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# **University of Alberta**

Peroxisome Dynamics in the Yeast *Saccharomyces cerevisiae*

By

Franco Joseph Vizeacoumar



A thesis submitted to the Faculty of Graduate Studies and Research in partial fulfillment of the requirements for the degree of Doctor of Philosophy

Department of Cell Biology

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> > Ä

**DATE**

**"I can do everything through Him who gives me strength"**

**Philippians 4:13**

# **TO MY DEAREST**

# APPA, AMMA, FREDDY and JESS

for their unwavering support and encouragement

#### **Abstract**

This thesis describes the identification and characterization of five novel *PEX* genes, namely *PEX28, PEX29, PEX30, PEX31* and *PEX32.*

The peroxins *PEX28* and *PEX29* were identified because of their high sequence similarity to Pex24p of the yeast *Y. lipolytica.* Peroxisomes of cells deleted for either or both of the *PEX28* and *PEX29* genes are increased in number, exhibit extensive clustering, are smaller in area, often exhibit membrane thickening between adjacent peroxisomes and have a decreased buoyant density compared to peroxisomes isolated from wild-type cells. Our data suggest that Pex28p and Pex29p. together with Pexllp, Pex25p and Vpslp, regulate peroxisome number, size and distribution in *S. cerevisiae.*

The peroxins *PEX30, PEX31* and *PEX32* were identified because of their high sequence similarity to Pex23p of the yeast *Y. lipolytica.* Cells deleted for *PEX30* exhibit increased numbers of peroxisomes, while cells deleted for *PEX31* and *PEX32* exhibit enlarged peroxisomes. PexSOp, Pex31p and Pex32p interact within themselves and with Pex28p and Pex29p. *PEX28* and *PEX29* function upstream of *PEX30, PEX31* and *PEX32* in the regulation of peroxisome size and number.

Additionally, we have addressed the role of Pexl9p in the assembly of the PMPs, PexSOp and Pex32p. Systematic truncations from the carboxyl terminus, together with inframe deletions of specific regions, have identified the mPTSs essential for the targeting of Pex30p and Pex32p to peroxisomes. Both Pex30p and Pex32p interact with Pexl9p in regions that do not overlap with their experimentally identified mPTSs. However, Pexl9p is required for localizing PexSOp and Pex32p to peroxisomes, because mutations that disrupt the interaction of Pex30p and Pex32p with Pex19p lead to their mislocalization to a compartment other than peroxisomes.

Pex19p also interacts with the dynamin-like protein Vps1p. Mutation of one of the putative Pexl9p-binding regions does not affect vacuolar assembly but leads to cells with reduced numbers of enlarged peroxisomes, the phenotype of cells lacking Vps1p. Our data suggest that the interaction of Pex30p, Pex32p and Vpslp with Pexl9p is required for their roles in peroxisome biogenesis and are consistent with a role for Pexl9p in stabilizing membrane proteins in peroxisomes.

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**CHAPTER ONE**

 $\sim$ 

**INTRODUCTION**

## <span id="page-23-0"></span>**1.1 Principles of organelle biogenesis**

Eukaryotes have evolutionarily organized various cellular processes into functionally distinct, membrane enclosed compartments called organelles. The ability to keep numerous chemical processes separate from one another in distinct organelles has permitted increased metabolic complexity and the concomitant diversification of cellular functions in eukaryotes. Hence, understanding the functionality of these organelles has increasingly become more complex, especially, as these structures are often derived from each other, share many of the same components with one another, communicate with the rest of the cell and behave dynamically. Organelle biogenesis requires at least three conceptually distinct processes: the formation of lipid bilayer, the insertion of membrane proteins into this lipid bilayer and the import of soluble proteins across the membrane into the matrix.

Most cellular proteins encoded by nuclear genes are transported to their destined organelles by certain cis-acting sequences that are recognized by cytosolic chaperones/receptors. On the surface of the organelles protein translocation machinery moves the protein across the membrane or inserts it into the membrane. The intricate details that occur at the surface of mitochondria (Hartl *et al.*, 1989; Schatz and Dobberstein, 1996; Schwartz and Neupert 1994; Rapaport 2003; Koehler 2004), endoplasmic reticulum (ER) (Blobel and Dobberstein, 1975; Blobel, 1980; Sabatini *et al*., 1982; Lyman and Schekman, 1995; Haigh and Johnson, 2002; Romisch *et al.,* 2003) and the chloroplast (Robinson and Klösgen, 1994) are now well-understood. The corresponding events that occur at the peroxisomal membrane are just beginning to be

elucidated (Hettema and Tabak, 2000; Veenhuis *et al.,* 2002; Eckert and Erdmann 2003; Schliebs and Kunau, 2004). Additionally, for efficient segregation, cells must have a mechanism to ensure the growth and partitioning of these functionally intact organelles. This mechanism will help the cell to regulate the number, size, composition and distribution of organelles to meet varying cellular demands.

This chapter will focus on the molecular mechanisms involved in these processes during the biogenesis of peroxisomes.

# <span id="page-24-0"></span>**1.2 Peroxisomes**

Peroxisomes are ubiquitous organelles found in most eukaryotes. Peroxisomes, together with the glyoxysomes of plants and glycosomes of trypanosomes, make up the "microbody" family of organelles. They play diverse roles in the cell, compartmentalizing many activities related to lipid metabolism and functioning in the decomposition of toxic hydrogen peroxide. It was Rhodin in 1954, who first identified these microbodies in ultra-structural studies in mouse kidney cells. Later de Duve and colleagues coined the term "peroxisome" as they found both  $H_2O_2$ -forming oxidase and degrading catalase during their studies on rat liver (de Duve and Baudhuin, 1966). Since then, related organelles were described in plants (glyoxysomes) and in trypanosomes (glycosomes) (Breidenbach and Beevers, 1967; Opperdoes and Borst, 1977). Morphologically, peroxisomes are spherical in shape ranging from 0.1 um to 1 um in diameter, bound by a lipid bilayer membrane and are of high equilibrium density in Nycodenz (~1.23 g/cm<sup>3</sup>) at maturation. In the filamentous fungi *Neurospora crassa*. specialized peroxisomes that contain a hexagonal crystalloid core have been described (Jedd and Chua 2000). Unlike mitochondria, peroxisomes do not contain DNA (Kamiryo *et al.,* 1982) and all peroxisomal proteins are encoded in the nucleus.

## <span id="page-25-0"></span>**1.2.1 Functions of Peroxisomes**

Peroxisomes perform diverse metabolic roles including certain specialized roles in different organisms (reviewed in Titorenko and Rachubinski, 2001; Eckert and Erdmann, 2003). Two widely distributed and well-conserved functions are fatty acid  $\beta$ oxidation and  $H_2O_2$ -based respiration. These processes appear to be ubiquitous, occurring in the peroxisomes of animals (Lazarow and de Duve, 1976), plants (Beevers *et al.*, 1969) and yeasts (Tanaka *et al.,* 1982). Additionally, in humans, peroxisomes are involved in the synthesis of ether lipids (plasmalogens) that are abundant in nervous tissue such as myelin (Hajra *et al*., 1979; Hajra and Bishop, 1982), cholesterol and bile acid synthesis (Keller *et al.,* 1985, 1986; Krisans, 1996), and in the degradation of long chain fatty acids and phytanic acid (Mannaerts and Van Veldhoven, 1996; Olivier *et al.,* 2000). In fungi, these organelles are involved in the biosynthesis of penicillin. In higher plants, glyoxysomes house enzymes of the glyoxylate cycle, photorespiration and ureide biosynthesis (Van den Bosch *et al.,* 1992). In some yeast species, peroxisomes are involved in methanol and/or amine oxidation and assimilation (Veenhuis *et al.,* 1983). Also recent developments have shown that peroxisomes function as an intracellular signaling compartment and as an organizing platform that orchestrates certain developmental decisions from inside the cell (Titorenko and Rachubinski, 2004). Remarkably, the peroxisomal sorting of HIV Nefl is mandatory for the life cycle of the

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HIV virus and is essential for the development and manifestation of AIDS (Cohen et al., **2000).**

## <span id="page-26-0"></span>**1.2.2 Peroxisomes in Human health**

Goldfisher in 1973, established the link between the absence of peroxisomes (although 'peroxisomal ghosts' were reported in later studies) and Zellweger syndrome. Later various groups reported other peroxisomal abnormalities such as increased amount of very long chained fatty acids and deficiency of plasmalogen in Zellweger patients (Brown *et al*., 1980; Heyman *et al*., 1983). These findings triggered intense research in the field of peroxisome biology. Rapid progress has largely depended on extensive clinical, metabolic, cellular and molecular studies in different model organisms and in patients with inborn errors of these processes. Genetic disorders of peroxisome biogenesis and function are broadly classified into Peroxisomal Biogenesis Disorders (PBDs) and Peroxisomal Metabolic Disorders (PMDs). PBDs are a genetically heterogeneous group of autosomal recessive human disorders in which peroxisomes fail to assemble properly and do not import critical matrix proteins (Lazarow and Moser, 1994; Brosius and Gartner, 2002). PBDs comprise 12 complementation groups and are characterized by deficiency of multiple peroxisomal functions. At present, all genes responsible for each of these complementation groups have been identified. Clinically, PBDs are classified into two broad spectra namely, Zellweger spectrum (ZS) accounting for about 80% of the PBD patients and Rhizomelic chondrodysplasia punctata (RCDP).

Zellweger spectrum includes three overlapping phenotypes comprising Zellweger syndrome, neonatal adrenoleukodystrophy, and infantile Refsum disease of which

Zellweger syndrome is the most severe. They are characterized by severe neurological and hepatic dysfunction, mental retardation, craniofacial abnormalities, and hypotonia. Patients have very high levels of phytanic acid and very-long chain fatty acids in the blood and cannot synthesize plasmalogens, a very important class of phospholipids in nervous tissue (Schutgens *et al.,* 1986). Survival beyond infancy for these patients is extremely rare. Peroxisomes in the cells of this group of patients are unable to import proteins targeted by either PTS1 or PTS2 (Peroxisomal Targeting Signals), and thus most of their peroxisomal metabolic functions are impaired.

Rhizomelic chondrodysplasia punctata is characterized by severe growth defects, rhyzomelia, cataracts, proximal limb shortening and ichthyosis (Lazarow and Moser, 1995). These patients have normal blood levels of very-long chain fatty acids but have increased levels of phytanic acid and also lack plasmalogens. Unlike ZS patients, peroxisomes in the cells of RCDP patients are deficient in the import of PTS2 proteins only and have normal PTS1 import (Braverman *et al.,* 1997; Motley *et al.,* 1997; Purdue *etal.,* 1997).

Peroxisomes in cells of ZS patients were originally thought to be absent (Goldfischer *et al.,* 1973), but later studies revealed the existence of peroxisomal structures in cell lines from several patients (Santos *et al.,* 1988). In fact, studies showed that while matrix proteins containing PTS1 or PTS2 were mislocalized to the cytosol in these cell lines, peroxisomal membrane proteins (PMP) were correctly targeted. This indicates that the machinery involved in the matrix protein import is different from that of peroxisomal membrane proteins (Slawecki *et al.,* 1995; Hettema *et al.,* 2000).

PMDs do not involve a general mislocalization of peroxisomal matrix proteins. Rather, the milder symptoms associated with this class of disorder are the result of mutations in single genes which affect the function or targeting of individual peroxisomal enzymes. Examples of PMDs are X-linked adrenoleukodystrophy, acatalasaemia (catalase deficiency) and adult Refsum disease.

#### <span id="page-28-0"></span>**1.3 Assembly and proliferation of peroxisomes**

Assembly and proliferation of peroxisomes are controlled by a set of proteins called *peroxins* and the genes that encode them are called *PEX* genes. During the commencement of this work, only 24 *PEX* genes were known. An update of peroxins and their characteristic functions are listed in the Table 1-1. Peroxisome biogenesis was once believed to occur similar to that of endosymbiotic mitochondria and chloroplasts. However, identification of a number of core components has greatly changed this view stressing the significance of diverse mechanisms of transport of proteins into the organelles. They differ from other organelles in that, first, unlike mitochondria, peroxisomes do not contain DNA and peroxisomal proteins are encoded by the nucleus. Second, peroxisomes can arise newly in cells that apparently lack peroxisomal remnants. Third, protein unfolding is not a prerequisite for import of matrix proteins, suggesting a novel mechanism for the translocation of oligomers across the peroxisomal membrane. Currently, understanding the mechanism of membrane biogenesis, matrix protein import and the division and segregation of mature peroxisomes from the mother cells to the daughter cells are areas of intense research.

## **Table 1 1 Peroxins and their characteristic features**





# <span id="page-30-0"></span>**1.3.1 Endoplasmic reticulum as the template for origin**

Early studies suggested that peroxisomes are formed by budding off from the endoplasmic reticulum, primarily because of studies suggesting that peroxisome membranes were spatially associated with the ER *in vivo* (Novikoff and Shin, 1964). This concept was then abandoned leading to the generally accepted growth and division model

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of Lazarow and Fujiki (1985), which states that new peroxisomes arise exclusively by budding and fission of pre-existing peroxisomes. Evidence in support of this model came from the demonstration that the peroxisomal proteins were synthesized in the cytosol and then post-translationally imported into the peroxisome (reviewed in Lazarow and Fujiki, 1985). Also, studies showed that peroxisomal matrix proteins lacking PTSs were mislocalized to the cytosol and not the ER (Gould *et al.,* 1989; Elgersma *et al.*, 1996).

However, intense research has dramatically increased our knowledge in the biogenesis of peroxisomes resulting in the acceptance of the original hypothesis that peroxisomes indeed arise from the ER. Some of the major evidence for the ER origin of peroxisomes are listed here. In rat liver, the predominantly peroxisome-associated integral membrane protein PMP50 is preferentially synthesized on ER-bound ribosomes (Bodnar and Rachubinski, 1991). In *H. polymorpha*, the N -terminal portion of the peroxisomal membrane protein (PMP) Pex3p directs a fused reporter protein to the ER when overexpressed (Baraends et al., 1996). Also, observations like overproduction of *H*. *polymorpha* PMPs Pex3p and Pexl4p, *S. cerevisiae* PMP PexlSp and human PMP Pex3p leading to a profound proliferation of ER membranes, strongly suggests in a link between ER and peroxisomes (Elgersma *et al.,* 1997; Kammerer *et al.,* 1998; Komori *et al.,* 1997).

The C-terminal portion of *S. cerevisiae* Pexl5p, produced at a normal level, directs a fused reporter protein to the ER, and when overproduced, accumulates in the ER and is O-mannosylated in the ER lumen (Elgersma *et al.,* 1997). Unfortunately, in most of these experiments the trafficking of the protein from the ER to the peroxisome was not tested and the protein was found in the ER only when overproduced. Some proteins like Pex3p, Pex8p and Pex14p of *H. polymorpha* and the peroxisomal ascorbate peroxidase

(pAPX) of cottonseed accumulate in the ER in brefeldin A (BFA)-treated cells and can be chased from the ER to peroxisomes after removal of BFA (Salomons *et al.*, 1997; Mullen *et al.,* 1999). Recruitment of ADP-ribosylation factor and COPI in vitro by rat liver peroxisomes requires the PMP Pexl lp (Passreiter *et al.,* 1998). *S. cerevisiae* Pexl5p and cottonseed pAPX post-translationally insert *in vitro,* in an ATP-and cytosolic Hsp70 chaperone-dependent manner, only into ER membranes but not into peroxisomal or any other purified organelle membrane (Mullen *et al.,* 1999). In fact, pAPX was largely detected in the sub domains of rough ER in the cells of wild type Arabidopsis (Lisenbee) *etal,* 2003).

In the yeast *Y. lipolytica,* Titorenko and co-workers have demonstrated that mutations in certain secretory genes, as well as in two *PEX* genes *{PEX1* and *PEX6),* can result in the accumulation of the PMPs Pex2p and Pex16p in the ER (Titorenko *et al.*, 1997). Specifically, mutations in sec238A and srp54, were found to result in the accumulation of coated vesicles containing,  $Pex2p$ ,  $Pex16p$ , and the COPII components Secl3p and Sec23p (Titorenko and Rachubinski, 1998). Although Ben Distel and coworkers have shown that COP-I and COP-II vesicles are not necessary for targeting proteins to peroxisomes (Voom-Brouwer *et al.,* 2001), the same group has eventually shown by three-dimensional image reconstruction of peroxisomes, that ER serves as a major source from which peroxisomal membrane is derived (Geuze *et al.,* 2003). Recently, it has been shown that Pex10p of *Arabidopsis* is localized within the sub domains of ER from where it participates in peroxisome formation (Flynn *et al.*, 2005). Taken together, these observations strongly suggest that the ER serves as a template in peroxisome biogenesis. Yet, interestingly enough, a requirement for a known ERprotein import and export pathways, is still left to be established.

#### <span id="page-33-0"></span>**1.3.2 Import of matrix proteins into peroxisomes**

Peroxisomal matrix proteins have been found to be targeted to peroxisomes by peroxisome targeting signals (PTS). PTS1 is a conserved tripeptide located at the extreme C-terminal end of a protein (Gould *et al.*, 1987, 1989). It consists of the sequence Ser-Lys-Leu or conserved variants of these residues. Most peroxisomal matrix proteins contain a PTS1 (Gould *et al.,* 1989; Aitchison *et al.,* 1991; Swinkels *et al.,* 1992; Motley *et al.,* 1995; Elgersma *et al.,* 1996). Mutants defective only in the import of PTS1 proteins led to the identification of the PTS 1 receptor, Pex5p, in *P. pastoris* (McCollum *et al.,* 1993; Terlecky *et al.,* 1995). Subsequently, Pex5p was identified in several other organisms (van der Leij *et al.,* 1993; Szilard *et al.,* 1995; van der Klei *et al.,* 1995), including humans (Dodt *et al.,* 1995). All Pex5p orthologs share a region comprised of six to seven tetratricopeptide repeats (TPRs) within the C-terminal half of the protein, which binds the cargo proteins (Brocard *et al.,* 1994; Dodt *et al.,* 1995; Fransen *et al.,* 1995; Otera *et al.,* 2002). The TPR domains form two clusters hinged by TPR 4 and the PTS1 signal binds to the groove between the TPR clusters (Gatto *et al.,* 2000). Pex5p functions as a mobile receptor, shuttling between the cytosol and the peroxisomes (Dammai and Subramani, 2001). These authors demonstrated proteolytic cleavage of a Pex5p fusion protein exclusively within the peroxisomes and detected the cleaved Pex5p in the cytosol.

A second PTS, PTS2, was also identified and is a conserved N-terminal nonapeptide (Arg/Lys) (Leu/Val/Ile) Xaas (His/Gin) (Leu/Ala). Although, many mammalian proteins were shown to contain this signal, *Saccharomyces* seem to have only one protein, 3-ketoacyl-CoA thiolase, which is sorted through this signal (Swinkels *et al.,* 1991; Glover *et al.,* 1994b). In the nematode C. *elegans,* the PTS2 import pathway is completely absent (Motley *et al.,* 2000; Petriv *et al.,* 2004). Mutants defective solely in the import of PTS2 proteins led to the identification of the PTS2 receptor, Pex7p, in several organisms (Marzioch *et al.,* 1994; Rehling *et al.,* 1996; Elgersma *et al.,* 1998), including humans (Braverman *et al.,* 1997). Human Pex7p deficiency causes rhizomelic chondrodysplasia punctata (Braverman *et al.,* 1997; Motley *et al.,* 1997; Purdue *et al.,* 1997). Pex7p contains six WD repeats, which bind the cargo.

Redundant proteins Pexl8p and Pex21p, which are predominantly cytosolic, interact with Pex7p while it binds the cargo, facilitating the PTS2 pathway (Purdue *et al.,* 1998). In the yeast *Yarrowia lipolytica*, Pex20p seems to fulfill the task of Pex7p as well as Pexl8p and Pex21p; since this organism lacks Pex7p (Complementation studies of Einwachter *et al.,* 2001). Like Pex5p, Pex7p is also proposed to function as a mobile receptor (reviewed in Purdue and Lazarow, 2001). Apart from binding their ligands, Pex5p and Pex7p, bind to the components of the docking complex, Pex13p and Pex14p (Elgersma et al., 1996; Albertini et al., 1997; Girzalsky et al., 1999; Brocard et al., 1997; Will *et al.*, 1999). The interaction of Pex5p with Pex14p is much stronger when Pex5p is loaded with cargo. In contrast, the interaction with Pexl3p is stronger when Pex5p is not loaded with cargo (Urquhart *et al.,* 2000). Similarly, thiolase, a PTS2 protein, interacts with Pexl4p, via Pex7p, and not with Pexl3p (Stein *et al.,* 2002). These data suggest that Pexl4p acts as the initial docking site for the cargo loaded receptors, which is subsequently transported to other components of the import machinery. This is supported by observations that Pex5p is cytosolic in mammalian CHO *pex!4A* cells, but membrane associated in  $per13\Delta$  cells, and that Pex5p accumulates on peroxisomal membrane when Pexl4p (but not Pexl3p) is overexpressed (Otera *et al*., 2000). However, Pexl3p has been reported to be required for the targeting of  $Pex14p$ , thus suggesting both  $Pex13p$  and Pexl4p form the initial docking complex. Recently, Schell-Steven *et al.,* (2005) have shown that the association of Pex13p with the docking complex is essential for matrix protein import and this association is stabilized by Pex5p.

Pex5p was also found to interact with Pex2p, PexlOp and Pexl2p (Chang *et al.,* 1999a; Reguenga *et al.*, 2001). The level of Pex5p on the membrane seems to be unaffected by the effect of deleting or overexpressing these peroxins (Chang *et al.*, 1999a; Okumoto *et al.*, 2000). However, a mutation in Pex12p causes an accumulation of Pex5p in the lumen of the peroxisomes (Dodt and Gould 1996). Thus, these three RING finger peroxins are suggested to play a role in translocation of peroxins.

A few peroxins such as Pexlp, Pex4p, Pex6p and Pex22p have been suggested to act after matrix protein translocation, as the steady state level of Pex5p was affected in these mutants (Collins *et al.,* 2000). Based on this, it was suggested that these peroxins are involved in later events of import, probably in the recycling of the receptor. But this idea certainly deserves deeper investigation because in *Yarrowia lipolytica.* Pexlp and Pex6p are required for the fusion of small peroxisomal vesicles (Titorenko and Rachubinski, 2000; Titorenko and Rachubinski, 2001).
Another remarkable feature of peroxisomal protein import is that folded and assembled multimeric proteins can enter the organelle (Walton *et al.,* 1992, 1995; Glover *et al.,* 1994a; McNew and Goodman, 1994; Flynn *et al.,* 1998). The oligomerization of some of these multimeric proteins is facilitated by specific chaperone molecules, as in the case of the soluble peroxisomal matrix protein thiolase and its chaperone Pex20p (Titorenko *et al.*, 1998) or is self-assisted, as in the case of the heteropentameric fatty acyl-CoA oxidase (AOX) complex of the yeast *Yarrowia lipolytica* (Titorenko *et al.*, 2002). For Aox, the association of each subunit with FAD occurs in the cytosol, prior to transportation into the peroxisomes. However, monomers of *H. polymorpha* alcohol oxidase are transported into the peroxisomes and FAD is added inside peroxisomes (Evers *et al.,* 1994, 1996; Waterham *et al.,* 1996; Stewart *et al.,* 2001). This requires specific chaperones in the lumen of the peroxisomes. It is interesting to note that, although few chaperones have been identified to be involved in peroxisome biogenesis (Preisig-Muller *et al.*, 1994; Pause *et al.*, 1997; Hettema *et al.*, 1998), only hsp70 has been localized to peroxisomes (Walton *et al.,* 1994; Corpas and Trelease, 1997; Diefenbach and Kindi, 2000).

#### **1.3.3 Import of membrane proteins into peroxisomes**

The sorting of peroxisomal membrane proteins is much less understood and appears to be independent of matrix protein import. Evidence for this came from the identification of peroxisomal ghosts, spherical membranous structures containing peroxisomal membrane proteins (Hettema *et al.,* 2000). The same study has shown that Pex3p and Pex19p are the only two peroxins that are required for membrane biogenesis in *S. cerevisiae.* Similar results were obtained in human fibroblasts cell lines silenced for *PEX3* or *PEX19* (Matsuzono *et al.,* 1999; Shimozawa *et al.,* 2000). From these results, Pex3p and Pexl9p were thought to define essential components involved in the early biogenesis of the membrane. Pex3p is an integral membrane protein with its Cterminus exposed to the cytosol. It interacts with Pexl9p, a famesylated peroxin both *in vivo* and *in vitro* (Sacksteder *et al.,* 2000; Snyder *et al.,* 1999; Gotte *et al.,* 1998).

Pexl9p is mostly cytosolic with a smaller fraction of it found to be associated with the peroxisomes. It interacts with a number of peroxisomal integral membrane proteins (PMPs) (Snyder *et al.,* 1999; 2000). Shibata *et al.,* (2004) has recently shown that mammalian Pexl9p interacts with Pex3p through its flexible N-terminal half while its C-terminal region is responsible for its interaction with other PMPs. The cytosolic fraction of Pexl9p has been suggested to act as a shuttling receptor for peroxisomal membrane proteins, while its peroxisome-associated form could function as an assembly factor, assisting the assembly of multimeric complexes following their binding to the peroxisome membrane (reviewed in Subramani, 1998; Hettema *et al.,* 1999; Terlecky and Fransen, 2000; Subramani *et al.,* 2000; Purdue and Lazarow, 2001; Titorenko and Rachubinski, 2001; Schleibs and Kunau, 2004). However, other protein sorting mechanisms may also exist. Several peroxins undergo N- or O- linked glycosylation, suggesting that these proteins are sorted to peroxisomes via ER. This is especially true in the yeast *Yarrowia lipolytica* where, Pex2p and Pexl6p are N-Glycosylated (Titorenko and Rachubinski 1998).

Recent evidence from Gould's lab suggested that Pexl9p could be bifimctional acting both as a cytosolic PMP chaperone and as a PMP import receptor (Jones *et al..* 2004). Also, the same group has shown that Pex3p acts as a docking factor for Pexl9p (Fang et al., 2004). When considering all the above mentioned roles of Pex19p, a number of concerns are raised. First, Pexl9p's (unction as PMP receptor is questionable as a number of studies have shown independently that the Pex19p binding region of PMPs does not overlap with their m-PTS signal (Fransen *et al.,* 2001, 2004; Snyder *et al.,* 2000; Biermanns and Gartner 2001). Second, no convincing data has so far been obtained determining if Pexl9p recognize nascent PMPs. Snyder *et al.,* (2000) showed in *Pichia pastoris* that blocking protein synthesis followed by cross linking, the amount of PMP immunoprecipitated by Pexl9p remained unchanged compared to the untreated cells and have concluded that Pexl9p interacts with preexisting and not with newly synthesized pool of PMPs. Also, by subcellular fractionation of the cross linked cells, they found that the steady state of interaction occurs at the peroxisome and not in the cytosol. Thirdly, Pexl9p is an extremely hydrophilic protein and it is intriguing as to how it could shield the TMDs (Trans Membrane Domains) to act as a chaperone. This role also requires that Pex19p must bind at multiple regions of the PMPs. Fourth, in *Candida boidinii*, Pex19p binding to PMP47 is oleic acid induced and that in non-inducing conditions, another nonredundant mPTS targets PMP47 to peroxisomes (Wang *et al.,* 2004). Fifth, in *Yarrowia lipolytica,* Pexl9p is not involved in the membrane biogenesis as Pexl9p null mutants appear to contain peroxisome-like structures (Lambkin and Rachubinski 2001). Sixth, overproduction of Pex3p in a *PEX19* deletion strain results in the formation of peroxisomal protein-containing vesicles (Otzen *et al.,* 2004). Thus, understanding the exact functional role of Pex19p in membrane biogenesis certainly need extensive investigation.

#### **1.3.4 Regulation of peroxisomal abundance – fission and segregation**

Organelle division is a dynamic process that takes place once the organelle is inherited. Division is orchestrated by multi-component protein complexes that assemble and drive the constriction and fission of the organellar membranes (Osteryoung, 2001; Shaw and Nunnari, 2002). Peroxisomes are highly plastic and their size, number and shape vary dynamically based on the organism and environmental ques (Chang *et al.,* 1999b; Gould *et al.,* 2001; Purdue and Lazarow, 2001). Certain naturally occurring lipids and xenobiotics have been shown to increase peroxisome number and induce expression of genes encoding peroxisome matrix and membrane proteins (Reddy *et a l,* 1986; Reddy and Hashimoto, 2001; Zomer *et al.*, 2000). Evidence for metabolic control of peroxisome division has also been presented (Poll-The *et al.,* 1988; Sacksteder and Gould, 2000; Smith *et al.*, 2000) and might be mediated by signals derived from the B-oxidation of fatty acids (Chang *et al.,* 1999b; van Roermund *et al.,* 2000).

An increase in peroxisome number is usually achieved by maturation and division of preexisting peroxisomes (South and Gould, 1999; Gould *et al.,* 2001). It has been suggested that peroxisomes can undergo constitutive division in the absence of a stimulator or a regulated division under induced conditions (Marshall *et al.,* 1996). But weather the components required for these different types of division are one and the same or different is not clear.

