Coassembly of Mgm1 isoforms requires cardiolipin and mediates mitochondrial inner membrane fusion

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Two dynamin-related protein (DRP) families are essential for fusion of the outer and inner mitochondrial membranes, Fzo1 (yeast)/Mfn1/Mfn2 (mammals) and Mgm1 (yeast)/Opa1 (mammals), respectively. Fzo1/Mfn1s possess two medial transmembrane domains, which place their critical GTPase and coiled-coil domains in the cytosol. In contrast, Mgm1/Opa1 are present in cells as long (l) isoforms that are anchored via the N terminus to the inner membrane, and short (s) isoforms were predicted to be soluble in the intermembrane space. We addressed the roles of Mgm1 isoforms and how DRPs function in membrane fusion. Our analysis indicates that in the absence of a membrane, l- and s-Mgm1 both exist as inactive GTPase monomers, but that together in trans they form a functional dimer in a cardiolipin-dependent manner that is the building block for higher-order assemblies.

Introduction

Mitochondrial fusion is a conserved process whose fundamental function is likely to create a more connected compartment that facilitates content exchange and access to mitochondrial DNA (Hoppins et al., 2007). Two dynamin-related protein (DRP) families are essential for fusion: Fzo1 (yeast)/Mfn1/Mfn2 (mammals) and Mgm1 (yeast)/Opa1 (mammals), which drive outer and inner mitochondrial membrane fusion, respectively (Meeusen et al., 2004, 2006). Outer and inner membrane tethering is mediated by the self assembly of mitochondrial fusion DRPs via intermolecular interactions (Ishihara et al., 2004; Koshiba et al., 2004; Meeusen et al., 2004, 2006; Griffin and Chan, 2006). Analysis of mutant alleles of the fusion DRPs indicates that membrane tethering is separable from subsequent lipid content mixing and that fusion DRPs are essential at each stage (Meeusen et al., 2006).

The localization and topologies of the mitochondrial outer and inner membrane DRPs are distinct. Fzo1/Mfn1/2 possess two medial transmembrane domains that target and anchor them in the mitochondrial outer membrane and place the critical GTPase and coiled-coil regions in the cytosol, with a short loop in the intermembrane space (Hermann et al., 1998; Rapaport et al., 1998). Mgm1/Opa1 are targeted to the mitochondrial inner membrane via an N-terminal stop-transfer signal, placing the GTPase domain proximal to the membrane (Herlan et al., 2003).

Two isoforms of Mgm1/Opa1 are generated during their biosynthesis by divergent proteolytic mechanisms: long (l) isoforms are anchored via the N terminus to the inner membrane, and short (s) isoforms are predicted to be soluble in the intermembrane space. We addressed the roles of Mgm1 isoforms and how DRPs function in membrane fusion. Our analysis indicates that in the absence of a membrane, l- and s-Mgm1 both exist as inactive GTPase monomers, but that together in trans they form a functional dimer in a cardiolipin-dependent manner that is the building block for higher-order assemblies.

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Figure 1. s-Mgm1 assembly is regulated by CL. (A) Purified l- and s-Mgm1 (40 pmol of each) analyzed by SDS-PAGE. Schematic representations of Mgm1 isoforms are shown (right). (B) Hydrodynamic analysis of l- and s-Mgm1. (C) s-Mgm1 preferentially associates with IMC liposomes. 0.5 µM s-Mgm1 was incubated with OMC 6% CL, IMC 0% CL, or IMC 20% CL liposomes and analyzed by floatation in sucrose gradients. A representative SDS-PAGE and Western analysis of float (F) and pellet (P) fractions is shown. Quantification from three experiments is shown as the mean + SEM (error bars).
Results and discussion

l- and s-Mgm1 isoforms exist as inactive monomers

To understand their roles in fusion, we expressed and purified s- and l-Mgm1 and characterized their kinetic and structural properties (Fig. 1 A). l-Mgm1 uniquely required detergent and glycerol to maintain its solubility (1.5% vs. critical micelle concentration of 1.5–2.0%). We examined the ability of s- and l-Mgm1 to hydrolyze GTP, which for DRPs depends on self-assembly (Dunino and Hinshaw, 2001). We observed no detectable GTPase activity for s- or l-Mgm1 over a range of protein and GTP concentrations (unpublished data; s-Mgm1 0.03 min⁻¹; l-Mgm1 0.00 min⁻¹). Hydrodynamic analysis of s- or l-Mgm1 by sucrose gradient centrifugation and gel filtration chromatography revealed that both exist as monomers (Fig. 1 B). In contrast to Mgm1, the membrane division DRPs—Dnm1 and dynamin—exist as stable dimers, which in their unassembled form possess a basal rate of GTP hydrolysis (Ingerman et al., 2005; Ramachandran et al., 2007).

s-Mgm1 preferentially associates with liposomes containing the inner membrane-enriched anionic lipid cardiolipin (CL), which stimulates self assembly-driven GTP hydrolysis

The significance of monomeric species of Mgm1 was revealed when we examined how s-Mgm1 interacts with liposomes, as assessed by flotation using equilibrium sucrose gradient centrifugation. This analysis indicated that s-Mgm1 preferentially associates with liposomes whose composition mimics the inner mitochondrial membrane composition (IMC) as compared with liposomes with an outer membrane composition (Fig. 1 C, OMC 6% CL vs. IMC 20% CL, composition described in Materials and methods). These data indicate that the critical lipid species required for s-Mgm1 association with IMC liposomes was the inner membrane-enriched dianionic phospholipid, CL (Fig. 1 C, IMC 20% CL vs. IMC 0% CL; and not depicted).

