



Differential calmodulin-modulatory and electron transfer properties of neuronal nitric oxide synthase mu compared to the alpha variant



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ABSTRACT

Neuronal nitric oxide synthase μ (nNOS μ) contains 34 additional residues in an autoregulatory element compared to nNOS α . Cytochrome c and flavin reductions in the absence of calmodulin (CaM) were faster in nNOS μ than nNOS α , while rates in the presence of CaM were smaller. The magnitude of stimulation by CaM is thus notably lower in nNOS μ . No difference in NO production was observed, while electron transfer between the FMN and heme moieties and formation of an inhibitory ferrous-nitrosyl complex were slower in nNOS μ . Thus, the insert affects electron transfer rates, modulation of electron flow by CaM, and heme–nitrosyl complex formation.

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Nitric oxide synthases (NOSs) are heme- and flavin-containing enzymes that catalyze the synthesis of nitric oxide (NO) from L-arginine (L-Arg). There are three isoforms encoded by different genes, neuronal (nNOS), endothelial (eNOS), and inducible (iNOS). NO has been implicated in neurotransmission, hemodynamic control, and the immune response, depending on the isoform and the cell type in which it is being expressed (for review, see [1–3]). All NOSs are functional dimers, with each monomer containing an N-terminal oxygenase domain followed by a reductase domain containing at least two regulatory elements – the autoregulatory region (AR), a ~40 residue sequence in the FMN-binding subdomain [4–7] and the C-terminal tail region (CT) [8,9]. These oxygenase and reductase domains are connected by a CaM-binding site, to which CaM must be bound for electron transfer into the heme. Roman and Masters [10] proposed a model of NOS regulation in

which the AR, in combination with the CT, stabilizes either a closed or an open form of the enzyme, depending on the binding of NADP⁺/NADPH or CaM, which allows electron transfer between the FAD and FMN or between the FMN and the heme, respectively.

All NOS isoforms are expressed to some extent in mammalian skeletal muscle. In addition to nNOS α , skeletal muscle contains an alternatively spliced nNOS isoform, nNOS μ , which contains 34 additional amino acids within the AR [11]. The physiological role of either variant of nNOS in skeletal muscle is unclear, but in nNOS knockout (KN1) mice, skeletal muscle bulk and tetanic force production were decreased, and muscles exhibited increased susceptibility to contraction-induced fatigue [12,13], implicating nNOS as having a role in the maintenance and function of muscle.

Despite the 34-residue difference, initial reports indicated that nNOS μ and nNOS α shared similar spectral properties, substrate and CaM binding values, and NO synthesis activity [11,14]. In the present work, we probed deeper into biochemical differences between these two proteins, demonstrating that the insertion affects CaM modulation of electron transfer of nNOS μ relative to nNOS α , and also alters the formation of the heme nitrosyl adduct formed during NO synthesis. Moreover, detailed FMN–heme interdomain electron transfer (IET) kinetics studies suggested differences in

Abbreviations: NO, nitric oxide; NOS, nitric oxide synthase; nNOS, neuronal NOS; CaM, calmodulin; IET, interdomain electron transfer; L-Arg, L-arginine; H₄B, tetrahydrobiopterin; AR, autoregulatory region; CT, C-terminal tail region

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the FMN domain conformational equilibrium between the two proteins.

1. Materials and methods

1.1. Plasmids

The plasmid rat nNOS α pCW was as described [15]; that of nNOS μ pCW was made using the same protocol and oligonucleotides, with rat nNOS μ cDNA in pCDNA3 as template, kindly provided by David Bredt of Johnson and Johnson (San Diego). The plasmid encoding calmodulin, CaM ACMIP, was kindly provided by Anthony Persechini of the University of Missouri (Kansas City).

1.2. Protein expression and purification

Rat nNOS α and rat nNOS μ were expressed and purified as previously described [10], with a few modifications. After sonication, the lysate was applied to a 50 ml DEAE Sepharose column (Sigma–Aldrich) equilibrated in 20 mM Tris–HCl, pH 7.4, 100 mM NaCl, 0.1 mM EDTA, 0.1 mM DTT, and 10% glycerol (Buffer B). Flow-through was loaded onto a 30 ml 2′5′-ADP Sepharose affinity column equilibrated in Buffer B, washed with 300 ml of Buffer B and eluted with 50 ml Buffer B containing 500 mM NaCl and 5 mM 2′-AMP. Calmodulin was prepared as described [16].

