Structural basis for the clamping and Ca\textsuperscript{2+} activation of SNARE-mediated fusion by synaptotagmin

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Synaptotagmin-1 (Syt1) interacts with both SNARE proteins and lipid membranes to synchronize neurotransmitter release to calcium (Ca\textsuperscript{2+}) influx. Here we report the cryo-electron microscopy structure of the Syt1-SNARE complex on anionic-lipid containing membranes. Under resting conditions, the Syt1 C2 domains bind the membrane with a magnesium (Mg\textsuperscript{2+})-mediated partial insertion of the aliphatic loops, alongside weak interactions with the anionic lipid headgroups. The C2B domain concurrently interacts the SNARE bundle via the 'primary' interface and is positioned between the SNAREpins and the membrane. In this configuration, Syt1 is projected to sterically delay the complete assembly of the associated SNAREpins and thus, contribute to clamping fusion. This Syt1-SNARE organization is disrupted upon Ca\textsuperscript{2+}-influx as Syt1 reorients into the membrane, likely displacing the attached SNAREpins and reversing the fusion clamp. We thus conclude that the cation (Mg\textsuperscript{2+}/Ca\textsuperscript{2+}) dependent membrane interaction is a key determinant of the dual clamp/activator function of Synaptotagmin-1.
The rapid Ca$^{2+}$-triggered release of neurotransmitters at the synapse is a highly orchestrated process\textsuperscript{1-3}. The proteins involved are known and well-characterized\textsuperscript{4-5}. This includes the SNARE (Soluble N-ethylmaleimide-sensitive factor attachment protein receptor) proteins (VAMP2, Syntaxin, and SNAP25) that catalyze synaptic vesicle (SV) fusion, as well as chaperones Munc13, Munc18, and regulators Complexin and Synaptotagmin\textsuperscript{4-5}. Synaptotagmin-1 (Syt1) is a key component involved in all stages of the process, including SV docking and priming\textsuperscript{6-9}, preventing un-initiated SV fusion\textsuperscript{10-11} and triggering fusion upon Ca$^{2+}$ influx\textsuperscript{12-13}. Remarkably, such broad specialization has not resulted in structural complexity of the protein.

Genetic analysis shows that Syt1 also plays a critical role in ‘clamping’ spontaneous release and Ca$^{2+}$-evoked asynchronous release\textsuperscript{11,15,32}. Similarly, the deletion of Complexin (Cpx) shows an increase in spontaneous release frequency and de-synchronization of evoked responses\textsuperscript{33-35}. This suggests that Syt1 and Cpx act ‘synergistically’ to clamp un-initiated/delayed release components to ensure Ca$^{2+}$-dependent membrane interactions. Under in vitro conditions, Syt1 and Cpx have been shown to independently clamp SNARE-mediated vesicle fusion under physiologically relevant conditions\textsuperscript{13,21,24,25}. The overall protein appearance suggests that upon Ca$^{2+}$-binding, the Syt1 C2 domains reorient into the membrane, likely displacing the associated SNAREpins from the primary interface. Our data indicates that the Ca$^{2+}$ trigger of SV fusion possibly involves large conformational changes in the associated proteins.

**Results**

Syt1\textsuperscript{C2AB}–SNARE complex helically organizes on lipid tubules. Our attempts to co-crystallize Syt1 and SNARE on membranes using individual proteins were unsuccessful. So, we used a protein construct wherein the minimal C2B domains of Syt1 (Syt1\textsuperscript{C2AB}) are covalently linked to the SNARE complex, which was proven to be successful in 3D crystal formation\textsuperscript{41}. Negative stain EM analysis of Syt1\textsuperscript{C2AB}–SNARE protein complex incubated with liposomes containing negatively charged lipids (DOPC/DOPS/PIP2 60/34/6) revealed the formation of protein-coated lipid tubules with helical diffraction peaks (Supplementary Fig. 1). These protein-coated tubular projections, formed in the presence of 1 mM free Mg$^{2+}$, occurred in low frequency and were quite variable in size, with diameters ranging between ~40 nm and ~60 nm (Supplementary Fig. 1). Cryo-EM analysis further showed that the lipid tubules have inconsistent protein decoration, wherein some were partially decorated with protein, while other tubules were devoid of protein (Supplementary Fig. 1).

To improve the quantity and quality of the protein coated lipid tubes suitable for the helical 3D reconstruction process, we applied Syt1\textsuperscript{C2AB}–SNARE protein to pre-formed lipid nanotubes (LNT) (Supplementary Fig. 2). The LNTs of fairly uniform size (diameter ~35 nm) were assembled with the inclusion of glycolipid (20% galactosylceramide, GalCer 24:1) into the vesicle lipid mixture\textsuperscript{49}. Glycolipid derived nanotubes are routinely employed as substrate for preparing 2D-crystals, as they promote the formation of helical arrays of adsorbed proteins and macromolecular assemblies\textsuperscript{50}. For example, they were recently used to helically organize and reconstruct the membrane bound blood coagulation factor VIII containing C2 domains by Cryo-EM\textsuperscript{51}.
Correspondingly, negative stain EM analysis showed a large number of homogeneous protein coated LNTs, with a strong diffraction pattern corresponding to a helical organization of protein molecules on the lipid surface. These protein-coated LNTs, formed in presence of 1 mM free Mg$^{2+}$, were analogous to the tubules derived from vesicles both in their appearance and diffraction peaks (Supplementary Fig. 2). But frozen LNT samples analyzed by CryoEM revealed a majority of unorganized or poorly diffracted tubes (Supplementary Fig. 2). Since the partial protein decoration was never observed by negative stain EM for the vesicle-derived or pre-assembled tubules, we reasoned that the freezing likely weakens the Syt1-membrane interaction and the proteins were falling off the tube surface during the Cryo-EM grid preparation. So, to ensure tight binding of the proteins on the surface and to improve their packing, we increased the acidic lipid content and simplified the lipid composition of the LNTs. The optimum composition was found to be DOPE: GalCer at 80:20 ratio, which vastly improved both the overall frequency and quality of the helically organized Syt1C2AB-SNARE coated tubes (Fig. 1a). This enabled us to collect a homogenous dataset suitable for high-resolution reconstruction of the Syt1-SNARE complex bound to membranes under resting conditions (1 mM Mg$^{2+}$).

