The Cytosolic Domain of Pex22p Stimulates the Pex4p-Dependent Ubiquitination of the PTS1-Receptor

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Abstract

Peroxisomal biogenesis is an ubiquitin-dependent process because the receptors required for the import of peroxisomal matrix proteins are controlled via their ubiquitination status. A key step is the monoubiquitination of the import receptor Pex5p by the ubiquitin-conjugating enzyme (E2) Pex4p. This monoubiquitination is supposed to take place after Pex5p has released the cargo into the peroxisomal matrix and primes Pex5p for the extraction from the membrane by the mecano-enzymes Pex1p/Pex6p. These two AAA-type ATPases export Pex5p back to the cytosol for further rounds of matrix protein import. Recently, it has been reported that the soluble Pex4p requires the interaction to its peroxisomal membrane-anchor Pex22p to display full activity. Here we demonstrate that the soluble C-terminal domain of Pex22p harbours its biological activity and that this activity is independent from its function as membrane-anchor of Pex4p. We show that Pex4p can be functionally fused to the trans-membrane segment of the membrane protein Pex3p, which is not directly involved in Pex5p-ubiquitination and matrix protein import. However, this Pex3(N)-Pex4p chimera can only complement the double-deletion strain pex4Δ/pex22Δ and ensure optimal Pex5p-ubiquitination when the C-terminal part of Pex22p is additionally expressed in the cell. Thus, while the membrane-bound portion Pex22(N)p is not required when Pex4p is fused to Pex3(N)p, the soluble Pex22(C)p is essential for peroxisomal biogenesis and efficient monoubiquitination of the import receptor Pex5p by the E3-ligase Pex12p in vivo and in vitro. The results merge into a picture of an ubiquitin-conjugating complex at the peroxisomal membrane consisting of three domains: the ubiquitin-conjugating domain (Pex4p), a membrane-anchor domain (Pex22(N)p) and an enhancing domain (Pex22(C)p), with the membrane-anchor domain being mutually exchangeable, while the Ubc- and enhancer-domains are essential.

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Data Availability: The results merge into a picture of an ubiquitin-conjugating complex at the peroxisomal membrane consisting of three domains: the ubiquitin-conjugating domain (Pex4p), a membrane-anchor domain (Pex22(N)p) and an enhancing domain (Pex22(C)p), with the membrane-anchor domain being mutually exchangeable, while the Ubc- and enhancer-domains are essential.

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Introduction

Ubiquitination is one of the most prominent posttranslational modifications within the cell. It regulates the targeting and in many cases also the stability of proteins. The attachment of this modification requires a three-step enzyme cascade. First, the ubiquitin-activating enzyme (E1) forms an AMP-ubiquitin intermediate before it binds the activated ubiquitin moiety via a thioester-bond. Thereafter, ubiquitin is bound to the active-site cysteine of an ubiquitin-conjugating enzyme (E2). The final transfer of the ubiquitin moiety to the substrate usually requires the activity of an ubiquitin-protein ligase (E3). The E3 enzyme binds both the Ub-charged E2 enzyme as well as the target protein and either enables a direct transfer (RING-type E3) or requires a thioester-linked Ub-E3 intermediate (HECT-type and RBR-type E3s) in order to finally attach the ubiquitin moiety to the target amino acid of the substrate [1–3]. In recent years examples of additional, non-canonical ubiquitination factors have been described, comprising e.g. additional E3 enzymes (E4) like Ufd2p [4] or E2-activators like Cue1p [5,6].

Ubiquitination plays a central role in the biogenesis of peroxisomes. In fact, the first E2 enzyme required for the formation of an organelle, namely Pex4p (Pas2p/Ubc10p), has been found to be functionally linked to peroxisomal matrix protein import [7].

Peroxisomes are present in almost all eukaryotic cells and their dysfunction in humans is associated with a spectrum of severe peroxisomal disorders [9–10]. More than 50 different enzymes have been described to be compartmentalized in the peroxisomal matrix [11] which link this organelle to diverse biochemical reaction pathways and physiological functions. The most prominent task is the breakdown of fatty acids and the detoxification of hydrogen-peroxide [12,13].
The operativeness of this organelle is governed by dynamically operating import machineries for peroxisomal membrane and matrix proteins. All peroxisomal proteins are synthesized on free ribosomes in the cytosol and usually harbour a peroxisomal targeting sequence (PTS). Peroxisomal membrane proteins contain the mPTS, which comprises the binding site to the membrane protein-receptor/chaperone Pex15p as well as a membrane-association region [14,15]. Matrix proteins harbor either a PTS1 or a PTS2 sequence, which is recognized by the soluble PTS1-receptor Pex3p or the PTS2-receptor Pex7p, respectively [12,13,16]. Most peroxisomal matrix proteins are imported by the PTS1-receptor Pex5p. This cycling receptor binds its cargo proteins in the cytosol and ferries them to the peroxisome. The cargo is thought to be transported over the membrane via a transiently opened import pore [17] and finally released into the peroxisomal matrix. The PTS1-receptor is exported back to the cytosol in order to facilitate further rounds of matrix protein import [18]. This dislocation step is accomplished by the concerted action of peroxisomal membrane subcomplexes, collectively referred to as the peroxisomal exportomer [19]. This molecular machinery comprises enzymes required for the ubiquitination as well as the ATP-dependent extraction of the receptor from the membrane. A prerequisite for the export step is the monoubiquitination of the PTS1-import receptor Pex5p on a conserved cysteine [20–23]. Also the PTS2-receptor module has been shown to be regulated by monoubiquitination [24–26]. In yeast, the monoubiquitination of the PTS1-receptor Pex5p is catalyzed by the E2 enzyme Pex4p [22,23]. Pex4p cooperates with the peroxisomal RING-type E3 complex, from which Pex12p has been demonstrated to directly ubiquitinate Pex5p [27]. The monoubiquitinated Pex5p is recognized by the AAA-type ATPases Pex1p and Pex6p and dislocated to the cytosol [20–22,28,29]. In case this export reaction is affected, e.g. in a pex1Δ or pex4Δ strain, Pex5p becomes a substrate of a quality control pathway, resulting in its polyubiquitination on lysine residues and subsequent degradation by the 26S proteasome [22,23,30,31]. Interestingly, the deletion of the membrane protein Pex22p displays a similar phenotype [30,31], most likely due to its known role as the peroxisomal E2 unit. The Peroxisomal E2 Unit Consists of Three Functionally Distinct Parts