1.3. Spectrophotometric methods

CO difference spectra were obtained as described [15]. The molar protein concentrations for nNOSs were determined based on heme content via reduced CO difference spectra, where $\epsilon = 100 \text{ mM}^{-1} \text{ cm}^{-1}$ for $\Delta A_{445-470}$. All spectral analyses were performed using a Shimadzu Model 2401PC UV/visible dual-beam spectrophotometer.

1.4. Activity measurements

Nitric oxide formation and cytochrome *c* reduction were measured at 23 °C as described [17,18], in pH 7.4 buffer containing 50 mM Tris–HCl, 100 mM NaCl, and 200 μM CaCl_2 . Rates of NO synthesis and cytochrome *c* reduction were determined using extinction coefficients of $60 \text{ mM}^{-1} \text{ cm}^{-1}$ at 401 nm and $21 \text{ mM}^{-1} \text{ cm}^{-1}$ at 550 nm, respectively.

Oxidation of NADPH was monitored at 340 nm at 23° in pH 7.4 buffer containing 50 mM Tris–HCl, 100 mM NaCl, and 100 μM NADPH, with or without added l -arginine and CaM, as indicated. The rate was determined using an extinction coefficient of $6.2 \text{ mM}^{-1} \text{ cm}^{-1}$ at 340 nm for NADPH.

1.5. Stopped-flow spectrophotometry

Stopped-flow reactions were performed aerobically under turnover conditions at 23 °C, as described [9,19], using an Applied Photophysics SX.18MV diode array stopped-flow spectrophotometer. Reactions contained 1.5 μM enzyme, 100 μM NADPH, 10 μM H_4B , and 100 μM l -arginine in pH 7.4 buffer containing 50 mM Tris–HCl, 100 mM NaCl and, where indicated, 15 μM CaM. Heme nitrosyl formation and flavin reduction were monitored at 436 nm and 485 nm, respectively.

1.6. Laser flash photolysis

CO photolysis experiments were conducted as described [3]. Briefly, a solution ($\sim 350 \mu\text{l}$) containing 20 μM 5-deazariboflavin (dRF) and 5 mM fresh semicarbazide in pH 7.6 buffer (40 mM

Bis–Tris propane, 400 mM NaCl, 2 mM l -Arg, 20 μM H_4B , 1 mM Ca^{2+} and 10% glycerol) was degassed in a laser photolysis cuvette by a mixture of 1:3 CO/Ar for 90 min. Concentrated NOS was injected through a septum to the desired concentration, kept in ice, and further purged by passing the CO/Ar mixture over the surface for 60 min. The protein was illuminated for an appropriate period to obtain a partially reduced form of $[\text{Fe(II)}\text{--CO}][\text{FMNH}\cdot]$, then flashed with a 446 nm laser excitation to trigger the FMN–heme IET, which was followed by the loss of absorbance of Fe(II) at 460 nm [20].

2. Results

The absorption, EPR, and fluorescence spectra of the nNOS μ and nNOS α proteins are very similar (Figs. S1 and S2 in Supporting information), indicating that the insertion in nNOS μ likely does not perturb the protein environments of the heme and flavin moieties.

The presence of an additional 34 amino acids in nNOS μ in a known electron transfer regulatory region, the AR, might be expected to alter the rate of electron transfer through the reductase domain and/or into the oxygenase domain. Modulation of this activity by CaM, which both increases the electron transfer rate through the reductase domain and permits reduction of the heme, might also be altered. To examine this, NO synthesis activity, which requires electron transfer through the entire enzyme, and cytochrome *c* reduction, which probes electron transfer through the reductase domain only, were measured (Tables 1 and 2).

No difference in the rate of NO formation was observed between the variants (Table 1). Under optimal, fully coupled conditions, NO production requires 1.5 NADPH molecules per NO molecule formed. Deviation from this optimum indicates that reactive oxygen species are being formed at the expense of product (i.e., uncoupling). NADPH oxidation was monitored during NO synthesis to assess coupling of this process to product NO formation. No difference in the NADPH oxidation rate was observed (Table 1), suggesting no difference in the degree of coupling between the variants.

