NADPH oxidase links endoplasmic reticulum stress, oxidative stress, and PKR activation to induce apoptosis

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Introduction

A common feature of many disease processes is pathological cell death leading to tissue dysfunction. Two processes that are becoming increasingly recognized as inducers of pathological cell death are ER and oxidative stress (Chandra et al., 2000; Kim et al., 2008). Many of the diseases that feature ER and oxidative stress are associated with aging and/or obesity, including diabetes, atherosclerosis, renal disease, and neurodegenerative disease, and are becoming epidemic among humans in modern day society (Martínez, 2006; Kregel and Zhang, 2007; Kim et al., 2008). Nonetheless, progress in translating this understanding into useful therapeutic strategies has been disappointing. As one example, there is good evidence that the clinical progression of advanced atherosclerosis involves both ER and oxidative stress (Glass and Witztum, 2001; Tabas, 2010). However, ER stress involves many signaling pathways with differing pathological and physiological functions (Malhotra and Kaufman, 2007b; Ron and Walter, 2007), which makes targeting this process for atherosclerosis complex and challenging. Likewise, there are many sources of cellular oxidative stress in atherosclerotic lesional cells, and human trials using vitamin E as an antioxidant have failed to suppress the incidence of cardiovascular disease (Kris-Etherton et al., 2004; Williams and Fisher, 2005). Similar issues exist with other diseases driven by ER stress–induced apoptosis and oxidative stress.

An important approach to this problem is to gain a more in-depth understanding about molecular and cellular mechanisms and links between ER stress and oxidative stress, particularly in the process of cell death and in settings relevant to disease processes (Malhotra and Kaufman, 2007a). In this study, we focus on a previously elucidated ER stress–apoptosis pathway whose molecular components have been identified and documented to cause apoptosis in vivo. We now show that nicotinamide adenine dinucleotide phosphate reduced oxidase (NOX) and NOX-mediated oxidative stress are induced by this pathway and that apoptosis is blocked by both genetic deletion of the NOX subunit NOX2 and by the antioxidant N-acetylcysteine. Unexpectedly, NOX and oxidative stress further amplify CCAAT/enhancer binding protein homologous protein (CHOP) induction through activation of the double-stranded RNA–dependent protein kinase (PKR). In vivo, NOX2 deficiency protects ER-stressed mice from renal cell CHOP induction and apoptosis and prevents renal dysfunction. These data provide new insight into how ER stress, oxidative stress, and PKR activation can be integrated to induce apoptosis in a pathophysiologically relevant manner.
investigate these mechanisms using cultured macrophages as an in vitro model and the kidney in ER-stressed mice as an in vivo model. The macrophage model is relevant to advanced atherosclerosis (Tabas, 2010), and the kidney model is relevant to several renal diseases (Kitamura, 2008). With regard to macrophages, ER stress, often in combination with other proapoptotic hits, leads to apoptosis in part through the ER stress effector CCAAT/enhancer binding protein homologous protein (CHOP; Feng et al., 2003a; Thorp et al., 2009; Tabas, 2010). An important mechanism linking CHOP with apoptosis involves induction of the ER oxidase ERO1α, which causes activation of the ER calcium channel inositol 1,4,5-triphosphate (IP3) receptor (IP3R) and release of ER calcium (Li et al., 2009). The released calcium activates the enzyme calcium/calcmodulin-dependent protein kinase II (CaMKII), which triggers both death receptor and mitochondrial pathways of apoptosis (Timmins et al., 2009). Oxidative stress can also be a trigger for apoptosis in macrophages, although the detailed molecular mechanisms are not well understood (Forman and Torres, 2002). There is evidence that ER stress–induced macrophage apoptosis occurs in advanced atherosclerosis and contributes to plaque necrosis, a critical process in atherothrombotic vascular disease (Feng et al., 2003b; Thorp et al., 2009; Tabas, 2010). Oxidative stress has also been implicated in atherosclerosis, but its importance in advanced lesional macrophage death and plaque necrosis is not known (Glass and Witztum, 2001; Bonomini et al., 2008). With regard to the renal model, several different inducers of renal dysfunction result in ER stress–induced apoptosis of renal tubular epithelial cells, podocytes, and other cells in the kidney (Kitamura, 2008). Renal disease can also be associated with oxidative stress (Locatelli et al., 2003), but, as mentioned above, its possible links to ER stress are not known.

Using these two models, we show here that the aforementioned branch of ER stress signaling involving calcium and CaMKII induces NADPH oxidase (NOX) and oxidative stress, which are necessary for ER stress–induced apoptosis. We also provide evidence that CHOP is amplified by NOX/oxidative stress–mediated activation of a double-stranded RNA–dependent protein kinase (PKR)–CHOP pathway and that this amplification pathway is important for the final apoptotic response.