Members of the Pexl 1p family of peroxins, including Pex25p (Smith *et al.*, 2002) and Pex27p (Tam *et al.,* 2003; Rottensteiner *et al.,* 2003a) of *S. cerevisiae*, have been shown to affect peroxisome division in different organisms (Erdmann and Blobel, 1995; Marshall *et al.,* 1995; Sakai *et al.,* 1995; Li and Gould, 2002; Li *et al.,* 2002). In the yeast

*Y. lipolytica*, the membrane-bound pool of Aox interacts with Pex16p, a membraneassociated protein that negatively regulates the division of early intermediates in the assembly pathway. This interaction inhibits the negative action of Pexl6p, thereby allowing mature peroxisomes to divide (Guo *et al.,* 2003; Titorenko and Rachubinski, 2004).

Another family of proteins that were thought to be involved in organelle fission are the dynamins. Dynamin and dynamin related proteins are highly conserved family of large GTPases involved in a variety of cellular processes such as endocytosis, intracellular protein trafficking and organelle partitioning (Hinshaw, 2000; Danino and Hinshaw, 2001). The yeast genome contains three genes coding for dynamin related proteins. Studies in mitochondria have demonstrated the involvement of one of the members, Dnm1p in mitochondrial fission. Dnm1p structures assemble at the outer membrane and at a rate-limiting step in the divisional pathway, mediate the division of the mitochondrial membranes (Osteryoung and Nunnari, 2003).

Recently, Hoepfner *et al.* (2002) has convincingly demonstrated that Vps1p is involved in peroxisome fission and that peroxisomes are transported via actin cables to the bud in a Myo2p-dependent process. Vps1p is more related to DLP1, a human homologue of Vps1p, than to conventional dynamins. In mammalian cells, it has been shown that peroxisome elongation and constriction can occur independently of DLP1, while the fission of peroxisomes requires it (Koch *et al.*, 2003; 2004). Pex11p has been suggested to indirectly recruit DLP1 to the peroxisomes, although the authors were unable to detect any physical interaction between these two proteins (Li and Gould 2003). V ps1p has also been suggested to regulate actin cytoskeleton structure through its

interaction with Slalp (Yu and Cai, 2004). Slalp is known to play an important role in the regulation of actin cytoskeleton through its interaction with several proteins capable o f promoting actin assembly, such as Panlp, Lasl7p, and Abplp (Tang *et al.,* 1997; Howard *et al.,* 2002; Warren *et al.,* 2002; Gourlay *et al.,* 2003). Interestingly, *vpslArholA* double mutants showed accumulation of actin patches on peroxisomes, which suggests that the majority of actin is reorganized/disassembled before organelle fission (Marelli et *al.*, 2004). How may Vps1p and the Pex11p family of proteins regulate the fission of peroxisomes is still unclear. Thus, as it appears molecular mechanism of peroxisome division and segregation with high fidelity, especially as it progresses through cell cycle is poorly understood.

### **1.4 Experimental approaches to study peroxisome biogenesis**

The molecular mechanism of peroxisome assembly has been essentially conserved from yeasts to humans. Studies combining classical cell biological analyses and standardized biochemical analyses with global biology approaches such as transcriptome profiling, organellar proteomics, database mining and genomic comparative analyses, to determine novel genes, were applied in various model organisms.

### **1.4.1 Yeast as a model system**

Easy and cost effective cultivation has made yeast an ideally suited model system for studying the mechanisms of various cellular dynamics. In the yeast cells, peroxisomes

are the only sites of fatty acid  $\beta$ -oxidation (Tanaka *et al.*, 1982) and induction of peroxisome proliferation occurs when the cells are grown on certain "peroxisomerequiring" carbon sources such as oleate (Veenhuis *et al.*, 1987; Kunau *et al.*, 1988) or methanol (Fukui *et al.*, 1975). Peroxisomes proliferation in yeast cells can either be accelerated by carbon sources such as oleic acid or repressed by growing on carbon sources such as glucose, which do not require the metabolizing functions of peroxisomes for conversion to a usable energy source. This remarkable quality is exploited to construct mutants, thereby facilitating the cloning of genes required for peroxisome assembly. Also, the presence of large and numerous peroxisomes facilitate their isolation for biochemical studies and morphological analysis. Many different yeast species are currently being used as model systems for peroxisome biogenesis. For our studies, we use *Saccharomyces cerevisiae* as numerous molecular tools are available to manipulate the genome.

### **1.4.2 C lassical genetic screening**

Early approaches in studying peroxisome biogenesis used yeast genetic screens to identify mutants of peroxisome assembly. Yeast cells are mutagenized in order to make *pex* mutants defective in peroxisome biogenesis that are isolated through either positive or negative selection procedures. These mutants are then rescued by transformation with a genomic library in order to complement the mutant phenotype. The complementing gene is isolated and its gene product characterized (Elgersma and Tabak, 1996). Fujiki and colleagues developed an alternative method to identify mammalian genes that depends on functional complementation of PBD-CHO cells (Peroxisome defective

Chinese Hamster Ovary) with human cDNA expression libraries. These methods have been powerful, as 32 individual *PEX* genes have been isolated to date from CHO cells (Tsukamoto *et al.,* 1990; Shimozawa *et al.,* 1992; Suzuki *et al.,* 2001) and a variety of yeast species, including *S. cerevisiae* (Erdmann *et al.*, 1989; Vizeacoumar *et al.*, 2004), *P. pastoris* (Gould *et al.,* 1992), *H. polymorpha* (Cregg *et a l,* 1990) and *Y. lipolytica* (Nuttley *etal,* 1993).

### **1.4.3 Live cell im aging**

One of the rapidly developing techniques to screen mutants and identify novel genes is by live cell imaging-4D microscopy. Fluorophores like GFP (Green Fluorescent Protein) and its variants, cyan fluorescent protein (CFP), yellow fluorescent protein (YFP) and *Discosoma* sp. red fluorescent protein (DsRed) are used as reporters to identify various cellular compartments (Ward *et al.,* 1998). An improved photo-stable GFP+ is also now available (Scholz et al., 2000). A library of yeast strains genomically tagged with GFP has been extensively used to identify localization of over 70% of proteins (Huh *et al.,* 2003). This study has facilitated to identify a novel protein, Ymr204p, as a component involved in the inheritance of peroxisomes. Currently, further characterization o f this protein is in progress in our laboratory

### **1.4.4 Systems biology approaches**

Bioinformatics and systems biology tools have revolutionized and have greatly replaced the traditional approach of single gene analysis. Homology probing has exploited the evolutionary conservation of genes from yeast to mammals to identify mammalian genes and genes in other organisms. *PEX28, PEX29, PEX20, PEX31* and *PEX32* were identified this way in *S. cerevisiae.* Proteomics approaches were used to identify components of the yeast protein translocation apparatus (Erdmann and Blobel, 1996). Microarray screening was used to identify oleic acid-induced yeast genes and this way, *PEX25* and its homolog *PEX27* were identified (Smith *et al,* 2002; Tam et *al.,* 2004). And more recently, ICAT (isotope-coded affinity tag) coupled with mass spectrometry has revealed that the small GTPase, Rholp to be enriched in the peroxisomal fraction. Further analysis has revealed that Rholp is involved in peroxisome movement along actin (Marelli *et al.,* 2005). Genome wide phenomics has facilitated to identify Ymr204p as a novel protein that localizes to peroxisomes. Further characterization of this protein is under progress in our laboratory. Genome wide interaction data has facilitated the identification of Ymr163p, as another component of peroxisome biogenesis that interacts with Pexl9p.

### **1.5 Focus of this thesis**

The identification of genes involved in peroxisome assembly and elucidation of the roles of the proteins they encode would provide greater understanding of the molecular bases of lethal disorders like Zellweger syndrome. Completion of the *S. cerevisiae* genome sequencing project proved invaluable in that it provides the opportunity to identify these components through sequence similarity between proteins of unknown function encoded by the *S. cerevisiae* and proteins already shown to be required for peroxisome biogenesis in other organisms. Rachubinski's laboratory has extensively used the model organism *Y. lipolytica* to identify such novel components necessary for peroxisome biogenesis. Thus taking advantage of this, I report the identification and characterization of five genes in *S. cerevisiae*, of previously unknown function and localization, now renamed *PEX28, PEX29, PEX30, PEX31* and *PEX32.* Additionally, I report on how their protein products are targeted to the peroxisomal membrane defining a role for Pexl9p in this process.

# **CHAPTER TWO**

# **EXPERIMENTAL PROCEDURES**

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# **2.1 Materials**

# **2.1.1 List of reagents and chemicals**





# **2.1.2** List of enzymes



### **2.1.3 Multicomponent systems**



## **2.1.4 Plasmids**



### **2.1.5 Molecular size markers**



### 2.1.6 Antibodies

### **2.1.6.1 Primary antibodies**



'Dilutions are for use in western blotting. Ten-fold higher concentrations are used for immunofluorescense.

### **2.1.6.2 Secondary antibodies**



Secondary antibodies were either conjugated to horseradish peroxidase (HRP) for immunoblotting or to a fluorescent dye molecule (FITC (fluorescein) or TRITC (rhodamine)) for indirect immunofluorescence

### **2, L7 Software used in this study**



### **2.1.8 Oligonucleotides**

Oligonucleotides were either synthesized on an Oligo 1000M DNA synthesizer

(Beckman) or ordered from SIGMA. Oligonucleotides used in this study are described in

the following table.

### **Table2-1 Oligonucleotides used in this study**





pGBT9-PEXl9



 $32$ 







# **2.1.9 Standard buffers and solutions**



### **2.2 Microorganisms and culture conditions**

### **2.2.1 Y east strains and culture conditions**

The yeast strains and media components used in this study are listed in Table 2-4 and Table 2-2, respectively. All strains were cultured at 30°C as 10 ml cultures in glass tubes in a rotating wheel. For subcellular fractionation and similar experiments requiring large quantity of cells, cells were grown in flasks in a rotary shaker at 180 rpm. For all experiments, an overnight culture was subcultured so that cells grew to an  $OD_{600}$  of 0.75.

### **2.2.2 Bacterial strains and culture conditions**

*E. coli* strains and culture media used in this study are described in Tables 2-2 and 2-1, respectively. Bacteria were grown at 37°C unless stated otherwise. Cultures of 5 ml or less were grown in culture tubes in a rotary shaker at  $\sim$ 180 rpm. Cultures greater than 5 ml were grown in flasks in a rotary shaker at 250 rpm. Culture volumes were approximately 20% of flask volumes.

**Table 2-2 Media components**

<b>Medium</b>	Composition	Reference
LB.	1% tryptone, 0.5% yeast extract, 1% NaCl	Maniatis et al., 1982
<b>SOB</b>	2% tryptone, 0.5% yeast extract, 10 mM NaCl, 2.5 mM KCl	Maniatis et al., 1982
<b>TYP</b>	1.6% tryptone, 1.6% yeast extract, 0.5% NaCl, 0.25% K <sub>2</sub> HPO <sub>4</sub>	Promega Protocols and Applications
<b>SCIM</b>	0.67% YNB, 0.5% yeast extract, 0.5% peptone, 0.5% Tween (w/v) 40, 0.1% glucose, 0.15% (v/v) oleic acid, 1 $\times$ CSM	Guide, 1989/1990 Erdmann et al., 1989
Sporulation	1% potassium acetate, 0.1% yeast extract, 0.05% glucose	Rose et al., 1988
medium <b>CSM</b>	0.67% YNB, 2% glucose, $1 \times$ CSM without histidine and/or leucine	<b>Bio101</b>
<b>YEPA</b>	1% yeast extract, 2% peptone, 2% sodium acetate	Brade, 1992
<b>YEPD</b>	1% yeast extract, 2% peptone, 2% glucose	Rose et al., 1988
<b>YNA</b>	0.67% YNB, 2% sodium acetate	Brade, 1992
<b>YND</b>	0.67% YNB, 2% glucose	Rose et al., 1988
<b>YNO</b>	0.67% YNB, 0.05% (w/v) Tween 40, 0.1% (v/v) oleic acid	Nuttley al., et 1993
<b>YPBO</b>	0.3% yeast extract, 0.5% peptone, 0.5% $K_2HPO_4$ , 0.5% KH <sub>2</sub> PO <sub>4</sub> , 0.2% (w/v) Tween 40 or 1% (v/v) Brij 35, 1% (v/v) oleic acid	Kamiryo al. et 1982

### **Table 2- 3 Antibiotics used in this study**





## **Table 2-4 Strains used in this study**



 $GALI<sub>TATA</sub>$ -*lacZ*, *MELI* 



 $\mathcal{L}$ 





### **2.3 Introduction of DNA into microorganisms**

DNA was introduced into microorganisms via chemical transformation or eletroporation, and transformants were selected by antibiotic resistance or other genetic markers. Electroporation was performed in microelectroporation chambers (width -0.15 cm) using a Cell-Porator connected to a Voltage Booster (BRL).

### **2.3.1 Chemical transformation of** *E. coli* **(Heat shock)**

Plasmid DNA was introduced into transformation-competent DH5 $\alpha$  cells (subcloning efficiency) as described in a protocol from Invitrogen. Generally, 1 to 2  $\mu$ L of a ligation reaction or 0.25  $\mu$ g of plasmid DNA was added to 25  $\mu$ L of cells. The mixture was incubated on ice for 30 min, subjected to heat shock at  $37^{\circ}$ C for 20 sec and then returned to ice for 2 min. 1 ml of LB was added, and the cells were incubated in a rotary shaker at  $37^{\circ}$ C for 45 min. Cells were spread onto LB agar plates containing ampicillin or another antibiotic for selection. When necessary, 75  $\mu$ L of 2% X-gal in DMF and  $50 \mu L$  of 100 mM IPTG were spread onto plates prior to the plating of cells to select for blue/white colonies carrying recombinant plasmids. Plates were incubated at  $37^{\circ}$ C for  $\sim$ 16 h for colony formation.

### **2.3.2 Electroporation of** *E. coli*

High-efficiency electroporation-competent *E. coli* cells were prepared according to Maniatis *et al.* (1982). Freshly grown *E. coli* were used to inoculate 50ml of S.O.B. medium (Table 2-1) and were grown overnight at 37°C. Cultures were then diluted into

500 ml S.O.B. and grown for 2 to 3 h at  $37^{\circ}$ C until the OD<sub>600</sub> was approximately 0.8. Cells were harvested by centrifugation and washed twice with 500 ml of sterile, ice-cold 10% (v/v) glycerol. The pellet was then resuspended in 10% (v/v) glycerol to a final volume of 2 ml, and 20 µl aliquots were subjected to electroporation using a Voltage Booster (Whatman Biometra) at 395 V (amplified to  $\sim$ 2.4 kV) at a capacitance of 2  $\mu$ F and a resistance of 4 k $\Omega$ . Cells were then diluted to 1 ml with LB medium, grown for 1 h at 37°C and plated onto LB-agar plates containing ampicillin.

#### **2.3.3 Chemical transformation of** *S. cerevisiae*

This method is largely used to transform plasmid DNA into yeast cells (Gietz and Woods, 2002). Essentially, 25  $\mu$ l of cells was scraped from a freshly grown plate with a sterile toothpick and resuspended in 1 ml of water. Cells were washed twice by centrifugation, resuspended in 1 ml of 100 mM lithium acetate, and incubated at 30°C for 10 min. Cells were again harvested by centrifugation, and the following components were added on top of the cell pellet in the following order: 240  $\mu$ l of 50% (w/v) PEG, 36  $\mu$ l of 1 mM lithium acetate, 50  $\mu$ l of 2 mg sheared, denatured salmon sperm DNA/ml, 1  $\mu$ l of plasmid DNA and 20 µl of water. Alternatively, a master mix can also be made without the plasmid, if different DNA molecules were to be transformed. The mixture was vortexed vigorously for 1 min and incubated at 42°C for 20 min. Cells were harvested by centrifugation, resuspended gently in 200 µl of water and plated onto CSM (Complete Synthetic Medium) for selection. Plates were incubated at 30°C for 3 days for colony formation.

### **2.3.4 Electroporation of** *S. cerevisiae*

Yeast cells were made electrocompetent following the method of Ausubel *et al.* (1996). Cells were grown overnight in 10 ml of YEPD. 5 ml of culture was added to 50 ml of YEPD, and cells were grown to an  $OD_{600}$  of  $~0.75$ . Cells were harvested by centrifugation, resuspended in Transformation buffer (pH 7.5) (Table 2-4) containing 10 mM lithium acetate and 20 mM DTT and incubated for Ih. Cells were harvested by centrifugation and washed successively with 50 ml each of ice-cold water, ice-cold 1M sorbitol. Cells were resuspended in a minimal volume of ice-cold sorbitol buffer. 20  $\mu$ L of cells was added to  $1 \mu L$  of plasmid DNA, and the cells were suspended between the bosses of an ice-cold electroporation chamber (Whatman Biometra). Cells were subjected to 250V (amplified to ~1.6 kV) at a capacitance of 2  $\mu$ F and a resistance of 4 k $\Omega$ . Cells were immediately added to 100  $\mu$ L of ice-cold 1 M sorbitol and plated onto CSM agar plates (Table 2-6). Plates were incubated at 30°C for 2 to 3 days for colony formation.

### **2.4 Isolation of nucleic acids**

Chromosomal DNA from yeast cells and plasmid DNA from *E. coli* cells were isolated as follows:

#### **2.4.1 Plasmid DNA isolation from** *E. coli*

Plasmid DNA was isolated from small-scale bacterial cultures according to the alkaline lysis mini-prep procedure (Ausubel *et al.,* 1996) or by using the QIAprep Spin Miniprep Kit according to the manufacturer's instructions. Single colonies were inoculated into 2 ml of LB containing ampicillin and grown over night. Plasmid DNA

was isolated from 1.5 ml of culture by the alkaline lysis method (Maniatis *et al.*, 1982) or by using a QIAprepMiniprep Kit (Qiagen). This method employs essentially the same principles as the alkaline lysis method, except that after precipitation of proteins with potassium acetate solution, plasmid DNA is absorbed onto a silica-gel membrane in a high-salt environment. Chaotropic salts are passed over the column to remove contaminating proteins. Residual salts are removed, and plasmid DNA is eluted in 50  $\mu$ L of 10 mM Tris-HCl (pH 7.5).

### **2.4.2 Genomic DNA isolation from** *S. cerevisiae*

Total genomic DNA was isolated from yeast strains using the glass bead disruption technique (Ausubel *et al.,* 1989) in breakage buffer. Cells were grown overnight in 10 ml cultures of YEPD, washed three times with water and then resuspended in breakage buffer. Glass beads were added, tubes were vortexed three times for 1 min each, and then cell debris was removed by centrifugation for 5 min at maximum speed in a microfuge. Nucleic acids were separated from proteins with two phenol/chloroform/isoamyl alcohol (25:24:1) extractions and one chloroform/isoamyl alcohol  $(24:1)$  extraction. Total nucleic acids were precipitated by the addition of 2.5 volumes of absolute ethanol, pelleted by centrifugation in a microfuge for 4 min at maximum speed, and subsequently washed twice with 70% ethanol to remove contaminating salts. The nucleic acid pellet was then resuspended in approximately  $50 \mu l$ of TE/RNase A (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 20  $\mu$ g RNase A/ml) and incubated at 37°C to allow for RNA digestion and pellet dissolution. If plasmid DNA is to be recovered, subsequent electroporation into *E. coli* is performed and recovered.

### **2.5 Standard DNA manipulation**

Unless otherwise stated, reactions were in 1.5 ml microcentriuge tubes, and microcentrifugation was done in an Eppendorf microcentrifuge at 16,000 x *g.*

### **2.5.1 Amplification of DNA by the polymerase chain reaction**

PCR was used to amplify specific DNA fragments, to introduce restriction endonuclease sites within a DNA molecule or to facilitate the construction of hybrid DNA molecules. Integration of DNA fragments into the genome or deletion of specific sequences from the genome were also verified by PCR. PCR conditions (including primer design, cycling conditions and reaction components) were according to standard procedure (Innis and Gelfand, 1990; Saiki, 1990). Reactions were performed in 0.6 ml microcentrifuge tubes and typically contained 1.25 U of Platinum *Pfx* DNA polymerase, or *Taq* polymerase, 0.1 pg of template DNA, 20 to 100 pmol of each primer, 1 mM  $Mg_2SO_4$ , 25 mM each of dATP, dCTP, dGTP and dTTP in 50 $\mu$ L of reaction buffer. Alternatively, Ready-To-GO PCR beads were used according to the specifications of the manufacturer (Amersham Biosciences). Reactions were cycled in a Robocycler 40 (Stratagene) with a Hot Top attachment.

### **2.5.2 Restriction endonuclease digestion**

DNA was digested according to the restriction enzyme maufacturer's instructions. For diagnostic and preparative digests,  $0.5$  to  $1\mu$ g and 2 to  $3\mu$ g of DNA, respectively, were digested respectively for 1 to 1.5 h at  $37^{\circ}$ C by 1 U of enzyme under optimal buffer concentration. Double digests were done according to the instructions supplied by NEB.

### **2.5.3 Dephosphorylation and phosphorylation of 5'ends**

Prior to ligation, the 5' ends of linearized plasmid DNA molecules were usually dephoshorylated to prevent intramolecular ligations. Essentially, after plasmid digestion, reactions were mixed with 5 U of CIP and incubated for 30 min at 37°C. Also prior to ligation, the 5' termini of DNA molecules amplified by PCR were phosphorylated. Reactions were mixed with 10 U of T4 polynucleotide kinase and ATP and PNK buffer (to 10 mM and  $1 \times$  respectively) and incubated at 37 $\degree$ C for 1 h. Desphosphorylation and phosphorylation reactions were terminated by agarose gel electrophoresis of the DNA fragments.

### **2.5.4 Separation of DNA fragments and PCR products**

DNA fragments produced in the course of molecular cloning and polymerase chain reaction (PCR) products were first separated by electrophoresis using 1% agarose in  $1 \times$  TBE buffer containing 5 µg ethidium bromide/ml. Gels were run at 8-10 v/cm of gel. Prior to loading, samples were mixed with 6 x Gel Dye. Fragments 100 bp in length or smaller were separated on 3% agarose gels consisting of 0.5% SeaKem Genetic Technology Grade (GTG) agarose and 2.5% NuSieve GTG agarose. Large DNA fragments were separated on  $0.7$  to  $1.5\%$  agarose gels. The bands of interest were

visualized by ultraviolet illumination (Photodyne, Model 3-3006) and excised from the gel. The QIAquick gel extraction kit was used to recover the DNA.

### **2.5.5 Purification of DNA fragments from agarose gels**

DNA fragments were isolated from agarose gels using a QIAquick Gel Extraction Kit (Qiagen) according to the manufacturer's instructions. The gel fragment containing the DNA of interest is dissolved, the DNA is adsorbed to a silica-gel surface, contaminants are removed by washing and the DNA is eluted in 30  $\mu$ L of 10 mM Tris-HCl (pH 8.5).

### **2.5.6 Purification of DNA from solution**

Salts and enzyme contaminants present with the DNA in solutions, obtained after digestion can be removed by using the QIAquick PCR Purification Kit as described by the manufacturer (Qiagen). The principle of this method is similar to that of the QIAquick Gel Extraction Kit, except that no dissolution of agarose gel was involved. DNA was usually eluted in 30 to 50 µl of 10 mM Tris-HCl (pH 8.5).

#### **2.5.7 Ligation of DNA fragments**

DNA fragments treated with restriction endonucleases and purified were ligated using  $1 \mu l$  of T4 DNA ligase in the buffer supplied by the manufacturer (NEB). The reaction was typically in a volume of 10  $\mu$ l, with the molar ratio of vector to insert being between 1:3 and 1:10, and incubated overnight at  $4^{\circ}$ C. Alternatively, 1 µl of Quick T4

DNA ligase (NEB) in  $1 \times$  Quick Ligation Buffer was used in a reaction volume of 20 pi. The reaction was incubated at room temperature for 10 min.

### **2.5.8 Molecular cloning and subcloning by ligation**

PCR products are conveniently cloned by using pGEMT vector system. The 3'-T overhang present at the ends greatly improve the efficiency of ligation of a PCR product into the plasmid by preventing the recircularization of the vector. The multiple cloning site is flanked by specific recognition sites for restriction enzymes and a single or double digestion may be used to release the insert from the vector. All molecular cloning was performed according to manufacturer's instructions (Promega).

#### **2.5.9 DNA sequencing**

Sequencing was performed with an ABI Prism 310 Genetic analyzer, an automated DNA sequencer, using BigDye Terminator Cycle Sequencing  $v1.1/3.1$  Ready Reaction Kit as described by the manufacturer (Applied Biosystems). The system uses the method of Sanger (Sanger *et al*, 1977). It involves random incorporation of fluorescent dideoxy terminators during the elongation of DNA sequences with a modified version of *Taq* DNA polymerase. The reaction master mix contains  $3 \mu L$  of Terminator Ready Reaction Mix, 1  $\mu$ L of plasmid DNA, 3.2 pmol of primer and 2.5  $\mu$ L of the supplied  $5 \times$  buffer in a total volume of 20  $\mu$ l. The reaction was subjected to cycle sequencing using the Robocycler 40 with a Hot Top attachment (Stratagene) with the following conditions: 1 cycle at 96°C for 2 min; 25 cycles at 96°C for 46 sec, 50°C for 51 sec and  $60^{\circ}$ C for 4 min 10 sec; 1 cycle at  $6^{\circ}$ C to hold until ready to purify. Reaction products were precipitated with 80 µl of 75% isopropanol for 20 min at room temperature, subjected to microcentrifugation at  $16,000 \times g$  for 20 min, washed twice with 250 µl of 75% isopropanol, dried in a rotary vacuum dessicator (Labconco, Centrivap concentrator) and resuspended in  $15 \mu l$  of Template Suppression Reagent. Following a brief vortexing, the reaction mixture was then heated at 95°C for 2 min and immediately cooled on ice. This is loaded onto the ABI 310 Genetic analyzer, which separates by capillary electrophoresis, and the emitted fluorescence was detected and recorded to obtain the sequence.

### **2.6 Analysis of Proteins**

### **2.6.1 Determination of protein concentration**

Total protein was measured according to Bradford (1976) using a protein assay kit (BioRad) and bovine serum albumin as the standard. A standard curve was made by adding 1 ml of Bio-Rad Protein Assay Dye to 100 µl aliquots of water containing a series of different concentrations of BSA. Samples were incubated for 5 min at room temperature and briefly vortexed, and absorbance was measured at 595 nm using a Beckman DU640 spectrophotometer. Absorbance values were plotted against the BSA concentrations to generate a standard curve. Absorbance of a protein sample was measured in the same way as for BSA standards, and the protein concentration was estimated by comparing the absorbance value with the standard curve.
## **2.6.2 Electrophoretic separation of proteins**

Protein samples were resolved according to the method of Laemmli (1970). Samples were denatured by boiling 5 min in SDS-PAGE sample buffer and then separated by discontinuous SDS-PAGE on a 10% gel. Samples were run at 50 to 200 V in 50 mM Tris-HCl, pH <sup>8</sup> .<sup>8</sup> , 0.4 M glycine, 0.1% SDS until the dye front reached the bottom of the gel. Otherwise, proteins were electrophoretically transferred to nitrocellulose with either a BioRad western transblotter apparatus overnight at 100 mA in 20 mM Tris-HCl, pH 7.5, 150 mM glycine, 20% (v/v) methanol or by semi-dry electrophoretic transfer (Tyler Research Instruments, Edmonton, AB) for 1.5 h at 0.75  $mA/cm<sup>2</sup>$  of transfer area, as described (Kyshe-Anderson, 1984).

## **2.6.3 Precipitation of proteins**

Proteins were precipitated from solution by adding TCA to a final concentration o f 10% and incubation on ice for 30 min to overnight. Precipitates were collected by microcentrifugation at  $16,000 \times g$  for 30 min at 4<sup>o</sup>C. The pellet was washed twice with 1 ml of ice-cold acetone, dried in a rotary vacuum dessicator and dissolved in  $2 \times$  sample buffer

## **2.6.4 Detection of proteins**

Proteins extracted from the cells and separated on gels were detected and quantitated by the following ways.

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## **2.6.4.1 Staining with Coomassie blue staining**

Proteins in polyacrylamide gels were visualized by staining with 0.1% Coomassie Brilliant Blue R-250, 10% (v/v) acetic acid, 35% (v/v) methanol for 1 h with gentle agitation. Unbound dye was removed by multiple washes in  $10\%$  (v/v) acetic acid,  $35\%$ (v/v) methanol. Gels were dried for 1 h at 80°C on a Bio-Rad Model 583 gel drier. Proteins were directly visualized by staining in 0.1% Coomassie Brilliant Blue (R-250),  $10\%$  (v/v) acetic acid, 35% (v/v) methanol.

## **2.6.4.2 Staining with Ponceau S**

Ponceau S (sodium salt) is used to prepare a stain for rapid reversible detection of protein bands on nitrocellulose membranes (Western blotting). Common stain formulations include 0.1% (w/v) Ponceau S in 5% acetic acid or 2% (w/v) Ponceau S in 30% TCA and 30% sulfosalicylic acid. Ponceau S stain is easily reversed with water washes, facilitating subsequent immunological detection.

