Supplemental Information

An Imbalancing Act: Gap Junctions Reduce the Backward Motor Circuit Activity to Bias C. elegans for Forward Locomotion

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Figure S1

A moving transgenic C. elegans expressing Ca²⁺ sensors

Real-Time Processing
- raw fluorescent images
- image processing
- stage control
- XY-coordination
- image data

Post Imaging Processing
- object tracking
- fluorescent intensity measurements
- ratiometric measurements
- motion measurements

B

wild-type

AVA

YFP

CFP

YFP/CFP

Time (s)

C

wild-type

AVA

RIM

YFP

CFP

YFP/CFP

Time (s)

D

wild-type

VA8

VB9

YFP

CFP

YFP/CFP

Time (s)
Figure S1. Calcium imaging in C. elegans premotor interneurons and motoneurons

A. A schematic diagram of the imaging system for simultaneous calcium imaging and motion, and a flow chart of image processing and data analysis. Image capture, stage control and data storage are processed in real-time to track moving objects during recordings with in-house developed software. Raw data are processed post image capture by other customized algorithms to obtain profiles for neuronal activity and movement; B, An imaging profile of AVA in wild-type animals expressing a genetic calcium sensor cameleon. YFP and CFP signals show reciprocal changes when YFP/CFP ratio increases rapidly in the AVA premotor interneurons, indicating that these ratio changes reflect true FRET responses; C, Simultaneous calcium imaging of RIM and AVA showed similar calcium transient profiles, which supports their co-activation; D, Left panel, calcium transients of VB9 (red trace) and VA8 (blue trace) in a moving wild-type animal as shown in Figure 2A. This animal frequently switched directions during the recording. The large YFP/CFP ratio changes exhibited by VA8 (the blue trace in the left panel), often coinciding with switching directions, reflected FRET signals, as shown by reciprocal changes between YFP and CFP signals. a.u. arbitrary unit.
Figure S2

A

B

Duration of movement (s)
0 10 20 30

% of Time
0 20 40 60 80 100

wild-type

unc-7

control

+ silence A

fwd (≥1 pixel)
bwd (≥1 pixel)
pause (<1 pixel)

Duration of movement (s)
0 10 20 30

% of Time
0 20 40 60 80 100

wild-type

control

+ silence B

+ silence A/VB

fwd (≥1 pixel)
bwd (≥1 pixel)
pause (<1 pixel)
Figure S2. A, Restoring the B>A pattern rescued continuous forward motion in innexin mutants. The averaged duration time for forward and backward movements (upper graphs), and the total percentage of time, or the propensity of animals in forward and backward motion or pause (lower graphs) for each genotype. *unc-7* animals expressing the A motoneuron silencing transgene (*Punc-4::TWK-18(gf)*) exhibited an increased duration of continuous forward movement and decreased duration for backing (upper graphs), as well as an increased propensity for forward motion and a reduced propensity for backing (lower graphs). B. Reducing B motoneuron output in wild-type animals reduced forward motion and increased backing. Wild-type animals with a reduced B motoneuron output, by reducing either B activity (*Pacr-5::TWK-18(gf)*), or AVB activity (*Psra-11::TWK-18(gf)*), exhibited a decreased duration of continuous forward movement (upper graphs), a decreased propensity for forward movement, concomitant with an increased propensity for backing (lower graphs). Propensity analyses were carried out using pooled data from animals (N>10) per genotype. ***P<0.001, **P<0.01, *P<0.05 by the Mann-Whitney U test. Error bars: SEM.
Figure S3

(A) AVE

Ca\textsuperscript{2+}-insensitive control

wild-type

unc-7

unc-9

Time (s)

YFP/CFP

(A') A'

B

Ca\textsuperscript{2+}-insensitive control

wild-type

unc-7

Time (s)

YFP/CFP

(A'') A''

B

Ca\textsuperscript{2+}-insensitive control

wild-type

unc-7

Time (s)

YFP/CFP

(B) AVA

Cameleon

Tubulin

Pglr-1

Pnmr-1

Psra-11

D

Density (% of WT)

