Hemoglobin conformation couples erythrocyte S-nitrosothiol content to $O_2$ gradients

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It is proposed that the bond between nitric oxide (NO) and the Hb thiol Cys-$\beta^3$ (SNOHb) is favored when hemoglobin (Hb) is in the relaxed (R, oxygenated) conformation, and that deoxygenation to tense (T) state destabilizes the SNOHb bond, allowing transfer of NO from Hb to form other (vasoactive) S-nitrosothiols (SNOs).

However, it has not previously been possible to measure SNOHb without extensive Hb preparation, altering its allostery and SNO density. Here, we have validated an assay for SNOHb that uses carbon monoxide (CO) and cuprous chloride (CuCl)-saturated Cys. This assay is specific for SNOs and sensitive to 2–5 pmol. Uniquely, it measures the total SNO content of unmodified erythrocytes (RBCs) (SNOHb), preserving Hb allostery. In room air, the ratio of SNOHb to Hb in intact RBCs is stable over time, but there is a logarithmic loss of SNOHb with oxyHb desaturation (slope, 0.043). This decay is accelerated by extraerythrocytic thiol (slope, 0.089; P < 0.001). SNOHb stability is uncoupled from O$_2$ tension when Hb is locked in the R state by CO pretreatment. Also, SNOHb is increased ~20-fold in human septic shock (P = 0.002) and the O$_2$-dependent vasoactivity of RBCs is affected profoundly by SNO content in a murine lung bioassay. These data demonstrate that SNO content and O$_2$ saturation are tightly coupled in intact RBCs and that this coupling is likely to be of pathophysiological significance.

Evidence has accumulated for an S-nitrosothiol (SNO)-based vascular signaling system in which hemoglobin (Hb) reactions with nitric oxide (NO) transduce redox gradients into bioactivities (1–6). There is agreement that human Hb undergoes S-nitrosylation at Cys-$\beta^3$ (3, 7–11). Erythrocytes are proposed to couple O$_2$ tension to the distribution of NO activities (such as control of blood flow) by linking the allosteric transition of Hb (12, 13) to conformational-dependent changes in the redox activity of this Cys-$\beta^3$ (13–18) and the stereochemistry of this SNO bond at Cys-$\beta^3$ (6, 7). Indeed, Cys-$\beta^3$ SNO in human Hb (SNOHb) can be crystallized only with the Hb tetramer in the relaxed (R, oxygenated) conformation; the SNO bond is unstable with Hb in the tense (T, deoxygenated) conformation (7). These observations support a paradigm in which NO binding to Cys-$\beta^3$ is favored in the R state and NO binding to Fe(II) (and/or transnitrosation to an alternate thiol) is favored in the T state (19–21). Thus, the change in stability of Cys-$\beta^3$ SNO during Hb transition between R and T states may serve to couple regional O$_2$ gradients to the deployment or quenching of NO bioactivities in the microcirculation (2, 6, 22).

However, assaying SNOHb has been problematic. First, detection of the SNO bond has required dilution and/or pretreatment of Hb to (i) control for artificial identification of nitrite and Fe-nitrosyl species and (ii) prevent autocapture of NO on Fe during analysis (8, 19, 23–25). As a result, attempts to quantify Cys-$\beta^3$ SNO density can be biased by shifts in Hb conformation during sample preparation, thus altering the intramolecular disposition of NO groups and subverting deductions regarding allostery. Also, the p50 of isolated Hb is significantly lower than that of intraerythrocytic Hb, making it technically difficult to desaturate and study purified, extraerythrocytic Hb under gas tensions that are relevant to physiology (1, 6, 19, 26). Moreover, several general issues make it challenging to study Hb–NO interactions, including (i) conformational polymorphisms of Hb; (ii) numerous allosterically interrelated heme–NO redox interactions at the $\alpha$ and $\beta$ hemes [with Fe(II), Fe(III), and Cys-$\beta^3$] (1, 2, 19, 27); and (iii) uncoupling of conformational transition and pO$_2$ because of weakening of the Fe–axial imidazole bond in the $\alpha$ chains by NO binding to $\alpha$ heme (particularly in the presence of inositol hexaphosphate) (28, 29).

