Munc13-1 is a Ca\textsuperscript{2+}-phospholipid-dependent vesicle priming hub that shapes synaptic short-term plasticity and enables sustained neurotransmission

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**In brief**
Using novel knockin mouse models, Lipstein et al. show that Ca\textsuperscript{2+}-phospholipid binding activates the presynaptic protein Munc13-1 to fine-tune the rate of synaptic vesicle replenishment according to synaptic activity. This process determines short-term synaptic plasticity and the temporal fidelity of synaptic transmission in the auditory brainstem and the hippocampus.

**Highlights**
- The Munc13-1 C\textsubscript{2}B domain controls synaptic vesicle replenishment rates
- Blocking Ca\textsuperscript{2+}-phospholipid-C\textsubscript{2}B signaling attenuates vesicle replenishment
- Enhancing Ca\textsuperscript{2+}-phospholipid-C\textsubscript{2}B signaling accelerates vesicle replenishment
- This process determines short-term plasticity and fidelity of synaptic transmission
Munc13-1 is a Ca²⁺-phospholipid-dependent vesicle priming hub that shapes synaptic short-term plasticity and enables sustained neurotransmission

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SUMMARY

During ongoing presynaptic action potential (AP) firing, transmitter release is limited by the availability of release-ready synaptic vesicles (SVs). The rate of SV recruitment (SVR) to release sites is strongly upregulated at high AP frequencies to balance SV consumption. We show that Munc13-1—an essential SV priming protein—regulates SVR via a Ca²⁺-phospholipid-dependent mechanism. Using knockin mouse lines with point mutations in the Ca²⁺-phospholipid-binding C₂B domain of Munc13-1, we demonstrate that abolishing Ca²⁺-phospholipid binding increases synaptic depression, slows recovery of synaptic strength after SV pool depletion, and reduces temporal fidelity of synaptic transmission, while increased Ca²⁺-phospholipid binding has the opposite effects. Thus, Ca²⁺-phospholipid binding to the Munc13-1-C₂B domain accelerates SVR, reduces short-term synaptic depression, and increases the endurance and temporal fidelity of neurotransmission, demonstrating that Munc13-1 is a core vesicle priming hub that adjusts SV re-supply to demand.

INTRODUCTION

Neuronal signaling at chemical synapses requires transmitter release by synaptic vesicle (SV) fusion. A fraction of SVs residing at molecularly defined release sites constitutes the pool of readily releasable SVs (RRP) (Kaeser and Regehr, 2017), which are in a molecularly mature—“primed”—state to allow rapid membrane fusion upon an action potential (AP)-induced increase in cytosolic Ca²⁺ ([Ca²⁺]ᵢ). To maintain synaptic transmission during continuous activity, empty release sites must be re-populated with fusion-competent SVs (SV recruitment [SVR]).

SVR is generally slow, with a time constant of several seconds (Fuhrmann et al., 2004; Pyott and Rosenmund, 2002; Stevens and Tsujimoto, 1995; von Gersdorff et al., 1997; Wesseling and Lo, 2002), but can be transiently accelerated by an order of magnitude following bouts of synaptic activity (Sakaba and Neher, 2001a; Stevens and Wesseling, 1998), most likely due to the buildup of presynaptic [Ca²⁺]ᵢ (Dittman and Regehr, 1998; Sakaba and Neher, 2001a). This enables synapses to maintain high levels of transmitter release during presynaptic AP firing, resulting in reduced synaptic short-term depression (STD), and to speed up the recovery of synaptic strength following activity while [Ca²⁺]ᵢ decays back to resting levels (Dittman et al., 2000; Fuhrmann et al., 2004; Wang and Kaczmarek, 1998). By counteracting SV pool depletion, the SVR speed is a key determinant of synaptic strength and short-term plasticity (STP) and affects multiple complex brain processes (Zucker and Regehr, 2002).

Because [Ca²⁺]ᵢ controls several steps of the SV cycle and regulates numerous Ca²⁺-binding proteins (for review, see Alabi and Tsien, 2012 and Neher and Sakaba, 2008), the identity of proteins that mediate activity-dependent SVR acceleration has remained enigmatic. Moreover, beyond their [Ca²⁺]ᵢ sensitivity, SVR-regulating pathways also respond to lipid second messengers. During activity, the presynaptic membrane lipid composition changes as a consequence of SV fusion and the activity of lipid-modifying enzymes, some of which are Ca²⁺ regulated. In neuroendocrine cells, for example, vesicle fusion occurs at membrane sites rich in PIP₂ (phosphatidylinositol-4,5-bisphosphate) and PIP₂ augments RRP replenishment (Kabachinski et al., 2014; Milosevic et al., 2005; Walter et al., 2017). PIP₂ conversion to diacylglycerol (DAG) regulates transmitter release in cultured neurons (Rhee et al., 2002; Wierda et al., 2007) and in situ (Lou et al., 2008; Malenka et al., 1987).
The present study was designed to demonstrate a Ca\(^{2+}\)- and phospholipid-controlled step in SVR and STP regulation. We focused on the SV priming protein Munc13-1, a presynaptic active zone (AZ) component that is essential for transmitter release (Augustin et al., 1999b; Varoqueaux et al., 2002). Munc13-1 operates by regulating the conformation of the soluble N-ethylmaleimide-sensitive-factor attachment receptor (SNARE) fusion protein syntaxin, thus promoting partial SNARE complex formation and close SV-plasma membrane contact (Imig et al., 2014; Lai et al., 2017; Ma et al., 2011). The Munc13-1 MUN domain, which mediates SV priming, is located downstream of three regulatory domains: a Ca\(^{2+}\)-calmodulin (Ca\(^{2+}\)-CaM) binding motif (Junge et al., 2004; Lipstein et al., 2012a, 2013; Piotrowski et al., 2020) a DAG-binding C\(_2\) domain (Betz, 1998), and a C\(_2\) domain (C\(_2\)B) that binds phospholipids in a Ca\(^{2+}\)-dependent manner (Michelassi et al., 2017; Shin et al., 2010). These regulatory domains profoundly affect Munc13-1 activity in vitro (Junge et al., 2004; Rhee et al., 2002; Shin et al., 2010). However, a block of CaM binding to Munc13-1 has only subtle effects on SV priming during and after AP trains in intact circuits (Lipstein et al., 2013), demonstrating that major signaling pathways involved in SVR regulation are still unknown. In this context, the C\(_2\)B domain (Shin et al., 2010) and characterized functional consequences in the calyx of Held synapse. We demonstrate that presynaptic activity activates Munc13-1 by Ca\(^{2+}\)-phospholipid binding to its C\(_2\)B domain. This causes distinctly higher SV rates and thus shapes STP, enables sustained transmission, and enhances the temporal fidelity of synaptic signaling.

RESULTS

Generation of Munc13-1 C\(_2\)BDN and C\(_2\)BKW KI mice

The Ca\(^{2+}\)-binding site in the Munc13-1 C\(_2\)B domain is composed of negatively charged residues that chelate Ca\(^{2+}\), thus allowing negatively charged phospholipids to bind. We exchanged two such aspartic acid residues by asparagine (D705N and D711N) to create the Munc13-1 C\(_2\)BDN KI line (Shin et al., 2010). This exchange abolishes Ca\(^{2+}\) binding, slightly increases phospholipid binding at low [Ca\(^{2+}\)] (likely due to less membrane repulsion), and abolishes Ca\(^{2+}\)-mediated phospholipid binding at high [Ca\(^{2+}\)] (Shin et al., 2010). To increase the C\(_2\)B domain affinity to phospholipids, we created the Munc13-1 C\(_2\)BKW KI, targeting a lysine (K706) in the outer rim of the Ca\(^{2+}\)-binding site. In the synaptotagmin-1 (Syt1) C\(_2\)A domain, the corresponding residue is large and hydrophobic (M173), inserts into membrane bilayers, and enhances Ca\(^{2+}\)-dependent phospholipid binding (Chapman and Davis, 1998; Gerber et al., 2002; Shin et al., 2010; Zhang et al., 1998). We replaced K706 with tryptophan (K706W), which increases PIP\(_2\) binding without changing the Ca\(^{2+}\) affinity of the C\(_2\)B domain (Shin et al., 2010; Figures 1A–1D).

Homozygous mutant (mut) KIs of both lines were viable and fertile but showed an ~30% body weight reduction at 2 weeks of age (Figure 1E), which did not persist. Neither KI line showed signs of distress or gross behavioral abnormalities. Analyses of expression levels of a selected set of presynaptic proteins did not reveal any genotype-related differences (Figures 1F and S1A–S1C). The localization of the mutant Munc13-1 variants in presynaptic compartments contacting principal neurons (PNs) of the medial nucleus of the trapezoid body (MNTB) was examined by immunostaining (Figure S1D). A similar typical pattern of Munc13-1-positive structures co-localizing with the AZ marker Bassoon and surrounding MNTB PN somata was observed in wild-type (WT) and KI samples (Figures S1D–S1F; Chen et al., 2013; Lipstein et al., 2013).

In sum, we generated two novel KI mouse lines with specific point mutations in the Munc13-1 C\(_2\)B domain that abolish (C\(_2\)BDN) or increase (C\(_2\)BKW) Ca\(^{2+}\)-dependent phospholipid binding without altering Munc13-1 expression or localization.

Basal transmission and presynaptic Ca\(^{2+}\) influx in Munc13-1 C\(_2\)BDN and C\(_2\)BKW synapses

To assess the role of the Munc13-1 C\(_2\)B domain in synaptic transmission and STP, we used the calyx of Held synapse, as it is accessible to pre- and postsynaptic recordings (Borst et al., 1995; Forsythe, 1994). Recordings were obtained from homozgyous mut and WT littermates (mut\(_{\text{DN}}\) and WT\(_{\text{DN}}\) for C\(_2\)BDN KIs; mut\(_{\text{KW}}\) and WT\(_{\text{KW}}\) for C\(_2\)BKW) at postnatal day (P) 14–P17, i.e., after hearing onset, when Munc13-1 is functionally dominant (Chen et al., 2013). Only minor differences were observed between WT\(_{\text{DN}}\) and WT\(_{\text{KW}}\) (Table S1).

To assay synaptic strength and evoked excitatory postsynaptic current (eEPSC) kinetics, we recorded unitary eEPSCs in voltage-clamped MNTB PNs (Figure 2A). Unless stated otherwise, recordings were obtained in the presence of 1 mM kynurenic acid (kyn) (STAR Methods; Figure S2). In C\(_2\)BDN mice, we observed larger eEPSCs in mut DN as compared to WT DN (Figures 2A–2C; Table S1), whereas in C\(_2\)BKW mice, eEPSC amplitudes were indistinguishable between mut\(_{\text{KW}}\) and WT\(_{\text{KW}}\). eEPSC kinetics were unaltered (Figures 2D and 2E; Table S1). Scatter-plots of eEPSC rise time versus eEPSC half-width revealed a positive correlation (Figure S2), reflecting a developmental eEPSC shortening (Joshi et al., 2004; Koike-Tani et al., 2005; Taschenberger and von Gersdorff, 2000). This correlation was similar in all genotypes, indicating no adverse effects of KI mutations on developmental synapse refinement.

We next measured paired-pulse ratios (PPRs) (PPR = eEPSC\(_2\)/eEPSC\(_1\)) of two consecutive eEPSCs evoked at inter-stimulus intervals (ISIs) of 5 ms–2 s (Figures 2D and 2E). In many synapses, including the calyx of Held (Debanne et al., 1996; Dobrunz and Stevens, 1997; Taschenberger et al., 2016), PPRs correlate with initial eEPSCs size, indicating that differences in synaptic strength arise, at least partly, from differences in release probability. Regression lines fitted to log-linear plots of 1 – PPR versus ISI provide estimates for average release probabilities (\(\overline{p}\)) and average SV pool replenishment rate constants (\(K_\text{r}\)) (Betz, 1970). We observed lower PPRs at all ISIs in mut\(_{\text{DN}}\) synapses (Figure 2E1; Table S1) as compared to WT\(_{\text{DN}}\) synapses, indicative of elevated \(\overline{p}\) in the former. In
contrast, PPRs were similar in mutKW and WT KW synapses, consistent with unchanged $\bar{P}$. However, a steeper slope of the log-linear plot of $1/C_0$ PPR versus ISI indicated a faster $k_+$ in mutKW synapses (Figure 2E).

To test whether enhanced presynaptic Ca$^{2+}$ influx contributes to the elevated $\bar{P}$ in mutDN synapses, we recorded pharmacologically isolated voltage-gated Ca$^{2+}$ currents ($I_{Ca(V)}$) in voltage-clamped calyx terminals (Figures 2F and 2G). Peak $I_{Ca(V)}$ amplitudes were similar in mut and WT terminals, indicating similar voltage-gated Ca$^{2+}$ channel (VGCC) densities (Figure 2G; Table S1). To assess VGCC gating kinetics, we quantified the charge of $I_{Ca(V)}$ ($Q_{Ca(V)}$) elicited by short AP-like depolarizations, assuming that altered activation or deactivation of $I_{Ca(V)}$ will affect its current integral (Figure 2F; Li et al., 2007), and detected no differences (Figure 2G; Table S1).

