A Transient N-terminal Interaction of SNAP-25 and Syntaxin Nucleates SNARE Assembly*

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The SNARE proteins syntaxin, SNAP-25, and synaptobrevin play a central role during Ca\(^{2+}\)−dependent exocytosis at the nerve terminal. Whereas syntaxin and SNAP-25 are located in the plasma membrane, synaptobrevin resides in the membrane of synaptic vesicles. It is thought that gradual assembly of these proteins into a membrane-bridging ternary SNARE complex ultimately leads to membrane fusion. According to this model, syntaxin and SNAP-25 constitute an acceptor complex for synaptobrevin. In vitro, however, syntaxin and SNAP-25 form a stable complex that contains two syntaxin molecules, one of which is occupying and possibly obstructing the binding site of synaptobrevin. To elucidate the assembly pathway of the synaptic SNAREs, we have now applied a combination of fluorescence and CD spectroscopy. We found that SNARE assembly begins with the slow and rate-limiting interaction of syntaxin and SNAP-25. Their interaction was prevented by N-terminal but not by C-terminal truncations, suggesting that for productive assembly all three participating helices must come together simultaneously. This suggests a complicated nucleation process that might be the reason for the observed slow assembly rate. N-terminal truncations of SNAP-25 and syntaxin also prevented the formation of the ternary complex, whereas neither N- nor C-terminal shortened synaptobrevin helices lost their ability to interact. This suggests that binding of synaptobrevin occurs after the establishment of the syntaxin–SNAP-25 interaction. Moreover, binding of synaptobrevin was inhibited by an excess of syntaxin, suggesting that a 1:1 interaction of syntaxin and SNAP-25 serves as the on-pathway SNARE assembly intermediate.

Following a Ca\(^{2+}\) influx into the synaptic terminal, neurotransmitter is rapidly released from synaptic vesicles that fuse with the plasma membrane. The synaptic vesicle protein synaptobrevin 2 (also referred to as vesicle-associated membrane protein 2, or VAMP 2) and the plasma membrane proteins syntaxin 1a and SNAP-25\(^{1}\) play a central role during this process. They belong to the family of so-called soluble N-ethylmaleimide-sensitive factor attachment protein receptor (SNARE) proteins, which are essential for all vesicular trafficking steps. It is believed that SNAREs generally initiate the process of membrane fusion by the sequential formation of a stable heteromeric “trans” complex between the vesicular and the target membranes. This basic SNARE machinery is thought to be tightly controlled by a complex protein network (for a review, see Refs. 1–5).

Syntaxin and synaptobrevin each contain a single SNARE complex-forming coiled-coil region, termed the SNARE motif, directly adjacent to their C-terminal transmembrane domain. Syntaxin 1a contains an additional regulatory N-terminal region, called the Habc domain. SNAP-25 is composed of two SNARE motifs connected by a long linker and is attached to the membrane by palmitoyl modifications in this region. Upon SNARE complex formation, major conformational changes occur, with mostly unstructured proteins forming a stable α-helical complex (6–10). The mechanical force of assembly could potentially be used to overcome the repulsive forces between membranes (7). The assembled core complex consists of an elongated four-helix bundle with synaptobrevin and syntaxin each contributing one helix and SNAP-25 contributing two helices (11). In the interior, 16 layers of mostly hydrophobic amino acids are formed; however, an unusual hydrophilic layer of three glutamine residues and one arginine residue is located in the center of the bundle. Each helix can be assigned to a different SNARE subfamily. Accordingly, the SNARE helices contributing a glutamine to the zero-layer are named Qa- (syntaxin), Qc- (first half of SNAP-25; SN1), and Qd-SNARE (second half of SNAP-25; SN2), whereas synaptobrevin is the R-SNARE (12, 13). A very similar structure was found in the late endosomal SNARE complex, which is only distantly related to the synaptic complex (14). In both complexes, the SNARE helices are aligned in parallel, suggesting that assembly between vesicle and target membrane originates at the membrane-distal N-terminal ends of the respective SNARE motifs and proceeds in a zipper-like fashion toward the membrane anchors (15, 16). That SNARE proteins indeed constitute a minimal membrane fusion machinery was inferred from experiments in which SNAREs were shown to slowly catalyze fusion between liposomes (17).

