Glutamate spillover in *C. elegans* triggers repetitive behavior through presynaptic activation of MGL-2/mGluR5

Menachem Katz¹, Francis Corson², Wolfgang Keil¹,³,⁵, Anupriya Singhal¹, Andrea Bae¹, Yun Lu¹, Yupu Liang⁴ & Shai Shaham¹

Glutamate is a major excitatory neurotransmitter, and impaired glutamate clearance following synaptic release promotes spillover, inducing extra-synaptic signaling. The effects of glutamate spillover on animal behavior and its neural correlates are poorly understood. We developed a glutamate spillover model in *Caenorhabditis elegans* by inactivating the conserved glial glutamate transporter GLT-1. GLT-1 loss drives aberrant repetitive locomotory reversal behavior through uncontrolled oscillatory release of glutamate onto AVA, a major interneuron governing reversals. Repetitive glutamate release and reversal behavior require the glutamate receptor MGL-2/mGluR5, expressed in RIM and other interneurons presynaptic to AVA. *mgl-2* loss blocks oscillations and repetitive behavior; while RIM activation is sufficient to induce repetitive reversals in *glt-1* mutants. Repetitive AVA firing and reversals require EGL-30/Gαq, an mGluR5 effector. Our studies reveal that cyclic autocrine presynaptic activation drives repetitive reversals following glutamate spillover. That mammalian GLT1 and mGluR5 are implicated in pathological motor repetition suggests a common mechanism controlling repetitive behaviors.

¹Laboratory of Developmental Genetics, The Rockefeller University, 1230 York Avenue, New York, NY 10065, USA. ²Laboratoire de Physique Statistique, Ecole Normale Supérieure, CNRS, Université Pierre et Marie Curie, Université Paris Diderot, 75005 Paris, France. ³Center for Studies in Physics and Biology, The Rockefeller University, 1230 York Avenue, New York, NY 10065, USA. ⁴Research Bioinformatics, The Rockefeller University, 1230 York Avenue, New York, NY 10065, USA. ⁵These authors contributed equally: Francis Corson, Wolfgang Keil. Correspondence and requests for materials should be addressed to S.S. (email: shaham@rockefeller.edu)
Glutamate spillover following synaptic release has been extensively studied as a mechanism for engaging non-synaptic glutamate receptors\(^1\). Such spillover can have physiological roles, such as feed-forward disinhibition of GABAergic interneurons\(^3\); and pathological consequences, including a suggested role in drug-seeking behavior\(^4\). Spillover is often studied by inducing high levels of extracellular glutamate, conditions that can also drive neurotoxicity\(^5,6\). Thus, teasing apart whether consequences of spillover are due to toxicity or to extrasynaptic signaling effects on neural circuits is challenging, particularly in the context of long-term behavioral studies.

The *Caenorhabditis elegans* nervous system is composed of only 302 neurons\(^2\), and viability of these cells is not affected by excess glutamate\(^8\). Thus, this animal provides an excellent arena with synapses are illustrated (inset). Through cyclic autocrine presynaptic activation. Our findings also

To establish a glutamate spillover paradigm in *C. elegans* that might be broadly informative, we aimed to determine whether the CEPsh glia also control glutamate levels; whether this control is exerted through active glutamate uptake, as in astrocytes\(^1,5,9\); and, if so, whether it has an effect on synaptic activities. We used transcriptome profiling to demonstrate molecular similarities between *C. elegans* CEPsh glia and murine astrocytes, and found that mRNAs for glt-1, encoding a homolog of GLT1/EAAT2, are enriched in these glia. Postembryonic ablation of CEPsh glia, or GLT-1 loss, promotes two different glutamate-dependent behavioral defects: an increased overall rate of locomotory reversal events, and a surprising clustering of reversals in repetitive bouts.

In vivo measurements of extracellular glutamate dynamics near postsynaptic sites of the interneuron AVA, a major regulator of reversal behavior, show that GLT-1 is required to prevent oscillations in synaptic glutamate release, and consequent oscillations in AVA activity, which correlate with repetitive reversal events in frequency. Furthermore, both oscillations in AVA neuron activity and repetitive behavior are mediated by activation of the metabotropic glutamate receptor MGL-2/mGluR5 in glutamatergic interneurons presynaptic to AVA. Aberrant repetitive AVA activity and reversal behavior require the excitatory egl-30/Gaq signaling pathway.

Our studies suggest a simple, yet general, mechanism by which aberrant glutamate spillover can induce repetitive behavior through cyclic autocrine presynaptic activation. Our findings also...
suggest a plausible circuit origin for repetitive behavior in the context of human neuropsychiatric illness.

Results
CEPsh glia share molecular similarities with astrocytes. To establish a glutamate spillover paradigm in C. elegans, we first sought to examine molecular similarities between CEPsh glia and astrocytes, particularly in the context of glutamate homeostasis. We therefore sequenced CEPsh glia mRNAs, and assessed their abundance and enrichment relative to RNA levels of all other C. elegans cells. We then compared the list of CEPsh-enriched genes with lists of mouse homologs enriched in different brain cell types using hierarchical clustering. We found that CEPsh glia cluster together with mouse neurons, astrocytes, and oligodendrocytes, and separately from microglia and endothelial cells (Fig. 1b). This is in line with the developmental origins of these cells: C. elegans CEPsh glia, and mammalian neurons, astrocytes, and oligodendrocytes, are all ectodermally derived, whereas microglia and endothelial cells are of mesodermal origin. Rank-ordering mRNAs, according to fold-enrichment (FE) within a cell type, reveals that the more highly enriched a gene is in CEPsh glia, the more likely is its ortholog to be enriched in murine astrocytes, but not in oligodendrocytes (or microglia; Fig. 1c, Supplementary Data 1). For example, some genes enriched in both cell types include jfa-4, homologous to the glial fibrillary acidic protein (GFAP), known to affect astrocyte morphology and activity; fmi-1, homologous to Flamingo (Celsr1), a regulator of axon migration and fasciculation; and glt-1, homologous to the astrocyte-expressed Slc1a2/GLT1/EAAT2 glutamate transporter.5,6 (Fig. 1d, Supplementary Data 1). Thus, of the different mouse brain cells in our comparisons, CEPsh glia are most similar to astrocytes by molecular, morphological, and functional criteria.

