Termination of Isoform-Selective Vps21/Rab5 Signaling at Endolysosomal Organelles by Msb3/Gyp3

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Traffic through endosomes and lysosomes is controlled by small G-proteins of the Rab5 and Rab7 families. Like humans, Saccharomyces cerevisiae has three Rab5s (Vps21, Ypt52 and Ypt53) and one Rab7 (Ypt7). Here, we elucidate the functional roles and regulation of the yeast Rab5s. Using GFP-tagged cargoes, a novel quantitative multivesicular body (MVB) sorting assay, and electron microscopy, we show that MVB biogenesis and thus MVB cargo sorting is severely impaired in *vps21* Δ *ypt52* Δ double mutants. Ypt53, the third Rab5 paralog, is hardly expressed during normal growth but its transcription is strongly induced by cellular stress through the calcineurin-Crz1 pathway. The requirement for Rab5 activity in stress tolerance facilitated identification of Msb3/Gyp3 as the principal Rab5 GAP (GTPase accelerating protein). In vitro GAP assays verified that Vps21 is a preferred Gyp3 target. Moreover, we demonstrate that Gyp3 spatially restricts active Vps21 to intermediate endosomal compartments by preventing Vps21 accumulation on lysosomal vacuoles. Gyp3, therefore, operates as a compartmental insulator that helps to define the spatial domain of Vps21 signaling in the endolysosomal pathway.

Key words: endosome, ESCRT, GAP, G-protein, LUCID, lysosome, MVB, Rab, vacuole, Ypt, autophagy

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Membrane traffic through the secretory and endolysosomal systems is highly dynamic. Within these systems small G proteins of the Arf and Rab families regulate compartmental identity and control rates of membrane influx and egress from each compartment (1–4). Rabs are anchored to membranes through C-terminal prenyl anchors. In their inactive GDP-bound state, Rabs can be extracted from membranes and relocated by the cytoplasmic chaperone GDI (GDP dissociation inhibitor). Activated GTP-bound Rabs and Arfs bind effector proteins and

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complexes that execute diverse functions in membrane traffic including vesicle budding, transport, and fusion. Rab signaling is stringently regulated. Like other small G proteins within the Ras superfamily, Rabs bind GDP tenaciously and are activated through catalyzed exchange of GTP for GDP mediated by guanosine nucleotide exchange factors (GEFs). Rabs generally have slow intrinsic GTPase activity and are inactivated by GTPase accelerating proteins (GAPs). All known Rab GAPs contain TBC domains. In ternary Rab-GTP-GAP complexes, the TBC domain supplies catalytic Arg and Gln residues which stabilize the hydrolytic transition intermediate (5,6). Hence, the integrated activities of GEFs, GAPs and chaperones define the spatial and temporal domains of Rab signaling and activity.

Anterograde transport from early to late endosomes, and then to terminal lysosomes, is principally controlled by members of the Rab5 and Rab7 families, with Rab5 paralogs operating at earlier compartments and Rab7 at later compartments (7). Recent studies also suggest that Rab5 and Rab7 small G proteins coordinate retrograde traffic to the Golgi through interactions with the retromer complex (8,9). Saccharomyces cerevisiae, like humans, has three Rab5 paralogs that reside on endosomes (Vps21, Ypt52 and Ypt53), and one Rab7 (Ypt7) found mainly on the vacuolar lysosome (10-13). No clear role for Ypt53 has been established, and the relative roles of Vps21, Ypt52 and Ypt53 in multivesicular body (MVB) biogenesis and cargo sorting have not been fully characterized. During maturation of the late endosome Rab5 is rather abruptly replaced by Rab7 in a process termed endosomal Rab conversion (14). The acquisition of Rab7 is thought to confer competence for fusion with terminal endolysosomal compartments.

An important waypoint within the endolysosomal system is the late endosomal MVB. At the MVB, ESCRTs (endosomal sorting complexes required for transport) package specific cargo into intraluminal vesicles (ILVs) that sequester cargo away from the cytoplasm (15,16). When mature MVBs fuse with lysosomes, ILVs and their cargos are dumped into the lysosome lumen and destroyed, while lipids and transmembrane proteins not packaged into ILVs are delivered to the lysosomal limiting membrane.

The relative timing and regulatory dependencies of ESCRT activity, Rab7 activation and Rab5 inactivation are not fully understood. Rab7 recruitment and signaling require the activities of the Vps-C effector complexes HOPS and CORVET (14,17) and the GEF complex Ccz1-Mon1 (18,19). In contrast, it has been unclear when and where GAPs negatively regulate Rab5 in the endolysosomal Rab cascade. Multiple yeast GAPs have been reported to target

Vps21 *in vitro* (5,20), but it has been unclear which (if any) of these GAPs terminate Rab5 signaling during normal endolysosomal transport.

We performed genetic, biochemical and cell biological analyses to elucidate the roles and regulation of the three *S. cerevisiae* Rab5 paralogs. We now report that Rab5 signaling (by either Vps21 or Ypt52) is essential for MVB cargo sorting and biogenesis, and show that Ypt53 is a stress-induced functional homolog of Vps21. We exploit a general requirement for Rab5 signaling during ionic stress to identify Msb3/Gyp3 as the principal *in vivo* GAP of Vps21, and demonstrate that Gyp3 (Gyp, GAP for Ypt/Rab proteins) has a pivotal role in the endosomal Rab cascade: it spatially restricts Vps21 signaling to prevacuolar endosomal compartments by preventing accumulation of active Vps21 on the terminal vacuole.

Results

Roles of yeast Rab5 paralogs in Golgi-endosome traffic

Correct targeting of the soluble vacuolar hydrolase CPY (Figure 1A) requires cycling of the CPY receptor Vps10 between late Golgi and endosomal compartments (21,22). Vps10 or CPY missorting results in CPY mistargeting into the extracellular space. Of the three *S. cerevisiae* Rab5 paralogs, Vps21 is known to function in CPY traffic (12),



Figure 1: Relative contributions of three *S. cerevisiae* Rab5 paralogs in Golgi-endosome traffic. A) Cartoon depicts how carboxypeptidase Y (CPY) sorting defects depend largely upon Golgi-endosome traffic of the CPY receptor, Vps10. B) Measurement of extracellular CPY-invertase in cells lacking Rab5 paralogs or Rab5 family GEF, Vps9. Bars indicate mean \pm SD of triplicate samples from a representative experiment.

as is its GEF Vps9 (23). Conversely, the loss of two other Rab5 paralogs, Ypt52 and Ypt53, did not strongly impair CPY sorting (10). Using a chimeric CPY-invertase fusion to quantify CPY secretion (Figure 1B), we verified the sorting defects of *vps21* Δ and *vps9* Δ single mutants and observed a strong synthetic defect in a *vps21* Δ *ypt52* Δ double mutant. Vps21 and Ypt52, therefore, have both distinct and complementary activities (10,11). A *vps9* Δ mutant exhibited a less severe CPY sorting defect than a *vps21* Δ *ypt52* Δ double mutant, suggesting that residual Rab5 signaling still occurs in the absence of Vps9-catalyzed nucleotide exchange.

Rab5 signaling is required for MVB biogenesis and sorting

Previous work suggested that Vps21 operates upstream of the MVB to accept incoming endosomes and Golgi-derived vesicles (12,24–26). In cells lacking the Vps9 GEF, missorting of the endocytic MVB cargo Ste3 to cytoplasmic punctae further suggested a Rab5 requirement in MVB cargo sorting or biogenesis (27). Hence, we were surprised to observe correct sorting of both an ubiquitin-independent cargo, Sna3-GFP (Figure 2A) and an ubiquitin-dependent MVB cargo, GFP-CPS (carboxypeptidase S; unpublished results) to the vacuole lumen in $vps21\Delta$ or $ypt52\Delta$ single mutant cells. The persistence of cargo sorting in Rab5 single mutants is supported by TEM (transmission electron microscopy) observations of morphologically normal MVBs in $vps21\Delta$ cells, albeit at reduced frequency versus the wild-type (28).

In contrast to Rab5 single mutants, $vps21\Delta$ $vpt52\Delta$ double mutant cells strongly missorted both Sna3-GFP (Figure 2A) and GFP-CPS (Figure 2B). In the double mutant cells Sna3-GFP was present throughout the cytoplasm and occasionally (<10% of cells) at large, bright FM 4-64-positive punctae near the vacuole (Figure 2A, closed arrowheads). By light microscopy, this phenotype resembled the class E phenotype of cells with defective ESCRT function (e.g. $vps4\Delta$, Figure 2A). However, by TEM $vps21\Delta$ $ypt52\Delta$ double mutants were devoid of both MVBs (compare to wild-type, Figure 2C) and stacked class E compartments. Instead, $vps21\Delta$ $ypt52\Delta$ cells occasionally accumulated unstacked electron-dense structures (Figure 2E-G), both in the cytoplasm (Figure 2G) and within the vacuole limiting membrane (Figure 2F). These electron-dense structures were not observed in wild-type cells or in $vps21\Delta$ single mutants. The perivacuolar and intravacuolar locations of the electron-dense structures suggest that they correspond to Sna3-GFP and FM 4-64 positive structures observed by light microscopy (Figure 2A), as described below. These results suggest that MVB biogenesis is largely or completely abrogated in the double mutant cells.

