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Stress exposure results in increased peroxisomal levels of yeast Pnc1 and Gpd1, which are imported via a piggy-backing mechanism

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1. Introduction

Saccharomyces cerevisiae Gpd1 (Glycerol-3-phosphate dehydrogenase 1) and Pnc1 (nicotinamidase) are peroxisomal enzymes that are induced upon exposure of cells to various stress conditions [1-4]. Gpd1 is a nicotinamide adenine dinucleotide-hydrogen (NADH)-dependent enzyme that converts dihydroxyacetone phosphate (DHAP) into glycerol-3-phosphate, which subsequently can be converted into glycerol [5]. At osmotic stress conditions Gpd1 activity increases, which leads to enhanced cellular glycerol levels and osmotic stress resistance [6]. At normal conditions, Gpd1 contributes to reducing cellular DHAP levels, which prevents the spontaneous conversion of DHAP into methylglyoxal, a highly toxic compound that damages proteins and contributes to aging [7,8]. In addition, Gpd1 may play a role in controlling the redox balance by reoxidation of NADH to NAD+. Pnc1 functions in the NAD + salvage pathway and catalyses deamination of nicotinamide to nicotinic acid [9]. Overexpression of Pnc1 has been reported to positively affect the replicative lifespan of yeast [1]. It has been suggested that this is the result of Pnc1-mediated nicotinamide depletion in the nucleus, which activates the histone deacetylase Sir2, resulting in silencing of pro-ageing genes [1,10]. Pnc1 levels are elevated under conditions known to extend replicative lifespan, including osmotic and heat stress, conditions that also enhance Gpd1 levels [1]. Interestingly, examination of transcriptional profiles revealed that PNC1 expression is strongly correlated with that of GPD1 [11].

Although the above functions of Gpd1 and Pnc1 are expected to take place in the cytosol and nucleus respectively, at normal growth conditions both enzymes are predominantly localised to peroxisomes in conjunction with a minor portion to the cytosol [1,11]. When Gpd1 and Pnc1 levels were enhanced upon exposure of cells to osmotic stress, a much higher portion of these enzymes became cytosolic concomitant with a relative decrease in the peroxisomal protein levels [11]. However, it has also been reported that Pnc1 mainly remains peroxisomal in cells exposed to stress [1].

The peroxisomal localisation of Gpd1 depends on its N-terminal peroxisomal targeting signal 2 (PTS2) and the PTS2-receptor Pex7 [11]. It has been suggested that import of Gpd1 into peroxisomes is stimulated by the phosphorylation of two serine residues adjacent to the PTS2 sequence. The enhanced cytosolic Gpd1 level in cells exposed to stress was proposed to be due to decreased phosphorylation of these residues [11,12].

Sequence analysis did not reveal any predicted peroxisomal targeting signals in Pnc1. However, sorting of Pnc1 requires the PTS2 import machinery, because the protein mislocallises to the cytosol in Δpex7 cells [1]. Computational analysis revealed that Pnc1 may form a complex with Gpd1 [13]. If so, Pnc1 may import into peroxisomes in complex with Gpd1 by so called piggy-back import.
Here we show that Pnc1 and Gpd1 proteins indeed physically interact in vivo and that this allows piggy-back import of Pnc1 with Gpd1. Furthermore, quantitative fluorescence microscopy analysis revealed that exposure of the cells to osmotic stress the peroxisomal as well as the cytosolic levels of both proteins increased. Our data indicate that the appearance of cytosolic Gpd1 and Pnc1 most likely is caused by a general decrease in the rate of PTS2–matrix protein import in yeast cells exposed to stress, probably because this pathway is easily saturated.

2. Materials and methods

2.1. Strains and growth conditions

The *S. cerevisiae* strains used in this study are listed in Supplementary Table 1. Yeast cells were grown at 30 °C on mineral medium (MM; van Dijken et al. [14]) containing 0.25% ammonium sulphate and 2% glucose. MM was supplemented with the required amino acids or uracil to a final concentration of 20 μg ml−1 (histidine and methionine) or 30 μg ml−1 (leucine, lysine, and uracil). YPD medium (1% yeast extract, 1% peptone, 1% glucose) supplemented with 2% agar was used for growth on plates. *Escherichia coli* DH5α was used for cloning purposes and was cultured at 37 °C on LB medium supplemented with the appropriate antibiotics.

