Cellular/Molecular

Membrane Penetration by Synaptotagmin Is Required for Coupling Calcium Binding to Vesicle Fusion *In Vivo*

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The vesicle protein synaptotagmin I is the Ca $^{2+}$ sensor that triggers fast, synchronous release of neurotransmitter. Specifically, Ca $^{2+}$ binding by the C $_2$ B domain of synaptotagmin is required at intact synapses, yet the mechanism whereby Ca $^{2+}$ binding results in vesicle fusion remains controversial. Ca $^{2+}$ -dependent interactions between synaptotagmin and SNARE (soluble *N*-ethylmaleimide-sensitive fusion protein attachment receptor) complexes and/or anionic membranes are possible effector interactions. However, no effector-interaction mutations to date impact synaptic transmission as severely as mutation of the C $_2$ B Ca $^{2+}$ -binding motif, suggesting that these interactions are facilitatory rather than essential. Here we use *Drosophila* to show the functional role of a highly conserved, hydrophobic residue located at the tip of each of the two Ca $^{2+}$ -binding pockets of synaptotagmin. Mutation of this residue in the C $_2$ A domain (F286) resulted in a \sim 50% decrease in evoked transmitter release at an intact synapse, again indicative of a facilitatory role. Mutation of this hydrophobic residue in the C $_2$ B domain (I420), on the other hand, blocked all locomotion, was embryonic lethal even in syt I heterozygotes, and resulted in less evoked transmitter release than that in syt^{null} mutants, which is more severe than the phenotype of C $_2$ B Ca $^{2+}$ -binding mutants. Thus, mutation of a single, C $_2$ B hydrophobic residue required for Ca $^{2+}$ -dependent penetration of anionic membranes results in the most severe disruption of synaptotagmin function *in vivo* to date. Our results provide direct support for the hypothesis that plasma membrane penetration, specifically by the C $_2$ B domain of synaptotagmin, is the critical effector interaction for coupling Ca $^{2+}$ binding with vesicle fusion.

Introduction

During synaptic transmission, ${\rm Ca}^{2^+}$ influx into the presynaptic terminal triggers fast, synchronous neurotransmitter release. This process requires mechanisms both for sensing ${\rm Ca}^{2^+}$ and subsequently fusing transmitter-filled synaptic vesicles with the presynaptic membrane. The SNARE (soluble *N*-ethylmaleimidesensitive fusion protein attachment receptor) proteins are required for vesicle fusion events in both constitutive and ${\rm Ca}^{2^+}$ regulated vesicle transport pathways (Jahn and Scheller, 2006). At the nerve terminal, however, additional levels of regulation are required to prevent the fusion of synaptic vesicles with the presynaptic membrane before ${\rm Ca}^{2^+}$ entry and then expedite the synchronous fusion of multiple vesicles when ${\rm Ca}^{2^+}$ levels rise. The synaptic vesicle protein synaptotagmin (syt) is the major

Ca²⁺ sensor that both participates in reducing spontaneous fusion events in the absence of Ca²⁺ and triggers the synchronous fusion of synaptic vesicles with the presynaptic membrane during Ca²⁺ influx (DiAntonio and Schwarz, 1994; Geppert et al., 1994; Littleton et al., 1994; Chicka et al., 2008). Thus, synaptotagmin is a key regulator of SNARE-mediated fusion during synaptic transmission

Specifically, Ca²⁺ binding by the C₂B domain of synaptotagmin is required to trigger fast synchronous vesicle fusion (Mackler et al., 2002; Nishiki and Augustine, 2004). Mutations within the C₂B Ca²⁺-binding motif (see Fig. 1, C₂B, D₁-D₄) result in more severe deficits than complete removal of synaptotagmin I (Geppert et al., 1994; Loewen et al., 2001; Mackler et al., 2002; Nishiki and Augustine, 2004). Such mutations not only block Ca²⁺-evoked, synchronous release at an intact synapse and result in lethality but also dominantly inhibit the ability of wild-type (WT) synaptotagmin to trigger fusion when these proteins are coexpressed (Mackler et al., 2002). However, the mechanism that couples Ca²⁺ binding by synaptotagmin with vesicle fusion remains controversial. Any downstream, effector interactions required for coupling Ca²⁺-bound synaptotagmin to vesicle fusion should cause as severe an effect as mutation of the C₂B Ca²⁺binding site itself. Biochemical experiments and work on cultured cells suggest that Ca2+-dependent interactions with the SNARE complex and Ca²⁺-dependent interactions with the presynaptic membrane are likely candidates for effector interactions

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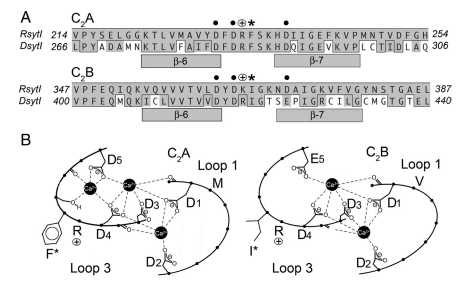


Figure 1. Both the C_2A and C_2B domains of synaptotagmin have conserved hydrophobic residues at the tip of the Ca $^{2+}$ -binding pocket. **A**, Alignment of synaptotagmin I from rat and *Drosophila*. Bars indicate β -sheets, asterisks indicate the conserved hydrophobic residues mutated in this study, dots indicate Ca $^{2+}$ -binding residues, and open circles with a plus indicate conserved basic residues. Within the alignment, conserved residues are shown in gray and identical residues are boxed. **B**, Schematic representation of loops 1 and 3 in *Drosophila* that form the Ca $^{2+}$ -binding pockets of each C_2 domain. D_1 – D_5 and E_5 indicate Ca $^{2+}$ -binding residues. F*, M, I*, and V indicate hydrophobic residues at the tip of the Ca $^{2+}$ -binding pockets.

(Chapman et al., 1995; Gerona et al., 2000; Fernández-Chacón et al., 2001; Bai et al., 2002; Rhee et al., 2005; Herrick et al., 2006; Lynch et al., 2007). However, all mutations that disrupt these interactions that have been examined at synapses result in only a partial disruption in fusion (Fernández-Chacón et al., 2001; Li et al., 2006; Paddock et al., 2008); none result in lethality. Here we report the most severe disruption of function by any synaptotagmin mutation identified to date.