Because ILV cargoes are missorted and ILVs fail to form in $vps21\Delta$ ypt52 Δ double mutants, we asked whether ESCRT localization is perturbed in these cells. ESCRT recruitment to endosomes depends largely on binding



Figure 2: Effects of Rab5 pathway defects on MVB cargo sorting and biogenesis. A–B) Fluorescence micrographs show Sna3-GFP (A) or GFP-CPS (B) and FM 4-64, an endocytic tracer that marks the vacuole, in cells with genotypes as indicated. Open arrowhead, perivacuolar colocalization of GFP and FM 4-64 at class E compartment puncta. Closed arrowhead, colocalization of GFP and FM 4-64 at puncta that appear to be inside the vacuole limiting membrane. Scale bars, 2 m. C–G) Thin-section electron micrographs of wild-type (C), $vps4\Delta$ (D) and $vps21\Delta$ $ypt52\Delta$ (E–G) cells, including electron-dense structures both within (arrows) and outside (arrowheads) the vacuole limiting membrane in $vps21\Delta$ $ypt52\Delta$ cells. F–G) Increased magnification of highlighted regions in (E). Scale bars span 100 nm. Vac, vacuole.

of Vps27 (ESCRT-0) to PI(3)P (phosphatidylinositol-3phosphate) (29). PI(3)P is on endosomes and required for efficient ILV formation (30). PI(3)P localization was probed with a GFP-FYVE (Fab1, YOTB, Vac1 and EEA1) probe. The endocytic membrane tracer FM 4-64 was used (without a chase step) to label both endocytic organelles and the vacuole (Figure 3A). In wild-type cells GFP-FYVE localized efficiently to FM 4-64-stained perivacuolar punctae as well as to the vacuole limiting membrane, indicating the presence of PI(3)P in cytoplasmic leaflets of both endosomal and vacuolar membranes (31,32). Cells lacking $vps4\Delta$ accumulated PI(3)P at FM 4-64-stained class E compartments. In striking contrast to both wild type and class E mutant cells, both $vps21\Delta$ and $vps21\Delta$ ypt52∆ mutants lacked discrete cytoplasmic GFP-FYVE punctae, and exhibited strong GFP-FYVE accumulation on the vacuole limiting membrane. The accumulation of

Traffic 2012; 13: 1411-1428

GFP-FYVE at the vacuole limiting membrane in *vps21* Δ and *vps21* Δ *ypt52* Δ cells argues against a global defect in PI(3)P synthesis and suggests the presence of an endosome-specific defect. Consistent with this finding, mammalian endosomal PI-3-kinase is stimulated by Rab5 signaling (33). Thus, in *vps21* Δ single and *vps21* Δ *ypt52* Δ double mutants a key signal for ILV formation is depleted from prevacuolar endosomal compartments and mislocalized to the terminal vacuole.

ILV scission involves ESCRT-III assembly on endosomal membranes followed by Vps4-catalyzed ESCRT-III disassembly (16). We therefore assessed the localization of Did2, perhaps the only ESCRT-III subunit that retains function when fused to GFP (34,35). Endosomal GFP-Did2 punctae were abundant in wild-type cells, while in $vps4\Delta$ cells GFP-Did2 concentrated at large class E punctae



Figure 3: Rab5 signaling is required for PI(3)P and ESCRT-III localization at endosomes. A–B) Confocal fluorescence micrographs show PI(3)P probe GFP-EEA1_{FYVE} ('GFP-FYVE') (A) or ESCRT-III component GFP-Did2 (B) in cells pulsed with endocytic tracer dye FM 4-64 for 30 min without chase. Scale bars, 2 m. C) Box plot quantification of observed GFP-Did2 fluorescent foci. 130–150 cell profiles from three independent experiments.

(Figure 3B) (34,35). $vps21\Delta$ $ypt52\Delta$ double mutants had few or no GFP-Did2 punctae (Figure 3B,C) despite increased FM 4-64 stain in the cytoplasm relative to wild-type, indicating delayed FM 4-64 traffic through endocytic compartments. $vps21\Delta$ single mutants had an intermediate phenotype with fewer GFP-Did2 punctae relative to wild-type (Figure 3C), consistent with the reduced frequency of endosomal luminal vesicles in $vps21\Delta$ cells (28). We emphasize that although Rab5 deficiency delays FM 4-64 trafficking to the vacuole the endocytic marker dye does reach the vacuole, indicating that at least a subset of endocytic vesicles retain a vacuolar itinerary. Together, these data support working models in which Rab5-deficient cells have endosomes, but these organelles do not support proper ESCRT activity and cannot properly form ILVs.

Rab5 Specificity and Regulation in Yeast



Figure 4: LUCID assay of MVB cargo missorting in Rab5 mutants. A) Schematic diagrams show predicted Sna3-FLuc sorting itineraries in wild-type and mutant cells. B–D) Validation of the LUCID assay. B) Effect of Sna3^{Y109A}-FLuc mutation. C) Partial rescue of early ESCRT cargo missorting defect by overexpression of ESCRT-II. D) Mutants with escalating degrees of disrupted ESCRT-III dynamics and cargo missorting display escalating degrees of Sna3-FLuc signal. E) MVB cargo missorting phenotypes in Rab signaling-deficient mutants. In panels (B–D), bars show mean \pm SD of triplicate samples from representative experiments. In panel (E) bars show mean \pm SEM of four independent experiments (except *ypt7* Δ , *n*=3). RLuc, Renilla luciferase. MVB, multivesicular body. 2 , yeast high-copy plasmid.

LUCID: quantitative assessment of Rab5 function in MVB sorting

The above experiments suggested that MVB formation and MVB cargo sorting are profoundly compromised in *vps21*Δ *ypt52*Δ mutant cells. To assess the severity of this defect, we sought more quantitative methods. Existing assays of ILV cargo sorting have significant limitations. Localization of GFP-cargo fusions to the vacuole lumen versus the limiting membrane is not readily quantifiable, and while pulse-chase analyses of cargo maturation resolve subtle differences in rates of transport to the vacuole/lysosome, the proteolytic maturation of MVB cargos does not necessarily result from correct sorting into lumenal vesicles (36). To address some of the limitations of existing assays, we constructed a reporter system called LUCID (*luc*iferase reporter of *i*ntraluminal *d*eposition).

The LUCID vector has two components. The reporter component encodes a chimeric fusion of firefly luciferase to

Traffic 2012; 13: 1411-1428

the MVB cargo Sna3 (Sna3-FLuc; Figure 4A). Sna3-FLuc is readily guantified using luciferase activity assays. Correct targeting of Sna3-FLuc to ILVs results in sequestration and eventual destruction of the FLuc reporter. Because the FLuc tag faces the cytoplasm, defects in Sna3-FLuc traffic to the MVB, or failure to package Sna3-FLuc into ILVs, result in the accumulation of active Sna3-Fluc. To enable ratiometric correction for variations in cell number, plasmid copy number, protein synthesis rate, or global protein turnover, the LUCID vector's normalization component encodes a soluble cytoplasmic *Renilla* luciferase (RLuc) (37,38). Sna3-FLuc and RLuc are sequentially assayed from each cell lysate using a microplate luminometer that incorporates robotic solution injectors. In pilot experiments, we found that signal-to-background is optimal when cells are lysed after a 20-30 min cycloheximide chase. The chase blocks new protein synthesis and allows nascent Sna3-FLuc to reach the lumen, resulting in lower FLuc activity in wild-type cells, while mutants defective for MVB targeting exhibit relatively higher FLuc activity.

Sna3 is sorted into ILVs through both ubiquitin-dependent and -independent mechanisms (39). To validate the LUCID assay we tested a mutant (Sna3^{Y109A}) that is defective in both mechanisms and inefficiently sorted to ILVs (40). Sna3^{Y109A}-FLuc accumulated to levels at least 10 times higher than the wild-type Sna3-FLuc reporter (Figure 4B). The signal-to-background ratio of LUCID therefore exceeds 10:1. Importantly, LUCID reliably reports intermediate defects in MVB cargo sorting. Deletion of the ESCRT-I subunit Vps23 substantially increased Sna3-Fluc activity, while overproduction of the ESCRT-II holocomplex partially suppressed the $vps23\Delta$ defect (Figure 4C), consistent with work using other assays (41). Previous work connected defects in disassembly or remodeling of ESCRT-III oligomers to proportional defects in MVB cargo sorting (34,35,42,43). We observed a corresponding phenotypic progression of signal intensity (Figure 4D) in mutants with compromised ESCRT-III dynamics, ranging from partial (vta1 Δ ; did2 Δ) to complete (vps4 Δ) defects.

