Extraocular Motoneuron Pools Develop Along a Dorsoventral Axis in Zebrafish, *Danio rerio*

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ABSTRACT
Both spatial and temporal cues determine the fate of immature neurons. A major challenge at the interface of developmental and systems neuroscience is to relate this spatiotemporal trajectory of maturation to circuit-level functional organization. This study examined the development of two extraocular motor nuclei (nIII and nIV), structures in which a motoneuron’s identity, or choice of muscle partner, defines its behavioral role. We used retro-orbital dye fills, in combination with fluorescent markers for motoneuron location and birthdate, to probe spatial and temporal organization of the oculomotor (nIII) and trochlear (nIV) nuclei in the larval zebrafish. We describe a dorsoventral organization of the four nIII motoneuron pools, in which inferior and medial rectus motoneurons occupy dorsal nIII, while inferior oblique and superior rectus motoneurons occupy distinct divisions of ventral nIII. Dorsal nIII motoneurons are, moreover, born before motoneurons of ventral nIII and nIV. The order of neurogenesis can therefore account for the dorsoventral organization of nIII and may play a primary role in determining motoneuron identity. We propose that the temporal development of extraocular motoneurons plays a key role in assembling a functional oculomotor circuit. J. Comp. Neurol. 525:65–78, 2017.

INDEXING TERMS: oculomotor; development; cranial nerve; nIII; nucleogenesis; cranial nucleus; RRID:SCR_002285; RRID:SCR_010279; RRID:ZFIN_ZDB-GENO-060619-2; RRID:ZFIN_ZDB-GENO-030919-1

Spatiotemporally regulated neurogenic processes underlie the production of diverse neuronal classes and subclasses including those of the vertebrate retina (Cepko, 2014), cortical projection neurons (Desai and McConnell, 2000), and interneurons (Sousa and Fishell, 2010), and individually identifiable cells within the lineage of a *Drosophila* neuroblast (Novotny et al., 2002). Trajectories of sensory neuron development were recently linked to their mature sensitivity profiles (Li et al., 2012). In the motor system, links between spatiotemporal organization and functional circuitry exist among interneuron subtypes involved in motor circuits (Lewis and Eisen, 2003). In the zebrafish spinal cord, interneuron birth order is related not only to dorsoventral topography but also to electrophysiological properties and recruitment order during progressively faster swimming behaviors (Fetcho and McLean, 2010). Spatiotemporal organization has likewise been observed for mammalian spinal cord interneuron populations with distinct functional roles (Tripodi et al., 2011). Similar evidence at the level of motoneuron organization is sparse, but would be welcome given the clear correspondence of motoneuron-to-muscle wiring and behavioral outputs. This study seeks to address the outstanding question of developmental organization of extraocular motoneuron pools.

The oculomotor periphery consists of three cranial motor nuclei whose connectivity has been well characterized across vertebrate species, thanks to their participation in highly conserved goal-driven and reflexive behaviors (Büttner-Ennever, 2006): the oculomotor (nIII), trochlear (nIV), and abducens (nVI) nuclei. nIII
comprises motoneuron populations innervating four of the six extraocular muscles and is the only one of these nuclei to house multiple pools. Motoneurons targeting each extraocular muscle are organized into pools within nIII, the coherence of and overlap between pools varying between species (Evinger, 1988). Such clustering also exists among motoneurons in the spinal cord, which form pools that innervate distinct muscles (Jessell et al., 2011). Pool clustering is dependent on cadherin signaling but independent of underlying motoneuron identity, or choice of muscle target (Price et al., 2002; Demireva et al., 2011). The development of nIII motoneuron identity is controversial. One proposal for how individual motoneurons come to acquire their identity is post-hoc self-identification following random extraocular muscle innervation (Glover, 2003), consistent with target-derived signals regulating motoneuron connectivity (Ladle et al., 2007) but not with a stereotyped order of subpopulation birth (Shaw and Alley, 1981). Others propose that a caudal-to-rostral order of differentiation of both extraocular muscle and motoneuron groups allows each newly-born pair to match (Evinger, 1988), although this conflicts with the observed order of motor nucleus development (Altman and Bayer, 1981; Varela-Eschavarria et al., 1996). We set out to elucidate the process of motoneuron fate determination in larval zebrafish.

A model vertebrate, the larval zebrafish has contributed much to our understanding of the organization and development of oculomotor circuitry and behavior (Miri et al., 2011; Ma et al., 2014). Recent work described the timecourse of axonal pathfinding and synaptogenesis between oculomotor neurons and their target extraocular muscles (Clark et al., 2013), but no characterization of either nIII motoneuron pool organization or its development yet exists. As part of a rich behavioral repertoire (Fero et al., 2010), 3–4-day-old larval zebrafish begin to perform eye movements (Easter and Nicola, 1997) involving all six extraocular muscles: two for nasal/temporal horizontal saccades (Schoonheim et al., 2010), and the remaining four for the vertical and torsional vestibulo-ocular reflexes (Riley and Moorman, 2000; Mo et al., 2010; Bianco et al., 2012). We took advantage of the external development, transparency, and genetic tractability of the larval zebrafish to probe nIII development using transgenic reporters of motoneuron location (Higashijima et al., 2000) and methods to mark birthdate in vivo (Caron et al., 2008).

