The Hsp70-Hsp90 co-chaperone Hop/Stip1 shifts the proteostatic balance from folding towards degradation

Kaushik Bhattacharya¹, Lorenz Weidenauer², Tania Morán Luengo³,⁴, Ellis C. Pieters³,⁴, Pablo C. Echeverría¹,⁶, Lilia Bernasconi¹, Diana Wider¹, Yashar Sadian⁵, Margreet B. Koopman³,⁴, Matthieu Villemin¹, Christoph Bauer⁵, Stefan G. D. Rüdiger³,⁴, Manfredo Quadroni² & Didier Picard¹

Hop/Stip1/Sti1 is thought to be essential as a co-chaperone to facilitate substrate transfer between the Hsp70 and Hsp90 molecular chaperones. Despite this proposed key function for protein folding and maturation, it is not essential in a number of eukaryotes and bacteria lack an ortholog. We set out to identify and to characterize its eukaryote-specific function. Human cell lines and the budding yeast with deletions of the Hop/Sti1 gene display reduced proteasome activity due to inefficient capping of the core particle with regulatory particles. Unexpectedly, knock-out cells are more proficient at preventing protein aggregation and at promoting protein refolding. Without the restraint by Hop, a more efficient folding activity of the prokaryote-like Hsp70-Hsp90 complex, which can also be demonstrated in vitro, compensates for the proteasomal defect and ensures the proteostatic equilibrium. Thus, cells may act on the level and/or activity of Hop to shift the proteostatic balance between folding and degradation.
Homoeostasis of the proteome (proteostasis) is essential for health and longevity. Proteotoxic stresses lead to protein misfolding and aggregation, which trigger protein quality control mechanisms. The major ones, all assisted by molecular chaperones, are protein refolding and the degradation of misfolded and aggregated proteins by the proteasome and by autophagy. Defects in any of these mechanisms can cause severe proteotoxicity, which in turn can contribute to neurodegenerative disorders such as Huntington’s, Parkinson’s, and Alzheimer’s, premature aging, and other diseases.

In eukaryotes, the Hsp70 and Hsp90 molecular chaperone machines are major contributors to proteostasis by providing a platform for folding of both nascent polypeptides and misfolded, structurally labile and mutated proteins, collectively called “clients.” For folding and assembly of clients, both Hsp70 and Hsp90 undergo large conformational changes and collaborate with co-chaperones. One of these co-chaperones is the Hop, an organizing protein (Hoop), encoded by the gene STIP1 in mammals. It is an adaptor molecule between Hsp70 and Hsp90, which facilitates the folding, stabilization, or assembly of clients by promoting their transfer to Hsp90 after the initial recognition and binding of clients by Hsp70 in collaboration with its J-domain containing co-chaperone Hsp40. Hop forms a ternary complex with Hsp70 and Hsp90 using its tetraicropetide repeat (TPR) domains. Two of its three TPRs, TPR1 and TPR2A, specifically bind the extreme C-terminal sequences EEEV and MEEVD of Hsp70 and Hsp90, respectively. While these are the primary interaction surfaces, additional contacts serve to stabilize the complexes and to facilitate dynamic rearrangements.

Proteins, whose folding or refolding fails, are degraded by the proteasome, a highly conserved and regulated eukaryotic protease complex. It is a 1.6 to 2.5 MDa complex consisting of a 20S proteolytic core particle (CP) and a 19S regulatory particle (RP); the CP can be capped by one or two RPs resulting in 26S or 30S particles, respectively. The RP is divided into a lid and a base and has unique regulatory functions; it recognizes ubiquitinated substrates produced by the E1-E2-E3 ubiquitination system, promotes their deubiquitination and unfolding, the subsequent gate-opening of the CP, and finally the loading of the processed substrates into the proteolytic chamber. Dedicated chaperones for the assembly of CP and the RP base are well known, whereas RP lid assembly is still not well understood. Hsp90 has been proposed to be an assembly chaperone for the RP lid complex based on genetic interactions in the budding yeast and the reconstitution of the RP lid complex in E. coli co-expressing yeast Hsp90.

Prokaryotes and eukaryotic organelles do have Hsp70 and Hsp90 orthologs but lack a Hop-like protein; their Hsp70 and Hsp90 physically and functionally interact directly. In eukaryotes, Hop is not absolutely indispensable as mutant budding yeast, worms, and flies are viable with only mild phenotypes. In contrast, the deletion of STIP1 is lethal early in embryonic development in the mouse, possibly indicating that the function of Hop might be cell type-specific or dependent on specific cellular states or requirements.

In this study, we have explored Hop in prokaryotes, what its critical functions are, and whether and how the eukaryotic Hsp70-Hsp90 molecular chaperone machines may function without Hop to ensure proteostasis. Our studies on the functions of Hop as a co-chaperone of the Hsp70-Hsp90 molecular chaperone machines led us to the discovery of alternative cellular strategies that ensure proper protein folding and proteostasis in human and yeast cells lacking this co-chaperone. These findings highlight the persistence of evolutionarily more ancient mechanisms in eukaryotic cells that may contribute to balance protein folding and degradation under certain conditions.

### Results
Human Hop knock-out cells maintain cellular fitness and proteostasis and are not hypersensitive to proteotoxic stress.

To study the functions of Hop in eukaryotic cells, we knocked out its gene STIP1 in several human cell lines with the CRISPR/Cas9 technique. Quantitation of the STIP1 mRNA of the knock-out (KO) clones by Q-PCR showed a drastic reduction (Supplementary Fig. 1a), and the absence of full-length Hop protein was confirmed by immunoblotting using a specific antibody to Hop (Fig. 1a). We did notice that the HEK293T clone KO1 expresses a residual low level of a truncated form of Hop, which we characterized by mass spectrometry (MS) (Supplementary Data 1); it only retains the Hsp90-binding domain TPR2A. In subsequent experiments, KO1 proved to behave essentially like the other HEK293T clone (KO33), which is devoid of any detectable trace of Hop. Morphological examination revealed no obvious differences between wild-type (WT) and KO cells (Supplementary Fig. 1b). Growth rates of KO cells are only moderately reduced (Supplementary Fig. 1c); this observation was supported by cell cycle analyses in that KO cells show less cyclic phase cells (S + G2/M) and more cells in the G0/G1 resting phase (Supplementary Fig. 1d). The flow cytometric analyses of cells stained with annexin V and propidium iodide (PI) showed that none of the KO cell lines have any obvious viability issues (Supplementary Fig. 1e).

A label-free MS quantitation of the whole-cell proteome indicated that the vast majority of the proteome is not affected by the KO (Fig. 1b and Supplementary Data 2). With our statistical cutoffs (see “Methods”), only a minor proportion of the identified proteins of HEK293T and HCT116 cells are altered by the KO. Gene ontology (GO) term analyses of the statistically significant changes reveal a strong enrichment of stress response processes and protein folding/refolding mechanisms in Hop KO cells (Fig. 1c). This prompted us to determine how the KO cells survive in various stress conditions. We exposed them to thapsigargin, DTT, and A23187, which all induce the unfolded protein response of the endoplasmic reticulum, and to the oxidative stress inducer H2O2. Remarkably, KO cells are clearly not hypersensitive to these stress agents since these lead to comparable numbers of dead cells (Fig. 1d) and to a rather reduced impact on cell morphology compared to WT cells (Supplementary Fig. 1f). Upon exposure to heat shock (HS), a stress which induces protein misfolding and aggregation, KO cells showed a similar sensitivity in the HEK293T background (Supplementary Fig. 1g), but a markedly reduced sensitivity in the HCT116 and A549 backgrounds (Fig. 1e). We also challenged both WT and KO cells with azetidine-2-carboxylic acid (AZC); the incorporation of this proline analog into nascent proteins causes misfolding and subsequent protein aggregation. KO cells are more resistant to this proteotoxic stress, based on the reduced number of apoptotic cells (Fig. 1f) and morphological examination (Supplementary Fig. 1h). Thus, the absence of Hop does not compromise the overall proteostatic equilibrium; instead, if anything, proteostatic buffering of KO cells becomes more robust and resilient to proteotoxic stress conditions.

The proteasome and the Hsp70-Hsp90 molecular chaperone axis are differentially required in KO cells. Cells cope with proteotoxic stresses by using two important protein quality control mechanisms: degradation of misfolded proteins by the proteasome and in some circumstances by autophagy, and refolding and stabilization of misfolded proteins by molecular
chaperones, including the Hsp70-Hsp90 molecular chaperone machines (Fig. 2a). To assess how these mechanisms are operating in KO cells, we pharmacologically blocked Hsp90, Hsp70, the proteasome, and ubiquitination. We observed that KO cells are hypersensitive to Hsp90 inhibition with geldanamycin (GA), as demonstrated by enhanced cell cycle arrest in the G2/M phase and accumulation of apoptotic cells (Fig. 2b and Supplementary Fig. 2d). Remarkably, these chemically different Hsp90 inhibitor PU-H71 (Supplementary Fig. 2b). KO cells also showed a higher sensitivity to the effects of GA on the accumulation of dead cells (Supplementary Fig. 2a). This observation was further confirmed by monitoring the effects of GA on the accumulation of dead cells (Supplementary Fig. 2b). KO cells also showed a higher sensitivity to the chemically different Hsp90 inhibitor PU-H71 (Supplementary Fig. 2c) and the C-terminal Hsp90 inhibitor novobiocin (Supplementary Fig. 2d). Remarkably, these findings are reminiscent of the observation with yeast that Hsp90 inhibition and Δstl1 are...
molecular chaperone machines are either enriched or only actors of Hop itself. The components of the Hsp70-Hsp90 be interactors of Hsp70 and/or Hsp90 rather than direct interaction on the ability of Hop to bind Hsp70 and Hsp90 and may mostly Thus, proteins of the Hop interactome are largely dependent enriched with WT Hop (Fig. 3a and Supplementary Data 3). mutant, ~41% were identified by mass spectrometry (IP-MS) analysis with transfected HEK293T KO cells. After an initial quality control by sodium dodecyl sulfate chromatography (Fig. 3b), samples were subjected to label-free liquid polyacrylamide gel electrophoresis (SDS-PAGE) (Supplementary mass spectrometry (LC/MS-MS) analysis. By synthetically lethal, indicating that this is an evolutionarily conserved characteristics. Similarly, the Hsp70 inhibitor JG-98 blocks the growth of KO cells more efficiently than that of WT cells (Fig. 2c and Supplementary Fig. 2e). We conclude that both Hsp90 and Hop continue to be functionally required in KO cells, which appear to be even more dependent on them. We checked the other side of the coin of proteostasis with proteasomal inhibitors. The treatments with both MG132 and bortezomib resulted in reduced cytotoxicity with all KO cell lines (Fig. 2d and Supplementary Fig. 2f, g). KO cells also displayed a higher resistance to the ubiquitin-activating enzyme inhibitor PYR41 (Fig. 2e and Supplementary Fig. 2h). These results suggested that KO cells are less dependent both on ubiquitination and proteasomal degradation. Moreover, transient overexpression of WT Hop in KO cells completely reversed the sensitivity to Hsp90 and proteasomal inhibitors (Fig. 2f–h). Since the proteostatic equilibrium could be influenced by changes in protein synthesis, we determined whether the latter is affected. A time-course experiment revealing puromycin-labeled and -released nascent chains showed that the translation rate is only minimally affected (Supplementary Fig. 2i). These results led us to hypothesize that proteostasis in KO cells is ensured by an alternative equilibrium between protein degradation by the proteasome and protein stability/refolding supported by the Hsp70-Hsp90. This raises two questions: (1) Is the function of the ubiquitin-proteasome system compromised in the absence of Hop? (2) How can the Hsp70-Hsp90 molecular chaperone machines function efficiently in the absence of Hop?

