Characterization of TPM1 Disrupted Yeast Cells Indicates an Involvement of Tropomyosin in Directed Vesicular Transport

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Abstract. Disruption of the yeast tropomyosin gene TPM1 results in the apparent loss of actin cables from the cytoskeleton (Liu, H., and A. Bretscher. 1989. Cell. 57:233–242). Here we show that TPM1 disrupted cells grow slowly, show heterogeneity in cell size, have delocalized deposition of chitin, and mate poorly because of defects in both shmooing and cell fusion. The transit time of α-factor induced α-agglutinin secretion to the cell surface is longer than in isogenic wild-type strains, and some of the protein is mislocalized. Many of the TPM1-deleted cells contain abundant vesicles, similar in morphology to late secretory vesicles, but without an abnormal accumulation of intermediates in the delivery of either carboxypeptidase Y to the vacuole or invertase to the cell surface. Combinations of the TPM1 disruption with sec13 or sec18 mutations, which affect early steps in the secretory pathway, block vesicle accumulation, while combinations with secl, sec4 or sec6 mutations, which affect a late step in the secretory pathway, have no effect on the vesicle accumulation. The phenotype of the TPM1 disrupted cells is very similar to that of a conditional mutation in the MY02 gene, which encodes a myosin-like protein (Johnston, G. C., J. A. Prendergast, and R. A. Singer. 1991. J. Cell Biol. 113:539–551). The myo2-66 conditional mutation shows synthetic lethality with the TPM1 disruption, indicating that the MY02 and TPM1 gene products may be involved in the same, or parallel function. We conclude that tropomyosin, and by inference actin cables, may facilitate directed vesicular transport of components to the correct location on the cell surface.

Tropomyosin has been found in many eucaryotes, including mammals, insects (Karlick and Fyrberg, 1986), nematodes, and recently the slime mold Physarum and the yeasts Saccharomyces cerevisiae and Schizosaccharomyces pombe (Liu and Bretcher, 1989a). In all cases tropomyosin has been found associated with actin filaments, although its function is only really understood in skeletal muscle.

Skeletal muscle tropomyosin is a rod-shaped coiled-coil dimer assembled from highly α-helical monomers. In striated muscle, tropomyosin dimers bind along the actin filament in a head-to-tail fashion, with each molecule spanning seven actin monomers. Associated with each tropomyosin dimer is a troponin complex; together the two protein complexes confer Ca2+ sensitivity on the acto-myosin interaction (Ebashi et al., 1969). Distinct tropomyosin subunits are expressed in smooth muscle cells, where they may also be involved in thin filament regulation (Cummins and Perry, 1974). Nonmuscle cells display an amazing heterogeneity in expression of tropomyosin isoforms, which results mainly from alternative splicing of transcripts from a small number of genes (Leesmiller and Helfman, 1991). The diversity and tissue-specificity of tropomyosin isoforms expressed in higher eucaryotes suggests that this protein may play an important role in the utilization of microfilaments for divergent cell-type specific functions. However, how tropomyosin does this remains to be elucidated.

The yeast Saccharomyces cerevisiae is an attractive system for studying a primitive cytoskeleton (Huffaker et al., 1987; Barnes et al., 1990; Solomon, 1991; Huffaker and Bretscher, 1991). It contains a single essential actin gene, designated ACT1, that encodes a protein with 88% sequence identity to actins from higher cells (Shortle et al., 1982; Ng and Abel, 1980; Gallwitz and Sures, 1980). Actin shows a polarized distribution suggesting an involvement in oriented cell surface growth (Kilmartin and Adams, 1984; Adams and Pringle, 1984). At the beginning of the cell cycle, cortical actin patches with emanating cables are found at the site of bud emergence. As the bud grows, cortical patches are found enriched in the bud, and actin cables extend into the mother cell. Later in the cycle, cortical patches form around the bud neck and may be involved in cytokinesis. A number of studies indicate a role for the actin cytoskeleton in the determination of cell polarity, secretion, and localized surface growth. Novick and Botstein (1985) explored the phenotype of cells carrying the temperature sensitive actin alleles act1-1 and act1-2. At the restrictive temperature the cells lost actin cables, the distribution of actin patches were less polarized, chitin deposition was delocalized, and the cells accumulated...
both the secretory protein invertease and secretory vesicles. The phenotypic consequences of disruptions in genes encoding actin-binding proteins are beginning to be assessed. These include the genes for conventional myosin (encoded by MYO1; Watts et al., 1984, 1987), tropomyosin (TPMI; Liu and Bretscher, 1989a,b), profilin (PFY1; Magdolen et al., 1988; Haarer et al., 1990), capping protein (CAP1; Amatruda et al., 1990), fimbrin (SAC6; Adams and Botstein, 1989; Adams et al., 1989, 1991), and a 65-kD actin binding protein (ABP1; Drubin et al., 1988, 1990). Disruption of any of these genes is not lethal, but confers phenotypes of different severities. In preliminary studies of the TPM1, PFY1 and CAP1 gene disruptions, phenotypes similar to those of the conditional actin mutants were observed. In all cases, the actin cytoskeleton was altered, cells were larger and more spherical, and chitin deposition was abnormal, implicating the involvement of the actin cytoskeleton in cell polarity and cell wall synthesis. However, no detailed characterization of any of these mutants has appeared yet.

We have described the purification and characterization of a protein from Saccharomyces cerevisiae with properties characteristic of a tropomyosin. These properties, together with immunological cross-reaction to bovine brain tropomyosin and protein sequence and structural homology to tropomyosins from higher cells, provided strong evidence for classifying the yeast protein as a tropomyosin (Liu and Bretscher, 1989a,b). In yeast, tropomyosin is found associated with the actin cables that extend from the daughter bud into the mother cell; it is not found associated with the cortical actin patches that localize mostly in the daughter cell and at the bud neck during cytokinesis. Tropomyosin is encoded by a single gene in yeast that contains no introns. In an earlier report we showed that elimination of this tropomyosin from yeast by gene replacement did not confer lethality, but resulted in slower growing cells with no detectable actin cables (Liu and Bretscher, 1989b). Since cells from which tropomyosin had been genetically eliminated are viable, a phenotypic comparison between the TPM1 deleted and wild-type cells might uncover the function of tropomyosin in yeast.

