A role for Vps1p, actin, and the Myo2p motor in peroxisome abundance and inheritance in *Saccharomyces cerevisiae*

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n vivo time-lapse microscopy reveals that the number of peroxisomes in *Saccharomyces cerevisiae* cells is fairly constant and that a subset of the organelles are targeted and segregated to the bud in a highly ordered, vectorial process. The dynamin-like protein Vps1p controls the number of peroxisomes, since in a *vps1Δ* mutant only one or two giant peroxisomes remain. Analogous to the function of other dynamin-related proteins, Vps1p may be involved in a membrane fission event that is required for the regulation of peroxisome abundance. We found that efficient segregation of peroxisomes from mother to bud is dependent on the actin cytoskeleton, and active movement of peroxisomes along actin filaments is driven by the class V myosin motor protein, Myo2p: (a) peroxisomal dynamics always paralleled the polarity of the actin cytoskeleton, (b) double labeling of peroxisomes and actin cables revealed a close association between both, (c) depolymerization of the actin cytoskeleton abolished all peroxisomal movements, and (d) in cells containing thermosensitive alleles of *MYO2*, all peroxisome movement immediately stopped at the nonpermissive temperature. In addition, time-lapse videos showing peroxisome movement in wild-type and *vps1Δ* cells suggest the existence of various levels of control involved in the partitioning of peroxisomes.

Introduction

Eukaryotic cells are compartmentalized into organelles that execute unique and essential functions. Two classes of organelles can be distinguished based on their origin. The first class comprises the autonomous organelles such as ER, mitochondria, and chloroplasts, and the second class comprises the organelles that are derived from the autonomous organelles, such as endosomes, lysosomes/vacuole, and secretory vesicles. It is of particular importance to segregate the set of autonomous organelles properly during each cell division. This is a reflection of the accepted rule that the autonomous organelles multiply by growth and division and cannot be regenerated by de novo synthesis (Nunnari and Walter, 1996; Warren and Wickner, 1996). How do peroxisomes fit into this picture?

Peroxisomes are small single-membrane-bounded organelles containing a set of enzymes allowing them to participate in cellular metabolism (van den Bosch et al., 1992). Loss of peroxisomal function is the cause of a number of diseases in man, ranging from relatively mild single enzyme deficiencies to severe syndromes in which the biogenesis of the whole organelle is compromised (Purdue and Lazarow, 1994). The severe peroxisomal biogenesis disorders (PBDs)* are caused by a loss of protein import into peroxisomes, marking most of the PBDs as “protein-trafficking” diseases. The identification of a large number of proteins required for trafficking of peroxisomal proteins now forms the basis for a detailed molecular analysis of the import process (Hettema et al., 1999; Subramani et al., 2000). In contrast, not much data are available as to how peroxisomes multiply and are inherited upon cell division. Current opinion holds that peroxisomes multiply by growth and subsequently divide by fission, and also that they constitute a branch of the autonomous organelle fam-

*Abbreviations used in this paper: CFP, cyan fluorescent protein; GFP, green fluorescent protein; Lat-A, Latrunculin A; PBD, peroxisomal biogenesis disorder; PTS1, peroxisomal targeting signal type I; SPB, spindle pole body; YFP, yellow fluorescent protein.
ily and are segregated from mother to daughter cell (Laz- 
arow and Fujiki, 1985). More recently, however, observa-
tions suggest that a peroxisome–ER connection exists, 
indicating that the ER might contribute to the biogenesis of 
peroxisomes. It has been postulated that peroxisomes can be 
formed from ER membranes (Erdmann et al., 1997; 
Titorenko and Rachubinski, 1998; Tabak et al., 1999). Al-
though in many cell types peroxisomes are present in appre-
ciable numbers and a stochastic segregation principle could 
suffice, all the evidence obtained thus far indicates that the 
proper distribution of all primary organelles between mother 
and daughter cells depends on specialized segregation pro-
cesses involving the cytoskeleton and motor proteins (Car-
lett and Weisman, 2000).

