V3 Interneurons Are Active and Recruit Spinal Motor Neurons during In Vivo Fictive Swimming in Larval Zebrafish

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Abstract
Survival for vertebrate animals is dependent on the ability to successfully find food, locate a mate, and avoid predation. Each of these behaviors requires motor control, which is set by a combination of kinematic properties. For example, the frequency and amplitude of motor output combine in a multiplicative manner to determine features of locomotion such as distance traveled, speed, force (thrust), and vigor. Although there is a good understanding of how different populations of excitatory spinal interneurons establish locomotor frequency, there is a less thorough mechanistic understanding for how locomotor amplitude is established. Recent evidence indicates that locomotor amplitude is regulated in part by a subset of functionally and morphologically distinct V2a excitatory spinal interneurons (Type II, nonbursting) in larval and adult zebrafish. Here, we provide direct evidence that most V3 interneurons (V3-INs), which are a developmentally and genetically defined population of ventromedial glutamatergic spinal neurons, are active during fictive swimming. We also show that elimination of the spinal V3-IN population reduces the proportion of active motor neurons (MNs) during fictive swimming but does not alter the range of locomotor frequencies produced. These data are consistent with V3-INs providing excitatory drive to spinal MNs during swimming in larval zebrafish and may contribute to the production of locomotor amplitude independently of locomotor frequency.

Key words: locomotion; motor neuron recruitment; spinal interneuron; zebrafish

Significance Statement
Currently, there is a limited understanding about the cellular and spinal network properties that produce locomotor amplitude, defined as limb displacement in limbed animals or tail-bend in nonlimbed animals during locomotion. Here, we show, directly for the first time in a vertebrate, that V3 interneurons (V3-INs) in zebrafish larvae are active during in vivo fictive locomotion, and that targeted ablation of the spinal V3-IN population reduces the proportion of active motoneurons during fictive swimming. Importantly, ablation of V3-INs does not affect locomotor frequency (speed), which clarifies their role in motor control rather than rhythm generation. Thus, we propose that the V3-IN population is a source of excitation in the vertebrate locomotor neural circuitry that may participate in regulating locomotor amplitude.

Introduction
Efficient locomotion is set by a combination of kinematic properties, including the frequency and amplitude of rhythmic motor output. These properties combine to determine various locomotor features, such as distance traveled, speed, and vigor (Webb, 1975; Jayne and
Lauder, 1995; Donley and Dickson, 2000; Tytell, 2004). Locomotor frequency (e.g., limb or tail-beat rhythmic activity) is a well-studied kinematic variable, which has been shown to increase linearly with speed (Bainbridge, 1958; Hunter and Zweifel, 1971; Webb et al., 1984; Müller and van Leeuwen, 2004). Less well studied is locomotor amplitude, which is measured by limb displacement in limbed animals during locomotor events (walking, trotting, galloping, hopping; Heglund et al., 1974) or by tail-bend in nonlimbed animals during anguilliform swimming (Wardle et al., 1995; Videler et al., 1999). Although there is a fundamental understanding of how identified subsets of excitatory spinal interneurons help to establish the frequency of motor output (Ausborn et al., 2012; Eklöf-Ljunggren et al., 2012; Kimura et al., 2013; Ampatzis et al., 2014; Ljunggren et al., 2014; Menelaou et al., 2014), a more complete understanding of the cellular and network properties that regulate locomotor amplitude in vertebrate animals is lacking.

Across vertebrate species from fish to mammals, the neural circuitry that controls locomotion is located in the ventromedial spinal cord (Grillner and Jessell, 2009; Kiehn, 2011). In vertebrates, developmental gene expression specifies at least five cardinal classes of ventral neuronal progenitor cells. These progenitors give rise to motor neurons (MNs) and four classes of ventral interneurons, referred to as V0 to V3 neurons (Ericson et al., 1997; Pierani et al., 1999). Over the past decade, a population of identified excitatory spinal interneurons (V2a) has been shown to control the speed of locomotion in larval and juvenile/adult zebrafish (Ausborn et al., 2012; Ampatzis et al., 2014). More recent studies indicate a role for a subset of V2a interneurons (Type II, nonbursting) in regulating locomotor amplitude (Menelaou and McLean, 2019) and/or vigor (Song et al., 2018) in larval and adult zebrafish, respectively. Both studies reveal a hierarchical control of MN recruitment by a diverse V2a interneuron population, which the authors conclude is the basis for regulating locomotor amplitude and/or vigor.

A cardinal class of spinal interneurons in the ventral cord that selectively regulates locomotor amplitude has been elusive. In the mammalian spinal cord, a genetically defined population of premotor glutamatergic neurons, called V3 interneurons (V3-INS), contributes to limb coordination, motor burst duration, and muscle spasms, but does not have a clear role in locomotor pattern generation (Chopek et al., 2018; Danner et al., 2019; Lin et al., 2019). Recently, neurons derived from the p3 progenitor domain were shown to modulate locomotor adaptation in larval zebrafish (Böhm et al., 2022). We hypothesized V3-INS provide excitatory drive during locomotor activity to regulate locomotor frequency, amplitude, or both. The work presented here shows directly that V3-INS in zebrafish larvae are active during in vivo fictive locomotion (swim-active), and that elimination of the V3-IN spinal population reduces the proportion of active MNs during fictive swimming but does not alter the range of locomotor frequencies produced. These data are consistent with V3-INS providing excitatory drive to spinal MNs during swimming in larval zebrafish and may contribute to the production of locomotor amplitude independently of locomotor frequency. This is conceptually important because it clarifies a role of the V3 cardinal class of spinal interneurons in motor control rather than rhythm generation.