Materials and Methods

Strains, Media, and Genetic Techniques

The genotypes of all yeast strains used in this paper are described in Table 1. The TPM1 gene was replaced by either the URA3 or LEU2 gene. The construction of the tpmlΔ::URA3 has been described (Liu and Bretscher, 1989b). For the LEU2 replacement, the URA3 containing DNA fragment from the HindIII to the Scal site in the tpmlΔ::URA3/YIp352 plasmid was replaced by the LEU2 gene of the HindIII-Scal fragment from a Ylp351 plasmid (Hill et al., 1986). This plasmid was linearized at the unique Scal site and transformed into diploid yeast cells (ABY365). This results in the replacement of the DNA from -236 to +225 at the TPM1 locus by LEU2. This replacement was confirmed by absence of detectable tropomyosin in Leu+ segregants as determined by immunoblotting with antibodies to yeast tropomyosin (Liu and Bretscher, 1989c). E. coli strain DH5α was used for all bacterial manipulations. Yeast media and genetic techniques were used as described by Sherman et al. (1974). The LiAc method was used for all the yeast transformations (Ito et al., 1983).

Mating Assay

Two methods were used to measure the efficiency of diploid formation (Trueheart et al., 1987). (i) Quantitative mass mating. Haploid cells were grown in YEPD (1% yeast extract, 2% peptone, 2% dextrose) medium at 30°C to no more than 5 × 10⁵ cells/ml 2 × 10⁶ α and α cells were mixed and filtered onto a sterile 25-mm HA filter (pore size 0.45 μm; Millipore Corp., Bedford, MA) (Duchett and Hartwell, 1983). The filter was transferred onto a YEPD plate without cells and incubated. After 4.5 h at 30°C, cells were spread on one side of a YEPD plate and yeasts were micromanipulated to a new area of the plate. Colonies arising from the yeasts were replica plated to test whether the yeasts had formed diploids or remained as haploids.

Invertase Assay

Cells were grown in YEP medium containing 5% glucose at room temperature to ~5 × 10⁶ cells/ml, washed, and resuspended into YEP medium containing 0.1% of glucose at 37°C. At each time point, 10 ml of the culture was removed, the cells were pelleted and washed in 10 mM NaN₃. The cell wall was removed by incubating the cells in 0.1 ml of 1.4 M sorbitol, 25 mM potassium phosphate (pH 7.5), 25 mM β-mercaptoethanol, 5 mM NaN₃, 25 μg/ml Zymolyase (ICN Immunobiologicals, Lisle, IL) for 1 h at 37°C. The spheroplasts were pelleted, resuspended in 0.1 ml of 25 mM Tris-phosphate (pH 6.7), 1 mM DTT, 1 mM EDTA, and lysed by vortexing with glass beads. Cell debris was pelleted at 15,000 g for 2 min. Proteins from the periplasm and the spheroplasts were fractionated on a 5.5% polyacrylamide native gel and the gel was stained for invertase activity (Gabriel and Wang, 1969).

Fluorescence Microscopy

To visualize DNA, cells were stained with 1 μg/ml 4',6'-diamidino-2-phenylindole (DAPI) (Duchett and Hartwell, 1983). Chitin was visualized after staining in 0.1% Calcofluor for 3 min and washing in distilled water (Slout and Pringle, 1978). Actin was stained with rhodamine-phalloidin (Adams and Pringle, 1984). To localize a-agglutinin, a-agglutinin antiserum (Watzke et al., 1988), generously provided by Dr. W. Tanner, was used after preabsorption on MAga cells. MAgα cells were grown in YEPD to ~2 × 10⁶ cells/ml at room temperature. β-Mercaptoethanol was added to the culture to final concentration of 1% and incubated for 1.5 h. This step reduces the background staining dramatically and also removes some constitutively expressed a-agglutinin from the cell wall seen in some strains. The cells were washed three times with YEP medium and resuspended in fresh YEPD (pH 4) medium to recover for 30 min at room temperature. α-Factor was added to the culture to 10 μg/ml and cells were incubated at room temperature and at 37°C. At each time point, aliquots of α-factor induced cells were fixed by adding formaldehyde to 3.7% and incubated for 1 h. The fixed cells were washed with 10 mM Tris-HCl, 140 mM NaCl, 5 mM EDTA, pH 7.5, and labeled with a-agglutinin antibodies and then FITC-conjugated second antibodies in the same saline solution (Watzke et al., 1988).

Electron Microscopy

Cells were prepared for thin section electron microscopy as described (Walworth and Novick, 1987) and viewed in an electron microscope (model EM300i; Philips Electronic Instrs. Co., Mahwah, NJ) operating at 80 kV.

Results

Disruption of the TPM1 Gene Reduces the Growth Rate and Alters Cell Morphology

Disruption of the TPM1 gene resulted in a reduced growth rate. The doubling time of haploids harboring the TPM1 gene disruption (tpmlΔ) at 30°C in rich media was ~140 min, whereas the doubling time for the corresponding TPM1+ cells is ~100 min.
haploids was 80 min. The slower growth of the tpmlΔ cells was not because of the death of a proportion of the cell population, since total cell and viable counts matched closely for both wild type and tpmlΔ cells (Fig. 1A). However, at elevated temperatures the tpmlΔ cells hardly grew and were temperature-sensitive, with ~20% cell survival after 6 h at 37°C (Fig. 1B). On plates, this temperature sensitivity was not very tight and varied with strain backgrounds. The tpmlΔ cells invariably reached stationary phase at a lower cell density than wild-type cells.

