Heterodimerization of UNC-13/RIM regulates synaptic vesicle release probability but not priming in C. elegans

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Abstract

UNC-13 proteins play an essential role in synaptic transmission by recruiting synaptic vesicles (SVs) to become available for release, which is termed SV priming. Here we show that the C2A domain of UNC-13L, like the corresponding domain in mammalian Munc13-1, displays two conserved binding modes: forming C2A/C2A homodimers, or forming a heterodimer with the zinc finger domain of UNC-10/RIM (C2A/RIM). Functional analysis revealed that UNC-13L's C2A promotes synaptic transmission by regulating a post-priming process. Stimulus-evoked release but not SV priming, was impaired in unc-10 mutants deficient for C2A/RIM heterodimerization, leading to decreased release probability. Disrupting C2A/C2A homodimerization in UNC-13L-rescued animals had no effect on synaptic transmission, but fully restored the evoked release and the release probability of unc-10/RIM mutants deficient for C2A/RIM heterodimerization. Thus, our results support the model that RIM binding C2A releases UNC-13L from an autoinhibitory homodimeric complex to become fusion-competent by functioning as a switch only.

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Introduction

Neurotransmitters are released by Ca²⁺-triggered synaptic vesicle (SV) exocytosis, a process mediated by the SNARE complex, and regulated by synaptic proteins such as Munc18s, Munc13s, RIMs, and synaptotagmins (Südhof and Rizo, 2011; Jahn and Fasshauer, 2012). Before final fusion with the plasma membrane, SVs undergo two crucial steps: docking and priming. These processes ensure that the SVs are translocated to the plasma membrane and become fusion-competent at the active zone (Sudhof, 2004). Defects in docking or priming produce corresponding decreases in SV exocytosis (Aravamudan et al., 1999; Augustin et al., 1999; Richmond et al., 1999; Hammarlund et al., 2007; Imig et al., 2014). However, post-priming regulation of SV release has also been reported (Pang et al., 2006; Bacaj et al., 2015; Luo et al., 2015). Various studies have demonstrated that Munc13-1, a protein with multiple functional domains, is required for the docking and priming (Aravamudan et al., 1999; Augustin et al., 1999; Richmond et al., 1999), as well as post-priming Ca²⁺-triggered SV release (Basu et al., 2007; Chen et al., 2013; Hu et al., 2013). The functions of Munc13-1 in these processes are believed to be mediated by its central MUN domain, which directly interacts with syntaxin and promotes SNARE assembly (Ma et al., 2011; Yang et al., 2015).

Multiple domains of Munc13-1 have been implicated in regulating the MUN domain’s SV priming activity (Michelassi et al., 2017; Xu et al., 2017). In mammals, four Munc13 isoforms have been...
identified, two of which contain an N-terminal C2A domain (Munc13-1 and ubMunc13-2) (Brose et al., 1995; Song et al., 1998; Betz et al., 2001). Betz et al. first found that the C2A domain binds to the zinc finger (ZF) domain of RIM, an active zone protein that is required for SV priming (Betz et al., 2001). A double mutation in rat RIM2α’s ZF domain (K97/99E) disrupts C2A/RIM heterodimerization (Dulubova et al., 2005) and was subsequently shown to produce a large reduction in the readily releasable pool (RRP) of SVs (Deng et al., 2011; Camacho et al., 2017), indicating that C2A binding to RIM plays an important role in SV priming. Apart from forming a heterodimer with RIM, the C2A domain also forms a homodimer, thereby promoting formation of Munc13-1 homodimers (Lu et al., 2006). This homodimerization, however, is not necessary for SV priming and post-priming SV release (Camacho et al., 2017).

In contrast, studies in chromaffin cells have shown that a truncated Munc13-1 lacking the N-terminal C2A domain is fully able to mediate vesicle priming (Betz et al., 2001), suggesting that the role of Munc13-1 in priming is independent of the C2A domain and its binding with RIM. Similarly, a recent study in C. elegans found that removing the N-terminal C2A domain from UNC-13L, a Munc13-1 homolog, does not cause a change in the size of the RRP (Zhou et al., 2013a). These conflicting findings have called into doubt whether binding of the C2A domain with RIM is indeed playing a role in SV priming, prompting us to re-analyze the function of both C2A/RIM heterodimerization and C2A/C2A homodimerization using an in vivo system.

Here, we examined the functional importance of the C2A domain by generating a transgenic line of UNC-13L with a point mutation in the C2A domain (K49E) that blocks formation of C2A/C2A homodimers, and a mutant carrying two mutations in the unc-10/RIM ZF domain (K77/79E) that disrupts C2A/RIM heterodimer formation. We found that tonic neurotransmitter release at a low Ca\(^{2+}\) concentration and stimulus-evoked release were significantly reduced in the unc-10(nu487 K77/79E) mutants, whereas SV release was normal in the UNC-13L(K49E)-rescued unc-13 mutants. Moreover, the hypertonic sucrose-evoked responses were unaltered following disruption of homodimerization and heterodimerization, suggesting that SV priming is not determined by the state of the C2A domain. Interestingly, simultaneously disrupting C2A/C2A homodimerization in the unc-10(nu487 K77/79E) mutant fully restored both tonic and evoked release. These results suggest that the synaptic transmission defect in the unc-10(nu487 K77/79E) mutant is most likely caused by formation of more C2A/C2A homodimers, and that the role of UNC-10/RIM binding UNC-13 C2A domain is to release the C2A from an autoinhibitory state, allowing it to become a fusion-competent monomeric form. Finally, we show that UNC-13’s C2A domain increases release probability but has little effect on SV priming.

Results
C2A/C2A homodimerization and C2A/RIM heterodimerization are conserved in C. elegans
The UNC-13L C2A domain and the UNC-10/RIM ZF domain share high sequence identity with the corresponding domains of mouse Munc13-1 (50%) and RIM (48%) (Figure 1A,B), suggesting that UNC-13’s C2A would form homo- and heterodimers in a manner similar to mouse Munc13-1. To test this idea, we directly assessed homo- and heterodimer formation by the UNC-13 C2A domain. Gel filtration analysis indicated that recombinant C2A forms a homodimer, and that this homodimerization was disrupted by a mutation (K49E) analogous to that which disrupts Munc13-1 homodimerization (Figure 1A,C) (Lu et al., 2006; Deng et al., 2011). To determine whether UNC-13L forms a heterodimer with UNC-10/RIM, we performed isothermal titration calorimetry (ITC) and co-immunoprecipitation experiments. Our results showed that the UNC-13L C2A domain tightly binds the UNC-10/RIM ZF domain (Kd = 1.29 μM, Figure 1D,E), and that this interaction was eliminated by a double mutation in the ZF domain (K77/79E), which is analogous to mutations that disrupt binding of the corresponding mammalian Munc13-1 and RIM domains (Dulubova et al., 2005). Thus, our findings confirmed that UNC-13L C2A has the same two binding modes exhibited by mouse Munc13-1.
Figure 1. UNC-13L exhibits conserved homodimerization and heterodimerization in C. elegans. (A, B) Sequence alignment of the C2A domain between worm UNC-13L and rat Munc13-1, and the zinc finger (ZF) domain between worm UNC-10 and mouse RIM1. The conserved residues in the C2A domain (K49) and the ZF domain (K77, K79) that disrupt homodimer and heterodimer formation are indicated by stars. (C) Separation of purified UNC-13L C2A monomer and homodimer by gel filtration. The wild-type C2A forms a homodimer that runs at 12.57 mL (blue trace, corresponding to an apparent molecular weight of 66 kDa). (D) Immunoblot analysis of purified UNC-13L C2A monomer and homodimer. (E) Kinetic analysis of the corrected heat rate and normalized fit for the homodimeric interaction. (F) Kinetic analysis of the corrected heat rate and normalized fit for the homodimeric interaction.
weight of 89 kDa; this homodimer is disrupted in the monomeric UNC-13L(K49E) mutant which runs at 13.88 mL (apparent weight 52 kDa). Elution volumes of gel filtration standards are indicated underneath the plot, as follows: 11.62 mL for alcohol dehydrogenase (150 kDa), 12.85 mL for bovine serum albumin (66 kDa), and 15.45 mL for carbonic anhydrase (29 kDa), with the column void volume being 10.78 mL. (D) Co-immunoprecipitation (co-IP) shows that HA-LNTD (UNC-13L N-terminal domain, aa 1–605) binds wild-type FLAG-UNC-10 (aa 1–601) but not UNC-10(K77/79E) or GST. These results are representative of three independent experiments. (E, F) Isothermal titration calorimetry (ITC) analysis of binding of wild-type or mutated (K77/79E) UNC-10 with UNC-13L.

