Role of Three Rab5-like GTPases, Ypt51p, Ypt52p, and Ypt53p, in the Endocytic and Vacuolar Protein Sorting Pathways of Yeast

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Abstract. The small GTPase rab5 has been shown to represent a key regulator in the endocytic pathway of mammalian cells. Using a PCR approach to identify rab5 homologs in Saccharomyces cerevisiae, two genes encoding proteins with 54 and 52% identity to rab5, YPT51 and YPT53 have been identified. Sequencing of the yeast chromosome XI has revealed a third rab5-like gene, YPT52, whose protein product exhibits a similar identity to rab5 and the other two YPT gene products. In addition to the high degree of identity/homology shared between rab5 and Ypt51p, Ypt52p, and Ypt53p, evidence for functional homology between the mammalian and yeast proteins is provided by phenotypic characterization of single, double, and triple deletion mutants. Endocytic delivery to the vacuole of two markers, lucifer yellow CH (LY) and α-factor, was inhibited in Δypt51 mutants and aggravated in the double ypt51ypt52 and triple ypt51ypt52ypt53 mutants, suggesting a requirement for these small GTPases in endocytic membrane traffic. In addition to these defects, the here described ypt mutants displayed a number of other phenotypes reminiscent of some vacuolar protein sorting (vps) mutants, including a differential delay in growth and vacuolar protein maturation, partial missorting of a soluble vacuolar hydrolase, and alterations in vacuole acidification and morphology. In fact, vps21 represents a mutant allele of YPT51 (Emr, S., personal communication). Altogether, these data suggest that Ypt51p, Ypt52p, and Ypt53p are required for transport in the endocytic pathway and for correct sorting of vacuolar hydrolases suggesting a possible intersection of the endocytic with the vacuolar sorting pathway.

IN eukaryotic cells endocytosis is a basic process by which extracellular molecules are internalized by invaginating and pinching off parts of the plasma membrane. This membrane transport pathway has been extensively studied also in the budding yeast Saccharomyces cerevisiae using two endocytic markers, Lucifer yellow CH (LY) to follow fluid-phase endocytosis and α-factor to follow receptor-mediated endocytosis (for review see Riezman, 1993). The availability of a large pool of mutants and the ease of isolation of novel ones has made the budding yeast S. cerevisiae an excellent system to study the in vivo functions of proteins in the process of endocytosis and to identify novel components of the endocytic machinery. For example, the requirement for clathrin heavy chain (Payne et al., 1988), actin and the actin-binding protein Sac6p (Kübler and Riezman, 1993) in the step of endocytic internalization has been recently established by such analysis. Genetic screens for mutants defective in the internalization of endocytic markers and their delivery to the vacuole have furthermore resulted in the identification of End3p and End4p (Raths et al., 1993) and Renlp/Vps2p (Davis et al., 1993), whose precise functions in endocytic membrane traffic are however yet unknown.

Another class of proteins that is believed to be required for endocytic transport is the family of small GTPases, known as Ypt family in yeast and rab family in mammalian cells. Their key regulatory role in intracellular membrane traffic was initially demonstrated with the help of conditional lethal yeast mutants, sec4 and ypt1, in the secretary pathway (Salminen and Novick, 1987; Segev et al., 1988; Schmitt et al., 1988). In mammalian cells a large number of proteins belonging to the rab family have been identified due to their similarity in primary structure to Ypt1p and Sec4p (Haugbruck et al., 1987; Touchot et al., 1987; Zahraoui et al., 1990).
Materials and Methods

Members of the rab family that are specifically associated with elements of the endocytic apparatus are rab4, rab5, rab7, and rab9. Whereas rab4 and rab5 are located on early endosomes (Chavrier et al., 1990; van der Sluijs et al., 1991) and have a regulatory function in early steps of endocytosis (Bucci et al., 1992; van der Sluijs et al., 1992), rab7 and rab9 are associated with late endosomes and therefore are involved in later steps of the pathway (Chavrier et al., 1990; Lombardi et al., 1993). In S. cerevisiae a homolog of rab7, Ypt7p, has recently been identified (Wichmann et al., 1992). The ypt7 null mutant is characterized by a delay of endocytic delivery of α-factor to the vacuole and by defects of vacuolar protein sorting and maturation (Wichmann et al., 1992).

In contrast to the proposed involvement of rab7 and Ypt7p in later steps of endocytosis, studies on rab5 have suggested a key regulatory role of this small GTPase in early endocytic traffic. Rab5 was found to be required for the fusion of early endosomes in vitro (Gorvel et al., 1991). In vivo, overexpression of the wild-type protein in BHK cells led to increased internalization of endocytic markers and to the appearance of very large early endosomal structures (Bucci et al., 1992). Conversely, expression of a mutant form of rab5, rab5le133, inhibited endocytosis and resulted in fragmentation of early endosomes.

In the fission yeast Schizosaccharomyces pombe a homolog of rab5, ypt5p, has been identified recently (Armstrong et al., 1993). However, no experimental data are yet available that suggest any function for this protein in endocytosis in the fission yeast (Armstrong et al., 1993).

Here we report the isolation of three genes from the budding yeast S. cerevisiae that encode Ypt proteins with extensive homology to mammalian rab5 and the S. pombe ypt5p. The defects associated with single, double, and triple null mutants of YPT51, YPT52, and YPT53 suggest an important function of the encoded proteins in the delivery to the vacuole of two endocytic markers, α-factor and LY. Furthermore, due to a number of vacuole-associated defects found for the here described ypt mutants, such as protein sorting, acidification and morphological changes, a requirement of these Ypt proteins for vacuole biogenesis can be envisaged.

Materials and Methods

Plasmids, Strains, and Growth Conditions

The yeast genomic library, kindly provided by M. Hall (Heitman et al., 1991), contains ~5-10 kb fragments resulting from partial digestion of S. cerevisiae genomic DNA with Sau3A (strain MH145-4B [A(mat)130-141::CEN1-A, mpf2-loc2, rme, ura3, leu2, his4, met, ade5, HMLa]). The fragments were inserted into the BamHI site of the polylinker of plasmid pSEYS, a high-copy number 2/~m URA3 vector (Emr et al., 1986).

Table I Strains Used

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The Escherichia coli strains used in these experiments were M11 (araD139, lacX74, galE, galK, his, rpsL) (provided by M. Hall, Bio-center, Basel, Switzerland), SE10 (Aehstetter et al., 1988), and XL1-Blue (Stratagene, La Jolla, CA), which were grown in LB medium (1% NaCl, 1% peptone, 0.5% yeast extract). LB plates contained additionally 2% agar. Strains carrying plasmids were grown in the presence of 100 μg/ml ampicillin.

The strains of S. cerevisiae used in these experiments are listed in Table 1. Unless otherwise indicated yeast strains were grown in complete medium (1% yeast extract, 2% peptone, and 2% glucose) (YPD) to early logarithmic phase (0.5-2 × 107 cells/ml) at 24°C on a rotary shaker. Synthetic growth medium (SD medium) has been described elsewhere (Dulic et al., 1991).

Reagents

35S-labeled α-factor was prepared from biosynthetically labeled yeast cells overproducing the pheromone (Dulic et al., 1991). LY-CH was purchased from Fluka (Buchs, Switzerland), quinacrine was from Sigma Chemical Co. (St. Louis, MO), and carboxydiethylfluorescein diacetate (CDFDA) was from Molecular Probes Inc. (Eugene, OR). Yeast tRNA, herring sperm DNA, dNTPs and dNTPs were obtained from Boehringer Mannheim (Mannheim, Germany). Sequenase Version 2.0 was from United States Bio-chem. corp. (Cleveland, OH). 35S-H2SO4 (carrier free), [35S]methionine, o25S-dATP, and o25P-dCTP were purchased from Amersham Intl. (Buckinghamshire, England). Klenow-enzyme was from Boehringer Mannheim. Protein A-Sepharose CL-4B and Sepharose CL-4B were from Pharmacia (Uppsala, Sweden). Chemicals for gel electrophoresis were obtained from Bio Rad Laboratories (Richmond, CA) and National Diagnostics (Atlanta, GA). EXTENSIFY used for fluorography was from Dupont–NEF (Boson, MA).

