Structures of the yeast dynamin-like GTPase Sey1p provide insight into homotypic ER fusion

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Homotypic membrane fusion of the endoplasmic reticulum is mediated by dynamin-like guanosine triphosphatases (GTPases), which include atlastin (ATL) in metazoans and Sey1p in yeast. In this paper, we determined the crystal structures of the cytosolic domain of Sey1p derived from Candida albicans. The structures reveal a stalk-like, helical bundle domain following the GTPase, which represents a previously unidentified configuration of the dynamin superfamily. This domain is significantly longer than that of ATL and critical for fusion. Sey1p forms a side-by-side dimer in complex with GMP-PNP or GDP/AlF4− but is monomeric with GDP. Surprisingly, Sey1p could mediate fusion without GTP hydrolysis, even though fusion was much more efficient with GTP. Sey1p was able to replace ATL in mammalian cells, and the punctate localization of Sey1p was dependent on its GTPase activity. Despite the common function of fusogenic GTPases, our results reveal unique features of Sey1p.

Introduction

The ER network consists of a continuous membrane system of tubules and sheets (Shibata et al., 2006). ER membranes are interconnected by homotypic fusion to maintain a reticular pattern and proper functioning. In metazoans, fusion is mediated by a class of dynamin-like GTPases called atlastin (ATL; Hu et al., 2009; Orso et al., 2009). In other eukaryotic organisms lacking ATL, a similar class of GTPases, including Sey1p in Saccharomyces cerevisiae (Hu et al., 2009; Anwar et al., 2012) and RHD3 in Arabidopsis thaliana (Zhang et al., 2013), has been identified. Deletion of Sey1p or depletion of ATL proteins in cells results in unbranched ER (Hu et al., 2009), delayed ER fusion (Anwar et al., 2012), or even ER fragmentation (Orso et al., 2009). When purified and reconstituted in vitro, members of both classes have been shown to mediate membrane fusion (Orso et al., 2009; Bian et al., 2011; Anwar et al., 2012; Zhang et al., 2013).

GTPase-based ER fusogens play critical physiological roles. In Candida albicans, deletion of Sey1p results in decreased virulence (Yamada-Okabe and Yamada-Okabe, 2002). In A. thaliana, lack of RHD3 leads to short and wavy root hairs and cell expansion defects (Schiefelbein and Somerville, 1990; Chen et al., 2011; Stefano et al., 2012), and the deletion of another family member causes lethality (Zhang et al., 2013). In Drosophila melanogaster and Danio rerio, depletion of ATL causes neuronal defects (Lee et al., 2008, 2009). In humans, mutations in ATL1, the dominant form of ATL in the central nervous system, are tightly associated with hereditary spastic paraplegia, a neurodegenerative disease characterized by axon shortening in corticospinal motor neurons and progressive spasticity and weakness of the lower limbs (Zhao et al., 2001; Salinas et al., 2008).

ATL, Sey1p, and RHD3 are anchored in the ER membrane by two closely spaced transmembrane (TM) segments, exposing both termini to the cytosol. The mechanism for homotypic ER fusion has been partially unveiled using ATLs (Hu et al., 2011; Lin et al., 2012). It is very different from SNARE or viral protein-mediated fusion. Recent structural and biochemical studies indicate that the GTPase domain of ATL forms a dimer upon GTP binding, initiating contact between apposing membranes (Bian et al., 2011; Byrnes and Sondermann, 2011). The region following the GTPase domain is a three-helix bundle (3HB) and can dock to its own GTPase domain or that of a neighboring molecule, undergoing major conformational changes during the GTP cycle to drive fusion (Bian et al., 2011;
Membrane tethering has also been shown to require continuous GTP hydrolysis of ATL (Saini et al., 2014; Liu et al., 2015). Fusion is facilitated by a membrane-binding amphipathic helix in the C-terminal tail and the TM segments (Liu et al., 2012; Faust et al., 2015).

Identical membrane topology and conserved signature motifs in the GTPase domains of ATL and Sey1p suggest that Sey1p may behave similarly to ATL during membrane fusion. However, the predicted helical domain between the GTPase domain and TM segments is much longer (≈300 amino acids more) in Sey1p compared with ATL. Furthermore, Sey1p is concentrated in punctae along ER tubules, sometimes at three-way junctions of tubules, whereas ATL1 is mostly distributed evenly on the ER membranes (Hu et al., 2009). Therefore, it is unclear whether Sey1p acts the same way as ATL.

Here, we elucidate how Sey1p mediates homotypic membrane fusion. The proposed mechanism demonstrates a common scheme for fusion mediated by dynamin-like GTPases and reveals that the stalk domain of Sey1p possesses unique functional features.

**Results**

**Crystal structures of the cytosolic domain of Sey1p**

To investigate the mechanism of Sey1p-mediated membrane fusion, we determined the structures of the cytosolic domains (cyt-Sey1p) of Sey1p from *C. albicans* (caSey1p), caSey1p shares 43% sequence identity with Sey1p from *S. cerevisiae* (scSey1p; see Fig. S1 A for sequence alignment). Residues 1–692 (truncation before the first TM region; Fig. 1 A) were expressed in *Escherichia coli*, purified, and crystallized first as an orthorhombic form in the presence of GDP plus AlF$_4^-$ (GDP/AlF$_4^-$). A structure at 2.8-Å resolution was determined by single anomalous diffraction (Table 1), and an electron density map of the nucleotide is shown in Fig. S2 A. Only one caSey1p molecule was present in the asymmetric unit, but a crystallographic dimer occurred along one of the twofold symmetry axes, similar to the hexagonal crystal form of human ATL1 (hsATL1, form 1; Fig. S1 B).

