A Cell-free Assay Allows Reconstitution of Vps33p-dependent Transport to the Yeast Vacuole/Lysosome

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Abstract. We report a cell-free system that measures transport-coupled maturation of carboxypeptidase Y (CPY). Yeast spheroplasts are lysed by extrusion through polycarbonate filters. After differential centrifugation, a 125,000-g pellet is enriched for radiolabeled proCPY and is used as “donor” membranes. A 15,000-g pellet, harvested from nonradiolabeled cells and enriched for vacuoles, is used as “acceptor” membranes. When these membranes are incubated together with ATP and cytosolic extracts, ∼50% of the radiolabeled proCPY is processed to mature CPY. Maturation was inhibited by dilution of donor and acceptor membranes during incubation, showed a 15-min lag period, and was temperature sensitive. Efficient proCPY maturation was possible when donor membranes were from a yeast strain deleted for the PEP4 gene (which encodes the principal CPY processing enzyme, proteinase A) and acceptor membranes from a PEP4 yeast strain, indicating intercompartmental transfer. Cytosol made from a yeast strain deleted for the VPS33 gene was less efficient at driving transport. Moreover, antibodies against Vps33p (a Sec1 homologue) and Vam3p (a Q-SNARE) inhibited transport 90%. Cytosolic extracts from yeast cells overexpressing Vps33p restored transport to antibody-inhibited assays. This cell-free system has allowed the demonstration of reconstituted intercompartmental transport coupled to the function of a VPS gene product.

Key words: carboxypeptidase Y • lysosome • membrane fusion • Saccharomyces cerevisiae • vacuole

The secretory and endocytic pathways in eukaryotic cells comprise a series of intercompartmental transport events. The directed movement of protein and lipid from the endoplasmic reticulum to the plasma membrane or from the plasma membrane to the lysosome necessarily engages transfer between multiple organelles. In most cases, carrier vesicles mediate the traffic of protein and lipid cargo from one subcellular compartment to another.

Two integral approaches, genetics and biochemistry, continue to contribute preeminently in elucidating the molecular details of vesicle-mediated transport in the secretory and endocytic pathways. Mutant isolation screens and selections in such diverse organisms as Drosophila melanogaster (Swanson et al., 1998), Caenorhabditis elegans (Brenner, 1974), and Saccharomyces cerevisiae (Novick and Schekman, 1979; Deshaies and Schekman, 1987) have not only uncovered hundreds of genes, but also helped reveal the ubiquitous nature of secretion and endocytosis among eukaryotic organisms. These genetic efforts were pioneered in yeast and include >20 secretion (sec) mutants (Novick et al., 1980), >40 mutants defective for vacuolar protein sorting (vps) (Robinson et al., 1988; Rothman et al., 1989), and >10 mutants defective for endocytosis (end) (Raths et al., 1993; Mann and Riezman, 1994; Mann et al., 1995). Biochemical efforts using reconstitution assays have also uncovered many proteins involved in vesicle-mediated transport. These assays are focused on anterograde and retrograde transfer between the ER and Golgi complex (Balch et al., 1988; Baker and Schekman, 1989; Balch, 1989; Spang and Schekman, 1998), intra-Golgi transport (Balch et al., 1984), fusion of secretory vesicles with the plasma membrane (Martin and Kowalchyk, 1997), and recycling from late endosomes to the trans-Golgi network (Goda and Pfeffer, 1988).

Despite the progress made in other vesicle-mediated events, a poorly understood intercompartmental step in eukaryotic cells continues to be transfer of proteins from prelysosomal compartments (PLC) to the lysosome. The PLC, or late endosome, plays a pivotal role in protein trafficking since it is the organelle where the secretory and endocytic pathways converge. Resident lysosomal proteins

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pass through the PLC after being sorted away from secretory proteins in the trans-Golgi network (Pfeffer, 1991). Similarly, cell surface proteins destined for degradation in the lysosome pass through the late endosome after endocytosis. Thus, eukaryotic cells must blend a variety of events for proper sorting, targeting, and delivery of proteins from the PLC. Several factors have concealed the molecular details of protein transport from the PLC to the lysosome. Perhaps the greatest source of confusion about the PLC revolves around whether carrier vesicles transport material to lysosomes or if the PLC undergoes a maturation process, changing into a lysosome (Futter et al., 1996). Evidence has been presented and interpreted to support both models. Even a hybrid of the vesicle-shuttle vs maturation models is suggested for PLC to lysosome transport in macrophages (Racoon and Swanson, 1993).

Saccharomyces cerevisiae contains not only a lysosome-like vacuole, but also a PLC-like prevacuolar compartment (PVC) (Raymond et al., 1990; Davis et al., 1993; Vida et al., 1993). The PVC is central to the function of nearly all VPS genes. The vps mutants either cause defects in anterograde or retrograde sorting/transport between the late Golgi complex and the PVC or in transport between the PVC and the vacuole (Bryant and Stevens, 1998). Subsequent studies have focused on uncovering the function of VPS genes and are beginning to reveal aspects on the biochemistry of transport to and from the PVC because several of the gene products have biochemical activity in vitro. For example, the VPS1 gene product is a dynamin-like protein that can bind and hydrolyze (Herman and Emr, 1990; Stack et al., 1993). However, coupling the catalytic activity of a VPS gene product to intercompartmental protein transport in a reconstituted assay remains elusive. A permeabilized yeast spheroplast system (Vida et al., 1990) has not allowed analysis of VPS gene product function because the membranes are not depleted for any VPS protein that has been examined.

