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Inhibition of protein translocation at the endoplasmic reticulum promotes activation of the unfolded protein response

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INTRODUCTION

In eukaryotic cells, the ER (endoplasmic reticulum) is the major site for the synthesis of membrane and secretory proteins. These are co-translationally translocated across or integrated into the membrane through a proteinaceous channel, the Sec61 complex [1]. Once exposed to the ER lumen, polypeptides are folded, assembled and may undergo post-translational modification, such as N-linked glycosylation, prior to being transported to their site of function. However, folding is an intrinsically error-prone process, and polypeptides may also fail to attain their native state due to errors in translation, genetic mutation, absence of partner subunits or unfavourable environmental conditions. Such misfolded or misassembled polypeptides are potentially harmful, and in order to prevent their deployment within the cell, they are retained in the ER by a quality control system [2]. To avoid congestion and potential interference with productive protein folding, misfolded proteins must be cleared from the ER, and this is achieved via a process known as ERAD (ER-associated degradation) [3]. ERAD is mediated by the cytosolic proteasome, which means that proteins selected for ERAD must be retrotranslocated back across the ER membrane into the cytosol prior to degradation [3]. Ubiquitin ligases at the cytosolic face of the ER membrane ubiquitinate the ER substrate, directing it to the UPS (ubiquitin–proteasome system). Ubiquitination also attracts p97, an ATPase that facilitates extraction of many ERAD substrates from the ER membrane. The ERAD substrate is then deglycosylated and deubiquitinated, before entering the catalytic core of the proteasome where it is broken down into small peptides [3].

Situations that perturb the balance between protein synthesis, folding and degradation pathways (termed proteostasis [4]), can lead to a build-up of misfolded proteins in the ER, causing ER stress. Eukaryotic cells possess a signalling network known as the UPR (unfolded protein response), to detect and manage such imbalances in ER proteostasis [5]. UPR signalling pathways function to reduce the rate of protein synthesis and up-regulate the expression of chaperones and ERAD factors, thereby reducing levels of misfolded proteins in the ER and restoring homeostasis. If homeostasis is not re-established, prolonged activation of the UPR can initiate programmed cell death [5]. Disturbances in ER proteostasis and UPR signalling are implicated in the pathogenesis of a diverse range of diseases, including diabetes mellitus, cancer, neurodegeneration, and bone and joint diseases, as well as classical ER protein folding diseases such as cystic fibrosis [6]. As a result, there is considerable interest in the therapeutic potential of selective small molecules to manipulate these pathways in order to ameliorate disease [4].

The small molecules ES8 (eeyarestatin I) and ES9 (eeyarestatin II) are ERAD inhibitors, first identified in a high-throughput screen based on stabilization of GFP (green fluorescent protein)-tagged MHC I heavy chain, which is degraded via an ERAD-like pathway in cells expressing the viral protein US11 [7]. The degradation of a conventional ERAD substrate, TCRα (T-cell receptor α subunit), was also inhibited by ES8 [7], and it was suggested that the compound acts at an early stage in the ERAD pathway, prior to retrotranslocation. Subsequent work provided evidence that a later stage in ERAD, namely the p97 ATPase and an associated deubiquitinating activity, may also be inhibited by ES8 [8,9]. ES8 has also been shown to activate the UPR and induce cell death [10]. Intriguingly, the cytotoxic activity of ES8 appears to be particularly effective against cancer cells, and synergizes with that of bortezomib (Velcade), a proteasome inhibitor with proven anticancer properties [10]. Together with accumulating evidence that UPR activation plays a role in a variety of...
cancers [11], these observations highlight a potential therapeutic application for ES₄-related compounds. Both activation of the UPR and perturbation of cellular ubiquitin homeostasis have been proposed to cause ES₄ cytotoxicity [10]. However, the underlying molecular mechanism(s) through which ES₄ activates the UPR and exerts its effects is not yet clear.

We have previously shown that ES₄ inhibits Sec61-mediated translocation of a range of proteins across the ER membrane, albeit to various degrees depending on the precursor studied [12]. The consequences of ES₄-dependent perturbations of ER translocation in cells are not known, raising the possibility that inhibition of translocation contributes to the potent cellular effects of ES₄. In the present study we show that ES₄ treatment results in the production of mislocalized membrane and secretory proteins, and demonstrate that accumulation of these species correlates with activation of the UPR. Using an alternative inhibitor of translocation, we provide evidence that accumulation of non-translocated polypeptides in the cytosol promotes activation of the UPR.

EXPERIMENTAL

Materials

The M2 anti-FLAG antibody was from Sigma–Aldrich. The P4D1 anti-ubiquitin antibody was from Santa Cruz Biotechnology. Derlin-1 and PDI (protein disulphide-isomerase) polyclonal rabbit antisera were made by Eurogentec. Anti-p97 antiserum was from Professor Bernhard Dobberstein (Center for Molecular Biology, University of Heidelberg, Heidelberg, Germany). Plasmids encoding TCRα were from Professor Ron Kopito (Department of Biology, Stanford University, Stanford, CA, U.S.A.) and Professor Emmanuel Wieritz (Leiden University, Leiden, The Netherlands). Bovine opsin cDNA was subcloned from the pZEO vector into pcDNA5/FRT/TO (Invitrogen). The preprolactin–mCherry construct was from Professor Viki Allan (Faculty of Life Sciences, University of Manchester, Manchester, U.K.).

Synthesis of ES₁, ES₅ and ES₃₅₅

Synthesis of the ESs (see the Supplementary methods section at http://www.BiochemJ.org/bj/442/bj4420639add.htm) commenced with nitroso-compound 1 (see Supplementary Figure S1 at http://www.BiochemJ.org/bj/442/bj4420639add.htm), which was prepared in a straightforward fashion by the reaction of iso-buten with nitrosyl chloride (generated in situ from isoamyl nitrite and HCl) [13]. The reaction of compound 1 with methyl glycinate proceeded smoothly to give a very good yield of oximino ester 2 [14]. Exposure of compound 2 to two equivalents of 4-chlorophenyl isocyanate or 1-naphthyl isocyanate gave the bis-adducts 3a and 3b [3], which were subsequently converted into their corresponding acyl hydrazides 4a and 4b by treatment with an aqueous methanolic solution of hydrazine (acyl hydrazide 4a is the compound also referred to as inactive analogue ES₃₅₅₅). Finally, condensation of hydrazides 4a and 4b with E-3-(5-nitro-2-furyl)acrylaldehyde in methanol gave the corresponding ESs (ES₁ and ES₅) in excellent yields. The double-bond geometry of the final products was indicated as E by the magnitude of the vicinal coupling constant between the olefinic protons (15.8 Hz for ES₁ and 15.4 Hz for ES₅).

Cell culture and transfection

HeLa cells were cultured in DMEM (Dulbecco’s modified Eagle’s medium) containing 10% (v/v) FBS (fetal bovine serum) and 2 mM L-glutamine. Cells were transfected using Lipofectamine™ 2000 (Invitrogen) and used in experiments after 24 h.