CytSey1p consists of an N-terminal GTPase domain and a long, stalk-like, helical domain connected by a linker region (Fig. 1 B). In the dimer, the GTPase domains interact with one another in such a way that the nucleotide binding sites face each other. The linker regions of the two Sey1p molecules cross one another, allowing a close association of the tops of the stalk domains. The stalk domain is composed of four 3HBs, with the last helix of each bundle extending into the first helix of the next bundle (see Fig. 1 D for topology). The first 3HB (3HB-1) is positioned under the GTPase domain of the pairing protomer. If Sey1p ended here, it would resemble the configuration of the crossover dimer of hsATL1 (form 1; Fig. S1 B).
al., 2011), but the configuration of the Sey1p stalk domain has not been previously identified. The scale of the domain is comparable to that of bacterial dynamin-like protein (BDLP) and is longer than Dynamin-1 or human guanylate-binding protein 1 (GBP1; Fig. S1 C). In BDLP structures, the stalk domain is composed of two four-helix bundles connected by a flexible hinge (Low and Löwe, 2006; Low et al., 2009). Major conformational changes, namely bending around the hinge, have been proposed for BDLP. In Dynamin-1 structures, the stalk consists of the bundle signaling element and one four-helix bundle, and it plays critical role in oligomerization (Faebler et al., 2011; Ford et al., 2011). The stalk of GBP1 starts with two 3HBs arranged similarly as 3HB-1 and -2 in Sey1p, followed by one long helix that reaches back to the bottom of the GTPase domain (Ghosh et al., 2006). Unlike Sey1p, the stalk domains of BDLP, GBP1, and Dynamin-1 interact extensively with their GTPase domains, suggesting that the GTPase–stalk linkage may have different roles in these proteins.

The same crystal form was obtained when GMP-PNP, a nonhydrolyzable analogue of GTP, was added during purification and crystallization. Overall, the Sey1p structure in GMP-PNP is similar to that of GDP/AlF₄⁻, with a root mean square deviation of only 0.3709 Å (Fig. S1 C). In contrast, a different conformation of cytSey1p was observed when the protein was crystallized with GDP (Fig. 1 C). This structure was determined from monoclinic crystals at 2.3-Å resolution using molecular replacement with the GDP/AlF₄⁻ structure as a search model (Table 1). One Sey1p molecule is found in each asymmetric unit, but unlike the first structure, the stalk domain (mainly 3HB-1) is associated with the GTPase domain of the same molecule. In one of the crystallographic dimers, the GTPase domain of another Sey1p molecule comes in close proximity (Fig. S1 E). Such a crystallographic “dimer” would be reminiscent of form 2 of hsATL1 (Fig. S1 B), in which the 3HBs point in opposite directions and the GTPase domains form a dimer interface of 670 Å². However, this Sey1p “dimer” buries only 455 Å² of total surface area, suggesting less substantial contact between the two GTPase domains of the symmetry mates. These results imply that, despite the overall structural similarity between Sey1p and ATL, these proteins likely undergo nucleotide-dependent dimerization differently.

### Nucleotide binding and GTP hydrolysis

The GTPase domain of caSey1p folds in a similar manner as other dynamin-like proteins. It is composed of a central β sheet surrounded by six α helices. The nucleotide at the active site is coordinated mainly by a magnesium ion and four signature motifs: the P-loop (G1, β1–α1), the switch 1 region (G2, α1–β1), the switch 2 region (G3, β3–α2), and the G4 motif (after β5, Fig. S2 B). Most residues involving GTP binding are conserved between Sey1p and ATLs (Fig. S2 B). The two residues of ca-

### Table 1. Data collection and refinement statistics

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Numbers in parentheses are corresponding values for the highest resolution shell. r.m.s.d., root-mean-square deviation; Res., residues.

![](image-url)
Sey1p’s G4 motif (R204 and D205) interact with the ribose of the nucleotide (Fig. S2 C) and are only seen in a subclass of the dynamin family that includes ATLs, Sey1p, RHD3, and GBPs. However, S68 in the P-loop of calSey1p is equivalent to S41 in Dynamin-1 (Chappie et al., 2010), which is required for GTP hydrolysis (Fig. S2, B and D). When compared with dynamin-1, the side chain of S68 does not point directly at the γ-phosphate bond of GTP and does not rotate during the transition state. Furthermore, a bridging sodium ion was found between S41 in dynamin-1 and the nucleotide (Fig. S2 D) but was not seen in Sey1p. In ATLs, an Arg finger (R77 in hsATL1) plays the same role at the position of S68 in calSey1p (Bian et al., 2011; Byrnes et al., 2013). This catalytic residue adopts a different rotamer conformation after the cleavage of the γ-phosphate bond, which in turn causes some minor local rearrangements (Fig. S2 D).

It is known that the GTPase activity of dynamin-1 is stimulated more than 100-fold, when self-assembled on lipid templates. Such dramatic increase was not seen with full-length scSey1p, when its GTPase activity in detergent was compared with that in reconstituted vesicles (Fig. S2 E). As expected and similar to dynamin-1, mutant S47A in scSey1p (equivalent to S68A in calSey1p) was less efficient in hydrolyzing GTP (Fig. S2 F). In addition, the side chain of calSey1p T95 and the backbone nitrogen of T94 and T95 engage the nucleotide. Consistent with the structural observations, mutation of scSey1p T75A (equivalent to T95A in calSey1p) reduces the enzymatic activity of the GTPase (Fig. S2, C and F).

**Dimerization-induced membrane fusion**

Nucleotide-dependent dimer formation is a common feature of dynamin-like GTPases. ATL dimerization is physiologically significant because ATL1 dimer mutants are defective in GTP hydrolysis and membrane fusion, and one “hot spot” of the hsATL1 mutations causing hereditary spastic paraplegia is localized in the GTPase domain near the dimer interface (Bian et al., 2011). Sey1p dimerization has been reported (Anwar et al., 2012) and was confirmed by our structural analysis (Fig. 1 B). Notably, hsATL1 dimers were seen in all structures, regardless of which nucleotide was added (GMP-PNP, GDP/AlF4−, and GDP), whereas calSey1p dimers only formed in the presence of GMP-PNP or GDP/AlF4−.

