S-Nitrosothiols signal hypoxia-mimetic vascular pathology

Lisa A. Palmer,1 Allan Doctor,1 Preeti Chhabra,1 Mary Lynn Sheram,1 Victor E. Laubach,2 Molly Z. Karlinsey,3 Michael S. Forbes,1 Timothy Macdonald,3 and Benjamin Gaston1

1Department of Pediatrics and 2Department of Surgery, University of Virginia School of Medicine, and 3Department of Chemistry, University of Virginia, Charlottesville, Virginia, USA.

NO transfer reactions between protein and peptide cysteines have been proposed to represent regulated signaling processes. We used the pharmaceutical antioxidant N-acetylcysteine (NAC) as a bait reactant to measure NO transfer reactions in blood and to study the vascular effects of these reactions in vivo. NAC was converted to S-nitroso-N-acetylcysteine (SNOAC), decreasing erythrocytic S-nitrosothiol content, both during whole-blood deoxygenation ex vivo and during a 3-week protocol in which mice received high-dose NAC in vivo. Strikingly, the NAC-treated mice developed pulmonary arterial hypertension (PAH) that mimicked the effects of chronic hypoxia. Moreover, systemic SNOAC administration recapitulated effects of both NAC and hypoxia. eNOS-deficient mice were protected from the effects of NAC but not SNOAC, suggesting that conversion of NAC to SNOAC was necessary for the development of PAH. These data reveal an unanticipated adverse effect of chronic NAC administration and introduce a new animal model of PAH. Moreover, evidence that conversion of NAC to SNOAC during blood deoxygenation is necessary for the development of PAH in this model challenges conventional views of oxygen sensing and of NO signaling.

Introduction

NO transfer reactions between protein and peptide cysteines have been proposed to represent regulated signaling processes (1, 2). For example, NO transfer from deoxygenated erythrocytes to glutathione ex vivo forms S-nitrosoglutathione (GSNO) (3). GSNO can signal acute vascular and central ventilatory effects characteristic of oxyhemoglobin desaturation (3–4) that are regulated by γ-glutamyl transpeptidase (GGT), GSNO reductase (GSNOR), and other enzymes (1, 3–6). However, direct measurement of S-nitrosothiol signaling in vivo has proven challenging because of the metabolism and tissue-specific localization of endogenous S-nitrosothiol species (1, 3, 4, 6). We have addressed these challenges by using N-acetylcysteine (NAC) as a bait reactant, allowing the stable NO transfer product, S-nitroso-N-acetylcysteine (SNOAC), to be distinguished by mass spectrometry (MS) from endogenous S-nitrosothiols. We report that NAC is converted to SNOAC in mice in vivo. Furthermore, chronic, systemic administration of either NAC or SNOAC to mice causes hypoxia-mimetic pulmonary arterial hypertension (PAH). These data reveal a previously unappreciated vascular toxicity of NAC and of S-nitrosothiols. Moreover, they suggest that S-nitrosothiol transfer reactions can signal hypoxia in vivo.

PAH is characterized by increased pressure in the pulmonary arteries (PAs), increased RV weight, and thickening and remodeling of small PAs. Untreated human PAH can progress to right heart failure and death (7, 8). Chronic hypoxia can cause PAH, and hypoxia-regulated genes in the pulmonary endothelium play a role (7, 9). The partial pressure of O2 (pO2) of blood in the hypoxic PA is not as low as that used to study the expression of hypoxia-modified genes in vitro (10). However, endothelium-derived NO in erythrocytes can be transferred as a nitrosonium (NO+) equivalent to cysteine thiolates on glutathione (GSH) and anion exchange protein 1 (AE1) — forming GSNO and S-nitroso-AE1 — during erythrocytic oxyhemoglobin desaturation in vitro; this effect is dependent on erythrocytic oxyhemoglobin saturation rather than pO2 (3, 4, 11, 12). S-Nitrosothiols, in turn, can increase the expression of hypoxia-associated gene regulatory proteins in vitro (5, 13, 14), an effect regulated by S-nitrosothiol metabolic enzymes (5, 14). Here, we report a hypoxia-mimetic effect, pulmonary vascular remodeling, that is associated with chronic systemic conversion of NAC to SNOAC in vivo. Strikingly, chronic exogenous administration of SNOAC — a vasodilator (15) — recapitulates the effects of hypoxia on the lung, causing hypoxia-mimetic murine PAH. Furthermore, experiments in eNOS-deficient mice reveal that eNOS is necessary for NAC, but not SNOAC, to cause PAH. Taken together, these data demonstrate an unanticipated potential toxicity of chronic, systemic NAC and SNOAC administration and suggest a novel paradigm for understanding hypoxia-associated disease.

Results

NAC and SNOAC cause hypoxia-mimetic PAH. Three weeks of normobaric hypoxia (10%) increased RV systolic pressure, RV weight (relative to LV and septum [LV+S]) and muscularization of small (<80-μm) pulmonary arterioles in both C57BL/6 and C57BL/6/129SvEv mice (n = 4–14 each; Figure 1, A–E). The effects of 3 weeks’ exposure to either NAC (10 mg/ml [52 mM]) or SNOAC (1 mM) in the drinking water were similar to those of 3 weeks’ exposure to hypoxia in C57BL/6/129SvEv mice, increasing RV pressure (n = 3–4 each, P < 0.001 relative to normoxia), RV/LV+S weight ratio (n = 6–35 each; P < 0.02 relative to normoxia), and muscularization of the small pul-
monary arterioles. \( n = 3–8; P < 0.003 \) relative to normoxia) (Figure 1, A–E). Of note, PA pressures in the treatment groups were greater than 125% those in controls in the hypoxia, NAC, and SNOAC groups, despite anesthesia, which can blunt PAH (ref. 8; Figure 1C). Similar results were obtained in C57BL/6 mice \( n = 4–12 \) each); however, the increase in RV/LV+S ratio was not significant in these mice in the case of SNOAC (Figure 1, A and B).