These data show that basal synaptic transmission is intact in C2BDN and C2BKW KI synapses. No indications of developmental abnormalities or postsynaptic modifications were observed, and normal eEPSC kinetics indicate unaltered SV fusion kinetics. The C2BDN mutation leads to enhanced synaptic strength, partly due to elevated $\bar{P}$, but neither mutation alters VGCC expression or kinetics. Thus, the increased $\bar{P}$ in mutDN synapses is either caused by a subtle change in AP waveform not affecting eEPSC kinetics or, more likely, by a mechanism downstream of Ca$^{2+}$ influx that increases the apparent Ca$^{2+}$ sensitivity of SVs in mutDN terminals. The C2BKW mutation
does not affect eEPSC amplitudes but accelerates the recovery from STD induced by single eEPSCs.

The Munc13-1 C2B domain regulates steady-state rates of quantal release during repetitive synaptic stimulation

During AP trains, calyx of Held synapses typically show STD at low and intermediate (<50 Hz) stimulus frequencies (\(f_{stim}\)) and sometimes transient facilitation followed by STD at higher \(f_{stim}\) (≥ 50 Hz; Grande and Wang, 2011; Taschenberger et al., 2016). Quantal release at steady state is limited by the SVR rate. To examine the role of the Munc13-1 C2B domain in determining STD, we recorded eEPSC trains for a range of \(f_{stim}\) (0.5–200 Hz, 35 APs; Figures 3A1 and 3A2). Normalized average eEPSC train amplitudes showed stronger STD at high \(f_{stim}\) for mutDN as compared to WTDN synapses (Figure 3B1; Table S1), whereas reduced STD was seen for all but the highest \(f_{stim}\) in mutKW as compared to WTKW synapses (Figure 3B2; Table S1).

The relationship between steady-state quantal release and \(f_{stim}\) is illustrated in Figure 3C by plotting eEPSC\(_{ss}\) and average eEPSC train amplitudes showed stronger STD at high \(f_{stim}\) for mutDN as compared to WTDN synapses (Figure 3B2; Table S1). The number of readily releasable SVs is not altered in Munc13-1 C2BDN and C2BKW synapses

The number of readily releasable SVs is not altered in Munc13-1 C2BDN and C2BKW synapses

Given the key role of Munc13-1 in RRP establishment and maintenance (Augustin et al., 1999b; Varoquaux et al., 2002), we next tested whether the C2BDN and C2BKW mutations affect the RRP. Pool size estimates corresponding to the subpool of “fast releasing” SVs (FRP) were obtained from cumulative eEPSC amplitudes measured during high-frequency trains and corrected for ongoing STD (Schneggenburger et al., 1999; Figure 4A). An estimate for \(\bar{p}\) is obtained from the ratio eEPSC/FRP. FRP estimates increased and \(\bar{p}\) values decreased with increasing \(f_{stim}\), likely because of more complete pool depletion (Figures 4A and 4B). We therefore termed these estimates FRP\(_{corr}\) and \(\bar{p}\). When plotting 1/FRP\(_{corr}\) and \(\bar{p}\) versus ISI, we observed a roughly linear relationship between these quantities and ISI. Assuming this relationship holds for the entire ISI range, we obtained corrected estimates for pool size (FRP\(_{corr}\)) and release probability (\(\bar{p}_{corr}\)) by extrapolation to ISI = 0 s. FRP\(_{corr}\) values were slightly larger when comparing C2BDN and C2BKW mut with littermate WT synapses (Figure 4B; Table S1) but statistically not significantly different, although \(\bar{p}_{corr}\) was significantly higher in mutDN synapses (Figure 4B; Table S1), consistent with their larger initial eEPSCs and reduced PPR (Figures 2C and 2E).

To exclude the possibility of an overcorrection when estimating FRP\(_{corr}\), we performed three controls. (1) Assuming a simple SV pool depletion model, an estimate for release probability can be obtained by plotting eEPSC amplitudes during high-frequency trains versus the cumulative sum of previous eEPSCs (Elmqvist and Quastel, 1965; Neher, 2015). Such plots show steeper negative slopes for mutDN as compared to WTDN synapses, consistent with their larger initial eEPSCs and reduced PPR (Figures 2C and 2E).
(legend on next page)
with more rapid SV consumption and therefore higher $\bar{p}$ in the former (Figure S3A1). In contrast, slopes and, therefore, $\bar{p}$ estimates were similar in WT KW and mut KW synapses (Figure S3A2). Both findings confirm our earlier conclusions (Figure 4B). (2) For a second set of control experiments, we recorded 200-Hz eEPSC trains in WT synapses before and after bath application of tetraethylammonium (TEA), a blocker of voltage-gated potassium channels that broadens calyceal APs and increases AP-evoked Ca$^{2+}$ influx and $\bar{p}$ (Ishikawa et al., 2003; Figure S3B), and observed an apparent FRP increase of ~$-40\%$ in the presence of TEA. This is in accord with the average ratio between FRP$_{corr}$ and FRP$_{200\text{ Hz}}$ in both WT$_{DN}$ and also WT$_{KW}$ synapses (Figure S3C), indicating that our strategy to correct FRP estimates for incomplete pool depletion (Figure 4B) does not result in a substantial overestimate. (3) Finally, we assayed SV fusion by measuring $\Delta C_{fr}$ in response to depolarizing voltage steps of incrementing duration in voltage-clamped calyces. $\Delta C_{fr}$-based pool estimates do not rely on postsynaptic responses and thus are not susceptible to AMPA receptor (AMPAR) saturation or desensitization. However, long-lasting presynaptic depolarizations—during which presynaptic [Ca$^{2+}$], spatially equilibrates—trigger release from an additional subpool of “slowly releasing” SVs (SRP; Sakaba and Neher, 2001b), while during short and spatially restricted AP-evoked [Ca$^{2+}$] elevations, SVs of the SRP fuse to a much lesser extent (Sakaba, 2006). Increasing $I_{c$ahl} duration led to larger $\Delta C_{fr}$ (Figure 4C). The average relationship between quantal release and Ca$^{2+}$ influx duration was very similar in WT and mutant C$_{2BDN}$ or mutant C$_{2BKW}$ calyces with respect to both $\Delta C_{fr}$ amplitudes and release kinetics (Figure 4D, Table S1). Despite the caveat that $\Delta C_{fr}$ represents the sum of fast and slowly releasing SVs (FRP + SRP), these experiments do not reveal changes in SV pool size in mut$_{DN}$ or mut$_{KW}$ synapses.

These data show that the Munc13-1 C$_{2BDN}$ and C$_{2BKW}$ mutations do not affect the number of release-ready SVs in calyx terminals. Thus, changes in SV pool size are unlikely to contribute to the observed differences in steady-state release rates and STP.

### Munc13-1 C$_{2BDN}$ and C$_{2BKW}$ mutations change the average rate constants of SV recruitment

Knowing steady-state release rates and the total number of release-ready SVs in resting calyces allowed us to approximate the relationship between $f_{\text{stim}}$ and the average FRP replenishment rate constant ($\bar{k}_{r}$), which represents the rate of release site refilling per empty site averaged over one inter-stimulus interval. Figure 4E plots $\bar{k}_{r}$ versus $f_{\text{stim}}$ for WT and C$_{2B}$ mutant synapses. For WT$_{DN}$ and WT$_{KW}$ synapses, a similar relationship between $\bar{k}_{r}$ and $f_{\text{stim}}$ was observed (Figure 4E1). At the highest $f_{\text{stim}}$ of 200 Hz, $\bar{k}_{r}$ is >10-fold above its value estimated for $f_{\text{stim}} = 0.5$ Hz. The reciprocal of the maximum $\bar{k}_{r}$, of $\approx 4 \text{ s}^{-1}$ ($\bar{k}_{r} = 0.250 \text{ ms}$) agrees well with the rapid time constant of FRP replenishment measured following prolonged presynaptic depolarizations ($\tau_{1} = 270 \text{ ms}$, Lipstein et al., 2013; $\tau_{1} = 360 \text{ ms}$, Sakaba and Neher, 2001a). Figure 4E shows that the C$_{2BDN}$ and C$_{2BKW}$ mutations alter the relationship between $\bar{k}_{r}$ and $f_{\text{stim}}$ so that the dynamic range of $\bar{k}_{r}$ is reduced. However, in mut$_{DN}$ synapses, we observed a decreased $\bar{k}_{r}$ for all $f_{\text{stim}} > 2$ Hz as compared to WT values (Figure 4E2). In contrast, higher $\bar{k}_{r}$ values were estimated for all stimulation frequencies $\leq 20$ Hz in mut$_{KW}$ synapses (Figure 4E2).

These data demonstrate lower SV rate constants for mut$_{DN}$ and mut$_{KW}$ synapses mainly at high $f_{\text{stim}}$ and higher SV rate constants for mut$_{KW}$ synapses for low and intermediate $f_{\text{stim}}$, consistent with the changes observed during steady-state depression.

### Abolishing Ca$^{2+}$ binding to the Munc13-1 C$_{2B}$ domain eliminates a fast component of eEPSC recovery following high-frequency conditioning

Having established that Munc13-1 C$_{2B}$ mutations alter $\bar{k}_{r}$ during ongoing stimulation, we next tested eEPSC recovery after conditioning trains. When calyces are conditioned with low-frequency trains, eEPSCs recover with a slow time course that is well described by a single exponential (\(\tau = 4 \text{ s}\); Iwasaki and Takahashi, 2001; von Gersdorff et al., 1997). Upon high-frequency conditioning, an additional fast recovery component is observed at calyces (Wang and Kaczmarek, 1998) and other synapses (Cho et al., 2011; Gomis et al., 1999; Wang and Manis, 2008; Yang and Xu-Friedman, 2008). This fast recovery is presumed to depend on elevated [Ca$^{2+}$], and Ca$^{2+}$-CaM binding to Munc13-1 participates in the molecular signaling pathway that mediates it (Lipstein et al., 2013; Sakaba and Neher, 2001a).

To examine SVR, we applied 100-Hz (25 stimuli) or 200-Hz (50 stimuli) trains to induce strong FRP depletion and monitored recovery of synaptic strength by measuring eEPSC test amplitudes at different intervals (Figure 5A1). When plotting fractional recovery (eEPSC$_{\text{test}}$ – eEPSC$_{\text{off}}$)/eEPSC$_{\text{1}}$ – eEPSC$_{\text{off}}$) versus recovery interval (Figure 5A3; Table S1), we observed for WT$_{DN}$ synapses...
a biphasic time course requiring a fast component to account for the recovery during the initial 500 ms (Figure 5A3; Lipstein et al., 2013, their Figure 5), which was selectively and completely abolished in mutDN synapses (Figures 5A2 and 5A3). After 16 s, full recovery of eEPSCtest was seen in both mutDN and WT DN synapses. Closer inspection of the time course of fractional recovery revealed values slightly below zero in mutDN synapses for the two shortest intervals (125 and 250 ms; Figure 5A3). This can be explained by acknowledging that release probability increases during conditioning stimulation and that this synaptic facilitation decays faster than FRP recovery proceeds. Such an effect is likely occluded by the fast SVR component in WT synapses but uncovered in mutDN synapses that lack this SVR component.

These data show that a fast SVR component is absent in mutDN synapses with abolished C2B domain binding to Ca2+ and phospholipids. Because a fast SVR component is observed in WT DN synapses only after high-frequency conditioning, we expected the recovery time course after 10- or 20-Hz conditioning to be largely unaltered in mutDN as compared to WT DN synapses. This is indeed the case (Figure S4), demonstrating that the slow component of eEPSC recovery is unaffected in mutDN synapses.

**Ca2+-CaM binding to Munc13-1 is independent of Ca2+ binding to the C2B domain**

The effects of the C2BDN mutation on SVR resemble those reported previously for a Ca2+-CaM-binding-deficient Munc13-1 mutant (W464R; CaMWR; WTWR and mutWR for WT and mut littermates, respectively; Lipstein et al., 2013). To compare STP at C2BDN and CaMWR calyces under identical conditions, we performed recordings similar to those described above also for WTWR and mutWR synapses and analyzed normalized eEPSC amplitudes during stimulus trains (Figure S5A), FRP, and estimated parameters (Figures S5B–S5D); the relationship between Fstim and steady-state release (\( r \times q \)) and the apparent average replenishment rate constant \( \tilde{r}_4 \) (Figures S5E and S5F) for WTWR and mutWR synapses. While FRP and estimated parameters were unchanged, we observed lower steady-state release during high-frequency stimulation in mutWR synapses, which corresponded to reduced \( \tilde{r}_4 \). Accordingly, the dynamic range for \( \tilde{r}_4 \) was reduced as seen in mutDN synapses (Figure 4E2). As reported (Lipstein et al., 2013), we observed slower recovery after depleting stimulus trains in mutWR synapses (Figures S5G1 and S5G2). However, contrary to the C2BDN mutation and consistent with Lipstein et al. (2013), differences in the eEPSC recovery time course were less pronounced after 200- as compared to 100-Hz conditioning. The presence of residual fast SV replenishment after 200-Hz conditioning in mutWR calyx synapses indicates that the Ca2+-CaM pathway operates also in the absence of the Ca2+-CaM pathway and partially compensates for its loss.