**Materials and Methods**

**Yeast strains and culture conditions**

*The Saccharomyces cerevisiae* strain UTL-7A (MATa, ura3–52, trp1, leu2–3/112) was used as wild-type strain for the generation of several isogenic deletion strains by the short flanking homology method as described [36]. The resulting deletion strains were pex4Δ [7], pex5A [37], pex22Δ and pex4Δ/pex22Δ [31]. Yeast complete (YPD) and minimal media (SD) as well as oleic acid medium (YNO) have been described previously [38].

**Recombinant proteins**

Genes coding for recombinant proteins were expressed in *E. coli* BL21 (DE3). The procedures for the preparation of GST-Pex12p(aa293–399) [39], GST-Pex4p [22], His-Pex3p [40], DsRed-PTS1 as well as DsRed-Ant1p [41] have been described previously. In order to generate the Pex3(1–45)-Pex4p-GFP construct, Pex4p was amplified from genomic DNA using the primer pair RE2417/RE2416. The obtained product was cloned with BamHI/SalI into the pAH37 vector (pUG35-Pex3(1–45)-GFP) [41]. The Pex3(1–45)-Pex4p construct was generated by the same procedure, with the only difference that Pex4p was amplified with its stop codon using the primer pair RE2418/RE2416. In order to generate the Pex22(1–35)-Pex4p-GFP construct, Pex4p was amplified from genomic DNA using the primer pair RE2417/RE2416. The obtained product was cloned with BamHI/SalI into the pAH44 vector (pUG35-Pex22(1–35)-GFP) [41]. The Pex3(1–45)-Pex4p construct was generated by the same procedure, with the only difference that Pex4p was amplified with its stop codon using the primer pair RE2418/RE2416. The Pex22p(36–180) construct was generated via amplification of the Pex22p-fragment from genomic DNA using the primer pair RE1222/RE3441. The isolated fragment was cloned via BamHI/XbaI restriction in a pRS415 vector, which contains a SacI/SpeI-inserted MET25-promoter.

In order to generate the His6-Pex22(36–180) construct, the fragment coding for the soluble domain of Pex22p was amplified from genomic DNA using the primers KU1164 and KU1165. The PCR fragment was cloned via NeoI and BamHI into pET9d (Novagen).

**Primer (Eurogentec) used in this study:**

- **KU1164** TAACGATGGAAAAAGTGACAGTGCAAAG
- **KU1165** TAAGGATCCCTTATTGCTAAATGCTAAT
- **RE1222** AAGTGCAGACCTAGATGCAACATTG CAAAGGAAGATA
- **RE2416** AAGGTACCATGCGAACATCTTTCCATTCT
- **RE2417** CCGTCGACATGTGTTGTGATCCGCCT
- **RE2418** AAGGTACCTAATGGTTGGTTGATCCGCCT
- **RE3441** AAGGATCC TTAATTGCTAAAGTGCTAATCAGGC

**In vitro ubiquitination**

Autoubiquitination assays of Pex4p were carried out as described previously [22] with the difference that 0.8 µg His-Pex22/Cp were added to the indicated reactions. *In vitro* ubiquitination of recombinant His-Pex5p contained 0.5 µg of.

**Plasmids, cloning strategies and recombinant proteins**

The generation of the constructs coding for GST-Pex12p(aa293–399) [39], GST-Pex4p [22], His-Pex3p [40], DsRed-PTS1 or as DsRed-Ant1p [41] have been described previously. Recombinant yeast E1 and ubiquitin were purchased from Sigma (München, Germany). Recombinant proteins

The functionality of peroxisomes was monitored by measuring the OD600 of the cells grown in YNO. Cells were first precultured for 16 h in 25 ml SD medium and transferred to 500 ml YNO with a starting OD600 of 0.1. OD600 was measured after 24 hrs and data were corrected for the measured value of the negative control. The corrected data are displayed in % complementation activity compared to wild-type cells.