Materials and Methods

Fish lines and maintenance

All animal procedures were performed in accordance with the animal care committee’s regulations. Wild-type (Segrest Farms) and transgenic adult zebrafish (Danio rerio) were maintained in the animal facility. The transgenic lines used in these experiments were: Tg(nkx2.2a: mEGFP)ru17 (Kirby et al., 2006), Tg(vglut2a: DsRed)mns9 (Miyasaka et al., 2009), Tg(vglut2a:Galo4ff)mns20 (Satou et al., 2013), Tg(UAS:GCaMP6s)ynk13a (Muto et al., 2017), and parga2m2Et (Balcıunas et al., 2004). Out-crossed transgenic larvae at 4–6 d postfertilization (dpf) were used in these experiments. Larval zebrafish were maintained in Petri dishes filled with embryo water (60 µg/ml Instant Ocean salt mix) and 0.00005% Methylene Blue in a 28.5°C incubator with a 14/10 h light/dark cycle (Zeitgeber time 0 = 8:30 A.M.).

V3-IN soma distribution analysis

Quantification of the distribution of V3-IN cell bodies was performed using Fiji and custom MATLAB code (MathWorks). A confocal microscope (Olympus, FV-100) was used to collect overlapping stacks of images documenting the expression of Tg(vglut2a: DsRed)mns9 and transmitted light morphology. Images were acquired at high magnification (40×/NA 0.8 objective) and a z-axis resolution of 1.3 µm. The fields of view in each stack overlapped with their neighbors by at least 10% of the width to permit accurate stitching and included the full width of the spinal cord and dorsal root ganglia (DRG) in the field of view. The collection of stacks was stitched into a single 3-D image of the caudal hindbrain and spinal cord (Preibisch et al., 2009). The Fiji Cell Counter tool (Schindelin et al., 2012) was used to identify the coordinates of all V3-INS in the rostral 2 mm of the larval spinal cord (out of a total length of ~3 mm), the coordinates of the DRG neuron clusters, and the position of the dorsal and ventral edges of the spinal cord. This work was supported by National Institutes of Health Grants R01 NS065054 (to M.A.M.), R01 NS094176 (to M.A.M.), F31 NS083110 (to T.D.W.), and R25 NS083059; Regenerative Medicine Minnesota Grant RMM 091718 DS 006 (J.E.M.); Grant-in-Aid of Research, Artistry, and Scholarship (M.A.M.), and the University of Minnesota Foundation Bridge Award B-0621-01 (to M.A.M.).

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Whole-mount immunohistochemistry

The markers tracing the dorsal and ventral edges of the spinal cord were used to calculate local tangent lines to the dorsal and ventral edges of the spinal cord at the rostrocaudal position of each V3-IN. The equation of a line perpendicular to the ventral tangent and intersecting the V3-IN was calculated, referred to as the “V3 plumb line.” The rostrocaudal distance of each V3-IN was calculated as the distance along the ventral surface of the spinal cord between the spinal cord/hindbrain junction [segment (s)2/3 boundary] and the V3 plumb line intersection with the ventral surface. The dorsoventral height of the spinal cord at the location of each V3-IN was calculated as the distance between the intersections of the V3 plumb line with the dorsal surface and the ventral surface, respectively. The distance of the V3-IN from the ventral surface of the spinal cord was also measured along the plumb line. The z-axis position of DRG neurons on each side of the spinal cord were used as boundary points of the lateral edges of the spinal cord, and a linear fit to the DRG neurons on each side of the animal was used to calculate the edges of the spinal cord at each rostrocaudal position in the imaging area. The midline slice and spinal cord width were calculated at the rostrocaudal position of each V3-IN, along with the distance from midline of the V3-IN.

**Kaede photoconversion**

*In vivo* photoconversion of genetically encoded Kaede was performed in Tg(vglut2a:Gal4ff)mn20;Tg(UAS:Kaede)s1999t larvae using a fixed-objective, upright microscope (Olympus FV-1000, 60×/NA 1.0 objective). Larvae were anesthetized in zebrafish Ringer’s solution (Wiggin et al., 2012, 2014) with 0.02% tricaine-S (MS-222; Western Chemical), embedded laterally in a fluorodur (World Precision Instruments) filled with 1.5% low-melting point agarose in zebrafish Ringer’s solution and covered with additional saline containing anesthetic. Photoconversion of Kaede was performed using the bleach function of the FV-1000 to direct a 405-μm laser spot onto a Kaede-expressing cell targeted for conversion (5% laser intensity, 7 s, sinusoidal laser spot onto a Kaede-expressing cell targeted for conversion (5% laser intensity, 7 s, sinusoidal). The embedding procedure resulted in small variations in the orientation of the spinal cord such that the larvae were not precisely level and square with respect to the imaging plane. Following confocal stack acquisition, Fiji (Schindelin et al., 2012) was used to rotate the image to correct these errors.

**Whole-mount immunohistochemistry**

Five dpf parga<sup>mn2ET</sup> larvae were killed with 0.2% tricaine-S and fixed overnight in 4% paraformaldehyde at 4°C. Whole-mount larvae were permeabilized with Proteinase K, post-fixed, and blocked with 0.2% BSA, 10% normal donkey serum, 0.5% Triton X-100 in PBS. Goat polyclonal anti-choline acetyltransferase (ChAT; 1:50; AB144P) was added to the blocking solution and incubated overnight at 4°C. Fixed larvae were incubated with Alexa Fluor 633 donkey anti-goat secondary antibody (1:500; A-21082, Thermo Fisher Scientific Inc.) in secondary blocking solution (0.2% BSA, 1% normal donkey serum, 0.5% Triton X-100 in PBS) for three nights at 4°C.