Here, we describe a method for assaying the total SNO content of intact RBCs (SNO$_{RBC}$) with no pretreatment other than washing. This method involves NO reduction in CO. We first validated this method by using fluorescence- and colorimetric-based assays of isolated SNOHb. We then used the assay to show that, in intact RBCs, decreasing Hb O$_2$ saturation (Hb SO$_2$) is coupled to a decrease in RBC SNO content (SNO$_{RBC}$), whereas SNO$_{RBC}$ is unrelated to O$_2$ tension when Hb is locked in the R state by pretreatment with CO. We also show that SNO$_{RBC}$ is increased in human sepsis and that the O$_2$-dependent vasoactivity of RBCs varies with SNO$_{RBC}$. These data support the proposed role for allosterically governed SNOHb metabolism in the regulation of blood flow distribution in health (2, 3, 6) and suggest a role for RBCs in the pathogenesis of circulatory dysfunction in inflammatory states (9, 30). We propose that this assay will be valuable for (i) characterizing the biochemistry of SNO$_{RBC}$ signaling in physiology; (ii) monitoring SNO metabolism in therapeutic trials; and (iii) identifying abnormal interactions between NO and Hb in human disease processes.

Methods

Preparation of SNO-Loaded Hb. Hb (Hb$_{A_0}$) purified from human blood (Curacaye, Durham, NC) (31) was dialyzed overnight against 2% acetic acid (0.5 mM EDTA, pH 9.2). S-nitrosocysteine (CSNO; 0.5 M) was prepared immediately before use by reacting 1 M NaNO$_2$ in H$_2$O with 1.1 M L-Cys in 0.5 N HCl, 0.5 mM EDTA. Hb was S-nitrosylated by incubation (5–10 min at 25°C) with a 10-fold excess of CSNO (pH 7.4). The reaction was stopped on a Sephadex G-25 column. The total (Hb) and percentage of metHb were determined by the cyanomethemoglobin method (32). Preparations with >5% metHb were discarded. Samples were protected from light and stored at −80°C. The molar ratio of SNO to Hb for isolated Hb (referred to as SNO/Hb) was then measured as described below.
Preparation of SNO-Loaded RBCs. To avoid oxidative side reactions or S-nitrosylation of erythrocytic proteins other than Hb by CSNO, SNOHb was synthesized in intact RBCs by (i) addition of aqueous NO to fully deoxygenated RBCs to yield Fe-nitrosylHb [HbFe(II)NO]; (ii) washing under anaerobic conditions; and (iii) reoxygenation, effecting intracellular intramolecular transfer of NO from heme [Fe(II)NO] to Cys-γ93 (5). The specific preparation of SNO-labeled RBCs is detailed in the supporting information, which is published on the PNAS web site.

Colorimetric Assay. Sulfanilamide [SA; 3.4% (wt/vol)] in 0.4 M HCl was prepared with and without 1% (wt/vol) HgCl2, as was 0.1% (wt/vol) of N-(1-naphthyl)ethylene diamine (NED) (33). Equal volumes of SNOHb were added to SA with or without HgCl2 and then reacted with NED. [SNO] was determined from the difference in absorbance (540 nm).

4,5-Diaminofluorescein (DAF-2) Fluorescence Assay. An assay for biological SNOs using transfer of NO equivalents to DAF-2 (Calbiochem) to yield a fluorescent triazolofluorescein (DAF-2T; excitation and emission, 485 and 520 nm) (34) has been adapted for detection of SNOHb (35). Hb preparations in 10 mM PBS (pH 7.4), were incubated (10 min) with and without HgCl2 (at a constant molar ratio to Hb of 6.1, because HgCl2 alters DAF-2T fluorescence and must be held constant between wells) and treated with acid (0.4 N HCl, final). After exclusion of Hb (10-kDa filters spun at 10,600 × g for 20 min; Centricon), filtrates were mixed with 150 μM DAF-2 in 10 mM PBS (pH 7.4) in black microplates with the final titration to pH 8 (NaOH) to maximize DAF-2T fluorescence. Plates were read in an Fx-800 fluorimeter (Bio-Tek Instruments, Winooski, VT) and compared with synthetic DAF-2T (Calbiochem) and SNOHb standards (at 1:1 SNO/Hb).

