Cell-free Reconstitution of Microautophagic Vacuole Invagination and Vesicle Formation

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Abstract. Many organelles change their shape in the course of the cell cycle or in response to environmental conditions. Lysosomes undergo drastic changes of shape during microautophagocytosis, which include the invagination of their boundary membrane and the subsequent scission of vesicles into the lumen of the organelle. The mechanism driving these structural changes is enigmatic. We have begun to analyze this process by reconstituting microautophagocytosis in a cell-free system. Isolated yeast vacuoles took up fluorescent dyes or reporter enzymes in a cytosol-, ATP- and temperature-dependent fashion. During the uptake reaction, vacuolar membrane invaginations, called autophagic tubes, were observed. The reaction resulted in the transient formation of autophagic bodies in the vacuolar lumen, which were degraded upon prolonged incubation. Under starvation conditions, the system reproduced the induction of autophagocytosis and depended on specific gene products, which were identified in screens for mutants deficient in autophagocytosis. Microautophagic uptake depended on the activity of the vacuolar ATPase and was sensitive to GTP$_\gamma$S, indicating a requirement for GTPases and for the vacuolar membrane potential. However, microautophagocytosis was independent of known factors for vacuolar fusion and vesicular trafficking. Therefore, scission of the invaginated membrane must occur via a novel mechanism distinct from the homotypic fusion of vacuolar membranes.

Key words: budding • autophagocytosis • membrane dynamics • proteolysis • scission

Introduction

Eukaryotic cells have developed multiple pathways for the intracellular degradation of proteins. The proteasome system and direct translocation of proteins through the lysosomal membrane account for only part of the protein turnover (Cuervo and Dice, 1998). A major route of degradation is autophagocytosis, a process of controlled self-digestion of cells. Autophagocytosis is constitutive, but it is strongly enhanced under conditions of stress, starvation, or redifferentiation (Seglen and Bohley, 1992; Knop et al., 1993). A wide range of eukaryotic cells are capable of autophagocytosis, indicating that the process may be a general characteristic of eukaryotic life.

Current models differentiate two forms of autophagocytosis, macro- and microautophagocytosis. Macrautophagocytosis is characterized by the appearance of a unique type of vesicles. These vesicles (autophagosomes) sequester portions of the cytosol. They can also engulf entire organelles such as mitochondria or peroxisomes. Autophagosomes are surrounded by two or sometimes several membranes that distinguish them from other cellular vesicles. The origin of the autophagosomal membranes is controversial (Dunn, 1990; Yamamoto et al., 1990a,b). The outer membrane of autophagosomes can fuse with lysosomes, delivering the inner membrane with its cytosolic contents (called an autophagic body) into the lysosomal lumen for degradation (Seglen and Bohley, 1992; Takeshige et al., 1992).

Microautophagocytosis is defined as the uptake of cytosolic components by direct invagination of the lysosomal membrane. It results in vesicles budding into the lumen of the lysosome. The resulting autophagic bodies are finally degraded. Also, the microautophagic pathway is capable of delivering not only soluble proteins, but entire organelles into lysosomes for degradation (Veenhuis et al., 1983; Tuttle et al., 1993; Yokota, 1993; Tuttle and Dunn, 1995). The mechanisms that drive the profound changes in lysosomal structure during invagination and scission of the nascent microautophagic vesicle are unknown.

Different approaches have been taken to analyze autophagocytosis in various systems. Pharmacological studies have defined agents inducing or inhibiting autophagocytosis in cell culture systems, perfused organs, or entire animals (Seglen and Bohley, 1992; Mortimore and Kadowaki, 1994). Considerable efforts were also made to iso-
late autophagosomes (Stromhaug et al., 1998), a prerequisite for analyzing their protein composition, identifying marker proteins, and analyzing their origin.

Pioneering genetic screens in *Saccharomyces cerevisiae* have revealed at least 16 complementation groups of mutants defective in the induction of autophagocytosis, providing a solid basis for the molecular analysis of the process (Tsukada and Ohsumi, 1993; Thumm et al., 1994). These complementation groups overlap strongly with a set of mutants defective in the cytoplasm to vacuole targeting (Cvt) pathway (Harding et al., 1996; Scott et al., 1996). The Cvt pathway leads to the selective and rapid import of aminopeptidase I into vacuoles via vesicular intermediates structurally related to autophagosomes (Baba et al., 1997). Most of the genes found in these screens have now been identified by exploiting the fact that autophagocytosis in yeast is essential for surviving starvation and sporulation. They encode presumptive regulatory components, such as a protein kinase (autophagocytosis [Apg]1p/autophagocytosis [Atu]3p) (Matsuura et al., 1997; Straub et al., 1997). A ubiquitin-like protein conjugation system was found comprising an activation enzyme (Apg7p), a ligase (Apg10p), and two proteins (Apg12p and Apg5p), which become covalently attached to each other by these enzymes (Mizushima et al., 1998; Kim et al., 1999; Shintani et al., 1999). A ubiquitin-activating enzyme is also involved in the autophagic pathway of mammalian cells (Lenk et al., 1992). Homologomers of Apg16p bind to the Apg5p–Apg12p conjugate (Mizushima et al., 1999). Aut2p and Aut7p/Apg8p were proposed to bind to microtubules and mediate the delivery of autophagosomes to the vacuole (Lang et al., 1998). However, this interpretation was challenged by the recent finding that disassembling microtubules by nocodazole treatment does not block autophagocytosis in yeast (Kirisako et al., 1999). Aut7p/Apg8p is a peripheral component on the membrane of autophagosome precursors and is the first marker protein for autophagosome induction in yeast (Kirisako et al., 1999). Genetic screens in *Pichia pastoris* and *Hansenula polymorpha* have identified another set of mutations affecting peroxisome degradation (Takeshige et al., 1995; Sakai et al., 1998; Yuan et al., 1999). One of the affected genes was Apg7, which provides genetic evidence that autophagocytosis of soluble proteins and peroxisomes may employ a common machinery (Kim et al., 1999; Yuan et al., 1999).

