Soluble guanylyl cyclase is a target of angiotensin II-induced nitrosative stress in a hypertensive rat model

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1Department of Pharmacology and Physiology, New Jersey Medical School, University of Medicine and Dentistry of New Jersey, Newark, New Jersey; 2G. P. Livanos Laboratory, Department of Critical Care and Pulmonary Services, University of Athens, Greece; and 3Laboratory of Molecular Pharmacology, Department of Pharmacy, University of Patras, Patras, Greece

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Crassous PA, Couloubaly S, Huang C, Zhou Z, Baskaran P, Kim DD, Papapetropoulos A, Fioramonti X, Durán WN, Beuve A. Soluble guanylyl cyclase is a target of angiotensin II-induced nitrosative stress in a hypertensive rat model. Am J Physiol Heart Circ Physiol 303: H597–H604, 2012. First published June 22, 2012; doi:10.1152/ajpheart.00138.2012. —Nitric oxide (NO) by activating soluble guanylyl cyclase (sGC) is involved in vascular homeostasis via induction of smooth muscle relaxation. In cardiovascular diseases (CVDs), endothelial dysfunction with altered vascular reactivity is mostly attributed to decreased NO bioavailability via oxidative stress. However, in several studies, relaxation to NO is only partially restored by exogenous NO donors, suggesting sGC impairment. Conflicting results have been reported regarding the nature of this impairment, ranging from decreased expression of one or both subunits of sGC to heme oxidation. We showed that sGC activity is impaired by thiol S-nitrosation. Recently, angiotensin II (ANG II) chronic treatment, which induces hypertension, was shown to generate nitrosative stress in addition to oxidative stress. We hypothesized that S-nitrosation of sGC occurs in ANG II-induced hypertension, thereby leading to desensitization of sGC to NO hence vascular dysfunction. As expected, ANG II infusion increases blood pressure, aorta remodeling, and protein S-nitrosation. Intravital microscopy indicated that cremaster arterioles are resistant to NO-induced vasodilation in vivo in anesthetized ANG II-treated rats. Concomitantly, NO-induced cGMP production decreases, which correlated with S-nitrosation of sGC in hypertensive rats. This study suggests that S-nitrosation of sGC by ANG II contributes to vascular dysfunction. This was confirmed in vitro by using A7r5 smooth muscle cells infected with adenoviruses expressing sGC or cysteine mutants: ANG II decreases NO-stimulated sGC activity in the wild-type but not in one mutant, C516A. This result indicates that cysteine 516 of sGC mediates ANG II-induced desensitization to NO in cells.

SOLUBLE GUANYLYL CYCLASE (sGC) is the main receptor for nitric oxide (NO) and the enzyme responsible for the conversion of GTP into cGMP. As such, the NO-receptor sGC is crucially involved in the physiology of the cardiovascular system by modulating vessel tone. Indeed, mice lacking this receptor are hypertensive (5). Oxidative stress is associated with cardiovascular diseases (CVDs) such as hypertension, atherosclerosis, and diabetes. Most oxidative CVDs are accompanied by endothelial dysfunction and impaired vascular reactivity with decreased NO bioavailability. Nonetheless, it should be pointed out that oxidative stress affects as well the smooth muscle cell (SMC) layers where sGC is expressed. It has been reported that reactive oxygen species (ROS) alter sGC expression and activity (19, 25, 29). We (21) have previously shown that in vitro and in vivo S-nitrosation of sGC impairs its ability to be activated by NO. In particular, we established that infusion of low therapeutic doses of nitroglycerin in rats for 3 days induces S-nitrosation of sGC and correlates with desensitization of sGC to NO stimulation. Importantly, the nitroglycerin treatment was associated with reduced relaxation of arterioles to exogenous NO donors, thus implicating directly sGC activity (22). Association between protein S-nitrosation and diseases such as diabetes (2, 30) has been reported but association with other CVDs such as hypertension has not been directly addressed. Angiotensin II (ANG II) is a peptide involved in blood pressure regulation and renovascular hypertension (11, 14, 18) and elevated levels of ANG II induce hypertension. ANG II infusion or treatment, which is clearly established as an inducer of oxidative stress via activation of NADPH oxidases in vivo (17) and in vascular smooth muscle cells such as A7r5 (31), was recently reported to generate global S-nitrosation (3).

