miR-216b regulation of c-Jun mediates GADD153/CHOP-dependent apoptosis

Zhenhua Xu1, Yiwen Bu1, Nilesh Chitnis1, Costas Koumenis2, Serge Y. Fuchs3 & J. Alan Diehl1

The ability of the unfolded protein response, UPR, to regulate cell homeostasis through both gene expression and protein synthesis has been well documented. One primary pro-apoptotic protein that responds to both PERK and Ire1 signalling is the CHOP/GADD153 transcription factor. Although CHOP deficiency delays onset of cell death, questions remain regarding how CHOP regulates apoptosis. Here, we provide evidence demonstrating that CHOP/GADD153-dependent apoptosis reflects expression of micro-RNA, miR-216b. MiR-216b accumulation requires PERK-dependent induction of CHOP/GADD153, which then directly regulates miR-216b expression. As maximal expression of miR-216b is antagonized by Ire1, miR-216b accumulation reflects the convergence of PERK and Ire1 activities. Functionally, miR-216b directly targets c-Jun, thereby reducing AP-1-dependent transcription and sensitizing cells to ER stress-dependent apoptosis. These results provide direct insight into the molecular mechanisms of CHOP/GADD153-dependent cell death.
The endoplasmic reticulum (ER) serves as a nexus for the folding and maturation of proteins that transit the secretory pathway. Proteins are imported co-translationally whereupon they undergo additional energy intensive post-translational modifications before folding and maturation. Protein folding is both regulated and sensed by ER resident chaperones, such as Grp78/Bip and Grp94 (refs 1–5). Under conditions where protein production outpaces protein folding capacity, for example, under nutrient-deficient conditions as might occur during neoplastic growth, misfolded proteins accumulate in the ER triggering a stress sensing/adaptive pathway referred to as the unfolded protein response (UPR).

Mammalian cells contain three ER transmembrane proteins that function as proximal effectors of the UPR. Inositol-requiring enzyme, Ire1, isoforms (α, ubiquitously expressed; β tissue restricted) are composed of a luminal domain that senses stress, a single transmembrane domain, and a cytosolic tail that contains both a protein kinase domain and an RNase domain6,7. Ire1 triggers increased expression of numerous ER chaperones through the activation of the X-box-binding protein 1 (Xbp1) transcription factor8,9. Accumulation of Xbp1 is mediated by the RNase and splicing function of Ire1; stress-induced splicing generates a shorter Xbp1 mRNA that is more efficiently translated10,11. Protein kinase RNA-like ER kinase (PERK) is activated in a manner analogous to the Ire1; it catalyses serine 51 phosphorylation of eIF2α resulting in repression of protein synthesis12–14. The third signalling component are the transmembrane transcription factors ATF6α/β (refs 15,16). Although normally tethered to the ER, upon stress, ATF6 migrates to the trans-Golgi, where it is processed by S1P and S2P proteases to release the N-terminal DNA-binding transcription factor domain17–19. Activation of PERK, Ire1 and ATF6 is mediated by sequences within their respective luminal domains to which the ER chaperone, Bip, binds20. Activation is triggered by increased unfolded proteins, which compete for Bip binding.

The ability of the UPR to regulate protein synthesis, gene expression and contribute to cell homeostasis has been well documented. However, an added layer of communication that entails the regulation of small non-coding RNAs, micro-RNAs, was recently revealed. Micro-RNAs are 20–22 nucleotide molecules that generally impact protein expression by virtue of their capacity to degrade target mRNAs or reduce their efficiency of translation21,22. During the past 4–5 years, work from several groups has revealed that all three branches of the UPR regulate specific subsets of micro-RNAs23–27. The modes of regulation include Ire1-mediated micro-RNA degradation23, ATF4-dependent and ATF6-dependent transcription27. The consequences of micro-RNA induction include the modulation of protein expression as would be expected, while certain micro-RNAs also directly contribute to regulated gene expression27.

The UPR can communicate both pro-adaptive and pro-apoptotic signals. Ambiguities remain regarding whether an adaptive or apoptotic signal is generated given that all three transducers have been associated with both survival and apoptosis under varying conditions (9,28–34). One primary pro-apoptotic protein that responds to both PERK and Ire1 signalling is the CHOP/GADD153 transcription factor. Although CHOP deficiency delays onset of cell death, questions remain regarding how CHOP regulates apoptosis. The data presented here reveal evidence for CHOP-dependent micro-RNA, miR-216b, which subsequently suppresses c-Jun expression thereby, triggering cell death.

