SNARE bundle and syntaxin N-peptide constitute a minimal complement for Munc18-1 activation of membrane fusion

Jingshi Shen,¹ Shailendra S. Rathore,¹ Lavan Khandan,² and James E. Rothman²

¹Department of Molecular, Cellular, and Developmental Biology, University of Colorado at Boulder, Boulder, CO 80309
²Department of Cell Biology, Yale University School of Medicine, New Haven, CT 06520

SEC1/MUNC18 (SM) proteins activate intracellular membrane fusion through binding to cognate SNAP receptor (SNARE) complexes. The synaptic target membrane SNARE syntaxin 1 contains a highly conserved $H_{abc}$ domain, which connects an N-peptide motif to the SNARE core domain and is thought to participate in the binding of Munc18-1 (the neuronal SM protein) to the SNARE complex. Unexpectedly, we found that mutation or complete removal of the $H_{abc}$ domain had no effect on Munc18-1 stimulation of fusion. The central cavity region of Munc18-1 is required to stimulate fusion but not through its binding to the syntaxin $H_{abc}$ domain. SNAP-25, another synaptic SNARE subunit, contains a flexible linker and exhibits an atypical conjoined $Q_{bc}$ configuration. We found that neither the linker nor the $Q_{bc}$ configuration is necessary for Munc18-1 promotion of fusion. As a result, Munc18-1 activates a SNARE complex with the typical configuration, in which each of the SNARE core domains is individually rooted in the membrane bilayer. Thus, the SNARE four-helix bundle and syntaxin N-peptide constitute a minimal complement for Munc18-1 activation of fusion.

Introduction

Intracellular membrane fusion is mediated by two conserved families of proteins, the SNAREs and the Sec1/Munc18 (SM) proteins (Südhof and Rothman, 2009). Fusion is initiated when the v-SNARE pairs with its cognate t-SNAREs to form a thermally stable trans-SNARE complex (SNAREpin) that brings two membranes into close apposition to fuse (Söllner et al., 1993; Sutton et al., 1998; Weber et al., 1998; Hu et al., 2003; Wickner and Schekman, 2008). One universal feature of the trans-SNARE complex is a four-helix coiled-coil bundle assembled from the core motifs of individual t- and v-SNAREs. N to C zipper of this SNARE core bundle is believed to provide the driving force required to merge two membrane bilayers (Melia et al., 2002; Pobbati et al., 2006; Walter et al., 2010). SM proteins are soluble factors of 60–70 kD that control vesicle fusion by directly interacting with specific subsets of SNAREs (Novick and Schekman, 1979). One critical outstanding question in cell biology is how SNAREs and SM proteins act in concert to fuse membrane bilayers.
The H\textsubscript{abc} domain is critical for the inhibitory interaction of Munc18-1 with the "closed" syntaxin 1 monomer, although formation of this high-affinity syntaxin–SM dimer appears to be restricted to regulated exocytosis (Gerber et al., 2008). Whether and how Munc18-1 interacts with these regulatory sequences during fusion remain unknown.

**Results and discussion**

Munc18-1 stimulation of fusion does not require a direct interaction with the H\textsubscript{abc} domain of syntaxin 1

The role of syntaxin H\textsubscript{abc} domain in Munc18-1 stimulation of fusion is not known. The highly conserved H\textsubscript{abc} domain connects the N-peptide motif to the core motif (Misura et al., 2000; Burkhardt et al., 2008). The H\textsubscript{abc} domain is critical for the inhibitory interaction of Munc18-1 with the “closed” syntaxin 1 monomer, although formation of this high-affinity syntaxin–SM dimer appears to be restricted to regulated exocytosis (Gerber et al., 2008). Whether and how Munc18-1 interacts with these regulatory sequences during fusion remain unknown.
Munc18-1–associated SNAREs. Reconstituted t- and v-SNAREs drove a basal fusion reaction that was strongly accelerated by Munc18-1. The stimulation was abolished when the N-peptide of syntaxin 1 was removed (Fig. 1, B and C). The potent effect of Munc18-1 on SNARE-mediated membrane fusion (approximately ninefold increase in initial rate) was in line with the severe defects in neuronal release (80% to >90% reduction) observed in Munc18-1–null neurons (Verhage et al., 2000; Voets et al., 2001; Weimer et al., 2003).

