Clathrin-dependent Localization of α1,3 Mannosyltransferase to the Golgi Complex of Saccharomyces cerevisiae

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Abstract. Posttranslational modification of yeast glycoproteins with α1,3-linked mannose is initiated within a Golgi compartment analogous to the medial Golgi cisternae of higher eukaryotes. We have characterized the synthesis, posttranslational modification, and localization of the yeast α1,3 mannosyltransferase (Mnnlp) using antibodies prepared against a segment of this protein expressed in bacteria. Mnnlp is initially synthesized as a 98.5-kD, type II integral membrane glycoprotein that is modified with both N- and O-linked oligosaccharides. It is subject to a slow, incremental increase in molecular mass that is dependent upon protein transport to the Golgi complex. Self-modification of Mnnlp with α1,3 mannose epitopes, primarily on O-linked oligosaccharides, is at least partly responsible for the incremental increase in molecular mass.

Mnnlp is a resident protein of the Golgi complex and colocalizes with guanosine diphosphatase to at least two physically distinct Golgi compartments by sucrose gradient fractionation, one of which may be a late Golgi compartment that also contains the Kex2 endopeptidase. Surprisingly, we found that a significant fraction of Mnnlp is mislocalized to the plasma membrane in a clathrin heavy chain temperature sensitive mutant while guanosine diphosphatase remains intracellular. A mutant Mnnlp that lacks the NH2-terminal cytoplasmic tail is properly localized to the Golgi complex, indicating that clathrin does not mediate Mnnlp Golgi retention by a direct interaction with the Mnnlp cytoplasmic tail. These results indicate that clathrin plays a broader role in the localization of Golgi proteins than anticipated.

The initial events of protein glycosylation in yeast and mammalian cells are essentially the same, but Saccharomyces cerevisiae lacks the glycosyltransferases for complex sugars and so only produces glycoproteins with extended chains of mannose. Maturation of oligosaccharides on yeast glycoproteins requires the sequential action of α1,6, α1,2, and α1,3 mannosyltransferases within the Golgi complex of the secretory pathway. These enzymes catalyze the linkage-specific transfer of mannose from a GDP-mannose donor to N- and/or O-linked oligosaccharides. It is subject to a slow, incremental increase in molecular mass that is dependent upon protein transport to the Golgi complex. Self-modification of Mnnlp with α1,3 mannose epitopes, primarily on O-linked oligosaccharides, is at least partly responsible for the incremental increase in molecular mass.

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The mnnl mutant was isolated by an immunological enrichment procedure in which mutagenized cells were precipitated with an antiserum specific to α1,3 linked mannose epitopes. Nonreactive cells remained in suspension and were enriched for mutants that failed to display the α1,3 linked mannose epitope on their cell walls (46). The mnnl mutant lacks α1,3 mannosyltransferase activity, suggesting that the MNN1 gene encodes this enzyme (5). The MNN1 gene was recently cloned and its sequence predicts a type II integral membrane protein of 88.6 kD (66). While yeast has only one α1,3 mannosyltransferase, there appear to be multiple α1,6 and α1,2 mannosyltransferases (22). The genes that encode an initiating α1,6 mannosyltransferase (OCH1) (36), and an α1,2 mannosyltransferase (MNT1/KRE2) (19, 23) have been cloned and sequenced. Surprisingly, these proteins do not exhibit sequence homology to each other or to Mnnlp, although all three proteins are predicted to be type II integral membrane proteins with short cytoplasmic tails. This structure is also a feature of several mammalian glycosyltransferases (43).

In previous work, we proposed that the yeast Golgi complex is divided into at least three functionally distinct compartments (16). Within these compartments are catalyzed, from cis to trans, the α1,6 mannosylation, α1,3 mannosylation, and Kex2p-mediated proteolytic processing of glycoproteins. This conclusion was based on the requirement for the Sec18/NSF intercompartmenal protein transport factor for each successive modification in vivo (16). These experiments indicated that the α1,3 mannosyltransferase activity is

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first encountered in a medial-Golgi compartment. Based on these observations, we reasoned that Mnnlp should be a resident marker protein for the yeast medial- and perhaps trans-Golgi compartments.

One approach towards understanding the biogenesis of the Golgi complex is to ask how the resident proteins are localized to this organelle. In the case of the late Golgi enzymes Kex1p, Kex2p, and dipetidylaminopeptidase A (DPAP A)\(^{1}\) that are required for the final proteolytic maturation of the \(\alpha\)-factor mating pheromone precursor, it was found that all of these proteins have Golgi localization signals within their cytoplasmic tails (10, 39, 63). The Kex2p and DPAP A Golgi localization signals are short peptide segments containing essential tyrosine or phenylalanine residues that are very similar to the signals within mammalian receptor proteins that mediate clustering into clathrin coated pits (reviewed in 40, 64). A role for clathrin in the Golgi localization of Kex2p and DPAP A was demonstrated using yeast strains harboring null or temperature-sensitive alleles of the clathrin heavy chain gene (chcl) (44, 53). Loss of clathrin function in these strains results in the mislocalization of Kex2p and DPAP A to the plasma membrane. Guanosine diphosphatase (GDPane), another Golgi enzyme that appears to mark earlier compartments of the Golgi complex, is not mislocalized to the plasma membrane in the clathrin mutants (53). It is thought that binding of the cytoplasmic tails of Kex2p and DPAP A to clathrin and associated protein (AP-1) coat complexes either mediates direct retention in the late Golgi by a tethering mechanism, or the recycling of these proteins from either the endosome or the yeast equivalent of a condensing secretory granule. It does not appear that these proteins are recycled to the Golgi complex by endocytosis from the plasma membrane (reviewed in 64). Surprisingly, deletion of the Golgi localization signals of Kex1p, Kex2p and DPAP A result in the mislocalization of these proteins to the vacuole rather than the plasma membrane. In addition, overexpression of these proteins also leads to their partial mislocalization to the vacuole (10, 48, 63). These observations have led to the proposal that in yeast, the default (signal-independent) destination of integral membrane proteins is the vacuole, rather than the plasma membrane as is the case in mammalian cells (48).

