A GABAergic and peptidergic sleep neuron as a locomotion stop neuron with compartmentalized Ca2+ dynamics

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Animals must slow or halt locomotion to integrate sensory inputs or to change direction. In Caenorhabditis elegans, the GABAergic and peptidergic neuron RIS mediates developmentally timed quiescence. Here, we show RIS functions additionally as a locomotion stop neuron. RIS optogenetic stimulation caused acute and persistent inhibition of locomotion and pharyngeal pumping, phenotypes requiring FLP-11 neuropeptides and GABA. RIS photoactivation allows the animal to maintain its body posture by sustaining muscle tone, yet inactivating motor neuron oscillatory activity. During locomotion, RIS axonal Ca2+ signals revealed functional compartmentalization: Activity in the nerve ring process correlated with locomotion stop, while activity in a branch correlated with induced reversals. GABA was required to induce, and FLP-11 neuropeptides were required to sustain locomotion stop. RIS attenuates neuronal activity and inhibits movement, possibly enabling sensory integration and decision making, and exemplifies dual use of one cell across development in a compact nervous system.

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Animals actively stop locomotion in order to await certain events or to avoid potentially dangerous situations. In order to quickly resume locomotion after the stop, they must keep their muscle tone. This is in contrast to phases of behavioral quiescence, or sleep, where vertebrates typically lose their muscle tone and assume a relaxed body posture\cite{1,2}. In limbbed animals, multi-layered neuronal systems control locomotion\cite{3}. Central pattern generators (CPGs) in the spinal cord mediate (1) rhythm generation, usually by networks of excitatory neurons that oscillate and cause mutual inhibition via interneurons, and (2) pattern generation that regulates motor neurons (MNs), and thus muscle action, to orchestrate coordinated movements underlying locomotion. For left-right coordination during walking, inhibitory and excitatory commissural interneurons are required\cite{4,5}. Locomotion is triggered by excitatory signals descending from supraspinal regions of the mid-brain or hindbrain (in mammals\cite{5}) or of the brain stem (in tadpoles\cite{6}), which “call” the spinal CPG networks into action. Recently, a class of interneurons in the murine brainstem was shown to induce a stop command for the pattern generation systems\cite{7}. These V2a “stop” neurons project to excitatory and inhibitory spinal cord neurons, inducing locomotion halt likely via inhibition of rhythm-generating neurons. The stop neurons do not reduce muscle tone and do not inhibit MNs. Thus the animal does not collapse but rather can quickly resume locomotion. Equivalents of these “stop neurons” and systems for slowing were identified in non-limbed vertebrates\cite{8,9,10}, and recently also in *Drosophila*\cite{11}, where descending interneurons induce locomotion stop during navigation of odorant gradients, while activity of other neurons causes slowing\cite{12}. However, molecular identities of stop cells are only partly known, and also different organisms appear to use different mechanisms and partly redundant circuitry to induce locomotion stop\cite{13}. Thus it is unclear whether “stop” systems evolved several times or whether a primordial locomotion stop system diversified into the different systems present today in different organisms.

In the nematode *Caenorhabditis elegans* with its much smaller number of neurons, rhythm generation resides in the MNs. Ventral cord excitatory MNs coordinate the undulatory behavior for forward and backward locomotion (B- and A-class, respectively). They exhibit oscillatory activity patterns that are entrained by proprioceptive feedback as well as bi-directional coupling by premotor interneurons (PINs) in the ventral nerve cord\cite{14,15,16}. In addition, the AS-class of asymmetric MNs exhibits oscillatory activity, interacts with PINs, and contributes to propagation of the body wave\cite{18}.

Activity of vertebrate stop neurons contrasts descending pathways that are active during sleep, which halt locomotion and affect muscle relaxation through inhibitory reticulospinal neurons\cite{19}. Sleep also occurs in *C. elegans* (1) Lethargus, also called developmentally timed sleep (DTS), occurring during larval molt transitions\cite{20,21,22} and (2) stress-induced sleep (SIS), in response to cellular insults\cite{21,22,23,24,25}. Both states are similar with a lack of locomotion and feeding, as well as increased arousal threshold. They are regulated by distinct neuropeptidergic pathways and neurons, i.e., NLP-8, FLP-24, FLP-13 (neuropeptide-like protein, FMRFamide-like peptide) neuropeptides and the ALA (anterior lateral A) neuron for SIS\cite{20,23,26,27}, while the GABAergic and neuropeptidergic RIS (ring interneuron S) neuron, as well as NLP-22 neuropeptides, are required for DTS\cite{24,25}. RIS photoactivation induced lethargus in larvae, independent of GABA, while FLP-11 neuropeptides were required for RIS-induced sleep, and their overexpression sufficed to cause lethargus\cite{26}. In *Ascaris*, FLP-11 was shown to be inhibitory to muscle cells\cite{27}. Also in other species, sleep and arousal are encoded by neuropeptidergic and biogenic amine neurotransmitters\cite{28}. In mammals, sleep is mainly regulated in the ventrolateral preoptic nucleus, where GABAergic and galanin-releasing neurons inhibit orexin/hypocretin-releasing neurons during sleep, while these modulators are released during (and promote) wakefulness\cite{29}. Mutations in the orexin receptor or abnormal activity in the hypothalamic area may lead to narcolepsy or cataplexy\cite{30}. Orexin homologs were found in insects (allatotropin) but not in *C. elegans*, where instead FLP-21 and NLP-49 neuropeptides as well as pigment-dispersing factor mediate arousal\cite{31,32,33}. Stop neurons were not identified in *C. elegans* to date.

Adult *C. elegans* show behavioral quiescence during satiety, starvation, or recovery from stress, e.g., heat shock\cite{22,23,24}. These states are associated with the RIS and ALA neurons, but it is unclear whether RIS has functions other than as a pure sleep neuron. In the compact *C. elegans* nervous system, neurons often multitask, thus RIS might be utilized for additional functions. *C. elegans* frequently interrupts its predominantly forward locomotion by brief reversals, then resumes forward locomotion with a change in direction. It is only partially understood which neurons orchestrate this pirouette behavior and in which sequence they may act\cite{35,36}. Forward locomotion slows down before the animal briefly stops, and this part of the pirouette could be actively controlled by neuronal activity\cite{37}. Thus, may the RIS neuron function like vertebrate stop neurons, i.e., inducing a brief locomotion stop while maintaining muscle tone, to enable directional changes? Furthermore, may functions of sleep and stop neurons have been combined in one cell in the compact worm nervous system, and could this thus represent an evolutionary ancient mechanism from which sleep and stop systems diversified into distinct systems?

