Up-regulation of Osh6 boosts an anti-aging membrane trafficking pathway toward vacuoles

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ABSTRACT Members of the family of oxysterol-binding proteins mediate non-vesicular lipid transport between membranes and contribute to longevity in different manners. We previously found that a 2-fold up-regulation of Osh6, one of seven yeast oxysterol-binding proteins, remedies vacuolar morphology defects in mid-aged cells, partly down-regulates the target of rapamycin complex 1 (TORC1), and increases the replicative lifespan. At the molecular level, Osh6 transports phosphatidylserine (PS) and phosphatidylinositol-4-phosphate (PI4P) between the endoplasmic reticulum (ER) and the plasma membrane (PM). To decipher how an ER-PM working protein controls vacuolar morphology, we tested genetic interactions between OSH6 and DRS2, whose protein flips PS from the lumen to the cytosolic side of the Golgi, the organelle between ER and vacuoles in many pathways. Up-regulated OSH6 complemented vacuolar morphology of ∆drs2Δ and enriched PI4P on the Golgi, indicating that Osh6 also works on the Golgi. This altered PI4P-enrichment led to a delay in the secretion of the proton ATPase Pma1 to the PM and a rerouting of Pma1 to vacuoles in a manner dependent on the trans-Golgi network (TGN) to late endosome (LE) trafficking pathway. Since the TGN-LE pathway controls endosomal and vacuolar TORC1, it may be the anti-aging pathway boosted by up-regulated Osh6.

INTRODUCTION The family of oxysterol-binding proteins is conserved in all eukaryotes. They mediate non-vesicular lipid transport between membranes and thus play critical roles in cell growth and development [1]. Knocking out all seven oxysterol-binding proteins encoded by OSH genes in yeast leads to highly fragmented vacuoles and cell death [2]. Late endosomes, the organelle immediately upstream of vacuoles (mammalian lysosomes) in multiple membrane trafficking pathways, are also targets of oxysterol-binding proteins. In Caenorhabditis elegans, knocking down the expression of all four oxysterol-binding proteins leads to enlarged late endosome (LE) [3]. In HeLa cells, knocking down the expression of oxysterol-binding protein ORP1L leads to enlarged LE [3]. Since enlarged LE is usually a sign of senescent cells [4], normal functions of oxysterol-binding proteins are thus critical for longevity.

We previously found that a 2-fold up-regulation of Osh6 (Pess-Osh6) remedies vacuolar morphology defects in mid-aged cells and extends the replicative lifespan. Interestingly, the level of Osh6 protein in wild type cells declines with age. Moreover, up-regulation of Osh6 partly represses the target of rapamycin complex 1 (TORC1) but requires TORC1 for longevity [5]. This two-way relationship with TORC1 is supported by a recent study on the spatial dissection of TORC1. In yeast cells, the vacuolar TORC1 stimulates protein synthesis and cell growth while the late endosomal TORC1 inhibits autophagy, an anti-aging process through which vacuoles (mammalian lysosomes) degrade damaged and obsolete proteins, lipids, and organelles for longevity [6]. This spatial dissection of TORC1 offers us a new route to explore how Osh6 contributes to longevity.

Osh6 transports phosphatidylinositol-4-phosphate (PI4P) from the plasma membrane (PM) to the endoplasmic reticulum (ER), where Sac1 dephosphorylates PI4P, and in turn transports phosphatidylserine (PS) from the ER to the PM [7]. Other Osh proteins also relay PI4P to Sac1 in the ER but extend the lifespan when depleted from the cell [8]. PS is a unique ligand of Osh6 and its closest paralog Osh7 [7]. Down-regulation of the PS synthesis in ER leads to fragmented vacuoles [9]. PS synthesized in the ER has to
be transported to the Golgi and/or other organelles to control vacuolar morphology [10]. Disrupting the flip of PS from the lumen of the Golgi to the cytosolic side also leads to highly fragmented vacuoles [11]. A key enzyme for such flipping is Drs2 [12]. The fact that up-regulated Osh6 promotes vacuole fusion [5] and that Osh6 transports PS and PI4P [7] led us to hypothesize that Osh6 may mediate PS/PI4P transport in intracellular membranes.

In support of the above hypothesis, we found an accumulation of PI4P on the Golgi in cells of \( P_{\text{ERG6}}\)-OSH6. The accumulation site is likely the trans-Golgi network (TGN) since \( P_{\text{ERG6}}-\text{OSH6} \) mimicked mutants defective in Golgi PS and PI4P trafficking [12] in rerouting a portion of the PM protein Pma1 to vacuoles. Intriguingly, this rerouting of Pma1 was dependent on the TGN-to-LE trafficking pathway. Based on these new findings and other published results, we propose that up-regulation of Osh6 speeds up vesicle trafficking between the TGN and LE by adjusting the local concentration of PI4P and PS on the TGN.