Most studies have focused on endothelial dysfunction in ANG II-linked hypertension, yet a decreased NO responsiveness, e.g., NO resistance, has been observed and could be due to scavenging of NO by reactive oxygen species, to decreased sGC activity, or to a decreased sGC expression, as reported in overactive renin-angiotensin system of hypertensive TGR rats (7). Based on our finding that sGC is desensitized to NO by S-nitrosation leading to decreased relaxation in vivo and that ANG II generates nitrosative stress, we hypothesized that sGC activity could be impaired via S-nitrosation in ANG II-induced hypertension thereby contributing to decreased vascular reactivity. Thus we assayed in ANG II-induced hypertension rat model, the level of S-nitrosation of the sGC, whether sGC was desensitized to NO, and measured vascular NO resistance in vivo (so far studies by others on NO responsiveness and vascular reactivity have been done on isolated vessels in organ chamber). In parallel, we explored the molecular mechanism of sGC desensitization by ANG II in A7r5 SMC, which are depleted for endogenous sGC, by infecting the SMC with adenoviruses expressing wild-type (WT) sGC and Cys mutants of sGC and measuring NO-stimulated sGC activity in the infected cells treated or not with ANG II.

MATERIALS AND METHODS

Animal model. Sprague Dawley (SD) male rats (160–230 g, Charles River) were treated with ANG II (0.7 mg·kg−1·day−1, 7 days) or with vehicle (0.9% NaCl, 7 days) via mini-osmotic pump (Alzet, model 2002,
Durect; Cupertino, CA) implanted subcutaneously on animal anesthesia with sodium pentobarbital (50 mg/kg ip). The procedure was initiated when the withdrawal reflex was no longer present indicating an adequacy of anesthesia; the respiratory rate was monitored as well. Buprenorphine hydrochloride (0.03 mg/min) was administered as analgesic after surgery. Seven days after pump implantation, rats were anesthetized (sodium pentobarbital sodium; 50 mg/kg ip) and bled by abdominal aorta incision. Tissues (aorta, lungs, and mesenteric vasculature) were removed and washed in PBS and snap frozen in liquid nitrogen for storage or processed for cryosection as detailed below.

All experimental procedures conform to the Guide for the Care and Use of Laboratory Animals published by the United States National Institutes of Health and were approved by University of Medicine and Dentistry of New Jersey Institutional Animal Care and Use Committee (#10085D0114).

**Blood pressure.** Mean arterial pressure (MAP) was determined by tail-cuff plethysmography (Kent Scientific, CODA system) on nonanesthetized rats trained for a week before surgery of pump implantation. Measurements were performed before implantation of the osmotic pump and from 2 to 7 days after. Blood pressure was also monitored on anesthetized rats (pentobarbital, 50 mg/kg ip) via a catheter implanted in left carotid artery and connected to a Power Lab Pressure Monitor (AD instruments, Colorado Springs, CO) 7 days after pump implantation.