72 kD

54 kD

Pglr-1

Pnmr-1

Psra-11

wild-type

unc-7

unc-9

wild-type

unc-7

wild-type

unc-7

wild-type

unc-7

wild-type

unc-7

wild-type

unc-7
Figure S3. Imaging of AVE and AVA separately under a restricted movement condition also showed an increased activity of these interneurons in *unc-7* mutants

A-B, Representative calcium transient traces in AVE (A) and AVA (B) of each genotype recorded under identical experimental conditions; A'-B', Raster plots of calcium traces, each horizontal line corresponding to the YFP/CFP ratio of an animal of the respective genotypes as in the left panels during five minute recording, pseudo-colored according to the calcium transient level over time in AVE (A') and AVA (B'); A''- B'', Mean and distribution of the calcium transient level of each genotype in AVE (A'') and AVA (B''). Each data point represents the averaged YFP/CFP ratio during the five-minute recording for each animal. N for each genotype is shown in brackets. *** p < 0.001, ** p < 0.01, * p < 0.05 by the Mann-Whitney U test; C-D, Expression levels of calcium sensors did not changed in *unc-7* and *unc-9* mutant background. (C) a representative result for western blot analyses of synchronized strains expressing interneuron cameleon calcium sensors used in this study; (D) the density ratio cameleon/Tubulin in each genotype, normalized against that of the ratio in wild-type animals, for each reporter. N=2. Error bars represent SEM.
Figure S4

A

wild-type: AVA

YFP/CFP

0 100 200 300

unc-7: AVA

YFP/CFP

0 100 200 300

B

wild-type: AVE

YFP/CFP

0 100 200 300

unc-7: AVE

YFP/CFP

0 100 200 300
Figure S4. The Pnmr-1::TWK-18(gf) transgene effectively silences the activity of backward premotor interneurons in wild-type animals and unc-7 mutants.

Representative calcium transient traces of premotor interneurons, AVA (A) and AVE (B), in animals without (left panels), or with (right panels) the Pnmr-1::TWK-18(gf) transgene. TWK-18(gf) expression in imaged neurons was monitored and confirmed by the RFP signal co-expressed from the bi-cistronic transcript.
Figure S5

A

**unc-7** wild-type

Duration of movement (s)

0 5 10 15 100

% of Time

20 40 60 80

20

bwd (≥1 pixel)

fwd (≥1 pixel)

pause (<1 pixel)

B

A

B

**unc-7**

wild-type

Duration of movement (s)

% of Time

fwd (≥1 pixel)

bwd (≥1 pixel)

pause (<1 pixel)

Figure S5. Manipulating premotor interneuron activity does not rescue kinking in
innexin mutants. A, the representative body posture during movement, shown by
curvature analyses, of wild-type (upper panels) and unc-7 (lower panels) animals that
expressed different neuronal silencer transgenes. TWK-18(gf) was expressed either by
Pnmr-1 to reduce the activity of the backward circuit premotor interneurons and PVC (+
silence AVA/D/E/PVC), or by both Pnmr-1 and Psra-11 to reduce the activity of all
premotor interneurons (+ silence AVA/D/E/PVC/AVB). kinking, represented by
distorted, discontinuous, deep body bends, could not be rescued by either transgene in
these innexin mutants. B, a quantification of directional movement for animals of the
respective genotypes in A. Upper graphs represent the averaged duration time of forward
and backward movements; the matching lower graphs represent the total percentage of
time that these animals spent in forward or backward motion (movement ≥1 pixel) or
pausing (movement <1 pixel). In wild-type animals, reducing the activity of backward
premotor interneurons and PVC (silencing AVA/D/E/PVC) led to a switch to an almost
exclusive forward movement (black bars). Innexin mutants expressing the same
transgene however failed to execute continuous forward movements (black bar, upper
graph), despite reducing their propensity and duration for backing (grey bars, upper and
lower graph). In unc-7 mutants, reducing the activity of all premotor interneurons
decreased the propensity and duration for backing (grey bars, upper and lower graph), but
it did improve the duration of forward movement (black bar, upper graph). Blocking
chemical synaptic output from backward premotor interneurons (AVA, AVE, AVD) and
PVC (+Pnmr-1::TeTx) also reduced propensity and duration of backing in unc-7 mutants.

