Yeast pex1 cells contain peroxisomal ghosts that import matrix proteins upon reintroduction of Pex1

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Introduction

Peroxisomes are ubiquitous and versatile cell organelles that are involved in a large variety of metabolic pathways. Conserved functions are hydrogen peroxide metabolism and fatty acid β-oxidation (Smith and Aitchison, 2013). Peroxisomes proliferate in response to various internal or external cues, thus ensuring that organelle abundance continuously adapts to cellular needs (Mast et al., 2015).

In higher eukaryotes, peroxisome deficiency is lethal (Fujiki et al., 2012; Hu et al., 2012). However, yeast mutants that show a defect in peroxisome biogenesis are normally viable and capable to grow on media containing glucose, but not on substrates that are metabolized by peroxisomal enzymes (e.g., oleic acid and methanol). This unique property enabled using simple yeast genetic screens to identify genes (PEX genes) that play a role in peroxisome formation (Erdmann and Kunau, 1992).

Upon reintroduction of the deleted PEX genes in yeast peroxisome-deficient (pex) mutants, peroxisomes invariably reappear. So far, different mechanisms of peroxisome reintroduction have been described. Deletion of a PEX gene encoding a protein involved in peroxisomal matrix protein import (e.g., Pex14) results in cells containing peroxisomal membrane remnant structures, designated ghosts, in conjunction with mislocalization of matrix proteins in the cytosol. Peroxisomal membrane proteins (PMPs) are normally present in these ghosts because sorting and insertion of PMPs is independent of matrix protein import. Upon reintroduction of the corresponding PEX gene, these preexisting ghosts develop into normal peroxisomes by importing matrix proteins.

For a long time, it was generally accepted that yeast mutants affected in peroxisomal membrane formation (i.e., pex3 or pex19 mutants) lack peroxisomal membrane remnants (Hettema et al., 2000). However, we recently showed that yeast pex3 and pex19 cells do contain small preperoxisomal vesicles (PPVs), which contain only a subset of PMPs, whereas other PMPs are mislocalized and very instable (Knoops et al., 2014). Upon reintroduction of the corresponding genes, the latter PMPs are also sorted to the PPVs, which results in the formation of a functional peroxisomal importomer and hence matrix protein import, thus leading to the maturation of PPVs into normal peroxisomes.

Recently, an alternative pathway of peroxisome reintroduction has been described for yeast pex1 and pex6 cells. According to this model, two types of ER-derived vesicles fuse upon reintroduction of Pex1 or Pex6, before the formation of normal peroxisomes (van der Zand et al., 2012). These vesicles each carry half a peroxisomal translocon complex, namely either proteins of the receptor docking complex (Pex13 and Pex14) or the RING complex (Pex2, Pex10, and Pex12) together with Pex11. This would imply that in yeast pex1 and pex6 cells, two types of biochemically distinct vesicles accumulate. Upon Pex1 or Pex6 reintroduction, heterotypical fusion of these vesicles would lead to the assembly of the full peroxisomal translocon, thus allowing PMP import.

Here we analyzed the ultrastructure of yeast pex1 and pex6 mutant cells and the mode of peroxisome reintroduction in depth using advanced, high-resolution microscopy techniques, i.e., electron tomography (ET), immunolabeling, and...
correlative light and electron microscopy (CLEM). The results of these studies are contained in this paper.

Results and discussion

Components of the docking and RING complex colocalize in pex1 and pex6 cells

We first analyzed the localization of PMPs of the docking and RING complex by fluorescence microscopy (FM). PMPs were chromosomally tagged to create endogenously expressed C-terminal fusions with the monomeric red fluorescent protein mCherry (Pex2 and Pex10) or monomeric green fluorescent protein mGFP (Pex13 and Pex14). FM revealed that the fluorescent spots of the docking and RING proteins overlapped in *Saccharomyces cerevisiae* BY4742 *pex1* and *pex6* cells, similar as observed in wild-type (WT) controls (Fig. 1 A and S1 A). In addition, the spots of Pex11-mCherry, a PMP involved in peroxisome fission, coincided with Pex14-mGFP spots (Fig. S1 A).

