Mechanisms tailoring the expression of heat shock proteins to proteostasis challenges

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All cells possess an internal stress response to cope with environmental and pathophysiological challenges. Upon stress, cells reprogram their molecular functions to activate a survival mechanism known as the heat shock response, which mediates the rapid induction of molecular chaperones such as the heat shock proteins (HSPs). This potent production overcomes the general suppression of gene expression and results in high levels of HSPs to subsequently refold or degrade misfolded proteins. Once the damage or stress is repaired or removed, cells terminate the production of HSPs and resume regular functions. Thus, fulfillment of the stress response requires swift and robust coordination between stress response activation and completion that is determined by the status of the cell. In recent years, single-cell fluorescence microscopy techniques have begun to be used in unravelling HSP-gene expression pathways, from DNA transcription to mRNA degradation. In this review, we will address the molecular mechanisms in different organisms and cell types that coordinate the expression of HSPs with signaling networks that act to reprogram gene transcription, mRNA translation, and decay and ensure protein quality control.

For organisms to grow and function properly, they must maintain specific internal cellular conditions that allow proteins to acquire their functional conformations and cells to achieve protein homeostasis (proteostasis) (1). Maintaining proteostasis becomes critical when facing abrupt changes in the external conditions, such as an increase in temperature, which can lead to protein misfolding and aggregation, and consequently, cellular dysfunction (2). Thus, organisms must sense, rapidly respond, and adapt to new environmental conditions for survival. Organisms from bacteria to mammals have evolved similar and varying stress responses to cope with protein misfolding and maintain proteostasis successfully. Some of these strategies include modulations of signaling cascades, changes in transcriptional programs, and regulation of translation, posttranslational modifications, and the dynamic assembly of RNA and protein condensates (ribonucleoprotein [RNP] granules) through liquid–liquid phase separation (1, 3–7). Several of these molecular mechanisms converge to sustain proteostasis in response to sudden and acute changes in environmental conditions.

Increases in the environmental temperature is a universal proteostasis challenge encountered by most organisms. For historical reasons, thermal stress has been used as a paradigm to study the stress response. Nowadays, these studies have an additional relevance due to the increased exposure of organisms to heatwaves derived from climate change (8–10). Increased thermal energy in the cells can result in heat-induced denaturation of proteins and thermally altered metabolic activity leading to an increase in reactive oxygen species, which can damage all biological macromolecules, including proteins (11). Cells cope with an increased load of unfolded and misfolded proteins by modulating the expression of specific molecular chaperones, also known as heat shock proteins (HSPs) (12–15). The heat shock response (HSR) refers to the activation of the expression of HSPs, and it is the most common and widely studied cell response to thermal stress. HSPs play a central role in the lifecycle of proteins because they promote the folding of nascent polypeptides into their native/functional configurations and prevent protein misfolding and aggregation (12, 15, 16). HSPs also collaborate with the quality control mechanisms, the ubiquitin–proteasome system, and autophagy, to target misfolded proteins and aggregates whose native functional state cannot be recovered for degradation (5, 17).

Given that HSPs are central to the cellular proteostasis network, cells undertake several gene expression adaptations to favor the synthesis of HSPs at the expense of decreasing most cellular functions (Fig. 1). Biochemical and molecular biology approaches highlight the unique regulation of HSP gene expression. The spatiotemporal resolution of such precise regulation is now being uncovered using high-resolution quantitative fluorescence microscopy. Gene expression adaptations during stress act together to protect macromolecules and promptly resume the cytoplasmic and nuclear activities once permissive conditions are restored (3). The regulation of HSP expression coordinates with other cell protective mechanisms, like the formation of RNP condensates and the activation of the integrated stress response (ISR) to repress translation initiation. The ISR and HSR also coordinate their actions with the unfolded protein response in the endoplasmic reticulum (ER) and the mitochondria to preserve proteostasis across cellular compartments. All organisms ranging from bacteria to...
Plants and mammals have genes encoding for HSPs. HSPs are grouped into families based on an apparent molecular weight (18, 19). The HSP70 and HSP90 families are the most functionally relevant HSPs in the cell (15, 20). They are ATP-dependent chaperones that cooperate with small HSPs and HSP110. Cochaperones of the J-domain family of proteins modulate HSP70 activity by accelerating ATP hydrolysis, participating in substrate recognition and substrate folding or refolding (Fig. 2)(21–27). HSPs are further categorized as constitutive or inducible based on their steady-state expression levels. The expression of all inducible and some constitutive HSPs is upregulated to some extent upon heat stress. Among them, the inducible HSP70 genes are the fastest and most upregulated (23, 24, 27). Interestingly, they are highly conserved among species having an amino acid similarity of 50% between *Homo sapiens* and *Escherichia coli*, while some domains are 96% similar, which highlights its vital role in cell adaption to changing environmental conditions (28).

In this review, we frame the molecular regulation of the HSR to the context of the gene expression changes undertaken by eukaryotic cells in response to an increase in temperature. We compare the response mounted by different organisms and cell types and suggest new technological approaches to overcome the gap in our knowledge on the HSPs expression.

**Transcriptional regulation of inducible HSPs versus constitutive genes**

**Transcriptional upregulation of stress-inducible HSPs**

The robust transcriptional induction of genes of the HSP70 family is one of the main and fastest response to heat stress. Their transcriptional induction occurs at the expense of a general transcriptional downregulation of constitutively expressed genes. Most inducible HSP70 genes are short (around 2500 nucleotides) and intronless, and their promoter contains one or more binding sites, known as heat shock elements (HSEs), for the association of the master transcription factor heat shock factor 1 (HSF1) (29). Under physiological conditions, the inducible HSP70 genes are not expressed. However, their loci are neither present in a compact heterochromatin domain nor marked by repressive epigenetic histone modification. The promoter and 3’ end of HSP70 gene is...
nucleosome-free while its gene body is covered by nucleosomes. The promoter is bound by a paused RNA polymerase II (RNAPII) (30). These characteristics prevent the stable transcriptional repression of HSP70 genes and facilitate their prompt activation in response to the binding of HSF1.

