

Clathrin Functions in the Absence of the Terminal Domain Binding Site for Adaptor-associated Clathrin-Box Motifs

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Clathrin is involved in vesicle formation in the *trans*-Golgi network (TGN)/endosomal system and during endocytosis. Clathrin recruitment to membranes is mediated by the clathrin heavy chain (HC) N-terminal domain (TD), which forms a seven-bladed β -propeller. TD binds membrane-associated adaptors, which have short peptide motifs, either the clathrin-box (CBM) and/or the W-box; however, the importance of the TD binding sites for these motifs has not been tested *in vivo*. We investigated the importance of the TD in clathrin function by generating 1) mutations in the yeast HC gene (*CHC1*) to disrupt the binding sites for the CBM and W-box (*chc1-box*), and 2) four TD-specific temperature-sensitive alleles of *CHC1*. We found that TD is important for the retention of resident TGN enzymes and endocytosis of α -factor; however, the known adaptor binding sites are not necessary, because *chc1-box* caused little to no effect on trafficking pathways involving clathrin. The *Chc1*-box TD was able to interact with the endocytic adaptor Ent2 in a CBM-dependent manner, and HCs encoded by *chc1-box* formed clathrin-coated vesicles. These data suggest that additional or alternative binding sites exist on the TD propeller to help facilitate the recruitment of clathrin to sites of vesicle formation.

INTRODUCTION

The efficient delivery of proteins and lipids to specific cellular compartments of the secretory and endocytic pathways is achieved by transport in membrane-bound vesicles. Selective recruitment of cargo and formation of vesicles is mediated by coat proteins. The coat protein clathrin is present in all eukaryotic cells and is an important component in the membrane trafficking events of the *trans*-Golgi network (TGN)/endosomal system and in endocytosis at the plasma membrane (PM) (reviewed in Brodsky *et al.*, 2001).

The assembly unit of the clathrin coat is the triskelion, which is composed of three 190-kDa clathrin heavy chains (HCs) and three 25- to 28-kDa clathrin light chains (LCs). Clathrin HCs trimerize at their C-termini, and each leg ex-

tends outward radially from a central vertex to generate distinct regions, including proximal and distal legs, a linker region, and finally a globular N-terminal domain (TD). A single LC noncovalently associates with each HC along the proximal leg region near the vertex, and proximal and distal regions from neighboring triskelions interdigitate to form the polygonal lattices characteristic of clathrin-coated membrane structures, with the TD facing inward toward the membrane surface (Fotin *et al.*, 2004).

Clathrin assembly at sites of vesicle formation requires adaptor proteins to sort protein cargo into developing vesicles and additional assembly factors to facilitate lattice formation. Several adaptor proteins exist in eukaryotic cells, including the heterotetrameric adaptor protein complexes (APs), monomeric adaptor proteins (GGAs, epsins, and AP180s), and cargo-specific adaptor proteins. APs and monomeric adaptors have cargo-binding, phospholipid-binding, and clathrin-binding capabilities to help facilitate clathrin recruitment and coated vesicle formation. In addition, many accessory proteins also bind clathrin directly to facilitate lattice assembly (reviewed in Owen *et al.*, 2004).

Adaptor proteins and assembly factors bind to the TD regions of clathrin triskelions via short, linear peptide sequences. The first consensus clathrin-binding sequence (SLLDLD) was identified in studies of the β 3a subunit of the heterotetrameric adaptor complex AP-3 (Dell'Angelica *et al.*, 1998), and subsequent alignment of sequences from the clathrin-binding regions of numerous adaptor proteins revealed the presence of a consensus clathrin-binding motif (L Φ p Φ p) called the clathrin-box motif (CBM), where Φ represents a hydrophobic residue and p represents a polar

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Abbreviations used: 5-FOA, 5-fluoroorotic acid; AP, heterotetrameric adaptor protein complex; Ape1, aminopeptidase I; CBM, clathrin-box motif; CCV, clathrin-coated vesicle; Cvt, cytoplasm-to-vacuole transport; HC, heavy chain; LAT-A, latrunculin-A; LC, light chain; PM, plasma membrane; TD, N-terminal domain; TGN, *trans*-Golgi network; ts, temperature sensitive; TIRFM, total internal reflection fluorescence microscopy.

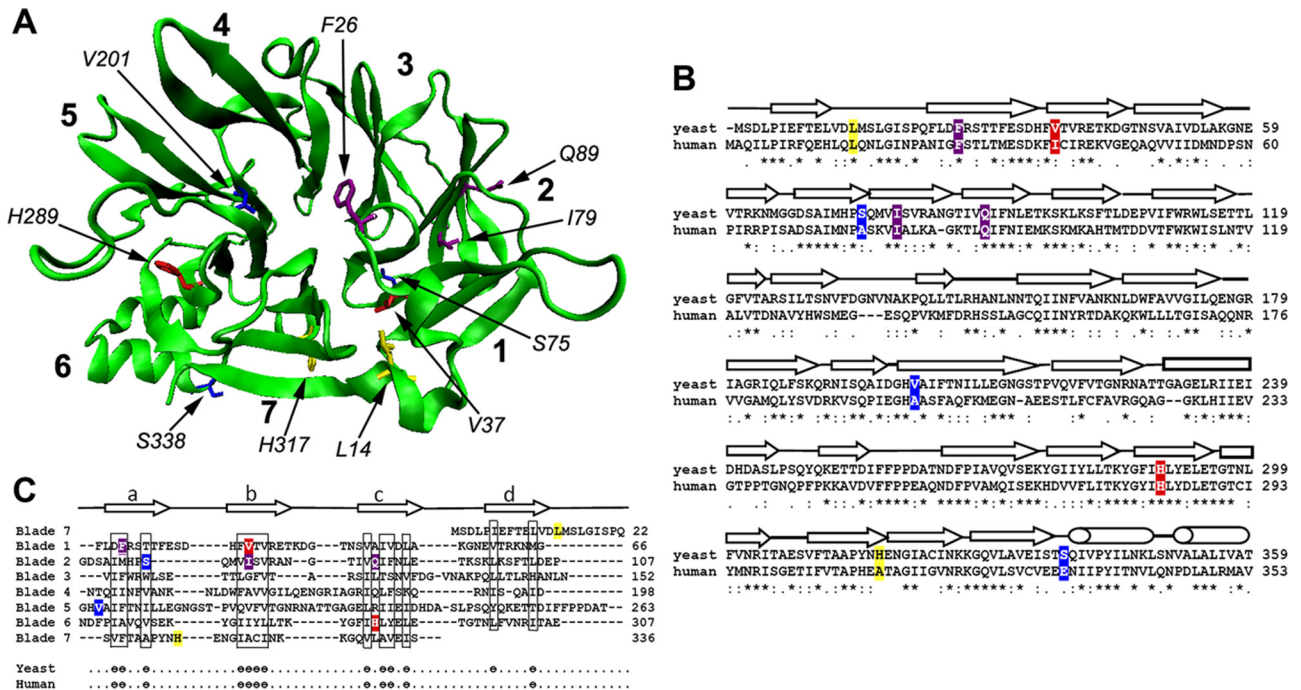


Figure 1. Structure of the clathrin heavy chain TD. (A) A homology model of the yeast TD (green) with blades 1–7 of the β -propeller numbered in bold. Side chains from amino acid residues that were mutated in the *chc1-TD-ts* alleles and the *chc1-box* allele are shown and color coded for each allele as follows: *chc1-1* and *chc1-3* (dark blue), *chc1-2* (red), *chc1-4* (yellow), and *chc1-box* (purple). (B) Sequence alignment of the yeast TD (residues 1–359) and human TD (residues 1–353) regions of clathrin heavy chain. Residues that were mutated in the *chc1-TD-ts* alleles and the *chc1-box* allele are highlighted according to the same color scheme used in A. An asterisk (*) denotes identical residues between the yeast and human sequence, a colon (:) denotes a conserved substitution between sequences, and a period (.) denotes a semiconserved substitution between sequences. (C) Alignment of the seven blades of the TD β -propeller structure. The approximate boundaries for every strand in each blade are denoted above the alignment. θ denotes conserved hydrophobic residues in both the yeast and human sequence. The S75P mutation of *chc1-1* is also present in *chc1-3*.

residue (reviewed by Dell’Angelica, 2001). More recently, a second clathrin-binding motif called the W-box motif (PWXXW), where X represents any amino acid, was identified in amphiphysin and sorting nexin 9 (SNX9), which are required for efficient endocytosis in mammalian cells (Ramjaun and McPherson, 1998; Drake and Traub, 2001; Lundmark and Carlsson, 2003; Miele *et al.*, 2004).

X-ray crystallographic studies have revealed the location of the binding sites on the TD for the CBM and W-box motif peptides present in clathrin adaptors. The TD forms a seven-bladed propeller structure where the fold of each blade resembles a WD40 repeat, a protein–protein interaction module found in heterotrimeric G proteins that forms a four-strand β sheet (ter Haar *et al.*, 1998) (Figure 1A). A binding pocket in the groove between blades 1 and 2 of the β -propeller structure accommodates the CBM peptides from the AP-3 β -subunit (LLDL) and the G protein-coupled receptor-specific adaptor β -arrestin 2 (LIEFE) (ter Haar *et al.*, 2000). A W-box motif peptide from amphiphysin (PWDLW), an accessory protein that facilitates membrane curvature during endocytic vesicle formation, fits into a deep pocket formed at the center of the top surface of the propeller structure (Miele *et al.*, 2004). The contribution of these binding sites to clathrin-mediated protein trafficking events has not been tested *in vivo*.

