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A conformational RNA zipper promotes intron ejection during non-conventional XBP1 mRNA splicing

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Abstract

The kinase/endonuclease IRE1 is the most conserved signal transducer of the unfolded protein response (UPR), an intracellular signaling network that monitors and regulates the protein folding capacity of the endoplasmic reticulum (ER). Upon sensing protein folding perturbations in the ER, IRE1 initiates the unconventional splicing of XBP1 mRNA culminating in the production of the transcription factor XBP1s, which expands the ER's protein folding capacity. We show that an RNA-intrinsic conformational change causes the intron of XBP1 mRNA to be ejected and the exons to zipper up into an extended stem, juxtaposing the RNA ends for ligation. These conformational rearrangements are important for XBP1 mRNA splicing in vivo. The features that point to such active participation of XBP1 mRNA in the splicing reaction are highly conserved throughout metazoan evolution, supporting their importance in orchestrating XBP1 mRNA processing with efficiency and fidelity.

Keywords: endoribonuclease; ER stress; RNA conformational change; unfolded protein response; XBP1 splicing

Subject Categories: Protein Biosynthesis & Quality Control; RNA Biology

Introduction

Protein folding deficiencies in the endoplasmic reticulum (ER) result in accumulation of un- or mis-folded polypeptides, a stress condition that triggers the unfolded protein response (UPR). The UPR is a network of signal transduction pathways that regulate the coordinated reduction of client protein load into the ER and the expansion of the ER folding capacity, thereby ensuring that the organelle remains in homeostasis [1]. In metazoans, the UPR is orchestrated by three principal ER-resident sensors/signal transducers of the ER folding status: the membrane tethered transcription factor ATF6 and the transmembrane kinases PERK and IRE1 [2]. Activation of ATF6 results in its translocation to the Golgi apparatus where it is processed by resident proteases, liberating its cytosolic domain as a soluble transcription factor from the ER membrane [3]. Activation of PERK results in global translational attenuation after phosphorylation of the eukaryotic translation initiation factor 2 (eIF2) [4]. This reduces the load of proteins entering the ER. eIF2 phosphorylation also leads to the production of the transcription factor ATF4, which augments the gene expression program driven by ATF6. Lastly, activation of IRE1 results in activation of a conserved downstream transcription factor, XBP1 [5–7]. The IRE1-mediated branch is the most evolutionarily conserved: IRE1 is the only existing ER stress sensor/transducer outside of metazoans.

IRE1 is a type 1 transmembrane protein that consists of a sensor domain residing in the ER lumen and a Ser/Thr kinase domain fused to a ribonuclease (RNase) domain residing in the cytosol. ER stress leads to oligomerization, trans-autophosphorylation, and allosteric activation of IRE1’s RNase domain [8,9], which transmits the UPR signal to the cell nucleus by a unique splicing mechanism: IRE1 cleaves HAC1 (yeast) or XBP1 (metazoans) mRNAs at non-conventional splice sites [5,7,10,11]. The free exons are joined by the tRNA ligase Trl1 in yeast [12], or the RTCB tRNA ligase complex in metazoans [13–15]. The spliced HAC1 and XBP1 mRNAs encode potent transcription factors that activate several hundred genes that correct ER folding defects [16,17]. Besides splicing, metazoan IRE1 also cleaves ER-bound mRNAs, preventing their translation and thereby diminishing the protein folding load in the ER. This additional functional output is known as regulated IRE1-dependent mRNA decay (RIDD) [18,19]. IRE1 recognizes RNA hairpins in its splicing substrates, cleaving a scissile bond 3’ of a guanosine always found in position 3 of a 7-mer loop [20]. The recognition motif of RIDD substrate RNAs is less conserved, and the determinants that shunt specific mRNAs into the RIDD pathway remain obscure. Biochemical and structural evidence suggest that oligomerization precedes activation and that at least one IRE1 dimer is required for a single cleavage event.

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How the splicing reaction, requiring two cleavage events and the subsequent ligation of the correct exon ends, is orchestrated with fidelity also remains an outstanding question. Here, we provide evidence that XBP1 mRNA is not a passive substrate but an active protagonist in the splicing reaction.

**Results**

**Conserved features in metazoan XBP1 mRNAs are required for recognition and cleavage by IRE1**

To investigate the RNA features required for XBP1 mRNA splicing, we performed multiple-sequence-alignment analyses of RNA segments adjacent to the splice sites from evolutionarily distant metazoans (Figs 1A and EV1A). These analyses identified the conserved exon–intron boundaries conforming to the C(C/U)G|CAGC consensus sequence of the splice junctions separated by the non-conventional intron (Figs 1A and EV1A and see [20]). RNA secondary-fold predictions suggested that these sequences fold into a conserved bifurcated stem-loop (BSL) structure comprised of three stems (Fig 1B, left structure, and Fig EV1B, top structures): a central stem, S1, and two arm stems, S2 and S3. S1 is formed by exon–exon pairing. Note that Fig 1B only shows the portion of the stem that is proximal to the bifurcation. S1 can invariably be extended to form a long stem interrupted by bulges (Fig EV1C), whereas S2 and S3 are short stems formed by exon–intron pairing. Analyses of recently published datasets obtained using two different genome-wide methods to determine RNA secondary structures in living cells [22,23] support our computational predictions of the secondary fold of the XBP1 BSL in vertebrates (Fig EV2).