*** P<0.001 by the Mann-Whitney U test. Error bars: SEM.
Figure S6

A

hplS179; hplS190

AVA

AVB

AVD

RIM

n.d.

AIA, AIY

B

akls11; hplS179; hplS190

AVA

AVB

n.d.

AIA, AIY

C

akls11; hplS179; hplS190

after laser ablation of the left and right AVB

n.d.

AIA, AIY
Figure S6. The anatomic removal of five pairs of premotor interneurons in wild-type animals. A, premotor interneurons in the head region (arrowheads) were labeled with fluorescent signals in a double transgenic animal hpIs190 (Pnmr-1::D3cpv); hpIs179 (Psra-11::D3cpv); B, in a triple transgenic animal akIs11 (Pnmr-1::ICE); hpIs190; hpIs179, backward premotor interneurons and PVC (located in the tail region, not shown) were absent, but AVBs were intact; C, a young adult animal with the AVB neuron pair laser ablated in the akIs11; hpIs190; hpIs179 background during the second larval stage. n.d. an unidentified pharyngeal neuron, possibly I4. Scale bar: 5μm.
Figure S7. Silencing the A-class motoneurons by TWK-18(gf) failed to reduce the activity of AVA premotor interneurons in wild-type animals. A, Representative traces of the AVA calcium transients in wild-type animals without (left panel) or with (right panel) the same Punc-4::TWK-18(gf) transgene that effectively reduced A-motoneuron activity (Figure 4D) and prevented backward movement; B, Raster plot of the calcium transients for wild-type animals without and with the Punc-4::TWK-18(gf) transgene; C, a mean and distribution of the calcium transient level of wild-type animals without and with the Punc-4::TWK-18(gf) transgene. There was no decrease of the AVA calcium transient level in the presence of the hyperpolarizing transgene Punc-4::TWK-18(gf).
Experimental Procedures:
A list of constructs and transgenes generated for this study

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<th>Transgenes Used For Data in Figures</th>
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### Molecular Biology

All promoters used in this study were generated by PCR-amplification using *C. elegans* N2 genomic DNA as a template. Promoters included a 5.3 kb *Pglr-1* (Maricq et al., 1995), 5.1 kb *Pnmr-1* (Brockie et al., 2001), 2.5 kb *Popt-3* (Fei et al., 2000), 4.8 kb *Prig-3* (Wormbase), 2.8 kb *Psra-11* (Troemel et al., 1995), 3.4 kb *Pacr-2* (Jospin et al., 2009), 2.2 kb *Psto-6* (Po and Zhen, unpublished), 2.5 kb *Punc-4* (Miller et al., 1992) and 4.3 kb *Pacr-5* (Winnier et al., 1999) genomic DNA sequence upstream of the ATG start codon of each gene, respectively. The *Pnmr-1* promoter used in this study excludes a 2 kb fragment encoding *cex-1* that interferes with calcium imaging (Kawano and Zhen, unpublished observation). Intercistronic region consisting of a U-rich element and Splice Leader sequence (UrSL) between *gpd-2* and *gpd-3* was PCR amplified with OZM2301 (AAG CTA GCG AAT TCG CTG TCT CAT CCT ACT TTC ACC) and OZM2302 (AAG GTA CCG ATG CGT TGA AGC AGT TTC CC) using pBALU14 (Tursun et al., 2009) as a template, and inserted into the *Psto-6::wCherry* construct to construct a bi-cistronic expression vector. A series of bi-cistronic expression vectors were generated by