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**Disrupting C2A/RIM heterodimerization reduces mEPSCs/miPSCs in low Ca\textsuperscript{2+}

We next examined the functional roles of the C2A/C2A homodimerization and C2A/RIM heterodimerization in tonic neurotransmitter release by measuring miniature excitatory/inhibitory postsynaptic currents (mEPSCs/miPSCs) at the neuromuscular junction. Both the mEPSCs and the miPSCs were recorded in the presence of 0 mM and 1 mM Ca\textsuperscript{2+}, as we have previously found that the miPSCs are almost arrested in 0 mM Ca\textsuperscript{2+} but are normal 1 mM Ca\textsuperscript{2+} in some mutants (Liu et al., 2018), suggesting that different Ca\textsuperscript{2+} concentrations should be tested to provide an accurate and thorough evaluation of tonic release. Our results showed that the mEPSCs and miPSCs were completely abolished in the unc-13(s69) null mutant, and were fully restored by neuronal expression of full-length UNC-13L (Figure 2). Deleting the C2A domain (UNC-13LΔC2A) significantly decreased mEPSC frequency in both 0 mM and 1 mM Ca\textsuperscript{2+} (Figure 2B,E, Table 1), demonstrating an important role of the C2A domain in tonic acetylcholine (ACh) release. The miPSC frequency was significantly decreased in the UNC-13LΔC2A rescue animals in 0 mM Ca\textsuperscript{2+} (Figure 2C, Table 1), but was unchanged in 1 mM Ca\textsuperscript{2+}, compared to that in the UNC-13L rescue animals (Figure 2F). Thus, the UNC-13L C2A domain is important for tonic release of both ACh and GABA; however, the requirement for C2A in GABA release can be bypassed by the presence of extracellular calcium, presumably because of increased release probability. The differential impact of deleting C2A on ACh and GABA release in 1 mM Ca\textsuperscript{2+} could reflect differences in release probability at these two types of synapses (Liu et al., 2018). In fact, SV release from inhibitory synapses displays a higher release probability than excitatory synapses in the mammalian central nervous system (Zhou et al., 2013b; Zhou et al., 2013c). Deleting UNC-13L’s C2A domain could also alter other aspects of synapse structure, such as SV trafficking, and coupling of SVs to Ca\textsuperscript{2+} entry at the release sites.

We next asked whether C2A/C2A homodimerization or C2A/RIM heterodimerization accounts for the role of the C2A domain in tonic release. To investigate this, we first examined the mEPSCs and miPSCs in the unc-13 mutants rescued by UNC-13L carrying the K49E mutation, which blocks C2A/C2A homodimerization (Figure 1B). The mEPSC and miPSC frequencies in the UNC-13L(K49E) rescue animals were comparable to those in the UNC-13L rescue animals in both 0 mM and 1 mM Ca\textsuperscript{2+} conditions (Figure 2A–F, Table 1), indicating that homodimerization does not affect tonic ACh or GABA release. To determine the role of C2A/RIM heterodimerization, we used crispr to introduce a K77/79E double mutation in the UNC-10 ZF domain, unc-10(nu487 K77/79E). Compared to wild-type animals, unc-10(nu487 K77/79E) mutants display slightly uncoordinated behaviour (data not shown). Analysis of tonic release showed that the mEPSCs and miPSCs were markedly reduced in unc-10(nu487 K77/79E) mutants in 0 mM Ca\textsuperscript{2+}, whereas both were unaltered in 1 mM Ca\textsuperscript{2+} (Figure 2A–F), suggesting that C2A/RIM heterodimerization is only required for tonic release in low Ca\textsuperscript{2+} when release probability is reduced. To confirm this, we measured mEPSCs and miPSCs in two additional Ca\textsuperscript{2+} concentrations (0.05 mM and 0.1 mM). As shown in Figure 2G, increasing the Ca\textsuperscript{2+} concentration from 0 mM to 0.05 mM and 0.1 mM caused robust enhancement of the mEPSC and miPSC frequencies in wild-type animals. In unc-10(nu487 K77/79E) mutants, a significantly lower miPSC frequency was observed in 0.05 mM Ca\textsuperscript{2+} but not in 0.1 mM Ca\textsuperscript{2+}, whereas mEPSCs were unaltered in both 0.05 mM and 0.1 mM Ca\textsuperscript{2+} (Figure 2G). Our results therefore demonstrate that C2A/RIM heterodimerization promotes tonic release but that this effect is bypassed when release probability is increased by raising external Ca\textsuperscript{2+} concentrations.
Figure 2. Tonic release is reduced by disrupting C2A/RIM heterodimerization in low Ca\(^{2+}\). (A) Representative traces of mEPSCs and mIPSCs recorded at 0 mM Ca\(^{2+}\) from indicated genotypes. (B, C) Averaged frequency and amplitude of the mEPSCs and mIPSCs. (D) Representative traces of mEPSCs and mIPSCs recorded at 1 mM Ca\(^{2+}\) from the same genotypes in (A). (E, F) Quantification of the mEPSC and mIPSC frequency and amplitude from the genotypes in (D). Data are mean ± SEM (##, p < 0.01, ###, p < 0.001, n.s., non-significant when compared to UNC-13L rescue; ***, p < 0.001 when compared to unc-10 (K7779E) rescue).
Table 1. Comparison of SV release.