Treatment with compounds

Stock solutions (10 nM) of ES₃₅₅, ES₅, ES₁, cpd A (translocation inhibitor compound A) and PS2 (proteasome inhibitor 2; benzoylcarbonyl-Leu-Leu-Phe-aldehyde; Calbiochem) were made up in DMSO. ES₃₅₅, ES₅ and ES₁ were added to a final concentration of 8 μM, PS2 and cpd A to 10 μM, and CHX (cycloheximide) to 100 μg/ml, then cells were incubated at 37°C.

Metabolic labelling and IP (immunoprecipitation)

Cells were incubated in DMEM lacking methionine and cysteine (Invitrogen) for 20 min, and were labelled with 22 μCi/ml EasyTag [³⁵S]Met/Cys (PerkinElmer) for 40 min at 37°C. Cells were washed twice in PBS and either harvested immediately or chased in DMEM supplemented with 2.5 mM unlabelled methionine/cysteine plus the indicated compounds. Cells were harvested in 50 μl of ice-cold IP buffer (140 mM NaCl, 10 mM Tris/HCl, pH 7.5, 1 mM EDTA and 1% Triton X-100) plus protease inhibitor cocktail (Sigma). For IP, lysates were denatured in 1% SDS, then 5 vol. of IP buffer containing 10 mM non-radioactive methionine/cysteine, 1 mM PMSF and 4% pansorbin (Calbiochem) was added. Samples were rotated for 1 h at 37°C, centrifuged at 15 000 g for 10 min, and the supernatant was incubated overnight at 4°C with an anti-FLAG antibody. Immune complexes were collected on Protein A–Sepharose beads, washed with IP buffer, eluted in reducing SDS/PAGE sample buffer and resolved by SDS/PAGE. Radioactive gels were analysed by FLA-3000 phosphorimaging (Fuji) and quantified using AIDA v3.52 software (Raytest Isotopenmessgerate).

Western blotting

To detect polyubiquitinated material, cells cultured in 12-well dishes were lysed in 50 μl of IP buffer and then denatured in SDS/PAGE sample buffer at 70°C for 10 min. Half of the sample was run on SDS/10% PAGE gels, analysed by blotting with an anti-ubiquitin antibody and then visualized by enhanced chemiluminescence.

XBP1 (X-box-binding protein 1) splicing

Total RNA was extracted using TRizol® reagent (Invitrogen), and first-strand cDNA was synthesized with an oligo-p(dT)₁₅ primer. cDNA was used as a template for PCR using primers flanking the XBP-1 intron (FWD, 5’-ACAGCCTTGGGATGGATG-3’; REV, 5’-TGACTGGTGCCAAGTTGTCC-3’), and PCR products were analysed on 2% agarose gels.

Fluorescence microscopy

HeLa cells incubated with 8 μM ES₁ in the medium for 0–8 h were fixed in methanol for 4 min at −20°C, then probed with primary antibodies against PDI and fluorophore-conjugated secondary antibodies (Molecular Probes). For visualization of preprolactin–mCherry, HeLa cells transfected using PEpTAG reagent (Peqlab), were incubated with 100 μg/ml CHX for 3 h at 37°C. Cells were washed three times for 1 min with 1 ml of PBS, then treated with 8 μM ES₁, 10 μM PS2 or 8 μM ES₁ and 10 μM PS2 for 6 h. Following treatment, cells were washed in PBS and fixed in 3% paraformaldehyde in PBS for 25 min at room temperature (21–23°C). Images were obtained
using an Olympus BX60 upright microscope with a MicroMax cooled charge-coupled device camera (Roper Scientific) driven by Metamorph software (Universal Imaging Corporation).

Subcellular fractionation and EndoH (endoglycosidase H) treatment

Cells in 10 cm dishes were rinsed twice in PBS, harvested by scraping and resuspended in 100 μl of hypotonic buffer (20 mM Heps, pH 7.6, 5 mM KCl, 2.5 mM EDTA and 1 mM PMSF) on ice. Cells were sonicated for three 10 s pulses in a sonicating waterbath. Lysates were centrifuged at 50 000 g for 30 min at 4°C. The membrane pellet was resuspended in 100 μl of 100 mM Na2CO3, pH 11.5, incubated on ice for 20 min, then centrifuged at 100 000 g for 1 h at 4°C. The supernatant was removed, and the pellet was solubilized in 100 μl of IP buffer for 10 min at 4°C, followed by a final centrifugation step at 100 000 g for 1 h at 4°C. The supernatant was removed and the Triton X-100-insoluble pellet was resuspended in IP buffer containing 0.1 % SDS. Opin was immunoprecipitated from the carbonate supernatant, Triton X-100 supernatant and Triton X-100-insoluble fractions, and was resolved by SDS/PAGE. Where indicated, immunoprecipitated material was incubated with 500 units of EndoH (New England Biolabs) for 2 h at 37°C.

RESULTS

ESI induces accumulation of polyubiquitinated proteins and induces ER stress

ES1, ESα and a related compound ESβ, designed to provide a negative control treatment (cf. [9]), were synthesized according to the scheme outlined in Supplementary Figure S1. The inhibitory effect of ES1 on the ERAD pathway was confirmed in two different human cell lines (see Supplementary Figure S2 at http://www.BiochemJ.org/bj/442/bj4420639add.htm). Notably, we found that acute (< 1 h) treatment of cells with ES1 inhibited degradation of the model substrate TCRα (Supplementary Figure S2A), showing that the action of ES1 on ERAD is rapid and does not require prolonged treatment. In contrast, ESα treatment did not reproducibly delay TCRα degradation, and ESβ had no detectable inhibitory effect (Supplementary Figure S2B). ES1 treatment also resulted in the appearance of substantial amounts of higher-molecular-mass forms of the model ERAD substrate (Supplementary Figure S2B). Less pronounced high-molecular-mass species were also visible in cells treated with a proteasome inhibitor (PS2) and at the start of the chase in untreated cells (Supplementary Figure S2B), suggesting that they might represent polyubiquitinated forms of TCRα en route to degradation. Consistent with this interpretation, the total levels of polyubiquitinated proteins detected in cells were also markedly increased upon ES1 treatment (Figure 1A, lane 4, and Supplementary Figure S2C). A similar, although less pronounced, effect was observed following treatment with ESα and PS2 (Figure 1A, lanes 3 and 5), whereas ESβ had no discernable effect (Figure 1A, lane 2). These high-molecular-mass polyubiquitin conjugates accumulated with time, and became apparent after 1 h of exposure to ES1 (Figure 1B). Increased levels of polyubiquitinated material following ES1 treatment have been observed previously and have been proposed to reflect an inhibition of deubiquitination [8].