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<th>Constructs</th>
<th>HP Expiration</th>
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substitution of the promoter fragments described above. TWK-18(gf) cDNA was PCR amplified with OZM2376 (AAG GAT CCA AAT GGC GAT TGT TGC) and OZM2377 (TTG GAT CCA CTA GAT GTC ATG CTC TAG) using pCFJ70 as a template and inserted to bi-cistronic vectors. wCherry (a modified version of mCherry codon-optimized for expression in C. elegans) provided by A. Desai (University of California, San Diego), was used as the bi-cistronic expression reporter. unc-7 cDNA was amplified by PCR using OZM412 (GCG GGA TCC GTC AGT CTA TCG TCC CTT GAC CGT) and OZM413 (GCC GGA TCC CCA TGC TCG GCT CCT CCA GCA ATC C) with cDNA library (Vidal cDNA library) as template. unc-9 cDNA was amplified by PCR using OZM417 (GGG GAT CCG GAT GAG TAT GCT ATT GTA TTA TTT CGC G) and OZM418 (GGG GAT CCT CAC ACG TCG TGC ATT TTT CCT TC) with the cDNA library as template. The sequences of resulting DNA clones were confirmed by sequencing. These cDNAs were inserted into bi-cistronic expression vectors described above to generate rescuing constructs for innexin mutants. The YFP region of YC2.12 (EallQ) was substituted with cpVenus to construct pJH1737, a calcium insensitive control for YC3.60. The same E all Q substitutions were introduced into the mtCalmodulin fragment of D3cpv by PCR mutagenesis to generate calcium-insensitive D3cpv control plasmid pJH2157.

Curvature analysis

High contrast images of crawling young adults on OP50 seeded plates were captured using a stereo-microscope or a custom built worm tracker which acquired images at 5 fps (Stephens et al., 2008). Image processing code was written in MATLAB
(Mathworks, Natick, MA). To calculate curvature of the animal, the “spine” was extracted by a modified center-line detection algorithm (Guberman et al., 2008). First the curvature of the animal’s boundary is calculated. The “tail” and the “head” are found by selecting the boundary points with the smallest (tail) and the second smallest (head) radius of curvature. The boundary is then divided into the dorsal and ventral sides. For each point inside the boundary, we find the shortest distances between the point and two sides, and if the difference between the distances is smaller than a certain threshold, the point is considered part of the spine. When all the points on the spine are extracted, the spine is re-sampled to 37 equally spaced points from head to tail. The curvatures of the points are then calculated as $k = (x'y'' - y'x'') / (x'^2 + y'^2)^{3/2}$ where the discrete derivatives are defined as $f'(j) = f(j+1) - f(j-1)$ and $f''(j) = f(j+2) - 2f(j) + f(j-2)$. Images of any animal that crossed over itself were not processed.

**Forward and backward movement analysis**

L4 stage animals were transferred to a NGM plate seeded with a thin layer of OP50. After one minute, a two-minute movie of the crawling animal was recorded using a standard digital camera installed on a Leica MS5 dissecting microscope. Images were sampled from the movies at 1fps and processed by in-house developed ImageJ plugins. The longest distance of the outline of a detected object was calculated, and points giving this value were assigned as the head or tail positions. The anterior to posterior direction of the assigned points was manually determined and tracked automatically by the plugins afterwards to determine the direction of movements. The distance between the center points of the head-to-tail positions between two consecutive frames was calculated to
determine the status of movements. Less than 1 pixel shift in position was defined as "pausing" in this analysis. The percentage of total frames exhibiting forward motion, backward motion and pausing was calculated from the pooled image frames obtained from multiple animals of each genotype. The duration of forward and backward motion, the mean number of video frames with an animal moving in the same direction, was determined for each animal during the two-minute recording. N>10 animals for each genotype were analyzed for the averaged duration.

**Electrophysiology**

Membrane potentials of AVA premotor interneurons were recorded in whole-cell configuration by a Digidata 1440A and a MultiClamp 700A amplifier using the Clampex 10 software, processed with Clampfit 10 (Axon Instruments, Molecular Devices, USA). Data were digitized at 10–20 kHz and filtered at 2.6 kHz. The dissection of *C. elegans* was performed with a modified protocol described previously (Gao and Zhen, 2011). Briefly, one or two day old hermaphrodite adults were glued to a sylgard-coated coverglass covered with bath solution. The cuticle in the head region was cut along the dorsal side of the animal and pinned down to the coverglass to expose neurons of the central nervous system. All recorded animals carried two transgenes expressing two fluorescence markers: a wCherry marker driven by the *Prig-3* promoter (*hpEx2414*) and a cameleon reporter driven by *Pnmr-1* promoter (*hpIs190*). The AVA neurons were identified as neurons expressing both markers.