Since GLT1 is a known astrocyte regulator of glutamate homeostasis5, we further characterized glt-1 function in CEPsh glia. To confirm glt-1 gene expression in CEPsh glia, we generated animals carrying a transgene in which glt-1 upstream regulatory sequences drive expression of GFP. Strong GFP expression is observed in all four CEPsh glia (Fig. 1e). Weaker expression is also detected in muscles, consistent with a previous report18.

CEPsh glia-expressed GLT-1 regulates reversal behavior. Decreasing or increasing glutamate signaling in C. elegans using mutations blocking presynaptic glutamate release (e.g. in eat-4/VGLUT19), or mutations activating glutamate receptors20, respectively, is known to decrease/increase locomotory reversal rates, respectively. Thus, to study glial glt-1 regulation of glutamate signaling, we recorded the locomotion of animals exploring their environment, a behavior characterized by infrequent reversals21. By analyzing these movies using a Hidden Markov Model with data-derived transition probabilities between forward and backward locomotion states (Supplementary Fig. 1a), we found that postembryonic ablation of CEPsh glia (which does not affect neuronal viability or nerve ring architecture; Supplementary Fig. 2a–c), or a mutation in glt-1, aberrantly increases animal reversal rates to similar extents (Fig. 2a and Supplementary Fig. 1b–e; consistent with previous observations).8 These excessive reversals are attenuated in animals also carrying a mutation in the vesicular glutamate transporter eat-4/VGLUT (Fig. 2a). This is consistent with the notion that excessive reversals are dependent on presynaptic glutamate release. While CEPsh glia ablation induces additional locomotion defects15 (Supplementary Movies 1–2), glt-1 loss does not (Supplementary Movie 3). Thus, glt-1 modulates glutamate-dependent reversal behavior.

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Fig. 2 C. elegans reversal behavior is controlled by CEPsh glia-expressed glt-1. a Reversal rates in wild type (WT, black circles; n = 9 movies), CEPsh-glia-ablated (black boxes; n = 9 movies), glt-1(ok206) (black triangles; n = 9 movies), eat-4(ky5) (open circles; n = 8 movies), eat-4(ky5); CEPsh-glia-ablated (open boxes; n = 8 movies) and eat-4(ky5); glt-1(ok206) (open triangles; n = 7 movies) strains. b A deconvoluted optical section. GLT-1 fused to GFP expressed in CEPsh glia using hlh-17 promoter. Synapses are marked by glt-1(ok206):NLG-1::mCherry. Scale bar, 4 μm. c Cell-specific glt-1(ok206) rescue studies. Reversal rates normalized relative to minimum value of WT controls and maximum value of glt-1 mutants. Wild type (WT, open circles; n = 23 movies), glt-1(ok206) (black boxes; n = 25 movies), GLT-1 transgene expressed in neurons (magenta triangles; eat-4; n = 11), CEPsh glia (hlh-17, green triangles; n = 14), muscles (myo-3, blue diamonds; n = 18). a, c Bars, mean ± SEM, ANOVA Tukey’s HSD post hoc test, **p < 0.005, ***p < 0.0005. Source data are provided as a Source Data file.
behavior\(^\text{18}\), and lack of glia-expressed GLT-1 is likely the primary cause of reversal behavior defects in CEPsh glia-ablated animals.

Consistent with the idea that \(glt-1\) influences synapic glutamate signaling, we found that most glutamatergic synapses, marked by mCherry-tagged neuroligin (NLG-1) expressed in ionotropic glutamate receptor (\(glt-1\))-positive neurons, are within 0.5 \(\mu\)m of glial processes expressing GLT-1::GFP (Fig. 2b, Supplementary Fig. 3 and Movie 4). To determine the functional relevance of this synaptic proximity, we assessed the rescue of \(glt-1\) mutant behavioral defects in animals expressing GLT-1 protein specifically in either CEPsh glia (within 0.5 \(\mu\)m of synapses; using the \(hlh-17\) promoter element), or in muscles (within 2 \(\mu\)m of synapses; using the \(myo-3\) promoter element), whose arms synapse to motor neurons at the inner facet of the nerve ring, but do not penetrate the neuropil\(^7\). We found that glial expression of GLT-1 restores normal reversal rates more effectively than muscle-specific expression (Fig. 2c). Further supporting a proximity model, forcing GLT-1 expression directly at synapses, by ectopic expression in glutamatergic neurons (using the \(eat-4\) promoter element), where \(glt-1\) is normally not endogenously expressed, results in the most efficient rescue (Fig. 2c).

In summary, our data suggest that loss of \(glt-1\) provides an appropriate setting for studying the effects of increased extracellular glutamate, allowing us to interrogate effects of glutamate spillover on circuit function and behavior.

GLT-1 functions to prevent repetitive reversal behavior. To examine the behavioral consequences of \(glt-1\) loss in detail, we examined the temporal distribution of spontaneous reversal events of \(glt-1(–)\) animals moving on agar plates. Specifically, we analyzed the probability, \(p(t)\), that an animal performing a reversal event will not initiate a reversal over a subsequent period of time, \(t\). In wild-type animals, \(p(t)\) is well fit by a decaying single exponential function, consistent with a random Poisson process in which the probability of reversal per unit time is fixed (Fig. 3a). In \(glt-1(ok206)\) mutants, we find two deviations from the wild-type reversal pattern. First, because \(glt-1\) mutants have a higher average reversal rate (Fig. 2a), an exponential fit based on this average rate has a steeper slope than seen in the wild type (black line, compare Fig. 3a, b). Second, and more important, the experimentally observed \(glt-1(ok206)\) reversal distribution is not well fit by a single exponential function (compare red and black lines in Fig. 3b). In these mutants, the probability of reversal within 25–30 s following a reversal event is high, and the proportion of long reversal-free intervals following an initial reversal is reduced (Fig. 3b). Indeed, modeling \(glt-1\) mutant reversals with a fixed reversal rate, boosted by a perturbation at short times after a reversal, fits our data well (Supplementary Fig. 4a, b; see Methods). The weight of the perturbation defines a natural repetitive behavior index (RI), which we use to quantify the deviation from the Poisson process (Fig. 3d and Supplementary Fig. 4a, b; see Methods). This defines a general approach to