**CryoEM structure of Syt1C2AB-SNARE on lipid surface.** We evaluated the quality of protein coating by analyzing the power spectra of selected tubes and their suitability for helical indexing by the Fourier-Bessel method.$^{32}$ Protein decorated LNTs around ~41 nm (outer diameter) showed helical diffraction corresponding to homogenous helical crystal (Fig. 1b), so this subset of tubes was selected for further structural analysis. Helical indexing of the representative tube defined the Syt1C2AB-SNARE arrangement on the surface as 14,−4 start helix with a helical rise (Δz) of 7.35 Å and a helical twist (Δϕ) of 78.48° (Fig. 1b). These parameters were then used for the iterative helical real-space reconstruction (HRSR)$^{33}$. The resulting 3D reconstruction at 10.4 Å resolution showed protein decoration on the tube surface with prominent rod-like features (SNARE proteins) arranged almost parallel to the tube axis and with globular structures (C2 domains) attached to the membrane surface (Fig. 1c, d). SNARE complexes were well resolved with distinguishable densities corresponding to the α-helices of SNARE proteins, however the orientation of SNAREs and relative positioning of the Syt1 C2A and C2B domains was not clear. Noteworthy, the proteins are arranged parallel to the nanotube axis (Fig. 1c, d) and thus, locally experience relatively flat-bilayer surfaces.

We noticed a striking visual similarity of protein organization between the 2D-crystal on the membrane and the recently published 3D crystal of the same complex$^{41}$. So, we performed rigid body docking with separate C2A, C2B, and SNAREs derived from the Syt1-SNARE crystal structure (PDB code: 5CCI$^{41}$). The crystal structures were unambiguously docked into the EM map and showed SNAREpins arranged in an antiparallel fashion with membrane-distal C-terminal interaction. The Syt1 protein located between the SNARE complex and membrane surface (Fig. 2, Supplementary Fig. 3, Supplementary Movie 1), with the C2B domain interacting with the middle portion of SNARE complex. Relative position of C2B over the SNAREs was very similar to the recently described ‘primary’ binding site, involving the t-SNARE helices$^{31,42}$. The linker between the C2A and C2B domains was fully extended, with the C2A domain locating close to but not
interacting with the N-terminal end of the same SNAREpin (Fig. 2, Supplementary Fig. 3). In parallel, we also aligned the crystal structure of a single Syt1C2AB–SNARE complex (primary interface unit) from 5CCI X-ray structure onto our CryoEM-derived Syt1C2AB–SNARE reconstruction (Supplementary Fig. 4). It showed that in relation to the SNARE complex, the Syt1 C2B domain is identically positioned and nearly superimposable, but the C2A position deviates slightly compared to the crystal structure (Supplementary Fig. 4). This indicates that on the membrane surface, the primary interface between C2B and SNARE proteins is preserved, but the C2A domain is likely more flexible. Overall, the Syt1C2AB–SNARE complexes were more densely organized on the lipid surface, and by conforming to the curved surface of the LNTs their higher order organization deviated from the 3D crystal lane (Supplementary Fig. 4).

The Syt1C2AB–SNARE complex adopted a previously undescribed orientation on the membrane surface with the C-terminal end of the SNAREs pointing away from the membrane and the Syt1 C2A and C2B calcium loops oriented towards the membrane (Fig. 2). Fitting the C2A and C2B domain crystal structure showed that one of the Ca2+-coordinating aliphatic loops (loop 3) on both C2 domains is partially inserted into the phospholipid heads plane, while loop 1 (black asterisk) stays close to the lipid surface. The conserved polybasic patch (residue K326/327 in blue) in the C2B domain is also positioned close to the lipid head group, suggesting possible interaction.

**Fig. 2** Organization of Syt1C2AB–SNARE on a lipid membrane surface. The rigid body docking of single C2A, C2B, SNARE crystal structures from Syt1C2AB–SNARE 3D crystal (5CCI) onto the Cryo-EM map. SNAREpins (multi-color) form the top-most protein layer and are positioned in antiparallel fashion with their C-terminus pointing away from the membrane (yellow). The Syt1 C2B domains (gray) are positioned between SNAREpins and the outer membrane’s phospholipid heads plane. The linker between the C2A (cyan) and C2B domain is extended and clearly visible (red asterisk) allowing unambiguous identification of C2A and C2B domains within the same Syt1C2AB unit. Both C2 domains adopt a previously unseen orientation relative to the membrane surface. One of the Ca2+-coordinating aliphatic loops (loop 3) on both C2 domains is partially inserted into the phospholipid heads plane, while loop 1 (black asterisk) stays close to the lipid surface. The conserved polybasic patch (residue K326/327 in blue) in the C2B domain is also positioned close to the lipid head group, suggesting possible interaction.

**Mg2+ orchestrates membrane interaction of Syt1C2AB.** To gain further insight into the previously unseen membrane orientation...
of the Syt1C2AB–SNARE complex, we investigated the consequence of targeted mutations altering the membrane interaction of the Syt1 C2B domain. Specifically, we assessed its ability to form an organized structure on the LNT surface based on visual appearance and helical diffraction pattern (Fig. 3). Neutralizing the conserved polybasic motif (K326A/K327A) of the C2B domain did not significantly affect the helical organization of Syt1C2AB–SNARE on the LNTs, which were visually identical to the wild type (Fig. 3). This suggested that the interaction of the polybasic patch with acidic lipids does not solely determine the attachment of the C2B domain to the membrane surface. Consequently, we looked into the effect of altering the physiochemical properties of the Ca\textsuperscript{2+}-loops. Polar mutations of aliphatic residues in the Ca\textsuperscript{2+}-loop (V304N/Y364N/G368N; Syt1\textsuperscript{3N}), intended to decrease its hydrophobicity and hinder its membrane insertion, abrogated the helical organization even though the protein still bound to the LNTs (Fig. 3). Conversely, enhancing the hydrophobicity of the Ca\textsuperscript{2+}-loops, by neutralizing the Ca\textsuperscript{2+}-coordinating Asp residues (D309/D363/D365A; Syt1\textsuperscript{3A}), stabilized the organized structures even in the absence of Mg\textsuperscript{2+} (Fig. 3). This was striking as the wild type Syt1 protein strictly required Mg\textsuperscript{2+} to form organized structures on LNTs. In fact, Mg\textsuperscript{2+} was crucial for the wild type Syt1C2AB–SNARE complex to stay bound to the LNT surface during the freezing procedure. This indicated that the Mg\textsuperscript{2+} stabilizes the Syt1–membrane interaction under these conditions, likely by coordinating the conserved Asp residues with the acidic lipid head group. Altogether, these results show the importance of Mg\textsuperscript{2+}-dependent C2B domain orientation with partial membrane insertion of the aliphatic loops for the Syt1C2AB–SNARE complex helical arrangement.