The Hsp70-Hop-Hsp90 complex is physically and functionally associated with the proteasome. To determine whether Hop must associate with Hsp70 and Hsp90 for optimal proteasomal function, we carried out immunoprecipitation (IP)-MS experiments with Hop mutants that are defective for Hsp70 or Hsp90 binding. Briefly, we used the TPR mutants K8A and K229A, which do not bind Hsp70 and Hsp90, respectively, and the double mutant, and confirmed the expected interaction patterns by IP (Supplementary Fig. 3a). Based on this result, we defined the Hop-specific interactome by an immunoprecipitation-mass spectrometry (IP-MS) analysis with transfected HEK293T KO cells. After an initial quality control by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (Supplementary Fig. 3b), samples were subjected to label-free liquid chromatography–mass spectrometry (LC/MS-MS) analysis. By comparison with the proteins associated with the TPR double mutant, ~41% were identified only with WT Hop and ~57% were enriched with WT Hop (Fig. 3a and Supplementary Data 3). Thus, proteins of the Hop interactome are largely dependent on the ability of Hop to bind Hsp70 and Hsp90 and may mostly be interactors of Hsp70 and/or Hsp90 rather than direct interactors of Hop itself. The components of the Hsp70-Hsp90 molecular chaperone machines are either enriched or only present with WT Hop, including Hsp70 (HSPA1), Hsc70 (HSPA8), Hsp90a (HSP90AA1), and Hsp90β (HSP90AB1) (Fig. 3b). By comparison, Grp94 (HSP90B1), the endoplasmic reticulum-specific Hsp90 isoform, and not a known interactor of Hop, yields the lowest enrichment (~1.5-fold, Fig. 3b) in this subset of proteins. When we performed a GO term enrichment analysis, we observed that proteasomal and proteasome-associated ubiquitin-related proteins are overrepresented (Supplementary Fig. 3c–e), because these proteins are enriched or only present with WT Hop (Fig. 3c). Interestingly, out of the 19 identified proteasomal components, 16 are associated with the RP of the proteasome (Fig. 3c). We confirmed the interactions of several RP components with Hop-containing complexes by a targeted co-IP analysis with WT HEK293T cells (Fig. 3d). We reevaluated a published MS dataset of IPs of a panel of proteasomal components from yeast. Remarkably, three of the bait proteins jointly pulled down Stil (yeast Hop), Hsp82/Hsc82 (yeast Hsp90s), and Ssa1/2 (yeast Hsp70s), compatible with the conclusion that the ternary complex associates with the proteasome in yeast (Supplementary Fig. 3f). Thus, the Hsp70-Hop-Hsp90 ternary complex is physically associated with proteasomal components, most notably with RP proteins.

The next experiments were designed to test whether the Hsp70-Hop-Hsp90 ternary complex is functionally relevant for the proteasome. We performed an in vitro proteasomal activity assay with extracts and noticed that the total steady-state activity of the 26S/30S proteasome is reduced across all KO cell lines (Fig. 4a and Supplementary Fig. 4a). In contrast, the rate of proteasomal activity is not significantly altered (Fig. 4b and Supplementary Fig. 4b). These results suggested that functional 26S/30S proteasome particles might be less abundant in KO rather than less functional on a per particle basis. We confirmed these in vitro results with an in vivo activity assay for the ubiquitin-proteasome system (UPS); this involved the transient expression and flow cytometric quantitation of a degradation-prone ubiquitin-GFP fusion protein (Ub-R-GFP) in parallel to its prone ubiquitin-GFP fusion protein (Ub-M-GFP) as a control (Fig. 4c and Supplementary Fig. 4c). Upon transient expression of WT and TPR mutant Hop in KO cells, the in vitro proteasomal activity is rescued by WT Hop, whereas TPR single and double mutants are not as efficient and completely dead, respectively (Fig. 4d and Supplementary Fig. 4d). Interestingly, it had been shown that the genetic suppression of Hsp90 function in Drosophila shifts the protein degradation balance from proteasome-mediated degradation to autophagy. However, this is unlikely to compensate for the proteasomal defect of Hop KO cells as the autophagic flux remains unchanged (Supplementary Fig. 4e); moreover, we did not see any upregulation of lysosomal or autophagy-related proteins in our whole-cell proteomic data (Supplementary Data 2). These results established that the Hsp70-Hop-Hsp90 ternary complex is not only physically associated with proteasomal components, but also functionally required for proteasomal activity.

Previous studies in different models had indicated that reduction or deletion of Hsp90 could negatively influence the proteasomal activity. When we deleted the genes encoding ...
Hsp90α (HSP90AA1) or Hsp90β (HSP90AB1) in HEK293T cells, we found a reduced level of steady-state proteasomal activity (Supplementary Fig. 4f). The pharmacological inhibition of Hsp70 or Hsp90 in WT cells similarly led to a reduction of proteasomal activity, but suboptimal concentrations did not show any additive or synergistic effects (Supplementary Fig. 4g). However, proteasomal activity was not affected when GA was added to the extracts after cell lysis (Supplementary Fig. 4h). These results collectively indicate that functional Hsp70 and Hsp90, including the possibility to form a ternary Hsp70-Hop-Hsp90 complex, are essential for optimal proteasomal activity. Furthermore, the requirement for the ternary complex for full proteasomal activity is conserved in yeast: analogously to mammalian cells, the steady-state activity, but not the rate, is reduced in a Δsti1 strain (Fig. 4e and Supplementary Fig. 4i); note that the apparent slight differences in the rates of proteasomal activity are statistically not significant (Supplementary Fig. 4i).
Stability of individual proteasomal components is independent of the Hsp70-Hop-Hsp90 complex. We considered two possibilities to explain the impact of the Hop KO on the proteasome: (1) Proteasomal components are clients of the ternary complex, and in the absence of Hop, they become unstable and subsequently degraded; (2) the ternary complex is required for 26S/30S proteasome assembly and/or maintenance. To address the first possibility, we reanalyzed published datasets of whole-cell proteomic experiments, where cells had been treated with Hsp90 inhibitors. As expected, the levels of intracellular clients of Hsp90 are decreased and Hsp90-associated molecular chaperone and co-chaperone proteins are increased by Hsp90 inhibition. In contrast, Hsp90 inhibitors do not reduce the protein levels of any of the known core proteasomal proteins. Similarly, we did not find any striking and cell line-independent differences in protein levels of proteasomal proteins between WT and KO cells in our own whole-cell proteomic datasets. Thus, it seems unlikely that proteasomal components are dependent on Hsp90 for accumulation and/or stability.

Overall composition of the assembled proteasome is similar in KO cells. To evaluate the impact on proteasome assembly/maintenance, we purified 26S/30S proteasome particles. These were pulled out by an affinity purification strategy targeting the RP protein S5a (human gene name PSMD4); note that this scheme enriches for single- (26S) and double-capped (30S) CPs and discards free CPs or unassembled components. The integrity of purified 26S/30S was analyzed by native gel electrophoresis (Supplementary Fig. 5a, b), and by functional assays (Supplementary Fig. 5c, d). To characterize the composition of the purified proteasome, we performed a comparative label-free LC/MS-MS analysis. We did not find any consistent cell line-independent differences of any identified stoichiometric components of the proteasome between WT and KO cells (Supplementary Fig. 5e, f and Supplementary Data 4). Although we noticed that three substoichiometric components and proteasome chaperones were reduced in preparations from HEK293T KO cells (Supplementary Fig. 5f), we did not further consider them since they were not changed in HCT116 cells (Supplementary Fig. 5e). The aforementioned Hop IP-MS analysis showed that many proteasomal proteins are associated with WT Hop (Fig. 3c). Interestingly, in our own proteomic analyses of purified proteasome from WT cells, all components of the Hsp70-Hop-Hsp90 ternary complex could be identified, albeit at very low substoichiometric levels (Supplementary Data 4). Thus, the association of the Hsp70-Hop-Hsp90 ternary complex with the proteasome, while specific, may only be transient and regulatory. We concluded from these experiments that overall the composition of fully assembled particles is similar without Hop. However, we cannot absolutely rule out the possibility that a minor fraction of purified free RP influenced the overall proteasome composition.

