Transition from hemifusion to pore opening is rate limiting for vacuole membrane fusion

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Fusion pore opening and expansion are considered the most energy-demanding steps in viral fusion. Whether this also applies to soluble N-ethyl-maleimide sensitive fusion protein attachment protein receptor (SNARE)- and Rab-dependent fusion events has been unknown. We have addressed the problem by characterizing the effects of lysophosphatidylcholine (LPC) and other late-stage inhibitors on lipid mixing and pore opening during vacuole fusion. LPC inhibits fusion by inducing positive curvature in the bilayer and changing its biophysical properties. The LPC block reversibly prevented formation of the hemifusion intermediate that allows lipid, but not content, mixing. Transition from hemifusion to pore opening was sensitive to guanosine-5’-[(γ-thio)triphosphate. It required the vacuolar adenosine triphosphatase V₀ sector and coincided with its transformation. Pore opening was rate limiting for the reaction. As with viral fusion, opening the fusion pore may be the most energy-demanding step for intracellular, SNARE-dependent fusion reactions, suggesting that fundamental aspects of lipid mixing and pore opening are related for both systems.

Introduction

Protein-mediated fusion reactions are sensitive to membrane lipid composition (Chernomordik and Kozlov, 2003). Lipids such as phosphoinositides control the activity and concentration of fusion proteins (Simonsen et al., 2001). Lipids also directly influence bilayer mixing via their biophysical properties, i.e., charge and conformation, which determine the spontaneous curvature of membranes (Chernomordik et al., 1997). A physical influence on the membrane may also form the basis for effects of phosphatidylethanolamine on p97/p47–mediated Golgi–liposome fusion (Pecheur et al., 2002) and of cholesterol and sphingolipids, which are required for fusion of Semliki forest virus (Ahn et al., 2002).

Lysophosphatidylcholines (LPCs) exhibit positive spontaneous curvature (Fuller and Rand, 2001). They inhibit fusion of protein-free bilayers (Chernomordik et al., 1995; Wu et al., 1996), viral fusion (Chernomordik et al., 1997; Melikyan et al., 2000), exocytosis (Vogel et al., 1993), and microsome fusion (Chernomordik et al., 1993). LPCs show increasing inhibitory potency with increasing hydrocarbon chain length (Chernomordik et al., 1997), and they show equivalent effects in different fusion systems using very different proteinaceous machines. Therefore, LPCs probably act via lipid conformation and membrane curvature rather than by direct influence on fusion-relevant proteins.

Consequently, a universal physical intermediate of all bilayer fusion reactions was postulated that should be independent of the protein machinery inducing fusion and cannot be overcome in the presence of LPC. This intermediate may be a fusion stalk (Chernomordik and Zimmerberg, 1995; Kozlovsky et al., 2002; Chernomordik and Kozlov, 2003). In a stalk, the outer leaflets of two fusing bilayers are merged, whereas the inner leaflets remain separated. The stalk is a structure of negative membrane curvature. It can expand into a hemifusion state in which lipids from the inner monolayers associate to produce a new bilayer, the hemifusion diaphragm. Hemifusion is detectable for viral fusion systems, particularly if the normal fusion pathway is disturbed (Kemble et al., 1994; Melikyan et al., 1995, 2005; Chernomordik et al., 1998; Munoz-Barroso et al., 1998). It can also be generated in the polyethylene glycol–, hypotonic stress–, and cation-induced fusion of liposomes (Chanturiya et al., 1997; Lee and Lentz, 1997; Meers et al., 2000). Also, SNARE-dependent fusion can generate hemifusion states. Depending on the conditions, this can be unproductive alternative outcomes or intermediates on the pathway to full fusion (Giraudo et al., 2005; Lu et al., 2005; Reese et al., 2005; Xu et al., 2005).

Homotypic fusion of yeast vacuoles is a model reaction of SNARE-dependent intracellular fusion for which a cell-free...
fusion assay is available (Mayer, 2002; Wickner, 2002). Vacuole fusion proceeds in several steps and depends on numerous proteins. In brief, the following events are relevant to this study: priming is characterized by the ATP-dependent rearrangements of SNAREs, GTPases, and tethering factors and is a prerequisite for subsequent steps. cis-SNARE complexes are disassembled by the ATPase Sec18p/NSF and its cofactor Sec17p/α-SNAP, which is released from the vacuole (Mayer et al., 1996; Ungermann et al., 1998a). The second step, membrane attachment, requires the Rab-GTPase Ypt7p and leads to the formation of trans-SNARE complexes between the fusion partners (Mayer and Wickner, 1997; Ungermann et al., 1998b). Trans-SNARE pairing triggers a measurable release of calcium from the vacuole (Peters and Mayer, 1998; Merz and Wickner, 2004b). \( V_0 \), the membrane integral part of the vacuolar H\(^+\)-ATPase, is required to complete fusion (Peters et al., 2001; Bayer et al., 2003; Thorngren et al., 2004).

