Grp170 and Hsp110 proteins constitute two evolutionary distinct branches of the Hsp70 family that share the ability to function as nucleotide exchange factors (NEFs) for canonical Hsp70s. Although the NEF mechanism of the cytoplasmic Hsp110s is well understood, little is known regarding the mechanism used by Grp170s in the endoplasmic reticulum. In this study, we compare the yeast Grp170 Lhs1 with the yeast Hsp110 Sse1. We find that residues important for Sse1 NEF activity are conserved in Lhs1 and that mutations in these residues in Lhs1 compromise NEF activity. As previously reported for Sse1, Lhs1 requires ATP to trigger nucleotide exchange in its cognate Hsp70 partner Kar2. Using site-specific cross-linking, we show that the nucleotide-binding domain (NBD) of Lhs1 interacts with the NBD of Kar2 face to face, and that Lhs1 contacts the side of the Kar2 NBD via its protruding C-terminal α-helical domain. To directly address the mechanism of nucleotide exchange, we have compared the hydrogen-exchange characteristics of a yeast Hsp70 NBD (Ssa1) in complex with either Sse1 or Lhs1. We find that Lhs1 and Sse1 induce very similar changes in the conformational dynamics in the Hsp70. Thus, our findings demonstrate that despite some differences between Hsp110 and Grp170 proteins, they use a similar mechanism to trigger nucleotide exchange.

The Hsp70 chaperones are essential components of the cellular machinery that controls the conformational states of proteins. They are involved in diverse functions, including folding of proteins, transport of proteins across membranes, and regulation of signal transduction components (1, 2). The common activity of Hsp70 chaperones that underlies these diverse functions is the transient binding to peptide segments of protein substrates. The association of Hsp70 with substrates is controlled by its ATPase cycle. When ATP is bound to the N-terminal nucleotide-binding domain (NBD), interdomain communication ensures that the C-terminal substrate-binding domain (SBD) exhibits low affinity for substrates, which consequently bind and release with high rates. Hydrolysis of ATP converts Hsp70 to the ADP-bound conformation, which traps the substrate with higher affinity. Exchange of ADP for ATP and concomitant substrate release completes this chaperone cycle.

The chaperone cycle of Hsp70s is regulated by cofactors that facilitate either ATP hydrolysis or exchange of ADP for ATP. ATP hydrolysis is stimulated by J-domain-harboring co-chaperones of the Hsp40 class that provide substrate specificity to the Hsp70 chaperone by delivering clients. Release of substrate from Hsp70 is facilitated by nucleotide-exchange factors (NEFs) that accelerate nucleotide dissociation, thereby allowing ATP to rebind. NEFs function by associating with and stabilizing specific conformations of the Hsp70 NBD that exhibit low affinity for nucleotide. Rebinding of ATP to the Hsp70 results in dissociation of the NEF.

Several structurally unrelated proteins function as NEFs for Hsp70s. Recent structural and biochemical analysis of the NEFs GrpE, Bag1, Bag2, HspBP1, and Hsp110 have revealed that each of these NEFs uses a unique Hsp70 NBD interaction interface (3–9). However, despite unique modes of binding, the actual mechanisms used to trigger nucleotide release fall into two principle classes. GrpE, Bag1, Bag2, and Hsp110 all promote nucleotide release by stabilizing similar Hsp70 NBD conformations that involve tilting NBD lobe II outwards. This tilting opens the NBD structure and facilitates nucleotide exchange (3–5, 7–9). In contrast, HspBP1/Fes1 associates with lobe II subdomain IIB and elicits a near global loss of tertiary structure of the NBD and thereby promotes nucleotide exchange (6, 9).

The eukaryotic Hsp110 proteins have recently been characterized as NEFs for Hsp70s in the cytosol (10–12). The Hsp110s are members of the Hsp70 superfamily of proteins and share overall structure with canonical Hsp70s but differ by an extended SBD with an acidic loop region inserted between the terminal strands of their β-sheet subdomain as well as by an extended flexible C terminus (13, 14). The yeast Saccharomyces

*This work was supported in part by Deutsche Forschungsgemeinschaft Grant SFB638 and a grant from the Fonds der Chemischen Industrie (to B. B.). The on-line version of this article (available at http://www.jbc.org) contains supplemental Figs. S1 and S2. To whom correspondence should be addressed: Stockholm University, The Wenner-Gren Institute, Cell Biology, Svante Arrheniusväg 16–18 ES, SE-10691 Stockholm, Sweden. Tel.: 46-8-159837; Fax: 46-8-159837; E-mail: claes.andreasson@cellbio.su.se. Supported by a fellowship from the Boehringer Ingelheim Fonds.

