Lipid-anchored Synaptobrevin Provides Little or No Support for Exocytosis or Liposome Fusion*

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SNARE proteins catalyze many forms of biological membrane fusion, including Ca\(^{2+}\)-triggered exocytosis. Although fusion mediated by SNAREs generally involves proteins anchored to each fusing membrane by a transmembrane domain (TMD), the role of TMDs remains unclear, and previous studies diverge on whether SNAREs can drive fusion without a TMD. This issue is important because it relates to the question of the structure and composition of the initial fusion pore, as well as the question of whether SNAREs mediate fusion solely by creating close proximity between two membranes versus a more active role in transmitting force to the membrane to deform and reorganize lipid bilayer structure. To test the role of membrane attachment, we generated four variants of the synaptic v-SNARE synaptobrevin-2 (syb2) anchored to the membrane by lipid instead of protein. These constructs were tested for functional efficacy in three different systems as follows: Ca\(^{2+}\)-triggered dense core vesicle exocytosis, spontaneous synaptic vesicle exocytosis, and Ca\(^{2+}\)-synaptotagmin-enhanced SNARE-mediated liposome fusion. Lipid-anchoring motifs harboring one or two lipid acylation sites completely failed to support fusion in any of these assays. Only the lipid-anchoring motif from cysteine string protein-α, which harbors many lipid acylation sites, provided support for fusion but at levels well below that achieved with wild type syb2. Thus, lipid-anchored syb2 provides little or no support for exocytosis, and anchoring syb2 to a membrane by a TMD greatly improves its function. The low activity seen with syb2-cysteine string protein-α may reflect a slower alternative mode of SNARE-mediated membrane fusion.

Nearly all biological fusion mediated by SNARE proteins employs at least one t-SNARE and one v-SNARE anchored to the membrane by a transmembrane domain (TMD)\(^3\) (1–3). Ca\(^{2+}\)-triggered exocytosis of neurotransmitter and hormone from many cells requires the vesicle-SNARE synaptobrevin 2 (syb2) and plasma membrane-SNARE syntaxin, both of which have a TMD (4, 5). Mutations in the TMD alter flux through exocytotic fusion pores in a manner consistent with structural models of fusion pores formed by the TMDs of syntaxin (6, 7) and syb2 (8). Yeast t- and v-SNAREs anchored to membranes by geranylgeranyl moieties instead of a TMD support docking but not fusion (9). SNAREs anchored to membranes by lipid instead of a TMD support lipid mixing of liposomes only when the lipid moiety is long enough to span a lipid bilayer or multiple lipid moieties are present (10) or an accessory protein is present (11).

Although early work supported a role of SNARE TMDs in membrane fusion, this issue has become controversial. Recent work on fusion of yeast vacuoles suggests that some SNAREs with lipid anchors can support lipid mixing, content mixing, or both, but other SNAREs cannot function when their TMD is replaced by a lipid anchor (12). Lipid-anchored forms of syntaxin and syb2 were reported to rescue neurotransmitter release at synapses as effectively as wild type (WT) SNAREs (13). This uncertainty about the role of membrane attachment raises important questions about the mechanism of membrane fusion. Many models of membrane fusion posit an active role for the TMD, beyond simply tethering the SNARE motif to the membrane. The initial fusion pore could not be proteinaceous if the TMD can be dispensed with (14, 15), and the transmission of force from the SNARE complex to the membrane is difficult to conceptualize without a TMD (16–19).

In this study, we sought to test the role of membrane attachment in the function of syb2 in diverse milieus. Four different lipid-anchored variants of syb2 were generated and assayed in Ca\(^{2+}\)-triggered release of catecholamine from chromaffin cells, spontaneous release of excitatory neurotransmitter from synapses, and Ca\(^{2+}\)-triggered fusion of reconstituted proteoliposomes. These lipid-anchored SNAREs supported little or no fusion above baseline levels. Only one construct harboring a cysteine string protein-α (CSPα) motif that can accept many acyl moieties supported a level of fusion significantly above background. None of the four lipid-anchored SNARE constructs supported fusion nearly as well as WT syb2. This work indicates that the form of membrane attachment of syb2 plays an important role in membrane fusion in synapses, endocrine cells, and proteoliposomes. A TMD may provide better support for fusion by forming a nascent fusion pore more readily or by more efficient force transmission to the lipid bilayer.

Experimental Procedures

Synaptobrevin 2 and Cellubrevin Double Knock-out Chromaffin Cells and Hippocampal Neurons—Syb2/cellubrevin double knock-out (DKO) mice were obtained by crossing syb2...
heterozygous knock-out mice with cellubrevin homozygous knock-out mice (20). Litters were one-quarter DKO (expressing neither syb2 nor cellubrevin), whereas the remaining three-quarters, taken as littermate controls, had one or two WT syb2 alleles and no cellubrevin. Chromaffin cells were cultured from E17.5–18.5 DKO embryos; individual embryos were genotyped and cultured in separate dishes.

