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Whole Body UVA Irradiation Lowers Systemic Blood Pressure by Release of Nitric Oxide From Intracutaneous Photolabile Nitric Oxide Derivates

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Rationale: Human skin contains photolabile nitric oxide derivates like nitrite and S-nitroso thiols, which after UVA irradiation, decompose and lead to the formation of vasoactive NO.

Objective: Here, we investigated whether whole body UVA irradiation influences the blood pressure of healthy volunteers because of cutaneous nonenzymatic NO formation.

Methods and Results: As detected by chemoluminescence detection or by electron paramagnetic resonance spectroscopy in vitro with human skin specimens, UVA illumination (25 J/cm²) significantly increased the intradermal levels of free NO. In addition, UVA enhanced dermal S-nitrosothiols 2.3-fold, and the subtraction of dermal S-nitrosoalbumin 2.9-fold. In vivo, in healthy volunteers creamed with a skin cream containing isotopically labeled 15N-nitrite, whole body UVA irradiation (20 J/cm²) induced significant levels of 15N-labeled S-nitrosothiols in the blood plasma of light exposed subjects, as detected by cavity leak out spectroscopy. Furthermore, whole body UVA irradiation caused a rapid, significant decrease, lasting up to 60 minutes, in systolic and diastolic blood pressure of healthy volunteers by 11±2% at 30 minutes after UVA exposure. The decrease in blood pressure strongly correlated (R²=0.74) with enhanced plasma concentration of nitrosated species, as detected by a chemiluminescence assay, with increased forearm blood flow (+26±7%), with increased flow mediated vasodilation of the brachial artery (+68±22%), and with decreased forearm vascular resistance (−28±7%).

Conclusions: UVA irradiation of human skin caused a significant drop in blood pressure even at moderate UVA doses. The effects were attributed to UVA induced release of NO from cutaneous photolabile NO derivates. (Circ Res. 2009;105:1031-1040.)

Key Words: nitric oxide ■ nitrite ■ nitroso compounds ■ UVA ■ decomposition ■ photolysis ■ human skin

A part from its effects on stroke, renal failure, and peripheral arterial disease, systemic arterial hypertension is a major risk factor for cardiovascular complications, including coronary artery disease, heart failure and sudden cardiac death.¹,²

Interestingly, mean systolic and diastolic pressures and the prevalence of hypertension vary throughout the world. Many data suggest a linear rise in blood pressure at increasing distances from the equator. Similarly, blood pressure is higher in winter than summer.³ Previously, it has been hypothesized that reduced epidermal vitamin D₃ photosynthesis associated with decreased UV light intensity at distances from the equator, alone or when coupled with decreased dietary calcium and vitamin D, may be associated with reduced vitamin D stores and increased parathyroid hormone secretion.⁴ These changes may stimulate growth of vascular smooth muscle and enhance its contractility by affecting intracellular calcium, adrenergic responsiveness, and/or endothelial function. Thus, UV light intensity and efficiency of epidermal vitamin D₃ photosynthesis may contribute to geographic and racial variability in blood pressure and the prevalence of hypertension.⁵

However, there might exist another or additional supporting mechanism, respectively, by which ambient electromagnetic radiation may affect blood pressure. Furchgott et al noted as long ago as 1961 that exposure to sun light relaxed isolated arterial preparations,⁶ although other types of smooth muscle tissue were much less sensitive.⁷ The vascular photorelaxation was wavelength-dependent, increasing as wavelength was reduced from the visible into the UV range, and it

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was independent of the endothelium.7 Furthermore, photorelaxation was markedly potentiated by solutions containing nitrite,8–10 indicating that under certain circumstances nitrite may exhibit relaxing activities comparable to NO.

Nitrite is a constituent of sweat, assumed to be formed on the skin surface by commensural bacteria.11 Furthermore, in human skin NOS-dependent production of nitric oxide (NO) potentially occurs in all dermal cell types.12,13 Some of the NO molecules formed remain at or close to the point of their origin as nitroso compounds, eg, S-nitrosothiols (RS-NO) or mercuric chloride–nonsensitive nitroso compounds or as the oxidation products nitrite and nitrate.14 UVA is known to penetrate deep enough into skin to reach the microvessels.15,16 Thus, in human skin photosensitive NO derivates like RS-NOs or nitrite may undergo photodecomposition when irradiated with UVA light,17–19 resulting in the formation of bioactive NO.14,20 Previously, we have demonstrated that UVA exposure of healthy skin specimens leads to an enzyme-independent high-output NO formation, reaching concentrations comparable or higher than found with maximal activity of the inducible NO synthase in cytokine-activated human keratinocyte cultures in vitro.21 We now extend these previous results by investigating the effect of whole body UVA exposure on the systemic blood circulation in humans.

Methods

Details regarding materials and experimental procedures with respect to materials, volunteers, UV sources, cell cultures, human skin samples, UVA-induced decomposition of nitrite and S-nitroso albumin formation, detection of S-nitroso proteins by immunohistochemistry, Western blot analysis of S-nitrosothiols proteins in human dermis, collection of blood samples and determination of blood pressure, cGMP measurements, analysis of cutaneous vascular parameters, sample preparation for detection of 15N-labeled nitroso compounds in human blood plasma by cavity leak out spectroscopy (CALOS), detection of NO, quantification of nitrite and nitroso compounds by chemoluminescence detection (CLD), electron paramagnetic resonance (EPR) spectroscopy, detection of 15NO by CALOS, and statistical analysis are in the expanded Methods section in the Online Data Supplement, available at http://circres.ahajournals.org.

UVA Irradiation of Human Skin Reduces Blood Pressure

Immediately after UVA irradiation, as well as up to 60 minutes after the light stimulus, the values of systolic as well as diastolic blood pressure were reduced in all subjects as compared to control values determined before the irradiation procedure. Figure 1 shows that mean arterial blood pressure (MAP) was significantly lowered after UVA illumination. The effect persisted for a considerable duration: relaxation toward previous resting state was observed on the timescale of about an hour (–5.6±3.2% immediately after UVA, –11.9±1.8% 15 minutes after UVA, and –5.9±2.1% 45 minutes after UVA); P<0.005 as compared to the controls).

UVA Irradiation of Human Skin Increases Plasma Nitroso Compounds and Nitrite Concentrations

The blood plasma of UVA-irradiated volunteers showed significantly enhanced nitroso compound (RX-NO) (Figure 2), as well as nitrite concentrations (Figure 3), in the time interval of 15 to 45 minutes after illumination (RX-NO: 74±16% 15 minutes after UVA and 53±19% 45 minutes after UVA; P<0.005 as compared to the controls; nitrite: 43±22% 15 minutes after UVA, 59±32% 45 minutes after UVA, and 40±26% 75 minutes after UVA; P<0.005 as compared to the controls). As shown in Figure 2D, UVA-induced decreases in blood pressure highly correlated with plasma RX-NO (R²=0.74) but did not correlate with plasma nitrite (R²=0.0071) concentration (Figure 3D).