We address these questions by (opto-)genetics and by imaging activity of RIS during locomotion. Genetic ablation of RIS reduces reversal and stop events. Specific RIS photoactivation induces a full locomotion stop and also affects other rhythmic behaviors, like pharyngeal pumping. RIS stimulation is accompanied by sustained Ca$^{2+}$ levels in body wall muscle (BWM) cells and inhibition of oscillatory activity in MNs. RIS-dependent behavioral responses are largely blocked without neuropeptidergic transmission, while interfering with GABA transmission affects their kinetics, and eliminating gap junctions uncovers further functions of RIS within circuits coordinating forward/backward transitions. The major determinant of RIS effects in adults, just as in larval sleep, is the FLP-11 neuropeptide. RIS shows compartmentalized axonal Ca$^{2+}$ transients. In the nerve ring process, the onset of these signals correlates with the onset of locomotion slowing, while in an axonal branch, they are correlated with the induction of reversals and require FLP-11 signaling. Our work dissects the function of a *C. elegans* “stop” neuron, providing new insights into the roles and circuits of such neurons. It may help to understand such neurons, identified only phenotypically\cite{12}, and emphasizes that stop cells may exist widely across locomotion systems.

**Results**

Single-cell-specific expression and photoactivation of channelrhodopsin-2 (ChR2) in RIS induces locomotion stop. We achieved conditional expression of ChR2 and green fluorescent protein (GFP) in the single RIS neuron. Animals showed fluorescence in a single cell body located in the ventral ganglion, next to the pharyngeal posterior bulb, on the right side of the head. RIS has a single process extending anteriorly toward the nerve ring, with a short branch reaching into the ventral nerve cord, while the axonal process wraps around the isthmus of the pharynx (Fig. 1a). When animals expressing RIS::ChR2 were cultivated in presence of all-trans retinal (ATR) and illuminated with blue light, all locomotion behavior stopped: On average, velocity dropped not only by 80% within 4–5 s ($\tau = 1.67$ s; Fig. 1b;
Supplementary Movie 1) but also quite immediate for individual animals, i.e., 1–2 s (Supplementary Fig. 1A). We analyzed the body posture of the animals, i.e., bending angles demarcated by three adjacent points along the body axis (Fig. 1c)39. Forward locomotion (sinusoidal body wave propagating antero-posteriorly) stopped upon RIS photostimulation: During the 10 s light stimulation, bending angles were “frozen”, i.e., the body posture was maintained. This contrasts the stop in locomotion upon photostimulation of all GABAergic MNs40, where animals resumed behavior after a few seconds of photostimulation, though being uncoordinated (Supplementary Movie 2). GABA neuron photostimulation causes overall body elongation by 4%40. Also RIS photostimulation induced body elongation, however, only ca. 2.5%, affecting only the anterior third, which elongated ca. 10% (Fig. 1d). Thus RIS may inhibit neurons driving the undulations, with some anterior muscle relaxation. RIS photoactivation also attenuated responses to mechanical stimulation, where a harsh touch to the head region led mostly to short reversals smaller than one body length; conversely, after RIS photostimulation, almost all animals reacted with a long reversal (Fig. 1e). During DTS, mechanical stimulation was shown to activate RIS, likely to suppress wake behaviors41, thus RIS photostimulation might mimic this larval behavior inhibition. To ask whether RIS is sufficient or required for reversals and stops, we ablated it using cell-specific overexpression of the apoptosis inducer EGL-1 (“egg-laying defective”), with GFP as a marker22. Animals lacking RIS showed significantly fewer long reversals and stops than wild type (WT; Fig. 1f), demonstrating a requirement of RIS for these behaviors.

Last, rhythmic pumping of the pharynx, the muscular feeding organ, ceased during 30 s RIS photostimulation depending on light intensity (Supplementary Fig. 1B). In electrophysyngograms (EPGs), i.e., extracellular recordings of electrical activity associated with pharyngeal contractions42, RIS photostimulation generally evoked a complete absence of electrical transients and on average significantly reduced the number of pump events (Supplementary Fig. 1C, D). Thus RIS activation, presumably through GABA and neuromodulator release, may inhibit pharyngeal pumping during locomotion reversals, in parallel to another established pathway for pumping inhibition using serotonin, the SER-2 receptor, and Gαo signaling37.