**Complementation activity**

Autoubiquitination assays of Pex4p were carried out as described previously [22] with the difference that 0.8 µg His-Pex22/Cp were added to the indicated reactions. *In vitro* ubiquitination of recombinant His-Pex5p contained 0.5 µg of.

**Primer (Eurogentec) used in this study:**

- **KU1164** TAACGATGGAAAAAGTGACAGTGCAAAG
- **KU1165** TAAGGATCCCTTATTGCTAAATGCTAAT
- **RE1222** AAGTGCAGACCTAGATGCAACATTG CAAAGGAAGATA
- **RE2416** AAGGTACCATGCGAACATCTTTCCATTCT
- **RE2417** CCGTCGACATGTGTTGTGATCCGCCT
- **RE2418** AAGGTACCTAATGGTTGGTTGATCCGCCT
- **RE3441** AAGGATCC TTAATTGCTAAAGTGCTAATCAGGC

**Recombinant proteins**

Genes coding for recombinant proteins were expressed in *E. coli* BL21 (DE3). The procedures for the preparation of GST-Pex12p(aa293–399) and GST-Pex4p [22] as well as for His-Pex22p(aa36–180) and His-Pex5p [27] have been described previously. Recombinant yeast E1 and ubiquitin were purchased from Sigma (München, Germany).

**Materials and Methods**

**Yeast strains and culture conditions**

*The Saccharomyces cerevisiae* strain UTL-7A (MATa, ura3–52, trp1, leu2–3/112) was used as wild-type strain for the generation of several isogenic deletion strains by the short flanking homology method as described [36]. The resulting deletion strains were pex4Δ [7], pex5A [37], pex22Δ and pex4Δ/pex22Δ [31]. Yeast complete (YPD) and minimal media (SD) as well as oleic acid medium (YNO) have been described previously [38].

**Complementation activity**

The functionality of peroxisomes was monitored by measuring the OD600 of the cells grown in YNO. Cells were first precultured for 16 h in 25 ml SD medium and transferred to 500 ml YNO with a starting OD600 of 0.1. OD600 was measured after 24 hrs and data were corrected for the measured value of the negative control. The corrected data are displayed in % complementation activity compared to wild-type cells.
His-Pex5p, 0.5 μg of GST-RING fusion proteins, 0.1 μg of yeast E1, 0.8 μg E2-enzyme as indicated, 3 μg of ubiquitin-species as indicated and 10 μl buffer containing 2 mM ATP, 50 mM Tris-HCl pH 7.5, 2 mM MgCl₂, 2 mM ZnCl₂ and 0.1 mM dithiothreitol. After incubated at 30°C for 90 min with gentle shaking, the reaction was stopped by addition of SDS-sample buffer and heating at 95°C for 5 min.

**In vivo ubiquitination**

Preparation of polyubiquitinated Pex5p from oleate-induced yeast was based on a trichloroacetic acid precipitation and has been described previously [31]. Preparation of monoubiquitinated myc-Pex5p was performed by immunoprecipitation with myc-antibodies and Dynabeads anti-mouse-IgGs (Life Technologies) in the presence of 20 mM NEM.

**Microscopy**

Analysis of live cells for dsRed and GFP fluorescence was performed with a Zeiss Axioplan microscope and AxioVision 4.1 software (Zeiss, Jena, Germany). Before inspection, cells were grown for 2 days on solid minimal medium containing oleic acid as a sole carbon source [41].

**Mass Spectrometry**

The sample of the complex isolation was deluted by LDS-Puffer (‘‘NuPAGE LDS Sample Buffer 4×’’ von Invitrogen/Life Technologies, Carlsbad, USA), heated for 5 minutes on 95°C and separated by SDS-PAGE with a 4–12% NuPAGE Bis-Tris Gel (Invitrogen). After 18 minutes of separation at 50 V, the gel was stained with Imperial Protein Stain (Thermo Scientific) and the bands were cut, destained and in-gel digested with trypsin (Promega). Peptide extraction, sample preparation and liquid chromatography-mass spectrometry (LC-MS/MS) was performed as described [42].

**Immunodetection**

Polyclonal rabbit antibodies were raised against Pex5p [43], Pex22p (this study) and ubiquitin (Sigma, München, Germany). Monoclonal mouse antibodies were used for detection of GST (Sigma, München, Germany). Immuno-reactive complexes were visualized using the IRDye 800CW goat anti-rabbit IgG or IRDye 680 goat anti-mouse secondary antibody (Li-COR Bioscience, Bad Homburg, Germany) followed by detection using the “Infrared Imaging System” (Li-COR Bioscience, Bad Homburg, Germany).

**Results**

The chimeric Pex3(N)-Pex4p fusion protein is targeted to peroxisomes and attracts the soluble Pex22(C)p

The soluble ubiquitin-conjugating enzyme Pex4p associates with peroxisomes via its interaction to the C-terminal part of the peroxisomal membrane protein Pex22p [32]. The binding to Pex22p enhances the activity of Pex4p [34,35]. We wanted to analyze the functional impact of this interaction on the assembly of the Pex4p/Pex22p complex as well as on the Pex12p-dependent ubiquitination of the PTS1-receptor Pex3p.