**Intravital microscopy.** SD male rats (160–230 g, Charles River) were anesthetized with sodium pentobarbital (50 mg/kg ip). The right cremaster muscle was prepared for intravital microscopy (10, 26). Pentobarbital sodium was continuously administered (0.03 mg/min, 0.1 ml/h iv) via catheter implanted in right jugular veins to maintain anesthesia. To assess the adequacy of anesthesia during the whole experiment, parameters such as responsiveness, blood pressure, respiratory rate, and heart rate were monitored. The recording system comprises an Optronics TEC-470 microscope camera (Optronics, Goleta, CA), a Sony monitor, and a MetaMorph image system (Universal Imaging, Downingtown, PA) for computer recording directly from the microscope camera and for image processing. Arteriolar luminal diameter was measured as the width of the transilluminated blood column using the MetaMorph image system. Two to three arterioles were studied per animal. Baseline diameters were normalized to a value of one. For each vessel, the experimental diameter was allowed to return fully to basal level, diameters were measured at the same place in the arteriolar lumen. After the excess of NEM was washed out, luminode was expressed as the ratio of baseline diameter (relative to three arterioles were studied per animal. Baseline diameters were measured directly from the microscope camera and for image processing. Arteriolar luminal diameter was measured as the width of the transilluminated blood column using the MetaMorph image system. Two to three arterioles were studied per animal. Baseline diameters were normalized to a value of one. For each vessel, the experimental diameter was allowed to return fully to basal level, diameters were measured at the same place in the arteriolar lumen. After the excess of NEM was washed out, luminode was expressed as the ratio of baseline diameter (relative

**S-nitrosation detection in tissues.** Aortas from control or ANG II-treated rats were embedded in Tissue-Tek O.C.T compound (Sakura Finetek) and frozen. Ten-micrometer transversal cryosections were mounted on glass slides (Superfrost plus, VWR) and processed for S-nitrosation detection.

To determine the level of S-nitrosation in situ, transversal cryosections were fixed and permeabilized with 3% paraformaldehyde and 0.2% Triton (10 min) and then incubated in 1% BSA for 1 h. Slides were treated overnight with primary antibody (rabbit anti-S-nitrosation, Alpha Diagnostic no. NISC 11-A at 1/100 dilution in PBS with 1% BSA) followed by 1 h incubation with secondary antibody (Alexafluor 594 Invitrogen, 1/100 dilution in PBS with 1% BSA). A coverslip was applied on top of preparation with Prolong gold anti-fading containing DAPI (Invitrogen). Cryosections of aorta from ANG II-treated rats were used for negative control as follows: slides were treated with ascorbate (10 mM, 1 h) to reduce S-nitrosated cysteine of proteins prior to incubation with anti-S-nitrosation and secondary antibodies as described by Ckless et al. (4).

To confirm S-nitrosation in situ, aorta transversal cryosections were fixed and permeabilized with 3% paraformaldehyde and 0.2% Triton for 10 min. After wash, slides were incubated in PBS, 0.4 mM EDTA, 0.04 mM neocuproine, containing 1% SDS and 40 mM N-ethylmaleimide (NEM) for 30 min at 40°C. After the excess of NEM was washed, S-nitrosation bonds were switched to S-biotin bonds: slides were incubated in PBS-EDTA-Neocuproine as above with 10 mM ascorbate and 0.1 mM HPDP-biotin for 1 h. After wash and blocking (PBS, 1% BSA, 1 h), biotin was detected by a primary antibody anti-biotin (Abcam, 1/250, overnight) followed by secondary antibody Alexafluor 594. Images were obtained with 200M Axiovit fluorescence microscope (Zeiss) and analyzed with Axiovision software.

**sGC activity.** cGMP production in tissues and sGC activity in cells were assayed as previously (22). cGMP production was measured by radioimmunoassay in lungs from control or ANG II-treated rats, incubated with 0.5 mM IBMX (a wide spectrum phosphodiesterase inhibitor) and treated with NO donor [0.1 mM diethylamine-NO (DEA-NO) or 0.01 mM S-nitroso-N-acetyl-penicillamine (SNAP)] for 1 h at 37°C. The reaction was stopped with 2.5% perchloric acid. Following neutralization with 0.75 N KOH, cGMP and cAMP were measured by radioimmunoassay. cAMP, measured in the same samples, was used to normalize cGMP data as cAMP is a good indicator of tissue amount used in the assay and is independent of SNAP or DEA-NO stimulation and because the increase in cAMP levels in response to NO is negligible compared with increased NO-dependent cGMP levels. assays of protein concentration (before perchloric acid lysis) correlated with estimated cAMP levels. sGC activity was determined by formation of [α-32P]GMP from [α-32P]GTP under basal (unstimulated) and NO-stimulated (DEA-NO; 1 μM) conditions for 5 min at 33°C. Typically, 40 μg of cytosol were used in each assay reaction. Reaction mix contained 50 mM HEPES pH 8.0, 5 mM MgCl2, and 0.5 mM GTP.

