Resolving kinetic intermediates during the regulated assembly and disassembly of fusion pores

Debasis Das1,2, Huan Bao1,2, Kevin C. Courtney1, Lanxi Wu1 & Edwin R. Chapman1*

The opening of a fusion pore during exocytosis creates the first aqueous connection between the lumen of a vesicle and the extracellular space. Soluble N-ethylmaleimide-sensitive factor attachment protein receptors (SNAREs) mediate the formation of these dynamic structures, and their kinetic transitions are tightly regulated by accessory proteins at the synapse. Here, we utilize two single molecule approaches, nanodisc-based planar bilayer electrophysiology and single-molecule FRET, to address the relationship between SNARE complex assembly and rapid (micro-millisecond) fusion pore transitions, and to define the role of accessory proteins. Synaptotagmin (syt) 1, a major Ca2+-sensor for synaptic vesicle exocytosis, drove the formation of an intermediate: committed trans-SNARE complexes that form large, stable pores. Once open, these pores could only be closed by the action of the ATPase, NSF. Time-resolved measurements revealed that NSF-mediated pore closure occurred via a complex ‘stuttering’ mechanism. This simplified system thus reveals the dynamic formation and dissolution of fusion pores.
The mechanism by which proteins catalyze the fusion of cellular membranes remain a central unanswered question in cell biology. In all eukaryotic cells, the majority of fusion events are mediated by soluble N-ethylmaleimide-sensitive factor attachment protein receptor (SNARE) proteins. The cytoplasmic domains of vesicular SNAREs (v-SNAREs) bind to cognate domains on target membrane SNAREs (t-SNAREs), forming trans-complexes that are necessary and sufficient for fusion. Here we focus on the SNAREs that mediate synaptic vesicle (SV) exocytosis in nerve terminals. A key intermediate in this pathway is called the fusion pore, which represents the first aqueous connection between the lumen of a secretory vesicle and the cell exterior. These are nanometer-scale transient structures, lasting only milliseconds before they either close or dilate as the vesicle membrane collapses into the plasmalemma. Their structure remains unknown, recent studies indicate that exocytotic pores are composed of both the transmembrane domains (TMDs) of SNARE proteins and phospholipids. Because they are crucial intermediates, fusion pores are a focal point for the action of regulatory proteins. In nerve terminals, fusion pore opening is thought to be triggered by the binding of Ca^{2+} ions to synaptotagmin 1 (syt1). Following complete fusion, v- and t-SNAREs lie within the plasma membrane, in cis-complexes, and must be disassembled to allow for segregation of SNAREs for future rounds of fusion. Disassembly occurs when the soluble factor, a-SNAP, binds cis-SNARE complexes and recruits the AAA+ ATPase, NSF, to hydrolyze ATP by NSF drives disassembly. While NSF clearly disassembles cis-SNARE complexes, it remains unclear as to whether the trans-SNARE complex also serves as a substrate. This is a crucial question; if trans-SNARE complexes are substrates, then the action of NSF should result in the closure of fusion pores. This would have ramifications in, for example, kiss-and-run exocytosis.

The regulated assembly and disassembly of fusion pores have been difficult to study because they are short-lived, so it has been challenging to capture intermediate states. Kinetic analysis of these transitions would provide crucial information concerning the action of regulatory factors that help to assemble, and potentially disassemble, pores. To date, most mechanistic studies of reconstituted fusion machines rely on assays with limited time resolution (i.e., seconds to minutes), thus obscuring dynamics.

To delve into the relationship between the status of trans-SNARE complexes and fusion pore dynamics, we developed a nanodisc (ND)—black lipid membrane (BLM) system. Trans-SNARE pairing results in the formation of individual recombiant fusion pores that can be studied for extended periods because NDs trap pores in intermediate states. The strength of this approach is that pores can be interrogated electrophysiologically, thus affording microsecond time resolution to reveal kinetic intermediates that could not be previously observed in defined systems.

Here we combine ND-BLM experiments and single-molecule fluorescence resonance energy transfer (smFRET) measurements to determine the impact of key regulatory factors on the structure and kinetic properties of fusion pores and trans-SNARE complexes. The first goal was to assess whether Ca^{2+}–syt1 directly regulates SNAREs to trigger pore opening. Indeed, Ca^{2+} and syt1 affected not only the occurrence but also the size and kinetic stability of purely recombinant fusion pores. The observation that Ca^{2+}–syt1 dilates pores prompted experiments to address the state of the trans-SNARE complexes that formed them. These experiments demonstrated that Ca^{2+}–syt1 mediates the formation of an intermediate: committed trans-SNARE complexes that can only be disassembled by the action of NSF. The time resolution afforded by the ND-BLM system resolved short-lived fusion pore states and indicated that disassembly occurred via a complex kinetic mechanism. These fusion pore intermediates correspond with trans-SNARE complex status and reveal coupling of SNARE complex assembly with fusion pore transitions.

