Giant Peroxisomes in Oleic Acid-induced *Saccharomyces cerevisiae*

lacking the Peroxisomal Membrane Protein Pmp27p

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**Abstract.** We have purified peroxisomal membranes from *Saccharomyces cerevisiae* after induction of peroxisomes in oleic acid–containing media. About 30 distinct proteins could be discerned among the HPLC- and SDS-PAGE-separated proteins of the high salt–extracted peroxisomal membranes. The most abundant of these, Pmp27p, was purified and the corresponding gene *PMP27* was cloned and sequenced. Its primary structure is 32% identical to PMP31 and PMP32 of the yeast *Candida boidinii* (Moreno, M., R. Lark, K. L. Campbell, and M. J. Goodman. 1994. *Yeast.* 10:1447–1457). Immunoelectron microscopic localization of Pmp27p showed labeling of the peroxisomal membrane, but also of matrix-less and matrix containing tubular membranes nearby. Electronmicroscopic data suggest that some of these tubular extensions might interconnect peroxisomes to form a peroxisomal reticulum. Cells with a disrupted *PMP27* gene (Δpmp27) still grew well on glucose or ethanol, but they failed to grow on oleate although peroxisomes were still induced by transfer to oleate-containing media. The induced peroxisomes of Δpmp27 cells were fewer but considerably larger than those of wild-type cells, suggesting that Pmp27p may be involved in parcelling of peroxisomes into regular quanta. Δpmp27 cells cultured in oleate-containing media form multiple buds, of which virtually all are peroxisome deficient. The growth defect of Δpmp27 cells on oleic acid appears to result from the inability to segregate the giant peroxisomes to daughter cells.

ONLY a few integral proteins of the peroxisomal membranes have been molecularly characterized in uni- and multicellular eukaryotes, using either genetic or reverse genetic methods. Pas3p from *Saccharomyces cerevisiae*, as well as PAF1 and PMP70 from mammalian cells, have been shown to be essential for peroxisome formation (Kamiyo et al., 1990, 1992; Höhfeld et al., 1991; Tsukamoto et al., 1991; Shimozawa et al., 1992). A defect in the ALD protein causes X-linked adrenoleukodystrophy (Mosser et al., 1993). Functions for mammalian PMP22 (Kaldi et al., 1993) and PMP31, PMP32, and PMP47 from *Candida boidinii* (McCammon et al., 1990a; Moreno et al., 1994) have not yet been described. Hence, the peroxisomal membrane remains among the least characterized of the cellular membranes. A further challenge arises from the hypothesis by Lazarow et al. (1980) that peroxisomes of mammals are interconnected and form a peroxisomal reticulum. Lazarow proposed that this peroxisomal reticulum consists of at least two distinct domains: the classical bulbous peroxisomes and tubular extensions that interconnect the bulbous regions. Morphological evidence consistent with this proposal has been presented (Gorgas, 1984; Lazarow and Fuyiki, 1985; Yamamoto and Fahimi, 1987; Lazarow, 1988; Baumgart et al., 1989; Leurs et al., 1993). At present, it is not known whether these morphologically distinct membrane domains are also biochemically and functionally distinct.

In yeasts grown under normal laboratory conditions, peroxisomes are among the least conspicuous organelles, consisting only of a few small and irregularly shaped entities with no evidence for a peroxisomal reticulum (Avers and Federman, 1968; Osumi et al., 1974; Veenhuis et al., 1979, 1987; Veenhuis and Goodman, 1990). Not surprisingly, these structures have resisted purification by conventional cell fractionation methods since their size and density are indistinguishable from other membrane-derived vesicles found in a postnuclear supernatant of a cell homogenate (Szabo and Avers, 1969). However, the situation changes dramatically after induction of peroxisome proliferation as a result of growing certain yeast species on certain carbon sources (Osumi et al., 1974; van Dijken et al., 1975; Roggenkamp et al., 1975; Fukui et al., 1975; Veenhuis and Harder, 1991). This peroxisome proliferation is characterized by an increase in the size and number of the organelles. The large size and dense matrix of induced peroxisomes (often containing paracrystalline bodies) has made it possible to separate them efficiently from other organelles of a postnuclear supernatant (Roggenkamp et al., 1975; Kamiyo et
brane proteins, Pmp27p (peroxisomal membrane protein of *S. cerevisiae* that were growth impaired on oleate. Unlike wild-type cells, *Δpmp27* cells showed only a few, but very large peroxisomes. Moreover, α-Pas3p antibodies decorate Pas3p on immunoblots, they failed to give detectable signals by immuno- fluorescence or immunoelectron microscopy (Höfeld et al., 1991). Hence, an ultrastructural characterization of the membranes delimiting the peroxisomal compartment of *S. cerevisiae*, using antibodies as probes, has not yet been accomplished.

In this paper, we report the purification and characterization of peroxisomes from oleate-induced *S. cerevisiae*. The gene coding for one of the most abundant peroxisomal membrane proteins, Pmp27p (peroxisomal membrane protein of 27 kD), was cloned and sequenced. Immunolocalization of Pmp27p allowed us to detect membranes that extended from bulbous peroxisomes and that are likely to represent tubular peroxisomal appendices. Deletion of the *PMP27* gene yielded cells that grew normally on glucose or ethanol, but that were growth impaired on oleate. Unlike wild-type cells, where oleate induction typically yields 20–30 small peroxisomes, oleate-induced *Δpmp27* cells showed only a few, but very large peroxisomes. Moreover, *Δpmp27* grown on oleate showed multiple buds, most of which were peroxisome deficient. These data suggest that Pmp27p is essential for the maintenance of peroxisome morphology and inheritance in oleic acid–induced *S. cerevisiae*.

