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Pan-neuronal imaging in roaming *Caenorhabditis elegans*

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We present an imaging system for pan-neuronal recording in crawling *Caenorhabditis elegans*. A spinning disk confocal microscope, modified for automated tracking of the *C. elegans* head ganglia, simultaneously records the activity and position of ~80 neurons that coexpress cytoplasmic calcium indicator GCaMP6s and nuclear localized red fluorescent protein at 10 volumes per second. We developed a behavioral analysis algorithm that maps the movements of the head ganglia to the animal's posture and locomotion. Image registration and analysis software automatically assigns an index to each nucleus and calculates the corresponding calcium signal. Neurons with highly stereotyped positions can be associated with unique indexes and subsequently identified using an atlas of the worm nervous system. To test our system, we analyzed the brainwide activity patterns of moving worms subjected to thermosensory inputs. We demonstrate that our setup is able to uncover representations of sensory input and motor output of individual neurons from brainwide dynamics. Our imaging setup and analysis pipeline should facilitate mapping circuits for sensory to motor transformation in transparent behaving animals such as *C. elegans* and *Drosophila* larva.

C. elegans | *Drosophila* | calcium imaging | thermotaxis

Understanding how brain dynamics creates behaviors requires quantifying the flow and transformation of sensory information to motor output in behaving animals. Optical imaging using genetically encoded calcium or voltage fluorescent probes offers a minimally invasive method to record neural activity in intact animals. The nematode *Caenorhabditis elegans* is particularly ideal for optical neurophysiology owing to its small size, optical transparency, compact nervous system, and ease of genetic manipulation. Imaging systems for tracking the activity of small numbers of neurons have been effective in determining their role during nematode locomotion and navigational behaviors like chemotaxis, thermotaxis, and the escape response (1–6). Recordings from large numbers of interconnected neurons are required to understand how neuronal ensembles carry out the systematic transformations of sensory input into motor patterns that build behavioral decisions.

Several methods for fast 3D imaging of neural activity in a fixed imaging volume have been developed for different model organisms (7–14). High-speed light sheet microscopy, light field microscopy, multifocus microscopy, and two-photon structured illumination microscopy have proved effective for rapidly recording large numbers of neurons in immobilized, intact, transparent animals like larval zebrafish and nematodes (15–19). However, these methods are problematic when attempting to track many neurons within the bending and moving body of a behaving animal. Panneuronal recording in moving animals poses higher demands on spatial and temporal resolution. Furthermore, extracting neuronal signals from recordings in a behaving animal requires an effective analysis

pipeline to segment image volumes into the activity patterns of discrete and identifiable neurons.

Here, we use high-speed spinning disk confocal microscopy—modified for automated tracking using real-time image analysis and motion control software—to volumetrically image the head ganglia of behaving *C. elegans* adults at single-cell resolution. Our setup can simultaneously track ~80 neurons with $0.45 \times 0.45 \times 2\text{-}\mu\text{m}$ resolution at 10 Hz. Activity was reported by the ultrasensitive calcium indicator GCaMP6s expressed throughout the cytosol under the control of the pan-neuronal *rgef-1* promoter (a gift from D. Pilgrim, University of Alberta, Edmonton, Alberta, Canada) (20). To facilitate segmentation into individual identifiable neurons, nuclei were tracked using calcium-insensitive, nuclear-bound red fluorescent protein (RFP), TagRFP, under the control of another pan-neuronal *rab-3* promoter (a gift from O. Hobert, Columbia University, New York) (21). We developed an image analysis pipeline that converts the gross movements of the head into the time-varying position and posture of the crawling worm, and converts fluorescence measurements into near simultaneous activity patterns of all imaged neurons.

A similar approach to brainwide imaging in moving *C. elegans* using the same transgenic strain has recently been reported (22). Although both setups use customized spinning disk confocal microscopes, the strategies for tracking the moving neurons and analyzing behavioral and neural activity patterns are different. Nguyen et al. (22) use a low power objective to track the posture of the animal and a high power objective to locate and image the

Significance

A full understanding of sensorimotor transformation during complex behaviors requires quantifying brainwide dynamics of behaving animals. Here, we characterize brainwide dynamics of individual nematodes exposed to a defined thermosensory input. We show that it is possible to uncover representations of sensory input and motor output in individual neurons of behaving animals. Panneuronal imaging in roaming animals will facilitate systems neuroscience in behaving *Caenorhabditis elegans*.