1 The abbreviations used are: NBD, nucleotide-binding domain; SBD, substrate-binding domain; NEF, nucleotide exchange factor; HX, hydrogen amide exchange; BMH, bis(maleimido)hexane; FOA, 5-fluoroorotic acid; UPR, unfolded protein response; ER, endoplasmic reticulum; MABA-ADP, N<sub>6</sub>-(4-N'-methylanthraniloylaminobuty)l-8-aminoadenosine 5'-diphosphate.
cerevisiae harbors two highly homologous members of the Hsp110 family, Sse1 and Sse2 (15).

Sse1-ATP has been crystallized both alone and in complex with mammalian Hsp70s (7, 8, 13). The crystal structures together with biochemical analysis (9) of the interaction with its cognate Hsp70 Ssa1 have revealed how Hsp110 and the Hsp70 partners interact in the nucleotide exchange reaction. Sse1 has to bind nucleotide to fold into a conformation that is capable of associating with the Hsp70 NBD (26). In the complex formed, the NBDs of Hsp70 and Hsp110 interact face to face in such a way that lobe I of the Hsp70 NBD contacts lobe II of the Hsp110 NBD. A second contact is formed between the side of Hsp70 NBD lobe II and the Hsp110 α-helical SBD. As an outcome Hsp110 embraces the Hsp70 NBD. The multiple site binding results in a tilting of Hsp70 NBD lobe II leading to an opening of the Hsp70 structure.

In the endoplasmic reticulum, Sil1 and Grp170s are NEFs that function together with the Hsp70 chaperone BiP/Kar2 (16, 17). Sil1 is a homologue of HspBP1, whereas the Grp170s are members of the Hsp70 superfamily. Grp170s carry extensions of their primary sequence compared with Hsp70s including a predicted loop structure in their SBD and an extended C terminus (14). In yeast, the single representative of the Grp170s is termed Lhs1 (18). Cells carrying lhs1 inactivating mutations are viable but display partial defects in Hsp70-dependent translocation and an induction of the unfolded protein response (UPR) (18–20). ER folding stress is sensed by the kinase Ire1, which then triggers the UPR (21, 22), resulting in up-regulation of a number of genes including SIL1 and LHS1 (18, 20, 23). Cells carrying double lhs1 sil1 mutations are not viable suggesting a strict requirement for NEF activity for proper Hsp70 function in the ER (23).

Currently, it is not clear how Grp170s interact with Hsp70s and by what mechanism they induce nucleotide exchange. Given that both the cytoplasmic Hsp110s and the ER-localized Grp170s are members of the Hsp70 superfamily it is tempting to speculate that they utilize a common NEF mechanism. However, available phylogenetic comparisons do not suggest that Grp170 and Hsp110 are more related to each other than they are to other members of the Hsp70 superfamily (14). Moreover, their ATPase activities are oppositely regulated upon interaction with their cognate Hsp70s; Lhs1 displays stimulation of its ATPase activity when interacting with Kar2 (17), whereas the ATPase activity of Sse1 is dramatically reduced upon interaction with Ssa1 (10). Hence, it is not clear whether Hsp110s and Grp170s utilize common or distinct mechanisms to trigger nucleotide exchange of their respective Hsp70s. In this study, we employed a combination of genetic, biochemical, and biophysical approaches to show that Hsp110s and Grp170s employ a similar mechanism to trigger nucleotide exchange in Hsp70s.

**EXPERIMENTAL PROCEDURES**

**Yeast Strains and Plasmids**—CAY1171 is a derivative of strain Y04922 (EUROSCARF, Germany) generated by introducing the complementing LHS1-plasmid pCA716 and deleting SIL1 using an hphMX4 cassette PCR amplified from pAG32 (24) with primers, GGTCGGATTCTCTCCCATATAA-TTTCGAGCGCCTATCTTCGAGCTGAAGCTTCGT-AGCG and CTATGAGCCATGGGTGCGCAAAAGATCTAA-GTGTCGCTGCTGATAAGCATAGCCACTAATGTAGTGA-TCTG. CAY1172 was constructed by replacing IRE1 with a PCR-generated LEU2 fragment (primers, ATCGGTCTACT-TCGAAGAAACATTTGATTTGACATGCTGTTT-GTGTAAGAAATATCTTGCCCGAG and GGTGAAGT-AATCGTAAAATCCATCGGGTACCGGCCCCATTAGT-GTCGCTATATCTCACCCTATGAACATATGCC). Plasmids pCA715 (HHS3 LHS1) and pCA716 (URA3 LHS1) contain similar LHS1 locus fragments amplified from genomic DNA using primers, GGCGGGGATCCTGCCCTTGTTTTGTCATAGTC and GGCGGACTAGTAAAGATGAAACAGTGTTT-AACT, and BamHI/Spel-cloned into pCA503 (9) and PRS316 (25), respectively. Mutations in LHS1 were introduced in pCA715 using oligonucleotide-based site-directed mutagenesis (lhs1–2, N608Y/E611A, GTTACCTGCAAGACGTAT-AATTTCCTTG; Ihs1–3, A325T/N326A, CTTGCCCTA-GAAGGCGTGCTTAATTTAAC; Ihs1–4, N415V/N417G, TCATGACAGGAAACTCCTCAACAC). Plasmid pCA708 is a derivative of the His6-Smt3 vector pCA528 (26) and contains the Kar2 NBD coding sequence (amino acid residues 43–426) that was PCR amplified using primers, CCAGTGGGTCTCAGGTGGTGCCGATGATGT-AGAAAACATCG and GGCGGGGATCCTAGTAAAGACAGTGTTT-AACT, and cloned using Bsal/BamHI. Plasmid pCA709 contains LHS1 (from amino acid residue 21) and was cloned similarly using primers CCAGTGGGTCTCAGGTGGTG-TCGCTTATTAGGTGGTATACG and GGCGGCGGAT- CCTAATAATCATCATGCAAAATGT. Plasmid pCA717 expresses Lhs1 carrying a C-terminal Strep-Tag II and was constructed by ligating a NdeI/BamHI-restricted PCR product (primers, CCGCCCCCATATGGGTGGCGGTATTAGGTGG-TATTACG and CCGCGGGATCTCCTCTTCCTGAACCTCGG- GTGGTGCGCTCAAGCGTAAATTTATCTCACTAGCAAAAT- TGCTTC) into pET24a (Novagen).