In fungi, two cytoskeletal systems involved in the organiza-
tion, maintenance, and segregation of cellular components 
have been identified: microtubules and the actin cytoskeleton. 
Microtubules are organized by the spindle pole body (SPB), 
the functional homologue of the microtubule organizing cen-
ter of higher eukaryotic cells. Since S. cerevisiae performs a 
closed mitosis without breakdown of the nuclear envelope, 
microtubules can be classified in nuclear microtubules and cy-
toplasmic astral microtubules (Byers and Goetsch, 1975; By-
ers, 1981). Nuclear microtubules are involved in assembly of a 
bi bipolar spindle and segregation of the chromosomes (Jacobs 
et al., 1988; Straight et al., 1997), astral microtubules function 
to position, move, orient, and finally segregate the nucleus be-
tween mother and daughter cell (Palmer et al., 1992; Sullivan 
and Huffaker, 1992; Carminati and Stearns, 1997; Shaw et 
al., 1997; Tirnauer et al., 1999; Hoepfner et al., 2000).
Whereas polarized vesicular movement was also shown to be 
dependent on astral microtubules in animal cells (Rogalski et al., 
1984; Schliwa, 1984; McNiven and Porter, 1986; Vale 
et al., 1986; Rindler et al., 1987) and filamentous fungi 
(Howard and Aist, 1980; Howard, 1981; Steinberg, 1998), 
nor such involvement could be demonstrated so far in S. cerevi-
siae (Jacobs et al., 1988; reviewed by Bretsch et al., 1994).

The actin cytoskeleton in yeast consists of two different 
structures, patches and cables. Patches are cortical actin–rich 
structures that generally cluster near regions of active secre-
tion and therefore mark sites of growth (Adams and Pringle, 
1984). Cables are long bundles of actin filaments that can 
span the whole cell (Adams and Pringle, 1984). Both types 
of actin structures behave in a cell cycle–dependent manner.

Figure 1. Peroxisome morphology as seen in GFP-PTS1–labeled cells. In wild-
type cells, up to nine individual round-
shaped peroxisomes of different size and 
signal intensity are discernible per cellular compartment. In this static 
picture no localization pattern is 
apparent. (B) Peroxisome morphology 
as seen in GFP-PTS1–labeled vps1Δ 
cells. The number of individual peroxi-
somes is significantly reduced. Most cells 
only carry one peroxisome. Peroxisomes 
appear as large tubular structures. In 
budded cells, the peroxisomal tubes 
frequently localize near the bud neck re-
region. Five Z-axis planes spaced by 0.8 
μm have been acquired and merged into 
one plane. (C) Dynamics of GFP-PTS1– 
labeled peroxisomes in wild-type cells. 
Before bud emergence, peroxisomes 
appear randomly localized (0°). After 
bud emergence, two to three peroxi-
somes localize to the growing tip and 
three remain in the mother cell body 
(10.5°). The clustered peroxisomes in the 
bud remain clustered at the growing tip 
and the peroxisomes in the mother cell 
maintain a relatively fixed position 
(36°–90°). Before cytokinesis, the peroxi-
somes in the bud spread and relocalize 
from the bud tip to the site of cytokinesis. 
Similar localization to the bud neck of 
peroxisomes can be observed in the 
mother cell body (141°). After cell separa-
tion, new bud emergence of the daughter cell is observed and the peroxisomes again relocalized to the growing bud tip (159°). The described 
characteristics can be better observed in the Video 1 (where this sequence was extracted from) and Video 2. (D) Dynamics of GFP-PTS1–labeled 
peroxisomes in vps1Δ cells. One large, round-shaped peroxisome localizes to the site of previous cytokinesis (0°). After new bud emergence, the 
peroxisome glides along the cell cortex to the site of the growing tip and transforms into an elongated structure that finally reaches into the bud neck 
and vigorously bends out of plane in the bud (25°–103°). After fission of the peroxisomal tube, two separated, round-shaped peroxisomes appear, 
one located in the mother and one in the daughter cell body (110°). After cytokinesis, the peroxisomes associated with the site of previous cell 
separation start to elongate and reposition to the new incipient bud sites (120°). After bud emergence and bud growth, the peroxisomes repeated the 
characteristic movements described for the previous cell cycle. The described characteristics can be better observed in the Video 3 (where this 
sequence was extracted from) and Video 4. Videos are available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1. Bars, 5 μm.
(Kilmartin and Adams, 1984; Ford and Pringle, 1991; for review see Madden and Snyder, 1998). Transport along the actin cytoskeleton of cargo such as mitochondria, vacuoles, and late Golgi compartment elements is dependent on myosin motor proteins (for review see Sellers, 2000). Myosins move in a defined directionality along the actin cytoskeleton, the polarity of which is apparent by the localization of the patches. So far, only plus end–directed myosins toward the patches have been described in S. cerevisiae.

In this study, we investigated the mechanisms supporting faithful inheritance of peroxisomes using in vivo time-lapse microscopy in S. cerevisiae. It represents a highly polarized cell in which vectorial transport of organelles and vesicles is an important feature in rapidly growing cells. We observed peroxisomal fission events and identified a mutant impaired in this process. In addition, we observed specific dynamics during peroxisomal segregation which suggests that this is a well controlled process. Furthermore, we found that peroxisome segregation requires a polymerized actin cytoskeleton and depends on the actin-associated motor protein, Myo2p.