Association of s-Mgm1 with CL-containing IMC liposomes stimulated GTP hydrolysis in a CL-dependent manner, to a maximal velocity of 1.3 min⁻¹ (Fig. 1 D, compare IMC 20% CL vs. IMC 0% CL). When compared with other DRPs this is relatively slow, but it is similar to recently published kinetic data for assembled s-Mgm1 (Meglei and McQuibban, 2009). Although CL is enriched in the inner membrane, biochemical analysis of mitochondria indicates that it may also comprise up to 6% of the outer membrane (Sperta-Gottlieb et al., 1988; Daum and Vance, 1997). In the presence of OMC liposomes containing up to 6% CL, s-Mgm1 was significantly less active as compared with s-Mgm1 in the presence of IMC 20% CL liposomes, which is consistent with its role in inner membrane fusion (Fig. 1 D; Meeusen et al., 2006).

These observations suggest that CL-containing liposomes stimulate self-assembly of s-Mgm1. To test this possibility, we chemically cross-linked s-Mgm1 in the presence and absence of CL-containing IMC membranes with bis(sulfosuccinimidyl)suberate (Fig. 1 E, BS3). Analysis of cross-linked products revealed that s-Mgm1 dimers and tetramers form in a CL-specific manner (Fig. 1 E). These data indicate that dimeric s-Mgm1 is the building block for the assembly of larger structures, which is consistent with previous cross-linking analysis of Mgm1 and Opa1, and further suggests that the formation of a DRP dimer interface is required for GTP hydrolysis and is a general property of the DRP family (Frezza et al., 2006; Gasper et al., 2009; Meglei and McQuibban, 2009).

Thus, in contrast to membrane division DRPs, our data indicate that the s-Mgm1 monomer-to-dimer assembly step is regulated by CL. CL-saturated detergent micelles did not activate s-Mgm1 GTP hydrolysis, indicating that CL stimulates s-Mgm1 self assembly only in the context of a lipid bilayer, which may serve as a 2D platform for the stimulation of s-Mgm1 assembly (unpublished data). This CL membrane regulatory step has likely been harnessed in vivo to couple inner membrane targeting of Mgm1 to its assembly and activation, which is critical given that s-Mgm1 has access to both mitochondrial membranes.

Short Mgm1 assembles into a parallel dimer that further assembles into a novel DRP structure

To gain insight into organization of assembled s-Mgm1 structures, we examined IMC liposomes containing s-Mgm1 by negative-stain EM. This analysis revealed that in the absence of nucleotide, s-Mgm1 associates with CL-liposomes and self-assembles into an extended, organized lattice (Fig. 1 F). The structural features of this lattice are striking in their novelty for a DRP. Division DRPs form curved filaments or helical structures in the absence or presence of GTP, respectively (Ingerman et al., 2005). In the presence of GTP or nonhydrolyzable GTP analogues, s-Mgm1 remained associated with IMC liposomes, but s-Mgm1 lattices were not observed, indicating that GTP binding and hydrolysis facilitates conformational changes (Fig. 1 F and not depicted).

The uniform organization of the s-Mgm1 lattice prompted us to attempt 2D crystallization of s-Mgm1 on CL-containing monolayers (Fig. 2). Crystals of ~1 μm were obtained and imaged as negative-stained preparations in the transmission EM (Fig. 2 A). The calculated power spectrum of recorded 2D crystal images showed diffraction spots with up to 3 nm resolution before image unbending. Comparison of phase residuals with the ALLSPACE program revealed a P3 symmetry. The unit cell parameters were a = b = 200 Å, γ = 120°. A total of eight images...
higher-order structures (Fig. 1 E). A contour level was chosen to include the volume of the s-Mgm1 dimer of 2 × 84 kD, assuming a density of 0.82 D/Å³ (Fig. 2 C).

The GTPase domain of Mgm1 has a high sequence homology to both dynamin A and dynamin 1 (from Dictyostelium discoideum and Rattus norvegicus, respectively) and the remainder of Mgm1 exhibits weak homology to Nostoc punctiforme bacterial dynamin-like protein, whose structure has been determined by x-ray crystallography (Low and Löwe, 2006). To obtain an atomic model for s-Mgm1, we used a combination of homology and threading modeling. The resulting model contains higher-order structures (Fig. 1 E). A contour level was chosen to include the volume of the s-Mgm1 dimer of 2 × 84 kD, assuming a density of 0.82 D/Å³ (Fig. 2 C).

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from the nontilted images were merged in 2dx_merge, yielding a projection map at 3.1 nm resolution (overall phase residual: 29.4° [IQ weighted]; Fig. 2 B). Six images of 30° tilted and nine images of 45° tilted negatively stained s-Mgm1 2D crystals were collected and merged with the nontilted data, yielding a 3D reconstruction at 3.1 nm resolution, applying P3 symmetry (Fig. 2 C). This 3D reconstruction showed a trimer of densities that clearly separated into dimers at higher contouring thresholds that were assembled in a hexameric ringlike structure. This structural model is consistent with our cross-linking data, which indicates that a dimer is the building block for self assembly of.

