

2

3

6

9

10

ARTICLE IN PRESS

Available online at www.sciencedirect.com



BBAMCR-15764; No. of pages: 12; 4C:



Biochimica et Biophysica Acta xx (2008) xxx-xxx

Nuclear translocation of the phosphoprotein Hop (Hsp70/Hsp90 organizing protein) occurs under heat shock, and its proposed nuclear localization signal is involved in Hsp90 binding

Sheril Daniel^a, Graeme Bradley^{a,1}, Victoria M. Longshaw^a, Csaba Söti^b, Peter Csermely^b, Gregory L. Blatch^{a,*}

^a Chaperone Research Group, Department of Biochemistry, Microbiology and Biotechnology, Rhodes University, Grahamstown, South Africa ^b Department of Medical Chemistry, Semmelweis University, Puskin street 9, H-1088, Budapest, Hungary

Received 14 September 2007; received in revised form 11 January 2008; accepted 14 January 2008

11 Abstract

The Hsp70-Hsp90 complex is implicated in the folding and regulation of numerous signaling proteins, and Hop, the Hsp70-Hsp90 Orga-12 nizing Protein, facilitates the association of this multichaperone machinery. Phosphatase treatment of mouse cell extracts reduced the number of 13 Hop isoforms compared to untreated extracts, providing the first direct evidence that Hop was phosphorylated in vivo. Furthermore, surface 14 plasmon resonance spectroscopy (SPR) showed that a cdc2 kinase phosphorylation mimic of Hop had reduced affinity for Hsp90 binding. Hop 1516 was predominantly cytoplasmic, but translocated to the nucleus in response to heat shock. A putative bipartite nuclear localization signal (NLS) 17 has been identified within the Hsp90-binding domain of Hop. Although substitution of residues within the major arm of this proposed NLS abolished Hop-Hsp90 interaction as determined by SPR, this was not sufficient to prevent the nuclear accumulation of Hop under leptomycin-B 18 treatment and heat shock conditions. These results showed for the first time that the subcellular localization of Hop was stress regulated and that 19 the major arm of the putative NLS was not directly important for nuclear translocation but was critical for Hop-Hsp90 association in vitro. We 20propose a model in which the association of Hop with Hsp90 and the phosphorylated status of Hop both play a role in the mechanism of nucleo-21 cytoplasmic shuttling of Hop. 22

23 © 2008 Elsevier B.V. All rights reserved.

25 Keywords: Hop; Heat shock protein; Phosphorylation; Hsp90; Co-chaperone; Nuclear localization signal

26

24

27 **1. Introduction**

The *H*sp70–Hsp90 *O*rganizing *P*rotein, Hop, was first identified in yeast during a genetic screen for proteins involved in the heat shock response [1]. Hop has insignificant chaperoning abilities [2] and upon heat shock (HS) and viral transformation, no corresponding change in the steady state levels of this protein was detected, although an increase in mRNA levels of Hop in mammalian cells was observed [3,4]. Hop acts as a scaffolding protein, mediating the interaction of the molecular chapero- ³⁵ nes Hsp70 and Hsp90 through specific tetratricopeptide-repeat ³⁶ (TPR)-rich binding domains [5,6] to form the Hsp70–Hop– ³⁷ Hsp90 chaperone heterocomplex. These TPR domains char- ³⁸ acterize the various homologues of Hop, found in humans [3], ³⁹ mice [7], rats [8], insects [9], plants [10], parasites [11] and ⁴⁰ viruses [12]. Hsp70 binding is mediated primarily through the ⁴¹ N-terminal TPR domain of Hop (TPR1) [13] while a central TPR ⁴² motif-containing region (TPR2A) is known to be essential for ⁴³ Hsp90 binding [14,15]. The TPR-acceptor site on both Hsp70 ⁴⁴ and Hsp90 comprises an EEVD motif at the C-terminus [15,16]. ⁴⁵

In the yeast system, it was recently shown that the third TPR 46 domain, TPR2B is also directly involved in Hop's interactions 47 with Hsp70 and Hsp90, and that an isolated TPR2A domain is 48

^{*} Corresponding author. Tel.: +27 46 603 8262; fax: +27 46 622 3984. *E-mail address:* g.blatch@ru.ac.za (G.L. Blatch).

¹ Current Address: Department of Biochemistry and Microbiology, University of Fort Hare, Alice, South Africa.

unable to interact with Hsp90 unless combined with TPR2B
[17]. Deletion of a specific amino acid in the TPR2B domain of
yeast Hop (A438) showed the disruption of in vivo interaction
with Hsp90 [18]. The possibility of a network of interactions
between Hop and the chaperones Hsp70 and Hsp90, involving
domains other than the TPRs, have been suggested, which
support its functionality as a scaffolding protein [15,19].

Early studies reported the presence of human Hop in the 56 Golgi apparatus and small vesicles of normal cells, and in the 57nucleolus of SV40-transformed cells [3]. Immunoprecipitation 58studies using membranous fractions from mouse brain showed 59that a population of Hop is localized at the cell surface, where it 60 interacts with Prp^c (a protein whose expression is crucial to the 61 propagation of neurological disease, in particular, Prion disease) 62 both in vitro and in vivo. It has been proposed that Hop is taken 63 into the plasma membrane as part of a Prp^c -protein complex 64 [20]. Alternatively, Hop may be secreted by a pathway that is 65 distinct from the classical route of membrane translocation 66 through the ER and Golgi apparatus [21] since a transmembrane 67 domain or signal peptide for membrane transport is not found 68 within the amino acid sequence of Hop [4,20]. 69

An investigation into the subcellular localization of Hop in 70 mouse cells showed a predominantly cytoplasmic localization 71 under normal growth conditions [4]. Leptomycin-B treatment 72(nuclear export inhibition) of mouse fibroblast cells, resulted in 73a predominantly nuclear accumulation of Hop, suggesting that 74 the protein shuttled between the nucleus and the cytoplasm, 75with the export of Hop from the nucleus occurring at a faster 76 rate than its import [22]. Moreover, Longshaw et al. identified a 77 78 potential bipartite nuclear localization signal (NLS), which when fused to enhanced green fluorescent protein (EGFP), 79 resulted in the localization of EGFP to the nucleus [22], pro-80 viding evidence that this NLS was indeed functional on its own. 81 This putative NLS comprised of a short lysine arm (K222-82 K223), a 13-amino acid spacer region, followed by a major 83 lysine-rich arm (K237–K239). 84

Early studies done on yeast Hop [1] identified two major and 85 two minor isoforms of the protein with pIs ranging from 5.75 to 86 6.05, which implied that Hop was differentially phosphorylated. 87 Other studies showed shifts in the isoform composition of Hop 88 to the more acidic range after viral transformation and heat 89 shock [3,4]. Immediately upstream of the putative bipartite 90 NLS, two potential phosphorylation sites for casein kinase II 91 (CKII; S189) and cell division cycle 2 (cdc2; T198) kinase were 92identified using in vitro assays [23]. Despite speculations that 93 94 Hop is a phosphoprotein, based on indirect evidence, thus far there has been no direct evidence to show that Hop exists as a 95 phosphoprotein in vivo. 96