Anti-ChAT-labeled parga<sup>mn2ET</sup> larvae were embedded laterally in low melting-point agarose. An Olympus Fluoview FV1000 confocal microscope was used to collect images through a 60× objective centered on body S15. Confocal stacks contained the entire dorsoventral extent of the spinal cord and spanned approximately three body segments (285 μm). Images were processed in Fiji and labeled cells from a single side of the spinal cord (approximately three hemisegments) were quantified using the Cell Counter plugin. Photoshop CS5 (Adobe Systems) was used to adjust brightness and contrast across entire z-projected images.

**In vivo Rhod-2 AM dye loading of MNs**

For *in vivo* characterization of MNs in parga<sup>mn2ET</sup> larvae, a 0.25% Rhod-2 AM (Life Technologies) stock was made by dissolving 50 μg of dye in 20-μl Pluronic F-127:DMSO (1:4; Life Technologies). Next, the skin was removed from the midbody region and a 1:50 dilution of the Rhod-2 AM stock was bath applied to the exposed muscles to permit sequestration of indicator dye by MNs via the peripheral motor axons. Sensory neurons were not strongly loaded by this procedure. Following 1-h incubation, the Rhod-2 AM solution was thoroughly washed-out using zebrafish Ringer’s solution. Confocal images were collected, and cells were quantified as described for whole-mount immunohistochemistry (above).

**Peripheral nerve (PN) recordings and analysis**

Zebrafish larvae were anesthetized, the skin was carefully removed from the recording region, and larvae were paralyzed with 0.1 mM α-bungarotoxin (Tocris) to prevent muscle contractions. The anesthetic was removed to allow the larvae to perform fictive behavior, and a suction electrode (9–15-μm tip diameter) was placed over the intermyotomal cleft to record MN axons. Fictive motor bursts were identified and grouped into episodes using custom MATLAB code. The program detected the presence or absence of activity at each voltage sample \( v(n) \). For each \( v(n) \), the algorithm determined a voltage autocorrelation \( c_n(k) \) over a small window (3 ms) centered at \( v(n) \). These “windowed” autocorrelations were computed as:

\[
c_n(k) = \sum_{i=-N_0}^{i=N_0} v(n-i) v(n-i-k)
\]

where 3-ms windowing was implemented in Equation 1 by setting \( N_0 = (3 \text{ ms} \cdot f_{\text{sam}})/2 \), where \( f_{\text{sam}} \) is the sampling frequency, and by setting \( v(j) = 0 \) for \( j \) outside the interval \([n-N_0, n+N_0]\).

A subset of the autocorrelation values (lags) from Equation 1 were used to compute a test statistic for each \( v(n) \) with the same lags \( (k = k_1, k_2, \ldots, k_m) \) used for all voltage samples. Building on Equation 1, for each \( v(n) \), a test statistic \( c_n \) was computed as:

\[
c_n = \sum_{k=1}^{k_m} \sum_{i=-N_0}^{i=N_0} v(n-i) v(n-i-k)
\]

where Equation 2 is the sum of the \( c_n(k) \) from Equation 1 specified by \( k = k_0 \) was set at \( k_0 = [1, 2] \) because we found that these values effectively separate the distributions of the test statistics \( \{c_n\} \) for samples of noise and
samples of activity across a broad range of recording quality.

Finally, activity was considered present at \( v(n) \) only when \( c_0 \) was greater than a detection threshold \( T \). \( T \) was set as the maximum of a set of \( \{c_n\} \) corresponding to the \( \{v(n)\} \) in one contiguous second of the voltage recording where activity was confirmed to be absent (typically the first second of the recording), and was set this way for each individual voltage recording to account for differences in baseline noise levels. fictive locomotor bursts were detected and grouped into episodes, and the burst and episode properties were determined as follows. Episode duration is the time from the onset of the first burst of an episode to the offset of the final burst in the same episode. Burst duration is the time from the onset to the offset of each burst, as defined by \( c_o \) and described above. Burst frequency is the inverse of the interburst period (IBP), which is defined for each pair of bursts within episodes as the time from the onset of the first burst to the onset of the second burst. Peak burst frequency is the inverse of burst period, which is defined as the shortest time duration from onset of one burst to onset of the subsequent burst in an episode of fictive swimming. IBPs between episodes are excluded from burst frequency analysis.

**Calcium imaging and analysis**

For calcium imaging of V3-INs, Tg(vglut2a:Gal4ff\(^{\text{rms20}}\); Tg(UAS:GCaMP6s)\(^{\text{pjk103a}}\) zebrafish larvae were used. For calcium imaging of MNs, Calcium Green-1 AM (Life Technologies) dye was loaded into MNs using the protocol for Rhod-2 AM dye loading described above. Larvae were transferred to a fixed-stage Olympus BX51 WI upright microscope with either a 20\( \times \)/NA 0.5 or a 40\( \times \)/NA 0.8 water dipping objective lens. A field of view containing GCaMP6s-expressing V3-INs in transgenic larvae or Calcium Green-1 AM-loaded MNs in wild-type larvae was selected for calcium imaging and a simultaneous PN recording was acquired. Calcium indicator fluorescence was excited with an E-Cite Series 120 Q lamp (Excélitas Technologies) and acquired at 20 Hz with a Retiga EXi camera and QCapture Pro 6.0 software (QImaging). Data were acquired during spontaneous fictive swimming in 10- to 60-s durations for both V3-IN and MN trials. Multiple trials (typically 4) were acquired for each field of view, with at least 5 min between trials. Movement artifacts during data acquisition were corrected in Fiji by registering frames from all trials to a common reference image (Preibisch et al., 2009). Optical and electrophysiological data acquisition were synchronized by using the sync output of the stimulator to trigger acquisition of the PN recording.

