ABSTRACT: Synaptotagmin I (syt), an integral protein of the synaptic vesicle membrane, is believed to act as a Ca\(^{2+}\) sensor for neuronal exocytosis. Syt’s cytoplasmic domain consists largely of two C2 domains, C2A and C2B. In response to Ca\(^{2+}\) binding, the C2 domains interact with membranes, becoming partially embedded in the lipid bilayer. We have imaged syt C2AB in association with lipid bilayers under fluid, using AFM. As expected, binding of C2AB to bilayers required both an anionic phospholipid [phosphatidylserine (PS)] and Ca\(^{2+}\). C2AB associated with bilayers in the form of aggregates of varying stoichiometries, and aggregate size increased with an increase in PS content. Repeated scanning of bilayers revealed that as C2AB dissociated it left behind residual indentations in the bilayer. The mean depth of these indentations was 1.81 nm, indicating that they did not span the bilayer. Individual C2 domains (C2A and C2B) also formed aggregates and produced bilayer indentations. Binding of C2AB to bilayers and the formation of indentations were significantly compromised by mutations that interfere with binding of Ca\(^{2+}\) to syt or reduce the positive charge on the surface of C2B. We propose that bilayer perturbation by syt might be significant with respect to its ability to promote membrane fusion.
forms aggregates but does not seem to adopt a single stoichiometry. Intriguingly, the binding of C2AB induces the formation of stable indentations in the bilayer, suggesting that it is able to significantly perturb the bilayer structure. The possible significance of this perturbation with respect to Ca\(^{2+}\)-triggered membrane fusion is discussed.

**EXPERIMENTAL PROCEDURES**

**DNA Constructs.** cDNAs encoding the syt cytoplasmic domain (C2AB, residues 96–421), the C2A domain (residues 96–265), and the C2B domain (residues 248–421) were expressed as GST-tagged proteins and cleaved with thrombin, as described previously (32). A C2AB mutant in which aspartate residues crucial to binding of Ca\(^{2+}\) to both C2A and C2B (D230, -232, -363, and -365) had been neutralized by conversion to asparagines, designated C2AM–C2BM (11, 13, 22, 23, 32), and C2AB harboring a mutation in C2B in which two lysine residues in a normally positively charged region (K326-K327) had been replaced with alanines, designated C2AB(KK) (11, 13, 22, 32), were also used. For all experiments, bacterial contaminants were removed from recombinant proteins by treatment with 1 M NaCl and RNase/DNase, as described previously (32).

**Formation of Supported Lipid Bilayers.** 1,2-Dioleoyl-sn-glycero-3-phosphatidylcholine (DOPC) and brain L-
all experiments, bacterial contaminants were removed from recombinant proteins by treatment with 1 M NaCl and RNase/DNase, as described previously (32).

**AFM Imaging.** AFM imaging was carried out at room temperature (20 °C) using a Digital Instruments Multimode atomic force microscope equipped with a J-scanner and a Nanoscope IIIa controller with an in-line electronics extender module (Veeco/Digital Instruments, Santa Barbara, CA). All images were collected using tapping mode in fluid and using oxide-sharpened silicon nitride probes (DNP-S; Veeco/Digital Instruments). The V-shaped cantilevers (typically exhibiting a spring constant of 0.58 N/m and an estimated curvature radius of 10 nm) were tuned to 10–20% below the peak of the resonance frequency, generally found between 7.5 and 9 kHz. The drive amplitude was set to generate a root-mean-square amplitude of 0.3–0.6 V. The microscope was engaged with a 0-nm scan area to allow for tuning and a setpoint adjustment of 100–200 nm above the surface before scanning of the sample was performed. The setpoint was adjusted to the highest setting that allowed imaging with little noise, to minimize the force applied to the sample. Images were captured at a scan rate of 1 Hz (unless otherwise noted), and with 512 scan lines per area. Data analysis was performed using commercially available software (NanoScope III software, Digital Instruments; Scanning Probe Image Processor, Image Metrology, Lyngby, Denmark).

**Determination of Molecular Volumes.** The molecular volumes of the protein particles were determined from particle dimensions based on AFM images. The particles tend to adopt the shape of a spherical cap. The heights and half-height radii were measured from multiple cross sections of the same particle, and the molecular volume was calculated using the following equation

\[
V_m = (\pi h/6)(3r^2 + h^2)
\]

where \(h\) is the particle height and \(r\) is the radius.