Results

A Ca^{2+} switch converts syt1 from fusion clamp to activator. In cells, manipulation of a number of syt isoforms alters the occurrence and properties of individual fusion pores, but it was not clear whether these were direct or indirect effects. To address this question, we used our recently described ND-BLM electrophysiology assay to study pores with microsecond time resolution. We reconstituted full-length syt1, along with syb2, into NDs (Fig. 1a) and t-SNAREs (syntaxin-1A and SNAP-25B heterodimers) into BLMs (Supplementary Fig. 1a), pore formation between the ND and BLM was monitored via the currents that were detected (Fig. 1b). Three kinds of ND preparations were used: ND3s, ND3l, and ND9L, where 3 and 9 refer to the syb2 (and syt2) copy number per ND and subscripts S and L refer to the small (13 nm) and large (30 nm) diameters of the NDs. We first describe representative traces, followed by quantitative analysis of the pores that were formed.

We began with ND3s and confirmed our earlier observation that these NDs gave rise to pores that remained mostly in the closed state but transiently flickered open to yield small, transient currents; Ca^{2+} had no obvious effect (Fig. 1b). When syt1 was co-reconstituted with syb2 into ND3s, the pore remained mostly in the closed state. However, addition of Ca^{2+} (500 µM [Ca^{2+}]_free in all experiments, unless otherwise indicated) resulted in the formation of large, stable pores (Fig. 1b). We note that the properties of individual pores did not differ at the beginning, middle, or end of a recording, but all pores closed within ~90 min. Once the terminal closure occurred, there were no further openings or flickers. Closure might involve reversion to a hemifused state. When syt1 alone was reconstituted (three copies) into NDs (ND0), pores failed to form either in the presence or absence of Ca^{2+} (Supplementary Fig. 1b), confirming that pore formation required trans-SNARE pairing. For the Ca^{2+}-free conditions, BAPTA was used to chelate any residual Ca^{2+} present in the buffers; in most of our experiments, we subsequently added Ca^{2+} to yield the indicated [Ca^{2+}]_free (note: we confirmed that BAPTA binds Ca^{2+} with a stoichiometry of 1:1 in Supplementary Fig. 1c).

Next, we examined ND3L (Fig. 1a) but were unable to detect pore formation in the absence of syt1 (Fig. 1c). These finding suggest that v-SNARE density (ND3S, 0.022 syb2/nm², ND3L: 0.0042 syb2/nm²) and not just copy number is a crucial parameter for pore formation. Indeed, a recent modeling study suggested that restricting the mobility of SNAREs, to increase their relative density, facilitates pore formation. In sharp contrast, when syt1 was co-reconstituted into ND3s, robust pore formation occurred upon addition of Ca^{2+} (Fig. 1c); however, these pores exhibited rapid, dramatic flickering behavior.

We then assessed ND9L (Fig. 1a) lacking syt1; these NDs yielded pores that remained mostly in the open state but transiently flickered closed (Fig. 1d). In contrast, ND3s, which has a somewhat higher syb2 density, gave rise to pores that were mostly in closed state but transiently flickered open. We conclude that the higher syb2 copy number in ND9L serves to hold the pore in a stable open state. Strikingly, inclusion of syt1 in ND9L, potently inhibited pore formation in the absence of Ca^{2+} (Fig. 1d). These conditions appear to recapitulate the clamping activity of syt1 that has been observed in neurons. Subsequent addition of Ca^{2+} triggered the opening of relatively stable, large pores (Fig. 1d).
To conduct quantitative analysis, we carried out 20–40 trials under each of the conditions described above. We first calculated the frequency of pore formation; these findings are represented as "percentage of occurrence" (Fig. 1e). This analysis demonstrated that the occurrence of pores formed by SNAREs alone was unaffected by Ca\(^{2+}\) (Fig. 1e). Moreover, inclusion of syt1 in both ND3S and ND9L reduced the occurrence of pores in the absence of Ca\(^{2+}\) (Fig. 1e). Interestingly, this clamping activity was the most apparent using ND9L, a condition that most closely mimics the SNARE TMDs, the pore diameter values, based on conductance measurements, are only approximations. For ND3S and ND9L in the absence of syt1 and Ca\(^{2+}\), the mean conductance was 170 ± 32 and 299 ± 74 pS, respectively, in the absence of Ca\(^{2+}\) and in the presence (+) of Ca\(^{2+}\) and in the presence (+) and absence (−) of syt1 were compared. Three independent sets of NDs of each type were used, and the total number of measurements obtained under each condition (n) is indicated. Pearson’s \(\chi^2\) analysis of pores formed by ND3S, ND3L, and ND9L was performed; *p < 0.05, **p < 0.001.