#### **2.6.4.3 Immunoblot analysis**

Nitrocellulose was hydrated in water for 5 min and then equilibrated in Western transfer buffer. Proteins separated by SDS-PAGE were transferred to nitrocellulose in Western transfer buffer, using either a wet or semi-dry apparatus. For wet blotting, the transfer was carried out at 100 mA for approximately 16 h at room temperature or at 400 mA for approximately 4 h using a cooling coil, in a blotting tank (Bio-Rad Trans-Blot or Hoefer TE Series Transphor electrophoresis unit). For semi-dry blotting, the transfer was carried out at 0.8 mA per cm<sup>2</sup> of gel in an ET-20 electrophoretic transfer system (Tyler Research Instruments, Edmonton, Alberta). Unoccupied protein binding sites on the nitrocellulose were blocked by incubating the blot in TBST-milk (1% skim milk powder in TBST) for 30 min with gentle agitation. Primary antiserum was diluted in TBST-milk at their respective concentrations. The blots were incubated with the primary antiserum for 60 min at room temperature. Excess primary antibody was removed by three 10 min washes in TBST. The appropriate secondary antibody conjugated to HRP was diluted in TBST-milk at a concentration of 1:30,000 and incubated with the blots for 60 min. Excess secondary antibody was removed by three 10 min washes in TBST. The blot was covered with ECL detection solution (a 1:1 mixture of the two ECL detection reagents) for 1 min, placed in a transparent plastic folder and exposed to Kodak XK-1 X-ray film. For quantitation of signals, densitometry was performed with a Bio-Rad GS800 calibrated densitometer.

## **2.7 Cell Disruption and Subcellular fractionation**

Subcellular fractionation is an approach universally used with all cell types and tissues for the isolation of organelles. This method has been widely used to identify the different protein compositions of isolated cellular compartments.

## **2.7.1 Preparation of whole cell lysates**

Total cell lysates were prepared by different methods such as the glass bead lysis method or the alkaline lysis method.

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#### 2.7.1.1 Glass bead lysis method

Total cellular protein was isolated by glass bead disruption of cells. Cells grown in YEPD or YPBO were washed twice with water and pelleted by centrifugation at 2,500 x *g* and resuspended in ice-cold Disruption buffer containing protease inhibitors and 1 mM DTT (Needleman and Tzagoloff, 1975). Glass beads are added to the meniscus of cell suspension and vortexed for 5 min at 4°C. The suspensions were subjected to centrifugation in a microfuge for 5 min to pellet unbroken cells, cell debris and glass beads. Supernatants were removed and stored at -20°C or loaded onto the gels after mixing with the gel loading dye and boiling for 10 min.

#### 2.7.1.2 Magic method

Alternatively, alkaline and reducing agents are used to prepare yeast cell lysates. Cells were harvested by centrifugation at  $2,000 \times g$  for 5 min, transferred to a microcentrifuge tube, and resuspended in 240  $\mu$ l of 1.85 M NaOH and 7.4%  $\beta$ mercaptoethanol. The cell suspension was incubated on ice for 5 min and mixed with an equal volume of pre-cooled 50% TCA by vortexing. The mixture was further incubated on ice for 5 min and subjected to microcentrifugation at  $16,000 \times g$  for 10 min at 4<sup>o</sup>C. The pellet was washed once with water, resuspended first in 100 µl of Magic A (1 M unbuffered Tris-HCl and 13% SDS) and then in an equal volume of Magic B (30%  $(v/v)$ ) glycerol, 200 mM DTT and 0.25% bromophenol blue). The mixture was boiled for 10 min and then subjected to microcentrifugation at  $16,000 \times g$  for 30 seconds to pellet unbroken cells and cell debris. Supernatants were removed and stored at -20°C or loaded onto the gels.

## **2.7.2 Subcellular fractionation**

Subcellular fractionation and peroxisome isolation were done essentially as described (Bonifacino *et al.,* 2000; Vizeacoumar *et al*., 2003). Cells were grown in YEPD for 12 h, shifted to oleic acid-containing medium and induced for 9 h. Cells were harvested by centrifugation at  $800 \times g$  in a Beckman JA10 rotor at room temperature and washed twice with water. Cells were resuspended in perforation buffer, at a concentration of 10 ml per gram of wet cells, and incubated at  $30^{\circ}$ C for 30 min at 70 rpm to loosen the outer mannoprotein layer. Cells were collected by centrifugation at  $2,500 \times g$  in a precooled Beckman JS13.1 rotor for 8 min at 4°C and washed once with Zymolyase buffer without Zymolyase. Cells were then resuspended in Zymolyase buffer containing 0.125 mg of Zymolyase  $100T/ml$  at a concentration of 8 ml per gram of wet cells and incubated at  $30^{\circ}$ C for 45 min to 1 h at 70 rpm to obtain spheroplasts. The rest of the treatment is done at 4°C or on ice bucket. Spheroplasts were harvested by centrifugation at 2,200  $\times$  *g* in a Beckman JS13.1 rotor for 8 min at 4°C and washed once with washing buffer (1.2 M sorbitol, 5 mM MES, pH 6.0, 1 mM EDTA). They were then resuspended in buffer H containing  $1 \times$  complete protease inhibitor cocktail (Roche) at a concentration of 3 ml per gram of wet cells. Resuspended spheroplasts were transferred to a homogenization mortar and disrupted by 10-15 strokes of a Teflon pestle driven at 1,000 rpm by a stirrer motor (Model 4376-00, Cole-Parmer). Disruption is monitored in a microscope for the release of organelles. Cell debris, unbroken cells and nuclei were pelleted by centrifugation at  $1,000 \times g$  in a Beckman JS13.1 rotor for 8 min at 4°C. The resultant supernatant is the postnuclear supernatant (PNS). The PNS was subjected to four to five additional centrifugations at  $1,000 \times g$  in a Beckman JS13.1 rotor for 8 min at 4°C. The PNS was

fractionated by centrifugation at 20,000  $\times$  g in a Beckman JS13.1 rotor for 30 min at  $4^{\circ}$ C into pellet (20KgP) and supernatant (20KgS) fractions. The 20KgP fraction is enriched in organelles. Aliquots of PNS, 20KgS and 20KgP were stored at -20°Cfor later analysis. Proteins in equal portions of each fraction were separated by SDS-PAGE and analyzed by immunoblotting.

## **2.7.3 Isolation of organelles by isopycnic centrifugation**

The 20KgP fraction was resuspended in buffer H containing 11% Nycodenz and  $2 \times$  PINS (Protease Inhibitors), and a volume containing 5 mg of protein was overlaid onto either a 30-ml discontinuous gradient consisting of 17%, 25%, 35% and 50% (w/v) Nycodenz (6.6 ml of 17%, 16.5 ml of 25%, 4.5 ml of 35% and 3 ml of 50% (w/v) Nycodenz in buffer H) or a 30-ml continuous gradient of 25% to 50% (w/v) Nycodenz, both in buffer H containing PINS. Organelles were separated by centrifugation at 100,000  $\times g$  for 90 min in a VTi50 rotor (Beckman). Fractions of 2 ml were collected from the bottom of the tube by puncturing the optiseal tube using an 18 gauge needle. 18 fractions of 2 ml each were collected from the bottom of the gradient. The 20KgS fraction was subfractionated by ultracentrifugation at  $200,000 \times g$  in a Beckman TLA120.2 rotor for 1 h at 4°C into a pellet (200KgP) fraction enriched for high-speed pelletable organelles and a supernatant (200KgS) fraction enriched for cytosol. Aliquots of fractions were mixed with  $6 \times$  dye, boiled for 10 min and loaded onto gels.

## **2.7.4 Membrane extraction**

Peroxisomal membrane fractions were prepared by extraction of purified peroxisomes isolated from Nycodenz gradients, as described (Fujiki *et al.,* 1982; Nuttley *et al.*, 1990). Essentially, organelles in the 20KgP fraction (50 µg of protein) were lysed by quick freezing and thawing (3 times) followed by incubation in 10 volumes of Ti8 buffer (10 mM Tris-HCl, pH 8.0) containing  $3 \times$  PINS on ice for 1 h and separated into pellet (Ti8P) and supernatant (Ti8S) fractions by centrifugation at 245,000  $\times g$  for 1 h at 4°C in a TLA120.2 rotor (Beckman). The Ti8P fraction was resuspended in Ti8 buffer to a final protein concentration of 0.5 mg/ml, and a portion of the resuspended fraction was extracted with 0.1 M Na<sub>2</sub>CO<sub>3</sub>, pH 11.3, for 1 h on ice and then separated into supernatant (CO<sub>3</sub>S) and pellet (CO<sub>3</sub>P) fractions by centrifugation at 245,000  $\times$  *g* in a TLA120.2 rotor at  $4^{\circ}$ C for 1 h. Proteins in the Ti8S, Ti8P, CO<sub>3</sub>S and CO<sub>3</sub>P fractions were precipitated by addition of 50% TCA, and the precipitates were washed with acetone. Proteins in equal portions of each fraction were separated by SDS-PAGE and analyzed by immunoblotting. Alternatively, the purified peroxisomal fractions were also subjected to treatment with Ti8 or alkaline carbonate buffer.

## **2.8 Microscopy**

## **2.8.1 Immunofluorescence microscopy**

Imunofluorescence of yeast cells was performed as described (Pringle *et al.*, 1991; Smith, 2000). This technique was performed either to localize protein A chimeras or endogenous proteins. Strains encoding protein A chimeras were transformed with the plasmid pDsRed-PTSl, grown in SM medium for 12 h and then incubated in YPBO medium for 8 h. Following induction, cells were fixed in 3.7% (v/v) formaldehyde for 30 min at room temperature in an orbital shaker. Cells were then pelleted by centrifugation at 2,000  $\times$  *g* for 5 min, washed with 2 ml of solution B, and resuspended with the same solution B at a concentration of 1 ml per 100  $\mu$ l of wet cells. 1 ml of cell suspension was mixed with  $l$ mg of Zymolyase 100 T/ml and 28 mM  $\beta$ -mercaptoethanol and incubated for 1 h at 30°C with gentle rotation. Spheroplasts were spotted onto slides precoated with poly Z-lysine and allowed to dry at room temperature. Spheroplasts were permeabilized by immersion of the slides for 6 min in -20 $^{\circ}$ C methanol and for 30 sec in -20 $^{\circ}$ C acetone, and allowed to air dry. For the rest of the treatment, slides were put in a dark humid box at room temperature. Spheroplasts were covered with 50 pi of blocking solution (1% milk in TBST) for 1 h. Slides were incubated with primary antibody diluted in blocking solution for 1 h, washed 10 to 20 times with  $1 \times TBST$ , and then incubated with secondary antibody conjugated to fluorescein or rhodamine diluted in blocking solution for 1 h. Cells were washed again 10 to 20 times with  $1 \times T$ BST and covered with 1 drop of mounting medium. Coverslips were placed on top of slides, and the edges were sealed with nail polish. Protein A chimeras were detected using rabbit antiserum to mouse IgG (ICN, Aurora, OH) and fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG. Primary antibodies listed in Table 2.1.6 were used to detect endogenous proteins. Images were captured on a LSM510 META (Zeiss) laser scanning microscope or an Olympus BX50 microscope with a digital camera (Spot Diagnostic Instruments, Sterling Heights, MI).

#### **2.8.2 Staining of the yeast nucleus with Hoechst dye**

Cell nuclei of live yeast cells can be visualized using Hoechst 33342 dye. Common stain formulations include 1 mM dye (56 mg) in 1 % (v/v) ethanol (Stock). Cells expressing GFP chimeras were grown in YEPD, induced with YPBO for 8 h and were treated with 10 nM Hoechst dye (Sigma) per µl of cells at 30°C for 5 to 15 min in a rotary shaker at 150 rpm. Cells were washed once, and images were obtained using an Olympus BX50 microscope.

## **2.8.3 Staining of the yeast vacuole with FM4-64 dye**

Cells were grown overnight and subcultured to reach an  $OD<sub>600</sub>$  of 0.5. To about 100  $\mu$ l of cells, 1  $\mu$ l of 8 mM FM4-64 dye was added, and cells were incubated for 30 min at 30°C. Cells were washed and reincubated in fresh YPD medium for 1.5 h. These cells were washed with water twice before examination. Images were captured on an Olympus BX50 microscope equipped with a digital fluorescence camera.

## **2.8.4 Staining of yeast mitochondria with MitoTracker Red**

Mitochondria of live yeast cells were stained with MitoTracker Red CMXRos according to the manufacturer's instructions (Molecular Probes). Cells grown in 10 ml of YPBO (Table 2-9) were harvested, washed once with water and resuspended in 10 ml of pre-warmed YPBO. The cell suspension was mixed with  $2 \mu$  of 1 mM MitoTracker Red CMXRos and incubated at 30°C for 10 to 15 min in a rotary shaker at 150 rpm. 1.5 ml of

culture was pelleted, washed twice with water, and viewed on an Olympus BX50 microscope equipped with a digital fluorescence camera.

#### **2.8.5 Electron microscopy**

Whole cells and subcellular fractions were processed for EM as described (Eitzen *et al.,* 1997; Eitzen, 1997; Goodman *et al.,* 1990). All microcentrifugations were performed at  $16,000 \times g$  for 1 min, and all incubations were done in 1.5 ml microcentrifuge tubes at room temperature with agitation in an orbital shaker, unless indicated otherwise.

#### **2.8.5.1 Processing of whole cells**

Cells were harvested and washed twice with water. Approximately  $100 \mu l$  of cell pellet was fixed in 1 ml of 3% freshly prepared  $KMnO<sub>4</sub>$  for 20 min, washed twice with water, and incubated in 1 ml of 1% sodium periodate for 10 min. Cells were pelleted, washed once with water, and incubated with 1 ml of 1% NH4CI for 15 min. Cells were again pelleted, washed once with water, and subjected to serial dehydration in 60%, 80%, 95%, and 100% ethanol and in propylene oxide. Incubation in the graded alcohol was for 5 min and in propylene oxide for 3 min. Incubation in propylene oxide was repeated three times. Cells were collected and incubated in 1 ml of a 1:1 mixture of propylene oxide and resin (a mixture of TAAB 812 resin, specially distilled DDSA, methyl nadic anhydride and 2,4,6,-tri-(dimethylaminomethyl) phenol in proportions suggested by the manufacturer (Marivac)) for 1 h. Cells were next pelleted and resuspended in 1 ml of resin. Incubation in resin was carried out for 1 h with agitation and 3 h in a fume hood with caps opened. Finally, cells were pelleted by microcentrifugation for 8 min, and small portions of cells were transferred to embedding capsules (EMS) containing resin. Embedding capsules were placed in an oven at 60°C to allow the resin to polymerize. Ultra-thin sections were cut (performed by Honey Chan, Department of Cell Biology, University of Alberta) using an Ultra-Cut E Microtome (Reichert-Jung) and viewed on a Phillips 410 electron microscope. Images were captured with a digital camera (Soft Imaging System). For time point experiments, cells were prefixed in 1 ml of 3% glutaldehyde prepared in 0.1 M cacodylate buffer, pH 7.2, for 15 min at 4°C with agitation. In this way, cells could be stored at 4°C until needed and processed for electron microscopy as described above.

## **2.8.5.2 Processing of purified organelles**

To the 2 ml fractions of purified organelles in Nycodenz, 25% glutaraldehyde was added to a final concentration of 0.5%, and organelles were fixed for 15 min at 4°C with agitation. These fixed organelles were pelleted using the Type 70Ti rotor (Beckman), and the pellet was covered with 2.5% glutaraldehyde in 0.1M cacodylate buffer. This is left for further fixation overnight at 4°C with very gentle agitation. Care must be taken that the pellet does not get dispersed in the fixative. Once fixed, the pellet is washed three times with 0.1M cacodylate, followed by incubating with 10% osmium tetroxide for 5 min. The pellet turns brown and can be washed with water and dehydrated with a series of alcohol and propylene oxide, as is done for processing whole cells. Embedding was

done as described for whole cells. Ultrathin sections were cut and mounted onto carbon-coated nickel grids. For better contrast, samples were also poststained with 2% uranyl acetate.

## **2.8.6 Morphometric analysis of peroxisomes**

For each strain analyzed, electron micrographs of 50 randomly selected cells at  $\times$ 17,000 magnification were developed (captured on film and printed) and scanned, and the areas of individual cells and of individual peroxisomes were determined by counting the number of individual pixels in a particular cell or peroxisome with Image Tool for Windows, Version 2.00 (University of Texas Health Sciences Center, San Antonio, TX). Alternatively, for each strain analyzed, electron microscopic images of 50 randomly selected cells at  $\times$ 17,000 magnification were captured with a digital camera (Soft Imaging System), and the areas of individual cells and of individual peroxisomes were determined by the program analySIS 3.1 (Soft Imaging System). To determine the average area of a peroxisome, the total peroxisome area was calculated and divided by the total number of peroxisomes counted. To quantify peroxisome number, the numerical density of peroxisomes (number of peroxisomes per  $\mu$ m<sup>3</sup> of cell volume) was calculated by the method of Weibel and Bolender (1973) for spherical organelles as follows. First the total number of peroxisome profiles was counted and reported as the number of peroxisomes per cell area assayed  $(N_A)$ . Next, the peroxisome volume density  $(V_V)$  was calculated for each strain (total peroxisome area/total cell area assayed). Using the values

 $V_V$  and  $N_A$ , the numerical density of peroxisomes was determined (Weibel and Bolender, 1973).

## **2.9 Construction of plasmids**

## **2.9.1 Construction of plasmids for epitope tagging**

CFP- and YFP-containing plasmids were constructed in the same manner. Genes encoding fluorescent proteins with different auxotrophic markers that could functionally complement *S. cerevisiae* genes were used (Lys1 of *S. pombe*, Leu-1 of *S. pombe* and Ura-4 of *S. pombe*) to clone into the *SacII* site of pGEMT (Promega) to generate the following plasmids: pCFP-ura, pYFP-ura, pCFP-leu, pYFP-leu, pCFP-lys and pYFP-lys. All the constructs carry a small section of protein A and Ura3. These cassettes were used to PCR from these plasmids with homologous region to the genes of interest and integrated into the genome by homologous recombination.

## **2.9.2 Construction of pNLS-PEX19 for nuclear targeting of Pexl9p**

The plasmid pMadlNLS-GFP constructed in pRS316 with an entire expression cassette under the control of the Pho4 promotor was a gift from Dr. Wozniak's laboratory. Two constructs under the Pho4 promoter containing the expression cassette Madl-NLS and PEX19 (pNLS-PEX19) in-frame or just Madl-NLS, GFP and *PEX19* (pNLS-GFP-PEX19) in-frame were constructed. The original construct was digested with EcoRI and *Notl* to release the GFP and *PEX19* was amplified with similar sites and cloned. For the second construct, the original plasmid was digested with *Sac\* and *Xbal* to open the plasmid, and *PEX19* was ligated downstream of GFP. In this way, although GFP

loses two amino acids along with the stop codon, the expression of GFP was found to be unaffected.

#### **2.9.3 Construction of plasmids for overexpression**

The plasmids pDsRed-PTSl (Smith *et al.,* 2002) and pProtA/HIS5 (Rout *et al.,* 2000) have been described. Genes to be overexpressed were amplified by PCR and cloned into the *BamHI* site of YEp13 (Broach *et al.*, 1979). For ovexpression, the *YHR150w* gene included 492 bp of upstream and 398 bp of downstream sequence, the *YDR479c* gene included 642 bp of upstream and 335 bp of downstream sequence, the *PEX11* gene included 599 bp of upstream and 329 bp of downstream sequence, the *PEX25* gene included 753 bp of upstream and 319 bp of downstream sequence, and the *VPS1* gene included 555 bp of upstream and 324 bp of downstream sequence. For overexpression, the *YLR324w* gene included 609 bp of upstream and 348 bp of downstream sequence, the *YGR004w* gene included 625 bp of upstream and 217 bp of downstream sequence, and the *YBR168w* gene included 592 bp of upstream and 381 bp of downstream sequence.

## **2.10 Construction of a yeast-gene deletion**

Separation of the four ascospores from individual asci by micromanipulation is required for the construction of deletion strains with different auxotrophic markers. Although targeted integrative disruptions of specific genes were used (Longtine *et al.*, 1998), construction of double and triple mutants can be easily achieved by tetrad dissection. Various gene deletion strains were generated by mating, sporulation and tetrad dissection as described by Sherman and Hicks (1991).

### **2.10.1 Mating, sporulation and tetrad dissection**

To generate diploid strains, haploid strains of opposite mating types were streaked in single straight lines on separate YEPD agar plates and incubated overnight at 30°C. They were then replica plated in such a way that streaks of cells of opposite mating types were perpendicular to each other and incubated overnight. Diploids were selected by replica plating these overnight grown colonies onto YND agar supplemented with the auxotrophic requirements of the diploid strain. Diploid cells grow at the intersection of the streaks after overnight incubation. The diploid strain is selected and grown in 5 ml of YND medium supplemented with its auxotrophic requirements. Cells were harvested by centrifugation and washed twice with 10 ml of water. 10  $\mu$ l of cell pellet was added to 3 ml of sporulation medium and incubated for  $3$  to  $7$  days. Formation of tetrads was examined by light microscopy. When approximately 10-15% of the cells formed tetrads, 1 ml o f cells was transferred to a microcentrifuge tube and washed twice with water. The cell pellet was resuspended in 1 ml of water, and 10  $\mu$ l of cells was taken from this for dissection. This 10  $\mu$ l of cells was suspended in 1ml of water containing 3 to 5  $\mu$ g of Zymolyase 20T and incubated at  $30^{\circ}$ C in a rotating wheel for 15 min. 20  $\mu$ l of spheroplasted cells was spread in a single line on one side of a thin YEPD plate. Tetrads were dissected using a Zeiss Axioskop 40 microscope equipped with a Tetrad Manipulator System (Carl Zeiss). Isolated spores were incubated for 2 days at 30°C.

Based on the auxotrophic marker, the spores that have grown will be checked for the required gene deletion by PCR from the genomic DNA.

## **2.10.2 Construction of double deletion mutants**

The homozygous deletion diploid strain *yhrl50wA-HD* (Giaever *et al.,* 2002) was spomlated, and the tetrads were dissected to select for the haploid *MATa* strain. This strain was mated to the haploid *MATa* deletion strain*ydr479A* by replica plating to obtain a heterozygous diploid strain harboring deletions for both *YHR150w* and *YDR479c.* The diploid strain was sporulated, and tetrads from 10 heterozygous diploids were dissected by micromanipulation. All spores were grown in YPD medium, and DNA was extracted. Haploid strains carrying deletions in both the *YDR479c* and *YHRlSOw* genes were selected by PCR analysis. Strains harboring different double deletions of the *YLR324w, YGR004w, YBR168w, PEX28* and *PEX29* genes were constructed in the manner described above for a strain deleted for the *YHRlSOw* and *YDR479c* genes. In this way, ten strains containing the following double gene deletions were constructed: *yhrl50A/ydr479A* (*DKO*), *ylr324A/ygr004A* (.*OKI*), *ylr324A/ybrl68A* (*DK2*), *ygr004A/ybrl 68A* (.*DK3*), *pex28A/ylr324A* (*CD1*), *pex28A/ygr004A* (*CD2), pex28A/ybrl68A* (*CD3*), *pex29A/ylr324A* (*CD4*), *pex29A/ygr004A (CDS),* and *pex29A/ybrl68A* (*CD6*).

## **2.10.3 Construction of a triple deletion mutant**

The gene conferring kanamycin resistance used to disrupt the *YBR168w* gene in the *MATa* strain of *ybrl68A* was replaced with the gene encoding resistance to the drug nourseothricin (Werner BioAgents, Jena, Germany) from the plasmid pHN15 (Krügel *et al.,* 1993). This nourseothricin-resistant *MATa* strain deleted for *YBR168w* was crossed with the *MATa* strain *DK1* (YLR3244/YGR0044), and the resultant diploid was sporulated. The strain *TKO* deleted for the *YBR168w, YLR324w and YGR004w* genes was selected on agar plates containing both kanamycin and nourseothricin, and deletion of the three genes in this strain was confirmed by PCR analysis.

# **2.11 Epitope tagging**

Genes were genomically tagged with the sequence encoding *Staphylococcus aureus* protein A or with the sequence encoding GFP+ by homologous recombination using PCR-based integrative transformation into parental *BY4742* haploid cells (Aitchison *et al.,* 1995; Dilworth *et al.,* 2001). The functionality of fusion proteins was confirmed by the ability of transformed strains to grow and to proliferate peroxisomes like the wildtype strain *BY4742* in medium containing oleic acid as the sole carbon source. A schematic representation of the technique is shown in Figure 2-1. Strains constructed this way are listed in Table 2-4.

## **2.12 Construction of in-frame deletion mutants**

#### **2.12.1 Construction of** *FD30* **and** *FD32* **mutants**

Yeast strains carrying in-frame deletions of the putative Pexl9p binding regions in the gene of interest were constructed by overlapping PCR. This method uses a similar strategy as described in Longtine *et al.,* (1998). Briefly, the *FD30* strain was constructed

as follows. Two individual PCR fragments were initially generated. Using the genomic DNA extracted from Pex30(1-230)p as template, a forward primer upstream of the 20 amino acid of interest and a reverse primer 5 ' CCCTT GGTCCTT ATTT ATT CCT CTT GCT ACTTT AG ACC AT G, which consists of nucleotide sequences before and after (overhang) the 20 amino acids, a first PCR fragment was generated. Thus this fragment does not have the 20 amino acids of interest. A second PCR fragment was generated using the genomic DNA extracted from Pex30p-GFP as template with a forward primer annealing immediately after the 20 amino acids of interest and a reverse primer downstream of the GFP-His5 integration (3'UTR). Thus this second PCR product has an overlapping region with the first PCR fragment. These two products were amplified together to generate a final fragment lacking the 20 amino acids of interest. This final fragment is cloned and sequenced to verify the construct and transformed into the wild type strain. The strain is once again checked by sequencing for correct in-frame deletion. Strain *FD32* was also generated in a similar manner. A schematic representation of the technique is shown in Figure 2-2.

## **2.12.2 Construction of** *S30* **and** *S32* **mutants**

Strains carrying only the amino acid residues of interest tagged with GFP+ were constructed by overlap PCR. Here the genes of interest were cloned in plasmids and were used as template so that the integration cassette could be engineered under the control of the endogenous gene promoter (if genomic DNA were to be used, during the PCR, the homologous promoter region will anneal and the entire region will be amplified). Briefly,



Figure 2-1. Schematic representation of genomic integration for epitope tagging genes of interest. The photograph shows the result of a strain checked for genomic integration.



Figure 2-2. Over lapping PCR strategy. Two different PCRs will be done, one amplifying the gene of interest downstream the domain to be deleted and the second to amplify an auxotrophic marker containing sharing region with the 3'terminator region of the gene of interest. These two PCR products will be used as templates for an overlapping PCR with a forward primer without the domain of interest. The final fragment will be cloned in pGEMT easy vector (Promega) and sequenced. After verification, it will be integrated into the genome of the yeast strain and selected with the His marker.

*S30* strain was constructed as follows. Two individual PCR fragments were initially generated. Using the plasmid with the gene of interest as template, forward primer 0507SG, GTTGCAATTTAGGATTCGAGCTGTCTAGTTGATCCTCCGGAGTGTA AAAACTGATTTTCAATGAGGCTTTTATGGAAGTTTAAG (which consists of 60bps of promoter region, ATG and 21bps of the beginning of the 20 amino acids of interest), and the reverse primer 0508SG, GAAAAGTTCTTCTCCT TTGCTA GCCATACCA AGGTCTAAACCCGTGAC (which consists of an overhanging GFP fragment and the last few bases of the 20 amino acid of interest), a PCR reaction was performed. A second PCR was performed using the pHis5-GFP+ plasmid as template, forward primer 0509SG, TTCTACGTCACGGGTTTAGACCTTGGTATGGCTAGCAAAGGAGAAGAA (which consists of an overhanging region of the last few bases of 20 amino acid of interest and the first 21 bases of GFP+) and a regular reverse primer normally used to make integration constructs. Thus this second PCR product has an overlapping region with the first PCR fragment. These two products were amplified together to generate a final fragment containing the 20 amino acids of interest. This final fragment is cloned and sequenced to verify the construct and transformed into the wild type strain. The strain is once again checked by sequencing for correct in-frame integration. Strains *S32* was also generated in a similar manner. A schematic representation of the technique is shown in Figure 2-2.

## **2.12.3 Construction of** *FDV1* **and** *FDV2* **mutants**

Yeast strains carrying in-frame deletions of the putative Pexl9p binding regions in the gene of interest were constructed by overlap PCR. Briefly, *FDV1* strain was

constructed as follows. Two individual PCR fragments were initially generated. Using the genomic DNA extracted from wild-type strain as template, a forward primer downstream of the amino acids of interest and a reverse primer which consists of nucleotide sequences of the last few bases of Vps1p with the stop codon, a first PCR fragment was generated. A second PCR fragment was generated using the plasmid pProtA/His5 as a template, with a forward primer containing the last few bases of Vps1 gene and a reverse primer downstream of the His5 integration. Thus this second PCR product has an overlapping region with the first PCR fragment. These two products were amplified together to generate a final fragment using a forward primer AGGT AT CCT GCT CT AAG AG AAGCG ATTT CT AAT C AGTTC ATTC AGTTCTT AAA GGATGCTCAAACTTACATCAATACAGCC lacking the amino acids of interest. This final fragment is cloned and sequenced to verify the construct and transformed into the wild type strain. The strain is once again checked by sequencing for correct in-frame deletion. Strain *FDV2* was generated in a similar manner. A schematic representation of the technique is shown in Figure 2-2.