Under physiological conditions, HSF1 shuttles between the nucleus and cytoplasm, and it is kept as an inactive monomer by constitutive members of HSP90 and HSP70 families. Upon stress, HSF1 is released from HSPs, trimerizes, and localizes in the nucleus where it binds to the HSE, which is comprised of at least three nGAAn repeats organized head to tail in the promoters of genes encoding HSPs and other gene products (31, 32) (Fig. 3). HSF1 has three domains, an oligomerization domain next to the DNA binding domain at the N terminus, a trans-activation domain at the C terminus that induces transcription initiation and elongation, and a regulatory domain in the middle that negatively regulates the function of the trans-activation domain in nonstress conditions. By forming a trimer, the affinity of HSF1 for the HSE increases as each HSF1 of the trimer binds to a nGAAn repeat through its DNA binding domain. The binding of HSF1 to HSE is not sufficient to activate transcription and has to be accompanied by extensive posttranslational modifications. HSF1 undergoes hyperphosphorylation of serine and threonine residues that cover up to 90% of the regulatory domain (33–36). However, only a few of these phosphorylation sites, like serines 230 or 326, are necessary for the activity of HSF1 (35, 37). Concomitantly, sumo groups that have an inhibitory effect on transcription are removed from HSF1 (38). HSF1 acetylation at lysines 116 and 118 favors its transcriptional activity, whereas acetylation at several other lysine residues regulates its nuclear localization and oligomerization (31). Acetylation of HSF1 occurs a few hours after heat shock to decrease its DNA affinity and the transcriptional response (39). In summary, HSF1 undergoes extensive posttranslational modifications, which are regulated under various stresses. Although the function of some of these modifications has been identified, the role of many others, as well as the proteins responsible for their regulation, remains to be elucidated.

The combination of posttranslational modifications and titration of HSPs by misfolded proteins have been demonstrated to activate HSF1. Recent work in the yeast, Saccharomyces cerevisiae, has allowed building a simple mathematical model that points to the dissociation of HSP70/HSP90 from HSF1 as the first “switch on” step to activate HSP70 transcription, which feedback to HSF1 to switch it off or repress it (40, 41). Zheng et al. (41) identified 70 phosphorylation sites on HSF1 upon heat shock and were able to model that these
phosphorylations have no effect on HSF1 activation but instead increase its transcriptional activity by favoring its association with the mediator complex. Additionally, the translation factor eEF1A and the noncoding RNA HSR1 are among the factors activating HSF1. They act together to form a nucleoprotein complex with HSF1 and stimulate HSF1 trimerization (42). Following heat shock, HSF1 recruits multiple cofactors to HSE (43–46), including SGO2, which recruits the subunit mediator complex MED12, essential for the strong transcriptional induction of HSP genes (47). SGO2 binding to hypophosphorylated RNAPII targets it to the promoter of HSP genes by forming a complex with HSF1. Transcription is then induced by other transcription factors like P-TEFb, recruitment of which are mediated by HSF1 (48). P-TEFb is sufficient to induce the phosphorylation of the serine 2 in the C-terminal domain of RNAPII, which leads to transcription elongation (48, 49). However, a strong transcriptional induction requires the nucleosomes positioned along the HSP70 gene body to be removed. The chromatin remodelers SWI/SNF in mammals and FACT together with the histone chaperone Spt6 in Drosophila melanogaster (D. melanogaster) are recruited by HSF1 to the HSP70 genes within minutes after heat shock to remove the nucleosomes (31).

Besides the activation of HSF1, heat shock induction of HSP70 in mammalian cells depends on the relocation of the HSP70 loci from the nuclear membrane to speckles (50, 51). The rapid, active, and unidirectional movement of HSP70 loci is mediated by nuclear actin polymerization. The association of the HSP70 locus with speckles depends on the promoter sequence and determines the robust transcriptional activation of HSP70 upon heat shock stress. Although speckles contain serine 2–phosphorylated RNAPII and other components of the transcriptional machinery, the specific speckle factors critical for the transcriptional activation of HSP70 have not yet been identified. In yeast, genes encoding for different HSPs coalesce in discrete spots in the nucleus upon transcription stimulation. This interallelic clustering leads to the interaction between HSP104 and HSP12 loci and depends on the activation of their transcription. This result suggested the presence of specific transcriptional factories formed in response to heat stress, which could be coregulated by HSF1 (52).

**Transcriptional downregulation of nonstress genes**

The transcriptional induction of HSP genes during heat stress is accompanied by the upregulation of other non-HSP genes encoding for cytoskeleton and oxidative stress proteins and a massive downregulation of thousands of genes (For review: (53)). Detailed analysis of the position of the RNA polymerases, chromatin modifications, and domains in D. melanogaster and mammalian cells suggest that changes in the chromatin landscape cannot explain the rapid changes in transcriptional preferences upon heat shock (54, 55).

Heat shock does not induce a global chromatin remodeling nor modifications of topology associated domains in human or D. melanogaster S2 cells (56) or rearrangements of topology associated domain borders in D. melanogaster Kc167 cells (57). The addition of DTT to induce protein unfolding in...
**D. melanogaster** S2 cells did not trigger a global decrease in nucleosome occupancy over the induced genes, their promoters, or enhancers, as detected by micrococcal nuclease sequencing (54). However, the accessibility of the chromatin is increased at upregulated genes as measured with ATAC-seq. These results could be explained by nucleosomes undergoing an increased turnover due to their increased acetylation and ongoing transcription. Mueller et al. also reported a decrease in nucleosome occupancy and an increase in accessibility in a few genes, like the constitutively expressed chaperone HSC70. Similar observations were done in human K562 cells following 30 minutes of heat shock. The level of histone 3 lysine 27 acetylation (H3K27Ac) increased at the promoter of all transcribed genes, which also experienced an increase in RNAPII occupancy in the promoter and to a lesser extent in the gene body (55). Conversely, a decrease of polymerase occupancy is observed along the gene body of constitutively expressed genes whose transcription is downregulated upon stress. Hence, it is suggested that the extra available RNAPII quickly replaced the RNAPII undertaking transcription elongation in stress-regulated genes.