**Results**

**ER stress stimulates oxidative stress in macrophages, which is dependent on CHOP, proapoptotic calcium signaling, and NOX**

Using two atherosclerosis-relevant inducers of ER stress, cholesterol loading and 7-ketocholesterol (Tabas, 2002; Myoishi et al., 2007), we determined whether oxidative stress was part of the previously elucidated CHOP–calcium–CaMKII pathway of apoptosis (Li et al., 2009; Timmins et al., 2009). Dihydrorhodaminefluorescein (DCF) staining was used as a measure of intracellular peroxide accumulation (Wolber et al., 1987). We found that both inducers caused a substantial increase in the percentage of DCF-positive cells in wild-type (WT) macrophages, and the increase was much less in Chop−/− macrophages (Fig. 1 A). We next examined the calcium arm of the pathway using the intracellular calcium chelator BAPTA-AM. The increase in DCF-positive cells was decreased substantially by the chelator, which paralleled its inhibitory effect on annexin V staining as a measure of apoptosis (Fig. 1 B). DCF positivity and annexin V staining were also blocked by the CaMKII inhibitor KN93 but not by the structurally related inactive homologue KN92 (Fig. 1 C). We also used RNAi to silence CaMKII-γ, which is the isoform expressed in macrophages (Timmins et al., 2009). Two separate Camk2g siRNAs, which decrease Camk2g by ~60 and 75% (Timmins et al., 2009), gave similar results as KN93 (Fig. 1 D). Thus, oxidative stress, like apoptosis, is stimulated by ER stress in macrophages and is dependent on CHOP, intracellular calcium, and CaMKII.

One well-documented source of oxidative stress in macrophages is NOX (Iles and Forman, 2002). To test the relevance of NOX in ER-stressed macrophages, we first determined whether expression of the major NOX isoform in macrophages, NOX2, was affected by ER stress. We found that both cholesterol loading and 7-ketocholesterol induced Nox2 mRNA (Fig. 2 A, black bars). To test the causative role of NOX2 in oxidative stress and annexin V staining, we used siRNA against Nox2, which was effective in decreasing the mRNA to non–ER stress levels in the cholesterol- and 7-ketocholesterol–treated cells (Fig. 2 A, gray bars). This level of Nox2 suppression significantly decreased both DCF staining and annexin V staining in ER-stressed macrophages (Fig. 2 B). As an independent approach, we compared macrophages isolated from WT versus Nox2−/− mice and found similar effects on oxidative stress and apoptosis, using either annexin staining or TUNEL for the apoptosis assay (Fig. 2, C and D). These data indicate that NOX plays an important role in the oxidative stress and apoptotic responses of macrophages subjected to ER stress.

The role of both CaMKII and NOX in ER stress–induced oxidative stress and apoptosis raised the question as to whether the two enzymes were in the same or different pathways. Consistent with their being in the same pathway, inhibition of CaMKII activity with KN93, or genetic targeting of the kinase through the use of Camk2g−/− macrophages, blocked Nox2 mRNA induction by cholesterol loading or 7-ketocholesterol (Fig. 3 A). We showed previously that CaMKII activation requires an ERO1α–IP3R1–cadium release pathway (Li et al., 2009), which predicts that IP3R1 and ERO1α would play a role in Nox2 induction. For this purpose, we transfected macrophages with an Ero1α siRNA that decreases ERO1α expression by ~50% or an Ip3r1 siRNA that decreases IP3R1 expression by ~60% compared with scrambled RNA (Li et al., 2009) and found that each was able to suppress Nox2 mRNA induction (Fig. 3, B and C).

Our previous work showed that a key downstream effector of CaMKII-induced apoptosis is the mitogen-activated protein kinase JNK (Timmins et al., 2009). We found here that the JNK inhibitor SP600125 blocked Nox2 mRNA induction, DCF positivity, and annexin V staining in ER-stressed macrophages (Fig. 3, D and E). The Nox2 mRNA data were confirmed using macrophages from Jnk2−/− mice (Fig. 3 D, right graph). In summary, the data in Figs. 1–3 suggest that NOX plays a critical yet previously...
absence of NOX2 actually suppressed both CaMKII activation/phosphorylation and CHOP induction in cholesterol-loaded or 7-ketocholesterol–treated macrophages (Fig. 4 A). Note that the transcriptional inducer of CHOP in the phospho-eIF2α (P-eIFα) pathway, ATF4, was also suppressed in Nox2−/− macrophages, whereas neither phospho–PKR-like ER kinase (PERK) nor Xbp1 splicing was suppressed in Nox2−/− macrophages (Fig. 4, A and B). These data are consistent with the conclusion that NOX2 deficiency was specifically suppressing a non-PERK–activated P-eIF2α pathway rather than global ER stress. We therefore looked at another eIF2α kinase, PKR, and found that phosphorylation of this kinase, which is a measure of its activation, was substantially suppressed in Nox2−/− macrophages (Fig. 4 A).

unknown role in the ER stress–CHOP–calcium–CaMKII–JNK pathway of apoptosis.