The dimerization of calSey1p involves a hydrophobic patch on top of the GTPase domain, including a conserved L257 (L233 in scSey1p and L274 in hsATL1) within the guanine cap, and several hydrophobic interactions along the GT-Pase interface (Fig. 2 A). The stalk domains also pack against each other in the dimer. The closest association was in 3HB-2: N407 and N410 from one molecule interact with K467 from the other, and E417 makes a salt bridge with K475 (Fig. 2 B). The bottoms of the stalk domains were also in proximity to each other, and E417 makes a salt bridge with K475 (Fig. 2 B). The same conclusion was drawn with the FRET-based assay (Liu et al., 2015). These results imply that, unlike ATL, Sey1p relies more on GTP binding than hydrolysis to achieve stable dimerization.

To probe the nucleotide requirement for Sey1p-mediated fusion, we used an in vitro lipid-mixing assay. In this assay, purified full-length scSey1p was reconstituted into donor and acceptor vesicles as judged by flotation experiments (Fig. S4 A). The donor vesicles contained lipids labeled with nitrobenzoxadiazole (NBD) and rhodamine at quenching concentrations; fusion with the unlabeled acceptor vesicles leads to fluorophore dilution and dequenching. Wild-type scSey1p resulted in efficient fusion in the presence of GTP and Mg2+. No fusion was seen with GDP or in the absence of Mg2+ (Fig. 2 D). Interestingly, the addition of GMP-PNP caused moderate but reproducible fusion (Fig. 2 D). Such activity was not observed for Drosophila ATL (dmATL) in the presence of GMP-PNP or GDP/AlF4− (Fig. S4 C). Collectively, these results suggest that the linkage of opposing membranes through GTP binding-induced dimerization may be sufficient to promote Sey1p-mediated fusion.

To confirm that GTP binding without hydrolysis can cause fusion, we performed a visual assay of membrane tethering and fusion. Wild-type full-length scSey1p was reconstituted into vesicles containing phospholipids with either a fluorescent Oregon green or Rhodamine dye attached to the head groups. Samples of the reactions were diluted and visualized with a confocal microscope. The vesicles initially appeared as green or red punctae (Fig. 2 F, left). As recently reported for ATL, large yellow punctae (i.e., colocalization of the two dyes) appeared after the addition of GTP, indicating that the differently colored vesicles were associated or fused with one another (Fig. 2 F, middle). When EDTA, which chelates magnesium ions needed for nucleotide binding, was added, some of these large punctae disappeared or became smaller, indicating that some of the vesicles were unethered (Fig. 2 F, right). Very little tethering was seen in the presence of GDP/AlF4− (Fig. 2 H). In the presence of GMP-PNP, a low level of fusion was observed and was not strongly affected by EDTA addition (Fig. 2 G). These results confirmed that the addition of GMP-PNP promotes some Sey1p-mediated fusion.

Next, we tested the effect of dimer mutations on Sey1p activity. Mutation of L233 (equivalent to L257 in caSey1p shown in Fig. 2 A) drastically reduced the dimerization (Fig. S3 C), GTPase activity (Fig. S2 F), and fusion activity of scSey1p (Fig. 2 E and Fig. S4 B for reconstitution controls), whereas mutation of the corresponding L274 in ATL1 resulted in weaker defects in dimerization (Fig. S3 D). Deletion of 3HB-2 to 3HB-4 (residues 378–675) in the stalk domain (Δ3/4 stalk) of scSey1p, which abolishes most of the stalk interactions in the dimer and mimics the size of ATL1 (Fig. S1 C), strongly inhibited fusion activity (Fig. 2 E). scSey1p was also inactive when 3HB-2 was retained but 3HB-3 and -4 (residues 462–675) were truncated (Δ1/2 stalk; Fig. 2 E). Conversely, the removal of a long loop (residues 613–636) within 3HB-4 (ΔLoop) did not significantly affect fusion activity (Fig. 2 E). Nevertheless, all three mutations in the stalk domain had GTPase activity at wild-
Figure 2.  
Nucleotide-dependent dimerization of Sey1p.  
(A) The interface between the two GTPase domains. The cartoon representation is colored as in Fig. 1. Key residues are indicated as sticks. The components of the second protomer (gray) are labeled with a prime symbol. (B) As in A, but for the interface between the two stalk domains. (C) The size of cyt-scSey1p (40 µM, theoretical molecular mass 80.8 kD) or cyt-hsATL1 (40 µM, theoretical molecular mass 51.7 kD) was determined by AUC in the presence of the indicated nucleotides. The estimated molecular masses are given above the peaks in kilodaltons. The data shown are from a single representative experiment out of three repeats. (D) Full-length wild-type (wt) scSey1p was reconstituted at equal concentrations into donor and acceptor vesicles (protein/lipid molar ratio of 1:200). Fusion of donor and acceptor vesicles was followed by the dequenching...
type levels (Fig. S2 G). These results suggest that the length of the stalk domain is important for fusion.

**Conformational change induced by P, release**

The structures suggest that Sey1p retains its configuration during hydrolysis of the γ-phosphate bond but undergoes at least two conformational changes upon P, release: one in the active site and one involving repositioning of the stalk domain relative to the GTPase domain. In the GDP/AlF₄⁻ or GMP-PNP structure, switch 1 and 2 wrap around the nucleotide (Fig. 3 A).

On the other hand, both loops are distant from the active site and partially disordered in the GDP structure (Fig. 3 A). These movements of switch 1 and 2 are typical among GTPases and explain why the dimer is stable with GMP-PNP or GDP/AlF₄⁻ but not with GDP, as the switches are at the dimerization interface (Figs. 2 A and 3 A).