Results

MiR-216b is induced by the UPR. To identify micro-RNAs that are induced following prolonged ER stress, RNA harvested from NIH3T3 cells challenged with tunicamycin (Tu) was subjected to micro-array analysis. MiR-211, not represented in this figure, induction was assessed as an internal control. MiR-216b and miR-217 exhibited increased expression throughout the time course (Fig. 1a). Several groups have evaluated miR-216b expression and potential function in various cancers including colorectal cancer, hepatocellular carcinoma and nasopharyngeal carcinoma35–37; we therefore chose to assess miR-216b regulation and function during ER stress. To verify the microarray data, RNA was collected from Tm-challenged NIH3T3 cells, assessed for CHOP induction as a control for UPR activation and for miR-216b. CHOP induction was noted by 2 h and consistent with our microarray data, we observed an accumulation of miR-216b (Fig. 1b). Accumulation of miR-216b was evident following the exposure of cells to an independent inducer of ER stress, thapsigargin (TG; Fig. 1c). MiR-216b expression was also induced in response to chemotherapeutic agents that generate ER stress38 and physiologically relevant low glucose (Fig. 1d,e).

PERK-dependent induction of miR-216b. Because PERK was previously implicated as a regulator of micro-RNA accumulation during ER stress, we assessed its contribution to the regulation of miR-216b expression. Wild-type or PERK deficient murine embryonic fibroblasts39 were treated with TG (500 nM), and expression of miR-216b was assessed by quantitative PCR (qPCR). PERK deficiency was confirmed by western blot analysis using a PERK-specific antiserum and through assessment of CHOP accumulation following stress (Fig. 2a). MiR-216b accumulation was noted in wild type but not PERK−/− cells (Fig. 2a). To corroborate the dependence of miR-216b expression on PERK activity, we used GSK2606414 (GSK414), a highly specific PERK small-molecule inhibitor40,41. GSK414 effectively inhibited PERK activation and miR-216b induction (Fig. 2b).

PERK-dependent regulation of gene expression is often dependent upon increased accumulation of the ATF4 transcription factor, whose accumulation is in turn dependent upon eIF2α phosphorylation13,14,42–44. To address the contribution of eIF2α and ATF4, wild-type mouse embryonic fibroblasts (MEFs) or MEFs isolated from eIF2αS51A knockin embryos45 or from ATF4−/− embryos46 were challenged with TG. MiR-216b induction was absent in both eIF2αS51A and ATF4−/− backgrounds but was induced in wild-type MEFs (Fig. 2c,d) demonstrating a role for ATF4 in the regulation of miR-216b expression.

As ATF4 accumulates within the first 2–4 h of ER stress46,47, and miR-216b accumulation occurs primarily between 5 and 10 h, we considered the possibility that ATF4-dependent regulation was indirect. We focused our attention on CHOP, a direct transcriptional target of ATF4 (ref. 43). MiR-216b induction was abolished in chop knockout MEFs (Fig. 2e). Let7a accumulation was assessed as a nonspecific control (Supplementary Fig. 1). Consistent with CHOP regulation, restoring CHOP expression in chop−/− MEFs rescued stress-dependent miR-216b expression (Fig. 2f). These data demonstrate that ER stress-dependent induction of miR-216b reflects a pathway minimally composed of PERK, eIF2α, ATF4 and CHOP.

We also assessed miR-216b expression in a panel of MMTV-neu tumours that were derived in mice with either wild-type PERK or from mice in which PERK was excised from mammary epithelium with Cre recombinase35,48. In this cohort, miR-216b expression was significantly reduced in PERK-deficient tumours and this correlated with c-Jun expression levels, providing evidence for PERK-dependent induction of miR-216b in vivo (Fig. 2g,h).
MiR-216b is a direct CHOP transcriptional target. Interestingly, ATF4 and CHOP share a set of target genes, with the preference of binding to the similar motifs (GCATCAT/G)47. Using an unbiased motif search of 2 kbp upstream of mouse miR-216b, we found a potential binding site for ATF4 and/or CHOP at -266 to -259 (Supplementary Fig. 2a). To evaluate potential binding to this motif, we performed chromatin immunoprecipitation (ChIP). ChIP with either anti-CHOP (Fig. 3a, right and left panel) or anti-ATF4 (Supplementary Fig. 2b) followed by qPCR revealed that TG treatment triggered a significant enrichment of CHOP on the miR-216b promoter relative to unstressed cells and control IgG. Under the same conditions, there was no apparent enrichment of ATF4 on the miR-216b promoter, although binding of ATF4 to the CHOP promoter was confirmed (Supplementary Fig. 2b,c).