Table I Strains Used

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PCR Amplification of Ypt51 and Ypt53 Fragments and Isolation and Sequencing of Full Length Clones

Degenerate primers representing the protein sequence motif GXXXGXS/T (Chavrier et al., 1992) and LARPYR (GTG/T/G/A/C-TAC- (G/T)AC-G(T/T)ATACGAGTGAATATAT) found in nab5A, rab5B, rab5C (Chavrier et al., 1992; Simons and Zerial, 1993) were used for amplification of yeast genomic DNA prepared from the strain RHA144. The initial PCR reaction was carried out in a volume of 20 μl and consisted of 30 cycles of 1 min each at 94°C, 2 min at 60°C, and 1 min at 72°C, with a final 5 min extension at 72°C. The reaction volume was increased to 100 μl and a second round of amplification was performed using the same conditions as described above. The reaction product of ~200 bp was gel purified and labeled with α-32P-dCTP (3,000 Ci/mmol) using the random primer labeling kit (Boehringer Mannheim). To isolate full length clones of the respective genes, ~40,000 E. coli colonies (transformed with the genomic library) were screened with the α-32P-labeled PCR reaction product. Plating of bacteria onto nitrocellulose filters on LB-Ampicillin plates and preparation of replica-filters were done by standard procedures (Sambrook et al., 1989). After lysing the cells with base and baking the filters for 2 h at 80°C, they were extensively washed with 3× SSC, 0.1% SDS at 68°C to remove bacterial debris. Prehybridization was performed for 2 h at 68°C in 6× SSC, 5× Denhard’s solution, 100 μg/ml herring sperm DNA and 0.5% SDS. Hybridization was done for 12 h at 68°C in 6× SSC, 1× Denhard’s solution, 100 μg/ml yeast tRNA and 32P-labeled PCR reaction product, at approximately 3 × 106 cpm/ml. Filters were washed with 2× SSC, 0.1% SDS at 68°C four times for 15 min and exposed against x-ray films (XAR5; Eastman Kodak Co., Rochester, NY). All positive clones identified in the initial screening were isolated and streaked out to obtain single colonies. 5–10 of these colonies were subjected to a second round of screening (as described above). Positive clones were restranded and single colonies were subjected to a third round of rescreening as described above. Plasmids were isolated from the positive clones using Qiagen columns (Diagen, Düsseldorf, Germany) as described by the manufacturer.

To determine the nucleotide sequences of the respective clones, a degenerate oligonucleotide, representing a protein sequence motif that can be found in most Ypt/rab proteins, was used as a primer in a primer deoxy sequencing of double-stranded plasmid DNA. Sequencing was done according to the method of Winship (1989) using Sequenase Version 2. This revealed that out of the 15 clones isolated, nine represented YPT51 (four different subclones), three represented YPT53 (one subclone), and three represented so far unidentified clones (different subclones). The coding sequence of YPT51 and YPT53 were determined from both strands by using nested primers.

Disruption of YPT51, YPT52, and YPT53

Large internal parts of the genes YPT51, YPT52, and YPT53 were deleted from the genome and replaced with the LYS2, the URA3, and the LEU2 genes, respectively. Using the polymerase chain reaction (PCR), the genomic flanking regions on either side of YPT51 were amplified from pSEY8-B56 to generate a 205-nucleotide fragment corresponding to the 5′ flanking region (5′FR) (nucleotides 385–589) and a 267-nucleotide fragment corresponding to the 3′ flanking region (3′FR) (nucleotides 1083 to 1349). The “inner” oligonucleotides, i.e., those immediately flanking the YPT51 gene, were designed to create restriction sites for HindIII. The “outer” oligonucleotides contained restriction sites for ClaI (5′FR) and EcoRI (3′FR). The two PCR products were purified from agarose gel and used as a template in a second round of PCR, which used as primers the two outer oligonucleotides used in the first PCR reactions. The major product of this reaction was digested with ClaI and EcoRI and subcloned into pBSK(−). The resulting plasmid was opened with HindIII and a 4.7-kb LYS2 fragment was ligated between the 5′FR and the 3′FR. Plasmid pBS-YPT51LYS2, containing the predicted insert, was linearized with ClaI, cloned into the polylinker of pBSK(−), and used to transform the diploid strain RH1201 in a one step gene replacement (Orr-Weaver et al., 1981). After selecting for growth on plates in the absence of lysine, colonies were sporulated at 24°C. Tetrad dissection was then performed and the cells derived from different spores were grown up at 24°C and examined as described below. The disruption of YPT53 was performed by the same strategy as described for YPT51. The genomic flanking regions on either side of YPT53 were amplified from pSEY8-HSV1 to generate a 196 nucleotide 5′FR (nucleotides 418 to 613) and a 206 nucleotide spanning 3′FR (nucleotide 1150 to 1355). The inner oligonucleotides were designed to create restriction sites for XhoI, the outer oligonucleotides contained restriction sites for BamHI (5′FR) and PstI (3′FR). After two rounds of PCR (see above) the reaction product was subcloned into pGEMI (Promega Corp., Madison, WI), opened with XhoI and a 2.2-kb XhoI/SalI fragment containing the LEU2 marker was introduced. The resulting plasmid (pGEM-YPT53SEU2) was linearized with BamHI and PstI. Transformation and disruption of LEU2 transformants was carried out as described above.

To create a loss-of-function allele of YPT52, a 2.4-kb YPT52-containing genomic fragment resulting from exonuclease/Xbal digestion was subcloned into pBSK(−) to generate pBS-YPT52 and a 799-nucleotide EcoRV/EcoRV fragment containing the entire coding sequence of the gene was replaced by a 1.2-kb HindIII/HindIII fragment containing the URA3 gene. The resulting plasmid (pBS-YPT52URA3) was linearized with KpnI (contained in pBSK) and XbaI and used for subsequent transformation of diploids.

The null alleles of YPT51 and YPT53 contained a deletion of codon 31 to 195 (inclusive) and codon 31 to 209 (inclusive), respectively. The null allele of YPT52 did not contain any coding sequence information.

To determine integration of the respective constructs by site-specific recombination, DNA was prepared from cells derived from the four spores of complete tetrads was prepared (Philippsen et al., 1991). PCR was performed with a primer of the respective oligonucleotide that binds 5′ from the sequence that was used for homologous recombination of the three respective genes and a primer that binds to sequences within the 5′ region of the respective marker genes. With the mutants, amplification of a correct construct was observed by gel electrophoresis. Southern hybridization was performed with wild-type and mutant strains to confirm proper integration of the respective marker genes. For this, chromosomal DNA of the various strains was digested with EcoRI, separated on 0.8% (to analyze disruption of YPT51 and YPT35) or 0.6% (to analyze disruption of YPT52) agarose gels and transferred to GeneScreen Plus membranes (Dupont-NEN) as suggested by the manufacturer. Probes for YPT51 and YPT53 were prepared by PCR, using the respective oligonucleotides described before, gel purified, and labeled with digoxigenin as suggested by the manufacturer (Boehringer Mannheim). To analyze disruption of YPT52 two probes were prepared: a digoxigenin-labeled EcoRV/EcoRV fragment containing the coding region of YPT52, and a KpnI/BglII fragment containing the 5′ non-coding region of YPT2. Prehybridization and hybridization of probes with the filter and detection using anti-digoxigenin Fab labeled with alkaline phosphatase was performed according to the protocol provided by Boehringer Mannheim. Correct integration and disruption of the respective YPT genes were indicated by the following results: replacement of a ~6.8- and 3.1-kb fragment in a wild-type strain; appearance of a ~6.8- and 3.1-kb fragment in a ypt51 mutant versus an ~8.6-kb fragment in the wild-type strain; appearance of a ~8.2- and 3.1-kb fragment in a ypt52 mutant versus a ~7.8-kb fragment in the wild-type strain using the KpnI/BglII probe, or disappearance of the 7.8-kb band in the ypt52 mutant using the EcoRV/EcoRV YPT52 probe.