UVA Irradiation of Human Skin Alters Cardiovascular Parameters

Furthermore, UVA-induced decrease in blood pressure was paralleled by increased forearm blood flow (FFB), increased flow-mediated vasodilatation of the brachial artery (FMDΔ%), as well as decreased forearm vascular resistance. As shown in Figure 4, 15 minutes after UVA, a significant increase in FBF (26.1±7.3%) and FMDΔ% (68±22%) and a significant decrease in forearm vascular resistance (–28.1±7.5%) was detected. UVA challenge had no significant effects on heart rates of irradiated volunteers.

Plasma From UVA-Irradiated Volunteers Exerts NO-Dependent Biological Activity

cGMP responses of RFL-6 cells in the presence of superoxide dismutase (500 U/mL) and isobutyl methylxanthine (0.6 mmol/L) were used to determine the bioactivity of plasma obtained from nonirradiated as well as UVA-irradiated volunteers. As shown in Figure 4H, incubation of RFL-6 cells with plasma that was collected from UVA-irradiated volunteers showed a significantly higher response in cGMP formation than plasma collected from nonirradiated volunteers (7.07±1.89 versus 2.65±0.63 nmol/L cGMP per milligram of protein). These increases were significantly lower in the presence of the NO scavenger 1H-imidazol-1-yl-oxy-2-(4-carboxyphenyl)-4,5-dihydro-4,4,5,5-tetramethyl-3-oxide (cPTIO) (1.95±1.23 nmol/L cGMP/mg protein).
Effects of Skin Temperature

During UVA irradiation, the ventral and lateral skin areas remained open to ambient air, and the skin temperatures of volunteers did not differ from controls (both 30.9 ± 1.3°C). The dorsal skin areas were not ventilated by ambient air. Here, the skin temperature of irradiated volunteers (37.9 ± 0.3°C) was slightly higher than that of controls (35.7 ± 0.8°C) (Figure 5A). To exclude a possible artifact from skin temperature on blood pressure, we investigated the effect of a 15 minutes bath in 38°C instead of UVA irradiation. The warm bath did not affect blood pressure at any time points up to 105 minutes post bath. (Figure 5B).

Additionally, we measured capillary–venous oxygen saturation, blood filling, blood flow, and flow velocity in superficial (1 mm deep) and deeper (6 mm deep) microvessels of human skin before, immediately after and 30 minutes after exposure to UVA (20 J/cm²) or 41°C warm water. As compared to nonirradiated skin, UVA exposure (20 J/cm²) had no effects on the mentioned cutaneous vascular parameters (Figure 5C and 5D). As positive control, exposure of human skin for 10 minutes to 41°C warm water significantly enhanced blood flow and blood velocity of superficial (1 mm deep) and deeper (6 mm deep) microvessels of human skin (Figure 5E and 5F).

Concentrations of Nitrite and S-Nitrosothiols in Skin Specimens and in Plasma of Volunteers

In parallel, immunohistological analysis of human skin specimens revealed consistently the ubiquitous presence of S-nitrosated proteins (Figure 6A), whereas UVA irradiation of skin specimens leads to a consistent strong increase in S-nitrosothiols (Figure 6B). In normal human dermis, S-nitroso thiols can be found at a concentration of 3.2 ± 0.9 μmol/L. The amount of S-nitroso thiols significantly increases after UVA challenge by 2.3-fold to 7.5 ± 1.2 μmol/L (Figure 6D). A similar UVA-induced 2.9-fold enhancement was found for S-nitrosoalbumin in skin specimens (Figure 6E). In the dermis of humans skin specimens incubated for 12 hours with 100 μmol/L nitrite, UVA irradiation leads to a 4.5-fold increase in S-nitrosoalbumin levels as compared to control specimens (Figure 6E).
To demonstrate UVA-dependent nonenzymatic NO formation from nitrite as well as UVA-induced nitrite-dependent S-nitroso-thiol formation in vitro, we irradiated nitrite-containing (100 μM) and/or BSA-containing (10 mg/mL) solutions (PBS, pH 7.4) with UVA and detected nonenzymatic NO formation by CLD and S-nitroso-BSA formation by Western blot. As shown in Figure 6F, at the physiological pH 7.4, UVA radiation led to an apparent nitrite decomposition and a significant formation of NO. Furthermore, in the presence of BSA this UVA-dependent nonenzymatic NO production from nitrite led to an significant increase in S-nitroso-BSA formation, as detected by the S-nitrosocysteine-specific antiserum (Figure 6G).

**Figure 2.** Effects of UVA irradiation of human skin on plasma concentrations of nitrosated compounds. Healthy volunteers (○, no. 1; ●, no. 2; ●, no. 3; ●, no. 4; △, no. 5; ▲, no. 6; □, no. 7) were irradiated for 15 minutes (gray area in A and B) with UVA light (20 J/cm²) or control-treated. Then, 15, 45, 75, and 105 minutes after irradiation concentrations of nitrosated compounds (RX-NO) of plasma were detected by CLD. A, Plasma RX-NO concentrations of UVA-irradiated volunteers. B, Plasma RX-NO concentrations of control-treated volunteers. C, Relative alterations in plasma RX-NO concentrations of irradiated (gray bars) and control volunteers (black bars) as compared to initial control values (0 minutes) indicated in A and B. Values are the means±SD of 7 individual experiments. D, Correlation blot between MAP and plasma RX-NO concentration. With each volunteer, the calculated values of relative alterations of MAP after UVA irradiation as indicated in Figure 1G were correlated to the calculated values of relative alterations of plasma RX-NO as indicated in 2C. The correlation coefficient (R²) is R²=0.7419. *P<0.001.

**Figure 3.** Effects of UVA irradiation of human skin on plasma nitrite concentrations. Healthy volunteers (○, no. 1; ●, no. 2; ●, no. 3; ●, no. 4; △, no. 5; ▲, no. 6; □, no. 7) were irradiated for 15 minutes (gray area in A and B) with UVA light (20 J/cm²) or control-treated. Then, 15, 45, 75, and 105 minutes after irradiation nitrite concentrations of plasma were detected by CLD. A, Plasma nitrite concentrations of UVA-irradiated volunteers. B, Plasma nitrite concentrations of control-treated volunteers. C, Relative alterations in plasma nitrite concentrations of irradiated (gray bars) and control volunteers (black bars) as compared to initial control values (0 minutes) indicated in A and B. Values are the means±SD of 7 individual experiments. D, Correlation blot between MAP and plasma nitrite concentration. With each volunteer, the calculated values of relative alterations of MAP after UVA irradiation, as indicated in Figure 1G, were correlated to the calculated values of relative alterations of plasma nitrite concentrations as indicated in 3C. The correlation coefficient (R²) is R²=0.0071. *P<0.001.

