

# Phosphorylation of Complexin by PKA Regulates Activity-Dependent Spontaneous Neurotransmitter Release and Structural Synaptic Plasticity

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## SUMMARY

Synaptic plasticity is a fundamental feature of the nervous system that allows adaptation to changing behavioral environments. Most studies of synaptic plasticity have examined the regulated trafficking of postsynaptic glutamate receptors that generates alterations in synaptic transmission. Whether and how changes in the presynaptic release machinery contribute to neuronal plasticity is less clear. The SNARE complex mediates neurotransmitter release in response to presynaptic  $\text{Ca}^{2+}$  entry. Here we show that the SNARE fusion clamp Complexin undergoes activity-dependent phosphorylation that alters the basic properties of neurotransmission in *Drosophila*. Retrograde signaling following stimulation activates PKA-dependent phosphorylation of the Complexin C terminus that selectively and transiently enhances spontaneous release. Enhanced spontaneous release is required for activity-dependent synaptic growth. These data indicate that SNARE-dependent fusion mechanisms can be regulated in an activity-dependent manner and highlight the key role of spontaneous neurotransmitter release as a mediator of functional and structural plasticity.

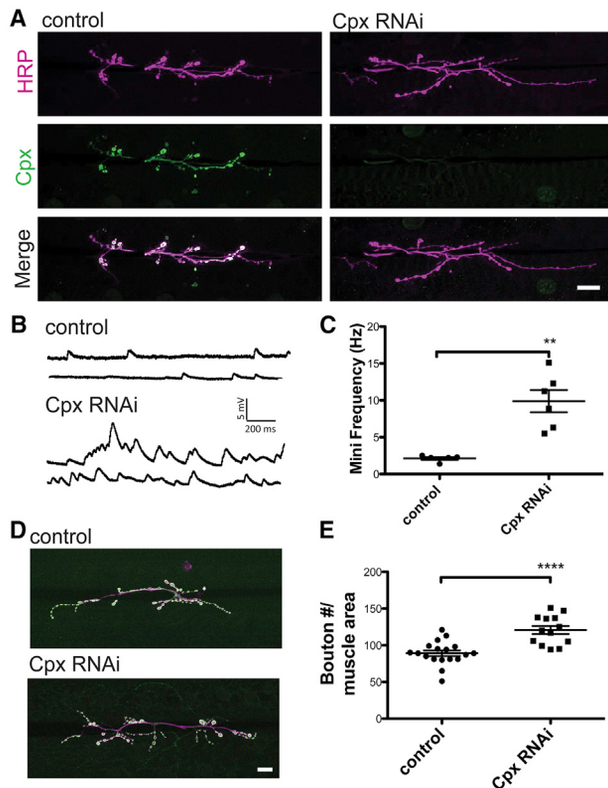
## INTRODUCTION

Synaptic transmission is initiated by the rapid fusion of docked synaptic vesicles via  $\text{Ca}^{2+}$  influx in response to an action potential (AP). Synaptic vesicles can also fuse independent of APs, resulting in individual spontaneous quantal events termed minis (Fatt and Katz, 1952). Recent studies suggest spontaneous release may function independently as a regulated channel for neuronal signaling distinct from AP-driven release. Minis have been shown to activate specific biochemical signaling pathways that regulate structural and homeostatic plasticity (Andreae and Burrone, 2015; Choi et al., 2014; Lee et al., 2010; McKinney et al., 1999; Sutton et al., 2006). Moreover, minis modulate AP firing in

certain neurons, suggesting spontaneous release can contribute to the integration of signals within neuronal ensembles (Carter and Regehr, 2002). Finally, accumulating evidence suggests that regulation of spontaneous versus evoked release can occur via distinct molecular interactions at the level of the fusion machinery (Cho et al., 2010; Kaeser-Woo et al., 2012; Kavalali, 2015; Martin et al., 2011).

Synaptic vesicle fusion is mediated by the regulated assembly of the SNARE complex to generate docked and primed synaptic vesicles (Südhof and Rothman, 2009). Zippering of the SNAREs into a four-helix bundle provides the energy necessary to drive bilayer fusion (Weber et al., 1998). At the synapse, SNARE assembly and zippering is tightly controlled to achieve the precision and speed required for effective neurotransmission. One such regulator is Synaptotagmin 1 (Syt 1), a  $\text{Ca}^{2+}$  sensor for AP-dependent synaptic vesicle fusion (Geppert et al., 1994; Littleton et al., 1993). In addition to Syt 1, the SNARE-binding protein Complexin (Cpx) has emerged as an important co-regulator for vesicle fusion (McMahon et al., 1995; Südhof, 2013). In *in vitro* cell-cell fusion and lipid mixing assays, Cpx reduces membrane fusion, acting as a clamp to block premature exocytosis in the absence of  $\text{Ca}^{2+}$  (Giraud et al., 2006; Malsam et al., 2012; Schaub et al., 2006). Genetic studies in *Drosophila* and *C. elegans* revealed a dramatic increase in minis in *cpx* null mutants (Hobson et al., 2011; Huntwork and Littleton, 2007; Jorquera et al., 2012; Martin et al., 2011). In addition to preventing spontaneous release, Cpx facilitates AP-dependent vesicle fusion (Hobson et al., 2011; Huntwork and Littleton, 2007; Martin et al., 2011; Xue et al., 2008). Structure-function studies indicate Cpx regulates minis and AP-dependent release via distinct molecular mechanisms (Cho et al., 2010, 2014; Hobson et al., 2011; Kaeser-Woo et al., 2012; Martin et al., 2011; Maximov et al., 2009). Binding of the Cpx central helix to the SNARE complex is required for both clamping and stimulating properties (Cho et al., 2010; Xue et al., 2010), while the C terminus is required for clamping spontaneous fusion (Buhl et al., 2013; Cho et al., 2010; Kaeser-Woo et al., 2012; Wragg et al., 2013). Interestingly, the C terminus of mammalian Cpx I and II is phosphorylated *in vivo*, suggesting its function might be regulated through post-translational modifications (Shata et al., 2007).

The *Drosophila* neuromuscular junction (NMJ) undergoes acute forms of activity-dependent structural and functional plasticity, providing a model synapse to examine the role of



**Figure 1. Cpx Knockdown Larvae Exhibit Increased Spontaneous Release and Enhanced Synaptic Growth**

(A) Expression of Cpx protein was reduced in Cpx RNAi knockdown lines ( $w^{1118}; c164-GAL4/+; Cpx\ RNAi/+$ ) compared to control ( $w^{1118}; c164-GAL4/+$ ) using a motor neuron driver ( $c164-GAL4$ ) to express Cpx-specific RNAi. Larvae were co-stained with anti-HRP and anti-Cpx specific antisera. Images were obtained from muscle 6/7 NMJs of 3<sup>rd</sup> instar larvae.

(B) Sample traces of spontaneous release from control and Cpx RNAi knockdown larvae.

(C) Mean mini frequency (Hz  $\pm$  SEM) recorded for each line.

(D) Representative images of NMJs from control and Cpx RNAi knockdown larvae. Larvae were stained with anti-HRP- (magenta) and anti-GluRIII-specific (green) antisera to identify synaptic boutons.

(E) Mean synaptic bouton number for control and Cpx RNAi knockdowns. Bouton number was normalized using muscle 6/7 area. Indicated comparisons were made using Student's *t* tests. Mean  $\pm$  SEM is indicated in all figures. Scale bar, 20  $\mu$ m.

the release machinery in synaptic plasticity (Ataman et al., 2008; Ballard et al., 2014; Berke et al., 2013; Frank et al., 2009; Goold and Davis, 2007; McCabe et al., 2003; Piccioli and Littleton, 2014). We previously identified a retrograde signaling pathway that drives synaptic plasticity at this glutamatergic synapse. High-frequency stimulation of motor neurons induces a long-lasting enhancement of spontaneous release, a process termed high-frequency-induced miniature release (HFMR). At embryonic NMJs, HFMR induction results in a 100-fold increase in minis for several minutes before returning to baseline (Yoshihara et al., 2005). This process is dependent on the presynaptic cAMP-PKA kinase signaling pathway and postsynaptic Syt 4, a Ca<sup>2+</sup>-sensitive regulator of postsynaptic vesicle fusion (Yoshihara et al., 2005). HFMR is also expressed at mature larval

NMJs and is required for activity-dependent synaptic growth (Barber et al., 2009; Piccioli and Littleton, 2014; Yoshihara et al., 2005). Interestingly, *cpx* null mutants exhibit enhanced synaptic growth in addition to enhanced minis (Huntwork and Littleton, 2007). Indeed, minis have recently been found to be necessary and sufficient to drive developmental synaptic growth at *Drosophila* NMJs (Choi et al., 2014). These results suggest that HFMR and synaptic growth signals may be mechanistically linked through Syt 4-dependent retrograde signaling.

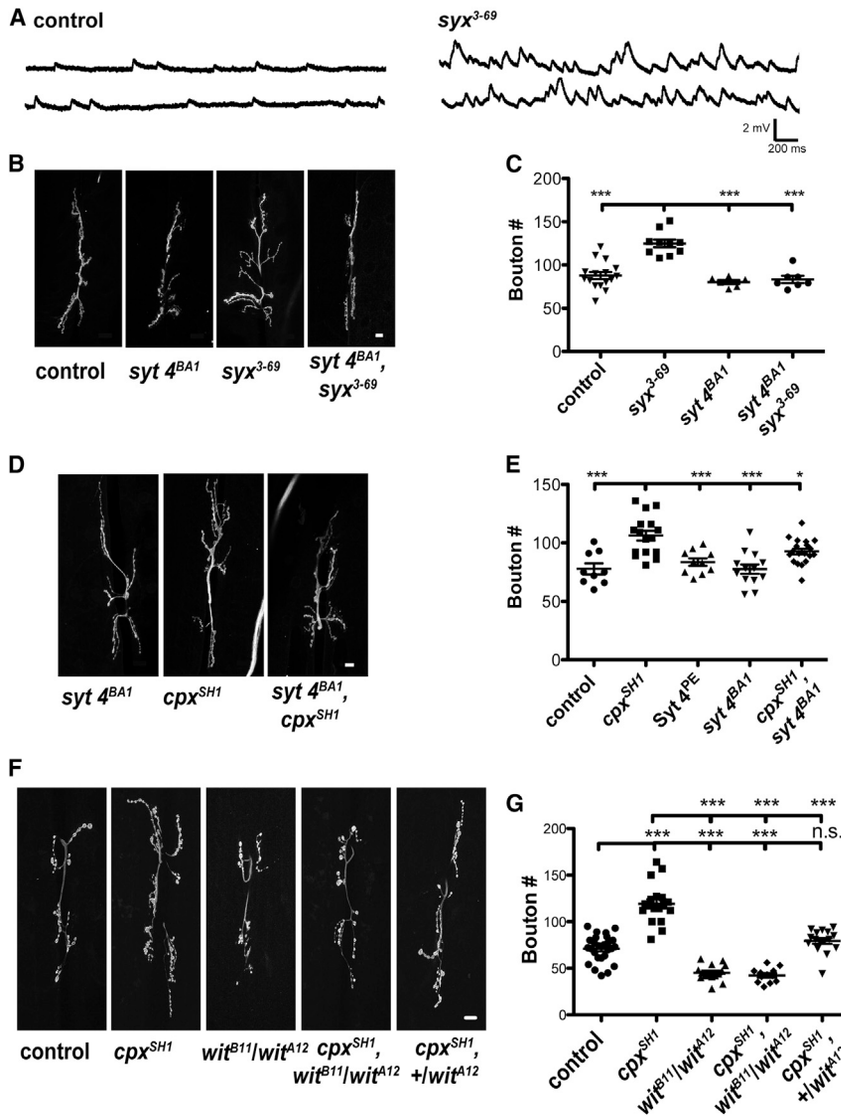
Here we demonstrate that minis are acutely regulated by activity-dependent phosphorylation of the C terminus of the fusion clamp, Cpx. PKA-dependent phosphorylation of Cpx is required for synaptic structural plasticity and growth. Our findings suggest that Syt 4-dependent retrograde signaling regulates pre-synaptic PKA-dependent phosphorylation of Cpx to modulate its clamping properties, allowing short-term increases in spontaneous neurotransmitter release to control functional and structural plasticity.