Optimal proteasomal assembly requires the Hsp70-Hop-Hsp90 complex. We studied the structural integrity of the purified proteasome particles by negative staining transmission electron microscopy (TEM). Two-dimensional (2D) class averaging of all visible TEM structures led to four different proteasomal structural projections. We could see side views of single-capped 26S and double-capped 30S proteasome particles, and the two expected top views: an uncapped form with a central hole and a capped form corresponding to CP and RP face up, respectively (Fig. 5a, b). Based on the purification scheme and the relative levels of different forms in the native-PAGE analysis (Supplementary Fig. 5a, b), we did not expect any free CP. We measured the dimensions of all four proteasomal projections and confirmed that regardless of the presence of Hop, they are similar to the known values. The main difference is the higher abundance of the double-capped proteasome (30S) in preparations from WT cells while the single-capped proteasome (26S) is more prevalent in preparations from KO cells (Fig. 5c); the opposite situation applies for the relative abundance of uncapped versus capped particles (Fig. 5d). Collectively, we concluded that the individual proteasomal components are not clients of the Hsp70-Hop-Hsp90 complex, but rather that it is involved in the capping/assembly and/or maintenance of the proteasome (Fig. 5e). We next evaluated the ensemble of proteasome particles in whole-cell extracts by native-PAGE. We detected CP and 26S/30S proteasome particles using a specific antibody against the CP component Psmα3, whose total levels are not significantly altered (Fig. 4h and Supplementary Fig. 6b). We found that the abundance of the 26S/30S particles but not CP is reduced in KO cells (Fig. 5f). As expected, only WT Hop and not the TPR double mutant can rescue proteasome assembly (Supplementary Fig. 6d, e). Note that the results of this rescue experiment and those with the purified proteasome (Supplementary Fig. 5a, b) argue against a technical problem underlying the low ratio of free CP to 26S/30S particles that we see with extracts from HEK293T and A549 cells (Fig. 5f and Supplementary Fig. 6a). Indeed, this ratio varies widely depending on the source, but independently of this, the difference is clearly genotype-dependent in our experiments. Moreover, the reduced proteasome activity in Δstil yeast is mirrored by the reduced assembly and/or stability of proteasome particles, but, as in mammalian cells, not the levels of the individual protein components (Fig. 5g and Supplementary Fig. 6f, g). The assembly defect of the proteasome in Hop KO cells is not due to altered subcellular localization of the proteasomal subunits either.
Thus, the Hsp70-Hop-Hsp90 complex is essential for efficient proteasomal capping and/or for optimal proteasomal stability/maintenance (Fig. 5h).

KO cells are less dependent on the proteasome even with proteotoxic stress. So far, we had compared the Hop requirements for the proteasome under normal conditions. We now turned to investigate the effects of proteotoxic stresses. When cells are treated with the Hsp90 inhibitor GA, more proteins form aggregates (Supplementary Fig. 6), and during a HS, more insoluble and ubiquitinated material accumulates in KO cells (Supplementary Fig. 7); this demonstrates that the reduction of proteasomal activity, and dependency of KO cells on it, is not a symptom of compromised ubiquitination. Upon inhibiting the proteasome in the recovery phase after a 6-h treatment with GA, we observed that WT cells are dying significantly more than KO (Supplementary Fig. 6h, i).
cells (Supplementary Fig. 7c-e). Similarly, inhibiting the proteasome after HS kills WT cells strikingly more than KO cells (Supplementary Fig. 7f, g).

We blocked the proteasomal activity with MG132 for 24 h and found that significantly less ubiquitinated substrates accumulated in KO cells (Supplementary Fig. 7h). This is consistent with a reduced degradation flux through the proteasome in KO cells since overall ubiquitination is not compromised and deubiquitination not augmented in KO cells (see above and Supplementary Fig. 7b). Intriguingly, an inverse and seemingly paradoxical relationship between the resistance to proteasome inhibitors, and proteasomal integrity and even flux had previously been reported. This further supports our conclusions that WT cells not only have a higher abundance of assembled proteasome particles, but that they are also more dependent on proteasomal function, and even more so in stressed conditions.

Hsp70 and Hsp90 functionally collaborate without Hop in vivo. Even though proteasomal assembly and function are compromised in KO cells, overall they nevertheless seem to maintain proteostasis (see Fig. 1b). To substantiate this conjecture, we biochemically fractionated soluble and insoluble proteins from WT and KO cells. Irrespective of genotype, we
found similar amounts of ubiquitinated proteins and protein aggregates (Supplementary Fig. 8a; see also untreated samples in Supplementary Fig. 7a, b). Therefore, more efficient chaperoning functions may compensate for the proteasomal defects of KO cells. We started to explore this with in vivo luciferase refolding assays; we found a higher rate of refolding of heat-inactivated luciferase in KO cells (Fig. 6a and Supplementary Fig. 8b), which could be reverted to that of WT cells by pharmacological inhibition of Hsp90 or Hsp70 (Fig. 6b and Supplementary Fig. 8c). Thus, Hop appears to restrain the refolding activity of Hsp70-Hsp90. Rescue experiments with overexpressed WT and TPR mutants of Hop in KO cells showed that only WT Hop inhibits luciferase refolding whereas the TPR mutants, and most significantly the double mutant, do not affect it (Fig. 6c). Hence, in the absence of Hop/Sti1, Hsp70 and Hsp90 are functional and responsible for enhanced folding of a model substrate.

From the literature, Δstil yeast cells are known to be heat-sensitive, unlike what we have found for human KO cells (Fig. 1e and Supplementary Fig. 1g). To check whether the Hop/Sti1-independent chaperoning mechanism is evolutionarily conserved in yeast, we performed an in vivo luciferase refolding experiment with the HS-sensitive Δstil yeast strain of the W303 background (Supplementary Fig. 8d). Despite its heat sensitivity, this strain displayed a faster rate of luciferase refolding during the early recovery phase (Supplementary Fig. 8e). To mirror the experiments with our HS-resistant human KO cells, we also performed the experiment with a Δstil strain of the BY4741 strain background, in which the mutant is as heat-resistant as the WT (Supplementary Fig. 8f). Here, the enhanced luciferase refolding in the absence of Hop/Sti1 is even more striking (Fig. 6d), and there is a much higher residual luciferase activity after a milder HS (Fig. 6e).

Another aspect of chaperoning is keeping aggregation-prone misfolded proteins in a soluble state. We checked the solubility of the aggregation-prone polyglutamine model protein Q74-EGFP along with a non-aggregating control (Q23-EGFP). We observed a reduction of the aggregated form of Q74-EGFP in HEK293T KO cells (Fig. 6f and Supplementary Fig. 8g). This improved solubility seems to be Hsp90-dependent as it could be substantially suppressed by GA (Fig. 6g). In HCT116 cells, we did not see any difference between the two genotypes (Supplementary Fig. 8h), possibly indicating that cell lines may differ with respect to their intrinsic anti-aggregation activities. Intriguingly, these beneficial effects of the absence of Hop were recently identified by a genetic screen in Drosophila: a Hop knockdown was shown to reduce the aggregation of a Huntingtin mutant with a polyglutamine expansion.

Considering that cell viability and proliferation are relatively unperturbed and if indeed Hsp70 and Hsp90 function even better for some activities in the absence of Hop, we expected only a minimal impact on Hsp90 clients and co-chaperones. To test this hypothesis, we filtered our whole-cell MS dataset for proteins of the Hsp90 interactome (https://www.picard.ch/downloads/Hsp90interactors.pdf). The vast majority of the identifiable Hsp90 interactors are unaffected (Fig. 7a, b and Supplementary Fig. 9a, Supplementary Data 2). Interestingly, the heat-inducible isofrom of Hsp70 (encoded by HSPA1) and Hsp110 (encoded by HSPH1), a nucleotide exchange factor (NEF) of Hsp70, are moderately upregulated in the absence of Hop (Fig. 7a and Supplementary Fig. 9a), whereas the levels of other co-chaperones of the Hsp70-Hsp90 chaperone systems, including several J-proteins, remain unaltered (Fig. 7a and Supplementary Fig. 9a).

To exclude the formal possibility that Hsp90 clients accumulate to normal levels but in an inactive conformation, we checked the activities of different classes of clients. As there are many tyrosine and serine/threonine kinases amongst the clients of Hsp90, we compared total levels of phosphorylated proteins; we could not see any significant global differences between the two genotypes (Supplementary Fig. 9b). We also did not observe any strong differences in the phosphorylation of specific sites indicative of activation of the Hsp90 client c-Src, and of its downstream kinases Erk1/2 (Supplementary Fig. 9c). Using luciferase reporter assays, we checked the transcriptional activities of several transcription factors, which are either Hsp90 clients by themselves or downstream of clients. We found that either the activity is enhanced (in HEK293T cells) or not strikingly compromised (in HCT116 cells) in the absence of Hop (Supplementary Fig. 9d). Thus, Hsp90 clients not only maintain their steady-state levels in KO cells, but they largely also maintain their activity compared to WT cells.

Hsp90 determines a unique spectrum of Hsp90 client proteins.

Since the vast majority of Hsp90 clients is unaltered in KO cells, we wondered whether challenging their proteostasis by overexpression of clients could reveal vulnerabilities. We overexpressed several steroid receptors, which need Hsp90 for proper folding, stability, and transcriptional activity. Indeed, the accumulation and transcriptional activity of the glucocorticoid receptor is compromised in KO cells compared to WT cells (Fig. 7c, e, and Supplementary Fig. 9e). Accumulation and transcriptional activities of other steroid receptors, the estrogen receptor α and the progesterone receptor, are only moderately affected (Fig. 7c–e). v-Src, a viral tyrosine kinase client of Hsp90, accumulated to lower levels associated with reduced
**Fig. 5** Hsp70-Hop-Hsp90 ternary complex is required for proteasome assembly. **a, b** 2D class-averaged images of purified proteasome particles obtained by negative staining TEM. The top 10 and 5 classes are presented here for the side views (panel **a**) and top views (panel **b**), respectively, of both the WT and KO samples. Representative side views of double-capped 30S and single-capped 26S particles, and representative top views of uncapped (CP face up, central hole visible) and capped (RP face up, central hole invisible) particles are indicated. Scale bar, 10 nm. **c, d** Quantitation of the fold changes of 30S over 26S (related to panel **a**) and capped over uncapped (related to panel **b**) proteasome particles between in WT and Hop KO samples. n, total numbers of structural projections of proteasome particles statistically analyzed with the software Relion. **e** Schematic representation of the involvement of the Hsp70-Hop-Hsp90 complex in proteasome particle assembly/maintenance rather than for individual proteasomal proteins. **f, g** Abundance of different proteasomal particles of human (n = 5 independent samples over three independent experiments) and yeast cells (n = 5 independent samples over 2 independent experiments) as indicated displayed by 4% native-PAGE and subsequent immunoblotting (with antibodies against the indicated proteasome component). Positions of 26S/30S proteasome particles and free 20S CP are indicated using the bands of purified proteasome particles as standards. Different exposures, for a given cell line, are from the same immunoblot. Nitrocellulose filters stained with Ponceau S indicate equal loading of proteins. **h** Models of how the Hsp70-Hop-Hsp90 ternary complex could enhance the abundance and stability of assembled 26S/30S proteasome particles.
In vivo luciferase refolding in HCT116 KO cells exogenously overexpressing WT or TPR mutants of Hop. Non-HS controls for each sample are set 1-fold (n = 4 biologically independent samples). d) In vivo refolding of heat-denatured luciferase in WT and Δstl1 yeast cells (BY4741 strain background; two different transformants each). ***p < 0.001 and **p < 0.01 statistically significant differences between WT and Δstl1 (n = 4 biologically independent samples). e) Residual in vivo luciferase activity immediately after mild HS (n = 2 biologically independent samples). Luciferase activity before HS is set to 100%. Residual in vivo luciferase activity immediately after mild HS (HS: 44 °C, 20 min).