**Release of Gaseous NO From Intact Skin and NO Spin Trapping in Human Skin Specimens**

In a further experiment, an airtight chamber (16 cm²) with a UVA transparent front window was placed on the forearm of
volunteers. A gas flow of helium collected the gaseous NO emanating from the skin and was fed into the CLD analyzer (Figure 7A). In absence of UVA, a basal release of 29±25 fmol of NO per second per square centimeter was detected. Under UVA illumination with 20 J/cm², the release of gaseous NO was enhanced fourfold to 148±55 fmol of NO per second per square centimeter (P<0.001). After application of skin cream containing 10 mmol/L nitrite, the photoinduced yield of gaseous NO was again significantly enhanced to 334±112 fmol of NO per second per square centimeter after 1 hour of incubation with plasma obtained from nonirradiated (gray bars), as well as UVA-irradiated (20 J/cm²), volunteers (black bars, blood samples were collected 30 minutes after UVA challenge). White bars represent the constitutive cGMP production of RFL-6 cells alone. Additionally, incubations were performed in the presence of the NO scavenger cPTIO (40 μmol/L). Values represent the means±SD of 3 individual experiments. *P<0.001 as compared to the controls; #P<0.001 as compared to the respective samples incubated in the absence of cPTIO.

After illumination of Fe²⁺-DETC–loaded human skin specimens for 30 minutes with UVA light (25 J/cm²), small sections of 200 to 250 mg were cut, immersed in strong HEPES buffer and snap frozen in liquid nitrogen. Before EPR analysis, the skin samples were reduced with dithionite (50 mmol/L for 15 minutes) to remove EPR signals from Cu²⁺-DETC complexes. The EPR spectra of figure 7B at test formation of mononitrosyl-iron complex (MNIC) adducts (¹⁵N-NO-Fe²⁺-DETC, hyperfine triplet at g = 2.035) in human skin. Spectra of mamma skin specimens (Figure 8A through 8C) routinely showed additional signal from nitrosylated ferrous hemoglobin (paramagnetic NO-Fe²⁺-Hb) and ceruloplasmin. From comparison with calibrated reference samples, we estimated formation of 63±7 pmol MNIC in 200 mg male abdomen skin after 30 minutes UVA illumination. In absence of UVA, the MNIC yield remains below the EPR detection limit of ≈20 pmol. The MNIC yield could be enhanced to a massive 500 pmol by applying nitrite-loaded cream to the apical side of the skin specimens before UVA.

After splitting the skin samples horizontally with a razor blade, the apical outer layer had roughly threefold higher MNIC content than the endothermal inner layer. It shows that the outer layer is the main source of NO, as expected. Significantly, a large fraction of the total UVA induced NO has been trapped in the deeper skin layers, presumably because of diffusion of free NO through the skin tissue (The Fe-DETC traps and MNIC adducts themselves are immobilized in the lipid and protein compartments). After 30 minutes UVA, the MNIC concentration in the upper layers of male abdomen skin was ≈0.5±0.1 μmol/L. When the skin was pretreated with nitrite-spiked cream, the upper layers reached 6-fold higher MNIC concentration of ≈3.1±0.4 μmol/L.

These data suggest that NO is released from nitrite anions in the skin. Decomposition of nitrite was proven by application of cream with¹⁵N-nitrite (I=1/2) before UVA. The isotopic doublet structure of Figure 8 proved that the¹⁵NO ligand of MNIC derived from the¹⁵N-nitrite of the cream. After subtraction of an experimental¹⁵NO-Hb spectrum, we quantified the formation of 460 pmol¹⁵NO-Fe²⁺-DETC in 240 mg of mamma skin (Figure 8b).

Figure 4. UVA irradiation of human skin alters cardiovascular parameters. Healthy volunteers (○, no. 1; ●, no. 2; ◇, no. 3; □, no. 7) were irradiated for 15 minutes with UVA light (20 J/cm²). Prior and 15 minutes after (post) irradiation plasma RX-NO concentration (A), MAP (B), forearm blood flow (FFB) (C), forearm vascular resistance (FVR) (D), the diameter of the brachial artery (FMD%) (E), and heart rate (F) was detected in parallel. Values are the means±SD of 4 or 12 individual (in B and F) experiments, respectively. G, Relative alterations in plasma RX-NO concentration, MAP, forearm blood flow (FFB), forearm vascular resistance (FVR), the diameter of the brachial artery (FMD%), and heart rate of UVA-irradiated, as well as control-treated, volunteers as compared to initial control values (0 minutes). Values are the means±SD of 4 or 12 individual experiments, respectively. H, cGMP production of RFL-6 cells (3×10⁶ cells) after 1 hour of incubation with plasma obtained from nonirradiated (gray bars), as well as UVA-irradiated (20 J/cm²), volunteers (black bars, blood samples were collected 30 minutes after UVA challenge). White bars represent the constitutive cGMP production of RFL-6 cells alone. Additionally, incubations were performed in the presence of the NO scavenger cPTIO (40 μmol/L). Values represent the means±SD of 3 individual experiments. *P<0.001 as compared to the controls; #P<0.001 as compared to the respective samples incubated in the absence of cPTIO.
UVA Irradiation of Human Skin Induces Transmigration of Nitrite-Derived NO From Skin Tissue Into Plasma

Additional experiments were performed to identify the source of the NO moiety in the metabolites circulating in the blood of irradiated volunteers. Following application of skin cream with $^{15}$N-nitrite (20 mL containing 100 μmol Na$^{15}$NO$_2$, 5 mmol/L), whole body UVA irradiation (20 J/cm$^2$) led to the formation of significant quantities of plasma $^{15}$N-nitrite (40±4 nmol/L in controls versus 62±4 nmol/L in irradiated subjects, $P<$0.001) and $S$-nitrosothiols (RS-$^{15}$NO) (0.4±0.4 nmol/L in controls versus 1.7±0.9 nmol/L in irradiated subjects, $P<$0.001) (Figure 8E and 8D). These quantities were determined by the isotope-sensitive CALOS method. The fraction of labeled to unlabeled nitrite or nitroso compounds remained undetermined in this experiment.

Discussion

The key finding of the present study is that UVA irradiation of healthy human skin significantly increases intracutaneous NO and $S$-nitrosothiol concentrations via decomposition of cutaneous photolabile NO derivates with the result of significantly enhanced concentration of plasma nitroso compounds and a pronounced decrease in blood pressure.

Our observations of systemic UVA response can be plausibly explained by a mechanism comprising 3 elemental steps. First, UVA liberates NO from photolabile intracutaneous NO metabolites. Second, a fraction of the highly mobile NO diffuses toward the outer surface, where it escapes into the ambient atmosphere. (This fraction is detectable with the airtight skin chamber.) Another NO fraction diffuses to deeper tissue layers, where it enters the capillary vessels and enhances local levels of RS-NO. These nitrosated species may be low-molecular-weight, such as glutathione-$S$-NO, or protein-bound high-molecular-weight, such as albumin-$S$-NO. Third, the fairly stable nitroso compounds are distributed via the blood circulation, where it may elicit a systemic response like a drop in blood pressure. We note that the vasodilating and hypotensive properties of $S$-nitrosothiols are well documented. The observed release of free NO from UVA-irradiated skin lends strong support to this mechanism. Using isotopically labeled $^{15}$N-nitrite skin cream, CALOS spectroscopy demonstrated unequivocally that the photolysis of a photolabile NO derivate, here $^{15}$N-nitrite, in the epidermis by UVA contributes to the formation of nitrite and RS-NO species in the systemic blood circulation of volunteers. It provides proof of principle that NO moieties generated in the upper skin layers may migrate to the interior and translocate to NO moieties in the blood circulation for our proposed mechanism in vivo.

