Yeast Miro GTPase, Gem1p, regulates mitochondrial morphology via a novel pathway

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Cell signaling events elicit changes in mitochondrial shape and activity. However, few mitochondrial proteins that interact with signaling pathways have been identified.Candidates include the conserved mitochondrial Rho (Miro) family of proteins, which contain two GTPase domains flanking a pair of calcium-binding EF-hand motifs. We show that Gem1p (yeast Miro; encoded by YAL048C) is a tail-anchored outer mitochondrial membrane protein. Cells lacking Gem1p contain collapsed, globular, or grape-like mitochondria. We demonstrate that Gem1p is not an essential component of characterized pathways that regulate mitochondrial dynamics. Genetic studies indicate both GTPase domains and EF-hand motifs, which are exposed to the cytoplasm, are required for Gem1p function. Although overexpression of a mutant human Miro protein caused increased apoptotic activity in cultured cells (Fransson et al., 2003. J. Biol. Chem. 278:6495–6502), Gem1p is not required for pheromone-induced yeast cell death. Thus, Gem1p defines a novel mitochondrial morphology pathway which may integrate cell signaling events with mitochondrial dynamics.

Introduction

Mitochondria are essential organelles whose morphology and activity adapt to physiological stresses and the changing metabolic state of the cell. Balanced mitochondrial division and fusion events help determine mitochondrial morphology in most organisms, and conserved proteins that mediate these processes have been identified (Hermann and Shaw, 1998; Yaffe, 1999; Jensen et al., 2000; Yoon and McNiven, 2001; Shaw and Nunnari, 2002; Westermann, 2002; Mozdy and Shaw, 2003; Osteryoung and Nunnari, 2003; Scott et al., 2003; Westermann, 2003; Chen and Chan, 2004). Much less is known about pathways that facilitate communication between mitochondria and other cellular compartments. Such pathways must exist, because mitochondrial division and cytochrome c release are essential for caspase-dependent apoptosis (Frank et al., 2001), mitochondria buffer cytosolic calcium levels (Brini and Carafoli, 2000; Sayer, 2002; Rizzuto et al., 2003; Jacobson and Duchen, 2004), and mitochondrial mass increases in response to cell growth. However, few mitochondrial proteins that interface with these pathways have been characterized.

The conserved mitochondrial Rho (Miro) family of proteins has the potential to coordinate cellular responses with mitochondrial dynamics and function. Miro proteins contain two GTPase domains, a pair of calcium binding EF-hand motifs (Fransson et al., 2003), and a COOH-terminal transmembrane domain (Wolff et al., 1999). The two Miro family members in mammalian cells, Miro-1 and Miro-2, localized to mitochondria in indirect immunofluorescence experiments (Fransson et al., 2003). Overexpression of Miro-1 containing a putative GTPase I activating mutation caused mitochondrial aggregation and increased apoptotic index (Fransson et al., 2003). These observations suggest that Miro proteins facilitate apoptosis and/or regulate mitochondrial dynamics.

Models for Miro function must take into account the behavior of cells lacking Miro protein, the importance of conserved functional domains, and the submitochondrial localization and topology of Miro proteins. For example, functional EF-hand motifs exposed to the cytoplasm would be in a position to sense changing cytosolic calcium levels, whereas localization of these motifs to the mitochondrial matrix could allow Miro to monitor organellar calcium. To address these issues and learn more about the cellular role of Miro, we analyzed the function of the single Miro homologue, Gem1p, in S. cerevisiae. Our studies demonstrate that Gem1p is a tail-anchored outer mitochondrial membrane protein required for normal mitochondrial dynamics. In addition, both GTPase domains and EF-hand
motifs, which are exposed to the cytoplasm, are required for function. Based on our findings, we suggest that Gem1p defines a novel pathway controlling mitochondrial dynamics.

**Results**

Yeast Miro (Gem1p) belongs to a novel gene family with predicted GTPases domains, EF-hand motifs, and a COOH-terminal transmembrane domain

The single Miro protein in *S. cerevisiae*, encoded by YAL048C, shares 30% identity and 48% similarity with the human Miro-1 protein. GEM1 (for GTPase EF-hand protein of mitochondria) encodes a 662–amino acid protein with predicted molecular mass of 75.2 kD (Fig. 1 A). The GTPase I domain contains G1, G2, G4, and G5 motifs characteristic of Ras and Rho-like proteins, but lacks an apparent consensus G3 motif and the Rho-specific sequence insert (Bourne et al., 1991; Wennerberg and Der, 2004). The GTPase II domain is not closely related to Ras, Rho, or other GTPase families, but contains recognizable G1, G2, G4, and G5 motifs (unpublished data).

Gem1p contains a pair of CaM-like EF-hand motifs, each consisting of two α-helices that flank a 12 residue loop (Nakayama and Kretsinger, 1994; Ikura, 1996; Lewit-Bentley and Rety, 2000). In CaM-like EF-hand proteins, carboxylate side chains in positions 1, 3, 5, 9, and 12 of the loop act as electron donors for calcium coordination (Lewit-Bentley and Rety, 2000). This differs from a second class of EF-hand motifs (e.g., the first motif in the protein S100A), where calcium is coordinated via backbone oxygen atoms (Fig. 1 C).