## RESULTS

### Synaptic Growth Correlates with Spontaneous Neurotransmitter Release Rate

We previously observed that *cpx*<sup>SH1</sup> null mutants exhibit enhanced minis at 3<sup>rd</sup> instar larval NMJs. Increased spontaneous release was correlated with enhanced synaptic growth (Choi et al., 2014; Huntwork and Littleton, 2007), suggesting that minis may play a unique role in regulating the synaptic growth machinery at *Drosophila* synapses. To examine the role of minis in regulating synaptic growth, we used UAS-RNAi against Cpx to specifically remove the protein in motor neurons using the *c164-GAL4* driver (Figure 1A). This approach allowed Cpx to be reduced in a motor neuron-specific manner, with minis enhanced to more physiological levels compared to the  $\sim$ 70–80 Hz range observed in the complete absence of Cpx. Electrophysiological recordings from 3<sup>rd</sup> instar larvae revealed enhanced mini frequency in Cpx knockdowns compared to controls ( $w^{1118}; c164-GAL4/+; Cpx\ RNAi/+ = 9.9 \pm 1.5$  Hz [ $n = 6$ ] versus  $w^{1118}; c164-GAL4/+ = 2.1 \pm 0.2$  Hz [ $n = 5$ ];  $p < 0.01$ ; Figures 1B and 1C). Similar to synaptic overgrowth observed in *cpx*<sup>SH1</sup> nulls (see also Figures 2E, 2G, and 5G), Cpx knockdown larvae exhibited enhanced synaptic growth at muscle 6/7 of 3<sup>rd</sup> instar larvae ( $w^{1118}; c164-GAL4/+; Cpx\ RNAi/+ = 120.8 \pm 5.5$  boutons [ $n = 13$ ] versus  $w^{1118}; c164-GAL4/+ = 89.2 \pm 3.8$  boutons [ $n = 18$ ];  $p < 0.0001$ ; Figures 1D and 1E).

To determine whether enhanced synaptic growth was unique to the *cpx* mutant and Cpx knockdowns that exhibit elevated mini frequencies, we examined synapses of an independent mutant that also exhibits elevated spontaneous release. *syx*<sup>3-69</sup> mutants harbor a point mutation in the t-SNARE syntaxin 1A (Littleton et al., 1998) that gives rise to elevated mini rates ( $w^{1118}; syx^{3-69} = 14.2 \pm 1.3$  Hz [ $n = 8$ ]) similar to those observed in Cpx RNAi animals ((Bykhovskaia et al., 2013; Lagow et al., 2007) (Figure 2A). Consistent with the *cpx*<sup>SH1</sup> mutant and RNAi Cpx knockdown, synapses in *syx*<sup>3-69</sup> mutants exhibited enhanced growth compared to control ( $w^{1118}; syx^{3-69} = 124.8 \pm 4.4$  boutons [ $n = 10$ ] versus control [ $w^{1118}$ ] = 87.8  $\pm$  3.9 boutons [ $n = 16$ ];  $p < 0.001$ ; Figures 2B and 2C). Together, these results suggest



that elevated rates of spontaneous release positively correlate with synaptic growth at *Drosophila* NMJs.

### A Syt 4-Mediated Retrograde Signaling Pathway Is Required for Mini-Dependent Synaptic Growth

Retrograde signaling from postsynaptic compartments regulates both synaptic growth and homeostatic plasticity during larval development (Goold and Davis, 2007; Marqués et al., 2002; McCabe et al., 2003; Paradis et al., 2001). We previously identified a retrograde signaling pathway that controls spontaneous release rates at *Drosophila* NMJs mediated by Syt 4, a  $Ca^{2+}$ -sensitive regulator of postsynaptic vesicle fusion. To examine if this pathway is required for the enhanced synaptic growth in animals that exhibit enhanced minis, we generated double mutants of *syx*<sup>3-69</sup> with *syt 4* null mutants (*syt 4*<sup>BA1</sup>). Introducing the *syt 4*<sup>BA1</sup> mutant allele into the *syx*<sup>3-69</sup> background suppressed synaptic growth ( $w^{1118}; syx^{3-69} = 124.8 \pm 4.4$  boutons [n = 10] versus  $w^{1118}; syx^{3-69}, syt 4^{BA1} = 80.2 \pm 2.3$  boutons

### Figure 2. Synaptic Growth Associated with Enhanced Minis Is Suppressed by Synaptotagmin 4 (*syt 4*<sup>BA1</sup>) and BMP Receptor (*wit*) Mutants

(A) Representative traces of spontaneous release recorded from syntaxin 1A mutant ( $w^{1118}; syx^{3-69}$ ) and control ( $w^{1118}$ ) larval muscle 6 NMJs.

(B) Representative images of muscle 6/7 NMJs of control ( $w^{1118}$ ), *syx*<sup>3-69</sup> mutant ( $w^{1118}; syx^{3-69}$ ), *syt 4*<sup>BA1</sup> null ( $w^{1118}; syt 4^{BA1}$ ), and *syx*<sup>3-69</sup>, *syt 4*<sup>BA1</sup> double mutant ( $w^{1118}; syx^{3-69}, syt 4^{BA1}$ ).

(C) Mean synaptic bouton number measured for indicated genotypes.

(D) Representative images of muscle 6/7 NMJs of the indicated genotypes.

(E) Mean synaptic bouton number measured for control ( $w^{1118}; Cpx^{PE}$ ), *cpx*<sup>SH1</sup> null ( $w^{1118}; cpx^{SH1}$ ), control *Syt 4*<sup>PE</sup> ( $w^{1118}; Syt 4^{PE}$ ), *syt 4*<sup>BA1</sup> null ( $w^{1118}; syt 4^{BA1}$ ), and *syx*<sup>3-69</sup>, *syt 4*<sup>BA1</sup> double mutant ( $w^{1118}; syt 4^{BA1}, cpx^{SH1}$ ).

(F) Representative images of muscle 6/7 NMJs from control ( $w^{1118}; Cpx^{PE}$ ), *cpx*<sup>SH1</sup> null, *wit* null ( $w^{1118}; wit^{A12}/wit^{B11}$ ), *wit*, *cpx*<sup>SH1</sup> double mutant ( $w^{1118}; cpx^{SH1}, wit^{A12}/cpx^{SH1}, wit^{B11}$ ), and *wit* heterozygotes in *cpx*<sup>SH1</sup> null background ( $w^{1118}; cpx^{SH1}, wit^{A12}/cpx^{SH1}$ ).

(G) Mean synaptic bouton number measured for the indicated genotypes. Indicated comparisons were made using ANOVA with post hoc Tukey analysis. Mean  $\pm$  SEM is indicated in all figures. Scale bars, 10  $\mu$ m.

[n = 6];  $p < 0.001$ ; Figures 2B and 2C). We next examined if Syt 4 is similarly required for synaptic growth in the *cpx*<sup>SH1</sup> null background. *cpx*<sup>SH1</sup> nulls exhibit greatly enhanced minis ( $w^{1118}; cpx^{SH1} = 70.4 \pm 3.5$  Hz [n = 7]) as well as enhanced synaptic growth compared to control ( $w^{1118}; cpx^{SH1} = 106.3 \pm 4.2$  boutons [n = 16] versus  $w^{1118}; Cpx^{PE}$  [precise excision control for *cpx*<sup>SH1</sup>] =  $77.8 \pm 2.8$  boutons

[n = 9];  $p < 0.001$ ; Figures 2D and 2E). Loss of Syt 4 alone did not show any elevation in synaptic growth compared to both *cpx*<sup>SH1</sup> and *syt 4*<sup>BA1</sup> controls ( $w^{1118}; syt 4^{BA1} = 77.5 \pm 4.0$  boutons [n = 13] versus ( $w^{1118}; Cpx^{PE} = 77.8 \pm 4.7$  boutons [n = 9] and  $w^{1118}; Syt 4^{PE}$  [precise excision control for *syt 4*<sup>BA1</sup>] =  $83.6 \pm 3.2$  boutons [n = 10]);  $p > 0.05$  for both comparisons; Figures 2D and 2E). Introducing the *syt 4*<sup>BA1</sup> allele into the *cpx*<sup>SH1</sup> background reduced synaptic growth compared to *cpx*<sup>SH1</sup> allele alone ( $w^{1118}; syt 4^{BA1}, cpx^{SH1} = 92.6 \pm 2.6$  boutons [n = 19] versus  $w^{1118}; cpx^{SH1} = 106.3 \pm 4.2$  boutons [n = 16];  $p < 0.05$ ; Figures 2D and 2E). Introduction of the *syt 4*<sup>BA1</sup> allele did not suppress enhanced minis in the *cpx*<sup>SH1</sup> or *syx*<sup>3-69</sup> backgrounds (see Table S2). These results indicate that synaptic overgrowth induced by elevated spontaneous release requires the Syt 4-dependent retrograde signaling pathway.

One well-characterized retrograde growth signal at the *Drosophila* NMJ is the bone morphogenic protein (BMP), Gbb (Aberle et al., 2002; Marqués et al., 2002; McCabe et al., 2003).

Gbb is released from muscle and activates the presynaptic type-II BMP receptor wishful thinking (Wit). *wit* null mutants (*wit<sup>A12/wit<sup>B11</sup></sup>*) exhibit reduced synaptic growth at larval NMJs, indicating the BMP retrograde signaling pathway is required for synapse development (Marqués et al., 2002). In addition to its role in synapse development, BMP signaling is also important for acute activity-regulated forms of synaptic structural plasticity through modulation of presynaptic actin cytoskeletal dynamics (Piccioli and Littleton, 2014). Given BMP signaling contributes to both developmental synapse formation and acute structural plasticity, we examined if BMP signaling is required for the enhanced synaptic growth in the context of elevated minis. In the absence of the BMP receptor (*wit<sup>A12/wit<sup>B11</sup></sup>*), synapses are undergrown compared to control (*w<sup>1118</sup>*; *wit<sup>A12/wit<sup>B11</sup></sup>* = 45.1 ± 2.2 boutons [n = 15] versus *w<sup>1118</sup>*; *Cpx<sup>PE</sup>* = 71.1 ± 2.8 boutons [n = 26]; p < 0.001; Figures 2F and 2G), consistent with previous studies (Aberle et al., 2002; Marqués et al., 2002). Synaptic overgrowth in the *cpx<sup>SH1</sup>* null background was strongly suppressed by the *wit* null mutant (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>* = 119.2 ± 5.0 boutons [n = 18] versus *w<sup>1118</sup>*; *cpx<sup>SH1</sup>*, *wit<sup>A12/cpx<sup>SH1</sup></sup>*, *wit<sup>B11</sup>* = 42.4 ± 2.2 boutons [n = 12]; p < 0.001). Introduction of a single *wit* allele in the *cpx<sup>SH1</sup>* background was also sufficient to suppress growth (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>* = 119.2 ± 5.0 boutons [n = 18] versus *w<sup>1118</sup>*; *cpx<sup>SH1</sup>*, *wit<sup>A12/cpx<sup>SH1</sup></sup>* = 79.4 ± 3.1 boutons [n = 18]; p < 0.001), indicating a dose-dependent genetic interaction. Introducing *wit* alleles did not suppress elevated minis in the *cpx<sup>SH1</sup>* null background (see Table S2). In summary, these results indicate that Syt 4 and BMP signaling pathways contribute to enhanced mini-dependent synaptic growth.