Materials and Methods

Site-directed mutagenesis

Phenylalanine 286 (Fig. 1, F*) and isoleucine 420 (Fig. 1, I*) of *Drosophila* synaptotagmin were mutated using PCR. For F286, mutant antisense oligonucleotide CGCCGATCTGGTCGTGCTTCGATTCGCGATCGAAG TCGAAAATGGC was used to incorporate a glutamate residue (syt^{A-FE}), and CGCCGATCTGGTCGTGCTTCGAGTAGCGATCGAAGTCGAAAA TGGC was used to incorporate a tyrosine residue (syt^{A-FY}). For I420, mutant antisense oligonucleotide TGCAGCGGCCGATGGGTTCGGAGGTGC-CTTCACGATCGTAGTCCACGACGGTCACAACG was used to incorporate a glutamate residue (syt^{B-IE}). Mutant PCR products were ligated into otherwise wild-type *Drosophila synaptotagmin I c*DNA in pBluescript II, sequenced to ensure that the only change introduced by PCR was the desired mutation, and subcloned into a pUAST vector to place the mutant syt under the control of the upstream activating sequence (UAS) promoter (Brand and Perrimon, 1993; Yao and White, 1994).

Generation of mutant transgenic lines

Drosophila embryos were transfected with the mutant pUAST plasmids as described previously (Paddock et al., 2008) or by Genetic Services. Expression of the transgene was localized to the nervous system using the pan-neuronal promoter elav to drive galactosidase-4 (Gal4) expression, and the Gal4/UAS system was used to amplify expression of the syt transgenes (Brand and Perrimon, 1993; Yao and White, 1994). Because no gender selection was used, a mix of male and female animals were used for all experiments. Lines used included the following: yw; syt^{null}elav-GAL4/CyO, GFP (source of Gal4 to drive transgene expression in the nervous system), yw; syt^{null}/CyO, y+; P[UAS syt^{A-F286F}] lines 3 and 5 (P[syt^{A-FY}] lines), yw; syt^{null}/CyO, y+; P[UAS syt^{B-I420E}] lines 4 and 6 (P[syt^{B-IE}] lines), and yw; syt^{null}/CyO, y+; P[UAS syt^{WT}] (transgenic control, P[syt^{WT}]). Stan-

dard genetic techniques were used to cross the transgenes into the *syt*^{null} background; selected larvae were green fluorescent protein (GFP) negative and y negative without regard to gender. Unless specifically noted, experiments were performed in the *syt*^{null} background to express the transgene in the absence of endogenous synaptotagmin I.

Analysis of dominant-negative lethality phenotype

The effect of the syt B-IE mutant protein was examined both in the absence and presence of expression from the wild type synaptotagmin gene. Drosophila lines used were: yw; syt^{null}elavGAL4/CyO, GFP: yw; syt^{null}/CyO, y+; P[UAS syt^{B-1420E}] lines 4 & 6: and yw; syt^{null}/ CyO, y+; $P[UAS syt^{WT}]$. Parental lines contain the native syt gene on the CvO balancer chromosome to maintain healthy stocks, and transgene expression is under the control of a UAS promoter that requires Gal4 to promote expression. The following lines were crossed on molasses plates, and progeny were monitored for the ability to hatch: yw; syt^{null}/CyO , y+; P[UASsyt]/P[UASsyt] were crossed to yw; syt*null elavGal4/CyO*, *GFP*; +/+. The four possible progeny are of the following genotypes: yw; sytnull/sytnullelavGal4; P[UASsyt]/+ (transgenic syt alone, no y+, no GFP); yw; syt^{null}/CyO,

GFP; P[UASsyt]/+ (native syt alone attributable to CyO and lack of the transgene driver elavGal4, no y+, GFP present); yw; $syt^{null}elavGal4/CyO$, y+; P[UASsyt]/+ (native + transgenic syt attributable to CyO and transgene driver elavGal4, y+, no GFP); and yw; CyO, y+/CyO, GFP; P[UASsyt]/+ (y+, GFP present). Balancer chromosome homozygotes, CyO, y+/CyO, GFP, are embryonic lethal and do not hatch. $P[UASsyt^{B-IE}]$ lines were crossed in parallel with $P[UASsyt^{WT}]$ lines.

Immunolabeling and immunoblotting

Saline used was HL3 [in mm: 5 KCl, 1.5 CaCl₂, 70 NaCl, 20 MgCl₂, 10 NaCHO₃, 5 HEPES, 115 sucrose, and 5 trehalose (Stewart et al., 1994)]. Immunolabeling and immunoblotting of third-instar larvae were performed by standard techniques (Paddock et al., 2008) using the anti-synaptotagmin antibody Dsyt-CL1 (Mackler et al., 2002). For immunolabeling of embryos, mutants were dechorionated in 50% bleach and glued (GLUture; Abbott Laboratories) to Sylgard-coated dishes containing Ca²⁺-free HL3. The cuticle was perforated and opened along the dorsal midline with a glass needle, and the gut was removed to expose the CNS. After fixation in 4% paraformaldehyde in PBS, synapses were labeled with either Dsyt-CL1 (diluted 1:200) or an anti-cysteine string protein (CSP) monoclonal antibody (diluted 1:50) (Zinsmaier et al., 1994) overnight at 4°C. During an overnight incubation at 4°C, Dsyt-CL1 was labeled with an Alexa Fluor-488 goat-anti-rabbit antibody (diluted 1:500; Invitrogen), whereas CSP was labeled with an Alexa Fluor-488 rabbit anti-mouse antibody (diluted 1:200; Invitrogen).

Electrophysiology

Third-instar recordings. Evoked and spontaneous excitatory junction potentials [EJPs and miniature EJPs (mEJPs)] were recorded from ventral longitudinal muscle 6 of segments 3 and 4 in HL3 saline as described previously (Loewen et al., 2001; Mackler and Reist, 2001; Paddock et al., 2008). Briefly, third-instar larvae were dissected in Ca $^{2+}$ -free HL3 to expose the body wall musculature. After changing to HL3 containing 1.5 mM Ca $^{2+}$, muscle 6 was impaled with a 10–40 MΩ electrode filled with a solution of three parts 2 M potassium citrate to one part 3 M potassium chloride. Evoked EJPs were generated by stimulating segmental nerves with 1 ms pulses of 30–50 nA at 0.05 Hz. Intracellular recordings of spontaneous mEJPs were collected and analyzed as described previously (Mackler and Reist, 2001). One-way ANOVA and Tukey range test were used for statistical comparisons.

Embryonic recordings. To record synaptic currents, *Drosophila* embryos (22–24 h after egg laying) were dechorionated with bleach and manually dissected as described previously (Featherstone et al., 2009). Embryonic ventral longitudinal muscle 6 from segment 3 or 4 was then whole-cell voltage clamped at -60 mV in bath solution [containing in mm: 135 NaCl, 5 KCl, 4 MgCl₂, 1.8 CaCl₂, 72 sucrose, and 5 *N*-[Tris(hydroxymethyl)methyl]-2-

aminoethanosulfonic acid (TES), pH 7.2], as described previously (Chen et al., 2009). The patch pipette solution contained the following (in mm): 120 KCl, 20 KOH, 4 MgCl₂, 0.25 CaCl₂, 5 EGTA, 36 sucrose, and 5 TES. Evoked synaptic currents were triggered by 5–10 V electrical stimulation of the appropriate segmental nerve using a suction electrode filled with bath solution. Data acquisition and analysis were performed using pClamp 10 (Molecular Devices). A Student's *t* test was used for statistical comparison of *syt*^{null} and *P*[*syt*^{B-IE}] mutants.