The macroautophagic pathway in *S. cerevisiae* has been extensively characterized by light and EM (Takeshige et al., 1992; Baba et al., 1994, 1995). The pathway depends on Vam7p and Vam3p, two SNARE (soluble N-ethylmaleimide–sensitive factor attachment protein receptor) proteins that are presumably involved in fusing the outer membrane of autophagosomes with the vacuole (Darsow et al., 1997; Sato et al., 1998). Therefore, macroautophagocytosis seems to involve the conserved fusion machinery that also mediates membrane fusion at many other steps of vesicular trafficking (Rothman, 1994). We have recently discovered a microautophagic pathway in *S. cerevisiae* (Müller et al., 2000, this issue), in which a specialized structure, the autophagic tube, mediates invagination of the vacuolar membrane and scission of vesicles into the lumen. This tube displays a striking lateral heterogeneity of membrane structure. The base of the tube has a normal content of transmembrane proteins, whereas the zone of vesicle formation at the tip of the tube is devoid of visible transmembrane particles. The smooth appearance of this zone matched that of the membrane of autophagic bodies. We also found small areas with very low content of transmembrane particles on the vacuolar surface. These smooth areas often invaginate and appear to be precursors of autophagic tubes.

To study the biochemical basis of the mechanisms behind these profound changes of membrane structure and organellar morphology, we have reconstituted the microautophagic membrane invagination in vitro. We have developed a system based on isolated yeast vacuoles and devised fast assays, which allow light microscopic tracing of the reaction, as well as convenient quantitation. We have used this system to characterize the requirements of microautophagic protein uptake into vacuoles.

### Materials and Methods

#### Yeast Strains and Genetic Manipulations

The strains used were: DBY7574 MATa ade2-1 lys2 his3 trpl1 ura3-53 cdc15-1::TRP ade3::HIS3::CMD1 (David Botstein, Stanford University, Stanford, CA); DBY7574-16 MATa pep4::LEU2 ade2-1 lys2 his3 trpl1 ura3-53 cdc15-1::TRP ade3::HIS3::CMD1, and DBY7574-19 MATa pho8::URA3 ade2-1 lys2 his3 trpl1 ura3-53 cdc15-1::TRP ade3::HIS3::CMD1. The following strains are derivatives of DBY7574 obtained by replacement of the indicated genes with a KAN resistance cassette: YTS1 (aut1Δ), YTS2 (aut2Δ), YTS3 (aut7Δ), YTS4 (apg10Δ), YTS5 (apg11Δ), and YTS6 (apg13Δ). The strains used for cytosol preparation were derived from K91-1A (MATa pho8::pAL134 pho13::pPH13 lys1; from Y. Kaneko, Osaka University, Osaka, Japan) by replacing the indicated gene with a KAN resistance cassette: YTS11 (aut1Δ), YTS12 (aut2Δ), YTS13 (aut7Δ), YTS14 (apg10Δ), YTS15 (apg11Δ), and YTS16 (apg13Δ). RSY271 (sec18-1; eve1-1) and the corresponding wild-type RSY249 were from Randy Schekman (University of California, Berkeley, CA). Pho8 and Pep4 were deleted by one-step gene replacement using plasmid pAR2 (J. Shaw, University of Utah, Salt Lake City, UT) and pTS17 (T. Stevens, University of Oregon, Eugene, OR), respectively.