To ascertain that the partially membrane inserted state with Mg\textsuperscript{2+} is not an artifice of crystallography, we examined if this can occur under dilute conditions using isolated proteins. We employed a well-established fluorescence assay with an environment-sensitive probe IAEDANS [5–9]([2-(iodoacetyl) amino)ethyl]amino)naphthalene-1-sulfonic acid introduced a tip of the Ca\textsuperscript{2+}-loops of Syt1 C2A and C2B domains to track their localization\textsuperscript{5,20}. In the presence of negatively charged liposomes (60% PC, 34% PS, 6% PIP2), the IAEDANS probe introduced at the loop 3 of both C2A (residue 235) and C2B (residue 367) exhibited a Mg\textsuperscript{2+}-dependent increase in fluorescence intensity (~40% compared to the EDTA condition) and blue shift (~15 nm) in the emission spectrum. This is consistent with IAEDANS probe locating within a hydrophobic (i.e., lipid membrane) environment (Fig. 4). In contrast, very little to no change in fluorescence signal was observed for IAEDANS probes introduced into loop 1 (residue 173 on C2A; or residue 304 on C2B) (Fig. 4, Supplementary Fig. 6). Noteworthy, the fluorescence analysis matched perfectly with the CryoEM structure showing a partial membrane insertion of C2 domain loop 3, but not loop 1 (Fig. 2).

The fluorescence profiles were markedly different in the presence of 1 mM Ca\textsuperscript{2+}. The IAEDANS probes on both loops 1 and 3 of either C2 domain exhibited a strong increase in fluorescence intensity (~95% compared to EDTA control) and a larger blue shift (~25 nm) in emission spectrum corresponding to stronger and deeper membrane association (Fig. 4, Supplementary Fig. 6). These results demonstrate that in the presence of 1 mM Mg\textsuperscript{2+}, loop 3 of the Syt1 C2A and C2B domains are submerged into the membrane even under non-crystallizing conditions but not as deep compared when Ca\textsuperscript{2+} is present. Importantly, it implies that the Mg\textsuperscript{2+}-induced partial-membrane insertion of Syt1 is an intrinsic feature, distinct from the Ca\textsuperscript{2+}-bound state, even though it involves the same structural elements. This is consistent with recent surface forces apparatus

| Protein construct | Intended effect                     | Outcome                           | Image | FFT |
|-------------------|------------------------------------|-----------------------------------|-------|-----|
| Wild type         | Control                            | Helical organization              | ![Image](https://example.com/image1.png) | ![FFT](https://example.com/FFT1.png) |
| K326/327A         | Reduced polybasic patch charge     | Helical organization              | ![Image](https://example.com/image2.png) | ![FFT](https://example.com/FFT2.png) |
| D309A/D363A/ D365A| Enhanced hydrophobicity of Ca\textsuperscript{2+} loops | Helical organization (even in absence of Mg\textsuperscript{2+}) | ![Image](https://example.com/image3.png) | ![FFT](https://example.com/FFT3.png) |
| V304N/Y364N/ I367N| Decreased hydrophobicity of Ca\textsuperscript{2+} loops | LNT binding, No organization     | ![Image](https://example.com/image4.png) | ![FFT](https://example.com/FFT4.png) |
| Syntaxin1 H239D/D250H| Disrupt SNARE c-terminal interaction | LNT binding, No organization     | ![Image](https://example.com/image5.png) | ![FFT](https://example.com/FFT5.png) |

**Fig. 3** Mutations in the Syt1C2AB and SNARE complex used to identify and validate critical crystallographic contacts. The mutations were assessed for their ability to form a helical crystal on the membrane surface by visual appearance and by helical diffraction pattern of corresponding Fourier transforms (shown as acquired at the vertical tubes orientation) of the decorated LNTs by Cryo-EM analysis. Disruption of the C2B polybasic patch (K326A/K327A) had no effect on helical organization, but alteration of the Ca\textsuperscript{2+}-loops showed a strong effect highlighting its critical role. Disrupting the C-terminal interaction of the SNAREpin also interfered with the helical organization indicating its role in stabilizing the helical formation.
measurements showing that Mg\textsuperscript{2+} increases the binding energy of Syt1 to anionic membranes as compared to EGTA, but is weaker than the Ca\textsuperscript{2+}-bound state (~6 kBT for EGTA, ~10 kBT with Mg\textsuperscript{2+}, ~18 kBT with Ca\textsuperscript{2+}).

So, it emerges that the Mg\textsuperscript{2+}-driven Syt1 orientation determines the relative positioning of the SNARE complex on the lipid membrane. As such, the SNARE complex bound to the Syt1 C2B domain via the primary interface is oriented with its C-terminus pointed away from the membrane surface (Fig. 2, Supplementary Fig. 3). If this Syt1–SNARE arrangement were to form under physiological conditions, it might sterically hinder the complete assembly of the membrane-anchored SNAREs and prevent fusion under resting (1 mM Mg\textsuperscript{2+}) conditions (see Discussion).

In our helical structure (similar to the 3D crystal), we also observed an interaction between neighboring SNAREpins mediated by charged residues in the Syntaxin1 and VAMP2 C-termini (Supplementary Fig. 3). Disrupting this interaction, using a mutation on Syntaxin (H239D/ D250H), resulted in complete loss of helical organization on LNTs (Fig. 3), highlighting its equal importance in the helical arrangement. IAEDANS-based fluorescence experiments, including analysis of Syt1 3A/3N proteins, shows that helically organized structures are stably formed only under conditions wherein Syt1 adopts a partially inserted state (Supplementary Fig. 7). Hence, we reason that the Syt1 C2B guides the SNAREpin orientation, and the C-terminal SNARE–SNARE interaction in turn stabilizes the membrane...

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**Fig. 4** Mg\textsuperscript{2+} induces the partial insertion of loop 3 on both C2 domains. a, b Fluorescence analysis using environmentally-sensitive probe, IAEDANS introduced at the tip of the aliphatic loops of the C2 domains shows the Mg\textsuperscript{2+} (red curve) induces an insertion of loop 3 (residue S235 of C2A & residue I367 of C2B) as evidenced by an increase in the fluorescence intensity and blue-shift in the emission maxima as compared to an EDTA control (black curve). This Mg\textsuperscript{2+}-induced membrane interaction is distinct, with shallower positioning of the aliphatic loops in the lipid membrane as compared to the Ca\textsuperscript{2+}-bound state (blue curve) (c, d) Normalized fluorescence signal at the emission maxima shows that Ca\textsuperscript{2+}-induces deep insertion of both loop 1 (residue M173 in C2A and V304 in C2B) and loop 3, but Mg\textsuperscript{2+} specifically triggers partial insertion of loop 3 alone in both C2 domains. Representative fluorescence curves are shown, along with averages and standard deviations from 4 to 5 independent experiments.
inserted state of the Syt1 C2 domains and the helical arrangement on the membrane surface. Correspondingly, we were unable to crystalize the Syt1–SNARE complex with truncated SNARE complexes.