The molecular volume based on molecular mass was calculated using the equation

\[
V_c = (M_0/N_0)(V_1 + dV_2)
\]

where \(M_0\) is the molecular mass, \(N_0\) is Avogadro’s number, \(V_1\) (0.74 cm\(^3\)/g) and \(V_2\) (1 cm\(^3\)/g) are the partial specific volumes of the particle and water, respectively, and \(d\) (0.4 g of water/g of protein) is the extent of protein hydration (33).

**Stopped-Flow Rapid Mixing Experiments.** DOPC, synthetic 1,2-dioleoyl-sn-glycero-3-PS, and 1,2-dioleoyl-sn-glycero-3-phosphethanolamine-N-[(5-dimethylamino-1-naphthalenesulfonyl; dansyl-PE) were obtained from Avanti Polar Lipids. A phospholipid mixture, composed of 5% dansyl-PE, 25% PS, and 70% DOPC was dried and resuspended in HBS. Large (~100 nm) unilamellar liposomes were prepared by extrusion, as described previously (18).

Rapid mixing experiments were carried out using an Applied Photophysics (Leatherhead, U.K.) SX.18MV stop-flow spectrometer. Ca\(^{2+}\)-dependent interactions between wild-type [C2AB(WT)] or mutant C2AB [C2AB(KK)] and PS-containing liposomes were monitored by fluorescence resonance energy transfer. Native tryptophan and tyrosine residues in C2AB (donors) were excited at 285 nm, and the fluorescence intensity of the dansyl-PE (acceptors) in liposome membranes was collected using a 470 nm band-pass filter. The dead time of the instrument was ~1.2 ms.

For the assembly experiments, increasing concentrations of liposomes were premixed with 0.2 mM Ca\(^{2+}\) and then rapidly mixed (1:1) with 4 μM C2AB. Single-exponential functions were used to determine the observed rate constants (\(k_{\text{on}}\)) of Ca\(^{2+}\)-dependent C2AB–liposome interactions. The on- and off-rate constants (\(k_{\text{on}}\) and \(k_{\text{off}}\), respectively) for the interaction of Ca\(^{2+}\)-C2AB with liposomes were calculated, assuming pseudo-first-order kinetics (18, 34): \(k_{\text{obs}} = [\text{liposome}]k_{\text{on}} + k_{\text{off}}\). The dissociation constants (\(K_d\)) for C2AB–liposome interactions in the presence of Ca\(^{2+}\) were calculated as \(k_{\text{off}}/k_{\text{on}}\). All experiments were carried out at 25.3 °C.

For the disassembly experiments, 44 nM liposomes (5% dansyl-PE/25% PS/70% PC) were premixed with 4 μM C2AB in the presence of 0.2 mM Ca\(^{2+}\) and then rapidly mixed with an equal volume of HBS buffer containing excess EGTA (2 mM). The chelation of Ca\(^{2+}\) by EGTA resulted in a rapid loss of the fluorescence signal. The rate constants
for the apparent change in molecular volume. We also tested between groups was calculated using a Student’s t-test. The statistical significance of differences in unpaired data.

Results
We first imaged C2AB(WT) (2.5 nM) in association with mica, which presents a negatively charged surface at pH 7.6. As shown in Figure 1A, in the absence of Ca\(^{2+}\) (i.e., with 2 mM EGTA), C2AB appeared as a fairly homogeneous population of particles. The molecular volumes of a representative sample of particles were determined from the particle dimensions, according to eq 1. Frequency distributions of the molecular volumes are shown in panels B and D. The peaks of the two distributions are indicated.

Statistical Analysis. All errors are reported as standard deviations (SD). The statistical significance of differences between groups was calculated using a Student’s t-test for unpaired data.

For disassembly (k_{dis}) were estimated by fitting the kinetic traces using a single-exponential function. All experiments were carried out at 14.5 °C. (The disassembly of C2AB liposome complexes was too fast to monitor when the temperature was ~25 °C.)

Figure 2: Appearance of protein-free supported lipid bilayers. Liposomes composed of either DOPC only (A and B) or DOPC/PS (3:1; C and D) were deposited on mica in the presence of Ca\(^{2+}\) (500 µM), and the resulting supported lipid bilayers were imaged under fluid. Vertical cross sections (white lines) at the edge of the bilayers allow the heights of the bilayer to be determined: 5.8 nm in panel B and 4.8 nm in panel D. The shade-height scale refers to both panels A and C.