Analysis of Golgi localization signals within two mammalian glycosyltransferases (34, 37) and the coronavirus E1 protein (58) has resulted in the unexpected finding that transmembrane domains in these proteins can target a reporter enzyme to the Golgi complex. The localization of these proteins is not easily saturable by overexpression, suggesting that specific interaction with a receptor may not be involved in the retention mechanism. Evidence exists to support two hypotheses for the mechanism of transmembrane domain mediated Golgi retention. One model suggests that the length of a transmembrane domain will specify Golgi localization by selectively associating with membranes of the appropriate thickness. The lipid and sterol composition of the Golgi membrane would determine the membrane thickness and would consequently specify the membrane proteins that are retained within each compartment (7). The second model suggests that Golgi resident membrane proteins form aggregates within the appropriate compartment resulting in a protein matrix that is too large to enter into budding transport vesicles (kin recognition hypothesis, references 38, 45). Neither model would require trans-acting protein receptors such as the Erd2 protein which recycles HDEL containing ER proteins from the Golgi back to the ER (32, 54). Moreover, it would seem unlikely that clathrin would mediate the Golgi retention of these mammalian glycosyltransferases via a tethering or recycling mechanism, because the primary localization signal is within the transmembrane domain rather than the cytoplasmic tails of these proteins.

In this work, we provide the first detailed analysis of the synthesis, posttranslational modification, and sorting of a yeast mannosyltransferase. Our data indicate that Mnnlp is a type II integral membrane, resident protein of the yeast Golgi complex. The Golgi marker enzyme GDPase colocalizes with Mnnlp in sucrose density gradients, but Kex2p colocalizes with only a denser membrane fraction of Mnnlp and GDPase. We also present evidence that clathrin is required for efficient Golgi localization of the Mnnlp by a mechanism independent of the Mnnlp cytoplasmic tail.

Materials and Methods

Strains and Media

The yeast strains used were XCY42-30D (MATa ura3 leu2-3,112 trpl lys2 ade2-101 ade6 suc2-\(\Delta 9\)), XCY42-30D Dmnnl::LEU2 (MATa ura3 leu2-3,112 trpl lys2 ade2-101 ade6 suc2-\(\Delta 9\) Dmnnl::LEU2 (this study)), SEY-6210 (MATa ura3-52 leu2-3,112 his3-D200 trpl-\(\Delta 901\) lys2-801 suc2-\(\Delta 9\) [50]), TVY1 (SEY6210 Adep4::LEU2 [57]) SEY5188 (MATa sec18-1 leu2-3,112 ura3-52 suc2-\(\Delta 9\) [16]), SF274 3A (MATa sec17-2), SF294-2B (MATa sec7-1), HMSC-1 (MATa sec-1), SF292-1A (MATa sec4-3 [Randy Schekman, University of California, Berkeley, CA]), GPY382 (MATa ura3-52 leu2-3,112 his3-D200 trpl dpp2p: HIS3 chcl-\(\Delta 10\) LEU2 YeChCl102) GPY-383 (MATa ura3-52 leu2-3,112 his3-D200 trpl dpp2p: HIS3 chcl-\(\Delta 10\) LEU2 YeChCl51), TGY31 (XCY42-30D/XCY42-30D p2V236), TGY32 (XCY42-30D/XCY42-30D), TGY33 (XCY42-30D/XCY42-30D Dmnnl::LEU2/XCY42-30D Dmnnl::LEU2). Isogenic MATa diploid strains were prepared by transforming the corresponding haploid strain with the HO gene carried on pHO-\(\Delta 2\) (51) to induce mating type switching and mating within transformed colonies. After restreaking transformants twice on selective plates, individual colonies were picked randomly and tested for the secretion of a mating factors on lawns of supersensitive yeast. Diploid colonies that secreted neither mating factor were cured of pHO-\(\Delta 2\) to give TGY31-TGY33.

Standard rich (YPD) and synthetic minimal (SD) media for yeast was used (55). The SD medium was supplemented with 0.2% yeast extract (SDYE) and other supplements as needed for growing cells in liquid culture before labeling experiments. Standard rich medium for Escherichia coli (33) was used.

Reagents

Zymolyase-100T (Kirin Brewery Co.) was obtained from Seikagaku Kogyo Co. (Tokyo, Japan), endoglycosidase H (endo H) was from New England Nuclear Corp. (Boston, MA), DNA modifying enzymes were from New England Biolabs, Inc. (Beverly, MA), proteinase K was from Boehringer Mannheim Biochemicals (Indianapolis, IN), Protein A-Sepharose and CNBr activated Sepharose 4B was from Pharmacia (Piscataway, NJ), Trans\(^{35}\)S-label was from ICN radiochemicals (Irvine, CA), the ECL detection kit was from Amersham (Arlington Heights, IL) and 0.2-0.3 mm glass beads were from Glen Mills Inc. (Maywood, NJ). All other chemicals were purchased from Sigma Chemical Co. (St. Louis, MO). Anti-sr to \(\alpha\)-mannose linkages, alkaline phosphatase, GDPane, and Ochlp were from R. Schekman (UC Berkeley, Berkeley, CA), Gregory Payne (UCLA, Los Angeles, CA), Carlos Hirschberg (University of Massachusetts, Worcester, MA) and Yoshifumi Jigami (National Institute of Bioscience and Human

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1. Abbreviations used in this paper: ALP, alkaline phosphatase; DPAP A, dipetidylaminopeptidase A; endo H, endoglycosidase H; GDPase, guanosine diphosphatase; SD, synthetic minimal media; TEA, triethanolamine acetate; YE, yeast extract; YPD, standard rich medium.
Technology, Tsukuba, Ibaraki, Japan), respectively. The preparation of antiserum to CPY has been previously described (Klionisky et al., 1988). Secondary antibodies were from Jackson Immunoresearch (West Grove, PA).

**Plasmid Construction**

To prepare pMNNI-XCT, a 1.4-kb BamHI-EcoRI fragment containing the 5' end of the MNNI gene was subcloned from pVZ236 into the BamHI site of pRS426ARI to generate pRS426MNNI and TGY222A to generate pMNNI-XCT (2.8 μm). The BamHI-EcoRI fragment from pVZ236-XCT was used to replace that of pATH2-MNNI at the BamHI-EcoRI site of pTGY222A, to generate pMNNI-XCT (2.8 μm).

**Immunological Techniques**

To prepare a bacterially expressed Mnnlp antigen, E. coli harboring pATH2-MNNI were induced to synthesize the trpE-Mnnlp fusion protein as described by Kjeld et al. (27) and the fusion protein was purified as described by Pavlincic et al. (42). Approximately 0.25 mg of gel-purified fusion protein was emulsified with Freund's complete adjuvant and injected intramuscularly into a male New Zealand white rabbit. The rabbit was boosted subcutaneously every three weeks with 0.05 mg of fusion protein in Freund's incomplete adjuvant. Bleeds were collected one week after each boost.