Because of the similarity of the functional deficits between mutDN versus mutWR synapses and the spatial proximity of the C2B domain and the Ca2+-CaM binding site, we tested whether blocked Ca2+ binding to the C2B domain perturbs Ca2+-CaM binding. Co-immunoprecipitation experiments showed unaltered Ca2+-CaM binding of the C2BDN mutant Munc13-1 (Figure 5B).

These data indicate that the Munc13-1 regulation via the C2B domain is likely downstream of or independent from the regulation by Ca2+-CaM and that Ca2+ binding to the C2B domain is not a pre-requisite for Ca2+-CaM binding. Whether these pathways operate independently or synergistically to regulate Munc13-1 function and SVR remains to be studied.

**The Munc13-1 C2BKW mutation accelerates eEPSC recovery following conditioning trains**

We next analyzed SVR in C2BKW synapses. The recovery time course in mutKW synapses was profoundly accelerated as compared to WT KW (Figure 6A) and well described by a single fast exponential time constant. Little differences in fractional recovery were observed between mutKW and WT KW synapses for the two shortest recovery intervals (125 and 250 ms), while already 2 s after conditioning, eEPSCtest had either nearly completely (100 Hz) or completely (200 Hz) recovered in mutKW synapses (Figures 6A2 and 6A3; Table S1). For 200-Hz conditioning, we noticed average amplitudes of eEPSCtest slightly larger than those of eEPSC1.

To exclude that accelerated eEPSCtest recovery in mutKW synapses reflects \( \tilde{r}_4 \) augmentation rather than faster SVR, we recorded eEPSCtest pairs to monitor PPRs. If \( \tilde{r}_4 \) were indeed higher in mutKW synapses at 2- or 4-s recovery intervals as compared to 16 s, we would expect different PPRs at these time points.

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**Figure 4. Total number of releasable SVs and average initial release probability in C2BDN and C2BKW calyx synapses**

Estimates for the readily releasable SV pool were derived from high-frequency eEPSC trains (FRP; A and B) and presynaptic \( \Delta C_{rel} \) measurements (FRP + SRP; C and D).

(A) Mean cumulative eEPSC amplitudes recorded in response to 50-, 100-, and 200-Hz trains (35 APs) in C2BDN (A1) and C2BKW (A2) synapses of WT (left) and mut (right) littermates. Solid, broken, and dotted lines represent regression lines correcting for ongoing SVR for 200-, 100-, and 50-Hz trains, respectively; assuming a constant average recruitment of WT (left) and mut (right) littermates. Solid, broken, and dotted lines represent regression lines correcting for ongoing SVR for 200-, 100-, and 50-Hz trains, respectively; assuming a constant average recruitment of WT (left) and mut (right) littermates. Solid, broken, and dotted lines represent regression lines correcting for ongoing SVR for 200-, 100-, and 50-Hz trains.

(B) 1/FRP and apparent average release probability for eEPSC, \( \bar{p} = \text{eEPSC}_{1}/\text{FRP} \), plotted versus inter-stimulus interval for C2BDN (B1) and C2BKW (B2) calyx synapses. Gray and colored symbols represent mean values for WT and mut synapses, respectively. Solid lines represent linear regressions to the scatterplots. Intersections of the line fits with the abscissa (at ISI = 0 ms) represent corrected estimates for 1/FRPcorr and \( \bar{p}_\text{corr} \). (C) Traces of presynaptic \( \Delta C_{rel} \), (left) and \( \Delta C_{rel} \), (right) elicited by step depolarizations of 1, 2, 5, 10, and 20 ms duration in voltage-clamped C2BDN (C1) and C2BKW (C2) calyx terminals of WT (top row) and mut (bottom row) littermates. (D) Average \( \Delta C_{rel} \) values plotted versus step duration for C2BDN (D1) and C2BKW (D2) calyx terminals. The numbers of SVs, obtained by assuming a single SV capacitance of ~80 aF (Sakaba, 2006), are shown on the right axis.

(E) Relationships between stimulus frequency \( f_{\text{stim}} \) and average replenishment rate constant \( \tilde{r}_4 \) for WT C2BDN (filled circles) and WT C2BKW (empty circles) calyx synapses (E1) and for mut C2BDN (red) and mut C2BKW (blue) calyx synapses (E2). As the two WT datasets are nearly indistinguishable, the gray symbols in E2 represent their average. Axis and tick mark labels in the graphs shown in the bottom rows of (A), (B), and (D) also apply to the graphs of the top rows of the respective panels but were omitted for clarity.

Data depict mean ± SEM; n values and statistical significance are summarized in Table S1. See also Figures S3 and S6.
measured after 2- and 4-s recovery in mutKW synapses were indistinguishable from those measured at 16 s in either mutKW or WT KW synapses, indicating similar $p$ values at the respective recovery intervals (Figure 6B).

To further corroborate that faster recovery from STD in mutKW synapses primarily stems from faster SVR, we assayed SV pool recovery directly by presynaptic $D_{\Delta C_m}$ recordings. The duration of presynaptic depolarizations was limited to 3 ms to limit fusion of slowly releasing SVs (Sakaba and Neher, 2001b). Recovery of $D_{\Delta C_m}$ was noticeably faster in mutKW as compared to WT KW calyx terminals (Figure 6C) and resembled the accelerated eEPSC recovery time course (Figure 6A).

We then tested whether accelerated recovery from STD in mutKW synapses is only observed after high-frequency conditioning (100 and 200 Hz), leading to substantial temporal summation of AP-evoked presynaptic global $[Ca^{2+}]_i$ transients, or if it is also seen following low-frequency conditioning (10 and 20 Hz), during which individual AP-evoked $[Ca^{2+}]_i$ transients decay nearly completely (Müller et al., 2007 and see below). We found that lowering the number of conditioning stimuli together with their frequency considerably slows down eEPSC recovery (Figures S4A3 and S4B3). However, at any tested conditioning frequency, recovery from STD occurred faster in mutKW as compared to WT KW synapses (Figures 6A and S4C). A detailed analysis of the relationship between the estimated steady-state $K_0$, during stimulus trains versus the fractional recovery at different intervals after a conditioning train for all three mutants at the level of individual synapses is presented in Figure S6.

These data show that, in the Munc13-1 C2BKW mutant with enhanced $Ca^{2+}$-dependent phospholipid binding to the C2B domain, SVR is accelerated after low- and high-frequency conditioning. Faster recovery of eEPSC amplitudes is not driven by an elevated release probability during the recovery period.

Augmenting presynaptic AP-evoked $Ca^{2+}$ influx does not mimic the acceleration of SV recruitment caused by the C2BKW mutation

The accelerated eEPSC recovery after high-frequency trains in mutKW synapses is reminiscent of faster recovery from STD upon increased presynaptic $Ca^{2+}$ influx in calyx synapses (Wang and Kaczmarek, 1998). We tested whether widening of calyceal APs by TEA, thus augmenting presynaptic $Ca^{2+}$ influx, induces fast SVR in WT synapses similar to that seen in mutant C2BKW synapses. Bath application of TEA in the presence of kyn enhanced eEPSCs >4-fold and strongly increased STD (Figure 6D). Unexpectedly, though, we observed only a moderate acceleration of the eEPSC recovery time course (Figure 6D2). To resolve the apparent discrepancy between our results and those of Wang and Kaczmarek (1998), we repeated our experiments in the presence of the high-affinity, slowly dissociating AMPAR antagonist NBQX instead of the low-affinity, fast-dissociating antagonist kyn. With 100 nM NBQX, adequate voltage-clamp is ensured by reducing eEPSCs to a size comparable to that observed with kyn in the bath, while eEPSCs are still subject to AMPAR saturation and desensitization (Wadiche and Jahr, 2001). Augmentation of eEPSCs following TEA application was considerably smaller (<3-fold; Figures S7A and S7B) in the presence of NBQX instead of kyn, indicative of postsynaptic AMPAR saturation (Taschenberger et al., 2002, their Figure 8).
(200-Hz conditioning; Figure S7A1), which is unexpected if SVR during steady state was strongly enhanced by TEA.

These data show that, under conditions that limit AMPAR saturation and desensitization, SVR acceleration by enhanced Ca²⁺ influx is substantially smaller than that seen in mutKW synapses. We conclude that the C₂BKW-mutation-induced SVR enhancement is not mimicked solely by augmenting AP-evoked Ca²⁺ influx. Rather, it reflects a combined regulatory effect of Ca²⁺ and phospholipid binding on Munc13-1 activity, whose magnitude is unique.

**AP-evoked [Ca²⁺]i transients in calyx terminals are unaltered in C₂BDN and C₂BKW synapses**

To rule out that changes in the dynamics of presynaptic [Ca²⁺]i due to differences in Ca²⁺ buffering or clearance contribute to the altered eEPSC recovery kinetics in mutDN and mutKW synapses, we measured [Ca²⁺]i transients in response to afferent-fiber stimulation in nearly unperturbed calyx terminals preloaded with the low-affinity Ca²⁺ indicator dye Cal520FF (~1 min; Figures 7A and 7B; Habets and Borst, 2005; Müller et al., 2007). Maximum [Ca²⁺]i amplitudes during trains were on average ~7 (100 Hz) and 16–18 (200 Hz) times larger than those of single AP-evoked transients. As described (Müller et al., 2007), calyceal [Ca²⁺]i transients decayed bi-exponentially with a slow τ in the hundreds of millisecond range, which became more prominent after repetitive AP firing. When superimposing [Ca²⁺]i transients, transients recorded in either mut or WT C₂BDN or C₂BKW terminals, their waveforms were indistinguishable (Figure 7C; Table S1).

Having established the average time course of global [Ca²⁺]i transients induced by conditioning trains, we modeled the recovery from SV pool depletion in C₂BDN and C₂BKW synapses by considering a SVR process that obeys a first-order rate law (Hosoi et al., 2007) and by using the previously established relationship between fₘₚ and Kᵣ (Figure 4E). We assumed that the latter also predicts the relationship between global [Ca²⁺]i and Kᵣ, i.e., we postulated a linear relationship between fₘₚ and [Ca²⁺]i at steady state. Numerical simulations (Figure 7D) capture the essential features of the C₂B-mutation-induced changes, indicating that the altered dynamic regulation of Kᵣ allows us to predict changes in pool recovery time course that correspond to the experimental observations. On the other hand, the simulations do not reproduce the initial “negative” or the “overshooting” eEPSC recovery observed in mutKW (Figure 5A3) and mutBKW (Figure 6A3) synapses, respectively, indicating that a more detailed model, covering the heterogeneity in p as well as changes in p during and after conditioning stimulation, is required to more faithfully reconstruct the eEPSC recovery time course.

The data above—along with the facts that Iᵣ(Δt) amplitudes, VGCC gating kinetics, and Iᵣ(Δt) facilitation time course during AP-like stimulus trains are unaltered—indicate that changes in presynaptic Ca²⁺ signaling do not contribute to altered SVR kinetics in C₂BDN and C₂BKW synapses. Numerical simulations based on measured parameters predict the major features of C₂B-mutation-induced changes in the pool recovery time course.

**Temporal precision of information processing at calyx synapses deteriorates in the absence of fast SV recruitment**

Calyx of Held synapses operate with high temporal precision, even at high transmission rates, leading to reliable AP firing of MNTB PNs (Guinan and Li, 1990; Kopp-Scheinpflug et al., 2008; Lorteije et al., 2009). In addition to other features (Borst and Soria van Hoeve, 2012; Schneggenburger and Forsythe, 2006; von Gersdorff and Borst, 2002), an activity-dependent SVR upregulation may critically contribute to sustained and temporally precise transmission. We thus tested whether the Munc13-1 mutations affect information processing at calyx synapses by analyzing postsynaptic AP timing following afferent fiber stimulation.

