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Structural Basis for Isoform-Selective Inhibition in Nitric Oxide Synthase

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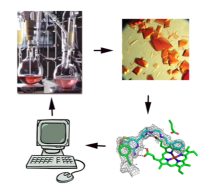
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CONSPECTUS

Itric oxide synthase (NOS) converts L-arginine into L-citrulline and releases the important signaling molecule nitric oxide (NO). In the cardiovascular system, NO produced by endothelial NOS (eNOS) relaxes smooth muscle which controls vascular tone and blood pressure. Neuronal NOS (nNOS) produces NO in the brain, where it influences a variety of neural functions such as neural transmitter release. NO can also support the immune system, serving as a cytotoxic agent during infections.

Even with all of these important functions, NO is a free radical and, when overproduced, it can cause tissue damage. This mechanism can operate in many neurodegenerative diseases, and as a result the development of drugs targeting nNOS is a desirable therapeutic goal. However, the active sites of all three human isoforms are very similar, and designing inhibitors specific for nNOS is a challenging problem. It



is critically important, for example, not to inhibit eNOS owing to its central role in controlling blood pressure.

In this Account, we summarize our efforts in collaboration with Rick Silverman at Northwestern University to develop drug candidates that specifically target NOS using crystallography, computational chemistry, and organic synthesis. As a result, we have developed aminopyridine compounds that are 3800-fold more selective for nNOS than eNOS, some of which show excellent neuroprotective effects in animal models.

Our group has solved approximately 130 NOS-inhibitor crystal structures which have provided the structural basis for our design efforts. Initial crystal structures of nNOS and eNOS bound to selective dipeptide inhibitors showed that a single amino acid difference (Asp in nNOS and Asn in eNOS) results in much tighter binding to nNOS. The NOS active site is open and rigid, which produces few large structural changes when inhibitors bind. However, we have found that relatively small changes in the active site and inhibitor chirality can account for large differences in isoform-selectivity. For example, we expected that the aminopyridine group on our inhibitors would form a hydrogen bond with a conserved Glu inside the NOS active site. Instead, in one group of inhibitors, the aminopyridine group extends outside of the active site where it interacts with a heme propionate. For this orientation to occur, a conserved Tyr side chain must swing out of the way. This unanticipated observation taught us about the importance of inhibitor chirality and active site dynamics.

We also successfully used computational methods to gain insights into the contribution of the state of protonation of the inhibitors to their selectivity. Employing the lessons learned from the aminopyridine inhibitors, the Silverman lab designed and synthesized symmetric double-headed inhibitors with an aminopyridine at each end, taking advantage of their ability to make contacts both inside and outside of the active site.

Crystal structures provided yet another unexpected surprise. Two of the double-headed inhibitor molecules bound to each enzyme subunit, and one molecule participated in the generation of a novel Zn^{2+} site that required some side chains to adopt alternate conformations. Therefore, in addition to achieving our specific goal, the development of nNOS selective compounds, we have learned how subtle differences in dynamics and structure can control protein—ligand interactions and often in unexpected ways.

Introduction

Structure based approaches to drug design date back to the 1970s with the development of compounds

designed to regulate hemoglobin^{1,2} and the antihypertensive drug, captopril.³ However, the wider acceptance of structure based methods coincided with the birth of

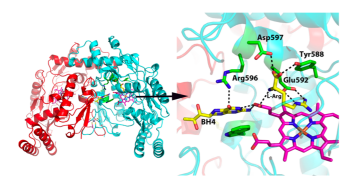


FIGURE 1. Rat nNOS heme domain dimer struture (10M4). L-Arg is held in place by a series of H-bonds to conserved active site residues as well as one heme propionate. The cofactor, BH_4 , is positioned at the dimer interface and also H-bonds with the same heme propionate.

the biotechnology industry in the early 1980s. With the availability of more interesting recombinant proteins, crystallographers had new proteins for structure determination, many of which were important drug targets. The hope was that structure based approaches would streamline drug discovery. In practice, however, the expense of determining crystal structures did not compare favorably with more rapid combinatorial chemistry approaches. To get around this problem was one of the major incentives of the so-called protein structure initiative, generously funded by NIH but met with justifiable skepticism.⁴ The basic idea is to dramatically lower the cost of structure determination and rapidly provide the structure of drug targets for structure based drug design. This would enable the rational design approach to effectively compete with more "random" synthetic chemistry approaches. It remains to be seen the long-range contributions of the protein structure initiatives, but we can ask if the basic approach of structure based drug design works which includes the development of clinically useful molecules. The answer is yes with perhaps the most widely known success story being the HIV protease. 5 This also is a spectacular example of what can be achieved by close collaborative efforts to move very quickly in the face of a health emergency.