Table 1
Rates of NO synthesis and NADPH oxidation in the presence of substrate.

	NO synthesis (min^{-1})	NADPH oxidation (min^{-1})	Efficiency of coupling (NADPH/NO)
nNOS α (+) CaM	53.4 $\pm$ 5.0	113.8 $\pm$ 4.5	2.1
nNOS μ (+) CaM	53.7 $\pm$ 1.1	125.1 $\pm$ 13.2	2.3

Conditions are as described in Section 1. Final nNOS and CaM concentrations are 0.1 and 1 μM , respectively. Rates reported are the average of at least 3 reactions $\pm$ S.E.M.

Table 2
Rates of cytochrome *c* reduction in the absence and presence of CaM.

	Cytochrome <i>c</i> reduction (min^{-1})	CaM stimulation
nNOS α (–) CaM	852 $\pm$ 58	9.7 $\times$
(+) CaM	8290 $\pm$ 54	
nNOS μ (–) CaM	1280 $\pm$ 45	3.5 $\times$
(+) CaM	4503 $\pm$ 380	

Conditions are as described in Section 1. Final nNOS and CaM concentrations are 1 and 50 nM, respectively. Rates reported are the average of at least 3 reactions $\pm$ SEM.

The rate of cytochrome *c* reduction is dependent on the rate of electron flow through the reductase domain. In the absence of CaM, the rate of cytochrome *c* reduction is approximately 50% higher for nNOS μ over nNOS α , and is stimulated to a much lower extent by CaM in nNOS μ . However, the rate in the presence of CaM is 1.8-fold higher in nNOS α than nNOS μ (Table 2). Thus, electron transfer through the reductase domain of nNOS μ is stimulated significantly less by CaM than that of nNOS α (3.5-fold vs. 9.7-fold for nNOS μ and nNOS α , respectively).

NO synthesis was measured at different NOS concentrations (25, 50, 75, and 100 nM) in the presence of increasing amounts of CaM (molar ratios of CaM:nNOS ranging from 0.25 to 5) to determine whether activation by CaM differs between nNOS α and nNOS μ . The data were analyzed as described [21], which is based on evaluation of tightly binding inhibitors [22]. The relationship between fractional velocity and the AC_{50} for CaM is given in Eq. (1):

$$y = v_{\text{fmax}} / (1 + (AC_{50}/x)^n) \quad (1)$$

where *y* is the fractional activity (rate/maximum rate), *x* is the concentration of CaM, v_{fmax} is the maximum fractional activity (when concentration of CaM approaches infinity), *n* is the Hill coefficient. The data for all nNOS enzyme concentrations were fit by Eq. (1) (Fig. 1a and b) to derive AC_{50} values. The calculated AC_{50} values for nNOS α and nNOS μ were then plotted as a function of nNOS concentration. The estimated activation constant (K_{act}) for each variant is the ordinal intercept of the linear fit of this plot (Fig. 1c). The K_{act} values were calculated to be 2.45 and 4.65 nM for nNOS α and nNOS μ , respectively. Thus, the 34-amino acid insertion in the AR appears to have little to no effect on CaM binding to nNOS μ , as measured indirectly by catalytic activity.

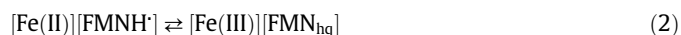
The rates of NO formation by nNOS α and nNOS μ at varying L-arginine concentrations with different CaM concentrations were measured to determine the effect of CaM on substrate binding (Fig. S3). The apparent K_m values were independent of CaM concentration ($3.19 \pm 1.2 \mu\text{M}$ and $3.68 \pm 1.95 \mu\text{M}$ ($\pm$ std. dev.) for nNOS α and nNOS μ , respectively), indicating that the interactions of both splice variants with L-arginine are independent of CaM binding.

In addition to NO synthesis with L-Arg, NOSs oxidize NADPH in the absence of substrate, forming reactive oxygen species [23,24]. We compared rates of NADPH oxidation by the two variants in the absence of substrate L-Arg. The NADPH oxidation rates of nNOS α and nNOS μ in the absence of CaM are similar and very low (3 ± 1 and $6 \pm 1 \text{ min}^{-1}$, respectively). In the presence of CaM, however, nNOS α oxidizes NADPH at approximately twice the rate of nNOS μ (490 ± 10 and $260 \pm 4 \text{ min}^{-1}$, respectively), once again

demonstrating that stimulation of activity by CaM is much lower for nNOS μ than nNOS α (45-fold vs. 154-fold, respectively).

