Ergosterol is required for targeting of tryptophan permease to the yeast plasma membrane

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It was known that the uptake of tryptophan is reduced in the yeast erg6 mutant, which is defective in a late step of ergosterol biosynthesis. Here, we show that this is because the high affinity tryptophan permease Tat2p is not targeted to the plasma membrane. In wild-type cells, the plasma membrane localization of Tat2p is regulated by the external tryptophan concentration. Tat2p is transported from the Golgi apparatus to the vacuole at high tryptophan, and to the tryptophan concentration. Tat2p is transported from the Golgi apparatus to the vacuole at high tryptophan, and to the plasma membrane at low tryptophan. However, in the erg6 mutant, Tat2p is missorted to the vacuole at low tryptophan. The plasma membrane targeting of Tat2p is dependent on detergent-insoluble membrane domains, suggesting that sterol affects the sorting through the organization of lipid rafts. The erg6 mutation also caused missorting to the multivesicular body pathway in late endosomes. Thus, sterol composition is crucial for protein sorting late in the secretory pathway. Tat2p is subject to polyubiquitination, which acts as a vacuolar-targeting signal, and the inhibition of this process suppresses the Tat2p sorting defects of the erg6 mutant. The sorting mechanisms of Tat2p that depend on both sterol and ubiquitin will be discussed.

Introduction

Understanding of protein sorting mechanisms in the secretory pathway is important to address how organelles establish their identities. Proteins are sent to their destinations along the pathway by being loaded onto particular kinds of transport vesicles. In the case of membrane proteins, this is often achieved through recognition of cytoplasmic sorting signals by vesicle coat or adaptor proteins. It is also known that some membrane proteins have their sorting signals in the membrane-spanning regions. For a typical example, yeast ER proteins Sec12p and Sec71p contain Golgi-to-ER retrieval signals in their transmembrane domains, and the Golgi protein Rer1p acts as the sorting receptor by recognizing these proteins as specific ligands and sending them back to the ER (Sato et al., 1996, 1997, 2001). In a different view, it is also plausible that lipid interacts with transmembrane domain signals, and thus the lipid composition affects the localization of proteins. In support of this idea, several mechanisms of sorting by lipids have been postulated. A well-known example is the sorting mediated by a sphingolipid- and cholesterol-rich membrane domain called “raft” (Simons and Ikonen, 1997). These lipids tend to cluster in the bilayer to form microdomains, which are not solubilized by detergents. A special set of proteins such as influenza virus HA and GPI-anchored proteins, which are targeted to the apical plasma membrane in epithelial cells, are associated with lipid rafts. This association appears to depend on protein–lipid interactions in the bilayer. HA requires both its transmembrane domain and cholesterol for segregation into rafts (Scheiffele et al., 1997). One of the roles of rafts is proposed to be serving as sorting platforms that emerge in the trans-Golgi and move to the apical surface.

For gaining further insights into the sterol-dependent sorting processes, the yeast Saccharomyces cerevisiae is an attractive organism. The structure of the major sterol in yeast, ergosterol, is slightly different from cholesterol, but its biosynthetic pathway has been almost completely understood (for review see Daum et al., 1998). In terms of membrane trafficking, the sterol composition has been shown to affect endocytosis in yeast (Heese-Peck et al., 2002). Evidence is also presented that yeast does have lipid rafts that are important for protein sorting (Bagnat et al., 2000, 2001). To further understand the role of sterols in traffic, we decided to start a study paying attention to yeast erg mutants, which are defective in the ergosterol biosynthesis. We examined phenotypes of the erg mutants to find potential defects in protein sorting. We were aware that the erg6 mutant was known to show reduced uptake of tryptophan from the
medium (Gaber et al., 1989). The ERG6 gene encodes S-adenosylmethionine Δ24 methyltransferase, which acts as a late step of the ergosterol biosynthetic pathway by converting zymosterol to fecosterol. The tryptophan uptake defect raised the possibility that the high affinity tryptophan permease Tat2p (Schmidt et al., 1994) is not correctly targeted to the plasma membrane. In this article, we will show the results of our detailed analysis of the localization of Tat2p and its post-Golgi sorting, with a particular focus on the roles of ubiquitination and lipid raft association.

Results

Localization of Tat2p is regulated by the tryptophan concentration in the medium

Tryptophan-auxotrophic yeast trp1 strains were used in most experiments. Growth of trp1 cells depends on uptake of tryptophan from the medium, which is largely performed by Tat2p (Schmidt et al., 1994). Due to the reduced uptake of tryptophan (Gaber et al., 1989), Δerg6 cells were unable to grow at 20 μg/ml of tryptophan (Fig. 1), the standard concentration of this amino acid in synthetic media (Sherman, 1991). At a high concentration of tryptophan (200 μg/ml), the growth was restored. This severe tryptophan auxotrophy seemed to stem from the impaired function of Tat2p because overexpression of TAT2 by a multicopy YEpl vector completely restored the growth of Δerg6 cells even at a low concentration of tryptophan (2 μg/ml). This result gave us a warning that we should be very cautious about the expression level of Tat2p.

Three copies of the HA epitope or the GFP was appended to the COOH terminus of Tat2p, which is predicted to orient to the cytoplasm (Beck et al., 1999). The TAT2 own promoter was chosen to express these variants. Either TAT2–3HA or TAT2–GFP on a single-copy YCp vector could support the growth of Δtat2 cells even at low tryptophan (Fig. 2), indicating that these fusion proteins are functional. In the following experiments, YCp TAT2–3HA or YCp TAT2–GFP was introduced into cells of the Δtat2 mutant and Δerg6 mutant, which is defective in the endosomal compartment. The yeast equivalent of late endosomes (Piper et al., 1995). Double labeling with Pep12p, the yeast syntaxin that marks late endosomes (Becherer et al., 1996), showed that Tat2–3HAp and Pep12p were clearly colocalized in the punctate structures (Fig. 3 B, bottom). These results indicate that Tat2–3HAp is localized to late endosomes when the tryptophan concentration is high in the medium. ER localization of HA–Tat2p was reported before (Beck et al., 1999), but we consider that this was due to overproduction.

The endosomal localization of Tat2–3HAp may result from plasma membrane targeting and rapid endocytosis. However, in the Δend3 mutant, which is defective in the endocytic internalization (Raths et al., 1993), Tat2–3HAp was still not detected on the plasma membrane at high tryptophan (Fig. 3 C), indicating that Tat2–3HAp is directed to late endosomes without detouring to the plasma membrane.

When wild-type cells were shifted from high to low tryptophan medium, staining of cell periphery became evident.

<table>
<thead>
<tr>
<th>200 μg/ml (High Trp)</th>
<th>20 μg/ml (Standard Trp)</th>
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<tr>
<td>WT (vector)</td>
<td>Δerg6 (vector)</td>
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<td>Δerg6 (YEpl TAT2)</td>
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Figure 2. Epitope tagging of Tat2p. KUY121 (Δtat2) cells harboring the indicated plasmids were grown and spotted onto the low tryptophan medium as described in Fig. 1.
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(Fig. 3 D), indicating that Tat2–3HAp is now targeted to the plasma membrane. This is reasonable because under the low tryptophan condition, Tat2–3HAp should be on the plasma membrane for efficient uptake of tryptophan.

These results demonstrate that the plasma membrane localization of Tat2–3HAp is regulated by the tryptophan concentration in the medium. Tat2–3HAp is targeted to late endosomes at high tryptophan, and to the plasma membrane at low tryptophan.

