NADPH oxidase 2 inhibitors CPP11G and CPP11H attenuate endothelial cell inflammation & vessel dysfunction and restore mouse hind-limb flow

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ABSTRACT

First described as essential to the phagocytic activity of leukocytes, Nox2-derived ROS have emerged as mediators of a range of cellular and tissue responses across species from salubrious to deleterious consequences. Knowledge of their role in inflammation is limited, however. We postulated that TNFα-induced endothelial reactive oxygen species (ROS) generation and pro-inflammatory signaling would be ameliorated by targeting Nox2. Herein, we \textit{in silico}-modelled two first-in-class Nox2 inhibitors developed in our laboratory, explored their cellular mechanism of action and tested their efficacy in \textit{in vitro} and mouse \textit{in vivo} models of inflammation. Our data show that these inhibitors (CPP11G and CPP11H) disrupted canonical Nox2 organizing factor, p47\textsuperscript{phox}, translocation to Nox2 in the plasma membrane; and abolished ROS production, markedly attenuated stress-responsive MAPK signaling and downstream AP-1 and NFκB nuclear translocation in human cells. Consequently, cell adhesion molecule expression and monocyte adherence were significantly inhibited by both inhibitors. \textit{In vivo}, TNFα-induced ROS and inflammation were ameliorated by targeted Nox2 inhibition, which, in turn, improved hind-limb blood flow. These studies identify a proximal role for Nox2 in propagated inflammatory signaling and support therapeutic value of Nox2 inhibitors in inflammatory disease.

1. Introduction

NADPH oxidases (Noxs) comprise a family of seven isozymes whose only known function is the generation of reactive oxygen species (ROS) [1–7]. Besides their evermore-accepted role as signaling molecules, under pathophysiological conditions, excessive ROS production contributes to cell and tissue damage by oxidative stress and, as a consequence, to the development of myriad diseases. Noxs play a key role in stress and attendant disease, including varied cardiovascular maladies [7–11], cancer [12,13], and neurodegenerative disorders [14,15]. These encompass many of the major leading causes of death in the United States [16] and around the world. As the demand for selective Nox inhibitors for pharmacological and therapeutic purposes has become increasingly clear, drug development both from pharmaceutical industry and academia has intensified in recent years. Still, the need for well-characterized, isoform-specific inhibitors has not been met [7,17,18].

Our laboratory identified two bridged tetrahydroisoquinolines as small molecule inhibitors with high selectivity for the Nox2 isozyme [19]. In that study, the inhibitory potential of these molecules was preliminarily characterized using COS- and HEK- heterologous systems that expressed components of replete Nox isozyme systems and a non-Nox ROS generator. While these inhibitors demonstrated high Nox2 selectivity in a system of Nox subunit overexpression, their inhibitory profile \textit{in vivo} and in parenchymal primary cells \textit{in vitro} were neither interrogated nor their mechanism of action in cellular processes explored.

In this study, we beta-tested novel Nox2 inhibitors CPP11G and CPP11H [previously referred to as compounds 11g and 11h [19]] in \textit{in vitro} and \textit{in vivo} human and mouse inflammatory models, respectively, and examined the impact of Nox2 on a wide array of oxidant-sensitive signaling pathways leading to adhesion molecule expression, monocyte

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adhesion, vascular dysfunction and disrupted hind-limb blood flow. Herein, we model, in silico, the interaction of these compounds with 2 crucial binding domains within the canonical Nox2 (comprised of Nox2, p22phox, p47phox, p67phox, and Rac1/2) and report for the first time that they effectively block Nox2 activity by interfering with p47phox cytosol to membrane translocation and its interaction with cytochrome b558 heavy chain (CYBB). Furthermore, our findings reveal that TNFα-stimulated Nox2-derived ROS, MAPK activation, AP-1 and NFκB phosphorylation and nuclear translocation, adhesion molecule expression, endothelial dysfunction and impaired peripheral blood flow are all ameliorated by the Nox2 inhibitors. In aggregate, the findings hold meaningful promise for these agents and their derivatives as viable therapeutics in the treatment of myriad Nox2-actuated disorders.

2. Results

2.1. In silico analysis of Nox2 inhibitors modeling disruption of critical Nox2 oxidase subunit p47phox-p22phox interactions, and in vitro proof of blockade of p47phox binding to cytochrome b558 heavy chain (Nox2)

In silico interaction analysis of CPP11G and CPP11H with atomic structures of key components of the Nox2 oxidase was exploited (Fig. 1A). Modeling predicted that they interfere with a well-defined p47phox-SH3 “super groove” domain key interaction with a proline-rich (PRR) domain on the C-terminus of p22phox that is established as pivotal for the docking of p47phox to the membrane-spanning cytochrome b558 light chain p22phox [6,20–23]. This informed the hypothesis that these compounds would interfere with cytosolic p47phox interactions with membrane-integrated cytochrome components (either p22phox or Nox2) upon Nox activation. To interrogate this potential interference, COS-phox cells overexpressing the Nox2 isozyme subunits were classically stimulated with a phorbol ester. As shown in Fig. 1B, levels of p47phox at the plasma membrane were significantly elevated by phorbol 12-myristate 13-acetate (PMA, 5 μmol/l), a known protein kinase C activator.

Fig. 1. Nox2 inhibitors disrupt p47phox membrane translocation and interaction with the Nox2-p22phox complex. (A) In silico 3D-modeling based on PDB 1OV3 showing (i) both Nox2 inhibitors CPP11G & H interfering with p22phox C-terminal PRR domain interaction with the p47phox SRC homology 3 (SH3) domain (super groove) (overlay); (ii) the p22phox C-terminus alone interacting with the p47phox SH3 super groove, (iii, iv) Selective Nox2 inhibitors binding individually to the p47phox super groove. (B) Western blotting of membrane fraction showing the effects of CPP11G (20 μmol/l) and CPP11H (20 μmol/l) on PMA (5 μmol/l)-induced p47phox membrane translocation in COS-phox cells overexpressing Nox2 subunits. The band density of p47phox protein in the membrane fraction was normalized to the density of Nox2 band detected in the same sample, n = 5 (*p < 0.05 vs. control, Ctrl, ##p < 0.01 vs. PMA.) (C) Western blotting showing the effects of CPP11G (20 μmol/l) and CPP11H (20 μmol/l) on PMA (5 μmol/l)-stimulated interaction between p47phox and Nox2 in COS-phox cells overexpressing Nox2 subunits. p47phox was immunoprecipitated (IP) from COS-phox homogenates and detected by Western blot for the presence of Nox2. The densities of Nox2 protein bands were normalized to the levels of p47phox detected in the same samples, n = 6 (*p < 0.05 vs. Ctrl, ###p < 0.001 vs. PMA).
which causes phosphorylation of p47\textsuperscript{phox} and triggers its membrane translocation. This response was inhibited in the presence of CPP11G and CPP11H (Fig. 1B). To further investigate their effects on the interaction between p47\textsuperscript{phox} and Nox2, co-immunoprecipitation was applied on a preparation of COS-phox cells, where PMA (5 μmol/l) stimulation increased Nox2 to p47\textsuperscript{phox} binding (Fig. 1C). Consistent with the results on p47\textsuperscript{phox} membrane translocation, both agents diminished PMA-induced interaction between Nox2 and p47\textsuperscript{phox}. In summary, CPP11G and CPP11H suppressed p47\textsuperscript{phox} cytosol-to-membrane translocation, thus blocking the Nox2 isozyme complex formation and by extension ROS-producing activity.

