Sly1 protein bound to Golgi syntaxin Sed5p allows assembly and contributes to specificity of SNARE fusion complexes

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Fusion of transport vesicles with their target organelles involves specific membrane proteins, SNAREs, which form tight complexes bridging the membranes to be fused. Evidence from yeast and mammals indicates that Sec1 family proteins act as regulators of membrane fusion by binding to the target membrane SNAREs. In experiments with purified proteins, we now made the observation that the ER to Golgi core SNARE fusion complex could be assembled on syntaxin Sed5p tightly bound to the Sec1-related Sly1p. Sly1p also bound to preassembled SNARE complexes in vitro and was found to be part of a vesicular/target membrane SNARE complex immunoprecipitated from yeast cell lysates.

This is in marked contrast to the exocytic SNARE assembly in neuronal cells where high affinity binding of N-Sec1/Munc-18 to syntaxin 1A precluded core SNARE fusion complex formation. We also found that the kinetics of SNARE complex formation in vitro with either Sly1p-bound or free Sed5p was not significantly different. Importantly, several presumably nonphysiological SNARE complexes easily generated with Sed5p did not form when the syntaxin was first bound to Sly1p. This indicates for the first time that a Sec1 family member contributes to the specificity of SNARE complex assembly.

Introduction

The prelude to and the execution of membrane fusion in exo- and endocytosis require sets of stage-specific core proteins, which are highly conserved from yeast to man (for review see Bennett and Scheller, 1993; Jahn and Südhof, 1999). Among them are donor and target membrane receptors, SNAREs (Sollner et al., 1993), SNARE-binding Sec1 family proteins (Hata et al., 1993; Grabowski and Gallwitz, 1997; Carr et al., 1999; Misura et al., 2000), and small GTPases of the Ypt/Rab family, which appear to act as regulators during fusion complex assembly (for review see Lazar et al., 1997; Novick and Zerial, 1997; Schimmöller et al., 1998).

SNAREs are type II membrane proteins with the membrane-spanning domain at or close to their COOH-terminal ends. Adjacent to this, they contain a conserved coiled-coil region, also termed the SNARE motif, which during SNARE pairing adopts an α-helical fold. In a fully assembled core fusion complex, four such domains donated by three or four individual SNAREs form a rigid four-helix bundle (Poirier et al., 1998; Sutton et al., 1998; Antonin et al., 2002). Instead of subdividing the membrane receptors into vesicular SNAREs (v-SNAREs) and target membrane SNAREs (t-SNAREs) (Sollner et al., 1993), it was proposed recently to classify them as Q- and R-SNAREs, depending on whether a glutamine or an arginine, two highly conserved amino acids residues, occupy the center position of the ionic layer of the four-helix bundle. It was also proposed that core fusion complexes are generally made up of three Q-SNAREs and one R-SNARE. (Passhauer et al., 1998). Although SNARE pairing in solution is rather promiscuous (Passhauer et al., 1999; Yang et al., 1999), yeast SNAREs inserted into liposomes were shown to interact in a remarkably specific way and to lead to membrane fusion, although at low rate (McNew et al., 2000).

It is unclear presently how within living cells SNAREs are activated, how the correct trans-SNARE interactions are established, and how membrane fusion efficiency is regulated. However, it is likely that Sec1 family proteins contribute to these events, and yeast genetic data regarding vesicular transport in exo- and endocytosis argue in favor of a positive rather than a negative regulatory function of these proteins (Novick and Scheckman, 1979; Ossig et al., 1991; Grote et
The t-SNARE Sed5p but not the v-SNAREs Bet1p, Bos1p, or Sec22p bind to Sly1p in vitro

We have shown previously (Grabowski and Gallwitz, 1997) that the yeast Golgi syntaxin Sed5p binds to the Sec1 family member Sly1p with the same nanomolar affinity as the neuronal plasma membrane-localized syntaxin 1A to N- Sec1/Munc18 (Pevsner et al., 1994). N-Sec1 binding requires syntaxin 1A to be in a closed conformation, and this bimolecular complex does not allow the assembly of certain SNARE complexes with other v-SNARE combinations that are, however, generated efficiently in the absence of Sly1p.
form the core fusion complex (Yang et al., 2000). Therefore, we sought to investigate the Sly1p-SNARE binding properties in more detail with the goal to come closer to an understanding of Sly1 protein function in ER to Golgi transport.

To begin with, in vitro binding studies were performed with a glutathione S-transferase (GST)–Sly1 fusion protein and NH2-terminally His6-tagged t/v-SNAREs Sed5p, Bos1p, Sec22p, and Bet1p, each lacking the COOH-terminal membrane anchor and all being produced in *Escherichia coli* and purified by affinity chromatography (Fig. 1 A). Glutathione agarose beads with bound GST–Sly1p were incubated at 4°C for 2 h with the individual SNAREs, and after extensive washing with binding buffer proteins bound to the beads were separated by SDS-PAGE. As expected, the synaptobrevin Sed5p bound efficiently to Sly1p, whereas none of the v-SNAREs exhibited binding to the Sec1 family member in this assay (Fig. 1 B).