To seek further support for this PMP colocalization, we performed quantitative FM analysis. All mCherry spots present in 25 randomly acquired FM images were selected, and their distance to the closest mGFP spot was measured (Fig. S1 B). High colocalization values were obtained for all PMP pairs tested in glucose-grown cells of BY4742 *pex1*, *pex6*, and WT controls (Fig. 1 B). Similar results (Figs. 1 B and S1 A) were obtained for a *pex1* *pex6* double-deletion strain, for cells grown on oleic acid, or when WT and mutant strains of the FY1679 parental strain were used (as was used by van der Zand et al., 2012; Fig. S1 A). These data suggest that all PMPs analyzed colocalize in the absence of Pex1 or Pex6 as they do in WT control cells.

Quantification of the number of Pex14-GFP spots revealed a mean number of 1.5 ± 0.04 (n = 441) per cell in glucose-grown *pex1* cells, which is similar to what was previously reported (see Fig. 3 A in van der Zand et al., 2012). This number did not increase significantly upon incubation of cells in oleic acid medium (mean number of 1.6 ± 0.20 Pex14-mGFP spots per cell; n = 540).

*pex1* and *pex6* cells contain peroxisomal membrane ghosts

To identify the nature of the structures to which the fluorescent PMPs were localized, we performed CLEM using cryosections of *pex1* cells. CLEM of oleic acid-grown *pex1* cells producing Pex10-mCherry revealed that at the site of a single red fluorescent spot, clusters of small membrane structures were present (Fig. 2 A). Immunocytochemistry using specific antibodies against Pex14 revealed that these membranes contained Pex14, suggesting that they represented peroxisomal membranes (Fig. 3 E). Similar Pex14-containing structures were observed in oleic acid-induced *pex6* cells (Fig. 3 F). These structures were not detected in WT controls, in which α-Pex14–specific label was confined to peroxisomes (Fig. 3 D). To test whether the Pex14-labeled structures also contained the RING protein Pex2 or the fission protein Pex11, we performed double-labeling experiments using cryosections of *pex1* cells producing either HA-tagged Pex2 or Pex11. These experiments revealed that in *pex1* cells producing Pex2-HA, both α-HA– and α-Pex14–dependent specific labeling was present at the same membrane structures (Fig. 3 G), indicating that the cells did not contain two different types of vesicles. Similarly, Pex14 colocalized with Pex11-HA (Fig. 3 H) at the same membrane structures in *pex1* cells. Specific α-Pex14 and α-HA label was invariably confined to the membrane structures and never observed at any other structure in the cell. Double-labeling experiments revealed that peroxisomal ghosts containing both Pex11 and Pex14 were also present in glucose-grown *pex1* cells (Fig. S2 A) but were difficult to detect because of their low abundance. These data suggest that the ghosts proliferate upon incubation of cells in oleic acid media, similar to peroxisomes in WT cells. Proliferation most likely occurs by fission because in cells of a *pex6* *pex11* double-deletion strain, invariably a single ghost structure was observed (Fig. S2 C).

The peroxisomal matrix protein thiolase (Pot1) did not localize to the membrane structures in oleic acid–induced *pex1* cells (Fig. 3 I), which suggests that the remnants are defective in matrix protein import. Therefore, we considered the remnants to represent peroxisomal ghosts that contain all PMPs and can proliferate but are unable to import matrix proteins.

