Multiple Pathways for Protein Transport to Peroxisomes

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Abstract

Peroxisomes are unique among the organelles of the endomembrane system. Unlike other organelles that derive most if not all of their proteins from the ER (endoplasmic reticulum), peroxisomes contain dedicated machineries for import of matrix proteins and insertion of membrane proteins. However, peroxisomes are also able to import a subset of their membrane proteins from the ER. One aspect of peroxisome biology that has remained ill defined is the role the various import pathways play in peroxisome maintenance. In this review, we discuss the available data on matrix and membrane protein import into peroxisomes.

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Introduction

Peroxisomes are organelles found in almost all eukaryotic cells. They are bounded by a single membrane and are usually spherical. Peroxisomes are involved in a variety of metabolic pathways that vary between species, cell types and environmental conditions. All peroxisomes contain enzymes that catalyze oxidation reactions, including a fatty acid oxidation system, as well as enzymes that protect cells from oxidative damage, such as catalase. It is not surprising that peroxisomes are essential for human metabolism and development, and this is illustrated by severe inherited diseases caused by impairments in one or more peroxisomal functions [1,2].

Peroxisomes are unique among single-membrane organelles in that peroxisomes are capable of importing both their matrix and membrane proteins directly from the cytosol. Peroxisomal matrix proteins are nuclear encoded and are synthesized on free polyribosomes [3]. As is the case for other targeting and translocation systems, peroxisomal matrix proteins contain specific targeting signals that are recognized by receptors that deliver them to the membrane translocation site. Translocation is dependent on consumption of energy in the form of ATP. Several aspects of peroxisomal protein transport distinguish it from other translocation systems. For example, peroxisomal proteins can fold, acquire cofactors and assemble into oligomers in the cytosol prior to import; another difference relates to the targeting receptors, which bind cargo in the cytosol but imbed into or cross the membrane as part of their targeting cycle.

Peroxisomal membrane proteins (PMPs) are also nuclear encoded. However, the mechanism of their import into peroxisomes is less understood. What is known is that PMPs and matrix proteins use different machineries for import. Most peroxisome biogenesis mutants contain membrane structures that are devoid of content, reflecting the large number of proteins required for import of matrix proteins (Table 1). The proteins required for peroxisome biogenesis are called peroxins (PEX) [4]. Only two PEX mutants in yeast (and three in mammals) are devoid of peroxisomal membrane structures.

Below, we review the targeting and import of matrix proteins separately from that of membrane proteins. We will discuss some of the discrepancies in the literature and the outstanding questions in the field.
detailed discussions on peroxisome multiplication, we refer to recent reviews [5–9].

Matrix Protein Import

Import of peroxisomal enzymes can be divided in multiple steps: (1) recognition in the cytosol by receptors, (2) docking of receptor–cargo complex at the peroxisomal membrane, (3) cargo translocation and (4) recycling of receptors (see Fig. 1).

Table 1. Peroxins required for peroxisomal matrix protein import in S. cerevisiae and their human orthologues

| S. cerevisiae | Homo sapiens | Process | Complex |
|--------------|--------------|---------|---------|
| Pex1         | Pex1         | Export of receptors | AAA+ complex |
| Pex2         | Pex2         | Ubiquitination of receptors | RING ubiquitin ligase complex |
| Pex4         | UbcH5a/UbcH5b/UbcH5c | Receptor ubiquitination | Ubiquitin conjugation complex |
| Pex5         | Pex5S/Pex5L  | Targeting, translocation | PTS1 receptor, import pore |
| Pex6         | Pex6         | Export of receptors | AAA+ complex |
| Pex7         | Pex7         | Targeting | PTS2 receptor |
| Pex8         |             | Translocation/cargo release | Intraperoxisomal peroxin |
| Pex10        | Pex10        | Receptor ubiquitination | RING ubiquitin ligase complex |
| Pex12        | Pex12        | Receptor ubiquitination | RING ubiquitin ligase complex |
| Pex13        | Pex13        | Docking | Docking complex |
| Pex14        | Pex14        | Docking, translocation | Docking complex, import pore |
| Pex15        | Pex26        | Export of receptors | AAA+ complex |
| Pex17        |             | Docking | Docking complex |
| Pex18/Pex21  | Pex5L        | Targeting | PTS2 coreceptor |
| Pex22        |             | Receptor ubiquitination | Ubiquitin conjugation complex |

As mentioned above, matrix protein import into peroxisomes differs from import into other organelles in that peroxisomes can import folded proteins and even protein complexes.

Fig. 1. Model for import of peroxisomal matrix proteins containing a PTS1 in S. cerevisiae. (a) A newly synthesized PTS1-containing peroxisomal matrix protein is recognized in the cytosol by the receptor Pex5. (b) Cargo-loaded receptor docks onto the docking complex consisting of Pex13, Pex14 and Pex17. (c) Receptor inserts in the membrane in complex with Pex14 and cargo is released into the lumen. (d) The ubiquitin ligase complex (Pex2, Pex10 and Pex12) in conjunction with the ubiquitin conjugation complex (Pex4 and Pex22) modifies Pex5 on a cysteine residue. (e) Ubiquitinated Pex5 is extracted from the membrane by the extraction complex (Pex1, Pex6 and Pex15) and (f) subsequently deubiquitinated for another round of targeting.
import is based on various experiments: (1) radioactive pulse-chase analyses show that newly synthesized β-oxidation enzymes are released in the cytosol before associating with peroxisomes [3], (2) in vitro synthesized peroxisomal matrix proteins are imported into purified peroxisomes (e.g., see Refs. [10–12]) and (3) import of purified peroxisomal proteins occurs in semi-permeabilized fibroblasts and after microinjection into the cytoplasm [13–15]. Indeed, the presence of a carboxy-terminal targeting signal in most peroxisomal matrix proteins is compatible only with a posttranslational targeting mechanism.

Whereas proteins targeted posttranslationally to mitochondria, chloroplasts and ER (endoplasmic reticulum) are kept in a partially unfolded conformation by chaperones and targeting factors, peroxisomal matrix proteins can fold and assemble into their active, cofactor-containing conformation in the cytosol before targeting (for a review, see Ref. [16]). For example, catalase is a haem-containing tetramer protein. In fibroblasts from patients with a peroxisome biogenesis defect, catalase is mislocalized to the cytosol. Fusion of peroxisome-deficient cells belonging to different complementation groups (i.e., having defects in different genes) results in import of the existing cytosolic pool of catalase [17]. Several studies have shown import of proteins that were folded prior to import [14,15,18]. Subunits of a homodimer or homooligomers artificially lacking a peroxisomal targeting signal (PTS) are co-imported with subunits containing a PTS [19–23]. Large structures (including gold particles or chemically cross-linked albumin) are imported into peroxisomes when coated with peptides containing a PTS after microinjection into the cytosol of fibroblasts [24].

However, folding is not a prerequisite for import [25], and not all peroxisomal proteins are multimers. Although tetrameric catalase can be imported in vivo [17,23], an in vitro study reported that catalase monomers preferentially bind the import receptor that inhibits oligomerization. This would suggest that catalase is imported mainly as a monomer if the concentration of the receptor is high enough to bind newly synthesized catalase [26]. Likewise, pumpkin isocitrate lyase monomers are imported preferentially over tetramers in vitro [27]. Other oligomeric proteins that assemble into their quaternary structures inside peroxisomes include Hansenula polymorpha alcohol oxidase (an octamer of 600 kDa) [28,29] and porcine heart peroxisomal carbonyl reductase (a tetramer). The latter protein cannot be imported as a tetramer as the PTSs are no longer accessible for recognition by the PTS receptor [30].

Therefore, although peroxisomes are capable of importing fully assembled oligomeric proteins, this may not reflect the way these proteins are imported under physiological conditions.

**Recognition by Targeting Receptors**

The carboxy-terminal PTS1 was identified first in firefly luciferase and has since been found in most matrix proteins. The C-terminal tripeptide SKL was found to be both necessary and sufficient to target a protein to peroxisomes [31]. Variations of this signal have been described, resulting in the consensus -(SAC)-(KRH)-(LM) that is evolutionarily conserved [32–35]. Some PTS1s do not conform to the consensus, and amino acid residues immediately preceding the tripeptide appear to contribute to targeting in those cases [36–38]. The PTS1 is recognized by the receptor Pex5, which is a modular protein containing a C-terminal tetratricopeptide repeat domain: it is the tetratricopeptide repeat domain that interacts with the PTS1 [39,40].

A second evolutionarily conserved PTS is found close to the amino-terminus of a subset of peroxisomal matrix proteins. This PTS2 was first described as a nonapeptide present in the amino-terminal cleavable presequence of rat peroxisomal thiolase [41–43]. Most PTS2s fit the following consensus -R-(LIVQ)-X-X-(LIVQH)-(LSGA)-X-(HQ)-(LA)- [43]. The PTS2 is recognized by the receptor Pex7. The PTS2-dependent targeting route is absent in Caenorhabditis elegans, Drosophila melanogaster and diatoms [44–46].

Whereas Pex5 is sufficient for targeting PTS1-containing proteins to the peroxisomal membrane, many fungi require the coreceptor Pex20 for PTS2 targeting via Pex7 [47–49] or the Pex20-like paralogues Pex18 and Pex21 in Saccharomyces cerevisiae [50]. These coreceptors are absent from mammals and plants, where PTS2 import requires both Pex7 and Pex5. Human cells express at least two isoforms of Pex5: Pex5S and Pex5L. The latter has an additional exon that is required for binding of Pex7 to Pex5. Human cells express at least two isoforms of Pex5: Pex5S and Pex5L. The latter has an additional exon that is required for binding of Pex7 to Pex5. This binding site is conserved in the Pex20-like proteins and plant Pex5 and is required for PTS2 import [47,51–53]. Besides the Pex7 binding sites, the carboxy-terminal domain of the Pex20-like proteins shows additional sequence similarities to the N-terminus of Pex5. Therefore, Pex5L is considered a Pex7 coreceptor.