Fig. 1 Syt1 regulation of single fusion pores measured via planar lipid bilayer electrophysiology. a Illustration of the nanodisc-black lipid membrane (ND-BLM) system, drawn to scale, indicating the different ND preparations used in this study. The number indicates the syb2/syt1 copy number (1:1) per ND; the subscript S stands for small, 13 nm, NDs and the subscript L stands for large, 30 nm, NDs. Traces of single pores with/without syt1 are shown for ND3S (b), ND3L (c), and ND9L (d). In each trace, minus (−) Ca\(^{2+}\) contains 1 mM BAPTA (in all bilayer recording experiments in this study) and plus (+) Ca\(^{2+}\) contains 500 μM [Ca\(^{2+}\)]\(_{tot}\). Closed (C) and open (O) states are shown; the current and time scale, for all traces, is shown in the inset in b. O1 and O2 indicate open state currents obtained before and after addition of Ca\(^{2+}\), respectively. e Fraction of trials in which a fusion pore was detected, plotted as percentage of occurrence. ND3S, ND3L, and ND9L, in the presence (+) or absence (−) of Ca\(^{2+}\) and in the presence (+) and absence (−) of syt1 were compared. Three independent sets of NDs of each type were used, and the total number of measurements obtained under each condition (n) is indicated. Pearson’s \(\chi^2\) analysis of pores formed by ND3S, ND3L, and ND9L was performed; *p < 0.05, **p < 0.001.
distribution to shorter life times in BAPTA while addition of Ca\(^{2+}\) drastically increased the open life time (Fig. 2b, upper panel). In the case of ND9\(_L\) pores, the findings were more complicated (Fig. 2b, lower panel). Here Ca\(^{2+}\) had a small effect on individual ND9\(_L\) pores lacking syt1, and inclusion of syt1 shifted the distribution to shorter life times in BAPTA. However, as in the case with ND3\(_S\)/syt1, addition of Ca\(^{2+}\) markedly increased the open life time of ND9\(_L\)/syt1 pores (Fig. 2b, lower panel). Together, these results demonstrate that syt1 has dramatic, direct effects on the kinetic properties of individual, recombinant, fusion pores. Ca\(^{2+}\)/syt1 not only increases the size of pores but also promotes membrane fusion by significantly stabilizing the open state.

In order to gain insights into the number of kinetic steps during pore transitions, the cumulative distribution functions (CDFs) for the opening and closure of individual pores were fitted with single or multiple exponential functions (Supplementary Figs. 4–6). CDFs of the closed and open time distributions reflect the kinetics of pore opening and closure, respectively. In the absence of syt1, pore opening was described by multiple exponentials for both the ND3\(_S\) and ND9\(_L\) (Fig. 2c); inclusion of apo-syt1 had little effect (Fig. 2c). In sharp contrast, Ca\(^{2+}\)/syt1 resulted in mainly single exponential kinetics for pore opening, despite the SNARE copy number heterogeneity in our ND preparations (Fig. 2c and Supplementary Figs. 4–6). These findings indicate that Ca\(^{2+}\)/syt1 drives fusion pores into a single open state with little or no involvement of intermediates.

Regarding pore closure, in the absence of syt1, single exponential kinetics were observed for ND3\(_S\), while ND9\(_L\) pores followed multi-exponential kinetics. In contrast, inclusion of

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**Fig. 2 Quantification of syt1-regulated fusion pore properties.**

a Cumulative distribution functions (CDF) of single-channel conductances across different trials for each experimental condition. b Open dwell time histograms for ND3\(_S\) and ND9\(_L\), plus or minus syt1 and Ca\(^{2+}\), are shown. c Pie diagrams showing the number of exponentials required to fit the open- and closed-state CDFs for individual pores under the indicated conditions. Details are described in Supplementary Fig. 4. d Conductance values (γ) (upper panel), open time (middle panel), and percentage of occurrence (lower panel) at each indicated [Ca\(^{2+}\)]\(_{0}\) are plotted. n = 16, 10, 10, 15; three different sets of ND preparations were used. In the open time plot, mean values for the open-state dwell times of individual traces were quantified. Error bars indicate SEM. e Percentage of occurrence of pore formation, plotted for WT syt1 versus a Ca\(^{2+}\) ligand mutant (CLM), D230/232N/D363/365N, that fails to bind Ca\(^{2+}\) via either C2 domain. The number of independent trials (n) are indicated in e. Pearson’s χ² analysis was performed; ***p < 0.001.
apo-syt1 resulted in predominantly single exponential kinetics for both ND3a and ND9, even though the number of SNAREs remained the same. Finally, in the presence of syt1 and Ca^{2+}, pore closure for ND3a and ND9 was best described by multiple exponentials (Fig. 2c and Supplementary Figs. 4–6). These findings indicate that pore closure, in the presence of Ca^{2+}/syt1, occurs via a complex multi-state mechanism.

Ca^{2+} dependencies of fusion pore parameters. We conducted [Ca^{2+}] dose–response experiments to determine whether the effects of Ca^{2+} on fusion pore conductance, opening, and dilation described in Fig. 2 could be dissociated from one another. Representative traces of individual ND3/syt1 pores as a function of [Ca^{2+}]free are shown in Supplementary Fig. 7a. The Ca^{2+}-dependent effect of syt1 on pore conductance was saturated at 125 μM [Ca^{2+}]free (Fig. 2d, top panel); we were unable to reliably determine conductance values at lower [Ca^{2+}], so these data were not fitted to estimate an EC_{50} or Hill slope. The Ca^{2+} titration measurements were carried out using the same Ca^{2+} stocks and buffers. The different Ca^{2+} requirements for each of the fusion pore parameters suggest that different Ca^{2+}-binding sites in syt1 (which binds 4–5 Ca^{2+} ions)^{32,33} might subserve distinct functions during exocytosis.