To further evaluate whether the upstream region from -266 to -259 region is responsible for miR-216b regulation, we used a reporter assay system where 1 kbp of upstream miR-216b

**Figure 1 | ER stress induced miR-216b expression.** (a) Heat map depicting differentially expressed microRNAs (at least two-fold) following exposure to tunicamycin (Tm) as indicated. (b,c) qPCR analysis of CHOP mRNA levels and miR-216b expression in NIH3T3 cells treated with Tm (1μg ml⁻¹) or thapsigargin (TG, 500 nM) for indicated times. Averages are calculated from three independent experiments. (d) qPCR analysis of miR-216b expression and CHOP mRNA levels in NIH3T3 cells cultured in low-glucose medium for indicated times. (e) qPCR analysis of miR-216b expression and CHOP mRNA levels in U2OS cells treated with paclitaxel (100 nM) for indicated times. Data are mean ± s.d. of three independent experiments. Statistical analysis was analysed by Student’s t-test. (*P<0.05, treatment versus non-treatment).
sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c). Luciferase sequence was cloned upstream of the firefly luciferase gene (Fig. 3b). Following expression in 293T cells, luciferase activity was induced following TG treatment (Fig. 3c). Luciferase activity was abolished by GSK414 (Fig. 3c).

Figure 2 | PERK-dependent miR-216b induction. (a) PERK+/+ and PERK−/− MEFs were treated with 500 nM TG for indicated times. MiR-216b was assessed by qPCR (left graph); PERK and CHOP were assessed by immunoblot (right). (b) MiR-216b levels were quantified by qPCR following exposure of cells to thapsigargin and a small-molecule PERK inhibitor (left). PERK, eIF2α, and CHOP induction were assessed by immunoblot (right). (c-e) MEFs of the indicated genotype were treated with TG (500 nM) for indicated intervals. Protein extracts from these cells were immunoblotted for the proteins as indicated (lower panels) and miR-216b levels were quantified by qPCR (upper panels; n = 3). (f) CHOP−/− MEFs were transfected with vector or CHOP and 2 days later treated with TG (500 nM) for indicated intervals. Protein extracts from these cells were immunoblotted for CHOP and miR-216b levels quantified by qPCR (n = 3). (g) MiR-216b expression and (h) c-Jun mRNA levels were analysed in MMTV-Neu tumours from either PERK+/+ or PERK−/− background. Data represent mean ± s.d. of three independent observations. Statistical significance was analysed by Student’s t-test. (*P<0.05, WT versus −/−.)

CHOP-binding site through site-directed mutagenesis abrogated UPR-dependent induction (Fig. 3c-white bars). Actinomycin D (2 µg ml−1) treatment before exposure to TG treatment suppressed miR-216b expression, consistent with transcription dependence of miR-216b induction (Fig. 3d). To determine
whether miR-216b biogenesis is Dicer dependent we utilized inducible Dicer Knockout (Dicerfl/fl) or heterozygous (Dicer+/–) MEFs after deletion of the floxed allele(s) with hydroxytamoxifen. After treatment with TG, miR-216b expression was significantly lower in Dicer-KO cells than that in Dicerfl/fl or Dicer+/– cells demonstrating Dicer dependence (Fig. 3e).

Ire1α inhibits miR-216b expression. Mammalian Ire1α initiates downstream signalling through unconventional splicing of the transcription factor Xbp1 (ref. 10), phosphorylation of downstream targets or via regulated Ire1-dependent decay (RIDD) of multiple substrates50. Ire1α RIDD activity also contributes to the regulation of microRNA accumulation during ER stress25. To determine the contribution of Ire1α in miR-216b expression, we treated Ire1α knockout MEFs (Ire1α−/−) and wild-type MEFs with TG. A significant increase in miR-216b induction in Ire1α knockout cells relative to wild type was observed (Fig. 4a). As an independent analysis, we used three different short hairpin RNAs (shRNAs) targeting Ire1α; notably, all three shRNAs reduced Ire1α expression thereby permitting increased accumulation of miR-216b following ER stress (Fig. 4b).

As the miR-216b stem-loop contains a putative Ire1α cleavage site (CUGCAG (ref. 51); Fig. 4c), we considered whether miR-216b might be a RIDD substrate for Ire1α. Recombinant Ire1α (aa 465–977), did not cleave a labelled miR-216b precursor (incubation up to 6 h), but did cleave Xbp1 (Xbp1-mini, 5 ’-CUG AGUCCGAGCAGCUGAG-3 ’; Fig. 4d). As an independent approach, we transfected HA-tagged Ire1α wt, Ire1α-509A and Ire1α-907A into 293T cells followed by immunoprecipitation with an anti-HA antibody. Immune-purified Ire1α wt cleaved the Xbp1 single stem-loop mini-substrate, whereas Ire1α-509A and Ire1α-907A were inactive as expected (Supplementary Fig. 3). No specific cleavage of miR-216b was noted suggesting that Ire1α-dependent suppression of miR-216b is indirect. Published work has demonstrated that CHOP expression is elevated in Ire1α−/− cells52, providing a possible explanation for increased miR-216b expression. Indeed, CHOP expression was more abundant in Ire1α−/− cells (Fig. 4e) correlating with increased miR-216b levels. Finally, to address whether Xbp1 suppress miR-216b accumulation, we enforced expression of spliced Xbp1 expression in Ire1α−/− cells (Fig. 4f). Enforced sXbp1 resulted in reduced miR-216b consistent with sXbp1-suppressing miR-216b (Fig. 4g).