**Tat2p is missorted to the vacuole in the Δerg6 mutant**

Next, we examined the localization of Tat2–3HAp in Δerg6 cells (Fig. 4 A). At high tryptophan, Tat2–3HAp was localized to punctate structures as in wild-type cells. However, when the Δerg6 cells were shifted to the low tryptophan medium, the staining of Tat2–3HAp did not change to the plasma membrane pattern, and the fluorescence within the cells became very faint.

The amounts of Tat2–3HAp were examined by immunoblotting (Fig. 4 B). In wild-type cells, a larger amount of Tat2–3HAp was detected when incubated in the low tryptophan medium for 2 h than when kept in high tryptophan. In the vacuolar proteinase–deficient Δpep4 mutant, a high level of Tat2–3HAp was detected regardless of the tryptophan concentration. Together with the localization, these results indicate the regulated sorting of Tat2–3HAp in the secretory pathway. At high tryptophan, Tat2–3HAp is transported to the vacuole via late endosomes and eventually degraded. The degradation is slowed down at low tryptophan because Tat2–3HAp is rerouted to the plasma membrane. In contrast, in Δerg6 cells, the amount of Tat2–3HAp was markedly reduced at low tryptophan (Fig. 4 B), consistent with the faint signal in the immunostaining. This reduction is due to vacuolar degradation because the disruption of PEP4 in Δerg6 prevented the loss of Tat2–3HAp at low tryptophan. Thus, in Δerg6 cells, Tat2–3HAp is missorted to the vacuole and quickly degraded under the low tryptophan condition.

The missorting of Tat2p implies that the severe tryptophan auxotrophy of Δerg6 can be suppressed if the vacuolar delivery is blocked. By using the pep12 mutation that inhibits the traffic to late endosomes and thereby redirects
vacuolar proteins to the cell surface (Becherer et al., 1996), we show this is indeed the case. As shown in Fig. 4 C, the \( \Delta \text{erg6} \) mutant did not grow below 100 \( \mu \text{g/ml} \) of tryptophan. However, this defect was clearly suppressed by \( \Delta \text{pep12} \), although not completely. The \( \Delta \text{erg6} \Delta \text{pep12} \) double mutant grew well at 100 \( \mu \text{g/ml} \) and slowly at 50 \( \mu \text{g/ml} \).

### Sorting of Tat2-GFP late in the secretory pathway

To follow the route of Tat2p in more detail, the localization of another fusion construct, Tat2-GFP, was examined in various mutants defective in late steps of the secretory pathway. The results are shown in Fig. 5. In wild-type cells grown at high tryptophan, Tat2-GFP was localized to the vacuole as well as to perivacuolar late endosomes. When the cells were shifted to low tryptophan, Tat2-GFP was localized to the plasma membrane. The advantage using Tat2-GFP in living cells is that the plasma membrane was much more clearly visualized than in the fixed cells by immunostaining (compare with Fig. 3 D). This is probably because the enzymatic removal of the cell wall can be omitted if Tat2-GFP is used. Again, relocalization of Tat2-GFP to the plasma membrane in response to low tryptophan was not observed in \( \Delta \text{erg6} \) cells, with prominent fluorescence in the vacuole.

The localization of Tat2-GFP was also examined in \( \text{TRP1} \) cells. The GFP fluorescence was clearly observed in these tryptophan prototrophs, indicating that Tat2p is expressed whether cells can synthesize tryptophan or not. Like in the \( \text{trp1} \) cells, the localization of Tat2-GFP in \( \text{TRP1} \) cells was regulated by external tryptophan. In wild-type \( \text{TRP1} \) cells, Tat2-GFP was localized to the vacuole and perivacuolar late endosomes in the high tryptophan medium, and targeted to the plasma membrane in the tryptophan-free medium. In contrast, when \( \Delta \text{erg6} \) \( \text{TRP1} \) cells were grown in the tryptophan-free medium, Tat2-GFP was not localized to the plasma membrane.

To test whether Tat2-GFP is targeted to the plasma membrane by the exocytic pathway, temperature-sensitive \( \text{sec} \) mutants were examined. \( \text{SEC14} \) is required for the exit from the Golgi (Stevens et al., 1982). When the \( \text{sec14} \) mutant was shifted to the low tryptophan medium and grown at the permissive temperature 23°C, plasma membrane localization of Tat2-GFP was observed. However, when the low tryptophan medium was kept at the nonpermissive temperature 37°C, Tat2-GFP was not targeted to the plasma membrane and stayed in intracellular compartments. \( \text{SEC6} \) encodes a component of the “exocyst” (TerBush et al., 1996), which is required for the fusion of Golgi-derived vesicles with the plasma membrane. At 37°C in the low tryptophan medium, Tat2-GFP was not targeted to the plasma membrane of the \( \text{sec6} \) mutant. These results

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**Figure 4.** Tat2–3HAp is missorted to the vacuole in the \( \Delta \text{erg6} \) mutant. (A) KUY153 (\( \Delta \text{erg6} \)) cells harboring YCpTAT2–3HA were grown in the high tryptophan medium. Cells were washed and shifted to high or low tryptophan for 2 h. Bar, 2 \( \mu \text{m} \). (B) KUY121 (WT), KUY154 (\( \Delta \text{pep4} \)), KUY153 (\( \Delta \text{erg6} \)), and KUY156 (\( \Delta \text{erg6} \Delta \text{pep4} \)) cells harboring YCpTAT2–3HA were grown as described in A. Cells were lysed and subjected to immunoblotting with the anti-HA antibody. (C) YPH500 (WT), KUY200 (\( \Delta \text{pep12} \)), KUY136 (\( \Delta \text{erg6} \)), and KUY204 (\( \Delta \text{erg6} \Delta \text{pep12} \)) cells were grown on MCD supplemented with uracil, adenine, and the indicated concentrations of tryptophan at 26°C.
appear to indicate that Tat2-GFP follows the exocytic pathway to the plasma membrane at low tryptophan.

In Δpep12 cells, Tat2-GFP was localized to the plasma membrane irrespective of the tryptophan concentration. Thus, the inhibition of the vacuolar delivery by Δpep12 resulted in constitutive plasma membrane targeting of Tat2-GFP. Consistent with the suppression of Δerg6 by Δpep12 (Fig. 4 C), Tat2-GFP was also targeted to the plasma membrane in Δerg6 Δpep12 cells.

The vacuolar protein sorting (VPS)* pathway represents the direct vesicular traffic from the trans-Golgi to late endosomes. The VPS1 gene product is involved in this pathway and is considered to be necessary for the vesicle formation from the trans-Golgi (Nothwehr et al., 1995). Unlike in the Δpep12 mutant, Tat2-GFP was not missorted to the plasma membrane in Δpsl1 cells at high tryptophan. Tat2-GFP was seen in the vacuole as well as perivacuolar dots. When the Δpsl1 cells were shifted to low tryptophan, plasma membrane localization of Tat2-GFP was observed. Δpsl1 did not suppress the severe tryptophan auxotrophy of Δerg6, either (unpublished data).

The result with Δpsl1 indicates that Tat2-GFP does not follow the normal VPS pathway to reach late endosomes at high tryptophan. The Δpep12 mutant is defective not only

*Abbreviations used in this paper: MVB, multivesicular body; MVL, mevalonic acid lactone; VPS, vacuolar protein sorting.
in the VPS pathway, but also in the endocytic pathway. Traffic from early to late endosomes is blocked in Δpep12 (Gerrard et al., 2000). Thus, Tat2p must have taken the route from the trans-Golgi to late endosomes at high tryptophan via early endosomes. We would suggest that the tryptophan-dependent sorting of Tat2p occurs in early endosomes, and the Δerg6 mutant is defective in this sorting process (see Discussion and Fig. 9).

