A miR-494 dependent feedback loop regulates ER stress

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Abstract

Defects in stress responses are important contributors in many chronic conditions including cancer, cardiovascular disease, diabetes, and obesity-driven pathologies like non-alcoholic steatohepatitis (NASH). Specifically, endoplasmic reticulum (ER) stress is linked with these pathologies and control of ER stress can ameliorate tissue damage. MicroRNAs have a critical role in regulating diverse stress responses including ER stress. Here we show that miR-494 plays a functional role during ER stress. ER stress inducers (tunicamycin & thapsigargin) robustly increase the expression of miR-494 in vitro in an ATF6 dependent manner. Surprisingly, miR-494 pretreatment dampens the induction and magnitude of ER stress in response to tunicamycin in endothelial cells. Conversely, inhibition of miR-494 increases ER stress de novo and amplifies the effects of ER stress inducers. Using Mass Spectrometry (TMT-MS) we identified 23 proteins that are downregulated by both tunicamycin and miR-494. Among these, we found 6 transcripts which harbor a putative miR-494 binding site. We validated the anti-apoptotic gene BIRC5 (survivin) as one of the targets of miR-494 during ER stress. Finally, induction of ER stress in vivo increases miR-494 expression in the liver. Pretreatment of mice with a miR-494 plasmid via hydrodynamic injection decreased ER stress in response to tunicamycin in part by decreasing inflammatory chemokines and cytokines. In summary, our data indicates that ER stress driven miR-494 may act in a feedback inhibitory loop to dampen downstream ER stress signaling. We propose that RNA-based approaches targeting miR-494 or its targets may be attractive candidates for inhibiting ER stress dependent pathologies in human disease.
Introduction

The endoplasmic reticulum (ER) is the site of mRNA translation alongside proper folding and post-translational modifications of proteins destined for secretion or localization to various cellular membrane systems. In addition, the ER is the center of lipid biosynthesis, detoxification, homeostasis of intracellular Ca^{2+} and redox balance. Several pathologies including neurodegenerative diseases, diabetes, atherosclerosis and cancers have attributed ER stress as a critical driver of disease. When the folding capacity of the ER is challenged, the accumulation of un- or mis-folded proteins in the lumen of the ER triggers the unfolded protein response (UPR) (1,2). This response activates a trio of transducers: the PKR-like ER kinase (PERK), inositol requiring enzyme 1 α (IRE1α) and activating transcription factor 6 (ATF6), which work synergistically to control transcriptional and translational programs to either alleviate the burden of unfolded proteins and return to protein homeostasis or initiate apoptosis. It is thought that acute ER stress can trigger feedback mechanisms that protect cells from death by suppressing global translation and increasing ER chaperone levels, whereas persistent UPR activation or chronic unmitigated ER stress leads to increased oxidative stress, inflammation and eventual apoptosis (3,4).

Endothelial cells (ECs) encounter a variety of stressors and stimuli during development and disease (5). Recent studies have implicated ER stress and the UPR as drivers of endothelial dysfunction in cardiovascular disease (6,7). For instance, ER stress is thought to promote both EC inflammation and apoptosis in atherosclerosis (8-10). Furthermore, ER stress has been shown to contribute to vascular dysfunction and cardiac damage in preclinical models of hypertension (11). Similarly, oxidative stress and ER stress pathways have been shown to be interlinked in ECs in metabolic syndromes such as diabetes and non-alcoholic fatty liver disease (NAFLD) (12-16). Therefore, ECs have adopted several mechanisms to regulate cell fate decisions in response to both acute and chronic ER stress (17).
MicroRNAs (miRs) are small non-coding RNAs that are critical regulators of physiological and pathological stress responses (18,19). We, and others have shown that specific miRs regulate EC responses to numerous effectors including: angiogenic growth factors (20,21), hypoxia (22,23), DNA damage (24), and oxidative stress (25). Furthermore, specific miRs have been identified as modulators of ER stress in different models of disease (26-29). Indeed Kassan et al., identified miR-204 as a promoter of ER stress in ECs by targeting the SIRT1 pathway (30). Our previous work identified a cohort of miRs that are induced by DNA damage in ECs (24,31,32). We further characterized one of these miRs, miR-494, as a regulator of endothelial senescence in response to genotoxic stressors (24). In this study, we show that ER stress is a potent inducer of miR-494 likely via ATF6. Surprisingly, we find that miR-494 operates in a feedback loop to dampen the ER stress response potentially by targeting the anti-apoptotic protein survivin. Our in vivo experiments suggest miR-494 pretreatment diminishes tunicamycin-induced ER stress in the liver. Overall, our studies illuminate a novel function for miR-494 and open new avenues for further investigations into mechanisms by which miRs modulate stress responses.

