The Hsp70 Homologue Lhs1p Is Involved in a Novel Function of the Yeast Endoplasmic Reticulum, Refolding and Stabilization of Heat-denatured Protein Aggregates

Nina Saris,* Heidi Holkeri,* Rachel A. Craven,‡ Colin J. Stirling,‡ and Marja Makarow*

*Institute of Biotechnology, University of Helsinki, Helsinki, Finland; and‡School of Biological Sciences, University of Manchester, Manchester M13 9PT, United Kingdom

Abstract. Heat stress is an obvious hazard, and mechanisms to recover from thermal damage, largely unknown as of yet, have evolved in all organisms. We have recently shown that a marker protein in the ER of Saccharomyces cerevisiae, denatured by exposure of cells to 50°C after preconditioning at 37°C, was reactivated by an ATP-dependent machinery, when the cells were returned to physiological temperature 24°C. Here we show that refolding of the marker enzyme Hsp150Δ–b-lactamase, inactivated and aggregated by the 50°C treatment, required a novel ER-located homologue of the Hsp70 family, Lhs1p. In the absence of Lhs1p, Hsp150Δ–b-lactamase failed to be solubilized and reactivated and was slowly degraded. Coimmuno-precipitation experiments suggested that Lhs1p was somehow associated with heat-denatured Hsp150Δ–b-lactamase, whereas no association with native marker protein molecules could be detected. Similar findings were obtained for a natural glycoprotein of S. cerevisiae, pro-carboxypeptidase Y (pro-CPY). Lhs1p had no significant role in folding or secretion of newly synthesized Hsp150Δ–b-lactamase or pro-CPY, suggesting that the machinery repairing heat-damaged proteins may have specific features as compared to chaperones assisting de novo folding. After preconditioning and 50°C treatment, cells lacking Lhs1p remained capable of protein synthesis and secretion for several hours at 24°C, but only 10% were able to form colonies, as compared to wild-type cells. We suggest that Lhs1p is involved in a novel function operating in the yeast ER, refolding and stabilization against proteolysis of heat-denatured protein. Lhs1p may be part of a fundamental heat-resistant survival machinery needed for recovery of yeast cells from severe heat stress.

When Saccharomyces cerevisiae cells are shifted from physiological temperature (24°C) to 50°C, they die. However, if preconditioned at the temperature where heat shock genes are activated, 37°C, they survive a subsequent thermal insult at 50°C and form colonies at 24°C. All organisms initiate a protective response to heat stress at characteristic temperatures in a similar way (1, 24, 28). Though the primary lesions causing cell death from thermal insult are not known, denaturation of proteins in various subcellular locations can be lethal, unless it is reversible (28, 30, 32, 34). Heat shock proteins, which are upregulated during the preconditioning period, have been assumed to function in the acquisition of thermotolerance (28, 30). However, little is known of how they protect or repair heat-damaged target proteins, though a wealth of information is available on their roles in folding of newly synthesized proteins under physiological conditions (12). Moreover, the Hsp26, Hsp70, and Hsp90 classes of heat shock proteins do not significantly affect thermotolerance in yeast (3, 31, 50). In contrast, Hsp104, an ATP-binding member of the Hsp100 (Clp) family of stress proteins, is essential for acquisition of thermotolerance in S. cerevisiae (35). In its absence, cytoplasmic protein aggregates generated by thermal insult to the cells at 50°C fail to be solubilized, and heat-inactivated mRNA splicing is not reactivated, resulting in cell death (29, 47).

The ER is the folding compartment for proteins destined to secretory organelles and to the exterior of the cell (14). When a newly synthesized polypeptide chain emerges in the ER lumen, it adopts a conformation dictated by its amino acid sequence. Folding is thought to be assisted by chaperones like the Hsp70 family member BiP/Kar2p, apparently by prevention of incorrect associations between nonnative chains (10). We showed recently that a fusion protein with β-lactamase activity, Hsp150Δ–β-lactamase, was inactivated in the yeast ER upon shift of the cells to 50°C after preconditioning at 37°C. When the cells were returned to 24°C, the enzyme was reactivated in an ATP-dependent way, even in the absence of de novo protein.
synthesis (21). In these studies the marker enzyme was accumulated in the ER of the temperature-sensitive secretion mutants sec23 or sec18 before the thermal insult at 50°C. When the marker enzyme was preaccumulated in the Golgi in sec7 or sec14 mutants, it failed to be reactivated, demonstrating that the refolding apparatus was confined to the ER (21). We show here that conformational repair of heat-damaged protein in the ER involves a novel member of the Hsp70 family, Lhs1p. Lhs1p was needed for reactivation, solubilization, and stabilization against proteolysis of the marker enzyme Hsp150Δ-β-lactamase, which was denatured and aggregated by thermal insult to living cells. Lhs1p appeared to have no significant role in folding and secretion of newly synthesized Hsp150Δ-β-lactamase molecules.