Tail-anchored outer mitochondrial membrane proteins contain positively charged residues flanking a hydrophobic transmembrane sequence (Wattenberg and Lithgow, 2001; Borgese et al., 2003). Although our sequence alignments indicate that this targeting motif is also present in hMiro-1 (Fig. 1 D) and Gem1p (Wolff et al., 1999), the possibility that Miro family members are embedded in the outer mitochondrial membrane has not been tested. As in other known tail-anchored mitochondrial proteins like Fis1p (Wattenberg and Lithgow, 2001; Habib et al., 2003; Horie et al., 2003), hMiro-1 and Gem1p contain several positively charged residues adjacent to the transmembrane domain (Fig. 1 D, bold). These combined structural features suggest that Gem1p has calcium binding and GTPase activities and is tail-anchored in the outer mitochondrial membrane.

Cells lacking Gem1p respire poorly on synthetic glycerol medium

To determine whether Gem1p is required during mitotic growth, we replaced the entire GEM1 ORF with a HIS3 cassette in the W303 strain background. This gem1Δ strain was viable, but grew at a slightly reduced rate relative to wild type on solid medium containing dextrose (Fig. 2 A, SDextrose 30°C). The gem1Δ strain grew significantly slower than wild type on minimal media containing the nonfermentable carbon source glycerol (Fig. 2 A, SGlycerol 30°C). This defect was enhanced when the strain was grown at 37°C (Fig. 2 A). Introduction of a plasmid encoding wild-type Gem1p restored wild-type growth in gem1Δ cells (Fig. 2 A). Interestingly, DAPI staining indicated that mtDNA nucleoids were retained in ~75% of gem1Δ cells cultured in dextrose-containing medium overnight. Even after extended log-phase growth in dextrose-containing medium, 43% of gem1Δ cells retain detectable mtDNA (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200405100/DC1). These findings suggest that loss of Gem1p may cause secondary defects in mtDNA expression or integrity.
gem1Δ cells have distorted mitochondrial morphology but maintain inner membrane cristae structures

In wild-type yeast, mitochondria form a branched tubular network located at the cell cortex (Stevens, 1981; Koning et al., 1993). When visualized with a mitochondrial-targeted GFP (mito-GFP), 86.6% of cells in a wild-type population displayed a tubular mitochondrial network (Fig. 3 A). In contrast, cells lacking Gem1p had pronounced defects in mitochondrial distribution and morphology. As shown in Fig. 3 C, 53.7% of gem1Δ cells contained mitochondria that were globular with an irregular perimeter. Aberrant mitochondria of this type were never observed in wild-type strains (n > 1000). These large mitochondria did not appear to be collapsed tubules, as labeling with an outer membrane targeted form of GFP resulted in rim staining that enclosed matrix-targeted RFP (unpublished data). Other mitochondrial morphologies, including grape-like clusters (24%; Fig. 3 D) and collapsed mitochondria (16%; Fig. 3 B) were also observed in gem1Δ cells. The grape-like mitochondrial clusters were larger and fewer in number than mitochondrial fragments observed in a mitochondrial fusion mutant (fzo1Δ; Fig. 3 E). Collapsed mitochondria (Fig. 3 B) were frequently thicker than wild-type mitochondrial tubules and had variations in diameter along the length of the tubule. Because these collapsed mitochondria retain their tubular structure, they were not included in the “mutant” category during phenotypic quantification. Globular (Fig. 3 C), grape-like (Fig. 3 D), and fragmented (Fig. 3 E) mitochondria were scored in a single category, designated “Percent mutant” (Table I), unless otherwise indicated. A plasmid containing GEM1 expressed from its own promoter restored tubular networks in 69.3% of the gem1Δ population (Table I). In the FY strain background, deletion of gem1Δ had a less severe affect on mitochondrial morphology (unpublished data). Overexpression of full-length Gem1p or Gem1p(1-632) lacking the putative transmembrane domain from a galactose-inducible promoter did not cause defects in mitochondrial morphology in wild-type cells (unpublished data).

To determine whether inner mitochondrial membrane ultrastructure was disrupted in gem1Δ cells, we performed transmission electron microscopy (TEM). Both wild-type and gem1Δ sections contained mitochondrial profiles with well-developed inner membrane cristae (Fig. 4). In wild-type cells, mitochondria in longitudinal sections appeared tubular (Fig. 4 A) and in cross section (Fig. 4 B) were surrounded by a double membrane. gem1Δ sections often exhibited large mitochondrial profiles with double membranes and lighter matrix staining, consistent with the idea that these organelles are swollen (Fig.
4, C–F). These findings provide additional evidence that globular mitochondria in gem1Δ/H9004 cells are not composed of aggregated tubules. Interestingly, gem1Δ/H9004 mitochondria in a few TEM sections appeared partially or completely engulfed by vacuoles, suggesting that aberrant mitochondria are turned over via autophagy (unpublished data).

Disruption of actin cytoskeleton or ER organization can cause secondary defects in mitochondrial morphology and inheritance (Drubin et al., 1993; Hermann et al., 1997; Prinz et al., 2000; Singer et al., 2000). However, mitochondrial morphology defects in gem1Δ cells are not due to disruption of these structures. Alexa-Phalloidin staining studies showed actin cytoskeletal organization was normal in 99.5% of GEM1 and gem1Δ cells (n = 200). Moreover, actin-based transport of mitochondria remained intact as medium and large-budded gem1Δ cells always inherit mitochondria (Fig. 5). The lack of mitochondria observed in some small-budded gem1Δ cells is likely a delay in movement of large globular organelles to the bud. Vacuole inheritance, another actin-based process in yeast (Hill et al., 1996), was not disrupted in gem1Δ strains. In pulse-chase experiments, FM 4-64–labeled vacuoles were inherited by 98.3 and 95.8% of wild-type and gem1Δ buds, respectively (n = 120). In addition, yeast nuclei (visualized by DAPI staining) and ER morphology (visualized by expression of Sec63-GFP) were normal in the absence of Gem1p (unpublished data). Vacuolar morphology and endocytic function also appeared wild type when tested by monitoring internalization of the vital dye FM 4-64 (unpublished data). Together, these data suggest that disruption of GEM1 causes primary defects in mitochondrial morphology and function but does not disturb organization of other cellular structures.