**Ca\(^{2+}\) disrupts Syt1\(^{C2AB}\)–SNARE organization on the membrane surface.** The Syt1\(^{C2AB}\)–SNARE complex did not form any organized structures on LNTs in the presence of 1 mM free Ca\(^{2+}\) (Fig. 5). Instead, we observed repeating densities on the membrane surface (red arrows in Fig. 5) with unorganized auxiliary densities (blue arrows in Fig. 5) on the top. Given the dimensions of the observed densities and known physiology\(^5,19,20\), we infer that the dense packing on the membrane (with smeared diffraction peaks) correspond to the Syt1\(^{C2AB}\) protein, and the SNARE complex displaced from its primary binding site represents the surrounding density. This is in stark contrast to 3D X-ray crystallography results, which shows similar Syt1\(^{C2AB}\)–SNARE organization in the Ca\(^{2+}\)-bound state\(^41\). This leads us to conclude that the Ca\(^{2+}\)-induced reorientation of Syt1 into lipid membranes\(^5,19,20\) disrupts the Syt1–SNARE organization found under the resting conditions. Furthermore, it appears that the deep membrane insertion of the C2 domain aliphatic loops\(^19,20\) dislodges the associated SNAREpins from the primary interface. Consistent with this premise, the Syt1\(^{3A}\) protein that lacks the ability to bind Ca\(^{2+}\) and thus, insert deep into the membranes (Supplementary Fig. 7) helically organized on LNTs even in the presence of Ca\(^{2+}\) (Supplementary Fig. 8). Also, Ca\(^{2+}\) restored the ability of the Syt1\(^{3N}\) mutant to organize on LNTs (Supplementary Fig. 8). Noteworthy, IAEDANS fluorescence experiment indicates that Ca\(^{2+}\) induces partial insertion of the Syt1\(^{3N}\) aliphatic loops mimicking the Mg\(^{2+}\)-bound state of the wild type Syt1 (Supplementary Fig. 7). In sum, we find that Ca\(^{2+}\)-binding induces a large-scale conformational rearrangement of the Syt1–SNARE complex on the lipid membrane surface, disrupting the pre-fusion architecture.

**Discussion**

Data presented establish that under resting conditions (1 mM Mg\(^{2+}\)), Syt1\(^{C2AB}\) adopts a previously unknown, molecularly distinct orientation on the membrane surface with partial insertion of the aliphatic loops into the phospholipid membrane. The structure further reveals that the Syt1 C2B domain simultaneously binds the t-SNARE helices on the opposite surface via the recently described ‘primary’ interface and that Syt1 is sandwiched between the SNARE complex and the lipid membrane. In fact, the SNARE complex sitting atop Syt1 is positioned such that its C-terminal end is furthest away from the membrane (Fig. 2). Considering that the SNARE proteins are anchored in the opposing membranes via the C-terminal TMDs, this implies that the Syt1 C2B domain creates a steric constraint to delay complete SNARE assembly by holding the membranes apart. In fact, modeling shows zippering beyond the +5 hydrophobic layer.
is disrupted following Ca2+
both as an oligomer or monomer can block full-zippering of the SNARE complex and in fact, rigid-body fitting shows that zipperung beyond layer + 4 would be impeded due to the separation imposed by the Syt1 oligomers. Besides the steric block, the oligomers would also radially retain the zipperung SNAREpins and balance the assembling forces to generate a stable clamp. This oligomeric clamp may be further augmented by an independent C2B (Syt1 or Syt7) bound via the ‘tripartite’ site42 and is positioned to interact with the SV membranes to provide a ‘vice-like’ dual clamp44,48. Note: Only a part of the ring-like oligomer is shown for clarity. It is possible that these Syt1 oligomers dynamically break and re-form, and thus are not always intact. Under these conditions, the dissociated Syt1 monomers would re-orient in the membrane corresponding to the Mg2+-bound state, with partial membrane insertion of the aliphatic loops, but largely still preserving the SNARE clamped architecture. Thus, Syt1 C2B domains, both as an oligomer or monomer can full zipperung of the SNARE complex and vis a vis fusion under resting conditions. This Syt1-SNARE organization is disrupted following Ca2+-influx, as Syt1 reorient into the PM, releasing the attached SNAREpins to fully-assembly and drive fusion

Fig. 6 Molecular model for the regulation of SNARE-mediated fusion by Syt1. Upon docking of the synaptic vesicles, Syt1 C2B domains self-assemble into oligomeric structures triggered by PIP2 on the PM, with the aliphatic loops locating to the dimer interface44,65. Positioning of the SNAREpin on these Syt1 oligomers via the ‘primary’ binding site41,42 would induce steric impediment to the complete zipperung of the SNARE complex and in fact, rigid-body fitting shows that zipperung beyond layer + 4 would be impeded due to the separation imposed by the Syt1 oligomers. Besides the steric block, the oligomers would also radially retain the zipperung SNAREpins and balance the assembling forces to generate a stable clamp. This oligomeric clamp may be further augmented by an independent C2B (Syt1 or Syt7) bound via the ‘tripartite’ site42 and is positioned to interact with the SV membranes to provide a ‘vice-like’ dual clamp44,48. Note: Only a part of the ring-like oligomer is shown for clarity. It is possible that these Syt1 oligomers dynamically break and re-form, and thus are not always intact. Under these conditions, the dissociated Syt1 monomers would re-orient in the membrane corresponding to the Mg2+-bound state, with partial membrane insertion of the aliphatic loops, but largely still preserving the SNARE clamped architecture. Thus, Syt1 C2B domains, both as an oligomer or monomer can full zipperung of the SNARE complex and vis a vis fusion under resting conditions. This Syt1-SNARE organization is disrupted following Ca2+-influx, as Syt1 reorient into the PM, releasing the attached SNAREpins to fully-assembly and drive fusion

(middle portion of c-termini) will be sterically impeded due to the separation imposed by Syt1 (Fig. 6). Hence, the Syt1–SNARE configuration on the membrane could inherently contribute to the strength of an overall fusion clamp under resting (1 mM Mg2+) conditions (Fig. 6). A recent X-ray structure of the Syt1–Cpx–SNARE complex shows that two Syt1 C2B molecules can bind a single SNARE complex—one via the primary site and a second by the Cpx-dependent ‘tripartite’ site42. Thus, we speculate that the above described steric clamp might be augmented by an independent C2B domain (derived from Syt1 or Syt7)44,58 bound to the tripartite interface juxtaposed to the SV membrane (Fig. 6). The resultant ‘dual’ clamp arrangement44,48,58 would trap each SNAREpin in a vice-like grip between the two membranes, held both from above (tripartite C2B on SV) and from below (primary C2B on PM) (Fig. 6).

