Kinesin-3 mediated delivery of presynaptic neurexin stabilizes growing dendritic spines and postsynaptic components in vivo

Devyn Oliver, Shankar Ramachandran, Alison Philbrook, Christopher M. Lambert, Ken C. Q. Nguyen, David H. Hall, *Michael M. Francis

1Department of Neurobiology, University of Massachusetts Medical School, Worcester, MA 01605
2Department of Neuroscience, Albert Einstein College of Medicine, Bronx, NY 10461

*Corresponding author: michael.francis@umassmed.edu

Key words:
dendritic spines, neurexin, kinesin, C. elegans, synapse development, synaptic maintenance, active zone, protein trafficking
Contributions Summary

D.O. generated strains, transgenic lines, molecular constructs, confocal and electron microscopy images, and data analysis. S.R. performed all calcium imaging experiments and analysis. C.M.L. generated destination vectors and nrx-1 constructs. D.H.H. and K.C.Q.N. aided in electron microscopy experiments and analysis. M.M.F and D.O designed and interpreted results of all experiments and wrote the manuscript.

Acknowledgements

Nematode strains were provided by the Caenorhabditis Genetics Center which is funded by the NIH National Center for Research Resources. Thank you to Dori Schafer for access to 3D rendering Imaris software. Thank you to Dirk Albrecht’s laboratory for advice with hydrogel immobilization. Thank you to the laboratories of: Dong Yan, Kang Shen, Andres Maricq, Michael Nonet, Peri Kurshan, and Gert Jansen for providing reagents. Thank you to William Joyce and Michael Gorczyca for technical assistance. Thank you to Steve Cook for help with analysis of electron micrographs and John White for use of N2U electron micrographs. Lastly, we thank members of the Francis laboratory for manuscript comments.

Funding

This research was supported by NIH NINDS RO1NS064263 (MMF), R21NS101649 (MMF), NIH OD 01943 (DHH), and F31NS103365 (DO). The Philips CM10 TEM electron microscope used in this study was acquired through a NIH Shared Instrumentation Grant (1S10OD016214-01A1) at Albert Einstein College of Medicine.
Abstract

A high degree of cell and circuit-specific regulation has complicated efforts to precisely define roles for synaptic adhesion proteins in establishing circuit connectivity. Here, we take advantage of the strengths of *C. elegans* for cell-specific analyses to investigate molecular coordination of pre- and postsynaptic development. We show that developing dendritic spines emerge from the dendrites of wild type GABAergic motor neurons following the localization of active zone proteins and the formation of immature synaptic vesicle assemblies in presynaptic terminals. Similarly, clusters of postsynaptic receptors and F-actin are visible in GABAergic dendrites prior to spine outgrowth. Surprisingly, these developmental processes occur without a requirement for synaptic activity. Likewise, the initial stages of spine outgrowth and receptor clustering are not altered by deletion of the *C. elegans* ortholog of the transsynaptic adhesion protein, neurexin/NRX-1. Over time, however, dendritic spines and postsynaptic receptor clusters are destabilized in the absence of presynaptic NRX-1/neurexin and collapse prior to adulthood. The kinesin-3 family member, UNC-104, delivers NRX-1 to presynaptic terminals and ongoing UNC-104 delivery is required into adulthood for the maintenance of postsynaptic structure. Our findings provide novel insights into the temporal order of synapse formation events *in vivo* and demonstrate a requirement for transsynaptic adhesion in stabilizing mature circuit connectivity.
Introduction

The capabilities of neural circuits to perform specific functions arise from the patterns of synaptic connections between their partnering neurons. The organization of these connections is circuit-specific and established through a complex process that involves the coordinated assembly and maturation of specialized pre- and postsynaptic structures on appropriate partnering neurons, and their maintenance in the mature nervous system. We now have a generalized understanding of synapse structure. Active zone (AZ) proteins and neurotransmitter-filled synaptic vesicles position near the presynaptic membrane for rapid release while neurotransmitter receptors are clustered at high density in apposition to these sites in order to ensure the fidelity of synaptic communication. Genetic studies have identified numerous mutations that alter circuit connectivity or affect the overall structural organization of synapses (Hendi et al., 2019) (Zhen & Jin, 1999) (Richmond et al., 1999). However, for many of the synapse-associated proteins affected by these mutations, we do not yet have a mechanistic understanding of their roles in establishing synaptic connections, or how their disruption may lead to alterations in synapse development and structure. Gaining an enhanced understanding of the sequence of events involved in synapse assembly, maturation, and maintenance, and their relative timing in vivo is critical for addressing these questions.

Several prior studies have examined molecular events during synaptogenesis. In vivo studies in C. elegans and Drosophila have largely focused on the formation of the presynaptic active zone. Collectively, these studies provide compelling evidence for a model where active zone assembly occurs sequentially. While there is some variability across synapse type, the early stages of this process are generally organized by the highly conserved synaptic scaffolds SYD-2/Liprin-α and SYD-1/Rho GTPase, which then recruit additional key conserved AZ proteins such as ELKS-1/Bruchpilot, Piccolo family members, and UNC-10/RIM for subsequent stages of assembly, including clustering of Ca^{2+} channels and the recruitment of synaptic
vesicles (Zhen & Jin, 1999) (Bohme et al., 2016) (Fouquet et al., 2009) (Patel et al., 2006) (Dai et al., 2006). Remarkably, nascent AZs can assemble quite rapidly (within minutes) but then often undergo a more extended period of maturation that can last for several hours (Nagerl et al., 2007) (Lipton et al., 2018) (Fouquet et al., 2009) (McDonald et al., 2020). The relationship of postsynaptic development to presynaptic assembly remains less clear. Live imaging studies of cultured mouse hippocampal neurons have suggested that the assembly of presynaptic components precedes recruitment of postsynaptic receptors and scaffolds (Friedman et al., 2000). More recently, 2-photon imaging of organotypic slice cultures from rat hippocampus showed that new spines accumulate glutamate receptors concurrently with their growth and are competent to participate in transmission within a few hours after outgrowth (Zito et al., 2009).

Comparatively few in vivo studies have investigated coordinated pre- and postsynaptic development and significant questions remain about molecular links between these processes.

Evolutionarily conserved synaptic adhesion proteins, such as neurexins, are prime candidates for coordinating pre- and postsynaptic events during synaptogenesis. Their importance is underscored by the fact that neurexin alterations are associated with cognitive disease (Sudhof, 2008). For example, mutations in human Neurexin1 are associated with several neuropsychiatric disorders, including schizophrenia and autism spectrum disorders (Kasem et al., 2018). Neurexins are typically localized presynaptically and linked to postsynaptic binding partners such as dystroglycans, LRRTMs, neuroligins, and cerebellins, through extracellular laminin and EGF-like repeats to establish transsynaptic connections. Neurexin mediated transsynaptic signaling has been implicated in key aspects of synapse development and function. For example, loss of presynaptic neurexin at the fly neuromuscular junction increases the length of the presynaptic density, and also alters the size and molecular composition of apposed muscle glutamate receptor clusters (J. Li et al., 2007) (Owald et al., 2012). The broad brain distribution of neurexins and the many neurexin isoform variants in the
nervous system enable complex, cell-type specific functions for neurexins at synapses (Missler & Sudhof, 1998) (Sudhof, 2017). For example, conditional deletion of mouse neurexins revealed strikingly different functions across synapses formed by cortical inhibitory interneurons and those formed by cerebellar climbing fibers onto Purkinje neurons (L. Y. Chen et al., 2017). Genetic tools that enable cell-type specific analysis are therefore critical for uncovering precise functional roles of synaptic adhesion proteins within the context of individual neural circuits. Importantly, molecular mechanisms for their trafficking and delivery to synapses also remain incompletely defined.

We previously identified finger-like dendritic protrusions from the dendritic processes of C. elegans DD GABAergic motor neurons (Figure 1A) (Philbrook et al., 2018). Characterization of these structures by our laboratory and others pointed towards the idea that they receive synaptic input from presynaptic cholinergic motor neurons and serve analogous roles to dendritic spines in the mammalian brain (J. G. White et al., 1976) (Cuentas-Condori et al., 2019). Deletion of nrx-1, the sole C. elegans ortholog of neurexin, disrupts these dendritic spines and impairs proper localization of cholinergic receptor clusters to postsynaptic sites on GABAergic dendrites (Philbrook et al., 2018). Here we define the order and timing of pre- versus postsynaptic development at spine-associated synapses and elucidate the role of neurexin in the coordination of these processes in vivo. We find that clusters of both presynaptic proteins and postsynaptic receptors are clearly visible prior to spine outgrowth. In the absence of neurexin, immature spines and receptor clusters form initially but then disappear, indicating that neurexin is required for synapse stabilization and maturation rather than synaptogenesis. These maintenance and maturation processes are supported by kinesin-3/UNC-104 motor dependent transport of NRX-1 to presynaptic terminals. Together, our results suggest that early and ongoing axonal delivery of neurexin to sites of synapse formation is critical for stabilization of postsynaptic structures. Our findings cleanly dissect a key role for neurexin mediated adhesion...
in sculpting connectivity, specifically in promoting the stabilization and maturation of growing synapses and spines.
Results

We initially explored morphological features of mature spines using 3D rendering of spines from confocal imaging. We identified several morphological classes (Figure 1B) based on the number of spine heads and the ratio between the length of the spine to the width of the spine head and spine neck. We found that a majority of DD spines can be categorized into four morphological classes (percentage ± SD): mushroom (5 ± 1%), branched (5 ± 1%), stubby (73 ± 3%), and thin (15 ± 1%). The morphological diversity we observed by light microscopy was also evident in ultrastructural studies using serial electron microscopy (Figure S1.1). Clusters of small synaptic vesicles were also evident in neurites (of cholinergic VA/VB neurons, presumably) immediately adjacent to the tips of DD spines, indicating putative sites of synaptic input.

To explore how the neuronal cytoskeleton may contribute to spine morphology, we examined the localization of F-actin and tubulin in DD dendrites using DD neuron-specific expression GFP::UtrCH (GFP fused to the Utrophin calponin homology domain) (Chia et al., 2014) and TBA-1::GFP (GFP-tagged α-tubulin) (Yan et al., 2013) respectively. We found that the two markers occupy distinct territories. Coexpression studies showed that F-actin is strongly enriched within dendritic spines (Figure 1C-E), while tubulin occupies the dendritic shaft (Figure 1F,G) with more variable localization to the spine base (Figure S1.2). Previous findings for vertebrate neurons indicate a similar segregation of actin and tubulin to the spines and shaft of the dendrite respectively (Gu et al., 2008) (Hu et al., 2008) (Jaworski et al., 2009).

DD spines are positioned immediately adjacent to the axon terminals of cholinergic neurons that are predicted to be their presynaptic partners by electron microscopy (Figure S1.1) (J. G. White et al., 1976) (J. G. White et al., 1986) (Cuentas-Condori et al., 2019). We previously showed stimulation of cholinergic motor neurons elicited Ca^{2+} responses in
GABAergic motor neurons (Philbrook et al., 2018). To confirm that spines on DD GABAergic motor neurons represent sites of functional synaptic contacts, we asked whether stimulation of presynaptic cholinergic neurons was sufficient to elicit calcium responses in DD spines. We performed in vivo calcium imaging of evoked responses in spines using combined expression of a membrane-associated GCaMP6f calcium sensor in DD GABAergic neurons and a red shifted channelrhodopsin, Chrimson, in cholinergic neurons (Philbrook et al., 2018) (Figure 1H, I, J) (Figure S1.3). Following 5 seconds of baseline recording (488 nm, 100 ms exposure), we measured calcium responses to presynaptic depolarization (5 s, 625 nm, 30 mW/cm²) (Figure 1I). We noted significant fluorescence increases in spines that were timed with the onset of stimulation and were not present in the absence of retinal. These typically occurred simultaneously across multiple spines and returned to baseline within 5 s following stimulation. Importantly, mutation of the cholinergic vesicular transporter/VACHT unc-17, reduced evoked calcium responses by 81%, indicating the Ca²⁺ responses we observed in dendrites were dependent on presynaptic acetylcholine release (Figure 1H, I, J). Our findings are consistent with those of another recent study (Cuentas-Condori et al., 2019), and indicate that spines on DD GABAergic neurons are primary sites of synaptic contact.