Additionally, noncoding RNAs have been shown to be a transcriptional repressor of non-HSP genes during heat shock. For instance, noncoding transcripts such as Alu RNA derived from short interspersed nuclear elements bind to RNAPII during heat shock to inhibit transcription of other mRNAs, such as actin (58–60). Interestingly, it has recently been shown that the long noncoding RNA, heat-enhanced antisense transcript (Heat) binds to HSF1 in vitro and in vivo via a trans-acting manner to attenuate the expression of stress genes. Experiments on mouse embryonic fibroblasts suggest that Heat uses HSF1 as a carrier by forming an RNP complex to target stress genes. While the exact mechanism by which Heat suppresses transcription is not known, Ji et al. (61) suggest the attenuation of the HSR by Heat involves an m6A modification and the nuclear m6A reader protein YTHDC1. Furthermore, the global downregulation of transcription induced by heat shock is suggested to be caused by activation of cryptic intronic polyadenylation sites in introns. Intronic polyadenylation sites led to premature transcriptional termination and new short mRNAs accumulating in the nucleus (62). Thirty percent of the stress-induced genes have HSF1 bound to the promoter. Several genes upregulated by heat stress but not bound by HSF1 are frequently contacted by distal regions bound by HSF1 as shown by chromosome conformation capture techniques, Hi-C performed in human and Drosophila cells (56). However, the regulatory mechanisms that coordinate the transcriptional induction of HSF1-dependent and -independent genes with the transcriptional repression of more than 6000 genes in humans remains to be uncovered. Overall, heat shock and other stresses dramatically affect global transcriptional regulation, leading to potent induction of genes encoding for prosurvival proteins.

### Translational regulation during the HSR

To preferentially synthesize HSPs during stressful events, cells have adapted a mechanism whereby non-HSP transcripts are retained in the nucleus, and HSP transcripts are selectively exported to and translated in the cytoplasm. The exact mechanism of this selective process is not known. However, work conducted on *S. cerevisiae* has shown that nuclear export of non-HSP mRNA involves RNA adaptor proteins, Npl3, Gbp2, Hrb1, and Nab2, cotranscriptionally loaded onto the pre-mRNA, which then recruits Mtx67–Mtr2 (TAP–p15 in humans), an essential heterodimeric receptor mRNA export factor (63). These adaptor proteins have an mRNA quality control function that prevents the nuclear export of incorrect, possibly improperly processed or assembled mRNAs (64). During stress, Mtx67 and the adaptor proteins are dissociated from non-HSP transcripts to prevent nuclear export. However, HSP transcripts do not require adaptive proteins and are loaded directly with Mtx67 via HSF1 (64). Thus, HSP transcripts bypass the adaptor–protein–mediated quality control mechanism to be rapidly exported and translated.

The newly synthesized HSP mRNAs encounter a cytoplasmic environment in which translation is repressed. Cells sense the load of misfolded proteins and repress translation by different pathways to decrease the load of unfolded and misfolded proteins. However, the translation of mRNA encoding for specific HSPs, such as the inducible HSP70, HSP82, and HSP27, is specifically favored (65). Regulation of translation enables the cells to rapidly adapt their proteome to stress conditions (66). From the three stages of translation: initiation, elongation, and termination, stress conditions repress cap-dependent translation initiation and elongation (67). The association of translation initiation factors (eIF4A, eIF4B, Ded1) and ribosomal proteins with mRNAs is decreased immediately upon stress (68). Additionally, stress conditions promote the recruitment of translation factors and regulatory RNA binding proteins to stress-induced cytoplasmic structures known as stress granules (SGs) and processing bodies (PBs), limiting their availability (69–71) (Fig. 1).

#### Regulation of translation initiation and elongation

Eukaryotic translation initiation is a highly regulated multistep process and the rate-limiting step in translation. It involves the assembly of the ternary complex (TC) and binding of the eIF4F complex to the 5′ m7G cap structure in the mRNA. Both steps are downregulated during stress by different signaling pathways (72).

The TC, GTP-eIF2α-initiator methionine tRNA, preloaded with eIF1, eIF1A, eIF3, and eIF5, assembles on the 40S ribosomal subunit to form the 43S preinitiation complex (73–75). Stress precludes the formation of the TC by the reversible phosphorylation of the Ser51 in eIF2α. This phosphorylation event limits the pool of available eIF2α for the TC formation. (Fig. 1). The phosphorylation of eIF2α in response to stress is part of the ISR (76). Depending on the stress stimulus, one of four serine-threonine kinase enzymes catalyzes the phosphorylation of Ser51. These kinases are the heme-regulated inhibitor and the general control nonderepressible 2, which are conserved in eukaryotes, the protein kinase R (PKR), which
is specific to vertebrates, and the PKR-like ER kinase, which is absent in fungi. Heme-regulated inhibitor is activated to cope with heme deficiency in red blood corpuscles, heat and osmotic shock, oxidative and mitochondrial stress, cytosolic protein aggregation, and arsenite treatment in cells other than red blood corpuscles (66, 77–82). General control non-repressible 2 phosphorylates eIF2α in conditions of nutrient depletion (83) and UV irradiation (84, 85). PKR mediates the phosphorylation of eIF2α in response to the detection of double-stranded viral RNA and hyperosmotic stress (dsRNA) (82, 86). PKR-like ER kinase activates upon ER stress, aggregation of misfolded proteins in the ER lumen, hypoxia, hypoglycemia, ischemia, oxidative stress, and perturbation in Ca²⁺ levels (87, 88). The conversion of the eIF2-GDP binary complex to the translation competent eIF2-GTP is mediated by the eIF2-specific guanine nucleotide exchange factor, eIF2B (80, 89). Two copies of eIF2B forms an active decameric complex that interacts with eIF2 and loads a molecule of GTP on eIF2. Phosphorylated eIF2α acts as a noncompetitive inhibitor by sterically hindering the access of eIF2 to the catalytic domain of eIF2B and sequestering it. Consequently, the recycling of eIF2α is decreased, which in turn decreases the abundance of TC (80, 90, 91) and impairs cap-dependent translation promoting metabolic dormancy to survive through the stress (92).