Because stimulation by CaM affects electron transfer into the NOSs, as well as transfer between the two flavins and from FMN to the heme moiety, the rates of flavin reduction in the presence and absence of CaM were measured using stopped-flow spectrophotometry. This is a biphasic reaction, with a fast phase representing the initial input of electrons from NADPH [9,25] and a slow phase presumably reflecting comproportionation of the electrons between FAD and FMN. The fast rate of flavin reduction is equivalent for nNOS α and nNOS μ in the absence of CaM (Table 3, Fig. S4 in Supporting information). In the presence of CaM, however, the fast rate quadruples for nNOS α , but only doubles for nNOS μ , again showing that CaM stimulation of nNOS μ is half that of nNOS α . Similar behavior occurs in the slow phase of flavin reduction (Table 3).

As the interdomain electron transfer (IET) between the FMN and heme is essential for NO production at the catalytic heme site, IET kinetics of nNOS μ were measured using a laser flash photolysis approach [26,27]. As expected, CO dissociation by laser photolysis of the $[\text{Fe(II)}-\text{CO}][\text{FMNH}^+]$ form of CaM-bound nNOS μ results in a spectral “transition” (i.e., a change in direction of absorbance changes over time) in the 460 nm traces of the nNOS μ protein. Fig. S5 shows a typical kinetic trace of the nNOS μ protein at 21 °C: a rapid decay (with a rate constant k_1 of $33.4 \pm 0.4 \text{ s}^{-1}$) below the pre-flash baseline, followed by a much slower recovery toward the baseline with a rate constant k_2 of $2.7 \pm 0.1 \text{ s}^{-1}$ (due to CO rebinding to Fe(II)). The rapid absorbance decay (Fig. S5a) is due to net oxidation of heme in the following IET process, resulting in the loss of absorbance of Fe(II) at 460 nm [20,28,29]:



where FMN_{hq} stands for the FMN hydroquinone species. No IET was observed in the absence of CaM (data not shown), confirming that IET in nNOS μ is activated upon CaM-binding, as in nNOS α . However, the FMN-heme IET rate constant for nNOS α at room temperature is 47 s^{-1} [27], about 40% larger than that of nNOS μ .

To compare activation parameters of IET in nNOS μ and nNOS α , IET rate constants k_{et} were determined over the temperature range 283–304 K (Table S1 in Supporting information). An appreciable increase in the IET rate constant was observed with increasing temperature. Importantly, the obtained rate constant of the rapid decay (Fig. S5a) is independent of the signal amplitude in the range of temperature studied (data not shown), confirming an intra-protein IET process.

Viscosity studies indicated that the NOS FMN-heme IET process is conformationally gated [30]. The temperature dependence of k_{et}