Disruption of the $TPM1$ gene is known to alter cell morphology and eliminate detectable actin cables from exponentially growing cells (Liu and Bretscher, 1989b). tpmlΔ cells were more spherical in shape and more heterogeneous in cell size than wild type controls, with many larger but also some smaller cells (Fig. 2, A and B). This phenotype was more severe in $tpmlA/tpmlA$ diploid cells. DNA staining revealed that most of the mutant cells contained only one nucleus like wild-type cells; however, multiple nuclei were observed in some of the larger cells (Fig. 2, C and D). In agreement with the nuclear staining pattern, data from DNA content analyses showed that 6% of the $tpmlA/tpml$ cells were tetraploid at 30°C and the percentage of tetraploids increased after shifting the cells to 37°C for 5 h (data not shown). A small percentage of aploid cells was also seen at 37°C. This result, together with the cell size heterogeneity, suggests that tpmlΔ cells are partially defective in polarized cell growth, giving rise to smaller than normal daughter cells and larger than normal mother cells.

$TPM1$ disrupted cells generate petites at a much higher frequency than wild-type cells. More than 50% of the cells in a tpmlΔ culture grown in YEPD that were derived from a single $p^+$ colony were $p^-$, whereas only ~10% of the iso-

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**Table 1. Yeast Strains Used in This Study**

| Strains     | Genotype                                      | Source             |
|-------------|-----------------------------------------------|--------------------|
| CUY25       | MATa ade2 his3-Δ200 leu2-3,112 ura3-52        | T. Huffaker       |
| CUY28       | MATa his3-Δ200 leu2-3,112 lys2-801 trp1-1 (am), ura3-52 | T. Huffaker       |
| CUY29       | MATa his3-Δ200 leu2-3,112 lys2-801 ura3-52 GAL+ | T. Huffaker       |
| ABY320      | MATa ade2 his3-Δ200 leu2-3,112 ura3-52 tpmlΔ::LEU2 | This study       |
| ABY321      | MATa his3-Δ200 leu2-3,112 lys2 ura3-52 tpmlΔ::LEU2 | This study       |
| ABY179      | MATa his4-539am ura3-52 tpmlΔ::URA3           | This study       |
| ABY365      | CUY25 × CUY28                                 | A. Adams          |
| ABY366      | ABY320 × ABY321                               | CSH*              |
| DBY2000     | MATa act1-3 ura3-52                           | P. Novick         |
| CGY339      | MATa pep4 his4-29 ura3-52 GAL*                | P. Novick         |
| NY3         | MATa ura3-52 secl-1                           | P. Novick         |
| NY10        | MATa ura3-52                                 | P. Novick         |
| NY13        | MATa ura3-52                                 | P. Novick         |
| NY17        | MATa ura3-52 sec6-4                           | P. Novick         |
| NY405       | MATa ura3-52 sec4-8                           | P. Novick         |
| NY414       | MATa ura3-52 sec13-1                          | P. Novick         |
| NY430       | MATa ura3-52 sec14-3                          | P. Novick         |
| NY431       | MATa ura3-52 sec18-1                          | P. Novick         |
| ABY409      | MATa ura3-52 secl tpmlΔ::URA3                 | This study       |
| ABY410      | MATa ura3-52 tpmlΔ::URA3                       | This study       |
| ABY411      | MATa ura3-52 tpmlΔ::URA3                       | This study       |
| ABY412      | MATa ura3-52 sec6-4 tpmlΔ::URA3                | This study       |
| ABY413      | MATa ura3-52 sec14-3 tpmlΔ::URA3               | This study       |
| ABY414      | ura3-52 sec4-3 tpmlΔ::URA3                     | This study       |
| ABY415      | ura3-52 sec13-1 tpmlΔ::URA3                    | This study       |
| ABY416      | ura3-52 sec18-1 tpmlΔ::URA3                    | This study       |
| JPTA        | MATa adel his3-Δ200 leu2-3,112 ura3-52 myo2-66 | G. Johnston       |
| JPTB        | MATa adel his3-Δ200 leu2-3,112 trp1-289 ura3-52 myo2-66 | G. Johnston       |
| ABY100      | MATa his3-Δ200 leu2-3,112 lys2-801 ura3-52 ade2 | This study       |
|             | tpmlΔ::LEU2 (TPM1 URA3)†                      |                    |

* Obtained by A. Bretscher at the Cold Spring Harbor Laboratory course in yeast genetics.  
† Genes listed in parentheses are carried on an autonomously replicating 2μ plasmid.

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Figure 1. Growth and viability of tpmlΔ of tpmlΔ cells. Haploid wild-type (CUY25) and tpmlΔ cells (ABY321) were grown in YEPD at 30°C (A) or after shifting to 37°C (B) and the cells/ml was determined directly by counting, and the viable counts were determined after plating aliquots on YEPD plates. Wild-type cell counts (○), viable counts (□); tpmlΔ cell counts (●), viable counts (●).
genic wild-type cells became $\rho^-$ under the same conditions. This caused heterogeneity in colony formation of tropomyosin deficient cells as colonies derived from a $\rho^+$ cell had an uneven edge, surface, and size, whereas those from $\rho^-$ cells were smaller, smooth, and uniform.

**Disruption of the TPM1 Gene Impairs Cell Mating**

Since $tpml\Delta$ cells lack actin cables and are more spherical than wild-type cells, actin cables may be responsible for the generation of the asymmetric shape. The most dramatic
shape change in yeast cells occurs during mating, when cells of opposite mating types respond to mating pheromones by shmooing. To assess the ability of wild type and \( \text{tpml} \Delta \) cells to shmoo in response to \( \alpha \)-factor, \( \text{MATa} \) cells were incubated with various concentrations of the mating pheromone and the induced shmoo formation was scored. The resulting dose-response curves (Fig. 3) show that \( \text{tpml} \Delta \) cells required \( \sim 5-10 \) times higher concentrations of \( \alpha \)-factor to shmoo than the wild-type cells. Upon treatment with \( \alpha \)-factor, \( \text{tpml} \Delta \) cells became larger and arrested in the unbudded portion (G1) of the cell cycle. However, even at \( \alpha \)-factor concentrations \( 10 \) times higher than required to induce shmoo in all wild-type cells, \( < 60 \% \) of the \( \text{tpml} \Delta \) cells had recognizable shmoo projections. In addition to this lower pheromone sensitivity, the \( \text{tpml} \Delta \) shmooing cells formed less pronounced, thicker projection tips (Fig. 3 B). Longer exposure of the cells to \( \alpha \)-factor gave similar results.

