Yeast Kex1p Is a Golgi-associated Membrane Protein: Deletions in a Cytoplasmic Targeting Domain Result in Mislocalization to the Vacuolar Membrane

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Abstract. We have investigated the localization of Kex1p, a type I transmembrane carboxypeptidase involved in precursor processing within the yeast secretory pathway. Indirect immunofluorescence demonstrated the presence of Kex1p in a punctate organelle resembling the yeast Golgi apparatus as identified by Kex2p and Sec7p (Franzusoff, A., K. Redding, J. Crosby, R. S. Fuller, and R. Schekman. 1991. J. Cell Biol. 112:27-37). Glycosylation studies of Kex1p were consistent with a Golgi location, as Kex1p was progressively N-glycosylated in an MNNl-dependent manner.

To address the basis of Kex1p targeting to the Golgi apparatus, we examined the cellular location of a series of carboxy-terminal truncations of the protein. The results indicate that a cytoplasmically exposed carboxy-terminal domain is required for retention of this membrane protein within the Golgi apparatus. Deletions of the retention region or overproduction of wild-type Kex1p led to mislocalization of Kex1p to the vacuolar membrane. This unexpected finding is discussed in terms of models involving either the vacuole as a default destination for membrane proteins, or by endocytosis to the vacuole following their default localization to the plasma membrane.

Secre ted biologically active proteins and peptides are classically produced as precursors which undergo both endo- and exoproteolytic processing to release the mature species while traversing the secretory pathway. The KEX1 gene product (Kex1p) is a carboxypeptidase, specific for basic amino acid residues, and is responsible for processing proteins secreted by Saccharomyces cerevisiae (Korfhagen et al., 1987; Cooper and Bussey, 1989). In conjunction with the proteases Kex2p and Dipeptidyl aminopeptidase A (DPA A)1 (Ste13p), Kex1p proteolytically matures proteins such as α-factor and K1 killer toxin from their precursors (for reviews see Bussey, 1988; Fuller et al., 1988).

Proteins that enter the secretory pathway are thought to be transported to the cell surface by default via a “bulk flow” mechanism unless they contain additional targeting information (Pfeffer and Rothman, 1987; Rothman, 1987; Wieland et al., 1987; Karrenbauer et al., 1990). Such targeting information is found in soluble proteins resident in the ER which maintain their localization by containing a retention signal at their carboxy termini (Munro and Pelham, 1987; Pelham et al., 1988). Deletion of the retention signal results in secretion of the soluble ER resident proteins to the cell surface. Soluble proteins destined for the mammalian lysosome receive a mannose-6-phosphate signal which targets them to the lysosome in a receptor-mediated manner. The absence of the mannose-6-phosphate signal from these soluble proteins results in their secretion (for review see Kornfield and Mellman, 1989). Secretion also occurs when the targeting signals for yeast-soluble proteins destined for the lysosome-like vacuole are altered (Vail et al., 1987; Johnson et al., 1987; Klionsky et al., 1988).

In addition to soluble proteins, the cell surface appears to be the default destination for mammalian ER, Golgi, and lysosomal membrane proteins as removal of their respective retention/targeting signals results in their delivery to the plasma membrane (Machamer and Rose, 1987; Jackson et al., 1990; Williams and Fukuda, 1990). As yet, no targeting or retention signals have been definitively assigned to yeast Golgi or vacuolar membrane proteins, nor is it known where these proteins are delivered upon perturbation of their targeting signals.

Kex1p is predicted to be a type I transmembrane protein with a large amino-terminal protease domain in the lumen of the secretory pathway, a single membrane-spanning domain, and a smaller carboxy-terminal domain positioned cytoplasmically. The observation that KEX1 cells intracellularly retain Kex1p activity prompted an analysis to determine in which secretory compartment Kex1p resided, and how it achieved such retention. Kex1p was found to be localized to the yeast Golgi apparatus, with retention mediated via the cytoplasmic domain of the protein. Removal of this domain...
or overproduction of wild-type Kexlp resulted in delivery of the protein to the vacuole rather than to the cell surface. This surprising result raises the possibility that the vacuole is the default destination for yeast Golgi membrane proteins.

Materials and Methods

Strains, Growth Media, and Procedures

Escherichia coli strains and associated DNA manipulations were as described previously (Cooper and Bussey, 1989).

Saccharomyces cerevisiae strains cSAB1386 (S86), Sc25k, and Sc25k-13 (Sc25k -4lex-Δ2) have been described previously (Cooper and Bussey, 1989). Other strains used were S6 (a strain sensitive to Kt killer toxin), TC106a (MATa leu2 his1 trp1 ura3 [K11-K1]), SEY5016a (sec1-1 leu3 ura3 gal10), XCY42-3D0 (MATa ade2-101 ade X [a putative second ade mutation that results in white colonies]), and 136A5 (MATa ade2-101 ade X [a putative second ade mutation that results in white colonies]).

Restriction endonucleases, T4 DNA polymerase, T4 DNA ligase, and Klenow fragment were purchased from either Bethesda Research Laboratories, (Gaithersburg, MD) or New England Biolabs (Beverly, MA), and were used as recommended by the suppliers. Unless otherwise stated, reagents used in these experiments were obtained from Sigma Chem. Co., St. Louis, MO.