Evidence for CHOP amplification through NOX/oxidative stress-mediated activation of PKR

The simplest way to integrate our previous findings (Li et al., 2009; Timmins et al., 2009) with the new data described thus far would be a linear pathway in which CHOP and CaMKII are upstream of NOX as follows: ER stress→CHOP→IP3R activation→ER calcium release→CaMKII activation→NOX activation→oxidative stress. However, the possibility of a bidirectional pathway was suggested by surprising data showing that absence of NOX2 actually suppressed both CaMKII activation/phosphorylation and CHOP induction in cholesterol-loaded or 7-ketocholesterol–treated macrophages (Fig. 4 A). Note that the transcriptional inducer of CHOP in the phospho-eIF2α (P-eIFα) pathway, ATF4, was also suppressed in Nox2−/− macrophages, whereas neither phospho–PKR-like ER kinase (PERK) nor Xbp1 splicing was suppressed in Nox2−/− macrophages (Fig. 4, A and B). These data are consistent with the conclusion that NOX2 deficiency was specifically suppressing a non-PERK–activated P-eIF2α pathway rather than global ER stress. We therefore looked at another eIF2α kinase, PKR, and found that phosphorylation of this kinase, which is a measure of its activation, was substantially suppressed in Nox2−/− macrophages (Fig. 4 A).
significantly decreased in the siRNA-treated macrophages (Fig. 4 E, right). Finally, we reasoned that a mechanism by which NOX2 activated PKR might be oxidative stress. In support of this idea, we found that the increases in both phospho-PKR and CHOP were also partially suppressed by the antioxidant N-acetylcysteine (NAC; Fig. 4 F). These data provide strong support for the concept that NOX2-induced oxidative stress amplifies CHOP expression through PKR activation and that this process is important for the induction of Nox2 itself as well as for oxidative stress and apoptosis.

**NOX is necessary for activation of IP3R and the mitochondrial pathway of apoptosis in ER-stressed macrophages**

The fact that CaMKII activation was suppressed in Nox2−/− cells (Fig. 4 A) suggested that NOX2 might be important in a key process upstream of CaMKII, IP3R activation (Li et al., 2009), as well an important proapoptotic process downstream of CaMKII, activation of the mitochondrial pathway of apoptosis (Timmins et al., 2009). To test the first prediction, we conducted experiments using Pkr siRNA. As shown in Fig. 4 C, the siRNA was effective in suppressing PKR, and this was associated with suppression of P-eIF2α and CHOP. Moreover, Pkr siRNA was able to block the ER stress–induced increment in Nox2 induction (Fig. 4 D).

In terms of apoptosis, we first asked whether the partial suppression of CHOP seen in Nox2−/− macrophages (Fig. 4 A) would be enough to decrease apoptosis. We therefore looked at apoptosis in Chop−/− macrophages, which express ~50% of the level of CHOP seen in WT macrophages (not depicted), and found that annexin V staining was indeed partially suppressed (Fig. 4 E, left). To test links among PKR, oxidative stress, and apoptosis, we tested the effect of Pkr siRNA on DCF fluorescence and annexin V staining. We found that both endpoints were significantly decreased in the siRNA-treated macrophages (Fig. 4 E, right). Finally, we reasoned that a mechanism by which NOX2 activated PKR might be oxidative stress. In support of this idea, we found that the increases in both phospho-PKR and CHOP were also partially suppressed by the antioxidant N-acetylcysteine (NAC; Fig. 4 F). These data provide strong support for the concept that NOX2-induced oxidative stress amplifies CHOP expression through PKR activation and that this process is important for the induction of Nox2 itself as well as for oxidative stress and apoptosis.

These data raised the possibility that NOX, through activation of PKR–P-eIF2α signaling, might amplify CHOP signaling and thus be necessary not only for CaMKII activation (Fig. 1, C and D) but also for induction of Nox2 itself (i.e., positive feedback loop), oxidative stress, and, most importantly, apoptosis. To explore this possibility, we conducted a series of experiments using Pkr siRNA. As shown in Fig. 4 C, the siRNA was effective in suppressing PKR, and this was associated with suppression of P-eIF2α and CHOP. Moreover, Pkr siRNA was able to block the ER stress–induced increment in Nox2 induction (Fig. 4 D).