RIS suppresses oscillatory activation of BWM by affecting cholinergic neurons. To explore the effects of RIS on the locomotion system, we used Ca2+ imaging. RcaMP1h, a red-fluorescent genetically encoded Ca2+ indicator43, was specifically expressed in BWMs in addition to ChR2 in RIS, and animals were physically immobilized44 (Fig. 2a, Supplementary Movie 3). Ca2+ signals were monitored over time, either in dorso-ventrally opposite regions (Fig. 2a, b) or in line scans along the animals’ dorsal muscles, assessed as “kymograms” spanning the body length (Fig. 2a, c). Despite the absence of dynamic proprioceptive feedback, muscular Ca2+ levels visibly fluctuated along the body. Ca2+ levels were high in bent regions, in line with muscular activity underlying the body bend, and with the proprioceptive coupling of MNs in one body segment to the anterior segment45. The Ca2+ signals oscillated at low frequency (ca. 0.12 Hz) in a dorso-ventrally reciprocal fashion (Fig. 2b). They could thus reflect rhythmic activity of the cholinergic MNs innervating the respective muscle cells. Oscillations were much slower than in free-moving animals (~0.36 Hz46; note that, due to immobilization, no traveling of the Ca2+ wave is observed). During RIS ChR2 activation, this oscillatory activity essentially stopped, while BWM Ca2+ levels did not obviously change (Fig. 2c; for Δ[Ca2+] over time, see Supplementary Movie 3B). Since oscillations were asynchronous across animals and to enable comparisons independent of actual signal intensities, we used sample auto-correlation, where the autocorrelation period reflects the extent of oscillatory activity (Fig. 2d). For animals grown without ATR, the period duration did not differ between the dark and lit periods, while it significantly increased with ATR present (Fig. 2e), indicating robust slowing of oscillations (since during 20 s stimulation often no full oscillations occurred, we assumed a 20 s minimum of the period).
Fig. 2 RIS photoactivation stopped muscular Ca$^{2+}$-dynamics. **a** Maximum intensity projection of RCAmp imaging in BWM cells of an immobilized animal (arrow: pharynx, pmyo-2::mCherry marker). Boxed regions: Violet, blue: regions of interest for dorso-ventrally alternating activity; white: region of interest for kymographic analysis of dorsal muscle Ca$^{2+}$ signals along the body in c, scale bar = 100 µm, A, P, V, D: anterior, posterior, ventral, dorsal, respectively. **b** Ca$^{2+}$ dynamics in both regions of interest from a (V = ventral, D = dorsal muscle cells) measured before, during, and after RIS::ChR2 photostimulation, denoted by the blue bar. F$_0$ defined as the mean RCAmp intensity during the first 4.5 s of the recording. **c** Kymograph representation of the Ca$^{2+}$ dynamics along the dorsal side. The RCAmp signal was normalized for visualization purposes. RIS photoactivation: blue bar. Scale bar = 10 s. **d** Example of the RCAmp signal autocorrelation for a specific point in the BWM over time, before, during, and following illumination. Ventral and dorsal BWMs were analyzed. Red lines: Peak autocorrelation of two consecutive Ca$^{2+}$ waves and their time lag (if no consecutive Ca$^{2+}$ signals were detected during the stimulation period, stimulus duration was taken as lower bound). **e** Distribution of the mean change in the muscular Ca$^{2+}$ oscillation period per animal, compared to before RIS photoactivation. When no oscillations occurred, the duration of photostimulation (20 s) was assumed as minimal period. Compared are animals without and with ATR, number of animals indicated in gray numbers. ***p ≤ 0.001; **p ≤ 0.01; statistical significance tested by ANOVA, Bartlett’s test, and Bonferroni’s multiple comparison test

Next, we assessed these effects in body muscle, i.e., downstream of MNs, by electrophysiology. Miniature postsynaptic currents (mPSCs) report on release of neurotransmitter from single synaptic vesicles (Fig. 3a). If RIS photodepolarization reduced muscular activity by inhibiting MNs, we would expect reduced mPSC rates. However, RIS::ChR2 photostimulation neither abolished nor reduced mPSCs and their frequency or amplitude (Fig. 3a–c). This indicates that MNs did not reduce their activity during RIS signaling and released the same amount of transmitter. Possibly they altered their relative activities, e.g., if the frequency of oscillation of the CPGs they constitute is altered, and thus MN activity may have become asynchronous. We thus analyzed the frequencies of the observed mPSCs (Fig. 3d). Predominantly low frequencies were populated in this analysis, and RIS photoactivation caused a significant reduction of their power. We also assessed muscular action potentials, which occurred at low frequency, not obviously silenced by RIS photostimulation (Supplementary Fig. 2). Thus, muscle tone is maintained during RIS activity, while MN activity is desynchronized.
As the observed muscle activity is evoked by MNs even in restrained animals, we wanted to analyze MNs directly. If RIS effects on BWM Ca\(^{2+}\) fluctuations occur at the MN level, Ca\(^{2+}\) fluctuations in MNs may seize during RIS photoactivation. We expressed RCaMP in a large subset (132/160) of cholinergic neurons, encompassing all synaptic cholinergic partners of RIS (VB, DB, AS, RMD, SMD, SDQ, PVC, AVE, including cells in the head ganglia, but excluding SAB; Supplementary Fig. 3A; see “Methods”), and analyzed Ca\(^{2+}\) dynamics before and during RIS photoactivation (Fig. 3e, f, Supplementary Movie 4). We observed spontaneous fluctuations in the signals across head MNs of ~0.17 Hz. RIS photoactivation increased the mean relative autocorrelation period 1.6-fold (Fig. 3g), i.e., activity in the head cholinergic spontaneous fluctuations in the neuronal network, which is the likely reason for the observed reduction of muscle oscillations.

**RIS photostimulation effects are accelerated by GABA transmission.** Effects of RIS on various behaviors could depend on different types of neurotransmission: RIS is GABAergic, peptidergic, and makes gap junctions, which may all be driven by ChR2 stimulation. In RIS’ control of DTS, GABA played no role, as lethargus still occurred in GABA-defective mutants, and was instead instructed by peptidergic transmission. We tested mutants of these signaling pathways by analyzing the locomotion state as forward, reverse, or stop categories, defined by a threshold of ±45 μm/s on the velocity trajectory. WT animals stopped locomotion during RIS photoactivation (Fig. 4a). After the stimulus ended, a significant proportion the animals resumed locomotion within 2 s, however, inducing reversals, from which they gradually returned to mostly forward locomotion. Mutants lacking the vesicular GABA transporter (“uncoordinated” *unc-47* (e307) animals) still exhibited the RIS-induced stop response, with delayed onset, and the post-stimulation reversal. RIS induced body elongation that was significantly stronger in *unc-47* mutants than in WT (Fig. 4b). This appears paradoxical, yet, compared to WT, *unc-47* animals are pre-contracted due to absence of GABA at the neuromuscular junction, thus inhibition of cholinergic dynamics by RIS (Fig. 3) likely causes more pronounced body elongation. In sum, GABA is not involved in maintaining RIS-induced locomotion stop phenotypes or the post-stimulation reversal but rather speeds up the slowing and