Affinity purified anti-Mnnlp antibodies were prepared by chromatography on a trpE-Mnnlp column. Approximately 0.8 mg of trpE-Mnnlp was coupled to CNBr activated Sepharose 4B following the manufacturer's protocol. 3 ml of anti-Mnnlp antiserum was diluted 1:1 with buffer A as defined by Redding et al. (47) and circulated through the fusion protein column for 3 h. The column was washed as previously described (47), and eluted with 0.1 M glycine-HCl, pH 2.75. The eluate fractions were neutralized by the addition of 0.1 M Tris-HCl, pH 9.4, and tested against dot blots of the fusion protein to identify antibody containing fractions. These fractions were pooled and stored in aliquots at −75°C.

Indirect immunofluorescence analysis was done as described by Redding et al. (47) with the following modifications. Fixed spheroplasts were permeabilized by incubation in SHA buffer (1 M sorbitol, 0.1 M Na Hepes pH 7.5, 5 mM Na Azide) plus 1% SDS for 10 rain at room temperature. Fixed, permeabilized cells were incubated for 3–4 h at room temperature with a 1:50 dilution of affinity purified α-Mnnlp antibodies and bound primary antibody was detected using FITC conjugated goat anti-rabbit secondary antibodies (1:1000 dilution). Cells were observed using a Zeiss Axiopt microscope with a 100x objective and photographed with a Tmax 400 film.

For labeling experiments, yeast cells were grown to mid-logarithmic phase in SD-YE, washed twice in water and resuspended in SD media with appropriate supplements at 5 OD600/ml. To initiate labeling, Tran35S-labeled proteins were blocked by adding a 30X chase solution (50 mM methionine, 10 mM cysteine and 5% yeast extract) to a 1X concentration. The chase was subsequently terminated by adding TCA to a final concentration of 10%. Processing of TCA pellets for immunoprecipitation and size fractionation of Mnnlp in 7% SDS-polyacrylamide gels was done as previously described (17).

For endo H treatment of immunoprecipitated, washed protein A-Sepharose immune pellets were dried, then resuspended in 64 μl of 1% β-mercaptoethanol, 0.2% SDS and heated at 95°C for 4 min. 16 μl of 250 mM Na citrate buffer, pH 5.5, was added to each tube, then 0.5 μl of endo H was added to 4 μl of each sample and incubated overnight in a 37°C incubator. The reactions were stopped by the addition of 4X Laemmli sample buffer and heating to 95°C for 5 min. For the experiment shown in Fig. 4, a Molecular Dynamics Phospholaser was used to quantitate the amount of 35S present within defined areas of the polyacrylamide gels. For the experiments shown in Figs. 1, 5, 7, and 8, an LKB laser densitometer was used to quantitate band intensities on autoradiograms.

**Subcellular Fractionation**

The association of Mnnlp with membranes and its accessibility to exogenously added protease were determined as follows. Strain XCY42-30D was grown in SD-YE to mid-logarithmic phase and labeled with the 35S amino acid labeling mix for 10 min at 30°C (20 OD600 U). Chase solution, an equivalent volume of 2X stop/spheroplast buffer (2 M sorbitol, 50 mM Tris–Cl, pH 7.5, 40 mM Na azide, 40 mM Na fluoride, 20 mM DTI) and 0.02 mg of Zymolyase were added and incubated for 30 min at 30°C. The spheroplasts were harvested, then resuspended in 0.18 ml of 1 M sorbitol, 20 mM trithionamide acetate (TEA), pH 7.2, and diluted to 1.8 ml with ice-cold 20 mM TEA, pH 7.2, to lyse the spheroplasts. One sixth of the samples, the remainder was centrifuged at 45,000 rpm for 30 min in a TL100.3 rotor. One fifth of the supernatant was TCA precipitated and the remainder was discarded, then the pellet was resuspended in 0.25 ml of 0.1 M sorbitol, 20 mM TEA, pH 7.2. 0.05-ml aliquots of the pellet fraction were added to: (a) 0.95 ml of 10% TCA; (b) 0.95 ml of 0.1 M Na carbonate, pH 11.0; (c) 0.95 ml of 1% Triton X-100, 20 mM TEA, pH 7.2; (d) 0.25 ml of 0.2 mg/ml proteinase K, 0.1 M sorbitol, 20 mM TEA, pH 7.2; and (e) 0.25 ml of 0.2 mg/ml proteinase K, 0.1 M sorbitol, 20 mM TEA, pH 7.2, 1% Triton X-100. All samples were incubated on ice for 30 min, then samples (b) and (c) were centrifuged as above and the pellet and supernatant fractions were TCA precipitated. The proteinase K incubations were terminated by adding PMSF to 1 mM and TCA to 10%. After precipitation, the TCA pellets were solubilized and subjected to immunoprecipitation as previously described (17).

Sucrose gradient fractionation of Golgi membranes was performed by adaptation of previously published protocols (6, 60). Strain XCY42-30D was grown in 1 L of YPD at 30°C to an OD600 of 1.0-1.4, then harvested and washed once with 10 mM Na azide. The cells were converted to spheroplasts as previously described (59), except 10 mM Na azide was added to all buffers and Zymolyase was used at 2 μg per OD600 of cells to digest the cell wall. The spheroplasts were washed once with spheroplasting buffer (1 M sorbitol, 20 mM Tris–Cl, pH 7.5, 10 mM Na azide), then resuspended in 1 M sorbitol, 10 mM TEA, pH 7.5, 1 mM EDTA at 100 OD600/ml and stored at −75°C in 1.5-ml aliquots. The spheroplasts (300 OD600) were thawed, then diluted seven fold with ice-cold lysis buffer (0.1 M sorbitol, 10 mM TEA, pH 7.5, 1 mM EDTA) and after 5 min on ice, were subjected to Dounce homogenization (25 strokes). The lysate was centrifuged at 1,000 g for 6 min to generate the PI (pellet) and the SI (supernatant) fractions, and the latter was centrifuged at 13,000 g to generate the PI3 and SI3 fractions. The SI3 was layered onto a two step sucrose cushion consisting of 1% 66% sucrose and 1 ml of 20% sucrose, then centrifuged in an SW41 rotor at 31,000 rpm for 2 h at 4°C. All sucrose solutions were prepared with/wi in 10 mM Na Hepes, pH 7.5, 1 mM Na azide (gradient buffer). The membranes present at the 20-66% sucrose interface were collected in a small volume as possible and the refractive index of the sample was measured using a Bausch and Lomb refractometer. The sample was then diluted with gradient buffer to ~20% sucrose and 1.5 ml of the membrane sample was layered on top of a sucrose step gradient prepared as described (59). The gradients were centrifuged in an SW41 rotor at 31,000 rpm for 17 h at 4°C. 16 fractions (<0.78 ml) were collected starting from the top of the gradient using a Buchner Auto-Densi Flow II and a Gilson fraction collector. Each fraction was diluted to 4.0 ml with gradient buffer and centrifuged again in a Beckman 50Ti or Sorvall Ti70 rotor at 40,500 rpm for 1.5 h at 4°C. After aspirating off the supernatants, the pellets were resuspended in 0.2 ml of gradient buffer and stored.