In this study, we have modulated bilayer conformation by LPC and explored the effects on the fusion pathway of yeast vacuoles. The assay of lipid perturbation promises important insights into the fusion mechanism, such as the timing of bilayer merger and factors required to induce it. We performed a kinetic analysis to map the LPC-sensitive step and correlated these data to molecular marker events for docking and post-docking events in order to determine the rate-limiting stage of vacuole fusion.

**Results**

**LPCs inhibit in vitro fusion of yeast vacuoles**

In vitro fusion of yeast vacuoles recapitulates the in vivo fusion of these organelles but provides accessibility to low and high molecular weight inhibitors. The reaction can be traced by a content-mixing assay relying on transfer of a luminal vacuolar protease between the fusion partners. Vacuoles are prepared from two strains. One strain (BJ3505) expresses pro-alkaline phosphatase (ALP) in its vacuoles, and the other (DKY6281) expresses the respective maturation enzyme proteinase A. Upon fusion between the vacuoles, proteinase A from one vacuole can mature pro-ALP from the other fusion partner and activate it. The resulting enzymatic activity can be assayed by spectrophotometry. It serves as a quantitative readout of fusion.

We investigated the effects of LPC on vacuole fusion by titrating LPCs of different acyl chain lengths into fusion reactions. LPCs spontaneously incorporate into lipid bilayers. They lack the fatty acid at the sn-C2 position of the glycerol backbone, conferring an inverted cone shape. At the C1 position, different fatty acids can be attached via ester linkages. We tested 1-capryl, 1-capryl, 1-lauroyl, and 1-myristoyl LPCs, carrying saturated fatty acids with 8, 10, 12, or 14 carbon atoms, respectively. All LPCs inhibited the fusion of yeast vacuoles (Fig. 1 A). The IC\(_{50}\) values increased from 30 \( \mu \)M for LPC-14 (1-myristoyl-2-hydroxy-sn-glycero-3-phosphocholine) to 3 mM for LPC-8 (1-capryl-2-hydroxy-sn-glycero-3-phosphocholine), depending on the hydrocarbon chain length of the respective LPC. A similar trend was reported for influenza HA fusion (Chernomordik et al., 1997). This suggests that LPCs inhibit vacuole fusion and viral fusion by the same principle. Because the proteins involved in both reactions are very different, LPCs most likely inhibit via a structural influence on the lipid bilayer rather than a deleterious effect on a fusion protein.

Recently, Thorngren et al. (2004) found that addition of the recombinant t-SNARE subunit Vam7p (rVam7p) leads to the accumulation of excess trans-SNAREs and renders vacuole fusion independent of several early acting components, such as ATP and Sec18p/NSF-mediated priming. LPC suppressed rVam7p-mediated fusion (Fig. 1 B), suggesting that it acts on a late step of vacuolar membrane fusion.

**LPCs block vacuole fusion after vacuole docking**

The latest reaction phase blocked by an inhibitor can be inferred from the kinetics with which the reaction becomes insensitive to the inhibitor. In such an experiment, a fusion reaction is...
incubated at 27°C. At certain time points, aliquots are removed, supplemented either with buffer (control; no change in the final fusion activity should be observed) or with inhibitor, and incubated further at 27°C until the end of a standard 70-min reaction period. Progression of an uninhibited reaction is monitored by transferring one aliquot to 0°C at every time point. This stops the reaction. The ice curve is indicative of fusion pore opening because there is no significant delay between contents mixing and the processing of the reporter pro-ALP (Merz and Wickner, 2004a). The reaction became insensitive to antibody inhibition of the priming ATPase Sec18p/NSF within 15 min (Mayer et al., 1996; Fig. 2). Gdi1p (guanine nucleotide disassociation inhibitor), which binds and partially extracts the Rab-GTPase Ypt1p from the membrane (Mayer and Wickner, 1997) and thereby blocks docking, inhibited up to 30 min. Inhibitors blocking the reaction between docking and fusion pore opening, such as the chelator 1,2-bis-(o-aminophenoxy)-ethane-N,N′,N′,N′′-tetraacetic acid (BAPTA) and guanosine-5′-(γ-thio)triphosphate (GTPγS), yield inhibition curves between those of Gdi1p and ice (Mayer et al., 1996; Peters and Mayer, 1998; Peters et al., 1999; Reese et al., 2005). LPC-14 and -12 showed such inhibition kinetics. The time courses for both inhibitors overlapped, suggesting that they affect the same late step of the reaction and act by the same mechanism. Because LPCs inhibit other fusion systems at the lipid-mixing stage (Chermomordik et al., 1998; Melikyan et al., 2000), this result suggests that lipid mixing is kinetically distinguishable from docking and significantly precedes contents mixing.