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**Fig. 3** GLT-1 prevents repetitive reversal behavior. a Probability that following a reversal, a wild-type animal will move forward for the time indicated on the horizontal axis. Exponential fits are plotted in black ± SEM, light gray lines are probability distributions of individual movies (blue, mean plot; \(n = 9\) movies). b Same as (a), except in \(glt-1\) mutants (red, mean plot; \(n = 9\) movies). a, b \(\chi^2\)-test used to determine if exponential fit differs from mean observed data (colored lines; n.s., nonsignificant; **\(p < 10^{-15}\)). c The residual difference between the mean exponential fit and the mean observed reversal probability for wild-type (WT; blue) and \(glt-1\) (red) mutants at the indicated time intervals. d \(glt-1\) mutants exhibit repetitive reversal behavior (from data in (a) (circles), (b) (boxes), Student’s \(t\) test, **\(p < 0.0005\)). e GLT-1 in CEPsh glia rescues repetitive reversals (ANOVA Tukey’s HSD post hoc test, *\(p < 0.05\), **\(p < 0.005\); WT (circles) \(n = 6\), \(glt-1(ok206)\) (boxes) \(n = 6\), \(glt-1(ok206)\); \(hlh-17\) promoter GLT-1 (triangles) \(n = 14\) movies). f \(glt-1\) mutants respond to mechanical stimuli with repetitive reversals (Student’s \(t\) test, **\(p < 0.005\); WT \(n = 16\), \(glt-1(ok206)\) \(n = 13\) animals). d-f Bars, mean ± SEM. Source data are provided as a Source Data file.
GLT-1 blocks oscillatory glutamate release and AVA activity.

To uncover the underlying circuit disruptions leading to repetitive behavior in glt-1 mutants, we focused on AVA, a major C. elegans interneuron driving backward locomotion. AVA expresses several glutamate receptors, and increased extracellular glutamate near AVA postsynaptic sites correlates with AVA calcium responses that induce reversal. We recorded spontaneous AVA activity in transgenic animals expressing both the fluorescent glutamate sensor iGluSnFR and the calcium indicator RCaMP. Animals were trapped in a microfluidic chamber, paralyzed using a muscle cholinergic receptor antagonist (tetrodotoxin), and continually exposed to food supernatant, which inhibits reversal behavior, and allows synchronization of our recordings. Wild-type animals initially show no glutamate release or calcium induction events in our assay. However, after 4 min ± 24 s (mean ± SEM) of habituation to the food cue, spontaneous glutamate release events onto AVA ensue, and these are highly correlated with AVA calcium transients (average peak cross-correlation 0.8; Fig. 4a and Supplementary Fig. 5a, Supplementary Movie 7). glt-1 mutants habituate more rapidly than wild-type animals (2 min ± 22 s on average), as might be expected for animals that have a higher propensity for activating glutamatergic signaling. Remarkably, glt-1 mutants often exhibit correlated long-duration oscillations of glutamate and AVA calcium signals following food cue habituation, suggesting repeated activation of AVA (Fig. 4b and Supplementary Fig. 5b, Supplementary Movie 8; average peak cross-correlation 0.7).

AVA calcium and glutamate signals are highly dynamic, with transient oscillations of varying frequencies and intensities, but quantitatively measure what has been anecdotally termed repetitive behavior in other settings. Confirming the importance of glia in glutamate clearance, expression of GLT-1 in CEPsh glia inhibits repetitive reversals in glt-1(ok206) mutants (Fig. 3e).

To investigate if GLT-1 is a general inhibitor of repetitive reversals, we examined reversals associated with the escape response to aversive stimuli. Anterior mechanical stimuli applied to wild-type C. elegans induce backward movement for ~3 body bends, followed by forward movement in a different heading. Similar stimulations of glt-1 mutants result in more consecutive reversals than in wild-type animals (Fig. 3f, Supplementary Movies 5,6). Thus, repetitive reversals in glt-1(ok206) mutants are induced in two distinct behavioral contexts (exploration and escape), demonstrating that GLT-1 functions generally to prevent repetitive reversal behavior.

Fig. 4 GLT-1 inhibits MGL-2-dependent oscillations in glutamate secretion and AVA activation. a Representative traces of spontaneous glutamate (light blue) and calcium (dark blue) dynamics near and in a wild-type AVA neuron, respectively. b As in (a), glt-1(ok206) mutant (glutamate, light red; calcium, dark red). c Time-dependent frequency spectrum of calcium traces of the wild-type animal in (a). d Time-averaged frequency amplitudes from the wavelet spectrum in (c). e, f As in (c, d), glt-1(ok206) mutant. g Mean time-averaged frequency amplitudes for WT (blue line; 20 traces), glt-1(ok206) (magenta line; 22 traces), mgl-2(tm355) (green line; 15 traces) and glt-1(ok206); mgl-2(tm355) (orange line; 17 traces). h Frequency amplitude differences between indicated strain pairs. Shading, areas of significant difference p < 0.05, permutation test from bootstrapped ensembles, n = 104 bootstrap samples. i Representative trace of head motion in nonparalyzed glt-1 mutant animal between forward (positive) and backward (negative) positions. Traces are decorated in green, magenta for increased, decreased calcium signals, respectively. Circles, transition points. Source data are provided as a Source Data file.
exhibit common features among animals of the same genotype. To quantify these common features, we used wavelet analysis to assign local oscillation frequencies at each time point, generating time-dependent frequency spectra (see Methods). This analysis reveals the presence of mostly low frequencies (long duration between peaks) for the calcium responses seen in wild-type animals (<0.02 Hz; Fig. 4a, c, d; Supplementary Fig. 5a, c). The repetitive calcium activities seen in the glt-1 mutant, however, exhibit pronounced enrichment of oscillations at ~0.04 Hz (Fig. 4b, e, f; Supplementary Fig. 5b, d). This enrichment can be clearly seen in time-averaged frequency amplitudes, averaged over all recorded animals (Fig. 4g, compare magenta and blue curves), demonstrating significant enrichment of oscillations above 0.03 Hz (peak at 0.04 Hz; Fig. 4b, top left). Importantly, these frequencies correspond to calcium induction events peaking at ~25 s intervals, matching the time interval between consecutive reversal events in our behavior assays (Fig. 3c). Supporting this correlation, nonparalyzed animals in our microfluidic setup show strong association between AVA calcium oscillations and attempted transitions between reversal/forward movements (phi coefficient = 0.71; Fig. 4i). Together, these studies are consistent with the idea that glutamate oscillations drive repetitive reversal behavior events in glt-1 mutants.

