Excitatory neurons sculpt GABAergic neuronal connectivity in the *C. elegans* motor circuit

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Keywords: GABA synapse, neural circuit, neural development, E/I balance
Summary

Establishing and maintaining the appropriate number of GABA synapses is key for balancing excitation and inhibition in the nervous system, though we have only a limited understanding of the mechanisms controlling GABA circuit connectivity. Here, we show that disrupting cholinergic innervation of GABAergic neurons in the *C. elegans* motor circuit alters GABAergic neuron synaptic connectivity. These changes are accompanied by a reduced frequency and increased amplitude of GABAergic synaptic events. Acute genetic disruption in early development—during the integration of post-embryonic born GABAergic neurons into the circuit—produces irreversible effects on GABAergic synaptic connectivity that mimic those produced by chronic manipulations. In contrast, acute genetic disruption of cholinergic signaling in the adult circuit does not reproduce these effects. Our findings reveal that GABAergic signaling is regulated by cholinergic neuronal activity, likely through distinct mechanisms in the developing and mature nervous system.
Introduction

Neurons in the brain are organized into circuits, and the activity of these circuits is regulated through the integration of excitatory and inhibitory synaptic inputs. Thus, appropriate numbers and proper positioning of excitatory and inhibitory synaptic connections are key factors governing circuit performance. Indeed, disruptions in the balance of excitatory and inhibitory synaptic activity are associated with debilitating brain diseases. For instance, impaired maturation of inhibitory signaling can produce circuit hyperexcitability and a predisposition to seizure activity that is thought to be important in epilepsy (Cossart et al., 2001; Powell et al., 2003; Briggs and Galanopoulou, 2011).

Though extensive studies of excitatory synapses have produced advances in our understanding of their development and regulation, far less is known about the processes that govern inhibitory synaptogenesis and patterning. Gamma-aminobutyric acid (GABA) is the principal inhibitory neurotransmitter in the mature mammalian brain. GABAergic inhibition controls spatiotemporal patterns of activity throughout many brain areas, and robust local GABAergic innervation of excitatory neurons is central for this role. Recent work in mammals has suggested that inhibitory connectivity is shaped by sensory experience, and identified activity-dependent transcriptional pathways important for this process (Morales et al., 2002; Chattopadhyaya et al., 2004; Jiao et al., 2006; Bloodgood et al., 2013; Pieraut et al., 2014). Despite these important advances, many aspects of inhibitory network development remain inadequately understood.

C. elegans provides a useful model in which to study neural circuit development and GABAergic function in vivo. Like mammals, inhibitory GABAergic signaling shapes C. elegans neural circuit activity and behavior. The identity, location and connectivity of all GABAergic neurons have been defined experimentally, and a wealth of genetic tools is available for manipulating GABAergic activity (Schuske et al., 2004). Informative genetic studies have previously identified genes involved in GABAergic synapse formation in C. elegans, however the extent to which these genes identify mechanisms specific to GABAergic synapses versus more generalized roles remains unclear (Zhen and Jin, 1999;
Schaefer et al., 2000; Zhen et al., 2000; Hallam et al., 2002; Liao et al., 2004; Ackley et al., 2005; Dai et al., 2006; Grill et al., 2007; Najarro et al., 2012). Here we investigate the role of neuronal activity in the establishment and maintenance of inhibitory synaptic connections using the *C. elegans* motor circuit as a model. Prior work has provided evidence that activity-dependent processes regulate the timing of GABAergic developmental remodeling in *C. elegans* (Thompson-Peer et al., 2012; Han et al., 2015). Our studies here provide some of the first evidence that activity-dependent processes help to shape the mature pattern of synaptic connectivity in post-embryonic born GABAergic neurons.
Results

**UNC-3 transcriptional regulation coordinates cholinergic synaptic connectivity with body wall muscles**

The adult pattern of synaptic connectivity in the motor circuit has been well defined. Body wall muscles receive synaptic inputs from both excitatory (ACh) and inhibitory (GABA) motor neurons and these inputs shape muscle activity and sinusoidal movement (Figure 1A). Cholinergic motor neurons also provide excitatory synaptic drive onto inhibitory GABA motor neurons. GABA motor neurons then make inhibitory synaptic connections onto opposing musculature. The ventral D (VD) class GABAergic motor neurons that are the focus of our studies are integrated into the motor circuit post-embryonically following the first larval stage, and mediate inhibitory transmission onto ventral muscles in adults (Sulston, 1976; Han et al., 2015).

To address whether excitatory motor neurons are important for shaping functional connectivity in this circuit, we first examined *unc-3* mutants. *unc-3* encodes the sole *C. elegans* homolog of the COE family of transcription factors and, in *unc-3* mutants, most ventral cord motor neurons completely lack expression of genes required for cholinergic function (e.g. *cha-1* and *unc-17*) and exhibit additional defects in their organization (e.g. variable and/or disorganized process extension)(Prasad et al., 1998; Prasad et al., 2008; Kratsios et al., 2012). Thus, cholinergic transmission onto body wall muscles should be severely disrupted in these animals. To confirm this, we made whole-cell patch clamp electrophysiology recordings of synaptic events from ventral body wall muscles of adult *unc-3* mutants (1 mM Ca²⁺) and found significant defects in excitatory transmission (Figure 1B, C). Specifically, the frequency of endogenous excitatory post-synaptic currents (EPSCs) is significantly reduced compared to wild type (79 ± 6% reduction, p<.001). The remaining cholinergic transmission may reflect activity of cholinergic VC motor neurons, which do not require *unc-3* for expression of cholinergic markers (Kratsios et al., 2012). Prior work identified a requirement for *unc-3* in the clustering of postsynaptic ionotropic (nicotinic) acetylcholine receptors (iAChR) in muscles (Kratsios et al., 2015). Confirming these
observations, we find that the distribution of muscle iAChRs (visualized with ACR-16::GFP) is disrupted by mutation of unc-3 (Figure 1D, E). While the total number of ACR-16::GFP clusters is only modestly reduced compared with wild type, large accumulations of receptor are often present in the shafts of muscle arms in unc-3 mutants, suggesting a redistribution to non synaptic locations. Thus, impaired terminal differentiation of cholinergic motor neurons leads to predicted defects in iAChR clustering and excitatory transmission at the cholinergic neuromuscular junction (NMJ).