In dose-response experiments, higher-dose NAC (10 mg/ml) increased RV weight relative to normoxia at 3 weeks in C57BL/6/129SvEv mice (Figure 1A); however, at a lower dose (1 mg/ml), RV/LV+S was normal at 3 weeks \( 0.27 ± 0.005; n = 5; P = \text{NS compared with no treatment} \). On the other hand, low-dose NAC increased RV systolic pressure at 3 weeks to 27.5 mmHg \( n = 4; P < 0.04 \) compared with normoxia).

In time course experiments performed using C57BL/6/129SvEv mice, there was a significant increase in RV pressure after 1 week of NAC exposure (10 mg/ml; mean RV pressure with NAC, 22.8 ± 3.3 mmHg, versus without NAC, 13.7 ± 0.7 mmHg; \( n = 5–8 \) each; \( P < 0.01 \)), persisting at 2 and 3 weeks. RV/LV+S increased more slowly: a significant difference between NAC-exposed and -unexposed animals was not observed until 3 weeks \( n = 5–8 \) each at 1 and 2 weeks; \( P = \text{NS} \). Concomitant 3-week exposure to both hypoxia and 10 mg/ml NAC did not result in a greater increase in mean RV/LV+S \( 0.35 ± 0.082 \) after NAC and hypoxia together versus 0.34 ± 0.020 after hypoxia alone; \( n = 4–6 \) each; \( P = \text{NS} \) or RV pressure \( 21.4 ± 1.30 \) mmHg after NAC plus hypoxia versus 19.9 ± 1.03 mmHg after hypoxia alone; \( n = 4–6 \) each; \( P = \text{NS} \), suggesting functional overlap between hypoxia- and NAC-stimulated pathways.

Whole-lung expression of hypoxia-inducible mitogenic factor (HIMF), fibronectin, and eNOS, proteins associated with the development of PAH in some models (16–20), was increased by hypoxia and by 3 weeks of NAC treatment (10 mg/ml; \( n = 3–6 \) each; \( P < 0.05 \) [HIMF and fibronectin] and \( P < 0.01 \) [eNOS]; Figure 2, A–D). Of note, VEGF-A and endothelin 1 protein expression did not increase significantly with hypoxia or NAC in our model (Figure 2, E and F). Though these proteins can be upregulated by low pO\(_2\) in vitro, results are variable in PAH models (21–26), suggesting that PAH-associated vascular remodeling is more complex than would be predicted based on the effects of low pO\(_2\) alone. Indeed, VEGF and
eNOS expression can be associated with both protection against PAH and development of PAH, depending on the model and time course (18–29). NAC did not change expression of neuronal or inducible NOS isoforms at 3 weeks (data not shown). NAC exposure increased whole-lung HIMF mRNA expression and transiently increased eNOS mRNA (Figure 2, D and G). Interestingly, parallel increases in lung protein and mRNA were not observed for the expression of some genes known to be upregulated in hypoxia in vitro, either because of nonvascular expression measured in whole-lung homogenates and/or because of the complexity of pathways involved in the response to hypoxia and the development of PAH in vivo. For example, mRNA levels for VEGF-A increased after 3 weeks of NAC treatment (n = 3; P < 0.04; Figure 2G), though there was no significant change in VEGF-A protein levels (Figure 2E). On the other hand, the change in mRNA expression for fibronectin was not significant (n = 2–5 each; P = NS; Figure 2G), though expression of the corresponding protein increased (described above; Figure 2A). Further, mRNA for glucose transport protein–1 (Glut-1), conventionally upregulated by hypoxia, did not increase (n = 3; P = NS). Consistent with this complexity, our time course analysis revealed an NAC-induced increase in whole-lung eNOS mRNA that preceded the NAC-induced increase in eNOS protein expression but was transient (Figure 2D).

Three weeks of NAC (10 mg/ml) did not affect systemic or portal vascular morphology (Supplemental Data; supplemental material available online with this article; doi:10.1172/JCI29444DS1) or hemoglobin (Hb) (12.5 ± 0.64 g/dl after NAC treatment versus 12.1 ± 0.65 g/dl in controls). Likewise, 3 weeks of oral SNOAC (1 mM) did not affect the systemic or portal vasculature (Supplemental Data).

NAC is converted to SNOAC in vivo and during erythrocytic deoxygenation in vitro and in vivo. The chemical mechanisms, including both inhibition of nitrosative/oxidative stress (30–32) and NO transfer chemistry (1, 33), by which NAC could cause PAH were investigated. NAC did not affect pulmonary vascular immunostaining for 3-nitrotyrosine (Supplemental Data), suggesting that its ability to cause PAH was not simply the result of tissue injury associated with altered nitrosative or oxidative stress.

