Subunit organization and Rab interactions of Vps-C protein complexes that control endolysosomal membrane traffic

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ABSTRACT Traffic through late endolysosomal compartments is regulated by sequential signaling of small G proteins of the Rab5 and Rab7 families. The Saccharomyces cerevisiae Vps-C protein complexes CORVET (class C core vacuole/endosome tethering complex) and HOPS (homotypic fusion and protein transport) interact with endolysosomal Rab5s to coordinate their signaling activities. To better understand these large and intricate complexes, we performed interaction surveys to assemble domain-level interaction topologies for the eight Vps-C subunits. We identified numerous intersubunit interactions and up to six Rab-binding sites. Functional modules coordinate the major Rab interactions within CORVET and HOPS. The CORVET-specific subunits, Vps3 and Vps8, form a subcomplex and physically and genetically interact with the Rab5 orthologue Vps21. The HOPS-specific subunits, Vps39 and Vps41, also form a subcomplex. Both subunits bind the Rab7 orthologue Ypt7, but with distinct nucleotide specificities. The in vivo functions of four RING-like domains within Vps-C subunits were analyzed and shown to have distinct functions in endolysosomal transport. Finally, we show that the CORVET- and HOPS-specific subunits Vps3 and Vps39 bind the Vps-C core through a common region within the Vps11 C-terminal domain (CTD). Biochemical and genetic experiments demonstrate the importance of these regions, revealing the Vps11 CTD as a key integrator of Vps-C complex assembly, Rab signaling, and endosomal and lysosomal traffic.

INTRODUCTION

Membrane traffic entails cycles of vesicle budding, transport, tethering, docking, and fusion with target membranes. These processes are regulated by the Rab-Arf family of small G proteins, which act as switches that confer specificity and directionality and as timers that control rates of membrane ingress and egress at each compartment. In current models, a patch of membrane moving (or maturing) through a sequence of compartments is signified by the presence of a specific associated Rab. As the patch matures, a handoff occurs in which one Rab is replaced by the next. This an arrangement is called a Rab cascade or relay (Grosshans et al., 2006). Traffic through the endocytic-lysosomal pathway is controlled by members of the Rab5 and Rab7 families, which operate at earlier and later compartments, respectively. The Rab5-to-Rab7 handoff is also controlled by signaling state of each Rab and by the Vps-C regulatory complexes CORVET and HOPS (Rink et al., 2005; Peplovska et al., 2007). The Rab5-to-Rab7 handoff is also controlled by a second evolutionarily unrelated complex, Mon1(SAND1)-Ccz1 (Kucharczyk et al., 2001, 2009; Kinchen and Ravichandran, 2010; Nordmann et al., 2010; Poteryaev et al., 2010), which has also been called “HOPSII” (Wang et al., 2003).

The Vps-C protein complexes were discovered in S. cerevisiae (Rieder and Emr, 1997; Srivastava et al., 2000; Nickerson et al., 2009). They contain a four-subunit core (Pep5/Vps11, Vps16, Pep3/ Vps18, and Vps33) and two additional, compartment-specific subunits. CORVET, which contains the Vps-C core plus Vps3 and Vps8 (Raymond et al., 1999; Horazdovsky et al., 1996; Peterson and Emr, 2001; Peplovska et al., 2007), operates at late endosomes and interacts with the Rab5 orthologue Vps21 (Horazdovsky et al., 1996;
Peplowska et al., 2007). HOPS, which contains the Vps-C core plus Vps39 (also called Vam6) and Vps41 (Seals et al., 2000), operates at the lysosomal vacuole and interacts with the Rab7 orthologue Ypt7 (Seals et al., 2000; Wurmser et al., 2000; Brett et al., 2008).

CORVET controls traffic into late endosomes, whereas HOPS controls all traffic—including late endosomes, autophagosomes, and Golgi-derived AP-3 vesicles—into the vacuole. Accordingly, the loss of any Vps-C core subunit results in the absence or malformation of all late-endosomal compartments, as well as severe conditional growth defects. Conversely, the loss of CORVET- or HOPS-specific subunits causes selective defects at late endosomes or vacuoles (Nickerson et al., 2009). HOPS promotes Ypt7-dependent membrane tethering (Stroup et al., 2006) and SNARE-mediated membrane fusion (Sato et al., 2000; Mima et al., 2008). The HOPS subunit Vps39 was reported to be an activating guanosine nucleotide exchange factor (GEF) of Ypt7 (Binda et al., 2009; Wurmser et al., 2000), but more highly purified preparations of Vps39 and HOPS do not exhibit intrinsic nucleotide exchange activity toward Ypt7 (Nordmann et al., 2010; Ostrowicz et al., 2010; Lobingier, Brett, and Merz, unpublished data). Moreover, strong genetic (Kucharczyk et al., 2000, 2001; Wang et al., 2002) and biochemical (Nordmann et al., 2010) evidence indicates that another complex, Mon1-Ccz1, is a bona fide Ypt7 GEF. The Vps41 subunit of HOPS directly interacts with Ypt7, and it is essential for stable HOPS binding to Ypt7-GTP (Brett et al., 2008). Functions analogous to HOPS have been proposed for CORVET (Peplowska et al., 2007), although CORVET is not known to tether membranes directly or to promote SNARE assembly. Hybrid complexes containing both CORVET- and HOPS-specific subunits (Peplowska et al., 2007) raise the possibility that Vps-C complexes are remodeled on-the-fly, concomitantly with the endolysosomal Rab cascade. However, dynamic subunit exchange has been neither demonstrated nor linked to specific endolysosomal trafficking events.