**Methods**

**Cell culture and reagents**

HUVECs (Cat: NC9946677, Lonza) were cultured in EGM-2 media (Cat: CC-3162, Lonza) supplemented with 10% fetal bovine serum (Cat: FB-21, Lot# 649116, Hyclone) and were maintained at 37°C with 5% CO₂. Cells used for experiments were low passage number (3-8). Tunicamycin (Cat: T7765, Sigma) was dissolved in DMSO (Sigma).

**Cell transfection**

HUVECs (50% confluence) were transfected with 50nM mimic/inhibitor RNA using the lipofectamine RNAiMax reagent (Cat: 13778-150, Invitrogen) according to the manufacturer’s
instructions. Specifically, mirVana miR mimic negative control (Cat: 4464061, Ambion), hsa-
mir-494-3p mimic (MIMAT002816) (Cat 4404066, Assay ID MC12409, Ambion), mirVana miR
inhibitor negative control #1 (Cat 4464076, Ambion), hsa-mir-494-3p inhibitor (MIMAT002816)
(Cat 4464084, Assay ID MH12409, Ambion). Negative Control LNA mimic (Cat: YM00479903,
Qiagen), hsa-miR-494-3p miRCURY LNA™ microRNA Mimic (YM00472106, MIMAT002816,
Qiagen), miR-control (CAT#4464058), miR-494 inhibitor (CAT#4464084/MIMAT0004975), and
miR inhibitor negative control (CAT#4464076) were purchased from Ambion/Life Technologies.

For gene knock down, HUVECs were transfected with 10nM siRNA using the lipofectamine
RNAiMax reagent (Cat: 13778-150, Invitrogen) according to the manufacturer’s instructions.
Specifically, siRNA for ATF6 (Cat: SR307883A, Origene), EIF2AK3 (Cat: SR306267A, Origene),
ERN1 (Cat: SR301457A&B, Origene), siRNA Negative Control (Cat: SR30001, Origene).

**Gene expression**

Total mRNA was isolated from cells and tissues using the miRNeasy Mini Kit (cat: 217004,
Qiagen). Reverse transcription was performed using High Capacity cDNA Reverse
Transcription Kit (Cat: 4368814, Applied Biosystems) according to the manufacturer’s
instructions. Gene expression was measured using real-time quantitative PCR (qRT-PCR) with
TaqMan™ Master Mix II no UNG (Cat 4440049, Thermofisher Scientific) with the following
primer/probe sets: miR-494-3p (Cat: 4427975, Assay ID: 002365), human primary miR-494
(Cat: 4427013, Assay ID: Hs04225959_pri) U6 snRNA (Cat: 4440887, Assay ID: 001973),
sno234 (Cat: 4440887, Assay ID: 001234) GAPDH (Cat 4351370 Assay ID: Hs02758991_g1)
according to the manufacturer’s instructions. SYBR Green qRT-PCR assays were conducted
using PowerUp SYBR Green Master Mix (Cat A25741, Thermofisher Scientific) with primers
shown in Table 1. Fold change was calculated using the $2^{-\Delta\Delta C_T}$ method relative to an internal
control (GAPDH, β-Actin (ActB), sno234 or U6).

Table 1: Oligonucleotide primers used for SYBR Green based qRT-PCR assays

| Primer          | Sequence                              |
|-----------------|---------------------------------------|
| mmu-DDIT3 For   | CCA CCA CAC CTG AAA GCA GAA           |
| mmu-DDIT3 Rev   | AGG TGA AAG GCA GGG ACT CA            |
| mmu-GAPDH For   | GCC TGG TCA CCA GGG CTG C             |
| mmu-GAPDH Rev   | CTC GCT CCT GGA AGA TGG TGA TGG      |
| hu-sXBP1 For    | CTG AGT CCG AAT CAG GTG CAG           |
| hs-sXBP1 Rev    | ATC CAT GGG GAG ATG TTC TGG           |
| hs-Total XBP1 For| TGG CCG GGT CTG CTG AGT CCG           |
| hs-Total XBP1 Rev| ATC CAT GGG GAG ATG TTC TGG           |
| hs-DDIT3 For    | AGA ACC AGG AAA CGG AAA CAG A         |
| hs-DDIT3 Rev    | TCT CCT TCA TGC GCT GCT TT            |
| hs-ERN1 For     | TGC TTA AGG ACA TGG CTA CCA TCA       |
| hs-ERN1 Rev     | CTG GAA CTG CTG GTG CTG GA            |
| hs-EIFAK3 For   | AAT GCC TGG GAC GTG GTG GC            |
| hs-EIFAK3 Rev   | TGG TGG TGC TTC GAG CCA GG            |
| hs-AFT6 For     | ATG AAG TTG TGT CAG AGA ACC           |
| hs-AFT6 Rev     | CTC TTT AGC AGA AAA TCC TAG           |
| hs-DUT For      | TGC ACA GCT CAT TTG CGA ACG G         |
| hs-DUT Rev      | CCA GTG GAA CCA AAA CCT CCT G         |
| hs-MINA For     | ACT TTG GCT CCT TGG TTG G             |
| hs-MINA Rev     | CCC GGC TTC AGC ATA AAC               |
| hs-DHFR For     | CAT GGT TGG TTC GCT AAA CTG C         |
Nanostring® Immune Profiling