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using the ATP–purinergic receptor pathway as a tool to acutely generate IP3 (Charest et al., 1985). As we previously reported (Li et al., 2009), IICR was increased by cholesterol loading in a standard intact-cell IP3R assay in which IP3-induced calcium release (IICR) is assayed at a given time point after exposure to a stimulus, in this case 6 h of cholesterol loading,
Figure 4. Evidence for CHOP amplification through NOX/oxidative stress–mediated activation of PKR. (A) Macrophages from WT or Nox2−/− mice were incubated under cholesterol-loading conditions (CHOL) or with 7-ketocholesterol (7KC) for the indicated times. Lysates were then immunoblotted for phospho-CaMKII (p-CaMKII), total CaMKII (T-CaMKII), CHOP, phospho-PERK (p-PERK), phospho-PKR (p-PKR), and GAPDH. (B) Macrophages from WT or Nox2−/− mice were incubated under cholesterol-loading conditions for 8 h (ATF4 experiment) or 9 h (XBP1 experiment). Nuclei were isolated and immunoblotted for ATF4 and nucleophosmin (Np) loading control, or RNA was extracted and assayed for spliced and unspliced Xbp1 and Gapdh loading control. (C) Macrophages were transfected with scrambled RNA (scrRNA) or Pkr siRNA. After 72 h, the cells were incubated for the indicated times under cholesterol-loading conditions. Lysates were then immunoblotted for PKR, phospho-eIF2α (P-eIF2α), CHOP, and β-actin loading control. (D) Macrophages were transfected with scrambled RNA or Pkr siRNA. After 72 h, the cells were incubated for an additional 8 h without sterol (Con) or with 7-ketocholesterol and then assayed for Nox2 mRNA. The data are displayed as Nox2 mRNA levels relative to those for the control scrambled RNA group. (E, left) Macrophages from WT, Chop+/−, or Chop−/− mice were incubated for 14 h without sterol, under cholesterol-loading conditions, or with 7-ketocholesterol and then assayed for annexin V staining. (E, right) Macrophages were transfected with scrambled RNA or Pkr siRNA. After 72 h, the cells were incubated for an additional 30 h without sterol, under cholesterol-loading conditions, or with 7-ketocholesterol and then assayed for DCF fluorescence and annexin V staining. In this graph, * indicates P < 0.05 and ** indicates P < 0.01. (F) Peritoneal macrophages were pretreated in the absence or presence of 5 µM of the antioxidant N-acetylcysteine (NAC). The cells were then incubated under cholesterol-loading conditions or with 7-ketocholesterol for the indicated time periods, also in the absence or presence of NAC. Lysates were then immunoblotted for phospho-PKR, CHOP, and β-actin. n = 3 for each experimental group. Data are presented as means ± SEM.
WT macrophages (Fig. 5 A, left, left pair of bars). However, there was no increase in IICR in cholesterol-loaded Nox2 \(^{-/-}\) macrophages (Fig. 5 A, left, right pair of bars). In theory, these increases and decreases in IICR could simply be driven by increases and decreases in ER luminal calcium, respectively, rather than by primary changes in IP3R activity. However, ER luminal calcium, as assessed by measuring thapsigargin-releasable calcium, was lower in the high-IICR cholesterol-loaded WT macrophages and higher in the low-IICR Nox2 \(^{-/-}\) macrophages (Fig. 5 A, right). Thus, the changes in IICR seen in Fig. 5 A (left)
were caused by changes in the fraction of ER calcium being released via differential IP3R activation. These data provide evidence that NOX2 is necessary for IP3R-mediated calcium release, which is consistent with the finding that NOX amplifies CHOP signaling (Fig. 4 A) in the context of the previously described CHOP–ER01/–IP3R pathway (Li et al., 2009).

A key proapoptotic process triggered by CaMKII activation is mitochondrial calcium accumulation and loss of mitochondrial membrane potential (Yao and Tabas, 2001; Timmins et al., 2009). To assess the role of NOX in these processes, we compared cholesterol-loaded WT and Nox2−/− macrophages for mitochondrial staining by Rhodamine-2 (Rhod-2) staining, a measure of mitochondrial calcium (Trollinger et al., 1997), and by MitoTracker red (chloromethyl-X-rosamine [CMXRos]), a measure of mitochondrial membrane potential (Poot et al., 1996). Cholesterol loading led to an increase in Rhod-2 staining in WT macrophages, and this was decreased substantially in Nox2−/− macrophages (Fig. 5 B). MitoTracker staining was diminished by cholesterol loading, which is consistent with its ability to cause mitochondrial membrane depolarization (Yao and Tabas, 2001), and this effect was also blocked in Nox2−/− macrophages (Fig. 5 C). The cholesterol-induced loss of MitoTracker red staining was also markedly prevented by NAC (Fig. 5 D). Mitochondrial membrane depolarization can lead to permeabilization of the outer mitochondrial membrane and release of proapoptotic factors, notably cytochrome c (Green and Reed, 1998). This process occurs in ER-stressed macrophages and is important for apoptosis (Yao and Tabas, 2001; Timmins et al., 2009). We show here that the increase in cytosolic cytochrome c and decrease in mitochondrial cytochrome c induced by cholesterol loading in WT macrophages were diminished in Nox2−/− macrophages (Fig. 5 E). Thus, activation of NOX plays an important role in the mitochondrial pathway of apoptosis in ER-stressed macrophages.

