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The Role of Endoplasmic Reticulum Stress in Human Pathology

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Abstract

Numerous genetic and environmental insults impede the ability of cells to properly fold and posttranslationally modify secretory and transmembrane proteins in the endoplasmic reticulum (ER), leading to a buildup of misfolded proteins in this organelle—a condition called ER stress. ER-stressed cells must rapidly restore protein-folding capacity to match protein-folding demand if they are to survive. In the presence of high levels of misfolded proteins in the ER, an intracellular signaling pathway called the unfolded protein response (UPR) induces a set of transcriptional and translational events that restore ER homeostasis. However, if ER stress persists chronically at high levels, a terminal UPR program ensures that cells commit to self-destruction. Chronic ER stress and defects in UPR signaling are emerging as key contributors to a growing list of human diseases, including diabetes, neurodegeneration, and cancer. Hence, there is much interest in targeting components of the UPR as a therapeutic strategy to combat these ER stress–associated pathologies.

Keywords

protein misfolding; unfolded protein response; apoptosis; neurodegeneration; cancer; diabetes

THE ENDOPLASMIC RETICULUM: A PROTEIN-FOLDING FACTORY

One of the largest organelles in eukaryotic cells, the endoplasmic reticulum (ER) is a network of branching tubules and flattened sacs interconnected through an enclosed space called the ER lumen, which is separated from the surrounding cytosol by a single intracellular lipid bilayer, the ER membrane. Thus, the ER membrane is a boundary between the cytosol and the ER lumen and governs the passage of molecules between these two compartments (1).

The ER plays a major role in the synthesis, folding, and structural maturation of more than a third of all proteins made in the cell (2). In particular, nearly all proteins destined for residence in the ER, plasma membrane, Golgi apparatus, and lysosomes are translated on ER membrane-bound ribosomes and injected into the ER lumen. Likewise, most proteins that are ultimately secreted from the cell begin their journey in the ER. Proteins targeted to the ER have an N-terminal signal sequence that directs them to the ER membrane while they are still being synthesized on ribosomes. These proteins are cotranslationally translocated through the translocon complex, whereupon the signal sequence is removed by a protease as translation of the polypeptide is completed. Once in the ER lumen, proteins must be folded into their unique three-dimensional shapes and undergo various posttranslational modifications, including glycosylation and disulfide bond formation. These processes are catalyzed by abundant ER-resident enzymes such as chaperones, glycosylating enzymes, and oxidoreductases (3, 4). Furthermore, the ionic and electronic milieu of the ER is ideally suited for these protein-folding activities. Compared with the cytosol, the ER maintains a much higher calcium concentration and a more oxidizing redox potential (5, 6). Cells expend large amounts of energy to maintain the unique environment and functions of the ER. Chaperones bind to and help client proteins negotiate and avoid energetically unfavorable twists and turns on the way to their final native conformation (7). In many cases, protein folding also involves the covalent addition and trimming of sugars onto proteins through the process of glycosylation. Together, these enzymatic processes ensure that secretory proteins are properly folded, modified, and assembled into multiprotein complexes in the ER before they traffic further downstream in the secretory pathway.

Despite the effort of these protein-folding machines, the success rate for proper folding is quite low (under 20%) for many proteins translocated into the ER. Because proteins of the secretory pathway often mediate crucial signaling roles (e.g., cell surface receptors, transporters, or polypeptide hormones), incompletely folded forms are not tolerated by the cell and instead are disposed of by stringent quality control systems. Through a process called ER-associated degradation (ERAD), unfolded proteins are removed to the cytosol for subsequent ubiquitylation and degradation by the 26S proteasome (8–10). Ultimately, this may lead to a deficiency of important proteins and a loss of the functions they serve. Moreover, folding efficiency for an individual protein can be further compromised by the presence of an inherited mutation in the gene that encodes it. For example, specific mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) cause problems in its folding within the ER, leading to depletion of this essential ion channel that normally transports chloride across epithelial tissues and to the life-threatening disease cystic fibrosis (11).

ENDOPLASMIC RETICULUM STRESS: WHEN THE CAPACITY TO FOLD PROTEINS WITHIN THE ENDOPLASMIC RETICULUM FAILS TO KEEP UP WITH DEMAND

The capacity for folding proteins within the ER varies greatly among cells types. Cells with the potential to secrete high protein loads are able to do so because they contain a large, well-developed ER. For example, each β -cell of the endocrine pancreas is capable of

synthesizing and secreting up one million molecules of insulin per minute; in insulinresistant states, this enormous protein synthetic load becomes even greater (12). Moreover, plasma cells can secrete their own weight in antibodies every day (13). In contrast, cells that do not routinely secrete large protein loads conserve resources by maintaining a relatively small ER with limited protein-folding capacity.

Regardless of the size of their ER, cells appear to work near the limits of their secretory capacity and often encounter conditions during which the workload imposed on the ER protein-folding machinery exceeds its capability. When the ER protein-folding capacity is overwhelmed, cells are said to be experiencing ER stress (14). A wide range of cellular disturbances can disrupt the efficiency of protein folding in the ER and lead to the accumulation of misfolded proteins within this organelle, including nutrient deprivation, hypoxia, point mutations in secreted proteins that stabilize intermediate folding forms or cause aggregation, and loss of calcium homeostasis with detrimental effects on ER-resident calcium-dependent chaperones (7, 15, 16). In the case of pancreatic β -cells, ER stress can occur as a result of the inability to fold the increased levels of insulin intermediates needed to maintain blood glucose (17). In other cells, such as neurons, the chronic expression of folding-defective secretory proteins can put unsustainable demands on the protein-folding machinery and lead to ER stress (18). Under ER stress, secretory proteins start to accumulate in improperly modified and unfolded forms within the organelle. Therefore, cells have evolved a sophisticated surveillance system to sense and respond to ER stress before it becomes a threat to their survival.