**RESULTS**

**OSH6 genetically interacts with DRS2, whose protein works on the Golgi**

To explore how Osh6 affects vacuoles and longevity, we tested potential genetic interactions between OSH6 and genes involved in the metabolism of PS and PI4P, ligands of Osh6. Drs2 flips PS from the lumen of the Golgi to the cytosolic side and leads to fragmented vacuoles when mutated [11, 12]. We over-expressed OSH6 and other OSH genes in a \( \text{drs2}^{\Delta} \) mutant (Fig. 1A, 1B). Interestingly, over-expression of OSH6 by a galactose promoter assisted the growth of \( \text{drs2}^{\Delta} \) cells marginally but over-expression of its closest paralog \( OSH7 \) severely delayed the cell growth at 30°C (Fig. 1A). At 17°C, over-expression of \( OSH6 \) or \( OSH3 \) by a high copy plasmid complemented the cold sensitivity of \( \text{drs2}^{\Delta} \) (Fig. 1B). We further checked the vacuolar morphology of the \( \text{drs2}^{\Delta} \) mutants with over-expressed OSH genes and found that over-expression of OSH6 significantly promoted vacuole fusion in \( \text{drs2}^{\Delta} \) while over-expression of \( OSH7 \) only marginally affected the morphology (Figs. 1C, 1D, and Fig. S1A).

Different from their effects on \( \text{drs2}^{\Delta} \), over-expression of OSH7/7 did not obviously affect the growth of \( \text{sac1}^{\Delta} \) (Fig. S1B). For vacuolar morphology, over-expression of OSH6 did not show an obvious impact on \( \text{sac1}^{\Delta} \) (Fig. S1C). The different impact of OSH6 and OSH7 genes on \( \text{drs2}^{\Delta} \) and \( \text{sac1}^{\Delta} \) mutants suggest that Osh6 and Osh7 proteins have their own working locations in addition to their common ER-PM contact site. To search for such unique working places of Osh6, we then monitored the localization of PS and PI4P in cells of different mutants.

**Up-regulation of OSH6 causes accumulation of PI4P on the Golgi**

We first checked the localization of PS by the GFP-Lact-C2 probe, which was already used by many other labs [10, 13]. To use this plasmid, we upregulated the expression of OSH7 by replacing its endogenous promoter on its chro-

mosome by a short version of the promoter of ERG6 to obtain \( P_{\text{ERG6}}-\text{OSH7} \) as we did for the up-regulation of OSH6 [5]. Similar to a previous report [13], up-regulation of Osh6 enriched PS on the PM of small buds (Fig. S2). We did not notice any obvious intracellular accumulations of PS in either \( P_{\text{ERG6}}-\text{OSH6} \) or \( P_{\text{ERG6}}-\text{OSH7} \) cells when compared with wild type cells (Fig. S2). This lack of differences in PS localization seems inconsistent with the genetic interactions between OSH6 and DRS2 (Fig. 1). This apparent inconsistency is likely due to the limitation of our current assay method for PS localization.

Then we monitored the localization of PI4P. We chose the probe 2XPH-OSBP-GFP because it tends to bind Golgi localized PI4P due to the pH of the cytoplasm [14-16]. Like previously reported, 2XPH-OSBP-GFP visualized PI4P-decorated punctates, typical Golgi structures (Fig. 2A). In the wild type, some cells showed plasma membrane PI4P with this probe (see arrowhead-pointed cell in Fig. 2A). Expectedly, deletion of Sac1 (\( \text{sac1}^{\Delta} \)) accumulated PI4P (Figs. 2A, 2B). Interestingly, \( P_{\text{ERG6}}-\text{OSH6} \) accumulated PI4P inside cells (Figs. 2A, 2B). \( P_{\text{ERG6}}-\text{OSH7} \) cells also showed punctate PI4P, but with less brightness (Figs. 2A, 2B). The fraction of cells carrying high intracellular PI4P levels was much higher in \( \text{sac1}^{\Delta} \) and \( P_{\text{ERG6}}-\text{OSH6} \) than that in wild type and \( P_{\text{ERG6}}-\text{OSH7} \) (Fig. 2B).

