Coupled ER to Golgi Transport Reconstituted with Purified Cytosolic Proteins

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Abstract. A cell-free vesicle fusion assay that reproduces a subreaction in transport of pro-α-factor from the ER to the Golgi complex has been used to fractionate yeast cytosol. Purified Sec18p, Uso1p, and LMA1 in the presence of ATP and GTP satisfies the requirement for cytosol in fusion of ER-derived vesicles with Golgi membranes. Although these purified factors are sufficient for vesicle docking and fusion, overall ER to Golgi transport in yeast semi-intact cells depends on COPII proteins (components of a membrane coat that drive vesicle budding from the ER). Thus, membrane fusion is coupled to vesicle formation in ER to Golgi transport even in the presence of saturating levels of purified fusion factors. Manipulation of the semi-intact cell assay is used to distinguish freely diffusible ER-derived vesicles containing pro-α-factor from docked vesicles and from fused vesicles. Uso1p mediates vesicle docking and produces a dilution resistant intermediate. Sec18p and LMA1 are not required for the docking phase, but are required for efficient fusion of ER-derived vesicles with the Golgi complex. Surprisingly, elevated levels of Sec23p complex (a subunit of the COPII coat) prevent vesicle fusion in a reversible manner, but do not interfere with vesicle docking. Ordering experiments using the dilution resistant intermediate and reversible Sec23p complex inhibition indicate Sec18p action is required before LMA1 function.

In eukaryotic cells, transport between many distinct membrane-bound compartments proceeds through a vesicular intermediate that buds from one membrane and fuses selectively with another. A framework for this transport process has emerged whereby soluble coat proteins are recruited to the donor membrane forming vesicles that uncoat before fusion with an acceptor compartment (Rothman, 1994). With respect to the membrane fusion step, a set of general factors including N-ethylmaleimide–sensitive fusion protein (NSF) and soluble NSF attachment protein (SNAP) are required at many intracellular compartments (Block et al., 1988; Eakle et al., 1988; Clary and Rothman, 1990; Clary et al., 1990). These general factors are used in concert with compartment specific factors such as v/t-SNAP receptors (SNAREs) and small GTPases that appear to be structurally related, yet impart specificity in membrane docking and fusion. The basic mechanisms of intracellular membrane fusion appear to be conserved within a species and from yeast to mammals (for review see Pfeffer, 1996). Although several of the molecules that catalyze specific intracellular membrane fusion reactions have been identified through biochemical and genetic approaches, a molecular description of this process is incomplete.

Using the yeast Saccharomyces cerevisiae as a model eukaryote for the investigation of intracellular transport, a cell-free assay that faithfully reproduces ER to Golgi transport has been developed based on the well-characterized processing and maturation of pro-α-factor, the precursor of a secreted pheromone in yeast (Baker et al., 1988). The application of an in vitro transport assay, together with genetic analyses of sec mutant strains has allowed the division of ER to Golgi transport into three distinct steps: vesicle budding, vesicle targeting, and membrane fusion (Kaiser and Schekman, 1991; Rexach and Schekman, 1991). The first step in this sequence, vesicle budding, has been reconstituted in vitro with a set of soluble factors (Sec1p, Sec23p complex, and Sec13p complex) that collectively form a membrane coat that drives budding from the ER. The vesicles formed with these purified protein fractions, termed COPII-coated vesicles, are competent for fusion with the Golgi apparatus in a reaction that requires cytosol and ATP (Barlowe et al., 1994). This cytosol provides multiple protein components required for fusion, some of which are...
defined by known secretion defective mutants (Lupashin et al., 1996).

To characterize the molecular events associated with intracellular membrane fusion, the long-term goal is to reconstitute this event with a minimal set of membrane bound and soluble proteins. In this report, the soluble factors required for cell-free docking and fusion of ER-derived vesicles to the Golgi complex have been defined. This vesicle fusion reaction in the presence of ATP and GTP requires purified Sec18p, Us10p, and a protein complex referred to as LMA1, a heterodimer composed of thioredoxin and IB2 (Xu and Wickner, 1996; Xu et al., 1997). Although these purified factors are sufficient for vesicle targeting and fusion to the Golgi complex, overall ER to Golgi transport in yeast semi-intact cells is dependent on the addition of COPII proteins. Thus, six purified, soluble proteins are shown to catalyze anterograde ER to Golgi transport and are ordered in distinct steps of vesicle budding, docking, and fusion.

Materials and Methods

Strains and Reagents

Strains RSY607 (MATa leu3-3,112 ura3-52 pep4::URA3), RSY919 (MATa ura3-1 mnn1 mnn2), RSY949 (MATa leu2-201 trpl ura3-52 usol-1), CBY300 (MATa his3Delta0 leu2Delta1, lys2a202 ura3-52 trplDelta6 sol-1), and CBY324 (MATa his3Delta0 ura2Delta1, lys2a202 ura3-52 trplDelta3 usol-1 pUSO1-Myc) were used in these studies. Rabbit antisera specific for α-1,6-mannose linkages was prepared by intravenous injection of heat-treated RSY919 cells as previously described (Ballou, 1970). Antibodies directed against Sec7p (Franzusoff et al., 1991), Sec13p (Salama et al., 1993), Sec18p (Haas and Wickner, 1996), Sec19p (Haas et al., 1995), Sec23p (Hicke and Schekman, 1989), Sec26p (Duden et al., 1995), Trx1p (kindly provided by Z. Xu, Dartmouth Medical School, Hanover, NH), Ypt1p (Rexach et al., 1994) and monoclonal 9E10 anti-myc (Evan et al., 1985) were used in these studies. In vitro-translated pre-pro-α-factor was synthesized (Baker et al., 1988) using translation grade [35]S-methionine (Amersham Corp., Arlington Heights, IL). Methods for SDS-PAGE (Laemmli, 1970), silver staining (Bloom et al., 1987), immunoblotting (Towbin et al., 1979), and immunodetection by enhanced chemiluminescence (Amer- sham Corp.) have been described. Deuterium oxide, Nycodenz, ATP, GTP, creatine kinase, and phosphocreatine were purchased from Sigma Chemical Co. (St. Louis, MO).