Recordings from MNTB PNs in cell-attached configuration represent a minimally invasive form of monitoring postsynaptic Na⁺ spikes with high signal-to-noise ratio, allowing analyses of AP timing with microsecond precision (Figures 6A and 8B; Lorteije et al., 2009). Extracellularly recorded APs measured during the recovery period following conditioning stimulation are shown superimposed in Figures 8A2 and 8B2. In WT synapses, a clear right shift in the AP timing relative to stimulus onset is seen for short recovery intervals, showing that small evoked excitatory postsynaptic potentials (eEPSPs), generated while the FRP is

**Figure 6. C₂BKW synapses recover faster from synaptic depression after high-frequency conditioning**

(A) Similar experiments as in Figure 5A but in C₂BKW synapses. (A1) Traces of 200-Hz trains (left column) and eEPSCaut recorded at different recovery intervals (right column) for a WT (top row) and a mutKW (bottom row) calyx synapse. (A2) Bar graphs and scatter dot plots representing average values and individual synapses, respectively, of the recovered fraction of eEPSCaut amplitude for different recovery intervals. (A3) Average time course of eEPSC recovery from synaptic depression after 100-Hz (top) or 200-Hz (bottom) conditioning trains. (B) Comparison of PPRs during the recovery time course in C₂BKW synapses. (B1) Traces of eEPSCaut pairs (10 ms ISIs) following 100-Hz conditioning trains (left) recorded at recovery intervals of either 2 s (middle) or 16 s (right) in a mutKW synapse. (B2) Bar graphs and scatter dot plots representing average values and individual synapses, respectively, of PPRs measured at 2, 4, and 16 s recovery in mutKW synapses (left bars, blue symbols) in comparison to 16 s recovery in WT synapses (right bar, gray symbols). (C) Recovery from SV pool depletion assayed by presynaptic Jₘp recordings. SV pool depletion was induced by applying step depolarizations of 3 ms duration from Vₘ = −80 mV to 0 mV, and recovery was monitored at intervals ranging from 0.125 s to 16 s after the end of the depolarization. (C1) Sample Jₘp traces for control and 0.25-s, 2-s, and 16-s recovery intervals recorded in a WT (top) and a mutKW (bottom) calyx terminal are shown. For clarity, Jₘp traces in this panel were low-pass filtered with a moving average kernel of width 7. (C2) Average recovery time course of ΔCₘp measured in WT (gray) and a mutKW (blue) calyx synapse. (D) (D1) Similar experiment as illustrated in (A) but recorded in a WT synapse in the absence (top row) and presence (bottom row) of 1 mM TEA to broaden presynaptic APs and enhance presynaptic Ca²⁺ influx. All recordings were obtained in the presence of 1 mM kyn. (D2) Average time course of eEPSC recovery after 100- (top) or 200-Hz (bottom) conditioning trains recorded in the absence (gray symbols) or presence (black symbols) of 1 mM TEA. Enhancing presynaptic Ca²⁺ influx accelerated recovery of eEPSCaut but to a much lower extent than in C₂BKW synapses (compare A3 and D2). Data depict mean ± SEM; n values and statistical significance are summarized in Table S1. See also Figures S4, S6, and S7.
Figure 7. AP-evoked global volume-averaged \([\text{Ca}^{2+}]_i\) transients in calyx terminals

(A) Fluorescence images of a calyx terminal pre-loaded with the low-affinity Ca\(^{2+}\) indicator dye Cal520FF (K_D = 9.8 \(\mu\)M) during a brief whole-cell episode (~1 min; pipette concentration 400 \(\mu\)M; final cytosolic concentration ~200 \(\mu\)M) at rest (top) and during 200-Hz stimulation (50 APs, center). The difference image is shown in the bottom panel.

(B) Presynaptic \([\text{Ca}^{2+}]_i\) transients \((\Delta F/F_0)\) elicited by a single AP (top) or trains of 25 (100 Hz, center) or 50 APs (200 Hz, bottom), corresponding to stimulus trains used for Figures 5A, 6A, and 6D. Each trace represents the average of 6 (single AP) or 4 (AP trains) repetitions, and the gray areas represent ±SEM. AP discharge pattern recorded in response to afferent fiber stimulation under current-clamp conditions during the preloading period is illustrated in the top right insets. Red traces represent double-exponential fits to the decay of the \([\text{Ca}^{2+}]_i\) transients. Fast- and slow-decay time constants are given next to the \([\text{Ca}^{2+}]_i\) transients. The single AP response is shown superimposed to the train responses in the center and bottom panels for comparison.

(C) Normalized average presynaptic \([\text{Ca}^{2+}]_i\) transients recorded in WT (back) and mut (red and blue) C2BDN (C1) and C2BK (C2) terminals in response to a single AP (left), a 100-Hz AP train (25 APs, center), or a 200-Hz AP train (50 APs, right).

(D) Numerical simulations of SV pool recovery after 100-Hz (left) and 200-Hz (right) trains assuming first-order kinetics and a \([\text{Ca}^{2+}]_i\)-dependent \(\tau_+\). For simplicity, we approximated the relationship between \(\tau_+\) and \([\text{Ca}^{2+}]_i\) by assuming a linear relationship between \(t_{\text{stim}}\) and \([\text{Ca}^{2+}]_i\) at steady state. The decay of \(\tau_+\) immediately after conditioning trains (corresponding to \(t = 0\) s in D) was modeled with a double-exponential function using decay time constants established by fitting the average \([\text{Ca}^{2+}]_i\) transients shown in (C).
largely depleted, mostly trigger postsynaptic APs with longer latencies. Strikingly, mutDN synapses, which lack the fast component of eEPSC recovery, tended to show even longer latencies between stimulus and postsynaptic spike for short recovery intervals.

Temporal jitter and timing of postsynaptic APs was further analyzed in whole-cell configuration to obtain stable long-term recordings and to collect more repetitions of a given protocol (Figures 8C and 8D). In rare cases, individual aberrant postsynaptic APs were triggered during conditioning, presumably by asynchronous release events generating supra-threshold eEPSPs, so that the number of postsynaptic APs exceeded the number of stimuli in some trials (Figure 8C1). As in cell-attached recordings, we observed longer AP latencies at short recovery intervals in mutDN as compared to WTDN synapses. In addition, we found that the temporal jitter increased (Figure 8D2). Conversely, mutKW synapses showed improved temporal precision as compared to WT synapses, so that AP latencies were shorter and spike latency jitter was reduced for a range of short recovery intervals (0.5–4 s; Figure 8D3). Figures 8E and 8F plot the average increase in spike timing relative to those measured for the longest recovery interval (16 s) and the mean of the standard deviation of the timing of AP peaks as a measure of spike timing variability. In both WT and mutKW synapses, AP latencies were ~200 μs longer for the shortest recovery interval (0.125 s) than those after 16 s recovery. In mutDN synapses, the spike latency increase for short recovery intervals was nearly twice as large and remained larger for all but the 4-s and 8-s intervals. In mutKW synapses, on the other hand, AP latencies declined much faster with increasing recovery intervals. For 2–16 s recovery, no differences in spike timing were measurable, consistent with the fact that, for recovery intervals, >2 s eEPSCs had nearly fully recovered after conditioning 100-Hz stimulation (Figure 8A3). A similar picture emerged for the temporal jitter of AP latencies. For intervals ≤1 s, the jitter was larger in mutDN but generally smaller in mutKW synapses as compared to WT synapses (Figure 8F).

These data show that the modulation of SVR speed mediated by Ca²⁺-phospholipid binding to the Munc13-1 C₂B domain influences synapse fidelity. This phenomenon is expected to alter synaptic computation at the level of single synapses and circuits.

**Faster rebound from synaptic depression at hippocampal synapses upon acceleration of SV recruitment**

To show that the rate of Munc13-1-dependent SVR also determines STP at synapses that are less prone to rapid SV pool exhaustion at their typical firing rates (Mizuseki and Buzsáki, 2013), we analyzed transmission at P16–P21 hippocampal Schaffer collateral/commisural synapses with CA1 neurons (SC/C-CA1), focusing on C₂BKW mice (Figure 8G). We conditioned SC/C-CA1 synapses with 10-Hz trains. This stimulus frequency was chosen to limit the contribution of synaptic augmentation during recovery from STD and to avoid posttetanic potentiation. We observed a slight initial paired-pulse facilitation (1.09 ± 0.05 and 1.05 ± 0.03 in WT KW and mutKW synapses, respectively), followed by depression, during which steady-state eEPSC amplitudes were reduced to ~55% as compared to the maximum eEPSC size measured during trains, indicating an RRP depletion of ≥45%. Following conditioning, eEPSCs recovered within 5–10 s to an amplitude that was ~20% larger than eEPSCt of the conditioning train (Garcia-Perez and Wesseling, 2008). While levels and decay time constants (~9 s) of synaptic augmentation were similar in WT KW and mutKW synapses, the rebound from STD was accelerated in mutKW synapses, particularly at 1- to 2-s recovery intervals (Figures 8H1 and 8H2). To corroborate a more complete SV pool recovery after 2 s recovery from STD, we approximated in a subset of synapses the relative RRP occupancy by estimating the ratio of $RRP/RPP_{0.1}$, assuming that pool recovery has completed after 16 s recovery. The corresponding ratio was larger in mutKW (96% ± 3%) as compared to WT KW (82% ± 5%) SC/C-CA1 synapses (Figure 8H3).

These data show that SVR acceleration by Munc13-1 affects short-term plasticity also at strongly augmenting synapses, indicating a fascinating interplay between different short-term plastic processes that shapes the distinct features of different synapse types.

**DISCUSSION**

Munc13s are essential SV priming proteins, in whose absence no fusion-ready SVs are formed and synaptic transmission ceases (Sigler et al., 2017; Varoquaux et al., 2002). Studies with cultured neurons showed that Munc13 activity can be regulated by Ca²⁺-CaM binding to an amphipathic helix motif (Junge et al., 2004; Lipstein et al., 2012), by DAG binding to a central C₁ domain (Betz et al., 1998; Rhee et al., 2002), and by Ca²⁺-phospholipid binding to an adjacent C₂B domain (Shin et al., 2010). What has remained unknown is how these regulatory processes affect synapse function downstream of SV priming and whether they are used purposefully in intact networks to shape computational synapse properties. We show here that Munc13-1 is regulated by convergent Ca²⁺- and phospholipid-dependent signaling to adjust SV priming rates, enable sustained transmission, and ensure temporal fidelity of synaptic signaling in intact circuits.

To assess the role of Ca²⁺- and phospholipid-dependent regulation of Munc13-1 function in intact circuits, we generated KI mice that express Munc13-1 variants with abolished (Munc13-1D706/711H) or increased Ca²⁺-dependent phospholipid binding to the C₂B domain (Munc13-1K706W; Shin et al., 2010) and characterized functional consequences in the calyx of Held synapse. Our data demonstrate an activity-dependent activation of Munc13-1 via its C₂B domain. In the absence of Ca²⁺-regulated phospholipid binding to the Munc13-1 C₂B domain, fast SVR during and immediately after high-frequency AP trains is strongly attenuated, resulting in reduced transmitter release and perturbed temporal precision of synaptic transmission following bursts of activity. Conversely, increasing the Ca²⁺-dependent PIP₂ affinity of the Munc13-1 C₂B domain enhances SVR, augments synaptic strength during and following presynaptic AP trains, and improves temporal precision of transmission. The fact that the mutation-induced bidirectional manipulation of Ca²⁺-phospholipid binding to the Munc13-1 C₂B domain leads to corresponding bidirectional changes in SVR rates demonstrates the existence of a Ca²⁺-phospholipid-sensing regulatory process in SV priming within intact
Figure 8. Munc13-1-dependent SV priming affects the AP timing in MNTB principal neurons during recovery from synaptic depression and accelerates the rebound from synaptic depression at hippocampal Schaffer collateral-CA1 synapses

(A–F) Recordings in brainstem slices obtained in the absence of kyn.

(A and B) Cell-attached recordings of postsynaptic APs triggered in MNTB PNs in response to afferent fiber stimulation during a 100-Hz (25 stimuli) train (A1 and B1) and at various recovery intervals (A2 and B2) in a WTDN (A) and a mut DN (B) synapse. All eight recovery intervals (0.125, 0.25, 0.5, 1, 2, 4, 8, and 16 s) were tested in a single sweep to shorten the duration of the recording protocol. APs recorded during recovery from synaptic depression are shown superimposed after alignment to the stimulus onset. Traces shown in B2 are temporally slightly offset to align the negative peak of the AP recorded after 16 s recovery (dotted line) in the mutDN with that of the corresponding WT DN synapse to facilitate latency fluctuation comparison.

(C) Similar experiment as in (A) and (B), but postsynaptic APs were recorded intracellularly under current-clamp conditions during the 100-Hz trains (left) and for various test intervals (superimposed, right).

(D) Latency fluctuations for APs recorded 0.25 s (left) and 16 s (right) after conditioning 100-Hz trains. 21 consecutive trials are shown superimposed.

(E and F) Summary data for latencies of intracellularly recorded postsynaptic APs (E) and fluctuations of AP timing (F) plotted versus recovery interval. Data from WT synapses of both KI lines were similar and therefore pooled. AP latencies are expressed relative to the timing of the AP peak for 16-s recovery interval ($t_{AP16\,\text{s}}$) (E). Changes in latency fluctuations are expressed as the ratio of the SD of $t_{AP}$ relative to the SD of $t_{AP16\,\text{s}}$ (F).

(G and H) Recordings in P16–P21 hippocampal slices, obtained using 2 mM Ca$^{2+}$ and 1 mM Mg$^{2+}$ in the bath solution and in the absence of kyn.

(G) Recovery from STD induced by 10-Hz stimulus trains (20 APs) at hippocampal SC/C-CA1 synapses of a C$\nu$B WT$^{KW}$ (G1) and a C$\nu$B mut$^{KW}$ (G2) mouse (left panels). Recovery was probed by recording a single eEPSC$\text{cond}$ evoked at increasingly longer recovery intervals (0.125, 0.25, 0.5, 1, 2, 4, 8, and 16 s), which are shown superimposed (right panels).

(H) Average time course of eEPSC amplitudes during conditioning 10-Hz stimulation and the subsequent recovery from STD, obtained from WT$^{KW}$ (H1) and mut$^{KW}$ (H2) SC/C-CA1 synapses. At least three trials were averaged for each synapse tested. Smooth traces represent a model of synaptic plasticity fitted to the data. The model consisted of an SV pool depletion and a stimulus-induced augmentation of the release probability, which slowly decayed in the absence of AP firing. Estimated SV pool recovery time constants $\tau_{R}$ were 2.40 s (WT$^{KW}$) and 1.76 s (mut$^{KW}$). Assuming that pool recovery has completed after 16 s, the relative pool occupancy at 2 s recovery was estimated as the ratio of $R_{PP2\,\text{s}}$/$R_{PP16\,\text{s}}$ for a subset of WT$^{KW}$ and mut$^{KW}$ synapses (H3).