**Mutation of unc-3 impairs post-synaptic iAChR clustering in GABA neurons and reduces inhibitory transmission**

Given that cholinergic motor neurons also make extensive synaptic contacts with inhibitory motor neurons, we next sought to determine whether the disruption of cholinergic innervation associated with mutation of unc-3 affected inhibitory activity. Using conditions that isolated inhibitory transmission, we measured endogenous inhibitory synaptic currents (IPSCs) from unc-3 mutants. The rate of endogenous IPSCs (5 mM Ca\(^{2+}\)) is significantly lower in unc-3 mutants compared to wild type (63% reduction, p<0.0001) (Figure 2A,B). Interestingly, mutation of unc-3 also produces a shift toward larger amplitude endogenous IPSCs (Figure 2C), suggesting either altered muscle responsiveness to GABA release or increased GABA content in synaptic vesicles. Our analysis supports the notion that cholinergic neurons provide direct excitatory drive onto GABAergic motor neurons and perhaps also regulate functional attributes of GABA synaptic outputs onto muscles.

Given these findings, we next asked whether unc-3 is required for establishing proper synaptic connectivity between cholinergic and GABAergic motor neurons. The nAChR subunit ACR-12 is a constituent of GABA neuron iAChRs. ACR-12::GFP clusters localize opposite ACh release sites and relocate appropriately during developmental remodeling, suggesting these clusters report mature post-synaptic structures (Petrash et al., 2013; He et al., 2015). We examined the distribution of ACR-12::GFP in the dorsal nerve cord where the majority of chemical synaptic inputs to VD neurons occur (VD synaptic contacts with
muscles are exclusively ventral) (White et al., 1986). We found that receptor clustering is severely disrupted in two independent unc-3 mutant strains (n3435 and e151) (Figure 2D, E). ACR-12 clusters are unevenly distributed and significantly reduced in number compared to wild type (e.g. 51% decrease in unc-3(e151), p<0.001). These effects are rescued with expression of the wild type unc-3 cDNA under control of a 1kb unc-3 promoter, but not by either GABA or muscle-specific expression (Figure 2D, E). This unc-3 promoter region drives expression in cholinergic, but not GABAergic, motor neurons (Figure S1A). Together, these observations suggest that unc-3 expression in cholinergic neurons is required for proper ACR-12 clustering in postsynaptic GABAergic neurons.

madd-4/Punctin encodes a synapse-associated extracellular scaffolding protein (Pinan-Lucarre et al., 2014). Secretion of MADD-4 from cholinergic motor neurons is required for proper iAChR localization at neuromuscular synapses (Pinan-Lucarre et al., 2014). Prior work showed that UNC-3 transcriptional regulation of madd-4/Punctin is essential for this process (Kratsios et al., 2015). To test whether MADD-4 is similarly required at synaptic connections between cholinergic and GABAergic motor neurons, we examined ACR-12 clustering in GABAergic neurons of madd-4/Punctin mutants. The madd-4(ok2854) allele is a 962 bp deletion that eliminates the two largest madd-4 exons common to all madd-4 isoforms, has been previously reported to be a null (Seetharaman et al., 2011; Maro et al., 2015), and produces significant defects in muscle AChR clustering similar to those previously observed for deletion of the entire madd-4 locus (Pinan-Lucarre et al., 2014)(Figure S1B, C). Mutation of madd-4/Punctin produces a slight decrease in the number of ACR-12 clusters in the dorsal nerve cord compared to wild type (18% decrease, p<0.05)(Figure 2D, E). Notably, this effect is much less severe than observed for mutation of unc-3, indicating that the post-synaptic defects in GABAergic neurons of unc-3 mutants cannot be accounted for by decreased madd-4 expression. Taken together, our results indicate that UNC-3 transcriptional control of cholinergic neuron development is essential for proper iAChR clustering in GABAergic motor neurons, though MADD-4 has a comparatively minor role.
Morphological development of body wall muscles and GABA motor neurons is not dramatically affected in *unc-3* mutants

To address whether the connectivity changes we observed might reflect alterations in the gross development or assembly of the motor circuit, we examined the cellular morphology of body wall muscles and GABA motor neurons. In wild type animals, body wall muscles project membrane extensions, called muscle arms, to the nerve cords where muscle arm termini bifurcate and project lengthwise along the nerve cord, forming *en passant* synapses with motor neurons (Dixon and Roy, 2005). We visualized muscles by specific expression of membrane bound mCD8::GFP (*him-4p::mCD8::GFP*) (Dixon and Roy, 2005; Collins and Koelle, 2013) (Figure 3A). Variable minor morphological defects (e.g. overgrowth, membrane blebbing) are apparent in the body wall muscles of *unc-3* mutants (Figure 3B). However, consistent with a prior study (Kratsios et al., 2015), we found that overall muscle morphology is preserved—muscle arms extend normally to the nerve cord and the extent of muscle membrane contact with the nerve cord region is not appreciably altered (Figure 3D).

We visualized the morphology of GABA motor neurons using cell-specific expression of mCherry (*unc-47p::mCherry*) (Figure 3A1). Similar to the case for muscle cells, disruption of cholinergic innervation does not produce significant defects in GABA motor neuron number, positioning or gross morphology (Figures 3B1 and S2A-C). Although in some cases we noted regions of nerve cord defasciculation in *unc-3* mutants (and sometimes one fewer commissure was visible), we did not observe disruptions in the continuity of the dorsal nerve cord processes. Thus, our findings provide evidence that gross morphological development of body wall muscles and GABA motor neurons proceeds to completion when cholinergic development is disrupted, arguing that the iAChR clustering and GABA transmission defects we observed in *unc-3* mutants are unlikely to arise secondary to impaired morphological development of these cell types.
Cholinergic innervation refines GABA synaptic outputs

Our above results support the idea that UNC-3 regulation of cholinergic neuron differentiation is important for the establishment of synaptic connectivity with GABAergic neurons. We next asked whether cholinergic neurons may similarly contribute to the development of GABA synaptic outputs onto muscles. We examined GABA synapses in a transgenic strain expressing the synaptic vesicle marker mCherry::RAB-3 in GABA motor neurons together with the GFP-tagged GABA$_A$-like receptor UNC-49 in muscles (Bamber et al., 1999; Klassen and Shen, 2007)(Figure 4A). In wild type animals, GABA synapses are evenly spaced along the ventral nerve cord, with presynaptic vesicles and post-synaptic GABA receptor clusters in close apposition (Figure 4B and S3A, D). Mutation of unc-3 produces striking changes in GABA synapses (Figure 4C-D, G-H), a phenotype consistent across independent unc-3 mutant strains (n3435 and e151). Specifically, the number of pre- and post-synaptic clusters is significantly decreased, while the average size of postsynaptic UNC-49::GFP puncta is increased (e.g. by 95% in unc-3(e151)). Further, we noted significant regions of the ventral nerve cord lacking GABA synaptic vesicle clusters in unc-3 mutants (e.g. 56 ± 4% in unc-3(e151), compared to 20 ± 2% in the wild type, p<0.01). A prior study reported diffusely distributed and enlarged clusters of the active zone marker SYD-2/α-Liprin in GABA neurons of unc-3 mutants (Yeh et al., 2005), consistent with our findings.