For this purpose, we investigated whether the function of Pex22p as Pex4p-anchor can be uncoupled from its function as Pex4p-enhancer. To this end, we generated a chimeric protein, consisting of Pex4p and the N-terminal amino acids 1–45 of Pex3p (Pex3p(N)-Pex4p) and GFP for detection (Fig. 1a). In *S. cerevisiae*, Pex3p takes part in the process of membrane protein insertion, *de novo* formation of peroxisomes, pexophagy and inheritance, but is not directly involved in matrix protein import [44–48]. The N-terminal 45 amino acids of Pex3p contain its mPTS and membrane anchor sequence [41,49] (Fig. 1a). Thus fusion of this region to Pex4p-GFP should direct the protein to peroxisomes and anchor the fusion protein to the peroxisomal membrane.

Accordingly, we analyzed the peroxisomal targeting of the Pex3(N)-Pex4p-GFP fusion protein to peroxisomes in wild-type, pex4Δ and pex3Δpex22A strains (Fig. 1b). For this, the GFP-fused Pex4p-chimera was assayed for co-localization with the DsRed-tagged peroxisomal adenine nucleotide transporter Ant1p. The fluorescence microscopic analysis revealed that Pex3(N)-Pex4p-GFP was targeted to peroxisomes in all tested strains, which indicated that the Pex3(N)-anchored Pex4p-GFP form is targeted to the peroxisomal membrane in cells that completely lack Pex22p (Fig. 1b). Thus, under these circumstances Pex22p is dispensable for peroxisomal targeting of Pex4p, which now is membrane anchored by the Pex3p(1–45).

Furthermore, we tested, if the detached soluble carboxyl-terminal part of Pex22(aa36–180)p (=Pex22(C)p) can bind to the peroxisomal Pex4p-chimera when it is additionally expressed within the cell. The Pex22(C)p-fragment mainly contains the binding region required for the interaction with Pex3p [34], but no membrane domain (Fig. 1a). To test for the binding capabilities, we isolated Pex3(N)-Pex4p-GFP via immunoprecipitation with GFP-antibodies from *pex4Δpex22A* cells that also expressed Pex22(C)p and analyzed the isolated complex by mass-spectrometry. Peptides derived from Pex22(C)p were detected in samples derived from cells that contained Pex3(N)-Pex4p-GFP, while no Pex22(C)p was detected in samples from strains that had not been transformed with the Pex4-GFP-chimera (data not shown). Therefore, we can conclude that Pex22(C)p binds via Pex3(N)-Pex4p-GFP to peroxisomes.

Taken together these data demonstrate that Pex4p can target to peroxisomes independently of Pex22p when it is fused to the membrane-anchor Pex3(N)p and that Pex22(C)p can associate with this assembly.

The function of Pex22(N)p is dispensable for matrix protein import, while Pex22(C)p is essential for biological activity

In order to determine the functionality of the Pex4p-chimera complex in biogenesis of peroxisomes, we tested the complementation activity of Pex3(N)-Pex4p in combination with Pex22(C)p in *pex4Δ* and *pex4Δpex22A* strains in comparison to wild-type cells (Fig. 2a). The cells were incubated in growth media that contained oleate as sole carbon source. Under these conditions peroxisomes become essential for the viability of the cells because they are the only site where fatty acids can be degraded via beta-oxidation [50]. Growth of the wild-type is set as 100% (Fig. 2a). As *pex4Δ* and *pex4Δpex22A* strains do not grow on oleate-medium, the *pex22D* strains in comparison to wild-type cells.

The Peroxisomal E2 Unit Consists of Three Functionally Distinct Parts
membrane insertion. When this construct is expressed in \textit{pex}4\textsuperscript{Δ} cells together with the chimeric \textit{pex}3\textsuperscript{N}-\textit{pex}4\textsuperscript{p}, the cells regained the ability to grow on oleate medium (Fig. 2a). This restoration does not reach the wild-type level, which is likely caused by the lower efficiency of the peroxisomal targeting of \textit{pex}22\textsuperscript{C}\textsuperscript{p}. In a control experiment, we verified that the cellular level of overexpressed \textit{pex}22\textsuperscript{C}\textsuperscript{p} is the same in all tested strains (Fig. 2c). Therefore, complementation can only be achieved when the peroxisomal E2-complex can form (\textit{e.g.} in \textit{pex}1\textsuperscript{Δ}\textit{pex}22\textsuperscript{Δ}+\textit{pex}3\textsuperscript{N}–\textit{pex}4\textsuperscript{p}+\textit{pex}22\textsuperscript{C}\textsuperscript{p}) and is not possible when \textit{pex}4\textsuperscript{p} is mislocalized, even if \textit{pex}22\textsuperscript{C}\textsuperscript{p} is present (\textit{e.g.} in \textit{pex}22\textsuperscript{Δ}+\textit{pex}22\textsuperscript{C}\textsuperscript{p}).

Importantly, these data demonstrate that membrane-anchored \textit{pex}4\textsuperscript{p} can perform its function during peroxisomal biogenesis independently of full-length \textit{pex}22\textsuperscript{C}\textsuperscript{p} as long as it can associate with \textit{pex}22\textsuperscript{C}\textsuperscript{p}, even when its membrane-spanning part \textit{pex}22\textsuperscript{N}\textsuperscript{p} is completely absent.

