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Erv41p and Erv46p: New Components of COPII Vesicles Involved in Transport between the ER and Golgi Complex

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Abstract. Proteins contained on purified COPII vesicles were analyzed by matrix-assisted laser desorption ionization mass spectrometry combined with database searching. We identified four known vesicle proteins (Erv14p, Bet1p, Emp24p, and Erv25p) and an additional nine species (Yip3p, Rer1p, Erp1p, Erp2p, Erv29p, Yif1p, Erv41p, Erv46p, and Emp47p) that had not been localized to ER vesicles. Using antibodies, we demonstrate that these proteins are selectively and efficiently packaged into COPII vesicles. Three of the newly identified vesicle proteins (Erv29p, Erv41p, and Erv46p) represent uncharacterized integral membrane proteins that are conserved across species. Erv41p and Erv46p were further characterized. These proteins colocalized to ER and Golgi membranes and exist in a detergent-soluble complex that was isolated by immu-

noprecipitation. Yeast strains lacking Erv41p and/or Erv46p are viable but display cold sensitivity. The expression levels of Erv41p and Erv46p are interdependent such that Erv46p was reduced in an *erv41Δ* strain, and Erv41p was not detected in an *erv46Δ* strain. When the *erv41Δ* or *erv46Δ* alleles were combined with other mutations in the early secretory pathway, altered growth phenotypes were observed in some of the double mutant strains. A cell-free assay that reproduces transport between the ER and Golgi indicates that deletion of the Erv41p–Erv46p complex influences the membrane fusion stage of transport.

Key words: ER • Golgi • vesicles • coat proteins • trafficking

Introduction

Membrane-bound transport vesicles are central intermediates for intracellular trafficking of proteins and lipids. The production of many types of transport vesicles depends on coat protein complexes that form vesicles and select the desired set of cargo (Schekman and Orci, 1996). Although the importance of vesicle carriers has been appreciated, the molecular mechanisms underlying distinct stages in vesicle-mediated transport are not well understood. Several approaches have been used to identify and characterize this protein machinery, including the isolation and characterization of specific vesicle carriers involved in synaptic transmission, endocytosis, exocytosis, and intra-Golgi transport (for reviews see Söhlner and Rothman, 1996; Arvan and Castle, 1998; Foletti et al., 1999; Smith and Pearse, 1999). We have undertaken a comprehensive analysis of ER-derived transport vesicles to elucidate the molecular events associated with transport between the ER and Golgi compartments.

Transport between these early compartments of the secretory pathway is bidirectional and mediated by the

COPI and COPII coat complexes. In general, it is thought that the COPII coat buds vesicles from the ER for anterograde transport, whereas the COPI coat is responsible for retrograde transport of recycled proteins from Golgi and pre-Golgi compartments back to the ER (Mellman and Warren, 2000). Formation of COPII-coated vesicles may be reproduced in a cell-free reaction with purified soluble components (the Sar1p GTPase, the Sec23p complex, and the Sec13p complex) and washed ER membranes (Salama et al., 1993; Barlowe et al., 1994). A highly purified preparation of uncoated ER-derived vesicles can be obtained through a scaled up version of the cell-free budding assay. Examination of purified vesicles on protein-stained gels reveals a characteristic set of polypeptides that are solubilized by detergents but not by an elevated pH treatment (Barlowe et al., 1994; Rexach et al., 1994). Some of the abundant ER vesicle (Erv)¹ proteins have been characterized (Barlowe et al., 1994; Schimmöller et al., 1995; Belden and Barlowe, 1996; Powers and Barlowe, 1998) and are found to cycle between the ER and Golgi compartments and function in the processes of vesicle formation and/or

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¹Abbreviations used in this paper: CPY, carboxypeptidase Y; Erv, ER vesicle; gp- α -F, glyco-pro- α factor; HA, hemagglutinin.

site-specific membrane fusion. In this report, we extend our studies to identify additional Erv proteins contained on ER-derived vesicles by mass spectrometry coupled to database searching. Several new Erv proteins were identified and two of these, Erv41p and Erv46p, were characterized further. Erv41p and Erv46p exist in a complex that is probably conserved across species and appears to influence the vesicle fusion stage of transport between the ER and Golgi compartments.

Materials and Methods

Yeast Strains and Media

Strains used for this study are listed in Table I. Unless noted otherwise, cultures were grown at 30°C in rich medium (YPD, 1% Bacto-yeast extract, 2% Bacto-peptone, 2% dextrose) or in minimal medium (YMD, 0.67% yeast nitrogen base without amino acids, 2% dextrose) containing the appropriate amino acid supplements. Standard yeast (Sherman, 1991) and cloning protocols (Ausubel et al., 1987) were used.

Plasmid Construction

Primer sequences are listed in Table II. The sequences of *ERV41* (YML067c) and *ERV46* (YAL042w) were obtained from the *Saccharomyces* Genome Database. To generate plasmid pRS424-*ERV41*, the *ERV41* sequence, and ~300 bp of its flanking upstream and downstream, sequences were cloned from genomic DNA prepared from strain RSY255 using primers YML067c-NotI and YML067c-BamHI. The product was ligated into the NotI and BamHI restriction sites of the pRS424 vector (Christianson et al., 1992). *ERV46* and its flanking regions of ~300 bp were amplified using primers YAL042w-NotI and YAL042w-BamHI and inserted into the NotI and BamHI sites of the pRS426 vector (Christianson et al., 1992) to yield plasmid pRS426-*ERV46*. Correct amplification and integration were verified by DNA sequencing.

For the overexpression of fusion proteins bearing NH₂-terminal 6x histidine tags, fragments of *ERV41*, *ERV46*, and *OCH1* were inserted into the BamHI and HindIII restriction sites of the pQE-30 vector (QIAGEN) to yield plasmids pQE-30-*ERV41*, pQE-30-*ERV46*, and pQE-30-*OCH1*, respectively. Primers YML067c-pQE-F and YML067c-pQE-R for *ERV41*, YAL042w-pQE-F and YAL042w-pQE-R for *ERV46*, and primers OCH1-pQE-F and OCH1-pQE-R for *OCH1* were used to amplify portions of these genes from genomic RSY255 DNA.

Strain Construction

ERV41 was targeted for disruption with the *HIS3* gene (Baudin et al., 1993). An *erv41::HIS3* construct was amplified using primers YML067c-KO-F and YML067c-KO-R and pHISKO as a template. The product contains the *HIS3* gene flanked by 43-bp upstream of the *ERV41* start codon and 45-bp downstream of the *ERV41* stop codon and was used to transform CBY453 cells. Several transformants showing histidine prototrophy were screened by PCR using primers YML067c-NotI and *HIS3*. One (CBY763) tested positive and was sporulated. Strains carrying *erv46Δ*, *rer1Δ*, or *erv29Δ* null alleles disrupted with a *KAN* marker were generated by a deletion project (Winzeler et al., 1999; Research Genetics) and crossed several times through the FY833/FY834 background to yield an isogenic set of spores.

Erv46p, Erv41p, and Erv29p were tagged at their NH₂ termini by the addition of three repeated influenza virus hemagglutinin (HA) epitopes and put under control of the *GALI* promoter (Longtine et al., 1998); plasmid pFA6a-His3MX6-PGAL1-3HA and primers YAL042w-F4 and YAL042w-R3 were used to amplify the construct targeted to the *ERV46* locus. CBY453 cells were transformed with this product and selected for histidine prototrophy. Several isolates were grown in YP with 2% galactose and 0.2% glucose and screened for expression of tagged protein by Western blot of membrane fractions using an anti-HA antibody. Approximately 50% of transformants tested positive, and one (CBY767) was sporulated to yield strains CBY770 and CBY771. *ERV41* was targeted with a PCR product obtained using plasmid pFA6a-TRP1-PGAL1-3HA and primers YML067c-F4 and YML067c-R3. Transformants were selected for tryptophan prototrophy and screened for expression of HA-tagged protein. One positive diploid strain (CBY782) was sporulated

to obtain strains CBY783 and CBY784. Primers YGR284c-F4 and YGR284c-R3 and plasmid pFA6a-His3MX6-PGAL1-3HA were used to amplify the construct directed to the *ERV29* locus. CBY453 cells were transformed and selected for growth on minimal media lacking histidine. Three transformants were screened by Western blot with the anti-HA antibody, and one tested positive (CBY950). Yif1p was tagged by the insertion of three HA epitopes at its COOH terminus (Longtine et al., 1998), using primers MHF2 and MHR1 and plasmid pFA6a-3HA-His3MX6 as template. FY834 cells were transformed with the PCR product and selected for histidine prototrophy. 7 out of 10 transformants expressed tagged protein, and one was analyzed further (CBY801).

Genetic Analyses

To generate strains carrying multiple mutations, *erv14Δ*, *erv41Δ*, and *erv46Δ*, strains were mated with other mutants. If possible, diploids were selected using markers of the parent strains, otherwise, zygotes were picked under the microscope. Diploid strains were sporulated, and asci were dissected on YPD plates using a micromanipulator. Plates were incubated at room temperature, and germinated spores were scored for mating types, markers, and growth on YPD plates at 16°C, room temperature, 30°C, 36°C, and 38°C. In cases where several loci had been replaced by the *HIS3* gene, deletions were scored by PCR using YML067c-NotI and the *HIS3* internal primer for the detection of the *erv41::HIS3* allele and the *ERV14* specific primer GP3 (Powers and Barlowe, 1998) and the internal *HIS3* primer to detect the *erv14::HIS3* deletion. In cases where one of the parent strains had a genetic background different from FY833/FY834, spores were backcrossed several times to obtain isogenic strains.

Antibodies and Immunoblotting

Polyclonal antibodies were raised against 6x histidine-tagged NH₂-terminal fusion proteins of fragments of Erv41p (amino acid positions 75–274), Erv46p (amino acid positions 80–296), and Och1p (amino acid positions 302–480) expressed from plasmids pQE-30-*ERV41*, pQE-30-*ERV46*, and pQE-30-*OCH1*, respectively. All of the recombinant proteins localized to the insoluble fraction of cells disrupted in a French Press. These fractions were solubilized with 8 M urea, and the fusion proteins were purified on Ni-NTA agarose (QIAGEN) as recommended by the manufacturer. The recombinant proteins were used to immunize rabbits according to standard procedures. For Western blotting, these antisera were diluted 1:1,000.

Antibodies directed against carboxypeptidase Y (CPY), Emp47p, Erv14p, Erv25p, Kar2p, plasma membrane ATPase, Sec12p, Sec23p, Sec61p (Powers and Barlowe, 1998), Bos1p (Cao and Barlowe, 2000), Rer1p (Boehm et al., 1997), and Yip1p (Yang et al., 1998) were described earlier. A monoclonal anti-HA antibody was obtained from Berkeley Antibody Co. Western blots were developed using the ECL method (Amersham Pharmacia Biotech). For densitometric analysis, films were scanned and plotted using NIH Image 1.52.