**Materials and Methods**

**Strains, Growth Conditions, and General Methods**

The yeast strains used in this study were *S. cerevisiae* wild-types UTL-7A (MATα, 2α, ade2-1, his3-115, leu2-3,112), SKQ2N (MATαα, ade2-1, his3-115, leu2-3,112, his3-115, his3-115, leu2-3,112, ade2-1, ade2-1, his3-115, his3-115, leu2-3,112, ade2-1, his3-115, leu2-3,112, ade2-1, his3-115, leu2-3,112, can1-100/can1-100). Yeast complete (YPD) and minimal (SD) media have been described previously (Erdmann et al., 1991). Hence, an ultrastructural characterization of the membranes delimiting the peroxisomal compartment of *S. cerevisiae*, using antibodies as probes, has not yet been accomplished.

In this paper, we report the purification and characterization of peroxisomes from oleate-induced *S. cerevisiae*. The gene coding for one of the most abundant peroxisomal membrane proteins, Pmp27p (peroxisomal membrane protein of 27 kD), was cloned and sequenced. Immunolocalization of Pmp27p allowed us to detect membranes that extended from bulbous peroxisomes and that are likely to represent tubular peroxisomal appendices. Deletion of the *PMP27* gene yielded cells that grew normally on glucose or ethanol, but that were growth impaired on oleate. Unlike wild-type cells, where oleate induction typically yields 20–30 small peroxisomes, oleate-induced *Δpmp27* cells showed only a few, but very large peroxisomes. Moreover, *Δpmp27* grown on oleate showed multiple buds, most of which were peroxisome deficient. These data suggest that Pmp27p is essential for the maintenance of peroxisome morphology and inheritance in oleic acid–induced *S. cerevisiae*.
Purification and Amino Acid Sequencing of Pmp27p

High salt–extracted peroxisomal membranes were prepared from oleic acid–induced SKQ2N cells. Further separation of the peroxisomal membrane proteins was achieved by reverse-phase HPLC. The peroxisomal membrane proteins were solubilized in 4% SDS, 10 mM Tris/HCl pH 7.4, diluted fivefold in 200 mM Tris/HCl pH 7.4, 20 mM DTT, and heated for 10 min at 60°C. Insoluble debris was removed by centrifugation in a microfuge for 10 min, and the soluble fraction was loaded onto an Aquapore butyl (C4) column (100 x 10 mm; Brownlee Labs, Applied Biosystems, Inc., Foster City, CA) equilibrated with 60% formic acid. After a 5-min linear increase to 9.9% acetonitrile in 60% formic acid, the column was eluted with a linear gradient of 32 ml 9.9%-33.0% acetonitrile in 60% formic acid, and 0.4-ml fractions were collected. Eluted fractions were dried in a Speed Vac concentrator (Savant Instruments Inc., Hicksville, NY), and pellets were solubilized in SDS sample buffer and analyzed by SDS-PAGE.

For sequencing of Pmp27p, the SDS samples of HPLC fractions containing this protein (fractions 52–57) were pooled and separated on a 12% polyacrylamide gel. Polypeptides were electrophoretically transferred on a polynvinylidene difluoride membrane and visualized with 0.1% amidobisocin in 10% acetic acid. Pmp27p was excised and subjected to NH2-terminal sequence analysis on a gas phase sequenator (Applied Biosystems).

Isolation and Sequencing of PMP27

A PMP27–specific probe was generated by the PCR. According to the obtained NH2-terminal sequence of Pmp27p, degenerate sense [5'ATGGATCCGA(CT)AGTT(CT)AGTTA(CT)CA(CT)CC3'] and antisense [5'AGAAATTC(CT)AAIAC(CT)TT(CT)CC(CT)TGT(AGCT)CC3'] oligonucleotide primers were synthesized, and the corresponding genomic region of the PMP27 gene was amplified by the polymerase chain reaction with yeast genomic DNA (100 μg; Promega Corp., Madison, WI) as template. The amplification product of the expected size was isolated and subcloned into pBluescript SK+ (Stratagene, La Jolla, CA), resulting in pkP27. The authenticity of the insert was confirmed by sequencing. The PMP27-specific probe was generated by the PCR. According to the obtained NH2-terminal sequence of Pmp27p (see Fig. 3), the organelle or membrane pellets were fixed as described above, dehydrated in ethanol, and embedded in Epon 812. Silver sections were stained with uranyl acetate (Farquhar and Palade, 1965). The pellets were dehydrated with ethanol, treated with propylene oxide, and embedded in Epon 812. Silver sections were stained with uranyl acetate and lead citrate (Reynolds, 1965).