Data depict mean ± SEM.
circuits and identifies Munc13-1 as its major target. We propose that Munc13-1 integrates Ca\(^{2+}\) and PIP\(_2\) signaling to tune SVR speed according to the requirements imposed on the release machinery. We demonstrate that this regulatory process is relevant in calyx of Held and Schaffer collateral/commissural-CA1 synapses. Given that Munc13-1 is expressed in essentially all neurons of the brain (Augustin et al., 1999a), this regulatory principle is likely to operate in many brain circuits to control information processing.

The Munc13 C2B domain
Five aspartic acid residues of the Munc13 C2B domain coordinate two Ca\(^{2+}\) ions (Shin et al., 2010). Binding assays with the isolated C2B domain indicate an apparent Ca\(^{2+}\) EC\(_{50}\) of 5.5 \(\mu\)M to promote phospholipid binding. Such high [Ca\(^{2+}\)]\(_{i}\) is likely only reached near open VGCCs (Eggermann et al., 2011; Neher and Sakaba, 2008) or during high-frequency AP firing (Korogod et al., 2005; Lin et al., 2017). Our findings, showing that Ca\(^{2+}\)-dependent phospholipid binding to the C2B domain promotes SVR, imply that the C2B domain of Munc13s “sees” high [Ca\(^{2+}\)]\(_{i}\) levels during AP firing.

The lipid binding properties of the Munc13-1 C2B are rather unique, with PIP and PIP\(_2\) binding preferentially and equally well. This may allow the domain to respond to subtle temporal and spatial changes in phosphoinositide (PI) second messenger levels, which occur during ongoing presynaptic activity, e.g., via Ca\(^{2+}\)-dependent regulation of phospholipases or PI kinases/phosphatases or upon activation of cell-surface receptors (Brown and Sihra, 2008). These considerations, and the fact that Munc13-1 acts as an AZ organizer (Sakamoto et al., 2016), lead to the notion of a dynamic interplay between the lipid and protein composition of SV fusion sites that is regulated by synaptic activity.

While the RRP is completely eliminated upon Munc13 loss (Siksou et al., 2009; Varoqueaux et al., 2002), several mutations of Munc13-1 regulatory domains reduce RRP size. This was observed in KI neurons expressing a DAG-insensitive Munc13-1 (Rhee et al., 2002) and subsequently with C2A and C2C deletion mutant Munc13-1 variants. The latter finding led to the notion that Munc13-1 may not only regulate SNARE complexes but also bridge SV and AZ plasma membranes, involving the C1- C2B tandem (Liu et al., 2016; Quade et al., 2019). The present study did not yield evidence for RRP changes upon elimination or enhancement of C2B-lipid interactions, indicating that the C2B mutations we introduced do not interfere with any membrane bridging function.

Certain Munc13 C2B mutations change presynaptic [Ca\(^{2+}\)]\(_{i}\) transients in cultured neurons, presumably via VGCC modulation (Calloway et al., 2015). We observed no defects in calyceal I\(_{Ca}\) of C2BDN and C2BKW mutants. Likewise, AP-evoked global [Ca\(^{2+}\)]\(_{i}\) transients in dye-preloaded calyces were unaltered, indicating that the major spatial aspects of AZ organization and AP-evoked presynaptic Ca\(^{2+}\) influx are intact in mut\(_{DN}\) and mut\(_{KW}\) synapses.

Phospholipid-dependent regulation of SV priming and fusion
Phospholipids are presumed to regulate Ca\(^{2+}\)-dependent vesicle fusion in neurons and neuroendocrine cells. Many presynaptic proteins have lipid-binding domains (Pinheiro et al., 2016), and PIP\(_2\) is a major component of the plasma membrane at release sites (Kabachinski et al., 2014; Milosevic et al., 2005; Walter et al., 2017). PIP\(_2\) likely clusters syntaxin (Honigmann et al., 2013; van den Bogaart et al., 2011) and other AZ components (de Jong et al., 2018), acts as a regulator of synaptotagmin-driven SV fusion (van den Bogaart et al., 2012), and is involved in presynaptic clathrin-mediated endocytosis (Jung and Haucke, 2007), also in the calyx of Held (Eguchi et al., 2012).

A major challenge in studying phospholipid-dependent regulation of synaptic protein function is that the dynamic lipid composition at AZs is unknown and difficult to manipulate experimentally. Most relevant data were obtained with in vitro analyses of neuroendocrine cells, where vesicle fusion occurs over the entire plasma membrane, and by pharmacologically manipulating phospholipid composition. The time course of such manipulations is slow, and they often lack specificity. Fast-acting, light-activated lipid compounds exist, but their activation is difficult to restrict to AZs (Frank et al., 2016; Walter et al., 2017). To circumvent these challenges, we chose a genetic approach, i.e., the mutagenesis of a single lipid-sensor protein. Interfering with the Ca\(^{2+}\)-phospholipid-dependent regulation of Munc13-1 strongly affects SVR speed. As pharmacologically enhancing AP-evoked Ca\(^{2+}\) influx alone only moderately speeds up SVR, we propose that AZ lipid composition modulates Munc13 activity to regulate SVR.

Phospholipid-binding properties of C2BKW resemble those of the synaptotagmin-1 C2A domain, with preference for PIP and largely unaltered Ca\(^{2+}\) sensitivity (Shin et al., 2010). In mut\(_{KW}\) synapses, initial synaptic strength, PPR, and the rate of eEPSC depression during the onset of high-frequency trains are unaltered, indicating that \(\tilde{\beta}\) is unchanged. However, for all but the three highest stimulus frequencies (50–200 Hz), we determined increased steady-state replenishment rate constants \(\tilde{R}\). Elevated PIP\(_2\) affinity of the mutated Munc13-1 C2BKW domain thus augments SVR. During high-frequency stimulation, the elevated PIP\(_2\) affinity of the mutated C2BKW domain may become less important because elevated [Ca\(^{2+}\)]\(_{i}\) enhances phospholipid binding to the C2B domain also in WT synapses. Indeed, for high-frequency stimulation, enhancing AP-evoked presynaptic Ca\(^{2+}\) influx pharmacologically only moderately speeds up eEPSC recovery, indicating that \(\tilde{R}\), during 100- and 200-Hz conditioning is close to maximum even in WT synapses.

The C2BDN mutation perturbs Munc13-1 function by replacing two aspartic acid residues, abolishing Ca\(^{2+}\) binding (Shin et al., 2010). We observed higher initial synaptic strength in mut\(_{DN}\) synapses as compared to WT. This is likely caused by higher \(\tilde{R}\), as PPRs are lower in Munc13-1 C2BDN synapses, and synaptic depression during high-frequency trains proceeds faster while RRP is unaffected. An explanation for this unexpected finding is that the C2BDN variant shows significant membrane binding in the absence of Ca\(^{2+}\) (Shin et al., 2010), which may lead to increased Munc13-1 C2BDN activity at resting [Ca\(^{2+}\)].

The interplay of regulatory domains of Munc13s
Three Ca\(^{2+}\)- and second-messenger-sensitive regulatory domains allow Munc13s to respond to a wide range of [Ca\(^{2+}\)], and lipid second messengers. Ca\(^{2+}\)-CaM binding to Munc13s occurs
at sub-micromolar [Ca\textsuperscript{2+}]. (Dimova et al., 2006; Zikich et al., 2008), while the C\textsubscript{2}B domain requires 2–10 μM [Ca\textsuperscript{2+}] (Shin et al., 2010). Thus, Ca\textsuperscript{2+}-CaM-dependent regulation of Munc13s can operate at lower [Ca\textsuperscript{2+}], than C\textsubscript{2}B domain activation. As Munc13-1 is localized at AZs near VGCCs, it is possible that the C\textsubscript{2}B domain preferentially senses brief local [Ca\textsuperscript{2+}] transients in addition to large global [Ca\textsuperscript{2+}] changes during prolonged high-frequency firing, while the high Ca\textsuperscript{2+} affinity of the Ca\textsuperscript{2+}-CaM pathway may allow efficient sensing of longer lasting and lower global [Ca\textsuperscript{2+}] transients. DAG binding to the C1 domain allows Munc13 activation at timescales that exceed the duration of [Ca\textsuperscript{2+}] transients as its termination depends on enzymatic DAG removal. It is therefore likely that the regulatory domains of Munc13s operate in a successive, cooperative, and synergistic fashion rather than redundantly.

We previously examined functional effects of a point mutation (Munc13-1\textsuperscript{W464R}; mut\textsubscript{WR}) that renders Munc13-1 Ca\textsuperscript{2+}-CaM insensitive (Lipstein et al., 2013). Mut\textsubscript{WR} calyces show much slower SV pool recovery following long presynaptic depolarizations, but recovery of AP-evoked eEPSCs following STD is less perturbed. The latter finding led to the conclusion that additional, possibly more effective, molecular pathways for [Ca\textsuperscript{2+}]-dependent SVR regulation must exist (Lipstein et al., 2013). The present study identifies Ca\textsuperscript{2+}-dependent phospholipid binding to Munc13-1 as such a pathway. The loss of fast eEPSC recovery in mut\textsubscript{DN} synapses demonstrates that it plays a major role in Ca\textsuperscript{2+}-dependent SVR. This mechanism likely operates in concert with Ca\textsuperscript{2+}-CaM and DAG binding to adjust presynaptic function on demand, stressing the notion of Munc13s as a regulatory hub in SV priming.

**The function of the Munc13 C2B domain—consensus and discrepancies**

Multiple previous studies focused on Munc13 C\textsubscript{2}B function, establishing it as a Ca\textsuperscript{2+}-phospholipid binding domain that controls Munc13 priming function and thus dynamically regulates transmitter release (Kabachinski et al., 2014; Michelassi et al., 2017; Shin et al., 2010). Beyond this, Munc13 C\textsubscript{2}B interactions with AZ scaffold proteins (Brockmann et al., 2020) and VGCCs (Calloway et al., 2015) were reported. Unfortunately, a direct comparison of previous studies with ours is complex as different paralogs, mutations, neuronal preparations, synapse types, or organisms were studied.

Nevertheless, at least three general commonalities emerge. First, effective and specific blockade of Ca\textsuperscript{2+}-phospholipid binding to Munc13 C\textsubscript{2}B reduces SV priming activity and attenuates synaptic transmission, albeit to different degrees under different stimulation conditions (Kabachinski et al., 2014; Michelassi et al., 2017; Shin et al., 2010). Second, certain mutations in the Ca\textsuperscript{2+}-phospholipid-binding pocket of Munc13 C\textsubscript{2}B can cause increased transmitter release, likely due to an increase in Ca\textsuperscript{2+}-independent phospholipid binding (Kabachinski et al., 2014; Michelassi et al., 2017; Shin et al., 2010). Third, Ca\textsuperscript{2+}-phospholipid-dependent regulation via the C\textsubscript{2}B domain can modulate the dynamics of synaptic transmission upon prolonged high-frequency stimulation (Shin et al., 2010; present study), with different consequences for short-term plasticity. The remaining differences in the context of these commonalities likely arise due to differences in the neuronal membrane lipid composition present in the various preparations and synapse types studied.

Equally important are the facets that distinguish the present findings from previously published ones. Most importantly, the conclusions drawn from our work versus that of Shin et al. (2010) differ fundamentally. Whereas Shin et al. (2010) did not report evidence for a regulation of SVR rate by Munc13 C2B, we provide clear evidence for such a regulation. Aspects of this discrepancy might be due to different experimental conditions. This notwithstanding, our findings demonstrate that the presynaptic process of dynamic SV priming must ultimately be studied in intact circuits as they appear in vivo to determine its exact molecular features and computational role in circuits.

**Synaptic depression and recovery at the calyx of Held synapse**

Several mechanisms, including pool depletion, negative feedback via presynaptic metabotropic receptors, and postsynaptic receptor desensitization, contribute to synaptic depression (Zucker and Regehr, 2002). Synaptic transmission at post-hearing onset calyx synapses shows only limited sensitivity to auto-inhibition via mGluRs (Renden et al., 2005) or AMPARs desensitization and saturation (Taschenberger et al., 2002). Because we recorded eEPSCs in the presence of 1 mM kyn, further attenuating possible postsynaptic effects, we assume pool depletion as the principal cause for synaptic depression in our experiments.

STD is determined by a balance of SV recruitment and consumption. Recovery from synaptic depression is slow (τ = 4 s) at calyx synapses when measured after stimuli that do not strongly increase global [Ca\textsuperscript{2+}]. (Iwasaki and Takahashi, 2001; von Gersdorff et al., 1997). Considering θ = 1/τ = 0.25 s\textsuperscript{-1} and a total number of SV docking sites (N) that approximately equals the size of the FRP (1,000–2,000 SVs), the SVR rate would be limited to 250–500 SV/s. For 200-Hz trains and q ∼ 60 pA (Chang et al., 2015), this equates to an eEPSC\textsubscript{ss} of only 75–150 pA (∼8–17 pA for recordings in 1 mM kyn), which is much less than experimentally observed. In fact, we estimated a 16-times-higher θ\textsubscript{d} of ∼4 s\textsuperscript{-1} to predict the observed steady-state release rates during 200-Hz trains. Our θ\textsubscript{d} estimate rests on the assumption that N is approximately equal to the FRP size, i.e., that the occupancy (ρ\textsubscript{occ}) of docking sites in resting calyces is close to 100%. If ρ\textsubscript{occ} were substantially lower (Malagon et al., 2020), we would need to postulate a larger N and, consequently, lower θ\textsubscript{d}.