Absence of the γ-phosphate not only breaks the caSey1p dimer but appears to alter the relative positions of the GTPase and stalk domains (Fig. 3 B). When GMP-PNP or GDP/AlF₄⁻ is bound, the linker connecting the GTPase domain and stalk domain attaches to the α₇ of the other protomer. Major contacts include L312 and L318 from one molecule, and V319 and F322 from the other molecule (Fig. 3 C). In addition, Q315 forms a homotypic interaction, and D309 from the linker makes salt bridges to K323 and R376 of the other protomer (Fig. 3 C). In the GDP state, α₇ docks to α₆ of the same Sey1p molecule using mostly the same set of residues as in the GMP-PNP or GDP/AlF₄⁻ state. However, V319 and F322 engage M295 in α₆, Q315 hydrogen bonds with Q303 and R299, and binding occurs between K323 and D291, and R376 and the carbonyl oxygen of A52 (Fig. 3 D).

A cross-linking assay was used to confirm the conformational change inferred from the structures. 3HB-1 of one cyt-Sey1p molecule and α₆ of the other molecule came in proximity to one another in the presence of GMP-PNP or GDP/AlF₄⁻ but not GDP or in the absence of nucleotide. As expected, a cross-linked dimer formed when the scSey1p mutant L297A in scSey1p (as L318A in caSey1p) was treated with diamide to induce disulfide bridge formation (Fig. 3 C). In addition, Q315 forms a homotypic interaction, and D309 from the linker makes salt bridges to K323 and R376 of the other protomer (Fig. 3 C). In the GDP state, α₇ docks to α₆ of the same Sey1p molecule using mostly the same set of residues as in the GMP-PNP or GDP/AlF₄⁻ state. However, V319 and F322 engage M295 in α₆, Q315 hydrogen bonds with Q303 and R299, and binding occurs between K323 and D291, and R376 and the carbonyl oxygen of A52 (Fig. 3 D).

**Analysis of Sey1p in mammalian cells**

To analyze the function of Sey1p, we tested whether it can replace ATLs in mammalian cells. We used siRNA to deplete the major ATL proteins, mainly ATL2 and ATL3 in COS-7 cells, and visualized the ER morphology using calreticulin as a marker. Consistent with previous results (Hu et al., 2009; Wu et al., 2015), when ATLs were knocked down in COS-7 cells (Fig. 4 A), the ER appeared as unbranched morphology in a majority of cells (Fig. 4, B and D), indicating defects in ER fusion. As expected, wild-type human ATL1, but not ATL1 with a mutation in the P-loop (K80A), efficiently restored the ER network in double-depleted cells (Fig. 4, C and D; and Fig. S5 A). When scSey1p was expressed at a comparable level, most cells had normal ER morphology (Fig. 4, C and D; and Fig. S5 A), suggesting that Sey1p and human ATL1 are equally active in maintaining the ER network. K50A (P-loop mutant), L233A (dimer interface mutant), L297A, V298A (both GTPase–stalk interface mutants), and Δ1/2stalk (stalk mutant) all failed to rescue the defects in ER morphology (Fig. 4 D and see Fig. S5 B for expression levels). Mutation M274A largely reduced the percentage of abnormal ER (Fig. 4 D), likely because it only affects the stabilization of the GDP-bound conformation and not Sey1p dimerization. These results confirm that Sey1p plays an important role in the maintenance of ER morphology.

To test whether caSey1p and scSey1p have similar functions in cells, we expressed caSey1p in ATL-depleted COS-7 cells and monitored ER morphology (Fig. 4 C). As expected and similar to scSey1p, wild-type caSey1p but not GTP-binding defective mutant K71A successfully replaced ATLs in mammalian cells (Fig. 4, C and D). To further analyze the function of caSey1p, we expressed caSey1p under the control of the endogenous SEY1 promoter in a CEN vector (Fig. S5 C) in yeast cells lacking Sey1p and Yop1p, an ER tubule-forming protein, and exhibiting abnormal ER morphology. caSey1p was mostly capable of substituting scSey1p in the maintenance of ER morphology (Fig. S5 D). These results confirmed that caSey1p is a functional homologue of scSey1p.

Sey1p protein has been observed to be concentrated in punctae along the ER tubules in yeast cells, mainly at the three-way junctions of the tubules (Hu et al., 2009). The localization of scSey1p in COS-7 cells appears to be similar to that of yeast cells, regardless of whether ATLs are present (Figs. 4 C and 5 A). Punctae were also seen when caSey1p was expressed in ATL-depleted COS-7 cells (Fig. 4 C). Interestingly, scSey1p mutant with defective dimerization and fusion lost its punctate pattern (L233A;
Figure 3. Nucleotide-dependent conformational changes of Sey1p. [A] Cartoon representation of the active site. The GDP/AlF$_4^-$ state is colored in red and the GDP state in gold. The four signature motifs are highlighted. Note that switch 1 and switch 2 in the GDP state are partly disordered. The pairing protomer in the GDP/AlF$_4^-$ state is shown in gray. [B] Superposition of the Sey1p structures in the GDP/AlF$_4^-$ state (red) and the GDP state (gold). The movement of the GTPase domain is highlighted. [C] The GTPase-linker interface in the GDP/AlF$_4^-$ state. The cartoon representation is colored as in Fig. 1. Key residues are shown in sticks. [D] As in C, but for the GTPase–stalk interface in the GDP state. [E] Purified cytosolic domains of scSey1p with cysteines at positions 297 and 298 or at positions 274 and 298 were treated with the oxidant diamide in the presence of the indicated nucleotides. Non–cross-linked protein (single asterisks) and cross-linked dimer (double asterisks) are indicated. Where indicated, the disulfide bridge was reduced with β-mercaptoethanol (BME) before nonreducing SDS/PAGE. [F] As in E, but with wild type (wt) or mutant (cysteines at positions 407 and 467) of the cytosolic domain of ca-Sey1p. [G] Fusion assays of the M274A mutant of scSey1p. Wild-type samples were included for comparison. [H] As in G, but with the mutants of L297A and V298A. The data shown in G and H represents independent experiments with three independently purified batches of protein.
Fig. 5 B). ATL1 mainly distributed evenly along the ER tubules (Fig. 5 C), but ATL3 formed punctae like Sey1p (Fig. 5 D). Given that ATL3 was recently reported to be slower than ATL1 (Hu et al., 2015), GTPases with relatively low hydrolysis activity should tend to form punctae in cells. To test this possibility, we mutated the Arg finger of ATL1 (R77) to Ala. It has been reported that ATL1 R77A can still bind to GTP and form dimers, but GTP hydrolysis occurs much more slowly (Byrnes et al., 2013). We found that ATL1 R77A formed punctae in COS-7 cells (Fig. 5 E). The same results were obtained when these proteins were expressed in ATL-depleted cells (Fig. S5 B). These data suggest that punctae formation is dependent on GTPase activity.

