RESEARCH ARTICLE

Single amino acid mutations in the *Saccharomyces cerevisiae* rhomboid peptidase, Pcp1p, alter mitochondrial morphology

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Abstract

Key to mitochondrial activities is the maintenance of mitochondrial morphology, specifically cristae structures formed by the invagination of the inner membrane that are enriched in proteins of the electron transport chain. In *Saccharomyces cerevisiae*, these cristae folds are a result of the membrane fusion activities of Mgm1p and the membrane-bending properties of adenosine triphosphate (ATP) synthase oligomerization. An additional protein linked to mitochondrial morphology is Pcp1p, a serine protease responsible for the proteolytic processing of Mgm1p. Here, we have used hydroxylamine-based random mutagenesis to identify amino acids important for Pcp1p peptidase activity. Using this approach we have isolated five single amino acid mutants that exhibit respiratory growth defects that correlate with loss of mitochondrial genome stability. Reduced Pcp1p protease activity was confirmed by immunoblotting with the accumulation of improperly processed Mgm1p. Ultra-structural analysis of mitochondrial morphology in these mutants found a varying degree of defects in cristae organization. However, not all of the mutants presented with decreased ATP synthase complex assembly as determined by blue native polyacrylamide gel electrophoresis. Together, these data suggest that there is a threshold level of processed Mgm1p required to maintain ATP synthase super-complex assembly and mitochondrial cristae organization.

Keywords: ATP synthase; blue native-PAGE; cristae structure; mitochondrial morphology; Pcp1p; peptidase activity

Introduction

In eukaryotic cells, mitochondria are important organelles involved in the production of energy in the form of adenosine triphosphate (ATP), apoptosis, and calcium regulation (Chandel, 2014). The maintenance of mitochondria morphology is integral to these functions (Sun et al., 2007; Cogliati et al., 2013). The structure of mitochondria in wild-type *Saccharomyces cerevisiae* cells is that of a tubular network that extends around the cell periphery (Nunnari et al., 1997; Rafelski et al., 2012). The ultra-structure of mitochondria generally consists of an outer membrane, an inner membrane that folds to form cristae, an intermembrane space, and the matrix. The inner membrane comes in close contact with the outer membrane before folding into cristae, with the area at the base of the cristae fold known as a cristae junction (Mannella et al., 2008; Zick et al., 2009b). A number of proteins embedded in the inner membrane are key to maintaining mitochondrial morphology via regulated fusion and fission-based events. Fusion of the outer and inner membranes with neighboring mitochondria forms the reticular network found in normal yeast cells and fission allows for the partitioning of mitochondria into daughter cells during cellular division (Nunnari et al., 1997). A balance between fusion and fission events is required for the maintenance of mitochondrial structure. For example, cells lacking expression of Dnm1p, a protein responsible for fission of the outer membrane, results in a large, branched mitochondrial network. In contrast, deletion of a protein involved in outer membrane fusion, Fzo1, results in fragmented mitochondria (Sesaki and Jensen, 1999). However, these defects in morphology can be overcome if the opposing activity is removed, as evidenced by the double Δfzo1 Δdnm1 cells containing normal mitochondria (Sesaki and Jensen, 1999). Fusion
and fission of the inner membrane can also direct cristae formation. Under the stress of induced matrix swelling, fusion of the inner membrane results in very few cristae junctions and large intra-cristae spaces. Upon retraction, most cristae returned to their normal state while some mitochondria contained inner membrane vesicles that are hypothesized to result from fission of those large cristae structures observed during swelling (Mannella et al., 2008). Cristae structures have also been shown to be lost during the progression of apoptosis (Sun et al., 2007).

In S. cerevisiae, Pcp1p is a multispanning transmembrane protein of the mitochondrial inner membrane. As a serine protease, Pcp1p cleaves substrate proteins within the membrane environment, an activity linked to regulating mitochondrial structure (Herlan et al., 2003; McQuibban et al., 2003; Sesaki et al., 2003a). Pcp1p has two known substrates: cytochrome c peroxidase, or Ccp1p (Esser et al., 2002; Michaelis et al., 2005), and the dynamin-like GTPase, Mgm1p (Wong et al., 2000; Herlan et al., 2003; McQuibban et al., 2003; Wong et al., 2003; Sesaki et al., 2003a). Ccp1p, functions in the reduction of cytochrome c to oxidize H₂O₂ (Yonetani and Ohnishi, 1966) which is toxic to cells and can build up as a by-product of oxidative phosphorylation. The second substrate, Mgm1p, has been shown to be required for mitochondrial fusion (Wong et al., 2000, 2003) as cells deleted for MGM1 give rise to fragmented mitochondria, lose mitochondrial DNA (mtDNA) (Guan et al., 1993; Herlan et al., 2004), and lack cristae ultrastructure (Sesaki et al., 2003b; Amutha et al., 2004). Deletion of PCP1 also leads to fragmented mitochondria, mitochondria absent of cristae, loss of mtDNA, as well as the inability to grow on nonfermentable carbon sources (McQuibban et al., 2003; Sesaki et al., 2003a) all of which is a result of the inability of these cells to process Mgm1p.

ATP synthase is another protein located within the mitochondria inner membrane with a role in mitochondrial cristae maintenance. This protein complex uses the proton gradient generated by electron transport chain activity for ATP generation. It is composed of a catalytic F₁ “head” found on the matrix side of the inner membrane and a F₀ stalk that is embedded in the inner membrane. The majority of the ATP synthase subunits are encoded by nuclear DNA with the exception of three F₀ subunits (ATP6, ATP8, and ATP9), which are encoded on the mitochondrial genome (Ackerman and Tzagoloff, 2005). The ATP synthase complex contains several proteins that aid in the association between the F₀ and F₁ complexes as well as those that promote the formation of ATP synthase dimers and higher-order oligomers (Arnold et al., 1998; Arselin et al., 2003; Amutha et al., 2004; Wagner et al., 2010). Recent work has shown that the formation of the ATP synthase super-complex is crucial to cristae structure (Arselin et al., 2003; Thomas et al., 2008; Davies et al., 2012). Tim11p, or subunit e of ATP synthase, has been shown to be one of the components required for ATP synthase dimerization (Arnold et al., 1997; Arselin et al., 2003; Everard-Gigot et al., 2005). Deletion of TIM11 leads to fragmented mitochondria that display “onion-like” cristae and an increased tendency to lose mtDNA (Giraud et al., 2002; Paumard et al., 2002; Arselin et al., 2003). F₁F₀ ATP synthase monomers are linked together through Tim11p at the base of the stalk. These dimers then assemble to form higher-order oligomers, which promote inner membrane bending to form the rim of the cristae (Arselin et al., 2003; Thomas et al., 2008; Davies et al., 2012). The observation that Tim11p is not detectable in mgm1 or pcp1 deletion mutants (Amutha et al., 2004) suggests a possible connection between mitochondria structure and function influenced directly or indirectly by Pcp1p activity.