Since the two Q-SNAREs syntaxin and SNAP-25 mainly reside in the plasma membrane, they are presumed to cooperate in providing the binding site for the vesicular R-SNARE synaptobrevin. This concept was substantiated by the fact that both Q-SNAREs form a stable complex in vitro. A closer inspection of the syntaxin–SNAP-25 complex, however, revealed that it already consists of a similar four-helix bundle, in which the

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The abbreviations used are: SNAP-25, soluble N-ethylmaleimide-sensitive factor attachment protein of 25 kDa; SNARE, N-ethylmaleimide-sensitive factor attachment protein receptor; FRET, fluorescence resonance energy transfer; OG, Oregon Green; TR, Texas Red; IAANS, anilino+naphthalenesulfonate iodacetamide; PBS, phosphate-buffered saline; Syb, synaptobrevin 2; BONT/A and BONT/E, botulinum neurotoxin A and E, respectively; Syx, syntaxin.
binding site for synaptobrevin is taken by a second syntaxin molecule (7, 10, 18, 19). Although one cannot exclude the possibility that the second syntaxin is necessary to stabilize the Q-SNARE interaction, it seems likely that a 1:1 syntaxin-SNAP-25 interaction would provide a more straightforward synaptobrevin binding site in vivo. The latter scenario is corroborated by studies on a homologous Q-SNARE complex involved in exocytosis in yeast, for which a 1:1 interaction of Snop (syntaxin homologue) and Sec9p (SNAP-25 homologue) is believed to provide the binding site for Snecp (synaptobrevin homologue) (20–22).

The core region of the yeast Q-SNARE complex assembles with a relatively slow rate of ~6000 m⁻¹ s⁻¹ (21). This rate of Q-SNARE assembly was found to be rate-limiting for the assembly of the ternary complex. Assembly of the synaptic SNAREs is similarly slow. Furthermore, the two synaptic Q-SNAREs syntaxin and SNAP-25 have to interact to allow for subsequent synaptobrevin binding (23). These slow SNAP assembly rates probably allow for a tight control of complex formation in vivo, but to catalyze fast membrane fusion the reaction obviously needs accelerating factors.

Strikingly, there are no obvious obstacles such as proline residues or disulfide bonds in the relatively simple four-helix bundle structure that could account for the slow assembly rates. Nevertheless, experiments at elevated temperatures revealed that assembly could not take place in the harsh conditions in which the complex falls apart (23). It seems possible that this marked hysteresis is caused by a rather delicate nucleation reaction, which might render SNAP assembly slow. But what is the structural reason for the delicacy of SNAP nucleation? An answer to this question requires deeper insights into the molecular mechanisms of synaptic SNAP assembly. Therefore, we now have designed fluorescence assays to follow the assembly steps of the core four-helix bundle of the synaptic SNAP complex. Our results suggest that a 1:1 interaction of the Q-SNARE proteins syntaxin and SNAP-25 precedes binding of synaptobrevin. The assembly of the SNAPs is rather complex, since it requires the precise N-terminal nucleation of three helices.

MATERIALS AND METHODS

Protein Constructs—The basic SNAP expression constructs, cysteine-free SNAP-25A (residues 1–206) (27), the first helix of SNAP-25A (SN1; residues 1–83), the second helix of SNAP-25A (SN2; residues 120–206), the syntaxin 1A SNAP-motif (SyxH3; residues 180–262), and synaptobrevin 2 (Syb; residues 1–96) (8) have been described before. Shortened versions of these constructs were cloned into the pET28a vector. The following shortened constructs were generated: SNAP-25A, BoNTAa fragment (residues 1–197), SN2/5C (residues 1–180), ΔNSSN25 (residues 39–206), SN1short (residues 7–83), ΔNSSN1 (residues 39–83), SN1C (residues 7–66), SN2short (residues 141–204), ΔNSSN2 (residues 159–206), and SN1C (residues 141–188); SyxH3, Δ NSSyxH3 (residues 212–262) and SyxH3A (residues 183–240); SyxSb 1–81 (residues 1–81), SyxH3C (residues 1–70), and Δ NSSybx (residues 42–96).

In addition, we utilized a variety of single cysteine variants of all three SNAP proteins that had been previously characterized by EPR spectroscopy (10). The following single cysteine variants of SyxH3 (residues 183–262) were used: Cys^225, Cys^226, and Cys^228. Synaptobrevin 2 (residues 1–96) Cys^99 and full-length SNAP-25A Cys^142 were also used.

Protein Purification—The different synaptic SNAP protein fragments were isolated from E. coli and purified by Ni²⁺-nitrophenyl-acetamide affinity chromatography followed by ion exchange chromatography on an Akta system (Amersham Biosciences) essentially as described (8, 27). The purity was greater than 95% as determined by SDS-PAGE. All ternary SNAP complexes were assembled overnight from purified individual proteins and purified using a Mono Q-column. Protein concentration was determined by absorption at 280 nm in the presence of guanidine HCl or for the individual SNAP-25 helices and the C-terminally truncated synaptobrevin constructs, which do not possess aromatic residues, using the Bradford assay.