Table I. Mitochondrial morphology in GEM1 and gem1Δ cells

| Genotype | Plasmid       | Percent tubular network | Percent collapsed | Percent mutant |
|----------|---------------|-------------------------|-------------------|---------------|
| GEM1     | None          | 86.6 ± 2.6              | 6.7 ± 2.7         | 6.7 ± 2.3     |
| gem1Δ    | None          | 3.4 ± 1.1               | 16.1 ± 3.2        | 80.5 ± 3.3    |
| GEM1     | pRS416        | 85.8 ± 1.8              | 7.4 ± 3.0         | 6.8 ± 1.9     |
| gem1Δ    | pRS416        | 2.2 ± 1.3               | 19.6 ± 6.2        | 78.2 ± 6.9    |
| GEM1     | pRS416-GEM1   | 69.3 ± 5.0              | 14.8 ± 4.6        | 15.8 ± 3.8    |
| gem1Δ    | pYX142        | 84.3 ± 1.5              | 9.0 ± 1.0         | 6.7 ± 1.2     |
| GEM1     | pYX142        | 0.9 ± 0.8               | 20.7 ± 5.3        | 78.5 ± 4.5    |
| gem1Δ    | pYX142-GFP-GEM1 | 57.9 ± 11.3          | 24.1 ± 11.6       | 18.0 ± 3.2    |

n_cells ≥ 500.
Mitochondria were visualized by expressing mito-RFP or mito-GFP.
Percent tubular network contains wild-type morphology shown in Fig. 3 A.
Percent collapsed contains aberrant morphology shown in Fig. 3 B.
Percent mutant category contains aberrant morphologies shown in Fig. 3 (C–E).

Figure 4. Inner membrane cristae are present in globular gem1Δ mitochondria. Mitochondria in wild-type cells (JSY7000) are tubular in longitudinal sections (A) and spherical in cross section (B) and contain inner membrane cristae. Profiles of globular mitochondria in gem1Δ strains (JSY7002, C–F), are larger than wild type but contain well-developed cristae. n, nucleus. Bars, 1 μm.

Figure 5. Mitochondrial inheritance is not blocked in the absence of Gem1p. Mito-GFP-labeled mitochondria were observed in strains grown to mid-log phase. Large- or small-budded cells containing mitochondria in the bud were scored. n > 250.
Gem1p is a tail-anchored outer membrane protein with its functional domains exposed to the cytoplasm

To determine where Gem1p localizes in vivo, we fused GFP to the NH$_2$ terminus of a constitutively expressed Gem1p (pYX142-GFP-GEM1). This construct complements the glycerol growth (Fig. 2 B) and mitochondrial morphology defects (Table I) of gem1 null cells at 30°C. When expressed in wild-type cells, GFP-Gem1p uniformly labeled mitochondrial tubules that were cosstained with MitoFluor red 589 (Fig. 6).

Subcellular fractionation studies confirmed that Gem1p is a mitochondrial protein (Fig. 7 A). When whole cell extracts (WCEs) from wild-type cells expressing GFP-Gem1p were separated into post-mitochondrial supernatant (PMS) and mitochondrial pellet (MITO) fractions, GFP-Gem1p was found in the MITO, along with the integral outer mitochondrial membrane protein porin. In contrast, GFP-Gem1p was not detected in the PMS, which contained the cytoplasmic protein 3-phosphoglycerate kinase (3-PGK).

Protease protection assays indicated that GFP-Gem1p is an outer mitochondrial membrane protein (Fig. 7 B). When intact mitochondria were treated with proteinase K (PK), both Fzo1p (an outer membrane protein) and GFP-Gem1p were efficiently degraded. In contrast, the intermembrane space protein Cyb2p was protected from externally added PK, indicating that the outer membrane was not compromised. PK did degrade Cyb2p when mitochondria were subjected to osmotic shock, a treatment that causes mitochondria to swell, rupturing the outer membrane while leaving the inner membrane intact. When membranes were solubilized using Triton X-100 in the presence of PK, the matrix protein mtHsp70 was also degraded. Because the GFP tag is fused to the NH$_2$ terminus of the protein, these results demonstrate that Gem1p is localized to the outer mitochondrial membrane with its NH$_2$ terminus exposed to the cytoplasm.

Although the membrane association of Miro proteins has never been analyzed, several observations suggest that these GTPases are tail-anchored outer membrane proteins. First, like other mitochondrial outer membrane proteins, Miro family members do not contain a classical mitochondrial targeting signal. Second, previous studies showed that, like other tail-anchored proteins, COOH-terminal–tagged forms of Gem1p were not expressed (Wolff et al., 1999). Third, Miro family members, including Gem1p, contain a predicted COOH-terminal transmembrane domain (Fig. 1 D). To test whether Gem1p is an integral membrane protein, we performed carbonate extraction on isolated mitochondria. Upon sodium carbonate treatment under alkaline conditions, peripheral membrane proteins and soluble components can be released from mitochondria into the supernatant, whereas integral membrane proteins remain membrane associated. The mitochondrial membrane pel-
Gem1p is not required for mitochondrial division and fusion events

The yeast mitochondrial network is maintained by balanced division and fusion events. Mutations in genes required for fusion, including FZO1 and UGO1, cause mitochondrial fragmentation due to unopposed division (Bleazard et al., 1999; Sesaki and Jensen, 1999, 2001). In contrast, mutations in genes required for mitochondrial division, like DNM1, result in net formation due to ongoing fusion (Bleazard et al., 1999; Sesaki and Jensen, 1999). Because Gem1p affects mitochondrial morphology, we asked whether Gem1p is required for Dnm1p-mediated mitochondrial division events or Fzo1p-mediated fusion events.