LUCID experiments with Rab5 mutant cells (Figure 4E) revealed moderate MVB sorting defects in the $vps21\Delta$ and $vps9\Delta$ single mutants and severe defects in the $vps21\Delta$ ypt52 Δ double mutant, with missorting at levels comparable to strong vps class E mutants that fail to form lumenal vesicles (e.g. $vps4\Delta$). Among the Rab5 signaling-deficient mutants, the relative severities of MVB missorting (Figure 4E) correlated with the pattern seen in analyses of CPY secretion (Figure 1B). Hence, we conclude that both formation of morphologically normal MVBs (Figure 2) and sorting of MVB cargos to ILVs (Figures 2A, B and 4E) exhibit strong dependence on Rab5 signaling.

Rab5 signaling is required for efficient autophagy

Cells lacking VPS21 and YPT52 sometimes accumulated endocytic markers at foci near or within the vacuole (Figure 2), and we wondered whether these structures might mark attempts to clean up stalled endocytic traffic through autophagy. Atg8 marks nascent and mature autophagic membranes (44,45). Upon fusion of autophagosomes with the vacuole, some Atg8 is recycled to the cytosol and some is deposited in the vacuole lumen. While intravacuolar membrane accumulations appeared infrequently in $vps21\Delta$ $ypt52\Delta$ mutants (estimation <10%), we observed colocalization of GFP-Atg8 with intravacuolar foci of both FM 4-64 (Figure S2A) and the MVB cargo mCherry-CPS (Figure S2B), in cells both with and without diffuse GFP-Atg8 signal in the vacuole lumen. In contrast, GFP-Atg8 did not colocalize with FM 4-64stained class E compartments in $vps4\Delta$ cells (Figure S2A). Thus, a key autophagic component colocalizes with aberrant endosomal structures in $vps21\Delta$ ypt52 Δ cells.

Recent genetic studies indicate that traffic through the Golgi-endosome system is crucial for CVT vesicle and autophagosome assembly (46). We therefore asked whether autophagy is globally up-regulated, unaffected,

or defective in Rab5-deficient cells. Proteolytic maturation of vacuolar aminopetidase I (Ape1) is a hallmark of constitutive cytosol-to-vacuole transport (CVT) and macroautophagy (44). Unstressed vps4 Δ and vps21 Δ cells contained precursor and mature Ape1 at approximately wild-type levels (Figure S2C, top panel). In marked contrast, unstressed vps21 Δ ypt52 Δ cells exhibited little or no maturation of Ape1, indicating a penetrant defect in CVT transport.

Rapamycin stimulates autophagy by inhibiting TOR (Target of rapamycin) kinase, resulting in dephosphorylation of the TOR substrate Atg13 (47). Wild-type, $vps4\Delta$, $vps21\Delta$ and $vps21\Delta$ $ypt52\Delta$ cells all contained similar levels of slow-migrating phospho-Atg13, and in all cases rapamycin treatment depleted phospho-Atg13 with comparable kinetics (Figure S2D). Thus, TORC1 signaling to the autophagy machinery is not markedly perturbed in Rab5-deficient cells. Nevertheless, rapamycin treatment only partially restored Ape1 maturation in $vps21\Delta$ $ypt52\Delta$ cells (Figure S2C, bottom panel). It is important to note that vacuoles in $vps21\Delta$ $ypt52\Delta$ cells are proteolytically competent (10). Thus, the accumulation of precursor Ape1 indicates that $vps21\Delta$ $ypt52\Delta$ mutants have significant defects in CVT and macroautophagy.

Rab5 signaling is required for tolerance of ionic stress

Many endolysosomal trafficking mutants are sensitive to salt and osmotic stress, and numerous endocytic trafficking genes were shown to be genetic modifiers of Ca^{2+} influx (48). To see whether ionic sensitivity could be used to probe the *in vivo* requirement for Rab5 signaling, we assaved the growth of mutants defective in Rab5 on media containing various concentrations of extracellular Ca²⁺. Mutants lacking any single Rab5 ortholog grew as well as the wild-type at up to 400 mM Ca²⁺ (Figure S3A,B; Table 1). In contrast, $vps21\Delta$ $ypt53\Delta$ double mutant cells had strong synthetic sensitivity to high Ca²⁺ and were somewhat more Ca²⁺-sensitive than cells lacking the Vps9 GEF. $vps21\Delta$ $ypt52\Delta$ double mutants were also hypersensitive to extracellular Ca²⁺, as were cells lacking the Rab5 effectors Vps3 and Vac1. Deletion of a third effector, Vps8, had little effect on Ca²⁺ tolerance. Triple mutants lacking all three Rab5 orthologs (vps21A $ypt52\Delta ypt53\Delta$) were hypersensitive to Ca²⁺, even more so than cells lacking Crz1, a key calcineurin-responsive transcription factor (49). The triple mutants grew poorly even in low extracellular [Ca²⁺] (Figure S3A; Table 1), again demonstrating a functional role for Ypt53 in stress tolerance. To our knowledge, these are the first strong phenotypes found to be associated with ypt53 deletion.

Ypt53 is a stress-induced functional homolog of Vps21

 $ypt53\Delta$ single mutants lack obvious growth or trafficking phenotypes (10) and *YPT53* has evaded identification in forward genetic screens. *YPT53* is transcribed at low levels in unstressed cells, while *VPS21* and *YPT52*

Table 1: Growth rates of Rab5 family mutants under calcium stress. Inflection doubling time (min), mean \pm SD, n=5 each.

Genotype	YPD	$YPD+200\;mM\;CaCl_2$
Wild-type	90.3 ± 7.2	117.0 ± 5.2
$vps21\Delta$	83.3 ± 4.7	114.0 ± 2.8
ypt52 Δ	83.1 ± 3.2	107.5 ± 5.6
ypt53 Δ	92.7 ± 2.7	114.1 ± 4.2
vps21 Δ ypt52 Δ	89.9 ± 7.0	255.0 ± 22.6
vps21 Δ ypt53 Δ	82.6 ± 6.8	170.3 ± 11.6
ypt52 Δ ypt53 Δ	99.4 ± 4.5	122.9 ± 5.5
vps21 Δ ypt52 Δ ypt53 Δ	122.1 ± 9.4	277.1 ± 50.2
vps9Δ	94.7 ± 3.4	152.9 ± 10.8

transcripts are considerably more abundant (50). However, *YPT53* was identified in microarray experiments as a target of the calcineurin stress response pathway (49). Using a luciferase reporter to measure transcription from the *YPT53* promoter, we verified that *YPT53* exhibits dose-dependent induction in elevated extracellular Ca²⁺ (Figure 5A). Moreover, *YPT53* transcription was strongly induced in Rab5-deficient mutants, even when grown in standard media (Figure 5B). *YPT53* induction by either extracellular Ca²⁺ or Rab5 deficiency was abolished in mutants lacking the calcineurin-regulated transcription factor Crz1, indicating that responses to both stresses converge on the calcineurin-Crz1 pathway (Figure 5A,B).

Synthetic phenotypes observed in *vps21 ypt53* double mutants (Figure S3; Table 1) raised the possibility that the Vps21 and Ypt53 proteins are functional homologs, but differentially expressed. To test this hypothesis, we expressed *YPT53* from the *VPS21* promoter. *VPS21*_{PR-}*YPT53* efficiently suppressed loss of Vps21 alone or Vps21 and Ypt52 together (Figure 5C–E). It restored sorting of CPY and of MVB cargo (Figure 5C,E) and almost completely restored growth in the presence of high extracellular Ca²⁺ (Figure 5D).

Roy1 (repressor of *ypt52*) was recently reported to oppose Ypt52 signaling (51). *VPS21pr::YPT53* suppressed CPYinvertase secretion in *vps21* Δ cells more completely than did loss of Roy1 (Figure 5C). *VPS21pr::YPT53* also restored characteristic MVB ultrastructure to *vps21* Δ *ypt52* Δ cells (Figure 5F,G). However, MVBs were now twice as abundant as in wild-type cells (0.82 versus 0.37 MVB per cell; n=50 cell profiles each), and more often appeared tethered or clustered (<30 nm apart) than in wild-type cells (71% clustered, n=41 MVBs versus 18% clustered, n=40 MVBs). Thus, when expressed from the same promoter, the Vps21 and Ypt53 proteins are largely but not totally interchangeable.

Gyp3 opposes Rab5 signaling

We next sought to identify additional regulators of Rab5 signaling through genetic tests. Cells lacking the Rab5 GEF Vps9 had less severe trafficking and growth defects than $vps21\Delta$ ypt52 Δ double mutants, implying that $vps9\Delta$

cells contain a reduced but functionally significant pool of active Rab5. We reasoned that the removal of Rab5 inhibitors might enrich the pool of activated Rab5, and thereby suppress the defects of $vps9\Delta$ mutants. Thus, we deleted each of the eight known *S. cerevisiae* Rab GAPs in a $vps9\Delta$ genetic background and tested the resulting double mutants for growth on high Ca²⁺ media. The loss of two GAPs, Msb3/Gyp3 and Gyp7, partially suppressed the Ca²⁺ sensitivity of $vps9\Delta$ cells (Figure 6A).