We used the Nanostring PanCancer Mouse IO360™ Gene Expression Panel containing 770 genes involved in the tumor, surrounding microenvironment and immune response in cancer. Livers were homogenized and total RNA was isolated using the miRNeasy Mini Kit (cat: 217004, Qiagen). 100ng of RNA was used to perform the generate the expression profile as per the manufacturer’s instructions. Data was analyzed using the nSolver™ Analysis software. Raw data was normalized to a panel of housekeeping genes.

Immunofluorescence

For immunofluorescence analysis, HUVECs were seeded (12,000 cells/well) on coverslips in a 24-well plate and transfected with mimic/inhibitor for 24h as described above. The following day, the media was aspirated and replaced with fresh media containing tunicamycin (10µg/mL TCN, Sigma) or DMSO (Sigma) for 24-48h. The media was aspirated and cells were rinsed in ice cold
PBS for 2 min followed by fixation in 4% paraformaldehyde diluted in PBS (Cat: NC9658705, FisherSci) for 10 min at room temperature (RT). Coverslips were rinsed in PBS three times and incubated in serum free DAKO Protein Block (X0909, DAKO) for 1h at RT. Next, coverslips were incubated overnight at 4°C in anti-survivin primary antibody diluted in PBS/5% BSA. After rinsing in PBS three times, the coverslips were incubated in the dark with Alexa Fluor® 488 goat anti-rabbit secondary antibody diluted in PBS/5% BSA. Refer to table 2 for antibody information and dilutions. After rinsing in PBS three times, the coverslips were mounted onto glass slides with Aqua-Poly/Mount (Cat No. 18606, Polysciences Inc). Slides were left to cure overnight at RT prior to imaging. Coverslips were imaged using a Nikon/Yokogawa CSU-W1 Spinning Disk Confocal Microscope. All imaging settings remained constant throughout the imaging sessions. The percentage stained area was analyzed using the open-access Image J software (NIH).

Table 2: Antibodies used in this study

| Antibody          | Source                  | Identifier | Concentration |
|-------------------|-------------------------|------------|---------------|
| Survivin          | Cell Signaling Technology | 2808       | 1:25          |
| CHOP              | Novus                   | NB600-1335 | 1:250         |
| XBP1-s            | Cell Signaling Technology | 83418     | 1:25          |
| GINS4             | Novus                   | NBP2-16659 | 1:25          |
| Alexa Fluor® 488  | Cell Signaling Technology | 4412       | 1:100         |
| (goat anti-rabbit)|                         |            |               |

Simple western blot

HUVECs were seeded in 6 well plates (200,000 cells/well) and transfected as described above. After 24h, cells were treated with 10µg/mL TCN for 48h. After treatment, the media was aspirated and the cells were washed twice in ice cold PBS and lysed directly in the plate in RIPA buffer (cat: PI89900, Pierce) containing Protease Inhibitor Mini Tablets (1 tablet/10mL RIPA buffer, Cat: A322953, Pierce) with phosphatase inhibitor cocktail 2 & 3 (1:1000, Cat:
P5726 and P0044) for 15 min on ice. Lysates were rotated at 4°C for 30 min-2h and then centrifuged at 12,000 x g at 4°C for 40 min. The supernatant was collected and protein concentration was determined using the Pierce BCA Protein assay kit (Cat No. 23225). Samples were diluted to 0.75µg/mL with 1× Sample Buffer (ProteinSimple). Protein quantification was performed using a 12–230 kDa 25 lane plate (PS-MK15; ProteinSimple) in a ProteinSimple Wes Capillary Western Blot analyzer according to the manufacturer’s instructions. The standard Simple Western protocol was altered to increase sample load time to 13s and a separation time of 33 min.