Here we found that larval zebrafish extraocular motoneuron pools in nIII are organized along a dorsoventral axis, and that a similar axis defines their birth order. We first measured the distribution of extraocular motoneurons across nIII and nIV labeled in Tg(isl1:GFP) larvae, and used these data to construct a spatial framework for motoneuron localization. Using targeted retro-orbital dye fills, we localized extraocular motoneuron somata in nIII and nIV of larval zebrafish. We found that nIII is divided into dorsal and ventral halves; nIV aligns with the dorsal half of nIII. Dorsal nIII contains mostly inferior/medial rectus (IR/MR) motoneurons, while inferior oblique (IO) and superior rectus (SR) motoneurons occupy largely distinct regions of ventral nIII. Next, we measured the time of terminal differentiation of motoneurons across nIII and nIV, and discovered a dorsoventral order to birthdate, complementary to the spatial organization of motoneuron pools. We propose that birth order is the primary determinant of the spatial organization and fate of extraocular motoneurons in nIII.

MATERIALS AND METHODS

Fish husbandry

All protocols and procedures involving zebrafish were approved by the New York University Langone School of Medicine Institutional Animal Care & Use Committee (IACUC). All larvae were raised at 28.5°C at a density of 20–50 larvae in 25–40 mL of buffered E3 (1 mM HEPES added). Older larvae, >7 days postfertilization (dpf), were raised in the facility on a standard 14/10 hour light/dark cycle, and fed twice daily.

Transgenic lines

Experiments were done on the mitfa/- background to remove pigment. Larvae homozygous for Tg isl1:GFP) (Higashijima et al., 2000; RRID:ZFIN_ZDB-GENO-030919-1) were used to label extraocular motoneurons for retro-orbital fill. Homozygotes were identified by level of GFP fluorescence relative to heterozygous siblings. Larvae for photoconversion experiments were doubly-heterozygous for Tg(elavl3:Kaede)rw0130a (Tg(huC:Kaede)) (Sato et al., 2006; RRID:ZFIN_ZDB-GENO-060619-2), which labels postmitotic neurons, and Tg(isl1:GFP).

Retro-orbital fills

Larvae were anesthetized in 0.02% Ethyl-3-aminobenzoic acid ethyl ester (MESAB, Sigma-Aldrich E10521, St. Louis, MO) at 4–5 dpf or 13 dpf until no longer responsive to touch/dish taps. Anesthetized larvae were mounted in 2% low-melting point agar (ThermoFisher Scientific 16520, Pittsburgh, PA), positioned with the right eye up and oriented for best access to nerve targeted (Fig. 1A). Larvae were oriented predominantly dorsal up for superior oblique (SO) and superior rectus (SR) neuron fills, or predominantly
ventral up for inferior oblique (IO) and inferior/medial rectus (IR/MR) fills, tilted slightly off-axis in either case. Agar was cleared from the area above and around the muscle innervated by the nerve of interest. An electrochemically sharpened tungsten needle (10130-05, Fine Science Tools, Foster City, CA) was used to create an incision in the skin at the position of the muscle, closely following the outside of the eye (Fig. 1A2–1B). Any excess liquid was removed from the area. A second sharpened tungsten needle was used to hold a piece of solidified dextran-conjugated Alexa Fluor 647 dye (10,000 MW, ThermoFisher Scientific D-22914) into the incision, until the dye had spread such that the incision location was fully colored (Fig. 1A3). Larvae were left for a minimum of 5 minutes following dye application, at which point E3 was added over the agar drop. Larvae were then freed from the agar and allowed to recover in E3 for a minimum of 4 hours, and usually overnight. Dye spread at the orbit was not systematically characterized, as dextran-conjugated dye is expected to wash out except where taken into structures severed by the incision. Dye was sometimes observed to extend to extraocular muscles neighboring the target site, indicating off-target incision; for instance, IO muscle fibers (but not necessarily IO motoneurons) occasionally stained following an IR fill.

Data collection and analysis
Assignation of filled cells as specific motoneurons
Larvae were anesthetized in 0.02% MESAB and mounted dorsally in 2% agar for image collection. Larvae were imaged between 5–7 dpf, or at 14 dpf, using the following confocal microscopes: Zeiss LSM 510 (SO and SR data, 40x/0.8NA objective), Leica SP8 in photon counting mode (IO data, 20x/1.0NA objective), Zeiss LSM 710 (IR/MR data, 40x/1.1NA objective), and Zeiss LSM 800 (Kaede data, 40x/1.0NA objective). Image analysis was performed using ImageJ (Fiji) (Schindelin et al.,...
of extraocular motoneurons (Fig. 1C, D), which are highly conserved across vertebrates (Evinger, 1988); and second, the known lack of GFP expression in IO motoneurons in the Tg(isl1:GFP) background (Higashijima et al., 2000). Somata filled contralaterally to the right eye (left side of the brain) were defined as SO motoneurons if found in nIV, or SR motoneurons if found in nIII (Fig. 2A). Somata filled ipsilaterally to the right eye (right side of the brain) were defined as IR/MR motoneurons, if GFP+, or IO motoneurons, if GFP– (Fig. 2B). We observed three putative contralateral IO motoneurons, which were excluded.

