Peroxisomal Remnant Structures in *Hansenula polymorpha* pex5 Cells Can Develop into Normal Peroxisomes upon Induction of the PTS2 Protein Amine Oxidase

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We have analyzed the properties of peroxisomal remnants in *Hansenula polymorpha* pex5 cells. In such cells PT51 matrix protein import is fully impaired. In *H. polymorpha* pex5 cells, grown on ethanol/ammonium sulfate, conditions that repressed the PTS2 protein amine oxidase (AMO), peroxisomal structures were below the limit of detection. In methanol/ammonium sulfate-grown cells, normal peroxisomes are absent, but a few small membranous structures were observed that apparently represented peroxisomal ghosts since they contained Pex14p. These structures were the target of a Pex10p.myc fusion protein that was produced in pex5 cells under the control of the homologous alcohol oxidase promoter (strain pex5::P$_{AUX}$PEX10.MYC). Glycerol/methanol/ammonium sulfate-grown cells of this transformant were placed in fresh glucose/methylamine media, conditions that fully repress the synthesis of the Pex10p.myc fusion protein but induce the synthesis of AMO. Two hours after the shift Pex10p.myc-containing structures were detectable that had accumulated newly synthesized AMO protein and which during further cultivation developed in normal peroxisomes. Concurrently, the remaining portion of these structures was rapidly degraded. These findings indicate that peroxisomal remnants in pex5 cells can develop into peroxisomes. Also, as for normal peroxisomes in *H. polymorpha*, apparently a minor portion of these structures actually take part in the development of these organelles.

In eukaryotic cells most organelar proteins are synthesized in the cytosol and sorted to their target compartments by unique signals. These signals are recognized by specific receptor proteins, which are either located in the cytosol (e.g. signal recognition particle for the endoplasmic reticulum (1)) or bound to the membrane of the organelle (e.g. TOM proteins at the mitochondrial outer membrane (2)).

Proteins destined for the peroxisomal matrix are encoded by nuclear genes and synthesized in the cytosol. Specific topogenic signals have been detected at the extreme C terminus (the PTS1) or in the N terminus of the protein (termed PTS2) (3). It is now widely accepted that the PTS1 and PTS2 are recognized in the cytosol by their specific receptor molecules, Pex5p and Pex7p, and that this interaction in fact initiates the actual protein import process. Both pathways converge at the docking site, consisting of at least Pex13p, Pex14p, and Pex17p. Maintenance of the correct stoichiometry of the components of the docking complex seems to be important since relatively minor alterations in the level of Pex14p already affect normal import (4). Another peroxin complex, consisting of the two zinc-binding proteins Pex10p and Pex12p, most likely binds Pex5p at a later stage after docking (5, 6). It can be envisaged that both complexes reflect an import cascade with different affinities for the receptor-cargo complex. These interactions could lead to the dissociation of the cargo from the receptor, and the hand over of the cargo molecule to the translocation machinery. Alternatively, the receptor-cargo complex may be completely translocated across the peroxisomal membrane, followed by dissociation of the complex in the peroxisomal lumen (7). Indeed, in *Hansenula polymorpha* we have obtained evidence for this pathway in which Pex4p, an ubiquitin-conjugating enzyme, plays a role (8).

In yeast, deletion of the PEX5 gene results in a general block in PT51 matrix protein import. However, at PT52 protein-inducing conditions (e.g. oleate, in case of *H. polymorpha* primary amines), small peroxisomes were observed in these mutants since both the PT52 import and the insertion of peroxisomal membrane proteins are independent of Pex5p function (9, 10). When *H. polymorpha* pex5 mutant cells were grown in carbon-limited chemostats on glucose/choline, conditions that strongly induce peroxisome proliferation, a few small peroxisomes were observed. These organelles contained the PT52 protein amine oxidase (AMO, induced by choline), whereas the PT51 proteins alcohol oxidase, dihydroxyacetone synthase, and catalase were mislocalized in the cytosol (11). However, under AMO-repressing conditions during growth of cells on methanol/ammonium sulfate, peroxisomes are fully absent in pex5 cells (12). Here we analyzed whether such cells contained peroxisomal membrane structures and normal peroxisomal membrane proteins. We have used the integral peroxisomal membrane protein Pex10p, carrying a C-terminal Myc tag (Pex10p.myc) as a marker protein for the detection of these structures. Also, we studied whether peroxisome development upon induction of AMO protein initiates from such structures or whether alternative mechanism existed. The results of these studies are the focus of this paper.
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**MATERIALS AND METHODS**

**Organisms and Growth Conditions—**The *H. polymorpha* strains used in this study are listed in Table I. The cells were cultivated in mineral media (13) containing 0.5% (v/v) glucose, 0.5% (v/v) ethanol, or a mixture of 0.15% (v/v) glycerol and 0.2% (v/v) methanol as carbon sources supplemented with 0.25% (w/v) ammonium sulfate, 0.25% (w/v) ethylamine, or 0.25% (w/v) methyamine as respective nitrogen sources at 37 °C. For the selection of mutants, solid YND media were used, containing 2% (w/v) yeast nitrogen base (Difeco) and 1.5% (w/v) agar.