Here we present domain-level dissections of Vps-C subunit interactions with one another and with Rab small G proteins. We employ yeast two-hybrid (Y2H) miniarrays to systematically search for Vps-C subunit–subunit and subunit–Rab contacts and purify three dimeric subassemblies. Two subassemblies are composed of HOPS- or CORVET-specific subunits; we show that they function as integrated Rab-interaction modules. Through genetic analyses, we show that the core subunit Vps11 is a key scaffold for Vps-C complex assembly and demonstrate that its C-terminal domain (CTD) is of special importance, physically and functionally linking the HOPS and CORVET Rab-interaction modules to the Vps-C core.

**RESULTS**

**Domain architecture of Vps-C subunits**

We analyzed yeast and human Vps-C subunits using the ROBETTA suite of structure prediction algorithms (Kim et al., 2004; Chivian et al., 2005). Previously, ~200 sequences of respiratory homology to clathrin proximal leg were detected in C-terminal regions of Vps8, Vps11, Vps18, Vps39, and Vps41 (Conibear and Stevens, 1998; Ybe et al., 1999). Vps39 was reported to contain a short motif similar to a COPI (coat protein complex I) outer-shell coat subunit (Conibear and Stevens, 1998). Vps41 was shown to contain WD40 motifs (repeats of ~40 residues, often terminating in W and D), which typically form β-propellers (Rehling et al., 1999). Our analyses (Figure 1 and Supplemental Table 1; Nickerson et al., 2009) revealed a pervasive similarity to vesicle coat and nuclear pore proteins: seven of eight subunits contain a predicted N-terminal β-propeller followed by a C-terminal α-solenoid. In addition, Vps8, Vps11, and Vps18 have C-terminal RING (really interesting new gene) motifs, and Vps39 has a partial RING motif, although this region lacks residues that would coordinate a second Zn2+ ion. Similar predictions were obtained for human and yeast orthologues (Supplemental Table 1). These findings provided the basis for our domain-level dissection of Vps-C subunits.

**Domain-level interaction topology of Vps-C complexes**

HOPS and CORVET are sufficiently stable to withstand affinity isolation, chromatographic purification, and sedimentation (Rieder and Emr, 1997; Seals et al., 2000; Stroup et al., 2006; Peplowska et al., 2007; Angers and Merz, 2009), but only a few direct intersubunit contacts have been identified, using disparate approaches and
organisms. To obtain a broader inventory of intersubunit contacts, we performed exhaustive Y2H analyses.

Y2H can be very sensitive, potentially reporting bridging interactions mediated by proteins not encoded by the query constructs, so we used a de-tuned Y2H system originally designed to minimize false-positive signals in high-throughput screens (Uetz et al., 2000). This system uses single-copy bait and prey vectors rather than the more commonly used multicopy vectors. Recent quantitative assessments demonstrate that use of single-copy Y2H vectors, in combination with the HIS3 reporter gene and 3-aminotriazole (3-AT) in colony growth assays, nearly eliminates false-positive signals in Y2H interaction experiments (Venkatesan et al., 2009; Chen et al., 2010).

Identical highly stringent assay conditions were used throughout the present study.

The Y2H-derived Vps-C interaction topology is summarized in Figure 1; the full results are reported in Supplemental Table S2. Our survey replicated (and in most cases mapped to higher resolution) almost all published interactions within Vps-C complexes across all organisms studied to date (Nakamura et al., 1997; Sevrioukov et al., 1999; Srivastava et al., 2000; Wurmser et al., 2000; Peterson and Emr, 2001; Pulipparacharuvil et al., 2005; Peplovska et al., 2007). This finding suggests that not only the subunit folds but also the interaction topologies and functional dynamics of Vps-C complexes are broadly conserved. Our survey also revealed candidate interactions that to our knowledge have not previously been reported. The minimal regions sufficient for each interaction are largely nonoverlapping, strongly suggesting that most interactions detected in this survey are direct rather than bridged through other subunits.

Vps11, particularly in its C-terminal region, is a densely connected hub (Figure 1E), interacting with Vps18, Vps8, Vps3, and Vps39. Vps39 and Vps3 share substantial sequence homology, with the region of highest similarity within a stretch of ∼100 residues near their C-termini. The Vps3 and Vps39 CTDs interacted strongly with residues 736–926 in the Vps11 CTD, suggesting that they may bind to a single site within Vps11 (Figure 1, C and D). The HOPS subunit Vps39 also interacts with the HOPS subunit Vps41. The Vps39–Vps41 interaction was relatively weak, but it was detected in both bait–prey and prey–bait configurations; this was the strongest interaction detected between Vps41 and any other HOPS subunit.