Plasmid Construction

The c-myc epitope was fused to the COOH terminus of Usol1p by modification of a 2 µm -URA3 plasmid containing the USO1 gene pSK47 (Sapperstein et al., 1995). Oligonucleotides U1 (CGGAATCT CATG CACCATC AAGGACTGC) and U2 (CGGAATTCT CAAGCTTCT TT-CAGAATA AGTTTGTIG CTCTGCA CTGTTTCTTC TC-CTCG) were used to amplify a portion of the USO1 gene, which was then digested with the restriction enzymes Xbal and SacI and inserted into pBluescript (Stratagene, La Jolla, CA) digested with Xbal and SacI. After confirmation of this sequence, the 1.4-kb Xbal-SacI fragment was inserted into pSK47 digested with SacI and partially digested with Xbal. The resulting construct (pUSO1-myc) fully complements an usol-1 strain (CBY300) for growth at 37°C.

Cytosol and Membrane Preparations

Strain RSY607 was grown to mid-log phase (OD600 = 1) (determined with a spectrophotometer; Beckman Instruments, Inc., Fullerton, CA) in 15 liters of yeast extract, peptone, dextrose (YPD) medium. The cells were harvested and washed once with cold H2O, and once with buffer 88 (B88) (20 mM Hepes, pH 7.0, 0.25 M sorbitol, 0.15 M KOAc, 5 mM MgOAc). Approximately 80 g (wet wt) of cells were recovered from this procedure and are resuspended with 15 ml of B88, and then quick frozen by dropwise addition to liquid nitrogen. The frozen cells were mixed with liquid nitrogen in a Waring blender for 10 min to prepare a cell lysate (Dunn and Wobbe, 1989). This lysate was thawed on ice and diluted with 10 ml of B88 and adjusted to final concentrations of 1 mM DTT and 1 mM PMSF. A medium speed supernatant (MSS) fraction was prepared by centrifugation at 25,000 g for 15 min (SS34 rotor; Sorvall, Newtown, CT), and then decanted from centrifuge tubes carefully excluding loosely sedimented membranes. Aliquots of this MSS fraction were quick frozen in liquid nitrogen and stored at ~75°C.

Cytosol used for transport reactions or for fractionation on Mono Q was prepared from thawed aliquots of MSS by centrifugation at 89,000 g for 15 min (50,000 rpm in a TL120.2 rotor; Beckman Instruments, Inc.). The resulting supernatant fluid was recovered and mixed with two volumes of B88 containing 1 mM DTT and 1 mM PMSF (B88D/P), and then centrifuged at 175,000 g for 15 min (70,000 rpm in a TL120.2 rotor). The clarified portion of this supernatant fraction was removed with a pipette and frozen in liquid nitrogen for storage at ~75°C. This cytosol fraction serves as the starting material for fractionation procedures and was ~5 mg/ml protein, using BSA as a standard (Bradford, 1976).

Acceptor membranes were prepared from the same MSS as above, except after the first centrifugation step (at 89,000 g) the resulting membrane pellet was processed. First, loosely sedimented membranes were aspirated from this pellet, leaving a compact translucent pellet. This pellet was resuspended by douche homogenization in B88D/P to a volume equal the initial MSS volume. The membranes were collected again by centrifugation at 89,000 g and the B88D/P wash was repeated. This final pellet was resuspended by douche homogenization in B88D/P with one-fifth of the initial MSS volume, and aliquots were frozen in liquid nitrogen and stored at ~75°C. This preparation of acceptor membranes was ~3 mg membrane protein/ml.

Fractionation of Cytosol by Mono Q Chromatography

Starting with 7 ml of MSS, 15 ml of cytosol was prepared as described above and loaded at 0.5 ml/min onto a Mono Q HR 10/10 column (Pharmacia Biotechnology Inc., Piscataway, NJ) that was equilibrated in buffer A (20 mM Hepes, pH 7.0, 0.15 M KOAc, 1 mM MgOAc, 0.1 mM DTT, 0.1 mM PMSF, and 0.01 mM ATP). Fractions (2 ml) were collected from the Mono Q flowthrough. The column was washed with 15 ml of buffer A, followed by step elutions with 15 ml of buffer A containing 0.75 M KOAc, and 15 ml of buffer A containing 1.5 M KOAc. The protein peaks were contained in fraction 6 (flow through [QFT] ~4 mg/ml), fraction 20 (intermediate ionic strength [Q75] ~10 mg/ml), and fraction 27 (high ionic strength [Q1.5] ~3 mg/ml). These fractions were dialyzed against B88D/P, frozen in liquid nitrogen, and then stored at ~75°C.

Protein Purification

Proteins contained in the Mono Q fractions were resolved on a Superox 6 HR 10/30 column (Pharmacia Biotechnology Inc.), equilibrated, and then eluted with 20 mM Hepes, pH 7.0, 150 mM KOAc, 1 mM MgOAc. Samples (0.2 ml) were loaded at a flow rate of 0.3 ml/min, and 0.5-ml fractions were collected after the initial 5 ml had been discarded. The void volume for this column corresponds to 7.5 ml. The c-myc-tagged version of Usol1p was purified from strain CBY324 as follows. 1 liter of cells was grown under selective conditions (yeast nitrogen base, 2% dextrose, and supplement minus uracil), and then transferred to 12 liters of YPD for growth to an OD600 = 1.0 and processed as for the preparation of cytosol from RSY607. Proteins contained in this cytosol were bound to the Mono Q HR 10/10 column as above except a 20-ml linear gradient was delivered from 0.75 to 1.5 M KOAc instead of a step elution. The peak of activity (Fraction 27, high ionic strength [Q1.5] ~3 mg/ml). These fractions were dialyzed against B88D/P, frozen in liquid nitrogen, and then stored at ~75°C.