| Tonic release | Evoked EPSC | RRP | P_{wr} |
|---------------|------------|-----|--------|
| mEPSC (0 mM Ca^{2+}) | mEPSC (1 mM Ca^{2+}) | mIPSC (0 mM Ca^{2+}) | mIPSC (1 mM Ca^{2+}) |
| Frequency (Hz) | Amplitude (pA) | Frequency (Hz) | Amplitude (pA) | Amplitude (pA) | Amplitude (nA) | Charge (-pC) | Charge (-pC) |
| Wild-type | 1.87 ± 0.16 | 23.2 ± 1.6 | 43.5 ± 4.3 | 23.9 ± 1.1 | 11 ± 1.1 | 25.9 ± 1.1 | 42.5 ± 4.1 | 22.5 ± 2.1 | 2.27 ± 0.2 | 20.6 ± 2.1 | 221 ± 23.1 | 0.12 ± 0.01 |
| unc-13 (nu487) | 0.05 ± 0.05 | 20.2 ± 3 | 2 ± 1 | 21.1 ± 2.1 | 0.05 ± 0.05 | 22.1 ± 4.1 | 1.1 ± 0.5 | 21.5 ± 3.2 | 0.02 ± 0.01 | 0.05 ± 0.01 | 25.5 ± 7.1 | 0 |
| UNC-13L | 1.51 ± 0.45 | 19.3 ± 1.9 | 43.4 ± 3.7 | 23.9 ± 1.1 | 10.1 ± 0.9 | 22.3 ± 2.2 | 38.1 ± 3.5 | 22.4 ± 1.1 | 1.7 ± 0.14 | 15.4 ± 2.6 | 237 ± 25.4 | 0.06 ± 0.01 |
| UNC-14LaC2A | 0.64 ± 0.07 | 20.5 ± 1.4 | 20.2 ± 3 | 23.3 ± 2.2 | 5.4 ± 0.5 | 28.2 ± 1.9 | 34.4 ± 3.2 | 25.7 ± 1.8 | 1.08 ± 0.12 | 8.4 ± 1.6 | 248 ± 66.3 | 0.034 ± 0.006 |
| UNC-13L (K49E) | 1.9 ± 0.48 | 19.4 ± 1.1 | 43.3 ± 4.3 | 19.3 ± 2.4 | 9.8 ± 1.2 | 23.4 ± 1.5 | 40.6 ± 5.1 | 23.7 ± 3.2 | 2.1 ± 0.27 | 17.3 ± 2.3 | 243 ± 38.3 | 0.07 ± 0.01 |
| unc-10 (K77/79E) | 0.57 ± 0.1 | 20.2 ± 1.8 | 45.4 ± 3.7 | 21.1 ± 1.2 | 3.5 ± 0.5 | 27.3 ± 1.8 | 39.7 ± 4.1 | 26.7 ± 1.1 | 1.18 ± 0.11 | 11.5 ± 1.5 | 245 ± 28.2 | 0.046 ± 0.006 |
| K49E; K77/79E | 0.94 ± 0.18 | 23.1 ± 1.5 | 40.8 ± 4.4 | 22.2 ± 1.6 | 9.5 ± 1 | 26.3 ± 1.4 | 43.7 ± 6.3 | 26.5 ± 1.8 | 1.81 ± 0.15 | 18.1 ± 2.2 | 260 ± 40 | 0.07 ± 0.008 |

All data were collected from three independent experiments to ensure the observed phenotype are replicable.

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A monomeric C2A domain is fully functional for tonic release

The decreased mEPSC frequency observed in the UNC-13LΔC2A rescue animals at 1 mM Ca^{2+} indicates that the C2A domain may have an additional function in regulating tonic release, particularly at high Ca^{2+}. We therefore examined whether simultaneously blocking the homo- and heterodimerization of the C2A domain could account for this additional function. To do that, we made UNC-13L (K49E); unc-10(nu487 K77/79E) double mutant by crossing the unc-10(nu487 K77/79E) into the UNC-13L(K49E) rescue strain. In 0 mM Ca^{2+} recordings, the UNC-13L(K49E); unc-10(nu487 K77/79E) double mutant mIPSC rate was significantly higher than that observed in unc-10(nu487 K77/79E) single mutants, suggesting that blocking homodimer formation was sufficient to prevent the decreased tonic GABA release caused by the unc-10(nu487 K77/79E) mutation (Figure 2C). By contrast, blocking homodimerization did not significantly improve mEPSC rates in 0 mM Ca^{2+} in unc-10(nu487 K77/79E) mutants (Figure 2B). Thus, heterodimerization increases tonic GABA release by preventing homodimer formation, whereas heterodimerization promotes tonic ACh release by a distinct mechanism. In addition, the tonic ACh and GABA release were at wild-type level in the UNC-13L(K49E); unc-10(nu487 K77/79E) double mutant in 1 mM Ca^{2+}, similar to those in the UNC-13L(K49E) rescue and the unc-10(nu487 K77/79E) mutant (Figure 2E,F). Taken together, these results demonstrate that blocking C2A homo- and heterodimerization does not decrease tonic ACh and GABA release. These results differ from those obtained in the UNC-13LΔC2A rescue animals, suggesting that the...
effect of deleting the C2A domain on the tonic release is not a result of a lack of homo- and heterodimerization.

**Stimulus-evoked release is reduced when C2A/RIM heterodimerization is blocked**

Next, we analysed the impact of C2A interactions on stimulus-evoked ACh release. Compared to UNC-13L, UNC-13LΔC2A exhibited a smaller evoked EPSC (Figure 3A, Table 1), consistent with previous findings (Hu et al., 2013; Zhou et al., 2013a). The evoked EPSCs were significantly smaller in unc-10(nu487 K77/79E) mutants, exhibiting decreased amplitude and total charge transfer compared to wild-type animals (Figure 3A–C). The percentage decrease of the evoked charge transfer in the unc-10(nu487 K77/79E) mutant (44%) was similar to that observed in the UNC-13LΔC2A rescue animals (46%), suggesting that C2A/RIM heterodimerization accounts for the role of the C2A domain in evoked release. Thus, our results revealed that C2A/RIM heterodimerization is required for evoked neurotransmitter release. It should be noted that the evoked EPSCs were recorded in 1 mM Ca\(^{2+}\) solution in which, however, the mEPSCs were unchanged (Figure 2E). These different observations on tonic and evoked neurotransmitter release may reflect differential release probabilities of the two release modes at various Ca\(^{2+}\) levels. Indeed, the mEPSCs and the stimulus-evoked EPSCs exhibit large differences in their Ca\(^{2+}\) dependence (Williams et al., 2012).

Next, we asked whether C2A homodimers regulate evoked responses. The evoked EPSCs mediated by UNC-13L(K49E) were indistinguishable from those rescued by wild-type UNC-13L, indicating that C2A homodimers are not required for evoked responses (Figure 3A–C). To determine whether homodimers inhibited evoked ACh release, we analyzed UNC-13L(K49E); unc-10 (nu487 K77/79E) double mutants. We found that preventing C2A/C2A homodimerization in the unc-10(nu487 K77/79E) mutant fully restored the evoked EPSC to a wild-type level (Figure 3A–C). These results suggest that the reduced evoked EPSC caused by blocking C2A/RIM heterodimerization is mediated by increased formation of C2A homodimers. These results are consistent with prior analysis of mouse synapses, which suggest that Munc13-1 homodimers inhibit SV release (Deng et al., 2011). These results also demonstrate that the monomeric C2A domain is sufficient to support evoked release, consistent with the observation of tonic release (Figure 2, Table 1). Thus, the RIM ZF domain appears to function only as a switch that releases the C2A domain from the inhibitory homomeric complex.

**The C2A domain is not required for SV priming**

A decrease in evoked EPSCs may arise from a priming defect or a change in release probability. We therefore examined whether SV priming is regulated by the C2A domain and its two different states (homodimer and heterodimer). SV priming was measured by the pulsed application of hypertonic sucrose solution (1M, 2 s) onto the ventral nerve cord, and the size of the RRP was calculated by integrating the sucrose-evoked current. The current was almost eliminated in the UNC-13L(K49E) rescue animals and the RRP was calculated by applying hypertonic solution in which, however, the mEPSCs were unchanged (Figure 2E). These different observations on tonic and evoked neurotransmitter release may reflect differential release probabilities of the two release modes at various Ca\(^{2+}\) levels. Indeed, the mEPSCs and the stimulus-evoked EPSCs exhibit large differences in their Ca\(^{2+}\) dependence (Williams et al., 2012).