Subcellular Fractionation

Membrane fractions were prepared by the bead-beat method in lysis buffer (25 mM Hepes, pH 7.0, 50 mM potassium acetate, 2 mM EDTA, 1 mM PMSF). Pellets were resolved on 12.5% polyacrylamide gels. Microsomes (Wuestehube and Schekman, 1992) and semiintact cells (Baker et al., 1988) were prepared as described. Sucrose gradient fractionation of membrane organelles was performed according to Powers and Barlowe (1998). To determine whether Erv41p and Erv46p are integral membrane proteins, semiintact FY834 cells were suspended in buffer (20 mM Hepes, pH 7.0, 150 mM potassium acetate, 2 mM EDTA), in buffer with 1% Triton X-100, or in 0.1 M sodium carbonate, pH 11.0, 2 mM EDTA, incubated on ice for 10 min and centrifuged at 60,000 rpm in a TLA100.3 rotor (Beckman Coulter) for 12 min. Equivalent amounts of the total, supernatant, and pellet fractions were resolved on a 12.5% polyacrylamide gel.

In Vitro Vesicle Budding and Transport Assays

The synthesis of COPII vesicles was performed on a preparative scale by incubating washed microsomes in the absence or presence of the proteins Sar1p, Sec13–31p complex, and Sec23–24p complex as described previously (Barlowe et al., 1994; Belden and Barlowe, 1996). After collection of fractions from nycodenz density gradients, peak fractions were diluted fourfold with buffer 88 and centrifuged at 60,000 rpm for 25 min in a TLA100.3 rotor (Beckman Coulter) to collect vesicles. The membrane pel-

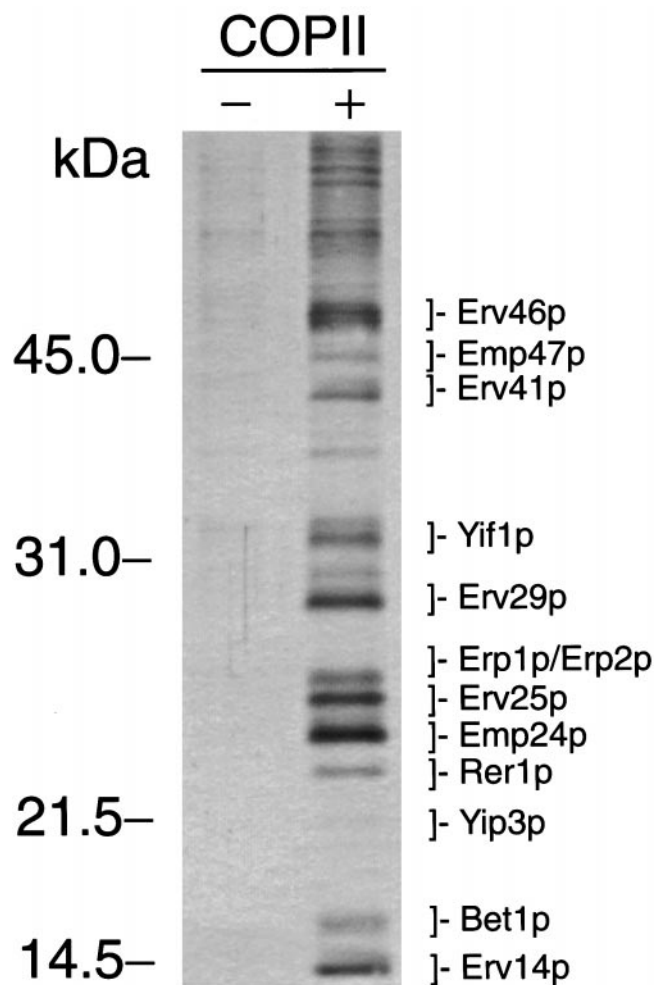


Figure 1. Identification of Erv proteins. COPII-coated vesicles were synthesized *in vitro* from ER membranes in the presence of COPII proteins (+), were solubilized in sample buffer, and were resolved on a 15% polyacrylamide gel. As a negative control, a mock reaction without COPII proteins (–) was performed. Proteins were silver stained for this figure. For mass spectroscopy, proteins were stained with colloidal blue, and individual bands were excised.

lets were dissolved in 40 μ l of sample buffer, and solubilized proteins were resolved on a 15% polyacrylamide gel (Novex). Gels were developed with a colloidal blue staining kit (Novex), and polypeptide staining bands were excised for analysis by mass spectrometry as described (Jensen et al., 1999). Analytical scale budding reactions were performed as described (Barlowe et al., 1994), and packaging efficiencies were determined by densitometry of scanned blots. Vesicle tethering and fusion assays following 35 S-labeled glyco-pro- α factor (gp- α -F) were described by Cao et al. (1998). The data plotted in these experiments are the average of duplicate determinations, and the error bars represent the range. Pulse-chase experiments were performed according to Belden and Barlowe (1996).

Immunoprecipitation Experiments

Microsomes were solubilized on ice with buffer 88 containing 2% Triton X-100 and 1 mM PMSF for 5 min, followed by a centrifugation at 14,000 g for 5 min. Portions (25 μ l) of the supernatant were mixed with 1 ml of IP buffer (15 mM Tris/HCl, pH 7.5, 150 mM NaCl, 1% Triton X-100), 25 μ l of a 50% protein A-Sepharose slurry (Amersham Pharmacia Biotech), and a saturating amount of anti-HA antibody and were incubated at 4°C for 2 h. The precipitates were washed four times with cold IP buffer, eluted from the beads by heating at 95°C in sample buffer, and resolved on polyacrylamide gels.

Results

Identification of ERV Proteins

A protocol to generate ER-derived vesicles *in vitro* from washed microsomes and purified COPII components (Barlowe et al., 1994) was scaled up to prepare sufficient amounts of material to observe polypeptides on protein-stained polyacrylamide gels (Belden and Barlowe, 1996). A set of polypeptides was observed (Fig. 1), and individual bands were subjected to tryptic peptide mass mapping by matrix-assisted laser desorption/ionization mass spectrometry followed by sequence database searching as described (Shevchenko et al., 1996; Jensen et al., 1999). Several proteins were identified, some that had been previously described and others that had not been characterized. Importantly, three of these proteins (Erv14p, Emp24p, and Erv25p) had been identified in our initial approaches using automated NH₂-terminal sequencing (Belden and Barlowe, 1996; Powers and Barlowe, 1998). Therefore, the isolated ER-derived vesicles are comparable to previous preparations, which suggests that this method should allow for the identification of additional Erv proteins. The SNARE protein Bet1p (Newman and Ferro-Novick, 1987) was also identified in this preparation and had been previously detected on ER-derived vesicles by immunoblot analysis and is required for transport between the ER and Golgi (Newman et al., 1992; Cao and Barlowe, 2000).

Nine other polypeptides identified in this preparation had not been localized to ER-derived vesicles before. Yip3p, Rer1p, Erp1p, Erp2p, Yif1p, and Emp47p are conserved proteins that had been characterized and are thought to operate in transport through the early secretory pathway (Sato et al., 1995; Schröder et al., 1995; Andrulis et al., 1998; Yang et al., 1998; Marzioch et al., 1999). Erv29p, Erv41p, and Erv46p represent uncharacterized proteins. Peptides derived from the Erv29p band correspond to the ORF YGR284c that encodes a nonessential 35-kD integral membrane protein terminating in the sequence KKKIY. According to its apparent molecular weight on polyacrylamide gels, we have designated this protein Erv29p. YML067c encodes an uncharacterized protein that has recently been immunisolated from an early Golgi compartment (Cho et al., 2000). We have designated this protein Erv41p according to its molecular weight. YAL042w encodes a protein of unknown function that was named Fun9p (“function unknown”) (Coleman et al., 1986; Diehl and Pringle, 1991) but not further characterized. We propose to name this protein Erv46p referring to its localization and molecular weight.

Selective Packaging of Erv Proteins *In Vitro*

Authentic Erv proteins should be selectively and efficiently packaged into ER-derived vesicles in the presence of COPII proteins. To test specific packaging of the uncharacterized proteins, we first generated strains that contained epitope-tagged versions of these proteins. Strain CBY801 expresses a COOH terminally HA-tagged version of Yif1p that fully complements for *YIF1* function. Erv29p (CBY950), Erv41p (CBY782), and Erv46p (CBY767) were modified to contain the 3HA epitope at their NH₂ termini and were placed under the control of