In this study, we have generated pcp1 mutants that contain single amino acid changes that result in a range of catalytic activities in vivo. It was determined by transmission electron microscopy (TEM) and blue native polyacrylamide gel electrophoresis (BN-PAGE) that the ability of each mutant to process Mgm1p correlates with mitochondrial architecture but not necessarily ATP synthase oligomerization. Given the established link between ATP synthase complex assembly and mitochondrial cristae formation, these mutants can be used to dissect the relationship among Pcp1p peptidase activity, Mgm1p-mediated inner membrane fusion, and mitochondrial architecture independent of ATP synthase.

Materials and methods

Strains, media, and reagents

All yeast strains used in this study were derived from BY4741 (Mata his3Δ1 leu2Δ0 met15Δ0 ura3Δ0) (Thermo Fisher Scientific). A pcp1 shuffle strain was constructed by targeted integration, in which the PCP1 open-reading frame was disrupted with the introduction of LEU2 (pRS405) as described (Rothstein, 1991). Correct integration was confirmed by genomic polymerase chain reaction (PCR) and the resulting diploid transformed with a URA3-marked PCP1 plasmid, sporulated, and micro-dissected. Covered ypcp1::LEU2 haploid mutants were identified by growth on uracil and leucine-deficient selective plates.

An epitope-tagged versions of MGM1 were generated by homologous recombination using a PCR-based approach as described (Longtine et al., 1998). Transformants capable of growth on 0.2 g/L G418 YPD plates were confirmed by genomic PCR analysis.

Yeast media was prepared according to the methods previously described (Sherman, 1991). To ensure retention
of a HIS3-marked plasmid, strains were grown in supplemented 0.67% yeast nitrogen base lacking histidine (-His) and containing 2% glucose, 2% raffinose, or 2% ethanol and 3% glycerol as a carbon source. The \( \Delta \text{tim11} \) strain was grown in YP media (2% peptone, 1% yeast extract, 2% carbon source). Strains for TEM analysis were grown in histidine-selective media containing 2% raffinose.

**pcp1 mutant generation**

The *pcp1* mutants were generated by random mutagenesis of a plasmid-based copy of the *PCP1* open-reading frame under the endogenous *PCP1* promoter in the HIS3-marked pRS413 vector. Nearly, 500 ng of purified plasmid (DG295) was treated with 1 M hydroxyamine hydrochloride in 0.45 M sodium hydroxide for 20 and 25 h at 37°C (Rose and Fink, 1987). The reaction was stopped with the purification of the plasmid using the Qiagen miniprep kit according to the manufacturer’s instructions. The resulting pool of mutagenized plasmid was transformed into the *PCP1* shuffle strain (DGY43) (Schiestl and Gietz, 1989). For identification of mutants, the covering wild-type *PCP1* plasmid was removed by 3 days of growth on 5-fluoroorotic acid (5-FOA)-containing plates (Sherman, 1991). Loss of the *PCP1* URA3-marked plasmid was confirmed as no growth on Uracl plates. Defects in Pcp1p activity were selected as poor growth after replica printing to ethanol/glycerol plates. Plasmids were isolated from the cultures grown overnight at 30°C in 5 mL of selective media as follows. Cells isolated by centrifugation were pretreated in 100 mM Tris pH 9.0 for 10 min at 30°C prior to spheroplast formation in spheroplast buffer (1 M sorbitol, 20 mM Tris pH 7.5, 10 mM ethylenediaminetetraacetic acid (EDTA) pH 8.0; 0.15 µg/µL 100 T zymolyase, and 0.2% β mercapto-ethanol) for 1 h at 30°C. Spheroplasts were isolated by centrifugation at 2,000g for 10 min, resuspended Qiagen P1 buffer, and subjected to two freeze-thaw cycles. Spheroplasts were mechanically lysed by vortexing with glass bead. Qiagen P2 buffer was added and incubated at room temperature for 5 min. Qiagen N3 buffer was added and the lysate clarified by centrifugation at 21,130g for 10 min. Plasmid DNA was isolated by isopropanol precipitation and used to transform NEB Turbo competent *Escherichia coli* (New England Biolabs). Nucleotide mutations were identified by Sanger sequencing using four primer pair sets that covered the *PCP1* promoter and open-reading frame (Arizona State University, CLAS DNA Laboratory). Following sequencing, all mutants were found to contain a common T962C nucleotide change that resulted in a Valine to Alanine mutation at amino acid position 321. This mutation was also found in the original plasmid and was determined to have no impact on Pcp1p activities (data not shown).

When possible, a site-directed mutagenesis approach was used to correct this nucleotide mutation (QuikChange II; Agilent).

**Serial dilution spotting assay**

Nearly \( 10^7 \) cells from exponentially growing cultures were isolated by centrifugation and the resulting cell pellet was resuspended at \( 10^5 \) cells/µL in sterile water. Cells were placed in the first column of a 96-well microtiter plate and serially diluted 10-fold by transferring 20 µl cells to 180 µl media. Three microliters from each well were spotted onto selective plates and growth recorded after 3 days at 30°C.