Deletions of *Aut* and *Apg* genes were obtained as follows: using the oligonucleotides listed below, DNA fragments for the chromosomal replacement of the corresponding *Aut* or *Apg* genes with a *LoxP-KAN*–*LoxP* cassette were created by PCR on plasmid pUG6 (Guldener et al., 1996). The strains YTS1–YTS6 and YTS11–YTS16 were made by transforming the amplified single *LoxP-KAN*–*LoxP* cassettes into DBY7574 and K91-1A. The oligonucleotides used were: aut1ΔKAN1, 5′-GGT TCT AGA AGA ACG GAG ATA GGC CAC TAG CTA GAA GTT AGG AAC AAA GAA GTA CCA CCA GCT GAA AGC TCT GTA GCC-3′; aut1ΔKAN2, 5′-GTT TAT CCT GCT TTT TTA CCA CCT GCC TTG CTA ATG AGC TAC TGA AAG ATT CCC TAC CTT ATC CGT GGC AGC AGC TCT GTA GCC-3′; aut2ΔKAN1, 5′-ATG AAG TCT ACA TTT AAG TCT TAG CTT ATG ATC AAT ACG ACT TGC GAA GCC TCT GTA GCC-3′; aut2ΔKAN2, 5′-GTA TAT CTT TGC TTT TTA GTA ATG ATC AAT ACC TAG ATG ATG GGC TCT CTG GTA GCCT GC-3′; aut7ΔKAN1, 5′-ATG AAG TCT ACA TTT AAG TCT TAG CTT ATG ATC AAT ACC TAG ATG ATG GGC TCT CTG GTA GCCT GC-3′; and aut7ΔKAN2, 5′-GTA TAT CTT TGC TTT TTA GTA ATG ATC AAT ACC TAG ATG ATG GGC TCT CTG GTA GCCT GC-3′. The following strains were also transformed: YTS11-LoxP-KAN1, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′; and YTS16-LoxP-KAN2, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′; and YTS11-LoxP-KAN2, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′; and YTS16-LoxP-KAN2, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′; and YTS11-LoxP-KAN2, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′; and YTS16-LoxP-KAN2, 5′-GGT CAT TTG TAC TTA ATA AGA CAT GAT GTA GGA TCT GTA GCC-3′.
Growth of Cells

Yeast was precultured in YPD (1% yeast extract, 2% Bacto peptone, 2% glucose) for 6–8 h at 30°C and then diluted for logarithmic growth overnight (14–16 h, 30°C, 225 rpm) in 2-liter Erlenmeyer flasks with 1 liter of YPD medium. For starving cells, overnight cultures were harvested at an optical density at 600 nm (OD_{600}) of 2, centrifuged (4 min, 3,800 g, 4°C, JLA 10.500 rotor; Beckman), washed with sterile water, resuspended in 1 liter of SD (–N) (0.67% Difco yeast nitrogen base without amino acids and without ammonium sulfate, 2% glucose), and incubated (up to 3 h, 30°C, 225 rpm).

Cytosol Preparation

Cytosol was prepared from strain K91-1A or its derivatives. Overnight cultures were harvested at OD_{600} = 4.5, centrifuged (4 min, 3,800 g, 4°C, JLA 10.500 rotor), washed with sterile water, harvested again, resuspended in 1 liter of SD (–N) and incubated for 3 h at 30°C and 225 rpm. For preparation of cytosol from nonstarved cells, SD (–N) was replaced by YPD in this incubation. Cells were harvested and washed as described above, first with water, and then with one pellet volume of chilled lysis buffer (40 mM Pipes-KOH, pH 6.8, 0.5 mM MgCl₂, 150 mM KCl, 200 mM sorbitol, 0.1 mM DTT, 0.2 mM PMSE, 0.1 mM pefabloc SC, 0.5 mM pepstatin A, 0.1 μg/ml leupeptin, 50 μM O-phenanthroline). After centrifugation (5 min, 3,800 g, 4°C, JLA 10.500 rotor), the pellet was resuspended in a small volume of lysis buffer so that a thick slurry resulted. The suspension was frozen as little nuggets in liquid nitrogen and blended (six to eight times for 30 s) in a Waring blender filled with liquid nitrogen. Cells were thawed and the lysate was centrifuged (10 min, 12,000 g, 4°C, JA 25.500 rotor; Beckman). The supernatant was centrifuged (20 min, 125,000 g, 4°C, TLA45 rotor; Beckman). The fatty top fraction was discarded, and the clarified cytosol was recovered. The protein concentration of this cytosolic fraction was adjusted to 25–30 mg/ml with lysis buffer. Aliquots were frozen in liquid nitrogen and stored at −80°C.

Vacuole Preparation

Cells were grown in YPD overnight to OD_{600} = 2.5–3.0, harvested (4 min, 3,800 g, 4°C, JLA 10.500 rotor), and resuspended in 50 ml of 30 mM Tris-HCl, pH 8.9, with 10 mM DTT. Cells were incubated (5 min, 30°C), centrifuged as above, resuspended in 15 ml of spheroplasting buffer (50 mM potassium phosphate, pH 7.5, 600 mM sorbitol in YPD with 0.2% glucose and oxaoylycine [3,600 U/ml for DBY5734-16, 2,400 U/ml for DBY5734-19, and 3,000 U/ml for DBY5734]) and transferred into 30 ml Corex tubes. Cells were incubated (23 min, 30°C), reisolated (4°C, 1 min at 750 g and 1 min at 1,500 g: JA 25.500 rotor) and resuspended in 2.5 ml 15% Ficoll 400 in PS buffer. The spheroplasts were incubated (2 min at 0°C, then 80 s at 30°C), chilled again, transferred to an SW41 tube, and overlaid with 3 ml 8% Ficoll 400, 5 ml 4% Ficoll 400, and 1.5–2 ml 0% Ficoll 400 in PS buffer. After centrifugation (90 min, 154,000 g, 2°C, SW41 rotor), vacuoles were harvested from the 0–4% interphase.