In order to analyze whether the observed complementation of the growth phenotype correlates with restoration of peroxisomal biogenesis and peroxisomal protein import, we monitored transformed wild-type and mutant cells for the intracellular localization of the model cargo DsRed-PTS1 by direct fluorescence microscopy (Fig. 2b). In wild-type cells, the punctate pattern indicates a complete import of the PTS1-containing fluorophore into peroxisomes (Fig. 2b), while the deletion of peroxin genes prevents the import and results in a cytosolic mislocalization as seen here for non-complemented \textit{pex}4\textsuperscript{Δ} cells. The punctate pattern observed for DsRed-PTS1 in the \textit{pex}4\textsuperscript{Δ} strain transformed with \textit{pex}3\textsuperscript{N}\textsuperscript{-pex}4\textsuperscript{p}+GFP demonstrates that these cells display a functional matrix protein import and thus restoration of the peroxisome biogenesis defect (Fig. 2b). However, \textit{pex}3\textsuperscript{N}\textsuperscript{-pex}4\textsuperscript{p}+GFP was not able to rescue the DsRed-PTS1 import in the \textit{pex}4\textsuperscript{Δ} background. The additional presence of the \textit{pex}22\textsuperscript{C}\textsuperscript{p} construct displayed a heterogenous phenotype. Even though a significant amount of DsRed-PTS1 was still mislocalized to the cytosol, the peroxisomes were nevertheless found to be import competent (Fig. 2b). These data correlate with our finding that the complementation of \textit{pex}3\textsuperscript{N}\textsuperscript{-pex}4\textsuperscript{p} combined with the \textit{pex}22\textsuperscript{C}\textsuperscript{p} fragment in the \textit{pex}1\textsuperscript{Δ}\textit{pex}22\textsuperscript{Δ} background is only partial (Fig. 2a).

To conclude, the data corroborate the complementation assays by growth analysis in that they demonstrate that the chimeric \textit{pex}3\textsuperscript{N}\textsuperscript{-pex}4\textsuperscript{p} is functional in the presence of \textit{pex}22\textsuperscript{p} and that under these conditions the \textit{pex}22\textsuperscript{C}\textsuperscript{p} fragment can account for the requirement for \textit{pex}22\textsuperscript{p}. These data clearly show that \textit{pex}22\textsuperscript{p} is not merely a docking protein for the ubiquitin conjugating \textit{pex}4\textsuperscript{p} but that its C-terminal domain maintains an important function in peroxisome biogenesis.

\textbf{Pex22\textsuperscript{C}\textsuperscript{p} stimulates the autoubiquitination of \textit{pex}4\textsuperscript{p} as well as the \textit{pex}12\textsuperscript{p}-dependent ubiquitination of the \textit{PTS1}-receptor \textit{in vitro}}

As \textit{pex}22\textsuperscript{p} interacts with \textit{pex}4\textsuperscript{p}, we investigated whether the detected function of \textit{pex}22\textsuperscript{C}\textsuperscript{p} in matrix protein import is connected to \textit{pex}4\textsuperscript{p}-mediated ubiquitination events. First, we tested the effect of \textit{pex}22\textsuperscript{C}\textsuperscript{p} on the \textit{in vitro} autoubiquitination of \textit{pex}4\textsuperscript{p} (Fig. 3a). For this purpose, we purified heterologously expressed GST-tagged \textit{pex}4\textsuperscript{p}, His-tagged \textit{pex}22\textsuperscript{C}\textsuperscript{p} as well as His-tagged \textit{pex}3\textsuperscript{p} and the RING-domain of \textit{pex}12\textsuperscript{p} from \textit{E. coli} and analyzed them in an \textit{in vitro} ubiquitination assay in different combinations. The generated protein samples were subjected to immunoblot analysis with GST- and ubiquitin-antibodies. The data set shows that the \textit{in vitro} autoubiquitination reaction of GST-\textit{pex}4\textsuperscript{p} depends on the presence of \textit{E. coli} and ubiquitin (Fig. 3a). Furthermore, His-\textit{pex}22\textsuperscript{C}\textsuperscript{p} does not display any ubiquitination activity on its own, but its addition to the reaction sample containing GST-\textit{pex}4\textsuperscript{p}, \textit{E. coli} and ubiquitin significantly enhances...
the autoubiquitination of the peroxisomal E2 enzyme. Thus, even though His-Pex22(C)p does not display intrinsic ubiquitination activity, it is capable to stimulate the autoubiquitination activity of GST-Pex4p (Fig. 3a).

Next we wanted to analyze whether Pex22(C)p also has an effect on the ubiquitination of the PTS1-receptor Pex5p, which is a well-known target protein for Pex4p-dependent ubiquitination [22,23]. The in vitro ubiquitination assays demonstrated that the ubiquitination of His-Pex5p depends on the presence of E1, GST-Pex4p, ubiquitin as well as the RING-domain of the E3-enzyme Pex12p [27] (Fig. 3b). Under these conditions a small portion of Pex5p is ubiquitinated (Fig. 3b, last lane). Moreover, the data reveal that His-Pex22(C)p does not ubiquitinate His-Pex5p itself, as no modification is visible when GST-Pex4p is omitted from the His-Pex22(C)p-containing sample (Fig. 3b, lane 6). However, in the presence of GST-Pex4p and His-Pex22(C)p, a clear increase in the intensity of the band representing ubiquitinated Pex5p is observed in the presence of Pex4p and His-Pex22(C)p (Fig. 3b, lane 7). The result shows that His-Pex22(C)p stimulates the GST-Pex4p-mediated in vitro ubiquitination of Pex5p (Fig. 3b). We conclude that Pex22(C)p can not only stimulate the autoubiquitination of Pex4p, but also that this enhancing effect can be directly transferred to the Pex12p-dependent ubiquitination of the PTS1-receptor Pex5p.