By contrast, similar structure prediction analyses performed on the corresponding spliced mRNA molecules from which we removed the intron and adjoined the exons computationally suggested that the spliced RNA folds into an extended stem loop (Fig 1B, right structure, and Fig EV1B, bottom structures). In this structure, the exon–exon base pairing of S1 is preserved but now is extended by a previously unrecognized new stem ES1 that results from base pairing between the exons. ES1 is a short stem interrupted by embedded bulges. Strikingly, we found that ES1 is conserved even among distant metazoans (Fig EV1B, bottom structures).
structures), suggesting that it may be part of an evolutionarily conserved architectural requirement for splicing.

Three-dimensional structure predictions on human XBP1 mRNA suggested that the loops containing the exon–intron boundaries are placed within ~32 Å of each other (Fig 1C), which renders cleavage of both sites by a single IRE1 dimer sterically improbable. We surmise that IRE1 assemblies that are obligatorily larger than a dimer are necessary to concomitantly engage both splice sites (see Discussion for details).

To determine the functional importance of the features thus identified, we constructed a short RNA transcript harboring the BSL of human XBP1 mRNA (XBP1-BSL) (Fig EV3A, see Materials and Methods for details). Incubation of XBP1-BSL with recombinant human IRE1α-KR43 (containing IRE1α's kinase and RNase domains and a 43-residue portion of the N-terminal linker that tethers the kinase domain to IRE1's transmembrane domain) yielded the expected cleavage products on denaturing polyacrylamide gels (Fig 2A). In agreement with previous findings [7,24], replacement of the guanosine residues 5′ of the scissile bond at either or both splice junctions by adenosines abrogated the cleavage reaction (Fig 2B–D), while not disrupting the predicted secondary structure fold (Fig 2E). Moreover, these results showed that the cleavage of each splice site proceeds independently and with comparable rates (compare Fig 2B and C), suggesting that IRE1 does not cleave the splice junctions in an obligate order.

Because of the repeated consensus motif at both splice junctions, IRE1 cleavage and exon–exon ligation restores this sequence element. Secondary structure prediction of the spliced XBP1-BSL indicated that the newly generated CCC/CAGC motif would be constrained in a 6-mer loop rather than the conserved 7-mer loop of a-BSL (Fig 2F), demonstrating that, in order to be cleaved by IRE1α, the consensus sequence must be in the correct structural context. Ribonuclease T1 mapping further confirmed the presence of ES1 in the secondary structure of XBP1-BSLspliced (Fig EV3B). In additional support of this notion, phylogenetically distant BSLs likewise predicted the restored conserved placebo in 5- or 6-membered loops after splicing (Fig EV1B). Thus, formation of ES1, which restrains this loop, serves to render the spliced product resistant to re-cleavage by IRE1.

Formation of ES1 requires intron displacement and conformational rearrangements that may facilitate the completion of the splicing reaction after IRE1 cleavage. These changes are driven by the new base pairing between exons, which resembles a zipper-like mechanism. Indeed, the predicted tertiary structure of the spliced RNA suggested that the conformational rearrangements introduced after formation of ES1 would allow the spliced XBP1 stem loop to sway away after its engagement with IRE1 to allow handover for ligation (Fig 2G). We reasoned that such zipper-up of the exons during formation of ES1 might provide an RNA-intrinsic switch important for intron removal.

To test this concept, we resolved the IRE1α-KR43 cleavage products of XBP1-BSL by native polyacrylamide gel electrophoresis (Fig 3A). Indeed, we observed that, upon IRE1α-KR43-catalyzed cleavage, the intron was ejected even in the absence of ligation, while the 5′ and 3′ exons remained base-paired to each other (Fig 3B). Thus, the extended base pairing of the exons via formation of ES1 may serve as the driving force for melting the exon–intron base pairing in S2 and S3. Hence, exon–exon base pairing may have an additional function than promoting adherence of the ends to be joined together, as previously proposed [24].

**Formation of ES1 is required for intron ejection and efficient splicing of XBP1 mRNA**

We next completed the RNA splicing reaction in vitro by inclusion of tRNA ligase (Fig 3A). First, we cleaved XBP1-BSL with IRE1α-KR43 and then added mammalian RTCB tRNA ligase complex (RTCB) or yeast tRNA ligase (Trl1) and confirmed the identity of the splice products by sequencing (Fig EV3C). We observed the production of circularized introns irrespective of thermal denaturation prior to ligase addition, indicating that the reaction conditions remained compatible with the ligation reaction (Fig 3C, lanes 4 and 7, and lanes 10–11, 13–14). By contrast, efficient exon–exon ligation only occurred in the absence of thermal denaturation, indicating that the exons must stay together for splicing to occur (Fig 3C, compare lanes 3–4 with 6–7 and 10–11 with 13–14).