Hippocampi from DKO E17.5–18.5 embryos were dissected in DMEM with 5 mm HEPES, dissociated in 0.05% trypsin for 30 min at 37 °C, and homogenized by rapid pipetting with 1-ml pipette tips. Dissociated neurons were then plated on 12-mm coverslips coated with 0.2 mg/ml poly-l-lysine. Hippocampal neurons were cultured in medium containing Neurobasal-A, Glutamax, penicillin/streptomycin, and B-27.

**Molecular Biology, Viral Infection, and Protein Purification—** Mouse syb2 constructs were amplified by PCR adding XhoI 5' and EcoRI 3' motifs were added to the 3' end primer right after residue 94. The construct containing residues 1–92 of syb2 and the C-terminal domain of CSPα (Syb2ATMR1# (13)) was provided by T. Südhof and used as the template for PCR. The PCR fragments of syb2 constructs were digested with XhoI and EcoRI and ligated into the lentiviral vector pLOX-CMV-EGFP to make EGFP-syb2 fusion constructs in which the sequence encoding EGFP was fused to the N terminus of syb2 by the linker sequence SGLRSRG. WT syb2, ΔTMD-CSPα, and ΔTMD-CIIIL coding sequences without an N-terminal EGFP tag were subcloned into a bicistronic lentiviral vector pLox-CMV-ires2-EGFP (modified from Ref. 21). To create syb2-RST-mVenus, a megaprimer containing the sequence of RST-mVenus and junction regions targeting to the vector pLox-CMV-syb2 was first amplified and then used for amplification and generation of pLox-CMV-syb2-RST-mVenus. All constructs were confirmed by sequencing.

Lentiviral particles were generated in HEK 293T cells as described previously (21). Chromaffin cells were infected by viral particles on the day of dissection, and hippocampal neurons were infected on the 7th day. Viral particles on the day of dissection, and hippocampal neurons were infected on the 7th day of dissection, and hippocampal neurons were infected on the 7th day of dissection, and hippocampal neurons were infected on the 7th day.

Bacterial expression vectors to generate full-length syb2 and its cytoplasmic domain were provided by J. E. Rothman. The cDNAs to generate full-length t-SNARE heterodimers were subcloned into pRSFduet as described previously (22). The cDNAs for ΔTMD-LC, ΔTMD-CIIIL, ΔTMD-CAC, and ΔTMD-CSPα described above were subcloned into pTrcHis for bacterial expression and Ni2+–Sepharose affinity purification. All proteins were expressed in *Escherichia coli* as His-tagged fusion proteins and purified as described previously (23).

**Amperometry Recording and Analysis—** To detect catecholamine release from chromaffin cells, a carbon fiber electrode (ALA Scientific Instruments, Westbury NY) was positioned at the surface of a cell under visual control and polarized to 650 mV. Amperometry current was recorded with a VA-10 amplifier (ALA Scientific Instruments). Chromaffin cells 1–3 days *in vitro* were bathed in a solution composed of (in mM) 150 NaCl, 4.2 KCl, 1 NaH2PO4, 0.7 MgCl2, 2 CaCl2, 10 HEPES, pH 7.4, at 22 °C. Transfected cells were visualized by EGFP fluorescence, and secretion was triggered by ejecting depolarizing solution (in mM: 105 KCl, 5 NaCl, 1 NaH2PO4, 0.7 MgCl2, 2 CaCl2, 10 HEPES, pH 7.4) from an ~2-μm tipped glass pipette positioned 10–15 μm from the cell. Current was recorded in 23-s episodes, beginning with a 3-s baseline, a 6-s period of depolarization, and 14 s after the end of depolarization. Secretion cells were stimulated up to six times. Spike frequency was determined for each secreting cell as the mean number of events with amplitudes ≥4 pA in the first episode, divided by 20 s. To construct cumulative spike plots, the events following depolarization in the first episode from all cells expressing a given construct were pooled, binned in 1-s intervals, and divided by cell number. The initial segment of each cumulative plot between 10 and 70% of the plateau was fitted by linear regression to obtain the 10–70% rising slope. One-way analysis of variance was applied to frequency measurements (Fig. 2B) and slopes (Fig. 2D), and with p < 0.05, the Tukey post-test was used to identify mutants that differed from WT syb2 or DKO. Bar graphs display mean ± S.E.