The analysis in the preceding paragraph does not rule out the possibility that LPCs additionally inhibit earlier reaction steps. Therefore, we scored molecular marker events for priming and docking. Priming separates vacuolar cis-SNARE complexes from opposing membranes. The in vitro fusion reaction of yeast vacuoles offers the opportunity to assay this association and to determine whether trans-SNARE pairing correlates to fusion activity (Ungermann et al., 1998a,b; Merz and Wickner, 2004b). To this end, vacuoles are prepared from strains expressing only the vacuolar t-SNARE Vam3p or the v-SNARE Nyv1p. After mixing these two populations of vacuoles and incubating them in the presence of postdocking inhibitors of fusion, the membranes are diluted and solubilized. The trans-association of v- and t-SNAREs can then be assayed by co-immunoprecipitation of the v-SNARE Nyv1p with antibodies to the t-SNARE Vam3p. In the absence of a fusion inhibitor, these complexes represent a mixture of three species: trans-SNARE complexes between vacuoles that did not complete fusion; trans-SNARE complexes converted into cis-SNARE complexes because they triggered successful fusion of the vacuoles; and cis-SNARE complexes formed independently of docking by reassociation of solitary v- and t-SNAREs subsequent to the completion of fusion, i.e., as a simple result of membrane mixing between the v- and t-SNARE–carrying membranes. In the
presence of late-acting fusion inhibitors, only the first species should accumulate. We tested for the formation of trans-SNARE complexes in the presence of LPCs. Without an ATP-regenerating system, priming and docking cannot occur and no Nyv1p could be coimmunoprecipitated with Vam3p (Fig. 4). In the presence of ATP, SNARE complexes formed. Simultaneous addition of the docking inhibitor Gdi1p completely blocked the formation of SNARE complexes, indicating that they resulted from docking. Similar to GTPγS, an established late-acting inhibitor of the postdocking phase, both LPC-14 and -12 allowed trans-SNARE pairing (Fig. 4), albeit only up to 50% of the uninhibited control. GTPγS permits docking as well as lipid mixing. Thus, the levels of trans-SNARE pairing reached in the presence of GTPγS should suffice to trigger lipid mixing. To assay the correlation between trans-SNARE pairing and fusion activity by independent means, we used another, recently identified inhibitor of vacuole fusion, myristoylated alanine-rich C kinase substrate effector domain peptide (MED; Fratti et al., 2004). MED sequesters phosphatidylinositol-4,5-bisphosphate, binds calmodulin with a $K_d$ of 4 nM, and penetrates the acyl chain zone of lipid bilayers with its five phenylalanines (Sundaram et al., 2004). All three effects could contribute to its inhibitory activity in vacuole fusion. We observed that MED strongly promoted mutual binding and cluster formation of vacuoles (unpublished data). In accordance with this, MED permitted SNARE pairing to at least the same levels as in the uninhibited control, and in some experiments even twofold more (Fig. 4 B). Nevertheless, MED blocked content as well as lipid mixing (see next paragraph). Together, these results suggest that trans-SNARE complexes can efficiently form in the absence of lipid mixing. Furthermore, they show that LPCs and MED permit trans-SNARE association but interfere with vacuole fusion downstream of docking.

**LPC inhibits lipid mixing in vacuolar membrane fusion**

To determine whether LPC inhibited merging of the membranes or fusion pore formation, we used a novel assay for vacuolar membrane fusion that detects the fusion-dependent flow of lipids from one fusion partner to the other (Reese et al., 2005). This assay is based on the concentration-dependent change in fluorescence intensity of lissamine rhodamine B 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine (Rh-PE; rhodamine dye coupled to dipalmitoyl-phosphatidylethanolamine). We incorporated self-quenching concentrations (3 mole percent) of Rh-PE into the membranes of vacuoles isolated from one of our standard fusion strains (DKY6281) and mixed them with a sixfold excess of unlabeled vacuoles isolated from the other fusion strain (BJ3505). Upon fusion, the labeled phospholipids are diluted into the unlabeled BJ3505 membrane, which increases the relative fluorescence of rhodamine. Fusion reactions of identical composition can be tested for content mixing via the common vacuole fusion assay, which is based on the activation of pro-ALP in BJ3505 vacuoles by processing proteases provided by DKY6281 vacuoles. We titrated LPC-12 and MED into fusion reactions to test in parallel how LPC influences lipid and content mixing (Fig. 5). LPC-12 and MED inhibited both content- and lipid-mixing signals to comparable degrees. Thus, LPC and MED already suppress the lipid-mixing step of vacuolar membrane fusion.