THE UNFOLDED PROTEIN RESPONSE: A SIGNALING PATHWAY THAT DETERMINES CELL FATE UNDER ENDOPLASMIC RETICULUM STRESS

To ensure that protein-folding capacity is in balance with demand, cells constantly monitor the amount of misfolded protein in the ER lumen and initiate corrective responses. When misfolded proteins in the ER accumulate above a critical threshold, this accumulation signals incipient problems in protein folding and sets in motion a signal transduction pathway called the unfolded protein response (UPR) that attempts to remedy the situation. An ancient signaling pathway conserved all the way from yeasts to mammalian cells, the UPR is initiated by three ER transmembrane proteins: IRE1a (inositol-requiring enzyme 1a), PERK (pancreatic endoplasmic reticulum kinase), and ATF6 (activating transcription factor 6) (19, 20). The common theme among these three ER stress sensors is that they all contain an ER-luminal domain believed to be capable of directly or indirectly sensing misfolded proteins when the latter reach critically high concentrations (Figures 1–3) Luminal domain sensing of misfolded proteins leads to changes in the oligomerization state of each sensor and activation of their associated downstream activities, thus transducing a signal from the ER lumen into the cytoplasm. For IRE1a and PERK, luminal domain dimerization is sufficient to initiate their activation, and this event may normally be prevented in unstressed cells by the binding of an ER chaperone called BiP. This inhibition of dimerization may be relieved when BiP is titrated away from the luminal domain by its high affinity for misfolded proteins (21). Furthermore, through direct binding to the luminal domain of IRE1a and PERK, misfolded proteins may act as activating ligands for these ER

stress sensors—a mechanism analogous to a wide range of extracellular ligands that activate various receptors present on the plasma membrane (22, 23).

All three UPR sensors have outputs that first attempt to realign protein-folding demand and capacity back into homeostasis so that the cell can continue to survive and function (24). In order to increase protein-folding capacity, the homeostatic UPR expands the size of the ER through increased biogenesis of components (e.g., protein and lipid) and upregulates the transcription of ER chaperones. Concurrently, the combined outputs of the homeostatic UPR increase transcription of ER-resident enzymes and structural components that lead to the removal and degradation of misfolded proteins from the ER lumen. However, despite their common mechanism of luminal activation, each UPR sensor signals in a unique way and has a distinct set of targets. The cytoplasmic tail of IRE1a has two enzymatic activities—a serine/threonine kinase domain and an endoribonuclease (RNase) domain-that work together as a team (25, 26). Upon luminal binding of misfolded proteins, IRE1a's kinase becomes activated and *trans*-autophosphorylates; this event leads to allosteric activation of its adjacent RNase domain (27). When active, IRE1a's RNase excises a 26-nt intron from the mRNA encoding the XBP1 (X-box protein 1) transcription factor. Cytosolic splicing of the two resulting mRNA fragments by a yet-to-be-identified ligase produces the homeostatic transcription factor XBP1s that contains a transactivation domain encoded in the altered reading frame (28, 29). XBP1s then translocates to the nucleus and induces transcription of many genes that augment ER size and function (30). In contrast, PERK contains a single cytosolic kinase that phosphorylates eukaryotic translation initiation factor 2a (eIF2a) (31, 32). Phosphorylation inhibits eIF2a activity and hence slows down global protein translation, giving the cell extra time to attempt to fold the backlog of proteins already present in the ER lumen (33). In the presence of misfolded proteins, ATF6 translocates to the Golgi and is cleaved by the Site-1 and Site-2 proteases to release the ATF6(N) transcription factor contained within its cytoplasmic tail (34). Together with XBP1s, ATF6(N) increases transcription of targets that expand ER size and increase its protein-folding capacity (30). Collectively, these transcriptional events act in a coordinated fashion as homeostatic feedback loops to temper ER stress. If successful in reducing the amount of misfolded protein, UPR signaling is attenuated and the cell survives.

However, if its adaptive responses prove inadequate to restore protein-folding homeostasis, UPR signaling will be sustained, indicating high or chronic ER stress. In the setting of irremediable ER stress, the UPR transforms into an alternate signaling platform called the terminal UPR that actively promotes cell death (35). Although the molecular details are still being worked out, evidence suggests that each of the three UPR sensors has a distinct set of proapoptotic outputs that contribute to cell demise if ER stress cannot be resolved. For example, although a temporary pause in protein translation due to eIF2a phosphorylation can be beneficial for cells under ER stress, a protracted block in translation from sustained PERK signaling is incompatible with survival. Moreover, PERK hyperactivation can upregulate the CHOP (C/EBP-homologous protein; also known as GADD153) transcription factor, which inhibits expression of the gene encoding antiapoptotic BCL-2 to hasten cell death (36, 37). Likewise, when hyperactivated by chronic ER stress, phosphorylated IRE1a transitions from homodimers into high-order oligomers, allowing its RNase to acquire affinity for additional RNA substrates aside from XBP1 mRNA. Under sustained

oligomerization, IRE1a's RNase causes endonucleolytic decay of hundreds of ER-localized mRNAs containing an N-terminal signal sequence, which depletes ER cargo and protein-folding components to further worsen ER stress (38, 39). In addition to reducing key protein-folding components in this way, IRE1a oligomerization has been shown to induce activation or upregulation of a number of proinflammatory and prodeath proteins. For example, when hyperactivated, IRE1a's RNase reduces the levels of select microRNAs (possibly by directly cleaving their precursors at the ER membrane) that normally repress proapoptotic targets such as the pro-oxidant protein TXNIP (thioredoxin-interacting protein), leading to their rapid upregulation (40, 41). Increased TXNIP expression then activates the inflammasome and its Caspase-1-dependent prodeath pathway (41). Finally, sustained IRE1a oligomerization may serve as an activation platform for ASK1 (apoptosis signal–regulating kinase 1) and its downstream target JNK (c-Jun NH₂-terminal kinase) (42, 43). Phosphorylation by JNK has been reported to both activate proapoptotic BIM and inhibit antiapoptotic BCL-2. ATF6a likely has proapoptotic targets as well, but these have not been well defined.