There is evidence to suggest that the localization of Hop is 97regulated by phosphorylation. Hydroxyurea (G1/S arrest) and 98 olomoucine treatment (cdc2 kinase inhibition) of mouse fi-99 broblast cells resulted in cytoplasmic and nuclear localization of 100Hop [22]. Experiments with phosphorylation mimics of Hop 101 (Hop-S189E and Hop-T198E) suggested that phosphorylation 102 of Hop by cdc2 kinase promoted the cytoplasmic retention of 103 Hop, whereas phosphorylation of Hop by CKII promoted nu-104 105clear localization [22]. Both these putative phosphorylation sites were found proximal to the Hsp90-binding TPR2A domain 106 of Hop, suggesting that Hop–Hsp90 interactions may also be 107 modulated by phosphorylation. We also noted that the proposed 108 NLS of Hop overlapped with its TPR2A domain. Moreover 109 residues within the spacer region of the NLS (K229, N233 and 110 K301) were shown to be crucial for Hop–Hsp90 interactions 111 [6,15]. Therefore, we hypothesize that the putative bipartite 112 NLS of Hop, and its phosphorylation status, play a significant 113 role in its subcellular localization and association with Hsp90. 114 In this report we have further characterized the putative bipartite 115 NLS of Hop, provided the first direct evidence that Hop was 116 phosphorylated in vivo, showed that heat shock affected the 117 subcellular localization and phosphorylation of Hop, and pro- 118 vided evidence that Hop–Hsp90 interactions involved residues 119 in the putative NLS and may be influenced by phosphorylation. 120

2. Experimental procedures

2.1. General

121 122

152

Common molecular biology procedures such as agarose gel electrophoresis, 123 ligation reactions, competent bacterial cell preparation, plasmid DNA isolation 124 and restriction enzyme digestion were performed according to standard proto- 125 cols [24]. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS- 126 PAGE) was according to Laemmli [25], and Western blotting according to 127 Towbin [26]. Luminol-based chemiluminescence detection reagents (ECL Ad- 128 vance Western Blotting Detection Kit; Amersham Biosciences) were used for 129 Western analysis. Protein estimation was done using the 2-D Quant Kit (Amer- 130 sham Biosciences). 131

2.2. Site-directed mutagenesis 132

All mutations were generated from the plasmids pGEX3X2000 and pSK- 133 mSTI1-EGFP, which contain the cDNAs encoding the full-length proteins mSTI1 134 (mouse homolog of Hop) as a C-terminal fusion with glutathione-S-transferase 135 (GST) and mSTI1 as an N-terminal fusion with enhanced green fluorescent protein 136 (EGFP), respectively. All mutations were carried out by site-directed mutagenesis 137 using a double-stranded whole plasmid linear amplification procedure (Quik- 138 Change mutagenesis kit; Stratagene). For screening purposes, silent mutations 139 were engineered to create restriction sites except where the desired codon change(s) 140 automatically generated restriction sites. The following mutant proteins were 141 generated: GST-Hop-K239A, GST-Hop/K237/238/239A, GST-Hop-T198E, GST- 142 Hop-S189E, Hop-K239A-EGFP, and Hop-K237/K238/K239A-EGFP. Mutations 143 were confirmed both by restriction enzyme analysis and by DNA sequencing. For 144 ease of manipulation, the mSTI1-EGFP encoding fragment in pSK-mSTI1-EGFP 145 was used as a template for mutagenesis before subcloning into pBCMGSNeo [27] 146 XhoI/NotI sites to produce pB-mSTI1-EGFP and its mutant derivatives, as de- 147 scribed previously [22]. Note: All plasmid constructs listed retain their original 148 cloning nomenclature which refers to mSTI1 (mouse homolog of Hop); however 149 the proteins they encode have been labeled Hop for this study so as to limit the 150 number of acronyms used for the same protein. 151

2.3. Production and purification of GST-Hop fusion proteins

Exponentially growing *Escherichia coli* XLI Blue cells carrying pGEX3X 153 derived plasmid constructs were induced for 5–6 h at 37 °C with 1 mM isopropyl- 154 1-thio- β -D-galactopyranoside (IPTG). The cells were harvested and lysed by 155 mild sonication in 0.01 culture volume of ice-cold phosphate-buffered saline 156 (PBS; 137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄, 1.4 mM KH₂PO₄, pH 157 7.3) containing 1 mM final concentration of phenylmethylsulfonyl fluoride 158 (PMSF). The sonicate was incubated, with gentle agitation, for 30 min at room 159 temperature after addition of Triton X-100 to 1% final concentration. The 160 extracts were clarified by centrifugation at 12,000 g for 20 min at 4 °C. 161 Aliquots of clarified extracts were added to 50% (w/v) slurry of glutathione- 162

163 agarose beads (2 mL bed volume per 100 mL of extract) previously equi-164 librated with PBS. Binding was allowed to occur for 1 h at 4 °C with gentle rocking. The beads were washed extensively with ice-cold PBS before the 165166 bound GST fusion proteins were eluted by adding appropriate volume of the elution buffer (10 mM reduced glutathione in 50 mM Tris-HCl, pH 8.0). 167 168 Eluted GST fusion proteins were analyzed on SDS-PAGE (12% acrylamide/ 169bis-acrylamide w/v). GST-Hop and its mutant derivatives were all successfully 170expressed and found to be soluble proteins, which were purified to at least 90% 171 purity.

172 2.4. Surface plasmon resonance spectroscopy

173Surface plasmon resonance (SPR) spectroscopy was performed using 174 a Biacore X apparatus (BIACORE, Sweden). A mixture of NHS/EDC (N-hydroxysuccinimide and 1-ethyl-3-[3-dimethylaminopropyl] carbodii-175 mide) was used to activate the surface of a CM5 sensor chip by modifica-176 tion of the carboxymethyl groups to N-Hydroxysuccinimide esters. Purified 177 178 rat liver Hsp90 (untagged) was injected into the first flow cell, such that the 179N-Hydroxysuccinimide esters reacted spontaneously with the amines on the 180 Hsp90 to form covalent links.

181 The second flow cell was used as a control with no protein immobilized on 182 it. Approximately 9165 response units (RU) of Hsp90 were immobilized onto the chip. After equilibration with running buffer (10 mM Hepes buffer con-183 184 taining 140 mM KCl and 0.001% Tween), increasing concentrations (0.2 µm, 1 µm and 5 µm) of GST-Hop and its derivatives were passed over the 185186 immobilized Hsp90 at a flow rate of $10 \,\mu L \,min^{-1}$. GST was passed through both cells and any non-specific binding to Hsp90 and/or the chip was monitored. All 187 experiments were performed at room temperature. The CM5 Sensor Chip 188 (Research Grade) was purchased from BIACORE (Sweden). The data was 189 190 analyzed using the BIAevaluation software version 3.0.

191 2.5. Hop isoform analysis by two-dimensional gel electrophoresis

To satisfactorily determine the in vivo phosphorylated isoform status of Hop, NIH 3T3 fibroblast cells were treated in various ways prior to preparation of a lysate for analysis by two-dimensional (2D) gel electrophoresis. The cells were either treated with phosphatase inhibitors (PI) okadeic acid (30 nM; specific for serine/threonine phosphatases) and sodium vanadate (2 mM; specific for tyrosine phosphatases), to limit or eliminate any dephosphorylation of Hop, or treated with shrimp alkaline phosphatase (SAP; 10 U) to dephosphorylate any 198 phosphorylated isoforms of Hop. The NIH 3T3 fibroblast cells treated for 199 isoform analysis were harvested by scraping into ice-cold PBS. The cell pellets 200 obtained were resuspended in two-dimensional (2D) lysis buffer (40 mM Tris– 201 HCL, 8 M Urea, 4% Triton X-100, 1 μ g/mL aprotinin and 100 μ g/mL PMSF) and 202 lysed on ice for 1 h. The lysate was centrifuged at 13,000 rpm for 30 min on a 203 benchtop microfuge to remove insoluble cell debris. Protein concentration was 204 determined using the 2D-Quant kit (Amersham Biosciences, USA), before 205 the protein samples were precipitated using the 2D clean-up kit (Amersham 206 Biosciences, USA).