### 2.2. CPP11G and CPP11H inhibit Nox2-derived ROS production, MAPK/SAPK signaling and AP-1 activation in response to TNFα

To evaluate the effectiveness of these Nox2 inhibitors under more physiological conditions, primary human aortic endothelial cells (HAECs) were employed to establish an acute inflammatory model of TNFa stimulation (10 ng/ml) in which the agents were optimally beta-tested. As expected, HAECs challenged with TNFa yielded increased amounts of O\textsubscript{2}\textsuperscript{•–}, which was measured using the fluorescent hydropropidine (HPr\textsuperscript+), a cell-impermeant probe for detecting extracellular O\textsubscript{2}\textsuperscript{•–}. Importantly, both CPP11G and CPP11H (10 μmol/l) obliterated this response (Fig. 2A). Next, as O\textsubscript{2}\textsuperscript{•–} is rapidly dismutated to H\textsubscript{2}O\textsubscript{2}, levels of H\textsubscript{2}O\textsubscript{2} stimulated by TNFa and the effects of the Nox2 inhibitors were assessed using a recently optimized coumarin-7-boronic acid (CBA) as well as the Amplex Red assay on whole cells and cell homogenates, respectively (Figs. 2B and C). Consistent with the O\textsubscript{2}\textsuperscript{•–} results, TNFa-stimulated elevation in H\textsubscript{2}O\textsubscript{2} was effectively blocked by both compounds (each at 10 μmol/l). Moreover, excessive generation of O\textsubscript{2}\textsuperscript{•–} gives rise to peroxynitrite (ONOO\textsuperscript–), a highly reactive and often destructive ROS that oxidizes lipoproteins and nitrosates tyrosine residues in many proteins, forming 3-nitrotyrosine (3-NT) [24], a well-established “footprint” biomarker of oxidative damage. Indeed, HAECs treated with TNFa (10 ng/ml) for 24 h exhibited increased (~2 fold vs. control, Ctrl) 3-NT formation as evidenced by stronger immunofluorescent staining. In comparison, HAECs that were pretreated with CPP11G (10 μmol/l) or CPP11H (10 μmol/l) revealed significantly lower levels of 3-NT than cells with TNFa alone (Fig. 2D), indicating by a fourth independent method that these inhibitors are capable of limiting Nox2-derived ROS in HAECs.

Next, we examined their effects on TNFa-induced Nox2-mediated mitogen-activated protein kinase (MAPK) signaling. TNFa (10 ng/ml) triggered a rapid and transient phosphorylation of p38 MAPK, which peaked at 10 min (6.9 ± 0.6 fold vs. Ctrl) and dropped below baseline by 30 min (Supplementary Fig. 1A). Similarly, SAPK/JNK phosphorylation in response to TNFa (10 ng/ml) was also transiently time-dependent, which achieved a maximum but transitory stimulation at 20 min with TNFa stimulation (10.3 ± 0.6 fold vs. Ctrl) (Supplementary Fig. 1B). Unexpectedly, no stimulations in ERK1/2 (p44/42) MAPK phosphorylation were observed with the first hour or even after prolonged TNFa (10 ng/ml) treatment (up to 24 h) (Supplementary Figs. 1C and D), suggesting that TNFa and subsequent Nox-derived ROS differentially regulate downstream MAPK signaling.

![Fig. 2. Nox2 inhibitors attenuate TNFa-induced reactive oxygen species (ROS) production in human aortic endothelial cells (HAECs).](image-url)
As TNFα-induced intracellular events appeared rapid and transient, Nox2 inhibitor effects were evaluated at the time of peak activation. Phosphorylation of p38 by TNFα (10 ng/ml, 10 min) was markedly suppressed by CPP11G and CPP11H by ≥50% in a concentration-dependent manner (Fig. 3A). Consistently, activation of SAPK/JNK was also robustly and concentration-dependently inhibited by CPP11G and CPP11H (Fig. 3B).

Activated MAPK, particularly SAPK/JNK, phosphorylates the transactivation domain of c-Jun, a key component of the transcription factor AP-1, thereby facilitating AP-1 to initiate inflammatory gene expression [25,26]. Thus, we next investigated the effects of our small molecule Nox2 inhibitors on c-Jun phosphorylation and nuclear translocation. With 1 h of TNFα (10 ng/ml) challenge, the ratio of active (phosphorylated) c-Jun to total c-Jun in whole cell lysates was enhanced (2.7 ± 0.2 fold vs. Ctrl). Both CPP11G and CPP11H significantly inhibited c-Jun phosphorylation concentration-dependently (Fig. 3C). Moreover, we were able to show that c-Jun translocated to the nucleus, where it positively regulates genes involved in endothelial activation [25,26]. Consistently, we found that TNFα (10 ng/ml, 1 h) substantially elevated the levels of phospho-c-Jun in the nuclear fraction of endothelial cells (~5-fold) wherein CPP11H exhibited a notably higher efficacy in diminishing c-Jun nuclear translocation by 60.7 ± 8.0% (p-c-Jun/Histone); CPP11G displayed a tendency toward an inhibition (not significant; Fig. 3D). Collectively, these results

Fig. 3. Nox2 inhibitors attenuate TNFα-induced mitogen-activated protein kinase (MAPK) activation in HAECs. (A) Representative Western blots (left) and cumulative data (right) showing concentration dependent inhibition by CPP11G and CPP11H of TNFα (10 ng/ml, 10 min)-stimulated p38 MAPK activation. The densities of phospho-p38 were normalized to the levels of total p38 detected in the same samples, n = 6–8. (****p < 0.0001 vs. unstimulated, #p < 0.05 vs. TNFα, ##p < 0.01 vs. TNFα, ####p < 0.0001 vs. TNFα). (B) Representative Western blots (left) and cumulative data (right) showing concentration dependent inhibition by CPP11G and CPP11H of TNFα (10 ng/ml, 20 min)-stimulated JNK phosphorylation. The densities of phospho-JNK were normalized to the levels of total JNK detected in the same samples, n = 7. (****p < 0.0001 vs. unstimulated, #p < 0.05 vs. TNFα, ##p < 0.01 vs. TNFα). (C) Representative Western blots (left) and cumulative data (right) showing concentration dependent attenuation by CPP11G and CPP11H of TNFα (10 ng/ml, 1 h)-stimulated c-Jun phosphorylation. The densities of phospho-c-Jun were normalized to the levels of total cJun detected in the same samples, n = 6. (****p < 0.0001 vs. unstimulated, #p < 0.05 vs. TNFα, ##p < 0.01 vs. TNFα, ####p < 0.0001 vs. TNFα). (D) Representative Western blots (left) and cumulative data (right) showing the effects of CPP11G (10 μmol/l) and CPP11H (10 μmol/l) on TNFα (10 ng/mL, 1 h)-modified phospho-cJun levels on the nuclear fraction. The densities of phospho-cJun were normalized to the levels of histone detected in the same samples, n = 3. (***p < 0.001 vs. control, ###p < 0.001 vs. TNFα).
indicate that both CPP11G and CPP11H are broadly effective in suppressing TNFα-induced Nox2-mediated MAPK/SAPK and AP-1 signaling.