Functional role of NOX in ER stress-induced pathology in vivo

ER stress–induced macrophage apoptosis plays an important role in plaque necrosis in very advanced atherosclerosis, which in mouse models of atherosclerosis requires 4 mo or more of sustained hypercholesterolemia. However, Nox2−/− mice on either the Apoe−/− or Ldlr−/− backgrounds or in a bone marrow transplant model lose weight and become ill after 8 wk of hypercholesterolemia, likely because of infection (Pollock et al., 1995; unpublished data). Therefore, currently available genetic models cannot be used to assess the role of NOX in advanced atherosclerosis. Nonetheless, consistent with our in vitro data, we did find that atherosclerotic lesions from Chop−/−,Apoe−/− mice, which are partially protected against advanced atherosclerotic macrophage apoptosis (Thorp et al., 2009), had 36% less dihydroethidium (DHE) staining as a measure of peroxide accumulation (P < 0.05; Laurindo et al., 2008) and 37% less Nox2 mRNA, as assayed by laser capture microdissection/reverse transcription quantitative PCR (qPCR), compared with lesions from Chop−/−,Apoe−/− (P < 0.05 for both endpoints; unpublished data).

To directly test the role of NOX2 in vivo, we used a short-term model of systemic ER stress that utilizes injection of subtoxic amounts of the unfolded protein response (UPR) activator tunicamycin and monitors effects on renal tubular epithelium, a pathophysiologically relevant site for ER stress–induced apoptosis and tissue dysfunction (Zinszner et al., 1998; Kitamura, 2008). In particular, ER stress–induced renal cell apoptosis is a common pathway in several renal diseases (Kitamura, 2008). To test the link between CaMKII and NOX2 expression, we immunostained kidney slices from ER-stressed WT and Camk2g−/− mice for NOX2 protein and stained them with DHE for evidence of superoxide accumulation. As with cultured macrophages, both NOX2 and DHE staining were increased in the setting of ER stress, and the ER stress–induced increment in both parameters was much lower in the absence of CaMKII–γ (Fig. 6, A and B). To examine apoptosis, we used the TUNEL stain and found that the kidney was markedly protected from ER stress–induced apoptosis in Nox2−/− mice (Fig. 6 C). Most importantly, Nox2−/− mice were protected against renal dysfunction, as measured by serum creatinine and urinary albumin (Fig. 6 D). Consistent with these data and the positive feedback cycle revealed in the cell culture experiments, the induction of CHOP in the kidney of ER-stressed WT mice was also blunted in Nox2−/− mice (Fig. 6 E). Thus, the key role of NOX in ER stress–induced apoptosis occurs and is functionally important in vivo.

Discussion

ER stress–induced apoptosis is a critical process in several widespread chronic disease processes (Kim et al., 2008), and although oxidative stress has been implicated as a proapoptotic process in ER stress–induced cell death, the source of oxidative stress and its molecular connections to ER stress represent major gaps in our knowledge in this area (Malhotra and Kaufman, 2007a). The data in this study address these gaps by showing that NOX and NOX–induced oxidative stress are triggered by a specific arm of ER stress signaling involving calcium and CaMKII, and the resulting oxidative stress participates in an amplification pathway involving the PKR–CHOP pathway. As such, a positive feedback loop results in which CHOP induces Nox2 and NOX2 activates CHOP. However, a simple cycle involving all molecules in the pathway is not supported by our previous finding that silencing of ERO1α does not suppress CHOP (Li et al., 2009). Note that ERO1α and CHOP would be the two most separated points in a clockwise cycle because CHOP directly induces Ero1α to trigger the IP3R1–calcium–CaMKII–NOX2–apoptosis cascade (Li et al., 2009). Thus, there are many nodes between ERO1α and CHOP in the clockwise direction, and we reason that CHOP expression is likely maintained in ERO1α–silenced cells by one or more additional inputs in these areas.

The findings reported here raise new questions about signal transduction reactions downstream of the UPR and upstream of PKR. For example, with regard to Nox2 induction, the data in Fig. 3 D raise the possibility that JNK may be a transcriptional activator of the Nox2 gene. This hypothesis is consistent with a previous finding in smooth muscle cells that Nox4, another member of the NOX family, is induced by 7-ketocholesterol through an IRE1α–JNK pathway (Pedrizzetti et al., 2004). In view of this work, we considered a role for NOX4-mediated oxidative stress
Figure 6. NOX2 deficiency protects against ER stress–induced renal cell apoptosis, renal dysfunction, and CHOP induction. (A–C) WT, Camk2g−/−, and Nox2−/− mice were injected i.p. with 1 mg/kg tunicamycin (TUN) or vehicle (Veh) control. The kidneys were harvested 40 h later and stained using anti-NOX2 (A), DHE (superoxide accumulation; B), or TUNEL (apoptosis; C). All sections were also stained with DAPI to visualize nuclei. Bars, 20 µm. The bar graphs show quantification of the mean fluorescent intensities from three mice in each group. (D) Serum creatinine levels and urine albumin levels (normalized to urine creatinine) were determined for all groups of mice. (E) Kidney lysates were assayed for CHOP and β-actin by immunoblotting. For A–D, bars with the same symbols are not significantly different from each other, whereas bars with different symbols are significantly different from each other. n = 3 for each experimental group. Data are presented as means ± SEM.
in the pathway described here, but Nox2 mRNA is far more abundant than Nox4 mRNA in primary macrophages (unpublished data). In terms of PKR activation in our model, ER stress has been shown to activate a molecule called PKR-activating protein, leading to sustained PKR-mediated CHOP expression (Lee et al., 2007). In another setting, there is evidence of ER stress–induced PKR activation in the adipose and liver of obese mice (Nakamura et al., 2010), and although the mechanism has not been reported, it is intriguing to consider the separate finding that obese adipose tissue has increased expression of NOX and increased oxidative stress (Furukawa et al., 2004). In the model described here, oxidative stress plays a role in PKR activation (Fig. 4 F), and a previous study showed hydrogen peroxide treatment of Jurkat cells induces PKR through a pathway involving Janus kinase–signal transducer and activator of transcription signaling and interferon-γ induction (Pyo et al., 2008). Whether NOX is involved in any of these pathways remains to be explored.

