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NO synthase isoforms specifically modify peroxynitrite reactivity

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Nitric oxide synthases (NOSs) are multi-domain hemothiolate proteins that are the sole source of nitric oxide (NO) in mammals. NOSs can also be a source or a sink for peroxynitrite (PN), an oxidant that is suspected to be involved in numerous physiopathological processes. In a previous study, we showed that the oxygenase domain of the inducible NOS (iNOS_{oxy}) reacts with PN and changes its oxidative reactivity [Maréchal A, Mattioli TA, Stuehr DJ & Santolini J (2007) *J Biol Chem* **282**, 14101–14112]. Here we report a similar analysis on two other NOS isoforms, neuronal NOS (nNOS) and a bacterial NOS-like protein (bsNOS). All NOSs accelerated PN decomposition, with accumulation of a similar heme intermediate. The kinetics of PN decomposition and heme transitions were comparable among NOSs. However, their effects on PN reactivity differ greatly. All isoforms suppressed PN two-electron oxidative activity, but iNOS_{oxy} enhanced PN one-electron oxidation and nitration potencies, the oxygenase domain of nNOS (nNOS_{oxy}) affected them minimally, and bsNOS abolished all PN reactivities. This led to the loss of both NOS and PN decomposition activities for nNOS_{oxy} and iNOS_{oxy}, which may be linked to the reported alterations in their electronic absorption spectra. Bacterial bsNOS was affected to a lesser extent by reaction with PN. We propose that these differences in PN reactivity among NOSs might arise from subtle differences in their heme pockets, and could reflect the physiological specificity of each NOS isoform, ranging from oxidative stress amplification (iNOS) to detoxification (bsNOS).

Introduction

Nitric oxide synthases (NOSs) play an atypical role among the vast family of hemoproteins. Like cytochromes P450 and other hemothiolate proteins, their heme moiety performs highly selective and effective oxidation of substrates via the activation of dioxygen. However, the catalytic properties of NOSs are such that they lead to the biosynthesis of nitric oxide (NO) [1–3].

NOSs are homodimers comprising two distinct domains with specific catalytic roles. The reductase domain [4] is a flavoprotein with a ferredoxin-NADP-reductase motif that shuttles electrons [5,6] from NADPH via two co-factors (flavin adenine dinucleotide and flavin mononucleotide) to the oxygenase domain where the catalytic reaction takes place. The catalytic site comprises a heme B that is linked to the

Abbreviations

Arg, L-arginine; bsNOS, NOS-like protein isolated from *Bacillus subtilis*; CplI, compound II of the chloroperoxidase and catalase catalytic cycle; DHR, dihydrorhodamine; DTPA, diethylenetriaminepentaacetic acid; eNOS, endothelial nitric oxide synthase; H₄B, tetrahydrobiopterin, (6R)-5,6,7,8-tetrahydro-L-biopterin; HPA, 4-hydroxyphenylacetic acid; HPA-NO₂, 3-nitroHPA; di-HPA, HPA dimer; iNOS, inducible nitric oxide synthase; nNOS, neuronal nitric oxide synthase; NO, nitric oxide; NOS, nitric oxide synthase; NOS_{oxy}, oxygenase domain of NOS; PN, peroxynitrite; RNS, reactive nitrogen species.

protein backbone by a proximal cysteine ligand [7]. The NOS heme catalyzes the two-step oxidation of L-arginine (Arg) [8] using 6R-tetrahydrobiopterin (H_4B) as an additional electron donor [9]. The first step, presumably similar to the mono-oxygenation mechanism of cytochromes P450, converts Arg into N^{ω} -hydroxy-L-arginine [10]. The second step, specific to NOSs, converts N^{ω} -hydroxy-L-arginine into citrulline, with NO being formed as a by-product [11,12]. This subtly regulated mechanism [13] is shared by numerous NOS isoforms and by a number of known NOS-like proteins (e.g. bacterial NOS-like proteins [14]). Despite similar 3D structures and catalytic mechanisms [1,7,15–18], NOSs display different if not contrary physiological activities. Constitutive mammalian NOSs (endothelial and neuronal NOSs, eNOS and nNOS) are involved in cGMP signaling cascades, and are implicated in the regulation of vascular tone or in neuronal communication [19,20]. Inducible NOS (iNOS) is involved in cytotoxic activities, such as the non-specific immune response [21,22]. The roles of bacterial NOS-like proteins remain mostly unknown [23–26], but several pieces of evidence indicate that they do not share the same catalytic mechanism as mammalian NOSs [27,28].

The variety of biological roles among NOS isoforms may be the result of the rich chemistry of NO [29]. Indeed, generation of secondary products called reactive nitrogen species (RNS) has been suggested to explain the various physiological effects of NO. For instance, peroxynitrite (PN), a powerful oxidative and nitrative agent, is commonly associated with the cytotoxic and pathological activities of NOSs [30–33]. PN is believed to form in cells via the stoichiometric reaction of NO and superoxide anion ($O_2^{\circ-}$) [34,35]. However, NOS may also produce PN directly, either through ‘uncoupling’ conditions, in which both NO and $O_2^{\circ-}$ are formed [36–38], or via oxidation of the transient $Fe^{II}NO$ complex [39,40].