Figure 2. Structural analysis of s-Mgm1. (A) EM analysis of a negatively stained 2D crystal of s-Mgm1. The power spectrum calculated from the raw image is shown in the inset and shows strong diffraction up to 3 nm resolution. (B) P3 symmetrized projection map of s-Mgm1, calculated from merged data from nine processed images of negatively stained s-Mgm1 2D crystals. 2 × 2 unit cells are shown. One unit cell has dimensions of a = b = 200 Å, γ = 120°. (C) 3D reconstruction from images of tilted negatively stained 2D crystals of s-Mgm1 as seen from the direction perpendicular to the membrane plane. 2 × 2 unit cells are shown. (D) Homology model for s-Mgm1. This model was created using Modeller (Martí-Renom et al., 2000) and is based on the structures of dynamin A, dynamin-1, and a bacterial dynamin-like protein (PDB accession nos. 1JWY, 2AKA, and 2J69). The monomer is depicted as seen from the dimer interface. The model is color coded as follows: the GTPase domains is yellow, the GED is blue, and the pair of helices that putatively bind the membrane are orange. GDP-Mg complexes are depicted as spheres. (E) Schematic representation of the proposed parallel s-Mgm1 dimer bound to a lipid bilayer. The homology-modeled dimer and the corresponding 3D reconstruction based on the 2D crystallographic data are shown at the same scale. One monomer is colored as in D. Bars: (A) 200 nm; (C) 10 nm; (E) 5 nm.
682 residues spanning the GTPase domain and part of the C-terminal coiled-coil domain (from residue 145 to 829) in which the N and C termini are colocated in an antiparallel helix bundle (Fig. 2 D). The homology model was docked into the 3D reconstruction, as described in Materials and methods. The available data are most consistent with a parallel dimeric arrangement of two s-Mgm1 models (Fig. 2 E).

The propensity of s-Mgm1 to readily self-assemble into an ordered 2D lattice indicates that it possesses distinct interfaces that mediate higher-order self-assembly interactions: a monomer–monomer interface for dimer formation and a dimer–dimer interface for the assembly of a higher-order hexameric s-Mgm1 ring. Our structural model of assembled s-Mgm1 also predicts that an open interface exists, which contains the helical region that is likely orthologous to HR2 in Mfn1, proposed to mediate trans interactions responsible for membrane tethering (Koshiba et al., 2004).

\section*{l-Mgm1 preferentially associates with and reconstitutes into liposomes containing \textit{CL}, but cannot hydrolyze GTP}
Similar to s-Mgm1, l-Mgm1 was preferentially inserted into the bilayer of IMC liposomes containing CL (Fig. 3 A, IMC 20\% CL vs. IMC 0\% CL). Stable insertion of l-Mgm1 into IMC liposomes was assessed by treatment with high salt followed by OptiPrep gradient centrifugation, which indicated that the efficiency of l-Mgm1 membrane insertion was \(\sim 74 \pm 1.9\%\), \(n = 3\) (Fig. 3 A, IMC 20\% CL, 0.5 M NaCl). Protease protection analysis of reconstituted l-Mgm1 indicated that the membrane-inserted form of l-Mgm1 is in its native topology, with its GTPase and coiled-coil regions facing outwards (Fig. 3 B).

In contrast to s-Mgm1, the GTPase activity of inserted l-Mgm1 in IMC liposomes was undetectable (Fig. 3 C, INS). However, l-Mgm1 GTPase activity was observed upon treatment of inserted l-Mgm1 IMC liposomes with concentrations of the detergent MEGA-8 that saturated the IMC membranes, and under noninsertion conditions produced by the addition of MEGA-8–solubilized l-Mgm1 to IMC liposomes (Fig. 3 C, INS and NI, 0.5\% MEGA-8, respectively). These observations indicate that insertion of l-Mgm1, placing its GTPase domain proximal to the membrane, likely constrains and attenuates its ability to hydrolyze GTP (Fig. 1 A). The topology of l-Mgm1 and its effects on GTPase activity are also likely responsible for the coevolution of divergent proteolytic pathways in yeast and mammalian cells that function to create the active GTPase, s-Mgm1. These data further suggest that a heterotypic s-Mgm1/l-Mgm1 dimer is the functional unit for fusion, where l-Mgm1 uniquely contributes a transmembrane region required for accurate inner membrane targeting and other fusion activities, whereas s-Mgm1 contributes an active GTPase domain.