Labeling—The sulfhydryl-reactive fluorescent probes anilinonaphthalene sulfonate (SNAS) (IAANS) Oregon Green 488iodoacetamide and Texas Red C5 bromoacetamide were purchased from Molecular Probes, Inc. (Eugene, OR). Before labeling, SNAP proteins (1–4 mg/ml) containing a single cysteine were dialyzed against degassed PBS buffer (20 mM sodium phosphate, pH 7.4, 100 mM NaCl). About a 10–20-fold molar excess of the respective fluorescence probe in N,N-dimethylformamide was added and incubated for 2 h. The reaction was stopped by adding 10 mM dithiothreitol. After 1 h, excess dye was removed by gel filtration on a Sephadex G-50 column. Afterwards, the labeled protein was dialyzed against PBS buffer. The concentration of the labeled protein was determined from its absorption at 327 nm (IAANS), 488 nm (Oregon Green), or 595 nm (Texas Red) using the molar extinction coefficients of the dyes. The protein concentration was determined by the Bradford assay.

Fluorescence Spectroscopy—All measurements were carried out in a Fluoromax 2 spectrometer equipped with autopolariizers (Jobin Yvon) in PBS buffer at 25 °C in 1-cm quartz cuvettes.

For measurements of IAANS-labeled molecules, the excitation wavelength was 397 nm and the emission spectrum was measured from 370 to 600 nm. The slit width of the excitation light was set at 2 nm, and that of the emission was set from 2.5 to 4 nm. Initially, the spectrum of the labeled molecule was measured and compared with the spectrum after the addition of unlabeled, interacting protein(s). Kinetic measurements were then performed using the emission wavelength at which the maximum change had been observed.

Changes of anisotropy upon complex formation using proteins labeled with Oregon Green were measured at an excitation wavelength of 488 nm and an emission wavelength of 520 nm. The slit widths were set to 2–4 nm, and the integration time was set at 1 s. At first, the G factor was determined according to $G = \frac{I_G}{I_{G0}}$, where $I$ is the fluorescence intensity, and the first subscript letter indicates the direction of the exciting light and second subscript letter is the direction of emitted light. Then the intensities of the vertically and horizontally polarized emission light after excitation by vertically polarized light were measured. The anisotropy (r) was calculated automatically according to $r = \frac{I_{VH} - G \times I_{VH} + G \times I_{GV}}{G \times I_{VH} - I_{GV}}$. At the beginning of the reaction, data points were recorded at the smallest interval possible (~5 s). Later, the intervals were extended accordingly.

For fluorescence resonance energy transfer (FRET) measurements, the emission spectrum was recorded from 500 to 700 nm at an excitation of 488 nm. For kinetic measurements, the change of the fluorescence intensity of the acceptor was recorded.

CD Spectroscopy—CD measurements were performed using a Jasco model J-720 instrument. For spectral measurements, different protein concentrations at a combination of about 5–10 μM in 20 mM Tris, pH 7.4, 100 mM NaCl (for details, see the appropriate figure legend) were incubated overnight. Hellma quartz cuvettes with a path length of 0.1 cm were used. The spectra were obtained by averaging over 5–50 scans using steps of 0.2 nm with a scan rate of 50 nm/min essentially as described (8, 27). The measurements were carried out at 25 °C.

Kinetic measurements were carried out in PBS buffer using 1-cm quartz cuvettes at 25 °C. Proteins were mixed, and the change in the CD signal was followed at 220 nm.

For thermal denaturation experiments, the purified ternary SNAP complexes were dialyzed against PBS buffer. The ellipticity at 220 nm was recorded between 25 and 95 °C at a temperature increment of 30 °C.