Materials and Methods

Yeast Strains and Media

The following yeast strains were grown at 24°C in shake flasks overnight to early logarithmic phase in YPD medium or synthetic complete (SC) medium lacking histidine (H645, H647, H656, and H660): H393 (Matα sec18-1 uaa3-52 trp1-289 leu2-3,112 URA3:HSP150-β-lactamase [43]), H486 (Matα sec18-1 his3-11,15 trp1-2 ade2-1 can1-100 hsp104:LEU2 [35]), H489 (Matα sec18-1 trp1-2 ade2-1 ura3-52 URA3:HSP150-β-lactama [12]), H503 (RCY127 Matα sec18-1 trp1-1 ade2-1 his3 ura3-52 lhis1::TRP1 [6]), H540 (RCY104 Matα ura3-52 ade2-1 trp1-1 lys2 his3 lhis1::TRP1 [6]), H541 (TR2 Matα ura3-52 ade2-1 trp1-1 lys2 his3 [27]), H602 (Matα ura3-52 ade2-1 trp1-1 lys2 his3 lhis1::TRP1:URA3:HSP150-β-lactamase), H604 (Matα ura3-52 ade2-1 trp1-1 lys2 his3::URA3:HSP150-β-lactamase), H621 (Matα sec18-1 ura3-52 ade2-1 trp1-1 his3 lhis1::TRP1 URA3:HSP150Δ-β-lactamase), H645 (Matα sec18-1 ura3-52 ade2-1 trp1-1 his3 lhis1::TRP1 URA3:HSP150Δ-β-lactamase [pRC42]), H656 (Matα sec18-1 ura3-52 ade2-1 trp1-1 his3 lhis1::TRP1 URA3:HSP150Δ-β-lactamase [pRC51]), H606 (Matα ura3-52 ade2-1 trp1-1 lys2 his3 lhis1::TRP1 [pRC51]), and H600 (Matα sec18-1 trp1-1 ade2-1 his3 ura3-52 lhis1::TRP1 [pRC51]).

Strains H602, H604, and H621 were created by integrating the HSP150Δ-β-lactamase gene in plasmid pKTH4544 (43), into the genome of strains H540, H541, and H503, respectively. Centromeric plasmid pRC2 containing the LHIS gene (6) was introduced into strain H621, creating strain H645, and centromeric plasmid pRC51 containing the c-myc-LHIS gene (6) was introduced into strains H621, H640, and H503 to create strains H647, H649, and H650, respectively. Transformations were done with the lithium acetate method (15). Strain H534 (Matα sec18-1 ade2-1 his3-11,15 trp1 hsp104:LEU2 URA3:HSP150Δ-β-lactamase) was created by mating strains H393 and H645 and dissecting the spores of the resulting diploid. The 4-kb BamHI/SacI fragment containing the LHIS gene was removed from pRC40 (6) and ligated into BamHI/SacI-digested pRS313 (CEN HIS3 [42]), to create pRC42. A 4.1-kb BamHI/SacI fragment containing a modified copy of the LHIS gene encoding Lhs1p tagged with the c-myc epitope was removed from pRC45 (6) and ligated into pRS313 (CEN HIS3) to create pRC51.

Metabolic Labeling and Immunoprecipitation

Metabolic labeling of cells (10⁵/200 μl) was with 20 μCi of [35S]methionine/cysteine (1,000 Ci/mmol; Amersham International, Buckinghamshire, UK) in SC medium lacking methionine and cysteine. The labelings were terminated with NaN₃, and the culture medium and cell lysate samples were immunoprecipitated as described (19). Briefly, cells were lysed mechanically with glass beads in NET buffer (0.05 M Tris-HCl, pH 8.0, containing 0.1 M NaCl, 5 mM EDTA, 1% NP-40, and 100 U/ml of aprotinin) in the presence of 2 mM PMSF. The lysates were boiled and preclariified for 1 h at 4°C with protein A-Sepharose (Pharmacia LKB Biotechnology, Inc., Piscataway, NJ). The samples were immunoprecipitated with anti-β-lactamase antiserum (1:100) or anti-carcboxypeptidase Y (CPY) antiserum (1:100) and protein A-Sepharose for 2 h at 4°C. After washing with diluted NET buffer (1:1), wash buffer (0.1 M Tris-HCl, pH 7.5, containing 0.2 M NaCl, 2 M urea, and 0.5% Tween-20), and with 0.1% SDS, the precipitates were analyzed by SDS-PAGE.

Coimmunoprecipitation and Western Analysis

Cells were lysed with glass beads in the presence of protease inhibitors as above, but in the presence of Triton X-100 and absence of SDS and boiling. The lysates were precleared with preimmune serum (1:100) and protein A-Sepharose and incubated overnight with anti-β-lactamase antiserum (1:100), anti-Cpy antiserum (1:100), or preimmune serum (1:100) at 4°C, followed by a 3-h incubation with protein A-Sepharose at 4°C. The precipitates were washed twice with NET buffer containing 10 mM CaCl₂ and lacking EDTA, and twice with the same buffer diluted 1:1 and subjected to SDS-PAGE. Proteins were blotted from gel onto nitrocellulose membrane (Hybond-C Extra; Amersham International) and immunostained with anti-β-lactamase antiserum (1:6,000) or anti-Cpy antiserum (1:1,000) and alkaline-phosphatase-conjugated anti-rabbit antibody (1:7,500; Promega Corp., Madison, WI), or with monoclonal anti-c-myc antibody (1:10,000; Chemicon International, Temecula, CA) and alkaline-phosphatase-conjugated anti-mouse antibody (1:3,000; Promega Corp.).