Many, but not all, of the prodeath signals sent from the UPR sensors ultimately converge on the mitochondrial apoptotic pathway. This apoptotic pathway is initiated when toxic mitochondrial proteins, such as cytochrome *c* and Smac/Diablo, are forcibly released into the cytoplasm, where they lead to activation of downstream effector caspases (e.g., Caspase-3) (44). The BCL-2 family, a large class of both pro- and antideath proteins, governs the intrinsic apoptotic pathway by regulating the integrity of the outer mitochondrial membrane (45). This pathway is engaged when cell injury leads to the expression and/or posttranslational activation of one or more BH3-only proteins, a structurally diverse collection of prodeath proteins that contain a short α-helix known as the BH3 (Bcl-2 homology 3) domain necessary for their killing activity (46). Once activated, BH3-only proteins disable mitochondrial protecting proteins (e.g., BCL-2, BCL-XL, MCL-1) and in some cases directly trigger the multidomain proapoptotic BAX and BAK proteins to permeabilize the outer mitochondrial membrane.

The terminal UPR has been reported to activate at least four distinct BH3-only proteins (BID, BIM, NOXA, PUMA) that then signal the mitochondrial apoptotic machinery (47–49). Each of these BH3-only proteins is activated by ER stress in a unique way. For example, BIM is transcriptionally upregulated by PERK and its protein product stabilized through JNK dephosphorylation in response to ER stress (48). However, it remains unknown whether these BH3-only proteins are simultaneously set in motion by all forms of ER stress or whether only a subset is activated under specific forms of pathological insult that injure this organelle. Researchers are working hard to unravel the molecular details of how ER damage is communicated to the apoptotic machinery, as these signals may represent therapeutic targets in some of the ER stress–related diseases discussed below (50).

THE ROLE OF ENDOPLASMIC RETICULUM STRESS IN DISEASE

Over the past decade, cell injury secondary to chronic ER stress has been increasingly implicated as a central contributor to the pathophysiology of a wide range of prevalent human diseases (20). For example, ER stress and sustained UPR signaling have been well

documented in affected tissues in diabetes, neurodegeneration, stroke, pulmonary fibrosis, viral infection, inflammatory disorders, cancer, and heart disease. The common theme among these seemingly disparate diseases is the presence of intracellular and/or extracellular conditions that disrupt protein folding and lead to the accumulation of misfolded proteins in the ER. In strong support of the notion that ER stress can contribute to pathology, inherited mutations in the UPR pathway have been associated with rare forms of diabetes and other diseases in humans (see below). For many of the above-mentioned diseases, genetic manipulation of specific UPR components has been shown to influence disease outcome in rodent models. The existing preclinical data linking ER stress to disease and the emergence of potential targets in the UPR will likely lead to human clinical trials of UPR-directed drugs within the next few years. A few of the diseases most strongly associated with ER stress are discussed below.

Diabetes Mellitus

Pancreatic β -cells synthesize, store and secrete large amounts of the polypeptide hormone insulin; in fact, it is estimated that each human β -cell produces on average about one million molecules of insulin every minute (51). In response to increases in ambient blood glucose levels, prepackaged insulin in secretory granules is released by the β -cell and is replenished by synthesis. A signal transduction cascade ensues when insulin binds its receptor on insulin-responsive cells in peripheral tissues. Upon insulin binding to a target cell, glucose consequently enters, causing energy production. Simultaneously, as blood glucose levels normalize, the stimulus for further insulin release from pancreatic β -cells is removed. This glucostatic cycle is dysregulated in the disease diabetes mellitus, ultimately because of an insufficient mass of functioning β -cells needed to produce the requisite amounts of insulin in the fasted and postprandial states in order to maintain normoglycemia (Figure 4).

To support high-level insulin secretion, β -cells contain highly developed ERs. Insulin biogenesis requires a complex series of molecular biosynthetic events that initiate in the β cell ER (52). The precursor to insulin, preproinsulin, is cotranslationally translocated into the ER lumen, where its signal sequence is subsequently clipped off, yielding proinsulin. ER-resident oxidoreductases catalyze the formation of three intramolecular disulfides in proinsulin, which allow it to fold to its native shape. This critical oxidative folding step is interrupted in the Akita diabetic mouse mutant, which expresses a proinsulin variant gene, Ins2 (C96Y)—Akita insulin. Ins2 (C96Y) lacks a cysteine needed to form one of the intramolecular disulfide bonds that helps it fold in the ER; its trafficking is therefore impeded, unlike wild-type proinsulin, which is properly trafficked to downstream Golgi and secretory granules, where it is further processed by endoproteases that remove its C-peptide to generate mature insulin (53–55, 56).

Akita vividly links ER stress to the death of β -cells and diabetes. Despite retaining three normal insulin gene copies (mice possess two distinct insulin-encoding genes), Akita mice suffer from insufficient insulin production secondary to β -cell loss. Akita insulin dominantly causes a toxic gain-of-function diabetic syndrome (55). By accumulating in the ER as a conformationally altered immature species, Akita insulin acts as a proteotoxin that exhausts homeostatic UPR outputs and instead triggers a terminal UPR (53). This causes β -cells in

the Akita mouse to deterministically enter the apoptotic pathway, leading to frank diabetes within 4 to 5 weeks after birth (15). Rare infantile diabetes–causing Akita-like insulin mutations have been recently described in humans (57). Interestingly, genetic removal of either CHOP, a proapoptotic transcription factor downstream of PERK, or the IRE1 α target TXNIP ameliorates β -cell loss and diabetes in the Akita background, emphasizing the central role the terminal UPR plays in β -cell degeneration (15, 41).

A recessive example of diabetogenic UPR dysregulation is evident in *Perk*-knockout mice. Homozygous deletion in mice of the gene encoding the UPR sensor PERK causes massive and rapid β -cell apoptosis, leading to infantile diabetes (58, 59). *Perk*-knockout mice further develop pancreatic exocrine insufficiency and exhibit growth defects early in life. These defects are believed to be secondary to the dysfunction and death of several different important professional secretory cell types; intriguingly, diabetes mellitus is one of the earliest and most severe phenotypes in the mutant animals, again highlighting the susceptibility of the β -cell to ER stress. A rare human diabetic syndrome caused by *PERK* null gene mutations (called Wolcott–Rallison syndrome) phenocopies many of the features of the *Perk*-knockout animals.