Discussion

Our results suggest how Sey1p causes homotypic fusion of ER membranes (Fig. 6 A). First, two Sey1p molecules in different membranes bind GTP and form a dimer, thereby tethering the membranes together. It is possible that the stalk domains in the dimer do not associate initially but eventually reach the stable side-by-side conformation. To efficiently juxtapose the two membranes, we speculate that the Sey1p dimer does not come in between the membrane but rather sits on the orbit of two closely attached membrane regions with its GTPase domains pointing away from the tethered area. Flattened, closely spaced membranes were previously captured by cryo-EM during membrane tethering mediated by ATL or SNAREs (Diao et al., 2012; Hernandez et al., 2012; Saini et al., 2014). Multiple dimers may self-associate, perhaps through oligomerization of their TM regions and form a ring to stabilize the tethered state, during which time the two membranes begin to fuse. GTP hydrolysis and P release causes a conformational change, probably triggered by changes in the switch 1 region that is part of the GTPase domain dimer interface, and the switch 2 region that propagates at the interaction surface between the GTPase domain of one Sey1p molecule and the linker and stalk domain of the other. The GTPase domain would then be released and move freely until docking to its own stalk domain, resulting in disruption of the Sey1p dimer. Regardless of whether the membrane merging occurs before the monomerization of Sey1p, these molecules would start a new fusion attempt by searching for a dimer partner on opposing membranes.

Surprisingly, we observed membrane fusion by Sey1p even with GMP-PNP, indicating that fusion can occur without GTP hydrolysis. However, most of the Sey1p dimers formed with GMP-PNP would be on the same membrane, i.e., cis-dimer, and are futile. Fusion is much more efficient when GTP is added, as GTP hydrolysis would allow Sey1p cycling between dimer and monomer states. Consequently, more Sey1p monomers would be available for trans-interactions between different membranes. Such competition between cis- and trans-interactions has also been recently reported for ATL (Liu et al., 2015).

Sey1p- and ATL-mediated fusion differs in that the power stroke occurs at different steps of the GTP cycle. Stable trans-dimers of ATL (in form 3) are in a crossover configuration and would not be possible without GTP hydrolysis (Fig. 6 B). To achieve the tethered state, ATL molecules from apposing membranes have to bind GTP and dimerize into form 2 first; the two GTPase domains can then rotate and recapture the 3HBs from the pairing molecule to adopt form 3. The GTP-bound ATL dimer may switch between the two conformations until GTP hydrolysis drives the equilibrium toward form 3. Consistent
**Figure 5.** **Sey1p punctae formation.** (A) Myc-scSey1p was expressed in COS-7 cells and its localization determined by anti-Myc antibodies and compared with that of calreticulin using indirect immunofluorescence and confocal microscopy. The right image shows an enlargement of the boxed region centered on the tubular ER network. Bars, 10 µm. (B) As in A, but with cells expressing Myc-scSey1p L233A. (C) As in A, but with cells expressing Myc-hsATL1. (D) As in A, but with cells expressing Myc-hsATL3. (E) As in A, but with cells expressing Myc-hsATL1 R77A. The arrowheads indicate the puncta localized at the three-way junctions of the ER network.

**Figure 6.** **Comparison of homotypic membrane fusion mediated by Sey1p and ATL.** (A) A model of Sey1p-mediated fusion. See Discussion for details. GTP and GDP molecules are indicated as cyan and blue spheres, respectively. (B) As in A, but for ATL. TMs, transmembrane segments; CT, cytosolic tail.
with these models, ATL dimer is most stable when in complex with GDP/$\text{AlF}_4^-$, whereas Sey1p dimer is strongest with GMP-PNP. Such a difference would not be explained by the structures but is likely determined by residues that sense the subtle change in distance between $\beta$- and $\gamma$-phosphate.

The formation of punctae at the three-way junctions or along ER tubules is seen in some, but not all, GTPase-based ER fusogens. Our results suggest that the kinetics of GTP hydrolysis play a role in punctae formation. Slow GTPases, such as Sey1p and ATL3, are prone to clustering. However, mutants with further reduced activity, which do not sustain fusion, will lose the punctate pattern and become evenly distributed on the ER membrane. One scenario would be that effective fusion requires the simultaneous participation of several GTPase molecules. Alternatively, slow hydrolysis would prolong the existence of a tethered complex, allowing the observation of punctae.

The proposed mechanism reveals that homotypic membrane fusion could be achieved by different approaches in Sey1p and ATL but with a similar paradigm. Some of the mechanism could apply to RHD3 and its homologues in the fusion of plant ER membranes, and to mitofusin/Fzo1p (Hermann et al., 1998; Chen et al., 2003), another member of the dynamin family, in the fusion of mitochondrial outer membranes. Three point mutations were previously identified in RHD3 (Wang et al., 1997). Based on sequence alignment with caSey1p, I134T and D185N would reside near the GTPase dimer interface, and A575V would be placed in the hydrophobic core of 3HB-4 (Fig. S1 D). The localization of these mutations confirms that dimerization of the GTPase domain and integrity of the stalk domain is critical for these dynamin-like proteins.