**Construction of *H. polymorpha* WT and pex5 Strains, Producing Pex10p.myc—**A leucine-auxotrophic PEX5 deletion strain, suitable for transformation with ScLEU2-based plasmids (14), was constructed as follows. By PCR, a BamHI site was introduced downstream from the 5'-untranslated region of PEX5, using the primer 5'-GGG GAA TCC ATT GAT GGT TTT TGCA AG-3' (BamHI site underlined). A 2.2-kb BglII (sticky ends filled-in using Klenow enzyme)-BamHI DNA fragment, containing the HpURA3 gene (15), was inserted between the engineered BamHI site and the genomic NruI site at position +1504 in the PEX5- open reading frame. By using BsiWI and NcoI, a PEX5 disruption cassette was obtained, containing the HpURA3 gene flanked by 0.2-kb PEX5-5'-untranslated region and 0.3-kb PEX5 open reading frame and 3'-untranslated region, which was used to transform *H. polymorpha*NCYC495 leu1 ura1. One transformant was selected which was uracil prototrophic, methanol utilization-deficient and showed the expected genomics changes after Southern blot analysis.

The chimeric gene, encoding Pex10p.myc, was constructed by PCR-mediated amplification of the PEX10 gene of pET4 (Table I) using the following primers (oligonucleotides were obtained from Life Technologies, Inc.): upstream 5'-AGA GGA TCC ATG TTA AAG CTT CTC AGA CAA CTT TCA AGA-3' (BamHI site underlined) and downstream 5'-AGA GAA TCC TTA CAA CTT CTC TGG TCA GAG CAA CAG CTC GCT-3' (EcoRI site underlined and the nucleotides coding for the Myc epitope (16) are depicted in italics). The PCR product was ligated as a 0.94-kb BamHI-EcoRI fragment in pBluescript II KS+, digested with BamHI and EcoRI. The resulting plasmid, pFAS01, was digested with BamHI and XhoI, and the 0.94-kb fragment was inserted into BamHI-SalI-digested pHIPX4 (Table I) under control of the alcohol oxidase promoter, resulting in plasmid pHIPX4-PEX10.MYC. The plasmid pHIPX4-PEX10.MYC was introduced into a *H. polymorpha* pex10-1 mutant strain by electroporation (17) to check the functionality of the hybrid protein. Also it was linearized with StuI and integrated at the *P*aox locus of *H. polymorpha* pex strain (17). Transformants were selected on YND plates and site-directed integration of the plasmid at the *P*aox locus was confirmed by Southern blot analysis, pex5 strains containing two copies of the pHIPX4 derivatives were selected for further studies.

**Biochemical Methods—**Crude extracts of *H. polymorpha* cells were prepared as detailed by Baerends et al. (18). Cell fractionation procedures (19) and cytochrome c oxidase activities measurements (20) were performed as described. Protein concentrations were determined using the Bio-Rad Protein Assay system (Bio-Rad) using bovine serum albumin as a standard. SDS-polyacrylamide gel electrophoresis (21) was carried out as described, and gels were subjected to Western blotting (22). Nitrocellulose blots were decorated using specific polyclonal antibodies against various *H. polymorpha* proteins.

**Electron Microscopy—**Cells were fixed and prepared for electron microscopy as described previously (23). Immunolabeling was performed on ultrathin sections of unicyrilled-embedded cells, using specific antibodies against various proteins and gold-conjugated goat anti-rabbit or goat anti-mouse antibodies (23).