Clathrin-mediated sorting is evolutionarily conserved, and many of the proteins that are critical for these pathways have homologues in lower eukaryotic cells. The yeast *Saccharomyces cerevisiae* has one clathrin heavy chain gene (*CHC1*), encoding a protein with nearly 50% identity to mammalian HC, and a light chain gene (*CLC1*). Clathrin-

deficient yeast grow poorly, and in some strains a null allele is lethal (Lemmon and Jones, 1987; Munn *et al.*, 1991). They redistribute Kex2 and dipeptidylaminopeptidase (Ste13), two enzymes involved in the processing and maturation of the mating pheromone α -factor, from the TGN to the plasma membrane (Payne and Schekman, 1989; Seeger and Payne, 1992). Clathrin is also important for endocytosis because the internalization of α -factor bound to its receptor Ste2, a G protein-coupled receptor, is reduced in cells lacking clathrin (Tan *et al.*, 1993; Huang *et al.*, 1997). There are many different TGN/endosomal adaptor proteins in yeast (AP-1, Gga1, and Gga2 and the epsin-related proteins Ent3 and Ent5), and disruption of these adaptor genes results in TGN/endosomal sorting phenotypes (Costaguta *et al.*, 2001; Mullins and Bonifacino, 2001; Duncan *et al.*, 2003; Friant *et al.*, 2003; Eugster *et al.*, 2004). However, in studies in which adaptor proteins containing CBM mutations were analyzed, no significant phenotypes were observed (Mullins and Bonifacino, 2001; Yeung and Payne, 2001). At least four endocytic adaptors have been studied in yeast—the epsin-related proteins Ent1 and Ent2 and the AP180 homologues Yap1801 and Yap1802—and all four possess C-terminal CBMs in which the carboxylic acid at the end of the polypeptide substitutes for the final polar residue in the motif. Yeast strains carrying a deletion in all four endocytic adaptor genes complemented with Ent1 lacking its CBM show no defects in uptake of the lipophilic dye FM4-64 or the α -factor receptor Ste3 (Baggett *et al.*, 2003; Maldonado-Baez *et al.*, 2008). The absence of major trafficking phenotypes in studies of CBM-deficient adaptor proteins could be due to the utilization of alternative binding site(s) on clathrin by these mutated adaptors, or

Table 1. Yeast strains used in this study

Strain	Genotype	Source
BJ3502	<i>MATa ura3-52 his3 can1^r pep4Δ::HIS3 prb1-Δ1.6R</i>	
BJ3556	<i>MATa sst1-2 ade2-1 his6 met1 cyh2 rme1 ura1 can1</i>	
PJ694α	<i>MATα SL2793</i>	
SL12	<i>MATa leu2 ura3-52 trp1 chc1Δ::LEU2 scd1-v</i>	
SL13	<i>MATα leu2 ura3-52 trp1 chc1Δ::LEU2 scd1-v pSL6 (URA3, CEN, CHC1)</i>	
SL82	<i>MATα leu2 ura3-52 trp1 his1 ade6 chc1Δ::LEU2 scd1-i pAP4 (TRP1, CEN, CHC1)</i>	
SL119	<i>MATα leu2 ura3-52 trp1 chc1Δ::LEU2 scd1-v</i>	
SL1462	<i>MATa leu2 ura3-52 trp1 his3-Δ200</i>	
SL1463	<i>MATα leu2 ura3-52 trp1 his3-Δ200</i>	
SL2567	<i>MATα ura3-52 leu2,3-112 his3Δ200 trp-Δ901 lys2-801 suc2-Δ9 vac8-Δ::HIS3</i>	
SL2793 (PJ694a)	<i>MATa leu2-3,112 ura3-52 his3-Δ200 trp1-901 gal4Δ ade2 gal80Δ GAL2-ADE2 LYS2::GAL1-HIS3 met2::GAL7-lacZ</i>	
SL3593	<i>MATa leu2 ura3-52 trp1 bar1-1 chc1Δ::LEU2</i>	
SL5089	<i>MATa leu2 ura3-52 trp1 his3-Δ200 ABP1-mRFP:KanMX</i>	
SL5171	<i>MATα ura3 chc1Δ::LEU2 scd1-i pJRC3 (TRP1, CEN, chc1-2 [V37A,H289R])</i>	
SL5177	<i>MATa ura3 his1 chc1Δ::LEU2 scd1-i pJRC2 (TRP1, CEN, chc1-1 [S75P])</i>	
SL5178	<i>MATa ura3 chc1Δ::LEU2 scd1-i pJRC5 (TRP1, CEN, chc1-4 [L14P,H317R])</i>	
SL5181	<i>MATα ura3 chc1Δ::LEU2 scd1-i pJRC4 (TRP1, CEN, chc1-3 [S75P,V201A,S338P])</i>	
SL5538	<i>MATa ura3-52 trp1 his3Δ200 leu2 chc1Δ::LEU2 ABP1-mRFP:KanMX GFP-CLC1</i>	
SL5634	<i>MATα leu2 ura3 trp1 chc1Δ::LEU2 scd1-i pJRC19 (TRP1, CEN, chc1-box [F26W,I79S,Q89M])</i>	
SL5719	<i>MATa leu2 ura3-52 trp1 his3-Δ200 ade2 chc1Δ::LEU2 chs6Δ::HIS3</i>	
YRV19	<i>MATa chs6Δ::HIS3 ade2-101oc his3-Δ200 leu2-Δ1 trp1 ura3-52</i>	R. Schekman

other adaptors are able to compensate for the mutated adaptors and promote clathrin function.

Because no major TGN sorting or endocytic phenotypes have been observed upon removal of the CBMs from various yeast adaptor proteins, we decided to directly test the importance of their clathrin binding sites on the TD in vivo by introducing mutations into *CHC1* that disrupt the conserved binding sites for both the CBM and the W-box motif (*chc1-box*). In addition, TD-specific, temperature-sensitive (ts) alleles of the *CHC1* gene were generated and analyzed for their effects on clathrin function. In this report, we demonstrate that the TD is critical for overall clathrin function but that the previously identified adaptor binding sites on the TD are not necessary, suggesting that additional interactions on the propeller mediate adaptor association and facilitate clathrin assembly at sites of vesicle formation.

MATERIALS AND METHODS

Strains, Media, and Growth Methods

Strains used in this study are listed in Table 1. YEPD and synthetic dropout media were prepared, and yeast mating, sporulation, and tetrad dissections were performed as described previously (Guthrie and Fink, 1991). Transformation of yeast was performed using the lithium acetate method of Gietz *et al.* (1995). For growth tests, log phase cultures were diluted to 10^7 cells/ml, and serial fourfold dilutions were spotted onto C-TRP plates and grown for 2–3 d at the indicated temperatures. To assess calcofluor sensitivity, cultures were diluted to 10^7 cells/ml, and serial fourfold dilutions were spotted onto YEPD plates with 50 μg/ml calcofluor white and grown for 3–5 d at 25 or 30°C where indicated. For the α-factor halo assay, a YEPD plate was first seeded with 5×10^5 BJ3556 cells in an agar overlay to generate a *MATa sst1* lawn. Liquid cultures of *MATα* test cells and controls were diluted to 10^7 cells/ml and spotted onto the previously seeded plate and then incubated at 30°C for 48 h.

Generation of Mutant *chc1* Alleles

To generate *chc1-TD-ts* alleles, the TD coding sequence (residues 1–363) of pAP4 (*CEN, TRP1, CHC1*, described in Lemmon *et al.*, 1991) was replaced with the His3MX6 module flanked on either side by unique NotI restriction sites to generate pJRC1. Next, error-prone PCR mutagenesis (Bumbulis *et al.*, 1998) was carried out with oligonucleotides (oligos) corresponding to 50 bp 5' to the *CHC1* ATG start codon and 50 bp 3' to the end of the TD coding sequence (corresponding to amino acid 363), respectively, by using pAP4 as the template. NotI-digested pJRC1 and products of error-prone polymerase chain

reaction (PCR) were cotransformed into SL13 (*chc1Δ::LEU2* + pSL6 [*CEN, URA3, CHC1*]) (Lemmon and Jones, 1987) and repaired plasmids containing TD-specific point mutations were selected for on –URA-TRP-selective medium. To screen for ts alleles, colonies were replica plated onto two –TRP plates containing 5-fluoroorotic acid (5-FOA) to select for loss of the *URA3, CHC1* plasmid and plates were incubated at 25 or 37°C. Plasmid DNAs from transformants that grew at 25°C but not at 37°C were isolated, retransformed into SL13, and retested for growth on –TRP + 5-FOA plates at 25 and 37°C to verify the ts nature of these alleles. DNA sequencing confirmed the generation of *CEN, TRP1* plasmids pJRC2 (*chc1-1*), pJRC3 (*chc1-2*), pJRC4 (*chc1-3*), and pJRC5 (*chc1-4*).

Generation of the *chc1-box* allele was as follows. First, plasmid pRTH7 was generated using PCR amplification of the TD + ankle coding region (residues 1–500) from pAP4 with primers to allow for gap repair into the two hybrid GAL-binding domain plasmid pGBDU-C1 (James *et al.*, 1996). Next, a BamHI/SalI fragment from pRTH7 was ligated into pGEX-4T1 (GE Healthcare, Chalfont St. Giles, Buckinghamshire, United Kingdom) to generate pTMN7. Targeted mutagenesis of the TD to generate the *chc1-box* allele was performed using the QuikChange system (Stratagene, La Jolla, CA) with pTMN7 as the template. A TD fragment containing the three mutations (F26W, I79S, and Q89M) was amplified by PCR and cotransformed with Bsu36I and SexAI-digested pSL6 for gap repair in yeast to generate pSL6-box (*CEN, URA3, chc1-box*). NotI-digested pJRC1 and a TD fragment containing the *chc1-box* mutations amplified by PCR were cotransformed into yeast to generate pJRC19 (*CEN, TRP1, chc1-box*).

Plasmids

pTMN38 (2μ, *URA3, GBD-ENT1*) has been described in Newpher and Lemmon (2006). Plasmids pJRC6 (2μ, *URA3, GBD-CHC1-TD[1-363]*) and pJRC8 (2μ, *LEU2, GAD-CHC1-TD[1-363]*) were generated using PCR amplification of the TD coding region from pAP4 with primers to allow for gap repair into two-hybrid GAL-binding domain (pGBDU-C1) and GAL-activation domain (pGAD-C1) plasmids (James *et al.*, 1996), respectively. Plasmids pJRC14 (2μ, *LEU2, GAD-CHC1-TD[S75P]*), pJRC15 (2μ, *LEU2, GAD-CHC1-TD[L14P,H317R]*), pJRC16 (2μ, *LEU2, GAD-CHC1-TD[F26W,I79S,Q89M]*), pJRC17 (2μ, *LEU2, GAD-CHC1-TD[S75P,V201A,S338P]*), and pJRC18 (2μ, *LEU2, GAD-CHC1-TD[V37A,H289R]*) were generated in identical manner to pJRC8 by using the appropriate template DNAs for each mutant allele. Plasmids pJRC20 (2μ, *URA3, GBD-CHC1-TD[S75P]*), pJRC21 (2μ, *URA3, GBD-CHC1-TD[V37A,H289R]*), pJRC22 (2μ, *URA3, GBD-CHC1-TD[S75P,V201A,S338P]*), pJRC23 (2μ, *URA3, GBD-CHC1-TD[L14P,H317R]*), and pJRC24 (2μ, *URA3, GBD-CHC1-TD[F26W,I79S,Q89M]*) were generated in identical manner to pJRC6 by using the appropriate template DNAs for each mutant allele. Cloning of yeast open reading frames into the GAL4 activation domain plasmid pOAD have been described in Hudson *et al.* (1997). To generate pOAD-Ent2(ΔCBM), a PCR product was generated using primers designed to introduce a stop codon following serine 609, immediately before the beginning of the C-terminal CBM, and to allow for gap repair into pOAD-Ent2 ORF. All PCR amplified regions were sequenced in resultant plasmids to verify the correct alleles and no other mutations were present.