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See Commentary on page 1965.

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nerve ring. The advantage of our single objective setup is that it affords the flexibility, for example, to deliver thermosensory inputs using an opaque temperature controlled stage below the animal. The advantage of low-magnification imaging is that it provides a direct measurement of animal posture, which we must infer. These new technologies for pan-neuronal imaging in roaming animals now enables correlating brainwide dynamics to sensory inputs and motor outputs in transparent behaving animals like *C. elegans* and *Drosophila* larvae.

Results

Automated Volumetric Tracking of Head Ganglia in a Moving Worm.

We built our imaging system into a spinning-disk confocal microscope (Fig. 1A). A custom-designed thermoelectric cooler was used to sinusoidally modulate the temperature of the stage over the course of an experiment to provide a defined thermosensory stimulus (Fig. 1B) (23). Individual young adult *C. elegans* were placed on a thin sheet of agar, covered with a large coverslip, and allowed to crawl between the agar and coverslip. A fast scientific complementary metal-oxide semiconductor (CMOS) camera (200 frames/s) and piezoelectric objective scanner (bidirectional rastering at 2 $\mu\text{m}/\text{step}$ with 20 steps) provided an imaging rate of 10 volumes/s. We captured the image of each z section in both red and green fluorescence emission channels using a dichroic to simultaneously project each channel onto different regions of the camera's sensor. When imaging with a 40 \times , 0.95-NA air objective; each volume represents 115 \times 115 \times 40 μm with pixel-limited resolution of 0.45 \times 0.45 \times 2 μm . These imaging volumes and speeds were made possible using custom software written in MATLAB to control the microscope hardware, carrying out all functions including image capture, motion control, stimulus control, and data streaming.

A motorized stage kept a targeted image volume in the field of view at all times. This tracking was achieved by real-time analysis of image volumes that sought and recognized the head ganglia in the green (GCaMP6s) channel. Fluctuations in the coordinates of the target in x , y , and z were used as difference signals to adjust the voltages applied to the stage and objective drivers to automatically keep the target in the center of the acquired volume. Feedback in xy ran at 10 Hz (every volume), and feedback in z ran at

2 Hz (every five volumes). The system was able to continuously image head ganglia of a crawling animal for up to 5 min without missing frames. The tracking system is described in detail in *Methods*.

Reconstructing a Worm's Motor Behavior from Head Movement. To identify neural correlates of locomotory behavior, it is important to know both the posture and position of the body over time. Our setup tracks the position and orientation of the head ganglia of a crawling worm at high magnification. Some setups for tracking neural activity at high resolution implement a second low-magnification objective for simultaneous imaging of the worm body (24, 25). The advantage of single objective experimental designs is a simpler optical path that allows imaging the worm on an opaque substrate like our thermally controlled platform. A possible concern of single-objective imaging at high magnification is that our body measurement is indirect.

When worms crawl on an agar surface, the undulating body follows the contours of the groove made by the head during forward movement, and the head follows the groove made by the body during backward movement (26). Thus, the posture of the animal can largely be estimated from the trajectory of the head.

First, we converted the $\{x(t), y(t)\}$ position of the head ganglia into coordinates for locomotion, $\{v(t), \omega(t)\}$, corresponding to the forward velocity of the worm ($\mu\text{m}/\text{s}$) and the angular velocity of the head in the ventral direction (rad/s). Next, we propagated the translational and curving movements of the head into the translation and curvature of all points along the body using phase lags dependent on fractional distance from the head, projecting all measurements onto the four-dimensional subspace of worm locomotory postures (called eigenworms) to predict the smoothly time-varying posture of the body (27, 28). We developed this algorithm (*Methods*) to perform this estimation based on the 2D trajectory of the head ganglia recorded by our tracking microscope. We verified the ability of our algorithm to correlate head coordinates with body posture using low magnification imaging (4 \times). Using exclusively the velocity of the head ganglia, we were able to reconstruct the posture of a worm at all points along a trajectory (Fig. 2A and *Movie S1*). The trajectory and posture of an animal that was reconstituted solely from high-resolution recordings (40 \times) are also shown (Fig. 2B).