Table I. Strain List

Strain	Genotype	Reference
FY833	<i>MATa his3200 ura3-52 leu21 lys2202 trp163</i>	Winston et al. (1995)
FY834	<i>MATα his3200 ura3-52 leu21 lys2202 trp163</i>	Winston et al. (1995)
RSY255	<i>MATα ura3-52 leu2-3,-112</i>	Kaiser and Schekman (1990)
RSY263	<i>MATα sec12-4 ura3-52 leu2-3,112</i>	Kaiser and Schekman (1990)
RSY265	<i>MATα sec13-1 ura3-52 his4-619</i>	Kaiser and Schekman (1990)
RSY267	<i>MATα sec16-2 ura3-52 his4-619</i>	Kaiser and Schekman (1990)
RSY277	<i>MATα sec21-1 ura3-52</i>	Kaiser and Schekman (1990)
RSY281	<i>MATα sec23-1 ura3-52 his4-619</i>	Kaiser and Schekman (1990)
RSY309	<i>MATα sec12-1 leu2-3,112</i>	Kaiser and Schekman (1990)
RSY962	<i>MATa sec35-1 lys2-801</i>	Wuestehube et al. (1996)
RSY976	<i>MATa ura3-52 ypt1-3</i>	Wuestehube et al. (1996)
CBY99	FY834 with <i>emp24::LEU2</i>	Belden and Barlowe (1996)
CBY263	<i>MATα trp1-1 ade2-1 ura3-1 leu2-3,112 can1-100 sed5-1</i>	Cao et al. (1998)
CBY300	FY834 with <i>uso1-1</i>	Barlowe (1997)
CBY356	FY834 with <i>erv14::HIS3</i>	Powers and Barlowe (1998)
CBY358	FY833 with <i>erv14::HIS3</i>	Powers and Barlowe (1998)
CBY453	FY833 x FY834	Powers and Barlowe (1998)
CBY763	CBY453 diploid with <i>ERV41/erv41::HIS3</i>	This study
CBY767	CBY453 diploid with <i>ERV46/ERV46::HIS3MX6-PGAL1-3HA</i>	This study
CBY770	FY834 with <i>ERV46::HIS3MX6-PGAL1-3HA</i>	This study
CBY771	FY833 with <i>ERV46::HIS3MX6-PGAL1-3HA</i>	This study
CBY782	CBY453 diploid with <i>ERV41/ERV41::TRP1-PGAL1-3HA</i>	This study
CBY783	FY834 with <i>ERV41::TRP1-PGAL1-3HA</i>	This study
CBY784	FY833 with <i>ERV41::TRP1-PGAL1-3HA</i>	This study
CBY794	FY833 with <i>erv41::HIS3 erv46::KAN</i>	This study
CBY795	FY834 with <i>erv41::HIS3 erv46::KAN</i>	This study
CBY796	FY833 with <i>erv41::HIS3</i>	This study
CBY797	FY834 with <i>erv41::HIS3</i>	This study
CBY798	FY833 with <i>erv46::KAN</i>	This study
CBY799	FY834 with <i>erv46::KAN</i>	This study
CBY801	FY834 with <i>YIF1::HIS3MX6-3HA</i>	This study
CBY822	FY833 with <i>erv14::HIS3 erv46::KAN</i>	This study
CBY823	FY834 with <i>erv14::HIS3 erv46::KAN</i>	This study
CBY825	FY834 with <i>erv14::HIS3 erv41::HIS3</i>	This study
CBY826	FY833 with <i>erv14::HIS3 erv41::HIS3</i>	This study
CBY829	FY834 with <i>ypt1-3</i>	This study
CBY831	FY833 with <i>emp24::LEU2 erv41::HIS3 erv46::KAN</i>	This study
CBY832	FY834 with <i>emp24::LEU2 erv41::HIS3 erv46::KAN</i>	This study
CBY836	CBY822 containing pRS426-ERV46	This study
CBY840	FY833 with <i>erv41::HIS3 ypt1-3</i>	This study
CBY841	FY834 with <i>erv41::HIS3 ypt1-3</i>	This study
CBY842	FY833 with <i>erv46::KAN ypt1-3</i>	This study
CBY843	FY834 with <i>erv46::KAN ypt1-3</i>	This study
CBY844	FY833 with <i>erv41::HIS3 erv46::KAN ypt1-3</i>	This study
CBY845	FY834 with <i>erv41::HIS3 erv46::KAN ypt1-3</i>	This study
CBY847	CBY826 containing pRS424-ERV41	This study
CBY848	FY833 with <i>erv41::HIS3 sec21-1</i>	This study
CBY849	FY834 with <i>erv41::HIS3 sec21-1</i>	This study
CBY850	FY833 with <i>erv46::KAN sec21-1</i>	This study
CBY851	FY834 with <i>erv46::KAN sec21-1</i>	This study
CBY852	FY834 with <i>erv41::HIS3 erv46::KAN sec21-1</i>	This study
CBY853	FY833 with <i>erv41::HIS3 sed5-1</i>	This study
CBY854	FY833 with <i>ade2-1 erv46::KAN sed5-1</i>	This study
CBY855	FY834 with <i>erv46::KAN sed5-1</i>	This study
CBY856	FY833 with <i>erv41::HIS3 erv46::KAN sed5-1</i>	This study
CBY857	FY834 with <i>ade2-1 erv41::HIS3 erv46::KAN sed5-1</i>	This study
CBY859	FY833 with <i>erv41::HIS3 sec12-1</i>	This study
CBY860	FY834 with <i>erv41::HIS3 sec12-1</i>	This study
CBY861	FY833 with <i>erv46::KAN sec12-1</i>	This study
CBY862	FY834 with <i>erv46::KAN sec12-1</i>	This study
CBY863	FY833 with <i>erv41::HIS3 erv46::KAN sec12-1</i>	This study
CBY877	FY833 with <i>erv14::HIS3 rer1::KAN</i>	This study
CBY878	FY834 with <i>erv14::HIS3 rer1::KAN</i>	This study
CBY884	FY833 with <i>erv41::HIS3 nce102::KAN</i>	This study
CBY885	FY834 with <i>erv41::HIS3 nce102::KAN</i>	This study

Table 1. continued

Strain	Genotype	Reference
CBY893	FY833 with <i>erv14::HIS3 erv41::HIS3 erv46::KAN</i>	This study
CBY894	FY834 with <i>erv14::HIS3 erv41::HIS3 erv46::KAN</i>	This study
CBY909	FY833 with <i>erv46::KAN uso1-1</i>	This study
CBY910	FY834 with <i>erv46::KAN uso1-1</i>	This study
CBY911	FY833 with <i>erv41::HIS3 uso1-1</i>	This study
CBY912	FY834 with <i>erv41::HIS3 uso1-1</i>	This study
CBY913	FY833 with <i>erv41::HIS3 erv46::HIS3 uso1-1</i>	This study
CBY914	FY834 with <i>erv41::HIS3 erv46::HIS3 uso1-1</i>	This study
CBY915	FY833 with <i>erv46::KAN sec23-1</i>	This study
CBY916	FY834 with <i>erv46::KAN sec23-1</i>	This study
CBY917	FY833 with <i>erv41::HIS3 sec23-1</i>	This study
CBY918	FY833 with <i>erv41::HIS3 erv46::KAN sec23-1</i>	This study
CBY919	FY834 with <i>erv41::HIS3 erv46::KAN sec23-1</i>	This study
CBY920	FY833 with <i>erv46::KAN sec16-2</i>	This study
CBY921	FY834 with <i>erv46::KAN sec16-2</i>	This study
CBY922	FY833 with <i>erv41::HIS3 sec16-2</i>	This study
CBY923	FY834 with <i>erv41::HIS3 sec16-2</i>	This study
CBY924	FY834 with <i>erv41::HIS3 erv46::KAN sec16-2</i>	This study
CBY930	FY834 with <i>erv41::HIS3 erv46::KAN sec13-1</i>	This study
CBY931	FY833 with <i>erv41::HIS3 erv46::KAN sec13-1</i>	This study
CBY932	FY834 with <i>erv41::HIS3 sec13-1</i>	This study
CBY933	FY834 with <i>erv46::KAN sec13-1</i>	This study
CBY934	FY833 with <i>erv46::KAN sec13-1</i>	This study
CBY940	FY833 with <i>erv46::KAN sec35-1</i>	This study
CBY941	FY834 with <i>erv46::KAN sec35-1</i>	This study
CBY942	FY833 with <i>erv41::HIS3 sec35-1</i>	This study
CBY943	FY834 with <i>erv41::HIS3 sec35-1</i>	This study
CBY944	FY833 with <i>erv41::HIS3 erv46::KAN sec35-1</i>	This study
CBY945	FY834 with <i>erv41::HIS3 erv46::KAN sec35-1</i>	This study
CBY948	FY833 with <i>erv41::HIS3 rer1::KAN</i>	This study
CBY949	FY834 with <i>erv41::HIS3 rer1::KAN</i>	This study
CBY950	CBY453 diploid with <i>ERV29/ERV29::HIS3MX6-PGAL1-3HA</i>	This study
CBY963	FY833 with <i>erv14::HIS3 rer1::KAN</i>	This study
CBY964	FY834 with <i>erv14::HIS3 rer1::KAN</i>	This study
CBY965	FY833 with <i>erv29::KAN</i>	This study
CBY966	FY834 with <i>erv29::KAN</i>	This study
CBY967	FY833 with <i>erv29::KAN erv41::HIS3</i>	This study
CBY968	FY834 with <i>erv29::KAN erv41::HIS3</i>	This study
CBY969	FY833 with <i>erv14::HIS3 erv29::KAN</i>	This study
CBY970	FY834 with <i>erv14::HIS3 erv29::KAN</i>	This study

the *GAL1* promoter (Longtine et al., 1998). Galactose induction resulted in stable expression of these tagged proteins and localization to ER membranes. However, we cannot assess if these proteins are fully functional because *ERV29*, *ERV41*, and *ERV46* are not essential. In vitro budding experiments were performed with microsomes from these strains (Fig. 2 A), and the relative packaging efficiencies were determined by comparing the lanes that represent 10% of the complete reactions (T) with the lanes that contain vesicles synthesized in the presence of COPII (+). Yif1p-3HA (8%), 3HA-Erv29p (13%), 3HA-Erv46p (10%), and 3HA-Erv41p (13%) were incorporated into vesicles at levels comparable to the positive controls Erv25p (14% for CBY801 and 5% for CBY950) and the v-SNARE Bos1p (11 for CBY767 and 5% for CBY782), whereas the ER resident proteins Sec12p and Sec61p were excluded. These results indicate that Yif1p, Erv29p, Erv41p, and Erv46p are selectively packaged into COPII vesicles.

Next, we prepared or obtained specific polyclonal antibodies against several of the newly identified Erv proteins

to measure their packaging efficiencies when endogenously expressed. As seen in Fig. 2 B, vesicles budded from wild-type microsomes contained Erv41p and Erv46p at levels (20% and 26%, respectively) that are higher than those of HA-tagged versions. Yip1p was packaged very efficiently (65%), as was Rer1p (51%; Sato et al., 1995). The early Golgi marker Och1p (Nakayama et al., 1992) was packaged to a significantly lesser extent (8%). The positive controls Erv14p, Erv25p, and Bos1p were efficiently packaged, whereas the ER marker proteins Sec12p, Sec61p, and Kar2p were excluded. Thus, Erv29p, Erv41p, Erv46p, Yif1p, Yip1p, and Rer1p meet the initial requirement of Erv proteins as they are selectively exported from ER membranes under reconstituted vesicle budding conditions.

Molecular Characterization of Erv41p and Erv46p

For the remainder of this report, we investigate the Erv41p and Erv46p proteins and their potential role in transport through the early secretory pathway. Erv29p will be described elsewhere. Erv41p is encoded by the ORF YML067c on chromosome XIII, and conceptual transla-

Table II. Primer Sequences

Primer	Sequence
YML067c-NotI	5'-ATAAGAATGCGGCCGCAACTCCCCTTCTCCAGACTTG-3'
YML067c-BamHI	5'-CGCGGATCCGCGAAGTATGCGGGACGCTG-3'
YAL042w-NotI	5'-ATAAGAATGCGGCCGCTGCAGATGACATTGCGCTGC-3'
YAL042w-BamHI	5'-CGCGGATCCGCCATGATCTCTCGGGTTGG-3'
YML067c-pQE-F	5'-CGCGGATCCACGAAATGTGATTGGTTGC-3'
YML067c-pQE-R	5'-CCCAAGCTTCACCTTTAGCAGCGACATC-3'
YAL042w-pQE-F	5'-CGCGGATCCTGTGACCTGGTGAATCTCG-3'
YAL042w-pQE-R	5'-CCCAAGCTTCCTGTCCGGGAACACC-3'
OCH1-pQE-F	5'-CGCGGATCCGATCCAAGATTTGAAGAAG-3'
OCH1-pQE-R	5'-CCCAAGCTTGGGTATGATGAAAGGAGAG-3'
YML067c-KO-F	5'-TGCTTTTGTCTGTAACGTACAGGCAGAGACTTCAGGAGTATATAGGCCTCTCTAGTACACTC-3'
YML067c-KO-R	5'-GAAACGTTATCTTGGGTATACTGCATTCTCTTTTCTTTTACATATGCGCGCTCGTTCAAGATG-3'
HIS3	5'-GCCTCATCCAAAGGCGC-3'
YAL042w-F4	5'-CGCCATTATAAAGGAAAGCTAGTTTATGTCTCTGATACAGAATTCGAGCTCGTTTAAAC-3'
YAL042w-R3	5'-TAGCGAATGCGTCCAGCGACAGCAACGTGGACCTTTTCATGCACTGAGCAGCGTAATCTG-3'
YML067c-F4	5'-GATATTATCAATGTCTATTCTTACTGAGAAAAGCGTTCCGAATTCGAGCTCGTTTAAAC-3'
YML067c-R3	5'-ATAACATACGAAACGCATCAAAATGTCTTCAATCCTGCCATGCACTGAGCAGCGTAATCTG-3'
YGR284c-F4	5'-ACGAAAGATCAAAAGGTGTCCTTATTTACTTACAATAGCTGGAATTCGAGCTCGTTTAAAC-3'
YGR284c-R3	5'-GGCATACCGCCAAAATTTCCAATAGTCTCTGTAAGACATGCACTGAGCAGCGTAATCTG-3'
MHF2	5'-TGGCTTCATTGCGCAAAATGTTCTAATGTGGTAAATGGGTCGGATCCCCGGGTAAATTA-3'
MHR1	5'-GCATGAAATTAATCTCTCTTTGATCTCTTCAATCAAGAGAATTCGAGCTCGTTTAAAC-3'