Integration of GFP-CLC1

To generate cells expressing GFP-Clc1, a 3.5-kb EcoRI/NotI fragment from pTMN17 (Newpher *et al.*, 2005) containing the coding sequence for GFP-Clc1 was ligated with EcoRI/NotI digested pRS306 to generate pJRC7. pJRC7 was digested with PacI to linearize the plasmid in the *CLC1* region for targeted integration. This was transformed into a diploid strain resulting from the mating of SL13 with SL5089 in which the *URA3*, *CHC1* plasmid had been dropped after mating. Cells were grown on synthetic complete medium + 5-FOA to select for excision of the *URA3* gene and plasmid sequences, and candidate yeast colonies were screened for GFP-Clc1 expression by fluorescence microscopy. To verify removal of the *URA3* gene, green fluorescent protein (GFP)-positive colonies were tested to confirm their inability to grow on -URA plates. Tetrad dissection resulted in the generation of SL5538.

Biochemical Procedures

For immunoblots of clathrin HC, extracts were prepared from 10 ml of log-phase cells (5×10^6 cells/ml) grown at the indicated temperatures by glass bead lysis in 250 μ l of 0.1 M Tris, pH 7.5, containing 1 mM phenylmethylsulfonyl fluoride and a protease inhibitor cocktail (Stepp *et al.*, 1995). Lysates were centrifuged for 20 min at $10,000 \times g$, and the supernatants were analyzed by SDS-polyacrylamide gel electrophoresis (PAGE), and immunoblotted using anti-Chc1 mouse monoclonal antibodies (Lemmon *et al.*, 1988) and anti- α -tubulin rat monoclonal antibodies (ab6161) as a loading control (Abcam, Cambridge, MA). To monitor aminopeptidase I (Ape1) processing, extracts from cells grown at 30°C were prepared as described previously (Huang *et al.*, 1999), and aminopeptidase was detected by immunoblotting using a rabbit-anti-Ape1 polyclonal antiserum at a 1:10,000 dilution (gift from Dan Klionsky, University of Michigan, Ann Arbor, MI). Clathrin-coated vesicles (CCVs) were isolated from SL5067 (*CHC1*) and SL5551 (*chc1-box*) yeast grown at 30°C by Sephacryl S-1000 column chromatography as described previously (Lemmon *et al.*, 1988). Polyclonal rabbit antiserum to Apm1 has been described previously (Stepp *et al.*, 1995). Detection of all proteins was performed with the Odyssey Infrared Imaging System (Li-Cor, Lincoln, NE) by using secondary antibodies labeled with infrared fluorophores.

α -Factor Endocytosis Assays

35 S- α -factor uptake was performed as described previously (Dulic *et al.*, 1991). Cells were incubated for 15 min at 37°C before the addition of radiolabeled α -factor. Plots show the best-fit trend lines generated in Excel (Microsoft, Redmond, WA). Data are the average of five independent experiments \pm SD. Endocytic rates were determined by taking the average slope of the best-fit trend lines from the 3- to 9-min time points.

Microscopic Analysis of Cortical Clathrin Recruitment

Live cell imaging of GFP-Clc1p was performed using overnight cultures of cells grown to early to mid-log phase at 25°C in synthetic media. SL5538 (*chc1 Δ ABP1-RFP GFP-CLC1*) transformed with expression plasmids for *CHC1* or *chc1-TD-mut* alleles were treated with 250 μ M latrunculin A (LAT-A) dissolved in dimethyl sulfoxide for 20 min. Actin disassembly was visualized in all cells using Abp1-red fluorescent protein (RFP) (data not shown). Clathrin accumulation was defined by the appearance of intense GFP-Clc1p cortical rim localization or by the appearance of discrete GFP-Clc1p cortical puncta. For each strain, 100–150 cells were counted, and data are from three experiments. Microscopy was performed using a fluorescence BX61 microscope (Olympus, Melville, NY) equipped with Nomarski differential interference contrast optics, a Uplan Apo 100 \times objective (numerical aperture [NA] 1.35), a CoolSNAP HQ camera (Photometrics, Tucson, AZ), Lambda 10-2 excitation and emission filters and Lambda LS 175-W xenon remote light source with liquid light guide (Sutter Instrument, Novato, CA).

To perform total internal reflection fluorescence microscopy (TIRFM), SL5538 (*chc1 Δ ABP1-RFP GFP-CLC1*) transformed with *CHC1* or *chc1-TD-mut* expression plasmids were grown to early to mid-log phase at 25°C in synthetic media, washed in phosphate-buffered saline (PBS), and 250 μ l of cells was added to polylysine-coated dishes (MatTek, Ashland, MA) for 5 min. Excess cells were washed away, and 50 μ l of fresh synthetic complete medium was added before coverslips were applied to prevent drying. TIRFM was performed using an IX-81 scope (Olympus), using a 100 \times /1.46NA PlanApo total internal reflection fluorescence (TIRF) objective (Olympus). Samples were visualized by illumination with 488 nm argon krypton and 543 nm helium neon lasers and captured by a three-color charge-coupled device (Hamamatsu, Bridgewater, NJ). Four-minute movies were taken at 2-s intervals by using 250-ms exposures for both RFP and GFP signals. For each strain, GFP-Clc1p patches were sorted into three categories: “mobile” patches that acquired the late endocytic actin marker protein Abp1-RFP before disappearing from the cell cortex; “immobile” patches that persisted at the cortex for >1 min and disappeared without acquiring actin; and “unproductive” patches, which appeared for <1 min but disappeared without receiving actin. Wide-field and TIRF images were acquired using SlideBook 4.01 (Intelligent Imaging Innovations, Santa Monica, CA).

Table 2. TD mutations in *chc1* alleles

<i>chc1</i> allele	Mutation
<i>chc1-1</i>	S75P
<i>chc1-2</i>	V37A, H289R
<i>chc1-3</i>	S75P, V201A, S338P
<i>chc1-4</i>	L14P, H317R
<i>chc1-box</i>	F26W, I79S, Q89M

Two-Hybrid Interaction Assays

For investigation of TD interaction with Ent1, overnight cultures of the two-hybrid reporter strain SL2793 cotransformed with bait and prey plasmids were diluted to 5×10^6 cells/ml and spotted onto complete synthetic medium lacking leucine and uracil (C-LEU-URA) to test for general growth and medium lacking leucine, uracil, and adenine (C-LEU-URA-ADE) to test for induction of the *GAL2-ADE2* reporter. Cells were grown at 30°C for up to 5 d.

For investigation of TD interaction with Ent2 and Apl2, the PJ694 α “bait” reporter strain transformed with each pGBD-CHC1-TD plasmid (or pGBD alone) was mated with the PJ694 α “prey” reporter strain transformed with the corresponding pOAD-adaptor plasmid (or pOAD alone), and diploids were selected for on C-LEU-URA plates. Overnight cultures of each diploid strain were diluted to 5×10^6 cells/ml and spotted onto C-LEU-URA-ADE plates. Plates were incubated at 30°C for up to 5 d.

Homology Modeling of the Yeast TD

A homology model of the yeast TD (residues 1–363) was generated from the rat TD structure (PDB 1BPO) by using Swiss Model and visualized using the Visual Molecular Dynamic viewer (University of Illinois, Urbana-Champaign, IL). A sequence alignment of the yeast and human TD coding sequence was performed using ClustalW (European Molecular Biology Laboratory-European Bioinformatics Institute).

RESULTS

Generation of Heavy Chain N-terminal Domain-specific Alleles

Previous studies on clathrin function in *S. cerevisiae* have focused on observable phenotypes associated primarily with the null allele (*chc1 Δ*), or the *ts* allele *chc1-521* (*chc1-ts*) containing multiple point mutations in the C-terminal part of the protein that affect LC binding and HC trimerization (Pishvaei *et al.*, 1997). To investigate the importance of the TD region on HC function, we sought to generate TD-specific *chc1-ts* alleles. Using random PCR mutagenesis and standard gap repair and plasmid shuffling techniques (see *Materials and Methods* for details), we isolated four *chc1-TD-ts* alleles [*chc1-(1-4)*]. A list of the mutation(s) present in each allele is provided in Table 2, and the locations of mutated residues within the TD structure are denoted in Figure 1A.