Peroxisomal ghosts in *pex1* cells are empty peroxisomal membranes that form a rounded membrane structure

Further morphological analysis revealed that the ghosts were rounded in shape, which in ultrathin cross sections appeared as double-membrane rings (Fig. 2 B and Fig. 3, B and C). These ghosts measured up to 100 nm in diameter. In longitudinal sections, a single hole was observed in the structure (Fig. 2 B, III, black arrowhead). Our data support a model in which *pex1* cells contain empty peroxisomes, which flatten and curve into rounded structures. The structures do not represent autophagosomes, as they were also observed in cells of a *pex1 atg1* double-deletion strain (Fig. S2 B). The ghosts strongly resemble those previously described in *S. cerevisiae* (Hettema et al., 2000) and *Hansenula polymorpha* *pex1* and *pex6* cells (Koek et al., 2007), as well as in *Pex6*-deficient CHO cells (Hashiguchi et al., 2002).

Pex1 reintroduction results in the import of matrix proteins into preexisting peroxisomal ghosts

To study peroxisome reintroduction in *pex1* cells, we created a conditional allele of Pex1 by tagging it with the yeast-optimized auxin-inducible degron (AID*) 6HA tag (AID*-6HA; Morawaska and Ulrich, 2013). Control experiments confirmed that upon growth of Pex1-AID* cells in media lacking auxin, Pex1-AID*-6HA protein was present (Fig. 4 A) in conjunction with normal import of DsRed-SKL into peroxisomes (Fig. 4 B). However, when cells were grown in the presence of auxin, Pex1-AID*-6HA protein was not detectable and DsRed-SKL mislocalized to the cytosol (Fig. 4, A and C).

For reintroduction experiments, Pex1-AID* cells were first precultivated in glucose media in the presence of auxin and subsequently incubated for another 4 h in oleic acid media supplemented with auxin to induce peroxisomal proteins. Next, the cells were washed twice and further cultivated in oleic acid media lacking auxin. FM revealed that after 4 h of further cultivation in the absence of auxin DsRed-SKL, spots had reappeared, indicative of peroxisome formation (Fig. 4 C).

We then analyzed the cells at different stages of Pex1 reintroduction by EM. Before the removal of auxin, the cells contained the typical membrane ghosts to which Pex14 was localized (Fig. 4 D), similar to those observed in *pex1* cells (compare with Fig. 3 B). Also, thiolase did not accumulate
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in these structures (Fig. 4 D). ET indicated that the ghosts in Pex1-depleted cells represented rounded structures with tightly opposed membranes. As in pex1 cells, these ghosts contained a single opening connecting the lumen of the structure with the cytosol (Fig. 4 D). Generally, these structures were localized in close vicinity of the plasma membrane and cortical ER (Fig. 5 A). ET analysis indicated that the membranes of the ghosts were not continuous with the ER (Fig. 4, D and E), which was underscored by FM analysis, which showed that Pex14-mGFP spots exist that do not colocalize with the ER marker Sec63-mRFP (Fig. 5 B).

Already 2 h after shifting the Pex1-AID* cells to oleic acid media lacking auxin, most ghosts had undergone a morphological change and different intermediate stages toward normal peroxisomes could be distinguished (Fig. 4 E). First, the space in-between the membranes expanded, particularly near the
opening, because of the import of matrix protein, as was evident from immuno-labeling experiments using antibodies against thiolase (α-Pot1; Fig. 4 E, early). Next, structures were observed containing globular extensions that are likely the result of further import of matrix proteins (Fig. 4 E, medium). Still, these large extensions were connected to the initial membranes by a sheet of tightly opposed membranes and reached similar diameters as the peroxisomes that were ultimately generated (Fig. 4 E, medium). This indicates that the peroxisomes (Fig. 4 E, late) most likely are formed by fission of the ghost structure.

In summary, our results demonstrate that *S. cerevisiae* pex1 and pex6 cells contain peroxisomal membrane ghosts.
cells contain peroxisomal ghosts

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that harbor the docking proteins Pex13 and Pex14 as well as the RING proteins Pex2 and Pex10 together with Pex11 (van der Zand et al., 2012) using our fluorescence and immunocytochemical methods.