A Fisher Exact test showed that the fraction of cells with high PI4P intensity (>50 AU/cell) in \( \text{sac1}^{\Delta} \) (31/64) (\( p=0.043 \)) and \( P_{\text{ERG6}}-\text{OSH6} \) (42/83) (\( p=0.016 \)) are significantly higher than that in wild type. Although both \( P_{\text{ERG6}}-\text{OSH6} \) and \( \text{sac1}^{\Delta} \) accumulated PI4P, they distribute PI4P differentially. \( P_{\text{ERG6}}-\text{OSH6} \) only showed punctate structures of PI4P, while \( \text{sac1}^{\Delta} \) cells also had PI4P on large membrane encircled organelles (arrow-pointed structure in Fig. 2A). Moreover, about 1/2 of \( \text{sac1}^{\Delta} \) cells showed bud-enriched PI4P, whereas only about 1/5 of \( P_{\text{ERG6}}-\text{OSH6} \) cells showed such an enrichment (Fig. 2C). In wild type cells, only about 1/10 of cells had such bud-enriched PI4P. The polarized secretion to the budding tip and small buds depends on dephosphorylation of PI4P by Sac1 [17]. Deletion of Sac1 likely caused an accumulation of PI4P in polarized secretory vesicles. The observation that \( P_{\text{ERG6}}-\text{OSH6} \) had a mild accumulation of PI4P in small buds (Fig. 2D) suggests that \( P_{\text{ERG6}}-\text{OSH6} \) has more polarized secretion and/or partial defect in dephosphorylation of PI4P in small budded cells. In support, \( P_{\text{ERG6}}-\text{OSH6} \) cells have more polarized actin cables [5], along which secretory vesicles are transported.

To check whether the punctate PI4P is on the Golgi in \( P_{\text{ERG6}}-\text{OSH6} \), we integrated the Sec7-mCherry coding DNA into the \( \text{URA3} \) gene [18] in \( P_{\text{ERG6}}-\text{OSH6} \) and \( P_{\text{ERG6}}-\text{OSH7} \) cells and then checked the colocalization of PI4P and Sec7 (a Golgi marker; Fig. 2D). In both \( P_{\text{ERG6}}-\text{OSH6} \) and \( P_{\text{ERG6}}-\text{OSH7} \) cells, PI4P did overlap with the Sec7-mCherry (see arrowhead-pointed structures in Fig. 2D), suggesting that up-regulation of Osh6 enriches PI4P on the Golgi. The higher enrichment of PI4P on the Golgi in \( P_{\text{ERG6}}-\text{OSH6} \) than in \( P_{\text{ERG6}}-\text{OSH7} \) may be a contributor for longevity, since \( P_{\text{ERG6}}-\text{OSH7} \) did not extend the replicative lifespan (data not shown).
FIGURE 1: OSH6 genetically interacts with DRS2. (A) Growth of drs2Δ cells with vector, P_{gal}-OSH6 (pCB248), or P_{gal}-OSH7 (pCB247) on SC-URA with glucose or galactose media at 30°C for two days. (B) Growth of drs2Δ cells with vector, high copy OSH6 (pCB237), or high copy OSH3 (pCB238) on SC-URA at 30°C for two days or 17°C for ten days. For A and B, 5 µl of serially diluted cells (0.1 OD/ml for the left) were spotted on the plate and then incubated. (C) Vacuolar morphology of drs2Δ cells with vector or the indicated plasmid. Overnight cultures were labeled with FM4-64 for one hour and chased at 30°C for three hours and then photographed. (D) Quantitative analyses of vacuolar morphology from Fig. 1C. Cells were divided into three categories based on the number of vacuolar vesicles/cell. Sample sizes are 138 for drs2Δ (vector), 180 for drs2Δ (P_{gal}-OSH6) and 194 for drs2Δ (P_{gal}-OSH7). A one-way ANOVA analysis shows that the fraction of cells with one vesicle/cell of P_{gal}-OSH6 is significantly different from that of vector (p<0.0001). Differences between P_{gal}-OSH7 and wild type is also significant (p=0.012).
FIGURE 2: Up-regulation of Osh6 traps PI4P on the Golgi. (A) Comparison of the localization of PI4P in wild type (BY4742), sac1\(\Delta\), \(\text{P}_{\text{PERS}-\text{OSH}6}\) (FTY536), and \(\text{P}_{\text{PERS}-\text{OSH}7}\) (FTY521). The indicated strains were transformed with the 2XPH-OSBP-GFP plasmid, grown to early log phase and photographed. The arrowhead points to a cell with PI4P on the PM in addition to intracellular punctate. The arrow points to a cell with PI4P-decorated circular organelles in sac1\(\Delta\). (B) Distribution of cells with different PI4P intensities. The intensity of PI4P of each cell was measured by the ImageJ software. Then cells were grouped based on the relative PI4P intensity with a ten arbitrary unit (reported by ImageJ) interval (X-axis). The fraction of each group in the whole set is presented on Y-axis. Sample sizes are 61 for wild type, 64 for sac1\(\Delta\), 83 for \(\text{P}_{\text{PERS}-\text{OSH}6}\), and 69 for \(\text{P}_{\text{PERS}-\text{OSH}7}\). (C) Distribution of cells with bud-enriched PI4P in different strains. A cell was counted as ‘bud enriched’ if its bud PI4P signal/(bud signal + mother signal) is larger than 0.6. Sample sizes are 31 for wild type, 34 for sac1\(\Delta\) and 35 for \(\text{P}_{\text{PERS}-\text{OSH}6}\). Fisher exact test shows that sac1\(\Delta\) is significantly different from wild type (\(P=0.029\)) and that \(\text{P}_{\text{PERS}-\text{OSH}6}\) is not significantly different from wild type. (D) PI4P colocalized with the trans-Golgi network marker Sec7-mCherry. Cells of the Sec7-mCherry integrated version of \(\text{P}_{\text{PERS}-\text{OSH}6}\) (FTY624) and \(\text{P}_{\text{PERS}-\text{OSH}7}\) (FTY625) strains were transformed with the 2XPH-OSBP-GFP plasmid. Early log phase cells of the transformants were photographed under FITC or Texas Red filter. Representative images are shown here. Arrow heads point to areas where PI4P overlapped with Sec7-mCherry.