For cryoimmunogold labeling of whole cells, cells were fixed in 3% paraformaldehyde/0.5% glutaraldehyde in 0.1 M cacodylate, pH 7.4. The cell wall of fixed cells was removed by incubation with 0.5 mg/ml zymolase 20T (ICN) in 1.2 M sorbitol, 0.1 M phosphate-citrate, pH 7.0. Pellets were washed with 0.6 M sorbitol in the same buffer, embedded in 10% gelatin, and refixed as above. Pellets were incubated with 2.3 M sucrose in PBS, and samples were frozen in liquid nitrogen until use (Tokuyasu, 1973). Ultrathin sections were made with glass knives in a Reichert-Jung FC-4E cryoultramicrotome. The sections were collected on Formvar–carbon coated nickel grids, treated with 1% BSA in PBS, and incubated with rabbit antihistidine detection (dilution = 1:5), or monoclonal 12CA5 antiserum against the HA-tag (dilution = 1:20), and goat anti-rabbit or anti-mouse IgG-gold (5 or 10 nm; Amersham Life Science, Arlington Heights, IL). The grids were processed and stained according to Griffiths et al. (1983).

For the immunolabeling of purified peroxisomes (see Fig. 1) and high salt–extracted peroxisomal membranes, pellets were fixed in 2.5% glutaraldehyde in 0.1 M cacodylate, postfixed with 1% osmium tetroxide, and stained with uranyl acetate (Farquhar and Palade, 1965). The pellets were dehydrated with ethanol, treated with propylene oxide, and embedded in Epon 812. Silver sections were stained with uranyl acetate and lead citrate (Reynolds, 1965).

For epimethion labeling of whole cells, cells were fixed in 3% paraformaldehyde/0.5% glutaraldehyde in 0.1 M cacodylate, pH 7.4. The cell wall of fixed cells was removed by incubation with 0.5 mg/ml zymolase 20T (ICN) in 1.2 M sorbitol, 0.1 M phosphate-citrate, pH 7.0. Pellets were washed with 0.6 M sorbitol in the same buffer, embedded in 10% gelatin, and refixed as above. Pellets were incubated with 2.3 M sucrose in PBS, and samples were frozen in liquid nitrogen until use (Tokuyasu, 1973). Ultrathin sections were made with glass knives in a Reichert-Jung FC-4E cryoultramicrotome. The sections were collected on Formvar–carbon coated nickel grids, treated with 1% BSA in PBS, and incubate with rabbit antihistidine detection (dilution = 1:5), or monoclonal 12CA5 antiserum against the HA-tag (dilution = 1:10). After buffer washes, the grids were incubated with goat anti-rabbit or anti–mouse IgG-gold (5 or 10 nm; Amersham Life Science, Arlington Heights, IL). The grids were processed and stained according to Griffiths et al. (1983).

For the immunolabeling of purified peroxisomes (see Fig. 1) and high salt–extracted peroxisomal membranes (see Fig. 3), the organelle or membrane pellets were fixed as described above, dehydrated in ethanol, and embedded in Lowicryl. Pale gold sections were collected on Formvar–carbon coated nickel grids. The sections were blocked with PBS containing 1% BSA and incubated with rabbit anti-Poxlp, anti-Fox3p (dilution = 1:500), or monoclonal 12CA5 antiserum against the HA-tag (dilution = 1:5). Characterization of the anti-Poxlp antibodies will be published elsewhere (Will G., and W. H. Kunau, personal communication). After washings, the grids were incubated with goat anti–rabbit or anti–mouse IgG-gold 10 nm gold (Amersham) and stained with uranyl acetate.

Electron Microscopy

For the morphology of peroxisomes and high salt–extracted peroxisomal membranes, pellets were fixed in 2.5% glutaraldehyde in 0.1 M cacodylate, postfixed with 1% osmium tetroxide, and stained with uranyl acetate (Farquhar and Palade, 1965). The pellets were dehydrated with ethanol, treated with propylene oxide, and embedded in Epon 812. Silver sections were stained with uranyl acetate and lead citrate (Reynolds, 1965).

Epitope Tagging of Pmp27p

For immuno localization studies on Pmp27p, an epitope tag encoding 10 amino acid residues from the influenza virus hemagglutinin antigen (HA)1 (Wilson et al., 1984; Field et al., 1988) plus two flanking glycines as a spacer was inserted at the COOH terminus of the protein. For the insertion of the epitope tag into Pmp27p, a BglII site was introduced in front of the stop codon of the PMP27 gene by the polymerase chain reaction. In parallel, a BamHI site was introduced behind the stop codon to facilitate further subclonings. The two complementary oligonucleotides, yHA-1 and yHA-2 (Wozniak et al., 1994), with the following sequence 

1. Abbreviations used in this paper: HA, hemagglutinin antigen; PMP, peroxisomal membrane protein.
Figure 1. Time course of induction of a peroxisomal membrane protein (Pas3p) and a peroxisomal matrix protein (Fox3p). Cells were shifted to oleic acid-containing medium and cultured for the indicated time points. (A) Whole-cell extracts were prepared for SDS-PAGE and Pas3p, Fox3p, and Kar2p (a marker for the endoplasmic reticulum; Rose et al., 1989) were detected by immunoblot analysis. The amounts analyzed were 1% of the homogenates from 30 mg of cells. (B) Relative concentrations of the proteins as determined by laser densitometry of the immunoblot data. The concentration at 24 h postinduction was set at 100%.

Immunoblots

Western blot analysis was performed according to standard protocols (Towbin et al., 1979) using anti-rabbit or anti-mouse IgG-coupled HRP as secondary antibody (Amersham). Protein–antibody complexes were visualized by treatment with HRP chemoluminescence developing reagents (ECL system; Amersham). Western blots were quantitated by laser densitometry. Polyclonal rabbit antibodies against Fox3p (Erdmann and Kunau, 1994), Kar2p (Rose et al., 1989), and p32 (Pain and Blobel, 1990) were used at dilutions of 1:50,000. Pas3p was detected with a nondiluted affinity-purified rabbit antiserum against the protein (Höhfeld et al., 1991). HA-tagged Pmp27p was detected with monoclonal 12CA5 antiserum against the HA-Tag (RAbCO; dilution = 1:3,000).

