A GABAergic/peptidergic sleep neuron also functions as a locomotion stop neuron with compartmentalized Ca\(^{2+}\) dynamics

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Supplementary Information
Supplementary figures

Supplementary Figure 1: Stop behavior and pharyngeal inhibition, elicited by RIS photostimulation. A) Locomotion stop delay is variable in WT animals. A fast (blue) and a slow (red) responder are highlighted. All other traces are shown in grey, the mean (thick line) ± SEM (thin lines) of all animals is shown in black. Illumination period is indicated by a blue bar. Zoomed in from data in Fig. 1B. B) Pharyngeal pumping inhibition upon RIS::ChR2 depolarization depends on illumination intensity. Pharynx pumping was visually inspected during RIS photostimulation, at different light intensities. The fraction of animals stopping pumping and locomotion at the indicated light intensities was registered for the indicated number of animals in three independent experiments. C) Electropharyngeograms obtained from cut-head preparation of RIS::ChR2 expressing animals, raised either with or without ATR. Blue bar indicates illumination (3 mW/mm², 470 nm). Spontaneous pumping was achieved by 5 min incubation with 2 μM serotonin prior to data acquisition. D) Normalized pump frequency of animals expressing RIS::ChR2 raised with ATR. Pump frequency aggregated per animal and 30 s bin as shown in (C). Box plot with Tukey whiskers. ***p≤0.001: Student's T-test with Bonferroni correction, n = 10.
Supplementary Figure 2: RIS photostimulation did not inhibit action potentials in BWMs: A) Overlay of the seven single current clamp measurements recorded from BWM at the ventral side, anterior to the vulva, as shown in B). RIS::ChR2 photostimulation denoted by blue bar and/or shading.
**Supplementary Figure 3: Neuron positions in the head and connectome of the RIS neuron.**

A) Representation of the cholinergic cell bodies with the ROI scored in Fig. 3E, depicted as green shaded area. Pink cell bodies are cholinergic neurons outside of the imaged region, while their processes might influence the recorded signal as well. Based on Gendrel, M., Atlas, E.G. & Hobert, O. A cellular and regulatory map of the GABAergic nervous system of C. elegans. *eLife* 5, e17686 (2016).


C) As in (B) but omitting gap junctions mediated by UNC-9.
Supplementary Figure 4. Initial steps in image processing for RIS axon Ca$^{2+}$ imaging: B) Image analysis involved cropping (I), binarization (II), repositioning the soma (III), and determining the angular orientation of the axon (IV). D, dorsal; V, ventral; P, posterior; A, anterior. Scale bar: 20 μm.

Supplementary Figure 5. RIS Ca$^{2+}$ signals are aligned to the moment of reversal (zero velocity), compared in wild type and flp-11(tm2706) mutants: A) Reversal-aligned normalized GCaMP fluorescence intensity (green) and animal velocity in μm/s (blue). A significant increase in Ca$^{2+}$ (n=75 reversals; p <0.001) was observed at the time before and after the moment of reversal, calculated for time periods indicated by brackets, flanking the point of zero velocity during reversal (blue lines). B) as in A), but in flp-11(tm2706) mutants. Significant increase in calcium (n=100 reversal events). Box plots with Tukey whiskers, ***,p≤0.001.
Supplementary Figure 6. RIS Ca\textsuperscript{2+} signals, or their time derivative, aligned to the peak, and concomitant animal speed or acceleration, compared in wild type and flp-11(tm2706) mutants: A, C) Ca\textsuperscript{2+} peak aligned normalized fluorescence (A), or (C) Ca\textsuperscript{2+} change rate (dF/dt) (green), and the animal speed (A) or acceleration (C) in µm/s\textsuperscript{2} (blue). n=45 Ca\textsuperscript{2+} peaks were aligned. Significant reductions in speed or acceleration were observed at the times before and during the onset of Ca\textsuperscript{2+} rise (indicated by green vertical lines). Differences were calculated for time periods indicated by brackets. Box plots with Tukey whiskers, ***p≤0.001; *p≤0.05. B, D) as in A, C), but for flp-11(tm2706) mutants (n=25 Ca\textsuperscript{2+} peaks).
Supplementary Movies

Supplementary Movie 1. **RIS photostimulation leads to a block of locomotion behavior:** Illumination of the anterior portion with a tracking and illumination system (Stirman et al., 2011) leads to a stop of the locomotion. Left half shows video stream, on the right, the illumination pattern sent to the projector is shown.

Supplementary Movie 2. **Photostimulation of GABA neurons leads to a transient inhibition of locomotion:** Stimulation of ChR2, expressed in GABA neurons (punc-47 promoter), causes a transient stop followed by resumed, yet uncoordinated locomotion.

Supplementary Movie 3. **Ca^{2+} fluctuations in the BWMS are suppressed by RIS photostimulation:** A) Animal expressing RCaMP1h in BWMS and ChR2 in RIS is immobilized and imaged for RCaMP fluorescence, shown in ‘fire’ lookup table. Video is accelerated 5 times. B) Difference video corresponding to A, showing fluorescence increase and decrease in red and blue, respectively.

Supplementary Movie 4. **Ca^{2+} oscillations in cholinergic MNs are suppressed by RIS photostimulation:** Animal expressing RCaMP1h in cholinergic MNs and ChR2 in RIS is immobilized and imaged for RCaMP fluorescence in the head ganglia. Difference video, showing fluorescence increase and decrease in red and blue, respectively. Video is accelerated 20 times.

Supplementary Movie 5. **Depiction of workflow for video processing to deduce localized Ca^{2+} dynamics in the RIS axon:** Upper row, from left to right: cropping, binarization, repositioning the soma, determining the angular orientation of the axon. Lower row: reorienting the raw image, masking unspecific gut fluorescence, fitting a parabola, measuring fluorescence intensity in perpendicular rectangular ROIs.

Supplementary Movie 6. **Example video of RIS activity during free locomotion, including slowing and reversal events:** Left half: RFP fluorescence (pharynx marker for tracking); right half: GCaMP6s image. Upper left corner: Reoriented and cropped image of RIS in the GCaMP6s channel.

Supplementary Movie 7. **RIS Ca^{2+} activity in the nerve ring axon and in the branch during locomotion.** Left: original video of the head region of free moving *C. elegans* expressing GCaMP6s in RIS. Upper right: Extracted RIS neuron fluorescence, with indicated (right, below) morphology (CB: cell body, Br: branch, NR: nerve ring portion of the axon, with overlaid false-colored Ca^{2+} fluorescence \( \Delta F/F_0 \) signal (blue – white hues: low – intermediate signal, yellow – red hues: intermediate – high signal. Scale and arrow on the right indicate locomotion speed and direction (green: forward, red: reverse).
Supplementary Movie 8. Example video of RIS branch activity during free locomotion. A) RIS GCaMP6s fluorescence signal false colored coded during a reversal. B) Analogous to A), but during a stop without subsequent reversal.

Supplementary Information files:

Excel sheets with the original / processed data for each of the figure panels.