In addition to eIF2α phosphorylation, the binding of the eIF4F complex to the 5' m⁷G cap structure is impaired by different means. The eIF4F complex is made of the cap recognizing factor eIF4E, the scaffold protein eIF4G, and the ATP-driven RNA helicase eIF4A that unwinds secondary structures in the 5' untranslated region (UTR) of mRNAs (75, 93). The binding of eIF4E to the cap is partially regulated by the mammalian target of the rapamycin (mTOR) pathway (93–96). mTOR is a kinase that phosphorylates the downstream targets eIF4E binding protein 1, preventing its binding to eIF4E, which allows its binding to the cap under favorable conditions (93, 97). During stress, mTOR is inactivated by TSC1/2, leading to the dephosphorylation of eIF4E binding protein 1, which readily sequesters eIF4E and suppresses the eIF4F complex formation (98–101). Additionally, the newly synthesized HSP27 binds eIF4G with high affinity preventing the formation of the eIF4F complex (102). Together, these mechanisms prevent the assembly and binding of eIF4F to the m⁷G cap structure and cap-dependent translation.

The elongation step of translation is also regulated to enforce the translation repression of nonstress mRNAs during conditions that challenge protein homeostasis. During translation elongation, the GTP-bound elongation factor eEF1A brings the aminoacyl-tRNA corresponding to the codon in the ribosomal A site, and eEF1A-GDP is released upon codon–anticodon base pairing. The ribosomal RNA in the peptidyl transfer center catalyzes the peptide bond formation, and the translocase eEF2–GTP triggers the mRNA–tRNA movement with the expense of GTP, and eEF2-GDP is released (103–105). Stress regulates translation elongation by a major downstream effector of mTORC1, the S6 kinase. The S6 kinase phosphorylates Ser366 of the eEF2 kinase and inactivates it (106). Inhibition of mTORC1 upon stress leads to the activation of eEF2 kinase that phosphorylates eEF2 at Thr56 of the GTP binding domain, pausing translation elongation (107–113). It should be noted that the regulation of eEF2 and its kinase is more complex than what we have described here, as they can be phosphorylated at different residues through different pathways (111, 114).

Besides eEF2 regulation, studies from two independent laboratories have identified chaperone-mediated regulation of translation elongation during stress. The cytoplasmic chaperone HSP70 and HSP90 interact with nascent polypeptides, favoring their cotranslational folding as they emerge out the ribosome exit tunnel (115, 116) (Fig. 1). During severe stress, the prevalence of unfolded proteins titrates out the chaperones leaving the nascent polypeptides unassisted for cotranslational folding. Hence, an arrest in elongation was observed after the synthesis of the first 65 amino acids, which corresponds to the length of the nascent polypeptide that fits in the ribosome exit tunnel. Consequently, stalled ribosomes were observed in several different mRNAs at nucleotide position 195. The lack of chaperone–ribosome interaction impairs the cotranslational folding of nascent polypeptides accounting for the elongation pausing and global translation repression during severe heat stress (117, 118). Overall, HSP encoding mRNAs should be equipped to overcome the several steps at which translation is shut by the cell.

**SGs and PB formation**

Translation repression is accompanied by changes in the physical properties of the cytoplasmic milieu. Stress triggers RNAs and proteins to phase separate and form SGs and PBs that contain untranslated, long, and highly unstructured RNAs and proteins to phase separate and form SGs and PBs 

Translation repression is accompanied by changes in the physical properties of the cytoplasmic milieu. Stress triggers RNAs and proteins to phase separate and form SGs and PBs that contain untranslated, long, and highly unstructured RNAs and proteins that participate in translation, transcription, splicing, and decay (71, 119–121). SGs and PBs are dynamic membraneless structures assembled by liquid–liquid phase transition that favor stress tolerance and promote cellular fitness (122–126). While the number and size of PBs increase upon stress, SGs are formed under stress conditions that invoke arrest in translation initiation (127). SGs and PBs have different mRNA and protein compositions. PBs are made of RNA processing factors (e.g., eIF4E, DDX6, and Ded1p), decapping and deadenylation enzymes, exoribonucleases, and factors mediating mRNA stability (For review: 124, 125). Based on their enrichment in decay-related factors, PBs were believed to be mRNA degradation sites. However, mRNAs retained in PBs can return to the cytoplasm and engage in translation (124, 128). Thus, it is widely accepted that PBs could serve as a reservoir of nontranslating mRNAs and inactive decay enzymes (129, 130).

SGs form under stress conditions that induce eIF2α phosphorylation (131, 132), but they can also form independently of eIF2α phosphorylation. For example, puromycin treatment and inhibition of eIF4A with hippuristanol stimulate SG assembly (133–137). SGs are enriched in translation initiation factors, mRNA binding proteins, 40S ribosomal subunit, and mRNAs encoding for house-keeping genes (131, 132, 138, 139). mRNAs recruited to SGs are translationally
repressed and undergo compaction as the elongating ribosomes are released from the transcripts (140, 141). The SGs were speculated to function in blocking protein synthesis by trapping several initiation factors and stabilizing translationally inactive constitutive mRNAs so that they can reengage in translation upon recovery (136, 142–144). Certain mRNAs are shown to have paused ribosomes at the start codon to quickly re-initiate the protein synthesis as the cells recover from stress (145). Even so, the lack of SGs does not change mRNA translational repression or stability (142). Additionally, mRNAs-encoding stress-regulated proteins like activating transcription factor 4 have been shown to translate inside the SGs (146). HSP mRNAs skip the localization in SGs and PBs as their ongoing translation prevents them from condensing (119, 147).