## **2.13 Oligonucleotide directed mutagenesis**

To construct yeast strains with desired point mutations in the gene of interest, standard PCR-based site-directed mutagenesis (Quick change XL Site directed mutagenesis kit, Stratagene, La Jolla, CA) on the plasmid pGBT9 with *PEX30* and *PEX32* cloned individually was used. Manufacturer's instructions were followed to make these constructs, briefly, 50 ng of the plasmid containing the gene and 125 ng of sitedirected mutant primer were used in the mutagenesis reaction, followed by Dpnl

digestion and transformed into E. coli strain. All constructs were sequenced through the coding regions of *PEX30* or *PEX32* or *VPS1*.

# **2.14 CP-Y filter assay**

CP-Y is cotranslationally translocated into the ER, where it gets glycosylated. It is then transported to the Golgi where a sorting signal present in its propeptide form is recognized and then finally sorted to the vacuoles. Defect in this vacuolar sorting, leads to the secretion of CP-Y, through the secretory pathway, outside the cell (Marcusson *et*) *al.,* 1994).The CPY secretion assay was performed according to the method of Roberts *et al.* (1991) with minor modifications. Briefly, cells were grown to an  $OD_{600}$  of 0.5 and spotted on YEPD plates with serial dilution and allowed to dry before filter overlay and incubation. Nitrocellulose sheet was spread on these plates spotted with diluted cells and left to grow for 24 h. The membrane is carefully lifted and rinsed to remove the cells stuck to it. Immunoblotting using the CP-Y antibody is done on the membrane and the secreted protein is detected by immunoblotting.

## **2.15 Two-hybrid Analysis**

#### **2.15.1 Construction of Chimeric genes**

Physical interactions between Pex30p, Pex31p, Pex32p, Pex28p and Pex29p were detected using the Matchmaker Two-Hybrid System (Clontech). Chimeric genes were generated by amplifying the ORFs of the *PEX30. PEX31. PEX32, PEX28* and *PEX29* genes by PCR and ligating them in-frame and downstream of the DNA encoding the transcription-activating domain (AD) and the DNA-binding domain (DB) of the GAL4 transcriptional activator in the plasmids  $pGAD424$  and  $pGBT9$ , respectively. Cells of S. *cerevisiae* strain *SFY526* were transformed simultaneously with a pGAD424-derived plasmid and a pGBT9-derived plasmid.

## **2.15.2 Two-hybrid analysis**

Transformants were grown on SM medium lacking tryptophan and leucine and tested for activation of the integrated *lacZ* construct using a β-galactosidase filter assay (Smith and Rachubinski, 2001). Plasmid pairs encoding AD and DB fusion proteins were transformed into *S. cerevisiae* strain *SFY526.* Transformants were grown in SM medium (Table 2-9). Possible interaction between AD and DB fusion proteins were detected by testing for activation of the integrated *IacZ* construct using  $\beta$ -galactosidase filter and liquid assays according to the instructions of Clontech, with modifications. For filter assays, cells were streaked directly onto filter paper placed on solid media and broken by 4 freeze-thaw cycles at -80 $\degree$ C. Z buffer was used to detect the  $\beta$ -galactosidase activity.

## **2.16 Computer aided analysis**

Major contribution to this study was brought about by computer aided analysis. Protein sequences of *YI*Pex23p and *YIPex24p* were compared to other sequences in the database using the BLAST algorithms [\(http://www.ncbi.nlm.nih.gov/BLAST](http://www.ncbi.nlm.nih.gov/BLAST)) (National Center for Biotechnology Information) and aligned using the ClustalW program (http://www.edi.ac.uk/clustalw/) (EMBL-EBI). This led to the identification of Pex28p, Pex29p, Pex30p, Pex31p and Pex32p.

The promoter regions of *PEX30*, *PEX31* and *PEX32* contain sequences that resemble the canonical sequence  $CCGN_3TNAN_{8-12}CGG$  of the oleic acid response element (ORE) (Rottensteiner *et al*., 2002; 2003a), which acts to increase gene transcription in *S. cerevisiae* in the presence of oleic acid as a carbon source through the binding of the transcription factors Pip2p and Oaf1p (Rottensteiner *et al.*, 1996; Karpichev *et al*., 1997). This was identified by computer analysis. Whether these sequences actually do function as OREs is yet to be determined.

To identify the putative Pexl9p binding site using the consensus sequence developed by Rottensteiner *et al.,* 2004, we used Protein Sub-string Match Analysis (PSMA) software developed in  $C^{++}$  on Linux operating system. It uses pattern matching algorithm from pair wise alignment of multiple sequence alignment (MSA). This matches the consensus motifs which may include unknown or multiple combinations of different factors and extracts from any given sequence. Using this, we could identify any consensus sequence of interest present in any protein of interest.

Also, protein sequences were analyzed using the CBS Prediction Servers [\(http://www.cbs.dtu.dk/services/TMHMM/](http://www.cbs.dtu.dk/services/TMHMM/)) (Center for Biological Sequence Analysis) for identification of transmembrane domains. Apart from this, Yeast Proteome Database (<http://www.proteome.com/>) (Incyte genomics) and the *Saccharomyces* Genome Database ([http://www.veastgenome.org/\)](http://www.veastgenome.org/) (Stanford University) were extensively used for a number of purposes.

# **CHAPTER THREE**

# *PE X28* **AND** *PEX29* **ENCODE PEROXISOMAL INTEGRAL MEMBRANE PROTEINS INVOLVED IN THE REGULATION OF PEROXISOME NUMBER, SIZE, AND DISTRIBUTION**

A portion of this chapter has previously been published as *"YHRlSOw* and *YDR479c* encode peroxisomal integral membrane proteins involved in the regulation of peroxisome number, size, and distribution in *Saccharomyces cerevisiae"* (Franco J. Vizeacoumar, Juan C. Torres-Guzman, Yuen Yi C. Tam. John D. Aitchison, and Richard A. Rachubinski. 2003. *J Cell Biol.* 2003 Apr 28; 161(2):321-32). Reprinted with permission.

## **3.1 Overview**

The peroxin Pex24p of the yeast *Yarrowia lipolytica* exhibits high sequence similarity to two hypothetical proteins, Yhrl50p and Ydr479p, encoded by the *Saccharomyces cerevisiae* genome. Like 7/Pex24p, both Yhrl50p and Ydr479p have been shown to be integral to the peroxisomal membrane, but unlike Y*Pex24p*, their levels of synthesis are not increased upon a shift of cells from glucose- to oleic acid-containing medium. Peroxisomes of cells deleted for either or both of the *YHR150w* and *YDR479c* genes are increased in number, exhibit extensive clustering, are smaller in area than peroxisomes of wild-type cells, and often exhibit membrane thickening between adjacent peroxisomes in a cluster. Peroxisomes isolated from cells deleted for both genes have a decreased buoyant density compared to peroxisomes isolated from wild-type cells and still exhibit clustering and peroxisomal membrane thickening. Overexpression of the genes *PEX25* or *VPS1,* but not the gene *PEX11,* restored the wild-type phenotype to cells deleted for one or both of the *YHR150w* and *YDR479c* genes. Together, our data suggest a role for Yhrl50p and Ydr479p, together with Pex25p and Vpslp, in regulating peroxisome number, size and distribution in *S. cerevisiae.*

## **3.2 Yhrl50p and Ydr479p share extensive similarity with YlPex24p**

7/Pex24p is an integral peroxisomal membrane protein that has been shown to be required for peroxisome assembly in the yeast *Y. lipolytica* (Tam and Rachubinski, 2002). A search of protein databases with the GENEINFO(R)BLAST Network Service of the National Center for Biotechnology Information revealed two proteins encoded by the ORFs *YHR150w* and *YDR479c* of the *S. cerevisiae* genome that exhibit extensive sequence similarity to *Yl*Pex24p (Figure 3-1). The alignment and homology identification was done by Dr. Richard A. Rachubinski.  $YIPex24p$  and Yhr150p exhibit 21.1% amino acid identity and 44.4% amino acid similarity, 7/Pex24p and Ydr479p exhibit 20.8% amino acid sequence identity and 42.1% amino acid similarity, while Yhrl50p and Ydr479p exhibit 20.4% amino acid identity and 41.3% amino acid similarity. YhrI50p is predicted to be a protein of molecular weight 66,145 Daltons and to have two transmembrane helices at amino acid residues 246-268 and 393-415 (http://www.cbs.dtu.dk/services/TMHMM-2.0/) (Krogh et al., 2001). Ydr479p is predicted to be a protein of molecular weight 63,533 Dalton and to have four transmembrane helices at amino acid residues 146-165, 172-194, 265-284 and 291-308. Some potential functional redundancy between Yhrl50p and Ydr479p may have prevented them from being identified as being involved in peroxisome assembly in *S. cerevisiae* by procedures involving random mutagenesis and negative selection for growth of yeast on oleic acid-containing medium.

## **3.3 Synthesis of Yhrl50p and Ydr479p are not induced**

The synthesis of many peroxisomal proteins is induced by the incubation of yeast cells in oleic acid-containing medium. Genomically encoded protein A chimeras of Yhrl50p and Ydr479p were monitored to analyze the expression of *YHRlSOw* and *YDR479c*, respectively, under the control of their endogenous gene promoters. Yeast strains synthesizing Yhrl50p-prA, Ydr479p-prA, thiolase-prA and Pexl7p-prA were





grown in glucose-containing YPD medium and then shifted to oleic acid-containing YPBO medium. Aliquots of cells were removed at various times after the shift to YPBO medium, and their lysates were subjected to SDS-PAGE and immunoblotting (Figure 3- 2). Yhrl50p-prA and Ydr479-prA, as well as Pexl7p-prA (Huhse et al., 1998), were detected in glucose-containing YPD medium at the time of transfer, and their respective levels did not increase with time of incubation of cells in YPBO medium. In contrast, thiolase-prA was barely detectable in cells at the time of transfer to YPBO medium, and its levels were substantially increased with time of incubation in YPBO medium.

## **3.4 Yhrl50p and Ydr479p are integral membrane proteins**

A carboxyl-terminal PTS1 is sufficient to direct a reporter protein to peroxisomes (for a review, see Purdue and Lazarow, 2001). A fluorescent chimera between *Discosoma sp.* red fluorescent protein (DsRed) and the PTS1 Ser-Lys-Leu has been shown to target to peroxisomes of *S. cerevisiae* (Smith et al., 2002). Genomically encoded protein A chimeras of Yhr150p, Ydr479p, the peroxisomal peroxin Pex17p (Huhse et al., 1998) and the mitochondrial translocon protein Tom20p (Lithgow et al., 1994) were localized in oleic acid-induced cells by indirect immunofluorescence microscopy combined with direct fluorescence from DsRed-PTSl to identify peroxisomes (Figure 3-3). Yhrl50pprA, Ydr479p-prA and Pexl7p-prA colocalized with DsRed-PTSl to small punctate structures characteristic of peroxisomes by confocal microcopy. As expected, Tom20pprA did not colocalize with DsRed-PTSl, as the respective individual green and red signals for these proteins remained separate in confocal microscopy.



Figure 3-2. Yhrl50p-prA and Ydr479p-prA remain at constant levels during incubation of *S. cerevisiae* in oleic acid-containing medium. Cells were grown for 16 h in glucosecontaining YPD medium and then transferred to, and incubated in, oleic acid-containing YPBO medium. Aliquots of cells were removed from the YPBO medium at the times indicated, and total cell lysates were prepared. Equal amounts of protein from the total cell lysates were analyzed by SDS-PAGE and immunoblotting to visualize the protein A fusions. Antibodies directed against glucose-6-phosphatase (G6PDH) were used to confirm the loading of equal protein in each lane.



Figure 3-3. Yhrl50p-prA and Ydr479p-prA are peroxisomal proteins by microscopy. The subcellular distributions of protein A chimeras were compared to that of DsRed-PTSl in oleic acid-incubated cells by double labeling, indirect immunofluorescence microscopy. Yhrl50p-prA and Ydr479p-prA colocalize with DsRed-PTSl in punctate structures characteristic of peroxisomes. There is no colocalization of DsRed-PTS1 and the protein A chimera of the mitochondrial protein, Tom20p. Protein A chimeras were detected with rabbit antibodies to mouse IgG and FITC-conjugated goat anti-rabbit IgG secondary antibodies.

Subcellular fractionation and organelle extraction were used to establish if Yhr150p and Ydr479p are associated with peroxisomes and to determine their suborganellar locations. Cells expressing Yhrl50p-prA, Ydr479p-prA and Pexl7p-prA were incubated in oleic acid-containing medium and subjected to subcellular fractionation to yield postnuclear supernatant (PNS) fractions. The PNS fractions were subjected to further centrifugation to yield a supernatant fraction (20KgS) enriched for cytosol and a crude organellar pellet fraction (20KgP). Equal portions of the PNS, 20KgS and the 20KgP were analyzed by immunoblotting. Yhrl50p-prA, Ydr479p-prA and Pexl7p-prA all preferentially localized to the 20KgP fraction (Figure 3-4 A). Peroxisomes were isolated from the 20KgP fractions of each of the strains expressing  $Yhr150p-prA$ ,  $Ydr479p-prA$ and Pex17p-prA. The gradients were fractionated, and equal portions of each fraction were analyzed by immunoblotting (Figure 3-4 B). Yhrl50p-prA and Ydr479p-prA coenriched with the peroxisomal peroxin Pexl7p-prA and not with the mitochondrial protein, Sdh2p. Therefore, both microscopic analysis and subcellular fractionation showed Yhr150p and Ydr479p to be peroxisomal proteins. Some amount of Ydr479pprA was always present in the 20KgS fraction and in the lighter fractions during the gradient isolation of peroxisomes. Whether this represents a selective liberation of a soluble form of Ydr479p-prA during the isolation of peroxisomes remains undetermined. It is unlikely that Ydr479p-prA is found in a cellular compartment in addition to peroxisomes, as immunostaining for Ydr479p-prA yielded exclusively a punctate pattern corresponding to the punctate pattern of peroxisomes defined by DsRed-PTSl (Figure 3- 3). In addition, this punctate compartment does not correspond to mitochondria, as the fluorescence pattern generated with the mitochondria-specific dye MitoTracker did not

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**Figure 3-4** A **and B.** Yhrl50p-prA and Ydr479p-prA are primarily peroxisomal proteins. (A) A PNS fraction was divided by centrifugation into a supernatant (20KgS) fraction enriched for cytosol and a pellet (20KgP) fraction enriched for peroxisomes and mitochondria. Equivalent portions of each fraction were analyzed. Immunoblotting detected the protein A chimeras shown, including that of the peroxisomal protein Pexl7p. (B) Yhrl50p-prA and Ydr479p-prA cofractionate with peroxisomes. Organelles in the 20KgP fraction were separated by isopycnic centrifugation on a discontinuous Nycodenz gradient. Fractions were collected from the bottom of the gradient, and equal portions of each fraction were analyzed by immunoblotting. Fractions enriched for peroxisomes and mitochondria were identified by immunodetection of the protein A chimera of  $Pex17p$ and Sdh2p, respectively.



Figure 3-4 C. Yhrl50p-prA and Ydr479p-prA are not mitochondrial proteins by microscopy. The subcellular distributions of protein A chimeras were compared to that of fixable dye Mitotracker in oleic acid-incubated cells by double labeling, indirect immunofluorescence microscopy. Tom20p-prA colocalize with Mitotracker in elongated tubular structures characteristic of mitochondria. Protein A chimeras were detected with rabbit antibodies to mouse IgG and FITC-conjugated goat anti-rabbit IgG secondary antibodies.

overlap with the punctate pattern generated by detection of Ydr479p-prA by immunofluorescence (Figure 3-4 C).Peroxisomes were hypotonically lysed by incubation in dilute alkali Tris buffer and subjected to centrifugation to yield a supernatant (Ti8S) enriched for matrix proteins and a pellet (Ti8P) enriched for membrane proteins (Figure 3-4 D). The chimeras of Yhrl50p and Ydr479p localized primarily to the Ti8P fraction, as did the chimeras of the peripheral peroxisomal membrane protein Pex17p (Huhse et al., 1998) and the integral peroxisomal membrane protein Pex3p (Hohfeld et al., 1991). The soluble peroxisomal matrix protein thiolase was found almost exclusively in the Ti8S fraction. The reproducible presence of some Ydr479p in the Ti8S fraction may again be representative of a soluble form of this protein that is selectively liberated during the isolation of peroxisomes (see Figure 3-4 A, B). The Ti8P fractions were then extracted with alkali sodium carbonate and subjected to centrifugation (Figure 3- 4 D). This treatment releases proteins associated with, but not integral to, membranes (Fujiki et al., 1982). Under these conditions, Yhrl50p-prA and Ydr479p-prA fractionated with Pex3pprA to the pellet fraction enriched for integral membrane proteins, while Pexl7p-prA fractionated to the supernatant fraction enriched for soluble proteins, including peripheral membrane proteins. These data suggest that Yhrl50p and Ydr479p are both primarily integral peroxisomal membrane proteins, as has been shown for F/Pex24p (Tam and Rachubinski, 2002).


**Figure 3-4 D.** Yhrl50p-prA and Ydr479p-prA are integral peroxisomal membrane proteins. Peroxisomes purified by isopycnic density gradient centrifugation were lysed by treatment with 10 mM Tris-HCl, pH 8.0, releasing matrix proteins to a supernatant fraction (Ti8S) after centrifugation. The membrane-containing pellet fraction (Ti8P) was treated with  $0.1$  M Na<sub>2</sub>CO<sub>3</sub>, pH 11.3, and then subjected to centrifugation to yield a supernatant fraction (C03S) enriched for peripherally associated membrane proteins and a pellet fraction (CO3P) enriched for integral membrane proteins. Equal portions of the respective supernatant and pellet fractions were analyzed by immunoblotting. Immunodetection of thiolase, Pexl7p-prA and Pex3p-prA marked the fractionation profiles of a matrix, peripheral membrane and integral membrane protein, respectively.

# **3.5 Abnormal peroxisomes are found in both single and double mutants of yhrl50w and ydr479c**

We next investigated the functionality of peroxisomes in the deletion mutants of these genes. Dr. Juan Carlos Guzman Torres constructed the double deletion mutant and checked for their growth defect. Both the single and double deletion mutants grew like the wild-type cells on oleic acid-containing medium (Figure 3-4 E). Import of matrix proteins in these mutants were unaffected (Figure 3-4 F). We next investigated the ultrastructure of cells incubated in oleic acid-containing YPBO medium by transmission EM. Wild-type cells (Figure 3- 5 A) consistently showed individual peroxisomes well separated from one another. In contrast, cells of the *yhr150* $\Delta$  (Figure 3-5 B), *ydr479* $\Delta$ (Figure 3-5 C) and, particularly, the *yhrl50A/ydr479A* (Figure 3-5 D) strains contained peroxisomes that exhibited clustering. 28.0%, 19.2% and 20.4% of peroxisomes of cells o f the *yhrl50A/ydr479A, ydr479A* and *yhrl50A* strains, respectively, showed clustering, in contrast to 4.0% of peroxisomes of wild-type cells - a cluster of peroxisomes was operationally defined as 3 or more adherant peroxisomes (Figure 3-5 F and G). The clustered peroxisomes often showed evidence of membrane thickening between adjacent peroxisomes in the cluster. Morphometric analysis showed that cells of the deletion strains contained a greater number of peroxisomes than did wild-type cells and that, on average, these peroxisomes were smaller in size than those of wild-type cells (Table 3-1). Cells of the deletion strains contained much greater numbers of peroxisomes with areas of 0.02  $\mu$ m<sup>2</sup> or less than did wild-type cells (Figure 3-5 E). Nycodenz density gradient centrifugation analysis showed that peroxisomes purified from *yhrl50A/ydr479A* cells



Figure 3-4 E. Growth of different strains on YPBO medium. All strains were spotted after serial dilution and grown over a period of 4 days at 30C. This figure was contributed by Dr.Juan carlos Guzman Torres.



**Figure 3-4 F.** Distribution of matrix proteins in the mutants. Subcellular localization of matrix proteins were analysed by differential fractionation. A PNS fraction was divided by centrifugation into a supernatant (20KgS) fraction enriched for cytosol and a pellet (20KgP) fraction enriched for peroxisomes and mitochondria. Equivalent portions of each fraction were analyzed.



Figure 3-5. Peroxisomes are smaller, more abundant and exhibit clustering in cells deleted for either or both of the YHR150w and YDR479c genes. Ultrastructure of wildtype *BY4742* (A), *yhrl50A* (B), *ydr479A* (C) and *y h rl50A/ydr479A* (D) cells. Cells were grown in YPD medium overnight, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for EM. P, peroxisome. P\*, peroxisome cluster. Bar, 0.5 pm. The double deletion mutant was constructed by Dr.Juan Carlos Torres Guzman.



**Figure 3-5 E.** Morphometric analysis of peroxisomes. Oleic acid-incubated wild-type *(WT) BY4742* and deletion mutant cells were analyzed. For each strain analyzed, electron micrographs of 50 randomly selected cells at  $\times$  17,000 magnification were scanned, and the areas of individual peroxisomes were determined by counting the number of individual pixels in a peroxisome with Image Tools for Windows, Version 2.00. The peroxisomes were then separated into size categories. A histogram was generated for each strain depicting the percentage of total peroxisomes occupied by the peroxisomes of each category. The numbers along the x-axis are the maximum sizes of peroxisomes in each category (in  $\mu$ m<sup>2</sup>).



Figure 3-5  $F$  and G. Analysis of the clustering of the peroxisomes. (F) The number of cells exhibiting clusters were counted and represented graphically. (G) The number of different peroxisomal clusters that were present in the wild type and various mutants quantitated were represented graphically.

G

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<b>Strain</b>	Cell area assayed $(\mu m^2)$	Peroxisome count <sup>a</sup>	<b>Numerical</b> density of peroxisomes <sup>b</sup>	Average area of peroxisomes <sup>c</sup> $(\mu m^2)$	
WT(BY4742)	404	0.37	1.71	0.055	
$yhr150\Delta$	424	0.51	2.59	0.048	
$ydr$ 479 $\varDelta$	382	0.56	3.20	0.039	
yhr150A/ydr479A	381	0.80	4.38	0.040	

Table 3-1. Average area and numerical density of peroxisomes in wild-type and deletion strains

"Number of peroxisomes counted per  $\mu$ m<sup>2</sup> of cell area on micrographs.

 $^{\circ}$ Number of peroxisomes per  $\mu$ m<sup>3</sup> of cell volume (Weibel and Bolender, 1973).

"Average area on micrographs.

have a greatly reduced density (peak fraction 8, 1.19  $g/cm<sup>3</sup>$ ) compared to peroxisomes from wild-type cells (peak fraction 1, 1.22  $g/cm^3$ ) (Figure 3-6 A). Peroxisomes isolated from cells deleted for *YHRlSOw* or *YDR479c* were also less dense than wild-type peroxisomes, although the differences in density were less than that observed between peroxisomes *from yhr 15OA/ydr479A* cells and wild-type peroxisomes (Figure 3-6 A). EM analysis showed that the peroxisomes purified from *yhr 15OA/ydr479A* cells still exhibited clustering and evidence of thickened peroxisomal membranes, while peroxisomes purified from wild-type cells were largely well separated from one another, with no evidence of membrane thickening (Figure 3-6 B).

### **3.6 Overexpression of** *PEX11***,** *PEX25* **and** *VPS1*

Since cells deleted for one or both of the *YHR150w* and *YDR479c* genes are compromised in their regulation of peroxisome number, size and distribution, we investigated the effects of overexpression of three genes previously shown to be involved in these processes in *S. cerevisiae.* Cells of strains mutant for the genes encoding the peroxisomal peroxins Pex l 1 p (Erdmann and Blobel, 1995; Marshall et al., 1995, 1996), Pex25p (Smith et al., 2002) and the dynamin-like protein Vps1p (Hoepfner et al., 2001) have reduced numbers of enlarged peroxisomes compared to wild-type cells (Figure 3-13). Overexpression of *PEX25* (Figure 3-7 and Figure 3-11) or *VPS1* (Figure 3-7 and Figure 3-12) in cells deleted for one or both of the *YHR150w* and *YDR479c* genes led to a partial restoration of the wild-type peroxisomal phenotype, with overexpressing cells containing increased numbers of separate, individual peroxisomes and reduced numbers of peroxisomal clusters. In contrast, overexpression of *PEX11* did not appear to have any



\_ -- — - —— - *—^mmtl-Thiott&i:* 1 2 .1 4 > 6 N V to t | 12 1.1 14 1? *y h r l5 0 J /* and the mutant strain *<sup>97</sup>* **I 2 3** 4 **N ft N «> 10** 1 **I** *12 IX* **14 I>** *y d r 4 7 9 J* — . - — *a-Ihioltzst' yhrl50A/ydr479A* were 1 2 .1 4 \* ft *"* X *\*\** 1(1 11 12 13 14 !?•  $\mu$ -*SDH* 

Figure 3-6. Peroxisomes isolated from *yhrlSOA* dense than isolated wildtype peroxisomes and retain a clustered  $a\text{-}SKL$  phenotype. (A) The wild-——*Z.* - type ( *WT)* strain *BY4742* grown overnight in YPD medium, transferred to oleic acid-containing YPBO medium and

incubated in YPBO medium for 8 h. A PNS fraction was prepared from cells of each strain and divided by centrifugation into 20KgS and 20KgP fractions. Organelles in the 20KgP fraction were separated by isopycnic centrifugation on a continuous 30% to 45% Nycodenz gradient. Fractions were collected from the bottom of the gradient, and equal portions of each fraction were analyzed by immunoblotting with antibodies to the PTS1, Ser-Lys-Leu, to detect peroxisomes.

<span id="page-118-0"></span>

Figure 3-6 B. Electron micrographs of purified peroxisomes. Peak peroxisomal fractions from cells of the wild-type strain  $(WT)$  BY4742 (Fraction 1) and the mutant strain *yhr 150Δ/ydr 479Δ* (Fraction 8). Bar, 0.5 μm.



Figure 3-7. Peroxisome morphology in cells of gene overexpression strains. Cells were grown in SM medium overnight, transferred to YPBO medium and incubated in YPBO medium for 8 h. Peroxisomes were detected by double labeling, indirect immunofluorescence microscopy with antibodies to the PTS1 Ser-Lys-Leu (SKL) and FITC-conjugated goat anti-rabbit IgG secondary antibodies, and with guinea pig antibodies to the PTS2-containing protein thiolase and rhodamine-conjugated donkey anti-guinea pig IgG secondary antibodies. The genetic backgrounds of the different yeast strains are given at *Top*, and the genes that are overexpressed are denoted at *Left.*

effect on the abnormal peroxisome morphology observed in cells mutant for the *YHR150w* and *YDR479c* genes (Figure 3-7, Figure 3-10). It should be noted that overexpression of the *YHR150w* gene led to restoration of wild-type peroxisome morphology in *yhrl50A* cells but not in *ydr479A* cells or in cells deleted for both the *YHRI50w* and *YDR479c* genes (Figure 3-7, Figure 3-8). In contrast, overexpression of *YDR479c* led to restoration of wild-type peroxisome morphology not only in *vdr479* cells, but also in *yhrl50A* cells and cells deleted for both genes (Figure 3-7, Figure 3-9). Therefore, it appears that Yhrl50p may be in large part redundant in its functions in regulating peroxisome dynamics vis-a-vis Ydr479p. Attempts at demonstrating physical interactions, either direct or indirect, between Yhrl50p and Ydr479p or between these two proteins and Pexl 1p, Pex25p or Vpslp were unsuccessful.

As reported previously (Erdmann and Blobel, 1995; Marshall et al., 1995, 1996; Hoepfher et al., 2001; Smith et al., 2002), cells deleted individually for the *PEX11*, *PEX25* and *VPS1* genes contained reduced numbers of enlarged peroxisomes (Figure 3-7 and Figure 3-13). Overexpression of the *PEX11* gene in wild-type cells or in cells deleted for *PEX11*, *PEX25* or *VPS1* led to a proliferation of smaller peroxisomes (Figure 3-7 and Figure 3-10). However, overexpression of *PEX25* or *VPS1*, while resulting in normal peroxisome morphology in their respective mutant backgrounds, did not affect the peroxisome morphology in the opposite mutant background (Figure 3-7, Figure 3-11 and Figure 3-12). Overexpression of *PEX25*, but not *VPS1*, led to only a nominal increase in the number of normal peroxisomes in cells deleted for the *PEX11* gene (Figure 3-7 and Figure 3-11). Moreover, the  $p \in x / l \leq l$  cells retained their enlarged peroxisomal phenotype when either *YHR150w* or *YDR479c* was overexpressed (Figure 3-7, Figure 3-8 and Figure

**9 9**

#### Table 3-2. Summary of results of gene overexpression



 ${}^{3}$ The (+) symbol denotes the presence of a particular peroxisomal morphological phenotype. Increased numbers of (+) symbols denote increased prevalence of a particular peroxisomal morphological phenotype. The absence of a (+) symbol denotes the absence of particular peroxisomal morphological phenotype.



Figure 3-8. Peroxisome morphology in cells of gene overexpression strains

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## **Overexpression of** *YDR479c*



Figure 3-9. Peroxisome morphology in cells of gene overexpression strains



Figure 3-10. Peroxisome morphology in cells of gene overexpression strains



Figure 3-11. Peroxisome morphology in cells of gene overexpression strains

## **Overexpression of** *VPSJ*



**Figure 3-** 12. Peroxisome morphology in cells of gene overexpression strains



# **Deletion Strains**

Figure 3-13. Peroxisome morphology in deletion mutants

3-9). Our results suggest that Pex11p plays a central role in regulating peroxisome division. The results of gene overexpression studies are summarized in Table 3-2.