### HFMR Is Transient and Its Expression Is Syt 4 and PKA Dependent

High-frequency stimulation of motor neurons induces a long-lasting enhancement of spontaneous release (HFMR) that is dependent on postsynaptic Syt 4 and the presynaptic cAMP-PKA kinase pathway (Yoshihara et al., 2005). HFMR is also expressed at more mature 3<sup>rd</sup> instar larval NMJs in a Syt 4-dependent manner (Barber et al., 2009). To examine if Cpx regulation of spontaneous release might contribute to this process, we further characterized HFMR in 3<sup>rd</sup> instar larvae. In response to strong stimulation (1 s 100 Hz stimulation 4x, spaced 2 s apart), the frequency of mini events rapidly doubled compared to the baseline prior to stimulation (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>* and *w<sup>1118</sup>*; *Syt 4<sup>PE</sup>* genotypes = 1.9 ± 0.2 fold [n = 6] 100 s after induction; Figures 3A–3C). To determine the kinetics of this activity-dependent plasticity, we extended our recordings up to 8 min after stimulation. HFMR was transient, with spontaneous release rates returning to baseline within ~4 min following induction (Figure 3B). Consistent with prior work, *syt 4<sup>BA1</sup>* null animals did not exhibit HFMR in response to stimulation within our 8 min recording session, indicating HFMR expression is not simply delayed in the absence of Syt 4 (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>* and *w<sup>1118</sup>*; *Syt 4<sup>PE</sup>* genotypes = 1.6 ± 0.1 fold [n = 7] versus *w<sup>1118</sup>*; *syt 4<sup>BA1</sup>* = 1.1 ± 0.1 fold [n = 7] at 100 s after induction; p < 0.001). Given the requirement for the presynaptic cAMP-PKA pathway in embryonic HFMR, we assayed whether PKA was required for activity-induced elevations in mini frequency at mature NMJs as well. Using a pharmacological approach to inhibit the cAMP-PKA pathway, we found

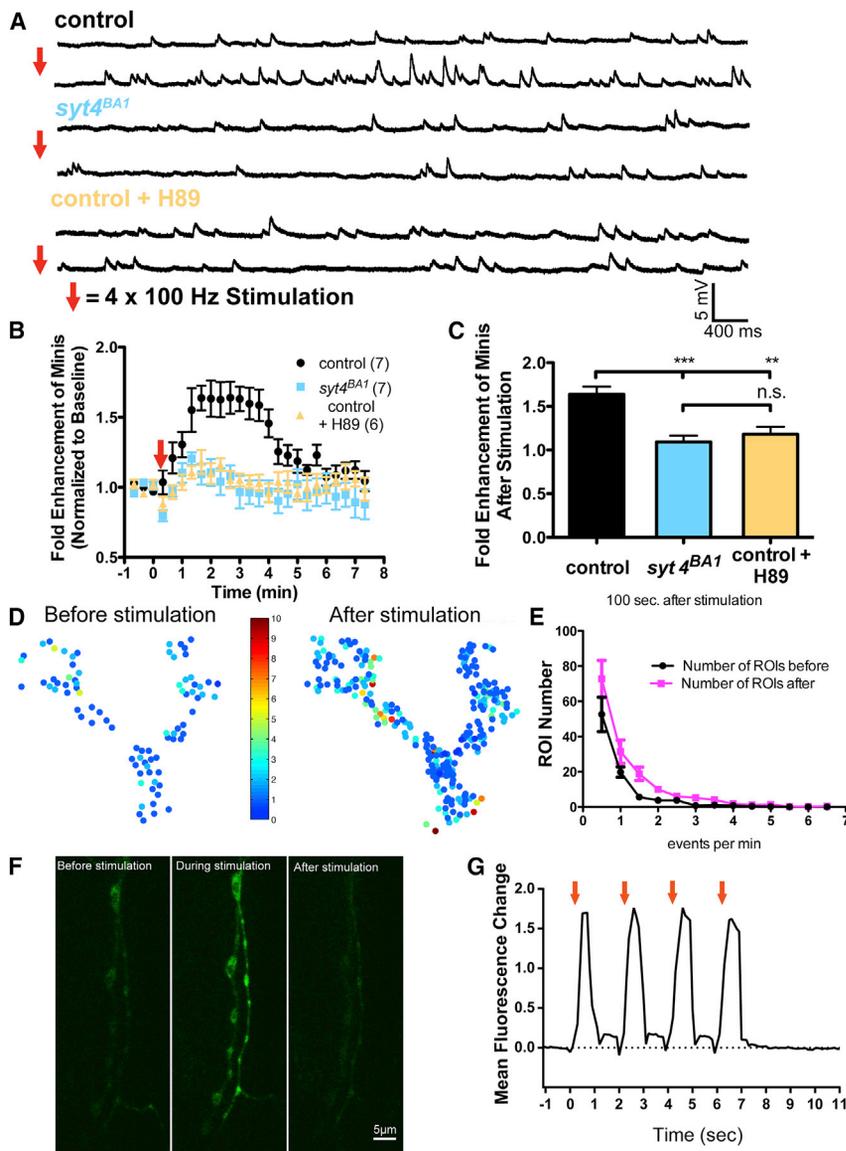
that bath application of H89, a PKA inhibitor, strongly suppressed HFMR compared to controls (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>* and *w<sup>1118</sup>*; *Syt 4<sup>PE</sup>* genotypes = 1.6 ± 0.1 fold [n = 7] versus *w<sup>1118</sup>*; *Syt 4<sup>PE</sup>* + H89 = 1.2 ± 0.1 fold [n = 6] at 100 s after induction; p < 0.01). Together, these results suggest that HFMR expression at mature NMJs requires Syt 4 and PKA.

We next assayed how HFMR is expressed at individual release sites. *Drosophila* NMJs contains spatially segregated active zones that are directly apposed to distinct clusters of glutamate receptors in postsynaptic muscle. By measuring postsynaptic Ca<sup>2+</sup> influx from individual glutamate receptor clusters following release, single fusion events at individual active zones can be monitored (Melom et al., 2013; Peled and Isacoff, 2011). To measure Ca<sup>2+</sup> influx at postsynaptic sites, we employed a genetically encoded myristolated (membrane-tethered) variant of a Ca<sup>2+</sup>-sensitive GFP (GCaMP6s). Postsynaptic expression of UAS-GCaMP6s using the muscle driver *mef2-GAL4* allowed robust detection of postsynaptic responses from both evoked and spontaneous synaptic vesicle fusion events (Melom et al., 2013). Using this method, we quantified individual release events at NMJ synapses for 2 min before and 2 min after HFMR induction (Figures 3D and 3E). A large increase in spontaneous release over a broad population of individual release sites was detected following HFMR induction, including the recruitment of release sites that were previously inactive (Figures 3D–3F). Using this optical method to detect individual release events, we conclude that HFMR is mediated by a broad increase in spontaneous fusion probability over a large population of release sites at the NMJ.

Intracellular presynaptic Ca<sup>2+</sup> modulates mini frequency at certain synapses (Xu et al., 2009). To examine if a long-lasting increase in basal presynaptic Ca<sup>2+</sup> levels occurs after HFMR induction that contributes to enhanced minis beyond retrograde signaling, we monitored presynaptic intracellular Ca<sup>2+</sup> levels using a presynaptically expressed myristolated UAS-GCaMP6s. Intracellular presynaptic Ca<sup>2+</sup> levels rapidly increased during HFMR induction, peaking with each stimulation train (Figures 3F and 3G). Immediately after the HFMR induction protocol, GCaMP6s signal rapidly returned to background levels within a second, indicating prolonged elevated presynaptic Ca<sup>2+</sup> levels are not the cause of the long-lasting increase in mini release that occurs over minutes. The transient local increase in Ca<sup>2+</sup> in response to HFMR induction suggests that long-lasting synaptic modifications downstream of presynaptic Ca<sup>2+</sup> influx, such as PKA phosphorylation of substrates associated with the vesicle fusion machinery, are likely to mediate HFMR.

### PKA Phosphorylation of Cpx Modulates Its Function as a Fusion Clamp

These data indicate retrograde signaling impinges on activation of presynaptic PKA to regulate HFMR. Additionally, the elevation of release rates over a broad population of active zones within minutes of stimulation suggest HFMR expression mechanisms might occur at the post-translational level at individual release sites. The downstream presynaptic targets of PKA that modulate HFMR are unknown but may include components of the SNARE-complex fusion machinery (Südhof and Rothman, 2009). Based



### Figure 3. Activity-Dependent Enhancement of Minis Requires Syt 4 and PKA Activity

(A) Representative traces of minis before (top trace) and after (bottom trace) tetanic stimulation recorded from control ( $w^{1118};; Cpx^{PE}$  and  $w^{1118};; Syt 4^{PE}$ ) and  $syt 4^{BA1}$  nulls ( $w^{1118};; syt 4^{BA1}$ ). Representative traces of minis are also shown for control animals pre-treated with the PKA-specific inhibitor H89.

(B) Mean mini frequency plotted as a function of time. Tetanic stimulation occurs at the 0 s time point (Red Arrow). Mini frequency is normalized to average baseline mini frequency 60 s prior to stimulation for each preparation.

(C) Average fold enhancement of mini frequency compared to baseline mini frequency 100 s after tetanic stimulation. Statistically significant differences in fold enhancement of minis were observed in control animals compared to  $syt 4^{BA1}$  nulls and H89-treated controls at each indicated time point starting at 100 s following HFMR stimulation up to 300 s, when minis started to return to pre-stimulus baseline mini frequency.

(D) Pseudo-color activity heatmaps showing the distribution of spontaneous fusion rates for individual release sites at the same NMJ of muscle 4 prior (left panel) and after (right panel) tetanus stimulation. Each circle is a single region of interest (ROI) that represents the detection of a release event indicated by postsynaptic myrGCaMP6s fluorescence. Color-coding in the heatmaps indicate the number of mini events detected over a period of 2 min before and after HFMR stimulation.

(E) Number of ROIs exhibiting indicated event frequency before (black) and after (magenta) HFMR stimulation.

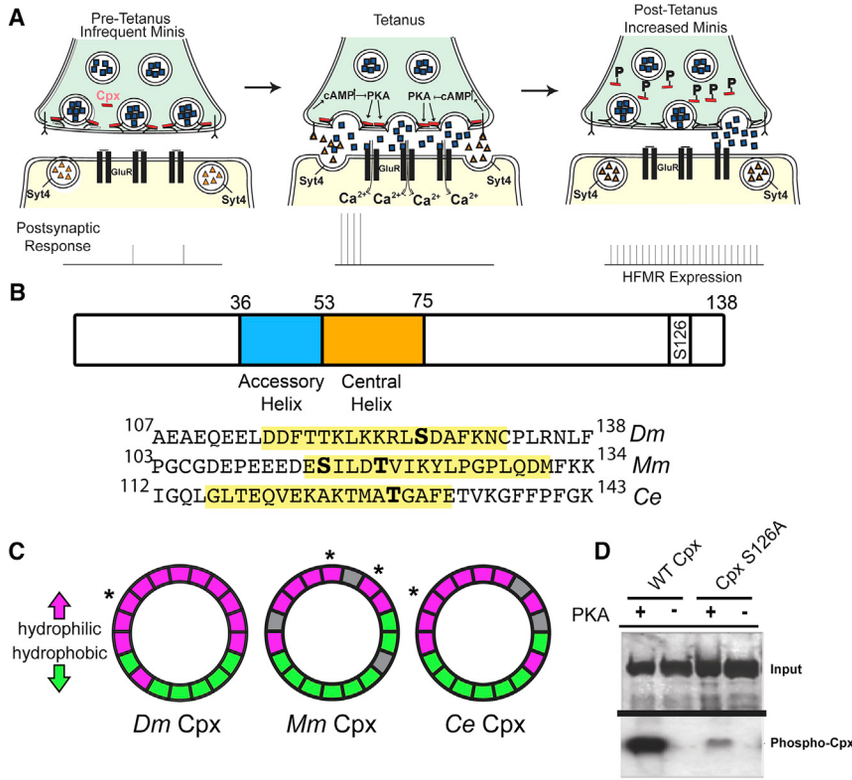
(F) Representative images of  $Ca^{2+}$  dynamics detected by presynaptic myrGCaMP6s expression before, during, and after stimulation.

(G) Mean fluorescence change as a function of time during tetanus is shown for one representative NMJ stimulation trial. Red arrows indicate each 100 Hz stimulation event. Mean  $\pm$  SEM is indicated in all figures. Indicated comparisons were made using ANOVA with post hoc Tukey analysis.

on Cpx's ability to directly regulate mini release frequency as a fusion clamp, we examined whether Cpx is a downstream target of the Syt 4 retrograde signaling pathway and if its synaptic function as a fusion clamp is regulated by PKA phosphorylation in response to stimulation that leads to HFMR expression at NMJs (Figure 4A).

Acute and local regulation of Cpx function as a vesicle clamp at the post-translational level is likely to involve the C terminus of Cpx (Figure 4B). We previously identified two transcripts that arise from the *cpx* locus that result in the expression of Cpx isoforms with distinct C-terminal domains—one containing a CAAX prenylation motif (termed 7A) and one lacking this lipid tethering domain (7B) (Buhl et al., 2013; Cho et al., 2010). The C terminus of Cpx is required to inhibit spontaneous vesicle fusion, likely via its ability to interact with lipid membranes (Cho et al., 2010; Iyer et al., 2013; Kaeser-Woo et al., 2012; Snead et al., 2014; Wragg et al., 2013; Xue et al., 2009).

Although Cpx 7B does not contain a CAAX motif, it contains a predicted amphipathic helix that likely mediates protein/lipid association important for Cpx function (Wragg et al., 2013). Interestingly, S126 in the C terminus of Cpx 7B is predicted to be a PKA phosphorylation site (<http://mendel.imp.ac.at/pkaPS/>) based on the consensus sequence R-R-X-S/T-Φ (Figure 4B) (Ubersax and Ferrell, 2007). This phosphorylation site flanks the hydrophobic face of the predicted amphipathic region near the end of the Cpx C terminus (Figures 4B and 4C). The amphipathic region is conserved in other Cpx orthologs, and mammalian and *C. elegans* phosphorylation sites that similarly flank the respective hydrophobic face of the amphipathic region have either been identified or predicted (see Figures 4B and 4C) (Shata et al., 2007). Phosphorylation of Cpx 7B (hereafter referred to as Cpx) at this site could disrupt C-terminal membrane interactions occurring along the hydrophobic face of the amphipathic helix. To assay if Cpx is indeed



**Figure 4. Cpx Is a Substrate for PKA Phosphorylation**

(A) Proposed retrograde Syt 4-dependent signaling pathway that is activated by strong stimulation. Postsynaptic Ca<sup>2+</sup> elevation following stimulation drives release of a retrograde signal that activates the presynaptic PKA pathway. PKA is hypothesized to phosphorylate Cpx, altering its clamping properties. In addition, PKA may phosphorylate additional targets to facilitate structural growth.