To explore whether Munc18-1 stimulation of fusion involves a specific interaction with the syntaxin H_{abc} domain, we replaced the latter with the corresponding H_{abc} domain of syntaxin 7, an endosomal SNARE exhibiting limited sequence homology with syntaxin 1 (Fig. 1 A; Antonin et al., 2002). This H_{abc}7 chimeric mutant is expected to retain the overall structure of syntaxin. Unexpectedly, the SNARE complex containing this mutant syntaxin was activated by Munc18-1 at a similar level to that of wild-type (WT) SNAREs (Fig. 1, B and C). We next replaced the H_{abc} domain with the yeast small ubiquitin-like modifier (SUMO; SUMO/Smt3p) protein that has no sequence similarity to syntaxin (Fig. 1 A). SUMO contains a β-barrel structure flanked by α-helices (Bergink and Jentsch, 2009), which is markedly distinct from the three-helix conformation of the syntaxin H_{abc} domain (Fernandez et al., 1998). Again, Munc18-1 robustly activated the fusion of liposomes reconstituted with SNAREs containing this syntaxin-SUMO chimera (Fig. 1, B and C). Importantly, all of the mutants tested in this study drove comparable basal reactions, suggesting that SNARE assembly remained intact (Fig. 1 B). These data indicate that Munc18-1 does not require a specific interaction with the syntaxin H_{abc} domain to activate SNARE-dependent membrane fusion.

Munc18-1 activates a mutant SNARE complex in which the syntaxin N-peptide is directly attached to the core bundle without the H_{abc} domain.

We next considered the possibility that the binding of Munc18-1 to the SNARE complex requires only a physical hinge between syntaxin N-peptide and the core bundle, rather than engaging a specific interaction with the H_{abc} domain. The H_{abc} domain of syntaxin 7 or even an unrelated SUMO protein may serve this hinge role as adequately as the native H_{abc} domain of syntaxin 1. Indeed, all of these chimeric proteins have similar linker lengths (∼9 nm) between the N-peptide and SNARE bundle. To this end, we removed the entire H_{abc} domain from syntaxin 1 such that N-peptide became directly attached to the core motif without the H_{abc} hinge (Fig. 2 A). The SNAREs lacking the H_{abc} domain (ΔH_{abc} mutant) drove a basal fusion reaction comparable with that of WT SNAREs and, strikingly, were fully activated by Munc18-1 (Fig. 2, B and C). The maximum stimulation was reached with 5 µM Munc18-1, similar to the dose response of WT SNAREs (Fig. S1). Munc18-1 acceleration of fusion was abolished when VAMP2 was substituted with VAMP8,
could not be stimulated by Munc18-1 (Fig. 2, B and C), resulted in a full activation of membrane fusion. The central cavity region of Munc18-1 is critical for stimulating fusion but not through binding to the syntaxin H\textsubscript{abc} domain. The dispensability of the syntaxin H\textsubscript{abc} domain in Munc18-1 stimulation of fusion is a surprising finding because Munc18-1 was previously suggested to require the H\textsubscript{abc} domain for activating vesicle fusion (Rodkey et al., 2008; Deák et al., 2009). In particular, the glutamate 59 residue (E59) of Munc18-1 was found to be critical for the association of Munc18-1 with the SNARE complex (Deák et al., 2009). Mutation of E59 into lysine (E59K) strongly reduced SNARE–Munc18-1 binding affinity and caused severe defects in neuronal release (Deák et al., 2009). The E59 residue is located within the central cavity of Munc18-1 and interacts with arginine 114 (R114) of syntaxin 1 in the syntaxin–Munc18-1 binary complex (Fig. 4 B; Misura et al., 2000). Because the R114 residue is within the H\textsubscript{abc} domain of syntaxin 1 (Fig. 4 A), it was suggested that Munc18-1 E59 also interacts with the syntaxin H\textsubscript{abc} domain when activating the SNARE complex (Deák et al., 2009). Consistent with the in vivo observation, we found that the E59K mutation dramatically