Fig. 6 Hsp70 and Hsp90 are functional in vivo even without Hop. a) In vivo refolding of heat-denatured luciferase. Luciferase activity before HS is set to 100%. Left panel, n = 4; right panel, n = 3 biologically independent samples. b) In vivo luciferase refolding in HCT116 KO cells treated with inhibitors before and during the recovery phase (n = 3 biologically independent samples). c) In vivo luciferase refolding in HCT116 KO cells exogenously overexpressing WT or TPR mutants of Hop. Non-HS controls for each sample are set 1-fold (n = 4 biologically independent samples). d) In vivo refolding of heat-denatured luciferase in WT and Δstl1 yeast cells (BY4741 strain background; two different transformants each). ***p < 0.001 and **p < 0.01 statistically significant differences between WT and Δstl1 (n = 4 biologically independent samples). e) Residual in vivo luciferase activity immediately after mild HS (n = 2 biologically independent samples). Luciferase activity before HS is set to 100%. f) Solubility of aggregation-prone polyglutamine model protein. Immunoblots of Q23-EGFP (bands marked with *) and Q74-EGFP (**) from soluble and insoluble protein fractions. GAPDH and H2A.Z serve as loading controls for the insoluble protein fractions. For the line graphs, the data are represented as mean values ± SEM. For box plots, data are represented as the median values and edges of the box plots represent the range of the data. The statistical significance between the groups was analyzed by two-tail unpaired Student’s t-tests. Source data are provided as a Source Data file.

kinase activity in KO cells (Supplementary Fig. 9f, g). Several other Hsp90 clients (Hif-1α, Hif-2α, and androgen receptor (AR)) are not compromised upon overexpression in KO cells (Supplementary Fig. 9h). For endogenous GR as well, both expression and transcriptional activity are remarkably reduced in A549 KO cells (Fig. 7f). Overall, these results suggested that affected steroid receptors, kinases such as v-Src, and the proteasome are rather the exceptions amongst Hsp90 clients with regards to their pronounced Hop-dependence; anecdotally, GR and v-Src are also the very same Hsp90 clients that were the first ones to be discovered. In agreement with a previous publication showing the plasticity of the Hsp90 co-chaperone network for folding of exogenous Hsp90 clients in yeast, we hypothesized that Hop might determine unique criteria of client selectivity for Hsp90 even in higher eukaryotes.

An alternative prokaryote-like Hsp70-Hsp90 binary complex maintains the Hsp90 interactome. KO cells maintain proteostasis by more efficient Hsp70- and Hsp90-dependent chaperoning (Fig. 8a). Since bacterial and organellar Hsp90 and Hsp70 orthologs can interact without a Hop-like protein, we investigated the possibility that an alternative Hop-independent Hsp70-Hsp90 complex might form in the cytosol of human cells. With an Hsp90 IP-MS analysis, we found Hsp70/Hsc70, encoded by HSPA1 and HSPA8, among the top hits even in KO cells, despite a 4-10-fold reduction (Fig. 8a and Supplementary Fig. 10a, b). In vivo luciferase refolding in HCT116 KO cells treated with inhibitors before and during the recovery phase (n = 3 biologically independent samples). c) In vivo luciferase refolding in HCT116 KO cells exogenously overexpressing WT or TPR mutants of Hop. Non-HS controls for each sample are set 1-fold (n = 4 biologically independent samples). d) In vivo refolding of heat-denatured luciferase in WT and Δstl1 yeast cells (BY4741 strain background; two different transformants each). ***p < 0.001 and **p < 0.01 statistically significant differences between WT and Δstl1 (n = 4 biologically independent samples). e) Residual in vivo luciferase activity immediately after mild HS (n = 2 biologically independent samples). Luciferase activity before HS is set to 100%. f) Solubility of aggregation-prone polyglutamine model protein. Immunoblots of Q23-EGFP (bands marked with *) and Q74-EGFP (**) from soluble and insoluble protein fractions. GAPDH and H2A.Z serve as loading controls for the insoluble protein fractions, respectively. g) Immunoblot of Q74-EGFP from soluble and insoluble protein fractions of KO cells treated with GA. Hsp90 even in higher eukaryotes.
Transfected KO only affects few specific Hsp90 clients. a Heat maps of the normalized fold changes of the levels of Hsp90 clients (left), and molecular chaperones and co-chaperones (right) identified by whole-cell MS analyses. b Immunoblots of Hsp90 clients, Hsp90 partners and co-chaperones (n = 3 independent experiments). c Immunoblots of Flag-tagged GR (F-GR) and estrogen receptor α (F-ER) overexpressed in HEK293T cells (n = 2 independent experiments). d Immunoblots of the progesterone receptor (PR) overexpressed in HEK293T cells (n = 2 independent experiments). e Transcriptional activities of overexpressed GR (induced by dexamethasone (DEX)), ERα (induced by β-estradiol (E2)), and the PR (induced by progesterone (PROG)), as assayed with specific luciferase reporter genes (n = 2 biologically independent samples); data are represented as fold change relative to those of WT cells (set to 1). f Immunoblots of the endogenous levels of GR (top, n = 2 independent experiments), and its DEX-induced transcriptional activity determined with a transfected luciferase reporter (bottom, n = 2 biologically independent samples). For box plots, data are represented as the median values and edges of the box plots represent the range of the data. Source data are provided as a Source Data file.

Supplementary Data 2). We confirmed this Hop-independent interaction of Hsp70 and Hsp90 by a targeted co-IP experiment (Fig. 8c and Supplementary Fig. 10b).

In search of a mechanism promoting a Hop-independent interaction of Hsp70 and Hsp90, we checked whether Hop activity might be functionally redundant in human cells. We revisited our own Hsp90 IP-MS datasets and extracted the data for proteins with a Hop-like architecture. There are only very few proteins with multiple TPRs, which are enriched with Hsp90 in the absence of Hop, and they are very moderately so (Supplementary Fig. 10c). Most importantly, all of these “enriched” TPR proteins are present at very low levels compared...
to the abundance of Hsp90 and Hsc70/Hsp70 in Hsp90 immunoprecipitates of either WT or KO cells, and to Hop in WT cells (Supplementary Fig. 10d). Thus, it is very unlikely that any of these proteins could substitute for Hop for the formation of another Hsp70-(multiple TPR protein)-Hsp90 ternary complex.

Using recombinant proteins, we tested the notion that human Hsp70 and Hsp90 could directly interact as they do in bacteria\(^28,29\) and in \(\Delta\textit{stl1}\) yeast\(^60\). To minimize client-type interactions of Hsp70, we used the substrate-binding mutant V438F of Hsp70. The co-IP experiment revealed a direct interaction between Hsp90 and Hsp70 both in the presence and absence of ATP.
The prokaryote-like human Hsp70-Hsp90 complex is also more efficient in vitro. Substrate folding can be obtained by the collaboration of bacterial Hsp70 with Hsp90 in vitro. To evaluate the chaperoning activity of the binary Hsp70-Hsp90 complex, we performed in vitro luciferase refolding experiments with purified components. Human Hsp70 was complemented with a J protein, the Nterminal Acp2, and human Hsp90a. We measured luciferase refolding as a function of increasing concentrations of Hop. To our surprise, but in agreement with our in vivo experiments (Fig. 6a and Supplementary Fig. 8b), we discovered that the Hsp70-Hsp90 molecular chaperone systems refold heat-denatured luciferase most efficiently in the absence of Hop (Fig. 9b). Increasing concentrations of Hop gradually decrease the final yield achieved by the Hsp70-Hsp90 system (Fig. 9b). Moreover, only WT but not the Hsp70-binding mutant K418A,K419A of Hsp90a stimulates Hsp70-mediated refolding of luciferase in the absence of Hop (Fig. 9c), confirming the functionality of the human prokaryotic-like binary complex of Hsp70-Hsp90 (Fig. 9d).

Discussion
The proteostatic equilibrium can be maintained by alternative mechanisms. For example, HS or inhibition of Hsp90 strongly induce the Hsp70-Hsp40 chaperone system and small Hsps to meet the new cellular requirements. Here, we establish cellular models with human cell lines and yeast, which, in the absence of Hop/Sti1, adopt the more ancient and more efficient mechanism of chaperoning of bacteria and thereby compensate for proteasomal defects, reestablishing an alternative proteostatic equilibrium. Thus, depending on cell-specific requirements or external inputs, eukaryotic cells may still be able to shift to a more prokaryotic-like or organellar mode of operation for the Hsp70-Hsp90 systems. We speculate that the "invention" of Hop during the evolution from prokaryotes to eukaryotes may have promoted a shift from a proteostatic system centered on refolding to a more extensive use of proteasomal degradation.

Hsp90 itself had already been linked to proteasomal integrity and activity. Our data complement this early evidence by demonstrating that it is the Hsp70-Hop-Hsp90 ternary complex that is required for optimal 26S/30S assembly and/or maintenance. According to current models, the RP ATPase-ring docks to the CP α-ring to form the functionally active holoenzyme. The substoichiometric proteasome component Ecm29, which itself is not dependent on Hop (Supplementary Fig. 5e, i), has been claimed to be a tethering factor for RP and CP. We propose that the ternary complex plays a similar role; further studies are required to define more clearly how the ternary complex promotes RP-CP docking and what the interplay with other factors might be. Our MS analysis showed that all components of the ternary complex are associated with 26S/30S particles at substoichiometric levels; hence, it is conceivable that the ternary complex acts like other bona fide proteasome-dedicated chaperones, which dissociate from the functionally mature proteasomal holoenzyme. Although activity and assembly are known to be regulated by posttranslational modifications of its components, it is not likely that this can help to explain the proteasomal defects of KO cells; the vast majority of proteins, in general, and several of the known posttranslational regulators of proteasomal components such as MAPKs, mTORC1-related proteins, and protein kinase A, in particular, are unaltered in KO cells (Supplementary Data 2). Thus, we conclude that Hop, as part of the Hsp70-Hop-Hsp90 complex, primarily binds to the CP, facilitates the docking with the CP, and stabilizes the assembled 26S/30S proteasome.