α-Factor Internalization and Degradation Assays

Pheromone internalization and degradation assays were carried out as described by Dulic et al. (1991), using biosynthetically labeled 3H-α-factor. Both assays were performed with the various strains grown overnight at 24°C. Binding of the pheromone was done for 1 h on ice, internalization at 30°C. The disappearance of intact, internalized α-factor (pH 1.2 sample) was quantified by densitometric scanning as described (Singer and Riezmann, 1990).

LFCH Accumulation

LY internalization experiments were essentially performed as described by Dulic et al. (1991). The various strains were grown to early logarithmic stage overnight at 24°C and LY accumulation (using 16 mg/ml final concentration) was analyzed after internalization at 30°C for 1 h. The cells were observed using a Zeiss Axioskop microscope equipped with fluorescence and Nomarski optics. Photographics were made using Kodak T-Max 400 films (Eastman Kodak Co.) exposed to ASA 1,600. Exposures for fluorescence were 30 s. To quantify the results obtained with ypt52pse53 mutants as compared with wild type, cells with a clear vacuolar staining positive for LY were counted as positive, cells without vacuolar staining, but with recognizable vacuole by Nomarski were counted as negative. 150 cells were counted.
Staining of Vacuoles with Quinacrine and CDCFDA

The different strains used for the analysis were grown overnight at 24°C to early logarithmic phase. For staining with quinacrine, 5 × 10⁷ cells were collected and resuspended in 500 μl YPD, 50 mM Na₂HPO₄, pH 7.6. Quinacrine was added to 0.2 mM final concentration and staining was performed at RT for 5 min. The cells were washed once in 2% glucose, 50 mM Na₂HPO₄, pH 7.6, and immediately viewed on concanavalin A-coated coverslips as described by Roberts et al. (1991), using fluorescence and Nomarski optics. Photographs were made using Kodak T-Max 200 film and exposures for fluorescence were done for 30 s.

For staining with carbodiimide-fluorescein diacetate (CDCFDA), 10⁷ cells were resuspended in 100 μl SD medium, 50 mM citric acid, pH 5, 20 mM CDCFDA. After 20 min at 30°C the cells were immediately viewed using either the Axiopt microscope or, alternatively, the EMBL confocal microscope. Excitation wavelength 476 nm. Photographs were made using Kodak T-Max 400 films exposed to ASA 1600 (Axiopt) and Kodak T-Max 100 films (confocal microscope). Exposures for fluorescence were 30 s except for analysis of ytp53pΔ52 and ytp53pΔ52pΔ53 mutants for which the exposure was reduced to 5 s.

Thin Section Electron Microscopy

Electron microscopy of cells that had been grown at 30°C to early logarithmic phase was performed as described previously by Ossig et al. (1991).

Carboxypeptidase Y and Alkaline Phosphatase Biogenesis

Cells of the various strains used in the analysis were grown overnight at 24°C in YPD medium to ~1 × 10⁶ cells/ml. Cells (5 × 10⁷ per time point) were harvested, washed twice with H₂O and resuspended to 2.5 × 10⁶ cells/ml in SD medium containing 2% glucose, 100 μM ammonium sulfate and nutrient supplements (SD low sulfate). After incubation at 24°C for 5 h the cells were collected, washed once with H₂O and resuspended to 2.5 × 10⁶ cells/ml in minimal medium as described above, but without sulfate (SD no sulfate) and with 0.5 mg/ml bovine serum albumin.

To label whole cells, a pulse was initiated at 30°C by the addition of [³⁵S]methionine (40 μCi/time point, 1 μCi/66 μl) and metabolic labeling conducted for 5 min at 30°C. 1 ml of cells was removed (and processed as described below), ammonium sulfate was added to 2 mM, and methionine and cysteine to 30 μg/ml, each, to initiate the period of chase. 1-ml samples were taken after 2.5, 5, 10, 20, 40, and 60 min. To terminate the reaction, the 1-ml samples were added to NaN₃ and NaBP (final concentration 5 mM each) and cysteine to 30 μg/ml, each, to initiate the period of chase. 1-ml samples were added to NaN₃ and NaBP (final concentration 5 mM each) and cysteine to 30 μg/ml, each, to initiate the period of chase.

The precipitates were collected by centrifugation, washed twice with TNET, once with 10 mM 1HCl, pH 6.8, 10 mM NaCl. The supernatant was diluted with 1 ml TNET was added and considered as 100% total.

To determine extracellular secretion of vacuolar enzymes, cells (2 × 10⁶ cells/ml) were treated in 0.1 M Tris/HCl, pH 6.8, 1.2 M sorbitol, 1 mg/ml each of BSA and ovalbumin and 10 μl lysylase. After incubation at 30°C for 45 min, the supernatants were collected by centrifugation for 5 min at 1,500 rpm, resuspended in the same medium except for lysylase, and pulse labeling was initiated by the addition of 6 μl [³⁵S]methionine (1 μCi/66 μl). After 15 min at 30°C, a chase period of 30 min followed as described above. The reaction was stopped by the addition of NaN₃ and NaBP to 5 mM final concentration, each, and by chilling the spheroplasts to 4°C. The spheroplasts were collected by centrifugation for 10 min at 1,500 rpm and the supernatant was withdrawn. Both, medium and spheroplasts, that were resuspended in the medium used for labeling, were frozen and processed for subsequent immunoprecipitation with anti-CPY and anti-ALP antibodies as described above. Quantitation was performed on short exposures of fluorograms by densitometric scanning. The values for pCPY and mCPY in the intracellular and extracellular fraction were added and considered as 100% total.

Cell extracts from overnight cultures grown at 24°C were prepared by collecting cells (50 OD units), washing them once with water and resuspending them to 1-2 OD/ml in 0.2N NaOH, 0.5% β-mercaptoethanol. After a 10-min incubation on ice, TCA was added to 10% final concentration and proteins were precipitated on ice for 30 min. The precipitates were collected by centrifugation, washed once with cold acetone and resuspended in Laemmli sample buffer (10 OD/ml). After SDS-PAGE on 8% gels the proteins were transferred to nitrocellulose membranes (Towbin et al., 1979).