Preparation of ER-derived Vesicles Containing 35S-labeled, Core-glycosylated Pro-α-factor

Vesicles were synthesized from a microsomal preparation (Wuestehube and Schekman, 1992) with purified Sar1p, Sec23p complex, and Sec13p.
complex (Barlowe et al., 1994). First, posttranslational translocation of pre-pro-a-factor into microsomal membranes was performed in 0.4 ml of B88 containing 0.7 mg of membrane protein, 35S-labeled gp-a-factor (5 \times 10^6 cpm), and an ATP regeneration system (Baker et al., 1988) at 10°C for 15 min. After chilling on ice, membranes were diluted in B88, centrifuged at 12,000 g for 3 min, and then washed twice with B88 by gentle resuspension in buffer before centrifugation. The final pellet was resuspended in 0.1 ml of B88 and mixed with 5 mg each of purified Sar1p, Sec13p complex, and Sec23p complex in 0.6 ml total vol of B88 containing the ATP regeneration system and 0.1 mM GTP. Budding reactions were incubated at 20°C for 10 min, and then placed on ice for 5 min. Vesicles were separated from microsomal membranes by centrifugation at 14,000 g for 5 min, and 0.55 ml of the supernatant fluid–containing vesicles was removed and mixed with 0.8 ml of a 60% Nycodenz (wt/vol) in D2O containing 20 mM K-Hepes, 150 mM KOAc, and 5 mM MgOAc. This mixture was layered on the bottom of a SW60 tube (344062; Beckman Instruments Co.) followed by 0.9-ml layers of 25% Nycodenz, and 20% Nycodenz in the same D2O-containing buffer. A final layer of B88 (1 ml) was placed on top, and the tube was centrifuged at 50,000 rpm in a SW60 rotor for 3 h. The top 0.8 ml was discarded, and then 0.15-ml fractions were taken and the vesicle peak was determined by scintillation counting. Typically, the vesicle peak was recovered in 0.3 ml and contained 35S-labeled, core-glycosylated pro-a-factors (~2,000 cpm/μl) that was protease protected, and precipitable with concanavalin A–Sepharose beads (Pharmacia Biotechnology Inc.).

**Vesicle Fusion Assay**

The fusion of purified, ER-derived vesicles with the Golgi compartment was reproduced in vitro by incubation of acceptor membranes (12 μg), vesicle-containing 400 cpm of 35S-labeled pre-pro-factor, and protein fractions (as indicated in figures) in a 30-μl reaction volume. Assays were performed at 23°C in B88, which contained an ATP regeneration system and GTP (0.1 mM). After specified times (standard reactions were 60 min), 50 μl of SDS (2%) was added and tubes were heated to 95°C for 4 min. 1 ml of IP buffer (25 mM Tris-Cl, 150 mM NaCl, 1% Triton X-100) was added, followed by 15 μl of anti-a-1,6-mannose-specific serum and 35 μl of a 20% vol/vol solution of protein A–Sepharose (Pharmacia Biotechnology Inc.). Immunoprecipitation reactions were gently rotated at room temperature for 2 h, and immune complexes were processed as previously described (Baker et al., 1988). Vesicle fusion measured in these experiments reflects the amount of 35S-labeled gp-a-factor that acquired the Golgi-specific α-1,6-mannose outer-chain modification (εmmunoprecipitated) as a percentage of the total 35S-labeled gp-a-factor, as determined by precipitation with concanavalin A linked to Sepharose beads. Data points represent the average of duplicate determinations, where each duplicate set varied by <10%.

**ER to Golgi Transport Assays**

Yeast semi-intact cells were prepared from log phase cultures of strain RSY607 (Baker et al., 1988). Spheroplasts were perforated using low osmotic support buffer and ER to Golgi transport reactions performed in two stages as described (Rexach and Schekman, 1991). After stage I (translocation of 35S-labeled pre-pro-a-factor into ER membranes), cells were placed on ice, centrifuged at 15,000 g for 2 min in a refrigerated centrifuge (5417; Eppendorf Inc., Madison, WI). Cells were then gently resuspended and washed three times in B88. Transport reactions (30 μl) containing semi-intact cells, GTP, ATP regeneration system, and various protein fractions were incubated at 23°C for 45 min, and then chilled on ice for 5 min. A 5-μl aliquot was removed and treated with trypsin followed by solubilization in 1% SDS and precipitation with conconavalin A–Sepharose to determine total protease protected 35S-labeled gp-a-factor (Baker et al., 1988). Transport to the Golgi is expressed as the percentage of this conconavalin A–precipitable, 35S-labeled gp-a-factor that has acquired Golgi-specific α-1,6-mannose outer chain modification determined by precipitation with anti-a-1,6-mannose serum as described above for the vesicle fusion assay. Parallel (but separate) tubes were processed to quantify the amount of freely diffusible, ER-derived vesicles present under each condition. After 5 min on ice, the samples were spun at 18,000 g for 3 min in a refrigerated centrifuge (5417; Eppendorf Inc.). Aliquots (15 μl) were taken from the resulting supernatant fluid (medium-speed supernatant) and the amount of 35S-labeled gp-a-factor that was determined after protease treatment and precipitation with conconavalin A–Sepharose (Rexach and Schekman, 1991). Budded vesicles are expressed as the percentage of total 35S-labeled gp-a-factor that is contained in the medium-speed supernatant fraction. Data points represent the average of duplicate determinations, where each duplicate set varied by <10%.

**Results**

**Fractionation of Cytosolic Components Required for Vesicle Fusion**

The formation of transport vesicles from the ER has been reproduced in a cell-free assay using purified soluble factors (Sar1p, Sec13p complex, and Sec23p complex) and washed membranes. The isolated vesicle intermediates produced in this reaction are distinct from the donor membrane fraction and are competent for fusion with the Golgi complex (Barlowe et al., 1994). This serves as a starting point for the isolation of soluble factors contained in a yeast cytosol preparation that catalyze vesicle fusion in an assay that is independent of factors required for vesicle budding. A facile and reproducible assay to measure this process is established in this report whereby the components of this assay (purified ER-derived vesicles, yeast cytosol, and Golgi membranes) could be prepared in large quantities and stored at −75°C for extended periods of time with minimal loss of activity.