Reduction in CO-Saturated Cu–Cys (CO/Cu/Cys): Chemiluminescence Assay. SNOs were selectively converted to NO in a cupric chloride (CuCl)-saturated, 1 mM Cys solution (pH 6.5) (36); NO was then detected by chemiluminescence (NOA 280, Sievers, Boulder, CO). In this assay, Cys has two functions. First, it forms CSNO by reaction with Cys to form Cys-SNO by reaction with Cu(I), forming NO [and Cu(II)]. The transnitrosation equilibrium favors CSNO because of the excess of Cys and rapid loss of product. Second, Cys reduces Cu(II), regenerating Cu(I). We modified this assay by adding carbon monoxide (CO) to the inert gas flow through the reflux chamber, preventing NO autoconversion by heme Fe(II). Metal carbonyls (∼0.7 ppm in research grade CO) must be removed, as both Ni- and Fe-carbonyls chemiluminescence in the presence of O2. Therefore, the CO source gas was passed through iodine crystals and (in series) activated charcoal (37) and then blended with the He stream in a gas proportioner (Aalborg, Orangeburg, NY). Note that (i) oxidized Cys should be replaced and residual Hb removed by refreshing the reflux chamber after each sample injection; and (ii) Nonidet P-40 and Triton X-100 both produce artifactual signal with this assay. This CuCl/Cys/CO (3C) assay was linear for SNOHb; the limit of detection was ∼2 pM for SNOHb, and specificity for SNOHb was nearly complete (see the supporting information), with 99–100% loss of signal after preincubation of sample with 10-fold excess HgCl2 (see Fig. 6 and supporting information). The coefficient of variation for sequential injections of 20 nM SNOHb [∼1/2 of the lowest (SNOHb) measured in human blood; see Fig. 4] was 0.055. Consistent with the measurements in ref. 36, the 3C technique did not detect nitrite, nitrate, or peroxynitrite in the presence of Hb. In all studies, signal in 3C attributed to SNOs was confirmed by loss after reaction with HgCl2.

3C Assay Validation Studies. We measured the effect of the ratio of HbFe(II) to SNO on assay sensitivity for low-mass SNOs as well as SNOHb. (i) Sensitivity for GSNO was determined in CuCl/Cys with and without CO and repeated after preinjection of SNO-depleted Hb into the reflux chamber. The starting SNO concentration of the depleted Hb was less than the limit of sensitivity of the colorimetric and DAF assays (that is, <500 nM; see supporting information); when diluted into the reaction vessel, the [SNO] was ∼25 nM and no signal was observed in the 3C assay. (ii) The assay linearity and the threshold for SNOHb detection were determined for synthetically SNO-loaded Hb (starting SNO/Hb ratios of 0.4, 0.2, and 0.06). These same dilutions were assayed by using both the colorimetric and the fluorescence-based assays. (iii) Sensitivity for SNO was measured while varying SNO/Hb while holding total [Hb] fixed at 100 μM; such stability is critical in the testing of biological samples. SNO-loaded Hb (SNO/Hb = 0.08) was diluted in Hb to yield samples with SNO/Hb ranging from 0.0006 to 0.01 and assayed for SNO content in 3C and DAF.

Correlation of SNO Content, O2 Tension, and Hb Conformation in Unaltered Erythrocytes. The following protocols were approved by the Human Investigation Committee of the University of Virginia, and written informed consent was obtained from all subjects. Washed arterial RBCs were prepared as above and resuspended (Hb, 1 mM) either with or without glutathione (GSH) (GSH/Hb ratio, 1:1,000), [Hb] pH, oxy-, deoxy- and met-Hb (%) were measured, and the suspension was then placed in a custom septated glass tonometer joined via a vacuum-tight, motorized couple (Buchler Instruments; Fort Lee, NJ) to source gas and a purge vent to permit continuous flushing and rotation of the chamber as well as sample extraction without atmospheric exposure. RBC O2 content was reduced steadily by flushing with argon. Aliquots were withdrawn by using gas-tight syringes (Hamilton) at 5-min intervals over 30 min for measurement of O2 tension, Hb cooxymetry, and total SNO_{RBC} (3C assay). In control experiments, O2-independent loss of SNO_{RBC} over time was studied by repeating the protocol without argon flushing. Also, Hb conformation was locked in the R state by using CO (until exclusively carboxyHb); deoxygenation and 3C assay were repeated.