Human skin tissue is known to contain significant quantities of nitrite (4 to 6 μmol/L), RS-NO (~2.6 μmol/L) and mercuric chloride-resistant, as well as UVA-resistant, nitroso species (1.3 μmol/L). These concentrations exceed the human plasma concentrations by several orders of magnitude (nitrite ~20-fold, RS-NO ~300 fold). Every cell type in human skin is able to produce NO by at least one of three NO synthases. Therefore, enzymatically generated NO represents an important source of cutaneous photolabile NO derivates.

Nevertheless, recently data presented by Mowbray et al gave evidence that dietary nitrite and nitrate represent a more important source for cutaneous NO derivates. Because dietary nitrate increases circulating nitrite concentrations, it appears possible and feasible that dietary nitrate may also represent an effective way to boost skin reservoirs of photolabile NO species.

Using EPR spectroscopy, we, for the first time, give direct evidence here for UVA-induced intracutaneous NO formation via photodecomposition of endogenous sources of pho-
S-nitrosothiol formation in human skin specimens, as well as in vitro and of UVA-induced photodecomposition of nitrite in aqueous solutions. In resting, as well as UVA-irradiated (25 J/cm²), human skin specimens, obtained from mammoplasty surgery, S-nitrosation of proteins was detected by the S-nitrosocysteine-specific antiserum. A. Genuine human skin. B. UVA-irradiated skin specimens. C. For negative control, cryostat sections were denitrosated by a reducing solution (16 hours of incubation with 25 μmol/L CuCl₂ plus 1 mmol/L ascorbic acid in PBS, pH 7.4) before the antibody staining. A through C. Shown are representative pictures of 5 individual experiments. D. Detection of S-nitrosothiols in dermal tissue of genuine and UVA-irradiated human skin specimens detected by CLD in homogenates of genuine and UVA-irradiated human skin specimens. Values are the means±SD of 5 individual experiments. *P<0.001. E. Western blot analysis for S-nitroso-cysteine-specific antiserum.

S-nitroso protein formation in human dermis of genuine or UVA-irradiated (25 J/cm²) human skin specimens maintained in the presence or absence of NaN₃ (100 μmol/L). Shown is 1 representative graph of 3 individual experiments. F. In vitro nonenzymatic NO formation from UVA-irradiated (84 mW/cm²) nitrite-containing solutions (10 μmol/L sodium nitrite in PBS, pH 7.4) detected by CLD. G. UVA-induced nitrite-dependent S-nitroso-thiol formation in vitro. UVA irradiation (25 J/cm²) of PBS-containing (pH 7.4) nitrite (10 μmol/L sodium nitrite) and 10 mg/mL BSA resulted in an apparent S-nitroso-BSA formation, as detected by Western blot using a S-nitroso-cysteine-specific antiserum.

tolabile NO derivates. The action of NO is largely determined by its rapid diffusion and its ability to penetrate cell membranes. The diffusion coefficient of NO at 37°C has been found to be 1.4-fold higher than that of oxygen or carbon monoxide and thus a diffusion distance of 500 μm was calculated for NO in tissue.29 Thus, not surprisingly, with nitrite-enriched skin specimens, UVA-induced NO liberation could be found by EPR spectroscopy not only in apical skin regions but also in 2- to 3-mm deep regions of the dermis.

The penetration of photons into the skin strongly depends on the wavelength. It is known that UVA penetrates the epidermis and reaches even the deeper dermal regions down to 1 mm.16 Approximately half of the UVA intensity can reach the depth of melanocytes and the dermal compartment and it has been estimated that the total solar energy deposited into the lower epidermis and upper dermis is 2 orders of magnitude higher for UVA than for UVB. In vitro studies have shown that UVA light at 340 to 360 nm induces the formation of NO by photolysis of nitrite anions, as well as S-nitrosated compounds, in aqueous solutions.32–35 As shown by us previously,14 UVA-induced photodecomposition of nitrite results in a modest but sustained release of NO. In contrast, irradiation of RS-NOs leads to a much elevated release of NO because of the far higher extinction coefficient of this species. Under high-UVA intensities, the release of NO is short-lived because of rapid depletion of RS-NO (photobleaching). It should be noted that neither nitrite nor HgCl₂-resistant nitroso compounds, probably N-nitrosated species (RNNOs), contribute to UVA-provoked NO release from human skin.34 Detailed analysis of the mechanism of light-induced nitrite decomposition revealed the formation of very reactive and potentially cytotoxic radical species like O₂⁻, OH·, or NO₂⁻.17,32 The radical NO₂⁻ recombines rapidly (k ≈ 4.5×10⁴ mol/L per second) with NO to N₂O₃. N₂O₃ and the catalytic action of transition metal ions represent very efficient nitrosating systems, in particular for thiols.36,37 Via this reaction, NO₂⁻ decreases the yield of free NO from UVA-induced nitrite decomposition. In the presence of thiols such as glutathione, however, the NO-trapping capacity of NO₂⁻ will be counteracted via 3 reactions. First, N₂O₃ efficiently nitrosates thiols to RS-NO, which by itself is efficiently photolysed to NO and thyl radicals (‘S’) under illumination by UVA. Secondly, NO₂⁻ will directly be reduced to nitrite by thiolate like GS⁻. Thirdly, ‘S’ reacts efficiently with GSH to yield NO and a disulfide. In contrast, simple recombination of GS⁻ and NO’ has not been observed.39 Therefore, reaction of thiols with both NO₂⁻ and with N₂O₃ will increase the formation of NO. The reaction of thiolate anions with NO₂⁻ is ~10 times faster than the reaction with N₂O₃ (5×10⁸ versus 6×10⁷ mol/L per second) (reviewed elsewhere38,40).
Cutaneous formation of NO-Fe\textsuperscript{2+} bated for 30 minutes with 1 mmol/L DETC–loaded skin specimens from male abdomen were incubated for 30 minutes with 1 mmol/L DETC–loaded skin specimens from male abdomen were incubated for 30 minutes with UVA irradiation of human skin (NO\textsubscript{2} \textsuperscript{-} iminoethyl-L-ornithine in N\textsubscript{6} pmol of MNIC. In the presence of nitrite, UVA irradiation of human skin tissue (UVA) induces the appearance of the EPR-typical triple signal for NO and a MNIC signal attributable to UVA-induced, nonenzymatic NO formation were detectable (white bar). Under UVA illumination with 20 J/cm\textsuperscript{2}, the release of gaseous NO was enhanced 4-fold to 148 \pm 334 10 \textsuperscript{-10} pmol of NO per second per square centimeter (gray bar). After application of skin cream containing 10 \mu mol/L nitrite, the photoinduced yield of gaseous NO was again significantly enhanced to 334 \pm 112 fmol of NO per second per square centimeter (black bar). \textit{P}<0.001 as compared to the control (white bar). B, Fe\textsuperscript{2+}–DETC–loaded skin specimens from male abdomen were incubated for 30 minutes with 1 mmol/L N\textsubscript{6}–iminooethyl-L-ornithine in the absence or presence of nitrite (100 \mu mol/L NaNO\textsubscript{2}) and then were irradiated for 30 minutes with UVA light (25 J/cm\textsuperscript{2}). Intra-cutaneous formation of NO–Fe\textsuperscript{2+}–DETC–complexes (MNIC) attributable to UVA-induced, nonenzymatic NO formation were detected by EPR spectroscopy. EPR spectra at 77 K of human skin specimens in HEPES buffer. The specimens are \textbackslash ~200 \pm 10 mg each. In nonirradiated skin (control), MNIC signals are below the detection limit (bd) of \textasciitilde 20 pmol. This spectrum shows the presence of \textasciitilde 0.3 nmol of paramagnetic Cu\textsuperscript{2+}–DETC complexes. UVA irradiation of human skin tissue (UVA) induces the appearance of the EPR–typical triple signal for NO and a MNIC signal representing 63 \pm 200 pmol of MNIC. In the presence of nitrite, UVA irradiation of human skin (NO\textsubscript{2} \textsuperscript{-} +UVA) leads to a MNIC signal, corresponding to 500 \pm 50 pmol of MNIC.