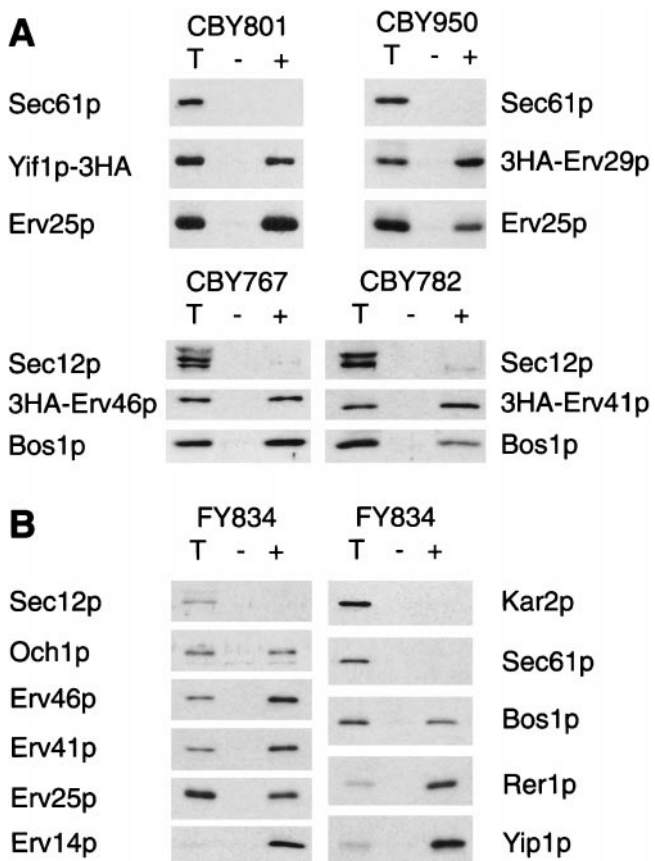


Figure 2. Selective packaging of Erv proteins into COPII-coated vesicles in vitro. (A) In vitro budding reactions with microsomes prepared from strains expressing tagged versions of Erv proteins: Yif1p-3HA (CBY801), 3HA-Erv29p (CBY950), 3HA-Erv46p (CBY767), and 3HA-Erv41p (CBY782). One tenth of a total reaction (T), budded vesicles isolated after incubation with COPII proteins (+), or a mock reaction without COPII proteins (-) were separated on a 12.5% polyacrylamide gel. Tagged proteins were visualized by immunoblot with an anti-HA antibody, and

tion yields a protein of 352-amino acid residues and a predicted molecular weight of 40.6 kD. The ORF YAL042w on chromosome I encodes Erv46p, a protein predicted to be composed of 415-amino acid residues with a molecular weight of 46 kD. Database alignments showed that Erv41p and Erv46p exhibit significant sequence similarity. Both proteins also have one putative homologue each in *Caenorhabditis elegans*, *Drosophila melanogaster*, and humans, however, no known functions have been reported for these proteins. Sequence identity scores (Table III) between Erv41p and its homologues (CDA14, EG:65F1.1 and C18B12.6) are higher than those between Erv41p and Erv46p. Similarly, Erv46p and its homologues in other species (the 43.2-kD protein, CG70, and K09E9.2) show even higher identity scores. The Erv46p group is further characterized by eight conserved cysteine residues in their NH₂-terminal half. Erv46p also terminates in the COPI binding motif KKXX (Jackson et al., 1990; Cosson and Letourneur, 1994), a feature that is conserved across species. These results suggest that a conserved set of Erv41p-Erv46p proteins are found in other species and are likely to perform a similar function.

ERV41 as well as *ERV46* were amplified from genomic DNA and inserted into the multicopy shuttle vectors pRS424 and pRS426, respectively, to yield plasmids pRS424-*ERV41* and pRS426-*ERV46*. Polyclonal antisera were raised against recombinant forms of Erv41p and Erv46p and then tested on wild-type, deletion, and over-

Sec61p or Sec12p (ER resident proteins) as negative controls and Erv25p (Erv protein) or Bos1p (v-SNARE) as positive controls were detected using polyclonal antisera. (B) In vitro budding reactions with FY834 wild-type microsomes. The same budding protocol as in A was used. Proteins were detected with polyclonal antisera against Sec12p, Kar2p, and Sec61p (ER residents), Bos1p (v-SNARE), Och1p (early Golgi marker), and the Erv proteins Erv46p, Erv41p, Erv25p, Erv14p, Rer1p, and Yip1p.

Table III. Percent Sequence Identities between *Erv46p*, *Erv41p*, and Their Homologues

	Erv46p	Erv41p	K09E9.2	C18B12.6	CG7011	EG:65F1.1	43.2 kD	CDA14
Erv46p (<i>S. cerevisiae</i>)	100	28	38	12	38	27	41	25
Erv41p (<i>S. cerevisiae</i>)		100	32	25	28	32	31	30
K09E9.2 (<i>C. elegans</i>)			100	20	41	27	43	32
C18B12.6 (<i>C. elegans</i>)				100	20	20	17	21
CG7011 (<i>D. melanogaster</i>)					100	27	47	34
EG:65F1.1 (<i>D. melanogaster</i>)						100	31	35
43.2 kD protein (<i>H. sapiens</i>)							100	38
CDA14 (<i>H. sapiens</i>)								100

Full-length sequences were aligned using the CLUSTAL V algorithm of GeneInspector v1.5 with the BLOSUM62 table and default settings (k-tuple 1, maximum gap length 5, gap penalty 3, 5 top diagonals).

producing strains to confirm their specificity (Fig. 3). No immunoreactive species were detected in the deletion strains, whereas the heterozygous diploid strains carrying one wild-type and one tagged copy of the respective genes expressed two immunoreactive species at the expected size. This result also demonstrated that expression levels from the *GALI* promoter were comparable to wild-type expression levels at a concentration of 1.5% galactose. Strains harboring *ERV41* or *ERV46* on 2 μ based multi-copy plasmids expressed slightly higher levels of Erv41p or Erv46p than wild-type strains.

Both Erv41p and Erv46p contain two segments of sufficient length and hydrophobicity to form transmembrane domains, and the short NH₂- and COOH-terminal regions of both proteins are predicted to be located on the cytoplasmic side with a large luminal domain separating the two transmembrane regions. The hydrophilicity plots of both proteins are superimposable (Fig. 4 A), and a similar topology is predicted for all of the known homologues. There is no evidence for a cleavable signal sequence. To confirm that Erv41p and Erv46p are indeed integral membrane proteins, we examined their fractionation behavior under conditions that release luminal and peripherally bound membrane proteins (carbonate buffer, pH 11.0) or solubilize integral membrane proteins (1% Triton X-100). The fractionation profile for Erv41p and Erv46p was the same as for the integral membrane protein Bos1p, as they could only be solubilized by detergent and not by carbonate treatment (Fig. 4 B).

The abundant Erv proteins are hypothesized to cycle between the ER and Golgi (Belden and Barlowe, 1996; Powers and Barlowe, 1998) because they localize to both of these membrane compartments, and, in contrast to abundant secretory proteins, their levels are not diminished by cycloheximide treatment (Yeung et al., 1995). Therefore, we examined the subcellular localization of Erv41p and Erv46p by resolution of membrane organelles on sucrose gradients (Fig. 5). Erv41p and Erv46p sedimented in two peaks, one that coincided with the Golgi marker Emp47p and the other with the ER marker Kar2p. This subcellular localization pattern was similar to that of Erv14p (Powers and Barlowe, 1998) and Erv25p (Belden and Barlowe, 1996). Rer1p has been shown to be predominantly localized to the Golgi (Sato et al., 1995) and to recycle between the Golgi and ER (Boehm et al., 1997). Consequently, we find that the majority of Rer1p cosedimented with Emp47p, and a second smaller peak cosedimented with Kar2p. In contrast, Och1p was localized almost exclusively to the Golgi. The localization of the epitope-tagged Erv46p was also examined by indirect immunofluorescence using the anti-HA antibody (data not shown). We observed a perinuclear staining pattern that partially overlapped with the ER marker Kar2p and was similar to that described for Erv14p (Powers and Barlowe, 1998). We conclude that Erv41p and Erv46p localize to the early secretory pathway.

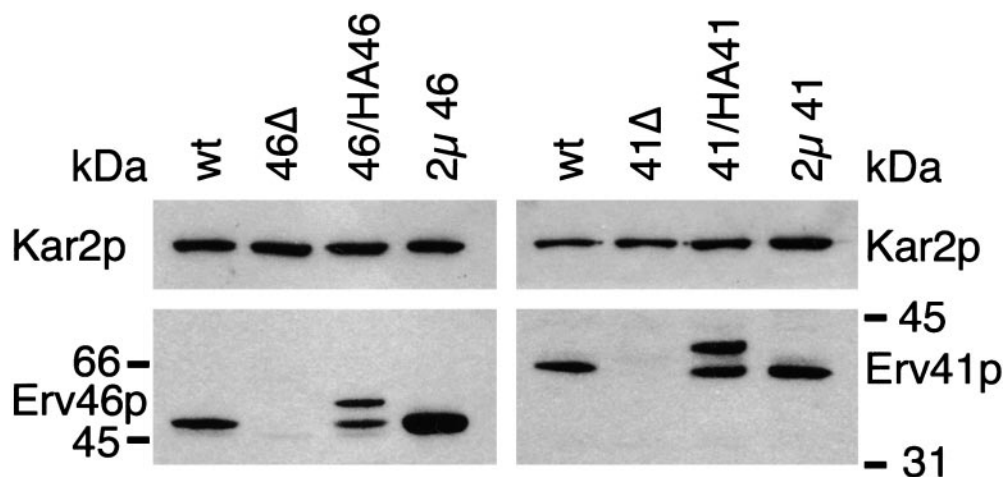


Figure 3. Immunoblot analysis of wild-type, *erv41Δ*, and *erv46Δ* deletion and over-producing strains. FY834 wild-type (wt), *erv46Δ* (46Δ, CBY799), and *erv41Δ* (41Δ, CBY797) strains were grown in YPD medium. Heterozygous diploid strains expressing NH₂ terminally 3HA-tagged Erv46p (46/HA46, CBY767) or Erv41p (41/HA41, CBY782) were grown in YP with 1.5% galactose and 0.5% glucose. *erv14Δ* *erv46Δ* deletion strains car-

rying pRS426-*ERV46* plasmid (2 μ 46, CBY836) or pRS424-*ERV41* plasmid (2 μ 41, CBY847) were grown in minimal media lacking uracil or tryptophan, respectively. Membrane fractions were resolved on a 12.5% polyacrylamide gel and immunoblotted with an anti-Kar2p antiserum as loading control and with polyclonal antisera raised against Erv46p or Erv41p.