For all calcium imaging experiments, raw fluorescence values were determined by drawing a region of interest around each identified V3-IN or MN soma and using the Plot z-axis Profile command in Fiji. Fluorescence from a region of interest containing background signal was subtracted from raw soma fluorescence measurements to reduce the effect of photobleaching. Baseline fluorescence (\( F_0 \)) was defined as the mean fluorescence in the first 10 frames of the recording. To calculate \( \Delta F/F \), \( F_0 \) was subtracted from fluorescence for each frame \( [F(t)] \), then divided by \( F_0 \); \( \Delta F/F = (F(t) - F_0)/F_0 \). A neuron was considered active if \( \Delta F/F \) was 10% or greater.

**Targeted laser ablation of V3-INs**

Acute ablation of the V3-INs was performed using a Micropoint pulsed nitrogen pumped dye laser (Andor Technology). Light (\( \lambda = 345 \text{ nm} \)) was delivered via an Olympus BX51 WI microscope using a 60\( \times \)/NA 1.0 water dipping objective lens. The laser was focused to apply maximal light intensity to the imaging plane of the microscope, the intensity of the laser was calibrated to produce a 4.5- to 5.0-\( \mu \text{m} \) hole in a mirror slide (this optimal intensity was determined empirically), and the x-y coordinates of the laser point in the field of view were indicated on a digital display. Three to 6 dpf Tg(vglut2a:DsRed\(^{\text{rms9}}\) zebrafish larvae were anesthetized in 0.02% tricaine in zebrasfish Ringer’s solution and embedded laterally in a gel composed of 3% methylene-cellulose and 0.02% tricaine in zebrasfish Ringer’s saline. A coverslip was placed over the gel and secured at the edges with agarose to stabilize the position of the larvae.

Targeted, bilateral laser ablation of DsRed\(^+\) V3-INs in the ventral spinal cord was performed by positioning the microscope objective such that a neuron was in the x-y position at which the laser was focused. Laser pulses were applied to each V3-IN soma at 33 Hz for 10 s. Laser intensity was empirically determined a priori such that >95% of the targeted V3-INs were not present 24 h post-ablation. Following the train of laser pulses, the microscope was repositioned to focus on another identified V3-IN, and the protocol was repeated. Larvae were embedded for ~50 min to perform bilateral ablation of ~140 identified V3-INs between body S5 and S20. To confirm that laser ablation of targeted V3-INs was effective and persistent, the lack of DsRed fluorescence in the lesioned spinal cord of Tg(vglut2a:DsRed\(^{\text{rms9}}\) larvae was confirmed ~24 h postablation. Sham larvae were treated identically to V3 ablation larvae, but the x-y position of the laser was offset by 5-20 \( \mu \text{m} \) from each V3 cell, producing nonspecific damage to neurons other than the V3 population. Rather than kill a specific cell type as a sham control which might have a specific effect, nonspecific laser damage was preferred.

**Statistical analysis**

Statistical analysis of data were performed using two-sample \( t \) test, ANOVA test, Holm–Sidak post hoc test, Kolmogorov–Smirnov test, and least-squares linear regression. Statistical tests were conducted using SigmaPlot 12 and 14 software (SyStat Software), MATLAB (MathWorks), or Microsoft Excel (Microsoft). An \( \alpha \) level of 0.05 was used to determine statistical significance. Data are reported as means and SD.

**Software accessibility**

Custom MATLAB code for the quantification of the distribution of V3-IN cell bodies and PN recording analysis is available on request.
Results

V3-INs are individually identifiable in the larval zebrafish spinal cord

To characterize the functional roles of V3-INs in vitro, it was necessary to reliably target the neurons for recording and perturbation. While there are no published transgenic zebrafish lines that specifically label neurons expressing sim1a (the zebrafish V3-IN marker; Schäfer et al., 2007), it is possible to infer V3-IN identity from their other characteristics. V3-INs proliferate ventral to the central canal and are glutamatergic (Zhang et al., 2008; Yang et al., 2010).

In zebrafish, there is a restricted to the region ventral to the central canal (Fig. 1A). At 4–6 dpf, nkk2.2a:mEGFP expression was retracted to the region ventral to the central canal (Fig. 1B). The p3 domain of the zebrafish spinal cord also produces Kolmer–Agduhr neurons (Yang et al., 2010), and likely other neuronal types, so it was expected that there would be more nkk2.2a:mEGFP-expressing neurons than vglut2a:DsRed-expressing neurons (Fig. 1C). Approximately 94% of vglut2a:DsRed-expressing neurons ventral to the central canal also expressed nkk2.2a:mEGFP (n = 6 larvae, 162 neurons), and were therefore confirmed to be V3-INs (Fig. 1D). There was not a significant difference between the dorsoventral position of the vglut2a:DsRed-expressing neurons that co-expressed nkk2.2a:mEGFP and those that did not (two-sample t-test; $t = 1.6; p = 0.10$), so there was not a way to reliably segregate the identified V3-INs, from those that did not express nkk2.2a:mEGFP. However, the overwhelming majority of the DsRed$^+$ neurons ventral to the central canal were confirmed to be V3-INs. The lack of confirmed transgene overlap in the remaining cells may be because of variegated transgene expression rather than the presence of a non-V3-IN ventral glutamatergic neuronal population. Based on this experiment, we treat the ventral vglut2a:DsRed-expressing neurons as the V3-IN population, with the caveat that a small fraction (~6%) may be misidentified.