As a control, we tested a mutant form of syt1 in which two acidic Ca^{2+} ligands, in each C2-domain of syt1, were neutralized by substitution with asparagine residues, thus abolishing Ca^{2+}-binding activity^{34,35}. This Ca^{2+} ligand mutant was unable to couple Ca^{2+} to fusion pore opening (Fig. 2c).

Syt1–PIP2 interactions stabilize fusion pores. It is established that syt1 must penetrate membranes that harbor acidic phospholipids in order to drive membrane fusion in vitro^{36} and to trigger exocytosis in cells^{26,28,37}. There is evidence that PIP2, which is localized to the plasma membrane, interacts with syt1 under resting conditions to steer the Ca^{2+}-triggered membrane penetration of its C2-domains toward the target membrane^{38} to regulate fusion in vitro^{36}. Indeed, PIP2 plays an essential role in exocytosis in neuroendocrine cells^{39,40} and appears to have a key role in at least some modes of SV release^{41}. However, before the development of the fully defined ND-BLM system, it had not been possible to directly explore the role of PIP2 on the biophysical properties of individual fusion pores, as previous work relied on cell-based measurements. We therefore carried out ND-BLM measurements using BLMs with and without PIP2. These experiments, using ND3/syt1, revealed that, in the absence of PIP2, pores became less stable (Fig. 3a–c), with the appearance of a partially open state (Fig. 3a). Similar results were obtained using ND3/syt1 but with a more prominent partially open state (Supplementary Fig. 7b, c). As a control, PIP2 had no effect on pores formed by ND3 in the absence of syt1 (Supplementary Fig. 7d, e).

These experiments confirm that syt1–lipid interactions play a key role in the regulation of fusion pores.

**Ca^{2+}•syt1 commits trans-SNARE complexes.** We next turned to the question of how syt1 affects trans-SNARE complex assembly^{42}, using a previously described smFRET approach^{19}. It was not technically feasible to conduct smFRET using v-SNARE NDs bound to t-SNARE BLMs, so we used v- and t-SNARE NDs. The strength of this approach is that the assembly state of full-length, membrane-embedded SNAREs can be assessed, but we note that the t-SNARE NDs are more constrained as compared to the BLM. Nonetheless, this system revealed that apo- and Ca^{2+}-bound syt1 have distinct, direct effects on the assembly of trans-SNARE complexes.

Syb2 was labeled with a donor fluorophore (cy3), and syntaxin-1A was labeled with an acceptor (cy5). Monitoring multiple trans-SNARE complexes by smFRET is extremely challenging, so we used ND1 for these experiments. For each SNARE, two positions were selected, based on the crystal structure of the cis-complex^{39,44}: one pair at layer +7 near the N-termini of the SNARE motifs (N-N FRET pair), and the other between layer +4 and +5 near the C-termini of the SNARE motifs (C-C FRET; Supplementary Fig. 9a; Fig. 4). Labeled trans-SNARE complexes were first formed using v- and t-SNARE NDs in the absence of syt1. Under these conditions, the FRET ratio (R) distribution of the N-terminal pair peaked at ~0.7, while this value was only 0.2 for the C-C pair; Ca^{2+} had no effect (Fig. 4). Thus single trans-SNARE complexes formed between two NDs in the absence of syt1 but exist in a partially zipped state: the N-termini of the SNARE motifs assembled together, but the C-termini remained unzipped. These findings indicate that one SNARE pair is not sufficient to drive fusion in this system, a finding that is also consistent with the inability of ND3 to form viable pores (Fig. 1e). In the presence of apo-syt1 (reconstituted in the v-ND), the N-N pair remained in the high FRET state, whereas the C-C pair resulted in three different FRET states: 0.2, 0.5, and 0.7 (Fig. 4). These results demonstrate that apo-syt1 drives further assembly of the SNARE complex. We suggest that this structure corresponds to the primed, clamped conformation of the trans-SNARE complex. Subsequent addition of Ca^{2+} to these samples had only subtle effects on the smFRET signals (Fig. 4), probably because the C-terminal end of the SNARE motif had largely assembled into a primed and clamped structure^{29,45}. However, an increase in the highest C-C FRET peak was apparent, revealing that Ca^{2+}•syt1 drives further assembly of the SNARE complex toward the membrane anchors. For completeness, representative raw time-based FRET traces, showing all three FRET states, are shown in Supplementary Fig. 9b. We note that our smFRET experiments are in agreement with previous studies, based on force measurements, that examined the assembly of trans-SNARE complexes^{46–48} (Supplementary Fig. 9a). In short, the ability of syt1 to clamp membrane fusion prior to the Ca^{2+} signal is associated with its ability to drive assembly of trans-SNARE complexes into a more zipperred, yet inhibited, state. Then addition of Ca^{2+} drives further assembly and thus fusion pore opening. These findings are consistent with cell-based studies which indicate that the C2B-domain of apo-syt1 is a potent fusion clamp^{36,49}. This clamping activity appears to be controlled by conformational changes that determine the relative disposition of the tandem C2-domains. Namely, apo-syt1 clamps fusion when the tandem C2-domains of apo-syt1 are askew^{26}. Then, upon binding Ca^{2+}, the C2-domains reorient and point in the same direction to trigger exocytosis^{26}.

