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Atg18 phosphoregulation controls organellar dynamics by modulating its phosphoinositide-binding activity

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Introduction

In eukaryotes, various biological processes are regulated by phosphoinositide (PI) signaling (Di Paolo and De Camilli, 2006). PIs contain inositol phosphate head groups that are specifically recognized by PI-binding domains on partner proteins, e.g., FYVE, PX, PH, and GRAM domains (Lemmon, 2008). However, the posttranslational regulation of lipid-binding activity in these domains has been scarcely investigated at this point.

PROPPIN family members, including Atg18, Atg21, and Hsv2, are proteins with a lipid-binding domain comprising a seven β-propeller motif that binds to phosphatidylinositol 3-monophosphate (PI(3)P) and/or phosphatidylinositol 3,5-bisphosphate (PI(3,5)P2; Dove et al., 2009). Atg18 is one of the autophagy-related molecules responsible for autophagic processes, and is conserved from yeast to higher organisms, including the human proteins WIPI1–WIPI4 (Mizushima et al., 2011). In autophagy, nutrient starvation leads to bulk degradation of intracellular components in a sequestering vesicle called the autophagosome. This remodeling is mediated by the TORC1 complex and PI(3)P signaling (Mizushima et al., 2011). Pexophagy, one type of selective autophagy involving peroxisomes, is classified into two distinct pathways: micropexophagy and macropexophagy (Manjithaya et al., 2010; Oku and Sakai, 2010). In the methylotrophic yeast, Pichia pastoris, the carbon source used for pexophagy induction determines which pathway is used: glucose and ethanol adaptation induce micropexophagy and macropexophagy, respectively. During micropexophagy, a vacuole engulfs the peroxisomal cluster by repeating septation, resulting in the formation of a vacuolar sequestering membrane (VSM). Simultaneously, a micropexophagy-specific membrane apparatus (MIPA) is necessary to complete the incorporation of peroxisomes into the vacuole. Similar to pexophagosome formation, Atg8 is localized to MIPA, and its assembly requires vacuolar membrane, Atg18 was rephosphorylated, and the vacuoles fused and formed a single rounded structure. Vacular dynamics were regulated according to osmotic changes, oxidative stresses, and nutrient conditions inducing micropexophagy via modulation of Atg18 phosphorylation. This study reveals how the phosphoinositide-binding activity of the PROPPIN family protein Atg18 is regulated at the membrane association domain and highlights the importance of such phosphoregulation in coordinated intracellular reorganization.
the conventional Atg proteins, Atg11 and Atg30 (Farré et al., 2008; Oku and Sakai, 2010). *P. pastoris* Atg18 (PpAtg18) is necessary for VSM formation (Guan et al., 2001); however, its involvement in MIPA formation has not yet been reported. In contrast, in the budding yeast, *Saccharomyces cerevisiae*, Atg18 is required for the regulation of vacuolar shape via its PI(3,5)P$_2$-binding activity. Vacuolar fission is induced under certain environmental stresses, such as hyperosmotic pressure and high glucose concentrations. PI(3,5)P$_2$, synthesized by certain environmental stresses, such as hyperosmotic pressure and high glucose concentrations. PI(3,5)P$_2$, synthesized by the PI(3)P 5-kinase, Fab1, is required for this process (Dove et al., 1997, 2009). By regulating vacuolar shape, Atg18 is suggested to function as an effector of PI(3,5)P$_2$ signaling (Dove et al., 2004).

Collectively, these studies suggested that Atg18 has dual physiological functions, i.e., autophagy regulation mediated by PI(3)P signaling, and control of vacuolar shape via PI(3,5)P$_2$ signaling. However, the molecular mechanisms underlying Atg18’s role in these signaling pathways were not known. In this paper, two phosphorylation regions were found in the loops in the β-propeller structures. Based on the recently revealed molecular structure of the PROPPIN family member Hsv2, from the yeast *Kluyveromyces lactis* (KIHsv2), the two phosphorylation sites were thought to play critical roles in the protein’s affinity for phosphoinositides (Baskaran et al., 2012; Krick et al., 2012; Watanabe et al., 2012). However, phosphorylation of PpAtg18 was found to decrease the PI-binding activity of PpAtg18. Remarkably, the level of PpAtg18 phosphorylation in the cells correlated with the intracellular localization of PpAtg18. Further analyses revealed that the phosphorylation of PpAtg18 governed vacuolar shape during adaption to various environmental stresses, including situations that induced micropexophagy.

**Results**

**PpAtg18 is phosphorylated in *P. pastoris***

We overproduced GST-tagged PpAtg18 under the control of the constitutive GAPDH promoter in the original host *P. pastoris*. The purified fraction from a Glutathione Sepharose 4B (GS 4B) column exhibited two distinct bands corresponding to the molecular size of the GST-PpAtg18 protein as determined by Coomassie brilliant blue staining of SDS-PAGE (Fig. 1 A). Because phosphatase treatment abolished the upper band, leaving a single band on the gel, we concluded that the purified GST-PpAtg18 protein contained both phosphorylated and nonphosphorylated forms (Fig. 1 A and B).

To exclude the possibility that phosphorylation of PpAtg18 was caused by overexpression of the protein, we expressed functional PpAtg18 tagged with five repeats of the Flag peptide (5xFlag) under the control of the original *PpATG18* promoter. Again, we observed two bands via immunoblot analysis of cell-free extracts from *P. pastoris* (grown in glucose). Phosphatase treatment of the cell-free extract yielded a single band that was cross-reactive with the Flag tag (Fig. 1 C). These experiments indicated that PpAtg18 was present in both phosphorylated and nonphosphorylated forms in *P. pastoris* under physiological conditions.

**Phosphorylation of PpAtg18 inhibits PI(3,5)P$_2$-binding activity**

Atg18 proteins were reported to bind to PI(3)P and PI(3,5)P$_2$ (Dove et al., 2004). Purified GST-PpAtg18 protein containing both phosphorylated and nonphosphorylated isoforms was subjected to a variety of lipid binding tests, including the phosphatidylinositol phosphate (PIP) strip assay, the PIP array assay, a liposome pull-down assay, and surface-plasmon resonance analysis to determine the specificity of lipid binding. The PIP strip, PIP array, and liposome pull-down assays identified significant PI(3,5)P$_2$-binding and weaker PI(3)P-binding activities (Fig. 1, D–F). However, we could not detect significant PI(3)P binding activity using Biacore surface-plasmon resonance analysis (Fig. S2 B). These data were in accord with previous results obtained using Atg18 from *S. cerevisiae* (Dove et al., 2004).