**Translation of HSP70 mRNAs under stress conditions**

Several inducible HSP mRNAs are highly translated in spite of the general repression of translation during heat stress (148). Among them, the inducible HSP70 is the most synthesized HSP, as shown in 3H-leucine pulse labeling experiments (65). Hence, the HSP70 mRNA should have specific features favoring its translation. In 1985, Klemenz et al. (149) were the first to indicate that the 5′UTR of the HSP70 mRNA is required for its preferential translation during heat stress. McGarry and Lindquist also reported that the 5′ leader sequence of the HSP70 mRNAs is the region detected by *D. melanogaster* cells to preferentially translate it during heat stress (150). Given that both deletions and insertions within the 5′ UTR rendered HSP70 mRNA untranslated during heat stress and actively translated during recovery from stress, the authors suggested the presence of a secondary structure in the 5′ UTR (150). However, it was not until 1988 that the internal ribosome entry sites (IRESs) were discovered to mediate the translation of picornaviral mRNAs (151, 152), and until 1991 that the first cellular IRES was identified in the mRNA encoding for the immunoglobulin heavy chain binding-protein BiP or GRP78 (153). GRP78 is indeed an HSP70 that localizes and functions in the ER. Unlike the picornavirus mRNAs, the GRP78 mRNAs are capped, and their translation is favored when cap-dependent translation is halted under conditions of stress (149, 154, 155). Studies in HeLa cells also indicated that the factors eIF4E and eIF4G are dispensable for translation of HSP70 mRNA, suggesting a cap-independent mechanism of translation initiation (156). Following these discoveries, several labs have attempted to characterize the IRES structure and trans-regulatory factors required to translate HSP70 mRNA in different organisms (157–159). However, those attempts were unsuccessful in finding an IRES in the 5′ UTR of HSP70 mRNA and provided early evidence of an HSP70 mRNA cap-independent translation mechanism (160). Overall, these studies indicated that the mammalian HSP70 mRNA translation has reduced dependence on the eIF4F cap-binding complex during stress, HSP70 mRNA is translated in a cap-independent manner upon mTOR inhibitions, and the 5′ UTR is required for HSP70 mRNA translation during heat stress (157, 161) (Fig. 4).

Two different translational initiation control mechanisms have been suggested for the mammalian HSP70 mRNA: ribosome shunting (157) and recruitment of eIF3 by...
N6-methyladenosine (m6A) modification (161, 162). During ribosome shunting, the 40S ribosomal subunit skips a large portion of the HSP70 5' UTR and shunts to a region proximal to the canonical start codon when the cap-dependent translation is inhibited during heat shock (157, 159). More recently, it was found that HSP70 mRNAs are cotranscriptionally imprinted at adenosine 103 of their 5'UTR by methylation, m6A (161, 162). This methylation supports translation initiation by binding to the initiation complex eIF3, which recruits the ribosome to the mRNA. A follow-up paper concluded that cells activate m6A-mediated translation through the factor ABCF1. ABCF1 serves as an alternative recruiter for the TC to HSP70 mRNA during noncanonical translation upon heat shock (163). Further, heat-stress-mediated O-GlcNAcylation of eIF4GI has been reported essential for the translation of HSP70 mRNAs (164). Additionally, the escape of HSP70 mRNA from the shutoff of global protein synthesis was explained by the existence of specialized stress ribosomes. These ribosomes bear the cytoplasmic version of a mitochondrial protein, MRPL18, synthesized upon stress and might facilitate the recruitment of factors involved in translation elongation (165). The main caveat of these publications is the timing at which translation of HSP70 mRNA was studied. They used 4 h of recovery following 1 h of heat stress. At this time point, cap-dependent translation resumes, and SGs are resolved. Hence, they provide solid evidence on the regulation of HSP70 mRNA translated during recovery. Whether the same factors participate in its translation during stress remains to be elucidated (Fig. 4).

A relevant outcome of this research is the suggestion of cotranscriptionally imprinting of the mRNA that provides an advantage for its translation in the cytoplasm (163, 165). Paradigm shifting studies in yeast support the role of the HSE sequence in the promoter of HSP genes in determining the translation of inducible HSP mRNAs in the cytoplasm of glucose-starved yeast (166). We have previously reported that eIF1A1 links HSP70 transcription to translation in mammals, implicating the evolutionary conservation of this “remote control” mechanism of translational regulation (167). Hence, newly synthesized HSP mRNAs might arrive at the cytoplasm equipped for translation. These factors, together with the intrinsic characteristics of the 5'UTR, allow HSP mRNAs to engage in translation at the expense of housekeeping mRNAs. For example, the Ded1 helicase is recruited to condensates upon heat stress, precluding the translation of housekeeping mRNAs with secondary structures and favoring the translation of HSP mRNAs that have little structure in yeast (168). However, the 5'UTR sequence varies among HSP70 inducible genes and species. While D. melanogaster contains mostly AU-rich (70%) sequence and no secondary structure formation, like in yeast (158), the mammalian sequence has higher GC content (63%) which is likely to favor a formation of stable secondary structures. Even though we have not yet put together all the regulatory elements and the cascade of events that lead to the preferential synthesis of the inducible HSP70 during stress, they might differ among species.

**Recovery from stress and the degradation of HSP70 mRNA**

The newly synthesized HSP70 allows cells to resume their normal functions gradually by recovering proteostasis through the folding of misfolded proteins, which also prevents their aggregation, and by participating in the disassembly of SGs (169, 170). Accordingly, the synthesis of a nonfunctional HSP70 or inhibition of HSP70 mRNA translation during stress delays the recovery of global translation in D. melanogaster cells (148, 171–173). However, the synthesis of HSP70 should be repressed when cells return to optimal conditions. The persistent expression of the inducible HSP70 under nonstress conditions causes growth defects in D. melanogaster (174) and promotes malignancy in mammalian cells (175). Hence, the HSP transcription repression is accompanied by the rapid degradation of their mRNAs during recovery (171, 176). HSP70 mRNAs are the first to undergo translational repression and degradation. The timing and rate of its degradation during recovery depends on the severity and duration of heat treatment (176, 177). When cells are returned to optimal conditions following heat shock, they restore global protein synthesis, and constitutive mRNAs engage in translation (176, 177).

