completed extension to the ventral aspect of the muscle segment [2]. The specific temporal expression of *ina* immediately before and during axogenesis in CaP, VaP and VeLD places it in the right position to be involved in the acquisition of axon morphology.

Indeed, we have shown that when *ina* is forcibly misexpressed, VaP axons have significantly more processes than they do normally, indicating a role for *ina* in axon morphology of VaP MNs. The mechanism by which this axon morphology phenotype arises is unclear – *ina* may normally prevent VaP from reacting to some cue that promotes axon process formation. It is intriguing that although *ina* mRNA is expressed in CaP, VaP and VeLD, its disruption has an effect on only the VaP axon. It is unclear why this differential effect exists. *ina* may be interacting with as-yet-unknown gene products specific to VaP MNs. We also do not know if there is differential expression of Inab protein. Although a working antibody against Inab does not currently exist, it would be interesting to learn whether Inab protein is differentially expressed in CaP and VaP. Regulation at the protein level could contribute to the differential effect of *ina* misexpressing in these primary motoneurons.

Also worth noting is that although VaP axon morphology is changed when *ina* is disrupted, VaPs still typically die by 36 hpf, as they do in controls. This observation suggests that Inab is not involved in mediating VaP survival. This observation also indicates that the supernumerary processes of VaP axons are insufficient to overcome the normal interactions with CaP and muscle pioneer cells that are required to prevent VaP from extending a CaP-like axon and ultimately lead to VaP death [26,27].

**Supporting Information**

**Figure S1** RNA samples contain spinal cord tissue. (A–C) RT-PCR results confirming that all RNA samples contain spinal cord tissue. “Spinal cord” = *dbx1*, “hindbrain” = *krox20*, “somite” = *myod*. Wild-type RNA sample contains spinal cord tissue (asterisk, A), as well as some contaminating hindbrain tissue and little-to-no somite tissue (A). *dab7* injected (“supernumerary”) RNA sample contains spinal cord tissue (asterisk, B), as well as some contaminating hindbrain and somite tissues (B). *smoothed* mutant (“decreased”) RNA sample contains spinal cord tissue (asterisk, C), as well as some contaminating hindbrain tissue (C). (TIF)

**Figure S2** *ccdc85al* and *m22f1b* are expressed in the zebrafish spinal cord. (A–B) Single confocal slices of 24 hpf embryos labeled with riboprobe. *ccdc85al* is expressed in cells just dorsal to the *islet1* MNs (A), *m22f1b* is expressed broadly throughout the spinal cord, including expression in *islet1* MNs (circles, B-B’). Scale bar: 10 μm. (TIF)

**Figure S3** *inaa* knockdown does not affect ventral motor nerve or VeLD axons. (A–B) Z-projections of confocal images of control and *inaa* MO-injected *mnx1:GFP* transgenic embryos. At 20 hpf, both ventral CaP axons (arrows, A–B) and descending VeLD...
Conceived and designed the experiments: LVR LS JSE. Performed the experiments: LVR LS. Analyzed the data: LVR LS JSE. Contributed reagents/materials/analysis tools: LVR LS JSE. Wrote the paper: LVR LS JSE.

Acknowledgments

We thank Chris Doe and Monte Westerfield for comments on the manuscript, Steve Seredick for transgenic fish, and Jacob Lewis, Dayna Lamb, and the UO Zebrafish Facility for animal husbandry.