Our most surprising finding is that cells lacking Hop/Sti1 compensate the proteasomal defect by improved protein folding. In view of the well-established role of Hop for substrate transfer between Hsp70 and Hsp90, and as allosteric regulator of the Hsp90 ATPase activity, this is the last thing one would have expected. Hop function may be beneficial or even essential to allow the folding or assembly of some substrates such as GR, v-Src, the proteasome, and possibly of some particularly labile clients such as the ΔF508 CFTR mutant. TPR mutants of Hop, including the residual fragment of HEK293 KO1 cells, are unable to rescue Hop functions in KO cells. It should be acknowledged here that we do not have formal biochemical proof that Hop acts as part of a ternary complex in these contexts, but the fact that it must be able to form such a complex makes it very likely. In contrast to the aforementioned clients, for most Hsp90
clients, the temporary slow-down of the Hsp90 molecular chaperone machine by Hop during substrate transfer may be counterproductive. What defines the Hop requirements of Hsp90 clients remains to be deciphered, but it is clear from our data that the vast majority of Hsp90 clients can be efficiently processed by the prokaryote-like Hsp70-Hsp90 binary complex. We assume that this core molecular chaperone complex still requires the support of other co-chaperones like Hsp40, Cdc37, Aha1, and p23 to select, to process and to release Hsp90 clients. Compromising the function of these co-chaperones could be particularly detrimental for Hop KO cells running on a somewhat hyperactive Hsp90 machine; indeed, the deletion of *SBA1*...
This could be a useful strategy to reequilibrate the proteostatic balance in diseases or altered physiological states with proteasomal dysfunctions. Neurodegenerative disorders such as Huntington’s and Parkinson’s diseases, which may be associated with inefficient chaperoning, could potentially benefit from improved chaperoning induced by Hop inhibitors. Some compounds have been reported to inhibit specifically the interaction between Hsp90 and the TPR2A domain of Hop, and this is promising leads for such a therapeutic strategy.

Methods

Additional methods are available in Supplementary Information.

Reagents and resources

Details on all reagents and resources are provided in Supplementary Data 5.

Cell lines and cell culture

HEK293T human embryonic kidney cells, HCT116 human colorectal carcinoma cells and A549 human lung epithelial carcinoma cells (as well as the corresponding Hop KO cell lines) were maintained in Dulbecco’s Modified Eagle Media (DMEM) supplemented with GlutaMAX, 10% fetal bovine serum (FBS) and penicillin/streptomycin (100 µg/ml) with 5% CO₂ in a 37 °C humidified incubator.

Yeast strains

Yeast strain BY4741 (his3Δ1 leu2Δ0 met15Δ0 ura3Δ0) and its Δstl1 variant Y101803 (his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 Δstl1::kanMX4), strain W303 (ΔBMA64-1A) (ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1) and its Δstl1 variant W303 Δstl1 (ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 Δstl1::kanMX4) were maintained on yeast extract peptone dextrose (YPD) agar plates or in broth at 30 °C. To generate the W303 Δstl1 strain, W303 cells were transformed with a PCR fragment encompassing the Δstl1::KanMX locus of BY4741 and selected with geneticin (G418, 200 µg/ml) on YPD agar plates. Deletion of STI1 was verified by PCR amplification and by immunoblot analysis with a specific antibody to the Sti1-TPR2A/B domain. BY4741 and W303 yeast cells were transformed with plg/LUC, a galactos-inducible luciferase expression plasmid, to produce yeast strains expressing firefly luciferase.

Plasmids

Site-directed mutagenesis was performed on pCDNA3.1(+) (Hop) (WT) to generate human Hop mutants K8A (codon change: AAG > GCG), K229A (AAA > GCA), and the double mutant K8A/K229A, each with a C-terminal HA-tag. Site-directed mutagenesis was performed on the bacterial expression vector pCAS28-Hsp90α (WT) and mammalian expression vector pFLAG-CMV2/Hsp90α (WT) to generate the plasmids for the human Hsp90α mutants K418A (codon change: AAA > GCA), K419A (AAA > GCA), and the double mutant K418A/K419A, each with either a 6x-His or FLAG tags. Sequences of all oligonucleotides (from Microsynth) are described in Supplementary Data 5.

Genome engineering

Hop (STI1) KO HEK293T, HCT116, and A549 cells, as well as Hsp90α (HSP90AA1) and Hsp90β (HSP90AB1) KO HEK293T cells were generated by the CRISPR/Cas9 gene editing technology. The guide RNAs (gRNAs) were identified by the corresponding oligos designed using the ATUM (previously DNA2.0) CRISPR/Cas9 design tool (https://www.atum.bio/). The gRNA sequences corresponding to the 5th exon of STI1, 1st exon of HSP90AA1, and 6th exon of HSP90AB1 are listed in Supplementary Data 5. Sense and antisense oligos were synthesized (Microsynth), annealed and cloned into the Bsal site of px459 (Addgene plasmid #48139) as described previously. WT HEK293T, HCT116, and A549 cells were transiently transfected with the gRNA plasmids using polyethyleneimine (PEI). At 48 h post-transfection, transfected cells were selected with puromycin (3–5 µg/ml). Surviving cells were further cultured in the absence of puromycin until visible cellular foci formed. Cellular foci were individually picked and analyzed by immunoblotting using primary antibodies specific to Hop, Hsp90α, and Hsp90β. Clones which did not express Hop, Hsp90α or Hsp90β were

(continuing from previous text...)

...cellular or organismic fitness could nevertheless be negatively affected in the long run. Indeed, even though Hop KO worms are viable, their lifespan is reduced. Furthermore, Hop levels drop in the aging brain, and aging is correlated with reduced proteasomal and chaperoning activity. Since the absence of Hop generates a mixed outcome in different experimental models, tissue- and cell type-specific functions of Hop for the maintenance of proteostasis should be further studied.

Our discoveries may also have translational potential. If Hop could be specifically inhibited, this might promote superior chaperoning by the prokaryote-like Hsp70–Hsp90 binary complex.
considered KO cells and frozen in liquid nitrogen. Some KO clones were also validated by MS analysis. The frequency of obtaining Hop KO clones with these human cell lines ranged between 23–46% indicating that Hop is not absolutely essential in human cells similarly to what had previously been found for the budding yeast Saccharomyces cerevisiae.32 The Hsp90α and Hsp90β KO cells will be described in more detail elsewhere.

Flow cytometry. For all flow cytometric analyses, a minimum of 10,000 cells were analyzed for each sample. We used a FACScalibur (BD Biosciences) with the software package CellQuest Pro and a FACS Gallios (Beckman Coulter), and data were analyzed by the software packages FlowJo and CellQuest Pro. Additional details are given in the following paragraphs. Gating and data analysis strategies are presented in Supplementary Figs. 11 and 12.

Annexin V-PI staining. Human cells seeded at a density of 5 × 10^4 per 2 ml were harvested by trypsinization, washed in phosphate-buffered saline (PBS) and resuspended in 100 µl annexin V-binding buffer (10 mM HEPES pH 7.4, 150 mM NaCl, 2.5 mM CaCl2). Annexin V-FITC (5 µl) and propidium iodide (PI; 2.5 µg/ml) were added to the cells and incubated for 15–20 min at 4°C. In the dot plot analyses, the first quadrant represented healthy, unstained cells; the second quadrant represented early apoptotic cells with only annexin V-FITC staining; the third quadrant represented late apoptotic cells with both annexin V-FITC and PI staining; and the fourth quadrant represented necrotic cells with PI staining.

Cell death assays. Human cells were resuspended in 100–200 µl of PBS-containing PI (2.5 µg/ml) for 15–20 min at room temperature (RT) before flow cytometric analysis.

Cell cycle analyses. Cells were harvested as outlined above. Cells were fixed with 70% ice-cold ethanol, washed, treated with 100 µg/ml RNase A at RT for 5 min, then incubated with 50 µg/ml PI for 15–20 min at RT before flow cytometric analysis. Apoptotic cells were identified by the quantitation of the subG0 cell population.

Autophagic flux measurement. Cells seeded at a density of 4 × 10^4 per 2 ml were transfected with the autophagy reporter plasmid FWU mCherry-GFP-LC3 using PEI. At 48 h post-transfection, cells were harvested by trypsinization, and GFP- and mCherry-positive cells were measured by flow cytometry. The autophagic flux was measured by calculating the ratio of the mean fluorescence intensity of GFP-negative and mCherry-positive cells. A lower relative ratio of GFP/mCherry is indicative of a higher autophagic flux.

Cell proliferation assay. Human cells seeded at a density of 4 × 10^4 per 200 µl were grown for 24, 48 or 72 h after which, 100 µg/ml MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) was added to the culture medium for 3 h. The culture medium was removed, the formazan crystals were dissolved in DMSO, and the OD measured at 550 nm with a plate reader (Tecan Sunrise).

Biochemical fractionation of soluble and insoluble proteins. To analyze the polyglutamine (polyQ) protein aggregation within cells, cells seeded at a density of 7 × 10^3 per 2 ml were transfected with plasmids pEGFP-Q74 and pEGFP-Q23 with PEI. At 48 h post-transfection, cells were harvested for biochemical fractionation. For some experiments, at 24 h post-transfection, cells were treated with GA (0.75 µM) for 24 h. For the comparative analysis between WT and KO cells, soluble and insoluble fractions were biochemically separated from 4 × 10^6 cells for each genotype as described previously. Briefly, cells were lysed in a lysis buffer with mild detergents (20 mM Tris-HCl pH 7.4, 2 mM EDTA, 150 mM NaCl, 1.2% sodium deoxycholate, 1.2% Triton X-100, 200 mM iododecametamide, protease inhibitor cocktail [PIC]), sonicated (low power, three cycles of 10 s pulses), and centrifuged at 16,100 × g for 20 min. The supernatant was collected as the soluble fraction. The precipitate (insoluble fraction) was washed 5–6 times with PBS and solubilized in NuPAGE protein sample buffer (Life Technologies, cat. no. NP0008) complemented with 10 mM DTT. Both biochemical fractions were analyzed by immunoblotting.