**Biotin switch assay.** sGC S-nitrosation was determined by the biotin switch assay (9) as modified by us; 0.5 to 1 mg/ml proteins from lungs or cells lystate were incubated in blocking buffer (250 mM HEPES, 150 mM NaCl, 0.1 mM neocuproine, 1 mM EDTA, 2.5% SDS, and 50 mM NEM) at 40°C for 30 min in the dark. Proteins were

**Fig. 1. ANG II induces aorta remodeling.** Width of the media of thoracic aorta from control (CTRL; n = 4) and ANG II-treated (ANG II; n = 5) rats were determined on transversal cryosections (bright field, ×10 magnification). **P < 0.0005.
precipitated in 3 vol of acetone to remove NEM and resuspended in 25 mM HEPES containing 1% SDS and then treated with HPDP-biotin (0.2 mM) with or without ascorbate (1 mM) for 1 h at room temperature. After addition of 2 vol of neutralizing buffer (20 mM HEPES, 100 mM NaCl, 1 mM EDTA, and 0.5% Triton), biotinylated proteins were purified with streptavidin-agarose resin beads (Thermo Scientific) for 1 h and eluted in Laemmli reducing buffer (Sigma). Proteins incubated with HPDP-biotin without ascorbate were used as negative control for the experiment, e.g., specificity of biotinylation.

Western blot. Proteins from lungs or mesenteric vasculature from control or ANG II-treated rats or from biotin switch assay or cell lysates were separated by SDS-PAGE and transferred to nitrocellulose membrane. After 1 h blocking in PBS containing 5% BSA, membranes were incubated with sGC anti-α-subunit 1/10,000 (Sigma), sGC anti-β-subunit 1/1,000 (Cayman Chemical), or β-actin 1/5,000 (Sigma). The appropriate secondary antibody coupled with peroxidase (anti-rabbit or anti-mouse 1/10,000, Amersham) allowed protein detection by chemiluminescence with ECL kit (GE Healthcare). Signal quantification of blots were performed with ImageJ software (W. S. Rasband; National Institutes of Health, Bethesda, MD).

Cell culture, infection and lysates preparation. Rat primary vascular SMC (A7r5; ATCC), which do not express detectable sGC, were used between passages 4 to 10. They were grown in DMEM (with L-glutamine and 5% BSA) until 80% confluence and infected with 1 to 4 multiplicity of infection of adenoviruses expressing α-WT or αC516A-subunit with β-WT subunit of sGC. 48 h after infection, cells were harvested in “sGC activity buffer” [50 mM HEPES, 150 mM NaCl, antiprotease inhibitors (tablet, Roche), and 35 μg/ml PMSF] or “biotin switch buffer” [250 mM HEPES, 150 mM NaCl, 1 mM EDTA, 0.1 mM neocuproine, antiprotease inhibitor (tablet, table)] and processed for biotin switch assay.