**Patch Clamp Recording and Analysis—** Cultured hippocampal neurons were bathed in a solution composed of (in mM) 140 NaCl, 5 KCl, 5 CaCl2, 1 MgCl2, 10 HEPES, and 10 glucose, with pH adjusted to 7.4 by NaOH. The bathing solution also contained 0.5 μM tetrodotoxin, 10 μM SR95531, and 50 μM amino phosphonovalerate to isolate mEPSCs mediated by α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid-type glutamate receptors. Patch pipettes were filled with internal solution containing (in mM) 120 cesium gluconate, 8 CsCl, 2 NaCl, 10 EGTA, 5 HEPES, 5 phosphocreatine, 2 Mg-ATP, and 0.3 Na-GTP, with pH adjusted to 7.2 by CsOH. Patch pipettes containing this solution had resistances of 3–8 mehoms. All recordings were performed at room temperature. The voltage was held at −70 mV with an EPC-7 patch clamp amplifier (HEKA). Current was filtered at 3 kHz, digitized at 50 kHz, and read into a computer with PClamp software. Frequency was the mean of the average number of events in each cell divided by time. Differences in mEPSC frequencies were evaluated with the Mann-Whitney *U* test. Bar graphs display mean ± S.E.

**Immunocytochemistry and Imaging—** DKO chromaffin cells were transfected with neuropeptide Y-DsRed (NPY-DsRed) and WT or mutant EGFP-syb2 constructs. Three days after transfection, cells were fixed with 4% paraformaldehyde for 20 min, permeabilized by 0.3% Triton X-100 for 10 min, blocked in 10% goat serum and 0.1% Triton X-100 for 10 min, and immunostained with mouse anti-syb2 antibody (104211, Synaptic Systems) and rabbit anti-RFP antibody (ab62341, Abcam) for 1 h. After washing three times with phosphate-buffered saline (PBS), cells were treated with Cy2-goat anti-mouse and Alexa Fluor 568-goat anti-rabbit secondary antibodies for 1 h, followed by three washes with PBS. All the above steps were carried out at room temperature. Images of immunostained chromaffin cells were acquired at room temperature on a Nikon Eclipse Ti confocal microscope with a ×100, 1.49 numerical aperture oil-immersion lens and Andor DV-897 camera. The laser intensity was set to 8% for both 488- and 561-nm illumination, and the exposure time was between 100 and 200 ms. Background subtraction was performed with ImageJ software. Pearson’s coefficient and total fluorescence intensity (area of cell–mean cell intensity − background) were quantified using.
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Nikon Eclipse Ti analysis software. One-way analysis of variance with Tukey post test was used for Pearson’s coefficients and total fluorescence intensity to evaluate the significance of differences between syb2 mutants and WT syb2 or syb2 mutants and ΔTMD-94.

Neurons on coverslips were fixed using 4% paraformaldehyde in PBS for 20 min. After washing with PBS, the neurons were permeabilized, blocked in blocking buffer containing 0.3% Triton X-100, 3% goat serum, and 2% BSA at room temperature for 1 h, and incubated at 4 °C overnight with polyclonal guinea pig anti-synaptophysin 1 (1:1000) (101004, Synaptic Systems) and monoclonal mouse anti-syb2 (1:500) (104211, Synaptic Systems). After rinsing with PBS, the coverslips were incubated at room temperature with Cy3-conjugated rabbit anti-guinea pig IgG (1:300) and Cy2-conjugated goat anti-mouse IgG (1:500) for 45 min. All antibodies were diluted with blocking buffer. Fluorescent signals were detected using an Olympus IX83 microscope with 60 × 1.49 numerical aperture oil-immersion objective and ORCA flash 4.0 camera. Images were acquired with MetaMorph software and analyzed with Fiji ImageJ software.

Reconstituted Proteoliposomes—Proteoliposomes, generated as described previously (23), were composed of 25% phosphatidylserine, 45% phosphatidylcholine, 30% phosphatidylethanolamine and harbored the t-SNAREs - syntaxin-1A and SNAP-25B. Briefly, lipids were dried under a stream of nitrogen gas while rotating. The lipid film was further lyophilized for at least 1 h to remove any residual organic solvent. The lipid film was solubilized with proteins in 25 mM HEPES-KOH, pH 7.4, 100 mM KCl, 10% glycerol, 1% octyl glucoside, 1 mM DTT. Vesicles were formed by rapid dilution of the octyl glucoside with 2 volumes of reconstitution buffer (25 mM HEPES-KOH, 100 mM KCl, 10% glycerol, 1 mM DTT) and overnight dialysis against 4 liters of reconstitution buffer. Proteoliposomes were collected by density gradient centrifugation and analyzed by SDS-PAGE and Coomassie Blue staining. Vesicles harboring full-length syb2 were prepared as described above; however, the phosphatidylethanolamine content was reduced from 30 to 27% to accommodate the addition of 1.5% rhodamine-phosphatidylethanolamine and 1.5% NBD-phosphatidylethanolamine to the lipid mixture.