The localization of other synaptic markers, including the synaptic vesicle marker SNB-1/synaptobrevin (Nonet, 1999) and the active zone marker UNC-10/RIM, are similarly altered by mutation of unc-3 (Figure 4I-L). The size and distribution of SNB-1 clusters are both affected similarly to RAB-3. The number of UNC-10 clusters is decreased significantly but we do not observe a change in their average area, suggesting that synaptic vesicle clustering may be more dramatically affected than the size of the active zone itself. Despite the altered distribution of synaptic markers, we did not observe obvious changes in the apposition of presynaptic mCherry::RAB-3 and postsynaptic UNC-49::GFP fluorescent...
signals (Figure S3D) (wild type: 88 ± 3%, unc-3(e151): 82 ± 4% in apposition), suggesting that trans-synaptic signaling important for establishing proper registration between pre- and post-synaptic specializations is maintained.

Most aspects of these GABA synapse defects are reversed with expression of the wild type unc-3 cDNA in cholinergic neurons (Figure 4E, G-H). In particular, the decreased number of GABAergic presynaptic RAB-3 clusters and the increases in synapse size are each rescued with unc-3p::unc-3 expression, but not by GABA-specific expression. While decreases in the number of GABAR clusters are not significantly rescued by unc-3p::unc-3 expression, GABA- or muscle-specific expression enhances the severity of this phenotype, arguing against a requirement for unc-3 expression in these cells. Finally, we noted that muscle-specific expression of unc-3 appears to partially reverse the increases in synapse size; however, decreases in synapse number are not rescued. Together, our transgenic rescue experiments provide additional support for the idea that unc-3 expression in cholinergic neurons is required for establishing proper GABAergic connectivity.

Surprisingly, unc-3p::unc-3 cDNA expression is not sufficient to restore movement to unc-3 mutants. In contrast, expression of an unc-3 minigene containing the first three unc-3 introns under control of the same promoter improves movement and restores synaptic connectivity (Figure S4), suggesting that additional aspects of unc-3 regulation are important for motor function and that unc-3 intronic regions are critical in this process.

To further test the idea that cholinergic innervation influences the number and positioning of GABAergic synapses, we ablated ACh motor neurons by expressing a putative gain-of-function variant of the acetylcholine receptor subunit acr-2, acr-2(L/S), that produces cell-autonomous death of cholinergic motor neurons during late embryogenesis (Barbagallo et al., 2010). Similar to unc-3, we did not observe gross morphological changes in GABA neurons or reduced muscle membrane contact with the nerve cord (Barbagallo et al., 2010) (Figure 3C, D). Nonetheless, genetic ablation of ACh motor neurons produces striking defects in inhibitory synapses, similar to those observed for unc-3 mutants, including a significant decrease in the number of both presynaptic mCherry::RAB-3 and postsynaptic
UNC-49::GFP clusters, as well as significant increases in their size (Figure 4F-H, S3C).

Thus, impaired differentiation or genetic ablation of ACh motor neurons does not affect the gross morphology of GABA neurons but instead alters the density and size of GABA synaptic outputs. Our analysis of inhibitory synapses in unc-3 mutants and acr-2(L/S) transgenic animals supports the hypothesis that presynaptic cholinergic neurons play important roles in both clustering iAChRs in GABA neurons and in shaping GABA synaptic outputs.

Vesicular release from cholinergic neurons shapes GABA synapses

We next asked whether vesicular release from cholinergic neurons is required to properly establish GABA neuron connectivity. We disrupted vesicular release by specific expression of tetanus toxin light chain (TetTx) (Figure 5 and S5). TetTx expression in cholinergic motor neurons does not affect morphology or outgrowth, but produces changes in sensitivity to the cholinesterase inhibitor aldicarb consistent with decreased transmission (Figure S5). Cholinergic expression of TetTx does not alter ACR-12 clustering in GABA neurons (Figure 5A, B). We did, however, observe significant abnormalities in GABA synaptic outputs (Figure 5C-F). These include enlargement of both presynaptic mCherry::RAB-3 and postsynaptic receptor clusters, as well as gaps in fluorescence that are similar to those observed for unc-3 mutants and acr-2(L/S) transgenic animals. Thus, synaptic structures between cholinergic and GABAergic neurons appear properly established when vesicular release from cholinergic motor neurons is impaired. Despite this, the distribution of GABAergic synaptic outputs is disrupted, implicating vesicular signals from ACh motor neurons in contributing to this process.

To investigate the possibility that ACh itself is an important signal, we examined GABA synapses in mutant strains with specific defects in cholinergic transmission. unc-17/vAChT mutants are defective in a vesicular transporter (vAChT) that loads ACh into synaptic vesicles (Brenner, 1974; Rand and Russell, 1984; Rand, 1989). The unc-17(e113) mutation disrupts the UNC-3 binding site in the unc-17 promoter, disrupting UNC-17 and CHA-1
expression in cholinergic motor neurons (J. Rand, personal communication), and these mutants are presumably null for ACh release from cholinergic motor neurons.

ACR-12::GFP clustering in the dorsal nerve cord is not significantly altered in unc-17(e113) mutants or in madd-4;unc-17 double mutants (Figure 6A, B), consistent with our above results for TetTx expression. In contrast, GABAergic pre- and post-synaptic clusters are significantly decreased in number in unc-17(e113) mutants (by 65% and 64% respectively, p<0.001) (Figure 6D, G). A second strongly hypomorphic unc-17 allele (e245) produced similar effects. The e245 mutation replaces a glycine with an arginine (G347R) in the ninth transmembrane domain of the vAChT protein (Alfonso et al., 1993) and decreases UNC-17/vAChT protein levels (Sandoval et al., 2006). ACR-12::GFP clustering is unchanged in unc-17(e245) compared to wild type (Figure 6B), but significant defects in GABA synaptic outputs are apparent. These include gaps in the distribution of synapses as well as enlarged pre- and post-synaptic clusters (Figure 6E, G-H). The increased size of GABAergic synapses in unc-17(e245) mutants, but not unc-17(e113) mutants, may suggest that partial loss of ACh release triggers changes in GABA synapse size. Alternatively, ACh release from cholinergic neurons in which unc-17 transcriptional regulation occurs independently of unc-3 (e.g. VC neurons) may influence GABA synapse size. Together, our analysis of unc-17 mutants provides evidence for the involvement of cholinergic transmission in directing the development of GABA synaptic outputs. Nonetheless, these effects are less pronounced than observed for either TetTx expression or mutation of unc-3, suggesting participation of additional ACh-independent signals.