Results

Identification of Three S. cerevisiae Genes, YPT51, YPT52, and YPT53, That Encode Proteins with Extensive Homology to rab5 and S. pombe ypt5p

Mammalian cells express rab5A (to which we refer as rab5) and two isofoms, rab5B and rab5C (Chavrier et al., 1992; Buccì, C., A. Lütcke, V. Olkkonen, P. Dupree, K. Simons, and M. Zerial, manuscript in preparation) which share sequence identity in the range of 85%. Recent data indicate that, similar to rab5, rab5B and rab5C regulate trafficking in the early endocytic pathway. To identify putative homologs of rab5 in S. cerevisiae we first searched for an amino acid sequence motif specific for the rab5 subgroup. Comparative analysis of all the available protein sequences revealed that the LAPMYYR motif is shared by all three mammalian rab5 proteins whereas it is absent from all other rab/Ypt proteins. This sequence is located within the helix α2 of small GTPases (nomenclature according to Pai et al., 1989), which is highly conserved between mammalian rab proteins and their yeast counterparts, such as rab1A, rab1B and Ypt1p, and which contributes to the functional specificity of rab5 (Stenmark et al., 1994). Two degenerate oligonucleotides corresponding to the LAPMYYR sequence and to the GXXXGKS/T motif found in the phosphate-binding loop of all rab/Ypt proteins (Chavrier et al., 1992) were used in the polymerase chain reaction (PCR) to amplify a putative YPT5 gene(s) from S. cerevisiae genomic DNA. As determined by Southern blot analysis, upon labeling with ³²P, the PCR reaction products hybridized to two distinct genomic DNA fragments (not shown). The respective full-length clones were isolated from a genomic library and shown to represent two novel sequences. Analysis of the two identified clones revealed two intronless genes encoding small GTPases of the...
Amino acid sequence of Ypt51p (A), Ypt52p (B) and Ypt53p (C) and numbers of identity/homology shared between rab5 and Ypt proteins (D). The boxed clear regions in A, B, and C represent the a2 domain, the hatched segments correspond to the effector domain (see Results). The nucleotide sequence data are available from EMBL/GenBank/DDBJ under the accession numbers X76173 SCYPT51, X76174 SCYPT52, and X76175 SCYPT53, respectively. In D, the values of identity (upper figure)/homology (lower figure) for the different protein pairs were determined by the gap program of the GCG package.

Figure 1. Amino acid sequence of Ypt51p (A), Ypt52p (B) and Ypt53p (C) and numbers of identity/homology shared between rab5 and Ypt proteins (D). The boxed clear regions in A, B, and C represent the a2 domain, the hatched segments correspond to the effector domain (see Results). The nucleotide sequence data are available from EMBL/GenBank/DDBJ under the accession numbers X76173 SCYPT51, X76174 SCYPT52, and X76175 SCYPT53, respectively. In D, the values of identity (upper figure)/homology (lower figure) for the different protein pairs were determined by the gap program of the GCG package.

rab/Ypt protein family (Fig. 1) with the closest identity and phylogenetic relationship to rab5 and S. pombe ypt5p (data not shown). Based on these features, but in keeping with the conventional nomenclature used for S. cerevisiae genes, the two genes were named YPTS1 and YPT53 (see below). Independently, sequencing of the S. cerevisiae chromosome XI revealed an open reading frame, that codes for another protein with a high sequence homology to rab5, but which was distinct from YPTS1 and YPT53. This gene was subsequently named YPT52 (Fig. 1). Although YPT52 contains the nucleotide sequence encoding the LAPMYYR motif, it was not identified by the PCR approach. This can probably be explained by three mismatches at the 3' end of the degenerate oligonucleotide representing the GXXGXGKS/T motif. Because all gene products were equally similar to rab5, the gene with the major importance in respect to phenotypic alterations (see below) was called YPT51, followed by YPT52 and YPT53.

The overall sequence identity/homology between all possible pairs of rab5/Ypt proteins was very similar and in the range of 54-70% identity, and 65-87% homology, respectively. Computer analysis of all available rab/Ypt protein sequences to calculate their phylogenetic relationship revealed that mammalian rab5A, B, and C, S. pombe ypt5p and S. cerevisiae Ypt51p, Ypt52p and Ypt53p constitute a group among the members of this family (data not shown). Further support for their close relationship is provided by the striking similarity in the effector domain, which in the case of p21ras is known to facilitate interactions with the GTPase-activating protein (Adari et al., 1988; Cales et al., 1988). Ypt52p and S. pombe ypt5p contain identical effector domains (RES-TIGAAF) (Fig. 1), which differ from rab5 in only one amino acid. Ypt51p and Ypt53p, which contain identical effector domains, have two amino acid differences when compared with Ypt52p and rab5, respectively (Fig. 1).

Analysis of the codon usage, which is a reliable indicator of protein abundance in yeast (Bennetzen and Hall, 1982), revealed that on a scale in which 1 represents exclusive use of preferred codons, YPTS1 scored 0.18, YPT52B scored 0.23, and YPT53 scored 0.07. This suggests that YPT53, most likely, encodes an extremely rare protein, whose putative low abundance is slightly above that of transcription factors (0-0.05), whereas YPTS1 and YPT52 encode proteins which lie in the typical range of other Ypt proteins, like Ypt7p (0.14) and Sec4p (0.21).

These data indicate that S. cerevisiae YPTS1, YPT52, and YPT53 are novel sequences which code for small GTPases sharing structural features with mammalian rab5A, B, and C proteins.

Disruption Of YPTS1 And YPT52 Has A Synergistic Inhibitory Effect on Cell Growth

To determine the function of Ypt51p, Ypt52p, and Ypt53p, we first constructed mutants with null alleles of YPTS1, YPT52, and YPT53 alone, all possible pairs of double mutants and the triple ypt51ypt52ypt53 null mutant. To generate single null mutants the disrupted genes and a selectable marker on a linear fragment were used to replace the respective wild-type genes in diploid strains, that were auxotrophic for the different markers. After sporulation and tetrad dissection it was observed that haploid progeny carrying either wild-type or the disrupted gene for YPTS1, YPT52, or YPT53 formed colonies of similar size after growth at 24°C, suggesting that all three YPT genes are non-essential. All possible combinations of double mutants and the triple mutant were obtained by crossing haploid null mutants, followed by tetrad dissection of the respective diploids. The haploid double and triple mutants also did not show any growth defect upon incubation at 24°C. However, when precultures of wild-type and mutant cells grown at 24°C were diluted into YPD medium at 37°C, Δypt51 mutants were delayed in growth, whereas Δypt52 and

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\(\Delta ypt53\) mutants propagated like wild-type cells (Fig. 2 A). Disruption of \(YPT53\) in combination with \(YPTS1\) or \(YPTS2\) did not change the growth properties of the latter two single mutants (Fig. 2 B). In contrast, in the double \(ypt51ypt52\) mutant growth was further inhibited at 37°C (Fig. 2 B) as compared with the single \(ypt51\) null mutant, and a mild delay of growth could already be observed at 30°C (not shown). The triple \(ypt51ypt52ypt53\) mutant was viable and had similar growth properties to the double \(ypt51ypt52\) mutant (compare Fig. 2 B with A).

**ypt51, ypt52, and ypt53 Mutants Internalize \(\alpha\)-Factor Normally, but Are Defective in Pheromone Degradation**

To investigate a possible role of the three \(Ypt\) proteins in endocytosis, the internalization of the mating pheromone \(\alpha\)-factor was analyzed. \(^{35}\)S-\(\alpha\)-Factor internalization assays were performed at 30°C with cells that had been grown at 24°C. At those temperatures no major growth defects could be observed, which might negatively influence the assay. The results showed that none of the mutants were defective for \(\alpha\)-factor binding and internalization (data not shown). Both the rate of internalization and the total amount of endocytosed pheromone in all three single and the triple null mutants were indistinguishable from wild-type cells.

The degradation of internalized \(\alpha\)-factor was then analyzed by separating the intact and degraded forms of \(^{35}\)S-labeled pheromone by thin layer chromatography. Similar to wild-type cells, that had completely degraded the pheromone after 30 min of internalization (Fig. 3 A), single null mutants of \(YPTS2\) and \(YPTS3\) degraded the pheromone with normal kinetics (data not shown). \(ypt52ypt53\) mutants revealed a minor delay in pheromone degradation, since small amounts of intact \(\alpha\)-factor were still present after 30 min of incubation at 30°C (Fig. 3 B). In contrast, pheromone degradation was strongly inhibited in \(ypt51\) mutants (Fig. 3 C). The \(t_{1/2}\) for \(\alpha\)-factor degradation was prolonged to about 50 min as compared with 10 min in wild-type cells. \(ypt51ypt52\) (Fig. 3 E) and \(ypt51ypt3\) (Fig. 3 D) mutants displayed a further delay in \(\alpha\)-factor degradation as compared to single \(ypt51\) mutants, with a \(t_{1/2}\) of pheromone degradation of about 60 and 75 min, respectively. Triple \(ypt51ypt52ypt53\) mutants (Fig. 3 F) seemed to be almost blocked in pheromone degradation, since the \(t_{1/2}\) was more than 120 min.