RESULTS

During SNAP assembly, major structural rearrangements occur, with mostly unstructured monomers forming complexes with high α-helical content. This allows us to monitor the assembly reaction using CD spectroscopy (6, 7, 23). This approach alone, however, does not provide the spatial information necessary for a detailed analysis of the assembly pathway, since intermediate products already exhibit similar changes in the secondary structure (8). To overcome these shortcomings, we have now introduced fluorescence probes into each of the three SNAP core complexes. Using the crystal structure (11) as a guide, several positions in each SNAP protein were chosen for fluorescence labeling. Throughout the study, only the soluble regions of the three synaptic SNAPs that form the core complex were used. An overview of the different labeling positions is given in Fig. 1.
A Second Syntaxin Competes with Synaptobrevin for Binding to a 1:1 SyxH3-SNAP-25 Intermediate—In our experiment, we mainly utilized the fluorophore IAANS, which exhibits environmentally sensitive fluorescence properties. Only a few positions of those tested exhibited a pronounced change in the fluorescence intensity that allowed us to follow SNARE assembly. As an example, SNAP-25 labeled at position 84 (SNAP-25 84IAANS) showed a strong increase in fluorescence intensity upon binding to syntaxin (Fig. 2B). At pseudo-first-order conditions, the rate of assembly was ~6000 s⁻¹ (Fig. 2D). Comparable rates were obtained when the SNARE motif of syntaxin (SyxH3) was labeled with IAANS at position 225, 240, or 248 and mixed with SNAP-25 (~7600, ~6800, and ~7400 s⁻¹, respectively) (Fig. 2C). Upon titration, saturation of fluorescence was reached at a molar ratio of about two SyxH3 and one SNAP-25 molecule, whereas in the presence of Syb, a molar ratio of 1:1 between SyxH3 and SNAP-25 was observed (data not shown). However, in all experiments in which the reaction between SyxH3 and SNAP-25 was monitored, no clear indication for a two-step assembly process was observable. A possible explanation might be that a transient 1:1 intermediate of SyxH3 and SNAP-25 creates a new binding site with increased affinity for the second SyxH3.

In the presence of Syb, the entire reaction appeared to be slower (Fig. 2, A and B) and to consist of at least two phases as we had previously observed using CD spectroscopy (23). As a different read-out for ternary complex formation, we then used Syb labeled at position 79 (Syb 79IAANS), which exhibited a clear rise in fluorescence intensity only upon the addition of both SyxH3 and SNAP-25. Interestingly, complex formation was faster when increasing amounts of SNAP-25 were added and SyxH3 was kept at a constant concentration (Fig. 3A), whereas increasing amounts of SyxH3 rather slowed down assembly (Fig. 3B). Thus, it appears that an excess of SyxH3 inhibits Syb binding, most likely because both molecules compete for the same binding site offered by a transient 1:1 SyxH3-SNAP-25 intermediate. Nevertheless, ternary complex formation was only slightly faster when SyxH3 and SNAP-25 were allowed to assemble beforehand (data not shown). This accelerating effect was most obvious when more SNAP-25 than SyxH3 was used. This probably suggests that only a small portion of premixed Q-SNAREs reside as transient 1:1 complexes, whereas the majority appears to exist as stable 2:1 complexes, which do not appear to serve as the primary binding site for synaptobrevin.

In another set of experiments, we added SNAP-25 to a mix of SyxH3 225OGK and SyxH3 225STR (i.e. syntaxins labeled at position 225 with the fluorescence dyes Oregon Green or Texas Red, respectively). Although this approach allowed only for a random integration of the differently labeled syntaxins, assembly gave rise to clear FRET, confirming that the Q-SNARE assembly contained two syntaxin molecules (Fig. 4, inset). At pseudo-first-order conditions, a Q-SNARE assembly rate of ~5900 m⁻¹ s⁻¹ was observed (data not shown), which is similar to the ones assessed with IAANS-labeled Q-SNAREs (see above). Upon the addition of an excess of Syb to the preassembled Q-SNARE complex, the FRET signal disappeared (Fig. 4), suggesting that one of the syntaxin molecules of the preformed 2:1 complex had been exchanged for Syb. The addition of an excess of unlabeled SyxH3 to the preformed Q-SNARE complex led to a comparable disappearance of the FRET signal (Fig. 4), which probably reflects the dissociation of the second syntaxin. Since Syb binding was not faster than the off-rate of the second syntaxin, it is likely that Syb does not actively replace one of the two syntaxins from the 2:1 Q-SNARE assembly.

The Syntaxin-SNAP-25 Interaction Nucleates N-terminal-ly—To assess which regions of the SNAREs are sufficient for assembly, N- or C-terminal parts of the SNARE motifs were removed (see Fig. 1). Remarkably, N-terminally truncated SNAP-25 (residues 39–206; ΔNSN25) as well as N-terminally truncated SyxH3 (residues 212–262; ΔNSyxH3) appeared to have lost their ability to undergo Q-SNARE interaction (Fig. 5, A and B). A similar result was obtained by CD spectroscopy, where no increase in α-helical content was observed for a mix of each of the constructs with the respective intact Q-SNARE binding partner (Fig. 5C). Furthermore, these constructs almost completely lost their ability to form a ternary complex containing Syb; only at high concentrations (>5 μM), a stable ternary complex slowly formed (~24 h) (Fig. 5C). This indicates that when the Q-SNARE interaction is perturbed, assembly occurs slowly due to a high energy barrier.