We also tested for a direct interaction between Munc18-1 and SNAREs in a copurification assay. Results obtained from this system agree well with those from other binding assays such as gel filtration and nuclear magnetic resonance (Dulubova et al., 2007; Shen et al., 2007). We found that Munc18-1 was associated with stoichiometric amounts of SNARE complexes containing both WT and ΔH\textsubscript{abc} syntaxin 1 but not with a mutant complex lacking the entire N terminus of syntaxin (Fig. 3 A). The identities of the components in the complex were confirmed by Western blotting (Fig. 3 B). Together, these results demonstrate that Munc18-1 does not require the syntaxin 1 H\textsubscript{abc} domain to interact with the SNARE complex or to activate fusion. Thus, addition of a short N-peptide to the SNARE bundle, which otherwise could not be stimulated by Munc18-1 (Fig. 2, B and C), resulted in a full activation of membrane fusion.

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Munc18-1 activation of fusion does not require the linker or the Q_{bc} configuration of SNAP-25

Although the interactions of Munc18-1 with syntaxin 1 and VAMP2 have been demonstrated (Figs. 1–3; Shen et al., 2007), little is known about Munc18-1 binding to SNAP-25. SNAP-25, lacking a transmembrane domain, contains two core motifs (Q_{bc} and Q_{cd}) joined together by a flexible linker (Fig. 5A). This Q_{bc} organization is an exception to the typical configuration of the SNARE fusion machinery, in which each of the four SNARE core domains (Q_{a}, Q_{b}, Q_{c}, and R) is individually rooted to the lipid bilayer through C-terminal transmembrane domains (Jahn and Scheller, 2006).

We examined whether Munc18-1 stimulation of fusion requires the flexible linker or the conjoined Q_{bc} configuration of SNAP-25. To this end, each of the core domains of SNAP-25 (25N or 25C) was fused to a generic transmembrane domain derived from the PDGF receptor (PDGFR; Fig. 5A). These two chimeric SNAREs, 25N-PDGFR (Q_{a}) and 25C-PDGFR (Q_{b}), were individually expressed and purified without the linker region or lipid modification found on the native SNAP-25 protein. 25N-PDGFR and 25C-PDGFR proteins were mixed with syntaxin 1 to assemble the t-SNARE complex. Proteoliposomes reconstituted with this engineered t-SNARE complex fused reduced the stimulatory activity of Munc18-1 on membrane fusion (Fig. 4, C and D). However, mutation of R114 of syntaxin 1 into glutamate (R114E) had little effect on Munc18-1 activation of fusion (Fig. 4, C and D), suggesting that the E59–R114 interaction does not occur when Munc18-1 is associated with the SNARE complex. Although unexpected, this result is consistent with our discovery that the entire H_{abc} domain of syntaxin 1 is dispensable for Munc18-1 activation of fusion (Fig. 2).

Thus, the residue E59 of Munc18-1 binds to a SNARE epitope other than the H_{abc} domain when stimulating fusion. Further support of our conclusion came from recent observations that the SNARE four-helix bundle competes with the syntaxin H_{abc} domain to occupy the central cavity region of Munc18-1 such that the binding to the region by one bundle is mutually exclusive of the other (Xu et al., 2010; unpublished data). Once occupied by the SNARE bundle (Shen et al., 2007; Xu et al., 2010), the central cavity region of Munc18-1 will have no room to accommodate the three-helix H_{abc} domain. Our findings also raise the intriguing possibility that residues lining the central cavity region of Munc18-1 such as E59 may engage in a dual mode of SNARE interaction: with the SNARE core bundle in the context of SNARE complex binding or with the H_{abc} domain in the context of binding to the closed syntaxin 1 monomer.

Figure 4. The central cavity domain of Munc18-1 is critical for stimulating fusion but not through binding to the syntaxin H_{abc} domain. (A) Diagram of syntaxin 1 showing the location of arginine 114 (R114). TMD, transmembrane domain. (B) Crystal structure of syntaxin 1–Munc18-1 dimer showing the interaction of syntaxin R114 with Munc18-1 glutamate 59 (E59; Misura et al., 2000). The structure was derived from reference (derived from Misura et al. [2000]) and edited in PyMOL (DeLano Scientific LLC). (C) Fusion of t-SNARE liposomes containing WT or mutant syntaxin 1 with VAMP2 liposomes in the absence or presence of 5 µM WT or E59K mutant Munc18-1. (D) Fold activation of the fusion reactions in C. The dashed line indicates basal fusion level (with no Munc18-1 activation). Error bars indicate standard deviation.
complex represents the minimal complement required for Munc18-1 acceleration of fusion. Previous studies suggested that the syntaxin H concatenated domain can interact with Munc18-1 (Dulubova et al., 2007; Rodkey et al., 2008). However, our data clearly show that this interaction, if existent, is dispensable for Munc18-1 activation of the SNARE complex. SM proteins exhibit conserved structures and similar loss of function phenotypes (abrogation of fusion), implying that we have likely uncovered a conserved regulatory mechanism of SM proteins. This conclusion is further supported by our observation that Munc18-1 activates an engineered SNARE complex with the typical configuration.