\section*{Short and long Mgm1 act together in trans to create a heterodimeric functional unit that mediates mitochondrial fusion}
We tested this hypothesis in vivo by engineering separate versions of s- and l-Mgm1, and used these constructs to individually create s-Mgm1\(^{S224A}\) and l-Mgm1\(^{S224A}\) mutants (Fig. 4 A; Herlan et al., 2004). S224A is a recessive mutation in the G1 motif of the GTPase domain that is predicted to abolish GTPase activity. This mutation does not interfere with the expression and targeting of the protein, but is unable to support mitochondrial fusion in cells (Wong et al., 2003). Our biochemical analyses indicate that neither s-Mgm1\(^{S224A}\) nor l-Mgm1\(^{S224A}\) possess significant, detectable GTPase activity. We observed that mitochondrial fusion is restored in \(\Delta mgm1\) cells when expressing a combination of s-Mgm1 and l-Mgm1\(^{S224A}\), but not when expressing the reciprocal combination of s-Mgm1\(^{S224A}\) and l-Mgm1, as assessed by the ability of cells to grow on glycerol and by the presence
Figure 4. The GTPase activity of l-Mgm1 is not essential for fusion in vivo. (A) Schematic of in vivo l- and s-Mgm1 constructs. Isoforms were constructed by deletion of the first hydrophobic domain (HD1) to produce s-Mgm1 and the second hydrophobic domain (HD2) to produce l-Mgm1. l-Mgm1S224A is functional in vivo as assessed by growth of yeast strains on glycerol media (B) and by mitochondrial morphology (C). Bar, 1 µm. A quantification of mitochondrial morphology in Δs-gm1 cells expressing the indicated l- and s-Mgm1 combinations is shown. Data were normalized to Δs-gm1 + l-Mgm1 + s-Mgm1 and are shown as the mean + SEM (error bars; three experiments, >50 cells/experiment).
Figure 5. l-Mgm1 stimulates s-Mgm1 GTPase activity. (A) Kinetics of GTP hydrolysis of l-Mgm1, s-Mgm1, and l + s-Mgm1. A representative kinetic plot fit to the Hill equation is shown. (B) Kinetic parameters for l-Mgm1, s-Mgm1, and l + s-Mgm1. (C) GTP hydrolysis of s-Mgm1 is stimulated by reconstituted l-Mgm1. The indicated combinations of s- and l-Mgm1/IMC liposomes were analyzed as described in Materials and methods. l- and s-Mgm1–stimulated phosphate release was calculated by subtracting phosphate released for each individual protein from phosphate released when combined. Data from three experiments are shown as the mean + SEM (error bars). (D and E) l- and s-Mgm1 interact in mitochondrial outer membrane fused intermediates in vitro. In outer membrane fused intermediates in vitro, l-Mgm1HA (D) or s-Mgm1HA (E) was immunoprecipitated using α-HA antibodies, and coimmunoprecipitated proteins were analyzed by SDS-PAGE and Western analysis with the indicated antibodies as described (see Materials and methods). l-Mgm1HA/s-Mgm1Flag (cis), l-Mgm1Flag/s-Mgm1Flag (trans), l-Mgm1HA/l-Mgm1Flag (trans), and s-Mgm1Flag/s-Mgm1Flag (trans) complexes were detected and are depicted schematically.
of tubular mitochondrial structures in cells (Fig. 4, B and C). These observations are in agreement with recently published data indicating that the GTPase domain of l-Mgm1 is not required for fusion in vivo (Zick et al., 2009).

To further test whether s- and l-Mgm1 interact, we examined the ability of a mixture of wild-type recombinant s- and l-Mgm1 proteins to hydrolyze GTP in the presence of IMC liposomes. In reactions with both s-Mgm1 and inserted l-Mgm1, we observed a synergistically stimulated maximal rate of GTP hydrolysis, as compared with maximal GTP hydrolysis rates observed in independent l-Mgm1 or s-Mgm1 reactions (Fig. 5, A–C). Full kinetic analysis of GTP hydrolysis revealed that either s-Mgm1 with liposomes or s-Mgm1 together with inserted l-Mgm1 are positively cooperative with respect to GTP, which is consistent with their coassembly (Fig. 5 B).

To assess which isoform was stimulated under these conditions, we determined maximal rates of GTP hydrolysis in reactions containing combinations of wild-type and S224A mutant l- and s-Mgm1 proteins. Consistent with our in vivo analysis, synergistic stimulation of GTP hydrolysis was observed in reactions containing s-Mgm1 and l-Mgm1 S224A, but not in those containing the reciprocal combination of s-Mgm1 S224A and l-Mgm1 (Fig. 5 C). These data are consistent with our model that l- and s-Mgm1 assemble together to form a functional unit required for mitochondrial fusion. Our data further demonstrate that within assembled l-Mgm1/s-Mgm1 structures, the s-Mgm1 GTPase domain is activated. Thus, although l-Mgm1 is not able to hydrolyze GTP, it can assemble with and activate the GTPase domain of s-Mgm1.

We also examined how l- and s-Mgm1 interact within mitochondria during fusion in vitro. Previously, under outer membrane fusion conditions in vitro, we observed a physical Mgm1–Mgm1 interaction from opposing inner membranes (Wong et al., 2003; Meeusen et al., 2004, 2006). Using our Mgm1 constructs engineered to separately express s- and l-Mgm1 (Fig. 2 A), we resolved whether l- and s-Mgm1 interactions occur on the same membrane and/or on opposing inner membranes in outer membrane fused mitochondria. We mixed mitochondria isolated from cells expressing l-Mgm1-FLAG and s-Mgm1-FLAG with mitochondria expressing l-Mgm1-FLAG, and after chemical cross-linking under stage 1 conditions, we immunoprecipitated Mgm1 using anti-FLAG antibodies (Fig. 5 D, see schematic). Western analysis indicated that in outer membrane fused intermediates, l-Mgm1 interacts with l-Mgm1 on opposing inner membranes, and s-Mgm1 interacts with l-Mgm1 on the same membrane (Fig. 5 D). In analogous immunoprecipitation experiments, s-Mgm1 interacted with s-Mgm1 on opposing inner membranes (Fig. 5 E). These data confirm that s- and l-Mgm1 interact with each other across adjacent inner membranes, as well as within the same membrane in mitochondrial fusion intermediates that are arrested at and poised to undergo inner membrane fusion.

Together, our genetic, biochemical, and structural data suggest a model in which the assembly of heterotypic l-s-Mgm1 structures facilitates the fusion of the mitochondrial inner membrane. EM analysis, however, indicated that in contrast to s-Mgm1, higher-ordered structures are not observed on IMC liposomes containing inserted l-Mgm1 alone or inserted l-Mgm1 in combination with s-Mgm1 in the absence or presence of GTP or nonhydrolyzable GTP analogues (unpublished data). Thus, based on our structural analysis, we postulate that l- and s-Mgm1 self assemble together into shorter range structures, not detectable by negative stain EM analysis, that function to mediate mitochondrial membrane fusion by locally deforming the inner membrane. In this context, we think it is likely that l-Mgm1, as the integral membrane protein, functions to both tether inner membranes together and harness GTP-dependent conformational changes of s-Mgm1 that are needed to destabilize lipid bilayers for fusion.