Underlying mechanisms of cell degeneration due to PERK deficiency are understood in considerable detail, although many important questions remain. As mentioned above, PERK is an ER transmembrane kinase that dimerizes under ER stress, which consequently increases phosphorylation activity against its downstream substrate, the eIF2a translation initiation factor. Activated PERK phosphorylates eIF2a on the Ser51 residue, which causes translation to cease globally because it depletes the eIF2·GTP·Met-tRNA; complex needed to initiate cap-dependent mRNA translation. It has been proposed that in the absence of PERK, β -cells cannot properly attenuate translation to match ER protein-folding capacity. As a consequence, they suffer deposition of unfolded proteins in the ER (58). Consistent with this notion, the ERs of β -cells in *Perk*^{-/-} mice are distended with electron-dense proteinaceous material, and the islets exhibit a high rate of apoptosis. Additionally, there appears to be a marked decline in β -cell proliferation early in neonatal life in the knockout animals. The inability to compensate for increased rates of apoptosis through increasing β-cell proliferation is likely to be an important component in the endocrine pancreatic failure in the $Perk^{-/-}$ animals (60). The homeostatic arm of the IRE1a pathway has also been found to be necessary for β -cell function. For example, mice with a β -cell-specific deletion of the *Xbp1* gene show impaired proinsulin processing and decreased insulin secretion (61). Likewise, ATF6 is critical to prevent β -cell loss in a mouse model of diabetes (62).

Translation attenuation through phosphorylation of eIF2 α by PERK kinase may be used by the β -cell as a glucose-sensing mechanism to prime the secretory apparatus for upcoming synthesis and structural maturation of proinsulin in response to changes in glucose levels in the blood (63). However, the strategy of limiting translation is fraught with the danger that should upstream stress remain unrelieved, cells may never resume translation at levels needed to recover viability. Thus, there are safety valves that extinguish PERK signaling after a time window has elapsed.

The importance for glycemic control of an escape from this global translational block is seen in two mouse mutant models. The first, a knock-in mouse mutant expressing an unphosphorylatable eIF2a version (S51A), develops a severe wasting syndrome shortly after birth life due to arrested hepatic gluconeogenesis in the homozygote (64, 65), and a milder insulin-resistant hyperglycemic syndrome in the heterozygote (65). The second model involves p58^{IPK}, a cochaperone produced several hours after the UPR has been initiated. p58^{IPK} may help to close a timing loop necessary to turn off the UPR by inhibiting eIF2a kinases, including PERK. Homozygous loss of the gene encoding p58^{IPK} causes diabetes (66–68). By the time p58^{IPK} is produced, if cells have not yet returned to homeostasis, the UPR may switch from promoting homeostasis to an apoptotic mode. In the absence of p58^{IPK}, a continued translational block through PERK may signal a frustrated UPR cycle. Together, the p58^{IPK}, eIF2a (S51A), and PERK genetic models offer fascinating insights into the importance of temporal control of UPR signaling for cell fate.

Another genetic example of ER stress–induced diabetes comes from the $Wfs1^{-/-}$ mouse, and in humans with Wolfram syndrome. WFS1 is an ER transmembrane protein whose loss causes early-onset diabetes, neurodegeneration, and optic and auditory defects (69). The WFS1 protein is widely expressed in diverse tissues and is thought to aid protein assembly and ERAD (70) as well as control processing of the UPR sensor ATF6 (70, 71). As with PERK and p58^{IPK} deficiency, the earliest defects manifest in pancreatic β -cells.

The aforementioned rare disease examples vividly show that accumulation of unfolded protein in the ER and removal of key UPR functions promote apoptosis in β -cells to cause diabetes. Although fascinating from an academic standpoint, the excitement of these experimental rare diseases stems from the question of whether they inform the understanding of common human diabetic syndromes—i.e., type 1, type 2, and gestational diabetes. In all three of these diseases, underlying mechanisms of β -cell functional shutdown and degeneration are evident. As individual units, functioning β -cells in a pancreas may experience increasing and unresolvable ER stress as they compensate for neighboring β -cells that have become dysfunctional through disease and aging. Studies confirm that β -cells of mice may already be functioning (even in healthy states) at levels of UPR activation that are significantly greater than in other professional secretory cells (72). Therefore, without a wide margin for further homeostatic adjustment, β -cells could quickly cross a threshold that puts them at risk for dedifferentiation and apoptosis through a terminal UPR. Thus, as percell ER stress levels rise, and terminal UPR outputs stochastically cause the death of individual cells, vicious cycles leading inexorably to whole pancreatic organ failure may set in. In this unifying scheme, the upstream stresses differ for type 1 and type 2 diabetes, but the downstream outcomes will be common: For type 1 diabetes, a disease that results from autoimmune attack by T lymphocytes against β -cells, as β -cell function degenerates, remaining cells in islets would necessarily have to compensate by overworking and may themselves experience critical thresholds of ER stress. Thus, UPR-mediated apoptosis from within the β -cell may synergize with prodeath processes initiated by autoimmune attack from without through cells of the innate and adaptive immune system (73).