In mammalian cells, HSP70 mRNA transitions from being a stable mRNA during heat stress to a short-lived mRNA with a half-life of around 50 min during recovery from heat shock (172, 178–180). While the instability of HSP70 mRNA is also characteristic of nonstress cells, other stresses like incubation of cells with sodium arsenite or inhibition of protein synthesis initiation with pactamycin also stabilize HSP70 mRNA. Therefore, it has been suggested that a trans-regulator synthesized by the cells under nonstress or recovery conditions could destabilize HSP70 mRNA by binding to its 3' UTR. Deletion of the HSP70 mRNA 3'UTR or substitution with that of the alcohol dehydrogenase (adh) 3'UTR sequence stabilizes the transcript during recovery from stress (173, 177, 181). In mammalian cells, PKR directly or indirectly, for example through AUFI protein, associates to the 3'UTR of HSP70 mRNA through its AU-rich elements (AUUUA) that destabilize the transcript during recovery (182). However, AU-rich elements become dispensable for the fast deadenylation of HSP70 transcripts during recovery in D. melanogaster cells, which suggests that RNP s mediating HSP70 mRNA stability might differ among species (183). Like most cellular mRNAs, most HSP70 transcripts undergo a fast deadenylation mediated by the CCR4-NOT complex followed by decapping and degradation by the exonuclease Xrn1 in the 5'-to-3' direction, as described in D. melanogaster. HSP70 mRNA fragments shorter than the full-length mRNAs were identified by northern blot suggesting that some molecules might undergo degradation after an endonucleolytic cleavage by the exosome in the 3'-to-5' direction (183) (Fig. 5).
How does the cell recognize the HSP70 transcripts as the ones to be degraded? Within minutes of heat shock, hundreds of these mRNAs are synthesized, and when the conditions become optimal, they are selectively degraded with high efficiency. Is there a link between the translational status of the cell and HSP mRNA turnover? Polysome fractionation to study the translational profile of HSP70 mRNA in D. melanogaster and mammalian cells show a fraction of transcripts retained in higher polysomes while a subset of them is translationally inactivated (177). Are both populations degraded by the exact decay mechanism? It might also be possible that RNPs associated with HSP70 mRNA undergo posttranslational modifications, like arginylation, mediated by ATE-1, to regulate the stability of HSP transcripts, providing protection to cells upon heat shock (184).

Proteostasis on specific organisms, cell types, and conditions

While the HSR is a universal survival response to changes in the environment, there are variations in the regulation of the HSR among species and even within cell types of the same organisms. Additionally, the adaptation to abrupt or long-term changes influence the cellular response differently (8, 16, 181–183).

Variation of the HSR among organisms

The number of genes encoding for HSPs and cochaperones has increased over evolution, probably reflecting the increased number of proteins and complexity of functions undertaken by more evolved cells (185, 186). Multicellular organisms have not only expanded on the HSPs encoded in the genome of unicellular organisms but also adapted them to their proteostasis needs. For example, the mammalian genome lacks the disaggregase HSP104 present in S. cerevisiae (in eubacteria ClpB), which reverses stress-induced protein aggregation (187–189) and has been suggested to serve as a potential therapeutic agent to disaggregate toxic misfolded aggregates characteristic of neurodegenerative diseases (NDs) (187). While it is unknown why mammalians lack HSP104, in vitro data have shown that protein disaggregation in humans (and other metazoans such as nematodes) relies on the molecular machinery comprised of HSP70, HSP110, and J-proteins (188, 190, 191). These J-proteins are critical in driving HSP70–HSP110–based disaggregase by concomitantly interacting with both substrates and HSP70 partner proteins via single or mixed cooperating J-protein cochaperones of class A and B which relocate to protein aggregates following heat shock promote specific or broad-range aggregate targeting (188, 190–192). Overall, the core chaperones of the HSP70 and HSP90 families are similar among species and their organization as constitutive or inducible HSPs depends on their basal expression level (Table 1).

There is a variation in the temperature threshold needed to activate the HSF1 and the HSR across multicellular and unicellular eukaryotic species (193, 194) (Table 1). For example, while D. melanogaster induces the HSR at 30 °C, humans do so at 40 °C. This variability in temperature threshold has been suggested to be based on the environmental temperature of the organism and its capacity to maintain a constant body temperature (195, 196). Indeed, organisms occupying moderate variable thermal environments can modify the constitutive levels of HSPs and adjust their HSR to a higher onset temperature. In contrast, those organisms from a stable or highly...
Table 1
Comparison of HSP genes and ranges of temperatures across different organisms

| Yeast                  | C. elegans | Drosophila | Mammals | E. coli |
|------------------------|------------|------------|---------|---------|
| **Temperature range of HSR** |            |            |         |         |
| 29–37°C (238)          |            |            | (238)   |         |
| DT: 7°C (238)          |            |            |         |         |
| **Number of HSP genes** |            |            | (>142 (18) |         |
| 63 (238)               |            |            | (239)   | (239)  |
| **Transcription of HSPs** |            |            |         |         |
| Initiation (242)       |            |            |         |         |
| Peak (241)             |            |            | (239)   | (239)  |
| Initiation (239)       |            |            | (245)   | (246)  |
| Peak (248)             |            |            | (248)   |         |
| 1–5 min at 37°C        |            |            | 1–2 min at 37°C | 60 min at 35°C |
| Hsp70 and Hsp80:       |            |            |         |         |
| 60 min at 35°C         |            |            |         |         |
| Hsp70 and Hsp80:       |            |            |         |         |
| 2 h at 35°C            |            |            |         |         |
| Hsp70:                  |            |            |         |         |
| 4 min at 37°C          |            |            |         |         |
| 1 – 2 min at 37°C      |            |            |         |         |
| Hsp14:                  |            |            |         |         |
| ~25–12 min at 42°C    |            |            |         |         |
| Hsp16:                  |            |            |         |         |
| 1 h at 43°C            |            |            |         |         |
| Hsp81:                  |            |            |         |         |
| 30 min at 42°C         |            |            |         |         |
| Hsp16:                  |            |            |         |         |
| 33°C at 42°C           |            |            |         |         |
| Hsp18:                  |            |            |         |         |
| 33°C at 35°C           |            |            |         |         |
| Hsp19:                  |            |            |         |         |
| 5 min at 37°C          |            |            |         |         |
| Hsp70 (DnaK):          |            |            |         |         |
| 1–4 min at 42°C        |            |            |         |         |
| Hsp70:                  |            |            |         |         |
| ~15 min at 42°C        |            |            |         |         |
| Hsp82:                  |            |            |         |         |
| 1 h at 43°C            |            |            |         |         |
| Hsp70 (DnaK):          |            |            |         |         |
| 5–10 min at 42°C       |            |            |         |         |
| Hsp60 (GroEL) and Hsp70 (DnaK): | | | | |
| 3–5 min at 42°C        |            |            |         |         |

HSP, heat shock protein.
variable thermal environment are not as capable of readjusting the levels of their HSPs or onset temperature as they are already meeting their maximum thermal limit.