In this report, we describe an intercompartmental protein transport assay using partially purified organelles. This cell-free system measures proteolytic maturation of soluble vacuolar proenzymes such as carboxypeptidase Y and proteasome A after transfer from the PVC to the vacuole. The reaction is sensitive to membrane dilution, requires ATP, and cytosol. Importantly, cytosol made from a vps33Δ strain is deficient at stimulating transport in the new cell-free system. Furthermore, antibody raised against Vps33p can inhibit the assay >90% and cytosolic extracts made from strains overexpressing Vps33p can restore this inhibition. Thus, we have developed a transport-coupled assay for the function of a VPS gene product.

Materials and Methods

Media

Yeast strains were maintained on YPD media (1% yeast extract, 2% peptone, 2% dextrose, and 2.5% bacto-agaro). Liquid media for radiolabeling and plasmid maintenance was Wickerham’s minimal proline (WIMP) (Wickerham, 1946) media supplemented with 0.5% yeast extract.

Strains and Plasmsids

The yeast strains used in this study include BGY 3300 (Gerhardt et al., 1990) [α ura3-52 leu2-3,112 his3-A2000 sop2-19 lys2-801 sec2-19 vps33Δ·HIS5, SEY 6210 (Robinson et al., 1998) TVY 614 MAT α ura3-52 leu2-3,112 his3-A2000 sop2-19 prc1Δ·HIS5 prb1Δ·hisG pep4Δ·LEU2; and TVY 1 (Gerhardt et al., 1990) MAT α ura3-52 leu2-3,112 his3-A2000 sop2-19 prc1Δ·HIS5 pep4Δ·LEU2. Several constructs were made to put the VPS33 gene under control of the glycerolaldehyde-3-phosphate dehydrogenase promoter (P GDP 1).], First, site-directed mutagenesis (D eng and N ickolf, 1992) was used to place a BamHI site at the second codon of the VPS33 gene and a SalI site ~200 bp from the stop codon in pPRP33-100, which contains the complete VPS33 gene in pBluescript KS (Stratagene, Inc.). The resulting plasmid (pBG33BbSe) was digested with BamHI and SalI and subcloned into pGPD 426 (Mumberg et al., 1995) to generate pGPD·BbSe-2. A six-histidine tag was placed at the NH₂-terminus of VPS33 with the PCR using 5'-TACGATCCATGAGAAGCTGATCATCCAATCATCAAATCAGGTTTGATTTGGAATCAG-3' as the forward primer and 5'-CAAAAAATGCTTCTTGGTCAAGAAG-3' as the reverse primer. The amplicon was digested with BamHI and CiaI and subcloned into pGPD·BbSe-2 to generate pGPD·HIS633-2.

Antibody Production

Two previously described tpe-VPS33 fusion constructs (Banta et al., 1990) were expressed in E. coli and the insoluble fraction of cell lysates was prepared (Kerner et al., 1991). A titer SDS-PAGE, the tpe-Vps33 fusion proteins were cut out of the gel and the gel slice used as antigens in rabbits at Cocalico Biologicals, Inc. Antiserum against Vam3p was a gift from Professor Trung Dinh at the University of California, Irvine. A protein A-Sepharose column was used to purify total IgG from pre- and immune sera. Vps33p rabbit sera.

Preparation of Cytosol

Yeast strain TVY 614 was grown at 30°C in YPD (with 5% glucose) to an OD₆₀₀ of 4-6 (usually 800-1,600 total OD₆₀₀ units of cells were used). The cells were harvested with centrifugation at 1,500 g for 15-30 min. After harvesting the cells (1,500 g for 5 min), they were rinsed once with 25 ml of sterile water and then agitated for three 30-s intervals in a Mini Bead-Beater (BioSpec Equipment Co., Inc.) for 15 min at 4°C. All tubes were subjected to centrifugation at 1,500 g for 10 min. The supernatants were pooled with the first supernatant and subjected to centrifugation in a Beckman TLA 100.3 rotor at 50,000 rpm (z 103,000 g) for 30 min. The supernatants were further centrifuged at 100,000 g for 1 h. The supernatant was removed, and the pellet was resuspended in the same volume of water and then agitated for three 30-s intervals in a Mini Bead-Beater (BioSpec Products, Inc.) at 4°C. A ll tubes were subjected to centrifugation at 15,000 g for 2 min. The supernatant was removed from each tube and the pellet was rinsed with 1 ml of TB, agitated briefly on a vortex mixer, and subjected to centrifugation at 15,000 g for 2 min. The second supernatant was pooled with the first supernatant and subjected to centrifugation in a Beckman TLA 100.3 rotor at 50,000 rpm (~103,000 g average) for 10 min. The supernatant was removed, dispensed into small aliquots, and snap-frozen in liquid nitrogen. The protein concentration of all cytosolic extracts ranged from 25–50 mg/ml.