For type 2 diabetes, a disease provoked by peripheral insulin resistance, β -cells are forced to compensate by increasing insulin production to abnormally high levels (74). Successful β -

cell compensation may prevent progression to frank diabetes, but in some insulin-resistant patients the dysfunction and death of enough β -cells (roughly half of the original β -cell mass) occurring over many years may lead to a tipping point and organ failure beyond which normoglycemia cannot be maintained. The inability to compensate for declining numbers of β-cells through cell proliferation in some populations may contribute to this process. Interestingly, although a prediabetic (compensated) state may exist stably for years, progression to diabetes occurs on a much shorter timescale (perhaps weeks) (75, 76), indicating that an acute-on-chronic organ shutdown may be occurring; the often dramatic initial presentation of type 1 diabetes is also consistent with such dynamics. Intriguingly, both disorders may have a long metastable period of failing compensation, which perhaps could be exploited through intervention in those at risk (as in the Diabetes Prevention Program) (77). Although type 1 diabetes has traditionally been regarded primarily as a discrete and homogeneous disorder (when compared with the greater heterogeneity evident in β -cell function of patients with type 2 diabetes), recent findings of persistent C-peptide production and preserved β-cell function decades after diagnosis in distinct groups of patients may indicate inherent differences in populations with regard to the ability to tolerate stress in the β -cell (78).

Neurodegeneration

A pathologic hallmark of many neurodegenerative diseases is the accumulation of misfolded proteins and protein aggregates within affected neurons and surrounding supporting cells (79). For example, in Alzheimer's disease (AD), intracellular deposits of tau are observed in neurofibrillary tangles, and extracellular aggregates of amyloid β are seen in senile plaques. The pathology of progressive supranuclear palsy (PSP) shows intracellular tangles of tau protein throughout the neocortex, basal ganglia, and brainstem. In Parkinson's disease (PD), ubiquitinated protein deposits called Lewy bodies (composed of a-synuclein) are seen in neuronal cytosol. Some cases of inherited amyotrophic lateral sclerosis (ALS) are caused by toxic, gain-of-function point mutations in SOD1 (superoxide dismutase 1) that lead to its aggregation. Other neurodegenerative disorders, such as Huntington's disease (HD), result from mutant proteins (e.g., huntingtin) containing expanded glutamate repeat sequences. Both mutant SOD1 and mutant huntingtin proteins aggregate, exhaust 26S proteasome activity, and result in secondary accumulations of misfolded proteins in the ER (80, 81). Prions are well known to organize into protein aggregates and are causally involved in the pathogenesis of a group of transmissible spongiform encephalopathies, including Creutzfeldt-Jakob disease and kuru (82). In addition to rare inherited mutations in a single protein that disrupt its proper folding, degenerating neurons are exposed to numerous other insults (e.g., oxidative stress, inflammation, metabolic disturbances) that can compromise protein folding and lead to ER stress.

How protein aggregates contribute to selective neuronal loss in each of these neurodegenerative diseases remains under active investigation. There is evidence that large protein aggregates (e.g., inclusion bodies) in neurodegenerating tissues are protective, whereas smaller misfolded protein species (e.g., fibrils) are toxic (83, 84). Accumulation of toxic protein species can kill neurons (85), and there is growing evidence that ER stress is an important mechanism driving this neurotoxicity (Figure 5) (86–88). IRE1a activation and

UPR induction are present in postmortem brain and spinal cord tissues in AD (89-92), PD (93, 94), and ALS (95). Moreover, the accumulation of protein aggregates in cellular and animal models of HD (43), PD (96), and ALS (97-99) strongly correlate with UPR activation. A recent genome-wide association study for PSP found a risk locus that mapped to the gene encoding PERK (100), and an independent study found evidence for PERK hyperactivation in disease-affected brain regions in PSP (101). Brain samples from patients who have succumbed to Creutzfeldt-Jakob disease show activation of a number of ER chaperones and other ER stress markers. Spinal cord segments taken at autopsy of patients with sporadic ALS show that ER stress results in the induction of the UPR, chaperones, and apoptotic markers (95, 102). Using an elegant in vivo reporter system in mutant SOD1 mice, researchers found that the UPR is activated in selectively vulnerable (but not diseaseresistant) motor neurons 25-30 days before the earliest signs of neuromuscular denervation (103). Coupled with evidence that mutant SOD1, but not wild-type SOD1, leads to secondary accumulations of misfolded proteins in the ER, these findings strongly suggest that affected neurons attempt to manage the accumulation of misfolded proteins in the ER by activating the UPR (98). Importantly, UPR upregulation is observed prior to the onset of symptoms, suggesting an active role in the disease (97).

Based on these data, researchers have recently begun to manipulate the UPR in mouse models of neurodegeneration and have uncovered some promising results. For example, conditional deletion of *Perk* ameliorated synaptic dysfunction in the APP/PSI transgenic model of AD (104). Furthermore, a recent study found that oral administration of a highly selective PERK inhibitor that efficiently crosses the blood-brain barrier significantly reduced neurodegeneration and clinical disease in prion-infected mice (105). In combination with the strong evidence for UPR activation in human patients, studies such as these are bringing great attention to the idea that pharmacologic manipulation of the UPR may have disease-modifying benefits for a variety of neurodegenerative diseases.

Heart Disease, Stroke, and Ischemia-Reperfusion Injury

The link between ER stress and ischemia-reperfusion injury has been implicated on a number of levels (106). Reductions in blood flow as a result of arterial occlusion or hypotension cause tissue hypoxia and hypoglycemia, two conditions that rapidly induce protein misfolding and ER stress. When blood flow is restored, reperfusion of the affected tissues leads to oxidative stress and alterations in the redox status of the ER that disrupt protein disulfide formation and cause ER protein misfolding. UPR activation has been documented in atherosclerotic plaques in both humans and various animal models (107, 108). There is also evidence that high cholesterol, fatty acids, and oxidative stress can trigger ER stress-induced apoptosis of macrophages and endothelial cells associated with atherosclerotic plaques and worsen the progression of atherosclerosis (109). Cardiac myocytes within and adjacent to the field of myocardial infarction activate the UPR. Moreover, genetic deletion of Ask1 in mice partially preserves left ventricular function after coronary artery ligation, implicating the terminal UPR as an important contributor to myocyte loss during myocardial infarction (110). Similarly, brain regions affected by stroke also show evidence of ER stress-induced apoptosis, and mice deficient in Chop show decreased neuronal loss after stroke injury compared with wild-type controls (111).