Clearly, the interaction of C2AB with mica has minimal physiological relevance. We therefore turned our attention to the interaction of C2AB with supported lipid bilayers under fluid, conditions that closely mimic those prevailing in vivo. Liposomes were produced and deposited onto mica in the presence of Ca\(^{2+}\) (500 µM). Under these conditions, the liposomes collapse and merge to form a single supported bilayer. When the liposomes were prepared from dioleoylphosphatidylcholine (DOPC) only, the resultant bilayer was smooth and featureless (Figure 2A). There were occasional gaps in the bilayer (e.g., dark area at the bottom right of Figure 2A), and a section through the bilayer revealed that the step in height down to the mica support was 5.8 nm (Figure 2B). The expected thickness of a lipid bilayer composed of DOPC is 3.5 nm (36). The additional thickness is likely due to the presence of a hydration layer between the bilayer and the mica; such a layer has been reported previously (37). Bilayers composed of DOPC and phosphatidylserine (PS) (3:1) were similarly featureless (Figure 2C) and now had a height step down to the mica support of 4.8 nm, likely indicating a closer interaction between the PS-containing bilayer and the mica (Figure 2D).

To study the interaction of C2AB with lipid bilayers, liposomes were preincubated with a low concentration of the protein (2.5 nM), and then supported bilayers were produced. Free C2AB was washed away before AFM imaging. When liposomes composed of DOPC were incubated with C2AB in the presence of Ca\(^{2+}\) (500 µM), the
resulting bilayer was almost featureless (Figure 3A), which was seen with DOPC bilayers produced in the absence of protein (Figure 2A). Figure 3B shows an image of a bilayer composed of DOPC and PS (3:1) that had been preincubated with C2AB in the absence of Ca\(^{2+}\). Again, few features were seen on the bilayer, as in the absence of protein (Figure 2C). In contrast, when the same experiment was repeated in the presence of Ca\(^{2+}\) (500 \(\mu\)M), the bilayer was now decorated with many raised features of various sizes (Figure 3C). Note also the presence of a large number of small particles attached to the uncoated areas of mica. These are presumably C2AB molecules primarily in dimeric form, as seen in Figure 1B. The lipid-bound particles are clearly larger than those attached to the mica and must therefore represent oligomers.

Figure 4 illustrates the interaction of C2AB, at a high concentration (1 \(\mu\)M), with bilayers containing DOPC and PS at two molar ratios (3:1 and 1:1). When the DOPC:PS ratio was 3:1, there was a heterogeneous array of particles attached to the bilayer (Figure 4A). A molecular volume distribution of the bound particles is shown in Figure 4B. The peak of the distribution was at a molecular volume of 588 nm\(^3\). In a previous electron microscopic study of C2AB bound to lipid monolayers, structures were observed that appeared to be barrel-shaped heptamers of the C2AB molecule (32). Interestingly, the size of the molecular volume peak obtained in our study would be consistent with structures of this type (predicted size of a monomer 80 nm\(^3\); 7 \(\times\) 80 nm\(^3\) = 560 nm\(^3\)). However, our data indicate that the bound protein particles are rather variable in size. Figure 4A also shows the presence of very large structures (>50 nm wide), which are likely to be liposomes bound to the bilayer but not fused (which were seen only rarely on protein-free bilayers). In an attempt to distinguish between these liposomes and bound C2AB particles, we routinely applied an arbitrary cutoff at 3000 nm\(^3\) to all of our molecular volume analyses. When the DOPC:PS ratio was 1:1, the binding density of C2AB was increased (Figure 4C), and the peak particle size increased from 588 to 951 nm\(^3\). These results emphasize the importance of PS in mediating the binding of C2AB to the lipid bilayer and are also consistent with a previous study indicating that C2 domain-containing proteins affect the distribution of PS within the plane of the bilayer, recruiting the lipid and binding to it in the form of protein patches (38). This previous study would predict that the mean particle size should increase along with the proportion of PS in the bilayer, which is what we observed. Some of the larger structures associated with the bilayer appeared as large, flat sheets. At higher magnifications, these sheets appeared to consist of two-dimensional aggregates of
globular particles (data not shown). Note that our observation of aggregation of C2AB contrasts with the results of a previous study (26), which found no evidence of aggregation of C2AB on liposomes, using an assay based on fluorescence resonance energy transfer. One possible explanation of this discrepancy is that C2AB interacts differently with planar bilayers (as in our experiments) and liposomes.