Once produced, PN may be activated by Lewis acids such as protons, metals or metalloproteins [41,42]. PN activation has been reported for several hemoproteins [43–45]. However, their effect on PN chemical reactivity varies from PN isomerisation and scavenging for globins [43] to increased nitration activity for hemothiolate proteins [46]. This apparent discrepancy remains unexplained despite several reports on the mechanism of PN activation [47,48].

In a previous study, we showed that the heme-containing oxygenase domain of inducible NO synthase (iNOSoxy) was able to activate PN, accelerating its decomposition rate and changing its chemical reactivity to promote one-electron oxidation and nitration

[49]. Here we report a similar analysis on two other NOS isoforms: the constitutive nNOS and a NOS-like protein from *Bacillus subtilis* (bsNOS). We analyzed their capacity to accelerate PN decomposition, the kinetics of NOS-induced PN activation, and the related heme transitions. We also investigated the effect of such activation on PN reactivity and NOS properties.

Results

All NOS isoforms activate PN decomposition

We studied the effects of the heme environment on PN decomposition kinetics by comparing three NOS isoforms (iNOSoxy, nNOSoxy and bsNOS) that exhibit different physico-chemical properties. NOS solutions were rapid-mixed with PN at room temperature, and PN decomposition was monitored by absorbance changes at 302 nm (see Experimental procedures). Figure 1A shows the PN apparent decay rates (k_{obs}) as a function of nNOSoxy and bsNOS concentrations. Data previously obtained for iNOSoxy [49] are superimposed for comparison. Both nNOSoxy and bsNOS accelerate PN decomposition, with second-order rate constants for PN activation (k_{act}) of $23.5 \pm 2.0 \times 10^4$ and $13.3 \pm 1.5 \times 10^4 \text{ M}^{-1}\cdot\text{s}^{-1}$, respectively (Table 1). As expected, no significant changes in PN spontaneous decomposition rates (y intercept) were noted as all data were obtained at the same pH value.

NOS isoforms share similar mechanism of PN decomposition

Superimposed absorption spectra showed that the kinetics of PN decomposition correlate with nNOSoxy and bsNOS heme spectral changes (Fig. S1B–D), as reported for iNOSoxy [49]. This confirms the involvement of NOS heme in the mechanism of PN decomposition for all NOSs. Analysis of the kinetic phases of PN activation by nNOSoxy at pH 6.4 (Fig. S1A,B) and by bsNOS at pH 7.4 (Fig. S1C,D) led to observations similar to those reported for iNOSoxy [49]. A series of transitions takes place during an initial phase (within a few milliseconds) that precedes the transient accumulation of a similar reaction intermediate (Fig. S1B–D) for all NOSs. The UV-visible absorption spectrum of the intermediate (Fig. 1B) is characterized by a Soret band between 442 and 447 nm and the presence of Q- β and $-\alpha$ absorption bands at approximately 550 and 590 nm. In the case of bsNOS, the Soret band is slightly shifted to the red (447 nm), and is broader (full width at half maximum of 52 nm) than

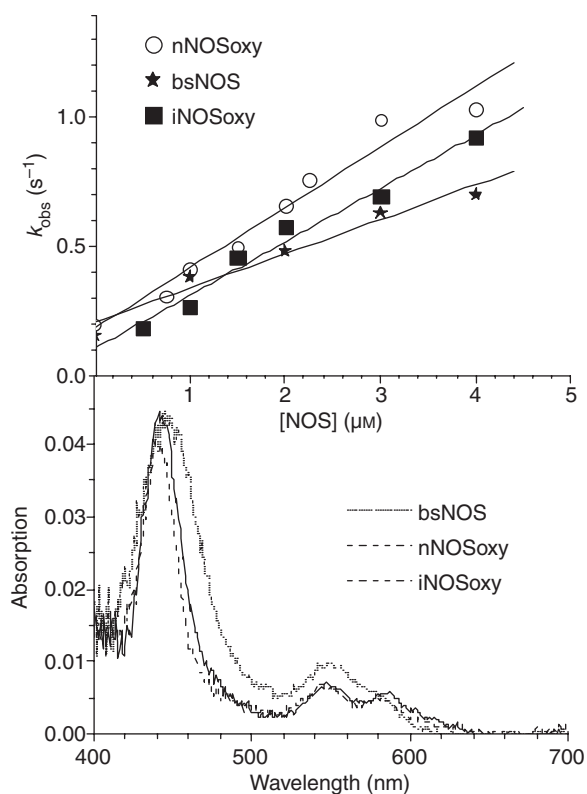


Fig. 1. Activation of peroxynitrite decomposition by the various NOSs. Top panel: 100 μM PN were rapid-mixed with increasing concentrations of nNOSoxy or bsNOS at pH 7.4. Conditions were as described in Experimental procedures. Kinetics of the 302 nm absorption decay were fitted to mono-exponential functions. The apparent decay rate constants (k_{obs}) are plotted as a function of the concentrations of nNOSoxy (open circles) and bsNOS (stars). Curves were fitted to a linear function to determine the rate constant for PN decomposition activation (k_{act} , slope) and PN spontaneous decomposition rate (k_{dec} , y intercept). Values are reported in Table 1. Data for iNOSoxy are from [49]. Lower panel: extracted spectra of the intermediates observed on reaction of 100 μM PN with 4 μM bsNOS at pH 7.4 (dotted line), nNOSoxy at pH 6.4 (dashed line) and iNOSoxy at pH 6.4 (solid line, data taken from [49]).