Cancer

Tumor cells often invade or metastasize into foreign environments where unfavorable conditions, such as hypoxia, glucose deprivation, lactic acidosis, oxidative stress, and inadequate amino acid supplies, compromise protein folding in the ER (112–115). Consequently, many studies have found evidence of sustained and high-level activation of all three branches of the UPR (PERK, ATF6, IRE1a) in a wide range of primary human tumor types, including glioblastoma, multiple myeloma, and carcinomas of the breast, stomach, esophagus, and liver (22, 116–120). Genomic screens have identified rare somatic mutations in *IRE1a* in a small percentage of human solid tumors (121). Numerous reports have also shown that components of the ER protein-folding machinery, most notably the chaperone Bip/GRP78, are overexpressed in cancers at levels that correlate with disease progression (122, 123).

However, despite the overwhelming evidence of ongoing ER stress and UPR activation in many forms of cancer, whether these processes ultimately inhibit or promote tumor growth in patients remains an area of intense inquiry. Most of the evidence arguing that the UPR supports tumor growth comes from xenograft studies in mice, in which genetically deleting one or more branches of the UPR or altering the expression of the ER chaperone Bip/GRP78 inhibits the in vivo growth of tumor cells (124–127). For example, genetic deletion of *IRE1a* in a human glioma cell line resulted in reduced angiogenesis and decreased tumor growth when these cells were subsequently injected into mice (128). The IRE1a-XBP1 signaling pathway has been found to induce a number of proangiogenic factors, such as VEGF (vascular endothelial growth factor), which may be one mechanism through which this arm of the UPR can promote the growth of solid tumors (129). These findings suggest that not only is the homeostatic UPR frequently activated in tumors, but it may be necessary for the survival and/or growth of the cancer cells under conditions that stress the ER.

Myeloma, a highly secretory tumor composed of malignant plasma cells, is one cancer for which the UPR is frequently mentioned as a potentially attractive target on the basis strong evidence that this pathway is essential for plasma cell development. In mice, IRE1a and its homeostatic target XBP1 are both required for the differentiation of B lymphocytes into plasma cells (130, 131), illustrating a critical role for the secretory pathway in the health of this cell type. Interestingly, up to 50% of primary myelomas show unusually high levels of XBP1s (116). Moreover, mice expressing a transgene of *Xbp1s* (that is missing the 26-nt intron and hence requires no further processing by IRE1a) in B lymphocytes develop a plasma cell malignancy closely resembling myeloma (116). There is also evidence to suggest that proteasome inhibition with bortezomib (Velcade), which is approved by the US Food and Drug Administration as first-line therapy for myeloma, leads to myeloma cell death in part by preventing disposal of misfolded proteins through the ERAD pathway and thus triggering ER stress-induced apoptosis (132, 133). On the basis of these findings, several pharmacologic inhibitors of the IRE1a RNase activity have recently been tested on human myeloma xenografts and were found to have antimyeloma activity (134, 135); however, the specificity and off-target effects of these pharmacological agents are not yet well understood.

Although the above findings suggest an oncogenic role for XBP1s in the development of myeloma, recent data have emerged that challenge this notion. First, downregulation of XBP1s expression in myeloma correlates with resistance to bortezomib (136, 137). Second, using a combination of whole-genome and whole-exome sequencing of primary tumors from 38 myeloma patients, researchers discovered *XBP1* mutations in two of these patients (138). On further analysis, these mutations were shown to inactivate XBP1s, arguing against an obligate role for this transcription factor in myeloma. Finally, it was recently reported that genetic knockdown of IRE1a or XBP1 in human myeloma cell lines is well tolerated and leads to bortezomib resistance (139), challenging the rationale for using IRE1a inhibitors in this disease. Overall, the lessons from myeloma to date suggest that the effects of the UPR (or at least its IRE1a-XBP1 branch) on tumor development and maintenance are more complicated and nuanced than originally anticipated.

Does the Unfolded Protein Response Present Drug Targets?

Given the evidence of UPR deregulation across a range of human disease, there is great interest in the possibility of pharmacologically modulating its outputs to control cell fate under ER stress. For cell degenerative diseases such as type 2 diabetes and neurodegeneration, the biasing of the UPR's homeostatic-apoptotic switch to favor cell survival may potentially be disease modifying. However, the parallel and cross-wired networking of the UPR may require multiple nodes to be targeted simultaneously to achieve the desired benefits. One approach may be to prolong the adaptive phases of the UPR to maximize chances of recovery (examples of such targets would include the XBP1 and ATF6 transcription factors); such a regime may involve preconditioning the secretory pathway by preemptive UPR activation to make it more robust. A different approach is to inhibit key mediators of apoptosis (CHOP, TXNIP). Perhaps potential timers of the UPR, such as GADD34 and p58^{IPK}, may also pose attractive targets. In this vein, a small molecule called salubrinal was shown to block phosphatases mediating eIF2a dephosphorylation and hence to enhance cell survival under ER stress (140).

Preemptive preconditioning was demonstrated to be partially cytoprotective in proof-ofconcept experiments using synthetic dimerizable modules of PERK in combination with small-molecule dimerizers in cell culture models of ER and oxidative stress (141). It is unclear, however, whether long-term PERK activation, with its attendant consequences of inhibiting translation, would be efficacious in vivo. Related to such an approach, we (142) and others (143) have demonstrated that preempting IRE1a's homeostatic activation mode can partially prolong cell survival under ER stress. The basis for this particular approach rests on a highly unusual relationship that we discovered between IRE1a's two catalytic domains (144); the kinase domain of IRE1a can be engaged with a designer kinase inhibitor called 1NM-PP1 to enforce an active ATP-binding conformation while bypassing autophosphorylation (145), which spontaneously triggers RNase activity even without ER stress and forces splicing of XBP1 mRNA, leading to production of the prosurvival Xbp1 transcription factor. Cells subjected to these maneuvers preemptively enjoy a small, but significant, measure of cytoprotection when they are challenged to ER stress (142). It remains to be seen whether these proof-of-concept strategies will be applicable to the wildtype versions of IRE1a and PERK.