**MGL-2 drives oscillatory glutamate release and AVA activity.** To further support this idea, and to elucidate the mechanism of oscillatory glutamate release onto AVA, we sought mutants abolishing glt-1-dependent glutamate release onto and calcium oscillations in AVA. We reasoned that repeated glutamate secretion in glt-1 mutants could be mediated by extra-synaptic neuronal glutamate receptors responding to glutamate spillover. The enrichment of 0.03—0.1 Hz oscillations (indicating activity peak separation of seconds to tens of seconds) suggest that receptors with slow activation/signaling kinetics, such as metabotropic glutamate receptors (mGlus)27,28, may be involved. C. elegans encodes three mGlur homologs, MGL-1, MGL-2, and MGL-3, corresponding to mammalian subgroups II, I, and III, respectively29. Loss-of-function mutations in mgl-1, -2, or -3 do not result in significant changes in glutamate release onto or calcium accumulation in AVA (Fig. 4g, h and Supplementary Fig. 6a, c, d). However, loss of mgl-2 (but not mgl-1 or mgl-3) significantly suppresses the high-frequency oscillations of glt-1 mutants (Fig. 4g, h and Supplementary Fig. 6b—d). Thus, glutamate oscillations and subsequent AVA calcium transients require MGL-2 activity. Importantly, loss of mgl-2 in glt-1 mutants also significantly, but not completely, ameliorates repetitive reversal behavior (Fig. 5a), supporting the idea that glutamate oscillations drive repetitive reversal bouts. The incomplete reduction in repetitive reversal may indicate that additional glutamate-related mechanisms are involved in this process.

Together, our observations support the notion that repetitive reversals in glt-1 mutants are caused by oscillations in glutamate release onto AVA.

**Fig. 5 Glutamate spillover induces repetitive reversals by presynaptic activation of MGL-2.** a mgl-2 mutation ameliorates increased repetitive reversals of glt-1 mutants. The effect is reversed by re-expression of MGL-2 specifically in RIM neurons (Kruskal–Wallis ANOVA with Dunn’s multiple comparison test, n.s., nonsignificant, **‘p < 0.0005, ’p < 0.05; WT, black circles, n = 30, mgl-2(tm355), black boxes, n = 13, tdc-1pro::MGL-2, black triangles, n = 25, glt-1(ok206), open circles, n = 21, glt-1(ok206); mgl-2(tm355), open boxes, n = 24, glt-1(ok206); mgl-2(tm355); tdc-1pro::MGL-2, open triangles, n = 25 movies). b MGL-2 expression in AIB neurons rescues repetitive reversals (WT, black circles, n = 21, glt-1(ok206), black boxes, n = 21, glt-1(ok206); mgl-2(tm355), black triangles, n = 22, glt-1(ok206); mgl-2(tm355); inx-1pro::MGL-2, open triangles, n = 10 movies). c Channelrhodopsin activation of RIM promotes GLT-1-mediated repetitive reversals (WT, black circles, n = 11, glt-1(ok206), black boxes, n = 12, glt-1(ok206); mgl-2(tm355), black triangles, n = 10 movies). d Bars, mean ± SEM. e ANOVA Tukey’s HSD post hoc test, **’p < 0.0005, ’p < 0.05. In (c), while not significant at the level of p < 0.05, the trend towards fewer repetitive events in animals also lacking mgl-2 is consistent with our observations in (a, b). d GFP-tagged MGL-2 localizes to RIM neurite (left), but not to mCherry-labeled synapses (middle); merge (right). Scale bar, 5 μm; nerve-ring region is indicated with dotted line; 2-μm-thick sum intensity projections. Source data are provided as a Source Data file.
MGL-2 acts extrasynaptically in neurons presynaptic to AVA. If glutamate spillover drives oscillatory glutamate release and consequently repetitive behavior in *glt-1* mutants, we might expect MGL-2 to function in neurons presynaptic to AVA. Indeed, we found that *mgl-2* is strongly expressed in RIM, a glutamatergic interneuron presynaptic to AVA, and more weakly in AIB, RIA, ADA and AIM, additional glutamatergic presynaptic neurons (Supplementary Fig. 7a and Supplementary Table 1). MGL-2 is not expressed in AVA or in CEPsh glia. Expression of MGL-2 specifically in RIM using a *tdc-1* promoter::mgl-2 transgene, can significantly increase repetitive behavior in *mgl-2*; *glt-1* double mutants to a level that is similar to the behavior of *glt-1* mutants alone (Fig. 5a). Similarly, MGL-2 expression in AIB, using an *inx-1* promoter element, also increases repetitive reversals in the double mutant, but to a lesser extent (Fig. 5b). These observations suggest a role for MGL-2 as a general amplifier of repetitive behavior upon glutamate spillover. Ectopic overexpression of MGL-2 in RIM can even induce repetitive reversals in animals with intact *glt-1*, probably due to hyper-sensitization of RIM to lower levels of extracellular glutamate (Fig. 5a). Moreover, MGL-2 expression in RIM restores glutamate and calcium oscillations in *mgl-2*; *glt-1* double mutants (Supplementary Fig. 7b, compare animals with and without transgene array). Rescue variability in these latter studies is high, likely because of variations in transgene copy number and expression.

We next asked whether RIM activation is sufficient to induce repetitive reversals when glutamate clearance is impaired. In agreement with a previous report, light activation of channelrhodopsin-expressing RIM promotes single reversal events in a retinal-dependent manner in wild-type animals (Fig. 5c and Supplementary Fig. 7c). RIM activation in *glt-1* mutants, however, can induce 2–3 short reversal events during light exposure, and these repetitive events appear to be partially reduced by loss of *mgl-2* (Fig. 5c). This observation indicates that spillover of glutamate, autonomously secreted by RIM, can feedforward onto RIM to induce repetitive behavior. It also suggests that RIM is unlikely to modulate activity of its presynaptic neurons to induce repetitive behavior.