(B) Structure of *Drosophila* (*Dm*) Cpx and the location of the predicted PKA phosphorylation site (S126) in the C terminus. The N terminus and C terminus flank the indicated accessory and central helix domains. The alignment of C-terminal regions of *Dm* Cpx, mouse (*Mm*) Cpx I, and *C. elegans* (*Ce*) Cpx is shown below. Predicted amphipathic helix domains are highlighted in yellow, and predicted phosphorylation sites are indicated in bold. S115 of *Mm* Cpx I is phosphorylated by protein kinase CK2 (Shata et al., 2007). T119 of *Mm* Cpx I is a predicted substrate for PKC, and T129 of *Ce* Cpx is a predicted substrate for AKT (Wong et al., 2007).

(C) Helical wheel diagrams of the amphipathic region for *Dm* Cpx, *Mm* Cpx I, and *Ce* Cpx. Hydrophilic residues are indicated in magenta, hydrophobic residues in green, and nonpolar and uncharged residues in gray. Predicted phosphorylation sites are indicated by \*.

(D) Cpx<sup>S126A</sup> exhibited greatly decreased [<sup>32</sup>P] incorporation compared to WT Cpx (Cpx<sup>S126A</sup> [<sup>32</sup>P] incorporation was reduced to 35.2% ± 10.6%

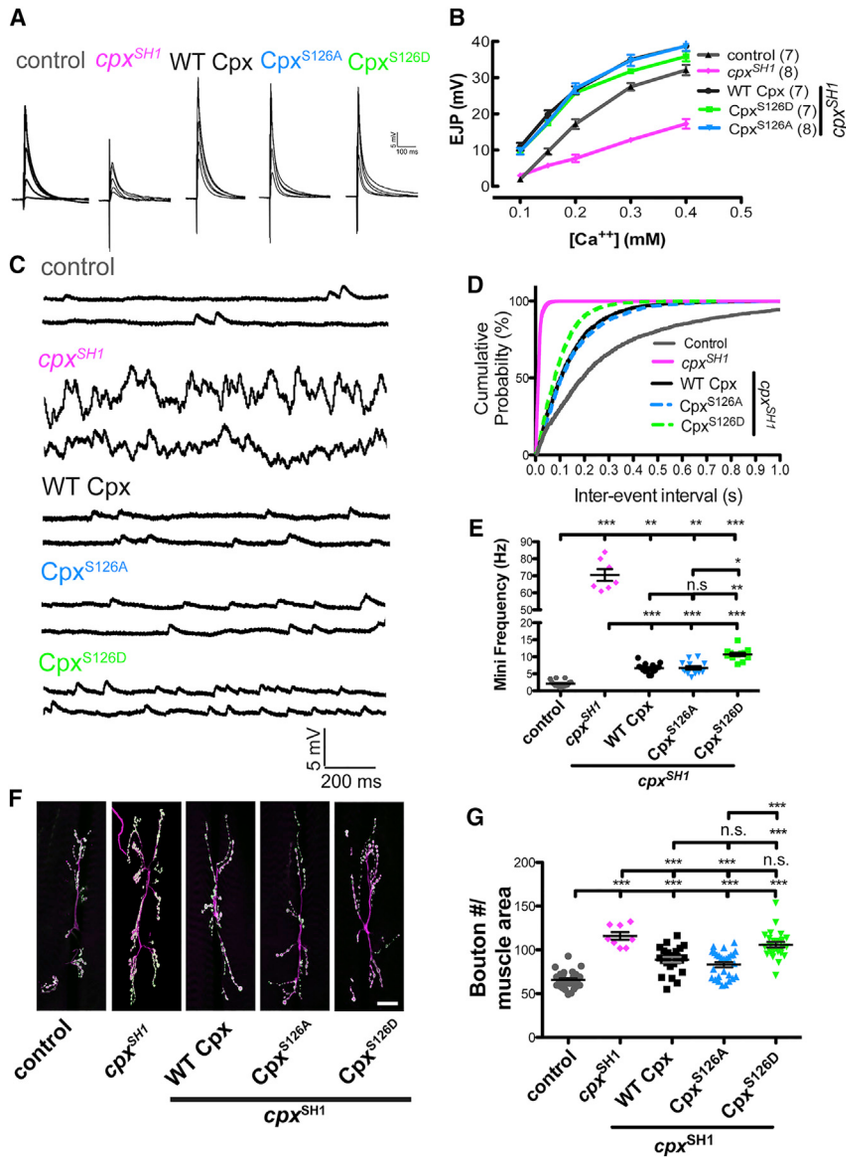
[mean ± SEM] relative band intensity of WT Cpx [<sup>32</sup>P] incorporation [n = 4]). Recombinant control Cpx (WT) and phosphoincompetent Cpx<sup>S126A</sup> were incubated with [<sup>32</sup>P]ATP and recombinant active PKA. Phosphorylation was visualized by autoradiography after proteins were separated by SDS-PAGE (bottom panel). Gels were also stained with Coomassie blue to assay protein loading (top panel).

a substrate for PKA phosphorylation, recombinant Cpx was purified and incubated with PKA and [<sup>32</sup>P]ATP. In the presence of PKA, Cpx readily incorporated <sup>32</sup>P, indicating Cpx is a substrate for PKA in vitro (Figure 4D). Mutation of the predicted PKA phosphorylation site S126 to a phosphoincompetent alanine residue (Cpx<sup>S126A</sup>) resulted in greatly reduced PKA-dependent phosphorylation compared to wild-type (WT) Cpx (<sup>32</sup>P incorporation into Cpx<sup>S126A</sup> is reduced to 35.2% ± 10.6% of WT Cpx [<sup>32</sup>P] incorporation [n = 4]), suggesting PKA phosphorylates Cpx primarily at the predicted PKA phosphorylation residue S126.

To examine how Cpx phosphorylation at S126 might alter its in vivo function, we generated transgenic animals that neuronally express WT Cpx or Cpx with S126 substitutions in the *cpx<sup>SH1</sup>* null background. We generated serine → alanine (Cpx<sup>S126A</sup>) and serine → aspartic acid (Cpx<sup>S126D</sup>) substitutions to mimic phosphoincompetent and phosphomimetic Cpx states, respectively. We first assayed the ability of the Cpx<sup>S126</sup> mutants to rescue basal electrophysiological properties compared to WT Cpx rescues by recording from 3<sup>rd</sup> instar NMJs and measuring minis and evoked responses. *cpx<sup>SH1</sup>* null larvae exhibited reduced evoked responses across all Ca<sup>2+</sup> concentrations compared to neuronally expressed WT Cpx rescues (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>* = 7.7 ± 1.0 mV [n = 8] versus C155; WT Cpx, *cpx<sup>SH1</sup>* = 26.6 ± 1.1 mV [n = 7] at [Ca<sup>2+</sup>] = 0.2 mM; p < 0.001; Figures 5A and 5B). Both neuronally expressed Cpx<sup>S126A</sup> and Cpx<sup>S126D</sup> point mutants

rescued the *cpx<sup>SH1</sup>* evoked release phenotype, with responses comparable to WT Cpx ([C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* = 27.1 ± 1.3 mV (n = 8) and C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>* = 25.8 ± 0.8 (n = 7)] versus C155; WT Cpx, *cpx<sup>SH1</sup>* = 26.6 ± 1.1 mV (n = 7) at [Ca<sup>2+</sup>] = 0.2 mM; p > 0.05). These results indicate C-terminal modification at S126 does not alter evoked release properties regulated by Cpx.

We next examined the ability of phosphoincompetent and phosphomimetic Cpx to rescue the *cpx<sup>SH1</sup>* null mini phenotype. *cpx<sup>SH1</sup>* null larvae exhibited an elevated rate of minis that are significantly reduced by neuronal expression of WT Cpx (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>* = 70.4 ± 3.4 Hz [n = 7] versus C155; WT Cpx, *cpx<sup>SH1</sup>* = 6.6 ± 0.4 Hz [n = 15]; p < 0.001; Figures 5C–5E). Phosphoincompetent Cpx<sup>S126A</sup> rescue lines exhibited mini frequencies similar to WT Cpx rescues (C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* = 6.7 ± 0.5 Hz [n = 14] versus C155; WT Cpx, *cpx<sup>SH1</sup>* = 6.6 ± 0.4 Hz [n = 15]; p > 0.05), indicating the basal clamping function of Cpx<sup>S126A</sup> is intact. In contrast, Cpx<sup>S126D</sup> rescue animals exhibited a significant elevation in mini frequency compared to WT Cpx and Cpx<sup>S126A</sup> rescues (C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>* = 10.7 ± 0.5 Hz (n = 13) versus [C155; WT Cpx, *cpx<sup>SH1</sup>* = 6.6 ± 0.4 Hz (n = 15); p < 0.001 and C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* = 6.7 ± 0.5 Hz (n = 14); p < 0.01]). These results suggest that PKA phosphorylation of Cpx<sup>S126</sup> may selectively regulate Cpx's ability to act as a vesicle fusion clamp, leaving its function in evoked release unperturbed.



We next assayed if WT Cpx and Cpx<sup>S126</sup> mutant rescues altered synaptic growth. *cpx<sup>SH1</sup>* nulls exhibit enhanced synaptic growth that correlate with enhanced synaptic minis (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>* = 115.8 ± 4.3 (n = 8) boutons versus control (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>* = 65.6 ± 1.8 boutons [n = 28]; p < 0.001; Figures 5F and 5G). WT Cpx and both Cpx<sup>S126A</sup> rescue lines show a mild elevation of mini frequency due to the lack of a complete rescue of the null phenotype (Figures 5C–5E). Consistent with a positive correlation of minis and synaptic growth, WT Cpx and Cpx<sup>S126A</sup> mutant rescues showed modest enhancements in synaptic growth compared to control (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>* = 65.6 ± 1.8 boutons [n = 28] versus C155; WT Cpx, *cpx<sup>SH1</sup>* = 88.7 ± 3.0 boutons [n = 24] and C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* = 83.1 ± 2.9 boutons [n = 27]; p < 0.001 for both comparisons; Figures 5F and 5G). Strikingly, the elevated mini frequency in the Cpx<sup>S126D</sup> rescue animals compared to WT Cpx and Cpx<sup>S126A</sup> rescues (Figures 5C–5E) was correlated with significantly enhanced synaptic growth

growth, suggesting acute regulation of Cpx function via PKA phosphorylation may regulate synaptic growth and plasticity.

### HFMR Expression Requires PKA and Cpx Phosphorylation Site S126

Having established that the PKA phosphorylation site in Cpx (S126) can regulate the clamping properties of the protein, we examined if Cpx<sup>S126</sup> is required for HFMR expression. We first determined if WT Cpx rescues were capable of expressing HFMR. HFMR induction in larval NMJs neuronally expressing WT Cpx in a *cpx<sup>SH1</sup>* null background resulted in HFMR that was blocked by the PKA inhibitor, H89 (C155; WT Cpx, *cpx<sup>SH1</sup>* = 1.6 ± 0.1 fold [n = 7] versus C155; WT Cpx, *cpx<sup>SH1</sup>* + H89 = 1.0 ± 0.1 fold [n = 6] at 100 s after induction; p < 0.001; Figures 6A–6C). To examine the requirement of S126 for HFMR expression, we neuronally expressed Cpx<sup>S126</sup> mutants in the *cpx<sup>SH1</sup>* null background. Compared to WT Cpx rescues, Cpx<sup>S126A</sup> did

### Figure 5. Mimicking Phosphorylation of Cpx at S126 Selectively Modulates Cpx Function as a Fusion Clamp and Regulates Synaptic Growth

(A) Averaged traces of EJPs from control (*w<sup>1118</sup>*; *Cpx<sup>PE</sup>*), *cpx* null (*w<sup>1118</sup>*; *cpx<sup>SH1</sup>*), WT Cpx (C155; WT Cpx, *cpx<sup>SH1</sup>*), and Cpx<sup>S126A</sup> and Cpx<sup>S126D</sup> mutant rescues (C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* and C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>*, respectively) recorded over a range of external Ca<sup>2+</sup> concentrations (0.1, 0.15, 0.2, 0.3, and 0.4 mM). Numbers in parentheses (n) represent individual muscle recordings at the indicated [Ca<sup>2+</sup>].