Kex2p and GDPhase were assayed as previously described (15, 65). Protein concentration was determined using the bicinchoninic acid protein assay kit from Sigma. Detection of Mnnlp on Western blots of gradient fractions was done by overnight incubation of the blot at 4°C with affinity purified α-Mnnlp antibodies (1:400 in TBST, 5% freeze dried milk) followed by incubation with horseradish peroxidase conjugated anti-rabbit antibodies and detection using the ECL kit from Amersham following the manufacturer's protocol.

**Results**

**Synthesis and Posttranslational Modification of Mnnlp**

To analyze the biosynthesis and intracellular localization of Mnnlp, we prepared a rabbit polyclonal antiserum against a
Figure 1. Synthesis and posttranslational modification of Mnnlp. 

(A) Strains XCY42-30D ΔMnnl (Δ), XCY42-30D (WT) and XCY42-30D pZV236 (2μ) were labeled for 10 min at 30°C as described in Materials and Methods and immunoprecipitations were performed using anti-Mnnlp (a-Mnnlp) or preimmune serum with anti-alkaline phosphatase (a-ALP) as indicated above each lane. pZV236 is a multicopy, 2μ plasmid harboring the MNN1 gene. Lane 3 was loaded with 1/10 the amount of sample as that in lanes 1, 2 and 4. 

|        | 1   | 2    | 3    | 4   |
|--------|-----|------|------|-----|
| Mnnlp  |     |      | -    |     |
| ALP    |     |      |      |     |
| Δ WT   |     | -    |      | 2μ  |

98.5 kD 76 kD 72 kD

(B) Strain XCY42-30D (wild type) was labeled for 10 min at 30°C, then chased for 0, 1.5, or 3 h (lanes 1-3) and processed for coimmunoprecipitation with antisera to Mnnlp and alkaline phosphatase (ALP). 

| Chase (h) | 0   | 1.5  | 3.0  |
|-----------|-----|------|------|
| Mnnlp     | 106 | 102.5| 98.5 |
| ALP       | 76  | 72   |      |

(C) Strain XCY42-30D (wild type) was labeled for 10 min at 30°C and chased for the times indicated. Labeled Mnnlp was recovered from each sample by immunoprecipitation and was eluted from the primary antibody by boiling in 1% SDS, 20 mM Tris–Cl, pH 7.5. Each sample was split in half and 

| Time (min) | 0   | 20  | 40  | 60  | 80  |
|------------|-----|-----|-----|-----|-----|
| % Mnnlp with α1,3 mannose | 0   | 20  | 40  | 60  | 80  |

The late Golgi enzymes Kex1p and Kex2p are subject to a slow posttranslational modification that results in a gradual increase in apparent molecular mass of these proteins (9, 62). As shown in Fig. 1 B, Mnnlp also exhibited a slow, incremental increase in molecular mass during a pulse/chase analysis. The 98.5-kD Mnnlp had increased in relative molecular mass to 102.5 kD at 1.5 h and 106 kD at 3 h of chase (Fig. 1 B, lanes 1–3). This is in contrast to ALP for which the 76-kD precursor undergoes a rapid PEP4-dependent proteolytic processing step to produce the 72-kD mature form (29), but does not increase in molecular mass thereafter (Fig. 1 B, lanes 1–3). In other experiments in which more time points were taken, we found that the gradual increase in mass of the Mnnlp was linear over a 3-h chase with a slope of ~2.5 kD per hour at 30°C. To test for the presence of α1,3 linked mannose on Mnnlp, we subjected this protein to a second immunoprecipitation using an antiserum specific to α1,3 linked mannose epitopes. Wild-type cells were labeled for 10 min and chased for the times indicated in Fig. 1 C. Mnnlp was recovered from the cells by immunoprecipitation and then was eluted from the primary antibody by boiling in 1% SDS. The samples were split in half and immunoprecipitated a second time with either anti-Mnnlp antiserum, or the linkage-specific antiserum. The amount of Mnnlp recovered in each pair of immunoprecipitates were compared over time. We found that the percentage of Mnnlp that could be precipitated with the linkage-specific antiserum also increased gradually with increasing time after synthesis (Fig. 1 C). The correlation of the slow rate of α1,3 mannose epiti...
tope acquisition with the slow rate of change in molecular mass suggests that self-modification of Mnnlp (auto-mannosylation) is at least partly responsible for the observed increase in molecular mass.

To determine if the auto-mannosylation of Mnnlp was occurring on O- or N-linked oligosaccharides, we treated Mnnlp with endo H which specifically cleaves N-linked oligosaccharides. A sec18 strain was labeled and chased for 0 and 3 h at the permissive temperature, then Mnnlp was recovered from cell lysates by immunoprecipitation and half of each sample was treated with endo H (Fig. 2, lanes 1–4). At 24°C, Mnnlp increased in apparent molecular mass from 98.5 kD (0 h, lane 1) to 104 kD (3 h, lane 2). If this posttranslational increase in molecular mass of Mnnlp was due to elaboration of N-linked oligosaccharides, then both forms of the enzyme should be converted to the same molecular mass after endo H treatment. This was not the case. Endo H treatment converted the 98.5-kD form to 92.5 kD (lanes 1 and 3) and the 104-kD form to 98 kD (lanes 2 and 4). The endo H treated Mnnlp from the 0 and 3 h chase times still differed by 5.5 kD (compare lane 3 with lane 4), as did the untreated 0 and 3 h forms. The molecular mass of both forms decreased by ~6 kD, which suggests that three of the four predicted N-linked glycosylation sites are used in vivo. After removal of N-linked oligosaccharides by endo H treatment, Mnnlp from the 3 h chase point was still precipitable with the α1,3 linkage specific antiserum (although not as efficiently as the untreated protein) indicating the presence of O-linked oligosaccharides on Mnnlp (data not shown).