**Protein Expression and Purification**—Lhs1 (pCA709) and Sse1 and the NBDs of Kar2 (pCA708) and Ssa1 were expressed with an Ulp1 cleavable N-terminal His6-Smt3 tag and purified using Ni-IDA matrix (for details, see Ref. 26). During Ulp1 cleavage of His6-Smt3-Lhs1 1.75 M urea was added to the buffer. Proteins were made nucleotide-free by dialysis against buffer containing 50 mM EDTA for 24 h and then dialyzed extensively against EDTA-free buffer. Complexes of the Ssa1 NBD and Lhs1 or Sse1 for hydrogen amide exchange (HX) experiments were purified using a double tag strategy (26). Briefly, crude lysates from cells expressing His6-Smt3-Ssa1 and Lhs1 (pCA717) or Sse1 carrying a Strep-Tag II at their C termini were mixed and formed complexes were isolated by serial Ni-IDA and StrepTactin (IBA, GmbH, Goettingen, Germany) chromatography.

**Nucleotide Release Assays**—Nucleotide release was measured using the fluorescently labeled analogue of ADP, N2-(4-′N′-methyleneanthraniloylaminobuty)-8-aminoadenosine 5′-diphosphate (MABA-ADP; TriLink Technologies Inc., San Diego, CA), and the stopped flow instrumentation SX-18MV from Applied Photophysics (Surrey, UK). The method has been described elsewhere (26, 27).
**Interaction Experiments**—To assess complex formation between Lhs1 and Kar2 NBD, 30 μM Lhs1, supplemented with and without a 2-fold excess of ATP, was incubated with 30 μM His6-Smt3-Kar2 NBD (made nucleotide free using EDTA containing buffer) for 10 min at 30 °C in LBW150 buffer (40 mM Hepes-KOH, pH 7.4, 150 mM KCl, 5 mM MgCl2, 5% (v/v) glycerol) + 0.1% (v/v) Triton X-100. Alternatively, 39 μM Kar1 NBD was incubated with 13 μM Lhs1 wild-type or Lhs1 mutant proteins in the presence of 30 μM ATP. Co2+-NTA-Sepharose was added and proteins were allowed to bind for 20 min at 4 °C. After washing two times with LWB150 and two times with LWB500 (40 mM Hepes-KOH, pH 7.4, 500 mM KCl, 5 mM MgCl2, 5% (v/v) glycerol) + 0.1% (v/v) Triton X-100, proteins were eluted by digestion with Ulp1 protease for 10 min at 30 °C and analyzed on 10% SDS-PAGE that was stained with Coomassie Brilliant Blue.

**Cross-linking Experiments**—Complexes for cross-linking with bis(maleimido)hexane (BMH) (Thermo Scientific) were obtained by incubating 6 μM Lhs1/Sse1 with equimolar concentrations of Kar2/Ssa1 NBDs overnight at 4 °C in the presence of 1 mM ATP and 6 μM tris(2-carboxyethyl)phosphine. Cross-linking was carried out on ice by addition of 74 μM BMH and the reactions were stopped after 60 min by addition of sample buffer containing 50 mM dithiothreitol. The reactions were analyzed on 10% SDS-PAGE stained with Coomassie Brilliant Blue.