Finally, the repetitive behavior of *glt-1* mutants can be elicited by glutamate spillover from RIM output synapses, MGL-2 should localize near but not at these synaptic sites. Indeed, GFP-tagged-MGL-2 exhibits punctate expression in the RIM neurite, adjacent to, but not overlapping glutamatergic synapses (Fig. 5d).

Taken together, these observations suggest that MGL-2 functions extrasynaptically in glutamatergic neurons presynaptic to AVA to mediate repetitive reversals caused by glutamate spillover.

**Repetitive reversal behavior is mediated by Gaq signaling.** How might MGL-2 effect glutamate release? The postsynaptic signaling mechanism of mammalian mGluR5, related to MGL-2, through Gaq has been extensively explored. However, how mGluR5, which localizes to both dendritic and axonal sites, functions presynaptically is not well understood. We therefore investigated whether Gaq might mediate the presynaptic effects of MGL-2. In *glt-1* mutant animals also defective in *egi-30/Gaq* signaling, repetitive reversals are nearly entirely abrogated (Fig. 6a, 6b). In these double mutants, and even single *egi-30/Gaq* mutants hardly show calcium activity (Supplementary Fig. 8b; hence waveform analysis cannot be applied for these activity traces). These observations identify EGL-30/Gaq as a required mediator of both reversal initiation and repetitive reversal behavior. To directly examine these roles, we expressed EGL-30 in RIM neurons of either *egi-30* single mutants or *egi-30; glt-1* double mutants. Transgenic *egi-30* single mutants exhibit a low reversal rate similar to nontransgenic animals (Supplementary Fig. 8c), suggesting that *egi-30* expression in RIM is not sufficient to stimulate reversal initiation. However, RIM::EGL-30 expression in *egi-30; glt-1* double mutants does significantly increase repetitive
behavior (Fig. 6b). Thus, EGL-30 functions in RIM to induce repetitive behavior, and outside of RIM to promote reversal initiation. Consistent with these observations, the goa-1/Gao pathway in C. elegans inhibits the egl-30/Gaq pathway33, and loss-of-function goa-1(n1134) mutants exhibit increased reversal rates (Supplementary Fig. 8a), glutamate secretion, and AIB activation events (Fig. 6c, d and Supplementary Fig. 8b), even when glt-1 is intact. Importantly, a strong increase in repetitive reversals is observed in goa-1; glt-1 double mutants compared to glt-1 single mutants alone (Fig. 6a), consistent with increased propensity for glutamate release. These findings indicate that repetitive reversals are induced through presynaptic excitations promoted by egl-30/Gaq and inhibited by goa-1/Gao.

Discussion
Taken together, our data suggest a model (Fig. 7) in which conditions permitting glutamate spillover beyond synapses, such as impaired clearance by glial GLT-1, allow glutamate to bind extrasynaptically localized MGL-2/mGluR5 on presynaptic neurons. This induces presynaptic neuron activation, via Gaq, promoting a stimulus-independent glutamate secretion event that leads to firing of the postsynaptic reversal command interneuron AVA. The cycle then repeats, driving repetitive reversal behavior. Termination of repetitive bouts could occur once synaptic vesicle pools are depleted. Consistent with such an autocrine excitatory feedback model, calcium increases in the presynaptic RIM or AIB and postsynaptic AVA neurons are known to be correlated30.

To generate oscillations, our model predicts that glutamate clearance from the glt-1(−) synapses should occur more rapidly than signaling through MGL-2/mGluR5. This idea is plausible, as glutamate has been estimated in other settings to diffuse at millisecond rates (~0.4 μm²/sec34), whereas signaling kinetics downstream of metabotropic glutamate receptors have been measured at seconds or tens of seconds27,28.

While the studies we present here mainly dissect RIM-AVA synapses, the proximity of CEPsh glia membranes to other glutamatergic synapses, coupled with expression of MGL-2 in other glutamatergic neurons, suggests that repetitive reversal behavior could be initiated at other synapses, whose activity is then propagated through the circuit to AVA. A recently described method for global analysis of neuron activation in C. elegans26 could be used to test this idea.

The studies we present here implicates uncontrolled extrasynaptic glutamate levels as a driver of repetitive behavior. In that context, it is interesting that impaired glutamate signaling in humans has also been implicated in the pathophysiology of disorders accompanied by repetitive behaviors35,36. Clearance of extracellular glutamate by active transport might be very relevant here. Linkage and copy number variation analyses implicate chromosome regions 11p12–p13 in Autism Spectrum Disorder (ASD)35, with SLCIA2 (GLT1/EAAT2) located close to maximal linkage peaks. Consistent with this, WAGR (Wilms tumor, Aniridia, Genitourinary malformations and mental Retardation) syndrome patients show features of autism and have deletions in this transporter gene37. Furthermore, mutations in the endosomal cation/proton antiporter NHE9, which promotes GLAST/EAAT1 membrane trafficking in astrocytes, confer susceptibility to ASD38. SLC1A2(EAAC1), a neuronal glutamate transporter gene, is associated with Obsessive Compulsive Disorder (OCD)38 and ASD39, and clinical trials for OCD treatment involve drugs that target glutamate signaling, which might affect glial uptake of glutamate36.

Murine repetitive grooming is a common model for the study of mechanisms regulating pathological repetitive behavior. Consistent with the repetitive behavior mechanism we describe here, astrocyte-specific knockout of GLT-1 in mice produces repetitive grooming behavior40. Additionally, mGluR5 inhibitors rapidly suppress repetitive grooming in mouse models for OCD and ASD41,42. A previous connection between GLT-1 and mGluR5, however, had not been noted. Furthermore, the underlying circuitry driving repetitive grooming behavior mediated by modulating these mouse proteins is not understood in detail. That the same proteins act together in repetitive behaviors in C. elegans and mice is striking.

Thus, while the specific motor behaviors subject to repetitive engagement differ dramatically among humans, mice, and C. elegans (e.g. head banging, grooming, and reversals, respectively), our results suggest that a common underlying physiology may be at play. Our results also suggest that physiological and pathological repetitive behaviors may be appropriate settings in which to investigate the behavioral consequences of glutamate spillover in mammals.