Next, we treated the purified PpAtg18 fraction (including both phosphorylated and nonphosphorylated forms) with phosphatase (Fig. 1 A, lane 2), and PI-binding activities were compared between phosphatase-treated and nontreated samples. As determined using the PIP strip assay, phosphatase treatment dramatically increased PpAtg18 binding activity toward PI(3,5)P$_2$ as compared with PI(3)P (Fig. 1 D). This increase in affinity to PI(3,5)P$_2$ after dephosphorylation was confirmed using the PIP array and the liposome-binding assay (Fig. 1, E and G). These experiments suggested that the affinity of PpAtg18 toward PI(3,5)P$_2$ is reduced by phosphorylation. However, the affinity of PpAtg18 toward PI(3)P was too weak to draw comparisons between phosphatase-treated and nontreated samples.

**PpAtg18 has two distinct phosphorylation regions in the loops of blades 6 and 7 that affect PI binding activity**

MS analysis of purified PpAtg18 protein identified two phosphorylated peptides, one with two putative phosphorylation sites. Further MS/MS analyses identified these as 387-SSTTS-391 and 492-SSTS-495 (Fig. S1, A and B). Next, we compared the sequence around the phosphorylated sites with those of other PROPPIN family members by BLAST analysis. The crystal structure of KIHsv2 revealed consensus sequences for each blade (Fig. 2 A; Baskaran et al., 2012; Krick et al., 2012). The phosphorylated region in blade 6 was found in a loop that is conserved in all PROPPIN family proteins, including WIPI1 and KIHsv2 (Fig. 2, A and B). This loop is thought to be associated with the membrane via a hydrophobic bond (Baskaran et al., 2012); therefore, phosphorylation in this loop is speculated to inhibit lipid binding. However, the region close to the C terminus (between β1 and β2 of blade 7) was conserved in the methyloctrophic yeast *Hansenula polymorpha*, and partially conserved in *S. cerevisiae*, but was not retained in higher organisms.

To confirm the negative effect of phosphorylation in vivo, the putative phosphorylated residues of Ser and Thr were mutated to Ala and expressed in *P. pastoris* (Table 1). Cell-free extracts were subjected to normal and Phos-tag SDS-PAGE. Both PpAtg18 SA and AS mutations caused increased mobility of bands in both normal and Phos-tag SDS-PAGE; the PpAtg18 AA mutant produced the fastest migrating band among the tested
subjected to SDS-PAGE. The purified PpAtg18 DD protein migrated slower and the PpAtg18 AA protein faster, the former and the latter nearly corresponding to the mobility of the phosphorylated and dephosphorylated forms, respectively (Fig. S2 A). Next, these samples were subjected to lipid binding experiments (Fig. 1, D–G; and Fig. S2 B). Both the PIP array and the liposome pull-down assay revealed that the phosphorylation-defective mutant had a higher affinity to PI(3,5)P2, whereas the phosphorylation mimic PpAtg18 DD protein showed an affinity...

Figure 1. Phosphorylation of PpAtg18 modulates PI(3,5)P2-binding activity. (A) Evaluation of purified GST-PpAtg18 from P. pastoris by SDS-PAGE and Coomassie brilliant blue staining. Lane 1: Purified GST-PpAtg18 was eluted from a GS 4B column using the reduced form of glutathione. Lane 2: Purified GST-PpAtg18 was treated with λ-phosphatase on the GS 4B column. This sample was used for the lipid binding assay. (B) Phosphatase treatment of purified PpAtg18 from P. pastoris. Heat, λ-phosphatase was inactivated at 65°C for 1 h. (C) Phosphatase treatment of cell-free extracts from the strain expressing PpAtg18-5×Flag under the original PpATG18 promoter. The cell lysate was treated with λ-phosphatase at 37°C for 1 h. (D) PIP strip analysis. Membranes were incubated with 0.5 µg/ml protein and detected using the Light Capture II system (ATTO). GST-PpAtg18Wt and GST-Atg18Wt with λ-phosphatase were acquired simultaneously to ensure equal exposure times. (E) PIP array analysis. Membranes were incubated with 1.0 µg/ml protein and detected simultaneously to ensure equal exposure times. (F) Liposome pull-down assay. Purified GSTPpAtg18 (1.0 µg) was incubated with each liposome preparation and centrifuged at 16,000 g for 20 min. The pellets were washed twice, suspended in sample buffer, and analyzed by immunoblotting to detect GST-PpAtg18. No lipid, no liposomes; No PI, liposomes without PIs; PI(3)P, liposome containing PI(3)P; PI(3,5)P2, liposome containing PI(3,5)P2. (G) Liposome pull-down with dephosphorylated PpAtg18. Each protein (1.0 µg) was incubated with liposomes containing P[3,5]P2, and analyzed by immunoblot to detect GST-PpAtg18. The bands were analyzed by densitometry, and band intensities were normalized to samples not treated with phosphatase. Error bars indicate mean ± SEM. **, P < 0.05.

mutants (Fig. 3, A and B). Therefore, we concluded that both the Ser and Thr residues at positions 387–391 and 491–495 were phosphorylated in P. pastoris. Similarly, we also constructed PpAtg18 mutants that mimicked phosphorylation by replacing Ser or Thr residues within these regions with Asp (Table 1).

We purified the phosphorylation-defective PpAtg18 AA protein and the phosphorylation mimic PpAtg18 DD protein as GST fusion proteins, along with the wild-type PpAtg18 protein from P. pastoris cells. Molar equivalents of each sample were subjected to SDS-PAGE. The purified PpAtg18 DD protein migrated slower and the PpAtg18 AA protein faster, the former and the latter nearly corresponding to the mobility of the phosphorylated and dephosphorylated forms, respectively (Fig. S2 A). Next, these samples were subjected to lipid binding experiments (Fig. 1, D–G; and Fig. S2 B). Both the PIP array and the liposome pull-down assay revealed that the phosphorylation-defective mutant had a higher affinity to P[3,5]P2, whereas the phosphorylation mimic PpAtg18 DD protein showed an affinity...
after SDS-PAGE with or without Phos-tag (Fig. S2 D). As a result, we identified four amino acid residues critical for Atg18 phosphorylation: two in the loop of blade 6, S388 and S391; and two in the loop of the C terminus, S492 and S495 (Fig. 2). Phosphorylation of PROPPIN, as observed for PpAtg18, may mediate the physiological signal specific to each PROPPIN by modulating its PI-binding activity.

Phosphorylation of PpAtg18 correlates with vacuolar shape under various environmental stresses

Atg18 was previously shown to regulate vacuolar shape during adaptation to various environmental stresses; e.g., nutrient and osmotic change. Furthermore, this regulation was mediated by the PI-binding activity of Atg18 (Dove et al., 2004; Krick et al., 2006; Efe et al., 2007; Obara et al., 2008). In S. cerevisiae, hypo-osmotic stress and carbon-source limitation lead to homotypic vacuolar fusion. In contrast, hyperosmotic stress and high glucose concentrations lead to vacuolar fission (Dove et al., 2009; Li and Kane, 2009).