Cortical actin patches have been found concentrated at the shmoo tip of wild-type cells with actin cables extending into the body of the cell (Hasek et al., 1987). Actin cables are clearly evident in the shmoo neck of the wild-type cells, but were not seen in the \( \text{tpml} \Delta \) cells (Fig. 3, C and D). In wild-type cells incubated longer with the pheromone, actin cables disappeared from the first shmoo tip and were reorganized to form dots and cables at a second shmoo tip (Fig. 3 C). After an extended period in \( \alpha \)-factor, no actin cables and cortical patches were seen in the shmooing cells (not shown). This indicates that actin cables might be responsible for shmoo shape formation.

In addition to a defect in shmooing, \( \text{tpml} \Delta \) cells showed a partial defect in cellular fusion during zygote formation. When cells of opposite mating types were mated on rich media, most of the zygotes formed between mating \( \text{tpml} \Delta \) cells had a phase-dense plate between the mating partners, which was not seen in zygotes of isogenic wild-type cells (Fig. 4). The plate seen in the \( \text{tpml} \Delta \) mating pairs is similar to that of \( \text{fus}1 \text{fus}2 \) prezygotes (Trueheart et al., 1987) and DNA staining of the \( \text{tpml} \Delta \) prezygotes showed that the two nuclei had not fused. After extended mating (13 h) the nuclei in many of the \( \text{tpml} \Delta \) prezygotic pairs had migrated to either side of the plate but were unable to fuse. This suggests a partial defect in cell fusion, rather than nuclear migration. To measure the magnitude of this defect in fusion, individual zygotes were picked by micromanipulation and the percentage giving rise to diploids was determined (Table II). About half of the zygotes from matings between \( \text{tpml} \Delta \) cells failed to form diploids, and also a higher percentage failed to form viable colonies. These results suggest that the mating pairs derived from \( \text{tpml} \Delta \) cells fail to fuse about half the time.

The combined defects in shmooing and cellular fusion lead to a much lower frequency of diploid formation. This was assayed in mass matings on nitrocellulose filters (Table III). In relation to the frequencies observed for matings between wild-type cells, crosses between \( \text{tpml} \Delta \) and \( \text{TPM}^{+} \) cells reduced the formation of diploids by \( 80 \% \), and between \( \text{tpml} \Delta \) and \( \text{tpml} \Delta \) cells by \( 98 \% \). In addition, \( \text{tpml} \Delta /\text{tpml} \Delta \) diploids that were able to form, sporulate very poorly.

**Disruption of the TPM1 Gene Causes Delocalized Chitin Deposition**

A ring of chitin is formed at the neck of a budding cell and remains on the mother as a bud scar after cell division. In haploid cells the site of bud emergence is adjacent to the previous bud emergence site. Chitin rings provide a record of the site of earlier budding cycles. Strains carrying condi-
Figure 4. Morphology (A and B) and DNA localization (C and D) of conjugating cells. (A and C) TPM/+ × TPM/+ (CUY28 × CUY25); (B and D) tpmlΔ × tpmlΔ (ABY320 × ABY321) after 4.5 h at 30°C. Cells were photographed under Nomarski optics (A and B) or after staining with DAPI (C and D). Arrows indicate the phase dense plate between mating partners. Bar, 15 μm.

Table II. Diploid Formation in Micromanipulated Zygotes

| Strains crossed     | Diploids formed* |
|---------------------|------------------|
| a TPM1 (CUY25) × α TPM1 (CUY28) | 100 (39)         |
| a TPM1 (CUY29) × α tpmlΔ (ABY320) | 91 (36)          |
| a tpmlΔ (ABY321) × α tpmlΔ (ABY320) | 46 (27)          |

* The percentage of diploids among the colonies that grew from the micromanipulated zygotes (the actual number of the colonies that grew is shown in parentheses). In each cross, 48 zygotes were picked after cells of opposite mating types were mixed for 4.5 h at 30°C.

Localization of chitin in tpmlΔ cells at room temperature and at 37°C revealed similarities to the actin conditional mutants (Fig. 5). At room temperature, diffuse chitin rings and generalized overall staining was seen in tpmlΔ cells. The staining was quite heterogeneous and seemed to correlate

Table III. Efficiency of Diploid Formation in Crosses with tpmlΔ Cells

| Strains crossed                  | Diploids formed |
|----------------------------------|-----------------|
| a TPM1 (CUY25) × α TPM1 (CUY28) | 49              |
| a tpmlΔ (ABY179) × α TPM1 (CUY28) | 10              |
| a tpmlΔ (ABY321) × α tpmlΔ (ABY320) | 0.6             |

The numbers are the percentage of diploids formed after 6 h at 30°C.
with cell size, with larger cells having more delocalized staining. At 37°C, the staining was bright and mostly delocalized.