An interesting example is multicellular organisms lacking the induction of an HSR, like the Antarctic marine invertebrate ciliates and the Antarctic fish of the suborder Notothenioidei (8, 197). The absence of the HSR is suggested to be due to these animals living in a highly stabilized cold environment; thus, they evolved to adapt to this nontransient environment. However, this is the opposite of the Notothenioidei cold-temperate relatives in New Zealand, which have been shown to induce an HSR following heat stress. Another example of an animal with no HSR is the freshwater cnidian species Hydra oligactis, which is also highly sensitive to minor thermal variations. Its congener, Hydra vulgaris, can tolerate greater thermal ranges because it induces thermal tolerance following induction of HSPs synthesis (8, 197). Despite the lack of expression of inducible HSPs, these organisms express constitutive HSPs. They might have adapted the expression of their constitutive HSPs to help them overcome challenges faced in their environmental niche (8, 198). Thus, constitutive HSPs might play an essential role in overcoming proteostasis challenges without an inducible HSR. Examining the potential coordinated network of constitutive and inducible HSPs in promoting a competent HSR would provide a better understanding of the network of HSPs participating in the adaptation to changes in temperature.

Multicellular endothermal organisms tolerate higher ranges in temperature than unicellular or stenothermal multicellular organisms. However, endothermal organisms required a lower increase over their body temperature to induce the HSR (Table 1). There are unicellular organisms from the Archaeal species that can tolerate extreme temperatures such as those between 50 °C to 70 °C (thermophiles) and 80 °C or higher (hyperthermophiles) before the HSR is induced (199). In addition to the HSR, bacteria rely on a structural liability in response to temperature changes which create biological temperature sensors, such as DNA- and RNA-based environmental temperature sensors. Temperature changes will alter gene expression at transcriptional and posttranscriptional steps via DNA and RNA thermosensors to maintain proteostasis. Furthermore, changes in the secondary or tertiary structure can also be used by “RNA thermometers” in bacteria to regulate the translation efficiency of heat shock mRNAs. In this RNA ‘zipper-like’ thermosensory mechanism, the mRNA will adopt a thermolabile stem-loop structure in the 5’UTR which will either close to block translation at low temperatures or open at high temperatures to favor ribosome binding and translation (3, 10). These RNA thermosensing mechanisms are also important in the translation control of some HSPs in eukaryotic cells when cap-dependent translation initiation is inhibited (3, 10). For example, HSP90 mRNA in D. melanogaster becomes actively translated in response to heat shock but is inefficient in normal growth temperatures. It has been suggested that the HSP90 mRNA translation is substantially activated by heat shock due to the presence of a long stem in the AUG initiation proximal half of the 5’UTR, which serves as a heat-sensitive inhibitory element in the UTR that impedes access to the initiation codon (200). However, in heat stress, the stem undergoes a thermal destabilization, which allows the ribosomal subunits to recognize the region (200). While the expression of HSP90 has been shown to be essential for restoring folding yield when HSP70 levels are high, excess HSP90 (as well as HSP70) produced during heat shock may be detrimental to folding (201). Thus, the presence of a thermosensor is critical in mediating the translation of HSPs that are important for protein folding during heat shock. This preferential heat shock translation occurs in a similar translation mechanism as in bacteria wherein the start codon (AUG) will respond to differences in temperature similarly to bacterial RNA thermosensors. A similar mechanism of translation control has been proposed for HSP70 mRNA in human cells but has yet to be examined (3). These examples suggest that multicellular organisms conserved some of the modes of adaptation to heat stress used by bacteria, but they have changed the regulatory factors sustaining them.

Additionally, endothermal organisms readjust their physiology to increasing temperatures (197). The repeated exposure to elevations in core temperature due to natural environmental heat stress causes heat acclimatization (202). An increase in temperature is associated with a change in the pattern of HSPs expressed due to proteins being able to adapt their structures to varying temperatures, e.g., altering hydrophobicity, charge, noncovalent interactions, volume, and cooperativity. Indeed, patterns of adaptive variation in the structural and functional properties of proteins from organisms that have adapted to different temperatures have been reported (203, 204). A similar phenomenon has been recently described in the yeast S. Cerevisiae in response to a long-term temperature shift (205). Interestingly, a key feature of long-term temperature adaptation is the disappearance of protein aggregates, whereas acute heat shock induces protein aggregates. Thus, yeast may have adapted to persistent high temperatures by reducing the load of thermolabile proteins and relocating some proteins to minimize protein misfolding/unfolding at high temperatures (205). These findings have important implications in the context of global climate change as the current temperature changes we are experiencing can be considered as long-term temperature adaptation to the constant heat.

**Variation of the HSR among cell types**

The requirements of cellular proteostasis also vary across cell types in multicellular organisms, which maintain a relatively stable internal environment to sustain proteostasis among specialized tissues and organs (1, 2, 206). These multicellular organisms have highly specialized functions performed by distinct proteomes, in which proteostasis, assisted by a network of core and cell-specific chaperones, becomes challenging (186, 206). There are two hypotheses as to how multicellular organisms maintain proteostasis following an environmental perturbation like temperature. The first is that molecular chaperones are expressed in all cell types to guide folding and prevent misfolding, and thus, they
can buffer unexpected folding challenges. This hypothesis requires all cells to invest energy to have a reserve of chaperones for emergencies (1, 197, 205–208). As described in this review, the constitutive HSPs provide immediate assistance in coping with misfolded proteins upon acute stress (197, 206). Whereas de novo synthesized HSPs favor recovery from stress and fit the cell to overcome subsequent and more detrimental stress stimuli. This phenomenon, known as stress preconditioning, is used for medical purposes and suggests that cells have a certain buffer of HSPs to handle mild changes in environmental conditions (205, 207, 209–212).