**RESULTS**

Characterization of Peroxisomal Structures in *H. polymorpha* pex5 Cells—The presence of putative peroxisomal membrane structures in *H. polymorpha* pex5 cells was first determined in cells, grown at moderate peroxisome-induction conditions on ethanol/ammonium sulfate. Under these conditions, WT cells contain few organelles, characterized by the presence of isocitrate lyase and malate synthase, key enzymes of the glyoxylate cycle. Electron microscopic analysis showed that in such pex5 cells peroxisomal structures were not detectable despite extensive research also including analysis of serial sections (Fig. 1A). This was remarkable since the levels of the membrane protein Pex14p was equal to WT control cells and over 5-fold enhanced, compared with cells grown on glucose (Fig. 2). Pex3p was also detectable but not enhanced compared with glucose-grown WT cells, independent whether ammonium sulfate or ethylamine was used as the nitrogen source (Fig. 2). Ethylamine metabolism is mediated by AMO, a PTS2 protein of *H. polymorpha*. At the morphological level, however, replacement of ammonium sulfate by ethylamine led to the rapid development of peroxisomes in ethanol-grown pex5 cells. Typically, one or very few peroxisomes developed within 1 h of cultivation in the new ethylamine environment (Fig. 1B). As expected, these organelles were characterized by the presence of AMO protein. This raises the question of the origin of these new organelles. A reason for the virtual absence of peroxisomal remnants in ethanol/ammonium sulfate-grown cells could be that these structures were below the limit of detection of electron microscopy. To enhance the level of peroxisomal proteins, pex5 cells were subsequently grown on glycerol/methanol/ammonium sulfate. Since *H. polymorpha* pex strains cannot grow on methanol alone (24), glycerol was applied as the growth substrate using methanol as additional energy source and peroxisome inducer. In this way growth conditions are created that are similar for both pex5 and WT control cells, at least until the mid-exponential growth phase. WT cells grown under these conditions contain several large peroxisomes (25). Western blot analysis revealed that the levels of Pex3p and Pex14p in glycerol/methanol-grown pex5 and WT cells were comparable but strongly enhanced compared with the levels in ethanol- or glucose-grown cells (Fig. 2). This indicated that deletion of the PEX5 gene did not affect the amounts of peroxisomal membrane proteins in the cells.

Electron microscopic analysis of glycerol/methanol/ammonium sulfate grown pex5 cells revealed the presence of a cluster of membranous structures (Fig. 1C). These structures appar-
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A. overall morphology of ethanol/ammonium sulfate-grown H. polymorpha pex5 cells. In these cells no peroxisomal structures are visible. After 60 min of cultivation on ethanol/ethanolamine, a small peroxisomal remnant was observed (B). C, overall morphology of methanol/ammonium sulfate-induced pex5 cells that contain few small peroxisomal remnants (arrow = peroxisomal remnants), characterized by the presence of Pex14p, as was evident upon labeling using α-Pex14p antibodies (C, inset). On glycerol/methanol/methanolamine the cells contained few peroxisomes (arrow) that were formed within 2 h (D). By using specific α-AMO antibodies, these structures were labeled (D, inset). A–D, KMnO4 fixation; insets, glutaraldehyde fixation.

To test the function of Pex10p.myc in H. polymorpha, we introduced the PEX10.MYC expression cassette in the original H. polymorpha pex10-1 mutant (27). The resulting transformant grew on methanol at WT rates (data not shown). This suggested that Pex10p.myc functionally complemented the peroxisomal defect in pex10-1 cells, since any defect in PTS1 (and thus alcohol oxidase) import would prevent growth of cells on methanol (24). This was confirmed by electron microscopic analysis, which revealed that methanol-grown cells of this transformant contained normal peroxisomes. Moreover, immunocytochemistry showed that the location of Myc protein was confined to peroxisomal membranes, confirming that indeed Pex10p.myc was normally sorted to the peroxisomal membrane (not shown).

Next, we introduced the P_{A09}PEX10.MYC cassette in the H. polymorpha pex5 strain. A transformant with two copies of P_{A09}PEX10.MYC integrated in the genome was selected and cultivated in glycerol/methanol/ammonium sulfate-containing media to induce Pex10p.myc and peroxisomal protein synthesis. Electron microscopy demonstrated that these cells contained clusters of membranous structures (Fig. 3A) that behaved like normal cell constituents in that they were donated to newly developing buds (not shown; compare Fig. 9A). These structures were the sole sites of gold labeling in immunocytochemical experiments, using α-Myc antibodies (Fig. 3B). Similar results were obtained when α-Pex3p, α-Pex10p, and α-Pex14p antibodies were used (shown for Pex10p; Fig. 3C), indicating that these structures represent peroxisomal membrane structures. Analysis of series of consecutive sections revealed that these clusters were invariably associated with the nucleus and generally include over 20 small vesicles (Fig. 4). Taken together, these data imply that the Pex10p.myc marker protein had specifically accumulated at the peroxiso-
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normal membrane remnants in glycerol/methanol/ammonium sulfate-grown pex5 cells (Fig. 1C) and, thus, is a suitable component to tag these structures.