In parallel to UVA-induced intracutaneous NO formation, we observed a strong increase in cutaneous S-nitrosothiol formation in the epidermis, as well as in the deeper regions of the dermis. As shown by Western blot analysis, the dermal fraction of S-nitrosated compounds predominantly represent S-nitroso-albumin, which, because of absent circulation activity in the skin specimens, reflect the blood or serum filling, respectively, of cutaneous microvasculature. In vivo, of course, because of the excellent capillarization of the Stratum papillare, synthesized dermal S-nitroso-albumin will imme-

diately leave the skin compartment. Functioning as a transport form for NO, S-nitroso-albumin will favor its rapid systemic distribution as well as its vasoadailability. S-nitroso-albumin has been previously proposed to act as a reservoir of NO within the circulation, transporting and releasing NO into vascular beds to cause vasodilation.\textsuperscript{41,42}

Photoproduction of NO has been observed previously at these wavelengths in vascular tissue of rats,\textsuperscript{43} and the action spectra of this photoproduction implicated endogenous S-nitrosothiols and nitrite as the source of NO. The UVA dose of 20 J/cm\textsuperscript{2}, as used here, was applied by using a commercial tanning facility. This dose remains significantly below the minimal erythimal UVA dose of 66 \pm 10 J/cm\textsuperscript{2} reported for fair-skinned persons\textsuperscript{43} and correlates with a sun exposure time of \textasciitilde 45 minutes in a temperate climate zone.

UVA-induced effects on cardiovascular parameters, as well as the timescale of alterations, are in reasonable agreement with previous observations. Recently, Rassaf et al demonstrated that intravenous slow infusion of NO in healthy volunteers increased plasma levels of RS-NO and induced systemic hemodynamic effects at the level of both conduit and resistance vessels, as reflected by dilator responses in the brachial artery and forearm microvasculature, and elicits a simultaneous and significant drop in mean blood pressure. Interestingly, slow infusion of NO had no significant effects on heart rates of the treated volunteers.\textsuperscript{44} These findings demonstrate that in humans, the pharmacological delivery of NO solutions results in the transport and delivery of NO as RS-NO along the vascular tree. Furthermore, in a pig model, Vilahur et al could show that low doses of S-nitroso glutathione (GS-NO), slowly administered, significantly reduced blood pressure.\textsuperscript{45} In accordance with our observations, in both studies, heart rates were not significantly affected, neither by an NO nor low-dose RS-NO injection. In this context, it should be noted that the systemic response of the vascular system depends on whether the given dose is administrated by bolus injection or gradually with slow infusion. Thus, in the same study by Rassaf et al, an intravenous bolus injection of higher GS-NO amounts led to significantly enhanced heart rates.\textsuperscript{44} Considering the time scale of UVA exposure, as well as of light-induced cardiovascular changes in our experimental setup, the underlying mechanism of our observations is less related to the high-dose GS-NO experiment of Rassaf et al but more to the mentioned NO and low dose RS-NO experiments.

As already mentioned, UVA radiation penetrates up to 1 mm into the skin. Therefore, hemodynamic changes shown here cannot be a direct result of cutaneous UVA exposure but rather are mediated by an UVA-induced factor. This assumption is strengthened by our observation that at the UVA doses used in our study, irradiation of skin did not show any significant local effects on cutaneous vasodilation or blood flow. Furthermore, we observed that an isolated irradiation of an arm, did not show any significant effects on blood pressure that was detected on this arm. On the other side, blood pressure detected on a nonirradiated arm of an otherwise UVA-irradiated volunteer shows the same results that were detected on the irradiated arm of the same volunteer (these
data are shown in the expanded Results section in the Online Data Supplement).

Furthermore, our control data strongly argue against an involvement of augmented ambient air temperature or skin temperature as an etiologic parameter for the effects on blood pressure observed after UVA challenge. In contrast to control-treated subjects, with UVA-irradiated volunteers, the permanent air stream exposure of ventral and lateral body parts, because of evaporation cooling, significantly decreases skin temperature. The surface of UVA-irradiated dorsal skin (not ventilated by cooling air), had a mean temperature of approx. 38°C. This is slightly higher than the skin temperature of fully covered subjects (35.7±0.8°C). Measuring capillary–venous oxygen saturation, blood filling, blood flow, and velocity of superficial (1 mm deep) and deeper (6 mm deep) microvessels of human skin clearly reveal that UVA exposure (20 J/cm²) had no effects on the mentioned cutaneous vascular parameters, whereas, as positive control, exposure of human skin for 10 minutes to 41°C warm water significantly enhanced blood flow and blood velocity of superficial, as well as deeper, cutaneous microvessels. Moreover, mimicking skin temperature increases by a full-body bath in 38°C warm water for 15 minutes, none of the volunteers showed significant alterations in blood pressure. Thus, the influence of skin temperature-dependent effect on blood pressure during UVA challenge can be neglected.

In conclusion, here we give evidence that whole body UVA irradiation NO-dependently decreases blood pressure of healthy volunteers. These systemic effects are correlated with increased concentrations of nitroso compounds in the systemic circulation. We attribute the observed effects to photolysis of cutaneous nitrite and show that the physiological response may be enhanced by loading the skin with photolabile NO derivates before irradiation. Alternatively, endogenous photosensitive NO derivates may be modulated by control over dietary nitrate and nitrite intake. These findings reveal the impact of light as an environmental parameter contributing to the phenomenon of “French paradox” and thus might have potential for the therapeutic applications in diseases with hypertension.

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Disclosures

None.

References

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Supplement Material

EXTENDED MATERIALS AND METHODS

Materials
If not other indicated all chemicals were from Sigma (Deisenhofen, Germany), the peroxidase-conjugated goat anti rabbit IgG antibody and the isotype-matched non-relevant rabbit anti-serum were from Calbiochem (Luzern, Switzerland). Gaseous NO was determined using the chemiluminescence detector (CLD 77 Amsp) from Eco Physics (Ann Arbor, MI, USA).