Not all peroxisomal matrix proteins are targeted to peroxisomes under all conditions. For instance, the activity of the PTS2 of S. cerevisiae GPD1 depends on phosphorylation [54].

**Non-Classical Import Mechanisms**

Interestingly, not all matrix proteins contain a PTS1 or a PTS2 [20]. CuZn superoxide dismutase 1 is found mainly in mitochondria and cytosol but is also localized to peroxisomes. This non-PTS1/PTS2-containing protein is piggy back imported into peroxisomes through its direct interaction with the PTS1-containing copper chaperone for superoxide dismutase [22].
Others, however, are not imported via the classical PTS1, PTS2 or even by a piggy-back pathway. Instead, proteins such as acyl-CoA oxidase (Pox1) in *S. cerevisiae* are recognized via the N-terminal domain of Pex5 rather than via its PTS1 binding domain [55] (for review, see Ref. [56]).

### The Recycling Receptor Concept

Early analysis of human Pex5 revealed it to be localized mainly to the cytosol, with a small fraction associated with peroxisomes. Localization to peroxisomal membranes is increased in the skin fibroblasts from peroxisome biogenesis disorder patients with a mutation in Pex2, Pex10 or Pex12 and in wild-type cells upon ATP depletion and incubation at low temperature. On restoring translocation conditions, Pex5 returned to the cytosol, after which it could be trapped again on peroxisomes by depleting ATP and incubating at low temperature. These experiments show clearly that Pex5 is a cycling receptor and that this cycle is modulated by factors required for peroxisomal protein import [57].

Different models have been proposed for Pex5 shuttling between cytosol and peroxisomal membrane, including a model whereby it dips into the membrane and releases its cargo at the luminal side of the membrane (shuttle model) [58] and another model whereby Pex5 enters the peroxisomal lumen before being exported back to the cytosol (extended shuttle model) [59]. Pex7 also behaves similar to a cycling receptor and was proposed to follow the extended shuttle model [60]. It is clear from these studies that Pex5 and Pex7 are exposed to the lumen of the peroxisome, but whether they are released into the lumen is not clear.

### Docking of Cargo–Receptor Complexes

The docking complex consists of Pex13 and Pex14 (and Pex17 in yeasts) and is required for both PTS1 and PTS2 import [61]. The docking complex proteins interact with the PTS receptors and each other (for detailed reviews, see Refs. [62–64]). The exact roles of Pex13 and Pex14 in docking are unclear. Both proteins have been shown to interact with cargo-loaded PTS1 and PTS2 receptors [65–67]. In mammals, cargo-loaded PTS1 receptor associates first with Pex14 before being translocated to a high-molecular-weight complex containing Pex13 [68]. The cargo-loaded PTS2 receptor complex also docks initially onto Pex14 via its coreceptor Pex5L [65,69]. Subsequently, the Pex7-cargo-loaded complex binds to Pex13 independently of Pex5L [65,67].

In *S. cerevisiae*, the PTS2 receptor complex appears to dock first onto Pex13 [66].

### Cargo Translocation and Release

After docking of the cargo-loaded receptors, the cargo must be translocated across the membrane and released inside the peroxisome. These processes are poorly understood. *In vitro* import studies using Pex5 are starting to give new insights into the translocation mechanism, which needs to allow for translocation of oligomeric proteins without disturbance of the internal milieu of the peroxisome. Several observations are compatible with a model whereby translocation occurs through a dynamic transient pore that disassembles after translocation. This pore is proposed to be composed of the targeting receptors plus components of the docking complex [70]. Indeed, Pex5 behaves as a soluble protein in the cytosol, as well as a carbonate-extraction-resistant and partially protease protected protein when in its cargo-loaded form at the peroxisomal membrane: as a receptor–cargo complex, Pex5 becomes an integral part of the translocation pore and behaves as an integral membrane protein. The C-terminal domain containing the PTS1 binding site is then exposed to the lumen of the peroxisome [71–73]. Pex5 has been reported to form homooligomers and to insert spontaneously into membranes [74,75].

Pex5 together with Pex14 are the only peroxins required for import of Pex8 into peroxisomes, and they therefore constitute the minimal translocon [76]. *In vitro*, the *S. cerevisiae* Pex5–Pex14 complex can form an ion-conducting channel with a variable pore size of up to 9 nm [77], with the size of the pore depending on the size of the cargo. *Leishmania* Pex14 has also been shown to have pore-forming activity [78]. The actual architecture of the pore is unknown, and many questions remain, such as the driving force for translocation, whether PTS2 proteins are transported via the same or a separate pore and whether the Pex7 coreceptors will form part of the pore.

The mechanism of release of cargo from receptors has remained elusive. Recent *in vitro* studies suggest that Pex14 in mammals and Pex8 in *Pichia pastoris* may be involved [26,79]. In plants, Pex14 has been suggested to be required for release of PTS2 cargo [80]. Furthermore, binding/release of cargo to Pex5 has been suggested to be redox dependent [79]. The development of an *in vitro* import system that follows the import of a specific cargo rather than that of the targeting receptors [81] is an important step that will likely resolve some of the mysteries.

### Export of Receptors

After release of their cargo, the receptors are exported back to the cytosol in an ATP-dependent
ubiquitin ligases Pex2, Pex10 and Pex12. Monoubiquitination occurs after insertion of Pex5 into the membrane and cargo release and is essential for recycling of Pex5 and therefore for PTS1 import. Monoubiquitination is catalyzed by the dedicated ubiquitin-conjugating enzyme Pex4 that is localized to the cytosolic side of the peroxisomal membrane. In mammals, Pex4 is absent, and Pex5 monoubiquitination depends on a family of related ubiquitin-conjugating proteins (Ubc5a–Ubc5c). Pex5 monoubiquitination also depends on an integral membrane protein complex containing the RING ubiquitin ligases Pex2, Pex10 and Pex12.

In vitro export assays revealed that monoubiquitinated Pex5 is exported by the AAA-type ATPases Pex1 and Pex6. The hexameric AAA+ complex consists of three subunits each of Pex1 and Pex6. The complex associates at the cytosolic side of the peroxisomal membrane via the tail-anchored membrane protein Pex19. Pex19 acts as a soluble recycling receptor/chaperone that picks up newly synthesized PMPs in the cytosol and subsequently docks on Pex3 in the peroxisomal membrane. The current model for direct posttranslational targeting of PMPs to peroxisomes proposes that Pex19 binds to newly synthesized PMPs, and those that are not targeted by Pex19 are called class 2 or mPTS2.

Membrane Protein Targeting

The machinery required for PMP import is distinct from that required for matrix protein import. In almost all patients with peroxisome biogenesis disorders, matrix proteins are mislocalized to the cytosol, while PMPs are present in membrane structures called peroxisomal membrane ghosts. The exceptions to this are two mutants in yeasts and three mutants in mammals (pex3, pex16 and pex19) that lack peroxisomal membrane ghosts. For this reason, Pex3, Pex16 and Pex19 are thought to be involved in the formation of peroxisomal membranes and/or insertion of PMPs into peroxisomal membranes.

Biochemical and microscopy studies of pex3, pex16 and pex19 mutants suggest that at least two routes exist by which PMPs can reach peroxisomes: one direct route and one via the ER. Below, we discuss the various routes and machineries involved in PMP insertion and provide examples to illustrate that a PMP may not be confined to a single route.

Direct Targeting Pathway

Most PMPs are synthesized on free polyribosomes and imported posttranslationally from the cytosol. Many of these PMPs possess a membrane PTS (mPTS) that consists of a cluster of positively charged residues or a mixture of positively charged and hydrophobic residues flanked by one or two transmembrane segments that are recognized and bound by Pex19. However, there is another group of PMPs that does not bind Pex19 (Jones et al., 2004). Thus, PMP membrane targeting signals have been classified by their ability to be recognized and targeted to peroxisomes by Pex19. Those that are not targeted by Pex19 are called class 1 or mPTS1, and those that are not targeted by Pex19 are called class 2 or mPTS2.

A model for direct targeting to peroxisomes

The current model for direct posttranslational targeting of PMPs to peroxisomes proposes that Pex19 acts as a soluble recycling receptor/chaperone that picks up newly synthesized PMPs in the cytosol and subsequently docks on Pex3 in the peroxisomal membrane. In line with this model, Pex19 binds to newly
synthesized PMPs to keep them soluble and stable in the cytosol. This is based on in vitro studies where Pex19 was found to bind newly synthesized PMPs and keeps them soluble [125,126,140–142]. In line with this, peroxisome-less cells (Pex3-deficient fibroblasts) overexpressing Pex19 stabilize PMPs in the cytosol and prevent them from being degraded [113,143]. Furthermore, Pex19 not only binds PMPs but also transports them to different subcellular locations. For instance, the overexpression of Pex19 tagged with a nuclear localization signal resulted in the accumulation of PMPs in the nucleus [143]. Pex19 cannot insert, on its own, PMPs into the membrane for this, the PMP membrane receptor protein Pex3 is also required. Together with the cargo, Pex19 binds to Pex3 on the peroxisomal membrane to form a trimeric complex of Pex3, Pex19 and cargo [125]. The importance of the Pex3–Pex19 interaction for PMP targeting is illustrated by mutations that prevent Pex3–Pex19 interaction that also prevents PMP insertion into peroxisomes [126,144,145].

Pex3 is anchored in the peroxisomal membrane by its N-terminal transmembrane domain, with the remainder of the protein forming an α-helical bundle that protrudes into the cytosol [144,146]. This domain contains a hydrophobic surface located near the base of the protein close to the lipid bilayer. It is thought to mediate binding to liposomes in vitro [147] and has been suggested to locally deform the peroxisomal membrane [142]. Mutagenesis studies of this hydrophobic patch revealed the importance of its hydrophobicity and its shape for PMP insertion [145] but not Pex19 binding [142]. At the apex of the bundle lies a hydrophobic groove that mediates binding of an N-terminal amphipathic helix of Pex19 [144,146]. The region on Pex19 that binds mPTSs is located in its C-terminal domain [138,139,148]. An additional amphipathic helix in Pex19 is thought to facilitate the handover of transmembrane domains to Pex3.