A yeast mating assay can be used to monitor mitochondrial fusion in vivo. After zygote formation, mitochondria from each haploid cell fuse, and their contents mix (Nunnari et al., 1997). When mitochondria from each haploid parent are labeled either with mito-RFP or mito-GFP, mixing of contents in fused mitochondria appears as complete overlap of red and green fluorescence. Previous studies showed that zygotes lacking the division gene DNM1 (dnm1Δ) fuse as efficiently as wild type (Bleazard et al., 1999). However, zygotes lacking the fusion genes UGO1 (ugo1Δ) (Sesaki and Jensen, 2001) or FZO1 (Hermann et al., 1998) contain fragmented mitochondria that fail to fuse. Disruption of DNM1 in ugo1Δ and fzo1Δ strains blocks fragmentation but does not restore fusion (Bleazard et al., 1999; Sesaki and Jensen, 2001).

As shown in Table II, mitochondrial fusion occurs at near wild-type levels (GEM1 DNM1 UGO1, 98.1%) in zygotes lacking Gem1p (gem1Δ DNM1 UGO1, 85.9%). Absence of Dnm1p did not dramatically change the efficiency of mitochondrial fusion (gem1Δ dnm1Δ UGO1, 80.9%). In contrast, mitochondrial fusion was completely blocked in zygotes lacking both Ugo1p and Dnm1p (GEM1 dnm1Δ ugo1Δ, 0.0%). The slightly reduced rate of mitochondrial fusion in the absence of Gem1p is likely due to morphology changes that prevent efficient mitochondrial collision in zygotes. Nevertheless, aberrant mitochondria in gem1Δ cells are clearly competent for fusion. Moreover, in gem1Δ dnm1Δ strains, some mitochondrial nets can be observed, indicating that mitochondrial fusion is occurring (unpublished data).

| Strains mated | No. of fused mitochondria /total scored | Percent fusion observed |
|---------------|----------------------------------------|------------------------|
| GEM1 DNM1 UGO1 × gem1Δ DNM1 UGO1 | 157/160 | 98.1 |
| gem1Δ DNM1 UGO1 × gem1Δ DNM1 UGO1 | 177/206 | 85.9 |
| gem1Δ dnm1Δ UGO1 × gem1Δ dnm1Δ UGO1 | 131/162 | 80.9 |
| GEM1 dnm1Δ ugo1Δ × GEM1 dnm1Δ ugo1Δ | 0/157 | 0.0 |

Large-budded zygotes containing both mito-RFP and mito-GFP were scored at 2.5–4 h after mating on YPD.

Figure 8. The COOH-terminal 45 aa of Gem1p are sufficient for mitochondrial targeting. Images of GEM1 cells (JSY7000) expressing GFP alone (A–D) or GFP-Gem1p (aa618-662) (E–H) are shown. Mitochondria were visualized simultaneously with mito-RFP. v, vacuole. Bar, 5 μm.
Mitochondrial fragmentation in an fzo1 mutant requires active mitochondrial division. To test whether Gem1p is required for mitochondrial division, we asked if gem1Δ could prevent division and fragmentation in an fzo1Δ mutant strain. In both the gem1Δ fzo1Δ test strain and the GEM1 fzo1Δ control strain, mitochondria fragment in the majority of the population (91.1%, 99.0% respectively, Table III). Similar results were obtained when the temperature-sensitive fzo1-1 allele was used for these studies (unpublished data). Moreover, in gem1Δ cells, components of the division machinery, including Dnm1p, Mdv1p, and Fis1p (Otsuga et al., 1998; Bleazard et al., 1999; Fekkes et al., 2000; Mozdy et al., 2000; Tieu and Nunnari, 2000; Cerveny et al., 2001) were properly localized as mitochondrial puncta (Dnm1p, Mdv1p; Fig. 9, A and B) or uniformly on the mitochondrial network (Fis1p; Fig. 9 C), supporting the notion that division is not impaired in this strain. Steady-state levels of Fis1p, Dnm1p, Mdv1p, and Fzo1p in gem1Δ cells were wild type (unpublished data), suggesting that Gem1p does not regulate stability of these mitochondrial morphology proteins.

Both GTPase domains I and II and the EF-hand motifs are required for Gem1p function

To determine whether the predicted functional domains of Gem1p were required for its role in maintenance of tubular mitochondrial morphology, we generated mutations in conserved residues of each GTPase domain (Fig. 1 B) or the EF-hand motifs (Fig. 1 C). Similar mutations in known GTPases and EF-hand motifs have been shown to abrogate function. Mutation of K16 in Ras disrupts GTP nucleotide binding (Sigal et al., 1986). Mutation of S17 in Ras reduces affinity for GTP by 20–40-fold (Feig and Cooper, 1988). Finally, mutation of the conserved threonine (T35) residue in Ras disrupts GTPase activating protein (GAP) binding such that GTP hydrolysis cannot be stimulated efficiently (Cales et al., 1988). In GEM1, none of these mutations caused dominant mitochondrial morphology defects in wild-type strains (unpublished data). GFP-tagged versions of all mutant Gem1 proteins were expressed and localized to mitochondria (unpublished data).