2.2. Construction of yeast strains

Plasmids and primers used in this study are listed in Supplementary Tables 2 and 3, respectively.

2.2.1. Construction of Apnc1 and Δgpd1 strains

*GPD1* was deleted in a strain producing Pnc1–GFP obtained from GFP fusion collection [15], by replacing the ORF with the *KanMX4* gene from pUG6 [16] using primers PGD1F and PGD1R. Pnc1–GFP producing cells were transformed with the PCR product, colonies were selected on YPD plates containing 200 μg ml−1 G418, and positive clones were checked by colony PCR. Correct integration was confirmed by Southern blotting. PNC1 deletion in Gpd1–GFP producing cells obtained from GFP fusion collection [15] was obtained by replacement of the ORF with the *KanMX4* gene from pUG6 using primers PNC1F and PNC1R. Cells were transformed with the PCR product and transformants were selected on YPD plates containing 200 μg ml−1 G418, positive clones were checked by colony PCR and Southern blot.

2.2.2. Strains producing NΔGpd1 and NΔgpd1–mCherry

To construct a strain lacking 17 N-terminal amino acid residues (+4 to +54 bps) of Gpd1 after the start codon, a plasmid pAG25-N_del_Gpd1 was cloned and transformed into Δgpd1 strain obtained from Euroscarf. To this end, the *GPD1* coding region +55 (+1 is A of the start codon) from the start codon (without PTS2 sequence) and the *GPD1* promoter region −540 from the start codon were amplified from *S. cerevisiae* genomic DNA using the primers pairs GPD1.OL-1/GPD1.OL-2.1 and GPD1.OL-1/GPD1.OL-2.2 respectively. The two PCR products of 1457 bps and 472 bps were joined together by overlap PCR and the combined fragment was further amplified using primer pair GPD1.OL-1/GPD1.OL-2.1, which resulted into a fragment of 1984 bps. This fragment was digested with HindIII and cloned in plasmid pAG25 [17] resulting in plasmid pAG25-N_del_Gpd1. The insert was sequenced to exclude the presence of errors. The plasmid was linearized with SbfI and transformed to *S. cerevisiae* Δgpd1 cells. Transformants were selected on YPD plates containing 100 μg ml−1 nourseothricin (WERNER BioAgents). Correct integration in the genomic DNA was checked by colony PCR and Southern blotting. The resulting strain that produces NΔgpd1 but not the endogenous Gpd1 was designated as nΔgpd1.

2.2.3. Construction of a strain producing NΔGpd1–mCherry and Pnc1–SKL

To obtain a plasmid containing the zeocin resistance gene and a gene encoding Pnc1–SKL, PNC1 genomic DNA starting from −700 was amplified using primers P.SKL_F_BamHI and P.SKL_R_HindIII. The PCR product was digested with BamHI/HindIII and cloned into pSP34 resulting in pPNCI–SKL. To add the PTS1 tripeptide -SKL to endogenous Pnc1, the region encoding the 3′-end of the Pnc1 gene along with the -SKL (PTS1) coding region and the zeocin resistance gene were amplified from pPNCI–SKL–1 using primers PNC1.1 and PNC1.GFP.SKL–2.2. The PCR product was used to transform nΔgpd1 cells and clones were selected on YPD/zeocin plates. mCherry was introduced at the C-terminus of NΔGpd1 in the resulting strain as follows: the mCherry–hph (Hygromycin R) fragment was amplified from pARM001 (see below) using primer pair GPD1_MC_F/GPD1_MC_Rev. The PCR product was used to transform nΔgpd1–SKL. Positive clones were checked by colony PCR. The resulting strain produces Pnc1–SKL and NΔGpd1–mCherry but not endogenous Gpd1.