In addition to the generation of the *chc1-TD-ts* alleles, we also generated a fifth HC allele (*chc1-box*) in which three point mutations (F26W, I79S, and Q89M) were introduced into the TD coding region. Glutathione transferase pulldown experiments have demonstrated previously that for the mammalian TD the F27W mutation, which corresponds to F26 in yeast TD, blocks the interaction of TD with a W-box motif-containing peptide from amphiphysin (Miele *et al.*, 2004). The Q89M mutation has been shown previously to significantly disrupt binding of β -arrestin 2 to the TD (Goodman *et al.*, 1997), and based on the x-ray crystal structures of the TD in complex with a clathrin-box peptide from either β -arrestin 2 or from the hinge region of the β -subunit of AP-3 (ter Haar *et al.*, 2000), the I79S and Q89M mutations in combination should block clathrin-box motif-containing adaptor interaction with the TD. Although no W-box-containing adaptors in yeast have been identified, these

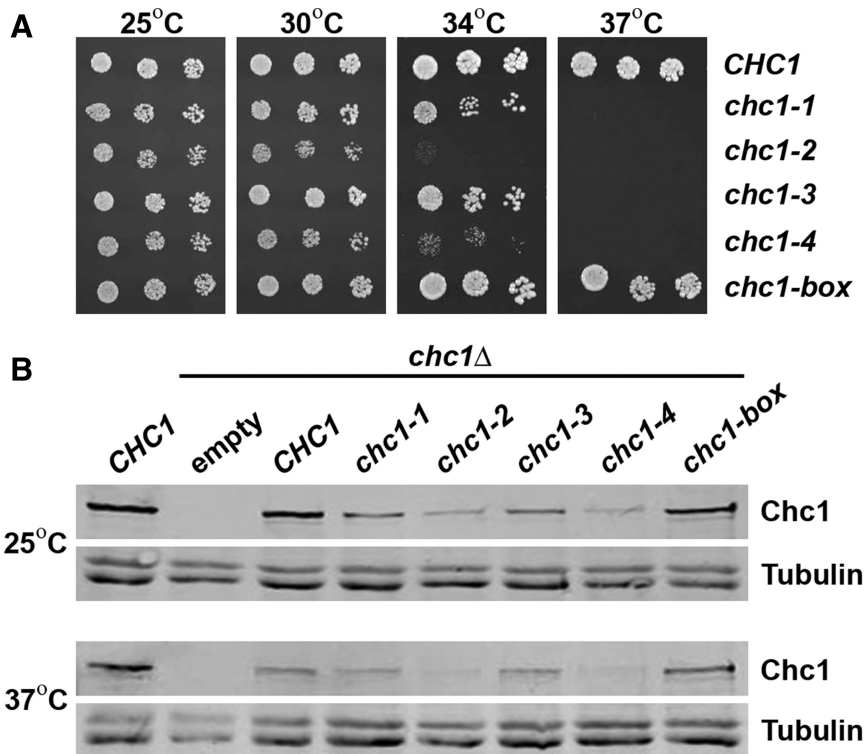


Figure 2. The *chc1-box* mutant is not temperature-sensitive for growth. (A) SL82 (*chc1* Δ + pAP4[*CEN, CHC1*]), SL5177 (*chc1* Δ + pJRC2[*CEN, chc1-1*]), SL5171 (*chc1* Δ + pJRC3[*CEN, chc1-2*]), SL5181 (*chc1* Δ + pJRC4[*CEN, chc1-3*]), SL5178 (*chc1* Δ + pJRC5[*CEN, chc1-4*]), and SL5634 (*chc1* Δ + pJRC19[*CEN, chc1-box*]) cells were diluted to 10^7 cells/ml, and fourfold serial dilutions were spotted onto plates and grown at the indicated temperatures for 3 d. (B) Immunoblots of extracts from *chc1* Δ yeast (SL12) transformed with pUN30 (*CEN, empty*), pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), or pJRC19 (*CEN, chc1-box*) were detected with anti-Chc1 antibodies. Cells were grown continuously at 25°C, and then one-half of each sample was removed and incubated for 90 min at 37°C before harvesting. To verify equal protein loading, blots also were probed with antibodies to tubulin.

mutations were all combined together to generate a TD that should have impaired binding to both motifs. Sequence alignment of the TD regions of yeast and human HC show that these residues are identical between species (Figure 1B).

To compare the severity of the ts alleles and to test for any growth phenotype associated with the *chc1-box* allele, spot dilutions of cultures were grown on plates over a range of temperatures. Yeast whose only source of HC was from expression of any of the four *chc1-TD-ts* alleles were able to grow at the permissive temperature of 25°C (Figure 2A), and similar to *chc1* Δ cells, were inviable at 37°C. However, at the intermediate temperatures of 30 or 34°C, *chc1-2* and *chc1-4* yeast displayed more severe growth phenotypes than *chc1-1* and *chc1-3* yeast (Figure 2A). Surprisingly, yeast expressing HC from the *chc1-box* allele grew normally at all temperatures (Figure 2A).

HC proteins encoded by these *chc1-TD-ts* alleles might be unstable as the point mutations introduced into the TD region could result in aberrant folding of the β -propeller structure and destabilization of the HC. To determine whether full-length HC was produced, protein extracts from yeast encoding HC from each of the *chc1-TD-ts* alleles and the *chc1-box* allele were prepared, and HC protein levels were examined by immunoblot analysis using monoclonal antibodies to Chc1. When cells were grown at 25°C, all of the HC mutant proteins were expressed and detected as full-length proteins of 190 kDa (Figure 2B). The amount of HC expressed from each ts allele was significantly reduced relative to the wild-type allele (20–25% of wild-type for *chc1-1* and *chc1-3* alleles and 10–14% of wild-type for *chc1-2* and *chc1-4* alleles; $n = 3$), but HC expression from the *chc1-box* allele was comparable with wild type (~90% of wild type; $n = 3$). Next, we tested whether these HC proteins would be stable after a shift to the nonpermissive temperature of 37°C for 90 min. The full-length protein was still present in cells

expressing HC from all alleles (Figure 2B), although levels were significantly less in the more severe ts alleles (*chc1-2* and *chc1-4*) suggesting protein destabilization might reduce their function.

TGN Sorting in *chc1-TD* Mutants

Clathrin-deficient *MAT α* yeast are defective in the secretion of the mature form of the mating pheromone α -factor due to the mislocalization of α -factor processing enzymes, including Kex2, a subtilisin-like protease, and Ste13, a dipeptidyl-aminopeptidase, from the TGN to the plasma membrane (Payne and Schekman, 1989; Seeger and Payne, 1992). To determine whether HCs encoded by any of the *chc1-TD* mutant alleles could rescue the α -factor processing phenotype of *chc1* Δ cells, halo assays were performed in which *MAT α* yeast expressing different HCs were tested for their ability to inhibit the growth of *MAT α* cells supersensitive to mature α -factor at the semipermissive temperature of 30°C. Strong halos were observed in *chc1* Δ cells complemented with a plasmid carrying the *CHC1* allele (Figure 3A, 4) indicative of the secretion of mature α -factor and restoration of proper TGN sorting. Yeast expressing HC from any of the four *chc1-TD-ts* alleles did not fully rescue the TGN sorting phenotype of *chc1* Δ (Figure 3A, 5–8) as the halos produced by each of these strains were smaller than the halo generated by cells complemented with wild-type HC. The most severe phenotypes were seen with *chc1-2* and *chc1-4* (Figure 3A, 6 and 8, respectively), consistent with the stronger growth phenotype observed at intermediate temperatures. *chc1-box* yeast produced a halo very comparable to the wild-type allele, indicating TGN retention/sorting seems normal (Figure 3A, 9).

Ape1 Processing in *chc1-TD* Mutant Cells

Whereas most resident vacuolar hydrolases are delivered to the vacuole via the secretory pathway, at least one hydro-

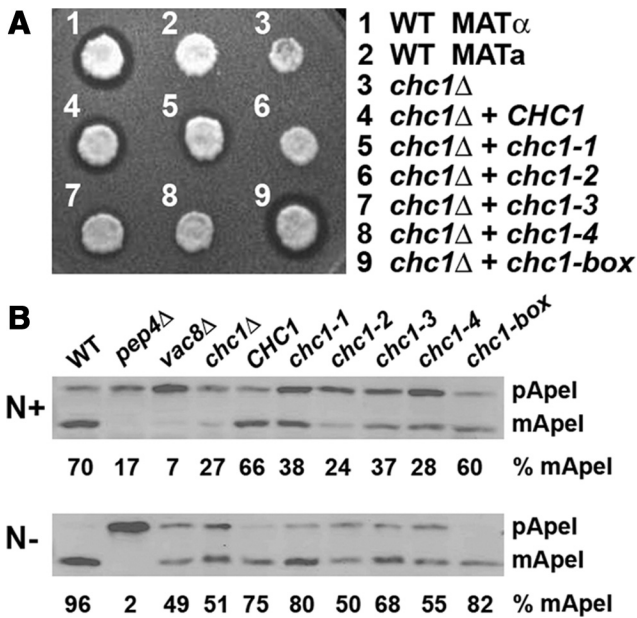


Figure 3. *chc1-TD-ts* alleles cause TGN sorting and Ape1 processing defects. (A) Halo assays for MAT α *chc1* Δ yeast (SL119) transformed with pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), pSL6-box (*CEN, chc1-box*), and wild-type controls, SL1462 (*MATa*) and SL1463 (*MATa*) spotted onto a lawn of *MATa* (BJ3556) and grown at 30°C for 2 d. A zone of growth inhibition demonstrates secretion of mature α -factor. (B) Ape1 processing: immunoblots of wild-type (SL1463), *pep4* Δ (BJ3502), *vac8* Δ (SL2567), *chc1* Δ (SL12), and SL12 transformed with pUN30 (*CEN, empty*), pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), or pSL6-box (*CEN, chc1-box*), grown in normal medium (N+) or nitrogen starvation medium (N-) at 30°C and blotted with anti-Ape1 antibodies. The amounts of precursor (pApe1) and mature aminopeptidase (mApe1) were determined by densitometry, and the percentage of mApe1 is reported relative to total mApe1 + pApe1 for each.

lase, Ape1, is delivered through the cytoplasm-to-vacuole (Cvt) targeting pathway (Harding *et al.*, 1995). In this pathway, Ape1 is initially synthesized as a precursor protein in the cytoplasm, and upon delivery to the vacuole is processed in a Pep4-dependent manner to generate the active protease. Many of the genes involved in the Cvt pathway also are involved in autophagy (Khalfan and Klionsky, 2002), and our lab showed previously that clathrin deficiency causes Ape1-processing defects (Huang *et al.*, 1999). To determine whether the TD mutations affect Ape1 maturation, *chc1-TD-ts* and *chc1-box* cells were grown at the semi-permissive temperature of 30°C in nitrogen-rich medium (Cvt pathway growth conditions) or nitrogen starvation medium (which induces autophagy), and Ape1 processing was monitored by the blotting of extracts with aminopeptidase I antibodies.