For the first dimension separation by isoelectric focusing (IEF), the pre- 208 cipitated total protein samples were diluted in rehydration buffer (9 M Urea, 2% 209 Triton X-100, 2% Ampholytes, 0.002% bromophenol blue, 100 mM DTT) to a 210 final concentration of 1 µg/µL. A total protein sample (120 µg) was then applied 211 onto an immobilized linear pH gradient (IPG; pH 4-7) strip within the chambers 212 of a Zoom® IPGRunner[™] Cassette (Invitrogen) and rehydrated overnight as per 213 the manufacturer's instructions. Rehydrated IPG strips were focused by per- 214 forming IEF in distilled water. The applied electric potential (V) was increased in 215 a step-wise fashion as follows: 200 V (20 min), 450 V (15 min), 750 V (15 min) 216 and 1000 V (120 min), with a total power of 0.1 W strip⁻¹ and current, 0.05 mA 217 strip⁻¹. The second dimension was resolved by SDS-PAGE (12% acrylamide/ 218 bis-acrylamide w/v) at 200 V for 45 min. Western analysis was carried out on the 219 resolved proteins using antibodies specific to Hop (short chain recombinant 220 antibodies made to the C-terminal region of mSTI1, C-334; Antibodies By 221 Design, MorphoSys®; Germany). 222

2.6. Cell culture, transfections and preparation of cell lysates 223

Baby Hamster Kidney (BHK-21) cells and NIH 3T3 mouse fibroblast cells 224 were maintained in Dulbecco's modified Eagle's medium (Cambrex, USA) 225 supplemented with 10% calf serum, penicillin (100 U mL⁻¹) and streptomycin 226 (100 U mL⁻¹) (Highveld Biological PTY. Ltd., South Africa) in a humidified 227 atmosphere, at 37 °C with 5% CO₂. 228

Transient transfections were conducted on BHK-21 cells using IBA fect (for 229 magnet assisted transfection) and MA Lipofectin Enhancer (IBA, Germany) on 230 cells seeded to 60% confluency, according to the manufacturer's instructions. 231 Leptomycin-B (Synexa Life Sciences, South Africa) was used to arrest the 232 nuclear export of proteins; cells were treated with 2 ng mL⁻¹ of leptomycin-B 233 overnight at 37 °C. 234



Fig. 1. Hop was differentially phosphorylated in vivo at 37 °C and 42 °C. (A) The isoform composition of Hop after resolution by 2D gel electrophoresis in the pH range 4–7 of 120 μ g of total protein extract from NIH 3T3 cells (prepared as described in Experimental procedures), followed by Western analysis for the detection of Hop. The isoelectric focusing (IEF) and SDS-PAGE steps are shown. The upper panel shows the Western of lysates prepared from cells that were grown under 37 °C and treated with phosphatase inhibitors (37 °C+PI); the lower panel shows the Western of lysates prepared from cells that were also grown at 37 °C and treated with shrimp alkaline phosphatase (SAP) but not treated with PI (37 °C — PI+SAP). Approximate pIs for the prominent isoforms were estimated and are indicated at the top of each panel. The position of the isoforms with assigned pIs did not always coincide due to technical discrepancies such as slight changes in gel size. The major isoform under normal conditions, with an approximate pI of 6.1 was arbitrarily given the label '0'. Any isoform that was found to be more acidic than '0' was labeled with increasing positive numbers. Additional isoforms which were not assigned a pI value were labeled with an asterisk. This figure was representative of the results obtained from at least three independently conducted experiments. (B). The two panels show cell extracts prepared from cells that were incubated for 30 min at 42 °C. The upper panel shows the isoform composition of Hop in cells lysed in the absence of SAP but presence of PI and the lower panel, cells lysed in the presence of SAP. This figure was representative of the results obtained from at least three independently conducted experiments.

235

236

237 238

239

ARTICLE IN PRESS

S. Daniel et al. / Biochimica et Biophysica Acta xx (2008) xxx-xxx

Cells were fixed in 4% paraformaldehyde for 10 min and nuclei stained with

2 ng m L^{-1} 4',6-diamidino-2-phenylindole (DAPI; Sigma) in PBS. Slides were

mounted in fluorescent mounting solution (DAKO) and the fluorescent images

were viewed and captured using the Zeiss LSM 510 Meta confocal microscope

(40×oil immersion objective) and LSM 510 software (Zeiss).

2.7. Biochemical fractionation

BHK-21 cells were chemically fractionated using the ProteoExtract[®] Sub- 241 cellular Proteome Extraction Kit (Calbiochem) according to the manufacturer's 242 instructions. Proteins obtained from each fraction were resolved by SDS-PAGE 243



Fig. 2. Hop was localized to the nucleus under leptomycin-B treatment and heat shock conditions; the nuclear localization of Hop was not dependent on the major arm of the NLS. (A) BHK-21 cells were transiently transfected with pB-mSTI1-EGFP (encoding Hop-EGFP) and two Hop-NLS mutants, pB-mSTI1-K239A-EGFP (encoding Hop-K239A-EGFP), and grown at 37 °C. Cells were treated with leptomycin-B, fixed, stained with DAPI (blue), mounted and then visualized by confocal laser fluorescence microscopy. The labels on top of each panel refers to the filters used for visualization on the confocal microscope (FITC for EGFP, DAPI for nuclei, and MERGED). Scale bars, 10 μ m. (B). BHK-21 cells grown and transiently transfected in the same manner described above, were subjected to acute heat shock by incubation with pre-warmed media 30 min at 42 °C. The cells were then fixed and visualized as described above. Scale bars, 10 μ m. (C) BHK-21 cells were transiently transfected and treated as mentioned in A and B. Cells showing predominantly cytoplasmic fluorescence, cytoplasmic and nuclear fluorescence and predominantly nuclear fluorescence were quantified. Average values were taken from six different fields and the error bars represent standard deviations. (D) Western analysis using antibodies against β -actin (cytoplasmic and membrane marker), Hsp72/73, Hsp40, Hsp90 and Hop, after the biochemical fractionation of whole BHK-21 cell lysate obtained from cells grown at 37 °C and cells incubated for 30 min at 42 °C. Identical cell numbers were processed at 37 °C and 42 °C, and volume equivalence of each subcellular fraction was loaded proportional to the identical number of cells they were extracted from. The results shown are representative of three independent experiments.

Please cite this article as: S. Daniel, et al., Nuclear translocation of the phosphoprotein Hop (Hsp70/Hsp90 organizing protein) occurs under heat shock, and its proposed nuclear localization signal is involved in Hsp90 binding, Biochim. Biophys. Acta (2008), doi:10.1016/j.bbamer.2008.01.014

240

S. Daniel et al. / Biochimica et Biophysica Acta xx (2008) xxx-xxx



244 (12% acrylamide/bis-acrylamide w/v), and then subjected to Western ana-245 lysis, using antibodies specific for β -actin (A-2228, Sigma), Hsp72/73 (SPA-246 820, StressGen), Hsp40 (Hdj1 homologue; SPA-400, StressGen), Hsp90 α and β 247 (SPA-835, StressGen), and Hop (Antibodies by Design, MorphoSys[®]; Germany).

248 **3. Results**

249 3.1. Hop was phosphorylated in vivo

Various different lysates were prepared from NIH 3T3 cells
 treated with phosphatase inhibitors (PI) or a phosphatase (SAP).

This allowed the potential phosphorylated state of Hop in vivo to 252 be analyzed after 2D gel electrophoresis and Western analysis 253 for the detection of Hop isoforms (Fig. 1A). Previously pub- 254 lished phosphorylation studies of mouse Hop were performed 255 using cell lysates that were treated with a tyrosine phosphatase 256 inhibitor (sodium vanadate) but no serine/threonine phosphatase 257 inhibitors [4]. As preliminary work to the study presented here, 258 the 2D-electrophoretic experiments described by Lässle et al. [4] 259 were repeated but without using any phosphatase inhibitors. The 260 results were identical to those previously obtained [4] suggesting 261 that Hop was phosphorylated at sites other than tyrosine. Thus 262

ARTICLE IN PRESS

the present study made use of serine/threonine phosphatase inhibitors, in addition to the tyrosine phosphatase inhibitor.