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**Fig. 4** The stop phenotype induced by RIS photoactivation requires GABA and neuropeptide signaling. a Animal locomotion analyzed before, during, and after photoactivation of RIS (in *lith-1(e374)* background) to eliminate unspecific photophobic responses and the proportion of animals in distinct state (forward (green), stop (white), reversal (magenta)) deduced from the animal velocities is represented in color, over time, across all animals analyzed (number of animals and genotypes indicated above each data set). Significant change in stop proportion during RIS photoactivation versus WT indicated; blue bar and blue shade: illumination period; scale bar: 2 s. b Relative body elongation during RIS photoactivation: box plot with Tukey whiskers, numbers of animals, and genotypes are indicated below. c, d Mean ± SEM locomotion speed (c) or body length (d) before, during, or after photoactivation of RIS-CHR2 (blue bars) compared in egl-3(gk238) mutants raised with or without ATR. Number of animals depicted in gray. e Mean normalized angular velocity in anterior quarter of the animal for WT and *unc-9(e101)* mutants expressing RIS-CHR2, with and without ATR. Box plot with Tukey whiskers. ***p ≤ 0.001; **p ≤ 0.01; *p ≤ 0.05; ns: non-significant; statistical significance tested by ANOVA, Kruskal-Wallis with Dunn’s Multiple Comparison Test in (a, black, versus WT in b, e) or Wilcoxon Signed Rank Test, versus no body length change (red, in b)
RIS photostimulation induces subsequent reversals via gap junctions. RIS forms electrical synapses with five neuron types (Supplementary Fig. 3B): AIB, AVJ, DB, RIM, and SMD. We analyzed whether effects observed during and after RIS::ChR2 activation may be due to concomitant depolarization of these neurons in mutants lacking the innexin (gap junction subunit) UNC-9, expressed in RIS and forming homotypic or heterotypic gap junctions with itself or with UNC-7. Based on its expression pattern, lack of UNC-9 should affect gap junctions between RIS and AIB, AVJ, DB, SMD, and RIM (Supplementary Fig. 3C). RIM inhibits reversals. Conversely, reversals can be induced by AIB neurons, which inhibit RIM, via disinhibition mutants on average showed no relaxation (Fig. 4b). RIS was found to affect ChR2 signaling by analyzing mutants lacking the RIS photostimulation effects require neuropeptides. We next assessed the role of neuropeptides in photovoked RIS:: ChR2 signaling by analyzing mutants lacking the Ca\textsuperscript{2+}-dependent activator protein for secretion (CAPS, encoded by unc-31) or the pro-protein convertase EGL-3. UNC-31 is required for secretion of (many of the) mature neuropeptides, while EGL-3 mediates processing of most if not all neuropeptide precursors. RIS photostimulation in unc-31(e1304) mutants still evoked stopping effects induced by RIS activation. This may be via RIB neurons, which were shown to increase locomotion speed, and to which RIS makes synapses\textsuperscript{58,49} (Supplementary Fig. 3B).

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Ca\textsuperscript{2+} activity in the RIS axon and soma in freely moving animals. Photostimulated RIS affects locomotion, pharyngeal pumping, and withdrawal after mechanical stimuli via GABA, FLP-11, and possibly other neuropeptides acting on different cells. During intrinsic behavior, RIS is active along with other neurons. Singular stimulation of RIS, evoking the observed phenotypes, could thus artificially exaggerate only aspects of behaviors that occur when RIS is (co-)activated with/by other cells. We explored this by analyzing RIS activity in free-moving animals by Ca\textsuperscript{2+} imaging. We modified a previously described tracking system. A four-quadrant photomultiplier controls a stage, keeping a fluorescent spot in the center, and a high-magnification fluorescence video is acquired. An infrared behavior video is acquired at low magnification, stage positions are recorded, and the combined data streams allow behavioral quantification and correlation with neuron activity. We expressed GCaMP6s in RIS, along with a red fluorescent protein in the pharyngeal terminal bulb (TB), to track the head region (Fig. 5a). As not RIS but the TB was tracked, the RIS image rotates around the center of the picture, while the animal changes direction. Custom-written software for image processing (1) registered images on the RIS cell body, (2) cropped a region of interest (ROI) containing the entire RIS morphology and rotated it such that the axon was oriented (Fig. 5b, Supplementary Fig. 5 and Movies 5 and 7), (3) defined a smaller ROI for each image, depicting only the RIS soma and axon, (4) fitted, along the spine of this ROI, a parabola, defined 100 equally spaced perpendicular segments, and quantified their fluorescence.

RIS axonal Ca\textsuperscript{2+} activity during locomotion is correlated with slowing and the onset of reversals. RIS Ca\textsuperscript{2+} activity, assessed on short time scales (in contrast to previous analyses during sleep), was transient, or lasted for several seconds, and coincided with slowing and/or reversals (Fig. 5c), for which the magnitude of the Ca\textsuperscript{2+} signal was indicative. Ca\textsuperscript{2+} rose stepwise while the animal slowed, until high levels were reached (Fig. 5d; Supplementary Movie 6) and the animals exhibited subsequent reversals. During those reversals, Ca\textsuperscript{2+} dropped and forward movement resumed a few seconds later. Across all events recorded, the main change in Ca\textsuperscript{2+} signal was observed in the nerve ring region of the RIS axon (Fig. 5e); sometimes activity was also observed in the branch (Fig. 5c, d; Supplementary Movies 7 and 8). Axonal Ca\textsuperscript{2+} transients are expected to be most indicative of relevant neuronal activity, likely correlated with transmitter release. Analyzing axonal events further helped to exclude noise from intestinal fluorescence. Thus we used only the fluorescence of the anterior
half of the neuronal ROI, comprising the axon around the nerve ring, unless we also analyzed the axonal branch. Slowing and reversal events (as shown in Fig. 5c) were identified by analyzing locomotion based on the x, y position of the tracked animal and its body posture, allowing to derive directional velocity along the mid-body axis. The relative occurrence of reversals and Ca\(^{2+}\) peaks are shown in Fig. 6a, as probability distributions, aligned to the nearest Ca\(^{2+}\) peak or nearest reversal, respectively. The skew of these distributions suggested that reversals followed the onset of a Ca\(^{2+}\) rise. We aligned events recorded from 20 animals, either to Ca\(^{2+}\) peaks (Fig. 6b, d, 45 events) or to the moment of reversal (Supplementary Fig. 5A, 75 reversals). For Ca\(^{2+}\) peak-aligned data, we either analyzed velocity (becoming negative upon reversal), to probe if RIS Ca\(^{2+}\) may be a determinant of reversals (Fig. 6b), or absolute speed, to explore if RIS may be a speed sensor (Supplementary Fig. 6A).