Isolation of three Vps-C subcomplexes

To examine the ability of subunits to heterodimerize in the absence of the HOPS and CORVET holocomplexes, pairs of subunits were coexpressed in insect cells. In each case, one subunit was expressed as a glutathione S-transferase (GST) fusion, the other as a His6 fusion, and complexes were purified using the two affinity tags in sequence. From several tested pairs, three subcomplexes were identified (Figure 2). Vps16 is required for Vps33 association with Vps11 and Vps18 (Rieder and Emr, 1997), and a region corresponding to yeast Vps16 residues 451–595 mediates binding between Drosophila Vps16A and B and Vps33A and B (Pulipparacharuvil et al., 2005). Our Y2H survey mapped an overlapping site within Vps16 (residues 479–798; Figure 1) and indeed Vps16 and Vps33 form a subcomplex when coexpressed (Figure 2A). The CORVET subunits Vps3 and Vps8 also form a subcomplex (Figure 2B). This binding interaction has not to our knowledge been reported previously, and it was not detected in our Y2H survey. Like Vps3 and Vps8, the HOPS subunits Vps39 and Vps41 formed a binary complex (Figure 2, C and D); the interacting domains were mapped by Y2H to the N-termini of both proteins (Figure 1C; Supplemental Table S2). The Vps39 CTD was sufficient to interact with Vps11 and was necessary for stable Vps39 association with HOPS (Figure 1; Supplemental Table S2; Wurmser et al., 2000). However, the Vps39 CTD was not needed for Vps39 binding to Vps41 (Figure 2D). Both subcomplexes exhibited signs of lability, suggesting that their interactions are stabilized when they reside within the HOPS and CORVET holocomplexes.

Y2H identification of Vps-C Rab-interaction sites

CORVET interacts with the Rab5 orthologue Vps21, and HOPS interacts with the Rab7 orthologue Ypt7. Vps39 binds GDP- or GTP-Ypt7 and was previously reported to harbor Ypt7 GEF activity (Wurmser et al., 2000; Binda et al., 2009). Vps41 binds directly to Ypt7 and is essential for stable binding of HOPS to Ypt7-GTP (Brett et al., 2008). Until recently, no Rab-binding sites were definitively identified within CORVET, although there was genetic evidence that Vps3 and Vps8 contribute to the CORVET Vps21 binding activity (Horazdovsky et al., 1996; Peplovska et al., 2007; Markgraf et al., 2009; Pawelec et al., 2010). To identify Rab-binding sites, Y2H miniarrays comprising the 11 yeast Rab s were prepared. In addition to wild-type Rabs, Rab point mutants were included in the arrays, including mutants favoring specific nucleotide states or lacking C-terminal sites of covalent prenylation (lipid-free). Rab miniarrays were used to probe Vps-C miniarrays in both bait and prey configurations (Supplemental Table S3). At least six candidate interactions were identified by Y2H. Vps21 interacted with CORVET subunits Vps3 and Vps8, and with the N-terminal domain of Vps-C core subunit Vps11 (Supplemental Table S3). Ypt7 interacted with HOPS subunits Vps39 and Vps41 (Figure 3A; Supplemental Table S3; Brett et al., 2008; Wurmser et al., 2000) and weakly with the core subunit Vps33.

The HOPS–Ypt7 interaction module

In pulldowns, native HOPS interacts preferentially with Ypt7-GTP/S versus Ypt7-GDP (Price et al., 2000; Seals et al., 2000; Peplovska et al., 2007; Brett et al., 2008). However, under conditions that yielded highly selective binding of native HOPS to Ypt7-GTP, there was no GDP versus GDP selectivity when purified Vps41 was tested. Further studies indicated that nucleotide stripping under our previous conditions was inefficient. With a more stringent strip-exchange procedure, purified Vps41 selectively bound Ypt7-GTP/S (Figure 3B and Supplemental Figure S1), establishing Vps41 as a direct effector of Ypt7-GTP and explaining why it is required (Brett et al., 2008) for the stable association of HOPS with Ypt7-GTP.

In previous studies, Vps39 was reported either to bind Ypt7-GDP (Wurmser et al., 2000) or to show no nucleotide selectivity (Brett et al., 2008). Under our improved nucleotide-loading conditions, purified Vps39–860A bound Ypt7 without any detectable GNP selectivity (Figure 3, C and D). Taken together, our Y2H and biochemical data show that the N-terminal predicted β-propellers of Vps39 and Vps41 interact with one another and with the vacuolar Rab Ypt7, with Vps41 showing strong selectivity for Ypt7-GTP and Vps39 showing little or no nucleotide selectivity. Because Vps39 was the only HOPS subunit that interacted with Vps41 in our Y2H experiments, we asked whether Vps41 could remain associated with the remaining HOPS subunits in the absence of Vps39. Coimmunoprecipitation experiments using anti-Vps11 antibodies (Figure 3E) demonstrated that the five remaining HOPS subunits remain stably associated in the absence of Vps39. This result indicates that, in addition to Vps39, Vps41 has additional binding interactions with the core that were not detected in our Y2H survey. In contrast to Vps39, the core subunit Vps18 has a crucial role in the assembly of HOPS and the Vps-C core, because deletion of Vps18 prevented...
Coimmunoprecipitation of Vps11 and Vps16, Vps18, Vps33, or Vps41 (Figure 3E). As the region of Vps39 required for its interaction with Vps11 maps to the Vps39 C-terminus (residues 860–979), we analyzed the in vivo phenotypes of two C-terminal Vps39 deletion mutants. In vivo homotypic fusion was evaluated by examining vacuole morphology with the endocytic tracer dye FM4-64 (Figure 4A). Traffic to the vacuole was evaluated by subcellular fractionation followed by immunoblotting for two cargo proteins, CPY (carboxypeptidase Y, Prc1) and alkaline phosphatase (ALP) (Figure 4B). In this procedure, Golgi membranes and small vesicles fractionate to the 100,000 × g pellet (P100). Endosomes are found in the 13,000 × g pellet (P13) and P100 fractions. Vacuoles are found in the P13 fraction almost exclusively. Immunoblotting of fractions with fiduciary markers for vacuolar, endosomal, and soluble markers was performed for each experiment, and proper fractionation was confirmed for each mutant (unpublished results). CPY is transported from the late Golgi to endosomes, and then to the vacuole. ALP is delivered from the Golgi to the vacuole through the AP-3 pathway, which bypasses endosomes. Both ALP and CPY are synthesized as inactive proenzymes (p) that are proteolytically processed into active mature (m) enzymes upon arrival at the vacuole. Transport defects result in the accumulation of slow-migrating ALP or CPY proenzymes in the P100 and P13 fractions (Figure 4B).