Animal experiments

For in vivo experiments, male C57Bl/6 mice (aged 12-20 weeks) were purchased from Jackson labs. Animals were housed in a controlled animal facility with access to normal rodent chow and water ad libitum. All experimental protocols were approved by the Institutional Animal Care and Use Committee (IACUC) at Oregon Health and Science University (OHSU). For in vivo studies, the LentimiRa-GFP-hsa-miR-494-3p Vector (Cat: mh10739, Applied Biological Materials, abmgood.com) and pLenti-III-miR-GFP Control Vector (Cat: M001, Applied Biological Materials) were purified using the EndoFree Plasmid Maxi Prep Kit (Cat: 118379, Qiagen). Plasmids (20-30µg in PBS) were injected using a hydrodynamic injection method into the tail vein with a volume of 1.5-2mL/animal in 7-10s as described (33,34). After 24h, animals were injected i.p. with tunicamycin (2 µg/g Cat: T7765, Sigma) which was prepared by dissolving the compound in a small volume of DMSO (100µL, Sigma) and then diluting in 150mM Sucrose (Cat: 57-50-1, FisherSci) in dH₂O. After 6-20h, livers were harvested and total RNA was isolated using the miRNeasy Mini Kit (Cat: 217004, Qiagen).

TMT-Mass Spectrometry

TMT labeling and mass spectrometry were performed by the OHSU proteomics core facility as described in detail elsewhere (35). Briefly, HUVECs were treated with microRNAs or
tunicamycin as described above. After 24h-48h, samples were lysed in 50 mM triethyl ammonium bicarbonate (TEAB) buffer (50 µg of protein/ sample) followed by a Protease Max digestion, a microspin solid phase extraction and TMT labeling. Multiplexed TMT-labeled samples were separated by two-dimensional reverse-phase liquid chromatography. Tandem mass spectrometry data was collected using an Orbitrap Fusion Tribrid instrument (Thermo Scientific). RAW instrument files were processed using Proteome Discoverer (PD) (Thermo Scientific). Searches used a reversed sequence decoy strategy to control peptide false discovery and identifications were validated by Percolator software. Search results and TMT reporter ion intensities were processed with in-house scripts. Differential protein abundance was determined using the Bioconductor package edgeR.

**Statistical Analysis**

All statistical analysis was performed using Prism software (GraphPad Software, San Diego, CA). Differences between pairs of groups were analyzed by Student’s t-test. Comparison among multiple groups was performed by one-way ANOVA followed by a post hoc test (Tukey’s or Holm-Sidak). In the absence of multiple comparisons, Fisher’s LSD test was used. Values of n refer to the number of experiments used to obtain each value. For mouse studies where the data was not normally distributed, we used two-tailed Mann-Whitney U test. Values of p ≤ 0.05 were considered significant.

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Author Contributions

N.C, C. E-D and S.A. designed experiments, N.C, C.E-D performed experiments, analyzed the data, N.C and S.A. wrote the manuscript.
Results:

ER stress induces miR-494 in vitro

We previously showed that miR-494 is responsive to radiation and chemical inducers of genotoxic stress and functions to increase endothelial senescence during DNA damage responses (24). Given the intricate relationship between radiation, oxidative stress and ER stress (14,17), we asked if ER stress affected miR-494 expression and function. First, we confirmed a robust ER stress response to known inducers, tunicamycin (TCN) and thapsigargin (TG), in human umbilical vein endothelial cells (HUVECs) by measuring the level of the transcription factors spliced XBP1 (mRNA: sXBP1, protein: XBP1s) and DDIT3 (CHOP), which are well characterized markers of the ER stress response (Fig. 1A-B). We observed that TCN significantly increased the levels of mature miR-494 (Fig. 1C) and to a lesser extent, the primary unprocessed miR-494 transcript (Fig. 1D). Similarly, TG also induced sXBP1 and DDIT3 in parallel with the primary and mature forms of miR-494 (Supplementary Fig. 1). We also saw a comparable increase in both ER stress and the primary miR-494 transcript in another EC line (Human Microvascular Endothelial cells - HMVECs) (Supplementary Fig. 2).

A triumvirate of transducers (ATF6, IRE1α & PERK) co-ordinate ER stress response programs to promote cell recovery, or in the case of excessive or chronic stress, initiate apoptosis. We asked which of these transducers was responsible for miR-494 induction in response to ER stress using siRNAs. We found that knockdown of ATF6 but not IRE1α (ERN1) or PERK (EIF2AK3) significantly decreased the induction of miR-494 6h after TCN treatment (Supplementary Fig. 3). These data indicate that ER stress induces the expression of miR-494 in an ATF6 dependent manner.

miR-494 diminishes ER stress

To understand the functional relevance of miR-494 during ER stress, we performed gain- and
loss-of function experiments utilizing miR-494 mimics or inhibitors, respectively. Pretreatment of HUVECs with miR-494 mimic suppressed the TCN based induction of ER stress responsive genes DDIT3 (Fig. 2A). Conversely, inhibition of miR-494 robustly increased levels of DDIT3 (Fig. 2B). Similarly, miR-494 mimic decreased levels of sXBP1 (Fig. 2C) whereas, inhibition of miR-494 increased sXBP1 mRNA both de novo and in combination with TCN (Fig. 2D). We validated that the miR-494 mediated decrease in these transcription factors also persisted at the protein level (Fig. 2E). Taken together, our data suggest that miR-494 diminishes the induction of ER stress-associated transcription factors and plays a protective role in HUVECs during ER stress.