(B) Summary of mean EJP amplitude (mV ± SEM) plotted at the indicated [Ca<sup>2+</sup>].

(C) Representative traces of spontaneous release events from muscle 6 of control, *cpx<sup>SH1</sup>* null, and WT Cpx, Cpx<sup>S126A</sup> and Cpx<sup>S126D</sup> rescue lines.

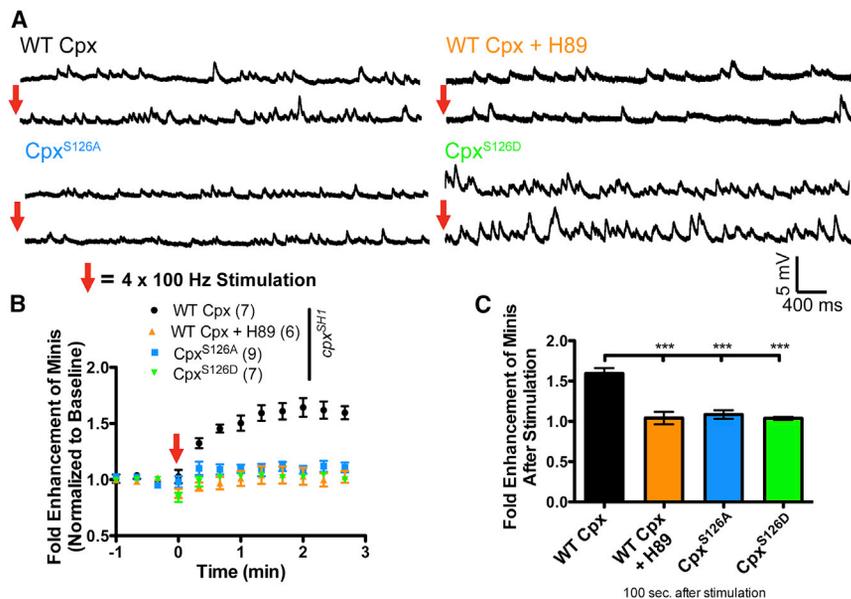
(D) Cumulative probability plot of inter-event intervals (seconds) between mini events recorded indicated genotypes.

(E) Summary of mean mini frequency (Hz ± SEM) for indicated genotypes.

(F) Representative images of muscle 6/7 NMJs of control, *cpx<sup>SH1</sup>* null, and WT Cpx, Cpx<sup>S126A</sup>, Cpx<sup>S126D</sup> rescue lines. Larvae were stained with anti-HRP (magenta) and anti GluRIII (green) specific antisera to identify synaptic boutons. Scale bar, 20 μm.

(G) Summary of mean synaptic bouton number normalized to muscle area for the indicated genotypes. Indicated comparisons were made using ANOVA with post hoc Tukey analysis. Mean ± SEM is indicated in all figures.

(C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>* = 105.8 ± 3.2 boutons (n = 25) versus [C155; WT Cpx, *cpx<sup>SH1</sup>* = 88.8 ± 3.0 boutons (n = 24) and C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* = 83.1 ± 2.9 boutons (n = 27)]; p < 0.001 for both comparisons). These data demonstrate that phosphomimetic Cpx<sup>S126D</sup> displays elevated mini release and increased synaptic



**Figure 6. Cpx PKA Phosphorylation Site S126 Is Required for HFMR Expression**

(A) Representative traces of minis before (top) and after (bottom) tetanic stimulation recorded from *cpx<sup>SH1</sup>* rescue lines neuronally expressing WT Cpx rescues (C155; WT Cpx, *cpx<sup>SH1</sup>*) with or without PKA inhibitor H89, and Cpx<sup>S126A</sup> and Cpx<sup>S126D</sup> mutant rescues (C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* and C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>*, respectively).

(B) Mean mini frequency plotted as a function of time. Tetanic stimulation occurs at the 0 s time point (Red Arrow). Mini frequency is normalized to average baseline mini frequency prior to stimulation for each preparation.

(C) Average fold enhancement of mini frequency compared to baseline 100 s after tetanic stimulation. Indicated comparisons were made using ANOVA with post hoc Tukey analysis. Mean  $\pm$  SEM is indicated in all figures.

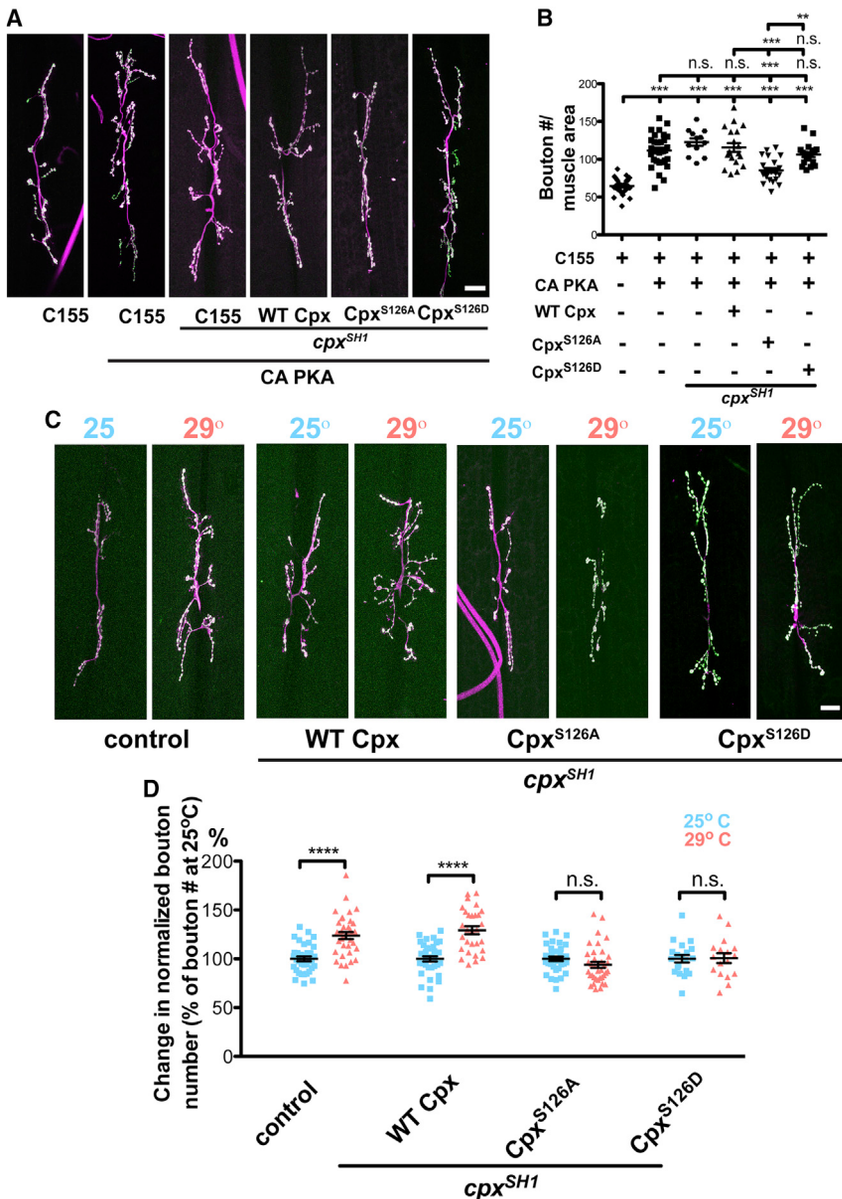
not exhibit significant HFMR (C155; WT Cpx, *cpx<sup>SH1</sup>* =  $1.6 \pm 0.1$  fold [ $n = 7$ ] versus C155; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>* =  $1.1 \pm 0.1$  fold [ $n = 9$ ] 100 s after induction;  $p < 0.001$ ), demonstrating that S126 is necessary for HFMR expression. The phosphomimetic Cpx<sup>S126D</sup> exhibited elevated mini frequency under basal conditions (see Figures 5C–5E). If activity-dependent phosphorylation of Cpx<sup>S126</sup> is necessary for HFMR expression, we hypothesized that the phosphomimetic Cpx<sup>S126D</sup> would occlude HFMR. Examination of HFMR in Cpx<sup>S126D</sup> rescues revealed that Cpx<sup>S126D</sup> animals failed to express HFMR compared to WT Cpx rescues (C155; WT Cpx, *cpx<sup>SH1</sup>* =  $1.6 \pm 0.1$  fold [ $n = 7$ ] versus C155; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>* =  $1.0 \pm 0.1$  fold [ $n = 7$ ] 100 s after induction;  $p < 0.001$ ), demonstrating that mimicking phosphorylation of Cpx at S126 is sufficient to elevate spontaneous release rate without stimulation and to occlude HFMR expression. Together, these results suggest that dynamic phosphorylation of Cpx at residue S126 is necessary for HFMR expression.

### PKA Phosphorylation of Cpx S126 Is Required for Activity-Dependent Structural Plasticity

Our results suggest PKA phosphorylation of Cpx regulates its function as a vesicle fusion clamp to express HFMR. The cAMP-PKA pathway also regulates synaptic morphology at the *Drosophila* NMJ. cAMP metabolism mutants, *dunce* (*dnc*) and *rutabaga* (*rut*), exhibit synaptic growth alterations in addition to the behavioral learning defects by which they were originally identified (Davis et al., 1996; Guan et al., 2011; Zhong et al., 1992; Zhong and Wu, 1991). We previously observed that expression of a constitutively active PKA (CA-PKA) results in synaptic NMJ overgrowth (Yoshihara et al., 2005). 3<sup>rd</sup> instar larvae that express presynaptic CA-PKA exhibit enhanced synaptic growth compared to control (C155; CA-PKA/+ =  $111.6 \pm 3.6$  boutons [ $n = 33$ ] versus control (C155) =  $64.3 \pm 1.8$  boutons [ $n = 30$ ];  $p < 0.001$  Figures 7A and 7B). To examine the contribution of Cpx<sup>S126</sup> to PKA-dependent synaptic growth, we presynaptically co-expressed CA-PKA in the *cpx<sup>SH1</sup>* null

background with either WT Cpx or the phosphoincompetent Cpx<sup>S126A</sup>. Co-expression of CA-PKA with WT Cpx results in synaptic overgrowth similar to CA-PKA expression alone (C155; CA-PKA/+; WT Cpx, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $115.6 \pm 5.8$  boutons [ $n = 19$ ] compared to control (C155) =  $64.3 \pm 1.8$  boutons [ $n = 30$ ];  $p < 0.001$ ). However, co-expression of CA-PKA with PKA phosphoincompetent Cpx<sup>S126A</sup> suppressed synaptic overgrowth compared to WT Cpx rescues (C155; CA-PKA/+; Cpx<sup>S126A</sup>, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $85.4 \pm 3.1$  boutons [ $n = 24$ ] versus C155; CA-PKA/+; WT Cpx, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $115.6 \pm 5.8$  boutons [ $n = 19$ ];  $p < 0.001$ ). We next examined synaptic growth when co-expressing CA-PKA and phosphomimetic Cpx<sup>S126D</sup>. We found that co-expressing CA-PKA with phosphomimetic Cpx<sup>S126D</sup> exhibited synaptic growth similar to WT Cpx rescues (C155; CA-PKA/+; Cpx<sup>S126D</sup>, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $106.3 \pm 3.5$  boutons [ $n = 18$ ] versus C155; CA-PKA/+; WT Cpx, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $115.6 \pm 5.8$  boutons [ $n = 19$ ];  $p > 0.05$ ). Interestingly, CA-PKA expressed in the *cpx<sup>SH1</sup>* null background did not exhibit significant synaptic growth beyond CA-PKA alone or WT Cpx rescues (C155; CA-PKA/+; *cpx<sup>SH1</sup>* =  $122.8 \pm 5.0$  boutons [ $n = 12$ ] versus [C155; CA-PKA/+ =  $111.6 \pm 3.6$  boutons ( $n = 33$ ) and C155; CA-PKA/+; WT Cpx, *cpx<sup>SH1</sup>*/*cpx<sup>SH1</sup>* =  $115.6 \pm 5.8$  boutons ( $n = 19$ )]);  $p > 0.5$  for both comparisons). This result suggests a shared signaling pathway between PKA and Cpx in regulating synaptic growth rather than parallel pathways. Overall, these results demonstrate that S126 is required for PKA-dependent synaptic growth and suggest Cpx is a downstream target of the PKA signaling pathway that regulates morphological growth and structural plasticity of synaptic terminals.