In contrast, a C-terminally truncated SyxH3 (residues 183–240; SyxH3ΔC) was still able to interact with SNAP-25 (Fig. 5A). Likewise, a C-terminally truncated SNAP-25 (residues 1–197), which resembles the botulinum neurotoxin A (BoNT/A) cleavage product, formed a complex with SyxH3 (data not shown). However, further shortened SNAP-25 construct (residues 1–180; SN25ΔC, which resembles the BoNT/E cleavage product) only slowly interacted with SyxH3 (Fig. 5B). Never-
theless, upon the addition of Syb, SN25ΔC readily formed a ternary SNARE complex (Fig. 5B). The molecular mass of the purified SyxH3-SN25ΔC complex determined by multiangle laser light scattering indicated a 2:2 stoichiometry (data not shown). An equivalent 2:2 complex has been described for the isolated first helix of SNAP-25 (SN1) and SyxH3 (24). Thus, in the absence of Syb, the much shortened second helix of the SN25ΔC construct may not suffice to prevent the competing but slow process of dimerization (for illustration, see Fig. 6). To test whether Syb truncations affected SNARE assembly, we utilized the increase in fluorescence anisotropy of SyxH3225OG upon complex formation. Upon assembly with SNAP-25, anisotropy of SyxH3225OG exhibited only a small increase, which might be caused by the proximity of two OG fluorophores in the SyxH3-SNAP-25 complex. A much more pronounced rise, however, was observed upon ternary SNARE complex forma-
tion. Interestingly, N- or C-terminal truncations of Syb had no significant effect on SNARE assembly (Fig. 5D). Similar results were obtained by CD spectroscopy (Fig. 5C). Taken together, these results substantiate the view that Syb binding occurs after Q-SNARE nucleation.

Although all of the above SNARE truncations removed a substantial part of the core complex-forming region, about three hydrophobic coiled-coil layers (see Fig. 1), their core complexes remained highly stable (Table I). Consequently, the pronounced effect of the truncations on assembly cannot simply be explained by the fact that the resulting core complex is less stable. Rather, the stability of the Q-SNARE intermediate is likely to be mostly affected. The SNARE complex containing ΔNSyxH3 did not refold during the time period tested, confirming our results that assembly of this fragment is extremely slow (data not shown). Remarkably, complexes containing either N- or C-terminally truncated Syb still exhibited a strong hystere-

sis (data not shown), suggesting that fragments of the Syb helix are sufficient to produce an “irreversible” SNARE interaction.

The Q-SNARE Helices Work Together for Nucleation—The SNARE protein SNAP-25 contains two SNARE motifs connected by a flexible linker region (10), which spans the entire length of the SNARE bundle in the core complex. Therefore, the question arises whether both SNAP-25 helices are necessary for successful nucleation of the Q-SNARE complex. For a detailed analysis of this process, we applied the same fluorescence approaches using the two helices of SNAP-25 as independent fragments. Neither of the two individual SNAP-25 helices produced a significant increase in fluorescence intensity upon mixing with SyxH3 and the indicated amounts of SNAP-25 (A) or upon mixing with 1.2 μM SNAP-25 and the indicated amounts of SyxH3 (B).

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played a structural change upon mixing with syntaxin (data not shown). Similarly, both shortened syntaxin variants (SyxH3\textsubscript{C} and \DeltaNSyxH3) did not exhibit a structural change upon mixing with SN1 (data not shown). Whereas we had expected, in the light of our above mentioned results, that the N-terminally truncated fragments would not be able to assemble, we had not anticipated the additional inability of the C-terminally truncated helix to assemble. One possible explana-