Fig. 5 Ca\(^{2+}\) activity measured along the RIS axon in freely moving animals correlates with slowing and reversals: a Strain used for tracking and Ca\(^{2+}\) imaging RIS in moving animals, expressing a red fluorescent marker in the pharyngeal terminal bulb (for tracking) and GCaMP6s in RIS. Micrographs of red (II) and green (III) fluorescence and merged color channels (I). Scale bar: 50 µm. b Image analysis after binarization and repositioning the soma involved reorienting the raw image (I), masking unspecific gut fluorescence (II), fitting a parabola (III), and measuring fluorescence intensity in perpendicular rectangular ROIs (IV). Dorsal is up, anterior to the left; CB cell body, NR nerve ring. c, d Upper panels: Representative traces of animal velocity (blue) and fluorescence intensity in the RIS nerve ring portion (green). Lower panels: Corresponding heat maps displaying the normalized fluorescence dynamics along the axon over time. RIS pictograms on the left indicate morphology including nerve ring (NR), branch (Br), and cell body (CB), the distance along the axon as well as the region of the nerve ring (green box) used for calculating the \(\Delta F/F_0\) traces in the upper panels, while dashed region shows extent of ROIs analyzed in lower panels (also in e). Distinct Ca\(^{2+}\) rise events in the branch region are boxed. e Mean normalized fluorescence heat map of \(n = 45\) acquired Ca\(^{2+}\) events along the entire length of RIS, partially excluding the soma, by aligning time windows 6 s prior and post Ca\(^{2+}\) peaks (\(N = 11\) animals)
Fig. 6 RIS Ca\(^{2+}\) activity induces decreased forward locomotion and increased reversal probability, which requires FLP-11 neuropeptides: a Conditional probability density function of the shortest unbiased time lag to a reversal given a Ca\(^{2+}\) peak event (aligned nearest reversal events depicted as blue lines) and vice versa for the probability of a Ca\(^{2+}\) peak event given a reversal (green lines: time lag to the nearest reversal) in WT animals. The dotted vertical line indicates the mean onset of a Ca\(^{2+}\) rise, the black solid line indicates peak Ca\(^{2+}\). b Ca\(^{2+}\) peak-aligned normalized GCaMP intensity (green, mean ± SEM) in the nerve ring region of the RIS axon (as depicted in Fig. 4c) and animal velocity in μm/s (blue, mean ± SEM, n = 45; a significant reduction in the two periods before and during the Ca\(^{2+}\) rise is shown on the right, boxplot, p < 0.001). Shown in red is the mean first derivative (dF/dt rise rate, s\(^{-1}\)) of all Ca\(^{2+}\) signals. c Mean ± confidence intervals of time-shifted cross-correlation (red) of animal velocity aligned to Ca\(^{2+}\) signal rise rate, (as in b). Anticorrelation is significantly different from 0 and shows a negative time lag (t = −0.15). d Ca\(^{2+}\) peak-aligned analysis of the proportion of the animal population in one of the four behavioral states: 1 moving forward and accelerating (v+a), positive velocity and acceleration, dark green; for acceleration data, see Supplementary Fig. 6C, D), 2 moving forward and decelerating (v–a, light green), 3 moving backwards, but accelerating (v–a+, pink), and 4 moving backwards and slowing (v–a–, dark red). Number of reversing animals increased significantly during RIS Ca\(^{2+}\) events (T test, p < 0.001). Green line indicates mean onset of RIS Ca\(^{2+}\) rise. i Scatter plot with means of the data in d, h; statistical differences analyzed for the time periods indicated by black boxes (before and during Ca\(^{2+}\) rise). e–h As in a–d, but in flp-11(tm2706) background. Boxplots in b, f compare the average in the time windows indicated by black brackets below the traces. ***p ≤ 0.001; *p ≤ 0.05; ns: not significant; statistical significance tested by paired T test in b, f and unpaired T test in c, g. ANOVA with Tukey multiple comparisons test in i.
RIS Ca\(^{2+}\) signals required \(-1.2\) s from detectable onset to peak (Fig. 6b). Concomitantly, a drop in velocity occurred, which in 80% of cases led to a reversal (mean velocity thus approached zero but remained positive). Animals significantly decreased velocity (Fig. 6b, box plot). We also assessed the onset of Ca\(^{2+}\) rise (determined from dF/dt), which likely coincides with transmitter release (Fig. 6b). Cross-correlation of the Ca\(^{2+}\) rise rate and velocity drop showed a low but significant coefficient of \(-0.15\) and a time lag of \(-0.8\) s (Fig. 6c). Thus the onset of the Ca\(^{2+}\) rise (green vertical line, Fig. 6b) preceded slowing, which is likely regulated by RIS. Correlating the moment of reversal and the Ca\(^{2+}\) signal indicated that RIS Ca\(^{2+}\) determines duration of reversals (Supplementary Fig. 5A). In sum, a rise in RIS Ca\(^{2+}\) preceded slowing and reversals, as also indicated by the event probabilities (Fig. 6a). We conclude that RIS activity precedes the behavioral change and may determine it.

We further analyzed how RIS activity is associated with behavioral changes by assessing the fraction of animals performing particular locomotion behaviors at and around the timing of RIS Ca\(^{2+}\) peaks (Fig. 6d,i). Before RIS became active, there was a similar propensity for animals to accelerate or to slow down. During the Ca\(^{2+}\) rise, animals were more likely to decelerate, and following the Ca\(^{2+}\) peak, reversal probability increased. Thus the likely moment of RIS releasing GABA and FLP-11 neuropeptides is correlated with, and most likely induces, inhibition of locomotion, preceding reversals, analogous to our observations upon RIS photostimulation (Fig. 1b). We conclude that RIS fulfills a crucial role in neuronal programs controlling forward–backward transitions.