**Hydrogen-Deuterium Exchange Experiments and Mass Spectrometry**—HX experiments were performed in a similar manner as described previously (9, 28, 29). Briefly, 100–350 pmol of purified monomeric Ssa1 NBD, or Ssa1 NBD complex with Lhs1 or Sse1 were preincubated for 3 min at 30 °C and diluted 20-fold into D2O-based buffer (25 mM Hepes-KOH, pH 7.6, 50 mM KCl, 5 mM MgCl2) to initiate amide proton-deuterium exchange. The exchange reaction was stopped at defined times by addition of 1 volume of ice-cold quench buffer (0.4 M potassium phosphate buffer, pH 2.2). Quenched samples were immediately injected into the high pressure liquid chromatography setup, subjected to online peptic digest, and analyzed on an electrospray ionization quadrupole time-of-flight mass spectrometer (QSTAR Pulsar; Applied Biosystems) as described (29, 30). The data processing was performed according to Ref. 29. D2O buffer for the HX experiment was prepared by using 99.85% D2O (Euriso-top), lyophilized, and redissolved five times in equal D2O volumes.

**Sequence Analysis**—The primary sequence of Lhs1 and Lhs1 orthologues from *Kluyveromyces lactis*, *Ashbya gossypii*, and *Candida albicans* were analyzed using PELE protein structure prediction. ClustalW (31) was used to globally align the sequences as well as for local alignments of sequences corresponding to SBDα and SBDβ (defined by secondary structure prediction and the fact that the sequences follow NBD sequences). Manual motif screening of Lhs1 and Sse1 ClustalW alignments identified the conserved pairs Ala325/Asn326, Asn415/Asn417, and Asn608/Glu609.

**RESULTS**

**Lhs1 Shares Subdomain Organization with Sse1**—We wanted to perform biochemical analysis of Lhs1 and were therefore in need of a model that defined the subdomain organization of Lhs1. Initially, we attempted to use global sequence alignment methodology to gain information regarding potential sequence conservation between Hsp110s and Grp170s. Although sequence alignment programs could successfully align the more conserved NBD sequences of Hsp70s, Hsp110s, and Grp170s, they failed to properly align sequences derived from the poorly conserved SBDs. We hypothesized that Grp170s shared SBD subdomain structure with Hsp110s and tested this hypothesis by employing local instead of global sequence alignments of the SBD sequences of Lhs1 and Sse1 (see “Experimental Procedures”). Moreover, we applied secondary structure predictions to define SBDα and SBDβ. The result from this analysis is consistent with Lhs1 sharing subdomain organization with Sse1 (Figs. 1A and supplemental S1). Specifically, we find that Lhs1 carries a putative flexible loop structure right before the last β-sheet of SBDβ (Fig. 1A, L), a motif shared with Hsp110s that is not present in Hsp70s. Moreover, the sequence directly following the SBDβ appears to constitute the three α-helices B, C, and DE that are present in Hsp70s and Hsp110s.

We asked if residues involved in the nucleotide exchange activity of Sse1 also are conserved in Lhs1. Polier et al. (8) documents three pairs of Sse1 residues involved in the interaction between Sse1 and Hsp70. Two of these pairs are localized in the NBD and one pair in the α-helix B of the SBDα (Fig. 1B). Sse1 NBD residue pairs Ala280/Asn281 and Thr365/Asn367 correspond to Ala326/Asn327 and Asn415/Asn417, respectively, in the Lhs1 sequence. Note that Sse1 Thr365 is not perfectly conserved with Lhs1 Asn415. However, in other fungal homologues, for example, such as *C. albicans* Lhs1, position 415 carries a threonine residue (data not shown). The Sse1 SBDα residue pair Asn572/Glu575 also appears to be conserved in fungal Lhs1 homologues despite the considerable variation in the α-helix B sequence (Fig. 1A, Asn608/Glu611). Our analysis suggests that Lhs1 carries corresponding residues at positions known to be important for Sse1 NEF activity.

**Residues Conserved between Lhs1 and Sse1 Are Important for NEF Activity**—We tested if the identified residues in Lhs1 were important for NEF activity by mutational analysis. First, we introduced mutations in the *LHS1* coding sequence on a yeast single copy plasmid in the context of the native promoter and terminator sequences. All three alleles (*lhs1–2*, *N608Y/E611A*, SBDβ: *lhs1–3* A326T/N327A, NBD; and *lhs1–4*, N415V/N417S, NBD) carry corresponding substitution mutations used in the previous functional analysis of Sse1 (see supplemental Fig. S1 and Ref. 8). We tested if the alleles could complement the *lhs1Δ sil1Δ* non-growth phenotype of strain CAY1171. This strain carries a complementing *URA3*-marked Lhs1-expressing plasmid that can be counterselected on 5-fluoroorotic acid (FOA) containing medium. Hence, cells will only grow on 5-FOA medium if they are supplied with a plasmid expressing at least partially active Lhs1. We introduced single copy plasmids harboring *lhs1–2* (N608Y/E611A), *lhs1–3* (A326T/N327A), and *lhs1–4* (N415V/N417S) in the context of the endogenous *LHS1* locus and found that all three alleles complemented the phenotype (Fig. 2A, left panels). However, *lhs1–3* cells appeared to undergo a phase of adaptation right after they
lost expression of wild-type \textit{LHS1} because they transiently formed unevenly sized colonies when they were transferred to selective FOA medium (data not shown). We reasoned that this weak phenotype might represent cellular adaptation to low \textit{LHS1} NEF activity via the UPR regulation. Importantly, the \textit{LHS1} promoter itself is under UPR control (18, 20), which potentially allows \textit{lhs1} alleles encoding low activity to up-regulate their own expression. We tested this possibility by performing the complementation analysis of \textit{lhs1–2, lhs1–3, and lhs1–4} in \textit{lhs1–2, lhs1–3, and lhs1–4} in \textit{lhs1–4} cells that lack a functional UPR pathway (22, 23). Both \textit{lhs1–2} and \textit{lhs1–4} complemented the non-growth phenotype fully, whereas \textit{lhs1–3} cells exhibited a lower plating efficiency on selective FOA medium compared with nonselective SC medium (Fig. 2A, right panels). Upon closer inspection we observed that \textit{lhs1–3} cells exhibit a slow-growth phenotype (data not shown) that explains why FOA\textsuperscript{K} cells were underrepresented in the plasmid shuffle experiment, manifested as a lower plating efficiency on FOA medium.