**Pex22(C)p reduces polyubiquitination and stimulates monoubiquitination of Pex5p in vivo**

We wanted to analyze whether the detected stimulating effect of Pex22(C)p on the E2 activity of Pex4p and the in vitro ubiquitination of Pex5p are of relevance in vivo. Our data on the specific role of Pex22(C)p in peroxisomal biogenesis and PTS1-protein import so far showed that Pex22(C)p can partially
complement the \textit{pex4\textDelta} deletion strain when Pex4p is anchored by another protein (Fig. 2). It is known that the inhibition of the monoubiquitination of Pex5p caused by the deletion of \textit{PEX4} and/or \textit{PEX22}, or, alternatively, mutation of the AAA-peroxins, induces polyubiquitination and proteasomal degradation of Pex5p [22,23,30,31,51]. Because of this, the occurrence of polyubiquitinated Pex5p can serve as a biochemical \textit{in vivo} tool to monitor a potential defect in Pex5p export. Here we show that the polyubiquitination of Pex5p seen in \textit{pex4\textDelta} cells (Fig. 4a, lane 2) can be nearly completely abolished when the mutant cells express Pex3(N)-Pex4p (Fig. 4a, lane 3). When the Pex4p-chimera is expressed in the \textit{pex4\Delta pex22\Delta} strain, the amount of polyubiquitinated Pex5p is only very weakly reduced in comparison to \textit{pex4\Delta pex22\Delta} cells (Fig. 4a, lane 4 and lane 5). However, when in addition to Pex3(N)-Pex4p the Pex22(C)p-fragment is expressed in the \textit{pex4\Delta pex22\Delta} strain, the polyubiquitination of Pex5p is significantly reduced. This indicates that the presence of Pex22(C)p can indeed partially suppress the export defect of Pex5p.

The physiological more important ubiquitination of Pex5p is its monoubiquitination, which represents the canonical signal for export of Pex5p [20–22]. In yeast, this modification is catalyzed by Pex4p [22,23] and we wanted to investigate whether Pex22(C)p also has an effect on this process. To this end, we isolated Pex5p in the presence of NEM in order to inhibit deubiquitination [22,51]. Moreover we used myc-Pex5p construct, because the N-terminal presence of this tag is known to block polyubiquitination of Pex5p, which would normally occur in \textit{pex4\textDelta} or \textit{pex22\textDelta} strains [22]. This enabled us to analyze the effect of the Pex4p- and Pex22p-constructs on the monoubiquitination of the PTS1-receptor. We found that Pex3(N)-Pex4p can monoubiquitinate myc-Pex5p in the \textit{pex4\Delta} strain with an efficiency that is comparable to the wild-type situation (Fig. 4b). The level of monoubiquitinated myc-Pex5p is reduced significantly when the Pex4p-chimera is expressed in the \textit{pex4\Delta pex22\Delta} strain. This effect is due to the complete absence of Pex22p, but also indicates that the basic E2-activity of Pex4p does not essentially depend on the presence of Pex22p "per se". However, when Pex22(C)p was expressed in addition, the level of monoubiquitinated Pex5p increased significantly (Fig. 4b). This demonstrates that Pex22(C)p is required for full activation of Pex4p and efficient monoubiquitination of the PTS1-receptor.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Pex22(C)p enhances autoubiquitination of Pex4p and Pex4p/Pex12p-catalyzed ubiquitination of the PTS1-receptor \textit{in vitro}. (a) Autoubiquitination of GST-Pex4p. The effect of Pex22(C)p on the autoubiquitination of GST-Pex4p was examined \textit{in vitro} using heterologously expressed and purified proteins as indicated. Autoubiquitination of GST-Pex4p depends on the presence of E1 and ubiquitin (Ub). As shown by immunoblot analysis with antibodies against GST and Ub, the autoubiquitination of GST-Pex4p is significantly enhanced in the presence of His-Pex22(C)p. (b) Ubiquitination of the PTS1-receptor Pex5p. Recombinant proteins were used for \textit{in vitro} ubiquitination of Pex5p, which depends on the presence of E1, GST-Pex4p, Ub as well as the RING-domain of the E3 enzyme Pex12p and was significantly enhanced in presence of His-Pex22(C)p.}
doi:10.1371/journal.pone.0105894.g003
\end{figure}
The stimulating effect of Pex22(C)p on Pex5p monoubiquitination and matrix protein import cannot be enforced or replaced by Pex22(N)p.