While the reaction catalyzed by Trl1 requires a tripartite mechanism consisting of modifying the cleaved ends (i.e. opening the 2′–3′ cyclic phosphate produced by Ire1 and 5′-end phosphorylation) followed by ligation [25], RTCB employs a direct mechanism where the ends generated by IRE1 are rejoined without prior modification [26]. Regardless of the ligase we used, splicing was always abrogated after thermal denaturation, indicating that the base pairing of the exons is a prerequisite irrespective of the biochemistry of the ligation reaction (Fig 3C).

To explore the role of ES1 formation in these reactions, we engineered an XBP1-BSL mutant, termed XBP1-BSLNZ (“non-zippering”), in which base substitutions prohibit the zippering of the exons and formation of ES1 after cleavage by IRE1, but maintain the predicted secondary and tertiary structures of XBP1-BSL (Figs 4A and EV4A). Cleavage of XBP1-BSLNZ by IRE1α-KR43 was indistinguishable from that of the wild-type XBP1-BSL, validating the design (Fig 4B). Ribonuclease T1 mapping confirmed that the wild-type XBP1-BSL and XBP1-BSLNZ assume similar structures, as predicted by our computational modeling (Fig EV4B). However, by contrast to the wild-type XBP1-BSL, the splicing efficiency of XBP1-BSLNZ was severely compromised (Fig 4C and D, compare lanes 3–7 to lanes 10–14). This impairment occurred independent of which ligase we used.

Interestingly, we noted that, whereas ligation of the wild-type XBP1-BSL by Trl1 proceeded to the anticipated splicing product, XBP1-BSLNZ was efficiently re-ligated at the exon–intron boundaries to restore the input RNA (Fig 4C, lanes 3–7). By contrast, RTCB produced heterogeneous products, resulting from re-ligation of either single exon to the intron (Fig 4D, lane 7, labeled by open arrowheads), both exons to the intron restoring the substrate (Fig 4D, lane 7, labeled by an asterisk), or exon-to-exon resulting in some fully spliced product (Fig 4D, lane 7, labeled by a closed arrowhead). We surmise that the difference between the ligases used may be due to the presence of the additional subunits in RTCB, which include DDX1, a helicase that might assist, albeit inefficiently, in intron removal even when ES1 cannot form.

To show directly that formation of ES1 is important for intron ejection, we next cleaved XBP1-BSLNZ with IRE1α-KR43 and analyzed the reaction products on native polyacrylamide gels.
Figure 2. Cleavage of XBP1 mRNA by IRE1.

A TBE–urea–PAGE gel showing the IRE1-mediated cleavage pattern of the short XBP1-BSL transcript of human origin.

B–D TBE–urea–PAGE gels showing the IRE1-mediated cleavage pattern of human XBP1-BSL cleavage-site mutants. Red crosses over the schematic of the RNA structures on the right side of each gel indicate G-to-A substitutions in the IRE1 cognate sequence within the loops (see E for details).

E Secondary structure of cleavage-site mutants of human XBP1-BSL. Substituted bases are indicated in red. Arrowheads: scissile bonds.

F TBE–urea–PAGE gel showing that IRE1 does not cleave the spliced XBP1-BSL RNA. The RNAs in (A–D, F) were incubated with 0.5 μM of IRE1-x-KR43 for the indicated times.

G Superimposition of the predicted tertiary structures of unspliced and spliced XBP1-BSL RNAs used in (A–D, F). Arrowheads: scissile bonds (unspliced structure); exon–exon junction (spliced structure). The intron is colored in grey, and the exons are colored in blue (unspliced structure) or gold (spliced structure). The guanosines 3′ of the scissile bond are colored in red.

H Pictogram key.

(Fig 4E and F). Under these conditions, the cleavage products of XBP1-BSLNZ co-migrated at the same position as the input RNA unless the sample was heat-denatured (Fig 4E and F, compare lanes 2 and 4), indicating that XBP1-BSLNZ indeed failed to eject the intron. Adding Trl1 to XBP1-BSLNZ in a subsequent reaction after the cleavage reaction was complete led to efficient re-ligation to
XBP1-BSL\textsuperscript{NZ} exhibited a melting temperature that was 5°C higher than that of wild-type XBP1-BSL (Fig EV4C). These results suggest that the added structural plasticity of XBP1-BSL may be important to initiate the zipperung of exons leading to ES1 formation and intron ejection. Similar analyses on the spliced XBP1-BSL yielded an even higher melting temperature (~84°C), indicating the formation of a thermodynamically stable fold after splicing (Fig EV3D).