Sucrose Gradient Centrifugation

For determination of β-lactamase activity, duplicate cell samples (25 × 10⁵/0.5 ml of YPD or SC medium) were incubated at indicated temperatures. After termination of the incubation with NaN₃, the cells were separated from the medium, and cell-associated and secreted β-lactamase activity was assayed using nitrocefin as a substrate as described (43). Invertase synthesis was induced by incubating cells (5 × 10⁴/0.5 ml) in YPD medium containing 0.1% glucose, and intracellular and cell wall-bound invertase activity of duplicate samples was measured as described in (19). Thermotolerance was assayed by incubating duplicate cell samples (3 × 10⁶/ml) in YPD medium (except strain H647 in SC medium lacking histidine) in Wassermann tubes successively at 37 and 50°C (35). After cooling on ice, the cells were diluted and plated for 3–4 d at 24°C. Cycloheximide (CHX), DTT, and NaN₃, were from Sigma Chemical Co. (St. Louis, MO) and used in final concentrations of 100 μg/ml, 20 mM, and 10 μM, respectively. BSA and porcine thromboglobulin were from Sigma Chemical Co. SDS-PAGE was in 8% reducing gels.

Results

Effect of Lhs1p on Reactivation of Heat-denatured Hsp150Δ-β-lactamase In Vivo

Our assay to study heat-denaturation and refolding of ER-located proteins in living S. cerevisiae cells is presented at the top of Fig. 1. Cells grown at 24°C are incubated in liquid medium for an h at 37°C. The marker protein Hsp150Δ-β-lactamase is expressed under the heat-activated HSP150 promoter in a temperature-sensitive mutant (sec18), where ER-derived transport vesicles are unable to fuse with Golgi membranes at 37°C. The 37°C incubation thus preconditions the cells to survive a subsequent thermal insult at 50°C, upregulates the synthesis of the marker protein, and causes its retention in the pre-Golgi compartment. The cells are then incubated for 20 min at 50°C, resulting in inactivation of the β-lactamase fusion protein, and shifted to 24°C to monitor reactivation of the marker enzyme. In tens of experiments, 20–70% of the...
original β-lactamase activity was resumed in 3–6 h at 24°C in the presence of CHX (21). The variation in the reactivation yields evidently reflects the vulnerability of the cells to extreme temperature. Hsp150Δ-β-lactamase consists of Escherichia coli β-lactamase fused to the COOH terminus of the Hsp150Δ-carrier, an NH₂-terminal 321–amino acid fragment of the yeast secretory glycoprotein Hsp150 (20). The carrier is needed for translocation and to promote proper folding of foreign proteins that by themselves are not secretion-competent in yeast (25, 43).

To study whether Lhs1p has a role in refolding of proteins denatured by thermal insult to living cells, Hsp150Δ-β-lactamase was expressed in S. cerevisiae sec18 cells lacking a functional LHS1 gene because of targeted gene disruption (strain H621, lhs1-sec18). When H621 cells were incubated at 37°C, β-lactamase activity accumulated inside of the cells (Fig. 1 A), to a somewhat lower level than in wild-type cells (Fig. 1 B). Hsp150Δ-β-lactamase adopts an enzymatically active conformation upon translocation into the ER and concomitant disulfide formation. The ER form of Hsp150Δ-β-lactamase is primary O-glycosylated at many serines and threonines of the carrier portion and migrates in SDS-PAGE like a protein of ~110 kD, whereas the mature protein with extended O-glycans migrates like a 145-kD protein (21, 43). Shift of the cells from 37 to 50°C abolished the β-lactamase activity. When CHX was added to inhibit protein synthesis and the cells were shifted to 24°C for recovery, very little of the initial β-lactamase activity was resumed in 6 h (Fig. 1 A, closed circles). Parallel cells, which received NaN₃ at the time of shift to 24°C, served as a negative control: no reactivation was observed in the absence of ATP (open circles). H393 cells, which harbored the wild-type genomic LHS1 gene and the sec18 mutation, served as a positive control: 38% of the initial β-lactamase activity, which had accumulated in the ER before inactivation at 50°C, was recovered in 6 h (Fig. 1 B, closed circles), and NaN₃ inhibited the reactivation (open circles). We returned one copy of the wild-type LHS1 gene on a centromeric plasmid to strain H621, creating strain H645 (lhs1-LHS1+sec18). Reactivation of heat-denatured Hsp150Δ-β-lactamase was rescued to the level observed in the wild-type strain H393, again in an ATP-dependent fashion (Fig. 1 C). In none of the strains was β-lactamase activity found in the culture medium. These experiments show that Lhs1p did not protect the marker protein from thermal inactivation but was involved in reactivation of the heat-denatured marker protein.