We chose to use macrophages as a cell culture model to explore mechanisms linking ER stress and NOX because of pathophysiologic relevance in advanced atherosclerosis (Ibrahimii and Abumrad, 2002). There is also ample evidence of oxidative stress in advanced plaques (Bonomini et al., 2008), and the expression of NOX subunits, including NOX2, is up-regulated in atherosclerotic lesions (Sorensen et al., 2002). Moreover, we found that Nox2 mRNA and DHE staining, a measure of superoxide accumulation, were lower in Chop−/− lesions (unpublished data), which was consistent with our data. Although there are several informative studies on early-to-mid–stage atherogenesis using Nox2-null mice (Hsic et al., 2000; Barry-Lane et al., 2001; Vendrov et al., 2007; Judkins et al., 2010), the endpoint in such studies—lesion area—is entirely different from the atherosclerosis endpoints relevant to the signaling pathways described here—advanced lesion macrophage apoptosis and plaque necrosis. Unfortunately, advanced atherosclerosis is not amenable to in vivo studies using Nox2-null mice because during the months it takes to develop very advanced lesions, the null mice lose weight and become ill (unpublished data) and have decreased plasma cholesterol and triglyceride (Kirk et al., 2000), which may be caused by infection (Pollock et al., 1995).

The ultimate goal of elucidating signaling pathways linked to pathophysiologic processes is to suggest novel, mechanism-based approaches to disease prevention or treatment. Uninformed efforts at using antioxidant therapy in human diseases have been generally disappointing (e.g., the failure of vitamin E to prevent atherosclerotic heart disease; Williams and Fisher, 2005). However, understanding the interplay between ER stress and a specific cause of intracellular oxidative stress, as revealed here, could suggest new therapeutic strategies that are targeted, more specific, and mechanism based. In the case of the many disease processes involving both ER and oxidative stress, including advanced atherosclerosis and renal disease, our findings herein suggest that drugs targeting the CHOP branch of the UPR, and perhaps PKR in particular, and/or NOX may have particular promise. Conversely, attempts to kill cancerous cells can be thwarted by resistance to ER stress— and oxidative stress—induced apoptosis (Healy et al., 2009; Verrax et al., 2009), and knowledge of the pathways revealed here may suggest new mechanisms that could explain this resistance and new therapies for overcoming it.

Materials and methods

Reagents

Cell culture media and reagents were obtained from Invitrogen. HPCL-grade organic solvents were purchased from Thermo Fisher Scientific. Low-density lipoprotein (LDL; density, 1.020–1.063) was isolated from fresh human plasma by preparative ultracentrifugation as described previously (Havel et al., 1955). Our experiments with purchased human plasma were exempt from Institutional Review Board approval because the sources were anonymous. Acetyl-LDL was prepared by reaction of LDL with acetic anhydride (Basu et al., 1976). KN93 [2-(N[2-hydroxethyl]-N-[4-oxo-4-methylbenzenesulfonyl]amino-N-[4-chlorocinnaminyl]-N-methylbenzylamine], KN92 ([N-[4-oxo-4-methylbenzenesulfonyl]amino-N-[4-chlorocinnamyl]-N-methylbenzylamine phosphate), and SP600125 [anastro[1,9-c]pyrazol-4(1H)]-1 were obtained from EMD. All other chemical reagents, including the acyl-coenzyme A:cholesterol acyltransferase inhibitor 58035 [3-[decyl(dimethylsilyl)-N-[2-(4-methylphenyl)-1-phenylethyl] propanamide], were purchased from Sigma-Aldrich unless otherwise specified.

Mice

C57BL6/J WT mice and Nox2−/− mice on the C57BL6/J background were purchased from The Jackson Laboratory. Camk2g−/− mice on the C57BL6/J background were provided by J. Backs (University of Heidelberg, Heidelberg, Germany) and E. Olson (University of Texas Southwestern Medical Center, Dallas, TX) as described previously (Backs et al., 2010).