Author Contributions

References

1. Lewis K, Eisen J (2003) From cells to circuits: development of the zebrafish spinal cord. Progress in Neurobiology 69: 419–449.
2. Myers P, Eisen J (1986) Development and axonal outgrowth of identified motoneurons in the zebrafish. The Journal of Neuroscience 6: 2276–2289.
3. Eisen J, Myers P, Westerfield M (1986) Pathway selection by growth cones of identified motoneurons in live zebrafish embryos. Nature 320: 269–271.
4. Hutchinson S, Eisen J (2006) Id1delta and Id2elta have equivalent abilities to promote motoneuron formation and to specify motoneuron subtype identity. Development 133: 2131–2147.
5. Hutchinson S, Chereciss S, Hale L, Boone J, Eisen J (2007) Nkx6 proteins specify one zebrafish primary motoneuron subtype by regulating late id1 expression. Development 134: 1671–1677.
6. Scheder S, Van Rosyvlo L, Hutchinson S, Eisen J (2012) Zebrafish Mux proteins specify one motoneuron subtype and suppress acquisition of interneuron characteristics. Neural Development 7.
7. Lariviere R, Julien J (2009) Functions of intermediate filaments in neuronal development and disease. Journal of Neurology 58: 131–148.
8. Cochard P, Paulin D (1984) Initial expression of neurofilaments and vimentin in the central and peripheral nervous system of the mouse embryo in vivo. The Journal of Neuroscience 4: 2080–2094.
9. Chang L, Goldmann R (2004) Intermediate filaments mediate cytoskeletal crosstalk. Nature Reviews Molecular Cell Biology 5: 601–613.
10. Lindahl U, Zimmerman L, McKay R (1990) CNS stem cells express a new class of intermediate filament protein. Cell 66: 585–595.
11. Pachter J, Liem R (1985) alpha-Internexin, a 66-kD intermediate filament protein of Xenopus laevis, shares sequence features with neuronal intermediate filament protein, are regulated by the tectum during optic tract formation. Development 123: 3001–3008.
12. Niloff M, Dunn R, Raff M (1998) The levels of retinal mRNA for gefiltin, a neuronal binding protein from mammalian central nervous tissues. The Journal of Cell Biology 101: 1072–1084.
13. Meijering E, Jacob M, Sarria J, Steiner P, Hirling H, et al. (2004) Design and validation of a tool for neurite tracing and analysis in fluorescence microscopy images. Cytometry Part A 58: 167–176.
14. Kwan K, Fujimoto E, Grabher C, Mangum B, Hardy M, et al. (2007) The Toll2kit: A Multisite Gateway-Based Construction Kit for Toll2 Transposon Transgenesis Constructs. Developmental Dynamics 236: 3088–3099.
15. Langhammer C, Previtera M, Sweet E, Seaton S, Chen M, et al. (2010) Automated Sholl analysis of digitized neuronal morphology at multiple scales: whole cell Sholl analysis versus Sholl analysis of arbor subregions. Cytometry Part A 77A: 1160–1168.
16. Meijering E, Jacobs J, Sarric J, Steiner P, Hirling H, et al. (2004) Design and validation of a tool for neurite tracing and analysis in fluorescence microscopy images. Cytometry Part A 58: 167–176.
17. Rodriguez A, Ehlenberger D, Hof P, Warneke S (2006) Rayburst sampling, an algorithm for automated three-dimensional shape analysis from laser scanning microscopy images. Nature Protocols 1: 581–585.
18. Lenth RV (1989) Algorithm AS 243: Cumulative Distribution Function of the Non-central t Distribution. Journal of the Royal Statistical Society, Series C 38: 185–189.
49. Appel BE, Eisen JS (1998) Regulation of neuronal specification in the zebrafish spinal cord by Delta function. Development 125: 371–380.

50. Lewis KE, Eisen JS (2001) Hedgehog signaling is required for primary motoneuron induction in zebrafish. Development 128: 3485–3495.

51. Bernhardt R, Patel C, Wilson S, Kuwada J (1992) Axonal trajectories and distribution of GABAergic spinal neurons in wildtype and mutant zebrafish lacking floor plate cells. The Journal of Comparative Neurology 326: 263–272.

52. Tallafuss A, Eisen J (2006) The Met receptor tyrosine kinase prevents zebrafish primary motoneurons from expressing an incorrect neurotransmitter. Neural Development 3.

53. Nakatani Y, Takeda H, Kohara Y, Morishita S (2007) Reconstruction of the vertebrate ancestral genome reveals dynamic genome reorganization in early vertebrates. Genome Res 17: 1254–1265.

54. Braasch I, Postlethwait J (2012) Fish polyploidy and the teleost genome duplication. In: Soltis PSS, D.E., editor. Polyploidy and Genome Evolution. Berlin, Heidelberg: Springer. pp. 341–383.

55. Sato-Maeda M, Tawarayama H, Obinata M, Kuwada J, Shoji W (2006) Sema3a1 guides spinal motor axons in a cell- and stage-specific manner in zebrafish. Development 133: 937–947.