The focus of this Account is our collaborative effort with Prof. Rick Silverman at Northwestern University to develop nitric oxide synthase inhibitors targeting neurodegenerative disorders. A similar review was published in 2009 with a focus on the medicinal chemistry end of this project.⁶ Here we focus on the protein structural end with an emphasis on new discoveries made since 2009.

NOS Structure

NOS catalyzes the oxidation of L-arginine to L-citrulline and nitric oxide (NO).

The first step of the reaction is very similar to cytochromes P450 with the exception that the tetrahydrobiopterin (BH_4) cofactor serves as a source of an electron.^{7,8}

The mechanism for the second step, *N*-hydroxy-L-arginine to NO and L-citrulline, remains unsettled and is still under active investigation.

Like cytochrome P450, the source of electrons is an FMN/FAD reductase which shuttles electrons from NADPH via the flavins to the heme. With NOS, however, the FMN/FAD reductase is fused to the C-terminal end of the heme domain which gives a large polypeptide over 1000 residues which dimerizes through the heme domains. Although the crystal structure of holo-NOS has not yet been determined, the structures of the individual heme $^{10-13}$ and reductase domains are known. The structures show that the substrate, L-Arg, is held in place by a series of H-bonds including the conserved active site Glu (Figure 1). The BH4 is situated at the dimer interface and H-bonds to the same heme propionates accepting H-bonds from L-Arg.

NOS as a Drug Target

Humans produce three isoforms of NOS: endothelial NOS (eNOS), neuronal NOS (nNOS), and inducible NOS (iNOS). ¹⁵ The active site structure is the same for all three with some key differences to be highlighted further on. NO produced by eNOS and nNOS binds to the heme of guanylate cyclase which activates the cyclase to produce cyclic GMP, the final signaling molecule. ¹⁶ eNOS helps to control proper blood

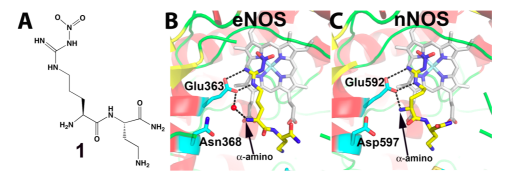


FIGURE 2. (A) Structure of one of the early dipeptide lead compounds, **1**, that exhibits excellent isoform selectivity. (B, C) show the crystal structures of the dipeptide inhibitor **1** in the active site of eNOS (PDB: 1P6L) and nNOS (PDB: 1P6H). In nNOS, the inhibitor "curls" which enables the inhibitor α -amino group to interact with both Glu592 and Asp597. In eNOS, Asn368 is the homologue to nNOS Asp597.

pressure while nNOS-derived NO is involved with neuronal signaling. iNOS is controlled at the level of transcription and is produced by the immune system where NO serves as a cytotoxic agent.

The over production of NO is associated with many neurodegenerative disorders, 17-22 and thus, it is highly desirable to develop nNOS inhibitors. The problem is selectivity. eNOS is critical for maintaining proper blood pressure, and thus, generic NOS inhibitors might well block the neurotoxic effect of too much NO in the brain but would also result in unwanted hypertension owing to the inhibition of eNOS. The goal of our joint project has been to use crystallography, computational methods, and synthetic chemistry to develop inhibitors that selectively block nNOS but not eNOS.