Cell Preparation for Donor and Acceptor Membranes

All steps are reported for the preparation of donor membranes from 25 OD₆₀₀ units of cells. When preparing more than this amount, volumes were scaled up proportionally. Yeast cells were grown in Wickerham's minimal proline (Wickerham, 1946) media supplemented with 0.5% yeast extract at 30°C to an OD₆₀₀ of 0.5-1.2. The cells were harvested with centrifugation at 1,500 g for 5 min and washed once with 25 ml of sterile distilled water. A sterile water washing the cells (1,500 g for 5 min), they were resuspended in 2.5 ml of 0.1 M Tris-HCl pH 9.4, plus 10 mM EDTA and incubated with shaking at 30°C for 15–30 min. A sterile water washing the cells (1,500 g for 5 min), they were resuspended in Wickerham’s minimal proline media
containing 0.2% glucose, 1.0 M sorbitol and 25 mM Tris-HCl, pH 7.5, to a total volume of 1 ml. The cells were converted to spheroplasts using 25 µg of Zymolyase 100T (Seikagaku Kogyo Co., Ltd) and 0.5% (vol/vol) glucose
lase (NE N Du Pont) with gentle agitation for 20-30 min at 30°C. The cells were harvested (1,500 g for 5 min) and resuspended in 2.5 ml of Wickerham’s minimal proline media containing 2% glucose and 1.0 M sorbitol. The cells were incubated with gentle shaking at 30°C for 15 min and then pulse-labeled with Tran3-S-label (ICN, Inc.) at 200 µCi/ml for 5 min. A filter paper was washed with methanol (5 mM final), cyclohexane (1 mM final), and yeast extract (0.5% final) were added and the cells were chased for 2 min. A filter paper that was transferred to 10 ml of ice-cold 1.0 M sorbitol, 20 mM Hepes-KOH, 150 mM potassium acetate, and 5 mM magnesium acetate (freezing buffer) and incubated on ice for 5 min. The cells were harvested (1,500 g for 5 min) and washed two times with 1 ml freezing buffer. The washed samples were centrifuged (1.7 ml), and the supernatant (of the microcentrifuge tube) was removed at 16,000 g at 4°C. An approximatively 45 µl of freezing buffer was added to the washed cells. They were then resuspended, placed in a Nalgene™ 1-ml cryofreeze tube with isopropanol (prechilled to 4°C), and the cryofreeze was incubated at −70°C for at least 45 min. AII steps to prepare cells for acceptor membrane were identical to the above steps for donor membranes except for the following changes. Rich media,YPD, was used instead of W1/M P and after cell wall removal, the spheroplasts were incubated at 30°C (at 10 OD650 units/ml) in YPD plus 1.0 M sorbitol for 60 min without shaking.

Preparation of Donor and Acceptor Membranes

If using frozen cells, they were thawed in a 25°C circulating water bath for 1 min and placed on ice. 600 µl of 0.6 M Sorbitol with 5 mM HEPES-KOH, pH 7.5 (lysis buffer) was added and the cells were resuspended thoroughly. The cells were harvested by centrifugation for 1 min at 16,000 g and then resuspended to 8 OD650 units/ml in lysis buffer. The resuspended cells were pushed through a 13-mm polycarbonate filter (Nucleopore™; Corning) with 3-µm pores using a 3-ml syringe. The filter effluent was subjected to centrifugation at 440 g for 5 min to generate a P1 pellet and S1 supernatant fraction. The S1 supernatant was subjected to centrifugation at 15,000 g for 10 min to generate a P2 pellet (acceptor membranes) and S2 supernatant fraction. The S2 supernatant was subjected to centrifugation at 125,000 g for 10 min to generate a P3 pellet (donor membranes) and S3 supernatant fraction.

Cell-free Assays

Radiolabeled donor membranes and nonradiolabeled acceptor membranes were resuspended in TB. Standard conditions for assays were 50 µl total volume containing donor membranes (equivalent to 5 OD650 units of cells), 100-125 µg of acceptor membranes, 1 mM ATP, 40 mM creatine phosphate, 0.2 mg/ml creatine kinase, and 5 mg/ml cytosol. All reactions were assayed on ice and then incubated at 25°C in a circulating water bath for 60 min. To stop the reactions, 3 µl of 100 mM PMF, 25 µl of 8.0 M urea, 5% SDS, and 5% NP-40 was added and they were boiled for 5 min. All SDS was removed for immunoprecipitation, SDS-PA G E, and autoradiography as previously described (Vida et al., 1990). All samples were digitized with a Epson Expression 636 flatbed scanner and quantitation of the protein bands was done using NIH Image software (v 1.61).

Microscopy

All light microscopy images were obtained as previously described (Gerhardt et al., 1998). Yeast cells were stained with dichloroarboxyfluorescein diacetate and FM 4-64 (Molecular Probes, Inc., Eugene, OR) as previously described (Vida and Emr, 1995). For electron microscopy, sample membranes pellets were fixed as previously described (Vida et al., 1993). A filter fixation, the samples were washed and treated with Miliopore-filtered, cacodylate-buffered 0.1% tannic acid, postfixed with buffered 1% osmium tetroxide, and stained en bloc with Millipore-filtered aqueous 1% uranyl acetate, postfixed with buffered 1% osmium tetroxide, and stained en bloc with Millipore-filtered aqueous 1% uranyl acetate. The samples were dehydrated in increasing concentrations of ethanol, infiltrated, and embedded in microcentrifuge tubes in Spurr’s low viscosity medium. The samples were then polymerized in a 60°C oven for 2 d. Ultrathin sections were cut in an LKB Nova ultramicrotome (Leica), stained with uranyl acetate and lead citrate in an LKB Ultratome, and then examined in a J EOL 1200-EX transmission electron microscope at an acceleration voltage of 80 kV.