Lipid-anchored Proteoliposomes—Syb2 proteins lacking their TMD were attached to liposomes via cysteine residue(s). Briefly, the syb2 proteins were dialyzed overnight at 4 °C against cross-linking buffer (25 mM HEPES-KOH, pH 7.4, 200 mM NaCl, 5 mM tris(2-carboxyethyl)phosphine). The protein was quantified by SDS-PAGE and stained with Coomassie Blue; concentrations were determined using a BSA standard curve. Lipids stored in chloroform were mixed (45% palmitoyloleoyl-phosphatidylcholine, 25% POPE, 25% dioleoylphosphatidylserine, 5% POPE-maleimide) and dried under a stream of nitrogen and lyophilized as mentioned above. The lipid film was hydrated to 3 mM (total lipid) with cross-linking buffer and formed into vesicles by extrusion through a 100-nm filter (Avanti Polar Lipids). Extruded vesicles (500 μl) were incubated with 50 μM protein overnight at 4 °C; the reaction was quenched by addition of 5 mM DTT. Proteoliposomes were floated through an Accudenz (Accurate Chemical) density gradient to remove unincorporated proteins as described previously (23, 24). POPE-maleimide was purchased from NOF America Corp. All other lipids were purchased from Avanti Polar Lipids.

In Vitro Fusion Assay—Lipid mixing experiments were carried out as described previously (24). Briefly, 100-μl reactions containing 4.5 μl of t-SNARE liposomes, 0.5 μl of v-SNARE liposomes, 1 μM synaptotagmin 1 C2AB, and 0.2 mM EGTA were incubated at 37 °C for 20 min, followed by the addition of 1 mM Ca2+ (arrows in Fig. 5). Fluorescence signals were measured using a BioTek synergy plate reader equipped with 460/40 excitation and 530/25 emission filters. Addition of 25 μl of 2.5% n-dodecyl maltoside was used to obtain the maximal dequenching signal to normalize the data. ΔTMD-94 (5 μM) was included in a subset of reactions to inhibit SNARE-mediated fusion.

Results

We tested four lipid-anchored syb2 constructs, two cytosolic syb2 constructs, full-length syb2, and constructs with fluorescent protein tags at either the C or N terminus (Fig. 1: “Experimental Procedures”). Three lipid-anchored constructs were...
truncated at the last residue of the linker before the TMD (residue 94) and had various cysteine-harboring sequences appended to the C terminus. ΔTMD-CIIL contains a geranylgeranylation site (9); ΔTMD-LC was used previously in tests of liposome fusion (10), and ΔTMD-CAC contains a well known prenylation motif. We also generated a CSPα construct designated ΔTMD-CSPα by subcloning the intact sequence of Syb2ΔATMR1 (used by Zhou et al. (13)) into our expression vector. The connection between syb2 and the C terminus of CSPα was not modified. In this construct the first 92 residues of syb2 (from the N terminus) were followed by 81 residues of the CSPα C terminus, which includes a palmitoylation sequence, CYYCCCLCCCFNCCCGKC. These proteins provide single (ΔTMD-CIIL and ΔTMD-LC), double (ΔTMD-CAC), and multiple (ΔTMD-CSPα) sites for post-translational acylation of long hydrocarbon chains.

We truncated syb2 at amino acids 84 and 94 to generate two cytosolic syb2 mutants (ΔTMD-84 and ΔTMD-94) that respectively omit or include the linker between the SNARE motif and TMD. We generated N-terminal EGFP-tagged constructs for all lipid-anchored syb2, cytosolic syb2 constructs, and WT to provide an expression marker. To control for the N-terminal EGFP tag, we subcloned WT syb2, ΔTMD-CSPα, and ΔTMD-CIIL without tags into a bicistronic vector from which EGFP was expressed from an IRES2. In addition to full-length syb2 with an N-terminal tag (EGFP-Syb2), we tested the control protein used by Zhou et al. (13), WT syb2 with a C-terminal mVenus tag (Syb2-RST-mVenus) in which the sequence RST served as the linker.

Endocrine Release—To assay syb2 constructs in endocrine release, we prepared cultured chromaffin cells from double knock-out (DKO) mice lacking the major dense-core vesicle SNAREs syb2 and cellubrevin. In contrast to chromaffin cells from WT mice, DKO chromaffin cells display very low secretion (20). We used amperometry recording to assay catecholamine release from individual chromaffin cells in response to depolarization by local application of a solution containing 105 mM KCl. We confirmed the poor secretion of DKO chromaffin cells (Fig. 2A) and found that DKO cells expressing WT syb2, with or without an N-terminal EGFP tag, secreted vigorously at levels comparable with littermate control cells; these cells all produced similar vesicle fusion frequencies of nearly 1 event/s (Fig. 2B) compared with ~0.1 events/s in DKO cells with no syb2. The residual release in this background likely reflects the action of a non-canonical v-SNARE, as has been found to be the case for residual spontaneous release at synapses (23). However, the molecules responsible for residual release in chromaffin cells lacking syb2 and cellubrevin have not been identified. These data thus indicate that DKO chromaffin cells provide a suitable background for testing syb2 mutants. A large number of syb2 mutants with TMD substitutions also rescue Ca2+−triggered exocytosis in this system (8). By contrast, the syb2 construct with a C-terminal mVenus tag, Syb2-RST-mVenus (13), only weakly rescued secretion, increasing the frequency of release events ~2-fold over background in DKO cells (Fig. 2B).