Determination of RBC SNOHb Content in Human Sepsis and Lung Injury. Consensus criteria were used to identify patients with systemic inflammatory response syndrome (SIRS) (38) and acute respiratory distress syndrome (ARDS) (39). Mixed venous blood was drawn into gas-tight glass vials and protected from light (4°C). Samples were analyzed for pO2, pCO2, pH, total (Hb), oxyHb, deoxyHb, methHb (%), and SNOHb within 60 min. To prevent altering Hb conformation, samples were processed in an O2-controlled glove box with FiO2 set to 0.1% (by using 5% CO2 to raise the pCO2 to 500 nM; see supporting information). RBCs were washed as described above and protected from light before 3C assay for SNO_{RBC}.

Perfused Murine Lung (40) Modified for Bioassay of O2-Dependent SNOHb Vasoreactivity. This protocol was approved by the Animal Care and Use Committee of the University of Virginia in accordance with the Guide for the Care and Use of Laboratory Animals (41). Preparation of the isolated-perfused lung is detailed in the supporting information. Baseline pulmonary artery (PA) perfusion pressure (PAP) (∼8–10 cm H2O) was established at constant buffer flow during normoxic ventilation (21% FiO2/5% FiCO2/95% N2) and recorded for 10 min. Preparation viability was tested by establishing (i) exchange of CO2 and O2 across the lung, (ii) stable lung mechanics, and (iii) a ∼5% increase in PAP during a hypoxic ventilatory challenge (0% FiO2/5% FiCO2/95% FiN2). In viable preparations, normoxic ventilation was resumed and baseline PAP was reestablished for 10 min. Hb preparations were infused into the PA cannula at a rate to yield an intrapulmonary [Hb] of 100 μM of either SNO-depleted free Hb, SNO-loaded free Hb (SNO/Hb, 1:1; [SNOHb], 100 μM), or SNO-loaded RBCs (SNO/Hb, 0.1:1) in 10 mM PBS, pH 7.4. Total perfusate flow was kept constant and not recirculated. After initiation of the Hb infusion,
a PAP baseline was established over 5 min followed by hypoxic ventilation. The peak percentage increase in PAP from baseline during hypoxia was taken as a measure of the hypoxic pulmonary vasoconstriction (HPV) response. In additional controls, (i) the supernatant from the last wash of both SNO-depleted and SNO-loaded RBCs was infused (rather than the RBC suspensions) and HPV quantified; and (ii) the pressor response during normoxia to CO did not alter sensitivity for GSNO in controls were compared by the Mann–Whitney rank sum test. HPV quantified; and (ii) the pressor response during normoxia to CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb (Fig. 1). CO did not alter sensitivity for GSNO in the absence of Hb. GSNO, were accurate when compared with the sensitivity limit of the colorimetric and DAF2 assays, as confirmed by Bland–Altman analysis (see supporting information). [SNO] detected by 3C was linear over the range of [Hb] of 10 nM to 800 μM and SNO/Hb ratios of 10−6 to 1 (Fig. 2 and supporting information). As SNO/Hb fell below 0.1, Bland–Altman analysis revealed a minor bias between DAF and 3C: [SNO] detected by DAF was depressed in concert with an increase in the relative abundance of heme to SNO (see supporting information). We confirmed that there was a lack of such bias for 3C, detection was within the range of SNO/Hb ratios that have been reported for native human Hb samples (3, 8, 35, 43, 44) (Fig. 2), and the coefficient of variation for a [SNOHb] of 20 nM was 0.055.