Results

A New Method to Lyse Yeast Spheroplasts

The usefulness of cell-free assays cannot be overstated in their contribution to our understanding of mechanisms in protein transport, secretion, and endocytosis. Since the development of a permeabilized cell assay for transport to the yeast vacuole (Vida et al., 1990), a longstanding goal has been to establish an intercompartmental transport assay using separate subcellular fractions in a cell-free system. The previous permeabilized cell assay is not cell-free. Because the conditions maintain organelle structural integrity while simultaneously dissociating membrane aggregates (Vida et al., 1993). However, these membranes were routinely devoid of transport activity (data not shown). We solved this problem and used another lysis method using polycarbonate filters with a defined pore size to gently shear away the plasma membrane.

The technique of passing cells through a small orifice to generate a crude lysate from shear forces was first used for mammalian cells. For example, cell homogenates have been prepared from PC12 cells by passing cell suspensions 15 times through a narrow clearance (10 µm) stainless steel ball homogenizer (Martin and Kowalchyk, 1997). Stainless steel ball homogenizers have been instrumental in reconstituting several steps in the secretory pathway such as fusion of secretory vesicles with the plasma membrane and ER to Golgi transport (Balch and Rothman, 1985). Rather than use a steel ball homogenizer, we used polycarbonate filters to shear yeast spheroplasts. Intact spheroplasts were suspended with an osmotic support of 0.6 M sorbitol giving them a diameter in the range of 5-8 µm. The cells were then forced through a 3-µm polycarbonate filter from a syringe (Fig. 1A). Typically, in a single pass through the filter, >98% of the cells lysed to generate a crude homogenate.

We performed centrifugation techniques on crude lysates after extrusion through polycarbonate filters and examined each supernatant and pellet fraction for marker proteins. Simple differential centrifugation allows separation of a variety of yeast organelles and membranes (Bowser et al., 1992; Vida et al., 1993; Rieder and Emr, 1997). Subjecting the lysate to 440 g produced a pellet (P1) containing insignificant amounts of CPY, PrA, and A LP (Fig. 1B). A plot not shown in this experiment, <2% of a cytosolic marker protein (glucose 6-phosphate dehydrogenase) fractionated with the P1 pellet, indicating that polycarbonate filter lysis was very efficient. Subjecting the postnuclear supernatant (S1) to 15,000 g produced a pellet (P2) containing ~5% of the total p1CPY, ~50% of p2CPY, and ~95% of the sedimentable mCPY, suggesting...
the presence of ER, early Golgi membranes (p1CPY), and intact vacuoles (mCPY, Fig. 1 B). The majority (>90%) of the p2CPY in the postvacuolar supernatant (S2) was found in the pellet (P3) after subjecting the postvacuolar supernatant to centrifugation at 125,000 g. Although the latGolgi complex marker, Kex2p, also fractionated with the 125,000-g membrane pellet, <10% of p2CPY cofractionated with Kex2p activity on sucrose gradients (data not shown). This suggested that very little p2CPY was localized in the late Golgi complex and most likely resided in the PVC as previously shown using comparable pulse-chase radiolabeling conditions (Vida et al., 1993). Overall, these fractionation characteristics of membranes obtained from extrusion through polycarbonate filters were very similar to those observed after dissociating permeabilized cells (Vida et al., 1993).

Various steps from the filter lysis procedure were also examined with microscopy. To follow the vacuole, we prestained yeast cells with FM 4-64 and CDCFDA. As expected, the P1 pellet was devoid of unbroken cells and was enriched in cell wall remnants (Fig. 2). As expected from the marker protein analysis, the P2 pellet was enriched in intact vacuoles, since many FM 4-64–stained membranes containing CDCFDA were observed (Fig. 2). In contrast, the 125,000-g P3 pellet was devoid of vacuoles and instead was enriched for very small particulate structures. Importantly, if cells were stained with FM 4-64 at 15°C, many of the small particulate structures in the 125,000-g pellet exhibited fluorescence (Fig. 2, inset). Additionally, membrane fluorescence in the 15,000-g P2 pellet was markedly reduced at 15°C (Fig. 2, inset). Since FM 4-64 is kinetically trapped in prevacuolar compartments at 15°C (Vida et al., 1993), the membrane fluorescence in the 125,000-g pellet suggests that these differential centrifugation conditions separated vacuoles from prevacuolar compartments. Although not shown in this experiment, the P1 pellet also was enriched with intact nuclei after first staining yeast cells with DNA dyes (i.e., 4′,6-diamidino-2-phenylindole, DAPI). The P2 pellet was enriched for mitochondria after first staining cells with the mitochondrial vital dye 2,4-(4-(dimethylamino)styryl)-N-methylpyridinium iodide (DASPMI). We also examined the P2 and P3 pellets with electron microscopy. The P2 pellet comprised numerous electron-dense 1,000–1,500-nm membrane-delineated structures, which was consistent with the size expected for vacuoles (data not shown). In contrast, the P3 pellet was devoid of the relatively large, electron-dense membranes and instead was composed of 50–100 nm and 250–400 nm membrane-delineated structures (data not shown).