The vps39Δ-860Δ mutation completely phenocopies the defects in CPY and ALP localization and maturation, and vacuole morphology, conferred by vps39Δ or vps41Δ mutations (Figure 4; Supplemental Table S2). In contrast, vps39Δ-979Δ mutant cells produce Vps39 with an intact Vps11-binding site and exhibit only mild trafficking defects (Figure 4B). The vps39Δ-860Δ mutant is not due to poor expression or accelerated turnover of Vps39-860 protein. Because Vps39 residues 860–979 are necessary and sufficient for Vps39 binding to Vps11 and Vps39 association with HOPS, but not required for Vps39 interactions with Ypt7 or Vps41, our results suggest that Vps39 function depends on its interactions with HOPS through both Vps11-binding sites and the Vps39 C-terminus, and that Vps39–860 is not functional in the absence of an interacting partner whose tag can bind the affinity resin. Vps39–860 is not functional in the presence of noncognate affinity beads.

![Figure 2: Purification of Vps-C subcomplexes.](image)

(A–C) In each panel, the gel on the left shows isolation of a subcomplex, while the gel on the right shows pulldown specificity controls. Lanes 1 and 2, GST-tagged or His6-tagged subunits were expressed individually and purified on either Ni2+-NTA resin (Ni2+) or glutathione-agarose (GSH) resin. Lanes 3 and 4, pairs of GST- and His6-tagged subunits were coexpressed, captured on GSH resin and eluted with reduced glutathione (lanes 3), and then recaptured on Ni2+ resin (lanes 4). Lanes 5–8, pulldown specificity controls showing that Ni2+-NTA beads retain little or no GST fusion protein, and that glutathione-sepharose beads (GSH) retain little or no His6 fusion protein when these proteins are expressed individually (i.e., in the absence of an interacting partner whose tag can bind the affinity resin). Lanes 6 and 8, material retained on the noncognate affinity beads. Lanes 5 and 7, unbound (flowthrough) material. The SDS–PAGE gels were silver-stained.

(D) Copurification of His6- and His6-Vps41 associated with GST-Vps39 or GST-Vps39 truncation mutants.

**Figure 2:** Purification of Vps-C subcomplexes. (A–C) The indicated subunits were expressed individually or in pairs in insect cells and isolated by affinity chromatography. In each panel, the gel on the left shows isolation of a subcomplex, while the gel on the right shows pulldown specificity controls. Lanes 1 and 2, GST-tagged or His6-tagged subunits were expressed individually and purified on either Ni2+-NTA resin (Ni2+) or glutathione-agarose (GSH) resin. Lanes 3 and 4, pairs of GST- and His6-tagged subunits were coexpressed, captured on GSH resin and eluted with reduced glutathione (lanes 3), and then recaptured on Ni2+ resin (lanes 4). Lanes 5–8, pulldown specificity controls showing that Ni2+-NTA beads retain little or no GST fusion protein, and that glutathione-sepharose beads (GSH) retain little or no His6 fusion protein when these proteins are expressed individually (i.e., in the absence of an interacting partner whose tag can bind the affinity resin). Lanes 6 and 8, material retained on the noncognate affinity beads. Lanes 5 and 7, unbound (flowthrough) material. The SDS–PAGE gels were silver-stained. (D) Copurification of His6- and His6-Vps41 associated with GST-Vps39 or GST-Vps39 truncation mutants.

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but not with Vps21-S21L (a GDP-biased mutant) and any other Rab (Figure 5, A and B; Supplemental Table S3). Additional controls validated the Y2H Rab constructs: the Vps21 effector Vac1 (Peterson et al., 1999) had Rab interaction patterns identical to Vps3 and Vps8, whereas the Vps21 GEF Vps9 interacted strongly with Vps21-S21L but not with wild-type Vps21 or Vps21-Q66L (Hama et al., 1999; Plemel and Merz, unpublished data). Vps3, like Vps41 and Vps39, interacted with its cognate Rab through a predicted N-terminal β-propeller. In contrast, Vps8 interacted with Vps21 through the Vps8 C-terminal predicted α-solenoid (Figure 5A; see also Pawelec et al., 2010). The Vps3–Vps21 interaction was validated biochemically. Vps3 in cell lysates, presumably as a subunit of CORVET, bound to Vps21-GTP7S with high specificity (Figure 5C; Peplowska et al., 2007). Moreover, purified Vps3 bound to Vps21 with strong selectivity for Vps21-GTP versus -GDP (Figure 5D). These experiments suggest that both Vps3 and Vps8 contain Vps21-binding sites, and establish that Vps3 is a direct, GTP-selective effector of Vps21.