In Vitro Autophagocytosis Assay

A standard reaction had a volume of 180 μl and was composed of: vacuoles (0.2 mg/ml from strain DBY5734, 105 mM KCl, 7 mM MgCl₂, 2 mM ATP, 10 mM DTT, 30 mM diacetyl phosphate, 140 μM creatine kinase, 1% proteinase inhibitor cocktail (PIC: 75 μM pefabloc SC, 75 ng/ml leupeptin, 37.5 μM O-phenanthroline, 375 μg/ml pepstatin A), 17 μg/ml firefly luciferase, 3 mg/ml cytosol from starved cells, 200 mM sorbitol, and 10 mM Pipes-KOH, pH 6.8. This mixture was incubated for 1 h at 27°C. A 70-μl sample was chilled on ice, diluted with 300 μl 150 mM KCl in PS buffer, centrifuged (6,800 g, 4 min, 2°C, fixed angle–table top centrifuge), washed twice with 300 μl 150 mM KCl in PS buffer, and resuspended in 60 μl 150 mM KCl in PS buffer. 10 μl was used to check for complete vacuole recovery by determining the protein concentration or, alternatively, the activity of endogenous Pho8p, as described previously (Haas et al., 1994). 40 μl of the suspension was treated with proteinase K (15 min, 0°C, 275 μg/ml, added from a 5-mg/ml stock in PS) to digest surface-bound luciferase. Digestion was terminated by adding an equal volume of freshly prepared PMSF solution (0.5 mM PMSF and 150 mM KCl in PS buffer). Luciferase activity was determined from a 25-μl aliquot in a luminometer using a luciferase kit (Berthold Detection Systems), according to the instructions of the supplier.

In Vitro Fusion

When both fusion and autophagocytosis were to be determined, the standard assay was performed as above, but a 1:1 mixture of vacuoles from a Δrep1 and a Δpho8 strain were used. The former carries proalkaline phosphatase (pro-Pho8p), and the latter only the appropriate maturation enzyme (Pep4p). Alkaline phosphatase activity is only generated upon mixing the contents of both vacuole types, resulting in fusion (Haas et al., 1994). Fusion was quantitated by using a 30-μl aliquot out of the standard assay (after incubation for 1 h at 27°C) and determining alkaline phosphatase activity, as described previously (Haas et al., 1994).

Purification of Fluorescein Dextran 500,000

10 mg fluorescein dextran 500,000 (Molecular Probes) was dissolved in 1 ml of PS buffer, concentrated, and rehydrated at least five times by ultrafiltration in Centricron 30 devices. The retentate was used for the experiments if low molecular weight fluorescein was no longer detectable in the filtrate by fluorescence spectrophotometry in a PerkinElmer LS 50-B spectrometer.

Microscopic Assay of Autophagocytosis In Vitro

The standard assay was modified by using 1 μM purified fluorescein dextran 500,000 (20-μM stock), instead of luciferase. Samples were taken, and the vacuoles were reisolated and washed, as described for the chemical assay. The vacuoles were resuspended in PS buffer with 50 μM FM4-64 to a protein concentration of ~1 mg/ml. The suspension was mixed with an equal volume of 0.4% Seaplaque agarose in PS (kept liquid at 35°C). 12 μl was transferred to a slide, chilled at 4°C for 5 min to immobilize the vacuoles, and analyzed by confocal fluorescence microscopy on a Leica TCS system, or by conventional fluorescence microscopy.

Results

Reconstitution of Vacular Inagination and Solute Uptake In Vitro

Microautophagocytosis occurs by direct invagination of lysosomal membranes, which is an “inverted” budding process into the lumen of the organelle. Soluble material is entrapped inside the invaginating vesicle and, upon scission of the membrane, becomes inaccessible to the surrounding medium. We have reconstituted this process in vitro using purified vacuoles from yeast. These organelles can be isolated in good purity and yield (Wiemken, 1975) and can be readily observed by light microscopy. We incubated isolated vacuoles in the presence of a cytosolic extract, an ATP-regenerating system, and appropriate salt and buffer conditions. To monitor vesicular uptake of soluble material from the medium, the incubation was performed in the presence of high molecular weight dextran coupled to FITC. Dye that had not been taken up was removed by reisolating the vacuoles. Uptake was analyzed by fluorescence microscopy. After 60 min of incubation almost a quarter of the vacuoles contained one or two punctate structures (Fig. 1, A and B) that were mobile in the lumen. Formation of the FITC-dextran–containing vesicles was ATP dependent. Upon longer incubation, these vesicles disappeared. The vacuolar lumen became homogeneously stained (Fig. 1 A), indicating that the vesicles had been lysed inside the vacuoles and their contents dispersed in the vacuolar lumen. Vacuoles were also stained with the membrane dye FM4-64 to assay for the presence of au-
trophagic tubes, the tubular membrane invaginations from which vesicles bud into the lumen of vacuoles in vivo (Müller et al., 2000, this issue). At the beginning of the incubation, <1% of the vacuoles had autophagic tubes (Fig. 1 C). After a lag phase of 15 min, the frequency of autophagic tubes increased to almost 10%. Beyond 60 min of incubation, the tubes disappeared again. The formation of autophagic tubes was ATP dependent, indicating that it was an active process. We conclude that the in vitro reaction reconstitutes the formation of autophagic tubes and the vesicular uptake of soluble material into vacuoles.