## **3.7 Discussion**

#### **3.7.1 Database mining as a tool to identify novel** *PEX* **genes**

Completion of the *S. cerevisiae* genome sequencing project proved invaluable for the identification of a novel *PEX* gene, *PEX25*, by transcriptome profiling of cells grown in oleic acid-containing medium versus cells grown in glucose-containing medium (Smith et al., 2002). The completed *S. cerevisiae* genome sequence is added value in that it provides the opportunity to identify novel proteins required for peroxisome biogenesis in *S. cerevisiae* through sequence similarity between proteins of unknown function encoded by the *S. cerevisiae* and proteins already shown to be required for peroxisome biogenesis in other organisms. The requirement of the peroxisomal integral membrane protein Pex24p for peroxisome assembly in the yeast *Y. lipolytica* was recently demonstrated (Tam and Rachubinski, 2002). Y/Pex24p was shown to share extensive sequence similarity to two proteins of unknown function and unknown localization encoded by the ORFs *YHR150w* and *YDR479c* of the *S. cerevisiae* genome. In this manuscript, genomically encoded protein A chimeras of Yhr150p and Ydr479p were shown by a combination of microscopic and subcellular fractionation analyses to be peroxisomal proteins. In their response to extraction by different salts, Yhr150p and Ydr479p act primarily as integral membrane proteins. However, a fraction of Ydr479p consistently acts as a soluble protein. The exact nature of this soluble form of Ydr479p and its origin (Is it in equilibrium with the peroxisomal membrane form of Ydr479p?) remain under investigation.

#### **3.7.2 A role for the novel genes in peroxisome biogenesis**

The proteins encoded by the *YHRlSOw* and *YDR479c* genes are not required for peroxisome assembly, as cells harboring deletions for one or both of these genes still contain peroxisomes. These peroxisomes were functional, at least to some degree, as the cells containing one or both of the gene deletions were able to grow in oleic acidcontaining medium with essentially the same kinetics as the wild-type strain (Figure 3-4 E). However, the peroxisomes in the deleted strain are not normal and show phenotypic characteristics distinct from those of wild-type peroxisomes. The peroxisomes of cells deleted for one or both of the *YHR150w* and *YDR479c* genes are more abundant and smaller, and show extensive clustering, as compared to wild-type peroxisomes. In addition, the membranes of the clustered peroxisomes of the gene deletion strains are often thickened in appearance. These characteristics of peroxisomes of the deletion strains are consistent with a role for *YHR150w* and *YDR479c* in the control of peroxisome size, number and distribution within cells. However, it does not appear that *YHRlSOw* and *YDR479c* are required for peroxisome inheritance per se, as all cells deleted for one or both of these genes still contained peroxisomes after numerous cell divisions. Also, if *YHR150w* or *YDR479c* had a direct role in the inheritance of peroxisomes, one might expect that a loss of peroxisomes from cells over time resulting from impaired segregation of peroxisomes into daughter cells would lead to decreased kinetics of growth in oleic acid-containing medium for the deletion strains vis-a-vis the wild-type strain, which, as reported above, was not observed. It is interesting to note that *Y.*

*lipolytica* cells deleted for the *PEX24* gene also show evidence of abnormal peroxisomal divisional control. These cells lack mature peroxisomes but do accumulate membrane structures that contain both peroxisomal matrix and membrane proteins (Tam and Rachubinski, 2002). However, these membrane structures are not functional peroxisomes in *Y. lipolytica*, as *pex24A* cells cannot grow on medium containing oleic acid as the sole carbon source. Therefore, although T/Pex24p, like Yhrl50p and Ydr479p, most likely has a role in the regulation of peroxisome division, Y/Pex24p probably does not function identically to Yhrl50p or Ydr479p or is modulated in its actions differently than are Yhrl50p and Ydr479p.

#### **3.7.3 Regulation of peroxisome division**

The size, number and distribution of peroxisomes are tightly controlled by the cell. Loss of the enzymatic activities of individual peroxisomal  $\beta$ -oxidation enzymes has been shown to result in pronounced changes in peroxisome size and/or number (Fan et al., 1998; Chang et al., 1999; Smith et al., 2000; van Roermund et al., 2000), due primarily to the increased levels of the remaining peroxisomal  $\beta$ -oxidation enzymes. The molecular mechanisms underlying this so called metabolic control of peroxisome abundance (Chang et al., 1999) remain essentially unknown.

In contrast, members of the Pexl 1 family of peroxins have been implicated as effectors of peroxisome division in multiple species (Erdmann and Blobel, 1995; Marshall et al., 1995, 1996; Sakai et al., 1995; Abe and Fujiki, 1998; Lorenz et al., 1998; Passreiter et al., 1998; Schrader et al., 1998; Li and Gould, 2002). The recently reported *PEX25* gene has also been implicated in the regulation of peroxisome size and number in

*S. cerevisiae* (Smith et al., 2002), as has the dynamin-like protein Vpslp (Hoepfher et al., 2001). Like other dynamin-related proteins, Vpslp was proposed to be involved in a membrane fission event required for the regulation of peroxisome size and abundance. The thickened membranes between some peroxisomes of a peroxisome cluster seen in cell deleted for the *YHR150w* and/or *YDR479c* genes are also suggestive of a role for Yhr150p and Ydr479p in controlling fission of the peroxisomal membrane.

#### **3.7.4 Mechanism of peroxisome division**

How might Yhr150p, Ydr479p, Pex11p, Pex25p and Vps1p act and interact to control the abundance, size and distribution of peroxisomes in the *S. cerevisiae* cell? We sought to get some insight into this question by determining the effects on peroxisome morphology of overexpressing the genes for these proteins in wild-type cells and cells deleted for the different genes.

Overexpression of the  $PEX11$  gene in the  $pex11\Delta$  genetic background has been reported to result in large numbers of small peroxisomes (Marshall et al., 1995). In contrast, overexpression o f the *PEX25, VPS1, YHR150w* and *YDR479c* in their respective gene deletion backgrounds does not result in the production of large numbers of small peroxisomes but instead restores the wild-type peroxisomal phenotype. Considering the proliferation of peroxisomes as a two step pathway, namely division of peroxisomes and separation of peroxisomes, overexpression of the *PEX11* gene either in wild-type cells or in cells of the various deletion strains leads to a significant proliferation of peroxisomes, which remain for the most part adherant to one another. Thus, Pex 1 1 p plays a central and positive regulatory role in the division step of peroxisome proliferation but has little or no readily apparent role in the separation step of the process. The presence of reduced numbers of enlarged peroxisomes in *pex25* $\Delta$  cells (Smith et al., 2002; Figure 3-13) and *vps14* cells (Hoepfner et al., 2001; Figure 3-13) suggests that Pex25p and Vps1p function in addition to Pex11p in the divisional step of peroxisome proliferation.

Upon completion of peroxisome division, peroxisomes must be separated from one another. YhrlSOp and Ydr479p are two proteins required for this process, as their absence leads to an arrest or retardation of the peroxisome proliferation pathway, leading to the presence of clusters of peroxisomes with evidence of thickened membranes sometimes occurring between adjacent peroxisomes. Significant recovery of the wildtype peroxisomal phenotype by overexpression of *PEX25* or *VPS1* in cells deleted for one or both of the *YHR150w* or *YDR479c* genes implies that Pex25p and Vps1p have roles in the separation of peroxisomes in addition to their roles in peroxisome division discussed above. In contrast, overexpression of *PEX11* in cells deleted for one or both of the *YHR150w* or *YDR479c* genes did not result in the reappearance of wild-type peroxisomes, and peroxisomes remained clustered and sometimes exhibited membrane thickening between adjacent peroxisomes, as in the original strains deleted for *YHRlSOw*and/or *YDR479c*. Therefore, Pex11p appears to function primarily or only at the divisional step of peroxisome proliferation but not at the separation step.

Organelles are highly dynamic structures that undergo fission and fusion processes to allow cells to respond to intracellular and extracellular cues and to allow for their correct segregation at cell division. The maintenance of compartmental integrity in the eukaryotic cell therefore requires tight control mechanisms for these events. In the control of peroxisome number, size and distribution, our data suggest that Pexl Ip plays a pre-eminent role in controlling peroxisome division, while Pex25p, Vpslp and the newly identified peroxisomal proteins Yhrl50p and Ydr479p all play a prominent role in controlling the separation of peroxisomes from one another. The challenge for the future lies in understanding further the interplay amongst these proteins and the signaling events they respond to and initiate in order to control peroxisomal dynamics in the cell.

## **CHAPTER FOUR**

# *PEX30, PE X 31***, AND** *PEX32* **ENCODE A FAMILY OF PEROXISOMAL INTEGRAL MEMBRANE PROTEINS REGULATING PEROXISOME SIZE AND NUMBER**

A portion of this chapter has previously been published as "Pex30p, Pex31p. and Pex32p form a family of peroxisomal integral membrane proteins regulating peroxisome size and number in *Saccharomyces cerevisiae"* (Franco J. Vizeacoumar, Juan C. Torres-Guzman, David Bouard, John D. Aitchison. and Richard A. Rachubinski. 2003. *Mol Biol. Cell.* 2004 Feb; 15(2):665-77. Reprinted with permission

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### **4.1 Overview**

The peroxin Pex23p of the yeast *Yarrowia lipolytica* exhibits high sequence similarity to the hypothetical proteins Ylr324p, Ygr004p and Ybrl68p encoded by the *Saccharomyces cerevisiae* genome. Ylr324p, Ygr004p and Ybrl68p are integral to the peroxisomal membrane and act to control peroxisome number and size. Synthesis of Ylr324p and Ybrl68p, but not of Ygr004p, is induced during incubation of cells in oleic acid-containing medium, the metabolism of which requires intact peroxisomes. Cells deleted for *YLR324w* exhibit increased numbers of peroxisomes, while ceils deleted for *YGR004w* or *YBR168w* exhibit enlarged peroxisomes. Ylr324p and Ybrl68p cannot functionally substitute for one another or for Ygr004p, while Ygr004p shows partial functional redundancy with Ylr324p and Ybrl68p. Ylr324p, Ygr004p and Ybrl68p interact within themselves and with Pex28p and Pex29p, which have been shown also to regulate peroxisome size and number. Systematic deletion of genes demonstrated that *PEX28* and *PEX29* function upstream of *YLR324w, YGR004w* and *YBR168w* in the regulation of peroxisome proliferation. Our data suggest a role for Ylr324p, Ygr004p and Ybr168p – now designated Pex30p, Pex31p and Pex32p, respectively – together with Pex28p and Pex29p in controlling peroxisome size and proliferation in *S. cerevisiae.*

# **4.2 Ylr324p, Ygr004p and Ybrl68p share extensive sequence similarity with YIPex23p**

T/Pex23p is an integral peroxisomal membrane protein that has been shown to be required for peroxisome assembly in the yeast *Y. lipolytica* (Brown *et al.,* 2000). A search of protein databases with the GENEINFO(R)BLAST Network Service of the National Center for Biotechnology Information revealed three proteins encoded by the ORFs *YLR324w, YGR004w* and *YBR168w* of the *S. cerevisiae* genome that exhibit extensive sequence similarity to YIPex23p (Figure 4-1). YIPex23p and Ylr324p exhibit 36.3% amino acid sequence identity and 20.9% amino acid similarity (at positions of nonidentity), y/Pex23p and Ygr004p exhibit 35.6% amino acid sequence identity and 19.3% amino acid similarity, F/Pex23p and Ybrl68p exhibit 17.6% amino acid identity and 24.4% amino acid similarity, while Ylr324p, Ygr004p and Ybrl68p exhibit 9.3% amino acid identity and 20.7% amino acid similarity among themselves. Ylr324p is predicted to be a protein of molecular weight 59,461 and to have two transmembrane helices at amino acids 3-22 and 191-209 [\(http://www.cbs.dtu.dk/services/TMHMM-2.0/\)](http://www.cbs.dtu.dk/services/TMHMM-2.0/) (Krogh *et al*., 2001). Ygr004p is predicted to be a protein of molecular weight 52,942 and to have four transmembrane helices at amino acids 93-111, 110-129, 178-195 and 226-244. Ybrl68p is predicted to be a protein of molecular weight 48,578 and to have six transmembrane helices at amino acids 45-62, 69-86, 102-121, 178-197, 205-224 and 230-249. Partial functional redundancy among Ylr324p, Ygr004p and Ybrl68p (discussed below) may have prevented them from being identified as being involved in peroxisome biogenesis in *S. cerevisiae* by procedures involving random mutagenesis and negative selection for growth of yeast on oleic acid-containing medium.

#### **4.3 Synthesis of Ylr324p andYbrl68p are inducible**

The culturing of yeast cells in oleic acid-containing medium elicits a dual cellular response in that it promotes the proliferation of peroxisomes and induces the expression



Figure 4-1. Sequence alignment of *Yarrowia lipolytica* Pex23p with the proteins Ylr324p, Ygr004p and Ybrl68p encoded by the *Saccharomyces cerevisiae* genome. Amino acid sequences were aligned with the use of the ClustalW program (EMBL-European Bioinformatics Institute, <http://www.ebi.ac.uk/clustalw/>). Identical residues in at least three of the proteins are shaded black, while identical residues in two proteins are shaded blue. Similar residues are shaded gray. Similarity rules:  $G = A = S$ ;  $A = V$ ;  $V = I$  $= L = M$ ;  $I = L = M = F = Y = W$ ;  $K = R = H$ ;  $D = E = Q = N$ ; and  $S = T = Q = N$ . This figure was aligned by Dr. Richard A. Rachubinski.

of many genes encoding peroxisomal proteins. Genomically encoded protein A chimeras of Ylr324p, Ygr004p and Ybr168p were monitored to analyze the expression of *YLR324w, YGR004w* and *YBR168w*, respectively, under the control of their endogenous gene promoters. Yeast strains synthesizing Ylr324p-prA, Ygr004p-prA and Ybr168p-prA were grown in glucose-containing YPD medium and then shifted to oleic acid-containing YPBO medium. Aliquots of cells were removed at various times after the shift to YPBO medium, and their lysates were analyzed by SDS-PAGE and immunoblotting (Figure 4- 2). Ylr324p-prA, Ygr004p-prA and Ybr168p-prA were all detected in glucose-containing YPD medium at the time of transfer. The levels of Ylr324p-prA and Ybr168p-prA increased with time of incubation of cells in YPBO medium, but not as dramatically as the levels of Potlp (peroxisomal thiolase). The levels of Ygr004p-prA did not show any apparent increase with time of incubation of cells in YPBO medium. It is noteworthy that the promoter regions of *YLR324w*, *YGR004w* and *YBR168w* contain sequences that resemble the canonical sequence  $CCGN_3TNAN_{8-12}CGG$  of the oleic acid response element (ORE) (Table 4-1) (Rottensteiner *et al.,* 2002; 2003a), which acts to increase gene transcription in *S. cerevisiae* in the presence of oleic acid as a carbon source through the binding of the transcription factors Pip2p and Oaflp (Rottensteiner *et al.*, 1996; Karpichev *et al.,* 1997). Whether these sequences actually do function as OREs remains to be determined.



Figure 4-2. The levels of Ylr324p-prA and Ybr168-prA, but not of Ygr004p-prA, are increased during incubation of *S. cerevisiae* in oleic acid-containing medium. Cells were grown for 16 h in glucose-containing YPD medium and then transferred to, and incubated in, oleic acid-containing YPBO medium. Aliquots of cells were removed from the YPBO medium at the times indicated, and total cell lysates were prepared. Equal amounts of protein from the total cell lysates were analyzed by SDS-PAGE and immunoblotting to visualize the protein A fusions and Potlp. Antibodies directed against glucose-<sup>6</sup> phosphatase (G6PDH) were used to confirm the loading of equal protein in each lane.

Gene	<b>ORE Canonical Sequence</b> $(CCGN3TNAN8-12CGG)$	
PEX7	$CAGN_{10}TNAN_3CCG$	
PEX13	$CGGN_{12}TNAN_3CCG$	
YLR324w	CGGN <sub>12</sub> TNAN <sub>3</sub> CCG	
YGR004w	$CGGN_{13}TNAN_{8}CCG$	
YBR168w	CAGN <sub>2</sub> TNAN <sub>13</sub> CCG	

Table 4-1. Putative OREs in the promoter regions of the *YLR324w*, *YGR004w* and *YBR168w* genes

## **4.4 Ylr324p, Ybrl68p and Ygr004p are integral membrane proteins**

A carboxyl-terminal PTS1 is sufficient to direct a reporter protein to peroxisomes. A fluorescent chimera between *Discosoma sp.* red fluorescent protein (DsRed) and the PTS1 Ser-Lys-Leu has been shown to target to peroxisomes of S. *cerevisiae* (Wang *et a l*, 2001; Smith *et al.,* 2002). Genomically encoded protein A chimeras of Ylr324p, Ygr004p, Ybr168p and the peroxisomal matrix protein Pot1p were localized in oleic acid-induced cells by indirect immunofluorescence microscopy combined with direct fluorescence of DsRed-PTSl to identify peroxisomes (Figure 4-3A). Ylr324p-prA, Ygr004p-prA, Ybr168p-prA and Potlp colocalized with DsRed-PTS1 to small punctate structures characteristic of peroxisomes by confocal microcopy.

Subcellular fractionation and organelle extraction were used to establish if Ylr324p, Ygr004p and Ybr168p are associated with peroxisomes and to determine their suborganellar locations. Ylr324p-prA, Ygr004p-prA and Ybr168p-pA, like Potlp, preferentially localized to the 20KgP fraction enriched for peroxisomes (Figure 4-3B). Peroxisomes were isolated from the 20KgP fractions of each of the strains expressing Ylr324p-prA, Ygr004p-prA or Ybr168p-prA. The gradients were fractionated, and equal portions of each fraction were analyzed by immunoblotting (Figure 4-3C). Ylr324p-prA. Ygr004p-prA and Ybr168p-prA coenriched with the peroxisomal matrix protein thiolase (Potlp) and not with the mitochondrial protein, Sdh2p. Therefore, both microscopic analysis and subcellular fractionation showed Ylr324p, Ygr004p and Ybr168p to be peroxisomal proteins. Some amount of Ylr324p-prA was always present in the lighter fractions during the gradient isolation of peroxisomes, while Ygr004-prA consistently enriched in fraction 9 at a density lighter than that of peroxisomes. Whether there is a



Figure 4-3 A. Ylr324p-prA, Ygr004p-prA and Ybr168p-prA are peroxisomal proteins by confocal microscopy. (A) The subcellular distributions of protein A chimeras were compared to that of DsRed-PTS1 in oleic acid-incubated cells by double labeling, indirect immunofluorescence microscopy. Ylr324p-prA, Ygr004p-prA and Ybr168-prA, along with the peroxisomal matrix protein Potlp, colocalize with DsRed-PTSl in punctate structures characteristic of peroxisomes. Protein A chimeras were detected with rabbit antibodies to mouse IgG and FITC-conjugated goat anti-rabbit IgG secondary antibodies.



Figure 4-3 B and C. Ylr324p-prA, Ygr004p-prA and Ybr168p-prA are peroxisomal proteins by sub cellular fractionation. (B) A PNS fraction was divided by centrifugation into a supernatant (20KgS) fraction enriched for cytosol and a pellet (20KgP) fraction enriched for peroxisomes. Equivalent portions of each fraction were analyzed. Immunoblotting with rabbit anti-Potlp antibodies detected both the protein A chimeras shown and Potlp. (C) Ylr324p-prA, Ygr004p-prA and Ybr168-prA cofractionate with peroxisomes. Organelles in the 20KgP fraction were separated by isopycnic centrifugation on a discontinuous Nycodenz gradient. Fractions were collected from the bottom of the gradient, and equal portions of each fraction were analyzed by immunoblotting. Fractions enriched for peroxisomes and mitochondria were identified by immunodetection of Potlp and Sdh2p, respectively.
selective liberation of a soluble form of Ylr324p-prA during the isolation of peroxisomes or there are vesicular elements containing either Ylr324p or Ygr004p remains to be determined. It is also noteworthy that Ygr004p-prA consistently migrated as two distinct molecular species in SDS-PAGE (see Figure 4-3C). The reason for this heterogeneity is unknown but could be due to some form of posttranslational modification, e.g. phosphorylation, of Ygr004p-prA.

Peroxisomes were hypotonically lysed by incubation in dilute alkali Tris buffer and subjected to centrifugation to yield a supernatant (Ti8S) enriched for matrix proteins and a pellet (Ti8P) enriched for membrane proteins (Figure 4-3D). The chimeras of Ylr324p, Ygr004p and Ybr168p localized primarily to the Ti8P fraction, as did the chimeras of the peripheral peroxisomal membrane protein Pexl7p (Huhse *et al.,* 1998) and the integral peroxisomal membrane protein Pex3p (Höhfeld *et al.*, 1991). The soluble peroxisomal matrix protein Potlp was found almost exclusively in the Ti8S fraction. The reproducible presence of some Ylr324p-prA in the Ti8S fraction may again be representative of a soluble form of this protein that is selectively liberated from peroxisomes during their isolation (see Figure 4-3, B and C). The Ti8P fractions were then extracted with alkali sodium carbonate and subjected to centrifugation (Figure 4- 3D). This treatment releases proteins associated with, but not integral to, membranes (Fujiki *et al.*, 1982). Under these conditions, Ylr324p-prA, Ygr004p-prA and Ybr168ppA fractionated with Pex3p-prA to the pellet fraction enriched for integral membrane proteins, while Pexl7p-prA fractionated to the supernatant fraction enriched for soluble proteins, including peripheral membrane proteins. These data suggest that Ylr324p,



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Figure 4-3 D. Ylr324p-prA, Ygr004p-prA and Ybr168p-prA are integral membrane proteins of peroxisomes. (D) Peroxisomes purified by isopycnic density gradient centrifugation were lysed by treatment with 10 mM Tris-HCl, pH 8.0, releasing matrix proteins to a supernatant fraction (Ti8S) after centrifugation. The membrane-containing pellet fraction (Ti8P) was treated with 0.1 M  $Na<sub>2</sub>CO<sub>3</sub>$ , pH 11.3, and then subjected to centrifugation to yield a supernatant fraction  $(CO_3S)$  enriched for peripherally associated membrane proteins and a pellet fraction  $(CO_3P)$  enriched for integral membrane proteins. Equal portions of the respective supernatant and pellet fractions were analyzed by immunoblotting. Immunodetection of Potlp, Pexl7p-prA and Pex3p-prA marked the fractionation profiles of a matrix, peripheral membrane and integral membrane protein, respectively.

Ygr004p and Ybr168p are primarily integral peroxisomal membrane proteins, as has been shown for J7Pex23p (Brown *et al.,* 2000).

# **4.5 Ylr324p, Ygr004p and Ybrl68p act to control the number and size of peroxisomes**

Immunofluorescence analysis of oleic acid-incubated wild-type *BY4742* cells with antibodies to the carboxyl-terminal PTS1 tripeptide Ser-Lys-Leu (SKL) or to the PTS2 containing enzyme Potlp showed a pattern of small punctate structures characteristic of peroxisomes (Figure 4-4). In contrast, the majority of cells of the *ylr324* $\Delta$  strain showed increased numbers of punctate structures, while cells of the *ygr004*∆ and *ybr168*∆ strains showed a reduced number of often enlarged punctate structures. There was no evidence of increased cytosolic immunofluorescence with either anti-SKL or anti-Potlp antibodies in cells deleted for one or more of the *YLR324w*, *YGR004w* and *YBR168w* genes, suggesting that these genes do not encode proteins required for the import of matrix proteins into peroxisomes (Figure 4-4). Dr. Juan Carlos Guzman Torres constructed the double deletion mutant *DK1* and *DK2* and checked for their growth defect. Both the single and double deletion mutants grew like the wild-type cells on oleic acid-containing medium (Figure 4-3 E).

In electron micrographs, wild-type cells grown in oleic acid-containing medium contained characteristic peroxisomes of  $0.3 \mu m$  to  $0.5 \mu m$  in diameter that were well separated from one another (Figure 4-5, A and I). In contrast, cells of the *ylr324* d strain (Figure 4-5 B) showed increased numbers o f peroxisomes (Table 4-2) that were similar in



Figure 4-3 E. Growth of different strains on YPBO medium. All strains were spotted after serial dilution and grown over a period of 4 days at 30°C. *DK1* and *DK2* were constructed by Dr.Juan Carlos Guzman Torres.



Figure 4-4. Cells deleted for one or more of the *YLR324w*, *YGR004w* and *YBR168w* genes exhibit altered peroxisome morphology. Cells of the wild-type strain *BY4742* and of the *ylr324A. ygr004A, ybr!68A, ylr324wA/ygr004wA (DK1*), *ylr324wA/ybrl68wA* (*DK2*), *ygr004wA/ybrl 68wA* (*DK3)* and *ylr324wA/ygr004wA/ybrl 68wA (TKO*) deletion strains were grown in YPD medium for 16 h, transferred to YPBO medium, and incubated for 8 h in YPBO medium. Cells were observed by immunofluorescence microscopy with antibodies to the PTS1 tripeptide SKL (SKL) or to the PTS2-containing protein, Potlp. Rabbit primary antibodies (SKL) were detected with FITC-conjugated secondary antibodies. Guinea pig primary antibodies (Potlp) were detected with rhodamine-conjugated secondary antibodies. Bar, 1 um.



Figure 4-5. Cells harboring deletions in one or more of the *YLR324W, YGR004W* and YBR168w genes exhibit peroxisomes that are

altered in number and/or size. Ultrastructure of wild-type  $\frac{BY4742}{A}$ ,  $\frac{y}{r}324\Delta$  (B), *ygr004A* (C) *ybrl68A* (D) *ylr324A/ygr004A* (E), *ylr324A/ybrl68A* (F), *ygr004A/ybrl68A* (G) and *ylr324A/ygr004A/ybrl68A* (H) cells. Cells were grown in YPD medium for 16 h, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for electron microscopy. P, peroxisome; P\*. peroxisome cluster. Bar, 1.0 pm.



Figure 4-5 I. Morphometric analysis of peroxisomes of oleic acid-incubated wild-type *(WT) BY4742* and deletion mutant cells. Fifty randomly selected cells from each strain were analyzed by the program analySIS 3.1 (Soft Imaging System) to determine the areas of individual peroxisomes. Peroxisomes were then separated into size categories. A histogram was generated for each strain depicting the percentage of total peroxisomes occupied by the peroxisomes of each category. The numbers along the x-axis are the maximum sizes of peroxisomes in each category (in  $\mu$ m<sup>2</sup>).

Strain	Cell area assayed $(\mu m^2)$	Average area of peroxisomes <sup>a</sup> $(\mu m^2)$	Average number of peroxisomes per cell	Peroxisome count <sup>b</sup>	Numerical density of peroxisomes <sup>c</sup>
WT (BY4742)	402	0.06	1.32	0.16	0.73
$Yl$ r324 $\varDelta$	447	0.05	3.02	0.34	1.75
ygr0044	434	0.09	1.68	0.19	0.70
ybr1684	461	0.13	1.72	0.19	0.56
DKI	478	0.07	2.20	0.23	0.95
DK <sub>2</sub>	490	0.08	2.64	0.27	1.02
DK3	463	0.12	1.98	0.21	0.66
<b>TKO</b>	498	0.08	6.24	0.63	2.43

Table 4-2. Average area and numerical density of peroxisomes in cells of wild-type and mutant strains

<sup>a</sup>Average area on micrographs.

 $\Omega_{\text{Number}}$  of peroxisomes counted per  $\mu$ m<sup>2</sup> of cell area on micrographs.

"Number of peroxisomes per  $\mu$ m" of cell volume (Weibel and Bolender, 1973).

size to peroxisomes of wild-type cells (Figure 4-5 I) (Table 4-2). Cells of the *ygr004A* (Figure 4-5 C) and *ybrl68A* (Figure 4-5 D) strains exhibited similar numbers of peroxisomes to wild-type cells (Table 4-2), but these peroxisomes were noticeably larger than wild-type peroxisomes, particularly in the case of  $vbr168\Delta$  cells (Figure 4-5 I) (Table 4-2). Cells of strain *DK1* (Figure 4-5 E) carrying deletions in the *YLR324w* and *YGR004w* genes exhibited a mixed phenotype of increased numbers of peroxisomes (Table 4-2) of normal to enlarged size (Figure 4-5 I) (Table 4-2). Cells of strain  $DK2$ (Figure 4-5 F) carrying deletions in the *YLR324w* and *YBR168w* genes showed increased numbers of peroxisomes (Table 4-2) of normal to enlarged size (Figure 4-5 I), some of which exhibited clustering (Figure 4-5 J), while cells of strain *DK3* (Figure 4-5 G) deleted for the *YGR004w* and *YBR168w* genes also contained greatly enlarged peroxisomes (Figure 4-5 I) (Table 4-2). Cells of the strain *TKO* carrying deletions in all three genes showed an approximately 5-fold increase in the average number of peroxisomes per cell, but the size distribution of peroxisomes was similar to that of wildtype cells (Figure 4-5 I). Our results suggest that Ylr324p acts primarily as a negative regulator of peroxisome number, while Ygr004p and particularly Ybr168p act as negative regulators of peroxisome size.

#### **4.6 Overexpression studies of** *YGR004w***,** *YLR324w* **and** *YBR168w*

Because cells deleted for one or more of the *YLR324w, YGR004w* and *YBR168w* genes are compromised in their regulation of peroxisome size and number, we investigated the effects of overexpression of these genes on the overall peroxisome phenotype in cells harboring various combinations of deletions of these genes.