Mitochondria and other organelles are often localized near postsynaptic specializations in order to sustain synaptic function (Vos et al., 2010) (Karagas & Venkatachalam, 2019). To examine the distribution of organelles within DD dendrites and spines, we used fluorescent reporters labeling the Golgi apparatus (AMAN-2::GFP) (Edwards et al., 2013), mitochondria (Pre-Su9::GFP) (Sarasija & Norman, 2015), and endoplasmic reticulum (ER) (RFP::TRAM-1) (Rolls et al., 2002) (Edwards et al., 2013) (Figure S1.4). We found that Golgi is exclusively labeled in the DD cell bodies (Figure S1.4A). In contrast, mitochondria and rough ER are dispersed throughout the main dendritic shaft and near the base of spines (Figure S1.4B-E), perhaps suggesting involvement in spine function (Perez-Alvarez et al., 2020) (Z. Li et al.,
2004). Consistent with our light microscopy observations, we also observed mitochondria in the main dendritic process of DD GABAergic neurons near spines by electron microscopy (Figure S1.1B,C).

The development of synapses at DD GABAergic spines

Using the synapses at DD spines as a model, we next investigated the relative timing of pre- and postsynaptic events during synapse formation. *C. elegans* progress through four larval stages of development (L1-L4) prior to adulthood. DD neurons undergo a well-characterized program of synaptic remodeling such that the mature circuit organization is established during the transition from L1 to L2 stage (J. G. White et al., 1978). During this period of remodeling, newly born cholinergic neurons form new synaptic connections with the ventral dendrites of DD GABAergic neurons. This synaptic remodeling event offers a well-defined temporal window to investigate *de novo* formation of neuron-neuron synaptic connections *in vivo*. We focused on understanding the time course of spine formation relative to 2 key events in synaptogenesis: 1) the formation of presynaptic release sites, and 2) the clustering of postsynaptic receptors and F-actin (Figure 2).

The appearance of newly born ventral cholinergic motor neuron cell bodies and processes progressed anteriorly to posteriorly starting roughly 16 hours after hatch, consistent with the birth and integration of cholinergic neurons during this time frame (Figure S2.1) (Kurup & Jin, 2016). We analyzed the distribution of pre- and postsynaptic markers at timepoints before (12 hours after hatch), during (16, 18, 20 hours after hatch) and following (24, 32 hours after hatch, L4 stage/42-50 hours) DD synaptic remodeling (Figures 2, S2.1, and S2.2). We first assessed the formation of the active zone and presynaptic specialization by analyzing clustering of the Piccolo-like active zone protein, CLA-1, (GFP::CLA-1e) and the synaptic vesicle-associated protein synaptobrevin/SNB-1 (SNB-1::GFP) (Figure 2A, D-E) (Xuan et al., 2017).
Overall, we found CLA-1 is more discretely localized to putative synapses than SNB-1, both at earlier timepoints and throughout development, consistent with its specific localization to active zones. Immediately prior to remodeling (~12 hours after hatch), we did not observe significant localization of either presynaptic CLA-1 or SNB-1 adjacent to the DD process. Shortly after (16 hours after hatch), we noted the initial appearance of SNB-1- and CLA-1-associated fluorescence in presynaptic cholinergic processes. Over the next 4 hours (20 hours after hatch) individual CLA-1 clusters were more clearly distinguishable, suggesting CLA-1 association with developing active zone structures (Figure 2A, D, L). Synaptic vesicle fluorescence was initially diffuse at 16 hours after hatch and became more clearly organized into discrete puncta over a similar time course to CLA-1 (Figure 2A, E, L). We did not observe the emergence of DD dendritic spines until 24-32 hours after hatch, well after initial active zone formation and recruitment of synaptic vesicles (Figure S2A, C-E). Spines continued to mature through L4 stage, increasing in both length and number (Figure 2.2C).

We next examined the clustering of postsynaptic acetylcholine receptors in DD dendrites (ACR-12::GFP). Prior to remodeling (12 hours after hatch), we did not observe detectable levels of cholinergic receptors in the DD process. Surprisingly, we noted immature receptor clusters faintly visible in the dendritic shaft by 16 hours after hatch, well prior to the emergence of dendritic spines (Figure 2B, F, L). These receptor clusters increased in number and redistributed towards the tips of growing spines by 32 hr (Figure 2B, F, L). We found a similar developmental trend using DD neuron-specific labeling of the LEV-10 transmembrane auxiliary protein (Gally et al., 2004), previously shown to concentrate in spines (He et al., 2019) (Figure S2.3). By L4 stage, receptor and LEV-10 clusters are clearly visible at the tips of mature spines. Together, our analysis indicates that the initial stages of development of both pre- (CLA-1 and SNB-1 clusters) and postsynaptic (AChR and LEV-10 clusters) structures occur prior to spine outgrowth, raising questions about how these initial processes may be regulated.
To begin to address this question, we analyzed the distribution of F-actin in dendrites of the DD neurons during synapse formation **(Figure 2B, G-L)**. We noted that clusters of F-actin were evident in the ventral DD process prior to the completion of synaptic remodeling (12 hours after hatch), prior to presynaptic CLA-1 and synaptic vesicle accumulation, and before the clustering of postsynaptic receptors. Dendritic F-actin-based structures became more abundant coincident with increases in the number of presynaptic CLA-1 and postsynaptic receptor clusters. To investigate this process in real-time, we used live imaging to examine the dynamics of postsynaptic F-actin (GFP::UtrCH) in the developing DD dendrite. We found that F-actin is highly dynamic during early developmental stages (16-20 hours after hatch) compared to L4 stage where the circuit has completed maturation **(Figure S2.4, Video 1, and Video 2)**. In young animals, GFP::UtrCH clusters often shuttled out of the cell body to the main dendritic process, perhaps indicating delivery of postsynaptic F-actin to sites of postsynaptic assembly. By 24 hours after hatch, we observed clear co-localization of F-actin with newly formed AChR clusters in the dendritic shaft that are positioned for developmental translocation into growing dendritic spines. In mature animals, AChR clusters are stably sequestered at the tips of spines, while F-actin occupies the spine head and neck **(Figure 2H-K)**.

**Dendritic spines form in the absence of presynaptic activity**

Our above analysis indicated that the localization of synaptic vesicles and active zone proteins, such as CLA-1, occur prior to spine outgrowth. In the rodent brain, spine morphogenesis is clearly regulated by presynaptic activity (Sala & Segal, 2014) (Saneyoshi et al., 2010) (Engert & Bonhoeffer, 1999) (De Roo et al., 2008), but recent evidence suggests initial spine outgrowth may proceed independently of synaptic activity (Lu et al., 2013) (Sigler et al., 2017) (Sando et al., 2017). We therefore next asked whether presynaptic cholinergic activity is important for the formation of DD dendritic spines. To address this question, we analyzed the
number of dendritic spines (Pflp-13::mCherry or Pflp-13::myrGFP) and cholinergic receptors (Pflp-13::ACR-12::GFP) at L4 stage in strains carrying mutations that affect various aspects of synaptic function and neuronal excitability (Table 1, Figure S3.1).

Surprisingly, the abundance of spines and receptor clusters at L4 stage were not significantly affected by disruption of ACh synthesis in cholinergic motor neurons (mutation of the vesicular acetylcholine transporter unc-17) or by a strong reduction in synaptic vesicle exocytosis (mutation of the syntaxin binding protein unc-18). Similarly, loss-of-function mutations in genes required for dense core vesicle release (CAPS/unc-31) and Ca\(^{2+}\) signaling (VGCC/unc-2, CaM Kinase II/unc-43) did not significantly reduce the number of spines or receptor clusters in the mature circuit (Table 1, Figure S3.1). Our data indicate that alterations in synaptic activity do not affect the abundance of L4 stage dendritic spines. However, we cannot exclude the possibility of developmental delays in spine formation (Cuestas-Condori et al., 2019). Notably, many of the mutations we examined significantly altered spine length and the size of receptor clusters (Table 1), suggesting that activity may influence aspects of spine morphology and post-synaptic structure. Together, our findings point toward a model where initial spine formation and postsynaptic development proceeds without a strong requirement for synaptic activity, while subsequent spine morphogenesis may be affected.

Presynaptic NRX-1 stabilizes postsynaptic components

In previous work, we found that mutations in nrx-1, the sole C. elegans ortholog of neurexin, eliminate DD dendritic spines and disrupt cholinergic receptor clusters in the mature C. elegans circuit (Philbrook et al., 2018), indicating alternate activity-independent mechanisms may be critical for spine formation. Consistent with these findings, we found that mutations of nrx-1 disrupted dendritic calcium responses evoked by presynaptic stimulation of cholinergic motor neurons (Figure S3.2). To examine possible early involvement of presynaptic NRX-1 in
the establishment of spines we asked when NRX-1 first localizes to presynaptic terminals of cholinergic motor neurons. Axonal clusters of NRX-1::GFP are present by 16 hours after hatch (Figure 3A, B), approximately the same time at which assemblies of the presynaptic scaffold CLA-1 become visible (Figure 2). Diffuse synaptic vesicle material is also visible at this time (Figure 2) but has yet to be organized into discrete clusters.

The early arrival of NRX-1 to synaptic terminals could indicate a role in either initial synapse formation or subsequent stabilization and maturation. To distinguish between these possibilities, we analyzed the development of dendritic spines and cholinergic receptor clusters in nrx-1(wy778) null mutants. We quantified spine and AChR cluster number over a similar time course as previously completed for wild type. Surprisingly, we noted that immature spines and receptor clusters were clearly evident in nrx-1 mutants during early development, albeit at slightly reduced density relative to wild type (Figures 3C-F, S3.3). The density of spines in nrx-1 mutants increased significantly over the next several hours until 24 hours after hatch (Figure 3C,D). However, after this time, spine density decreased dramatically such that nrx-1 DD dendrites were almost completely devoid of spines by L4 stage, consistent with our previous work (Philbrook et al., 2018). Measurements of spine formation and disassembly in live imaging studies of wild type and nrx-1 mutants offered further support for this conclusion. Mature wild type spines (L4) were remarkably stable over 1-2 hours of recording, but were more dynamic in the developing circuit (16-20 hrs after hatch) (Figure S3.4, Videos 3-5). Developing spines in nrx-1 mutants were also highly dynamic. Notably, heightened spine dynamics persisted into more advanced developmental stages for nrx-1 mutants compared with wild type animals. For example, at 21-24 hours after hatch, almost 90% of wild type spines were stable over the recording period, while less than 50% of nrx-1 mutant spines remained stable, indicating decreased stability in the absence of NRX-1.
Receptor clustering followed a similar trend to spine formation. The number and fluorescence intensity of ACR-12 clusters at 16 hours after hatch in nrx-1(wy778) animals (6.8 ± 0.8 clusters/25 µm) were similar to wild type (5.6 ± 0.7 clusters/25 µm) (Figure 3E-F, S3.3). Within a few hours however, ACR-12 cluster number and intensity decreased significantly in nrx-1 mutant dendrites, and remained low throughout the remainder of development (Figure 3E-F, S3.3). Our time course analysis indicated that both spines and receptor clusters initially form normally in nrx-1 mutants, but rapidly disappear as animals proceed through development in the absence of NRX-1. Remarkably, we found that the organization of dendritic F-actin was similarly affected by nrx-1 deletion. GFP::UtrCH was arranged similarly in nrx-1 mutants and wild type in early development (16 hrs after hatch), positioned in discrete clusters along the main dendritic process (Figure 3G). By L4 stage, wild type F-actin was almost entirely associated with dendritic spines. This organization was strikingly altered in L4 stage nrx-1 mutants. GFP::UtrCH was diffusely localized along the length of nrx-1 mutant dendrites, indicating that presynaptic NRX-1 promotes dendritic F-actin assemblies (Figure 3G). Taken together, our results indicate that presynaptic NRX-1 is dispensable for the earliest stages of postsynaptic assembly and spine formation, but is critical for stabilizing dendritic spines and AChR clusters and promoting their maturation.