V3-INs are located in the ventromedial spinal cord and are distributed along the rostrocaudal axis of the spinal cord

In mice, the V3-INs migrate following differentiation to form spatially segregated dorsal and ventral populations that have different cellular properties and may be necessary for different behaviors (Borowska et al., 2013). To determine whether there are multiple populations of spatially segregated V3-INs in zebrafish larvae, we used confocal microscopy to create 3-D reconstructions of the spinal cords of 4–6 dpf Tg(vglut2a:DsRed)$_{nns9}$ zebrafish larvae (Fig. 2A,B). The center of each V3-IN soma in the most rostral 2 mm of the spinal cord was tagged, and the distribution of the neurons within the spinal cord was calculated based on the imputed borders of the spinal cord (see Materials and Methods). Neurons in the far caudal spinal cord were not counted because the dimensions of the spinal cord taper to the extent that it was difficult to
determine the location of the central canal and Mauthner axons. All V3-INs (n = 4 larvae; 797 neurons) were located medial of the Mauthner axon (Fig. 2C). V3-IN density was highest in the rostral spinal cord but remained close to 1 cell per 10 μm throughout the cord (Fig. 2D). The V3-IN somas were ~5 μm in diameter in the rostrocaudal axis, so a density of ~1 cell per 10 μm thoroughly tiled the ventral spinal cord. The majority of V3-INs were located in the most medial and most ventral 25% of the spinal cord (Fig. 2E,F). There was no parasagittal plane through the medial spinal cord that was unoccupied by V3-INs (Fig. 2E,F), which suggested that the V3-IN population may be considered a midline population. However, the density of V3-INs was not highest at the sagittal midline of the spinal cord, and when cell density was plotted along the population axis, the point of highest density was adjacent to the spinal cord midline (Fig. 2G). Therefore, we propose that distinct left and right populations of V3-INs are present, but that there are not multiple ipsilateral populations of spatially segregated V3-INs in larval zebrafish.

V3-INs project bilaterally and have descending processes

To characterize the morphologies and putative synaptic connectivity of the V3-INs, we examined the 3-D morphology of individual V3-INs using confocal microscopy (Fig. 3). The density of DsRed fluorescence is too high in the Tg(vglut2a:DsRed)nns9 transgenic line to accurately trace the processes of a single neuron, so we expressed a photoconvertible protein (Kaede) in Tg(vglut2a:Gal4)nns20;Tg(UAS:Kaede)s1999t larvae to identify and photconvert single V3-INs (Fig. 3A,B). Following photoconversion, it was possible to obtain morphologic images of the V3-INs (n = 5 neurons, 4 larvae; Fig. 3C). Based on this sample of V3 morphologies, it was evident that variation in the projection patterns of the V3-INs was present. For example, the V3-IN imaged in S10 had minimal projection density dorsal or rostral to the cell body, while the V3-IN in S18 predominantly projected rostrally and dorsally (Fig. 3C, lateral projection). Despite these differences, there were some common features of this sample of V3-IN morphologies. First, the imaged V3-INs possessed bilateral projections, but had a higher density of processes ipsilateral to the cell body (Fig. 3C, horizontal projection). Second, V3-INs had the greatest density of processes close to the cell body and longer caudal projections than rostral projections (Fig. 3C). Third, while the full extent of the descending processes of the S9 and S14 neurons is not presented in Figure 3C, V3-INs had a maximum detectable projection distance shorter than 300 μm. The limited projection length we observed may be because of the
Kaede photoconversion method used and should be considered a lower bound on the full extent of the projections of the V3-INs. The projection pattern of the V3-INs appeared to innervate the ventral spinal cord, which is populated by MNs, V2a-INs (Eklöf-Ljunggren et al., 2012; Menelaou and McLean, 2013), other V3-INs, KA neurons, and serotonergic neurons (McLean and Fetcho, 2004; Yang et al., 2010; Montgomery et al., 2018).

The majority of V3-INs are active during spontaneous fictive swimming

In mice, there is only indirect evidence supporting temporal correlation between V3-IN activity and locomotion (Borowska et al., 2013). To test the hypothesis that activity in the V3-INs is appropriately timed to provide excitatory drive to MNs during locomotion, we monitored the activity of the V3-INs during spontaneous fictive swimming using a genetically encoded calcium indicator (GCaMP6s; Fig. 4).

Figure 3. V3-INs are descending, bilaterally projecting cells. Kaede was expressed in vglut2a-expressing neurons in Tg(vglut2a:Gal4ff)\textsuperscript{rm20}, Tg(UAS:Kaede)\textsuperscript{1999t} double transgenic larvae (A). A single V3-IN was exposed to UV laser light, converting the Kaede protein from green fluorescence to red (B). C, Inverted contrast projections of Kaede-red fluorescence in five neurons. Each row of images is shown in a lateral z-projection (left) and horizontal z-projection (right). On the horizontal projections, the center of the cell body is indicated with white triangles to account for optical bloom and contralateral projections are indicated with black triangles. Scale bar: 20 μm.

The activity of V3-INs and fictive locomotor output during spontaneous swimming was measured via simultaneous calcium imaging and PN recording using 4–6 dpf Tg(vglut2a:Gal4ff)\textsuperscript{rm20}, Tg(UAS:GCaMP6s)\textsuperscript{mrk13a} larvae (Fig. 4B,C). We found that the majority, 44 of 48 (91.7%), of V3-INs were active (ΔF/F > 10%) during spontaneous fictive swimming episodes across a range of burst frequencies [mean burst frequency = 23.4 (SD 2.0) Hz, range = 13.9–28.6 Hz; mean peak burst frequency = 31.8 (SD 12.3) Hz, range = 20.2–74.6 Hz; n = 6 larvae, 14 trials, 81 swim episodes; Fig. 4D]. Therefore, this provides direct evidence that V3-INs are active during fictive locomotion, which supports the participation of V3-INs in producing locomotor activity.

