A guanine nucleotide exchange factor (GEF) limits Rab GTPase–driven membrane fusion

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The identity of organelles in the endomembrane system of any eukaryotic cell critically depends on the correctly localized Rab GTPases, which bind effectors and thus promote membrane remodeling or fusion. However, it is still unresolved which factors are required and therefore define the localization of the correct fusion machinery. Using SNARE-decorated proteoliposomes that cannot fuse on their own, we now demonstrate that full fusion activity can be achieved by just four soluble factors: a soluble SNARE (Vam7), a guanine nucleotide exchange factor (GEF, Mon1–Ccz1), a Rab–GDP dissociation inhibitor (GDI) complex (prenylated Ypt7–GDI), and a Rab effector complex (HOPS). Our findings reveal that the GEF Mon1–Ccz1 is necessary and sufficient for stabilizing prenylated Ypt7 on membranes. HOPS binding to Ypt7–GTP then drives SNARE-mediated fusion, which is fully GTP-dependent. We conclude that an entire fusion cascade can be controlled by a GEF.

Organelles of the endomembrane system continuously exchange membranes by fusion and fission reactions but maintain their overall identity. Rab GTPases cooperate with specific tethering complexes and organize the fusion of vesicles with organelles through membrane-localized SNAREs and thereby serve as signposts of organelle identity (1, 2). Newly made Rabs are posttranslationally lipidated by a geranylgeranyl transferase complex and kept soluble thereafter by a Rab escort protein (REP), which binds both the GTPase domain and the hydrophobic hydrocarbon tail (3). After delivery to the right membrane, which is paralleled by REP release, a Rab–specific GEF promotes GTP loading on the organelle surface. This enables the Rab to bind to specific effectors and promote fusion (2, 4, 5).

Turnover of Rabs on membranes requires a GTPase-activating protein, which completes the active catalytic site and stimulates GTP hydrolysis. The resulting Rab–GDP can then be either reactivated or extracted by the GDI, a functional homolog of REP (6), allowing for another cycle of membrane delivery and activation.

The exchange of consecutively acting Rabs as part of vesicle and organelle maturation is an essential part of organelle biogenesis both in exocytosis and endocytosis (2, 7–11). Such a process requires coordination between Rab recruitment and activation and subsequent inactivation of the previous Rab, a process named Rab cascade (12).

Despite extensive structural insights into Rab interactions with its regulators, the molecular mechanism of specific delivery of Rabs to the right surface is still poorly understood. Previous studies suggested that organelle-specific proteins other than GEFs are needed to determine efficient Rab delivery to membranes (13–18). However, GEFs that are relocalized to mitochondria in vivo also recruit their corresponding Rab (5, 19, 20), and, in vitro, GEFs can drive Rab delivery and activation on liposomes (21). Although this puts GEFs into the center, it remains unclear whether other cytosolic or membrane-bound factors may be involved in Rab localization during an authentic membrane fusion process.

To understand the underlying principles, we set out to determine the minimal machinery of Rab delivery, activation, and effector recruitment to membranes. We now present the functional reconstitution of Rab-driven membrane tethering and subsequent fusion based on just four soluble factors. Our data unravel the simplicity of how an organelle can become fusion-competent during its maturation and illustrate the importance of the correct spatiotemporal localization of GEFs.

Results

Activation of purified prenylated Ypt7 and Vps21

Organelles such as endosomes recruit Rabs from a cytosolic pool to their surface. To recapitulate this process in vitro and unravel the required components, we established protocols to generate functional Rab–REP or Rab–GDI complexes with the yeast Rab5 (Vps21) and Rab7 (Ypt7) using the published prelyation machinery of the two Rab geranylgeranyl transferase subunits Bet2 and Bet4, REP (Mrs6), GDI (Gdi1) for all purified proteins, see Fig. S1), and geranylgeranyl pyrophosphate (21–23). We routinely obtained up to 90% prelyation in our reactions (Fig. 1A). Initially, we tested membrane association of the
Rab–REP complexes and observed spontaneous partitioning of only the prenylated Ypt7 (pYpt7) and Vps21 (pVps21) into protein-free liposomes (Fig. 1B). To test for functionality, we used liposome-bound Rabs to monitor their GEF-dependent activation. For this, we preloaded the prenylated Rabs Ypt7 and Vps21 with MANT–GDP and recorded the loss of fluorescence upon nucleotide exchange (24, 25). We observed efficient activation of Rabs regardless of their prenylation or the presence of REP (Fig. 1C). Only when His-tagged Rabs were directly bound to liposomes carrying DOGS-NTA, Rabs were activated with higher efficiency, as reported before (21, 25). We thus conclude that the in vitro prenylated Rabs can be recognized and activated by the corresponding GEF protein.