Fig. 2. S-nitrosation (S-NO) is increased in ANG II-treated rats. A: representative images (×10) of transversal cryosection of thoracic aorta from control or ANG II-treated rats. Aortas were incubated with anti-S-NO and revealed with secondary antibody Alexafluor594. As control for specificity of S-NO detection, aortas from ANG II-treated rats were preincubated for 1 h with 10 mM ascorbate (Asc), which reduces S-NO bonds. Corresponding bright field images are shown at top. B: fluorescence image analysis from aorta cryosections of control (CTRL, n = 5), ANG II (n = 6), and ANG II + Asc (n = 5) indicates significant and specific S-NO by ANG II. **P < 0.005, ANG II vs. CTRL; *P < 0.05, ANG II vs. ANG II + Asc; NS: nonsignificant, CTRL vs. ANG II + Asc. Signal intensity was quantified at same exposure time between groups and normalized to control. C: representative images (×20) of S-NO level detected by biotin switch assay on transversal cryosection of thoracic aorta from control or ANG II-treated rats. Biotin switch assay was performed on aortic cryosections, and the level of biotin was detected by immunofluorescence using a primary antibody anti-biotin (Abcam) revealed by secondary antibody Alexafluor594, as described in MATERIALS AND METHODS. Same exposure times were used in each experiment. As control for specificity of S-NO detection, aortas from ANG II-treated rats were preincubated with 10 mM ascorbate (Asc). Corresponding bright field images are shown at top.
Roche), and 35 μg/ml PMSF] and sonicated on ice. Debris were removed by centrifugation at 16 000 g for 5 min at 4°C, and the supernatants (cytosolic fraction) were aliquoted and snap frozen. Protein concentration was determined by Bradford method. Adenoviral construction and site-directed mutagenesis were done as in (21, 32).

Statistical analysis. Data are expressed as means ± SE. For statistical analysis, unpaired Student’s t-test was used to compare ANG II-treated and control groups and one-way ANOVA (Tukey’s post hoc test) was used to compare more than two groups. Analysis was done with SigmaPlot 11.0. *P* < 0.05 was considered significant.

RESULTS

**ANG II-induced hypertension causes oxidative/nitrosative stress.** We analyzed MAP, oxidative/nitrosative stress level, and aortic remodeling in control rats and in rats treated with ANG II. Administration of ANG II significantly increased MAP relative to control rats [tail-cuff measurement: ANG II (n = 4) 140.9 ± 9.7 vs. control (n = 4) 100.8 ± 3.4 mmHg; *P* < 0.0005]. The increase in MAP in ANG II-treated rats was confirmed by recording carotid pressure directly via catheter on anesthetized rats (control 114 ± 3 vs. ANG II 158 ± 8 mmHg; *P* < 0.0005; n = 7). Vascular hypertrophy was demonstrated by the significant enlargement of the aortic media wall width in ANG II-treated rats (Fig. 1) with extensive remodeling of the aorta wall (Fig. 1, bottom). We have previously determined in vitro that S-nitrosation of sGC causes a decreased response to NO; thus we measured whether S-nitrosation following ANG II infusion occurs. Using immunocytochemistry with anti-S-nitrosothiol antibodies, we showed that ANG II significantly increases the level of S-nitrosation in the media of aorta. The specificity of S-nitrosation detection was confirmed by treatment with ascorbate (Fig. 2, A and B). This increase in S-nitrosation levels was confirmed by conducting a biotin-switch assay on the aortic tissues (Fig. 2C). These results indicate that at day 7 of ANG II treatment, rats have developed hypertension, nitrosative stress, and aorta remodeling. These results confirm previous studies (3, 16, 17, 27, 28) and show that ANG II-induced hypertension is an adequate model to study the relevance of oxidative and nitrosative stress-induced alterations on sGC activity and their impact on vascular pathology.

**NO resistance in arterioles of cremaster muscle.** To assay whether ANG II-induced hypertension correlates with reduced vascular reactivity to NO, we measured in vivo arterioles vasodilation to NO donors by intravital microscopy of cremaster muscle arterioles. S-nitrosylated tissue such as mesentery (Fig. 4, A) and lung (Fig. 4, B) were used for ANG II-treated rats. No significant change in sGC expression was observed between the two NO-donor applications. Relative luminal diameter from 2 to 3 arterioles was assessed for each preparation. Baseline mean diameters of the selected arterioles were 49 ± 2 μm in the control group and 46 ± 4 μm in the ANG II treated group and not significantly different. The reduced increase in relaxation in response to SNAP and DEA-NO in ANG II-treated rats was significant, *P* < 0.05 and *P* < 0.0001, respectively.