Combined \textit{Lhs1} mutations such as \textit{lhs1–2,3} and \textit{lhs1–2,4} also complemented the \textit{lhs1–2,3} and \textit{lhs1–2,4} non-growth phenotype (data not shown). This observation contrasts the predicted analogous \textit{sse1} mutations, which exhibited non-complementation (8), suggesting either that the residues are not essential for NEF activity of \textit{Lhs1} or that little residual NEF activity is sufficient to maintain cellular growth.

Next, we assessed the impact of the mutations on \textit{Lhs1} NEF activity \textit{in vitro} using MABA-ADP release assays (Fig. 2B). In these assays preloaded complexes consisting of the fluorescent nucleotide derivative MABA-ADP and the NBD of Kar2 (2 \textmu M) were rapidly mixed with \textit{Lhs1} and dissociation of MABA-ADP was followed by a decrease of its fluorescent signal in stopped flow measurements. Both \textit{Lhs1–2} and \textit{Lhs1–4} exhibited a low activity, corresponding to approximately half of the wild-type \textit{Lhs1} activity. The activity impairment of \textit{Lhs1–3} was even more pronounced; the protein exhibited only barely detectable activity under identical conditions. The low activity of \textit{Lhs1–3} \textit{in vitro} correlates with the inability of the mutant to fully complement the \textit{in vivo} phenotype. The residual activity is appar-
ently sufficient to support growth of \( lhs1\Delta \; sil1\Delta \; ire1\Delta \) cells, which might either mean that the *in vitro* assay underestimates the proteins activity *in vivo*, or that very little NEF activity is required for growth. In any case, we and others have previously observed similar partial complementation of growth phenotypes when testing \( sse1 \) NEF mutants *in vivo* (8, 9). All mutant \( Lhs1 \) proteins appeared to be properly folded as judged by CD measurements (data not shown).

Our *in vivo* and *in vitro* mutational analysis suggests that \( Lhs1 \) depends on similar residues as \( Sse1 \) for its NEF activity. The residual NEF activity of \( Lhs1−2 \) (in \( SBD\alpha \)) and \( Lhs1−4 \) (in \( NBD \)) is expected if \( Lhs1 \), like \( Sse1 \), employs a large surface with multiple contact points to interact with the Hsp70 (8).

**Nucleotide Is Required for NEF Activity of \( Lhs1 \)—**In contrast to other characterized NEFs, \( Sse1 \) requires ATP binding to reach an active conformation that is capable of interacting with Hsp70 and trigger nucleotide exchange. We wondered also if \( Lhs1 \) required nucleotide for functional interaction with Kar2 and investigated this possibility by testing if \( Lhs1 \) could form a complex with Kar2 in the absence or presence of ATP. The NBD of \( Kar2 \) was incubated with nucleotide-free \( Lhs1 \) in the absence or presence of added ATP and immobilized onto cobalt affinity resin by means of an N-terminal His\(_6\)-Smt3 tag. After washing, Kar2 and bound \( Lhs1 \) were eluted by proteolytic severing of the His\(_{52}\)-Smt3 tag and analyzed by SDS-PAGE. In the absence of nucleotide, little or no \( Lhs1 \) was bound to Kar2, whereas the presence of ATP resulted in formation of substantial amounts of complex (Fig. 3A).