**Prenylated Ypt7 drives membrane fusion**

To test whether the prenylated Rab can drive an authentic fusion reaction, we focused on the vacuolar Rab Ypt7 in the context of reconstituted vacuole fusion. Previous studies established that proteoliposomes carrying vacuolar SNAREs with transmembrane domains and co-reconstituted prenylated Ypt7, purified from yeast cells, can fuse in the presence of the soluble SNARE Vam7, the SNARE chaperones Sec17/α-SNAP.
and Sec18/NSF, which activate SNAREs in the presence of ATP, and the HOPS tethering complex (26–28). HOPS binds to Ypt7 on both membranes to drive tethering and assemble SNAREs (29–32) (Fig. 1D). Fusion was assayed by mixing of the luminal dyes biotin–phycoerythrin and Cy5-streptavidin, present in separate tester liposomes, which produce a strong FRET signal after luminal mixing (26).

We used previously established conditions of optimized lipid composition and physiological SNARE concentrations, which makes fusion entirely Ypt7- and HOPS-dependent (33, 34). We also used a heterotypic fusion assay, where the Qa and Qb SNAREs Vam3 and Vti1 reside on one membrane, whereas the R-SNARE Nyv1 is on the other proteoliposome (Fig. S2). However, we now omitted prenylated Ypt7 from the reconstitution but provided the prenylated Rab as a soluble factor, either in complex with REP or GDI. Thus, fusion should depend exclusively on the added Ypt7 but should require its GEF Mon1–Ccz1, which was included in our fusion assays.

We initially titrated the Ypt7–REP complex to the fusion reaction, which was generated either in the presence or absence of GGPP. Only prenylated Ypt7 was able to support fusion, indicating that our prenylated Ypt7 is fully functional (Fig. 1, E and F). We noticed that fusion was also observed in the presence of GDP, as reported previously (27, 34). We therefore extended our titration of pYpt7–REP to the fusion assay and observed clear GTP dependence at lower pYpt7–REP concentrations, whereas, at higher concentrations, both GDP and GTP stimulated fusion equally (Fig. 2A, top panels). HOPS also binds GDP-loaded Ypt7 in vitro (32, 34). However, cells possess GDI to extract GDP-loaded Rabs (35). Indeed, when GDI was included, GDP-dependent fusion was strongly suppressed (Fig. 2A, bottom panels).
To ask whether recruitment of prenylated Ypt7 onto membranes occurs exclusively by the corresponding Mon1–Ccz1 complex, we triggered nucleotide exchange by EDTA and Mg\(^{2+}\)/H\(_{11001}\) and observed the same Ypt7 triggered fusion as before (Fig. 2B). GDP-dependent fusion was likewise suppressed by addition of GDI. We noticed that, after addition of GDI, more pYpt7–REP complex was required to obtain the same fusion as observed in the absence of GDI. This indicates that the combination of GEF and GDI restricts the pool of readily available Ypt7 for fusion.

A GEF can act as a limiting factor to recruit pYpt7 for fusion

In the cell, cytosolic Rab proteins are in complex with GDI. We therefore tested the pYpt7–GDI complex to drive liposome fusion. With GDP as a nucleotide, no fusion was observed, regardless of whether we increased the amount of pYpt7–GDI complex (Fig. 3A, top panels, green lines, and B). However, in the presence of GTP, robust fusion was observed (Fig. 3A, top panel, red lines; quantified in Fig. 3B). This strongly differs from our observations of the pYpt7–REP complex (Fig. 2, A and B). When we used Mg\(^{2+}\) and EDTA instead of GEF, fusion was as efficient (Fig. 3A, bottom panels; quantified in Fig. 3B).

The analysis of the membrane association of pYpt7–REP and pYpt7–GDI in the presence or absence of Mon1–Ccz1 clearly illustrates the reason for the difference seen in the fusion assays (Fig. 3C). Although pYpt7–REP allows for spontaneous membrane association of the Rab GTPase, as observed before (Fig. 1B), pYpt7 bound to GDI only inserted into the membrane in the presence of both its GEF and GTP. Noteworthy is that the amount of Ypt7 associating with membranes from the REP complex did not increase in the presence of the GEF and GTP. This strongly suggests that the required GEF, Mon1–Ccz1, could be limiting in the course of events that finally lead to fusion.