The Vps3 CTD is highly homologous to the Vps39 CTD. As with Vps39, the Vps3 CTD is required for its interaction with the Vps11 catalytic core of the HOPS complex (190-residue region within the Vps11 CTD). (A) Vacuole morphology phenotypes of vps39 null mutant (Figure 5A; see also Pawelec et al., 2010). The Vps3–Vps21 interaction was validated biochemically. Vps3 in cell lysates, presumably as a subunit of CORVET, bound to Vps21-GTP7S with high specificity (Figure 5C; Peplowska et al., 2007). Moreover, purified Vps3 bound to Vps21 with strong selectivity for Vps21-GTP7S versus -GDP (Figure 5D). These experiments suggest that both Vps3 and Vps8 contain Vps21-binding sites, and establish that Vps3 is a direct, GTP-selective effector of Vps21.

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Δvps11–926

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ing region (Figure 6C and Supplemental Figure S2B). We sequenced constructs or with truncated constructs containing the Vps21-bind

Moreover, neither GTP preference nor selectivity for Vps21 versus

and Vps8 or Vps8–200 were used as baits (Supplemental Figure S2A).

The same enhanced interaction was ob

able to differences in the expression of the Vps8 and Vps8–200 prey

served in reciprocal experiments where Vps21 was used as the prey

A and B; Woolford

Vps11 to residue 445 causes conditional growth defects, complete

S3) is functionally important and suggest that the Vps8–Vps21 interaction is, like the Vps3–Vps21 interaction, direct. Truncation of Vps11 to residue 445 causes conditional growth defects, complete defects in biosynthetic traffic to late endosomes and vacuoles (Figure 6A; Woolford et al., 1998), and class C vacuole morphology (Figure 6B). The VPS8–200 allele suppresses several of these vps11–445A phenotypes: it rescues growth defects, restores endosomal (but not vacuolar) trafficking, and results in a class B vacuole morphology similar to vps39Δ, vps41Δ, or ypt7Δ mutant cells (Figure 6, A and B; Woolford et al., 1998). In Y2H screens we found that Vps21 interacted more strongly with Vps8–200 than with wild-type Vps8 (Figure 6C). This difference in interaction strength was not attributable to differences in the expression of the Vps8 and Vps8–200 prey constructs (Figure 6D). The same enhanced interaction was ob

erved in reciprocal experiments where Vps21 was used as the prey and Vps3 or Vps8 were used as baits (Supplemental Figure S2A). Moreover, neither GTP preference nor selectivity for Vps21 versus other Rabs was modified by the Vps8–200 mutation, and the difference between Vps8 and Vps8–200 was seen with either full-length constructs or with truncated constructs containing the Vps21-binding region (Figure 6C and Supplemental Figure S2B). We sequenced VPS8–200 and identified a single missense substitution at an evolutionarily conserved residue (L830 to R), which is near the middle of the Vps21 interaction region (Figure 5A). Taken together, our results indicate that Vps21 interacts with both Vps3 and Vps8, and suggest a biochemical basis—enhanced Vps21 binding to Vps8—for the genetic suppression of vps11–445A trafficking phenotypes by VPS8–200.

The Vps11 CTD organizes Vps-C assembly and signaling

Among the Vps-C subunits, Vps11 had the densest web of interactions with other subunits, particularly within its CTD (Figure 7A). Notably, both the CORVET-specific subunit Vps3 and its HOPS-specific paralogue Vps39 interact via their conserved CTDs with the Vps11 CTD; these regions in Vps3 and Vps39 are functionally indispensable (Figures 4 and 5E; Supplemental Table S4). To dissect the functional architecture of the Vps11 CTD, we prepared a set of Vps11 truncation alleles (Figure 7A) and evaluated their in vivo phenotypes.

Each vps11 allele was prepared as a chromosomal replacement driven by the VPS11 promoter. vps11Δ null mutants exhibited defects in endosomal delivery indicated by pro-CPY accumulation and CPY secretion into the extracellular medium (Figure 7B and Supplemental Table S4), defects in vacuolar delivery indicated by accumulation of pro-ALP and pro-CPY, and defects in late endosome and vacuole fusion, indicated by class C vacuole morphology (Figure 7C). Four C-terminal Vps11 truncations (350Δ, 445Δ, 505Δ, and 735Δ) completely phenocopied the vps11Δ null allele (Figure 7, B and C; Supplemental Table S4). The total loss of function in the VPS11–735Δ mutant is particularly important, because this mutant lacks binding sites for Vps3 and Vps39 (Figure 7A).

In contrast to the more severe truncation mutants, a mutant lacking the Vps11 RING motif (vps11–926Δ) exhibited relatively selective defects: slightly impaired CPY processing and limited CPY secretion from the cell, but a severe defect in ALP processing (Figure 7B and Supplemental Table S4). These phenotypes are diagnostic of defective docking or fusion at the vacuole, but not at the endo

sistent with this interpretation, vps11–926Δ cells had class B fragmented vacuoles similar to vps39Δ or vps41Δ cells (Figure 7C). Because the Vps11 RING motif interacted weakly with other Vps-C subunits including Vps16 (Supplemental Table S2), communoprecipitation experiments were performed to ascertain whether HOPS is intact in vps11–926Δ mutant cells (Figure 5D). The results show that Vps11, Vps16, Vps18, and Vps41 all coprecipitate at least as efficiently with Vps11–926Δ as with wild-type Vps11. Together, these results indicate that the Vps11 RING is not needed for HOPS complex assembly but instead functions predominantly to regulate fusion at the vacuole, presumably through binding interactions with other HOPS subunits or interactions with other fusion factors.