To directly assess if nucleotide is required for NEF activity of \( Lhs1 \), we measured MABA-ADP release from Kar2 under nucleotide-depleted conditions (no excess nucleotide to compete with the released MABA-ADP in the assay). In the absence of added nucleotide, 2 \( \mu \)M \( Lhs1 \) could only accelerate the basal release rate of Kar2 (2 \( \mu \)M) 1.4-fold, whereas preincubation of \( Lhs1 \) with 2 \( \mu \)M ATP resulted in 9-fold acceleration (Fig. 3B). Preincubation with 20 \( \mu \)M ATP further enhanced the activity to 13-fold acceleration of the basal release rate. The interaction experiments together with the activity determinations indicate that \( Lhs1 \) requires nucleotide to interact with Kar2 and trigger nucleotide release. During review of this manuscript a parallel study was published that also reported on an ATP requirement for \( Lhs1 \) NEF activity (32).

**\( Lhs1-Kar2 \) Association Depends on Residues Conserved between \( Lhs1 \) and \( Sse1 \)—**We wanted to test if the residues in \( Lhs1 \) that are important for NEF activity (Asn\(_{608}\)/Glu\(_{611} \), \( SBD\alpha \); Ala\(_{326}\)/Asn\(_{326} \), NBD; and Asn\(_{415}\)/Asn\(_{417} \), NBD) also are important for association of \( Lhs1 \) and Kar2. We undertook quantitative pulldown assays, where the Kar2 NBD was immobilized onto cobalt affinity resin by means of an N-terminal His\(_{52}\)-Smt3 tag (see above). We applied \( Lhs1 \) and our \( Lhs1 \) with amino acid substitutions that impair NEF activity to Kar2 and assayed the interaction after washing. \( Lhs1−2 \) (N608Y/E611A, \( SBD\alpha \)) and \( Lhs1−3 \) (A326T/N327A, \( NBD \)) exhibited approximately 15 and 4% ATP-dependent Kar2 binding compared with wild-type \( Lhs1 \) (Fig. 3C). \( Lhs1−4 \) (N415V/N417S, \( NBD \)) exhibited about 53% binding. In the case of \( Lhs1−3 \) and \( Lhs1−4 \) the Kar2 binding correlates directly with their determined NEF activity; \( Lhs1−3 \) is a poor binder and exhibits very little NEF activity and \( Lhs1−4 \) exhibits both substantial NEF activity and binding. Interestingly, \( Lhs1−2 \) displays NEF activity at the level of \( Lhs1−1 \) but a significantly lower level of complex formation. Hence, this observation implies that Hsp70 complex stability and NEF activity might not be directly coupled. A similar phenomenon has been reported previously for the \( Sse1-Hsp70 \) interaction (8).

**\( Lhs1 \) and \( Sse1 \) Contact the Hsp70 NBD Similarly—**We used site-specific cross-linking to investigate if \( Lhs1 \) and Kar2 form a complex with architecture reminiscent of the \( Sse1-Ssa1 \) complex. We have previously established that a cysteine residue replacing Asn\(_{283}\) in \( NBD \) lobe II of \( Sse1 \) forms a cross-link to a cysteine replacing Gln\(_{31}\) of the \( Ssa1 \) NBD using bismaleimide cross-linkers (9). This cross-link provides evidence that the NBD of \( Sse1 \) interacts face to face with the NBD of \( Ssa1 \). We used cysteine replacing Gln\(_{31}\) of the \( Ssa1 \) NBD using bismaleimide cross-linkers (9). This cross-link provides evidence that the NBD of \( Sse1 \) interacts face to face with the NBD of \( Ssa1 \), an observation consistent with recently published crystal structures (7, 8). We introduced cysteines at corresponding positions in \( Lhs1 \) (N326C) and the NBD of \( Kar2 \) (Q79C) and tested their ability to cross-link using BMH (10 Å length). Because both \( Lhs1 \) and \( Sse1 \) require ATP to form complexes with Hsp70s, the proteins were mixed with BMH in the presence and
absence of ATP to control for cross-linking specificity. We observed that Lhs1-N326C and Sse1-N283C formed ATP-dependent cross-links to both Kar2-C79C and Ssa1-Q31C (Fig. 4A), indicating that the NBDs of Lhs1 and Kar2 and Sse1 and Ssa1 are oriented similarly in their respective complexes. Furthermore, the data suggest that key interacting residues in Lhs1 and Sse1 are conserved.

Next, we wanted to investigate if Lhs1 SBD contacts the side of the Kar2 NBD lobe II in a fashion similar to how Sse1 contacts Ssa1 (see Fig. 1B). Aided by the recently published crystal structures of complexes between Sse1 and Hsp70s (7, 8), we introduced cysteines positioned in Lhs1 SBD (K643C) and at the side of Kar2 NBD lobe II (E349C) so that they potentially could be cross-linked using BMH. Again, we took advantage of the ATP requirement for efficient complex formation between Lhs1 and Kar2 to control for specificity and found that BMH cross-links Lhs1-K643C efficiently to Kar2-E349C in an ATP-dependent manner (Fig. 4B). Cross-links were not detected between Lhs1-K643C and Kar2-Q79C (Fig. 4B), providing another specificity control. Taken together, our cross-linking data indicate that Lhs1 interacts with Kar2 NBD using two contact surfaces similar to the architecture of the Sse1-Ssa1 complex.