Expressing lipid-anchored syb2 constructs resulted in little or no rescue of secretion in DKO chromaffin cells (Fig. 2A). DKO cells expressing ΔTMD-CIIL, ΔTMD-84, or ΔTMD-94 secreted as poorly as untransfected DKO cells (Fig. 2B). ΔTMD-CSPα, ΔTMD-CAC, and ΔTMD-LC produced very small apparent increases in spike frequency over untransfected DKO cells, but these changes were not statistically significant. Untagged ΔTMD-CSPα and ΔTMD-CIIL produced results indistinguishable from the corresponding EGFP-tagged constructs (data not shown). The fact that three untagged constructs (including WT) produced results similar to the corresponding N-tagged constructs indicates that fusing EGFP to the N terminus does not influence function.

More detailed analysis of secretion kinetics confirmed the lack of support for dense-core vesicle exocytosis by lipid-anchored syb2. Cumulative plots of spike number versus time from DKO cells expressing WT syb2 (N-tagged or untagged) exhibited a brief delay after the start of depolarization and then rose nearly linearly before reaching a plateau (Fig. 2C, bar indicates depolarization). However, the C-tagged syb2-RST-mVenus construct produced a slower rising phase. The slopes of the linear regions (between 10 and 70% of the plateau), determined by linear regression, were much lower for ΔTMD-CSPα, ΔTMD-CAC, ΔTMD-LC, and C-tagged syb2-RST-mVenus than for WT and N-tagged syb2, and only ~2-fold greater than that for untransfected DKO cells (Fig. 2D). Slopes from DKO cells expressing ΔTMD-CIIL, ΔTMD-84, and ΔTMD-94 were indistinguishable from slopes from untransfected DKO cells; slopes from DKO cells expressing untagged ΔTMD-CSPα and ΔTMD-CIIL were indistinguishable from slopes obtained with the corresponding N-tagged proteins (data not shown). Only ΔTMD-CSPα produced a significant increase in slope over background. This slope was comparable with that with C-tagged syb2-RST-mVenus, and both were about one-third that seen in cells expressing untagged or N-tagged WT syb2.

We logged the number of cells responding with one or more spikes during the first two depolarizations (Table 1). Spikes were observed in 83 and 89% of DKO chromaffin cells expressing untagged and N-tagged WT syb2, respectively, versus 64% of untransfected DKO cells, and 68–72% for Syb2-RST-mVenus and lipid-anchored syb2 constructs. As noted above, the residual secretion in the DKO background probably reflects the presence of a non-canonical v-SNARE, which can produce a low level of secretion in over half the DKO cells. Expression of ΔTMD-84 and ΔTMD-94 reduced the number to 52 and 46%, respectively, suggesting that these constructs can interfere with the function of the putative non-canonical v-SNAREs.

It is possible that the lack of activity observed with lipid-anchored constructs resulted from incorrect localization. To address this possibility, the expression of each construct was evaluated by confocal fluorescence microscopy and analyzed for colocalization with the dense-core vesicle cargo, NPY-DsRed (Fig. 3). All of the lipid-anchored ΔTMD constructs targeted to dense-core vesicles, but ΔTMD-CAC and ΔTMD-LC constructs, also displayed some diffuse cytosolic expression (Fig. 3A). The ΔTMD-CSPα and ΔTMD-CIIL constructs targeted vesicles as effectively as WT syb2. Consistent with the lack of a membrane-anchoring motif, ΔTMD-94 was entirely diffuse with no puncta. Syb2 staining could not be seen in untransfected DKO cells. Expression patterns revealed by line
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scans showed peaks for WT syb2 and all lipid-anchored constructs, but the line scan for ΔTMD-94 was relatively smooth. Thus, the lipid-anchored constructs were efficiently targeted to vesicle membranes and did not accumulate on the plasma membrane.