Figure 4- 5 J. Abnormal morphology of peroxisomes in a double deletion mutant, *DK2* showing clustering of peroxisomes with membrane thickening, suggesting that this could be a fission or a fusion mutant.

Overexpression of *YLR324w, YGR004w* and *YBR168w* (Figure 4-6) in their respective deletion backgrounds led to restoration of the wild-type peroxisomal phenotype. Overexpression of *YLR324w* in cells deleted for one or both of *YGR004w* and *YBR168w* did not lead to the complementation of the peroxisomal phenotype observed in cells deleted for either or both of *YGR004w* and *YBR168w* (Figure 4-6). Similarly, overexpression of *YBR168w* in cells deleted for one or both *YLR324w* and *YGR004w* did not lead to complementation of the peroxisomal phenotype seen in cells deleted for either or both of the *YLR324w* and *YGR004w* genes (Figure 4-8). In contrast, overexpression of the *YGR004w* gene in cells deleted for one or both of the *YLR324w* and *YBR168w* gene led essentially to the restoration of the wild-type peroxisomal phenotype, although evidence of peroxisomal clustering was still observed in cells deleted for both the *YLR324w* and *YBR168w* genes (Figure 4-7). Taken together, these data suggest that Ylr324p and Ybrl68p cannot functionally substitute for one another or for Ygr004p, while Ygr004p shows partial, but not complete, functional redundancy with Ylr324p and Ybrl68p.

### **4.7 Interacting partners of Ylr324p, Ygr004p and Ybrl68p**

A limited yeast two-hybrid screen was performed to identify physical interactions between Ylr324p, Ygr004p and Ybr168p and between these proteins and Pex11p, Pex25p, Pex28p, Pex29p and Vpslp, which have also been implicated in the control of peroxisome size and number (Erdmann and Blobel, 1995; Marshall *et al.,* 1995; Hoepfher *et al.,* 2001; Smith *et al.,* 2002; Vizeacoumar *et al.,* 2003). Others have used this methodology to detect interactions between peroxins (for examples, see Girzalsky *et al.,*



Figure 4-6. Peroxisome morphology in cells of gene overexpression strains

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*Overexpression ofYGR004w* P\_\_\_\_\_\_\_\_\_\_\_ *BY4742 ylr324A ygr004A ybr168A* \ **P\*** *J DK1 DK2* M\* i *i* DK3 **i** *TKO*  $P^*$ 

Figure 4-7. Peroxisome morphology in cells of gene overexpression strains

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Figure 4-8. Peroxisome morphology in cells of gene overexpression strains

1999; Smith and Rachubinski, 2001; Sichting *et al.*, 2003). Chimeric genes were made by ligating the ORFs of *YLR324w, YGR004w, YBR168w, PEX11, PEX25, PEX28, PEX29* and *VPS1* in frame and downstream of sequences encoding one of the two functional domains (AD or DB) of the GAL4 transcriptional activator. All possible combinations of plasmid pairs encoding AD and DB fusion proteins were transformed into *S. cerevisiae* strain *SFY526*, and β-galactosidase filter detection assays were performed. Interactions were detected between Ylr324p and Ygr004p, Pex29p and itself (Figure 4-9). Ygr004p also interacted with itself, while weak interactions were detected between Ybrl68p and Ylr324p and between Ybrl68p and Pex28p (Figure 4-9). No interactions were detected between Ylr324p, Ygr004p or Ybr168p and any of Pexl1p, Pex25p, Pex29p and Vpslp.

# **4.8** *PEX28* **and** *PEX29* **function upstream of** *YLR324w> YGR004w* **and** *YBR168w*

Since yeast two-hybrid analysis provided evidence of interaction between the *S*. *cerevisiae* homologs of YIPex23p and the *S. cerevisiae* homologs of YIPex24p and because cells of deletion mutants of the genes encoding these proteins are affected in the control of peroxisome size and number (data herein and Vizeacoumar *et al.*, 2003), we investigated the hierarchical organization of these genes. Strains containing systematic pairwise gene deletions of the *S. cerevisiae* homologs of *YIPEX23* and *YIPEX24* were made, namely *pex28A/ylr324A (CD 1), pex28A/ygr004A* (*CD2),pex28A/ybrl68A* (*CD3),*



Figure 4-9. Yeast two-hybrid analysis of the interaction partners of Ylr324p, Ygr004p and Ybrl68p. A p-galactosidase filter detection assay is presented. *SFY526* cells synthesizing both Gal4-AD (left columns) and Gal4-DB (right columns) fusion proteins were tested for  $\beta$ -galactosidase activity. The color intensities of transformants for each strain is shown.



Figure 4-10. Peroxisome morphology in *S. cerevisiae* cells deleted for a *YIPEX23* homolog and a *YIPEX24* homolog. Ultrastructure of  $pex28\Delta/y\text{lr}324\Delta$  (A), *pex28A/ygr004A* (B), *pex28A/ybr!68A* (C), *pex29A/ylr324A* (D), *pex29A/ygr004A* (E), and *pex29A/ybrJ68A* (F) cells. Cells were grown in YPD medium for 16 h, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for electron microscopy. P, peroxisome;  $P^*$ , peroxisome cluster. Bar, 1  $\mu$ m.

*pex29A/ylr324A* (*CD4*), *pex29A/ygr004A (CDS)* and *pex29A/ybr!68A* (*CD6).* Electron microscopy analysis (Figure  $4-10$ ) showed that the majority of cells containing double deletions of a *YIPEX23* gene homolog and a *YIPEX24* gene homolog exhibited the clustered peroxisomal phenotype typical of *pex28A* and *pex29A* cells (Vizeacoumar *et al.*, 2003), although cells of the *CD2* and *CD3* strains (Figure 4-10 B) and C, respectively) often exhibited enlarged peroxisomes that were well separated one from another. Together our data suggest that *YLR324w, YGR004w* and *YBR168w* function downstream of *PEX28* and *PEX29*.

#### **4.9 Discussion**

#### **4.9.1 Growth and division of peroxisomes**

Studies combining electron microscopy morphometry, pulse-chase analysis of peroxisomal protein trafficking *in vivo*, and the isolation and protein characterization of distinct peroxisomal subforms have shown that yeast peroxisomes do not grow and divide at the same time (Veenhuis and Goodman, 1990; Tan *et al.,* 1995; Titorenko *et al.,* 2000). There appears to be at least two different temporal patterns of peroxisome growth and division. In the yeast *Candida boidinii* (Veenhuis and Goodman, 1990), an initial event in peroxisome development is the extensive proliferation of immature peroxisomal vesicles containing only minor amounts of matrix proteins. This large increase in the number of immature peroxisomes by division precedes their growth through the import of membrane and matrix proteins and their conversion to mature organelles containing the complete set of peroxisomal proteins (Veenhuis and Goodman, 1990). The timing of

events of peroxisome growth and division is different in *Y. lipolytica.* In this organism, the growth of immature peroxisomal vesicles, which is accomplished by the import of matrix proteins, and their development into mature peroxisomes occur before completely assembled mature peroxisomes undergo division (Titorenko *et al.*, 2000). Similar temporal patterns of peroxisome growth and division have been observed for the yeast *Hansenula polymorpha (Tan et al.,* 1995). In human cells, both immature peroxisomal vesicles and mature peroxisomes are proposed to be able to divide (Gould and Valle, 2000). However, the division of immature peroxisomes prior to their growth and maturation by peroxisomal protein import can be seen only in some peroxin-deficient fibroblasts following reactivation or reexpression of an originally defective peroxinencoding gene (Matsuzono *et al.,* 1999; South and Gould, 1999; Sacksteder and Gould, 2000). In normal human cells, in contrast, conversion of immature peroxisomal vesicles to mature peroxisomes by membrane and matrix protein import may occur before peroxisomes undergo division (Gould and Valle, 2000).

#### **4.9.2 Molecular components in division**

Members of the Pex11 family of peroxins, including Pex25p (Smith *et al.*, 2002) and Pex27p (Tam *et al.,* 2003; Rottensteiner *et al.,* 2003b) of *S. cerevisiae*, have been shown to effect peroxisome division in different organisms (Erdmann and Blobel, 1995; Marshall *et al.*, 1995; Sakai *et al.*, 1995; Li and Gould, 2002; Li *et al.*, 2002). The dynamin-like protein Vps1p has also been implicated in this process (Hoepfner *et al.*, 2001), and we recently showed that the peroxisomal integral membrane proteins Pex28p and Pex29p are also involved in controlling the number, size and separation of peroxisomes in *S. cerevisiae* (Vizeacoumar *et al.,* 2003). Here, we have identified three novel peroxisomal proteins encoded by the ORFs *YLR324w, YGR004w* and *YBR168w* of *S. cerevisiae* and demonstrated that these proteins also act to control peroxisome size and number in this organism.

The identification of novel proteins required for peroxisome biogenesis in S. *cerevisiae* through their sequence similarity with known peroxins in other organisms enabled the identification of Pex28p and Pex29p (Vizeacoumar *et al.*, 2003). Y/Pex23p is a peroxisomal integral membrane protein required for peroxisome assembly in *Y. lipolytica* that shares extensive sequence similarity to three proteins of unknown function and unknown localization encoded by the ORFs *YLR324w, YGR004w* and *YBR168w* of the *S. cerevisiae* genome. Genomically encoded protein A chimeras of Ylr324p, Ygr004p and Ybr168p were shown by a combination of confocal microscopy and subcellular fractionation to be peroxisomal proteins. In their response to extraction by different chaotropic agents, Ylr324p, Ygr004p and Ybrl68p act primarily as integral membrane proteins.

#### **4.9.3 A role for the novel genes in peroxisome biogenesis**

Ylr324p, Ygr004p and Ybrl68p are not required for peroxisome assembly *per se,* as cells harboring deletions for one, two or all three of these genes still contain peroxisomes that are unaffected in their capacity to import PTS1- or PTS2-containing proteins. These peroxisomes are functional, at least to a degree, as the cells harboring deletions of these genes are able to grow in oleic acid-containing medium with essentially the same kinetics as wild-type cells (Figure 4-3 E). *YLR324w, YGR004w* and *YBR168w* are also apparently not required for peroxisome inheritance, as all cells deleted for one or more of these genes still contained peroxisomes after numerous cell divisions. Also, if *YLR324w*, *YGR004w* and *YBR168w* had a direct role in the inheritance of peroxisomes, one might expect that a loss of peroxisomes from cells over time resulting from the impaired segregation of peroxisomes into daughter cells would lead to inhibited growth in oleic acid-containing medium for the deletion strains as compared to the wildtype strain, which was not observed.

Peroxisomes in cells deleted for the YLR324w, *YGR004w* and *YBRJ68w* genes are not normal and show distinctive phenotypic differences from wild-type peroxisomes. *ylr324A* cells showed increased numbers of peroxisomes versus wild-type cells, while *ygr0044* and *ybr1684* cells contained not only greater numbers of peroxisomes but also enlarged peroxisomes (Figure 4-5). Cells deleted for two of the *YLR324w*, *YGR004w* and *YBR168w* genes contained increased numbers of generally enlarged peroxisomes. Cells of the strain deleted for all three genes contained increased numbers of smaller to normally sized peroxisomes. Morphometric analyses and quantification revealed a 5-fold increase in the numbers of peroxisomes on average per cell for the triple deletion strain than for the wild-type strain. Although an occasional enlarged peroxisome was evident in cells deleted for all three genes, the peroxisomal phenotype of these cells strongly resembled that of cells deleted for only the *YLR324w* gene. The characteristics of peroxisomes of cells of the deletion strains are consistent with a role for *YLR324w, YGR004w* and *YBR168w* in the control of peroxisome size and number within *S*. *cerevisiae* cells. Our results suggest that Ylr324p acts primarily acts as a negative regulator of peroxisome number, while Ygr004p and particularly Ybr168p act as negative

regulators of peroxisome size. Nevertheless, Ygr004p shares some redundancy of function with Ylr324p and Ybrl68p, but not *vice versa,* as overexpression of *YGR004w* in cells deleted for one or both of the *YLR324w* and *YBR168w* genes results essentially in the reestablishment of the wild-type peroxisome phenotype, but in contrast to wild-type cells, there is some evidence of peroxisome clustering. The reason why peroxisomes cluster in these overexpressing cells is unknown.

It is interesting to note that *Y. lipolytica* cells deleted for the *PEX23* gene also show evidence of abnormal peroxisomal divisional control. These cells lack mature peroxisomes but do accumulate small vesicular structures that contain both peroxisomal matrix and membrane proteins (Brown *et al.,* 2000). However, these membrane structures do not function as peroxisomes, as *pex23A* cells cannot grow on medium containing oleic acid as the sole carbon source. Therefore, although YIPex23p, like Ylr324p, Ygr004p and Ybrl68p, likely has a role in the regulation of peroxisome division, 7/Pex23p probably does not function identically to Ylr324p, Ygr004p or Ybrl68p in this process.

#### **4.9.4 Interaction of genes involved in the same process**

Pex28p and Pex29p have been implicated in the control of peroxisome size and number (Vizeacoumar *et al.,* 2003). A limited yeast two-hybrid screen revealed physical interactions among Ylr324p, Ygr004p, Ybrl68p, Pex28p and Pex29p. No interactions were detected between these five proteins and Pexl1p, Pex25p and Vpslp, which also have been shown to play a role in the control of peroxisome size and division. Further experimentation is required to determine whether these interactions are direct or bridged by other proteins. It is interesting to note that Ylr324p was shown to interact with Pex29p. Some amount of Ylr324p (Figure 4-3) and of Pex29p (Vizeacoumar et al., 2003) was always present in the 20KgS fraction in differential fractionation and in the less dense fractions during the gradient isolation of peroxisomes. Whether some portion of these proteins forms a complex and is localized to some compartment other than peroxisomes remains to be determined.

How might Ylr324p, Ygr004p, Ybrl68p, Pex28p and Pex29p act to control the abundance, size and distribution of peroxisomes in the *S. cerevisiae* cell? Cells systematically deleted for one of the *YLR324w, YGR004w* and *YBR168w* genes and one of the *PEX28* and *PEX29* genes exhibited clusters of peroxisomes typically observed in cells deleted for the *PEX28* or *PEX29* gene (Vizeacoumar *et a l,* 2003). Our data suggest that Pex28p and Pex29p act upstream of Ylr324p, Ygr004p and Ybr168p in controlling peroxisome abundance and size.

Organelles are highly dynamic structures that undergo fission and fusion to control their numbers and modify their morphology in response to intracellular and extracellular cues and to permit their correct segregation at cell division. As a consequence, the maintenance of compartmental integrity by the eukaryotic cell requires the tight coordination of mechanisms controlling these events. Many proteins, including those encoded by the genes *YLR324w, YGR004w* and *YBR168w* of *S. cerevisiae*, are involved in controlling peroxisome number and size in the cell. Because of their role in the control of peroxisome size and number, we propose that *YLR324w*, *YGR004w* and *YBRl68w* be designated as *PEX30, PEX31* and *PEX32,* respectively, and their encoded peroxins as Pex30p, Pex31p and Pex32p. The challenge remains to understand how the increasing number of proteins shown to be involved in controlling peroxisome number

and size interplay amongst themselves and signal to the cell how to control peroxisome dynamics.

### **CHAPTER FIVE**

## **THE Pexl9p-BINDING REGIONS OF THE PEROXINS Pex30p AND Pex32p ARE REQUIRED FOR THEIR PEROXISOMAL LOCALIZATION BUT ARE SEPARATE FROM THEIR PEROXISOMAL TARGETING SIGNALS**

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### **5.1 Overview**

The assembly of proteins at the peroxisomal membrane is a multi step process requiring their recognition in the cytosol, their targeting to the peroxisomal membrane, their insertion into that membrane and their stabilization within the lipid bilayer. The famesylated peroxin (protein required for peroxisome assembly) Pexl9p has been suggested to be the receptor that recognizes and targets newly synthesized peroxisomal membrane proteins (PMP). However, other evidence has implicated Pexl9p not as an import receptor for PMPs but as a chaperoning molecule required for the stabilization of PMPs at the peroxisomal membrane. Differentiation between these two purported roles for Pexl9p could be achieved by determining whether a PMP's targeting signal (mPTS) and region of Pex19p binding are one and the same or different. We have addressed the role of Pex19p in the assembly of two PMPs, Pex30p and Pex32p, of the yeast *Saccharomyces cerevisiae.* Pex30p and Pex32p are dispensable for normal peroxisome function, as cells lacking one or both of these proteins can still grow on oleic acidcontaining medium, the metabolism of which requires functional peroxisomes. Systematic truncations from the carboxyl terminus, together with in-frame deletions of specific regions, have identified the mPTSs essential for the targeting of Pex30p and Pex32p to peroxisomes. Both Pex30p and Pex32p interact with Pexl9p in regions that do not overlap with their experimentally identified mPTSs. However, Pexl9p is required for localizing Pex30p and Pex32p to peroxisomes, because mutations that disrupt the interaction of Pex30p and Pex32p with Pex19p lead to their mislocalization to a compartment other than peroxisomes. Mutants of Pex30p and Pex32p that localize to peroxisomes but exhibit the phenotypes of cells lacking these proteins demonstrate that

the peroxisomal targeting and functionality of these two proteins are separable. Together, our data show that the interaction of Pex30p and Pex32p with Pexl9p is required for their roles in peroxisome biogenesis and are consistent with a role for Pexl9p in stabilizing or maintaining membrane proteins in peroxisomes

# **5.2 Identification of the region containing the targeting information in Pex30p and Pex32p**

Pex30p and Pex32p are two newly identified peroxisomal integral membrane proteins (Vizeacoumar *et al.,* 2004). To delineate the mPTS of these two proteins, strains carrying deletions of varying lengths from the C-terminus regions were constructed. All these constructs were either fusion chimeras of protein A for biochemical analysis or GFP+ tagged for confocal microscopy. All constructs were made by Wanda N. Vreden. Several truncated mutants were constructed but only selective constructs will be discussed for simplicity. Subcellular fractionation was used to establish if the truncated mutants of Pex30p and Pex32p are associated with peroxisomes. In the case of Pex30p, the truncated protein Pex30(l-250)p, like the full length protein, preferentially localized to the 20KgP fraction enriched for peroxisomes. Further truncations of this protein preferentially accumulated in the 20KgS fraction (Figure 5-1 A) suggesting that the targeting information is present within the amino acid residues 230 to 250. In the case of Pex32p, the truncated protein Pex32(l-179)p, like the full length protein, preferentially localized to the 20KgP fraction enriched for peroxisomes. Further truncations of this protein preferentially accumulated in the 20KgS fraction (Figure 5- 1B) suggesting that the targeting information for Pex32p is present within the amino acid residues 159 to 179. Peroxisomes were isolated from the 20KgP fractions of each of the strains expressing the truncated mutant proteins by isopycnic density gradient centrifugation. The gradients were fractionated, and equal portions of each fraction were analyzed by immunoblotting (Figure 5- 1C and ID). All the truncated mutant proteins that enriched in the 20KgP fraction, co enriched with the peroxisomal matrix protein thiolase (Potlp). As a slight shift in the peaking of the purified peroxisomal fraction was observed, the localization of the truncated proteins was determined by tagging them with  $GFP+$  and performing a confocal microscopy analysis.

A fluorescent chimera of *Discosoma sp.* red fluorescent protein (DsRed) and the PTS1 Ser-Lys-Leu have been shown to target to peroxisomes of *S. cerevisiae* (Wang *et al.,* 2001; Smith *et al.,* 2002). Genomically encoded GFP+ chimeras of truncated proteins of Pex30p and Pex32p were localized in oleic acid-induced cells by fluorescence microscopy with DsRed-PTSl to identify peroxisomes (Figure 5- 2A and 2B). Pex30(l-282)p, Pex30(l-250)p Pex32(l-203)p and Pex32(l-179)p, like their respective full lengths, colocalized with DsRed-PTSl to small punctate structures characteristic of peroxisomes by confocal microcopy. Therefore, both microscopic analysis and subcellular fractionation showed that the truncated chimeric proteins are targeted to the peroxisomal compartment and that the targeting information lies in a span of 20 amino acid residues present in these proteins.



Figure 5-1 A-D. Subcellular fractionation of full length and truncated mutants. (A and B) Full length and truncated mutants of Pex30p and Pex32p are targeted to the organellar compartment. A PNS fraction was divided by centrifugation into a supernatant (20KgS) fraction enriched for cytosol and a pellet (20KgP) fraction enriched for peroxisomes. Equivalent portions of each fraction were analyzed. Immunoblotting with rabbit anti-Potlp antibodies detected the protein A chimeras shown. (C and D) Organelles in the 20KgP fraction were separated by isopycnic centrifugation on a discontinuous Nycodenz gradient. Fractions were collected from the bottom of the gradient, and equal portions of each fraction were analyzed by immunoblotting. Fractions enriched for peroxisomes were identified by immunodetection of Potlp. Strains expressing truncated protein were constructed by Wanda N. Vreden.



Figure 5- 2 A-D. Confocal analysis of full length and truncated mutants. (A and B) Full length and truncated mutants of Pex30p and Pex32p are targeted to the peroxisomal compartment. The sub cellular distributions of GFP+ chimeras were compared to that of DsRed-PTSl in oleic acid-incubated cells by double labeling. Full length Pex30p and Pex32p colocalize with DsRed-PTS1 in punctate structures characteristic of peroxisomes. (C and D) In-frame deletion of Pex30p and Pex32p (*FD30* and *FD32* respectively), shows mislocalization of these proteins largely to the cytosol. Only the 20 amino acid of interest(Pex30p and Pex32p constructs are denoted by *S30* and *S32* respectively) fused to GFP+ showed shows diffused pattern. Strains expressing truncated protein were constructed by Wanda N. Vreden.

# **5.3 The region containing the 20 amino acid residues constitute mPTS**

We next analyzed if the span of 20 amino acid residues is essential to target  $P$ ex $30<sub>p</sub>$ and Pex32p to the peroxisomal compartment. For this, strains carrying in-frame deletions of the 20 amino acid residues of interest were constructed by overlap PCR for Pex30p *(FD30)* and Pex32p *(FD32)*. The method of constructions of these strains is described in section 2.12.1. These constructs are under the control of the endogenous gene promoters. These constructs were made by Wanda N. Vreden. In the case of *FD30*, majority of the cells showed punctate pattern which colocalized with the peroxisomal marker, while a minor percentage of the cells exhibited a diffused pattern suggesting that the efficiency of targeting was greatly reduced. On the contrary, *FD32* cells, exhibited a diffused pattern with occasional punctate structures. This suggests that the region containing the 20 amino acid residues is essential for efficient targeting to the peroxisomal compartment.

To see if the 20 amino acid residues alone is sufficient for targeting, strains *S30* and *S32* were constructed. *S30* or *S32* carries only the 20 amino acid residues containing the targeting information fused to  $GFP+$ , with an ATG in the beginning of the segment while still remaining under the control of the endogenous gene promoter. These constructs were made by Wanda N. Vreden. Both *S30* and *S32* showed a diffused pattern with occasional punctate structures (Figure 5- 2 C and D). This suggests that the stretch of 20 amino acid residues alone is not sufficient for targeting of these peroxins. This was not surprising because these constructs lacked a transmembrane domain (TMD) that is essential for anchorage into the membrane. It has been shown that the TMD does not contain any

targeting information and that at least one TMD is essential for anchorage of the protein into the peroxisomal membrane (Mullen and Trelease 2000; Rottensteiner *et al.,*  $2004$ ). Remarkably, the amino acid residues within the 20 amino acid regions of Pex $30<sub>p</sub>$ and Pex32p were strikingly similar to the sorting sequences identified in many other PMPs. The similarity is shown in the table 5-1. Using the similarity, we developed a general consensus sequence, R/K,  $(X_{0.3})$ , R/K,  $(X_{0.3})$ , R/K, X, I/L, for the sorting of membrane proteins. Thus from these results and by analytical reasoning, we address these sequences as mPTS in the rest of the document. It will be interesting to identify if Pex30p and Pex32p has multiple targeting sequences as reported in CbPMP47 (Wang *et* al., 2001). Further analysis by truncating these proteins from the N-terminal will reveal the answer to this question. However, we have discussed the significance of this in the later part of the text.

### **5.4 Mapping of Pexl9p-binding region on Pex30p and Pex32p**

From our analysis, we speculate that there could be multiple factors involved in the targeting of these proteins to the peroxisomal compartment. Pex19p has been shown to interact with a number of integral membrane proteins. We investigated if Pexl9p interacts with Pex30p or Pex32p by yeast two hybrid analysis. Others have used this methodology to detect interactions between peroxins (for examples, see Girzalsky *et al.,* 1999; Sichting *et al.*, 2003; Vizeacoumar *et al.*, 2004). Chimeric genes were made by ligating the ORFs of *PEX30*, *PEX32* and *PEX19* in frame and downstream of sequences encoding one of the two functional domains (AD or DB) of the *GAL4* transcriptional activator. All possible combinations of plasmid pairs encoding AD and DB fusion



Table 5-1. A general consensus for mPTS signal present in most PMPs.

proteins were transformed into *S. cerevisiae* strains *HF7C* or *SFY526,* and tested for growth on selective plates lacking leucine, tryptophan and histidine or for the pgalactosidase filter detection assay. B-galactosidase filter detection assay yielded very weak or no interaction even in the known interactions such as  $P \text{ex11p}$  and  $P \text{ex19p}$ (Rottensteiner *et al.,* 2004). Hence we used the *His* reporter gene as selection marker. We found that Pex19p interacts with both Pex30p and Pex32p (Figure 5- 3A). None of the Gal4p fusion proteins alone were able to self activate the transcription of *His* reporter gene. However, since Pexl9p self activates as a fusion with BD, we were not able to reverse the positions of the prey and bait vectors. It will be interesting to verify the binding properties of bacterially expressed recombinant proteins of Pex30p and Pex32p with Pexl9p.

While our work was still in progress, Erdmann's group published (Rottensteiner *et al.,* 2004) an interesting consensus Pexl9p binding region present in most integral membrane proteins of peroxisomes. By using a computer aided sequence analysis search, (Protein Sub-string Match Analysis (PSMA); OS, C++, in Unix platform), we searched for the presence of the consensus sequence and identified multiple putative Pexl9p binding regions in Pex30p and Pex32p (listed in Table 5-2). It was interesting that the mPTS signals identified experimentally overlapped with one of the putative Pexl9p binding region of Pex30p. In the case of Pex32p, we found that none of the putative Pexl9p binding region overlapped with the experimentally determined mPTS signal. Based on this, we constructed truncated mutants to map the Pexl9p binding regions on Pex30p and Pex32p. To further delineate the exact site of Pex19p interaction and to see if there are multiple interacting regions, we systematically constructed point mutants to

disrupt the interaction of Pex30p/Pex32p with Pex19p. Rottensteiner et al., 2004, have shown that in *S. cerevisiae*, Pex11p and Pex13p interact with Pex19p at only one site by substituting proline within the core Pexl9p binding region. This core Pexl9p binding region, is suggested to form a small  $\alpha$ -helix. Hence we made similar substitutions with proline in the listed (Table 5-2) putative Pex19p binding site. In the case of Pex30p, we found that the entire interaction was abolished when a proline substitution was made in the position  $L104$  (Figure 5- 3B). Similarly, in the case of Pex32p, we found that a single substitution of L142P, abolished Pex19p interaction (Figure 5-3B). These point mutants L104P and L142P were still functional as they were able to interact with Pex28p and Pex29p respectively (Vizeacoumar *et al.*, 2004). Thus like Pex11p and Pex13p, Pex30p and Pex32p, interacts with Pexl9p at only one site and this region does not overlap with the experimentally identified mPTS signal (Figure 5- 3C).

#### **5.5 Pexl9p controls the cellular distribution of Pex30p and Pex32p**

Our result, thus far suggests that the mPTS signal and the Pexl9p binding region are non-overlapping. This was interesting because, if Pexl9p should act as a receptor for integral membrane proteins, then one would expect that the mPTS signal and the receptor binding region on the cargo must be one and the same. Since our data does not suggest that Pexl9p functions as an import receptor, we next investigated if Pexl9p has any role in controlling the distribution of these proteins to the peroxisomal compartment. To analyse this, we examined the distribution of  $Pex30p/Pex32p$  tagged with GFP+ in a strain expressing MadlNLS-PEX19 (refer section 2.9.2), a form of Pexl9p that is



Figure 5-3 A and B. Identification of Pex19p-binding site in Pex30p and Pex32p. (A) Interaction of full length and truncated mutants with Pexl9p in a yeast two-hybrid assay. **Full-length and various** truncations thereof were fused to the *GAL4* binding domain (Gal4p-BD) in vector pGBT9. The resulting plasmids were cotransformed into HF7C with a pGAD424-derived plasmid expressing a *PEX19-GAL4* activation domain (Gal4p-AD) fusion. As controls, empty pGBT9 or pGAD424 plasmids were used for transformation. Two independent sets of transformants were tested for growth on histidine, leucine and tryptophan dropout plates. Pexl9p interactions were not tested in the opposite orientation because of auto activity of the Pexl9p-Gal4p-BD fusion protein. (B) The effect of mutating the Pexl9p-binding site in vivo. A yeast two-hybrid assay was used to study the interaction of Pexl9p with full-length Pex30p and Pex32p, both mutated at position L104 and L142 respectively. Functionality of the mutated full-length-Gal4 BD fusions was tested for their known interaction with Pex29p and Pex28p respectively. Figure 5-3 A was contributed by Wanda N.Vreden.

Table 5-2. Putative Pexl9p binding sites in Pex30p and Pex32p.