In a second approach, we utilized the cytosolic domain of syb2 (cd-syb2), which competes with full-length syb2 in NDs for binding t-SNAREs in the BLM, to prevent or disrupt trans-SNARE pairing. In an earlier study, we found that, in the absence of syt1, cd-syb2 readily closes fusion pores, thus revealing that trans-SNARE complexes are not fully assembled, or committed,
at a stage in which fusion pores have opened to a relatively small conductance. Here we started with stable, open ND9l pores lacking syt1 and, consistent with our previous study, observed that 20 µM cd-syb2 efficiently closed individual pores (Fig. 5a, upper panel); Ca\textsuperscript{2+} was without effect. We then carried out the same experiments using ND9l/syt1. After the pore was opened by Ca\textsuperscript{2+} and robust currents, indicative of large pores, were detected, 20 µM cd-syb2 was added. Remarkably, pore closure was not observed, even over a 45-min recording period (Fig. 5a, lower panel). We then repeated this experiment but first added excess BAPTA to chelate Ca\textsuperscript{2+} and deactivate syt1; even then, subsequent addition of cd-syb2 was without effect (Fig. 5b and Supplementary Fig. 10a, b). Thus the action of Ca\textsuperscript{2+} is not required to maintain pore properties.

To formalize these observations, we calculated the fraction of time that individual pores were in the open state over a 45-min period. This fraction decreases in the presence of cd-syb2 when SNAREs alone were used (Fig. 5c). In contrast, after the action of Ca\textsuperscript{2+}/syt1, the fraction open was unaffected by cd-syb2 (Fig. 5c). These results further demonstrate that syt1 drives assembly of fusion pores formed by ND3/S/syt1 plus Ca\textsuperscript{2+} and deactivate syt1; even then, subsequent addition of cd-syb2 was without effect (Fig. 5b and Supplementary Fig. 10a, b). Thus the action of Ca\textsuperscript{2+}/syt1 appears to be terminal, as the ongoing presence of Ca\textsuperscript{2+} is not required to disassemble SNARE complexes in vitro and in vivo. NSF is recruited to SNARE complexes via an adaptor protein, α-SNAP. Then, upon ATP hydrolysis by NSF, SNARE complexes are disassembled. In vivo, NSF is thought to act on cis-SNARE complexes in the plasma membrane following fusion, where disassembly allows v- and t-SNAREs to be segregated into separate compartments for subsequent rounds of fusion. Whether NSF can act on trans-SNARE complexes has been the subject of debate. The ND-BLM assay makes it possible to determine directly whether NSF acts on trans-SNARE complexes in a functional manner, as disassembly would be evidenced by pore closure. As shown in Fig. 6, addition of α-SNAP/NSF-ATP to fusion pores formed by ND3/S/syt1/α-SNAP-γ-S (Fig. 6a, lower panel, 6a) or omission of ATPγ-S (Supplementary Fig. 11b) abolished the action of α-SNAP/NSF-ATP on pore closure, confirming that disassembly depended on efficient hydrolysis of ATP. Finally, the individual
in all experiments. Data were collected using three independent sets of NDs. In all smFRET experiments, Ca\(^{2+}\) based on SNARE complex assembly and disassembly stems from studies that have been reported\(^{43,44,52}\), which do not form until after priming but coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes. To study kinetic transitions in pores, we determined which of the distinct syt1-SNARE complex structures have been reported\(^{45,53,54}\) coincide with the primed but inactive state of SNARE complexes.

**Discussion**

At present, little is known concerning the structure and dynamics of trans-SNARE complexes because they have been difficult to trap in distinct functional states. Most of what is known regarding SNARE complex assembly and disassembly stems from studies based on cis complexes\(^{43,44,52}\), which do not form until after fusion. The major goal of the current study was to address the relationship between trans-SNARE complex assembly and fusion pore properties. We approached this question by determining how transitions in these structures are affected by two classes of regulatory proteins: one—syt1—is thought to trigger pore opening, while the other—α-SNAP/NSF—serves to disassemble SNARE complexes. To study kinetic transitions in pores, we applied a newly described ND-BLM method, which affords microsecond time resolution. This approach was combined with smFRET measurements to monitor SNARE zippering.

The first goal was to address the impact of syt1 on recombinant fusion pores. Under resting conditions, it has been proposed that apo-syt1 serves as a fusion clamp that prevents SV exocytosis until the arrival of an action potential and a concomitant increase in [Ca\(^{2+}\)]\(_{cyt}\)\(^{26,28,29,49}\). We observed that a clamping function for syt1 emerged in the ND-BLM system and was most prominent when fusion pores were formed using physiological densities of syb2\(^{30}\) (Figs. 1 and 2). smFRET measurements indicate that the ability of syt1 to clamp fusion, under resting conditions, was mediated by a direct action on trans-SNARE complexes (Fig. 4). In the absence of Ca\(^{2+}\), spontaneous openings still occurred to some degree, and kinetic analysis revealed that these openings followed multi-exponential kinetics (Fig. 2c), indicating the involvement of multiple intermediates. In contrast, in the presence of both Ca\(^{2+}\) and syt1, pore opening followed single exponential kinetics, suggesting that all trans-SNARE complexes were driven into the same primed state, making the pore opening the result of a single collective stroke. It will be interesting to determine which of the distinct syt1-SNARE complex structures that have been reported\(^{45,53,54}\) coincide with the primed but clamped state, and with the fusogenic state, of this protein complex.