Neuronal cell bodies were patched using fire-polished 8-12 MΩ resistant borosilicate pipettes (World Precision Instruments, USA). The pipette solution contains
(in mM): K-gluconate 115; KCl 25; CaCl₂ 0.1; MgCl₂ 5; BAPTA 1; HEPES 10; Na₂ATP 5; Na₂GTP 0.5; cAMP 0.5; cGMP 0.5, pH7.2 with KOH, ~320 mOsm. The bath solution consists of (in mM): NaCl 150; KCl 5; CaCl₂ 5; MgCl₂ 1; glucose 10; sucrose 5; HEPES 15, pH7.3 with NaOH, ~330 mOsm. Resting membrane potentials were recorded at 0 pA. Input resistance was determined measuring the voltage responses upon current injections. All experiments were performed at room temperatures (20-22°C). Subsequent analysis and graphing were performed using Excel (Microsoft), Igor Pro (Wavemetrics) and Clampfit (Molecular Devices). In this study, N number represents the number of animals recorded unless specified otherwise. All data are presented as means ± SEM.

**Immunofluorescent staining**

Transgenic animals *unc-9(fc16) unc-7(e5); lin-39(n709); hpEx2227[Pnmr-1::UNC-7::GFP+Punc-4::UNC-9- UrSL-wCherry+Podr-1::GFP]* were fixed in 2% paraformaldehyde for 2 hrs and processed as previously described (Finney and Ruvkun, 1990). *lin-39* eliminates VC neurons, where *Punc-4* also drives expression, to ensure that UNC-9 signals observed strictly came from the A-class motoneurons. A mouse monoclonal antibody against GFP (Roche), and a rabbit polyclonal antibody against UNC-9 (purified from the crude rabbit antisera raised against the last 77 amino acids, Chen et al., 2007) were used at 1:200 and 1:100 dilutions, respectively. Non-transgenic *lin-39(n709); unc-9(fc16) unc-7(e5)* progeny from the transgene population, serving as negative controls, showed no background staining.
Silencing, inactivation and ablation of premotor interneurons

Silencing of premotor interneurons was achieved by expressing Pnmr-1::TWK-18(gf) to inactivate AVA, AVE, AVD and PVC, or/and Psra-11::TWK-18(gf) to inactivate AVB. Both constructs co-express wCherry from a bi-cistronic transcript to allow the confirmation of the transgene expression.

Ablation of premotor interneurons was achieved through a combined genetic and laser ablation approach. The pair of AVB neurons was removed at the second larval (L2) stage with a MicroPoint laser ablation system (Photonic Instruments) as previously described (Bargmann and Avery 1995). The AVB laser ablations were performed in the background of akIs11; hpIs179; hpIs190, a strain that expresses an integrated Pnmr-1::ICE (human caspase, akIs11) to induce apoptosis in AVA, AVD, AVE and PVC (Zheng et al., 1999), Psra-11::D3cpv to label AVBs, and Pnmr-1::D3cpv to monitor the absence of the rest of premotor interneurons. The removal of all premotor interneurons was confirmed by the absence of hpIs179 and hpIs190 fluorescence signals in all premotor interneurons 24 hours post laser ablation. Curvature analysis was performed on ablated animals at the last larval (L4) stage. A total number of 6 such animals were obtained and examined for this experiment.

Blocking the chemical synaptic output of AVA, AVD, AVE and PVC was achieved by expressing Pnmr-1::Tetanus toxin in unc-7(e5) animals. This construct co-expresses wCherry from a bi-cistronic transcript to allow the confirmation of the transgene expression.
Statistical Analysis

For calcium imaging and motion analyses, p values were calculated by the two-tailed Mann-Whitney U test. For electrophysiology analyses, two-tailed Student’s t-test was applied to determine statistical differences. Statistical differences at p < 0.05 are considered to be significant (* P<0.05, ** P<0.01, *** p<0.001).

References


Hille B. (2001) Ion Channels of Excitable Membranes (Sinauer, Sunderland, MA), 3rd Ed