Our study suggests intriguing similarities between the C. elegans CEPsh glia and mammalian astrocytes. Like vertebrate glia, the CEPsh cells are derived from neuroepithelial precursors, but are not neurons themselves43. The four CEPsh glia tile around the nerve ring, the brain neuropil of the animal, suggesting morphological similarities with vertebrate astrocytes tiling across the central nervous system7,44. Here, we demonstrate that CEPsh glia are structurally elaborate, projecting fine processes that penetrate the nerve ring and that can be found near synapses, as are vertebrate astrocyte processes. Our gene-expression profile also reveals gene-expression similarities between C. elegans CEPsh glia and vertebrate astrocytes. Furthermore, our studies exploring the function of the CEPsh glia-enriched GLT-1 transporter demonstrate similar roles to those of astrocyte-enriched GLT1/EAAT2 in preventing repetitive behavior40. Previously, we found that like mammalian astrocytes, CEPsh glia also regulate sleep behavior15,45. Thus, C. elegans is a suitable setting for investigating aspects of astrocyte function in neuronal circuit activities and behavior. Moreover, these findings emphasize the limitations of predicting behavior solely based on mapping of neuronal connectomes, and suggest that glial architecture around synapses and their molecular properties should be integrated into such models.

Methods
Strains. C. elegans strains were cultured at 20 °C unless otherwise indicated. Wild-type (WT) animals were Bristol strain N2. Other strains used in this work are: OS1914: mls105 (hbl-17;pro::GFP) OS1965: mls105; mls108 (ptr-10).
DNA for gene expression. For cell-specific gene expression, the following promoter regions were used. DNA fragments were PCR amplified from C. elegans genomic DNA or existing plasmids:

\( hli-17 \): A 4.5 kb region upstream of the ATG

\( glt-1 \): A 3.5 kb region upstream of the ATG

\( mgl-3 \): A 2.4 kb region upstream of the ATG

\( eat-4 \): A 4 kb region upstream of the ATG

\( hli-17 \): A 4.5 kb region upstream of the ATG

\( glt-1 \): A 1 kb region upstream of the ATG

\( myo-3 \): A 2.4 kb region upstream of the ATG

\( spa-1 \): A 1.1 kb region upstream of the ATG

\( mgl-2 \): A 4.5 kb region upstream of the ATG

\( elt-2 \): A 4.5 kb region upstream of the ATG

\( elt-2 \): A 4.5 kb region upstream of the ATG

\( ptr-10 \): A 4.5 kb region upstream of the ATG

\( myrRFP \): A 4.5 kb region upstream of the ATG

\( hli-17 \): A 4.5 kb region upstream of the ATG

\( glt-1 \): A 4.5 kb region upstream of the ATG

\( glt-1 \): A 4.5 kb region upstream of the ATG

\( glt-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

\( glr-1 \): A 4.5 kb region upstream of the ATG

Glutamate and calcium imaging from AVA. Imaging of glutamate input to, and calcium signals in AVA neurons, in first-day adult animals, were performed in animals microinjected with 0.5 mM tetrodotoxin in a microfluidic chamber. Postembryonic cell ablation was performed using a reconstituted Caspase-3+48 were expressed in CEPsh glia starting in early L1 larvae. Ablation was determined by lack of fluorescence. Ablation was confirmed by lack of fluorescence. Ablation was confirmed by lack of fluorescence.
exhibited common features among animals of the same genotype. To characterize these features, we applied wavelet analysis to calcium signals, which more accurately captures the temporal and directional information than Fourier analysis. We used complex-valued Morlet wavelets composed of a Gaussian envelope and a plane wave

$$\phi(t, f) = \frac{1}{\sqrt{\pi} \sigma} e^{-t^2/(2\sigma^2)} e^{2\pi i f t}. \quad (2)$$

The wavelet spectrum $W(t, f)$ of a signal $S(t)$ is then given by

$$W(t, f) = \int dt' \phi(t-t', f) S(t'). \quad (3)$$

For a given frequency $f$, the width of the Gaussian was chosen as

$$\sigma = \frac{\xi}{2\pi f}. \quad (4)$$

with $\xi = 4$. This parameter choice enables detection of local frequency variations on scales larger than two periods while at the same time enabling robust estimation of the local frequency itself. Normalization with $1/\sigma$ ensures that wavelet coefficients can be compared across frequencies. We analyzed frequencies from 0.01 to 0.2 Hz. Edge effects of the wavelet transformations were minimized by padding the original signal with reflected signals on both ends.

To determine common temporal features of spontaneous activity traces for a given genotype, we first time-averaged the wavelet spectrum of each spontaneously active animal over the entire recording time and then subtracted it from the onset of spontaneous activity to the end of the 540 s recording period (see Fig. 4d, f). To obtain the mean time-averaged frequency amplitudes (e.g. Fig. 4g), we then averaged these time-averaged wavelet spectra over all animals of a given genotype, weighing each animal with the fraction of time it was spontaneously active during the recording period.

Plots were generated by an R script and include only the first 9 min of recordings, where the food cue was continuously present.

For correlation between animal head motion and AVA calcium signals, first-day adult animals were recorded for 8 min at 1 frame per second in the presence of food cue, without the addition of anesthesia in a microfluidic device. Head position was obtained through alignment of the position of the AVA neuron between consecutive frames using the Imagej Template Matching plugin, and extracting the difference in position between consecutive frames on the $x$-axis (motion on the $y$-axis is limited for animals trapped in the microfluidic chamber). Both the head position signal and the calcium signal were first Gaussian low-pass filtered (standard deviation, 3 s for position, 1.5 s for calcium). We then normalized the calcium signal to unit variance, calculated its first derivative and determined the times when the first derivative crossed a threshold of 0.1 from below or above. These times were considered transitions between high and low calcium states (magenta and green dots in Fig. 4i). We calculated the position difference during each high and low calcium state. For overall position increased, the animal was considered moving forward during a given calcium state and vice versa. Correlation between forward/backward motion and up/down states was assessed with Pearson’s coefficient.