2.2.4. Construction of a strain producing Pot1–GFP under control of the *GPD1* promoter

The GPD1 promoter starting from −700 bps of the start codon and the POT1 open reading frame were amplified from *S. cerevisiae* genomic DNA using primer pairs P–GPD1.PciI/GPD1.OL-Rev. and POT1.OL-Fw/POT1–BglII that resulted in PCR products of 731 bps and 1284 bps, respectively. Both DNA fragments were joined together by overlap PCR and combined fragment was further amplified using primer pair GPD1.PciI/POT1–BglII, which resulted in a DNA fragment of 1973 bps. The combined fragment was digested by restriction enzymes NciI/BglII and cloned into the pHP2–mGFP fusinatior plasmid [18], which resulted in pPGPD1–POT1–GFP. The plasmid was linearised by SfiI and transformed into *S. cerevisiae* strain producing DsRed1–SKL. Transformants were selected on YPD/zeocin plates and checked by colony PCR.

2.2.5. Construction of other plasmids

2.2.5.1. Construction of pSL33. To construct a plasmid producing DsRed–SKL under control of the MET25 promoter, the *P META25–DsRed–SKL–tyc1* fragment was amplified from pUG34–DsRed–SKL [25] using primer pair DsRed–1/DsRed–2. The obtained PCR product was digested with KpnI/XbaI and cloned into pBSII KS + resulting in pSL32. The nourseothricin resistance gene was amplified from pAG25 using primer pair Nat1.1/Nat1.2 and after digestion with SacII/KpnI the fragment was cloned into pSL32 that resulted in pSL33.

2.2.5.2. Construction of *pTDA1–GFP–SKL*. The promoter of the *TDH3* gene was amplified from *S. cerevisiae* genomic DNA by using primer pair TDH3–Nat1.1/TDH3–BamHI, resulting in a fragment of 716 bps that was digested with NotI/BamHI and cloned into pHPX7–GFP–SKL [21] resulting in pTDA1–GFP–SKL. To mark peroxisomes with GFP–SKL, the plasmid was linearised with BseYI and used to transform *S. cerevisiae* strains, and transformants were selected on YD plates without leucine. Correct integration was checked by colony PCR.

2.2.5.3. Construction of *pARM001*. The PEX14–mCherry region of pHIPN–PEX14–mCherry [20] was amplified using primer pair PRARM001_FWD/PRARM002_REV. The PCR product was digested with NotI/HindIII and cloned into pHIPH4 [22] that resulted into pARM001.
2.4 Western blotting

For the construction of plasmid pHIPZ4–mCherry fusinator, a PCR fragment of 700 bp was obtained by primer pair RSA10fw/RSA11rev on pCDNA3.1mCherry [23]. The resulting BglII–SalI fragment was inserted between the BglII and SalI of pANL31 [24].

2.5 Fluorescence microscopy

Antibodies conjugated to horseradish peroxidase were used for detection of G6PD, which was used as a loading control. Secondary antibodies (sc-9996) and rabbit polyclonal antiserum against glucose-6-phosphate dehydrogenase (G6PD) were used. Proteins of total cell extracts of trichloroacetic acid treated cells were separated by SDS-PAGE followed by Western blotting. Equal amounts of protein were loaded per lane. Blots were probed with rabbit polyclonal antiserum against GFP (Santa Cruz Biotechnology, sc-9996) and separately cloned into the vectors pBTM116-C and pVP16-C. The resulting plasmids were co-transformed with the indicated pVP16- and pBTM116-derived fusion constructs and transfectants were selected on synthetic medium lacking leucine and tryptophan. HIS3 reporter gene activation was detected by analysing growth on medium lacking histidine, leucine and 3-aminotriazole. From each co-transformation four independent transfectants were tested. Empty vectors were used to check for reporter self-activation. The well-established Hansenula polymorpha Pex3–H. polymorpha Pex19 interaction [18] was used as a positive control.