The brain plasma membrane syntaxin 1A requires the NH2-terminal variable region for high affinity binding to N-Sec1 (Kee et al., 1995). In contrast, Vam3p, the yeast t-SNARE essential for homotypic vacuole fusion (Nichols et al., 1997; Wada et al., 1997; Seals et al., 2000), appears to bind its cognate Sec1 family member Vps33p via the SNARE motif region (Dulubova et al., 2001). In a previous report, Sly1 protein binding was assigned to the NH2-terminal 78 amino acids of Sed5p (Kosodo et al., 1998). As in this study in which GST–Sly1 or MBP–Sly1 fusions were probed for binding with GST–Sed5 fusions, we performed an affinity study with untagged soluble Sly1p that was incubated with agarose bead-bound GST fusions of either the NH2-terminal domain or the SNARE motif region of Sed5p (Fig. 1 C). In accordance with the results of Kosodo et al. (1998), Sly1p bound efficiently only to the NH2-terminal region of Sed5p, whereas the v-SNAREs exhibited binding to the Sec1 family member in this assay (Fig. 1 B).

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### Efficient SNARE complex formation in vitro on Sly1p-bound Sed5p

Since a bimolecular complex of Sly1p and Sed5p could be easily formed on beads, we addressed the question of whether in this complex the syntaxin Sed5p was able to associate with cognate v-SNAREs, Bos1p, Sec22p, and Bet1p. Preformed GST–Sly1p–Sed5p complex was incubated at 4°C for 17–22 h with an excess of His6-Sly1p, the latter at equimolar ratio. As shown in Fig. 2 A, lane 2, extensively washed beads retained, in addition to GST–Sly1p and Sed5p, all three v-SNAREs at a stoichiometry of 1.0:0.7:0.8. Binding of the v-SNAREs to GST alone was not observed (Fig. 2 A, lane 1). To explore the significance of different v-SNAREs in the process of fusion complex formation in vitro, the GST–Sly1p–Sed5p subcomplex on agarose beads was incubated with an excess of His6-Sly1p and each v-SNARE at an equimolar ratio. As shown in Fig. 2 B, lane 2, the GST–Sly1p–Sed5p subcomplex on agarose beads was incubated with an excess of His6-Sly1p and each v-SNARE at an equimolar ratio.

The stoichiometry of Sly1p to Sed5p in the complexes formed with excess of Sly1p (lanes 5 and 6) was on average 1:0.85 as determined by densitometry. (C) GST–Sed5p (10 μg, lane 2) and the heterodimeric complex formed from GST–Sed5p (10 μg) and His6-Sly1p (50 μg) bound to beads were incubated at 4°C for 18 h with Bos1p (15 μg), Sec22p (15 μg), and Bet1p (5 μg). The complexes formed were analyzed as in A. The stoichiometry of the Sed5p–Bos1p–Sec22p–Bet1p in the complex was 1:1.3:1:3:2:1:5 (lane 2).
Sly1p binds to preassembled SNARE complex

One might argue that once a SNARE complex has formed in vitro or in vivo Sed5p would be in a conformation unable to associate with Sly1p. This in fact is the case for the neuronal syntaxin 1A, which after assembled into a SNARE complex does not bind N-Sec1 (Yang et al., 2000). Therefore, we generated a SNARE fusion complex, this time with GST–Sec22p and the His<sub>6</sub>-tagged SNARES Sed5p, Bos1p, and Bet1p. Preparation as above, were incubated with His<sub>6</sub>–Sec17p (20 μg) at 4°C for 3 h, and complexes on beads were characterized by SDS-PAGE and densitometry. The Sed5p to Sec17p stoichiometry in Sly1p-free and Sly1p-containing complexes was 1:3.3 and 1:2.7, respectively.

Sec17p binds to Sly1p-containing SNARE complexes

α-SNAP and its yeast homologue Sec17p have been shown to bind to neuronal and exocytic SNARE complexes, respectively, that were generated from recombinant proteins (Söllner et al., 1993; Rossi et al., 1997). We felt it to be of interest to investigate whether Sec17p could also associate with ER to Golgi SNARE complexes to which stoichiometric amounts of Sly1p were already bound.