**nIII/nIV template generation and motoneuron identification**

To provide a consistent spatial framework for placement of filled cells within the boundaries of nIII/nIV, we generated a template (Fig. 3). Two evenly aligned stacks from 5–7 dpf larvae, showing minimal roll or pitch deviations, were chosen to estimate stereotyped boundaries of GFP+ cells. Fifteen dorsoventral divisions of the nuclei were established, spaced ~6 μm apart (“z0–z14”). Maximum intensity projections were made using 12 μm of data centered at each dorsoventral division. Boundaries were manually drawn around the extent of GFP+ cells within each projection and expanded to encompass all cells across the two larvae to represent the expected extent of GFP+ cells at each division. The midbrain–hindbrain boundary delineated nIII and nIV. The template generation process was repeated using four stacks to generate boundaries for 14 dpf larvae, also resulting in 15 divisions spaced ~6 μm apart. To test how stereotyped GFP+ expression relative to anatomical landmarks was across larvae, we manually bounded both green somatic fluorescence and the ascending posterior mesencephalic central artery (PMcta) on each side of the midline, at the level of the medial longitudinal fasciculus (MLF), and measured the distance between the center of fluorescence intensity and the centroid of the corresponding PMcta (Fig. 3B).

When excited at 488 nm, Alexa Fluor 647 dye emits fluorescence detectable in the 500–550 nm collection channel at the gains necessary to resolve dim signal in Tg(isl1:GFP) larvae. We developed a thresholding procedure to differentiate GFP emission from dye emission. Control fills were first performed and imaged in Tg(isl1:GFP) negative siblings. Pixel values from brightly fluorescent dye-filled cells were used to establish an intensity floor for the GFP channel. For each data stack, imaged under identical experimental conditions, the intensity was adjusted to only display values above the floor. Applying this minimum intensity display threshold to stacks from homozygous Tg(isl1:GFP) larvae allowed us to distinguish dye-filled GFP– cells from faintly...
GFP+ cells. To define somatic location within nIII/nIV, individual filled cells were classified as particular motoneurons as described above and localized within the boundaries of the template at the appropriate dorsoventral plane. Somata were represented as circles ~4 µm in diameter, roughly the size of a small nIII/nIV neuron. Neurons with somata that sat between template planes were assigned based on the plane of brightest fluorescence. Each soma was only counted once.

**Birthdating of neurons using a photolabile protein**

Larvae heterozygous for Tg isl1:GFP and Tg(huC:Kaede) were exposed for 10 minutes to focused full spectrum light from a mercury light source at 22, 29, 36, 43, or 50 hpf (Fig. 4A) to photocovert green Kaede to red. Larvae were kept in the dark following collection until anesthesia and mounting for imaging at 5 dpf. To establish a threshold for red Kaede signal that corresponded to a postmitotic neuron, rather than basal photocooperation, we used data from stacks of Tg(huC:Kaede); Tg isl1:GFP siblings that had not been exposed to light. Control and experimental stacks were matched for overall Kaede expression. Seven experimental stacks that greatly exceeded control stacks in overall Kaede intensity were not used for analysis. Three stacks were used for neuronal birthdating at 22 and 43 hpf, four stacks were used for 50 hpf, and five stacks were used for 29 and 36 hpf. GFP+ neurons in the final image were placed into the map of nIII/nIV only if they contained supra-threshold red Kaede signal in the initial image.

**RESULTS**

**Dorsoventral distribution of isl1:GFP+ cells in cranial nuclei III and IV**

To establish a spatial framework to localize extraocular motoneurons, we employed the Tg isl1:GFP line, which labels SO, SR, IR, and MR motoneurons (Higashijima et al., 2000). Extraocular motoneurons were observed 100–200 µm deep relative to the dorsal surface in the midbrain and hindbrain. We divided a region that spanned about 90 µm in depth into 15 unique planes spaced every 6 µm, and defined boundaries around GFP+ motoneurons (Fig. 3). We tested the consistency of GFP+ expression relative to other anatomical landmarks across larvae by measuring the distance between the center of intensity of green somatic fluorescence on one side of the midline and the center of the corresponding ascending PMCtA at the level of the MLF (Fig. 3B). We measured this distance for each hemisphere in 10 larvae and recorded an average of 18 ± 4 µm (SD). We conclude that the distribution of GFP fluorescence is largely stereotyped relative to anatomical landmarks identifiable under visible illumination.