O-linked glycosylation is thought to be initiated in the ER by the addition of a single mannose residue to Ser and Thr residues which are then extended in the Golgi complex with α1,2 and α1,3 linked mannose (22). To determine if the posttranslational increase in molecular mass of Mnnlp required protein transport to the Golgi complex, Mnnlp was also immunoprecipitated from sec18 cells labeled at a restrictive temperature (34°C) where ER to Golgi transport is blocked. Under these conditions, Mnnlp failed to increase in molecular mass during the chase period, demonstrating that protein transport out of the ER is required for this posttranslational modification (Fig. 2, lanes 5–8). These data indicate that the slow posttranslational increase in molecular mass of Mnnlp is primarily due to auto-mannosylation of O-linked oligosaccharides in a post-ER compartment, presumably the Golgi complex. The slow increase in molecular mass of Kex2p has also been shown to result from modification of O-linked oligosaccharides on Kex2p, and the extent of this modification on Kex2p was decreased in a mnn1 mutant (62).

**Membrane Association of Mnnlp**

The sequence of the MNN1 gene predicts a 762-amino acid protein with a single NH2-terminal hydrophobic domain encompassing amino acids 19–37 (66). This signal sequence is predicted to be uncleaved by signal peptidase based on the empirical rules of von Heijne (61), suggesting a dual role for the transmembrane domain as a signal sequence and membrane anchor which would result in the protein adopting a type II integral membrane topology. We addressed the membrane association and topology of Mnnlp by extraction of intracellular membranes with alkaline carbonate buffer and Triton X-100, and by a protease protection assay. Wild-type cells were labeled for 10 min, then converted to spheroplasts and lysed by osmotic shock. The lysate was subjected to centrifugation at 100,000 g to pellet membranes and one fifth of the pellet (P) and supernatant (S) fractions were immediately TCA precipitated (Fig. 3, lanes 2 and 3). The remaining membrane pellet was divided into four equal portions which were then treated with 0.1 M Na carbonate, pH 11.0 (lanes 4 and 5), 1% Triton X-100 (lanes 6 and 7), proteinase K (lane 8), or proteinase K plus 1% Triton X-100 (lane 9). The carbonate and Triton X-100 treated samples were centrifuged again at 100,000 g to produce pellet and supernatant fractions and the proteinase K treated samples were stopped by the addition of PMSF and TCA as described in Materials and Methods. Mnnlp, carboxypeptidase Y and glucose-6-phosphate dehydrogenase were recovered from each sample by sequential immunoprecipitation and subjected to SDS-PAGE. Nearly all the Mnnlp fractionated with a crude membrane high speed pellet (Fig. 3, lanes 2 and 3), while 95% of a cytoplasmic enzyme, glucose-6-phosphate dehydrogenase, remained in the supernatant (data not shown). Mnnlp sedimented in the membrane pellet after treatment of the membranes with high pH carbonate buffer (lanes 4 and 5), but was extracted from the membrane into the supernatant fraction with the detergent Triton X-100 (lanes 6 and 7). Precursors of carboxypeptidase Y that were present in the ER and Golgi were completely extracted from the membranes by the carbonate buffer, indicating that the carbonate had effectively removed soluble components from the lumen of subcellular compartments (lanes 4 and 5) as expected (14). These data indicate that Mnnlp is membrane associated and suggests that the NH2-terminal signal sequence was not cleaved. In addition, a large fragment of Mnnlp was protected from degradation by proteinase K added to the cytoplasmic face of the membranes (Fig. 3, lane 8). A small decrease in molecular mass of Mnnlp was observed in the proteinase K treated sample, indicating a portion of Mnnlp crossed the membrane and was accessible to the cytoplasm. This molecular mass shift (~1 kD) is consistent with the removal of the exposed NH2-terminal cytoplasmic tail. In the presence of proteinase K and Triton X-100, Mnnlp was completely degraded (lane 9) showing that an intact lipid bilayer was required for protection from the protease, and that this protein is not inherently protease resistant. These results, together with the hydropathy analysis of the MNN1

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**Figure 2.** Endo H treatment of Mnnlp synthesized in a sec18 strain. Strain SEY5188 (sec18) was labeled at 24°C (lanes 1–4) or 34°C (lanes 5–8, nonpermissive temperature) and chased for 0 or 3 h. After recovery of Mnnlp by immunoprecipitation, the samples were split in half and one of each pair was treated with endo H to remove N-linked oligosaccharides as described in Materials and Methods.
Compartmental Site of Mnnlp Modification

In order to assess the subcellular compartment in which the slow incremental increase in molecular mass of Mnnlp was catalyzed, we analyzed the extent to which Mnnlp was subject to this modification in sec mutants that block protein transport at various stages of the secretory pathway (41). The sec mutants indicated in Fig. 4 were preincubated at 37°C for 30 min before labeling to inactivate the sec gene products. The sec12 mutant exhibits a block in ER to Golgi protein transport and would be expected to accumulate newly synthesized Mnnlp in the lumen of the ER (26). As expected, this mutant exhibited a complete block in the modification of Mnnlp (Fig. 4, sec12). The sec7 mutant exhibits a partial block in ER to Golgi protein transport as well as transport steps between Golgi cisternae (13). This mutant exhibited a nearly complete block in the gradual modification of Mnnlp. The sec14 mutant harbors a defective phosphatidylinositol/phosphatidylcholine exchange factor which results in an altered phospholipid composition of the Golgi at 37°C and an inability to form transport vesicles from the late Golgi complex (4). Although not as defective as the sec12 or sec7 mutants, the sec14 mutant also exhibited a partial block in modification of Mnnlp. The sec1 mutant, which exhibits a block in secretory vesicle fusion with the plasma membrane, did not exhibit a defect in the modification of Mnnlp. The complete block in Mnnlp modification exhibited by the sec12 and sec18 mutants (Figs. 2 and 4) demonstrate a requirement for transport of Mnnlp to the Golgi complex for this modification to occur. The partial inhibition of this modification in two mutants that disrupt Golgi function, and the lack of an effect in the sec1 mutant, strongly suggest that the slow modification of Mnnlp was catalyzed within the Golgi complex. Taken together, these data suggest that Mnnlp is a resident of the yeast Golgi where the auto-mannosylation with αl,3 mannose is catalyzed over a prolonged period of time.