Cytosol was bound to an anion exchange resin and three fractions were obtained: the flow through (QFT), a spectrum of proteins that elute at an intermediate ionic strength (Q.75), and proteins that elute at a high ionic strength (Q1.5). The individual fractions were dialyzed and tested in a cell-free assay for promotion of vesicle fusion (Fig. 1). Addition of individual fractions revealed that none of the fractions alone could drive vesicle fusion as efficiently as a crude cytosol, though a significant signal could be detected by adding saturating amounts of the Q1.5 fraction alone (Fig. 1, columns 1–4). A maximal signal (comparable to cytosol) could be obtained by combining all three of the fractions, and in fact, omission of either fraction resulted in a fusion efficiency below crude cytosol (Fig. 1, columns 5–8).

Since a number of the yeast proteins involved in intracellular transport have been identified, specific antibodies that recognize these species were used in an attempt to...
guide further purification efforts. An immunoblot characterizing some of the species contained in these fractions is shown in Fig. 2. A requirement for Sec7p in vesicle transport has been reported (Lupashin et al., 1996), and this protein was detected in the Q1.5 eluate. Sec23p, a subunit of the COPII complex required for vesicle budding (Hicke and Schekman, 1989) was also contained in the Q1.5 fraction. Sec26p, the β-COP subunit of yeast COPI complex (Duden et al., 1994) elutes at ~0.73 M KOAc (Hosobuchi et al., 1992) and was found in both the Q.75 and the Q1.5 fractions. Sec18p, the yeast homologue of NSF, was found exclusively in the Q.75 fraction; however, yeast thioredoxin, a protein involved in homotypic vacuolar membrane fusion (Xu and Wickner, 1996) was not bound to this anion exchange resin under these conditions and was contained in the flow through fraction. Together, these results indicate the column was not overloaded with cytosolic protein, and that several protein species were effectively resolved. In addition to the proteins shown in Fig. 2, Sec19p, which is the yeast GDP dissociation inhibitor (GDI) (Garrett et al., 1994), and the small GTPase, Ypt1p, were detected in both the QFT and the Q.75 fractions and monitored throughout the following purification procedures.

**Sec18p Replaces the Q.75 Fraction**

Sec18p/NSF is required for several intracellular transport steps and is considered a “general fusion factor” that uses ATP hydrolysis in fulfilling a role in membrane fusion (Rothman, 1994). Mutant sec18 yeast strains are defective for ER to Golgi transport in vivo and in vitro although this block has not been readily restored in various cell-free assays (Rexach and Schekman, 1991; Lupashin et al., 1996). Recombinant forms of NSF and Sec18p that contain six His residues at their COOH termini have been constructed, which may be overproduced in *Escherichia coli* (Whiteheart et al., 1994), and appear to be fully functional in cell-free assays (Whiteheart et al., 1994; Haas et al., 1996). Since Sec18p plays a central role in intracellular membrane fusion reactions, the six His-tagged protein was isolated in a buffer compatible with this assay and tested for promotion of vesicle fusion to the Golgi complex. Sec18p could completely substitute for the Q.75 fraction (Fig. 3 A) at a concentration similar to that contained in a saturating amount of this fraction (see Fig. 6 B). Furthermore, the purified Sec18p alone was not sufficient for fusion of ER-
derived vesicles but required the addition of both the QFT and the Q1.5 for a maximal signal in the vesicle fusion assay (Fig. 3 A).

**Uso1p Replaces the Q1.5 Fraction**

Resolution of proteins contained in the Q1.5 fraction by gel filtration chromatography on a Superose 6 column indicated a single peak of activity that migrated at ~800 kD could support vesicle fusion when combined with purified Sec18p and the QFT fraction (data not shown). Two previously characterized proteins could behave in this manner and have been implicated in ER to Golgi transport: Sec7p (Franzusoff and Schekman, 1989; Franzusoff et al., 1991; Franzusoff et al., 1992; Lupashin et al., 1996) and Uso1p (Nakajima et al., 1991; Seog et al., 1994; Lupashin et al., 1996; Sapperstein et al., 1996). A series of experiments were performed to determine if either or both of these proteins are functional components of the Q1.5 fraction. An antibody that recognizes Sec7p (Franzusoff et al., 1992) was used to analyze the fractions eluting from the Superose 6 column, and a broad peak of Sec7p immunoreactivity was detected that was offset from the activity peak (not shown). However, there was clearly Sec7p immunoreactivity contained in the peak activity fraction raising the possibility that Sec7p plus additional factors are provided from the Superose 6 activity peak. To exclude this possibility, a cytosol was prepared from a mutant yeast strain that is rendered temperature sensitive due to an amber mutation in the *USO1* gene (Seog et al., 1994). Cytosol prepared from the mutant strain was directly compared to a wild-type cytosol by fractionation on a Mono Q column (see Materials and Methods). In this experiment (Fig. 4), a linear gradient of increasing ionic strength was used to elute proteins from 0.75 to 1.5 M KOAc instead of the step gradient used to prepare the Q1.5 fraction. Under these conditions, two observations strongly suggest that Uso1p alone is the active component present in the Q1.5 fraction. First, assay of individual fractions across the elution profile of a wild-type cytosol reveal a single peak of activity.
when combined with purified Sec18p and QFT fractions while cytosol prepared from an uso1-1 strain did not contain detectable activity eluting at this ionic strength (Fig. 4). Second, immunoblot analysis of wild-type fractions indicated Sec7p immunoreactivity was resolved from the peak of vesicle fusion activity on this gradient and eluted before the activity peak (not shown).