The colocalization of syb2 variants with NPY-DsRed was quantified using the Pearson’s coefficient (Fig. 3B). WT and lipid-anchored syb2 constructs targeted dense-core vesicles significantly better than ΔTMD-94 (p < 0.0001 for WT and all lipid-anchored constructs). Only ΔTMD-CAC showed significantly lower colocalization than WT syb2, but the Pearson’s coefficient remained well above that for ΔTMD-94. Thus, even for the constructs that target vesicles slightly less efficiently than WT, targeting is still well above the negative control pro-

| Syb2-     | Percent responding | ΔTMD-CSPα | ΔTMD-CAC | ΔTMD-LC | ΔTMD-CIIL | ΔTMD-94 | ΔTMD-84 | None |
|-----------|--------------------|-----------|----------|---------|-----------|---------|---------|------|
| WT syb2   | 83% (44:53)        | 68% (26:38) | 67% (62:91) | 72% (52:72) | 67% (32:48) | 72% (34:47) | 46% (33:71) | 52% (46:88) | 64% (54:84) |
| EGFP-syb2 | 89% (49:55)        | 68% (26:38) | 67% (62:91) | 72% (52:72) | 67% (32:48) | 72% (34:47) | 46% (33:71) | 52% (46:88) | 64% (54:84) |
| Syb2-RST-mVenus | 68% (26:38) | 67% (62:91) | 72% (52:72) | 67% (32:48) | 72% (34:47) | 46% (33:71) | 52% (46:88) | 64% (54:84) |
| ΔTMD-CSPα | n.s.               | n.s.      | n.s.     | n.s.    | n.s.      | n.s.    | n.s.    | n.s. |
| ΔTMD-CAC  | ***                | ***       | ***      | ***     | ***       | ***     | ***     | *** |
| ΔTMD-LC   | n.s.               | n.s.      | n.s.     | n.s.    | n.s.      | n.s.    | n.s.    | n.s. |
| ΔTMD-CIIL | **                 | **        | **       | **      | **        | **      | **      | ** |

TABLE 1
The percent of secreting DKO chromaffin cells expressing the indicated construct, with the number of cells responding to depolarization over the number of cells tested in parentheses

FIGURE 2. Amperometry recording from chromaffin cells. A, amperometry traces from chromaffin cells: DKO cells, littermate controls, and DKO cells expressing untagged WT syb2, EGFP-Syb2, Syb2-RST-mVenus, lipid-anchored mutants, and ΔTMD-94. Untransfected DKO cells and DKO cells expressing ΔTMD-94 exhibited low secretion in response to depolarization by a solution containing 105 mM KCl (indicated by bars), but littermate and DKO cells expressing untagged WT syb2 or N-terminal tagged EGFP-Syb2 produced robust trains of spikes. Expression of C-terminal tagged Syb2-RST-mVenus construct in DKO cells produced much less support for exocytosis compared with N-tagged and untagged syb2. DKO cells expressing ΔTMD-CSPα, ΔTMD-CAC, ΔTMD-LC, and ΔTMD-CIIL all secreted poorly. B, spike frequencies of cells expressing the indicated protein during the first depolarization. Syb2-RST-mVenus, the lipid-anchored constructs, and ΔTMD constructs produced frequencies statistically indistinguishable from untransfected DKO cells. n = 26–62 cells from 2 to 7 DKO embryos. C, cumulative spike counts versus time in the first response to depolarization (indicated by the bar) from chromaffin cells expressing the indicated proteins. D, slopes of the rises in cumulative spike count plots from 10 to 70% of the plateau. The slopes of littermate control cells and DKO cells expressing N-tagged and untagged WT syb2 were similar and significantly higher than those from Syb2-RST-mVenus, lipid-anchored mutants, ΔTMD, and untransfected DKO cells. All lipid-anchored syb2 constructs were statistically indistinguishable from DKO except ΔTMD-CSPα. The comparison with EGFP-syb2 or DKO for one-way analysis of variance with post test is indicated by bars. **, p < 0.01; ***, p < 0.001; n.s., not significant. Paired comparisons of slopes with a t test were also performed in GraphPad Prism, and the slopes of Syb2-RST-mVenus and all mutants (except ΔTMD-94) were significantly higher than DKO.
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Expression and vesicle targeting in chromaffin cells. A, confocal images of untransfected DKO cells and DKO cells transfected with NPY-DsRed (red) and either WT syb2, or lipid-anchored syb2, or ΔTMD-94 (green). The inset and line scan panels correspond to the white squares and dashed lines in the merged panels. In line scan panels, black traces indicate syb2 variants, and red indicates NPY-DsRed. B, Pearson’s coefficient quantifies colocalization. Except for ΔTMD-CAC, lipid-anchored syb2 showed targeting comparable with WT syb2. WT and all lipid-anchored syb2 constructs targeted more efficiently than ΔTMD-94. p value for ΔTMD-LC is shown. The negative control construct ΔTMD-94 was well below all the other constructs (p value range 10^{-8}–10^{-4}). C, quantification of total fluorescence intensity indicates roughly similar expression levels for all constructs. n = 18–33 cells from three DKO embryos. **, p < 0.01; ***, p < 0.001. Scale bars, 3 μm.
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Individual events illustrate that regardless of the syb2 variant, spontaneous synaptic release in neurons. Expression of the C-tagged Syb2-RST-mVenus construct produced an ~2.5-fold increase over background that was similar to that seen with ΔTMD-CSPα and ΔTMD-CAC and was not statistically significant (Fig. 4, A–C).