Ablation of V3-INs does not affect burst frequency during spontaneous fictive swimming

The previous experiments demonstrated that V3-INs are positioned appropriately and have neurotransmitter...
phenotype and activity consistent with providing excitatory drive to MNs; however, demonstration of the function of the V3-INs or their necessity for behavior has not been described. To determine whether burst frequency is regulated by V3-IN activity, we performed laser ablation of a large percentage of the V3-IN population (95%; ~140 V3-identified somas across body S5–S20; Fig. 5). Three experimental groups were used. (1) The V3 ablation group, in which all V3-INs on both sides of the spinal cord between S5 and S20 were laser ablated (n = 6 larvae; 145 (SD 9.6) neurons per larva). Representative examples of targeted laser ablation efficacy within a restricted region (Fig. 5 A, B) and spatial localization (Fig. 5 C, D) are shown. (2) The sham ablation group, in which laser pulses were delivered 5–20 μm rostral to each identified V3-IN cell body (n = 6 larvae; 130 (SD 10.8) sham targets per larva). (3) The unmanipulated control group was not embedded or exposed to laser pulses (n = 6 larvae).

To measure the effects of V3-IN ablation on burst frequency during spontaneous fictive locomotion, fictive motor activity was measured using extracellular PN recordings 24 h after the ablation procedure. This delay between ablation and recording was necessary to confirm cell death by disambiguating the temporal properties of the loss of fluorescence caused by application of laser pulses. Specifically, laser pulses applied to targeted neurons that does not result in cell death will produce a temporary loss of fluorescence (photobleaching), which will recover since protein synthesis in the viable cells will produce additional fluorescent molecules. In contrast, persistent loss of fluorescence is consistent with absence of new protein synthesis and, thus, cell death.

Assessment of the mean burst frequency probability distributions showed that the Ablated group was significantly reduced compared with the Control group (Kolmogorov–Smirnov test, D = 0.34, p < 0.004; Fig. 6B). However, mean burst frequency was not significantly different between either the Control and Sham groups (Kolmogorov–Smirnov test, D = 0.33, p = 0.16; Fig. 6B) or the Sham and Ablated groups (Kolmogorov–Smirnov test, D = 0.21, p = 0.63; Fig. 6B). Further, analysis of variance showed that mean burst frequency was not significantly different between Control [29.0 (SD 3.7) Hz], Sham [28.0 (SD 4.2) Hz], and Ablated [27.2 (SD 4.7) Hz] groups (one-way ANOVA, F = 2.2, p = 0.2; Fig. 6C).

Assessment of the peak burst frequency probability distributions showed that there were no significant differences
between Control, Sham, and Ablation groups (Kolmogorov–Smirnov test, D = 0.14, p = 0.09 for all group comparisons; Fig. 6D). This was consistent with an analysis of variance that showed peak burst frequency was not significantly different between Control [39.0 (SD 8.5) Hz], Sham [39.3 (SD 8.8) Hz], and Ablated [36.4 (SD 10.7) Hz] groups (one-way ANOVA, F = 1.2, p = 0.3; Fig. 6E).

Finally, an effect on the coordination of rostrocaudal (ipsilateral) and side-to-side (contralateral) locomotor activity was not observed (data not shown). These results indicate that the population of spinal V3-INs do not play a significant role in establishing burst frequency (a proxy for tail-beat frequency) or coordination during fictive locomotion in larval zebrafish.

Ablation of V3-INs reduces the proportion of active MNs during spontaneous fictive swimming

Since V3-IN soma position (Fig. 2) and morphologic properties (Fig. 3) suggest that they project to the motor column in the spinal cord and targeted ablation of spinal V3-INs in zebrafish larvae did not affect burst frequency (Fig. 6), we tested the hypothesis that ablation of V3-INs decreases the number of active MNs during spontaneous fictive swimming. To ablate V3-INs without also photodamaging or bleaching MNs, we used a two-step neuronal targeting strategy. First, V3-INs were identified and ablated using the Tg(vglut2a:DsRed) nns9 transgenic line. Following recovery from laser ablation, MNs were loaded with a cell-permanent calcium indicator, Calcium Green-1 AM. To verify that spinal neurons loaded with AM dye (Calcium Green-1 and/or Rhod-2) were MNs, we first confirmed motor neuronal specificity of GFP expression in the pargamn2Etl enhancer trap line (Bertiuzzi and Ampatzis, 2018).

Parga mn2Etl larvae immunolabeled for the cholinergic marker ChAT revealed that most GFP-expressing neurons co-labeled with ChAT antibodies [n = 5; 91.3% (SD 2.1); Fig. 7A,A′′]. A small percentage of dorsally-positioned ChAT+ cells did not colocalize with GFP [4.1% (SD 2.8); Fig. 7A,A′′, arrowheads]; these cells were considered members of a subset of cholinergic spinal interneurons that modulate MN excitability recently identified in zebrafish (Bertiuzzi and Ampatzis, 2018). Thus, GFP expression in the parga mn2Etl enhancer trap line was specific to MNs.

Next, to confirm that loading of AM dye was restricted to MNs, the skin was removed from both sides of the body from parga mn2Etl larvae, in which GFP expression was shown to be limited to MNs (Fig. 7A,A′′), and MNs were labeled with Rhod-2 AM dye (n = 4; Fig. 7B,B′′). Rhod-2 AM, which was not used for calcium imaging experiments because it did not produce a strong calcium signal, was used as a proxy for Calcium Green-1 AM since the excitation/emission wavelengths of Calcium Green-1 AM overlapped with and could not be disambiguated from GFP expressed in parga mn2Etl MNs. The majority of GFP-expressing MNs co-labeled with Rhod-2 AM [84.3% (SD 4.3); Fig. 7B,B′′]. A small percentage of GFP-negative cells contained Rhod-2 AM [5.4% (SD 0.5); Fig. 7B,B′′, arrowheads]. Taken together, these results confirmed that MNs were selectively labeled with AM dye and, therefore, the cell permeable calcium indicator (Calcium Green-1 AM) could be used to monitor in vivo calcium activity in spinal MNs.