those observed for iNOSoxy and nNOSoxy (position 442 nm; full widths at half maximum of 36 and 30 nm, respectively). The accumulation and decay kinetics of this intermediate were fitted to a mono-exponential function for each isoform under the same concentration and pH conditions (data not shown). nNOSoxy and iNOSoxy intermediates showed similar formation and decay rates, and similar PN activation rates, whereas bsNOS exhibits two- to threefold slower formation and decay rates and a slower PN activation rate (Table 1). As observed for iNOSoxy [49], the last phase after PN consumption corresponds to the appearance of a modified ferric nNOSoxy species with

Table 1. Kinetic parameters for the reaction of PN with iNOSoxy, nNOSoxy and bsNOS. PN (100 μM) was rapid-mixed with increasing concentrations of NOS isoforms in gas-free 0.1 M KPi pH 7.4 buffer. Time traces of PN decay at 302 nm were fitted to a mono-exponential function. The observed rate of PN decomposition increases linearly with NOS concentration. The calculated slope represents the constant of activation of PN decomposition by NOS (k_{act}). Heme spectral changes were analyzed upon reaction of 100 μM PN with 4 μM NOS. Characteristic intermediates were observed at 443 nm (nNOSoxy and iNOSoxy) and 447 nm (bsNOS). Time traces at 443 and 447 nm were fitted to mono-exponential functions to obtain k_{build} (accumulation of intermediate) and k_{decay} (decay of intermediate).

	k_{act} ($10^4 \text{ M}^{-1}\text{s}^{-1}$)	k_{build} (s^{-1})	k_{decay} (s^{-1})
iNOSoxy	21.0 ± 2.0	14.8 ± 1.5	1.22 ± 0.04
nNOSoxy	23.5 ± 2.0	16.1 ± 4.1	1.02 ± 0.05
bsNOS	13.3 ± 1.5	6.0 ± 0.1	0.40 ± 0.01

absorption bands at 413 and 624 nm, distinct from native nNOSoxy, which is characterized by a Soret absorption band at approximately 417 nm and a Fe \rightarrow S charge transfer band at 650 nm (Fig. S1A,B). The bsNOS final spectrum is unaltered, with a Soret band at 404 nm and no absorbance change at approximately 624 nm (Fig. S1C,D). These results show that all NOSs bind PN at the heme and accelerate PN decomposition, with transient accumulation of a similar reaction intermediate. However, bsNOS does not show the same spectral modifications as mammalian NOSs upon PN activation.

NOS isoforms diversely modify PN oxidative properties

We investigated the effect of nNOSoxy and bsNOS activation on PN reactivity. The method was identical to that one previously reported for iNOSoxy [49] (see Experimental procedures). The two-electron oxidation reactivity of PN was assessed by comparing the production of rhodamine from dihydrorhodamine (DHR) in the absence and presence of NOS (Fig. 2A,B). The one-electron oxidation reactivity of PN was assessed by comparing the production of 4-hydroxy-phenylacetic acid dimers (di-HPA) from 4-hydroxy-phenylacetic acid (HPA) under the same conditions (Fig. 2C,D). As PN is known to be able to nitrate HPA to give HPA-NO₂, the same HPA probe was also used to characterize the nitrative power of PN in the absence and presence of NOS (Fig. 2E,F). All data were fitted to a linear function of PN concentration to quantify PN reactivity in the absence and presence of NOS isoforms. Figure 3 compares the effect of NOS isoforms on the oxidation