**Petite frequency determination**

For each strain, a volume of overnight culture equivalent to \( 10^7 \) cells was brought up to 1 mL in selective glucose media and serially diluted to achieve 1,000 cells/mL. Multiple 100 µL aliquots were plated onto -His glucose plates. After 3 days at 30°C, the numbers of colonies on each plate were counted and the colonies replica printed onto -His ethanol/glycerol plates. Colonies that grew on -His ethanol/glycerol plates were counted after an additional 3 days of growth at 30°C. This process was independently repeated five times for each strain. The petite frequency was calculated using the following equation: (colony number on glucose–the colony number on ethanol/glycerol)/colony number on glucose × 100.

**mtDNA detection by 4′,6-diamidino-2-phenylindole (DAPI) staining**

The cultures were grown overnight in a selective media at 30°C and fixed in 3.7% formaldehyde for 2 h at room temperature. The cells were isolated by centrifugation, washed with sterile distilled water, and resuspended in 50 mM phosphate-buffered saline (PBS) pH 7.4. Permeabilization was carried out in 70% ethanol on ice for 30 min, cells isolated by centrifugation, and the cell pellet resuspended in PBS. DAPI was added at 50 µg/mL for 5 min on ice. DAPI-stained cells were spotted onto a glass microscope slide and viewed under 100× oil immersion on a Nikon Eclipse-50 microscope. Images of at least 200 cells from each strain were captured using the same exposure settings with QCapture Pro software. This process was independently repeated three times for each strain. The percentages of cells containing mtDNA were determined, with a cell containing any extranuclear DAPI staining scored as containing mtDNA.

**TEM**

The TEM protocol was compiled using suggestions from Wright (2000). The *pcp1* strains were grown in 15 mL -His...
2% raffinose media at 30°C overnight with shaking to reach an OD600 between 0.5 and 1. The cells were pelleted at 1,500g for 10 min at 18°C and the cell pellet resuspended in 15 mL YP 2% raffinose and returned to 30°C with continuous shaking for 1 h. Each culture was fixed with the addition of an equal volume of 4% gluteraldehyde in 2× piperazine-N,N′-bis(2-ethanesulfonic acid) (PIPES) buffer (0.2 M PIPES pH 6.8, 0.2 M sorbitol, 2 mM MgCl2, 2 mM CaCl2) for 5 min at room temperature followed by 2% gluteraldehyde in 1× PIPES buffer overnight at 4°C. Fixed cells were incubated in 2% potassium permanganate, washed with water, and dehydrated with a series of graded ethanol washes (35% ethanol, 50% ethanol, and 70% ethanol). The cells were incubated en bloc in 1% uranyl acetate in 70% ethanol overnight. Next day, the cells were twice washed for 15 min with 95% ethanol followed by four 15 min washes in 100% ethanol.

The cells were gradually infiltrated with resin replacing ethanol with acetone using 25% Spurr’s Resin in 75% acetone, 50% Spurr’s Resin in 50% acetone, and 75% Spurr’s in 25% acetone overnight. Infiltration was completed in 100% resin for 24 h. The following day, cells were embedded in fresh Spurr’s Resin in a 68–70°C oven overnight. The blocks were trimmed, sectioned, and placed onto copper mesh grids for staining in 2% alcoholic uranyl acetate followed by lead citrate. TEM was performed on a JEOL JEM-1230 electron microscope at 80 kV, 60 μA, 140–180pA/cm2 using spot size three at a magnification of ×6,000–8,000 to image whole cells and ×12,000 to visualize individual mitochondria. Approximately 30 cells were photographed for each of the pcp1 mutants.

Mitochondria isolation

The cells (2 L) were grown overnight in the appropriate selective media containing 2% raffinose at 30°C with continuous shaking. Mitochondria were isolated following published procedures as described (Murakami et al., 1988). The final mitochondrial pellet was resuspended in 1× buffer A (20 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid [HEPES] pH 7.5, 0.6 M sorbitol, 10 mM trisylol, 1 mM p-aminobenzamidine, 0.1% bovine serum albumin, 1 mM phenylmethylsulfonylfluoride) and the mitochondrial protein content determined spectrophotometrically. Isolated mitochondria were diluted with an equal volume of 1× buffer A containing 20% dimethyl sulfoxide, aliquoted in 1–2 mg volumes, and stored at −80°C.

Sodium dodecyl sulfate PAGE (SDS-PAGE)

Isolated mitochondria were washed with 20 mM HEPES pH 7.4, 0.6 M sorbitol buffer and centrifuged at 20,400g for 2 min at 4°C. The mitochondrial pellets were resuspended in 20 mM HEPES pH 7.4, 0.6 M sorbitol buffer to achieve a concentration of 10 mg/mL. One hundred micrograms was added to 1× SDS loading buffer (300 mM Tris base, 4% SDS, 17% glycerol, 0.007% bromophenol blue, 90 mM dithiothreitol), sonicated, and heated at 55°C for 5 min. Proteins were separated by electrophoresis through a 12% or 15% discontinuous SDS polyacrylamide gel.

BN-PAGE

The following BN-PAGE protocol was modified based on various sources (Schägger and von Jagow, 1991; Schägger et al., 1994; Dekker et al., 1996; Wagner et al., 2010). One microgram of isolated mitochondria from selected strains was washed with 1 mL 20 mM HEPES pH 7.4, 0.6 M sorbitol buffer and centrifuged at 20,400g for 2 min at 4°C. Mitochondrial pellets were lysed in a 0.5% digitonin buffer (0.5% digitonin, 20 mM Tris-HCl pH 7.5, 1 mM EDTA, 50 mM NaCl, 10% glycerol, 1 mM phenylmethylsulfonylfuoride) to obtain a protein concentration of 4 mg/mL at a detergent to protein ratio of 1:4:1 (g/g). Different detergent to protein ratios were obtained by varying the percent weight by volume of digitonin used in the lysis buffer (0.5% digitonin = 1:4:1 g, 0.25% digitonin = 0.7:1 g). A soluble protein fraction was then isolated by centrifugation at 100,000g for 30 min at 4°C. 5.5 μL of 10× loading buffer (5% Coomassie Blue G, 500 mM 6-aminoacaproic acid, 100 mM Bis-Tris pH 7.0) was added to 200 μg of protein. After an additional spin at 20,400g for 2 min at 4°C, samples were either frozen at −20°C or immediately used. Samples were loaded onto a 3–13% blue native polyacrylamide gel and run at 4°C in 0.02% Coomassie Blue G-250, 50 mM tricine, 15 mM Bis-Tris pH 7.0 cathode buffer at 100 V for ~30 min, 160 V for about 3 h, and then 160 V in cathode buffer without Coomassie Blue G-250 for an additional 5 h. The resulting BN-PAGE was either used directly for determining ATP synthase activity using an ingel activity assay or transferred to polyvinylidene fluoride (PVDF) membrane for immunoblotting.