Reconstitution of p2CPY Maturation after Mixing the Donor and Acceptor Membrane Pellets

The polycarbonate filter lysis technique and simple differ-
Figure 2. Light microscopy of cell-free membrane pellets after polycarbonate filter lysis. Wild-type yeast cells (as in Fig. 1) were first stained with FM 4-64 (15-min pulse, 45-min chase) followed with dichlorocarboxyfluorescein diacetate (15 min at pH 4.0) to mark the vacuole membrane and lumen, respectively. The double-stained cells were then enzymatically converted to spheroplasts at 30 or 15°C (as indicated). The spheroplasts were extruded through a polycarbonate filter with 3-μm pores (as described in Fig. 1A). The lysate, P1, P2, and P3 pellets (as indicated) were then examined under a light microscope with differential interference contrast (DIC), phase contrast, and epifluorescence optics using a FITC and Texas red filter set (as indicated). The fluorescence images were digitally overlaid for a composite. The cells at 15°C (inset) were stained with just FM 4-64 for a 30-min pulse. Bar, 5 μm.
Wild-type yeast spheroplasts were radiolabeled (as in Fig. 1 B). The cells were subjected to lysis through a polycarbonate filter and cytosol. With lysis buffer and reharvested before incubation with ATP and 11, both donor and acceptor membranes were washed once with subsequent differential centrifugation to generate a 125,000 g P3 donor membrane pellet. The same yeast strain was used to make a 15,000 g, P2 acceptor membrane pellet from nonradiolabeled spheroplasts. The radiolabeled donor membranes (from ~5 × 10^7 spheroplasts per reaction) were incubated at 25°C for 60 min with various combinations of buffer, ATP (plus a regeneration system), cytosol (5 mg/ml), and acceptor membranes (~100 µg) in a total volume of 50 µl, as indicated. All reactions were sequentially immunoprecipitated for CPY and PrA, subjected to SDS-PAGE, and autoradiography. For the reactions in lanes 10 and 11, both donor and acceptor membranes were washed once with lysis buffer and reharvested before incubation with ATP and cytosol.

Characteristics of the Cell-free Assay

The characteristics of cell-free assays with P3 donor membranes and P2 acceptor membranes were examined to determine if they suggested that the reaction was intercompartmental. The first characteristic that we examined was dilution sensitivity. Normally, reactions were carried out in a 50-µl volume with the efficiency of p2CPY maturation ranging from 35 to 55%. To test the effect of dilution, the reaction volume was increased to dilute the concentration of donor/acceptor membranes while the concentration of ATP and cytosol was maintained at a constant level. An exponential decrease in p2CPY maturation efficiency was observed concomitant with an incremental increase in the reaction volume (Fig. 4 A). For example, a sixfold decrease in efficiency (38% vs 6%) took place with a 10-fold increase in reaction volume (from 50 to 500 µl). This suggested that the concentration of donor and acceptor membranes had a critical threshold for optimal reconstitution of p2CPY maturation. The second characteristic that we examined of the cell-free assay was the reaction kinetics. A prominent lag period was observed in the first 15–20 min (Fig. 4 B). A linear phase followed for the next 20 min and reached a plateau between 40 and 60 min (Fig. 4 B). Although not shown in this experiment, an increase in p2CPY maturation did not occur after a further 60 min incubation. This kinetic analysis suggested that a rate-limiting event(s) occurred early in the incubation, which might be the formation of a transport intermediate. The third characteristic that we examined of the cell-free assay was its temperature dependence. The maturation of p2CPY was undetectable when the incubation was carried out at 0 or 5°C (Fig. 4 C). The optimal efficiency occurred between 20 and 30°C and sharply tapered off at temperatures above 30°C (Fig. 4 C). Overall, the dilution sensitivity, kinetics, and temperature dependence of this new cell-free assay for p2CPY maturation indicated a complex event(s) was reconstituted after incubating P3 donor membranes and P2 acceptor membranes in the presence of ATP and cytosol.

The Cell-free Assay Reconstitutes Intercompartmental Protein Transport

To truly determine if this new cell-free assay reconstituted
intercompartmental protein transport, we performed reactions where the donor and acceptor fractions were prepared from yeast strains defective in vacular processing enzymes. A hallmark of most cell-free intercompartmental protein transport assays is using donor membranes deficient in the activity that marks the transport event. Two proteases are responsible for cleaving the propeptide from p2CPY, proteinase A (PEP4 gene) and proteinase B (PRB1 gene). In yeast strains mutant for the PEP4 gene (pep4-1, or pep4Δ), p2CPY travels to the vacuole but is not processed to the mature form of the protein (Stevens et al., 1982). In prb1 mutant strains, p2CPY also travels to the vacuole but instead of remaining unprocessed, active proteinase A (which can autoactivate) cleaves away a portion of the propeptide (Knop et al., 1993).

We took advantage of CPY processing characteristics in vivo (Fig. 5A) using PEP4 and pep4Δ yeast strains as a source of both donor and acceptor membranes in vitro. To confirm the genotype of the strains, we performed pulse-chase analysis and compared CPY processing. Both PEP4 and pep4Δ strains showed no significant differences for p1 and p2CPY after a 5-min pulse (Fig. 5B, lanes 1 and 3). However, after a 60-min chase the fate of the p1 and p2CPY precursors was different. The PEP4 strain produced mCPY and the pep4Δ strain did not produce any mCPY but the p2CPY precursor accumulated (Fig. 5B, lanes 2 and 4). With these phenotypes established, we prepared radiolabeled P3 donor membranes and unlabeled P2 acceptor membranes from the wild-type PEP4 and the pep4Δ mutant strains. These membranes were then mixed and incubated for cell-free assays in all combinations. Importantly, the radiolabeled reaction product took on the processing phenotype of the unlabeled acceptor membranes, not the radiolabeled donor membranes. For instance, PEP4 acceptor membranes gave rise to mCPY even from pep4Δ donor membranes (Fig. 5C, lanes 5 and 6). A acceptor membranes from the pep4Δ strain did not produce detectable p2CPY maturation (Fig. 5C, lanes 7 and 8). The small amount of mCPY (~9%) that occurred from mixing PEP4 donor membranes with pep4Δ acceptor membranes (Fig. 5C, lane 7) was present in the reaction where no acceptor membranes were added back (Fig. 5C, lane 3). This indicated that a trace amount of vacuoles contaminated the PEP4 donor membranes in this experiment. These reactions with donor and acceptor membranes from a strain deleted for the principal processing protease gene provided the strongest evidence that our new cell-free assay was indeed intercompartmental. This reconstitution was likely an intercompartmental transport process between the PVC and the vacuole.