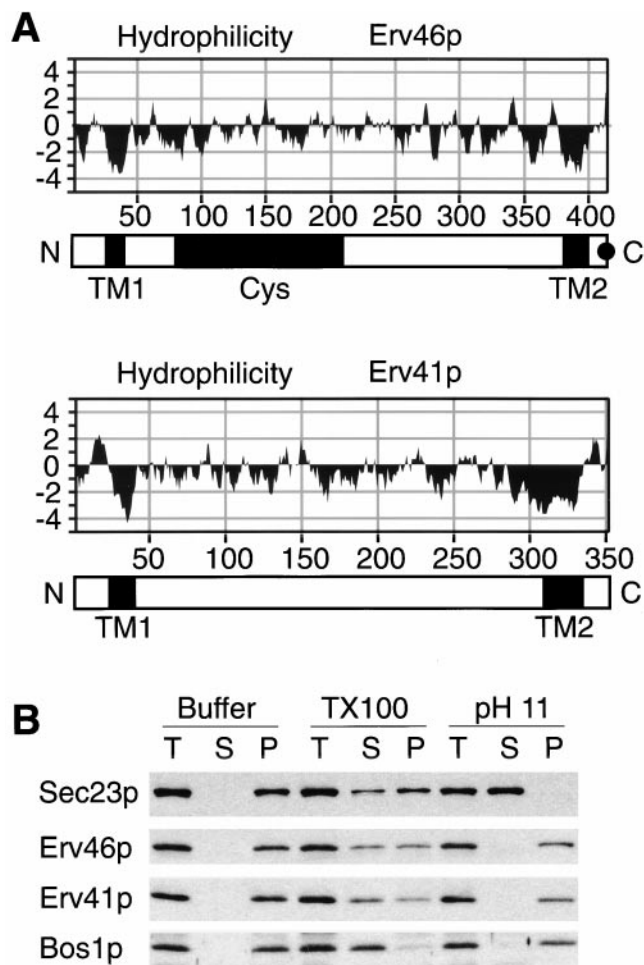


Figure 4. Erv46p and Erv41p are integral membrane proteins. (A) Hydrophilicity plots according to Kyte and Doolittle (1982) with a window size of 7. The schematic representations of the polypeptide chains indicate the positions of the putative transmembrane domains (TM1 and TM2) predicted by the HMMTOP program (Tusnady and Simon, 1998), the region containing conserved cysteine residues (Cys) and the KKXX motif (black dot). (B) Semiintact FY834 cells were treated with buffer (Materials and Methods), buffer containing 1% Triton X-100 (TX100), or 0.1 M Na_2CO_3 (pH 11) and centrifuged at 100,000 g. Totals before centrifugation (T), supernatant (S), and pellet (P) fractions were analyzed on a 12.5% polyacrylamide gel and immunoblotted for Sec23p (peripheral membrane protein), Erv46p, Erv41p, and Bos1p (integral membrane protein).

Analysis of the *erv41Δ* and *erv46Δ* Strains

To investigate the function of Erv41p and Erv46p, we analyzed strains bearing null alleles at the *ERV41* or *ERV46* loci. A PCR-based approach was used to direct the *HIS3* gene to the *ERV41* locus, replacing the entire *ERV41* ORF. A heterozygous *ERV41/erv41Δ* diploid strain (CBY763) was sporulated, and dissection of asci produced four viable spores. Tetrad analysis of these spores for the *HIS3* marker confirmed that *ERV41* was not essential for vegetative growth, although spores carrying the deletion were slightly delayed in their germination (data not shown). A haploid strain in which the *ERV46* reading frame had been replaced by a *KAN* marker (Winzeler et al., 1999) was crossed three times through the FY833/FY834 background to obtain the isogenic haploid *erv46Δ* strains CBY798 and CBY799.

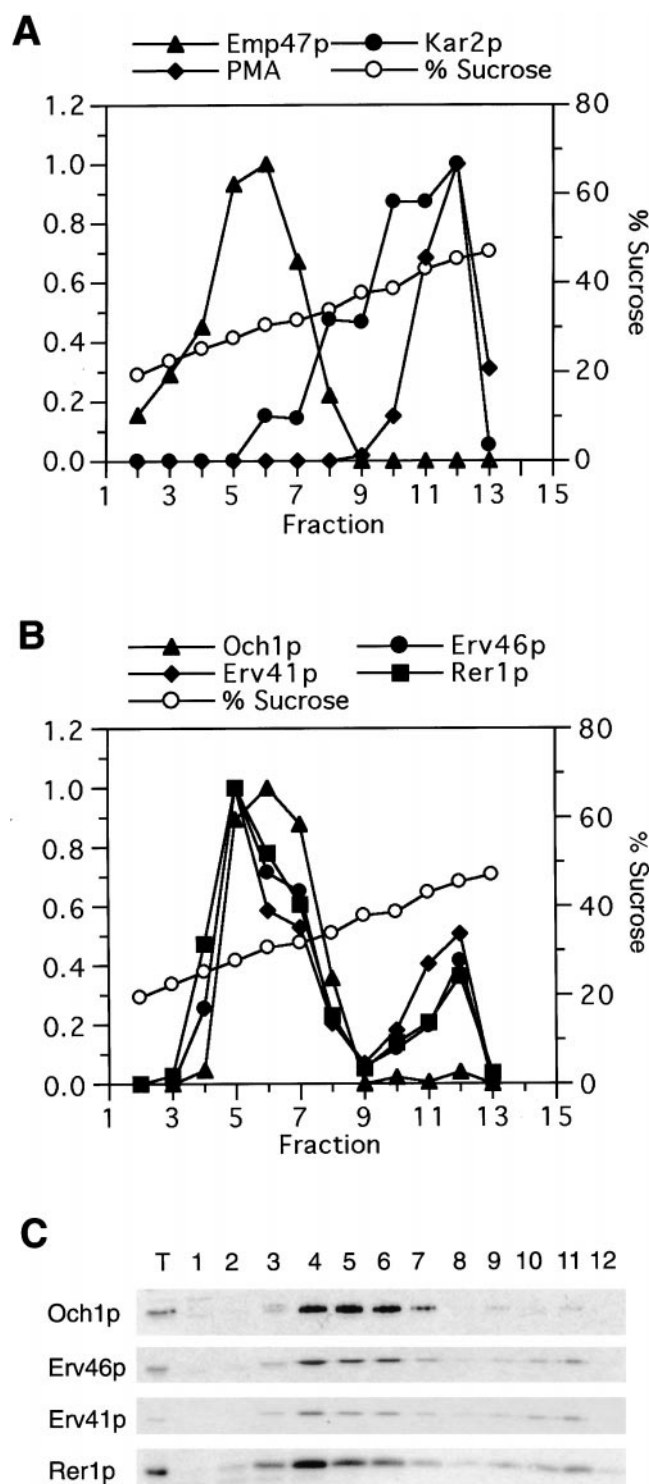


Figure 5. Sucrose gradient fractionation of Erv41p, Erv46p, Och1p, and Rer1p. An FY834 whole cell lysate was separated on an 18–60% sucrose density gradient, and fractions were collected, starting with fraction 1 at the top. (A) Relative levels of Emp47p (Golgi marker), Kar2p (ER marker), and plasma membrane marker (PMA) in each fraction were quantified by densitometry of immunoblots. (B) Relative levels of Och1p, Erv46p, Erv41p, and Rer1p as determined by densitometry of the immunoblots shown in C.

The haploid *erv41Δ* strains (CBY796 and CBY797) grew at rates comparable to wild-type strains at 30°C and 37°C but displayed a reduced growth rate at 16°C (Fig. 6 B). The cellular morphology as determined by light mi-

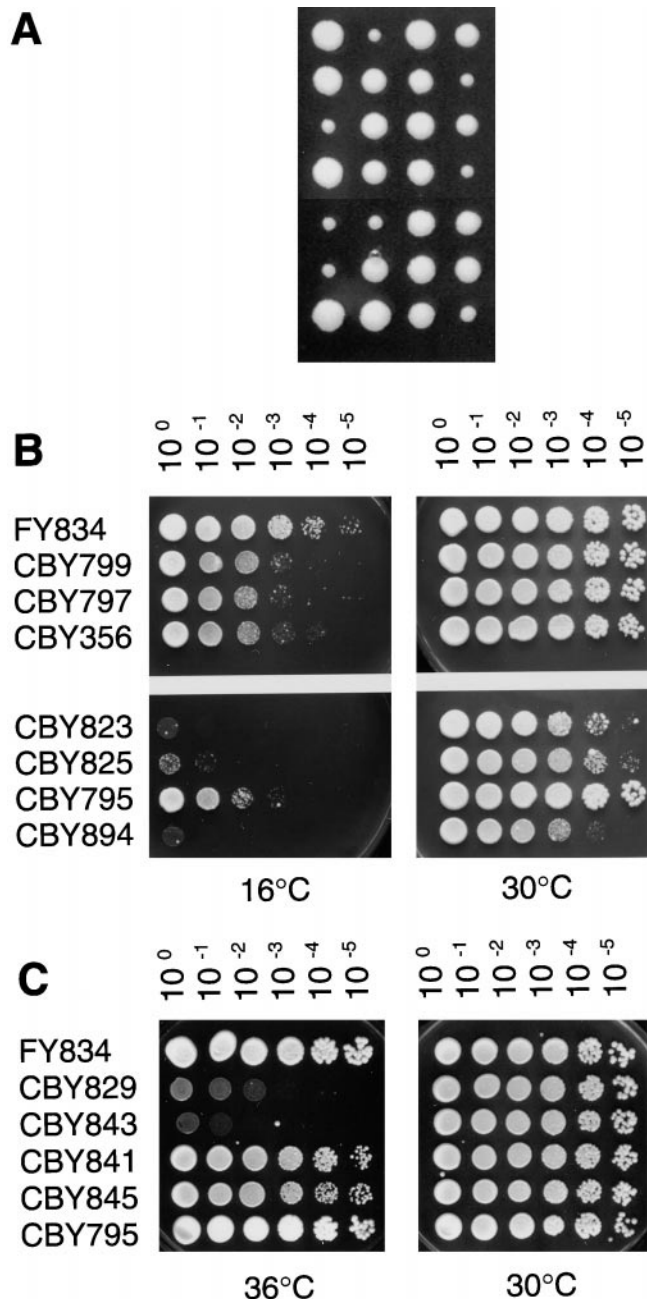


Figure 6. Genetic experiments with *erv41*Δ and *erv46*Δ strains. (A) An *erv14*Δ strain (CBY358) was mated with an *erv46*Δ strain (CBY799), and spores were dissected on a YPD plate. Spores that germinated and grew slower were shown to carry both deletions. (B) Cold sensitivity of *erv14*Δ, *erv41*Δ, and *erv46*Δ strains. Wild-type (FY834), *erv46*Δ (CBY799), *erv41*Δ (CBY797), *erv14*Δ (CBY356), *erv14*Δ *erv46*Δ (CBY823), *erv14*Δ *erv41*Δ (CBY825), *erv41*Δ *erv46*Δ (CBY795), and *erv14*Δ *erv41*Δ *erv46*Δ (CBY894) strains were grown to saturation in YPD, adjusted to an OD₆₀₀ of 3.0, and 5 μl of a 10-fold dilution series were spotted onto YPD plates. (C) Effects of *erv41*Δ and *erv46*Δ mutations on the *ypt1-3* mutation. Wild-type (FY834), *ypt1-3* (CBY829), *erv46*Δ *ypt1-3* (CBY843), *erv41*Δ *ypt1-3* (CBY841), *erv41*Δ *erv46*Δ *ypt1-3* (CBY845), and *erv41*Δ *erv46*Δ (CBY795) cells were spotted on YPD plates as in B.