Gyp7 is the major in vivo GAP of Ypt7, the yeast Rab7 ortholog (52). Suppression of $vps9\Delta$ by $gyp7\Delta$ is probably not due to a direct effect of Gyp7 on Rab5 signaling, as discussed below. At the plasma membrane, Gyp3 functions redundantly with Msb4/Gyp4 to regulate delivery of secretory vesicles by targeting the Sec4 Rab (53). As reported previously (20) and as we confirm here, Gyp3 has strong GAP activity against Vps21. To verify suppression of the $vps9\Delta$ growth defect by $gyp3\Delta$, we performed a competition experiment. Double mutant $vps9\Delta$ $gyp3\Delta$ cells had increased competitive fitness versus $vps9\Delta$ cells in Ca²⁺-rich media (Figure 6B), but there was not a fitness difference under standard (control) growth conditions. We also found that $gyp4\Delta$ deletion, in contrast to $qvp3\Delta$, did not suppress the $vps9\Delta$ mutant's Ca²⁺ sensitivity (Figure 6A). Thus, GYP3 and GYP4 function redundantly with respect to Sec4 (53), but only GYP3 interacts genetically with VPS9 and, presumably, Rab5 signaling.

To test the hypothesis that $gyp3\Delta$ suppression of the $vps9\Delta$ mutant's Ca²⁺-sensitivity depends on Vps21, we constructed a $vps9\Delta$ $gyp3\Delta$ $vps21\Delta$ triple mutant. The inability of this mutant to grow in the presence of elevated extracellular [Ca²⁺] demonstrated that $gyp3\Delta$ suppression of the $vps9\Delta$ growth defect requires an intact copy of VPS21. To test whether $gyp3\Delta$ suppresses trafficking defects caused by the $vps9\Delta$ mutation, we assayed cargo sorting. $vps9\Delta$ $gyp3\Delta$ double mutants secreted less CPY-invertase than $vps9\Delta$ single mutants (Figure 6C) and in LUCID assays the double mutants accumulated significantly less of the MVB cargo Sna3-FLuc (Figure 6D). Thus, we conclude that GYP3 functionally opposes Rab5 signaling *in vivo*.

Vps21 is a preferred target of the Gyp3 GAP

The above experiments led us to predict that one or more of the yeast Rab5 proteins should be a direct target of the Gyp3 GAP activity. Using purified proteins, we assayed the GAP activities of the Gyp1_{TBC} catalytic core, full-length Gyp3, and full-length Gyp7 against a panel of endolyso-somal Rabs (Figure 7). Single-turnover kinetics of GTP hydrolysis were measured using a real-time assay of inorganic phosphate evolution, with each GAP assayed across a range of concentrations. The results were analyzed by nonlinear fitting of a pseudo first-order Michaelis–Menten model, which allowed us to estimate the intrinsic hydrolysis rate for each Rab and the specificity constant (K_{cat}/K_{M}) for each Rab-GAP combination (Figure 7C). As



Figure 5: Ypt53 is a stress-induced Vps21 paralog. A) Transcription from the YPT53 promoter was monitored using FLuc transcriptional fusions in wild-type or *crz1* Δ cells grown in medium supplemented with different concentrations of Ca²⁺. Bars shown mean \pm SD of duplicate samples from a representative experiment. B) YPT53 expression is induced in cells deficient in Rab5 signaling. Bars show mean \pm SEM from independent experiments (n=5, 5, 4, 5, 3, 4 and 4, respectively). C–G) YPT53 driven by the VPS21 promoter rescues CPY trafficking (C), calcium growth tolerance (D), ILV cargo trafficking (E) and ILV biogenesis (F–G) in *vps21* Δ *ypt52* Δ mutants. C) Assay of extracellular CPY-invertase, normalized to the wild type. Bars indicate mean \pm SD of triplicate samples from a representative experiment. D) Growth curves in liquid culture, 30°C. Mean, n=4. E) Fluorescence microscopy of FM 4-64-stained cells expressing GFP-CPS (carboxypeptidase S). Scale bar, 2 m. F–G) Thin-section electron micrographs show restored MVB morphology. Vac, vacuole. Arrowheads, individual MVBs. Bars span 100 nm.

reported previously (5) Gyp1_{TBC} is relatively promiscuous and efficiently accelerated GTP hydrolysis by all four Rabs tested. Compared to Gyp1_{TBC}, Gyp7 and Gyp3 were more selective. Consistent with previous *in vitro* assays (5), cell-free fusion experiments (52,54) and multiple lines of *in vivo* evidence (52), Gyp7 strongly accelerated hydrolysis on Ypt7 but had low or undetectable activity against the Rab5 paralogs. Gyp3 had robust GAP activity against Vps21 and low but reproducible activity against Ypt53 and Ypt7. These results verify and extend previous work



Figure 6: Rab GAP Msb3/Gyp3 opposes endosomal Rab5 signaling. A) Limiting dilution plate growth assay, 30° C on YPD \pm 200 mM CaCl₂. B) *vps9* Δ *gyp3* Δ cells outcompete *vps9* Δ cells in mixed cultures. Bars indicate mean \pm SD, *n*=2. C) Assay of extracellular CPY-invertase, normalized to the wild-type. Mean \pm SD of four samples per point from a representative experiment. D) LUCID assay of ILV sorting. Bars indicate mean \pm SEM from six independent experiments; * p=0.024, two-tailed, paired *t*-test.

(5,20,52,54) and, together with our genetic experiments, support the hypothesis that Gyp3 operates *in vivo* as the major Vps21 GAP. Moreover, the observed GAP activities of Gyp3 and Gyp7 against Ypt7 are consistent with *in vivo* fragmentation of vacuoles when either *GYP3* or *GYP7* is overproduced, as previously reported by our group (52).

Gyp3 enforces the endolysosomal boundary

Rab GAPs are thought to regulate the timing and levels of Rab signaling, and also to police the spatial boundaries of Rab activity (2,3). During endolyosomal Rab conversion in mammals, Rab5 dissociates from late endosomes and is replaced by Rab7, a process that requires Mon1/SAND and the Vps3 ortholog hVam6 (14,17,19,55).

The Rab5 paralog Vps21 typically appears on cytoplasmic puncta, often adjacent to FM 4-64-stained vacuoles (Figure 8A), while downstream vacuoles are marked by the Rab7 homolog Ypt7. In $gyp3\Delta$ mutant cells we observed dramatic mislocalization of Vps21 to the

Traffic 2012; 13: 1411–1428

vacuole limiting membrane. In mutants lacking each of the other known Rab GAPs, Vps21 remained on endosomal punctate structures and was excluded from the vacuole (Figure 8B), indicating a highly selective role for Gyp3 in Vps21 inactivation. These results suggested a working model in which Vps21 is first activated on endosomes by Vps9 and then inactivated by Gyp3 during late endocytic Rab conversion. In the absence of Gyp3, Vps21 remains active through endosome fusion at the vacuole, resulting in steady-state Vps21 accumulation on the vacuole limiting membrane.

Two GFP-tagged Vps21 effectors, the CORVET subunits Vps3 and Vps8 (Figure S4) exhibited apparently normal localization to prevacuolar endosomes in $gyp3\Delta$ cells, indicating that CORVET localization is specified not only by the location of Vps21 but also by other as-yet unidentified cues (see Discussion). In $gyp3\Delta$ cells, the GFP-Vps21 perivacuolar punctae appeared brighter than in wild-type cells, somewhat reminiscent of class E compartments or clustered endosomes when viewed by fluorescence



Figure 7: Msb3/Gyp3 is a Vps21 GAP in vitro. A) Time course of solution-based GTP hydrolysis by Vps21, Ypt52, Ypt53 or Ypt7 in the presence of increasing concentrations of Gyp3, Gyp7 or Gyp1TBC. The data shown are representative of three independent experiments. B) Purified proteins were separated by polyacrylamide gel electrophoresis (2.5 g per lane) and stained with Coomassie brilliant blue. C and D) Kinetics curves were fitted to a pseudo-first-order Michaelis–Menten model function to obtain K_{cat}/K_{M} of a particular Rab-GAP pairing and the intrinsic GTP hydrolysis rate of each Rab. The plots represent the fitted mean ± SEM rate constants obtained from duplicate experiments.

microscopy. Ultrastructural examination of $gyp3\Delta$ cells by TEM revealed morphologically normal, unclustered MVBs (Figure S1), indicating that the Vps21 mislocalization in $gyp3\Delta$ cells does not have obvious morphological consequences. Similarly, $gyp3\Delta$ mutants exhibited nearnormal sorting of the Golgi-endosome-vacuole cargo CPY (Figure 6C).