2.6. Yeast two-hybrid assays

The LexA system was used for screening interactions between S. cerevisiae proteins using derivatives of the reporter strain S. cerevisiae L-40 (Takara Bio Inc.). Using S. cerevisiae genomic DNA as template, the entire coding sequences of PNC1 and GPD1 were amplified with primer combinations PNC1.BamHI/F/PNC1.EcoRI/R and GPD1.BamHI/F/GPD1.EcoRI/R, respectively. The PCR fragments were digested with BamHI/EcoRI and separately cloned into the vectors pBTM116-C and pVP16-C, which yielded plasmids pBTM116–PNC1, pVP16–PNC1, pBTM116–GPD1 and pVP16–GPD1. S. cerevisiae L-40 was co-transformed with the indicated pVP16- and pBTM116-derived fusion constructs and transfectants were selected on synthetic medium lacking leucine and tryptophan. HIS3 reporter gene activation was detected by analysing growth on medium lacking histidine, leucine and 3-aminotriazole. From each co-transformation four independent transfectants were tested. Empty vectors were used to check for reporter self-activation. The well-established Hansenula polymorpha Pex3–H. polymorpha Pex19 interaction [18] was used as a positive control.

2.4 Western blotting

Proteins of total cell extracts of [26] trichloroacetic acid treated cells were separated by SDS-PAGE followed by Western blotting. Equal amounts of protein were loaded per lane. Blots were probed with mouse monoclonal antiserum against GFP (Santa Cruz Biotechnology, sc-9996) and rabbit polyclonal antiserum against glucose-6-phosphate dehydrogenase (G6PD), which was used as a loading control. Secondary antibodies conjugated to horseradish peroxidase were used for detection. Blots were scanned using a densitometer (Biorad).

2.5. Fluorescence microscopy

All fluorescence images were acquired using a 100 × 1.30 NA Plan-Neofluar objective (Carl Zeiss). Wide-field microscopy images were captured by an inverted microscope (Observer Z1; Carl Zeiss) using AxioVision software (Carl Zeiss) and a digital camera (CoolSNAP HQ²; Photometrics). GFP signal was visualised with a 470/40-nm band pass excitation filter, a 495-nm dichromatic mirror, and a 525/50-nm band pass emission filter. To visualise DsRed fluorescence, a 546/12-nm bandpass excitation filter, a 560-nm dichromatic mirror, and a 575–640-nm bandpass emission filter were used. mCherry fluorescence was visualised with a 587/25-nm band pass excitation filter, a 605-nm dichromatic mirror, and a 647/70-nm band pass emission filter.

To analyse the acquired fluorescence images ImageJ software (US National Institutes of Health, Bethesda, MD, USA) was used. For quantification, a straight line was drawn using ImageJ’s “line tool” through the region of interest and pixel intensity along the line was measured. The measured mean fluorescence intensity of GFP on peroxisomes and in the cytosol was corrected for the background intensity and a box plot was made using Microsoft Excel.

3. Results

3.1. Pnc1 is targeted to peroxisomes via piggy-back import with Gpd1

Piggy-back import requires that Pnc1 and Gpd1 physically interact in vivo. To test this we performed yeast two-hybrid analysis. As shown in Fig. 1, activation of the reporter gene HIS3, indicated by the capacity of yeast transformants to grow in the absence of histidine, was observed when PNC1 was co-expressed with GPD1. Similarly, growth was observed in strains co-expressing H. polymorpha PEX3 and PEX19, which were used as positive controls. Growth was not observed in control experiments using empty plasmids or in strains expressing either GPD1 or PNC1.

Next, we tested whether the peroxisomal localisation of Pnc1 depends on Gpd1. As shown in Fig. 2A, B, in cells producing chromosomally tagged GFP-fusion proteins, Gpd1–GFP and Pnc1–GFP are predominantly co-localising with the peroxisomal marker protein DsRed–SKL, in conjunction with low fluorescence in the cytosol. Accumulation of the proteins in the nucleus was not observed. The peroxisomal localisation of Pnc1–GFP was fully abolished in Δgpd1 cells (Fig. 2C). Conversely, in Δpnc1 cells the peroxisomal localisation of Gpd1–GFP was unaffected (Fig. 2D). This result indicates that import of Pnc1 depends on the presence of Gpd1, consistent with piggy-back import.

Next, we tested the effect of the removal of the PTS2 from the N-terminus of Gpd1 on the localisation of ΔN-Gpd1–mCherry and Pnc1–GFP fusion proteins. As shown in Fig. 2E in cells producing the N-terminal truncated Gpd1 the peroxisomal localisation of both proteins was abolished.