Volunteers
The protocol was approved by the ethics committee of the Medical Faculty of the Heinrich-Heine University of Düsseldorf and conducted in compliance with the Declaration of Helsinki Principles.

Fair skinned volunteers with skin type 2-3, 2 female, 8 male, age 38±11 years, body-mass index 26±4 kg/cm² were recruited from the student population and staff members.

UV sources
Irradiation of human skin specimens in vitro was performed using a 4000 W mercury arc lamp unit from Sellas Medizinische Geräte (Gevelsberg, Germany) emitting a UVA1-spectrum (340 – 410 nm) with a maximum of intensity at 366 nm (the lamp was used in a distance of 35 cm from the sample corresponding to a radiant flux intensity of 84 mW/cm²). The UVA dose in the in vitro experiments was 25 J/cm².

The IK ERGOLINE 44 sun-tube (Ergoline GmbH, Windhagen, Germany) was used for whole-body irradiation. This air-conditioned tanning device was fitted with 44 Solarium Plus R 100 W fluorescence lamps (Wolff System AG, Riegel, Germany) emitting UVA-light (99.3% of UV at wavelengths >320 nm and 84% >340 nm) with a
maximum intensity at 355 nm. The integrated irradiance at skin level was 19.5± 0.9 mW/cm² for UVA and 0.05± 0.01 mW/cm² UVB (means from 40 measure sites). The UVA dosage for whole body irradiation was 20 J/cm². This dosage was given in line with the manufacturer's recommendations and corresponds to the UVA contents of approx. 45 minutes sun exposure in a temperate climate zone. During UVA-exposure volunteers lying in horizontal position were irradiated from all sides. While the cooling air stream was able to cool head, neck, arms, legs as well as the ventral and lateral parts of the body, the dorsal body areas of pad contact were not ventilated. Volunteers only wore goggles that were opaque to UV and visible light. Control-treatment was performed by UVA-irradiation of dressed volunteers encased with UV-light-impermeable cloths. During irradiation a constant ambient temperature-interval within the sunbed-tube of 29±1 °C was achieved by manual adjustment of the integrated air-condition. Skin surface temperature was measured within 0.5 °C with a contact thermometer (Testo, T-stripe, Vienna, AU). Every volunteer contributed to dressed and undressed experiments (crossover experiment).

Cell cultures
Rat lung fibroblastoma cells (RFL-6) were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% bovine calf serum, 100 U/ml penicillin, and 100 µg/ml streptomycin. Cells were grown on 10 cm culture dishes and were used for experiments after reaching confluency.

Human skin samples
Human skin specimens were derived from mammoplastic or abdominoplastic surgery, cut into 10-mm squares, embedded immediately in Tissue-Tek (Reichert-Jung, Vienna, Austria), and snap frozen in liquid nitrogen for immunohistochemical
characterization. In order to examine S-nitrosoprotein concentrations in human dermis, skin specimens were incubated overnight (15 h) with Dispase (15 mg/ml; Boehringer, Mannheim, Germany) at 4°C and then epidermis was detached from the dermis.

The concentrations of cutaneous NO-derived products or reduced thiols were analyzed in the supernatants of skin homogenates. For this purpose embedded skin specimens (10-mm squares) were cryo-cut parallel to the epidermis into 20 µm thin slices down to a depth of 2 mm into the dermis (100 sections). The material was then weighed, diluted in 3 w/vol. of NEM-buffer (PBS containing 5 mM N-ethyl-maleimide (NEM), 2.5 mM EDTA, protease inhibitor), and homogenized. After a short centrifugation step the supernatants were collected, diluted to a protein content of 10 mg/ml, and immediately used or frozen at -20°C for maximally two weeks.

In addition, aliquots of fresh skin specimens (5-mm squares) were taken in short-time organ culture and were maintained in the RPMI1640/20% FCS culture medium (pH 7.4) in the dark up to 96 hours without loss of cell viability or function exactly as described by us previously.²

**UVA-induced decomposition of nitrite and S-nitroso albumin formation**

In order to demonstrate non-enzymatic NO-formation from aqueous nitrite solutions as a result of UVA-irradiation we irradiated PBS (pH 7.4) containing sodium nitrite (10 µM) with UVA (84 mW/cm² in a distance of 35 cm from the sample). UVA-irradiation of the solutions (in total 20 ml) was performed in a quartz glass cylinder (120-cm³ glass cylinder with 3.3-cm diameter), permanently exhausting of the gases for detection of NO by CLD exactly as described previously.³

Additionally, in order to demonstrate nitrite-dependent S-nitroso-thiol formation as a result of UVA-irradiation we irradiated PBS (pH 7.4) containing sodium nitrite (10 µM) and 10 mg/ml bovine serum albumin (BSA) with UVA (25 J/cm²). S-nitrosation of BSA was detected by Western-Blot and using a S-nitrosocysteine-specific rabbit anti-
serum exactly as described below.

Detection of S-nitroso proteins by immunohistochemistry

In resting as well as UVA-irradiated (25 J/cm²) human skin specimens S-nitrosation of proteins was examined using a S-nitrosocysteine-specific rabbit anti-serum. Skin specimens were embedded, snap frozen, and cryostat sections (7 µm of thickness) were prepared using a micro-cryotom exactly as described earlier. Cryostat sections were fix by glutaraldehyde (0.2% in TBS, pH 7.0) for 15 min at 4°C in a moist chamber, followed by inhibition of endogenous peroxidase activity with 0.3% H₂O₂ in ethanol, and washed three times in TBS. In order to stabilize S-nitrosothiols, transnitrosation was inhibited by alkylation of reduced thiols. Therefore, fixed sections were incubated for two hours in the dark with 10 mM NEM plus 0.3 % Triton X-100 in PBS. After blocking of unspecific binding with 0.5% BSA in TBS for 30 min and rinsing, specimens were incubated with the previously described rabbit anti-S-nitrosocysteine anti-serum used in a 1:100 dilution in TBS (supplemented with 3% low-fat milk powder and 0.5% Tween 20, pH 7.4). As secondary antibody peroxidase-conjugated goat anti rabbit IgG was used in a final dilution of 1:30 for one hour in TBS. All steps were performed at 4°C. For negative control S-nitrosothiols on cryostat sections were denitrosated by incubating the sections overnight (16 h) in the dark at 25°C in PBS containing 1 % CuSO₄ or 0.2 % HgCl₂ plus 0.5 % Sulfanilamid (diluted in 1N HCl) plus 2 % SDS. Then sections were washed using the washing buffer (0.1 mM diethylene triamine pentaacetic acid, DTPA plus 0.3 % Triton X-100 in PBS). After an additional washing in TBS nuclei were stained with hematoxylin for 1 min. Then, samples were incubated with 0.05% diaminobenzidine + 0.015% H₂O₂ for 5 minutes at room temperature. For light microscopy, cell samples were dehydrated, cleared with xylene and embedded in Eukitt.
Western-Blot-analysis of S-nitrosothiol proteins in human dermis