To address whether the effects of decreased ACh release reflect a requirement for ACh signaling onto muscles, we examined GABA synapses in animals lacking functional iAChRs at the NMJ (unc-29;acr-16 double mutants)(Francis et al., 2005). Eliminating synaptic activation of muscles causes a slight reduction in the number of postsynaptic GABA receptor clusters (Figure 6F-H) but does not alter the density or positioning of presynaptic structures, suggesting that cholinergic activation of muscles is not essential for shaping GABAergic neuromuscular connectivity.
Acute reductions in cholinergic activity in adults lead to decreased GABA synapse size

Our previous genetic analysis indicated that cholinergic transmission helps to shape the distribution of GABA synaptic outputs, but did not distinguish whether these effects occur in the mature or developing nervous system. To address this question, we used a temperature sensitive allele (y226) of cha-1/choline acetyltransferase (ChAT) (Rand and Russell, 1984). When maintained at a permissive temperature (15°C), cha-1(y226)/ChAT mutants develop similarly to wild type, generating the full complement of cholinergic motor neurons (Figure S6A). When switched to a restrictive temperature (25°C), cha-1(y226)/ChAT mutants become uncoordinated within minutes, consistent with rapid impairment of cholinergic transmission, and these behavioral effects are reversible upon shifting back to the permissive temperature ((Zhang et al., 2008) and Figure S6B).

We first investigated the effects of reduced cholinergic transmission in adult animals. We shifted cha-1(y226)/ChAT mutants to 25°C for a 4 hr period in early adulthood, and visualized GABA synapses immediately following this shift (Figure 7A). Interestingly, this transient reduction in cholinergic transmission does not produce appreciable changes in the distribution or number of GABA synapses. Instead, we noted a significant decrease in the size of both pre- and post-synaptic clusters (Figure 7B-G), while GABA synapses in wild type animals remain unaffected. This temperature shift paradigm also produces a significant decrease in movement velocity in cha-1(y226) mutants, and these effects are reversible within a 4 hr recovery period at 15°C (Figure S6B). GABA synaptic clusters in cha-1 mutants are also restored to their normal size following a 4 hr recovery period (Figure 6D-G).

Increasing the duration of shifts to the restrictive temperature (8 hr) produces further decreases in the size of GABA pre- and post-synaptic clusters. Our findings indicate that acute changes in cholinergic activity during adulthood are sufficient to alter the size of GABA synapses, without producing obvious effects on GABA synapse positioning or density. The reversible nature of the synaptic alterations in adults provides evidence for active
mechanisms that regulate the size of GABA synapses in response to changing levels of cholinergic transmission.

**Acetylcholine release is required during a highly synaptogenic period for proper GABA synapse localization**

We next asked whether cholinergic activity during development might help to shape the density or distribution of GABA synapses. To address this question we again used *cha-1(y226)/ChAT* mutants, shifting animals to 25°C for 8 hr periods during the first larval (L1) stage, then imaging GABA synapses later in adulthood (Figure 8A). We noted a time window beginning approximately 20 hr after hatch (late L1 stage in wild type at 15°C) during which an 8 hr shift to the restrictive temperature produces significant defects by adulthood. Specifically, we found the number of pre- and post-synaptic clusters is decreased by 40% and 27% respectively in *cha-1* mutants subjected to temperature shift (Figure 8B-D). We also noted a significant increase in the size of presynaptic clusters (Figure 8E), although the size of postsynaptic clusters does not appear appreciably altered (Figure 8F). These effects do not occur in control animals exposed to the same shift in temperature, or in *cha-1* mutants grown continuously at 15°C. This temperature shift paradigm also produces a modest decrease in movement velocity in *cha-1(y226)* adult animals, suggesting that transient reduction of cholinergic transmission during development may be sufficient to produce behavioral defects that extend into adulthood (Figure S6C). The timing of the 8 hr temperature shift is predicted to overlap the L1/L2 transition for wild type animals during which GABAergic VD neurons are initially integrated into the motor circuit, although the precise timing of these events can vary in mutant and transgenic strains. Thus, reduced cholinergic activity early in the development of GABAergic VD neurons affects their synaptic distribution, implicating cholinergic signaling in defining the mature pattern of inhibitory connectivity.
Discussion

Establishing the appropriate pattern of GABAergic innervation is critical for proper inhibitory regulation of mature neural circuits. In this study, we explored the role of cholinergic motor neurons in sculpting the synaptic connectivity of GABAergic neurons in the *C. elegans* motor circuit. Our results show that integration of post-embryonic born GABAergic VD neurons into the motor circuit is extensively regulated by cholinergic motor neurons, complementing prior studies that provided evidence for activity-dependent synaptic remodeling of embryonic-born GABAergic DD neurons (Thompson-Peer et al., 2012; Han et al., 2015). First, mutation of the COE transcription factor *unc-3* that is required for terminal differentiation of cholinergic motor neurons disrupts post-synaptic development (iAChR clustering) in GABAergic neurons. These effects are not strongly dependent on either cholinergic vesicular release or expression of the synaptic organizer *madd-4*/Punctin, suggesting involvement of a novel molecular pathway. Second, mutation of *unc-3* produces defects in GABAergic VD neuron synaptic connectivity with muscles. Third, genetic ablation of cholinergic motor neurons or specific TetTx expression in cholinergic neurons similarly alters the size and distribution of GABA synaptic outputs to muscles. Mutation of the cholinergic vesicular transporter *unc-17* produces similar effects on GABAergic connectivity with muscles. Finally, the density of GABAergic synaptic outputs is decreased by acute reduction of cholinergic transmission during a developmental period that corresponds with the time of initial integration of GABAergic VD neurons into the circuit in wild type. Our findings suggest a model where vesicle-borne signals from cholinergic motor neurons, both ACh and additional signals, are important for shaping GABAergic connectivity.

Cholinergic motor neurons innervate both GABAergic motor neurons and body wall muscles. Disruption of cholinergic innervation did not impair gross morphological development or outgrowth in either of these cell types, indicating that cell intrinsic genetic programs are sufficient to guide normal morphological development. In contrast, additional activity-dependent processes appear to be required for establishing the mature pattern of GABAergic synaptic connectivity. These results may be indicative of a functional segregation...
between processes underlying activity-dependent regulation of synaptic connectivity and intrinsic genetic regulation of gross morphological features. Prior work showed that elimination of glutamatergic synaptic inputs has little effect on the morphological development of mouse hippocampal neurons (Lu et al., 2013), providing support for a similar segregation between these processes in mammals.

Mutation of unc-3 produced the most striking effects on synaptic connectivity in our experiments, both impairing post-synaptic ACR-12 iAChR clustering in VD neurons and decreasing their synaptic outputs. The requirement for UNC-3 in ACR-12 iAChR clustering is unlikely to indicate involvement of cholinergic activity-dependent regulation because neither TetTx expression nor mutation of unc-17/vAChT altered ACR-12 clustering. Similarly, mutation of the synaptic organizer madd-4/Punctin did not strongly affect ACR-12 iAChR clustering in VD neurons, either alone or in combination with mutation of unc-17/vAChT, suggesting that a novel as yet unidentified transcriptional target of UNC-3 may be important for this process.