Kinesin-3/UNC-104 transports presynaptic neurexin to cholinergic terminals in order to stabilize postsynaptic components

To investigate this model further, we sought to determine how presynaptic neurexin is transported to active zones in vivo. Prior work showed that both synaptic vesicles and CLA-1 depend on the Kinesin-3 motor UNC-104 for their delivery to synapses (Hall & Hedgecock, 1991) (Xuan et al., 2017) (Figures S4.1 and 4). In particular, disruption of unc-104 causes an accumulation of synaptic vesicles within neuronal somata and a corresponding loss of synaptic vesicles within axons (Hall & Hedgecock, 1991) (Figure S4.1). Consistent with potential
involvement of UNC-104 in NRX-1 trafficking, we found that UNC-104 and NRX-1 are partially colocalized in cholinergic axons (Figure S4.2). unc-104(e1265) carries a D1497N mutation in the PH domain of UNC-104 that impairs cargo binding and leads to a severe reduction in UNC-104 abundance in axons (Kumar et al., 2010). NRX-1 endogenously tagged with GFP is strongly visible in neuronal processes within the nerve ring and nerve cords of wild type animals (Figure S4.3). In contrast, NRX-1::GFP clusters were strikingly decreased in the nerve cords of unc-104(e1265) mutants (Figures 4A,C, S4.4). A similar deficit in axonal NRX-1::GFP localization was evident with specific cholinergic expression of NRX-1::GFP in unc-104 mutants (Figures 4B,C, S4.4). By contrast, NRX-1::GFP fluorescence was significantly increased (5.4-fold) in unc-104 mutant cholinergic somata compared to wild type (Figure S4.4). Accumulation of NRX-1 in cholinergic cell bodies, coupled with a decrease in axons, indicates a failure of NRX-1 transport in the absence of functional UNC-104. Consistent with this, cholinergic-specific expression of wild type unc-104 in unc-104 mutants restored axonal NRX-1::GFP localization, indicating a cell autonomous requirement (Figure 4 B,C). In contrast, mutation of the Kinesin-1 motor unc-116 did not produce significant accumulation of NRX-1::GFP in cholinergic somata and caused comparatively modest decreases in axonal NRX-1::GFP, demonstrating preferential involvement of Kinesin-3 for NRX-1 transport (Figure S4.5). Live imaging studies offered additional evidence that synaptic vesicle and NRX-1 trafficking may share a common dependence on UNC-104. We found that NRX-1::GFP trafficking events occurred with similar anterograde and retrograde velocities to SNB-1::GFP (labeling synaptic vesicles) trafficking events, though less frequent (Figure S4.6, Videos 6 and 7). Prior studies of active zone protein transport have noted these events occur at reduced frequency compared with synaptic vesicle trafficking events (Lipton et al., 2018). Together, our analyses point to a specific requirement for the UNC-104/KIF1A motor in delivery of NRX-1 to presynaptic terminals.
The density of spines and AChR clusters were also severely reduced in L4 stage unc-104(e1265) mutants (Figure 4 G-I), similar to nrx-1 mutants (Figure 4J). These effects were rescued by either native or cholinergic expression of wild type unc-104 in unc-104(e1265) mutants, but not by GABA- or muscle-specific unc-104 expression (Figure 4 G-I), demonstrating that presynaptic UNC-104-mediated transport is critical in cholinergic axons for postsynaptic spine development and receptor localization.

Notably, we observed that several other presynaptic AZ components also require UNC-104 for their delivery. In addition to CLA-1, transport of both ELKS-1/ELKS/CAST and UNC-10/RIM were severely disrupted by mutation of unc-104, indicating that Kinesin-3 cargo selection and transport is key for the synaptic delivery and assembly of several key active zone constituents in these neurons (Figure 4C-F). In contrast, the synaptic localization of these active zone proteins was not appreciably affected by nrx-1 deletion (Figure S4.7). We next asked how a failure in delivery of AZ components may contribute to the severe reductions in spines and receptor clusters we observed in unc-104 mutants. We compared postsynaptic organization in strains carrying deletion mutations in either nrx-1, cla-1, unc-10 or elk-1. Remarkably, only mutation of nrx-1 produced significant decreases in dendritic spines (Figure 4J), demonstrating that presynaptic ELKS, UNC-10 and CLA-1 are dispensable for postsynaptic development. Our results indicate a specific requirement for NRX-1 in the stabilization and maturation of postsynaptic structures and provide evidence that a failure in synaptic delivery of NRX-1 is a primary causal factor in the postsynaptic structural defects of unc-104 mutants.

The stabilization of mature dendritic spines requires ongoing synaptic delivery of NRX-1

Our results suggest UNC-104 mediated transport positions NRX-1 at the presynaptic terminal in the early stages of synapse formation where it acts to stabilize growing postsynaptic structures, including spines and receptor clusters. We next sought to address whether there is
an ongoing requirement for presynaptic UNC-104-dependent transport in the stabilization of mature spines. To address this question, we used the previously characterized temperature-sensitive allele, *unc-104(ce782)*. *unc-104(ce782)* animals carry a G105E missense mutation in the motor domain of UNC-104 (Edwards et al., 2015). When grown at permissive temperature (13.5°C), *unc-104(ce782)* axonal synaptic vesicle abundance and animal motility are modestly reduced compared to wild type. In contrast, synaptic vesicles are completely absent from *unc-104(ce782)* axons following growth at restrictive temperature (20-25°C), and animal motility is severely compromised within 12 hours of a shift to 23°C (Edwards et al., 2015). We raised *unc-104(ce782)* animals at the permissive temperature (13.5°C) to L4 stage (*Figure 5*). We then shifted L4 stage animals to the restrictive temperature (25°C) for 16-20 hrs, and quantified NRX-1::GFP localization and spine density immediately following this shift. Axonal NRX-1::GFP clusters were strikingly decreased in *unc-104(ce782)* mutants subjected to the temperature shift compared with control animals subjected to the same shift or *unc-104(ce782)* mutants raised continuously at the permissive temperature (*Figure 5B,C*). *unc-104(ce782)* mutants grown at the permissive temperature had a slightly reduced number of spines overall (*Figure 5E*), compared with wild type. A shift to the restrictive temperature at L4 stage also produced a striking reduction in spine density for *unc-104(ce782)* mutants (*Figure 5D, E*), but not for wild type animals subjected to the same temperature shift. We obtained similar results for *unc-104(ce782)* mutants using an earlier shift to the restrictive temperature (at L3 stage) (*Figure S5.1*). Taken together, our findings indicate an ongoing requirement for UNC-104 transport that extends well beyond the period of initial synapse formation and spine outgrowth. Among the UNC-104 cargoes we investigated, only deletion of *nrx-1* produces a significant reduction in spine density at L4 stage. We therefore propose a model where ongoing UNC-104 delivery of presynaptic NRX-1 is critical for postsynaptic maturation and maintenance of mature spines.
Discussion

**C. elegans GABAergic motor neurons are decorated with functional dendritic spines**

Dendritic spines are known to act as specialized sites for compartmentalizing neurotransmission and are widely observed across various classes of neurons in both vertebrates and invertebrates (Tonnesen & Nagerl, 2016). Though suggested by prior EM studies (J. G. White et al., 1976) (J. G. White et al., 1986), the presence of functional dendritic spines on *C. elegans* neurons had not been fully appreciated until recently (Philbrook et al., 2018) (Cuentas-Condori et al., 2019). Here we examined the developmental trajectory of postsynaptic structures (dendritic spines and cholinergic receptor clusters) located on GABAergic motor neurons relative to presynaptic release sites and showed that the stabilization of growing dendritic spines requires presynaptic NRX-1/neurexin.

Neural circuit organization and performance is dictated by the locations of synaptic connections and the identities of the interconnected neurons. Elucidation of the processes governing the construction of synapses and their coordination across presynaptic axons and postsynaptic dendrites are therefore fundamental for understanding circuit assembly and function. However, relatively few studies have monitored in real-time the coordinated assembly of pre- and postsynaptic specializations *in vivo*. Our work tackled this important question, taking advantage of the GABAergic DD spine model and the cellular precision offered by the *C. elegans* motor circuitry. We find that these dendritic spines share many of the hallmark features of vertebrate dendritic spines, but also have some key differences. (1) *C. elegans* spines have morphological similarities with mammalian spines and can be classified into comparable morphological categories. (2) F-actin is localized to spines while tubulin is mainly present in the dendritic shaft. F-actin is a major component of mammalian dendritic spines and is important for spine structural dynamics (reviewed in (Borovac et al., 2018)). (3) Mitochondria and rough
endoplasmic reticulum are positioned near spines, either in the spine neck or at the base of spines in the dendritic shaft. Dendritic organelles contribute to key processes involved in spine formation and spine metabolism (Liu & Shio, 2008). Ribosomes and mitochondria positioned near or within dendritic spines may aid in modifying dendritic spine activity, local protein synthesis, and act as synaptic ATP and Ca\(^{2+}\) sources (Steward & Levy, 1982) (Aakalu et al., 2001) (Chirillo et al., 2019) (Z. Li et al., 2004). (4) Presynaptic stimulation elicits calcium responses in spines on GABAergic dendrites. DD spines are located opposite cholinergic release sites and calcium responses are eliminated by mutation of the vesicular acetylcholine transporter. (5) Mature *C. elegans* spines display limited overt activity-dependent dynamics in wild type animals and appear comparatively stable relative to spines in rodents. This raises the important question of what is the functional role of spines in GABAergic neurons? Prior studies have provided evidence for compartmentalized calcium signals in *C. elegans* neurons (Hendricks et al., 2012). Therefore, one intriguing possibility is that, similar to the situation in mammals, spines serve to compartmentalize calcium and perhaps other biochemical signals in GABAergic neurons. Spines may also be required in order to achieve the unusual dyadic arrangement of synapses in the ventral nerve cord, where presynaptic specializations of cholinergic motor neurons are positioned for transmission onto both GABA motor neuron and muscle postsynaptic partners (Hall & Russell, 1991). In this case, GABAergic spines may have developed to intercept cholinergic release sites onto muscles, as suggested previously (J. White, 2018).

Ventrally-directed (VA/B) cholinergic neurons are born and integrated into the motor circuit post-embryonically, forming new synaptic connections with both ventral muscles and dorsally-directed (DD) GABAergic motor neurons. We found that F-actin is compartmentalized to discrete regions within the dendritic shafts of DD neurons very early in the development of the circuit (prior to the appearance of spines) and later becomes exclusively localized to dendritic
spines. F-actin labeling of spines is apparent as soon as they are detectable, suggesting that F-actin assemblies may participate in the earliest stages of spine development. Prior to remodeling of DD neurons, sites of GABA release are located on the ventral processes of DD neurons. As F-actin based structures are typically associated with release sites (Chia et al., 2012) (Meng et al., 2015), our finding that F-actin is clustered in ventral DD processes prior to remodeling may suggest that F-actin initially associated with release sites persists in the DD neurite through remodeling and is then repurposed for postsynaptic development, perhaps acting as a landmark or an actin-based mechanism for stimulating post-synaptic maturation.