**FLP-11 neuropeptides are required for RIS' effects on locomotion speed.** RIS affects locomotion slowing and is particularly active before reversals. As FLP-11 neuropeptides were required for stopping, we asked whether in *flp-11* mutants RIS Ca\(^{2+}\) activity occurs but may be unable to evoke behaviors. From 24 *flp-11* animals, we recorded 100 reversals (Supplementary Fig. 5B) and 25 Ca\(^{2+}\) events (Fig. 6e–h). Overall occurrence of reversals or Ca\(^{2+}\) events did not differ between WT and *flp-11* (Fisher’s exact tests, two sided, \(p = 0.865\) and 0.055, respectively, n.s.), thus network functions appeared normal. As in WT, *flp-11* mutants displayed similarly skewed probability distributions for reversals and Ca\(^{2+}\) peaks (Fig. 6e). Reversal induction per se is independent of FLP-11. Slowing, on average, was less pronounced during RIS Ca\(^{2+}\) rise events, which were less frequently paired to slowing or reversals; thus reversal velocity (Fig. 6f) and speed (Supplementary Fig. 6B) were not significantly different before and during the Ca\(^{2+}\) rise. No significant cross-correlation of the Ca\(^{2+}\) rise rate and velocity was found (Fig. 6g). *flp-11* mutants did not show significant changes in the proportion of forward, reverse, accelerating, or slowing subsequent to a Ca\(^{2+}\) transient, while these proportions were significantly different between WT and *flp-11*, and *flp-11* animals reversed more prior to RIS Ca\(^{2+}\) signals (Fig. 6d, h, i). In sum, locomotion, particularly slowing, is abnormally regulated in *flp-11* mutants. RIS tight control of the timing of reversals after a stop requires FLP-11 release, which also aids in sustaining stops (Fig. 4a), while it elicits fast slowing by GABA. Yet, as reversals occur after RIS photostimulation even without GABA and FLP-11, additional neurons must partake in inducing reversals.

**Compartmentalized Ca\(^{2+}\) dynamics in the RIS axon.** In *flp-11* mutants, RIS was on average less efficient in causing slowing. This was intriguing, since we could still observe occasional reversal events. We asked whether there is a specific feature of RIS Ca\(^{2+}\) activity that distinguishes such events. We segregated the RIS Ca\(^{2+}\)-aligned events into those paired with a reversal and those that merely led to a velocity reduction or stop. We then assessed Ca\(^{2+}\) signals spatiotemporally, from soma to nerve ring, encompassing also the branch, focusing on the 2 s centered on the Ca\(^{2+}\) event. To capture dynamic changes, Ca\(^{2+}\) signals were normalized to the first 150 ms of this time window, and significantly different Ca\(^{2+}\) levels were color coded (Fig. 7).

In the RIS nerve ring region, we observed Ca\(^{2+}\) signals for both types of events, i.e., paired (Fig. 5d, e.g., seconds 28–30; Fig. 7a) or unpaired to a reversal (Fig. 5d, e.g., seconds 21–23; Fig. 7b). Interestingly, events paired to a reversal showed significantly increased Ca\(^{2+}\) dynamics in the branch region (Fig. 7a; Supplementary Movies 7 and 8A), which were absent when animals only slowed or stopped (Fig. 7b; Supplementary Movies 7 and 8B). Furthermore, reversal–unpaired events showed a significant reduction in branch Ca\(^{2+}\) signals \(-750\) ms preceding the maximal nerve ring Ca\(^{2+}\) signal. In these events, no significant activity was observable in the branch, even when the nerve ring region was active (Fig. 7b). We conclude that RIS axonal Ca\(^{2+}\) dynamics are compartmentalized and that the branch region has a specific function in induction of, or concomitant with, reversals. RIS axonal Ca\(^{2+}\) in *flp-11(tm2706)* mutants had no significant dynamics during reversals (Fig. 7c), while events paired only to a stop still showed nerve ring activity, though no significant reduction in branch activity was observed (Fig. 7d). FLP-11 neuropeptides may directly or indirectly provide positive feedback to RIS during reversals.

**Discussion**

Neuronal circuits regulate and fine-tune locomotion. While in mammals this is orchestrated by whole-brain systems like motor and prefrontal cortex, cerebellum, and spinal cord neurons, much fewer neurons must fulfill these tasks in compressed nervous systems. Here we analyzed the role of one neuron, RIS, which orchestrates locomotion slowing and reversals in *C. elegans*. RIS does this by re-employing a peptidergic pathway used in sleep control, e.g., after larval molts or in response to stress. Here RIS uses also GABAergic signaling, not required during DTS induction. During optogenetic stimulation, GABAergic signaling from RIS induces the locomotion stop within seconds, possibly by inhibiting RIB neurons (Supplementary Fig. 3B, C), and sustains it by FLP-11 signaling (Fig. 8a). The observed axonal compartmentalization of Ca\(^{2+}\) dynamics during locomotion (Fig. 8b) jointly with the requirement of peptidergic and GABAergic signaling suggest that RIS uses both modalities in a spatiotemporally defined manner to control locomotion. RIS' role in both locomotion and sleep control represent two types of temporally different activity: (1) where RIS is employed to control locomotion with brief activity bouts (seconds), in coordination with other locomotion neurons, and (2) where DTS and SIS involve long-term RIS activity (minutes to hours), together with the sleep neuron ALA.  

RIS depolarization in the nerve ring results in fast GABA- and FLP-11 neuropeptide-mediated inhibition of neuronal activity and locomotion. Suppression of branch depolarization is permissive for locomotion halting, and conversely, if the RIS axon is not hyperpolarized in the branch before depolarization in the nerve ring, the most likely behavioral outcome is a reversal. This coincides with optogenetic experiments where RIS depolarization led to reversals and required both GABA and FLP-11 release. These transmitter(s) dampened cholinergic MN oscillations and desynchronized acetylcholine release, thus suppressing oscillations of body muscle activity. Occasionally reversals occurred also without RIS activity or in the absence of FLP-11. Thus RIS is sufficient but not essential for reversals and...
must act in coordination with other neurons. Roberts et al. described a four-state model of *C. elegans* locomotion, where forward–reverse and vice versa transitions were connected by brief pause states. RIS may induce this brief stop. Hints of this are found in our analyses, e.g., pause events at seconds 13, 44, and 56 in Fig. 5c, accompanied by RIS Ca$^{2+}$ peaks. Also, in analyses of mean Ca$^{2+}$ events, a brief phase of zero acceleration followed the moment of maximal slowing (Supplementary Fig. 6C).