**Aggregation and Solubilization of Heat-denatured Hsp150Δ-β-lactamase**

Next, we studied whether the thermal treatments caused aggregation of Hsp150Δ-β-lactamase. H393 cells (sec18) were 35S-labeled for 1 h at 37°C (Fig. 2 A), a parallel cell sample was incubated thereafter for 20 min at 50°C (Fig. 2 B), and a third sample was allowed to recover for 6 h at 24°C in the presence of CHX (Fig. 2 C). A fourth sample received NaN₃ for the time of recovery (Fig. 2 D). The cells were lysed under nondenaturing conditions and subjected to velocity sedimentation centrifugation in sucrose gradients. The fractions plus the pellet material (lane P) were subjected to immunoprecipitation with anti-β-lactamase antiserum and SDS-PAGE analysis. After labeling at 37°C, Hsp150Δ-β-lactamase was detected in the top fractions, indicating that it was soluble (Fig. 2 A). When immunoprecipitated from the gradient fractions, the protein migrated in SDS-PAGE as a doublet of ~95 and 110 kD. Thermal insult to the cells caused Hsp150Δ-β-lactamase to form aggregates since it was detected across the whole gradient and in the pellet fraction (Fig. 2 B).
covery period at 24°C, it sedimented again mainly in the top fractions, showing that it was largely solubilized. Solubilization required metabolic energy, since it was inhibited in the presence of NaN_3 (Fig. 2D). Since the sec18-1 mutation blocks fusion of ER-derived vesicles with the Golgi membranes, at least part of our marker protein may have resided in the vesicles in addition to the ER proper. Thus, we repeated the aggregation assay described above using sec23-1 cells (H486), where the marker is blocked quantitatively in the ER proper. Similar results were obtained as shown in Fig. 2, A–D, for the sec18-1 mutant. We have shown before that resumption of enzymatic activity of heat-inactivated Hsp150Δ-β-lactamase occurred equally efficiently in the sec23-1 and sec18-1 mutants (21).

Next we assayed the β-lactamase activity after fractionation. After incubation of H393 cells at 37°C, β-lactamase activity coincided with 35S-Hsp150Δ-β-lactamase in fractions 1–3 of the gradient (Fig. 3A). After the thermal insult, no activity could be detected in the fractions (Fig. 3B), whereas after 6 h at 24°C in the presence of CHX, ~22% of the original β-lactamase activity reappeared in the top fractions (Fig. 3C). We conclude that our marker protein was soluble when accumulated in the ER, aggregated by thermal insult to the living cells, and resolubilized in an ATP-dependent fashion, even in the absence of de novo protein synthesis, during recovery of the cells at 24°C.

In the absence of the LHS1 gene (strain H621, lhs1Δ sec18), 35S-Hsp150Δ-β-lactamase labeled at 37°C was soluble and migrated in SDS-PAGE again as a doublet of 95 and 110 kD (Fig. 4A). After the 50°C treatment, it was found in aggregates (Fig. 4B), and after the recovery period it was barely detectable (Fig. 4C). After the thermal insult and the recovery period, very little of the marker protein could be immunoprecipitated. Exposure time for B and C was five times longer than for A. Fractions 1–3 contained β-lactamase activity when the cells had been incubated at 37°C, whereas after thermal insult and recovery, no activity could be detected in the gradients (not shown). In the rescue strain H645 (lhs1Δ LHS1+ sec18), ATP-dependent solubilization of the Hsp150Δ-β-lactamase–containing aggregates occurred similarly as in strain H393.
(see Fig. 2; not shown). Thus, heat-denatured Hsp150Δ–β-lactamase remained in aggregates and was susceptible for proteolysis in the absence of Lhs1p. Degradation in this experiment occurred mainly by cellular proteases after cell lysis since less degradation occurred in vivo, as shown below. We conclude that thermal insult caused aggregation of ER-located Hsp150Δ–β-lactamase, whether Lhs1p was present or not. In the absence of Lhs1p, the Hsp150Δ–β-lactamase–containing aggregates failed to be solubilized and were susceptible to proteolytic degradation in vitro.

Coimmunoprecipitation of Lhs1p with Heat-denatured Hsp150Δ–β-lactamase and Pro-CPY

Coimmunoprecipitation experiments were performed to study whether Lhs1p was associated with Hsp150Δ–β-lactamase. H647 cells (hs1Δ c-myc–LHS1 sec18) were incubated successively for 1 h at 37°C (Fig. 5, A and B, lanes 1), 20 min at 50°C (lanes 2), and with CHX at 24°C for 3 h (lanes 3) or 6 h (lanes 4). The cells were lysed under non-denaturing conditions and divided in two. One lysate set was immunoprecipitated with anti–β-lactamase antiserum (Fig. 5, A and B, lanes 1–4) and the other with preimmune serum (lanes 6–9). Both sets were divided in two and subjected to SDS-PAGE and Western analysis using anti–c-myc antibody (Fig. 5, A) or anti–β-lactamase antiserum (Fig. 5 B). After incubation of the cells at 37°C, no precipitation of c-myc–Lhs1p with anti–β-lactamase antiserum could be detected (Fig. 5, A, lane 1). However, after thermal insult at 50°C and recovery at 24°C, c-myc–Lhs1p was coimmunoprecipitated with anti–β-lactamase antiserum (lanes 2–4). The total cell lysate samples (Fig. 5 A, lanes 5 and 10) show that a minor portion of Lhs1p was coprecipitated with Hsp150Δ–β-lactamase. Anti–β-lactamase antiserum precipitated Hsp150Δ–β-lactamase of ~110 kD from all samples (Fig. 5 B, lanes 1–4). Neither Lhs1p (Fig. 5 A, lanes 6–9) nor the marker protein (Fig. 5 B, lanes 6–9) were precipitated with preimmune serum. Thus, Lhs1p appeared to occur in association predominantly with heat-denatured marker protein molecules. Whether the associ-
Normally, CPY is synthesized as a precursor, which is for Hsp150 CPY of 62 kD. Similar results were obtained for pro-CPY reported to the vacuole, where it is processed to mature coreglycosylated in the ER (pro-CPY of 67 kD), and then trans-
tered to the vacuole, where it is processed to mature CPY of 62 kD. Similar results were obtained for pro-CPY as for Hsp150Δ-β-lactamase. After incubation of H660 cells (lhs1 Δ c-myc LHS1 sec18) at 37°C, very little c-myc-