2.3. Nox2 inhibitors suppress IKK-NFκB pathway induced by TNFα

In addition to the MAPK and AP-1 signaling cascades, TNFa is implicated in Nox2-dependent NFκB signaling to endothelial inflammation [27,28]. Hence, we characterized the effects of CPP11G and CPP11H on the NFκB signaling cascade. Mirroring the mode of MAPK activation, IκB kinases α & β (IKKα/β) transiently reached a ∼20-fold increase in phosphorylation (represented as p-IKK) within 10 min of TNFa stimulation (Supplementary Fig. 2A). Diminution in endothelial cell IκB (NFκB attenuator) levels was also observed with TNFa (10 ng/ml) treatment in whole cell lysates. This effect, in keeping with IκB ubiquitination and degradation, is widely reported [29–31]. Within 20 min, IκB levels reached its nadir, and appeared to partially recover at later time points (Supplementary Fig. 2B). These results illustrate that TNFa triggers the canonical NFκB pathway in HAEcs in a time-dependent fashion. More importantly, administration of the novel small molecule Nox2 inhibitors revealed that TNFa (10 ng/ml, 10 min)-activated IKKα/β was concentration-dependently attenuated by both CPP11G and CPP11H (Fig. 4A). Characteristically, TNFa-ablated IκB levels were partially but significantly rescued by both inhibitors at a concentration as low as 5 μmol/l (Fig. 4B). Consistent with these observations, NFκB p65 activation/phosphorylation in whole cell lysates was highly elevated by TNFa (10 ng/ml, 1 h) (4.3 ± 0.2 fold vs. Ctrl). Both agents suppressed this response (Fig. 4C). Upon interogation of NFκB p65 nuclear translocation using fluorescence microscopy (Fig. 4D), TNFa (10 ng/ml, 1 h) discernibly increased the levels of p65 in HAEc nuclei, which were abolished by CPP11H (10 μmol/l), and were trending towards an inhibition by CPP11G (10 μmol/l). To further validate these results, we compared the phospho-p65 levels in the nuclear extract isolated from endothelial cells. Phospho-p65 in the nuclear fraction (p-p65 vs. histone) was substantially potentiated by TNFa (10 ng/ml) (29.1 ± 2.2 fold vs. control). In accordance with our observations on immunofluorescent images, pretreating cells with CPP11H but not CPP11G significantly attenuated phospho-p65 in the nuclear extract (Supplemental Fig. 2E). These findings demonstrate that while both agents elicit inhibitory actions on the proximal NFκB pathway, CPP11H uniquely displays efficacy with respect to NFκB nuclear translocation.

2.4. CPP11G and CPP11H ameliorate endothelial inflammation and monocyte adhesion in response to TNFa

Upon activation of NFκB and AP-1 signaling cascade, inflammatory endothelial activation response ensues with upregulated expression of adhesion molecules [26], including vascular cell adhesion molecule-1 (VCAM-1), intracellular adhesion molecule-1 (ICAM-1) and E-selectin, all of which facilitate the recruitment of neutrophils and monocytes, and predispose for vascular inflammation and atherosclerosis [32]. We first investigated the ability of the tetrahydrosoquinolines [19] to suppress TNFa-adhesion molecule upregulation at the protein level. Treating HAEcs with TNFa for 24 h (a time point selected based on Supplementary Figs. 2C and D illustrating that VCAM-1 and ICAM-1 are stimulated in a time-dependent manner with maximal and robust inductions appearing at 24 h respectively) promoted the levels of VCAM-1 and ICAM-1 by ∼50-fold and ∼20-fold, respectively, which were sharply repressed by CPP11G and CPP11H in a concentration-dependent fashion. Remarkably, VCAM-1 and ICAM-1 expression were virtually abolished at the highest concentrations (10 μmol/l) (Figs. 5A and B). To confirm these observations, a monocyte recruitment assay was performed comparing calcein-labelled monocyte adherence to HAEcs in response to different treatments. As is shown in Fig. 5C, TNFa (10 ng/ml, 24 h) stimulation markedly promoted monocyte adhesion to HAEcs as indicated by a 2.2 ± 0.2-fold increase in fluorescence intensity of monocytes compared to control. Pretreating HAEcs with either inhibitor concentration-dependently diminished the TNFa-induced response, with 10 μmol/l CPP11G or CPP11H by 30.9 ± 1.3% or 29.1 ± 2.4%, respectively. These findings point to the novel Nox2 inhibitors as efficacious agents in ameliorating endothelial activation and chemotraction of monocytes.

2.5. Nox2 inhibition decreases ROS production and ameliorates endothelial dysfunction in response to TNFa in vivo

To test the clinical relevance of these agents in preventing ROS production and preserving vascular functionality, we employed an acute inflammatory mouse model as per cytokine challenge. Mice were initially injected with either vehicle (30% DMSO in PBS), CPP11G (15 mg/kg) or CPP11H (15 mg/kg) by i.v. bolus. Subsequently, TNFa (20 μg/kg) was administered by i.v. bolus in mice in the TNFa plus/minus CPP11G and CPP11H groups. To evaluate the effectiveness of CPP11G and CPP11H in ROS (hydrogen peroxide, H2O2) production in vivo, we employed Amplex Red to examine levels of H2O2 production in aortic homogenates. We observed that aortas taken from mice treated with TNFa alone generated 69.1 ± 12.5% higher H2O2 in comparison to the control group. Importantly, H2O2 levels were significantly lower in mice pretreated with CPP11H and trending toward a reduction by CPP11G (Fig. 6A). These observations suggest that the inhibitors are efficacious in limiting Nox2-derived ROS generation in vivo. Clinical significance of these novel Nox2 inhibitors was also exemplified by their beneficial effects on endothelial inflammation. In particular, ICAM-1 protein levels on the aortic intima were augmented considerably by TNFa stimulation (Fig. 6B). DAPI (blue) and α-actin (green) staining show the bulk of tissue staining in the media while merged images depict positive ICAM-1 (red) staining along the luminal lining of the vessel. ICAM-1 expression was visibly reduced in mice treated with both inhibitors and this decrease reached significance in the CPP11H group (Fig. 6B; lower magnification images shown in Supplementary Fig. 3).