### Analysis of worm locomotion

Twenty to forty first-day adult animals were placed on an NGM plate (1 mM NaCl; 1 mM CaCl; 1 mM MgSO₄; 1.7% agar; 0.25% glucose) and the velocity $v$ of the worm along the axis of its body. The orientation of the body (head vs. tail) is not visible in the images; thus, we arbitrarily choose the direction along the body axis that corresponds to positive increments in $x$. This choice was made for the first frame, then we keep track of the orientation such that the relation between changes in $x$ and direction of worm movement remains consistent. Whether forward movement corresponds to increasing or decreasing $x$ is determined as part of the inference.

The model comprises four behavioral states: forward crawling, backward crawling, forward turn, and omega turn. For the purpose of tracking the orientation of the body, the forward and backward crawling states are each subdivided into two states, depending on whether increasing $x$ corresponds to forward or backward movement (labelled $\rightarrow$ and $\leftarrow$). We do not attempt to keep track of the body orientation in the turning states, and they admit no such subdivision. In total, the model thus has six states (Supplementary Fig. 1a). Not all possible transitions are permitted: for instance, the sign ($\oplus$ or $\ominus$) must remain the same when the worm reverses, and an omega turn occurs only in the sequence backward crawling $\rightarrow$ omega turn $\rightarrow$ forward crawling.

For the inference of model states, we specify probability distributions for the observable variables in each state. Crawling and turning states are distinguished by the relative elongation $r$ of the body, defined as the ratio between the instantaneous elongation of the body and its average over a track. For a worm that is crawling forward or backward, $r$ is close to 1. When the worm makes a forward or omega turn, it bends its body and $r$ is less than one. Under the model, the distribution of $r$ for crawling worms is a Gaussian distribution with average one and standard deviation $\sigma$:

$$P(r(\text{crawling})) = \frac{1}{\sigma \sqrt{2\pi}} e^{-r^2/2\sigma^2} \quad (6)$$

where $\sigma = 0.075$ is close to the variance of the observed distribution in moving worms.

When the worm is turning, we simply take the distribution to be uniform over the interval $[0,1]$. For numerical reasons it is desirable for the distribution to be non-zero for $r > 1$, and we take this to distribute to decay as a Gaussian with the same standard deviation $\sigma$ as above:

$$P(r(\text{turning})) = \begin{cases} \frac{1}{\sigma \sqrt{2\pi}} e^{-r^2/2\sigma^2} & \text{if } r \leq 1 \\ e^{-r^2/\sigma^2} & \text{if } r > 1 \end{cases} \quad (7)$$

Forward and backward movements are distinguished according to the increment $\Delta x$ of a worm’s position between frames. Equivalent to specifying distributions for $\Delta x$ in each state, we specify probabilities for each state given a value of $\Delta x$.

Given a matrix of transition probabilities, the forward-backward algorithm allows efficient computation of the likelihood of state sequences and other quantities of interest, such as the number of transitions between pairs of states. The transition probabilities can thus be re-estimated from the output of the algorithm.

The Baum-Welch algorithm iterates rounds of the backward-forward algorithm to refine the state sequence estimates. The Baum-Welch algorithm increases monotonically at each step, guaranteeing convergence to a maximum. Thus, the transition probabilities are not free parameters of the model but estimated from the data.

To infer the absolute direction of movement, we use the fact that periods of backward movement are brief, much shorter on average than periods of forward movement. If we simply initialize the transition probabilities with a higher rate for transitions from backward to forward than from forward to backward, the inference procedure converges to a set of transition probabilities and state probabilities that assigns the direction (increasing or decreasing $x$) corresponding to the overall sequence of changes in $x$ of the worm during its forward movement with high probability (Supplementary Fig. 1b–d).

To characterize the duration of forward bouts, periods of backward or forward movement are defined as periods when the inferred probable of the backward crawling state is greater or smaller than one half, respectively. The distribution of forward and backward bout durations can be measured directly, but we may also estimate them when the worm exits the field of view or touches another worm. The distribution is thus estimated from the rule of change between forward and backward movement, as
function of time since the beginning of the bout. If \( P_r \approx (1 - \frac{t}{\tau}) \), the probability that a worm that moved forward for \( t \) time steps moves backward in the next step, is given by

\[
P_r(t) = \prod_{i=0}^{n-1} (1 - P_r(i)).
\]

(11)

Repetitive reversals were quantified based on the deviation between the observed distribution of forward bouts and the exponential distribution associated with a Poison process. To do so, the reversal rate \( r(t) \) observed after a time interval \( t \) moving forward was modeled as

\[
r(t) = r_0 + \frac{\mu}{\phi}
\]

(12)

where \( r_0 \) denotes a baseline, long-time reversal rate, and \( \phi \) is an enhancement in the reversal rate at short times, decaying over a time \( \tau \). The corresponding probability \( p(t) \) that a worm is still moving forward after a time interval \( t \) is given by

\[
\log[p(t)] = -r_0 t - \delta(t - e^{\frac{-t}{\tau}}).
\]

(13)

where \( \delta = \ln \phi \).

This minimal model for the transient enhancement of reversals following an initial reversal fits well with the behavior observed in glt-1 mutants, with a time scale \( \tau = 20 \) s, and provides a natural definition of a repetitive behavior index

\[
R_I = \frac{\delta}{\ln 2}.
\]

(14)

An RI of 1 means that, at long times, 50% fewer worms are still moving forward than would be expected according to the baseline reversal rate. Noting that a similar time scale \( \tau \) is found in wild type (although the deviation from a constant reversal rate is smaller), we imposed a fixed value \( \tau = 20 \) s for all lines, to keep a consistent definition of early vs. late reversal rates, and fit \( r_0 \) and \( \delta \) independently for each experiment.