Electron microscopy

Stage 17 embryos were dissected in ice-cold, Ca²⁺-free HL3 and transferred immediately to ice-cold primary fixative (1% acrolein and 2.5% glutaraldehyde in 0.1 M cacodylate buffer) for 1 h. They were postfixed in a reduced osmium solution (0.5% osmium tetroxide and 0.8% potassium ferricyanide in 0.1 M

cacodylate buffer) for 1 h, incubated in 2% aqueous uranyl acetate overnight, embedded in 2% agar to facilitate handling, dehydrated, and embedded in Embed 812/Araldite (Electron Microscopy Sciences). Sections, at 70 nm, were grid stained in 2% aqueous uranyl acetate for 15 min, followed by Reynold's lead for 2 min. Random sections through the ventral nerve cord were examined at 30,000× magnification in the electron microscope.

Co-sedimentation assay

Co-sedimentation assays were conducted as described previously (Wang et al., 2003). Briefly, synaptotagmin C_2AB or C_2B domains (4 μ M final concentration) were incubated with increasing concentrations of protein free liposomes (PS/PC/PE, 15/55/30%) for 15 min at room temperature in the presence of either 1 mM Ca^{2+} or 0.2 mM EGTA. The mixtures were then centrifuged at 70,000 rpm for 1 h. The supernatant of each sample was collected, mixed with SDS loading buffer, and boiled for 5 min. Samples were subjected to SDS-PAGE and stained with Coomassie blue.

Co-flotation assay

As described previously (Gaffaney et al., 2008), PS-free t-SNARE vesicles were mixed with the indicated syt proteins ($10~\mu\mathrm{M}$ final concentration) in the presence of either 1 mm Ca $^{2+}$ or 0.2 mm EGTA at room temperature for 30 min. Samples were mixed with an equal volume of 80% Accudenz and transferred to centrifuge tubes (Beckman Instruments). Accudenz, at 35, 30, and 0%, was sequentially added to form step gradients. Samples were centrifuged at 55,000 rpm for 105 min, collected at the interface between the 30 and 0% Accudenz layers, and analyzed by SDS-PAGE.

Glutathione S-transferase—syt pull-down assay

Glutathione S-transferase (GST)-tagged syt proteins (C_2AB , 3 μ M; C_2B , 6 μ M) were incubated with Glutatione Sepharose beads (GE Healthcare) for 1 h in buffer (20 mm Tris, 150 mm NaCl, and 1% Triton X-100). The mixtures were washed with buffer and then incubated with t-SNARE protein (3 μ M) in buffer containing either EGTA (0.2 mM) or Ca (1 mM) for 30 min at room temperature. The samples were spun down and washed with buffer three times. The pellets were boiled in sample buffer, subjected to SDS-PAGE, and stained with Coomassie blue.

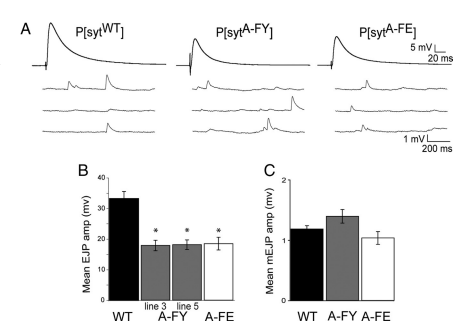


Figure 2. Evoked release is impaired in C_2A hydrophobic mutants. **A**, Representative voltage traces recorded from larval muscle fiber 6. Each top trace shows the mean of 30 consecutive evoked responses from the same muscle fiber. The bottom traces show 3 s of continuous spontaneous recordings. Line 3 shown for $P(syt^{A-FY})$. **B**, Compared with $P(syt^{WT})$, the mean EJP amplitude of $P(syt^{A-FY})$ lines 3 and 5, and $P(syt^{A-FE})$ were significantly decreased (*p < 0.005; $P(syt^{WT})$, n = 24; $P(syt^{A-FY})$ line 3, n = 20; $P(syt^{A-FY})$ line 5, n = 14; $P(syt^{A-FE})$, n = 10). No difference was found between $P(syt^{A-FY})$ and $P(syt^{A-FY})$ (p > 0.9). **C**, Compared with $P(syt^{WT})$, the mean mEJP amplitude of $P(syt^{A-FY})$ (line 3 shown) and $P(syt^{A-FE})$ showed no significant change (p > 0.05).

CD spectroscopy

CD spectra were measured with an AVIV stop flow Circular Dichroism spectropolarimeter at 192–260 nm using a 1 mm path-length cell. Samples containing 0.2 mg/ml of either wild-type or mutant C_2AB or C_2B domains were assayed at 25°C. For corrections of baseline noise, the signal from a blank run of buffer (50 mm sodium phosphate) was subtracted from all the experimental spectra.

Results

Each C_2 domain contains two highly conserved, hydrophobic residues around the rim of the Ca²⁺-binding pocket that penetrate the bilayer during Ca²⁺-dependent interactions between synaptotagmin and anionic membranes (Fig. 1 B, C_2A , F^* and M; C_2B , I^* and V) (Bai et al., 2002; Herrick et al., 2006). To determine whether membrane penetration by synaptotagmin is necessary for synaptic function, we mutated a single hydrophobic residue at the very tip of the Ca²⁺-binding pocket in either the C_2A or C_2B domain (Fig. 1, C_2A , F^* ; or C_2B , I^*) that has been shown to penetrate one-sixth of the way into membranes during Ca²⁺ binding (Bai et al., 2002; Herrick et al., 2006).

C₂A membrane penetration facilitates evoked release

In C_2A , we mutated the phenylalanine of loop 3 (Fig. 1, C_2A , F^*) to either tyrosine (syt A-FY) or glutamic acid (syt A-FE). The syt A-FY mutation maintains the general structure of the side chain but disrupts the hydrophobicity, whereas the negative charge of the syt A-FE mutation disrupts hydrophobicity and repels the head groups of anionic phospholipids. To indicate their transgenic origin, we will call the mutants $P[syt^{A-FY}]$ and $P[syt^{A-FE}]$ and the transgenic controls $P[syt^{WT}]$. Because these mutants were viable, all C_2A experiments were performed on third instars in the absence of the native protein.