A Role for Vps33p in the Cell-free Reconstitution Assay

One difficulty in reconstituting an intercompartmental transport event in our previous permeabilized cell assay was incomplete removal of many cytoplasmic VPS gene products such as Vps33p (Vida et al., 1990). For example, no transport defect has been observed when a cytosolic extract devoid of Vps33p from a vps33 null strain (vps33Δ) was added back to wild-type permeabilized cells (data not shown). However, a significant defect was observed in vps33Δ cytosol when it was added back to the cell-free...
transport assay. The transport efficiency was decreased ~2.5-fold compared with cytosol made from a wild-type VPS33 strain (Fig. 6, lanes 2 and 3, 5 and 6). Although the standard concentration of cytosol in our cell-free reactions was 5 mg/ml, these experiments also demonstrated that overall transport efficiency was remarkably consistent with the concentration of protein in crude, undiluted wild-type cytosol. For example, using extracts with a protein concentration of 50 mg/ml produced an average transport efficiency of 47.0% ± 1.3% (n = 10). We observed an average transport efficiency of 32.6% ± 2.5% (n = 10) with an undiluted cytosolic protein concentration of 35 mg/ml. The 30% decrease in transport efficiency correlated well with the 30% decrease in protein concentration, which suggested that the level of a soluble protein factor(s) was critical for driving intercompartmental transport.

Polyclonal antiserum (raised against Vps33p-trpe fusion proteins) also directly implicated Vps33p in playing a specific role during the cell-free assay. We prepared a new antiserum against Vps33p and it proved to be monospecific, recognizing a single polypeptide of ~72 kDa after immunoprecipitation of a total yeast cell lysate (Fig. 7 A, lane 2). The preimmune serum did not immunoprecipitate any proteins in this cell lysate (Fig. 7 A, lane 1). In pilot experiments, the Vps33p immune serum inhibited the cell-free assay while the preimmune had no effect. To avoid potential inhibitory problems from whole serum, we purified total IgG from both the preimmune and immune sera and measured the inhibition in titration experiments with the cell-free assay. As more of the immune IgG against Vps33p was added to the cell-free assay, we observed a proportional decrease in p2CPY transport (Fig. 7 B). At 128 μg and above, the immune IgG was able to block >90% of intercompartmental transport in the assay (Fig. 7 B, lane 8). Importantly, preimmune IgG was without any measurable inhibitory effect (Fig. 7 B, lanes 3-8). Using immune IgG against Vps33p as a specific inhibitor of its function, we determined where Vps33p was most active during the cell-free assay. To test this, we added back IgG against Vps33p at different points during a time course (similar to the kinetics in Fig. 4 B). After allowing antibody/antigen binding for 15 min on ice at each time point, we then continued the incubation for 60 min at 25°C (see Fig. 8 A). The results from this analysis suggested that the role of Vps33p in intercompartmental transport to the vacuole was executed at an early stage in the cell-free assay. For example, when the antibody was added back before incubation (at 0 min), >90% inhibition was observed (Fig. 8 C). Moreover, this inhibition was most effective during the first 10–15 min of the time course (Fig. 8 C). This interval of time in the cell-free assay was the latent period showing very little maturation of p2CPY (Figs. 8 B and 4 B). The inhibition from adding immune IgG against Vps33p during the cell-free assay time course was significantly less at the 15 min time point and beyond (Fig. 8 C). For example, at 15 min only 10% of intercompartmental transport took place (Fig. 8 B) and the inhibition was only 40% (Fig. 8 C). This effect was more notable at the 30 min time point where ~60% (Fig. 8 B) of intercompartmental transport occurred but the inhibition was only 10% (Fig. 8 C).

We also determined the possible involvement of another protein in the cell-free assay, Vam3p. Vam3p is a Q-SNARE protein (Fasshauer et al., 1998) of the vacuole that is directly implicated in homotypic vacuole fusion (Nichols et al., 1997) and delivery of vacuolar proteases through the biosynthetic pathway (Darsow et al., 1997). In contrast to antibodies against Vps33p, antiserum against Vam3p inhibited the assay when added at any time during the entire 45-min incubation (Fig. 8 C). The inhibition by anti-Vam3p serum decreased only ~25% in the first 15 min and remained ~60% throughout the time course (Fig. 8 C).
These results suggested that the function of both Vps33p and Vam3p was required for efficient transport in the cell-free assay. However, an early event(s) was more dependent on the function of Vps33p, particularly during the first 15 min, than later events and Vam3p appeared to be required both early and late in the assay. The ability to inhibit the assay with Vps33p-specific antibodies decreased nearly threefold faster than with antibodies against Vam3p during the first 15 min of the cell-free assay.