To quantify motoneurons in nIII and nIV, we characterized the distribution of GFP+ motoneurons within nIII and nIV in ten 5 dpf larvae (Fig. 5). We observed 104 ± 20 (SD) nIII motoneurons (max: 134, min: 68) and 47 ± 8 nIV motoneurons (max: 58, min: 35) per
side per larva. Note that nIII totals do not include IO motoneurons, which are not labeled in Tg(isl1:GFP) (Higashijima et al., 2000). GFP neurons were found in comparable numbers across larvae at every plane (Fig. 5B). As in avians and other teleosts, GFP motoneurons in nIII could be organized into dorsal and ventral sections (Evinger, 1988). Here, each division accounts for approximately half of the height and cell count of the nucleus. GFP motoneurons in nIV align with the dorsal half of nIII on the caudal aspect of the midbrain–hindbrain boundary (MHB), and number 45% as many as nIII motoneurons.

As the level of GFP varies across motoneurons, we tested the interobserver reliability. Motoneurons from the same five larvae were counted independently by two observers (M.G. and A.P.). A 3-way analysis of variance grouping data for dorsoventral sublevel, larva, and observer rejected the null hypothesis of equal means for all three variables \( P < 10^{-16}, P < 10^{-16}, P = 0.044, \ df = 14, 4, 1, F = 41, 16, 4.1 \). Since we cannot rule out an observer effect on motoneuron quantification, we measured interobserver differences. On average, M.G. found 9% more GFP cells, but interobserver counts were correlated with a pairwise linear correlation coefficient of 0.97. We conclude that while the absolute motoneuron counts may vary by experimenter, the relative spatial distribution of motoneurons is robust to interexperimenter variation. Boundaries representing the extent of isl1:GFP motoneurons are a suitable spatial framework to map motoneuron position.

**Superior rectus motoneurons are restricted to ventral nIII in both 5–7 and 14-day-old zebrafish**

To determine if Tg(isl1:GFP) is a veridical marker of extraocular motoneurons, we targeted SO motoneurons for retro-orbital fills. SO motoneurons comprise the entirety of nIV and project to the contralateral eye with distinct dorsal axonal morphology. Our fills labeled

![Figure 4](Image)

Birthdating Analysis by photoconverted fluorescent Protein Tracing In vivo, with Subpopulation Markers (BAPTISM) method (Caron et al., 2008) for identifying the time of terminal differentiation (birthdate) of motoneurons in Tg(huC:Kaede; isl1:GFP) zebrafish, which express photolabile fluorescent protein Kaede pan-neuronally. (A) Neurons born by the time of initial photoconversion (one of five timepoints shown) contain converted Kaede in initial image, taken at 5 dpf. Second photoconversion converts remaining Kaede, leaving GFP as the only green signal in Final image. Images are compared to determine birthdate of GFP motoneurons. (B) Dorsal plane (z12, dashed line) from a 5 dpf Tg(huC:Kaede; isl1:GFP) larva initially photoconverted at 36 hpf. Yellow arrowheads indicate an nIII motoneuron (isl:GFP, final image) born by 36 hpf (converted huC:Kaede, initial image). White arrowheads indicate a neuron born by 36 hpf not belonging to nIII/nIV. Orange arrowheads indicate an nIV motoneuron not born by 36 hpf. Scale bars = 20 \( \mu m \).
individual somata within a cluster just caudal to the MHB and contralateral to the targeted eye. Each dye-filled soma was also GFP+ in this area (Fig. 6, magenta). Dye-filled axons were often visible within the GFP+ IVth nerve, ascending dorsally and contralaterally. We concluded from their GFP expression, axonal morphology, and location in the hindbrain that these cells were successfully filled SO motoneurons. We never observed filled somata in nIV that were not GFP+, supporting the hypothesis that the Tg isl1:GFP is a veridical marker of motoneurons. On average, we labeled 42 ± 33 motoneurons per larva, 14 ± 10 of which were contralateral neurons in nIV, yielding a total of 606 SO motoneurons across 42 larvae. We observed a maximum of 43 SO motoneurons in a single larva. We conclude that the retro-orbital fill method is well suited to stochastically label extraocular motoneurons.

To examine nIII subpopulation distributions, we first defined SR-projecting motoneurons as all GFP+ nIII motoneurons filled on the contralateral side of the brain. SR motoneurons appeared solely in the ventral half of nIII and were particularly dense in medial and rostral portions of the nucleus (Fig. 6, blue). On average, we labeled 60 ± 44 neurons per larva, 10 ± 12 of which were contralateral GFP+ neurons in nIII, yielding a total of 229 SR motoneurons across 22 larvae. We observed a maximum of 41 SR motoneurons in a single larva; both dorsolateral and within-subdivision distribution persisted if this larva was removed from consideration. We did not observe any GFP-negative filled cells in this part of contralateral nIII, supporting the hypothesis that, as with SO motoneurons, Tg isl1:GFP reliably labels SR motoneurons.