For lysates from cells treated with PI and no SAP added. Hop 265 was found to exist not only as a major and a minor isoform, but 266 also as three more relatively acidic isoforms (Fig. 1A, upper 267panel). The major isoform with a calculated pI of approximately 2686.1 was labeled '0' and the minor isoform found at a pI of 269approximately 6.0 was labeled '+1'. A third isoform, ap-270proaching a pI of 6.0 and merging with isoform +1 was labeled 271'+2'. The two other relatively more acidic isoforms at pIs less 272than 5.9 were designated by asterisks and were not assigned 273approximate pIs. For lysates from cells treated only with SAP 274 there was a considerable reduction in isoforms +1 and +2275(Fig. 1A, lower panel), suggesting that they were depho-276sphorylated by the SAP treatment. These results provided 277 evidence of in vivo phosphorylation of at least two subpopula-278tions of Hop under normal conditions. The most acidic isoform 279obtained under normal conditions (designated by an asterisk) 280 disappeared after SAP treatment, indicating that this isoform 281 may have corresponded to a minor phosphorylated isoform of 282 Hop. A prominent isoform, designated '+3' was detected after 283 SAP treatment at an approximate pI of 5.9, and this isoform was 284 not prominent in lysate obtained from normally grown cells that 285 were treated with PI (Fig. 1A, upper panel). These results 286 directly related the occurrence of certain acidic isoforms of Hop 287to post-translational phosphorylation of the protein. 288

Mouse NIH 3T3 cells were subjected to heat shock, and 289shown to undergo induction of the stress response as evidenced 290by induction of Hsp72 protein (data not shown). The isoform 291 292composition of Hop after 30 min of heat shock (Fig. 1B, upper panel) differed from that of normal conditions in that there was a 293decrease in the amount of the major isoform 0. There was also 294 an increase of a number of minor acidic isoforms, the most 295prominent of which had an approximate pI of 5.9 and was 296 designated '+3'. This isoform was also seen in the SAP treated 297normal lysate (Fig. 1A, lower panel). The additional minor 298isoforms that appeared after heat shock were not assigned pIs, 299 and were rather indicated by asterisks. However, it was noted 300 that these isoforms were found at pIs of 5.9 or lower. Lysates 301 prepared after treatment of heat-shocked cells with SAP yielded 302 an isoform profile for Hop (Fig. 1B, lower panel) which in-303 cluded a slight increase of the previously major isoform '0', and 304 the complete absence of isoform +2. Isoform +3 showed a 305 considerable reduction after treatment with SAP. These results 306 were consistent with previous studies [4] that showed that heat 307 shock resulted in modifications of Hop. This data also provided 308 evidence for the first time that Hop was phosphorylated in vivo 309 after heat shock. 310

311 3.2. The major arm of the proposed bipartite NLS did not 312 contribute to the nuclear localization of Hop under 313 leptomycin-B treatment

We examined the subcellular distribution of Hop in baby hamster kidney cells (BHK-21), using a Hop-EGFP fusion protein, which could also be modified to determine the possible effects of mutations in the predicted NLS. The protein was transiently expressed in BHK-21 cells, fixed, and then analyzed 318 by confocal laser fluorescence microscopy. 319

Under normal conditions Hop-EGFP was found to localize 320 predominantly to the cytoplasm of most cells (Fig. 2A, I; B, I; C), 321 whereas the localization pattern of EGFP was found to be 322 distributed throughout the cell and therefore identical to pre- 323 viously published findings [data not shown; [4,22]]. Treatment 324 of transfected cells with leptomycin-B, a specific inhibitor of the 325 CRM-1 nuclear protein export pathway, showed the accumula- 326 tion of Hop-EGFP in the nucleus (Fig. 2A II; C), a finding that 327 correlates with previous literature [22].

We mutated the residues of the major arm of the proposed 329 NLS, a region in most bipartite NLSs that is critical for func- 330 tionality in vivo [28–30], and examined the subcellular loca- 331 lization patterns of the derivative proteins. Both NLS mutants of 332 Hop-EGFP also showed a predominantly cytoplasmic profile of 333 subcellular localization under normal growth conditions (Fig. 2A, 334 I; B, I). However, closer inspection of the various localization 335 patterns by quantitative analysis revealed that both of the mutant 336 Hop-EGFP derivatives showed a greater incidence of cytoplasmic 337 and nuclear localization compared to unmutated Hop-EGFP 338 (Fig. 2C). Leptomycin-B treated cells showed that both mutant 339 Hop-EGFP derivatives were 'arrested' within the nucleus, as was 340 the case for unmutated Hop-EGFP (Fig. 2A, II; C).

3.3. Hop was localized to the nucleus under heat shock and the 342 nuclear localization of Hop was not dependent on the major 343 arm of the proposed NLS 344

BHK-21 cells were treated with pre-warmed DMEM and heat 345 shocked at 42 °C for 30 min after transient transfection. In 346 response to heat shock, Hop and its putative NLS mutants showed 347 increased nuclear localization compared to its subcellular lo- 348 calization under normal conditions (Fig. 2B I and II; C). 349

Total cell extracts obtained from cells grown at 37 °C and 350 cells heat shocked at 42 °C for 30 min, were biochemically 351 fractionated into cytoplasmic, membrane and nuclear fractions. 352 Volume equivalence of each subcellular fraction was loaded 353 proportional to the identical number of cells they were ex- 354 tracted from. The proteins in each fraction were resolved by 355 SDS-PAGE and then subjected to Western blot analysis using 356 antibodies against β-actin, Hsp72/73, Hsp40, Hsp90 and Hop 357 (Fig. 2D). β -actin was found, as expected, in high amounts in 358 the cytoplasmic and membranous extracts of both normal and 359 heat-shocked extracts (Fig. 2D, Panel 1: lanes 1, 2, 3 and 4). No 360 β-actin was detected in the nuclear fraction from cells grown 361 under normal conditions (Fig. 2D, Panel 1: lane 5), suggesting 362 that this fraction was relatively free of contamination from the 363 cytoplasmic fraction. Although generally not nuclear, β -actin is 364 known to reversibly localize to the nucleus under heat shock 365 [31,32] and a faint band was thus visible in the nuclear extract 366 of heat-shocked extracts (Fig. 2D, Panel 1: lane 6). Hsp72/73 367 was found to be predominantly cytoplasmic and membranous 368 under normal and heat shock conditions (Fig. 2D, Panel 2: lanes 369 1-4). There was a slight presence of Hsp72/73 in the nuclear 370 fractions before heat shock and a definite nuclear fractionation 371 of Hsp72/73 under heat shock (Fig. 2D, Panel 2: lanes 5-6). 372



S. Daniel et al. / Biochimica et Biophysica Acta xx (2008) xxx-xxx

7



Fig. 3. Proposed NLS residues were critical for Hop–Hsp90 interaction and the phosphorylation of Hop by cdc2 kinase may perturb Hop–Hsp90 interactions. (A) Binding response curves of GST-Hop and its NLS mutants ($0.2-5 \mu$ M used, only 5 μ M trace shown for each; GST-Hop-K239A and GST-Hop-K237/238/239A) to Hsp90 (immobilized) were generated using surface plasmon resonance spectroscopy (SPR). (B) Ribbon representation of the structure of TPR2A of Hop (light grey) with the C-terminal MEEVD motif of Hsp90 shown in dark grey (1ELR; [6]). The space filled residues, K237, K238 and K239 form the major arm of the predicted NLS, and these are shown relative to the space filled residue, methionine, which forms part of the pentapeptide MEEVD of Hsp90. This figure was generated using Pymol Molecular Graphics Software [[52]; http://pymol.sourceforge.net]. (C) Binding response curves of GST-Hop and its phosphorylation mimics (GST-Hop-S189E, mimic of phosphorylation by CKII kinase; and GST-Hop-T198, mimic of phosphorylation by cdc2 kinase) to Hsp90 using SPR. GST was used as a control for non-specific binding in A and C.