In the following we tested if also the N-terminal region of Pex22p, thus Pex22(aa1–35)p, in the following designated Pex22(N)p, has an enhancer effect on Pex5p-monoubiquitination. We generated a fusion of Pex22(aa1–35) and full length Pex4p (Pex22(aa1–35)-Pex4p) (Fig. 5a). In a first step, we verified that the chimeric construct is targeted to peroxisomes by fluorescence microscopy. The peroxisomal localization of Pex22(1–35)-Pex4p-GFP is indicated by its congruent fluorescence pattern with the peroxisomal marker DsRed-Ant1p (Fig. 5b). Next, we could show by immunosolubilisation of Pex4p-complexes and their mass spectrometric analysis that Pex22(N)-Pex4p interacts with Pex22(C)p (data not shown). Furthermore, Pex22(1–35)-Pex4p can complement and thus restore growth ability on oleate medium of the pex4Δ strain but not of the pex3Δpex22Δ strain (Fig. 5c). However, a partial complementation of the pex4Δpex22Δ strain was achieved by combined expression of Pex22(1–35)-Pex4p and the Pex22(C)p fragment (Fig. 5c). These data demonstrate that the presence of the Pex22(1–35)p alone is not enough to compensate the loss of the full length Pex22p, even if it is genetically fused to Pex4p. Only the additional presence of the detached Pex22(C)p enables a partial restoration of peroxisome function.

We next investigated whether it makes a difference for the enhancing effect of Pex22(C)p when Pex4p is fused to the membrane anchor of either Pex3p or Pex22p. The direct comparison of pex4Δpex22Δ strains expressing Pex22(C)p and either Pex3(N)-Pex4p or Pex22(N)-Pex4p revealed that both display a similar complementation activity (Fig. 5c). This result demonstrates that Pex22(N)p is not able to replace or enforce the function of Pex22(C)p. This also accounts for the enhancing effect of the C-terminal domain of Pex22p for Pex5p-monoubiquitination as shown in Fig. 5D. The pex4Δpex22Δ strain containing Pex22(N)-Pex4p alone displays only a residual level of Pex3p-monoubiquitination, while modification of Pex5p is enhanced, when Pex22(C)p is additionally present.

Taken together, these results indicate that Pex22(N)p, just like Pex3(N)p, functions as the membrane anchor, but is not directly involved in the ubiquitin transfer reaction of Pex4p. The enhancer effect is facilitated only by Pex22(C)p.

**Discussion**

In this study, we have analysed the functional interplay of the ubiquitin-conjugating enzyme complex Pex4p/Pex22p and its physiological substrate, the PTS1-receptor Pex5p. We demonstrate that the soluble C-terminal Pex4p-binding domain of Pex22p stimulates the E2-activity of Pex4p both in vivo and in vitro and thereby enhances the monoubiquitination of Pex5p mediated by the RING-type E3 enzyme Pex12p and supports proper matrix protein import.

The membrane protein Pex22p has been described previously as membrane anchor of Pex4p [32,33] and it has been shown that the interaction of Pex4p with Pex22p is important for full activity of the E2 enzyme [34,35]. The crystal structure of Pex4p bound to the soluble part of Pex22p revealed a novel binding interface and block of this interaction resulted in a reduced autoubiquitination of Pex4p in vitro and decreased PTS1-receptor ubiquitination in vivo and impaired growth on oleate medium [34]. In our study, we analysed the molecular details of the enhancing effect of Pex22p on Pex4p and show that Pex22p stimulates the in vitro ubiquitination of its physiological substrate, the PTS1-receptor Pex5p. Thus, the Pex22-induced stimulation of Pex4p is directly transferred to an increased ubiquitination of Pex5p by the RING-ligase Pex12p. This stimulating effect was ascribed to the C-terminal domain of Pex22p. The only other E2 complex that seems to be assembled and regulated in a similar manner is the E2 enzyme Ubc7p and its binding partner Cue1p. The membrane-bound protein Cue1p was first described as anchor and stabilizer of Ubc7p at the endoplasmic reticulum [52]. Moreover, it has been demonstrated that it also functions as activator of the E2 activity of Ubc7p [5,6]. The truncated version of Cue1p, which contains the binding region to Ubc7p, can enhance the E2 activity by stimulating both the charging of Ubc7p with ubiquitin activated by E1 as well as the transfer of the ubiquitin to the substrate mediated by the interacting E3 ligase Hrd1p [53].

The resemblance of the two ubiquitination cascades is of special interest because there are certain similarities between the ERAD (endoplasmic reticulum associated degradation)-machinery and the peroxisomal exportomer [54–56]. However, the available crystal structures indicate that the Pex22p-Pex4p interaction does not display the structural features that resemble the situation of the Cue1p-Ubc7p complex [53]. Therefore, it will be important for further studies to identify a functional interconnection of the Pex22(C)p/Pex4p unit and its assembly with Pex12(RING)p or...
The Peroxisomal E2 Unit Consists of Three Functionally Distinct Parts

with the entire trimeric peroxisomal RING-complex [Pex12p, Pex10p, Pex2p] [57].