Consistent with previous work [10], we also found that ES1 induced ER stress in cultured cells. Splicing of Xbp1 mRNA, an early event in the IRE1 (inositol-requiring enzyme 1) branch of the UPR can be measured using RT (reverse transcription)—PCR. A single PCR product, representing the unspliced mRNA, was observed in DMSO-treated cells (Figure 1C, lane 2). A faster-migrating product, representing spliced Xbp1 mRNA, was apparent in cells treated with the reducing agent DTT (dithiothreitol) to perturb oxidative protein folding (Figure 1C, lane 1). ES1 also induced Xbp1 mRNA splicing (Figure 1C, lane 5), showing that treatment with this compound resulted in activation of IRE1. In contrast, no spliced Xbp1 was detected in cells treated with ESα or ESβ (Figure 1C, lanes 3 and 4), consistent with the minor effect of these compounds on ERAD and polyubiquitination (Supplementary Figure S2 and Figure 1A). The response to ES1 did not require prolonged treatment with ES1, and spliced Xbp1 was evident after 4 h (Figure 1D). Activation of the PERK [PKR (double-stranded-RNA-dependent protein kinase)-like ER kinase] branch of the UPR results in phosphorylation of eIF2α (eukaryotic initiation factor 2α), resulting in a decrease in the rate of protein synthesis under conditions of ER stress, as observed following treatment with the reducing agent DTT to perturb oxidative protein folding (Figure 1E, cf. lanes 1 and 2). Phosphorylation of eIF2α occurred rapidly upon ES1 treatment (Figure 1E, bottom two panels), and this effect correlated with a decrease in total protein synthesis (Figure 1E, top panel, lanes 3–8), showing that the PERK branch of the UPR was also activated. Similar effects of ES1 were also observed in HEK (human embryonic kidney)-293 cells (see Supplementary Figure S3 at http://www.BiochemJ.org/bj/442/bj4420639add.htm). Interestingly, ES1 treatment resulted in a more pronounced phosphorylation of eIF2α and greater translational attenuation in HEK cells than in HeLa cells (Supplementary Figure S3), suggesting that sensitivity to ES1 can vary between cell types (cf. [10]).

Our recent work suggests that ES1 delays certain intracellular trafficking pathways [13], and we therefore examined the effect of the ES compounds on cell morphology. Interestingly, large vacuole-like structures developed within 4–8 h of treatment with ES1 (Figure 1F, arrows), and these were also apparent by immunofluorescence microscopy using an antibody specific for ER-localized proteins (Figure 1F, G and H), suggesting that these structures were derived from the ER. ESβ had no obvious effect on cell morphology (Figure 1F), and although no effect of ESα was visible within 8 h (Figure 1F), prolonged treatment (12–16 h) did cause the appearance of large vacuoles (results not shown). A similar swelling of the ER occurs during paraptosis [14], a non-apoptotic form of cell death, and has also been observed under conditions of chronic ER stress [15,16]. Hence these changes in ER structure may reflect ES1-induced ER stress and/or commitment to cell death.

Inhibition of translocation promotes ER stress

The mechanisms through which ES1 induces ER stress are not known, although inhibition of ERAD and/or perturbation of ubiquitin homoeostasis could potentially play a role [10]. Indeed, inhibition of the proteasome is able to induce activation of the UPR in a variety of cell types (e.g. [17]). Interestingly, however, treatment of HeLa cells with PS2 for 8 h failed to induce Xbp1 mRNA splicing (Figure 2A, lane 3), despite causing a clear build-up of polyubiquitinated species and profound inhibition of ERAD (Figure 1A and Supplementary Figure S2). Similarly, Xbp1 splicing was not observed in cells treated with an alternative proteasome inhibitor MG132 (Figure 2A, lane 4), or in response to a small molecule that inhibits protein deubiquitination (N5C687852) (Figure 2A, lane 5), and causes a pronounced accumulation of polyubiquitin adducts [18,19] (see Supplementary Figure S4A at http://www.BiochemJ.org/bj/442/bj4420639add.htm). Thus
neither blockade of ERAD nor perturbation of global ubiquitin homeostasis (using a variety of different inhibitors) activated the UPR in HeLa cells over this time course. In contrast, parallel treatment with ES_{I} induced a robust ER stress response (Figure 2A, lane 2). These results are significant as they suggest that ES_{I} does not induce ER stress solely by blocking degradation of ERAD substrates or disrupting the UPS [10]. We therefore considered the alternative possibility that the ability of ES_{I} to activate UPR was related to its ability to perturb protein translocation [12]. In order to test this hypothesis, we used a second small-molecule inhibitor of co-translational translocation (referred to in the present study as cpd A), derived from a naturally occurring cyclic depsipeptide [20]. Like ES_{I}, the cyclodepsipeptide inhibitors cause a wide-ranging inhibition of translocation when used at low micromolar concentrations [21,22], but are structurally distinct and target a different stage of Sec61-mediated protein translocation across the ER membrane [20,23]. Strikingly, treatment of HeLa cells with this translocation inhibitor also induced Xbp1 mRNA splicing on a timescale comparable with ES_{I} treatment (cf. Figures 1C and 2B). In addition, cpd A induced phosphorylation of eIF2α and reduced the rate of translation (Figures 2C and 2D), demonstrating that both the IRE1 and PERK branches of the UPR were activated. This effect of the cyclodepsipeptide inhibitors has not previously been reported and supports the view that inhibiting Sec61-mediated translocation promotes ER stress. This effect could potentially be caused by the reduced biogenesis of specific ER factors such as chaperones, some of which are known to be relatively short-lived [24,25]. However, the global inhibition of protein synthesis using CHX did not induce Xbp1 splicing (Figure 2E, lane 5), suggesting that a lack of ER components resulting from a defect in co-translational translocation is unlikely to underlie the ability of ES_{I} and cpd A to activate the UPR. These observations raised the possibility that the induction of ER stress by these translocation inhibitors might instead be related to the production of non-translocated forms of membrane and secretory proteins. In order to address this issue further, we examined the fate of various ER-targeted proteins in the presence of ES_{I}. 

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Addition of N-linked glycans to polypeptides occurs within the ER lumen and was thus used to provide a readout of TCRα insertion into the ER (cf. [12,22]). In DMSO- and ES_{100}5-treated cells, FLAG–TCRα was efficiently N-glycosylated and migrated as a species of approximately 48 kDa (Figure 3A, lanes 1 and 2). As observed previously [12], levels of the N-glycosylated form of TCRα were reduced in ES_{10}5-treated cells (Figure 3A, lane 4). In addition, a more rapidly migrating species of the size expected for the unglycosylated protein was found to accumulate after 8 h of treatment with ES_{1} (Figure 3A, lane 4). A similar, although less pronounced, effect was seen upon ES_{0.5} treatment (Figure 3A, lane 3). These observations indicate that ES_{1} treatment results in the production of a form of TCRα lacking N-glycosylation.

We next examined the effect of ES, on the subcellular localization of a chimaeric secretory protein composed of fluorescent mCherry fused to the ER-targeting sequence of preprolactin (ppl–mCherry). In untreated cells, ppl–mCherry was predominantly localized to the secretory pathway, as demonstrated by the reticular and ribbon-like pattern of fluorescence, typical of the ER and Golgi apparatus respectively (Figure 3B, panel A). Cells were then treated with CHX to prevent further protein synthesis and allow existing protein to exit the ER. After washing to remove CHX, cells were incubated with ES_{1}, PS2 or a combination of ES_{0.5} and PS2 for a further 6 h, prior to imaging. This treatment regime (outlined in Figure 3B) allowed us to determine the localization of ppl–mCherry synthesized in the presence of ES_{1}, ppl–mCherry fluorescence was considerably lower after 3 h of CHX treatment, confirming that much of the protein had been secreted from the cells (Figure 3B, panel B).