**Disruption of the TPM1 Gene Affects Polarized Secretion of α-Agglutinin**

Secretion and deposition of newly synthesized cell wall components in wild-type cells is directed to the growing buds in a process suggested to involve actin cables (Kilmartin and Adams, 1984; Adams and Pringle, 1984). Since the mechanism of chitin ring formation is not known, chitin deposition cannot be used to probe the behavior of the secretory and targeting pathways. A method of negatively staining with FITC-concanavalin A was used to localize the area of newly synthesized mannoproteins (Adams and Pringle, 1984). In this method, cells are labeled with FITC-ConA, chased in unlabeled growth medium for 30 min, and the unlabeled halo is used as an indication of the site of new cell wall deposition. Using this assay, bud growth in *tpmlA* cells seemed normal. However, since this method does not allow the detection of newly synthesized materials that are incorporated into pre-existing cell walls, it can not reveal a partial defect in targeting. To detect any defects in localized secretion, the appearance of α-agglutinin at the cell surface was studied.

α-Agglutinin is an α-factor inducible cell surface glycoprotein in *MATa* cells. Upon α-factor treatment of exponentially growing *MATa* cells, the protein is first secreted exclusively to the growing bud and later to the tip of the pear-shaped shmoo (Watzele et al., 1988). α-Agglutinin was localized by immunofluorescence microscopy in *tpmlA* and wild-type cells at various times after α-factor addition (Fig. 6). At room temperature, the localization was similar in the two strains with α-agglutinin appearing at the cell surface 30 min after induction and concentrated in the buds (Fig. 6, A and B). At 180 min the staining was observed at the tips of the shmooing cells. However, the staining pattern of wild type and *tpmlA* cells at 37°C was different. α-Agglutinin was seen on the cell surface of wild type but not *tpmlA* cells at 30 min, and scattered patches appeared on the surface of the mother *tpmlA* cells at 60 and 90 min after induction, although staining in most cells was concentrated in the buds.

**Disruption of the TPM1 Gene Does Not Block Secretion**

The delayed appearance and partial delocalized incorporation of α-agglutinin on the cell surface could indicate an impairment in the secretory pathway. The status of the secretory pathway was assessed by examining the secretion of the periplasmic enzyme invertase and the delivery of a vacuolar enzyme carboxypeptidase Y (CPY).

Invertase is the product of the *SUC2* gene, which encodes
Figure 7. Invertase secretion and carboxypeptidase Y (CPY) delivery to the vacuole are normal in tpm1Δ cells. (A) Internal and external invertase. Wild-type (CUY25), tpm1Δ (ABY321) and act1-3 (DBY2000) cells were grown to early log phase at room temperature and switched from 5% glucose to 0.1% glucose containing media to induce invertase secretion. Cytoplasmic and periplasmic invertase were assayed after fractionation on a native gel after 0, 15, 30, 45, and 60 min induction at 37°C. The fully glycosylated secretory form and the constitutively expressed nonglycosylated cytoplasmic form of invertase are indicated. (B) Delivery of CPY to the vacuole is normal in tpm1Δ cells. Wild-type (CUY25), tpm1Δ (ABY321), pep4 (CGY339), and sec18Δ (NY431) cells were grown at room temperature and the pep4 and sec18Δ cells shifted to 37°C for 2 h. All cells were grown in YEPD + 0.5% BSA to prevent proteolytic cleavage of the secreted CPY (Stevens et al., 1986). Total cell extracts and total proteins in the media were subjected to electrophoresis and immunoblotted with CPY antibodies. (Lane 1) Wild type cells; (lane 2) tpm1Δ cells; (lane 3) pep4 cells; (lane 4) sec18Δ cells. Lanes 1'-4' are the media from the corresponding cells. The precursor (p) and mature (m) forms of CPY are indicated.

Figure 6. Localization of a-agglutinin at the cell surface at various times after α-factor induction. Wild-type (CUY25) and tpm1Δ (ABY321) cells were grown to early log phase at room temperature and exposed to α-factor for the indicated times at both room temperature and 37°C. Cells were then fixed and stained for a-agglutinin by indirect immunofluorescence microscopy. (A) Wild type at room temperature; (B) tpm1Δ cells at room temperature; (C) wild-type cells at 37°C; (D) tpm1Δ cells at 37°C. Arrows indicate shmoo tips. Bar, 15 μm.

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Figure 8. Electron micrographs of thin sections of wild-type (A) and \( \text{tpml} \Delta \) cells (B and C). Wild-type (NY13) and \( \text{tpml} \Delta \) cells (ABY411) were grown at room temperature and processed for EM. Bars, 1.5 \( \mu \text{m} \).

It is possible that the secretion defect in the \( \text{tpml} \Delta \) cells is minor and >1 h is necessary to result in any detectable accumulation. To explore this, cells were grown up in rich medium containing sucrose as carbon source for 20 h at 30°C and the internal and external levels of invertase were determined. Again, no greater accumulation of invertase could be detected in \( \text{tpml} \Delta \) cells than in wild-type cells. Cells carrying conditional mutations in the \( \text{SEC4}, \text{SEC14}, \text{or SEC18} \) genes accumulated invertase after shifting to 37°C for 2 h (data not shown). These combined data imply that invertase secretion is not significantly impaired in \( \text{tpml} \Delta \) cells.

CPY is an enzyme whose delivery to the vacuole can be monitored by changes in the apparent molecular weight of its precursors (Stevens et al., 1982). CPY is partially glycosylated in the ER to give a precursor designated pl, then fully glycosylated in the Golgi apparatus to give p2, and finally proteolytically cleaved in the vacuole by the \( \text{PEP4} \) gene product to yield mature CPY. To explore whether CPY is properly delivered to the vacuole in \( \text{tpml} \Delta \) cells, blots of total proteins from wild-type cells, \( \text{tpml} \Delta, \text{sec18}, \text{and pep4} \) cells were probed with antibody to CPY (Fig. 7B). The CPY in the \( \text{tpml} \Delta \) cells is essentially all in the mature form, indicating a normal vacuolar protein sorting pathway and a functional vacuole. Mutants defective in vacuolar protein sorting are unable to deliver CPY to the vacuole and, therefore, secrete it into the medium in the p2 form (Rothman and Stevens, 1986). A small amount of mature CPY is found in the medium of \( \text{tpml} \Delta \) cells suggesting that this fully processed product is released into the medium probably by cell lysis because actin could also be detected by immunoblotting in the medium (data not shown).