To explore effects on vascular function, we evaluated mouse hindlimb blood flow (Fig. 6C). Compared to mice in the control group, TNFa-stimulated mice exhibited a 25.7 ± 7.3% decrease in basal femoral artery blood flow (highest degree of flux shown in red). However, when mice were pre-injected with either agent, femoral artery blood flow was restored to or even slightly above basal levels, indicating that the Nox2 inhibitors are able to ameliorate TNFa-mediated blood flow disturbance in vivo.

Further, effects on endothelial dilatory function were interrogated using ex vivo two-pin myography. Aortic rings were isolated and mounted on two pins, and stimulated in myograph organ chambers with TNFa (100 ng/ml) for 2 h in the presence or absence of inhibitors. Acetylcholine (ACh: 10−8-10−5 mol/l) as a direct smooth muscle vasodilator displayed no change in aortic rings incubated with TNFa (Fig. 6D). In contrast, pretreating the aortic rings with either CPP11G (20 μmol/l) or CPP11H (20 μmol/l) prevented TNFa-induced endothelial dysfunction (Fig. 6D), pointing to the capability of both compounds to preserve endothelial cell function. Control experiments using sodium nitroprusside (SNP: 10−9.5, 10−8 mol/l) as a direct smooth muscle vasodilator displayed no change among the treatment groups (Supplementary Fig. 4). These results demonstrate that neither TNFa nor Nox2 inhibitors alter smooth muscle cell vasomotor tone.

3. Discussion

Herein, we interrogated the capacity of novel Nox2-selective small molecule inhibitors CPP11G and CPP11H to alter endothelial ROS production, vascular inflammatory responses and dilator function. The major findings of the current study are the ability of these inhibitors to...
1) disrupt canonical p47phox-p22phox interaction, p47phox membrane translocation and interaction with the membrane-bound Nox2 subunit in the heterologous Nox2 isozyme; 2) abrogate TNFα-stimulated Nox2-derived ROS, MAPK, AP-1 and NFκB pathway activation in human endothelial cells; 3) limit TNFα-stimulated endothelial adhesion molecules and monocyte adherence; 4) prevent in vivo TNFα-elicted ROS production and vascular endothelial inflammatory response; 5) ameliorate ex vivo endothelial dysfunction; and 6) preserve hind-limb blood flow in mice. Collectively, these findings indicate that the novel inhibitors are highly efficient at inhibiting Nox2-mediated signaling cascades and protecting vascular function against inflammatory insults both in human endothelial cells and in an in vivo mouse model of systemic inflammation.

The small molecule Nox2-selective inhibitors described herein were originally identified by high-throughput screening assay of a subset of approximately 600 small molecules from the University of Pittsburgh Chemical Methodologies and Library Development (UPCMLD) library, followed by structure-activity relationship (SAR) studies of newly
synthesized molecules containing a common backbone structure [19]. Among these molecules, CPP11G and CPP11H emerged as highly efficacious and selective inhibitors for the Nox2 isozyme [19] exhibiting no inhibition of Nox1, 4, or 5 isozymes or xanthine oxidase, thus demonstrating no direct ROS scavenging properties. By comparison, the contemporaneous existing tools for Nox2 inhibition, including diphenylene iodonium (DPI), apocynin, VAS2870 and ebselen congeners, presented limitations ranging from non-selective Nox inhibition to formidable off-target effects, prognostic of undesired consequences in vivo and a poor Nox2-selective therapeutic [17,18]. Therefore, with high specificity and no observed off-target effects, CPP11G and CPP11H held significant promise as therapeutic agents for Nox2-mediated pathologies. Their actions, however, in parenchymal cells in vitro and in vivo had not been studied.

Aiming to gain insights into the mechanisms underlying their Nox2 suppression, in silico analyses of various Nox2 subunit interactions were performed. Computational modeling predicts that CPP11G and CPP11H interfere with a critical interaction of a proline-rich binding domain within the C-terminus of p22phox which binds to a well-characterized Src-homology 3 (SH3) “super groove” in the p47phox subunit involved in requisite assembly and canonical Nox2 isozyme activation [20–23,33] (Fig. 1A). Consistent with this prediction, we found that both agents: 1) abrogated translocation of p47phox from the cytosol to plasma membrane where Nox2 (aka gp91phox) and p22phox are juxtaposed and reside (Fig. 1B); and 2) disrupted the canonical active enzyme-essential p47phox:Nox2 interaction (Fig. 1C). These findings corroborate that by way of impeded p47phox interaction with the membrane-bound cytochrome (CYBB & CYBA), these agents disrupt the assembly of the Nox2 isozyme complex, thus achieving inhibition. A similar mechanism of action has been reported for the previously described peptidic Nox2 inhibitor, Nox2ds-tat, which blocks the interaction between Nox2 and p47phox by binding to the latter and preventing p47phox membrane translocation [34,35]. The most salient perceived drawback of that agent is a limited oral bioavailability as a peptide in its current form. Another small molecule inhibitor, celastrol, appears to disrupt binding of a p22phox proline-rich peptide to the tandem SH3 domain of both p47phox and NoxO1 (homolog for p47phox in the Nox1 system) [36]. However, celastrol is expected to have multiple off-target effects, as it also inhibits cytosolic factor-independent Nox isoforms such as Noxs4 and 5 [36]. Furthermore, another group of small molecules, namely ebselen and congeners, appear to also interrupt the binding of the SH3 domain of p47phox to the proline-rich domain of p22phox, and prevent p47phox and p67phox translocation to the plasma membrane in neutrophils [37]. However, ebselen and its analogs exhibited poor selectivity for Nox2 over Nox1 with the exception of one congener, JM-77b, which showed relatively higher selectivity, namely a lower IC50 for Nox2 compared to Nox1 and Nox5 with no detectable inhibition of Nox4 [27]. In addition, ebselen and its analogs have been reported to affect numerous targets, including protein kinase C [38], endothelial nitric oxide synthase [39,40], lipoxigenases [41], c-Jun N-terminal kinase [40] and horseradish peroxidase [42]. Therefore, whether or not they directly prevented p47phox and p67phox membrane translocation or indirectly mitigated Nox activation remains undefined.

An important question that has not been addressed in the current...
study is whether CPP11G and CPP11H could disrupt Nox1 activity in a hybrid system that employs p47\textsuperscript{phox} as its organizing subunit [43]. That is, the crucial isozyme-activating p47\textsuperscript{phox}, p22\textsuperscript{phox} interaction not only sustains Nox2 isozyme activation, but is also operant in a hybrid Nox1 system wherein, in lieu of NoxO1, p47\textsuperscript{phox} is utilized as a cytosolic organizer. The hybrid Nox1 system has been reported to be functional in vascular smooth muscle cells [43,44] and in a variety of disease settings [17,45,46].