In cells treated with PS2 alone following CHX treatment, the ppl–mCherry distribution was similar to that in untreated cells (Figure 3B, cf. panels A and C), suggesting that the majority of the protein synthesized under conditions of proteasome inhibition was correctly targeted to the ER. A different pattern of fluorescence was seen in cells treated with ES_{1} (Figure 3B, panel D). Under these conditions, the reticular distribution of ppl–mCherry was replaced by more diffuse fluorescence distributed throughout the cell, suggesting that a proportion of the newly synthesized protein had not entered the ER, but was instead located in the cytosol. A variable amount of mCherry fluorescent protein was also observed in punctate structures adjacent to the nucleus (Figure 3B, panels D and E). Co-treatment with PS2 resulted in a much clearer accumulation of fluorescent protein in the cytosol (Figure 3B, panel E), suggesting that ppl–mCherry which failed to enter the ER in the presence of ES_{1} was degraded via the proteasome.

To validate these data, we took a complementary biochemical approach to explore the fate of a polytopic membrane protein synthesized in the presence of ES_{1}. Membranes isolated from HeLa cells transiently expressing opsin were extracted with sodium carbonate, pH>11, to remove peripherally associated proteins [26]. Membrane-integrated proteins are resistant to such treatment, and were re-isolated by centrifugation, then solubilized in Triton X-100. This method allowed separation of the ER luminal chaperone PDI (Figure 3C, lane 3), from the integral ER membrane protein derlin-1, which remained in the membrane-associated pellet after carbonate extraction (Figure 3C, lanes 2 and 3), confirming that this treatment effectively strips non-integral proteins from the ER membrane. Next, cells treated with DMSO, ES_{1}, ES_{0.5} or ES_{0.25} for 1 h were pulse-labelled in the continued presence of the compounds plus PS2, prior to carbonate extraction and opsin IP. In untreated cells, the large majority of the radiolabelled opsin was present in the carbonate-resistant pellet (Figure 3D, lanes 2 and 3, DMSO), demonstrating that it was stably integrated into the ER membrane. A smaller proportion of the radiolabelled non-glycosylated opsin was extracted in the carbonate supernatant (Figure 3C, lane 3), confirming that this treatment effectively strips non-integral proteins from the ER membrane.

In the presence of ES_{1}, however, radiolabelled opsin was almost exclusively found in the carbonate-resistant pellet and was non-glycosylated (Figure 3D, lane 1, ES_{1}). Virtually no radiolabelled opsin could be detected in the sodium carbonate-extracted pellet and there was no indication of any glycosylated protein chains (Figure 3D, lanes 2 and 3, ES_{1}). It is clear that ES_{1} inhibited the integration of opsin into the ER membrane, leading to production of non-glycosylated protein. Notably, the population of non-glycosylated opsin was only readily apparent in HeLa cells when PS2 was included during the pulse-labelling period (Figure 3E, cf. lanes 5 and 9), showing
that proteins prevented from entering the ER upon ES treatment are normally subjected to proteasomal degradation.

Taken together, these results support a model whereby an inhibition of translocation by ES leads to the production of mislocalized membrane and secretory proteins in cultured cells. The non-translocated forms of ppl–mCherry and opsin that failed to enter the ER in the presence of ES were degraded via the proteasome, suggesting that such polypeptides are targeted to the UPS. We therefore postulated that the polyubiquitinated species observed in ES-treated cells (Figure 1A; see also [8]), may include a variety of polypeptides that had failed to translocate correctly across the ER membrane and were en route to proteasomal degradation. Indeed, treatment of cells with ES, which has only a minor effect on protein translocation [12], caused a less pronounced accumulation of polyubiquitinated material (Figure 1A and Supplementary Figure 2C). Furthermore, we found that treatment with the cyclodepsipeptide translocation inhibitor cpd A also resulted in some accumulation of high-molecular-mass polyubiquitinated material (Figure 4A, lane 3). This previously unreported effect of cpd A provides additional evidence that inhibition of ER translocation in live cells promotes a build-up of polyubiquitin conjugates. Significantly, co-treatment of cells with CHX completely prevented the accumulation of polyubiquitin adducts caused by both cpd A and ES (Figure 4A, lane 4, and Figure 4B, lanes 4 and 5). This requirement for ongoing protein synthesis suggests that these species represent proteins newly synthesized in the presence of the translocation inhibitors.

In contrast, PS2 still caused accumulation of polyubiquitinated proteins in the presence of CHX (Figure 4B, lanes 8 and 9), suggesting that a significant proportion of these adducts were derived from pre-existing proteins. We also found that treatment with ES led to the appearance of higher-molecular-mass forms of EDEM-1 (ER degradation-enhancing α-mannosidase-like 1), a model ER-resident protein (Figure 4C, lane 2). These species migrated as diffuse bands near the top of the polyacrylamide gel, comparable with the polyubiquitin conjugates detected with anti-ubiquitin antibodies (Figure 4B). However, they were not observed in cells treated with PS2 (Figure 4C, lane 3), showing that they were not generated solely as a consequence of inhibiting proteasomal degradation. Accumulation of these higher-molecular-mass forms of EDEM-1 was prevented by cotreatment with CHX (Figure 4C, lane 6), showing that they represent polypeptides synthesized in the presence of ES, and not an ES-dependent modification of a pre-existing pool of EDEM-1 (cf. [27]). On the basis of these results, we conclude that the polyubiquitinated species induced by ES and cpd A most probably include non-translocated membrane and secretory proteins targeted for proteasomal degradation. Supporting this interpretation, endogenous ES-induced polyubiquitin conjugates were eliminated by cells following addition of CHX and a subsequent chase period (Figure 4D). However, since PS2 rapidly induces accumulation of polyubiquitinated proteins even in the presence of CHX (Supplementary Figure S4B), it was not possible to demonstrate that the loss of ES-induced
polyubiquitin conjugates was proteasome-dependent. Therefore we cannot exclude the possibility that this effect is the result of deubiquitination of the substrates rather than degradation.