MiR-216b regulates c-Jun accumulation during ER stress. Before searching for targets of miR-216b, a luciferase reporter gene containing three tandem miR-216b-binding sites in its 5 ’ untranslated region (UTR) was used to determine whether miR-216b induction was of functional consequence. Treatment of cells

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**Figure 3** CHOP binds to miR-216b promoter region. (a) Occupancy of CHOP on promoter of mouse miR-216b. NIH3T3 cells (left panel) and U2OS (right panel) were treated with/without TG for 5 h, followed by ChIP assay using IgG or an anti-CHOP antibody. Precipitated DNA was subjected to qPCR analysis. All ChIP data represent the mean and standard error of the mean from three independent experiments. (b) Schematic representation of miR-216b promoter reporter (216b-pro-pGL4; left panel). 293T cells were co-transfected with 216b-pro-pGL4, 216b-mut-pGL4 and Renilla-Luc as an internal control. 216b levels were measured with qPCR (mean ± s.d., n = 3). (c) Luciferase/renilla ratio to input (P1 P2). (d) Luciferase/renilla ratio to input (P1 P2). (e) Luciferase/renilla ratio to input (P1 P2).
expressing the reporter with TG resulted in a 50% reduction in luciferase activity (Fig. 5a; left two columns). Co-transfection with miR-216b mimic also resulted in an ~50% decrease in reporter gene expression; inclusion of TG reduced expression further by ~80% (Fig. 5a; middle columns). Luciferase activity increased 1.5-fold when co-transfected with an antagonir, A-miR-216b (Fig. 5a; right columns).

To identify miR-216b targets, a computational algorithm was utilized to search 3′UTR regions of annotated cDNAs for miR-216b seed sequence matches; TargetScan (http://www.targetscan.org), revealed c-Jun as a high relevance target (Fig. 5b). TG exposure resulted in a time-dependent decrease in c-Jun levels (Fig. 5c). To assess the contribution of miR-216b to stress-dependent inhibition of c-Jun, U2OS cell lines that stably overexpressed an shRNA vector encoding a miR-216b mimic or an anti-miR-216b were generated and subsequently exposed to a time-course of TG treatment. Cells expressing miR-216b mimic exhibited reduced basal c-Jun levels and increased kinetics of c-Jun loss relative to control (Fig. 5d; quantification cJun knockdown was confirmed by immunoblot, and miR-216b levels were quantified by qPCR. (c) Illustration of the potential Ire1α cleavage sites within pre-miR-216b. (d) γ-32P labelled pre-miR-216b or Xbp-1 oligonucleotides were incubated with recombinant protein Ire1α (aa 465–977) for 2, 4 and 6 h, and cleavage products were resolved by a urea-polyacrylamide gel electrophoresis and visualized by autoradiography. (e) Ire1α−/− and Ire1wt MEFs were treated with TG (500 nM) for indicated intervals. Ire1α, Bip and CHOP were assessed by immunoblot. (f) Xbp1s expression and (g) miR-216b levels in U2OS cells transfected with vector or Xbp1s were quantified by qPCR following exposure of cells to thapsigargin. All panels provide values that are the average of three independent experiments and error bars indicate standard deviation among all the replicates. Statistical analysis was analysed by Student’s t-test (*P<0.05).

In combination with c-Fos, c-Jun forms the AP-1 early response transcription factor53,54. It is expected that miR-216b induction should trigger decreased AP-1 activity. To test this hypothesis, a reporter construct wherein luciferase expression is driven by a minimal promoter and 3X AP-1 binding sites (TGAACCTCA) was utilized. AP-1-dependent activity was reduced by ER stress (Fig. 5e). The inhibition of AP-1-dependent reporter gene expression was amplified by co-expression of ectopic miR-216b and attenuated by A-miR-216b (Fig. 5e). We also reasoned that if miR-216b induction is CHOP dependent, CHOP deficiency should ablate c-Jun inhibition during ER stress. Consistently, CHOP−/− exposed to ER stress failed to suppress c-Jun after TG treatment (Fig. 5f). Finally,
assessment of expression of c-Jun target genes revealed that ER stress and miR-216b reduced expression, whereas A-miR-216b rescued expression (Fig. 5g–j; quantification of cJun Supplementary Fig. 4d). These results revealed a direct relationship between miR-216b expression and c-Jun downregulation following ER stress.