Turnover of Overexpressed Mnnlp

Kex2p has a half-life of ~80 min and is turned over in the vacuole by PEP4-dependent proteases. Overexpression of Kex2p leads to a two- to threefold increased rate of degradation in wild-type cells, and to an accumulation of Kex2p in the vacuole of pep4 mutant cells (63) which are pleiotropically deficient in the activities of the major vacuolar proteases (67). Kex1p and DPAP A also accumulate in the vacuole upon overexpression in pep4 cells (10, 48). We tested if overexpression would lead to mislocalization of Mnnlp and degradation in the vacuole. Wild-type cells with or without MNN1 on a multicopy plasmid were labeled and chased for 0, 1.5, or 3 h and then Mnnlp and ALP were immunoprecipitated. The level of overexpression from the multicopy plasmid ranged from 35–50-fold relative to Mnnlp expressed from a single copy gene in these experiments (see Fig. 1 A). While Mnnlp expressed from the genomic MNN1 gene was very stable (Fig. 1 B and Fig. 5, MNN1), overexpression led to a significant increase in the rate of degradation of this protein such that after three hours of chase ~80% of the labeled Mnnlp had been degraded (Fig. 5, 2μ MNN1). Even with the increased rate of degradation, there was still an 8–10-fold higher level of Mnnlp in the overexpressing strain relative to wild-type at the three hour time point. To assess the role of vacuolar proteases in degradation of overexpressed Mnnlp, the pulse chase analysis was also done in an isogenic pep4 strain. The degradation rate of overexpressed Mnnlp was nearly identical in the Pep4+ and Pep4− strains (Fig. 5, 2μ MNN1 vs. 2μ MNN1 pep4). Although the PEP4-dependent vacuolar proteases were not involved in the
Figure 5. Turnover of overexpressed Mnnlp in isogenic Pep4+ and Pep4− cells. Strains SEY6210 (MNN1), SEY6210 pZV236 (2µ MNN1) and TVY1 pZV236 (2µ MNN1 pep4) were labeled, chased, and subjected to immunoprecipitation as in Fig. 1 B. Bands on the autoradiograms were quantitated by densitometry as described in Materials and Methods and the recovery of Mnnlp was normalized to the recovery of ALP in the coimmunoprecipitates. ALP is subject to PEP4-dependent proteolytic processing and so the mobility of ALP in the gel also served as an internal control to confirm the genotype of the strains being used. The amount of Mnnlp in the immunoprecipitate at each time point was divided by the amount present at the 0 h chase point for each strain and expressed as the % initial Mnnlp.

Subcellular Localization of Mnnlp

Indirect immunofluorescent detection of overexpressed Mnnlp in wild-type diploid cells revealed a punctate staining pattern scattered throughout the cytoplasm, but excluded from the vacuole and the nucleus (Fig. 6). The number of distinct fluorescent spots ranged from 3 to 14 per cell in a given focal plane with the average being 7.2 (50 cells) in diploid cells overexpressing Mnnlp. The fluorescent signal was very faint from wild-type diploid cells, but the same punctate pattern was observed with an average of 5.1 distinct spots per cell (data not shown). This staining pattern is also seen for Kex2p, Kex1p, DPAP A, and Pmr1p, other integral membrane proteins thought to localize to the yeast Golgi complex (2, 10, 47, 49). We failed to see the ER, vacuole or plasma membrane stain with the anti-Mnnlp antibodies in cells overexpressing Mnnlp. The Amnnl strain showed a complete absence of staining demonstrating the specificity of the anti-Mnnlp antibody preparation (Fig. 6). When a haploid pep4 strain (TVY1) harboring the multicopy MNN1 plasmid was subject to immunofluorescence analysis, most of the cells did not exhibit vacuolar staining, but we did find a few cells (~1 in 500) that exhibited a brightly stained vacuole (data not shown). Based on the intensity of staining, this latter phenotype may represent a small population of cells in which the 2µ plasmid had replicated to very high copy number. Vacuo-
the p100 fraction as compared to the lysate (data not shown). Most of the ER, nuclei, vacuole membrane and plasma membrane are sedimented at 13,000 g (21, 25). The Golgi membranes collected on the sucrose cushion following the 100,000 g centrifugation step were loaded on top of a sucrose step gradient and centrifuged to equilibrium. We found three peaks of Mnnlp that migrated with sucrose densities of 1.11, 1.14, and 1.18 g/ml, respectively. GDPase exhibited the same distribution in these gradients, but membranes containing the Kex2 endopeptidase cofractionated with only the denser peaks of Mnnlp and GDPase (Fig. 7). These data suggest that the Mnnlp and GDPase are localized to at least two physically distinct compartments and that these enzymes may reside within the same, or at least very similar Golgi compartments. This is not surprising as α,3 mannosyltransferase activity is linked to GDPase function; GDPase is required to hydrolyze the GDP product of mannosyltransferase reactions to GMP which is thought to be exchanged for GDP-mannose by a specific cotransporter in the Golgi membrane. Strains harboring a null allele of the GDPase gene exhibit a defect in glycosylation (1) as well as GDP-mannose transport into Golgi vesicles (5). The two peaks in the more dense part of the gradient were not significantly different in the distribution of the three enzymes that we tested, so it is not clear if these peaks represent two distinct compartments or fragmented portions of the same compartment. We have also examined the distribution of Ochlp (an α,6 mannosyltransferase, reference 36) in these gradients and found that this protein was also present in the same membrane fractions as Mnnlp, but was somewhat more enriched in the lighter fraction (data not shown). These results suggest that the lighter membrane fraction that contained Mnnlp, GDPase and Ochlp, but lacked Kex2p, corresponds to the early Golgi compartments (cis and medial). From these experiments it appears that the late Golgi compartment that contains Kex2p may also contain a significant fraction of Mnnlp and GDPase.