In wild-type cells, the majority of Ape1 protein (~70%) is present as the 50-kDa mature protease (mApe1) under Cvt conditions (N+) (Figure 3B), but in *chc1* Δ cells only 27% was mature, similar to previous observations (Huang *et al.*, 1999). In strains expressing HC from any of the *chc1-TD-ts* alleles, more Ape1 was present in the precursor form compared with that from *CHC1* cells. Ape1 processing in the null (*chc1* Δ) and all *chc1-TD-ts* mutants was more complete upon stimulation of the autophagy pathway; however, *chc1-2* and *chc1-4* cells more closely resembled the *chc1* Δ phenotype,

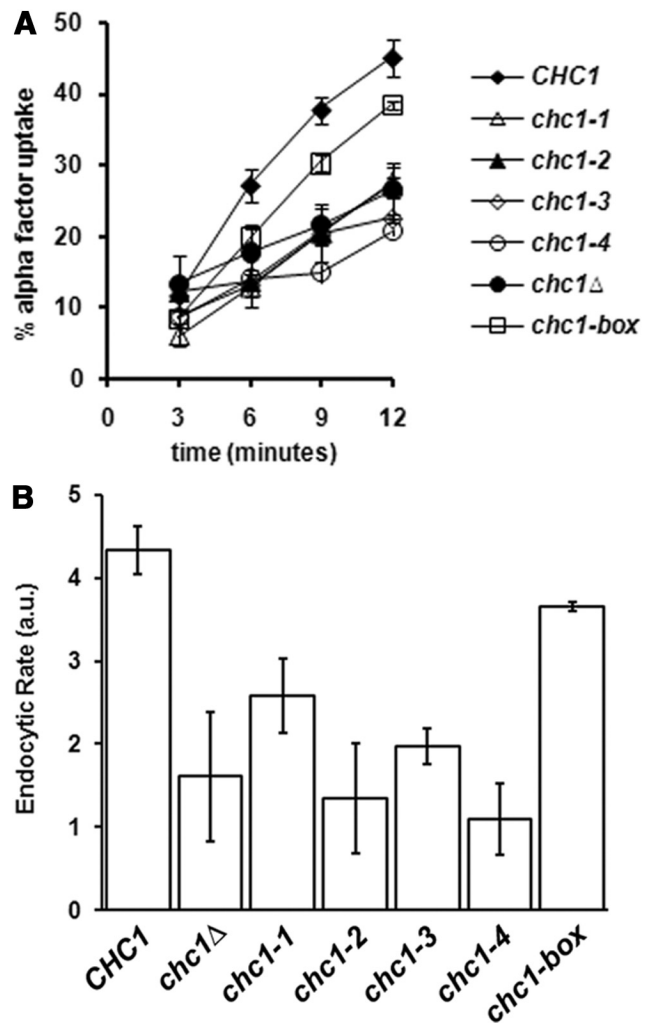


Figure 4. The *chc1-box* allele, but not *chc1-TD-ts* alleles, rescues the endocytic phenotype of *chc1* Δ . *MATa chc1* Δ *bar1-1* (SL3593) was transformed with pUN30 (*CEN, empty*), pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), or pJRC19 (*CEN, chc1-box*). α -factor internalization was analyzed at 37°C as described in *Materials and Methods*.

whereas *chc1-1* and *chc1-3* cells more closely resembled *CHC1*. *chc1-box* cells yielded mature protease comparable to *CHC1* under both Cvt (N+) and autophagy (N-) conditions (Figure 3B), indicating that Ape1 processing and Cvt/autophagy are normal in the *box* mutant.

Endocytic Analysis in *chc1-TD* Mutants

Studies of endocytosis in yeast have focused on the internalization of the mating pheromone α -factor bound to the G protein-coupled receptor Ste2. Clathrin-deficient yeast have partial defects in endocytosis, as α -factor internalization is reduced by ~ 50% (Tan *et al.*, 1993; Huang *et al.*, 1997). To determine whether the TD is critical for HC function in endocytosis, we performed α -factor internalization assays with cells expressing HC-TD mutations after shifting to 37°C. In *chc1* Δ yeast transformed with an empty *CEN* plasmid, the amount of internalized α -factor after 12 min of uptake was roughly 50% of that seen in *chc1* Δ yeast complemented with wild-type *CHC1* (Figure 4A), similar to previous studies of clathrin-deficient yeast (Tan *et al.*, 1993;

Huang *et al.*, 1997). Yeast cells expressing HC from any of the four *chc1-TD-ts* alleles were not able to suppress the α -factor internalization defect of *chc1* Δ cells (Figure 4A), but the *chc1-box* allele provides near complete rescue, with an endocytic rate that was $\sim 85\%$ of wild type (Figure 4B).

Visualization of Clathrin and Endocytic Dynamics in TD Mutants

GFP-LC is a suitable marker for localizing HC within yeast cells as GFP-Clc1 has been shown to completely colocalize with the fluorescent signal of Chc1-RFP (Newpher *et al.*, 2005). GFP-LC localizes to the TGN, endosomes, and small cortical patches. Cortical actin patches are the sites of endocytosis in yeast, and clathrin recruitment to these sites is dependent upon the endocytic adaptors Ent1/2 and Yap1801/2 (Newpher *et al.*, 2005), which have C-terminal clathrin-box motifs. However, the clathrin in these cortical patches is often masked by the intense fluorescent clathrin signal on TGN/endosomal structures.

To visualize clathrin and to enhance detection of cortical clathrin, we treated cells expressing GFP-Clc1, Abp1-RFP, and the TD mutant HCs with LAT-A. LAT-A is an actin monomer-sequestering drug that inhibits actin assembly, causing immobilization and accumulation of cortical patches containing early endocytic proteins including clathrin (Kaksonen *et al.*, 2003; Newpher *et al.*, 2005). In all TD mutants, GFP-LC was observed on internal structures (with and without LAT-A, Figure 5; data not shown), indicating that the HC from all alleles is able to bind endosomal/TGN compartments at 25°C. In cells expressing wild-type HC, $\sim 70\%$ of cells showed either discrete GFP-Clc1 puncta or rim GFP-Clc1 staining at the cortex after LAT-A treatment (Figure 5, *CHC1* + LAT-A, arrows). However, cortical clathrin accumulation in LAT-A-treated cells was significantly reduced (~ 10 – 23% of cells) in cells expressing HC from the *chc1-1*, *chc1-2*, and *chc1-3* alleles (Figure 5), consistent with the inability of these alleles to suppress the *chc1* Δ endocytic phenotype. Surprisingly, GFP-Clc1 was visualized at the cortex in a large percentage of cells ($\sim 50\%$ of cells) expressing HC from the *chc1-4* allele (Figure 5), but consistent with its endocytic defect at 37°C, its recruitment to cortical sites was ablated at elevated temperature (Supplemental Figure 1). The cortical accumulation of clathrin in *chc1-2* was also decreased to only a few percent of cells at 37°C as well (Supplemental Figure 1). Note that at elevated temperature there were still visible internal patches of clathrin, even though cortical accumulation was impaired; however, we cannot rule out that these are assemblies that form spontaneously and are not associated with membrane. In contrast, in the *chc1-box* mutant $\sim 50\%$ of cells had cortical clathrin, but this recruitment was not temperature dependent (Figure 5 and Supplemental Figure 1). This is consistent with the α -factor internalization assays where the *box* mutant internalized clathrin nearly as well as wild type at 37°C.

We next performed TIRF microscopy experiments to monitor the behavior of cortical/endocytic clathrin in these cells. In time-lapse movies of *CHC1* cells, three different types of GFP-Clc1 patch dynamics were observed. First, a majority of the patches at the cell surface recruited Abp1-RFP before moving away from the evanescent field and into the cell. These mobile patches represented 78% of the total number of events analyzed in *CHC1* cells ($n = 63$; Figure 6), in good agreement with previous observations (Newpher *et al.*, 2005). Second, occasional immobile GFP-Clc1 patches ($\sim 3\%$) would persist at the surface for an extended period (≥ 1 min), display very limited lateral movement, and never acquire Abp1-RFP before their disappearance (Figure 6).

Third, $\sim 19\%$ of events in wild-type cells consisted of shorter lived patches (< 1 min) where clathrin puncta formed but also did not acquire Abp1/actin before disappearing. We refer to these as unproductive events. When clathrin patch dynamics were observed in *chc1-TD-ts* and *chc1-box* cells at the permissive temperature, three different patterns emerged (Figure 6). Strikingly, *chc1-4* resembled the wild type, with $\sim 77\%$ of events scored as mobile, i.e., receiving Abp1 before disappearing. This pattern is consistent with the LAT-A analysis in Figure 5, showing substantial cortical recruitment of clathrin in *chc1-4* cells at 25°C. The three other *ts* alleles (*chc1-1*, *chc1-2*, and *chc1-3*) formed a second class, with ~ 40 – 50% unproductive patches, whereas the number of longer lived immobile patches was a smaller percentage (5–14%) of the total. The large percentage of unproductive patches may be reflected in the decreased recruitment in the presence of LAT-A (Figure 5). Significantly, the *chc1-box* allele forms a third class. It has fewer unproductive patches ($\sim 25\%$) compared with the second class (*chc1-1*, *chc1-2*, and *chc1-3*), but it also has an increased percentage of immobile patches that persist for > 60 s but disappear without receiving Abp1. Perhaps the immobile clathrin patches contribute to the cortical signal seen in LAT-A, thus explaining why *chc1-box* retains fairly strong cortical recruitment of clathrin.

Using wide field imaging, we also analyzed the dynamics of Sla2, an endocytic coat protein that is recruited after clathrin in the immobile phase of endocytosis. In each of the mutants, nearly all Sla2 cortical patches ($> 95\%$) had normal lifetimes and acquired Abp1 and internalized (data not shown). Also, there were very few of the Abp1-marked actin comet tails often seen in cells without clathrin. We infer from this that immobile and unproductive clathrin patches may abort before acquiring other later coat module factors, like Sla2, perhaps because of impaired ability to bind adaptors, engage other components of the endocytic machinery and/or cluster receptors. Abortive events during endocytic clathrin coat assembly have been clearly documented in mammalian cells (Ehrlich *et al.*, 2004; Loerke *et al.*, 2009).

Adaptor binding to HC-TD mutants

To investigate whether the TD regions encoded by these alleles could interact with endocytic adaptors, two-hybrid interaction assays between the endocytic adaptors Ent1 and Ent2, each containing a C-terminal CBM, and the HC-TD encoded by the *chc1-TD-ts* and *chc1-box* alleles were performed. In addition, two-hybrid interactions were examined between HC-TD proteins and Apl2, the large (β) subunit of the Golgi adaptor AP-1 that contains an internal CBM.