Our studies suggest that cholinergic activity may have a more prominent role in directing the distribution of GABAergic synaptic outputs. Our electrophysiological analysis of unc-3 mutants showed that GABAergic VD motor neurons are capable of sustaining inhibitory transmission. Notably however, the frequency of GABAergic synaptic events is dramatically reduced compared with wild type, and their average amplitude increased significantly, mirroring the changes we observed in GABAergic outputs using our synaptic markers. We noted similar structural changes in GABAergic synaptic outputs when cholinergic neurons were genetically ablated (acr-2(L/S)), and when cholinergic vesicular release was disrupted (ACh::TetTx). For all of these, presynaptic alterations were paralleled by similar effects on the clustering of post-synaptic GABA receptors, suggesting that these effects reflect a change in synapse patterning/maturation rather than direct impairment of synapse formation per se.

Although synapse dynamics continue throughout the life of an animal, there is often increased plasticity earlier in development (Hensch and Fagiolini, 2005). The effects of
cholinergic activity we observed might reflect an ongoing requirement for cholinergic transmission or a more specific requirement during a developmental period of increased plasticity. Our analysis of temperature-sensitive cha-1 mutants provides support for the latter possibility. A temporally defined reduction in cholinergic transmission that overlaps with the predicted timing of the highly synaptogenic L1/L2 transition in wild type produced irreversible defects in the patterning and size of GABAergic synaptic outputs. At the L1/L2 transition, thirteen GABAergic VD neurons are added to the motor circuit, forming synapses onto ventral muscles. Our findings suggest that cholinergic activity during this developmental period influences the patterning of GABA synapses, and deficits in transmission during this time period cannot be compensated for in later developmental stages. How might reduced cholinergic activity in early development alter GABAergic synaptic outputs in the adult? Synaptic inputs to the VDs primarily come from the dorsal A and B (DA/DB) classes of cholinergic motor neurons that are present from hatch, and are rewired to innervate VD neurons after the L1 stage. One interesting possibility is that a reduction in cholinergic activity impairs rewiring and DA/DB innervation of VD GABAergic neurons. According to this model, defects in GABA synapse patterning would then arise due to impaired DA/DB activation of GABAergic neurons. In contrast, acute reduction of cholinergic activity in the adult produces effects that are distinct from those during development, including changes in synapse size without decreases in synapse number, raising the possibility that homeostatic mechanisms are established after synapse patterning is completed. Similar scaling mechanisms have been reported at excitatory glutamatergic synapses in worms, flies and mammals (Grunwald et al., 2004)(Turrigiano, 2008; Davis, 2013).

GABA release itself has been suggested to be key for proper inhibitory synapse maturation in mammals (Chattopadhyaya, 2011). Notably, disruption of GABA biosynthesis (unc-25/GAD mutants) does not impair GABA synapse formation in worms (Jin et al., 1999; Gally and Bessereau, 2003). Likewise, mutations that produce global reductions in neurotransmission (e.g. mutations in unc-13/Munc13 or unc-64/Syntaxin) do not dramatically alter inhibitory synaptogenesis (Saifee et al., 1998; Richmond et al., 1999). For our studies
we used genetic tools that specifically affect cholinergic neurons. Under these conditions, inhibitory synapse formation appears properly initiated. Pre-synaptic specializations and post-synaptic receptor clusters are in close apposition, however the size and density of these synapses is altered, implicating vesicular signals from cholinergic neurons in the refinement or patterning of GABAergic synaptic outputs. While the mechanism underlying this process remains to be determined, one possibility is that vesicular signals from cholinergic neurons may regulate expression of synapse organizing molecules in GABA neurons. Several recent studies have identified genetic pathways important for GABA synapse development and regulation in worms (Maro et al., 2015; Tong et al., 2015; Tu et al., 2015). It will be interesting to investigate whether cholinergic neurons influence expression levels of these and other potential synaptic organizers. Notably, activity-dependent transcriptional cascades shape the development and placement of GABA synapses in mammals (Lin et al., 2008; Bloodgood et al., 2013). Our findings lead us to propose that similar processes may be important for integrating post-embryonic born GABAergic VD neurons into the motor circuit and patterning their synaptic outputs.
Materials and Methods

Strains

*C. elegans* strains were grown at room temperature (22-24°C) on nematode growth media (NGM) plates seeded with the *Escherichia coli* strain OP50 unless otherwise noted. Wild type represents the N2 Bristol strain. Transgenic strains were obtained by microinjection to achieve germline transformation (Mello et al., 1991). Multiple independent lines were obtained for each transgenic strain, and data presented are from a single representative transgenic line. Transgenic strains were identified either by co-injection of *lin-15(+) (pL15Ek)* into *lin-15(n765ts)* mutants or by co-injection of either *lgc-11p::GFP* or *lgc-11p::mCherry* (restricted expression to the pharynx). As required, transgenes were integrated using x-ray bombardment and outcrossed to wild type. A complete list of strains is available in the supplemental information (Table S1).

Molecular Biology

Plasmids were constructed using the two-slot Gateway Cloning system (Invitrogen) as described previously (Bhattacharya et al 2014).

him-4p::mCD8::GFP: Entry Vector: A *him-4p* promoter fragment (2156 bp) was amplified from pPRZ138.2 (a gift from Peter Roy) and cloned into pENTR-D-TOPO to create pENTR-3'-him4. Destination Vector: mCD8::GFP (1566 bp) was amplified from pKMC72 and ligated into pDEST-16 to create pDEST-20. pENTR-3'-him4 and pDEST-16 were recombined to generate pBB102 (*him-4p::mCD8::GFP*).

Tetanus toxin construct: Entry Vector: An *acr-2* promoter fragment (3386 bp) was amplified from pBB67 and subcloned into pENTR-D-TOPO to create pENTR-5'-acr2. Destination Vector: Tetanus Toxin Light Chain (1374 bp) was amplified from TetTx-pWD157 to create pDEST-34. pENTR-5'-acr2 and pDEST-34 were recombined to generate pBB112 (*acr-2p::TetTx*).

unc-3 reporter and rescue constructs: Transcriptional reporter: An *unc-3* promoter fragment (-1068 bp relative to start) was amplified from genomic DNA and cloned into pENTR-D-TOPO to generate pENTR-17. pENTR-17 was recombined with pDEST-94 to generate *unc-
3p::GFP (pAP192). cDNA rescue: Wild type unc-3 cDNA was RT-PCR amplified from total N2 RNA and inserted into a destination vector to create pDEST-140. pDEST-140 was recombined with pENTR-17 (unc-3 promoter), pENTR-3′-myo3 (muscle-specific promoter), and pENTR-3′-unc47 (GABA-specific promoter) to generate pAP194 (unc-3p::unc-3cDNA), pAP172 (myo-3p::unc-3cDNA), and pAP176 (unc-47p::unc-3cDNA), respectively. Minigene rescue: A 3154 bp fragment extending from the unc-3 start to exon 4 of the unc-3 genomic locus was amplified from N2 genomic DNA and ligated into pDEST-140 to create pDEST-142. pDEST-142 was recombined with pENTR-17 to create pAP196 (unc-3p::unc-3 minigene) containing the first three unc-3 introns (a 463 bp repetitive fragment within intron 1 (+2181 relative to start) was not included) and all unc-3 coding sequence.