Immunoblotting

The immunoblot analysis was carried out following established procedures (Harlow and Lane, 1988). BN gels were transferred to PVDF membrane, while SDS gels were transferred to nitrocellulose. Primary antibodies were used at the following dilutions: ATP-V α subunit at 1:5,000 (SDS) or 1:10,000 (BN-PAGE) (Cat# 459240, RRID: AB_2532234; Thermo Fisher Scientific); Hemagglutinin at 1:5,000 (Cat# MMS-101R, Antibody Registry: AB_291263; Covance); Ccp1p, Tom40p, and Tim11p at 1:5,000 (gift from Debkumar Pain, Rutgers University, NJ). Membranes were developed using horseradish peroxidase-conjugated secondary antibodies (Cat# NA9310 and NA934, Antibody Registry: AB_772193 and AB_772206; GE Healthcare) and
SuperSignal West Femto Chemiluminescence reagents (Thermo Fisher Scientific).

In-gel ATP synthase activity assay

Freshly processed mitochondrial proteins were separated by electrophoresis through a BN-PAGE, and the resulting gel was incubated with 20 mL ATP buffer (50 mM glycine-NaOH pH 8.4, 5 mM MgCl₂, 20 mM ATP) for 20 min at room temperature without shaking followed by 20 mL 10% CaCl₂ as described (Bornhövd et al., 2006). The positions of calcium phosphate precipitates were marked, a photo taken, and the gel incubated in destain solution (64% methanol, 3.1 M glacial acetic acid) overnight to visualize the protein standards.

Data analysis

The relative differences in protein levels were determined using ImageJ software with s-Mgm1p compared with Pgk1p and the ATP synthase α subunit and Tim11p levels compared to Tom40p. The resulting ratios were then normalized to that obtained for PCP1. Mitochondria cristae was quantified based on the ultrastructure present in TEM images. Area measurements for each mitochondrion were acquired using ImageJ software. The subsequent areas where binned into 20 nm² groupings for analysis. Statistical significance was calculated from measurements using a two-tailed T test assuming equal variances. The null hypothesis assumed the means from each pcp1 strain was equal to the wild-type (PCP1) mean with a 95% confidence level (α = 0.05).

Data obtained for mtDNA staining and Mgm1p processing was analyzed using a one-way analysis of variance with a post hoc Tukey’s HSD test with statistical differences having a P < 0.01.

Results

pcp1 mutants exhibit a range of growth defects

We took advantage of the finding that yeast cells deleted for PCP1 are unable to use nonfermentable carbon to sustain growth to identify alleles of PCP1 that resulted in reduced function (Sesaki et al., 2003a). To this end, a haploid pcp1 deletion strain transformed with a library of hydroxylamine mutagenized PCP1 plasmid was screened for colonies that exhibited poor growth on ethanol/glycerol containing selective plates. After multiple rounds of screening, plasmid isolation, and retesting, five mutants were selected for further analysis (Figure SAI1A). These mutants displayed a varying degree of growth defects and were found to contain nucleotide mutations that altered 1–2 amino acids in the PCP1 open-reading frame. All mutants are referred to by single-letter code along with their numeric amino acid position and include: Pcp1p (G120D), Pcp1p (G233D), Pcp1p (S252N), Pcp1p (G315D), and Pcp1p (G233S V321A). These mutations are found in highly conserved regions of the protein (Figure SAI1C) and are predicted to map to transmembrane helices (TMH) 1, 4, and 6, as well as a loop domain between helices 3 and 4 in the mature Pcp1 protein (Figure SAI1B).

As an initial approach to compare the impact each pcp1 mutation had on overall growth, a spotting assay was carried out. Variable growth defects were identified for the mutants under nonselective (glucose) and selective (ethanol-glycerol) testing conditions (Figure 1A). Least impaired was the Pcp1p (G233S V321A) mutant, followed by Pcp1p (G120R) and Pcp1p (S252N). The Pcp1p (G233D) mutant was found to have poor growth on ethanol-glycerol. The G315D mutant exhibited a growth pattern most similar to that of the Δpcp1 mutant, with reduced growth on glucose and no growth on ethanol-glycerol-containing plates suggesting that the mutant is likely a null allele. However, immunoblot analysis confirmed that Pcp1 protein was present in the G315D mutant, although at a reduced level compared with other mutants (Figure SAI2).

Yeast having lost the ability to utilize nonfermentable carbon for growth are referred to as petites (Day, 2013). To quantify the differences in growth defects seen for pcp1 mutants on ethanol-glycerol, the percentage of petites in an exponentially growing culture was determined using a replica printing approach. Similar to that seen by spotting, cultures of Δpcp1 and Pcp1p (G315D) were found to be 100% petite, while cultures of Pcp1p (G233S V321A) had a petite frequency similar to wild-type at ~2–3% (Figure 1B). The three remaining mutants, Pcp1p (S252N), Pcp1p (G120R), and Pcp1p (G233D), had calculated average petite frequencies of 14%, 59%, and 70%, respectively. Overall, the calculated petite frequencies matched the increased deficiencies seen for growth on ethanol-glycerol by spotting.