croscopy as well as the mating and sporulation efficiencies of an *erv41*Δ strain were indistinguishable from the isogenic wild-type strains. Intracellular transport of the secretory proteins CPY and Gas1p was not detectably al-

tered in an *erv41*Δ strain and the major secretory proteins contained in culture supernatants were also unchanged (data not shown). Therefore, deletion of *ERV41* does not appear to interfere with general secretion. We performed a similar set of analyses on *erv46*Δ strains and observed phenotypes that were identical to *erv41*Δ strains, notably a reduced growth rate at 16°C. Furthermore, *erv41*Δ *erv46*Δ strains did not exhibit any exacerbated phenotypes compared with the single deletions strains (Fig. 6 B).

Next, we investigated possible influences of the *erv41*Δ and *erv46*Δ deletions on other mutations that impede transport through the early secretory pathway (Table IV and Fig. 6). First, we examined phenotypes combined with null alleles of other known *ERV* genes. An *erv14*Δ strain (CBY356) also exhibits a delayed rate of spore germination, a slightly slower growth rate in YPD at 30°C (increase in doubling time of ~18% over the wild type), and cold sensitivity. These phenotypes are significantly exacerbated by either *erv41*Δ or *erv46*Δ. In *erv14*Δ *erv41*Δ (CBY825) and *erv14*Δ *erv46*Δ (CBY823) strains, germination is delayed by several days (Fig. 6 A), the doubling time is increased by ~40% compared with the wild type, and cold sensitivity is increased. The triple *erv14*Δ *erv41*Δ *erv46*Δ (CBY894) mutant does not show any further exacerbation of these phenotypes except that its germination is extremely delayed by ~1 wk at room temperature. In contrast to the *erv14*Δ effects, combining *emp24*Δ with *erv41*Δ or *erv46*Δ did not alter growth phenotypes.

Also, we investigated the influences of *erv41*Δ and *erv46*Δ deletions on thermosensitive mutations in known budding, tethering, and fusion genes. An *erv41*Δ *ypt1-3* strain (CBY841) showed a significantly decreased thermosensitivity compared with *ypt1-3* alone (CBY829). The same effect was observed in a *ypt1-3* *erv41*Δ *erv46*Δ strain (CBY845), whereas *erv46*Δ alone slightly exacerbated the thermosensitivity of *ypt1-3* (CBY843, Fig. 6 C). In contrast, no effect was observed with the other tethering mutations *uso1-1* and *sec35-1*. The *erv41*Δ or *erv46*Δ null mutations did not influence any of the growth phenotypes associated with mutations in the COPI (*sec21-1*), COPII (*sec12-4*, *sec13-1*, *sec16-1*, and *sec23-1*), or fusion (*sed5-1*) proteins. Finally, we did not observe any synthetic effects when the *erv41*Δ or *erv46*Δ mutations were combined with *rer1*Δ or *erv29*Δ.

Effects of *erv41*Δ and *erv46*Δ on Transport to the Golgi

Since *erv41*Δ and *erv46*Δ strains did not exhibit major defects in secretion, but the *erv14*Δ *erv46*Δ strains showed an exacerbated growth phenotype, we examined the transport kinetics in this strain compared with wild-type and the single *erv14*Δ and *erv46*Δ strains. A pulse-chase experiment was performed in which cells were grown in minimal media and pulsed for 7 min with [³⁵S]methionine and [³⁵S]cysteine to label newly synthesized proteins. Excess unlabeled cysteine and methionine were added for the chase phase, and the maturation of CPY was followed by immunoprecipitation with an anti-CPY antibody. CPY first appears in the ER as the P1 precursor form of 67 kD and is then modified in the Golgi to yield its P2 form of 69 kD and finally processed in the vacuole to the mature M form of 61 kD (Stevens et al., 1984). As seen in Fig. 7, a slight delay in transport of CPY was observed in an *erv14*Δ strain (Powers and Barlowe, 1998), and, when combined with *erv46*Δ, no further decrease in the transport rate was detected.

Table IV. Effects of *erv41* and *erv46*

Strain number	Genotype	Phenotype*
FY834	wild type	—
CBY356	<i>erv14Δ</i>	Delayed germination, slightly slower growth, cold sensitive
CBY799	<i>erv46Δ</i>	Cold sensitive
CBY797	<i>erv41Δ</i>	Delayed germination, cold sensitive
CBY795	<i>erv41Δ erv46Δ</i>	No effect
CBY823	<i>erv14Δ erv46Δ</i>	Slow germination, slow growth, cold sensitive
CBY825	<i>erv14Δ erv41Δ</i>	Very slow germination, slow growth, cold sensitive
CBY894	<i>erv14Δ erv41Δ erv46Δ</i>	Extremely slow germination, slow growth, cold sensitive
CBY832	<i>erv41Δ erv46Δ emp24Δ</i>	No effect
CBY841	<i>erv41Δ ypt1-3</i>	Significant decrease in temperature sensitivity
CBY843	<i>erv46Δ ypt1-3</i>	Slight increase in temperature sensitivity
CBY845	<i>erv41Δ erv46Δ ypt1-3</i>	Significant decrease in temperature sensitivity
CBY912	<i>erv41Δ usol-1</i>	No effect
CBY910	<i>erv46Δ usol-1</i>	No effect
CBY914	<i>erv41Δ erv46Δ usol-1</i>	No effect
CBY943	<i>erv41Δ sec35-1</i>	No effect
CBY941	<i>erv46Δ sec35-1</i>	No effect
CBY945	<i>erv41Δ erv46Δ sec35-1</i>	No effect
CBY849	<i>erv41Δ sec21-1</i>	No effect
CBY851	<i>erv46Δ sec21-1</i>	No effect
CBY852	<i>erv41Δ erv46Δ sec21-1</i>	No effect
CBY853	<i>erv41Δ sed5-1</i>	No effect
CBY855	<i>erv46Δ sed5-1</i>	No effect
CBY857	<i>erv41Δ erv46Δ sed5-1</i>	No effect
CBY860	<i>erv41Δ sec12-1</i>	No effect
CBY862	<i>erv46Δ sec12-1</i>	No effect
CBY863	<i>erv41Δ erv46Δ sec12-1</i>	No effect
CBY923	<i>erv41Δ sec16-2</i>	No effect
CBY921	<i>erv46Δ sec16-2</i>	No effect
CBY924	<i>erv41Δ erv46Δ sec16-2</i>	No effect
CBY932	<i>erv41Δ sec13-1</i>	No effect
CBY933	<i>erv46Δ sec13-1</i>	No effect
CBY930	<i>erv41Δ erv46Δ sec13-1</i>	No effect
CBY917	<i>erv41Δ sec23-1</i>	No effect
CBY916	<i>erv46Δ sec23-1</i>	No effect
CBY919	<i>erv41Δ erv46Δ sec23-1</i>	No effect
CBY964	<i>erv14Δ rer1Δ</i>	No effect
CBY949	<i>erv41Δ rer1Δ</i>	No effect
CBY970	<i>erv29Δ erv14Δ</i>	No effect
CBY968	<i>erv29Δ erv41Δ</i>	No effect

*Conditions tested: YPD at 16°C, 25°C, 30°C, 36°C, and 38°C.

To further investigate a possible defect in ER to Golgi transport, we used an in vitro transport assay that permitted us to differentiate between the budding, tethering, and fusion stages (Barlowe, 1997; Cao et al., 1998). We have

found this assay to be a more sensitive method to measure transport between the ER and Golgi. In some instances, mutant strains that display normal transport kinetics in pulse–chase experiments show defects in distinct stages of

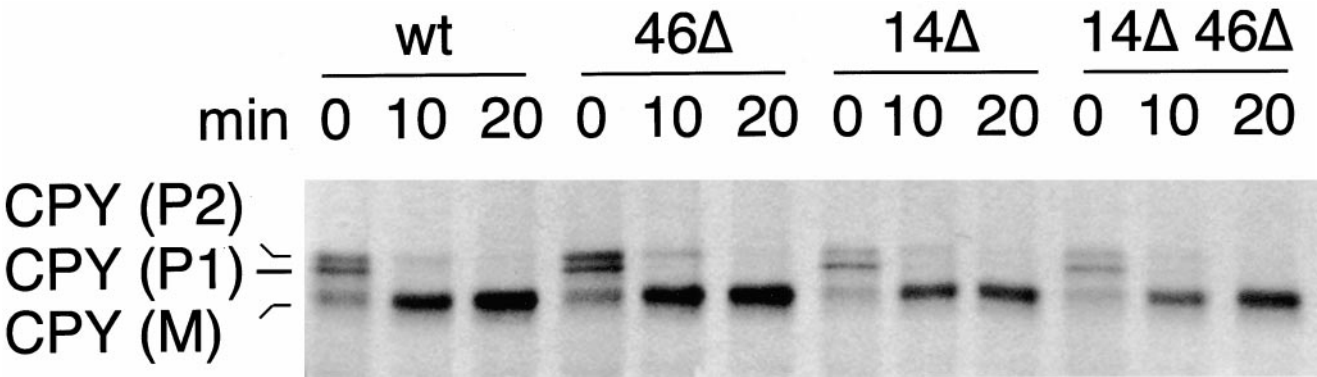


Figure 7. Pulse–chase analysis of CPY maturation in wild-type and *erv14Δ erv46Δ* deletion strains. Wild-type (wt, FY833), *erv46Δ* (46Δ, CBY798), *erv14Δ* (14Δ, CBY358), and *erv14Δ erv46Δ* (14Δ 46Δ, CBY822) strains were pulsed for 7 min with ³⁵S-labeled cysteine and methionine and then chased for 10 or 20 min. Labeled CPY was immunoprecipitated from cell extracts, resolved on a 10% polyacrylamide gel, and visualized by autoradiography.