In contrast, the ratio of total erythrocytic S-nitrosothiol concentration to Hb (SNOHb) (11) in NAC-treated mice (1.0 ± 10⁻⁴ ± 0.12 ± 10⁻⁴ for NAC and 1.0 ± 10⁻⁴ ± 0.12 ± 10⁻⁴ for controls, for 3 weeks) was higher than that in controls (1.0 ± 10⁻⁴ ± 0.12 ± 10⁻⁴ for NAC and 1.0 ± 10⁻⁴ ± 0.12 ± 10⁻⁴ for controls, for 3 weeks), suggesting that NAC caused PAH by increasing SNOHb levels and/or by increasing the amount of SNOAC formed in vivo.
Figure 3
SNOAC is formed from NAC in blood ex vivo and in vivo. (A) The SNO_{rbc} in heparinized LV blood (black bars), measured by reductive chemiluminescence (11), was lower than normal following 3 weeks of treatment with 10 mg/ml NAC (n = 3–4 each). In the same mice, plasma SNOAC levels (gray bars; measured by MS) increased from undetectable to approximately 2 μM over the same time (*P < 0.05). (B) Serum SNOAC, measured by MS, formed in NAC-treated mice (3 weeks). Left: liquid chromatogram; right: MS spectrum. NAC-treated mice had a SNOAC peak (m/z 193; red) coeluting with 15N-labeled SNOAC standard (m/z 194; black) that was absent in untreated animals (green) and was not detected in NAC-treated mice after serum pretreatment with HgCl₂ to displace NO• from the thiolate (blue). (C) Oxygenated erythrocytes were deoxygenated ex vivo (argon; ref. 11) in the presence of 100 μM NAC; supernatant SNOAC was measured by MS (above). SNOAC concentration increased with oxyhemoglobin (Oxy Hb) desaturation (co-oximetry: inset), being maximal at 59.3% saturation (blue), less at 77.2% saturation (green), and undetectable at 98% saturation. (D) SNOAC (filled circles) accumulated as the concentration of S-nitrosothiol–modified Hb (SNOHb; open circles) and oxyhemoglobin saturation (Hb SO₂; blue line) both decreased in heparinized whole blood using argon with 5% CO₂ (pH 7.3) in a tonometer. Both the increase in SNOAC and the loss of SNO_{rbc} between 0 and 20 minutes were significant (P < 0.01 by ANOVA followed by pairwise comparison to the maximum value; n = 3). *SNOAC levels were below the limit of detection when the oxyhemoglobin saturation was greater than 80%.
Chronic GSNO exposure in normoxia does not cause PAH. GSNO is an endogenous S-nitrosothiol that is similar to SNOAC in its kinetics of the homolytic decomposition (35). Normoxic CS7BL/6J 129SvEv mice receiving 1 mM GSNO in their drinking water for 3 weeks did not develop PAH as measured by any parameter (n = 3 in the GSNO group and 20 in the untreated control group; P = NS).

The molecular effects of S-nitrosothiols in vitro are hypoxia mimetic and recapitulate whole-lung effects of NAC in vivo. S-Nitrosothiols can have hypoxia-mimetic gene-regulatory effects involving the expression of transcription factors relevant to pulmonary vascular disease, including the hypoxia-inducible factor (HIF) (5, 13) and specificity protein (Sp) families (14). HIF 1α, HIF 2α, Sp1, and Sp3 are involved in regulating the expression of the genes upregulated in the lungs of mice treated chronically with NAC (9, 16, 17, 24, 25, 36). Because the expression and activity of the HIF and Sp families are affected by S-nitrosothiols in vitro (5, 13, 14, 37, 38), we tested whether chronic NAC or SNOAC could affect their pulmonary expression in mice. Interestingly, chronic NAC increased whole-lung Sp3 expression in vivo but not in primary pulmonary vascular endothelial cells in vitro (Figure 4, B and C). SNOAC, on the other hand, increased both intracellular S-nitrosothiol levels and nuclear Sp3 expression in primary murine pulmonary vascular endothelial cells (Figure 4, A and C). Thus, conversion of NAC to SNOAC (as shown in Figure 3) could be one explanation for the paradox that NAC can increase Sp3 expression in vivo but not in vitro.

Similarly, NAC increased HIF 1α activity in whole-lung extracts in vivo (Figure 5A), though NAC suppresses upregulation of HIF 1α, the oxygen-labile subunit of HIF 1, by both substance P (31) and by oxidized low-density lipoprotein (32) in vitro. As with Sp3, chronic NAC to SNOAC or HIF stabilizes HIF 1α expression, relative to β-actin, in primary pulmonary vascular endothelial cells in vitro (Figure 5B). This is consistent with previous work showing HIF 1α stabilization by GSNO (5, 13); indeed, we found that GSNO reduced HIF 1α monoubiquitination in bovine pulmonary arterial cells (BPAECs) expressing HA-tagged HIF 1α and a histidine-tagged, dominant-negative ubiquitin that stops ubiquitin chain propagation (K48R DN-Ub) (Figure 5C).

Mechanisms by which S-nitrosothiols inhibit HIF 1α ubiquitination and degradation are complex, involving both NO-radical reactions and transnitrosation reactions (1, 5, 13, 37–40). Low-micromolar SNOAC levels derived from NAC in vivo (Figure 3) would not generate concentrations of intravascular NO radical relevant to endothelial gene regulation (38) in the presence of millimolar concentrations of intravascular Hb (41); therefore, signaling through transnitrosation chemistry would seem more likely to be relevant to our in vivo model. NO transfer between thiols could stabilize HIF 1α through effects on prolyl hydroxylase, Akt signaling, and HIF 1α itself (13, 37–43). In addition, GSNO modifies E3 ubiquitin ligases through transnitrosation (44). Both SNOAC (Figure 5D) and GSNO (data not shown) inhibited the coimmunoprecipitation of HIF 1α with its E3 ligase, von Hippel–Lindau protein (pVHL) (45) in COS cells overexpressing both FLAG-tagged pVHL and HAtagged HIF 1α. Strikingly, SNOAC S-nitrosylated native pVHL in HeLa cells (Figure 5E), and SNOAC and GSNO increased pVHL

assayed by MS was undetectable). These findings are consistent with evidence that SNOAc distribution is, in part, dependent on oxyhemoglobin saturation (3, 11, 12) and that thiols accelerate the desaturation-induced loss of SNOAc content (11). SNOAC formation was pseudo-first order (in excess NAC) and relatively slow (k ~5.3 × 10−10 M/s).