**Fig. 5.** N-terminal truncations in the coiled-coil region of the Q-SNAREs prevent assembly. A, SNAP-25\textsuperscript{\textsubscript{240ANNS}} (~200 nm) was mixed with ~2 \textgreek{m}N- or C-terminally truncated syntaxin SNARE motifs (\DeltaNSyxH3, residues 212–262; SyxH3\textsubscript{C}, residues 183–240, respectively). Note that the overall increase in fluorescence intensity upon binding of SyxH3\textsubscript{C} (residues 183–240) was smaller than with the intact SyxH3 (residues 180–262; see also Fig. 2B), probably because fewer conformational changes extended into the C-terminal region of SNAP-25, which carried the fluorophore. B, the fluorescence change of SyxH3\textsuperscript{240ANNS} (~200 nm) was monitored upon mixing with ~1.2 \textgreek{m} full-length SNAP-25 (\textbullet{}), N-terminally (\DeltaNSN25; residues 39–206) (\textbullet{}), or C-terminally truncated SNAP-25 (SN25\textsubscript{C}, residues 1–180). C, anisotropy of SyxH3\textsuperscript{225OG} (~100 nm) was monitored after mixing with SNAP-25 alone or in the presence of Syb or N-terminally (\DeltaNSyb; residues 42–96) or C-terminally (Syb\textsubscript{C}; residues 1–70) truncated synaptobrevin variants. D, changes in the CD signal at 220 nm of the different truncated SNARE proteins. Black columns denote the theoretical noninteracting signal (i.e. the sum of the signal of the unmixed components), whereas white columns represent the observed signal after overnight mixing of equimolar concentrations of the indicated protein combinations. The first six combinations were mixed at about 6 \textgreek{m} and all others at about 9.3 \textgreek{m} concentrations of the individual proteins. \DeltaN and \DeltaC indicate the above mentioned N- and C-terminally truncated SNAREs, respectively.
N-terminal Nucleation of SNARE Assembly

Fig. 6. Schematic representation of the assembly pathway of a C-terminal truncated SNAP-25. A, the SNARE motif of syntaxin (SyxH3) and the first helix of SNAP-25 (SN1) slowly form a stable assembly. This assembly exhibits a 2:2 stoichiometry and consists of an extended parallel four-helix bundle similar to that of the core SNARE complex (24). The SyxH3–SN1 complex probably represents an off-pathway SNARE assembly, which forms via slow dimerization of a SyxH3–SN1 intermediate. B, C-terminally truncated SNAP-25 (SN25C) and SyxH3 assemble into a complex resembling the SyxH3–SN1 complex. Probably, the truncated SN2 helix of the SN25C construct does not prevent dimerization but is only loosely connected to the four-helix bundle. In the presence of Syb, however, SN25C supports the formation of the ternary ternary SNARE complex at the rate of full-length SNAP-25.

Table I
Melting temperatures of core SNARE complexes containing truncated fragments measured by CD spectroscopy

<table>
<thead>
<tr>
<th>Complexes</th>
<th>Temperatures</th>
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<tr>
<td>Sx 183–240</td>
<td>78 °C</td>
</tr>
<tr>
<td>Sx 212–262</td>
<td>72 °C</td>
</tr>
<tr>
<td>Sb 1–70</td>
<td>79 °C</td>
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<tr>
<td>Sb 42–96</td>
<td>72 °C</td>
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<tr>
<td>SN25C</td>
<td>65 °C</td>
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<tr>
<td>Core synaptic SNARE complex</td>
<td>82 °C (23)</td>
</tr>
<tr>
<td>SyxH3–SNAP-25 complex</td>
<td>44.5 °C at 7 μM (23)</td>
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tion for this finding is that the C-terminal regions of SN1 and of SyxH3 are necessary for dimerization into the α-helical SyxH3–SN1 complex (Fig. 5). Nevertheless, the addition of SN2 to a mixture of SN1AC and SyxH3 led to a slow induction of α-helical structure (i.e., Q-SNARE complex formation) (data not shown), whereas no such induction was found with the ΔNSN1 fragment. A structural change was also observed when both individual SNAP-25 helices were mixed with SyxH3AC but not with ΔNSyxH3 (data not shown). Similarly, the addition of the SN1AC fragment but not the ΔNSN1 fragment to a mix of SyxH3AC25 and SN2 showed a clear rise in fluorescence intensity (Fig. 7B). Taken together, this largely confirms that the N-terminal regions of the SyxH3 and SN1 are necessary for Q-SNARE assembly.