Inhibition of protein translocation contributes to the cellular actions of ESI

Non-translocated membrane and secretory proteins have potentially harmful properties, such as hydrophobic transmembrane domains and/or uncleaved signal peptides. Therefore we wondered whether the accumulation of such aberrant polypeptides could contribute to the cellular effects of ESI and cpd A. To address this issue, we exploited the observation that non-translocated polypeptides accumulated to a greater extent when proteasome activity is inhibited (Figure 3), whereas their production is effectively blocked by CHX (Figure 4). This allowed us to examine whether the ability of ESI and cpd A to induce ER stress was altered by conditions that promoted or inhibited the accumulation of non-translocated polypeptides. When protein synthesis was prevented by co-treatment with CHX, both Xbp1 mRNA splicing and the ER vacuolization induced by ESI were completely abolished (Figure 5A, lanes 5 and 7, and Supplementary Figure 4C). Similarly, CHX also substantially diminished Xbp1 splicing in response to cpd A (Figure 5B, lanes 3 and 5). In contrast, the ability of DTT to induce Xbp1 splicing was not prevented by CHX (Figure 5A, lanes 3 and 4), demonstrating that activation of this pathway is not dependent on the continued import of cargo proteins into the ER, nor mechanistically on protein synthesis itself. Thus activation of the UPR by both ESI and cpd A requires ongoing translation. This is an important finding as it suggests that polypeptides synthesized in the presence of these compounds are responsible for their ability to induce ER stress. In contrast, combinatorial treatment with ESI and proteasome inhibitors more effectively induced Xbp1 splicing (Figure 5C, lanes 5 and 6, and Supplementary Figure S4D) and enhanced ER vacuolization (Supplementary Figure S4E) when compared with ESI alone. The ability of cpd A to induce Xbp1 splicing was also increased by proteasome inhibition (Figure 5B, lanes 3 and 4). Thus conditions that inhibit the efficient degradation of non-translocated proteins (Figure 3) enhance the ability of ESI and cpd A to activate the UPR, supporting the hypothesis that accumulation of polyubiquitinated non-translocated proteins promotes activation of the UPR. Consistent with this interpretation, the accumulation of polyubiquitinated proteins in ESI-treated cells correlated well with the induction of ER stress (cf. Figures 1B and 1E).

On the basis of these findings, we propose a working model for the cellular actions of ESI. In this model (Figure 6), a proportion of nascent membrane and secretory proteins fail to enter the ER in the presence of ESI, or cpd A, and are instead polyubiquitinated...
Figure 5  Correlation between accumulation of mistranslocated polypeptides and induction of ER stress

(A) HeLa cells were treated with 100 μg/ml CHX for 8 h, 2 mM DTT and CHX for 2 h or 8 h, or with 8 μM ES1 or 10 μM PS2 in the presence or absence of 100 μg/ml CHX for 8 h. Xbp1 splicing was determined by RT–PCR. PCR products corresponding to unspliced (u) and spliced (s) Xbp1 mRNA are indicated. (B) HeLa cells were treated with DMSO, 2 mM DTT, or the indicated combination of 10 μM translocation inhibitor cpd A, 10 μM PS2 and 100 μg/ml CHX for 8 h. Xbp1 splicing was determined by RT–PCR. (C) HeLa cells were incubated with DMSO, 8 μM ES1, 10 μM PS2, 10 μM MG132, 8 μM ES1 and 10 μM PS2, or 8 μM ES1 and 10 μM PS2 for 8 h. Xbp1 splicing was determined by RT–PCR.

Figure 6  Proposed model for cellular actions of ES1

Proteins possessing an ER signal sequence are targeted to the ER and translocated through the Sec61 translocon (1). Properly folded proteins exit the ER and move along the secretory pathway. Misfolded proteins are retained by the ER quality control machinery (ER-QC) (2), handed over to the retrotranslocation machinery, polyubiquitinated and retrotranslocated (2). The retrotranslocated protein is deubiquitinated and deglycosylated (3) before being degraded by the proteasome. Accumulation of misfolded proteins in the ER lumen activates UPR signalling. ES1 inhibits translocation of proteins across the ER membrane, leading to the production of non-translocated proteins (NTP) at the cytosolic face of the ER membrane. These aberrant species are degraded by the UPS and also activate the UPR through an unknown mechanism. ES1 may also inhibit one or more steps of the ERAD pathway, including retrotranslocation or deubiquitination. Also shown are the points of action of CHX, PS2 and NSC687852 (NSC), a deubiquitinase inhibitor that induces accumulation of polyubiquitin in cells [18].

and may be subjected to proteasomal degradation. Since ES1 and cpd A only activate Xbp1 splicing if protein synthesis is ongoing, we suggest that accumulation of these non-translocated proteins in the cytosol contributes to activation of the ER stress response.

DISCUSSION

ER protein folding and degradation pathways have emerged as exciting targets for the treatment of a diverse array of diseases, from cancer and stroke to classical protein folding diseases. As a result, there is much interest in the therapeutic potential of selective small molecules to modulate ER protein folding homeostasis and the ER stress response. In the present study we describe the preparative synthesis and cellular activity of the ERAD inhibitor ES1 and two related compounds ES2 and ES35. We confirm that ES1 inhibits ERAD of TCRα, causes accumulation of polyubiquitinated proteins, and rapidly induces ER stress. We also show that ES1 reduces the efficiency of protein translocation at the ER in cultured cells, and provide evidence that membrane and secretory proteins that fail to enter the ER in the presence of ES1 are subjected to proteasomal degradation. Using a second small-molecule inhibitor of cotranslational translocation, we examine the consequences of reduced translocation efficiency in cultured cells. Intriguingly, our results suggest that the accumulation of non-translocated membrane and secretory proteins promotes activation of the ER stress response.

Originally identified as an ERAD inhibitor, ES1 was subsequently found to impair Sec61-mediated protein translocation at the ER for a range of substrates [12]. In the present study, we have employed different techniques to show that ES1 also perturbs the entry of a variety of proteins into the ER of living cells.
Thus the N-glycosylation of TCRα, the membrane integration and N-glycosylation of opsin, and the ER localization of ppl–mCherry were all impaired by treatment of cells with ESI. In the presence of ESI, non-glycosylated TCRα, non-integrated opsin and non-ER-localized ppl–mCherry continued to be synthesized, and could be observed in ESI-treated cells. However, the amount of these non-translocated species that accumulated in cells following ESI treatment was somewhat variable. These variations may reflect differences in the response of different cell types to ESI treatment, for example greater translational attenuation in HEK-293 cells, but could also be due to differences in the degree of translocation inhibition caused by ESI. Indeed, non-translocated forms of membrane proteins, including TCRα and MHC class I heavy chain, were not observed in previous studies [7,9], suggesting that ESI-mediated inhibition of translocation is incomplete and/or can be influenced by experimental variables such as the duration of drug treatment.