We also considered that NAC could be converted to circulating SNOAC by reacting with NO in endothelial cells. S-Nitrosothiol concentrations in whole-cell lysates (43 ± 3.4 nM) were not increased by 4 hours’ exposure to 50 μM NAC in the presence of 5 μM exogenous NO (45 ± 3.1 nM; P = NS; n = 3). However, cellular levels were increased by exposure to 5 μM SNOAC (2.0 ± 0.4 μM; P < 0.05 when compared with NAC alone and control; Figure 4A).

eNOS-deficient mice are protected from the hypoxia-mimetic effects of NAC. eNOS is proposed to be important for maintaining SNOAc content (34). We studied NAC-induced PAH in eNOS−/− mice. The SNOAc content of these mice was 0.39 × 10−4 ± 0.15 × 10−4 (nearly a log order lower than in wild-type mice; P = 0.004). At baseline, eNOS−/− mice had slight increases in RV pressure and RV weight relative to those of the wild-type background (C57BL/6) mice, as measured previously (refs. 27–29; Figure 1, A and B). Strikingly, eNOS-deficient mice were completely protected from NAC-induced PAH — suggesting that the chronic effects of NAC to increase PAH are eNOS dependent — but they were not protected from SNOAC-induced PAH (Figure 1, A and B).

NAC could cause PAH in eNOS-replete mice by deleting endothelial NO or depleting eNOS-derived SNOAc (ref. 34; Figure 3). However, (a) NAC was not converted to SNOAC in eNOS-replete pulmonary endothelial cells in culture, even in the presence of exogenous NO; (b) SNOAc levels in eNOS−/− mice at baseline were lower than those in wild-type, NAC-treated mice; and (c) SNOAC caused, rather than ameliorating, PAH in eNOS−/− mice. Therefore, SNOAC excess, rather than endothelial NO or SNOAc depletion, appears most likely to have caused PAH in our model.

**Figure 4**
SNOAC recapitulates in primary pulmonary arterial endothelial cells the hypoxia-mimetic whole-lung effect of chronic NAC administration on Sp3 expression in vivo. (A) One micromolar SNOAC, but not 50 μM NAC, treatment (4 hours each) increased intracellular S-nitrosothiol levels (assayed by Cu/cysteine chemiluminescence; ref. 11) in primary murine pulmonary endothelial cells (P < 0.05 compared with SNOAC treatment). (B) Immunoblot showing increased Sp3 expression relative to MAPK in the whole-lung homogenates of mice treated for 3 weeks with 10 mg/ml NAC but not in those of control mice. By densitometry, this increase was significant (P < 0.01). (C) Paradoxically, however, NAC (50 μM; 4 hours) did not increase Sp3 expression relative to β-actin in primary murine pulmonary endothelial cells in vitro, while both SNOAC (1 μM; 4 hours) and hypoxia (10%; 4 hours) did.
S-nitrosothiols prevent normoxic ubiquitination and degradation of HIF 1α. (A) NAC treatment (10 mg/ml; 3 weeks) increased whole-lung HIF 1α-DNA binding activity. Complexes were supershifted (ss) with anti-HIF 1α and eliminated with excess cold probe (P). (B) SNOAC (1 μM), like GSNO (5, 13, 38), increased normoxic HIF 1α expression in nuclear extracts isolated from primary murine pulmonary endothelial cells. NAC alone (50 μM) did not affect HIF 1α expression. β-Actin was used as a protein load control. (C) In BPAECs transfected with HA-tagged HIF 1α and dominant-negative His-6-Myc–tagged ubiquitin (DN-Ub), ubiquitinated proteins were isolated using a nickel column and immunoblotted for HIF 1α. Both hypoxia and GSNO (G; 10 μM) inhibited HIF 1α ubiquitination relative to normoxia. (D) In COS cells cotransfected with HA-tagged HIF 1α and FLAG-tagged pVHL, GSNO (10 μM) did not affect HIF 1α expression. (E) S-nitrosylation of pVHL by SNOAC (5 μM) in equal protein aliquots isolated from HeLa cells was identified by biotin substitution (49); in the absence of ascorbate, S-nitrosylated pVHL was not detected. (F) Similarly, SNOAC and GSNO (5 μM) increased pVHL S-nitrosylation in pVHL-overexpressing 786-O cells. (G) C162, but not C77, was identified by biotin substitution to be S-nitrosylated in BPAECs transfected with wild-type cysteine 77 to serine mutant (C77S), C162S, or combined C77S/C162S pVHL exposed to SNOAC (1 μM). Native pVHL and MAPK immunoblots represented the pVHL expression and protein load controls, respectively. All in vitro treatments were for 4 hours.