The mislocalization of Vps21 in $gyp3\Delta$ cells was verified and further characterized in biochemical fractionation experiments. Loss of a Rab's cognate GAP is predicted to result in enrichment of Rab-GTP relative to Rab-GDP, rendering the Rab resistant to membrane extraction by Rab GDI and increasing the steady-state level of membrane-associated Rab (52). About 50% of the Vps21 in the vacuole-enriched P13 membrane fraction from wildtype cells was vulnerable to extraction in incubations

a comparable fraction from $gyp3\Delta$ cells resisted GDI extraction (Figure 8D, compare lanes 2 and 4). We further predicted that purified Gyp3 should target the GDI-resistant (and presumably GTP-bound) population of Vps21 in membranes from $gyp3\Delta$ cells. To test this prediction, we isolated membranes from $gyp3\Delta$ cells and incubated them with recombinant purified Gyp3 prior to GDI extraction. Gyp3 rapidly converted almost all membrane-bound Vps21 to a GDI-extractable and presumably GDP-bound form (Figure 8D, compare lanes 4 and 5). The most straightforward interpretation of this result is that Gyp3 inactivates Vps21 on intact endolysosomal membranes, rendering Vps21 susceptible to GDI extraction. Gyp3 also enhanced GDI extraction of the vacuolar Rab Ypt7, consistent with its ability to stimulate GTP hydrolysis on Ypt7 in vitro (Figure 7)

with purified GDI; in contrast, most of the Vps21 in



Figure 8: Msb3/Gyp3 inactivates Vps21 and spatially constrains Vps21 activity. A) GFP-Vps21 contaminates the vacuole (marked by the fluorescent tracer FM4-64) in *gyp3* Δ mutants. B) Mislocalization of GFP-Vps21 to the vacuole occurs in cells lacking Gyp3, but not in cells lacking other yeast Rab GAPs. Scale bars, 2 m. C) Subcellular fractionation of wild-type and *gyp3* Δ cells. PGK and V-ATPase (Vph1) were used as fiduciary markers for cytosol and endolysosomal membrane fractions. T, total lysate after 1000 × *g* spin. P15, membrane-associated 15000 × *g* pellet fraction. S15, cytosolic 15000 × *g* soluble fraction. D) Extraction of Rab proteins Vps21 and Ypt7 by Rab GDI. P15 membrane pellets prepared from wild-type or *gyp3* Δ cells were resuspended in lysis buffer or buffer supplemented with 3 μ M His₇-MBP-Gyp3, and incubated for 15 min on ice. Indicated samples received 9 μ M recombinant GDI immediately prior to another 15000 × *g* spin in which all samples were again separated to membrane-bound (P) and soluble fractions (S). Alkaline phosphatase (ALP) and the vacuolar SNARE Vam3 served as fiduciary markers for sedimentation of vacuoles.

and its ability to cause vacuole fragmentation when overexpressed *in vivo* (52). In summary, our experiments show that Gyp3 acts to limit the total pool of activated Vps21 *in vivo*, and that it specifically targets Vps21 at the terminal vacuolar lysosome for inactivation. We conclude that Gyp3 restricts the territory of Vps21 activity to prevacuolar endosomal compartments, thereby enforcing a key aspect of compartmental identity.

Discussion

Our experiments show that Rab5 signaling is required not only for classical endosomal cargo sorting but also for normal MVB biogenesis, the CVT/autophagy pathway and cellular stress tolerance. The requirement for Rab5 signaling in survival of severe Ca²⁺ stress facilitated a suppression analysis that identified *GYP3* as a negative regulator of Rab5 signaling. Further experimentation demonstrated that Gyp3 is a strong and relatively selective Vps21 GAP, and revealed that Gyp3 functions

Traffic 2012; 13: 1411-1428

to spatially restrict Vps21 signaling to pre-vacuolar endosomal compartments. In parallel work, Lachmann and colleagues (56) performed a biochemical survey of Rab GAP selectivity, independently identifying Gyp3 as a Vps21 GAP, and similarly observed mislocalization of Vps21 to the vacuole in $gyp3\Delta$ cells.

The CORVET subunits Vps3 and Vps8 bind Vps21-GTP and are released from membranes in *vps21* Δ cells (57–60). However, in *gyp3* Δ cells Vps3 and Vps8 expressed from native promoters did not follow Vps21 to the vacuole (Figure S4). Overexpression of Vps8 in *gyp3* Δ cells resulted in partial mislocalization of Vps8 to vacuole membranes (56). Taken together, these results suggest that the presence of Rab5 is necessary but not sufficient for CORVET localization. We speculate that additional determinants including phosphinositides and SNAREs operate in concert with Rab5 to recruit CORVET, and indeed we have recently shown that CORVET directly binds the endosomal Qa-SNARE Pep12 (B. Lobingier, Alexey J. Merz, personal communication).

Rab 5 signaling was required not only for ILV cargo sorting (Figures 2 and 4), but for formation of morphologically normal MVBs (Figure 2). ESCRT-0, I, II and III drive these processes, and both ESCRT recruitment (29) and ILV biogenesis (30) are strongly promoted by PI(3)P. In mammals, endosomal PI-3-kinase is stimulated by Rab5 (33). Consistent with these findings, we found that both PI(3)P and ESCRTs are depleted from endosomes in Rab5-deficient yeast cells. The data suggest a working model in which loss of Rab5 signaling leads to loss of PI(3)P from endosomes, preventing efficient recruitment and activity of ESCRTs. In addition, both Rab5 and PI(3)P have important roles in endosomal membrane fusion. A mature MVB contains on the order of 20 times more total membrane than a typical endocytic vesicle or post-Golgi carrier. Thus, the biogenesis of a single mature MVB may require dozens of preceding homotypic and heterotypic membrane docking and fusion events, all orchestrated by Rab5 and PI(3)P signaling. Reduced membrane fusion in $vps21\Delta$ $ypt52\Delta$ double mutant cells might reduce endosome size and directly limit the quantity of membrane available to form ILVs.

Although the three yeast Rab5 proteins are partially redundant, they exhibit some biochemical and functional specialization. In recent work characterizing an intrinsic membrane tethering activity of yeast endosomal Rabs, we detected robust tethering mediated by Vps21 or Ypt53, but none with Ypt52 (61). Similarly, although Gyp3 had robust GAP activity against Vps21 in our assays, Gyp3 activity against Ypt52 and Ypt53 was lower (Figure 7). These results are in good qualitative and quantitative agreement with previous work (5,20). In contrast, an exhaustive assessment of all yeast Rab-GAP pairs (56) found relatively strong stimulation of Ypt52 GTP hydrolysis by Gyp3, and also reported anemic stimulation of Ypt7 GTP hydrolysis by Gyp7, where we and others (5) saw robust stimulation. These differences in apparent in vitro GAP selectivity may reflect divergent assay conditions or differences between time courses versus single endpoint measurements. We emphasize that every study to date has demonstrated strong activity of Gyp3 against Vps21.

Ypt52 is notable for its fast intrinsic hydrolysis rate, which results in hydrolysis of ≥90% of bound GTP within 60 min, without assistance from a GAP (Figure 7) (20). *In vivo*, Ypt52 (but not Vps21 or Ypt53) is selectively bound and inhibited by the F-box containing inhibitor Roy1 (51) (also A. J. M. and R. Plemel, unpublished data). *YPT53* is a stress-inducible gene that encodes a functional homolog of Vps21 (Figure 5). The relative immunity of Ypt53 to Gyp3 may account for increased tethering of MVBs when *YPT53* is expressed from the *VPS21* promoter in a *vps21∆ ypt52∆* genetic background (Figure 5F,G). Given its immunity to Gyp3, Ypt53 might also extend the spatial domain of active Rab5 signaling to the vacuole. Consistent with this hypothesis, overproduced Ypt53 localizes to both endosome and vacuole membranes (62).

In the Rab countercurrent model of GAP recruitment (2), an activated Rab recruits the GAP for the preceding upstream Rab. In this way, activation of the downstream Rab triggers inactivation of the upstream Rab, and sharp boundaries between Rab signaling domains are self-organized. In support of this idea, elegant work with Caenorhabditis elegans demonstrated that Rab7 is needed to position a Rab5 GAP, TBC-2 (63). In yeast, the situation is less clear. Gyp3 inhibits both endosomal and vacuolar Rabs, Vps21 and Ypt7 (Figure 7) (20,52,56). However, Gyp3 is considerably more active against Vps21 than it is toward Ypt7 (Figure 7). Thus, it is still possible that Ypt7 positions Gyp3 to inactivate Vps21 on the vacuole. Further work will be needed to evaluate this model. Alternatively, other factors may regulate Gyp3 localization. Rho family small G proteins control Gyp3 and Gyp4 recruitment to sites of polarized exocytosis (64). Rho1 and Cdc42 are also found on the vacuole, where they influence actin dynamics and membrane fusion (65,66). Thus, endolyosomal compartments and secretory patches at the plasma membrane might share common mechanisms for directing the localization and activity of Gyp3, and excluding Vps21, at the initial and terminal boundaries of the endocytic pathway. These models provide an obvious agenda for future experimentation. Note added in proof: As this manuscript was nearing completion, Zerial et al. reported knockdown of all three mammalian Rab5 isoforms in the livers of intact animals. The endocytic system was essentially absent in cells lacking Rab5 (67).