The structures most likely proliferate from existing ghosts by fission. These ghosts are defective in matrix protein import...
because of depletion of Pex1, but not in PMP insertion. Also, organelle fission is most likely not affected based on the finding that their proliferation depends on PEX11.

Upon reintroduction of Pex1, the ghosts imported matrix proteins and developed directly into peroxisomes essentially as described before for Pex6-deficient CHO cells upon genetic complementation (Hashiguchi et al., 2002; for a hypothetical model, see Fig. S3). Our data therefore support a model in which Pex1 and Pex6 play a role in matrix protein import (Platta et al., 2005) but cannot exclude an additional role for these proteins in vesicular fusion during peroxisome biogenesis (Tiforeno and Rachubinski, 2000; van der Zand et al., 2012).

**Materials and methods**

**Organisms and growth**

The yeast strains used in this study are listed in Table S1. Cells were grown in selective medium (0.17% yeast nitrogen base without amino acids and ammonium sulfate, 0.5% ammonium sulfate and 1% casamino acids) containing 2% glucose (YM2) or 0.1% oleic acid and ammonium sulfate, 0.5% ammonium sulfate and 1% casein hydrolysate (YM7). Amino acids and uracil were added when needed.

**DNA manipulations, cloning procedures, and strain constructions**

*S. cerevisiae* BY4742 WT, *pex1*, and *pex6* strains and the FY1679 WT strain were obtained from the Euroscarf collection (Table S1). All plasmids used in this study are described in Table S2. Gene fusions were made by PCR-based methods using the primers listed in Table S3. Correct introduction of the fusion genes was checked by colony PCR, Western blotting, and/or Southern blotting. All gene deletions were confirmed by PCR or Southern blotting.

For construction of the FY1679 *pex1* and *pex6* strains, the *PEX1* and *PEX6* deletion cassettes were PCR amplified from BY4742 *pex1* and *pex6* strains by using the primer pairs pKEK123/pKEK124 or pKEK127/pKEK128, respectively. The resulting PCR fragments of 2.6 and 2.3 kbp were transformed into FY1679 WT cells that were plated on YPD with selective antibiotics (GE Healthcare). For construction of the *pex1* strain, the *ATG1* deletion cassette was PCR amplified from strain WT *PEX1* using the primers pTER208/pTER209, resulting in a PCR fragment of 2.7 kbp, which was transformed into the BY4742 *pex1* strain, which was plated on YPD containing nourseothricin selective antibiotics (GE Healthcare). For construction of the *pex1 Δpex6* double-deletion strain, the *PEX6* deletion cassette was PCR amplified from pHyg-AID*-6HA* using the primer pair pKEK229/pKEK230, resulting in a PCR fragment of 1.7 kbp, which was transformed into a *pex1* strain already containing Pex2-mCherry and Pex14-mGFP. In the same way, *Pex6* was deleted in the Euroscarf *pex1* strain to construct the *pex1 Δpex6* double-deletion strain.

For construction of Pex14-mGFP strains, the PEX14-mGFP gene with downstream HIS marker was PCR amplified from strain WT *PEX14* using the primers pKEK036/pKEK041. The resulting PCR fragment of 2.3 kbp was integrated in yeast cells using standard transfection protocols. For construction of Pex13-mGFP strains, the PEX13-mGFP gene with downstream HIS marker was PCR amplified from strain WT *PEX13* using the primers pKEK164/pKEK165, resulting in a PCR fragment of 2.4 kbp.