Irremediable ER stress hyperactivates both PERK and IRE1 α , leading to entry into apoptosis. Thus, a diametrically opposite strategy is to inhibit the hyperactivated state by using inhibitors of PERK (146), RNase inhibitors of IRE1 α (134), or allosteric RNaseinhibitory type II kinase inhibitors of IRE1 α (145). Early reports suggest that a PERK inhibitor may protect against preclinical models of neurodegeneration (105), but much more work needs to be done to understand the potential benefits and risks of inhibiting ER stress– induced cell degeneration in vivo.

Finally, it has been demonstrated that small chemical chaperones such as tauroursodeoxycholic acid, which act as templates for protein folding and have global effects on stabilizing protein conformations, may also afford significant protection against preclinical models of diabetes (147, 148).

Although they are by no means exhaustive, these examples illustrate emerging concepts about the role of ER stress in disease and the therapeutic potential of targeting components of the UPR to control cell fate under conditions of ER stress.

CONCLUSIONS

The UPR is a highly conserved signal transduction pathway that is activated when cells are unable to keep up with the protein-folding demands on the ER—a form of cell injury called ER stress. In response to ER stress, the UPR initially sends out adaptive outputs that decrease the load and increase the capacity of the ER secretory pathway in an effort to restore ER homeostasis. However, under irremediable ER stress, the UPR assembles into a platform that sends out proinflammatory and prodeath signals to cause cell demise. Cell injury from chronic ER stress is emerging as central to the pathophysiology of a wide range of prevalent human diseases, including diabetes, neurodegeneration, stroke, and cancer. Recent advances in our understanding of how the UPR switches from life to death signaling will hopefully lead to new strategies to combat these ER stress–associated diseases.

TERMS AND DEFINITIONS

ASK1

apoptosis signal-regulating kinase 1

ATF6

activating transcription factor 6

Chaperones

a family of enzymes that bind maturing client proteins, prevent their aggregation, and catalyze their folding to native structures

eIF2a

eukaryotic translation initiation factor 2

ER stress

a form of intracellular stress that occurs whenever the protein-folding capacity of the ER is overwhelmed

ERAD

ER-associated degradation

Inflammasome

a multiprotein component of the innate immune system that in response to various signals activates procaspase-1 and inflammatory cytokines

IRE1a

inositol-requiring enzyme 1a

JNK

c-Jun NH2-terminal kinase

Mitochondrial apoptotic pathway

genetically controlled cell suicide program initiated when toxic mitochondrial proteins, such as cytochrome *c*, are released into the cytoplasm

PERK

pancreatic endoplasmic reticulum kinase

TXNIP

thioredoxin-interacting protein

Unfolded protein response (UPR)

intracellular signaling pathway that responds to an accumulation of misfolded proteins in the ER

XBP1

X-box protein 1

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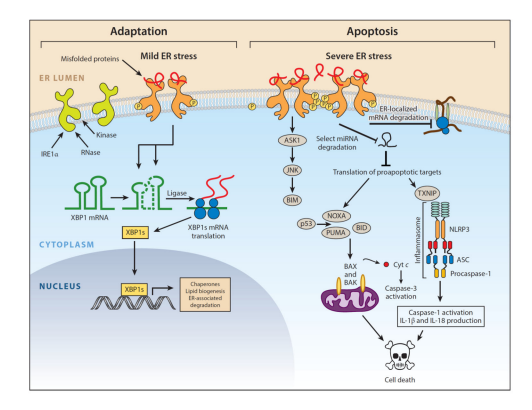


Figure 1.

IRE1a arm of the unfolded protein response. IRE1a is an endoplasmic reticulum (ER) transmembrane protein that becomes activated when misfolded proteins in the ER lumen bind to its luminal domains. The cytoplasmic tail of IRE1a has two enzymatic activities—a serine/threonine kinase domain and an endoribonuclease (RNase) domain. Upon luminal binding of misfolded proteins, IRE1a's kinase becomes activated and transautophosphorylates multiple serine/threonine residues on the cytosolic tail. IRE1a phosphorylation leads to allosteric activation of the adjacent RNase. The consequences of IRE1a activation vary depending on the level of ER stress. In response to low levels of ER stress, IRE1a's RNase excises a 26-nt intron from the mRNA encoding the XBP1 (X-box protein 1) transcription factor to produce the homeostatic transcription factor XBP1s. XBP1s then translocates to the nucleus and induces transcription of many genes that augment ER size and function in an attempt to restore ER homeostasis. However, if ER stress is irremediable, IRE1a becomes hyperactivated and undergoes homo-oligomerization. Under sustained oligomerization, IRE1a's RNase endonucleolytically degrades hundreds of ERlocalized mRNAs containing an N-terminal signal sequence, which depletes ER cargo and protein-folding components to further worsen ER stress. Moreover, when hyperactivated, IRE1a's RNase directly cleaves select microRNAs that normally repress proapoptotic targets. In addition to signaling through RNA substrates, IRE1a oligomerization has been shown to induce activation or upregulation of a number of proinflammatory proteins, including the pro-oxidant protein TXNIP (thioredoxin-interacting protein), to activate the inflammasome and its Caspase-1-dependent prodeath pathway. Finally, sustained IRE1a oligomerization serves as an activation platform for ASK1 (apoptosis signal-regulating kinase) and its downstream target JNK (c-Jun NH2-terminal kinase). Phosphorylation by

JNK has been reported to both activate proapoptotic BIM and inhibit antiapoptotic BCL-2. Once activated, BH3-only proteins such as BID and BIM disable mitochondrial protecting proteins (e.g., BCL-2, BCL-XL, MCL-1) and in some cases directly trigger the multidomain proapoptotic BAX and BAK proteins to permeabilize the outer mitochondrial membrane and release toxic mitochondrial proteins, such as cytochrome *c*, into the cytoplasm, where they lead to activation of downstream effector caspases (e.g., Caspase-3) and cell demise. Modified with permission from Reference 149. Copyright © 2014 by Elsevier.