The physiological significance of PpAtg18 phosphorylation was sought. Cells expressing PpAtg18-5×Flag were adapted to various environmental conditions, and the phosphorylation level of PpAtg18-5×Flag was assessed (Fig. 4 A). As similar to the wild-type protein (Fig. 3, C and D). Similarly, we further assessed PI(3,5)P2 binding activity of PpAtg18 SA and PpAtg18 AS proteins in comparison with the PpAtg18 wild-type protein to see dephosphorylation in which loop structure was responsible to the regulation of the affinity (Fig. S2 C). Because both mutations increased the affinity, we concluded that dephosphorylation of either phosphorylated region of PpAtg18 increased the affinity toward PI(3,5)P2.

We mutated each of putative phosphorylated residues of serine/threonine residues to alanine, and examined the phosphorylation levels of the variant forms by the immunoblot analysis after SDS-PAGE with or without Phos-tag (Fig. S2 D). As a result, we identified four amino acid residues critical for Atg18 phosphorylation: two in the loop of blade 6, S388 and S391; and two in the loop of the C terminus, S492 and S495 (Fig. 2). Phosphorylation of PROPPIN, as observed for PpAtg18, may mediate the physiological signal specific to each PROPPIN by modulating its PI-binding activity.

Table 1. Designation of phosphorylation-mutants

<table>
<thead>
<tr>
<th>Name</th>
<th>Phosphorylated regions in:</th>
<th>Blade 6</th>
<th>Blade 7</th>
</tr>
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<tbody>
<tr>
<td>Wt</td>
<td>383 PRRSSSTTLGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>AS</td>
<td>383 PRVVAAALGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>SA</td>
<td>383 PRRSSSTTLGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>AA</td>
<td>383 PRVVAAALGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>DS</td>
<td>383 PRVDDDDDLGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>SD</td>
<td>383 PRRSSSTTLGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
<tr>
<td>DD</td>
<td>383 PRVDDDDDLGSGQSQES</td>
<td>384 407</td>
<td>407 427</td>
</tr>
</tbody>
</table>

Wt, wild type.
cells were adapted to hyperosmotic stress (0.9 M NaCl) or high-glucose conditions (20% glucose), vacuolar fission occurred (Fig. 4 B), and the dephosphorylated form of PpAtg18 increased. When cells were subjected to hypo-osmotic stress (H2O) or glucose starvation (−glucose), vacuolar fusion occurred, and the phosphorylated form of PpAtg18 increased. The change from hyperosmotic stress to glucose starvation caused a drastic increase in phosphorylated PpAtg18, and adaptation in the opposite direction caused a decrease of phosphorylated PpAtg18 (Fig. 4 C). However, the phosphorylation levels did not change in a control experiment (transfer from synthetic dextrose [SD] medium to SD medium; Fig. 4 A).

Next, we observed vacuolar morphology and the localization of PpAtg18 under these various environmental conditions. PpAtg18 tagged with the fluorescent protein YFP was expressed under its own promoter. Under vacuolar-fission conditions, the vacuole became fragmented, and PpAtg18-YFP was recruited to the vacuolar membrane, especially to the sites of vertex domains (Fig. 4 B). Alternatively, under vacuolar fusion conditions, PpAtg18-YFP was released from vacuolar membrane, and diffused throughout the cytosol. In the control experiment, PpAtg18 was weakly detected on the vacuolar membrane, and the level of vacuolar fusion (counted as the vacuolar number) seemed to be intermediate between that observed under hyperosmotic and hypo-osmotic stress (Fig. 4 B). We also tested other stress conditions that induce changes in vacuolar morphology (e.g., heat shock and ethanol stress), and observed that PpAtg18 phosphorylation levels increased under vacuolar fusion conditions and decreased under fission conditions.

Interestingly, PpAtg18 was highly phosphorylated when cells were exposed to oxidative stresses that induced vacuolar fusion: the oxidants, H2O2, tBOOH, and diamide (Fig. 4, D and E). We further tested other environmental stresses (reducing agents, exogenous calcium levels) that did not change either PpAtg18 phosphorylation levels or vacuolar shape. These results were consistent with a correlation between PpAtg18 phosphorylation levels and vacuolar shape in P. pastoris.

**Dephosphorylation of PpAtg18 regulates vacuolar fission by increasing PI(3,5)P2-binding activity**

Fab1 synthesizes PI(3,5)P2 from PI(3)P. In Ppfab1Δ, PpAtg18-YFP, which is present on the vacuolar membrane in the wild-type strain when cells are cultured in glucose media or under hyperosmotic conditions, seemed to be released to the cytosol (Fig. 5 A). The finding that PpAtg18-YFP was recruited to the vacuolar membrane in a PI(3,5)P2-dependent manner led us to test whether PpAtg18 phosphorylation-defective mutants were localized differently as compared with wild-type protein. The phosphorylation-defective mutant PpAtg18AA-YFP showed stronger fluorescence at the vacuolar membrane in hypo-osmotic (SD + 0.9 M NaCl) or control (SD) conditions. In contrast, localization of the PpAtg18DD-YFP mutant, which retained PI(3,5)P2-binding activity, was similar to the wild-type PpAtg18 (Fig. 5 B). However, both PpAtg18 AA and DD mutant proteins were released into the cytosol under vacuolar fusion conditions (SD – glucose); the cells in these experiments displayed a single spherical vacuole. To investigate whether PpAtg18AA localized to the vacuolar membrane affected vacuolar shape, we next assessed the level of vacuolar fission as the number of vacuolar compartments per cell. As a result, PpAtg18AA displayed more vacuolar fission than the wild-type and PpAtg18DD when cultured both in SD media and under hyperosmotic conditions (Table 2). Together with the results from the in vitro PI(3,5)P2-binding experiment (Fig. 1 and Fig. 3), we concluded that dephosphorylation of PpAtg18 positively regulates vacuolar fission by increasing binding to PI(3,5)P2.
Figure 4. *PpAtg18* phosphorylation correlates with the vacuolar dynamics observed under a variety of environmental conditions. (A) *PpAtg18*-5×Flag from cells grown under various conditions was detected by immunoblotting with the anti-FLAG antibody. Cells were shifted from SD (0 min) to SD medium containing 0.9 M NaCl or 20% glucose, to water, to YNB medium without amino acids (−glucose), to methanol-containing medium, or (as a control) to SD medium for the indicated times (10, 20, 40, or 60 min). (B) Fluorescence microscopy analysis of *PpAtg18* localization and vacuolar morphology. Growing conditions were the same as in Fig. 4 A. *PpAtg18*-tagged YFP was expressed under the original promoter and vacuoles were stained with FM 4-64. ([C] Pulse-chase experiment. Cells were shifted from SD medium to either SD + 0.9 M NaCl or SD-glucose, and then shifted again into the opposite medium (see arrows). Analysis was performed by immunoblotting. (D) Oxidative stress induces *PpAtg18* phosphorylation. t-BOOH, t-butylhydroperoxide. Analysis was performed by immunoblotting. (E) Morphological analysis and morphometric analysis of vacuoles in cells with or without oxidants. Cells
increased its vacuolar association, this result may represent the fact that phosphorylated PpAtg18 is mainly localized to the vacuolar membrane.