Using GST–Sec22p as a tool for SNARE complex isolation on glutathione beads, complexes formed with Bos1p, Bet1p, and either free or Sly1p-bound Sed5p interacted equally efficiently with purified Sec17p. Densitometric analysis revealed that the multimeric complexes (Fig. 3 A, a complex of the four SNARE proteins could be formed in solution to which Sly1p did bind in stoichiometric amounts. Here again the Sec1 binding properties of the ER to Golgi and the neuronal exocytic SNARE complexes differ, but the former appear to resemble the yeast exocytic SNARE complex, which by immunoprecipitation experiments has been shown to associate with Sec1p in cellular extracts (Carr et al., 1999).

To further analyze the molar ratio of individual components within the SNARE complex assembled in vitro, a fixed amount of GST–Sed5p (Fig. 1 A) was incubated with increasing amounts of NH<sub>2</sub>-terminally His<sub>10</sub>-tagged Sly1p. The bimolecular complexes were purified on glutathione agarose beads. As can be seen in Fig. 1 B; even with an eight-fold excess of Sly1p over Sed5p, a complex of ~1:1 stoichiometry was formed. Assembly of the fusion complex was then performed with the cognate v-SNAREs, and the molar ratios of Sly1p, Sed5p, Bos1p, Sec22p, and Bet1p in the purified complex were determined by densitometry (Fig. 2 C, lane 3). Assuming that equal amounts of Bos1p and Sec22p (that did not clearly separate in gels) were in the complex, a stoichiometry of 1.2:1.0:1.4:1.4:1.3 was determined.

This result indicated that in sharp contrast to what has been found with mammalian syntaxin 1A bound to N-Sec1 (Yang et al., 2000) a core fusion complex can be formed in vitro on Sed5p bound to the Sec1 family member Sly1p. This might indicate that Sly1p associates with Sed5p being in an open conformation.
Figure 4. Sly1p is a component of the core SNARE fusion complex in vivo. (A) Cleared detergent lysates (1.2 mg of protein) from wild-type yeast strain MSUC1A (lane 2, untagged) and from yeast strain RPY137 expressing a protein A–Sly1 fusion protein (lane 3, ProA-SLY1) were incubated with IgG-Sepharose beads at 4°C overnight. Proteins on beads (lanes 2 and 3) were separated by SDS-PAGE along with a fraction of either the original lysate (lane 1; representing 3.2% of protein applied to beads) or the protein not bound to beads (lane 4; representing 4% of protein applied to beads) and subjected to immunoblot analysis with anti-SNARE and anti-Sly1p antibodies. Note that ProA-Sly1p and Sed5p were partially degraded. (B) A cleared detergent lysate of yeast strain MSUC1A was subjected to immunoprecipitation with affinity purified anti-Bos1p antibody. The immunoprecipitate (lane 2) along with a fraction of the original lysate (lane 1) and the non-precipitated protein (lane 3) representing 3 and 3.25% of the material used for immunoprecipitation, respectively, were subjected to immunoblot analysis with antibodies to proteins indicated to the right. Sed5p is somewhat obscured by the presence of the IgG heavy chain.

Figure 5. Sly1p does not affect stability of Sed5p in vivo or of Sed5p-containing ER to Golgi SNARE complexes generated in vitro. (A) GSF4 (GAL10-SLY1) was grown in galactose-containing medium and then transferred to glucose medium to switch off transcription of the SLY1 gene. At the indicated times, aliquots of the cultures were removed, and the same amount of cells from each sample were lysed with NaOH and subjected to immunoblot analysis with affinity purified antibodies against Sly1p, Sed5p, and Bos1p. Densitometric data analysis revealed that cells at 20 h in glucose contained ~2.4% of Sly1p compared with the level of wild-type cells, which corresponds to the 10-h time point. The Sed5p level (set at 1.0 at the 0-h time point) was calculated to be 1.0, 1.1, 1.5, 1.3, 1.2, 0.8, 1.0, 0.8, 1.0, and 1.3 for the 2–20 h time points. (B) SNARE complexes generated without (lanes 1 and 2) or with Sly1p (lanes 3 and 4) and purified on glutathione beads as described in the legend to Fig. 2 C were subjected to treatment with sample buffer containing 2% SDS at room temperature (–) or 95°C (+) for 5 min. Proteins were separated on a 16% SDS-polyacrylamide gel and visualized by Coomassie blue staining. As a control, the in vitro assembled neuronal minimal core complex (lanes 5 and 6) was treated and processed in the same way. Note the SDS resistance of this complex (lane 5).
tained an approximately threefold higher molar concentration of Sec17p compared with the individual SNAREs. Importantly, binding of Sec17p was independent of whether Sly1p was part of the preassembled SNARE complex or not.