Notably, we also observed punctate cholinergic receptor fluorescence in GABAergic dendrites several hours prior to the formation of dendritic spines. These immature receptor clusters colocalize with dendritic F-actin assemblies suggesting that receptors are trafficked into dendrites and positioned with F-actin to rapidly populate growing spines. Indeed, we observed that receptor clusters are visible in growing spines as soon as spines can be clearly resolved. This is consistent with recent findings from time-lapse 2-photon imaging of hippocampal organotypic slice cultures showing that receptor accumulation in spines occurs concurrently with spine outgrowth (Zito et al., 2009). Similarly, we found that synaptic material accumulates in presynaptic axons prior to the emergence of spines from dendritic processes. In particular, initial accumulations of the presynaptic scaffold CLA-1 and the synaptic vesicle marker synaptobrevin/SNB-1 were visible in the axon with a similar time-course to the appearance of dendritic cholinergic receptor clusters. Presynaptic material became more clearly localized to discrete puncta over the next 4-6 hours, occurring roughly coincident with receptor accumulation at the tips of growing spines. Notably, our findings do not support a strong requirement for presynaptic neurotransmitter release in spine formation. The abundance of spines was either mildly affected or unaffected across several mutant strains with severely impaired presynaptic function.
Transsynaptic NRX-1/Neurexin signaling stabilizes dendritic F-actin assemblies to promote postsynaptic maturation

Numerous studies across a variety of model systems have examined roles for neurexins in synapse development and function. However, no single consensus view of neurexin function has emerged. Instead, the function of neurexin at specific synapses are thought to be dictated by cellular and molecular context. Mammalian genomes encode three Nrxn genes that can each give rise to α-, β-, and γ-Nrxn isoforms. These isoforms share common intracellular and transmembrane regions but differ in their extracellular domains. α-Nrxn has six extracellular LNS domains interleaved with three EGF-like repeats. β-Nrxn has only one LNS domain, while γ-Nrxn lacks all identifiable extracellular domains. Conditional triple knockout of mouse neurexin indicated synapse loss in parvalbumin-positive prefrontal cortex interneurons, but not somatostatin-positive interneurons (L. Y. Chen et al., 2017). At mouse calyx synapses, conditional triple neurexin panneuronal knockout indicated a role in active zone organization and Ca$^{2+}$ and BK channel regulation, but no requirement in synapse formation or maintenance (Luo et al., 2020). The multi-faceted roles for neurexins at synapses likely emerge as a consequence of the many neurexin isoforms that are generated by alternative splicing in vertebrates, the complexity of their cellular expression, and the potential for these isoforms to selectively interact with distinct postsynaptic partners.

The C. elegans genome encodes a single neurexin gene, nrx-1 which generates two isoforms, a long α and a short γ isoform (Haklai-Topper et al., 2011). As is the case in other systems, C. elegans NRX-1 has roles in presynaptic organization, for example in calcium channel clustering at the active zone (Kurshan et al., 2018). Interestingly, the short γ-NRX-1 isoform that lacks identifiable ectodomains performs these roles (worms do not encode β-NRX-
1), but even in the absence of both isoforms, a level of presynaptic functionality is maintained, as indicated by the presence of evoked responses in muscle cells following presynaptic motor neuron stimulation (Philbrook et al., 2018). Our work here and our prior studies pointed toward the importance of NRX-1 at synapses between cholinergic and GABAergic motor neurons (Philbrook et al., 2018). Surprisingly, our time course studies here revealed that initial spine outgrowth and AChR clustering occurs normally in nrx-1 mutants. However, spines and AChRs in nrx-1 mutants become destabilized within hours and collapse, such that GABAergic dendrites are almost completely devoid of these postsynaptic components by L4 stage. Dendritic F-actin assemblies, that normally show discrete localization to spines, are also disorganized in nrx-1 mutants, and are diffusely distributed throughout GABAergic dendritic processes in nrx-1 mutants. We propose a model where presynaptic NRX-1 is required for the stabilization of dendritic spines and other postsynaptic structures in GABAergic dendrites. We suggest that presynaptic NRX-1 directs the organization of dendritic F-actin to stimulate maturation and stabilization of spines and postsynaptic receptor clusters. A similar form of F-actin based reorganization is a key step in presynaptic differentiation (Chia et al., 2012) (Chia et al., 2014).

Our work has parallels with recent studies in Xenopus (S. X. Chen et al., 2010) and Drosophila (Constance et al., 2018). In embryonic Xenopus brain, presynaptic β-neurexin stabilizes dendritic filopodia through an adhesive partnership with neuroligin to direct the development of dendritic arbors (S. X. Chen et al., 2010). Similarly, during fly metamorphosis, neurexin/neuroligin-based adhesion promotes the growth of neurite branches independently of synaptic activity (Constance et al., 2018). In our studies, NRX-1 stabilization of dendritic spines also occurs independently of synaptic activity and without involvement of neuroligin (Philbrook et al., 2018), suggesting interaction with an alternate binding partner that to date remains unidentified.
UNC-104/Kinesin 3-mediated synaptic delivery of NRX-1/Neurexin is an essential early step required for synapse maturation and stabilization

Efficient trafficking of synaptic and active zone cargoes to presynaptic terminals is essential for synapse formation and neuronal communication. Long-range axonal transport of synaptic vesicles and active zone proteins is carried out by anterograde kinesin and retrograde dynein motors. However, significant questions remain about which synaptic proteins are trafficked together and their dependence on specific motors. We found a specific requirement for the kinesin-3 UNC-104/KIF1A motor to deliver NRX-1 to presynaptic terminals. The anterograde transport of synaptic and dense core vesicles are also strongly dependent upon UNC-104 (Hall & Hedgecock, 1991) (Zahn et al., 2004). In contrast, several studies suggest that other AZ proteins exhibit weaker requirements for UNC-104 in their delivery. For example, ELKS-1 and other AZ proteins localize to C. elegans HSN synapses in an UNC-104-independent manner (Patel et al., 2006). Similarly, UNC-10/RIM properly concentrates within the nerve cord of unc-104(e1265) mutants, while synaptic vesicles accumulate in neuronal cell bodies (Koushika et al., 2001). Here, we found that NRX-1, UNC-10/RIM, ELKS-1/ELKS and CLA-1/piccolo all require UNC-104 for anterograde transport to cholinergic motor neuron synapses suggesting cell-specific functions for UNC-104 mediated delivery of active zone molecules.

Prior studies of rodent cultured neurons also suggested a requirement for the Kinesin-3 motor in neurexin transport, indicating that mechanisms for NRX-1 delivery to synapses are conserved (Neupert et al., 2015). The effects of a failure in delivery on synapse structure and function were not previously assessed. We found that dendritic spines collapse and AChR clusters disperse when UNC-104 delivery of synaptic cargoes is disrupted. Of the UNC-104 cargoes we analyzed, only mutation of nrx-1 produced significant disruption of dendritic spines. We therefore propose that NRX-1 is a key UNC-104 cargo required for spine stabilization. Our
live imaging points to similar rates of transport for synaptic vesicles and NRX-1. However, it remains unclear whether NRX-1 is delivered as a component of synaptic vesicles or may be segregated into a distinct vesicular population, such as synaptic vesicle protein transport vesicles (STVs). Our findings that mutations which severely impair synaptic vesicle release do not affect spine density point toward the latter possibility.

The prior identification of a temperature-sensitive allele of unc-104 allowed us to explore the temporal requirements for UNC-104 delivery of NRX-1. Importantly, we found that UNC-104 delivery was not solely required during early synaptogenesis but was critical throughout the developmental progression of the circuit toward maturity. These findings argue that ongoing delivery of presynaptic NRX-1 is required to maintain postsynaptic structure long after circuit assembly is complete. This raises interesting questions about the relationship between NRX-1 transport and synapse stability, perhaps suggesting that alterations in the rate of axonal NRX-1 transport may directly impact synaptic connectivity in mature animals. More broadly, our studies of the development and stabilization of synapses at dendritic spines in C. elegans provide a new view of the role of adhesive mechanisms in circuit connectivity and highlight a novel role for neurexin in the stabilization of mature synapses and dendritic spines.
Materials and Methods

Strains

All strains are N2 Bristol strain derivatives (wild type) and were maintained at room temperature (20-24°C) on nematode growth media plates (NGM) seeded with *E. coli* strain OP50. Transgenic strains were obtained by microinjection to achieve transformation (Mello et al., 1991) and identified using co-injection markers. Integrated lines were produced with X-ray irradiation and outcrossed to wild type/N2 Bristol eight times. Only hermaphrodites (L1-L4, young adults) were used in this study. A complete list of all strains used in this study is found in Supplemental File 1. Worms used for time course studies were staged by transferring embryos to seeded OP50 plates and transferring to 25°C (time point 0).

Molecular Biology

Plasmids were constructed using the two-slot Gateway Cloning system (Invitrogen) and confirmed by restriction digest and sequencing.

*Utrophin/F-actin reporter.* Sequence coding for mCherry was amplified from pDest-16 (and ligated into AgeI-HF/BspEI-digested PNYL183 (P*flp*-13::GFP::UtrCH, gift from Dong Yan’s laboratory) to make pCL87 (P*flp*-13::mCherry::UtrCH).

*TBA-1/tubulin reporter.* Sequence coding for GFP::TBA-1 was amplified from plasmid pYJ128 (gift from Kang Shen laboratory) and ligated into a destination vector to create pDest-173. pDest-173 was recombined with pENTR-5'-*flp*-13 to generate pDO61 (P*flp*-13::GFP::TBA-1).

*Golgi apparatus reporter.* pDest-161 (AMAN-2::GFP) was recombined with pENTR-3'-*flp*-13 to generate pDO47 (P*flp*-13::AMAN-2::GFP). pDest-161 was created by HindIII-HF digestion of Pgpa-4::AMAN::GFP (gift from Gert Jansen’s laboratory) and ligated with a destination vector.

*Mitochondria reporter.* pre-su9::GFP was isolated from Xmal/SphI-HF digested pDM1389 (gift from A.V. Maricq lab) and ligated into a destination vector to create pDest-91 (pre-su9::GFP). pDest-91 was recombined with pENTR-5'-*flp*-13 to generate pAP102 (P*flp*-13::pre-Su9::GFP).
**ER reporter.** Tag-RFP::TRAM-1 was amplified from plasmid pCT27 (gift from A.V. Maricq laboratory) and ligated to a KpnI-HF/NgoMIV digested destination vector to generate pDest-169. pDest-169 was recombined with pENTR-3'-flp-13 to generate pDO59 (Pflp-13::RFP::TRAM-1).

**Presynaptic vesicle reporter.** acr-5 promoter was amplified from plasmid pDM806 and ligated into pENTR-D-TOPO to generate pENTR-40. pENTR-40 was then recombined with pDest-5 to generate pAP264 (Pacr-5::SNB-1::GFP).

**B-type specific motor neuron reporter.** pDO45 (Pacr-5::GFP) was created by recombining pDest-94 (GFP) and pENTR-40 (acr-5 promoter).

**myr-GCamp6f.** myr-GCamp6f was amplified from plasmid pDest-164 and ligated into a destination vector to create pDest-180. pDest-180 was recombined with pENTR3'::flp-13 to create pDO69 (Pflp-13::SL2::myr-GCamp6f).

**CLA-1 reporter.** 3XnovoGFP::CLA-1e was isolated from SphI-HF/AgeI-HF digested pKP85 (gift from Peri Kurshan) and ligated with a destination vector to create pDest-307 pDest-307 was recombined with pENTR-3'::unc-17β to generate pSR58 (Punc-17β::3XnovoGFP::CLA-1e cDNA).

**unc-104 rescue constructs.** UNC-104cDNA::mCherry was isolated from SphI-HF/Nhel-HF digested pLL48 (gift from Kang Shen lab) and ligated with a destination vector to make pDest-308. pDest-308 was recombined with pENTR-unc-17β to generate pDO126 (Punc-17β::UNC-104cDNA::mCherry), pENTR-3'::unc-47 to generate pDO134 (Punc-47::UNC-104cDNA::mCherry), and pENTR-3'::myo-3 to generate pDO136 (Pmyo-3::UNC-104cDNA::mCherry).