Results
Peroxisomes segregate from mother to daughter cells
To investigate the partitioning strategy followed by peroxisomes we studied their behavior in budding yeast using time-lapse microscopy. Peroxisomes were marked with a green fluorescent protein (GFP) variant containing the peroxisomal targeting signal type I (PTS1), which is specifically imported into peroxisomes (Monosov et al., 1996; Hettema et al., 1998). Under the conditions of our experimental set up, cells contained on average 9 (3–15) fluorescent spots representing peroxisomes. Peroxisomes displayed an ordered migration behavior during the cell cycle and moved along the cortex before bud emergence (Fig. 1 A; Videos 1 and 2, with representative images shown in Fig. 1 C). A subset of peroxisomes then localized to the incipient bud site (Fig. 1 C, 0–10.5H11032). These peroxisomes subsequently moved into the nascent bud as soon as a new bud was clearly visible (Fig. 1 C, 10.5H11032). During bud growth, peroxisome dynamics were different in the mother cell and the bud. Whereas most peroxisomes in the mother cell body retained their cortical position (Fig. 1 C, 10.5’–90’), peroxisomes in the bud displayed very complex movements. Early during the bud-growth phase, peroxisomes clustered at the bud tip and also during this phase we observed peroxisomes passing through the bud neck as observable in the second cell cycle of the mother cell in Video 2. Later during the bud-growth phase peroxisomes spread over the whole bud cortex (Fig. 1 C, 10.5’–90’). Prior to cytokinesis and cell separation, subsets of peroxisomes frequently localized to the bud neck region (Fig. 1 C, 141’, Videos 1 and 2). We conclude from these observations that peroxisomes segregate during cell division in a well-defined series of events and that the average number of peroxisomes per cell is carefully maintained. Videos are available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1.

Vps1Δ deletion mutants contain a few giant peroxisomes
Based on recent work with mitochondria, members of the dynamin protein family are likely candidates for presumptive involvement in a process supporting peroxisome fission (van der Bliik, 1999, 2000; McNiven et al., 2000). The yeast genome contains three genes coding for dynamin-related proteins: Vps1p, Dnm1p, and Mgm1p. The deletion strains vps1Δ, dnm1Δ, and mgm1Δ expressing GFP-PTS1 were inspected for alterations in peroxisome morphology. Morphology and number was unaffected in dnm1Δ and mgm1Δ cells (unpublished data), however, vps1Δ cells showed only 1–3 peroxisomes per cell (Fig. 1 B). Moreover, the size of these peroxisomes was increased compared with those in wild-type cells (see below). To further analyze the larger GFP-PTS1–labeled structures, vps1Δ cells were subjected to immunogold electron microscopy using anti-GFP coupled to gold particles. Peroxisomal structures varied in morphology ranging from single enlarged organelles (un-
published data) to clusters and strings of small interconnected organelles as if they were caught in an abortive fission event (Fig. 2). We propose that multiplication of peroxisomes is dependent upon Vps1p in a manner analogous to the function of the other dynamins involved in mitochondrial fission and fusion. Attempts to locate Vps1p to peroxisomes gave suggestive results, but, due to the cytoplasmic abundance of the Vps1 protein, they remained inconclusive. For instance, association was observed in videos showing coordinated movement over several minutes of patches of the yellow fluorescent protein (YFP) signal of Vps1p-YFP with the cyan fluorescent protein (CFP) signal of CFP-PTS1–labeled peroxisomes (Fig. 3 and Video 5, available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1).

Segregation of peroxisomes is accurately controlled

We were surprised that in 500 observed vps1Δ/H9004 cells at least one peroxisome was present, although S. cerevisiae cells can tolerate the loss of all peroxisomes in certain pex mutants (Hettema et al., 2000). In vivo time-lapse imaging of 62 vps1Δ cells did not show a single case where insertion of at least one peroxisome into the daughter cell had failed (Videos 3 and 4, representative examples extracted from Video 3 are shown in Fig. 1 D). Frequently, we observed a tubular extension emerging from a single, large peroxisome present in the mother cell which passed through the bud neck. The extension was subsequently separated from the large peroxisomal structure resulting in two peroxisomal structures, one in the mother cell and one in the bud.

Together, these observations in wild-type and vps1Δ mutant cells imply that peroxisomes can divide in a Vps1p-dependent and -independent manner and that their distribution between mother and daughter cell is not a stochastic event, but instead depends on an as yet unidentified, highly controlled segregation machinery.

Peroxisomes move independently from microtubules

Since peroxisomes were reported to be microtubule-associated organelles in mammalian cells (Rapp et al., 1996; Wiemer et al., 1997), we tested whether peroxisomal movement depends on microtubules.