Fig. 1. In vivo interaction of Gpd1 and Pnc1. Full length Gpd1 and Pnc1 were tested for interaction using a yeast two-hybrid assay. Genes were fused to the LexA binding domain (BD) in vector pBTM116-C and a VP16 activation domain (AD) in vector pVP16-C. The resulting plasmids were co-transformed into S. cerevisiae L-40. The interaction between H. polymorpha Pex3 and Pex19 is used as positive control [18]. As negative controls, empty pVP16-C or pBTM116-C was used. HIS3 reporter gene activation was detected by analysis of growth on plates lacking histidine.
We also investigated whether we could restore import of NΔGpd1-mCherry into peroxisomes by the addition of a C-terminal PTS1 to Pnc1 (Pnc1–SKL). In Δgpd1 cells all NΔGpd1-mCherry was cytosolic (Fig. 3A; compare Fig. 2E). However, upon co-production of Pnc1–SKL a portion of the NΔGpd1–mCherry protein localised to peroxisomes (Fig. 3B). The import of only a minor portion of the total NΔGpd1–mCherry is most likely related to the fact that Gpd1 is present in large excess relative to Pnc1 (see below).

Fig. 2. The peroxisomal localisation of Pnc1 depends on Gpd1. Fluorescence microscopy images showing the localisation of chromosomally tagged Gpd1–GFP (A) and Pnc1–GFP (B) in S. cerevisiae cells producing DsRed–SKL as red peroxisomal matrix marker. Localisation of Pnc1–GFP in Δgpd1 cells (C) or Gpd1–GFP in Δpnc1 cells (D) both producing DsRed–SKL as peroxisomal marker. (E) Δgpd1 cells producing Pnc1–GFP and NΔGpd1–mCherry. The bar represents 5 μm.
Taken together, these results strongly suggest that Pnc1 is imported into peroxisomes via piggy-backing with Gpd1.

3.2. Pnc1 and Gpd1 stability

If both proteins form a stable complex in vivo, the absence of one component of the complex might cause instability of the other proteins and vice versa. To test this we analysed the levels of Pnc1–GFP in Δgpd1 cells and vice versa in cells in the stationary (T = 0), early exponential (T = 4), mid-exponential (T = 8 h) and late-exponential (T = 12 h) growth phase. The levels of Pnc1–GFP are comparable in Δgpd1 cells relative to the wild-type (PNC1–GFP) control strain (Fig. 4A). Similarly, the levels of Gpd1–GFP are not reduced in Δpnc1 cells (Fig. 4B). These results imply that Pnc1 and Gpd1 are not required for the stability of each other. In the absence of Gpd1, Pnc1 localises to the cytosol. As no major difference in Pnc1 levels is observed between wild-type and Δgpd1 cells the peroxisomal versus cytosolic localisation apparently also does not affect Pnc1 stability.

3.3. Gpd1 and Pnc1 are not present at a fixed stoichiometry

Previous transcriptional analysis indicated that GPD1 and PNC1 expression is strongly correlated [11]. In order to analyse whether the protein levels of Gpd1 and Pnc1 are correlated as well, we performed Western blot analysis using strains producing GFP fusion proteins and anti-GFP antibodies. Upon exposure of cells for 4 h to 1 M sorbitol,
Fig. 5. Modulations in Gpd1 and Pnc1 protein levels and localisation upon exposure of cells to stress. (A) Western blots showing the protein levels of Pnc1–GFP and Gpd1–GFP after exposure to various stress conditions for 4 h. Cells producing chromosomally tagged GFP-fusion proteins under control of their endogenous promoter were used for the experiments. Blots were probed with antibodies against GFP. G6PDH was used as loading control. (B) Quantification of Gpd1–GFP and Pnc1–GFP protein levels from 2 separate blots of 2 independent experiments. Error bar = +/- STDV. Fluorescence microscopy images of Pnc1–GFP (C) and Gpd1–GFP (D) cells at control conditions (unstressed) or upon exposure to stress for 4 h. The scale bar represents 5 μm. The box plot shows mean fluorescence intensities of Pnc1–GFP (E) and Gpd1–GFP (F) at peroxisomes or in the cytosol after 4 h of stress. Fluorescence intensities were measured using ImageJ. The box represents values from the 25 percentile to the 75 percentile; the horizontal line through the box represents the median value. The bar represents maximum and minimum values. For each experiment the fluorescence intensity of at least 100 peroxisomes and the cytosol of at least 44 cells were measured.
1 M NaCl or elevated temperature (37 °C), the levels of both proteins were enhanced (Fig. 5A). Quantification of the protein levels revealed that the ratio between Gpd1 and Pnc1 was approximately 7:1 in unstressed control cells, but increased considerably in cells exposed to stress (up to ~11:1 upon exposure to 1 M NaCl) (Fig. 5B). This result indicates that both proteins most likely do not form a hetero-oligomeric complex of fixed stoichiometry. Notably, at all conditions analysed, Gpd1 was present in large excess relative to Pnc1.