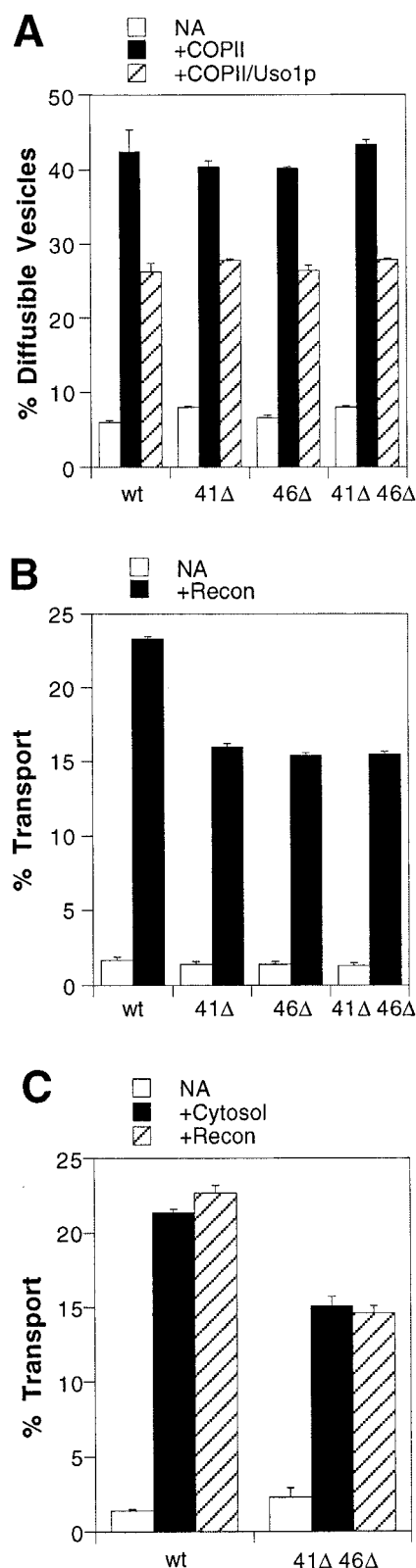


Figure 8. Influences of *erv41Δ* and *erv46Δ* mutations on ER-Golgi transport. (A) Vesicle budding and tethering in washed semiintact cells prepared from wild-type (wt, FY834), *erv41Δ* (41Δ, CBY797), *erv46Δ* (46Δ, CBY799), and *erv41Δ erv46Δ* (41Δ 46Δ, CBY795) strains. The levels of diffusible vesicles in reactions without reconstitution proteins (NA), with COPII proteins (+COPII), and with COPII proteins plus the tethering factor

cell-free transport (Conchon et al., 1999). Washed semiintact cells containing ^{35}S -labeled gp- α -F bud ^{35}S -labeled gp- α -F-containing vesicles in the presence of COPII proteins. Packaged ^{35}S -labeled gp- α -F in vesicles can be quantified by precipitation with concanavalin A-Sepharose, allowing us to assay budding efficiencies. The vesicle tethering stage may be monitored as the decrease in diffusible COPII vesicles upon addition of the tethering protein Uso1p. Lastly, reconstituted transport to the Golgi complex can be measured after addition of COPII, Uso1p, and LMA1 to semiintact cells by precipitation of Golgi-modified forms of ^{35}S -labeled gp- α -F with α 1,6-mannose-specific antiserum. As seen in Fig. 8 A, the budding and tethering stages of transport were not impaired in mutant strains compared with a wild-type strain. However, there was a modest but significant decrease in transport to the Golgi complex in the *erv41Δ*, *erv46Δ*, and the double *erv41Δ erv46Δ* membranes (Fig. 8 B). Notably, the effects of the single deletions were neither additive nor cooperative when combined in the double mutant strain. Together, these results suggest that the transport defect occurred during the fusion stage of this assay. Also, we examined transport efficiencies in the presence of a crude cytosol to determine if the *erv41Δ erv46Δ* membranes required additional factors not provided by purified reconstitution proteins. As shown in Fig. 8 C, a similar transport defect was observed for reactions using crude cytosol or purified proteins to drive transport.

Association of *Erv41p* and *Erv46p*

Based on the colocalization of *Erv41p* and *Erv46p*, the similarity in phenotypes displayed by *erv41Δ* and *erv46Δ* in our genetic experiments and the nonadditive effects of the *erv41Δ* and *erv46Δ* mutations on ER-Golgi transport assays, we suspected that these proteins act together and possibly form a complex. Therefore, we investigated whether the expression of *Erv41p* and *Erv46p* is interdependent by immunoblotting cells containing the *erv41Δ*, *erv46Δ*, and *erv41Δ erv46Δ* alleles with antisera against *Erv41p* and *Erv46p*. The *erv46Δ* strain did not express a detectable level of *Erv41p*, and the *erv41Δ* strain exhibited a reduced expression of *Erv46p* compared with the wild type (Fig. 9). Consistent with these findings, we observed that *Erv46p* could be expressed at higher levels from a 2 μ plasmid than *Erv41p* (Fig. 3). A strain carrying both *ERV41* and *ERV46* on 2 μ plasmids did not express either of them at levels higher than a strain just overexpressing *Erv46p* (not shown). Also, the expression level of endogenous *Erv41p* could be elevated (approximately twofold) if *Erv46p* was overproduced under control of the *GAL1* promoter (compare the total [T] lanes in Fig. 10). These re-

Usolp (+COPII/Uso1p) are indicated. (B) Overall transport of ^{35}S -labeled gp- α -F to the Golgi complex in the same strains. Semiintact cells were incubated alone (NA) or with the reconstitution proteins COPII, Uso1p, and LMA1 (+Recon). (C) Overall transport of ^{35}S -labeled gp- α -F factor to the Golgi complex in wild-type (wt, FY834) and *erv41Δ erv46Δ* (41Δ 46Δ, CBY795) strains. Semiintact cells were incubated alone (NA), with cytosol, or with the reconstitution proteins COPII, Uso1p, and LMA1 (+Recon).

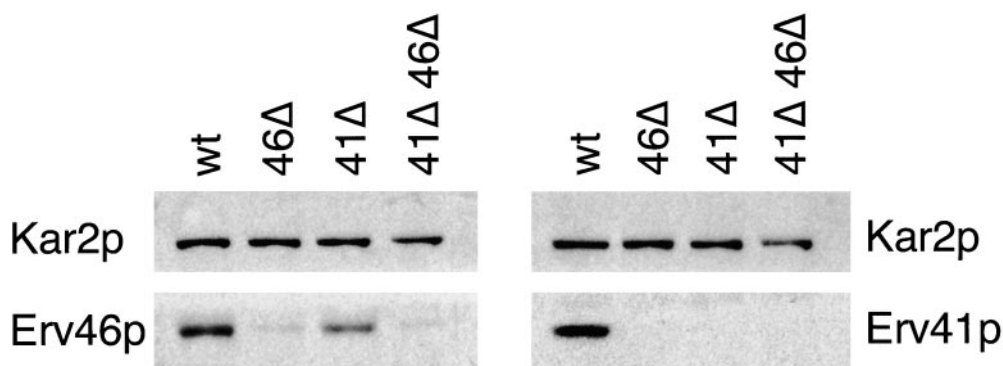


Figure 9. Erv41p and Erv46p depend on each other for their expression. wt, wild-type strain FY834; 46Δ, *erv46*Δ deletion strain CBY799; 41Δ, *erv41*Δ deletion strain CBY797; and 41Δ 46Δ, *erv41*Δ *erv46*Δ double deletion strain CBY795. Semi-intact cells were resolved on a 12.5% polyacrylamide gel and immunoblotted with polyclonal anti-Erv41p and anti-Erv46p antisera. Kar2p is shown as a loading control.

sults indicate that the expression of Erv41p is highly dependent on the presence of Erv46p, and to a lesser extent Erv46p expression depends on the presence of Erv41p. One interpretation of these results is that Erv41p and Erv46p are associated in a multisubunit protein complex, and when one of the members of this complex is absent, the other subunit(s) is destabilized.

To determine if Erv41p and Erv46p are physically associated, we performed a native immunoprecipitation experiment from detergent-solubilized membranes. Microsomes from a wild-type strain and from a strain expressing 3HA-Erv46p were solubilized with Triton X-100, and tagged Erv46p was precipitated by the addition of anti-HA antibodies. As seen in Fig. 10, Erv41p coprecipitated with 3HA-Erv46p in the tagged strain, whereas no Erv41p was precipitated from the untagged wild-type microsomes. As controls, neither Sec23p nor Erv25p were precipitated, indicating that the association of 3HA-Erv46p with Erv41p was specific. In other experiments, we found that HA-tagged Erv41p could be coimmunoprecipitated with Erv46p when Erv46p specific polyclonal antibodies were used (not shown). Finally, we observed that endogenous Erv46p was not detected in complexes that were isolated with 3HA-Erv46p (Fig. 10; Otte, S., unpublished observation). Collectively, the expression studies and immunoprecipitation results indicate that Erv41p and Erv46p are physically associated in a heteromeric complex and that these complexes probably contain single Erv46p and Erv41p subunits.

Discussion

Using a reconstituted budding assay combined with mass spectrometry, we have identified four known transmembrane protein constituents of COPII-coated vesicles (Erv14p, Emp24p, Erv25p, and Bet1p), and have localized six other characterized proteins (Yip3p, Rer1p, Erp1p, Erp2p, Yif1p, and Emp47p) and three additional novel proteins (Erv29p, Erv41p, and Erv46p) to these carrier vesicles. We went on to characterize Erv41p and Erv46p, two related proteins that are conserved across species. Both are integral transmembrane proteins and are predicted to have large luminal domains with shorter COOH- and NH₂-terminal cytoplasmic segments. Localization studies show that both proteins reside in the Golgi and ER at steady state. *erv41*Δ and *erv46*Δ strains are cold-sensitive, and these deletions exacerbate the *erv14*Δ

phenotype. The *erv41*Δ and *erv46*Δ mutations also influence the thermosensitivity of a *ypt1-3* strain. Strains carrying the *erv41*Δ and *erv46*Δ null alleles show normal vesicle budding and tethering but a reduced overall transport efficiency between the ER and Golgi, suggesting a defect downstream of the tethering stage. Expression of Erv41p and Erv46p is interdependent, and both proteins could be coimmunoprecipitated, indicating that these proteins are physically associated.