Closer inspection of images obtained in the presence of C2AB, PS, and Ca\(^{2+}\) revealed another feature that was absent when no C2AB binding occurred: the presence of many indentations in the surface of the supported bilayer (e.g., compare Figure 4A with Figure 2C). Further information about the origin of these indentations was obtained from repeated scanning of bilayers decorated with C2AB. Panels A and B of Figure 5 are images of the same area of a bilayer taken in successive scans, approximately 7 min apart. The protein binding pattern was very similar between the two scans. However, a small number of the particles were lost between scans (indicated by arrows). In many cases, when a protein dissociated it left behind an indentation in the bilayer. This effect is shown at a higher magnification in panels C and D of Figure 5. A section was taken through a bound particle and an indentation was taken at the position of the white line. (F) The section indicates that the depth of the indentation is 1.4 nm.

FIGURE 5: Dissociating C2AB particles leave behind indentations in the bilayer. Supported lipid bilayers with bound C2AB (initial concentration of 1 \(\mu\)M) were produced from liposomes composed of DOPC and PS (3:1). (A and B) The same area of a bilayer was scanned twice, with a 7-min interval. The appearance of the bilayer in the two scans is very similar; however, several particles have dissociated from the bilayer, leaving behind indentations (arrows). (C and D) At a higher magnification, the result of the dissociation of one particle is shown (arrow). (E) A vertical cross section through a bound particle and an indentation was taken at the position of the major indentation seen in the two successive scans. The depth of the indentation is 1.4 nm. Note that the indentation initially has a rather flat bottom and that its profile changes between scans. The depth of the indentation is \(\sim 3\) nm.

Our ability to identify indentations in the bilayer in the positions previously occupied by bound C2AB particles indicates that the mobility of the particles (and the indentations) in the plane of the bilayer is restricted. We have seen this restricted mobility previously when we imaged sphingomyelin/cholesterol-enriched domains in a DOPC bilayer (39). The shapes of the domains were largely preserved in successive scans but changed dramatically, even during the course of a single scan, in response to manipulation of the cholesterol content of the bilayer. Other workers have shown that influenza M2, a small tetrameric protein with a molecular mass of 50 kDa, diffuses freely within the plane of a supported lipid bilayer (40). The C2AB particles imaged here are larger and are therefore likely to move more slowly on the bilayer. In addition, the effect of Ca\(^{2+}\) on the PS-containing bilayer is likely to be significant. For instance, it has been shown by others that Ca\(^{2+}\) triggers a strong interaction between bilayer PS and the mica support (41) and can even cause the flipping of PS into the leaflet of the bilayer that is in contact with the mica (42). It is likely that these effects would considerably reduce the mobility of features in the bilayer, such as the indentations. We should emphasize that we cannot exclude the possibility that there is a population of monomeric C2AB that is moving so rapidly that it escapes detection by the scanning tip. We were interested in examining the stability of indentations in the bilayer, so we took successive scans of a suitable area. As shown in Figure 6, the same indentation was visible in successive scans taken 7 min apart (Figure 6A,B). However,
The scale bars refer to all images on either the left or the right. The shade bars refer to all images on either the left or the right. The shade-height scale refers to panels E and F.

We next asked whether the individual C2 domains, C2A and C2B, were able to form aggregates on PS-containing bilayers and whether they too were able to induce the formation of indentations. As shown in panels A and B of Figure 7, C2A bound efficiently to bilayers in the presence of Ca\(^{2+}\) and formed both multimeric particles and large sheetlike structures with a typical thickness of 2 nm. In addition, many indentations were seen, often in close association with attached particles, which seemed to line the edge of the indentations. C2B also formed large aggregates on the surface of the bilayers (Figure 7C,D). These were often elongated structures rather than sheets. C2B, too, caused the formation of indentations. Depths of the indentations seen with the individual C2 domains were 2.33 ± 0.60 nm (SD; \(n = 55\)) for C2A and 2.14 ± 0.66 nm (\(n = 61\)) for C2B. Both of these values are significantly larger than the depth of the indentations seen with C2AB (\(P < 0.001\) for C2A; \(P < 0.01\) for C2B). How does the presence of bound C2AB lead to the formation of the indentations? One possibility is that C2AB could bind tightly to the bilayer, and the scanning tip could then exert sufficient lateral force to dislodge both the protein particle and some attached lipid. This process could leave behind an area where one leaflet of the bilayer had been removed. However, the generation of such areas of a relatively stable lipid monolayer is thermodynamically unlikely. Alternatively, the protein could locally disrupt the structure of the bilayer, to produce a patch that is thin and therefore detectable by the tip as a depression. Such a patch might be sufficiently stable to persist for some time after the protein itself has dissociated. There are at least two reasons for favoring the latter explanation. First, the scanning was carried out in tapping mode, which is specifically designed to exert minimal lateral force on the scanned objects. It is therefore unlikely that the tip is able to push the protein particles off the bilayer. Second, areas of the bilayer are seen in initial scans that have little bound protein but many indentations. An extreme example is shown in panels E and F of Figure 7. This structure was observed in the first scan of this area of bilayer and therefore could not have resulted from the removal of bound particles by the scanning tip. We suggest that the incubation of C2AB with the liposomes might have brought about the multiple indentation of the bilayer. When the free protein was subsequently washed away, the resulting bilayer might have had a relatively low level of protein binding but might contain many residual indentations.