Assuming a strongly upregulated θ\textsubscript{d} during high-frequency conditioning, which then decays to its basal value after cessation of stimulation, we expect a biphasic eEPSC recovery time course. During low-frequency conditioning, θ\textsubscript{d} increases much less, which explains the absence of a fast component of eEPSC recovery. However, the mere existence of a biphasic eEPSC recovery alone is insufficient proof for a Ca\textsuperscript{2+} dependence of SVR (Hallemand et al., 2010), and for some synapses, experimental interference with either [Ca\textsuperscript{2+}] or its sensor binding did not indicate a Ca\textsuperscript{2+} dependence of SVR (Miyano et al., 2019; Ritzau-Jost et al., 2018). Nevertheless, the findings that, in calyx synapses, eEPSC recovery is accelerated upon augmenting presynaptic Ca\textsuperscript{2+} influx (Wang and Kaczmarek, 1996) and sensitive to
manipulations that interfere with Ca\(^{2+}\) binding to CaM (Sakaba and Neher, 2001a), with Ca\(^{2+}\)-CaM binding to Munc13-1 (Lippeinstein et al., 2013), or with Ca\(^{2+}\)-phospholipid binding to Munc13-1 (present study) support the notion that the fast component of pool recovery is regulated by elevated [Ca\(^{2+}\)].

For 200-Hz stimulation of WT synapses, we estimated a \(\bar{R}\), of \(-4\) pools/s at steady state. Assuming a linear relationship between global [Ca\(^{2+}\)], and SVR rate constant according to \(\bar{R} = k_{\text{basal}} + \alpha \times [\text{Ca}^{2+}]\), with \(k_{\text{basal}} = 0.1\) s\(^{-1}\) and a slope factor \(\alpha\) of \(-1\) pool/(μM \times s) (Hosoi et al., 2007), this corresponds to [Ca\(^{2+}\)] of \(-3.9\) μM. Such supra-micromolar global [Ca\(^{2+}\)] values of \(-1.4\) μM and in the range of 3–10 μM were previously reported for voltage-clamped WT calyces stimulated with 200-Hz AP waveform trains (Lin et al., 2017) and for dye-preloaded WT calyces during prolonged 100-Hz firing (Korogod et al., 2005), respectively.

**Munc13-1-controlled SV recruitment and information processing in single synapses and synaptic circuits**

Our analyses of postsynaptic AP latency and jitter show that altered SVR can affect information transfer at calyx synapses. Consistent with the “failsafe” operation of calyx synapses (Guinan and Li, 1990; Lorteije et al., 2009), we never observed spike failures during or following AP trains. In comparison to WT, mut\(_{\text{ON}}\) synapses show longer AP latencies and reduced temporal precision of transmission following conditioning trains. In contrast, mut\(_{\text{KW}}\) synapses show improved temporal precision and shorter AP latencies. This demonstrates that the SVR speed determines postsynaptic AP timing after bouts of presynaptic activity. This may have only subtle functional consequences at the calyx of Held, particularly with faster SVR at physiological temperature (Kushmerick et al., 2006). However, profound effects on the reliability of postsynaptic firing are likely at the many less failsafe synapses in the brain or in scenarios where neurons summate multiple synaptic inputs for triggering AP firing. In the corresponding circuits, the dynamic regulation of Munc13-1 activity by Ca\(^{2+}\) and phospholipids is expected to have substantial effects on information processing.

Our experiments on SC/C-CA1 synapses show that an acceleration of SVR speed also affects STP at small hippocampal synapses. In this case, accelerated recovery from STD in mut\(_{\text{KW}}\) synapses operates in parallel to strong augmentation of release probability, which is typical for these synapses. These data indicate that dynamically regulated SV priming by Munc13-13s likely interfaces with other presynaptic processes and STP phenomena to shape the unique characteristics of different synapse types throughout the brain.

**STAR+METHODS**

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**SUPPLEMENTAL INFORMATION**

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**AUTHOR CONTRIBUTIONS**

N.B. initialized the project. N.L. generated and validated Munc13-1 C\(_2\)B KIs. N.L., S.C., and H.T. designed and performed postsynaptic recordings in brainstem slices. K.-H.L. designed and performed presynaptic recordings and Ca\(^{2+}\) imaging in brainstem slices, with assistance of H.T. F.J.L.-M. designed and performed recordings in hippocampal slices and Munc13-1 immunolabeling analyses of brainstem sections, with assistance of H.T. N.L., S.C., K.-H.L., F.J.L.-M., and H.T. analyzed and interpreted data. H.T. supplied software routines and performed simulations. N.B. and E.N. provided conceptual input and advice. N.L., H.T., and N.B. wrote the manuscript. All authors provided text edits.

**DECLARATION OF INTERESTS**

The authors declare no competing interests. N.B. is a member of the Neuron advisory board.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-Bassoon antibody | Enzo Life Sciences | Cat# SAP7F407; RRID: AB_2313990 |
| Anti-Calmodulin antibody | Upstate Biotechnology | Cat# 05-173; RRID: AB_309644 |
| Anti-CAPS 1 antibody | Synaptic Systems | Cat# 262 013; RRID: AB_2619979 |
| Anti-Doc 2b antibody | Synaptic Systems | Cat# 174 103; RRID: AB_2619874 |
| MAP2 Antibody       | Novus  | Cat# NB300-213; RRID: AB_2138178 |
| Anti-Munc13-1, rabbit polyclonal | Generated in house | 41 |
| Anti-Munc13-1, mouse monoclonal | Generated in house | 3HS; Betz et al., 1998 |
| Anti-ubMunc13-2, rabbit polyclonal | Generated in house | 48 |
| Anti-bMunc13-2, rabbit polyclonal | Generated in house | 50 |
| Anti-Munc13-3, rabbit polyclonal | Generated in house | 52 |
| Anti-Munc13-1 antibody | Synaptic Systems | Cat# 126 103; RRID: AB_887733 |
| Anti-Munc18-1 antibody | Synaptic Systems | Cat# 116 002; RRID: AB_887736 |
| Anti-Rim1, rabbit polyclonal | Generated in house | Q703 |
| Anti-SNAP 25 antibody | Synaptic Systems | Cat# 111 011; RRID: AB_887794 |
| Anti-Synapsin 1 antibody | Synaptic Systems | Cat# 106 011; RRID: AB_2619772 |
| Anti-Synaptobrevin 2 antibody | Synaptic Systems | Cat# 104 211; RRID: AB_887811 |
| Anti-Synaptotagmin 1 antibody | Synaptic Systems | Cat# 105 001; RRID: AB_887831 |
| Anti-Syntaxin 1 antibody | Synaptic Systems | Cat# 110 011; RRID: AB_887844 |
| Peroxidase-AffiniPure Goat Anti-Mouse IgG (H+L) antibody | Jackson ImmunoResearch Labs | Cat# 115-035-146; RRID: AB_2307392 |
| Peroxidase-AffiniPure Goat Anti-Rabbit IgG (H+L) antibody | Jackson ImmunoResearch Labs | Cat# 111-035-144; RRID: AB_2307391 |
| Goat anti-rabbit IgG Secondary Antibody, Alexa Fluor 488 | Thermo Fisher Scientific | Cat# A-11008; RRID: AB_143165 |
| Goat anti-mouse IgG Secondary Antibody, Alexa 555 | Thermo Fisher Scientific | Cat# A21424; RRID: AB_141780 |
| Goat anti-Chicken Secondary Antibody, Alexa Fluor 633 | Invitrogen | Cat# A-21103; RRID: AB_2535756 |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| 2,3-Dioxo-6-nitro-1,2,3,4-tetrahydrobenzo[f]quinoxaline-7-sulfonamide disodium salt (NBQX) | HelloBio | Cat# HB0443 |
| Goat Serum | GIBCO | Cat# 16210-072 |
| Kynurenic acid | Sigma-Aldrich | Cat# K3375 |
| MemCode™ Reversible Protein Stain Kit | Thermo Scientific | Cat# 24580 |
| Mounting glue: Aqua-Poly/Mount | Polysciences | Cat# 18606-20 |
| Paraformaldehyde (PFA) | Serva | Cat# 31628.02 |
| Phorbol-12,13-dibutyrate (PDBu) | Calbiochem | Cat# 524390 |
| Recombinant Protein G - Sepharose 4B | Invitrogen | Cat# 101242 |
| Strychnine hydrochloride | Tocris Bioscience | Cat# 2785 |
| Tetraethylammonium chloride (TEA) | Sigma Aldrich | Cat# T-2265 |
| Triton X-100 | Roche | Cat# 10789704001 |
| **Experimental models: Cell lines** |        |            |
| Mouse: 129/ola embryonic stem cell line E14 | Hooper et al., 1987 | N/A |

(Continued on next page)
### RESOURCE AVAILABILITY

**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Nils Brose (Brose@em.mpg.de).

**Materials availability**

Mouse lines will be shared upon request within the limits of the respective material transfer agreements.

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| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental models: Organisms/strains** | | |
| Mouse: Unc13a\textsuperscript{tm5Bros} | This manuscript | C2BDN |
| Mouse: Unc13a\textsuperscript{tm6Bros} | This manuscript | C2BKW |
| Mouse: Unc13a\textsuperscript{tm4.1Bros} | Generated in house | Lipstein et al., 2013 |
| Mouse: C57BL/6NCrL | Charles River Laboratories | RRID:IMSR_CRL:027 |
| Mouse Gt(Rosa)26Sor\textsuperscript{tm1(FLP1)Dym} | Breeding in house | Farley et al., 2000 |
| **Oligonucleotides** | | |
| Sense prime: 5'-GGGTAGCTGCAGGATTTATTGTAT-3' | Generated in house | 25254, UNC13A gene intron 18 |
| Antisense prime: 5'-TGTGTCCAGTTGAGGGTC-3' | Generated in house | 35776, UNC13A gene intron 19 |
| **Recombinant DNA** | | |
| Plasmid: pTKPuroFRT-UNC13A Exons 13-26 Exon 18\textsuperscript{C2BDN} | Cloned in house | N/A |
| Plasmid: pTKPuroFRT-UNC13A Exons 13-26 C2BK\textsuperscript{C2BKW} | Cloned in house | N/A |
| **Software and algorithms** | | |
| GraphPad Prism 8 | GraphPad Software | https://www.graphpad.com; RRID: SCR_002798 |
| IgorPro 6.3.7.2 | Wavemetrics | https://www.wavemetrics.com; RRID: SCR_000325 |
| ImageJ | National Institutes of Health | https://imagej.nih.gov/ij; RRID: SCR_003070 |
| Microsoft Excel | Microsoft | https://www.microsoft.com/en-us/; RRID: SCR_016137 |
| PatchMaster v2x53/ Pulse v8.80 | HEKA / Harvard Bioscience | https://www.heka.com; RRID: SCR_000034 |
| Python based Relational Animal Tracking | Scionics Computer Innovation GmbH | https://www.scionics.com/pyrat.html; RRID:SCR_021014 |
| R: A Language and Environment for Statistical Computing | R Core Team | http://www.r-project.org/; RRID:SCR_001905 |
| **Other** | | |
| Beckman Ultracentrifuge L-70 | Beckman | N/A |
| Borosilicate glass with filament | Science Products | GB150F-8P |
| Cryostat | Leica | RRID:SCR_016844 |
| EPC 10 double patch clamp amplifier | HEKA Elektronik | RRID:SCR_018399 |
| Leica TCS SP5 II microscope | Leica | RRID:SCR_018714 |
| Vibratome VT1000S Leica | Leica | RRID:SCR_016495 |
| Olympus BX51 microscope | Olympus | RRID:SCR_018949 |
**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Mouse Generation**

The Munc13-1 knock-in (KI) mice were generated by homologous recombination in 129/ola embryonic stem cells (Hooper et al., 1987). A targeting vector containing UNC13A exons 13-26 with an insertion of an FRT-Puromycin-FRT cassette before exon 18 (Figure 1A) was cloned. To create the C2BDN line, three nucleotide exchanges were included, two leading to the exchange of aspartic acid residues in position 705 and 711 of Munc13-1 to asparagine (GAC to AAC), and one that eliminates a nearby BamHI site (GGA to GGC) but preserves the glycine residue (pTKPuroFRT-UNC13A Exons 13-26 Exon 18* C2BDN; Figures 1A and 1D). To create the C2BKW line, the targeting vector included two nucleotide exchanges, one leading to the replacement of a lysine residues in position 706 of Munc13-1 by a tryptophan (AAG to TGG), and one to eliminate a nearby BamHI site (GGA to GGC), preserving the glycine residue (pTKPuroFRT-UNC13A Exons 13-26 Exon 18* C2BKW; Figures 1A and 1D). The resulting vectors were used to electroporate stem cells, and the correct genomic integration of the cassette was identified by the acquisition of puromycin resistance (Thomas and Capecchi, 1987), and by long-range PCR amplification and DNA sequencing. Positive clones were amplified, injected into blastocysts, and the resulting mice were screened for germline transmission via long-range PCR amplification. To eliminate the puromycin resistance cassette, mice were crossed with Gt(ROSA)26Soa-bar(FLP1)Dym mice (Farley et al., 2000). Offspring were analyzed using genotyping PCR (Sense prime: 5′-GGTAGCAGAGGTTATGGATAT-3′; Antisense prime: 5′-TGTTGCCAGTTTCAGAGGC-3′), and sequencing (Figures 1B–1D), and animals in which a successful cre recombination had occurred were selected for further breeding with C57BL/6N mice for three generations. Mice were then cross-bred to produce homozygous and wt littermates for experiments.