**Punc-17β::NRX-1::GFP.** pAP112 was generated by recombining pENTR-3'::unc-17β with pDest-99 (NRX-1::GFP).

**ELKS-1::mCherry.** mCherry::ELKS-1cDNA was isolated from SphI-HF/Ascl digested pK043 (gift from Peri Kurshan laboratory) and ligated with a destination vector to create pDest-325. pDest-
325 was recombined with pENTR-3'-'unc-129 to generate pDO156 (\textit{Punc-129::ELKS-1::mCherry}).

\textit{UNC-10::GFP}. UNC-10::GFP was isolated from BamHI-HF/SbfI digested pRIM3 (gift from Michael Nonet laboratory) and ligated with a destination vector to generate pDest-304. pDest-304 was recombined with pENTR-3'-'unc-129 to generate pDO128 (\textit{Punc-129::UNC-10::GFP}).

\textbf{CRISPR/Cas-9 endogenously tagged \textit{NRX-1::GFP}}

Strain PHX3578 \textit{nrx-1(syb3578)} was generated in N2 animals by SunyBiotech. Linker sequence and GFP were inserted after exon 27 at the 3' end of C29A12.4a1/\textit{nrx-1}. Sequence flanking the GFP is below.

5': CGGGAAT\textbf{GGAGTCCGCAAAGAAAAGGATTTTAAGAGTGGTACGTA}aaggtaccgcgggccc
ggatccacccggtcgccaccatg

3': TCCATTTCCTTCAATCAAAACTCAATACAATGATGATTAAAAAATTCACTTTTGCTGCAAA

\textbf{PAM sequence is underlined}

\textbf{Synonymous mutations are bolded}

\textbf{Linker is in lower case letters}

\textit{nrx-1} gene is \textbf{CAPTIALIZED}

\textbf{Hydrogel solution}

Hydrogel was prepared as an aqueous solution of polyethylene glycol (20 % w/v) and the photoinitiator Irgacure (0.5% stock, 0.1% working concentration), and stored at 4°C in the dark.
**Confocal microscopy**

All strains were immobilized with sodium azide (0.3 M) on a 2% or 5% agarose pad. Images were obtained using an Olympus BX51WI spinning disk confocal equipped with a 63x objective.

For time course analyses, newly hatched larvae were transferred to a seeded OP50 plate and maintained at 25°C.

*Long term live imaging of dendritic spines and F-actin.* Nematodes were immobilized using 2 µL 50 mM muscimol in 10 µL 20% PEG hydrogel solution. Once paralyzed, hydrogel was curated using a handheld UV Transilluminator (312 nm, 3 minutes). Z-stacks were acquired every 5 minutes for at least one hour.

*Live imaging of synaptic vesicle and NRX-1 trafficking.* Nematodes were immobilized using 50 mM muscimol on a 5% agarose pad. Cholinergic commissures were imaged using Perkin Elmer spinning disk confocal equipped with a 63x objective using 100 ms exposure for 30 seconds.

**Confocal microscopy analysis**

All image analysis was conducted using ImageJ software (open source) within defined ROIs using intensity threshold values determined from control experiments. ROIs were located 15-30 µm anterior to DD1, DD2 or DD3 somata.

*Spine analysis.* Mature spines were quantified as protrusions from the dendrite >0.3 µm in length (measuring from the base to the tip of the protrusion). Spine density was defined as the number of spines/unit length within a selected ROI.

3D rendering was conducted using Imaris/bitplane 3D image analysis software. Morphological categories were determined based on criteria used in (Harris et al., 1992). The ImageJ line tool was used to measure the length and width of the spine 3D rendering. Based on these calculations, spines were placed into one of four morphological categories based on the
following criteria: Stubby: middle width > total length; Thin: middle width = tip width; Mushroom: tip width > middle width; Branched: > 1 spine head.

**Fluorescence intensity analysis**

*F-actin/spine and F-actin/tubulin.* The DD neurite was extracted using the line and line straighten functions. Total fluorescence intensity (utrophin/UtrCH, tubulin/TBA-1, ER/TRAM-1) on the dendritic shaft (line width 4) and below the dendritic shaft (line width 8) were measured. For soma fluorescence intensity, ROI was drawn around the perimeter of the soma.

**Synaptic marker and receptor cluster.** Background fluorescence was subtracted, and the number and size of synaptic/receptor puncta were measured using the ‘analyze particles’ function. Confocal montages were assembled using the ‘straighten to line’ function in a 50 µm region of the dorsal nerve cord.

**Calcium Imaging**

Animals were grown on plates seeded with OP50 containing 2.7 mM All-Trans Retinal (ATR). Plates were stored at 4°C under dark conditions and used within one week. Imaging was carried out using L4 animals immobilized in hydrogel (Burnett et al., 2018). Animals were transferred to 7.5 µL of the hydrogel mix placed on a silanized glass slide and covered with a glass slide. Hydrogel was cured using a handheld UV Transilluminator (312 nm, 3 minutes). Post-curing, the covering slide was removed and replaced with a coverslip. Imaging was carried out using a Yokogawa CSU-X1-A1N spinning disk confocal system (Perkin Elmer) equipped with EM-CCD camera (Hamamatsu, C9100-50) and 63X oil immersion objective. Chrismos photoactivation (~30 mW/cm²) was achieved using a TTL-controlled 625 nm light guide coupled LED (Mightex Systems). A 556 nm BrightLine single-edge short-pass dichroic beam splitter was positioned in the light path (Semrock) (Figure S1.3). Data were acquired at 10 Hz for 15 s using Volocity software and binned at 1×1 during acquisition. Analysis was performed using ImageJ. The DD
neurite process in each time series was extracted using the straighten function, background subtracted and photobleaching correction was carried out by fitting an exponential function to the data (CorrectBleach plugin). A smoothing function was applied to the data to enhance signal-to-noise. Individual spine ROIs were identified using the mCherry fluorescence. Post imaging processing, pre-stimulus baseline fluorescence ($F_0$) was calculated as the average of the data points in the first 4 s of the recording. Data was normalized to prestimulus baseline as $\Delta F/F_0$, where $\Delta F = F - F_0$. Peak $\Delta F/F_0$ was determined by fitting a Gaussian function to the $\Delta F/F_0$ time sequence using Multi peak 2.0 (Igor Pro, WaveMetrics). All collected data were analyzed, including failures (no response to stimulation). Control recordings were carried out in the absence of Retinal.

**Electron microscopy**

Staged L4 wild type (Pflp-13::ACR-12::GFP) hermaphrodites were subjected to high-pressure freeze fixation using a HPM10 high-pressure freezer. Hermaphrodites were slowly freeze substituted in 2% osmium tetroxide (OSO4), 0.1% uranyl acetate, 2% H2O in acetone fixative (Weimer, 2006) (Muller-Reichert et al., 2010). Samples were embedded in blocks, sectioned at 70 nm-thick serial sections and subsequently collected on copper slot grids. Samples were post-stained in 2.5% uranyl acetate in 70% methanol for 20 minutes and 1:6 lead citrate in H2O for 2 minutes at room temperature. Images were acquired using the Philips CM10 TEM. We identified and characterized 6 DD1 spines from a 35 µm region of the ventral head region where the neurite emerges from the cell body of DD1 GABAergic motor neuron.

**Temperature shift experiments**

Wild type and unc-104(ce782)ts animals eggs were raised at 13.5°C until L3 or L4 stage as indicated, then shifted to 25°C for 16-20 hrs prior to imaging. Animals were age-matched at the time of the temperature shift to account for developmental delays of unc-104(ce782)ts mutants.
References


Figure 1. *C. elegans* GABAergic DD neuron dendritic spines share characteristics with dendritic spines in vertebrates.

(A) Representative fluorescent images and cartoon representation of the ventral nerve cord/dendrite of a single L4 stage GABAergic DD (DD2) motor neuron with ventrally projecting dendritic spines. CB, cell body.

(B) Fluorescent image (top, inverted LUT) and 3D rendering (middle and insets) of DD dendritic spines from an animal expressing P*flp*-13::mCherry, shows diverse spine morphologies. DD dendritic spines share similar morphologies with mammalian dendritic spines: mushroom (6.02 ± 0.66%), branched (6.02 ± 0.66%), stubby (80.72 ± 3.09%), and thin (16.87 ± 1.21%). Arrowheads, spine insets. n=83 dendritic spines from 11 animals, measurements, percentage ± SD.

(C) Representative images of DD dendritic spines from animal co-expressing P*flp*-13::mCherry (labeling dendrite, red) and P*flp*-13::GFP::UtrCH (labeling F-actin, green). F-actin is highly enriched in DD dendritic spines.

(D) Scatterplot of F-actin/GFP::UtrCH fluorescence in dendritic shaft compared to dendritic spines, normalized to total fluorescence within the ROI. Bars indicate mean ± SEM. Student’s t-test, ***p<0.001, n=21 animals.

(E) Scatterplot of the percentage of spines with or without F-actin. Bars, mean ± SEM, n=21 animals.

(F) Fluorescent images of tubulin (P*flp*-13::TBA-1::GFP) (green) and actin (P*flp*-13::mCherry::UtrCH) (red) in DD neurons. Dendritic spines are highly enriched with F-actin while tubulin occupies the main dendritic process.
(G) Quantification of fluorescence ratio of actin/tubulin in the dendritic shaft versus dendritic spines. Student’s t-test, ****p<0.0001, n=31 spines from 6 animals.

(H) Calcium transients recorded from wild type GABAergic DD motor neuron dendrites expressing myrGCaMP6f (Pflp-13::myrGCaMP6f::SL2::mCherry) during stimulation of presynaptic cholinergic motor neurons (Pacr-2::Chrimson). Top, representative image showing dendritic spines identified by mCherry fluorescence (inverted LUT). Representative maximum intensity projection heat maps showing calcium fluxes immediately prior to stimulation (prestimulus, middle panel) and during stimulus (ON, bottom panel). Heat maps were generated by maximum intensity projection of myrGCaMP6f fluorescence from the first 4 seconds of recording (prestimulus) and the 5 s period of stimulation (red sharded bar in I). Numbering corresponds to representative traces in Figure 1I.

(I) Representative evoked responses for indicated dendritic spines in wild type animals (wild type 1, 2 and 3, Figure 1H), animals grown in the absence of retinal (-Retinal), and unc-17(e113) mutants. Calcium responses were continuously recorded at 10 Hz for 15 s. Prestimulus calcium fluxes were recorded for 5 s, followed by stimulation (625 nm, ~30 mW/cm²) of cholinergic motor neurons for 5 s. All data was normalized to prestimulus (ΔF/F₀). Red shaded bar indicates duration of stimulation. Pink traces indicate Gaussian fits to responses (ΔF/F₀) during stimulation.

(J) Scatter plot showing peak ΔF/F₀ of responses during stimulation in wild type or unc-17(e113) mutants. Bars, mean ± SEM. Statistical analysis, Student’s t-test, ****p<0.0001. n ≥ 10 animals.
Figure 2. Temporal order of molecular events during synapse and spine development.

(A) Fluorescent images of GABAergic dendrites at 12, 16, 20, 24, and 32 hours after hatch and at L4 stage (~42-50 hours post-hatch). Animals express a postsynaptic dendritic spine marker (Pflp-13::mCherry) with either presynaptic active zone (Punc-17β::GFPnovo2::CLA-1e) (top) or synaptic vesicle marker (Pacr-5::SNB-1::GFP) (bottom). Images are pseudo colored to indicate spines (magenta) and presynaptic CLA-1 or vesicle clusters (cyan).

(B) Fluorescent images of GABAergic dendrites at 12, 16, 20, 24, and 32 hours after hatch and L4 animals (~42-50 hours post-hatch). Animals express a postsynaptic dendritic spine marker (Pflp-13::mCherry) with either postsynaptic AChR (Punc-47::ACR-12::GFP, top) or postsynaptic actin (Pflp-13::GFP::UtrCH, bottom) markers. Images are pseudo colored to indicate spines (magenta) and postsynaptic AChR or F-actin (cyan). White asterisks (*) indicate images used in Figure 2C.