The second hypothesis states that cells do not store excess chaperones. Instead, the cellular concentration of the chaperones is regulated precisely according to the immediate cellular requirements. Hence, the folding environment in the cell is delicate, with little capacity for a flux of non-native species. This hypothesis requires the HSR to be rapidly tailored to the proteostatic demands of the cell (206). Additionally, cell types and tissues would need to exchange information on the status of internal cellular proteostasis to coordinate proteostasis at the organismal level (2). The second hypothesis is demonstrated in Caenorhabditis elegans, wherein the thermosensory amphid neurons with finger-like ciliated endings (AFDs) detect changes in the ambient temperatures and coordinate the response. This coordinated response involves communication between the different tissues of C. elegans and is regulated by neurons. More specifically, AFDs sense heat shock stress in the environment; these stress signals are then sent via neuroendocrine fashion to tissues such as muscle and intestinal cells to regulate their HSR by activating HSF1 and promoting the induction of HSP70 (C12C8) (2, 213).

At the same time, muscle and intestine cells have a transcellular chaperone signaling between nonneuronal tissues that sense local proteotoxic stress and enhance chaperone signaling at a distance by signaling back to the neurons. These AFD neurons along with their postsynaptic cells, AIY interneurons, further regulate the temperature-dependent behavior of these organisms, such as growth and reproduction (206, 213). Similar activation of HSF1 through neuroendocrine signaling from the hypothalamic–pituitary–adrenal axis operates in rats (214).

As neurons play an important role in this organismal-level coordination in eukaryotes, it is important to mention the current literature on neuronal proteostasis. Neurons are highly polarized cells that have the capacity to tune their proteome locally, at axons, dendrites, and synapses, through the regulation of local protein synthesis, degradation, and post-translational modifications (215). However, it is not fully understood how proteostasis is sustained in different neuronal subcompartments under heat stress conditions. Rodent hippocampal and motor neurons exhibit a lower HSR activation than nonneuronal cells (206, 216). This impaired HSR can make neurons vulnerable to the toxic accumulation of misfolded proteins that underlie age-related NDs (216–219). Hence, a potential therapeutic strategy for NDs is to promote the activation of the HSR in neurons (220–223). The overexpression of HSP70 in a mouse model of Alzheimer’s Disease exerted cytoprotective roles and ameliorated physiological and behavioral deficits (220–223). Promoting the activation of HSF1 has long been considered a promising treatment for NDs. However, the threshold to activate HSF1 in motor neurons is higher than in somatic nonneuronal cells (216). The chromatin environment of HSP genes in neurons does not favor the binding of HSF1, and the treatment with histone deacetylases can enhance the transcriptional activity of HSF1 in motor neurons undergoing specific stresses (224). Hence, neurons might have been wired differently to handle proteostasis challenges and rely on other quality control mechanisms, like ubiquitin-proteasome system and autophagy, to sustain a healthy and functional proteome (218, 225–229).

**Conclusion and perspectives**

The induction of HSPs represents the first line of defense toward an increase in protein unfolding (230). To date, the regulation of HSPs expression has been mostly studied by ensemble measurements in cultured cell lines and yeast. This extensive research has provided detailed information on the kinetics of HSPs transcription and mRNA translation and degradation and has identified regulatory factors involved in each step of the life cycle of HSP mRNAs. In the last decade, single-cell microscopy approaches have granted the spatial resolution needed to investigate changes in the localization of HSP loci upon stress (50) as well as the subcellular localization of newly synthesized mRNAs (166, 167). These latest studies indicate that the fate of HSP mRNAs is decided cotranscriptionally. Therefore, a finely tuned communication between the nucleus and the cytoplasm under stress conditions could enable cells to identify HSP mRNAs as the ones to be translated. These results also suggest that each step in the life cycle of HSP mRNAs highly influences the next one. Given that hundreds of HSP mRNAs molecules are rapidly synthesized, translated, and degraded, the use of single-molecule fluorescence microscopy techniques to detect single mRNAs and de novo protein synthesis will mind the gap of our knowledge in the impact of HSP mRNAs translation on decay and the localization of mRNA degradation (231–233).

Activation of the HSR requires a mechanism to sense the damage and gene expression reprogramming to prioritize the expression of HSPs. The transcription factor HSF1 directs the upregulation of HSPs transcription. Activation of HSF1 occurs in all eukaryotes under a wide range of stresses (31, 230, 234). Together with HSF1 activation, cells attempt to minimize protein unfolding by blocking general translation elongation (230, 235). It is possible that besides the increased load of unfolding proteins, this general ribosome stalling signals to HSF1 by a still unknown mechanism (166). Identifying the mechanisms that act to orchestrate a competent HSR will provide the means to interrogate neurons in their stunted HSR and relate it with the neuronal vulnerability to accumulate misfolded proteins. In this case, using pathophysiological conditions relevant to the neuronal activity will provide a better understanding of the neuronal response to proteostasis challenges. Expanding the pioneering research done in
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Abbreviations—The abbreviations used are: AFD, amphid neurons with finger-like ciliated ending; ER, endoplasmic reticulum; HSE, heat shock element; HSF1, heat shock factor 1; HSP, heat shock protein; HSR, heat shock response; IRESs, internal ribosome entry sites; ISR, integrated stress response; mTOR, mammalian target of the rapamycin; ND, neurodegenerative disease; PBs, processing protein; PKR, protein kinase R; RNAPII, RNA polymerase II; RNP, ribonucleoprotein; SGs, stress granules; TC, ternary complex; UTR, untranslated region.

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