Evidence for a Sly1p-bound SNARE complex in vivo
To complement the in vitro studies and to demonstrate that ER to Golgi SNARE complexes bound to Sly1p are also present in vivo, two different experimental approaches were followed. First, a yeast strain was constructed with the essential chromosomal SLY1 gene deleted and replaced by a vector-contained gene expressing a protein A–Sly1 fusion protein. This strain grew like wild-type yeast, showing that the Sly1 fusion protein was functionally active. A cleared cell lysate of this strain was incubated with IgG-Sepharose to capture the Sly1 fusion protein and proteins bound to it. After extensive washing of the affinity beads, bound proteins were separated by SDS-PAGE and subjected to immunoblotting with specific polyclonal antibodies. Under conditions where the protein A–Sly1 fusion protein of the cell extract was entirely bound to the beads, fractions of the total cellular SNAREs Sed5p, Bos1p, Sec22p, and Bet1p were copurified with Sly1p (Fig. 4 A). Except for Sed5p, of which a significant part copurified with the Sly1 fusion protein, only small fractions of the SNAREs were found in the immunocomplex. Quantification of band intensities with the help of a Lumimager revealed that \( \sim 74\% \) of total Sed5p but \( < 1\% \) of the cognate v-SNAREs coprecipitated with Sly1p. This would be consistent with the reasonable assumption that in growing cells only limited fractions of the SNAREs are in quaternary complexes (and bound to Sly1p) at any given time. However, consistent with a previous report (Søgaard et al., 1994) a significant percentage of the cellular Sed5 protein appears to be always associated with Sly1p. Small amounts of the exocytic v-SNAREs Snc1/2p but no vacuolar v-SNARE Nyv1p were detected to coprecipitate with Sly1p.

In a second set of experiments, immunoprecipitations from cleared cell lysates were performed with a polyclonal antibody directed against Bos1p. Under conditions where Bos1p was completely precipitated, fractions of total Sed5p, Bet1p, Sec22p, and, importantly, of Sly1p were coprecipitated. In contrast, the vacuolar v-SNARE Nyv1p was not detected in the anti-Bos1p immunoprecipitate (Fig. 4 B).

Sly1p does not stabilize Sed5p in vivo or core SNARE complexes generated in vitro
In a recent study, it was reported that the yeast Sec1 family member Vps45p stabilizes the endocytic t-SNARE Tlg2p by preventing its proteasomal degradation (Bryant and James, 2001). Since Sly1p to a large extent is complexed with Sed5p in living cells, we sought to investigate whether Sed5p would be prone to degradation in cells depleted of Sly1p. A haploid strain (GSF4) with the chromosomal SLY1 gene under transcriptional control of the GAL10 promoter and its isogenic wild-type strain were grown in galactose and then placed into glucose-containing medium to block Sly1p production in GSF4 cells. The mutant cells stopped multiplying around 12–14 h after medium shift. Lysates from GSF4 cells after different times after shutdown of SLY1 gene transcription and those from parallely grown wild-type cells were prepared and subjected to SDS-PAGE and immunoblot analysis. As shown in Fig. 5 A, the Sly1p level fell under that of wildtype cells \( \sim 10\% \) after shift to glucose medium, and after further 10 h Sly1p was hardly detectable with the antibody used. According to densitometric measurement of the intensity of immunoblot signals, no significant change of

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Figure 6. Sly1p does not influence the kinetics of assembly or the NSF-catalyzed disassembly of ER to Golgi SNARE complexes. (A) Samples of GST–Sec22p (5 μg) bound to glutathione agarose beads were incubated at 4°C for the indicated times with His-tagged Bos1p (15 μg), Bet1p (5 μg), and either free Sed5p (10 μg) or preformed Sed5p–Sly1p complex. Reactions were stopped with 5 vol of ice-cold buffer C, and beads were washed four times. Proteins bound to beads were characterized on Coomassie blue–stained gels. (B) SNARE complexes assembled with GST–Sec5p or with the heterodimeric GST–Sec5p–Sly1p complex were purified on glutathione agarose as described in the legend to Fig. 2 C. Complexes on beads were mixed with equal molar concentration of NSF and α-SNAP, 2 mM MgCl₂, 2.5 mM ATP in 20 mM Tris, pH 7.4, 100 mM NaCl, and 1 mM DTT for 20 min at 25°C. The reaction was stopped by dilution with ice-cold reaction buffer. As a control, the ATPase activity of NSF was inhibited by replacing MgCl₂ with 10 mM EDTA. GST–Sec5p and its associated proteins on glutathione agarose beads were analyzed by SDS-PAGE and Coomassie blue staining.
the cellular levels of either Sed5p or Bos1p could be observed over the time course followed (see legend to Fig. 5).

Neuronal core SNARE complexes assembled in vitro are resistant to SDS at room temperature and can be disassembled by heating only. In contrast, yeast SNARE complexes generated from recombinant proteins appear not to be resistant to SDS (Tsui et al., 2001). Therefore, we sought to examine whether Sly1p bound to in vitro–generated ER to Golgi SNARE complexes would gain resistance to SDS. As shown in Fig. 5 B, SNARE complexes exposed to SDS-containing buffer for only 5 min at room temperature readily disassembled regardless of whether Sly1p was bound or not. As expected, a ternary neuronal SNARE complex (Pabst et al., 2000) proved SDS resistant at room temperature under identical conditions. These results suggest that in living cells, the lifetime or the production of Sed5p is not dependent on Sly1p and that Sly1p does not confer SDS resistance to core SNARE complexes generated in vitro.