Although a wild-type TD prey interacted with an Ent1 bait (Figure 7A), none of the mutant TD prey, including one containing the *chc1-box* allele mutations, were able to bind (Figure 7B). Similar results were seen between HC-TD and Apl2 as only the wild-type TD bait interacted with the Apl2 prey, although the interaction with wild-type TD was fairly weak to begin with (Figure 7C). In this assay, the most robust interaction was observed between the wild-type TD and Ent2 (Figure 7C), but similar to Ent1, TDs containing any of the point mutations from the *chc1-TD-ts* alleles still disrupted this interaction. Surprisingly, a TD containing the *chc1-box* allele mutations was still able to interact with Ent2, although the interaction was much weaker compared with the wild-type TD (Figure 7C). The binding results were not caused by differences in stability of GBD-TD mutant proteins, as all were expressed to similar steady-state levels (see Supplemental Figure 2).

To assess whether Ent2 association with TD containing the *chc1-box* allele mutations was mediated by the Ent2 C-termi-

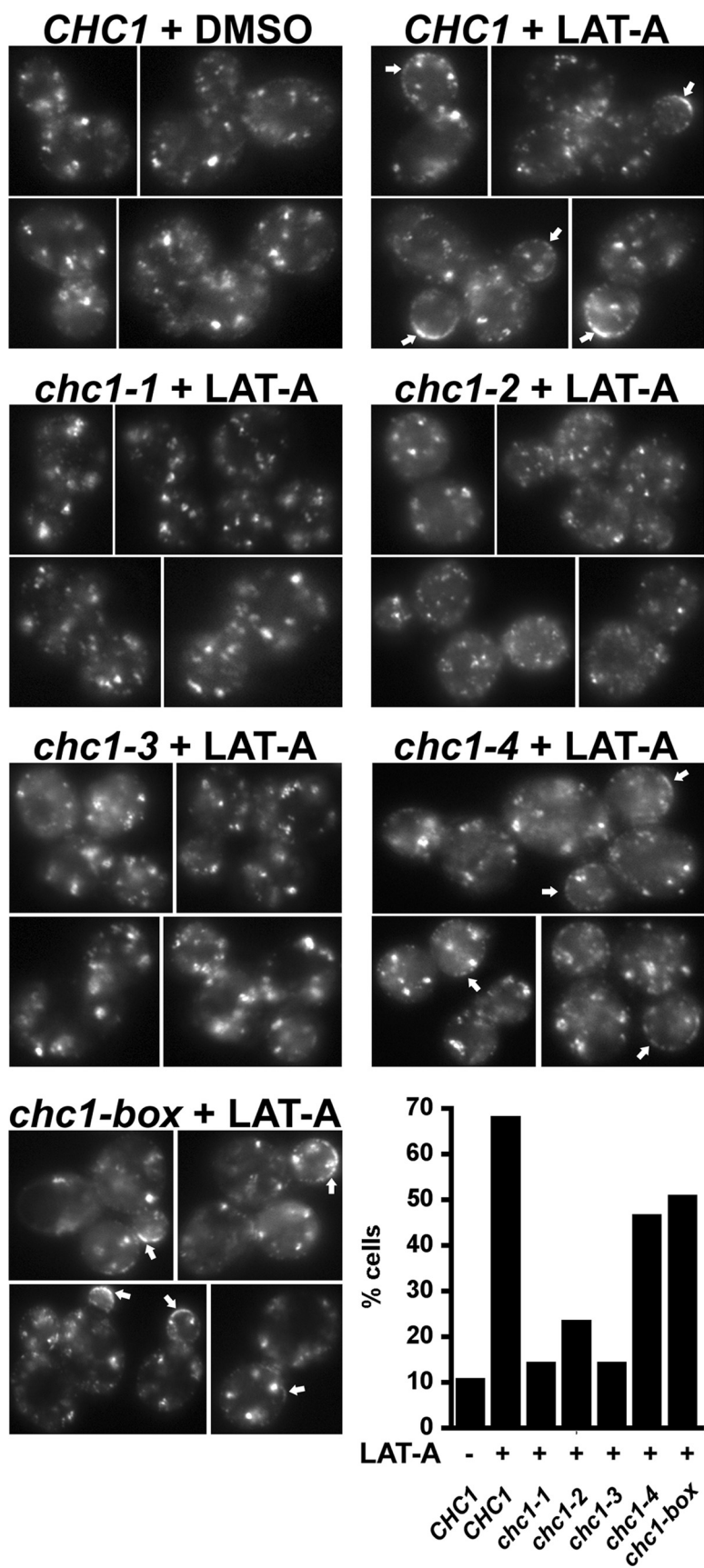


Figure 5. GFP-Clc1p accumulation at the cell cortex in *chc1-TD* alleles in the presence of LAT-A. *GFP-CLC1 ABP1-mRFP chc1Δ* (SL5538) was transformed with pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), or pJRC19 (*CEN, chc1-box*). Cells grown to log phase at 25°C were treated with LAT-A (250 μM) for 20 min at 25°C before imaging. Abp1-mRFP was completely cytosolic after 20 min in LAT-A, indicating the disassembly of actin (data not shown). Arrows indicate the accumulation of clathrin at the cell cortex. Cells that showed strong cortical puncta or rim GFP staining were considered to have accumulated clathrin. The percentage of cells for each condition that contained cortical GFP-Clc1 accumulation is shown in the graph (n = 100–150).

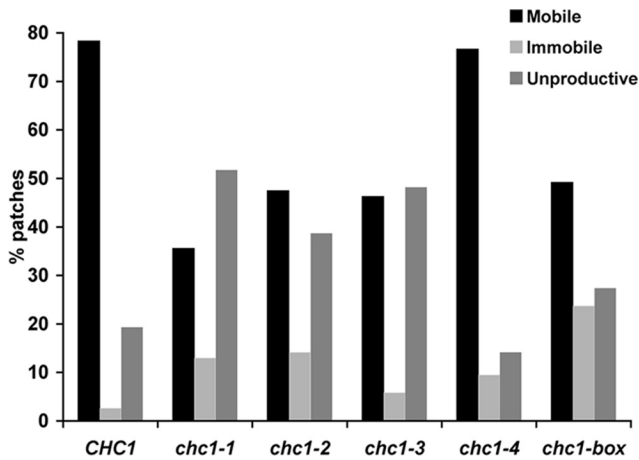


Figure 6. TIRFM of clathrin patch dynamics at the cell cortex. Graph shows percentages of GFP-Clc1 patch types observed in *CHC1* and *chc1-TD* mutants at 25°C. Between 50 and 80 GFP-Clc1 patches for each strain were defined as mobile, immobile or unproductive as described in *Materials and Methods*. Strains are the same as in Figure 5.

nal CBM, a truncated version of Ent2 lacking the final four residues (amino acids 610–613) that constitute the C-terminal CBM was generated (Ent2 Δ CBM) and tested in the two-hybrid assay. Ent2 Δ CBM was unable to interact with either wild-type TD or a TD containing *chc1-box* mutations (Figure 7C), indicating that the CBM is required for efficient Ent2 association with the TD.

Heavy Chain Encoded by *chc1-Box* Forms Clathrin-coated Vesicles

Recruitment of soluble clathrin triskelions to the membrane surface is mediated in part by interactions between the TD and clathrin-box motif containing adaptor proteins. The I79S and Q89M mutations of the *chc1-box* allele should disrupt TD-clathrin box adaptor protein interactions, but the lack of major growth, TGN sorting, α factor uptake, or Ape1 processing phenotypes for the *chc1-box* allele suggests that clathrin function is not significantly impaired in these cells. The microscopy data also indicates that Chc1-box protein is recruited to membranes even in the absence of the CBM-binding region. To further test for clathrin function, we subjected *chc1-box* cells to our standard protocol for isolation of CCVs. Cells were spheroplasted, lysed, and after differential centrifugation, a 100,000 \times g microsomal membrane pellet was resuspended and fractionated on a Sephacryl S-1000 column (Lemmon *et al.*, 1988). Column fractions were analyzed by SDS-PAGE followed by immunoblotting with monoclonal antibodies to Chc1. In *CHC1* cells, HC peaked in column fractions 34–36 (Figure 8). The heterotetrameric Golgi adaptor AP-1 also cofractionated with HC as shown by blotting for the μ chain Apm1 (Figure 8). When microsomal membranes from *chc1-box* cells were similarly fractionated, the HC protein peak was located in similar column fractions as was observed for *CHC1* cells (Figure 8), indicating that the HC protein encoded by *chc1-box* could form clathrin-coated vesicles. However, Apm1 did not cofractionate with clathrin isolated from *chc1-box* cells, but instead eluted in later fractions presumably as free AP-1 complex (Figure 8). This suggests that TD binding to AP-1 is critical for stable incorporation of this adaptor into clathrin-coated structures.