The co-localization of Pex10p.myc with the peroxisomal membrane remnants in pex5:PAOXPEX10.MYC cells was confirmed by cell fractionation experiments. A post-nuclear supernatant (PNS), prepared from homogenized protoplasts of glycerol/methanol/ammonium sulfate-grown cells of this strain, was subjected to sucrose density centrifugation (29).

Analysis of the various fractions obtained indicated that the bulk of the cellular protein was located at the top of the gradient, representing soluble, cytosolic proteins. Western blot analysis of these fractions revealed that Pex10p.myc migrated to a density corresponding to ~35% (w/w) sucrose (Fig. 5B, fractions 16–18). A similar distribution pattern was observed for Pex14p. At 53% (w/w) sucrose, where intact peroxisomes of H. polymorpha wild type cells normally sediment (8, 29), Pex10p.myc and Pex14p were not detectable. The protein peak at ~42% (w/w) sucrose contains mitochondria judged from the distribution of the mitochondrial marker enzyme cytochrome c oxidase. The minor cytochrome c oxidase peak, present at 30% (w/w) sucrose, most probably reflects fragmented and subsequently sealed protoplasts that were still present in the PNS.

From this we conclude that the peroxisomal membrane structures present in glycerol/methanol/ammonium sulfate-grown H. polymorpha pex5:PAOXPEX10.MYC cells sediment to a density of 35% (w/w) sucrose. These findings are in line with earlier data that revealed that peroxisomal remnants from other H. polymorpha pex mutants defective in matrix protein import also sedimented to relatively low densities in sucrose gradients (e.g. pex4 (8), pex8 (30), and pex14 (29)).

A Minor Number of the Peroxisomal Membrane Structures Accumulate Newly Synthesized AMO to Develop into Normal Peroxisomes—The approach to analyze whether the peroxisomal remnants in pex5:PAOXPEX10.MYC cells can develop into normal peroxisomes includes the initial induction of these structures by methanol, followed by the synthesis of the peroxisomal PTS2 protein AMO under conditions that the synthesis of Pex10p.myc is fully repressed (by glucose).

Pex5:PAOXPEX10.MYC cells were pre-cultivated on glycerol/methanol/ammonium sulfate to induce the synthesis of Pex10p.myc containing membrane remnants. These cells were harvested by centrifugation and incubated for 30 min at 37°C in mineral medium, lacking the C- and N-source to remove any Pex10p.myc mRNAs (31). Subsequently, methyamine (as N-source) was added to this medium to induce AMO synthesis together with glucose (as C-source) to concurrently block the synthesis of Pex10p.myc. At various time points after addition of these substrates, cells were harvested and analyzed. The protein patterns of AMO, Pex10p.myc, and Pex3p were determined by Western blotting (Fig. 6). As evident in Fig. 6A, AMO was rapidly induced within 2 h after the shift to glucose/methyamine medium. Identical blots, decorated with antibodies against the Myc epitope, revealed a drastic decrease in Pex10p.myc levels in time. Within 2 h after the shift the level of Pex10p-Myc had decreased to ~20% of the level in the inoculum cells. A similar reduction was observed for Pex3p (Fig. 6A).

To analyze the initial events of peroxisome formation, a post-nuclear supernatant (PNS) of glycerol/methanol/ammonium sulfate-grown pex5:PAOXPEX10.MYC cells, transferred for 2 h to glucose/methyamine media, was subjected to sucrose density centrifugation. Analysis of the various fractions obtained did not allow the unequivocal demonstration of the Myc epitope, most likely due to the diminished levels of Pex10p.myc protein. For this reason we tried to enrich the peroxisomal fraction by conventional differential fractionation of the PNS. Western analysis revealed that Pex10p.myc pelleted in the 30,000 × g (P3) organellar fraction (Fig. 7, inset). Next, this Pex10p.myc-enriched organellar fraction was subjected to sucrose density centrifugation (Fig. 7). The major protein peak present in this gradient (Fig. 7A, fractions 12 and 13) represents mitochondria, judged from the distribution of cytochrome c oxidase activity. Western blot analysis of the various fractions collected from the gradient revealed that bulk of the Pex10p.myc had migrated to a density corresponding to ~48% (w/w) sucrose (Fig. 7B, fractions 9 and 10). The major portion of the peroxisomal membrane protein Pex14p was found in the same fractions. Hence, these peroxisomal proteins migrated to higher densities than the peroxisomal remnants of methanol-induced pex5:PAOXPEX10.MYC cells (~35% (w/w) sucrose, Fig. 5) but not yet to the density of mature peroxisomes of WT methanol-grown H. polymorpha cells (~53% (w/w) sucrose (8, 29)). This location most probably reflects the relatively smaller size of the organelles, compared with WT peroxisomes. The bulk of the AMO protein co-fractionated with Pex10p.myc and Pex14p in fractions 9–10, suggesting that AMO protein may be present in the same compartment as Pex10p.myc and Pex14p.
Interestingly, a portion of Pex10p.myc was also found in fraction 8, which contained only a small amount of Pex14p. Hence these fractions might contain a slightly different type of structures. In addition, almost no Pex10.Myc was detected at lower densities (fractions 11–12), which did, however, contain Pex14p and AMO. Possibly these fractions represent very small, newly formed peroxisomes.