(C) Fluorescent images (inverted LUT) of GABAergic dendrites at 20 hours after hatch and L4 stage (~42-50 hours post-hatch) as indicated. Animals express a DD dendritic spine marker (Pflp-13::mCherry). Dashed red boxes indicate regions used in Figure 2B (asterisks).

(D-G) Quantification of spine number (purple) relative to clusters of active zone protein/CLA-1 (D), synaptic vesicle/SNB-1 (E), AChR/ACR-12 (F), and F-actin/UtrCH (G) (black) through development. Clusters of each of these markers are visible well prior to spine outgrowth. Blue shading indicates the approximate
period of DD remodeling. For this and subsequent figures, numbering indicates animals quantified at each timepoint for each genotype. Data points and bars indicate mean ± SEM.

(H) Fluorescent confocal images of DD1 AChR/ACR-12 (Pflip-13::ACR-12::GFP, magenta) and F-actin clusters (Pflip-13::mCherry::UtrCH, cyan) 24 hours after hatch. Dotted rectangle indicates high magnification image shown in 2I. Expanded view of DD1 dendrites 24 hours after hatch from Figure 2H.

(J) Fluorescent confocal images of DD1 AChR/ACR-12 (Pflip-13::ACR-12::GFP, magenta) and F-actin clusters (Pflip-13::mCherry::UtrCH, cyan) at L4 stage. Dotted rectangle indicates high magnification image shown in 2K.

(K) Expanded view of DD1 dendrites 24 hours after hatch from Figure 2J.

(L) Cartoon summary of the order of events during development of synapses onto DD neurons. In early development, presynaptic vesicles (SNB-1) and active zone protein (CLA-1) localize along presynaptic terminals while postsynaptic F-actin and postsynaptic receptors cluster along the postsynaptic process. Following their formation, dendritic spines begin to extend. Spines continue to elongate throughout development and into the mature circuit (L4).
Table 1. Analysis of dendritic spines and cholinergic receptors in mutant strains with altered synaptic activity.

Dendritic spine number, spine length, ACR-12/AChR cluster number, and AChR cluster size at L4 stage normalized to the same measurements from wild type are shown (mean ± SEM) for each genotype (ACR-2/nAChR UNC-2/CaV2α, UNC-13/MUNC-13, UNC-17/VAChT, UNC-18/MUNC-18, UNC-31/CAPS). Mutations that affected synaptic activity had little effect on the density of spines or AChR clusters, but had variable effects on spine length and AChR cluster size. One-way AVOVA, Dunnett’s multiple comparisons test, *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Spines</th>
<th>AChR/receptors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normalized spine number</td>
<td>Normalized ACR-12 cluster</td>
</tr>
<tr>
<td></td>
<td>± SEM (%)</td>
<td>number ± SEM (%)</td>
</tr>
<tr>
<td></td>
<td>N</td>
<td>P value</td>
</tr>
<tr>
<td>wild type</td>
<td>100.0 ± 4.1</td>
<td>58 ns</td>
</tr>
<tr>
<td>acr-2(ok1887)</td>
<td>86.2 ± 9.1</td>
<td>24 ns</td>
</tr>
<tr>
<td>acr-2(n2420)gf</td>
<td>52.9 ± 5.9</td>
<td>26 ****</td>
</tr>
<tr>
<td>unc-2(e55)</td>
<td>81.8 ± 7.6</td>
<td>29 ns</td>
</tr>
<tr>
<td>unc-2(zf35)gf</td>
<td>81.7 ± 7.2</td>
<td>22 ns</td>
</tr>
<tr>
<td>unc-13(e51)</td>
<td>62.1 ± 8.1</td>
<td>18 **</td>
</tr>
<tr>
<td>unc-17(e113)</td>
<td>104.5 ± 6.8</td>
<td>12 ns</td>
</tr>
<tr>
<td>unc-18(e234)</td>
<td>76.9 ± 10.0</td>
<td>11 ns</td>
</tr>
<tr>
<td>unc-31(e169)</td>
<td>97.6 ± 5.7</td>
<td>22 ns</td>
</tr>
</tbody>
</table>
Figure 3. Dendritic spines and postsynaptic components form initially but subsequently collapse without the synaptic adhesion protein, NRX-1.

(A) Fluorescent confocal images of dendritic spines (Pflp-13::mCherry, magenta) and cholinergic expression of NRX-1::GFP (Punc-17/β::NRX-1::GFP, cyan) at 16, 20, and 24 hours after hatch and L4 stage (~42-50 hours post-hatch). Presynaptic clusters of NRX-1::GFP are visible by 16 hours after hatch and increase in size until roughly 24 hours after hatch.

(B) Quantification of the average number of spines (purple) and NRX-1 clusters (black) at the time points indicated. Blue shading indicates the approximate period of DD remodeling. Numbers indicate the number of animals quantified at each timepoint. Data points indicate mean ± SEM.

(A) Fluorescent images (inverted LUT) of DD1 dendritic spines (Pflp-13::myrGFP) in wild type (left) and nrx-1(wy778) (right) animals at 16, 18, 20, and 24 hours after hatch and at L4 stage. Immature dendritic spines clusters form initially but then collapse in nrx-1 mutants.

(B) Quantification of the percentage of wild type (black) and nrx-1(wy778) (purple) animals with 2 or more DD dendritic spines at the timepoints indicated. Numbers indicate the number of animals quantified at each timepoint.

(C) Fluorescent images of ACR-12/AChR clusters (Pflp-13::ACR-12::GFP) in wild type (left) and nrx-1(wy778) mutants (right) 16, 20, and 24 hours after hatch and at L4 stage (~42-50 hours post-hatch). Immature AChR clusters initially form but then disperse in nrx-1 mutants.
(D) Quantification of the number of ACR-12::GFP clusters in wild type (black) and \textit{nrx-1}(wy778) mutants (purple). Two-way ANOVA, Sidak’s multiple comparisons, \(*p<0.05, **p<0.01, ***p<0.001.\)

(E) Fluorescent confocal images of dendritic spines (\textit{Pfip-13::mCherry}) and F-actin (\textit{Pfip-13::GFP::UtrCH}) in wild type and \textit{nrx-1}(wy778) animals at 16 hours after hatch and L4 stage.
A NRX-1::GFP (knock-in)

B ACh::NRX-1::GFP

C % of control (number of puncta)

D ACh::CLA-1::GFP

E ACh::UNC-10::GFP

F ACh::ELKS-1::mCherry

G Spines (GABA::myrGFP)

H AChR (GABA::ACR-12::GFP)

I # rescued lines/total

J Spines (GABA::mCherry or GABA::myrGFP)
Figure 4. UNC-104/KIF1A motor-dependent transport of NRX-1 to presynaptic terminals is required for stabilization of postsynaptic structures.

(A) Representative confocal images of NRX-1::GFP (endogenous nrx-1 knock-in) from dorsal cords (inverted LUT) of wild type and unc-104(e1265) mutants. Images on each line are from different animals (5 shown for each genotype). Mutation of unc-104 impairs axonal NRX-1::GFP localization.

(B) Representative confocal images of cholinergic NRX-1::GFP (Punc-129::NRX-1::GFP, inverted LUT) from the dorsal nerve cord of wild type, unc-104(e1265) and unc-104(e1265) mutants rescued by cholinergic expression of wildtype unc-104 (Punc-17β). Images on each line are from different animals (5 shown for each genotype).

(C) Scatterplot of the average number of NRX-1::GFP (endogenous or transgenic), CLA-1::GFP, UNC-10::GFP, or ELKS-1::mCherry clusters per 50 µm of dorsal nerve cord in unc-104(e1265) mutants or rescue as indicated. Values are normalized to their respective controls. Red dotted line indicates 50% of wild type values. Bars indicate mean ± SEM. Student's t-test, ****p<0.0001. n ≥ 11 animals. Raw values with respective controls are shown in Figure S4.6.

(D) Representative confocal images of cholinergic CLA-1 (Punc-17β::CLA-1::GFP, inverted LUT) from the dorsal nerve cord of wild type or unc-104(e1265) mutants. Images on each line are from different animals (5 shown for each genotype).

(E) Representative confocal images of cholinergic UNC-10 (Punc-129::UNC-10::GFP, inverted LUT) from dorsal nerve cords of wild type and unc-104(e1265) mutants. Images on each line are from different animals (5 shown for each genotype).
(F) Representative confocal images of cholinergic ELKS-1 (Punc-129::ELKS-1::mCherry, inverted LUT) from dorsal nerve cords of wild type and unc-104(e1265) mutants. Images on each line are from different animals (5 shown for each genotype).

(G) Fluorescent images (inverted LUT) of DD dendritic spines (Pflp-13::myrGFP) from wild type, unc-104(e1265), and unc-104(e1265) mutants rescued by cholinergic expression of wild type unc-104.

(H) Fluorescent images of AChR/ACR-12 clusters in DD dendrites (Pflp-13::ACR-12::GFP) of wild type, unc-104(e1265), and unc-104(e1265) mutants rescued by cholinergic expression of wild type unc-104.

(I) Scatterplot showing quantification of DD dendritic spines per 15 µm (Pflp-13::myrGFP) (purple) and DD cholinergic receptor clusters (Pflp-13::ACR-12::GFP) (green) in wild type, unc-104(e1265), and indicated cell-specific rescue lines. Inset, number of rescuing lines/total transgenic lines tested for each rescue construct. Bars, mean ± SEM. One-way ANOVA, Dunnett’s multiple comparisons test, ****p<0.0001. n ≥ 10 animals.

(J) Fluorescent images of DD dendritic spines (Pflp-13::myrGFP or Pflp-13::mCherry) from wild type, nrx-1(wy778), cla-1(ok937), unc-10(md1117), or elks-1(ok2762) mutants. Only mutation of nrx-1 affects dendritic spines.
Figure 5. UNC-104/KIF1A transport of NRX-1 is required in the mature circuit to maintain postsynaptic structure.

(A) Cartoon depiction of experimental timeline. Animals were grown at 13.5°C until L4 stage (approx. 144 hours in unc-104(ce782)ts mutants, 120 hours in wild type.
animals) before shifting animals to the restrictive temperature of 25°C for 16-20 hours and imaging.

(B) Line scan fluorescent images of NRX-1::GFP (knockin to endogenous locus, inverted LUT) from dorsal cords of wild type and unc-104(ce782) mutants grown continuously at the permissive temperature (13.5°C) (middle panel) or shifted to the restrictive temperature (25°C) at L4 stage for 16-20 hours before imaging (bottom panel). Each line scan indicates an individual animal.

(C) Quantification of NRX-1::GFP clusters along a 50 µm region of the dorsal nerve cord in wild type and unc-104(ce782)ts animals. Two-way ANOVA, Tukey’s multiple comparisons test, ns, not significant, *p<0.05, n ≥ 11 animals. Data points indicate mean ± SEM.

(D) Fluorescent images of DD spines in wild type and unc-104(ce782)ts animals grown continuously at the permissive temperature (13.5°C) (middle panel) or shifted to the restrictive temperature (25°C) at L4 stage for 16-20 hours before imaging.

(E) Quantification of the number of spines per 15 µm in wild type and unc-104(ce782)ts animals. Two-Way ANOVA, Tukey’s multiple comparisons test, ns, not significant, **p<0.001. n ≥ 13 animals. Data points indicate mean ± SEM.
Figure S1.1 Ultrastructure of DD dendritic spines.

(A) Cartoon depiction of the anterior serial cross-sections used for electron microscopy studies of the ventral nerve cord, focusing on the dendrite of the DD1 neuron.