Several points regarding Hb interactions with CO and Cu are worth noting. When saturated, the 3C solution (4 ml in the reflux chamber) contained ≈8.3 mM CO, in excess by a factor of 109 over the NO released from fully nitrosylated Hb (50 μl sample at 100 μM), and by a factor of 1012−14 over the [NO] at physiologic NO/Hb ratios. This ratio, and the fact that CO is carried continuously through the reaction mixture, may account for our complete SNO yield, despite the relatively greater affinity of heme for NO over CO (45). Note that the relative affinities of CO and NO for Hb under the conditions in the reflux chamber (where Hb tetramer instantly dissociates) are not known. Interestingly, an internal e− transfer pathway has been reported between Fe(II) in β-chain hemes and Cu bound to β3 sulfhydryls (15); also, Cu(II) binding at Cys-β3 accelerates reduction of β chain Fe(III) by CO by a factor of 105 (46). Together, these features may account for SNO signal fidelity in 3C.

GSNO Content Is Coupled to Hb Conformation in Intact Human RBCs. Suspensions of washed, fresh RBCs isolated from the arterial blood of healthy subjects were steadily deoxygenated with and without extracellular GSH (ratio of Hb to GSH = 1.000:1). The natural logarithm of the ratio of SNOHb to Hb was modeled as a function of HbSO2; the presence/absence of extracellular GSH was included as a covariate in the model. In both conditions, the ratio of SNOHb to Hb and Hb SO2 fell in tandem (R2 = 0.92), suggesting allosteric coupling. The SNOHb ratio fell more precipitously as the percentage of oxyHb crossed the range that is typically seen in arterio-venous traversal, yet halted short of full...
ventilation and vasopressor support. All patients had circulating pathogens [Escherichia coli (2), Enterobacter cloacae (1), Klebsiella pneumoniae (1), Staphylococcus aureus (1), and group A Streptococcus (1)]. None were receiving nitrovasodilators. In patients with SIRS and ARDS, the SNO_{RBC}/Hb ratio was 1.48 × 10^{-3} ± 7.9 × 10^{-4} (Hb SO_2, 60.2 ± 18.5%), 21-fold higher than normal: 7.07 × 10^{-5} ± 4.3 × 10^{-5} (Hb SO_2: 68.2 ± 7.2%) (n = 6; P = 0.002) (Fig. 4).

Sepsis and SIRS are characterized by inflammation and high levels of NO metabolites in general (nitrate, nitrite, and nitrotyrosine) (48, 49) and of SNOs in particular (30, 50, 51). Indeed, SNOHb is formed in vitro in RBCs exposed to inducible NOS (iNOS) (11) and SNOHb levels are high in rats exposed to LPS (50) or subjected to cecal ligation and puncture (9). Note that mice deficient in GSNO reductase have increased basal levels of RBC SNOs and have worsened tissue damage and mortality after LPS or bacterial challenge (30), suggesting a role for SNO metabolism in vascular instability associated with severe systemic inflammation.

**O_2-Dependent SNOHb Vasactivity in the Isolated Murine Lung.** O_2-dependent vasactivity of Hb (constriction in normoxia and dilation in hypoxia) is observed in systemic vascular ring preparations (1, 2, 6, 19, 20, 35), augmenting the intrinsic systemic vascular response to hypoxia. However, in the isolated lung preparation, putative RBC-based vasodilation (provoked by RBC traversal of a falling O_2 gradient) would counter the intrinsic pulmonary vascular response to hypoxia (HPV reflex). We hypothesized that SNOLoading of Hb and RBCs would augment O_2-dependent, RBC-based vasodilation and attenuate HPV amplitude in the isolated lung. Therefore, O_2-dependent dilation (quantified as a reduction in PAP during constant flow) can be wholly attributed to RBC-based NO bioactivity, as opposed to an intrinsic response. Also, the isolated lung model permits the creation of a steep O_2 gradient across the microcirculation (pO_2 fell from 500 to ~20 torr from PA to left atrium). Last, because the perfusate does not contact the cut edge of a vessel ring in this preparation, RBC-based bioactivity must act through the endothelium.