With that said, one limitation of this work is that to-date CPP11G/H mechanism of action studies have only been conducted in a cell system that expresses the canonical Nox2 subunits (Nox2, p22\textsuperscript{phox}, p47\textsuperscript{phox}, p67\textsuperscript{phox}, and Rac 1/2). Thus, whether CPP11G and CPP11H inhibit the hybrid Nox1-p47\textsuperscript{phox} interaction warrants further investigation.

A process wherein Nox2-dependent ROS production has been implicated is inflammation [47–49]. A defense response to injury that involves a cascade of cellular signaling responses, inflammation and its mediators initiate endothelial activation and expression of membrane adhesion molecules, all of which are needed for monocyte recruitment and infiltration [50]. However, when ROS production is sustained, it is noted to give rise to chronic inflammation and endothelial dysfunction, and underlie the pathophysiology of many cardiovascular and non-cardiovascular disorders [49]. One of the most widely studied pro-inflammatory cytokines, tumor necrosis factor alpha (TNFa), has been shown to induce endothelial activation by increasing Nox-derived ROS production and by engaging the NFkB pathway to increase expression of inflammatory genes such as those for cytokines and adhesion molecules [47,51,52].

With respect to the current study, several lines of evidence...
suggested that the Nox2 isozyme is a source of ROS and a mediator of an inflammatory response in endothelial cells. Specifically, TNFα failed to induce oxidant generation in lung vascular endothelial cells from Nox2 null mice [53], a response normally characterized by NfκB activation and inflammation in wild type mice. Furthermore, TNFα-induced $O_2^{-}\cdot$ generation in pulmonary microvascular endothelial cells was shown to be inhibited by selective Nox2 inhibitor, Nox2ds-tat, but not by Nox1-selective inhibitor NoxA1ds, concomitant with a suppressed adhesion molecule expression and monocyte adherence [52]. Still, other lines of evidence show that TNFα also induces ROS production from alternate sources including Nox1 [54] and Nox4 [55]. With that said, further testing of the hypothesis with an additional Nox2-selective agent that is delivered acutely and avoids compensatory roles of other sources spurred the current studies. Moreover, the nature of the current agents as small molecules, their selectivity, their potential for facile oral bioavailability, and alterability for enhanced pharmacokinetic and SAR properties, justified their beta-testing in the current study. On these premises, the characterization of CPP11G and CPP11H on inflammatory pathways was conducted in both human endothelial cells in culture and mouse endothelial cell and vascular tissue both in vivo and ex vivo.

We report that CPP11G and CPP11H are highly effective in reducing TNFα-stimulated ROS production in human aortic endothelial cells (HAECs, Fig. 2). We employed HPr$^+$ (a cell-impermeable $O_2^{-}\cdot$-specific probe [56]) to quantify $O_2^{-}\cdot$ generation as the initial product of Nox2 isozyme activity. Our data show that both compounds abrogated the HPr$^+$ signal in these cells, in accordance with a highly effective enzymatic blockade. Coumarin-7-boronic acid (CBA [57]) was used for comprehensive detection of $H_2O_2$ (the rapidly dismutated metabolite of $O_2^{-}\cdot$) on the whole cell level and similar results were observed. Employing a third assay of direct ROS detection (Amplex Red), we were able to show that in cell homogenates, the rise in $H_2O_2$ was abolished by the novel Nox2 inhibitors. These results corroborate by three independent biochemical assays that ROS generation was maximally inhibited. Furthermore, these findings were confirmed by significant blockade of “footprint” oxidation and nitration marker 3-nitrotyrosine (3-NT) in whole cells in culture (Fig. 2). In aggregate, the data are strongly supportive of the ability of CPP11G and CPP11H to ablate Nox2 activity in these cells.

Our data also show that TNFα triggers activation of inflammation-related MAPK/AP-1 signaling and the IKK-NfκB pathway in HAECs. For one, we observed that both compounds inhibited p38 and JNK activation albeit to varying degrees. Subsequent phosphorylation/activation of another key player involved in these pro-inflammatory pathways, c-Jun, that in conjunction with c-Fos, comprises the AP-1 transcription factor complex, was effectively blocked by both CPP11G and CPP11H (Fig. 3). In addition to the MAPK-AP1 signaling cascade, TNFα-trig- gered IKK-NfκB pathway, characterized by IkκB activation, IkB degradation and NfκB (p65) phosphorylation, was significantly mitigated by CPP11G and CPP11H (Fig. 4), recapitulating a primary role of Nox2-ROS in mediating TNF-induced endothelial inflammation reported by others [28,52]. Intriguingly, our results on the c-Jun and p65 nuclear translocation consistently show that only CPP11H, but not CPP11G, (Figs. 3 and 4 and Supplementary Fig. 2) was able to significantly reduce their levels in the nucleus. The fact that CPP11G and CPP11H are equally effective preventing ROS generation and enzyme assembly but differ in some downstream endpoints could be explained by the distinct time points at which the measurements are taken, thus reflecting a difference in binding affinity/stability between both compounds. In fact, in Fig. 2D, 3-NT footprint assessed at 24 h supports the idea that CPP11H effects endure more strongly at later time points than those of CPP11G. The reason for this disparity, however, remains unclear but could also suggest a differential penetrability of the two compounds to compartments that connect Nox2 with c-Jun or p65.

Further studies in terms of pharmacokinetic and pharmacodynamic properties of both compounds are required to assess the possibility of unexpected off-target effects or effects on Nox-independent functions of p47$^\text{HOX}$. It is also worth noting that even relatively high concentrations of CPP11G and CPP11H (10 μmol/l) were not able to completely abolish these TNFα-activated signaling cascades. This may be attributed to a need to test even higher concentrations of the drug that approach or exceed their previously reported IC$\text{50}$ values [19]. Of course, the possibility also exists that TNFα alternatively triggers these pathways independently of Nox2-derived ROS. Several lines of evidence show that TNFα also induces ROS production from other sources in different primary cell types, such as mitochondria [58,59], Nox1 [54] and Nox4 [55]. However, whether they are subjected to TNFα regulation in human aortic endothelial cells, and whether they play a role in endothelial activation and inflammation remains undefined. In addition, though MAPKs and NfκB are generally sensitive to the redox state of their environment, it has also been reported that MAPKs and NfκB can be activated in a ROS-insensitive manner [60,61] as ROS scavengers fail to completely abrogate receptor-induced MAPK activation, and that activation of NfκB signaling can be achieved by direct binding of TNFα receptor with components of the NfκB cascade through TNF receptor-associated factor 2 (TRAF2) [62]. It is therefore plausible that TNFα-induced endothelial activation is mediated primarily but not exclusively by Nox2/ROS.