Figure 8 shows the effect of two types of mutation on the interaction of C2AB with the supported lipid bilayer. Panels A and B of Figure 8 show the binding of C2AB harboring mutations in the Ca\(^{2+}\) binding regions of both C2A and C2B (C2AM\(^{-}\)-C2BM; see above). It is clear that the binding density was low, compared with that of wild-type C2AB (Figure 3C), and that the typical particle size was smaller. The frequency distribution of molecular volumes (Figure 8C) had two peaks, a sharp one at 120 nm\(^3\) and a broader one at 240 nm\(^3\), indicating the presence of mainly monomers and dimers. Note also that the bilayer was almost free of indentations. Panels D and E of Figure 8 show the binding of C2AB harboring a mutation in C2B in which two lysine residues (K326 and K327) in a normally positively charged region have been replaced with alanines [C2AB(KK); 11, 13, 22, 32]. As for the C2AM\(^{-}\)-C2BM mutant, the binding density of C2AB(KK) was low, and the typical particle size was small; the peak of the frequency distribution shown in Figure 8F is 219 nm\(^3\), consistent with a predominantly dimeric structure. Again, there are almost no indentations in the bilayer. The effect of mutations in the Ca\(^{2+}\) binding regions of the C2 domains on the interaction of C2AB with the supported lipid bilayer is unsurprising, since these regions are known to become embedded in the bilayer (9–11, 13, 18, 27–30), and the effects of these mutations on C2AB binding to liposomes are well-documented (e.g., ref 11). The effect of the KK mutation is more difficult to explain, especially as the polybasic region of C2B where the two lysines reside is quite distant from the Ca\(^{2+}\)-binding loops and has been reported not to make intimate contact with the surface of the membrane (30).
of C2AB(WT) and C2AB(KK) are not significantly different (P > 0.05), in contrast to the large difference in their behavior in our AFM imaging experiments. Similar discrepancies have arisen previously in experiments designed to investigate the association of isolated C2B with PS-containing liposomes, and it has become clear that the reported behavior of C2B depends on the assay system used. For instance, it was found that C2B, immobilized as a GST fusion protein, binds radiolabeled liposomes only weakly in pull-down assays (5, 32), whereas C2B efficiently cosediments with liposomes (32). The effect of the KK mutation is also assay-dependent. For example, it had no effect on the interactions of C2B with PS-containing membranes in the presence of Ca²⁺. Error bars represent SDs of three independent experiments. Values of kₐ and kₐ₋d for C2AB(WT) and C2AB(KK) are not significantly different (P > 0.05). (D) Summary of the values of kₐ, kₐ₋d, and the dissociation constant (K_d). (E) Representative trace of the disassembly kinetics of the C2AB(WT)—liposome complex. The rate constant for the fluorescence change, kₐ₋d, was calculated by fitting the kinetic traces with a single-exponential function. (F) Representative trace of the disassembly kinetics of the C2AB(KK)—liposome complex. Values of kₐ₋d for C2AB(WT) and C2AB(KK) are not significantly different.
the polybasic region might be involved in the anchoring of the synaptic vesicle membrane to the plasma membrane during exocytosis (26). To account for the discrepancies between the results produced using different assays, it has been suggested that cosedimentation of syt with liposomes requires an additional property (possibly the ability to aggregate) that depends on the polybasic region (32). In this context, it should be noted that, as in our experiments, the KK mutation abolished the oligomerization of C2AB on lipid monolayers, as visualized by electron microscopy (32). It should also be borne in mind that in the kinetic experiments, where the lipid:protein ratio is relatively high, most C2AB molecules will be bound to the membrane. Under these conditions, the small difference in the affinity of binding between C2AB(WT) and C2AB(KK) might not be detected. In contrast, in most of the AFM experiments, the lipid:protein ratio will be lower so that only a fraction of the C2AB molecules will be bound to the bilayer. Under these conditions, differences in behavior between C2AB(WT) and C2AB(KK) might be more obvious.