Disruption of the KEX1 Locus

Disruption of the KEX1 locus with the URA3-based construct pl6 resulted in the deletion of the allele kexl-Δ2. An alternative disruption using the LEU2-containing plasmid pl7 produced the allele kexl-Δ3. Pl7 was constructed by digesting pl6 (Dmochowska et al., 1987) with EcoRV to remove the URA3 gene and into this blunt-ended fragment was ligated the NheI nonsense codon linker 5'-CTAGCTAGCTAG 3'. To aid in identifying mutant clones an HpaI site was introduced by the mutagenesis. Introduction of the various KEX1 mutations into pVT103-L produced the following plasmids: pKX1-18AS, pKX1-18Hpa, pKX1-18Bel, and pKX1-18Hinc.

A 3.1-kb HindIII fragment containing the KEX1 gene was inserted into YCp50 to create pKX1-20AS, pKX1-20AMS, pKX1-20Hpa, and pKX1-20Hinc. pAD81 (Dmochowska et al., 1987), which contained the KEX1 gene, was digested with XhoI, the site was filled in with Klenow enzyme, and into this blunt-ended fragment was ligated the Nhel nonsense codon linker 5'-CTAGCTAGCTAG 3'.

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The diploids were then transformed into HAB595 (mnn9 ura3) to create the plasmid YCP50 to create pKX1-20Hpa.

Site-directed Mutagenesis of KEX1

Mutagenesis of the carboxy-terminal region of KEX1 was performed on the same single-stranded DNA template and in the same manner as described (Cooper and Bussey, 1989). All of the mutations were confirmed by DNA sequencing (Sequenase, United States Biochem. Corp., Cleveland, OH).

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Immunoprecipitations and Carbonate Extractions

Antiserum production and procedures for sodium carbonate extractions, radio-labeling cells, and associated immunoprecipitations have been described previously (Cooper and Bussey, 1 Affinity Purification of Kexlp Antibodies and Indirect Immunofluorescence Studies

β-Galactosidase-Kexlp fusion protein (Cooper and Bussey, 1989) was isolated by preparative SDS-PAGE; electroelution into 0.1 M NH4CO3, 0.1% SDS; lyophilised; resuspended in coupling buffer (0.5 M NaCl, 0.1 M NaHCO3, pH 9.0); and then dialyzed against coupling buffer + 0.25% SDS + 80 mg ml-1 PMSF. The extract was then coupled to 2 g of activated CNBr-Sepharose 4B (Pharmacia LKB Biotechnology Inc., Piscataway, NJ) overnight at room temperature. The coupling efficiency was judged to be > 80% as determined by comparing unbound protein with the starting material. The Sepharose was incubated with blocking buffer (1 M ethanolamine, 0.1 M NaHCO3, pH 8.0, 0.5 M NaCl) for 4 h at room temperature. The resulting column (bed volume, 2 ml) was then treated with successive washes of blocking buffer and acetate buffer (0.1 M sodium acetate, pH 4.0, 0.5 M NaCl), and finally with PBS. A similar column (bed volume, 2 ml) was made
with an extract made from the granules of E. coli cells producing β-galactosidase.

Both the β-galactosidase and β-galactosidase-Kexlp columns were washed successively with buffer A (50 mM Hepes, pH 7.5, 150 mM NaCl, 1 mM EDTA), buffer B (buffer A + 1 M guanidine-HCl), and buffer C (50 mM Hepes, pH 7.5, 15 mM NaCl, 4.5 M MgCl₂). The two columns were then arranged so that the eluate from the β-galactosidase column flowed into the β-galactosidase-Kexlp column. All buffers used from this point on contained in addition: 1 mg/ml BSA, 0.2% NaN₃, 1 mM PMSF. Columns were equilibrated with buffer A; Kexlp antiserum was loaded onto the linked columns and was circulated for 3 h at 4°C. The columns were disconnected and the β-galactosidase-Kexlp column was washed first with buffer B and then with buffer A. Antibodies were then eluted from this column with buffer C; fractions were collected and immediately dialyzed against buffer A at 4°C. Eluted fractions were tested for the presence of antibodies with a double Western approach and the positive fractions tested for their ability to immunoprecipitate [³⁵S]methionine-labeled Kexlp from yeast. These fractions were concentrated by dialysis against buffer A containing 15% glycerol, and stored at −70°C.

Immunofluorescence studies were performed following the procedure of Redding et al. (1991) and used either affinity-purified anti-Kexlp antibodies or a mouse monoclonal (3D11) which is directed against the 60-kD subunit of the yeast vacuolar membrane H⁺-ATPase (Kane et al., 1992). The cells were observed using an Axiopt microscope (Carl Zeiss, Inc., Thornwood, NY) (equipped for epifluorescence at excitation wavelengths appropriate for the described fluoros) with a 100X objective and photographed with T-Max 400 film (Eastman Kodak Co., Rochester, NY). Mitochondria and nuclei were identified by 4,6-diamidino-2-phenyl-indole staining while vacuoles were identified by differential interference contrast (Nomarski) optics.

Results

Kexlp Is Glycosylated within the Golgi Apparatus

Upon translocation into the yeast ER, proteins that are destined for Asn-linked glycosylation receive a core oligosaccharide which can be elaborated in the Golgi apparatus by the addition of further mannose residues to produce the “outer chain.” The outer chain consists of a backbone of α(1→6) linked mannose residues to which are attached mannose sidechains in α(1→2) linkages which then terminate in α(1→3) linkages (Fig. 1; Ballou, 1982). The extension of the core is heterogenous in nature with different proteins receiving varying degrees of elaboration.