Materials and methods

Constructs, strains, and antibodies

The N-terminal cytosolic domains of \textit{C. albicans} Sey1p (residues 1–692), codon-optimized \textit{S. cerevisiae} Sey1p (residue 1–680), or hsATL1 (residues 18–447) were inserted into the NheI–Xhol sites of pET-28a (EMD Millipore), which includes a thrombin-cleavable, N-terminal HA tag and 300 bp upstream and downstream sequences of scSey1p and full-length dual-amyase were cloned into the BamHI–Xhol sites of pGEX6p-1 vector with a GST tag followed by a PreScission protease digestion site at the N terminus. For fusion assay, codon-optimized full-length scSey1p and full-length dual-amyase were cloned into the BamHI–Xhol sites of pGEX6p-1 vector with a GST tag followed by a PreScission protease digestion site at the N terminus. For mammalian expression, Myc-scSey1p, Myc-caSey1p, or Myc-hsATL1 was PCR-amplified and cloned into pcDNA4/TO (Invitrogen) by KpnI–BamHI, BamHI–Xhol, and KpnI–XhoI sites. Myc-hsATL3 was PCR amplified and cloned into pGW1. For endogenous level expression of scSey1p and caSey1p in yeast cells, the coding region of scSey1p and caSey1p plus an N-terminal HA tag and 300 bp upstream and downstream sequences of scSey1 were amplified and inserted into the KpnI–NotI site of pRS316 (a URA/CEN plasmid). The plasmid pK59, encoding SeC6-GFP was used to visualize ER morphology. All point mutations were generated using the QuikChange Site-Directed Mutagenesis kit (Agilent Technologies). The truncations (Δ3/4 stalk, deletion of residues 378–675; Δ1/2 stalk, deletion of 462–675; ΔLloop, deletion of 613–636) were generated using overlap PCR primers. All constructs were confirmed by DNA sequencing.

The strain JHY4 (BY4741 \textit{sey1Δ::kanMX4 yop1Δ::HIS3MX6; Hu et al., 2009}) was used to visualize ER morphology. Mouse anti-Myc and rabbit anti-Calreticulin antibodies were purchased from Abcam, and mouse anti-HA and mouse anti-phosphoglycerate kinase antibodies were purchased from Sigma-Aldrich.

Protein expression and purification

The cyt-caSey1p, cyt-scSey1p, and cyt-hsATL1 constructs were transformed into \textit{Escherichia coli} strain BL21 (DE3), and cultures were grown in Luria-Bertani media at 37°C to an OD600 of 0.6. Protein expression was induced by the addition of 0.4 mM IPTG for 20 h at 16°C. Cells were harvested, resuspended in lysis buffer (20 mM Tris, pH 8.0, 150 mM NaCl, 4 mM MgCl$_2$, and 20 mM imidazole), and lysed by an ultra-high pressure cell disrupter (JN-3000 PLUS; JNBI0). The lysate was centrifuged at 16,000 rpm for 1 h, and the supernatant was collected. The protein was isolated by Ni-NTA chromatography (GE Healthcare) and further purified by ion-exchange chromatography (Resource-Q or HiTrap Q; GE Healthcare) and gel filtration chromatography (Superdex-200; GE Healthcare). For crystallization, corresponding nucleotides at 2 mM were included in the gel filtration buffer (Yan et al., 2011; Chen et al., 2012).

For expression of selenomethionine-substituted (Se-Met) Sey1p, cells were grown in M9 medium at 37°C to an OD600 of 0.6. Solid amino acid supplements (50 mg/liter Lys, Thr, and Phe; 25 mg/liter Leu, Ile, and Val; 30 mg/liter Se-Met) were added to the medium before induction. The Se-Met protein was purified by the same procedure as the native protein.

Full-length dual-amyase or scSey1p were expressed and purified with a GST tag. The proteins were solubilized and isolated in Triton X-100 (Anatrace), and the GST tag was cleaved before reconstitution (Bian et al., 2011; Anwar et al., 2012).

X-ray crystallography

All native and selenomethionine-substituted proteins were concentrated to 8 mg/ml and crystallized by vapor diffusion methods at 16°C. Crystals of Sey1p with GDP/$\text{AlF}_4^-$ were grown within days in 160 mM lithium sulfate monohydrate, 80 mM Tris, pH 8.5, 20% (wt/vol) polyethylene glycol (PEG) 3350, 40 mM ammonium acetate, 20 mM sodium citrate tribasic dehydrate, pH 5.6, 6% (wt/vol) PEG 4000, and 20 mM spermidine. Crystals of Sey1p with GMP-PNP appeared around 6 d in 120 mM DL-malic acid, pH 7.0, 16% (wt/vol) PEG 3350, 20 mM bicine, pH 9.0, 400 mM magnesium chloride hexahydrate, and 6% D-(+)-trehalose dihydrate. Crystals of Sey1p with GDP complex appeared around 2 mo in 20 mM Tris, pH 8.5, 160 mM potassium sodium tartrate tetrahydrate, 4 mM GDP, and 21% (wt/vol) PEG 3350.

All crystals were transferred to a cryo-protectant solution (4 M sodium formate) before being flash frozen in liquid nitrogen for storage. The native data for Sey1p with GDP was collected in a beamline BL17U at the Shanghai Synchrotron Radiation Facility (China). Other data were collected at 100 K using a charge-coupled device detector (Q270; Area Detector Systems Corporation) on beamline BL1A at Photon Factory (Japan). All data were processed and scaled using HKL2000 package 24 (Otwinowski and Minor, 1997).

The structure of Sey1p with GDP/$\text{AlF}_4^-$ was solved by single anomalous diffraction of Se-Met using SHELX (Schneider and Sheldrick, 2002). An initial model was built by the program coot and used in molecular replacement to determine the structure of Sey1p with GDP or GMP-PNP (Emsley and Cowtan, 2004). For model refinement, the Phenix refinement program (phenix.refine) was used (Afonine et al., 2012). Interface and surface area were calculated using PDBePISA.