These results indicate that disruption of the \(YPTS1\) gene does not affect \(\alpha\)-factor internalization but causes a severe inhibition of pheromone degradation. Furthermore, this effect is aggravated by the additional disruption of \(YPTS2\) and \(YPTS3\).

**ypt51, ypt52, and ypt53 Mutants Are Defective in Vacuolar LY Accumulation**

The observed lack of pheromone degradation in the here described \(ypt\) mutants could be explained by an inhibition of transport from an endocytic compartment to the site of degradation, the vacuole. To test this hypothesis, we investigated the ability of the various mutants to deliver LY to the vacuole. As compared with wild-type cells (Fig. 4 A), single mutants for \(YPTS2\) (Fig. 4 B) and \(YPTS3\) (not shown) revealed no defect in the accumulation of the fluorescent dye in the vacuole. In contrast, \(\Delta ypt51\) mutants were characterized by a strong reduction in the amount of LY accumulation in the vacuole (Fig. 4 C). The majority of cells showed a very weak or no detectable fluorescence in the region of the vacuole, which was generally recognizable by Nomarski optics. Only in rare cases did LY accumulate in the vacuole in similar amounts as observed for wild-type cells (Fig. 4 C). Interestingly, disruption of both \(YPTS1\) and \(YPTS2\) further decreased LY transport to the vacuole (Fig. 4, E and F), since even the occasionally observed weak fluorescence of vacuoles could not be detected. This phenotype was especially noteworthy in light of the fact that the majority of mutant cells contained the typically large vacuole, as seen in wild-type cells. Occasionally, cells of double as well as triple mutants revealed a remarkably bright intracellular fluorescence. This bright staining could sometimes be resolved into tiny spots (Fig. 4, D, E, and F, arrowheads). As detected by Nomarski optics, these particular cells did not contain a large vacuole like wild-type cells, but rather exhibited a granular surface similar to class C vps mutants lacking vacuoles (Banta et al., 1988). \(ypt51ypt53\) mutants appeared very similar to \(ypt51\) mutants, in that occasionally a weak vacuolar staining by LY could be observed (Fig. 4 D). However, in difference to the latter mutant a few cells showed the spotty fluorescence as described for the \(ypt51ypt52\) mutant (Fig. 4, E and F, arrowheads). Therefore, additional disruption of \(YPTS3\) in \(\Delta ypt51\) mutants resulted in a phenotype which was stronger than observed for \(ypt51\) and weaker than...
Figure 3. α-factor degradation by ypt mutants as compared to wild-type cells. α-cells of the respective strains were grown overnight at 24°C to early logarithmic stage. After incubation with 35S-α-factor for 1 h at 0°C the cells were collected and resuspended in YPD medium and transferred to a 30°C waterbath to start internalization of the pheromone. Total cell-associated (pH 1.2 washed cells) (i) and internalized α-factor (pH 6 washed cells) (d) was extracted at the times indicated and resolved by thin layer chromatography. The positions of the intact (i) and degraded (d) pheromone are indicated. The fluorograms were exposed for 20 d at -70°C. A, wild type; B, ypt52ypt53; C, ypt51; D, ypt51ypt53; E, ypt51ypt52; F, ypt51ypt52ypt53.

observed for ypt51ypt52 mutants. Although the majority of cells with a disrupted copy of both YPT52 and YPT53 showed no defect in LY accumulation (data not shown), it appeared that the number of cells without dye accumulation was increased as compared to wild-type cells. Whereas typically ~95% of wild-type, ypt52 and ypt53 cells revealed staining of the vacuole with LY, this number was reduced to about 50% in double ypt52ypt53 mutants.

ypt51, ypt52, and ypt53 Mutants Display a Delay in the Maturation of a Soluble and a Membrane-bound Vacuolar Enzyme and also Partially Missort a Soluble Vacuolar Marker Protein

As in mammalian cells, the endocytic and the vacuolar sorting pathways in S. cerevisiae have been proposed to intersect in a prevacuolar, endosomal compartment (Vida et al., 1993; Schimmöller and Riezman, 1993; Davis et al., 1993). The finding that the here described Ypt proteins are required for transport of LY and possibly of α-factor to the vacuole suggests that delivery of enzymes to the vacuole via the secretory pathway could also be affected in the ypt51, ypt52, and ypt53 mutants. To test this, transport of the vacuolar hydrolyases carboxypeptidase Y (CPY) and alkaline phosphatase (ALP) to the vacuole was analyzed. The transit of soluble CPY through the secretory pathway can be monitored by the appearance of its different intermediate forms. Cleavage of the signal peptide and core glycosylation in the ER generate a 67-kD form, p1CPY, which is transported to the Golgi complex, where the core oligosaccharides are extended, yielding p2CPY (69 kD). Finally, just before or upon arrival of p2CPY in the vacuole the NH2-terminal prosequence is removed to yield mature (m) CPY (61 kD) (Stevens et al., 1982) (Fig. 5). During biosynthesis of membrane-bound ALP, conversion of the 76-kD Golgi precursor (pALP) into...
Figure 4. Accumulation of LY-CH by wild-type (A), Δypt52 (B), Δypt51 (C), ypt51ypt53 (D), and ypt51ypt52 (E and F) cells. After growth of cells overnight at 24°C to early logarithmic stage LY-CH internalization was performed at 30°C for 1 h. The cells were subsequently washed, mounted in low melting agarose, and visualized using Nomarski and fluorescence optics. Bar, 5 μm.
Figure 5. Carboxypeptidase Y biogenesis in ypt mutants and wild-type cells. Cells were labeled for 5 min at 30°C using [35S]methionine and the chase was subsequently initiated by the addition of ammonium sulfate, unlabeled methionine and cysteine. At 0 (lane 1), 2.5 (lane 2), 5 (lane 3), 10 (lane 4), 20 (lane 5), 30 (lane 6), and 60 (lane 7) min of chase aliquots of cells were removed and processed for immunoprecipitation with anti-CPY antibodies. Immunoprecipitates were analyzed on 8% gels by SDS-PAGE and fluorography. The fluorograms were exposed for 5 d at -70°C. p, precursor; m, mature form. Other background bands are nonspecific. a, wild type; b, ypt52; c, ypt52ypt53; d, ypt51; e, ypt51ypt52; and f, ypt51ypt52ypt53.

The 71-kD mature form (mALP) occurs in the vacuole by PEP4-dependent processing (Klionsky and Emr, 1989).

Pulse-chase labeling with [35S]methionine was performed with cells that had been grown at 24°C. After a pulse of 5 min at 30°C, the cells were chased for various periods of time at 30°C, and the radioactive CPY and ALP were successively immunoprecipitated from total cellular protein and subsequently analyzed by SDS-PAGE. Conversion of p1 to p2CPY and maturation to mCPY took place with normal kinetics in Δypt52 (Fig. 5) and Δypt53 (not shown) mutants as compared with wild-type cells. In ypt52ypt53 double mutants, despite of a 5-min delay in the conversion from p1 to p2 CPY, processing to the mature form occurred normally. Interestingly, Δypt51 mutants exhibited a clear delay in the processing of p2 to mCPY. While in wild-type cells ~50% of p2CPY was processed to mCPY by 10 min of chase, in mutant cells a similar degree of processing was only observed after 20–30 min of chase. ypt51ypt52 and ypt51ypt52ypt53 mutants were characterized by a further delay in this maturation step, since a chase of approximately 60 min was required to yield 50% of both p2 and mCPY (Fig. 5).