Disruption of the \( \text{TPM1} \) Gene Results in the Accumulation of Vesicles

Electron micrographs of thin section of \( \text{tpml} \Delta \) cells revealed that they contained an abnormally large number of vesicles (Fig. 8). These resemble the secretory vesicles that accumulate in yeast cells conditionally defective in a late step of the secretory pathway (Novick et al., 1980). As with its other phenotypes, the accumulation was heterogeneous and varied with genetic background and growth conditions. Between 20 and 48% of haploid \( \text{tpml} \Delta \) cells accumulated vesicles at room temperature and a greater percentage accumulated them when the cells were shifted to 37°C for 2 h. Diploid \( \text{tpml} \Delta \) cells had a more severe phenotype, with ~60% of the cells grown at 30°C showing accumulated vesicles. Accumulation seemed to be correlated with cell size, but since serial sections have not been collected, a small cell in a thin section is not necessarily a small cell in reality. However, very large \( \text{tpml} \Delta \) cells were packed with vesicles. The accumulation was not caused by the property of \( \text{tpml} \Delta \) cells to yield \( \rho^- \) progenies at high frequency, as a similar fraction of \( \rho^- \text{tpml} \Delta \) cells contained vesicles.

The accumulation of vesicles morphologically similar to late secretory vesicles appears to contradict the finding that \( \text{tpml} \Delta \) cells do not accumulate internal invertase. To determine whether the vesicles are in fact intermediates in the secretory pathway, the membrane traffic of cells carrying both the \( \text{tpml} \Delta \) disruption and conditional mutations in various \( \text{SEC} \) genes was examined. The \( \text{SEC} \) mutants have been classified into three major groups according to the membrane bound compartments that they accumulate at the restrictive temperature (Novick et al., 1980). Mutations in early genes (such as \( \text{SEC13}, \text{SEC18} \)) accumulate an exaggerated ER; mutations in genes involved in transport from the Golgi apparatus to the vacuole and to secretory vesicles (such as \( \text{SEC14} \)) develop an enlarged Golgi apparatus; and mutations in genes necessary for the exocytosis of secretory vesicles (such as \( \text{SEC1}, \text{SEC4}, \text{and SEC6} \)) accumulate secretory vesicles. If the \( \text{TPM1} \) gene product is involved in one or more steps in the secretory pathway, disruption of the \( \text{TPM1} \) gene in combination with a conditional \( \text{sec} \) mutation necessary for that step might be lethal. Such synthetic lethality between \( \text{sec} \) mutations has been very informative at identifying genes specifying interacting components of a particular step in the secretory pathway (for example see Salminen and Novick, 1987; Kaiser and Schekman, 1990). In addi-
Table IV. Organelle Accumulation in tpm1Δ sec Double Mutants

| Relevant genotype (strain) | Accumulation at 23°C | Accumulation at 37°C* |
|---------------------------|----------------------|-----------------------|
| Wild type (NY13)          | None                 | None                  |
| tpm1Δ (ABY411)            | None                 | None                  |
| tpm1Δ, sec13 (ABY415)     | Vesicles in 48% cells| ER                    |
| tpm1Δ, sec18 (ABY416)     | None                 | ER                    |
| tpm1Δ, sec14 (ABY413)     | Vesicles in 14% cells| Golgi apparatus and some vesicles |
| tpm1Δ, sec1 (ABY409)      | Vesicles in 49% cells| Vesicles in all cells  |
| tpm1Δ, sec4 (ABY414)      | Vesicles in 30% cells| Vesicles in all cells  |
| tpm1Δ, sec6 (ABY412)      | Vesicles in 40% cells| Vesicles in all cells  |

* Cells were shifted to 37°C for 2 h.

tion, if the vesicles seen in tpm1Δ cells are late secretory vesicles, a block at an early step of the pathway might be expected to reduce the accumulation of the late secretory vesicles. To explore double mutant combinations, the TPM1 gene was disrupted in strains carrying the conditional mutations sec1-1, sec4-8, sec6-4, sec13-1, sec14-3, and sec18-1. No synthetic lethality was found among any of the double mutation combinations. In the tpm1Δ strains also harboring sec1, sec4, or sec6 mutations, which affect a late step in the pathway, 30–50% of the cells accumulated vesicles when grown at room temperature (Table IV). Highly packed vesicles were seen in all the cells after shifting to 37°C due to the phenotypic expression of the conditional sec mutation (Fig. 9). The vesicles were indistinguishable from those accumulated at room temperature. In tpm1Δ, sec14-3 cells, vesicles were found in ~14% of the cell population after growth at room temperature (Table IV and Fig. 9). This double mutant accumulated both Golgi apparatus and vesicles in most of the cells at 37°C. Since TPM1+sec14-3 cells accumulate both Golgi apparatus and vesicles at 37°C, it is not clear if the vesicles seen in the tpm1Δ, sec14-3 construct are a consequence of the sec14-3 mutation or the tpm1Δ disruption. In contrast to the mutations affecting late steps in the secretory pathway, vesicles were rarely seen in the tpm1Δ sec13-1 or tpm1Δ sec18-1 mutants grown at room temperature (Table IV, Fig. 9). Our interpretation is that at the permissive temperature the mutations in these SEC genes reduce the flux through the pathway and so the defect that leads to late secretory vesicle accumulation is not seen. This supports the idea that the vesicles that accumulate in tpm1Δ are vesicular intermediates in the late secretory pathway.

Although the sec13 and sec18 mutations were able to phenotypically suppress the accumulation of vesicles seen in tpm1Δ cells, they did not suppress either the slower growth rate or the heterogeneous cell size (not shown).