FIGURE 5: CORVET subunits Vps3 and Vps8 are a Vps21 interaction module. (A) Summary of Vps3 and Vps8 Y2H interactions. The position of the L830R mutation in the VPS8–200 mutant is indicated. (B) Vps3 and Vps8 interact with Vps21-GTP in Y2H miniaarray experiments. The prey strains are indicated at left and were mated with strains containing either (middle) Vps3 or (right) Vps8 bait constructs. Yeast colony growth on synthetic medium (-Trp, -Leu, -His, +3-AT) indicates a positive Y2H interaction. Note that Vps3 and Vps8 interacted with Vps21 and the GTP-biased mutant Vps21-S21L but did not interact with any Rab other than Vps21 and did not interact with the GDP-biased Vps21-S21L mutant. (C) Native Vps3 in yeast cell lysates is selectively retained on a Vps21-GTP affinity resin. Note that most Vps3 in cell lysates is found within the CORVET complex. A similar pulldown, employing purified Vps3 rather than a cell lysate, is shown in Figure 4B. Cell lysates were prepared as described previously (Brett et al., 2008). TAP (tandem affinity purification)-tagged Vps3 was identified by immunoblotting with peroxidase-antiperoxidase (Upstate-Millipore, Billerica, MA). (D) Vps3 binds directly to Vps21-GTP. GST–Rab pulldown experiments were performed using purified recombinant His6–Vps3 and beads decorated with GST–Ypt7 or GST–Vps21 loaded with GDP or GTPγS by procedure 2 (Materials and Methods). (E) In vivo vacuole morphology of strains carrying Vps3 and Vps39 truncations that cannot interact with the Vps11 CTD. Vacuoles were stained using an FM4–64 pulse-chase (Materials and Methods).
The Vps11 CTD (residues 736–926) interacts with ~100-residue CTDs of high similarity in Vps3 and Vps39, so we asked whether the vps11–735Δ truncation might be complemented in trans by the missing Vps11-(736–1029) fragment. Remarkably, coexpression of this fragment rescued the vacuole morphology (Figure 8A) of both vps11–736Δ and vps11–936Δ cells, but it failed to rescue in vps11–350Δ, Δ–445, or -505A cells. Identical rescue results were obtained with expression of green fluorescent protein (GFP)-tagged or untagged Vps11-(736–1029). The CYP and ALP maturation defects of vps11–735A cells were also rescued by Vps11-(736–1029) (Figure 8B; Supplemental Table S4). We also evaluated the formation of multivesicular endosomes (MVEs) in vps11–735A cells expressing Vps11-(736–1029). GFP-CPS (carboxypeptidase S, Cps1) is sorted through the MVE pathway into the vacuole lumen. Defects in the formation of MVEs result in mis-sorting of GFP-CPS to the vacuole limiting membrane. The two-fragment complemented strain correctly sorted GFP-CPS to the vacuole lumen, indicating that MVE function is intact in these cells (Supplemental Table S4). Together, the results indicate that endolysosomal traffic is essentially normal in cells expressing both Vps11-(735A) and Vps11-(736–1029).

In every vps11 mutant strain tested, the GFP-Vps11-(736–1029) fragment exhibited a discrete subcellular localization (Figure 8A). In vps11–735A or vps11–926A cells, it localized predominantly to the vacuole limiting membrane and to perivacuolar foci, which are probably late endosomes. In vps11–350Δ, -445A, and -505A cells, GFP-Vps11-(736–1029) usually localized to a single bright punctate spot. In wild-type cells, however, GFP-Vps11-(736–1029) was exclusively cytoplasmic, indicating that the C-terminus of native Vps11 saturates the site(s) required for localization of Vps11-(736–1029). This was confirmed in coimmunoprecipitation experiments (Figure 8C). GFP-Vps11-(736–1029) co precipitated Vps16, Vps18, Vps33, and Vps41 in the truncation mutants (Figure 8C, lanes 2–6). In cells containing full-length Vps11, coisolation of other subunits with GFP-Vps11-(736–1029) was almost eliminated (Figure 8C, lanes 1), consistent with its diffuse localization in VPS11 cells (Figure 8A). Together, these data show that Vps11 can be split in two and retain nearly its full function. However, in the absence of a central domain (e.g., when 1–505 and 736–1029 are coexpressed), the phenotype is equivalent to a vps11A null, even though a HOPS complex of apparently normal composition is assembled (Figure 8C, lanes 4 and 5). Thus the central domain (residues 351–735) is also crucial for Vps11 function. The split N- and C-terminal fragments of Vps11 probably assemble on Vps18 (Figure 7A). This conclusion is supported by our finding that the integrity of the entire wild-type complex is lost when Vps18 is deleted (see Figure 3E, lanes 4). Taken together, these experiments underscore the complex and critical functions of the Vps11 CTD in both HOPS and CORVET function.

**DISCUSSION**

Our main conclusions are that the CORVET subunits Vps3 and Vps8, and the HOPS subunits Vps39 and Vps41, form discrete complex-specific subassemblies that function in Rab recognition. The activities of these subassemblies are coordinated with the core through contacts between the CTDs of Vps3 or Vps39 and the CTD of Vps11. The core subunits Vps16 and Vps33 also form a stable subassembly. Vps33 belongs to the Sec1-Munc18 (SM) family of cofactors (Sudhof and Rothman, 2009), which are universally required for SNARE-mediated membrane fusion, so Vps16 and Vps33 likely link the Rab signaling activities of HOPS and CORVET to SNARE-mediated membrane fusion.