However, such a clamp is likely to be only meta-stable as the SNAREpins could twist out of position by the radial force generated by SNARE zipperung44. Additionally, this arrangement cannot account for the observed synchronicity and co-operativity of neurotransmitter release59–62. Therefore, while this mechanism may contribute to the overall strength of the fusion clamp, it is unlikely to be the only mechanism at work. So, we envision a concerted mechanism involving the recently described Ca2+-sensitive oligomers of Syt156,63,64, which have been shown to be essential to generate a Ca2+-sensitive fusion clamp under reconstituted conditions38 and for Ca2+-control of vesicular exocytosis in PC12 cells65. Syt1 polymerization is driven by the C2B domain, with the Mg2+/Ca2+-binding aliphatic loops locating to the dimer interface56,63,64. We suspect that under our crystallization conditions, the curved geometry of the LNTs, combined with the C-terminal SNARE interaction stabilizes the membrane-inserted geometry of the Syt1 monomer precluding the oligomerization. But under the physiological configuration, with the relatively planar plasma membrane surface and partially zipperung SNARE complex, the Syt1 oligomerization should be more favorable. In fact, Syt1 oligomers are observed on lipid monolayer surfaces even in the presence of 1 mM free Mg2+56,63.

We posit that Syt1 self-organizes at the site of SV docking triggered by PIP2 clusters on the PM44,56,63 and the SNAREpins are positioned on the Syt1 oligomers via the primary interface, similar to the Syt1 monomer (Fig. 6). Indeed, the ‘primary’ binding site is accessible and free to interact with the SNAREpins in the Syt1 oligomer configuration44,56. Consequently, the Syt1 oligomer could serve as the template to link multiple SNAREpins to enable rapid and co-operative release44. Besides the steric impediment, the Syt1 oligomers will also radially restrain the assembling SNAREpins and symmetrically balance the SNARE assembling forces to create a stable clamp on SV fusion44. The oligomeric clamp could be further reinforced by the dual clamp arrangement44,58 (Fig. 6). Supporting this, recent cryo-electron microscopy analysis in PC12 cells revealed a symmetrical organization of the exocytotic machinery under docked vesicles, likely templated by the Syt1 ring-like oligomers66. It is possible that the C2B oligomers dynamically break and re-form and intact ring-like oligomers are not always present or predominate in vivo. Under these conditions, dissociated monomers by re-orienting into the membrane, could still maintain the half zipperung SNAREpins. Thus, Syt1 organization on the membrane, both as a monomer or an oligomer, lowers the probability of fusion of the docked vesicle in the absence of the Ca2+-trigger (Fig. 6). Likewise, Cpx is an essential component of this fusion clamp. The Cpx central helix is required to generate the ‘tripartite’ binding site for the second C2B domain42 and thus, critical for the ‘dual-clamp’ arrangement44 (Fig. 6). Additionally, the Cpx accessory helix has been shown to directly compete with VAMP2 c-terminus zipperung and thus, stabilize the half zipperung SNAREpins67–69. However, the precise molecular arrangement that enables the accessory-helix based clamp is still unclear.
The Syt–SNARE organization on the LNT surface is disrupted upon Ca\(^{2+}\) addition (Fig. 5). It appears that as the Syt1 C2 domains reorient into the membrane following Ca\(^{2+}\) binding, the associated SNARE complexes are displaced from the primary binding site. Modeling suggests that indeed the Syt1 reorientation could produce a significant steric clash between the SNARE complex bound to the primary site and the vesicle/plasma membranes (Supplementary Fig. 9). Previous fluorescence analyses suggest that some form of Syt1–SNARE interaction is maintained during the Ca\(^{2+}\)-activation process\(^{5,27}\). Given the low signal-to-noise ratio of the EM map, we cannot rule out this possibility, but the precise nature of this interaction remains unclear. Nonetheless, our data argues against the en bloc activation model\(^{41,42,51,52}\) and suggests that a significant conformational rearrangement is associated with the Ca\(^{2+}\)-activation process. Incidentally, the Ca\(^{2+}\)-binding also reorients the Syt1 from the ring oligomer geometry, disrupting the oligomers\(^{56,63}\).

Thus, it emerges that the Ca\(^{2+}\) binding, the Syt1 C2B domain, which is physiologically required for triggering synaptic transmission\(^{21,24,25}\), co-constitutively reorients the Syt1 clamp and liberates the partially assembled SNAREpins to complete zipperig and open the fusion pore cooperatively. Although speculative, the above described model provides a plausible framework to explain the kinetics of Ca\(^{2+}\)-triggered SV fusion in molecular terms. Additional focused high-resolution structural and functional analysis is required to ascertain its relevance.

In summary, our data shows that the divalent cation (Mg\(^{2+}/\) Ca\(^{2+}\)) coupled membrane interaction is a key element of Syt1 function as both a clamp and an activator of SNARE-mediated fusion. Under resting conditions (with Mg\(^{2+}\)), Syt1 locates between the SNARE complex and the plasma membrane creating a SNARE complex-specific steric constraint to prevent full zipperig (Fig. 6). Ca\(^{2+}\)-triggered reorientation into the lipid membrane allows the associated SNAREpins to complete zipperig and trigger fusion. In this manner, Syt1 couples Ca\(^{2+}\)- influx to SNARE-mediated SV fusion and neurotransmitter release.

### Methods

#### Materials

Lipids. 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC), 1,2-dioleoyl-sn-glycero-3-phospho-L-serine (sodium salt) (DOPS). L α-phosphatidylino-sitol-4,5-bisphosphate (Brain, Porcine) (ammonium salt) (PIP2), D-galactosylβ-1,1′ N-nervonoyl-D-erythro-sphingosine (GC) were purchased from Avanti Polar Lipids (Alabaster,AL). Thiol reactive N-ethylmaleimide (EMI) and N-ethylmaleimide-4-aminobenzenesulfonic acid (EMBSA) were synthesized in-house. All oligonucleotides were purchased from Sigma (St. Louis, MO) and self-assembled LNTs were stored at 4°C.

Model membranes and lipid nanotube preparation. Vesicles were prepared with lipid composition of DOPC/DOPS/PIP2 in molar ratio 60/40/6. The lipid stocks were mixed together in chloroform and methanol mix and the solvent was evaporated under N\(_2\) and the samples were vacuum desiccated for 1 hour. The resulting dried lipid film was hydrated for 1 h at room temperature with constant vortexing in the appropriate buffer. For structural studies, the buffer composition was 100 mM Tris-HCl pH 7.4, 1 M KCl, 0.5 mM TCEP, 10% Glycerol. For EM studies the buffer was prepared using multiple freeze-thaw cycles and then extruded (Mini-Extruder, Avanti Polar Lipids) through a 400 nm pore filter (Whatman) for structural studies and a 30 nm pore filter for fluorescence experiments.

To prepare lipid nanotubes (LNT), Galactyl ceramide (GalCer 24:1) was included in the lipid composition. Two different compositions of LNTs were used: DOPC/GC/DOPS/PIP2 and GC/DOPS with lipid molar ratio 40/20/34/6 and 20/80, respectively. The lipids were mixed in chloroform, dried and then hydrated using 20 mM MOPS pH 7.4, 5 mM KCl, 1 mM EDTA, 0.5 mM TCEP buffer. The samples were briefly sonicated using a bath sonicator (Branson Ultrasonics, Danbury, CT) and self-assembled LNTs were stored at 4°C.