**Phosphoregulation of PpAtg18 during adaptation to methanol medium, and during micro- and macrophagocytosis**

Regulation of vacuolar shape. Previously, we reported that vacuoles fused to form a single spherical vacuole in a wild-type strain, the level of PpAtg18 phosphorylation increased under vacuolar fusion conditions, and decreased under vacuolar fission conditions. Interestingly, the level of PpAtg18 phosphorylation was lower in the **Ppfab1Δ** strain and the PpAtg18 FTTG mutant (Dove et al., 2004; Krick et al., 2006; Obara et al., 2008) as compared with the wild-type protein (Fig. 5 C). In these mutants, however, the level of dephosphorylation was comparable to that observed in wild type (Fig. 5 C). Because the PI(3,5)P$_2$-binding activity of PpAtg18 was shifted to SD (control) or to SD with 1 mM H$_2$O$_2$, 15 mM t-BOOH, or 4 mM diamide. Vacuoles were stained with FM 4-64. Error bars indicate mean ± SEM. *, $P < 0.01$ (compared with SD [control]). Bars, 2 µm.
Collectively, these data suggest that VSM formation during in PI(3,5)P2-binding activity, showed a drastic decrease in VSM PpVac14 observed in the MIPA formation. In contrast, neither VSM nor MIPA were vacuoles and YFP-PpAtg8 formed a cuplike structure representing type strain, VSMs labeled with FM 4-64 budded from maternal kinase complex. Both kinase component mutants, published data). Vac14 and Fab1 are components of the PI(3)P 5-kinase complex. During this process, the formation of autophagic membranes during mechanisms underlying PpAtg18 regulation of vacuolar dynam- coordinately regulated. Therefore, we wanted to understand the processes, i.e., VSM formation and MIPA formation, should be required for the early stage of micropexophagy, we speculate that PpAtg18 is involved in the formation of VSM concomitant with vacuolar fission (Guan et al., 2001). Also, the two distinct processes, i.e., VSM formation and MIPA formation, should be coordinately regulated. Therefore, we wanted to understand the mechanisms underlying PpAtg18 regulation of vacuolar dynamics, including the formation of autophagic membranes during methanol adaptation and pexophagic conditions.

First, we examined MIPA and VSM formation during micropexophagy in the Ppatg18Δ strain (Fig. 6 B). In the wild-type strain, VSMs labeled with FM 4-64 budded from maternal vacuoles and YFP-PpAtg8 formed a cuplike structure representing MIPA formation. In contrast, neither VSM nor MIPA were observed in the Ppatg18Δ strain. In the Pp fab1Δ cells, MIPA-like structures were formed, but VSM formation was not observed. We performed morphometric analysis of VSM formation by counting the number of vacuoles during micropexophagy (unpublished data). Vac14 and Fab1 are components of the PI(3)P 5-kinase complex. Both kinase component mutants, PpFab1Δ and PpVac14Δ, along with the PpAtg18 mutant that was impaired in PI(3,5)P2-binding activity, showed a drastic decrease in VSM formation (unpublished data).

Next, we investigated the intracellular localization of PpAtg18-YFP. In cells grown in methanol medium (at both late log phase and early lag phase) or under macropexophagic conditions, PpAtg18-YFP localized to the cytosol (Fig. 6 C). In contrast, during micropexophagy, PpAtg18-YFP was recruited to the vacuolar membrane. This vacuolar recruitment was lost in the Pp fab1Δ strain (Fig. 6 C), which indicates that PpAtg18 was recruited to the vacuolar membrane in a PI(3,5)P2-dependent manner during micropexophagy. Collectively, these data suggest that VSM formation during micropexophagy requires the PI(3,5)P2-dependent recruitment of PpAtg18 to the vacuolar membrane.

Then, we examined the phosphorylation levels of PpAtg18 under various conditions. At early phases of methanol adaptation (1–2 h), the phosphorylated form of PpAtg18 became dominant concomitant with vacuolar fusion, and PpAtg18-YFP diffused into the cytosol (Fig. 4, A and B). After 8–12 h of methanol adaptation, we could detect both phosphorylated and dephosphorylated forms of PpAtg18 and observed huge peroxisomes (Fig. 6 D). Therefore, although PpAtg18 was phosphorylated upon vacuolar fusion at early log phase, part of the phosphorylated PpAtg18 became dephosphorylated during cellular proliferation. This may reflect the fact that vacuolar inheritance from mother to daughter cells includes both vacuolar fission and fusion, which may be controlled by phosphoregulation of PpAtg18 (Weisman, 2006; Tamura et al., 2010).

Cells cultured in methanol at late-log phase were induced to undergo micro- and macroexophagy by transfer to glucose and ethanol medium, respectively. Then, PpAtg18 phosphorylation and localization (using PpAtg18-YFP) were observed. When cells were shifted to ethanol medium to induce macropexophagy, the phosphorylated form became dominant and the dephosphorylated form decreased significantly, suggesting that the phosphorylated form of PpAtg18 functioned in pexophagosome formation (Fig. 6 D). In contrast, both the phosphorylated and dephosphorylated forms of PpAtg18 were observed during micropexophagy, which included both MIPA formation and vacuolar fusion for VSM formation (Fig. 6 D).