Formation of ES1 is required for efficient XBP1 splicing in cells

We next asked if the RNA zipperung mechanism leading to ES1 formation is important for XBP1 mRNA splicing \textit{in vivo}. To this end, we introduced a XBP1::GFP\textsuperscript{YFP} reporter bearing the non-zippering mutations in the BSL into HEK293T cells (Fig 5A). Indeed, as measured by semi-quantitative multiplex reverse transcription PCR, upon induction of ER stress by thapsigargin (which inhibits calcium re-uptake into the ER) splicing was severely compromised in cells harboring the XBP1-BSL\textsuperscript{NZ} version of the reporter when compared to cells harboring the wild-type version (Fig 5B, compare lanes 1–3 to 4–6). Note that PCR products derived from endogenous XBP1 mRNA provided a convenient internal control (Fig 5B, lanes 7–9). To ascertain the impact of the compromised splicing on the synthesis of the encoded protein product, we conducted immunoblot analyses of lysates of the cells transfected with the aforementioned constructs. Because the constructs encode N-terminal FLAG epitope tags, we could analyze the accumulation of protein products encoded by the spliced and unspliced RNAs (Fig 5C). Cells expressing the mutant XBP1-BSL\textsuperscript{NZ} version of the reporter were compromised in generating the product encoded in the spliced mRNA and instead accumulated the product encoded in the unspliced one (Fig 5D, compare lanes 1–2 to 4–5). By late time points, the cells expressing the XBP1-BSL\textsuperscript{NZ} reporter showed a mild increase in the spliced product (Fig 5D, compare lanes 4–5 to lane 6); however, this accumulation was much reduced when compared to the product encoded by the wild-type reporter (Fig 5D, compare lanes 3 and 6). This result is consistent with an inefficient conversion rate of the unspliced transcript into its spliced form. Thus, both \textit{in vitro} and \textit{in vivo} experiments converge in support of the model that an RNA conformational rearrangement via ES1 formation and intron ejection ensures efficient XBP1 mRNA splicing.

Discussion

The non-conventional splicing of XBP1 mRNA requires coordinated cleavage and ligation events. Since both exon–intron splice junctions are comprised of stem-loop structures, base pairs need to be melted to eject the intron prior to ligation of the exons. Here, we show that an RNA-intrinsic structural rearrangement allows the severed exons to engage in the pairing of bases (leading to the formation of ES1) that before cleavage were engaged in pairing to the intron (in stems S2 and S3). In this way, the exons zipper up into an extended stem (S1-ES1), juxtaposing the RNA ends to be ligated but constraining the resulting loop so that no functional IRE1 cleavage-site results. This mechanism ensures the correct RNA ends are presented to the ligase, suggesting that the ligase itself does not discriminate between exon and intron ends as corroborated by the observed exon–intron re-ligation in the absence of intron ejection.

Figure 3. Formation of ES1 is required for XBP1 mRNA intron ejection.
A Schematic of experimental steps for PAGE gels shown in (B, C).
B Native PAGE gel showing intron ejection and exon base pairing after IRE1 cleavage. The grey box indicates the unligated ends generated by IRE1 cleavage: 2′-3′ cyclic phosphate and 5′ hydroxyl. The RNAs were incubated with 0.5 μM of IRE1α-KR43 for 10 min prior to thermal denaturation.
C TBE–urea PAGE gel showing the ligation of IRE1-cleaved XBP1-BSL after incubation with the RTCB complex or Trl1 ligase after thermal denaturation. Arrowheads: spliced products. Circ. intron: circularized intron. The RNAs were incubated with 0.5 μM of IRE1α-KR43 for 5 and 30 min. All reactions with RTCB were supplemented with recombinant archease.

restore the input substrate (Fig 4E, lane 5). By contrast, when we used wild-type XBP1-BSL as the substrate, addition of Trl1 led to completion of the splicing reaction, consistent with the substrate’s capacity to form ES1 and eject the intron (Fig 4E, lane 10). Adding the RTCB complex to XBP1-BSL\textsuperscript{NZ} after cleavage led to the production of incomplete splice products (Fig 4F, lane 5, labeled by a diamond) and restored some of the input (Fig 4F, lane 5, labeled by an asterisk), whereas the same experiment using wild-type XBP1-BSL as a substrate exclusively produced a spliced product, despite the reaction not reaching completion (Fig 4F, lane 10). These results are in agreement with those presented in Fig 4C and D.

Independent biochemical methods suggested that the XBP1-BSL adopts a predominant structure (Figs EV2 and EV4B). However, computational predictions showed that even before IRE1 cleavage, XBP1-BSL can adopt more than one secondary structure, which raises the possibility that additional conformers can establish an equilibrium between alternative structures (Fig EV5). By contrast, XBP1-BSL\textsuperscript{NZ} was predicted to assume only one conformation (Figs 4A and EV4A), locking it into a more rigid conformation that mimics the most stable XBP1-BSL conformer (Fig EV4B).
Figure 4. Formation of ES1 is required for XBP1 mRNA splicing.

A Secondary structure of the mutant XBP1-BSLNZ RNA transcript. The base substitutions indicated in red disrupt exon–exon base pairing. Arrowheads: scissile bonds.

B TBE–urea–PAGE gels showing cleavage of the XBP1-BSLNZ or XBP1-BSLWT RNAs by IRE1. The RNAs were incubated with 0.5 μM of IRE1a-KR43 for the indicated times.

C, D TBE–urea–PAGE gels showing IRE1-mediated cleavage of the XBP1-BSLWT or XBP1-BSLNZ RNAs and their splicing after incubation of the cleaved RNAs with Trl1 (C), or the RTCB complex (D). All reactions with RTCB were supplemented with recombinant archease. Circ. intron: circularized intron. Asterisk: input RNA (both exons religated to intron). Open arrowheads: incomplete splice products (5′ or 3′ exons ligated to the intron). Closed arrowhead: spliced product. The RNAs in (C, D) were incubated with 0.5 μM of IRE1a-KR43 for cleavage and with the depicted tRNA ligase for the indicated times. The light grey pictograms on the right of the gels indicate the partially cleaved or incompletely spliced RNAs (containing a single exon and the intron).