Synaptic expansion at the NMJ is regulated by the overall level of motor neuron activity. Genetic manipulations that enhance or decrease neuronal activity at the NMJ alter synapse formation at *Drosophila* NMJs (Budnik et al., 1990; Schuster et al., 1996). In addition to genetic manipulations that modulate activity, larvae reared under variable environmental temperatures modulate



**Figure 7. Cpx PKA Phosphorylation Site S126 Is Required for PKA and Activity-Dependent Synaptic Growth**

(A) Representative images of muscle 6/7 NMJs from control (C155 *elav-GAL4*) and lines neuronally overexpressing constitutively active PKA alone (C155; CA-PKA/+ ) or in the *cpx<sup>SH1</sup>* null (C155; CA-PKA/+; *cpx<sup>SH1</sup>*), WT Cpx rescue (C155; CA-PKA/+; WT Cpx, *cpx<sup>SH1</sup>/cpx<sup>SH1</sup>*), *Cpx<sup>S126A</sup>* rescue (C155; CA-PKA/+; *Cpx<sup>S126A</sup>, cpx<sup>SH1</sup>/cpx<sup>SH1</sup>*), and *Cpx<sup>S126D</sup>* rescue (C155; CA-PKA/+; *Cpx<sup>S126D</sup>, cpx<sup>SH1</sup>/cpx<sup>SH1</sup>*) backgrounds. (B) Summary of mean synaptic bouton number normalized to muscle area for the indicated genetic backgrounds. Indicated comparisons were made using ANOVA with post hoc Tukey analysis.

(C) Representative images of control (*w<sup>1118.</sup>*; *Cpx<sup>PE</sup>*) and WT Cpx rescues (C155; WT Cpx, *cpx<sup>SH1</sup>*), *Cpx<sup>S126A</sup>* rescues (C155; *Cpx<sup>S126A</sup>, cpx<sup>SH1</sup>/cpx<sup>SH1</sup>*), and *Cpx<sup>S126D</sup>* rescues (C155; *Cpx<sup>S126D</sup>, cpx<sup>SH1</sup>/cpx<sup>SH1</sup>*) reared at 25° or 29°. (D) Summary of percent change in synaptic bouton number normalized to muscle area for the indicated genetic backgrounds and rearing temperatures. Synaptic bouton number was normalized to bouton number of larvae reared at 25° for each respective genotype to reflect percent change. Indicated comparisons were made using Student's t test. Scale bars, 20 μm. Mean ± SEM is indicated in all figures.

*w<sup>1118.</sup>*; *Cpx<sup>PE</sup>* at 25°C = 100.0 ± 2.5 boutons [n = 35]; p < 0.0001; Figures 7C and 7D). Rescue larvae expressing presynaptic WT Cpx in the *cpx<sup>SH1</sup>* null background exhibited elevated synaptic growth at 29°C compared to growth at 25°C, similar to controls (C155; WT Cpx, *cpx<sup>SH1</sup>* at 29°C = 129.2 ± 4.1 boutons [n = 29] versus C155; WT Cpx, *cpx<sup>SH1</sup>* at 25°C = 100.0 ± 2.7 boutons [n = 36]; p < 0.0001). We next examined if phosphorylation of S126 might regulate this form of structural plasticity. Unlike control and WT Cpx rescues, phosphoincompetent *Cpx<sup>S126A</sup>* rescues

failed to exhibit enhanced synaptic growth at elevated temperatures (C155; *Cpx<sup>S126A</sup>, cpx<sup>SH1</sup>* at 29°C = 93.9 ± 3.2 [n = 36] boutons versus C155; *Cpx<sup>S126A</sup>, cpx<sup>SH1</sup>* at 25°C = 100.0 ± 2.3 boutons [n = 35]; p > 0.05). Similar to earlier experiments in which HFMR expression was occluded in *Cpx<sup>S126D</sup>* rescues, phosphomimetic *Cpx<sup>S126D</sup>* rescues did not express temperature-induced synaptic growth (C155; *Cpx<sup>S126D</sup>, cpx<sup>SH1</sup>* at 29°C = 100.7 ± 5.0 [n = 17] versus C155; *Cpx<sup>S126D</sup>, cpx<sup>SH1</sup>* at 25°C = 100.0 ± 3.8 boutons [n = 20]; p > 0.05). These results indicate that the PKA phosphorylation site at S126 in Cpx is required for expression of temperature-induced structural plasticity at *Drosophila* NMJs. Taken together, these data suggest that a PKA-dependent retrograde signaling pathway mediated by postsynaptic Syt 4 converges on Cpx to regulate its function as a synaptic vesicle fusion clamp and that this regulation is required for

locomotor activity (Sigrist et al., 2003; Zhong and Wu, 2004). At elevated temperatures, enhanced larval locomotion is correlated with enhanced synaptic NMJ growth, suggesting locomotor experience and NMJ growth are directly coupled (Sigrist et al., 2003). Interestingly, enhancement of synaptic growth at elevated temperatures is suppressed in adenylyl cyclase mutants, indicating that increased neuronal activity and synaptic growth induced environmentally requires PKA signaling (Zhong and Wu, 2004). To determine if Cpx phosphorylation contributes to this form of activity-dependent structural plasticity, we examined temperature-induced synaptic growth in WT Cpx and *Cpx<sup>S126A</sup>* mutant rescue animals. Similar to previously published results, rearing control larvae at 29°C resulted in a significant increase in synaptic growth compared to animals reared at 25°C (*w<sup>1118.</sup>*; *Cpx<sup>PE</sup>* at 29°C = 123.8 ± 3.6 boutons [n = 36] versus

activity-dependent functional and structural plasticity that occurs downstream of enhanced spontaneous release.

## DISCUSSION

These findings indicate that the SNARE-binding protein Cpx is a key PKA target that regulates spontaneous fusion rates and pre-synaptic plasticity at *Drosophila* NMJs. We demonstrate that Cpx's function can be modified to regulate activity-dependent functional and structural plasticity. In vivo experiments using Cpx phosphomimetic rescues demonstrate that Cpx phosphorylation at residue S126 selectively alters its ability to act as a synaptic vesicle fusion clamp. In addition, S126 is critical for the expression of HFMR, an activity-dependent form of acute functional plasticity that modulates mini frequency at *Drosophila* synapses. Our data indicate a Syt 4-dependent retrograde signaling pathway converges on Cpx to regulate its synaptic function (Figure 4A). Additionally, we find that elevated spontaneous fusion rates correlate with enhanced synaptic growth. This pathway requires Syt 4 retrograde signaling to enhance spontaneous release and to trigger synaptic growth. Moreover, the Cpx S126 PKA phosphorylation site is required for activity-dependent synaptic growth, suggesting acute regulation of minis via Cpx phosphorylation is likely to contribute to structural synaptic plasticity. Together, these data identify a mechanism of acute synaptic plasticity that impinges directly on the presynaptic release machinery to regulate spontaneous release rates and synaptic maturation.

How does acute phosphorylation of S126 alter Cpx's function? The Cpx C terminus associates with lipid membranes through a prenylation domain (CAAX motif) and/or the presence of an amphipathic helix (Cho et al., 2010; Iyer et al., 2013; Kaeser-Woo et al., 2012; Snead et al., 2014; Wragg et al., 2013; Xue et al., 2009). The *Drosophila* Cpx isoform used in our study (Cpx 7B) lacks a CAAX-motif but contains a C-terminal amphipathic helix flanked by the S126 phosphorylation site. S126 phosphorylation does not alter synaptic targeting of Cpx (Figures S1A and S1B) or its ability to associate with SNARE complexes in vitro (Figures S2, S3A, and S3B; Table S3). As such, phosphorylation may instead alter interactions of the amphipathic helix region with lipid membranes and/or alter Cpx interactions with other proteins that modulate synaptic release. Given the well-established role of the Cpx C terminus in regulating membrane binding and synaptic vesicle tethering of Cpx, phosphorylation at this site would be predicted to alter the subcellular localization of the protein and its accessibility to the SNARE complex (Buhl et al., 2013; Snead et al., 2014; Wragg et al., 2013). However, we did not detect large differences between WT Cpx and Cpx<sup>S126D</sup> in liposome binding (Figure S4). This assay is unlikely to reveal subtle changes in lipid interactions by Cpx, as we found that C-terminal deletions (CTDs) maintained its ability to bind membranes. The ability of the CTD versions of *Drosophila* Cpx to associate with liposomes is unlike that observed with *C. elegans* Cpx (Wragg et al., 2013) and indicate domains outside of *Drosophila* Cpx's C terminus contribute to lipid membrane association as well, potentially masking effects from S126 phosphorylation that might occur in vivo. Alternatively, phosphorylation of the Cpx C terminus could alter its as-

sociation with other SNARE complex modulators such as Syt 1 (Jorquera et al., 2012; Tang et al., 2006).

Our data indicate that enhanced minis regulate synaptic growth through several previously identified NMJ maturation pathways (Figure 4A). We find that the Wit signaling pathway is required for synaptic growth in the background of enhanced minis. The *wit* gene encodes a presynaptic type II BMP receptor that receives retrograde, transsynaptic BMP signals from postsynaptic muscles (Marqués et al., 2002; McCabe et al., 2003; Piccioli and Littleton, 2014). Consistent with our data, other studies have demonstrated that downstream signaling components of the BMP pathway are necessary and sufficient for mini-dependent synaptic growth at the *Drosophila* NMJ (Choi et al., 2014). Additionally, we found that the postsynaptic Ca<sup>2+</sup> sensor, Syt 4, is required for enhancing spontaneous release and increasing synaptic growth (Barber et al., 2009; Yoshihara et al., 2005). Our data do not currently distinguish the interdependence of the BMP and Syt 4 retrograde signaling pathways, and other retrograde signaling pathways might contribute to mini-dependent synaptic growth as well (Ballard et al., 2014). Recently, several retrograde pathways have been identified that regulate functional homeostatic plasticity at the *Drosophila* NMJ (Davis and Müller, 2015). Future work will be required to fully define the retrograde signaling pathways necessary to mediate mini-dependent synaptic growth.

How might elevated spontaneous release through Cpx phosphorylation regulate synaptic growth? We hypothesize that the switch in synaptic vesicle release mode to a constitutive fusion pathway that occurs over several minutes following stimulation serves as a synaptic tagging mechanism. By continuing to activate postsynaptic glutamate receptors in the absence of incoming APs, the elevation in mini frequency would serve to enhance postsynaptic calcium levels by prolonging glutamate receptor stimulation (as observed in Figure 3D). This would prolong retrograde signaling that initiates downstream cascades to directly alter cytoskeletal architecture required for synaptic bouton budding (Piccioli and Littleton, 2014). Given that elevated rates of spontaneous fusion still occur in *cpx* and *syx*<sup>3-69</sup> in the absence of Syt 4 and BMP signaling, yet synaptic overgrowth is suppressed in these conditions, it is unlikely that spontaneous fusion itself directly drives synaptic growth. Rather, the transient enhancement in spontaneous release may serve to prolong postsynaptic calcium signals that engage distinct effectors for structural remodeling that fail to be activated in the absence of elevated spontaneous release. Results from mammalian studies indicate spontaneous release can uniquely regulate postsynaptic protein translation and activate distinct populations of NMDA receptors (Atasoy et al., 2008; Sutton et al., 2006) compared to evoked release, so it is possible that spontaneous fusion may engage unique postsynaptic effectors at *Drosophila* NMJs as well.

In the last few decades, intense research efforts have elucidated several molecular mechanisms of classic Hebbian forms of synaptic plasticity that include long-term potentiation (LTP) and long-term depression (LTD), alterations in synaptic function that lasts last minutes to hours (Mayford et al., 2012). The most widely studied expression mechanism for these forms of synaptic plasticity involve modulation of postsynaptic AMPA-type glutamate receptor (AMPA) function and membrane trafficking

(Huganir and Nicoll, 2013). In contrast, molecular mechanisms of short-term synaptic plasticity remain poorly understood. Several forms of short-term plasticity have been described, such as post-tetanic potentiation (PTP), which involves stimulation-dependent increases in synaptic strength, including changes in mini frequency (Habets and Borst, 2005; Korogod et al., 2005; Lou et al., 2008). Short-term plasticity expression is likely to impinge on alterations to the presynaptic release machinery downstream of activated effector molecules (Zucker and Regehr, 2002). For example, Munc 18, a presynaptic protein involved in the priming step of vesicle exocytosis via its ability to associate with members of the SNARE machinery, is dynamically regulated by  $Ca^{2+}$ -dependent protein kinase C (PKC), and its regulation is required to express PTP at the Calyx of Held (Genc et al., 2014). Here we demonstrate that the presynaptic vesicle fusion machinery can also be directly modified to alter spontaneous neurotransmission via activity-dependent modification of Cpx function by PKA. Protein kinase CK2 and PKC phosphorylation sites within the C terminus of mammalian and *C. elegans* Cpx have been identified (see Figures 4B and 4C) (Shata et al., 2007). Therefore, activity-dependent regulation of Cpx function via C-terminal phosphorylation may be an evolutionarily conserved mechanism to regulate synaptic plasticity. Moreover, Cpx may lie downstream of multiple effector pathways to modulate various forms of short-term plasticity, including PTP, in a synapse-specific manner. Interestingly, Cpx is expressed both presynaptically and postsynaptically in mammalian hippocampal neurons and is required to express LTP via regulation of AMPAR delivery to the synapse, suggesting Cpx-mediated synaptic plasticity expression mechanisms may also occur postsynaptically (Ahmad et al., 2012).