Discussion

In vitro evidence has suggested that NO transfer reactions occurring during erythrocyte deoxygenation may signal hypoxia (3–5, 11, 12). However, the relevance of these reactions to hypoxia-associated processes at the whole-organism level has not previously been directly demonstrated. Here, we used the pharmaceutical antioxidant NAC as a bait reactant to study this biochemistry in vivo. We report that chronic NAC administration to normoxic mice can cause increased RV pressure, RV hypertrophy, and pulmonary vascular remodeling that are indistinguishable from the effects of chronic hypoxia.

**NAC must be converted to SNOAC to cause PAH.** Several observations suggest that the chronic pulmonary vascular effects of NAC require conversion of NAC to SNOAC. To begin with, chronic treatment with both NAC and SNOAC caused PAH (Figure 1). Thus, PAH must have resulted from 1 of 4 pathways: (a) NAC was converted to SNOAC to cause PAH; (b) SNOAC was converted to NAC to cause PAH; (c) both NAC and SNOAC caused PAH, each by a separate mechanism; or (d) the generation of a third metabolite from both NAC and SNOAC caused PAH.

Second, eNOS+/- mice had decreased circulating SNOAC content and were protected from NAC-induced PAH but were not protected from SNOAC-induced PAH. These data suggest that SNOAC formation from high concentrations of NAC and eNOS-derived NO, rather than reduction of lower concentrations of NAC and eNOS-derived NO, was necessary for the development of PAH. They suggest that neither NAC alone nor a common NAC/SNOAC metabolite causes PAH.

Third, we identified an oxyhemoglobin saturation–dependent mechanism by which NAC was converted to SNOAC under physiological conditions in blood. SNOAC concentrations, measured by MS, increased with the loss of SNOAC content during whole-blood deoxygenation in the presence of NAC ex vivo (Figure 3), consistent with previously reported NO transfer from deoxygenated erythrocytes to GSH to form GSNO (3). Neither SNOAC nor GSNO (3) were formed in the presence of oxygenated blood: deoxygenation was necessary. Alternative mechanisms by which NAC could be converted to SNOAC involve reactions in which NO radical either: (a) is oxidized in nonerythrocytic cells to an NO* equivalent—a slow reaction under physiological conditions—that, in turn, reacts with previously reported NO transfer from deoxygenated erythrocytes to GSH to form GSNO (3).
with NAC thiolate; or (b) reacts directly with NAC as a thyl radical. These mechanisms are not kinetically favored in the context of competing reactions with excess hemc or in the setting of hypoxia (1, 11, 40), and they were not observed experimentally in the current study: NAC had minimal reactivity with endogenous or exogenous endothelial NO. Taken together, these data suggest that erythrocyte deoxygenation resulted in conversion of NAC to SNOAC.

Fourth, NAC given systemically to the mice was converted to SNOAC in vivo, as identified by MS in RV (deoxygenated) blood. Levels of SNOAC were undetectable in LV (oxygenated) blood, suggesting that SNOAC was lost across the pulmonary vasculature and/or (consistent with our data) not formed from NAC in the presence of oxygenated erythrocytes. Further, chronic NAC administration depleted SNOAC content in vivo (Figure 3). Thus, the transnitrosation of an NO$^+$ equivalent from erythrocyte to NAC can explain SNOAC formation from NAC in vivo, consistent with previous observations (3, 11).

Note that these data, coupled with evidence that low levels of NO radical react minimally with thiols in isolated endothelial cells and in oxygenated erythrocytes, argue against a model in which NAC causes PAH simply by depleting endothelial NO. They also argue against a model in which SNOAC depletion causes NAC-associated PAH. Humans with hypoxia-associated PAH are deficient in S-nitrosothiol-modified Hb (34); the relevance of this deficiency to PAH in hypoxic humans is not contradicted by our data. Indeed, we show that eNOS$^{−/−}$ mice have a modest baseline increase in PA pressure associated with SNOAC depletion. However, chronic SNOAC treatment itself caused PAH in excess of that caused by SNOAC depletion.

Last, NAC-derived SNOAC is hypoxia-mimetic at the cellular level, whereas NAC itself opposes cellular effects associated with hypoxia (31, 32). SNOAC increases S-nitrosothiol levels in primary pulmonary vascular endothelial cells (Figure 4). S-nitrosothiols can upregulate hypoxia-associated genes in vitro through HIFs and/or Sp1’s (5, 13, 14, 38). In whole-lung homogenates, significant increases in mRNA and protein expression for genes associated with hypoxic pulmonary vascular remodeling were not uniform. This may reflect the use of whole-lung homogenates and the expression of genes in nonvascular cells. Nevertheless, mRNA and protein expression increased for several genes regulated by HIF 1, Sp3, and other hypoxia-associated factors. In the NAC-treated animals, it is unlikely that these effects on gene expression represented an effect of NAC alone: NAC (a) prevents hypoxia-mimetic cell injury in the absence of erythrocytes in vitro (31, 32); (b) did not cause PAH in the eNOS$^{−/−}$ mice (Figure 1); and (c) did not increase Sp3 or HIF 1$\alpha$ expression in vitro (Figures 4 and 5). Taken together, these data suggest that conversion of NAC to SNOAC was necessary for the hypoxia-mimetic effects of chronic, systemic NAC exposure.