**Reversal response to mechanical stimuli.** To determine the response to mechanical touch, the first-day adult animals were placed individually on 6 cm NGM-agar plates, freshly covered with a thin bacterial layer. Touch was applied to the anterior part of animals that are either in forward motion or paused states, using a fine glass capillary with an angle of 30°, for a total of six injections, at least eight times, with at least 15-min interval between consecutive stimuli. Animals were scored blindly based to three criteria: no response, or short pausing (pass); reversal response composed of backward bout of about 2–3 body bends (Supplementary Movie 5); animals responding with multiple reversal events that are separated by a short pause or forward bouts (repetitive). A worm was considered to have responded if it had more than one body bend. A total of 8, as measured by a Bioanalyzer (Agilent). Specificity of the RNA was determined by using –7% of purified RNA for first strand cDNA synthesis with SuperScript III kit (Invitrogen), followed by qPCR (LightCycler 480, Roche) analysis, using lnm-1 as a control to standardized the samples, and hli-17 to measure differential expression between samples derived from the CEPch glia to the control. All of the data were performed for control samples by the Rockefeller Genomics Resource Center. Briefly, mRNA amplification and cDNA preparation were performed using the SMARTer mRNA amplification kit (Clontech). Labeled samples were sequenced using an Illumina HiSeq 2000 sequencer using either 50- or 100- base read protocols.

**RNA-Seq quality and differential expression (DE) analyses.** Fastq files were generated with CASAVA v1.8.2 (illumina), and examined using the FASTQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) application for sequence quality. Reads were aligned to customized build genome that combine C. elegans WS244 reference genome (ftp://ftp.wormbase.org/pub/wormbase/releases/ WS244/species/c_elegans/PRJNA137586/c_elegans.PRJNA137586.WS244. genomics.fa.gz) and GFP using the STAR v2.3 aligner with parameters55 (--out-filterMultimapScoreRange 1). Mapping rate was >99% with >49 million uniquely mapped reads. The alignment results were evaluated through RNA-SeQC v1.17 to make sure all samples had a consistent alignment rate and no obvious 5’ or 3’ bias56. Aligned reads were summarized through featureCounts57 with gene models from Ensemble (Caenorhabditis_elegans. WBC26-37.gi) at gene level unstranded option, with uniquely mapped reads (NH “tag” in bam file) that overlapped with an exon (feature) by at least 1 bp on either strand were counted and then the counts of all exons annotated to an Ensemble gene (meta feature) were summed into a single number. rRNA genes, mitochondrial genes and genes with length <40 bp were excluded from downstream analysis. Experiment was done with three independent replicates. DESeq2 was applied to normalize count matrix and to perform differential gene-expression analysis, comparing RNA counts derived from the CEPch cells (GFP positive) to RNA counts that were derived from all other C. elegans cells, using negative binomial distribution58.

**Gene-expression comparison between C. elegans and mouse cells.** For gene-expression comparisons between C. elegans and mouse, RNA-sequencing data for the various mouse brain cells were downloaded from GEO (GSE52564), and fold-enrichment (FE) values were calculated as previously described16. RNA-sequencing data of oligodendrocyte precursor cells (OPC), newly formed oligodendrocytes (GC) and myelinating oligodendrocytes (MOG) were merged and averaged to generate a list of oligodendrocyte genes. A comparison between C. elegans and mouse genes was downloaded from PANTHER (ftp://ftp.pantherdb.org/ortholog/14.0/), including both least diverged orthologs (LDO, 5117 records) and all other orthologs (O, 16,487 records). To find unique identifiers for mouse genes, gene symbols from the RNA-sequencing data were converted into Mouse Genome Informatics (MGI) ID numbers to match the data of PANTHER. WormBase gene ID number was used as a unique identifier for C. elegans genes. To perform heatmap hierarchical clustering of mouse brain cells and CEPch glia-expressed genes, mouse genes were converted into their corresponding C. elegans orthologs based on the PANTHER database. Next, FE values in each list were converted to rank order. Gene ranking within each column was used as the input matrix and the heatmap was generated through R (https://cran.r-project.org/web/packages/heatmapplus) using correlation as the distance function. To test for cell type-specific enrichment of the various mouse glia gene lists in the CEPch glia-list, we used step-wise fold-enrichment (FE) cutoff values (for CEPch-glia, log FE = 1.1, 1.5, 2.5, 3.5, 4). To achieve gene-expression specificity for the various mouse glia, we only 5 μM filter to remove undigested animal fragments, and immediately sorted by FACS.
considered genes that are uniquely present in a cell type for any given fold change cutoff value. Each C. elegans gene was counted as related to a mouse glial cell type if the C. elegans gene had any ortholog present on the list of enriched genes from the mouse brain RNA-seq data. For each fold change cut-off pair, we calculated the enrichment score. Specifically, we compared the frequency of CEPsh glia genes with mouse brain-enriched orthologs to the frequency of C. elegans genes in the genome with mouse brain-enriched orthologs, normalized by the number of genes in each list with a mouse ortholog.

\[
\text{Enrichment score} = \frac{\text{Proportion of CEPsh glia enriched genes with a mouse ortholog}}{\text{Proportion of worm genes with a mouse ortholog}}.
\]

(16)

Proportion of CEPsh glia enriched genes with a mouse ortholog = \frac{6 \text{ of CEPsh glia enriched genes with a mouse brain enriched ortholog}}{\# of CEPsh enriched genes with a mouse ortholog};

(17)

Proportion of worm genes with a mouse ortholog = \frac{2 \text{ of worm genes with a mouse brain enriched ortholog}}{\# of worm genes with a mouse ortholog}.

(18)

A \chi^2 test between the various mouse brain cells at the various FE cutoff values was used to correct a potential bias ortholog-mapping may have introduced, and the ratio between the expected value of the enrichment score (given there is no cell type specificity) and the observed enrichment score was calculated as the association score (see Supplementary Data 1).

Statistics. Unpaired two-sided Student’s t test was used to determine the statistical significance between two groups, or between several samples compared to a control group with Bonferroni correction for multiple comparisons. For multiple comparisons ANOVA Tukey’s HSD post hoc test was used, and in cases where the variance between samples were too high (Brown Forsythe test, \(F\) values), Fisher’s exact test was used.

\[ \Delta \text{variance between samples were too high} < \Delta \text{variance between samples were high}. \]

Unpaired two-sided Students t-test was used to determine the statistical significance between two groups, or between several samples compared to a control.

The RNA-seq data sets generated during the current study have been deposited in the European Nucleotide Archive (ENA) with the primary accession code PRJEB31134. The RNA-seq data sets generated during the current study have been deposited in the European Nucleotide Archive (ENA) with the primary accession code PRJEB31134.

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