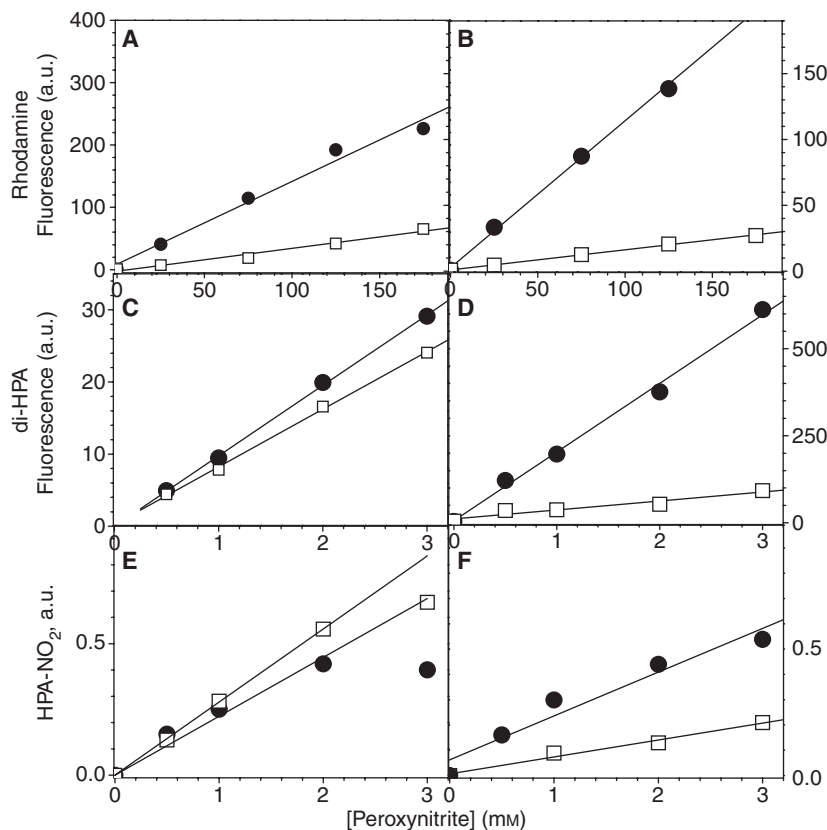


Fig. 2. Modification of peroxynitrite oxidative and nitrative properties by nNOSoxy and bsNOS. (A,B) DHR two-electron oxidation as a function of PN concentration. Fluorescence spectra of rhodamine ($\lambda_{\text{excitation}}$ 500 nm) were recorded after reaction with various concentrations of peroxynitrite (0, 25, 75, 125 and 175 μM). Fluorescence emission at 525 nm was plotted as a function of PN concentration in the absence of NOS (filled circles) or in the presence of 20 μM nNOSoxy (A, open squares) or bsNOS (B, open squares). (C,D) HPA dimerization as a function of PN concentration. Fluorescence spectra of di-HPA ($\lambda_{\text{excitation}}$ 326 nm) were recorded for various concentrations of peroxynitrite (0, 0.5, 1, 2 and 3 mM). Fluorescence emission at 405 nm was plotted as a function of PN concentration in the absence of NOSs (filled circles) or in the presence of 20 μM nNOSoxy (C, open squares) or bsNOS (D, open squares). (E,F). Analysis by reverse-phase HPLC of HPA-NO₂ production upon reaction of HPA with peroxynitrite. Conditions are described in Experimental procedures. HPA-NO₂ production was plotted as a function of PN concentration in the absence of NOS (filled circles) or in the presence of 20 μM nNOSoxy (E, open squares) and bsNOS (F, open squares). All curves were fitted to linear functions to compare the oxidative properties of free and NOS-activated peroxynitrite. a.u., arbitrary unit.

and nitration properties of PN. The bars indicate the percentage of reactivity when PN is activated by NOS compared to PN alone. Data obtained for iNOSoxy are included for comparison [49]. As previously reported, iNOSoxy quenches the PN two-electron oxidation capacity (-56%) but increases its one-electron oxidation and nitration properties ($+40\%$ and $+90\%$, respectively). Although nNOSoxy suppresses the PN two-electron oxidation ability (-72%), it does not drastically modify its one-electron oxidation and nitration properties (-18% and $+20\%$, respectively). On the contrary, bsNOS appears to suppress all chemical reactivities of PN by 86%, 83% and 70%, respectively (Fig. 3).

Effect of NOS-induced PN activation on NOS activity

We analyzed the effect of PN addition and NOS-induced activation on the NOS absorption spectrum. Ten to fifteen successive additions of four NOS equivalents of PN were mixed with NOS samples in the absence of substrate and co-factor (see Experimental procedures). UV-visible absorption spectra of the NOSs were recorded after each PN addition (Fig. 4). PN addition resulted in significant modification of the iNOSoxy absorption spectrum (Fig. 4C). A new absorption band appeared at 624 nm, together with flattening of the α/β region, resulting in disappearance

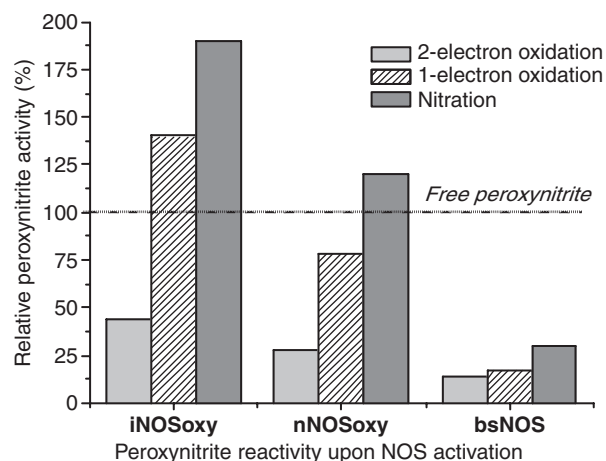


Fig. 3. Differential effects of NOS isoforms on PN-activated reactivity. Two-electron, one-electron and nitration properties of PN in the presence of bsNOS or nNOSoxy are derived from data in Fig. 4. Data for iNOSoxy are derived from [49]. PN reactivity in the presence of NOS was normalized to the control experiment performed in the absence of NOS (100% reactivity of free peroxynitrite).