### Helical crystallization of Syt1(C2A,B)-SNARE complex. Purified Syt1(C2A,B)-SNARE complexes were made up to a stock concentration to ~40 μM so that the final protein and salt concentration (~25 mM KCl) were similar under all experimental conditions. Protein dilution was performed in low-salt buffer (20 mM MOPS pH 7.4, 5 mM KCl, 1 mM EDTA, 0.25% (w/v) Trehalose (Sigma), 0.5 mM TCEP and mixture of MgCl\(_2\) or CaCl\(_2\) to provide a 1 mM final concentration of free ions (based on MaxChelator online calculator). For the liposome samples, pre-formed liposomes (~0.5 mM lipids) were applied to glow-discharged grids and then flash-frozen on tip of a Fe(III) protein and KCl concentration were ~4 μM and ~30 mM respectively. After 30 min incubation at room temperature negative staining samples were negatively stained or flash-frozen on Vitrobot Mark III. Helical crystals were observed at 100 mK KCl and confirmed by negative staining microscopy, but those samples showed high sensitivity to the freezing process since most of the tubes in the Cryo-EM samples were naked.

Fluorescent labeling. Syt1(C2A,B) constructs with single cysteine residues (MY73C, S323C, V304C and I367C) were mixed and incubated with a 10-fold molar excess of the IAEDANS dye (freshly prepared) in HEPES buffer. Prior to the addition of the IAEDANS dye, the protein concentration was determined by Bradford assay using BSA as a standard. Labeling efficiency was ~85% in all cases.

Fluorescent measurements. Labelled protein was mixed with liposomes in 1:1000 molar ratio (0.5 μM protein: 0.5 mM lipids) and incubated with 1 μM TCEP and allowed to equilibrate in the lysis buffer with 0.1% Triton X-100. Beads were harvested and washed with lysis buffer with 0.1% Triton X-100 and incubated with the lysis buffer for 30 min before centrifugation and 10 μg/mL RNase A and 10 μg/mL DNase I. The supernatant was loaded onto Ni-NTA beads (Thermofisher Scientific, Waltham, MA). The Syt1(C2A,B)-SNARE complex was eluted in the lysis buffer supplemented with 30 mM Imidazole. The Syt1–SNARE complex was eluted at room temperature by incubating 30 min in a 25°C water bath. Steady-state fluorescence measurements were made at 20°C using a Cary Eclipse spectrophotometer (Varian, CA) overnight to allow for complete binding. Prior to the fluorescence measurements, the samples were brought up to room temperature by incubating 30 min in 25°C water bath. Steady-state fluorescence measurements were made at 20°C using a Cary Eclipse spectrophotometer (Varian, CA) overnight to allow for complete binding. Prior to the fluorescence measurements, the samples were brought up to room temperature by incubating 30 min in 25°C water bath.
Electron microscopy sample preparation and data acquisition. Negative staining of samples was done using home-made carbon coated copper grids (400 mesh). Vesicles samples were stained right after the incubation time, lipid nanotubes samples were applied onto glow-discharged grids for 1 min. Both samples were stained with uranyl acetate solution (1% w/v) and air dried. Negatively stained samples were imaged using FEI Tecnai T12 operated at an acceleration voltage of 120 kV, equipped with LaB6. Micrographs were recorded using 4k x 4k Gatan UltraScan 4000 CCD camera with an effective magnification of ×42000 and under low-dose conditions (~25e−/Å²) with defocus range of 1–2.5μm. Micrographs were binned by a factor 2 at a final sampling of 5.6 Å per pixel. For the Syt(C2A)SNARE decorated vesicles, the samples on the grids were pre-washed with buffer without trehalose, blotted with paper towel in an instantaneously adding 2.5 μl of the same buffer, transferred into a Vitrobot Mark III held at 20°C with 100% humidity, blotted for ~5 s and plunge-frozen in liquid ethane cooled by liquid nitrogen. For the Syt(C2A)SNARE decorated lipid nanotubes, the C-flat holey carbon grids (Protophics, CF-1.2/1.3-T) were plasma cleaned with Solarus Model 950 plasma cleaning system (Gatan) using O2/Ar mixture for 15 s. Then 2.5 μl of the sample was applied to the grids, blotted for ~7 s and plunge-frozen in liquid ethane cooled by liquid nitrogen using a Vitrobot Mark III at 20°C with 100% humidity. Images were recorded on FEG equipped FEI Tecnai F20 microscope operating at 200 kV at a nominal magnification of 25,000 on Gatan K2 Summit direct electron microscopy sample preparation and data acquisition, with specimen on holey carbon grids (Protochips, CF-1.2/1.3–T) were manually selected using boxer from EMAN1 package75 implemented in SPARX.

Cryo-EM data processing. Movies were motion-corrected, dose-weighted and summed using IMAGIC. Micrographs were evaluated for presence of helical structures with similar dimensions as SNARE-pins (rise, φ = 7.35 Å and twist, Δφ = 7.35 Å) and using SPIDER76. Parameters that gave reconstruction with pronounced, elongated densities with outer diameter ~41 nm showed helical diffraction pattern corresponding to homogenous helical crystal and was suitable for obtaining real-space helical parameters as rise (Az) and twist (Δϕ). Calculated possible parameters of helical lattice were checked by low resolution IHRSR reconstruction using SPARX76. Briefly, using boxes from EMAN1 package75 implemented in SPARX particles were extracted with 97% overlap and then times binned and used in SPARX-IHRSR reconstruction using featureless hollow cylinder created using SPIDER76. Parameters that gave reconstruction with pronounced, elongated densities with similar dimensions as SNAP-25s (rise, Δz = 7.35 Å and twist, Δϕ = 7.848°, 14–14 start helix) were used for further high-resolution reconstruction in RELION 2.078. For the final reconstruction all tubes with an outside diameter —41 nm (as identified above) were manually selected using e2helixboxer.py from EMAN279 and boxes were extracted with an inter-box distance of 3 asymmetrical units (96% overlap) using RELION 2.0. Defocus values were estimated by CTFFIND4.0. The resulting 2844 particles subjected to 2D classification and particles belonging to blurry classes were discarded, leaving 2082 particles that were further subjected to 3D refinement using a featureless hollow cylinder as an initial reference. The resulting 3D volume was post-processed using soft defocus with a B-factor of 300. The average resolution estimated using a gold standard FSC = 0.143 was 10.4 Å (Supplementary Fig. 10). The local protein resolution estimated using blocco program (BSOFT) was in the range of 8–12 Å (Supplementary Fig. 10). Only one handedness gave density consistent with the X-ray structures of SNAREs, Synaptotagmin C2A and C2B domains. The selected handedness defines a left-handed helix on the tube’s surface (Δϕ = −78.48°, Δz = 7.35 Å).