**Mass spectrometry identifies putative targets of miR-494 relevant for ER stress**

miRs typically are thought to regulate numerous targets in a context dependent manner. We have previously shown that miR-494 targets the Mre11a, Rad50, Nbn (MRN) complex in the DNA damage repair pathway in response to genotoxic stressors (24). To identify the targets relevant for miR-494 in the context of ER stress, we undertook a proteomics-based approach. We used Tandem mass tag-based mass spectrometry (TMT-MS) to compare changes in the proteome with either miR-494 mimic treatment or TCN treatment. We found 23 proteins were commonly downregulated between the TCN and miR-494 treatment conditions (Fig. 3A). Among these, we identified six proteins whose mRNAs harbored putative miR-494 binding sites in their 3'UTRs as predicted by the TargetScan algorithm. We validated the expression of these six genes using qRT-PCR (Fig. 3B and Supplementary Fig. 4A-D). All six targets were validated and demonstrated a substantial decrease in expression in response to miR-494 gain-of-function activity. Conversely, inhibition of miR-494 restored BIRC5, GINS4, MINA and to a lesser extent, UHRF1 mRNA levels in HUVECs treated with TCN (Supplementary Fig 4E). Since BIRC5 and GINS4 were consistently regulated in a miR-494 dependent fashion, we chose to further validate these two targets at the level of protein expression. We found survivin (gene: BIRC5),
and GINS4 protein levels were significantly decreased upon treatment with miR-494 mimic alone, TCN treatment alone, and the sequential combination of miR pre-treatment (24h) followed by TCN (24h) (Fig. 3C). Finally, we used immunofluorescence staining and confocal microscopy to evaluate survivin expression in HUVECs. Consistent with our western blot data, we observed a significant decrease in the amount of survivin in cells transfected with either miR-494 mimic or TCN treatment alone or in the sequential combination (Fig. 3D). Multiple studies have demonstrated that miR-494 binds to the 3'UTR and leads to subsequent degradation of the survivin (BIRC5) transcript (36-38). Our observations support these data and further demonstrates this relationship in ECs during ER stress.

**miR-494 pretreatment ameliorates ER stress in mouse liver**

ER stress is thought to be a significant driver of liver diseases including hepatic fibrosis, non-alcoholic fatty liver disease (NAFLD) that can progress to steatohepatitis (NASH) and hepatocellular carcinoma (3,39,40). Endothelial dysfunction plays a critical role in the pathology of NAFLD and NASH (41). Indeed, liver sinusoidal endothelial cells (LSECs) can function to provoke inflammation and fibrosis in NAFLD (16). To delve further into the function of miR-494 during ER stress *in vivo*, we utilized a well characterized murine model of NASH (42). It has been shown that injection of TCN (2µg/g) increases acute ER stress in the liver with upregulation of CHOP and spliced XBP1 (42). We observed that TCN treatment induced endogenous miR-494 expression in mouse livers (Fig. 4A). To address the relationship between miR-494 and ER stress, we used hydrodynamic injection of plasmid DNA to transiently increase miR-494 levels in the liver. Compared to the GFP control plasmid, miR-494 plasmid injection significantly increased the levels of miR-494 (Fig. 4B). Importantly, when followed by TCN treatment 24h later, this increase in miR-494 levels was sufficient to decrease ER stress in the livers as assessed by DDIT3 levels (Fig. 4C).

ER stress induction by TCN treatment has been shown to induce inflammation in the liver,
which is a hallmark of NASH (3,7). To explore the mechanisms by which miR-494 may impact the activity of TCN in vivo, we utilized a Nanostring® IO360 mouse profiling array to evaluate the immune response and inflammation in these animals. Compared to the plasmid control, miR-494 treated mice exhibited a significant decrease in several immune response and inflammatory genes in the liver (Supplementary Fig. 5A). Gene Ontology analysis identified cytokine response, immune cell migration and immune cell adhesion to vascular endothelium as the most significantly downregulated pathways in the miR-494 treatment group (Supplementary Fig. 5B).