The non-translocated species we observed in ESI-treated cells were labile, and inhibitor studies indicate that they are degraded by the proteasome. Similarly, the cytosolic form of the VCAM1 (vascular cell adhesion molecule 1) precursor generated in the presence of translocation inhibitor cpd A is also normally degraded via a proteasome-dependent pathway [20]. Other non-translocated proteins generated under various conditions of reduced translocation efficiency are also rapidly removed by the proteasome [22,28,29]. Taken together, these observations strongly support the existence of an efficient cellular quality control system that functions to eliminate proteins that fail to be properly translocated across the ER membrane. Indeed, recent work has begun to identify the cytosolic machinery involved in such a pathway [30]. Non-translocated species will typically expose hydrophobic transmembrane domains and/or uncleaved signal peptides, and it is easy to envisage that their accumulation in the cytosol might be particularly harmful. Indeed, aberrant interactions of non-translocated prion protein have been proposed to contribute to neurodegeneration in prion diseases [31,32]. Our results suggest that non-translocated polypeptides may also participate in cellular dysfunction by promoting ER stress. Thus we find that both ESI and another small-molecule inhibitor of co-translational translocation induce accumulation of polyubiquitinated proteins and activate the UPR. Strikingly, co-treatment with CHX, which blocks accumulation of polyubiquitin adducts, also abolishes the ER stress response induced by these compounds. This requirement for ongoing translation suggests that it is actually the polyubiquitinated non-translocated polypeptides synthesized in the presence of the inhibitors that promote activation of the UPR. In support of this model, the ability of both ESI and cpd A to induce ER stress is enhanced under conditions of proteasome inhibition that prevent efficient degradation of non-translocated proteins. Our working model (Figure 6) is consistent with the known actions of ESI and cpd A, and would also explain the previously observed synergy between ESI and the proteasome inhibitor bortezomib [10]. Although we cannot formally exclude the possibility that ESI and cpd A activate the UPR by ‘off-target’ (ie. non-translocon) effects, the facts that they are structurally distinct and inhibit different stages of co-translational translocation suggest that this is unlikely. Furthermore, the observation that CHX prevents UPR activation in response to ESI and cpd A, but does not affect DTT-induced UPR, suggests that the translocation inhibitors do not induce ER stress through a direct effect on ER chaperones, ERAD machinery or UPR components [7,27,33]. In addition to perturbing ER translocation, ESI also inhibits p97 and an associated deubiquitinase activity [9,10]. Interfering with p97 function is known to inhibit ERAD and can also induce swelling of the ER (cf. [34]). It thus seems likely that reduced p97 function may also contribute to the pronounced ER stress phenotype induced by ESI. However, the ability of a second small-molecule inhibitor of the Sec61 translocon to recapitulate many features of ESI-induced UPR activation supports a model whereby products of failed translocation promote ER stress.

Since the UPR is typically triggered by the presence of misfolded proteins in the ER lumen, it is not immediately obvious how the generation of non-translocated polypeptides in the cytosol might promote this pathway (Figure 6). One potential explanation is that polyubiquitination of these species reduces the availability of free ubiquitin, thereby indirectly inhibiting removal of misfolded proteins from the ER and inducing the UPR. This would be consistent with the suggestion that inhibition of deubiquitination by ESI causes a global ubiquitin deficiency [10]. However, several distinct inhibitors of the UPS, namely PS2, MG132 and N6C87852, failed to induce Xbp1 mRNA splicing despite causing comparable accumulation of polyubiquitinated material. On this basis, we believe that perturbation of ubiquitin homeostasis is not the underlying cause of ESI-induced UPR. Non-translocated proteins could also disrupt protein homeostasis in the cytosol, for example by sequestering molecular chaperones. Cytosolic chaperones such as Hsp70 (heat-shock protein 70) play an important role in the biogenesis of some membrane proteins [35], and therefore interfering with their function may perturb folding at the ER and promote ER stress. Alternatively, the accumulation of non-translocated polypeptides, which have a high propensity to aggregate and initiate non-native interactions, at the cytosolic face of the ER could potentially interfere with the removal of misfolded proteins from the ER. This would result in a build-up of misfolded proteins in the lumen and activation of the UPR, and is consistent with the suggestion that ESI inhibits retrotranslocation [7]. Indeed, certain misfolded cytosolic proteins have been shown to engage in inappropriate interactions with cytosolically oriented components of the ERAD machinery, thereby inhibiting retrotranslocation and activating the UPR [36,37]. We speculate that perturbation of such an early stage in the ERAD process may have a particularly severe effect on levels of misfolded proteins in the ER lumen and hence ER stress. In contrast, inhibition of the proteasome does not always block retrotranslocation, and several ERAD substrates can still exit the ER when the proteasome is inhibited [38–41]. Alternatively, non-translocated polypeptides may also activate UPR signalling through some other, as yet undefined, mechanism [33,42].

The possibility that modulating protein translocation at the ER might have therapeutic potential is further supported by the finding that apratoxin A, a cyclodepsipeptide with anticancer properties [43], inhibits co-translational translocation [44]. In this case, depletion of cancer-associated transmembrane receptors was proposed to underlie the cytotoxic activity of apratoxin A towards cancer cells [44]. Thus the results of the present study further extend the prospect that inhibitors of ER translocation may offer novel therapeutic opportunities in the future, and underline the potential for using small-molecule inhibitors to understand the consequence of defects in membrane translocation events at the cellular level [45].

AUTHOR CONTRIBUTION

Eileithyia Swanton and Stephen High conceived and supervised the project. Eileithyia Swanton wrote the paper. Craig McKibbin, Alina Mares, Peristera Roboti, Marjo Paumalainen and Karolina Lesiak-Mleczkowska performed the experiments, analysed the data, and contributed to preparation of the Figures and paper. Anna Callan performed pilot experiments. Michela Piacenti and Helen Williams performed chemical syntheses and wrote synthetic protocols. Roger Whitehead and Sabine Flitsch conceived and supervised
the synthetic chemistry. Hanna Harant and Stig Linder provided reagents and contributed to analysis and interpretation of the results.

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SUPPLEMENTARY ONLINE DATA

Inhibition of protein translocation at the endoplasmic reticulum promotes activation of the unfolded protein response

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EXPERIMENTAL

Synthetic protocols

The synthesis of ES1 and ES2 commences with nitroso-compound 1, which was prepared according to a previous procedure by reaction of isobutene with nitrosyl chloride (generated in situ from isoamyl nitrite and HCl) [1]. Displacement of the chlorine atom in 1 by the amino group of methyl glycinate then smoothly proceeded smoothly to give oximino ester 2 at a very good yield [2]. Exposure of 2 to two equivalents of its corresponding isocyanate gave the bis-adducts [3] 3 and 4, which were subsequently converted into their acyl hydrazides 5 and 6 by treatment with an aqueous methanolic solution of hydrazine. Finally, coupling to commercially available (E)-3-(5-nitro-2-furyl)acrylaldehyde gave ES1 and ES2 (7 and 8 respectively).

General experimental

Synthetic reagents were purchased from Sigma–Aldrich or Alfa Aesar and used as supplied. Solvents were dried and distilled prior to use. IR spectra were recorded on a PerkinElmer 881 spectrometer, an AT1-Matson Genesis Series FTIR (Fourier-transform infrared) spectrometer or a PerkinElmer Spectrum BX FTIR spectrometer. 1H and 13C spectra were recorded on a Bruker Avance II 500 MHz spectrometer, a Bruker Avance III 400 MHz spectrometer, an AT1-Matson Genesis Series FTIR (Fourier-transform infrared) spectrometer or a PerkinElmer Spectrum BX FTIR spectrometer. Mass spectra were recorded on a Micromass Platform II (electrospray) spectrometer. Melting points were recorded using a Sanyo Kofler Block or a Buchi B-545 Capillary Hot Stage.