Kexlp is a glycoprotein which receives Asn-linked glycosylation in a multistep process (Cooper and Bussey, 1989). The initial event occurs in the ER where core oligosaccharides are attached to the protein. The observed difference in molecular mass of ~3–4 kDa between Kexlp labeled for 10 min and that of Kexlp produced in tunicamycin-treated cells suggests that two of the three predicted lumenal Asn acceptor residues are glycosylated. The Asn-linked core oligosaccharides undergo a modification in a post-ER compartment that increases the mass of Kexlp by 1–2 kDa, demonstrating that Kexlp does not receive an extensive outer chain (Cooper and Bussey, 1989). Such a modification was reduced in a sec1 mutant which blocks intra-Golgi transport, and was unaffected in a sec1 mutant that prevents secretory vesicle fusion with the plasma membrane (Novick et al., 1981; Cooper and Bussey, 1989).

The yeast Golgi apparatus has been functionally subdivided into several compartments on the basis of Asn-linked oligosaccharide addition (Frantzusoff and Schekman, 1989; Graham and Emr, 1991). Glycoproteins receive mannose outer chain modifications while traversing these Golgi compartments. Identification of the enzymes responsible for the post-ER modification(s) may indicate to which Golgi compartments Kexlp has been exposed, and thereby localize Kexlp within the secretory pathway. The basis of the post-ER progressive carbohydrate modification of Kexlp was analyzed using strains with mutations at various mnn loci. Such mutant strains show defects in outer chain glycosyl elaboration resulting in various truncations of the outer chain (Ballou, 1982; Kukuruzinska et al., 1987; Fig. 1). Several mnn strains were radiolabeled and the degree of modification of immunoprecipitated Kexlp was examined. Kexlp produced in mnn9 cells received most, if not all, of the wild-type glycosyl modification (Fig. 2, lanes 5 and 6) suggesting that most of the carbohydrate addition to Kexlp was not in the elaboration of the outer chain as mnn9 blocks initiation of outer chain synthesis (Ballou, 1982; Kukuruzinska et al., 1987). A likely candidate for the mnn9-independent modification was the Golgi-localized α(1→3) mannosyltransferase which has been shown to be responsible for α(1→3) mannoside addition to both the core and outer chains (Fig. 1; Nakajima and Ballou, 1975; Frantzusoff and Schekman, 1989; Graham and Emr, 1991). This enzyme activity has been found to be deficient in an mnn1 strain, and Kexlp synthesized in a strain disrupted at the mnn1 locus received only a small modification of its core oligosaccharide (Fig. 2, lanes 3 and 4). The α(1→3) mannosyltransferase is therefore responsible for producing the majority of the post-ER glycosylation modification to Kexlp.

Carboxy-terminal Truncations of Kexlp

The modification of Kexlp by the Golgi α(1→3) mannosyltransferase suggested that Kexlp reached the Golgi compartment containing the mannosyltransferase, yet little Kexlp-dependent activity was detected at the cell surface (see below) and indicated that Kexlp was retained within the secretory pathway.

An analysis of mutant forms of Kexlp was undertaken to examine which domain(s) of Kexlp was responsible for its function.
retention. The hydrophobic domain near the carboxy terminus of Kexlp was previously shown to be responsible for conferring membrane association, as deletion of this 17-amino acid residue region resulted in a soluble protein, Kexlp-AMS (Cooper and Bussey, 1989). Cells expressing this protein showed a significant increase in Kexlp-dependent activity in the cell medium (see below). That Kexlp-AMS was secreted rather than retained suggests that one of several domains may be responsible for retention of Kexlp: (1) the membrane-spanning domain which was absent in Kexlp-AMS, (2) the carboxy-terminal domain which was no longer exposed in the cytoplasm in Kexlp-AMS, and (3) a domain amino-terminal to the membrane-spanning domain which is perturbed in Kexlp-AMS when the cytoplasmic domain was juxtaposed adjacent to it.

A number of carboxy-terminal truncations of Kexlp were produced to test the above possibilities. The construction of these mutations via site-specific mutagenesis or nonsense linker insertion is described in Materials and Methods and the predicted proteins are shown in Fig. 3. Immunoprecipitations of the mutant Kexlp proteins were performed to confirm that they were correctly synthesized by the cell (Fig. 4). All of the truncated proteins were smaller than Kexlp by the predicted amount and entered the secretory pathway as shown by the addition of Asn-linked glycosylation (data not shown). A concern was that the mutants designed to remain membrane associated may not attain a stable type I orientation and remain in the ER. The most severely truncated mutant in this class, Kexlp-Hpa, lacks the entire endogenous cytoplasmic domain; it was further analyzed and shown to receive Asn-linked glycosylation which was further modified within the Golgi apparatus (Fig. 4). Previous sodium carbonate extractions of whole cell extracts demonstrated that Kexlp fractionated with the membrane containing pellet fraction whereas the soluble protein, Kexlp-AMS, fractionated with the supernatant (Cooper and Bussey, 1989). Carbonate treatment of whole cell extracts showed that Kexlp-Hpa fractionated with the membrane pellet (Fig. 4, lanes 5 and 6), consistent with it being an integral membrane protein. In addition, we have shown that CPY, a soluble Kexlp-like protein, is fully soluble under these extraction conditions (data not shown).