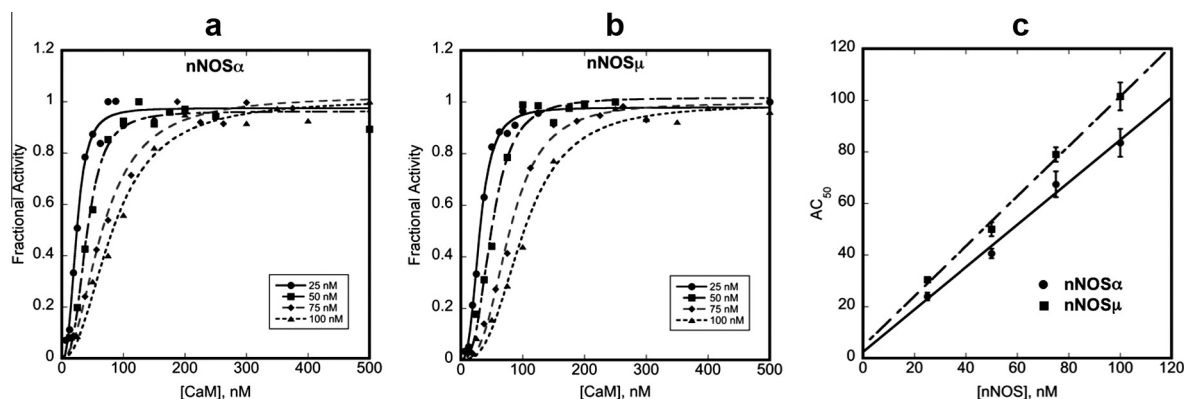


Fig. 1. Determination of CaM activation constant (K_{act}) of nNOS α and nNOS μ . The rates of NO production by nNOS α and nNOS μ were measured at different enzyme concentrations (25 nM, 50 nM, 75 nM and 100 nM) with varying CaM concentrations (6.25–500 nM). The AC_{50} values were calculated for each nNOS concentration for both nNOS α (a) and nNOS μ (b) using Eq. (1) ($R > 0.99$ for all fits). AC_{50} values were plotted against nNOS concentration (c) to determine K_{act} for CaM. $R = 0.9949$ and 0.9975 for the linear fits of nNOS α and nNOS μ data, respectively.

Table 3
Rapid kinetics measurements of flavin reduction and ferrous-nitrosyl complex formation.

	Flavin reduction (fast) (s ⁻¹)	Flavin reduction (slow) (s ⁻¹)	CaM stimulation (fast/slow)	Heme nitrosyl formation (s ⁻¹)
<i>nNOSα</i>				
(–) CaM	118.98 ± 2.52	7.90 ± 0.15	3.4×/3.5×	nd ^a
(+) CaM	401 ± 14.79	27.82 ± 0.42		41.85 ± 1.11
<i>nNOSμ</i>				
(–) CaM	124.72 ± 2.69	10.67 ± 0.14	1.5×/1.7×	nd ^a
(+) CaM	192.24 ± 4.57	18.26 ± 0.25		24.46 ± 1.4

Conditions are as described in Section 1. Rates reported are derived from 3 to 5 reactions averaged together and fit by a double exponential for flavin reduction and a triple exponential for heme nitrosyl formation.

^a Not detectable.

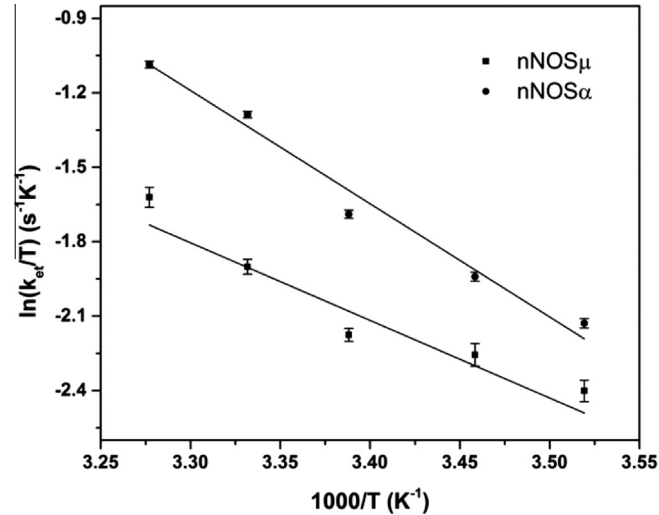


Fig. 2. Eyring plots showing the temperature dependence of k_{et} for *nNOSμ* (squares) and *nNOSα* (circles). The obtained ΔH and ΔS values are listed in Table 4.

Table 4
Eyring parameters from temperature dependence analysis of observed rate constants for the FMN–heme IET in *nNOSμ* holoenzymes, along with the FMN–heme IET rates k_{et} at room temperature.

	ΔH (kJ/mol)	ΔS (J/mol/K)	k_{et} (s ⁻¹)
Rat <i>nNOSμ</i> holoenzyme	25.7 ± 4.5	–127.2 ± 28.2	33.4 ± 0.4
Rat <i>nNOSα</i> holoenzyme	37.9 ± 3.2	–82.2 ± 7.9	47 ± 4 ^a

^a Data taken from Ref. [27].

was thus analyzed by transition state theory using the Eyring equation:

$$\ln(k_{et}/T) = -\Delta H^\ddagger/RT + \Delta S^\ddagger/R + \ln(k_B/h) \tag{3}$$

where $\Delta H^\ddagger$ is activation enthalpy, $\Delta S^\ddagger$ is activation entropy, h is Planck's constant, k_B is the Boltzmann constant, and R is the gas constant. The parameters of the *nNOSμ* data fit by Eq. (2) (Fig. 2) are listed in Table 4, along with those of the *nNOSα* protein. The obtained activation values are comparable to those of inter-flavin electron transfer in human cytochrome P450 reductase [31], which is homologous to NOS reductase domain.