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Next, we asked whether C2A homodimers regulate evoked responses. The evoked EPSCs mediated by UNC-13L(K49E) were indistinguishable from those rescued by wild-type UNC-13L, indicating that C2A homodimers are not required for evoked responses (Figure 3A–C). To determine whether homodimers inhibited evoked ACh release, we analyzed UNC-13L(K49E); unc-10 (nu487 K77/79E) double mutants. We found that preventing C2A/C2A homodimerization in the unc-10(nu487 K77/79E) mutant fully restored the evoked EPSC to a wild-type level (Figure 3A–C). These results suggest that the reduced evoked EPSC caused by blocking C2A/RIM heterodimerization is mediated by increased formation of C2A homodimers. These results are consistent with prior analysis of mouse synapses, which suggest that Munc13-1 homodimers inhibit SV release (Deng et al., 2011). These results also demonstrate that the monomeric C2A domain is sufficient to support evoked release, consistent with the observation of tonic release (Figure 2, Table 1). Thus, the RIM ZF domain appears to function only as a switch that releases the C2A domain from the inhibitory homomeric complex.
Figure 3. Effects of homodimerization and heterodimerization on evoked neurotransmitter release. Electronic stimulus and hypertonic sucrose-evoked EPSCs were recorded from the body wall muscle of adult animals in 1 mM Ca$^{2+}$. (A) Example traces of the evoked EPSCs from indicated genotypes. (B–D) Quantification of the amplitude, charge transfer and 20–80% risetime of the evoked EPSCs in (A). Example traces and averaged charge transfer of the sucrose-evoked EPSCs from the same genotypes in (A) are shown in (E) and (F). (G) Quantification of the probability of SV release ($P_r$). Data are Figure 3 continued on next page.
currents was significantly reduced in the ΔC2A rescue and the unc-10(K77/79E) mutant (Figure 3G). It should be noted that recording of the sucrose-evoked current requires a tight gigaohm seal between the recording pipette and the cell, as a loose seal often leads to an unreal current. We therefore analyzed the leak currents for each sucrose recording. Our data showed that all recordings were controlled well with acceptable leak currents (Figure 3—figure supplement 1). Together, our results demonstrate that the UNC-13L C2A domain regulates release probability by interacting with the UNC-10/RIM, rather than by regulating SV priming.

The decreased release probability in the UNC-13LΔC2A rescue and the unc-10(K77/79E) mutant could arise from a change in the expression level of UNC-13, or a change in sub-localization of the UNC-13 proteins at the nerve terminals. We next analyzed the UNC-13 protein levels from different transgenic rescue lines. A fluorescence protein (mApple) was fused to the C terminus of the various UNC-13L constructs (WT, K49E, and ΔC2A), and expressed in unc-13 null mutants. Our prior study showed that c-terminal mCherry tags do not interfere with UNC-13’s function in promoting SV exocytosis (Hu et al., 2013). All mApple-tagged UNC-13L constructs rescued the SV release in the unc-13 mutants, and the phenotypes were similar to those rescued by corresponding untagged UNC13L constructs (Figure 4—figure supplement 1). By quantifying the mApple fluorescence intensity from different rescue lines, we showed that there were no significant differences (Figure 4A,B), indicating that the UNC-13 proteins were expressed at similar levels in these lines. These results suggest that the decreased tonic and evoked release in UNC-13LΔC2A rescued animals are unlikely to result from reduced UNC-13L abundance. Prior studies have shown that the C2A domain is crucial for synaptic localization of UNC-13 (Melom et al., 2013). Therefore, it is likely that the decrease in release probability in the UNC-13LΔC2A rescue animals was caused by mis-localization of UNC-13 at the active zone.

We next asked if the decreased release probability in the unc-10(K77/79E) mutant was caused by a change in the sub-localization of UNC-13L. To determine this, we compared colocalization of UNC-13L and an active zone marker (ELKS-1) in wild-type controls and in unc-10(K77/79E) mutants. UNC-13L and ELKS-1 were strongly colocalized in wild-type (Figure 4C,D), and this colocalization was not obviously altered in unc-10(K77/79E) mutants (Figure 4E,F). These results suggest that disrupting UNC-13L binding to RIM does not prevent synaptic localization of UNC-13L.

**Disrupting heterodimerization does not affect synaptic depression and recovery**

Because of the reduced P_{vr} in the unc-10(K77/79E) mutant, we asked whether the synaptic depression and recovery would be altered by disruption of C2A/RIM heterodimerization. An integrated line expressing ChlEF, a variant of channelrhodopsin-2 (ChR2) in the cholinergic motor neurons was used to record the EPSCs evoked by a train or a paired light stimulus (Liu et al., 2009; Watanabe et al., 2013). Both the control animals and the unc-10(K77/79E) mutants exhibited synaptic depression following a 1 Hz or 5 Hz train stimulus (Figure 5A,C). The depression rate was described by normalizing the amplitudes of all the EPSCs (EPSC_i) to the amplitude of the first EPSC (EPSC_1). As shown in Figure 5F, no difference in the recovery rate was observed between the control animals and the unc-10(K77/79E)
mutants. This indicates that disrupting C2A/RIM binding does not affect refilling of the SVs at synapses.

Figure 4. Expression of UNC-13 transgenic constructs and effects of disrupting UNC-13L/RIM heterodimerization on sub-localization of UNC-13L. (A) Presynaptic distribution of UNC-13 at the dorsal nerve cord in unc-13 animals rescued by the indicated transgenic constructs. Scale bar, 5 μm. (B) Quantification of the mApple fluorescence intensity from the transgenic rescue lines in (A). (C) Representative confocal Z-stack images for UNC-13L and ELKS-1 in wild-type background. Scale bar, 5 μm. (D) Line scans along the dorsal nerve cord. (E) Representative confocal Z-stack images for UNC-13L and ELKS-1 in unc-10(K77/79E) mutants. (F) Line scans along the dorsal nerve cord. Data are mean ± SEM (n.s., non-significant; one-way ANOVA). The number of worms analyzed for each genotype is indicated in the bar graphs.

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The following figure supplement is available for figure 4:

Figure supplement 1. UNC-13 proteins with C-terminal tags are fully functional in mediating SV release.

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Figure 5. Synaptic depression and recovery are unaltered by disrupting C2A/RIM heterodimerization. Synaptic depression and recovery were investigated by applying a train (1 Hz or 5 Hz) or a paired light stimulus onto the ventral nerve cord of the adult worms with expression of ChIEF in cholinergic motor neurons. (A, C) Example traces of 1 Hz and 5 Hz light train stimulus-evoked EPSCs (black, wild-type; red, unc-10(K77/79E) mutants). (B, D) Quantification of synaptic depression by normalizing the EPSC amplitude (EPSC \(_i\)) to the first EPSC amplitude (EPSC \(_1\)). (E) Evoked EPSCs
Discussion

In this study, we analyzed the functional importance of the C2A domain of UNC-13L, a Munc13-1 homolog in C. elegans, for SV priming and release probability. Prior studies in mouse and C. elegans have reported contradictory results regarding the role of the C2A domain in priming (Betz et al., 2001; Zhou et al., 2013a; Camacho et al., 2017). This motivated us to investigate the function of this domain, with a particular focus on its two binding modes, in SV priming and post-priming Ca2+-triggered exocytosis. Below we discuss our findings.