The morphological adaptations of pex5::PAOXPEX10.MYC cells shifted from glycerol/methanol/ammonium sulfate into fresh glucose/methylamine media were analyzed by electron microscopy (Fig. 8). After 2 h, the fraction of peroxisomal membrane remnants had strongly reduced. Peroxisomal membrane structures could still be detected (Fig. 8, B and C) but were frequently seen surrounded by additional layers of membranes, which is a typical feature for vacuolar uptake of peroxisomes to be degraded (Fig. 8D). Immunocytochemistry demonstrated that Pex10p.myc was still detectable on the remaining peroxisomes (Fig. 8, inset). After 4 h of incubation on glucose/methylamine, few small peroxisomes could be observed (Fig. 8D), which contained the PTS2 protein AMO, as was evident from labeling experiments using specific α-AMO antibodies (Fig. 8D, inset). Apart from these peroxisomes, membrane remnant structures were no longer detectable. Remarkably, the number of growing organelles was low compared with the number of vesicles originally present in the cells. Also, after a shift of glycerol/methanol/ammonium sulfate to glycerol/methanol/methylamine (thus only changing the nitrogen source), only few (up to maximally 5–6) organelles developed (data not shown).

Double labeling of ultrathin sections of the cells using α-Myc and α-AMO antibodies revealed that Pex10p.myc and AMO protein co-localized on the peroxisomal structures (Fig. 9). This strongly suggests that the peroxisomal membrane remnants in glycerol/methanol/ammonium sulfate grown pex5::PAOXPEX10.MYC cells had accumulated newly synthesized AMO.

The biochemical and morphological observations indicate that our shift experiment probably had initiated two opposite processes that occurred simultaneously, glucose-induced degradation of peroxisomal membrane remnants (32) and peroxisome development due to synthesis and import of newly induced AMO protein (33). To analyze the glucose-induced degradation process separately from the import process of AMO, glycerol/methanol-induced pex5::PAOXPEX10.MYC cells were shifted to glucose/ammonium sulfate-containing media. Crude extracts were prepared at various time points after the shift and were analyzed. The protein patterns of Pex10p.myc and Pex3p were determined by Western blotting experiments (Fig. 6B). These data indicated that the levels of both proteins rapidly decreased in time (Fig. 6B), indicating that the Pex10p.myc-tagged peroxisomal membrane remnants are subject to degradation under these conditions. This degradation process resembles the selective autophagy that has been described for peroxisomal remnants in other H. polymorpha pex mutants (34).
Notably, in glucose/methylamine media the Pex10p.myc and Pex3p reduction is slightly less than in glucose/ammonium sulfate cultures lending support to the view that indeed fraction of these membranes takes part in new peroxisome development. Another indication for this is the morphological observation that in other H. polymorpha pex deletion strains re-introduction of the corresponding gene invariably results in the development of one (or infrequently few) new small peroxisome in the vicinity of the cell wall.