DISCUSSION

We have shown that syt associates with PS-containing lipid bilayers in the form of aggregates of varying stoichiometry. These aggregates are seen at both low (2.5 nM) and high (1 \(\mu\)M) concentrations of syt. Hence, aggregation appears to be independent of the extent of saturation of the bilayer by syt. In addition, the binding of syt to the bilayers caused the perturbation of the bilayer structure, producing relatively stable indentations. It has been shown previously, using changes in the emission characteristics of strategically placed fluorescent probes, that both C2A and C2B are able to penetrate the bilayer in response to Ca\(^{2+}\) binding (9–11, 13, 18, 27). Specifically, loops 1 and 3 of both C2 domains dip into the bilayer in response to Ca\(^{2+}\) binding. Fluorescence quenching experiments revealed that the tip of C2A can penetrate to a depth of \(\sim 1.5\) nm (9, 27), similar to the depth of the indentations seen with C2AB in our experiments. In addition, others have concluded, on the basis of studies using site-directed spin labeling of C2AB, that both C2 domains penetrate to a depth of \(\sim 1\) nm (30).

Since it is known that perturbation of the bilayer structure can under some circumstances trigger the fusion of lipid bilayers (e.g., ref 43), the bilayer perturbation caused by C2AB might be significant with respect to the role of syt as a Ca\(^{2+}\) sensor for neurotransmitter release. It should be emphasized that there is a large difference between the conditions under which our experiments were conducted and those prevailing in vivo; for instance, in the nerve terminal, syt is anchored to the synaptic vesicle membrane through its transmembrane domain, rather than being free in solution, as in our studies. It is possible that during neurotransmitter release syt is targeted to the presynaptic plasma membrane through its interaction with the t-SNAREs and phosphatidylinositol 4,5-bisphosphate (13, 14, 22), and that in response to Ca\(^{2+}\) a cluster of syt molecules causes membrane destabilization at the site of vesicle docking, which then leads to membrane fusion.

Most other proteins studied by AFM in association with supported lipid bilayers by us (37, 44–46) and by others (e.g., refs 40 and 47) have not been reported to cause the indentations seen here with C2AB. Interestingly, however, the 50-kDa C-terminal region of the tetanus toxin heavy chain, which mediates the binding of the toxin to the ganglioside GT1b, has recently been shown by AFM analysis to cause the formation of 40–80 nm diameter and 1.5 nm deep indentations in supported lipid bilayers (48). These indentations developed slowly (i.e., over \(\sim 12\) h) and were very stable. Total internal reflection fluorescence microscopy conducted in parallel with the AFM analysis indicated that the toxin fragment was present within the indentations. The ganglioside binding region of this tetanus toxin fragment has a predominantly positively charged surface. Significantly, a number of previous studies have found that other polybasic molecules cause the production of indentations in lipid bilayers similar to those seen in our experiments. In some cases, for example with poly-L-lysine and poly(ethylenimine), these features are actually transbilayer holes that have been shown to be associated with the permeabilization of the bilayers (49). In other cases, for example with diethylaminomethylxextran, the indentations are too shallow to completely penetrate the bilayer, as seen in our experiments (49). These polybasic molecules [e.g., poly(amidoamine)] have also been found to line the edges of the indentations (50), as we have found, particularly with the isolated C2 domains. Furthermore, the binding of the polybasic antimicrobial peptide MSI-78 to dimyristoyl-sn-glycero-3-phosphocholine bilayers has been shown by AFM to cause thinning of the bilayer by just more than 1 nm (51). Of course, both C2 domains of syt have predominantly positively charged surfaces in their Ca\(^{2+}\)-bound states (4, 5).

In addition, the C2B domain contains a prominent polybasic region (2, 3). Reduction of the positive charge on this region by the introduction of neutralizing mutations is known to interfere with syt’s ability to support neurotransmitter release in vivo (14, 52, 53). In our experiments, these mutations significantly reduce both the level of association of C2AB with the supported lipid bilayers and the extent of generation of bilayer indentations. The importance of the positively charged surfaces of Ca\(^{2+}\)-liganded syt with respect to its ability to interact with bilayers and to function in vivo supports the suggestion that membrane perturbation by syt’s C2 domains might be key to its physiological function.

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