To quantify the correctness of our algorithm, we tested it against an online repository of postural analysis of freely crawling worms (28). Plugging the time-varying position of a coordinate fixed to the location of the head along the body into our algorithm, we reconstructed the complete posture of the animal. We were then able to compare the reconstructed posture to the documented posture available on the online repository. We calculated correlation coefficients at each point along the body. Our algorithm does not capture the variance in the movements anterior to the head ganglia. This discrepancy is expected because the high-frequency exploratory movements of the worm's nose are largely uncorrelated with the movements of the body. However, our algorithm effectively captures much of the variance of the rest of the worm body (Fig. 2C). Because the estimate provided by our algorithm is not perfect, it may not be useful for applications where fine postural information is required, but may be useful in studies of navigational behavior, for example, where strategies are largely based on motor decisions that originate within the head and can be inferred from the movements of the head (dorsal or ventral bending for steering, or forward and backward movement).

Semiautomated Registration of Individual Nuclei in the Moving Head Ganglia.

As an animal crawls, the relative location of neurons will change, sometimes dramatically, over the course of image acquisition (*Movie S2*). Without fiducial markers that capture this changing geometry, it is impossible to identify regions that correspond to signals from specific neurons. One clever solution was to

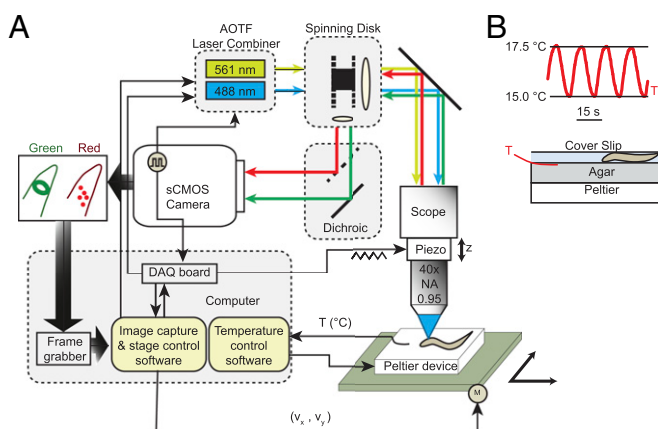


Fig. 1. Experimental setup. (A) Microscopy schematic. Data were acquired using a spinning disk confocal microscope with red and green channels captured side by side on a CMOS sensor at 200 frames/s. A volume was imaged every 10 s using a piezo objective scanner. Every 20 frames were grouped together to form an image stack. Custom software controlled the stage and objective movements to maintain the tracked target at the center of the image volume. (B) In all experiments, the animal was provided with a sinusoidal temperature stimulus using a custom thermoelectric stage (described in ref. 23). Temperature was recorded using a microthermocouple embedded at the agar surface.