To test for the minimal requirements of pYpt7–GDI–dependent fusion, we selectively omitted each of the fusion components from the assay. Fusion was entirely blocked without pYpt7–GDI, Mon1–Ccz1, Vam7, and HOPS (Fig. 4, A and B). Neither ATP nor Sec17 and Sec18 were required under these conditions (Fig. 4, C and D), as explained by the setup of our heterotypic fusion assay, where SNARE assembly is driven by the soluble Vam7. This shows that prenylated Ypt7 in its active state drives HOPS-dependent fusion with all previously established characteristics of content mixing.

Of the four required factors, each may support Rab localization to membranes. Conceptually, the GEF is the most upstream factor, as it activates the prenylated Rab and therefore allows for effector binding and prevents Rab extraction by GDI. GEFs should thus be able to time and localize a reaction cascade in vivo. To test this, we isolated the corresponding proteoliposomes after fusion. Prenylated Ypt7 was only found in the pellet of the liposome fraction, when Mon1–Ccz1 GEF was present (Fig. 4E, lanes 8–10). In the presence of GDP, little Ypt7 was bound (Fig. 4E, lane 5), whereas HOPS addition stabilized Ypt7 on membranes (Fig. 4E, lanes 8 and 9). Interestingly, HOPS
binding to membranes was also observed without Ypt7 but did not contribute to fusion (Fig. 4E, top panel). This indicates that Ypt7-binding may orient the HOPS for fusion on membranes. In agreement with these observations, we could completely limit fusion by sequentially titrating down Mon1–Ccz1 (Fig. 5A; quantified in Fig. 5B). Our data thus reveal that the GEF controls membrane localization and activation of prenylated Ypt7 and, thus, the entire fusion cascade.

Discussion

Here we uncover the minimal conditions under which organelle membranes can be made fusion-competent. Just four soluble factors are needed for Rab–GTP–dependent fusion: the GEF complex Mon1–Ccz1, the pYpt7–GDI complex, the HOPS tethering complex, and the SNARE Vam7. We demonstrate that the GEF is the most upstream factor limiting the fusion reaction by regulated spatiotemporal activation of pYpt7. Subsequently, HOPS interacts with membrane-bound pYpt7–GTP and triggers membrane fusion.

These observations agree with several in vivo and in vitro studies. GEF relocalization to mitochondria relocalizes the corresponding Rab as well (19, 20). Furthermore, the GEF complex TRAPP can only recruit and activate pYpt1–GDI on liposomes (21). We now extend these studies by embedding Rab activation into a fusion cascade that is entirely GEF-dependent.

We and others (21) did not find evidence that additional factors are required for dissociation of the Rab–GDI complex. However, we cannot exclude that additional factors facilitate this process in vivo to provide another layer of regulation; for instance, accelerated Rab membrane association. Indeed, Pra1 stimulates reconstituted Rab5-dependent fusion (18), and Pra1-mediated GDI displacement has been postulated to be a critical intermediate of Rab delivery (13–17). Other mechanisms, such as posttranslational modifications, may enhance GDI displacement and membrane recruitment of Rabs in vivo (36). At least Ypt7 and Vps21 efficiently partitioned by themselves into membranes out of the REP complex (Figs. 1 and 3C), and Ypt7 just requires Mon1–Ccz1 to be stabilized on membranes by nucleotide exchange in the presence of GDI (Figs. 3C and 4E), as observed for Ypt1 and Ypt32 in the presence of the TRAPP complexes (21).

In the context of Rab–effector interactions, GDI is critical to buffer any available Rab–GDP, making Rab–GTP the exclusive effector binding site on membranes. At least for Ypt7, we quantified the relative abundance of Rabs and effectors in the cell. Here the amount of available Rab appears to be limiting compared with the intracellular amount of effector proteins.3 The main role of the GTPase-activating protein is then a constant shift of uncomplexed Rab–GTP to the GDI-extractable Rab–GDP form (5, 37). This, in turn, also strengthens the role of GDI in the cell by not only providing a membrane shuffling mechanism but also by giving higher specificity to membrane-bound Rab–GTP in terms of the Rab–effector interaction. The coop-

3 P. Carpio Malia, L. Langemeyer, A. Perz, D. Kümmel, and C. Ungermann, unpublished observations.
GERT-limited membrane fusion

Figure 5. Mon1–Ccz1 drives activation of Ypt7 and, therefore, fusion. A, the amount of Mon1–Ccz1 was titrated down in the preincubation time of the fusion assay. As before, fusion was started after allowing for nucleotide exchange of 80 nM pYpt7–GDI for 15 min. B, quantification of two independent fusion reactions.

eration of GEF and GDI thus sharpens the boundaries between GDP- and GTP-bound forms (2, 5).