In early studies, HOPS was identified as an effector of Ypt7-GTP (Price et al., 2000; Seals et al., 2000) and the Vps39 subunit was...
reported to be a Ypt7 GEF (Wurmser et al., 2000; Binda et al., 2009). Vps41 binds Ypt7-GTP directly (Figure 3B), which explains why Vps41 is essential for stable binding of the HOPS holocomplex to Ypt7-GTP (Brett et al., 2008). However, although Vps39 shows a lack of nucleotide specificity in Ypt7 binding, as might be expected for a GEF, highly purified preparations of Vps39 or Vps39–860 failed to exhibit any nucleotide exchange activity in fluorescence and radioisotopic assays that readily detected the activities of bona fide GEFs (Nordmann et al., 2010; Ostrowicz et al., 2010; Lobingier, Brett, and Merz, unpublished results). Instead, the major Ypt7 GEF activity appears to reside within the Ccz1/Mon1 complex (Kucharczyk et al., 2000; 2001; Nordmann et al., 2010). The Vps39 CTD is dispensable for Vps39 binding to Ypt7 and Vps41 (Figures 1 and 2D; Supplemental Table S2), and for binding of HOPS to Ypt7-GTP (Brett and Merz, unpublished results). The Vps39 CTD is, however, essential for Vps39 function in vivo, because its deletion phenocopies a vps39Δ null mutant (Figure 4 and Supplemental Table S2), and for binding of HOPS to Ypt7-GTP but also the ratio of Ypt7-GTP to Ypt7 GDP. The functional consequences of the divergent Rab-binding modalities of HOPS and CORVET are not yet understood.

As membrane traffic traverses successive endolysosomal organelles, the Rab5 orthologue Vps21 is replaced by the Rab7 orthologue Ypt7 (Figure 9). CORVET and HOPS, operating in concert with the GEFs Vps9 and Ccz1-Mon1, appear to control this Rab cascade (Rink et al., 2005; Vonderheit and Helenius, 2005; Peploowska et al., 2007). Vps11 appears to be a central scaffold linking Rab signaling to Vps-C outputs, including the SNARE machinery, which mediates fusion at the endosome and vacuole. In either case, the Vps11 CTD is now the prime candidate for the location at
mids were constructed either by conventional cloning or by gap-repair methods. Y2H assays
Two-hybrid vectors were constructed using gap-repair and homologous recombination (Uetz et al., 2000). Prey domains were cloned into the plasmid pOAD and transformed into the yeast strain PJ-9-4a. Bait domains were cloned into pOBD-2 and transformed into PJ69-4a. Clonal isolates were obtained and verified by PCR and in many cases by DNA sequencing. Interaction tests were performed by mating haploid strains containing the bait and prey vectors. Liquid cultures of the bait and prey strains were grown in selective media then mixed in 96-well plates and pinned to YPD (yeast extract peptone dextrose) plates supplemented with adenine using a 48-spoke inoculating manifold. The mating plates were grown at 30°C overnight, then diploid cells were selected by replica plating onto medium lacking tryptophan and leucine and supplemented with adenine. Diploid colonies were grown at 30°C for 2 d, then tested for Y2H interactions by replica plating to medium lacking

which switching between the HOPS and CORVET Vps-C configurations might occur. Consistent with the idea that the Vps11 CTD is a key element in this cascade, the C-terminal RING domain of Vps11 appears to function predominantly in fusion at the vacuole (Figure 7 and Supplemental Table S4). Moreover, we recently isolated new vps11 alleles bearing mutations solely in the Vps11 CTD and exhibiting selective defects in traffic through late endosomes, to the vacuole, or both (Nickerson, Fawcett, and Merz, unpublished results). We are now working to understand how Rab signaling through Vps-C complexes drives fusion and other "output" processes (Nickerson et al., 2009) and how Vps-C complexes signal in time and space to transport cargoes derived from early endosomes, the late Golgi, autophagosomes, and cytoplasm-to-vacuole transport (CVT) vesicles through the endolysosomal system.

MATERIALS AND METHODS
Strains and plasmids
Yeast strains used are described in Supplemental Table 1. Plasmids for Y2H screening are listed in Supplemental Tables S2 and S3. Plasmids were constructed either by conventional cloning or by gap-repair methods.

Y2H assays
Two-hybrid vectors were constructed using gap-repair and homologous recombination (Uetz et al., 2000). Prey domains were cloned into the plasmid pOAD and transformed into the yeast strain PJ-9-4a. Bait domains were cloned into pOBD-2 and transformed into PJ69-4a. Clonal isolates were obtained and verified by PCR and in many cases by DNA sequencing. Interaction tests were performed by mating haploid strains containing the bait and prey vectors. Liquid cultures of the bait and prey strains were grown in selective media then mixed in 96-well plates and pinned to YPD (yeast extract peptone dextrose) plates supplemented with adenine using a 48-spoke inoculating manifold. The mating plates were grown at 30°C overnight, then diploid cells were selected by replica plating onto medium lacking tryptophan and leucine and supplemented with adenine. Diploid colonies were grown at 30°C for 2 d, then tested for Y2H interactions by replica plating to medium lacking
tryptophan, leucine, and histidine, supplemented with adenine and various concentrations of 3-amino-1,2,4-triazole (3-AT), ranging from 1.5 to 25 mM. The use of low-copy vectors (pOAD and pOBD-2), and the assay of histidine auxotrophy in the presence of 3-AT at $>1$ mM, together almost totally suppresses false-positive signals in Y2H experiments (Braun et al., 2009; Chen et al., 2010). After 5 d at 30°C, plates were scored for growth. Bait–prey interactions were scored as follows: very strong (3), strong (2), weak (1), or none (0). Mean scores are presented; in a large majority of cases these mean scores represent results from two or three independent interaction tests. In many cases, positive hits from the Y2H surveys were also subjected to further rounds of screening, which are not reflected in the survey scores presented in Supplemental Tables S2 and S3.