Using these microscopical assays, the concentrations of all components were optimized to yield maximal uptake activity (data not shown). The results of these optimizations were identical to those obtained with the chemical uptake assay described below. We developed this assay, which is based on the uptake of firefly luciferase, to provide a means for more convenient and sensitive quantitation of vesicle formation. Isolated vacuoles were incubated with cytosol and ATP in the presence of purified firefly luciferase. The organelles were reisolated, washed, and treated with increasing concentrations of proteinase K to digest residual luciferase potentially adhering to the surface of the vacuoles. Only luciferase that had been taken up into the vacuoles would resist this treatment. A strong signal of proteinase K–protected luciferase was recovered from samples incubated in the presence of ATP (Fig. 2 A). In the absence of ATP, no protease-protected luciferase cofractionated with the vacuoles. The protease-protected luciferase became accessible to proteinase K after lysis of the vacuoles by detergent (Fig. 2 A), or after rupturing the membranes by freeze–thawing (Fig. 2 B). This demonstrates that protease protection was due to sequestration in the vacuolar lumen. Therefore, the behavior of lu-
Luciferase matched that of FITC-dextran and could be used as a measure for membrane invagination and solute uptake from the medium.

The uptake reaction proceeded for \( \sim 60 \) min, with a lag in the first 30 min, and then entered a plateau at 90 min (Fig. 3 A). This correlates with the time course of formation and disappearance of autophagic tubes (see Fig. 1 C). Longer incubation in the absence of new uptake led to a significant reduction of the luciferase signal inside the vacuoles (Fig. 3 B). Luciferase incubated under the same conditions in reaction buffer remained stable. This is consistent with the fact that the invaginated vesicles became lysed inside the vacuoles (compare with Fig. 1 A). Luciferase released from the vesicles into the vacuolar lumen could then be degraded by vacuolar proteases. Luciferase uptake was temperature dependent, with a narrow optimum at 27°C (Fig. 4 A). The reaction had a strong requirement for cytosol, with the optima ranging from 2 to 4 mg/ml, depending on the individual batch of cytosol (Fig. 4 B). The reaction also required ATP and Mg\(^{2+}\), with optima at 2 and 7 mM, respectively (Fig. 4, C and D). The optima matched those found independently by the visual assay for vesicular uptake of FITC-dextran (data not shown), confirming that both assays measure the same process.
Vesicle Formation Depends on Starvation and on Gene Products Required for Autophagocytosis

Autophagocytosis in *S. cerevisiae* is a constitutive process that is induced under conditions of stress or starvation (Takeshige et al., 1992; Noda et al., 1995). Many mutations influencing this process have been isolated (Tsukada and Ohsumi, 1993; Thumm et al., 1994; Harding et al., 1995). These mutants affect macroautophagocytosis (Tsukada and Ohsumi, 1993; Thumm et al., 1994; Harding et al., 1995) and the formation of autophagic tubes (Müller et al., 2000, this issue) although they thoroughly blocked the homotypic fusion process necessary to pinch off the invaginated membrane and release vesicles into the vacuolar lumen. An in vitro uptake reaction was performed in the presence of affinity-purified antibodies to Sec17p, Sec18p, Vam3p, Vam7p, or Nny1p, or in the presence of Gdi1p, which extracts Ypt7p from the membrane. One aliquot of the reaction was used to quantify the uptake of luciferase into the vacuoles. A second aliquot of the same sample was used to monitor homotypic vacuolar fusion and control for the effect of the inhibitors. Vacuole fusion occurs under the conditions of the uptake reaction, albeit at only 20–30% of the efficiency under optimized conditions (Mayer et al., 1996). None of the inhibitors had a strong influence on the uptake of luciferase into the vacuolar lumen (Fig. 6 A) although they thoroughly blocked the homotypic fusion of vacuoles.

This finding was confirmed by results with a mutant carrying the temperature-sensitive sec18-1 allele. Vacuoles and cytosols from sec18-1 cells, which had been exposed to the restrictive temperature after their isolation from the cells, showed the same uptake activity as those from wild-type cells (Fig. 6 B). Successful inactivation of Sec18-1p was controlled by performing a visual fusion assay. Samples with sec18-1 vacuoles and cytosols had no fusion activity (not shown), whereas those with wild-type components showed many large fusion products (Haas et al., 1994). Like vacuole fusion (Haas et al., 1994), luciferase

Complemented by wild-type cytosol. Therefore, the Apg/Aut pathway does not obviously produce persistent changes on the vacuolar membrane. Its influences can be reset by the cytosol during the incubation. In summary, isolated vacuoles perform the uptake of macromolecular solutes into vesicular structures inside the organelle. The in vitro system reproduces the induction of vacuole invagination by starvation and the influence of the Apg/Aut proteins on this process, which was seen in vivo (Müller et al., 2000, this issue). Therefore, the in vitro reaction reconstitutes microautophagocytosis.