UNC-13L primes SVs independently of the C2A domain

The fact that the UNC-13/Munc13 is required for SV priming has been reported in both invertebrates and vertebrates (Aravamudan et al., 1999; Augustin et al., 1999; Richmond et al., 1999). Sucrose-evoked responses are almost eliminated in neurons lacking UNC-13 or Munc13. Thus far, however, it is still unclear how the priming activity of UNC-13/Munc13 is achieved and regulated. It is believed that the MUN domain plays an essential role in SV priming. Removing the C2A domain from Munc13-1 leads to a marked decrease in the RRP of SVs in cultured hippocampal neurons, suggesting a role for C2A in priming (Camacho et al., 2017). However, we found that unc-13 mutants lacking a C2A domain (UNC-13L C2A rescued animals) exhibit a normal RRP, arguing that the C2A domain is not required for priming (Figure 3E). Zhou et al. showed that an isolated unc-13 mutant (n2609) with a point mutation in the C2A domain of UNC-13L exhibits a similar reduction in evoked EPSCs to UNC-13L C2A, whereas docked SV numbers are normal in the unc-13(n2609) mutant. This suggests that the C2A domain is not required for docking (Zhou et al., 2013a), consistent with our observations in relation to priming. It should be noted that all the UNC-13 transgenes in this study are extrachromosomal arrays, and that the UNC-13 proteins are overexpressed in these transgenic animals. The overexpression of UNC-13 may cause differential effects with wild-type expression of this protein. Despite this, our findings (e.g., ΔC2A) are similar to the results of Zhou et al., which were obtained from unc-13 mutants rescued by a single copy of UNC-13.

Our results imply that UNC-13L regulates SV priming via a different mechanism, and that this mechanism is independent of the C2A domain. Prior studies have shown that a point mutation in the C1 domain of Munc13-1 accelerates hypertonicity-induced SV release without changing the size of the RRP (Basu et al., 2007). Transgenic C. elegans rescued by a truncated UNC-13L lacking its C1 or C2B domain also exhibit a normal RRP (Michelassi et al., 2017). Expression of a mutated Munc13-1 (P827L, located in the linker domain between C2B and MUN) leads to a significant increase in release probability without causing an increase in the size of the RRP (Lipstein et al., 2017). These observations appear to support the notion that the regulatory domains in UNC-13L (e.g. C2A, C1 and C2B) alter release probability but are not required for SV priming. Thus, our results, together with the findings from other studies in C. elegans and mouse, suggest that UNC-13 has multiple regulatory effects on synaptic strength and plasticity, which are achieved by differential control of release probability by its various functional domains.

State of the C2A domain and release probability

The reduced evoked EPSCs and unchanged RRP in the UNC-13LΔC2A rescue animals led to a significant decrease in release probability (Pv), consistent with previous studies (Camacho et al., 2017). Similar to these observations, the unc-10(K77/79E) mutant also exhibits an impairment in evoked EPSCs but a normal RRP, thereby leading to a decreased Pv (Figure 3). These results demonstrate that deleting the C2A domain or blocking C2A/RIM heterodimerization decreases release probability. It should be noted that, release probability remains unchanged in the hippocampal neurons when C2A/RIM heterodimerization is disrupted, because of a similar decrease in the evoked EPSCs and the RRP (Camacho et al., 2017), different from our findings in this study.
The function of the C2A domain in regulating the release probability appears to be determined by its state. It has been proposed that the Munc13 C2A domain has three binding states, homodimer, heterodimer, and monomer, and that these three forms may co-exist in synaptic terminals. The decrease in evoked EPSC and release probability in the unc-10(nu487 K77/79E) mutant was blocked by disrupting C2A/C2A homodimerization (Figure 3). This indicates that the synaptic transmission defect caused by disrupting C2A/RIM heterodimerization occurs because of an increase in C2A/C2A homodimerization, which is proposed to comprise an autoinhibitory state. Again, this is post-priming regulation, as SV priming is normal in both the UNC-13L(K49E) rescue and UNC-13L (K49E); unc-10(nu487 K77/79E) animals (Figure 3E). Although our findings are similar to the observations reported in cultured hippocampal neurons in which the reduced evoked EPSCs caused by the disruption of heterodimerization were partially recovered when homodimerization and heterodimerization were disrupted simultaneously (Camacho et al., 2017), we found that UNC-13L with a monomeric C2A domain is fully functional for both the priming and post-priming Ca\textsuperscript{2+}-triggered tonic and evoked release. Thus, our results support the notion that homodimerization of the C2A domain inhibits SV release, and that RIM releases the C2A domain from the autoinhibitory homodimeric complex by competitively binding the C2A domain to form a heterodimer, thereby allowing the C2A domain to become fusion-competent. The heterodimer itself, however, does not produce an additional effect on synaptic transmission.

**Distinct priming mechanisms between worm and mouse?**

RIM binding to Munc13-1 is a key determinant in SV priming in the mouse central nervous system. In fact, deleting Munc13-1 C2A or disrupting C2A/RIM heterodimerization leads to a similar reduction in the size of the RRP to that observed in RIM knockout neurons (Deng et al., 2011; Camacho et al., 2017). Whereas SV priming in worm does not rely on the UNC-13 C2A and the heterodimerization of UNC-13/RIM, although this interaction is highly conserved between worm and mouse. This raises a question of whether the UNC-10/RIM is required for priming in worm. Electron microscopy results have revealed that the number of docked SVs is slightly decreased in the unc-10 mutant (Koushika et al., 2001; Weimer et al., 2006; Gracheva et al., 2008), which cannot account for the far more dramatic SV fusion defect, suggesting a post-docking role of RIM in the active zone. Consistent with these relatively minor docking defects, we found that the size of the RRP in unc-10 mutants (assessed by sucrose evoked currents) is comparable to that in the wild-type animals (data not shown), demonstrating that UNC-10/RIM is not required for SV priming. These results suggest that UNC-10/RIM primarily regulates SV fusion by increasing release probability and not by promoting SV priming. UNC-10/RIM increases release probability via its ability to bind UNC-13 but also through its requirement for recruiting CaV2 calcium channels to presynaptic terminals (Han et al., 2011; Kaeser et al., 2011; Müller et al., 2012).

Why does RIM play different roles at worm and mouse synapses? SV priming is thought to consist of two sequential steps, termed molecular and positional priming (Neher and Sakaba, 2008). Molecular priming is proposed to account for a slow component of release, which consists of fusion of SVs that are docked far from calcium channels driving release. Positional priming is proposed to mediate a faster form of release, which comprises fusion of SVs that are docked very close to calcium channels. One possibility is that worms rely less on positional priming, which presumably requires UNC-13/RIM interaction, and instead rely more on molecular priming. In this scenario, worm unc-10 RIM mutants would have decreased release probability but relatively normal RRP size. By contrast, mice may require positional priming to maintain the RRP, and thus RIM mutants would have significant priming defects.