Early in development, the extraocular motoneuron somata may still be motile (Puelles-Lopez et al., 1975). To determine if the spatial organization at 5–7 dpf represented stable localization of motoneurons within the
nucleus, we filled SO and SR motoneurons in 13 dpf larvae. Contralaterally filled cells were distributed with similar frequency among dorsoventral subdivisions at 14 dpf as at 5 dpf (Fig. 6B) in similar positions within subdivisions, supporting the hypothesis that the organization evident among motoneurons at 5–7 dpf is stable across early development.

**Inferior oblique motoneurons are found in ventral nIII, largely distinct from superior rectus motoneurons, while inferior/medial rectus motoneurons are found in dorsal nIII**

We next defined IO-projecting motoneurons as all GFP– somata filled on the ipsilateral side of the brain. Similar to SR motoneurons, the large majority of IO neurons lay in the ventral half of nIII at 5–7 dpf. A few scattered IO cells were also present in dorsal nIII, interspersed within the GFP+ cell population. Most of the ventral IO motoneurons were located on the lateral or caudal edges of the GFP+ cell cluster, although some were found closer to the midline (Fig. 7, red). In all, 154/328 labeled neurons were GFP– in 30 larvae; all but one of the larvae had GFP+ neurons labeled as well. On average, we labeled 11 ± 7 neurons per larva, 5 ± 4 of which were GFP– neurons in nIII. The maximum number of IO neurons filled in a single larva was 13. The low number is likely due to the conservative criteria used to define a soma as GFP–, given the presence of GFP+ neuropil and the axial resolution of the confocal microscope.

Data from adult teleosts suggests that IO and SR motoneuron pools are spatially distinct (Graf and McGurk, 1985). We compared the positions of IO and SR motoneurons within ventral nIII subdivisions (Fig. 7A). As in adult teleosts, we observed that IO and SR populations are largely spatially segregated into caudolateral and rostromedial divisions of ventral nIII, respectively.

The remaining nIII subpopulations, IR and MR motoneurons, both innervate ipsilateral extraocular muscles, and are both labeled in Tg isl1:GFP. Furthermore, both the IR and MR muscles are found near the dye insertion site (Kasprick et al., 2011). Thus, we were unable to distinguish IR from MR motoneurons in our fills. To optimally differentiate IR/MR motoneurons from IO motoneurons, we only evaluated larvae that had no dye in GFP– somata. IR/MR motoneurons were located throughout the entirety of dorsal nIII at 5–7 dpf, and infrequently in ventral nIII (Fig. 7A, brown). On average, we labeled 10 ± 11 neurons per larva, 9 ± 11 of which were ipsilateral GFP+ neurons in nIII, yielding 160 IR/MR motoneurons across 17 larvae. The maximum number hit in single larva was 41; dorsoventral distribution remained similar if this larva was removed from consid-

![Figure 7.](image-url) Inferior oblique motoneurons are located mainly in ventral nIII in 5–7-day-old zebrafish, in a distinct caudal region relative to superior rectus motoneurons, while inferior/medial rectus motoneurons are located mainly in dorsal nIII. (A) Location of IO (red), SR (blue), and IR/MR (brown) motoneurons at 5–7 dpf across nIII/nIV. GFP– neurons in nIII ipsilateral to the filled eye were defined as IO motoneurons; GFP+ neurons in nIII ipsilateral to the filled eye were defined as IR/MR motoneurons. SR data is mirrored across the midline from Fig. 4. (B) Probability distributions of dye-filled IO and IR/MR motoneurons across nIII/nIV. IO motoneurons: n = 30 larvae (5–7 dpf), 154 cells. IR/MR motoneurons: n = 17 larvae (5–7 dpf), 160 cells.

**Development of extraocular motoneuron organization in nIII proceeds along a dorsoventral order**

To determine if the dorsoventral organization that we observed in nIII was related to the time of terminal differentiation, we localized motoneurons born before a series of five timepoints (22, 29, 36, 43, and 50 hpf). “Birthdating” was accomplished with a photolabile fluorescent protein in a second transgenic line,
Tg(huC:Kaede), using a validated technique, BAPTISM, detailed in the Materials and Methods and Figure 4 (Caron et al., 2008). Timepoints spanned most of nIII/nIV nucleogenesis (Higashijima et al., 2000; Clark et al., 2013), allowing us to localize the somata of the majority of neurons in nIII/nIV with respect to their final mitotic division. We used the Tg(isl1:GFP) marker to define nIII/nIV motoneurons, and thus could only localize SR, IR, MR, and SO, but not IO, motoneurons.

We mapped extraocular motoneurons born at each timepoint into the boundaries of nIII/nIV (Fig. 8). Overall, 25–33% of the Tg(huC:Kaede)+ dorsal nIII population evaluated at 5 dpf was born by 22 hpf. Nearly all of dorsal nIII was born by 29 hpf. Almost no ventral nIII motoneurons were born until after 36 hpf; ventral nIII motoneuron birth appeared complete by 50 hpf. We first saw postmitotic motoneurons in nIV at the 36 hpf conversion, and nIV motoneuron birth also appeared complete by 50 hpf. We conclude that the dorsoventral organization of nIII reflects the birth order of motoneurons.