Fig. 2. Models for PMP import pathways. PMPs are either inserted directly from the cytosol into the peroxisomal membrane (a–c) or inserted into the ER and subsequently sorted to peroxisomes (d–f). (a) The main mechanism of direct import of PMPs into peroxisomes dependent on Pex3 and Pex19. Pex19 acts as the cytosolic chaperone while Pex3 is the membrane anchor. Some PMPs have multiple mPTS sequences that likely bind multiple Pex19 to allow for the import of multi-membrane spanning PMPs. This mechanism seems universal. (b) Direct insertion of Pex3 into the peroxisomal membrane dependent upon Pex16 and Pex19 in mammals. (c) Similar to other PMPs, TA PMPs are inserted directly via the Pex3/Pex19 complex in mammals. (d) Insertion of Pex3 and other PMPs into the ER via the ER translocon. This process has been shown in S. cerevisiae. It is not known whether a cytosolic chaperone is required for this process. (e) In mammalian cells, Pex3 insertion into the ER requires Pex16. (f) TA PMPs in S. cerevisiae have been shown to use the Get complex. However, the mammalian homologue TRC40 complex is not involved in the import of TA PMPs.
Subsequent transmembrane insertion occurs where the Pex3 hydrophobic surface deforms the peroxisomal membrane [142].

PMPs with multiple transmembrane segments have multiple Pex19 binding domains: these PMPs may bind multiple Pex19s simultaneously and dock onto the same Pex3 sequentially or to multiple Pex3s (Fig. 2a) to allow their insertion into the bilayer.

**Indirect Targeting Pathway (via the ER)**

The ER-to-peroxisome pathway was first described in *Yarrowia lipolytica* [149,150]. One of the key observations was that Pex2 and Pex16 are core-glycosylated PMPs. This result suggested that these PMPs travel via the ER to peroxisomes bypassing the Golgi complex. No other endogenous PMPs have been shown to be glycosylated in any other model system, which is unfortunate as this would provide a useful tool to study ER to peroxisome transport. Since then, several PMPs have been described to travel to peroxisomes via the ER in a number of different systems, including ascorbate peroxidase and Pex16 in plants and Pex3 and Pex16 in mammals, as well as Pex3, Pex15 and Pex22 in *S. cerevisiae* [117,151–155]. Of these PMPs, transport of Pex3 has been studied the most intensively.

Pex3, the protein that mediates direct insertion of PMPs into the peroxisomal membrane together with Pex19, is itself a PMP. This leads to an apparent conundrum of how Pex3 is itself inserted into the peroxisomal membrane. It has been known for some time that Pex3 targeting to peroxisomes differs from that of other PMPs as it contains an mPTS2 instead of an mPTS1 such as most PMPs [143]. This targeting quagmire was resolved by a number of groups who simultaneously described the transport of Pex3 via the ER to peroxisomes in *S. cerevisiae* – [156–158]. In Pex3-deficient cells, conditional expression of Pex3-GFP allowed for the visualization of de novo peroxisome formation from the ER. Pex3 was seen first in the ER and associated puncta, after which the puncta lost their ER association and matured into peroxisomes [156]. Also, when Pex3 is forced to insert cotranslationally into the ER by addition of the invertase signal peptide, not only it complemented pex3 cells but also, more importantly, the signal peptide was cleaved and the processed Pex3 was shown to end up in newly formed peroxisomes [157], thus showing that Pex3 traveled to peroxisomes via ER.

More recent studies suggest that Pex3 import into the ER in *S. cerevisiae* involves the ER posttranslational Sec61/Sec62/Sec63 translocon (Fig. 2d) [159]. The mPTS2 of Pex3 of *S. cerevisiae*, which was previously shown to be required for its peroxisomal localization [158], was dissected further into signals that direct Pex3 (1) into the ER, (2) sort it from there to a punctate ER subdomain (pER) and (3) sort it from the pER onto peroxisomes [159,160]. Interestingly, Pex3 appears not to be the only PMP to be imported into the ER by the posttranslational translocon. For instance, partial depletion of the ER translocon subunits Sec62 and Sec63 in *S. cerevisiae* showed a partial mislocalization of Pex13-GFP to the cytosol [161]. Additionally, Pex13 and Pex14 fail to associate with membranes in pex3 mutants depleted for Sec62/Sec63 as revealed by differential centrifugation [162]. Together, these observations make a strong case for a requirement of the Sec61 translocon for peroxisome biogenesis and membrane insertion of some PMPs in *S. cerevisiae*.

**Targeting of TA PMPs in *S. cerevisiae*: A Tale of Two Pathways**

TA proteins destined for the endomembrane system insert into the ER membrane via a variety of partially redundant machineries [163,164]. The ATP-dependent insertion via the Get pathway is the best studied of these mechanisms and appears to be a major contributor to insertion of TA proteins with a hydrophobic tail anchor region. The Get/TRC machinery consists of a cytosolic ATPase (Get3/TRC40) that binds the transmembrane segment of newly synthesized TA proteins and delivers them to the ER membrane [165,166]. Get1 and Get2 are integral ER membrane proteins that function as Get3–cargo complex receptors in yeast [166].

Overexpression of the peroxisomal TA protein Pex15 results in its accumulation in the ER [166,167]. In Get-deficient cells, a pulse of a GFP-Pex15 fusion protein fails to label peroxisomes and the ER but instead labels a cytosolic aggregate initially, which is followed by mitochondrial localization at later time points [166]. Pex15 appended with an opsin tag that is a 23-amino-acid residue is N-glycosylated and reaches peroxisomes [154]. Together, these observations indicate that Pex15 is initially targeted to the ER by the Get complex before being routed to peroxisomes. However, Pex15 can also reach peroxisomes independent of the Get machinery. In contrast to pex15 cells [167], get1/get2 and get3 cells have peroxisomes [162,166]. A short pulse of YFP-Pex15 expression showed low levels of YFP-Pex15 associated with peroxisomes in get3 cells [162]. As Get3 and Pex19 have been shown to interact with the tail anchor region of Pex15 [166,168], it is tempting to speculate that the direct pathway may also contribute to targeting *S. cerevisiae* Pex15 to peroxisomes.

In mammals and in *Neurospora crassa*, the Pex15 homologue, the TA PMP Pex26, is targeted and inserted independently of the Get/TRC machinery. Instead, it relies on the Pex19-dependent direct targeting pathway [126,142]. Similarly, the TA
protein Fis1 requires Pex19 and its Pex19 binding sites for targeting to peroxisomes [169]. These studies of TA proteins clearly show that this type of PMP can be targeted to peroxisomes using different machineries and pathways. In yeast, the predominant but not exclusive route appears to be the indirect pathway via the ER, whereas in mammalian cells and in N. crassa, the direct import into peroxisomes appears to dominate.

### Pex3 Targeting in Mammals

As observed in S. cerevisiae cells, human Pex3 fused to a signal peptide can complement Zellweger syndrome fibroblasts deficient in Pex3. This implies that, similar to yeast Pex3, human Pex3 can traffic to peroxisomes via the ER [153,155]. However, using an in vitro targeting assay in semi-permeabilized cells, Fujiki’s group has demonstrated that Pex3 can insert directly into peroxisomes by a pathway similar to that shown for other PMPs in that it requires the PMP cytosolic receptor Pex19. Since Pex3 lacks an mPTS1 but instead contains a targeting signal that does not bind Pex19 [143], the role of Pex19 in this transport is unclear, but may reflect its chaperone function.

Pex3 targeting directly to peroxisomes also requires Pex16, which is considered to function as a peroxisomal receptor for newly synthesized Pex3 (Fig. 2e) (Matsuzaki, 2008). Pex16 has a dual localization as it is also found in the ER membrane [117]. In live cells, Pex3 was shown to target to the ER when Pex16 was localized there even in cells deficient in functional Pex19 (Fig. 2b) [117,145]. However, in the absence of Pex16, Pex3, like many other PMPs, localized to mitochondria even in the presence of Pex19 [117,143,153]. Furthermore, in addition to Pex3, a number of other PMPs are recruited to the ER by Pex16 [117,155]. These data suggest a model whereby Pex3 and maybe some additional PMPs are targeted directly to peroxisomes via a Pex16-dependent, Pex19-dependent route, as well as to the ER by a Pex16-dependent, Pex19-independent route. The steady-state distribution of Pex16 and the availability of Pex19 binding sites on cargo molecules may determine which route is taken in mammalian cells.

How Pex16 itself is imported into a membrane is a point of contention. It has been reported that Pex16 is imported by the Pex3/Pex19 direct pathway [143], while another group suggests that Pex16 is cotranslationally inserted into the ER membrane before being routed to peroxisomes [117]. The different conclusions may result from differences in the Pex16 constructs used. The report showing direct targeting of Pex16 to peroxisomes is based largely on the ability of Pex19 to bind and redirect the last third of the Pex16 polypeptide to the nucleus and peroxisomes. However, the first half of Pex16 has been shown to be necessary and sufficient to target to peroxisomes in control cells [133]. In vitro targeting experiments suggest that Pex16 is inserted cotranslationally into the ER membrane [117], most likely by the ER translocon. Neither Pex3 nor Pex19 is required for its ER insertion. Furthermore, recent work that quantified PMP import kinetics showed that the import rate of Pex16 into peroxisomes was low compared to that of Pex3 and PMP34. It is proposed that direct import into peroxisomes is faster than insertion into the ER membrane and subsequent sorting to peroxisomes. Indeed, Pex16 transport kinetics are similar to those of Pex3 fused to a signal peptide, a construct that is first forced into the ER by an ER targeting signal sequence before being routed to peroxisomes [155]. It is also possible that mammalian Pex16 has two independent targeting sequences, one in the N-terminal part of the protein for cotranslational insertion into the ER membrane and one in the C-terminal part of the protein for direct insertion into the peroxisomal membrane. Since the ER targeting signal will emerge first from the ribosome, we propose that most Pex16 is cotranslationally inserted into the ER.