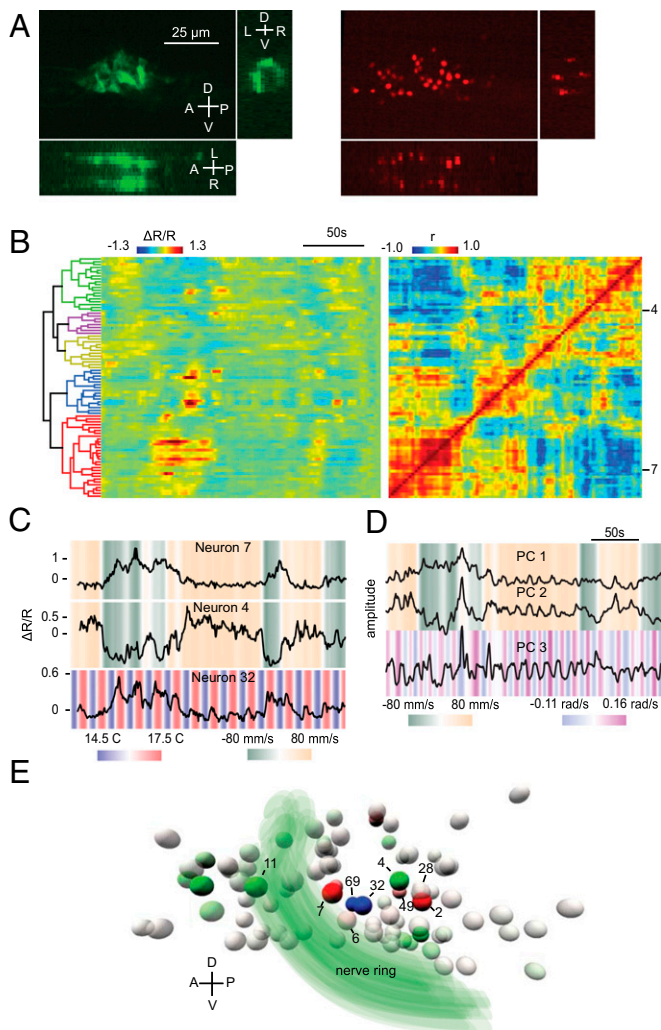


Fig. 3. Multineuronal activity patterns. (A) Images obtained in a 5-ms exposure with our setup in the red channel showing TagRFP and green channel showing cytosolic GCaMP6s. Each image is a slice through the acquired volume. Neurons are resolvable in the green channel, and nuclei are well resolved in the red channel in all dimensions. Signals are attenuated further from the objective. (B) Normalized calcium dynamics of 84 neurons and matrix of correlation coefficients from the trajectory shown in Fig. 2B. Neuronal activity patterns are grouped and ordered by agglomerative hierarchical clustering. Neurons corresponding to the two largest clusters are highlighted in red and green on the tree diagram. Indexes corresponding to the neurons with the strongest signals within each cluster are indicated. (C) Sample traces from a cell activated during forward movement (#7), a cell activated during backward movement (#4), and a cell correlated with temperature oscillations (#32). (D) The first three principal components of the whole-brain neuronal activity correlate most strongly with behavioral output. The first two correlate with velocity, and the third component reports the angular velocity of the head. (E) A volume showing the relative location of segmented neurons in one image volume. Neurons corresponding to the green and red clusters in Fig. 3B are colored accordingly. Neurons discussed in this figure along with Fig. 4 are annotated for reference. Two neurons correlated with thermosensory input are colored in blue. The position of the nerve ring is drawn for reference. The right side of the animal is into the page.

variance in brainwide neuronal dynamics, appear to be representations of worm directionality. The third principal component, which encoded 9%, represented the angular displacement of the head.

Assigning Neuronal Identities to Indexed Nuclei. Unambiguous identification of the neuronal identity is essential for interpreting their

activity patterns. Lineage tracing using differential interference contrast (DIC) microscopy has shown that most cells (i.e., neuronal somas) in *C. elegans* are found in reproducible relative positions (30, 31). Assuming stereotypy, each automatically assigned index should correspond to a unique neuron. However, for neurons that could have variable positions, the same cell may be assigned a different index in different animals. We asked whether our analyses can uncover activity patterns of specific cells with stereotyped locations by performing pan-neuronal recording and subsequent image analyses in three additional crawling worms (Fig. 4A and B). Across all four animals, indexes 32 and 69 were assigned to two neurons that responded to temperature changes. Indexes 2, 6, 7, 28, and 49 were assigned to neurons that were activated by backward movement in all animals. Indexes 4 and 11 were assigned to neurons that were deactivated by backward movement. Because all index assignments were made blind to calcium dynamics, this result highlights the consistency of activity patterns and neuronal location across animals. To demonstrate the fraction of our signal that may be due to noise from the acquisition and analysis procedure, we prepared a worm with GCaMP6s replaced by GFP and collected traces from moving animals studied in the same way (Fig. 4B). In control animals, no significant temperature or motion correlated signals were observed in any neurons.

We sought the identity of neurons that exhibited strong calcium signals with high stereotypy. The unique morphology of *C. elegans* neurons could be readily observed by DIC microscopy. By examining high-resolution DIC images of an adult animal from the pan-neuronal imaging strain, overlaid with the red and green fluorescence signals from the neuronal reporters, we were able to identify cells that correspond to the nine most highly stereotyped signals across the four animals (Fig. S1). The two neurons correlated with temperature inputs across the animals were the AFD thermosensory neurons: both AFDL (index 32) and AFDR (index 69) are activated by warming and deactivated by cooling during a sinusoidal temperature variation, consistent with previous reports (Fig. 4B) (32). The neurons consistently encoding forward locomotion were the AVB (index 4) premotor interneuron and the RMEL (index 11) head motor neuron. The consistent backward coding neurons were the AIBR (index 49) interneuron, and the AVAL (index 7), AVDL (index 28), and RIML (index 2) premotor interneurons. All of these activity patterns were consistent with previous reports of their activity patterns in moving animals (1, 2, 6, 33–36). The activity traces of all segmented neurons in the animals that we studied are shown in Fig. S2. We note that most neurons identified were located on the left side of the animal, where we recorded the strongest signals because the four animals crawled on their right sides (Fig. 3A).