Electrophysiology

Endogenous postsynaptic currents (PSCs) were recorded from body wall muscles as described previously (Petrash et al., 2013). The extracellular solution consisted of the following (in mM): 150 NaCl, 5 KCl, 4 MgCl₂, 1 or 5 CaCl₂ (as indicated), 15 HEPES, and 10 glucose, pH 7.4, osmolarity adjusted with 20 sucrose. To facilitate analysis of unc-3 mutants that display very low rates of GABAergic activity, recordings of GABAergic currents were performed in 5 mM CaCl₂. The intracellular fluid consisted of the following (in mM): 115 K-gluconate, 25 KCl, 0.1 CaCl₂, 50 HEPES, 5 Mg-ATP, 0.5 Na-GTP, 0.5 cGMP, 0.5 cAMP, and 1 BAPTA, pH 7.4, osmolarity adjusted with 10 sucrose. At least 60–90 s of continuous data were used in the analysis. Data analysis was performed using Igor Pro (Wavemetrics) and Mini Analysis (Synaptosoft) software.
Aldicarb Assay

Aldicarb assays were performed on adult animals (24 hours post L4) at room temperature (22-24 °C) with the researcher blind to genotype. Staged animals (10/plate) were transferred to NGM plates containing 1 mM Aldicarb (ChemService) and assessed for movement every 15 minutes for two hours.

Microscopy and Image analysis

For all imaging, nematodes were immobilized with sodium azide (0.3 M) on a 5% agarose pad. Each n represents analysis of the nerve cord from an independent animal. Confocal microscopy of GABA NMJ synapses was performed using an LSM Pascal 5 confocal microscope (Zeiss). Images were analyzed using ImageJ software (open source). Images were obtained using a 63x oil lens and aligned using Photoshop (Adobe) or ImageJ.

Synapse density was calculated in a 50 µm section of the ventral nerve cord directly posterior to the vulva. A minimum fluorescence threshold was set to identify synaptic peaks (1500 a.u. for cholinergic synapses and 500 a.u. for GABAergic synapses). Peaks were identified as two or more consecutive data points above threshold and spaces were defined as two or more data points below threshold. Synapse area was calculated by separating particles using the ImageJ analyze particles function. Particles ≤ 0.2 µm² were excluded as background.

To quantify orphan puncta at the GABAergic NMJ, the fluorescence intensities of both GFP and mCherry signals in an 85 µm region of the ventral nerve cord were analyzed. Intensity data were exported to Microsoft Excel and normalized to the corresponding highest value. Each peak of fluorescence was categorized as a colocalized puncta, pre-synaptic orphan puncta (mCherry signal without overlapping GFP signal), or post-synaptic orphan puncta (GFP signal without overlapping mCherry signal).

The extent of muscle membrane coverage of the ventral nerve cord region was calculated by measuring the length of the ventral midline of the posterior body wall muscles.
The length of muscle membrane along the ventral midline (visualized by him-4p::mCD8::GFP) was quantified and used to calculate a percentage of total nerve cord length.

ACR-12::GFP (ufls92), ACR-16::GFP (ufls8), and UNC-29::GFP (ufls2) images were acquired using a 3i (Intelligent Imaging Innovations) Everest spinning-disk confocal microscope and collected at 0.27 μm/step using a 63X objective.

Confocal montages of ACR-12::GFP were assembled by imaging the dorsal nerve cord posterior to the vulva in adult hermaphrodites. Identical image and laser settings were used for each genotype. Straightened dorsal cords were extracted from these images using the straighten to line function.

For quantifying ACR-12::GFP, ACR-16::GFP, and UNC-29::GFP puncta along the nerve cord, background fluorescence was first subtracted by calculating the average intensity of each image in a region devoid of puncta. Puncta were quantified across 80 µm of the nerve cord region with ImageJ analyze particles, using an intensity threshold specific for each marker. Particles ≤ 0.04 µm² were excluded as background.

**unc-3 movement analysis**

For locomotion assays, one-day old adults were allowed to acclimate for 1 minute on NGM plates thinly seeded with a bacterial lawn. 30 s digital videos of individual worms were then captured and analyzed using WormLab software (MBF Bioscience).
**cha-1 time course**

**Microscopy**

*Adult Shifts and Recovery:* Wild type and cha-1 animals were cultured at 15°C. Animals were synchronized at the L4 stage, and then placed at 15°C for an additional 24 hours. Young adult animals were then shifted to 25°C for either four or eight hours and imaged immediately. A third group of animals were shifted to 25°C for four hours, then placed back at 15°C for four hours (recovery) before imaging.

*Developmental Shift:* Embryos were staged at the 21+ cell stage and allowed to develop at 15°C for 20 hours. Animals were then shifted to 25°C for eight hours and returned to 15°C. Animals were synchronized at the L4 stage and allowed to grow for an additional 24 hours at 15°C for prior to imaging.

**Behavior**

Temperature shifts were performed as described above. For behavioral analysis, animals were transferred individually to plates without food for a period of five minutes. Movement was quantified using Wormlab software to track animals and measure their average forward velocity over a period of five minutes.
Acknowledgments

We would like to thank Claire Bénard and members of the Francis lab for critical reading of the manuscript, Will Joyce for technical assistance, Mei Zhen, Kevin Collins, Bruce Bamber and Peter Roy for sharing reagents. Some nematode strains used in this work were provided by the Caenorhabditis Genetics Center, which is funded by the NIH National Center for Research Resources (NCRR).
References


Collins, K. M. and Koelle, M. R. (2013). Postsynaptic ERG potassium channels limit muscle...
excitability to allow distinct egg-laying behavior states in *Caenorhabditis elegans*. *J Neurosci* 33, 761-775.