Ubiquitination and sorting of Tat2p
Evidence is rapidly accumulating that ubiquitin acts as a sorting signal at multiple steps in post-Golgi traffic. In the case of the yeast general amino acid permease Gap1p, sorting is affected by its ubiquitinated status (Helliwell et al., 2001; Soetens et al., 2001). Polyubiquitination of Gap1p by the Rsp5p ubiquitin ligase complex results in sorting to the vacuole instead of the plasma membrane. In other words, polyubiquitin is recognized as a vacuolar-targeting signal. This prompted us to examine the ubiquitination status of Tat2p. Tat2–3HAp was immunoprecipitated from cells expressing myc-tagged ubiquitin (Hochstrasser et al., 1991), and the precipitated materials were detected with the anti-myc or anti-HA antibody. To prevent degradation of Tat2–3HAp in the vacuole, Δpep4 strains were used. As shown in

Figure 6.  
Ubiquitination and sorting of Tat2p. (A) KUY211 (Δpep4 TAT2; lane 1), KUY310 (Δpep4 TAT2–3HA; lane 2 and lane 3), and KUY314 (Δbul1 Δpep4 TAT2–3HA; lane 4) cells were grown in the high tryptophan medium. The strains harbored either pKU105 (Ub) or pKU106 (myc-Ub). In lane 5, KUY177 (sec14) cells harboring YCpTAT2–3HA and YEp105 (myc-Ub) were grown in the high tryptophan medium at 23°C, and then shifted to 37°C for 2 h. (B) KUY277 (Δbul1 Δtut2) cells harboring YCpTAT2–3HA were grown in the high tryptophan medium and shifted to low tryptophan for 2 h. Cells were processed for immunostaining with the anti-HA antibody. Bar, 5 μm. (C) YPH500 (WT), KUY136 (Δerg6), KUY266 (Δerg6 Δbul1), and KUY253 (Δerg6 Δdoa4) cells were grown in the high tryptophan medium. Cells were washed and adjusted at the density of 10^7 cells/ml. 5-μl aliquots of 10-fold serial dilutions were spotted on MCD supplemented with uracil, adenine, and the indicated concentrations of tryptophan. (D) KUY154 (Δpep4) and KUY156 (Δerg6 Δpep4) cells harboring YEp105 (myc-Ub) and a plasmid for the indicated variant of Tat2p were subjected to the detection of ubiquitination as described in A. Plasmids used to express Tat2p were YCpTAT2 in lane 1, YCpTAT2–3HA in lane 2, and YCpTAT2^{Ile8,3HAP} in lane 3 and lane 4.
the right panel of Fig. 6 A, Tat2–3HAp was specifically precipitated (compare lane 1 with lanes 2–4). High mol wt myc–ubiquitin conjugates of Tat2–3HAp were detected in Δpep4 cells grown under the high tryptophan condition, (Fig. 6 A, left panel, lane 3), indicating that Tat2–3HAp is polyubiquitinated. It is known that ubiquitination of cargo proteins, such as the yeast pheromone receptor Ste2p, occurs in the plasma membrane on endocytic internalization (Hicke and Riezman, 1996). However, because Tat2–3HAp does not take the detour to the plasma membrane by exocytosis and endocytosis under the high tryptophan condition (Fig. 3), the place of its polyubiquitination must be somewhere else. On the other hand, the polyubiquitination of Tat2–3HAp was not detected in sec14 (Fig. 6 A, lane 5), indicating that the polyubiquitination reaction takes place after Tat2–3HAp has left the Golgi.

BUL1 and BUL2 encode components of the Rsp5p ubiquitin ligase complex (Yashiroda et al., 1996, 1998). The deletion of these genes causes efficient plasma membrane delivery of Gap1p by decreasing its polyubiquitination (Helliwell et al., 2001). Similarly, we found that the myc–ubiquitin conjugation to Tat2–3HAp was markedly decreased by deletion of BUL1 (Fig. 6 A, lanes 3 and 4). We also examined the localization of Tat2–3HAp in Δbul1 cells and found that Tat2–3HAp was targeted to the plasma membrane even at high tryptophan (Fig. 6 B). At low tryptophan, Tat2–3HAp was localized to the plasma membrane very efficiently. Altogether, these results indicate that Tat2–3HAp is polyubiquitinated mostly by the Rsp5p–Bul1p ubiquitin ligase complex, and polyubiquitinated Tat2–3HAp is delivered to the vacuole without detouring to the plasma membrane. As has been reported for Gap1p (Soetens et al., 2001), the inhibition of polyubiquitination by Δbul1 would have dual roles for the marked accumulation of Tat2–3HAp in the plasma membrane; efficient targeting and inhibition of endocytic internalization.

Next, we examined the effect of Δbul1 on the tryptophan auxotrophy of Δerg6. Surprisingly, the Δerg6 Δbul1 double mutant could grow at as low as 10 μg/ml tryptophan (Fig. 6 C). Similarly, the deletion of DOA4, which reduces the efficiency of overall protein ubiquitination, also suppressed the severe tryptophan auxotrophy of Δerg6, although weakly. Δerg6 Δdoa4 cells could grow at 20 μg/ml of tryptophan (Fig. 6 C). These results led us to the hypothesis that Tat2p is inappropriately polyubiquitinated in Δerg6, resulting in the missorting to the vacuole.

There is another line of evidence that indicates aberrant polyubiquitination of Tat2p in Δerg6. Many lysine residues are present in the cytoplasmic domains of Tat2p, among which Beck et al. (1999) identified five lysine residues (10, 17, 20, 29, and 31) in the NH2-terminal domain as the ubiquitin acceptor sites on nutrient starvation. We confirmed that the three lysine residues (10, 17, and 20) are indeed the major ubiquitin acceptor sites of Tat2p. The variant of Tat2–3HAp, in which these three lysine residues were replaced by arginine (Tat2K10,17,20-3HAp), was little ubiquitinated in the Δpep4 background (Fig. 6 D, compare lane 2 and lane 3). However, in Δerg6 Δpep4 cells, Tat2K10,17,20-3HAp was again clearly ubiquitinated (Fig. 6 D, lane 4). The ubiquitination of Tat2p in Δerg6 must have occurred on improper lysine residues.

**Missorting to the multivesicular body sorting pathway in the Δerg6 mutant**

In immunofluorescence staining of Δpep4 strains to visualize vacuolar localization of Tat2–3HAp, we noticed an interesting difference between ERG6 and Δerg6 cells (Fig. 7 A). In Δpep4 cells, the vacuolar-limiting membrane was clearly stained, regardless of the tryptophan concentration. In contrast, in the Δerg6 Δpep4 cells, the fluorescence of Tat2–3HAp was not detected on the vacuole-limiting membrane, but almost exclusively in the lumen, either at high or low tryptophan. Such luminal staining would indicate that Tat2–3HAp entered the multivesicular body (MVB)–sorting pathway, which transfers a subset of cargo proteins to the invaginating vesicles in yeast late endosomes (Odorizzi et al., 1998). To test this possibility, we examined the effect of VPS27 disruption. VPS27 is one of the class E VPS genes, all of which are required for MVB formation (Odorizzi et al., 1998). As shown in Fig. 7 A, Tat2–3HAp was localized to the vacuole-limiting membrane and the exaggerated class E compartment in Δerg6 Δups27 Δpep4. These results indicate that Tat2–3HAp is efficiently sorted to the MVB pathway in Δerg6, regardless of the tryptophan concentration.