Syt1 is unlikely to clamp fusion by preventing the docking of NDs to the BLM via steric effects, as this protein has been reported to facilitate docking in reconstituted systems and in synapses\(^{36,55}\). Moreover, we used physiologically relevant syb2 and syt1 densities that approximate the densities found on native SVs\(^{56}\). Indeed, SVs contain a myriad of additional proteins, so are more crowded than our NDs. Finally, the observation that syt1 inhibits pore activity, at least in part, by reducing the open life time in a dose-dependent manner (Supplementary Fig. 2) indicates that aspects of the inhibitory/clamping activity of syt1 are mediated by the intrinsic properties of the protein.

Upon addition of Ca\(^{2+}\), syt1 efficiently triggered fusion pore opening in the ND-BLM system, under all conditions that were tested. So we conducted experiments to probe for changes in the
structure of the underlying SNARE complexes. We previously observed, using SNAREs alone, that addition of cd-syb2 resulted in the disassembly of \textit{trans}-SNARE complexes and pore closure\textsuperscript{16}. These findings prompt the question of how far pores must progress such that SNAREs assemble into highly stable state that requires the action of NSF in order to be disassembled. Indeed, Ca\textsuperscript{2+}•syt1 not only opened pores but also drove pore dilation (Supplementary Fig. 12), and this was associated with further assembly of \textit{trans}-SNARE pairs as revealed by smFRET. Remarkably, once pores were opened by Ca\textsuperscript{2+}•syt1, they became resistant not only to a Ca\textsuperscript{2+} chelator but also to high concentrations of cd-syb2. Apparently, \textit{trans}-SNARE complexes had entered into a committed state such that cd-syb2 could no longer displace membrane-anchored syb2, within SNARE complexes, even over relatively long time frames (45 min). Committed fusion pores still flickered, and this might reflect their partially lipidic structure\textsuperscript{7,56,57}. We also note that the interaction of syt1 with PIP\textsubscript{2}\textsuperscript{36,38}, a plasma membrane lipid that plays a key role in exocytosis, served to stabilize the Ca\textsuperscript{2+}-triggered opening of fusion pores, so lipids also impact—directly or indirectly—fusion pore transitions.

Given that fusion pores triggered to open by the action of Ca\textsuperscript{2+}•syt1 were functionally committed, we asked whether \textit{cis}-SNARE complexes after fusion, whether these factors can disassemble \textit{trans}-complexes remains unresolved\textsuperscript{13,14}. The ND-BLM system provides a functional read-out that relates SNARE complex disassembly with fusion pore transitions in real time. Indeed, upon hydrolysis of ATP, NSF and α-SNAP did in fact close fusion pores formed by \textit{trans}-SNARE complexes. This experiment further demonstrates that SNARE zipper and unzipping reactions underlie structural and kinetic transitions in individual fusion pores. Moreover, these findings support the conclusion that we are studying \textit{trans}, and not \textit{cis}, complexes, as disassembly of \textit{cis} complexes would not be expected to close pores. It is possible that the TMD of syb2 binds to the MSP, preventing zipper opening with the TMD of syntaxin\textsuperscript{13}, but we did not observe binding in pull-down assays (Supplementary Fig. 13). However, since these experiments were performed in the presence of detergent, it remains formally possible that interactions might occur within lipid-filled NDs. It is also possible that the curvature of the membranes, or the proteinaceous aspects of the pore, prevent the v- and t-SNARE TMDs from coalescing.