**Mouse Maintenance**

Mutant mouse generation and animal experiments were approved by the responsible authorities of the local government (Lower Saxony State Office for Consumer Protection and Food Safety (LAVES; permit 33.19-42502-04-15/1817). Animals were maintained in groups in accordance with European Union Directive 63/2010/EU and ETS. Animal health was controlled daily by caretakers and by a veterinarian, and a quarterly health monitoring was done according to FELASA recommendations with either NMRI sentinel mice or animals from the colony (serological analyses; microbiological, parasitological, and pathological examinations). Abnormal findings were not made during the period of the study. Mice were kept in individually ventilated cages, under specific pathogen-free conditions, 21 ± 1°C, 55% relative humidity, 12 h/12 h light/dark cycle). Food and tap water, as well as bedding and nesting material, were provided ad libitum, and cages were changed once a week. P14–17 littermate homozygous Munc13-1 KI mice and wt controls of either sex were analyzed. Littermates were used to minimize effects of genetic background, although only minor differences were observed between wt lines (Table S1). Mice were routinely genotyped by PCR (Figures 1B–1D). We did not observe large deviations from the expected Mendelian ratios among offspring, although for C2BKW mice a slightly lower than expected fraction of homozygous KI mice was obtained, possibly indicating a slightly increased perinatal lethality. C57BL/6N mice were used for the experiments shown in Figures 6D, S3B, S3C, and S7.

**METHOD DETAILS**

**Western Blot Analyses**

P2 crude synaptosomal fractions were obtained from cortical brain tissue of P16 KI mice and wt littermates as previously described (Lipstein et al., 2013). Between 2-20 μg protein were separated on 4%–12% gradient Bis-Tris polyacrylamide gels (Invitrogen) and blotted onto nitrocellulose membranes. The protein load per lane was quantified by a reversible membrane staining protocol (MemCode; Pierce) and quantified using ImageJ. The following antibodies were subsequently used to identify the indicated proteins: rabbit polyclonal (rp) anti-Munc13-1 (40), anti-ubMunc13-2, and anti-bMunc13-2 (Varoqueaux et al., 2005), rp-anti-Doc2 (SySy; 174 103), rp-anti-CAPS-1 (SySy 262 013), rp-anti-Munc18-1 (SySy; 116 002), mouse monoclonal (mm)-anti Synapsin 1 (SySy 106 011), mm-anti-Syntaxin 1A/B (SySy 110 011), mm-anti-Synaptobrevin 2 (SySy 104 211), mm-anti-Synaptotagmin (SySy 105 001), rp anti-Rim (lab antibody; Q703), mm anti-SNAP25 (SySy; 111 011). After incubation with corresponding secondary antibodies ( Peroxidase-AffiniPure Goat Anti-Mouse IgG or peroxidase-AffiniPure Goat Anti-Rabbit IgG antibody), the western blot signal was detected with an INTAS imager (INTAS Science Imaging), quantified using ImageJ, and normalized to the protein load. Between 3-4 independent analyses were performed per condition. Quantification is presented as mean ± SEM.

**Co-Immunoprecipitation Experiments**

Co-immunoprecipitation was performed according to Lipstein et al. (2013). Briefly, crude synaptosomal fractions obtained from cerebral cortic of adult (8–11 weeks) wt and KI mice were solubilized and ultracentrifuged using a Beckmann L-70 at 100,000 g to remove
insoluble material. A sample ("input") was collected, and the remaining fraction was incubated with a rp-anti-Munc13-1 (40). Sepharose-Protein G beads (Invitrogen) were added to capture the antibody and associated proteins, washed to remove background, and eluted using denaturating Laemmli buffer ("IP" samples). The samples were loaded on 4%–12% gradient Bis-Tris polyacrylamide gels and blotted according to a modified version of the Sigma protocol (product C7055) with mm-anti-Calmodulin (Upstate Biotechnology). Munc13-1 was blotted using the mm-anti Munc13-1 (HS5; Betz et al., 1998).

Immunostaining

Immunostaining experiments were performed on P15 coronal brainstem sections using primary antibodies against Munc13-1, Bassoon, and MAP-2. Brains were rapidly frozen in isopentane (−35°C). Sixteen μm-thick coronal cryosections of the MNTB region were cut and mounted on Superfrost slides, air-dried for 15 min, and immersion-fixed in ice-cold 4% paraformaldehyde solution (4% PFA in 0.1 M PB, pH 7.4) for 5 min at room temperature (RT). To ensure similar fixation and labeling conditions between genotypes, sections of mutant and corresponding wt mice were mounted together. Sections were incubated for 90 min at RT in blocking solution (0.1 M PB, 5% normal goat serum, 0.1% cold water fish skin gelatine, 0.5% Triton X-100, pH 7.4) before being treated overnight at 4°C with the primary antibodies rp-anti-Munc13-1 Ab (SySy 126 103, 1:400), mm-anti-Bassoon (Enzo Life Sciences SAP7F407 1:400), and chicken polyclonal anti-MAP2 (Novus NB300-213, 1:600), diluted in incubation buffer (0.1 M PB, 3% normal goat serum, 0.1% cold water fish skin gelatine, 0.3% Triton X-100, pH 7.4). After washing in PB, sections were incubated for 2 h at RT in the dark with the fluorescent secondary antibodies Alexa 488-coupled goat anti-rabbit, Alexa-555-coupled goat anti-mouse, and Alexa-633-coupled goat anti-chicken (Invitrogen, 1:1000) diluted in incubation buffer. Coverslips were washed with Aqua-PolyMount (Polysciences). Confocal laser scanning micrographs of presynaptic compartments surrounding MNTB PNs were acquired with a Leica TCS SP5 II confocal microscope. An HCX PL APO lambda blue 63 x water immersion objective (NA = 1.2) and a pinhole setting of 0.38 AU were used to obtain single-plane micrographs (512 x 512; x-y pixel spacing = 48.1 nm) in sequential scanning mode. Laser power and gain were adjusted to ensure that signals were in the linear range of detection. Confocal images were subjected to deconvolution using two ImageJ (National Institutes of Health; Bethesda, MD) plugins: point spread functions were generated using -Diffraction PSF 3D plugin, and deconvolution was performed using DeconvolutionLab plugin (Biomedical Imaging Group, EPFL; Lausanne, Switzerland). Individual Munc13-1 immunoreactive puncta were detected using a thresholding algorithm (ImageJ). For all Munc13-1-positive puncta within ~1 μm wide regions of interest (ROIs) drawn around the perimeter of MNTB PNs and approximately delineating the presynaptic compartments, the center of mass was determined and those coordinates were exported for further processing. For each ROI, the average density of Munc13-1-positive puncta per μm perimeter was obtained and the mean nearest-neighbor distance was calculated as a parameter characterizing the distribution of Munc13-1-positive puncta within a given ROI.

Slice Preparation

Acute brainstem slices of postnatal (P14–P17) mice of either sex were prepared as previously described (Chang et al., 2015). After decapitation, brains were immersed in ice-cold low-Ca2+/Mg2+ artificial CSF (aCSF) containing (in mM): 125 NaCl, 2.5 KCl, 1.25 MgCl2, 25 glucose, 25 NaHCO3, 1.25 NaH2PO4, 0.4 ascorbic acid, 3 myoinositol, and 2 Na-pyruvate (pH 7.4), bubbled with 95% O2, 5% CO2. The brainstem was glued onto the stage of a VT1000S vibratome (Leica), and 200 μm-thick coronal slices containing the medial nucleus of the trapezoid body (MNTB) were cut. Slices were incubated for 40 min at 35°C in a chamber containing normal aCSF (identical to low-Ca2+/Mg2+ aCSF, except that 3 mM MgCl2 and 0.1 mM CaCl2 were replaced with 1.3 mM MgCl2 and 1.7 mM CaCl2). Slices were kept at RT (21–24°C) for up to 5 h after recovery. Acute parasagittal hippocampal slices (300 μm thick) were prepared from P16–P21 mice of either sex using a VT1000S vibratome and ice-cold low-Ca2+/high-Mg2+ sucrose-based cutting solution containing (in mM): 120 sucrose, 64 NaCl, 25 NaHCO3, 2.5 KCl, 1.25 NaH2PO4, 10 glucose, 0.5 CaCl2 and 7 MgCl2 (pH 7.4, bubbled with 95% O2, 5% CO2). Slices were incubated for 40 min at 35°C in a chamber filled with normal aCSF containing 1 mM MgCl2 and 2 mM CaCl2. Slices were kept at RT for up to 5 h after recovery.

Electrophysiology

Whole-cell patch-clamp recordings were made from calyx of Held terminals and principal neurons (PNs) of the MNTB at RT using an EPC-10 amplifier controlled by Pulse or PatchMaster software (HEKA Elektronik). Patch-pipettes (Science Products) were coated with dental wax in order to minimize fast capacitative transients during voltage-clamp experiments and to reduce stray capacitance. For postsynaptic recordings, pipettes with an open-tip resistance of 2.5–4 MΩ were filled with a solution containing (in mM): 100 K-glucuronate, 60 KCl, 5 Na2-phosphocreatine, 10 HEPES, 5 EGTA, 0.3 Na2-GTP, and 4 ATP-Mg, pH 7.3, with KOH. During experiments, slices were continuously perfused with normal aCSF solution containing 1.3 mM MgCl2 and 1.7 mM CaCl2 and supplemented with 5 μM strychnine, to block glycineergic inputs. Cells were visualized by infrared-differential interference contrast microscopy through a 40X water-immersion objective using an upright BX51WI microscope (Olympus). All experiments were performed at RT. A bipolar stimulation electrode was used to evoke presynaptic APs (stimulus intensity ≤ 20 V, 100 μs duration). Series resistance (Rs) was ≤ 8 MΩ and compensated ≥ 82%. Holding potential (Vh) and leak current were ~700 mV and ≤ 200 pA, respectively. Sampling interval and low-pass filter settings were 20 μs and 5.0 kHz, respectively, eEPSC peaks measured in P14-17 mouse calyx synapses frequently exceed amplitudes of 10 nA (Chang et al., 2015; Joshi and Wang, 2002). Assuming a maximum Rp = 8 MΩ that is 80% compensated by the amplifier Rp-compensation circuitry, such peak amplitudes cause a transient voltage escape that corresponds to a >20% drop in driving force at Vh = −70 mV for a synaptic conductance having a reversal potential of ~0 mV. To reduce
eEPSC amplitudes for improved voltage-clamp and in order to attenuate postsynaptic AMPAR saturation and AMPAR desensitization, all experiments were performed in the continuous presence of 1 mM of the low-affinity GluR antagonist kynurenic acid (kyn), unless explicitly stated otherwise. The blocking ratio eEPSC_{kyn}/eEPSC_{ctrl} is shown for a subset of the experiments in Figure S1. Furthermore, voltage-clamp errors caused by remaining uncompensated Rs were corrected offline by a software routine (see offline analysis).

Presynaptic voltage-clamp recordings were performed using patch pipettes with an open-tip resistance of 3–4.5 MΩ. Series resistance was ≤ 15 MΩ and Rs was compensated 60%–65%. For measuring h_{Ca(V)} and membrane capacitance (ΔC_m), pipettes were filled with a Cs-gluconate based solution containing (in mM): 100 Cs-gluconate, 30 TEA-Cl, 30 CsCl, 10 HEPES, 0.05 EGTA, 5 Na_2-phosphocreatine, 4 ATP-Mg, 0.3 GTP, pH 7.3 with CsOH. The bath solution was supplemented with 1 μM TTX, 1 mM 4-AP, and 40 mM TEA-Cl to suppress voltage-gated sodium and potassium currents. Calyx terminals were visualized by oblique illumination (Dodt gradient contrast) through a 60 x water-immersion objective using an upright BX51WI microscope (Olympus). All experiments were performed at RT. The size of the readily releasable pool of SVs was estimated by monitoring ΔC_m using the sine + DC technique (Lindau and Neher, 1988) using the software lock-in amplifier implemented in PatchMaster (HEKA Elektronik) by adding a 1 kHz sine-wave voltage command (peak-to-peak amplitude ± 35 mV) to V_h = −80 mV. Presynaptic recordings with a leak current > 200 pA were excluded from the analysis. In eEPSC train recordings and during recordings of eEPSC recovery from depression, ≥ 3 repetitions per protocol were recorded for each cell included in the final analysis.