To confirm that Uso1p was the active component and to guide purification efforts, a version of Uso1p was constructed whereby the COOH terminus of the protein contains an additional 11 amino acid residues comprising a c-myc epitope recognized by mAb 9E10 (Evan et al., 1985). Expression of Uso1p-myc from a multicopy vector in a uso1-1 temperature sensitive strain fully complements when grown at 37°C indicating the c-myc extension does not interfere with Uso1p function. This strain was used to overproduce and purify Uso1p-myc by a two-step procedure using anion exchange and gel filtration chromatography as described under Materials and Methods. The final purification step (elution from the Superose 6 column) is shown in Fig. 5. The peak of fusion activity that elutes from the Superose 6 column at 800 kD coincided with a single 210-kD polypeptide species observed on SDS–polyacrylamide gel and c-myc immunoreactivity. These fractionation properties are consistent with previous studies suggesting Uso1p forms a nonglobular oligomer due to an extended coiled-coil rod structure similar to a related protein in mammalian cells, termed p115 (Waters et al., 1992; Seog et al., 1994; Sapperstein et al., 1995). The peak fractions eluting from the Superose 6 column were pooled (16–17) for use in later experiments, and neither Sec7p nor Sec26p (β-COP) could be detected in this pooled fraction by immunoblot analysis.

LMA1 Replaces the QFT

Addition of saturating amounts of Sec18p and Uso1p to vesicle fusion reactions resulted in a fusion efficiency below that catalyzed by crude cytosol, yet addition of the QFT fraction stimulated to near cytosolic levels (data not shown). Fractionation of QFT by Superose 6 chromatography revealed a broad peak of activity in the ~20–40 kD range however, maximal activity could not be recovered perhaps because of extensive dilution through these combined purification steps. Immunoblot analysis of these fractions suggested the presence of Ypt1p, Sec19p (GDI), and thioredoxin: a subunit of a heteroligomeric complex termed LMA1 (low molecular weight activity 1) required for vacuole membrane fusion (Xu et al., 1997). Addition of fractions enriched in Ypt1p or Sec19p did not stimulate nor replace the QFT requirement when combined with purified Uso1p and Sec18p (not shown). However, the addition of purified LMA1 stimulated the cell-free vesicle fusion reaction and when saturating levels of protein were added, fusion efficiencies at or above cytosolic levels were obtained (Fig. 6 A). The amounts of Sec18p and LMA1 required for saturation are comparable to levels contained in a crude cytosol (Fig. 6 B). Saturating levels of LMA1 alone did not support vesicle fusion in the absence of Uso1p and Sec18p (Fig. 7 A).

These purified protein fractions were titrated to obtain an optimal concentration for fusion efficiency and each component was then tested alone and in various combina-
tions for support of vesicle fusion (Fig. 7A). Neither of the fractions alone provided significant levels of vesicle fusion activity, whereas addition of Sec18p and Uso1p resulted in half the maximal level of vesicle transport but could be stimulated by addition of LMA1. In Fig. 7B, vesicle fusion catalyzed by crude cytosol or the purified factors was monitored over time at relevant temperatures. The reconstituted reaction showed time and temperature dependence that was very similar to a reaction driven by crude cytosol. Therefore in this fusion assay, purified Sec18p, Uso1p, and LMA1 together represent a minimal set of proteins required to catalyze the fusion of ER-derived vesicles with the Golgi in a manner indistinguishable from crude cytosol. Requirements for these individual factors now allow for further characterization in the processes of vesicle docking and membrane fusion.

**Requirements for COPII and Fusion Factors in ER to Golgi Transport**

The initial cell-free assay for ER to Golgi transport (Baker et al., 1988) required semi-intact yeast cells and cytosol. If the vesicle fusion assay used in this current study represents an authentic subreaction of the overall transport process, these isolated fusion factors should be required for overall transport. Furthermore, if vesicle budding is a prerequisite for membrane fusion, the isolated fusion factors should be most effective in the presence of the COPII proteins (Sar1p, Sec23p complex, and Sec13p complex) that drive vesicle formation (i.e., fusion is coupled to budding). Washed semi-intact cells were first incubated with levels of COPII proteins that promote maximal budding efficiency in the presence of purified fusion factors. Overall transport could not be detected above background under these initial conditions. However, a titration revealed that high concentrations of COPII proteins inhibit transport whereas lower concentrations of the coat constituents stimulate transport, even though suboptimal for vesicle budding (Fig. 8A). Transport of 35S-labeled gp-α-factor to the Golgi was maximally 21% efficient in the reconstituted reaction compared to a background of 3%. As observed in Figs. 8 and 9, both COPII (2 ng/μl) and purified fusion factors were necessary for overall ER to Golgi transport in semi-intact cells indicating vesicle budding is a prerequisite for membrane fusion. To determine if the production of vesicle intermediates simply satisfies a spatial separation of the ER and Golgi present in semi-intact cells, a similar experiment was performed using diffusible ER (microsomes) and acceptor membranes incubated with fusion factors in the presence or absence of COPII (2 ng/μl) as in Fig. 8A. Under this condition, addition of both fusion factors and COPII were again required for a maximal stimulation of transport (not shown), similar to that observed in semi-intact cells. These results suggest that COPII budding activates factor(s) involved in membrane fusion and that direct fusion of ER with Golgi membranes is not efficient.

Elevated concentrations of COPII inhibit ER to Golgi transport (Fig. 8A). One interpretation of this result is that under high concentrations of COPII, ER-derived vesicles retain bound COPII proteins thus hindering access to the Golgi docking and/or fusion machinery. Indeed, under conditions where the COPII coat remains locked on ER-derived vesicles because of inhibition of the Sar1p GTPase, vesicles fail to fuse with the Golgi complex (Barlowe et al., 1994). Interestingly, only one of the COPII components, the Sec23p complex, was required for potent inhibition of vesicle fusion (Fig. 8B). The inhibition caused by elevated concentrations of Sec23p complex could be at vesicle docking or bilayer fusion. A method to monitor these separate events was devised in the following experiment.