| Lanes | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 |
|-------|---|---|---|---|---|---|---|---|---|
| IP α-CPY | Lhs1p |
| IS α-c-myc | |

| Lanes | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 |
|-------|---|---|---|---|---|---|---|---|---|
| IP α-CPY | CPY |
| IS α-CPY | IgG |

Figure 6. Coimmunoprecipitation of Lhs1p with pro-CPY. (A and B) H660 (lhs1 Δ c-myc LHS1 sec18) cells were incubated as in Fig. 5 for 1 h at 37°C (lanes 1 and 5), 20 min at 50°C (lanes 2 and 6), and at 24°C with CHX for 3 h (lanes 7 and 7) or 6 h (lanes 4 and 8). After lysis of the cells under nondenaturing conditions, the samples were divided into two. One set of samples was immu-
oprecipitated (IP) with anti-CPY antiserum (α-CPY, A and B, lanes 1–4) and the other with preimmune serum (pim; A and B, lanes 5–8). Lanes 9 contained total cell lysate. After SDS-
PAGE and blotting, the proteins were immunostained (IS) using anti-c-myc antibody (α-c-myc; A, lanes 1–9) or anti-CPY antiserum (B, lanes 1–9). c-myc Lhs1p (Lhs1p), pro-CPY (CPY), and IgG are indicated. Molecular mass markers (kD) are indicated on the left.

Stability of Heat-denatured Marker Protein

To study the stability of heat-aggregated Hsp150Δ-β-lactamase in the absence of Lhs1p in vivo, strain H621 (lhs1 Δ sec18) was labeled with [35S]methionine/cysteine at 37°C (Fig. 7 A, lane 1). Parallel cells were then incubated at 50°C (Fig. 7, lane 2), or thereafter at 24°C for 2–8 h (lanes 3–6). The cell lysates were immunoprecipitated with anti-
β-lactamase antiserum, followed by SDS-PAGE analysis. Hsp150Δ-β-lactamase persisted for about 4 h at 24°C,
proteins for hours after shift to 24°C, and the amount of the marker protein remained virtually unchanged. In the absence of DTT, mature Hsp150Δ–β-lactamase of 145 kD was immunoprecipitated from the culture medium (Fig. 9 A, lane 1), and some 66- and 145-kD protein from the cell lysate (lane 2). The intracellular reduced molecules were difficult to detect (Fig. 9 A, lane 4), as noted before (43). When cells labeled under reducing conditions were washed and chased in the absence of DTT and presence of CHX, secretion of the marker protein was resumed, since it could be detected in the culture medium (Fig. 9 A, lane 5). Some 66-, 110-, and 145-kD proteins remained cell associated (Fig. 9 A, lane 6). Similar results were obtained for the parental strain H604: DTT prevented reversibly the secretion of the marker protein (Fig. 9 B), but less intracellular proteins were detected in the cell lysates as compared to H602 (Fig. 9 B, lanes 2, 4, and 6).

The role of Lhs1p in conformational maturation of pro-CPY was then studied. Both in strains H602 (lhs1Δ; Fig. 9 C) and H604 (WT; Fig. 9 D), pro-CPY was accumulated in the ER in the presence of DTT (lanes 1), and converted similarly to CPY after DTT was replaced by CHX (lanes 2 and 3). In lanes 4, the cells were labeled in the absence of DTT, and thus mature CPY could be detected because of its rapid transport to the vacuole. From these data we conclude that Lhs1p had no significant role in translocation or conformational maturation of Hsp150Δ–β-lactamase and pro-CPY under our experimental conditions.

Acquisition of Thermotolerance in the Absence of Lhs1p

Next we studied whether Lhs1p affected the ability of yeast cells to acquire thermotolerance. Strain H540 (lhs1Δ), the parental strain H541 (WT), and strain H656 (lhs1Δ c-myc–LHS1+) were incubated for different times at 37°C growing in size apparently because of some increase in O-glycosylation. During the next 4 h, it was largely degraded. In the rescue strain H647 (lhs1Δ c-myc–LHS1+ sec18), the amount of the marker protein remained virtually unchanged (Fig. 7 B), as shown before for strain H393 harboring the genomic LHS1 gene (21). Degradation of the marker in the absence of Lhs1p was due to abrupt cell death, since heat-treated H621 cells were capable of protein synthesis for hours after shift to 24°C (see below). To confirm this, we studied the stability of Hsp150Δ–β-lactamase in a strain lacking a functional HSP104 gene (H534), which does not acquire thermotolerance at 30°C in spite of preconditioning at 37°C (35) (see below). In this strain, the thermal treatments inhibited protein synthesis irreversibly (see below). Hsp150Δ–β-lactamase 35S-labeled at 37°C remained after the 50°C treatment apparently undegraded at 24°C at least for 8 h (Fig. 7 C). These data show that Lhs1p suppressed in vivo degradation of the heat-denatured marker protein.