We only observed a representation of sensory input in two neurons: AFDL (index 32) and AFDR (index 69). We asked whether activity patterns of additional neurons that correlated to temperature might be found by eliminating movement altogether. To do this, we completely immobilized worms from our pan-neuronal imaging strains using microbeads (37). With immobilized neurons, the ability to segment activity is improved, so we can determine whether additional neurons with significant temperature correlated responses might have escaped our analysis in moving worms. We calculated the correlation coefficient of each neuronal trace throughout the nerve ring of immobilized animals and again found that the two neurons with the strongest temperature correlated responses were AFDL (index 32) and AFDR (index 69) (Fig. S3). In the immobilized worm, small but significant correlation was also found in two neurons in the anterior ganglion that we were unable to unambiguously identify. Nevertheless, these results support our conclusion that the representation of sensory input is largely localized to the AFD neurons.

Tracking Thermosensory Neurons in the Crawling *Drosophila* Larva. Single neuron tracking in moving *C. elegans* has been made tractable by the

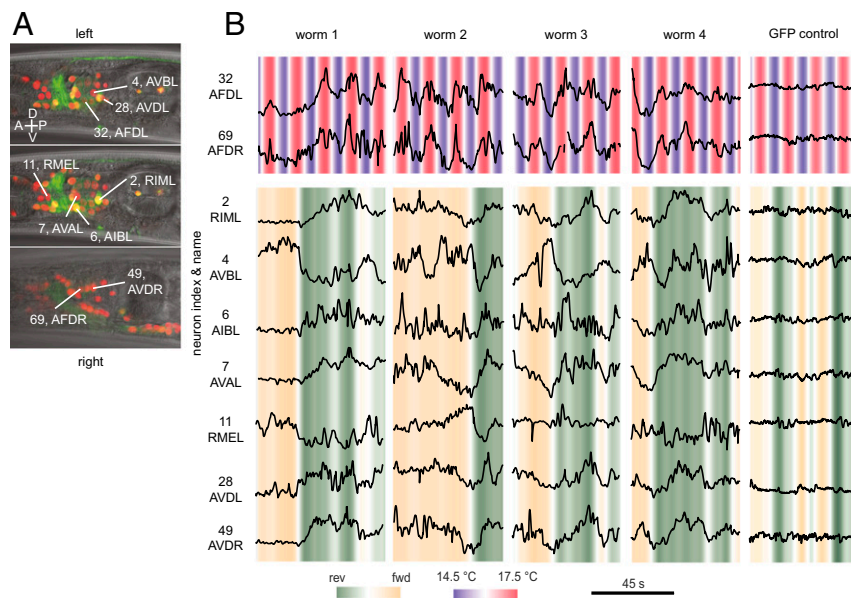


Fig. 4. Stereotyped responses of indexed neurons. (A) High-resolution stacks with simultaneous red (nucleus), green (cytoplasm), and DIC information were used to match the indexes with stereotyped activity patterns to their cellular identities (also see Fig. S1). (B) Consistent signals across animals from temperature-sensitive neurons (32-AFDL and 69-AFDR) and several neurons correlated with forward/backward movement, suggesting that the indexes assigned using DIC are the same as those recorded in roaming animals. Four worms (1–4) represent our standard imaging conditions with pan-neuronal GCaMP6s. The GFP control worm was subjected to the same imaging and analysis procedure, but using a transgenic worm that expressed pan-neuronal GFP. The residual signal in the control worm provides an estimate of total noise in the system caused by experimental measurement and the analysis pipeline.

fact that their movement is largely restricted to two dimensions. It has not yet been possible to record calcium dynamics in other unrestrained animals such as *Drosophila* larva owing to their faster movements in all dimensions. Because *Drosophila* larva crawl using peristalsis waves, neurons undergo substantial z axis movements with each wave. When the larva picks a new direction during forward movement, it lifts and swings its head until it selects a new direction (38, 39). Because our system is capable of both 3D tracking and volumetric imaging, we asked whether it could meet the challenge of capturing calcium dynamics of neurons in the head of crawling *Drosophila* larva.