(B) Serial cross-sections (225/491, 226/491, 227/491, 228/491) of the ventral nerve cord. Teal indicates DD1 dendrite and dendritic spines (226/491 and 227/491) dipping into the ventral nerve cord to meet presynaptic cholinergic terminals (presumably VA/VB neurons, magenta). Asterisks indicates mitochondria within the dendritic shaft, see Figure S1.4. Scale bar, 100 nm.

(C) Serial cross-sections (443/597, 444/597, 445/597, 446/597) of the ventral nerve cord of John White’s N2U series (J. G. White et al., 1976). Teal indicates DD1 dendrite and dendritic spines dipping into the ventral nerve cord to meet presynaptic cholinergic terminals (magenta) (sections 444-446). Asterisks indicates mitochondria within the dendritic shaft, see Figure S1.4.

(D) Ventral nerve cord electron micrographs. Dendritic spines (teal) have varied morphology and sizes. Two representative micrographs from serial sections in which the entire extent of the spine is visible are shown. Asterisk indicates mitochondrion within the dendritic shaft. Also see Figure S1.4B. Scale bar, 100 nm.
**Figure S1.2. Microtubules localize at the base of spines.**

Fluorescent images of tubulin (Pflip-13::TBA-1::GFP) and F-actin (Pflip-13::mCherry::UtrCH) in DD neurons. White arrows indicate dendritic regions where tubulin dips into the base of a dendritic spine.
**Figure S1.3. Cartoon illustration of calcium imaging recording platform.**

Imaging was carried out using a Yokogawa CSU-X1-A1N spinning disk confocal system (Perkin Elmer) equipped with EM-CCD camera (Hamamatsu, C9100-50) and 63X oil immersion objective. Chrimson photoactivation (~30 mW/cm²) was achieved using a TTL-controlled 625 nm light guide coupled LED (Mightex Systems), permitting illumination of the entire immobilized animal, while simultaneously recording GCaMP6f fluorescence (excitation 488 nm, emission 525 nm). 625 nm light was prevented from reaching the
camera using a 556 nm edge BrightLine single-edge short-pass dichroic beam splitter positioned in the light path (Semrock).
Supplementary Figure 1.4 Examination fluorescent organelle markers in DD dendrites and dendritic spines.

A) Golgi

B) Mitochondria

C) TRAM-1::RFP fluorescence intensity / total fluorescence (AU)

D) Rough ER

E) GABA::mCherry

pre-Su9::GFP fluorescence intensity / total fluorescence (AU)

Mitochondria

Rough ER

TRAM-1::RFP fluorescence intensity / total fluorescence (AU)

Golgi

merge

merge

merge

merge

merge

merge

**

***

****
**Figure S1.4. Examination of fluorescent organelle markers and F-actin interacting proteins in DD dendrites and dendritic spines.**

(A) Fluorescent images of DD1 dendritic spines (Pflp-13::mCherry) and Golgi marker (Pflp-13::AMAN-2::GFP) in DD neurons. Golgi apparatus is primarily localized to the DD1 soma. Dotted white line traces the neuronal cell body.

(B) Fluorescent images of DD1 dendritic spines (Pflp-13::mCherry) and mitochondria (Pflp-13::pre-Su9::GFP). Mitochondria are localized within the main dendritic process of DD neurons near spines.

(C) Quantification of mitochondria fluorescence intensity in dendritic shaft and dendritic spines. Bars, mean ± SEM. Student’s t-test, ****p<0.0001.

(D) Fluorescent images of DD1 dendritic spines (Pflp-13::myrGFP) and rough endoplasmic reticulum (ER) (Pflp-13::TRAM::RFP). Dotted white line traces the neuronal cell body. ER is primarily localized within the main dendritic process of DD neurons and occasionally at the spine base.

(E) Quantification of rough ER fluorescence intensity (Pflp-13::RFP::TRAM-1) in dendritic shaft and dendritic spines. Bars, mean ± SEM. Student's t-test, ***p<0.001. n=9 animals.
**Figure S2.1. Development of post-embryonic born ventral cholinergic motor neurons.**

(A) Fluorescent images of B-type cholinergic neurons (DB/VB) (Pacr-5::GFP) (green) and DD GABAergic neurons (Pflp-13::mCherry) (red) at 16 and 24 hours after hatch. VB cholinergic neurons are born post-embryonically in an anterior to posterior order (blue). Note that at 16 hours after hatch Pacr-5::GFP fluorescence indicating VB1 and VB2 cell bodies is visible. By 24 hours after hatch Pacr-5::GFP fluorescence indicating VB3 is visible. White dotted circles outline the neuronal cell bodies.

(B) Fluorescent images of the anterior ventral nerve cord in animals co-expressing the synaptic vesicle marker Pacr-5::SNB-1::GFP in B-type cholinergic neurons (DB/VB) (green) with Pflp-13::mCherry labeling DD GABAergic neurons (red) at 12, 16, and 24 hours after hatch. At 12 hours, little SNB-1::GFP fluorescence is visible in the ventral nerve cord. Embryonic born, dorsally directed B-type (DB) cholinergic motor neurons predominantly make synaptic contacts in the dorsal nerve cord. Ventrally-directed, B-type (VB) cholinergic motor neurons are born post-embryonically and have not yet completed their maturation at this time. SNB-1::GFP fluorescence in the ventral nerve cord is faintly visible by 16 hrs after hatch and becomes more prominent by 24 hrs after hatch, coincident with maturation of VB motor neurons. White dotted circles indicate outlines of neuronal cell bodies. Yellow arrows indicated presynaptic vesicle clusters, SNB-1.
(C) Cartoon representation of the developmental timing of post-embryonic born ventral cholinergic motor neurons, spine outgrowth, and cholinergic synaptic vesicle localization.
Figure S2.2. Spine outgrowth progresses anteriorly to posteriorly.

(A and B) Fluorescent images (inverted LUT) of DD1 and DD2 dendritic spines 24 hours after hatch (A) and at L4 (~42-50 hours after hatch) stage (B). Animals express Pflp-13::mCherry to label dendritic spines. Red dashed rectangle indicates inset anterior
to DD1 soma. Blue dashed rectangle indicates inset anterior to DD2 soma. Note that DD1 spines form prior to DD2 spines.

(C) Quantification of the length of growing dendritic spines at 24 and 32 hours after hatch and at L4 (~42-50 hours after hatch) stage. Data points indicate mean ± SEM.
**Figure S2.3 LEV-10 localizes to putative postsynaptic sites prior to spine formation.**

(A) Fluorescent confocal images showing dendritic spines (magenta) and the postsynaptic CUB transmembrane domain LEV-10 (cyan). Animals express Pflip-13::mCherry with DD neuron specific LEV-10::GFP using a strategy for cell-specific labeling of endogenous LEV-10 (NATF) (He et al., 2019).

(B) Quantification of the number of DD dendritic spines and LEV-10 clusters at 16, 20, and 24 hours after hatch, and L4 stage (~42-50 hours post-hatch).
Figure S2.4 Actin dynamics during development.

(A,C,E) Line scans displaying relative fluorescence intensity of F-actin (Pflp-13::GFP::UtrCH) at 16 hours post-hatch (A), 24 hours post-hatch (C), and L4 (~42-50 hours post-hatch) animals (E). Each color indicates a line scan of fluorescence intensity for the same DD dendrite ROI acquired at 5-minute intervals. Note the variable distribution of fluorescence intensities across line scans from images acquired near 16 hours after hatch compared to later time points, indicating increased F-actin dynamics during early development.

(B,D,F) Confocal images (inverted LUT) showing Pflp-13::GFP::UtrCH fluorescence (labeling F-actin) in the DD dendrite of 16 (B), or 24 (D) hours after hatch, or at L4 stage (~42-50 hours post-hatch) (F). For each, sequences of fluorescent images separated by 30 minutes are shown.
**Video 1:** Confocal live imaging video of F-actin dynamics (labeled by Pflp-13::GFP::UtrCH) in the DD1 dendrite of wild type at 16 hours after hatch. For display, movies are shown at 3 fps. Images were acquired every 5 minutes.

**Video 2:** Confocal live imaging video of F-actin dynamics (labeled by Pflp-13::GFP::UtrCH) in the DD1 dendrite of wild type at L4 stage. For display, movies are shown at 3 fps. Images were acquired every 5 minutes.
**Figure S3.1. Spines and AChR localize independently of presynaptic activity.**

(A,B) Fluorescent confocal images (inverted LUT) of DD1 dendritic spines (*Pflip-13::*mCherry (A) or *Pflip-13::*myrGFP (B)) in L4 stage wild type and selected mutant strains where synaptic activity is affected.

(C) Fluorescent confocal images of AChR clusters (*Pflip-13::*ACR-12::*GFP) in the DD dendrite of L4 stage wild type and selected mutant strains where synaptic activity is affected.
Figure S3.2. Dendritic calcium responses are severely reduced in nrx-1 mutants.

Scatter plot showing peak $\Delta F/F_0$ responses measured from GABAergic DD motor neuron dendrites during a 5s period cholinergic photostimulation stimulation in wild type, acr-16(ok789); acr-12(ok367), nrx-1(wy778), nrx-1(nu485), and nrx-1(ok1649) mutants. All genotypes co-express Pflp-13::myrGCaMP6f::SL2::mCherry for measurement of dendritic calcium responses with Pacr-2::Chrimson for cholinergic neuron depolarization. Bars indicate mean ± SEM. One-way ANOVA, Dunnett’s multiple comparisons, ****$p<0.0001$. Wild type control is the same as Figure 1J. n ≥ 10 animals.
Figure S3.3. Spine and AChR localization in developing wild type and \textit{nrx-1(wy778)} mutants.

(A) Quantification of the number of DD spines/15 \( \mu \text{m} \) in wild type (black) and \textit{nrx-1(wy778)} (purple) animals 16, 20, and 24 hours after hatch and at L4 stage (~42-50 hours post-hatch). Two-way ANOVA, Sidak’s multiple comparisons test, **\( p<0.01 \), ****\( p<0.0001 \). Data points indicate mean ± SEM. Numbers indicate animals quantified for each timepoint.

(B) Quantification of ACR-12::GFP fluorescence intensity in wild type (black) and \textit{nrx-1(wy778)} (purple) animals at 16, 20, and 24 hours after hatch and L4 stage (~42-
50 hours post-hatch). Two-way ANOVA, Sidak’s multiple comparisons test, 

**p<0.01, ***p<0.001. Data points indicate mean ± SEM.
Figure S3.4. Spine dynamics during development in wild type and \textit{nrx-1(wy778)} mutants.

(A) Quantification of the percentage of spine events in wild type and \textit{nrx-1(wy778)} animals at 16-20 hours after hatch, 21-24 hours after hatch, and at L4 compared to the total number of events. An event is defined as an incidence of spine destabilization/pruning, elongation/formation, or a stable spine during an entire live imaging recording. Z-stacks of DD1 dendrites were acquired every 5 minutes for
60-160 minutes. Numbers indicates number of events for each genotype at each timepoint.

(B) Fluorescent image (inverted LUT) of DD dendritic spines at timepoint zero in a wild type animal 16-20 hours after hatch. Dashed red box indicates region shown in Figure S3.4C.

(C) Fluorescent images (inverted LUT) of an individual wild type DD dendritic spine (Pflp-13::mCherry) at 0, 45, and 70 minute timepoints during live imaging of area indicated by red box in Figure S3.4B. Red dotted line outlines spine dynamics.

(D) Fluorescent image (inverted LUT) of nrx-1(wy778) mutant DD1 dendritic spines (Pflp-13::mCherry) at timepoint zero acquired 16-20 hours after hatch. Dashed red box indicates region shown in Figure S3.4E.

(E) Fluorescent images (inverted LUT) of an individual nrx-1(wy778) mutant DD dendritic spine (Pflp-13::mCherry) at 90, 95, and 100 minute timepoints during live imaging of area indicated by red box in Figure S3.4D. Red dotted lines outline spine dynamics.

(F) Fluorescent image (inverted LUT) of wild type DD dendritic spines (Pflp-13::mCherry) at timepoint zero at L4 stage. Dashed red box indicates region shown in Figure S3.4G.