NO-stimulated conditions in lungs. Lungs were chosen because they are highly vascularized tissues with high expression of sGC, facilitating cGMP detection. Figure 4A indicates that basal cGMP production was similar in lung of control and ANG II-treated rats whereas cGMP production in response to the NO donors SNAP (100 μM) and DEA-NO (10 μM) was drastically reduced in ANG II-treated rats. No significant change in sGC expression was observed between the Ctrl and ANG II-treated rat lungs (Fig. 4B). There was also no change in sGC expression in other vasculature tissues such as mesentery (Fig. 4B). These results indicate that NO-stimulated sGC activity per se is impaired in ANG II-induced hypertensive rats.

**sGC is nitrosated in ANG II hypertensive rats.** We next investigated whether sGC was S-nitrosated under these conditions. Using the biontin-switch assay (see MATERIALS AND METHODS and Ref. 9), we detected a strong S-nitrosation of sGC in the lungs of ANG II-treated rats compared with untreated rats (Fig. 5, A and B). Similar levels of sGC in input (biotinylated samples from ANG II and untreated rats) confirm that ANG II infusion did not modify sGC expression (Fig. 5B). Those data indicate that ANG II treatment leads to S-nitrosation of sGC.

These results show a correlation between in vivo blunted NO-induced relaxation of arterioles (NO resistance), decreased NO-dependent sGC activity, and increased sGC S-nitrosation in ANG II-induced hypertensive rats. To determine the potential causal relationship between ANG II treatment and NO resistance via thiol oxidation of sGC, we used an in vitro adenosviruses/cell system.

**sGC is desensitized to NO in A7r5 cells treated with ANG II.** A7r5 SMC, which do not express detectable level of sGC, were infected with adenoviruses expressing the α- and β-WT subunits of sGC. After 48 h, cells were treated or not with 100 nM ANG II for 4 h and their cytosolic fractions were prepared (32). As shown on Fig. 6A, basal sGC activity was similar in the
cytosolic fraction of untreated and ANG II-treated samples. On the other hand, NO-stimulated sGC activity was markedly inhibited (49.7 ± 4.4%) by ANG II treatment, while expression of sGC in both cytosolic fractions was similar (inset). In parallel, biotin switch assay showed that ANG II treatment increased S-nitrosation of sGC (data not shown). Those results indicate that ANG II affects directly NO-stimulated sGC activity, potentially via S-nitrosation, in agreement with results obtained in vivo.

Cysteine 516 (C516) in the α-subunit of sGC mediates ANG II-dependent desensitization to NO. To establish a causal relationship between S-nitrosation of sGC by ANG II and its desensitization to NO, we conducted site directed mutagenesis of C516, a cysteine newly identified by mass spectrometry as