Ca²⁺-evoked neurotransmitter release was decreased to 63% of control in both $P[syt^{A-FY}]$ and $P[syt^{A-FE}]$ mutants (Fig. 2A,B). Because each of the C₂A mutants are independent

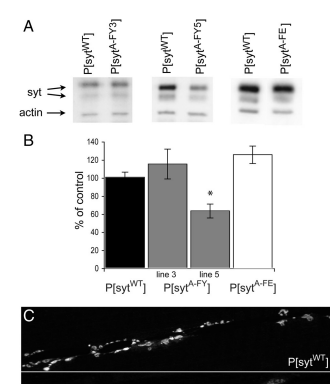


Figure 3. The C_2A hydrophobic mutations do not disrupt synaptotagmin expression or targeting. **A**, Representative Western blots from $P[syt^{A-FY}]$ line 3, line 5, and $P[syt^{A-FE}]$ and their $P[syt^{WT}]$ controls. **B**, $P[syt^{A-FY}]$ line 5 (n=17) expressed less transgenic protein than $P[syt^{WT}]$ (n=28) and the other two C_2A mutants ($P[syt^{A-FY}]$ line 3, n=7; $P[syt^{A-FE}]$, n=11; p<0.001). **C**, Transgenic synaptotagmin was localized to synaptic sites in all lines (line 5 shown for $P[syt^{A-FY}]$). Scale bar, 20 μ m.

P[syt^{A-FE-}

transgenic lines, the random insertion site of the transgene does not contribute to the decrease in evoked release. The finding that the mean amplitude of spontaneous release events (mEJP) is unchanged (Fig. 2C) demonstrates that the mutations do not disrupt either vesicle filling or responsiveness of the postsynaptic membrane. Protein misexpression also cannot account for the decrease in evoked release. Although one of the C₂A hydrophobic mutants had less transgene expression (Fig. 3 A, B), all three lines provided similar levels of synaptotagmin function during evoked transmitter release (Fig. 2B). In addition, no difference in synaptotagmin localization to synaptic sites was observed (Fig. 3C). Thus, the decrease in evoked transmitter release in these mutants is specific for the mutation. Interestingly, despite the difference in charge between tyrosine (polar but neutral) and glutamic acid (negatively charged), no difference in the amplitude of evoked release was observed between the $P[syt^{A-FY}]$ or $P[syt^{A-FE}]$ mutants (Fig. 2B), suggesting that disruption of penetration by C_2A into the hydrophobic core of the presynaptic membrane, regardless of electrostatic effects, causes the impairment of synaptic transmission.

Table 1. The C₂B hydrophobic mutant results in embryonic lethality

Line	Transgenic syt alone (%)	Native syt alone (%)	Native + transgenic syt (%)
$P[syt^{B-IE}]$			
Line 4	0	100	0
Line 6	0	100	0
P[syt ^{WT}]	34.1	34.8	31.1

Progeny are expressed as a percentage of the total larvae that hatched. Total larvae counted per genotype were as follows: $P[syt^{B-lE}]$ line 4, 265 larvae; line 6, 237 larvae; $P[syt^{WT}]$, 1178 larvae.

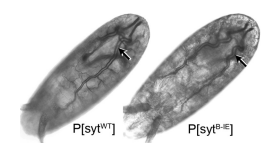


Figure 4. syt ^{B-IE} mutant embryos complete embryogenesis and exhibit gross morphology indistinguishable from wild-type embryos. Like controls, $P[syt^{B-IE}]$ mutants were able to develop through stage 17, as demonstrated by the development of trachea (arrows).

C₂B membrane penetration is essential for synaptotagmin function

To assess the physiological significance of the homologous hydrophobic residue in C₂B, we mutated the isoleucine of loop three (Fig. 1, C₂B, I*) to glutamic acid (syt ^{B-IE}). This introduction of negative charge should both prevent Ca²⁺-dependent membrane penetration and repel the head groups of the anionic phospholipids. Expression of the syt B-IE protein in the absence of native synaptotagmin resulted in embryonic lethality (Table 1, compare $P[syt^{B-IE}]$ to $P[syt^{WT}]$, transgenic syt alone). The embryos complete embryogenesis (Fig. 4) but fail to hatch. When manually removed from the egg case, the embryos are virtually immobile. There are microscopic movements within the body wall musculature, but these generally are insufficient to displace the embryo and all visible movement stops within hours (our unpublished observation). The only other synaptotagmin mutation to date that causes embryonic lethality is the $P[syt^{B-D1,2N}]$ mutant in which Ca2+ binding by the C2B domain has been blocked by mutating two of the aspartate residues (Fig. 1B, C₂B, D₁ and D₂) required for coordinating calcium (Mackler et al., 2002). Thus, this single hydrophobic residue at the tip of the C_2B Ca²⁺-binding pocket is the first putative effector residue to be as critical to the organism as the C₂B Ca²⁺-binding residues, suggesting that membrane penetration by the C₂B domain is required in vivo.

The syt $^{B-1E}$ mutation directly causes the embryonic lethality without disrupting synaptic vesicle biogenesis. Because lethality was observed in multiple independent lines (Table 1, $P[syt^{B-IE}]$ line 4 and line 6, transgenic syt alone), it cannot be caused by the random insertion of the mutant transgene. Similar levels of transgenic protein are expressed in mutant and control embryos (Fig. 5A, $P[syt^{WT}]$ vs $P[syt^{B-IE}]$, line 6 shown), so the deficit is not attributable to insufficient transgene expression. Although the absence of synaptotagmin results in a dramatic depletion of synaptic vesicles (Jorgensen et al., 1995; Reist et al., 1998; Loewen et al., 2006a), the syt $^{B-IE}$ mutation does not. Ultrastructural examination reveals abundant small clear vesicles within the synaptic neuropil (Fig. 5B). In addition, immunolabeling of the mutant embryos for a synaptic vesicle marker, CSP (Mastrogiacomo et

al., 1994), demonstrates that the overall population of synaptic vesicles in the neuropil is correctly localized to synaptic regions (Fig. 5C), indicating that this mutation does not disrupt the formation or targeting of synaptic vesicles. Finally, the syt B-IE mutant protein is also appropriately localized to synaptic sites. In the central nervous system, syt B-IE staining is appropriately localized to the synaptic neuropil (Fig. 5D). In the peripheral nervous system, syt B-IE staining is appropriately localized to the neuromuscular junction (Fig. 5E), demonstrating that this mutation does not disrupt synaptotagmin targeting. Indeed, because syt^{null} mutants can survive through to adulthood (Loewen et al., 2001), the embryonic lethality demonstrates that the syt B-IE transgene is expressed and is dominantly inhibiting function.