**LPC blocks formation of $V_o$-trans-complexes**

$V_o$ sectors undergo a transformation in the final phase of vacuole fusion, which can be diagnosed by the association of $V_o$ sectors from opposing membranes (Peters et al., 2001; Bayer et al., 2003). $V_o-V_o$ association preferentially affects a population of...
V₀ sectors associated with the vacuolar t-SNARE Vam3p. It is a diagnostic criterion linked to the fusion cascade because it depends on vacuole priming and docking and is sensitive to the postdocking inhibitor BAPTA but insensitive to the latest acting inhibitor of vacuole fusion, GTPγS. Lipid transition mapped to the same stage as V₀ transformation, i.e., it was insensitive to GTPγS but sensitive to BAPTA (Reese et al., 2005). This suggests that V₀ transformation coincides with the induction of lipid flow. If so, changing lipid bilayer conformation by LPC might in turn influence V₀ transformation. We tested the formation of V₀ trans-complexes in the presence of LPCs or MED (Fig. 6). Vacuoles were prepared from two strains carrying either an HA or an AU tag on the V₀ subunit Vph1p. The membranes were incubated under fusion conditions in the presence of various inhibitors. They were then solubilized, and Vph1p-HA was precipitated with a monoclonal anti-HA antibody. Vph1p-AU that was coimmunoprecipitated with Vph1p-HA was detected by Western blotting against the AU tag. Without ATP, a condition preventing priming and docking, no V₀ trans-association was detected (Fig. 6 A). With ATP, Vph1p-AU coimmunoprecipitated with Vph1p-HA, indicating a stable association of the proteins. Gdi1p, the docking inhibitor that prevents trans-SNARE pairing, suppressed the formation of these V₀ trans-complexes as completely as MED. In contrast, GTPγS, which inhibits after lipid mixing, permitted formation of V₀ complexes, although it reduced fusion by >95%. In line with previous observations (Peters et al., 2001), this confirmed that the V₀-V₀ complexes observed arose from trans-association of V₀-V₀ sectors from two fusion partners before completion of fusion. LPC prevented V₀ trans-association (Fig. 6 A).

We also checked whether LPC destabilized an existing trans-association of V₀. To this end, we ran fusion reactions for 40 min in the presence of GTPγS to generate V₀ trans-complexes. LPC was added and, after 10 min of further incubation, the vacuoles were solubilized and assayed for V₀ trans-complexes. The association of preformed V₀ trans-complexes was only slightly affected by subsequent addition of LPCs (Fig. 6 B), demonstrating that LPCs do not destabilize V₀ trans-complexes but interfere with their formation.

**LPC arrests vacuole fusion at a productive intermediate state**

Because LPCs reversibly incorporate into membranes, we tried to rescue the fusion block by removing LPCs again (Chernomordik et al., 1997; Melikyan et al., 2000). Fusion reactions were started in the presence of LPC. After 40 min, LPC was extracted from the membranes by floating the vacuoles through fatty acid–free bovine serum albumin. New fusion reactions were set up with these reisolated organelles and tested for their sensitivities toward inhibitors (Fig. 7). Good fusion activity was recovered after removing LPC. This shows that the vacuoles were not irreversibly damaged by LPC. Fusion after reversing the LPC block was insensitive to the docking inhibitor Gdi1p, as expected from the kinetic analysis (Fig. 2) and the docking assay (Fig. 4). The reaction stayed sensitive, however, to the postdocking inhibitors BAPTA and GTPγS. This indicates that the LPC block is reversible, that the intermediate accumulating in the presence of LPC is productive, and that the BAPTA- and GTPγS-sensitive steps cannot be passed before the LPC-sensitive stage.
To test the effect of LPC addition on V₀ trans-complexes that had formed,
inhibitors were used: 1

Figure 6. V₀ trans-complexes in the presence of LPC. (A) 15× standard
fusion reactions with cytosol were started using vacuoles from strains B) Vph1-HA and -AU and the indicated inhibitors. One sample did not con-
tain an ATP-regenerating system and was kept on ice. After 50 min, the
reactions were stopped by chilling on ice and adding 1 ml PS buffer. Vacuoles were pelleted by centrifugation [21,000 g, 2 min, 4°C]. Pellets were
solubilized in 1.3 ml of precipitation buffer (150 mM KCl, 0.5 mM
MnCl₂, 1% [wt/vol] Triton X-100, and 1 mM PMSF in PS), shaken for 5
min at 4°C, and pelleted by centrifugation (21,000 g, 3 min, 4°C). To 1.1
ml of the solubilisate, 5 μl of monoclonal anti-HA antibody [raw ascites;
5–7 mg/ml] and 20 μl of protein G-Agarose was added. After 1 h of
gentle shaking at 4°C, samples were washed with precipitation buffer, sol-
ubilized in SDS sample buffer, and analyzed by SDS-PAGE and Western
blotting. (B) Stability of existing V₀ trans-complexes against LPC-12. Fusion
reactions were run as in A but without LPC-12. 40 min after the start of
fusion, two of the samples (–10) were supplemented with LPC-12 in order
to test the effect of LPC addition on V₀ trans-complexes that had formed
during the first 40 min. All samples were incubated for another 10 min at
27°C, and then assayed for V₀ trans-complexes as in A. The following
inhibitors were used: 1 μM Gdi1p, 500 μM LPC-12, 10 μM MED, and 4 mM GTPyS.