In summary, these results indicate minis serve as an independent and regulated neuronal signaling pathway that contributes to activity-dependent synaptic growth. We previously found Cpx's function as a facilitator and clamp for synaptic vesicle fusion is genetically separable, demonstrating distinct molecular mechanisms regulate evoked and spontaneous release (Cho et al., 2010, 2014). Evoked and spontaneous release are also separable beyond Cpx regulation, as other studies have demonstrated that minis can utilize distinct components of the SNARE machinery, distinct vesicle pools, and distinct individual synaptic release sites (Kavalali, 2015; Melom et al., 2013). These findings suggest diverse regulatory mechanisms for spontaneous release that might be selectively modulated at specific synapses.

## EXPERIMENTAL PROCEDURES

### *Drosophila* Genetics and Molecular Cloning

*Drosophila melanogaster* were cultured on standard medium at 25°C. QuickChange (Stratagene) was used for site directed mutagenesis on the existing cloned Cpx (Isoform 7B) (Buhl et al., 2013; Huntwork and Littleton, 2007) to generate specified Cpx mutants. PCR products were subcloned into a modified pValum construct, allowing use of the Gal4/UAS expression system (Brand and Perrimon, 1993). These constructs were injected into a *yv; attP 3rd* chromosome docking strains by Genetic Services Inc. Homozygous third chromosome UAS lines and as well as mutant alleles, *syt 4<sup>BA1</sup>* and *wit*, were then recombined into the *cpx<sup>SH1</sup>* null mutant background. The C155 *elav*-GAL4 driver was used for neuronal expression of transgenes unless otherwise indicated. Cpx RNAi knockdown line (transformant 21477) was obtained from the Vienna *Drosophila* Resource Center (Dietzl et al., 2007).

### Immunostaining

Immunostaining was performed on wandering 3<sup>rd</sup> instar larvae reared at 25°C, unless otherwise indicated, as previously described (Huntwork and Littleton, 2007). See Supplemental Information for more information.

### In Vitro Kinase Assay

In vitro kinase assays were performed using the purified recombinant Cpx protein and the catalytic subunit of PKA (New England Biolabs), according to manufacturer supplied instructions. Briefly, approximately 10 mg of purified GST-fusion protein (WT Cpx and Cpx<sup>S126A</sup>) was used per reaction and incubated with 2,500 units of recombinant kinase and [<sup>32</sup>P]ATP (Perkin Elmer). Reaction products were separated by SDS-PAGE. Gels were stained with Bio-Safe Coomassie Blue (Bio-Rad), dried, and exposed to autoradiography film at room temperature. Mean integrated density of each band (mean intensity/pixel multiplied by the area of the band) was quantified using ImageJ, and relative density of mutant Cpx was calculated by normalizing to WT Cpx band intensity.

### Electrophysiological Analysis

Electrophysiological analysis of wandering 3<sup>rd</sup> instar larva was performed in *Drosophila* HL3.1 saline containing (in mM) 70 NaCl, 5 KCl, 4 MgCl<sub>2</sub>, 10 NaHCO<sub>3</sub>, 5 Trehalose, 115 sucrose, and 5 HEPES-NaOH [pH 7.2] and indicated concentrations of  $Ca^{2+}$ . Evoked EJPs were recorded intracellularly from muscle fiber 6 of segments A3 and A4 using an Axoclamp 2B amplifier (Axon Instruments). Data acquisition and analysis was performed using Clampfit 9.0 software (Axon Instruments) and Mini Analysis 6.0.3 (Synaptosoft, Inc.) as previously described (Cho et al., 2010). Nerve stimulation at indicated frequencies in each experiment was performed using a programmable stimulator (Master-8; AMPI). PKA inhibitor H89 ([25 μM] in HL3.1) was bath applied to the dissected larvae and incubated for 10 min prior to stimulation (Tocris Bioscience). Mini frequency (number of individual mini events/second) was calculated from recordings of at least 2 min. HFMR was induced as previously described (Barber et al., 2009; Yoshihara et al., 2005). Briefly, HFMR was induced by four trains of 100 Hz stimuli spaced 2 s apart in 0.3 mM extracellular  $Ca^{2+}$ . Mini events 1 min before and up to ~10 min after HFMR induction were continuously collected. Mini frequency at indicated time points were calculated in 20-s bins. Fold enhancement was calculated by normalizing to the baseline mini frequency recorded prior to HFMR induction. Each n value represents a single muscle recording, with data generated from at least three individual larvae of each genotype.

### Ca<sup>2+</sup> Imaging

Spontaneous release events were detected as previously described (Melom et al., 2013). Briefly, wandering 3<sup>rd</sup> instar larvae expressing postsynaptic UAS-myrgCaMP6s using the *mef2-GAL4* driver were dissected in  $Ca^{2+}$ -free HL3.1. The motor neurons were cut just below the ventral nerve cord, washed several times with HL3.1 containing 1.5 mM  $Ca^{2+}$ , and allowed to rest for 5 min. Images were acquired with a PerkinElmer Ultraview Vox spinning disc confocal microscope equipped with a Hamamatsu C9100-13 Imagem EM CCD at 8–35 Hz with a 40 × 0.8 NA water-immersion objective (Carl Zeiss). A single confocal plane of muscle 4 NMJ in segment A3 was continuously imaged for 2 min before and after HFMR stimulation. Presynaptic  $Ca^{2+}$  influx during HFMR stimulation was monitored in larvae expressing presynaptic UAS-myrgCaMP6s driven by the pan neuronal C155 *elav*-GAL4 driver.

### Statistical Analysis

All error bars are SEM. Statistical significance was determined using one-way ANOVA with post hoc Tukey analysis for multiple comparisons and Student's t test when only two groups were compared (n.s.:  $p > 0.05$ , \* $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ , and \*\*\*\* $p < 0.0001$ ).

### ACCESSION NUMBERS

The Genbank accession number for the Cpx sequence reported in this paper is Genbank: NP\_525019.

## SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, three tables, and Supplemental Experimental Procedures and can be found with this article online at <http://dx.doi.org/10.1016/j.neuron.2015.10.011>.

## AUTHOR CONTRIBUTIONS

R.W.C. performed and analyzed the electrophysiology, confocal imaging, biochemistry, and genetics. L.K.B. performed and analyzed confocal imaging and designed experiments. D.V. performed the *in vitro* kinase experiment. Y.A. performed and analyzed the GCaMP6s experiments. A.T. and R.W.C. performed and analyzed the Cpx/SNARE pull-down experiments. F.L. performed ITC experiments. J.T.L. supervised the project. R.W.C. and J.T.L. designed experiments and wrote the paper.

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## REFERENCES

- Aberle, H., Haghghi, A.P., Fetter, R.D., McCabe, B.D., Magalhães, T.R., and Goodman, C.S. (2002). wishful thinking encodes a BMP type II receptor that regulates synaptic growth in *Drosophila*. *Neuron* 33, 545–558.
- Ahmad, M., Polepalli, J.S., Goswami, D., Yang, X., Kaeser-Woo, Y.J., Südhof, T.C., and Malenka, R.C. (2012). Postsynaptic complexin controls AMPA receptor exocytosis during LTP. *Neuron* 73, 260–267.
- Andraea, L.C., and Burrone, J. (2015). Spontaneous Neurotransmitter Release Shapes Dendritic Arbors via Long-Range Activation of NMDA Receptors. *Cell Rep.* February 12, 2015. <http://dx.doi.org/10.1016/j.celrep.2015.01.032>.
- Ataman, B., Ashley, J., Gorczyca, M., Ramachandran, P., Fouquet, W., Sigrist, S.J., and Budnik, V. (2008). Rapid activity-dependent modifications in synaptic structure and function require bidirectional Wnt signaling. *Neuron* 57, 705–718.
- Atasoy, D., Ertunc, M., Moulder, K.L., Blackwell, J., Chung, C., Su, J., and Kavalali, E.T. (2008). Spontaneous and evoked glutamate release activates two populations of NMDA receptors with limited overlap. *J. Neurosci.* 28, 10151–10166.
- Ballard, S.L., Miller, D.L., and Ganetzky, B. (2014). Retrograde neurotrophin signaling through Tollo regulates synaptic growth in *Drosophila*. *J. Cell Biol.* 204, 1157–1172.
- Barber, C.F., Jorquera, R.A., Melom, J.E., and Littleton, J.T. (2009). Postsynaptic regulation of synaptic plasticity by synaptotagmin 4 requires both C2 domains. *J. Cell Biol.* 187, 295–310.
- Berke, B., Wittnam, J., McNeill, E., Van Vactor, D.L., and Keshishian, H. (2013). Retrograde BMP signaling at the synapse: a permissive signal for synapse maturation and activity-dependent plasticity. *J. Neurosci.* 33, 17937–17950.
- Brand, A.H., and Perrimon, N. (1993). Targeted gene expression as a means of altering cell fates and generating dominant phenotypes. *Development* 118, 401–415.
- Budnik, V., Zhong, Y., and Wu, C.F. (1990). Morphological plasticity of motor axons in *Drosophila* mutants with altered excitability. *J. Neurosci.* 10, 3754–3768.
- Buhl, L.K., Jorquera, R.A., Akbergenova, Y., Huntwork-Rodriguez, S., Volfson, D., and Littleton, J.T. (2013). Differential regulation of evoked and spontaneous neurotransmitter release by C-terminal modifications of complexin. *Mol. Cell. Neurosci.* 52, 161–172.
- Bykhovskaia, M., Jagota, A., Gonzalez, A., Vasin, A., and Littleton, J.T. (2013). Interaction of the complexin accessory helix with the C-terminus of the SNARE complex: molecular-dynamics model of the fusion clamp. *Biophys. J.* 105, 679–690.
- Carter, A.G., and Regehr, W.G. (2002). Quantal events shape cerebellar inter-neuron firing. *Nat. Neurosci.* 5, 1309–1318.
- Cho, R.W., Song, Y., and Littleton, J.T. (2010). Comparative analysis of *Drosophila* and mammalian complexins as fusion clamps and facilitators of neurotransmitter release. *Mol. Cell. Neurosci.* 45, 389–397.
- Cho, R.W., Kümmel, D., Li, F., Baguley, S.W., Coleman, J., Rothman, J.E., and Littleton, J.T. (2014). Genetic analysis of the Complexin trans-clamping model for cross-linking SNARE complexes *in vivo*. *Proc. Natl. Acad. Sci. USA* 111, 10317–10322.
- Choi, B.J., Imlach, W.L., Jiao, W., Wolfram, V., Wu, Y., Grbic, M., Cela, C., Baines, R.A., Nitabach, M.N., and McCabe, B.D. (2014). Miniature neurotransmission regulates *Drosophila* synaptic structural maturation. *Neuron* 82, 618–634.
- Davis, G.W., and Müller, M. (2015). Homeostatic control of presynaptic neurotransmitter release. *Annu. Rev. Physiol.* 77, 251–270.
- Davis, G.W., Schuster, C.M., and Goodman, C.S. (1996). Genetic dissection of structural and functional components of synaptic plasticity. III. CREB is necessary for presynaptic functional plasticity. *Neuron* 17, 669–679.
- Dietzl, G., Chen, D., Schnorrer, F., Su, K.C., Barinova, Y., Fellner, M., Gasser, B., Kinsey, K., Oettel, S., Scheiblaue, S., et al. (2007). A genome-wide transgenic RNAi library for conditional gene inactivation in *Drosophila*. *Nature* 448, 151–156.
- Fatt, P., and Katz, B. (1952). Spontaneous subthreshold activity at motor nerve endings. *J. Physiol.* 117, 109–128.
- Frank, C.A., Pielage, J., and Davis, G.W. (2009). A presynaptic homeostatic signaling system composed of the Eph receptor, ephexin, Cdc42, and CaV2.1 calcium channels. *Neuron* 61, 556–569.
- Genc, O., Kochubey, O., Toonen, R.F., Verhage, M., and Schneggenburger, R. (2014). Munc18-1 is a dynamically regulated PKC target during short-term enhancement of transmitter release. *eLife* 3, e01715.
- Geppert, M., Goda, Y., Hammer, R.E., Li, C., Rosahl, T.W., Stevens, C.F., and Südhof, T.C. (1994). Synaptotagmin I: a major Ca<sup>2+</sup> sensor for transmitter release at a central synapse. *Cell* 79, 717–727.
- Giraudo, C.G., Eng, W.S., Melia, T.J., and Rothman, J.E. (2006). A clamping mechanism involved in SNARE-dependent exocytosis. *Science* 313, 676–680.
- Goold, C.P., and Davis, G.W. (2007). The BMP ligand Gbb gates the expression of synaptic homeostasis independent of synaptic growth control. *Neuron* 56, 109–123.
- Guan, Z., Buhl, L.K., Quinn, W.G., and Littleton, J.T. (2011). Altered gene regulation and synaptic morphology in *Drosophila* learning and memory mutants. *Learn. Mem.* 18, 191–206.
- Habets, R.L., and Borst, J.G. (2005). Post-tetanic potentiation in the rat calyx of Held synapse. *J. Physiol.* 564, 173–187.
- Hobson, R.J., Liu, Q., Watanabe, S., and Jorgensen, E.M. (2011). Complexin maintains vesicles in the primed state in *C. elegans*. *Curr. Biol.* 21, 106–113.
- Huganir, R.L., and Nicoll, R.A. (2013). AMPARs and synaptic plasticity: the last 25 years. *Neuron* 80, 704–717.
- Huntwork, S., and Littleton, J.T. (2007). A complexin fusion clamp regulates spontaneous neurotransmitter release and synaptic growth. *Nat. Neurosci.* 10, 1235–1237.
- Iyer, J., Wahlmark, C.J., Kuser-Ahnert, G.A., and Kawasaki, F. (2013). Molecular mechanisms of COMPLEXIN fusion clamp function in synaptic exocytosis revealed in a new *Drosophila* mutant. *Mol. Cell. Neurosci.* 56, 244–254.
- Jorquera, R.A., Huntwork-Rodriguez, S., Akbergenova, Y., Cho, R.W., and Littleton, J.T. (2012). Complexin controls spontaneous and evoked neurotransmitter release by regulating the timing and properties of synaptotagmin activity. *J. Neurosci.* 32, 18234–18245.