In order to examine S-nitrosoprotein formation in human dermis, skin specimens were irradiated by UVA (25 J/cm²) in the presence or absence of NaNO₂ (100 µM), incubated overnight (15 h) with Dispase (15 mg/ml; Boehringer, Mannheim, Germany) at 4°C. After detaching the epidermis homogenate solutions (10 mg protein/ml) of the dermis were prepared. In each lane of a 10%-Bis-Tris NuPAGE Novex pre-cast polyacrylamide gel (Invitrogen, Karlsruhe, Germany) 35 µl (35 µg protein) of skin homogenates were separated by electrophoresis using the NuPAGE electrophoresis system (Invitrogen, Karlsruhe, Germany) and the MOPS-SDS running buffer system under non-reducing conditions. Then protein was blotted on a nitrocellulose membrane using the NuPAGE transfer buffer (25 mM Bis-Tris, 25 mM Bicine, 1 mM EDTA, 20% methanol, pH 7.2) and following the manufactures instructions. Protein was visualized by using S-nitroso-specific anti-sera. Further incubations of the blots were: 2 hours with blocking buffer (2% BSA, 5% non fat milk powder, 0.1% Tween 20 in PBS-buffer), 30 min at room temperature with a 1:100 dilution of the anti-S-nitroso anti-serum, and 30 minutes with a 1:2000 dilution of the secondary horseradish peroxidase conjugated antibody. Finally, blots were incubated for 5 minutes in ECL reagent (Pierce, Rockford, IL, USA), and exposed to an autoradiographic film.

Collection of blood samples and determination of blood pressure

During the whole experimental procedure - from 45 minutes prior until 120 minutes post UVA-challenge - volunteers were kept in supine position. Then, 30 minutes prior to UVA-irradiation as well as 15, 45, 75 and 105 minutes after the light exposure venous blood samples (2 x 10 ml) were collected from the right cubatal vein. Systolic and diastolic blood pressure was determined by a cardiologist at the time points -30, -15, 0 minutes prior the irradiation, immediately after irradiation as well as 15, 45, 75, and 105 minutes after UVA-exposure by the method of Riva-Rocci using
a mercury sphygmomanometer or the OMRON M6 Automatic Digital Blood Pressure Monitor (OMRON Medizintechnik, Mannheim, Germany). Blood pressure, calculated as average of three successive measurements, always was determined at the right upper arm. Mean arterial pressure (MAP) was calculated as MAP \approx DP + 1/3 (SP-DP) where SP is systolic pressure and DP is diastolic pressure.

Changes in forearm blood flow (FBF) were measured at 10-second intervals using standard techniques of mercury-in-rubber strain-gauge plethysmography. The diameter of the brachial artery (FMDΔ%) was measured with a 15-MHz linear array transducer proximal above the antecubital fossa at end diastole by an automated analysis system. We calculated forearm vascular resistance (FVR) by dividing the mean arterial pressure by FBF.

cGMP-measurement

Guanosine 3',5'-cyclic monophosphate (cGMP) responses of RFL-6 cells in the presence of superoxide dismutase (500 U/ml) and isobutylmethylxanthine (0.6 mM) were used to determine the bioactivity of plasma obtained from non-irradiated as well as UVA-irradiated volunteers NO. RFL-6 monolayers (3x10⁵ cells) were covered with 1 ml plasma prepared from blood samples of healthy volunteers or from blood samples from UVA-irradiated volunteers collected 30 minutes after UVA (20 J/cm²) exposure in the presence or absence of the NO scavenger 1H-imidazol-1-yloxy-2-(4-carboxyphenyl)-4,5-dihydro-4,4,5,5-tetramethyl-3-oxide (cPTIO, 40 µM; Alexis Biochemicals, Grünberg, Germany). After 60 min of incubation cells were scratched and lysed by repeated freezing and thawing. cGMP levels in the supernatants were detected using the cGMP-specific ELISA (R&D-Systems, Wiesbaden, Germany) following the manufactures recommendation and were calculated as nM cGMP/mg protein.
Analysis of cutaneous vascular parameters

Effects of UVA radiation and warm water bath of cutaneous vascular parameters were performed using the O2C-device (LEA Medizintechnik GmbH, Giessen, Germany). O2C is a diagnostic device for non-invasive determination of perfusion of tissue, capillary-venous oxygen saturation, blood flow velocity, and blood filling of microvessels in microcirculation of blood perfused tissues. O2C is a multiple channel system which makes it possible to determine perfusion quantities and oxygen values by two channel. Channel 1 records the mentioned parameters in the superficial skin regions, while channel 2 monitors the different values of deeper skin tissue.

Sample preparation for detection of $^{15}$N-labeled nitroso compounds in human blood plasma by CALOS

In order to analyze the mechanism of the translocation of non-enzymatically produced NO from the skin into the circulation, 20 ml of a basis cream (standard oil-in-water-cream) containing $^{15}$N-labeled nitrite (5 mM Na$^{15}$NO$_2$, 98%$_{atom}^{15}$N) was evenly spread on the entire skin of healthy volunteers. The cream totally entered into skin leaving an essentially dry surface within 15 minutes. After a residence time of 45 minutes test persons were UVA-irradiated or control-treated exactly as mentioned above. Venous blood samples (20 ml) were collected 10 minutes prior to the irradiation procedure (control) and 15 minutes after finishing the irradiation. Immediately, after collection, blood samples were centrifuged (10 min at 800 x g, 4°C), the cellular fraction was discarded, and plasma was mixed with two volumes of cold (-20°C) acetone. Precipitated proteins were separated by centrifugation (10 min at 3200 x g, 4°C), protein pellets were washed twice with cold (-20°C) acetone, and resolved in PBS. In the deproteinized plasma fraction acetone was removed by overnight vacuum evaporation. Then protein-containing solutions as well as the deproteinized plasma were incubated for 60 min with nitrate reductase (0.15 U/ml in PBS containing 2.7 µM NADPH, 1.35 mM glucose-6-phosphate, 0.4 U/ml glucose-6-
phosphate dehydrogenase). In order to detect the grade of protein S-nitrosation in the protein fraction or nitrite in the deproteinized plasma, respectively, we used the iodine/iodide reduction system as described above, whereas the liberated $^{15}$NO was detected by cavity leak out spectroscopy (CALOS) as described in detail below.

**Detection of nitric oxide**

Three methods were used for detection of NO. In one series of experiments, the release of gaseous NO from skin and skin specimens was collected in an oxygen-free chamber and quantified using the chemiluminescence detector (CLD 77 Amsp) from Eco Physics (Ann Arbor, MI, USA). In a second series of experiments, in-vivo release of NO in the skin tissue was determined by in-vivo NO-spin trapping with iron-diethylthiocarbamate (Fe-DETC) complexes. Upon trapping of NO, these complexes form a paramagnetic mononitrosyl-iron complex (MNIC). The yield of MNIC in skin biopsies was quantified using electron paramagnetic resonance spectroscopy (EPR). In the third series $^{15}$NO was detected by cavity leak out spectroscopy (CALOS).