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Up-regulation of OSH6 delays the Pma1 secretion, a function of the Golgi

Since both PS and PI4P on the Golgi are required for proper secretion of the PM protein Sur7 and Pma1 [12], the accumulation of PI4P on the Golgi in P_{ERG6-OSH6} (Fig. 2) may affect the secretion. Thus, we monitored the localization of PM proteins. For this purpose, we constructed P_{ERG6-OSH6} Sur7-GFP and P_{ERG6-OSH6} Pma1-mCherry strains by mating P_{ERG6-OSH6} with Sur7-GFP or Pma1-mCherry strains [19].

Up-regulation of Osh6 (P_{ERG6-OSH6}) did not show obvious effects on the PM Sur7 (Fig. S3), but affected the secretion of Pma1 (Fig. 3A). First, P_{ERG6-OSH6} cells accumulated more Pma1 inside cells; almost every cell had intracellular Pma1. The intracellular accumulation sites of Pma1 are likely in vacuoles, since the structure of the Pma1-accumulated organelle in some small-budded cells is very similar to vacuolar segregation structure during budding (see arrow pointed structure in Fig. 3A). Second, many small buds of P_{ERG6-OSH6} cells did not show discernible Pma1 (see arrowhead-pointed buds in Fig. 3A), suggesting an alteration in secretion and/or endocytosis in the bud.

To double check the effect of up-regulation of Osh6 on Pma1 in small buds, we conducted a zygote assay (Fig. 3B). We mated wild type Pma1-GFP with P_{ERG6-OSH6} Pma1-mCherry and then monitored the fluorescence of zygotes. As shown in Fig. 3B, the Pma1-GFP signal from the wild type parent arrived at the zygote bud earlier than the Pma1-mCherry signal from the P_{ERG6-OSH6} parent (see arrow-pointed buds in Fig. 3B), indicating that up-regulation of Osh6 delays the arrival of Pma1 to the bud PM.
ver, the intracellular Pma1 in P_{ERG6-OSH6} moved to the other parent and the bud, phenocopying what vacuoles do in such zygotye assays [20].

**Decreasing Golgi PI4P abrogates Osh6’s effects on Pma1 distribution**

To confirm the Golgi localization of PI4P in P_{ERG6-OSH6}, we monitored the PI4P intensity after glucose starvation (Fig. 4). Glucose starvation reallocates the ER-localized Sac1 to the Golgi and prevents the accumulation of PI4P on the Golgi [21]. In line with this previous report, glucose starvation decreased the intensity of PI4P punctate in wild type cells in our assays (compare the GFP pictures of row 1 and row 2 in Fig. 4A). Upon glucose starvation, P_{ERG6-OSH6} cells showed a clear decrease in PI4P intensity (compare the GFP pictures of row 1 and row 2 in Fig. 4B). In addition to Sac1 action, glucose starvation also changes cytosolic pH and decreases the binding of Golgi PI4P by PH domains [16]. Both possibilities (Sac1, pH) support that PI4P-decorated punctates in P_{ERG6-OSH6} cells are Golgi structures.

Along with the decrease of the Golgi PI4P levels, the secretion pattern of Pma1 was also changed upon glucose starvation. In log phase cells grown in a medium with 2% glucose, the Pma1-mCherry signal on the PM of the mother-bud neck is not as bright as other areas of PM of the mother cell (see hollow arrow pointed areas in wild type (Fig. 4A) and P_{ERG6-OSH6} (Fig. 4B)). Glucose starvation made Pma1 homogenously distributed on the mother PM (see solid arrow pointed cells in Fig. 4). Moreover, we found that treating yeast cells with 5 mM MnCl₂ also decreased the Golgi PI4P level (Fig. 4) and simultaneously altered the secretion of Pma1 similar to what glucose starvation did (Fig. 4). These observations further confirm that up-regulation of Osh6 causes accumulation of PI4P on the Golgi.