Sly1p does not influence the kinetics of SNARE complex assembly in vitro

The assembly of SNARE complexes in vitro is a slow process. This allowed us to easily follow the kinetics of their formation and to answer the question whether Sly1p might facilitate or slow down SNARE complex assembly. To GST–Sec22p on agarose beads, Bos1p, Bet1p, and either free Sed5p or Sly1p-bound Sed5p were added, and incubations at 4°C were allowed to proceed for up to 22 h. Complexes of the four SNAREs could be detected after ~3 h of incubation. About 20 h were needed for maximal complex formation. Although there was some variability in the extent of SNARE assembly in different experiments, no clear difference in the kinetics of complex formation was observed depending on whether Sly1p was present or not. A representative experiment (Fig. 6 A) shows that the formation of both types of complexes started to be seen at the same time point (3 h) and that Sed5p alone and the Sed5p–Sly1p heterodimer were incorporated into SNARE complexes with comparable kinetics. This was monitored by the densitometric assessment of Sec22p to Sed5p stoichiometry at different time points, which amounted to 1:0.3 and 1:0.3 at 3 h, 1:0.5 and 1:0.44 at 5 h, and 1:1.1 and 1:0.99 at 22 h in the absence and presence of Sly1p, respectively.

From this we conclude that the speed of SNARE complex formation from recombinant proteins is not significantly influenced by Sly1p.

Sly1p does not protect core SNARE complexes from disassembly by NSF/α-SNAP

The simultaneous binding of Sly1p and Sec17p to in vitro–generated ER to Golgi SNARE complexes (Fig. 3 B) allowed us to investigate whether the Sec1 family protein affects the disassembly of core complexes. To address this question, SNARE complexes were formed with either GST–Sed5p or Sly1p-bound GST–Sed5p and the remaining three SNAREs. The complexes, purified on and bound to glutathione agarose beads, were then treated for 20 or 40 min with a 1:1 mix of recombinant rat NSF and α-SNAP in the presence of either Mg-ATP or EDTA. As can be seen in Fig. 6 B, α-SNAP and NSF bound to both SNARE complexes under conditions where ATP hydrolysis was inhibited. Importantly, Sly1p did not protect the core complexes from disassembly when NSF was active. At the shortest reaction time tested (20 min), the SNARE complexes were completely disassembled without affecting the Sed5p–Sly1p interaction.

Sly1p contributes to the specificity of SNARE complex formation

It has been shown previously that SNAREs are promiscuous with respect to their integration into complexes assembled in...
vitro and that mammalian syntaxin-containing cognate and noncognate SNARE complexes have similar biochemical and biophysical characteristics (Fasshauer et al., 1999; Yang et al., 1999). Since in contrast to N-Sec1–bound syntaxin Sly1p-bound Sed5p can be assembled into a core fusion complex, we inquired into the possibility that Sly1p might interfere with the generation of noncognate SNARE complexes.

First, we tried to replace one of the ER to Golgi v-SNAREs by the exocytic v-SNARE Snc2p. Of the three cognate v-SNAREs, only Bet1p was absolutely needed for complex formation with free Sed5p (Fig. 7 A). The combination of Snc2p–Bos1p–Bet1p allowed tetrameric SNARE complexes to be formed with free and Sly1p-bound Sed5p. Interestingly, the combination of v-SNARES Snc2p, Sec22p, and Bet1p efficiently assembled into a tetrameric complex with free Sed5p but not Sly1p-bound Sed5p (Fig. 7 A). In similar experiments, SNARE complex assembly was also probed with the Golgi v-SNARE Ykt6p and the vacuolar v-SNARE Nyv1p. Of all three possible combinations, only Nyv1p–Sec22p–Bet1p (Fig. 7 B) or Ykt6p–Sec22p–Bet1p (Fig. 7 C) readily formed tetrameric complexes with free Sed5p. Importantly, none of these complexes was generated when Sed5p was first bound to Sly1p.

These results indicate that at least in vitro the Sec1 family member Sly1p can prevent the formation of noncognate tetrameric SNARE complexes.