Materials and Methods

Strain and plasmid construction

Strains and plasmids are summarized in Table 2. Expression vector pDN616 was made by PCR amplification of the *CEN/ARSH4* locus from pRS415, using primers incorporating flanking, parallel LoxP sequences. Gap repair of AatII-digested pRS406 vector using the resulting PCR product yielded a low-copy replicative plasmid similar to pRS416, but with the ability to readily convert to a non-replicative, integrating plasmid after excision of the *CEN* locus by Cre recombinase. pDN526 was made in manner similar to pDN616, except that gap repair inserted the 2 locus PCR-amplified from pRS425 using primers that encoded a trio of flanking restriction sites (AatII-AvrII-SphI).

pDN224 was constructed by using sequence overlap extension (SOE)-PCR to fuse the *VPS21* promoter to the *YPT53* coding sequence and terminator using primers that also inserted a six-codon N-terminal FLAG epitope, followed by gap repair of Pvull-digested pDN616. pDN224 was passaged through NS2114Sm bacteria expressing Cre recombinase, removing the *CEN* locus to yield the integrating plasmid pDN225A. DNY512 and DNY515 were constructed by integrating PstI-digested pDN225A at the *ura3-52* locus in DNY471 and DNY438, respectively.

High-copy dual luciferase vector pDN228 was constructed by sequential gap repair of pDN526 using PCR products derived from plasmid template pVW31 (38) *PGK1pr::RLuc::GCY1term* and *FLuc::CYC1term* (lacking promoter) inserted at Xhol and Sacl cut sites, respectively. Promoter sequence (500 bp upstream plus 3 native codons) of *YPT53* was PCR-amplified from genomic DNA and inserted by gap repair into SaclI-digested pDN228, introducing promoters and start codons in-frame with *FLuc* to yield pDN247.

Table 2:	Strains and	plasmids	used in	this study
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Name	Genotype	Reference/source
S. cerevisiae		(00)
SEY6210	NIA Τα Teu2-3, TT2 Ura3-52 hts3-200 trp 1-901 Tys2-801 suc2-9	(68)
BHY10	SEY6210 CPY-Invertase::LEU2 (pBHY11)	(12)
GUYZ3	SEY6210 pro1A::LEU2 pep4A::LEU2	(69)
	SEY6210 VpS4A::////S2	(70)
	SETUZIU V μ SZ3 Δ π IS3	(71)
DINY 38	BHYTU dla2A::HIS3	(72)
		(72) This work
DN1404		This work
	PHV10 vps21A.vAN	This work
DNV433	$BHY10 \ vp521\Delta 0$	This work
	BHY10 ypt52AU BHY10 ypt52AUKAN	This work
DNY/71	$BHY10 vp(33\Deltavar)$	This work
DNY449	BHY10 $vp52\Lambda 0$ $vpt53\Lambda$ ··KAN	This work
DNY511	BHY10 v_{PS} 21 Λ ··· $IIBA3$ v_{PT} to S^{A} ··· pat^{R}	This work
DNY532	BHY10 vps21 Δ 0 vpt53 Δ · · · pat ^R	This work
DNY472	BHY10 v_{PS2} / $\Delta 0$ v_{PS2} / $\Delta 0$ v_{PS2} / $\Delta 0$ v_{PS2}	This work
DNY536	BHY10 vps21 Λ 0 vpt52 Λ 0 crz1 Λ ··K Δ N	This work
DNY441	BHY10 vp/2AKAN	This work
DNY347	BHY10 vac1A "TRP1	This work
DNY349	BHY10 vps3A ···HIS3	This work
DNY350	BHY10 $v_{DS}45\Lambda$ ··H/S3	This work
DNY462	BHY10 $v_{DS}8A$::KAN	This work
GOY223	BHY10 v_{DS} SA::HIS3	G Odorizzi (U Colorado)
DNY479	BHY10 v_{D} s9 Λ ::HIS3 $av_{D}1\Lambda$::KAN	This work
JMY22	BHY10 v_{P} so $HIS3 av_{P} \Delta ::nat^{R}$	This work
DNY494	BHY10 $vps9\Delta$::HIS3 $avp3\Delta$::KAN	This work
DNY495	BHY10 $vps9\Delta$::HIS3 $qvp4\Delta$::KAN	This work
DNY522	BHY10 $vps9\Delta$::HIS3 $gyp5\Delta$::KAN	This work
DNY496	BHY10 $vps9\Delta$::HIS3 $gyp6\Delta$::KAN	This work
DNY524	BHY10 $vps9\Delta$::HIS3 $gyp7\Delta$::KAN	This work
DNY521	BHY10 vps9A::HIS3 gyp8A::nat ^R	This work
DNY525	BHY10 vps9A::HIS3 roy1A::KAN	This work
DNY520	BHY10 <i>vps9</i> Δ:: <i>HIS3 vps21</i> Δ:: <i>URA3</i>	This work
DNY519	BHY10 vps9A::HIS3 gyp3A::KAN vps21A::URA3	This work
NSY1	SEY6210 VPS8-GFP::HIS3	G Odorizzi (U Colorado)
DNY523	NSY1 <i>gyp3∆::KAN</i>	This work
DNY239	BHY10 VPS3-GFP::KAN	This work
DNY539	BHY10 <i>VPS3-GFP::KAN gyp3∆::nat^R</i>	This work
DNY364	SEY6210 gyp1∆::KAN	This work
DNY377	SEY6210 gyp2∆::KAN	This work
DNY489	SEY6210 <i>gyp3∆::KAN</i>	This work
DNY490	SEY6210 gyp4∆::KAN	This work
DNY491	SEY6210 gyp5∆::KAN	This work
DNY504	SEY6210 gyp6∆::KAN	This work
DNY373	SEY6210 gyp7A::HIS3	This work
DNY366	SEY6210 gyp8A::KAN	This work
DNY503	SEY6210 gy/1A::KAN	This work
JMY16	BHY10 gyp3A::KAN	This work
DNY533	BHY10 gyp/A::KAN	This work
DNY531	GUY23 gyp3A::KAN	
DNY514	BHY10 vps21 \u00ed 0 vp321 \u00ed vp321 \u00	This work
DNY515	BHY10 $vps21\Delta UVPS21pr::Flag-YP153$ (pDN225A::URA3)	
	вні ти <i>vps21Δ0 урі52Δ0 vPS21pr::Fiag-YP153</i> (pDN225A::UKA3)	
	SETOZIU VPSZIA::KAN	G Udorizzi (U Colorado)
E. COII	Ellool In10/totBil more Almer had PLAC more PC/100/Z+NA4E +1VZA - D	In stars and a
IUPIUF	F[IdCI] = IIII U(IeU)] IIICIA Δ(IIII-IISUΠΙVIO-MCIBU)φ80/aCZ ΔIVI I 5 Δ/aCX / 4 deoK	invitrogen
BL21(DE3)	F^{-} omp gal dcm lon hsdS _B (r _B m _B) λ (DE3 [lacl lacUV5-T7 gene 1 ind1 sam7	Stratagene/Agilent
NC01140		(70)
NS2114Sm	F^{-} reca λ -cre rpsL	(73)

Traffic 2012; 13: 1411–1428

Table 2: Continued

Name	Genotype	Reference/source
Plasmids		
pRS416	URA3 CEN/ARSH4 Amp ^R	(74)
pRS406	URA3 Amp ^R	(74)
pDN616	URA3 LoxP::CEN/ARSH4::LoxP Amp ^R	This work
pDN606	URA3 LoxP Amp ^R	This work
pDN526	URA3 2μ Amp ^R	This work
pDN224	<i>VPS21pr::Flag-YPT53</i> (pDN616)	This work
pDN225A	VPS21pr::Flag-YPT53 (pDN606)	This work
pGO45	URA3 2µ Amp ^R GFP-CPS1	(69)
pGO47	TRP1 2μ Amp ^R GFP-CPS1	(69)
pSna3-GFP	URA3 CEN Amp ^R SNA3-GFP	(39)
pRP1	pRSF <i>kan^R His₇-MBP-(tev)</i>	This work
pDN190	pRSF <i>kan^R His₇-MBP-(tev) URA3 CEN</i>	This work
pDN193	His ₇ -MBP-(tev)-GYP7 (pDN190)	This work
pDN248	His ₇ -MBP-(tev)-Flag-GYP3 (pDN190)	This work
pMB175	LEU2 Amp ^R 2µ. VPS22 VPS25 VPS36	(41)
pVW31	URA3 2µ Amp ^R PGK1pr::RLuc GCN4pr::FLuc	(38)
pDN228	URA3 2µ PGK1pr::RLuc 0::FLuc	This work
pDN247	PGK1pr::RLuc YPT53pr::FLuc (pDN228)	This work
pDN251	URA3 CEN PGK1pr::RLuc 0::FLuc	This work
pDN252	PGK1pr::RLuc SNA3-FLuc (pDN251)	This work
pMM174	URA3 CEN Amp ^R sna3 ^{Y109A} -GFP	(40)
pDN263	PGK1pr::RLuc sna3 ^{Y109A} -FLuc (pDN251)	This work
pVJS3	Amp ^R GDI-intein-CBD	(75)
pRP2	VPS21pr::GFP-VPS21 (pRS416)	This work
pGO700	mCherry-CPS1 (pRS414)	G. Odorizzi (U Colorado)
pSL5916	GFP-Atg8 (pRS416)	S. Lemmon (U Miami)
pCB257	GFP-FYVE(EEA1) (pRS424)	C. Burd (Yale)
рСВ260	GFP-FYVE(EEA1) (pRS425)	C. Burd (Yale)
AMP219	Amp ^R GST-VPS21 (p//GST)	(52)
AMP220	Amp ^R GST-YPT52 (p//GST)	(52)
AMP217	Amp ^R GST-YPT53 (p//GST)	(52)
AMP218	Amp ^R GST-YPT7 (p//GST)	(52)
pDN125	GFP-DID2 (pRS416)	(35)
pPHY2427	URA3 2μ Amp ^R CUP1 _{pr} ::3xHA-ATG13 (pRS426)	P. Hermann (Ohio State)