#### *QCOQCFIL TVWjXXIA CFIL Q VWY][CIL VjXX[A CFIL VWY][IL QR VjXXX*



Figure 5-3 C. Schematic view of Pex30p and Pex32p showing that the Pex19p binding site and the mPTS signal are separable. A proposed TMD adjacent to the mPTS is also shown. Numbers indicate amino acid positions. Highlights in blue represent the Putative Pexl9p binding site. Highlights in green represent mPTS signal. Yellow box represent the transmembrane domain (TMD) adjacent to the mPTS signal. The position of the amino acid residues are indicated.

efficiently targeted to the nucleus. Similar approaches were adapted in other studies (Jones *et al.,* 2001 and 2004). This is done in *pex!9A* strain background. Briefly, cells expressing GFP+ chimeras of Pex30p or Pex32p in a *pex!9A* strain were transformed with MadlNLS-PEX19 plasmid. These cells were grown in minimal medium and induced for 8h with oleic acid-containing medium. Cells were washed and processed for fluorescence microscopy. Both Pex30p-GFP+ and Pex32p-GFP+ localized to punctate structures in a strain deleted for *PEX19* (Figure 5-4C). This suggests that these proteins are targeted to some other compartment as *pex!9A* strain harbors no peroxisomes (Figure 5- 4C). While in the case of MadlNLS-PEX19 expressing cells, Pex30p-GFP+ chimera and Pex32p-GFP+chimera localized to slightly elongate structures in the periphery of the nucleus (Figure 5-4 A and B). As a control, an NLS-GFP-PEX19 construct was used. This always showed a nuclear localization of Pexl9p chimera. Thus the distribution of Pex30p and Pex32p seems to be under the control of Pexl9p suggesting that Pexl9p is indeed essential for targeting these proteins. This experiment also suggests that this interaction occurs in the cytosol possibly with the nascent polypeptides.

#### **5.6 Pexl9p is essential to target PMPs to the peroxisomal compartment**

If Pexl9p controls the subcellular distribution of these PMPs, then the disruption of the Pexl9p binding site should lead to the original phenotype observed in the *pexl9A* strain. Hence we constructed GFP+ chimeric proteins carrying the point mutation that abolishes the interaction with Pexl9p. The mutant construct still remains under the control of the endogenous gene promoter. This strain was transformed with the



Figure 5-4 A and B. Sub cellular distribution of Pex30p and Pex32p in a strain expressing a form of Pex19p that gets targeted to the nucleus. (A and B) Cells expressing Pex30p-GFP+ and Pex32p-GFP+ in a *PEX19* deletion background were transformed with Mad1NLS-PEX19, were grown in SM medium (Ura-), induced in YNO medium for 8 h, stained with 10 nM per microlitres of cells with Hoescht 33342 (Sigma) for 2 min and images were obtained using Olympus BX50 microscope.



Figure 5-4 C. Cells expressing Pex30p-GFP+ and Pex32p-GFP+ in a *PEX19* deletion background or cells carrying a point mutation, disrupting Pexl9p interaction and transformed with MadlNLS-PEX19, were analysed the same way as described above.

MadlNLS-PEX19 construct, induced in oleic acid-containing medium for 8 h and observed in the microscope for the localization of Pex30p and Pex32p GFP+ chimeras. The point mutants localized to random punctate structures that do not associate with the nuclear compartment, suggesting that Pexl9p indeed controls the distribution of Pex30p and Pex32p (Figure 5-4 C). We performed biochemical analysis using protein A chimeras carrying the point mutations to see the localization of these mutants. Genomically encoding protein A chimeras of Pex30p and Pex32p carrying the point mutation in wild type cells and in *pexl9A* cells were fractionated and a purified organellar fraction (20KgP) and a cytosolic fraction (20KgS) was obtained. We found that both Pex30p and Pex32p, in a strain deleted for *PEX19,* were enriched in the 20KgP (Figure 5-4 D). This is not surprising as it has been shown in  $Hs$ Pex14p and Pex3p of CHO cells are targeted to mitochondria in a *pexl9A* strain (Otzen *et al*., 2004; Honsho *et al.,* 2002) suggesting that PMPs are mistargeted in the absence of the correct compartment. When the organellar fraction was subjected to isopycnic gradient centrifugation to isolate different compartments, Pex30(L104P) and Pex32(L142P) did not co-fractionate with the purified peroxisomal fractions. Instead, they peaked at fractions 5 and 6 at the density of 1.150  $g/cm<sup>3</sup>$  (Figure 5-4E). It is interesting to note that in the strain deleted for *PEX19* these two proteins fractionated at the density of  $1.156$  g/cm<sup>3</sup>. Note that Pot1p is completely mislocalized to the cytosolic fraction when *PEX19* is deleted as there are no morphologically identifiable peroxisomes in this strain (Figure 5-4 E). Thus, either when Pexl9p interaction is disrupted or when *PEX19* is deleted, Pex30p and Pex32p seem to be mistargeted.



Figure 5-4 D and E. Sub cellular fractionation of cells expressing protein A chimeras of Pex30p and Pex32p. (D) Sub cellular fractionation of cells expressing protein A chimeras of Pex30p and Pex32p. in a *PEX19* deletion background were found to be enriched in the organellar compartment. (E) Organelles in the 20KgP fraction were separated by isopycnic centrifugation on a discontinuous Nycodenz gradient. Fractions were collected from the bottom of the gradient, and equal portions of each fraction were analyzed by immunoblotting. Fractions enriched for peroxisomes were identified by immunodetection of Potlp. Note that these proteins are targeted to a less dense fraction (1.15g/cm3) either on the deletion of *PEX19* or upon making point mutations.

#### **5.7 Mapping the functional domains of Pex30p and Pex32p**

We next sought to identify the functional domains present in the PMPs, Pex30p and Pex32p. We processed all the strains expressing the truncated mutants for electron microscopy and analyzed the morphology of peroxisomes in all these strains. We found that in both the case of Pex30p and Pex32p, cells deleted for even the shortest sequence in the C-terminus, rendered the protein non functional even though the protein was still targeted to the peroxisomal compartment. Strains carrying Pex30(l-282)p and Pex30(l-250)p, although targeted these proteins to the peroxisomal compartments, they exhibited increased number of peroxisomes as observed in the full length deletion mutant (Figure 5-5). Similarly, Pex32(l-307)p, Pex32(l-203)p and Pex32(l-179)p although entered the peroxisomal membrane, exhibited enlarged peroxisomes as observed in *PEX32* deletion mutant (Figure 5- 6). Thus it seems that the C-terminus is essential for the functionality o f these proteins. If Pex30p and Pex32p were to be targeted in a Pexl9p dependent fashion, then the point mutants that do not interact with Pexl9p must produce the deletion mutant phenotype. Indeed, this was the case as we found deletion mutant phenotype in these mutants (Figure 5-5 E and Figure 5-6 F). It is noteworthy to mention that in the *S. cerevisiae* genome Pex30p, Pex31p and Pex32p are the only three proteins that contain an unknown dysferlin domain. This domain is present towards the C-terminal region of these proteins. Truncation of this domain renders these proteins non functional. It has been suggested that these domains can bind caveolin in other organisms. However, yeast does not have any caveolin related proteins and it is of future interest to investigate the function of this domain in relation to peroxisomes. Pex24p of *Y. lipolytica* is yet another protein that contains this dysferlin domain.



Figure 5-5. Peroxisome morphology in cells either deleted for *PEX30* or truncated mutants of *PEX30* exhibit peroxisomes that are altered in number and/or size. (A) *pex304* (B) *PEX30(1-282aa)-prA* (C) *PEX30(l-250aa)-prA* (D) *PEX30(l-230aa)-prA* (E) *P30- L104P-prA* (F) *BY4742.* Cells were grown in YPD medium for 16 h, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for electron microscopy. P, peroxisome. Bar, 1  $\mu$ m.



Figure 5-6. Peroxisome morphology in cells either deleted for *PEX32* or truncated mutants of *PEX30* exhibit peroxisomes that are altered in number and/or size. (A) *pex32A* (B) *PEX32(1-307aa)-prA* (C) *PEX32(l-203aa)-prA* (D) *PEX32(l-179aa)-prA* (E) *PEX32(l-l59aa)-prA* (F) *P32-Ll42P-prA.* Cells were grown in YPD medium for 16 h, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for electron microscopy. P, peroxisome. Bar, 1  $\mu$ m.

#### **5.8 Discussion**

#### **5.8.1 Role of Pexl9p in membrane biogenesis**

The assembly of proteins at the peroxisomal membrane is a multi step process requiring their recognition in the cytosol, their targeting to the peroxisomal membrane, their insertion into that membrane and their stabilization within the lipid bilayer. While there is a general agreement that *PEX19* is required for peroxisomal membrane biogenesis, concerns have been raised regarding its specific role as an import receptor for newly synthesized PMPs (Subramani *et al.,* 1998,2000; Schliebs and Kunau, 2004). One o f the debated issues is whether the mPTS signal and the Pexl9p binding region on the cargo are one and the same to validate Pexl9p as a bona fide receptor (Snyder *et al.,* 2000; Fransen *et al.*, 2001, 2004; Biermanns and Gartner, 2001).

#### **5.8.2 Pex30p and Pex32p are ideal candidates to study PMP targeting**

Here the role of Pex19p in the targeting of the newly identified peroxins namely Pex30p and Pex32p of the yeast *S. cerevisiae* has been addressed. Most approaches using mutation of a peroxisomal integral membrane protein to probe for sequences that can act to target it to the peroxisomes lead to conditional lethal phenotype. This is due to the failure of cells to assemble peroxisomes and has been a major drawback to the study of membrane assembly. Accordingly, the newly identified PMPs, represent excellent experimental systems with which to define an mPTS, since they are dispensable for normal peroxisome function, as cells lacking one or both of these proteins can still grow on oleic acid-containing medium, the metabolism of which requires functional

peroxisomes. Various truncated forms of Pex30p and Pex32p to identify a minimal sequence that contained the targeting information were constructed. Subcellular fractionation was used to establish if the truncated mutants of Pex30p and Pex32p are associated with peroxisomes. Analysis of truncated form of Pex30p revealed that the targeting information is present within the amino acid residues 230 to 250. In the case of Pex32p, the targeting information is present within the amino acid residues 159 to 179. All the truncated mutant proteins that enriched in the 20KgP fraction, co-enriched with the peroxisomal matrix protein Potlp (Figure 5- 1C). As we observed a slight shift in the peaking fraction, we determined the localization of the truncated proteins by tagging them with GFP+ and performing a confocal microscopy analysis.

Genomically encoded GFP+ chimeras of truncated proteins of Pex30p and Pex32p co-localized in oleic acid-induced cells by fluorescence microscopy with DsRed-PTSl. By this analysis, we confirm that the targeting information is present within a stretch of 20 amino acid residues as observed by subcellular fractionation. Therefore, both subcellular fractionation and microscopic analysis showed that the truncated chimeric proteins are targeted to the peroxisomal compartment and that the stretch of 20 amino acid residues is necessary for targeting.

#### **5.8.3 mPTS signals of Pex30p and Pex32p**

To verify if this stretch of 20 amino acid residues is essential, we constructed an inframe deletion of these specific residues and determined its localization by confocal microscopy. In the absence of the 20 amino acid residues, Pex32p was not targeted to the peroxisomal compartment. Although, Pex30p still targeted, the efficiency of the targeting was greatly reduced. This suggests that the stretch of 20 amino acid residues contain essential targeting information. To see if the information contained in this sequence is sufficient, we constructed two different strains containing only the 20 amino acid residues of interest fused to GFP+. We found that neither of the constructs was localized to the peroxisomal compartment. This suggests that this sequence alone is not sufficient for efficient targeting. This also indicates that there could be more factors playing a key role in the proper targeting of these novel peroxins. Interestingly, in mitochondria, for proteins that traverse the outer membrane once, it is now clear that the targeting signal is contained within the single TMD and its flanking region. To target proteins to mitochondria, this signal must contain a relatively short membrane spanning segment that has moderate hydrophobicity and is flanked by positively charged residues (Rapaport 2003). But in the case of peroxisomes, it has been shown that the TMD does not contain any targeting information. When a basic cluster of cottonseed ascorbate peroxidase (pAPX) containing the targeting information was fused to a synthetic membrane span, the reporter protein showed punctate pattern (Mullen and Trelease 2000). Hence we conclude that the region that contains this targeting information is the bona fide mPTS of these new peroxins. However, it has been shown that PMPs use multiple targeting signals (Dyer *et al.,* 1996; Jones *et al.,* 2001; Brosius *et al.,* 2002). Although it will be interesting to see if there are multiple mPTS signal present in these proteins by making N-terminal truncations, under the specific conditions used in these experiments, these mPTS signals seem to be functional. Especially, recent evidence from Goodman's laboratory suggest that multiple targeting modules on peroxisomal proteins are not redundant and that under different metabolic states, different mPTSs could function independently and efficiently

(Wang *et al.,* 2004). Interestingly, the amino acid residues within the mPTS regions o f Pex30p and Pex32p were strikingly similar to each other and to the sorting sequences identified in many other PMPs. The similarity is shown in the table 5-1. This has led us to redefine and identify a unique signal sequence to target peroxisomal membrane proteins to the peroxisomal compartment. A general idea regarding the mPTS is that it is a hydrophilic stretch of amino acids with few basic residues. Of specific interest, this consensus mPTS signal happens to be an integral part of  $Pex19$  binding site in  $Pex11p$ , Pexl3p and even in HsALDP. Using the similarity, we developed a general consensus sequence,  $R/K$ ,  $(X_{0-3})$ ,  $R/K$ ,  $(X_{0-3})$ ,  $R/K$ ,  $X$ ,  $V<sub>L</sub>$ , for the sorting of membrane proteins. The presence of this consensus pattern in the hydrophilic stretch of residues, could be beneficial in defining the membrane targeting signals of PMPs.

#### **5.8.4 Interaction of Pexl9p with the PMPs**

The fact that Pex30p could still be targeted, although with poor efficiency, hinted us to search for other factors involved in targeting this protein to the peroxisomal compartment. In an attempt to understand the role of Pexl9p, to target proteins effectively, as a first step, we performed yeast two-hybrid analysis. We found that both Pex30p and Pex32p interact with Pexl9p. While our work was still in progress, Erdmann's group published some exciting finding by peptide scan analysis, a consensus sequence, for the identification of putative Pex19p binding sites on most PMPs (Rottensteiner *et al.,* 2004). Using this consensus sequence, we found that there are multiple putative Pexl9p binding sites in these proteins (Listed in Table 5-2). It was interesting that the mPTS signals identified experimentally overlapped with one of the putative Pexl9p binding region of Pex30p. In the case of Pex32p, we found that the putative Pexl9p binding region was at a closer proximity to the mPTS signal. We constructed truncated mutants to map the Pexl9p binding regions on Pex30p and Pex32p. We also constructed some point mutants to delineate the exact site of Pex19p interaction and to see if there are multiple interaction sites. To do this, we systematically made point mutations in the sequences of putative  $Pex19p$  binding regions to disrupt the interaction of Pex30p/Pex32p with Pexl9p. We made similar substitutions as described by Rottensteiner *et al.*, 2004, with proline in the putative Pex19p binding site. In the case of Pex30p, we found that the entire interaction was abolished when a proline substitution was made in the position L104. Similarly, in the case of Pex32p, we found that a single substitution of L142P, abolished Pex19p interaction. Thus like Pex11p and Pex13p, Pex30p and Pex32p, interacts with Pexl9p at only one site in a region non-overlapping with the experimentally identified mPTS signal. Of particular interest is the interaction site on Pex32p with Pexl9p. The interacting region and the mPTS signal are at a closer proximity separated by a distance of 9 amino acids. This is beneficial for an important reason. Reports have claimed that *PEX19* can function as an import receptor (Jones *et al.,* 2004). This is largely based on the analysis of several constructs made by Jones *et al.* These constructs contained a minimum of 60 to a maximum of 160 amino acid residues. Using these, it was shown that Pexl9p interacts within this region and it is also sufficient to target proteins to the nuclear compartment in a Pexl9p dependent manner (Jones *et al.,* 2004). What Jones et al., has observed is probably a combinatorial action of two independent factors, the mPTS and its interaction with Pexl9p. Furthermore, the argument that multiple mPTS could be found on the same protein is often used as an

argument to support the fact that Pexl9p could probably interact in those other non redundant targeting modules. Here we provide evidence that Pexl9p interacts at only one site, as in the case of Pex11p and Pex13p (Rottensteiner *et al.*, 2004) and this site is different from the mPTS signal. Interestingly, Pex19p binding regions of Pex11p and Pexl3p were shown to be enough for targeting (Rottensteiner *et al,* 2004). The possible explanation for this is that, these proteins seems to have the mPTS signal motif as an integral part of the Pex19p binding region (Table  $5-1$ ) It has been shown that multiple mPTS signals are non-redundant. PMP47 in *Candida boidini* has two discrete mPTS signals. One of them is responsible for targeting to the peroxisomes in an induced state and the other is responsible in a non induced state. PMP47 also interacts with Pexl9p, but this association is only under inducible condition. Hence, without further molecular dissection, the data of Jones *et al.*, can only be interpreted as evidence favoring a role for *PEX19* as an import receptor.

#### **5.8.5 The role of Pexl9p in PMP targeting**

We next investigated if Pex19p has any role as a receptor for targeting these proteins to the peroxisomal compartment. For this we examined the distribution of Pex30p/Pex32p genomically tagged with GFP+ in a strain expressing NLS-PEX19, a form of Pex19p that is efficiently targeted to the nucleus. Both Pex30p-GFP+ and Pex32p-GFP+ localized to a punctate structures in a strain deleted for *PEX19.* This suggests that these proteins are targeted to some other compartment as *pex!9A* strain harbors no peroxisomes. Pex30p-GFP+ chimera and Pex32p-GFP+chimera in cells expressing Mad1-NLS-PEX19, localized to slightly elongate structures in periphery of the nucleus. To further validate these findings, two different biochemical approaches were undertaken; first we monitored the localization of Pex30p and Pex32p chimeras in a strain where the interaction of these proteins with Pexl9p was disrupted by point mutations. Second, we monitored these chimeric proteins in a *pexl9A* strain. We found that in both cases, either when *PEX19* was deleted or when the interaction is disrupted Pex30p and Pex32p are targeted to a distinct compartment of lesser density than that of wild-type peroxisomes  $(1.15g/cm3)$ . These results suggest that the distribution of Pex30p and Pex32p seem to be under the control of Pex19p. Since Pex30p and Pex32p are directed to the nuclear membrane, in a Pexl9p dependent manner, one could conclude that Pexl9p binds these proteins (nascent?) in the cytosol and directs them to the nuclear membrane. Alternatively, there could be other factors that could target these proteins to the membrane associated Pexl9p for further assembly and insertion. It will be interesting to isolate such complexes from the cytosolic fraction and to see if Pexl9p forms a part of this complex.

#### **5.8.6 Functional domains of Pex30p and Pex32p**

In addition, we have also mapped the region that is essential for the functionality of these two proteins. We found that in both the case of Pex30p and Pex32p, cells deleted for even the shortest sequence in the C-terminus, rendered the protein non functional even though the protein was still targeted to the peroxisomal compartment (Figure 5- 6A to G). Thus it seems that the C-terminus is essential for the functionality of these proteins. If Pex30p and Pex32p were to be targeted in a Pexl9p dependent fashion, then the point mutants that do not interact with Pexl9p should lead to the deletion mutant phenotype. Indeed, this was the case as we found deletion mutant phenotype in these mutants (Figure 5- 5E and 6F). It is noteworthy to mention that in the yeast genome Pex30p, Pex31p and Pex32p are the only three proteins that contain an unknown dysferlin domain. This domain is present towards the C-terminus region of these proteins. Truncation of this domain renders the proteins non functional. It has been suggested that these domains can bind caveolin in other organisms. However, yeast does not have any caveolin related proteins and it is of future interest of our laboratory to investigate the function of this domain in relation to peroxisomes. Pex24p of *Y. lipolytica* is yet another protein that contains this dysferlin domain.

#### **5.8.7 Factors governing targeting of PMPs**

In summary, we find that Pex19p is essential for targeting these novel peroxins and there seems to be more factors that essentially contribute for proper targeting and insertion of PMPs into the peroxisomal compartment. Three factors can be envisioned: the interaction of the PMPs with Pexl9p, the mPTS signal of the PMPs and a TMD, adjacent to the mPTS. As such, there is no credible evidence or argument that *PEX19* does not function as an assembly factor facilitating the insertion of membrane proteins or a cytosolic chaperone. Pex30p and Pex32p are not essential for early peroxisome membrane synthesis and hence the interaction of Pex30p and Pex32p cannot be attributed to a general role of Pex19 in peroxisome membrane synthesis. Therefore, Pex19p interactions may be connected with a transport function or to facilitate insertion of proteins into the peroxisomal membrane. From our data, we suggest that the mPTS could

be a separate entity or be an integral part of the Pex19p binding site. Also, in conjunction with an adjacent TMD, proper anchorage of the peroxin could be facilitated.

To obtain a more comprehensive picture of the biogenesis of membrane proteins, the following questions still need to be addressed. Are there any other cytosolic factors that contribute to the targeting of membrane proteins? Does the specific lipid composition of the membrane have any role in targeting? Finally, does the same mechanism apply to proteins irrespective of whether the signals are contained within the C- or N-terminal regions? Future research might reveal more exciting results on how the membrane is assembled.

### **CHAPTER SIX**

## **THE DYNAMIN LIKE PROTEIN Vpslp ASSOCIATES WITH PEROXISOMES IN A Pexl9p-DEPENDENT MANNER**

#### **6.1 Overview**

Organelle division is a dynamic process regulated by multimeric protein machineries, the details of which are just beginning to be unveiled. The mechanisms underlying the accurate partitioning of yeast peroxisomes, mitochondria and vacuoles are distinct, yet share common elements. These organelles all move along actin filaments and require fusion and fission to maintain normal morphology. Dynamins are thought to play important roles in the fission of peroxisomes and mitochondria. The dynamin-related protein V pslp is suggested to be involved in peroxisome fission, as cells lacking V pslp contain a few enlarged peroxisomes. But if and how Vps1p associates with the peroxisomal membrane is unclear. Pexl9p has been shown to function as an import receptor for peroxisomal membrane proteins (PMP) and/or as a chaperone acting to stabilize PMPs at the peroxisomal membrane. Pexl9p recognizes two regions in Vpslp. Mutation of these Pex19p-binding regions does not affect vacuolar assembly, while mutation of only the first, more amino-terminal region leads to cells with reduced numbers of enlarged peroxisomes, the phenotype of cells lacking Vps1p. Vps1p does not associate with peroxisomes once its interaction with Pexl9p is disrupted. However, overproduction of Pex 19p does not alter the normal distribution of Vps  $1p$  within the cell. Together, our data suggest that Pex19p stabilizes the association of Vps1p with peroxisomes and that this interaction is required for peroxisome fission.

#### **6.2 Pexl9p interacts with Vpslp**

Rottensteiner *et al.,* 2004, developed a consensus sequence of amino acid residues that seems to be recognized by Pexl9p in most peroxisomal membrane proteins. We identified two such putative Pex19p binding sites in Vps1p using the Unix based Protein Sub-string match analysis software (Table 6-1). Vpslp belongs to the dynamin protein family and is essential for sorting proteins to the vacuolar compartment. Deletion mutant of Vps1p contains few enlarged peroxisomes (Hoepfner *et al.*, 2001). Since we found two putative Pex19p binding sites, we hypothesized that the association of Vps1p with the peroxisomes depends on Pexl9p. Yeast two hybrid analysis was used to detect any interaction of these proteins. Others have used this methodology to detect interactions between peroxins (for examples, see Girzalsky *et al.*, 1999; Sichting *et al.*, 2003; Vizeacoumar *et al.*, 2004). Chimeric genes were made by ligating the ORF of *VPS1* in frame and downstream of sequences encoding one of the two functional domains (AD or DB) of the *GAL4* transcriptional activator. All possible combinations of plasmid pairs encoding AD and DB fusion proteins were transformed into *S. cerevisiae* strains *HF7C* or *SFY526,* and tested for growth on selective plates lacking leucine, tryptophan and histidine or for the  $\beta$ -galactosidase filter detection assay.  $\beta$ -galactosidase filter detection assay yielded a weak or no interaction even in the known interactions such as Pexl lp and Pexl9p (Rottensteiner *et al.,* 2004). Using *his* as the reporter gene, we found that Pexl9p indeed interacts with  $Vps1p$  (Figure 6-1). None of the Gal4p fusion proteins alone were able to self activate the transcription of *his* reporter gene. However, since Pex19p self activates as a fusion with BD, we did not reverse the positions of the prey and bait vectors.

### **Table 6-1 Putative Pexl9p binding regions in Vpslp**





Figure 6-1. Interaction of Vps1p with Pex19p in a yeast two-hybrid assay. Full-length **genes** were fused to the *GAL4* binding domain (Gal4p-BD) in vector pGBT9. The resulting plasmids were cotransformed into HF7C with a pGAD424-derived plasmid expressing a *PEX19-GAL4* activation domain (Gal4p-AD) fusion. As controls, empty pGBT9 or pGAD424 plasmids were used for transformation. Two independent sets of transformants were tested for growth on histidine, leucine and tryptophan dropout plates. Pex19p interactions were not tested in the opposite orientation because of auto activity of the Pexl9p-Gal4p-BD fusion protein.

# **6.3 Disrupting Pexl9p interaction causes** *VPS1* **deletion mutant phenotype**

We next wanted to see if the putative Pex19p binding regions, identified using the consensus Pexl9p binding sequence, are essential for the association of Vpslp with Pexl9p. For this, we constructed strains carrying in-frame deletions of these amino acids of interest by overlap PCR designated as  $FDVI$  for  $Vps1(\Delta 509-523)$  and  $FDV2$  for  $Vps1(\Delta 633-647)$ . These mutants are still under the control of the endogenous gene promoter. To analyse the morphology of peroxisomes, we performed an indirect immunofluorecence analysis using antibodies raised against thiolase and SKL-containing peroxisomal matrix proteins. *FDV1* mutant exhibited few enlarged peroxisomes as observed in a  $vps/\Delta$  deletion mutant (Figure 6-2). The other mutant,  $FDV2$ , exhibited normal peroxisomes like the wild type strain (Figure 6-2). This suggests that the amino acid residues 501-523 in Vps1p are essential for its functional contribution towards peroxisomal compartment.

#### **6.4 Vacuolar protein sorting function is normal in** *FDV1* **mutant**

By making an in-frame deletion of the putative Pex19p binding site, one of the possibilities is that the entire protein Vps1p is rendered non functional. Alternatively, the amino acid residues between 501 and 523 could be essential for the abnormal peroxisome phenotype. This might help to dissect the domain that is essential for its function in the sorting of proteins to the vacuolar compartment to its peroxisomal role. Vacuolar



Figure 6-2. Analysis of in-frame deletions of Vps1p by immunofluorescence microscopy (*FDV1* and *FDV2* respectively). Mutant strains were grown in YPD medium for 16 h, transferred to YPBO medium, and incubated for 8 h in YPBO medium. Cells were observed by immunofluorescence microscopy with antibodies to the PTS1 tripeptide SKL (SKL) or to the PTS2-containing protein, Potlp. Rabbit primary antibodies (SKL) were detected with FITC-conjugated secondary antibodies. Guinea pig primary antibodies (Potlp) were detected with rhodamine-conjugated secondary antibodies. Bar,  $1 \mu m$ .

homeostasis requires fusion, fission and actin-dependent transport of vacuolar membranes to the daughter cells. Mutations of proteins involved in these processes often results in aberrant vacuolar morphology (Seeley *et al.*, 2002). It has been known that at least 60% of the cells exhibit fragmented vacuolar morphology when *VPS1* is deleted (Seeley *et al.*, 2002; Peters *et al.*, 2004). Using the fluorescent vacuolar vital stain FM4-64 (Vida and Emr 1995), we analysed the morphology of vacuoles in the mutants that we have constructed. We found that both *FDV1* and *FDV2* exhibited normal vacuolar morphology, unlike the deletion mutant which showed higher percentage of fragmented vacuoles (Figure 6-3A). It has been suggested that the mixed phenotype observed in  $vps/\Delta$  showing enlarged and fewer vacuoles (fission mutants) and many small fragmented vacuoles (fusion mutants), is due to its ability to mutually control membrane fission and fusion (Peters *et al.*, 2004). We also analysed the sorting of the soluble vacuolar hydrolase carboxypeptidase Y (CP-Y). CP-Y is cotranslationally translocated into the ER, where it is glycosylated. It is then transported to the Golgi where a sorting signal present in its propeptide form is recognized and then finally sorted to the vacuoles. Defects in this vacuolar sorting lead to the secretion of CP-Y through the secretory pathway, outside the cell (Marcusson *et al.,* 1994). We found that *FDV1* had a similar background as that of wild-type strain suggesting that vacuole protein sorting is unaffected in this strain (Figure 6-3B). However, the other in-frame deletion mutant *FDV2*, exhibited a slightly increased secretion of CP-Y, suggesting that it might be partly impaired in vacuolar protein sorting. As expected, *VPS1* deletion showed high secretion of CP-Y outside the cell (Figure 6-3 B).