When gem1Δ cells contained a plasmid expressing full-length wild-type Gem1p, 59.9% of cells in a population had tubular mitochondrial networks, whereas only 14.6% contained mutant mitochondria (Table IV). Mutation of conserved residues in the nucleotide binding domain of GTPase I completely abrogated protein function. When K18 was mutated to alanine (K18A) in Gem1p, 75.8% of cells displayed mutant mitochondria, similar to gem1Δ cells containing the vector with no insert (83.3% mutant). Similarly, expression of Gem1p(S19N) did not rescue gem1 null cells (79.1% mutant). Mutation in the effector binding domain of GTPase I did not prevent Gem1p function, as 57.8% of gem1Δ cells expressing Gem1p(T33A) contained tubular networks. Based on mutational analysis of known GTPases, these data suggest that the nucleotide binding activity of GTPase I domain is required for function. Whether this domain requires accessory proteins for nucleotide exchange or hydrolysis remains to be determined.

Figure 9. Previously characterized mitochondrial morphology proteins are properly localized in gem1Δ cells. Colocalization of mitochondria and GFP-labeled Dnm1p [A], Mdv1p [B], Fis1p [C], and Mmm1p [D] was observed in log phase GEM1 and gem1Δ cells. Bars, 5 μm.
The GTPase II domain of Gem1p is also important for function (Table IV). Similar mutations of the conserved lysine and serine residues within the GTPase II P-loop and the conserved threonine in the effector binding domain were analyzed (Fig. 1B). In gem1Δ strains expressing Gem1p(S462N), 80.3% of cells contained mutant mitochondria, similar to gem1Δ cells containing vector with no insert (83.3% mutant). When gem1Δ strains expressing Gem1p(K461A) or Gem1p(T480A) were scored, the percentage of cells with mutant mitochondria (55.3 and 48.3%, respectively) was somewhat lower than vector alone (83.3%), suggesting that these mutant proteins may have retained partial function. Because Gem1p(S462N) cannot rescue gem1Δ mutant mitochondria, these data demonstrate that the GTPase II domain is essential for Gem1p function.

Finally, we examined the role of the EF-hand motifs in Gem1p function. Calcium binding activity of EF-hand motifs in CaM can be abrogated by mutation of a conserved loop glutamine to lysine (Feng and Stemmer, 2001). We analyzed a mutant protein in which either residue E225 of EF I or residue E354 of EF II were replaced with a lysine (K) residue (Fig. 1C). As some multiple EF-hand proteins require mutation of both EF-hand motifs to cause a marked decrease in protein function (Janssens et al., 2003), a protein containing both E225K and E354K mutations was also analyzed. As shown in Table IV, Gem1p containing only one EF-hand motif mutation had partial function (E225K 28.7% tubular, E354K 42.2% tubular). In contrast, the function of Gem1p(E225K,E354K) was dramatically impaired with only 14.3% of cells in a population displaying branched tubular mitochondria. Mutation of either or both of these conserved glutamine residues to alanine partially abrogated protein function similar to the lysine substitutions (unpublished data). These results establish that the EF-hand motifs are important for Gem1p function.

Gem1p is not required for cell death induced by α-factor mating pheromone
Overexpression of a putative hyperactive Miro-1 variant in COS-7 cells results in a high rate of spontaneous apoptosis (Fransson et al., 2003). Previous studies in yeast demonstrated that apoptosis-like cell death occurs after prolonged treatments with α-factor, a pheromone that induces mating and calcium signaling pathways that help to prevent cell death (Iida et al., 1990; Cyert et al., 1991; Cyert and Thorner, 1992; Foor et al., 1992; Moser et al., 1996; Withee et al., 1997; Muller et al., 2001; Severin and Hyman, 2002). To test whether Gem1p might perform a similar “pro-death” function in yeast, we measured the degree of cell death in wild-type and gem1 mutant cultures after a 10-h treatment with α-factor in the presence and absence of FK506, an inhibitor of calcineurin in the calcium signaling pathway. The levels of cell death in gem1 cultures were indistinguishable from those of wild-type cultures at all incubation times (Fig. 10 and not depicted). Thus, unlike Miro-1 in mammalian cells, Gem1p had no obvious role in these forms of yeast cell death.

Discussion
The mechanisms that allow mitochondria to receive or generate signals that modulate organelle shape and function are not well understood. Here, we show that cells lacking Gem1p have abnormal mitochondrial morphology and reduced respiratory function. Both GTPase domains and EF-hand motifs of Gem1p are required for its role in regulating mitochondrial dynamics. Gem1p is tail-anchored in the outer mitochondrial membrane, exposing these functional domains to the cytoplasm where they are available to interact with cell signaling pathways or other cellular structures. These findings suggest that Miro family members function in a novel pathway that integrates cell signaling with mitochondrial dynamics.