Suppression of c-Jun sensitizes cells to apoptosis. As c-Jun protects cells from ultraviolet and tumour-necrosis factor-α-induced apoptosis through cooperation with NF-κB55,56, we considered the possibility that miR-216b might regulate ER stress-dependent apoptosis through regulation of c-Jun. Initially, we determined whether modulation of miR-216b altered ER stress-dependent

Figure 5 | MiR-216b targets the c-Jun 3′ UTR. (a) MiR-216b reporter construct (left) was expressed in mimic control (ctrl), miR-216b or A-miR-216b-expressing U2OS cells. Luciferase activity was measured and normalized to internal transfection control Renilla. Error bars represent standard deviation for three independent experiments. (b) MiR-216b seed sequence matches in the c-Jun 3′ UTR. (c) NIH3T3 cells were treated with PERK inhibitor for 1 h and followed by 500 nM thapsigargin (TG); c-Jun and CHOP protein levels assessed by immunoblot. (d) U2OS stable cell lines expressing control, miR-216b or A-miR-216b were treated with 500 nM TG as indicated. Levels of the indicated proteins were measured by immunoblot. (e) An AP-1 luciferase reporter was transiently introduced into NIH3T3 cells along with a Renilla control. Cells were co-transfected as indicated with control, miR-216b or A-miR-216b. 48 h post transfection, cells were treated with TG before quantifying luciferase relative to Renilla control. (f) CHOP+/− and CHOP−/− MEFs were treated with 500 nM TG for indicated intervals. CHOP and c-Jun were immunoblotted. (g–j) qPCR analysis of CHOP mRNA levels downstream target genes of c-Jun in U2OS treated as in (d). Data present the mean of three independent experiments, error bars indicate the standard deviation. Statistical significance was analysed by Student’s t-test (*P<0.05).
apoptosis. Consistent with miR-216b as a regulator of cell survival, overexpression of miR-216b increased apoptosis induced by TG (Fig. 6a). In contrast, A-miR-216b-expressing cells were largely refractory to the TG-induced apoptosis (Fig. 6a). As an independent measure of cell survival, we exposed control, miR-216b- and A-miR-216b-expressing U2OS to TG for 0 or 3 h following which cells were supplied with fresh medium lacking TG. The cells were allowed to recover for 12 days, at which point surviving cells/colonies were quantified. miR-216b-expressing U2OS cells were significantly more sensitive to stress than control, whereas A-miR-216b-expressing cells were refractory to TG-induced death (Fig. 6b). The effect of miR-216b on recovery from ER stress was also observed with a second ER stressor, Tm (Supplementary Fig. 5). Cell doubling times were not affected by the expression of either A-miR-216b or overexpression of miR-216b mimic confirming that miR-216b is regulating cell viability not cell proliferation (Fig. 6c).

To investigate whether the effect of miR-216b is c-Jun-dependent, c-Jun was suppressed using three independent shRNAs targeting distinct regions of c-Jun. c-Jun loss triggered a significant increase in cleaved poly (ADP-ribose) polymerase (PARP) that correlated with the degree of c-Jun suppression (Fig. 7a). We compared the impact of overexpression of sh-c-Jun versus overexpression of miR-216b by serial increases in viral supernatants encoding either sh-c-Jun or miR-216b. This experiment revealed that while the artificial sh-c-Jun was significantly more potent in reducing c-Jun than miR-216b, the degree of cleaved caspase 3 triggered by either correlated with the degree of c-Jun knockdown (Fig. 7b). If loss of c-Jun triggers

Figure 6 | MiR-216b expression sensitizes cells to ER stress. (a) Representative FACS analysis histograms of U2OS stable cells expressing control, miR-216b or A-miR-216b after treatment with 500 nm TG. (b) U2OS cells stably expressing control, miR-216b or A-miR-216b were plated and treated with 400 nM TG for 1 or 3 h, and then returned to normal growth medium for 12 days (d). Plates were assessed for colony formation by staining with Giemsa staining (left). Quantification of colonies (right). (c) Cell doubling was quantified over 6 days. Values are means ± s.d. (n = 3) and statistical significance was analysed by Student’s t-test. (*P < 0.05; #P > 0.05).
apoptosis, overexpression of c-Jun should over-ride ER stress-dependent and miR-216b-dependent apoptosis as assessed by caspase 3 cleavage. U2OS cells were infected with retrovirus coding c-Jun, lacking a 3'UTR, followed by TG treatment. Enforced c-Jun expression in U2OS cells rescued cells from cell death induced by miR-216b expression and TG treatment (Fig. 7c,d). Finally, we reasoned that if miR-216b mediates CHOP-dependent apoptosis, enforced expression of miR-216b should sensitize CHOP \(^{-/-}\) MEFs to UPR-dependent cell killing. Consistently, enforced miR-216b expression resulted in increased cleaved caspase 3 in CHOP \(^{-/-}\) MEFs (Fig. 7e).