Q1

The detection of Hsp72/73 in the nuclear fraction after heat shock was consistent with what has been shown previously [33], and these data together with the actin data, suggested that the nuclear fractions had been correctly isolated.

The localization of Hsp40 (Hdj1 homologue) was also ana-377 lyzed, since it is a co-chaperone of Hsp70, and like Hsp70 378 translocates from the cytoplasm to the nucleus under heat shock 379 conditions [34]. Hsp40 was detected in the cytoplasmic frac-380 tions under normal and heat-shocked conditions, as was ex-381 pected, with an increased detection in the membranous fraction 382 after heat shock (Fig. 2D, Panel 3: lanes 1-4). Although Hsp40 383 could not be detected in the nuclear fractions obtained from 384 normally grown cells, it was visibly co-fractionated with Hsp70 385 within the nuclear proteins from heat-shocked cells (Fig. 2D, 386 387 Panel 3: lanes 5-6). Hsp90 was detected predominantly in the cytoplasmic fraction, with an increased detection in the mem-388 branous fraction after heat shock (Fig. 2D, Panel 4: lanes 1-4). 389 Although previous reports have shown the nuclear translocation 390 of Hsp90 during stress conditions [5,35,36], Hsp90 was not 391 detected in the heat-shocked nuclear fractions obtained in this 392 study (Fig. 2D, Panel 4: lanes 5-6). Hop also showed pre-393 dominantly cytoplasmic fractionation under normal and heat 394 shock conditions with some detection in the membranous 395 fraction. This increased after heat shock (Fig. 2D, Panel 5: lanes 396 1-4). The detection of Hop was evident in the nuclear fraction 397 of heat-shocked lysate (Fig. 2D, Panel 5: lane 6). Taken together 398 these results demonstrated that a population of Hop localized to 399 the nucleus when the cells were subjected to heat shock, and 400 that mutations of the major arm of the proposed NLS of Hop did 401 402 not disrupt this nuclear localization.

3.4. Proposed NLS residues were critical for Hop-Hsp90
 interactions and a cdc2 kinase phosphorylation mimic of Hop
 reduced its affinity for Hsp90

SPR was used to investigate Hop-Hsp90 interactions when 406 one or more residues of the major arm of the proposed NLS of 407 Hop were mutated. Hsp90 was immobilized on the sensor chip 408 and GST-Hop constructs were passed over the chip. GST was 409used as a negative control to monitor any non-specific binding 410 of GST-Hop and its derivatives to Hsp90. Affinity constants in 411 the form of $K_{\rm D}$ values were determined for the binding of GST-412 Hop to Hsp90 and found to be 1.4 µM, consistent with pre-413 viously published data [6,15]. Both GST-Hop-K239A and GST-414 Hop-K237/238/239A showed abrogated binding to Hsp90 and 415 were comparable to the negative control GST (Fig. 3A). These 416 results suggested that residues K237, K238 and K239 were 417 critical for the interaction of Hop with Hsp90. A three dimen-418 sional (3D) representation of the structure of the TPR2A domain 419 of Hop interacting with the C-terminal MEEVD pentapeptide of 420421 Hsp90 (Fig. 3B) showed that K237, K238 and K239 potentially made contact with the methionine of the Hsp90 MEEVD motif. 422We also investigated the effect of phosphorylation at sites 423 S189 and T198 of Hop on its interactions with Hsp90. Phos-424phorylation was mimicked at both these sites by substituting the 425respective serine and threonine residues with an acidic residue, 426 427 glutamic acid. SPR analysis was conducted to determine the

extent of interaction of both these Hop phosphorylation mimics 428 with Hsp90. The binding curves obtained by plotting response 429 units versus time (seconds) showed that the CKII kinase phos- 430 phorylation mimic of Hop (GST-Hop-S189E) did not show 431 much deviation from the unmutated GST-Hop (Fig. 3C) where- 432 as the cdc2 kinase phosphorylation mimic of Hop (GST-Hop- 433 T198E) exhibit diminished affinity for Hsp90. K_D values of 434 2.5 μ M and 1 μ M were obtained for GST-Hop-T198E–Hsp90 435 and GST-Hop-S189E–Hsp90 interactions respectively. These 436 values may or may not be of biological significance in vivo. 437

All SPR analyses were repeated independently by immobi- 438 lizing the GST-Hop constructs on a sensor chip and passing 439 over purified Hsp90. Similar results were obtained (data not 440 shown). Taken together, the results from these experiments 441 showed that the major arm of the proposed NLS was crucial for 442 Hop–Hsp90 interactions, and that mimicking phosphorylation 443 by cdc2 kinase reduced the affinity of Hop for Hsp90. 444

4. Discussion

This is the first report to have provided direct evidence of Hop 446 being a phosphoprotein under normal and stress conditions, and 447 that the subcellular localization of Hop was primarily stress 448 regulated. We have established that unlike previously reported 449 bipartite NLSs, the major arm of the proposed NLS of Hop was 450 not critical for its nuclear translocation. However these residues 451 were crucial for Hop–Hsp90 interaction, and through this in- 452 teraction, we hypothesize that this region could possibly serve to 453 retain Hop within the cytoplasm. Furthermore, phosphorylation- 454 mimic studies suggested that phosphorylation of Hop at a cdc2 455 kinase site may diminish the affinity of Hop for Hsp90, whilst 456 phosphorylation of Hop at a CKII site may not interfere with 457 Hop–Hsp90 interactions [22].

The presence of okadeic acid (serine/threonine phosphatase 459 inhibitor) in the phosphorylation study conducted, suggested 460 that phosphorylation of Hop occurred predominantly at serine or 461 threonine residues. This is further substantiated by evidence of in 462 vitro phosphorylation of Hop at S189 and T198 [22,23]. Heat 463 stress may exert inhibitory and activating effects on various 464 kinases, including Mitogen-activated protein (MAP) kinases, 465 and phosphatases [37,38,39,40]. Hop was found to have a po- 466 tential phosphorylation site for MAPKAP kinase 2 and two 467 potential sites for S6 kinases within its amino acid sequence [4]. 468 Hop may thus be phosphorylated by a stress kinase pathway in 469 vivo [4] in response to heat shock. The differentially phos- 470 phorylated isoforms of Hop may correspond to subpopulations 471 of Hop, which are localized to varying subcellular compartments 472 during normal and/or stress conditions. Since Hop is known to 473 be predominantly cytoplasmic under normal conditions in this 474 study and in others [4,22], the major '0' isoform of Hop resolved 475 by 2D analysis, may correspond to the cytoplasmic population of 476 Hop under normal conditions. 477

Acute heat shock resulted in increased incidence of cells 478 showing greater translocation of Hop-EGFP to the nucleus (but 479 not nucleolus). This was supported by the detection of endo- 480 genous Hop in the nuclear protein fractions after heat shock. 481 Short-term heat shock in mammalian cells is known to cause the 482

Please cite this article as: S. Daniel, et al., Nuclear translocation of the phosphoprotein Hop (Hsp70/Hsp90 organizing protein) occurs under heat shock, and its proposed nuclear localization signal is involved in Hsp90 binding, Biochim. Biophys. Acta (2008), doi:10.1016/j.bbamcr.2008.01.014

445

S. Daniel et al. / Biochimica et Biophysica Acta xx (2008) xxx-xxx

ARTICLE IN PRESS

transient arrest of the cell cycle at the G1/S stage [41]. The arrest of mouse fibroblast cells in the G1/S phase showed increased nuclear localization of Hop [22]. Since the heat shock conditions used here also increased the localization of Hop to the nucleus, these conditions probably promoted G1/S arrest of the cells. The nuclear localization of Hop may thus be primarily stress regulated and secondarily regulated by cell cycle events.