Initial Discoveries

Although the development of NOS inhibitors began very soon after NOS was discovered, structure based approaches could not be applied until the first crystal structures appeared in 1998. ^{10–13} In our lab, studies have focused on rat nNOS and bovine eNOS. From the perspective of isoform-selectivity, the NOS crystal structures were disappointing since the active sites for all three isoforms are nearly identical (Figure 1) which is not too surprising because all three catalyze exactly the same reaction. The earliest NOS inhibitors developed mimicked the substrate L-Arg. For example, prior to the availability of the crystal structures, L-nitroarginine was reported to be moderately selective for nNOS over iNOS. ²³

This provided a logical starting point for the development of more selective inhibitors and is where the Silverman lab stepped into the picture. It was reasoned that since the active site where the chemistry of ι -Arg oxidation occurs is the same in all NOS isoforms, selectivity is unlikely to be

achieved by simple variation on the basic L-Arg skeleton. However, further from the active site toward the surface of the enzyme, sequence diversity will be tolerated and thus it might be possible to add a "tail" to L-nitroarginine that extends out of the active site and possibly interact selectively to a unique patch on nNOS and not eNOS. Of course, there were no crystal structures although it was known that a conserved Glu is required for activity.²⁴ Therefore, a series of dipeptide inhibitors were developed using a substrate analogue, L- or D-nitroarginine, as the NOS active site recognition group linked by a peptide bond with a second amino acid or an other similar chemical group. This worked and resulted in dipeptide inhibitors that were about 1000-fold more selective for rat nNOS than bovine eNOS.^{25–27} Our laboratories then teamed up to try to understand the structural basis for selectivity. The best dipeptide inhibitors that exhibited about 1500-fold selectivity were chosen for initial crystal structure work. These structures provided the first Eureka moment and the answer to selectivity was shockingly simple.²⁸ As shown in Figure 2, the dipeptide inhibitor "curls" in nNOS but adopts an extended conformation in eNOS.

This is due to a single amino acid difference in the active site: where nNOS has Asp597, eNOS has Asn368. The α -amino group of the inhibitor maximally interacts with both Glu592 and Asp597 in nNOS if the inhibitor curls, thus providing additional electrostatic stabilization not possible in eNOS. This hypothesis was proven by swapping the Asp/Asn side chains in eNOS and nNOS, and, indeed, replacing the Asn with Asp in eNOS resulted in tighter binding and a curled conformation and just the opposite happened when the Asp in nNOS was replaced with Asn (Table 1).

The Importance of Chirality

The next generation of inhibitors has an aminopyridine head, a pyrrolidine bridge group, and a tail. A novel fragment

TABLE 1. K_i Values for Initial Dipeptide Inhibitors Used for Crystal Structure Determination²⁸

	$K_{\rm i}$ (μ M) of inhibitors		
	I	II	III
WT eNOS WT nNOS nNOS D597N eNOS N368D	107.0 0.30 67.0 9.5	80.0 0.15 34.0 4.6	110.0 0.10 21.0 5.1



Active site Glu

FIGURE 3. Initial design of aminopyridine-pyrrolidine inhibitors. The aminopyridine mimics the guanidinium group of L-Arg but exhibits a much lower pK_a , thus increasing bioavailability. In addition, the aromatic aminopyrridine should stack over the heme ring. The pyrrolidine NH group extends up toward the active site Asp597 in nNOS.

hopping computational method was used by the Silverman lab in the design process. The basic idea was for the aminopyridine to mimic the guanidinium group of L-Arg to H-bond to Glu592 while the pyrrolidine N atom, mimicking the α -amino N atom in the dipeptide, would be rigidly held by the ring between Glu592 and Asp597 (Asn in eNOS) thereby providing additional electrostatic stabilization in nNOS compared to eNOS (Figure 3).

Subsequent crystal structures showed that the initial design and docking studies closely matched the crystal structure.³⁰ These inhibitors have two chiral centers (Figures 3 and 4), and up to this point we were working with racemic mixtures.

Most importantly, a racemic mix of compound **2** (Figure 4) exhibits excellent protection from ischemic brain damage in a cerebral palsy animal model.³¹ The crystal structure of

enantiopure **2** complexed to nNOS provided a surprise.³² (3*S*,4*S*)-**2** bound as expected with the aminopyridine H-bonded to the active site Glu592. The *trans* versions of **2** also bound as expected. However, (3*R*,4*R*)-**2** flipped 180°, placing the aminopyridine in position to H-bond with the heme propionate. For this to happen, Tyr706 must adopt a new rotamer conformation. The reorientation of Tyr706 requires that the Tyr706-heme propionate H-bond be broken, a conserved interaction in all NOS isoforms for which structures are available. The new rotamer position enables the Tyr706 aromatic ring to stack against the inhibitor aminopyridine.