### ER-to-peroxisome transport

Since it is clear that some PMPs travel via the ER to peroxisomes, this raises the question as to how this transport is mediated. Two models explain how ER-inserted PMPs reach peroxisomes. In the first model, ER-inserted PMPs are sorted away from the secretory pathway and are delivered to peroxisomes, probably by vesicular transport. In the second model, peroxisomes form de novo from ER-derived vesicles thereby forming a new peroxisome. The two processes may be combined into a single process where a pre-peroxisomal vesicle either matures into a new peroxisome or fuses with an existing peroxisome. Which process dominates may vary among growth conditions, cell types or whether any peroxisomes are present.

In mammalian cells, the first model is supported by the finding that Pex16 is inserted into the ER membrane and sorted from there to peroxisomes [117,170]. This transport requires the secretory factor Sec16b [170]. Similarly, Pex3 “forced” into the ER by a signal sequence is transported to all pre-existing peroxisomes in mammalian cells [155]. This suggests that the ER is constantly providing lipids and proteins to pre-existing peroxisomes. The rate of transport of ER-targeted Pex3 was quite high, thus giving a possible explanation as to why PMPs are not readily found in the ER in wild-type cells. Further support for a model whereby existing peroxisomes continue to receive membrane material was provided by overexpression of a
Pex11β-YFP fusion protein that inhibits peroxisome fission. Newly synthesized matrix and membrane proteins are imported into subdomains of these elongated peroxisomal structures [171]. Additionally, using Halo-tag approaches, the Fransen laboratory showed that peroxisomes receive newly synthesized PMPs, including Pex16 (Huybrechts et al., 2009).

In cells that are devoid of peroxisomes due to loss of PMP import peroxins (Pex3 or Pex19), peroxisomes can form de novo on introduction of the corresponding gene. In S. cerevisiae, Pex19 is required for exit of Pex3 from the ER [156,158,172]. In vitro reconstitution of the initial step of de novo formation revealed that Pex19 is required for release of Pex3-containing vesicles from the ER [154,173]. However, how Pex19 mediates PMP targeting and budding from the ER is unclear. It is possible that Pex19 mediates recruitment to Pex3 at the ER of factors required for pre-peroxisomal vesicle formation. This may not be mechanistically different from the role of Pex19 in PMP targeting. For instance, in S. cerevisiae, Pex25, a member of the Pex11 family, has been reported to work with Pex3 to initiate formation of per-preoxisomal vesicles [174]. As Pex25 is an integral membrane protein, Pex19 may play a role in delivering it to Pex3 at the ER. Pex19 may also be involved in delivering to the ER factors required for vesicle formation. In general, vesicle budding from donor membranes involves recruitment of peripheral membrane proteins from the cytosol. Instead of handing the cargo over to Pex3 for insertion, peripheral membrane proteins may be handed over to other factors at the pre-peroxisomal membrane. Indeed, Pex19 has been shown to recruit the peripheral membrane proteins Vps1, Pex17 and Myo2 to peroxisomes via a motif resembling the mPTS1 [175–177].

Recently Tabak and colleagues proposed that, in S. cerevisiae, all newly synthesized PMPs insert into the ER membrane via the ER translocon or Get complex. The PMPs then sort to distinct ER subdomains that bud off and heterotypically fuse to bring together a complete set of PMPs, after which matrix protein import commences [161,162]. The peroxins Pex1 and Pex6 are proposed to mediate vesicle fusion. The import defect observed in pex1 and pex6 cells is proposed to result from a lack of fusion. According to this model, existing peroxisomes do not receive newly synthesized PMPs and therefore there is no need for a PMP insertion machinery in the peroxisomal membrane.

Although this model is attractive, it does not explain the strong experimental evidence that existing peroxisomes continue to receive newly synthesized PMPs and that peroxisomes multiply by dynamin-related protein fission [160,172,178–180]. Furthermore, since many S. cerevisiae PMPs contain Pex19 binding sites that are required for their targeting [137] and since the steady-state localization of Pex3 is at the peroxisomal membrane, it is very likely that, by analogy to mammalian cells, most PMPs are inserted directly into peroxisomal membranes. Pex3 that travels via the ER to peroxisomes may facilitate PMP insertion into the ER and into subsequent transport intermediates along the Pex3 trafficking pathway. The kinetics of ER to peroxisome transport and the availability of Pex3 for PMP import will determine whether newly synthesized PMPs are imported directly into peroxisomes or indirectly via the ER. This hypothesis is in line with a recent observation by the van der Klei group. In H. polymorpha cells lacking Pex3, non-ER membrane structures containing the PMPs Pex13 and Pex14 but lacking other PMPs were detected [118]. Upon reintroduction of Pex3, Pex3 accumulated in these structures and subsequently import of other PMPs commenced. Therefore, in this yeast, it is the localization of Pex3 that appears to direct the site of import.

The proposal that Pex1 and Pex6 are involved in fusion of precursor peroxisomes [161] is difficult to reconcile with previous studies. According to this model, the matrix protein import defect observed in pex1 and pex6 cells is a consequence of the docking complex (Pex13/Pex14) being present in separate vesicles from the exportomer complex (including Pex2/Pex10/Pex12). Many independent studies in various model systems have shown that Pex1 and Pex6 are required for recycling of ubiquitinated PTS receptors (see above). In pex1 and pex6 cells, PTS receptors accumulate at a postdocking stage at the peroxisomal membrane and are poly-ubiquitinated by the transmembrane RING finger ubiquitin ligases that are part of the exportomer complex, before their degradation. These observations indicate that docking complex and exportomer are present in the same membrane structures in pex1 and pex6 cells. On the other hand, in support of Tabak’s model, Y. lipolytica Pex1 and Pex6 have been suggested to function in membrane fusion events. Clearly, these observations are not easily explained and further investigations will be needed to resolve this apparent contradiction.

Concluding Remarks and Considerations

Although our understanding of peroxisome biology has been much advanced, there remain several huge gaps in our knowledge and in some unexplained observations. One such gap is the mechanism by which matrix proteins and membrane proteins are translocated across/into the peroxisomal membrane. A translocon for matrix proteins has been identified, but its transient and dynamic nature has hampered detailed biochemical characterization. Innovative approaches will be required to unravel the mysteries of translocation into the peroxisomal matrix.

With respect to delivery of PMPs to peroxisomes, there is evidence for two routes, one direct and one indirect via the ER. A particular PMP may follow
different routes to peroxisomes but the factors that decide which route it will follow are not clear. Mechanistic insights are limiting and it is entirely unclear how PMPs with multiple membrane spans are inserted into the peroxisomal membrane.

Other major gaps in our understanding include the mechanisms that regulate peroxisome dynamics. What determines whether peroxisomes multiply via growth and division or via de novo formation from the ER, and what are the signals for peroxisome degradation? Are there new machineries to be discovered or are all the peroxins known? Genetic approaches initially identified peroxins as factors for matrix protein import or membrane biogenesis. However, it is becoming clear that many peroxins have multiple functions. For instance, Pex3 has been implicated in peroxisome segregation and pexophagy in various yeast species [181–183]; Pex19 functions in peroxisome fission and segregation [175,177], and Pex14 has been implicated in pexophagy and peroxisome motility [184–186]. As discussed above, the exportomer factors Pex1 and Pex6 have been proposed to also mediate fusion events during peroxisome formation in Y. lipolytica and S. cerevisiae [161,187].

With many peroxins being involved in a variety of aspects of peroxisome dynamics and with different modes of peroxisome multiplication and various routes for PMP targeting that may display partial overlap, it is not surprising that studies on peroxisomal membrane biogenesis have resulted in controversies and unexplained observations. Careful dissection of the different functions of peroxins under controlled growth conditions is an important avenue to follow and may unravel the mechanisms that coordinate various aspects of peroxisome dynamics and resolve some of the unexplained observations discussed above.

For the last 20 years, the research focused on understanding peroxisome biogenesis has advanced the field significantly. However, several studies clearly demonstrate the need to understand the degradation process, as it appears that the cell has an intricate process of detecting and destroying misassembled peroxisomes [118,188]. This indicates that caution must be taken when interpreting data using peroxisome biogenesis mutants, as increased turnover of peroxisomes and/or pre-peroxisomal structures can potentially skew the results. Only by understanding both processes may we be able to comprehend better the mechanisms governing peroxisome dynamics.