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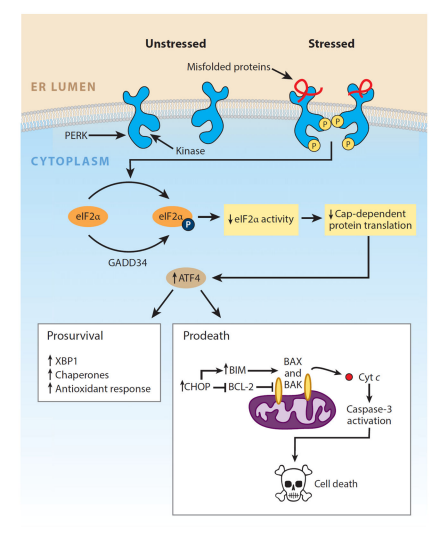


Figure 2.

PERK arm of the unfolded protein response. PERK is an endoplasmic reticulum (ER) transmembrane protein that contains a single cytosolic kinase. When its luminal domains are dimerized in the presence of misfolded proteins, PERK phosphorylates eukaryotic translation initiation factor 2α (eIF2α). Phosphorylation inhibits eIF2α activity and hence slows down global protein translation, giving the cell extra time to attempt to fold the backlog of proteins already present in the ER lumen. In contrast, translation of the transcription factor ATF4 (activating transcription factor 4) is selectively upregulated when the amount of active eIF2α is limiting. ATF4 expression transcriptionally upregulates CHOP (C/EBP-homologous protein; also known as GADD153), which tips the ER toward homeostasis through induction of a number of corrective genes, including XBP1 and chaperones. Although a temporary pause in protein translation due to eIF2α phosphorylation can be beneficial for cells under ER stress, a protracted block in translation from sustained PERK signaling is incompatible with survival. Moreover, high levels of CHOP/GADD153 transcription factor can inhibit the expression of antiapoptotic BCL-2 to hasten cell death and upregulate proapoptotic BIM to trigger activation of the mitochondria-dependent

apoptotic pathway. Modified with permission from Reference 149. Copyright $\ensuremath{\mathbb{O}}$ 2014 by Elsevier.



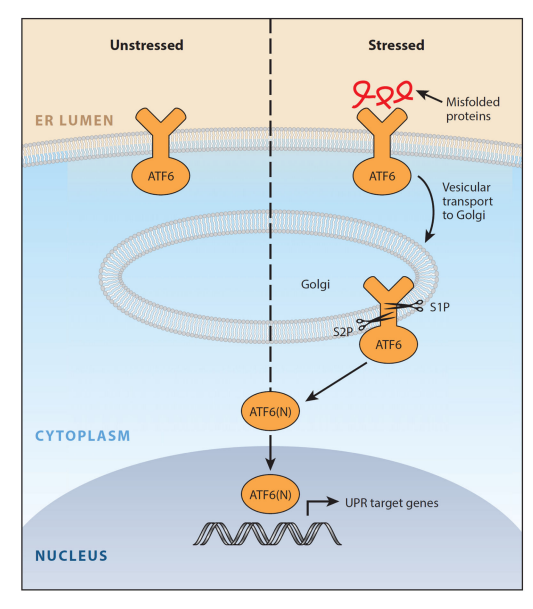


Figure 3.

ATF6 arm of the unfolded protein response. In the presence of misfolded proteins, ATF6 translocates to the Golgi and is cleaved by the Site-1 and Site-2 proteases to release the ATF6(N) transcription factor contained within its cytoplasmic tail. Together with XBP1s, ATF6(N) increases transcription of targets that expand endoplasmic reticulum (ER) size and increase its protein-folding capacity to promote cell survival. Modified with permission from Reference 149. Copyright © 2014 by Elsevier.

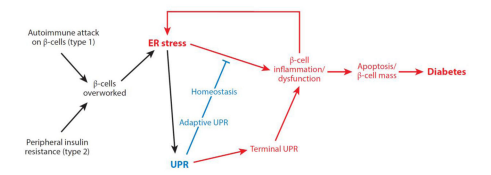


Figure 4.

Model for the role of endoplasmic reticulum (ER) stress in diabetes. ER stress is emerging as an important form of β -cell injury in both type 1 and type 2 diabetes. Overwork of β -cells under conditions of insulin resistance, such as secondary to obesity, may promote attrition of β -cells through unfolded protein response (UPR)-mediated apoptosis in type 2 diabetes. Similarly, as the islet becomes inflamed under autoimmune attack in type 1 diabetes, the percell workload of the remaining β -cells to secrete proinsulin increases. Taken together, these combined insults may lead to critically enhanced ER stress in the remaining β -cells, thus hastening their demise through a vicious cycle that ultimately leads to diabetes. Modified with permission from Reference 149. Copyright © 2014 by Elsevier.

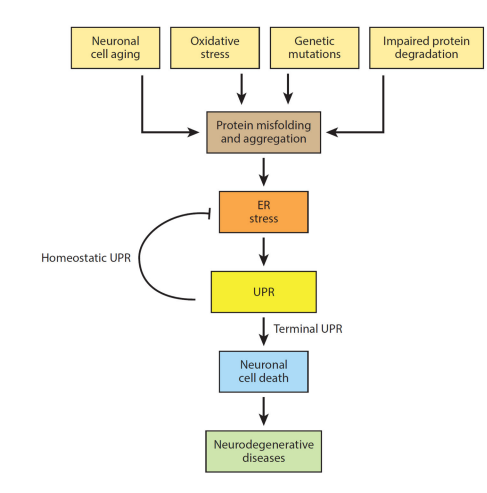


Figure 5.

Model for the role of endoplasmic reticulum (ER) stress in neurodegeneration. In addition to rare inherited mutations in a single protein that disrupt its proper folding, degenerating neurons are exposed to numerous other insults (e.g., oxidative stress, inflammation, metabolic disturbances) that can compromise protein folding and lead to ER stress. Abbreviation: UPR, unfolded protein response. Modified with permission from Reference 149. Copyright © 2014 by Elsevier.