To examine the effects of V3-IN ablation on the proportion of spinal MNs active during spontaneous fictive swimming episodes, MN activity in Tg(vglut2a:DsRed) nns9 larvae was measured using in vivo calcium imaging by loading MNs with Calcium Green-1 AM, while fictive

Figure 5. Targeted ablation of V3-INs is both efficient and spatially restricted. A, B, Confocal stacks of the same Tg(vglut2a:DsRed) nns9 zebrafish larva before (A) and 12 h after (B) laser ablation of the V3-INs in the region indicated (red dashed lines). C, D, A magnified view of two V3-INs before (left) and after (right) a single cell laser ablation. Scale bars: 20 μm (A, B) and 5 μm (C, D).
locomotor activity was measured using extracellular PN recordings (Fig. 8A, B). We quantified the proportion of active MNs during spontaneous fictive swimming for Control (50 swim episodes, four larvae), Sham Ablated (14 swim episodes, two larvae), and V3-IN Ablated (59 swim episodes, three larvae) groups.

Assessment of the probability distributions showed that the proportion of active MNs in the Ablated group was significantly reduced compared with both the Control (Kolmogorov–Smirnov test, $D = 0.68$, $p < 0.001$) and Sham (Kolmogorov–Smirnov test, $D = 0.54$, $p = 0.001$) groups (Fig. 8C). The proportion of active MNs was not

Figure 6. Targeted ablation of V3-INs does not affect burst frequency during spontaneous fictive locomotion. A, Representative traces of extracellular peripheral motor nerve recordings from Control (top), Ablated (middle), and Sham (bottom) groups. Summary plots of cumulative probability distribution (B) and mean burst frequencies (C). Summary plots of cumulative probability distribution (D) and peak burst frequencies (E).
significantly different between the Control and Sham groups (Kolmogorov–Smirnov test, $D = 0.24$, $p = 0.50$; Fig. 8C). Further, V3-IN ablation had a main effect of significantly reducing the proportion of active MNs during spontaneous fictive swimming [Control mean = 64.6% (SD 17.0); Sham mean = 56.6% (SD 24.2); one-way ANOVA, $F = 25.3$; $p < 0.001$; Fig. 8D]. Pairwise multiple comparisons (Holm–Sidak; Fig. 8D) revealed significant differences for the proportion of active MNs between both the Control and Ablated groups ($t = 7.02$, $p < 0.001$) and the Sham and Ablated groups ($t = 2.98$, $p = 0.007$), but significant differences were not found between Control and Sham groups ($t = 1.54$, $p = 0.10$). These results indicate that V3-INs may participate in regulating locomotor amplitude by recruitment of spinal MNs during spontaneous fictive locomotion.

Discussion

The goal of this study was to characterize the potential role of V3-INs in regulating locomotor frequency, amplitude, or both, in genetically tractable and optically accessible larval zebrafish. Locomotor amplitude is determined by the recruitment of motor units appropriate to the desired movement: small units for fine behaviors and large units for large movements (Zajac and Faden, 1985; Mendell, 2005). This recruitment pattern of small to large motor units, the so-called size principle, is well established in both limbed (mammalian; terrestrial locomotion) and nonlimbed (swimming) locomotor systems (Henneman et al., 1965; Gustafsson and Pinter, 1984a,b; Cope and Pinter, 1995; McLean et al., 2007; Gabriel et al., 2011). However, investigations into the premotor neural mechanisms that contribute to the regulation of locomotor amplitude have been difficult in mammals for two primary reasons. First, mammalian spinal locomotor circuits are complicated by the requirements to control and coordinate limbs and joints, rather than simply produce a smooth gradient of motion intensity (Grillner, 2003; Grillner and El Manira, 2020). Second, despite recent progress in tool development, in vivo functional manipulations of mammalian spinal locomotor circuits using mini-scopes and fiber-optics are both challenging and limited for long-term recordings in freely locomoting animals (Montgomery et al., 2015; Samineni et al., 2017; Canales et al., 2018). Zebrafish

Figure 7. Spinal neurons loaded with rhodamine-2 AM dye are definitively identified as MNs. **A**, The EGFP-expressing pargam2Et enhancer trap line (A, green) was labeled with antibodies to choline acetyl transferase (ChAT; A’, magenta). Colabeling (A’) of EGFP with ChAT antibodies indicated that EGFP expression in the pargam2Et line is a marker of spinal MNs. White arrowheads indicate ChAT-positive cells that do not contain EGFP (non-MNs). **B**, Live pargam2Et larvae were anesthetized, skinned, and incubated in rhodamine-2 AM dye (Rhod-2, AM). EGFP expression (B, green) and Rhod-2, AM labeling (B’, magenta), showed that Rhod-2, AM specifically labeled all MNs (B’’). White arrowheads indicate two Rhod-2, AM-labeled cells that did not contain EGFP. Scale bar: 20 $\mu$m.
V3-INS are a population of previously uncharacterized ventral vglut2a+ neurons in larval zebrafish

Here, we report that almost all ventrally-located neurons labeled by the Tg(vglut2a: DsRed)\textsuperscript{nms9} and Tg(vglut2a: Gal4ff)\textsuperscript{nms20} transgenic lines are considered V3-INS (Figs. 1, 2). Since there are no available or published transgenic zebrafish lines that specifically label V3-INS (e.g., sim1a-expressing neurons) in larval zebrafish, we recognize that definitive identification of the spinal V3-INS reported here is indirect (Fig. 1). However, we are confident that this approach correctly identifies spinal V3-INS based on the convergence of characteristics, including co-expression of vglut2a and the p3 progenitor cell marker nkk2.2a (Briscoe et al., 2000; Schäfer et al., 2007), soma position of V3-INS in the p3 domain ventral to the central canal (Figs. 1, 2), and V3-INS neurotransmitter phenotype indicated by the overlap of vglut2a and sim1a expression in the zebrafish ventral spinal cord (Yang et al., 2010).