Apart from our studies with respect to intracellular signaling pathways, translational relevance was demonstrated by our findings that CPP11G and CPP11H in vitro effectively suppressed upregulation of adhesion molecules in response to TNFα (Figs. 5A and B) on human cells, leading to significantly ameliorated monocyte attachment in the presence of CPP11G or CPP11H (Fig. 5C). Furthermore, in vivo validation in this study of the novel compounds is consistent with our in vitro results. That is, in vivo administration of TNFα prominently enhanced aortic $H_2O_2$ production and intimal ICAM-1 expression. Both these responses were more effectively suppressed by CPP11H than CPP11G (Figs. 6A and B). Indeed, it has been well established that the MAPK, AP-1 and NfκB pathways control adhesion molecule expression and monocyte adhesion [26,47,51]. In aggregate, ROS production, adhesion molecule expression and monocyte adhesion in vivo and ROS and adhesion molecule expression in mouse aortas in vivo were ameliorated significantly by CPP11H indicative of its effectiveness in alleviating Nox2-mediated vascular inflammation.

Endothelial activation has been shown to potentiate endothelial dysfunction not only by inhibiting eNOS expression [63], but also by decreasing NO bioavailability through induction of $O_2^{-}\cdot$ and generation of ONOO$^-$ [64,65]. In line with this notion as well as a number of studies by others [66–69], we also observed that incubation of mouse aortas with TNFα led to a significantly blunted vasodilatory response to the endothelium- and NO-dependent vasodilator acetylcholine. More importantly, the novel small molecule inhibitors were able to largely prevent TNFα-induced endothelial dysfunction (Fig. 6D). The protective role of Nox2 inhibition on endothelial function has been reported by several groups. For instance, in type 2 diabetic mouse models, Nox2ds-tat (aka gp91ds-tat)-treated mesenteric and coronary arteries exhibited improved vasodilatation to acetylcholine compared to untreated vessels [70]. Moreover, knocking out p47$^\text{HOX}$ as well as Nox2 prevented disruption of endothelium-dependent vasodilatation under pathological conditions [9,70].

In agreement with those findings, novel Nox2 inhibitors CPP11G and CPP11H appear to have conferred protection by abolishing TNFα-induced ROS production and ameliorating endothelial dysfunction. Further, disrupted endothelial function hampers the ability of arteries and arterioles to dilate in response to endogenous vasodilatory agents, and as a result, limits local and systemic blood flow. Indeed, we found that treating mice with TNFα significantly decreased hind limb blood flow, which is consistent with our results regarding endothelial dysfunction in aortas. Though these measurements were employed on different vascular beds (femoral arteries and aortas, respectively), it stands to reason that endothelial function in femoral arteries was also
impaired, leading to reduced hind limb blood flow. Surprisingly, though Nox2-selective inhibitor Nox2ds-tat was applied to cerebral circulation to identify the involvement of Nox2 in cerebral blood flow modulation [71,72], there are no other studies, to our knowledge, on Nox2-mediated systemic blood flow regulation under physiological or pathological conditions. Here, for the first time, we show that the novel Nox2 inhibitors CPP11G and CPP11H effectively preserved normal peripheral blood flow against inflammatory insults and protected vascular function. As a decrease in blood flow, particularly in resistance arteries, is associated with increase in blood pressure, our current result is partially supported by one of our previous studies in which Nox2ds-tat attenuated systolic blood pressure elevation in mice infused with AngII [35]. Both studies indicate that targeted Nox2 inhibition promotes vascular protection in various disease models, and that CPP11G and CPP11H hold significant promise clinically for preventing and treating Nox2-associated pathologies.

It is also noteworthy that CPP11H generally exhibits higher efficacy compared to CPP11G. In particular, in vivo results on mouse aortas illustrate that CPP11H reduced ROS production and ICAM expression perhaps to a higher degree than CPP11G. This might be attributed to the different structures of the side chains on the amine group: pentyl and thiophene functional groups for CPP11G and CPP11H, respectively [19]. These functional groups are distinct in their bulkiness, rigidity and hydrophobicity, all of which could influence drug-protein binding affinity. It is plausible that the thiophene functional group on CPP11H presents a better fit for the hydrophobic pocket of the “super groove” on the SH3 domain of p47phox for p22phox docking, thereby yielding higher efficacy in addition. The distinct functional groups could also affect drug absorption, distribution, metabolism and excretion in vivo, leading to varied efficacy. Interrogation of whether CPP11H possesses higher binding affinity, better tissue penetration, slower metabolism or less excretion than CPP11G will require thorough biochemical and pharmacokinetic studies that are currently beyond the scope this work.

Collectively, these data provide exciting new evidence for the role of Nox2 in vascular inflammation. They also illustrate effectiveness for novel small-molecule Nox2-selective inhibitors CPP11G and CPP11H in ameliorating TNFα-elicited endothelial inflammation and dysfunction in human cells in vitro and in mice in vivo. More development in the way of improving potency will likely be necessary to render more effective and clinically relevant agents. Nevertheless, their manifold and profound effects on key inflammatory pathways as well as cell and tissue function justify their development and appear to hold significant promise for this new class of therapeutics for Nox2-mediated inflammatory vascular diseases, such as atherosclerosis and restenosis.

4. Methods

4.1. Reagents

Superoxide dismutase (SOD) and catalase were purchased from Sigma-Aldrich (St. Louis, MO). Protease inhibitor and phosphatase inhibitor cocktail tablets were purchased from Roche Diagnostics GmbH (Mannheim, Germany). Antibodies for phospho-p38 (#4631), total p38 (#9228), phospho-SAPK/JNK (#9255), total SAPK/JNK (#9252S), phospho-p44/42 (#8544), total p44/42 (#4695), phospho-IκB/β (#2697), total IKKβ (#8943), total IκB (#4814), phospho-c-Jun (#3270), total cJun (#9165), phospho-NFκB p65 (#3033), total NFκB p65 (#8242) were purchased from Cell Signaling Technology (Danvers, MA). VCAM-1 (sc-13160), ICAM-1 (sc-19584), and β-Actin (sc-47778) antibodies were purchased from Santa Cruz Biotechnology (Dallas, TX). Nox2 antibody (ab129068) was purchased from Abcam (Cambridge, MA) and p47phox antibody (07-001) was purchased from Millipore (Burlington, MA). Rabbit (925-68070), mouse (925-68071) and goat (925-68074) secondary antibodies were purchased from LI-COR Biosciences (Lincoln, NE). The nuclear extract kit (40010) was purchased from Active Motif (Carlsbad, CA). Coumarin 7-boronic acid (CBA) (1357078-03-5) was purchased from Cayman Chemical (Ann Arbor, MI). Hydropropidine (HPr+) was generously provided by Dr. Jacke Zielonka (Medical College of Wisconsin).