Tpm1Δ and Myo2-66 Mutants Have Many Properties in Common and the Mutations Show Synthetic Lethality

Recently, Johnson et al. (1991) described the properties of a temperature-sensitive mutation, designated myo2-66, in a gene that encodes a myosin-like protein. The myo2-66 mutants are in many ways similar to the tpm1Δ strain described here. This mutant was isolated from a screen after enriching for abnormally large cells (Prendergast et al., 1990) and arrests as unbudded cells at the restrictive temperature. At the restrictive temperature the myo2-66 mutant has an aberrant actin cytoskeleton, has delocalized chitin deposition, and accumulates vesicles yet secretes invertase normally.

We have examined myo2-66 mutant cells and found that they do not accumulate more invertase at their restrictive temperature than wild-type cells when grown in sucrose as the carbon source (data not shown). The α-agglutinin secretion pattern in the myo2-66 cells was also similar to that in the tpm1Δ cells, normal at room temperature and slower and partially delocalized at 37°C (data not shown). These similarities, together with the fact that tropomyosin and myosin are both important components of the contractile machinery of higher cells, prompted us to explore the phenotype of tpm1Δ myo2-66 double mutants. A tpm1Δ::URA3 strain was crossed to a myo2-66Δ strain. Sporulation of the resulting diploid gave no complete tetrads of four viable spores. Analysis of viable spores showed that no tpm1Δ::URA3 myo2-66Δ haploids were recovered in 12 sets of tetrads, suggesting that tpm1Δ myo2-66 might be lethal. However, an excess lethality of tpm1Δ cells unrelated to myo2-66 seemed to exist. To show the synthetic lethality more definitively, a tpm1Δ::LEU2/TPM1 myo2-66Δ MYO2+ diploid was transformed with a TPM1/2μ(URA3) plasmid before sporulation. Four viable spores were now recovered, with the temperature sensitivity (myo2-66Δ) segregating 2:2 and the Leu+ (tpm1Δ) segregating 2:2. The segregants were placed on FOA plates at room temperature to select for Ura- cells that had lost the plasmid. Only the tpm1Δ::LEU2 myo2-66Δ segregants were unable to loose the TPM1/2μ(URA3) plasmid. Since the myo2-66 mutant grows well under these conditions, we conclude that the tpm1Δ myo2-66 double mutant is inviable at room temperature.

Discussion

Tropomyosin is an important microfilament-associated protein of cells from mammals to yeast, yet outside of striated muscle its function is not clearly understood. The availability of yeast mutants that lack tropomyosin allows an assessment of its function in at least this organism. We have shown that cells in which the TPM1 gene was functionally deleted had no detectable actin cables. This, and the finding of an association of tropomyosin with actin filaments in vitro and in vivo, led to the idea that tropomyosin is involved in the assembly or stabilization of actin cables in yeast (Liu and Bretscher, 1989b).

Cells lacking a functional TPM1 gene grow remarkably well despite their aberrant actin cytoskeleton. The lack of
tropomyosin imparts considerable heterogeneity into the population, with the larger cells having a more abnormal phenotype than the smaller cells, making a phenotypic characterization all the more difficult. Despite this difficulty, there appears to be a common thread that links all the results: cells lacking tropomyosin have a partial defect in the directed transport of components to their correct destination on the cell surface. The fact that the abnormal phenotype is more severe in larger cells, which are still viable, argues strongly for a partial defect in some form of intracellular transport.

In exponentially growing wild-type cells, all the growth is directed to the bud and the mother cell maintains a constant size during the cell cycle. In tpm/A cells, the growth is less directed and both the mother and daughter cells grow. This results in smaller than normal daughter cells and larger than normal mother cells. The defect in directed transport appears to be more pronounced in tpm/A diploids than haploids, presumably because their larger size makes the greater distance that materials have to be transported more critical.

Abnormalities in chitin deposition in tpm/A cells also seem to be size dependent, with smaller cells having a relatively normal staining pattern, and larger cells having more delocalized staining. The phenotype is more severe at 37°C with most of the cells having completely delocalized chitin. However, since the mechanism of chitin deposition is unknown, it is hard to evaluate the reason for chitin mislocalization in the tpm/A cells.

Perhaps the clearest physiological demonstration that the tpm/A cells are partially defective in directed secretion are our results examining the transit time and localization of a-agglutinin at 37°C. The transit time for a-agglutinin to reach the cell surface after induction is at least two times longer than in isogenic wild-type cells, and when it does come to the surface, some of it is inappropriately localized. However, the targeting of a-agglutinin looks perfectly normal at room temperature and most of it is localized in the bud even at 37°C. This mislocalization phenotype is not as strong as expected if the directed transport of secretory vesicles, presumably along actin cables, is the only mechanism for polarized cell surface growth. It is therefore very likely that there are also proteins specifically localized at sites of active surface growth that facilitate the fusion of the secretory vesicles with the plasma membrane. The absence of these proteins in the mother cell would make exocytosis inefficient and be expected to lead to the accumulation of vesicles in the mother cell. Conditional mutations affecting targeting proteins of this type might confer a defect in bud formation, and arrest as unusually large unbudded cells at the restrictive temperature, such as is found for the CDC24, CDC42, CDC43, and BEM1 gene (Sloot et al., 1981; Adams et al., 1990; Bender and Pringle, 1991). The targeting of the secretory protein acid phosphatase in cdc24 mutant cells has been shown to be completely delocalized (Field and Schekman, 1980). This same problem appears to afflict the mating response of tpm/A cells. When wild-type cells of opposite mating types are mixed, cells grow towards each other forming shmoo projections. Agglutinin is targeted to the shmooing tip that sticks cells of opposite mating types together. During this process, actin cables are oriented towards the shmoo tip. If this morphological change does not occur correctly, as in the case of tpm/A cells which do not assemble actin cables directed to the tip, the cells cannot conjugate efficiently. It is not yet clear why tpm/A cells require more mating pheromone to induce shmoo formation; they seem to be about as sensitive as wild-type cells to a-factor induced growth inhibition as determined by a halo assay. In the next step of zygote formation, removal of the cell wall between the two conjugating cells involves the correct targeting of the cell wall lysis and fusogenic materials; this too seems to be partially defective in tpm/A cells.