As shown in Fig. 6, inhibition of the ubiquitination by Δbul1 or Δdoa4 restored the tryptophan uptake in the Δerg6 mutant. Ubiquitin is also known to act as a sorting signal to the MVB (Katzmann et al., 2001), and the MVB sorting of cargo proteins is prevented by Δdoa4 (Losko et al., 2001; Reggiori and Pelham, 2001). We examined the effect of Δbul1 and Δdoa4 on the MVB sorting of Tat2–3HAp in Δerg6. The results are shown in Fig. 7 B. In contrast to Δerg6 Δpep4 cells, the vacuole-limiting membrane was clearly stained in Δerg6 Δbul1 Δpep4 and Δerg6 Δdoa4 Δpep4 cells, regardless of the tryptophan concentration. Plasma membrane staining at low tryptophan, which was not observed in Δerg6 Δpep4, was also appreciable in some Δerg6 Δbul1 Δpep4 and Δerg6 Δdoa4 Δpep4 cells, consistent with the suppression of the tryptophan auxotrophy (Fig. 6 C). Thus, the two defects of Tat2p sorting in the Δerg6 mutant, the cell surface–targeting defect and the MVB missorting, are simultaneously suppressed by the inhibition of ubiquitination.

In contrast to Tat2–3HAp, it may be noted that the fluorescence of Tat2-GFP is clearly seen in the vacuole lumen of wild-type cells (Fig. 5). We raised a specific antibody against Tat2p, and found that untagged Tat2p was detected on the vacuolar-limiting membrane in Δpep4, but in the vacuole lumen in Δerg6 Δpep4 (Fig. 7 C). This behavior is very similar to that of Tat2–3HAp, and therefore, the results with Tat2–3HAp may reflect the authentic nature of Tat2p. The sorting of Tat2-GFP into the vacuolar lumen was blocked in both Δpep4 and Δdoa4 cells (Fig. 7 D), indicating that it undergoes ubiquitin-dependent MVB sorting. Like in Δbul1 cells (Fig. 6 B), the plasma membrane signal of Tat2-GFP in Δdoa4 cells was obvious at high tryptophan, and become remarkable on the shift to low tryptophan.

Then why is Tat2-GFP efficiently sorted to the MVB in wild-type cells, even though it is functional and correctly targeted to the cell surface at low tryptophan? Tat2-GFP may be ubiquitinated more efficiently. Alternatively, vacuolar luminal
localization of the MVB vesicles could be detected more clearly by GFP fluorescence in living cells. Due to the fixation and subsequent permeabilization procedures, the MVB vesicles might look more obscure by immunofluorescence microscopy.

As another way to assess the MVB missorting in the \( \Delta \text{erg6} \) mutant, we looked at a different GFP marker, GFP-Pep12p. The results are shown in Fig. 7 E. As reported previously (Reggiori et al., 2000), GFP-Pep12p resides mostly on the vacuolar-limiting membrane in wild-type cells when overexpressed. However, in \( \Delta \text{erg6} \) cells, the GFP fluorescence was now evident in the vacuole lumen. The fluorescence on the limiting membrane still remained, indicating that GFP-
Pep12p is not completely relocated to the vacuole lumen. The luminal signal in Δerg6 cells disappeared by either Δups27 or Δdoa4. Thus, GFP-Pep12p in Δerg6 cells is also missorted to the MVB in a ubiquitin-dependent manner, indicating that a subset of cargo proteins is inappropriately ubiquitinated and then sorted to the MVB in Δerg6.

Transport of Tat2p to the plasma membrane depends on detergent-insoluble membrane domains

Because the deficiency of normal sterol in Δerg6 affected the sorting of Tat2p, we further investigated whether sterol-rich, detergent-insoluble membrane domains (so-called rafts) are involved in the plasma membrane delivery of Tat2p. Detergent insolubility of Tat2p was examined by treatment with CHAPS followed by a flotation analysis as diagrammed in Fig. 8 A. In wild-type cells (Fig. 8 B), a fraction of the GPI-anchored protein Gas1p floated to the interphase between 0 and 30% of OptiPrep™ (Fig. 8 B, fraction 2, arrowhead), whereas the vacuolar alkaline phosphatase Pho8p did not, as reported previously (Bagnat et al., 2000). Tat2–3HAp did not float to fraction 2 under the high tryptophan condition where it is sorted to late endosomes. However, Tat2–3HAp was clearly detected in fraction 2 under the low tryptophan condition where it is targeted to the plasma membrane. In the Δerg6 mutant (Fig. 8 C), Gas1p was still detected in the floating fraction 2. This is consistent with a recent report (Sievi et al., 2001), and indicates that the sterol intermediates accumulating in the Δerg6 mutant can replace ergosterol in the context of detergent-insoluble membrane domain formation. However, Tat2–3HAp in Δerg6 cells failed to float to fraction 2 even at low tryptophan (Fig. 8 C).

These results suggest that the association of Tat2p with the detergent-insoluble membrane rafts is required for plasma membrane delivery. Alternatively, the detergent insolubility of Tat2p could simply reflect the fact that rafts are the major lipid phase in the plasma membrane (Bagnat et al., 2001). We prefer the former possibility. First, Tat2–3HAp acquired the detergent insolubility even when its exit from the Golgi was blocked in the sec14 mutant (Fig. 8 D). When ER export was blocked by the sec12 mutation (Nakano et al., 1988), the detergent insolubility of Tat2–3HAp was not observed (Fig. 8 D), indicating that Tat2–3HAp is partitioned into lipid rafts after it reached the Golgi apparatus. Second and more importantly, when Tat2–3HAp was forced to localize to the plasma membrane in Δpex12 cells grown at high tryptophan, Tat2–3HAp was not found in the detergent-insoluble fraction (Fig. 8 D).

To confirm the role of rafts, we also attempted to disrupt the detergent-insoluble domain by inhibiting the initial step of the ergosterol biosynthesis. The ERG13 gene encodes HMG-CoA synthase. Δerg13 cells require mevalonate in the medium for growth. As reported before (Dimster-Denk et al., 1994), the growth of Δerg13 cells was arrested at a low concentration (5 mg/ml) of mevalonic acid lactone (MVL). At 10 mg/ml of MVL, Δerg13 cells were able to grow slowly. As shown in Fig. 8 E, only a small amount of Gas1p was found in the floating fraction 2 when MVL was supplied at 10 mg/ml. For simplicity, the distribution of Gas1p was compared between the detergent-insoluble (I) fractions (Fig. 8 A, defined as the mixture of fractions 2 and 3) and the detergent-soluble (S) fractions (Fig. 8 A, the mixture of fractions 7–9). As shown in Fig. 8 F, Gas1p gradually disappeared from the I fractions of Δerg13 cells according to the decrease of the supplementing MVL, indicating that the detergent-insoluble domains were significantly depleted when the flux of sterol synthesis was reduced. Exactly under the same condition, Tat2-GFP was inefficiently routed to the plasma membrane, and the vacuolar staining remained prominent (Fig. 8 G). In addition, Δerg13 cells were unable to grow at low tryptophan (Fig. 8 H), implying that the plasma membrane targeting of Tat2p is critical for growth under this condition. Strikingly, Δboll suppressed such severe tryptophan auxotrophy of the Δerg13 mutant (Fig. 8 H), indicating that the raft requirement for the plasma membrane targeting of Tat2p can be bypassed by the inhibition of polyubiquitination.

All these results strongly suggest that the partitioning of Tat2p into the rafts at low tryptophan is not the indirect consequence of the plasma membrane targeting, but is rather the cause of the sorting into the plasma membrane route. In other words, lipid raft sorting is very important for the cell surface delivery of Tat2p. However, it should be remembered that Tat2p can also be delivered to the plasma membrane in the absence of raft association under some conditions (for example, when the vacuolar trafficking pathway is blocked or the polyubiquitination is inhibited).