### Additional functions of the C2A domain

Despite the above findings, the actual mechanism whereby the UNC-13L with a monomeric C2A regulates SV release remains unclear. Based on the normal synaptic transmission in the UNC-13L (K49E); unc-10(nu487 K77/79E) double mutants, RIM binding to the UNC-13L C2A domain only appears to be playing a switch role. The C2A domain must therefore have additional functions for regulation of SV release. This is confirmed by our observations in relation to tonic release, which has been proposed to have a different regulatory mechanism from evoked release (Ramirez and Kavalali, 2011; Melom et al., 2013). The mEPSC frequency was reduced by 50% in the UNC-13LΔC2A
rescue animals at both 0 mM and 1 mM Ca\(^{2+}\), similar to the evoked release findings (Figure 2), revealing an important role of the C2A domain for tonic release at cholinergic synapses. However, this role cannot be explained by the state of the C2A domain (homodimer, heterodimer, or monomer), as the mEPSC frequencies were normal in the UNC-13L(K49E) rescue, unc-10(nu487 K77/79E) mutant, and UNC-13L(K49E); unc-10(nu487 K77/79E) double mutants in 1 mM Ca\(^{2+}\). We only observed a change in the Ca\(^{2+}\) sensitivity in very low Ca\(^{2+}\) at GABAergic synapses, and this phenotype disappeared when the Ca\(^{2+}\) levels were increased (Figure 2G). Our observations at cholinergic synapses therefore demonstrate that, although the C2A/C2A homodimer inhibits evoked release, it does not affect tonic release. There must still be an unknown mechanism by which the monomeric C2A domain regulates post-priming SV exocytosis.

Materials and methods

Strains
Strain maintenance and genetic manipulation were performed as previously described (Brenner, 1974). Animals were cultivated at room temperature on nematode growth medium (NGM) agar plates seeded with OP50 bacteria. On the day before experiments L4 larval stage animals were transferred to fresh plates seeded with OP50 bacteria for all the electrophysiological recordings. The following strains were used:

- Wild-type, N2 bristol
- KP6901 unc-13(s69)-l
- KP7503 nu487 unc-10(nu487 K77/79E), X
- KP6893 nuEx1515 [Psnb-1::UNC-13L; unc-13(s69)]
- ZTH6 unc-13(s69); hztEx106 [Psnb-1::UNC-13LΔC2A]
- ZTH454 unc-13(s69); hztEx91 [Psnb-1::UNC-13L(K49E)]
- ZTH7 unc-10(nu487 K77/79E); hztEx91 [Psnb-1::UNC-13L(K49E); unc-13(s69)]
- ZTX460 zxls6 [Punc-17::ChR2(H134R)::YFP + lin-15(+) + ] V
- ZTH71 unc-10(nu487 K77/79E); zxls6
- ZTH605 unc-13(s69); hztEx76 [Psnb-1::UNC-13L::mApple]
- ZTH606 unc-13(s69); hztEx77 [Psnb-1::UNC-13LΔC2A::mApple]
- ZTH607 unc-13(s69); hztEx78 [Psnb-1::UNC-13L(K49E)::mApple]
- JSD0025 tauls12 (9x) [Punc-129::ELKS-1::Cerulean::Cerulean LG]
- EG5793 oxSi91 [Punc-17::ChIEF::mCherry::unc-54UTR; unc-119(+)] II
- ZTH608 unc-10(nu487 K77/79E); oxSi91, MosSCI

‘Ex’ in the strain name indicates that the rescue experiments use extrachromosomal array.

Constructs, transgenes and germline transformation
A 5.451 kb cDNA corresponding to UNC-13L was amplified by PCR from the N2 cDNA library and inserted into MCSII of the JB6 vector between the KpnI and NotI sites. The snb-1 promoter (3 kb) was inserted into MCSII between the Sphl and BamHI sites. The C2A domain deletion removed the first 150 amino acids. The 5’ primer (GCTGCAGACTGTAGAATCAACAACTGT) and 3’ primer (ACAGTTGTTGATTCTACAGTCTGCAGC) were used to create the Psnb-1::UNC-13L(K49E) construct.

Transgenic strains were isolated by microinjection of various plasmids using either Pmyo-2::NLS-GFP (KP#1106) or Pmyo-2::NLS-mCherry (KP#1480) as the co-injection marker. Integrated transgenes were obtained by UV irradiation of strains carrying extrachromosomal arrays. All integrated transgenes were outcrossed at least seven times.

Electrophysiology
Electrophysiology was conducted on dissected C. elegans as previously described (Hu et al., 2012). Worms were superfused in an extracellular solution containing 127 mM NaCl, 5 mM KCl, 26 mM NaHCO\(_3\), 1.25 mM Na\(_2\)HPO\(_4\), 20 mM glucose, 1 mM CaCl\(_2\), and 4 mM MgCl\(_2\), bubbled with 5% CO\(_2\), 95% O\(_2\) at 22°C. The 1 mM CaCl\(_2\) was replaced by 1 mM MgCl\(_2\) to record mEPSCs and mLIPSCs in 0 mM of Ca\(^{2+}\). Whole-cell recordings were carried out at −60 mV for all EPSCs, including mEPSCs, evoked EPSCs, and sucrose-evoked responses. The holding potential was switched to 0 mV to record mLIPSCs. The internal solution contained 105 mM CH\(_3\)O\(_2\)SCs, 10 mM CsCl, 15 mM CsF, 4 mM
MgCl₂, 5 mM EGTA, 0.25 mM CaCl₂, 10 mM HEPES, and 4 mM Na₂ATP, adjusted to pH 7.2 using CsOH. Stimulus-evoked EPSCs were stimulated by placing a borosilicate pipette (5–10 μm) near the ventral nerve cord (one muscle distance from the recording pipette) and applying a 0.4 ms, 85 μA square pulse using a stimulus current generator (WPI). A pulsed application of sucrose (1M, 2 s) was applied onto the ventral nerve cord using a Picospritzer III (Parker). The glass pipette containing sucrose was placed at the end of the patched muscle (around half muscle distance from the recording pipette).

**Fluorescence imaging**

Animals were immobilized on 2% agarose pads with 30 mM levamisole. Fluorescence imaging was performed on a spinning-disk confocal system (3i Yokogawa W1 SDC) controlled by Slidebook 6.0 software. Animals were imaged with an Olympus 100 × 1.4 NA Plan-Apochromat objective. Z series of optical sections were acquired at 0.11 μm steps. Images were deconvolved with Huygens Professional version 16.10 (Scientific Volume Imaging, The Netherlands) and then processed to yield maximum intensity projections using ImageJ 1.51 n (Wayne Rasband, National Institutes of Health).

**Isothermal titration calorimetry (ITC)**

ITC experiments were performed using a Nano ITC (TA Instruments). ITC titrations were performed in degassed 20 mM HEPES pH 7.5, 150 mM NaCl at 25°C. After baseline subtraction, the area under each peak was integrated and data were fit to a one site independent binding model using NanoAnalyze software (TA Instruments) to determine the binding affinity (K), enthalpy (dH) and stoichiometry (n); binding was not observed for unc-10 62–121 K77/79E.