Table III. Mitochondrial division and fragmentation occurs in fzo1Δ lacking Gem1p

| Genotype            | Percent tubular network | Percent collapsed | Percent globular and large grapes | Percent fragmented |
|---------------------|-------------------------|-------------------|----------------------------------|--------------------|
| GEM1 fzo1           | 82.7 ± 5.7              | 0.0 ± 0.0         | 0.0 ± 0.0                        | 9.3 ± 3.1          |
| gem1 fzo1           | 0.0 ± 0.0               | 18.3 ± 6.6        | 68.5 ± 8.0                       | 13.2 ± 5.3         |
| GEM1 fzo1Δ          | 0.0 ± 0.0               | 1.0 ± 1.8         | 0.0 ± 0.0                        | 99.0 ± 1.8         |
| gem1 fzo1Δ          | 0.0 ± 0.0               | 3.2 ± 2.2         | 5.6 ± 6.3                        | 91.1 ± 6.9         |

n = 300.
Mitochondria were visualized by expressing mito-GFP.
Percent tubular network contains wild-type morphology shown in Fig. 3 A.
Percent collapsed contains aberrant morphology shown in Fig. 3 B.
Percent globular and large grapes contains aberrant morphologies shown in Fig. 3 (C and D).
Percent fragmented contains aberrant morphology shown in Fig. 3 E.
Members of the GTPase superfamily have evolved to regulate diverse processes by acting as binary molecular switches. Most GTPases contain a single nucleotide binding domain, raising the possibility that one of the Gem1p GTPase domains is an evolutionary relic and no longer functions. Our analysis demonstrates that both GTPase domains in a single Gem1p protein are required for activity. Specifically, mutations in the G1 motif (K to A or S to N) of either GTPase I or GTPase II abolish protein function. Thus, Gem1p function requires the nucleotide binding residues of both GTPase domains. Although changing the G2 conserved threonine to alanine dramatically alters Ras activity in vivo, similar mutations have no effect in GTPase I (Gem1p(T33A)), making it unlikely that this threonine residue mediates effector interactions required for maintenance of mitochondrial morphology. Moderate effects on Gem1p function were observed when a similar mutation was made in GTPase II (T480A; Table IV). Studies are in progress to characterize the nucleotide binding and hydrolysis activities of both Gem1p GTPase domains and identify potential Gem1p regulators.

EF-hand motifs have been shown to bind calcium, inducing a conformational change that alters protein activity (Yap et al., 1999; Lewit-Bentley and Rety, 2000; Ikura et al., 2002). Mutation of Gem1p EF-hand motifs demonstrated that conserved residues required for calcium coordination are important for Gem1p function (Table IV). Like other multiple EF-hand motif proteins (Janssens et al., 2003), mutation of both motifs produces a stronger phenotype than mutation of either alone. Whether calcium binding to Gem1p modulates protein–protein interactions, the GTPase cycle of either or both GTPase domains, or some other Gem1p function is not clear.

Several observations suggest that Gem1p is not an essential component of previously characterized pathways that control mitochondrial shape. Two processes, division and fusion, have been shown to control mitochondrial network morphology in fungi, invertebrates, and mammals. In tissue culture cells, overexpression of a hMiro-1 protein containing a putative activating mutation led to mitochondrial aggregation that could, in principle, result from decreased division and/or increased fusion (Fransson et al., 2003). Here, we demonstrate that mitochondrial fusion (Table II) and division (Table III) occur in gem1Δ cells, indicating that Gem1p is not a core component of the mitochondrial division or fusion machinery.

Studies have defined another yeast pathway that regulates mitochondrial shape. This pathway requires the function of four proteins: Mmm1p (Burgess et al., 1994), Mmm2p (Youngman et al., 2004), Mdm10p (Sogo and Yaffe, 1994), and Mdm12p (Berger et al., 1997). Mmm1p, Mdm10p, and Mdm12p have been shown to interact with each other, forming a complex on the outer mitochondrial membrane (Boldogh et al., 2003). Mmm2p is essential for formation of the Mmm1p-containing complex (Youngman et al., 2004), suggesting that all four proteins act in the same pathway. Additional studies indicate that this Mmm1p-containing complex is necessary for mtDNA replication and maintenance and spans both mitochondrial membranes (Burgess et al., 1994; Sogo and Yaffe, 1994; Berger et al., 1997; Aiken-Hobbs et al., 2001; Kondo-Omamoto et al., 2003; Meeusen and Nunnari, 2003).

When grown on dextrose medium, mutants lacking Mmm1p, Mdm10p, Mdm12p, or Mmm2p contain enlarged, spherical mitochondria somewhat similar to the globular organelles observed in 54% of dextrose-grown gem1Δ cells (Fig. 3). Although these findings suggest that Gem1p may be required for the function of these four proteins, several lines of evidence argue against this possibility. First, large spherical mitochondria occur in mmm1Δ, mmm2Δ, mdm10Δ, and mdm12Δ cells grown in a variety of media (Burgess et al., 1994; Sogo and Yaffe, 1994; Berger et al., 1997; Youngman et al., 2004). In contrast, mitochondria are not spherical or globular in most gem1Δ cells grown in synthetic glycerol medium. Instead, gem1Δ cells contain short, tubular mitochondria (unpublished data). Second, Mmm1p-GFP localizes properly in cells lacking Gem1p (Fig. 9D). Because Mmm2p, Mdm10p, and Mdm12p are required for proper Mmm1p lo-