**The effects of up-regulation of OSH6 rely on the Golgi-to-LE trafficking**

While multiple pathways transport post-Golgi vesicles to vacuoles, the pathway that depends on the level of Golgi PI4P is the TGN to LE (TGN-LE) pathway (also termed the CPY pathway) [22]. To test whether the TGN-LE pathway is used by Osh6 in rerouting Pma1 to vacuoles, we checked the impact of Gga2, which binds Golgi PI4P and catalyzes the formation of the CPY vesicles [22], and Vps13, which recycles materials from LE back to the Golgi [23] on the Pma1 localization in P_{ERG6-OSH6} cells (Fig. 5). As a control, we also tested the impact of Lag1, a ceramide synthase working in the ER [24]. In the double mutant P_{ERG6-OSH6 gga2Δ} or P_{ERG6-OSH6 vps13Δ}, the Pma1 localization was similar to that in the corresponding gga2Δ or vps13Δ single mutant, where the intracellular Pma1 is dispersed to almost the whole cytoplasm (Fig. 5A). Similar to the trafficking of Pma1, a growth phenotype of P_{ERG6-OSH6 gga2Δ} also followed the pattern of the gga2Δ single mutant; P_{ERG6-OSH6 gga2Δ} was as resistant as gga2Δ to 5 mM MnCl₂ and as sensitive as gga2Δ to 50 µM CdCl₂ (Fig. 5B). The dependency of Osh6 on Gga2 supports that Osh6 works through the TGN-LE pathway. Different from gga2Δ or vps13Δ mutants, the P_{ERG6-OSH6 lag1Δ} double mutant behaved similar to P_{ERG6-OSH6} in enriching Pma1 into certain organelles (Fig. 5A).

**Up-regulation of OSH6 has a unique role in restoring cell growth**

Pma1 is secreted to the PM by complexing with sphingolipids carrying very long chain fatty acids (VLCFA) in the ER and the Golgi [25]. Since sphingolipids with VLCFA are also required for functions of vacuolar ATPase and vacuolar morphology [26, 27], we studied interactions between Osh6 and Sur4, which synthesizes VLCFA and controls vacuolar morphology via the TGN-LE trafficking step [28], and Lag1, which incorporates VLCFA into ceramide, the precursor for sphingolipids. Over-expressing OSH6 by a high copy plasmid restored vacuole fusion in lag1Δ and sur4Δ while over-expression of OSH4 did not show such restoration (Fig. 6A and Fig. S4). The observations that high levels of Osh6 down-regulated the secretion of Pma1 (Fig. 3) while promoted vacuole fusion in mutants (lag1Δ and sur4Δ defective in enzymes in the ER (Figs. 6A, 6B, and Fig. S4) further support the idea that Osh6 works on the Golgi.

To further test the genetic interactions between OSH6 and LAG1, we tested growth phenotype of the P_{ERG6-OSH6 lag1Δ} double mutant on caffeine plates (Fig. 6C). Caffeine inhibits TORC1 in a manner similar to rapamycin [29]. Shocking cells by 0.2% caffeine at 37°C for two days led to growth arrest (see middle panel of Fig. 6C). The growth was recovered after transferring this plate to room temperature (24°C) and incubating for five days (see row 1 and row 2 in Fig. 6C). Although lag1Δ showed a weak recovery, the P_{ERG6-OSH6 lag1Δ} double mutant recovered 10- to 100-fold better than lag1Δ did (rows 3 and 4 in Fig. 6C). Different from its effects on lag1Δ, P_{ERG6-OSH6} only very mildly recovered the growth of a mutant missing Vps13, which facilitates the TGN-LE trafficking (rows 5 and 6 in Fig. 6C). The recovery of growth after TORC1 repression requires TORC1 activation to stimulate protein synthesis as well as TORC1 repression to relieve the inhibition on microautophagy, which affects vacuolar morphology [30]. Since LE TORC1 inhibits autophagy including microautophagy and vacuolar TORC1 stimulates cell growth [6], we interpret the recovery of growth of the P_{ERG6-OSH6 lag1Δ} mutant (Fig. 6C) and the promotion of vacuole fusion of Osh6 in lag1Δ and sur4Δ (Figs. 6A, 6B, and Fig. S4) as that up-regulation of Osh6 differentially affects the endosomal TORC1 and vacuolar TORC1 by altering the TGN-LE trafficking step.