We recently discovered three thermosensory neurons in the dorsal organ of *Drosophila* larva that are required for positive thermotaxis up temperature gradients (23). Like the AFD neuron in *C. elegans*, they can phase-lock calcium dynamics to an oscillating thermosensory input (32). We used the *R11F02* Gal4 driver line to specifically target the express of both GCaMP6s and RFP in the cytosol of these sensory neurons and monitored their calcium dynamics in a crawling larva subjected to sinusoidal temperature oscillations.

During a bout of spontaneous quiescence during the temperature cycle, we observed intracellular calcium signals that were characteristic of the sensory-evoked response of the thermosensory neurons. During episodes of head swinging, where these neurons moved dramatically in an xyz trajectory, our system was able to track them within the image volume. We found identical calcium dynamics precisely phase-locked with temperature changes, as during the quiescence period. Therefore, the calcium dynamics of these sensory neurons is a representation of sensory input without significant correlation to movement (Fig. S4).

Discussion

Recent years have seen rapid development in multineuronal imaging systems in diverse preparations (13). A formidable challenge is to simultaneously observe brain-wide activity and behaviors of animals with defined sensory inputs, the only scenario where neuronal activity patterns can be properly correlated with sensory

inputs and behavioral outputs. Our setup has uncovered patterns in sensorimotor transformation in the *C. elegans* nervous system that both buttress and extend recent findings from single or small number of neuron tracking in behaving animals, and brain-wide imaging in immobilized animals. The AFD neuron has been shown to be the dominant thermosensory neuron in *C. elegans*, required for all modes of thermotaxis (40, 41). Although spontaneous activity in other sensory neurons have been shown to be modulated by temperature changes, only AFD phase locks its calcium dynamics to a rapidly oscillating temperature waveform in a behaving animal (32, 42–44). Our recordings confirm that the thermosensory response is largely localized to AFDL and AFDR.

Brain-wide imaging of unstimulated and immobilized worms has revealed spontaneous activity that are clustered into groups with correlated and anti-correlated patterns (16, 17). Neurons that exhibited activity increase during periods of backward movement by single neuron tracking in behaving animals (e.g., AIB and AVA) exhibited strongly correlated activity pattern in the pan-neuronal imaging setup in immobilized animals. Thus, periods of fictive forward or backward movement might be inferred from the activity patterns of different groups of neurons. The advantage of our system is the ability to make a direct correlation between brain dynamics and behavioral dynamics from sensory input to motor output.

A limitation of recording calcium dynamics in regions surrounding nuclei is that we are restricted to somatic calcium transients. Several neurons in *C. elegans* exhibit less calcium activity in their soma, with higher activity compartmentalized to their processes (5, 32, 45). Because processes of the head ganglia are tightly bundled fascicles, our current setup does not allow isolation of signals corresponding to individual neuronal processes. In these cases, sparse labeling of smaller groups of neurons is required to quantify their calcium dynamics. Our system may still be useful in these studies, however, as it would deliver a 3D recording of calcium dynamics throughout a cell, allowing comparison and contrast of calcium dynamics in different cellular compartments.

Our system now allows the extraction of correlations between neurons in the head ganglia and their correlations with sensory input and motor output. Systematic analysis of brain-wide activity patterns during complex behaviors like thermotaxis and chemotaxis may uncover characteristics in the information processing and decision-making process during the sensory to motor transformation. Comparison of brain-wide activity patterns in animals with specific perturbations to the nervous system, e.g., by laser ablation or genetic inactivation of specific neurons, should reveal cellular and synaptic mechanisms for sensory to motor transformation.

Methods

Molecular Biology. GCamp6s was PCR amplified from pGP-CMV-GCaMP6s (Addgene) with BamHI and NotI restriction sites and subcloned in pCB101 behind the *rgef-1* promoter (gift of D. Pilgrim) (20). We cultivated transgenic worms at 22 °C on nematode growth medium (NGM) plates with OP50 bacteria. *zfls124* was generated by injection of Prgef-1::GCaMP6s at 50 ng/μL into *lin-15(n765ts)* worms along with the *lin-15* rescuing plasmid (pL15 EK) at 50 ng/μL. The extrachromosomal array was integrated using gamma-ray irradiation and outcrossed four times to WT N2. Strain QW1217 was made by combining *zfls124* with *otIs355(Prab-3::NLS::tagRFP)* (gift of O. Hobert) (21). Strain QW1473 (*rgef::GFP* and *rab3::NLS::tagRFP*) was made by combining strain *evIs111(Prgef::GFP; dpy-20)* with *otIs355(Prab-3::NLS::tagRFP)* (46).