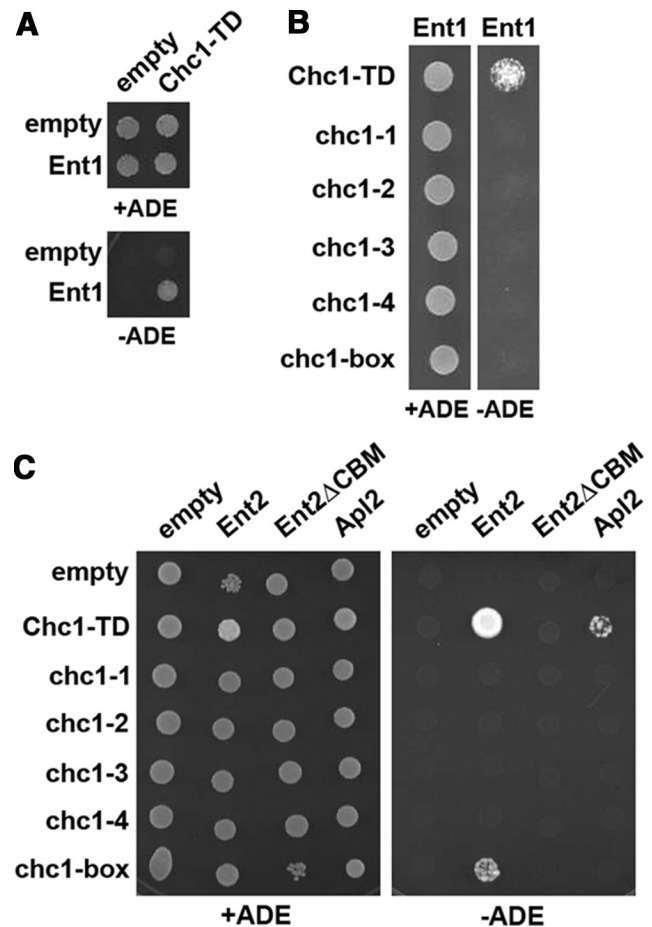


Figure 7. HC-TD mutations disrupt interactions with clathrin adaptors Ent1, Ent2, and Apl2. (A) A two-hybrid reporter strain (SL2793) was transformed with either the empty GAL4 binding domain (GBD) plasmid pGBDU-C1 or pGBD-*ENT1* in combination with either the empty GAL4 activation domain (GAD) plasmid pGAD-C1 or pGAD-*CHC1-TD*(1-363). Cells from each set of transformants were diluted to 5×10^6 cells/ml, spotted onto C-LEU-URA (total growth) and C-LEU-URA-ADE (activation of the *GAL2:ADE2* reporter) plates, and grown at 30°C for 3 d. (B) SL2793 was cotransformed with pGBD-*ENT1* and each of the following GAD-*CHC1-TD* plasmids: pJRC8 (GAD-*CHC1-TD*), pJRC14 (GAD-*CHC1-TD[chc1-1]*), pJRC15 (GAD-*CHC1-TD[chc1-2]*), pJRC16 (GAD-*CHC1-TD[chc1-3]*), pJRC17 (GAD-*CHC1-TD[chc1-4]*), or pJRC18 (GAD-*CHC1-TD[chc1-box]*) and plated and grown as described in A. (C) PJ694a was transformed with pGBDU-C1 or the following plasmids: pJRC6 (GBD-*CHC1-TD*), pJRC20 (GBD-*CHC1-TD[chc1-1]*), pJRC21 (GBD-*CHC1-TD[chc1-2]*), pJRC22 (GBD-*CHC1-TD[chc1-3]*), pJRC23 (GBD-*CHC1-TD[chc1-4]*), pJRC24 (GBD-*CHC1-TD[chc1-box]*) and mated with PJ694a transformed with either the empty GAD plasmid pOAD, pOAD-Ent2, pOAD-Ent2(Δ CBM), or pOAD-Apl2. Diploids were selected on C-LEU-URA plates, and then activation of the *ADE2* reporter was assayed as described above.

Chitin Synthase Trafficking in *chc1-TD* Mutant Strains

Because AP-1 was not stably associated with CCVs in the *chc1-box* mutant, we tested whether an AP-1-dependent pathway was affected in the TD mutants. The AP-1/clathrin pathway is involved in the transport of proteins, including chitin synthase III (Chs3) from early endosomes back to the TGN (Valdivia *et al.*, 2002). Chs3 can be found both intracellularly in Kex2-containing compartments and at the PM where it localizes to the mother-bud neck to deliver a ring of chitin around developing buds (Chuang and Schekman, 1996). The delivery of Chs3 from the TGN to the cell surface

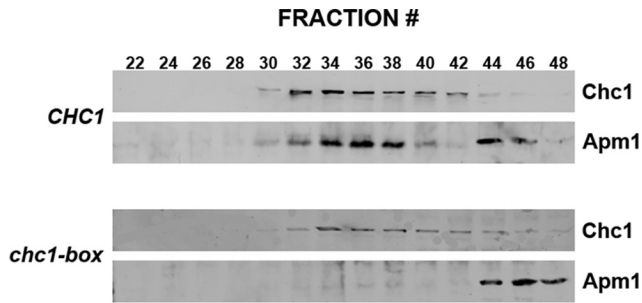


Figure 8. Clathrin HC encoded by the *chc1-box* allele forms coated vesicles. Clathrin-coated vesicles were prepared from *chc1Δ* cells (SL12) transformed with pAP4 (*CEN, CHC1*) or pJRC19 (*CEN, chc1-box*), grown at 30°C, and fractionated on a Sephacryl S-1000 column. Column fractions 22–48 for each preparation were immunoblotted with anti-Chc1 and anti-Apm1 antibodies.

is dependent upon the Chs5/Chs6 coat complex, as *chs5Δ* or *chs6Δ* cells show reduced chitin deposition at the bud site and intracellular retention of Chs3 (Santos and Snyder, 1997; Ziman *et al.*, 1998). *chs6Δ* yeast are therefore resistant to the toxic effects of the chitin-binding compound calcofluor white (Figure 9A) (Ziman *et al.*, 1998). Deletion of clathrin or AP-1 subunit genes eliminates intracellular Chs3 retention and restores calcofluor sensitivity in the *chs6Δ* background (Figure 9A) (Valdivia *et al.*, 2002).

Therefore, calcofluor sensitivity growth assays were performed with *chc1Δ chs6Δ* ($\Delta\Delta$) cells transformed with plasmids encoding wild-type or each of the *chc1-TD* mutants. $\Delta\Delta$ cells complemented with wild-type *CHC1* restored intracellular Chs3 retention and calcofluor resistance (Figure 9, A and B). Expression of HC from each *chc1-TD-ts* allele was not able to fully restore calcofluor resistance to $\Delta\Delta$ cells (Figure 9B), and consistent with the results obtained in the TGN sorting, mature α -factor secretion halo assay, the *chc1-2* and *chc1-4* mutants were more sensitive to calcofluor, although *chc1-4* retained some residual function at 25°C. The *chc1-box* allele restored calcofluor resistance in a manner similar to wild-type HCs (Figure 9B), indicating that Chs3 retrieval

from early endosomes to the TGN is efficient in these cells even though AP-1 association with CCVs is impaired.

DISCUSSION

Most previous studies examining the role of clathrin function in yeast have used either the null allele (*chc1Δ*) or a temperature-sensitive allele (*chc1-521*, also referred to as *chc1-ts*) that disrupts LC binding and/or trimerization of HC (Seeger and Payne, 1992; Pishvaei *et al.*, 1997). Additional clathrin HC mutations have been described previously (Lemmon *et al.*, 1991; Chen and Graham, 1998), but they also were localized in the C-terminus close to the trimerization domain. To directly assess the role of the TD in clathrin function, we generated a HC allele, *chc1-box*, containing substitutions in the binding sites for CBMs and W-box motifs present in known clathrin adaptor proteins. In addition, we isolated four *chc1-TD-ts* alleles in which random mutation(s) were introduced throughout the TD coding region. The TD seems to be necessary for overall clathrin function as the *chc1-TD-ts* alleles all demonstrated impairment of several sorting pathways that require clathrin. Surprisingly, HC expression from the *chc1-box* allele resulted in no major growth or trafficking phenotypes, suggesting that additional or alternative binding sites might exist on the TD for clathrin adaptor proteins.

chc1-TD-ts Alleles and the Location of Point Mutations

Although clathrin-dependent functions were perturbed in cells expressing HC from each *chc1-TD-ts* allele, the *chc1-1* and *chc1-3* mutants displayed less severe growth, TGN enzyme retention, endosome-to-TGN retrieval of Chs3, and Ape1 processing phenotypes compared with the *chc1-2* and *chc1-4* mutants. The severity of the phenotypes caused by each of these alleles may be due in part to differences in HC protein expression. In fact using siRNA-mediated knock-down of HC, Moskowitz *et al.* (2005) observed dramatic changes in endocytic rates of transferrin receptor with a narrow range of clathrin concentrations, and reduction of HC levels by as little as 30–40% significantly impaired internalization (Moskowitz *et al.*, 2005). The amount of HC

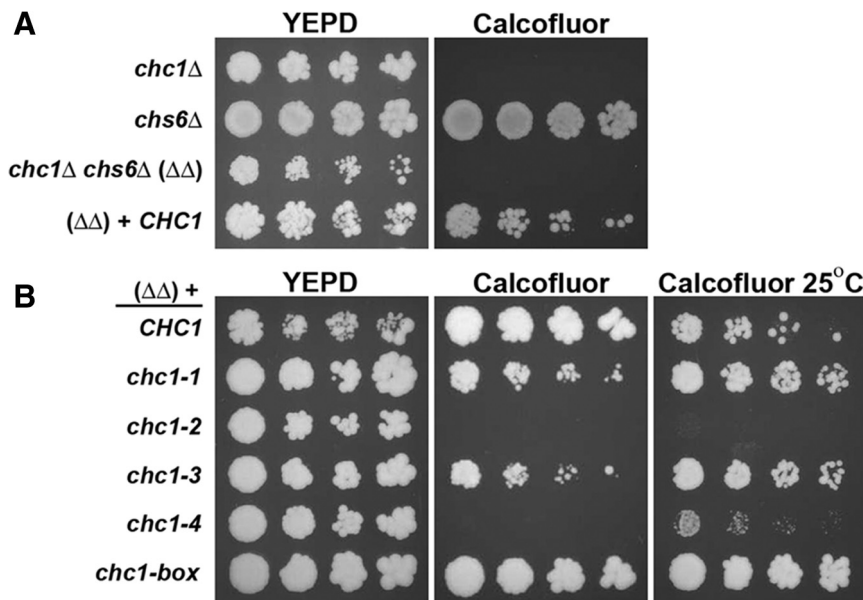


Figure 9. The *chc1-box* allele restores calcofluor resistance to *chs6Δ chc1Δ* ($\Delta\Delta$) cells. (A) SL119 (*chc1Δ*), YRV19 (*chs6Δ*), SL5719 (*chs6Δ chc1Δ* [$\Delta\Delta$]), and $\Delta\Delta$ + pAP4 (*CEN, CHC1*) cells were diluted to 1×10^7 cells/ml, and fourfold serial dilutions were plated onto both YEPD plates and YEPD plates containing 50 μ g/ml calcofluor white and incubated at 30°C for 3–5 d. (B) $\Delta\Delta$ cells transformed with pAP4 (*CEN, CHC1*), pJRC2 (*CEN, chc1-1*), pJRC3 (*CEN, chc1-2*), pJRC4 (*CEN, chc1-3*), pJRC5 (*CEN, chc1-4*), or pJRC19 (*CEN, chc1-box*) were diluted to 1×10^7 cells/ml, and fourfold serial dilutions were plated as described in A and incubated at 30°C (or 25°C where indicated).

expressed in *chc1-1* and *chc1-3* cells was reduced four- to fivefold compared with HC expression from the *CHC1* allele, and expression of HC from the *chc1-2* and *chc1-4* alleles was reduced 7- to -10-fold relative to *CHC1*. However, at the permissive temperature of 25°C, each mutant HC protein facilitated robust growth and clathrin was associated with internal membranes as visualized by fluorescence microscopy. In addition, though the *chc1-2* and *chc1-4* caused similar, more severe phenotypes in the several tests for clathrin function, strains with these two alleles behaved differently in a number of assays. For example, the *chc1-4* allele, but not the *chc1-2* allele, was able to partially restore clathrin function in TGN sorting halo assays and calcofluor resistance at 25°C (Figure 9B; data not shown). Also, *chc1-4*, but not *chc1-2*, showed significant cortical clathrin recruitment in LAT-A and near normal clathrin patch dynamics by TIRFM at 25°C. Moreover, we found that complete removal of the TD, causes a null-like phenotype, but this HC is expressed at two- to threefold higher levels than the HC encoded by *chc1-4* (unpublished observations). Therefore, protein expression levels cannot fully account for the differences observed for each *chc1-TD-ts* allele.