E, F Native PAGE gels showing IRE1-mediated cleavage of the XBP1-BSLWT or XBP1-BSLNZ RNAs and their splicing after subsequent incubation of the cleaved RNAs with Trl1 (E) or the RTCB complex (F). The RNAs in (E, F) were incubated with 0.5 μM of IRE1a-KR43 prior to thermal denaturation and ligation. The ligation reactions in (F) were supplemented with recombinant archease. Asterisk in (F): input RNA (both exons re-ligated to intron). Diamond in (F): incompletely spliced products (a single exon ligated to the intron). Note that the spliced product (exon–exon ligation) co-migrates with the incompletely spliced products, which makes it indistinguishable in native PAGE conditions.
Intron ejection occurs in the absence of ligase and was prevented in a mutant in which ES1 cannot form. Failure to form ES1 impaired mRNA splicing in vivo, indicating that the conformational rearrangements leading to intron ejection are important for XBP1 mRNA splicing in a living cell. These features are highly conserved throughout metazoan evolution, supporting their importance in orchestrating the splicing reaction with efficiency and fidelity.

The energetics of the RNA rearrangements are intriguing because, after IRE1-cleavage, the XBP1-BSL RNA settles into a conformation with fewer base pairs (5 base pairs in the newly formed ES1 versus the original 13 base pairs engaging the intron in the uncleaved XBP1-BSL) that a priori would seem less stable. However, melting temperature analysis of the spliced BSL indicates that this is not the case and that the BSL adopts a thermodynamically stable fold after splicing. Analysis of the melting temperature and the prediction of a dynamic equilibrium between multiple alternative conformational states of the uncleaved XBP1-BSL RNA suggest that intron ejection and ES1 formation are energetically favored because a more rigid, less plastic structure is formed. This could be attributable, at least in part, to the existence of conformers with partial zippering of the exons (see Fig EV5, middle structure, which shows incomplete formation of ES1). Moreover, both (i) the local concentration of the intron is reduced upon its ejection and, (ii) as suggested by our in vitro data, the intron is circularized by the tRNA ligase following its excision. Both mechanisms would render the back-reaction substantially less favorable.

Structural models depicting the interaction of yeast Ire1 with substrate RNA postulate recognition of a single stem loop by an RNase-active Ire1 dimer oriented in a back-to-back conformation [21]. Both human IRE1 and yeast Ire1 possess high structural similarity [27,28], suggesting a similar recognition mode, wherein each stem loop of the XBP1-BSL could be recognized and cleaved by an active enzyme dimer. Our tertiary structure predictions place the RNase-active sites. We speculate that the coordinated cleavage of both cleavage events by independent IRE1 dimers or oligomer in which the active sites are spaced by ~40 A˚ (yeast IRE1: PDB ID 3FBV) [8].

IRE1 exerts both corrective and pre-emptive tasks through XBP1 mRNA splicing and RIDD, respectively. Both functions rely on the processing of ER-bound mRNAs by IRE1: one involving a single or multiple non-coordinated endonucleolytic cleavage events with low sequence or structural requirements leading to mRNA decay as...
occurs during RIDD, and the other requiring two precisely coordinated cleavage events that must present the correct substrate for the ligase to complete the XBP1 mRNA splicing reaction. Taken together, our data support the concept of an evolutionarily conserved structural RNA rearrangement that is hard-wired in XBP1 mRNA as a fundamentally important element to shunt XBP1 mRNA into the splicing pathway. This defines XBP1 mRNA not as a static, passive substrate but as an active and dynamic element instrumental to the metazoan UPR.

Materials and Methods

Computational analyses

Homologous sequences containing the XBP1 BSL structure (intron sequences plus upstream and downstream flanking –20-mers) of 34 metazoans were retrieved from the National Center for Biotechnology Information (NCBI) using the Basic Local Alignment Search Tool (BLAST) [32] and assembled into a phylogenetically diverse list that includes vertebrates (mammals, birds, reptiles, amphibians and fishes), invertebrates, and nematodes for multiple sequence alignment using the web-based program ClustalW (European Molecular Biology Laboratory, European Bioinformatics Institute) [33]. The intron–exon boundaries with the XBP1 BSL sequences were visually annotated by direct comparison to those in the XBP1 BSL of human origin. Secondary structure predictions of conserved BSL sequences were performed with the mFold web server (The RNA Institute, University at Albany, State University of New York) [34]. We chose the secondary structures with the highest free energies computed as the sum of free energies assigned to all the loops and base pair stacks as determined by mFold for subsequent analyses. Secondary structure-guided three-dimensional structure predictions were performed using the RNAComposer modeling server (Bioserver, Institute of Computing Science, Poznan University of Technology and the Institute of Bioorganic Chemistry, Polish Academy of Sciences) [35]. Parallel analysis of RNA structure (PARS) scores were computed for the transcript encoding unspliced XBP1 of human origin (Accession NM_005080) as previously described [22, 23] using the dataset in the Gene Expression Omnibus accession GSE50676. icSHAPE reactivities for mouse XBP1 mRNA were retrieved from the Gene Expression Omnibus accession GSE64169. The custom tracks (BigWig files) were uploaded into the University of California at Santa Cruz (UCSC) Genome browser (http://genome.ucsc.edu, and [36]) and displayed on the 2011 mouse genome assembly (UCSC mm10; Genome Reference Consortium GRCm38).