We first analyzed if peroxisomes colocalized in vivo with astral microtubules. Therefore, we used a GFP-Tub1 construct to label all microtubules (Straight et al., 1997) and YFP-PTS1 to highlight the peroxisomes (Monosov et al., 1996; Brocard et al., 1997). Although GFP and YFP could not strictly be separated by fluorescent imaging techniques, the two different structures were easily identified by their typical morphology. The GFP signal of astral microtubules was very weak, preventing masking of the intensely labeled peroxisomes by the GFP signal of a microtubule. We analyzed 50 cells in different cell cycle stages from log phase cultures, but no direct interactions of microtubules and peroxisomes were apparent (Fig. 4 A). To further verify that peroxisomal dynamics did not...
any characteristic alterations in peroxisome dynamics in persisted (Fig. 4 B). Initial time-lapse analysis did not reveal faithful segregation into mother and daughter cells per-
spindles, the early bud localization of peroxisomes and crotubules and completely mispositioned and misoriented Admas, 1998). In spite of the absence of long astral mi-
which are unable to generate long astral microtubules

Directional peroxisomal movements parallel actin cytoskeleton polarity

The independence of peroxisomes on microtubules suggested the second cytoskeletal system in S. cerevisiae to be involved in peroxisome dynamics: the actin cytoskeleton. Therefore, we tested if the cell cycle–dependent movement and localization of peroxisomes spatially and temporally paralleled the organization of the actin cytoskeleton. To label the actin cytoske-
leleton, we fused YFP to the actin-capping protein Cap2, which highlights actin patches but does not label actin cables (Waddle et al., 1996). Direct GFP fusion to actin (Act1) or other proteins in order to visualize actin cables has so far not been successful (Doyle and Botstein, 1996). In addition, we labeled the peroxisomes with CFP-PTS1.

We analyzed 50 Cap2-YFP, CFP-PTS1–labeled cells in different cell cycle stages. Z-axis scans performed through the cells using Cap2-YFP as a cortical reference revealed almost exclusive cortical localization of peroxisomes; however, although the general distribution of actin patches and peroxisomes was very similar, no tight colocalization of patches and peroxisomes was apparent (Fig. 5, A–C). Nevertheless, the cortical localization and relative distribution was indicative of association of peroxisomes with actin cables. To test this hypothesis we performed time-lapse analysis of cells expressing both Cap2-YFP and CFP-PTS1 in wild-type cells. Since myosin/actin cable–based cargo movements have only been reported to occur toward the patch-covered plus end (the barbed end) in S. cerevisiae (Brown, 1997), we hypothe-
sized that long range peroxisome migration on actin cables would be directed toward actin patches. To test this hypo-
hesis, we analyzed the movement of CFP-PTS1–labeled peroxisomes relative to Cap2-labeled actin patches in 38 wild-
type cells. Long range movement of actin patches (transloca-
tions >1 μm) in the mother cell body strictly followed the actin polarity (Videos 6–8, available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1; representative examples extracted from Video 6 shown in Fig. 5 D). Peroxi-
somes exclusively moved toward the selected bud site. In small budded cells, they generally moved toward the bud tip where they formed clusters (Fig. 5 D, 0°–70°). Although we observed oscillations back and forth, long range retrograde movement only occurred after the isotropic switch and re-
orientation of the actin cytoskeleton toward the mother–daughter junction, before cytokinesis (Fig. 5 D, 70°–145°). When the subsequent bud site was selected at the distal pole, the observed reorientation of the actin cytoskeleton was always accompanied by a change in the direction at which at least several peroxisomes moved (Fig. 5 D, 145°–175°).
Peroxisomes colocalize with actin cables
Using a mild fixation method and staining the cells with phalloidin-rhodamine, it was possible to stain actin structures without losing the GFP signal of stained peroxisomes (see Materials and methods and the acknowledgment). Peroxisomes in both the mother cell and in large buds before cytokinesis colocalized with actin cables (Fig. 6). In small budded cells, however, peroxisomes again are detectable on actin cable structures. Actin cable–based peroxisomes are especially apparent in the cell indicated with an arrow and represented in larger magnification in D. Bars, 5 μm.

Actin depolimerization abolishes peroxisomal movements
To obtain further support for actin-based peroxisomal movement we incubated GFP-PTS1-labeled cells with Latrunculin A (Lat-A). This drug sequesters actin monomers and thereby disrupts the complete actin cytoskeleton (Coue et al., 1987). As a control we used Cap2-YFP, CFP-PTS1–labeled cells. After incubation for 15 min in 200 μM Lat-A at 30°C, the Cap2 label lost its defined localization and completely dispersed throughout the cell (unpublished data). We used these conditions to study the effect of Lat-A on peroxisomal movements. We tracked movements of individual peroxisomes over 15 min in Lat-A–untreated and -treated cells (Video 9, available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1). Peroxisomal movements in Lat-A–treated cells were drastically reduced to minor oscillations in the mother cell as well as in the bud (Fig. 8).