**Biochemical Complementation of VPS33 Function**

The cell-free assay has allowed us to directly implicate the function of a VPS gene product in a reconstituted intercompartmental transport event. The inefficient transport from vps33Δ cytosolic extracts and the inhibition by IgG against Vps33p demonstrate that the function of this protein is required during incubation of donor and acceptor membranes. We wanted to positively implicate the role of Vps33p in the cell-free assay, which would establish a transport-coupled biochemical assay for a VPS gene product.

To this end, we expressed Vps33p in bacteria and it was produced at high levels (data not shown). However, over a variety of induction conditions with changes in temperature, time, or inducer concentration, Vps33p repeatedly was insoluble in bacterial lysates (data not shown). To avoid the insolubility problems from overexpression in bacteria, we overexpressed the VPS33 gene in yeast. We placed Vps33p under control of the promoter for glyceraldehyde 3-phosphate dehydrogenase (GPD1) because it is one of the strongest promoters in *S. cerevisiae* (Mumberg et al., 1995). Indeed, a 100-200-fold increase in the amount of GPD1p-Vps33p was observed compared with endogenous levels of the protein and the overexpressed Vps33p behaved like a soluble protein (data not shown). The high level overproduction of soluble Vps33p in yeast allowed us to determine if we could reverse the antibody inhibition of the cell-free reconstitution assay. A fiter incubating the donor/acceptor assay with IgG against Vps33p, a transport efficiency of ~50% maximum was observed with a cytosolic extract from a strain expressing GPD1p-VPS33 (Fig. 9, lane 4). A significant level of transport was
not observed when a cytosolic extract from the vps33Δ strain was added back to the IgG-inhibited reaction (Fig. 9, lane 2). This suggested that restoration of p2CPY transport to the vacuole may be specific to Vps33p. A wild-type cytosol (i.e., VPS33Δ-VPS33) leads to a transport efficiency just under 20% maximum, further suggesting that reversal of inhibition reflected the level of Vps33p added back to the assay. The results of this experiment provide evidence for biochemical complementation of a Vps protein-dependent defect to the yeast vacuole.

**Discussion**

Gaining access to the cell cytoplasm is essential for detailed understanding of intracellular transport between organelles. Genetics and molecular biological approaches are able to obtain entrance into cells with the manipulation of genes and gene products. Although this control can often be very thorough, it is also often limited without augmentation using biochemical approaches in parallel. The biochemistry of transport between organelles requires working in a cell-free system. Frequently, severe limitations to cell-free analyses are not only maintaining organelle structure, but also (and more importantly) organelle function. These are the two most important criteria in successful cell-free reconstitution of intercompartmental transport.

Lysing yeast spheroplasts by extrusion through polycarbonate filters maintains function of organelles in the yeast vacuolar system. Polycarbonate filters are hydrophilic and contain uniform cylindrical pores. The diameter of these pores can be carefully controlled via ion etching, which allows for an even distribution across one plane over the entire exposed membrane surface. The ability to change the overall diameter of a yeast spheroplast with osmotic forces permits swelling of the cells to just greater than the diameter of the polycarbonate filter pores. Thus, in one simple step, the plasma membrane can be gently sheared away from cells and most organelles are free to pass through with little damage. In fact, we have used this method of lysis on mammalian cells (Chinese hamster ovary), which required a simple increase in pore size (from 3 to 8 μm). The yeast vacuole does undergo some loss of luminal content during extrusion through polycarbonate filters as expected from its labile structure. This loss is most likely from leakage rather than lysis and is inconsequential because the amount of soluble proteases is sufficient for processing of propeptides from vacuolarzymogens.

Proteolytic maturation within the donor compartment, presumably the PVC, does not appear to be an efficient process in vitro. A nother explanation for our cell-free assay that measures maturation of p2CPY could be intercompartmental activation of processing proteases. To a

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**Figure 8.** The function of Vps33p is required early during the cell-free reconstitution system and precedes the function of Vam3p. (A) Experimental strategy. (B) Transport kinetics. A standard 50-μl reaction (see Fig. 3) was scaled up, incubated at 25°C, and then aliquots were removed and stopped at the indicated times. (C) A 2nd degree polynomial was used to fit the curve for the inhibition data points (C).

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**Figure 9.** Biochemical reconstitution of Vps33p-dependent transport between donor and acceptor membranes. Radiolabeled donor membranes and nonradiolabeled acceptor membranes were prepared from wild-type yeast spheroplasts (Fig 1A). The donor and acceptor membranes were incubated for 15 min at 0°C with 128 μg of anti Vps33p IgG. Cytosol (5 mg/ml) was then added from a vps33Δ (lane 2), a VPS33 (lane 3), or a VPS33 strain containing pGPDHIS-633-2 (lane 4), and incubated at 25°C for 60 min (as indicated). All reactions were immunoprecipitated for CPY, subjected to SDS-PAGE, and autoradiography. The bar graph depicts average transport efficiency from three independent determinations and normalized to the percent of maximal transport.
first approximation, proPrA is contained in the same compartments as p2CPY. Unlike proCPY, the proPrA precursor has the ability to autoactivate (McClellan et al., 1988, 1987). Therefore, incubation of donor membranes with ATP and cytosol might lead to changes in luminal pH that would enhance autoactivation of proteinase A. Once proPrA becomes an active hydrolase it would begin a cascade of proteolytic events leading to the activation of Prb1p and ultimately maturation of p2CPY. However, under our cell-free assay conditions, maturation of p2CPY was undetectable after incubating donor membranes alone with ATP and cytosol. The inhibition of the cell-free assay after diluting the membranes further supports the conclusion that proteolytic processing of p2CPY does not occur within the donor membrane compartment. These points argue that the organelle containing p2CPY does not acquire the capacity to cleave propeptides, which indicates that organelle maturation may not be a prevalent mechanism to produce vacuoles/lysosomes.