Analytical Procedures

Catalase (EC 1.11.1.6) was assayed according to Moreno de al Garza et al. (1985). Protein was measured by laser densitometry of proteins separated by SDS-PAGE and stained with Coomassie blue, as well as by the method of Bradford (1976) with bovine serum albumin as standard.

Results

Oleate Induction of Peroxisomal Proteins

S. cerevisiae cells were shifted from growth in glucose-containing media to growth in oleate-containing media. At various time points, cell homogenates were prepared, and the proteins were separated by SDS-PAGE, transferred to nitrocellulose, and probed with antibodies against either Pas3p (a peroxisomal membrane protein), Fox3p (a peroxisomal matrix protein), or Kar2p (a matrix protein of the endoplasmic reticulum) (Fig. 1). Even before shift to oleate-containing media, cells contained readily detectable levels of Pas3p (Höhfeld et al., 1991; Fig. 1, lane 1 in this manuscript), whereas Fox3p was not detectable (Fig. 1, lane 1). During a 24-h induction period, Pas3p increased about four to fivefold, whereas Fox3p increased from nondetectable to clearly detectable levels (Fig. 1). The levels of Kar2p, serving as a control, remained essentially unchanged during the 24-h induction period.

Peroxisome Induction Causes Shift of Pas3p from Light to Heavy Membranes

As Pas3p could be used as a peroxisomal membrane marker that is detectable in noninduced cells, it became possible to assess physical characteristics of the peroxisomes before and after various times of induction. This was done by gradient centrifugation of postnuclear supernatants of cell homogenates. We found that before induction, most of the Pas3p-reactive membranes banded at a density of 1.15 g/ml (Fig. 2, fraction II). At the 3-h point, the bulk of Pas3p-reactive membranes shifted to a broad band of higher density between 1.16 and 1.20 g/ml (Fig. 2). This bulk shift to higher density continued at the 6-h point, and it was completed at the 12-h point when the bulk of the Pas3p-reactive membrane equilibrated at 1.21 g/ml (Fig. 2, fraction 3). However, a fraction of the Pas3p-reactive membrane remained at 1.15 g/ml and at the broad region between 1.15 and 1.21 g/ml (this was clearly evident after overexposure of the

Figure 2. Isopycnic gradient centrifugation analysis of Pas3p reactive membranes of cells shifted to oleate containing media and cultured for the indicated times. Cell homogenates were fractionated on continuous 15-35% Nycodenz gradients (see Materials and Methods), and gradient fractions were analyzed by immunoblots with αPas3p. Fraction numbers are indicated and each lane corresponded to 1.25% of the fraction volume. Peroxisomes from noninduced cells peaked at a density of 1.15 g/cm³ (fraction 11), but at later stages of induction, peroxisomes were mainly obtained at 1.21 g/cm³ (fraction 3).
Figure 3. Electron microscopy of isolated peroxisomes before (A) and after (B) extraction with high salt. (A) The ultrastructural appearance of purified peroxisomes from *S. cerevisiae* that had been induced for 9 h on oleic acid medium is characterized by the presence of finger-like membrane extensions (black arrowheads). Some of the peroxisomes appear to be broken (asterisks). Few mitochondria (m) were found in the preparation. A higher magnification of a peroxisome is shown in the inset, showing the continuity of the peroxisomal membrane between the bulbous peroxisome and its extended domain. Bars, 1 μm and 0.1 μm (inset). (B) Ultrastructural appearance of acyl-CoA oxidase crystals in "high salt"-extracted peroxisomal membranes (see Materials and Methods). The inset shows the localization of acyl-CoA oxidase within the crystals by immunoelectron microscopy. Sections were probed with a polyclonal antibody against the acyl-CoA oxidase from *S. cerevisiae* and goat anti-rabbit antibodies coupled to 10 nm gold. Bars, 0.5 μm.
branes (Fig. 3 B) without the dense cortical layer that is characteristically of nonextracted peroxisomes (Fig. 3 A). However, some of the membranes were associated with striking paracrystalline structures that must have formed during extraction (Fig. 3 B). These paracrystalline structures contain Poxlp as shown by immunogold labeling (Fig. 3 B, inset).

High salt extracted peroxisomal membranes were solubilized in SDS and the proteins separated by HPLC. Proteins in various HPLC fractions were separated by SDS-PAGE and visualized by Coomassie blue (Fig. 5). About 30 distinct polypeptides can be discerned. One of the major proteins of an apparent molecular mass of 27 kD (Fig. 5) was subjected to partial protein sequencing in preparation for DNA cloning and sequencing of the corresponding PMP27 gene (see Materials and Methods).