Approximately 80% of *nNOSα* has been estimated to be quickly converted to an inactive form due to ferrous-nitrosyl complex formation [32–34], functioning to down-regulate NO synthesis. To compare this behavior between the two *nNOS* variants, ferrous-nitrosyl complex formation was monitored by stopped-flow spectrophotometry. The kinetics of heme-nitrosyl formation is complex, involving several steps [34]. The resulting trace was best fit

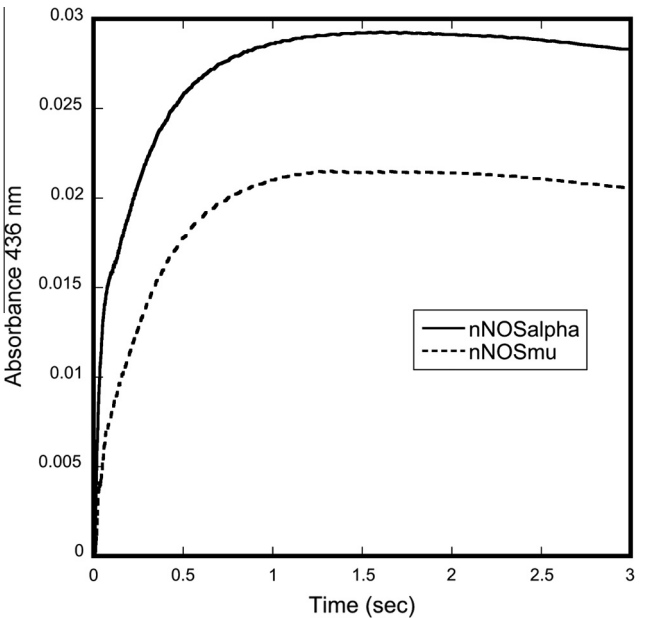


Fig. 3. Rate of ferrous nitrosyl complex formation monitored as a change in absorbance at 436 nm under turnover conditions using a stopped-flow apparatus. Conditions are as described in Section 1. Solid line: *nNOSα*, dashed line: *nNOSμ*. Traces shown represent an average of 3–5 experiments.

by a triple exponential; curve fits are shown in Fig. S5 of the Supporting information. The initial rate, i.e., the formation of the ferrous-nitrosyl complex, of *nNOSμ* is almost half that of *nNOSα* (Table 3 and Fig. 3), indicating that *nNOSμ* is inactivated at a much slower rate. In addition, the amplitude of the complex formation (Fig. 3) is lower for *nNOSμ*, indicating that a smaller proportion is in this inactivated state.

3. Discussion

The *nNOS* splice variant *nNOSμ* is found primarily in differentiated skeletal muscle and heart [11], while most other tissues express *nNOSα*. The expression of *nNOSμ* only in certain tissue types as well as the presence of the additional 34 amino acids in the auto-regulatory (AR) regions warrants a detailed mechanistic study of this variant in comparison to the much-studied variant, *nNOSα*. No major difference in the rate of NO production by *nNOSμ* vs. *nNOSα* was observed, corroborating prior results [14]. The rates of cytochrome *c* and flavin reduction in the absence of CaM were faster in *nNOSμ* than *nNOSα*, while the rates in the presence of CaM were smaller in *nNOSμ*. The magnitude of stimulation of the rate by CaM is thus notably lower in *nNOSμ*. The activation of *nNOSα* and *nNOSμ* by CaM shows little or no difference as the K_{act} val-

ues were 2.45 and 4.65 nM, respectively, indicating that the 34 amino acid insert in nNOS μ does not appear to alter the apparent affinity of CaM for the enzyme.