4.2. 3-D modeling

Smina [73] was used to dock CPP11G and CPP11H to p40phox (from a p40phox-p47phox complex, PDB 1W70) [74], p47phox (from a p47phox-p22phox complex, PDB 10V3) [23], and p67phox (from a p67phox-p47phox complex, PDB 1K4U) [75] at the protein-protein interface. The docking box was defined by the corresponding ligand peptide of each structure. Water and cofactors were removed from the receptor protein, and an exhaustiveness level of 50 was used with other parameters kept at their defaults. Poses docked to p47phox exhibited > 1 kcal/mol improved binding affinity compared to poses docked to p40phox and p67phox.

4.3. Cell culture

COS-phox cells, kindly provided by Dr. Mary Dinauer, were cultured in Dulbecco’s Modified Eagle Medium (DMEM) with 4.5 g/l glucose, 1-glutamine and sodium pyruvate containing 10% heat-inactivated fetal bovine serum (FBS), 100 units/ml penicillin and 100 μg/ml streptomycin (complete media) (Gibco, Carlsbad, CA). Human aortic endothelial cells (HAECs) from at least three different donors were obtained from Lonza (CC-2535; Walkersville, MD) and cultured in EBM-2 medium containing EGM-2 bullet kit components (CC-3182, Lonza) at 5% CO2, 37°C. Passages 3 to 6 were used for subsequent experiments. COS-phox cells and HAECs were grown to 75-80% confluency and subjected to serum deprivation by incubation in Opti-MEM (Gibco, Carlsbad, CA) and endothelial cell media containing 10% of supplied growth factors and 0.2% FBS, respectively. After 16 h, cells were treated with CPP11G, CPP11H or vehicle (DMSO) 20 min prior to the addition of PMA (5 μmol/l) or TNFa (10 ng/ml).

4.4. Measurements of extracellular superoxide production by hydropropidine assay

Hydropropidine (HPr+) probe preparation and methodology were adopted and modified from a publication by Zielonka et al. [56]. HAECs seeded in 96-well, clear-bottom, black-sided plates were serum starved overnight. On the following day, wells were washed with PBS, and incubated in 80 μl assay buffer consisting of Hank’s Balanced Salt Solution supplemented with 25 mmol/l HEPES, 1% BSA, 10 μmol/l DTPA, 100 μmol/l L-NAME and 1 mmol/l taurine (L-NAME and taurine were added to scavenge peroxynitrite and hypochlorous acid respectively, which react with the probe) for 30 min in an incubator (37°C, 5% CO2). 250 μmol/l hydropropidine (SOD) was added to select wells under each treatment as negative controls. After incubation, probe solution was added to each well to reach a final concentration of 100 μmol/l HPr+ and 0.5 mg/ml salmon sperm DNA was added at a final reaction volume of 125 μl. Plates were placed in a Biotek Synergy 4 hybrid multimode microplate reader (preheated to 37°C), and read kinetically (every minute for 2 h) at excitation 400 nm and emission 574 nm. The average rate of fluorescence generation was determined over the linear portion of the response normalized to SOD negative controls.

4.5. Measurements of hydrogen peroxide (H2O2) production by coumarin boronic acid (CBA) assay

CBA probe preparation was adopted and modified from methods described by Zielonka et al. [57]. HAECs seeded in 96-well, clear-bottom, black-sided plates were serum starved overnight, washed with PBS, and incubated in assay buffer consisting of EC starvation media (phenol red free) in an incubator (37°C, 5% CO2) for 30 min. Assay buffer was supplemented with 10 μmol/l DTPA, 100 μmol/l L-NAME and 1 μmol/l taurine. 1 KU/ml bovine liver catalase (Sigma-Aldrich, St.
Louis, MO) was added to select wells as negative controls. After incubation, probe solution was added to each well to reach a final concentration of 0.5 mmol/l CBA (Cayman Chemicals, Ann Arbor, MI) at a final reaction volume of 125 µl. Plates were placed in a Biotech Synergy 4 hybrid multimode microplate reader (preheated to 37 °C), and read kinetically (every minute for 2 h) at excitation 350 nm and emission 450 nm wavelengths. The average rate of fluorescence generation was determined over the linear portion of the response normalized to cat- halase negative controls.

4.6. Measurements of H$_2$O$_2$ production by Amplex Red Assay

HAECs were collected in lysis buffer (Hank’s Balanced Salt Solution with Complete Mini protease inhibitor and PhosStop phosphatase inhibitor from Roche), and were lysed by five freeze/thaw cycles and passed through a 30-gauge needle five times to disrupt cells. The cell lysates were centrifuged at 1000 g for 5 min at 4 °C to remove unbroken cells, nuclei, and debris. Throughout all procedures, extreme care was taken to maintain the lysate at a temperature close to 0 °C. Lysates of human aortic endothelial cells or mouse aortic tissue were resuspended in Amplex Red assay mixture (25 mmol/l Heps, pH 7.4, containing 120 mmol/l NaCl, 3 mmol/l KCl, 1 mmol/l MgCl2, 0.1 mmol/l Amplex red (Invitrogen, Grand Island, NY)), and 0.35 U/ml horseradish peroxidase (HRP) in the presence and absence of catalase (300 U/ml). The reaction was initiated by the addition of 36 µmol/l NADPH (MP Biomedicals, Grand Island, NY). Fluorescence measurements were made using a Biotek Synergy 4 hybrid multimode microplate reader with a 530/25-excitation and a 590/35-emission filter. The reaction was monitored at 25 °C for 1 h.

4.7. Western blotting

Western blot experiments were performed as previously described [76,77]. Tissue or cells were homogenized in ice-cold lysis buffer. Supernatants were collected and lysates quantified using a Bradford assay (Bio-Rad, Hercules, CA). Total protein (30 µg) from cell lysates was added to Tris-glycine SDS sample buffer, boiled, resolved by SDS-PAGE and transferred onto nitrocellulose membrane (Bio-Rad). Blots were blocked with the Odyssey Blocking Buffer (LI-COR Biosciences, Lincoln, NE) and probed using rabbit anti-phospho-p38 (1:5000), mouse anti-p38 (1:1000), rabbit anti-phospho-SAPK/JNK (1:500), rabbit anti-SAPK/JNK (1:1000), rabbit anti-phospho-p44/42 (1:5000), rabbit anti-44/42 (1:1000), rabbit anti-phospho-IκB (1:1000), rabbit anti-β-Actin (1:2000), rabbit anti-p47phox antibody (sc-17845, Santa Cruz Biotechnology) using anti-p47phox antibody (sc-17845, Santa Cruz Biotechnology) using protein A/G plus agarose slurry. Immunoprecipitates were then subjected to Western blotting using the anti-p47phox antibody (07-001, Millipore) and anti-Nox2 antibody (ab129068, Abcam).