In Lat-A–treated cells microtubule dynamics persisted whereas peroxisomal movements were completely absent (unpublished data). Together, our results demonstrate that peroxisomes colocalize with the actin cytoskeleton and that directed movement depends on a properly organized actin cytoskeleton.

Peroxisomal movement is dependent on the class V myosin Myo2
Movement along actin cables is mediated by myosin motors. In S. cerevisiae five myosin genes have been identified, two of class I, one of class II, and two of class V (for review see Sellers, 2000). Among these three classes, class V myosins are the
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most likely candidates for involvement in peroxisome segregation. They consist of an NH₂-terminal motor domain, a neck domain, a coiled-coiled domain involved in dimerization, and a COOH-terminal cargo binding domain (Cheney et al., 1993; for reviews see Hildebrandt and Hoyt, 2000; Reck-Peterson et al., 2000). The two class V myosins in S. cerevisiae are named Myo2 and Myo4p. Myo2 is an essential protein. Analyses of temperature-sensitive MYO2 alleles have shown that Myo2 is directly involved in vesicular transport, vacuolar inheritance, and nuclear spindle orientation (Johnston et al., 1991; Govindan et al., 1995; Hill et al., 1996; Santos and Snyder, 1997; Catlett and Weisman, 1998; Schott et al., 1999; Beach et al., 2000; Yin et al., 2000). Myo4 is required for transport of specific mRNAs from the mother cell to the daughter cell (Bobola et al., 1996; Long et al., 1997; Takizawa et al., 1997). We expressed GFP-PTS1 in wild-type MYO2, myo2–12, myo2–16, myo2–20, myo2–66, and myo4Δ mutants (Johnston et al., 1991; Schott et al., 1999).

myo2–66 cells already at the permissive temperature were shown to be defective in vacuolar dynamics (Catlett and Weisman, 1998). When analyzing the localization of peroxisomes in the myo2–66 cells we observed a striking delay in insertion of peroxisomes into the bud (Fig. 9). 87% of all small-budded myo2–66 cells lacked peroxisomes in the bud, whereas this was the case in only 12% of wild-type cells (n = 250). Analysis of the other myo2-ts and the myo4Δ cells did not show significant alterations in tip localization of peroxisomes.

In addition, we tracked peroxisomal dynamics over a time interval of 15 min in the myo2-ts mutants at 24°C and after 10 min of incubation at 37°C. Typical examples of peroxisomal tracks are depicted in Fig. 10. Wild-type cells showed indistinguishable dynamics at both temperatures similar to those described above. This was completely different when analyzing the dynamics of the myo2-ts cells. At 24°C peroxisomal distribution appeared indistinguishable from wild-type and fast movements occurred frequently in the buds, but at 37°C only minor oscillations of peroxisomes were detectable. This effect is reversible as cells shifted back to 24°C for 30 min resulted again in normal peroxisome movement.

In contrast to the myo2-ts mutants, the myo4Δ mutant did not display any observable alterations in peroxisomal dynamics. We conclude that peroxisomes migrate along actin cables using the class V myosin, Myo2.

Discussion

During the cell cycle, organelles must be accurately multiplied and partitioned to daughter cells (Nunnari and Walter,
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Figure 9. Peroxisome distribution in small budded cells at 24°C as seen in GFP-PTS1–labeled wild-type and myo2–66 cells. (A) Peroxisome insertion into the nascent bud at very early stages is typical for wild-type cells. (B) In the myo2–66 mutant most cells do not localize peroxisomes into the bud at early cell cycle stages. In addition, the cells accumulate fluorescent vacuoles not observed in the wild-type cells. Bar, 5 μm.

1996; Warren and Wickner, 1996; Yaffe, 1999). This is particularly so for the autonomously replicating organelles comprising nucleus/ER, mitochondria, and chloroplasts. Although other organelles comprising Golgi complex, endosomes, lysosomes/vacuole, and secretory vesicles are derived from the ER it becomes increasingly clear that special care is taken to ensure their efficient segregation too. Here we have shown that peroxisomes follow the same rule. Peroxisomes were visualized by transforming yeast cells with a gene coding for GFP with the PTS1 appended to its COOH terminus, which is efficiently imported into the organelles. This allowed us to follow their partitioning between mother and daughter cells for several generations by time-lapse microscopy. Under the conditions of growth cells maintained on average a number of nine peroxisomes. This apparently involves an active fission process that results in the formation of additional peroxisomes. We infer this from our observation that most cells with a deleted VPS1 gene maintain only a single enlarged peroxisome. Vps1p is a member of the dynamin protein family whose members are known to function in vesicle trafficking events involving the ability to constrict and sever membrane tubules into discrete vesicles (van der Bliek, 1999; McNiven et al., 2000). In yeast, Dnm1p and Mgm1p participate in fission of mitochondria (Otsuga et al., 1998; Sesaki and Jensen, 1999; Shekarchi and Yaffe, 1999; Mozdy et al., 2000; Tieu and Nunnari, 2000; Wong et al., 2000), whereas Vps1p is required for sorting of proteins to the vacuole (Rothman et al., 1991; Wilsbach and Payne, 1993). It is interesting that mitochondria, with their endosymbiotic history, have their own dedicated dynamins, whereas peroxisomes share a dynamin-like function with the vacuolar compartment. It adds additional thought to recent ideas that peroxisomes are semiautonomous organelles which are at least in part dependent on contributions from the ER (Tsorten and Rachubinski, 1998). This contribution may be particularly important in terms of donating the phospholipid membranes to the peroxisomal compartment.