For the construction of plasmid pRSA01, a PCR fragment of 700 bp was obtained by primers RSA10fw and RSA11rev on pCODNA3.1.mCherry. The resulting BglIII–SalI fragment was inserted between the BglIII and SalI of pANL31. For construction of Pex2-mCherry strains, the mCherry gene and Zeocin selection marker of the pRSA01 plasmid were amplified with primers pKEK182/pKEK183, thus yielding a PCR fragment of 2.3 kbp with 5′ overlap on the *PEX2* gene and 3′ overlap on the *PEX2* terminator. For construction of Pex10-mCherry strains, the mCherry gene and Zeocin selection marker of the pRSA01 plasmid were amplified with primers pKEK038/pKEK039, thus yielding a PCR fragment of 2.3 kbp with 5′ overlap on the *PEX10* gene and 3′ overlap on the *PEX10* terminator. For construction of Pex11-mCherry strains, the mCherry gene and Zeocin selection marker of the pRSA01 plasmid were amplified with primers pKEK184/pKEK185, thus yielding a PCR fragment of 2.3 kbp with 5′ overlap on the *PEX11* gene and 3′ overlap on the *PEX11* terminator.

For C-terminal tagging of *PEX11* with the HA tag, first the Zeocin marker was PCR amplified with 5′ overlap of the HA sequence (GYP YDVPDYASG) using the primers pKEK131/pKEK136, yielding a PCR fragment of 1.5 kbp. This PCR product was then amplified using 5′ overlap of the Pex11 C terminus and *PEX11* terminator using the primers pKEK137/pKEK138, then yielding a fragment of 1.6 kbp, which was then used to transform *pex1* cells. For C-terminal tagging of *Pex2* with the HA tag, the pHyg-Pex2-6HA plasmid was constructed by SmaI and EcoRI digestion of the pHyg-AID*-6HA* plasmid resulting in two fragments of 4,355 and 168 bp. The C-terminal part of Pex2 was amplified using primers pKEK202/pKEK203, yielding a product of 720 bp, which was then plated on LB containing ampicillin as selection marker. The final pHyg-Pex2-6HA plasmid was linearized with ApaI and transformed into yeast.
To obtain constitutive expression of a gene encoding DsRed containing a peroxisomal targeting signal 1, first the TDH3 promoter was amplified with primers TDH3_NotI/F/TDH3_BamHI.R, thus yielding a PCR fragment of 716 bp with 5′ NotI and 3′ BamHI digestion sites. The vector pHIPX7 GFP-SKL was digested with NotI and BamHI and assembled in pTEF promoter and flanked with the digested TDH3 promoter, thus resulting in the pPdh3 GFP-SKL plasmid. The plasmid pHIPX2 DsRed–SKL was digested with BamHI and SalI and the obtained DsRed–SKL fragment was subsequently ligated into the pPdh3 GFP-SKL plasmid, which was digested with BamHI and SalI, finally resulting in the pPdh3 DsRed-SKL plasmid. The plasmid was linearized with DraI and integrated into a TIR1 strain already containing Pex14-mGFP. The plasmid for integration of a C-terminal AID*–tag for Pex1 was created by Pex1 amplification with primers pKEK178/pKEK179, resulting in a product of 1697 bp, which was digested with HindIII and SalI and ligated in pHyg-AID*–6HA plasmid that was digested with the same enzymes. The resulting plasmid, pHyg-Pex1-AID*–6HA, was linearized with BstBI for integration in the TIR1 strain containing Pex14-mGFP, as well as DsRed-SKL. The TIR1 strain and AID*–plasmid were provided by H. Ulrich, Institute of Molecular Biology, Mainz, Germany.

In general, S. cerevisiae cells were transformed using PCR-amplified DNA. 2 μg PCR-purified DNA and 0.1 μg carrier DNA were added to 60 μl cells. Carrier DNA was denatured at 100°C for 10 min. A solution of PEG/LiAc/DTT was added with a volume of 300 μl every 100 μl of cell-DNA mixture. Heat shock was performed for 15 min at 42°C, followed by 2 min on ice. Then, the cells were centrifuged for 1 min at 5,000 rpm, the supernatant discarded, the pellet suspended in 5 ml YPD, and incubated for 3 h at 30°C with 200 rpm shaking. Again, the cells were centrifuged for 1 min at 5,000 rpm and the pellet suspended in 100 μl YPD or YND and plated on selective YPD/YND plates.