Synthesis of short XBP1-BSL transcripts

Long sense and antisense oligonucleotides containing a minimal T7 RNA polymerase promoter (5′-TAATACGACTCACTATAGG-3′) fused upstream of the sequence containing the XBP1 BSL of human origin (nucleotides 513–592 of NCBI Reference Sequence NM_005080.3) and an RNA polymerase III terminator sequence (5′-TGGCTTTTT-3′) of RNY4 of human origin (NCBI Reference Sequence NR_004393.1) fused downstream of the XBP1 BSL sequence and harboring 5′-EcoRI and 3′-BamHI overhangs were annealed and ligated into the cognate restriction sites of pUC19 (Invitrogen, Life Technologies). The resulting clone, pUC19-T7-hXBP1-BSL-Y4, was digested with BamHI, purified and used as a template for in vitro transcription reactions with T7 RNA polymerase using the HiScribe T7 high-yield RNA synthesis kit (New England Biolabs) following the manufacturer’s recommendations. The transcribed RNA was purified by urea-polyacrylamide gel electrophoresis (urea-PAGE), and the RNA, recovered from gel fragments by the crush-and-soak method, was precipitated with 300 mM NaOAc and 1 volume of isopropanol. No co-precipitants were employed. The precipitated RNA pellet was desalted by two washes with 70% ice-cold ethanol, air-dried and re-suspended in an appropriate volume of either nuclease-free water or RNA resuspension buffer (20 mM HEPEs, 100 mM NaCl, 1 mM Mg(OAc)2). Mutant XBP1 BSL probes were constructed by standard site-directed mutagenesis of the pUC19-T7-hXBP1-BSL-Y4 wild-type clone using mutagenic oligonucleotides and Phusion high-fidelity DNA polymerase (New England Biolabs). Mutant RNAs were transcribed in vitro and purified as described above.

RNA melting curves

Approximately 250–300 ng (for XBP1-BSLWT or XBP1-BSLmut) or 500 ng (for XBP1-BSLPspliced) of in vitro transcribed RNAs was employed for melting curve analyses with 1× SYBR Gold nucleic acid stain (Invitrogen, Life Technologies) in 20 μl reactions using RNA resuspension buffer (20 mM HEPEs, 100 mM NaCl, 1 mM Mg(OAc)2). The RNAs were heated to 90°C for 3 min and then cooled down to 25°C at a rate of 0.1°C/s. The RNAs were then melted in a CFX96 real-time PCR thermocycler (BioRad), and melting curve data points were taken every 0.5°C. To find the temperature of dissociation, the negative of the first derivative was plotted as a function of temperature.

Protein expression and purification

The cytosolic kinase/ribonuclease domain construct of IRE1-α (KR43) was expressed and purified as described previously [37].

The Chaetomium thermophilum tRNA ligase Trl1 (NCBI Entrez Gene ID: 18257519, CTHT_0034810 tRNA ligase-like protein) was cloned from C. thermophilum cDNA into pET15b (EMD Millipore) using the restriction enzymes NdeI and EcoRV. Recombinant, His6-tagged Ctrl1 was expressed in E. coli BL21-CodonPlus (DE3)-RIL (Agilent Technologies) and purified by Ni2+ affinity chromatography using a HisTrap FF column (GE Healthcare Life Sciences), followed by size-exclusion chromatography using a HiLoad 16/60 Superdex200 pg column (GE Healthcare Life Sciences) in 25 mM Tris/HCl pH 7.1, 300 mM NaCl, and 1 mM MgCl2.

Human archease constructs were cloned as described elsewhere [38]. His-tagged archease was expressed in E. coli BL21-CodonPlus (DE3)-RPL (Agilent Technologies) and purified by Ni2+ affinity chromatography using a HisTrap FF column (GE Healthcare Life Sciences), followed by tag removal with thrombin (Sigma-Aldrich) and a final size-exclusion chromatography step using a HiLoad 16/60 Superdex200 pg column (GE Healthcare Life Sciences) in 25 mM Tris/HCl pH 7.5, 100 mM NaCl, 5% glycerol, and 1 mM TCEP. The human RTCB complex was immunoprecipitated from HEK293 cells expressing FLAG-RTCB as described elsewhere [38].
All proteins were flash-frozen in liquid nitrogen directly after purification and kept at −80°C.