Fig. 4. NO-induced cGMP production is decreased in ANG II treated rats. A: lung of control (CTRL, n = 5) and ANG II-treated (ANG II, n = 5) rats were incubated with IBMX (500 μM) for 15 min, followed, or not (Basal), by incubation with NO donors (SNAP, 100 μM; DEA-NO, 10 μM) for 1 h at 37°C. cGMP production was measured by radioimmunoassay; *P < 0.05; **P < 0.001. B: Western Blots (WB) with antibodies against α- and β-subunit of sGC and β-actin indicate that sGC expression is similar in lysate from lung (top) and mesentery vasculature (bottom) of control and ANG II-treated rats (n = 8 in lungs and n = 4 in mesentery; P > 0.05, CTRL vs. ANG II).
S-nitrosated and highly conserved in the catalytic domain of sGC. Adenovirus containing the C516A mutation in the α subunit was coinfected with the WT β-subunit. Cytosolic fractions of the infected A7r5 SMC treated or not with ANG II were prepared as above. Figure 6B indicates that replacement of C516 with Ala confers resistance to NO desensitization, as no decrease in NO-stimulated sGC activity was observed following ANG II treatment. Expression of αC516A/β was unchanged by ANG II treatment (Fig. 6B, inset). This result shows that ANG II affects NO-stimulated sGC activity via a thiol-dependent mechanism. It should be noted that in the absence of ANG II, both NO-stimulated and ANG II treatment was similar between WT and C516A mutant (10,475 ± 1,435 vs. 9,874 ± 1,328 pmol cGMP·min⁻¹·mg⁻¹). This is important as it indicates that in the absence of ANG II-induced stress, the NO-stimulated sGC activity is not affected by the mutation C516A. We conducted the same experiment with adenovirus expressing mutant αC243A, as C243 is known to mediate sGC desensitization following treatment with S-nitrosocysteine (CSNO) or nitroglycerin (21, 22). However, ANG II affected NO-stimulated activity of C243A as it did for WT (not shown), suggesting that C243 does not participate in the ANG II desensitization effect.

**DISCUSSION**

In this study, we tested the hypothesis that sGC activity could be impaired by S-nitrosation leading to decreased vascular reactivity in vascular diseases associated with generation of oxidative/nitrosative stress. We used a model of ANG II-induced hypertension because it increases oxidative stress via activation of NAD(P)H oxidase (17) and S-nitrosation in tissues (3). Our data (Fig. 2) confirmed that nitrosative stress occurs in vascular SMC, the site of expression of sGC in the vascular system. Many studies have reported that overactivation of the renin-angiotensin system, as mimicked in ANG II-induced hypertension model, leads to endothelial dysfunction with decreased NO bioavailability. An impaired NO production (15) or NO being scavenged by reactive oxygen species has been proposed as an explanation of the decreased vascular reactivity and of the increased vascular wall remodeling that takes place in hypertension (for recent reviews see Refs. 20, 23). Meanwhile, it was reported that NO responsiveness was also decreased by this overactive renin-angiotensin system suggesting impairment downstream of NO production (1, 8). Some groups (1, 7, 15) have proposed that this vascular desensitization to NO was due to downregulation of sGC, and recently one group (13) observed that stimulation of sGC by the NO-independent activator (Bay 41–2272) could prevent cardiovascular remodeling in a similar ANG II hypertensive animal model. These observations pointed to a role of sGC in ANG II-induced hypertension, but the mechanism by which sGC was involved in decreased NO responsiveness remained controversial.

Our studies show that ANG II infusion induces NO resistance in arterioles of the rat cremaster muscle that was characterized by a significant decrease in vasodilation in response to two different types of NO donors. Importantly, we studied the vascular NO resistance in resistant arterioles (cremaster muscle arterioles) because they are largely involved in blood pressure regulation and in ANG II hypertensive effect (24) whereas most of previous investigations focused on conduit vessels (aorta and arteries). Indeed, remodeling of the arteriolar wall and changes in smooth muscle reactivity is associated with vascular resistance during the development of hypertension (12, 33). More importantly, to our knowledge, this is the first report showing that ANG II impairs vascular reactivity via decreased NO responsiveness in vivo, as previous studies (3, 15) used an ex vivo system in which isolated blood vessels reactivity is measured in organ chamber. Blunted arterioles responsiveness to NO in hypertensive rats is unlikely due to NO scavenging because of the high NO donors concentrations we used (10 μM SNAP and 1 μM DEA-NO). This strongly suggested an impairment of sGC and led us to investigate its activity.