Methyl 2-[3-(4-chlorophenyl)-4-[3-(4-chlorophenyl)-1-hydroxyureido]-5,5-dimethyl-2-oxoimidazolidin-1-yl]acetate (3)

A solution of 4-chlorophenyl isocyanate (1.01 g, 5.75 mmol) in freshly distilled tetrahydrofuran (2 ml) was added under an atmosphere of nitrogen to a solution of oxime (2) (0.57 g, 2.87 mmol) in tetrahydrofuran (14 ml). The resulting solution was stirred at room temperature for 18 h and then concentrated under reduced pressure. The product was crystallized from dichloromethane, collected by filtration, washed with diethyl ether and dried. The filtrate was concentrated under reduced pressure and the crystallization procedure was repeated twice. Combination of the batches of crystallized product gave the title compound as a colourless solid (1.03 g, 65%). Mp 86–86.9°C; νmax (film)/cm−1: 3081 (m), 2977 (m), 2874 (m), 2798 (m), 1750 (s); δH (CDCl3: 500 MHz) 7.28 (3H, s, CH3; 3.73 (3H, s, CO2CH3); 3.39 (2H, s, NCH2CO2CH3); 1.27 (6H, s, C(CH3)2); δC (CDCl3: 300 MHz) 7.28 (1H, s, CH = NOH), 3.73 (3H, s, CO2CH3), 3.39 (2H, s, NCH2CO2CH3), 1.27 (6H, s, C(CH3)2); δC (CDCl3: 125 MHz) 172.7 (CO2CH3), 154.8 (CH = NOH), 54.1 [C(CH3)2], 52.0 (CO2CH3), 44.7 (NCH2CO2CH3), 25.2 [C(CH3)2]; m/z (positive ion electrospray) 197 ([M + Na]+, 100%); (found 197.0899, C9H12N2NaO3, [M + Na]+, requires 197.0897).

Methyl 2-[4-(3-(4-chlorophenyl)-4-[3-(4-chlorophenyl)-1-hydroxyureido]-5,5-dimethyl-2-oxoimidazolidin-1-yl]acetate (4)

Glycine methyl ester hydrochloride (5.17 g, 41 mmol) was added, under an atmosphere of nitrogen, to a solution of 2-chloro-2-methyl-1-nitrosopropane (1) (5.0 g, 41 mmol) in freshly distilled acetonitrile (80 ml). Triethylamine (8.3 ml, 82 mmol) was then added dropwise. The reaction mixture was stirred at room temperature for 18 h and then quenched by the addition of a saturated aqueous solution of sodium bicarbonate (40 ml). Organic material was extracted into dichloromethane (4×50 ml) and the combined organic layers were then washed with brine (40 ml), dried over magnesium sulfate and concentrated under reduced pressure to yield the title compound as a pale yellow solid (5.75 g, 88%). Mp 86–86.9°C; νmax (film)/cm−1: 3292 (m), 3163 (m), 3081 (m), 2977 (m), 2874 (m), 2798 (m), 1750 (s); δH (CDCl3: 300 MHz) 7.38 (1H, s, CH = NOH), 3.73 (3H, s, CO2CH3), 3.39 (2H, s, NCH2CO2CH3), 1.27 (6H, s, C(CH3)2); δC (CDCl3: 125 MHz) 172.7 (CO2CH3), 154.8 (CH = NOH), 54.1 [C(CH3)2], 52.0 (CO2CH3), 44.7 (NCH2CO2CH3), 25.2 [C(CH3)2]; m/z (positive ion electrospray) 197 ([M + Na]+, 100%); (found 197.0899, C9H12N2NaO3, [M + Na]+, requires 197.0897).

3-(4-Chlorophenyl)-1-[3-(4-chlorophenyl)-1-[2-hydrazinyl-2-oxo-oxoethyl]-5,5-dimethyl-2-oxoimidazolidin-4-yl]-1-hydroxyurea (5, ES2)

An aqueous solution of hydrazine (62%, 1.18 ml, 24 mmol) was added to a solution of imidazolidinone (3) (0.567 g, 1.18 mmol)
in methanol (6 ml). The reaction mixture was stirred at room temperature for 48 h and then cooled to 0 °C. The resulting precipitate was collected by filtration, washed with cold methanol (10 ml) and diethyl ether (10 ml) and dried to yield the title compound as a colourless solid (0.358 g, 63%). Mp 197.5–199 °C; \( \nu_{max} \) (solid state)/cm\(^{-1} \) 3304 (m), 2058 (s), 1928 (s), 1729 (s), 1690 (s), 1469 (s), 1387 (s), 1252 (s), 1227 (s), 1209 (s), 1125 (s), 958 (s), 836 (s), 721 (s), 607 (s), 501 (m), 311 (w), requires 660.2212).

\[ \delta_{H} (CDCl_{3}; 500 MHz) \]

- 8.17 (1H, brs, NH or OH), 8.08 (1H, brd, J = 7.9 Hz, Ar-CH), 7.99 (1H, brs, NH or OH), 7.92 (1H, brd, J = 6.7 Hz, Ar-CH), 7.87–7.78 (4H, m, 4 × Ar-CH), 7.61 – 7.59 (2H, m, 2 × Ar-CH), 7.55–7.52 (1H, m, Ar-CH), 7.50–7.44 (5H, m, 5 × Ar-CH), 7.35–7.32 (1H, m, Ar-CH), 6.19 (1H, s, NCHN), 4.69 (1H, d, J = 18.2 Hz, CH\(_2\)CO\(_2\)CH\(_3\)), 3.88 (3H, s, CO\(_2\)CH\(_3\)), 3.79 (1H, d, J = 18.2 Hz, CH\(_2\)CO\(_2\)CH\(_3\)), 1.71 (3H, s, CH\(_3\)CH\(_2\)), 1.49 (3H, s, CH\(_3\)CH\(_3\)).

\[ m/z \] (negative ion electrospray) 511 ([M – H]\(^-\), 100%); (positive ion electrospray) 535 ([M + Na\(^+\)], 100%); (found 535.1949, C\(_{24}\)H\(_{24}\)N\(_4\)O\(_4\)Na ([M + H\(^+\)]\(^-\)), requires 535.1952).

1-[(2-Hydrizyl-2-oxoethy)-5,5-dimethyl-3-(naphthalen-1-yl)-2-oxoimidazolidin-4-yl]-1-hydroxy-3-(naphthalen-1-yl)urea (6)

An aqueous solution of hydrazine (62%, 0.88 ml, 23 mmol) was added to a solution of imidazolidinone, 4 (0.451 g, 0.88 mmol) in methanol (4.5 ml). The reaction mixture was stirred at room temperature for 48 h and then cooled to 0 °C. The resulting precipitated solid was collected by filtration and washed with cold methanol (10 ml) and diethyl ether (10 ml). The filtrate was concentrated under reduced pressure and the residual solid was recrystallized at room temperature from dichloromethane/diethyl ether (1:1), washed with cold methanol and diethyl ether and then combined with the first batch of precipitate to yield the title compound as a colourless solid (0.248 g, 55%). Mp 195–197°C; \( \nu_{max} \) (film)/cm\(^{-1} \) 3390 (m), 3271 (brm), 3054 (m), 2971 (m), 2857 (brw), 1686 (s), 1596 (m), 1520 (m), 1495 (s); \( \delta_{H} \) (CD,OD; 500 MHz) 7.12 (1H, brs, NH or OH), 7.83–7.75 (3H, m, 3 × Ar-CH), 7.75 (1H, d, J = 8.5 Hz, 2 × Ar-CH), 7.57–7.52 (1H, m, Ar-CH), 7.48–7.38 (5H, m, 5H), 6.28 (3H, s, CH\(_3\)CO\(_2\)CH\(_3\)), 1.83 (3H, s, CH\(_3\)CH\(_2\)).

\[ m/z \] (negative ion electrospray) 511 ([M – H]\(^-\), 100%); (positive ion electrospray) 535 ([M + Na\(^+\)], 100%); (found 535.2065, C\(_{24}\)H\(_{24}\)N\(_4\)O\(_4\)Na ([M + Na\(^+\)]\(^-\)), requires 535.2064).