3.4. Gpd1 and Pnc1 are localised to peroxisomes and the cytosol in cells exposed to stress

Jung and colleagues reported that exposure of cells to stress results in a decrease in peroxisomal Gpd1 and Pnc1 levels concomitant with a relative increase in cytosolic protein levels [11]. In contrast, Anderson and colleagues showed that Pnc1 mainly remains peroxisomal in cells exposed to stress [1].

We performed quantitative fluorescent microscopy to analyse the mean cytosolic and peroxisomal fluorescent intensities of Gpd1–GFP (Fig. 5D) and Pnc1–GFP (Fig. 5C) before and after exposure of cells to various stress conditions. Because we rarely observed significant accumulation of Gpd1 or Pnc1 in the nucleus at our experimental set up, we excluded nuclear GFP signal in our analysis.

Our data indicate that at all three stress conditions tested (1 M sorbitol, 1 M NaCl, heat stress) the fluorescence intensities of peroxisomal Pnc1–GFP and Gpd1–GFP increased relative to the controls (Fig. 5E; F; Table 1). All three stress conditions also resulted in an increase in cytosolic Gpd1–GFP, whereas enhanced cytosolic Pnc1–GFP was only detected upon exposure of cells to 1 M NaCl.

To test whether this behaviour is specific for Gpd1 and Pnc1, we performed a control experiment in which we produced the PTS2 protein Pot1 (thiolase) containing a C-terminal GFP under control of the GPD1 promoter. Western blot analysis of GFP fusion proteins using anti-GFP antibodies revealed that similar protein levels were obtained for Pot1–GFP and Gpd1–GFP upon exposure of cells to osmotic stress (Fig. 6A).

Quantitative fluorescent microscopy (images shown in Fig. 6B, C & D) indicated that similar to Gpd1–GFP the fluorescence intensity of Pot1–GFP also increased in both the cytosol and peroxisomes after treatment of cells with 1 M sorbitol or 1 M NaCl (Fig. 6E, F).

These results indicate that import of Gpd1 and Pnc1 is not blocked upon exposure of cells to stress. However, our data suggest that the capacity of the PTS2 import machinery is insufficient to fully import the enhanced levels of Gpd1 and Pnc1 at these conditions.

4. Conclusions

In this paper we show that yeast Pnc1 piggy-back imports with Gpd1 into peroxisomes. Almost all peroxisomal matrix proteins contain either a PTS1 or PTS2 sorting sequence. However, proteins lacking a PTS can be imported in complex with a PTS containing protein by so called piggy-back import. Many reported examples of peroxisomal piggy-back import are artificial as these involve import of a subunit of an oligomeric protein from which the PTS is removed in complex with subunits that still contain a PTS [27–29]. So far only a few examples of natural piggy-back import have been described. These include import of the PTS lacking Cu/Zn superoxide dismutase 1 (SOD1) with its PTS-containing chaperone in mammalian cells [30]. In plant, import of two PTS lacking subunits of heterotrimeric protein phosphatase depends on the PTS containing third subunit of this enzyme [31]. Here, we report natural piggy-back import of Pnc1 with the PTS2 protein Gpd1 into yeast peroxisomes. Effelsberg et al. [32] recently reported the same observation. In addition, these authors showed that import of Gpd1 requires the general Pex7 co-receptor Pex21, a protein that is constitutively produced. Instead the second Pex7 co-receptor Pex18 is induced by olate and selectively required for import of the β-oxidation enzyme thiolase.