**Figure 4.** Requirements for in vitro uptake. Standard uptake reactions were performed (60 min) (A) at different temperatures, or in the presence of different concentrations of (B) cytosol, (C) ATP, or (D) MgCl2. Luciferase uptake was determined as described in the legend to Fig. 3 A. In B, titration curves for four independent cytosol preparations are shown. The maximal uptake signal of each titration curve was set at 100%. In D, the sample drawn as 0 mM MgCl2 contained 10 mM EDTA to chelate free magnesium in the reaction.
uptake also depended on guanine nucleotides. Vesicle formation was prevented by the GTP analogue GTPγS, which is poorly hydrolyzed, with an IC₅₀ of 100 μM (Fig. 7 A), indicating that GTP hydrolysis is required. It was also prevented by the addition of the proton uncoupler carbonyl cyanid-4-trifluoromethoxyphenylhydrazon (FCCP) or the V-ATPase inhibitor concanamycin A (Fig. 7 B). Therefore, the activity of the vacuolar ATPase and the proton gradient across the membrane are prerequisites for the uptake reaction.

We conclude that the elementary reaction of invagination and scission of a vesicle into the vacuolar lumen involves the action of a GTPase and requires a membrane potential. It occurs independent of homotypic vacuolar fusion, of biosynthetic transport to the vacuole, and of macroautophagocytosis, but it is subjected to control via starvation and components of the Apg/Aut pathway.
The smooth ultrastructural appearance of this zone of this issue). The autophagic tube is a narrow and long invagination that is filled with cytosol and pinches off vesicles into the lumen of the vacuole. The tip of the tube, such as by repressing macroautophagocytosis in rich media, or by macroautophagic membrane invagination and vesicle scission may be independent of conventional fusion machinery altogether, although it is topologically a homotypic fusion event between opposing membranes. It should be noted that in the in vitro system, an individual vacuole does not go through many rounds of invagination and vesicle formation. Most of the vacuoles bud only one or two vesicles into their lumen, which can be readily scored (see Fig. 1; our unpublished observations). Therefore, the source membrane for the invagination process is probably not limiting in the in vitro system. In contrast, the in vivo situation involves repetitive budding into the lumen of the organelle. A permanent supply of new vesicles to the vacuolar boundary membrane will be necessary to maintain microautophagic membrane invagination. If this supply is cut, such as by repressing macroautophagocytosis in rich media, or by apg/aut mutations, vacuolar invagination will be impaired by the limitation of source membrane.

Discussion

Transport of proteins into yeast vacuoles can occur via multiple pathways: vesicular transport from endosomes along the alkaline phosphatase, carboxypeptidase Y, or multivesicular body pathways (Wendland et al., 1998); the Cvt pathway, defined by aminopeptidase I (Scott and Klionsky, 1998); the Vid pathway, leading to degradation of fructose-1,6-bisphosphatase by inclusion into a novel type of vesicle (Huang and Chiang, 1997); direct Hsp70p-dependent translocation across the vacuolar membrane (Horst et al., 1999); and macroautophagocytosis (Take-shige et al., 1999). We have recently identified a microautophagic pathway in S. cerevisiae that operates via a specialized structure, the autophagic tube (Müller et al., 2000, this issue). The autophagic tube is a narrow and long invagination that is filled with cytosol and pinches off vesicles into the lumen of the vacuole. The tip of the tube, where vesicles are formed, is free of transmembrane particles. The smooth ultrastructural appearance of this zone of vesiculation matches that of autophagic bodies, which are also devoid of transmembrane particles (Müller et al., 2000, this issue; Baba et al., 1995).

In vivo analysis indicated that autophagic tubes are only prominent when macroautophagocytosis is induced, that is, under conditions of massive membrane influx towards the vacuolar membrane (Müller et al., 2000, this issue). Also, autophagic bodies, which can be formed both by micro and macroautophagocytosis, can only accumulate inside the vacuoles of protease-deficient cells if the macroautophagic pathway is active (Darsow et al., 1997). Therefore, it appears that the microautophagic pathway depends on the induction of macroautophagy. These observations suggest that a major function of autophagic tubes could be the maintenance of membrane homeostasis, that is, to invaginate excess membrane that is transferred into the vacuolar boundary membrane by its continuous fusion with the outer membrane of newly formed macroautophagosomes.

Microautophagic vacuole invagination in vitro was independent of the t-SNAREs, Vam3p and Vam7p, which are required for macroautophagocytosis, as well as the Cvt, alkaline phosphatase, carboxypeptidase Y, and multivesicular body pathways to the vacuole (Darsow et al., 1997; Sato et al., 1998; Srivastava and Jones, 1998; Abeliovich et al., 1999). Furthermore, microautophagocytosis was independent of Sec18p/NSF, Sec17p/α-SNAP, of the vacuolar Rab-like GTPase Ypt7p, and of the vacuolar v-SNARE Nyv1p. This indicates that the core process of microautophagic membrane invagination and vesicle scission may be independent of conventional fusion machinery altogether, although it is topologically a homotypic fusion event between opposing membranes. It should be noted that in the in vitro system, an individual vacuole does not go through many rounds of invagination and vesicle formation. Most of the vacuoles bud only one or two vesicles into their lumen, which can be readily scored (see Fig. 1; our unpublished observations). Therefore, the source membrane for the invagination process is probably not limiting in the in vitro system. In contrast, the in vivo situation involves repetitive budding into the lumen of the organelle. A permanent supply of new vesicles to the vacuolar boundary membrane will be necessary to maintain microautophagic membrane invagination. If this supply is cut, such as by repressing macroautophagocytosis in rich media, or by apg/aut mutations, vacuolar invagination will be impaired by the limitation of source membrane.