The absence of growth on nonfermentable carbon is often due to the acquisition of deletions in, or complete loss of, the mitochondrial genome (Contamine and Picard, 2000). To determine whether the petite phenotype seen for each mutant was due to the complete loss of the mitochondrial genome, the presence of mtDNA was scored by DAPI staining. The cells in wild type and Pcp1p (G233S V321A) mutant cultures had extranuclear staining in over 93% of the cells, consistent with the majority of the culture retaining respiratory competence (Figure 1C). However, ~75% of the cells expressing Pcp1p (G120R) or Pcp1p (G233D) had staining consistent with mtDNA, suggesting that the petite phenotype in these strains was primarily due to the accumulation of mutations or deletions in the mitochondrial genome. Even cultures of the nonfunctional pcp1 mutants (Δpcp1 and G315D) had some cells that retained mtDNA staining despite all scoring phenotypically...
Figure 1 Reduced growth and loss of respiratory competence found for strains expressing mutant versions of PCP1. (A) Analysis of growth characteristics of haploid pcp1 mutants under nonselective (glucose) and selective (ethanol-glycerol) conditions. A six-point 10-fold serial dilution spotting assay identifies pcp1 mutants that are unable to support growth (Δpcp1 and G315D) or show reduced growth (G233D, S252N, and G120R) under selective conditions compared with PCP1. (B) Quantification of petite frequency reveals a range in respiratory competence. The difference in the number of cells in each culture that were able to grow on nonselective versus selective plates were determined for each strain with the average ± standard deviation reported for three independent experiments. (C) Quantification of 4',6-diamidino-2-phenylindole (DAPI) staining reveals a varying degree of extranuclear DNA staining for all pcp1 mutants. Cells viewed by fluorescence microscopy identified mitochondrial DNA staining that correlated with petite frequency findings. Over 200 cells were counted for each strain with data presented as the average ± standard deviation for three independent experiments. Statistical significance was determined using a one-way analysis of variance (ANOVA) with a post hoc Tukey’s HSD test, *P<0.01.
as petites, data consistent with prior observations for pcp1 deletion mutants (Herlan et al., 2003). The presence of mitochondrial DNA in these mutants were also supported by mitochondria-to-nuclear genome ratio data obtained by quantitative PCR (qPCR) analysis (Table SA1).

**pcp1 mutants are defective in substrate processing**

To identify the impact that each mutation had on Pcp1p activity, the processing of two Pcp1p substrates, Ccp1p and Mgm1p, was determined. In wild-type cells, Ccp1p is processed in two steps with Pcp1p activity catalyzing the final cleavage step to generate the mature form of the enzyme (Esser et al., 2002; Figure 2A, lane 1). In Δpcp1, the second cleavage step does not take place, resulting in the accumulation of an intermediate form of Ccp1p (iCcp1p), and in our hands, a faster migrating form that is likely a degradation product. Of the 5 pcp1 mutants, all retained some level of peptidase activity except for the Pcp1p (G315D) mutant, which primarily accumulated the intermediate form of Ccp1p. In wild-type cells, Mgm1p accumulates as both a long- and short-form (Figure 2B, lane 1) while only the long-form is found in Δpcp1 mutants (Herlan et al., 2003; Figure 2B, lane 2). Similar to that seen for Ccp1p processing, the Pcp1p (G315D) mutant appeared to essentially lack peptidase activity. While Mgm1p processing for the Pcp1p (G233S V321A) mutant was most similar to that of wild type, a significant reduction in the conversion of the long-to-short isofrom of Mgm1p was identified for the three remaining mutants including Pcp1p (S252N), Pcp1p (G233D), and Pcp1p (G120R) (Figure 2C). Despite these differences in Mgm1p processing, no statistically significant differences in the total amount of Mgm1p protein was found. Altogether, the pcp1 mutants exhibited defects in peptidase activity that correlated in severity with their growth patterns on ethanol-glycerol.

**pcp1 mutants have a variable impact on mitochondrial morphology**

A balance between the long and short isoforms of Mgm1p has been implicated in influencing mitochondrial fusion events in yeast (Herlan et al., 2003; Ishihara et al., 2006). Given the defects in Mgm1p processing detected for the pcp1 mutants, they are expected to impact mitochondrial morphology. The cursory analysis with fluorescence microscopy using mitochondrially targeted GFP or MitoTracker Orange CMTMRos found an increase in fragmented mitochondria for all of the Pcp1p mutants (data not shown; Xiao, 2013; Both and Gordon, 2015). To better define defects in mitochondrial architecture, TEM

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**Figure 2** Pcp1p mutants exhibit reduced peptidase activity. (A) Differences in Pcp1p-mediated processing of Ccp1p were determined by immunoblot analysis of protein obtained from mitochondria isolated from the indicated strains. Complete processing of Ccp1p generates a mature form of the protein (mCcp1p) while defects in Pcp1p-mediated cleavage of Ccp1p results in the accumulation of a slower migrating intermediate form (iCcp1p). *Improperly processed form of Ccp1p. Tom40p was used to confirm equal lane loading (bottom panel). (B) Differences in Pcp1p-mediated processing of Mgm1p were determined by immunoblot analysis of total protein extracts isolated from the indicated strains. Mgm1p is present as a long- and short-form (l-Mgm1p and s-Mgm1p, respectively) with the conversion of the long- to short-form mediated by Pcp1p activity. Defects in Pcp1p peptidase activity result in reduced levels of s-Mgm1p and the accumulation of an improperly processed form of the protein (indicated with an asterisk; Schäfer et al., 2010). Pgk1p was used to confirm equal lane loading (bottom panel). The normalized amount of s-Mgm1p present in each mutant relative to Pgk1p is shown (n = 3). (C) The percent each form is of the total Mgm1p detected in each strain was determined following densitometric analysis. Data presented as the average ± standard deviation from three experiments. Values significantly different from PCP1 are indicated. *P<0.05, **P<0.01.
Figure 3 Mitochondrial cristae structure is disrupted in pcp1 mutants as determined by transmission electron microscopy (TEM). A. Representative TEM images of cristae structures found include: (a) normal, (b) normal-vesicular, (c) vesicular, and (d) concentric. Images were taken at x12,000 magnification; scale bars = 500 nm. (B) Distribution of mitochondrial cristae organization identified by TEM analysis for each strain. Graphical representation of cristae morphology found in wild-type (PCP1), the deletion mutant (∆pcp1), and each of the five pcp1 mutant strains. Normal; white box, normal-vesicular; hatched box, vesicular; gray box, concentric; black box.
was used. The ultrastructure for all strains were analyzed and four different cristae categories were identified: normal, normal-vesicular, vesicular, and concentric (Sun et al., 2007) (Figure 3). Normal cristae are defined as those that had little intra-cristae space visible and often appeared to be protruding from the outer membrane as close proximity of the outer and inner membrane make it difficult to distinguish between the two (Figure 3a). Normal cristae were also fairly numerous and appeared to be the textbook example of wild-type mitochondrial morphology.