of the 543 nm band. In parallel, the Soret band shifts from 420 to 411 nm upon PN addition, as observed by Huhmer *et al.* [50]. Although the first transition is complete after five successive additions, the Soret shift is not yet complete after ten additions, suggesting that the transitions might correspond to distinct structural effects. The same experiments were repeated with nNOSoxy and bsNOS. The nNOSoxy spectral changes match those observed for iNOSoxy, although to a lesser extent: the Soret band is blue-shifted (413 nm after 15 additions), and a similar absorption bands appears at 620 nm (Fig. 4A). The pattern is notably different for bsNOS. We did not detect any significant increase in the absorbance at 624 nm, and the Soret band is shifted from 401 to 408 nm (Fig. 4B). These results match the kinetic analysis of steady-state PN decomposition (above), which showed similar differences between bsNOS and mammalian NOSoxy final states.

Using the same samples, we analyzed the effects of PN activation on NOS catalytic properties. The protocols are similar to those used previously for iNOSoxy [49] (see Experimental procedures). We measured nitrite production by nNOSoxy (300 nM) and bsNOS (1 μ M) after each PN addition. NOS activity was normalized to NOS nitrite production in the absence of PN addition. Figure 5A shows that PN addition strongly inhibits nitrite production by nNOSoxy in a way similar to that previously reported for iNOSoxy (Fig. 5B, from [49]). No inhibition or decrease of the

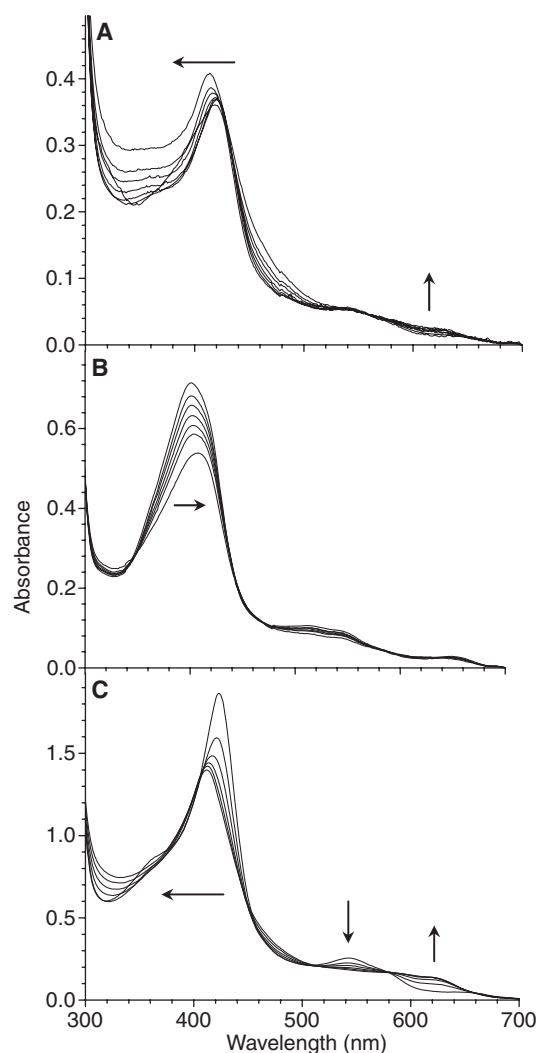


Fig. 4. Effect of activated peroxynitrite on NOS visible absorption spectrum. Superimposed UV-visible absorption spectra of the various NOSs after successive additions of PN. Successive additions (0–15) of four NOS equivalents of peroxynitrite were rapid-mixed with a 0.1 M KPi pH 7.4 solution containing nNOSoxy (6 μ M, A) bsNOS (10 μ M, B) or iNOSoxy (24 μ M, C). Horizontal arrows indicate the shift of the Soret band; vertical arrows indicate changes in the α/β band and the S \rightarrow Fe charge transfer band.

NO synthase activity was observed for bsNOS for up to 10 additions. We also characterized the effects of PN addition on NOS-induced PN activation. The kinetics of PN decay were monitored at 302 nm and fitted to a mono-exponential function after each successive PN addition (see Experimental procedures). Apparent decay rates were plotted as a function of the number of PN additions (Fig. 5). After several PN additions, nNOSoxy and bsNOS lose their ability to activate PN decomposition, as observed for iNOSoxy [49].