Previous work on the Legionella DrrA protein implied that the extraction of Rab from the GDI complex directly correlates with the GEF activity of the protein (38–41). Compared with intracellular GEFs, Legionella DrrA appears to be hyperactive and induces a dramatic conformational change to enforce efficient nucleotide exchange. This may be required for the pathogenicity of Legionella by restricting Rab1 to the Legionella-containing vacuole (39–41). Importantly, our work now extends these findings to intracellular GEFs such as the Mon1–Ccz1 complex.

Factors that regulate the membrane association of GEFs consequently have a strong impact on the correct spatiotemporal localization of effector proteins. For TRAPP, Arf1–GTP seems to be one critical factor (21); for Mon1–Ccz1, Rab5–GTP is a likely recruiter (42–46). Similarly, Mon1–Ccz1 localization will depend on corresponding membrane interactors, which need to be dissected in future experiments.

Importantly, GEFs so far require mainly peripheral membrane proteins for their binding, although membrane composition is likely also involved, such as phosphoinositides. This suggests that cytosolic proteins could sequentially recruit each other to a maturing organelle to complement the membrane-embedded SNARE proteins and thus specify fusion competence. We are confident that the dedicated orchestration of Rabs and their regulators can explain the principle of organelle maturation not only on endosomes but along the entire endomembrane system.

Materials and methods

Yeast strains

Yeast strains overexpressing Mon1–Ccz1 and HOPS have been described previously (24, 29).

Expression and purification of Rab GTPases

GST-TEV-Ypt7 and GST-TEV-Yps21 were expressed and purified with slight modifications as described previously (24). Cells were lysed in a Microfluidizer (model M-110L, Microfluidics, Newton, MA), and the lysate was clarified by centrifugation at 40,000 × g for 30 min at 4 °C and loaded onto a pre-equilibrated Protino GST/4B 1-ml column (Macherey & Nagel). Protein was extensively washed and eluted in lysis buffer (50 mM Tris (pH 7.4), 300 mM NaCl, and 2 mM MgCl2) containing either 20 mM glutathione or TEV protease for cleavage at 4 °C overnight. Proteins were then dialyzed into assay buffer (50 mM HEPES, NaOH (pH 7.4), 150 mM NaCl, and 2 mM MgCl2), changing the buffer twice.

Expression and purification of Rab GGTase and Rab escort protein

pGATEV-Bet2 and pET30a-Bet4 (47) were co-expressed in Escherichia coli BL21 Rosetta, induced with 0.25 mM isopropyl 1-thio-β-D-galactopyranoside for 16 h at 18 °C. The Rab escort protein Mrs6 (48) was expressed and purified the same way. Cells were lysed in 50 mM Tris (pH 8.0), 300 mM NaCl, 2 mM β-mercaptoethanol, and 1 mM PMSF as described above. The cleared lysate was loaded on a Hi-Trap nickel-Sepharose column (GE Healthcare) equilibrated with the lysis buffer. The column was washed extensively with lysis buffer containing 30 mM imidazole, and bound protein was eluted with a linear 30 mM to 300 mM imidazole gradient over 30 column volumes. Fractions containing GGTase-II and REP, respectively, were pooled and dialyzed against assay buffer, which was changed twice.

Expression and purification of Gdi1

Competent E. coli Rosetta BL21 cells were transformed with pGEX-6P-Gdi1 (kindly provided by C. Fromme, Weill Institute, Cornell University). A single colony was picked from selection plates, and 2 liters of culture was grown to an A600 of around 0.8. Expression was induced with 0.25 mM isopropyl 1-thio-β-D-galactopyranoside, and cells were incubated at 18 °C overnight. Cells were lysed in PBS containing 1 mM PMSF, 2 mM MgCl2, and 5 mM β-mercaptoethanol, and the cell homogenate was cleared as described above. Cleared lysate was loaded on a Proteino GST/4B 1-ml column (Macherey & Nagel). Protein was eluted after extensive washing by cleaving the GST tag by incubating with Precision protease at 16 °C for 2 h.