In vitro proteasomal activity assay. Human cells were harvested and washed. Cell pellets were resuspended in lysis buffer (25 mM Tris-HCl pH 7.4, 250 mM sucrose, 5 mM MgCl2, 1% NP-40, 1 mM DTT, 1 mM ATP) and incubated for 10–15 min on ice. Samples were centrifuged at 16,100 × g for 20 min and supernatants were collected for the proteasomal activity assay. Yeast cells were collected, washed with H2O and lysed mechanically with glass beads (3 × 30 s pulses) in lysis buffer (10 mM Tris-HCl pH 7.4, 50 mM NaCl, 10 mM MgCl2, 1 mM EDTA, 20% glycerol, 1 mM DTT, 1 mM ATP). Samples were centrifuged at 16,100 × g for 20 min and supernatants were collected for the proteasomal activity assay. Equal amounts of protein (25–50 µg) for each sample were diluted in proteasomal reaction buffer (50 mM Tris-HCl pH 7.4, 5 mM MgCl2, 1 mM Na-citrate buffer (pH 5.0) and 1 mM D-luciferin, before luminescence was measured with a luminescence plate reader. The luciferase activity of cells not subjected to HS was set to 100%.

In vivo UPS activity assay. Cells seeded at a density of 5 × 10^5 per 2 ml were transfected with plasmids Ub-M-GFP (stable GFP) and Ub-R-GFP (degradation-prone GFP) with PEI. At 48 h post-transfection, cells were harvested by trypsinization and GFP-positive cells were quantified by flow cytometry. The in vivo UPS activity was expressed as the % Ub-M-GFP-positive cells minus the % Ub-R-GFP-positive cells.
Immunoprecipitation (IP). Hsp90α, Hsp90β, Hop, and FLAG-Hsp90α IP. Human cells were resuspended in HSP90 complex lysis buffer (10 mM Tris-HCl pH 7.5, 50 mM NaCl, 1 mM EDTA, 1 mM DTT, 1% Triton X-100, 0.01% TNF-α, 0.01% Triton X-100, PICO) and lysed by sonication (30–40 cycles of 30 s) using a Bioruptor® sonicator (Diagenode). For Hsp90α or Hsp90β IP, 1 mg of clarified cell extract was mixed with 5 µg anti-Hsp90α (RDI) or anti-Hsp90β (HRH-10) antibodies. For FLAG-Hsp90α IP, 1 ng of cell extract was mixed with 10 µg anti-FLAG M2 antibodies. For Hop IP to detect associated proteasomal components, 2–3 mg of cell extract was mixed with 5 µg anti-Hop antibodies. For all experiments, IPs were incubated on 4 °C at 30 °C and the association of Hsp90 and Hsp70 was determined by a co-IP. Briefly, 5 µg anti-Hsp90α (RDI) or anti-Hsp90β (HRH-10) antibodies along with 25 µl of Dynabeads®-Protein G were added to the protein mixtures and incubated on 4 °C at 4 °C. Equal amounts of normal IgG were used for the corresponding control IP. Dynabeads were washed and boiled with the NuPAGE protein sample buffer complemented with 10 mM DTT, and eluates were collected. Further SDS-PAGE and immunoblotting were performed with eluted protein samples, using purified proteins as inputs.

Native PAGE analysis. For experiments to visualize proteosomal complexes in total cell lysates, human and yeast cells were lysed in 25 mM HEPES-KOH pH 7.4, 40 mM KCl, 5 mM MgCl2, 10% glycerol, 1 mM Na-molybdate, 0.1% Triton X-100, PICO) and lysed by sonication (30 cycles of 30 s) with a water bath sonicator and by shaking with glass beads with three 30 s pulses, respectively. Equal amounts of total cell extracts (40–75 µg) or purified proteosomes (2 µg) were separated by 4% Tris-HCl native-PAGE. For all samples, native-PAGE was performed for 3.5–4 h at 100–110 V in a 4 °C cold room. Separated proteins were transferred onto nitrocellulose membrane and visualized with proteasomal subunit-specific primary antibodies.

In vitro protein-protein interaction assay. Five micrograms of purified WT or point mutant Hsp90α or WT Hsp90β protein were mixed with 7.5 µg of purified Hsp90 (V438F) in association buffer (10 mM Tris-HCl pH 7.5, 50 mM NaCl, 1 mM EDTA, 1 mM DTT, 0.01% Triton X-100, 10% glycerol, 10 mM Na-molybdate, PICO) in the presence or absence of 5 mM ATP. The mixtures were incubated for 2–3 h at 30 °C and the association of Hsp90α and Hsp70 was determined by a co-IP. Five micrograms of purified Hsp90α (RDI) or anti-Hsp90β (HRH-10) antibodies along with 25 µl of Dynabeads®-Protein G were added to the protein mixtures and incubated O/N at 4 °C. Equal amounts of normal IgG were used for the corresponding control IP. Dynabeads were washed and boiled with the NuPAGE protein sample buffer complemented with 10 mM DTT, and eluates were collected. Further SDS-PAGE and immunoblotting were performed with eluted protein samples, using purified proteins as inputs.

Assay of protein translation rate. Cells seeded at a density of 5 × 10^6 per 10 ml were treated with puromycin (1 µM) for 2–6 h, harvested and lysed in 20 mM Tris-HCl pH 7.4, 2.4 mM EDTA, 150 mM NaCl, 1% Triton X-100, 0.2% Nonidet P-40, 75 µg) or purified cell lysate in 35 min). Full MS survey scans were performed at 120,000 resolution. All survey scans were externally calibrated using the 445.1200 background ion mass. Using a data-dependent acquisition strategy controlled by Xcalibur 2.1 software (Thermo Fisher), a maximum ion count number of multi-charged precursor ions was selected for tandem MS acquisition within a maximum cycle time of 3 s. Selected precursor ions were fragmented by Collision-Induced Dissociation (CID) and analyzed in the linear ion trap with an isolation window of 1.6 m/z. Selected ions were then dynamically excluded from further selection during 60 s.

Confocal microscopy. Hop WT and KO A549 cells were fixed with 4% formaldehyde and subsequently permabilized with 0.1% Triton X-100. Cells were blocked with 1% bovine serum albumin and incubated with primary antibodies against Psma3, Psmc5, and Psmde6 (all at a 1:200 dilution) O/N in a moist chamber at 4 °C. Appropriate AlexaFluor488-conjugated secondary antibodies were used at a 1:2000 dilution at RT for 1 h. After DAPI staining, cells were mounted with mounting medium and data acquisition was performed with a Leica TCS SP8 confocal microscope with a 63x oil immersion objective. Digital images were acquired with 2x digital magnification. Images were analyzed with the software ImageJ-Fiji.

Transcriptional activity assays. Human cells were seeded at a density of 6 × 10^4 per 0.5 ml in phenol red-free DMEM medium containing 5% charcoal-treated FBS in 24-well cell culture plates. The cells were co-transfected with ERTα, GR or PR mammalian expression plasmids (only for HEK293T cells), as well as both a luciferase reporter plasmid and a constitutive Renilla luciferase expression plasmid (pRL-CMV). In another experiment, cells were transfected with luciferase reporter plasmids as described for Supplementary Fig. 9d along with pRL-CMV. At 48 h post-transfection, cells were lysed with Passive Lysis Buffer, and Firefly and Renilla luciferase activities were measured using the Dual-Luciferase detection kit (Promega) with a bioluminescence plate reader. Renilla luciferase activity was used as control transcription.

RNA extraction and quantitative reverse transcription PCR (qRT-PCR). Cells were lysed with the guanidine-acid phenol method by using the TRI reagent (4 M guanidine thiocyanate, 25 mM sodium citrate, 0.5% N-lauroylsarcosine, 0.1 M 2-mercaptoethanol, pH 7). Then sequentially 2 M sodium acetate pH 4, aquaphenol and chloroform/isoamyl alcohol (49:1) were added to the lysates and mixed. Organic and aqueous phases were separated by centrifugation at 10,000 × g for 20 min and the aqueous phases were collected. RNA was precipitated by adding isopropanol and centrifugation at 12,000 × g for 20 min at 4 °C. RNA pellets were washed with 75% ethanol and resuspended in nucleic-acid free water. cDNA was prepared from RNA (400 ng) by using random primers (Promega), GoScript buffer (Promega), and reverse transcriptase (Promega) according to the manufacturer’s instructions. cDNAs were mixed with the GoTaq master mix (Promega) and specific primer pairs for STIP1 (Supplementary Data 5) for quantitative PCR with a Biorad CFX96 thermocycler. Expression of mRNA of the gene of interest was normalized with GAPDH as internal standard.

Bioinformatic analyses. In silico protein homology modeling. Beginning with the apo conformation of HspG (PDB ID: 2IQO90), homology models of the human Hsp90α and Hsp90β proteins were constructed using the I-TASSER structure prediction web server95. Briefly, monomeric structures were predicted according to the C-score and TM-score of Hsp90α (C-score = −1.53, TM-score = 0.53 ± 0.15) and Hsp90β (C-score = −0.99, TM-score = 0.59 ± 0.14). To generate dimer structures of Hsp90α and Hsp90β, full-length predicted monomeric structures of each protein were manipulated using HPLC system (Thermo Fisher Scientific, Bremen, Germany). Extracted peptide mixtures were separated on a custom-packed reversed-phase C18 nanocolumn (75 µm ID × 20 cm, 1.8 µm particle size, Reprosil Pur, Dr. Maisch) with a gradient from 5% to 55% acetonitrile in 0.1% formic acid for 120 min (for IP samples: same gradient in 35 min). Full MS survey scans were performed at 120,000 resolution. All survey scans were internally calibrated using the 445.1200 background ion mass. Using a data-dependent acquisition strategy controlled by Xcalibur 2.1 software (Thermo Fisher), a maximum ion count number of multi-charged precursor ions was selected for tandem MS acquisition within a maximum cycle time of 3 s. Selected precursor ions were fragmented by Collision-Induced Dissociation (CID) and analyzed in the linear ion trap with an isolation window of 1.6 m/z. Selected ions were then dynamically excluded from further selection during 60 s.
the ClusPro protein–protein docking web server programmed in dimer mode. Hydrophobic interaction driven dimer structures, with low free energy, were selected. The accessibility of surface residues of the dimeric Hsp90α and Hsp90β were predicted with the Swiss-PdbViewer 4.1.1 software. Note that our modeling and the surface-exposure of the highlighted residues are supported by a recent study of Tastan Bock and colleagues.

**Gene ontology enrichment analysis.** GO enrichment analyses with the whole-cell MS datasets were done with those proteins, which changed the most (fold change >0.4 or smaller than −0.4 and with a VIP value over (1) with the ClusterProfiler package for R. Overrepresentation of “Biological Process” GO terms in the data of Hop KO cells was determined by using the database “org.Hs.es.db” from Bioconductor. GO enrichment analyses for the database (HA)-IP-MS dataset was performed with the Enrichr web server (http://amp.pharm.mssm.edu/Enrichr), focusing on KEGG and WIKI pathway-annotated proteins. Significant biological processes were identified by lower p-values, a higher number of common genes/proteins responsible for a biological process, and a high combined score provided by the Enrichr web server.