DNA-deduced Primary Structure of Pmp27p

The PMP27 open reading frame encodes a polypeptide of 236 amino acid residues with a calculated molecular mass of 26,879 D (Fig. 6 A) in good agreement with the molecular mass estimated by SDS-PAGE (Fig. 5). Hydrophilicity plots according to Kyte and Doolittle (1982) revealed several hydrophobic regions, but none of them appeared to fulfill the requirements for a membrane spanning α helix (data not shown). However, using a manual β turn identification pro-

![Figure 4. Successive extraction of peroxisomes and immunological detection of peroxisomal marker proteins in subfractions. (A) Coomassie stain of total peroxisomal proteins (lane 1) and proteins of peroxisomal subfractions obtained by successive treatment of total purified peroxisomes with low salt, high salt, and pH 11.0 (see Materials and Methods). Lane 5 shows the protein content of the final membrane pellet. (B) Immunological detection of the peroxisomal matrix marker Fox3p and the peroxisomal membrane marker Pas3p by Western blot analysis of the subfractions. The amount of the peroxisomal subfractions loaded in lanes 2–5 corresponded to four equivalents of the total peroxisomal protein in lane 1.](image-url)
staining. The position of Pmp27p is indicated by an arrowhead. The position of acyl-CoA oxidase, identified by peptide sequencing (data not shown), is marked by an asterisk. The amount per lane corresponded to 12.5% of the total fraction.

Pmp27p is Essential for Growth on Oleate

One genomic copy of PMP27 in the diploid wild-type W303 was replaced with LEU2. The proper integration of the deletion construct into the genome was confirmed by Southern blotting (data not shown). Tetrad analysis showed a 2:2 segregation for growth ability. All spores derived from the tetrads were viable, indicating that Pmp27p is not essential for growth on YPD medium. However, on oleic acid medium, a 2:2 segregation for growth ability was observed. The growth defect on oleic acid medium cosegregated with the Leu+ marker, therefore indicating that the PMP27 gene product is essential for growth on oleic acid medium. However, Δpmp27 mutant cells grew normally on ethanol medium (data not shown). The impaired growth phenotype on oleate of a haploid Δpmp27 strain, derived from this tetrad analysis, is shown in Fig. 7A. Δpmp27 cells regained

Figure 6. Nucleotide and deduced amino acid sequence of the S. cerevisiae PMP27 locus (A) and comparison with PMP31 from C. boidinii (B). (A) A putative oleic acid responsive element (Einerhand et al., 1993) and a presumptive TATA sequence (Struhl, 1987) in the 5' noncoding region, as well as presumptive termination sequences in the 3' noncoding region (Zaret and Sherman, 1982), are underlined. The underlined amino acid sequence was obtained by peptide sequencing of the purified Pmp27p. (B) Identical residues are indicated by vertical bars. Two dots represent similar residues. The identity between both proteins is 32%, and the similarity 59%. Similarity rules: G=A=S, P=A=S, S=T=A, D=E=Q=N, K=R, V=I=L=M=F, F=Y. These sequence data are available from EMBL/GenBank/DDBJ under accession No. X81465.
Figure 7. Pmp27p is essential for growth on oleic acid medium. (A) Wild-type W303a, Δpmp27 mutant, and Δpmp27 mutant cells expressing Pmp27p or HA-tagged Pmp27p from single-copy plasmids were plated on oleic acid medium and incubated for 7 d at 30°C. (B) Growth curve of wild-type, Δpmp27, and complemented Δpmp27 in oleic acid liquid culture. The growth defect of the Δpmp27 mutant becomes obvious after 12 h, when wild-type and the complemented strain enter the logarithmic growth phase. At this timepoint, the growth of the mutant ceased.

the ability to grow on oleic acid medium when transformed with the wild-type copy of PMP27 (Fig. 7 A). The growth defect of Δpmp27 could also be complemented by expression of a tagged Pmp27p-HA, indicating that the tagging had no obvious effect on the function of the protein (Fig. 7 A). Fig. 7 B shows the rate of growth of the various strains after transfer from glucose-containing media to oleate-containing media. Complemented Δpmp27 cells grew as well as wild-type cells, whereas mutant Δpmp27 cells ceased to grow after transfer to oleate.

Immunolocalization of Pmp27p

The cellular location of epitope-tagged Pmp27p (Pmp27p-HA) was examined by immunofluorescence microscopy, immunoblot analysis of subcellular fractions, and immunoelectron microscopy. The tag, which consisted of a 10-amino acid epitope derived from the hemagglutinin antigen (HA), was introduced at the extreme COOH terminus of Pmp27p. The tagged protein was expressed in Δpmp27 cells, which contain a chromosomal disruption of the wild-type PMP27 gene (see Materials and Methods). Expression of the tagged Pmp27p was under the control of its own promoter on a single-copy plasmid. As expression resulted in functional complementation of the mutant phenotype of Δpmp27 (see Figs. 7, 12, and 14), the tagging apparently did not influence the function of Pmp27p. Thus, subcellular localization of the tagged Pmp27p can be expected to closely mirror that of wild-type Pmp27p.

The HA-tagged Pmp27p was readily identified among the SDS-PAGE-separated polypeptides of a total homogenate of cells grown for 12 h in oleate-containing medium with αHA monoclonal antibodies (Fig. 8 A).

To examine whether PMP27 expression is induced by growth in oleate-containing medium, cells were grown on low glucose medium and were subsequently shifted to oleate containing medium. At various time points, cell homogenates were prepared and analyzed by immunoblotting with αHA mAbs and αFox3p. Pmp27p was found to be highly inducible by oleate (Fig. 8 B) and its induction appeared to precede that of Fox3p by 1–2 h (Fig. 8 B). All further analyses were done with cells at the 12-h time point of induction.