**Soluble Forms of Kexlp Are Secreted to the Cell Surface**

Kexlp activity was assayed in whole cells and their growth media to detect any potential mislocalization of the truncated forms of Kexlp to the cell surface. To compare mutants, it was important that the KEXI gene copy number be constant. The Kexlp truncation constructs were, therefore, inserted into the vector YCp50, a centromeric-based plasmid which is normally maintained at a single copy per cell (Rose et al.,...
generations before assay. YEPD, a medium consisting primarily of yeast extract and peptone, should provide a substrate "buffer" against proteolytic degradation. Although levels of total activity for Kexlp-Xho, -AS, and -AMS was occurring external to the plasma membrane, then the addition of BSA to the growth medium may lessen the extent of degradation. The addition of relatively high levels of BSA did not, however, alter the levels of total activity for Kexlp-Xho, -AS, and -AMS.

A different approach to reduce potential degradation of truncated forms of Kexlp was taken where transformants were grown temporarily under nonselective conditions, plasmid loss was never >2% (data not shown). Activity partitioning data suggested that Kexlp-Xho, -AS, and -ΔMS were secreted from the cell while the other constructs (Kexlp, Kexlp-Hpa, and Kexlp-Hinc) remained intracellular (Table I). All the mutant forms of Kexlp had approximately the same total activity as the plasmid-borne wild type. The percentage of Kexlp extracellular activity for the membrane-associated forms of Kexlp was comparable to the percentage of cells that stained positively with the vital dye methylene blue, suggesting that such extracellular activity resulted from cell lysis. KEX1 was placed downstream from the GAL1 promoter which, upon induction by growth on galactose, resulted in a 70-fold increase over endogenous Kexlp activity. However, such overproduction did not increase the percentage of extracellular Kexlp activity relative to wild-type levels (Table I).

To demonstrate the secretion of the soluble forms of Kexlp, S86-16 (kex1-Δ2) was transformed with pXKI-18 (Kexlp) or pXKI-18ΔMS (Kexlp-ΔMS), and cells were then spheroplasted. The spheroplasts were radiolabeled (10 min), chased (30 min), and solubilized with SDS, and Kexlp was immunoprecipitated. Media, containing proteins exported

Table I. Activity Partitioning of Kexlp Truncations in YEPD

| Strain         | Activity in the medium | Activity in the periplasm | Intracellular activity | Total activity | % of Total activity | Units/Optical Density |
|----------------|------------------------|---------------------------|------------------------|----------------|---------------------|-----------------------|
| A. S86         | 4.8                    | 1.2                       | 94                     | 4.3            |                     |                       |
| S86-17         | 0                      | 0                         | 0                      | 0              |                     |                       |
| S86-17/ Xho    | 72                     | 9                         | 19                     | 0.45           |                     |                       |
| S86-17/ AS     | 67                     | 8                         | 25                     | 0.57           |                     |                       |
| S86-17/ ΔMS    | 70                     | 8                         | 22                     | 0.6            |                     |                       |
| S86-17/ Hpa    | 7                      | 3                         | 90                     | 0.65           |                     |                       |
| S86-17/ Hinc   | 1.9                    | 5.1                       | 93                     | 0.49           |                     |                       |
| S86-17/ WT     | 3.5                    | 3.5                       | 93                     | 0.44           |                     |                       |
| B. S86         | 9.5                    | 4.4                       | 86                     | 1.6            |                     |                       |
| S86/ GAL-KEX1  | 4                      | 10                        | 86                     | 110            |                     |                       |

The indicated strains and transformants were inoculated in either YEPD (A) or YEP + galactose (B) to a high cell density and grown to stationary phase. The cells were then assayed for Kexlp activity as described (Cooper and Bussey, 1989). A unit is defined as 1 pmol of product produced per minute at 30°C. (Error ± 5%).

1987). Construction of these plasmids resulted in the deletion of an upstream portion of the KEX1 promoter. This deletion resulted in reduced production of the encoded proteins relative to wild-type levels. These constructs (the pXKI-20 series of plasmids) were transformed into the yeast strain S86-17 (kex1-Δ3), a derivative of S86 in which the KEX1 gene had been disrupted. The transformants were grown in liquid selective media; and the media, whole cells, and solubilized lysed cells were assayed for Kexlp activity (data not shown). S86-17 produced no activity and the plasmid-borne wild-type KEX1 gene partially restored activity (10% of the level produced by the genomic allele of KEX1) consistent with the promoter truncation.

In comparing the partitioning of Kexlp activity from various truncation mutants with that of the wild-type, two separate groups became apparent. The Kexlp-Hpa and Kexlp-Hinc truncated proteins (those that remained membrane associated; Fig. 3) showed the same partitioning pattern as that of wild-type, whereas the proteins Kexlp-Xho, -AS, and -ΔMS (soluble proteins lacking the membrane-spanning domain) formed a different pattern with a 10-fold increase in activity at the cell surface relative to that of wild type. The total activity of each protein was approximately constant within a group. Kexlp-Hpa and Kexlp-Hinc had a total activity similar to that of wild type while the soluble forms of Kexlp had ~50% of wild-type activity. Although the decrease in activity may have been a direct consequence of the mutations, it was also possible that the reduced activity was due to secretion and subsequent degradation of the truncated soluble proteins. If such degradation of the soluble forms of Kexlp-Xho, -AS, and -ΔMS was occurring external to the plasma membrane, then the addition of BSA to the growth medium may lessen the extent of degradation. The addition of relatively high levels of BSA did not, however, alter the levels of total activity for Kexlp-Xho, -AS, and -ΔMS.