**Baseline Characteristics of the Preparation.** In nonrecirculating, constant flow conditions, baseline PAP during normoxic ventilation was 9.2 ± 0.4 cm H_2O (n = 23). Hypoxic challenge resulted in a 3.1 ± 0.4% increase in PAP (n = 23) (Fig. 5 A, B, D, and E). Baseline PAP (9.2 ± 0.5 cm H_2O; n = 23) was reestablished on

### Table 1. Human SNO_{RBC} values

<table>
<thead>
<tr>
<th>Condition</th>
<th>SNO_{RBC}/Hb</th>
<th>Whole-blood (SNO_{RBC})°</th>
<th>Hb SO_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sepsis</td>
<td>4.8 × 10^{-4} to 2.5 × 10^{-3}</td>
<td>(median, 1.3 × 10^{-4})</td>
<td>1.2–6.25 μM</td>
</tr>
<tr>
<td>Artery</td>
<td>1.0 × 10^{-4} to 2.2 × 10^{-4}</td>
<td>(median, 3.25 μM)</td>
<td>250–560 nM</td>
</tr>
<tr>
<td>Vein</td>
<td>1.9 × 10^{-4} to 1.3 × 10^{-4}</td>
<td>45–325 nM</td>
<td>68</td>
</tr>
<tr>
<td>Hypoxia (nadir)</td>
<td>1.08 × 10^{-5}</td>
<td>27.5 nM</td>
<td>34</td>
</tr>
</tbody>
</table>

°Estimated for 2.5 mM [Hb].
in baseline PAP (normoxia) on SNO loading of Hb (free or RBCs); (RBCs or free Hb attenuates the HPV response (hypoxic challenge; and (ii) during RBC perfusion and normoxic ventilation, SNO-loading did not alter the pressor response to U-46619 (100 nM) [SNO-depleted RBCs: 57% Δ cmH2O (n = 2); SNO-loaded RBCs: 62% Δ cmH2O (n = 1)].

Thus, SNO loading led to a Hb concentration-dependent change in vasoactivity. Specifically, it did not change vasoconstriction caused by perfusion with Hb or RBCs during normoxia, but it changed vasoactivity during perfusion with Hb and RBCs during hypoxia; SNOHb and SNO-RBC weakened the HPV response. SNO-RBC did not affect the normoxic pressor response to U-46619, nor did SNO-RBC supernatant affect HPV. Moreover, these data (after anaerobic exposure of RBCs to aqueous NO) highlight the exquisite responsiveness of RBC–NO reactions to O2 gradients, in that they complement data in which all allosteric control of NO delivery is absent after normoxic exposure of RBCs to excess CSNO (57), a SNO loading method that may lead to promiscuous transnitrosation of erythrocyte proteins. Indeed, vascular exposure to SNO-loaded RBC membranes results in vasodilation uncoupled from allosteric regulation by Hb O2 content (20); preexposure of RBCs to excess CSNO may mimic the pathophysiology of nitrosative stress in sepsis, confirmed here to increase SNO in vivo (Fig. 4), providing an explanation for the loss of vascular control in this condition.

Our measured ratios of SNO-RBC to Hb and estimated whole-blood (SNO-RBC) (Fig. 6 and Table 2) are consistent with reported values for human endogenous [(SNOHb) ranging 0.3–3 μM], as well as arterio-venous gradients and allosteric regulation of isolated SNOHb (3, 8, 35, 43, 44) made by some, but not all (52, 53), investigators. As reviewed in refs. 22, 54, and 55, differences among reports may relate to artifacts introduced during perfusion with Hb and RBCs during normoxic ventilation. The pressor response to hypoxia (HPV) was augmented during perfusion with SNO-depleted free Hb (Fig. 5 A and B, 10.4 ± 0.7 Δ cmH2O; n = 5), rather than for RBCs (Fig. 5 D and E, 6.5 ± 0.9 Δ cmH2O; n = 5; P < 0.05). The normoxic pressor effect was not altered by SNO-loading either free Hb (Fig. 5 A and B, 10.9 ± 1.1 Δ; n = 5; P value, not significant compared with SNO-depleted Hb) or RBCs (Fig. 5 D and E, 6.0 ± 1.2 Δ; n = 5; P value, not significant compared with SNO-depleted RBCs). SNO loading blunted the pressor effects of free Hb and RBCs during hypoxic ventilation. The pressor response to hypoxia (HPV) was augmented during perfusion with SNO-depleted free Hb (Fig. 5 A and B, 74.4 ± 11.1% Δ cmH2O; n = 5; P < 0.001 Δ cmH2O vs. hypoxia without Hb perfusion). This pressor response was less pronounced during perfusion with SNO-depleted RBCs (Fig. 5 D and E, 12.1 ± 3.2% Δ cmH2O; n = 5; P < 0.01 vs. hypoxia without RBC perfusion). SNO loading of either free Hb or RBCs attenuated HPV by ~50% (P < 0.05) (Fig. 5 C and F). In additional control studies, (i) perfusion with supernatant from the final wash of SNO-loaded RBCs, mixed into the buffer at the same rate as the RBC preparations, did not alter HPV (4.6 ± 0.8% Δ cmH2O; n = 2); and (ii) during RBC perfusion and normoxic ventilation, SNO-loading did not alter the pressor response to U-46619 (100 nM) [SNO-depleted RBCs: 57% Δ cmH2O (n = 2); SNO-loaded RBCs: 62% Δ cmH2O (n = 1)].