Regulation of autophagy. As shown in Fig. 6 B, PpAtg18 was necessary for MIPA formation in micropexophagy. In nutrient starvation–induced autophagy, PpAtg18 was reported to be recruited to the preautophagosomal structure (PAS), represented by PpAtg8 dots, and this localization required Atg2 and PpAtg8 (Suzuki et al., 2007). To visualize such colocalization in P. pastoris, we expressed CFP-tagged PpAtg18 under the PpACT1 promoter, and coexpressed YFP-PpAtg8 under its original promoter. Fluorescence microscopy revealed that CFP-PpAtg18 was colocalized with the PAS (Fig. S3 A) during micropexophagy. However, we also observed that PpAtg26, which is necessary for pexophagy, was dispensable for the recruitment of PpAtg18 to the PAS.
Figure 6. Phosphorylation of PpAtg18 regulates VSM formation in micropexophagy. [A] Intracellular structures observed during micropexophagy and macroexophagy. MIPA, micropexophagy-specific membrane apparatus; VSM, vacuolar sequestering membrane. (B) Fluorescence microscopy analysis during micropexophagy. Cells were shifted from methanol medium to glucose medium for 30–60 min to induce micropexophagy. Vacuoles were stained with FM 4-64, and MIPA was visualized by YFP-tagged PpAtg8 expressed under the PpATG8 promoter. (C) Intracellular localization of PpAtg18. YFP C-terminally tagged with PpAtg18 was expressed in Ppatg18 under the PpATG18 promoter. Cells were shifted from methanol medium to ethanol or glucose medium for 30–60 min to induce macropexophagy or micropexophagy, respectively. (D) Immunoblot detection of PpAtg18-5×Flag. PpAtg18-5×Flag was expressed in Ppatg18 under the PpATG18 promoter. Cells were shifted from methanol medium to ethanol or glucose medium to induce macropexophagy or micropexophagy, respectively. This blot was incubated with the anti-FLAG antibody. (E) Intracellular localization of PpAtg18 AA and DD during micropexophagy. YFP-tagged PpAtg18 mutants were expressed under the PpATG18 promoter. Cells were shifted from methanol medium to glucose medium for 60 min to induce micropexophagy. Bars, 2 µm. (F) PpPex12 degradation assay to assess micropexophagy and macropexophagy activity in PpAtg18 mutants. Strains were grown in methanol medium and then shifted to glucose or ethanol medium to induce micropexophagy or macropexophagy, respectively. Cell-free extracts were prepared and analyzed by immunoblotting with anti-PpPex12 or anti-Pgk1 antibodies. Optical density measurements showed no significant difference in growth between these mutants.
VSM formation was normal in the during micropexophagy and the phosphorylation of PpAtg18. Although the starvation, the phosphorylation levels of PpAtg18 (Fig. S3, suggested that mutations in the phosphorylated regions did not influ-
ence nitrogen starvation–induced autophagy using the YFP-
macropexophagy by regulating the intracellular distribution 
PpAtg18 was required for optimum activity of both micro- and 
process. Although PpPex12 degradation was retarded in the PpAtg18 AA mutant under micropexophagic conditions as compared with the wild-type and the PpAtg18 DD mutant, even though VSM formation was enhanced (Fig. 6, E and F). However, PpPex12 degradation was enhanced in the PpAtg18 DD mutant during macropexophagy when compared with the wild type and the PpAtg18 AA mutant (Fig. 6 F). These results indicated that phosphoregulation of PpAtg18 was involved in pexophagic activity.

Next, we monitored the intracellular localization of the phosphorylation-defective PpAtg18AA-YFP and the phosphorylation-mimic PpAtg18DD-YFP mutants during the pexophagic process. Although PpAtg18DD-YFP showed a similar localization to the wild-type protein, PpAtg18AA-YFP was recruited to the vacuolar membrane more efficiently than was the wild-type protein, which resulted in enhanced VSM formation (Table 3 and Fig. 6 E). These data indicated that phosphoregulation of PpAtg18 was required for optimum activity of both micro- and macropexophagy by regulating the intracellular distribution of PpAtg18.

We evaluated how PpAtg18 phosphorylation levels affected nitrogen starvation–induced autophagy using the YFP-
Phosphorylation of PpAtg18 interferes with the interactions of PpAtg18 and PI(3,5)P2 in vitro
One of the important findings in this study is that the dephosphorylation of PpAtg18 increases its affinity for PI. Atg18 is a member of the PROPPIN family of proteins that bind to PI(3)P and PI(3,5)P2 via a seven -propeller motif (Dove et al., 2004). Recently, the crystal structure of KHSv2 was solved, 2012; Watanabe et al., 2012). Specificity to PI was mediated by ionic interactions between the negative charge of the PI head group and the basic amino acid residues.

### Table 3. VSM formation during micropexophagy in PpAtg18 AA and DD mutant strains

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<tr>
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The difference in the average of VSMs per cell between PpAtg18Wt and PpATG18AA was significant (P < 0.05). Wt, wild type.

*PpAtg18Wt, AA, or DD tagged 5×Flag were expressed under the Ppatg18 promoter in PpAtg18A. Micropexophagy was induced by adapting cells from methanol to glucose medium for 30 min.

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of sites 1 or 2. Furthermore, the hydrophobic loop of blade 6, conserved in all PROPPINs, was able to embed in the membrane, thus facilitating the interactions between PROPPIN and the membrane (Baskaran et al., 2012).

Our results indicated that dephosphorylation of either region, in blade 6 or blade 7, was sufficient to increase the PI-binding activity (Fig. S2 C). The higher affinity of PpAtg18 AA protein toward PI(3,5)P₂ compared with PpAtg18 DD protein (Fig. 3) suggested that the charged moieties of the phosphorylated regions inhibited the lipid-binding activity. One plausible explanation is that the negative charge of the phosphorylated residues simply impeded the hydrophobic interactions between the hydrophobic loop of blade 6 and the membrane (Fig. 7). Although the lipid-binding PH domain of protein kinase B was known to be phosphoregulated (Powell et al., 2003), to our knowledge, this is the first study that addresses, at molecular and biochemical levels, how phosphorylation of a protein modulates its binding to PIs.

Role of PpAtg18 phosphoregulation in the maintenance of vacuolar shape

When the dephosphorylated form of PpAtg18 increased and PpAtg18 was recruited to the vacuolar membrane by its PI(3,5)P₂-binding activity, vacuolar fission occurred. Phosphorylation-defective PpAtg18 AA mutants exhibited more septated vacuoles than did the cells expressing wild-type PpAtg18. In contrast, when the phosphorylated form of PpAtg18 increased and PpAtg18 dissociated from the vacuolar membrane, the vacuoles fused and formed a single rounded compartment. From these results, we conclude that the phosphorylation of PpAtg18 is correlated to its intracellular localization and vacuolar shape.