Discussion

In this paper, we present clear evidence that sterol is crucial for the correct sorting of the high affinity tryptophan permease Tat2p in yeast cells. The routes that we propose Tat2p follows are illustrated in Fig. 9. In the wild-type cells, Tat2p goes to the plasma membrane at low tryptophan, and to the vacuole via late endosomes at high tryptophan. The localization analysis of Tat2p in mutants defective in post-Golgi traffic indicates that Tat2p is delivered to late endosomes at high tryptophan not directly from the trans-Golgi nor via endocytosis from the plasma membrane. It is probably once delivered from the trans-Golgi to early endosomes, and there the tryptophan-dependent sorting occurs. These elaborate trafficking regulations of Tat2p are compromised in the Δerg6 mutant. Tat2p is not delivered to the plasma membrane, but to the vacuole at low tryptophan (Fig. 9, Δerg6, arrow 1). This is why Δerg6 cells are unable to take up tryptophan efficiently. In addition, Tat2p is missorted to the MVB pathway in late endosomes (Fig. 9, Δerg6, arrow 2).

Regulated transport of Tat2p

The activity of the Tat2p permease is controlled by the regulated sorting of membrane traffic rather than by synthesis. When and where the commitment is made as to whether Tat2p is transported to the vacuole or to the plasma membrane is a very important question, which turned out to be not an easy one. The experiments using Δend3, Δups1, and Δpex12 mutants indicated that Tat2p takes the unconventional route, trans-Golgi to early endosomes to late endosomes under the high tryptophan condition. On the other hand, it is not clear how Tat2p is routed to the plasma mem-
Figure 8. **Detergent-insoluble membrane domain is involved in the plasma membrane transport of Tat2p.** (A) The flotation procedure to monitor the detergent insolubility. Detergent-insoluble membrane domain is floated to the interface (arrow) that corresponds to fraction 2. Cells were subjected to the flotation analysis. In B–D, arrowheads indicate fraction 2, which contains detergent-insoluble membrane domains. (B) KUY121 (WT) cells harboring YCpTAT2–3HA were grown in the high tryptophan medium, washed, and shifted to low tryptophan for 2 h. (C) KUY153 (Δerg6) cells harboring YCpTAT2–3HA were grown and analyzed as described in B. The amount of Tat2–3HA at low tryptophan is significantly decreased in this mutant (see Fig. 4 B), but an enhanced image is shown here. (D) KUY177 (sec14), KUY197 (sec12), and KUY202 (Δpep12) cells harboring YCpTAT2–3HA were grown in the high tryptophan medium at 23°C. The sec mutant cells were washed, shifted to the prewarmed (37°C) low tryptophan medium, and incubated for 2 h. (E) KUY256 (Δerg13) cells harboring YCpTAT2–3HA were grown in the high tryptophan medium supplemented with 10 mg/ml MVL. (F) Distribution of Gas1p between the detergent-insoluble (I; mixture of the fractions 2 and 3) and soluble (S; mixture of the fractions 7–9) fractions. The high Trp samples of B–D were used. (G) KUY255 (Δerg13) and KUY257 (Δerg13 Δbul1) cells were grown in the high tryptophan medium supplemented with 50 mg/ml MVL. Cells were washed and adjusted at the density of 10⁸ cells/ml. 5-μl aliquots of 10-fold serial dilutions were spotted on the high or low tryptophan medium supplemented with 10 mg/ml MVL, uracil, and adenine.
Sterol is required for sorting of Trp permease | Umebayashi and Nakano

Ubiquitin-dependent sorting of amino acid permeases

It has been known that intracellular sorting of yeast nutrient transporters, such as Gap1p and the ferrichrome transporter Arn1p, is regulated late in the secretory pathway (Roberg et al., 1997; Kim et al., 2002). In the case of Gap1p, the sorting depends on the nitrogen source in the medium. Gap1p is targeted to the plasma membrane when cells are grown on urea, but to the vacuole in the glutamate medium. The similarity between Tat2p and Gap1p regarding nutrient-dependent regulation of sorting led us to suspect the presence of a common mechanism (i.e., ubiquitination).

Kaiser’s group has shown that polyubiquitination by the Rsp5p ubiquitin ligase complex directs Gap1p to the vacuole instead of the plasma membrane (Helliwell et al., 2001). This also turns out to be the case with Tat2p. Bul1p, known as a component of the Rsp5p complex required for elongation of polyubiquitin chains (Yashiroda et al., 1996; Helliwell et al., 2001), plays a critical role in the regulation of ubiquitination status of Tat2p. Polyubiquitination of Tat2p is little detected in the Δbul1 mutant. Surprisingly, almost all the defects of Δerg6 in Tat2p sorting are simultaneously suppressed by the knockout of BUL1. Presumably, aberrant polyubiquitination in the Δerg6 mutant is alleviated by the Δbul1 mutation. The anomaly of ubiquitination in Δerg6 is also seen on the acceptor sites of ubiquitin. Tat2p is polyubiquitinated on inappropriate lysine residues in Δerg6.

MVB sorting of Tat2p

Δerg6 cells also show a peculiar behavior in the MVB sorting. Although Tat2p remains on the limiting membrane when it is finally targeted to the vacuole in wild-type cells, Tat2p is almost completely segregated into the lumen of the vacuole in Δerg6. Similarly, Pep12p is also missorted into the MVB in Δerg6 cells. This MVB mistargeting is blocked by the class E Δeps27 mutation, suggesting that normal mechanisms of MVB sorting by the ESCRT complexes (Katzmann et al., 2001; Babst et al., 2002a, 2002b) are operating in this process. Interestingly, a CHO cell mutant defective in cholesterol biosynthesis also shows MVB missorting of the cation-independent mannose 6-phosphate receptor (Miwako et al., 2001). This kind of missorting may be a general outcome caused by defects of normal sterol synthesis.

Our finding that the MVB missorting of Tat2p and Pep12p in Δerg6 is suppressed by either Δbul1 or Δdoa4 indicates that it occurs in a ubiquitin-dependent manner. The sequential sorting defects of Tat2p in Δerg6, namely in early and late endosomes, could be explained solely by ubiquitination. That is, Tat2p is inappropriately ubiquitinated in Δerg6, delivered from early to late endosomes, and then sequestered into the MVB by being caught by the ESCRT-1 complex, a putative sorting receptor for ubiquitinated cargoes (Katzmann et al., 2001).

In several cases, monoubiquitination has been shown sufficient for the entry of cargo into the MVB. Tat2p is polyubiquitinated even under the low tryptophan condition in Δpea4 cells (unpublished data). Beck et al. (1999) have also shown that Tat2p is polyubiquitinated under the starvation condition and found on the vacuolar-limiting membrane. Then the question is why Tat2p is not always sorted to the MVB pathway. Mono- or polyubiquitination could explain the difference. Alternatively, the position of ubiquitination may be important. For example, ubiquitin signals near the membrane could be recognized by the MVB-sorting machinery more easily than the distal ones (Reggiori and Pel-
Lipid raft–dependent sorting of Tat2p
We present two lines of evidence indicating that association with the detergent-insoluble membrane domain is required for the plasma membrane delivery of Tat2p. First, Tat2p became detergent-insoluble under the low tryptophan condition. Second, depletion of sterols by using the mevalonate auxotroph Δerg13 mutant disrupted the detergent-insoluble membrane domain and simultaneously blocked the plasma membrane targeting of Tat2p. The detergent insolubility and sterol dependence of this membrane domain fit well with the concept of the lipid raft (Simons and Ikonen, 1997). Our results with Δerg13 indicate that Tat2p is missorted to the vacuole in the absence of lipid rafts. Similarly, the proton ATPase Pma1p is delivered to the plasma membrane in association with rafts, and missorted to the vacuole when rafts are disrupted (Bagnat et al., 2001). It appears that raft and nonraft domains are segregated for the plasma membrane and late endosomal delivery, respectively.