Fusion proceeds with similar rates from early and late intermediates
The availability of several reversible and late-acting inhibitors
enabled us to compare the kinetics of vacuole fusion after re-
lease from these blocks. Should an early subreaction be rate
limiting, a reaction arrested at a stage past this rate-limiting
event should proceed significantly faster than a standard re-
adoption. In contrast, if a late event in the reaction cascade is rate
limiting, a reaction rescued from arrest before or at this rate-
limiting step should run with a similar rate as an uninhibited
reaction. We determined fusion kinetics after release from re-
versible blocks. Fusion was started in the presence of the inhib-
itors to allow the vacuoles to proceed up to the targeted
reaction stage. Then the vacuoles were resolated to remove the
inhibitors and used in new fusion reactions. After restarting the
reactions, aliquots were transferred to 0°C in 10-min intervals
to stop further fusion and monitor the progress of the reaction.
We compared the fusion rates after release from postdocking
blocks with LPC, GTPyS, and BAPTA to those of reactions
released from a high-salt block (250 mM KCl, which arrests
fusion at docking; unpublished data) and to a reaction without in-
hibitor, in which fusion had been prevented by omission of ATP.

Figure 7. The fusion block with LPC is reversible. A 20× standard fusion
reaction was started in the presence of 450 μM LPC-12. After 40 min at
27°C, 800 μl PS buffer with 15% (wt/vol) Ficoll and 0.5 mg/ml fatty acid-
free bovine serum albumin were added. 300 μl PS buffer with 4% (wt/vol)
Ficoll and 400 μl PS buffer were layered on top. Vacuoles were floated by
centrifugation (21,000 g, 5 min, 4°C), recovered from the 0%/4% inter-
face, and assayed for protein content. The recovered vacuoles were used in
new standard fusion reactions with the indicated inhibitors. After 80 min,
fusion was assayed. Control fusion reactions that had not been blocked
with LPC and released were run in parallel. Their fusion activity was deter-
mimed after 80 min at 27°C. Four independent experiments were averaged.
Ice values had been subtracted. For the blocked and released reactions, ice
values varied from 0.58 to 0.97 U and fusion activities varied from 2.2 to
4.2 U. For the unblocked reactions, ice values varied from 0.2 to 0.35 U
and control values varied from 2.5 to 4.2 U. The following inhibitors were used:
1 μM Gdi1p, 5 mM BAPTA, and 4 mM GTPyS.

MED was not used in this experiment because its inhibitory ef-
effect was not reversible by a simple washing of the membranes
(unpublished data). The rates after recovery from all reversible
blocks were similar to that of an uninhibited standard reaction
(Fig. 8). It appears unlikely that incomplete removal of the in-
hibitors after the block or slow dissociation of the inhibitors
from their targets could limit the fusion rate after rescue be-
cause the inhibitors are unrelated by chemical structure and by
mode of action. Therefore, the rate-limiting step of vacuole fu-
sion should lie at or beyond the stages sensitive to LPC,
BAPTA, and GTPyS. Because docking and lipid flow between
the fusion partners occurs in the presence of GTPyS (Reese et
al., 2005), our data suggests that the rate-limiting step in the fu-
sion pathway lies downstream of these events.

In sum, our results provide a basis to kinetically map the
requirement for lipid transition in the reaction sequence of vac-
uole fusion (Fig. 9). LPCs permit priming and docking. Lipid
conformation is crucial for the transition from docking to fu-
sion, i.e., to complete the BAPTA- and GTPyS-sensitive steps
and to induce the conformational change of V₀, which occurs
between docking and full fusion.