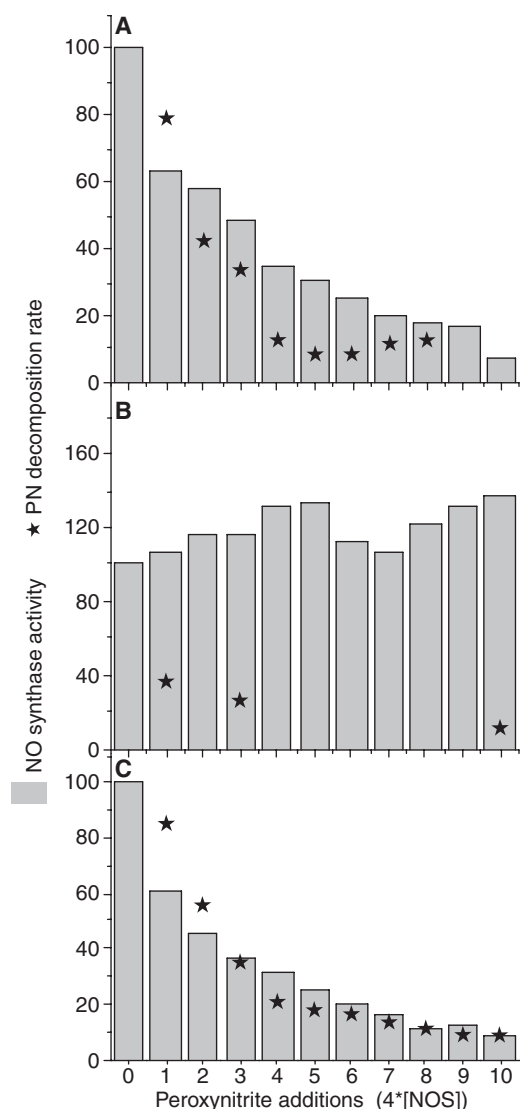


Fig. 5. Effects of activated peroxynitrite on NOS activities. (A) nNOS activity was measured after successive additions of PN (four NOS equivalents) using a standard Griess assay. Activities were normalized to control activity measured in the absence of PN, and are plotted as a function of the number of PN additions (histogram). The ability of nNOS to activate PN decomposition was measured after each addition of peroxynitrite. PN decay kinetics were fitted to a mono-exponential function. Apparent rates of PN decomposition were plotted as a function of the number of PN additions (stars). (B) As (A) but for bsNOS. (C) as (A) for iNOS; values are from [49].

Discussion

Fate and nature of the catalytic intermediate

All hemothiolate proteins (and proteins with a strong electron-donating proximal ligand) appear to catalyze the 'activation' of PN, i.e. the homolytic cleavage of

the PN O–O bond, to lead to the concomitant formation of a compound II (CpII) and a NO_2° radical [44]. This system has been suggested to account for the increased one-electron oxidation and nitration activities of PN, and we previously suggested that the 443 nm intermediate observed on reaction of PN with iNOSoxy was a CpII [49]. We report here that two other NOSs (nNOSoxy and bsNOS) also rapidly react with PN to form a similar 443 nm intermediate, but show different and at times contrasting oxidation properties upon reaction with PN. Compared to iNOSoxy, nNOSoxy does not significantly affect PN one-electron chemical reactivities, but bsNOS appears to suppress them. This is very intriguing, as the PN activation kinetics, the absorption spectrum of the reaction intermediate and its accumulation and decay rates are similar (Table 1 and Fig. S1), suggesting a common PN activation mechanism. There appears to be no correlation between accumulation of the 443 nm intermediate (common to all NOSs) and PN reactivity (different for each NOS). In fact, the differences observed in NOS/PN reactivity could occur downstream of CpII formation. Enhancement of the one-electron oxidation and nitration reactivities observed on PN activation by iNOSoxy could arise from a rapid escape of NO_2° from the heme pocket, as described for other hemoproteins [44,48,51]. In this context, the suppression of PN reactivity – as reported for bsNOS – could be explained by poor escape of NO_2° . This is reminiscent of the slow NO off-rate observed for bsNOS ($k_{\text{off}} = 0.23 \text{ s}^{-1}$, compared to 2.3 s^{-1} for iNOS), which has been attributed to the replacement of a valine residue by an isoleucine in the distal heme pocket of the bacterial homolog [28,52,53]. Thus, bsNOS is equally competent to produce the CpII– NO_2° pair but would promote its recombination with subsequent release of nitrate. This suggests that the product of the reaction between PN and NOSs actually depends on subtle differences in the heme pockets, which are reflected in their spectroscopic fingerprints [27,54,55].

Assignment of the 443 nm intermediate to a CpII for the reaction of NOS with PN is still awaiting experimental proof. It is supported by Newcomb *et al.*, who observed, by X-ray absorption spectroscopy, the formation of a CpII intermediate on reaction of PN with native cytochrome P450 CYP119 [56], but this has been challenged by Behan *et al.*, who suggested that the actual intermediate of the PN–P450 reaction could be a ferric-nitrosyl complex [57]. In addition, several reports have suggested that the reaction of PN activation might be too fast to permit characterisation of any reaction intermediates by conventional kinetic methods [58,59]. Consequently, the actual acti-

vation of PN by NOSs, leading to formation of the CpII-NO₂^o pair, might occur within the dead time of the reported kinetic analyses, and the 443 nm intermediate might result from side or end reactions forming other types of intermediates [51], such as the ferric-nitrosyl complex observed previously [57]. The series of successive transitions observed in the initial phase (first few milliseconds) might reflect several reactions occurring before accumulation of the 443 nm intermediate. Analysis of these extremely fast reactions, which is beyond the scope of this study, could allow identification of the very first reaction intermediates and characterisation of the molecular mechanism of PN activation.