Role of Lhs1p in De Novo Folding and Secretion of Hsp150Δ–β-lactamase and Pro-CPY

Lhs1p is required for translocation into the ER of a subset of proteins (2, 6). Thus, we studied the role of Lhs1p in translocation, folding, and secretion of de novo–synthesized Hsp150Δ–β-lactamase in more detail. Strain H602 (lhs1Δ) (Fig. 8 A) and its parental strain H604 (WT) (Fig. 8 B), both wild-type for secretion, were incubated at 37°C, and the β-lactamase activities of the culture medium samples (Fig. 8, closed circles, EX) and lysed cell samples (open circles, IN) were determined. In both strains, β-lactamase activity externalized to the culture medium increased linearly, a little faster in the parental strain than in the lhs1Δ mutant. In the parental strain, little intracellular activity accumulated (0.12 U/ml in 90 min), indicating efficient secretion of the marker enzyme. In the lhs1Δ mutant, twice as much intracellular activity accumulated during the incubation, and its level in the beginning of the experiment was high. This suggests that a fraction of the newly synthesized marker protein remained cell-associated in active form and gradually built up an intracellular pool in the absence of a functional LHS1 gene.

Proteins that normally acquire disulfide bonds are retained in the yeast ER if cotranslational disulfide formation is prevented by incubating cells with a reducing agent like DTT, though the secretory machinery remains functional. When oxidizing conditions are reestablished by removing DTT, the sulfhydryls are oxidized, and the proteins undergo conformational maturation resulting in resumption of secretion (19, 44). To study whether Lhs1p was involved in conformational maturation of newly synthesized proteins, H602 (lhs1Δ) cells were 35S-labeled in the presence of DTT to allow translocation, while folding of Hsp150Δ–β-lactamase was inhibited. Immunoprecipitation with anti–β-lactamase antiserum of the culture medium sample showed that very little of the marker protein was secreted (Fig. 9 A, lane 3). In the absence of DTT, mature Hsp150Δ–β-lactamase of 145 kD was immunoprecipitated from the culture medium (Fig. 9 A, lane 1), and some 66- and 145-kD protein from the cell lysate (lane 2). The intracellular reduced molecules were difficult to detect (Fig. 9 A, lane 4, as noted before (43). When cells labeled under reducing conditions were washed and chased in the absence of DTT and presence of CHX, secretion of the marker protein was resumed, since it could be detected in the culture medium (Fig. 9 A, lane 5). Some 66-, 110-, and 145-kD proteins remained cell associated (Fig. 9 A, lane 6). Similar results were obtained for the parental strain H604: DTT prevented reversibly the secretion of the marker protein (Fig. 9 B), but less intracellular proteins were detected in the cell lysates as compared to H602 (Fig. 9 B, lanes 2, 4, and 6).

Figure 8. Secretion of β-lactamase activity. Strains (A) H602 (lhs1Δ) and (B) H604 (WT) were incubated at 37°C. The β-lactamase activity of the culture medium (closed circles, EX) and lysed cells (open circles, IN) was determined and plotted against incubation time.

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and thereafter for 20 min at 50°C and plated at 24°C (Fig. 10 A). More than 50% of H541 (Fig. 10 A, open circles) and H656 cells (closed circles) formed colonies when pre-conditioned for 45 min at 37°C before the 50°C treatment, whereas only 4% of H540 cells (Fig. 10 A, squares) survived. Strain H454 lacking a functional HSP104 gene (Fig. 10 A, triangles) served as a negative control: only 0.1% of them were able to form colonies after the thermal treatments. When the assay was performed by preincubating strains H540 (Fig. 10 B, lhs1Δ; squares) and H656 (lhs1Δ LHS1Δ; circles) for 45 min at 37°C followed by different times at 50°C, lack of LHS1 again caused a 10-fold decrease in survival.

Though only 10% of lhs1Δ cells, as compared to wild-type cells, were able to form colonies after the thermal insult, most of them remained biochemically active for several hours after the 50°C treatment as shown by de novo synthesis of invertase during the recovery period. H621 cells (lhs1Δ sec18; Fig. 11 A) and H393 cells (sec18; Fig. 11 B) were incubated successively for 1 h at 37°C, 20 min at 50°C, and 3 h at 24°C. The cells were then shifted to low glucose medium to derepress the synthesis of invertase (Fig. 11 B, arrow), which is normally destined to the cell wall. Determination of invertase activity in the cell wall (Fig. 11, A and B, closed circles, EX) and inside of the cells (open circles, IN) showed that similar amounts of enzyme were synthesized and secreted in the presence and absence of Lhs1p. Both strains secreted similar amounts of active invertase also at 24°C in the absence of any thermal treatments (Fig. 11, C and D). In contrast, no invertase synthesis during a recovery period of 6 h could be detected in strain H454 (hsp104Δ) (not shown). We conclude that synthesis and secretion, but not repair of the heat-damaged marker protein, functioned for many hours during the recovery period at 24°C in the absence of Lhs1p. Decreased thermotolerance of lhs1Δ cells, as measured by plating efficiency, was thus
not due to abrupt cell death soon after the thermal insult, but to later events resulting finally in failure of colony formation in >90% of the cells.