Discussion
Observations from this study can be integrated with previous
data to define the order of events in the postdocking phase of
vacuole fusion. LPC-blocked fusion reactions remain sensitive
to the fast-acting Ca²⁺ chelator BAPTA and to GTPyS, sug-
gesting that the LPC-sensitive step precedes events targeted by
these two inhibitors. In turn, BAPTA-blocked vacuoles become resistant to LPC. Trans-SNARE pairing is not prevented by any of these three inhibitors. These data suggest that trans-SNARE pairing is followed by an ordered sequence of steps defined by sensitivities to LPC, BAPTA, and GTPγS. Molecular events can now be grouped into this scheme: the transformation of V₀, which is diagnosed by their capacity to form V₀-V₀ trans-complexes; lipid transition by hemifusion; and fusion pore opening and contents mixing. LPC (Figs. 5 and 6) and BAPTA (Peters et al., 2001) block V₀ transformation and lipid transition between the fusion partners, whereas GTPγS permits these reactions and only blocks final opening of the fusion pore (Reese et al., 2005). Thus, V₀ transformation and lipid flow kinetically map to the same stage between BAPTA and GTPγS. Because LPC and, by inference, bilayer conformation and spontaneous curvature influence V₀ transformation, all kinetic and molecular observations link V₀ to the induction of lipid transition.

When released from blocks with postdocking inhibitors, vacuoles arrested by LPC, BAPTA, or GTPγS completed fusion with comparable kinetics as uninhibited reactions. This strongly suggests that reactions blocked with either of the three inhibitors still have to pass the rate-limiting step that seems to be the same for all of them and hence should lie past the latest acting of these inhibitors, GTPγS. Several studies demonstrate that the assay system is not rate limiting. The vacuolar content-mixing assay requires activation of the luminal pro-ALP by two soluble proteases (Pep4p and Prb1p) from the other fusion partner. A detailed study on the kinetics of pro-ALP conversion by Pep4p and Prb1p (Merz and Wickner, 2004a) showed that this step is not rate limiting in the in vitro fusion assay. Also, in vivo mature ALP appears with a half time of 6 min (Klionsky and Emr, 1989), which includes transport through the biosynthetic pathway to the vacuole. Therefore, pro-ALP activation in vacuoles takes much less than 6 min, and the rate-limiting process must be inherent in the fusion cascade.

What could this rate-limiting step of fusion be? Theoretical considerations (Chizmadzhev et al., 2000; Cohen and Melikyan, 2004) and experimental observations on influenza HA-mediated fusion (Frolov et al., 2000; Markosyan et al., 2000) have suggested that opening and expansion of a fusion pore rather than induction of lipid transition may be the most energy demanding step of fusion. In line with this, our observations place the rate-limiting step past the GTPγS-sensitive intermediate, i.e., at the transition from lipid mixing to fusion pore opening (Reese et al., 2005). This suggests that fusion pore opening rather than trans-SNARE pairing or induction of lipid mixing is rate limiting for vacuole fusion. Our results hence provide experimental evidence from a physiological SNARE-dependent fusion system that is consistent with the theoretical predictions postulating that opening the pore may carry the highest cost of energy and be rate limiting. With our current methodology, we could not differentiate between substeps of fusion pore opening, e.g., the opening of initial small pores and their expansion. Our content-mixing assay requires transfer of an ~40-kD protease and hence measures a relatively large pore. It may not allow for distinguishing very small initial pores, which could already be observed in other systems (Breckenridge and Almers, 1987; Zimmerberg et al., 1987, 1994; Curran et al., 1993; Lolli et al., 1995; Chanturiya et al., 1997; Lindau and Alvarez de Toledo, 2003), from the final, fully expanded state.

Because both SNAREs and viral fusion proteins form helical bundles, mechanistic parallels should be considered (Chernomordik et al., 1997, 1998; Melikyan et al., 2000). Type I viral fusion proteins fold back onto themselves and form α-helical bundles (Eckert and Kim, 2001; Park et al., 2003). Formation of fully stabilized helical bundles need not precede but can even follow lipid transition and the formation of a fusion pore (Melikyan et al., 2000; Borrego-Díaz et al., 2003; Markosyan et al., 2003). Complete helical bundle formation by viral fusion proteins hence does not correlate with lipid mixing. It may rather assist in opening the fusion pore or stabilize it after its formation. Lipid mixing is induced by the membrane-perturbing activity of viral fusion peptides (Chernomordik et al., 1997, 1998; Melikyan et al., 2000).
mordik and Kozlov, 2003; Tamm et al., 2003; Cohen and Melikyan, 2004). SNAREs could act in a similar fashion, i.e., the helical bundle might serve to pin the membranes together and assist in fusion pore opening once lipid transition has been triggered by membrane-disturbing compounds.