We can also speculate on the potential roles of homotypic l- and s-Mgm1 structures in mitochondria. One obvious function for homotypic membrane-anchored l-Mgm1 dimers is in the maintenance of cristae structures that have previously been shown to require Mgm1/Opa1 function (Frezza et al., 2006; Meeusen et al., 2006). Our observation that the GTPase domain of l-Mgm1 is inactive and that fusion requires a heterodimeric l-Mgm1/s-Mgm1 structure predicts that l-Mgm1 dimers that form from adjacent membranes could function as membrane tethers that would not promote membrane fusion events. Such l-Mgm1 dimers could provide stability to cristae structures and proximity to adjacent inner membranes in the process of fusion. Although s-Mgm1 dimers possess intrinsic affinity for inner membranes, a role as an inner membrane tether seems unlikely given that they do not possess a transmembrane region. However, s-Mgm1 dimers could play a role in inner membrane organization and cristae structure by forming lattices to create a more lamellar inner membrane structure.

Materials and methods

Strains and plasmids

Strains used were W303 (ade1; leu2-3; his3-11, 15; trp1-1, ura3-1; can1-100 MatA) and W303 s-mgm1 (ade1; leu2-3; his3-11, 15; trp1-1, ura3-1; can1-100 MatA, rGM1::HIS3). pRS425 dnm1G385D, as described previously (Naylor et al., 2006), was subcloned into pRS316 using the XhoI and NotI sites, which adds an N-terminal 6xHis tag to yield pFastBacHT–6xHis–s-Mgm1 for purification of s-Mgm1. For expression and purification of l-Mgm1, full-length Mgm1 was first cloned into pFastBacHT–6xHis–s-Mgm1 and inserted into the NcoI and XbaI sites, which adds an N-terminal 6xHis tag to yield pFastBacHT–6xHis–l-Mgm1 for purification of l-Mgm1. For expression in baculovirus-infected insect cells, Mgm1 was PCR amplified beginning at residue A151 and cloned into pFastBacHT using primers of 20–30 nucleotides on either side of the altered nucleotides. Whole plasmid amplification was performed and PCR products were digested with DpnI for 2 h at 37°C to remove template DNA. The amplified product was then subcloned into pRS316 using the XhoI and NotI sites. To create epitope-tagged alleles of Mgm1, we engineered 3′ restriction sites with an amino acid linker (GCCGCGC) by PCR (Meeusen et al., 2006), and cloned the resulting fragment into pRS425. HA and FLAG tags were then subcloned from pFA6a-3HA-TRP1 and pFA6a3FLAG into pRS425-Mgm1-FLAG-CGC-GGC at the 3′ end (Longtine et al., 1998; Hoppins et al., 2009). s-Mgm1 FLAG was made by deleting the first hydrophobic domain (residues 172–179) by site-directed mutagenesis of pRS425-MGM1HA using a PCR-based method with complementary primers of 20–30 nucleotides on either side of the altered nucleotides. Whole plasmid amplification was performed and PCR products were digested with DpnI for 2 h at 37°C to remove template DNA. The amplified plasmid DNA was transformed into DH5α cells. Plasmid DNA was isolated from selected colonies and sequenced to confirm mutations. l-MGM1HA and l-MGM1FLAG were created by deleting the second hydrophobic domain (residues 135–148) by site-directed mutagenesis of pRS425-MGM1HA using a PCR-based method with complementary primers of 20–30 nucleotides on either side of the altered nucleotides. Whole plasmid amplification was performed and PCR products were digested with DpnI for 2 h at 37°C to remove template DNA. The amplified plasmid DNA was transformed into DH5α cells. Plasmid DNA was isolated from selected colonies and sequenced to confirm mutations. l-MGM1FLAG was created by deleting the first hydrophobic domain (residues 172–179) by site-directed mutagenesis of pRS425-MGM1FLAG using a PCR-based method with complementary primers of 20–30 nucleotides on either side of the altered nucleotides. Whole plasmid amplification was performed and PCR products were digested with DpnI for 2 h at 37°C to remove template DNA. The amplified plasmid DNA was transformed into DH5α cells. Plasmid DNA was isolated from selected colonies and sequenced to confirm mutations. l-MGM1HA and l-MGM1FLAG were created by deleting the second hydrophobic domain (residues 135–148) by site-directed mutagenesis of pRS425-MGM1FLAG using a PCR-based method with complementary primers of 20–30 nucleotides on either side of the altered nucleotides. Whole plasmid amplification was performed and PCR products were digested with DpnI for 2 h at 37°C to remove template DNA. The amplified plasmid DNA was transformed into DH5α cells. Plasmid DNA was isolated from selected colonies and sequenced to confirm mutations.

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For s-Mgm1, cells were thawed and resuspended in wash buffer (25 mM Hepes, pH 7.0, 25 mM Pipes, pH 7.0, 500 mM NaCl, 80 mM imidazole, pH 7.4, and 0.1% MEGA-8 [Dojindo]) with protease inhibitor cocktail I (EMD) and 2 mM PMSF. Cells were then lysed by sonication and passage through a 27-gauge/0.5-inch needle. The lysate was centrifuged at 60,000 g for 40 min. The cleared lysate was loaded onto a HiTrap metal chelating column (GE Healthcare) attached to an AKTA prime system (GE Healthcare). s-Mgm1 was eluted from the column by using a linear gradient of 25 mM Hepes, pH 7.0, 25 mM Pipes, pH 7.0, 500 mM NaCl, 500 mM imidazole, pH 7.4, and 0.1% MEGA-8. s-Mgm1 was further purified over a HiLoad 16/60 Superdex 200 preparative gel filtration column (GE Healthcare) in 20 mM Tris-Cl, pH 8.0, 2 mM EDTA, 375 mM NaCl, and 0.01% (v/v) β-mercaptoethanol. Glycerol was added to a final concentration of 20%, rendering the following short freezing buffer (SFB): 23 mM Hepes/Pipes, 450 mM imidazole, 1.4% MEGA-8, and 20% glycerol. The protein was aliquoted, flash frozen in liquid N2, and stored at −80°C.