**DISCUSSION**

While Osh6 and its homologs work on multiple organelles including the ER, the PM, secretory vesicles, and mitochondria [7, 13, 31, 32], it differs from other Osh proteins in its effects on longevity. OSH6 extends the replicative lifespan when up-regulated, but other OSH genes (OSH1, OSH2, OSH3, OSH4) extend the lifespan when deleted from the genome [5, 8]. The unique roles of Osh6 and the critical contribution of the TGN-LE trafficking to pro-longevity pathways [6] led us to propose that Osh6 accelerates the
FIGURE 4: Golgi PI4P controls Pma1 secretion. (A) The wild type Pma1-mCherry strain (QAY559) was transformed with the PI4P-labeling (2XPH-GFP) plasmid. Transformants were grown in SC-URA with 2% glucose to early log phase (first row), and then starved for glucose for 30 minutes (second row) or treated with 5 mM of MnCl2 for 30 min (third row) and photographed. (B) The $P_{erg6^\rightarrow osh6}$ Pma1mcherry strain (FTY520) was transformed with the PI4P-labeling (2XPH-GFP) plasmid. Transformants were grown in SC-URA to mid-log phase (first row), and then starved for glucose for 30 minutes (second row), or treated with 5 mM of MnCl2 (third row). Hollow arrows point to mother-bud neck where the Pma1 signal is not as bright as other PM of its mother cell. Solid arrows point to small or mid-size budded cells with homogeneously distributed Pma1 on the PM of the mother cell.
TGN-LE vesicle trafficking and differentially affects endosomal TORC1 and vacuolar TORC1 to increase longevity.

The TGN is a working place of Osh6

Our phenotypic and subcellular localization analyses along with other group’s results on Osh6 and PI4P suggest that a subset of Osh6 works on the TGN. Although both \( P_{\text{ERG6}}-\text{OSH6} \) and \( \text{sac1} \Delta \) accumulated PI4P inside cells, \( P_{\text{ERG6}}-\text{OSH6} \) caused an enrichment of PI4P on Golgi-like punctate while in \( \text{sac1} \Delta \) cells PI4P could also be found on other membranes that gave a faint signal (see arrow-pointed cell in Fig. 2A). In log phase cells, intracellular PI4P localizes to the Golgi membrane due to the function of the PI 4 kinase Pik1[33]. In \( \text{sac1} \Delta \) cells, PI4P also localizes to other organelles including endosomal/vacuolar membranes and leads to a large vacuole with multiple invaginations [34] (also see Fig. S1C). Different from \( \text{sac1} \Delta \) cells, \( P_{\text{ERG6}}-\text{OSH6} \) cells have normal vacuolar morphology, two to five vacuolar vesicles per cell for most cells [5]. Thus, \( P_{\text{ERG6}}-\text{OSH6} \) could not accumulate PI4P on endosomal/vacuolar membranes. Indeed, PI4P in \( P_{\text{ERG6}}-\text{OSH6} \) overlapped with the Golgi marker Sec7 (Fig. 2D). The localization data of Osh6 from Orin’s lab show that Osh6 can work on TGN. A chromosomal version of GFP-tagged Osh6 with its endogenous promoter localizes to the PM and intracellular organelles in small buds [35]. During yeast cell division, organelles such as ER, late Golgi elements, and vacuoles are transported to buds at an early stage of budding [20, 36, 37]. Since ER hosts Sac1 and degrades PI4P, we conclude that the enriched PI4P in small buds of \( P_{\text{ERG6}}-\text{OSH6} \) (Fig. 2C) is a sign of Osh6 working on late Golgi elements. In line with this conclusion, in \( P_{\text{ERG6}}-\text{OSH6} \) cells a portion of Pma1 was rerouted to vacuoles (Fig. 3), phenocopying mutants missing Drs2 or Osh4, two proteins working on the TGN [12].

Osh6 promotes TGN-LE transport to extend the lifespan

Three lines of evidence suggest that elevated Osh6 promotes TGN-LE membrane trafficking as depicted in Fig. 7. First, \( P_{\text{ERG6}}-\text{OSH6} \) caused an enrichment of PI4P on the Golgi (Fig. 2) and this relied on the TGN-LE protein Gga2, which binds Golgi PI4P for its effects (Fig. 5).
Second, over-expression of OSH6 restores vacuole fusion of mutants defective in TGN-LE trafficking. Deletion of Sur4 compromises the TGN-LE trafficking but does not affect the TGN-vacuole pathway [28]. In this work, we observed that over-expression of OSH6 but not OSH4 restored vacuole fusion in sur4Δ (Fig. S3) and lag1Δ (Fig. 4), cells which are defective in incorporating Sur4’s product into sphingolipids. The specific growth recovery effect of PERG6-OSH6 on lag1Δ (Fig. 6) suggests that the TGN-LE trafficking step is the most likely route for Osh6 to control vacuolar morphology and TORC1.