**Discussion:**

Here we show that endoplasmic reticulum (ER) stress is a potent inducer of miR-494 in endothelial cells (ECs) in vitro and in murine liver in vivo. Our data indicates that the ER stress-associated increase in expression of miR-494 occurs in an ATF6 dependent manner. Upon induction, miR-494 decreases the level of survivin (BIRC5)—an anti-apoptotic protein—amongst other genes important in DNA replication and cell proliferation. We show that miR-494 gain-of-function diminishes the magnitude of the unfolded protein response (UPR) induced by tunicamycin (TCN) in both in vitro and in vivo models of acute ER stress. Functionally, mice pre-treated with a miR-494 overexpression plasmid exhibit a decrease in numerous inflammatory genes and cytokines in response to TCN in the liver. This miR-494 mediated decrease in ER stress appears to impact inflammation. Overall, this report shows that miR-494 downregulates critical targets in cell proliferation and survival which function to dampen the ER stress response by mechanisms which have yet to be revealed (Fig. 4D).

Many chronic human diseases have an established link within the ER stress response signaling cascades. Mechanistically, ER stress and the unfolded protein response (UPR) are triggered by the accumulation of misfolded proteins (2,43). The UPR is initiated by the dissociation of the molecular chaperone protein BiP from the three sensors, PERK, IRE1α and ATF6. Dissociation
of BiP activates these proteins resulting either oligomerization or export from the ER. These activated sensors transduce signals to initiate pathways which work in a coordinated manner to halt protein translation, increase ER chaperone levels and clear misfolded proteins. Known chemical inducers of ER stress, TCN and thapsigargin (TG) generate a robust UPR in multiple models of disease including atherosclerosis and NASH (42,44). TCN is a toxic aminoglycoside antibiotic which triggers ER stress by preventing core oligosaccharide addition to nascent polypeptides, thereby blocking proper protein folding. Numerous studies have shown that TCN stimulates all three of the UPR signaling pathways causing an upsurge in transcription factor activity (eg. CHOP, XBP1 splicing), global inhibition of translation (eIF2α phosphorylation), and ATF6 translocation to the Golgi apparatus. We found that TCN treatment of ECs induced CHOP transcription and XBP1 splicing, which are indicative of an appropriate ER stress response. Further, TCN consistently generated a rapid and robust upregulation of miR-494 in multiple EC cell lines (Fig. 1 and Supplementary Fig. 2). TG, a non-competitive inhibitor of the Sarco/Endoplasmic Reticulum Calcium ATPase (SERCA) disrupts calcium homeostasis in the ER. In our experiments, TG treated ECs experienced similar signaling patterns, with an increase in both spliced XBP1, CHOP transcription and induction of miR-494 (Supplementary Fig 1). Interestingly, TCN but not TG induced a biphasic expression of mature miR-494 with a slight increase at 1h and a more significant (~30 fold) increase at 6h. Moreover, primary miR-494 transcript expression mirrored that of the mature transcript in TG treatment but not TCN (Fig. 1 & Supplementary Fig.1). It is unclear if these differences reflect differential regulation of miR processing pathways by these two inducers. Indeed, the canonical miR processing protein Dicer has been shown to localize and interact with proteins in the ER (45). Therefore, it is possible that ER stress pathways can differentially impact miR transcription and processing and it will be pertinent to investigate the impact of ER stress in the setting of EC-associated induction of miR-
Our data from gain-and loss-of-function experiments show that miR-494 decreases ER stress in ECs and in mouse liver (Fig. 2, 4). These results bolster other reports which also demonstrate miR-mediated responses to ER stress in different tissues (26,46-49). While miRs have a recognized role in liver disease (50), our studies are the first to show the ER stress reducing activity of miR-494 in the context of liver disease (NASH). miR-494 is known to be oncogenic and drives tumor progression and drug resistance in specific cancer types including colorectal cancer (51) and hepatocellular carcinoma (52,53). Furthermore, miR-494 has been shown to attenuate ischemia reperfusion injury in the liver by regulating the PI3K/Akt signaling pathway (54). We previously reported a unique role for miR-494 as a mediator of endothelial senescence by decreasing the MRN DNA repair protein complex. Other studies have shown that miR-494 is involved in vascular inflammation in atherosclerosis (55). Our data indicates a novel function of miR-494 in a complex pathological process.