**Uso1p Catalyzes Vesicle Docking**

Semi-intact yeast cells incubated with COPII proteins produce freely diffusible, ER-derived vesicles that remain in the supernatant fraction after centrifugation at 18,000 g for 3 min (Fig. 8). Upon addition of purified fusion factors, it was observed that a significant fraction of these vesicles (>50%) cosedimented with semi-intact cells and were not
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Figure 8. Excess COPII proteins inhibit ER to Golgi transport. (A) Budding and transport assays with semi-intact yeast cells in the presence of purified factors. Budding of 35S-labeled gp-α-factor into freely diffusible vesicles (■) was determined in the presence of increasing amounts of COPII proteins alone (no fusion factors added). COPII concentrations reflect individual protein concentrations such that the 2 ng/μl COPII condition is 2 ng/μl Sar1p, 2 ng/μl Sec23p complex, and 2 ng/μl Sec13p complex. Transport (○) was quantified in the presence of fusion factors (Sec18p [50 ng], Uso1p [75 ng], and LMA1 [50 ng]) and varying concentrations of COPII proteins in 30-μl reactions. (B) Transport in the presence of fusion factors (as in A) and COPII proteins (2 ng/μl), plus 8 ng/μl of Sar1p (5), or 8 ng/μl Sec13p complex (13), or 8 ng/μl Sec23p complex (23).

Although Uso1p reduces the level of freely diffusible vesicles produced by COPII, vesicle fusion as measured by Golgi specific glycosylation under this condition was modest (Fig. 9, hatched bar, column 12) and does not account completely for the decrease in diffusible vesicles. These observations suggest that vesicles dock, but require additional factors to promote bilayer fusion. Indeed, the addition of LMA1 resulted in a significant increase in vesicle fusion (Fig. 9, hatched bar, column 14) that was largely dependent on Uso1p (Fig. 9, hatched bar, compare to column 10). Addition of Sec18p has a minor effect on vesicle fusion in this assay that may reflect ample levels of endogenous Sec18p associated with semi-intact cell membranes.

The effects of inhibitory amounts of Sec23p complex on vesicle docking and fusion were also determined. As documented in Fig. 8 B, Sec23p complex is a potent inhibitor of ER to Golgi transport at high concentrations. Upon addition of this inhibitor, transport was blocked (Fig. 9, hatched bar, column 17) but the reduction in freely diffusible vesicles was quite similar to conditions of optimal targeting (Fig. 9, compare to column 16). Thus, Sec23p complex appears to interfere with some aspect of membrane fusion, perhaps binding to SNARE molecules, but does not affect vesicle docking. Together, these results suggest that components involved in Uso1p mediated vesicle attachment are distinct from the components involved in vesicle fusion.

Ordering Uso1p, Sec18p, and LMA1 Requirements

Ordering of the Uso1p, Sec18p and LMA1 requirements was pursued in the reconstituted vesicle fusion assay because the semi-intact cell assay does not show a strict dependence on exogenously added Sec18p. If Uso1p tethers ER-derived vesicle to the Golgi compartment, the action

Figure 9. Purified fusion factors mediate distinct steps in vesicle docking and fusion. Saturating amounts of fusion factors (Sec18p [50 ng], Uso1p [75 ng], LMA1 [50 ng]) and 2 ng/μl COPII proteins were mixed in various combinations with semi-intact cells. 35S-labeled gp-α-factor contained in freely diffusible vesicles (black bars) and Golgi modified forms (hatched bars) were determined after 45 min at 23°C. The incubation on the far right (+) contains an additional 8 ng/μl of Sec23p complex.

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Figure 8.

Figure 9.
of Uso1p may produce a dilution resistant intermediate. Indeed, results shown in Fig. 10 support this concept. In this experiment, the concentrations of vesicles, acceptor membranes, and purified proteins were lowered (see Fig. 10, figure legend) to manipulate this intermediate. The maximal fusion efficiency was reduced under these conditions but a clear sensitivity to 10-fold dilution was observed (Fig. 10, ○). Preincubation of vesicles and acceptor membranes with Uso1p alone generated a dilution resistant species that may be chased upon addition of Sec18p and LMA1 (Fig. 10, ■). In contrast, incubation with Sec18p or LMA1 alone does not produce dilution resistance even when adequate levels of the missing fusion factors are supplied in the diluent. These results again suggest Uso1p function is independent from and precedes Sec18p and LMA1 action. Incubation with Uso1p alone followed by dilution with Sec18p and LMA1 (Fig. 10, ■) was not the same as that observed for the complete reaction diluted with buffer containing ATP (Fig. 10, ○). This was due to dilution of the Sec18p and LMA1 proteins under the latter condition whereas active concentrations of these fusion factors were maintained under the former. Therefore, at early time points, Uso1p-docked vesicles efficiently chased due to maintenance of LMA1 and Sec18p throughout the second incubation but docked vesicles for the complete reaction (Fig. 10, ○) were not chased as efficiently because of dilution of Sec18p and LMA1. At later times (20 and 30 min), the fusion efficiency of the Uso1p-docked intermediate was not as efficient as that observed with the complete reaction. This may be because of lability of the docked intermediate in the absence of Sec18p and LMA1.

To determine the temporal requirements for Sec18p and LMA1, the Uso1p-tethered intermediate was generated in the presence of Sec18p or LMA1 for 20 min followed by dilution with the missing factor (Fig. 11). Incubation with Sec18p alone generated a dilution resistant intermediate. LMA1 and Sec18p are then required for the LMA1 requirement. Further support for this idea is shown in Fig. 11, columns 5–8 using a Sec23p blocked intermediate. In column 5, potent inhibition of vesicle fusion upon the addition of Sec23p complex was observed in the reconstituted reaction as was found in the semi-intact cell assay (Figs. 8 and 9). Dilution of this inhibited species with buffer alone or buffer containing Sec18p resulted in a modest increase in fusion efficiency. Strikingly, the addition of diluent containing LMA1 reverses the Sec23p stalled intermediate and promotes vesicle fusion to near maximal levels (Fig. 11, column 7). This result indicates Sec23p inhibition allows for Uso1p-mediated docking and Sec18p action but prevents LMA1 function that leads to vesicle fusion. In summary, these experiments demonstrate that Sec18p function precedes the LMA requirement.