Atg18 is thought to be a negative regulator of PI(3,5)P₂ synthesis, and the localization cycle of Atg18 between the vacuolar membrane and the cytosol regulates PI(3,5)P₂ levels at the vacuolar membrane (Dove et al., 2004, 2009; Efe et al., 2007). Under hyperosmotic conditions, the cytosolic phosphorylated form of PpAtg18 was dephosphorylated, and recruited to the vacuolar membrane by binding vacuolar PI(3,5)P₂. Once PpAtg18 is recruited to the vacuolar membrane, the PI(3,5)P₂ levels on the vacuolar membrane may be decreased. When the levels of PI(3,5)P₂ on the vacuolar membrane are decreased and/or PpAtg18 is phosphorylated, PpAtg18 will be released from the vacuolar membrane. We think that dephosphorylation of PpAtg18 mediates a signal inducing vacuolar fission, as PpAtg18 AA mutation was sufficient to induce and enhance the vacuolar fission. This result also indicated that the vacuolar PI(3,5)P₂ level is sufficient to recruit PpAtg18 (Dove et al., 2004; Krick et al., 2006; Efe et al., 2007). However, for dissociation of PpAtg18 from the vacuolar membrane to occur, their interactions should weaken. It was difficult to determine whether the phosphorylation of PpAtg18 or the decrease in vacuolar PI(3,5)P₂ was the key event allowing the dissociation of PpAtg18 and the signaling of vacuolar fusion, because low vacuolar PI(3,5)P₂ levels were difficult to follow experimentally, and the phosphorylation mimic PpAtg18 DD mutant behaved like the wild type, and did not induce vacuolar fusion. Although the detailed mechanism for vacuolar fission itself should be investigated further, this study first clarified that vacuolar fission is regulated by phosphorylation of PpAtg18 through its association to the vacuolar membrane.

We tried to identify the upstream kinases that were responsible for PpAtg18 phosphorylation. Our results suggested that phosphorylation occurred at the vacuolar membrane. Specifically, both the PpJak1 mutant and the PpAtg18 FTTG mutant, in which PpAtg18 was not recruited to the vacuolar membrane, had dephosphorylated PpAtg18 under low glucose conditions (Fig. 5 C), whereas wild-type cells had only phosphorylated PpAtg18 under the same conditions (Fig. 4 A). P. pastoris mutants with a single disruption of the kinases PpAtg1, PpTor1, PpHog1, or PpVps15 retained phosphorylation of PpAtg18 (Fig. S3 E). Depletion of yeast casein kinase PpYck3 caused a partial defect in PpAtg18 phosphorylation. It is noteworthy that, in contrast to PpAtg18, in S. cerevisiae, Vps41, a component of a tethering complex for vacuolar homotypic fusion (HOPS family), is phosphorylated by Yck3 under vacuolar fission conditions and released into the cytosol (LaGrassa and Ungermann, 2005; Cabrera et al., 2009). Vps41 contains an amphipathic lipid-packing sensor (ALPS) motif, which interacts with highly curved membranes, and the phosphorylation of this motif prevents membrane recruitment of Vps41 (Cabrera et al., 2010). Vps41 seems to serve as a counterpart to Atg18 in regulating vacuolar shape.

Role of PpAtg18 phosphoregulation in autophagic processes

Atg2, a binding partner of Atg18, was not involved in the phosphorylation of PpAtg18. Structural analysis revealed that Atg2 bound to blades 1 and 2 of Atg18, which are at opposite faces relative to the lipid binding sites and the blade 6 loop involved in membrane association (Watanabe et al., 2012; Rieter et al., 2013). During autophagic processes, Atg18 is thought to recruit and complex with Atg2 at the PAS. This interaction does not affect the lipid-binding activity of Atg18 (Kobayashi et al., 2012; Rieter et al., 2013). Because macroautophagy proceeds despite the phosphorylation defect in the PpAtg18 AA mutant, phosphorylation in the loop of PpAtg18, which affects the PI(3,5)P₂-binding activity necessary for its recruitment to the vacuolar

Figure 7. Phosphoregulation of PpAtg18 for maintenance of vacuolar shape.
membrane, does not seem to directly affect the PI(3)P-binding activity of PpAtg18 necessary for autophagy. It may be noteworthy to point out that the phosphorylated form PpAtg18 still binds to PI3, which may be sufficient to execute macroautophagy.

Phosphoregulation of PpAtg18 was necessary for optimal execution of microexophagy, which involved formation of both the VSM and MIPA (Fig. 6 A), but was not necessary for other autophagic processes. During microexophagy, although VSM formation was enhanced in the phosphorylation-defective PpAtg18 AA mutant, PpPex12 degradation was retarded. This result likely reflected the fact that efficient recruitment of the PpAtg18 AA mutant to the vacuolar membrane reduced the amount of PpAtg18 in the PAS, which is necessary for MIPA formation. The efficiency of MIPA formation could not be assessed using the PpAtg18 AA mutant, as macroexophagy was induced after glucose adaptation and nascent pexophagosomes could not be distinguished from MIPA. We suggest that, during macroexophagy, dephosphorylated and phosphorylated forms of PpAtg18 function in VSM and MIPA formation, respectively, and that PpAtg18 phosphorylation is regulated accordingly. Although a defect caused by impaired phosphoregulation was not observed in macroautophagy, PpAtg18 was observed to be mainly in a phosphorylated form, which suggests that the dissociation of PpAtg18 from the vacuolar membrane enhances the recruitment of PpAtg18 to PAS for autophagosome formation.

Phosphoregulation of PpAtg18 governs the intracellular re-arrangement and coordinated dynamics of multiple organelles, i.e., vacuoles, autophagosomes, MIPA, and peroxisomes, in response to various environmental conditions and stresses. An example of such a highly coordinated event may be seen in fungi living in nature. Previously, we showed that the proper execution of pexophagy, but also to oxidative and osmotic stresses leading to changes in organelle life span and PpAtg18 expression, are necessary for fungal cells to proliferate on the leaves of Arabidopsis (Asakura et al., 2009; Kawaguchi et al., 2011). In nature, cells have to adapt not only to nutrient stresses leading to autophagy, but also to oxidative and osmotic stresses leading to changes in vacuolar shape. It is likely that, in P. pastoris, microexophagy is the biological process through which cells adapt simultaneously to both nutritional and environmental stresses, whereby phosphoregulation of Atg18 plays a critical role under natural conditions.

Materials and methods

Strains, media, and antibodies

The strains used in this study are listed in Table S1. Yeast extract/peptone/dextrose (YPD) medium consisted of 1% yeast extract, 2% Bacto Peptone, and 2% glucose. SD medium contained 0.67% yeast nitrogen base without amino acids and 2% glucose. Synthetic methanol (SM) medium contained 0.67% yeast nitrogen base without amino acids and 0.8% methanol. These synthetic media were supplemented with the appropriate amino acids (100 µg/ml arginine and 100 µg/ml histidine) for auxotrophic strains. Anti-FLAG (Wako), anti-GFP (Molecular Probes), anti-GST (GE Healthcare), anti-Pgk1 (Molecular Probes), anti–mouse-HRP (EMD Millipore), anti–goat-HRP (Abcam), and anti–rabbit-HRP antibodies (Rockland Immunochemicals, Inc.) were used in this study. The anti-PpPex12 antibody was a generous gift from S. Subramani (University of California, San Diego, La Jolla, CA).