**Protein interaction network analysis.** Fold change values (WT vs KO) of proteins from the whole-cell MS analyses were analyzed as described previously. All nodes correspond to identified proteins. Cytoscape analyses were represented as heat maps of a log2 fold change of protein expression. Hsp90-specific clients, chaperones and co-chaperones were classified using our own web server for Hsp90 interactors (https://www.picard.ch/Hsp90Int/index.php).

**General data analyses.** Data processing and analyses were performed using GraphPad Prism (versions 7 and 8).

**Mass spectrometric data analyses.** Tandem MS data were processed by the MaxQuant software (ver.1.6.0.13) incorporating the Andromeda search engine. The UniProt human reference proteome database (of October 2017) was used (71,803 sequences), supplemented with sequences of common contaminants. All identifications were filtered at 1% false discovery rate at both the peptide and protein levels with default MaxQuant parameters. For the whole-cell proteome and Hsp90 IP-MS experiments, from the relevant MS datasets only protein groups were kept for the subsequent analyses, which met the criteria of at least two unique peptides, four peptides, and six MS/MS counts. Protein quantitation was performed either with the iBAQ values (for the IP-MS and proteasomal analyses) or with the LFQ values (for the whole-cell proteome analyses). In whole-cell proteome and Hsp90 IP-MS analyses, the log2 fold change >0.4 or ≤−0.4 of a protein was considered as the biologically significant difference. Proteins that were detected in at least one of three biological replicates (for the whole-cell proteome) were used in all further analyses.

In Hsp90 IP-MS analyses, the criteria for the validation of interactors was an average fold change of 8 (3 in log2 scale) or more, identified by MS/MS in all positive IP samples compared to control IP, and an adjusted p-value <0.05. The iBAQ values of interactors were normalized on the sum of the iBAQ of Hsp90α and Hsp90β.

In Hop (HA) IP-MS analyses, the interactor validation criteria was an average fold change of 40 or more identified by MS/MS in WT Hop IP samples compared to the control IP and at least 3 unique peptides. Since chaperone, co-chaperone and proteasomal proteins were our focus, we included three proteasomal components (PSMB1, PSMD7, and PSMD14) and three molecular chaperones (HSP90AB1, HSP6, and HSP7) in further analyses despite them being represented by only 2 (PSMB1, PSMD7, and PSMD14) and three molecular chaperones (HSP90AB1, HSP6, and HSP7) in further analyses despite them being represented by only 2 unique peptides. The iBAQ values of interactors were normalized either on the abundance of WT or TRP mutant Hop depending on the experimental set. PCA and OPLS-DA analysis of the whole cell proteome and Hsp90 IP-MS experiments were done using the ropls package for R. The variable importance in projection (VIP) values were derived from the “Orthogonal Projections to Latent Structures Discriminant Analysis” (OPLS-DA) model and used to select variables of interest with a VIP cut-off of 1.95.

**Proteasomal activity analyses.** For steady-state proteasomal activity measurements, AMC fluorescence of WT samples for each time point was considered as 100% proteasomal activity and the corresponding proteasomal activity of KO samples was calculated relative to that. For the proteasomal activity rate measurements, AMC fluorescence of each time point was normalized with AMC fluorescence at 5 min; data were represented as the fold change while slopes were considered as the rate of proteasomal activity over time.

**TEM image analyses.** Eight hundred twenty-seven and 705 micrographs were acquired for proteasome samples from Hop WT and KO HEK293T cells, respectively, at 2.245 Å/pix using an Eagle detector (Thermofisher). The micrographs were analyzed using Relion (version 3.1)130. 106,491 and 92,102 particles were picked for the WT and KO samples, respectively, using Gaussian picking implemented in Relion. The particles were initially classified for four rounds. For the WT sample, 17,342 and 7314 particles were selected for the top and side views, respectively, and for the KO sample, 20,120 and 4410 particles were selected for the top and side views, respectively, and each classified separately for another two rounds. Finally, the particles of good classes corresponding to each view and sample were selected and statistically analyzed.

**Cell death analysis.** The calculations of the percent dead cells for the experiments of Supplementary Fig. 7斐 were performed as follows. First, the background of dead cells in the controls were subtracted from the corresponding MG132-treated sets for both 37°C and HS-treated cells. Then the percent dead cells at 37°C of a particular genotype was subtracted from the percent dead cells of the same genotype under HS treatment and the final measurement was considered the percent cell death under the experimental condition of HS + MG132. This explains how some cells or some conditions can yield negative values.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

All data supporting the findings of this study are available from the corresponding authors upon reasonable request. The mass spectrometric proteomic data are available through the ProteomeXchange Consortium with the identifier PXD012774 and for a subset in Supplementary Data 1–4. Source data of uncropped immunoblots images and the individual data points are provided with this paper. Source data are provided with this paper.

Received: 8 June 2020; Accepted: 30 October 2020; Published online: 25 November 2020.

**References**

1. Balch, W. E., Morimoto, R. I., Dillin, A. & Kelly, J. W. Adapting proteostasis for disease intervention. Science 319, 916–919 (2008).

2. Hipp, M. S., Park, S. H. & Hartiala, F. U. Proteostasis impairment in protein-misfolding and -aggregation diseases. Trends Cell Biol. 24, 506–514 (2014).

3. Labbadia, J. & Morimoto, R. I. The biology of proteostasis in aging and disease. Annu. Rev. Biochem. 84, 435–464 (2015).

4. Wong, E. & Cuervo, A. M. Integration of clearance mechanisms: the proteasome and autophagy. Cold Spring Harb. Perspect. Biol. 2, a006734 (2011).

5. Chen, B., Retzlaff, M., Roos, T. & Frydman, J. Cellular strategies of protein quality control. Cold Spring Harb. Perspect. Biol. 3, a004374 (2011).

6. Balchin, D., Hayer-Hartl, M. & Hartl, F. U. In vivo aspects of protein folding and quality control. Science 335, aa4354 (2016).

7. Schmidt, M. & Finley, D. Regulation of proteasome activity in health and disease. Biochim. Biophys. Acta 1843, 13–25 (2014).

8. Echeverria, P. C. & Picard, D. Molecular chaperones, essential partners of steroid hormone receptors for activity and mobility. Biochim. Biophys. Acta 1803, 641–649 (2010).

9. Taipale, M. et al. Quantitative analysis of HSP90-client interactions reveals principles of substrate recognition. Cell 150, 987–1001 (2012).

10. Picard, D. Preface to Hsp90. Biochim. Biophys. Acta 1823, 605–606 (2012).

11. Schopf, F. H., Biebl, M. M. & Buchner, J. The Hsp90 chaperone machinery. Nat. Rev. Mol. Cell. Biol. 18, 345–360 (2017).

12. Morán Luengo, T., Mayer, M. P. & Rüdiger, S. G. D. The Hsp70/Hsp90 chaperone cascade in protein folding. Trends Cell Biol. 29, 164–177 (2019).

13. Radli, M. & Rudiger, S. G. D. Dancing with the Diva: Hsp90-Client Interactions. J. Mol. Biol. 430, 3029–3040 (2018).

14. Rosenzweig, R., Nillegoda, N. B., Mayer, M. P. & Bukau, B. The Hsp70 chaperone network. Nat. Rev. Mol. Cell. Biol. 20, 665–680 (2019).

15. Li, J., Soroka, J. & Buchner, J. The Hsp90 chaperone machinery: conformational dynamics and regulation by co-chaperones. Biochem. Biophys. Acta 1823, 624–635 (2012).

16. Mayer, M. P. & Le Breton, L. Hsp90: breaking the symmetry. Mol. Cell 58, 8–20 (2015).

17. Kirschke, E., Goswami, D., Southworth, D., Griffin, P. R. & Agard, D. A. Glucocorticoid receptor function regulated by coordinated action of the proteasome and hsp90. Cell 157, 1685–1697 (2014).

18. Scheuff, C. et al. Structure of TPR domain-peptide complexes: critical elements in the assembly of the Hsp70-Hsp90 multichaperone machine. Cell 101, 199–210 (2000).

19. Schmidt, A. B. et al. The architecture of functional modules in the Hsp90 co-chaperone Sti1/Hop. EMBO J. 31, 1506–1517 (2012).

20. Bhatattacharya, K., Bernasconi, D. A. Glucose response energy transfer between genetically encoded donor and acceptor for protein-protein interaction studies in the molecular chaperone Hsp70/Hsp90 complexes. Sci. Rep. 8, 2801 (2018).
21. Hatherley, R., Clitheroe, C. L., Faya, N. & Tastan Bishop, Ö. Plasmodium falciparum: a detailed analysis on complex formation with Hsp70 and Hsp90. Biochim. Biophys. Acta 1856, 440–445 (2015).

22. Alvira, S. et al. Structural characterization of the substrate transfer mechanism in Hsp70/Hsp90 folding machinery mediated by Hop. Nat. Commun. 5, 5484 (2014).

23. Gallastegui, N. & Groll, M. The 26S proteasome: assembly and function of a degradative machine. Trends Biochem. Sci. 35, 634–642 (2010).

24. Murata, S., Yashiro-Tsukagoshi, H. & Tanaka, K. Molecular mechanisms of proteasome assembly. Nat. Rev. Mol. Cell. Biol. 10, 104–115 (2009).

25. Collins, G. A. & Goldberg, A. L. The Logic of the 26S Proteasome. Cell 169, 792–806 (2017).

26. Immay, J., Maruya, M., Yashiro-Tsukagoshi, H., Yahara, I. & Tanaka, K. The molecular chaperone Hsc70 plays a role in the assembly and maintenance of the 26S proteasome. EMBO J. 22, 3557–3567 (2003).

27. Lander, G. C. et al. Complete subunit architecture of the proteasome. Nature 442, 186–191 (2012).

28. Genest, O., Hoskins, J. R., Kravats, A. N., Doyle, S. M. & Wickner, S. Hsp70 and Hsp90 of E. coli directly interact for collaboration in protein remodeling. J. Mol. Biol. 427, 3877–3889 (2015).

29. Kravats, A. N. et al. Interaction of E. coli Hsp90 with DnaK involves the DnaK binding region of DnaK. J. Mol. Biol. 429, 858–872 (2017).