The nearly complete absence of locomotion in the syt $^{\rm B-IE}$ mutants indicates that the lethality results from an inability to move out of the embryonic egg casing. However, this is an indirect assessment of synaptic activity. To directly test the impact of the syt B-IE mutation on synaptic transmission, excitatory junction currents (EJCs) were recorded from $P[syt^{B-IE}]$ mutant, syt^{null} mutant, and $P[syt^{WT}]$ transgenic control embryos. Our recordings are not directly comparable with previous reports (Broadie et al., 1994; Mackler and Reist, 2001; Yoshihara and Littleton, 2002) because of differences in the positive control $(+/+ \text{ vs } -/-;P[syt^{WT}])$ and/or saline (differing concentrations of

divalent cations) used. As shown in Figure 6, the amplitude of evoked transmitter release in $P[syt^{B-IE}]$ mutants was 17.5% of the transgenic $P[syt^{WT}]$ control. However, most importantly, expression of the syt B-IE protein was worse than no synaptotagmin I expression at all; the EJC amplitude in $P[syt^{B-IE}]$ mutants was only 50% of that seen in syt^{null} mutants (Fig. 6) (p < 0.02). Thus, the syt B-IE mutant is the only effector interaction mutation to date that results in a more severe inhibition of synaptic transmission than the syt^{null} mutation. The only other synaptotagmin mutants to exhibit such an effect are C_2B Ca^{2+} -binding motif mutants (Mackler et al., 2002).

Proper folding of the syt $^{B-IE}$ mutant protein was verified by CD spectroscopy. Figure 7A demonstrates that this mutation does not alter the spectra for either the C_2AB (left panel) or the C_2B (right panel) domains. But does mutation of this C_2B hydrophobic residue primarily block interactions with negatively charged membranes or does it inhibit interactions with SNAREs? Mutation of all four of the hydrophobic residues that penetrate membranes (Fig. 1 B, C_2A , F^* ; M, C_2B , I^* and V, mutated to alanines; C_2AB^{4A}) dramatically inhibited interactions with negatively charged phospholipids (Lynch et al., 2008). However, the effect of this quadruple mutation on C_2AB interactions with SNARE proteins has been inconsistent. In one study, C_2AB binding to t-SNAREs was not effected by the C_2AB^{4A} mutation (Lynch et al., 2008), indicating that even this quadruple mutation was properly folded and specific for Ca^{2+} -dependent

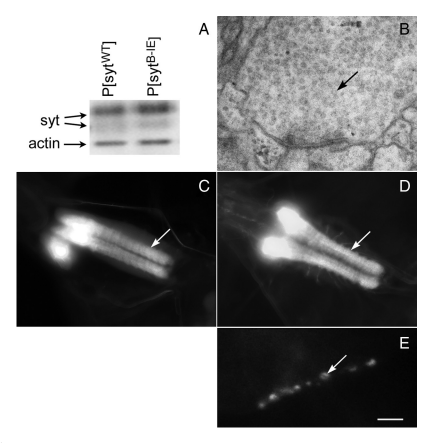


Figure 5. The C₂B hydrophobic mutation does not disrupt synaptic vesicle localization, synaptotagmin expression, or synaptotagmin targeting. **A**, A representative Western blot of embryos shows similar transgene expression levels for P[syt^{WT}] and P[syt^{B-IE}]. Blot was probed with an anti-actin antibody to confirm equal loading. **B**, A representative nerve terminal within the neuropil of a P[syt^{B-IE}] mutant ventral nerve cord exhibits abundant small clear vesicles (black arrow). **C**, Anti-CSP labeling was localized to the neuropil of the CNS in P[syt^{B-IE}] mutants demonstrating that synaptic vesicles are appropriately localized to synaptic regions. The white arrow indicates the neuropil of the ventral nerve cord; brain ganglia are out of the plane of focus. The mutant synaptotagmin was also correctly localized to the neuropil (**D**, white arrow) in the CNS, as well as to neuromuscular junctions (**E**, white arrow) in the peripheral nervous system. P[syt^{B-IE}] line 6 is shown. Scale bar: **B**, 180 nm; **C**, **D**, 40 μm; **E**, 5 μm.

phospholipid interactions. However, in another study, this same quadruple mutation resulted in a partial decrease in binding to t-SNAREs (Hui et al., 2009).

To determine the effector interactions disrupted by our single syt B-IE mutation, we conducted biochemical interaction assays using C₂AB or isolated C₂B domains. To determine the effect of the syt B-IE mutation on interactions with membranes, we measured the ability of protein-free, negatively charged liposomes (containing 15% PS) to bind C₂AB domains. The syt B-IE mutation caused a marked decrease in Ca²⁺-dependent interactions of C₂AB with PS liposomes, but significant binding remained (Fig. 7*B*). However, interactions with membranes may be mediated by both the C₂A and the C₂B domains (Bai et al., 2002; Shin et al., 2003; Rhee et al., 2005). To determine whether any membrane binding activity can be mediated by the mutated C₂B domain, we measured the ability of isolated C₂B domains to bind PS liposomes. As shown in Figure 7*B*, the syt ^{B-IE} mutation abolished Ca²⁺-dependent interactions with negatively charged liposomes. Thus, the complete block of membrane interactions by the C₂B domain could account for the severe inhibition of evoked release and lethality seen in these mutants.

To determine whether the syt ^{B-IE} mutation also impacted interactions with t-SNAREs in the absence of membranes, we measured the ability of GST–C₂AB or GST–C₂B fusion proteins to bind to full-length t-SNAREs solubilized in detergent (Fig. 8 *A*).

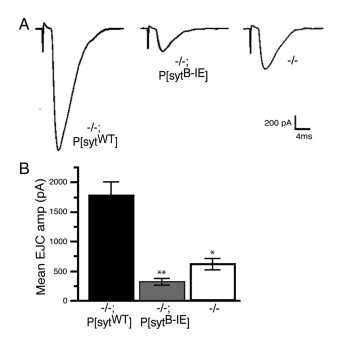


Figure 6. The C₂B hydrophobic mutation inhibits evoked transmitter release more severely than the syt^{null} mutation. **A**, Representative current traces recorded from embryonic muscle fiber 6. **B**, The mean EJC amplitude in transgenic control embryos was $1782 \pm 224 \,\mathrm{pA} \,(-/-; P[syt^{WT}], n=8)$, in $P[syt^{B-IE}]$ mutant embryos was $312.2 \pm 62.6 \,\mathrm{pA} \,(-/-; P[syt^{B-IE}], n=8)$, and in syt^{null} mutants was $625.2 \pm 89.0 \,\mathrm{pA} \,(-/-, n=11)$. Evoked release in $P[syt^{B-IE}]$ was significantly less than in syt^{null} mutants (* vs **, p<0.02).