**Quantification of nitrite and nitroso compounds by CLD**

The concentrations of nitrite and nitrosated compounds (RX-NO) or S-nitrosothiols (RS-NO) in blood and dermal tissue were quantified by reductive denitrosation of plasma or of skin homogenate supernatant samples using a mixture of iodine/iodide in glacial acetic acid and subsequent detection of the liberated NO by its gas-phase chemiluminescence reaction with ozone, essentially as described. Nitrite concentrations were determined by the difference in peak areas of untreated aliquots and those subjected to preincubation with 0.5% sulfanilamide/HCl, the latter representing total nitrosated species. Discrimination between S-nitrosated molecules (RS-NO) and other nitroso species was achieved by preincubation of sample aliquots with mercuric chloride (HgCl$_2$; 0.2%), which selectively cleaves S-NO
bonds while preserving N-nitroso moieties, followed by sulfanilamide. The chemiluminescence detector was calibrated weekly using a 100 ppb mixture of NO in N₂ or He and calculated NO amounts were validated by injection of freshly prepared nitrite standards into the reaction mixture.

Alternatively, nitrite and RS-NO was determined in culture supernatants using the diazotisation reaction as modified by Wood et al. and NaNO₂ as standard.

**EPR spectroscopy**

Fresh skin specimens were kept in DMEM medium at pH 7.4 and 37°C under a controlled atmosphere containing 20 % O₂, 5 % CO₂ and 75 % N₂. Within 1 hr of surgery, the small 1cm x 1cm section of skin were loaded with Fe-DETC complexes for NO trapping. Such complexes are hydrophobic and were produced in situ in the low polarity compartments (lipid and protein) of the skin by two successive soaking steps taking a total of 1 hr. In the first, iron was loaded by soaking for 30 min in DMEM containing 150 µM iron-citrate. After rinsing with fresh DMEM to remove free iron-citrate, the skin was soaked for 30 min in DMEM containing 300 µM diethyldithiocarbamate (DETC) ligands. After ca 15 min exposure to DETC, the inner dermal side of the skin showed a noticeable dark-pink hue indicating formation of the strongly absorbing dark Fe³⁺-DETC complexes in the tissue. Visual inspection of a tissue cross section confirmed that the dark-pink hue was not restricted to the outer dermal surface but extended into the interior tissue. It was verified by visual and microscopic inspection that formation of black particulates (solid insoluble Fe³⁺-DETC crystals) did not occur in the medium. After loading with DETC ligands, the skin sections were again rinsed with fresh medium prior to the start of the actual trapping experiment. Placing the skin sections on ice terminated the trapping experiments. Small (ca. 200 – 250 mg) sections were cut, immersed in strong HEPES buffer (150 mM, pH 7.4) and snap frozen in liquid nitrogen until EPR assay. In order to discriminate NO formation in different skin thicknesses, specimens were split in
horizontal sections of 1.0 - 1.5 mm by a razor blade and sections were treated exactly as described above. In some experiments, enzymatic NO production was inhibited by administration of 1 mM L-NIO. In this case, the L-NIO was present during the 30 min incubation with DETC ligand, as well as during the actual trapping experiment. Alternatively, in some experiments the apical side of this skin section had been treated with cream containing 15N-Nitrite (5 mM) prior to UVA. After numerical subtraction of an experimental spectrum of 15NO-Hb, the difference spectrum was calculated.

The incubation times for the trapping experiments were calculated from the time point when loading with Fe-DETC was completed. We note that some formation of MNIC adducts may already occur during the incubation with DETC ligands since Fe-DETC traps are formed as soon as the iron-loaded skin is brought into contact with DETC. The quantity of these preformed MNIC adducts in Fe-DETC loaded skin sections remained below the EPR detection limit of ca 30 pmol MNIC, and was neglected. The EPR spectra were measured at 77 K on a modified X-band ESP300 spectrometer (Bruker BioSpin, Rheinstetten, Germany) operating with 20 mW microwave power. The skin specimens weighed ca 200 - 250 mg, and were immersed in strong HEPES buffer (150 mM, pH 7.4) and snap frozen in liquid nitrogen as small frozen columns of 4.8 mm diameter and a volume of ca 400 ± 20 µl. These columns were placed at the bottom of a quartz liquid finger dewar filled with liquid nitrogen. The sample was carefully centered inside a Bruker ER4103TM cylindrical cavity operating in TM110 mode with unloaded Q ~ 10.000. The magnetic field was modulated at a frequency of 100 kHz with 5 G amplitude. The detector gain was 2·10⁶, time constant 82 ms, and ADC conversion time 82 ms. Up to four field sweeps were accumulated to improve signal to noise. With these spectrometer settings, the detection limit was ca 30 pmol MNIC. The MNIC yields in the tissue samples were quantified by comparison with frozen reference samples of paramagnetic NO-Fe²⁺-MGD complexes (10 µM) in PBS buffer. The absolute
accuracy of the MNIC yields is better than 10%.

NO spin trapping is widely used in biological systems and formation of paramagnetic MNIC adducts is usually taken as evidence for free NO radicals. This is not strictly true, since S-nitrosothiols may also transfer the NO moiety to Fe-dithiocarbamate traps in a direct trans-nitrosylation reaction. However, the reaction rate is 6 orders of magnitude smaller than the trapping rate of free NO radicals, and makes this alternative pathway of MNIC formation negligible in biological systems.

**Detection of $^{15}$NO by cavity leak out spectroscopy**

The cavity leak out spectroscopy (CALOS) set-up has been developed mainly for trace gas analysis in atmospheric and medical applications. The method is based on laser absorption spectroscopy, which utilizes the fact that molecules absorb light at distinct frequencies. For NO, the strongest absorption features are located in the mid-infrared wavelength region near 5 $\mu$m. Due to well separated vibronic absorption lines even differentiation of different isotopologues of NO is possible. We used a CO sideband laser operating at 5.26 $\mu$m (1900 cm$^{-1}$) and 5.30 $\mu$m (1874 cm$^{-1}$) for $^{14}$NO and $^{15}$NO detection, respectively, providing a sideband power of about 120 $\mu$W. The laser light is coupled into a high finesse cavity consisting of two high reflective mirrors (R >99.99%), which is used as absorption cell. The transmitted laser power (2.5% of the incident power) is detected by a LN$_2$-cooled InSb photodector (3.5 A/W at 1875 cm$^{-1}$).

The resulting effective absorption path length (>5 km), which can roughly be estimated by dividing the cavity length (0.5 m) by the mirror transmission, allows the determination of extremely low absorption coefficients. The noise-equivalent absorption coefficient is $1.2 \times 10^{-10}$ cm$^{-1}$ at an integration time of 100 s, corresponding to 18 ppt $^{14}$NO and 16 ppt $^{15}$NO. Shorter integration times lead to slightly lower sensitivity. The time resolution is limited by the gas exchange time of the absorption cell which is less than 800 ms.
Statistical analysis

Values were reported as mean ± standard deviations (SD). For statistical analysis we used ANOVA followed by an appropriate post-hoc multiple comparison test (Tukey method). A p<0.05 was considered significant.
REFERENCES