**Discussion**

A cell-free assay that measures vesicle fusion has been reconstituted with Golgi membranes, purified ER-derived vesicles and three soluble proteins: Sec18p, LMA1, and Uso1p. In the presence of ATP and GTP, these proteins fully substitute for a crude yeast cytosol in catalyzing vesicle fusion. In an overall transport reaction performed in yeast semi-intact cells, COPII proteins, and fusion factors are required for efficient transport. Even with saturating levels of budding factors (COPII) or fusion factors (Uso1p, Sec18p, and LMA1), transport remains coupled—budding must precede membrane fusion. Because direct membrane fusion of ER with the Golgi is not efficient in vitro, it is probable that vesicle formation activates components involved in fusion as well as bridging a spatial barrier separating these organelles. This result contrasts reconstituted intra-Golgi transport where membrane fusion can be driven in the absence of COPI, the membrane coat that forms Golgi-derived vesicles (Malhotra et al., 1989; Elazar et al., 1994).

COPII produces freely diffusible ER-derived vesicles from semi-intact cells (Salama et al., 1993) or microsomal membranes (Barlowe et al., 1994). In this report, it is shown that the addition of Uso1p in the presence of vesicles and Golgi acceptor membranes reduces the level of diffusible vesicles and produces a docked, dilution-resistant intermediate. LMA1 and Sec18p are then required for the
fusión de Uso1p docked vesicules. Elevatéd concentrations of the Sec23p complex are found to reversibly inhibit vesicle fusion but do not interfere with Uso1p-mediated docking. Normalmente, the Sec23p complex acts in vesicle formation and is proposed to bind cargo and targeting molecules for packaging into COPII-coated vesicles (Scheikman et al., 1996). Therefore, this inhibitory effect may be due to competition between Sec23p complex and component(s) of the fusion machinery. For example, Sec23p is detected in a specific complex with ER to Golgi v-SNAREs, including Sec22p (Scheikman et al., 1996) and Bet1p (Barlowe, C., unpublished observation). These observations suggest reasonable sites of inhibition to test, however there may be other unknown components of the fusion machinery that interact with the Sec23p complex and prevent vesicle fusion.

Chase of Uso1p-docked, dilution-resistant intermediates indicates a temporal order for the Sec18p and LMA1 requirements. Sec18p function precedes the LMA1 requirement. This was observed whether the dilution-resistant intermediate was accumulated in the presence of Uso1p and Sec18p or through Sec23p complex inhibition. These observations may be summarized in a model for vesicle docking and fusion with the Golgi complex as follows. First, Uso1p tethers freely diffusible vesicles to the acceptor compartment in a reaction that is independent of Sec18p and LMA1. Second, Sec18p activates SNARE molecules contained on the acceptor and/or vesicle membranes. Third, LMA1 action leads to specific SNARE associations that are prevented by elevated Sec23p complex concentrations. The features of Uso1p, Sec18p, and LMA1 are further discussed below in the context of this proposed model.

A requirement for purified Uso1p and a role in vesicle docking is consistent with much of the current literature (for review see Pfeffer, 1996) but has not been directly demonstated until this report. The function of Uso1p as a general fusion factor is not well established, but evidence indicates that a related protein in mammalian cells (termed p115) participates in multiple transport processes including intra-Golgi transport (Water et al., 1993; Sapperstein et al., 1995), the formation of Golgi cisternae after mitotic disassembly (Rabouille et al., 1995), and in transcytotic membrane traffic (Barroso et al., 1995). In yeast, genetic and biochemical evidence clearly implicates Uso1p in ER to Golgi transport, placing the requirement after vesicle formation but before SNARE complex assembly (Nakajima et al., 1991; Lupashin et al., 1996; Sapperstein et al., 1996). Although a role for Uso1p in other transport processes cannot be excluded, the essential role of this protein appears to be in early stages of the yeast secretory pathway since overproduction of specific ER to Golgi SNARE proteins are able to suppress uso1-1 temperature-sensitive mutations as well as an uso1 deletion (Sapperstein et al., 1996).

Based on biochemical and rotary shadowing electron microscopy experiments, p115 forms an extended homodimer with two globular NH2-terminal domains and a parallel coiled-coil rod domain (Sapperstein et al., 1995). Uso1p is a 206-kD protein that shares overall amino acid identity with p115 in both the NH2-terminal globular domains as well as portions of the coiled-coil domains and electron microscopy experiments indicate Uso1p is arranged in a similar parallel homodimer (Yamakawa et al., 1996). Uso1p possesses a predicted coiled-coil rod domain about twice the size of p115. Interestingly, p115 is reported to act in conjunction with a Golgi matrix protein (termed GM130) whose sequence predicts extended coiled-coil structures (Nakamura et al., 1995). It has been suggested that Uso1p represents a fusion of p115 and GM130 although this possibility has not been explored (Nakamura et al., 1997). Because a soluble form of Uso1p is required for vesicle docking in vitro, binding sites for Uso1p are likely contained on purified ER-derived vesicles; the identification of protein(s) comprising this binding site are currently under investigation.