The V3-INS, a numerous ventromedial population, are distributed across the rostrocaudal axis of the spinal cord (Fig. 2). It is unlikely that the glutamatergic V3-INS include other identified ventrally-located spinal neurons, such as the GABAergic Kolmer–Agduhr cells (Dale et al., 1987; Bernhardt et al., 1992; Montgomery et al., 2016) or the ventral serotonergic neurons (Van Raamsdonk et al., 1996; Brustein et al., 2003; McLean and Fetcho, 2004; Montgomery et al., 2018). The observed diversity of cell morphology in the V3-INS cell population in the larval zebrafish spinal cord (Fig. 3) is similar to the diversity of V3-INS in the mammalian spinal cord (Borowska et al., 2013; Deska-Gauthier et al., 2020). In mammals, the morphology of V3-INS subtypes is correlated with their circuit function (Borowska et al., 2013). However, we have not found evidence for differential recruitment of V3-INS during fictive swimming at various frequencies (Fig. 4D), which would indicate potential functional heterogeneity. It may be that in larval zebrafish V3-INS are a primitive class of excitatory spinal interneurons, as has been previously proposed for Engrailed1-expressing inhibitory neurons in zebrafish (Higashijima et al., 2004). However, analysis of cell morphology in zebrafish larvae is complicated by ongoing development in the nervous system and the rostrocaudal developmental gradient (Cole and Ross, 2001). Additional experiments are necessary to clarify whether different morphologies represent distinct functional subtypes of V3-INS, developmental stages of a single morphologic family, or regional specializations of the V3-INS.

Zebrafish V3-INS are active during fictive locomotion

Results indicating that V3-INS activity is directly correlated to fictive locomotor activity in vertebrate preparations has not been convincingly presented. In mice, there is indirect evidence indicating temporal correlation between V3-INS activity and locomotion (Borowska et al., 2013). Through simultaneous optical and electrophysiological recordings, we showed that activity in most V3-INS is correlated to fictive swimming (Fig. 4) and is therefore appropriately timed to provide excitatory drive to MNs during locomotion. Thus, this is the first direct evidence in any vertebrate preparation that V3-INS are active during fictive swimming.
ficative locomotion, which provides supporting evidence that they may participate in locomotor activity.

**V3-in activity is necessary for MN recruitment, but not necessary for rhythm generation**

Given that V3-INs have been shown to affect the regularity and coordination of locomotor activity in mice (Zhang et al., 2008), it was surprising that laser ablation of V3-INs in larval zebrafish did not affect burst frequency during spontaneous fictive swimming in larval zebrafish (Fig. 6). One interpretation of these findings is that despite similarities in gene expression and neurotransmitter profile, premotor interneuron function is divergent between axial and tetrapod locomotion. The latter argument has been advanced regarding the V2a-INs (Dougherty and Kiehn, 2010; Dougherty et al., 2013); the different effects (or lack thereof) of V3-IN ablation in zebrafish and mice may be a second example of the trend. An important difference between the current study and mouse knock-out experiments is the mechanism of ablation. In the current study, V3-INs were ablated acutely, and little time (~24 h) was given for circuit compensation. In the mouse experiments, on the other hand, V3-INs were ablated or silenced chronically (Zhang et al., 2008; Rabe Bernhardt et al., 2012), leaving opportunity for physiological compensation. Acute silencing experiments using optogenetics could be performed in both species (Arrenberg et al., 2009; Hägglund et al., 2013), and may demonstrate that this difference between species is because of the experimental manipulation rather than the underlying neural circuit.

Finally, a subset of identified V2a excitatory spinal interneurons has been shown to control the speed of locomotion in larval and juvenile/adult zebrafish (Ausborn et al., 2012; Ampatzis et al., 2014). Of particular interest, recent studies also ascribed a role for regulating locomotor amplitude/vigor to a morphologically and functionally distinct subset of the V2a-INs (Type II, nonbursting) in larval and adult zebrafish (Song et al., 2018; Menelau and McLean, 2019). Taken together, these results indicate that the general population of V2a-INs is involved in regulating both locomotor speed and amplitude. Importantly, a direct role of V2a-INs in swimming behavior in vivo was not shown in either study. To date, however, a population of spinal interneurons that selectively regulates locomotor amplitude has not been identified. Our work provides evidence that, in zebrafish larvae, V3-INs do not contribute to locomotor speed (burst frequency). Specifically, we show that targeted ablation of V3-INs (Fig. 5) reduces the proportion of active MNs during fictive swimming (Fig. 8) but does not affect the range of locomotor frequencies produced (Fig. 6). Thus, we propose that V3-INs are a source of excitation in the vertebrate locomotor neural circuitry that may regulate the amplitude of locomotor output independently of locomotor frequency.

In conclusion, the work presented here addressed an important concept regarding motor control; specifically, a limited understanding of the cellular properties that produce locomotor amplitude in vertebrates. We showed, directly for the first time, that V3-INs in zebrafish larvae are active during in vivo fictive locomotion (Fig. 4) and contribute to MN recruitment (Fig. 8). Importantly, targeted ablation of the V3-IN spinal cord population does not affect locomotor frequency (speed; Fig. 6), which suggests a role in motor control rather than rhythm generation. Thus, we propose that the V3-IN population may regulate locomotor amplitude independently of frequency, which is an important functional difference between the V3-INs and the general population of V2a-INs since the latter regulate both frequency and amplitude (Ausborn et al., 2012; Ampatzis et al., 2014; Song et al., 2018; Menelau and McLean, 2019).

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