The raft association is not obligatory for the plasma membrane targeting of Tat2p if its vascular sorting is inhibited by Δspp12 or Δbul1. The fact that the severe tryptophan auxotroph of the raft-deficient Δerg13 was suppressed by Δbul1 indicates that raft association and polyubiquitination have counteracting effects in the sorting of Tat2p. On aberrant polyubiquitination, Tat2p is probably diverted from the raft-dependent plasma membrane targeting pathway to the nonraft pathway to late endosomes. Because ERG6 is involved in a late step of the ergosterol biosynthetic pathway, sterols are not depleted in Δerg6, but intermediates such as zymosterol accumulate (Munn et al., 1999). These intermediates are still capable of forming rafts, judging from the detergent insolubility of the GPI-anchored protein in Δerg6. However, Tat2p cannot be associated with such altered rafts any more and missorted to the vacuole. Tat2p may be just unable to reside stably in the rafts with the unusual sterol composition, or could be excluded from the rafts due to its inappropriate polyubiquitination.

Experiments with the sec mutants grown at low tryptophan indicated that Tat2p becomes associated with rafts in the Golgi. Tat2p associates with rafts even at high tryptophan in the sec14 mutant (unpublished data). Although the altered phospholipid composition of this mutant (McGee et al., 1994) might indirectly affect the raft organization, this observation suggests that Tat2p can gain access to rafts in the Golgi. On the other hand, polyubiquitination of Tat2p was not detected in the sec14 mutant, indicating that the polyubiquitination occurs after the exit from the Golgi. Thus, Tat2p would be first partitioned into rafts, and then be subjected to the ubiquitin-dependent sorting, presumably in early endosomes.

How polyubiquitin acts as a sorting signal to the nonraft, vacuolar trafficking pathway remains to be resolved. Lafont and Simons (2001) have shown that the ubiquitin ligases Cbl and Nedd4 are partitioned into rafts. Interestingly, the yeast Nedd4 homologue Rsp5p is partially resistant to detergent extraction (Wang et al., 2001), implying that polyubiquitination of Tat2p by the Rsp5p–Bul1p complex could occur in the rafts. Sorting receptors such as Hrs (Raiborg et al., 2002) may bind to polyubiquitin and divert cargo proteins to the nonraft membrane domains. Alternatively, Tat2p might dissociate from rafts independently of ubiquitin. The dissociation could change the environment around the molecule and would then trigger its polyubiquitination and sorting to late endosomes.

That the slight alteration in sterol structure or composition can dramatically change the destination of a plasma membrane protein raises a possibility that similar regulation could be used for differentiation of the cell surface, for example, during the development in higher eukaryotes. Our future work will aim at understanding how sterols might be involved in such higher order regulations and how they are linked to ubiquitin, a key player in the post-Golgi traffic.

While this manuscript was in preparation, Bagnat and Simons (2002) reported that Fus1p, a plasma membrane protein required for yeast mating, is largely excluded from rafts and mislocalized to the vacuole in the Δerg6 mutant. This behavior of Fus1p in Δerg6 is similar to that of Tat2p, and supports the view that a subset of plasma membrane proteins are missorted in Δerg6 to cause pleiotropic phenotypes. Indeed, the mating deficiency (Gaber et al., 1989) and the drug hypersensitivity (Kaur and Bachhawat, 1999) of Δerg6 might all be explained by the missorting of plasma membrane proteins due to impaired raft association and inappropriate ubiquitination.

Materials and methods
Yeast strains and media
Yeast strains used in this study are listed in Table I. Yeast cells were grown in MCD medium, composed of 0.67% yeast nitrogen base without amino acids (Difco Laboratories), 0.5% casamino acids (Difco Laboratories), and 2% glucose. Casamino acid is the mixture of amino acids lacking tryptophan. Adenine and uracil were supplemented at 20 μg/ml. Tryptophan was supplemented at a high (200 μg/ml), standard (20 μg/ml), or low (2 μg/ml) concentration. Unless otherwise indicated, yeast cells were grown at 30°C.

Plasmids and antibodies
Details of the various plasmid constructions and antibodies are described in the supplemental materials and methods section (available at http://www.jcb.org/cgi/content/full/jcb.200303088/DC1).

Fluorescence microscopy
Immunofluorescence microscopy was performed essentially as described before (Nishikawa and Nakano, 1991), except that permeabilization of fixed cells was performed by spheroplasting buffer containing 1% (vol/vol) BSA and 0.1% (vol/vol) Triton X-100 for 10 min at RT. Cells were observed and photographed using a photomicroscope (model BX-60; Olympus). Alternatively, the same microscope equipped with a confocal laser scanner unit (model CSU10; Yokogawa Electronic Corp.) was used. Images were acquired by a high resolution digital charge-coupled device camera (model C4742–95; Hamamatsu Photonics) and processed by IPLab software (Scanalytics).

Detection of the ubiquitinated forms of Tat2–3HAp
Detection of ubiquitinated Tat2–3HAp was performed basically according to the method used for the case of Gap1p (Helliwell et al., 2001). To enhance the detection, the myc-tagged ubiquitin was exogenously expressed. The expression of myc-Ub was under the control of the CUP1 promoter, which was inducible by addition of CuSO4 to the medium (Ellison and Hochstrasser, 1991). However, myc-Ub conjugates were detectable even when the promoter was uninduced, as was reported previously (Hoch-
In this work, cells were grown at the basal expression level of the CLU1 promoter. 5 × 10⁵ cells were collected and treated with NaNO₃ and potassium fluoride at the final concentration of 20 mM each. The cells were resuspended in 125 μl lysis buffer (20 mM Tris-HCl, pH 7.4, 1 mM EDTA, 1.6% SDS, 6 M urea, 5 mM Nethylmaleimide, and 0.02% NaNO₃) containing a protease inhibitor mixture (1 mM PMSF, 5 μg/ml chymostatin, leupeptin, antipain, and pepstatin A), lysed by agitation with glass beads, and incubated at 37°C for 20 min. 875 μl IP dilution buffer (1.1% Triton X-100, 170 mM NaCl, 6 mM EDTA, 60 mM Tris-HCl, pH 7.4, 5 mM N-ethylmaleimide, and 0.02% NaN₃), the protease inhibitor mixture) was added to the cell lysates, and insoluble material was removed by centrifugation. 750 μl supernatant was mixed with 40 μg protein G Sepharose 4 Fast Flow (Amersham Biosciences), and preclarified by rotation at RT for 30 min. The samples were centrifuged, and 700 μl supernatant was mixed with 770 μl IP dilution buffer (1 mM Tris-HCl, pH 7.4) and once with high salt wash buffer (1% Triton X-100, 0.2% SDS, 50 mM NaCl, 5 mM EDTA, and 50 mM Tris-HCl, pH 7.4). The beads were washed with SDS-PAGE sample buffer (2% SDS, 5% β-mercaptoethanol, 10% glycerol, 50 mM Tris-HCl, pH 6.8, and 0.025% bromophenol blue) containing 6 M urea, and incubated at 37°C for 20 min. 24 μl of the sample was subjected to SDS-PAGE and immunoblotting with the anti-myc antibody (9E10) to detect myc-Ub conjugates. To detect Tat2-3HA, the sample was diluted 15-fold and 20 μl was loaded. Anti-HA antibody (16812) was used for immunoblotting.