The syt ^{B-IE} mutation decreased interactions with t-SNAREs, especially in the absence of Ca ²⁺. Ca ²⁺ increased t-SNARE binding to C₂AB domains by 0.2 units or 2.4-fold in wild type and by 0.112 units or 4.5-fold in syt ^{B-IE} mutants. When isolated C₂B domains were used, Ca ²⁺ increased t-SNARE binding by 0.116 units or 2.5-fold in wild type and by 0.105 units or 4.9-fold in syt ^{B-IE} mutants. The effect on Ca ²⁺-dependent t-SNARE binding to C₂B is minimal, yet the inhibition of Ca ²⁺-dependent phospholipid binding to C₂B is complete. Thus, the lethality seen in these mutants correlates best with the disruption in phospholipid interactions. However, a contribution attributable to altered t-SNARE binding cannot be definitively excluded. In addition, the finding that the Ca ²⁺-dependent increase in t-SNARE binding is essentially intact in the C₂B hydrophobic mutant suggests that, if this residue mediates any direct interaction with t-SNAREs (but see below), it likely occurs during the docking or priming of vesicles before Ca ²⁺ influx.

Interestingly, the Ca²⁺-dependent interaction between C₂B and membrane-embedded t-SNAREs requires this hydrophobic residue in C2B. When t-SNAREs are embedded in net neutral liposomes [PC/PE to prevent direct electrostatic interactions with synaptotagmin (Gaffaney et al., 2008)], Ca²⁺ caused an approximate twofold increase in binding (Fig. 8 B, C₂B; C, C₂AB). The syt B-IE mutation partially blocked the C₂AB (Fig. 8C, C₂AB, I420E) and completely blocked the C₂B Ca²⁺-dependent t-SNARE/ membrane interaction (Fig. 8B, C₂B, I420E), similar to the finding for binding to PS-containing liposomes (Fig. 7B). Thus, the presence of membrane altered the interactions with t-SNAREs. The finding that Ca²⁺-dependent t-SNARE interactions mediated by C₂B were intact in the mutants in the absence of membrane but are abolished in the presence of membrane suggests that the interaction shown in Figure 8B is actually membrane dependent. In addition, because the Ca²⁺-independent interaction with t-SNAREs that are embedded in membranes (as they are *in vivo*) is intact in the mutant (Fig. 8 *B*, *C*), this hydrophobic residue is unlikely to mediate a direct interaction with t-SNAREs during docking or priming. The simplest interpretation is that this t-SNARE interaction may require membrane penetration by the tip of the C_2B domain. Together, the biochemistry supports the hypothesis that Ca^{2+} -dependent membrane penetration by this hydrophobic residue on the tip of the C_2B domain is the essential effector interaction that couples Ca^{2+} binding by synaptotagmin with vesicle fusion.

Expression of the $P[syt^{B-IE}]$ transgene has a dominant-negative effect

To analyze the inhibitory effect of the syt B-IE protein, we expressed the transgenic mutant protein in the presence of native synaptotagmin. When transgene expression rescues synaptotagmin function, the hatching progeny are approximately evenly distributed between the "transgenic syt alone," "native syt alone," and "native + transgenic syt" genotypes (see Table 1, $P[syt^{WT}]$ and methods). If transgene expression did not enhance viability in the absence of native synaptotagmin yet did not impact the function of native synaptotagmin, we would expect few progeny in the transgenic syt alone genotype [syt^{null} mutants are weakly viable (Loewen et al., 2001)] and an approximately equal distribution of progeny between the native syt alone and native + transgenic syt genotypes (i.e., no dominant-negative effect). When more progeny are able to hatch of the native syt alone genotype than of the native + transgenic syt genotype (Table 1, $P[syt^{B-IE}]$), this indicates that the transgene is dominantly inhibiting the function of the native protein. Remarkably, expression of syt B-IE mutant protein was embryonic lethal even in the presence of native synaptotagmin (Table 1, compare $P[syt^{B-\hat{IE}}]$ to $P[syt^{WT}]$, native + transgenic syt). Previously, the only synaptotagmin mutations documented to result in a dominant-negative effect were tandem point mutations within the C_2B Ca^{2+} -binding motif [syt $^{B-D1,2N}$ and syt $^{B-D3,4N}$ (Mackler et al., 2002)]. However, in the presence of wild-type synaptotagmin, some of these tandem mutants were able to hatch and remained viable through the third-instar stage, although they did impair synaptic transmission. Thus, mutation of the hydrophobic residue at the tip of the C₂B Ca²⁺-binding pocket is the most severe mutation of synaptotagmin function to date.

Discussion

In our model for synaptotagmin function (Fig. 9), we propose that presynaptic membrane penetration by C₂B is the critical effector interaction for coupling Ca²⁺ binding by synaptotagmin with evoked transmitter release at the synapse. In agreement with models of SNARE-mediated fusion (Weber et al., 1998; Lang and Jahn, 2008; Rizo and Rosenmund, 2008), we propose that multiple synaptotagmin/SNARE complexes are localized in a ring around the site of synaptic vesicle/presynaptic membrane apposition (Fig. 9A, D). Before Ca²⁺ influx, interactions between the C₂B polylysine motif (Fig. 9B, yellow space-filled residues) and SNARE complexes help hold the vesicle in a docked/primed state (Rickman et al., 2004; Loewen et al., 2006b), whereas the negative charge of the Ca^{2+} -binding pockets (Fig. 9A, single – signs; B, cluster of - signs) repulses the negatively charged presynaptic membrane. During Ca²⁺ binding, the negative charge of the pockets is neutralized, which initiates the electrostatic switch (Davletov et al., 1998; Ubach et al., 1998; Zhang et al., 2002): a dramatic shift in the electrostatic potential of each C₂ domain. This increase in positive charge could strengthen (Loewen et al., 2006b), or initiate new (Lynch et al., 2007), interactions with

negatively charged regions of the SNAREs and permit an electrostatic interaction between the positively charged residues at the tips of the Ca²⁺-binding pockets (Fig. 9, blue space-filled residues, +) and the negatively charged presynaptic membrane (Fernández-Chacón et al., 2001; Wang et al., 2003; Paddock et al., 2008). Ca²⁺ binding would also permit the hydrophobic residues at the tips of each C2 domain (Fig. 9, gray space-filled residues) to penetrate into the hydrophobic core of the presynaptic membrane (Bai et al., 2002; Herrick et al., 2006), thereby inducing positive curvature in this membrane (Fig. 9C) (Martens et al., 2007; Hui et al., 2009). Membrane penetration would assist SNARE-mediated fusion by destabilizing the presynaptic membrane and bulging it toward the vesicle (Fig. 9D, large gray ovals, arrows) in a ring around the SNARE complex transmembrane domains (Fig. 9D, small gray circles, arrowheads).