### Table IV. Conserved GTPase domains and EF-hand motifs of Gem1p are required for function

| gem1Δ strain containing pRS416 plus: | Percent tubular network | Percent collapsed | Percent mutant |
|--------------------------------------|-------------------------|------------------|---------------|
| No insert                            | 0.9 ± 1.3               | 15.8 ± 3.4       | 83.3 ± 4.3    |
| GEM1                                 | 59.9 ± 6.2              | 25.4 ± 5.7       | 14.6 ± 7.1    |
| GTPase I domain mutations:           |                         |                  |               |
| gem1 (K18A)                          | 3.1 ± 1.3               | 21.1 ± 3.5       | 75.8 ± 3.2    |
| gem1 (S19N)                          | 2.3 ± 0.9               | 18.6 ± 7.8       | 79.1 ± 7.4    |
| gem1 (T33A)                          | 57.8 ± 6.1              | 28.5 ± 4.7       | 13.7 ± 5.4    |
| GTPase II domain mutations:          |                         |                  |               |
| gem1 (K461A)                         | 13.1 ± 1.8              | 31.6 ± 13.8      | 55.3 ± 12.0   |
| gem1 (S462N)                         | 3.2 ± 2.6               | 16.7 ± 1.6       | 80.1 ± 2.4    |
| gem1 (T480A)                         | 13.9 ± 11.7             | 37.9 ± 10.3      | 48.3 ± 5.9    |
| EF-hand motif mutations:             |                         |                  |               |
| gem1 (E225K)                         | 28.7 ± 2.9              | 33.8 ± 2.4       | 37.5 ± 2.2    |
| gem1 (E354K)                         | 42.2 ± 3.5              | 30.7 ± 7.6       | 27.0 ± 10.4   |
| gem1 (E225K, E354K)                  | 14.3 ± 6.7              | 30.3 ± 4.9       | 55.3 ± 7.0    |

n_sub > 300.
Mitochondria were visualized by expressing mito-GFP.
Percent tubular network contains wild-type morphology shown in Fig. 3 A.
Percent collapsed contains aberrant morphology shown in Fig. 3 B.
Percent mutant category contains aberrant morphologies shown in Fig. 3 C–E.
calization on mitochondria (Boldogh et al., 2003; Kondo-Okamoto et al., 2003; Youngman et al., 2004), it seems likely that all four proteins function normally in gem1Δ cells. Third, steady-state levels of Mnm1p and Mmmp2 are indistinguishable between GEM1 and gem1Δ strains (unpublished data). Fourth, when grown on dextrose medium, most gem1Δ, mmmΔ2, mdm10Δ, and mdm12Δ strains exhibit significant mtDNA instability and lose mtDNA (Berger et al., 1997; Youngman et al., 2004). In contrast, DAPI staining of cells grown for at least 35 generations in dextrose-containing media revealed that 43% of gem1Δ cells maintain visible mtDNA (Fig. S1). Fifth, although deletion of MMT1 causes abnormal inner mitochondrial membrane structure (Aiken-Hobbs et al., 2001), gem1 null mutants grown in glycerol- or dextrose-containing media maintain inner membrane cristae (Fig. 4 and not depicted). Together, these observations suggest that Gem1p is not essential for the function of Mmm1p-containing complexes.

Mitochondrial attachment to actin filaments is also required for normal mitochondrial morphology and transport of the organelle from mother to bud during cell division (Drubin et al., 1993; Lazzarino et al., 1994; Hermann et al., 1997; Boldogh et al., 1998, 2003). However, gem1Δ cells do not exhibit defects in mitochondrial inheritance (Fig. 5), suggesting that mitochondrial–actin interactions remain intact in this mutant. In addition, Gem1p does not appear to regulate actin organization as actin cables and patches are normal in gem1Δ mutants.

Although Fransson et al. (2003) suggested that a mutant Miro-1 protein had a pro-apoptotic function in tissue culture cells, whether Miro-1 was required to induce cell death was not tested. Here, we showed that Gem1p is not required for pheromone induced cell death in yeast. Although it is formally possible that Gem1p participates in other forms of yeast cell death (Madeo et al., 2002; Wysocki and Kron, 2004), our combined results support a primary role for Gem1p in mitochondrial morphological maintenance.

Gem1p does not appear to be essential for any of the known pathways controlling mitochondrial morphology in yeast. The localization and topology of Gem1p suggest that it may sense cytoplasmic signals (e.g., cytosolic calcium, other signaling molecules) and directly or indirectly alter mitochondrial morphology. Several observations support the notion that Gem1p is a regulatory molecule. First, a large portion of Gem1p is devoted to domains that act as molecular switches in other proteins, and these domains are required for Gem1p function. Second, unlike mutants which disrupt division, fusion, or the Mnm1p morphology pathway, gem1Δ cells exhibit pleiotropic mitochondrial morphologies (Fig. 3). Third, changing the cellular environment by altering strain background or carbon source alters the severity of the gem1Δ mitochondrial phenotype. One attractive hypothesis currently being tested is that Gem1p regulates mitochondrial morphological pathways in response to changes in cytosolic calcium levels. Identification of protein binding partners for Gem1p is in progress and will likely reveal the pathways in which Gem1p acts.

Materials and methods

Strains and plasmid constructions

Yeast strains were constructed in the W303 background. Standard methods were used to manipulate yeast (Sherman et al., 1986; Guthrie and Fink, 1991) and bacterial (Maniatis et al., 1982) strains. All mutations, disruptions, and constructs were confirmed by PCR, DNA sequencing, and, where appropriate, Western blotting.