**Lhs1 and Sse1 Trigger Nucleotide Exchange in Both Kar2 and Ssa1**—Because Lhs1 and Sse1 interacted in the cross-linking experiments with both the cognate and noncognate Hsp70s, we wished to test if they catalyze nucleotide exchange from both Ssa1 and Kar2. Using MABA-ADP release assays we found that 1 μM Lhs1 accelerated the basal nucleotide release rate from its cognate partner Kar2 6- and 11-fold from non-cognate Ssa1, both at 0.5 μM (Fig. 4C). Sse1 accelerated the basal release rate of cognate Ssa1 46-fold and of Kar2 3-fold under identical experimental conditions. Although both Lhs1 and Sse1 function less well with the non-cognate Hsp70s than the cognate interaction partners, we find that the basic features required for functional interaction with Lhs1 and Sse1 are conserved in both Kar2 and Ssa1.

**Lhs1 and Sse1 Employ the Same Mechanism to Trigger Nucleotide Exchange in Hsp70s**—The similarity in the interactions of Lhs1 and Sse1 with Hsp70s raises the possibility that the two NEFs employ the same mechanism to trigger nucleotide exchange in Hsp70s. We have previously devised a methodology that allows us to directly probe for NEF-induced conformational changes in the Hsp70 NBD using HX combined with mass spectrometry (9). Briefly, by measuring HX protection or deprotection of the Ssa1 NBD at a peptide level we can monitor conformational changes induced by the NEFs upon interaction with the Hsp70 NBD.

We wished to compare the specific patterns of HX protection and deprotection that Lhs1 and Sse1 induce in the NBD of Ssa1. The Ssa1 NBD was chosen for analysis because we have previously annotated all of its peptides and established the effect that several NEFs, including Sse1, induce on this NBD (9). Furthermore, our cross-linking analysis and activity assays demonstrate that Lhs1 functionally interacts with this particular Hsp70.

We purified nucleotide-free monomeric Ssa1 NBD and the NBD in complex with either Lhs1 or Sse1 (see “Experimental Procedures”) and subjected these proteins to HX in heavy water buffer for defined durations. Online pepsin digestion followed by mass spectrometry allowed us to identify Ssa1 NBD peptides and the extent of HX of each peptide at the different time points (Fig. 5A). Upon comparison of the HX behavior of each peptide derived from either monomeric Ssa1 NBD or the NBD in complex with either Lhs1 or Sse1 we found mainly three classes of peptides: (i) peptides that exhibited HX protection induced by both Lhs1 and Sse1 (Fig. 5A, left panel, peptide 23–39), (ii) peptides that were not affected by Lhs1 and Sse1 (Fig. 5A, middle panel, peptide 105–119), and (iii) peptides that exhibited HX deprotection induced by Lhs1 and Sse1 (Fig. 5A, right panel, peptide 215–229). Very few peptides exhibited different HX.
behavior induced by Lhs1 and Sse1. We mapped the changes onto a structural model of the Ssa1 NBD (9) derived from the crystallized bovine Hsc70 NBD (33) (Fig. 5B). Strikingly, for nearly all peptides analyzed, Lhs1 and Sse1 induced the same general pattern of HX protection and deprotection (Figs. 5B, and 6, and supplemental S2). Specifically, Lhs1 or Sse1 induced protection in subdomain I and deprotection of peptides in subdomain II, whereby deprotection was localized to the interface of subdomains IIA and IIB. The induced HX changes were more pronounced for Sse1 than for Lhs1, likely reflecting a higher affinity of Sse1 over Lhs1 for the Ssa1 NBD. One segment in subdomain IIB spanned by peptide 265/266–280 (Figs. 5B and 6), exhibited protection induced by Lhs1, whereas Sse1 induced a slight deprotection of the same segment. This is the only segment that exhibits differences in HX effects induced by Lhs1 and Sse1.

Taken together, the HX data for the interaction of Lhs1 with the Ssa1 NBD are consistent with the mechanism we previously derived for interaction of Sse1 with Ssa1 using HX and cross-linking (9). Two recently published crystal structures of the Sse1-Hsp70 complex show how Sse1 contacts the Hsp70 NBD and induces outward rotation of subdomain II (7, 8). Hence, our HX data indicate that Lhs1 and Sse1 employ a very similar mechanism to trigger nucleotide exchange from Hsp70s.

**DISCUSSION**

We have defined how the yeast Grp170 Lhs1 binds Hsp70s and how this interaction induces nucleotide release. Initially the published data on Lhs1 suggested some mechanistic differences to the Sse1/Ssa1 system (see below). Our study, however, demonstrates that the ER-localized Lhs1 shares both binding interface and NEF mechanism with the cytoplasmic Hsp110 Sse1. The main contact between Lhs1 and Kar2 is mediated via their NBDs so that they orient to face each other and Lhs1 NBD lobe II contacts lobe I of Kar2 (see Fig. 1B). A second contact between the protruding SBD/H9251 of Lhs1 and the side of Kar2 NBD lobe II induces tilting of lobe II thereby opening the Kar2 NBD. The resulting conformation is incompatible with high nucleotide binding affinity.