Response to GABA Affects Development and Behavior of the Locomotor Circuit of

_Caenorhabditis elegans._ Genetics


Figure 1. Cholinergic neuromuscular transmission is impaired in *unc-3(e151)* mutants

(A) The *C. elegans* motor circuit. Excitatory cholinergic motor neurons (gray) synapse onto inhibitory GABAergic motor neurons (blue) and onto body wall muscles. (B) Representative recordings of endogenous EPSCs from body wall muscles of wild type and *unc-3(e151)* mutant animals. Red traces are expanded views of time segments under the red bars in the upper traces. (C) The average frequency of endogenous EPSCs recorded from wild type and *unc-3(e151)* mutants. Each bar represents the mean ± SEM, and numbers in bars indicate the n for each genotype in this figure and for all subsequent figures. ***p<0.0001, student’s t-test. (D) Confocal images of one wild type (upper) and two *unc-3(e151)* (lower) animals expressing GFP-tagged AChRs in body wall muscles (*myo3p::ACR-16::GFP*). Scale bar, 5 µm. (E) Normalized receptor cluster number/75 µm of ventral nerve cord. *p<0.05, student’s t-test.
Figure 2. *unc-3* mutation causes defects in GABAergic transmission and iAChR clustering in GABAergic neurons

**(A)** Representative recordings of endogenous IPSCs from body wall muscles of wild type animals and *unc-3(e151)* mutants. Red traces are expanded views of time segments under the red bars in the upper traces. **(B)** The average frequency of endogenous IPSCs recorded from wild type and *unc-3(e151)* mutants. ***p<0.0001, student's t-test. **(C)** Cumulative...
distribution of amplitudes of endogenous IPSCs recorded from wild type and unc-3(e151) mutants. ***p<0.001, Kolmogorov-Smirnov test. (D) Confocal images showing the dorsal nerve cord of eight animals expressing ACR-12::GFP in GABA neurons for the genotypes indicated. Scale bar, 5 µm. (E) Quantification of the average number of receptor clusters/85 µm of dorsal nerve cord for the genotypes indicated, normalized to wild type. ****p<0.0001, *p<0.05, ANOVA with Dunnett’s multiple comparisons test.
Figure 3. Gross morphological development of GABA motor neurons and muscles does not require cholinergic innervation

(A-C) Confocal images of the ventral nerve cord and body wall muscles posterior to the vulva of an adult wild type (A), unc-3(e151) mutant (B) and acr-2(L/S) transgenic (C) animals expressing a membrane bound GFP in the distal row of body wall muscles (him4p::mCD8::GFP) (A-C) or mCherry in the GABA nervous system (unc-47p::mCherry) (A1-C1). A-C: scale bar, 10 µm. A1-C1: scale bar, 20 µm. Arrows indicate muscle arm shafts and arrowheads indicate muscle arm termini. Asterisks indicate GABA neuron commissures. (D) Quantification of the percentage of the ventral nerve cord region covered by muscle membrane. ns: not significant, ANOVA with Dunnett’s multiple comparisons test.
Figure 4. Impaired cholinergic innervation alters the distribution and size of GABA synapses

(A) Diagram depicting the location of fluorescent reporters used to label GABA pre- (unc-47p::mCherry::RAB-3) and postsynaptic (UNC-49::GFP) structures. (B-F) Confocal images of the ventral nerve cord in adult wild type (B), unc-3(n3435) mutants (C), unc-3(e151) mutants (D), unc-3(e151) mutants expressing an unc-3 cDNA rescuing array (unc-3p::unc-3 cDNA) (E), and transgenic acr-2(L/S) animals (F) co-expressing unc-47p::mCherry::RAB-3 in GABA motor neurons with UNC-49::GFP in body wall muscles. Note areas devoid of synaptic clusters (brackets) and enlarged clusters in unc-3 mutants and acr-2(L/S) transgenic animals. Motor neuron cell bodies are outlined in this figure and all subsequent figures. Scale bar, 20 µm. (G) Average number of pre- and post-synaptic clusters/50 µm, normalized to wild type. (H) Average size of pre- and post-synaptic clusters at GABA synapses. *p<0.05, **p<0.01, ****p<0.0001 compared with either control or unc-3 mutant as indicated, ANOVA with Dunnett’s multiple comparisons test. (I, J) Confocal images of the ventral cord in wild type or unc-3(e151) mutants expressing the GFP-tagged synaptic vesicle marker synaptobrevin (SNB-1::GFP) (I), or the Rim1 homolog (UNC-10::mCherry) (J). Scale bar, 20 µm. (K) Quantification of synaptic cluster number for each marker, normalized to wild type. (L) Quantification of presynaptic cluster area for genotypes indicated. **p<0.01, ***p<0.0001, student’s t-test.
Figure 5. Vesicular release from cholinergic motor neurons shapes GABA synaptic outputs

(A) Confocal images showing the dorsal nerve cord of 8 animals co-expressing ACR-12::GFP in GABA neurons and Tetanus toxin in cholinergic neurons. Scale bar, 5 µm. (B) Quantification of the average number of receptor clusters/85 µm of dorsal nerve cord for the genotypes indicated, normalized to wild type. (C, D) Merged confocal images of unc-47p::mCherry::RAB-3 and UNC-49::GFP labeling in the ventral nerve cord of adult wild type (C) and transgenic animals with specific TetTx expression in cholinergic neurons (D). Scale bar, 20 µm. (E) Average number of pre- and post-synaptic clusters/50 µm for each genotype, normalized to wild type. (F) Average area of pre- and post-synaptic clusters at GABA synapses for each genotype as indicated. ***p<0.001, ****p<0.0001, student’s t-test.
Figure 6. Reduced cholinergic transmission alters GABA synapse density and localization

(A) Confocal images showing the dorsal nerve cord of eight unc-17(e113) and madd-4(ok2854);unc-17(e113) animals expressing ACR-12::GFP in GABA neurons. Scale bar, 5 µm. (B) Quantification of the average number of receptor clusters/85 µm of dorsal nerve cord for the genotypes indicated, normalized to wild type. (C-F) Merged confocal images of...
unc-47p::mCherry::RAB-3 and UNC-49::GFP labeling in the ventral nerve cord region of adult wild type animals (C), unc-17(e113)/vAChT mutants (D), unc-17(e245)/vAChT mutants (E) or unc-29(x29);acr-16(ok789) double mutants that lack functional muscle iAChRs (F). Scale bar, 20 µm. (G) Average number of GABA pre- and post-synaptic clusters/50 µm for each genotype, normalized to wild type. (H) Average size of GABA pre- and post-synaptic clusters. *p<0.05, **p<0.01, ****p<0.0001, ANOVA with Dunnett’s multiple comparisons test.
**Figure 7. Acute reduction of cholinergic transmission during adulthood produces reversible decreases in GABA synapse size**