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References

[1] Fujiki Y, Yagita Y, Matsuzaki T. Peroxisome biogenesis disorders: molecular basis for impaired peroxisomal membrane assembly; in metabolic functions and biogenesis of peroxisomes in health and disease. Biochim Biophys Acta 2012; 1822:1337–42.
[2] Wanders RJ, Waterham HR. Biochemistry of mammalian peroxisomes revisited. Annu Rev Biochem 2006;75:295–332.
[3] Lazarow PB, Fujiki Y. Biogenesis of peroxisomes. Annu Rev Cell Biol 1985;1:489–530.
[4] Distel B, Erdmann R, Gould SJ, Blobel G, Crane DI, Gregg JM, et al. A unified nomenclature for peroxisome biogenesis factors. J Cell Biol 1996;135:1–3.
[5] Tabak HF, Braakman I, van der Zand A. Peroxisome formation and maintenance are dependent on the endoplasmic reticulum. Annu Rev Biochem 2013;82:723–44.
[6] Hettema EH, Erdmann R, van der Klei I, Veenhuis M. Evolving models for peroxisome biogenesis. Curr Opin Cell Biol 2014; 29C:25–30.
[7] Veenhuis M, van der Klei IJ. A critical reflection on the principles of peroxisome formation in yeast. Front Physiol 2014;5:110.
[8] Dimitrov L, Lam SK, Schekman R. The role of the endoplasmic reticulum in peroxisome biogenesis. Cold Spring Harb Perspect Biol 2013;5:a013243.
[9] Agrawal G, Subramani S. Emerging role of the endoplasmic reticulum in peroxisome biogenesis. Front Physiol 2013;4:286.
[10] Imanaka T, Small GM, Lazarow PB. Translocation of acyl-CoA oxidase into peroxisomes requires ATP hydrolysis but not a membrane potential. J Cell Biol 1987;105:2915–22.
[11] Small GM, Szabo LJ, Lazarow PB. Acyl-CoA oxidase contains two targeting sequences each of which can mediate protein import into peroxisomes. EMBO J 1988;7:1167–73.
[12] Lazarow PB, Thieringer R, Cohen G, Imanaka T, Small G. Protein import into peroxisomes in vitro. Methods Cell Biol 1991;34:303–26.
[13] Walton PA, Gould SJ, Rachubinski RA, Subramani S, Feramisco JR. Transport of microinjected alcohol oxidase from Pichia pastoris into vesicles in mammalian cells: involvement of the peroxisomal targeting signal. J Cell Biol 1992;118:499–508.
[14] Rapp S, Soto U, Just WW. Import of firefly luciferase into peroxisomes of permeabilized Chinese hamster ovary cells: a model system to study peroxisomal protein import in vitro. Exp Cell Res 1992;205:59–65.
Protein transport to peroxisomes

[15] Wendland M, Subramani S. Cytosol-dependent peroxisomal protein import in a permeabilized cell system. J Cell Biol 1993;120:675–85.

[16] Leon S, Goodman JM, Subramani S. Uniqueness of the mechanism of protein import into the peroxisome matrix: transport of folded, co-factor-bound and oligomeric proteins by shuttling receptors. Biochim Biophys Acta 2006;1763:1552–64.

[17] Brul S, Wiener EA, Westerveld A, Strijland A, Wanders RJ, Schram AW, et al. Kinetics of the assembly of peroxisomes after fusion of complementary cell lines from patients with the cerbro-hepato-renal (Zellweger) syndrome and related disorders. Biochem Biophys Res Commun 1988;152:1083–9.

[18] Walton PA, Gould SJ, Feramisco JR, Subramani S. Transport of microinjected proteins into peroxisomes of mammalian cells: inability of Zellweger cell lines to import proteins with the SKL tripeptide peroxisomal targeting signal. Mol Cell Biol 1992;12:531–41.

[19] McNew JA, Goodman JM. An oligomeric protein is imported into peroxisomes in vivo. J Cell Biol 1994;127:1245–57.

[20] Glover JR, Andrews DW, Rachubinski RA. Saccharomyces cerevisiae peroxisomal thiolase is imported as a dimer. Proc Natl Acad Sci U S A 1994;91:10541–5.

[21] Elgersma Y, Vos A, van den Berg M, van Roermund CW, van der Sluijs P, Distel B, et al. Analysis of the carboxyl-terminal peroxisomal targeting signal 1 in a homologous context in Saccharomyces cerevisiae. J Biol Chem 1996;271:26375–82.

[22] Islinger M, Li KW, Seitz J, Volki A, Luers GH. Hitchhiking of Cu/Zn superoxide dismutase to peroxisomes—evidence for a natural piggyback import mechanism in mammals. Traffic 2008;10:1711–21.

[23] Otera H, Fujiki Y. Pex5p imports folded tetrameric catalase by interaction with Pex13p. Traffic 2012;13:1364–77.

[24] Walton PA, Hill PE, Subramani S. Import of stably folded proteins into peroxisomes. Mol Biol Cell 1995;6:675–83.

[25] Brocard CB, Jedeszko C, Song HC, Terlecky SR, Walton PA. Protein structure and import into the peroxisomal matrix. Traffic 2003;4:74–82.

[26] Freitas MO, Francisco T, Rodrigues TA, Alencastre IS, Pinto MP, Grou CP, et al. PEX5 protein binds monomeric catalase, blocking its tetramerization and releases it upon binding the N-terminal domain of PEX14. J Biol Chem 2011;286:40509–19.

[27] Crookes WJ, Olsen LJ. The effects of chaperones and the influence of protein assembly on peroxisomal protein import. J Biol Chem 1998;273:17236–42.

[28] Waterham HR, Russell KA, Vries Y, Cregg JM. Peroxisomal targeting, import, and assembly of alcohol oxidase in Pichia pastoris. J Cell Biol 1997;139:1419–31.

[29] Faber KN, van Dijk R, Keizer-Gunnink I, Koek A, van der Klei J, Veenhuis M. Import of assembled PTS1 proteins into peroxisomes of the yeast Hansenula polymorpha: yes and no! Biochim Biophys Acta 2002;1591:157–62.

[30] Tanaka N, Aoki K, Ishikura S, Nagano M, Imamura Y, Hara A, et al. Molecular basis for peroxisomal localization of tetrameric carboxyl reductase. Structure 2008;16:388–97.

[31] Gould SG, Keller GA, Subramani S. Identification of a peroxisomal targeting signal at the carboxy terminus of firefly luciferase. J Cell Biol 1987;105:2923–31.

[32] Gould SJ, Keller GA, Subramani S. Identification of peroxisomal targeting signals located at the carboxy terminus of four peroxisomal proteins. J Cell Biol 1988;107:897–905.

[33] Gould SJ, Keller GA, Hosken N, Wilkinson J, Subramani S. A conserved tripeptide sorts proteins to peroxisomes. J Cell Biol 1989;108:1657–64.

[34] Gould SJ, Keller GA, Schneider M, Howell SH, Garratt LJ, Goodman JM, et al. Peroxisomal protein import is conserved between yeast, plants and mammals. EMBO J 1990;9:85–90.

[35] Lametschwandtner G, Brocard C, Fransen M, Van Veldhoven P, Berger J, Hartig A. The difference in recognition of terminal tripeptides as peroxisomal targeting signal 1 between yeast and human is due to different affinities of their receptor Pex5p to the cognate signal and to residues adjacent to it. J Biol Chem 1998;273:33635–43.

[36] Maynard EL, Gatto GJ, Berg JM. Pex5p binding affinities for canonical and noncanonical PTS1 peptides. Proteins 2004;55:856–61.

[37] Brocard C, Hartig A. Peroxisome targeting signal 1: is it really a simple tripeptide? Biochim Biophys Acta 2006;1763:1565–73.

[38] Chowdhary G, Kataya AR, Lingner T, Reumann S. Non-canonical peroxisome targeting signals: identification of novel PTS1 tripeptides and characterization of enhancer elements by computational permutation analysis. BMC Plant Biol 2012;12:142.

[39] Gatto GJ, Geisbrecht BV, Gould SJ, Berg JM. Peroxisomal targeting signal-1 recognition by the TPR domains of human PEX5. Nat Struct Biol 2000;7:1091–5.

[40] Stanley WA, Filipov FV, Kurasula P, Schiller N, Erdmann R, Schliebs W, et al. Recognition of a functional peroxisome type 1 target by the dynamic import receptor pex5p. Mol Cell 2006;24:653–63.

[41] Swinkels BW, Gould SJ, Bodnar AG, Rachubinski RA, Subramani S. A novel, cleavable peroxisomal targeting signal at the amino-terminus of the rat 3-ketoacyl-CoA thiolase. EMBO J 1991;10:3255–62.

[42] Osumi T, Tsukamoto T, Hata S, Yokota S, Miura S, Fujiki Y, et al. Amino-terminal presequence of the precursor of peroxisomal 3-ketoacyl-CoA thiolase is a cleavable signal peptide for peroxisomal targeting. Biochem Biophys Res Commun 1991;181:947–54.

[43] Petrivo OI, Tang L, Titorenko VI, Rachubinski RA. A new definition for the consensus sequence of the peroxisomal targeting signal type 1. J Mol Biol 2004;341:119–34.

[44] Motley AM, Hettema EH, Ketting R, Plasterk R, Tabak HF. Caenorhabditis elegans has a single pathway to target matrix proteins to peroxisomes. EMBO Rep 2000;1:40–6.

[45] Faust JE, Verma A, Peng C, McNew JA. An inventory of peroxisomal proteins and pathways in Drosophila melanogaster. Traffic 2012;13:1378–92.

[46] Fodor K, Wolf J, Erdmann R, Schliebs W, Wilmanns M. Molecular requirements for peroxisomal targeting of alanine-glyoxylate aminotransferase as an essential determinant in primary hyperoxaluria type 1. PLoS Biol 2012;10:e1001309.

[47] Einwachter H, Sowinski S, Kunau WH, Schliebs W, Yarrowia lipolytica. Yarrowia lipolytica Pex20p, Saccharomyces cerevisiae Pex18p/Pex21p and mammalian Pex5pL fulfil a common tripeptide recognition for a natural piggyback import mechanism in mammals. Traffic 2008;10:1711–21.

[48] Otera H, Fujiki Y. Pex5p imports folded tetrameric catalase by interaction with Pex13p. Traffic 2012;13:1364–77.

[49] Walton PA, Hill PE, Subramani S. Import of stably folded proteins into peroxisomes. Mol Biol Cell 1995;6:675–83.

[50] Brocard CB, Jedeszko C, Song HC, Terlecky SR, Walton PA. Protein structure and import into the peroxisomal matrix. Traffic 2003;4:74–82.

[51] Freitas MO, Francisco T, Rodrigues TA, Alencastre IS, Pinto MP, Grou CP, et al. PEX5 protein binds monomeric catalase, blocking its tetramerization and releases it upon binding the N-terminal domain of PEX14. J Biol Chem 2011;286:40509–19.

[52] Crookes WJ, Olsen LJ. The effects of chaperones and the influence of protein assembly on peroxisomal protein import. J Biol Chem 1998;273:17236–42.