Why Pnc1 piggy-back imports with Gpd1 and not with another PTS2 or PTS1 protein is highly speculative. Possibly this is related to the fact that Gpd1 and Pnc1 are both involved in stress response and nucleotide metabolism. Also their expression is regulated in a similar manner.

Although we observed that Pnc1 and Gpd1 interact in a two-hybrid assay, Pnc1 and Gpd1 most likely do not form a stable complex. This view is based on the finding that i) the absence of one protein did not affect the stability of the other and ii) the cellular ratio of both proteins is not constant. Also, we were unable to show a stable physical interaction between both proteins using a variety of in vitro approaches (data not shown). These observations imply that complex formation between Gpd1 and Pnc1 likely is transient and possibly only required for sorting of Pnc1 to peroxisomes.

Our data confirm earlier reports which indicated that upon exposure of yeast cells to stress conditions the protein levels of Gpd1 and Pnc1 increase. Our quantitative fluorescent microscopy analyses showed that under these conditions the intensities of Gpd1–GFP and Pnc1–GFP inside peroxisomes increased. This increased peroxisomal signal is in contract with previous observations of Jung and colleagues, who reported a decrease in peroxisomal signal for both proteins [11]. However, our data are in line with the report of Anderson and colleagues, who showed that Pnc1–GFP remains predominantly peroxisomal upon exposure of cells to stress.

At all stress conditions tested we observed an increase in cytosolic fluorescent intensities of Gpd1–GFP, but not of Pnc1–GFP, for which an increase in cytosolic fluorescence was only detected upon exposure of cells to NaCl. This could be due to the higher total Gpd1 protein levels upon exposure of cells to stress (see Western blots in Fig. 5A). Also the ratio of Gpd1–Pnc1 increased at stress conditions, thus rendering more Gpd1 molecules available to Pnc1 for piggy-back import.

When a control PTS2 protein (thiolase, Pot1–GFP) was produced under control of the GPD1 promoter, we also observed the appearance of cytosolic fluorescence when cells were exposed to stress. This result suggests that PTS2 protein import is inefficient in cells exposed to stress, probably because the PTS2 import pathway is easily saturated.

Consequently, both proteins most likely play their cellular function inside peroxisomes and not in the cytosol or nucleus as previously proposed. What their function is in peroxisomes is still very speculative. Peroxisomes are highly oxidative organelle and therefore maintenance of a proper redox environment is crucial for functioning of peroxisomal enzymes. Because both proteins are involved in nucleotide metabolism, possibly they play a role in maintaining the redox balance in the peroxisomal matrix.

Table 1

<table>
<thead>
<tr>
<th>Condition</th>
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Transparency document

The Transparency document associated with this article can be found in the online version.
Fig. 6. Localisation of Pot1–GFP. (A) Western blot analysis using antibodies against GFP showing the levels of Gpd1–GFP and Pot1–GFP before and after exposure of cells to stress. G6PDH was used as a loading control. (B–D) Fluorescence microscopy images of cells producing Pot1–GFP (under control of the GPD1 promoter) and Gpd1–GFP (also under control of the GPD1 promoter) after exposure to stress for 4 h. (B) Unstressed cells (C) 1 M sorbitol and (D) 1 M NaCl stress. Scale bar represents 5 μm. The box plots show the mean fluorescence intensities of Pot1–GFP and Gpd1–GFP in the cytosol (E) or peroxisomes (F). The box represents values from the 25 percentile to the 75 percentile and the horizontal line through the box represents the median value. The bar represents maximum and minimum values. Fluorescence intensities were measured using ImageJ. For cytosolic fluorescence at least 29 cells were measured per experiment. For fluorescence at peroxisomes, at least 89 organelles were analysed per experiment.
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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jbamcr.2015.10.017.

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