**Discussion**

The open reading frame of the LHS1 gene (YKL 073) was discovered in the Yeast Genome Sequencing Project. The translated amino acid sequence begins with a potential NH2-terminal signal peptide and terminates with an ER-retrieval signal HDEL. The ER-located Hsp70 cognate BiP/Kar2p has 25% identical and 50% similar amino acids with Lhs1p (33). LHS1 encodes a glycoprotein of 113–119 kD, is located in an ER-like compartment, and is required for translocation of a subset of proteins (2, 6). The LHS1 gene and Lhs1p are also called SSI1 and Ssi1p, respectively (2). We show here that Lhs1p is involved in conformational repair and stability of heat-damaged protein in the yeast ER.

**Figure 10.** Effect of Lhs1p on acquisition of thermotolerance. (A) Strains H541 (WT, open circles), H540 (lhs1−, squares), H656 (lhs1− c-myc–LHS1+, closed circles), and H454 (hsp104−, triangles) were incubated for different times at 37°C and then for 20 min at 50°C, and plated at 24°C. (B) Strains H656 (closed circles) and H540 (squares) were incubated for 45 min at 37°C and then for different times at 50°C, and plated at 24°C. The colonies were counted to determine the percentage of survival, which was plotted against incubation times.

**Figure 11.** Invertase synthesis during recovery at 24°C. (A) Strains H621 (lhst− sec18) and (B) H393 (sec18, indicated WT) were incubated successively for 1 h at 37°C, 20 min at 50°C, and 3 h at 24°C in YPD medium containing 2% glucose. The cells were shifted to YPD medium containing 0.1% glucose (arrow in upper scheme of experimental design) and incubated further at 24°C. In C and D, H621 and H393 cells, respectively, were incubated at 24°C in YPD medium containing 0.1% glucose (arrow in lower scheme). Cell samples were withdrawn for determination of intracellular (open circles, IN) and externalized (closed circles, EX) invertase activity. The relative activities are plotted against incubation time.
In the absence of a functional \textit{LHS1} gene, Hsp150\(\Delta\)-\(\beta\)-lactamase, aggregated and denatured by thermal insult to the cells, failed to be solubilized and reactivated like in wild-type cells.

According to coimmunoprecipitation experiments, Lhs1p was associated with heat-denatured Hsp150\(\Delta\)-\(\beta\)-lactamase, whereas association with native marker protein molecules could not be detected. Lhs1p could be immunoprecipitated even with a natural glycoprotein of \textit{S. cerevisiae}, pro-CPY, predominantly after thermal insult. Also, pro-CPY was damaged by the thermal insult to the cells since it became secretion incompetent (Saris, N., and M. Makarow, to be published). Whether association of the heat-damaged marker proteins with Lhs1p was mediated by other components is not known. Similar studies demonstrated earlier that BiP/Kar2p also could be coimmunoprecipitated with heat-denatured Hsp150\(\Delta\)-\(\beta\)-lactamase, and to a lesser extent with the native marker protein, whereas no association with protein disulfide isomerase could be detected (21). Heat-denatured Hsp150\(\Delta\)-\(\beta\)-lactamase thus occurred in aggregates containing at least Lhs1p and BiP/Kar2p. The role of BiP/Kar2p in refolding of heat-damaged Hsp150\(\Delta\)-\(\beta\)-lactamase could not be studied in vivo because it is required for translocation. Under conditions where BiP/Kar2p is nonfunctional, the marker protein is unable to translocate into the ER, and under normal conditions the marker protein escapes from the ER immediately after translocation (not shown). An ER-located hsp70-like stress protein grp170 has been found in association with BiP and immunoglobulin in CHO cells (5, 23). Sequence comparisons suggest that grp170 could be the mammalian homologue of yeast Lhs1p, raising the possibility that this novel molecular chaperone is ubiquitous amongst eukaryotes.

Though heat-damaged aggregated Hsp150\(\Delta\)-\(\beta\)-lactamase was solubilized and part of its enzymatic activity resumed, it remained in the ER at least for 6 h of recovery at 24°C (21). We have found that different secretory proteins, damaged by the thermal insult to the cells while residing in the ER, resume secretion competence with widely varying kinetics and efficiency (Saris, N., and M. Makarow, manuscript in preparation). Apparently, the structural features of Hsp150\(\Delta\)-\(\beta\)-lactamase were not compatible with secretion competence at the time points examined.

In the presence of Lhs1p, Hsp150\(\Delta\)-\(\beta\)-lactamase remained stable during the recovery period at 24°C, whereas it was slowly degraded in vivo in the absence of Lhs1p. Lately, degradation of mutant pro-CPY was shown to be retrotranslocated from the ER back to the cytosol for degradation by the proteasome complex (16). Since retrotranslocation of protein aggregates is unlikely, degradation of the heat-denatured marker protein probably occurred by some other mechanism, perhaps in the ER lumen or by autophagy (22, 46). Recently, a misfolded soluble protein was shown to be targeted to the vacuole for degradation (18). It has been suggested that heat shock proteins dissolve protein aggregates and facilitate protein degradation (13, 17). Newly synthesized misfolded proteins are degraded in the mitochondrial matrix by the PIM1 protease, if they are maintained in a soluble state by the chaperones mt-Hsp70 and Mdj1p (49). Lhs1p did not promote degradation of heat-damaged Hsp150\(\Delta\)-\(\beta\)-lactamase but suppressed its degradation. Exposed internal domains of pro-insulin were shown to display signals for degradation in the ER of mammalian cells. BiP has been suggested to mask these domains, suppressing the degradation of unfolded pro-insulin molecules (39).