Cells

Peroxioneal macrophages from WT or mutant mice were harvested 3 d after i.p. injection of concanavalin A or 4 d after i.p. injection of methyl-BSA in mice that had previously been immunized with this antigen (Cook et al., 2003). Macrophages were harvested by peritoneal lavage with ice-cold PBS and maintained in DME containing 10% FBS and 20% L cell-conditioned medium. The medium was replaced every 24 h until the cells reached 80–90% confluency. Cholesterol loading was achieved by incubating the cells with 50 µg/ml acetyl-LDL plus 10 µg/ml 58035 for the indicated times (Devries-Schouw et al., 2005). The medium was changed every other day and immediately before treatments.

Calcin imaging

Macrophages cultured on 25-mm cover slips were placed in 35-mm dishes and loaded with 4 µM Flu-3 acetoxyethyl ester (Invitrogen) and 0.08% Pluronic F-127 in HBSS at room temperature for 30 min. The cells were then washed twice with HBSS, incubated in HBSS for an additional 30 min, and then mounted on the stage of an inverted confocal microscope (Live5; Carl Zeiss, Inc.) equipped with a 40x objective. 250 µM sulphinpyrazone was included in all solutions to prevent excretion of the Flu-3 by the macrophages. To trigger IICR, 10 µM ATP was added directly to the cell culture medium (Charest et al., 1985; Li et al., 2009). In some experiments, 2 µM thapsigargin, a sarco-ER calcium-dependent ATPase inhibitor, was added to assess ER calcium stores at the end of the experiments (Li et al., 2009). Cells were excited using the 488-nm laser line, and images were acquired at 1-s intervals under the time-lapse mode and subsequently analyzed using ImageJ (National Institutes of Health). The data are presented as a ratio of fmax/f0, where fmax represents the maximum fluorescence intensity obtained with ATP stimulation and f0 indicates the baseline fluorescence intensity. These data were quantified as area under the curve of the increment in fmax/f0.

Apoptosis assay

Apoptosis was assayed by annexin V staining using an apoptosis assay kit (Vybrant Apoptosis Assay kit #2; Invitrogen) and by TUNEL staining using the In Situ Cell Death Detection kit, tetramethylrhodamine (TMR) red (Roche; Yao and Tabas, 2000). For annexin V, the cells were gently washed once with PBS and then incubated for 15 min at room temperature with 110 µl annexin-binding buffer (25 mM Hepes, 140 mM NaCl, 1 mM EDTA, pH 7.4, and 0.1% BSA) containing 10 µl Alexa Fluor 488– or 594–conjugated annexin V. The staining mixture was removed and replaced with 110 µl of binding buffer. The cells were viewed immediately at room temperature with an inverted fluorescent microscope (IX-70; Olympus) equipped with filters appropriate for fluorescein (Alexa Fluor 488) or rhodamine (Alexa Fluor 594). Images were obtained with a charge-coupled device camera (Cool Snap; Photometrics) equipped with imaging software obtained from Roper Industries. Three fields of ~700 cells per field were photographed for each condition, and the number of annexin V-positive cells in each field was counted and expressed as a percentage of the total number of cells. TUNEL staining was assayed using
the In Situ Cell Death Detection kit, TMR red. At the end of the incubation period, the cells were gently washed once with PBS and then fixed in 4% paraformaldehyde in PBS for 1 h at room temperature. Then the cells were rinsed once with PBS and permeabilized in 0.1% Triton X-100 with 0.1% sodium citrate for 2 min on ice. The cells were then rinsed twice with PBS, 100 µl TUNEL reaction mixture (10 µl of labeling solution and 90 µl of enzyme solution) was added, and the cells were incubated for 1 h at 37°C with 5% CO₂. The staining mixture was then removed, and the cells were washed three times with PBS and then incubated in fresh PBS. The cells were viewed immediately at room temperature, as for the aforementioned annexin V assay, using filters appropriate for TMR (rhodamine). 10 fields of ~100 cells per field were photographed for each condition, and the number of TUNEL-positive cells in each field was counted and expressed as a percentage of the total number of cells.

Fluorescent analysis of peroxide accumulation

After incubation under control or cholesterol-loading conditions, macrophages were loaded with 5-(and-6)-chloromethyl-2',7'-DCF diacetate acetyl ester (Invitrogen). After 30 min, the cells were washed and viewed immediately at room temperature with an inverted fluorescent microscope (IX-70), equipped with filters appropriate for fluorescein, and images were obtained with a charge-coupled device camera (Cool Snap) equipped with imaging software. Three fields of ~700 cells per field were photographed for each condition, and the number of DCF-positive cells in each field was counted and expressed as a percentage of the total number of cells.