Our model can account for the effects of synaptotagmin mutations seen at synapses. First, we will consider mutations that do not have a dominant inhibitory effect. Mutation of the C₂B polylysine motif (Fig. 9B, C, yellow space-filled residues) results in only a ~50% decrease in evoked transmitter release (Mackler and Reist, 2001; Li et al., 2006). Thus, the synaptic vesicle docking/ priming function of this motif facilitates release but is not essential; the Ca²⁺-binding pockets remain intact and can still penetrate the presynaptic membrane. Mutation of the positively charged residue at the tip of each Ca^{2+} -binding pocket (Figs. 1, \oplus ; 9, blue space-filled residues, +) results in a $\sim 50-$ 80% decrease of evoked transmitter release (Fernández-Chacón et al., 2001; Wang et al., 2003; Paddock et al., 2008). This decrease in function would result from a decrease in the electrostatic attraction of the presynaptic membrane, likely leading to the hydrophobic residues penetrating the membrane less deeply (Herrick et al., 2006). Mutation of the C₂A hydrophobic residue, syt^{A-FY} or syt A-FE, results in a ~50% decrease in evoked transmitter release (Fig. 2B), indicating that membrane penetration by the C₂A domain improves the efficacy of Ca²⁺triggered vesicle fusion but, like the other mutants discussed above, it is not required.

Mutations that disrupt interactions with the SNARE complex provide variable results. A mutation that impairs both ${\rm Ca}^{2+}$ -dependent and ${\rm Ca}^{2+}$ -independent SNARE interactions (R398A) by $\sim 50\%$ also impairs the ability of the C₂AB domain to stimulate SNARE-mediated liposome fusion *in vitro* by $\sim 35\%$ (Gaffaney et al., 2008), indicating that this synaptotagmin interaction with the SNARE com-

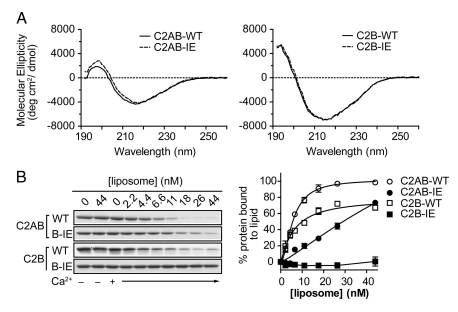


Figure 7. The syt ^{B-IE} mutation abolishes Ca ²⁺ -dependent, C₂B interactions with membranes. **A**, C₂AB or C₂B domains with the syt ^{B-IE} mutation are correctly folded. The CD spectra of the mutant domains (C₂AB-IE and C₂B-IE) did not show significant changes compared with wild type (C₂AB-WT and C₂B-WT, respectively). **B**, The syt ^{B-IE} mutation abolishes the ability of the C₂B domain to bind negatively charged liposomes. Left, A representative co-sedimentation assay; right, graph of three experiments. In the absence of Ca ²⁺, C₂AB and C₂B domains do not bind to liposomes containing 15% PS and thus remain in solution (left, —, WT and B-IE). In the presence of 1 mm Ca ²⁺, increasing concentrations of liposomes bind increasing amounts of control C₂AB or C₂B domains (C₂AB-WT and C₂B-WT). The syt ^{B-IE} mutation impairs this interaction with C₂AB (C₂AB-IE) but abolishes the interaction with the isolated C₃B domain (C,B-IE).

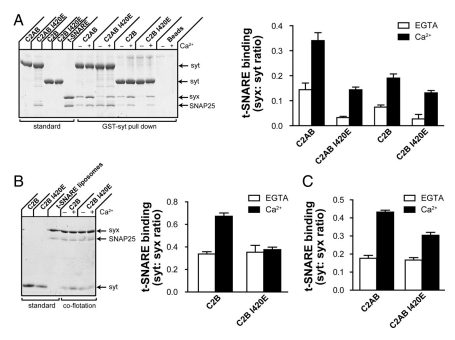


Figure 8. Direct, Ca $^{2+}$ -dependent C₂B interactions with t-SNAREs are intact in the syt $^{B-IE}$ mutation. **A**, The syt $^{B-IE}$ mutation does not inhibit Ca $^{2+}$ -dependent interactions with solubilized t-SNAREs. Left, A representative GST-syt pull-down assay; right, histogram of three experiments. The Ca $^{2+}$ -dependent increase in t-SNARE interactions (black bars relative to white bars) is not disrupted by the syt $^{B-IE}$ mutation. Ca $^{2+}$ -independent t-SNARE interactions are decreased (white bars, C₂AB vs C₂AB 1420E and C₂B vs C₂B 1420E). **B**, The syt $^{B-IE}$ mutation abolishes Ca $^{2+}$ -dependent C₂B interactions with membrane-embedded t-SNAREs. Left, A representative co-floatation assay; right, histogram of four experiments. The syt $^{B-IE}$ mutation does not alter the amount of C₂B domain that binds to PS-free, t-SNARE vesicles (t-SNARE) in the absence of Ca $^{2+}$ (-, left; EGTA, right). However, the syt $^{B-IE}$ mutation abolishes the increase in t-SNARE vesicles binding induced by 1 mm Ca $^{2+}$ (+, left; Ca $^{2+}$, right; p < 0.05). **C**, The syt $^{B-IE}$ mutation impairs C₂AB interactions with membrane-embedded t-SNAREs. Again, the syt $^{B-IE}$ mutation does not alter the amount of C₂AB domain that binds to PS-free, t-SNARE vesicles in the absence of Ca $^{2+}$ (EGTA, white bars). In the presence of 1 mm Ca $^{2+}$ (Ca $^{2+}$, black bars), the t-SNARE vesicles bind 70% of wild-type levels of C₂AB domain when the syt $^{B-IE}$ mutation is present (p < 0.001; n = 6).