Figure 6-3 A and B. Analysis of vacuolar morphology and protein sorting function. (A) Cells were grown overnight and sub cultured to reach an OD of 0.5. To about 100  $\mu$ l of cells, 1 µl of 8 mM FM4-64 dye was added and incubated for 30 minutes at 30 C. Cells were washed and re-incubated in fresh YPD medium for an hour and a half. Images were captured on an Olympus BX50 microscope (Olympus, Tokyo, Japan) equipped with a digital fluorescence camera (Spot Diagnostic Instruments, Sterling Heights, MI). (B) The CPY secretion assay was performed according to the method of Roberts *et al.* (1991) with minor modifications. Briefly, cells were grown to an OD of 0.5 and spotted on YEPD plates with serial dilution and allowed to dry before filter overlay and incubation. Nitrocellulose sheet was spread on these plates spotted with diluted cells and left to grow for 24 h. The membrane is carefully lifted and rinsed to remove the cells stuck to it. Immunoblotting using the CP-Y antibody is done on the membrane and the secreted protein is detected.

#### **6.5 Effect of point mutation in Vpslp**

Thus far, our results suggest that one of the putative Pex19p binding sequences in Vps1p is essential for its function in the fission of peroxisomes and we next wanted to investigate the exact site of interaction of between Pex19p and Vps1p. Hence we constructed point mutants to disrupt the interaction of VpsIp with Pex19p. Rottensteiner *et al.,* (2004), have shown that in *S. cerevisiae* Pexl lp and Pexl3p interact with Pexl9p at only one site by substituting proline at any position within the core Pexl9p binding region. Substitution of proline within the Pex19p binding region is likely to disrupt the  $\alpha$ helical conformation thought to enhance its association with Pexl9p. Hence we made similar substitutions with proline in the putative Pexl9p binding site at the residue 516  $(V516P)$  instead of valine. We performed a two hybrid analysis to see if the interaction was dismpted. Although the interaction with Pexl9p was disrupted (Figure 6-4A), the cells grew on drop- out plates over prolonged incubation. We then constructed a strain carrying this point mutation and monitored the morphology of peroxisomes by immunofluorescence and by electron microscopy. This way, the gene is still under the control of its endogenous gene promoter and does not lead to any aberrant expression. We found that the point mutant strain harbored a mixed phenotype of both enlarged peroxisomes and wild-type peroxisomes (Figure 6-4B and 6-5C). This suggests that either the point mutation substitution does not fully abolish its interaction with Pexl9p or there could be other potential factors that could still associate Vps1p with the peroxisomes.

## $\overline{A}$ GAL-4-BD GAL-4AD fused to fused to Pexllp - Pexl9p pGBT9 - Pex 19p traces and **Edmonwell** Vpslp - Pexl9p ^ \_\_\_\_\_*<* Vpsl(V512P)p - Pcx 19p





Figure 6-4 A and B. The effect of mutating the Pexl9p-binding site in vivo. (A) A yeast two-hybrid assay was used to study the interaction of Pex19p with Vps1p, mutated at position V516. (B) Peroxisomal morphology of point mutant. Cells were observed by immunofluorescence microscopy with antibodies to the PTS1 tripeptide SKL (SKL) or to the PTS2-containing protein, Potlp. Rabbit primary antibodies (SKL) were detected with FITC-conjugated secondary antibodies. Guinea pig primary antibodies (Potlp) were detected with rhodamine-conjugated secondary antibodies



**Figure 6-5.** Peroxisome morphology in wild-type cells and Vpslp mutants. (A) *BY4742* (B) *FDV1* (C) *vpslA* (D) *vps(V516P).* Cells were grown in YPD medium for 16 h, transferred to YPBO medium and incubated in YPBO medium for 8 h. Cells were fixed and processed for electron microscopy. P, peroxisome. Bar, 1  $\mu$ m.

We next attempted to find out if the distribution and association of Vpslp with peroxisomes was altered in a Pexl9p-dependent fashion. To analyze this, we examined the distribution of Vps1p genomically tagged with GFP+ in a strain expressing MadlNLS-PEX19, a form of Pexl9p that is efficiently targeted to the nucleus. Similar approaches were adapted in other studies (Jones *et al*., 2001 and 2004). This is done in *pexl9*Δ strain background. Briefly, cells expressing GFP+ chimera of Vps1p in a *pexl9*Δ, background were transformed with MadlNLS-PEX19 plasmid. Cells were grown in minimal medium and induced for 8 h with oleic acid-containing medium. Induced cells were washed and processed for fluorescence microscopy. Vpslp-GFP+ localized to punctate structures with a diffused cytosolic background in both the strains, either deleted for *PEX19* or in the case of cells expressing Mad1NLS-PEX19, a pattern observed in wild type strains (Figure 6-6 B). As a control, an NLS-GFP-PEX19 construct was used. Thus the distribution of Vps1p seems to remain unaffected by Pex19p.

## **6.7 V pslp is not recruited to the peroxisomes in a Pexl9p dependent manner**

Vps1p is predominantly a cytosolic protein and Vps1-GFP+ chimera often exhibit a diffused cytosolic stain with occasional punctate pattern. The nature of this punctate composite is unknown. Now that, we know Pex19p associates with Vps1p, we wanted to see if the localization pattern of Vps1p was altered in a strain overexpressing Pex19p. We hypothesized that overexpression of *PEX19*, could possibly lead to an increased,

detectable amount of Vps1p associating with the peroxisomes. We monitored the Vpsl-GFP+ chimera by fluorescence microscopy using DsRed-PTSl as the peroxisomal marker. A fluorescent chimera formed between *Discosoma sp.* red fluorescent protein (DsRed) and the PTS1 Ser-Lys-Leu have been shown to target to peroxisomes of S. *cerevisiae* (Wang *et al.,* 2001; Smith *et al.,* 2002). Briefly, cells expressing GFP+ chimera of Vps1p were transformed with the overexpression construct, *Yep13-PEX19* and the peroxisomal marker DsRed-PTSl into a strain deleted for *PEX19.* Cells were grown in minimal medium and induced for 8 h with oleic acid-containing medium. Induced cells were washed and processed for fluorescence microscopy. Vps1p-GFP $+$ localized to punctate structures with diffused cytosolic staining. These punctuate structures did not colocalize with the peroxisomal marker and that the subcellular distribution of Vps1p-GFP+ chimera remained unaltered (Figure 6-6A). From these data we conclude that Pex19p does not recruit Vps1p to the peroxisomal compartment.

## **6.8 V pslp associates with the membrane fraction in a Pexl9pdependent manner**

Pex19p does not seem to recruit Vps1p to the peroxisomes. To understand the significance of this interaction, we monitored the endogenous level of Vps1p in a strain where the interaction with Pexl9p was disrupted. Vpslp, being a cytosolic protein, was found to be preferentially enriched in the 20KgS fraction with a minimum amount in the 20KgP fraction (Figure 6-7). The control proteins G6PDH was found exclusively in the



Figure 6-6 A and B. Effect of Pex19p on Vps1p. (A) The sub cellular distributions of GFP+ chimeras of Vps1p were compared to that of DsRed-PTS1 in oleic acid-incubated cells by double labeling, fluorescence microscopy, in a strain over expressing *PEX19.* GFP+ chimera of Vps1p does not colocalize with DsRed-PTS1. (B) Sub cellular distribution of GFP+ chimera of Vps1p in a strain expressing a form of  $Pex19p$  that gets targeted to the nucleus. Cells expressing Vpslp-GFP+ in a *PEX19* deletion background were transformed with *MadlNLS-PEX19*, were grown in SM medium (Ura-), induced in YNO medium for 8 h, stained with 10 nM per microlitres of cells with Hoescht 33342 (Sigma) for 2 minutes and images were obtained using Olympus BX50 microscope.



Figure 6-7. Subcellular fractionation of full length, in-frame deletion mutant and the point mutant. A fraction of full length Vps1p is targeted to the organellar compartment. A PNS fraction was divided by centrifugation into a supernatant (20KgS) fraction enriched for cytosol and a pellet (20KgP) fraction enriched for peroxisomes. Equivalent portions of each fraction were analyzed. Immunoblotting with rabbit anti-Vpslp, anti-Sdh2p, anti-G6PDH and guinea-pig anti-Pot1p were used.

cytosol and the mitochondrial protein Sdh2p and the peroxisomal protein Potlp were found exclusively in the organellar pellet fraction. In the in-frame deletion mutant and the point mutant, no membrane associated V psl p was detected (Figure 6-7). This suggests that once Vps1p is recruited to peroxisomes, its interaction with Pex19p perhaps could stabilize its association with peroxisomes.

#### **6.9 Discussion**

#### **6.9.1 Organelle division**

Organelles are highly dynamic structures that undergo fission and fusion to control their numbers and modify their morphology in response to intracellular and extracellular cues and to permit their correct segregation at cell division. As a consequence, the maintenance of compartmental integrity by the eukaryotic cell requires the tight coordination of mechanisms controlling these events. Although much progress has been made, many questions concerning the mechanisms of organelle fission still remain unanswered. Division is regulated by multimeric protein machinery, the details of which are just beginning to be unveiled. It seems that the mechanism of accurate partitioning of yeast peroxisomes, mitochondria and vacuoles are distinct, yet they share common elements. All these organelles move along actin filaments and they all require fusion and fission to maintain normal morphology. Dynamin like proteins have been suggested to play a role in organelle fission.

In mitochondria, fission seems to occur by a multi-step pathway, including the recruitment of the dynamin like protein Dnm lp (and any accessory proteins) to sites on
the sides of mitochondrial tubules, the constriction of mitochondrial tubules at these sites, and the coordinated division of the outer and inner mitochondrial membranes to generate new tubule ends. While V ps1 p seems to regulate peroxisome fission. V ps1 p is a major cytosolic protein belonging to the dynamin protein family and deletion mutant of Vps1p contains few enlarged peroxisomes (Hoepfner *et al.*, 2001). Vps1p has been shown to regulate the cytoskeleton through its interaction with Slalp (Yu and Cai, 2004). Also a double deletion mutant of *vpsI* $\Delta$ *rhoI* $\Delta$  has been found to accumulate actin patches on peroxisomal membrane. These evidences suggest that Vps1p perhaps regulates fission by controlling the actin dynamics. How may Vps1p regulate peroxisomal fission and how it is recruited to the peroxisomal membrane is unclear.

### **6.9.2 Pexl9p and Vpslp**

Since Pexl9p was proposed to be an assembly factor, we hypothesized that Pex19p could recruit Vps1p to the peroxisomes and help in the membrane fission. We found two putative Pexl9p binding sites, using the Pexl9p binding consensus sequence (Rottensteiner *et al.,* 2004). By two hybrid analysis, we found that Pexl9p indeed interacts with  $V_{DS}$  (Figure 6-1).

If Pex 19 $p$  is essential for the recruitment of V $p$ s 1 $p$  to the peroxisomes, mutations disrupting the Pexl9p interaction region, should lead to the deletion mutant phenotype. Accordingly, in our immunofluorescence study, we found that *FDV1* mutant exhibit few enlarged peroxisomes as observed in a  $vps/\Delta$  deletion mutant (Figure 6-2). To verify if the other functionalities of this protein is affected, we immediately analysed the vacuolar sorting function of Vpslp, as it plays a role in vacuolar protein sorting as well. We found

that both *FDV1* and *FDV2* exhibited normal vacuolar morphology, unlike the deletion mutant which showed higher percentage of fragmented vacuoles (Figure 6-3A). It has been suggested that this mixed phenotype observed in *vpsl* A showing enlarged and fewer vacuoles (fission mutants) and many small fragmented vacuoles (fusion mutants) is due to mutual control of membrane fission and fusion by Vpslp (Peters *et al*., 2004). We also analysed the sorting of the soluble vacuolar hydrolase carboxypeptidase Y (CP-Y), as mutants involved in this could lead to the sorting of this protein to the secretory pathway. We found that *FDV1* had the same background as that of wild-type strain suggesting that vacuole protein sorting is unaffected in this strain (Figure 6-3B). However, the other inframe deletion mutant *FDV2* exhibited a relatively increased secretion of CP-Y suggesting that it might be partly impaired in vacuolar protein sorting. It is interesting to note that this mutation lies in the GED domain of Vps1p. This region has been proposed to form coiled coil structures and has been suggested to interact with Vam3p, an interaction that is essential for vacuolar fusion (Peters *et al.,* 2004).

#### **6.9.3 Distribution of Vpslp-GFP chimera**

Like, Pexl1p and Pexl3p, Vpslp interacts with Pexl9p at only one site as substitution mutation within the core Pexl9p binding region abolished the interaction to a considerable extent. In the strain carrying the point mutation, we found that the peroxisomes were of mixed phenotype of both wild-type peroxisomes and enlarged peroxisomes (Figure 6-4B and 5C). This suggests that either the point mutation substitution does not fully abolish its interaction with Pexl9p or there could be other factors that could potentially associate Vpslp with the peroxisomes. To see if Pexl9p recruits Vpslp to peroxisomes, we examined the distribution of Vpslp-GFP+ chimera in a strain expressing *MadlNLS-PEX19,* a form of Pexl9p that is efficiently targeted to the nucleus. Vpslp-GFP+ localized to punctate structures with a diffused cytosolic staining. Thus the distribution of Vpslp seems to remain unaffected by Pexl9p. We also conclude that Vps1p is not recruited to the peroxisomal compartment in a strain that overexpresses Pexl9p. It will be interesting to see if this recruitment is cell cycle dependent.

Subcellular fractionation was done to analyze the association of Vpslp with the peroxisomes. In the in-frame deletion mutant and the point mutant, no membrane associated Vps1p was detected (Figure 6-7). This suggests that Pex19p is essential for the association of Vps1p with the peroxisomes, perhaps in the stabilization of its interaction with peroxisomes. Since Vps1p is not integral to the peroxisomal membrane, this result also suggests that Pex19p does not necessarily play a role in the insertion of the membrane proteins.

#### **6.9.4 Vpslp and membrane fission**

The observation that peroxisomes do manage to divide at cytokinesis in *vpslA*, suggest that there is also dynamin-independent peroxisome fission (Hoepfher *et al*., 2001). In fact, this is re-inforced by our previous finding that more wild type peroxisomes were formed upon overexpression of  $PEX11$  in a  $vps1\Delta$  mutant (Vizeacoumar *et al.*, 2003). Also, Pexl lp has been suggested to indirectly recruit *DLP1* to the peroxisomes, although the authors were unable to detect any physical interaction between these two proteins (Li and Gould 2003). However, since we know Pexl lp is recruited in a Pexl9p dependent fashion (Rottensteiner *et al.,* 2004), we speculate that Vpslp recruitment could perhaps be coordinated through both these proteins.

Members of the Pexl 1p family of peroxins, including Pex25p (Smith *et al.*, 2002) and Pex27p (Tam *et al.*, 2003; Rottensteiner *et al.*, 2003) of *S. cerevisiae*, have also been shown to affect peroxisome division in different organisms (Erdmann and Blobel, 1995; Marshall *et al.,* 1995; Sakai *et al.,* 1995; Li and Gould, 2002; Li *et al.,* 2002). We have also shown that the peroxins Pex28p and Pex29p are necessary for the separation of peroxisomes, an event that follows division (Vizeacoumar *et al.*, 2003). Also the newly identified Pex30p, Pex31p and Pex32p are involved in controlling the number, size and separation of peroxisomes in *S. cerevisiae* (Vizeacoumar *et al.*, 2004). Their exact roles in peroxisome fission are yet to be determined and that understanding how constriction and fission of these membranes are accomplished will require more detailed analyses of these divisional components.

Another important issue that is addressed by our study is the role of Pexl9p in membrane protein sorting. It has been shown that Pex3p and Pexl9p are the only two peroxins that are required for membrane biogenesis in *S. cerevisiae* and other mammalian cells (Hettema *et al.,* 2000; Matsuzono *et al.,* 1999; Shimozawa *et al.,* 2000). Pexl9p is a major cytosolic protein with a smaller fraction of it found to be associated with the peroxisomes. It interacts with a number of peroxisomal integral membrane proteins (PMPs). The cytosolic fraction of Pexl9p has been suggested to act as a shuttling receptor for peroxisomal membrane proteins, while its peroxisome-associated form could function as a chaperone, assisting the assembly of multimeric complexes following their binding to the peroxisome membrane (reviewed in Subramani, 1998; Hettema *et al.,* 1999; Terlecky and Fransen, 2000; Subramani *et al.,* 2000; Purdue and Lazarow,

2001; Titorenko and Rachubinski, 2001; Schleibs and Kunau, 2004). Recent evidence from Gould's lab has suggested that Pexl9p could be bifunctional acting both as a cytosolic PMP chaperone and as a PMP import receptor (Jones *et al.,* 2004). In the light of many other evidences, whether  $Pex19p$  functions as an import receptor or an assembly factor, remains unclear (Fransen *et al.*, 2001, 2004; Snyder *et al.*, 2000; Biermanns and Gartner 2001; Wang *et al.,* 2004; Lambkin and Rachubinski 2001; Otzen *et al.,* 2004). Understanding the exact functional role of Pex19p needs more investigation and our results suggest that Pexl9p interacts with the cytosolic Vpslp and that it functions as an assembly factor. Upon certain maturation signals, Vps1p is recruited to the peroxisomes and Pexl9p stabilizes this association resulting in organelle fission. It still remains to be determined if the GTPase activity of Vps1p signals division.

The challenge remains to understand how the increasing number of proteins shown to be involved in controlling peroxisome number and size interplay amongst themselves and signal to the cell how to control its peroxisome dynamics.

**CHAPTER SEVEN**

**PERSPECTIVES**

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## **7.1 Synopsis:**

Peroxisome biogenetic pathway conceptually requires at least few distinct processes: the formation of lipid bilayer, the insertion of membrane proteins into this lipid bilayer, the import of soluble proteins across the membrane into the matrix and the division and segregation of the mature peroxisomes to the daughter cells. In this study, different experimental approaches were made to identify and characterize novel components involved in peroxisomal assembly, especially in the division and proliferation of peroxisomes. The peroxisomes of cells deleted for one or both of the *PEX28* and *PEX29* genes are more abundant and smaller, and show extensive clustering, as compared to wild-type peroxisomes. Cells deleted for two of the *PEX30, PEX31* and *PEX32* genes contained increased numbers of generally enlarged peroxisomes.

Additionally, we have analyzed the assembly of membrane proteins using Pex30p and Pex32p as experimental candidates. This has revealed that Pexl9p acts as a targeting factor for PMPs. Our approach to understand the role of Pexl 9p in membrane biogenesis has revealed that the dynamin like protein Vps1p associates with peroxisomes in a Pexl9p-dependent manner. The implications of these data in peroxisomal dynamics are discussed in the following section and represented in the model 7-1.

#### **7.2 Fusion or fission mutants?**

Upon completion of peroxisome division, peroxisomes must be separated from one another. Pex28p and Pex29p are two proteins required for this process, as their absence leads to an arrest or retardation of the peroxisome proliferation pathway, leading to the



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Figure 7-1. A composite model for protein targeting and the dynamic events that occur during the assembly of peroxisomes. PPV-Pre-Peroxisomal Vesicle, PPT-Pre Peroxisomal Template.

presence of clusters of peroxisomes with evidence of thickened membranes sometimes occurring between adjacent peroxisomes. The thickened membranes between some peroxisomes are also suggestive of a role for Pex28p and Pex29p in controlling fission of the peroxisomal membrane. One of the important aspects that needs to be scrutinized is that if they are peroxisomal fission or fusion mutants. So far, no study has identified components of peroxisomal fission or fusion mutants. One way to study this would be to mate a and  $\alpha$  strains of the same mutant (for example  $pex29\Delta$ ) carrying different peroxisomal fluorophores (pGAL-Dsred-SKL and pGAL-GFP-SKL) and monitor the fusion events by time lapse imaging. If *PEX29,* is a fusion mutant, then we would observe clustering of red and green structures. If it is not affected in fusion, then we would observe yellow punctate structures.

Similar approaches can be made to study mutants of *PEX30*, *PEX31* and *PEX32*. Especially, the observation of increased number of peroxisomes in  $pex30\Delta$  is suggestive o f a fusion mutant phenotype. Fusion and fission events might require more than one component. Identification of more components will facilitate to resolve the chronological and spatial arrangement of protein complexes during these processes. A more straightforward approach will be based on immunoprecipitation studies. Tandem affinity purification can be used to isolate protein complexes; however, to understand the dynamics of interaction, cell biological approaches using fluorescent reporter proteins will be of immense use.

### **7.3 Transcriptional regulation of PEX genes**

Pex30p-prA, Pex31p-prA and Pex32p-prA were all detected in glucose-containing YEPD medium at the time of transfer. The levels of Pex30p-prA and Pex32p-prA increased with time of incubation of cells in YPBO medium, but not as dramatically as the levels of Potlp (peroxisomal thiolase). The levels of Pex31p-prA did not show any apparent increase with time of incubation of cells in YPBO medium. The promoter regions of *PEX30, PEX31* and *PEX32* contain sequences that resemble the canonical sequence  $CCGN_3TNAN_{8-12}CGG$  of the oleic acid response element (ORE) (Rottensteiner *et al.*, 2002; 2003a), which acts to increase gene transcription in S. *cerevisiae* in the presence of oleic acid as a carbon source through the binding of the transcription factors Pip2p and Oaf1p (Rottensteiner *et al.*, 1996; Karpichev *et al.*, 1997). It will be interesting to verify if these sequences actually do function as OREs.

#### **7.4 Regulated division versus constitutive division**

Peroxisomes undergo either constitutive division to maintain the number of peroxisomes or a regulated division when there is an external stimuli. External stimuli can be oleic acid and methanol for yeasts or peroxisomal proliferators for mammalian cells. Pex11p has been suggested to play a role in both of these processes (Li and Gould, 2002). However, whether the machinery involved in these two processes is one and the same remains unclear. Pex30p is a constitutively produced protein and its levels are not altered by external stimuli, while Pex32p is induced upon shifting the cells to oleic acid containing medium. Two hybrid analysis has revealed that these two proteins interact

with each other. The significance of this interaction in constit machinery of peroxisomal division during these different processes n Also, since both these proteins are polytopic in nature, it will be nec the topology of these proteins to enhance our understanding of the co similar situation can be observed in the Pex11 family of proteins. W. Pexl lp and Pex25p are induced, synthesis level of Pex27p is not ir Pex25p interact within themselves and each other in a two hybrid an the spatial arrangements of these proteins and the nature of the com will elucidated the machinery involved in the two types of division.

### **7.5 The enigma of peroxisomal subforms**

There had been significant evidence that, in a single cell, per heterogeneous population of organelles that differ in their size, buoy composition and protein import capacity. Titorenko and Rachubinski demonstrated in the yeast *Y. lipolytica* that this heterogenous populat assembles in a systematic multi step pathway. On the basis of studies *pastoris* and human fibroblasts, different models, although with 1 intermediates, provides a unified support for a multi step processi selective uptake of different matrix and membrane proteins into di: along the peroxisomal biogenetic pathway. A major challenge for tl define the applicability of the models presented in, to various model *Saccharomyces cerevisiae.*

Significant amount of PexSOp and Pex29p were always present in the less dense fractions during the gradient isolation of peroxisomes. Of particular interest is that these two proteins interact with each other on a two hybrid analysis. Therefore it remains to be determined if some portion of these proteins form a complex and represent components *en route* to peroxisomes. Preliminary results in *S. cerevisiae* suggest existence of at least four different bonafide compartments containing different peroxisomal proteins (Figure 7-2). This is extremely exciting, as the mechanism of peroxisomal assembly has so far been an unsolved mystery. It still remains to be determined if these compartments assemble in a systematic pattern or if they randomly fuse to form a mature peroxisome. This has opened a new avenue for further future research challenging the old model of peroxisome biogenesis. In depth investigation requires studies on the molecular mechanisms regulating the formation, selective import competency, division and conversion of distinct intermediates along the peroxisome assembly pathway.

#### **7.6 Are there multiple m-PTS signals on the same PMP?**

Systematic truncation of the integral membrane proteins Pex30p and pex32p from the C-terminus region has led to the identification of a m-PTS signal for these proteins. However, a number of studies have demonstrated that multiple targeting signals are present on the same protein. Especially, recent evidence from Goodman's laboratory suggest that multiple targeting modules on peroxisomal proteins are not redundant and that under different metabolic states, different mPTSs could function independently and



•**Figure** 7-2. Subforms of peroxisomes? Lighter fractions where Pex30p and Pex29p constantly appeared were floated on a two step gradient, pelleted and processed for electron microscopy. (A) Fractions of Pex29p. (B) Fractions of Pex30p.

efficiently (Wang *et al.,* 2004). In *Candida boidinii* Pexl9p binding to PMP47 is oleic acid induced and that in non-inducing conditions, another mPTS targets PMP47 to peroxisomes. Truncation of Pex30p and Pex32p from the N-terminus will help to identify such additional signals present in these proteins. Further investigation will be needed to verify if these signals are redundant and to determine the conditions under which they function.

#### **7.7 Can Pexl9p function as a cytosolic PMP chaperone?**

Pexl9p is a mostly cytosolic protein with a small fraction of it associated with the peroxisomes. The cytosolic fraction of Pexl9p has been suggested to act as a shuttling receptor for peroxisomal membrane proteins, while its peroxisome-associated form could function as a chaperone, assisting the assembly of multimeric complexes following their binding to the peroxisome membrane (reviewed in Purdue and Lazarow, 2001; Titorenko and Rachubinski, 2001; Schleibs and Kunau, 2004). Recent evidence from Gould's lab has suggested that Pexl9p could be bifunctional acting both as a cytosolic PMP chaperone and as a PMP import receptor (Jones *et al.,* 2004). Snyder *et al.,* (2000), have shown in *Pichia pastoris* that upon blocking protein synthesis followed by cross linking, the amount of PMP immunoprecipitated by Pex19p remained unchanged like the untreated cells and thus have concluded that Pexl9p interacts with preexisting and not with newly synthesized pool of PMPs. For Pex19p to function as a cytosolic chaperone, it must bind nascent peroxisomal proteins. Hence, biochemical analysis can be done to isolate complexes from the cytosol that contains Pexl9p as a component.

#### **7.8 Which protein recruits Vpslp to the peroxisome?**

The results of our experiments are suggestive of transient recruitment of Vps1p to the surface of the peroxisomes and this association is stabilized by Pex19p. The components involved in the recruitment of Vps1p to the peroxisomal surface are unknown. Hence, biochemical analyses need to be undertaken to identify the interacting partners of Vps1p. If these interacting partners have a role to recruit Vps1p, peroxisome morphology will be affected in the mutants of these genes. Alternatively, association of Vpslp-GFP chimeras or Vpslp-prA with the peroxisomal compartment can be monitored in the peroxisomal fission mutants or the interacting candidates.

Also, Pex11p has been suggested to indirectly recruit DLP1, a mammalian homolog of Vps1p, to the peroxisomes, although the authors were unable to detect any physical interaction between these two proteins (Li and Gould 2003). However, since we know Pexl lp is recruited in a Pexl9p dependent fashion (Rottensteiner *et al*., 2004), we speculate that V ps 1 p recruitment could perhaps be coordinated through both these proteins. This is interesting because, deletion of *PEX11* has similar phenotype as the deletion of *VPS1*. Hence, tracking the Vps1p-GFP chimera in a  $pex11\Delta$  mutant might reveal interesting data.

### **7.9 How does V pslp regulate fission?**

Vpslp is a dynamin like protein containing a tripartite GTP-binding domain towards its N-terminus and a GTPase effector domain towards its C-terminus. How may Vps1p regulate peroxisomal fission is unclear. Experiments can be easily designed to understand the mechanism of this division. Essentially, expression of Vps1p-GFP chimera, under the *GAL* promoter using time lapse imaging can be monitored to visualize fission events in a strain with a peroxisomal reporter (DsRed-PTSl). This double labeling can show the formation of green ring like structures around the red (elongated?) peroxisomes. Similar observations were made in mitochondria] division.

Although Vps1p could be selectively recruited to the dividing peroxisomes, we believe that the GTPase activity of Vps1p could act as a signaling event in the fission of peroxisomes. This is supported by the fact that V ps l p has also been shown to regulate vacuole fusion (Christopher *et al.*, 2004) and Mgm<sub>l</sub> p (another dynamin like protein) has been shown to regulate mitochondrial fusion. Hence further investigation is needed to test this hypothesis.

#### **7.10 Is there a dynamin independent mechanism of peroxisome fission?**

The observation that peroxisomes manage to divide at cytokinesis in *vpsJA*, suggest that there is also dynamin-independent peroxisome fission (Hoepfner *et al.,* 2001). In fact, this is re-inforced by our previous finding that more wild type peroxisomes were formed upon overexpression of PEX11 in a *vps1* $\Delta$  mutant (Vizeacoumar *et al.*, 2003). Vps1p has been shown to regulate the cytoskeleton through its interaction with Slalp (Yu and Cai, 2004). Also a double deletion mutant of *vpsl* $\Delta$ *rhol* $\Delta$  has been found to accumulate actin patches on peroxisomal membranes. These findings instigate us to see if there is a dynamin independent mechanism of peroxisome division.

## **7.11 Conclusion**

The processes of protein import, membrane fission and fusion are important events that occur in all eukaryotes. Studies that focus on similar events in peroxisomes will definitely contribute to the understanding of fundamental processes that occur in membrane trafficking in the endocytic and secretory pathways. Prior to the work done in this thesis, only three components namely Pex11p, Pex25p and Pex27p were known to regulate peroxisome abundance. This thesis has broadened our understanding of peroxisome proliferation by identifying at least five more molecular players that coordinate this event. Work described here has opened new avenues of research, further understanding of which will produce exciting insights on organelle biogenesis.

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# **CHAPTER EIGHT**

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