30. Sung, N. et al. Mitochondrial Hsp90 is a ligand-activated molecular chaperone coupling ATP binding to dimer closure through a coiled-coil intermediate. Proc. Natl Acad. Sci. USA 113, 2952–2957 (2016).

31. Sun, M., Kotler, J. L. M., Liu, S. & Street, T. O. The endoplasmic reticulum (ER) chaperones BiP and Grp94 selectively associate when BiP is in the ADP conformation. J. Biol. Chem. 279, 6387–6396 (2019).

32. Nicolet, C. M. & Craig, E. A. Isolation and characterization of STI1, a stress-inducible gene from Saccharomyces cerevisiae. Mol. Cell. Biol. 9, 3638–3646 (1989).

33. Song, H. O. et al. C elegans STI-1, the homolog of Sis1/Hop, is involved in aging and stress response. J. Mol. Biol. 390, 604–617 (2009).

34. Ambegaokar, S. S. & Jackson, G. R. Functional genomic screen and network analysis reveal novel modifiers of tauopathy dissociated from tau phosphorylation. Hum. Mol. Genet. 20, 4947–4977 (2011).

35. Beraldo, F. H. et al. Stress-inducible phosphoprotein 1 has unique phosphorylation. J. Biol. Chem. 282, 589 (2007).

36. Liu, X. D., Morano, K. A. & Thiele, D. J. The yeast Hsp110 family of proteasome regulators. Biochim. Biophys. Acta 1856, 440–445 (2015).

37. Guerrero, C., Milenkovic, T., Przulj, N., Kaiser, P. & Huang, L. Characterization of the proteasome interaction network using a QTAX-based tag-team strategy and protein interaction network analysis reveals novel modifiers of tauopathy dissociated from tau phosphorylation. Hum. Mol. Genet. 20, 4947–4977 (2011).

38. Dantuma, N. P., Lindsten, K., Gläs, R., Jelline, M. & Masuccio, M. G. Short-lived proteins as quality control sensors for ubiquitin/proteasome-dependent proteolysis in living cells. Nat. Biotechnol. 18, 538–543 (2000).

39. Choukta, E. et al. Hsp83 loss suppresses proteasomal activity resulting in an upregulation of caspase-dependent compensatory autophagy. Autophagy 13, 1573–1589 (2017).

40. Vanduri, P., Hao, R., Fitzpatrick, T. & Yao, T. P. Chaperone-mediated 26S proteasome remodeling facilitates free K3 ubiquitin chain production and aggresome clearance. J. Biol. Chem. 290, 9455–9464 (2015).

41. Yamano, T. et al. Hsp90-mediated assembly of the 26 S proteasome is involved in major histocompatibility complex class I antigen processing. J. Biol. Chem. 283, 20606–20608 (2008).

42. Sharma, K. et al. Quantitative proteomics reveals that Hsp90 inhibition preferentially targets kinases and the DNA damage response. Mol. Cell. Proteom. 11, M111 014654 (2012).

43. Fierro-Monti, I. et al. Dynamic impacts of the inhibition of the molecular chaperone Hsp90 on the T-cell proteome have implications for anti-cancer therapy. PLoS ONE 8, e60425 (2013).

44. Wala, J. et al. 26S proteasome structure revealed by three-dimensional electron microscopy. J. Struct. Biol. 121, 19–29 (1998).

45. Unno, M. et al. The structure of the mammalian 20S proteasome at 2.75 Å resolution. Structure 10, 609–618 (2002).

46. Tai, H. C., Besche, H., Goldberg, A. L. & Schuman, E. M. Characterization of the brain 26S proteasome and its interacting proteins. Front. Mol. Neurosci. 3, 12 (2010).

47. Tsvetkov, P. et al. Compromising the 19S proteasome complex protects cells from reduced flux through the proteasome. elife 4, e08467 (2015).

48. Morozov, A. V. & Karpov, V. L. Proteasomes and several aspects of their heterogeneity relevant to cancer. Front. Oncol. 9, 761 (2019).

49. Averbaugh, R. D. et al. Paradoxical resistance of multiple myeloma to proteasome inhibitors by decreased levels of 19S proteasomal subunits. elife 4, e08153 (2015).
77. Hamazaki, J. et al. Rpn10-mediated degradation of ubiquitinated proteins is essential for mouse development. *Mol. Cell. Biol.* **27**, 6629–6638 (2007).
78. Sakao, Y. et al. Mouse proteasomal ATPases Pmc3 and Pmc7: genomic organization and gene targeting. *Genomics* **67**, 1–7 (2000).
79. Szlanka, T. et al. Deletion of proteasomal subunit 55α/Rpn10/p54 causes lethality, multiple mitotic defects and overexpression of proteasomal genes in Drosophila melanogaster. *J. Cell Sci.* **116**, 1023–1033 (2003).
80. Brokhin, V., Gheyseelink, J., Gagliardini, V., Genschik, P. & Grossniklaus, U. The RPN1 subunit of the 26S proteasome in Arabidopsis is essential for embryogenesis. *Plant Cell* **17**, 2723–2737 (2005).
81. Brehme, M. et al. A chaperome subnetwork safeguards proteostasis in aging and neurodegenerative disease. *Cell Rep.* **9**, 1135–1150 (2014).
82. Lackie, R. E. et al. Modulation of hippocampal neuronal resilience during aging by the Hsp70/Hsp90 co-chaperone STI1. *J. Neurochem.* **153**, 727–758 (2020).
83. Chakraborty, A. & Edkins, A. L. Hop depletion reduces HSF1 levels and activity and coincides with reduced stress resilience. *Biochem. Biophys. Res. Commun.* **527**, 440–446 (2020).
84. Ruckova, E., Muller, P., Nenutil, R. & Vojtesek, B. Alterations of the Hsp70/Hsp90 chaperone and the HOP/CHIP co-chaperone system in cancer. *Cell. Mol. Biol. Lett.* **17**, 446–458 (2012).
85. Yi, F. & Regan, L. A novel class of small molecule inhibitors of Hsp90. *ACS Chem. Biol.* **3**, 645–654 (2008).
86. Pimenta, G., Herbert, K. M. & Regan, L. A compound that inhibits the HOP-Hsp90 complex formation and has unique killing effects in breast cancer cell lines. *Mol. Pharm.* **8**, 2252–2261 (2011).
87. Ran, F. A. et al. Genome engineering using the CRISPR-Cas9 system. *Nat. Protoc.* **8**, 2281–2308 (2013).
88. Hjerpe, R. et al. UBQLN2 mediates autophagy-independent protein aggregate clearance by the proteasome. *Cell* **166**, 935–949 (2016).
89. Besche, H. C. & Goldberg, A. L. Affinity purification of mammalian 26S proteasomes using an ubiquitin-like domain. *Methods Mol. Biol.* **832**, 423–432 (2012).
90. Roy, A., Kucukural, A. & Zhang, Y. I-TASSER: a uni-
91. Echeverria, P. C., Bhattacharya, K., Joshi, A., Wang, T. & Picard, D. The protein structure and function prediction. *J. Chem. Inf. Model.* **50**, 278–279 (2010).
92. Cox, J. et al. Andromeda: a peptide search engine integrated into the MaxQuant environment. *J. Proteom.* **14**, 2028–2037 (2019).
93. Cox, J. & Mann, M. MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. *Nat. Biotechnol.* **26**, 1367–1372 (2008).
94. Cox, J. et al. Andromeda: a peptide search engine integrated into the MaxQuant environment. *J. Proteome Res.* **10**, 1794–1805 (2011).
95. Cox, J. et al. Accurate proteome-wide label-free quantification by delayed normalization and maximal peptide ratio extraction, termed MaxLFQ. *Mol. Cell. Proteom.* **13**, 2513–2526 (2014).
96. Theyenot, E. A., Roux, A., Xu, Y., Ezan, E. & Junot, C. Analysis of the human adult urinary metabolome variations with age, body mass index, and gender by implementing a comprehensive workflow for univariate and OPLS statistical analyses. *J. Proteome Res.* **14**, 3322–3335 (2015).
97. Galindo-Prieto, B., Eriksson, L. & Trygg, J. Variable influence on projection (VIP) for orthogonal projections to latent structures (OPLS). *J. Chemom.* **28**, 623–632 (2014).
98. Scheres, S. H. RELION: implementation of a Bayesian approach to cryo-EM structure determination. *J. Struct. Biol.* **180**, 519–530 (2012).
99. Galindo-Prieto, B., Eriksson, L. & Trygg, J. Variable influence on projection (VIP) for orthogonal projections to latent structures (OPLS). *J. Chemom.* **28**, 623–632 (2014).
100. Scheres, S. H. RELION: implementation of a Bayesian approach to cryo-EM structure determination. *J. Struct. Biol.* **180**, 519–530 (2012).

**Acknowledgements**

We are grateful to Alfred L. Goldberg, Matthias P. Mayer, Ueli Schübler, Marcello Maggioni, Donald McDonnell, Yoshikico Miyata and Adrienne Edkins for gifts of plasmids. We thank David O. Toft, and Martine Collart for gifts of antibodies, Jason Gestwicki for the gift of JG-98, Irene Olivan Mura for contributing to the in vitro refolding experiments, Stacey Mattison for critically reading the Methods section of this manuscript, and previous members of the Picard laboratory for miscellaneous reagents. This work has been supported by the Swiss National Science Foundation and the Canton de Genève.

**Author contributions**

K.B. conceived the study, designed and performed experiments, analyzed the data, prepared figures, and wrote the manuscript. I.W. and M.Q. conducted the proteomic analyses. T.M.L., E.C.P., M.B.K., and S.G.D.R. designed and performed in vitro luciferase refolding assays. P.C.E. performed bioinformatics analyses of proteomic datasets. M.V. generated Hsp90α- and Hsp90β-KO cells. L.B. contributed to experiments with recombinant proteins. D.W. helped with all yeast experiments. Y.S. and C.B. contributed to TEM experiments and analyses. D.P. conceived the study, contributed to designing the experiments and analyzing the data, supervised the work, and wrote and edited the manuscript. All authors provided critical analysis of the data and contributed to the editing of the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

Supplementary information is available for this paper at https://doi.org/10.1038/s41467-019-19783-w.

Correspondence and requests for materials should be addressed to D.P.

Peer review information *Nature Communications* thanks Andrew Truman, Sue Wickner and other, anonymous, reviewers for their contributions to the peer review of this work. Peer review reports are available.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2020