Behavioral Measurements, Microscopy Hardware, and Software. The day before behavioral measurements, *C. elegans* were moved to a 15 °C incubator to ensure that the thermotactic setpoint was in a range that would generate a calcium signal in our experimental setup where we subjected animals to sinusoidal temperature waveforms between 15 °C and 17.5 °C. *Drosophila* larvae were cultivated and handled as described recently before being introduced to the experimental setup (23).

The setup is built around a Nikon Eclipse LV100 upright microscope. We imaged worms using a 4×, 0.2-NA Nikon Plan Apo objective (Fig. 2A) or the nerve ring at high resolution using a 40×, 0.95 NA Nikon Plan Apo Lambda objective. Worms were placed on 3-mm-thick 2% agar surfaces and covered with a 48 × 65-mm #1 coverslip. We captured dual color images of a 115 × 115-μm field of view by projecting both images obtained using a Yokogawa CSU22 spinning disk confocal setup onto the sensor of an Andor Zyla 4.2 sCMOS camera using an Andor Optosplit II with an mCherry/GFP dichroic cube. The microscope stage was controlled by a Ludl BioPrecision2 XY motorized stage and MAC 6000 stage-controller, a PI P-721.LLQ high speed piezo, and a Ludl precision stepper motor with a linear encoder. During data acquisition, computer software kept the worm centered in the field of view via an automated feedback loop that provided real-time image analysis and motion control. All data were collected with 4 × 4 binning and a vertically centered 256 × 512 pixel area of interest. The left half of this area consisted of the red output from the dichroic and the right half consisted of the green output. The measured pixel size of the system was 0.45 μm.

During acquisition, the camera provided the primary clock, running at 200 Hz, with one frame of data generated per clock cycle. Volumes consisted of 20 frames spaced 2 μm apart. The rastering triangular waveform was modified to optimally compensate for hysteresis due to the inertial load of the microscope objective at these fast scanning rates. Optimal compensation was strongly objective dependent. Feedback in *xy* could occur after the acquisition and segmentation of each volume to find a bright centroid. However, feedback in *z* involved latency due to buffering of the voltage being output to the piezo. This buffering resulted in a feedback time of roughly 0.5 s in typical operation. Image analysis and motion control were managed by custom software written in MATLAB, which was run in a Windows 7 environment with real-time priority. The code will be made freely available on www.github.com on publication.

Postural Reconstruction Algorithm.

Inputs.

- $\hat{n}(t)$, the unit vector specifying the forward direction, evenly sampled
- $\vec{x}(t)$, the trajectory of the nerve ring, sampled with \hat{n}
- L , the length of the worm (in the same units as \vec{x})
- θ , an indicator of whether the worm is lying on its left or right side.

Output.

- $P(t)$, an array of the estimated *x* and *y* coordinates for each of 100 segments

A worm's posture is stored as a series of 100 ordered pairs evenly spaced between the animal's nose and tail. The animal's head is assumed to be point 15. Whether the worm was lying on its left and right side was evident based on the ventral position of its vulva and the overall geometry of the nerve ring.

A worm's posture is reconstructed by storing the history of the worm's head's location on a stack (length parametrized). When the worm moves forward, points are pushed to the stack. As it moves backward, points are popped. To determine the posture at a given time, the top 0.85L of the stack is projected on to the nearest eigenworm (28).

To check for goodness of fit, we downloaded data from an online database corresponding to N2 worms grown off food (26 animals, up to 15 min/animal) (28). Worms were split into 100 segments, and angles between neighboring segments were calculated. The variation explained in Fig. 2C is the square of the Pearson correlation coefficient between predicted and true angles.