Immunoprecipitations

Immunoprecipitations were performed to detect cis and trans Mgm1 interactions during outer membrane fusion (Meeusen et al., 2006). Mitochondria were isolated from cells expressing s-Mgm1/Flag-l-Mgm1/HA or l-Mgm1Flag/s-Mgm1 (for l-Mgm1 trans interactions), or s-Mgm1/HA/ l-Mgm1FLAG or l-Mgm1/s-Mgm1FLAG (for s-Mgm1 trans interactions), and then mixed and subjected to outer mitochondrial membrane fusion (Meeusen et al., 2004). After fusion, mitochondria were subjected to chemical cross-linking with 1 mM dithiobis(succinimidyl) propionate (Thermo Fischer Scientific) for 1 h on ice. Reactions were quenched with 100 mM glycine and proteins were subjected to TCA precipitation. Proteins were denatured and solubilized in 100 mM MURB (100 mM MES, pH 7, 1% SDS, and 3 M urea) followed by the addition of 900 µl TWIP (50 mM Tris-Cl, pH 7.5, 150 mM NaCl, 0.5% Tween 20, and 0.1% EDTA). Insoluble proteins were removed by centrifugation at 16,000 g at 4°C for 15 min. Lysate was preadsorbed with 75 µl TWIP-equilibrated protein A agarose beads (Santa Cruz Biotechnology, Inc.) for 30 min at 4°C, then incubated with 8 µl polyclonal HA antibody (Covance) for 1.5 h at 4°C followed by incubation with 75 µl TWIP-equilibrated protein A agarose beads for 45 min at 4°C. Agarose beads were washed three times with 500 µl TWIP and protein was eluted with MURB + 10% β-mercaptoethanol to reduce cross links. Analysis by SDS-PAGE and Western blotting was performed on the total fraction and the eluate fractions with monoclonal α-Flag antibody (Sigma-Aldrich) and monoclonal α-Porin (Covance).

Liposome preparation

Lipids were obtained from Avanti Polar Lipids in chloroform. The following lipids were used in biochemical assays: tetraoleoyl-CL, palmitoyl-oleoyl phosphatidylcholine, palmitoyl-oleoyl phosphatidylethanolamine, soybean phosphatidylglycerol, 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphate (POPA), and 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(serine). Liposomes were mixed in the appropriate ratios and dried under nitrogen gas. The dried lipids were placed under vacuum for at least 1 h, and hydrated in 100 mM NaCl for several hours to overnight. Hydrated liposomes were extruded through a 1-µm nanopore membrane (GE Healthcare) a minimum of 11 times. Compositions of liposomes used in this study were as follows. IM composition: 39% phosphatidylcholine (PC), 29% phosphatidylethanolamine (PE), 20% CL, 8% phosphatidylglycerol (PG), 2% phosphatidic acid (PA), 4% phosphatidylserine (PS), PC/PS/PA, 22% PE, 20% PI, 3% PA, 2% PS, and 6% CL, as described in Zinser and Daum (1995). When the percentage of CL was varied, compensatory changes were made in the percentage of PC present in the lipid mixture. Headgroup-labeled lissamine rhodamine B phosphatidylethanolamine was added to all reactions in trace amounts.

Purification of s- and l-Mgm1

To purify s- and l-Mgm1 in insect cells for expression, we first made bacmid by transforming the baculovirus expression vectors described in “Strains and plasmids” into DH10Bac cells. Bacmid was then transfected into Sf9 insect cells using Transfection Buffer Set A and B (BD). Primary virus was harvested after 5 d and amplified twice in Sf9 cells to make high-titer 3° virus. The high-titer virus was used to infect Hi5 suspension cells (a gift from B. Hammock, University of California, Davis, Davis, CA) for protein expression. Hi5 cells were harvested after 48 h and stored at −80°C.

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using Xhol and XbaI sites with a GGGGCC linker before the C-terminal 6xHis tag. Subsequently, the second hydrophobic domain of Mgm1 containing the processing site to generate s-Mgm1 was deleted by site-directed mutagenesis resulting in pFastBac1-l-Mgm1His.

For s-Mgm1, cells were thawed and resuspended in wash buffer (25 mM Hepes, pH 7.0, 25 mM Pipes, pH 7.0, 500 mM NaCl, 80 mM imidazole, pH 7.4, and 0.1% MEGA-8 [Dojindo]) with protease inhibitor cocktail I (EMD) and 2 mM PMSF. Cells were then lysed by sonication and passage through a 27-gauge/0.5-inch needle. The lysate was centrifuged at 60,000 g for 40 min. The cleared lysate was loaded onto a HiTrap metal chelating column (GE Healthcare) attached to an AKTA prime system (GE Healthcare). s-Mgm1 was eluted from the column by using a linear gradient of 25 mM Hepes, pH 7.0, 25 mM Pipes, pH 7.0, 500 mM NaCl, 500 mM imidazole, pH 7.4, and 0.1% MEGA-8. s-Mgm1 was further purified over a HiLoad 16/60 Superdex 200 preparative gel filtration column (GE Healthcare) in 20 mM Tris-Cl, pH 8.0, 2 mM EDTA, 375 mM NaCl, and 0.01% (v/v) β-mercaptoethanol. Glycerol was added to a final concentration of 20%, rendering the following short freezing buffer (SFB): 23 mM Hepes/Pipes, 450 mM imidazole, 1.4% MEGA-8, and 20% glycerol. The protein was aliquoted, flash frozen in liquid N2, and stored at −80°C.