The vesicular category was used to describe mitochondria, in which the mitochondrial inner membrane formed vesicle-like structures that did not appear to be attached to the outer membrane (Rabl et al., 2009) (Figure 3b). The mitochondria described as having normal-vesicular morphology had cristae with long inner membrane sheets that appeared to divide the mitochondria into sections that contained inner membrane vesicles (Figure 3c). The mitochondrial cristae morphology described as concentric contained mitochondria that contained no visible cristae or cristae that appeared “onion-like” as described in the literature (Paumard et al., 2002) (Figure 3d). Concentric cristae also contained almost no cristae junctions. All of the pcp1 mutant strains contained multiple cristae types and many of the strains often contained a single cell with mitochondria having more than one cristae type.

The number of mitochondria in each cristae category was determined for each pcp1 strain (Figure 3B). For the wild-type strain, the majority of the mitochondria were found to be normal (99.2%). Pcp1p (G233S V321A) also had the majority of its mitochondria in the normal category (88.7%). Pcp1p (S252N) and Pcp1p (G120R) both had a little over 50% normal mitochondria, while only 28.5% of the mitochondria in Pcp1p (G233D) were scored as normal. Cells deleted for pcp1 or containing the Pcp1p (G315D) mutation had no mitochondria with normal ultrastructure. However, Pcp1p (G315D) may exhibit some peptidase activity as a few of the mitochondria were found to have retained some cristae ultrastructure (7.6% normal-vesicular and 4.8% vesicular), a conclusion that is supported by data presented in Figure 2. Pcp1p (G233S V321A) and Pcp1p (S252N) both contained levels of concentric and normal-vesicular mitochondria, although these occurred in greater numbers in Pcp1p (S252N). Both Pcp1p (G120R) and Pcp1p (G233D) contained all four mitochondrial cristae morphologies with Pcp1p (G233D) appearing to have a more severe impact on cristae organization as 71.5% of the mitochondria lacked normal cristae compared to 41.8% in Pcp1p (G120R).

To determine whether there were differences in mitochondrial size between pcp1 mutants the area of each mitochondrion was measured. Only mitochondria believed to be in cross-section were analyzed for this purpose. The cross-sectional areas were measured for each strain and the data presented in Figures 4 and SA3. The majority of the mitochondria in Pcp1p (G233D), Pcp1p (S252N), and Pcp1p (G120R) were found within the 41–60 nm² range, consistent with the mitochondria found in the PCP1 strain. Mitochondria in Pcp1p (G315D) (P = 2.16 × 10⁻¹⁸) and Δpcp1 (P = 6.34 × 10⁻²¹) had a significantly smaller cross-sectional

Figure 4  The analysis of mitochondrial cross-sectional area identifies differences in mitochondria size. Transmission electron microscopy (TEM) images were analyzed for mitochondria presented in cross-section and their area measured. Data were grouped into 20 nm² increments and the percentage of mitochondria in each group presented graphically. The number of mitochondria measured for each strain were as follows: PCP1; n = 72, Pcp1p (S252N); n = 102, Pcp1p (G233D); n = 106, Pcp1p (G120R); n = 122, Δpcp1; n = 69, Pcp1p (G315D); n = 65, and Pcp1p (G233S V321A); n = 82.
area falling within 21–40nm². This is consistent with Amutha et al. (2004) who described Δmfm1 mitochondria as small, round, and lacking cristae folds. Surprisingly, the Pcp1p (G233S V321A) mutant was found to have a significantly larger mitochondria cross-sectional area, with the majority falling in the range of 61–80nm² ($P = 0.0002$).

**ATP synthase and Tim11p steady-state levels are altered in pcp1 mutants**

To understand how each of the Pcp1p mutants resulted in the accumulation of differently structured mitochondria, we focused on characterizing the oligomerization status of the ATP synthase complex in each mutant as it has been linked to cristae architecture (Paumard et al., 2002; Strauss et al., 2008; Davies et al., 2012). First, we measured the steady-state levels of the F₁ complex component, ATP synthase α, and a protein associated with the F₀ region of the ATP synthase complex, Tim11p (subunit ϵ). Immunoblot analysis on isolated mitochondria found no significant differences in the levels of ATP synthase α subunit for Pcp1p (S252N), Pcp1p (G233D), Pcp1p (G233S V321A), and Pcp1p (G120R) when compared with a wild-type PCP1 strain. However, Δpcp1 ($P = 0.03$) and Pcp1p (G315D) ($P = 0.05$) had significantly lower levels of ATP synthase α subunit compared with wild type (Figures 5A and 5B). As previously reported, the Δpcp1 mutant lacked detectable Tim11p (Amutha et al., 2004). The Pcp1p (G315D) also contained no detectable Tim11p, while Pcp1p (S252N), Pcp1p (G233D), and Pcp1p (G120R) had Tim11p levels not statistically different from wild type. Interestingly, the Pcp1p (G233S V321A) mutant was found to have higher Tim11p ($P = 0.008$) (Figures 5C and 5D).

**pcp1 mutants decrease ATP synthase super-complex assembly and activity**

Tim11p has been shown to be required for ATP synthase dimer formation and thus has a role in mitochondrial cristae formation (Arnold et al., 1997; Arselin et al., 2003; Amutha et al., 2004; Everard-Gigot et al., 2005; Bornhöved et al., 2006). To establish whether any of the Pcp1p mutations impacted the ability of ATP synthase to form dimers as well as higher-order complexes, mitochondrial lysates from each pcp1 mutant was analyzed by BN-PAGE using conditions shown to separate ATP synthase into oligomers, dimers, and monomers (Paumard et al., 2002; Arselin et al., 2003; Bornhöved et al., 2006). To identify subtle differences in ATP synthase complex stability, we compared complexes formed using digitonin-to-protein ratios of 1:4:1 (g/g) and 0.7:1 (g/g) (Paumard et al., 2002). Pcp1p (G315D) was omitted from these experiments as no ATP synthase complexes were readily detected (Figure 6C). The Δpcp1 and Pcp1p (G315D) mitochondria (Figure SA4B) were found to only contain free F₁, while all other pcp1 mutants had ATP synthase activity associated with higher-order super-complexes. As expected, when comparing ATP synthase activity between the detergent ratios, an increase in oligomer activity was found using the lower digitonin-to-protein ratio (Figure 6C).