Identifying Nuclear Positions. Nuclei in the red imaging channel were tracked using a combination of manual annotation and standard image registration techniques. The processing pipeline was as follows:

- i) Separate red and green volumes from each frame. Identify sporadic bad image volumes using coarse image statistics. Rotate and crop volumes to a standard size centered on the nerve ring using maximum intensity projections from the red and green channels.
- ii) Assign consistent indexes to nuclei in one or more reference frames. To obtain consistent indices across animals, we manually compared images between a registered index-labeled library and the unlabeled data set being analyzed. When neurons are in stereotyped positions, each index would thus be uniquely assigned to a specific cell type.
- iii) Select a nucleus for registration (the active nucleus). We want to ensure that frames are aligned to similar frames, and we want to avoid long-range error propagation. Registering frames sequentially in time can result in poor reference frame choices when the worm is moving dramatically. Often, we want to register a frame to a temporally distant but conformationally similar frame. Additionally, frames that are similar in the context of one nucleus may not be similar in the context of another due to long-range distortions through the animal's brain. Therefore, one should search for a different optimal registration order for each nucleus. To accomplish this, we used the following tree-based procedure for each active nucleus. (a) Identify up to four nearby completely registered nuclei (reference nuclei); (b) Using the coordinates of these reference nuclei, calculate a pairwise distance between frames as the L_1 norm of the differences of their coordinates. In the typical case of four reference nuclei, this would be the mean of the absolute values of 12 coordinate differences. (c) Use these pairwise distances to generate a complete, weighted, undirected graph for the frames. Insert a dummy node into this graph with zero-weight connections only to the frames where the active nucleus has already been registered (either manually or from a previous iteration of the registration routine). (d) Compute a minimum spanning tree for this graph rooted at the dummy node. By design, the children of the root node will be the previously registered frames. (e) Perform a breadth-first search of the tree to generate the active nucleus's registration order along with a reference frame for each unregistered frame (parents in the registration tree).
- iv) Compute the location of the active nucleus in each frame (ordered as described above). Use the following method (steps *e*, *f*, and *g* are partly redundant, but they increase the robustness of the method). (a) Identify the coordinates of the reference nuclei in the current frame's reference frame. (b) Estimate a rigid transformation that will map the positions of the reference nuclei from the reference frame to the current frame. (c) Compare maximum-intensity projections in *x*, *y*, and *z* around a small volume centered around the preimage (in the reference frame) and image (in the current frame) of the transformation found in *b*. (d) Perform pairwise image registration between the maximum intensity projections to obtain three transformations between the reference frame and the current frame. (e) Apply these transformations to the preimage coordinates of our active nucleus to obtain a new guess for its location in the current frame. The guesses for *x* and *y* are taken from the *z* projection; the guess for *z* is the average of the values returned from the *x* and *y* projections. (f) Search for a bright point near the current guess and record its position. Gaussian masks are used to enforce an elastic penalty that keeps the chosen location near the current guess and away from previously registered nuclei. (g) Mark the nucleus as registered.

- v) Perform proofreading to ensure that nuclei are correctly registered. Manual proofreading was performed for all neurons reported here (the 84 traces in Fig. 3B and the additional traces shown in Fig. 4B).

To visualize the quality of the locally rigid deformation that this algorithm implies (and identify regions that are being incorrectly warped), we can take several frames and fix their parent to some reference frame. Then we can apply our algorithm to each pixel in the original set of frames to see how the image is being deformed. This deformation is presented in Fig. 2D.

Extracting Calcium Dynamics. To extract signals, we first computed the mean pixel values $F(t)$ in $2 \times 2 \times 4\text{-}\mu\text{m}$ regions around each nucleus in the green channel. A slowly varying background $F_0(t)$ was computed and subtracted for each color channel to obtain the normalized signal: $f_g(t) = [F(t) - F_0(t)]/F_0(t)$. The same procedure was applied to the red channel to obtain $f_r(t)$. To mitigate motion artifacts that produce correlated signals between the red and green channels, we subtracted the correlated component between the red and green

channels quantified by linear regression on a neuron-by-neuron basis. Pairwise similarity between neuronal signals was measured using Pearson correlation coefficients, and complete-linkage hierarchical clustering was performed with the resulting pairwise distances. Finally, principal components were calculated using the singular value decomposition method (without additional centering of data). We imaged worms for 1–3 min each and selected from each a 60-s interval to analyze in detail for calcium activity. This segment was selected on the basis of behavior such that it would include periods of both forward and reverse motion. No selection was made on calcium activity, and all extracted data are presented in Fig. S2.

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