DISCUSSION

This study found that extraocular motoneurons in cranial nuclei III and IV of the larval zebrafish are spatially organized, in a manner consistent with their birth order. We localized extraocular motor pools within nIII and nIV of larval zebrafish, assigning identities and relative positions to retro-orbitally dye-filled somata using the characteristic patterns of extraocular muscle innervation and of Tg(isl1:GFP) expression in these nuclei. Our data show a spatial organization to nIII pools in the 5–7 dpf zebrafish: IR/MR motoneurons are predominantly found in dorsal nIII, while IO and SR motoneurons are predominantly found in ventral nIII (Fig. 9A). Since SR and nIV SO organization does not change between 5–7 and 14 dpf, pool location is likely stable throughout early larval development. By birthdating neurons in nIII/nIV, we discovered a complementary dorsoventral division in timing of neurogenesis (Fig. 9B). As both motoneuron identity and birthdate comparably delineate nIII/nIV, the time of terminal differentiation could account for the dorsoventral organization of nIII subpopulations. We propose that birth order plays a preeminent role in determining extraocular motoneuron fate, such that dorsal IR and MR motoneurons become postmitotic earlier than ventral SR motoneurons and SO motoneurons in nIV. Here we consider the limitations of our techniques, a general model for spatiotemporal development of extraocular motoneurons, and experiments to elucidate the mechanisms underlying such spatiotemporal organization.

Limits of the retro-orbital and BAPTISM techniques

Because the larval zebrafish extraocular muscles are too small to restrict dye placement, we classified...
motoneurons by whether their somata were ipsilateral or contralateral to the targeted eye, and on their overlap with Tg isl1:GFP expression. Consequentially, we would incorrectly label any ipsilaterally innervating SO or SR motoneurons or contralaterally innervating IO, IR, or MR motoneurons. Several earlier studies have reported infrequent ipsilateral innervation of SO or SR, by <5% of the total motoneuron pool in each case (Luiten and de Vlieger, 1978; Graf and Baker, 1985; Miyazaki, 1985; Murphy et al., 1986; Evinger et al., 1987; Sonntag and Fritzsch, 1987; Sun and May, 1993), and one study reported rare contralateral IR motoneurons (El Hassni et al., 2000). The low numbers seen in other organisms lead us to believe that few, if any, motoneurons were incorrectly labeled in our study. We conclude that the spatial organization we observe is likely robust to misclassification errors.

Our protocol does not distinguish IR from MR motoneurons, which may be spatially segregated. Selectively severing only the nasal part of the MR nerve, far from the IR/MR muscle bifurcation, could improve the odds of labeling MR motoneurons. This might allow us to discern a spatial organization to the GFP+ ipsilateral motoneuron population, with the caveat that MR would be incompletely labeled. IR and MR motoneurons do occupy distinct regions of nIII in other species, with varying degrees of overlap. For instance, in other teleosts (goldfish and carp), MR motoneurons have a wider dorsoventral distribution than mainly dorsal IR motoneurons; and MR motoneurons are found clustered ventral to IR motoneurons in flatfish (Luiten and de Vlieger, 1978; Graf and McGurk, 1985; Graf and Baker, 1985). Across vertebrates, IR is usually rostral to MR (Evinger, 1988); and in zebrafish, dorsal nIII may be divisible along the rostrocaudal axis at 50 hpf (Clark et al., 2013). We predict, by homology, that zebrafish IR motoneurons may be located rostrally and/or dorsally to MR motoneurons.

The BAPTISM procedure relies on a threshold to separate red Kaede fluorescence indicative of postmitotic neurons from the basal background level of spontaneous photoconversion. We chose conservative background red Kaede thresholds, which might exclude motoneurons that had only recently exited the cell cycle upon initial photoconversion, whose above-background red Kaede signal would still be faint. Similarly, we may also have missed a cluster of 10–15 postmitotic motoneurons in ventrocaudal nIII, or 5–8% of total nIII GFP+ cells, that systematically expressed low basal levels of Tg(huC:Kaede). Even accounting for these potential underestimates, our finding of patterned dorsal-to-ventral birth order is unaffected. Finally, IO motoneurons are not labeled by Tg isl1:GFP, and so are excluded from our birthdating dataset. We began to see postmitotic GFP− cells around the edges of ventral nIII at the 36 hpf conversion timepoint, but cannot positively identify these as nIII motoneurons. We conclude that the temporal order we observe is likely robust to technical constraints of the BAPTISM method.