The measured FMN–heme IET rate of nNOS α is also higher than in nNOS μ . Despite the differences in the rate of electron flow, the rates of NO production by nNOS α and nNOS μ are similar (Table 1); this may be explained by a difference in their rates of autoinhibition by NO (Fig. 3) as all NOS isoforms are autoinhibited by NO to some degree due to the binding of NO to the heme to form a ferrous-nitrosyl complex [32,34]. The rate of ferrous nitrosyl complex formation is higher in nNOS α than in nNOS μ , and the percent difference between the variants is similar to that of the FMN–heme IET rates, indicating that the heme–NO complex formation is directly linked to the heme reduction rate. Previous studies of a nNOS α S1412D mutant also support this connection [35]. Thus it is likely that, at any given time under steady-state turnover conditions, more nNOS α exists in an inhibited state compared to nNOS μ , which may have physiological significance for skeletal muscle function.

The FMN–heme IET rate constant of nNOS α is about 40% larger than that of nNOS μ , and the activation entropy of nNOS α is notably smaller than nNOS μ (Table 4). Interestingly, the order of the IET rate constant (nNOS α > nNOS μ) correlates with the order of the magnitude of the activation entropy (Table 4). This can be explained by a conformational sampling model, in which sampling of a continuum of conformational states gives a range of transient donor–acceptor complexes, only a subset of which are IET-competent [36–38]; this avoids the necessity for tight binding of the FMN domain to achieve efficient IET. During NOS catalysis, the FMN domain cycles between the interaction with the NADPH/FAD domain (to receive electrons) and the interaction with the heme-containing oxygenase domain (to deliver electrons to the heme) [39], favoring the interaction with the NADPH/FAD domain [40]. In the NOS holoenzyme, even in the presence of bound CaM, the dominant conformational state is the docked FAD/FMN state [41], which must undergo a transformation to an IET-active, docked FMN/heme conformation before the FMN–heme IET occurs. Interdomain FMN/heme interactions in the NOS enzymes are rather weak, and the docked IET complexes are short-lived [30,40]. The observed rate of heme reduction is thus limited by the relatively infrequent formation of the docked IET-competent complexes. The larger negative activation entropy of nNOS μ indicates that the FMN domain needs to search for the IET-active conformation among a larger number of conformations that are not IET-competent than the nNOS α protein, resulting in less frequent formation of the FMN/heme complex and slower subsequent IET.

How an insertion of 34 residues in a regulatory region of the reductase domain influences the rates of heme reduction and ferrous nitrosyl complex formation is unknown, but one possibility is that the insert affects the stabilities and thus the time spent in various conformational states. The transfer of electrons to the heme requires the FMN domain to shuttle between an input (closed) or electron-accepting state and an output (open) or electron-donating state [39]. It was proposed that the AR and CT regulatory elements interact in some manner to stabilize either the open or closed conformation, depending on whether CaM and/or NADPH/NADP⁺ are bound [10]. Indeed, deletion of the AR regulatory element decreased the amount of enzyme present in the open conformation, implicating the AR as playing a role in the stability of the open complex [27]. Thus, the rate of electron transfer within the reductase domain and from the reductase to the oxygenase domain can be readily influenced by the dwell time in either a closed or open conformation.

The structural and functional aspects behind the differences in NADPH oxidation by the two variants in the absence of substrate

L-Arg may have profound implications on downstream signaling mediated by NO and/or production of reactive oxygen species (ROS), both of which can be produced by NOSs [24]. Reactions between NO and various oxygen species form products such as peroxynitrite and nitrate, which also have downstream effects. The observed decrease in NADPH oxidation by nNOS μ might be of great physiological importance as nNOS μ is differentially expressed only in heart and skeletal muscle. It will be important to curb or minimize the possible oxidative stress caused by nNOS μ as its expression/activity is up-regulated under exercise conditions [42] or by insulin stimulation [43]. Because these two nNOS isoforms have very similar catalytic properties *in vitro*, the sequence differences between them may also allow for differential regulation through protein/protein interactions and/or differential subcellular localization *in vivo*.

In conclusion, detailed biochemical analysis of nNOS μ and its comparison with nNOS α demonstrates that the presence of 34 additional residues in the AR affects electron flow through the enzyme, the enzyme response to CaM, and the formation of an inhibitory heme-nitrosyl complex. In addition, these studies suggest differences in the conformational equilibrium of the FMN domain, an important determinant of electron flux in NOS.

Acknowledgments

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.febslet.2013.10.032>.

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