Mixing the N-terminally shortened second SNAP-25 helix (residues 159–206; ΔNSN2) with SN1 and SyxH3AC25ΔNSN2 did not produce a fluorescence change (Fig. 7B), suggesting that the N-terminal portion of all three Q-SNARE helices is required for assembly. Unexpectedly, however, the C-terminally shortened second SNAP-25 helix (residues 141–188; SN2ΔC) also appeared to be incapable of supporting the formation of a Q-SNARE complex (Fig. 7B). Yet, considering our observations made with the BoNT/E-fragment of SNAP-25 (SN25AC; Fig. 4B), it seemed likely that SN2ΔC could, in principle, nucleate Q-SNARE assembly but, like SN25AC, could not prevent the competing dimerization process into the SyxH3–SN1 complex (Fig. 6). To test whether SN2ΔC might be able to support assembly in the presence of Syb, we monitored the anisotropy of SyxH3AC25, since it primarily denotes the formation of the Syb-containing SNARE complex. The addition of both individual helices of SNAP-25 to SyxH3AC25ΔN did not produce a significant increase in anisotropy, whereas the addition of Syb produced a clear increase in anisotropy (Fig. 7C). C-terminal truncations of either one of the two SNAP-25 helices exhibited a similar increase in anisotropy, indicating that they indeed allowed SNARE assembly. In contrast, N-terminal truncations of either one of the two SNAP-25 helices abolished SNARE complex formation (Fig. 7C).

Altogether, our results suggest that assembly of SyxH3 and SNAP-25 requires the orchestrated coming together of the N-terminal portions of all three Q-SNARE helices. After successful nucleation, the free binding site is readily occupied, by a second SyxH3 or Syb molecule.

DISCUSSION

The SNARE proteins syntaxin 1, SNAP-25, and synaptobrevin 2 are widely believed to constitute the basic catalytic machinery of the complex neurotransmitter release apparatus. It has been suggested that they gradually assemble into a tight complex between the synaptic vesicle and plasma membrane. Final zipper of this complex would lead to the fusion of the two membranes (15, 16).

Using CD spectroscopy, we have previously shown that the interaction of syntaxin and SNAP-25 serves as an essential intermediate for the assembly of the synaptic SNARE complex (23). This approach, however, did not allow us to clarify this intermediate consists of a four-helix bundle syntaxin-SNAP-25, complex or an entity containing only one copy of syntaxin. To achieve a more detailed picture of the assembly steps and the involved intermediates, we have now used fluorescence spectroscopy.

We found that assembly of the SNARE motif of syntaxin (SyxH3) and SNAP-25 occurred with the relatively slow rate of ~7000 M−1 s−1. After assembly, a majority of the proteins resided in a complex with 2:1 stoichiometry. This SyxH3–SNAP-25 complex has been shown to resemble the four-helix bundle structure of the ternary SNARE complex, but with a second syntaxin molecule occupying the binding site of synaptobrevin (7, 10, 18, 19). Since we did not observe two distinct steps during the assembly of syntaxin and SNAP-25, it seems possible that the fluorescence techniques used were still not suited to detecting both binding events. Another, more likely explanation is that the first step, nucleation of a 1:1 Syx-SNAP-25 complex, is rate-limiting, concealing the faster binding of a second syntaxin molecule. Furthermore, syntaxin appeared to efficiently compete with synaptobrevin, suggesting that both molecules bind onto the same transient 1:1 intermediate. Consequently, the kinetics of ternary complex assembly were rather complex and did not allow us to directly assess the rate of synaptobrevin binding. Nevertheless, our data suggest that the transient 1:1 Syx-SNAP-25 interaction allows for fast binding of synaptobrevin. Further studies to test this possibility are in progress.

A similar assembly rate of ~6000 M−1 s−1 was observed for the interaction of the core regions of the exocytotic yeast Q-SNAREs Sso1p (syntaxin homologue) and Sec1p (SNAP-25 homologue) (21). The yeast Q-SNARE complex consists of a 1:1 heterodimer (20, 21), forming a three-helix bundle that is structured only in the N-terminal two-thirds of the three participating SNARE motifs (22). This heterodimer was suggested to
serve as a high affinity binding site for Snc1p (synaptobrevin homologue) (22). In line with these findings, it might be possible that the synaptic 1:1 Syx-SNAP-25 complex also consists of a three-helix bundle. In vitro, however, syntaxin has a tendency to occupy the fourth helix position, thereby obstructing the binding of synaptobrevin. However, it is unlikely that this “autoinhibitory” effect of syntaxin observed in the cuvette plays a major role in vivo. The majority of the abundant synaptic Q-SNAREs syntaxin and SNAP-25 appear to exist in distinct locations in the plasma membrane (25), suggesting that their interaction is tightly controlled. Nevertheless, the autoinhibitory effect of syntaxin on SNARE-mediated liposome fusion needs to be considered more critically, since it is likely that the Q-SNAREs in the membrane will also reside mainly as a 2:1 complex (4).