### Protein purification

His$_{60}$- and GST-fused Vps-C subunits were expressed in BTI-Tn5B1–4 (High-5) insect cells as described (Brett et al., 2008). Baculovirus vectors were constructed and propagated using the Bac-to-Bac system as recommended by the vendor (Invitrogen, Carlsbad, CA). Suspension-adapted or monolayer insect cells were infected with passage-3 or passage-4 virus, grown for 3 d at 27°C, and harvested. All subsequent steps were performed at 4°C or on ice. Cells were resuspended in lysis buffer (50 mM HEPES-[4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid], 400 mM NaCl, 5 mM 2-mercaptoethanol, pH 8.0) and various concentrations of 3-amino-1,2,4-triazole (3-AT), ranging from 1.5 to 25 mM. The use of low-copy vectors (pOAD and pOBD-2), and the assay of histidine auxotrophy in the presence of 3-AT at $>1$ mM, together almost totally suppresses false-positive signals in Y2H experiments (Braun et al., 2009; Chen et al., 2010). After 5 d at 30°C, plates were scored for growth. Bait–prey interactions were scored as follows: very strong (3), strong (2), weak (1), or none (0). Mean scores are presented; in a large majority of cases these mean scores represent results from two or three independent interaction tests. In many cases, positive hits from the Y2H surveys were also subjected to further rounds of screening, which are not reflected in the survey scores presented in Supplemental Tables S2 and S3.

### Protein sorting and vacuole morphology

Mutant yeast strains were propagated at room temperature to minimize selection for suppressor mutations. Liquid cultures were grown to log phase (OD$_{600}$ = 0.6–1.0) in complex or synthetic media at 26°C. Cell pellets were chilled on ice and washed in 10 mM Na$_3$3-AT at $>1$ mM, together almost totally suppresses false-positive signals in Y2H experiments (Braun et al., 2009; Chen et al., 2010). After 5 d at 30°C, plates were scored for growth. Bait–prey interactions were scored as follows: very strong (3), strong (2), weak (1), or none (0). Mean scores are presented; in a large majority of cases these mean scores represent results from two or three independent interaction tests. In many cases, positive hits from the Y2H surveys were also subjected to further rounds of screening, which are not reflected in the survey scores presented in Supplemental Tables S2 and S3.

### Immunoprecipitation

Cell lysates were prepared, and immunoprecipitations were performed as described (Angers and Merz, 2009).

### Protein purification

His$_{60}$- and GST-fused Vps-C subunits were expressed in BTI-Tn5B1–4 (High-5) insect cells as described (Brett et al., 2008). Baculovirus vectors were constructed and propagated using the Bac-to-Bac system as recommended by the vendor (Invitrogen, Carlsbad, CA). Suspension-adapted or monolayer insect cells were infected with passage-3 or passage-4 virus, grown for 3 d at 27°C, and harvested. All subsequent steps were performed at 4°C or on ice. Cells were resuspended in lysis buffer (50 mM HEPES-[4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid], 400 mM NaCl, 5 mM 2-mercaptoethanol, 10% glycerol, 0.5% Triton X-100, supplemented with pH 7.8, 400 μM phenylmethylsulfonyl fluoride, 1 μg/ml leupeptin, 1 μg/ml pepstatin, 100 μM Pefabloc SC [Roche, Mannheim, Germany], and 20 mM imidazole). The cells were lysed with detergent and brief sonication, and the lysate was clarified by centrifugation (18,000 × g, 25 min). Clarified lysates were incubated with glutathione sepharose 4B (GE Healthcare, Piscataway, NJ) for 2 h, and resin was subsequently washed three times with the lysis buffer and eluted with lysis buffer supplemented with 10 mM reduced glutathione. GST eluates were then incubated with pre-equilibrated Ni$^{2+}$-NTA resin (GE Healthcare) for 1 h, washed three times with lysis buffer supplemented with 35 mM imidazole, eluted with lysis buffer supplemented with 500 mM imidazole, and analyzed by SDS–PAGE.

### Rab pulldowns

GST-Rab fusion proteins were cloned, expressed, and purified as described (Brett et al., 2008). Nucleotide loading of Rab proteins was performed either (procedure 1) as described (Brett et al., 2008) or (procedure 2) by washing the resin once with nucleotide loading buffer (50 mM Tris, 200 mM NaCl, 7.5 mM EDTA, 5 mM 2-mercaptoethanol, pH 8.0), then incubating the resin with the nucleotide loading buffer and 20-fold molar excess of nucleotide for 2 h at room temperature. The reaction was terminated with an excess of quenching buffer (50 mM Tris, 200 mM NaCl, 15 mM MgCl$_2$, 5 mM, pH 8.0). The resin was washed three times with reaction buffer (50 mM Tris, 200 mM NaCl, 5 mM MgCl$_2$, 5 mM 2-mercaptoethanol, pH 8.0). Cell lysates or purified Vps-C subunits were incubated with the resin in reaction buffer for 1 h at 4°C, and the resin was then washed with reaction buffer three times. Bound protein was eluted from the beads in reaction buffer supplemented with 10 mM reduced glutathione, pH 8.0 (final), and the eluates were analyzed by SDS–PAGE.

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