In vitro cleavage and splicing assays

In vitro transcribed, PAGE-purified, refolded RNAs (50 nM) were incubated with 0.5 μM IRE1α-KR43 for the indicated times in RNA cleavage buffer (20 mM HEPES pH 7.5, 70 mM NaCl, 2 mM Mg(OAc)₂, 1 mM TCEP, and 5% glycerol). Stop solution (10 M urea, 0.1% SDS, 1 mM EDTA, 0.05% xylene cyanol, 0.05% bromophenol blue) was added at five-fold excess to stop the reactions followed by heating at 80°C for 3 min. The denatured samples were then loaded on 15% TBE–urea gels (Invitrogen, Life Technologies) and the gels stained with SYBR Gold nucleic acid stain (Invitrogen, Life Technologies).

For splicing assays, the cleavage reactions were performed as described above and subsequently stopped by addition of the IRE1 inhibitor 4μB8 (CAS No. 14003-96-4; Matrix Scientific, Cat. No. 037985) to a final concentration of 5 μM. Ligation of the cleaved RNA was initiated with 500 nM Cfl1l (1 mM ATP, 1 mM GTP) or mammalian RTCB complex (1:5 dilution of FLAG-RTCB eluate, 500 nM recombinant archease, 1 mM GTP, 250 μM MnCl₂). Reactions were analyzed by denaturing urea–PAGE (after addition of stop solution, as described above) or native PAGE (see next paragraph).

Native polyacrylamide gel electrophoresis

Cleavage and splicing reactions were carried out as described above, and the reaction products were loaded on 6% Tris-borate DNA retardation gels (Invitrogen, Life Technologies) using native RNA loading buffer (15% Ficoll w/v, 84% Tris-borate-EDTA (TBE), 0.5% bromophenol blue, 0.5% xylene cyanol). The gels were run in pre-chilled 0.5× TBE at 100 V (constant voltage) and 4°C for 100 min. The gels were subsequently stained with SYBR Gold nucleic acid stain (Invitrogen, Life Technologies) to visualize the RNA-containing complexes by ultraviolet trans-illumination.

Sequencing of splicing products

Splicing reactions using the XBP1-BSLWT RNA as a substrate were carried out and the samples separated in urea–PAGE gels as described in “In vitro cleavage and splicing assays”. A control reaction for sequencing the unspliced RNA was set up aside with no IRE1 and no ligase. The RNAs of interest (unspliced and complete splice product) were then cut out from the gels and purified by the crush-and-soak method followed by precipitation. GlycoBlue (Ambion, Life Technologies) was added as a co-precipitant. The purified RNAs were resuspended in 20 μl of nuclelease-free water, and 9 μl of the RNA solution was used for reverse transcription using Superscript III (Invitrogen, Life Technologies) in 20 μl reactions following the manufacturer’s recommendations. The oligonucleotide for reverse transcription was T7-hXBPI-Y4_hyb_M13R: 5'-GTGGTTGATCGCCCGGAACAAGGATCCAAAAAGGCGGACTGCT-3'. About 50% of the cDNA was the used as template for PCR amplification using Phusion high-fidelity DNA polymerase (New England Biolabs) and oligo-nucleotides T7-hXBPI-F: 5'-TAATACGACTCACTATAGGGAGAGGCTGTGGC-3' and T7-hXBPI-Y4_hyb_M13R. The resulting PCR products were purified using DNA Clean & Concentrator-5 columns (Zymo Research) and eluted in 12 μl. The purified PCR products (2 μl) were cloned into pCRII-Blunt-TOPO (Invitrogen, Life Technologies) following the manufacturer’s recommendations and transformed into competent DH5α E. coli. About 10 colonies from each transformation were sequenced using the Sanger method.

RNA structure determination using ribonuclease T1

Approximately 25–30 ng of in vitro transcribed RNAs were resuspended in RNA re-suspension buffer (20 mM HEPES, 100 mM NaCl, 1 mM Mg(OAc)₂) and refolded. The RNAs were incubated with increasing concentrations of biochemistry grade RNase T1 (Ambion, Life Technologies) for 15 min at 20°C, and the reactions were immediately stopped with stop solution (10 M urea, 0.1% SDS, 1 mM EDTA, 0.05% xylene cyanol, 0.05% bromophenol blue) at 2-fold excess followed by heating at 80°C for 3 min. The denatured samples were loaded on 15% TBE–urea gels (Invitrogen, Life Technologies) and the gels stained with SYBR Gold nucleic acid stain (Invitrogen, Life Technologies). The final RNase T1 concentrations were (in U/μl) as follows: 0.1, 0.03, 0.01, 0.003, and 0.001. RNA ladders included a mixture of three PAGE-purified synthetic RNA oligonucleotides (a 31-mer, GE Healthcare Dharmacon Inc.; a 21-mer, Integrated DNA Technologies; and a 17-mer, Integrated DNA Technologies) or IRE1-cleaved XBP1-BSLWT.