**FM**

Cells were fixed in 1% formaldehyde in PHEM buffer, consisting of 60 mM Pipes, 25 mM Hapes, 8 mM MgCl₂, and 40 mM EGTA, pH 6.9, for 1 h and embedded in 1% low-melting-point agarose in PHEM buffer to prevent cell movement during imaging.

To assess PMP colocalization, single plain images were acquired through the middle of the cells for bright-field, mGFP and mCherry on a Personal Deltavision (GE Healthcare) using a Photometrics CoolSNAP HQ2 digital camera and SoftWorks 5.5.1 software. All images were made at room temperature using 100× 1.30 NA Plan-Neofluar objectives (Carl Zeiss), resulting in a pixel size of 64 nm. The mGFP signal was visualized with a 470/40 nm band-pass excitation filter, a 495-nm dichromatic mirror, and a 525/50 nm band-pass emission filter. mCherry fluorescence was visualized with a 572/35 nm band-pass excitation filter and a 632/60 nm band-pass emission filter. Because the mCherry spots displayed lower fluorescence intensities relative to the mGFP spots, all mCherry spots present in 25 randomly acquired fluorescence microscopy images were selected and their distance to the closest mGFP spot was measured (Fig. S1 B). To extract the fluorescent spots from the original 16-bit TIF images, the coordinates of mCherry spots were determined semiautomatically using IMOD (Kremer et al., 1996), after which they were automatically boxed out for both channels resulting in images with a frame size of 32 × 32 pixels. The signal was enhanced and normalized with a fast-Fourier transform band-pass filter (between 16 and 4 pixels) in ImageJ (National Institutes of Health). Per channel, all processed images were corrected equally for background signal and converted to 8-bit TIF images. For the object-based colocalization analysis, spots were masked and labeled using the Dipimage toolbox in MATLAB (MathWorks). The center of gravity was determined in both mGFP and mCherry channels and used to calculate the distance between the mostly centered spots.

Sec63-mRFP and DsRed localization images were captured with a confocal microscope (LSM510; Carl Zeiss), equipped with photomultiplier tubes (Hamamatsu Photonics) and Zen 2009 software. All images were made at room temperature using 100× 1.30 NA Plan-Neofluor objectives (Carl Zeiss). For Sec63-mRFP, the cells were fixed in 1% formaldehyde in PHEM buffer and embedded in 1% LMP agarose in PHEM buffer. mGFP fluorescence was analyzed by excitation of the cell with a 488-nm argon ion laser (Lasos), and emission was detected using a 500–550-nm band-pass emission filter. The mRFP and DsRed signals were visualized by excitation with a 543-nm helium neon laser (Lasos), and emission was detected using a 655- to 615-nm band-pass emission filter. To reduce possible bleed through of mGFP into the mRFP/DsRed channel, the fluorescence images were acquired sequentially. The resulting 3D confocal stacks were median filtered using a 3D 2 × 2 × 2 kernel and merged in Z-direction by averaging.

**EM**

For morphological analysis, S. cerevisiae cells were fixed in 1.5% potassium permanganate, poststained with 0.5% uranyl acetate, and embedded in epon 812 (Serva, 21045). Ultrathin sections were viewed in a Philips CM12 TEM. For ET, 10-nm gold beads were layered on top of 400-nm-thick sections and acted as fiducial markers for ET. Two single-axis tilt series, each containing 141 images with 1° tilt increments, were acquired with a pixel size of 1.16 nm on a FEI Tecnai20 at 200 kV using the FEI automated tomography software and a cooled slow-scan charge-coupled device camera (4k Eagle; FEI Company) in 2 × 2 binned mode. The tilt series were aligned and reconstructed using the IMOD software package (Kremer et al., 1996) and analyzed using the AMIRA visualization package (TGS Europe). To generate 3D surface-rendered models in AMIRA, masks of organelles were first drawn manually and afterward improved by thresholding.