Using a proteomics approach, we identified six putative miR-494 targets that are relevant during the ER stress response in ECs. Of these, survivin (BIRC5) has been previously shown to be a bona fide target of miR-494 in different disease models (37,38). Interestingly, the cross-talk between ER stress and survivin has been shown to downregulate inflammatory genes in a mouse model of chronic ER stress in the colon (56). Consistent with our observations of a miR-mediated reduction in ER stress combined with a decrease in survivin (BIRC5) and markers of inflammation, Gundamaraju et al., demonstrated that pharmacological inhibition of survivin is comparable to ER stress inhibition and attenuates inflammatory gene expression (56).

We used a Nanostring Mouse PanCancer™ profiling array to query the in vivo response to miR-494 pre-treatment prior to induction of ER stress. Our results reveal that a number of inflammatory genes are downregulated in the liver with miR-494 plasmid pre-treatment. Very few of the array-identified genes harbor miR-494 binding sites and therefore, the mechanism(s)
by which miR-494 diminishes ER stress and subsequent inflammation is unclear. Further mechanistic complexity arises from recent reports which indicate miR-494 can be found in exosomes from patients with cirrhosis (57,58). Pertinent to this body of work, the relative contributions of miR-494 from hepatocytes in comparison to liver ECs requires further investigation. In this setting, it is conceivable that exosomal miR-494 from sinusoidal or other ECs in the liver can orchestrate inflammation and mediate paracrine effects during ER stress.

In summary, our work shows that miR-494 is induced during acute ER stress and functions to attenuate ER stress \textit{in vitro} and \textit{in vivo}. Our observations elucidate a new potential mechanistic role for miR-494 in the ER stress response pathway in ECs and likely in other cell types. These studies offer opportunities to inspire new hypotheses to understand the link between miR activity and the response to stressors which influence cell fate decisions in many human diseases.
Figure Legends

Figure 1: ER stress induces expression of the primary and mature forms of miR-494.
Relative mRNA expression of ER stress responsive genes as measured using qRT-PCR. A) spliced XBP1 and Total XBP1 and B) DDIT3 (CHOP) levels in HUVECs treated with 5µg/mL tunicamycin (TCN) over a time course. Relative expression of C) primary miR-494 (pri-miR-494) and D) mature miR-494 in HUVECs treated with TCN over a time course. Gene expression is normalized to GAPDH or U6 and mean fold change compared to vehicle control or time 0h is shown. Graphs are representative of one biological replicate from three independent replicates where values indicate mean ± standard deviation.

Figure 2: miR-494 is a negative regulator of ER stress in vitro.
Relative mRNA expression of ER stress responsive genes as measured by qRT-PCR. A-B) DDIT3 (CHOP), C-D) spliced XBP1 in HUVECs treated with 10µg/mL TCN 48h after transfection with A, C) miR-494 mimic or B, D) miR-494 inhibitor. Gene expression is normalized to GAPDH and mean fold changes compared to control treatments are shown. E) Simple Western blot analysis of HUVECs transfected with miR-494 mimic or control (24h) followed by TCN (10µg/mL) for 24h. Vertical dotted red line indicates non-adjacent lanes. Graphs are mean + SEM fold changes of biological replicates from n=3 independent experiments. * P<0.05, ** P< 0.01, ***P<0.001 by two-tailed Student’s T-test.

Figure 3. miR-494 regulates target genes in cell survival and DNA replication.
A) Venn diagram showing the number of downregulated target proteins in a Tandem Mass Tag labelled Mass Spectrometry profile from HUVECs treated with TCN or transfected with miR-494 compared to vehicle treatment or control miR respectively. B) Fold-change (compared to
respective controls) of protein or mRNA levels as assessed by Mass Spectrometry or qRT-PCR respectively for the 6 targets that were downregulated in both TCN and miR-494 groups. All 6 targets harbor miR-494 binding sites in their 3'UTRs. C) Representative Simple Western blot showing survivin and GINS4 levels in HUVECs 24h after miR-494 transfection followed by TCN treatment (24h). Right panels show quantitation of biological replicates. * p<0.05, ** p< 0.01, by two-tailed Student’s T-test. D) Immunofluorescence imaging showing survivin expression in HUVECs 24h after miR transfection and/or TCN treatment. Right panel shows Survivin fluorescence intensity quantifications via Image J in each cell. Each dot represents individual cells. Scale bar in white = 50 µm. * P<0.05, ** P< 0.01, ***P<0.001 by one-way ANOVA with post-hoc Tukey’s correction.

Figure 4: miR-494 regulates ER stress in vivo.