AUC

Purified cyt-scSey1p and cyt-hsATL1 were used for AUC in a buffer containing 50 mM Tris, pH 8.0, 300 mM NaCl, and 5 mM MgCl$_2$. Sedimentation velocity experiments were performed at 4°C in a proteomeLab XL-1 Protein Characterization System (Beckman Coulter). 2 mM GDP, 2 mM GMP-PNP, or 2 mM GDP/6 mM AlCl$_3$/30 mM NaF.
were added to 40 μM protein before centrifugation. All interference data were collected at a speed of 42,000 rpm in a rotor (An-60 Ti; Beckman Coulter) and analyzed by the program SEDFIT in terms of a continuous c(s) distribution. In sedimentation equilibrium experiments, the proteins were prepared at three different concentrations (0.2, 0.3, and 0.5 mg/ml). Interference data were collected at three different speeds (8,000, 12,000, and 15,000 rpm for cytATL1; 6,400, 9,600, and 12,000 rpm for cytSey1p) in a An-60 Ti rotor at 4°C and analyzed by SEDPHAT using a model of monomer–dimer self-association.

Cross-linking assays

1 μM wild-type or mutant cytSey1p protein in 50 mM Hepes, pH 7.4, 100 mM NaCl, and 5 mM MgCl₂, with or without 1 mM GDP, 1 mM GMP-PNP, or 1 mM GDP/3 mM AlCl₃/15 mM NaF was incubated with 10 μM diadime at room temperature for 30 min 500 μM N-ethylmaleimide was used to quench the reactions. The samples were analyzed by SDS-PAGE and Coomassie blue staining.

Fusion assays

Lipid-mixing assays were performed as previously described (Bian et al., 2011). In brief, lipids were dried down to a film, hydrated with A100 buffer (25 mM Hepes, pH 7.4, 100 mM KCl, 1 mM 2-mercaptoethanol, and 10% glycerol), and extruded through polycarbonate filters with a pore size of 100 nm. Proteins were reconstituted at a protein/lipid molar ratio of 1:200, except dmATL, which was reconstituted at a ratio of 1:1,000 (Fig. S4 C). The flotation of proteoliposomes was done on a sucrose gradient to determine the reconstruction efficiency. 20 μM proteoliposomes was mixed with 100 μl of 1.9 M sucrose and overlaid with 100 μl of 1.25 M sucrose and 30 μl of 0.25 M sucrose (all sucrose solutions were in 25 mM Hepes, pH 7.4, and 100 mM KCl). After centrifugation at 174,000 g for 75 min and 4°C in a rotor (TLS-55; Beckman Coulter), the gradient was fractionated into five fractions and analyzed by SDS-PAGE and Coomassie blue staining. Donor and acceptor liposomes were added at a 1:3 ratio. The initial NBD fluorescence was set to zero, and the maximum fluorescence was determined based on a standard curve.

Visual tethering/fusion assays

Full-length wild-type scSey1p was reconstituted into perovform liposomes with Rhodamine-DPPE (1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine) or Oregon green-DPPE (Invitrogen) in the same way as described in the methods of Fusion assays. The vesicles were mixed 1:1 in a reaction buffer containing 5 mM MgCl₂. Different nucleotides were added and the samples diluted 1:100 in reaction buffer or buffer with 10 mM EDTA. 5 μl samples were spotted onto a glass coverslip and visualized by confocal microscopy. Oregon green and Rhodamine dyes were excited with 488- and 561-nm lasers (100 mW), and their emissions selected with 530/40- and 620/50-nm filters, respectively. All images were captured under room temperature on a Zeiss LSM 710 using ZEN2012 software (Carl Zeiss). Changes were made linearly across the entire image using Photoshop (Adobe).

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Mammalian cell culture, transfections, and microscopy

COS-7 cells were maintained at 37°C with 5% CO₂ in DMEM containing 10% fetal bovine serum and passaged every 2–3 d. For siRNA transfections, COS-7 cells were plated at 50% confluency and transfected the next morning with 13 nM each siRNA oligonucleotide duplex using RNAiMAX (Invitrogen) for 6 h. Cells were then scraped for immunoblot analysis or fixed for immunofluorescence analysis 48 h after transfection. Specific siRNA oligonucleotides (Invitrogen) for ATL2 were targeted against the following sequences: 5′-GGAGCUAUCCCUAUGAACUUCAUA-3′; and that for ATL3 were 5′-GGGUAGAGAUGGAGUUCUCCUUUAU-3′. The control siRNA oligonucleotide was obtained from Invitrogen. The siRNA-resistant constructs were transfected into knockdown cells using Lipofectamine 3000 (Invitrogen) after 24 h of siRNA transfection.

Transfected cells were fixed for 20 min at room temperature with 4% formaldehyde (Bio Basic, Inc.) and permeabilized with 0.1% Triton X-100 (Bio Basic, Inc.). The cells were washed twice with PBS and probed with primary antibodies, including mouse anti-Myc antibody (Abcam) diluted 1:500 or rabbit anti-calreticulin antibody (Abcam) diluted 1:500, for 60 min in PBS containing 3% calf serum, followed by incubation with various fluorophore-conjugated secondary antibodies for an additional 45 min (Alexa Fluor 488–conjugated anti–rabbit or Alexa Fluor 594–conjugated anti–mouse (1:800 dilution; Invitrogen). For ER morphology rescue assay, yeast cells were imaged using Sec63p-GFP as a marker (Wu et al., 2015). All images were captured under room temperature on a confocal microscope (TCS SP5; Leica) with a 63×/1.40 numerical aperture Plan Apochromat oil immersion objective lens using LAS AF version 1.3.1 build 525 software (Leica).

Accession numbers

The coordinates and structure factors have been deposited in the Protein Data Bank under accession nos. 5CA8 (GDP bound state), 5CA9 (GDP/AIF4– bound state), and 5CB2 (GMP-PNP bound state).