Maturation of membrane-bound ALP was also perturbed in mutant cells lacking Ypt proteins. However, while in ypt51ypt52 and ypt51ypt52ypt53 mutants 50% of mALP was formed only after 20 min, as compared with 5–10 min in wild-type cells, in all other mutants the processing of this protein took place with wild-type kinetics (data not shown). Thus, maturation of ALP appeared to be less severely perturbed in the here described ypt mutants than maturation of CPY.

In view of the observed delay in the maturation of vacuolar enzymes, the possibility had to be considered that a fraction of vacuolar enzymes is not properly sorted to the vacuole, but instead missorted to the extracellular space, a phenotype associated with a number of vacuolar protein sorting (vps) mutants (Robinson et al., 1988; Rothman et al., 1989a). To examine this, spheroplasts of the various mutants and of wild-type cells were pulse labeled for 15 min and chased for 30 min. After termination of the chase period by addition of NaN3 and NaF, the intracellular and extracellular fractions were separated and two successive immunoprecipitations were performed with CPY and ALP antisera. As shown in Fig. 6 A, wild-type, ypt52, ypt53 and ypt52ypt53 cells contained mainly mCPY which was present in the intracellular fraction, suggesting proper targeting to the vacuole. In contrast, substantial amounts of Golgi-modified p2CPY (~40% of total) was found in the extracellular fraction of Δypt51 mutants, providing evidence for missorting of this soluble enzyme. In fact, vps21 (Robinson et al., 1988) is a mutant allele of YPT51 (Emr, S., personal communication).

Figure 6. Sorting of the vacuolar enzymes CPY (A) and ALP (B) and steady state levels of mCPY (C). (A and B) Cells of the respective strains were converted into spheroplasts, labeled for 15 min with [35S]methionine and chased for 30 min using conditions described in Fig. 8. To analyze secreted proteins the spheroplasts were separated from the medium by centrifugation at 1,500 rpm for 5 min. The cell and the medium factions were processed for successive immunoprecipitation using antibodies against CPY (A) and ALP (B). I, intracellular; E, extracellular; p, precursor; m, mature form. Other background bands are nonspecific. (C) Protein extracts of the various strains were prepared from overnight cultures grown at 24°C and separated by SDS-PAGE. Western analysis was performed with antibodies against CPY and the ECL detection system. m, mature CPY. a, wild type; b, ypt53; c, ypt51; d, ypt52; e, ypt51ypt53; f, ypt52ypt53; g, ypt51ypt52; h, ypt51ypt52ypt53.
tional disruption of either YPT52 or YPT53 increased the ratio of external to internal CPY suggesting enhanced missorting. Approximately 55 and 50% of total CPY was present extracellularly in ypt51ypt52 and ypt51ypt53 mutants, respectively. Furthermore, increasing amounts of intracellular p2CPY versus mCPY accumulated in ypt51ypt53 and even more in ypt51ypt52 mutants. The triple ypt51ypt52ypt53 mutants displayed the highest degree of CPY missorting (~60% of total) and the strongest defect in precursor accumulation.

Although upon pulse–chase labeling of ypt51ypt52 and ypt51ypt52ypt53 mutants small but clearly detectable quantities of mCPY could be detected, as shown in Fig. 5, in the experiment of Fig. 6 A the amounts of mCPY seemed to be reduced in favor of increased quantities of p2CPY. This apparent discrepancy in the results most likely can be ascribed to the fact that spheroplasts on the one hand (more precursor accumulation) and intact cells on the other hand (less precursor accumulation) were used. A similar result was obtained in studies of vacuolar protein maturation with vacuolar acidification (vps) mutants (Klionsky et al., 1992).

As expected for a membrane protein, missorting of ALP to the extracellular medium could not be detected in any of the different ypt mutants (Fig. 6 B). As seen in pulse–chase labeling experiments (data not shown), precursor accumulation could be detected only in ypt51ypt52 and ypt51ypt52ypt53 mutants.

We also determined the steady state levels of mCPY and mALP in the various mutants as compared with wild-type cells. This was done by Western analysis of protein extracts prepared from cultures grown overnight at 24°C. Surprisingly, ypt51 mutants revealed no detectable amounts of mCPY as compared with wild-type cells and the other single ypt mutants (Fig. 6 C). This is in apparent contrast to the results obtained by pulse–chase labeling experiments, in which the generation of mCPY could clearly be observed in the YPT51-disrupted cells, and suggests a reduced stability of mCPY over longer periods of time. The steady-state levels of mALP were reduced to ~50 % of wild-type levels in ypt51 and ypt51ypt52ypt53 mutants (data not shown).

To exclude the possibility that in the absence of Ypt51p, Ypt52p, and Ypt53p the secretory transport pathway was affected in a more general way, we also investigated the secretion efficiency of invertase and the mating pheromone α-factor. A comparison of wild-type cells with the different ypt null mutants revealed that both marker proteins were secreted with normal kinetics (data not shown).

In conclusion, while these results reveal no secretion defects and only slight perturbations of ALP maturation in ypt51ypt52 and ypt51ypt52ypt53 mutants, the kinetics of CPY maturation were increasingly impaired by the successive disruption of YPT51, YPT52, and YPT53, and missorting of CPY was simultaneously increased. Furthermore, it appeared that the steady state level of mCPY was strongly reduced in Δypt51 mutants.

**Acidification Properties Are Disturbed in the Here Described ypt Mutants**

In view of the missorting defect associated with some of the ypt mutants the possibility existed that this phenotype could be related to improper vacuole acidification. Precedents for this are a number of vps mutants (Banta et al., 1988; Preston et al., 1989; Rothman et al., 1989b) and specific mutants lacking subunits of the vacuolar ATPase (Nelson and Nelson, 1990; Preston et al., 1989; Yamashiro et al., 1990). Missorting of vacuolar enzymes can also be caused by raising the pH specifically of the vacuolar system by treatment with the drug bafilomycin A1 (Banta et al., 1988; Klionsky and Emr, 1989). Vacuole acidification can be microscopically detected by staining living yeast cells with the lysosomotropic agent quinacrine, a fluorescent dye that accumulates in acidified organelles (Weisman et al., 1987).

As shown in Fig. 7 A, wild-type cells revealed a bright vacuolar fluorescence, as did ypt52, ypt53, and ypt52ypt53 null mutants (not shown). In contrast, this staining was strongly reduced in cells with a disrupted copy of YPT51 (Fig. 7 B), and was virtually absent in ypt51ypt52 (not shown) and ypt51ypt52ypt53 mutants (Fig. 7 C). This clearly supports the idea that vacuole acidification is disturbed in the here described ypt mutants.
demonstrated that the absence of Ypt51p and Ypt52p results in a severe acidification defect of the vacuolar system.

To exclude the possibility that improper vacuole acidification could have been the primary reason for the observed lack of α-factor degradation rather than an endocytic delivery defect, pheromone degradation was analyzed in vat2 mutants disrupted for a subunit of the vacuolar ATPase. These cells display similar defects in vacuole acidification than YPT51-disrupted mutants. Interestingly, in spite of a delay in α-factor degradation to a t½ of 16 min in vat2 cells (data not shown) as compared with wild-type cells (10 min), this delay was much less severe than observed for ypt51, and especially for ypt51ypt53, ypt51ypt52, and ypt51ypt52ypt53 mutants (see above). It is therefore highly unlikely that the lack of α-factor degradation in the here described ypt mutants was mainly caused by a lack of acidification.