The functional compartmentalization of the RIS axon indicated that the branch is instructive for reversal onset. Since the effects of RIS activation are rather fast and the RIS axon extends only in the nerve ring and branch regions, with the branch being mainly post-synaptic, and the nerve ring process being dominated by output synapses. The branch connects to only three neuron types: AVJ, PVC, and SMD. SMD neurons are part of the network encoding the amplitude of events, a brief phase of zero acceleration followed the moment of maximal slowing (Supplementary Fig. 6C).

![Image](https://example.com/image1)

**Fig. 7** Compartimentalized Ca$^{2+}$ dynamics in the RIS process. **a, c** Ca$^{2+}$ events paired to a reversal in the 2 s time window flanking a Ca$^{2+}$ signal obtained from freely behaving animals (WT in **a**, *flp-11(tm2706)* in **c**). Top is the dorsal nerve ring (NR) region, passing through the branch region (Br) to the center of the cell body (CB). Only events significantly different from inactivity are shown, color coded for intensity, normalized and relative to the mean of the first 150 ms depicted. Blue and red hues indicate significant reductions and increases, respectively. Non-significant dynamics are omitted and shown in white. **b, d** Ca$^{2+}$ events unpaired to a reversal in the 2 s time window flanking a Ca$^{2+}$ signal, as in **a, c**. Number of events indicated. Significance level: *T* Test, *p* < 0.05.
brief activity and sleep during prolonged activity. In more complex brains, such functions dissociate into different systems, distinct for sleep control and locomotion stop, which reside in different cell types (molecular identities still need to be determined for most systems) and brain structures, and use different mechanisms. Here more elaborate control regimes have evolved, where sleep neurons induce relaxation, while stop neurons halt locomotion but do not relax muscle. RIS makes this distinction by brief and short signaling via the same molecules. RIS may thus represent a primordial neuronal pathway, e.g., for fast sensory integration, and may have evolved to more finely tuned stop neurons in vertebrates.

Methods

Molecular biology. pCaS6 (pغلر-2::flox::ChR2(H134R)::mCherry::SL2::GFP) was a gift from Cornelia Schmitt. The following plasmids were kindly provided by Navin Pokala (Bargmann lab, Rockefeller University, USA): pNP165 (pSM::pغلر-1::flox::ChR2(H134R)::mCherry), pNP260 (pSM::pmrn1::flox::ChR2::STAR::mCherry), and pNP259 (pSM::pgpa-14::Cre). pPD95.79 was a gift from Andrew Fire (Addgene plasmid # 1496; RRID:Addgene_1496). pOT15 (QUAS::JRCaMP1b), pOT16 (punc-17::QF), and pOT17 (punc-4::QF::mCherry) were provided by Oleg Tolstenkov. We used the plasmid pmyo-2::RCAmpMP1h for RCaM expression in BWM cells. pmyo-2::GCaMP3 was kindly provided by Karen Erbguth. pGP (pnc-4::GCaMP6::SL2::TagRFP) was kindly provided by Douglas S. Kim. pCG02 was linearized with SacII and ligated to the PCR sequence from pWSC28, both cut with SphI and Not-1 HF. pCG03 (pggl-2::flox::GCaMP6::SL2::RFP) was kindly provided by Oleg Tolstenkov and was linearized with SacI and ligated to the PCR product of pWSC24 with forward primer oCG11 (AAAGAATTCGGTACCGAT).
to pWSC15, cut with HindIII and KpnI. PCR for pharyngeal terminal bulb marker

Reagents

GCaMP3) pmyo-2::GCaMP3 was cut with MscI and SphI and ligated to the

supplemented to

that, at elevated temperature of 25 °C, conditional transgene expression for RIS was

AGGCACCA), A (ATGGTCTCAAAGGGTGAAG), D (ACGACGGCCAGTGAATTATC),

C (ATGGTCTCAAAGGGTGAAG), D (ACGACGGCCAGTGAATTATC),

GTACCT

AACGCGGACCAAGGACCA

GGT), C (ATGGTCTCAAAGGGTGAAG), D (ACGACGGCCAGTGAATTATC),

fl

ox::ChR2(H134R)::mCherry::SL2::GFP); pggr-1::nCre; pncx-10::mCherry

p-11(tm2706)X; lite-1(ce314)X; zxEx360[pggr-2::

fl

ox::GCaMP6s::SL2::tagRFP; pggr-1::nCre; pncx-10::mCherry]

Ca2+ imaging microscopy setp. An inverted fluorescence microscope (Axiovert 200, Zeiss, Germany) equipped with an MS 2000 motorized stage and PhotoTrack quadratic photomultiplier tube (both Applied Scientific Instrumentation, USA) was utilized for Ca2+ imaging, similar to a system described earlier. As excitation light sources, two high-power light-emitting diodes (LEDs; 470 and 590 nm wavelength, KSL 70, Rapp Optoelektronik, Germany) or a 100 W HBO mercury lamp were used. Simultaneous dual-wavelength activation was enabled by a Photometrics DualView-A beam splitter in combination with a Hamamatsu Orca Flash 4.0 CMOS camera controlled by HCImage Live (Hamamatsu) or MicroManager (version 1.4.13; http://micro-manager.org) software.

Ca2+ imaging in immobilized worms. Animals were immobilized on 10% M9 agar pads with polystyrene beads (Polysciences, USA) and imaged either by x25 (BWM) or x40 (nerve ring) objective lenses. The following light filter settings were used: GFP/mCherry Dualband ET Filterset (FS6-019, AHF Analysetechnik, Germany) was combined with 532/18 nm and 625/15 nm emission filters and a 565 long-pass beam splitter (FS9-833, FS9-624, and F48-567, respectively, all AHF). ChR2 stimulation was performed using 1 mW/mm² blue light. To measure Ca2+ fluorescence, animals were acquired at 31 ms exposure time and 20 fps. Light illumination protocols were acquired at 31 ms exposure time and 20 fps. Light illumination protocols were generated by a Lambda SC Smart shutter controller unit (Sutter Instruments, USA), using its TTL output to drive the LED power supply or to open a shutter when using the HBO lamp. Video synchronization was achieved by cropping the acquisition to obtain equally sized time bins before, during, and after blue light exposure. Image analysis was performed in ImageJ (NIH), called by a custom KNIME workflow, after manual annotation of ROIs. Spline ROIs were selected for the BWM cells or ventral nerve cord to nerve ring region and the kymographs were generated by averaging a 7 × 7 × 7 moving voxel (where time is the third dimension) across the ROI. A ventral and a dorsal BWM ROI were defined, and their periodicity data were averaged for the analysis of a single animal. Additionally, an elliptic ROI was selected for background fluorescence exclusion. Mean intensity values for each video frame were obtained and background fluorescence values were subtracted from the fluorescence values derived for RCaMP. Subtracted data were normalized to $\Delta F/F = (F_0 - F)/F_0$, where $F_0$ represents the intensity at the given time point and $F$ represents the average fluorescence of the first second of the acquisition.