**(A)** Diagram indicating predicted timeline of wild type *C. elegans* development at 15°C and temperature shifts employed. For 7A and 8A, arrow indicates approximate time of hatch. Temperature shifts to 25°C (red) and time of imaging (green) are indicated. E, embryo. **(B, C)** Merged confocal images of *unc-47p::mCherry::RAB-3* and *UNC-49::GFP* labeling in the ventral nerve cord region of adult wild type animals (B) or *cha-1(y226)* mutants (C) following a shift to 25°C for the durations indicated. Scale bar, 20 µm. **(D, E)** Average number of GABA presynaptic (D) and postsynaptic (E) clusters/50 µm for wild type (black) and *cha-1(y226)* mutants (gray). **(F, G)** Average size of GABA presynaptic (F) and postsynaptic (G) clusters for wild type (black) and *cha-1(y226)* mutants (gray). *p<0.05, **p<0.01, ***p<0.0001, ANOVA with Dunnett’s multiple comparisons test.
Figure 8. Reduced cholinergic neurotransmission during a period of GABAergic synaptogenesis alters GABAergic synaptic connectivity

(A) Diagram indicating predicted timeline of wild type *C. elegans* development at 15°C and temperature shifts employed. (B, C) Confocal images of the ventral nerve cord in adult wild type (B) and *cha-1*(y226) (C) temperature sensitive mutant animals co-expressing GABA
pre- and postsynaptic markers (unc-47p::mCherry::RAB-3 and UNC-49::GFP) grown at the permissive temperature and after an eight hour shift to the non-permissive temperature during the first larval stage. Note that the representative image for the cha-1 no shift control group shown in C is the same as that displayed in Fig. 7C. Scale bar, 20 µm. (D) Average number of GABA presynaptic and postsynaptic clusters/50 µm for wild type (black) and cha-1(y226) mutants (gray). (E,F) Average size of GABA presynaptic (E) and postsynaptic (F) clusters for wild type (black) and cha-1(y226) mutants (gray). *p<0.05, **p<0.01, ***p<0.0001, student’s t-test.
**Supplemental Information**

**Table S1. C. elegans strains used in this work.**

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Development 144: doi:10.1242/dev.141911: Supplementary information

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Figure S1. Analysis of unc-3 expression and madd-4(ok2854)

(A) Confocal images of the posterior ventral nerve cord in adult animals expressing a transcriptional reporter labeling cholinergic neurons (acr-2p::mCherry) and an unc-3 promoter construct fused with GFP (unc-3p::GFP). Asterisks indicate colocalization in cholinergic cell bodies. Scale bar, 5 µm. (B) Confocal images of UNC-29::GFP expressed specifically in body wall muscle (myo-3p::UNC-29::GFP) in wild type and madd-4(ok2854) mutant animals. Asterisks indicate UNC-29::GFP fluorescence in muscle arms. Scale bar, 5 µm. (C) Quantification of the average number of receptor clusters/80 µm of the ventral nerve cord for the genotypes indicated. Each bar represents the mean ± SEM. Numbers in bars indicate the n for each genotype. ****p<0.0001, student’s t test.
Figure S2. GABA neuron morphology is grossly normal in *unc-3* mutants

(A) Confocal images of the posterior dorsal nerve cord in adult animals expressing a transcriptional reporter labeling GABAergic (*unc-47p::mCherry*) neurons in wild type and *unc-3(e151)* mutant animals. Scale bar, 10 µm. (B, C) Quantification of the average number of GABAergic cell bodies (B) and commissures (C) for the genotypes indicated. Each bar represents the mean ± SEM. Numbers in bars indicate the n for each genotype. **p<0.001, student’s t test.
**Figure S3.** GABA synaptic distribution is altered with disruption of cholinergic neurons.

(A-C) Confocal images showing the distribution of GABA synaptic markers (mCherry::RAB-3 and UNC-49::GFP) in extended regions of the ventral nerve cord of adult wild type, *unc-3(e151)*, and transgenic *acr-2(L/S)* animals. Scale bar, 20 µm. (D) Representative line scans depicting colocalization of UNC-49::GFP and mCherry::Rab3.
in a 40 µm region of the ventral nerve cord in wild type and unc-3(e151) mutant animals. Note larger puncta and gaps in fluorescence in unc-3 mutants.

Figure S4. Expression of an unc-3 minigene construct rescues synapse and movement defects in unc-3(e151) mutants

(A) Quantification of the average number of receptors clusters/85 µm of dorsal nerve cord for the genotypes indicated, normalized to wild type. Each bar represents the mean ± SEM. Numbers in bars indicate the n for each genotype. ****p<0.0001, ANOVA with Dunnett’s multiple comparisons test. (B) Top: Average number of pre- and post-synaptic clusters/50 µm, normalized to wild type. Bottom: Average size of pre- and post-synaptic GABA Synapses.

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clusters at GABA synapses/50 µm. Each bar represents the mean ± SEM. Numbers in bars indicate the n for each genotype. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001 for comparisons with either wild type or unc-3 mutant values as indicated, ANOVA with Dunnett’s multiple comparisons test. For A and B, values for wild type, unc-3 mutant and cDNA rescue were reproduced from Figures 2 and 4 respectively. (C) Average distance travelled in 30 s for the genotypes indicated. Each bar represents the mean ± SEM. Numbers in bars indicate the n for each genotype. ****p<0.0001, ***p<0.001, ANOVA with Dunnett’s multiple comparisons test.

Figure S5. Cell specific expression of tetanus toxin in cholinergic neurons alters neuronal activity but not neuronal organization

(A) Schematic representation of the motor circuit showing connectivity of cholinergic (green) and GABAergic neurons (red). Sites of tetanus toxin action are denoted by X.

(B) Time course of paralysis of adult animals on 1 mM aldicarb, an inhibitor of cholinesterase. The percentage of animals paralyzed was calculated every 15 minutes for two hours. Each data point represents the mean ± SEM of n=10 or n=8 assays for
wild type or \textit{ACH::TetTx} respectively. (C, D) Confocal images of the posterior ventral nerve cord in adult animals co-expressing transcriptional reporters labeling cholinergic (\textit{unc-17p::GFP}) and GABAergic (\textit{unc-47p::mCherry}) neurons in wild type or with cell-specific expression of TetTx as indicated. The identities of motor neurons based on cell body position are indicated (green, cholinergic neurons) (red, GABAergic neurons).

\textbf{Figure S6. Neuronal development and locomotory behavior in \textit{cha-1(y226)} mutant animals}
(A) The number of GABA motor neuron cell bodies (visualized by unc-47p::mCherry expression) present in wild type or cha-1(y226) animals grown at 15°C at the time points indicated. All time points are taken from the 21+ cell stage. Graph represents the average number of cell bodies for 10-15 animals per time point for each genotype. (B) Forward velocity of adult wild type and cha-1(y226) animals following the shifts indicated. See Figure 7A for details of time shift. **p<0.001, ***p<0.0001, ANOVA with Tukey’s multiple comparisons test. (C) Forward velocity of young adult wild type and cha-1(y226) animals after being shifted to a non-permissive temperature for eight hours spanning the L1/L2 transition. See Figure 8A for details of temperature shift. *p<0.05, student’s t-test.