**Protein expression and purification (for ITC experiments)**: *C. elegans* UNC-13 long isoform residues 1–258 bearing the K49E mutation (which abolishes homodimer formation), *C. elegans* UNC-10 residues 62–121 WT or *C. elegans* UNC-10 K77/79E were expressed from pGEX6p1 as a GST fusion in Rosetta2 (DE3) cells. Cells were grown at 37°C in 6L 2xyt with carbenicillin (100 μg/mL), chloramphenicol (35 μg/mL) and ZnSO₄ (200 μM, Unc-10 constructs only), to O.D.600 ~ 0.5, then allowed to cool down to 18°C for an hour before induction with 1 mM IPTG overnight at 18°C. The cells were harvested the next day and lysed in 20 mM HEPES (pH 7.5), 150 mM NaCl, 1 mM PMSF, 1% Triton X-100, 1 mM DTT, 200 μM ZnSO₄ (UNC-10 constructs only); the cells were sonicated 3 × 40 s using a midi tip, and lysates clarified 45 mins @ 40,000 rpm, 4°C in a Sorvall Type 45 Ti rotor. Clarified lysates were bound to 1.5 mL (bed volume) glutathione sepharose beads tumbling overnight at 4°C. The next day, the beads were washed extensively, and HRV3C protease added for on-beads cleavage overnight at 4°C. The eluted soluble constructs were concentrated and subjected to gel filtration on a Superdex 75 10/300 against 20 mM HEPES pH 7.5 150 mM NaCl at 4°C. Single peaks corresponding to the monomeric proteins were concentrated to 21 μM (UNC-13L residues 1–258), to 140 μM (UNC-10 62–121 WT) and to 116 μM (UNC-10 62–121 K77/79E mutant), and used in the ITC experiment.

**Data acquisition and statistical analysis**

All recordings were obtained using a HEKA EPC10 double amplifier (HEKA Elektronik) filtered at 2 kHz, and analyzed by built-in programs in Igor seven software (WaveMetrics). Each set of data represents the mean ± SEM of an indicated number (n) of animals. Statistical significance was determined using student’s-t test or one-way ANOVA followed by Dunnett’s test to control for multiple comparisons.

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Author contributions

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References

Aravamudan B, Fergestad T, Davis WS, Rodesch CK, Broadie K. 1999. Drosophila UNC-13 is essential for synaptic transmission. Nature Neuroscience 2:965–971. DOI: https://doi.org/10.1038/14764, PMID: 10526334
Augustin I, Rosenmund C, Südhof TC, Brose N. 1999. Munc13-1 is essential for fusion competence of glutamatergic synaptic vesicles. Nature 400:457–461. DOI: https://doi.org/10.1038/22768, PMID: 10440375
Bacaj T, Wu D, Bürre J, Malenka RC, Liu X, Südhof TC. 2015. Synaptotagmin-1 and -7 are redundantly essential for maintaining the capacity of the Readily-Releaseable pool of synaptic vesicles. PLOS Biology 13:e1002267. DOI: https://doi.org/10.1371/journal.pbio.1002267, PMID: 26437117
Basu J, Betz A, Brose N, Rosenmund C. 2007. Munc13-1 C1 domain activation lowers the energy barrier for synaptic vesicle fusion. Journal of Neuroscience 27:1200–1210. DOI: https://doi.org/10.1523/JNEUROSCI.4908-06.2007, PMID: 17267576
Betz A, Thakur P, Junge HJ, Ashery U, Rhee JS, Scheuss V, Rosenmund C, Retig J, Brose N. 2001. Functional interaction of the active zone proteins Munc13-1 and RIM1 in synaptic vesicle priming. Neuron 30:183–196. DOI: https://doi.org/10.1016/S0896-6273(01)00272-0, PMID: 11343654

Brenner S. 1974. The genetics of caenorhabditis elegans. Genetics 77:71–94. PMID: 4366476

Brose N, Hofmann K, Hata Y, Südhof TC. 1995. Mammalian homologues of caenorhabditis elegans unc-13 gene define novel family of C2-domain proteins. The Journal of Biological Chemistry 270:25273–25280. DOI: https://doi.org/10.1074/jbc.270.42.25273, PMID: 7559667

Camacho M, Basu J, Trimbuch T, Chang S, Pulido-Lozano C, Chang S-S, Duluvova I, Abo-Rady M, Rizo J, Rosenmund C. 2017. Heterodimerization of Munc13 C2a domain with RIM regulates synaptic vesicle docking and priming. Nature Communications 8:15293. DOI: https://doi.org/10.1038/scomms15293

Chen Z, Cooper B, Kalla S, Varoquaux F, Young SM. 2013. The Munc13 proteins differentially regulate readily releasable pool dynamics and calcium-dependent recovery at a central synapse. Journal of Neuroscience 33:8336–8351. DOI: https://doi.org/10.1523/JNEUROSCI.5128-12.2013, PMID: 23658173

Deng L, Kaeser PS, Xu W, Südhof TC. 2011. RIM proteins activate vesicle priming by reversing autoinhibitory homodimerization of Munc13. Neuron 69:317–331. DOI: https://doi.org/10.1016/j.neuron.2011.01.005, PMID: 21262469

Dulubova I, Lou X, Lu J, Huryeva I, Alam A, Schneggerenburger R, Südhof TC, Rizo J. 2005. A Munc13/RIM/ Rab3 tripartite complex: from priming to plasticity? The EMBO Journal 24:2839–2850. DOI: https://doi.org/10.1038/sj.emboj.7600753, PMID: 16052212

Gracheva EQ, Hadwiger G, Nonet ML, Richmond JE. 2008. Direct interactions between C. elegans RAB-3 and rim provide a mechanism to target vesicles to the presynaptic density. Neuroscience Letters 444:137–142. DOI: https://doi.org/10.1016/j.neulet.2008.08.026, PMID: 18721860

Hammarslund M, Palfreyman MT, Watanabe S, Olsen S, Jorgensen EM. 2007. Open syntaxin docks synaptic vesicles. PLOS Biology 5:e198. DOI: https://doi.org/10.1371/journal.pbio.0050198, PMID: 17645391

Han Y, Kaeser PS, Südhof TC, Schneggerenburger R. 2011. RIM determines Ca2+ channel density and vesicle docking at the presynaptic active zone. Neuron 69:304–316. DOI: https://doi.org/10.1016/j.neuron.2010.12.014, PMID: 21262468

Hu Z, Hom S, Kudze T, Tong XJ, Choi S, Aramuni G, Zhang W, Kaplan JM. 2012. Neurexin and neulin regulate retrograde synaptic inhibition in C. elegans. Science 337:980–984. DOI: https://doi.org/10.1126/science.1224896, PMID: 22859820

Hu Z, Tong XJ, Kaplan JM. 2013. UNC-13L, UNC-13S, and tomosyn form a protein code for fast and slow neurotransmitter release in caenorhabditis elegans. eLife 2:e00967. DOI: https://doi.org/10.7554/eLife.00967, PMID: 23951547

Imig C, Min SW, Krinner S, Arancillo M, Rosenmund C, Südhof TC, Rhee J, Brose N, Cooper BH. 2014. The morphological and molecular nature of synaptic vesicle priming at Presynaptic active zones. Neuron 84:416–431. DOI: https://doi.org/10.1016/j.neuron.2014.10.009, PMID: 25374362

Jahn R, Fasshauer D. 2012. Molecular machines governing exocytosis of synaptic vesicles. Nature 490:201–207. DOI: https://doi.org/10.1038/nature11320, PMID: 23060190

Kaeser PS, Deng L, Wang Y, Dulubova I, Liu X, Rizo J, Südhof TC. 2011. RIM proteins tether Ca2+ channels to presynaptic active zones via a direct PDZ-domain interaction. Cell 144:282–295. DOI: https://doi.org/10.1016/j.cell.2010.12.029, PMID: 21241893

Kouchkina SP, Richmond JE, Hadwiger G, Weimer RM, Jorgensen EM, Nonet ML. 2001. A post-docking role for the SNARE complex. Nature Neuroscience 4:997–1005. DOI: https://doi.org/10.1038/nn732, PMID: 11559854