Phenotypic differences caused by the *chc1-TD-ts* alleles may arise from specific alterations in the TD β -propeller structure and/or binding ability caused by each set of mutations. Both the *chc1-1* and *chc1-3* alleles possess the S75P mutation. Serine 75 is located at the beginning of blade 2 near the central axis of the propeller structure (Figure 1, A and B), and the substitution of proline may result in aberrant folding. The *chc1-3* allele encodes two additional substitutions (V201A and S338P); however, because HC protein levels were similar in cells expressing these alleles and the *chc1-3* mutation did not confer more severe phenotypes relative to *chc1-1*, these two residues probably do not contribute to major structural alterations.

The *chc1-2* allele contains two mutations (V37A and H289R) that could contribute to destabilization of the propeller. Valine 37 is found in propeller blade 1, but the side chain does not participate in the formation of either hydrophobic pocket that accommodates the hydrophobic side chains of CBM residues. Replacing one hydrophobic residue with another hydrophobic residue is not very likely to alter protein folding and result in TD destabilization. Histidine 289, identical between yeast and humans (Figure 1B), is located within a region of propeller blade 6 with a high degree of sequence identity between yeast and mammalian TDs. The histidine side chain lines the groove formed between blades 5 and 6, and mutating this residue to arginine might affect the folding of blade 6 or alter the packing of blades 5 and 6 with respect to one another. Alternatively, it is possible that the groove formed between blades 5 and 6 functions as a binding pocket for CBM-containing adaptors. Evidence to suggest this possibility comes from studies on tachylectin-2, whose five-blade β -propeller has binding sites for *N*-acetylglucosamine between each blade of its propeller (Beisel *et al.*, 1999), but further characterization of this allele is necessary to distinguish between these two possibilities.

Two missense mutations (L14P and H317R) were encoded by the *chc1-4* allele. Leucine 14 is identical between yeast and human HC (Figure 1B) and resides in a short α -helical segment of the loop connecting the final β -strand of blade 7 to the first β -strand of blade 1. Mutating this leucine residue to proline might affect the orientation of blade 1 with respect to the other six propeller blades. Histidine 317 is located in blade 7 with its side chain facing the groove between blades 7 and 1 of the TD propeller. An arginine residue at this position might affect the folding of blade 7 or alter the packing of

blades 7 and 1. The GBD-*chc1-4* TD fusion exhibited a slower mobility on gels, suggesting there is some structural alteration in this TD.

The *chc1-4* allele is unusual among the *ts* alleles in that clathrin recruitment to the cortex in LAT-A was relatively strong at 25°C, and the percentages of clathrin patch types seen by TIRFM were most similar to wild-type. Thus, for endocytosis at least, the *chc1-4* allele has properties of a true *ts* allele, and its unique properties will make it interesting for further study.

chc1-box Allele Mutations and Clathrin Function

HC expression from the *chc1-box* allele was expected to produce significant phenotypes in clathrin-dependent processes as the mutations introduced in this allele have been shown to disrupt interaction with adaptor proteins in *in vitro* binding assays (Goodman *et al.*, 1997; Miele *et al.*, 2004). However, *chc1-box* cells showed robust growth, TGN/endosomal sorting (α factor processing and calcofluor sensitivity), Ape1 processing (an indicator of Cvt/autophagy function), and α factor internalization. HCs encoded by *chc1-box* were also able to form clathrin-coated vesicles. In more refined analysis, we did notice partial impairment of clathrin recruitment to cortical sites in LAT-A, but compared with *chc1-1*, *chc1-2*, and *chc1-3*, as well as *chc1-4* at 37°C, this recruitment was quite strong and was not temperature-dependent. Also the TIRFM showed these cells have a higher percentage of immobile clathrin patches and unproductive patches that disappear without acquiring Abp1 and completing internalization than *CHC1* or *chc1-4* cells. Nevertheless, although more events with these altered dynamics were observed, the *box* mutant still internalized α factor at near wild-type rates. Together, these data indicate that the critical residues of the TD that constitute the known binding site for CBM and W-box motif containing adaptors are dispensable for clathrin function in yeast.

One possible explanation of these data is that other clathrin binding motifs in adaptor proteins might be able to compensate for the inability of the CBM to associate with the TD. Multiple copies of a short peptide motif ([D/S]LL) were identified in AP180, a synapse-specific adaptor protein that is critical for synaptic vesicle endocytosis, and in the large subunits of heterotetrameric adaptors AP-1, AP-2, AP-3, and AP-4 (Morgan *et al.*, 2000). The dileucine sequence of the (D/S)LL motif may participate in hydrophobic interactions with other grooves of the TD β -propeller, in which the hydrophobic nature of each groove is conserved (ter Haar *et al.*, 1998); however, the exact interaction site(s) for this motif have not been defined. Many yeast adaptor proteins, including the large β and γ subunits of AP-1 (Apl2 and Apl4, respectively), possess one or more copies of the (D/S)LL sequence, and each motif might have different groove specificities because there is greater degeneracy in the surrounding sequence compared with the CBM.

Alternatively, a recently identified binding site within the ankle region of HC for appendage domains from three subunits of clathrin adaptor protein complexes (tetrameric adaptor subunits AP-1 γ , AP-2 α , and AP-2 β) and the appendage domain from the monomeric adaptor protein Gga1 (Knuehl *et al.*, 2006) could compensate for the disrupted TD-binding site and facilitate clathrin function. Modeling and mutagenesis studies indicated that the HC ankle-appendage interaction, unlike clathrin box and W-box association with the TD, occurred over an extended interface, and two surface residues within the HC ankle (C682 and G710) were essential in the appendage-HC ankle interaction (Knuehl *et al.*, 2006). Sequence alignment of the ankle do-

main regions from yeast and mammalian HC shows that G710 is present in the yeast HC ankle (residue G716), whereas C682 is not conserved in the yeast sequence. This interaction could be important for some adaptors, suggested by the differences in interaction of Ent1 and Apl2 compared with Ent2 with our TD fragment that lacked the ankle region.

We favor the idea that these *chc1-box* allele data could be explained by an additional binding site(s) within the yeast TD that facilitates CBM-mediated adaptor binding. A TD fragment containing the *chc1-box* mutations was able to interact with Ent2, but this interaction was not nearly as strong as the interaction observed between the wild-type TD and Ent2. Interaction of both the wild-type TD and TD(*box*) domains with Ent2 was dependent upon the C-terminal CBM of Ent2, suggesting that one or more other sites on the TD, possibly other grooves between the propeller blades, are being used by the Ent2 CBM for its association: 1) a higher affinity interaction with the blade 1/2 groove and 2) a much lower affinity interaction with a second TD binding site. TD fragments with *chc1-box* mutations did not interact with either Ent1 or Apl2, but because the strength of the interactions between the wild-type TD and these two adaptors was much weaker when compared with the TD-Ent2 interaction, it is possible that utilization of a second TD binding site in the TD(*chc1-box*) fragment was not strong enough to detect using this assay. Alternatively, we cannot exclude that the blade 1/2 interaction was not completely ablated by the *chc1-box* mutations.

HCs encoded by the *chc1-box* allele were able to form CCVs, but these were not enriched for the AP-1 adaptor. Unlike mammalian AP-1 β , yeast Apl2 lacks the β -appendage domain and could not interact with the HC ankle binding site even if present in the yeast HC. Yeast Apl4, the AP-1 γ chain, does contain a relatively conserved γ -appendage domain that could mediate binding to the HC ankle (Hirst *et al.*, 2000; Knuehl *et al.*, 2006). However, this potential interaction must not be sufficient for stable incorporation of AP-1 into CCV in the *box* mutant. Likewise, if the TDs containing *chc1-box* mutations only possess a lower affinity binding site for adaptor-associated CBMs, this low-affinity interaction between AP-1 and clathrin was easily disrupted during the cell fractionation procedure. In previous studies, AP-1 incorporation into CCVs was almost abolished in cells containing mutations to the two CBM sequences in the β -subunit (Apl2) and a C-terminal deletion of the clathrin-binding region in the γ -subunit (Apl4) (Yeung and Payne, 2001). Thus, the reduced AP-1 association with CCVs isolated from *chc1-box* cells is consistent with these observations and suggests that stable incorporation of AP-1 into CCVs is mediated by the interaction of CBM-containing AP-1 subunits with the TD. Nevertheless, AP-1 interaction with clathrin seems to operate in vivo to abolish calcofluor sensitivity in the *box* mutant, indicating that the AP-1/clathrin-dependent, early endosome-to-TGN pathway used for the intracellular retention of Chs3 was functional in these cells. Likely, the AP-1 interaction occurs in the context of other coat components that might stabilize or augment its function during cargo selection steps and membrane invagination. In general, the clathrin box mutant analysis presented here provides additional support for the prevailing view that clathrin-mediated function is facilitated by the cooperative interactions of multiple accessory factors. Further work will be directed toward identification of additional HC adaptor CBM binding sites that are predicted from our studies.

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