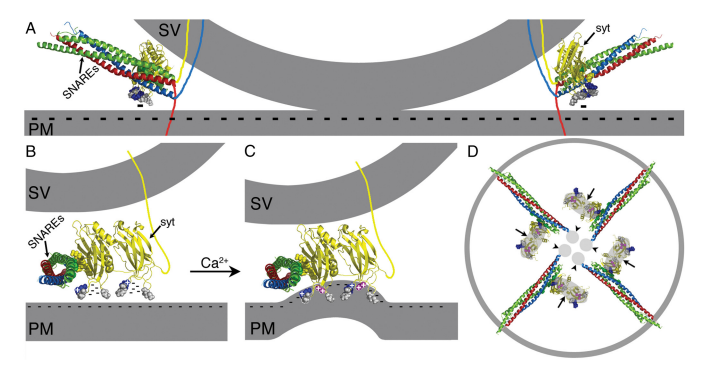


Figure 9. Model of the role of Ca²⁺-dependent penetration of the anionic presynaptic membrane by synaptotagmin. The crystal structure of the core complex [Protein Data Bank (PDB) file 1SFC, containing syntaxin (red), SNAP-25 (synapse associated protein of 25 kD) (green), and VAMP (vesicle-associated membrane protein)/synaptobrevin (blue)], the nuclear magnetic resonance structures of the C₂A (PDB file 1BYN) and C₂B (PDB file 1K5W) domains of synaptotagmin (yellow), and Ca²⁺ (pink "+") were rendered using PyMOL Molecular Graphics System (DeLano Scientific). The synaptic vesicle (SV), the presynaptic membrane (PM), the transmembrane domains, and the link between C₂A and C₂B were added in Adobe Photoshop. *A*, Docked synaptic vesicle with two synaptotagmin/SNARE complexes shown. *B*, syt/SNARE complex viewed end on from the site of synaptic vesicle/presynaptic membrane apposition (SNARE complex transmembrane domains in front of plane of section). Ca²⁺-independent priming between the C₂B polylysine motif (yellow, space-filled residues) and SNAP-25 [green, space-filled residues (Zhang et al., 2002; Rickman et al., 2004; Loewen et al., 2006b)] holds the C₂B Ca²⁺-binding site immediately adjacent to the SNARE complex and the C₂A Ca²⁺-binding site nearby. The negative charge of the Ca²⁺-binding pockets (cluster of — symbols) prevents interactions between the tips of the C₂ domains and the presynaptic membrane attributable to electrostatic repulsion. *C*, Ca²⁺ binding neutralizes the negative charge of the pockets resulting in a strong attraction of the negatively charged, phospholipid head groups of the presynaptic membrane by the bound Ca²⁺ (pink spheres +) and the basic residues at the tips of Ca²⁺-binding pockets (blue, space-filled residues) into the core of the presynaptic membrane and then triggers fusion by promoting a local Ca²⁺-dependent buckling of the plasma membrane (Martens et al., 2007; Hui et al., 2009). *D*, syt/SNARE complexes (small gray circles, arrowheads).

plex facilitates vesicle fusion. Another mutation that abolishes Ca²⁺-dependent SNARE interactions [R199A,K200A,K297A, K301A (Lynch et al., 2007)] showed no effect on the ability of the Ca²⁺-bound C₂AB domain to stimulate SNARE-mediated liposome fusion in vitro by one group (Gaffaney et al., 2008), indicating that this Ca²⁺-dependent SNARE interaction may not play a role. Yet another group found that this same mutation, when Ca²⁺ bound, supports a level of SNARE-mediated liposome fusion that is only halfway between Ca²⁺-bound, wild-type C₂AB and Ca²⁺-free, wild-type C₂AB (Lynch et al., 2007), indicating that this interaction provides a facilitatory effect similar to the interactions listed above. This mutant failed to rescue large dense-core vesicle (LDCV) fusion in PC12 cells, suggesting that the SNARE interaction plays a more important role in this system (Lynch et al., 2007). However, the properties of LDCV fusion are different from those of synaptic vesicle fusion in terms of timing and Ca2+ sensitivity (Kasai, 1999). Importantly, in PC12 cells, introduction of synaptotagmin I that contains aspartate to asparagine mutations in the C₂A Ca²⁺-binding motif causes a 60% decrease in evoked release (Wang et al., 2006), whereas the same mutations do not inhibit evoked release at synapses (Fernández-Chacón et al., 2002; Robinson et al., 2002). In addition, the positively charged residues at the tips of the C2 domains (Fig. 9, blue space-filled residues, +) are equally important during LDCV fusion [mutation of either results in a ${\sim}50\%$ decrease in fusion (Wang et al., 2003)], whereas this residue in C_2B is more critical than in C_2A for synaptic vesicle fusion at an intact synapse [80% inhibition in C_2B mutant and 50% inhibition in C_2A mutant (Paddock et al., 2008)]. These discrepancies indicate a shift in the relative role of the C_2A and C_2B domains between these systems with both C_2 domains participating in the Ca^{2+} -sensing function during LDCV fusion in PC12 cells, whereas C_2B fulfills the more critical role at the synapse. Although interactions with the SNARE complex are clearly important in facilitating vesicle fusion, whether they are required for fusion at the synapse has not been tested.

Synaptotagmin mutants that block ${\rm Ca}^{2+}$ -dependent membrane penetration would have a dominant-negative effect (i.e., inhibit the action of wild-type synaptotagmin) because multiple synaptotagmin:SNARE complexes are required to fuse a single synaptic vesicle (Fig. 9A,D) (Jahn and Scheller, 2006). As these nonfunctional mutants displace native synaptotagmin, synaptic efficacy declines. Our ${\rm C}_2{\rm B}$ hydrophobic mutant and the ${\rm C}_2{\rm B}$ ${\rm Ca}^{2+}$ -binding motif mutants fall into this category. Our finding that expression of the syt ${\rm ^{B-IE}}$ mutant results in embryonic lethality in the presence of native synaptotagmin (Table 1) indicates that disruption of ${\rm C}_2{\rm B}$ membrane penetration of even a fraction of the synaptotagmin molecules in the synaptotagmin:SNARE

complexes mediating fusion is lethal. However, C2B Ca2+binding motif mutants, which also disrupt Ca²⁺-dependent interactions with negatively charged phospholipids, reduce evoked transmitter release and only impair viability when coexpressed with native synaptotagmin (Mackler et al., 2002). This apparent discrepancy may be explained by the nature of the mutations of the C₂B Ca²⁺-binding motif. Aspartate to asparagine mutations (DtoN) would partially neutralize the negative charge of the Ca²⁺-binding pocket similar to the action of Ca²⁺. Indeed, syt DtoN mutations in C₂A partially mimic Ca²⁺ binding (Stevens and Sullivan, 2003). Because the positively charged residue at the tip of the pocket (Fig. 9, blue space-filled residue, +) remains intact in syt B-DtoN mutants to attract the presynaptic membrane, the hydrophobic residues at the tip of the C₂B Ca²⁺-binding pocket may be able to insert into the membrane, at least partially. Thus, syt B-DtoN mutations, by permitting partial membrane insertion, would have a less severe dominant-negative effect on native synaptotagmin than the syt B-IE mutation.

In summary, the syt^{B-IE} mutation is the first effector interaction mutation to inhibit evoked release below levels seen in syt^{mull} mutants, similar to the C_2B Ca^{2+} -binding motif mutations. The lethality exhibited in the $P[syt^{B-IE}]$ mutants, even in the presence of native synaptotagmin, demonstrates that mutation of a single hydrophobic residue in C_2B results in the most severe disruption of synaptotagmin function to date. These findings strongly support the hypothesis that membrane penetration by the C_2B domain of synaptotagmin is the key effector interaction required for triggering Ca^{2+} -dependent synaptic vesicle fusion.

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