3D model refinement and model building. First fitting was performed using separate crystal structures of C2A, C2R, and the SNARE-pin obtained from a Mg2+–binding motif of synaptotagmin is required for synaptic transmission in vivo. Nature 418, 336–340 (2002).

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References

1. Sudhof, T. C. Neurotransmitter release: the last millisecond in the life of a synaptic vesicle. Neuron 80, 673–690 (2013).
2. Sudhof, T. C. & Rothman, J. E. Membrane fusion: grappling with SNAREs and SM proteins. Science 323, 474–477 (2009).
3. Rizo, J. & Xu, J. The synaptic vesicle release machinery. Annu. Rev. Biophys. 44, 339–367 (2015).
4. Jahn, R. & Fasshauer, D. Molecular machines governing exocytosis of synaptic vesicles. Nature 490, 201–207 (2012).
5. Bai, J., Tucker, W. C. & Chapman, E. R. Pip2 increases the speed of response of synaptotagmin and steers its membrane-penetration activity toward the plasma membrane. Nat. Struct. Mol. Biol. 11, 36–44 (2004).
6. Parisetto, D., Malsam, J., Scheutzow, A., Krause, J. M. & Söllner, T. H. SNAP-25–SNARE assembly by Munc18-1 requires previous vesicle docking by synaptotagmin 1. J. Biol. Chem. 287, 31041–31049 (2012).
7. Honigmann, A. et al. Phosphatidylinositol 4.5-bisphosphate clusters act as molecular beacons for vesicle recruitment. Nat. Struct. Mol. Biol. 20, 679–686 (2013).
8. Park, Y. et al. Controlling synaptotagmin activity by electrostatic screening. Nat. Struct. Mol. Biol. 19, 991–999 (2012).
9. Wang, Z., Liu, H., Gu, Y. & Chapman, E. R. Reconstituted synaptotagmin I mediates vesicle docking, priming, and fusion. J. Cell Biol. 195, 1159–1170 (2011).
10. Littleton, J. T., Stern, M., Perin, M. & Bellen, H. J. Calcium dependence of neurotransmitter release and rate of spontaneous vesicle fusions are altered in Drosophila synaptotagmin mutants. Proc. Natl. Acad. Sci. USA 91, 10888–10892 (1994).
11. Xu, J., Pang, Z. P., Shin, O. H. & Sudhof, T. C. Synaptotagmin-1 functions as a Ca2+-sensor for spontaneous release. Nat. Neurosci. 12, 759–766 (2009).
12. Brose, N., Petrenko, A. G., Sudhof, T. C. & Jahn, R. Synaptotagmin: a calcium sensor on the synaptic vesicle surface. Science 256, 1021–1025 (1992).
13. Fernández-Chacón, R. et al. Structure/function analysis of Ca2+-binding to the C2A domain of synaptotagmin 1. J. Neurosci. 22, 8438–8446 (2002).
14. Geppert, M. et al. Synaptotagmin I: a major Ca2+-sensor for transmitter release at a central synapse. Cell 79, 717–727 (1994).
15. Littleton, J. T., Stern, M., Schulze, K., Perin, M. & Bellen, H. J. Mutational analysis of Drosophila synaptotagmin demonstrates its essential role in Ca2+-activated neurotransmitter release. Cell 74, 1125–1134 (1993).
16. Fusion, K. L., Montes, M., Robert, J. J. & Sutton, R. B. Structure of human synaptotagmin 1 C2AB in the absence of Ca2+ reveals a novel domain association. Biochemistry 46, 13041–13048 (2007).
17. Geppert, M. R. B., Davletov, B. A., Berghuis, A. M., Sudhof, T. C. &Sprang, S. R. Structure of the first C2 domain of synaptotagmin 1: a novel Ca2+-phospholipid-binding fold. Cell 80, 929–938 (1995).
18. Rizo, J. & Sudhof, T. C. C2 domains, structure and function of a universal Ca2+-binding domain. J. Biol. Chem. 273, 15879–15882 (1998).
19. Herrick, D. Z., Sterbing, S., Rasch, K. A., Hinderliter, A. & Ca2+ as a trigger for vesicle fusion. J. Neurosci. 23, 14437–14430 (2003).
20. Mackler, J. M., Drummond, J. A., Loewen, C. A., Robinson, S. B. & Schwarz, T. L. Synaptotagmins I and IV mediate synaptic vesicle release by binding the C2A–phospholipid binding motif of synaptotagmin is required for synaptic transmission in vivo. Nature 418, 340–344 (2002).