This cell-free system is easily manipulated to show a near absolute requirement for exogenous cytosol. With this cytosol requirement, we can tentatively assign the location of Vps33p function to cycling from the cytosol to either the donor or acceptor membrane fractions. Although Vps33p is predominately localized to the cytosol, a fraction of the protein sediments with membranes (Gerhardt et al., 1998). Since Vps33p binds ATP and readily interchanges between soluble and insoluble forms in an energy-dependent fashion (Gerhardt et al., 1998), the cell-free system is the best way to understand how ATP influences its function in transport to the vacuole.

One prospect for functional interaction of Vps33p with an insoluble component(s) is a SNAARE complex (Bennett, 1995). The VPS33 gene product is a member of the Sec1p family of proteins (Pevsner, 1996) and thus is expected to bind to a target SNAARE protein on the vacuole such as Vam3p (Darsow et al., 1997; Wada et al., 1997). Our results suggest that Vps33p acts earlier than Vam3p in the cell-free assay. This is distinct to the inhibition via antibodies against Vam3p, which still show a significant block late into the reaction. One possibility to explain these results is that the epitopes on Vps33p are exposed for antibody binding early during the assay but become inaccessible due to conformational changes later in the time course. On the other hand, Vps33p may have a catalytic or binding activity that is required early during inter compartmental transport but not at late stages. Although the precise cause for losing Vps33p function during inter compartmental transport to the vacuole is unknown, the results suggest that it may be independent of Vam3p function. This has implications on how we view the roles Sec1- and syntaxin-like proteins play in vesicle-mediated transport.

The Sec1p family has many members, suggesting that these proteins function at every vesicle-mediated step in eukaryotic cells (Pevsner, 1996; Pevsner et al., 1994). The Drosophila Sec1 homologue, ROP, and yeast Syl1p can negatively regulate neurotransmitter release (Schulze et al., 1994) and prevent v-SNARE/t-SNARE interactions in ER to Golgi transport, respectively (Lupashin and Waters, 1997). Mammalian Sec1-like proteins bind to syntaxin with high affinity (Hodel et al., 1994; Pevsner et al., 1994). Although this protein–protein interaction is most likely to be physiologically relevant, its biological meaning is far from clear. For example, the direct physical interaction between Sec1p-like proteins and t-SNARE proteins has only been demonstrated in vitro (Hodel et al., 1994; Pevsner et al., 1994). Furthermore, attempts to coimmunoprecipitate a Sec1p homologue-syntaxin homologue complex from cell extracts have never been successful (Garcia et al., 1995; Wu et al., 1998), suggesting their interaction in vivo is transient, weak, or both. Nonetheless, simultaneous overexpression of syntaxin and ROP in Drosophila suppresses the defects in neurotransmission that are observed when either is overexpressed individually (Wu et al., 1998). Moreover, recent genetic evidence in yeast suggests that Vps33p interacts with a Q-SNARE protein of the vacuole membrane, Vam3p (Darsow et al., 1997). A haploid double mutant strain with temperature-sensitive alleles in both vps33 and vam3 shows a ~50% defect in CPY maturation under conditions where the single haploid mutations are wild-type (Darsow et al., 1997). These are compelling examples that implicate a ROP/syntaxin and Vps33p/Vam3p functional interaction in vivo. However, studies involving overexpression or synthetic defects of two different genes are not of sufficient resolution to distinguish whether their products physically bind to one another or if they are part of a linear pathway. A direct physical interaction between ROP and syntaxin or Vps33p and Vam3p has not been demonstrated and instead has been inferred from analogy to studies with n-Sec1 and syntaxin (Pevsner et al., 1994). This inference may be appropriate for ROP because it is significantly more similar to n-Sec1 than Vps33p. Indeed, Vps33p shows characteristics not shared among the other Sec1 members such as ATP binding and energy insolubility (Gerhardt et al., 1998). These differences suggest that Vps33p may play a distinct role in vesicle-mediated transport.

The advent of our cell-free assay will help uncover biochemical activities of VPS gene products. The majority of these proteins do not show sequence similarity to proteins of known biochemical properties, although several VPS gene products have activities in vitro. Without exception, the ability to design assays for detection of these catalytic activities arose from sequence similarities to proteins that had been subjected to previous biochemical characterization. This underscores the importance that biochemistry plays in elucidating gene function and discovering new activities should progress rapidly with our inter compartmental assay. This assay will allow us to define biochemical functions of cytosolic and membrane-associated factors necessary to execute transport between the PVC and vacuole. Our results with the cell-free assay imply that a vesicle intermediate may truly shuttle cargo between the PVC and the vacuole in yeast. Preincubation of donor membranes (in the absence of acceptor membranes) with ATP and cytosol gives rise to a fraction that contains p2CPY and separates from donor membranes. This membrane-enclosed compartment can be used as a functional intermediate in a second incubation with acceptor membranes (Gerhardt and Vida, manuscript in preparation). Many of the factors involved in this process are likely VPS gene products and may play a role in vesicle formation, transport, targeting, and fusion with the yeast vacuole/lysosome.
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