### Analysis of lipid rafts

5 × 10⁵ cells were collected, treated with NaNO₃, and potassium fluoride at a final concentration of 20 mM each, and resuspended in 275 μl TNE buffer (25 mM Tris-HCl, pH 7.5, 150 mM NaCl, and 5 mM EDTA) containing a protease inhibitor mixture (1 mM PMSF and 5 μg/ml chymostatin, leupeptin, antipain, and pepstatin A). After adding glass beads, the suspension was vortexed for 30 s and was then chilled on ice for 30 s, repeating six times. Unbroken cells and debris were removed by centrifugation at 500 rpm for 5 min. The cleared lysate (175 μl) was mixed with equal volume of TNE containing 40 mM CHAPS (Sigma-Aldrich), and was then incubated at 4°C for 30 min. The tube was centrifuged at 5,000 rpm for 5 min, and 330 μl supernatant was mixed with 770 μl 50% OptiPrep (Nycomed Pharma/TNE) 200 mM CHAPS to give the final concentration of 35% OptiPrep. The solution was set on the bottom of a 3 PC tube (Hitachi Koki Co., Ltd.), and overlaid with 1.4 ml 30% OptiPrep/TNE 200 mM CHAPS and 0.5 ml TNE 200 mM CHAPS. The gradients were centrifuged at 4°C for 7.5 h using a rotor (model RPS65T; Hitachi Koki Co., Ltd.) at 35,000 rpm.
and nine fractions (320 μl each) were collected from the top. Each fraction was mixed with 288 μl 110 mM Tris-HCl, pH 6.8/4.4% SDS/22% glycerol and 32 μl β-mercaptoethanol, incubated at 37°C for 5 min, and subjected to SDS-PAGE.

Online supplemental materials
Plasmid construction, antibodies, and immunoblotting procedures are included in the online supplemental materials, available at http://www.jcb.org/cgi/content/full/jcb.200303088/DC1.

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Plasmids and antibodies
The 2.9-kb PstI-EcoRI fragment of TAT2 was prepared from YCpTAT2 (Schmidt et al., 1994; Nakamura et al., 2000), and was subcloned into YEpUC195 (Gietz and Sugino, 1988) to generate pKU68 (YEpTAT2).

Three tandem copies of the HA epitope were placed just before the stop codon of TAT2 as follows: the 2.9-kb PstI-EcoRI fragment of TAT2 was subclone into pUC118 to generate pKU1. Using pKU1 as a template, the XbaI site (underlined in the sequences below) was introduced by PCR just before the stop codon of TAT2. The upstream or downstream region of the stop codon was amplified using the primer combination 5′-GGCCACGTGATTGTGCAG-3′ and 5′-GTATTA TCTAGAACACAGAGCTGCTG-3′, or 5′-TTGGATCTAATACCAGAGAAGC-3′ and 5′-GTTATGCAATGATCT-3′, respectively. After annealing these two PCR products, the entire 0.57-kb fragment was amplified, cut with PmaCI and EcoT221, and then sequenced for confirmation. This fragment was used to replace the corresponding region of pKU1 to generate pKU42. The Nhel-NheI cassette containing three copies of the HA epitope was amplified from pYT11 (Takita et al., 1995), and inserted into the XbaI site of pKU42. The 3.0-kb PstI-EcoRI fragment of TAT2–3HA from pKU42 was subcloned into YCpplc33 to generate pKU46 (YCpTAT2–3HA).

GFP was fused to the COOH-terminus of Tat2 as follows: the ORF of GFP lacking the stop codon was amplified by PCR using mutated PCR primers. Primers used were 5′-AAACTAGTATGTTAGAGCAGGAGGCCC-3′ and 5′-AAAATCGTCTTGCTGCTG-3′, both of which contained the SpeI sites (underlined). The 0.7-kb SpeI-SpeI fragment of GFP was sequenced, and was inserted into the XbaI site of pKU42. TAT2-GFP was prepared as the 3.6-kb PstI-EcoRI fragment and subcloned into YCpplc33 to generate pKU76 (YCpTAT2–GFP).

Plasmids for disruption of TAT2 were based on pKU1. The 1.5-kb EcoT14I-EcoR1 fragment of TAT2 was replaced by ADE2 or hisG-URA3-hisG. ADE2 was prepared as the 2.2-kb BglII-BglII fragment from pAS11 (Stotz and Linder, 1990). HisG-URA3-hisG was prepared as the 3.8-kb BamHI-BglII fragment from pNKY51 (Alani et al., 1987). Either pKU41 (tat2Δ::ADE2) or pKU82 (tat2Δ::hisG-URA3-hisG) was digested with EcoRI and SplI, and then used for transformation. Yeast transformation was performed by the lithium thiocyanate method (Keszenen-Pereyra and Hieda, 1988). Correct integrants (tat2Δ::hisG-URA3-hisG) were grown on 5-fluoroorotic acid (5-FOA) medium to select ura3 auxotrophs (tat2Δ::hisG).

For integration of TAT2–3HA into the genome, the 1.2-kb StuI-EcoRI fragment of TAT2–3HA, which corresponds to the COOH-terminal and 3′ noncoding regions, was prepared from pKU46. This fragment was inserted between the Smal and EcoR1 sites of YEpplc21 (Gietz and Sugino, 1988). The resultant URA3-marked plasmid pKU51 was cut with PmaCI and used for transformation. The integrants were grown on 5-FOA medium to replace the TAT2 locus with TAT2–3HA. The integration and the replacement was confirmed by PCR and immunoblotting to detect Tat2–3HAp.

Three lysine residues (10, 17, and 20) of Tat2p were replaced with arginine by using mutated PCR primers. The upstream region was amplified using the primer combination 5′-TCGTTAATGGGTAGTAAGC-3′ and 5′-GTATGA TCTCCGCCTGTTGAAAGCGGCA-3′. The downstream region was amplified by 5′-TCTGTCAAGGGGTCAGTGGCAAGGGAAGGGAAGATCTCAGCTG-3′ and 5′-GTTAGATCTCCTCCTCCTACCTGTTGGAACAG-3′. Sequences where lysine codons are converted to arginine are underlined. After annealing these two PCR products, the entire 0.7-kb fragment was amplified and cut with SnaBI and BamHI. The SnaBI-BamHI fragment was used to replace the corresponding region of pKU1, and the resultant plasmid (pUC-TAT23K) was sequenced for confirmation. Ligation was performed among three fragments; the PstI-PvuI fragment of pKU1, and the resultant plasmid (pUC-TAT23K) was isolated from pSPRA12, and subcloned into pUC18. The 0.49-kb EcoT14I-EcoT14I fragment of PEP4 was replaced by the 1.8-kb BamHI-BamHI fragment of HIS3. Plasmid pKU60 (pep4Δ::His3) was digested with EcoRI and Sall, and then introduced into yeast cells.

VPS1 was amplified from the genome of YPH499. Primers used were 5′-AAAGCAGGACGATCATCTTCAAGACGCC-3′ and 5′-AAGCCCGGTCACGCGCAGA-3′. The SplI sites are introduced in the primers (underlined). The PCR product was cut with SplI, and the 3.1-kb fragment was subcloned into pUC19. The internal 1.8-kb BstPI-SpeI fragment of VPS1 was replaced by the 1.8-kb BamHI-BamHI fragment of HIS3. Plasmid pKU62 (vps1Δ::His3) was digested with SplI, and then introduced into yeast cells.
To disrupt \textit{BUL1}, the 5.6-kb EcoRI-Sacl fragment of \textit{BUL1} was prepared from pHY06 (Yashiroda et al., 1996), and subcloned into pBlueScript\textsuperscript{R} II KS+. The internal 1.8-kb BstI-SpeI region was replaced by the 1.8-kb BamHI-BamHI fragment of \textit{HIS3} or the 2.3-kb XhoI-SalI fragment of \textit{LEU2} to form pKU143 or pKU121, respectively. Before transformation, pKU143 was cut with EcoRI and SalI, and pKU121 with Apal and SalI.