Reconstitution of l-Mgm1

l-Mgm1 was reconstituted in 100-µl reactions containing liposomes by saturating 0.2 µg liposomes with 0.5% MEGA-8 detergent. 7.0 µM l-Mgm1 was added to the detergent liposomes so the final MEGAB-8 concentration was 0.9%. The lipid, protein, and detergent mixture was incubated for 1 h and then rapidly diluted 10-fold with 20 mM Tris, pH 8.0, and 100 mM NaCl. After reconstitution, l-Mgm1 was at a final concentration of 0.7 µM. Salt extraction of inserted l-Mgm1 was performed to remove associated protein by adding NaCl to l-Mgm1 to a final concentration of 0.5M. To remove uninserted protein, we performed equilibrium centrifugation using an OptiPrep gradient. Specifically, 860 µl of a 53% OptiPrep solution (0.5 M NaCl and 20 mM Tris; Sigma-Aldrich) was added to 500 µl of salt-washed l-Mgm1 to make a 34% OptiPrep solution. This solution was layered with 2 ml 23% OptiPrep floatation buffer (OFB; 20 mM Tris, and 100 mM NaCl, 1 mM 6% OFB, and 750 µl 0% OFB). Liposomes were floated at 200,000 g for 2.5 h (SW-55 rotor; Sorvall) and then extracted from the gradient in a 500-µl fraction.

GTPase assays

All assays were performed in 5 mM MgCl2, 100 mM NaCl, and 20 mM Tris, pH 8.0. The malachite green assay was used to detect free phosphate as described previously (Quan and Robinson, 2005). s-Mgm1 GTPase reactions were performed by first incubating 1 µM s-Mgm1 with 0.4 mg/ml liposomes for 15 min at RT. Buffer was added to give a final concentration of 0.2 mg/ml liposomes in 5 mM MgCl2, 20 mM Tris, pH 8.0, 100 mM NaCl, and 500 µM GTP (unless otherwise stated). The reaction was stopped after 1 h by addition of 125 mM EDTA. 200 µl of malachite green was added to 25 µl of reaction and the absorbance at 650 nm was determined using a microtiter Max Plus plate reader (MDS Analytical Technologies). GTP hydrolysis was determined by subtracting the amount of phosphate release in a no-protein control from that of the protein reactions. l-Mgm1 and l-Mgm1 + s-Mgm1 GTPase assays were done after reconstitution by adding 0.3 µM of inserted l-Mgm1 to 0.1 mg/ml liposomes and 0.5 µM s-Mgm1 or buffer. The proteins were incubated for 30 min at RT before adding reaction buffer (25 mM MgCl2 and 20 mM Tris-Cl, pH 8.0) with the nucleotide to yield 0.3 mg/ml liposomes, 0.2% MEGA-8, 5 mM MgCl2, 20 mM Tris, 100 mM NaCl, and 500 mM GTP (unless otherwise stated). GTP hydrolysis activity was determined using malachite green as described above.

Equilibrium sucrose gradient centrifugation of liposomes

Purified s-Mgm1 or freezing buffer was preincubated with 0.4 mg/ml liposomes for 20 min in a volume of 50 µl. The protein/lipid mix was diluted to 100 µl in floatation buffer (FIB) so that the final liposome concentration
was 0.2 mg/ml in 20 mM Tris, pH 8.0, 100 mM NaCl, 5 mM MgCl₂, and 300 μM nucleotide, then incubated for 1 h. 400 μl of FIB + 50% sucrose (FIB50) was added and mixed thoroughly, and the resulting 40% sucrose reaction mix was transferred to a centrifuge tube. The 40% sucrose layer was overlaid with 1 ml FIB30, 500 μl FIB10, and 250 μl FIB0.

Sucrose step gradients were centrifuged in a SW-55 rotor at 200,000 g for 2.5 h at 4°C. Five 450-μl fractions were pipetted from the top and the pellet was resuspended in 450 μl FIB, resulting in a total of six fractions. The top three fractions were pooled as the float fractions (F) because the majority of liposomes were found in these fractions, as determined by measuring the rhodamine fluorescence of each fraction using a Spectra Max Plus plate reader with the excitation and emission monochromators set at 550 nm and 590 nm, respectively. The bottom two fractions were combined with the pellet and were pooled as a single float fraction (P).

For quantification of the fraction of protein that floated with the liposomes, equal volumes of F and P fractions were analyzed by SDS-PAGE and Western analysis using the primary antibody, s-Mgm1, and an IRDye 800 CW secondary antibody (LI-COR Biosciences) for detection. The immunoreactive bands were detected with the Odyssey Infrared Imaging System (LI-COR Biosciences) and quantified using the accompanying software.