At the center of the vesicle fusion machinery is the SNARE four-helix bundle. The bundle not only supplies the energy required for the merging of two membranes, but it is also the primary binding target of SM proteins, which are fundamentally designed to grasp four-helix bundles using their central cavity regions (Scott et al., 2004; Latham et al., 2006; Mima et al., 2008; Toonen and Verhage, 2007; Rickman and Duncan, 2010). Although individual SNARE subunits exhibit diverse conformations, the structures of assembled four-helix bundles are highly conserved across pathways (Ungar and Hughson, 2003).

Conclusion

In this study, we demonstrate that the syntaxin N-peptide motif and the SNARE four-helix bundle are sufficient for Munc18-1 activation of fusion, whereas all other SNARE sequences are dispensable (Fig. 5 D). We propose that this reduced SNARE complex represents the minimal complement required for Munc18-1 acceleration of fusion. Previous studies suggested that the syntaxin H concatenated domain can interact with Munc18-1 (Dulubova et al., 2007; Rodkey et al., 2008). However, our data clearly show that this interaction, if existent, is dispensable for Munc18-1 activation of the SNARE complex. SM proteins exhibit conserved structures and similar loss of function phenotypes (abrogation of fusion), implying that we have likely uncovered a conserved regulatory mechanism of SM proteins. This conclusion is further supported by our observation that Munc18-1 activates an engineered SNARE complex with the typical configuration.

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C-terminal transmembrane domains of SNAREs form continuous α-helices with the core motifs and thus are integral parts of the SNARE bundle (Stein et al., 2009). Because N-peptide binding is not found among certain SNARE–SM pairs, the SNARE bundle is likely the conserved target of SM proteins. Structural modeling and biochemical analysis demonstrate that the SNARE bundle competes with the closed syntaxin 1 monomer to occupy the central cavity domain of Munc18-1 such that the binding to this region by one bundle is mutually exclusive of the other (Xu et al., 2010; unpublished data). This agrees well with our discovery that the syntaxin $H_{abc}$ domain is dispensable for Munc18-1 association with the SNARE complex: once occupied by the SNARE bundle, the central cavity region of Munc18-1 will not have room to accommodate the three-helix $H_{abc}$ domain of syntaxin. Our data also imply that the same residues lining the Munc18-1 cavity region such as E59 may engage a dual mode of SNARE interaction, with the SNARE core bundle in the context of SNARE complex binding or with the $H_{abc}$ domain in the context of closed syntaxin 1 monomer binding. In contrast to the closed syntaxin monomer that displays a relatively static conformation, the SNARE bundle is highly dynamic, and Munc18-1 may recognize multiple intermediates of the assembling SNARE bundle.

It should be noted that the regulatory sequences outside the minimal SNARE complement, albeit dispensable for Munc18-1 activation, may play important roles in membrane trafficking in the cell. For instance, the flexible linker and Q$_c$ configuration of SNAP-25 may be necessary for the proper subcellular localization of the protein. In contrast, the $H_{abc}$ domain of syntaxin 1 may fine-tune synaptic transmission in higher organisms when it folds back onto syntaxin’s own core motif to present a closed conformation (Misura et al., 2000; Gerber et al., 2008). The $H_{abc}$ domain may also serve as an organizing center allowing additional regulatory factors such as Munc13, CAPS/Unc-31, synaptotagmin, and complexin to be superimposed on the SNARE–SM fusion machinery to achieve temporal and spatial control of synaptic release (Basu et al., 2005; Giraudo et al., 2006; Jackson and Chapman, 2006; Tang et al., 2006; Martens et al., 2007; Zhou et al., 2007; Yoon et al., 2008; de Wit et al., 2009; James et al., 2009).