Shortage of source membrane is sufficient to explain the effects of the Apg/Aut components tested could regulate macro- and microautophagocytosis of soluble components and of peroxisomes in vivo (Müller et al., 2000, this issue). However, it does not readily explain why cytosols from Apg/Aut mutants support in vitro invagination of purified vacuoles significantly less than wild-type cytosols. This observation is consistent with a direct effect of the Apg/Aut components on the formation of autophagic tubes, on the formation of the vesicles at their tips, or on vesicle scission. Two different mechanisms are conceivable in which the same set of components could regulate macro- and microautophagocytosis of soluble and membrane-bound substrates at the same time. First, the Apg/Aut components tested could be part of a regulatory pathway that controls the induc-
tion of autophagy as a whole. It could regulate distinct machineries that are directly involved in the structural changes necessary for micro- and macroautophagy of soluble components or organelles. Second, the Apg/Aut components could themselves be part of an apparatus designed to bend the precursor membranes for autophagosomes and autophagic tubes and/or to sort their constituents. For both micro- and macroautophagy, similar membrane curvature must be induced in the future autophagic body membrane, that is, both reactions require topologically equivalent changes. In macroautophagy, large cisternae must be bent to encircle portions of cytosol or an organelle. In microautophagy, invagination of the vacuole requires the induction of an equivalent curvature in part of its boundary membrane. This could explain how similar components and mechanisms may be responsible for both reactions. Detailed morphological and biochemical analysis will be necessary to distinguish between these possibilities.

Ultrastructural analysis revealed that nascent microautophagic vesicles have a strikingly low density of intramembranous particles (Müller et al., 2000, this issue), that is, a very low content of large transmembrane proteins. This unusual structure correlates with the unexpected biochemical properties of vesicle formation, its independence of homotypic fusion factors. An attractive speculation is that scission may be lipid driven. Scission of synaptic vesicles undergoing endocytosis requires the concerted action of the GTPase dynamin and endophilin, a phosphatidic acid acyl transferase (Schmidt et al., 1999). The action of this enzyme converts an inverted cone-shaped lipid (lysophosphatidic acid) into a cone-shaped lipid (phosphatidic acid) in the cytoplasmic leaflet of the bilayer, a process that could support formation, as well as scission, of the nascent vesicle. It is unlikely that the same mechanism also acts on microautophagic vesicles, because microautophagic scission is topologically inverse. In endocytosis, enzymes involved in vesicle formation and pinching bind to the outer side of the vesicle and to its neck and induce a negative curvature of the cytoplasmic leaflet. However, in microautophagy they would have to bind to the inside of the nascent structure and locally increase the curvature of the cytoplasmic leaflet. Topologically, microautophagic vesicle formation is more related to the formation of multivesicular bodies, that is, the formation of internal endosomal vesicles. Those vesicles are strongly enriched in a specialized lipid, lysobisphosphatidic acid (Kobayashi et al., 1998). The smooth appearance of the fractured membrane of both vacuolar autophagic bodies and the zone of vesicle formation on autophagosomes suggests that both membranes are poor in transmembrane proteins and, therefore, rich in lipids. Since vesicle formation is also independent of the enzymes catalyzing homotypic vacuolar membrane fusion, lipids may play an important role in the budding of vesicles into the vacuolar lumen. The in vitro system described here will be an important tool to address this question.

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References

Abeliovich, H., T. Darsow, and S.D. Emr. 1999. Cytoplasm to vacuole trafficking of amnopenitase I requires a t-SNARE-Scclp complex composed of Tslg and Vps45. EMBO (Eur. Mol. Biol. Organ.) J. 18:6005–6016.

Müller, O., T. Sattler, M. Flötenmeyer, H. Schwarz, H. Plattner, and A. Mayer. 1999. New efficient gene disruption cassette for repeated use in budding yeast. Nucl. Acids Res. 24:2519–2524.

Haas, A., B. Conradt, and W. Wickner. 1994. G-protein ligands inhibit in vitro reactions of vacuolar inheritance. J. Cell Biol. 126:87–97.

Hurst, M., E.C. Knecht, and P.V. Schütz. 1999. Import into and degradation of cytosolic proteins by isolated yeast vacuoles. Mol. Biol. Cell. 10:2879–2889.

Huang, P.H., and H.L. Chiang. 1997. Identification of novel vesicles in the cytosol to vacuole protein degradation pathway. J. Cell Biol. 136:803–810.

Kim, J., V. Dalton, V. Eggerton, S.V. Scott, and D.J. Klionsky. 1999. Apg7p/Cvt2p is required for the cytoplasm-to-vacuole targeting, macroautophagy, and peroxisome degradation pathways. Mol. Biol. Cell. 10:1337–1351.

Kirisako, T., M. Baba, N. Ishihara, M. Kouchi, M. Ohsumi, T. Yoshimori, T. Noda, and Y. Ohsumi. 1999. Formation process of autophagosomes is traced with Apg8/Aut7p in yeast. J. Cell Biol. 147:435–446.

Knap, M., H.H. Schiffer, S. Rupp, and D.H. Wolf. 1993. Vacuolar/lysosomal proteolysate: proteases, substrates, mechanisms. Curr. Opin. Cell Biol. 5:990–996.