The dissection of the functional regions of Pex22p revealed that there molecular functions in peroxisome biogenesis are divided among the two domains of the protein. It turned out that the stimulating effect of Pex22p is limited to the C-terminal domain of the protein without contribution of the N-terminal region. The N-terminal region instead comprises the peroxisomal targeting signal and transmembrane domain, required for peroxisomal targeting and membrane-anchoring of the E2-complex. We have used the aa 1–35 fragment of Pex22(N)p for the fusion to Pex4p, while a previous study used Pex22(aa1–54)-Pex4p to complement the pex4Δ strain [34]. The complementation seems to be more efficient with the longer construct, possibly indicating a function as hinge-region for the aa 36–54. However, as our Pex22(1–35)-Pex4p construct was introduced in wild-type, pex4Δ and pex4Δpex22Δ cells as indicated and co-localization of the chimeric protein with the peroxisomal membrane marker DsRed-Ant1p was analyzed under by fluorescence microscopy. (c) The functionality of indicated constructs was tested by analysis of their capability to complement the growth defect of depicted strains. To this end, the Pex3(N)-Pex4p and/or Pex22(C)p or Pex22p encoding information was expressed in the indicated constructs was tested by analysis of their capability to complement the growth defect of depicted strains. To this end, the Pex3(N)-Pex4p and/or Pex22(C)p or Pex22p encoding information was expressed in the wild-type (+/+) or pex22Δ deletion strains, as indicated. The optical density (OD600) of cells grown in oleate medium was monitored (n = 3 experiments) and the results are presented in % in comparison to the wild-type (+/+−−− standard error of the mean). While Pex22(N)-Pex4p complements the growth defect of the pex22Δ strain, it is only capable to regain functionality in the pex4Δpex22Δ background, when Pex22(C)p is present. (d) The Pex22(N)-Pex4p fusion protein is capable to monoubiquitinate Pex5p in vivo. In the absence of full-length Pex22p, the level of monoubiquitinated Pex5p is significantly reduced, which is compensated in the presence of Pex22(C)p.

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Figure 5. The stimulating effect of Pex22(C)p on Pex5p ubiquitination and peroxisomal function cannot be enforced or replaced by Pex22(N)p. (a) Schematic representation of Pex22(N)-Pex4p. The Pex22(aa1–35)-fragment contains the transmembrane domain (TMD) in red and the intraperoxisomal part in orange fused to full-length Pex4p (blue) as well as green fluorescent protein (green). (b) Pex22(N)-Pex4p-GFP is targeted to peroxisomes. The Pex22(N)-Pex4p-GFP construct was introduced in wild-type, pex4Δ and pex3(N)-pex22Δ cells as indicated. (c) The functionality of indicated constructs was tested by analysis of their capability to complement the growth defect of depicted strains. To this end, the Pex3(N)-Pex4p and/or Pex22(C)p or Pex22p encoding information was expressed in the pex3(N)-pex22Δ, pex4Δ or pex22Δ deletion strains, as indicated. The optical density (OD600) of cells grown in oleate medium was monitored (n = 3 experiments) and the results are presented in % in comparison to the wild-type (+/+−−− standard error of the mean). While Pex22(N)-Pex4p complements the growth defect of the pex22Δ strain, but these cells cannot complement the pex4Δ or pex22Δ deletion strains, as indicated. The optical density (OD600) of cells grown in oleate medium was monitored (n = 3 experiments) and the results are presented in % in comparison to the wild-type (+/+−−− standard error of the mean). While Pex22(N)-Pex4p complements the growth defect of the pex22Δ strain, it is only capable to regain functionality in the pex4Δpex22Δ background, when Pex22(C)p is present. (d) The Pex22(N)-Pex4p fusion protein is capable to monoubiquitinate Pex5p in vivo. In the absence of full-length Pex22p, the level of monoubiquitinated Pex5p is significantly reduced, which is compensated in the presence of Pex22(C)p.

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membrane complexes seem to be restricted [19,56,58]. This connection has been demonstrated for A. thaliana, where a strain expressing an functionally impaired PEX6 mutant was only able to facilitate matrix protein import, when this late-acting mutant was combined with an early-acting-mutant, which was a low-expressed allele of the docking proteins PEX13 [59]. In line with this, a recently published stochastic computational model of the dynamics of Pex3p ubiquitination and matrix protein translocation in mammals suggests that both processes are cooperatively coupled [60]. Therefore, the residual monoubiquitination detected in our Pex4p-chimeras expressing pex14Δpex22A strains may most likely drastically reduce the export-rate of the receptors and block the import of new cargo-charged receptors.

This study focuses on the function of Pex22p/Cp by utilizing the PTS1-import pathway as experimental system. It will be of interest to elucidate if the described enhancer effect of Pex22(C)p has a similar impact on the ubiquitination of the PTS2-co-receptors and PTS2-protein import.

To conclude, our data demonstrate that the ubiquitination-activity of Pex4p and therefore the Pex12p-dependent mono-ubiquitination of the PTS1-receptor are stimulated by the C-terminal domain of Pex22p and that this enhancer effect is required to enable the monoubiquitination above a physiological threshold and therefore matrix protein import.

Further experiments will address the functional integration of this Pex22/Cp-mediated enhancer-effect on the translocation of folded proteins over the peroxisomal membrane.

Author Contributions
Conceived and designed the experiments: HWP FE AS RE. Performed the experiments: FE AS RB T. Mastalski LK. Analyzed the data: HWP FE RE WG AS. Contributed reagents/materials/analysis tools: HWP RE. T. Muller KM HEM. Contributed to the writing of the manuscript: HWP RE FE AS.

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