Materials and methods

Protein expression and purification

To produce recombinant VAMP2/synaptobrevin proteins with no extra residues, a full-length mouse VAMP2 gene was subcloned into a pET28a-based SUMO/Smt3p cloning vector. The sequence between the SUMO cleavage site and the first residue of VAMP2 was removed by site-directed mutagenesis. After binding to a nickel column (GE Healthcare) through an N-terminal His$_6$ tag, VAMP2 protein was released from SUMO by overnight on-column cleavage with SUMO protease at 4°C. The iSNARE complex of mouse His$_6$–SNAP-25 and rat syntaxin 1A (p14V34) was expressed and purified as previously described (Weber et al., 2000; Melia et al., 2002). Recombinant untagged Munc18-1 protein was produced in E. coli as previously described (Shen et al., 2007). SNARE and Munc18-1 mutants were generated by site-directed mutagenesis and standard molecular cloning and purified similarly to WT proteins.

Proteoliposome reconstitution

All lipids were obtained from Avanti Polar Lipids, Inc. For iSNARE reconstitution, 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (POPE), 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoserine (POPS), and cholesterol were mixed in a molar ratio of 60:20:10:10. For v-SNARE reconstitution, POPC, POPE, POPS, cholesterol, N7-nitro-2,1,3-benzoazadiazole-4-yli-1,2-dipalmitoyl phosphatidylethanolamine (NBD-DPPE), and N-lissamine rhodamine B sulfonyl-DPPE (rhodamine-DPPE) were mixed at a molar ratio of 60:17:10:10:1.5. iSNARE proteoliposomes were prepared by detergent dilution and isolated on an Accudenz density gradient flotation (Weber et al., 1998). SNARE proteins were kept at physiologically relevant densities, with protein/lipid ratios at 1:200 for v-SNAREs (similar to VAMP2 densities reported for native synaptic vesicles; Jahn and Südhof, 1994; Walch-Solimena et al., 1995) and at 1:500 for t-SNARE liposomes. Reconstituted liposomes were routinely monitored by dynamic light scattering and electron microscopy with negative staining.

Liposome fusion assay

Fusion reactions and data analysis were performed as previously described (Shen et al., 2007). A standard fusion reaction contained 45 µl unlabeled v-SNARE liposomes and 5 µl labeled v-SNARE liposomes and was conducted in a 96-well Nunc plate at 37°C. Fusion was followed by measuring the increase in NBD fluorescence at 538 nm (excitation 460 nm) every 2 min. At the end of the 2h reaction, 10 µl of 2.5% dodecyl-maltoside was added to the liposomes. The raw NBD fluorescence data were converted to fusions using an equation as previously described (Parlati et al., 1999). To assess the regulatory activity of Munc18-1, v- and t-SNARE liposomes were incubated with or without 5 µM Munc18-1 on ice for 1 h before the temperature was elevated to 37°C to initiate fusion. The maximum fusion rate within the first 20 min of liposome fusion was used to represent the initial rate of a fusion reaction. Full accounting of statistical significance was included for each figure based on at least three independent experiments.

Protein interactions in a copurification assay

SNARE–Munc18-1 interactions were probed in a copurification assay in which bacterial cells expressing both GST–Munc18-1 and SNAREs were lysed in protein-binding buffer (25 mM Heps, pH 7.4, 150 mM KCl, 10% glycerol, 1% CHAPS, and 1 mM DTT). Glutathione Sepharose (GE Healthcare) was added to the lysate to purify GST–Munc18-1 and associated proteins. After washing three times with binding buffer, protein complexes bound to the beads were resolved on SDS-PAGE and stained with a Coomasie blue staining kit (Thermo Fisher Scientific). Identities of the proteins in the complexes were confirmed by Western blotting with polyclonal anti–Munc18-1 (Sigma-Aldrich), monoclonal anti–SNAP-25 (Cl 71.2; Synaptic Systems GmbH), or anti-VAMP2 (Cl 69.1; Synaptic Systems GmbH) antibodies.

Online supplemental material

Fig. S1 shows the dose dependence of the stimulatory effects of Munc18-1 on the fusion reactions mediated by WT or ΔH$_{abc}$ SNAREs. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201003148/DC1.

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