The constant number of peroxisomes per cell, the correct segregation of peroxisomes to the buds, and the rather precise control over the number of peroxisomes retained by the mother and delivered to the bud suggest the existence of various components taking part in peroxisome partitioning. In this respect, peroxisomes resemble mitochondria. Here a “retention zone” in the mother cell distal from the bud was defined in which a subset of mitochondria was retained by interaction with the actin cytoskeleton (Yaffe et al., 1999). Precise partitioning is even more obvious in the vps1Δ mutant. In all cases we observed that the single remaining peroxisome was partitioned equally well between mother and daughter. Whether this is achieved by a remaining fission capacity of another protein than Vps1p or is in part an indirect, physical effect of pulling forces acting from opposite sites remains to be studied.

Moreover, during our time-lapse analysis it became apparent that in a few vps1Δ mutant cells early insertion of the peroxisomal compartment failed. These cells showed a striking delay in cytokinesis compared with the other cells in the time-lapse sequence. This observation is similar to those made in some mdm and vac mutants (Weisman et al., 1990; McConnell and Yaffe, 1993; Xu and Wickner, 1996). This suggests the existence of a surveillance mechanism preventing the onset of cytokinesis until all organelles are properly partitioned. The conditional dependency of S. cerevisiae cells on peroxisomes and the reduced number of peroxisomes in the vps1Δ mutant could provide a valuable basis for genetic screens to identify putative components of such an organelle checkpoint mechanism.

We have studied the interaction of Vps1p with peroxisomes using a Vps1-YFP fusion protein and CFP-marked peroxisomes. Microscopic analysis of the double-labeled cells showed moderate cytoplasmic localization of the Vps1-YFP

Figure 10. Analysis of peroxisomal dynamics in GFP-PTS1–labeled MYO2, myo2-ts, and myo4Δ cells at 24°C and 37°C by tracking of peroxisomal movements for 15 min. In all strains at 24°C, highly dynamic peroxisomal movements can be observed that are described in more detail in Fig. 1 C. At 37°C, MYO2 wild-type and myo4Δ deletion mutants exhibit peroxisomal movements similar to those observed at 24°C. The other mutants show significantly reduced peroxisomal translocations at 37°C and only minor oscillations are observable. A few peroxisomes in the myo2–66 mutants show nondirected translocations, probably due to detachment from the cell cortex and passive cytoplasmic random movement. Tracking of peroxisomes was based on 15 min time-lapse sequence with acquisition intervals of 15 s. 10 individual cells of each mutant at both temperatures have been analyzed and representative examples are shown in this figure.
protein with additional spotted accumulations, most likely in the Golgi and post-Golgi compartments as suggested in the literature (Rothman et al., 1990). Due to the disperse localization of Vps1p, minor fractions always seemed to colocalize with peroxisomes. Coordinated movement of Vps1p with individual peroxisomes occurred and persisted for several minutes. We surmise that Vps1p only transiently interacts with peroxisomes and that fission events of peroxisomes can occur without accumulation of high concentrations of the Vps1 protein at the peroxisome membrane.

The orderly, track-like movements of peroxisomes in dividing wild-type cells suggested the involvement of cytoskeletal components. All our evidence indicates that peroxisomes are bound in the mother cell by the cortical actin skeleton and that they actively slide along actin filaments to the site of bud appearance and subsequently into the bud itself. Our time lapse analyses of Cap2-YFP, CFP-PTS1–labeled cells revealed directed peroxisomal movement relative to the Cap2-labeled actin patches along the cell cortex strictly following the polarity of the actin cytoskeleton. In fixed cells stained for actin, all the GFP-PTS1 fluorescently labeled peroxisomes colocalize with actin polymers. The dynamic movements of peroxisomes in life cells were lost upon treatment of the cells with Lat-A, a poison that breaks down actin filaments. Finally, the protein responsible for tracking of peroxisomes through the cell is Myo2, a typical actin-based motor protein. This was inferred from the behavior of several thermosensitive alleles of Myo2. In all cases dynamic movements of peroxisomes stopped after bringing the cells to the nonpermissive temperature. In contrast, movements of peroxisomes after bringing the cells to the nonpermissive temperature.