Discussion

In the present study, we have used soluble ER to Golgi SNAREs for the assembly into specific complexes to address the question as to whether the Sec1 family member Sly1p, which binds with high affinity to the syntaxin Sed5p, affects SNARE complex formation or specificity. Surprisingly and in sharp contrast to the neuronal N-Sec1 and its associated syntaxin 1A, the Sed5p–Sly1p dimer proved as efficient as the free t-SNARE to assemble into equimolar complexes with its cognate v-SNAREs Bos1p, Sec22p, and Bet1p without affecting significantly the kinetics of complex formation or their NSF-catalyzed disassembly. Sly1p also bound to preassembled tetrameric SNARE complexes in vitro, and it could be precipitated with Sed5p, Bos1p, Sec22p, and Bet1p from yeast cell lysates. These results suggest that like the yeast exocytic fusion complexes (Carr et al., 1999) but unlike the neuronal ones (Yang et al., 2000) SNARE complexes of ER to Golgi transport are associated with its specific Sec1 family protein in vivo. From different studies, it now emerges that the interaction of Sec1 proteins with their cognate SNAREs (binding to NH2- and/or COOH-terminal region or to open or closed conformation of syntaxins) and/or with SNARE fusion complexes (formation of SNARE complexes on Sec1-bound syntaxins, association with preassembled SNARE complexes) is distinct for different family members. The results of our study also disfavor the often generalized view that Sec1 family proteins are negative regulators of trans-SNARE pairing, and they are at odds with the conclusion of previously published work (Lupashin and Waters, 1997) that Sly1p in vivo has to be released from Sed5p as a prerequisite for SNARE-mediated fusion of ER-derived vesicles with the cis-Golgi compartment.

In a previous in vitro study, a remarkable specificity of fusion-competent yeast SNARE complexes was found when SNARE proteins were fixed in lipid bilayers, although some complexes remained unexplained (McNew et al., 2000; Parlati et al., 2000). It was argued that analyzing specificity of complex assembly with soluble SNAREs lacking the COOH-terminal transmembrane domain might be less meaningful. However, the pairings of soluble SNAREs that we have observed in solution also are not simply random. For effective complex assembly, not every of the three ER to Golgi SNAREs could be replaced by nonsyntaxin SNAREs acting in different pathways. For example, the plasma membrane v-SNARE Snc2p, which in vivo is interchangeable with Snc1p, could replace Sec22p or Bos1p but not Bet1p. Ykt6p, which in vivo appears to act in Golgi transport and homotypic vacuole fusion (McNew et al., 1997; Ungermann et al., 1999), and the vacuolar v-SNARE Nyv1p could only replace Bos1p but not Sec22p or Bet1p. Interestingly, Snc2p in combination with Bos1p and Bet1p could readily assemble into SNARE complexes regardless of whether the syntaxin Sed5p was free or bound to Sly1p. Since Snc1/2p could also be recovered in immunoprecipitates from cell extracts with anti-Sly1p and anti-Bos1p antibodies, we take these findings as an indication that functional ER to Golgi SNARE fusion complexes in vivo can integrate Snc2p instead of Sec22p into the four-helix bundle. A similar conclusion was reached by Grote and Novick (1999).

However, presumably nonphysiological tetrameric complexes could be efficiently formed in vitro when Snc2p, Ykt6p, or Nyv1p, all R-SNAREs like Sec22p, replaced Bos1p. Complex assembly with two R-SNAREs was unexpected, since it has been proposed (Fasshauer et al., 1998) and backed in part by mutational analysis (Katz and Brennwald, 2000; Ossig et al., 2000) that functional SNARE complexes could accommodate only three glutamines and one arginine in the central layer of the four-helix bundle. We cannot exclude the possibility that some of these complexes form in vitro because the SNAREs used lack their membrane anchors. However, SNARE combinations that led to apparent nonphysiological core complexes in our study (Nyv1p replacing Bos1p, for example) were not investigated in the study with SNAREs embedded into lipid vesicles (McNew et al., 2000), precluding a direct comparison of the two investigations. Importantly, the presumably nonphysiological SNARE complexes could not be generated when Sed5p was prebound with Sly1p. This indicates that Sly1p tightly bound to the NH2-terminal helical region of syntaxin Sed5p could potentially interfere with the assembly of nonphysiological SNARE complexes and contribute to the specificity of SNARE fusion complexes in vivo. A chaperone-like function of Sec1 family members in preventing nonproductive SNARE complexes has indeed been discussed to explain phenotypes of certain yeast SNARE mutants (Katz and Brennwald, 2000). Preliminary experiments aimed at identifying nonphysiological SNARE complexes in yeast cells depleted of Sly1p did not give clear results. This might be explained by very small fractions of such complexes due to the fact that even 20 h after the switch-off of Sly1 gene expression a low level of Sly1p (~2.4% of wild-type level) (Fig. 5) still persists in mutant cells, which at this stage ceased multiplication and efficient protein transport activity (Ossig et al., 1991).
The first experimental evidence for a possible role of a Sec1 protein in adding to the specificity of SNARE complex formation is furnished by the results we have reported here. On the other hand, a general role for Sec1 family proteins to regulate the stability of their cognate syntaxins, as recently proposed (Bryant and James, 2001), could be excluded for Sly1p whose depletion from growing cells, as shown here, did not affect the cellular level of the syntaxin Sed5p to which it binds.