[53] Waterham HR, Russell KA, Vries Y, Cregg JM. Peroxisomal targeting, import, and assembly of alcohol oxidase in Pichia pastoris. J Cell Biol 1997;139:1419–31.

[54] Faber KN, van Dijk R, Keizer-Gunnink I, Koek A, van der Klei J, Veenhuis M. Import of assembled PTS1 proteins into peroxisomes of the yeast Hansenula polymorpha: yes and no! Biochim Biophys Acta 2002;1591:157–62.

[55] Tanaka N, Aoki K, Ishikura S, Nagano M, Imamura Y, Hara A, et al. Molecular basis for peroxisomal localization of tetrameric carboxyl reductase. Structure 2008;16:388–97.

[56] Gould SG, Keller GA, Subramani S. Identification of a peroxisomal targeting signal at the carboxy terminus of firefly luciferase. J Cell Biol 1987;105:2923–31.

[57] Gould SJ, Keller GA, Subramani S. Identification of peroxisomal targeting signals located at the carboxy terminus of four peroxisomal proteins. J Cell Biol 1988;107:897–905.
Protein transport to peroxisomes

PpPex20p: ubiquitin-dependent localization and regulation. J Cell Biol 2006;172:67–78.

[50] Purdue PE, Yang X, Lazarow PB. Pex18p and Pex21p, a novel pair of related peroxins essential for peroxisomal targeting by the PTS2 pathway. J Cell Biol 1996;143:1859–69.

[51] Matsumura T, Otera H, Fujiki Y. Disruption of the interaction of the longer isoform of Pex5p, Pex5pL, with Pex7p abolishes peroxisome targeting signal type 2 protein import in mammals. Study with a novel Pex5-impaired Chinese hamster ovary cell mutant. J Biol Chem 2000;275:21715–21.

[52] Dodt G, Warren D, Becker E, Rehling P, Gould SJ. Domain mapping of human PEX5 reveals functional and structural similarities to Saccharomyces cerevisiae Pex18p and Pex21p. J Biol Chem 2001;276:41769–81.

[53] Woodward AW, Bartel B. The Arabidopsis peroxisomal targeting signal type 2 receptor PEX7 is necessary for peroxisome function and dependent on PEX5. Mol Biol Cell 2005;16:573–83.

[54] Jung S, Morelli M, Rachubinski RA, Goodlett DR, Aitchison JD. Dynamic changes in the subcellular distribution of Gpd1p1 in response to cell stress. J Biol Chem 2010;285:6739–49.

[55] Klein AT, van den Berg M, Botterg G, Tabak HF, Distel B. Saccharomyces cerevisiae acyl-CoA oxidase follows a novel, non-PTS1, import pathway into peroxisomes that is dependent on Pex5p. J Biol Chem 2002;277:25011–9.

[56] van der Klei IJ, Veenhuis M. PTS1-independent sorting of peroxisomal matrix proteins by Pex5p. Biochim Biophys Acta 2006;1763:1794–800.

[57] Dodt G, Gould SJ. Multiple PEX genes are required for proper subcellular distribution and stability of Pex5p, the PTS1 receptor: evidence that PTS1 protein import is mediated by a cycling receptor. J Cell Biol 1996;135:1763–74.

[58] Kunau WH. Peroxisomes: the extended shuttle to the peroxisomal matrix. Curr Biol 2001;11:R659–62.

[59] Dammal V, Subramani S. The human peroxisomal targeting signal receptor, Pex5p, is translocated into the peroxisomal membrane and recycled to the cytosol. Cell 2001;105:187–96.

[60] Nair DM, Purdue PE, Lazarow PB. Pex7p translocates in and out of peroxisomes in Saccharomyces cerevisiae. J Cell Biol 2004;167:599–604.

[61] Agne B, Meindl NM, Niederhoff K, Einwachter H, Rehling P, Sickmann A, et al. Pex5p: an intraperoxisomal organizer of the peroxisomal import machinery. Mol Cell 2003;11:635–46.

[62] Azuvedo JE, Schliebs W. Pex14p, more than just a docking receptor. Biochim Biophys Acta 2006;1763:1574–84.

[63] Williams C, Distel B. Pex13p: docking or cargo handling protein? Biochim Biophys Acta 2006;1763:1585–91.

[64] Lanyon-Hogg T, Warriner SL, Baker A. Getting a camel through the eye of a needle: the import of folded proteins by peroxisomes. Biol Cell 2010;102:245–63.

[65] Mukai S, Fujiki Y. Molecular mechanisms of import of peroxisome-targeting signal type 2 (PTS2) proteins by PTS2 receptor Pex7p and PTS1 receptor Pex5pL. J Biol Chem 2006;281:37311–20.

[66] Grunau S, Schliebs W, Linnepe R, Neufeld C, Cizmowski C, Reinartz B, et al. Peroxisomal targeting of PTS2 pre-import complexes in the yeast Saccharomyces cerevisiae. Traffic 2009;10:451–60.

[67] Natsumaya R, Okumoto K, Fujiki Y. Pex5p stabilizes Pex14p: a study using a newly isolated pex5 CHO cell mutant, ZPEG101. Biochem J 2013;449:195–207.

[68] Miyata N, Fujiki Y. Shuttling mechanism of peroxisome targeting signal type 1 receptor Pex5: ATP-independent import and ATP-dependent export. Mol Cell Biol 2005;25:10822–32.

[69] Urquhart AJ, Kennedy D, Gould SJ, Crane DJ. Interaction of Pex5p, the type 1 peroxisomal targeting signal receptor, with the peroxisomal membrane proteins Pex14p and Pex13p. J Biol Chem 2000;275:4127–36.

[70] Erdmann R, Schliebs W. Peroximal matrix protein import: the transient pore model. Nat Rev Mol Cell Biol 2005;6:738–42.

[71] Gouveia AM, Reguenca C, Oliveira ME, Sa-Miranda C, Azuvedo JE. Characterization of peroxisomal Pex5p from rat liver. Pex5p in the Pex5p-Pex14p membrane complex is a transmembrane protein. J Biol Chem 2000;275:32444–51.

[72] Platta HW, Grunau S, Rosenkranz K, Girzalsky W, Erdmann R. Functional role of the AAA peroxins in dislocation of the cycling PTS1 receptor back to the cytosol. Nat Cell Biol 2005;7:817–22.

[73] Gouveia AM, Guimaraes CP, Oliveira ME, Sa-Miranda C, Azuvedo JE. Insertion of Pex5p into the peroxisomal membrane is cargo protein-dependent. J Biol Chem 2003;278:4389–92.

[74] Reguenca C, Oliveira ME, Gouveia AM, Sa-Miranda C, Azuvedo JE. Characterization of the mammalian peroxisomal import machinery: Pex2p, Pex5p, Pex12p, and Pex14p are subunits of the same protein assembly. J Biol Chem 2001;276:29935–42.

[75] Stanley WA, Wilmanns M. Dynamic architecture of the peroxisomal import receptor Pex5p. Biochim Biophys Acta 2006;1763:1592–9.

[76] Ma C, Schumann U, Rayapuram N, Subramani S. The peroxisomal matrix import of Pex6p requires only PTS receptors and Pex14p. Mol Biol Cell 2009;20:3680–9.

[77] Meinecke M, Cizmowski C, Schliebs W, Kruger V, Beck S, Wagner R, et al. The peroxisomal importome constitutes a large and highly dynamic core. Nat Cell Biol 2010;12:273–7.

[78] Cyr N, Madrid KP, Strasser R, Aurousseau M, Finn R, Ausio J, et al. Leishmania donovani peroxin 14 undergoes a marked conformational change following association with peroxin 5. J Biol Chem 2008;283:31488–99.

[79] Ma C, Hagstrom D, Polley SG, Subramani S. Redox-regulated cargo binding and release by the peroxisomal targeting signal receptor, Pex5. J Biol Chem 2013;288:27220–31.

[80] Lanyon-Hogg T, Hooper J, Gunn S, Warriner SL, Baker A. PEX14 binding to Arabidopsis PEX5 has differential effects on PTS1 and PTS2 SNARE cargo occupancy of the receptor. FEBS Lett 2014;588:2223–9.

[81] Francisco T, Rodrigues TA, Freitas MO, Grou CP, Carvalho AF, Sa-Miranda C, et al. A cargo-centered perspective on the PEX5 receptor-mediated peroxisomal protein import pathway. J Biol Chem 2013;288:29151–9.

[82] Oliveira ME, Gouveia AM, Pinto RA, Sa-Miranda C, Azuvedo JE. The energetics of Pex5p-mediated peroxisomal protein import. J Biol Chem 2003;278:39483–8.

[83] Platta HW, Hagen S, Reidick C, Erdmann R. The peroxisomal receptor dislocation pathway: to the exporter and beyond. Biochimie 2014;96:16–28.

[84] Kragt A, Voorn-Brouwer T, van den Berg M, Distel B. The Saccharomyces cerevisiae peroxisomal import receptor Pex5p is monoubiquitinated in wild type cells. J Biol Chem 2005;280:7867–74.

[85] Carvalho AF, Pinto MP, Grou CP, Alencastre IS, Fransen M, Sa-Miranda C, et al. Ubiquitination of mammalian Pex5p, the peroxisomal import receptor. J Biol Chem 2007;282:31267–72.

[86] Williams C, van den Berg M, Sprenger RR, Distel B. A conserved cysteine is essential for Pex4p-dependent ubiquitination of the peroxisomal import receptor Pex5p. J Biol Chem 2007;282:22534–43.
Debelyy MO, Platta HW, Saffian D, Hensel A, Thoms S, Meyer HE, et al. Ub1p5, a ubiquitin hydrolase associated with the peroxisomal export machinery. J Biol Chem 2011;286:28223–34.

Grou CP, Francisco T, Rodrigues TA, Freitas MO, Pinto MP, Carvalho AF, et al. Identification of ubiquitin-specific protease 9X (USP9X) as a deubiquitinase acting on ubiquitin-peroxin 5 (PEX5) thioester conjugate. J Biol Chem 2012;287:12815–27.