Figure 9. Spatiotemporal organization of extraocular motoneurons in nIII and nIV. (A) Aggregate figure showing relative locations of nIII/nIV motoneurons at two exemplar planes; data from Figs. 6, 7. (B) Comparison of motoneuron subpopulation birthdates (gradient bars) with development landmarks (Clark et al., 2013) from live imaging of extraocular motoneuron projections (timeline markers; events inside gray box). Solid color bars = periods between BAPTISM initial photoconversion timepoints.
How many motoneurons innervate each extraocular muscle?

Reports from other teleosts vary in the relative numbers accorded to each motor pool (Luiten and de Vlieger, 1978; Graf and McGurk, 1985). While we cannot specify the precise number of neurons that make up a particular motoneuron pool, we propose the maximum number filled in any one larva as an estimate. Based on our Tg(isl1:GFP) counts, nIV is composed of 47 ± 8 SO neurons; the maximum number of SO neurons filled in any one larva was 43, supporting our approach. Within nIII, the maximum number for SR was similar at 41; this would comprise close to the expected one-third (35, total n = 104 ± 20) of nIII motoneurons in Tg(isl1:GFP). For IO motoneurons the maximum was only 13, a likely underestimate reflecting the challenge of calling a soma GFP⁺ in the presence of dense GFP⁺ nIII neuropil and/or signal from surrounding GFP⁺ cells. Our best estimate for combined IR/MR pool numbers is the total number of GFP⁺ motoneurons in the dorsal half of nIII, or 55 ± 14 per side. This is less than the expected (70) two-thirds of GFP⁺ neurons in nIII and far less than would be expected relative to SO or SR numbers, given an even subpopulation distribution. The discrepancy between observed and expected numbers could reflect stochastic expression in the Tg(isl1:GFP) line or an unequal number of motoneurons innervating each extraocular muscle. As we never observed a filled motoneuron on the contralateral side that was not GFP⁺ (i.e., SO and SR motoneurons), Tg(isl1:GFP) expression likely reflects the full complement of motoneurons. Our data instead are consistent with a slight superior/inferior asymmetry, where superior muscles SO and SR may have larger motoneuron pools. At rest, larval zebrafish maintain constant tension in the superior muscles, enabling a slight upward gaze deviation (Bianco et al., 2012), potentially necessitating such an asymmetry.

Model for the spatiotemporal development of nIII/nIV

Our findings largely support and extend earlier work that defined the timeline for extraocular motoneuron axon projection and extraocular muscle innervation in larval zebrafish (compared in Fig. 9B) (Clark et al., 2013). Our data agree with their observation of neurogenesis in nIII preceding nIV, consistent with the order of isl1 expression in chick (Varela-Echavarría et al., 1996) but in contrast to their concurrent development in mammals (Altman and Bayer, 1981; Shaw and Alley, 1981). The oculomotor nerve begins projection at 30 hpf, before birth of SR motoneurons has begun. The projecting nerve therefore contains only IR/MR, and possibly IO, axons. nIII nucleogenesis is mostly complete by 48 hpf, when the axon front pauses at the edge of the orbit. Because most SR motoneurons are not postmitotic until after 43 hpf, their axons are likely delayed in reaching this location. Indeed, axon extension toward the IR, IO, and MR muscles resumes at 54 hpf, but Clark et al. (2013) do not observe a branch innervating the SR muscle until 66 hpf. This may reflect IO differentiation before SR in the zebrafish, in agreement with the order seen in rabbit (Shaw and Alley, 1981).

Our data speak to two prior hypotheses about the developmental organization of extraocular motoneurons. One proposition was that a shared caudal-to-rostral sequence of extraocular motoneuron and extraocular muscle development allows progressive innervation of earliest-born muscles by earliest-born extraocular motoneurons (Evinger, 1988). This hypothesis does not match the true order of events, since the IR and MR muscles, which are the latest to differentiate (Noden et al., 1999) and still a paired anlage when invaded by motoneuron axons (Clark et al., 2013), are innervated by the earliest motoneurons to differentiate. In contrast, nIV motoneurons innervating the earlier-differentiating SO muscle are born only after IR/MR motoneuron birth is complete. Nor do our results agree with a random innervation model of nIII development (Glover, 2003). Such a model predicts no correlation between motoneuron birthdate and muscle target, in contrast to the dorsal-to-ventral progression we observe. Other factors thus are likely to define the organization of nIII/nIV, at least in larval zebrafish.