Materials and methods

Strains, media, and yeast transformation

Yeast strains used in this study are listed in Table I. Yeast media were prepared as described by Guthrie and Fink (1991). The yeast transformation procedure was based on the protocol by Schiestl and Gietz (1989). After the heat shock step, cells were pelleted and resuspended in 5 ml of YPD and incubated for 2 h at 30°C. Cells were again pelleted, resuspended in 1 ml H2O and plated on selective YPD-G418 medium (200 mg/l geneticin). Lat-A (Molecular Probes) was added from a 200 mM stock in DMSO to exponentially growing YPD cultures (final concentration 200 μM). Further incubation for 15 min at 30°C was performed on a roller drum and the cells then analyzed by microscopy. E. coli XL1-blue (Bullock et al., 1987) was used to propagate plasmids.

DNA manipulations, cloning procedures, and strain constructions

DNA manipulations were performed as described by Sambrook et al. (1989). We applied a PCR-based method to construct gene deletion cassettes for transformations (Wach et al., 1994). DNA of E. coli plasmids pFA6-AkMX4 (Wach et al., 1994) and pFA6a-YFP-His3MX6 (Yeast Resource Center) served as template for preparative PCR reactions. Genomic integration of the corresponding construct was verified by analytical PCR (Huxley et al., 1990; Wach et al., 1994). Yeast strains were grown on YPD-G418 (200 mg/l geneticin) to select for transformants that had integrated KanMX4, or GFP-kanMX6 cassettes. Growth on SD plates lacking histidine, leucine, or tryptophane selected for YFP-HIS3MX6 or YIp-derived constructs (see below) integration. Plasmids pEW161, pEW171, and pEW213 were constructed as follows. pEW161: PCR amplification of the HIS3 promoter with 5'-GGGATCC-TATTACCCTGGCCTCTCC-3' introducing an EcoRI site at the 5' end and a SacI site at the 3' end with primer 5'-GGGACATGAAAGGCAGGAAAGAGG-3'. The fragment was subsequently introduced into YIp lac204 (Gretz and Sugino) (pEW156) and YIp lac211 (pEW157). GFP-PTS1 (from pEW88: Hettema et al., 1998) was cloned into pEW156 and pEW157 cut with BamHI and HindIII resulting in pEW160 and pEW161, respectively. pEW171: the Ncol-BstBI fragment GFP-PTS1 derived from pEW160 was swaped with that of CFP or YFP. pEW213: genomic Vps1 was PCRred with primer 5'-GGGATCC-TATTACCCTGGCCTCTCC-3' and primer 5'-GGGACATGAAAGGCAGGAAAGAGG-3'. This fragment has been ligated in frame with YFP in YIp lac211 using EcoRI and PstI.

For gene deletions we followed the EUROFAN guidelines (guidelines for EUROFAN B0 program ORF deletants, plasmid tools, and basic functional analyses are available at www.mips.biochem.mpg.de/proj/eurofan/index.html) for gene replacement in S. cerevisiae. We used the oligonucleotide pairs depicted in Table II for generation of the KanMX4, deletion cassettes, or YFP fusion constructs. To label microtubules we integrated plasmid pAFS125 into the ura3 or the tpi1 locus (Straight et al., 1997). To label peroxisomes we integrated pEW161 or pEW177 into the ura3 or the tpi1 locus. Microtubules or peroxisomes were clearly observable under the fluorescence microscope upon successful transformation. Label-
ing of Vps1 was performed by integrating pEW213 into the ura3 locus. Correct integration rescued the peroxisomal morphology defect and the temperature sensitivity of vps1A strains.

**Microtubule depolymerization**
Microtubules were depolymerized by adding 2.5 μM/ml culture of a 5 mg/ml stock solution of either nocodazole or benomyl (both Sigma-Aldrich) to growing cells. Microtubules could be identified, whereas a normal array of microtubules was present in DMSO-treated control cells.

**Acquisition of still images**
YFP-PTS1, CFP-PTS1, Cap2-YFP, and GFP-Tub1–labeled wild-type and strains were described in Schott et al., 1999. The video microscopy setup and in vivo time lapse procedures using GFP-PTS1, CFP-PTS1, and GFP-Tub1–labeled strains were described by Hoepfner et al. (2000). Imaging of the YFP/CFP variant of GFP was performed using filter sets 41028 and 31044v2 (Chroma Technology Corp.).