Schliebs W, Girzalsky W, Erdmann R. Peroxisomal protein import and ERAD: variations on a common theme. Nat Rev Mol Cell Biol 2010;11:885–90.

Hensel A, Beck S, El Magraoui F, Platta HW, Girzalsky W, Erdmann R. Cysteine-dependent ubiquitination of Pex18p is linked to cargo translocation across the peroxisomal membrane. J Biol Chem 2011;286:43495–505.

Liu X, Subramani S. Unique requirements for mono- and polyubiquitination of the peroxisomal targeting signal co-receptor, Pex20. J Biol Chem 2013;288:7239–40.

Rodrigues TA, Alencastre IS, Francisco T, Brites P, Fransen M, Grou CP, et al. A PEX7-centered perspective on the peroxisomal targeting signal type 2-mediated protein import pathway. Mol Cell Biol 2014;34:297–28.

Santos MJ, Imanaka T, Shio H, Small GM, Lazarow PB. Peroxisomal membrane ghosts in Zellweger syndrome—aberrant organelle assembly. Science 1988;239:1536–8.

Tager JM, Brul S, Wiemer EA, Strijland A, Van Driel R, Schutgens RB, et al. Genetic relationship between the Zellweger syndrome and other peroxisomal disorders characterized by an impairment in the assembly of peroxisomes. Prog Clin Biol Res 1990;321:545–58.

Honsho M, Tamura S, Shimozawa N, Suzuki Y, Kondo N, Fujiki Y. Mutation in Pex16 is causal in the peroxisome-deficient Zellweger syndrome of complementation group D. Am J Hum Genet 1998;63:1622–30.

Kinoshita N, Ghaedi K, Shimozawa N, Wanders RJ, Matuszono Y, Imanaka T, et al. Newly identified Chinese hamster ovary cell mutants are defective in biogenesis of peroxisomal membrane vesicles (peroxisomal ghosts), representing a novel complementation group in mammals. J Biol Chem 1998;273:24122–30.

Matuszono Y, Kinoshita N, Tamura S, Shimozawa N, Hamasaki M, Ghaedi K, et al. Human Pex19: cDNA cloning by functional complementation, mutation analysis in a patient with Zellweger syndrome, and potential role in peroxisomal membrane biogenesis. Proc Natl Acad Sci U S A 1999;96:2116–21.

South ST, Gould SJ. Peroxisome synthesis in the absence of preexisting peroxisomes. J Cell Biol 1999;144:255–66.

Sacksteder KA, Jones JM, South ST, Li X, Liu Y, Gould SJ. Pex19 binds multiple peroxisomal membrane proteins, is predominantly cytoplasmic, and is required for peroxisome membrane synthesis. J Cell Biol 2000;148:931–44.

Hettema EH, Girzalsky W, van Den Berg M, Erdmann R, Distel B. Saccharomyces cerevisiae pex3p and pex19p are required for proper localization and stability of peroxisomal membrane proteins. EMBO J 2000;19:223–33.

Muntau AC, Mayerhofer PU, Paton BC, Kamaroser S, Roscher AA. Defective peroxisomal membrane synthesis due to mutations in human Pex4 causes Zellweger syndrome, complementation group G. Am J Hum Genet 2000;67:967–75.

Koek A, Komori M, Veenhuis M, van der Klei U. A comparative study of peroxisomal structures in Hansenula polymorpha pex mutants. FEMS Yeast Res 2007;7:1126–33.
[118] Knoops K, Manivannan S, Cepinska MN, Krikken AM, Kram AM, Veenhuis M, et al. Preperoxisomal vesicles can form in the absence of Pex3. J Cell Biol 2004;204:659–68.

[119] Haza PP, Suriaipranata I, Snyder WB, Subramani S. Peroxisome remnants in pex3delta cells and the requirement of Pex3p for interactions between the peroxisomal docking and translocation subcomplexes. Traffic 2002;3:560–74.

[120] Fujiy Y, Rachubinski RA, Lazaro PB. Synthesis of a major integral membrane polypeptide of rat liver peroxisomes on free polyosomes. Proc Natl Acad Sci U S A 1984;81:7127–31.

[121] Suzuki Y, Oir T, Takiguchi M, Mori M, Hijikata M, Hashimoto T. Biosynthesis of membrane polypeptides of rat liver peroxisomes. J Biochem 1987;101:491–6.

[122] Diestelkotter P, Just WW. In vitro insertion of the 22-kD peroxisomal membrane protein into isolated rat liver peroxisomes. J Cell Biol 1993;123:1717–25.

[123] Imanaka T, Shina Y, Takan T, Hashimoto T, Osumi T. Insertion of the 70-kDa peroxisomal membrane protein into peroxisomal membranes in vivo and in vitro. J Biol Chem 1996;271:3706–13.

[124] Matsuzono Y, Fujiy Y. In vitro transport of membrane proteins to peroxisomes by shuttling receptor Pex19p. J Biol Chem 2006;281:36–42.

[125] Pinto MP, Grou CP, Alencastre IS, Oliveira ME, Sa-Miranda C, Fransen M, et al. The import competence of a peroxisomal membrane protein is determined by Pex19p before the docking step. J Biol Chem 2006;281:34982–902.

[126] Yagita Y, Hirokasa T, Fujiy Y. Tail-anchored PEX26 targets peroxisomes via a Pex19-dependent and TRC40-independent class I pathway. J Cell Biol 2013;200:651–66.

[127] Dyer JM, McNew JA, Goodman JM. The sorting sequence of the integral peroxisomal membrane protein PMP47 is contained within a short hydrophilic loop. J Cell Biol 1996;133:269–80.

[128] Baerends RJ, Faber KN, Kram AM, Kiel JA, van der Kleij U, Veenhuis M. A stretch of positively charged amino acids at the N terminus of Hansenula polymorpha Pex3p is involved in incorporation of the protein into the peroxisomal membrane. J Biol Chem 2000;275:9986–95.

[129] Jones JM, Morrell JC, Gould SJ. Multiple distinct targeting signals in integral peroxisomal membrane proteins. J Cell Biol 2001;153:1141–50.

[130] Honsho M, Fujiy Y. Topogenesis of peroxisomal membrane protein requires a short, positively charged intervening-loop sequence and flanking hydrophobic segments. study using human membrane protein PMP34. J Biol Chem 2001;276:9375–82.

[131] Wang X, Unruh MJ, Goodman JM. Discrete targeting signals direct Pmp47 to oleate-induced peroxisomes in Saccharomyces cerevisiae. J Biol Chem 2001;276:10897–905.

[132] Fransen M, Wylt T, Brees C, Mannata GP, Van Veldhoven PP. Human Pex19p binds peroxisomal integral membrane proteins at regions distinct from their sorting sequences. Mol Cell Biol 2001;21:4413–24.

[133] Honsho M, Hiroshige T, Fujiy Y. The membrane biogenesis peroxin Pex16p. Topogenesis and functional roles in peroxisomal membrane assembly. J Biol Chem 2002;277:44513–24.

[134] Brosius U, Dehmel T, Gartner J. Two different targeting signals direct human peroxisomal membrane protein 22 to peroxisomes. J Biol Chem 2002;277:774–84.

[135] Biermanns M, von Laar J, Brosius U, Gartner J. The peroxisomal membrane targeting elements of human peroxin 2 (PEX2). Eur J Cell Biol 2003;82:155–62.

[136] Murphy MA, Phillipson BA, Baker A, Mullen RT. Characterization of the targeting signal of the Arabidopsis 22-kD integral peroxisomal membrane protein. Plant Physiol 2003;133:813–28.

[137] Rottensteiner H, Kramer A, Lorenzen S, Stein K, Landgraf C, Vollmer-Engert R, et al. Peroxisomal membrane proteins contain common Pex1p-binding sites that are an integral part of their targeting signals. Mol Biol Cell 2004;15:3406–17.

[138] Muntau AC, Roscher AA, Kunau WH, Dodt G. Interaction of Pex3 and Pex19 visualized by fluorescence resonance energy transfer (FRET). Adv Exp Med Biol 2003;544:221–4.

[139] Fang Y, Morrell JC, Jones JM, Gould SJ. Pex3 functions as a Pex19 docking factor in the import of class I peroxisomal membrane proteins. J Cell Biol 2004;164:863–75.

[140] Shibata H, Kashiwayama Y, Imanaka T, Kato H. Domain architecture and activity of human Pex19p, a chaperone-like protein for intracellular trafficking of peroxisomal membrane proteins. J Biol Chem 2004;279:38486–94.

[141] Kashiwayama Y, Asahina K, Shibata H, Morita M, Muntau AC, Roscher AA, et al. Role of Pex19p in the targeting of PMP70 to peroxisomes. Biochim Biophys Acta 2005;1746:116–28.

[142] Chen Y, Piechot L, Loh RA, Yang J, Kari TM, Wong JY, et al. Hydrophobic handoff for direct delivery of peroxisomal tail-anchored proteins. Nat Commun 2014;5:5790.

[143] Jones JM, Morrell JC, Gould SJ. Pex19 is a predominantly cytosolic chaperone and import receptor for class I peroxisomal membrane proteins. J Cell Biol 2004;164:57–67.

[144] Sato Y, Shibata H, Nakatsu T, Nakano H, Kashiwayama Y, Imanaka T, et al. Structural basis for docking of peroxisomal membrane protein carrier Pex19p onto its receptor Pex3p. EMBO J 2010;29:4083–93.

[145] Schmidt F, Dietrich D, Eylestein R, Groemping Y, Stehle T, Dodt G. The role of conserved Pex3 regions in Pex19-binding and peroxisome biogenesis. Traffic 2012;13:1244–60.

[146] Schmidt F, Treibler N, Zocher G, Bjelic S, Steinmetz MO, Kalbacher H, et al. Insights into peroxisome function from the structure of Pex3 in complex with a soluble fragment of Pex19. J Biol Chem 2010;285:25410–7.