A different approach to reduce potential degradation of truncated forms of Kexlp was taken where transformants were grown in selective conditions (minimal media) and then transferred to nonselective YEPD media for several generations before assay. YEPD, a medium consisting primarily of yeast extract and peptone, should provide a substrate "buffer" against proteolytic degradation. Although transformants were grown temporarily under nonselective conditions, plasmid loss was never >2% (data not shown). Activity partitioning data suggested that Kexlp-Xho, -AS, and -ΔMS were secreted from the cell while the other constructs (Kexlp, Kexlp-Hpa, and Kexlp-Hinc) remained intracellular (Table I). All the mutant forms of Kexlp had approximately the same total activity as the plasmid-borne wild type. The percentage of Kexlp extracellular activity for the membrane-associated forms of Kexlp was comparable to the percentage of cells that stained positively with the vital dye methylene blue, suggesting that such extracellular activity resulted from cell lysis. KEX1 was placed downstream from the GAL1 promoter which, upon induction by growth on galactose, resulted in a 70-fold increase over endogenous Kexlp activity. However, such overproduction did not increase the percentage of extracellular Kexlp activity relative to wild-type levels (Table I).

To demonstrate the secretion of the soluble forms of Kexlp, S86-16 (kex1-Δ2) was transformed with pXKI-8 (Kexlp) or pXKI-18ΔMS (Kexlp-ΔMS), and cells were then spheroplasted. The spheroplasts were radiolabeled (10 min), chased (30 min), and solubilized with SDS, and Kexlp was immunoprecipitated. Media, containing proteins exported

Figure 5. Mislocalization of Kexlp-ΔMS to the periplasm. S86-16/pXKI-8 (Kexlp-WT) or S86-16/pKXI-18ΔMS (Kexlp-ΔMS) were spheroplasted and then radiolabeled for a 10-min pulse (P). Half of the spheroplasts and media were harvested while the remainder was chased for an additional 30 min before harvest. The forms of Kexlp were immunoprecipitated from both the media and the spheroplasts and analyzed by SDS-PAGE and fluorography. The arrow indicates the full length Kexlp-ΔMS.

Cooper and Bussey Yeast Kexlp Is a Golgi-associated Membrane Protein
Kexlp resides in the yeast Golgi apparatus while precursor to a lesser extent than Kexlp. Kexlp-Bcl and that a Kexlp signal could only be detected upon overproduction of Kexlp (Fig. 7). Kexlp was found to be localized to proteins differed. The expression of the proteins was such or vacuoles, but were characteristic of proteins localized to the yeast Golgi apparatus (Redding et al., 1991; Cleves et al., 1991; Franzusoff et al., 1991; Roberts et al., 1992).

Comparison of the Kexlp-Hpa localization with that of the vacuole (as determined by Nomarski optics) indicated that almost all of Kexlp-Hpa was in the vacuole (data not shown). It is likely that Kexlp-Hpa was transported to the vacuole, where, in a protease-deficient strain such as $86-16$, the protein was not degraded and remained active.

To confirm this possibility, colocalization studies were undertaken between the 60-kD subunit of the S. cerevisiae vacuolar membrane H+ -ATPase and either Kexlp or Kexlp-Hpa. The vacuole (as determined by Nomarski optics) correlated to the ring structure in which the 60-kD ATPase subunit was localized by indirect immunofluorescence. The level of Kexlp detected varied among cells due to the variable plasmid copy number (2 $\mu$m based). Observation of cells expressing both low and high levels of Kexlp-Hpa (as deter-

The Effect of Carboxy-terminal Truncations of Kexlp upon K1 Killer Toxin Processing

A stringent test of the effect of the truncations of Kexlp was to determine the ability of the mutant proteins to carry out the Golgi-based proteolytic processing of K1 killer toxin in vivo (for review see Bussey, 1988). Sc25k is a strain harboring the dsRNA virus that encodes K1 killer toxin, and produces a killing zone on plates seeded with a strain sensitive to the killer toxin (Bussey et al., 1983). The pKXI-20 series of plasmids (centromeric based) were transformed into Sc25k-13 (kex/-A/), and the transformants analyzed for their ability to process K1 killer toxin and produce a killing zone (Fig. 6 A). Sc25k-13 containing the vector plasmid produced no killing zone, whereas the introduction of a plasmid-based copy of KEXI (pKXI-20WT) enabled the strain to produce active K1 killer toxin as shown by the appearance of a zone (Fig. 6 B). This zone was smaller than that of Sc25k (KEXI) because of the KEXI promoter truncation in pKXI-20WT (discussed above). The truncated forms of Kexlp, both soluble and membrane associated, produced killer zones significantly smaller than that produced by pKXI-20WT (Fig. 6 A). This functional complementation assay was also conducted in a different K1 killer strain (TC106x17) with similar results.