These structures and inhibition data presented two key questions. First, why is (3R,4R)-2 some 3800-fold more selective for nNOS over eNOS while (35,45)-2 is only about 500-fold selective; and second, why is the K_i for (3R,4R)-2 10fold lower than that for (35,45)-2? The structure of eNOS complexed with (3R,4R)-2 exhibited exactly the same flipped binding mode as in nNOS, so unlike the dipeptide inhibitors the isoform-selectivity in this case is not attributed to differences in the inhibitor conformation and binding mode. The first question was addressed by determining the K_i of various nNOS mutants designed to mimic eNOS. The two most important are Asp597 and Met336 in nNOS which are Asn and Val, respectively, in eNOS. Although Asp597 does not directly contact 2 in the flipped binding mode, the negative charge on Asp597 should increase electrostatic stabilization of (3R,4R)-2 especially in the relatively low dielectric environment of the active site. In this mutagenetic analysis, we also included the effects of Tyr706 which must move for inhibitors to bind in the flipped mode. We have consistently found that, for various inhibitors that bind in the flipped mode, Tyr706 forms a better stacking interaction with the aminopyridine in nNOS than eNOS. The only way to test the relative effects of this Tyr side chain was to simply covert it to Ala. The mutant that was most important to compare is the triple D597N/M336 V/Y706A nNOS mutant with the Y477A eNOS mutant. For the nNOS triple mutant, $K_i = 1290$ nM, a 240-fold increase, while $K_i = 35200$ nM for the Y477A eNOS mutant, a modest 1.7-old increase. It thus appears that a combination of additional electrostatic interactions and better nonbonded contacts in nNOS contribute to isoform selectivity.

More recently, we found that some aminopyridinepyrrolidine inhibitors bind in the flipped mode in both eNOS and nNOS but the Tyr moves only in nNOS (unpublished). This must mean Tyr706 is more flexible in nNOS than in eNOS. As shown in Figure 5, when Tyr706 rotates out to

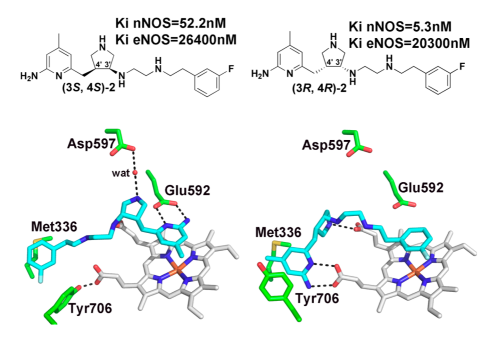


FIGURE 4. Crystal structures of the two enantiomers of cis-**2** bound to nNOS. (3*S*,4*S*)-**2** (PDB: 3JWS) binds as expected with the aminopyridine positioned over the heme near Glu592. However, (3*R*,4*R*)-**2** (PDB: 3JWT) binds in the flipped mode which places the aminopyridine near a heme propionate. This requires Tyr706 to adopt the "out" rotamer conformation.

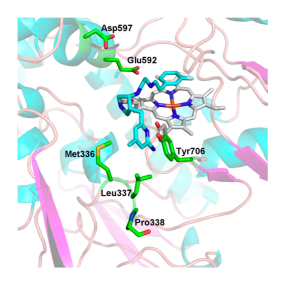


FIGURE 5. nNOS active site showing the inhibitor in the flipped mode (PDB: 3JWT). Note that the aminopyridine and Tyr706 are close to residues Met336 and Leu337. There is a break in the electron density right after Pro338, suggesting that this region may be particularly flexible in nNOS which enables Tyr706 to more readily adopt the out conformation relative to eNOS.

make room for the inhibitor, Tyr706 contacts Leu367, and we have found that nonbonded contacts in this region are, indeed, important.³³ Probing this region via mutagenesis, however, is problematic since the last well-defined residue is Pro338. After Pro338 nNOS electron density maps effectively disappear for about 10 residues clearly showing that this region is highly flexible. eNOS is similar with 10 residues

disordered after Pro108 (equivalent to Pro338 in nNOS) in most of the structures. However, with some inhibitor complexes, the electron density extends a few more residues, suggesting that this region of eNOS is slightly more rigid. If this region is more flexible in nNOS, then it might be easier for Tyr706 to adopt the rotamer conformation favorable for the flipped binding mode. In this scenario, Tyr706 is an equilibrium mix of both the "in" and "out" rotamers and the inhibitor binds best to the "out" rotamer. In nNOS the effective population of the "out" rotamer is higher than in eNOS which accounts in part for isoform selectivity.