Low-copy dual luciferase vector pDN251 derived from removal of the 2μ locus from pDN228 by restriction digestion with AatII and subsequent gap repair at the AatII cut site using a *LoxP::CEN/ARSH4::LoxP* insert. For trafficking studies, *SNA3* or *sna3*^{Y109A} were PCR amplified from genomic DNA or pMM174 (40) templates, respectively, then cloned in-frame into SacII-digested pDN251 to create chimeric FLuc reporters (pDN252 and pDN263).

Bacterial expression vector pDN190 was constructed by PCR amplifying a portion of the pRS416 vector containing both the *URA3* and *CEN/ARSH4* loci and inserting this sequence into Agel-digested AMP451 via gap repair in yeast. The resulting pRSF-family vector contains a T7 promoter to allow regulated expression of His₇-MBP (maltose binding protein) in bacteria, but also replicates stably in yeast. His₇-MBP-Gyp3 and His₇-MBP-Gyp7 bacterial expression plasmids were constructed by PCR-amplification of *GYP3* and *GYP7* coding sequences from yeast genomic DNA, and gap repair of BamHI/Sall-digested pDN190.

Culture and media

Standard methods for culture and media were used (76). Liquid media for experiments involving added $CaCl_2$ or NaCl was adjusted to pH 5. Synthetic media was prepared using NH₄Cl and buffered with 7.5 mm

1424

succinic acid in order to avoid precipitation of Ca²⁺ salts (49). Microbial growth in liquid cultures was assayed at 30°C with periodic shaking using a Bioscreen-C machine (Growth Curves USA) and 150 μ L cultures inoculated at OD₆₀₀ = 0.1.

Direct competition between $vps9\Delta$ and $vps9\Delta$ $gyp3\Delta$ strains was assessed by mixing independently grown cultures to achieve equal representation in a mixed starter culture, then diluting 1:1000 in fresh YPD (pH 5) with or without 200 mM CaCl₂ and shaking at 30°C. This procedure was iterated four times. Starter and overnight cultures were serially diluted to achieve a plate inocultur of ~150 colonies and applied to YPD plates, from which isolated colonies were restruck to YPD plates containing G418 in order to determine the presence of the $gyp3\Delta$::*KAN* marker.

Trafficking and dual luciferase assays

Colorimetric assays of CPY-invertase secretion were performed essentially as described (77). Cells for invertase assays were grown in media supplemented with 2% fructose instead of glucose. Levels of the MVB cargo Sna3 were analyzed using a Promega dual-luciferase assay system. Cells were grown to early log phase ($OD_{600} = 0.25 - 0.4$) at 30°C in synthetic medium (2% glucose) lacking uracil and supplemented with

casamino acids, cycloheximide (50 g/mL final) was added for 20-30 min, and cultures were harvested by low-speed centrifugation. Cell pellets containing 0.4 OD_{600} \times mL were resuspended in 500 μ L lysis buffer and incubated at RT for 25 min before vortexing with glass beads for a further 5 min at RT (22°C). Five microliter aliquots of cell lysate were analyzed in 96-well plate format using a Perkin Elmer Victor Light Model 1420 luminometer. After subtracting backgrounds, luciferase activity of Sna3-FLuc was normalized versus the activity of soluble RLuc, expressed from the same plasmid and driven by the constitutive PGK1 promoter. Luciferase assays to examine promoter activity were performed similarly, but without bead beating and with a cell input of 0.22 OD_{600} × mL. FLuc activity was further normalized by subtracting the FLuc:RLuc signal ratio resulting from promoter-less FLuc. Autophagy was initiated in log phase cells by addition of 200 ng/µL rapamycin (LC Laboratories). Plasmidborne CUP1pr::3xHA-ATG13 expressed adequately without added copper, permitting analysis without toxic effects associated with copper induction.

Light and electron microscopy

For microscopy, overnight yeast cultures were diluted in synthetic media and grown at 30 $^{\circ}\text{C},$ 225 rpm to OD_{600} ${\sim}0.4,$ then harvested by low-speed centrifugation. For FM 4-64 labeling of endolysosomal membranes (78), cell pellets were resuspended in 50 μ L synthetic media containing FM 4-64 (2 m) and incubated at 30°C for 15 min. Labeled cells were recovered by centrifugation and rinsed twice in media before resuspension in media and shaking at 30°C to permit growth to OD₆₀₀ ~0.5 before concentration of cells and imaging. Where specified, cells were labeled with FM 4-64 for 30 min without chase and imaged immediately. Fluorescence microscopy was performed using an IX71 Olympus microscope equipped with 100× UPlanFLN, NA 1.30 objective, Hg arc or custom LED illuminator and Andor iXon electron-multiplying charge-coupled device driven by Andor IQ v6 software. Confocal fluorescence microscopy was performed as described (28). Plate images were captured using a Sony DSC-W30 digital camera. Thin section TEM was performed essentially as described (28). Photoshop v9.0 (Adobe) and Canvas v9.0 (ACD) software were used for image processing and figure preparation.

Protein purification

Purifications of Gyp1_{TBC} (61) and Rab GDI (75) were as described. His₇-MBP-FLAG-Gyp3 and His7-MBP-Gyp7 were expressed in Escherichia coli BL21(DE3)-pRIL grown in TB medium with chloramphenicol and kanamycin to $OD_{600} = 1.0-1.5$. Protein expression was induced with 0.3 mm IPTG for 3.5 h at 30°C and cells were harvested by centrifugation. Cells containing His7-MBP-FLAG-Gyp3 or His7-MBP-Gyp7 were lysed by sonication on ice in Buffer A (50 mm sodium phosphate, pH 7.4, 500 mm NaCl and 20 mm imidazole, pH 7.4) or Buffer B (25 mm Tris-HCl, pH 7.9, 300 mm NaCl, and 40 mm imidazole, pH 7.9), respectively, in the presence of protease inhibitors (PMSF, leupeptin and pepstatin), lysozyme and DNasel. Cell lysates were clarified by centrifugation (18500 \times g, 20 min, 4°C). GAPs were affinity purified on HisTrap FF columns (GE Healthcare) and washed extensively with Buffer A or Buffer B prior to elution with Buffer A or Buffer B supplemented with 500 mm imidazole. His7-MBP-FLAG-Gyp3 fractions were subsequently diluted with 25 mm Tris-HCl, pH 7.5 to [NaCl] <100 mm and loaded onto a Source Q ion exchange column (GE Healthcare). His7-MBP-FLAG-Gyp3 eluted at 25 mm Tris-HCl and 220-240 mm NaCl. Eluate fractions were concentrated and then further purified by size exclusion on Superdex 200 pre-equilibrated with Buffer C (25 mM Tris-HCl, pH 7.5, 300 mm NaCl, and 2 mm 2-mercaptoethanol). Fractions collected from the major peak were concentrated and glycerol was added to 5% (w/v) prior to storage. His7-MBP-Gyp7 containing fractions eluted from HisTrap FF resin were concentrated and buffer exchanged on Superdex 75 pre-equilibrated with Buffer C, concentrated, and glycerol was added to 10% prior to storage. For both GAPs, aliquots in thin-wall PCR tubes were flash-frozen in liquid N₂ and stored at -80°C.

GST-Rab fusions were expressed in *E. coli* BL21(DE3)-pRIL grown in TB with chloramphenicol and kanamycin to $OD_{600} = 2-2.5$. Expression was induced with 0.1 mm IPTG overnight at 18°C. Cells were harvested by

Rab5 Specificity and Regulation in Yeast

centrifugation and resuspended into Buffer D (50 mm Tris-HCl, pH 7.9, 100 mm NaCl, 0.1% Triton-X-100 and 10% glycerol) supplemented with protease inhibitors (PMSF, leupeptin and pepstatin), lysozyme and DNasel. Cells were lysed with a pressure lysis apparatus (EmulsiFlex-C5; Avestin) with minimum peak pulses of 7500 psi, then clarified by centrifugation (18500 \times *g*, 20 min, 4°C). GST-Rabs were adsorbed to 2 mL of Glutathione Sepharose 4B Resin (GE Heathcare) for 3 h at 4°C. Bound GST-Rabs were washed three times with Buffer D, then Buffer D plus 500 mm NaCl, and again in Buffer D, all without Trition-X-100 or glycerol. GST-Rabs were eluted by incubating the resins in 2 mL of buffer E (50 M Tris-HCl, 7.9, 100 тм NaCl, 5 тм MgCl₂, 10 тм glutathione and 2 тм 2-mercaptoethanol) for 10 min at 4°C. Elution was repeated at least three times. Eluted fusion proteins were concentrated to 4-8 mg/mL and then glycerol was added to 10% (w/v) prior to storage. All concentration steps were performed using Amicon Ultra-15 concentrators (10 000 MWCO; Millipore). Aliquots were flash-frozen in liquid N₂ and stored at -80° C.