Third, the effect of up-regulation of Osh6 on Pma1 lo-

**FIGURE 6:** Up-regulation of OSH6 complements defects of lag1Δ. (A) Comparison of vacuole morphology of lag1Δ with the indicated plasmids listed on the left. Vector: YEp24, OSH4ox: YEp24-OSH4; OSH6ox: YEp24-OSH6. Log phase cells were labeled with FM4-64 for one hour and chased for three hours at 30°C and then photographed. (B) Quantitation of cells with different categories of vacuoles. Sample sizes are 98 for lag1Δ(vector), 105 for lag1Δ(OSH4ox) and 102 for lag1Δ (OSH6ox). A one-way ANOVA shows that the fraction with one vacuolar vesicle/cell of lag1Δ (OSH6ox) is significantly higher than that of lag1Δ (vector) (p< 0.0001). (C) Recovery of cell growth after caffeine and high temperature arrest. Overnight cultures of wild type (BY4742), PERG6-OSH6 (FTY373), lag1Δ, PERG6-OSH6 lag1Δ (FTY527), vps13Δ, and PERG6-OSH6 vps13Δ (FTY534) were serially diluted by 10-fold (starting at 0.1 OD600/ml from the left). Five µl of serial diluted cells of the indicated strains were spotted on YEPD (left) and YEPD + 0.2% caffeine plate (middle) and incubated at 37°C for two days. Then the caffeine plate was incubated at room temperature (24°C) for five days.
Osh6 may accelerate TGN-LE transport by adjusting the local PS and PI4P concentrations of the early TGN

Osh6's two lipid ligands, PS and PI4P, are both crucial for the formation of post-Golgi vesicles and post-Golgi membrane trafficking. Upon maturation of the trans-Golgi cisternae, vesicles carrying Pma1 and other cargoes are budded from the late TGN cisternae and transported to the PM (summarized in Fig. 7A). The Pma1-vesicles require balanced PI4P and PS [12]. During the process of TGN maturation, the early cisternae of the TGN forms vesicles destined to LE [41]. The formation of TGN-LE vesicles depends on Gga2-PI4P interactions [22]. In active growing cells, the TGN-LE transport is very slow [42]. This slow rate is likely caused by lack of PI4P on the early TGN cisternae, since the majority of Golgi PI4P is used for the formation of secretory vesicles and is consumed by Osh4 and other Osh proteins during secretion [12, 17, 31].

Elevated Osh6 could accelerate the TGN-LE transport by its PI4P/PS swapping activity (Fig. 7B). The formation of vesicles toward LE requires PI4P [22]. Moreover, PS on LE needs to be transported back to the TGN by the LE-TGN retrograde pathway [43, 44], since elevated PS or PS/phosphatidylethanolamine (PE) ratio on LE and vacuolar membranes leads to fragmented vacuoles [45]. Osh6's working on TGN may transport PI4P to the early TGN cisternae and remove PS from that area (dark arrows in Fig. 7B). A support for such removal of PS is the observation that over-expression of OSH6 complemented defects of drs2Δ (Fig. 1). This complementation suggests that Osh6 can glean PS for some essential Golgi functions that are normally achieved by Drs2-flipped PS. The potential PI4P/PS swapping on the TGN facilitates both the formation of vesicles toward LE and the LE-to-TGN retrograde trafficking. Such elevated retrograde trafficking also leads to a decrease of LE membrane and thus endosomal TORC1. Testing this model in future studies would reveal detailed mechanistic links between TGN-LE trafficking and endosomal/vacuolar TORC1 activities and hence provide insights on how oxysterol-binding proteins control TORC1 and longevity in other organisms.

MATERIALS AND METHODS

Strains, plasmids, yeast manipulations, and media

All yeast strains (listed in Table S1) are derivatives of BY4742. The PS-labeling GFP-Lact-C2 plasmid was kindly provided by...

FIGURE 7: A working model for Osh6’s role in TGN-LE trafficking. (A) A summary of the maturation of the trans-Golgi network (TGN) and post-Golgi trafficking. The formation of Pma1-vesicles for secretion occurs at late cisternae of the TGN while vesicles carrying carboxypeptidase Y (CPY) or other vacuolar hydrolases to late endosome (LE) occurs in early cisternae of the TGN [41] (see text for detail). (B) A hypothetical action of up-regulated Osh6. Up-regulation of Osh6 transports more PI4P to the early cisternae of the TGN, which facilitates the TGN-to-LE trafficking since the formation of the CPY-carrying vesicles depends on Gga2-PI4P binding. The accelerated TGN-LE trafficking refreshes fresh lipids to LE and vacuoles and thus support vacuolar functions. As a side effect, transporting PI4P out of the late cisternae of TGN delays the secretion of Pma1 and leads to the rerouting of Pma1 to LE and vacuoles.