Clathrin-dependent Localization of Mnnlp

The absence of clathrin function in a strain bearing a disruption of the clathrin heavy chain gene results in the mislocalization of Kex2 and DPAP A to the plasma membrane (44, 53). GDPase remains intracellular in the mutant cells suggesting that this protein is retained in the Golgi by a mechanism distinct from that employed by Kex2p and DPAP A (53). To test if Mnnlp is mislocalized to the plasma membrane when clathrin function is lost, a temperature-sensitive chcl mutant (chcl-ts) and an isogenic wild-type strain (CHC1) were shifted to 37°C for 0, 30, or 60 min, then subjected to cell surface iodination (Fig. 8 C) as previously described (52). Half of the cells treated at each temperature were lysed...
(Fig. 8 L) before iodination to provide an estimate of the total amount of Mnnlp present in these cells. After iodination, Mnnlp was recovered from the cells by immunoprecipitation and analyzed by SDS-PAGE (Fig. 8). In addition, a portion of each sample was also subject to immunoprecipitation with antiserum to GDPase to control for the integrity of the intact cells. We could not detect any Mnnlp on the cell surface of chcl-ts cells grown at the permissive temperature (Fig. 8, chc-ts O) or of wild-type cells incubated at 37°C for 1 h (Fig. 8, CHC 60). Yet, we found that Mnnlp appeared on the cell surface by 30 min after temperature shift and ~40% of the Mnnlp was mislocalized to the plasma membrane of the clathrin mutant after 60 min at 37°C. As previously reported (53), GDPase did not mislocalize to the cell surface in these experiments under any of the conditions tested (Fig 8, GDPase). In other experiments, as much as 80% of Mnnlp was mislocalized to the cell surface of the chcl-ts strain after 2 h at the nonpermissive temperature. The kinetics and extent of mislocalization of Mnnlp in the clathrin mutant is very similar to that previously described for Kex2p (44). In addition, Mnnlp was also mislocalized to the plasma membrane of a Δchcl strain to a similar extent as seen in the chcl-ts strain (data not shown).

Both Kex2p and DPAP A have large cytoplasmic tails with Golgi localization signals roughly defined as a (Y/F)X(Y/F)-XX(I/L) motif that is similar to clathrin coated pit localization signals found within the cytoplasmic tails of mammalian plasma membrane receptor proteins (reviewed in reference 40). Mutations in the cytoplasmic tails of Kex2p and DPAP A result in the mislocalization of these proteins to the vacuole (40). The Mnnlp cytoplasmic tail does not have an amino acid sequence that fits this motif well, although there is a FIL sequence that could possibly serve as this type of Golgi localization signal and mediate the clathrin-dependent Golgi localization of Mnnlp (Fig. 9 A). If this is the case, then removal of the Mnnlp cytoplasmic tail should result in the mislocalization of this protein to either the cell surface or vacuole. In order to maintain the type II topology of a Mnnlp cytoplasmic tail mutant, we exchanged the sequences that code for the Mnnlp 18-amino acid cytoplasmic tail for the six-amino acid cytoplasmic tail of dipeptidyl peptidase IV (gppl0, reference 24), a rat plasma membrane protein with the same topology as the Mnnlp, to produce pMNN1-XCT (Fig. 9 A). Other than the required initiator methionine, a threonine residue is the only common amino acid between these two cytoplasmic tails, and there are no tyrosines or phenylalanines in the dipeptidyl peptidase IV cytoplasmic tail.

As shown in Fig. 9 B by immunofluorescence, the Golgi retention of Mnnlp is not mediated through its cytoplasmic tail. Cells overexpressing this protein bearing the heterologous cytoplasmic tail exhibited a typical Golgi staining pattern (Fig. 9 B). We also found that the MNN1-XCT allele complemented a Δmnnl null allele when expressed from a single copy plasmid (data not shown). Moreover, the Mnnl-xct protein exhibited wild-type stability and slow posttranslational modification in pulse-chase experiments. We could find no differences in the activity, localization, stability or modification of the Mnnlp cytoplasmic tail mutant. These data argue that the Golgi localization of Mnnlp is not mediated by a direct interaction of clathrin/AP-1 complexes with the Mnnlp cytoplasmic tail.

**Discussion**

The Golgi complex plays a central role in the transport, modification and sorting of proteins in the secretory pathway. Towards a better understanding of the organization of this multicompartiment organelle and the mechanisms used to localize the intrinsic proteins of the Golgi complex, we have initiated an analysis of the yeast αl,3 mannosyltransferase. This work represents the first detailed characterization of the biosynthesis, modification and localization of a yeast mannosyltransferase. The predicted amino acid sequence of Mnnlp suggested that the protein would be a type II integral membrane protein (66). Consistent with this prediction, Mnnlp is resistant to extraction from membranes with alkaline carbonate buffer, but is readily extracted from the membrane with the detergent Triton X-100 indicating that it is tightly associated with the membrane. Proteinase K treatment of an intact membrane fraction indicates that the bulk of the Mnnlp is in the lumen of the Golgi, but also that this protein traverses the membrane such that a small cytoplasmic tail is susceptible to exogenously added protease (Fig. 4). These data indicate that the Mnnlp is an integral membrane protein, and combined with the hydrophathy analysis that shows a single NH2-terminal hydrophobic domain, suggests that this protein adopts a type II integral membrane topology.

Several lines of evidence demonstrate that Mnnlp is localized to the Golgi complex. (a) It has been shown previously that the modification catalyzed by the Mnnlp (αl,3 mannosyl addition to N- and O-linked oligosaccharides) is restricted...
The slow acquisition of $\alpha_1,3$ mannose epitopes on Mnnlp is surprising, as other yeast glycoproteins such as carboxypeptidase Y, proteinase A, invertase, and $\alpha$-factor, can be quantitatively immunoprecipitated with the antiseraum that has the appropriate concentration of substrates (ions, pH, inhibitors or activators) to allow this modification to occur. Consistent with this latter model, the Mnnlp appears to be localized to at least two distinct Golgi compartments, and may require recycling from the Kex2 compartment into the earlier compartment for auto-mannosylation. An $\alpha_1,3$ mannose residue is normally added onto an $\alpha_1,2$ linked mannose to extend the O-linked oligosaccharide chain. It is also possible that the addition of $\alpha_1,2$ mannose is limiting, and retrograde trafficking of Mnnlp into earlier Golgi compartments is required to form the appropriate substrate for auto-mannosylation.

We found that Mnnlp was very stable when expressed from a single copy gene, but showed a dramatic increase in the rate of degradation when overexpressed from a multicopy plasmid (Fig. 5). The rate of degradation of overexpressed Mnnlp was nearly the same in isogenic Pep4+ and Pep4- strains. The pep4 strain is pleiotropically deficient in the activities of the major vacuolar proteases; therefore, it is unlikely that the degradation of overexpressed Mnnlp was the result of mislocalization to the vacuole and degradation in this organelle. Moreover, immunofluorescence data indicated that the Mnnlp is not localized to the vacuole in wild-type or pep4 cells overexpressing this protein. At this time, we do not know where the degradation of overexpressed Mnnlp is occurring within the cell, but analysis of this event in sec mutants should define more precisely the organelle where the degradation is catalyzed.