Kobayashi, T., E. Stang, K.S. Fang, P. Demoerloose, R.G. Parton, and J. Gruenberg. 1998. A lipid associated with the antiphospholipid syndrome regulates endosome structure and function. Nature. 392:193–197.

Lang, J., E. Schaeffeler, D. Bernreuther, M. Bredschneider, D.H. Wolf, and M. Thumm. 1998. Aut2p and Aut1p, two novel microtubule-associated proteins, are essential for delivery of autophagic vesicles to the vacuole. EMBO (Eur. Mol. Biol. Organ.) J. 17:3597–3607.

Lenk, S.E., W.A. Dunn, Jr., J.S. Trausch, A. Ciechanover, and A.L. Schwartz. 1992. Ubiquitin-activating enzyme E1 is associated with maturation of autophagic vacuoles. J. Cell Biol. 118:301–308.

Mayer, A., W. Wickner, and A. Haas. 1996. Sec1lp (NSF)-driven release of Sec1lp (t-SNARE) precedes docking and fusion of yeast vacuoles. Cell. 85:83–94.

Matsurau, A., M. Tsukada, Y. Wada, and Y. Ohsumi. 1997. Apg1p, a novel protein kinase required for the autophagic process in Saccharomyces cerevisiae. Gene. 192:245–250.

Mizushima, N., T. Noda, T. Yoshimori, Y. Tanaka, T. Ishii, M.D. George, D.J. Klionsky, M. Ohsumi, and Y. Ohsumi. 1998. A protein conjugation system essential for autophagy. Nature. 395:395–398.

Mizushima, N., T. Noda, and Y. Ohsumi. 1999. Apg16p is required for the function of the Apg12p–Apg5p conjugate in the yeast autophagy pathway. EMBO (Eur. Mol. Biol. Organ.) J. 18:3888–3896.

Mortimore, G.E., and M. Kadowaki. 1994. Autophagy: its mechanism and regulation. In Cellular Proteolytic Systems, A.L. Ciechanover and A.L. Schwartz, editors. Wiley & Liss, New York. 65–87.

Müller, O., T. Sattler, M. Flötenmeyer, H. Schwarz, H. Plattner, and A. Mayer. 2000. Autophagic tubes: vacuolar invaginations involved in lateral membrane sorting and inverse vesicle budding. J. Cell Biol. 151:519–528.

Nichols, B.J., C. Ungermann, H.H.B. Pelham, W.T. Wickner, and A. Haas. 1997. Homotypic vacuolar fusion mediated by t- and v-SNAREs. Nature. 387:199–202.
Takeshige, K., M. Baba, S. Tsuboi, T. Noda, and Y. Ohsumi. 1992. Autophagy in yeast demonstrated with proteinase-deficient mutants and conditions for its induction. J. Cell Biol. 119:301–311.

Thumm, M., R. Egner, B. Koch, M. Schlumpberger, M. Straub, M. Veenhuis, and D.H. Wolf. 1994. Isolation of autophagocytosis mutants of Saccharomyces cerevisiae. FEBS Lett. 349:275–280.

Titorenko, V.I., I. Keizer, W. Harder, and M. Veenhuis. 1995. Isolation and characterization of mutants impaired in the selective degradation of peroxisomes in the yeast Hansenula polymorpha. J. Bacteriol. 177:357–363.

Tsukada, M., and Y. Ohsumi. 1993. Isolation and characterization of autophagy-defective mutants of Saccharomyces cerevisiae. FEBS Lett. 333:168–174.

Tuttle, D.L., and W.A. Dunn, Jr. 1995. Divergent modes of autophagy in the methylotrophic yeast Pichia pastoris. J. Cell Sci. 108:25–35.

Tuttle, D.L., A.S. Lewin, and W.A. Dunn, Jr. 1993. Selective autophagy of peroxisomes in methylotrophic yeasts. Eur. J. Biochem. 208:283–290.

Ungermann, C., and W. Wickner. 1998. Vam7p, a vacuolar SNAP-25 homolog, is required for SNARE complex integrity and vacuole docking and fusion. EMBO (Eur. Mol. Biol. Organ.) J. 17:3269–3276.

Veenhuis, M., A. Douma, W. Harder, and M. Osumi. 1983. Degradation and turnover of peroxisomes in the yeast Hansenula polymorpha induced by selective inactivation of peroxisomal enzymes. Arch. Microbiol. 134:193–203.

Wada, Y., Y. Ohsumi, and Y. Anraku. 1992. Genes for directing vacuolar membrane and autophagosomal membranes in rat hepatocytes. J. Histochem. Cytochem. 40:699–709.

Wendland, B., S.D. Emr, and H. Riezman. 1998. Protein traffic in the yeast endocytic and vacuolar protein sorting pathways. Curr. Opin. Cell Biol. 10:513–522.

Wiepkens, A.D., and W. van Doorn. 1978. Glucose-induced autophagy of peroxisomes in the yeast Hansenula polymorpha. Biochem. J. 173:39–45.

Yuan, W., P.E. Stromhaug, and W.A. Dunn. 1999. Glucose-induced autophagy of peroxisomes in Pichia pastoris requires a unique E1-like protein. Mol. Biol. Cell. 10:1553–1566.