(G) Fluorescent images (inverted LUT) of an individual wild type DD dendritic spine (Pflp-13::mCherry) at 0, 30, and 65 minute timepoints during live imaging of the area indicated by red box in Figure S3.4F.
**Video 3.** Wild type DD dendritic spines (Pflp-13::myrGFP) 16 hours after hatch. A z-stack was acquired every 5 minutes for at least 1 hr of imaging duration. For display, movies are shown at 3 fps

**Video 4.** *nrx-1(wy778)* mutant DD dendritic spines (Pflp-13::myrGFP) 16 hours after hatch. A z-stack was acquired every 5 minutes for at least 1 hr of imaging duration. For display, movies are shown at 3 fps

**Video 5.** Wild type DD dendritic spines (Pflp-13::myrGFP) at L4 stage. A z-stack was acquired every 5 minutes for at least 1 hr of imaging duration. For display, movies are shown at 3 fps
Figure S4.1. Mutation of *unc-104* impairs synaptic vesicle delivery to cholinergic axons in the dorsal nerve cord.

(A) Fluorescent images of cholinergic synaptic vesicles (Pacr-2::mCherry::RAB-3) in the dorsal nerve cord of young adult wild type and *unc-104(e1265)* mutants. Images on each line are from different animals (4 are shown for each genotype).
(B) Quantification of cholinergic synaptic vesicles (Pacr-2::mCherry::RAB-3) per 50 µm of the dorsal nerve cords of wild type and unca-104(e1265) mutants. Student’s t-test, ****p<0.0001. Bars, mean ± SEM.

(C) Fluorescent images of the soma in wild type and unca-104(e1265) mutants expressing cholinergic vesicle reporter (Pacr-2::mCherry::RAB-3). Dotted white lines outline the cell body.

(D) Quantification of soma cholinergic synaptic vesicle fluorescence intensity (AU) (Pacr-2::mCherry::RAB-3) of wild type and unca-104(e1265) mutants. Student’s t-test, ****p<0.0001. Bars, mean ± SEM.
Figure S4.2. NRX-1 colocalizes with UNC-104 in cholinergic axon terminals.

(A) Line scans depicting relative fluorescent intensity of NRX-1::GFP (green) and UNC-104::mCherry (red) for a 50 μm region of the dorsal nerve cord. Gray dotted rectangles indicate corresponding puncta in Figure S4.2B.

(B) Fluorescent images of the dorsal nerve cord in an adult animal expressing NRX-1::GFP (Punc-129::NRX-1::GFP) and UNC-104::mCherry (Punc-129::UNC-104::mCherry). Colocalization is indicated by white carets.
NRX-1::GFP localizes within neuronal processes of the nerve ring and shows punctate localization in processes of the ventral and dorsal nerve cords of L4 stage worms.

**Figure S4.3. NRX-1 endogenously tagged with GFP localizes to neuronal processes.**
Figure S4.4. NRX-1 localization to cholinergic axons is decreased in *unc-104* mutants.

(A) Quantification of NRX-1::GFP (endogenous knock-in) clusters in a 50 µm region of the dorsal nerve cord in wild type and *unc-104(e1265)* animals. Bars, mean ± SEM. Student’s t-test, ****p<0.0001. n ≥ 11 animals.

(B) Quantification of NRX-1::GFP (endogenous knock-in) axon fluorescence intensity in a 50 µm region of the dorsal nerve cord. Bars, mean ± SEM. Student’s t-test, ****p<0.0001. n ≥ 11 animals.
(C) Fluorescent images of NRX-1::GFP (endogenous knock-in) in somas of wild type and unc-104(e1265) mutants. Dotted white lines outline the neuronal cell body.

(D) Quantification of NRX-1::GFP soma fluorescence intensity. Bars, mean ± SEM. Student’s test, ****p<0.001.

(E) Quantification of the number of NRX-1::GFP clusters in a 50 μm region of the dorsal nerve cord in wild type, unc-104(e1265), and unc-104(e1265) mutants rescued with cholinergic expression of wild type unc-104. Bars, mean ± SEM. One-way ANOVA, Dunnett’s multiple comparisons test, ****p<0.0001. n ≥ 11 animals.

(F) Quantification of NRX-1::GFP fluorescence intensity in cholinergic axons of a 50 μm region of the dorsal nerve cord. Bars, mean ± SEM. One-way ANOVA, Dunnett’s multiple comparisons test, ****p<0.0001. n ≥ 11 animals.

(G) Fluorescent images of NRX-1::GFP in cholinergic somata of wild type, unc-104(e1265) and unc-104(e1265) mutants rescued with cholinergic expression of wild type unc-104. Dotted white lines outline the neuronal cell body.

(H) Quantification of NRX-1::GFP fluorescence intensity in cholinergic somata. Bars, mean ± SEM. One-way ANOVA, Dunnett’s multiple comparisons test, ****p<0.001.
Figure S4.5. NRX-1 localization in *unc-116*/KIF5A/C mutants.

(A) Fluorescent images (inverted LUT) of NRX-1::GFP (*Punc-129::NRX-1::GFP*) from the dorsal nerve cord of wild type and *unc-116(e2310)* mutants. Images on each line are from different animals (5 are shown for each genotype).

(B) Quantification of the number of NRX-1 clusters (*Punc-129::NRX-1::GFP*) in a 50 μm region of the dorsal nerve cord of wild type and *unc-116(e2310)* animals. Bars, mean ± SEM. Student’s t-test, ***p<0.001. n ≥ 14 animals.

(C) Fluorescent images of NRX-1 (*Punc-129::NRX-1::GFP*) from cholinergic neuron somata of wild type and *unc-116(e2310)* mutants. Dotted white lines outline cell bodies.

(D) Quantification of fluorescence intensity of NRX-1 (*Punc-129::NRX-1::GFP*) from cholinergic somas of wild type and *unc-116(e2310)* mutants. Bars, mean ± SEM. Student’s t-test, ns, not significant.
Figure S4.6. NRX-1 and SNB-1 have similar anterograde and retrograde velocities.

(A, B) Kymographs of synaptic vesicle (Punc-129::SNB-1::GFP) (A) and NRX-1 (Punc-129::NRX-1::GFP) (B) trafficking events recorded from cholinergic neuron commissures.
(C) Quantification of the anterograde velocity ($\mu$m/second) of SNB-1::GFP (blue) and NRX-1::GFP (orange) along the axonal commissure. ns, not significant. $n \geq 10$ animals for all panels.

(D) Quantification of the retrograde velocity ($\mu$m/second) of SNB-1::GFP (blue) and NRX-1::GFP (orange) along the axonal commissure. ns, not significant.

(E) Quantification of the total number of SNB-1::GFP (blue) and NRX-1::GFP (orange) trafficking events binned into retrograde and anterograde directions. Note that NRX-1 trafficking events occur significantly less frequently than SNB-1 events.
**Video 6:** NRX-1::GFP trafficking in a cholinergic commissure of a young adult animal (Punc-129::NRX-1::GFP). Videos were recorded at 100 ms exposure for 30 seconds. For display, movie is shown at 25 fps.

**Video 7:** SNB-1::GFP trafficking in a cholinergic commissure of a young adult animal (Punc-129::SNB-1::GFP). Videos were recorded at 100 ms exposure for 30 seconds. For display, movie is shown at 25 fps.
Figure S4.7 Active zone protein localization in nrx-1(wy778) and unc-104(e1265) mutants.

A) Ach::CLA-1::GFP

B) Ach::CLA-1::GFP fluorescence intensity (AU)

C) Ach::CLA-1::GFP fluorescence intensity (AU)

D) Ach::UNC-10::GFP

E) Ach::UNC-10::GFP fluorescence intensity (AU)

F) Ach::UNC-10::GFP fluorescence intensity (AU)

G) Ach::ELKS-1::mCherry

H) Ach::ELKS-1::mCherry fluorescence intensity (AU)

I) Ach::ELKS-1::mCherry fluorescence intensity (AU)
Figure S4.7. Active zone protein localization in *nrx-1(wy778)* and *unc-104(e1265)* mutants.

(A) Line scan fluorescent images (inverted, LUT) of active zone protein CLA-1 (*Punc-17β::CLA-1-1::GFP*, inverted) from cholinergic axons of dorsal nerve cords in wild type, *nrx-1(wy778)*, and *unc-104(e1264)* mutants. Images on each line are from different animals (5 are shown for each genotype). These data correspond to quantification in Figure 4C.

(B) Scatterplot of CLA-1::GFP puncta in a 50 µm region of the dorsal nerve cord in wild type, *nrx-1(wy778)* and *unc-104(e1265)* animals. One-way ANOVA, Dunnett's multiple comparisons test, ****p<0.0001. Bars, mean ± SEM.

(C) Scatterplot of CLA-1::GFP fluorescence intensity in a 50 µm region of the dorsal nerve cord in wild type, *nrx-1(wy778)* and *unc-104(e1265)* animals. One-way ANOVA, Dunnett’s multiple comparisons test **p<0.01, ****p<0.0001. Bars, mean ± SEM.

(D) Line scan fluorescent images (inverted, LUT) of active zone protein UNC-10 (*Punc-129::UNC-10::GFP*) from cholinergic axons of dorsal nerve cords in wild type, *nrx-1(wy778)* and *unc-104(e1265)* mutants. Images on each line are from different animals (5 are shown for each genotype). These data correspond to quantification in Figure 4C.

(E) Quantification of UNC-10::GFP puncta in a 50 µm region of the dorsal nerve cord in wild type, *nrx-1(wy778)* and *unc-104(e1265)* animals. One-way ANOVA, Dunnett's multiple comparisons test, **p<0.001, ****p<0.0001. Bars, mean ± SEM.
(F) Quantification of UNC-10::GFP fluorescence intensity in a 50 μm region of the dorsal nerve cord in wild type, \textit{nrx-1}(wy778) and \textit{unc-104}(e1265) mutants. One-way ANOVA, Dunnett’s multiple comparisons test, ****p<0.0001. Bars, mean ± SEM.

(G) Line scan fluorescent images (inverted, LUT) of active zone protein ELKS-1 (\textit{Punc}-129::ELKS-1::mCherry, inverted) from cholinergic axons of dorsal nerve cords in wild type, \textit{nrx-1}(wy778) and \textit{unc-104}(e1265) mutants. Images on each line are from different animals (5 are shown for each genotype). These data correspond to quantification in Figure 4C.

(H) Quantification of ELKS-1::mCherry puncta per 50 μm of the dorsal nerve cord in indicated genotypes. One-way AVOVA, Dunnett’s multiple comparisons test, ****p<0.0001, ns, not significant. Bars, mean ± SEM.

(I) Quantification of ELKS-1::mCherry fluorescence intensity in a 50 μm region of the dorsal nerve cords of wild type, \textit{nrx-1}(wy778) and \textit{unc-104}(e1265) mutants. One-way ANOVA, Dunnett’s multiple comparisons test, ***p<0.001, ns, not significant. Bars, mean ± SEM.
Figure S5.1 Shift to restrictive temperature during L3 in wild type and \textit{unc-104(ce782)ts} animals.

(A) Cartoon depiction of experimental timeline. Animals were grown at 13.5°C until L3 stage (~approx. 120 hours in \textit{unc-104(ce782)} mutants, 96 hours in wild type
animals) before shifting animals to their restrictive temperature of 25°C for 16-20 hours and imaging.

(B) Fluorescent images of DD spines from wild type (top) or unc-104(ce782)ts animals grown continuously at the permissive temperature (13.5°C) (middle panel) or shifted to the restrictive temperature (25°C) for 16-20 hours before imaging.

(C) Quantification of DD spines per 15 µm from wild type and unc-104(ce782)ts mutants. Two-way ANOVA, Tukey’s multiple comparisons test, ****p<0.0001, n ≥ 12 animals. Data points indicate mean ± SeM.