In discussing potential functions for Erv41p and Erv46p, it may be informative to consider other characterized vesicle proteins. We have now identified four p24 proteins on isolated ER-derived vesicles. Yeast has eight members of this family, Erv25p (Belden and Barlowe, 1996), Emp24p (Schimmöller et al., 1995), and Erp1p to Erp6p (Marzioch et al., 1999). These abundant, conserved, integral type I transmembrane proteins have been found in COPI and COPII vesicles and have been shown to shuttle between the ER and Golgi. Their cytosolic tails have a high affinity for coat proteins and have been proposed to act as a scaffold during the formation of the protein coat (Bremser et al., 1999), as transport receptors for secretory cargo (Schimmöller et al., 1995) or as negative regulators of vesicle budding that influence cargo sorting (Elrod-Erickson and Kaiser, 1996). However, a strain that lacks all eight p24 proteins is viable and did not show defects in overall COPI- or COPII-mediated transport (Kaiser, 2000; Springer et al., 2000), indicating that they do not perform an essential role in transport through the early secretory pathway in yeast. The phenotypes associated with p24 deletion strains are consistent with a role in protein and/or lipid sorting during vesicle budding either by association with cargo or through formation of specialized sorting regions or membrane compartments. The expression levels of the Erv25p, Emp24p, Erp1p, and Erp2p are interdependent, these proteins are found associated in a heterooligomeric complex, and strains lacking any one subunit of this complex exhibit similar phenotypes (Marzioch et al., 1999). Our detection of the Erp1p and Erp2p proteins on COPII vesicles is consistent with these observations, and we are testing the possibility that this heterooligomeric p24 complex is packaged en bloc during vesicle formation.

Yip3p and Yif1p, two previously identified proteins, were also encountered on ER-derived vesicles. The Yip proteins were initially discovered as Ypt1p interacting proteins (Yang et al., 1998) and Yif1p as a Yip1p interact-

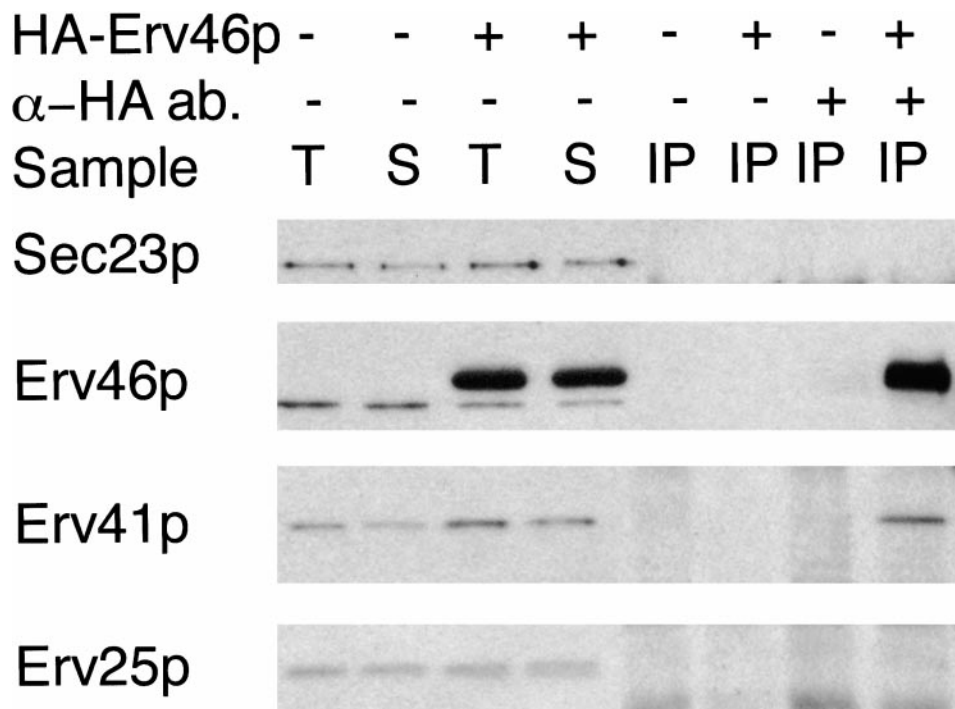


Figure 10. Erv41p is coimmunoprecipitated with HA-tagged Erv46p. Microsomes were prepared from FY833 (– HA-Erv46p) and CBY767 (+ HA-Erv46p) cells grown in YP with 2% galactose and 0.2% glucose and solubilized with buffer containing 2% Triton X-100 (T). The soluble extracts (S) were incubated in absence (– anti-HA ab) or presence (+ anti-HA ab) of the monoclonal anti-HA antibody. Precipitates were resolved on a 12.5% polyacrylamide gel and immunoblotted for Sec23p, Erv46p, Erv41p, and Erv25p.

ing factor (Andrulis et al., 1998). Ypt1p is a small GTPase required for vesicle transport through the early secretory pathway (Rexach and Schekman, 1991; Segev et al., 1988) and therefore the Yip and Yif proteins are thought to operate in conjunction with GTPases to catalyze vesicle fusion. Yip1p and Yif1p are integral membrane proteins that localize predominantly to Golgi membranes and possess hydrophilic NH₂-terminal domains facing the cytosol. Recently, a Yip1p–Yif1p complex has been proposed to function as a receptor in recruiting GTPases to specific membranes, perhaps acting as a GDI displacement factor (Yang et al., 1998; Matern et al., 2000). We find the Yip1p–Yif1p complex is efficiently packaged into COPII vesicles, suggesting that these proteins actively cycle between the ER and Golgi compartments. It seems possible that the Yip1p–Yif1p complex on vesicles could act directly to target vesicles to Ypt1p on the surface of Golgi membranes or Yip1p–Yif1p could perform a more general role in recruiting GTPases to several distinct intracellular membranes. Further studies will be needed to distinguish between these possibilities.

Rer1p is required for the ER localization of type II transmembrane proteins (e.g. Sec12p) and has been proposed to couple retrieved proteins to the COPI coat during retrograde Golgi to ER transport (Sato et al., 1995; Boehm et al., 1997; Sato et al., 1997). We have found that Rer1p is included in COPII vesicles with a high efficiency, suggesting this protein actively cycles between the ER and Golgi compartments. Our observation seems consistent with the proposed function of Rer1p in retrieval of ER residents from post-ER compartments. However, it remains to be determined if the COPII-dependent forward transport of Rer1p reflects recycling necessary to perform its function in retrieval of ER residents or whether this protein performs additional roles in anterograde transport.

We had expected that the abundant Erv proteins would

represent transport machinery involved in vesicle formation, cargo selection, and membrane fusion (Belden and Barlowe, 1996). To a large extent, our current study bears this out. However, we also found that the outer-chain mannosyltransferase (Och1p) was packaged into COPII vesicles as has been previously reported (Bednarek et al., 1995). Och1p was not detected in our mass spectral analysis, but antibodies against this protein were generated to allow for a comparison of packaging efficiencies with other Erv proteins. We find that Och1p is packaged into COPII vesicles, albeit at a lower efficiency than most other Erv proteins (Fig. 2), and is localized predominantly to Golgi membranes (Fig. 5). The packaged Och1p could represent newly synthesized protein in transit to the Golgi or may belong to a class of Golgi localized proteins that cycle between the ER and Golgi compartments at a significant rate as has been observed for mammalian galactosyltransferase (Zaal et al., 1999). It is interesting to note that other Golgi localized proteins such as Ypt1p and GDPase were not efficiently packaged into COPII vesicles (Cao and Barlowe, 2000). Furthermore, *in vivo* studies indicate that some Golgi-localized proteins cycle rapidly through the ER, whereas others do not (Wooding and Pelham, 1998; Barrowman et al., 2000). It remains to be determined if cycling rates necessarily reflect an important functional property. Regardless, the fact that some proteins typically thought to be Golgi residents are found in COPII vesicles suggests that uncharacterized Erv proteins could potentially belong to this group.

Erv41p and Erv46p are predicted to have large luminal domains and short NH₂- and COOH-terminal cytosolic tails that potentially interact with the COPII and/or COPI coats. Based on these and other considerations discussed above, we can envisage a few possible roles for the Erv41p–Erv46p complex that are consistent with our experimental observations. First, this complex could perform

a role similar to the p24 complex in sorting during vesicle formation. The complex could fulfill this role through direct interaction with cargo molecules or by establishing domains on the surface of budding membranes. In some respects, the heteromeric arrangement, the membrane topology, and the nonessential phenotypes exhibited by the Erv41p–Erv46p complex are reminiscent of the p24 deletion strains. We have not detected any specific secretory cargo that accumulates in the *erv41Δ* and/or *erv46Δ* deletion strains, however, we currently have a very limited capability of monitoring individual secretory cargo. If the Erv41p–Erv46p complex is required for the efficient transport of a small subset of nonessential cargo proteins, we may not easily detect an accumulation. Second, the Erv41p–Erv46p complex could operate in the retention and/or retrieval of transport machinery to the early secretory pathway. For example, as Rer1p acts in the retrieval of Sec12p and other ER residents, the Erv41p–Erv46p complex could operate in localizing proteins to Golgi membranes. In keeping with this idea, the *erv41Δ erv46Δ* strain displayed a modest defect in the fusion stage of in vitro transport between the ER and Golgi, suggesting the Erv41p–Erv46p complex could interact with or serve to correctly localize the membrane fusion machinery (e.g., SNAREs and SNARE regulatory proteins). Third, the Erv41p–Erv46p complex may not act in movement or localization of proteins, but could be involved in lipid transport. A majority of cellular phospholipid, glycolipid, and sterol synthesis occurs in the ER (for review see Daum et al., 1998), and it seems probable that specific proteins act to sort and transport these species to their proper location. Although nonsecretory routes exist for lipid transport, much of this lipid transport occurs through the classical secretory pathway and presumably through COPII vesicles. Therefore, the observed in vitro defect during the vesicle fusion stage could be caused by a suboptimal lipid composition of vesicle or acceptor membrane bilayers, leading to a reduction in membrane fusion efficiency. Finally, the Erv41p–Erv46p complex could perform a role in the posttranslational maturation of secretory proteins such as protein folding or glycosylation in the early secretory pathway. If this last possibility were the case, we again would speculate that the effect is on a subset of secretory cargo because we do not detect any general defects in folding, no extracellular secretion of kar2p (Otte, S., unpublished observation), and no obvious alterations in N- or O-linked oligosaccharide modifications.

Several recent reports suggest a role for COPII-dependent transport in regulation of cellular homeostasis (Niwa et al., 1999; Nohturfft et al., 2000; Travers et al., 2000). Although the underlying mechanisms of these regulatory pathways remain to be elucidated, our investigation of Erv proteins may shed light on these questions. Notably, several of the proteins we have identified on COPII vesicles, including the novel proteins Erv29p, Erv41p, and Erv46p, are induced upon activation of the unfolded protein response pathway (Travers et al., 2000). With respect to the Erv41p–Erv46p complex, it will be important to determine binding partners that potentially include additional members of the heteromeric Erv41p–Erv46p complex, vesicle coat proteins, and/or specific cargo molecules. We will combine these biochemical approaches with cell-free

transport assays and molecular genetic analyses of Erv41p–Erv46p to elucidate their function.

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