In all trials, NSF-mediated closure involved flickering behavior and the appearance of a subconductance state. This state was most commonly observed during pore transitions (Fig. 6a and Supplementary Fig. 11a). Because the partially open state was associated with both open-to-closed, as well as closed-to-open, transitions, it is not simply a product of vectorial disassembly. In this light, we note that cd-syb2 can give rise to subconductance states when used to close pores formed by SNAREs alone (Supplementary in ref. \textsuperscript{16}); this occurs via the formation of partially assembled.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5.png}
\caption{Ca\textsuperscript{2+}•syt1 drives \textit{trans}-SNARE complexes into a functionally committed state. \textbf{a} Representative recording of ND9\textsubscript{L} lacking or containing syt1. Once pores were open, 20 µM cd-syb2 was added to each reaction. Closed (C) and open (O) states are indicated; the current/time scale for all traces is shown in the inset, left. Right panel: respective current histograms are shown. \textbf{b} Representative recordings of ND9\textsubscript{L} bearing syt1 in 500 µM [Ca\textsuperscript{2+}]\textsubscript{free} are shown, followed by the addition of 1.5 mM BAPTA (arrow), further followed by 20 µM cd-syb2 (arrow). The current/time scale is shown on the bottom left. \textbf{c} Fraction of time individual pores were open over a 45-min period, plotted for each condition as indicated. Error bars indicate SEM from 12 and 8 independent BLMs for \textbf{a} and \textbf{b}, respectively; 3 independent sets of NDS were used.}
\end{figure}
SNARE complex intermediates. It is likely that α-SNAP/NSF acts in a similar manner, by allowing the forward assembly of partially assembled trans-SNARE complexes, during ongoing disassembly, to yield a subconduction state. According to this view, there is a tug-of-war in which disassembly and assembly oppose one another, even during the ongoing action of NSF. One model to explain this tug-of-war is that the fusion pore is formed by two SNARE complexes. The action of NSF on one SNARE pair would close the pore, but the other SNARE complex would hold the ND in place so that the partially or completely disassembled pair can reassemble and thus re-open the pore. In this model, complete closure would occur when both SNARE pairs are disassembled at the same time. Regardless of the detailed underlying model, these time-resolved measurements indicate that functional disassembly of a pore is a complex process. Moreover, our findings raise the possibility that this disassembly reaction might underlie aspects of kiss-and-run exocytosis8,15. Interestingly, binding of SM (Sec1-Munc18) proteins in synapses38 or the binding of HOPS (Homotypic fusion and protein sorting) complex in case of vacuolar fusion29 might protect trans-SNARE complexes from NSF/α-SNAP-mediated disassembly60. Future studies will reveal whether these factors affect NSF-mediated fusion pore closure in the ND-BLM system described here.

In summary, we report key kinetic intermediates that are formed during the regulated assembly and disassembly of fusion pores. During assembly, the major Ca2+ sensor for SV exocytosis, syt1, directly regulates the occurrence, size, and dynamics of fusion pores by promoting SNARE zipper formation into a functionally committed state. This committed state could only be disassembled upon ATP hydrolysis by NSF, resulting in pore closure. Thus the defined, reduced, reconstituted ND-BLM system makes it possible to measure, in real time, the entire SNARE cycle, while revealing the behavior of fusion pores as they are constructed and deconstructed.

Methods
Materials. 1-Palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (PC), 1,2-dioleoyl-sn-glycerol-3-phospho-1-mer (PS), 1,2-dioleoyl-sn-glycerol-3-phosphoethanolamine (PE), 1,2-dipalmitoyl-sn-glycerol-3-phosphocholine (DPhPC), 1-palmitoyl-2-oleoyl-sn-glycerol-3-phospho-(1′-rac-glycerol) (sodium salt) (PG), and brain PI (4,5)P2 were purchased from Avanti polar lipids; DDM (dodecyl α-N,N,N,N-tetradecylmaltoside) and OG (n-octyl glucoside) were from Gold Biotechnology; IPTG was from Promega; RNase (Sigma, 10 µg/ml) were then added and samples were sonicated in 35 ml batches on ice for 2× 45 s (50% duty cycle). Triton X-100 was added to 19,000 g for 30 min in a 1A-17 rotor (Beckman). The supernatant was then incubated for >2 h at 4 °C with Ni-NTA agarose (Qiagen; 0.5 ml of a 50% slurry per liter of cell culture) equilibrated in resuspension buffer. Beads were washed and incubated overnight with rotation at 4 °C before centrifugation of the cell lysate and incubation for >2 h at 4 °C with Ni-NTA agarose (Qiagen; 0.5 ml of a 50% slurry per liter of cell culture) equilibrated in resuspension buffer. Beads were washed extensively with resuspension buffer containing 1% Triton X-100 and then washed with OG wash buffer (25 mM HEPES-KOH [pH 7.4], 400 mM KCl, 10 mM imidazole, and 5 mM β-mercaptoethanol) and incubated for 20 min on ice after addition of 0.5 mg/ml lysosome protease inhibitor (1 mM PMSF), DNase I, and RNase (Sigma, 10 µg/ml) were then added and samples were sonicated in 35 ml batches on ice for 2× 45 s (50% duty cycle). Triton X-100 was added to 2.1% (v/v) and incubated overnight with rotation at 4 °C before centrifugation of the cell lysate at 19,000 g for 30 min in a 1A-17 rotor (Beckman). The supernatant was then incubated for >2 h at 4 °C with Ni-NTA agarose (Qiagen; 0.5 ml of a 50% slurry per liter of cell culture) equilibrated in resuspension buffer. Beads were washed extensively with resuspension buffer containing 1% Triton X-100 and then washed with OG wash buffer (25 mM HEPES-KOH [pH 7.4], 400 mM KCl, 50 mM imidazole, 10% glycerol, 5 mM β-mercaptoethanol, 1% octyl glucoside). The slurry was loaded onto a column, washed with 5–10 column volumes of OG wash buffer, and step-eluted with OG wash buffer containing 500 mM imidazole.