Drs. Gregory Fairn and Vanina Zaremberg. The PI4P-labeling 2XPH-OSBP-GFP plasmid pPGK1303 was kindly provided by Dr. Christina Mitchell. OSH plasmids (pCB237 ( Yep24-OSH6), pCB238 (Yep24-OSH3), pCB241 (Yep24-OSH4), pCB247 (Pgal-OSH7), pCB248 (Pgal-OSH6)) were kindly provided by Dr. Christopher Beh. Construction of Pgal-OSH7 is described below. Double and triple mutants of Pgal-OSH6 carrying Pma1-mCherry or Sur7-GFP were constructed by standard yeast mating of Pgal-OSH6 with Pma1-mCherry, Sur7-GFP strains provided by Dr. Mara Duncan [19], induction of meiosis, and tetrad dissection. Standard yeast media YEPD, yeast extract (1%)-peptone (2%)-dextrose (2%), was used unless otherwise stated.
Construction of \( \text{P}_{\text{ERG6}}-\text{OSH7} \) mutants

The strategies for constructing previously described \( \text{P}_{\text{ERG6}}-\text{OSH6} \) mutants [5] were employed to construct the \( \text{P}_{\text{ERG6}}-\text{OSH7} \) in this study. The \( \text{P}_{\text{ERG6}}-\text{OSH7} \) strain was constructed by switching the endogenous promoter of \( \text{OSH7} \) with the promoter of \( \text{ERG6} \) via in vivo recombination. The promoter of \( \text{ERG6} \) on the plasmid pRS316-ERG6 [46] was PCR-amplified by the primer \( \text{Perg6OSH7up} \) (5’TATCGAATTTAGGAAGTTAATCGTCGTGACTATTGTTATCTCC) and \( \text{Perg6OSH7down} \) (5’ACTGGTTGTAAGAAATGTCCTTGGTATGGAGGCCATCATCGATGTGGCCTACT). This PCR-amplified fragment contained the \( \text{URA3} \) gene followed by the basal \( \text{ERG6} \) promoter flanked by sequences homologous to the \( \text{OSH7} \) promoter. This fragment was transformed into BY4742. Verification of the correct insertion-replacement was confirmed by PCR with primers \( \text{RCERG6up} \) (5’ATAGTTCGGGTGTTTT) and \( \text{RCosh7down} \) (5’TGATCGATCTTCTTCATG). A 1.5 kb PCR-amplified fragment was used as the diagnostic band for the \( \text{P}_{\text{ERG6}}-\text{OSH7} \) replacement.

PS, PI4P, and vacuole labeling

A plasmid expressing the GFP-Lact-C2 fusion protein, which binds PS [10] or a plasmid expressing PI4P-binding marker proteins (pPGK1303/PH-OSBP-GFP) [15] was transformed into the wild type BY4742 or different mutant strains. Transformation was done with the LiAc protocol following all the steps listed in [47]. If the mutant strain carried the \( \text{URA3-P}_{\text{ERG6}} \) promoter, the \( \text{URA3} \) marker was replaced by a Kanamycin-resistance marker with BamHI-linearized M3927 obtained from Addgene [48] before PS and PI4P assay. Transformants were grown in SC-URA liquid media at 30°C to early log phase (OD600 between 0.2 to 0.4). Cells were observed under a 90i microscope and photographed under the FITC filter for GFP and Texas Red filter for mCherry. The resulting pictures were imported to the ImageJ software to report the fluorescent intensity of the region of interest (bud, mother) that was drawn manually. Intensities obtained for each cell were used for distribution analyses.

Vacuoles were labeled by FM4-64 and chased as described before [49].

The co-localization of PI4P with the Golgi marker Sec7-mCherry

The \( \text{URA3}::\text{Sec7-mCherry} \) DNA linearized by XcmI and Bst8I from YIpLac211-SEC7-mCherry28x [18] was transformed into \( \text{URA3-P}_{\text{ERG6}}-\text{OSH6} \) (FTY373) and \( \text{URA3-P}_{\text{ERG6}}-\text{OSH7} \) (FTY437) cells. The resulting transformants were selected on SC+URA+S-fluoroacetate acid plate. After confirming the lack of growth on SC-URA plates, the Sec7-mCherry version of \( \text{P}_{\text{ERG6}}-\text{OSH6} \) (FTY624) and \( \text{P}_{\text{ERG6}}-\text{OSH7} \) (FTY625) strains were transformed by the plasmid carrying the PI4P-binding probe (2XPH-OSBP-GFP). The resulting transformants were grown to early log phase in SC-URA and then photographed under the FITC (for PI4P) and Texas Red (for Sec7) filters.

Image quantitation and statistical analyses

All the pictures were analyzed by the ImageJ software to measure the fluorescent intensity of the region of interest (bud, mother) that was drawn. For quantitative analysis, we used Graph Pad Prism 8 software to generate all the graphs in this study. The significance of the difference in the mean values was determined by 1 WAY ANOVA that was applied on all the data by using Tukey’s multiple comparisons test, all those tests indicated P value < 0.0001 was considered as significant. Also, other statistics analysis such as Fisher exact test was conducted to judge whether the fraction of cells with bud-enriched PI4P is significant between two samples.

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SUPPLEMENTAL MATERIAL

All supplemental data for this article are available online at www.microbialcell.com.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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REFERENCES


entifies Ubx2 as a critical regulator of lipid saturation and lipid bilayer stress.