Plasmid construction

The primers are listed in Table S2. The 5xFlag tag was amplified by using FLAGORF+[G+]FWx5 and FLAGORF+[TGA]RVpSil primers, and the fragments were inserted into pNT204, resulting in the plasmid pNT206. PpATG18 original promoter and ORF were amplified with PpATG18promoterfwBamHI and PpATG18orfr[TAAT]KavXhoI, and the PCR product was inserted into pNT206 or pNT205. The plasmids were designated pNT1801 and pNT1808, respectively. The PpATG18orfr was amplified using PpATG18ORF[ATG]FwBamHI and PpATG18ORF[TAAT]KavXhoI and then introduced into pSYS03 or pNT401, resulting in the pNT1111 and pNT1112 plasmids, pNT1801, pNT1808, and pNT1112 were subjected to site-directed mutagenesis by using PpATG18 ST387-391AFw, PpATG18 ST387-391ARv, PpATG18 ST387-391DFw, PpATG18 ST387-391DRv, PpATG18 ST491-495AFw, PpATG18 ST491-495ARv, PpATG18 ST491-495DFw, PpATG18 ST491-495DRv, PpATG18 ST491-495TFTGfw, and PpATG18 ST491-495TRGv. These resulting plasmids were designated pNT1802–1807, pNT1809, pNT1810, pNT1813, pNT1814, pNT1816, pNT1826, and pNT1827, respectively. To construct a more efficient expression vector, the Kpn1–XhoI fragment including PpATG18ORF from pNT1812 was inserted into pGAP2Z, resulting in pNT1815. For single mutation analysis, pNT1801 was subjected to site-directed mutagenesis by using PpATG18 S387AFw, PpATG18 S387ARv, PpATG18 S388AFw, PpATG18 S388ARv, PpATG18 T389AFw, PpATG18 T389ARv, PpATG18 T390AFw, PpATG18 T390ARv, PpATG18 S391AFw, PpATG18 S391ARv, PpATG18 S492AFw, PpATG18 S492ARv, PpATG18 S493AFw, PpATG18 S493ARv, PpATG18 T494AFw, PpATG18 T494ARv, PpATG18 S495AFw, and PpATG18 S495ARv. These resulting plasmids were designated pNT1817–1825.

For disruption of each null mutant, we used the primers (pDV–) listed in Table S2. The amplified PCR products were inserted into the disruption vectors.

Immunoblot analysis

To prepare samples for immunoblot analyses, harvested cells were suspended in lysis buffer (50 mM Tris–HCl, pH 7.5, 1 mM PMSF, 1 mM EDTA, EDTA-free complete protease inhibitor cocktail [Roche], and phosphatase inhibitor PhosSTOP [Roche]) and then lysed using Multi-Beads Shocker (Yasui Kikai). Cell extracts were centrifuged at 8000 g for 5 min at 4°C to remove cell debris, and dissolved in sample buffer as described previously (Tamura et al., 2010). Each sample (5 µg for PpAtg18 18S-Flag detection or 15 µg for YFP-PpAtg8, PpPex12, or PpPgk1 detection) was electrophoresed on a 7.5–8.0% SDS-PAGE gel (YP-PpAtg8, PpPex12, and PpPgk1). The gels were transferred to nitrocellulose (Flag, YFP, and Pgk1) or PVDF membranes (PpPex12) by semidry blotting (ATTO). The blots were incubated with anti-Flag (diluted 1:10,000) for 1 h, or anti-Pgk1 (diluted 1:10,000) in TBS-T buffer for 1 h or overnight. Then, the membranes were washed three times with TBS-T buffer and incubated with anti–mouse-HRP (diluted 1:10,000) or anti–rabbit-HRP (diluted 1:10,000) for 1 h. Finally, bound secondary antibodies were detected using Western Lightning (PerkinElmer) and the signals were analyzed with a Light Capture system (ATTO).

Phosphatase with or without heat inactivation was added to cell lysates and incubated at 30°C for 1 h. The samples were precipitated with acetonitrile and analyzed by immunoblotting. For Phos-tag experiments, we followed the manufacturer’s instructions. An 8% SDS-PAGE was prepared using Phos-tag acrylamide (25 or 50 µM) and an equal volume of 10 mM MnCl₂. After electrophoresis, the samples were transferred to a PVDF membrane and analyzed by immunoblotting.

Protein purification from P. pastoris

For purification of PpAtg18 variants, we used P. pastoris cells expressing GST-tagged proteins. Cells were cultivated in YPD media up to stationary growth and harvested by centrifugation at 2,000 g for 5 min. Cells were washed with 1× PBS buffer once, and then suspended in lysis buffer (PBS, 1 mM PMSF, 10 mM MgCl₂, 2 µg/ml Pepstatin A, 10 µg/ml Leupeptin, 0.05% [w/v] Tween 20, and phosphatase inhibitor PhosSTOP). Next, cells were lysed by sonication and centrifuged at 6,000 g for 15 min to yield cell lysates. Cell lysates were incubated with GS 4B (GE Healthcare) at RT for 30 min. The column was washed with 5× PBS buffer multiple times, and the captured protein was treated with reduced glutathione (for PIP strips, PIP arrays, and liposome pull-downs), or cleaved with PreScission Protease (for surface plasma resonance; GE Healthcare). In the phosphatase treatment experiment, we incubated the captured GST-PpAtg18 with λ-phosphatase for 2 h on the column. Then, to remove λ-phosphatase from the sample, the column was washed with 1× PBS buffer multiple times before exchanging the buffer with liposome binding assay buffer: LB buffer, 10 mM HesperosKOH, pH 8.0, and 50 mM NaCl.
Lipid dot blot analysis
PIP strip and PIP array assays were performed according to the protocols of Echelon Bioscience, Inc. For the PIP strip, membranes were incubated with 0.5 µg/ml purified proteins for 1 h. For the PIP array, membranes were incubated with 1.0 µg/ml protein for 3 h. Anti-GST antibody diluted 1:5,000 was incubated for 1 h and anti–goat-HRP diluted 1:10,000 was incubated for 1 h. For both PIP strip and PIP array experiments, the signals were acquired using the same exposure time.