Signal autocorrelation analysis. Since spontaneous activity was not synchronized between animals, we introduced the metric of signal autocorrelation for calcium imaging as a means of describing perturbations in the Ca2+ dynamics of both BWM and RIS neurons. We observed that Ca2+ dynamics in immobilized animals were periodic, albeit without a fixed frequency. This periodicity could be obtained by analyzing the signal correlation to itself, where peaks in the autocorrelation function were indicative of the period of Ca2+ dynamics. This was performed by a custom script in R called by a KNIME workflow. The R script calculated the autocorrelation of a smooth spline fit of the $\Delta F/F$ signal and searched for its peaks in window of 20 s of data to calculate the period of the autocorrelation. If no period could be found, it was set as the maximum of 1 in every 20 s. Hence, this function

C. elegans strains. The following strains were used or generated for this study: N2 (WT isolate, Bristol strain) and transgenic animals were cultivated on either nematode growth medium (NGM) or high growth medium plates seeded with Escherichia coli (WT isolate, Bristol strain), prior to seeding for optogenetic experiments. Transgenic animals were cultivated on either nematode growth medium (NGM) or high growth medium plates seeded with Escherichia coli (WT isolate, Bristol strain)

ARTICLE
Axonal Ca\(^{2+}\) imaging of RIS in moving animals. For measuring spontaneous Ca\(^{2+}\) activity, a tracking system for single neuron Ca\(^{2+}\) imaging in moving animals\(^{59}\) was modified to allow axonal visualization with an improved temporal accuracy and signal-to-noise ratio. Animals expressing a GCaMP6 indicator exclusion filter (F39-019, AHF Analysentechnik, Germany) were used to visualize the RIS neuron. Fluorescent measurements of GCaMP6 and mCherry were enabled using a GFP/mCherry Dualband ET Filterset (F36-019, AHF Analyseentechnik, Germany), combined with 522/18 nm and 625/15 nm emission filters and a 565 long-pass beam splitter (E39-833, F39-624, and F48-567, respectively, all AHF). Tracking the animal’s head was established by the PhotoTrack system (Applied Scientific Instrumentation, USA) that automatically repositions the motorized XY stage to keep a bright fluorescent marker in center of the field of view (FOV). By keeping the relative signal strength from each of the four segments of the RIS neuron (by binarizing a smaller ROI after masking the gut). Subsequently, also the dorso-ventral axis was determined. In this way, an ROI containing regions of the axon (by binarizing a smaller ROI after masking the gut). Subsequently, also the dorso-ventral axis was determined. In this way, an ROI containing regions of the axon was always oriented in the same direction. At last, a parabola was fitted through the neuronally shaped image components. Alongside this parabola, about 25 equally spaced points were determined to generate rectangular ROIs perpendicular to it. Mean fluorescence and the brightest fluorescent intensity during the event-aligned 6 s time windows of both normalized fluorescent change and acceleration time shifted up to 3 s time lags.

Electrophysiology. Recordings from dissected BWM cells anterior to the vulva at the ventral side are described in ref. \(^{10}\). Light activation was performed using an LED lamp (W60-70, Rapp OptoElectronic, Hamburg, Germany; 470 nm, 8 mW/mm\(^2\)) and controlled by an EPC10 amplifier and Patchmaster software (HEKA, Germany). mPSC analysis was done by the Mini Analysis software (Synaptosoft, Decatur, GA, USA, version 6.0.7). Amplitude and mean number of mPSC events per second were analyzed for each 10 s before, 10 s during, and 10 s after illumination.

Electropharyngeograms. EPG recording was performed as previously described\(^{89}\). Animals were selected on the day prior to measurement as L4 larvae. The head was cut away from the body with a scalp at about one third to half of the body length. The tip of the worm head was sucked under 100-fold magnification into an EPG-suction electrode, connected via a silver chloride-coated silver wire to an EPC-10 amplifier (Heka, Germany). Prior to measurement, the preparation was incubated in 2 μM serotonin for 5 min to induce pharyngeal pumping. EPG recordings were performed by the PatchMaster software (Heka). We recorded spontaneous pumping for 30 s prior to 30 s of RIS-CHr2 depolarization via 3 mW/mm\(^2\) (470 nm) light and further 30 s post stimulus. We used the Review software (Bruxton Corporation, USA) to convert from PatchMaster to ABF files. Pump rate and duration were analyzed by AutoEPG58 (kindly provided by C. James, Embody Biosignals Ltd., UK).

Statistics. Statistical analysis was performed in Prism (Version 5.01, GraphPad Software, Inc.), Mathematica (version 10, Wolfram Research, Inc., Champaign, IL, USA), OriginPro 2015G (OriginLab Corporation, Northampton, USA), R (version 3.3.2), the latter called by RStudio (version 1.0.136, RStudio, Inc.), or KNIME (Desktop version 3.5, KNIME.com AG, Zurich, Switzerland). No statistical methods were applied to predetermined sample size. However, sample sizes reported here are consistent to data presented in previous publications. Data were tested for normality prior to statistical inference. Data are given as means ± SEM when not otherwise stated. Significance between data sets after paired or two-tailed Student’s t test is given as p value (*p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001), when not otherwise stated. For other statistical tests used, see figure legends.

Data availability
All data are provided as Supplementary Information. Reagents and code are available upon request.

Code availability
Codes are available upon request.

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