Double immunofluorescence microscopy using αFox3p and αPmp27p-HA showed an identical staining pattern, demonstrating that Pmp27p is a peroxisomal protein (Fig. 9).

Figure 8. Immunological detection of HA-tagged Pmp27p (A) and time course of Pmp27p induction by oleic acid (B). (A) Equal amounts of whole-cell lysates from oleic acid induced Δpmp27 cells expressing Pmp27p (lane 1) and HA-tagged Pmp27p (lane 2) were subjected to Western blot analysis with rabbit antiserum against Fox3p and mAb against the HA-tag (see Materials and Methods). The amount loaded per lane corresponds to 0.5% of extracts from 30 mg of cells. (B) Δpmp27 [Pmp27p-HA] cells expressing the HA-tagged Pmp27p were shifted to oleic acid-containing medium. At indicated timepoints, whole-cell extracts were prepared and subjected to Western blot analysis with mAb against the HA-tag.
Double immunofluorescence microscopy localization of Fox3p and HA-tagged Pmp27p. Oleic acid induced Δpmp27 cells were processed for double immunofluorescence microscopy using mAb against the HA-tag (anti-Pmp27p-HA) and a rabbit antibody against peroxisomal thiolase (anti-thiolase). Secondary antibodies were FITC-conjugated anti-mouse IgG and Texas red–conjugated anti-mouse IgG. Bar, 5 μm.

Immunoblot analyses of cell fractions that were obtained by differential centrifugation of a homogenate are shown in Fig. 10 A. Although most of the Pmp27p-HA–reactive membranes that were present in a postnuclear supernatant (lane 1) sedimented at 25,000 g (lane 3), a significant amount remained in the corresponding supernatant (lane 2). This latter material may represent tubular membranes that were sheared off from the peroxisomal membrane during homogenization. The membranes that were pelleted at 25,000 g were resuspended and further fractionated by sucrose gradient centrifugation. Immunoblot analysis of fractions probed with αHA, αFox3p, and αp32 (p32 being a mitochondrial marker) is shown in Fig. 10 B. As expected for peroxisomal components (see Fig. 2), most of the αPmp27p– and αFox3p–reactive membranes peaked at a density of 1.21 g/ml (Fig. 10 B, fraction 9) clearly separated from mitochondria that peak at a density of 1.17 g/ml (Fig. 10 B, fraction 17). However, a significant amount of the αPmp27p–HA–reactive membranes banded at lighter densities (Fig. 10 B). Again, these lighter membranes could be derived from tubular membranes sheared off from the peroxisomes.

Pmp27p-HA that is associated with the purified membrane fraction behaves as an integral membrane protein since most of it was not extracted at pH 11.0 (Fig. 10 C).

Immunoelectron microscopy of isolated peroxisomes using αPmp27p-HA showed gold labeling of the peroxisomal periphery consistent with Pmp27p being a peroxisomal membrane protein (Fig. 11 A). Immunoelectron microscopy

Figure 9. Double immunofluorescence microscopy localization of Fox3p and HA-tagged Pmp27p. Oleic acid induced Δpmp27 cells were processed for double immunofluorescence microscopy using mAb against the HA-tag (anti-Pmp27p-HA) and a rabbit antibody against peroxisomal thiolase (anti-thiolase). Secondary antibodies were FITC-conjugated anti-mouse IgG and Texas red–conjugated anti-mouse IgG. Bar, 5 μm.

Figure 10. Coenrichment of HA-tagged Pmp27p and peroxisomal thiolase during peroxisome isolation (A and B) and intraperoxisomal localization of Pmp27p-HA (C). (A) Immunoblot analysis of cell fractions that were obtained by differential centrifugation (see Materials and Methods) of the cell homogenate from oleic acid induced Δpmp27 cells expressing the HA-tagged Pmp27p. (B) The organelles of the 25,000-g pellet were separated on a 36–68% (wt/vol) sucrose gradient (see Materials and Methods). 1.2-ml fractions were collected from the bottom of the gradient. Localization of HA-Pmp27p, as well as peroxisomal thiolase and mitochondrial p32 (Pain et al., 1990), in fractions was monitored by immunoblot analysis. Peroxisomes peaked in fraction 9 at a density of 1.23 g/cm³. Mitochondria peaked in fraction 17 at a density of 1.17 g/cm³. Pmp27p-HA as well as Fox3p were mainly found in the peroxisomal peak fractions. (C) The isolated peroxisomes (fraction 9 in B) were extracted by low salt, high salt, and pH 11.0 treatments (see Materials and Methods). As most of the Pmp27p-HA was not extracted by either means (lane 5), it behaves as an integral membrane protein. Equivalent amounts of proteins were loaded per lane. HA-Pmp27p and thiolase amounts in peroxisomal subfractions were monitored by Western blot analysis (see Materials and Methods).
of frozen thin sections of spheroplasts using α Pmp27p-HA showed gold labeling in the peroxisomal periphery, as expected for a membrane protein (Fig. 11 B). However, there also was gold labeling of a region in the vicinity of the peroxisomes that coincided with tubular membrane loops (Fig. 11 B, black arrows). Double immunoelectron microscopy of frozen thin sections of cells using α Pmp27p-HA (10 nm gold) and α Fox3p (5 nm gold) showed the expected gold labeling of the peroxisomal membrane and the peroxisomal matrix, respectively (Fig. 11 C). In addition, there was labeling with only 10 nm gold (Fig. 11 C, black arrowheads) in what appear to be matrixless, peroxisome-attached tubules. In some sections, an interconnection of bulbous peroxisomes by a tubular matrix-containing region could be detected (Fig. 11 D).