Immunoblotting

Cell extracts were electrophoresed on 4–20% gradient SDS-PAGE gels (Bio-Rad Laboratories) and transferred to 0.22-µm nitrocellulose membranes. The membranes were blocked for 1 h at room temperature in Tris-buffered saline and 0.1% Tween 20 (TBS-T) containing 5% (wt/vol) nonfat milk. The membranes were then incubated with primary antibody in TBS-T containing 5% nonfat milk or BSA at 4°C overnight, followed by incubation with the appropriate secondary antibody coupled to horseradish peroxidase. Proteins were detected by ECL (Thermo Fisher Scientific). Anti-CHOP and anti-ATF4 antibodies were obtained from Santa Cruz Biotechnology, Inc.; anti-α-actin, anti-phospho-Thr287-CaMKII, and anti-phospho-PKR antibodies were purchased from Millipore; and anti-phospho-ERK and anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) antibodies were obtained from Cell Signaling Technology.

siRNA-mediated gene silencing

siRNAs against murine Ip3r1 (S0206863277) and murine Pkp (S1003189108) were purchased from Qiagen. For CaMKIIγ-silencing (Fig. 1 D), two Camk2g siRNAs were used as described previously (Timmins et al., 2009); Camk2g siRNA #1 and Camk2g siRNA #2 in Fig. 1 D represent the siRNAs referred to as siRNA 2 and siRNA 3, respectively, in this previous study. For Ip3r1 and Ero1a silencing, Ip3r1 siRNA #1 and Ero1α siRNA #3 were used as described previously (Li et al., 2009). Macrophages grown to 50% confluency were transfected with each siRNA using Lipofectamine 2000 (Invitrogen) and 10 pmol siRNA per square centimeter well area (~40,000 macrophages per well in 48-well plates) according to the manufacturer’s instructions. After 4 h of transfection, the media were replaced, and, 72 h later, the indicated experiments were conducted.

In vivo model of ER stress using i.p. injection of tunicamycin

C57BL6/J and Nax2+/− mice were injected i.p. with concanavalin A, and then, 3 d later, the mice were given an i.p. injection of tunicamycin at a dose of 1 mg/kg of body weight. After 40 h, the kidney was harvested, and cryosections were prepared. TUNEL staining was performed using the In Situ Cell Death Detection kit, TMR red, followed by nuclear counterstaining with DAPI. Sections were mounted with anti-fade reagent (ProLong Gold). DHE is a hydrophobic uncharged compound that can cross intracellular membranes and binds to superoxide, which can be monitored as a bright red fluorescence (Laurindo et al., 2008).

Renal function tests

Serum was collected after centrifuging clotted blood at 3,000 g for 20 min. The creatinine concentration in serum was measured using a commercially available kit (Creatinine Enzymatic Assay kit; Bio-quant, Inc.). The albumin concentration in the urine was measured by a competitive ELISA (Exocell) and normalized to urine creatinine levels, which were measured as using the Creatinine Enzymatic Assay kit.

Statistics

Data are presented as means ± SEM. For most of the bar graph data, n = 3 for each experimental group. However, for the bar graphs depicting f_, the i/p data, n = ~30 per group. Analysis of variance followed by Tukey posttest (Prism 4 version 4.03, GraphPad Software, Inc.) was used to determine statistical significance among all groups. Bars with the same symbols are not significantly different from each other, whereas bars with different symbols are significantly different from each other. P-values for statistically different values ranged from P < 0.001 to P < 0.0001.

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Mitochondrial membrane potential

At the end of the indicated treatment periods, macrophages on glass-bottom 35-mm culture dishes (MatTek) were washed two times with PBS and then stained with 100 µM MitoTracker red CMXROS (Invitrogen) for 15–30 min at 37°C. Fluorescent images were captured and quantified as detailed in Mitochondrial calcium uptake for the Rhod-2 method.

Laser capture microdissection and RT-QPCR

RNA from nonendothelial intimal cells was captured by laser capture microdissection using a laser microdissection system (PALM; Carl Zeiss, Inc.). Immunohistochemistry performed on sections that were separated by 6 µm from the captured sections revealed that >70% of the intimal cells were F4/80-positive macrophages. Total RNA was isolated using the RNAqueous-Micro kit (Applied Biosystems) and reverse transcribed into cDNA using SuperScript III First-Strand Synthesis SuperMix (Invitrogen). QPCR was performed as described previously (Feng et al., 2003a).

Immunohistochemistry and immunofluorescence analyses

Mouse kidneys were immersion fixed in 10% neutral-buffered formalin overnight followed by embedding in paraffin. 5-µm paraffin-embedded sections were immunostained using anti-NOX2 antibody (Santa Cruz Biotechnology, Inc.) and the rabbit staining system (ImmuNo Cruz; Santa Cruz Biotechnology, Inc.). Sections were counterstained with DAPI and mounted with antifade reagent (ProLong Gold). Images were captured using an inverted microscope (Axiovert 200M; Carl Zeiss, Inc.). For immunofluorescence detection of superoxide, the frozen kidney or aorta sections were incubated with 2 µM DHE for 30 min at 37°C and then washed for 5 min with PBS. Sections were counterstained with DAPI and mounted with antifade reagent (ProLong Gold). DHE is a hydrophobic uncharged compound that can cross intracellular membranes and binds to superoxide, which can be monitored as a bright red fluorescence (Laurindo et al., 2008).

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