Materials and methods

Yeast strains and growth conditions

Yeast strains used in this study were MSUC1A3D (MATa ura3/ara3 leu2/leu2 his3/3tp1/tp1 lys2/lys2 ade8/ade8), MSUC1A (MATa ura3 leu2 his3 tp1 ade8) and its isogenic strain RPY137 (MATa slv1: KanMX4 ura3 his3 tp1 ade8 pRS416-PROA-SLY1), SY6210 (MATa ura3-52 leu2-3 lys2-A200 tp1-Δ901 lys2-Δ801 suc2-Δ9) and its isogenic strain GF54 (MATa LEU2-GAL10-SLY1 ura3-52 leu2-3 his3-A200 tp1-Δ901 lys2-Δ801 suc2-Δ9). RPY137 was constructed using techniques described previously (Peng et al., 2000). Yeast was grown in standard YPD or in selective media with the necessary additions.

Plasmid construction

All plasmids were propagated in the E. coli strain DH5α. The coding sequences of Sly1p (amino acids 1-666), Sed5N (amino acids 1-250), Sed5C (amino acids 251-324), the whole cytosolic region of Sed5p (amino acids 1-324), and Ykt6p (amino acids 1-195) were amplified by PCR from Saccharomyces cerevisiae genomic DNA and inserted into pGEX-TT, a vector for NH2-terminal GST fusion, to create pGEX-SLY1 (GST–Sly1p), pGEX-SED5N (GST–Sed5N), pGEX-SED5C (GST–Sed5C), pGEX-SEDS (GST–Sed5p), and pGEX-YKT6 (GST–Ykt6p), respectively. Full-length Sly1p (NH2-terminally His6-tagged) was expressed from pET19B-SLY1. We expressed His6-Sed5p/22 (amino acids 1-188) and GST–Sec22p from pQE30-SEC22 and pGEX-SEC22, respectively. They carry the SEC22 gene lacking the sequence encoding the COOH-terminal transmembrane domain. Expression of His6-Sed5p was described previously (Peng et al., 1999). His6-Bet1p and His6-Bos1p were expressed from plasmids pET14b-BET1 (SFBN358) and pET11d-BOS1 (SFBN358) described previously (Stone et al., 1997). Plasmids expressing the His6-Sec17p (full length), GST–Nvy1p (full length), and GST–Snc2p (amino acids 1-93) were provided by A. Mayer (Max-Planck-Gesellschaft, Tübingen, Germany), T. Sollner (Memorial Sloan Kettering Cancer Center, New York, NY), and D. Banfield (The Hong Kong University of Science and Technology, Hong Kong, China), respectively. Plasmid pRS416-ProA carrying two copies of IgG-binding motif of protein A was a gift of M. Peterson (University of California at San Diego, San Diego, CA) and was used to construct pRS416-ProA-SLY1, which expresses the protein A fusion of Sly1p (ProA-Sly1p).

Antibodies

A polyclonal antibody recognizing Snc1p and Snc2p was provided by P. Brennwald (Cornell University, New York, NY), and anti-Nvy1p antibody was from A. Mayer. Polyclonal anti-Bos1p and anti-Bet1 antibodies were affinity purified with GST–Bos1p and GST–Bet1p from antiserum produced in rabbits. Antibodies against Sly1p and Sed5p were described previously (Peng et al., 2000).

Protein expression and purification

Plasmids used for protein expression were transformed into the E. coli strain C41. Transformed cells were grown at 37°C to an absorbance of 1.0 at 600 nm and induced for recombinant protein expression with 1 mM IPTG for 4 h at 37°C or for 6 h at 30°C (GST–Sly1p). Cells from one liter cultures were collected by centrifugation and lysed by sonication in 10 ml of buffer A (25 mM Tris-HCl, pH 7.5, 400 mM KCl, 10% glycerol, 4% Triton X-100, 5 mM β-mercaptoethanol, 1 mM PMSF). Lysates were clarified by centrifugation and incubated with 1 ml of 50% glutathione-agarose (Amersham Pharmacia Biotech) or Ni2+-NTA agarose slurry (QIAGEN). The binding was conducted at 4°C for 60 min before the mixture was loaded onto a minicolumn. Affinity matrices were then washed with 2 ml of buffer B (25 mM Tris-HCl, pH 7.5, 500 mM KCl, 10% glycerol, 1% Triton X-100, 2 mM β-mercaptoethanol, 1 mM PMSF). The bound proteins were eluted with buffer C (which is buffer B without PMSF and with 150 mM instead 500 mM KCl) containing either 10 mM reduced glutathione (for GST fusion proteins) or 250 mM imidazole (for His-tagged proteins). In case the GST moiety was removed by thrombin, the column was treated with elution buffer containing 0.015 U/µl of thrombin (Sigma-Aldrich) at 4°C overnight.