Mechanisms by which S-nitrosothiols could be hypoxia-mimetic at the cellular level. SNOAC could interact with cellular proteins to cause pulmonary vascular remodeling through: (a) homolitic cleavage to form NO radical, which reacts downstream with cellular free radicals or heme groups according to diffusion-limited kinetics; or (b) transnitrosation reactions, in which NO is transferred to a target cysteine thiol, conventionally represented as a covalent NO$^+$ transfer between thiolate anions (1). The interactions of S-nitrosothiols with gene-regulatory proteins such as HIF 1$\alpha$ are complex and likely involve both NO radical chemistry and transnitrosation chemistry (38). In the context of hypoxia and/or oxidative stress in vitro, both NAC alone and S-nitrosothiol–derived NO radical can affect HIF 1 activation through mechanisms involving consumption of free radicals (38, 40). However, transnitrosation chemistry may be more relevant to our in vivo model because plasma S-nitrosothiols likely evolve little or no bioavailable NO radical in the presence of blood (1, 11, 41).

There are several potential targets of transnitrosation chemistry by which S-nitrosothiols might affect hypoxia-associated gene regulation. HIF 1$\alpha$ and prolyl hydroxylases could be modified by transnitrosation chemistry (37–39); and there may be targets in the Akt signaling cascade (13). Our results demonstrate that pVHL is also a target for S-nitrosylation. This posttranslational modification of pVHL appears to alter pVHL–HIF 1$\alpha$ interaction and decrease HIF 1$\alpha$ ubiquitination. Of pVHL’s 2 critical reactive cysteines, our data suggest that C162, which is necessary for the interaction of pVHL with elongin C within the HIF–E3 ligase complex (46, 47), is the likely target for S-nitrosylation. Additional work will be required to determine which mechanisms are the most important determinants of the hypoxia-mimetic cellular effects of SNOAC in our in vivo model.

Specific pulmonary effect of SNOAC. Oral administration of GSNO, an endogenous S-nitrosothiol (35) formed during erythrocyte deoxygenation (3), did not cause PAH, though GSNO has hypoxia-mimetic effects in vitro (5, 13, 14). This suggests that GSNO toxicity is prevented by mechanisms that may be bypassed by SNOAC. GGT cleaves GSNO to S-nitrosocysteinyl glycine and glutamate, serving as a gatekeeper in certain cells to regulate intracellular GSNO effects (3). SNOAC can bypass this regulation. However, pulmonary vascular endothelial cells express functional GGT (5). Therefore, we propose that the pulmonary vasculature is protected from oral GSNO, in part, by gastrointestinal GSNO metabolism (6). Additionally, NAC and SNOAC did not affect the kidney or liver; systemic vascular beds may be insensitive to SNOAC.

Alternatively, slow transnitrosation kinetics (Figure 3) may cause SNOAC to be formed only gradually as hypoxic systemic blood returns to the right heart, favoring an effect of SNOAC on the pulmonary vascular bed, where it is consumed before returning to the systemic circulation. However, mice fed SNOAC for 3 weeks at a dose that caused PAH had no change in portal histology. Therefore, we speculate that the pulmonary vascular bed is selectively sensitive to SNOAC, perhaps reflecting differential pulmonary expression of S-nitrosothiol metabolic enzymes (6). In this regard, the pulmonary vascular bed is susceptible to remodeling in the face of increased flow, chronic inflammation, and chronic hypoxia (7); each of these conditions could lead to increased S-nitrosothiol delivery to the lung. A fuller understanding of regional S-nitrosothiol metabolism, perhaps in animals in which long-term intra-arterial S-nitrosothiols can be delivered, will likely prove to be important.

The role of eNOS in this model of PAH. eNOS has been reported to have opposing roles in PAH (18–21, 26–29). An increase in pulmonary eNOS expression, described in certain species early in the course of PAH, may have a compensatory pulmonary vasodilator role. Consistent with this paradigm, we observed a transient increase in eNOS mRNA as well as a later increase in eNOS protein in PAH. On the other hand, the upregulation of eNOS protein could also represent a feed-forward mechanism by which SNOAC worsens pulmonary vascular remodeling, since eNOS$^{−/−}$ mice were protected from NAC-induced PAH.

The role of eNOS in chronic PAH certainly involves effects in addition to an eNOS-associated increase in S-nitrosothiol deliv-
ery to the pulmonary vascular bed: eNOS−/− mice have increased PA pressure at baseline, suggesting a role of eNOS on pulmonary vascular smooth muscle tone. These potential opposing roles for eNOS in regulating pulmonary vascular gene expression and smooth muscle tone might help to explain paradoxical observations regarding pulmonary vascular eNOS expression and activity in the setting of PAH.

Conclusions. Chronic exposure to NAC and to SNOAC causes murine PAH. We propose that these observations may be understood according to the following pathway. In normoxic, NAC-treated animals, oxyhemoglobin desaturation in erythrocytes leads to SNOAC formation in systemic blood returning to the lungs. These reactions require the presence of endogenous SNOec formed, in part, by eNOS. Formation of SNOAC — like that of GSNO, S-nitrosothiol-AE1, and other S-nitrosothiols during erythrocyte deoxygenation — is dependent on transnitrosation reactions. In these reactions, an NO− equivalent, protected from heme autocapture, is transferred from one thiolate moiety to another as Hb changes conformation during deoxygenation (1, 3, 4, 11, 12). The demonstration that SNOAC is formed during blood deoxygenation in vitro is complemented by the direct demonstration that it is present in hypoxic, but not normoxic, blood in vivo. While the cellular effects of GSNO and other S-nitrosothiols are regulated (1, 3, 5, 6, 14), delivery of NO to the pulmonary endothelium by SNOAC appears to bypass this regulation, causing hypoxia-mimetic effects. These observations may help to explain hypoxia-mimetic effects of systemic NAC therapy observed in humans in vivo (48).