Hydrodynamic analysis
The Svedberg coefficients of s- and l-Mgm1 were determined as described previously (Tinggeman et al., 2005). 100 μl of 0.5 μM Mgm1 in 5% sucrose was loaded on top of a 2 ml 5–20% sucrose gradient and subjected to a 4-h ultracentrifugation spin at 200,000 g in an SW-55 rotor. Molecular mass markers with known sedimentation coefficients (17–0445-01; GE Healthcare) were simultaneously run over a separate gradient. 10 × 0.2-ml fractions were collected manually from the top of the gradients for analysis. Svedberg coefficients were estimated from a standard curve from fraction number versus known sedimentation coefficients of the protein standards.

To determine the Stokes’ radii of s- and l-Mgm1, purified protein at 0.5 μg/μl or gel filtration standards (Bio-Rad Laboratories) were run over an HiLoad 16/60 SuperSet 200 preparative gel filtration column using an AKTA prime system (GE Healthcare). For each protein component of gel filtration standards, values of Kₛ²/₃ were plotted versus Stokes’ radii to construct a Porath correlation standard curve. Molecular mass was calculated from combined sucrose gradient and gel filtration data using the method developed by Siegel and Monty (1966). l-Mgm1 was analyzed identically to s-Mgm1, except the 5–20% sucrose gradient contained 0.75% MEGA-8.

s-Mgm1 cross-linking
s-Mgm1 was dialyzed into 25 mM Hepes/Pipes, 300 mM NaCl, and 5% glycerol for 1 h at 4°C. A 10-fold molar excess of the nonreversible cross-linker bis(sulfosuccinimidyl)suberate (BS3; Thermo Fisher Scientific) was added to s-Mgm1 preincubated with 0.2 mg/ml liposomes or liposome buffer for 15 min at RT. The solutions were incubated with 80 mM Tris-Cl, pH 8.0, for 15 min at RT. Cross-linked products were resolved on a 3–8% Tris-acetate gel (NuPage Novel; Invitrogen) and visualized by Coomassie blue staining. HiMark molecular weight markers (Invitrogen) were used as standards.

Mgm1 structural analysis
2D crystals were grown under a lipid monolayer covering the air-water interface, as described by Lévy et al. (1999). In brief, ~60-μl wells in a Teflon block were filled with Tris 20 mM, pH 8.0, such that a flat air-water interface was formed. A 0.5-ml drop of IMC lipid mixture at 0.1 mg/ml concentration dissolved in a 9:1 chloroform/methanol mixture was spread on the water surface. After 24 h of incubation time at RT, 5 μl of 0.1 mg/ml s-Mgm1 suspension was added through a side well. After an additional 24-h incubation time, the monolayers were recovered by applying non-glowl discharged carbon-coated grids to the surface of the wells. Grids were removed, blotted, stained with 2% uranyl acetate, blotted again, air dried, and transferred into a transmission electron microscope (JEM-1230; JEOL, Ltd.), operated at an acceleration voltage of 120 kV. Images were recorded at 30,000x nominal magnification with a F214 CCD camera (2,048 x 2,048 pixels; TVIPS GmbH), which corresponds to a pixel size of 3.8 Å at the specimen level.

Images were processed using the 2dx program system (Gispson et al., 2007a,b), which is based on the MRC software (Crowther et al., 1996). A total of nine images of nontilted negatively stained sMgm1 2D crystals were processed and merged, yielding a projection map at 31 Å resolution. Images from six tilted negatively stained 2D crystals at a 30° sample tilt and eight tilted crystals at a 45° sample tilt were processed and merged into a 3D reconstruction, and the same 31 Å resolution cutoff was applied. The final map was created from 1,033 observed amplitudes and phases, and had an overall phase residual of 26.8°. The overall weighted phase residual was 9.6°.

Homology modeling of s-Mgm1
Homology modeling was performed with the program Modeller (Martí-Renom et al., 2000) using the structures of dynamin A, dynamin 1, and dynamin-like bacterial protein (PDB accession nos. 1JYW, 2AKA, and 2J69) as references. Sequence alignment for Mgm1, dynamin A, and dynamin 1 was performed within Modeller. Because of the low homology between Mgm1 and the dynamin-like bacterial protein, the sequence alignment for the latter was obtained from the mGenTHREADER server (McGuffin and Jones, 2003). The full alignment was checked with the mumsa program (Lassmann and Sonnhammer, 2002), and 1,000 models were generated using Modeller (Martí-Renom et al., 2000). The five models with the lowest DOPE score were manually inspected, and the best model that did not show knots in coil regions was selected. This model had a DOPE score of ~59,372.

The dimeric form of s-Mgm1 was obtained by 100 rounds of blind docking with the GRAMM-X server (Tovchigrechko and Vakser, 2005), followed by a local search with Rosetta (Gray et al., 2003). From the 1,000 docked models, the best 10 dimeric arrangements were selected by manual verification of the presence of a twofold symmetry and a head-to-head arrangement of the monomers. The model that showed the best fit with the experimental 3D reconstruction and the lowest number of atoms outside of the volume was manually selected (Fig. 4 D).

Figures of the 3D reconstruction were produced with Chimera (Pettersen et al., 2004) and the homology model was rendered using POVRay (http://www povray.org).

Negative stain EM
Reactions containing l-Mgm1, s-Mgm1, and l+s-Mgm1 or corresponding storage buffers were prepared as described for the GTPase assays, and nucleotide was added as indicated. When the reactions were complete, a glow-discharged carbon-coated grid was placed on a 1-µl drop of the reaction mix for 1 min. The grid was then washed with 2% uranyl acetate for 1.5 min, blotted, and air dried. The grids were imaged at 37,000x using a transmission EM (CM120 [Phillips] with a biotwin lens [FEI]) operating at 80 kV, and images were captured digitally using a MegaScan CCD camera (Gatan) and the Digital Micrograph software package (Gatan).

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