YEp96 and YEp105 (Ellison and Hochstrasser, 1991), \textit{TRP1}-marked plasmids with which ubiquitin and myc-tagged ubiquitin, respectively, are expressed by the copper-inducible \textit{CUP1} promoter and were provided by Mark Hochstrasser (Yale University, New Haven, CT). To convert the plasmid marker to \textit{URA3}, the 1.0-kb BamHI-ClaI fragment of \textit{P\textsubscript{CUP1}-Ub-T\textsubscript{Cyc1}} was prepared from YEp96 and subcloned into pRS426 (Sikorski and Hieter, 1989) to form pKU105. Similarly, pKU106 was constructed by preparing the \textit{P\textsubscript{CUP1}-myc-Ub-T\textsubscript{Cyc1}} fragment from YEp105.

Disruption of \textit{DOA4} was performed as follows: the upstream region of the \textit{DOA4} ORF was amplified by PCR using the primer combination 5'-\texttt{AAAAAGCTTCTATTCCCTCCTCCGCCTTTGGTCC-3'} and 5'-\texttt{AAAAAGATCCGTTATTTCTTGTTGGAAGGTGGCA-3'}, and then cut with Sacl and BamHI (each restriction site is underlined). The downstream region was amplified using 5'-\texttt{AAAAAGATCTTTTTGAGTGCATTCCTTGGTTGGAAGGTGGCA-3'} and 5'-\texttt{AAAAAGCTTCTATTCCCTCCTCCGCCTTTGGTCC-3'}, and then cut with BamHI and Sphi (each restriction site is underlined). Ligation was performed among these two fragments and pUC18 to generate a 0.8-kb Sacl-Sphi insert in pUC18. This plasmid was cut at the BamHI site, the junction between the two fragments, and the 1.8-kb BamHI-BamHI fragment of \textit{HIS3} was inserted. Plasmid pKU108 (\textit{doa4\Delta::HIS3}) was cut with Sacl and Sphi, and then was introduced into yeast cells.

\textit{VPS27} was amplified from the genome of YPH499. Primers used were 5'-\texttt{AAAAAGATGCTTTACCTGTGGATGTTT-3'} and 5'-\texttt{AAAAAGCTTCTATTCCCTCCTCCGCCTTTGGTCC-3'}. The PCR product was cut with BamHI and EcoRI, and the 3.2-kb fragment was subcloned into pUC18. The internal 1.4-kb BglII-ClaI fragment of \textit{VPS27} was replaced by the 1.8-kb BamHI-BamHI fragment of \textit{HIS3}. Plasmid pKU65 (\textit{vps27\Delta::HIS3}) was digested with EcoRI and Sall, and then introduced into yeast cells.

Plasmids for expression of GFP-Pep12p were constructed as follows. The \textit{PEP12} ORF was amplified by PCR using the primer combination 5'-\texttt{AAAAAGATGCTTTACCTGTGGATGTTT-3'} and 5'-\texttt{AAAAAGCTTCTATTCCCTCCTCCGCCTTTGGTCC-3'}. As indicated by the underlines, The BamHI or the HindIII site was introduced adjacent to the start or stop codon, respectively. The BamHI-HindIII fragment of \textit{PEP12} was subcloned into pSKY5, at the junction between \textit{EGFP} and the \textit{CMKI} terminator, to form pKU84. The plasmid pSKY5 is based on pRS316 (Sikorski and Hieter, 1989), and is designed for expression of EGFP fusion proteins under the control of the \textit{TDH3} promoter. As another plasmid marker (\textit{TRP1}), the 3.3-kb BamHI-SalI fragment of \textit{P\textsubscript{TDH3-EGFP-PeP12-CMKI}} was subcloned into pRS314 to form pKU144.

Disruption of \textit{ERG13} was performed as follows: the upstream region of the \textit{ERG13} ORF was amplified by PCR using the primer combination 5'-\texttt{AAAAAGATGCTTTACCTGTGGATGTTT-3'} and 5'-\texttt{AAAAAGCTTCTATTCCCTCCTCCGCCTTTGGTCC-3'}. As indicated by the underlines, The BamHI or the HindIII site was introduced adjacent to the start or stop codon, respectively. The BamHI-HindIII fragment of \textit{PEP12} was subcloned into pSKY5, at the junction between \textit{EGFP} and the \textit{CMKI} terminator, to form pKU84. The plasmid pSKY5 is based on pRS316 (Sikorski and Hieter, 1989), and is designed for expression of EGFP fusion proteins under the control of the \textit{TDH3} promoter. As another plasmid marker (\textit{TRP1}), the 3.3-kb BamHI-SalI fragment of \textit{P\textsubscript{TDH3-EGFP-PeP12-CMKI}} was subcloned into pRS314 to form pKU144.

The anti-HA (16B12) and the anti-myc (9E10) mAbs were purchased from BAbCO. The rabbit affinity-purified pAb against the HA epitope (Y-11) was purchased from Santa Cruz Biotechnology, Inc. To eliminate nonspecific staining in immunofluorescence microscopy, these antibodies were adsorbed with fixed and Triton X-100–permeabilized yeast cells not expressing the HA epitope (Roberts et al., 1991). Anti-Pep12p mAb (2C3-G4), anti-vacuolar alkaline phosphatase mAb (1D3-A10), goat anti–mouse Alexa Fluor\textsuperscript{R} 488, and goat anti–rabbit Alexa Fluor\textsuperscript{R} 568 were purchased from Molecular Probes, Inc. Anti-Gas1p antibodies were provided by R. Hirata (RIKEN, Saitama, Japan). In immunofluorescence microscopy, the preadsorbed anti-HA mAb was used at 1:300, and anti–mouse Alexa\textsuperscript{R} 488 at 3 \textmu g/ml. For double staining, the anti-Pep12p mAb and the preadsorbed anti-HA pAb were used at 10 \textmu g/ml and 1:500, respectively.

To prepare antibodies specific for Tat2p, a synthetic peptide that contains the 14 amino acids spanning from 248 to 261 residues (PDHEFIGAKYWHDP) of Tat2p was used as the antigen. Glycine and cysteine were added to the COOH terminus of this peptide. The peptide was coupled to KLH using the cross-linker \textit{N}-maleimidobenzoyl-\textit{N}-hydroxysuccinimide ester, and linked conjugates were dialyzed against PBS. The peptide–KLH conjugate (0.2 mg) was fixed with Freund’s complete adjuvant and injected into rabbits, followed by biweekly boosts with 0.2 mg of conjugate in Freund’s incomplete adjuvant. The antiserum against Tat2p was affinity purified, adsorbed to \textit{Δtat2} cells, and used for immunofluorescence microscopy at 1:2.5 dilution.

**Immunoblotting**

Cells were treated with NaN\textsubscript{3} and KF at the final concentration of 20 mM each. Preparation of protein extracts was performed by agitation with glass beads in 50 mM Tris-HCl (pH 6.8), 2% SDS, 10% glycerol, 1 mM EDTA, and 1 mM PMSF. Before SDS-PAGE, \textit{β}-mercaptoethanol was added at the final concentration of 5%, and the samples were heated at 37°C for 10 min. The anti-HA mAb was used at 1:1,000.
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