Western blot analysis of Mnnlp suggested that the steady-state level of overexpressed Mnnlp in these strains is 10-15-fold higher than wild-type strains (data not shown). Therefore, the mechanism used to retain Mnnlp in the Golgi complex is apparently not saturated by a 10-15-fold increase in the amount of this protein. These results are in contrast to the findings that a significant fraction of Kexlp, Kex2p and DPAP A all mislocalize to the vacuole when overexpressed, where they are subsequently degraded in a PEP4-dependent manner (10, 49, 63). This result is more similar to mammalian glycosyltransferases, which upon overexpression are predominantly Golgi localized and do not appear to mislocalize to the plasma membrane or lysosome, although ER accumulation was sometimes observed (38). Overexpression of Mnnlp does not significantly affect the turnover of Kex2p, nor does overexpression of Kex2 affect the turnover of Mnnlp (Chen and Graham, unpublished data). These data suggest that overexpression of Mnnlp does not lead to general turnover of Golgi membrane, and that Kex2p and Mnnlp are not competing for a limiting component of a common retention apparatus.

We have found that 40-80% of Mnnlp was mislocalized to the cell surface in a temperature sensitive chcl-ts strain incubated at the nonpermissive temperature for 1-2 h (Fig. 8). Clathrin is also required for the efficient retention of Kex2p and DPAP A in the yeast Golgi complex (44, 53). It has been suggested that clathrin mediates the retention of Kex2p and DPAP A by a direct association of clathrin coats with the cytoplasmic tails of these Golgi proteins (64). The cytoplasmic tail of Mnnlp is very short (18 amino acids) relative to Kex2p (115 amino acids) or DPAP A (118 amino acids). Although there is a single aromatic amino acid in the cytoplasmic tail of Mnnlp (F7), it does not appear to be in the appropriate sequence context to fit the Golgi localization motif described for Kex2 and DPAP A (40). The cytoplasmic tail of Mnnlp is more similar to that of GDPase (1), which is predicted to have a nine-amino acid NH2-terminal cytoplasmic tail with...
two aromatic residues (F5 and Y8). Moreover, Mnnlp cofractionated with GDPase in sucrose gradients which suggests that these two proteins are localized to the same Golgi compartments. However, GDPase is not mislocalized to the plasma membrane in the clathrin mutant cells (Fig. 8, reference 53), and defects in outer chain mannosylation has not been observed in clathrin mutants (44). Therefore, we were surprised to find that a significant fraction of Mnnlp was mislocalized to the plasma membrane of clathrin disrupted cells, and that the extent of Mnnlp mislocalization in this mutant was very similar to what had previously been shown for Kex2p (44).

The finding that clathrin is required for Golgi localization of Mnnlp suggested that the Mnnlp cytoplasmic tail may contain a localization signal that mediates this retention, analogous to the signals within the Kex2p and DPAP A cytoplasmic tails. To test this, we replaced the Mnnlp cytoplasmic tail with that of dipeptidyl peptidase IV, a mammalian plasma membrane protein with a six-amino acid tail containing no tyrosine or phenylalanine residues (24). The mutant Mnnl-xct protein was expressed in a Δmnl strain and was found to be localized normally to the Golgi complex (Fig. 9 B). If clathrin mediates the retention of the Mnnlp in the Golgi by a direct interaction with the cytoplasmic tail, then deletion of the tail should result in the mislocalization of the Mnnlp to either the plasma membrane, as occurs in the clathrin mutant, or to the vacuole as occurs with cytoplasmic tail mutants of Kex2p and DPAP A. These results suggest that the clathrin requirement for Mnnlp Golgi localization is not mediated by a direct interaction of clathrin coats with the cytoplasmic tail of this protein. Moreover, this data argues for a different clathrin-dependent mechanism for the Golgi localization of the Mnnlp than that employed by Kex2p and DPAP A.

There are three models we can suggest to explain how clathrin is required for Mnnlp Golgi localization. (a) The entire Kex2p compartment may be lost to the plasma membrane upon disruption of clathrin function. Therefore, any protein contained in the Kex2p compartment would be mislocalized to the plasma membrane in the clathrin mutant. (b) Localization of the Mnnlp to the Golgi complex is mediated by a protein(s) that directly requires clathrin for Golgi localization by an interaction of the cytoplasmic tail(s) of this hypothetical protein with clathrin coat complexes. (c) Proteins required to maintain the appropriate balance of lipid and luminal constituents of the late Golgi are mislocalized in the clathrin mutant causing significant changes in the physical characteristics of this compartment and possibly earlier compartments as well. These changes would then indirectly disrupt the mechanisms used to localize the Mnnlp. The first model seems least likely because our data suggests that a fraction of GDPase was also contained within the Kex2p compartment, but was not mislocalized to the plasma membrane in the clathrin mutant. In addition, protein transport through the secretory pathway is relatively unaffected in the clathrin mutants which would be surprising if an entire Golgi compartment was lost. Others have reported a greater extent of separation of GDPase and Kex2 in sucrose gradients (6, 8), but we have found a peak of GDPase activity that cofractionated with Kex2p in each of 15 sucrose gradient fractionation experiments. Models 2 and 3 are difficult to discriminate at present; however, we have recently found that a fusion protein containing only the transmembrane domain of the Mnnlp can function to localize a reporter enzyme to the yeast Golgi complex (manuscript in preparation). Therefore, the mechanism for Golgi localization of Mnnlp might be similar to that used by several mammalian glycosyltransferases (see introduction). Current models used to explain the transmembrane domain-mediated retention of Golgi enzymes do not implicate specific interactions with a receptor, or tethering protein, but do require specific local environments of the membrane or compartment. Clathrin may be required for these proposed retention mechanisms by localizing enzymes required to maintain the appropriate lipid or luminal environment of the late Golgi (model 3). It is also possible that Mnnlp forms transmembrane domain mediated aggregates with late Golgi proteins (the kin recognition hypothesis) that are stabilized by a direct interaction of clathrin with the cytoplasmic tails of a subset of these Golgi proteins. Loss of clathrin might destabilize these compartment-specific aggregates resulting in the mislocalization of the late Golgi proteins, including Mnnlp, to the plasma membrane (model 2). Additional studies currently in progress are aimed at distinguishing between these possible models for the mechanism of Mnnlp retention within the appropriate Golgi compartment(s).

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