Several observations are compatible with a role of SNAREs in fusion pore formation. First, in artificial liposomes the outcome of fusion depends on the density of SNAREs reconstituted into their membranes (Lu et al., 2005; Xu et al., 2005). Completion of fusion requires higher SNARE densities and, by inference, higher densities of trans-SNARE complexes than the induction of lipid flow. Second, flipped SNAREs expressed on the surface of cells can also mediate slow fusion of such cells. If their transmembrane helices are replaced by glycosylphosphatidylinositol anchors so that they cannot span the entire bilayer, fusion pore opening is inefficient and hemifusion becomes the frequent end state (Giraudo et al., 2005). Also, truncation of the authentic transmembrane helix of SNAREs favors hemifusion (Xu et al., 2005). That SNAREs themselves might form part of the fusion pore was proposed based on the observation that substitutions in the transmembrane anchors of t-SNAREs change the electrophysiological properties of exocytic fusion pores measured in living cells (Han et al., 2004). This was taken as an indication that the transmembrane anchors of multiple SNAREs may form a barrel that constitutes the initial fusion pore and dissociates upon fusion pore expansion. However, biochemical evidence for the existence of a SNARE barrel is still missing. The effects observed are also consistent with other arrangements of SNAREs, and the substitutions might influence the interaction of SNAREs with other fusion-relevant proteins at exocytic sites, of which there are many (Hong, 2005).

In contrast to the observations made mainly in liposome systems with reconstituted SNAREs, which are consistent with a direct role of SNAREs in fusion pore opening, other observations on physiological membrane systems are less compatible with this view. Proteolytic removal of SNAREs on sea urchin cortical vesicles does not abolish fusion if contact is established by other means (Coorssen et al., 2003; Szule et al., 2003). Alternatively, trans-SNARE complexes in this system can also be disassembled by adding low concentrations of Ca^{2+}. This does not prevent subsequent fusion either (Tahara et al., 1998). Also, vacuolar trans-SNARE complexes were reported to become dispensable for completing fusion from the docked state (Ungermann et al., 1998b).

In the mechanochemical fusion model (Sollner, 2004), trans-SNARE complexes were proposed to induce lipid mixing by progressively zipping up from their membrane distal to their membrane-proximal parts. An alternative hypothesis is that they could already be fully zipped at the docking stage, mainly pinning the membranes together but requiring other factors to efficiently induce fusion. Several observations argue for the latter model. First, in this study we observed that reduction or enhancement of trans-SNARE pairing does not lead to comparable reduction or enhancement of lipid mixing. Second, during trans-SNARE complex formation in proteoliposomes, the entire SNARE domains can pair in a single step, but fusion follows complete core-complex formation with a long lag time (Zhang et al., 2004). Third, in case of progressive helix zippering toward the membrane anchors, one might expect competitive inhibition by peptides corresponding to the membrane-proximal COOH-terminal part of the v-SNARE. This was not observed (Melia et al., 2002), although analogous viral peptides could block formation of the helical bundle (Eckert and Kim, 2001). Finally, additional proteins are required for the transition from trans-SNARE pairing to lipid mixing or complete vacuole fusion, such as Vac8p (Wang et al., 2001) and V_0 (Peters et al., 2001; Bayer et al., 2003). Also, fusion of docked secretory vesicles in Drosophila melanogaster requires V_0, as demonstrated recently (Hiesinger et al., 2005).

LPC and the positive curvature of the membrane affect a step past membrane docking and block subsequent transformation of V_0 and lipid mixing. Therefore, we propose that docking and lipid mixing are separable events in SNARE-dependent fusion reactions. Trans-SNAREs may hold the membranes in close proximity and possibly modify downstream factors, in particular V_0, which are required to trigger fusion. Such an interpretation is consistent with the picture from viral fusion where membrane apposition and fusion pore opening via helical bundles are separable from the induction of lipid mixing via the fusion peptide (Qiao et al., 1999; Melikyan et al., 2000). In both systems, the last step, opening the fusion pore, appears to be the most demanding and rate-limiting event.

**Materials and methods**

LPC-8, LPC-10 (1-capryl-2-hydroxy-sn-glycero-3-phosphocholine), LPC-12 (1-lauroyl-2-hydroxy-sn-glycero-3-phosphocholine), and LPC-14 were purchased from Avanti Polar Lipids, Inc. 20 mM of stock solutions of the LPCs in PS buffer (200 mM sorbitol and 10 mM Pipes/KOH, pH 6.8) were stored at −20°C. Rh-PE was obtained from Invitrogen. Fatty acid-free bovine serum albumin and analytical grade dimethyl sulfoxide were obtained from Sigma-Aldrich. 3 mM Rh-PE stock solutions in DMSO were stored at −20°C. Production and purification of polyclonal antibodies to Sec18p, Sec17p, Vam3p, and Nyv1p were described previously (Haas and Wickner, 1996; Mayer et al., 1996; Nichols et al., 1997). Anti-HA monoclonal mouse antibodies (raw ascites) and monoclonal mouse AU antibodies were purchased from BabCo. Monoclonal mouse antibody to AIP was purchased from Invitrogen. Gdi1p was recombinantly expressed and purified from Escherichia coli (Haas et al., 1995). GTPαS was obtained from Roche Molecular Biochemicals, and BAPTA was obtained from Calbiochem. PS buffer was used as a basis for most solutions. MED was produced by solid phase peptide synthesis and purified by HPLC.