A) miR-494 levels measured by qRT-PCR from whole liver harvested from male C57BL/6 mice injected i.p. with TCN (2µg/g, n=11) or vehicle control (n=10) for 6h. Bars show mean + SEM. P-values are from a Mann-Whitney U-test. B) miR-494 levels and C) ER stress levels measured by DDIT3 mRNA were assessed by qRT-PCR from whole livers harvested from male C57BL/6 mice (n=5-8 per group) after hydrodynamic injection of miR-494 or GFP control plasmids (20-30µg/animal, 48h) followed by TCN or vehicle treatment (24h). Livers of mice treated with Tunicamycin 24h after hydrodynamic injection of miR-494 or a control GFP plasmid. P=0.01 via one-way ANOVA with a Holm-Sidak’s post-hoc correction. Each dot represents an individual mouse. D) Schematic showing ER stress induction of miR-494 targets BIRC5 and functions to dampen ER stress.
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Chatterjee et al. Figure 2

(A) DDIT3

(B) DDIT3

(C) sXBP1

(D) sXBP1

(E) CHOP (DDIT3) protein and XBP1s protein levels with miR-494 mimic and control mimic treatment.
A) Endogenous miR-494 in mouse liver

miR-494 levels (Fold change vs U6 RNA)

|        | Control | Tunicamycin |
|--------|---------|-------------|
| miR-494 levels | p=0.03  |             |

B) miR-494 levels in liver

|        | GFP     | miR-494 | Vehicle | GFP | miR-494 | Tunicamycin |
|--------|---------|---------|---------|-----|---------|-------------|
| Relative Expression | p=0.01  |         |         |     |         |             |

C) ER Stress in liver: DDIT3

Relative Expression (vs GAPDH)

|        | GFP miR-494 | Vehicle | GFP miR-494 | Tunicamycin |
|--------|-------------|---------|-------------|-------------|
| p=0.01 |             |         |             |             |

D) ER Stress

- ATF6
- DDIT3 (CHOP)
- miR-494
- BIRC5 (Survivin)
- Cell death?
A miR-494 dependent feedback loop regulates ER stress

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Supplementary Figure 1: Thapsigargin induced ER stress drives miR-494 expression.
Relative mRNA expression of A) sXBP1 and B) DDIT3 (CHOP) mRNAs and C) primary and mature miR-494 in HUVECs treated with 0.1µM Thapsigargin. Gene expression is normalized to GAPDH or U6 and mean fold change compared to vehicle control or time 0h is shown. Graphs are representative of 1 of 3 biological replicates. Values indicate mean ± standard deviation.
Supplementary Figure 2: ER stress induces expression of miR-494 in human microvascular ECs. A. Relative mRNA expression of DDIT3 (CHOP) in HMVECs treated with 5µg/mL Tunicamycin. B) Relative expression of primary miR-494 treated with Tunicamycin. Gene expression is normalized to GAPDH and mean fold change compared to vehicle control or time 0h is shown. Graphs are representative of 1 biological replicate from 3 independent replicates where values indicate mean ± standard deviation. * P<0.05 using two-way ANOVA with Fisher’s LSD test.
Supplementary Figure 3: ER stress via ATF6 induces expression of miR-494.

A) HUVECs were transfected with indicated siRNAs. After 48h, cells were treated with vehicle or Tunicamycin (10µg/mL). Relative expression of primary miR-494 normalized to GAPDH is shown. Mean fold change compared to vehicle control is shown. Graphs are representative of 1 biological replicate from 2 independent replicates where values indicate mean ± standard deviation. * p<0.001 using two-tailed Student’s T-test.
Supplementary Figure 4: **miR-494 regulates several target genes in cell survival and DNA replication.**

A-D) Fold-change (compared to respective controls) of mRNA levels as assessed by qRT-PCR for the 4 targets that are downregulated in both Tunicamycin and miR-494 groups in Fig 3B. HUVECs were transfected and treated as described in 2A. Gene expression measured at 48h post treatment. E) HUVECs were transfected with either a control inhibitor or miR-494 inhibitor as shown in 2B. mRNA levels of indicated target genes were evaluated using qRT-PCR. Bars show mean + S.D. of one representative experiment out of three independent biological replicates. * P<0.05, ** P< 0.01, ***P<0.001 by two-tailed Student's T-tests comparing respective control groups.
**Supplementary Figure 5: miR-494 treatment in vivo impacts inflammatory gene expression**

A) Heatmap depicting normalized gene expression in mice treated with miR-494 plasmid (n=2) or a control plasmid (n=2) via hydrodynamic injection followed by Tunicamycin treatment from experiment shown in Fig 4B-C. Gene expression was assessed using a Nanostring IO360 panel containing 770 tumor microenvironment, immune response and inflammation genes. Heatmap depicts genes at least 2 fold downregulated or upregulated compared to control plasmid. B) Gene Ontology analysis based on the whole 770 gene panel using STRING database. All the GO enriched genes derived from the miR-494 downregulated gene cluster.