Although mutations within the major arm of the putative 490 NLS of Hop did not disrupt the nuclear translocation of the 491 Hop-EGFP construct, SPR analysis showed that these muta-492 tions resulted in minimal interaction of GST-Hop with Hsp90. A 493three-dimensional (3D) representation of the Hop TPR2A do-494 main showing the NLS residues that were mutated, together 495 with the C-terminal pentapeptide of Hsp90 (MEEVD motif), 496 497 illustrated the proximity of these NLS residues to the methionine of the Hsp90 peptide (Fig. 3B). One or more of these 498 residues could be involved in direct interactions with Hsp90. 499

We propose that the minor arm of this putative NLS could be the functional NLS, acting together with a second putative monopartite NLS found within amino acids 337–351 (Fig. 4). Mutation of the major arm of the putative bipartite NLS may have resulted in a modification of the structure of Hop such that the putative monopartite NLS was better positioned and exposed to importin binding, thereby facilitating more efficient nuclear localization than that obtained for unmutated Hop. A 507 second possibility is that the major arm of the putative bipartite 508 NLS of Hop might promote the cytoplasmic retention of Hop by 509 binding to Hsp90, thus explaining why mutation of residues 510 within Hop that rendered it incapable of interacting with Hsp90 511 in vitro, allowed for greater nuclear translocation of Hop in 512 vivo. Hsp90 has previously been reported to sequester the glu- 513 cocorticoid receptor to the cytoplasm through their interaction 514 within a complex [42].

A subpopulation of endogenous Hop was detected in the 516 membranous fractions under normal and heat shock conditions. 517 This finding was consistent with reports of the interaction of 518 Hop with Prp^c at the plasma membrane [20]. Interestingly, it 519 was also found that Prp^c interacts with Hop at residues 230–245 520 [43], which overlaps with the Hsp90 binding site [[15]; and this 521 study]. Prp^c binding to Hop at this site may act as a mechanism 522 to improve the efficiency of retaining a subpopulation of Hop at 523 the plasma membrane. This further supports the proposed role 524 of Hop-Hsp90 interactions in modulating the subcellular lo- 525 calization of Hop. Our data also showed the co-fractionation of 526 Hsp70, Hsp90 and Hsp40, along with Hop, in the membranes 527 under normal and heat shock conditions. This also suggests that 528 Hop may associate with one or more of its chaperone partners 529 within various membranous structures. 530



Fig. 4. Positioning of the proposed bipartite NLS with respect to TPR2A domain and a second potential NLS. Ribbon representation of the structure of TPR2A of Hop and a flanking C-terminal helix interacting with the C-terminal MEEVD motif of Hsp90 (1ELR; [6]). The residues, K229, N233 and K301 are those that are known to be critical for interaction with Hsp90. Residues denoted with a single asterisk correspond to those that form the major arm of the bipartite NLS (K237, K238 and K239). The residues denoted with a double asterisk, K222 and K223 (dark grey), form part of the minor arm of this NLS, which may be responsible for functionality of the NLS within the protein. A second NLS has been identified within the flanking C-terminal portion of Hop (337–351) shown in dark grey, which may be working in concert with the first predicted NLS. This figure was generated using Pymol Molecular Graphics Software [49]; http://pymol.sourceforge.net.

ARTICLE IN PRESS

Based on the results obtained from this study, together with 531what is currently known about Hop particularly regarding its 532nucleo-cytoplasmic shuttling properties, we propose a model 533detailed in Fig. 5. Within the normal cytoplasmic environment, 534Hop equilibrates between an Hsp90-complexed state and an 535 uncomplexed free state [44,45], and the equilibrium between 536 the two states may be regulated through phosphorylation of Hop 537(Fig. 5, arrow A). Phosphorylation of Hop by cdc2 kinase may 538disrupt Hop-Hsp90 interactions as indicated by the SPR ana-539lysis. This frees Hop, but retains it within the cytoplasm [[22]; 540Fig. 5, arrow C/cdc2?] so that it can interact with other proteins 541in complexes that are independent of Hsp90 [46,47]. 542

543 Under normal conditions, Hop–Hsp90 interactions within the 544 cytoplasm may serve to anchor Hop to this subcellular com-545 partment. The major arm of the proposed NLS of Hop may contribute to the overall localization of Hop by its interaction 546 with Hsp90. This facilitates the assembly of the Hsp70–Hsp90 547 chaperone complex, which has been shown to be cytoplasmic 548 (Fig. 5, arrow B). It also implies that Hop translocates into the 549 nucleus when it is not complexed with Hsp90, possibly pro- 550 moted by CKII phosphorylation of Hop (Fig. 5, arrow C/CKII?). 551 Previous reports have shown the nuclear translocation of Hsp90 552 during stress conditions including heat shock [5,35,36]. In the 553 present study Hsp90 was not detected in the nuclear fractions 554 under normal or heat shock conditions. Biochemical fractiona- 555 tion techniques are known to be disruptive and may have altered 556 the subcellular localization pattern of a protein such as Hsp90, 557 whose translocation into the nucleus may be transient or dyn- 558 amic in nature. This may also explain why relatively low levels 559 of Hop were detected in the nuclear fractions, when compared to 560



Fig. 5. A model reflecting the possible links between the subcellular localization of the phosphoprotein Hop and its functionality as a Hsp70/Hsp90 scaffolding protein. Hop exists on its own or in complex with Hsp90, in the cytoplasm under normal conditions. This may be regulated by phosphorylation, with cdc2 kinase phosphorylation of Hop disrupting its interaction with Hsp90 (cdc2? arrow; A). Interaction of Hop with Hsp90 is known to facilitate a number of other interactions, of which the most well established one is the interaction of the Hop–Hsp90 complex with Hsp70 (associated with its co-chaperone Hsp40 and substrate) in order to facilitate substrate transfer from Hsp70 to Hsp90 (B). This multichaperone complex then dissociates, freeing its various components. Hop is known to translocate to the nucleus (C) under stressful conditions, and its localization may be regulated by phosphorylation, with CKII phosphorylation possibly promoting nuclear localization (CKII? arrow) and cdc2 kinase phosphorylation possibly promoting cytoplasmic retention (cdc2? arrow). It is speculated that Hop may also be capable of moving into the nucleus in concert with Hsp90 (D) as a complex (arrows shown in dotted lines) by either the putative NLS (222–239), or through the functioning of multiple NLSs, and possibly also promoted by CKII phosphorylation (CKII? dotted arrow). It is already known that both Hsp70 (together with Hsp40) and Hsp90 translocate into the nucleus under heat shock (E and F respectively). Within the nucleus Hop may have a number of functions, including its basic function of interacting with Hsp70 and/or Hsp90 to form nuclear complexes (G). The 1, 2A, 2B, C and NLS annotations on Hop refer to its TPR1, TPR2A and TPR2B domains, C-terminal domain, and nuclear localization signal sequence, respectively. Hsp40, Hsp70 and Hsp90 are labeled as 40, 70 and 90, respectively.

the confocal microscopic data. Thus, if Hsp90 does indeed translocate into the nucleus during heat shock, the absence of functional NLSs within Hsp90 [36,48] suggests that Hsp90 may have to "piggyback" on an NLS-bearing protein. CKII phosphorylation was shown to promote nuclear localization of Hop [22] and now, evidence has been presented to suggest that CKII phosphorylation does not disrupt Hop–Hsp90 binding. Thus we cannot exclude the possibility of a subpopulation of Hop–Hsp90 translocating as a complex into the nucleus under specific regulatory conditions, such as phosphorylation by CKII (Fig. 5, dotted arrow D/CKII?). As other studies [34] and ours have shown the presence of Hsp70 and Hsp40 in heat-shocked nuclear fractions. This is suggestive of the nuclear assembly of the Hsp70/Hsp90 chaperone heterocomplex during heat stress (Fig. 5 arrows E, F and G).