The final question to address is why (3R,4R)-2 binds about 10-fold better to nNOS than (35,45)-2. Even though the binding modes are totally different, the structures alone did not provide much insight into why there should be a difference. Here is where computational approaches proved useful. The Molecular Mechanic/Poisson-Boltzmann Surface Area (MM-PBSA) procedure is a proven tool for calculating ligand binding free energies.34 The overall molecular mechanical internal energy (electrostatics, nonbonded, bond angles, distance, etc.), the electrostatic component of the solvation energy, and the nonpolar component of the solvation energy are calculated, and the results from all three are summed to give the total free energy of the complex, $G_{complex}$. Then the inhibitor is removed from the protein and exactly the same calculations are carried out to give the free energy of the protein (receptor) alone, $G_{receptor}$,

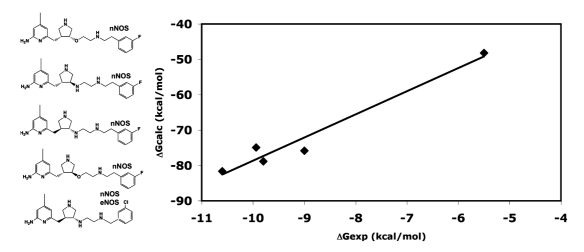


FIGURE 6. A total of seven crystal structures of the five different aminopyridine inhibitors bound to either nNOS or eNOS were used for the free energy calculations. The correlation between the relative ΔG_{calc} and experimental ΔG_{exp} (extracted from measured K_i values) is excellent.

and of the inhibitor alone, $G_{\text{inhibitor}}$. The total free energy of binding is then

$$\Delta G_{\text{bind}} = G_{\text{complex}} - G_{\text{receptor}} - G_{\text{ligand}} - T\Delta S$$

The most challenging part of these types of calculations is the entropy. As others have done³⁵ we ignored the entropy term which for the aminopyridine inhibitors is justified since the size and number of rotatable bonds is about the same for most of the inhibitors yet the affinities cover a wide range. This precludes the estimate of absolute ΔG but does provide a relative ΔG that can be compared to experimental values and enables the generation of "standard curves" as shown in Figure 6.

For such studies, parameters for the ligand are required which includes total charge. Compound 2 and related aminopyridines have 2-3 groups which can carry a full positive charge. For 2, this is the aminopyridine, the pyrrolidine, and the NH group closest to fluoro-phenyl ring so 2 can potentially carry a +3 charge. In the normal binding mode of (3S,4S)-2, the pyrrolidine N atom is about 4.1 Å from the aminopyridine. While both are interacting with Glu592, there is no net charge neutralization which raises the possibility that the aminopyridine, with a pK_a near neutrality, is only partially protonated. However, in the flipped binding mode, the pyrrolidine N atom and aminopyridine are 4.8 Å apart and both propionates of the heme are close so there is a net charge neutralization if the aminopyridine carries a full +1 charge. We used the MM-PBSA approach to compute the difference between these two models. In model 1, we assume that in both the flipped and normal binding modes the aminopyridine is fully protonated so the entire molecule

FIGURE 7. The initial design of double-headed symmetric inhibitors.

has a +3 charge. In model 2, we assume that the charge is +3 only in the flipped mode but in the normal binding mode the aminopyridine carries only a +0.5 charge so the net charge is +2.5. The MM-PBSA calculation was carried out for 11 aminopyridine-nNOS crystal structures that provided a mix of flipped and normal binding modes and then the calculated free energies plotted against experimental free energies. In the charge +3 model 1, the correlation coefficient of the plot was 0.52, but for the charge +2.5 model 2 the correlation coefficient increased to 0.81. The calculations and chemical intuition agree: the state of protonation of the inhibitor is highly dependent on inhibitor conformation and interactions with the protein and clearly contributes to binding affinity and isoform-selectivity.

Are Two Heads Better than One?

Since the aminopyridines can bind in two different orientations, the next obvious step was to build a double headed symmetric inhibitor of the form shown in Figure 7.