**Morphological Characterization of Δpmp27 Cells**

Immunofluorescence microscopy using α Fox3p was done with methanol-fixed spheroplasts of wild-type cells, Δpmp27 cells, and Δpmp27 cells complemented with a copy of the wild-type PMP27 gene (Fig. 12). Analysis was after cells were shifted to growth in oleate-containing media for a period of 0–24 h. When induced for 12 h, in wild-type spheroplasts, a characteristic punctate pattern of 10–20 small spots was observed (Fig. 12 A, upper panel). In the mutant Δpmp27 cells, however, there was a striking reduction in the number of spots with a considerable increase in the size of the individual spots (Fig. 12 A, middle panel). Complementation of the mutant cells with the wild-type gene gave a pattern (Fig. 12 A, lower panel) indistinguishable from that of wild-type cells (Fig. 12 A, upper panel). A time course of oleic acid induced peroxisome proliferation of Δpmp27 cells and complemented mutant cells is shown in Fig. 12 B. Peroxisomes were detected in both strains, even from the very beginning of induction, and no significant difference with respect to the size and number of detected organelles was observed (Fig. 12 B, 0 and 3 h). However, as early as 6 h after the start of induction, the mutant phenotype, indicated by fewer but bigger peroxisomes, became obvious. Most striking was the difference at later stages of induction, when the complemented strain was filled with small dots, but only one to a few giant peroxisomes were detected in the Δpmp27 mutant cells (Fig. 12 B, 24 h).

Electron microscopy (Fig. 13, A and B) and immunoelectron microscopy (Fig. 13, C and D) of thin sections of Δpmp27 mutant cells and complemented Δpmp27 cells using...
Figure 12. Immunofluorescence microscopy localization of thiolase in wild-type, Δpmp27 mutant, and Δpmp27 mutant cells expressing Pmp27p. (A) Cells were induced for 12 h on oleic acid medium. The panels show either thiolase localization by immunofluorescence, DAPI staining of DNA by fluorescence, or phase contrast micrographs of the same cells. Bar, 10 μm. (B) Comparison of oleic acid induced peroxisome proliferation in Δpmp27 mutant and Δpmp27 mutant cells expressing Pmp27p. Thiolase localization was determined using a polyclonal antiserum against the enzyme and Texas red-labeled donkey anti-rabbit antiserum. Bar, 5 μm.
Figure 13. Electron microscopy (A and B) and immunoelectron microscopy (C and D) of 12-h oleic acid-induced Δpmp27 mutant cells (A and C) and complemented Δpmp27 cells (B and D). The mutant phenotype (A and C) was characterized by the presence of fewer, but larger peroxisomes. Sections were probed with polyclonal antiserum against thiolase and goat anti-rabbit antibodies coupled to 10 nm gold. p, peroxisome; m, mitochondrion; n, nucleus; l, lipid droplet. Bars, 0.5 μm.

α Fox3p confirmed the immunofluorescent images of Fig. 12. Peroxisomes in the mutant cells were considerably larger and less abundant than those in the complemented strain.

Δpmp27 Cells Are Defective in Peroxisomal Inheritance

Δpmp27 cells cultured in oleic acid liquid medium showed multiple buds (Fig. 14). Electron microscopical investigation of these cells showed that giant peroxisomes could easily be detected in the majority of mother cells but no peroxisomes were found in cell buds. Other cell organelles (mitochondria, nuclei) showed normal morphology, and they were nearly always detected in the buds (data not shown). Immunofluorescence microscopy of the mutant cells incubated for 48 h in oleic acid medium showed the presence of one to a few giant peroxisomes in mother cells and confirmed the absence of peroxisomes in virtually all cell buds (Fig. 15).

Discussion

We have identified an integral membrane protein, Pmp27p, of S. cerevisiae peroxisomes. This protein served as a membrane marker for immunoelectron microscopy analysis of the peroxisomal compartment of S. cerevisiae, providing evidence for the existence of tubular extensions of the peroxisomal membrane. Yeast cells lacking Pmp27p possess giant peroxisomes, and they are defective for peroxisomal inheritance.

Pmp27p is only the second integral membrane protein of S. cerevisiae that has been molecularly cloned and sequenced, Pas3p (Höfeld et al. 1991) being the first. In peroxisomal membranes that were purified from cells induced for peroxisome proliferation by growth on oleate, Pmp27p was the most abundant peroxisomal membrane protein (Fig. 5). Analysis of its DNA-deduced primary structure did not yield any clues as to its function. So far we have no
data on the topology of Pmp27p in the peroxisomal membrane. Searches in the data banks showed similarity to two integral peroxisomal membrane proteins of C. boidinii, PMP31 and PMP32 (Moreno et al., 1994). It remains to be seen whether cbPMP31 or cbPMP32 can compensate for the loss of scPmp27p in ∆pmp27 cells. The function of cbPMP31p or cbPMP32p is unknown.