Truncation experiments showed that assembly of syntaxin and SNAP-25 requires the coordinated interaction of the Q_a (Syntaxin), Q_b (first helix of SNAP-25), and Q_c helices (second helix of SNAP-25). Nucleation into a productive “Q_{abc}” intermediate appears to be a prerequisite for subsequent synaptobrevin binding. Remarkably, only the N-terminal and not the C-terminal truncations of the Q-SNARE helices abolished nucleation. As discussed above, the structured region of the transient 1:1 Syx-SNAP-25 complex, in analogy to the Sso1p-Sec9p complex, might be restricted to its N-terminal portion. Obviously, perturbations in the N-terminal region would critically disturb this Q_{abc} intermediate and thus render the energetic barrier for nucleation essentially insurmountable. Thus, at first glance, our data strongly support the idea that assembly of the Q_{abc} intermediate starts with the contact of the N-terminal tips of the Q-SNARE motifs and proceeds in a zipper-like fashion toward the C-terminal membrane anchor region. However, it is also possible that the first contact is not restricted to the N-terminal tips but requires the entire N-terminal region of the participating Q-SNARE helices.

Interestingly, neither N- nor C-terminal truncations of synaptobrevin appeared to impair SNARE assembly. However, as discussed above, our approach did not allow us to directly analyze binding of synaptobrevin, which occurred after the formation of the Q_{abc} intermediate. Thus, in principal, binding of synaptobrevin onto the Q_{abc} intermediate could start from either direction. Interestingly, it was shown that synaptobrevin reconstituted into liposomes is inhibited from forming a SNARE complex, although its N-terminal portion appears to be free (26). This suggests a model in which binding of synaptobrevin starts with the C-terminal portion. However, it still seems more plausible that in vivo the free N terminus of synaptobrevin binds to the structured N-terminal portion of the Q-SNARE intermediate. Otherwise, to allow for a C-terminal start of assembly, the two membranes, assuming the zipper model is correct, would have to be pulled together by a so far unknown mechanism. Such a process would require energy that is actually thought to be produced by SNARE assembly.

In this study, we have focused on the soluble portions of the synaptic SNARE proteins. Thus, clearly, in the future, similar experiments need to be carried out in the presence of memb...
branes. Although the membrane fusion activity of SNAREs has been studied in some detail (17), so far knowledge of their activity and structure in membranes is rather sparse. Nevertheless, we believe that it is unlikely that the presence of the C-terminal transmembrane region will dramatically alter the properties of the distinct soluble portion (see Ref. 4 for a more extended discussion).

Given the relatively simple four-helix bundle structure, it had been previously unclear why SNARE complex formation is slow. Our findings suggest that the slow rate can be explained by a structurally complex, and possibly unique, assembly mechanism. For productive assembly, the N-terminal portions of all three Q-SNARE helices have to come together in a parallel orientation. Two of these helices are part of one protein, SNAP-25, in which they are connected by a long flexible linker. Partition of the two helices caused the reaction to be much slower. The merger of two helices into one protein might therefore have offered a kinetic advantage for the process of synaptic exocytosis. In several other SNARE complexes that exist in vivo, however, the two helices are provided by two independent SNARE proteins. For these assembly processes, establishment of a Qabc intermediate appears to be more challenging. On the other hand, the need to bring together three proteins to form an acceptor complex on one membrane may allow for a stricter control of the reaction. However, we cannot exclude the possibility that these SNAREs might utilize a somewhat different assembly pathway.

In vivo, Q-SNARE nucleation might be catalyzed by “priming” factors, which would help to arrange the Q-SNARE helices and stabilize the intermediate, rendering it available for vesicular R-SNARE binding. It is possible that the intricate assembly intermediate permits the tight control of the upcoming fusion process. It may also confer an element of Q-SNARE selectivity during the nucleation process.

Altogether, our data suggest that synaptic SNARE assembly starts with a rather complicated interaction of the N-terminal regions of all three Q-SNARE helices (Fig. 8). It is possible that only the structured N-terminal portion of this intermediate serves as high affinity binding site for a flexible synaptobrevin. Such a scenario would increase the capture radius of the proteins interacting between two still separated membranes. The relative instability of the Qabc intermediate would provide a means of withholding and tightly controlling the upcoming, nearly irreversible, interaction with the cognate R-SNARE. It is unclear whether synaptobrevin binding occurs in a single step or whether it proceeds through a stage in which only its N-terminal region is tightly bound. Such a partially assembled trans-complex might become stalled by an interaction with late acting factors, which eventually trigger complete assembly and thus neurotransmitter release upon the influx of Ca2+.

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REFERENCES