In patients meeting consensus criteria for SIRS and ARDS, the mean mixed venous SNOHb/Hb ratio was 1.48 ± 10−3 ± 7.9 ± 10−3, which is 21-fold higher than in normal volunteers (7.07 ± 4.3 ± 10−3; P = 0.002).

Fig. 4. In patients meeting consensus criteria for SIRS and ARDS, the mean mixed venous SNOHb/Hb ratio was 1.48 ± 10−3 ± 7.9 ± 10−3, which is 21-fold higher than in normal volunteers (7.07 ± 4.3 ± 10−3; P = 0.002).

Fig. 5. Bioassay for O2-dependent SNOHb vasoactivity in the isolated mouse lung. Representative PAP traces during perfusion with free Hb (A), free SNOHb (B), RBCs (D), and SNORBCs (E). (A) Experimental stages are identified including (a) PA cannulation and baseline during buffer perfusion and normoxic ventilation; (b) hypoxic challenge; (c) reestablishment of baseline after normoxic ventilation; (d) new baseline after addition of free Hb or RBCs to perfusate; (e) second hypoxic challenge; and (f) reestablishment of baseline after normoxic ventilation. Note (i) an increase in baseline PAP with Hb (free Hb > RBCs); (ii) no change in baseline PAP on SNO loading of Hb (free or RBCs); (iii) HPV amplitude during perfusion with Hb > RBCs (equimolar Hb); (iv) SNO loading of either RBCs or free Hb attenuates the HPV response (C and F). SNO loading causes an O2-dependent reversal of Hb and RBC vasoactivity, constriction in normoxia is maintained (baseline pressor response is unaltered), and dilation in hypoxia emerges (HPV is blunted with SNO loading).
during Hb isolation and pretreatment, which would not affect SNO content of intact RBCs. However, SNORBC decreases exponentially with decreasing Hb O2 saturation. These data confirm that SNORBC is allosterically regulated by O2 saturation, consistent with previous reports. Also, SNORBC is increased in human septic shock and is likely to be relevant to O2-dependent vascular control. We propose that the 3C assay and SNORBC deoxygenation slope constants will be applicable both to understanding pathophysiology and to monitoring therapeutic interventions in sepsis and other human disease states.

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Summary. The CO- and CuCl-saturated Cys (3C assay) is sensitive, does not detect HbFe(II)NO and has signal fidelity over an extensive range of SNO/Hb. It does not require sample pretreatment, permits SNO measurement in unaltered RBCs and eliminates the bias induced by shifting Hb conformation and/or the distribution of NO groups on Hb before assay. Using this method, we show that the SNO content of intact RBCs is stable over time and, when Hb is locked in R conformation by CO, over a broad range of PO2 levels. However, SNORBC decreases exponentially with decreasing Hb O2 saturation. These data confirm that SNORBC is allosterically regulated by O2 saturation, consistent with previous reports. Also, SNORBC is increased in human septic shock and is likely to be relevant to O2-dependent vascular control. We propose that the 3C assay and SNORBC deoxygenation slope constants will be applicable both to understanding pathophysiology and to monitoring therapeutic interventions in sepsis and other human disease states.