**DISCUSSION**

In this paper we provide evidence that peroxisomal remnants present in methanol-induced H. polymorpha pex5 cells may develop into normal peroxisomes upon the synthesis of the key enzyme in amine metabolism, the PTS2 protein AMO. The generally accepted view is that peroxisomes develop from already existing organelles by growth and fission (35). Our data indicate that this rule can now be extended in that peroxisomes may also develop from remaining residual structures in pex5 mutant cells. Whether this is also true for the other pex mutants, e.g. after re-introduction of the defective gene, is not yet clear. We have demonstrated that the peroxisomal remnants (“ghosts”) in H. polymorpha pex4 and pex14 cells can develop into normal peroxisomes upon overexpression of the PEX5 gene (8, 29). Peroxisome re-assembly also occurs in mutant strains, which are known to lack any peroxisomal membrane remnants, upon re-introduction of the defective gene. Examples of this include Pichia pastoris pex3 (36) and Saccharomyces cerevisiae pex19 (37) null mutants, H. polymorpha pex3 (38) and ts6 (31) mutants, as well as human pex16 mutants (39). This indicates that alternative modes of peroxisome assembly may exist. It has been suggested that new peroxisomes may arise from the endomembrane system, e.g. from specialized regions of the ER.
South and Gould (39) presented an interesting model for peroxisome re-assembly in human pex16 mutant cells and suggested that they may arise by conversion from an unknown pre-peroxisomal structure, mediated by Pex16p. It remains unclear whether this model reflects the WT situation or that it serves as rescue machinery to peroxisome formation. However, as in S. cerevisiae a PEX16 homologue has not been identified yet, and the human model may not be generally valid. Additional research is therefore required for elucidating this problem in both human and yeast.

Characteristic for various pex5 mutants (e.g. of P. pastoris (9), S. cerevisiae (40), Yarrowia lipolytica (41–43)) is the cytosolic location of PTS1 proteins together with the presence of peroxisomal membrane remnants or, in case PTS2 proteins are synthesized (e.g. during growth of yeast on oleate), small peroxisomes. A similar phenotype is described for H. polymorpha pex5 cells, grown under conditions that both PTS1 and PTS2 proteins are induced (11). Conversely, in H. polymorpha pex5 cells, grown on glucose or ethanol in the presence of ammonium sulfate as sole nitrogen source (conditions that fully repress synthesis of the PTS2 protein AMO), peroxisomal membrane remnants were below the limit of detection. On the other hand, peroxisomal remnants were detectable in H. polymorpha pex5, when grown at high peroxisome induction conditions on methanol. Therefore, the proliferation of these structures is dependent on the growth conditions and is most likely related to the ultimate level of essential peroxisomal membrane proteins.

The membrane remnants in H. polymorpha pex5 cells display several characteristics, typical for normal WT peroxisomes. They are the target for artificially produced Pex10p.myc, proliferate un-

**Figure 7.** Sucrose gradient prepared from an organellar pellet (P3) obtained from homogenized, glycerol/methanol/ammonium sulfate-induced pex5. P_{Aox}PEX10.MYC cells shifted to glucose/methylamine-containing medium (A), showing the distribution of the protein (C) in mg ml^{-1}, sucrose (+) expressed as percentages (w/w), and the activity of the mitochondrial enzyme cytochrome c oxidase (▲) expressed as percentages of the activity in the peak fraction, which was arbitrarily set at 100. After differential centrifugation of the PNS, Pex10p.myc is present in the 30,000 × g pellet (inset). P3 and S3 designate the 30,000 × g pellet and supernatant, respectively, while P4 and S4 represents the 100,000 × g pellet and supernatant respectively. B, Western blot analysis of the sucrose gradient demonstrate the distribution of the Myc epitope, Pex14p, and AMO protein. Pex10p.myc was detected in fractions 8–10 corresponding to a density of ~48% (w/w) sucrose. The peak of Pex14p colocalized with Pex10p.myc. Bulk of the AMO protein was detected in fractions 9–10. Equal volumes were loaded per lane.
under specific growth conditions (this study), and are susceptible to selective degradation (32, 34).

Our data demonstrated that newly produced AMO protein accumulated at the Pex10p.myc-containing, pre-existing membrane structures, originally present in the methanol/ammonium sulfate-grown cells. Since the new peroxisomes developed

FIG. 8. Morphological analysis of glycerol/methanol/ammonium sulfate-grown pex5. H. polymorpha PEX10-MYC cells after the shift to glucose/methylamine-containing medium. Before the shift, these cells contained few small peroxisomal membranes, indicated by the arrow (A). After incubation for 2 h on glucose/methylamine, these membrane structures could still be detected (C), although a small portion was surrounded by additional membrane layers and degraded by the vacuole (B, indicated by the arrow). Labeling of ultrathin sections of these cells using specific antibodies against the Myc-epitope showed that the peroxisomal membrane structures contained Pex10p-Myc (C, inset). After a longer incubation period (4 h), a few small peroxisomes could be observed (D), which contained the PTS2 protein AMO, as is evident from the immunolabeling using specific antibodies against this protein (D, inset). The abbreviations used are: M, mitochondrion; N, nucleus; V, vacuole. The electron micrographs (A–D) are from KMnO4-fixed cells and the insets are from glutaraldehyde-fixed cells. The bar represents 0.5 \( \mu \)m, and the bars in the insets of C and D represent 0.25 \( \mu \)m.