Lipstein N, Verhoeven-Duijf NM, Michelassi FE, Calloway N, van Hasselt PM, Pienkowska K, van Haften G, van Haelst MM, van Empelen R, Cuppen I, van Teeseling HC, Evelein AM, Vorstman JA, Thoms S, Jahn O, Duran KJ, Monroe GR, Ryan TA, Taschenberger H, Dittman JS, et al. 2017. Synaptic UNC13A protein variant causes increased neurotransmission and dyskinetic movement disorder. Journal of Clinical Investigation 127:1005–1018. DOI: https://doi.org/10.1172/JCI90259, PMID: 28192369

Liu Q, Hollopeter G, Jorgensen EM. 2009. Graded synaptic transmission at the caenorhabditis elegans neuromuscular junction. PNAS 106:10823–10828. DOI: https://doi.org/10.1073/pnas.0903570106, PMID: 19528650

Liu H, Li L, Wang W, Gong J, Yang X, Hu Z. 2018. Spontaneous vesicle fusion is differentially regulated by cholinerigic and GABAergic synapses. Cell Reports 22:2334–2345. DOI: https://doi.org/10.1016/j.celrep.2018.02.023, PMID: 29490270

Lu J, Machius M, Dulubova I, Dai H, Südhof TC, Tomchick DR, Rizo J. 2006. Structural basis for a Munc13-1 homodimer to Munc13-1/RIM heterodimer switch. PLOS Biology 4:e192. DOI: https://doi.org/10.1371/journal.pbio.0040192, PMID: 16732694

Luo F, Bacaj T, Südhof TC. 2015. Synaptotagmin-7 is essential for Ca2+-Triggered delayed asynchronous release but not for Ca2+-Dependent vesicle priming in retinal ribbon synapses. Journal of Neuroscience 35:11024–11033. DOI: https://doi.org/10.1523/JNEUROSCI.0759-15.2015, PMID: 26245964

Ma C, Li W, Xu Y, Rizo J. 2011. Munc13 mediates the transition from the closed syntaxin-Munc18 complex to the SNARE complex. Nature Structural & Molecular Biology 18:542–549. DOI: https://doi.org/10.1038/nsmb.2047, PMID: 21499244

Melom JE, Akbergenoyna Y, Gavornik JP, Littleton JT. 2013. Spontaneous and evoked release are independently regulated at individual active zones. Journal of Neuroscience 33:17253–17263. DOI: https://doi.org/10.1523/JNEUROSCI.3334-13.2013, PMID: 24174659
Michelassi F, Liu H, Hu Z, Dittman JS. 2017. A C1-C2 module in Munc13 inhibits Calcium-Dependent neurotransmitter release. Neuron 95:577–590. DOI: https://doi.org/10.1016/j.neuron.2017.07.015, PMID: 28772122

Müller M, Liu KS, Sigrist SJ, Davis GW. 2012. RIM controls homeostatic plasticity through modulation of the readily-releasable vesicle pool. Journal of Neuroscience 32:16574–16585. DOI: https://doi.org/10.1523/JNEUROSCI.0981-12.2012, PMID: 23175813

Neher E, Sakaba T. 2008. Multiple roles of calcium ions in the regulation of neurotransmitter release. Neuron 59:861–872. DOI: https://doi.org/10.1016/j.neuron.2008.08.019, PMID: 18817727

Pang ZP, Sun J, Rizo J, Maximov A, Südhof TC. 2006. Genetic analysis of synaptotagmin 2 in spontaneous and Ca2+-triggered neurotransmitter release. The EMBO Journal 25:2039–2050. DOI: https://doi.org/10.1038/sj.emboj.7601103, PMID: 16642042

Ramirez DM, Kavalali ET. 2011. Differential regulation of spontaneous and evoked neurotransmitter release at central synapses. Current Opinion in Neurobiology 21:275–282. DOI: https://doi.org/10.1016/j.conb.2011.01.007, PMID: 21334193

Richmond JE, Davis WS, Jorgensen EM. 1999. UNC-13 is required for synaptic vesicle fusion in C. elegans. Nature Neuroscience 2:959–964. DOI: https://doi.org/10.1038/14755, PMID: 10526333

Song Y, Ailenberg M, Silverman M. 1998. Cloning of a novel gene in the human kidney homologous to rat munc13s: its potential role in diabetic nephropathy. Kidney International 53:1689–1695. DOI: https://doi.org/10.1046/j.1523-1755.1998.00942.x, PMID: 9467201

Südhof TC. 2004. The synaptic vesicle cycle. Annual Review of Neuroscience 27:509–547. DOI: https://doi.org/10.1146/annurev.neuro.26.041002.131412, PMID: 15217342

Südhof TC, Rizo J. 2011. Synaptic vesicle exocytosis. Cold Spring Harbor Perspectives in Biology 3:a005637. DOI: https://doi.org/10.1101/cshperspect.a005637, PMID: 22026965

Watanabe S, Liu Q, Davis MW, Hollopeter G, Thomas N, Jorgensen NB, Jorgensen EM. 2013. Ultrafast endocytosis at Caenorhabditis Elegans neuromuscular junctions. eLife 2:e00723. DOI: https://doi.org/10.7554/eLife.00723, PMID: 24015355

Weimer RM, Gracheva EO, Meyrignac O, Miller KG, Richmond JE, Bessereau JL. 2006. UNC-13 and UNC-10/rim localize synaptic vesicles to specific membrane domains. Journal of Neuroscience 26:8040–8047. DOI: https://doi.org/10.1523/JNEUROSCI.2350-06.2006, PMID: 16885217

Williams C, Chen W, Lee CH, Yaeger D, Vyleta NP, Smith SM. 2012. Coactivation of multiple tightly coupled calcium channels triggers spontaneous release of GABA. Nature Neuroscience 15:1195–1197. DOI: https://doi.org/10.1038/nn.3162, PMID: 22842148

Xu J, Camacho M, Xu Y, Esser V, Liu X, Trimbuch T, Pan YZ, Ma C, Tomchick DR, Rosenmund C, Rizo J. 2017. Mechanistic insights into neurotransmitter release and presynaptic plasticity from the crystal structure of Munc13-1 C1C2BMUN. eLife 6:e22567. DOI: https://doi.org/10.7554/eLife.22567, PMID: 28177287

Yang X, Wang S, Sheng Y, Zhang M, Zou W, Wu L, Kang L, Rizo J, Zhang R, Xu T, Ma C. 2015. Syntaxin opening by the MUN domain underlies the function of Munc13 in synaptic-vesicle priming. Nature Structural & Molecular Biology 22:547–554. DOI: https://doi.org/10.1038/nsmb.3038

Zhou K, Stawicki TM, Goncharov A, Jin Y. 2013a. Position of UNC-13 in the active zone regulates synaptic vesicle release probability and release kinetics. eLife 2:e01180. DOI: https://doi.org/10.7554/eLife.01180, PMID: 24220508

Zhou P, Bacaj T, Yang X, Pang ZP, Südhof TC. 2013b. Lipid-anchored SNAREs lacking transmembrane regions fully support membrane fusion during neurotransmitter release. Neuron 80:470–483. DOI: https://doi.org/10.1016/j.neuron.2013.09.010, PMID: 24120845

Zhou P, Pang ZP, Yang X, Zhang Y, Rosenmund C, Bacaj T, Südhof TC. 2013c. Syntaxin-1 N-peptide and Habc-domain perform distinct essential functions in synaptic vesicle fusion. The EMBO Journal 32:159–171. DOI: https://doi.org/10.1038/emboj.2012.307, PMID: 23188083