The effect of chronic NAC exposure to cause pulmonary vascular pathology may be species dependent. It is reassuring that the NAC dose that caused PAH in mice was higher than the hypoxia-mimetic dose used in humans (48). However, PAH can be subclinical, even presenting as a terminal event (7, 8). In this context, it is noteworthy that NAC treatment may fail to decrease mortality in human clinical trials, despite beneficial antiinflammatory and antioxidant effects (30): PAH has not previously been considered as a NAC toxicity. Surveillance for this toxicity may be appropriate for long-term human trials with NAC.

In summary, these data: (a) provide the first in vivo measurements to our knowledge suggesting that NO can signal erythrocytic oxygen desaturation through S-nitrosothiol formation; (b) introduce a novel murine model of PAH, one in which transgenic animals are informative; (c) provide one explanation for previously reported, hypoxia-mimetic effects of NAC; and (d) suggest that PAH may be an unappreciated risk of systemic therapy with the pharmaceutical antiinflammatory and antioxidant agent NAC.

Table 1

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<tr>
<th>Gene</th>
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<tr>
<td>TNF-α (51)</td>
<td>Forward: 5′-GGTCCAGTGCATGATGAGACCCATAGA-3′</td>
</tr>
<tr>
<td>Reverse: 5′-CAGCTTCTACGTGGAGGCGATGTTGCGAC-3′</td>
<td></td>
</tr>
<tr>
<td>VEGF-A (52)</td>
<td>Forward: 5′-GCTCTCGTCACATGACAGGCAAGAC-3′</td>
</tr>
<tr>
<td>Reverse: 5′-GGTTCGACGTTACCTGTTGTA-3′</td>
<td></td>
</tr>
<tr>
<td>Glut-1 (53)</td>
<td>Forward: 5′-AAATAGGATGCGAGGGTGGTCT-3′</td>
</tr>
<tr>
<td>Reverse: 5′-AAGACAAGGCGAGGTGGGAAA-3′</td>
<td></td>
</tr>
<tr>
<td>eNOS</td>
<td>Forward: 5′-GCCAGGGGCAGGAAATAGTT-3′</td>
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</table>

Methods

Animal protocols. C57BL/6/129SvEv, C57BL/6, and eNOS−/− mice (10–12 weeks old) were treated in normoxia (21% O2) or hypoxia (10% O2, normobaric) as described (27) with or without 10 mg/ml NAC in the drinking water for 3 weeks (achieving serum NAC levels of 16.2 ± 4.3 μM as measured by liquid chromatography/MS). The plasma NAC level at this dose was similar to the target peak level in the human study of the hypoxia-mimetic effects of NAC (48), though the dose per weight was approximately 40-fold higher and the exposure continuous. Additional mice were treated for 3 weeks with 1 mM SNOAC in the drinking water (achieving venous plasma SNOAC levels of 350 ± 34 nM). The eNOS−/− mice used in these studies were on a C57BL/6 background, maintained by our group and backcrossed for 10 generations (27). Protocols were approved by the University of Virginia Animal Care and Use Committee.

RV pressure measurements. A Millar Mikro Tip catheter/transducer (1.4 F) was inserted through the external jugular vein of sedated (5 mg/ml pentobarbital) mice and threaded into the RV. RV pressure was determined from the average of 10–12 measurements (PulmoDyne; Hugo Sachs Elektronik) at a stable baseline (Figure 1C).

Immunohistochemistry, vessel morphology, and ventricular weight ratio measurements. Inflated lungs fixed in 10% formaldehyde were assessed for pulmonary vascular smooth muscle tone. These potential opposing roles for eNOS in regulating pulmonary vascular gene expression and smooth muscle tone might help to explain paradoxical observations regarding pulmonary vascular eNOS expression and activity in the setting of PAH.

Conclusions. Chronic exposure to NAC and to SNOAC causes murine PAH. We propose that these observations may be understood according to the following pathway. In normoxic, NAC-treated animals, oxyhemoglobin desaturation in erythrocytes leads to SNOAC formation in systemic blood returning to the lungs. These reactions require the presence of endogenous SNOec formed, in part, by eNOS. Formation of SNOAC — like that of GSNO, S-nitrosothiol-AE1, and other S-nitrosothiols during erythrocyte deoxygenation — is dependent on transnitrosation reactions. In these reactions, an NO− equivalent, protected from heme autocapture, is transferred from one thiolate moiety to another as Hb changes conformation during deoxygenation (1, 3, 4, 11, 12). The demonstration that SNOAC is formed during blood deoxygenation in vitro is complemented by the direct demonstration that it is present in hypoxic, but not normoxic, blood in vivo. While the cellular effects of GSNO and other S-nitrosothiols are regulated (1, 3, 5, 6, 14), delivery of NO to the pulmonary endothelium by SNOAC appears to bypass this regulation, causing hypoxia-mimetic effects. These observations may help to explain hypoxia-mimetic effects of systemic NAC therapy observed in humans in vivo (48).

The effect of chronic NAC exposure to cause pulmonary vascular pathology may be species dependent. It is reassuring that the NAC dose that caused PAH in mice was higher than the hypoxia-mimetic dose used in humans (48). However, PAH can be subclinical, even presenting as a terminal event (7, 8). In this context, it is noteworthy that NAC treatment may fail to decrease mortality in human clinical trials, despite beneficial antiinflammatory and antioxidant effects (30): PAH has not previously been considered as a NAC toxicity. Surveillance for this toxicity may be appropriate for long-term human trials with NAC.