- Kaesler-Woo, Y.J., Yang, X., and Südhof, T.C. (2012). C-terminal complexin sequence is selectively required for clamping and priming but not for Ca<sup>2+</sup>-triggering of synaptic exocytosis. *J. Neurosci.* *32*, 2877–2885.
- Kavalali, E.T. (2015). The mechanisms and functions of spontaneous neurotransmitter release. *Nat. Rev. Neurosci.* *16*, 5–16.
- Korogod, N., Lou, X., and Schneggenburger, R. (2005). Presynaptic Ca<sup>2+</sup> requirements and developmental regulation of posttetanic potentiation at the calyx of Held. *J. Neurosci.* *25*, 5127–5137.
- Lagow, R.D., Bao, H., Cohen, E.N., Daniels, R.W., Zuzek, A., Williams, W.H., Macleod, G.T., Sutton, R.B., and Zhang, B. (2007). Modification of a hydrophobic layer by a point mutation in syntaxin 1A regulates the rate of synaptic vesicle fusion. *PLoS Biol.* *5*, e72.
- Lee, M.C., Yasuda, R., and Ehlers, M.D. (2010). Metaplasticity at single glutamatergic synapses. *Neuron* *66*, 859–870.
- Littleton, J.T., Stern, M., Schulze, K., Perin, M., and Bellen, H.J. (1993). Mutational analysis of *Drosophila* synaptotagmin demonstrates its essential role in Ca(2+)-activated neurotransmitter release. *Cell* *74*, 1125–1134.
- Littleton, J.T., Chapman, E.R., Kreber, R., Garment, M.B., Carlson, S.D., and Ganetzky, B. (1998). Temperature-sensitive paralytic mutations demonstrate that synaptic exocytosis requires SNARE complex assembly and disassembly. *Neuron* *21*, 401–413.
- Lou, X., Korogod, N., Brose, N., and Schneggenburger, R. (2008). Phorbol esters modulate spontaneous and Ca<sup>2+</sup>-evoked transmitter release via acting on both Munc13 and protein kinase C. *J. Neurosci.* *28*, 8257–8267.
- Malsam, J., Parisotto, D., Bharat, T.A., Scheutzow, A., Krause, J.M., Briggs, J.A., and Söllner, T.H. (2012). Complexin arrests a pool of docked vesicles for fast Ca<sup>2+</sup>-dependent release. *EMBO J.* *31*, 3270–3281.
- Marqués, G., Bao, H., Haerry, T.E., Shimell, M.J., Duchek, P., Zhang, B., and O'Connor, M.B. (2002). The *Drosophila* BMP type II receptor Wishful Thinking regulates neuromuscular synapse morphology and function. *Neuron* *33*, 529–543.
- Martin, J.A., Hu, Z., Fenz, K.M., Fernandez, J., and Dittman, J.S. (2011). Complexin has opposite effects on two modes of synaptic vesicle fusion. *Curr. Biol.* *21*, 97–105.
- Maximov, A., Tang, J., Yang, X., Pang, Z.P., and Südhof, T.C. (2009). Complexin controls the force transfer from SNARE complexes to membranes in fusion. *Science* *323*, 516–521.
- Mayford, M., Siegelbaum, S.A., and Kandel, E.R. (2012). Synapses and memory storage. *Cold Spring Harb. Perspect. Biol.* *4*, <http://dx.doi.org/10.1101/cshperspect.a005751>.
- McCabe, B.D., Marqués, G., Haghighi, A.P., Fetter, R.D., Crotty, M.L., Haerry, T.E., Goodman, C.S., and O'Connor, M.B. (2003). The BMP homolog Gbb provides a retrograde signal that regulates synaptic growth at the *Drosophila* neuromuscular junction. *Neuron* *39*, 241–254.
- McKinney, R.A., Capogna, M., Dürr, R., Gähwiler, B.H., and Thompson, S.M. (1999). Miniature synaptic events maintain dendritic spines via AMPA receptor activation. *Nat. Neurosci.* *2*, 44–49.
- McMahon, H.T., Missler, M., Li, C., and Südhof, T.C. (1995). Complexins: cytosolic proteins that regulate SNAP receptor function. *Cell* *83*, 111–119.
- Melom, J.E., Akbergenova, Y., Gavornik, J.P., and Littleton, J.T. (2013). Spontaneous and evoked release are independently regulated at individual active zones. *J. Neurosci.* *33*, 17253–17263.
- Paradis, S., Sweeney, S.T., and Davis, G.W. (2001). Homeostatic control of presynaptic release is triggered by postsynaptic membrane depolarization. *Neuron* *30*, 737–749.
- Peled, E.S., and Isacoff, E.Y. (2011). Optical quantal analysis of synaptic transmission in wild-type and rab3-mutant *Drosophila* motor axons. *Nat. Neurosci.* *14*, 519–526.
- Piccoli, Z.D., and Littleton, J.T. (2014). Retrograde BMP signaling modulates rapid activity-dependent synaptic growth via presynaptic LIM kinase regulation of cofilin. *J. Neurosci.* *34*, 4371–4381.
- Schaub, J.R., Lu, X., Doneske, B., Shin, Y.K., and McNew, J.A. (2006). Hemifusion arrest by complexin is relieved by Ca<sup>2+</sup>-synaptotagmin I. *Nat. Struct. Mol. Biol.* *13*, 748–750.
- Schuster, C.M., Davis, G.W., Fetter, R.D., and Goodman, C.S. (1996). Genetic dissection of structural and functional components of synaptic plasticity. II. Fasciclin II controls presynaptic structural plasticity. *Neuron* *17*, 655–667.
- Shata, A., Saisu, H., Odani, S., and Abe, T. (2007). Phosphorylated synaphin/complexin found in the brain exhibits enhanced SNARE complex binding. *Biochem. Biophys. Res. Commun.* *354*, 808–813.
- Sigrist, S.J., Reiff, D.F., Thiel, P.R., Steinert, J.R., and Schuster, C.M. (2003). Experience-dependent strengthening of *Drosophila* neuromuscular junctions. *J. Neurosci.* *23*, 6546–6556.
- Snead, D., Wragg, R.T., Dittman, J.S., and Eliezer, D. (2014). Membrane curvature sensing by the C-terminal domain of complexin. *Nat. Commun.* *5*, 4955.
- Südhof, T.C. (2013). Neurotransmitter release: the last millisecond in the life of a synaptic vesicle. *Neuron* *80*, 675–690.
- Südhof, T.C., and Rothman, J.E. (2009). Membrane fusion: grappling with SNARE and SM proteins. *Science* *323*, 474–477.
- Sutton, M.A., Ito, H.T., Cressy, P., Kempf, C., Woo, J.C., and Schuman, E.M. (2006). Miniature neurotransmission stabilizes synaptic function via tonic suppression of local dendritic protein synthesis. *Cell* *125*, 785–799.
- Tang, J., Maximov, A., Shin, O.H., Dai, H., Rizo, J., and Südhof, T.C. (2006). A complexin/synaptotagmin 1 switch controls fast synaptic vesicle exocytosis. *Cell* *126*, 1175–1187.
- Ubersax, J.A., and Ferrell, J.E., Jr. (2007). Mechanisms of specificity in protein phosphorylation. *Nat. Rev. Mol. Cell Biol.* *8*, 530–541.
- Weber, T., Zemelman, B.V., McNew, J.A., Westermann, B., Gmachl, M., Parlati, F., Söllner, T.H., and Rothman, J.E. (1998). SNAREpins: minimal machinery for membrane fusion. *Cell* *92*, 759–772.
- Wong, Y.H., Lee, T.Y., Liang, H.K., Huang, C.M., Wang, T.Y., Yang, Y.H., Chu, C.H., Huang, H.D., Ko, M.T., and Hwang, J.K. (2007). KinasePhos 2.0: a web server for identifying protein kinase-specific phosphorylation sites based on sequences and coupling patterns. *Nucleic Acids Res.* *35*, W588–W594.
- Wragg, R.T., Snead, D., Dong, Y., Ramlall, T.F., Menon, I., Bai, J., Eliezer, D., and Dittman, J.S. (2013). Synaptic vesicles position complexin to block spontaneous fusion. *Neuron* *77*, 323–334.
- Xu, J., Pang, Z.P., Shin, O.H., and Südhof, T.C. (2009). Synaptotagmin-1 functions as a Ca<sup>2+</sup> sensor for spontaneous release. *Nat. Neurosci.* *12*, 759–766.
- Xue, M., Ma, C., Craig, T.K., Rosenmund, C., and Rizo, J. (2008). The Janus-faced nature of the C(2)B domain is fundamental for synaptotagmin-1 function. *Nat. Struct. Mol. Biol.* *15*, 1160–1168.
- Xue, M., Lin, Y.Q., Pan, H., Reim, K., Deng, H., Bellen, H.J., and Rosenmund, C. (2009). Tilting the balance between facilitatory and inhibitory functions of mammalian and *Drosophila* Complexins orchestrates synaptic vesicle exocytosis. *Neuron* *64*, 367–380.
- Xue, M., Craig, T.K., Xu, J., Chao, H.T., Rizo, J., and Rosenmund, C. (2010). Binding of the complexin N terminus to the SNARE complex potentiates synaptic-vesicle fusogenicity. *Nat. Struct. Mol. Biol.* *17*, 568–575.
- Yoshihara, M., Adolfsen, B., Galle, K.T., and Littleton, J.T. (2005). Retrograde signaling by Syt 4 induces presynaptic release and synapse-specific growth. *Science* *310*, 858–863.
- Zhong, Y., and Wu, C.F. (1991). Altered synaptic plasticity in *Drosophila* memory mutants with a defective cyclic AMP cascade. *Science* *251*, 198–201.
- Zhong, Y., and Wu, C.F. (2004). Neuronal activity and adenylyl cyclase in environment-dependent plasticity of axonal outgrowth in *Drosophila*. *J. Neurosci.* *24*, 1439–1445.
- Zhong, Y., Budnik, V., and Wu, C.F. (1992). Synaptic plasticity in *Drosophila* memory and hyperexcitable mutants: role of cAMP cascade. *J. Neurosci.* *12*, 644–651.
- Zucker, R.S., and Regehr, W.G. (2002). Short-term synaptic plasticity. *Annu. Rev. Physiol.* *64*, 355–405.