Fig. 6 α-SNAP/NSF-ATP disassemble trans-SNARE complexes formed by Ca2+ · syt1. a Representative recording of a pore formed by ND3/syt1 in 500 µM [Ca2+]free. After pore formation, 0.3 µM α-SNAP, 0.3 µM NSF, 1 mM ATP, and 5 mM MgCl2 were added (red arrow). In the lower panel, ATP was replaced with 1 mM ATP-γ-S. Two different epochs of each trace are shown; the time interval between each epoch is indicated. Closed (C), open (O), and partially open (P) states are marked; the current/time scale for all traces is provided in the inset. After addition of the disassembly factors, recordings were divided arbitrarily into five 720-s epochs; the fraction of time the pores were open during each epoch was quantified for individual traces and plotted beside each panel. n = 5 BLMs using ATP, and n = 3 BLMs using ATP-γ-S; two different sets of NDs were used for each condition. b Representative current histograms for the experiments described above are shown. c Illustration (drawn to scale) shows the α-SNAP/NSF/ATP/Mg2+-mediated disassembly of SNAREs, leading to fusion pore closure.
NSF, α-SNAP, the cytoplasmic domain of syb2, and membrane scaffold proteins (MSPIED3D and NW30) were also purified as hist-tagged proteins, as described previously11,12. In brief, a similar procedure as above was used to purify these proteins except all detergents were omitted from the wash buffers. The purified proteins were dialyzed against 25 mM HEPES-KOH (pH 7.4), 100 mM KC1, 10% glycerol, and 1 mM dithiothreitol (DTT).

**Proteoliposome reconstitution.** t-SNARE liposomes were prepared as described previously. In brief, t-SNARE heterodimers were mixed together with lipids (25% PE, 75% PG) in reconstitution buffer (25 mM HEPES, pH 7.5, 100 mM KC1, and 1 mM DTT) plus 0.02% DDM. Detergent was removed with BioBeads under gentle shaking (overnight, 4 °C). t-SNARE liposomes were then isolated by fractionation16, followed by dialysis against reconstitution buffer (overnight, 4 °C).

**ND reconstitution.** Reconstitution of syb2 into NDs was performed as described16,62. In some experiments, full-length recombinant syt136 was co-reconstituted with syb2 at a 1:1 molar ratio unless otherwise indicated. MSPIED3D1 was used to generate small, 13 nm, NDs (ND2). The ratios of MSP to lipid molecules were 2:120 for ND3 and 2:1000 for ND2; the MSP-to-syb2 ratios were 2:1 (ND3) and 2:8 (ND9). The copy number of syb2 and syt1 per ND refers to the total number of syb2 and syt1 molecules, not the number of copies per face of the ND. The lipid composition was 40% PS, 35% PE, 75% PG, and 15% PE. Briefly, reconstitution involved mixing syb2, MSP, and lipids, with or without syt1 in reconstitution buffer containing 0.02% DDM. Detergent was slowly removed with BioBeads (1/3 volume, BIORAD) with gentle shaking (overnight, 4 °C). The preparation was centrifuged (20 min at 100,000 × g) to remove aggregates, and the NDs in the supernatant were purified by gel filtration using a Superdex 200 10/300 GL column, equilibrated in reconstitution buffer plus 5% glycerol.

**Planar lipid bilayer electrophysiology.** Planar lipid bilayer recordings were performed using a Planar Lipid Bilayer Workstation from Warner Instruments (USA) as described16,63,64. Briefly, lipids (30% DOPE, 52% DPhPC, 16% DOPS and 2% PE, 75% PG) in reconstitution buffer (25 mM HEPES, pH 7.5, 100 mM KCl, and 1 mM DTT) were added to yield the indicated [Ca2+] conditions. Humsilencer) acquisition system (Molecular Devices Corp.). Single-channel recordings were acquired at 10 kHz using the pCLAMP 10 (Molecular Devices, LLC) software and were filtered at 1 kHz for display purposes. All recordings were conducted at room temperature. Experiments were initiated with 1 mM BAPTA in the cis chamber, followed by the sequential addition of reconstituted liposomes bearing t-SNAREs and then NDs. This was followed by the addition of 1.5 mM CaCl2 in the cis-chamber to yield 500 μM [Ca2+]in. To generate dose–response curves, different concentrations of Ca2+ were added to yield the indicated [Ca2+] in Fig. 2d and Supplementary Fig. 7a. Pore formation and dynamics were studied at Δψ = −60 mV.

**Statistical analysis.** The number of independent trials is provided in the figure legends, along with the statistical tests that were performed. Error bars represent SEM.

**Data availability.** Data supporting the findings of this manuscript are available from the corresponding author upon reasonable request. A reporting summary for this article is available as a Supplementary Information file. The source data underlying Figs. 1e, 2a, 2c, e, 3c, and 5c and Supplementary Figs. 3a, b, 7b, and 11a are provided as a Source Data file.

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Author contributions
D.D., H.B. and E.R.C. conceived of the project and designed the experiments. D.D., H.B., K.C.C. and L.W. performed the experiments. E.R.C. supervised the projects. D.D., H.B. and E.R.C. wrote the manuscript.

Competing interests
The authors declare no competing interests.

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Correspondence and requests for materials should be addressed to E.R.C.

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