FIG. 9. Immunocytochemical demonstration that the pre-existing peroxisomal remnants containing Pex10p-Myc are the target for newly synthesized AMO. Glycerol/methanol/ammonium sulfate-grown pex5::Pex5PEX10-MYC cells were transferred to glucose/methylamine medium and cultivated for 2 h. A double labeling experiment using antibodies against Myc (10 nm gold-conjugated goat anti-mouse) and AMO (5 nm gold-conjugated goat anti-rabbit) showed that Pex10p-Myc and AMO co-localized in the peroxisomal remnants (arrow = peroxisomal remnants). A small portion of these peroxisomal remnants was transferred to the bud of this cell. A more detailed image of the co-localization of Pex10p-Myc and AMO is provided by the inset (the arrow indicates the 5 nm gold-conjugated goat anti-rabbit). Electron micrographs were taken of glutaraldehyde-fixed cells, poststained with uranyl acetate, unless otherwise indicated. The abbreviations used are: M, mitochondrion; N, nucleus; V, vacuole. The bar represents 0.5 \( \mu \)m, and the bar in the inset represents 0.25 \( \mu \)m.
under conditions in which the synthesis of Pex10p.myc was fully repressed, the co-localization of Pex10p and AMO suggests that these organelles originate from Pex10p.myc-containing structures. Whether the development of the peroxisomal membrane remnants involved fusion with small pre-peroxisomal AMO-containing vesicles, as for instance demonstrated for matrix protein import in Y. lipolytica, is yet unknown (44).

The possibility that the failure of a portion of the peroxisomal structures to import AMO is due to the presence of Pex10p.myc is not very likely. We showed that Pex10p.myc is functional and can complement the pex10 mutant phenotype.

In conclusion, our data suggest that peroxisomal membrane remnant structures present in glycerol/methanol/ammonium sulfate-grown \textit{H. polymorpha} pex5 cells can develop into peroxisomes upon subsequent synthesis of the PTS2 protein AMO. Notably, this also happens when the cells are placed in conditions (shift from glycerol/methanol/ammonium sulfate to glucose/methylamine) for assistance in electron microscopy.

Whether the development of the peroxisomal remnants were able to gradually take up PTS1 proteins in reappeared after microinjection of Pex5p together with green fluorescent protein fused to a PTS1 signal. They observed green fluorescent protein fluorescence in particulate structures in which PMP70 co-localized, suggesting that all peroxisomal remnants were able to gradually take up PTS1 proteins in time. The reasons for the apparent discrepancy with the structures in \textit{H. polymorpha} pex5 cells are unknown. A possible explanation may be related to the fact that microinjection in fact confronts the cell with an excess of matrix proteins. In view of the finding that under WT conditions the maximal import capacity of peroxisomes apparently is not fully used (47), it can be envisaged that the sudden excessive amounts of matrix proteins disturbs the normal balance between matrix protein synthesis and import capacity thus leading to additional protein import in existing organelles. However, more research is required to solve this specific question.

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REFERENCES

1. Brodsky, J. L. (1998) in \textit{Protein Targeting and Translocation} (Phoenix, D. A., ed) pp. 169–191, Portland Press Ltd., London

2. Hovius, R. (1998) in \textit{Protein Targeting and Translocation} (Phoenix, D. A., ed) pp. 231–248, Portland Press Ltd., London

3. Erdmann, R., Veenhuis, M., and Kunau, W. H. (1997) \textit{Trends Cell Biol} 7, 400–407

4. Komori, M., Rasmussen, S. W., Kiel, J. A. K. W., Baerends, R. J. S., Cregg, J. M., van der Klei, I. J., and Veenhuis, M. (1997) \textit{EMBO J} 16, 44–53

5. Chang, C. C., Warren, D. S., Lametschwandtner, G., Christmann, J., Hartig, A., and Harada, W. H., and Erdmann, R. (1998) \textit{Mol. Cell. Biol} 18, 616–628

6. Baerends, R. J. S., Rasmussen, S. W., Hilbrands, R. E., van der Heide, M., Cregg, J. M., Kiel, J. A. K. W., and van der Klei, I. J., and Veenhuis, M. (1999) \textit{Proc. Natl. Acad. Sci. U. S. A} 96, 1509–1507