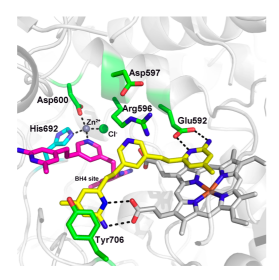


FIGURE 8. Crystal structure of **3** complexed to nNOS (3N5W). One molecule of **3** binds as predicted with one aminopyridine interacting with Glu592 and the other with the heme propionate. The second molecule of **3** displaces BH₄, thus enabling an aminopyridine to interact with the second heme propionate. This places the bridging pyridine in position to complete a tetrahedral coordination sphere around a Zn^{2+} ion. In order for the Zn^{2+} to bind, Arg596, which normally interacts with BH₄, must swing out of the way. In addition, the dimer interface must slightly tighten to enable His692 from molecule B of the dimer to move close enough for Zn^{2+} coordination.

Another advantage of these double headed compounds over the original parent inhibitors such as ${\bf 2}$ is an increase in lipophilicity which should present fewer problems in crossing the blood-brain barrier. We focus on compound ${\bf 3}$. The K_i for nNOS is 25 nM and for eNOS 2680 nM, giving a selectivity of about 107. Compound ${\bf 3}$ clearly is not as an effective inhibitor as ${\bf 2}$ nor is it as selective, but the better bioavailability might overcome these disadvantages. Indeed, compound ${\bf 3}$ exhibits an IC₅₀ of 5 μ M using a cell-based nNOS assay which is several fold better than ${\bf 2}$.

Once again, however, the crystal structures provided an unexpected surprise. The shown in Figure 8, there are two molecules of 3 bound to the nNOS active site. One of the molecules binds exactly as predicted. One aminopyridine interacts with the active site Glu while the other displaces Tyr706 and H-bonds with the heme propionate. The second molecule of 3 displaces the BH4 cofactor enabling one of the two aminopyridines to H-bond with the other heme propionate. Normally Arg596 H-bonds with BH4 but when the second molecule of 3 binds, Arg596 must swing out of the way where it now interacts with Asp597. This places the bridging pyridine of the second molecule of 3 in position to complete formation of a Zn^{2+} binding site. X-ray anomalous dispersion proves that this is a Zn^{2+} ion. In addition to the pyridine of 3, Asp600 and His692 together with a Cl^{-} ion

FIGURE 9. Variations on inhibitor **2** designed for better bioavailability.

complete the Zn^{2+} coordination sphere. This new Zn^{2+} site is close to the dimer interface and one of the ligands, His692, belongs to molecule B of the dimer. Thus, there is a slight tightening of the dimer interface in order to form the Zn^{2+} pocket. The affinity for Zn^{2+} must be fairly tight, since Zn^{2+} was not included in protein purification or crystallization.

With eNOS, only one molecule of **3** binds even though eNOS has exactly the same potential Zn²⁺ ligands as nNOS. Many mutants have been generated to try to understand this difference, but no consistent picture has emerged. One possible explanation is that the dimer interface in eNOS is tighter than in nNOS. As noted, the dimer interface must slightly tighten in order to enable His692 to be in position to coordinate Zn²⁺. If eNOS has restricted motion in this region, then the Zn²⁺ affinity will be much lower and the second molecule of **3** will not bind. This explanation is not particularly satisfying since there is no straightforward experimental test. Even so nNOS-**3** and additional structures of compounds that are variations of **3**³² provide the structural underpinning for alternative strategies in the development of isoform selective inhibitors.

Bioavailability

While the structural basis for isoform-selectivity is now understood and some of these compounds show great promise as therapeutic agents for neurodegenerative disorders, 31,38 bioavailability remains a problem. The main limitation is too much positive charge. Several well-known approaches have been taken in the Silverman lab to solve this problem. For example, introducing electron withdrawing fluorine atoms will lower the p K_a of a neighboring NH group and so compounds **4** and **5** (Figure 9) were synthesized. $^{39-41}$ The CF₂ group in **5** should lower the p K_a of the NH group to about 5.5–6 and turned out to be the most bioavailable inhibitor. The crystal structures of the NOS-**5** complex (PDB: 3NLX and 3NLU) showed that **5** binds