**Discussion**

A number of protein and protein complexes have been identified based on their role in organizing mitochondrial cristae. In budding yeast these include the inner membrane fusion protein Mgm1p, the F₁F₀ ATP synthase dimer, and the multi-subunit mitochondrial contact site and cristae organizing system (MICOS) complex (Strauss et al., 2008; Davies et al., 2012; Harner et al., 2016). In the case of Mgm1p, proteolytic processing of the polypeptide by the rhomboid peptidase, Pcp1p, has been shown to be critical for normal cristae formation (Herlan et al., 2003; Amutha et al., 2004). Targeted mutations that disrupt Pcp1p activity have been generated, primarily as an approach to confirm the catalytic site of the enzyme, focusing on the highly conserved serine and histidine residues at amino acid positions 256 and 313, respectively. Few mutations outside of these two regions have been analyzed (Esser et al., 2002; Sesaki et al., 2003a).

Using a random mutagenesis approach, coupled with a selection for reduced growth on a nonfermentable carbon source, we have identified additional amino acids within Pcp1p required for full peptidase activity. Two of these mutations mapped to an amino acid upstream and downstream to the catalytic dyad and included serine at
Figure 5 Pcp1 mutants have altered steady-state levels for select adenosine triphosphate (ATP) synthase subunits. (A) Nonfunctional pcp1 alleles have reduced levels of ATP synthase α subunit. Proteins from isolated mitochondria were separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) followed by immunoblot analysis using anti-ATPV α subunit and anti-Tom40 antibodies. Tom40p was used as a control to confirm equal loading of samples. (B) Densitometric analysis of the levels of ATP synthase α subunit relative to Tom40p and normalized to the PCP1 wild-type control. Data presented as the average ± standard deviation from four experiments. *P = 0.03 for Δpcp1 and 0.05 for G315D). (C) Tim11p levels vary across the pcp1 mutants. Proteins isolated as described above were analyzed by immunoblot for Tim11p and anti-Tom40 levels. (D) Densitometric analysis of the level of Tim11p relative to Tom40p and normalized to the PCP1 wild-type strain. Data represent the average ± standard deviation from six experiments. *P = 0.008.
position 252 and the glycine at position 315. Whereas the replacement of an isoleucine or an alanine for the serine had been reported previously to have no impact on Pcp1p function (Esser et al., 2002; Sesaki et al., 2003a), our data suggests that the introduction of an asparagine at position 252 interferes with the peptidase activity. Two additional amino acids important for Pcp1p activity that were identified in our screen included a glycine at positions 252 and 315. Whereas the replacement of an isoleucine or an alanine for the serine had been reported previously to have no impact on Pcp1p function (Esser et al., 2002; Sesaki et al., 2003a), our data suggests that the introduction of an asparagine at position 252 interferes with the peptidase activity. Two additional amino acids important for Pcp1p activity that were identified in our screen included a glycine at positions 252 and 315. Interestingly, we isolated two di

![Image](68x450 to 290x710)

Figure 6 Nonfunctional *pcp1* alleles have reduced adenosine triphosphate (ATP) synthase super-complexes formed. ATP synthase complex oligomerization and ATPase activity were measured following blue native-polyacrylamide gel electrophoresis (PAGE). (A) Representative immunoblot showing ATP synthase oligomerization status using ATP synthase α subunit antibodies. Mitochondrial proteins from the strains indicated were solubilized in buffer at a digitonin to protein ratio of 1.4:1 (g/g) and 0.7:1 (g/g). (B) The corresponding sodium dodecyl sulfate PAGE (SDS-PAGE) of Tom40p levels by immunoblot analysis for samples presented in (A) confirm equal loading. (C) Representative in-gel ATP synthase activity for the same *pcp1* strains under the two solubilization conditions described above (n = 3).

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position 252 and the glycine at position 315. Whereas the replacement of an isoleucine or an alanine for the serine had been reported previously to have no impact on Pcp1p function (Esser et al., 2002; Sesaki et al., 2003a), our data suggests that the introduction of an asparagine at position 252 interferes with the peptidase activity. Two additional amino acids important for Pcp1p activity that were identified in our screen included a glycine at positions 252 and 315. Interestingly, we isolated two different amino acid mutations at position 233 that varied in their severity with an aspartic acid less well tolerated than a serine. Sequence comparison of putative Pcp1p homologs from other fungi found glycine, alanine, or a serine at this position of the polypeptide (Figure SA1C), which may help explain the retention of functionality. However, given that this mutant also includes a second amino acid mutation at position 321, a valine to alanine, we cannot rule out the possibility that the effects that we have identified are due to the combined effect of these two alterations.