Whole-cell patch-clamp recordings from hippocampal CA1 pyramidal neurons were made at RT with patch-pipettes (open-tip resistance 2.5–3.5 MΩ) filled with a solution containing (in mM): 130 K-gluconate, 10 KCl, 2 MgCl_2, 2 Na_2ATP, 10 HEPES, 10 EGTA, pH 7.3 with KOH. During experiments, slices were continuously perfused with normal aCSF solution containing 1 mM MgCl_2 and 2 mM CaCl_2, supplemented with 25 μM bicuculine methiodide (HelloBio) to block GABA A,R-mediated IPSCs. Schaffer collateral/commissural fibers were stimulated with a glass electrode filled with aCSF and placed in stratum radiatum ≥ 80 μm away from the cell body. Transmitter release was evoked by applying brief electrical pulses (100 μs, 10–50 V). Stimulus intensity was adjusted to obtain peak EPSC amplitudes in the range of 80–300 pA (mean values 161 ± 18 pA, n = 13, and 172 ± 23 pA, n = 13, for wt KW and mut KW synapses, respectively) when stimulated with a single APs. Assuming a mean unitary EPSC amplitude of 10–20 pA, this corresponds to the recruitment of 4-30 Schaffer collateral/commissural fibers.

**Ca^{2+} Imaging in Nearly Unperturbed Terminals**

For measuring AP-evoked Ca^{2+} transients in nearly unperturbed calyces, terminals were preloaded with the low affinity Ca^{2+} dye Cal520-FF (K_d = 9.8 μM). Presynaptic patch pipettes were filled with a K-gluconate based solution containing (in mM): 100 K-gluconate, 60 KCl, 10 HEPES, 5 EGTA, 5 Na_2-phosphocreatine, 4 ATP-Mg, 0.3 GTP, pH 7.3 with KOH, to which 400 μM Cal520-FF were added from stock solution stored at −20 C. Calyx terminals were visualized by oblique illumination (Dodt gradient contrast) through a 60 x water-immersion objective (NA = 1.0) using an upright BX51WI microscope (Olympus). To preload calyx terminals with a final concentration of ≈200 μM Cal520-FF, the whole-cell recording configuration was established for a short period of 40–60 s during which the presynaptic firing reliability was tested by eliciting calyceal APs by afferent fiber stimulation using a single stimulus, a 100 Hz train (25 stimuli) and a 200 Hz train (50 stimuli). Presynaptic APs were recorded in the current-clamp mode of the EPC-10 after carefully adjusting the fast-capacitance cancellation in cell-attached mode. Thereafter the pipette was gently retracted to preserve the cytosolic environment. The Ca^{2+} indicator dye was excited at 490 nm using a custom-built LED light source. Fluorescent images were collected at a rate of 100 frames per s with a Grasshopper3 camera (CMOS, FLIR, Germany). The imaging hardware was controlled by Micro-Manager 2.0 software.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Time series image stacks were analyzed offline using ImageJ (Schneider et al., 2012) and Igor Pro (Wavemetrics). Offline analysis of electrophysiological data was performed using Igor Pro, R (R Project for Statistical Computing) or Microsoft Excel. Voltage-clamp errors caused by remaining uncompensated Rs were fully compensated by applying a software correction procedure similar to that described in Traynelis (1998) and courteously provided by E. Neher.

Statistical analysis was conducted using Igor Pro, R and Microsoft Excel. Data are expressed as mean ± standard error of the mean (SEM). Error bars in all graphs indicate SEM. SEM and 95% confidence intervals for FRP and δ estimates derived from eEPSC trains (see below) were obtained by bootstrap resampling analysis using a balanced bootstrap approach (every experimental observation appeared exactly the same number of times in the total population of 10,000 bootstrap samples) (Davison et al., 1986; Gleason, 1988). If not stated otherwise, a two-tailed Welch–Satterthwaite t test was used to test for statistical significance of differences between sample means (see p values in Table S1). In Table S1, the number of animals used in each experiment is indicated by a capital N, whereas the number of cells tested is indicated by a lowercase n.

**eEPSC Trains and Paired-Pulse Ratios (PPR)**

For a train of presynaptic APs, the prediction of a simple depletion model featuring a constant SV recruitment rate constant k_s = 1/N, for the number of occupied sites N_l immediately before arrival of the i+1th AP can be calculated recursively according to
where $N_{\text{total}}$ is the sum of all docking sites, $p_i$ is the release probability at AP arrival, $\Delta t$ is the inter-stimulus interval, and $\tau$ is the time constant of recovery.

For simplicity, we assume here $N_1 = N_{\text{total}}$.

Assuming that $p$ increases immediately following an AP from its current value by a constant fraction $a$ toward a maximum value of 1, we can express $p_i$ recursively as

$$p_{i+1} = (p_i + a \times (1 - p_i) - p_0) \times e^{-\Delta t/\tau} + p_0; \text{ for } i \geq 1$$

where $p_0$ is the release probability at rest, $\tau$ is the time constant of synaptic facilitation, and $\Delta t$ is the inter-stimulus interval.

For the paired-pulse ratio (PPR) of the first two consecutive eEPSCs we can therefore write

$$PPR = \frac{\text{EPSC}_2}{\text{EPSC}_1} \times \frac{N_2 \times p_2}{N_1 \times p_1} = \left(1 - p_0 \times e^{-\Delta t/\tau}\right) \times \left(1 + b \times e^{\Delta t/\tau}\right); b = a \times \left(\frac{1}{p_0} - 1\right)$$

Readily-Releasable SV Pool and Release Probability

The whole entity of SVs that can be released within about 50 ms in response to strong and prolonged presynaptic [Ca$^{2+}$] elevations elicited by presynaptic Ca$^{2+}$ uncaging or direct presynaptic depolarizations can be subdivided into two main SV subpools – ‘fast releasing’ (FRP) and ‘slowly releasing’ (SRP) SVs. The SRP only marginally contributes to AP-evoked eEPSCs primarily represent the FRP. There is good experimental evidence showing that the FRP itself is functionally inhomogeneous (Lee et al., 2013; Taschenberger et al., 2016). For simplicity, we neglect such functional heterogeneity of the FRP here and report average $p$ values ($\bar{p}$) instead. Assuming, for example, that the FRP of calyx terminals is composed of two population of SVs having different $p$, we can write

$$\text{EPSC} = q \times N \times \bar{p} = q \times \left(N_{\text{high}} \times p_{\text{high}} + N_{\text{low}} \times p_{\text{low}}\right) = q \times N \times \left(\frac{N_{\text{high}}}{N} \times p_{\text{high}} + \frac{N_{\text{low}}}{N} \times p_{\text{low}}\right)$$

where $N$ is the sum of occupied sites $N_{\text{high}} + N_{\text{low}}$, $q$ is the quantal size and the subscripts identify high- and low-$p$ release sites. In this case, $\bar{p}$ is simply a weighted average

$$\bar{p} = \frac{N_{\text{high}} \times p_{\text{high}} + N_{\text{low}} \times p_{\text{low}}}{N_{\text{high}} + N_{\text{low}}}. \text{ During train stimulation, } N_{\text{high}} \text{ decreases more rapidly than } N_{\text{low}} \text{ because } p_{\text{high}} > p_{\text{low}}. \text{ This causes } \bar{p} \text{ to decrease. This decrease is partially compensated by synaptic facilitation. For simplicity, we assume here that } \bar{p} \text{ remains approximately similar throughout the trains.}$$

We obtained an estimate for the FRP from cumulative eEPSCs measured in response to high-frequency trains (50, 100 and 200 Hz). The cumulative eEPSC amplitude was corrected for SV recruitment by fitting a line to the final 4 eEPSCs representing the steady state (eEPSC$_{\text{ss}}$) and back-extrapolating that line to eEPSC$_{1}$ (Neher, 2015; Schneggenburger et al., 1999). Such approach rests on the assumptions that (i) all release at steady state is balanced by SV replenishment which occurs at constant rate throughout a stimulus train and that (ii) the FRP is fully emptied after each AP under steady-state conditions. Neither of these two conditions is fully met: SV replenishment – the product of replenishment rate constant $k_r$ and number of empty docking sites $N_e$ – operates at considerably higher rate at steady state in comparison to the onset of the stimulus train because: (i) $k_r$ is upregulated during train stimulation and (ii) $N_e$ is still low at the beginning of stimulation but increases during ongoing stimulation. In addition, stimulation protocols featuring high frequency AP trains are unlikely to deplete the FRP completely and therefore the back-extrapolation approach measures only the depleted fraction of the FRP (for discussion see Neher, 2015).

Thus, the back-extrapolation approach is expected to underestimate the actual FRP size. We confirm this by observing that FRP estimates obtained at 50 and 100 Hz stimulation were consistently lower than those obtained at 200 Hz stimulation. We therefore refer to such estimates as apparent pool sizes $\text{FRP}_{50\text{Hz}}, \text{FRP}_{100\text{Hz}}$ and $\text{FRP}_{200\text{Hz}}$. When plotting $1/\text{FRP}$ for 50, 100 and 200 Hz as a function of $1/t_{\text{stim}}$, where $t_{\text{stim}}$ is the stimulation frequency, we notice that this relationship is nearly linear and we can therefore obtain a correct estimated $\text{FRP}_{\text{corr}}$ for the case $t_{\text{stim}} = 0$ s by linear regression as the inverse of intercept of such regression line with the abscissa (Figures 4B and 5C).

Steady-State Replenishment Rate Constant ($k_r$)

At steady state, the average rate of release is balanced by the average rate of replenishment $r_r = r_r$, with $r_r = N_e \times \bar{p} \times f_{\text{stim}}$ and $r_r = (N_{\text{total}} - N_{\text{ss}}) \times k_r$, where $N_{\text{total}}$ is the total number of sites (occupied or empty), $N_{\text{ss}}$ is the number of occupied sites at steady state immediately before AP arrival, and $N_{\text{ss}}$ and $k_r$ are the average number of occupied sites and the average replenishment rate constant.
during the inter-stimulus interval at steady state, respectively. For high-frequency stimulation that leads to strong SV pool depletion, a majority of docking sites is empty such that the difference between $N_{ss}$ and $\bar{N}_{ss}$ is negligible in comparisons to the number of empty sites $\bar{N}_{total} - \bar{N}_{ss}$ and we can write $\bar{N}_{ss} = N_{ss}/(N_{total} - N_{ss}) \times p_{ss} \times f_{stim}$.

With $EPSC_1 = N_{total} \times p_1 \times q$ and $EPSC_{ss} = N_{ss} \times p_{ss} \times q$, and if we further assume that $p_{ss} \approx p_1$, then we obtain $\bar{R}_s = EPSC_{ss} \times f_{stim} \times \frac{P_{ss}}{EPSC_{ss}}$. $\bar{R}_s$ is accurate only within the limits of the simplifications described above but nevertheless serves as a useful parameter for comparing the functional differences among $C_2BDN$, $C_2BKW$ and WR mutant mice.

**STP at Hippocampal SC/C-CA1 Synapses**

Synaptic responses ($R$) at SC/C-CA1 synapses were modeled as being proportional to the product of a depletable resource ($N$) and a probability by which this resource is consumed by an AP ($p$):

$$R_i \sim N_i \times p_i$$

Postsynaptic sensitivity to glutamate ($q$) was assumed to be invariant. All eEPSC train amplitudes were normalized to the respective peak amplitude of the first eEPSC during the conditioning 10 Hz stimulus trains. Changes in synaptic strength during conditioning trains and during subsequent recovery from depression were simulated using a simple model considering depletion of $N$ and a slowly decaying augmentation of $p$. A contribution of fast decaying paired-pulse facilitation was small under our experimental conditions and therefore neglected. The magnitude of $N$ immediately before arrival of the $i+1$th AP was calculated according to:

$$N_{i+1} = \left(1 - p_i\right) \times N_i + \left(N_{total} - (1 - p_i) \times N_i\right) \times \left(1 - e^{-\frac{\Delta t}{\tau}}\right)$$

$$p_{i+1} = (p_i + a \times (1 - p_i) - p_0) \times e^{-\frac{\Delta t}{\tau_a}} + p_0,$$ for $i \geq 1$

where $N_0$ designates the resting value of $N$ which for the analysis shown in Figure 8H equates to $1/p_1$ because of the normalization of all $R_i$ relative to $R_1$, $p_i$ is the release probability at arrival of the $i$th AP, $\Delta t$ is the inter-stimulus interval, and $\tau$ is the time constant of $N$. Assuming that $p$ increases immediately following an AP from its current value by a constant fraction $a$ toward a maximum value of 1, we can obtain $p$ recursively as:

$$p_{i+1} = (p_i + a \times (1 - p_i) - p_0) \times e^{-\frac{\Delta t}{\tau_a}} + p_0,$$ for $i \geq 1$

where $p_0$ is the resting value of $p$, $\tau_a$ is the decay time constant of augmentation, and $\Delta t$ is the inter-stimulus interval. Fit results for $p_0$ and $a$ were 0.06 and 0.002 in wtKW and 0.08 and 0.003 in mutKW synapses, respectively. The time constant of augmentation $\tau_a$ amounted to $\sim 9$ s for both wtKW and mutKW synapses.