4.9. Measurement of p47phox membrane translocation

COS- phox cells were serum starved in Opti-MEM (Gibco) for 16 h, and stimulated with 5 µmol/l PMA for 1 h in the absence of CPP11G, CPP11H or vehicle (DMSO). Cells were washed with ice-cold PBS and collected in lysis buffer (Hank’s Balanced Salt Solution with Complete Mini protease inhibitor and PhosStop phosphatase inhibitor from Roche Diagnostics). The cells were lysed by five freeze/thaw cycles and passed through a 30-gauge needle five times to disrupt cells. The cell lysate was centrifuged at 1000 g for 5 min at 4 °C to remove unbroken cells, nuclei, and debris. Throughout all procedures, extreme care was taken to maintain the lysate at a temperature close to 0 °C. After pre-washing with protein A/G plus agarose (sc-2003, Santa Cruz Biotechnology), total cell homogenates were immunoprecipitated with anti-p47phox antibody and subjected to Western blotting using the anti-p47phox antibody (07-001, Millipore) and anti-Nox2 antibody (ab129068, Abcam).

4.10. Immunofluorescent imaging

Immunofluorescence analysis was performed on HAECs grown on gelatin-coated glass coverslips as well as 5 µm sections of paraffin-embedded mouse aortas. Samples were antigen-retrieved, fixed in 2% paraformaldehyde, permeabilized with 0.1% Triton X-100 and washed using PBS. Sections were blocked in PBS with 2% bovine serum albumin (BSA) for 1 h at room temperature. HAECs on coverslips were then incubated with anti-3-nitrotirosine (ab61392, Abcam, 1:100) for 1 h at room temperature, or anti-total-NFκB p65 antibody (#8242, Cell Signaling, 1:100) overnight at 4 °C followed by Cy3-conjugated secondary antibody (Life Technologies Inc., 1:1000). Coverslips were then stained for nuclei with Hoechst dye, and cover-slipped using gelvatol mounting media (polyvinylalcohol, glycerol, H2O, sodium azide and Tris pH 8.5). Non-specific rabbit or goat IgG (5 µg/ml) was used in lieu of primary antibody as a negative control. Confocal images were captured on a Nikon A1 spectral confocal microscope (Nikon Instruments Inc. Melville, NY). For each experiment, 3-4 images per treatment group were captured. Three independent experiments were performed.

4.11. Monocyte adhesion assay

Confluent HAECs grown in black 96-well, clear-bottom plates were pretreated with CPP11G or CPP11H, then stimulated with TNF-α (10 ng/ml) for 24 h. THP-1 monocytes (ATCC, Manassas, VA) were tagged with the compound calcein-AM (Life Technologies, Carlsbad, CA) (10 µmol/L) for 30 min at 37 °C in RPMI-1640 (Sigma-Aldrich, St. Louis, MO) and were washed once with PBS and resuspended in RPMI-1640 at a concentration of 5 × 10⁵ cells/ml. The tagged THP-1 (5 × 10⁴ cells/well) were gently added to the HAEC monolayer and incubated for 1 h at 37 °C. Unbound THP-1 cells were discarded by washing with RPMI-1640 3 times followed by PBS washing once. HAEC-attached THP-1 cells were observed with a Zeiss Axiovert 40 CFL microscope (Carl Zeiss Microscopy, Thornwood, NY) at 10 X magnification. In parallel, fluorescence intensity was determined using a fluorescence microplate reader (BioTek) at 495 nm (excitation wavelength) and 515 nm (emission wavelength). Results were shown as re- lative monocyte adhesion levels normalized to the control group.
4.12. Animals

All animal experiments were approved by the Institutional Animal Care and Use Committee, University of Pittsburgh and are in accordance with National Institutes of Health guidelines. Mice purchased from Jackson Labs (Bar Harbor, ME, USA) were subjected to i.v. bolus injections of CPPI11G or CPPI11H (15 mg/kg, 100 μl) or vehicle (30% DMSO in PBS, 100 μl) into the jugular vein 15 min prior to TNF-α treatment (10 μg/kg, 100 μl). Animals were allowed to recover for 4 h, then sacrificed and aortic vessels collected.

4.13. Laser Doppler blood flow analysis

Femoral artery blood flow was assessed 4 h after i.v. bolus injection of TNFα (10 μg/kg) in the presence or absence of CPPI11G or CPPI11H (15 mg/kg). Briefly, mice were anesthetized with 2.5% isoflurane and placed in a supine position on a heating pad. Core temperatures were maintained at 37 °C and continuously monitored by rectal probe thermometer. In order to reveal the femoral artery and adjacent arterioles and capillaries, the inner hind limb was shaved and the skin removed to better expose subcutaneous tissue. Real-time blood flow was measured using laser Doppler imaging (MoorLDI-2λ; Moor Instruments, Devon, UK).

4.14. Vessel myography

Male C57/B6 mice were anesthetized with CO2 followed by cervical dislocation. Descending thoracic aortas were quickly isolated and cut into 4 rings each 2 mm in length. Aortic rings were placed on myograph stirrups (Danish Myo Technology, Atlanta, GA) in 5 ml physiological saline solution (PSS) buffer (containing 130 mmol/L NaCl, 4.7 mmol/l KCl, 1.17 mmol/l MgSO4, 1.18 mmol/l KH2PO4, 14.9 mmol/l NaHCO3, 5.5 mmol/l D-glucose, 1.6 mmol/l CaCl2, 0.026 mmol/l EDTA) maintained at 37 °C, pH 7.4, gassed with 95% O2 and 5% CO2, and brought to an optimal resting tension of 5000 mg by increasing tone by 500 mg every 5 min. Vessels were then allowed to equilibrate until baseline tension remained constant (30-60 min). Viability of the vessels was ascertained by a series of contractile responses to KCl (25 mmol/l, 50 mmol/l and 100 mmol/l KCl in PSS solution) for 5 min each. Concentration-response curves to phenylephrine were carried out and a concentration that produced 80% maximum contraction (EC80) was chosen for establishing vascular tone prior to additional treatments. In PE-preconstricted vessels, endothelium-dependent dilation in response to acetylcholine (10−9 to 10−5 mol/l) and endothelium-independent dilation in response to sodium nitroprusside (SNP) (10−9.5 to 10−6 mol/l) were then tested.

4.15. Statistical analyses

Data are expressed as mean ± SEM, and for comparison of results between two data sets, an unpaired Student’s t-test was performed. One-way ANOVA followed by Tukey’s multiple comparison test was used for comparison of results among more than two groups. Both analyses were performed using GraphPad Prism software (version 7.1). p < 0.05 was considered statistically significant.

4.16. Study approval

All animal experiments were approved by the Institutional Animal Care and Use Committee, University of Pittsburgh and are in accordance with National Institutes of Health guidelines.

Author contributions

YL, ECP, PJP, DNM collected and analyzed experimental data. YL, ECP, PJP wrote and/or edited the manuscript. All authors reviewed and approved the final version.

Conflicts of interest

The authors have declared that no conflict of interest exists.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.redox.2019.101143.

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