**Discussion**

Activation of the UPR under varying stress conditions triggers an alteration in cellular protein content through direct modulation of gene expression, micro-RNA biosynthesis and protein translation rates; this in turn impacts cell division and ultimately enforces cell fate determination. The final outcome, cell adaptation or cell death, is influenced by both stress intensity and duration. Importantly, how stress intensity is sensed by the UPR transducers, PERK/Ire1/ATF6, and transmitted into a fate decision remains to be elucidated. The complexity of this is increased by genetic evidence revealing that both genetic and pharmacologic inhibition of any branch can either sensitize cells to apoptosis or delay onset of apoptosis. Some of these paradoxical outcomes can be resolved when one considers either the type of stress or the cell/tissue type being stressed. However, such context still requires that downstream effectors of the primary transducers be selectively activated to achieve specific fates. The elucidation of how effectors interpret signals and regulate cell fate requires a detailed understanding of both pro-survival and pro-apoptotic signalling.

Accumulating evidence supports a role for differential regulation of micro-RNAs by the UPR in the regulation cellular fate decisions under conditions of stress. Indeed, all three UPR transducers have been associated with micro-RNA regulation. Thus far, Ire1 activity has been generally associated with inhibition of micro-RNA accumulation through its RNase function. Targeted micro-RNAs include miRs-17, 34a, 96 and 125b. In contrast, PERK-dependent activities appear to be associated with both micro-RNA induction, miR-
211/204, miR-708 (ref. 23) and suppression miR-106-25 (ref. 24). It is of interest that all of the PERK-dependent micro-RNAs have been linked to the pro-survival activities of PERK. In the work described here, miR-216b is demonstrated to accumulate via the coordinated activities of both PERK and Ire1. Strikingly, in contrast to other PERK-regulated micro-RNAs, miR-216b contributes to UPR-associated apoptosis through regulation of a newly identified target, c-Jun. Although c-Jun, a component of the AP-1 transcription factor64, has not previously been demonstrated to contribute to gene expression patterns associated with ER stress, activation of Jun kinase pathway (JNK), was previously linked with Ire1-dependent activation of apoptotic signalling kinase-1 (refs 59,60). Accordingly, activation of JNK is associated with increased apoptosis, a consequence of JNK-dependent inactivation of Bcl-2 (refs 61,62). Induction of miR-216b is not associated with alterations in JNK signalling; however, it also contributes to increased apoptosis through downregulation of c-Jun.

miR-216b accumulation peaks several hours into the stress response, suggesting it functions as cells commit either to survival or to cell death. This conclusion is supported by a number of experiments. First, miR-216b expression is directly dependent upon CHOP activity, another pro-apoptotic factor. Second, reduction in miR-216b activity increases cell resistance to UPR-dependent cell death. Third, overexpression of a miR-216b increases apoptosis. Finally, we demonstrate that modulation of c-Jun, a bona fide target of miR-216b during ER stress, also has the expected outcome on cell fate during an ER stress response including loss increasing cell death during ER stress.

One of the more vexing issues associated with UPR-triggered apoptosis has been the CHOP transcription factor activity. Genetic evidence from both cell and animal models has revealed CHOP to be a pro-apoptotic transcription factor during cellular stress, whether ER stress or DNA damage62,63. Several mechanisms have been suggested to mediate the pro-apoptotic activity of CHOP. These include suppression of anti-apoptotic Bcl-2 expression64, which phosphorylates and induces CHOP expression, but also depend upon ASK1-dependent activation of p38MAPK, which phosphorylates and induces CHOP65, demonstrating key points of Ire1 and PERK convergence on the establishment of an apoptotic fate under high stress conditions. Our data, which demonstrates CHOP-dependent induction of miR-216b, reveal yet another point of pro-apoptotic convergence wherein miR-216b antagonizes c-Jun accumulation and thereby promotes cell death.

The collective data suggest that CHOP-dependent apoptosis is unlikely to reflect a single functional target. Rather, as with many transcription factors, it is the additive function of several downstream targets that facilitate the establishment of cell fate. In this model, CHOP serves as a focal point downstream of PERK and Ire1 action. CHOP-dependent transcription of targets such as Bcl-2, BIM and miR-216b, the latter of which then targets an increasing number of transcripts, c-Jun as one example, thereby amplifying the pro-apoptotic activity of CHOP (Fig. 7f). As the action of miR-216b is delayed, by virtue of the time needed for its induction and targeting its substrates, this latency likely provides a needed opportunity for cells to repair and adapt or possibly prior to initiating apoptosis.

**Methods**

**Cell culture.** NIH3T3 mouse embryonic fibroblasts (American Type Culture Collection (ATCC)) and murine embryonic fibroblasts 293T human embryonic kidney cells (ATCC) were maintained in DMEM (Mediatech) with 10% heat-inactivated fetal bovine serum (BenchMark). In addition, MEF cells were supplemented with 10% fetal calf serum, 2 mM t-glutamate, 55 μM t-mercaptoethanol (containing 1 mM alamar blue, glutamate, glycoline, and serine), all from Invitrogen. Human U2OS human cervical adenocarcinoma cells (ATCC) were maintained in McCoy’s 5A (Mediatech) with 10% heat-inactivated fetal bovine serum, pen/strep.