**Morphologically Distinct Peroxisomal Domains**

Not surprisingly, immunoelectron microscopy yielded localization of Pmp27p-HA to the periphery of isolated peroxisomes, consistent with Pmp27p being a membrane protein (Fig. 11 A). However, immunoelectron microscopy of frozen thin sections of oleate induced cells showed not only labeling of the membrane surrounding the dense spherical peroxisomes, but also labeling of closely associated tubular regions (Fig. 11, B and C). These extensions might correspond to the reported catalase negative loops and tubular extensions of spherical peroxisomes that have been seen before in mammalian cells (Lazarow et al., 1980; Lazarow and Fujiki, 1985; Yamamoto and Fahimi, 1987; Baumgart et al., 1989). Thus, *S. cerevisiae* peroxisomes, as mammalian peroxisomes, appear to consist of morphologically distinct domains. Some of the tubular peroxisomals extensions observed might interconnect bulbous peroxisomes in *S. cerevisiae* (Fig. 11 D) to form a peroxisomal reticulum as found in mammalian organisms (Lazarow et al., 1980; Gorgas, 1984; Yamamoto and Fahimi, 1987).

Consistent with the presence of tubular extensions in peroxisomes of *S. cerevisiae* are the finger-like and vesicular extensions that we found to be associated with isolated peroxisomes (Fig. 3). Similar extensions of peroxisomal membranes have been observed in isolated mammalian peroxisomes (Lüers et al., 1993). These structures could be interpreted as buds pinching off mature peroxisomes. However, these structures could also be remnants of tubular extensions of peroxisomes or tubular connections between peroxisomes that remain attached to the peroxisomes during homogenization and cell fractionation. More distal regions of the tubular peroxisomal domains might be fragmented into small, light vesicles. Our data also suggest that peroxisomes of increased density arise from lighter preforms during induction (Fig. 2). One interpretation of this results could be that the observed light membranes represent small spherical organelles that increase in density during induction as a consequence of massive protein import and enlargement. However, if the light fraction would represent tubular membranes, much of the inconspicuous peroxisomal compartment in noninduced *S. cerevisiae* may consist of tubular peroxisomal domains rather than bulbous peroxisomes. During oleic acid induction in *S. cerevisiae*, bulbous peroxisomes might arise by local dilations of tubular peroxisomal segments.

Besides the finger-like extensions, the isolated peroxisomes also showed the presence of a dense submembranous layer that was absent in the finger-like extension (Fig. 3 A). This dense layer may prevent tight resealing of the surrounding peroxisomal membrane after injury during homogenization and cell fractionation, particularly if this layer is rigid and not readily deformable. The membrane leakiness that is ascribed to isolated peroxisomes could be caused by ruptured and incompletely resealed membranes overlying this cortical layer. Poxlp is among the major constituents of this cortical layer. It remains to be seen whether this layer represents a physiologically relevant interaction of the matrix with the overlying membrane.

**Function of Pmp27p**

Deletion of the PMP27 gene yielded cells with normal growth on glucose or ethanol, but with impaired growth on oleate. When grown under noninducing conditions, wild-type and ∆pmp27 mutant cells did not differ with respect to size and number of peroxisomes (Fig. 12 B), suggesting that Pmp27p is not required for the division and inheritance of
peroxisomes under noninducing conditions. During oleic acid induction, however, a progressive decrease in the number of peroxisomes per mutant cell was observed, and at later stages of induction, the few mutant peroxisomes were considerably larger than those in induced wild-type cells (Figs. 12 and 13). Failure of the Δpmp27 cells in parceling mature peroxisomes into smaller quanta may be the reason for their overproportional enlargement during induction. The Δpmp27 mutant phenotype is also characterized by the formation of multiple buds upon growth in oleic acid medium (Fig. 14). This result does not only indicate that the mother cells survived on oleic acid medium, but it also suggests that the cells metabolize oleic acid, which requires functional peroxisomes. Thus the giant peroxisomes observed might still fulfill their function in oleic acid metabolism. This is also in agreement with the comparison of the protein composition of purified wild-type and mutant peroxisomes which, except for the absence of Pmp27p, did not reveal significant differences (data not shown). As the majority of the cell buds of oleic acid-induced Δpmp27 mutant cells do not contain peroxisomes (Fig. 15), the inability of the mutant cells to grow on oleic acid is more likely caused by a defect in peroxisomal inheritance. However, as distribution of peroxisomes to daughter cells in Δpmp27 mutant cells still occurs to some extent at early time points of induction (data not shown), Pmp27p does not seem to be directly involved in migration of peroxisomes during mitosis. One simple explanation for the inheritance defect could be that the peroxisomes in Δpmp27 mutant cells may be too large to pass into the bud. However, even if the giant peroxisomes could become inherited, because of the suggested defect in parceling of mature peroxisomes, there would be no peroxisomes left to inherit after a few generations. Recently two yeast mutants with a defect in mitochondrial inheritance have been described (Sogo and Yaffe, 1994; Burgess et al., 1994). These mutants are characterized by the presence of giant mitochondrial structures that cannot be inherited to the daughter cells. In both cases, this phenotype is caused by defects in mitochondrial outer membrane proteins. It was suggested that mitochondrial morphology is maintained by binding of these proteins to the cytoskeleton. Likewise, if in S. cerevisiae the structure of peroxisomes or of a peroxisomal reticulum is maintained by the cytoskeleton, Pmp27p may serve as a peroxisomal membrane anchor to the cytoskeleton. The identification of proteins that interact with Pmp27p will provide further insight into molecular mechanisms regulating the morphology, proliferation, and inheritance of peroxisomes.

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