Vesicle Accumulation and Vacuolar Morphology in ypt51, ypt52, and ypt53 Mutants

As Ypt51p, Ypt52p, and Ypt53p are likely to act in the endocytic and vacuolar protein sorting pathways, a morphological investigation by electron microscopy of ypt51 and ypt51- ypt52ypt53 mutants as compared to wild-type cells was initiated. Potassium permanganate fixation and staining was chosen, as it allows visualization of membrane structures without the need to remove the cell wall. It was observed that the ypt51 null mutant (Fig. 8, B and D) as well as the triple

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**Figure 8.** Accumulation of vesicular structures in Ypt51p-depleted cells. Wild type cells (A), ypt51 (B and D), and ypt51ypt52ypt53 (C and E) null mutants were fixed with potassium permanganate and viewed by thin section electron microscopy. Arrows point to 40–60-nm vesicles. E, endoplasmic reticulum; G, Golgi; M, mitochondria; N, nucleus; V, Vacuole; W, cell wall. Bars: (A-C) 1 μm; (D and E) 0.5 μm.
yp51yp52yp53 mutant (Fig. 8 C and E) accumulated small vesicular structures with a diameter of ~40–60 nm. Compared with wild-type (Fig. 8 A), cells of both mutant strains contained on average 6–7-fold more of these vesicles. With respect to size, these structures resembled the vesicles accumulating in yeast mutants that are blocked in transport between the ER and the Golgi (Kaiser and Schekman, 1989; Becker et al., 1991). As we did not find any evidence for an impairment of CPY transport to the Golgi or for inhibition of protein secretion in Ypt51p-depleted cells, these accumulating type of vesicles are likely to represent transport intermediates between organelles other than the ER and the Golgi, perhaps endosomes.

Besides these morphological changes, thin section EM analysis did not reveal any other gross morphological changes, for example of the vacuolar system. However, due to the changed acidification properties and the appearance of small structures positive for LY in some of the ypt mutants, we also analyzed the vacuolar morphology on a larger cell population by light microscopy, using the vital stain carboxydichlorofluorescein diacetate (CDCFDA). Vacuolar labeling presumably results from diffusional uptake and subsequent hydrolysis of the dye by non-specific esterases, generating a fluorescent hydrolysis product (Pringle et al., 1989). In contrast to the microscopic analysis performed with LY, in which the cells were treated with energy poisons to stop endocytosis at a certain stage, these experiments were performed with metabolically active cells. Under these “optimal” conditions, typical vacuoles appear as multiple irregularly shaped organelles clustered in a certain region of the cell (Conradt et al., 1992) (Fig. 9, A, B, K, and L). During certain stages of the cell cycle a process termed vacuole inheritance induces the formation of so-called vacuolar segregation structures, which are projections formed by mother cell vacuoles leading into the developing bud (Weisman et al., 1987) (Fig. 9, B, C, and K, arrowheads).

Typical vacuoles and vacuolar segregation structures (arrowheads), as found in wild-type cells, were also observed in ypt52, ypt53 (not shown), and ypt52yp53 null mutants (Fig. 9, C and D). In contrast to the convoluted vacuolar shapes of wild-type cells, vacuoles of ypt51 mutants appeared completely spherical and as single structures (Fig. 9 E). Furthermore, the typical “tracks” formed by vacuolar material leading into the developing bud were never observed in these mutants, although larger buds clearly contained vacuoles. The spherical type of vacuole was also predominantly found in ypt52yp53 mutants (Fig. 9 F). However, numerous unusually small and bright structures were occasionally observed in these cells (Fig. 9 G). This type of staining became more frequent in ypt51yp52 and ypt51yp52yp53 mutants (Fig. 9, H and J). Analysis of these cells by Nomarski optics revealed that the typical larger vacuolar organelles were replaced by much smaller structures. Confocal microscopy further resolved these brightly fluorescent entities into structures that seemed partly interconnected to form a CDCFDA-positive network (Fig. 9, M and N).

In summary, while EM analysis revealed the accumulation of small vesicular structures in thin sections of ypt51 and ypt51yp52yp53 mutants, subtle and differential morphological alterations of the vacuolar system could be observed in a small subset of the various ypt mutants with the help of a vital stain and light microscopy, which enables analysis of a large fraction of cells.

Discussion

Structural and Functional Similarities between Ypt51p, Ypt52p, and Ypt53p and Mammalian rab5 Proteins

In the present study three new members of the Ypt/rab family, Ypt51p, Ypt52p, and Ypt53p, were identified in S. cerevisiae, that share a high degree of identity/homology to rab5. First, a significant identity exists between the sequences of each of these Ypt protein and rab5 (48–54%). The identity is not as high as that between Yptlp and rab1A (71%) (Haubruck et al., 1987), or Ypt7p and rab7 (63%) (Wichmann et al., 1992). On the other hand, rab8, which is required for transport from the TGN to the plasma membrane (Huber et al., 1993), thus providing evidence for functional similarity to Sec4p, is the closest structural homolog of Sec4p. Second, some of the structural elements that confer functional specificity to rab/Ypt proteins (located in the a2-helix [LAPMYVR] and effector domain [Stouten et al., 1993]) (Brennwald and Novick, 1993; Dunn et al., 1993; Haubruck et al., 1989; Hengst et al., 1990; Stenmark et al., 1994) are highly conserved between rab5 and Ypt51p, Ypt52p, and Ypt53p. Third, the phylogenetic analysis, which reflects the sum of similarities of individual sequence elements, clearly indicates that the rab5 proteins and the proposed S. pombe and S. cerevisiae homologs constitute a subgroup among all known small GTPases of the Ypt/rab family.

In addition to protein similarity, our data support the notion that, similar to rab5, the here described Ypt proteins are involved in the regulation of transport in the endocytic pathway. A strong inhibition of LY accumulation in the vacuole was observed for the single Δypt51 mutant, which was further aggravated by the additional deletion of YPT52 and YPT53. Furthermore, although ypt51, ypt51yp52, and ypt51yp52yp53 null mutants displayed normal a-factor internalization rates, they were strongly impaired in a-factor degradation. As this process occurs in the vacuole, also this phenotype could be explained by an endocytic transport defect. This hypothesis is supported by the finding that, so far,
endocytic mutants (with recognizable vacuole) have all been shown to be defective in transport of both LY and α-factor (Kübler and Riezman, 1993; Raths et al., 1993). However, the surprising finding that Δypt51 mutants displayed a dramatic reduction in the intracellular levels of mCPY under steady state conditions might suggest that inhibition of pheromone degradation could also be the result of a defect in vacuolar protease activity. While we cannot completely rule out this possibility, we think it is unlikely. The observation that proteinase A-dependent maturation of CPY and ALP is affected but does take place (even in ypt51ypt52 and ypt51ypt52ypt53 mutants) suggests that the processing and degradative potential of the various ypt mutants is not severely impaired. Therefore, since pheromone degradation also depends on proteinase A (Singer and Riezman, 1990), it should take place if α-factor were present in the vacuole. Thus, we favor the idea that the lack of α-factor degradation in the here described ypt mutants is due to impaired transport to the site of degradation.

In conclusion, these data suggest that Ypt51p, Ypt52p, and Ypt53p are structurally and functionally related to rab5, although, without complementation analysis, we cannot definitively state at the present time that they correspond to the yeast counterparts of rab5A, B, and C.

**Multiple Phenotypes of ypt51, ypt52, and ypt53 Mutants Suggest an Intersection of the Endocytic and Vacular Protein Sorting Pathways**

Besides the defect in the delivery of endocytic markers to the vacuole, the various ypt mutants displayed phenotypes that are reminiscent of vps mutants (Raymond et al., 1992). These include a delay in the maturation of vacuolar hydrolases, partial misrouting of the Golgi form of vacuolar CPY to the extracellular space, acidification defects and subtle morphological alterations of the vacuolar system. In fact, vps21 (Robinson et al., 1988) represents a mutant allele of *YP51* (Emr, S., personal communication).