7. South, S. T., and Gould, S. J. (1998) \textit{Trends Cell Biol} 18, 887–894

8. van der Klei, I. J., Hilbrands, R. E., Kiel, J. A. K. W., Rasmussen, S. W., Cregg, J. M., and Veenhuis, M. (1998) \textit{EMBO J} 17, 3608–3618

9. Malorail, R., Monosov, E., and Subramani, S. (1993) \textit{J. Cell Biol} 121, 761–774

10. van der Klei, I. J., Anke, M., and Veenhuis, M. (1994) \textit{Proc. Natl. Acad. Sci. U. S. A} 91, 1145–1149

11. van der Klei, I. J., and Veenhuis, M. (1999) \textit{J. Biol. Chem} 274, 17229–17236

12. Nitsch, W. A., and Kram, A. M., Kiel, J. A. K. W., and van der Klei, I. J., and Veenhuis, M. (2000) \textit{J. Biol. Chem} 275, 9896–9995

13. van der Klei, I. J., van der Heide, M., Baerends, R. J. S., Rechberger, K. B., Nicolay, K., Kiel, J. A. K. W., and Veenhuis, M. (1998) \textit{Curr. Genet} 34, 1–11

14. Dough, A. C., van der Klei, M., de Koning, W., Evers, M. E., and Harder, W. (1985) \textit{Arch. Microbiol} 143, 237–243

15. Leuvecchi, A. M. (1990) \textit{Nature} 345, 685

16. Kjyshe-Andersen, J. (1984) \textit{Biochem. Biophys. Methods} 10, 203–209

17. Waterham, R. H., Titorenko, V. I., Haima, P., Cregg, J. M., Harder, W., and Veenhuis, M. (1999) \textit{J. Cell Biol} 127, 737–749

18. van der Klei, I. J., Harder, W., and Veenhuis, M. (1991) \textit{Arch. Microbiol} 156, 15–23

19. Veenhuis, M., and Harder, W. (1987) in \textit{Peroxisomes in Biology and Medicine} (Fahimi, D., and Sies, H., eds) pp. 436–457, Springer Verlag, Berlin-Heidelberg

20. Faber, K. N., Haima, P., Gietl, C., Harder, W., AB, G., and Veenhuis, M. (1994) \textit{Ann. Rev. Genet} 28, 505–505

21. Kragler, F., Lametschwandtner, G., Christmann, J., Hartig, A., and Harada, W. H., and Erdmann, R. (1995) \textit{Biochem. Biophys. Res. Commun} 209, 15–15

22. Veenhuis, M., Komori, M., Salomons, F. A., Baerends, R. E., Hult, H., Baerends, R. J. S., Kiel, J. A. K. W., and van der Klei, I. J. (1996) \textit{FEBS Lett} 383, 114–118

23. Laserov, P. B., and Fujiy, K. (1985) \textit{Annu. Rev. Cell Biol} 1, 489–530

24. Wiemer, E. A. C., Nuttley, W. M., Bertolaet, B. L., Li, X., Francke, G., Cheese, H., and Rachubinski, R. A. (1995) \textit{Proc. Natl. Acad. Sci. U. S. A} 92, 9986–9995

25. Bruscato, C., Frager, K., Simon, M. M., Schuster, T., and Hartig, A. (1994) \textit{Biochem. Biophys. Res. Commun} 204, 1016–1022

26. Szilard, R. K., Titorenko, V. I., Veenhuis, M., and Rachubinski, R. A. (1995) \textit{Cell Biol} 131, 1453–1456

27. Kragler, F., Lametschwandtner, G., Christmann, J., Hartig, A., and Harada, W. (1996) \textit{Proc. Natl. Acad. Sci. U. S. A} 93, 13336–13343

28. Wiemer, E. A. C., Nuttley, W. M., Bertolaet, B. L., Li, X., Francke, U., Wheelock, M. J., Anne, U. K., Johnson, K. R., and Subramani, S. (1995) \textit{J. Cell Biol} 130, 51–65

29. Titorenko, V. I., Chan, H., and Rachubinski, R. A. (2000) \textit{J. Cell Biol} 148, 29–44

30. Veenhuis, M., Zwart, K., and Harder, W. (1981) \textit{Arch. Microbiol} 129, 35–41

31. Yasumatsu, M., Hashiguchi, T., Imamuka, T., Tsukamoto, T., and Osami, T. (1999) \textit{J. Biol. Chem} 274, 35283–35296

32. Boggenkamp, R., Didion, T., and Kowallik, K. V. (1989) \textit{Mol. Cell. Biol} 9, 988–994

33. Gleeson, M. A. G., and Sudbery, P. E. (1988) \textit{Yeast} 4, 293–303