[147] Pinto MP, Grou CP, Fransen M, Sa-Miranda C, Azvedo JE. The cytosolic domain of Pex3, a protein involved in the biogenesis of peroxisomes, binds membrane lipids. Biochim Biophys Acta 2009;1793:1669–75.

[148] Schweller N, Holton SJ, Fodor K, Milewski M, Konarev P, Stanley WA, et al. The peroxisomal receptor Pex19p forms a helical mPfTS recognition domain. EMBO J 2010;29:2491–500.

[149] Titorenko VI, Ogrydziak DM, Rachubinski RA. Four distinct secretory pathways serve protein secretion, cell surface growth, and peroxisome biogenesis in the yeast Yarrowia lipolytica. Mol Cell Biol 1997;17:3210–26.

[150] Titorenko VI, Rachubinski RA. Mutants of the yeast Yarrowia lipolytica defective in protein exit from the endoplasmic reticulum are also defective in peroxisome biogenesis. Mol Microbiol 1997;24:455–67.

[151] Mullen RT, Lisenbee CS, Miernyk JA, Trelease RN. Peroxisomal membrane ascorbate peroxidase is sorted to a membranous network that resembles a subdomain of the endoplasmic reticulum. Plant Cell 1999;11:2167–85.

[152] Halbach A, Rucktashel R, Rottensteiner H, Ermann R. The N-domain of Pex22p can functionally replace the Pex3p N-domain in targeting and peroxisome formation. J Biol Chem 2009;284:3906–16.
[153] Toro AA, Araya CA, Cordova GJ, Arnedondo CA, Cardenas HG, Moreno RE, et al. Pex3p-dependent peroxisomal biogenesis initiates in the endoplasmic reticulum of human fibroblasts. J Cell Biochem 2009;107:1083–96.

[154] Lam SK, Yoda N, Schekman R. A vesicle carrier that mediates peroxisome protein traffic from the endoplasmic reticulum. Proc Natl Acad Sci U S A 2010;107:21523–8.

[155] Aranovich A, Hua R, Rutenberg AD, Kim PK. Pex16 contributes to peroxisome maintenance by constantly trafficking Pex3 via the ER. J Cell Sci 2014;127:3675–86.

[156] Hoepfner D, Schildkret D, Braakman I, Philippson P, Tabak HF. Contribution of the endoplasmic reticulum to peroxisome formation. Cell 2005;122:85–95.

[157] Krat G, Voom-Brouwer T, van den Berg M, Distel B. Endoplasmic reticulum-directed Pex3p routes to peroxisomes and restores peroxisome formation in a Saccharomyces cerevisiae pex3Δdelta strain. J Biol Chem 2005;280:34350–7.

[158] Tam YY, Fagarasanu A, Fagarasanu M, Rachubinski RA. Pex3p initiates the formation of a preperoxisomal compartment from a subdomain of the endoplasmic reticulum in Saccharomyces cerevisiae. J Biol Chem 2005;280:34393–9.

[159] Thoms S, Harms I, Kales KU, Gartner J. Peroxisome formation requires the endoplasmic reticulum channel protein Sec61. Traffic 2012;13:599–609.

[160] Fakieh MH, Drake PJ, Lacey J, Muncck JM, Motley AM, Hettema EH. Intra-ER sorting of the peroxisomal membrane protein Pex3 relies on its luminal domain. Biol Open 2013;2:829–37.

[161] van der Zand A, Gent J, Braakman I, Tabak HF. Biochemically distinct vesicles from the endoplasmic reticulum fuse to form peroxisomes. Cell 2012;149:397–409.

[162] van der Zand A, Braakman I, Tabak HF. Peroxisomal membrane proteins insert into the endoplasmic reticulum. Mol Cell 2010;21:2057–65.

[163] Johnson N, Powis K, High S. Post-translational translocation into the endoplasmic reticulum. Biochim Biophys Acta 2013;1833:2403–9.

[164] Borgese N, Fasana E. Targeting pathways of C-tail-anchored proteins. Biochim Biophys Acta 2011;1808:937–46.

[165] Stefanovic S, Hegde RS. Identification of a targeting factor for posttranslational membrane protein insertion into the ER. Cell 2007;128:1147–59.

[166] Schuldiner M, Metz J, Schmid V, Denic V, Rakwalska M, Schmitt HD, et al. The GET complex mediates insertion of tail-anchored proteins into the ER membrane. Cell 2008;134:634–45.

[167] Elgersma Y, Kwast L, van den Berg M, Snyder WB, Distel B, Schuldiner M, Metz J, Schmid V, Denic V, Rakwalska M, Schmitt HD, et al. The GET complex mediates insertion of tail-anchored proteins into the ER membrane. Cell 2008;134:634–45.

[168] Delille HK, Agricola B, Guimaraes SC, Borta H, Luers GH, Fransen M, et al. Pex11pbeta-mediated growth and division of mammalian peroxisomes follows a maturation pathway. J Cell Sci 2010;123:2750–62.

[169] Motley AM, Hettema EH. Yeast peroxisomes multiply by growth and division. J Cell Biol 2007;178:399–410.

[170] Yonekawa S, Funano A, Baka T, Fujiki Y, Ogasaara Y, Yamamoto A, et al. Sec16β is involved in the endoplasmic reticulum export of the peroxisomal membrane biogenesis factor peroxin 16 (Pex16) in mammalian cells. Proc Natl Acad Sci U S A 2011;108:12746–51.

[171] Vizeacoumar FJ, Vreden WN, Fagarasanu M, Eltzen GA, Aitchison JD, Rachubinski RA. The dynamin-like protein Vps1p of the yeast Saccharomyces cerevisiae associates with peroxisomes in a Pex19p-dependent manner. J Biol Chem 2006;281:12817–23.

[172] Otzen M, Rucktaschel R, Thoms S, Emmrich K, Krikken AM, Erdmann R, et al. Pex19p contributes to peroxisome inheritance in the association of peroxisomes to Myo2p. Traffic 2012;13:947–59.

[173] Nagolu S, Saraya R, Otzen M, Veenhuis M, van der Klei IJ. Peroxisome proliferation in Hansenula polymorpha requires Dnm1p which mediates fission but not de novo formation. Biochim Biophys Acta 2008;1783:760–9.

[174] Menendez-Benito V, van Deventer SJ, Jimenez-Garcia V, Roy-Luzarraga M, van Leeuwen F, Neeffes J, Spatiotemporal analysis of organelle and macromolecular complex inheritance. Proc Natl Acad Sci U S A 2013;110:175–80.

[175] Knoblach B, Rachubinski RA. Doing the math: how yeast cells maintain their peroxisome populations. Commun Integr Biol 2013;6:e26901.

[176] Munck JM, Motley AM, Nuttall JM, Hettema EH. A dual function for Pex3p in peroxisome formation and inheritance. J Cell Biol 2009;187:463–71.

[177] Chang J, Mast FD, Fagarasanu A, Rachubinski DA, Eitzen GA, Dacks JB, et al. Pex3p peroxisome biogenesis proteins function for Pex3p in peroxisome formation and inheritance. J Cell Biol 2010;187:463–71.

[178] Huber A, Koch J, Kragler F, Brocard C, Hartig A. A subtle interplay between three Pex11 proteins shapes de novo formation and fission of peroxisomes. Traffic 2012;13:157–67.

[179] Vizeacoumar FJ, Vreden WN, Fagarasanu M, Eltzen GA, Aitchison JD, Rachubinski RA. The dynamin-like protein Vps1p of the yeast Saccharomyces cerevisiae associates with peroxisomes in a Pex19p-dependent manner. J Biol Chem 2006;281:12817–23.

[180] Girzalsky W, Hoffmann LS, Schemenewitz A, Nolte A, Kunau WH, Erdmann R. Pex19p-dependent targeting of Pex17p, a peripheral component of the peroxisomal protein import machinery. J Biol Chem 2006;281:19417–25.

[181] Agrawal G, Joshi S, Subramani S. Cell-free sorting of peroxisomes. Cell 2012;149:397–409.

[182] Delille HK, Agricola B, Guimaraes SC, Borta H, Luers GH, Fransen M, et al. Pex11pbeta-mediated growth and division of mammalian peroxisomes follows a maturation pathway. J Cell Sci 2010;123:2750–62.

[183] Motley AM, Hettema EH. Yeast peroxisomes multiply by growth and division. J Cell Biol 2007;178:399–410.

[184] Nuttall JM, Motley AM, Hettema EH. Deficiency of the yeast Saccharomyces cerevisiae. Proc Natl Acad Sci U S A 2011;108:9113–8.

[185] Schuldiner M, Metz J, Schmid V, Denic V, Rakwalska M, Schmitt HD, et al. The GET complex mediates insertion of tail-anchored proteins into the ER membrane. Cell 2008;134:634–45.

[186] Bellu AR, Komori M, van der Klei IJ, Kiel JA, Veenhuis M. Peroxisome biogenesis and selective degradation converge at Pex14p. J Biol Chem 2001;276:44570–4.

[187] Farre JC, Manjithaya R, Mathewson RD, Subramani S. PpAtg30 tags peroxisomes for turnover by selective autophagy. Dev Cell 2008;14:365–76.

[188] Bhat P, Schleibs W, Schievelbusch T, Neuhaus A, David C, Kock K, et al. PEX14 is required for microtubule-based peroxisome motility in human cells. J Cell Sci 2011;124:1759–68.

[189] Titorenko VI, Chan H, Rachubinski RA. Fusion of small peroxisomal vesicles in vitro reconstructs an early step in the in vivo multistep peroxisome assembly pathway of Yarrowia lipolytica. J Cell Biol 2000;148:29–44.

[190] Nuttall JM, Motley AM, Hettema EH. Deficiency of the exportomer components Pex1, Pex6, and Pex15 causes enhanced pexophagy in Saccharomyces cerevisiae. Autophagy 2014;10:835–45.