Methyl 2-[(4-1-hydroxy-3-(naphthalen-1-yl)ureido)-5,5-dimethyl-3-(naphthalen-1-yl)-2-oxoimidazolidin-1-yl)acetate (4)

A solution of 1-naphthyl isocyanate (1.01 g, 5.77 mmol) in freshly distilled tetrahydrofuran (2 ml) was added under an atmosphere of nitrogen to a solution of oxime (2) (0.57 g, 2.87 mmol) also in freshly distilled tetrahydrofuran (14 ml). The reaction mixture was stirred at room temperature for 15 h, when residual solvent was removed under reduced pressure. The product was crystallized from dichloromethane, collected by filtration, washed with diethyl ether (20 ml) and dried under vacuum. The filtrate was concentrated under reduced pressure and the crystallization procedure was repeated a further three times. Combination of the crystallized material yielded the title compound as a colourless solid (0.921 g, 61%). Mp 199–200 °C; \( \nu_{max} \) (solid state)/cm\(^{-1} \) 3436 (w), 3054 (brw), 2839 (brw), 1749 (s), 1658 (s), 1535 (s); \( \delta_{H} \) (CDCl\(_3\); 500 MHz) 8.17 (1H, brs, NH or OH), 8.08 (1H, brd, J = 7.9 Hz, Ar-CH), 7.99 (1H, brs, NH or OH), 7.92 (1H, brd, J = 6.7 Hz, Ar-CH), 7.87–7.78 (4H, m, 4 × Ar-CH), 7.61 – 7.59 (2H, m, 2 × Ar-CH), 7.55–7.52 (1H, m, Ar-CH), 7.50–7.44 (5H, m, 5 × Ar-CH), 7.35–7.32 (1H, m, Ar-CH), 6.19 (1H, s, NCHN), 4.69 (1H, d, J = 18.2 Hz, CH\(_2\)CO\(_2\)CH\(_3\)), 3.88 (3H, s, CO\(_2\)CH\(_3\)), 3.79 (1H, d, J = 18.2 Hz, CH\(_2\)CO\(_2\)CH\(_3\)), 1.71 (3H, s, CH\(_3\)CH\(_2\)), 1.49 (3H, s, CH\(_3\)CH\(_3\)).

\[ m/z \] (negative ion electrospray) 660 ([M – H]\(^-\), 100%); (positive ion electrospray) 684 ([M + Na\(^+\)], 100%); (found 660.2199, C\(_{24}\)H\(_{24}\)N\(_4\)O\(_4\)Na ([M – H]\(^-\)), requires 660.2212).
Inhibition of translocation promotes the UPR

Figure S1 Synthesis of ESs

Scheme illustrating the synthesis of ES, ESI, ESII and ESR35.

Figure S2 ES inhibits ERAD

(A) HeLa cells transiently expressing FLAG–TCRα were pulse-labelled with [35S]Met/Cys for 40 min then chased in the presence of DMSO or 8 μM ESR35, ESI or ESII. FLAG–TCRα was immunoprecipitated and analysed by SDS/PAGE and phosphorimaging. Glycosylated (TCRα,CHO) and unglycosylated (TCRα) FLAG–TCRα are indicated. Since cells were only exposed to the compounds after pulse-labeling, the FLAG–TCRα was predominantly in the glycosylated form in each case. (B) HEK cells transiently expressing FLAG–TCRα were treated with DMSO, 8 μM ES, 10 μM PS2 for 8 h, then incubated with 100 μg/ml CHX for 0–3 h. Lysates were analysed by Western blotting with an anti-FLAG antibody. (C) HEK cells were treated for 8 h with DMSO, 8 μM ESR35, ESI or ESII, or 10 μM PS2, and lysates were analysed by blotting with an anti-ubiquitin antibody. IB, immunoblot.
Figure S3  ES$_i$ activates the UPR in HEK cells

(A) HEK-293 cells were treated with 10 mM DTT for 30 min or 8 $\mu$M ES$_i$ for 0–6 h, then were labelled with [35S]Met/Cys for 40 min in the presence or absence of the respective compound. The total time of exposure to ES$_i$ is indicated above each lane. Lysates were analysed by SDS/PAGE and phosphorimaging (top panels), or by blotting with anti-peIF2$\alpha$ or anti-eIF2$\alpha$ antibodies (bottom two panels). (B) HEK-293 cells were left untreated (NT), or were treated with 2 mM DTT for 2 h or 8 $\mu$M ES$_i$ for the time indicated. Xbp1 mRNA splicing was determined by RT–PCR. Unspliced (u) and spliced (s) Xbp1 mRNA are indicated.

Figure S4  Relationship between ES$_i$ and the UPS

(A) HeLa cells were treated with 8 $\mu$M ES$_{355}$ or ES$_i$ or 0.4 $\mu$M NSC687852 (NSC) for 8 h, and lysates were analysed by blotting with an anti-ubiquitin (Ub) antibody. (B) HeLa cells were treated with DMSO for 4 h, 8 $\mu$M ES$_i$ or 10 $\mu$M PS2 with or without 100 $\mu$g/ml CHX for the times indicated, and lysates were analysed by blotting with the anti-ubiquitin antibody. N.B. Lanes 1–4 of this experiment are also shown in Figure 1(B) in the main paper. Molecular mass in kDa is shown on the left-hand side. (C) HeLa cells were treated with DMSO, 8 $\mu$M ES$_i$, or 8 $\mu$M ES$_i$ and 100 $\mu$g/ml CHX for 8 h, then visualized by phase-contrast microscopy. Arrows indicate vacuolar structures. (D) HeLa cells were left untreated (NT) or treated with 2 mM DTT for 2 h, or with 8 $\mu$M ES$_i$, 10 $\mu$M PS2, or 8 $\mu$M ES$_i$ and 10 $\mu$M PS2 for the time indicated. Xbp1 mRNA splicing was determined by RT–PCR. Unspliced (u) and spliced (s) Xbp1 mRNA are indicated. (E) HeLa cells were treated with 8 $\mu$M ES$_i$, 10 $\mu$M PS2, or 8 $\mu$M ES$_i$ and 10 $\mu$M PS2 for 8 h, then visualized by phase contrast microscopy. Arrows indicate vacuolar structures.

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