The details of the co-localization of Hop and Hsp90, as well 576as the scaffolding properties of Hop within the nucleus, under 577 conditions of heat shock remains to be investigated. Changes in 578the ionic state and conformation of Hop due to phosphorylation 579at S189 and T198, may have an effect on its interactions with 580other members of the chaperone heterocomplex, such as Hsp70, 581 and this also needs to be investigated. The mechanism of nu-582clear localization of the Hsp70-Hop-Hsp90 chaperone hetero-583complex under both normal and stress conditions is complex. 584 However, the putative phosphorylation-regulated NLS of Hop 585 appears to play an important role in the functionality of the 586 chaperone heterocomplex, not only on the level of its subcel-587 lular localization but also with regards to the assembly of the 588 chaperone heterocomplex. 589

590 Acknowledgements

561

562

563

564

565

566

567

568

569

570

571

572

573

574

575

This work was funded in part by a National Research Foun-591dation (NRF) Unlocking the Future Focus Area grant (South 592Africa; GUN No: 2053542) and a Wellcome Trust grant (UK; 593Grant No: 066705) awarded to GLB, and a NRF-Hungary 594Collaborative grant (South Africa; GUN No: 2067467) awarded 595to GLB and PC. SD was awarded an Andrew Mellon Scho-596 larship (Rhodes University, South Africa) during her doctoral 597studies, and a JRC Adhoc Postdoctoral Bursary (Rhodes Uni-598 599versity, South Africa) to facilitate the writing of this paper. CS is a Bolyai Research Scholar of the Hungarian Academy of Sci-600 ences. The authors thank Bronwyn McLean (Graphics Services 601 Unit, Rhodes University, South Africa) for her expert assistance 602 with the figures presented. We also thank the staff of the Elec-603 604 tron Microscope Unit at University of Kwazulu-Natal, Pietermaritzburg (South Africa) for their technical assistance. 605

606 **References**

- [1] C.M. Nicolet, E.A. Craig, Isolation and characterization of STI1, a stressinducible gene from *Saccharomyces cerevisiae*, Mol. Cell. Biol. 9 (1989)
 3638–3646.
- [2] R.J. Schumacher, R. Hurstll, W.P. Sullivan, N.J. McMahons, D.O. Tofts,
 R.L. Matts, ATP-dependent chaperoning activity of reticulocyte lysate,
 J. Biol. Chem. 269 (1994) 9493–9499.
- [3] B. Honoré, H. Leffers, P. Madsen, H.H. Rasmussen, J. Vandekerckhove, J.E.
 Celis, Molecular cloning and expression of a transformation-sensitive human

protein containing the TRP motif and sharing identity to the stress-inducible 615 yeast protein STI1, J. Biol. Chem. 267 (1992) 8485–8491. 616

- M. Lässle, G.L. Blatch, V. Kundra, T. Takatori, B.R. Zetter, Stress-inducible, 617 murine protein mSTI1. Characterization of binding domains for heat shock 618 proteins and in vitro phosphorylation by different kinases, J. Biol. Chem. 272 619 (1997) 1876–1884. 620
- [5] W.B. Pratt, D.O. Toft, Steroid receptor interactions with heat shock protein 621 and immunophilin chaperones, Endocr. Rev. 18 (1997) 306–360. 622
- [6] C. Scheufler, A. Brinker, G. Bourenkov, S. Pegoraro, L. Moroder, H. 623 Bartunik, F.U. Hartl, I. Moarefi, Structure of TPR domain–peptide com- 624 plexes: critical elements in the assembly of the Hsp70–Hsp90 multi- 625 chaperone machine, Cell 101 (2000) 199–210. 626
- [7] G.L. Blatch, M. Lässle, B.R. Zetter, V. Kundra, Isolation of a mouse cDNA 627 encoding mSTI1, a stress-inducible protein containing the TPR motif, 628 Gene 194 (1997) 277–282.
- [8] J. Demand, J. Luders, J. Hohfeld, The earboxy-terminal domain of Hsc70 630 provides binding sites for a distinct set of chaperone cofactors, Mol. Cell. 631 Biol. 18 (1998) 2023–2028.
 632
- M.D. Adams, S.E. Celniker, R.A. Holt, C.A. Evans, J.D. Gocayne, P.G. 633 Amanatides, S.E. Scherer, P.W. Li, R.A. Hoskins, R.F. Galle, et al., The 634 genome sequence of Drosophila melanogaster, Science 287 (2000) 635 2185–2195.
- Z. Zhang, M.K. Quick, K.C. Kanelakis, M. Gijzen, P. Krishna, Char- 637 acterization of a plant homolog of Hop, a cochaperone of Hsp90, Plant 638 Physiol. 131 (2003) 525–535.
- J.R. Webb, A. Campos-Neto, Y.A.W. Skeiky, S.G. Reed, Molecular char- 640 acterization of the heat-inducible LmSTI1 protein of Leishmania major, 641 Mol. Biochem. Parasitol. 89 (1997) 179–193. 642
- [12] C.H. Cheng, S.M. Liu, T.Y. Chow, Y.Y. Hsiao, D.P. Wang, J.J. Huang, H. 643
 H. Chen, Analysis of the complete genome sequence of the Hz-1 virus 644
 suggests that it is related to members of the Baculoviridae, J. Virol. 76 645
 (2002) 9024–9034.
- [13] J. van der Spuy, B.D. Kana, H.W. Dirr, G.L. Blatch, Heat shock cognate 647 protein 70 chaperone-binding site in the co-chaperone murine stress- 648 inducible protein 1 maps to within three consecutive tetratricopeptide re- 649 peat motifs, Biochem. J. 345 (2000) 645–651. 650
- [14] S. Chen, V. Prapapanich, R.A. Rimerman, B. Honore, D.F. Smith, In- 651 teractions of p60, a mediator of progesterone receptor assembly, with heat 652 shock proteins Hsp90 and Hsp70, Mol. Endocrinol. 10 (1996) 682–693. 653
- [15] O.O. Odunuga, J.A. Hornby, C. Bies, R. Zimmermann, D.J. Pugh, G.L. 654 Blatch, Tetratricopeptide repeat motif-mediated Hsc70–mSTI1 interaction. 655 Molecular characterization of the critical contacts for successful binding 656 and specificity, J. Biol. Chem. 278 (2003) 6896–6904. 657
- [16] A. Brinker, C. Scheufler, F. von der Mulbe, B. Fleckenstein, C. Herrmann, 658
 G. Jung, I. Moarefi, F.U. Hartl, Ligand discrimination by TPR domains. 659
 Relevance and selectivity of EEVD-recognition in hsp70.hop.hsp90 660
 complexes, J. Biol. Chem. 277 (2002) 19265–19275. 661
- [17] G. Flom, R.H. Behal, L. Rosen, D.G. Cole, J.L. Johnson, Definition of the 662 minimal fragments of Sti1 required for dimerization, interaction with 663 Hsp70 and Hsp90 and in vivo functions, Biochem. J. 404 (2007) 158–167. 664
- [18] G. Flom, J. Weekes, J.J. Williams, J.L. Johnson, Effect of mutation of the 665 tetratricopeptide repeat and asparatate-proline 2 domains of Sti1 on Hsp90 666 signaling and interaction in Saccharomyces cerevisiae, Genetics 172 667 (2006) 41–51. 668
- [19] P.E. Carrigan, D.L. Riggs, M. Chinkers, D.F. Smith, Functional com- 669 parison of human and drosophila Hop reveals novel role in steroid receptor 670 maturation, J. Biol. Chem. 280 (2005) 8906–8911. 671
- [20] S.M. Zanata, M.H. Lopes, A.F. Mercadante, G.N.M. Hajj, L.B. Chiarini, 672 R. Nomizo, A.R.O. Freitas, A.L.B. Cabral, K.S. Lee, M.A. Juliano, E. de 673 Oliveira, S.G. Jachieri, R. Linden, R.R. Brentani, V.R. Martins, Stress- 674 inducible protein 1 is a cell surface ligand for cellular prion that triggers 675 neuroprotection, EMBO J. 21 (2002) 3307–3316. 676
- [21] A. Muesch, E. Hartmann, K. Rohde, R. Rubartelli, R. Sitia, T.A. Rapoport, 677 A novel pathway for secretory proteins? Trends. Biochem. Sci. 15 (1990) 678 86–88.
- [22] V.M. Longshaw, J.P. Chapple, M.S. Balda, M.E. Cheetham, G.L. Blatch, 680 Nuclear translocation of the Hsp70/Hsp90 organizing protein mSTI1 is 681 regulated by cell cycle kinases, J. Cell. Sci. 117 (2004) 701–710.