Electron microscopy reveals that many of the cells in an exponentially growing population of the tpm/A mutant have an abnormal accumulation of vesicles, with a greater percentage of diploids than haploids showing this phenotype. These are reminiscent of the vesicles that accumulate in mutants blocked late in the secretory pathway. We have results to suggest that these vesicles are in the late secretory pathway. A combination of the tpm/A mutation with conditional mutations in genes required for late steps in the secretory pathway have an “additive” phenotype. That is, at the permissive temperature the cells are viable but many of them accumulate vesicles due to the tpm/A mutation. At the restrictive temperature all the cells accumulate vesicles due to both mutations. In contrast, combinations of the tpm/A mutation with either sec13-1 or sec18-1 show the phenotype of the sec mutant. At the permissive temperature, the cells are free of internal vesicles, whereas at the restrictive temperature they show the phenotype typical of their conditional mutation. These two genes have their earliest point of action in the secretory pathway between the endoplasmic reticulum and the Golgi apparatus (Kaiser and Schekman, 1990). Conditional mutations in these genes will presumably slow the flux through the secretory pathway even at the permissive temperature. Since they suppress the formation of vesicles by the tpm/A mutation, they presumably lie upstream of the partial defect imposed by the loss of tropomyosin. This places the site of action sensitive to the lack of tropomyosin as post-Golgi apparatus.

The phenotype of cells carrying the myo2-66 mutation is remarkably similar to tpm/A cells. Moreover, the myo2-66 and tpm/A mutations show synthetic lethality, strongly implicating their wild-type products as two interacting components of an important function. Very recently, Govindan et al. (1991) reported that myo2-66 shows synthetic lethality with mutations in a subset of late acting SEC genes (sec2, sec4, sec5, sec8, sec9, sec10 and sec15) and that duplicating SEC4 can partially suppress the phenotype of the myo2-66 mutant. This seems to imply that the MYO2 gene product

Figure 9. Ultrastructure of stains carrying both tpm/A and a sec mutation. Double mutants were grown at room temperature to ~5 x 10⁶ cells/ml and half the culture shifted to 37°C for 2 h. Cells were then processed for the preparation of thin section and examination by electron microscopy. A, C, E, and G are cells grown at room temperature. B, D, F, and H are cells after 2 h at 37°C. (A and B) sec13tpm/A cells (ABY415); (C and D) sec18tpm/A cells (ABY 416); (E and F) sec44tpm/A cells (ABY413); and (G and H) sec1tpm/A cells (ABY409). Bar, 1.5 μm.
acts in the late secretory pathway. The results presented here also imply that the TPM1 gene product may participate in this part of the pathway.

Despite this evidence for a defect in the late secretion pathway, there is no abnormally large accumulation of invertase found in either tpmlΔ or myo2-66 cells. If the products of the MYO2 and TPM1 genes are involved in vesicle transport and defects in them lead to the accumulation of secretory vesicles, biochemical intermediates in the pathway should also accumulate. Indeed, given the number of vesicles seen, particularly in the tpmlΔ cells, a very significant accumulation should have been found, yet none was. This paradox is not easy to resolve, and we have no completely satisfactory explanation. One possibility is that secretory vesicles only accumulate slowly and if they do not fuse with their target membrane their contents become degraded. Another possibility is that MYO2, TPM1, and some of the late SEC gene products are necessary for a vesicular transport step in another pathway and that the accumulated vesicles are intermediates in this pathway. For example, there may be two parallel secretory pathways from the Golgi apparatus to the plasma membrane, both requiring the late sec genes, but only one requiring the MYO2 and TPM1 gene products: this would be the route not carrying invertase. Alternatively, since many of the late, and some of the early, SEC genes are involved in endocytosis (Riezman, 1985), the accumulated vesicles may be intermediates in this pathway, although preliminary uptake studies with lucifer yellow CH do not support this possibility. Characterization of the vesicles that accumulate in tpmlΔ cells will help resolve this puzzle.

The results presented in this paper support a role for actin cables in vesicular transport and cell wall growth. This suggestion was first put forth by Kilmartin and Adams (1984), and Adams and Pringle (1984). It has consistently been supported by observations on the analysis of genes for many actin binding proteins (see introduction) and the analysis of the myo2-66 mutation described above. Support has also come from another direction. Novick et al. (1989) isolated allele-specific suppressors of the actl-1 mutation that led to the identification of five genes (designated SAC1 to SAC5) that may encode actin-binding proteins. Suppressors of a mutation in the SECl4 gene, which is necessary for transport from the Golgi to both the vacuole and secretory vesicles (Novick et al., 1981) and encodes a phospholipid transfer protein (Barkaitis et al., 1990), were also found to lie in the SAC1 gene (Cleves et al., 1989). The sacl alleles that suppressed the secl4 mutations also suppressed the actl-1 mutation. Additionally, the sacl mutations could also partially suppress some sec6 and sec9 mutations, and showed synthetic lethality with secl3 and sec20 mutations. The simple view of these studies is that the sacl mutations can suppress mutations in genes needed after the Golgi apparatus in the secretory pathway, whereas the same sacl mutations aggravate defects early in the pathway (ER to Golgi apparatus). Whatever the mechanism, these results again provide genetic evidence for a functional link between the actin cytoskeleton and the secretory pathway.

Despite these clear links to the secretory pathway, it should be noted that cells harboring mutations in actin-binding proteins have a more complicated phenotype than sec mutants. It appears that mutants such as tpmlΔ cells are also partially defective in delivering the secretory products to their correct destinations, which leads to a defect in morphogenesis, and may be defective in other functions yet to be appreciated. Only when appropriate mutations that affect specific aspects of microfilament function are available will it really become possible to pinpoint the precise function of microfilaments in vital processes.

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