Implications on NOS physiological specificities

This report highlights differences in heme-PN interactions between mammalian and bacterial NOSs. bsNOS appears to be the only hemothiolate protein that efficiently catalyzes the isomerisation of PN. The bsNOS active site appears to be specifically suited to rapid reaction with PN and its conversion into inert species, offering an effective means of PN detoxification. Moreover, unlike what has been observed for mammalian NOSs, the bsNOS structure is not dramatically altered by PN addition, and appears to be able to perform this detoxification task in the presence of a high concentration of PN. This protective role of bacterial NOS might also extend to other RNS and at first to NO [28,52]. Subtle differences in its heme pocket appear to allow bsNOS to trap and detoxify both NO and PN. This unique ability suggests another rationale for the physiological activity of bacterial NOS-like proteins that could be involved in defense of bacteria against the host immune response.

On the other hand, the iNOSoxy active site appears to promote PN one-electron oxidative and nitrative power, coinciding with its cytotoxic role, which is probably linked to production of highly reactive RNSs. iNOSoxy appears to catalyze its self-nitration and inhibition in the presence of large excess of PN, whereas activation of stoichiometric amounts of PN leaves iNOSoxy unaltered and functional. This may provide a direct way to control the extent of RNS production and therefore regulate the intensity of the cellular oxidative stress.

Our report highlights the remarkable ability of the NOS proteins to selectively adjust PN reactivity towards oxidation/nitration or isomerisation, which might be directly linked to their biological roles.

Experimental procedures

All chemicals were purchased from Sigma-Aldrich (St Louis, MO, USA) except dihydrorhodamine (DHR) and rhodamine, which were purchased from Calbiochem (San Diego, CA, USA), and PN, which was purchased from Cayman Chemical (Ann Arbor, MI, USA). PN solutions between 37 and 45 mM in 0.3 M NaOH were stored as 50 and 100 μ L aliquots at -80 °C. Before use, PN concentrations were measured by UV-visible spectroscopy ($\epsilon = 1670 \text{ M}^{-1}\text{cm}^{-1}$ at 302 nm). Argon gas was purchased from Messer France SA (Asnières, France).

Enzyme preparation

Wild-type iNOS and nNOS oxygenase domains (iNOSoxy and nNOSoxy) containing a six-histidine tag at their C-terminus were expressed in *Escherichia coli* BL21 using the PCWori vector as described previously [60–62]. The bacterial NOS-like protein from *Bacillus subtilis* (bsNOS) was over-expressed in *E. coli* BL21 as previously reported [63]. All proteins were purified in the absence of Arg and H₄B using Ni²⁺-nitrilotriacetate affinity chromatography as described previously [60,63]. NOS concentrations were determined by the absorbance at 444 nm of the heme ferrous OCO complex (Fe^{II}CO; $\epsilon_{444} = 74 \text{ mM}^{-1}\text{cm}^{-1}$ [64]).

Stopped-flow experiments

iNOSoxy, nNOSoxy and bsNOS were incubated in the absence of Arg and H₄B in a freshly anaerobized 100 mM KPi, 300 μ M diethylenetriaminepentaacetic acid (DTPA) buffer at the desired pH value. PN solutions were prepared just before use by dilution of a fresh stock solution in anaerobized 0.01 N NaOH. Enzyme solutions were rapidly mixed with the PN solution at 20 °C. The mixing sequence was defined to give determined concentrations of PN and NOSs. pH values were checked after mixing, and only negligible variations were noticed. Rapid-sampling stopped-flow experiments were performed on a Bio-Logic SFM-300 instrument (Bio-Logic Science Instruments SA, Claix, France) fiber-optically coupled to a Tidas spectrograph (WPI Inc., Sarasota, FL, USA) equipped with a rapid-sampling 1024 diode array spectrometer (3 ms per spectrum). The light source was a 150 W Xe lamp. The spectral range of the monochromator was 200–1015 nm, and the spectral resolution was 2 nm (0.8 nm per diode). The wavelength accuracy is quoted by the manufacturer as 0.1 nm. Seven hundred and fifty UV-visible absorption spectra were recorded at various time intervals to encompass the whole kinetics (ranging between 2.3 and 40 s). The apparatus dead time was estimated as < 2 ms. The above protocol was adapted to the kinetic analysis of PN decay and related heme spectral transitions.

PN decay kinetics

The PN decay kinetics were investigated for bsNOS and nNOSoxy using a protocol similar to that used for iNOSoxy [49]. PN decay was monitored by absorbance changes at 302 nm in the presence of increasing enzyme concentrations (from 0.5 to 4 μM). Three experiments were performed for each concentration. Time traces of the absorbance changes at 302 nm were multi-fitted to mono-exponential functions using ORIGIN 6.0 software (OriginLab Corporation, Northampton, MA, USA). This protocol was repeated three times. The derived apparent rate constants were averaged and plotted as a function of NOS concentration. The resulting data were fitted to a linear function to generate second-order rate constants.