A requirement for Sec18p, the yeast homologue of NSF, was expected from genetic experiments and from analyses of several cell-free membrane fusion reactions that require Sec18p/NSF (Block et al., 1988; Beckers et al., 1989; Diaz et al., 1989; Rexach and Scheikman, 1991; Söllner et al., 1993; Mayer et al., 1996). Sec18p/NSF is a homo-oligomeric ATPase that promotes bilayer fusion through interactions with membrane-bound SNAP (soluble NSF attachment protein) and SNAREs. While a role for Sec18p/NSF activity in separating SNARE protein complexes seems clear (Söllner et al., 1993; Sogaard et al., 1994; Otto et al., 1997), the placement of this activity in the context of membrane docking and fusion is debated. Sec18p/NSF function may be required to activate SNARE molecules before formation of v/t-SNARE complexes (Morgan and Burgoyne, 1995; Mayer et al., 1996; Otto et al., 1997), or to drive membrane fusion after assembly of the v/t-SNARE complex (Söllner et al., 1993; Sogaard et al., 1994). In this report, Sec18p function is not required for vesicle docking and acquisition of dilution resistance, however, Uso1p docking may be independent of SNARE protein function. In support of this notion, experiments with thermosensitive
sed5-1 and sly-1 membranes in the reconstituted assay demonstrate Uso1p-mediated docking does not depend on Sed5p or Sly1p function whereas vesicle fusion requires their action (Cao, X., and C. Barlowe, manuscript in preparation). Thus, the status of ER to Golgi SNARE proteins during different stages of this reconstituted reaction need to be determined before Sec18p/NSF function in activation or bilayer fusion can be unequivocally assigned.

For requirement for LMA1 in the fusion of ER-derived vesicles to the Golgi was unexpected. However, this affect is very reproducible and LMA1 activity appears to depend on activity or bilayer fusion can be unequivocally assigned. LMA1 was initially discovered by fractionation of a yeast cytosol required to drive homotypic vacuole fusion in vitro, and is composed of two polypeptides—thioredoxin plus IB2 (Xu et al., 1996). In cell extracts IB2 can be found in association with thioredoxin (LMA1) or as a monomeric species (LMA2), each capable of stimulating vacuolar membrane fusion (Xu et al., 1997). Additions of LMA1 and Sec18p replaces the requirement for crude cytosol in a fusion reaction between salt washed vacuoles. Results from this system indicate the action of LMA1 and Sec18p are coupled with a Sec18p requirement that precedes LMA1 function (Xu et al., 1997). Experiments with ER to Golgi transport described in this report indicate a similar order for Sec18p and LMA1. Both subunits of LMA1 are needed for normal vacuole inheritance in vivo although neither subunit is essential for cell viability, nor are there detectable delays of secretory protein transport in strains lacking the IB2 subunit (Barlowe, C., unpublished observation). However, there is a potential open reading frame (YHR138c) contained in the yeast genome that shares significant amino acid identity with IB2 and may compensate for the loss of IB2 function. Further in vivo analyses can address this question. Regardless, the involvement of LMA1 in ER to Golgi transport and homotypic vacuolar membrane fusion using very different cell free assays in addition to a functional relationship with Sec18p suggests a general role for this factor in trafficking.

Genetic and biochemical approaches have implicated additional soluble factors in the vesicle fusion stage of ER to Golgi transport in yeast including Ypt1p (Rexach and Schekman, 1991), Sly1p and Sec7p (Lupashin et al., 1996). Furthermore, ADP-ribosylation factor (ARF) and COPI are proposed to operate at this stage in mammalian cells (Aridor et al., 1995), and there is increasing evidence that the protein machinery of ER to Golgi transport is highly conserved between yeast and mammals (Orci et al., 1991; Kuge et al., 1994; Shaywitz et al., 1995; Dasher and Balch, 1996; Hay et al., 1997). Requirements for Ypt1p, Sly1p, Sec7p, ARF, and COPI are not detected through the biochemical approach used here either because of ample peripherally bound species on Golgi membrane preparations or the cell-free assay may not reproduce all of the in vivo requirements. Alternatively, these proteins may not be directly required for vesicle fusion. For Ypt1p, and Sly1p, it seems likely that sufficient protein is contained on acceptor membranes since >90% of these species sediment with the membrane fraction (Cao, X., and C. Barlowe, unpublished observation). In vivo experiments with a version of Ypt1p that contains a transmembrane segment on the COOH terminus, which converts the protein to an integral membrane species, is functional, and suggests that a soluble form of this protein is not essential for membrane fusion (Ossig et al., 1995). Significant pools of soluble Sec7p (Franzusoff et al., 1991) and COPI subunits (Hosobuchi et al., 1992; Stirling et al., 1992) are found in yeast cells. These proteins may cooperate in the production of COPI-coated vesicles since recent data indicates that Sec7p homologues encode guanine nucleotide exchange factors for ARF GTPases (Chardin et al., 1996; Morinaga et al., 1996; Peyroche et al., 1996) and it is this activated GTP-bound form of ARF that triggers assembly of COPI vesicles. In mammalian cells, experiments suggest there is a sequential coupling between COPI and COPII vesicle coats in ER to Golgi transport such that after budding from the ER, the COPII coat is shed and an additional round of coating and uncoating by COPI is required to complete transport of secretory proteins to the cis-Golgi (Aridor et al., 1995; Rowe et al., 1996). Purified, ER-derived vesicles from yeast are not enriched in ARF or COPI subunits (Barlowe et al., 1994). If COPI is required for sequential assembly onto purified, ER-derived vesicles, it must be provided from the acceptor membrane fraction in reconstitution experiments such as shown in Fig. 7 of this report, or there may be no direct requirement for COPI in this assay. Regardless, an acceptor membrane preparation depleted of COPI, ARF, and Sec7p will be required to rigorously address this issue.

The isolated fusion factors represent some, but not all of the genetic requirements for vesicle fusion established by analyses of secretion defect cells. Some of these factors (such as Sly1p and Ypt1p) are peripherally bound to membranes and have required schemes designed to selectively inactivate their function. In addition to the soluble factors and peripherally bound proteins, several of the genetically defined components of vesicle fusion encode integral membrane proteins such as Sec22p, Bet1p, Bos1p, and Sed5p (Dasher et al., 1991; Newman et al., 1990; Shim et al., 1991; Hardwick et al., 1992). To advance the long-term goal of reconstituting this process with pure protein and lipid fractions, the solubilization and reconstitution of ER-derived vesicles will be required.

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