Please cite this article as: S. Daniel, et al., Nuclear translocation of the phosphoprotein Hop (Hsp70/Hsp90 organizing protein) occurs under heat shock, and its proposed nuclear localization signal is involved in Hsp90 binding, Biochim. Biophys. Acta (2008), doi:10.1016/j.bbamcr.2008.01.014

ARTICLE IN PRESS

S. Daniel et al. / Biochimica et Biophysica Acta xx (2008) xxx-xxx

ARTICLE IN PRESS

- [23] V.M. Longshaw, H.W. Dirr, G.L. Blatch, M. Lässle, The in vitro phosphorylation of the co-chaperone mSTI1 by cell cycle kinases substantiates a predicted casein kinase II-p34cdc2-NLS (CcN) motif, Biol. Chem. 381 (2000) 1133–1138.
- [24] J. Sambrook, D.W. Russell, Molecular Cloning: a Laboratory Manual, 3
 ed., Cold Spring Harbour Laboratory Press, New York, 2001.
- [25] U.K. Laemmli, Cleavage of structural proteins during the assembly of the
 head of bacteriophage T4, Nature 227 (1970) 680–685.
- [26] H. Towbin, T. Staehelin, J. Gordon, Electrophoretic transfer of proteins
 from polyacrylamide gels to nitrocellulose sheets: procedure and some
 applications, Proc. Natl. Acad. Sci. U. S. A. 76 (1979) 4350–4354.
- [27] H. Karasuyama, F. Melchers, Establishment of mouse cells which constitutively secrete large quantities of interleukin 2, 3, 4, or 5, using modified cDNA expression vectors, Eur. J. Immunol. 18 (1988) 97–104.
- E. Conti, J. Kuriyan, Crystallographic analysis of the specific yet versatile
 recognition of distinct nuclear localization signals by karyopherin alpha,
 Structure 8 (2000) 329–338.
- [29] M.R. Fontes, T. The, B. Kobe, Structural basis of recognition of monopartite and bipartite nuclear localization sequences by mammalian importin-alpha, J. Mol. Biol. 297 (2000) 1183–1194.
- [30] B. Friedrich, C. Quensel, T. Sommer, E. Hartmann, M. Köhler, Nuclear
 localization signal and protein context mediate importin a specificity of
 nuclear import substrates, Mol. Cell. Biol. 26 (2006) 8697–8709.
- [31] K. Iida, S. Matsumoto, I. Yahara, The KKRKK Sequence is involved in heat shock-induced nuclear translocation of the 18-kDa actin-binding protein, Cofilin, Cell Struct. Funct. 17 (1992) 39–46.
- [32] A. Wada, M. Fukuda, M. Mishima, E. Nishida, Nuclear export of actin: a
 novel mechanism regulating the subcellular localization of a major cyto skeletal protein, EMBO J. 17 (1998) 1635–1641.
- [33] J.M. Velazquez, S. Lindquist, Hsp70: nuclear concentration during envi ronmental stress and cytoplasmic storage during recovery, Cell 36 (1984)
 655–662.
- [34] H. Hattori, Y-C. Liu, I. Tohnai, M. Ueda, T. Kaneda, T. Kobayashi, K.
 Tanabe, K. Ohtsuka, Intracellular localization and partial amino acid sequence of a stress-inducible 40-kDa protein in HeLa cells, Cell Struct.
 Funct. 17 (1992) 77–86.
- [35] M. Biggiogera, R.M. Tanguay, R. Marin, Y. Wu, T.E. Martin, S. Fakan,
 Localization of heat shock proteins in mouse male germ cells: an immu noelectron microscopical study, Exp. Cell Res. 229 (1996) 77–85.
- [36] T. Langer, S. Rosmus, H. Fasold, Intracellular localization of the 90 kDA
 heat shock protein (HSP90alpha) determined by expression of a EGFP-
- 765

HSP90alpha-fusion protein in unstressed and heat stressed 3T3 cells, Cell 724 Biol. Int. 27 (2003) 47–52. 725

- [37] G. Bagi, E.J. Hidvegi, Protein phosphorylation and kinase activities in 726 tumour cells after hyperthermia, Int. J. Radiat. Biol. 58 (1990) 633–650. 727
- [38] G.N. Somero, Proteins and temperature, Annu. Rev. Physiol. 57 (1995) 728 43–68. 729
- [39] J. Rouse, P. Cohen, S. Trigon, M. Morange, A. Alonso-Llamazares, A 730 novel kinase cascade triggered by chemical stress and heat shock which 731 stimulates MAP kinase-activated protein kinase-2 and phosphorylation of 732 the small heat shock proteins, Cell 78 (1994) 1027–1037. 733
- [40] F. Chen, M. Torres, R.F. Duncan, Activation of mitogen-activated protein 734 kinase by heat shock treatment in *Drosophila*, Biochem. J. 312 (1995) 735 341–349.
- [41] E.W. Khandijan, Heat treatment induces dephosphorylation of pRb and 737 dissociation of T-antigen pRb complex during transforming infection with 738 SV40, Oncogene 10 (1995) 359–367.
 [42] A. Guiochon-Mantel, K. Delabre, P. Lescop, E. Milgrom, Intracellular 740
- [42] A. Guiochon-Mantel, K. Delabre, P. Lescop, E. Milgrom, Intracellular 740 traffic of steroid hormone receptors, J. Steroid Biochem. Mol. Biol. 56 741 (1996) 3–9.
- [43] A.S. Coitinho, M.H. Lopes, G.N.M. Hajj, J.I. Rossato, A.R. Freitas, C.C. 743 Castro, M. Cammarota, R.R. Brentani, I. Izquierdo, V.R. Martins, Short- 744 term memory formation and long-term memory consolidation are enhanced 745 by cellular prion association to a stress-inducible protein 1, Neurobiol. Dis. 746 26 (2007) 282–290. 747
- [44] H.C.J. Chang, D.F. Nathan, S. Lindquist, In vivo analysis of the hsp90 748 cochaperone stil (p60), Mol. Cell. Biol. 17 (1997) 318–325. 749
- [45] M.P. Hernandez, W.P. Sullivan, D.O. Toft, The assembly and intermo- 750 lecular properties of the hsp70–Hop–hsp90 molecular chaperone complex, 751
 J. Biol. Chem. 277 (2002) 38294–38304. 752
- [46] T. Abbas-Terki, O. Donze, P.A. Briand, D. Picard, Hsp104 interacts with Hsp90 753 cochaperones in respiring yeast, Mol. Cell. Biol. 21 (2001) 7569–7575.
- [47] M. Gebauer, R. Melki, U. Gehring, The chaperone cofactor hop/p60 in- 755 teracts with the cytosolic chaperonin-containing TCP-1 and affects its nu- 756 cleotide exchange and protein folding activities, J. Biol. Chem. 273 (1998) 757 29475–29480.
- [48] K.I.L. Kang, J. Devin, F. Cadepond, N. Jibard, A. Guiochon-Mantel, E.-M. 759 Baulieu, M.G. Catelli, In vivo functional protein–protein interaction: 760 nuclear targeted Hsp90 shifts cytoplasmic steroid receptor mutants into the 761 nucleus, Proc. Natl. Acad. Sci. U. S. A. 91 (1994) 340–344. 762
- [49] W.L. DeLano, S. Bromberg, Pymol Users Guide, DeLano Scientific LLC, 763 California, 2004. 764