Tandem affinity purification

Purifications were conducted essentially as described previously (29, 49). Three liters of culture were grown at 30 °C to A600. Cells were harvested by centrifugation and lysed in buffer containing 50 mM HEPES-NaOH (pH 7.4), 150 mM NaCl, 1.5 mM MgCl2, 1× Protease-Inhibitor-Mix FY (Serva), 0.5 mM PMSF, and 1 mM DTT. Lysates were centrifuged for 90 min at 100,000 × g, and supernatants were incubated with IgG-Sepharose (GE Healthcare) for 1.5 h at 4 °C. Beads were sedimented by centrifugation at 800 × g for 5 min and washed with 15 ml of lysis buffer containing 0.5 mM DTT. Bound proteins were eluted by TEV cleavage overnight and analyzed on SDS-PAGE.
**In vitro prenylation of Rab GTPases**

The assay was performed as described previously (50) with slight modifications. Equimolar amounts of GTase preloaded with either GDP (Sigma-Aldrich) or MANT–GDP (Jena Bioscience) (51) and the Rab escort protein were incubated with a third of the amount of geranylgeranyl transferase and a 10-fold excess of geranylgeranyl pyrophosphate (Sigma-Aldrich, Munich, Germany) in assay buffer containing 1 mM DTT. The mixture was incubated at 37 °C for 30 min and used for subsequent assays. To obtain prenlated Ypt7–GDI, the prenylation reaction was performed as described previously (21).

**Membrane association assay**

Association of prenylated Rab GTPases to membranes was followed by incubation of 50 pmol of Rab GTase with 50 nmol of liposomes (61 mol % palmitoyl-oleoyl-phosphatidylcholine, 18 mol % palmitoyl-oleoyl-phosphatidylethanolamine, 10 mol % palmitoyl-oleoyl-phosphatidylserine, 2 mol % phosphatidylinositol 3-phosphate, 8 mol % ergosterol, and 1 mol % diacylglycerol) (25) for 30 min at 30 °C. Liposomes were pelleted by centrifugation at 20,000 X g at 4 °C. Proteins in the supernatant and pellet fractions were separated and analyzed by SDS-PAGE and Coomassie staining.

**Nucleotide exchange assays**

GEF assays were performed as described previously (51). 50 pmol of MANT–GDP (Jena Bioscience)-loaded Rab GTase was mixed with increasing amounts of GEF, and after baseline stabilization, exchange was triggered by addition of 0.1 mM GTP to the reaction. Nucleotide exchange was followed by measuring the decrease in fluorescence of the MANT fluorophore in a SpectraMax M3 multi-mode microplate reader (Molecular Devices). Where indicated, 50 nmol of liposomes was added to the reaction (25). Liposomes were composd as described above, and, where indicated, 10 mol % DOGS-NTA (1,2-dioleoyl-sn-glycero-3-[(N-(5-amino-1-carboxypentyl)iminodiacetic acid)succinyl]) was added and compensated by reduction of the palmitoyl-oleoyl-phosphatidylcholine amount.

**Fusion assay with reconstituted proteoliposomes (RPLs)**

Fusion of RPLs, including purification of the involved proteins, was performed as described previously (26, 52) with the indicated protein-to-lipid ratios. One set of RPL carried Nyv1 and the other set Vt1 and Vam3. Where indicated, RPLs were preincubated with prenylated Rab GTase and 100 nM Mon1–Cc21. The mixture was incubated for 15 min at 27 °C to allow for membrane association of the prenylated Rab–GTase and nucleotide exchange. Where indicated, 500 nM Gdi1 was added afterward, and reactions were incubated for another 5 min before triggering fusion. Fusion was triggered by addition of 50 nM HOPS complex, 50 nM Sec18, 600 nM Sec17, and finally 100 nM Vam7. To follow full fusion of liposomes, content mixing and the subsequent increase in fluorescence were monitored in a SpectraMax M3 multi-mode microplate reader (Molecular Devices).

**Fractionation of the RPL fusion assay**

To test for membrane association in the RPL fusion assay, the fusion reaction containing pYpt7–GDI was followed in the SpectraMax M3 multi-mode microplate reader (Molecular Devices) as above for 45 min. Afterward, samples were transferred to centrifugation tubes and spun for 15 min at 20,000 X g and 4 °C. Pellet and supernatant fractions were separated, and the volume was adjusted to 500 µl with assay buffer containing 0.1% Triton X-100. Proteins were TCA-precipitated and subjected to SDS-PAGE followed by Western blotting. Proteins were detected using antibodies against GST–Ypt7 and the TAP tag antibody (Invitrogen, CAB1001) directed against the TAP tag at Vps41 and Ccz1, respectively.

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GEF-limited membrane fusion

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