Liposome preparation
Lipids used in this study were purchased from Sigma-Aldrich: DOPC, 1,2-Dioleoyl-sn-glycero-3-phosphocholine; DSPE, 1,2-Dioleoyl-sn-glycero-3-phosphoethanolamine; DPPS, 1,2-Dipalmitoyl-sn-glycero-3-phospho-L-serine; PI(3)P, 1-phosphatidylinositol-3-monophosphate; PI(4)P, 1-phosphatidylinositol-4-monophosphate; PI(3,5)P2, 1-phosphatidylinositol-3,5-bisphosphate; PI(4,5)P2, 1-phosphatidylinositol-4,5-bisphosphate. For the liposome pull-down assay, 100:45:45:10 DOPC/DSPE/DSPE/Pls was mixed in lipid mixing buffer (14:5:1 chloroform/methanol/dissolved water) to give a final concentration of 1 mg/ml. Similarly, for surface plasmon resonance analysis, 97/3 DOPC/Pls was mixed. These mixtures were lyophilized, and then resuspended in an aliquot of LB buffer by multiple freeze/thaw steps and sonication. The lipid-containing mixture was passed through a Mini-Extruder set (Avanti Polar Lipids, Inc.) with a 100-nm pore size filter to make liposomes.

Liposome binding assay
The purified GST-PaAg18 variants (1 µg) were incubated with 50 µl of liposomes in 200 µl of LB buffer containing 2.0 × 10^{-2} (Avanti Polar Lipids, Inc.) with a 100-nm pore size filter to make liposomes.

Surface plasmon resonance analysis
A Biacore2000 (GE Healthcare) with an L1 sensor chip was used. Liposomes were suspended in 10 mM Hepes-KOH, pH 7.5, 150 mM NaCl, and 1 mM MgCl2 and were loaded up to 6,000 RUs (resonance units). Before injection, 0.1% (wt/vol) BSA was applied at 20 µl/min for 2 min. Atg18, from which the N-terminal affinity tag had been cleaved, was loaded and dissociated at 80 µl/min. Injection and dissociation were performed for 90 s and 240 s, respectively. Data were analyzed by BIAevaluator software (GE Healthcare).

Microscopy analysis
A motorized inverted microscope (IX81; Olympus) equipped with a Uplan-Apochromat 100x/1.35 NA oil objective lens using an XF104-3 filter set (Omega Optics) was used for YFP, and a U-MRPHQ unit (Olympus) was used for FM 4-64 as described previously (Tamura et al., 2010). Both YFP and FM 4-64 signals were acquired using a Plan-Fluar 100x lens (Carl Zeiss) with pinhole setting to 1.02 airy units for YFP acquisition. Image data were captured with charge-coupled device camera DP30BW (Olympus). These data were acquired and analyzed by using MetaMorph version 7.0 (Molecular Devices), and saved as TIFF files. The TIFF image files were optimized for their contrast on Photoshop CS3 (Adobe). The scale bars in all experimental microscopy figures are equal to 2 µm.

Morphometric analysis
The number of budded vacuoles from each cell (VSM) was counted after cells had been shifted from methanol to glucose medium. In each experiment, at least 200 cells were analyzed, and the experiment was repeated at least three times. Similarly, we counted budding vacuoles in cells grown on SD medium or under hyperosmotic conditions. When the cells were cultured with oxidants (1 mM H2O2, 15 mM tBOOH, or 4 mM diamide), the ratio of the number of cells possessing a single spherical vacuole was determined from the acquired images. At least 500 cells were counted for each time point, and the experiments were repeated at least three times. In all experiments, the vacuoles were stained with FM 4-64.

Sample preparation for mass spectrometry
The purified protein was applied to an anion-exchange column (HiTrap DEAE; GE Healthcare) equilibrated with 20 mM Tris-HCl, pH 8.0, and eluted with a linear gradient of 0–1 M NaCl. Purified protein (60 pmol) in 20 mM Tris-HCl, pH 8.5, was diluted to 200 µl with 100 mM ammonium bicarbonate, and 20 µl of acetonitrile was added. Each sample was treated with 10 µl of a reducing agent, 10 mM Tris[2-carboxyethyl]phosphine, at 37°C for 45 min, and alkylated with 10 µl of cysteine blocking reagent, 50 mM iodoacetamide, at 24°C for 1 h in the dark. Proteins were digested with 0.1% (wt/vol) trypsin (Promega) in 100 mM ammonium bicarbonate at 37°C overnight. The digestion reaction was stopped by addition of 250 µl of 1% trifluoroacetic acid. After vortexing, 200 fmol amounts of the resulting samples were directly subjected to liquid chromatography tandem mass spectrometry (LC-MS/MS) analysis.

LC-MS/MS
LC-MS/MS analysis of the digested samples was performed with an RPLC system (Paradigm MS4B, Michrom BioResources, Inc.) interfaced with a hybrid mass spectrometer (LTQ-Orbitrap, Thermo Fisher Scientific) using a nanoelectrospray ionization device (AMR). Samples were loaded onto the trap cartridge and washed with mobile phase A (98% H2O, 2% acetonitrile, and 0.1% formic acid) for concentration and desalting. Subsequently, peptides were eluted over 20 min from the analytical column via the trap cartridge using a linear gradient of 5–45% mobile phase B (10% H2O, 90% acetonitrile, and 0.1% formic acid) at a flow rate of 1 µl/min. To survey the phosphorylated peptides in each sample, the mass spectrometer was operated in the data-dependent mode to automatically switch between high-resolution MS scan (resolution, 60,000; scan range, m/z 400–1,600) by the Orbitrap hybrid mass spectrometer and up to three concurrent MS/MS scans in the LTQ for the three most intense peaks selected from each MS scan. For determination of the phosphorylation sites on the phosphorylated peptides, the mass spectrometer was operated in the targeted MS/MS scan mode to switch between high-resolution MS and high-resolution MS/MS scans (resolution, 60,000); both were acquired using the Orbitrap in profile mode. Mascot software (version 2.1.1; Matrix Science) was used for matching searches against the PaAg18 protein sequence without an initial Met and with an artificial Gly-Pro-Leu-Gly-Ser sequence at the N terminus. Peptide mass tolerance was 20 ppm, fragment mass tolerance was 0.8 D, and trypsin specificity was applied with a maximum of two missed cleavages. Carbamidomethylation of Cys was allowed as a fixed modification, whereas oxidation of Met and phosphorylation of Ser and Thr were allowed as variable modifications. Phosphorylation sites were determined by manually comparing datasets of product ions acquired from Orbitrap MS/MS spectra, and theoretical fragment ions were estimated using the MSProduct tool on the ProteinProspector web site (http://prospector.ucsf.edu/prospector/mshome.htm).

Online supplemental material
Fig. S1 shows identification of phosphorylated regions by LC-MS/MS. Fig. S2 shows purification of PaAg18 proteins from P. pastoris. Fig S3 shows the role of PaAg18 in autophagic pathways. Table S1 lists yeast strains used in this study. Table S2 lists oligonucleotides used in this study. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201302067/DC1. Additional data are available in the JCB DataViewer at http://dx.doi.org/10.1083/jcb.201302067.dv.

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