Online supplemental material

Fig. S1 shows comparison of Sey1p and other dynamin-like proteins. Fig. S2 shows characterization of the GT-Pase domain of Sey1p. Fig. S3 shows AUC analysis of Sey1p and ATL1. Fig. S4 shows controls for fusion assay of Sey1p. Fig. S5 shows additional analysis of Sey1p in cells. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201502078/DC1.
Figure S1. Comparison of Sey1p and other dynamin-like proteins. (A) C. albicans Sey1p (caSey1p) and S. cerevisiae Sey1p (scSey1p) were aligned. Predicted secondary structural elements, with α helices shown in bars and β strands given in arrows, and GTPase motifs of scSey1p are highlighted. Residues mutated in this paper are indicated by dots. TM, transmembrane domain; CTH, predicted C-terminal amphipathic helix. Residues important for hydrolysis and dimerization are indicated by black dots. (B) Human ATL1 dimers observed in crystal form 1 (PDB code: 3QNU, middle, GDP bound), form 2 (PDB code: 3QOF, top, GDP bound in the presence of inorganic phosphate), and form 3 (PDB code: 4IDP, bottom, GMP-PNP bound or GDP/AlF₄⁻ bound) are shown. The protomers in the dimer are shown in lime color and slate cartoon representation. Nucleotides are shown as orange sticks, and magnesium ions are shown as yellow spheres. (C) Structures of caSey1p in complexes with GMP-PNP or GDP/AlF₄⁻ are compared with other dynamin-like proteins, including GDP-bound BDLP (PDB code: 2W6D), nucleotide-free dynamin-1 (PDB code: 3ZVR), GMP-PNP–bound GBP1 (PDB code: 1F5N), and GDP-bound ATL1 (PDB code: 3QOF). All structures of dynamin superfAMILY members are shown in rainbow-colored cartoon representation. (D) Mutations of RHD3 are mapped into caSey1p dimer structure (blue spheres). (E) Crystallographic dimer of GDP-bound caSey1p. GDP is shown as orange sticks, and magnesium ions are shown as yellow spheres.
Figure S2. The GTPase of Sey1p. (A) The 2Fo – Fc electron density maps (1.6σ contour) are shown as wire mesh (blue) with the refined atomic model of the nucleotides superimposed. Magnesium ions are shown as green spheres. (B) Sequence alignment of the conserved GTPase motifs. Key residues are highlighted and numbered. (C) The catalytic center of caSey1p with GDP/AlF$_4^-$ bound is shown with key side chains in cyan sticks, GDP is given in orange, and AlF$_4^-$ is shown in gray and light blue. (D) Comparison of the catalytic centers in caSey1p, hsATL1, and dynamin-1. GMPPNP or GMP-PCP states are shown in gray, and GDP/AlF$_4^-$ is in cyan. The catalytic residue is shown in sticks. (E) GTPase activity of wild-type scSey1p. Left graph shows cytSey1p at different concentrations, and the right one shows full-length scSey1p (0.3 µM) in either proteoliposomes or detergents Triton X-100 (TX-100). (F) GTPase activity of Sey1p proteins. The activities were measured by phosphate release at saturating GTP concentrations (0.5 mM) using the cytosolic domains (1 µM). Each bar is the mean and SE of four measurements. All measurements are representative of at least three repetitions. wt, wild type.
Figure S3. **AUC analysis of Sey1p and ATL1.** (A) AUC analysis of wild-type scSey1p was performed at three concentrations and three different speeds in the presence of two different nucleotides as indicated. The absorbance profiles at sedimentation equilibrium were recorded and used to calculate the dissociation constant (highlighted in green boxes) of the dimers. The corresponding residuals for the fit are shown in the lower portion of each graph. (B) As in A, but with wild-type hsATL1. Data shown in A and B are representative of three repeats. (C) The size of cyt-scSey1p L233A (40 µM, theoretical molecular mass 80.8 kD) was determined by AUC in the presence of the indicated nucleotides. The estimated molecular masses are given above the peaks in kilodaltons. (D) As in C, but with cyt-hsATL1 L274A (40 µM, theoretical molecular mass 51.7 kD). All measurements shown in C and D are representative of three repetitions.
Figure S4. Controls for fusion assay of Sey1p. (A) Proteoliposomes containing wild-type (wt) full-length scSey1p were floated in a sucrose gradient and the fractions analyzed by SDS/PAGE and Coomassie staining. (B) Full-length wild-type scSey1p or the indicated mutants were purified and reconstituted into donor (D) and acceptor (A) vesicles. Aliquots of the proteoliposomes were floated in a discontinuous sucrose gradient and analyzed by SDS/PAGE and Coomassie blue staining. The same vesicles were used for the fusion assay in Figs. 2 (D and E) and 3 (G and H). (C) Fusion assay of the wild-type Drosophila ATL with indicated nucleotides. The data shown are representative of at least three repetitions.
Figure S5. Analysis of Sey1p in cells. (A) COS-7 cells used in Fig. 4 were double depleted for hsATL2 and hsATL3 and then transfected with Myc-hsATL1, Myc-scSey1p, or Myc-caSey1p. The levels of Myc-tagged proteins were determined by immunoblotting. Tubulin was used as a loading control. (B) Myc-tagged hsATL1 or scSey1p were expressed in COS-7 cells depleted of ATLs. Localization was determined by anti-Myc antibodies and compared with calreticulin using indirect immunofluorescence and confocal microscopy. Bar, 10 µm. (C) Yeast cells lacking Sey1p and Yop1p were transformed with CEN plasmids expressing HA-tagged scSey1p and caSey1p under the endogenous promoter. The levels of scSey1p and caSey1p were determined by immunoblotting. Phosphoglycerate kinase was used as a loading control. (D) The ER morphology of the cells used in C was visualized by confocal microscopy using a GFP fusion of the ER protein Sec63p, with the microscope focused either at the center or the periphery of the cell. Bar, 2 µm. The percentage of cells with abnormal ER (loss of reticular network in the cortex as shown on the right) was determined from ≥200 cells per sample. IB, immunoblot.