To determine whether the poor rescue of synaptic release was due to mistargeting of the expressed proteins, we examined colocalization with synaptophysin, a synaptic vesicle marker. All syb2 constructs colocalized with synaptophysin, indicating that these molecules targeted synapses (Fig. 4D). However, the ΔTMD-CAC construct showed some diffuse expression in addition to the punctate, synaptic labeling. Expression of ΔTMD-94 was entirely diffuse with no puncta, and appeared uniform in cell bodies and processes. Quantification of colocalization between syb2 and syp indicated that the targeting of lipid-anchored syb2 was very similar to WT syb2, although the Pearson’s coefficient for ΔTMD-CAC was slightly lower than that for WT syb2 (Fig. 4E).

ΔTMD-CAC showed a similar targeting of dense-core vesicles.
that was slightly lower than WT but well above ΔTMD-94 (Fig. 3, A and B).

Liposome Fusion—To characterize the role of the syb2 TMD in membrane fusion in vitro, we assayed fusion of proteoliposomes harboring syntaxin-1A/SNAP-25B heterodimer with proteoliposomes harboring a WT or lipid-anchored syb2 construct (identical to those in cellular assays but without an EGFP tag). The lipid-anchored ΔTMD constructs were linked chemically by their cysteines to liposomes with 5% maleimide-POPE. Proteoliposome protein content was analyzed by SDS-PAGE with Coomassie Blue (Fig. 5A). Fusion of liposomes leads to mixing of the 1.5% NBD-PE and 1.5% rhodamine-PE in the v-SNARE vesicles with the nonfluorescent lipids in the t-SNARE vesicles. This dilutes the FRET pair and dequenches NBD fluorescence to give a signal that can be monitored over time (28). To enhance fusion rate and extent, the cytosolic domain of synaptotagmin 1 was included, and the reaction was triggered by addition of Ca^{2+} (arrows in Fig. 5B) (23). ΔTMD-94 was used as a control to block SNARE-mediated fusion. As observed previously with SNARE-mediated fusion (10), replacement of the syb2 TMD with one or two lipid anchors failed to support Ca^{2+}-synaptotagmin enhancement of SNARE-mediated fusion (Fig. 5B). Interestingly, ΔTMD-CSPα, which harbors several potential lipid anchors (Fig. 1), supported a small but significant amount of SNARE-mediated lipid mixing. Overall, these data are consistent with the amperometry data from chromaffin cells and the patch clamp data from neurons and suggest that the TMD of syb2 is required for efficient membrane fusion in vitro as well as in vivo.

Discussion

This study tested the role of the syb2 TMD in three different assays as follows: Ca^{2+}-triggered dense-core vesicle exocytosis from endocrine cells, spontaneous synaptic vesicle exocytosis from neurons, and liposome fusion. In each assay, four different lipid-anchored syb2 variants failed to function as effectively as the WT protein. Proteins that can accept one or two lipidic acyl moieties, ΔTMD-LC, ΔTMD-CIL, and ΔTMD-CAC, clearly failed to support liposome fusion. Some of these constructs supported low levels of exocytosis that were statistically indistinguishable from background and 4–6-fold lower than the levels of exocytosis achieved with WT syb2. This failure to support exocytosis cannot be attributed to mislocalization, as the proteins correctly targeted dense-core vesicles in chromaffin cells and synapses in neurons. In contrast to these three lipid-anchored syb2 constructs, ΔTMD-CSPα displayed limited fusion activity. This construct produced low but statistically significant enhancement of dense-core vesicle and proteoliposome fusion, but this enhancement was well below that observed with WT syb2. This unique property of ΔTMD-CSPα may reflect the 13 cysteine residues of its lipid-anchor motif (Fig. 1), which enable it to accept many lipidic acyl moieties. SNAREs harboring multiple lipidic acyl moieties have been shown to support lipid mixing previously (10). With the exception of ΔTMD-CSPα, the present results indicate that attachment of syb2 to the membrane by a lipid anchor does not support biological fusion. Fusion requires more than just membrane proximity provided by the formation of a SNARE complex with lipid-anchored syb2.

The TMD of syb2 performs some additional functions in addition to anchoring to the membrane. It is unlikely that this additional function includes protein folding because the SNARE motif of syb2 is a random coil when not incorporated into a SNARE complex (29). The TMD may contribute to the availability of the SNARE motif and the manner in which the SNARE motif is presented. This could happen if the TMD influenced the conformation of the rest of the protein or if the TMD controlled membrane insertion of the juxtamembrane region (30), which has been shown to influence exocytosis (31). Microlocalization and dimerization are additional possible functional roles for the TMD, although the functional relevance of dimerization has been difficult to establish (32).