For immunogold labeling, cells were washed twice in PHEM buffer, then fixed overnight in a mixture of 0.2% glutaraldehyde and 2% formaldehyde in PHEM buffer at 4°C and subsequently incubated for 15 min in a solution of 0.4% sodium periodate and for 30 min in 1% ammonium chloride. Upon embedding in 12% gelatin in PHEM buffer, ~0.5-mm³ cubes were infiltrated overnight in 2.3 M sucrose in the same buffer. Cryosections of 70 nm were cut using a cryo diamond knife (Diatome) at −120°C using a Reichert Ultracut and mounted on carbon-coated formvar nickel grids. Immunolabeling of Pex14 and thiolase were performed using rabbit polyclonal antibodies followed by goat anti–rabbit antibodies conjugated to 6 or 10 nm gold (Auron). Pex11-HA and Pex2-6HA were localized using mouse monoclonal antibodies raised against HA (H9658; Sigma-Aldrich,) and goat-anti-mouse antibodies conjugated to 6 nm gold (Auron). Labeled sections were first stained for 2 min with 2% uranyl oxalate, pH 7, at room temperature and after a quick rinse on three drops of double distilled water stained and embedded in a mixture of 0.5% uranyl acetate and 0.5% methylcellulose (25 centipoise; Sigma-Aldrich) for 10 min on ice. Excess staining solution was drained, and the grids were left to dry before viewing them in a CM12 TEM (Philips).

For CLEM, the sample was prepared similarly as for immunogold labeling; however, 180-nm thin sections were cut and mounted on carbon-coated formvar copper grids that were over layered with 10-nm gold particles (Sigma 752584) for alignment of the tomograms. The grids were placed with section side facing the objective in a droplet of water on a coverslip. Fluorescence imaging was performed at room temperature using Axiovision 4.8.2 software on an Observer Z1 (Carl Zeiss) equipped with a 100× 1.30 NA Plan-Neofluor objective (Carl Zeiss) and an AxioCAM MRm camera (Carl Zeiss). mCherry fluorescence in both mGFP and mCherry channels was used to calculate the distance between the mostly centered spots.
was visualized with a 587/25-nm band pass excitation filter and a 647/70-nm band-pass emission filter. After fluorescence imaging, the grid was poststained and embedded in a mixture of 0.5% uranyl acetate and 0.5% methylcellulose. The area of interest was found back in the electron microscope using the bright-field images as maps. Low-magnification EM was used to align the EM images on the bright-field FM images. Acquisition of the double-tilt tomography series was performed manually in a CM12 TEM running at 80 kV and included a tilt range of 40° to −40° with 5° increments. Reconstruction of the tomograms was performed using the IMOD software package (Kremer et al., 1996).

Pex1 reintroduction

Cells producing Pex1 containing a C-terminal AID*-6HA-tag were precultivated for 16 h on YM2 medium containing 1 mM indole-3-acetic acid and subsequently inoculated to OD600 = 0.1 in YMO medium containing 1 mM indole-3-acetic acid salt (Sigma-Aldrich) and subsequently incubated at 40° with 5° increments. Reconstruction of the tomograms was performed using the IMOD software package (Kremer et al., 1996).

Online supplemental material

Fig. S1 shows FM analysis of colocalization of different PMPs in WT and mutant strains. Fig. S2 shows electron micrographs illustrating the presence of peroxisomal ghosts in glucose-grown pex1 cells as well as in oleic acid grown pex1 arg1 and pex6 pex11 double-deletion strains. Fig. S3 shows a hypothetical model of peroxisome reintroduction in yeast pex1 and pex6 cells. Tables S1, S2, and S3 contain the contributions and discussions and Leonard Bosgraaf for designing and preparing Fig. S3. We thank Egbert Boekema and Jan-Willem Veen for making their microscopy facilities available.

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