In light of the multiple phenotypes displayed by the here described ypt mutants, the question could be raised whether the defect in acidification may be responsible for the missorting of p2CPY and the inhibition of α-factor degradation. Previous studies have revealed that lack of vacular acidification has severe implications on intracellular transport pathways. Treatment of cells with bafilomycin A1, a specific inhibitor of the vacular ATPase, results in missorting of CPY (Banta et al., 1988) to the same extent as observed for the Δypt51 mutants. However, neither missorting of ALP (Klionsky and Emr, 1989) nor a major defect in α-factor degradation occurs (Singer and Riezman, 1990). In mutants disrupted for a gene encoding a subunit of the vacular ATPase (*vat2* or *vatb*) the extent of missorting of CPY to the extracellular medium was similar to that of bafilomycin A1-treated cells, while maturation of p2CPY to mCPY was further delayed and precursor ALP accumulated (Klionsky et al., 1992). Nevertheless, even if α-factor degradation was delayed in vat2 cells, its *t1/2* was only 16 min, whereas in Ypt51p-depleted cells it was prolonged to 48 min. Therefore, while it is possible that the acidification defect in the here described ypt mutants leads to missorting and precursor accumulation of vacuolar enzymes, it seems unlikely that it could be the primary reason for lack of α-factor degradation.

It rather seems that the phenotypes displayed by ypt51, ypt52, and ypt53 mutants could well be explained by a transport defect affecting an endocytic organelle which is connected with the vacuolar sorting pathway. Experimental evidence for the meeting of the two pathways comes from cell-fractionation analysis, which reveals that p2CPY and endocytosed α-factor cofractionation in a prevacuolar/endo- somal compartment (Vida et al., 1993). Independently, studies with the ypt7 null mutant have suggested that meeting of endocytosed α-factor with vacuolar hydrolases might occur before delivery to the vacuole, since a PEP4-dependent and energy-independent pheromone degradation can be observed in this mutant (Schimmöller and Riezman, 1993). Finally, the finding that Ren1p, a protein identified for its role in the delivery of α-factor receptor to the vacuole, is identical to Vps2p (Davis et al., 1993), which is required for vacuolar protein sorting, also provides further evidence for the intersection of the endocytic with the vacuolar protein sorting pathway.

**Site of Function of Ypt5 Proteins**

Due to the complexity of the ypt51-, ypt52-, and ypt53-associated phenotypes, it is too early to pinpoint the precise step controlled by these Ypt proteins. However, phenotypic comparison between Δypt7 and the here described ypt mutants provides some clues about the order in which these small GTPases might function. Ypt7p, the yeast homolog of mammalian rab7 which is associated with late endosomes (Chavrier et al., 1990), has been suggested by Wichmann et al. (1992) to play a role in the endocytic transport of α-factor to the vacuole. As in the here described ypt mutants, indirect evidence for this was a substantial delay of endocytosed pheromone degradation in a Δypt7 mutant. A number of observations support the idea that Ypt51p, Ypt52p, and Ypt53p proteins may act at an earlier step in the endocytic pathway than Ypt7p. First, by comparing the patterns of degradation in ypt7 and ypt51, ypt52, and ypt53 mutants obtained under the same experimental conditions, it becomes clear that α-factor degradation is more severely inhibited in the latter mutants. Second, vacuole fragmentation can be predominately observed in Δypt7 mutants, but is only occasionally seen in the here described ypt mutants. Third, since in yeast at least two kinetic endosomal intermediates (early and late endosomes) can be distinguished (Singer and Riezman, 1990; Singer-Krüger et al., 1993), it is interesting to note that recent studies on the ypt7 null mutant have revealed an accumulation of endocytosed α-factor in a gradient fraction enriched for late endosomes (Schimmöller and Riezman, 1993). This suggests that Ypt7p is required for transport from late endosomes to the vacuole (Schimmöller and Riezman, 1993). Conversely, preliminary cell fractionation experiments suggest that upon internalization of α-factor into the ypt51 null mutant, the pheromone accumulates in a prevacuolar endocytic compartment, most likely in early endosomes as defined by Singer-Krüger et al. (1993) (Singer-Krüger, B., and M. Zerial, unpublished observations). The proposed order of function of Ypt51p and Ypt7p would be in accordance with the proposed successive function of mammalian rab5 and rab7. Future experiments, like epistasis analysis with ypt51ypt7 double mutants, will hopefully help to test this working hypothesis.
While the internalization rate of transferrin can be modulated by rabs5 in mammalian cells (Bucci et al., 1992), it is at present still unclear why the deletion of YPT51, YPT52, and YPT53 does not affect the internalization rate of α-factor in yeast.

**Morphological Changes in ypt51, ypt52, and ypt53 Mutants**

Analysis of the vacuolar morphology of the various ypt mutants using CDFCA staining in living cells and light microscopy revealed that the ypt51 mutant contained a single large vacuole, which was completely spherical, in contrast to the convoluted vacuolar structures observed in wild-type cells. In addition, vacuolar segregation structures were not found in ypt51 mutants. The single, spherical type of vacuole was also found predominantly in the ypt51ypt52, ypt51ypt53, and ypt51ypt52ypt53 mutants. However, in the latter mutants, a small but significant number of cells displayed fragmented vacuolar structures. The observation that these structures were also labeled with LY would favor an endosomal origin. Simultaneous disappearance of typical vacuolar elements in these particular cells would then support the idea that vacuole biogenesis depends on proper endocytic and vacuolar protein transport, as discussed before.

Furthermore, the 40-60-nm vesicles which accumulate in ypt51 and ypt51ypt52ypt53 mutants, revealed by EM analysis, may correspond to transport intermediates of the endocytic and/or vacuolar protein sorting pathways. Their resemblance in size to vesicles accumulating in mutants that are blocked in transport between the ER and the Golgi might be rather coincidental, since no such transport defect could be observed in the here described ypt mutants. Definite proof for the origin of these vesicular structures will await more detailed ultrastructural and cell fractionation analysis. The finding that morphological changes of the vacuolar system were not observed by EM, most likely can be explained by the fact that first, these changes might be only visible in living cells, and second, that they are only apparent upon analysis of a large number of cells.

**Functional Redundancy of Ypt51p, Ypt52p, and Ypt53p?**

The existence of three similar Ypt proteins raises the question whether they might be functionally redundant or play specialized and distinct roles. Since in all respects the most severe defects were associated with the ypt51 mutant compared to the other two single mutants, it seems that Ypt51p is the most essential protein for the general endocytic pathway. On the other hand, detectable aggravation of all analyzed phenotypes (growth, endocytosis, vacuolar protein missorting, acidification, vacuolar morphology) upon additional disruption of YPT52 and YPT53 demonstrates that the three gene products have at least overlapping functions. The role of Ypt53p in endocytic traffic (or any other transport step) is more elusive than that for Ypt51p and Ypt52p. Its involvement in these transport pathways is, however, suggested in double ypt51ypt53, ypt52ypt53 and the triple ypt51ypt52ypt53 mutants, in which synergistic effects can be observed. Nevertheless, the sole presence of Ypt53p in ypt51ypt52 mutants does not complement the absence of the other two small GTPases, as no major phenotypic differences to the triple mutant could be detected. Therefore, it is possible that Ypt53p may have a very specialized, yet unresolved function. Interestingly, the severity of the mutant phenotypes correlates well with the predicted abundance of the respective proteins: Ypt51p and Ypt52p, which seem to be more important for the general endocytic pathway, have a higher value of theoretical protein abundance, whereas the low abundance calculated for Ypt53p would be in agreement with a specialized and less general function.

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Singer-Krüger et al. Ypt51p, Ypt52p, and Ypt53p and Endocytosis 297