**Plasmids and chemicals.** Chop (Plasmid #21914) and Xbp1s (PMV3-Flag: XBP1s) expression vectors were purchased from Addgene. Plasmids encoding sh-chop (TRCN0000039950 and TRCN0000039952) were purchased from Dharmacon. PERK inhibitor is purchased from Selleckchem (GSK-2606414).

**MicroRNA microarray and qPCR.** NIH 3T3 cells were challenged with 2 μg ml−1 Tm for 2, 5 or 10 h. Total RNA was purified using miRNAeasy Mini Kit (Qiagen). Microarray was carried out using GeneChip microRNA Arrays (Affymetrix) and TaqMan microRNA arrays (Applied Biosystems). MicroRNA was reverse transcribed using TaqMan MicroRNA Reverse transcription kit with specific Taqman RT primers for mmu-miR-216b and miR-202 (Cat. # 4427975, Applied Biosystems) as per the manufacturer’s protocol. RNA was hybridized with mmu-miR-216b and miR-202 Taqman primers using no AmpErase UNG Mix (Applied Biosystems). Primers for mouse c-Jun: forward 5′-TTGGGACATCA CCACCTCAGC-3′; reverse 5′-TCTTGGCTATCGACACTTGAGCC-3′. Primers for human c-Jun: forward 5′-GAAGCGGGCCGGCTGATAAAT-3′; reverse 5′-CCCTCC TCTCCAGTCACTC-3′. Primers for human COUL: forward 5′-AGGTTGG TTAGTGGCACAAC-3′; reverse 5′-GGGACCATATCTTACCAGCGGAC-3′. Primers for human DNM1T: forward 5′-CATAGTGACTGTGCGTGGCTG-3′; reverse 5′-CTCCACAGGAGGGAAGCAC-3′. Primers for human MT2: forward 5′-GTATCTG GGAAGGAGGTTG-3′; reverse 5′-TGTTCCTGCTTCGTCGCAAC-3′. Primers for human TMPI: forward 5′-GCTTCTATCCAGAGTACGTA-3′; reverse 5′-TG GAGCTACAGCTGCCTCTC-3′.

**Luciferase assays.** Oligonucleotides containing three copies of the miR-216b seed sequence were cloned into the 3′UTR of the firefly luciferase gene in pRLTK. The 1,000 bp upstream of mmu-miR-216b promoter was amplified by PCR and cloned into pGL4.11 using primers: forward 5′-GATCTGAGGTTTCCTAATCTGGCTT GCT-3′ and reverse 5′-CTGAAAGCTTTTGGTCTGAGTACATA-3′. The mutant mmu-miR-216b promoter mutant was generated using the QuikChange kit (Stratagene). Mutagenic primers used were as follows: forward 5′-GACTGGTGGCTTGTCGATC-3′; reverse 5′-GTGTGGATTGGCTTGGCT-3′. The 3X AP-1 binding sites (TGACTCA) upstream of a minimal promoter fragment containing a TATA box in the luciferase reporter site luciferase67. For luciferase assays, cells were transfected with the constructs and pRLTK (Addgene vector), to normalize for transfection efficiency and protein synthesis, with Lipofectamine plus method and treated with TG (500 nM) for the time indicated.

**Western blot analysis and ChIP.** Cells were harvested by gently scraping and resuspended in EBC buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 0.5% Nonidet P-40) supplemented with protease inhibitors. The cells were disrupted by sonication and then cleared by microcentrifugation at 10,000g for 10 min. Proteins were resolved by SDS–polyacrylamide gel electrophoresis, transferred onto nitrocellulose membrane and analysed by immunoblot. Antibodies used were as follows: PERK rabbit polyclonal antibody (Rockland, #100-401-962; 1,000 dilution), p-eIF2a mouse monoclonal antibody (Cell Signaling, #3897; 1,000 dilution), ATP4 antibody (Cell Signalling, #11815; 1,000 dilution), CHOP (Cell Signalling, #2895; 1,000 dilution), Ire1α (Cell Signalling, #3294; 1,000 dilution), c-Jun (Cell Signaling, #9661; 1,000 dilution), LUNB (Cell Signaling, #3733; 1,000 dilution), actin (Sigma, #A2228; 1,200,000 dilution), lamin A (Abcam, ab26300; 1,000 dilution), cleaved caspase 3 (Cell Signaling, #9661; 1,000 dilution), PARP (Sigma, #P7605; 1,000 dilution). Antibody binding was visualized by enhanced chemiluminescence (Perkin Elmer). Unscanned copies of western blots are provided in Supplementary Fig. 6.