Our understanding of how Lhs1 interacts with Hsp70s is based on a series of experiments directly comparing the well characterized Sse1 with Lhs1. Key observations include sequence conservation between the NEFs, the finding that Lhs1 and Sse1 can be cross-linked to the Hsp70 NBD at similar residues, and that both NEFs induce near identical conformational dynamics of an Hsp70 NBD as judged from HX experiments. Given the similarities we conclude that...
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Lhs1 undergoes a similar conformational activation event upon protein acquires the active conformation (26). Apparently, the finding (32) provides further evidence for the NEF activity of Sse1. Also Sse1 requires nucleotide for its NEF activity because substitution of the interacting Sse1 residues almost abolishes the activity in vitro and results in inefficient complementation in vivo (8). When we introduced the corresponding substitutions in Lhs1 (Lhs1–2), the complex formation and NEF activity were reduced, but approximately half the activity remained and the allele complemented in vivo. In addition, combining the mutations, which abolishes the NEF activity of Sse1 (8), was in the case of Lhs1 tolerated in vivo. Although the extensive interaction interface between Hsp70 and NEF may account for some of the resilience observed in Lhs1, our findings raise the possibility that Lhs1 might employ additional residues to contact the side of the Hsp70 NBD. Indeed, we observe a peptide located nearby (residues 266–280) that exhibits protection from HX when the Ssa1 NBD is complex with Lhs1 but not when in complex with Sse1. Hence, this peptide defines a potential contact point for Lhs1 that is not used by Sse1.

Curiously, the ATPase activities of Lhs1 and Sse1 offer examples of similarities and differences between the two proteins. In this study we document that ATP binding of Lhs1 is a prerequisite for NEF interaction with the Kar2 NBD. A parallel study has addressed the same issue and provides further evidence for the finding (32). Also Sse1 requires nucleotide for its NEF activity (10, 26) and we have previously documented that the molecular basis for this is a two-step folding event upon which the protein acquires the active conformation (26). Apparently, Lhs1 undergoes a similar conformational activation event upon ATP binding. Hydrolysis of the bound ATP does not play a role for the NEF activity of the two proteins (12, 17). Despite this fact, both Lhs1 and Sse1 hydrolyze the ATP and their ATPase activities are affected differently by complex formation with Hsp70s. When Lhs1 forms a complex with Kar2, its ATPase rate is stimulated (17), whereas Hsp70 potently inhibits the ATP hydrolysis of Sse1 (10). At this point, there is no explanation for this apparent difference between Lhs1 and Sse1.

Our finding that the Grp170 Lhs1 and the Hsp110 Sse1 employ a similar mechanism to trigger NEF in Hsp70s addresses a controversy in the field. One opinion has been that Hsp110s and Grp170s share a common origin and therefore are expected to employ a similar mechanism (see for example, Ref. 34). However, because experimental evidence or even rigorous sequence alignments have been missing, another idea has been that Grp170s and Hsp110s are more different (see for example, Ref. 14). In such a scenario the NEF mechanisms of the two protein families might differ. Our experimental analysis of Lhs1 supports the notion that Grp170s and Hsp110s are quite similar. Because both Lhs1 and Sse1 require nucleotide for NEF activity it is likely that this feature is mechanistically and evolutionarily linked to the NEF activity.

The recent development in the understanding of how NEFs promote nucleotide exchange from Hsp70s has previously allowed us to propose two general classes of nucleotide release mechanisms (9): (i) catalysis of nucleotide release through a global loss of tertiary structure of the Hsp70 NBD (HspBP1/Fes1) and (ii) catalysis of nucleotide release through opening of the nucleotide binding cleft by tilting lobe II. Our data regarding Lhs1 place the Grp170s in the second group together with the structurally related Hsp110s and the structurally unrelated GrpE and Bag1 proteins. During writing of this article the NEF mechanism for the mammalian Bag2 protein was published (5). The authors propose that Bag2 might have been mis-annotated as a classical Bag-family member and document that Bag2 induces outward tilting of Hsp70 NBD lobe II in a different manner than Bag1. Thus, according to our proposed classification, Bag2 also belongs to the same functional NEF group as GrpE, Bag1, Hsp110s, and now Grp170s. Apparently, the limited conformational repertoire of the Hsp70 NBD drives evolution to produce similar nucleotide release mechanisms despite different NEF-binding modes.

**Acknowledgments**—We thank Matthias P. Mayer for helpful discussions and the mass spectrometry facility of the ZMBH for support.
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APRIL 16, 2010 • VOLUME 285 • NUMBER 16 JOURNAL OF BIOLOGICAL CHEMISTRY 12453