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27. Stevens, C. F. & Sullivan, J. M. The synaptotagmin C2A domain is part of the calcium sensor controlling fast synaptic transmission. Neuron 39, 299–308 (2003).
28. Striegel, A. R. et al. Calcium binding by synaptotagmin’s C2A domain is an essential element of the electrostatic switch that triggers synchronous synaptic transmission. J. Neurosci. 32, 1253–1260 (2012).
29. Yoshuhara, M., Guan, Z. & Littleton, J. T. Differential regulation of synchronous versus asynchronous neurotransmitter release by the C2 domains of synaptotagmin 1. Proc. Natl Acad. Sci. USA 107, 14869–14874 (2010).
30. Bai, H. et al. Different states of synaptotagmin regulate evoked versus spontaneous release. Nat. Commun. 7, 10971 (2016).
31. Evans, C. S. et al. Functional analysis of the interface between the tandem C2 domains of synaptotagmin-1. Mol. Biol. Cell 27, 979–989 (2016).
32. Bacaj, T. et al. Synaptotagmin-1 and synaptotagmin-7 trigger synchronous and asynchronous phases of neurotransmitter release. Neuron 80, 947–959 (2013).
33. Huntsworth, S. & Littleton, J. T. A complex fusion clamp regulates spontaneous neurotransmitter release and synaptic growth. Nat. Neurosci. 10, 1235–1237 (2007).
34. Yang, X., Cao, P. & Sudhof, T. C. Deconstructing complexin function in activating and clamping Ca^{2+}-triggered exocytosis by comparing knockout and knockdown phenotypes. Proc. Natl Acad. Sci. USA 110, 20777–20782 (2013).
35. Cho, B. W. et al. Genetic analysis of the Complexin trans-clamping model for cross-linking SNARE complexes in vivo. Proc. Natl Acad. Sci. USA 111, 10317–10322 (2014).
36. Schaub, J. R., Lu, X., Doneske, B., Shin, Y. K. & McNew, J. A. Hemifusion arrest by complexin is relieved by Ca^{2+}–synaptotagmin 1. Nat. Struct. Mol. Biol. 13, 748–750 (2006).
37. Chicka, M. C., Hui, E., Liu, H. & Chapman, E. R. Synaptotagmin arrests the SNARE complex before triggering fast, efficient membrane fusion in response to Ca^{2+}. Nat. Struct. Mol. Biol. 15, 827–835 (2008).
38. Ramakrishnan, S. et al. Synaptotagmin oligomers are necessary and can be sufficient to form a Ca^{2+}–sensitive fusion clamp. FEBS Lett. 593, 154–162 (2019).
39. Malam, J. et al. Complexin arrests a pool of docked vesicles for fast Ca^{2+}–dependent release. EMBO J. 31, 3270–3281 (2012).
40. Diao, J. et al. Synaptic proteins promote calcium-triggered fast transition from point contact to full fusion. Elife 1, e00109 (2012).
41. Zhou, Q. et al. Architecture of the synaptotagmin-SNARE machinery for neuronal exocytosis. Nature 525, 62–67 (2015).
42. Zhou, Q. et al. The primed SNARE-complex-synaptotagmin complex for neuronal exocytosis. Nature 548, 420–425 (2017).
43. Gipson, P. et al. Morphologies of synaptic protein membrane fusion interfaces. Proc. Natl Acad. Sci. 114 9110–9115 (2017).
44. Rothman, J. E., Krishnakumar, S. S., Gruskin, K. & Pincet, F. Hypothesis – bioassayed rings assemble, clamp, and release SNAREpins for synaptic transmission. FEBS Lett. 591, 3493–3490 (2017).
45. Chapman, E. R. A Ca^{2+} Sensor for Exocytosis. Trends Neurosci. 41, 327–330 (2018).
46. Rizo, J. Mechanism of neurotransmitter release coming into focus. Protein Sci. 27, 1364–1391 (2018).
47. Brunger, A. T., Chou, U. B., Lai, Y., Leitz, J. & Zhou, Q. Molecular Mechanisms of Fast Neurotransmitter Release. Annu Rev. Biophys. 47, 469–497 (2018).
48. Brunger, A. T., Leitz, J., Zhou, Q., Choi, U. B. & Lai, Y. Ca^{2+}–Triggered Synaptic Vesicle Fusion Initiated by Release of Inhibition. Trends Cell Biol. 28, 631–645 (2018).
49. Kulkarni, V. S., Anderson, W. H. & Brown, R. E. Bilayer nanotubes and helical NATURE COMMUNICATIONS | (2019) 10:2413 | https://doi.org/10.1038/s41467-019-10391-x | www.nature.com/naturecommunications 50. Wilson-Kubalek, E. M., Brown, R. E., Celia, H. & Milligan, R. Lipid nanotubes. Nat. Struct. Mol. Biol. 15, 266–274 (2005).
51. Bollmann, J. H., Sakmann, B. & Borst, J. G. Calcium sensitivity of glutamate release in a calyx-type terminal. Science 293, 953–957 (2000).
52. Kaeser, P. S. & Regehr, W. G. Molecular mechanisms for synchronous, asynchronous, and spontaneous neurotransmitter release. Annu. Rev. Physiol. 76, 333–363 (2014).
53. Wang, J. et al. Calcium sensitive ring-like oligomers formed by synaptotagmin. Proc. Natl Acad. Sci. USA 111, 13966–13971 (2014).
54. Wang, J. et al. Circular oligomerization is an intrinsic property of synaptotagmin. Elife 6 (2017).
55. Bello, O. D. et al. Synaptotagmin oligomerization is essential for calcium control of regulated exocytosis. Proc. Natl Acad. Sci. USA 115, E7624–E7631 (2018).
56. Li, X. et al. Symmetrical organisation of proteins under docked synaptic vesicles. FEBS Lett. 593, 144–153 (2019).
57. Giraudon, C. G., Eng, W. S., Melia, T. J. & Rothman, J. E. A clamping mechanism involved in SNARE-dependent exocytosis. Science 313, 676–680 (2006).
58. Krishnakumar, S. S. et al. A conformational switch in complexin is required for synaptotagmin to trigger synaptic fusion. Nat. Mol. Biol. 18, 934–940 (2011).
59. Kummel, D. et al. Complexin cross-links prefusion SNAREs into a zigzag array. Nat. Mol. Biol. 18, 927–933 (2011).
60. Krishnakumar, S. S. et al. Conformational dynamics of calcium-triggered activation of fusion by synaptotagmin. Biophys. J. 105, 2507–2516 (2013).
61. Wang, S., Li, Y. & Ma, C. Synaptotagmin-1 C2B domain interacts simultaneously with SNAREs and membranes to promote membrane fusion. Elife 5, 1–21 (2016).
62. Mastronarde, D. N. Automated electron microscope tomography using robust prediction of specimen movements. J. Struct. Biol. 152, 36–51 (2005).
63. Grant, T. & Grigorieff, N. Automatic estimation and correction of anisotropic magnification distortion in electron microscopes. J. Struct. Biol. 192, 204–208 (2015).
64. Behrmann, E. et al. Real-space processing of helical filaments in SPARX. J. Struct. Biol. 177, 302–313 (2012).
65. Ludtke, S. J., Baldwin, P. R. & Chiu, W. EMAN: semiautomated software for high-resolution single-particle reconstructions. J. Struct. Biol. 128, 82–99 (1999).
66. Shaikh, R. T. et al. SPIDER image processing for single-particle reconstruction of biological macromolecules from electron micrographs. Nat. Protoc. 3, 1941–1974 (2008).
67. He, S. & Scheres, S. H. W. Helical reconstruction in RELION. J. Struct. Biol. 198, 163–176 (2017).
68. Scheres, S. H. W. RELION: implementation of a Bayesian approach to cryo-EM structure determination. J. Struct. Biol. 180, 519–530 (2012).
69. Tang, G. et al. EMAN2: an extensible image processing suite for electron microscopy. J. Struct. Biol. 157, 38–46 (2007).
70. Rohou, A. & Grigorieff, N. CTFFIN4: Fast and accurate defocus estimation from electron micrographs. J. Struct. Biol. 192, 216–221 (2015).
71. Cardone, G., Heymann, J. B. & Steven, A. C. One number does not do all: mapping local variations in resolution in cryo-EM reconstructions. J. Struct. Biol. 184, 226–236 (2013).
72. Pettersen, E. F. et al. UCSF Chimera—a visualization system for exploratory research and analysis. J. Comput. Chem. 25, 1605–1624 (2004).
73. Wrighgers, W. Conventions and workflows for using Situs. Acta Crystallogr. Sect. D. Biol. Crystallogr. 68, 343–351 (2012).

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Author contributions
K.G. designed experiments, collected, analyzed/interpreted CryoEM and fluorescence data; J.W. collected/analyzed CryoEM data; J.C. provided new reagents and collected fluorescence data; J.E.R., C.E.S., S.S.K. designed the experiments, provided supervision, analyzed/interpreted the data and wrote the manuscript. All authors read and revised the manuscript.

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