In vitro binding assay

In vitro binding assays were performed essentially as described (Peng et al., 1999). Briefly, purified GST or GST fusion proteins reconjugated onto glutathione agarose beads were incubated at 4°C in buffer C for the indicated time period with purified proteins with or without His tag. Proteins on the extensively washed beads were dissolved by boiling in SDS buffer, separated by SDS-PAGE, and visualized by Coomassie blue staining or immunoblotting using specific antibodies.

Assembly of Sed5p–Sly1p and Snc1p complexes

The Sed5p–Sly1p complex was formed by mixing GST–Sly1p conjugated on glutathione beads and His6-Sed5p of ~1:3 molecular ratio (Sly1p to Sed5p) or by His-tagged Sed5p and Sly1p of equal molar amounts. Complex formation proceeded for 90–180 min at 4°C on a rotation wheel. The complexes on the beads (GST–Sly1p–His–Sed5p) or in solution (His–Sly1p–His–Sed5p) were then incubated with cognate and noncognate v-SNAREs. Complex formation with single or multiple v-SNAREs was conducted at 4°C for 17–22 h in 100 µl buffer C (see above). After extensive washing with buffer C, bound proteins were analyzed by SDS-PAGE and Coomassie blue staining. For saturation experiments, GST–Sed5p was first bound to beads fusing one or two His6-Sed5p of 90 min, the beads were then washed and incubated with His–Sly1p. Although the concentration of GST–Sed5p on beads was kept constant at ~1.0 µM, the concentration of His–Sly1p was varied between 0.1 and 10 µM. The binding reactions were performed at 4°C for 3 h. For Sly1p and Sec17p binding experiments, SNARE complexes were preassembled and isolated by binding to glutathione agarose beads through GST–Sec22p. Beads were washed in buffer C and used for incubation with excess of His6-Sly1p or His6-Sec17p at 4°C for 3 h. The samples were processed by SDS-PAGE followed by Coomassie blue staining. A minimal neuronal SNARE complex was formed as described (Pabi et al., 2000).

The molecular ratios of individual proteins in complexes were quantitated by scanning the Coomassie blue–stained gels (on Linscon). Images were processed by Photoshop 3.0, and the signal intensities of protein bands were analyzed by a program of Advanced Image Data Analyzer (Alida, version 2.11; Raytest Isotopenmessgeraete GmbH). In gels where His6–Sed5p was not separated, the value for signal intensity was divided by two to arrive at the approximate stoichiometries of the two proteins.

Immunoprecipitation and Western blot analysis

For immunoprecipitation experiments, cells of strain MSUC1A were grown at 25°C in YPD to an optical density of 0.5–1.0 at 600 nm. Cells (20–25 U at OD600) were collected at 4°C and washed with ice-cold wash buffer (20 mM Tris-HCl, pH 7.5, 20 mM Na2SO4, 20 mM NaF) to deplete the cells of ATP. In a 2-ml tube, cells were suspended in 1 ml of ice-cold IP buffer (50 mM Hepes, pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.5% NP-40) in the presence of protease inhibitors and lysed with 1.7 g of glass beads in a minibeadbeater for 4 min at full speed. Lysates were centrifuged at 500 g for 5 min to separate the cell material from glass beads and unbroken cells. The supernatant was centrifuged again at 16,000 g for 30 min at 4°C and transferred to a new tube to determine protein concentration with a Bio-Rad Laboratories protein assay kit using mouse IgG (Pierce Chemical Co.) as standard. The samples (lysate) were adjusted to 4 mg/ml of total protein with IP buffer. To remove proteins unspecifically adhering to protein A–Sepharose, lysates were precleared with protein A–Sepharose beads at 4°C for 2–3 h with rocking. For immunoprecipitation, 1 ml of the pre-cleared lysate was incubated at 4°C overnight with affinity purified anti–Bos1p antibody. Beads were washed five times with 1 ml of IP buffer. Proteins were eluted from beads by boiling in SDS buffer, separated by SDS-PAGE, and visualized by immunoblotting with specific antibodies. For lysates containing ProA-Sly1p, samples were treated essentially the same except that they were precleared with Sepharose CL–4B (Amersham Pharmacia Biotech) followed by ProA-Sly1p binding to IgG-Sepharose fast flow matrix (Pharmacia Biotech). Serial dilutions of lysates were probed on blots with anti–Sed5p and v–SNARE antibodies to quantify the relative amounts of these proteins in immunoprecipitates through calculation of band intensities with a Lumimager (Boehringer).

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