Lhs1p thus appears to be involved both in translocation and in conformational repair of heat-damaged protein. Yeast BiP/Kar2p also has been assigned two functions. It is generally needed for translocation (36, 48) and for conformational maturation of newly synthesized pro-CPY (44) and Hsp150\(\Delta\)-\(\beta\)-lactamase (Holkeri, H., E. Paunola, E. Jämäsi, and M. Makarow, manuscript submitted for publication). In the latter studies, the marker proteins were translocated into the ER in reduced form in the presence of DTT and did not adopt a transport-competent conformation after reestablishment of oxidizing conditions unless BiP/Kar2p was functional. To compare the functions of BiP/Kar2p and Lhs1p, we studied the role of Lhs1p in folding of newly synthesized protein. Lack of Lhs1p did not inhibit translocation, conformational maturation, or intracellular transport of Hsp150\(\Delta\)-\(\beta\)-lactamase, pro-CPY, and invertase. Thus, Lhs1p differed from BiP/Kar2p in that it had no significant role in folding or secretion competence of our newly synthesized marker protein, but instead was required for its conformational repair and stability after heat denaturation. This suggests that the repair machinery in the ER may have distinct features as compared to the chaperone apparatus involved in folding of newly synthesized polypeptides. Newly synthesized unfolded polypeptides are fed to the chaperone machinery during translocation domain by domain, whereas the repair machinery deals mostly with full-length protein molecules that have been folded before stress-induced unfolding and aggregation. Frydman and Hartl (8) found that molecular chaperones interact differently with newly translated polypeptides and their chemically denatured counterparts in the eukaryotic cytosol. The in vivo kinetics of folding and conformational repair of Hsp150\(\Delta\)-\(\beta\)-lactamase differed strikingly: de novo folding occurs apparently in seconds, whereas conformational repair took several hours. Although recovery of yeast cells from thermal insult is efficient, it is remarkably slow. We found bulk protein synthesis to be inhibited for 2–3 h and fully recovered only after 6 h (21). Inhibition of protein synthesis for hours after severe heat stress has been noted also in mammalian cells (26). Evidently, in the absence of protein synthesis, survival requires refolding of heat-denatured proteins that perform vital functions.

Chaperone-dependent refolding of heat-denatured proteins has been shown before to occur in vivo in the cytosol of yeast and mammalian cells, and in \textit{E. coli}. Heat-denatured firefly luciferase was reactivated in \textit{S. cerevisiae} in an Hsp104-dependent manner (29). In \textit{E. coli}, refolding of \(\lambda\) repressor protein and luciferase were dependent on the chaperones DnaK (Hsp70), DnaJ (Hsp40), and GrpE (9, 40). In vitro, the DnaK/DnaJ/GrpE chaperone system and the chaperonins GroEL/GroES (Hsp60) are able to reactivate heat-damaged or chemically denatured protein (11, 45, 51). The human cytosolic chaperone Hsp90 was shown to convert chemically denatured \(\beta\)-galactosidase to a folding-competent state, whereafter the substrate protein was refolded by Hsp70 and hdj-1. It was suggested that Hsp90
maintains nonnative proteins in a soluble state in heat-stressed cells, whereas Hsp70 and hdj-1 refold them at normal temperature (7). As Bip/Kar2p and other Hsp70 chaperones require DnaJ/Hsp40 chaperones for activity (4, 37, 41), Lhs1p may need accessory proteins as well.

In the absence of Lhs1p, the ability of the yeast cells to form colonies after thermal insult was reduced 10-fold, whereas in the absence of Hsp104, thermotolerance is reduced 1,000-fold (35). Thermal insult did not result in abrupt death of the lhs1Δ mutant cells since they were capable of protein synthesis and secretion for many hours thereafter, and thus cell death must have occurred because of later events. In contrast, in the hsp104Δ mutant protein synthesis was completely and irreversibly inactivated by the 50°C treatment. The upstream sequence of LHS1 contains an unfolded protein response element. Indeed, LHS1 was upregulated under conditions where unfolded proteins accumulated in the ER, and moreover, the absence of Lhs1p caused induction of the unfolded protein response (2, 6). However, LHS1 is not upregulated at 37°C, whereas HSP104 is strongly induced by elevated temperature (35). A mitochondrial member of the Cip/Hsp100 family of S. cerevisiae, Hsp78, was recently found to be needed for maintenance of respiratory competence and resumption of mitochondrial protein synthesis after thermal insult at 50°C; however, it did not affect acquisition of cellular thermotolerance (36). Chaperones that repair vital heat-damaged proteins evidently occur in all compartments of the cell. Lhs1p located in the ER may be part of this fundamental survival system, which is needed for cells to recover from heat stress causing protein denaturation.

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