Kinetic analysis of heme spectral transitions

PN (100 μM) was rapid-mixed with 4 μM NOS at three pH values (6.4, 7.4 and 8.4). Time traces were monitored at wavelengths characteristic of specific species such as PN (302 nm), ferric enzyme (401–423 nm for the Soret band and 624 nm for the modified final state) and the major catalytic intermediate (441–447 nm). Time traces at 445 nm were used to determine the time (t) of maximum accumulation of the intermediate, i.e. between 200 and 300 ms according to the experimental conditions. The UV-visible absorption spectrum of the intermediate was systematically calculated by subtracting an estimated fraction of the initial spectrum from the spectrum recorded at time t . This allowed us to estimate the Soret band for each NOS. Accumulation and decay rates of the intermediate were obtained from the mono-exponential fit of the time traces of absorption increase (from 0 to 250 ms) and decrease (from 400 ms to the end of the kinetics) at this specific wavelength. Reported values correspond to the mean of three independent experiments.

HPA fluorescence and HPLC assays

The reaction of HPA with PN leads to formation of HPA dimers (di-HPA) and nitrated HPA (HPA-NO₂), which can be isolated, identified and quantified by fluorescence spectroscopy and HPLC. The protocols used to analyze the effects of bsNOS and nNOSoxy on PN one-electron oxidation and nitration properties were identical to those previously described for iNOSoxy [49]. HPA (10 mM) was added to 100 μL of a freshly degassed reaction buffer (0.1 M KPi, pH 7.4, 0.3 mM DTPA) in the absence or presence of 20 μM NOS isoform. PN (0.5, 1, 2 or 4 mM final concentration) was added to the buffer, and the solution was immediately mixed vigorously. The solution was kept at room temperature for 10 min to ensure completion of the reaction, and was diluted to 1 mL for analysis. The production of di-HPA ($\lambda_{\text{excitation}}$ 326 nm, $\lambda_{\text{emission}}$

400/405 nm) was determined by fluorescence spectroscopy using Cary Eclipse fluorescence spectrometer (Varian Inc., Palo Alto, CA, USA). Emission intensity was plotted as a function of PN concentration. This series of experiments was repeated three times. Each plot was fitted to a linear function using ORIGIN 6.0 software (OriginLab Corporation), and the slopes of all curves were averaged. HPA-NO₂ production during the HPA reaction with PN was quantified by HPLC using the 365 nm absorption maximum. The areas of the HPA-NO₂ chromatogram peak at 365 nm were plotted as a function of PN concentration, and the resulting curves were fitted to linear functions. Experiments were repeated twice.

Dihydrorhodamine fluorescence assays

Dihydrorhodamine (DHR) oxidation into fluorescent rhodamine was used to assess the two-electron oxidative properties of PN. The protocol used to analyze the effects of bsNOS and nNOSoxy on PN two-electron oxidative properties was identical to that previously described [49] for iNOSoxy. PN at increasing final concentrations of 0, 10, 20, 30, 40 or 50 μM was mixed vigorously with a degassed 0.1 M KPi pH 7.4, 0.3 mM DTPA buffer, containing 40 μM DHR, in the presence or absence of the appropriate NOS isoform. Production of rhodamine was determined by fluorescence spectroscopy ($\lambda_{\text{excitation}}$ 500 nm, $\lambda_{\text{emission}}$ 525 nm), and plotted as a function of PN concentration for each isoform. The curves were fitted to a linear function, and the slopes of three series of experiments were averaged.

Analysis of the effects of PN addition on the UV-visible absorption spectra of NOS isoforms

NOS isoforms in solution in 0.1 M KPi pH 7.4 buffer containing 0.3 mM DTPA were mixed with PN aliquots (four equivalents of NOS concentration in degassed 0.01 N NaOH) in a quartz cuvette. The number of successive additions varied between 10 (iNOSoxy) and 15 (nNOSoxy and bsNOS). UV-visible absorption spectra were recorded 1 min after each addition. The same experiments were reproduced in the presence of 23.5 $\text{mg}\cdot\text{L}^{-1}$ catalase to prevent any side reaction due to the presence/formation of hydrogen peroxide in solution.

Analysis of the effects of PN addition on NOS activity

The protocol used to analyze the effects of PN addition on bsNOS and nNOSoxy activities was identical to that previously described for iNOSoxy [49]. NOS samples (300 nM nNOSoxy and 1 μM bsNOS) were rapid-mixed with up to ten successive additions of PN aliquots at four equivalents

of NOS concentration. All samples were then washed by three successive cycles of concentration/dilution in a 0.1 M KPi pH 7.4 buffer, using a Millipore (Billerica, MA, USA) membrane filter concentrator (30 kDa cut-off) to remove contaminants. Nitrite production of each protein sample was then measured using the Griess assay. Experiments were repeated three times. The kinetics of PN decomposition in the presence of 10 μ M NOS isoforms were recorded for the same protein samples using the protocol described above. Absorbance decays at 302 nm were fitted to a mono-exponential function.

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Supporting information

The following supplementary material is available:

Fig. S1. Kinetic phases observed during PN activation by nNOS_{oxy} and bsNOS.

This supplementary material can be found in the online version of this article.

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