The rats treated with ANG II exhibited a significant decrease in NO-dependent cGMP production. Those experiments were done in the presence of IBMX, indicating that sGC activity per se was affected rather than activation of phosphodiesterase to explain the decreased cGMP (6). These results obtained in rat lungs are in agreement with the recently reported decrease of cGMP production in aorta of ANG II-induced hypertension mice (3). Of note, the level of expression of sGC was not

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**Fig. 5. ANG II induces S-NO of sGC.** A: representative WB with anti-sGC of biotin switch assay showing S-NO of sGC in lung of ANG II-treated rats (n = 7) vs. control (n = 6). Input shows the amount of sGC in each sample before streptavidin purification of the biotinylated samples (see MATERIALS AND METHODS). B: level of sGC S-NO in lung from control (n = 6) or ANG II-treated rats (n = 7) normalized to input and expressed as fold of CTRL; *P < 0.05.
decreased in lung or mesenteric vasculature of the ANG II-treated rats. This result is different from a previous study (15) showing that ANG II infusion led to decreased expression of sGC; this discrepancy might be due to a higher concentration used by the other group (1 mg vs. 0.7 mg·day^{-1}·kg^{-1}) and the tissues analyzed were aortic sections. Altogether these results indicate that the NO-dependent activity of sGC in SMC is directly impaired in the ANG II-induced hypertension model and is associated with a decrease in NO-dependent relaxation in vivo. Thus the next question was the mechanism by which the NO-stimulated sGC activity was decreased.

We (21) recently demonstrated that S-nitrosation of sGC causes desensitization to NO stimulation, and therefore, we hypothesized that S-nitrosation of sGC by ANG II could be the cause of the reduced NO responsiveness. In addition, global S-nitrosation was observed in homogenates of aortic tissues in ANG II-treated mice (3). We confirmed by another means, immunocytochemistry with anti-S-nitrosation antibodies and biotin switch assay in situ, that ANG II infusion led to significant S-nitrosation of aortic tissues (Fig. 2). More importantly, we specifically detected sustained S-nitrosation of sGC (Fig. 5), thus establishing a potential causal relationship between decrease NO vascular reactivity and S-nitrosation of sGC by ANG II.

To further examine whether the thiols of sGC were a specific target of ANG II-induced nitrosative stress and a potential explanation for decreased NO-stimulated activity, we used A7r5 cells that do not express detectable level of sGC and were previously shown to respond to ANG II treatment (31). Infection with wild-type sGC-expressing adenoviruses and treatment with ANG II leads to 50% desensitization of sGC to NO (the basal activity remained unchanged, Fig. 6A) and S-nitrosation of sGC. We established that this NO desensitization of sGC was thiol dependent as a single mutation of a cysteine in sGC (C516A) confers resistance to the ANG II-induced sGC desensitization in A7r5 SMC (Fig. 6B). This mutational analysis establishes, in vitro, a causal relationship between ANG II-induced NO desensitization and thiol oxidation of sGC. This study is the first to establish a link between ANG II treatment and thiol-dependent inactivation of sGC via S-nitrosation of sGC, yet further experiments using C516A knock-in animals will be necessary to establish a causal relationship in vivo in ANG II-induced hypertension model.

Interestingly, the replacement of another Cys in the α-subunit of sGC, C243, which is known to mediate desensitization induced by treatment with the nitrosating agent CSNO and with nitroglycerin (21, 22) does not prevent ANG II-induced desensitization as C516A does. This suggests that the S-nitrosation is specific and a function of the type of nitrosative stress. We speculate that C243 is nitrosated by low molecular weight S-nitrosothiols such as CSNO but might not be modified by ANG II-induced nitrosative stress, potentially in the form of N₂O₃. Alternatively, the thiol of C243 might not be as sensitive to oxidation (for creation of a reactive thiolate) as C516. Further experiments comparing sGC S-nitrosation status under various oxidative and nitrosative stresses are needed to address this question.

In summary, our in vivo and in vitro studies demonstrate that nitrosation of sGC correlates with NO resistance syndrome in ANG II-induced hypertension and that Cys 516 of the α-subunit of sGC is a specific target of ANG II-induced oxidative/nitrosative stress.

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