GAP assays

The GTP hydrolysis assay is modified from Pan et al. (6) and Lo et al. (61). GAPs were buffer exchanged into Buffer F (20 mm HEPES- NaOH, pH 7.5 and 150 mm NaCl) using Micro Bio-Spin columns (BioRad). For GTP loading, 3.0-3.5 mg of GST-Rab was suspended in 800 µL of Buffer D containing 5-10 mm reduced glutathione and supplemented with 1 mm DTT, 25× molar excess GTP and 15 mm EDTA. After 60 min at 25–28 $^{\circ}\text{C},\,\text{MgCl}_2$ was added to 20 mm and GST-Rab fusions were incubated for an additional 15 min on ice to terminate the GTP loading reaction. GTP-loaded Rabs were rapidly separated from free nucleotides on 5 mL HiTrap desalting columns (GE healthcare) pre-equilibrated with Buffer F and ran at 4°C. Eluted Rabs were concentrated using an Amicon Ultra-0.5 mL concentrator (30 000 MWCO; Millipore) to \geq 2.0 mg/mL. Inorganic phosphate released by single-turnover GTP hydrolysis was monitored in real time with the EnzChek Phosphate Assay (Invitrogen) in 100 µL reaction volume in 96well half area plates (Corning). To initiate the assay 60 µL of GTP-loaded GST-Rab was added to 40 µL of assay buffer with or without GAP. Each reaction contained a final concentration of 20 mm HEPES-NaOH, pH 7.5, 150 mм NaCl, 10 mм MgCl₂, 0.15 mм 2-amino-6-mercapto-7-methylpurine ribonucleoside, 0.75 U/mL purine nucleoside phosphorylase, 20 μ M GTPloaded GST-Rab, and GAPs at the indicated concentrations. Each reaction also contained 1% (w/v) bovine serum albumin (BSA), which improved assay signal-to-noise ratio and reproducibility. Qualitatively similar results were obtained in the absence of BSA. Before use BSA was exchanged into Buffer F by gel filtration on Superdex 75 (GE healthcare), and stored at 4°C. Inorganic phosphate evolution was monitored by measuring A360nm in a Perkin Elmer Victor 3 plate reader.

Subcellular fractionation, antibodies and western blotting

Cells (18 $OD_{600} \times mL$) grown to late log phase were harvested by lowspeed centrifugation and resuspended in 10 mm DTT, 0.1 m Tris-HCl pH 9.4 for 10 min. Cells were again pelleted and resuspended in spheroplasting buffer (50 mM HEPES-KOH pH 7.2, 1× yeast nitrogen base with casamino acids, 2% glucose, 1 M sorbitol) with Zymolyase 20T (Seikagaku) subjected to further purification by ion exchange chromatography. Spheroplasts were pelleted and rinsed once in spheroplasting buffer before resuspension in ice-cold lysis buffer [20 mm HEPES-KOH pH 7.2, 50 mm KOAc, 200 mm sorbitol, 0.1 mm Pefabloc SC, 1 mm PMSF, 0.01 mm chymostatin, 1 g/mL each of aprotinin, leupeptin and pepstatin, and 1X protease inhibitor cocktail (EDTA-free; Roche)] and lysis by 20 strokes in an ice-cold Dounce homogenizer. Lysates were cleared of cell debris and nuclear membranes by centrifugation at 1000 \times ${\it g}$ for 5 min before sedimenting the supernatant again at 15000 \times g to yield endolysosomal membrane pellet (P15) and supernatant (S15) fractions. P15 fractions were resuspended in lysis buffer and proteins were harvested from total. P15 and S15 samples by precipitation with 0.15% deoxycholic acid and 10% trichloroacetic acid followed by two washes in acetone. Protein pellets were dried under vacuum and resuspended in Laemmli sample buffer (40 µL sample buffer

per OD_{600} \times mL equivalent) before boiling 10 min. For each SDS-PAGE lane, 0.2 OD_{600} \times mL of cell suspension was analyzed.

Membrane extraction of Rabs by GDI was done by resuspending a P15 membrane pellet from 14 OD₆₀₀ × mL of cell suspension in 140 µL lysis buffer dividing the total into 25 µL (2.5 OD₆₀₀ × mL of cell suspension) aliquots to which were added lysis buffer with or without 3 µM recombinant Rab GAP. Samples were incubated on ice 15 min. Some tubes received 9 µM recombinant GDI immediately prior to sedimentation at 15000 × **g** for 15 min, after which pellets were resuspended in lysis buffer and proteins from both membrane-associated (P15) and soluble (S15) fractions were processed as described above. 0.2 OD₆₀₀ × mL of cell suspension per sample was analyzed by SDS-PAGE and western blot.

Western blotting was performed using an Odyssey fluorescence scanner (LI-COR Biosciences). Monoclonal anti-PGK (3'-phosphoglycerate kinase), anti-ALP (alkaline phosphatase) and anti-Vph1 antibodies were obtained from Invitrogen. Polyclonal anti-Ypt7 and -Vam3 antibodies were gifts of W. Wickner (Dartmouth College, Hanover, NH). Polyclonal anti-Vps21 antisera was a gift of B. Horazdovsky (Mayo Clinic, Rochester, MN). Polyclonal anti-Ape1 antisera were a gift of D. Klionsky (Univ. Michigan, Ann Arbor, MI).

Quantitative analyses

Quantification of western blots was performed using Odyssey software (LI-COR Biosciences). Summary statistics for raw data from yeast growth curves were derived using the YODA program (79). Other statistical analyses were performed using Microsoft Excel and GraphPad Prism (v. 4 and 5). To obtain Rab-GAP catalytic specificity constants (k_{cat}/K_{M}) and Rab intrinsic GTP hydrolysis rates ($k_{intrinsic}$), data collected from each Rab replica were adjusted to subtract baseline absorbance and then the kinetic curves for each Rab-GAP combination were simultaneously fitted by a nonlinear lest-squares method with a pseudo-first order Michaelis–Menten model (6):

$$\left(A_{\infty} - A_{0}\right) \left(1 - e^{-\left(k_{\text{intrinsic}} + \frac{k_{\text{Cat}}}{K_{\text{M}}}\left[\text{GAP}\right]\right)t}\right) + A_{0}$$

where $k_{\text{cat}}/k_{\text{M}}$ and $k_{\text{intrinsic}}$ were treated as non-zero global parameters. A_{∞} and A_0 were obtained from the Rab-Gyp1_{TBC} combination fit and treated as global parameters within each Rab replica.

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D. N. and A. M. conceived, and D. N. carried out, most experiments. S. L. designed and performed GTP hydrolysis assays. M. R. performed and analyzed electron microscopy. H. C. performed and analyzed assays of autophagy. J. M. and M. R. performed some of the fluorescence microscopy. All authors interpreted results. D. N. and A. M. wrote the manuscript.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1: Endosome morphology. A–C) Thin-section electron micrographs show endosomes in $ypt52\Delta$ (A), $ypt53\Delta$ (B) or $gyp3\Delta$ (C) cells, all possessing characteristic MVB morphology. Bar, 100 nm. Vac, vacuole (associated with Figure 2).

Figure S2: Autophagy in Rab5-deficient cells. A, B) Confocal fluorescence microscopy of GFP-Atg8 expressed in cells stained with FM 4-64 (A) or co-expressing MVB cargo mCherry-CPS (B). Scale bars, 2 m. (C) Western blot analysis of Ape1 maturation in cell lysates with or without rapamycin treatment. (D) Western blot analysis of a time course following rapamycin treatments, indicating dephosphorylation of the TOR kinase substrate Atg13. pApe1, premature Ape1. mApe1, mature Ape1. P-Atg13-HA, phospho-Atg13-HA (associated with Figure 2).

Figure S3: Rab5 signaling and effectors are required to resist Ca²⁺stress. A) Limiting dilution plate growth assay using YPD agar with the indicated concentrations of added Ca²⁺. B) Growth curves in YPD broth supplemented with 200 mM Ca²⁺. For all experiments, cells were grown at 30°C (associated with Figure 3).

Figure S4: Peri-vacuolar localization of Rab5 effector molecules in wild-type and *gyp3*∆ **cells.** A, B) Fluorescence microscopy of FM 4-64-stained cells expressing either Vps3-GFP (A) or Vps8-GFP (B). Scale bars, 2 m (associated with Figure 8).

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