**Chromatin was prepared using the truChIP Low Cell Chromatin Shearing Kit (Covaris) and sheared into 200–700 bp fragments using a Covaris S2 instrument (Covaris) and sheared into 200–700 bp fragments using a Covaris S2 instrument.**
5′-CAGTTGGCTTATGGACGACGAGC-3′, reverse: 5′-GTCATCTGGAGCCTGCC-3′; primer 1: 5′-TCTCTTCTCATGTTGCCGACC-3′, reverse primer 2: 5′-GGGCCAGTACATCTACAA-3′, reverse 2: 5′-CCTCCTGGTGGCAGAATA-3′; the primers for ATF4 binding to CHOP promoter, forward: 5′-AGCCCTTGGGGAGGACAGA-3′, reverse: 5′-AACAGGCTTACGCTTCGTT-3′.

**Mir-216b cleavage assay.** In vitro cleavage reactions were performed by using γ-32P-ATP labeled oligonucleotide. Xbp-1 mini (5′-CUGAAGGUCCAGACGUCCAC-3′) or miR-216b (5′-AACUUCCUCGAGCCAGAUAUGG-3′; Integrated DNA Technologies) was labeled with γ-32P-ATP using T4 polynucleotide kinase (New England Biolabs). The labelled Xbp-1 and miR-216b were purified with G-25 Sephadex mini column (Roche, #1127390001). One microgram recombinant Ire1α (aa 466–955–977; Sino Biological Inc.) was incubated with 1 pmol of 32P-ATP labeled Xbp-1 mini or miR-216b in cleavage buffer (20 mM HEPES, pH 7.5, 50 mM NaCl, 1 mM dithiothreitol, 1 mM ATP) at 37 °C for 2 h. RNA cleavage products were resolved on a urea-19% polyacrylamide gel electrophoresis without phenol/chloroform extraction and then exposed to Blue Autoradiol Film for 20–40 min.

**Clonogenic survival assay.** Clonogenic assays were performed in U2OS stable cells69. Briefly, cells were plated at 2 × 104 cells per 60 mm plates. Twenty-four hours later, they were exposed to 400 mM TG or 2.5 μg ml−1 Tm for the indicated times, then returned to growth medium. Fresh media were replenished every 2 days. Viable colonies were stained with 3 ml of 0.5% Giemsa (v/v) for 5–10 min at room temperature at 12 days. Colony counts were normalized to plating efficiency and represented as fraction surviving compared with control (dimethylsulfoxide).

**Retroviruses and stable U2OS cells.** 293T cells were transfected with PMDL, VSVG, REV and pEZX containing miR-216b mimic (MmiR-3190-MR03, GeneCopoeia) or anti-miR-216b (miR-AN0324-AM03, GeneCopoeia) or pEZX empty vector (mR-001-MR03, GeneCopoeia) as control using Lipofectamine Plus (Invitrogen). Viral supernatants were harvested 48 h after transfection, and were used to infect U2OS cells in the presence of 10 μg ml−1 polybrene. Retroviruses coding c-Jun or vector control were produced by co-transfection of ectopic helper retrovirus plasmid and pBabe vector into 293T cells70. Selection to create stably A-miR-216b overexpressing cells was conducted with puromycin at 1 μg ml−1.

**Sephadex mini column.** One microgram recombinant Ire1α (aa 466–955–977; Sino Biological Inc.) was labelled with 32P-ATP using T4 polynucleotide kinase (New England Biolabs). The labelled Ire1α was separated on a Sephadex mini column (Roche, #11273990001) with 20 mM HEPES, pH 7.5, 50 mM NaCl, 1 mM dithiothreitol, 1 mM ATP. The Evan’s blue dye was added to the eluate to confirm protein purity.

**Cloning of mammalian Ire1 reveals diversity in the ER stress sensor.** Cloning of mammalian Ire1 reveals diversity in the ER stress sensor. The mammalian ER stress sensor Ire1α is a member of the IRE1 family of stress-responsive genes. The mammalian ER stress sensor Ire1α is a member of the IRE1 family of stress-responsive genes. The mammalian ER stress sensor Ire1α is a member of the IRE1 family of stress-responsive genes. The mammalian ER stress sensor Ire1α is a member of the IRE1 family of stress-responsive genes. The mammalian ER stress sensor Ire1α is a member of the IRE1 family of stress-responsive genes.
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Author contributions
J.A.D. designed the experiments, analysed the data and wrote the manuscript. Z.X. performed the experiments, analysed the data and wrote the manuscript; Y.B. and J.A.D. designed the experiments, analysed the data and wrote the manuscript. Z.X. provided the intellectual input and reagents.

Additional information
Accession codes: All microarray data were deposited and can be accessed via GSE79455.
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