A complete disruption of YAL048C (GEM1) coding region was accomplished by homologous recombination using a HIS3 cassette (Rothstein, 1991) generating strain JSY6592 (MAT ade2-1 leu2-3 his3-11,15 trp1-1 ura3-1 can1-100 gem1::HIS3). A backcross of this strain to W303 wild type generated the wild-type JSY7000 (MAT ade2-1 leu2-3 his3-11,15 trp1-1 ura3-1 can1-100 gem1::HIS3) and JSY7002 (MAT ade2-1 leu2-3 his3-11,15 trp1-1 ura3-1 can1-100 gem1::HIS3). Double mutants of gem1 and other genes were generated by mating, sporulation, and dissection. To generate JSY77234 (MAT a ade2-1 leu2-3 his3-11 trp1-1 ura3-1 can1-100 gem1::HIS3 dnm1::HIS3) and JSY77236 (MAT ade2-1 leu2-3 his3-11 trp1-1 ura3-1 can1-100 gem1::HIS3 mcm7::TRP1) and JSY5692 was crossed to JSYS158 (MAT ade2-1 leu2-3 his3-11 trp1-1 ura3-1 can1-100 dnm1::HIS3). To make JSYS766 (MAT ade2-1 leu2-3 his3-11 trp1-1 ura3-1 can1-100 gem1::HIS3 fzo1::TRP1) and JSYS7768 (MAT ade2-1 leu2-3 his3-11,15 trp1-1 ura3-1 can1-100 gem1::HIS3 fzo1::TRP1), a PCR fragment encoding GFP and Flanked by HindIII and EcoRI was cloned into pYX142. GFP fusions in this vector are driven by the constitutively active TPI promoter. For pYX142-GFP-GEM1 and pYX142-GFP-GEM1(aa618-662), a PCR fragment cloned by HindIII and SacI sites was cloned into pYX142-GFP.

Plasmids for analysis of specific domain mutations were generated by site-directed mutagenesis (Stratagene) to generate pRS416-GEM1. For pYX142-GFP, a PCR fragment encoding GFP and Flanked by HindIII and EcoRI was cloned into pYX142. GFP fusions in this vector are driven by the constitutively active TPI promoter. For pYX142-GFP-GEM1 and pYX142-GFP-GEM1(aa618-662), a PCR fragment cloned by HindIII and SacI sites was cloned into pYX142-GFP.

For serial dilutions, strains JSY7000 GEM1 or JSY7002 gem1Δ containing the indicated plasmid were grown in SDextrose dropout medium to early log phase (OD600 0.5–1.0), pelleted, and resuspended to OD600 0.5. Aliquots of 1:5 serial dilutions were spotted onto SDextrose or Dextrose containing media revealed that 43% of wild-type and mutant strains. Double

Analytical methods

Yeast strains were constructed in the W303 background. Standard methods were used to manipulate yeast (Sherman et al., 1986; Guthrie and Fink, 1991) and bacterial (Maniatis et al., 1982) strains. All mutations, disruptions, and constructs were confirmed by PCR, DNA sequencing, and, where appropriate, Western blotting.

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For serial dilutions, strains JSY7000 GEM1 or JSY7002 gem1Δ containing the indicated plasmid were grown in SDextrose dropout medium to early log phase (OD600 0.5–1.0), pelleted, and resuspended to OD600 0.5. Aliquots of 1:5 serial dilutions were spotted onto SDextrose or SGlycerol medium lacking the appropriate amino acids and grown for 3 d at the indicated temperatures.

Analysis of mitochondrial morphology, mitochondrial fusion, and mtDNA maintenance, and actin cytoskeleton

Mitochondrial morphology was scored at 30°C in wild-type and mutant cells expressing a matrix targeted form of mito-GFP (pYX142-mtGFP::1) or mito-RFP (p[AD4GPD::mtRFP]). Yeast were grown in SDextrose or Dextrose containing medium, unless otherwise indicated. In some cases, mitochondria were labeled with Mitotracker red 589 (Molecular Probes, Inc.). Phenotypes were quantified in at least 100 cells in three or more independent experiments. Data reported are the average and SD of all experiments. Phenotypes were quantified in at least 100 cells in three or more independent experiments. Data reported are the average and SD of all experiments. Phenotypes were quantified in at least 100 cells in three or more independent experiments. Data reported are the average and SD of all experiments.
Subcellular and submitochondrial localization of GFP-Gem1p
JSY7000 cells containing pYX142-GFP-GEM1 were grown in lactate medium to log phase (OD600 0.6–1.0), spheroplasted, lysed, and subjected to differential sedimentation to generate an enriched mitochondrial fraction as described previously (Kondo-Okamoto et al., 2003). WCE, PMS, and MTO fractions were separated by SDS-PAGE and analyzed by Western blotting with antibodies specific for Porin (Molecular Probes, 1:8,000), 3-PGK (Molecular Probes, 1:1,000), and GFP (Covance, Inc., 1:500). HRP-conjugated secondary antibody (goat α mouse; Sigma-Aldrich, 1:10,000; goat α rabbit; Jackson ImmunoResearch Laboratories, 1:10,000) and ECL detection were used to visualize bands. MTO (30 μg protein were loaded) were analyzed by 10-fold more (cell equivalents) than WCE and PMS.

Mitochondria (200 μg protein) were analyzed by protease protection as described previously (Kondo-Okamoto et al., 2003). Samples were subjected to PK treatment, osmotic shock and PK treatment, detergent solubilization and PK treatment, or a mock treatment. Samples were denatured and analyzed by SDS-PAGE and Western blotting (one quarter of total reaction per lane). To perform carbonate extraction, mitochondria were subjected to PK treatment, osmotic shock and PK treatment, detergent solubilization, and observed using bright field microscopy. Results shown are the averages of two independent experiments in which n = 200 for each treatment.

Online supplemental material
mtDNA maintenance during growth in dextrose-containing medium is depicted in Fig. S1. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200405100/DC1.

We greatly appreciate discussions with all members of the Shaw lab. We thank W. Neupert, Munich, Germany, for a gift of Mdm12p, a component required for mitochondrial inheritance that is conserved between budding and fission yeast. We thank Nousheen Alasti for constructing pRS415-α-GFP-α-GEM1. We thank Nousheen Alasti for constructing pRS415-α-GFP-α-GEM1.

Submitted: 17 May 2004
Accepted: 31 August 2004

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