**In vivo microscopy procedures and techniques**
The video microscopy setup and in vivo time lapse procedures using GFP-PTS1, CFP-PTS1, Cap2-YFP, and GFP-Tub1–labeled strains were described by Hoeprchner et al. (2000). Imaging of the YFP/CFP variant of GFP was performed using filter sets 41028 and 31044v2 (Chroma Technology Corp.). For long term in vivo time-lapses we used diploid cells because spreading of the cells was better due to bipolar budding. Acquisition settings like interval time, exposure time, excitation light transmission, the number, and spacing of the z-axis planes are indicated in the video legends. To analyze the temperature-sensitive mutants at the nonpermissive temperature we used a temperature-adjustable stage (Biowerk). Acquisition and processing had to be optimized for each strain.

### Table I. Yeast strains used in this study

| Strain | Genotype | Source |
|--------|----------|--------|
| BJ1991 | MAT_ura3-251 leu2 trp1 prb1-1122 pep4-3 gal2 | Jones, 1977 |
| BSL1-2c | MAT_ura3-251 His4-519 leu2-3 leu2-112 | B. Lemire |
| FY 1679 | MAT::ura3-52::TRP1 leu2-112/LEU2 his3-Δ200::HIS3 | B. Dujon |
| DHY 339 | MAT_ura3-251::GFP-PTS1-URA3(pEW161) trp1 leu2 prb1-1122 pep4-3 gal2 | This study |
| DHY 340 | MAT_ura3-251::GFP-PTS1-URA3(pEW213) trp1 leu2 prb1-1122 pep4-3 gal2 | This study |
| DHY 341 | MAT_ura3-251::GFP-PTS1-URA3(pEW231) trp1 leu2 prb1-1122 pep4-3 gal2 | This study |
| DHY 342 | MAT_ura3-251::GFP-PTS1-URA3(pEW231) trp1 leu2 prb1-1122 pep4-3 gal2 | This study |
| DHY 343 | MAT_ura3-251::GFP-PTS1-URA3(pEW171) leu2 prb1-1122 pep4-3 gal2 | This study |
| DHY 344 | MAT_ura3-251::GFP-PTS1-URA3(pEW213) trp1 prb1-1122 pep4-3 gal2 | This study |
| DHY 345 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 346 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 347 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 348 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 349 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 350 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 351 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 352 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 353 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 354 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 355 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 356 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 357 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 358 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 359 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 360 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 361 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 362 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 363 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 364 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 365 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 366 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 367 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 368 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 369 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |
| DHY 370 | MAT::ura3-251::GFP-PTS1-URA3(pEW177) trp1-Δ63 leu2 Δ11 his3 Δ200 | This study |

### Table II. Oligonucleotides used in this work

| Oligonucleotide | Sequence |
|----------------|----------|
| vps1Δ1 | 5’-ACCACAATCAAGCACCCGATGAAAATGGTCGACATATTATATTAGAAT'C-3' |
| vps1Δ2 | 5’-TTTGCGATCATTTATATATATTATGATATTATAGAAT'C-3' |
| Cap2Fus | 5’-CACTTAGTCTGTCATGTATTATTATGATAACACGGCATGAAT'C-3' |
of images was performed using the Metamorph 4.1 program (Universal Imaging Corp.). Acquired z-axis stacks were merged into one plane using the “stack arithmetic: maximum” command of Metamorph. Stored images were then scaled and converted to 8-bit files. False color look-up tables were assigned to the individual channels as indicated in the figure legends. The phase-contrast and fluorescence 8-bit planes were then overlaid using the built-in “Overlay” command with default color balance.

Online supplemental material

For time-lapse analysis we assembled the picture files to videos in QuickTime format (Apple Computer) with a frame rate of 10 frames per second using the Premiere 4.2 program (Adobe Systems Europe). The videos are platform-independent and can be viewed using QuickTime movie player that can be downloaded at www.apple.com. Explanatory remarks and detailed information about acquisition settings are shown in the individual video legends associated with the figures showing representative still images of the time-lapse sequences. Videos are available at http://www.jcb.org/cgi/content/full/jcb.200107028/DC1.

We thank Anthony Bretscher for sending us the myo2-ts strains, Aaron F. Straight for the pAFS125 plasmid, and the team at the YRC Microscopy Department for plasmids pH43 and pH45. We are grateful to Barbara Winters for suggestions about the phalloloidin-rhodamine/GFP staining and Ben Distel for stimulating discussions. This work was supported by grants from the University of Basel and the Swiss Federal Office for Education and Science as well as the Netherlands Foundation for Scientific Research (NWO).

Submitted: 9 July 2001
Revised: 17 September 2001
Accepted: 16 October 2001

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