The reduced processing of K1 killer toxin by the soluble forms of Kexlp can be explained because of their secretion and consequent reduced concentration of Kexlp within the processing compartment. The membrane-associated truncated forms of Kexlp remain intracellular and, therefore, secretion cannot account for their reduced processing ability. The above results indicated that Kexlp-Hpa was membrane associated, had received glycosyl modifications in the Golgi apparatus, gave wild-type levels of total activity, and was retained intracellularly; yet processed the K1 killer toxin and consequent reduced concentration of Kexlp within the cell) which were not associated with mitochondria, nuclei, or vacuoles, but were characteristic of proteins localized to the yeast Golgi apparatus (Redding et al., 1991; Cleves et al., 1991; Franzusoff et al., 1991; Roberts et al., 1992).

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Figure 6. Production of active K1 killer toxin by the truncated forms of Kexlp. (A) Left column shows five different transformants of Sc25k-13 with pKXI-20WT. Right column shows Sc25k-13 transformed with one of the following plasmids: pKXI-20-Xho (Xho), pKXI-20-AS (AS), pKXI-20-AMS (AMS), pKXI-20-Hpa (Hpa), or pKXI-20-Hinc (Hinc). Transformants were grown to stationary phase in minimal media, harvested by centrifugation, washed in H2O, pelleted, and then resuspended in H2O to a concentration of 5 x 10^8 cells ml^-1. 10 $\mu$l was then placed onto a minimal plate (pH 4.7), seeded with S6 (a strain sensitive to K1 killer toxin). Plates were then incubated at 18°C for 24 h and zones of toxin killing examined. Approximate relative toxin activities, determined by comparison with a toxin dilution series were WT 100%, Xho 30%, AS 40%, AMS 15%, Hpa 55%, and Hinc 55%. (B) Sc25k-13 (ked), Sc25k (KEXI), and Sc25k-13/pKXI-20WT (WT) were grown and spotted onto a plate seeded with a strain sensitive to K1 killer toxin and treated as described above.

Kexlp Resides in the Yeast Golgi Apparatus While Membrane-associated Mutant Forms Are Mislocalized to the Vacuolar Membrane

Indirect immunofluorescent detection of Kexlp and Kexlp-Hpa was undertaken to determine if the location of the two proteins differed. The expression of the proteins was such that a Kexlp signal could only be detected upon overproduction of Kexlp (Fig. 7). Kexlp was found to be localized to a number of small punctate structures (on average, ~3–5 per cell) which were not associated with mitochondria, nuclei, or vacuoles, but were characteristic of proteins localized to beyond the plasma membrane, were concentrated and Kexlp was immunoprecipitated. Wild-type Kexlp remained associated with the spheroplasts, whereas Kexlp-AMS was shown to be secreted from the cell; the Kexlp-AMS signal associated with the cell fraction diminished with time while the signal from the medium showed the reverse trend (Fig. 5). Thus, the results of the pulse-chase analysis correlated with the activity data and indicated that Kexlp-AMS, a soluble form of Kexlp, was secreted from the cell.
Figure 8. Indirect immunofluorescence detection of truncated Kexlp in S. cerevisiae. S86-16/pKXI-18Hpa (Kexl-Hpa) cells were prepared as described in Fig. 7 above and viewed under FITC (which indicates the position of the 60-kD subunit of the vacuolar membrane H+-ATPase) and Texas red (which indicates the position of Kexlp-Hpa) excitation wavelengths.

Figure 7. Indirect immunofluorescence detection of Kexlp in S. cerevisiae. S86-16/pVT103-L (kexl-A), and S86-16/pKXI-8 (Kexl-WT) were prepared for immunofluorescence as described in Materials and Methods. The cells were viewed by Nomarski differential interference optics, and under 4',6-diamidino-2-phenyl-indole (indicates the position of the nucleus and mitochondria), FITC (indicates the position of the 60-kD subunit of the vacuolar ATPase), and Texas red (indicates the position of Kexlp) excitation wavelengths.

**Discussion**

**Kexlp Resides in the Yeast Golgi Apparatus**

An analysis of the oligosaccharide modification of Kexlp indicated the presence of secretory compartments to which the protein had been exposed, and the likely subcellular location of Kexlp. The modification of the Asn-linked oligosaccharide cores of Kexlp occurred within the Golgi apparatus and likely involves several sequential steps. Initially, in a pre-sec7 compartment, the core oligosaccharides are partially modified as judged by an increase in apparent molecular mass (Cooper and Bussey, 1989). Subsequently, in a post-sec7 compartment, the core is further modified in a MNN1-dependent manner. Invertase produced in sec7 cells at the restrictive temperature does not receive α(1→3) linked mannose residues to the core or outer chain (Franzusoff and Schekman, 1989), most likely because of the failure of secretory proteins to reach the compartment containing the α(1→3) mannosyltransferase. It is consistent that sec7 and mnn1 mutations result in a similar reduction in the glycosyl elaboration of Kexlp, as they would respectively either prevent Kexlp from reaching, or reduce the activity of, the α(1→3) mannosyltransferase. The Golgi-based modification of Kexlp can thus be explained as a result of two processes. The first occurs before the sec7 block and most likely involves the addition of an α(1→6) mannose residue followed by the attachment of an α(1→2) linked mannose residue (indicated by * in Fig. 1). The second step involves the MNN1-dependent addition of up to three α(1→3) linked mannose residues (indicated by a circle in Fig. 1) to the oligosaccharide (Ballou et al., 1990).