**Strains**

BJ3505 and DKY6281 were the standard fusion strains. For the assay of trans-SNARE pairing, we used deletion strains for Vam3p and Nyv1p (Nichols et al., 1997; Ungermann et al., 1998b). V_0 trans-complex formation was assayed with BJ3505 strains carrying either an AU [SBY129] or an HA tag [SBY119] on Vph1p (Peters et al., 2001).

**Vacuole isolation and fusion**

Yeast was precultured in yeast extract/peptone/dextrose (YPD) medium (6–8 h, 30°C). Overnight cultures were inoculated (30°C, 14–16 h; 225 rpm) in baffled 2-liter Erlenmeyer flasks with 1 liter of YPD medium. At an OD_600 of 2–5, cells were centrifuged (5 min, 4,000 g, 23°C; JA10 rotor), resuspended in 50 ml of 0.1 M Tris/Cl, pH 8.9, with 10 mM EDT, incubated for 10 min, 30°C, centrifuged (5 min, 4,000 g, 2°C; JA10 rotor), re-suspended in 15 ml SB (50 mM potassium phosphate, pH 7.5, and 600 mM sorbitol in YPD with 0.2% dextrose and 3,600 U/ml oxaetylase for BJ3505 and 1,800 U/ml for DKY6281), and transferred into 30-ml Corex tubes. Cells were incubated (20 min, 30°C), resuspended (1 min, 800 g, then 1 min, 1,500 g, 2°C; JA20 rotor), and resuspended in 2.5 ml of...
15% Ficoll 400 in PS buffer by gentle stirring with a glass rod. DEAE-dextran (200 μl for BJ3505 or 100 μl for DKY6281) was added from 0.4 mg/ml of a fresh stock in 15% Ficoll in PS buffer. The cells were incubated (2 min at 0°C and 75 s at 30°C), chilled again, transferred to a SW41 tube, and overlaid with 3 ml of 8% Ficoll, 3 ml of 4% Ficoll, and 1.5 ml of total volume of 31.5 ml of reaction buffer (20 mM Pipes/KOH, pH 6.8, 200 mM sorbitol, 125 mM KCl, 0.5 mM MgCl₂, 0.5 mM MnCl₂, 0.5 mM ATP, and 0.05 mM pefabloc SK, 7.5 ng/ml leupeptin, 3.75 μM α-amanitin, 37.5 ng/ml pepstatin A, 20 mM creative phosphate, and 35 U/ml creatine kinase). The ATP-regenerating system, containing MgCl₂, ATP, creatine phosphate, and creatine kinase, was added from a frozen 20× stock solution in PS buffer with 150 mM KCl. After 60 min at 27°C, fusion was assayed.

Content-mixing assay

Content mixing was determined via ALP activity. Mature ALP is generated by transfer of proteinase A (contained in DKY6281 vacuoles) into BJ3505 vacuoles (carrying pro-ALP). After completion of the fusion reaction, vacuoles were lysed with 500 μl of reaction buffer (20 mM Pipes/KOH, pH 8.9) for 4 min, 500 μl of 1 M glycine, pH 11.5, was added, and the OD was read at 400 nm. 1 U of fusion activity is defined as 1 μmol/l p-nitrophenol released per minute and microgram of BJ3505 vacuoles at 27°C. Ice values were subtracted.

Lipid-mixing assay

For the incorporation of Rh-PE into vacuolar membranes, 560 μl of freshly prepared vacuoles from strain DKY6281 were equilibrated in 800 μl PS buffer to 27°C for 1.5 min, 60 μl of 3 mM Rh-PE in DMSO was added under mild vortexing, and vacuoles were resolated. 500 μl PS buffer containing 15% (wt/vol) Ficoll were added and mixed. On top, 300 μl PS buffer containing 4% and 400 μl PS buffer containing no Ficoll were layered. After centrifugation (5 min, 4°C, 11,500 g), the labeled vacuoles were recovered from the 4%/0% interface. 3× standard fusion reactions contained a sixfold excess of unlabeled vacuoles from strain BJ3505 over Rh-PE-labeled vacuoles isolated from strain DKY6281. For fluorescence measurements, the reactions were transferred into black noncoated microtiterplates (Nunc). Rhodamine fluorescence was measured with a fluorescence plate reader (SpectraMax GeminiXS; Molecular Devices) at 538 nm for all samples. The average of the measurement after addition of Triton X-100 served as the fluorescence after infinite dilution, resulting in ΔF/F(TX-100) values. The resulting value for the 0-min time point was set to 0. Content mixing was measured in parallel samples via the established assay of ALP activity. To this end, 31.5 μl aliquots with an identical composition to the ones measured in the fluorescent plate reader were incubated at 27°C for 70 min.

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