Mammalian expression constructs and cell transfection

The coding sequence of an ER stress reporter construct consisting of an N-terminal FLAG epitope-tagged partial coding sequence of XBP1 of human origin containing the IRE1-cognate intron and the BSL structure, and fused to the venus variant of GFP (kind gift of Takao Iwawaki, RIKEN) [39], was used as a template to generate retroviral expression constructs containing both the wild-type XBP1-BSL and the non-zipper XBP1-BSLμN2 mutant. The wild-type sequence was amplified by PCR using Phusion high-fidelity DNA polymerase (New England Biolabs) using oligonucleotides with 5' and 3' EcoRI engineered restriction sites. The resulting PCR product was cloned into the cognate sites of the mammalian expression vector pLPX (Clontech). To generate the mutant XBP1-BSLμN2 construct, a fusion PCR strategy was employed. Left and right arm PCR products were generated with mutagenic oligonucleotides. The arms were then fused in a second PCR employing the outermost oligonucleotides with 5' and 3' EcoRI engineered restriction sites. The resulting PCR product harboring the mutant XBP1-BSLμN2, GFPVenus coding sequence was cloned into the cognate sites of the mammalian expression vector pLPX (Clontech). HEK293T cells were transfected with 1 μg of either plasmid using Lipofectamine 2000 (Life Technologies) following the manufacturer’s recommendations. The cells were diluted 1:3 24 h after transfection and re-seeded onto 6-well plates. After an additional 24 h, the cells were treated with the 200 nM of ER stress inducing agent thapsigargin (Sigma-Aldrich) for 2 or 4 h. The sequences of the oligonucleotides used are the following (restriction enzyme sites and mutant bases are underlined): Xhol-FLAG-Hs-XBP1-Fwd: 5'-ATTAATCTCGAGACCCACACAATTACGACTCATATAGGGAGAGGCTGTGGC-3'; Hs-XBP1-BSLμN2-Amo: 5'-ATTAATCTCGAGACCCACACAATTACGACTCATATAGGGAGAGGCTGTGGC-3'; GGCAGCGTCAGATCGTTCCAGCAGCCTCTCGTCGTCAACCT-3';
and Hs-XBP1-Bst.NS-Antisense: 5'-AGGTGCACGTAGTGAAGTGCTGTCGGACTCTCTGACCACCCTACT-3'.

cDNA generation and multiplex semi-quantitative PCR

Transfected cells exposed to DMSO or thapsigargin were collected in 1 ml of TRIzol reagent (Life Technologies), and total RNA was extracted following the manufacturer’s recommendations. To generate cDNAs, 500 ng of total RNA were reverse transcribed using the SuperScript VILO system (Life Technologies) following the manufacturer’s recommendations. The resulting 20 µl reverse transcription reactions were diluted to 200 µl with 10 mM Tris–HCl pH 8.2, and 1% of this dilution was used for multiplex semi-quantitative PCR. The multiplex PCR was set up using 1 µM of the common forward oligonucleotide and 0.5 µM of each of the gene-specific reverse oligonucleotide, 0.4 units of Taq DNA polymerase (Thermo Scientific), 0.2 mM of each dNTP, and 1.5 mM MgCl₂, in a 20 µl reaction using the following buffer system: 75 mM Tris–HCl pH 8.8, 20 mM (NH₄)₂SO₄, and 0.01% Tween-20. The oligonucleotide sequences are the following: Hs_XBP1_Fwd: 5’-GGAGTTCAGGGACCCCGCCTG-3’; Hs_XBP1_Rev: 5’-ACTGGGTCCAAGTTG-3’; eGFP_Rev: 5’-AAGTCGTGCTGCTTCATGTG-3’. PCR products were amplified for 28 cycles and resolved on 2.5% agarose gels (1:1 mixture of regular and low-melting point agarose) stained with ethidium bromide.

Immunoblotting

Total cell lysates were collected in SDS sample buffer (62.5 mM Tris pH 6.8, 10% glycerol, 2% SDS, 0.004% bromophenol blue). Lysates were sonicated for ~15 s to shear the genomic DNA. 2-mercaptoethanol was added to a final concentration of 5% to the lysates just prior to boiling and loading on SDS–PAGE gels. Proteins were separated by electrophoresis and transferred onto 2.0 µm pore nitrocellulose membranes. The blocked antibody membrane was incubated with a mouse monoclonal anti-FLAG antibody (clone M2, Sigma-Aldrich F1804, 1:1,000) or an anti-GAPDH rabbit polyclonal antibody (Abcam ab9485, 1:2,000). Immunoreactive bands were detected using HRP-conjugated secondary antibodies (Amersham, GE Healthcare Life Sciences NA931, NA934, 1:5,000) and luminol-detecting antibodies (Aldrich F1804, 1:1,000) or an anti-GAPDH rabbit polyclonal antibody. (Abcam ab9485, 1:2,000). Immunoreactive bands were detected using HRP-conjugated secondary antibodies (Amersham, GE Healthcare Life Sciences NA931, NA934, 1:5,000) and luminol-based enhanced chemiluminescence substrates (SuperSignal West Dura Extended Duration Substrate, Pierce, Life Technologies) and exposed to radiographic film or imaged directly in a digital gel imager (Chemidoc XR+, BioRad). Digital images were automatically adjusted for contrast using the photo editor Adobe Photoshop (Adobe Systems).

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Author contributions

JP and DAA performed experiments and analyzed data. ASM purified human Ire1-KR43 and helped with in vitro assays. JP, DAA, and PW designed the research and wrote the paper.

Conflict of interest

The authors declare that they have no conflict of interest.

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