In summary, these data: (a) provide the first in vivo measurements to our knowledge suggesting that NO can signal erythrocytic oxygen desaturation through S-nitrosothiol formation; (b) introduce a novel murine model of PAH, one in which transgenic animals are informative; (c) provide one explanation for previously reported, hypoxia-mimetic effects of NAC; and (d) suggest that PAH may be an unappreciated risk of systemic therapy with the pharmaceutical antiinflammatory and antioxidant agent NAC.

Table 1

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primers used for PCR experiments</th>
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</tbody>
</table>
Biotin substitution of S-nitrosothiol bonds. This was performed according to the method of Jaffrey et al. (49). EMDA. EMDAs were performed on nuclear extracts using oligonucleotides and antibodies (for supershift) as described previously (5).

Real-time PCR. Messenger RNA was isolated from whole-lung homogenates by an RNA Easy kit (QIAGEN) as per the manufacturer’s instructions. Expression of eNOS, VEGF, HIFM, fibroinectin, and Glut-1 mRNA was assessed by real-time PCR using the primers (Integrated DNA Technologies Inc.) noted in Table 1. The RT reaction was performed using 200 ng RNA under the following conditions: 25°C for 10 minutes, 48°C for 30 minutes, and 95°C for 5 minutes using the Reverse Transcriptase Kit (Applied Biosystems), as described by the manufacturer, in the presence of SYBR Green Supermix (Bio-Rad). PCR cycles for HMIF were: 95°C for 3 minutes, 95°C for 30 seconds, 63°C, 72°C for 30 seconds; for VEGF: 95°C for 3 minutes, 95°C for 30 seconds, 60°C for 1 minute, 72°C for 1 minute; for eNOS: 95°C for 3 minutes, 95°C for 30 seconds, 58°C for 1 minute, 72°C for 1 minute; for fibronectin: 95°C for 3 minutes, 95°C for 30 seconds, 55°C for 1 minute, 72°C for 1 minute; for Glut-1: the same conditions as for VEGF, each for 50 cycles. Relative gene expression was determined according to Livak and Schmittgen (50).

Commmunoprecipitation. COS cells were transiently transfected with HA–HIF-1α and FLG–pVHL using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions. Following S-nitrosothiol treatment, cells were lysed in lysis buffer (20 mM Tris pH 7.6, 150 mM NaCl, 10% glycerol, 2 mM EDTA 1% Triton X-100, 1 mM Na3VO4, 1 μg/ml leupeptin, 1 mM PMSF). Lysates were incubated with Flag M2 affinity gel (Sigma-Aldrich) (2 hours; 4°C). The gel was washed 4 times in lysis buffer. Samples were eluted with 50 μl 2× SDS buffer, heated (100°C; 3 minutes), and separated by 8% SDS-PAGE. HIF 1α was identified by immunoblot (5).

Nickel column isolation. BPAECs were transfected with HA–HIF-1α and pCW-8, a His-6 Myc–tagged ubiquitin containing the K48R mutation (gift from R. Kopito, Stanford University, Palo Alto, California, USA), using Lipofectamine 2000. Untransfected and transfected cells were grown in the absence or presence of S-nitrosothiol (21% O2) or in 10% O2. Cells were lysed in the absence of EDTA and lysates were passed over a ProBond Resin nickel column (Invitrogen) to isolate monoubiquitin-containing complexes. Complexes were eluted with imidazole and separated on 8% SDS-PAGE. HIF 1α protein was detected by immunoblot.

Site-directed mutagenesis of pVHL. pVHL was mutated using the Quick- Change Site-Directed Mutagenesis Kit (Stratagene) according to the manufacturer’s instructions using the following primers: pVHL C77S primer 1, 5′-CCCTCCGCAGGCTTCTCTTCTATGCAGTCGCCG-3′; primer 2, 5′-CGGACTGGGATTAGAAGAGATCAGGGGAGGG-3′; pVHL C162S primer 1, 5′-CTGAAAGGCAGATGCTCCAGTGGTCCGAGGC-3′; primer 2, 5′-GTCCTGGAACACTGGGAGGCTGCTTTACAG-3′. MS. Plasma (100 μl) spiked with 13N-labeled SNOAC was applied to a Phenomenex Strata XA 33 μm SPE cartridge (60 mg, 3 ml), preequilibrated (and washed following loading) with methanol/0.02% trifluoroacetic acid in water. Analytes were eluted with methanol and concentrated under reduced pressure (0°C). Partially purified samples were reconstructed in methanol, separated on a C8 column (5 mm, 2.1 × 150 mm) (Waters 2695 HPLC) with a gradient of 0.1% formic acid in water and methanol, and analyzed by electrospray ionization MS using a Finnigan LCQ system. (Finnigan Corp.) SNOAC cations were monitored at a mass to charge ratio of 192.5–193.5. Statistics. Multiple comparisons were made using ANOVA followed by pairwise analysis. Individual comparisons were made by Student’s t test if normally distributed; or otherwise by Mann-Whitney rank-sum test (using SigmaStat; Jandel). Relative gene expression by RT-PCR was quantitated as above (50). Except where noted, data are presented as mean ± SD. P < 0.05 was considered significant.

Acknowledgments

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Address correspondence to: Benjamin Gaston, Pediatric Respiratory Medicine University of Virginia Health System, Box 800386, Charlottesville, Virginia 22908, USA. Phone: (434) 924-1820; Fax: (434) 924-8388; E-mail: bmg3g@virginia.edu.