Graham and Emr (1991) provided evidence for at least three functional compartments in the yeast Golgi apparatus that contain, from cis to trans, the following activities: (1) α(1→6) mannosyltransferase, (2) α(1→3) mannosyltransferase, and (3) Kex2p endoprotease. Kexlp must reach the Golgi compartment housing the MNN1-dependent α(1→3) mannosyltransferase as it is modified by this enzyme. In addition, to process killer toxin, Kexlp must reach the proposed third Golgi compartment containing Kex2p as the toxin precursor substrate for Kexlp is created by a Kex2p-mediated endoproteolytic cleavage. No post-Kex2p compartment before secretory vesicles has been observed (Graham and Emr, 1991) and, therefore, it is likely that Kexlp resides with Kex2p in the same late Golgi compartment (Redding et al., 1991). It is interesting to note that both Kexlp and Kex2p
Kexlp-AMS which is also secreted. The results suggest that contains all but 11 residues of the 596-residue lumenal domain to the Golgi apparatus in both, are responsible for the retention of Kexlp within the Golgi apparatus.

The Cytoplasmic Domain of Kexlp Confers Retention in the Golgi Apparatus

Of the mutations that result in a soluble form of Kexlp (Kexlp-Xho, -AS, and -ΔMS), Kexlp-AS is the most informative with regard to determining which domain(s) of Kexlp is involved in its localization. Kexlp-AS is secreted, yet contains all but 11 residues of the 596-residue lumenal domain (Fig. 3). The 11 residues absent in Kexlp-AS are present in Kexlp-ΔMS which is also secreted. The results suggest that either the membrane-spanning or cytoplasmic domain, or both, are responsible for the retention of Kexlp within the Golgi apparatus.

Kexlp-Hpa contains both the luminal and transmembrane domains of Kexlp yet is mislocalized to the vacuolar membrane. Similar results were found for Kexlp-Bcl and Kexlp-Hinc which lack all or part of the cytoplasmic domain and strongly implicate this domain of Kexlp as being necessary for correct localization within the Golgi apparatus.

Previous work has demonstrated the role that cytoplasmic domains of transmembrane proteins play in the retention and targeting of proteins within the secretory pathway. A consensus retention motif has been identified at the termini of the cytoplasmically exposed domain of mammalian ER resident type I transmembrane proteins (Nilsson et al., 1989; Jackson et al., 1990). The cytoplasmic domains of the transferrin receptor, cation-independent mannose-6-phosphate receptor, and the lysosomal acid phosphatase are involved in the correct targeting or endocytosis of these proteins (Rothenberger et al., 1987; Lobel et al., 1989; Peters et al., 1990).

The three proteins involved in processing α-factor, Kexlp, Kex2p, and DPAP A, are all transmembrane proteins with cytoplasmic domains which are thought to be involved in retention of the respective proteins in similar if not identical compartments of the yeast Golgi apparatus (Fuller et al., 1989; Roberts et al., 1992). Attempts to identify a consensus yeast Golgi retention motif shared between these proteins have, however, showed no obvious amino acid sequence homology between the cytoplasmic domains of these three proteins.

**Retention of Membrane Proteins in the Golgi Apparatus**

Different mechanisms have been proposed for the retention of proteins within Golgi compartments. Interactions involving the cytoplasmic domain of Kexlp may result in the aggregation of the protein in the appropriate compartment, and thereby prevent it from exiting transport vesicles that are exiting the Golgi compartment (Pfeffer and Rothman, 1987). This retention via aggregation model is unlikely as it does not explain why aggregation would not occur earlier in the secretory pathway or why overproduction of a Kexlp results in mislocalization to the vacuole rather than greater aggregation in the Golgi compartment. A more likely explanation for the retention involves a receptor interacting with the cytoplasmically exposed domain of Kexlp (Fig. 9). High level expression of Kexlp would saturate the receptor with the result that excess Kexlp is diverted to the vacuole. Removal of the cytoplasmic domain would abolish such a receptor–ligand interaction, also resulting in mislocalization to the vacuole. Retention of soluble ER proteins has been shown to involve a receptor-mediated retrieval of proteins from a pre-cis-Golgi compartment (Semenzato et al., 1990; Lewis et al., 1990). A cytoplasmically based retrieval system analogous to the ER system can be envisaged in which Kexlp is retrieved from a trans compartment, possibly the plasma membrane, and returned to the correct Golgi compartment. To determine if Kexlp is retrieved from the plasma membrane an immunofluorescence experiment was performed by inducing expression of Kexlp in a *sec1* strain. *sec1* is a conditional mutation that, at the nonpermissive temperature, prevents the fusion of secretory vesicles with the plasma membrane (Novick et al., 1981). The delivery of Kexlp to its Golgi location was not blocked by *sec1*, and if KEX1 expres-
sion was constitutive before imposition of the sec1 block, the staining pattern of Kexlp was not altered even after 3 h at the restrictive temperature (data not shown). Therefore, if a receptor retrieval system exists, it is unlikely that Kexlp is retrieved from the plasma membrane as outlined in one of several proposals presented by Payne and Schekman (1989). If a retrieval system is responsible for the retention of Kexlp, then the salvage compartment may instead be located between the Golgi compartment and the vacuole where failure to bind the receptor would result in delivery to the vacuole. Alternatively, the receptor may remain anchored within the Golgi apparatus and not recycle. The clathrin heavy chain has been implicated in the retention of Kex2p (Payne and Schekman, 1989), raising the possibility that the receptor-based system that recognizes the cytoplasmic domain of Kexlp, and potentially Kex2p and DPAP A, might be one of the clathrin-associated adaptor proteins known to occur in yeast (Kirchhausen, 1990).