The conclusion of an important role for the TMD is consistent with reports that substitutions within the TMDs of t-SNAREs (6, 7) and v-SNAREs (8) alter flux through fusion pores. The gap junction-like structure composed of these TMDs can form a proteinaceous channel without bending the
adjacent lipid bilayers. Only when the fusion pore expands does it incorporate lipid and bend lipid bilayers (33, 34). The important functional role for the syb2 TMD indicated by the present results is consistent with the virtually universal presence of a TMD in v-SNAREs (1–3).

Previous work has shown that lipid-anchored SNAREs can only drive lipid mixing in an in vitro fusion assay when the SNARES harbor either a lipid moiety long enough to span a lipid bilayer or multiple lipid moieties (35). Accessory proteins can also enable lipid-anchored SNAREs to drive fusion (11). A recent study of vacuole fusion indicated that yeast SNAREs differ in their requirement for a TMD for both lipid and content mixing (12). Another recent study reported that lipid-anchored fer in their requirement for a TMD for both lipid and content mixing (12). A previous study (36) showed that positive charge in the syb2 TMD impairs function to a degree similar to that found in this study with the syb2-RST-mVenus construct. The mEPSC frequencies we reported here in hippocampal neurons for the DKO background, ΔTMD-CSPα and their WT control, syb2-RST-mVenus. Here, we showed that the syb2-RST-mVenus construct functions poorly in the rescue of release, in contrast to untagged and N-tagged WT syb2 constructs, which rescue release to levels seen in littermate control cells. The mEPSC frequencies we reported here in hippocampal neurons for the DKO background, ΔTMD-CSPα and syb2-RST-mVenus (Fig. 4), were all similar to those reported previously in cortical neurons (13). Our replication of these results points to the poor function of the syb2-RST-mVenus construct as a WT control and thus indicates this as the basis for the discrepancy between our conclusion of poor function by lipid-anchored syb2, and the previous conclusion that they are fully functional (13). The poor function of the syb2-RST-mVenus construct probably results from the presence of arginine in the linker, as a previous study (36) showed that positive charge in precisely the same location immediately after the C terminus of the syb2 TMD impairs function to a degree similar to that found in this study with the syb2-RST-mVenus construct.

This study clarifies the role of the syb2 TMD in synaptic release and presents the first test of lipid-anchored SNAREs in Ca^{2+}-triggered dense-core vesicle exocytosis, as well as Ca^{2+}-synaptotagmin enhancement of SNARE-mediated liposome fusion. TMDs are essential elements in models that posit the initiation of SNARE-mediated membrane fusion through formation of a proteinaceous fusion pore (14, 15, 37). This study indicates that lipid-anchored syb2 variants lacking a TMD either fail to support fusion or support fusion very poorly. This was observed not only in two different forms of biological fusion but also in proteoliposome fusion in vitro. Because one or two lipidic linkages failed to provide significant support for fusion in any assay, this would suggest that proximity is not sufficient and that the TMD of syb2 plays an important role. This role may be formation of a proteinaceous fusion pore (14, 15, 37) or transmission of force to the lipid bilayer (16–19). Lipid anchors do not support these functions or support them poorly. Whether some other forms of attachment of proteins to a membrane can support such functions to achieve the levels of fusion seen with WT syb2 remains to be tested.

The levels of fusion seen with ΔTMD-CSPα in dense-core vesicle release and liposome fusion support the idea that anchoring a SNARE protein to a lipid bilayer through a large number of lipid moieties provides limited support for fusion (10). Multiple lipidic linkages may enable SNAREs to create a defect and deform lipid bilayers sufficiently to drive fusion at a slow rate. Contacts between the syntaxin and syb2 TMDs (38) contribute to the stability of the SNARE complex (39), and removal of its TMD would prevent a SNARE from harnessing these contacts as a driving force for fusion. SNARE-mediated fusion can occur through hemifusion intermediates (40), presumably bypassing proteinaceous fusion pores. Further studies will be required to address the interesting question of whether the limited fusion seen with ΔTMD-CSPα reflects a different mechanism involving hemifusion.

The number of SNARE complexes needed to trigger fusion may be only one for liposomes (41, 42), three for dense-core vesicles (43), and two for synaptic vesicles (44). These numbers are well below the estimates of 5–10 based on pore conductance (7, 45), and it is difficult to envision how fewer SNARE complexes could form a purely proteinaceous fusion pore. Other proteins can act as accessories to enable fusion driven by lipid-anchored SNAREs (11), and it is possible that another integral vesicle membrane protein such as synaptophysin can donate TMDs to a fusion pore. The speed of SNARE-mediated membrane fusion differs dramatically between different cellular membrane compartments, and it is possible that SNAREs can fuse membranes by more than one mechanism. The speed with which SNAREs catalyze fusion may depend on whether the nascent fusion pore contains protein or lipid or both (46, 47). Thus, proteinaceous fusion pores may serve as an adaptation to provide speed and temporal control for synaptic transmission.

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