All of the *pcp1* mutants that we identified had reduced Pcp1p peptidase activity as measured by the altered ratio of short-to-long isoforms of Mgm1p (Figure 2). As shown previously, an imbalance in Mgm1p processing negatively impacts fusion of the mitochondrial inner membrane (Herlan et al., 2004). This is likely due to the different activities ascribed to each isoform with the long isoform responsible for tethering the fusion machinery to the inner membrane while the short isoform mediates the guanosine 5′-triphosphate (GTP)-dependent phospholipid rearrangement required for membrane bending and ultimately membrane fusion (Zick et al., 2009a; Rujiviphat et al., 2015). As expected, these *pcp1* mutants were found to vary in their accumulation of mitochondrial cristae structural defects with the severity in loss of cristae architecture correlating with the severity of Mgm1p processing deficiency. Given that the assembly of ATP synthase into dimers and higher-order oligomers has been shown to be important for the membrane curvature at the base of cristae (Paumard et al., 2002; Strauss et al., 2008; Davies et al., 2010), the assembly status of ATP synthase in the *pcp1* mutants was interrogated using BN-PAGE. Downstream analysis of oligomeric status by F₁ α-subunit western blot and ATPase activity assays were unable to identify ATP synthase complexes in the Pcp1p G315D mutant while a reduction in ATP synthase oligomers was found for the Pcp1p G233D mutant. These findings are consistent with the defects we identified in mitochondrial organization by electron microscopy. However, no substantial differences in ATP synthase assemblages were identified for the remaining *pcp1* mutants despite the accumulation of abnormal mitochondrial structures. Minor differences in the detection of ATP synthase complexes were noted when comparing immunoblot data with that of an ATPase activity assay. Generally, the F₁ α-subunit antibody was more sensitive at identifying ATP synthase oligomers at the higher digitonin: protein ratio compared to the ATPase activity assay while the in-gel activity assay was more sensitive at detecting free F₁ subunits than by immunoblotting. These differences may be attributed to the variability of epitope accessibility to antibody recognition/binding under the native conditions used in BN-PAGE while for the ATPase activity assay the higher detergent levels may interfere with nucleotide binding or hydrolysis activities without disrupting the overall complex stability.

Despite identifying defects in mitochondrial organization for all of the *pcp1* mutants, even the most severe *pcp1* mutants (Δpcp1 and G315D) retained some level of detectable mitochondrial DNA suggesting that loss of cristae folds preceded the loss of the mitochondrial genome. These results are consistent with the findings of
Harnar et al. (2016) using the *mgm1-5* temperature-sensitive mutant. Here, the authors found that only a short incubation at the nonpermissive temperature was required for loss of cristae morphology but the time period was likely too short as to have allowed for mtDNA loss. However, for each mutant a larger percentage of the cells were respiratory incompetent than would be predicted from the mitochondrial DNA staining data (Figures 1A and 1B) or qPCR analysis of genome ratios (Table S1), suggesting that defects in mitochondrial organization can give rise to nonfunctional mitochondrial genomes. This conclusion is confirmed by work from Osman et al. (2015) that found a yeast strain defective in mitochondrial fusion–fission activity accumulated deletions and rearrangements within the mitochondrial genome that resulted in loss of respiratory competence. The finding that ~25% of the cells expressing either the Pcp1p (G120R) or Pcp1p (G233D) mutation lacked detectable mtDNA by DAPI staining but had only a minor reduction in mitochondrial nuclear genome ratios is also consistent with these cells acquiring mitochondrial genomic deletions. The absence of mtDNA by fluorescence microscopy may reflect a property of the mitochondrial nucleoids in these mutants, such as a structural alteration in mtDNA packaging that reduces DAPI binding, or simply an issue associated with resolving the cytoplasmic staining of mtDNA from that of the nucleus, a conclusion that would be supported by the observation that fusion deficient mitochondria are distributed throughout the cell rather than at the cell periphery (McQuibban et al., 2003).

In addition to having defects in cristae morphology, mitochondria in the Pcp1p (G315D) strain were found to have a smaller mitochondrial cross-sectional area than that of the wild-type strain. A similar decrease in mitochondrial size has been previously reported for *MGM1* mutants and for cells deleted for the outer membrane fusion protein *FZO1* (Hermann et al., 1998; Sesaki et al., 2003b; Amutha et al., 2004). A surprising result was finding a larger average mitochondrial cross-sectional area for the Pcp1 (G233S V321A) mutant compared with that of wild-type cells. This mutant also had elevated levels of the ATP synthase dimerization component, Tim11p without an appreciable difference in ATP synthase oligomerization. How could increased Tim11p levels impact mitochondrial architecture? Recent work has shown that Tim11p, as part of the ATP synthase dimer, physically associates with the MICOS component, Mic10 (Eydt et al., 2017; Rampelt et al., 2017). This interaction occurs outside of the cristae junction and involves a subpopulation of Mic10 that is not part of the larger MICOS complex.

In yeast, the MICOS complex is composed of six different protein components (Mic60, Mic19, Mic10, Mic26, Mic27, and Mic12) responsible for the formation and/or stabilization of cristae junctions (review; Quintana-Cabrera et al., 2018). MICOS functions as two sub-complexes. One sub-complex contains the core constituent, Mic60, a transmembrane protein shown to interact with proteins of the preprotein import and sorting pathways and therefore involved in the formation of outer and inner membrane contact sites (Harnar et al., 2011; Hoppins et al., 2011; Zerbes et al., 2012). The second MICOS complex contains Mic10 whose membrane bending activity is responsible for cristae junction curvature (Barbot et al., 2015). Deletion of Mic60 or Mic10 results in the loss of cristae junctions and the formation of lamellar mitochondria structures (Bohnert et al., 2015; Eydt et al., 2017) and both are thought to influence F1F0-ATP synthase oligomerization (Rabl et al., 2009; Eydt et al., 2017; Rampelt et al., 2017). Therefore, it is tempting to speculate that the increased Tim11p in the Pcp1 (G233S V321A) mutant leads to an increase in Mic10 association with ATP synthase dimers, which, in turn, alters the balance for the roles of the MICOS complex and ATP synthase oligomers in cristae structure.

Changes in mitochondrial bioenergetics have been shown to influence Mgm1p processing providing a link between physiological parameters and mitochondrial structure (Herlan et al., 2004). The identification of Pcp1p mutants that vary in Mgm1p processing provide us with an additional tool to probe the link between these inner membrane complexes and cristae morphology.

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**Author contribution**

D.M.G. conceived and designed the experiments; N.X., M.E.H., A.P.B., and D.M.G. performed the experiments; N.X., M.E.H., A.P.B., and D.M.G. analyzed the data; M.E.H. and D.M.G. wrote the paper; M.E.H., A.P.B, and D.M.G. edited the paper.

**Conflict of interest**

The authors declare no conflict of interest. The funding sponsors had no role in the design of the study; in the
collection, analyses, or interpretation of data; in the writing of the manuscript; and in the decision to publish the results.

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Additional supporting information may be found online in the Supporting Information section at the end of the article.

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