Default Targeting of Yeast Membrane Proteins to the Vacuole

Truncated forms of Kexlp which lacked the cytoplasmic domain yet still remained membrane associated were not retained in the correct Golgi compartment, and even at subwild-type expression levels were diverted to the vacuole. Similar mislocalization results have been found with DPAP A, where overexpression of the protein results in mislocalization to the vacuole, as do mutations within its cytoplasmically exposed domain (Roberts et al., 1992). Differing results were obtained with Kex2p, where a mutation that deleted the cytoplasmic domain and part of the membrane-spanning domain resulted in a significant proportion of the Kex2p activity being mislocalized to the cell surface (Fuller et al., 1989). The Kex2p activity study did not, however, address whether the truncated protein produced was membrane associated; if not, then the resulting soluble protein would be expected to be secreted to the cell surface. In addition, the strain used for such Kex2p activity studies contained wild-type activity levels of vacuolar hydrolases (PEP4; Jones, 1984) and, therefore, any Kex2p potentially mislocalized to the vacuole might be degraded and hence undetected.

Mislocalization of membrane proteins to the lysosome does not appear to occur in mammalian systems where mutations that interfere with the localization of ER, Golgi, and lysosomal integral membrane proteins result in their delivery to the plasma membrane (Paabo et al., 1987; Machamer and Rose, 1987; Peters et al., 1990). One exception to this observation in mammalian systems is the coronavirus El protein, a type III membrane protein normally resident in the Golgi apparatus. Mutations that affect the retention of this protein resulted in delivery to the lysosome rather than to the cell surface (Armstrong et al., 1990).

A number of models could explain the observation that the membrane-associated mutant forms of Kexlp are delivered to the vacuole while the soluble truncated forms of Kex1 are exported to the cell surface (Fig. 9). (1) The first model suggests that membrane-associated truncated forms of Kexlp are misfolded, and as such may be targeted, via an unknown mechanism, to the vacuole for degradation. This model seems unlikely as all of the truncated forms of Kexlp, both soluble and membrane associated, have similar total protease activity to that of the wild-type protein, suggesting that at a minimum the catalytic domain of the mutants has folded to a conformation similar to that of wild type. In addition, these forms of Kexlp all exit the ER and reach the Golgi apparatus where they are both glycosylated and process killer toxin. The misfolding targeting model would be unusual in that it must explain the mislocalization of the membrane-associated mutant forms of Kexlp yet allow the soluble forms to be secreted. Also, such a garbage pathway for delivery of misfolded mutant forms of Kexlp cannot explain the vacuolar localization of highly expressed but presumably correctly folded Kexlp. (2) The second model proposes that the membrane-spanning domain of Kexlp contains a latent or cryptic targeting signal for the vacuole. Studies are currently under way to address this possibility. It should be noted that although Kexlp is homologous to the vacuolar protein CPY, the homology does not extend to include the vacuolar targeting signal found in proCPY. (3) The third model involves Kexlp-Hpa (-Bcl, -Hinc) being secreted to the plasma membrane by default, where it is then endocytosed and delivered to the vacuole. Such a "transient appearance" of Kexlp-Hpa (-Bcl, -Hinc) at the cell surface would not be detected by the activity assay used or by indirect immunofluorescence. Soluble Kexlp-AS, having reached the cell surface, would be released into the medium and, hence, would not be endocytosed to the vacuole. (4) The final model proposes that the vacuole is the direct default destination for membrane-associated proteins that enter the secretory pathway, such that membrane proteins lacking positive targeting/retention signals would be delivered to the vacuole.

Work with DPAP B, a type II vacuolar membrane protein homologous to DPAP A, has shown that no domain of the protein contains positive targeting information for the vacuole (Roberts et al., 1992). The targeting signal for a second vacuolar membrane protein, alkaline phosphatase, has not been identified although the lumenal domain is not required for the correct localization of this protein (Klionsky and Emr, 1990). The authors concluded that the vacuolar-sorting determinant must therefore reside in the cytoplasmic and/or membrane-spanning domain of alkaline phosphatase, but in light of the data presented here and elsewhere (Roberts et al., 1992), these results could be reinterpreted to suggest that the membrane association of alkaline phosphatase mediates its vacuolar delivery.

Given the results presented here, we favor models 3 and 4 which are variants of each other in that both result in the delivery of Kexlp to the vacuole by default (no positive targeting signal involved) but differ in the delivery route. Model 4, involving a direct Golgi-to-vacuole route, may be the most likely as delivery of the vacuolar membrane protein DPAP B does not involve transport to the plasma membrane and subsequent endocytosis (Roberts et al., 1992). In addition, Kexlp-Bcl contains no cytoplasmic tail to participate in classical endocytosis.

An important implication of such delivery is that yeast integral membrane proteins destined for the plasma membrane must have positive targeting information to either remain at the plasma membrane (proposal 3), or to avoid being diverted to the vacuole (proposal 4). It is pertinent that a mutation in the α-factor receptor (Ste3p), an integral plasma membrane protein, results in delivery of this protein to the vacuole, independent of the plasma membrane (Ho-
recka, J., and G. Sprague, personal communication), raising the possibility that a plasma membrane targeting signal has been destroyed.

Further work is proceeding to determine if the delivery route of Kex1p to the vacuole is via the plasma membrane, and whether plasma membrane proteins have positive targeting/retention information.

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