The dorsoventral organization of nIII in larval zebrafish is similar to that seen in avians and other teleosts (Evinger, 1988); embryonic avian nIIs feature particularly well-delineated dorsolateral IR, dorsomedial MR, and ventral IO (lateral) and SR (medial) regions. The timing of nIII development we observe likewise agrees with previous studies: Hasan et al. (2010) found a dorsal-to-ventral gradient of neurogenesis in the chick oculomotor complex (which includes Edinger-Westphal motoneurons). Since avian and zebrafish nIII dorsoventral spatial organizations are comparable, the dorsoventral temporal gradient in chick implies birth of IR/MR before IO/SR motoneurons, as we report here. Shaw and Alley (1981) found an IR-MR-IO-SR progression in the birth order of rabbit nIII subpopulations. In rabbits and other mammals, the dorsoventral spatial organization is inverted relative to avians and teleosts, with IR and MR populations generally ventral to IO and SR populations. The temporal order of motoneuron birth thus appears to be conserved, while migration pattern / eventual location
within the nucleus may not be. We propose that our finding in the larval zebrafish of motoneuron pools organized according to birthdate is likely general, even when the final spatial layout is different. This hypothesis could be tested in a species in which motoneuron identity and spatial order are partially uncoupled. In adult frog (*Rana esculenta*), SR motoneurons occupy most of ventromedial nIII, while the ipsilaterally innervating nII subpopulations overlap in dorsolateral nIII (Matesz and Székely, 1977). Correspondence of birth order and motoneuron identity in this preparation would provide strong evidence for the primacy of birthdate in fate determination.

### Potential mechanisms and consequences of spatiotemporal development

Spatiotemporal determination of identity is a fundamental theme of neurogenesis (Moore and Livesey, 2015). One possible mechanism for temporally ordered development is to ensure that motoneuron precursors sit in particular spatial locations within nIII, where external cues then specify identities of each subpopulation (Shaw and Alley, 1981). Experiments in which oculomotor complexes were ectopically induced laterally and rostrally in the chick midbrain, with normal dorsoventral positioning of viscerals, suggest that major spatial cues working on the developing nIII lie along the dorsoventral axis (Hasan et al., 2010). Other extrinsic contributions might resemble the negative feedback cues onto retinal cell progenitors that inhibit prolonged production of early-born retinal ganglion and amacrine neurons (Waid and McLoon, 1998; Belliveau and Cepko, 1999; Poggi et al., 2005), as retinoic acid produced by early-born spinal cord motoneurons acts on progenitors as a cue for late-born motoneuron generation (Sockananth and Jessell, 1998). However, cell-autonomous intrinsic factors such as the “temporal series” of transcription factors inherited by the offspring of *Drosophila* neuroblasts (Isshiki et al., 2001; Jacob et al., 2008) may also play a role. Transplantation of GFP+ neurons from Tg(isl1:GFP) donor embryos into particular spatial locations in the dorsal or ventral midbrain would allow us to differentiate intrinsic and extrinsic factors, as well as to discover the precise timing of extracocular motoneuron commitment. Similar experiments performed in cortex (McConnell and Kaznowski, 1991; Frantz and McConnell, 1996) and spinal cord (Appel et al., 1995) have defined time windows within which progenitors’ competence to respond to external cues shifts or narrows, representing a convergence of intrinsic and extrinsic cues.

As the final target for all oculomotor circuits, multiple upstream inputs must find partners within particular extracocular motoneuron pools. Our data support a model in which, by the time motoneurons occupy their final positions within nIII, they possess sufficient information to choose a target extracocular muscle. Our study thus supports molecular profiling of developing motoneurons at this time, specifically a comparison of dorsal and ventral GFP+ neurons from Tg(isl1:GFP) at later stages of nucleogenesis. We hypothesize that each subpopulation will express a unique complement of terminal effectors (Hobert, 2016) that allow it to respond appropriately to cues arising from potential muscle targets responsible for motoneuron matching. Similarly, our model suggests that motoneurons may be competent to signal upstream partners, even before synaptogenesis onto target extracocular muscles (72 hpf).

Since the motoneurons occupy fairly stereotyped regions, neurons projecting to the extracocular motor nuclei might themselves rely on spatial cues to aid targeting. This strategy is employed by sensory afferents projecting to the expected locations of motor pools in the spinal cord (Sürmeli et al., 2011), as well as in the *Drosophila* nerve cord at spatial scales on the order of 10 μm (Zlatic et al., 2009), suggesting the potential for sufficiently fine spatial patterning. Vestibular projection neurons mediating the “nose-down/eyes-up” response of the vestibulo-ocular reflex (joint activation of IO and SR motoneurons/muscles) might thereby be able to target ventral nIII with little chance of incorrect innervation (Bianco et al., 2012). Spatial segregation is likely to complement rather than replace more traditional molecular recognition cues, due to imperfect segregation of motoneuron pools within nIII.

Together, transplantation and molecular profiling experiments can uncover candidate mechanisms and factors downstream of *isl1* expression that define extracocular motoneuron subtypes. Understanding the contributions of temporal and spatial factors in nIII motoneuron identification should clarify how they find, and are found by, partners during circuit construction. Our data showing common dorsoventral organization between motoneuron identity and birth order provide crucial spatial and temporal guidance for this set of experiments. We propose that the larval zebrafish extracocular motor system is a powerful model to uncover the molecular basis for extracocular motoneuron organization, and by extension, the behaviors it subserves.

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CONFLICT OF INTEREST
The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS
Conceptualization: MG and DS. Methodology: MG and DS. Investigation: MG, AP, and KD. Validation: MG and AP. Visualization: MG. Writing: MG and DS. Funding Acquisition: DS. Supervision: DS.

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