in the flipped binding mode as expected. The computed free energy of binding of **5** best matched experimental free energies when the NH group was assigned a +0.5 charge. In fact, the agreement was remarkably good, $\Delta G_{\rm calc} = -10.4$ kcal/mol and $\Delta G_{\rm exp} = -10.2$ kcal/mol, especially since the fluorophenyl group adopted multiple conformations so it was necessary to compute each structure and then average the results. **5** exhibits a K_i = 36 nM for nNOS and 140 000 nM for eNOS so the selectivity is about 3800-fold. Intravenous dosing of rats showed that **4** (**5** without the two gemfluorines) was 0.33 h and oral bioavailability essentially zero while **5** exhibited a half-life of 7.5 h and a 22% oral availability.³⁹

Summary

Although challenging, it has been possible to generate inhibitors selective for nNOS. Similar approaches to those we have described here also have been employed to develop iNOS inhibitors that exhibit exceptional selectivity over eNOS. 42 The basis for selectivity is quite different than we found with nNOS and is due primarily to inhibitors promoting formation of a novel binding pocket presumably due to greater flexibility near the iNOS active site. 43 While the problem of isoform selectivity is basically solved, the next step of moving into human clinical trials faces the bioavailability hurdle. Even so, some of the compounds we have developed exhibit excellent neuroprotective effects in animal models. NuerAxon also has developed nNOS-selective inhibitors that work well in animal migraine models.44 These efforts provide examples of how a combination of computational modeling, crystallography, mutagenesis, organic synthesis, and close collaborative research can solve an important drug design problem. This effort also has provided some interesting insights into how unexpected structural changes contribute to isoform-selectivity. The role of chirality in inhibitor binding modes, the binding of two of the symmetric double-headed inhibitors resulting in the "creation" of a novel Zn²⁺ site, and the formation of a novel binding pocket in iNOS⁴³ were unexpected. NOS has thus provided a platform for both further sharpening the tools of structure-based drug design and for probing the more basic questions on protein-ligand interactions.

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BIOGRAPHICAL INFORMATION

Thomas L. Poulos was born in Monterey California on Feb. 1, 1947. After graduation from Carmel High School, he earned a B.A. degree in Zoology at the University of California, Santa Barbara in 1968 followed by a Ph.D. in Biology in 1972 at the University of California at San Diego (UCSD). He then moved to the Chemistry Department at UCSD for postdoctoral work in the protein crystallography lab of Joe Kraut. While at UCSD, he solved the first heme enzyme crystal structure, cytochrome c peroxidase, and initiated work on P450s. In 1983 he was recruited to Genex Corp. in Gaithersburg, Maryland where he held the position of Principal Research Scientist and then Director of Protein Engineering. It was during this time that he solved the first cytochrome P450 structure. In 1987 he moved to the University of Maryland where he was a Professor of Chemistry and Director of the Center for Advanced Research in Biotechnology. In 1992 he moved to the Department of Molecular Biology and Biochemistry at UCI where he now holds the title of Chancellor's Professor and joint appointments in the Departments of Chemistry and Pharmaceutical Sciences. In 1991 he won the Presidential Meritorious Service Award from the University of Maryland and in 2004 the Brodie Award from the American Society of Experimental Pharmacology and Therapeutics. His primary research interests are in heme enzyme structure and function and structure-based drug design.

Huiving Li was born on July 26, 1951 in Beijing, P.R. China. He received a Bachelor degree in Chemistry in 1977 from Shanxi University, Taiyuan, PRC. He next earned a Ph.D. in Physical Chemistry in 1989 from University of South Carolina. In 1990 he joined the Poulos lab as a postdoctoral fellow at the University of Maryland's Center for Advanced Research in Biotechnology. Here is where he initiated his work on cytochromes P450 and was the first to solve the structure of a P450 in both the "open" and "closed" forms which provided the initial structural insights into the dynamics involved in substrate binding. He moved with Prof. Poulos to the University of California, Irvine (UCI) in 1992 where he played a key role in establishing protein crystallography at UCI. Dr. Li initiated the early work on nitric oxide synthase crystallography which ultimately resulted in the crystal structure of all three mammalian isoforms. Dr. Li currently holds the title of Project Scientist where he leads the lab's effort in nitric oxide synthase.

FOOTNOTES

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