

HNO/NO Conversion Mechanisms of Cu-Based HNO Probes with Implications for Cu,Zn-SOD

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Supporting Information

ABSTRACT: HNO has broad biological effects and pharmacological activities. Direct HNO probes for in vivo applications were recently reported, which are Cu^{II}-based complexes having fluorescence reporters with reaction to HNO resulting in Cu^I systems and the release of NO. Their coordination environments are similar to that in Cu,Zn-superoxide dismutase (SOD), which plays a significant role in cellular HNO/NO conversion. However, none of these conversion mechanisms are known. A quantum chemical investigation was performed here to provide structural, energetic, and electronic profiles of HNO/NO conversion pathways via the first Cu^{II}-based direct HNO probe. Results not only are consistent with experimental observations but also provide numerous structural and mechanistic details unknown before. Results also suggest the first HNO/NO conversion mechanism for Cu,Zn-SOD, as well as



useful guidelines for future design of metal-based HNO probes. These results shall facilitate development of direct HNO probes and studies of HNO/NO conversions via metal complexes and metalloproteins.

SECTION: Biophysical Chemistry and Biomolecules

I NO plays significant roles in many biological processes, Ho page activity regulation, enzyme activity regulation, and neurological function regulation. $^{1-3}$ It has also been found to possess unique pharmacological effects.^{2,4–6} For instance, it has a more favorable vasodilative effect than the well-known signaling molecule NO and an increased contractility effect, which enables HNO donors as a promising new class of vasodialtors and heart failure treatment.⁷ HNO has also been reported to inhibit GAPDH, allowing it to be an effective anticancer agent.8 However, most HNO detection methods, though useful in many scientific studies, are indirect or inconvenient for in vivo uses.^{1,4,9-18} Recently, intriguing breakthroughs were reported regarding direct HNO probes for in vivo applications,^{19–21} which are Cu-based complexes coupled with fluorescence reporters. The first complex with high selectivity against other reactive nitrogen and oxygen species including NO is Cu^{II}[BOT1].¹⁹ As shown in Figure 1, it has a BODIPY-based fluorescence reporter (X) and a tripodal receptor coupled via a triazole bridge. Upon reaction with HNO, the Cu^{II} center is reduced to Cu^I to trigger strong "turnon" type fluorescence in the visible light range, together with the release of NO. More recently, a new complex, $Cu^{II}[COT1]$, with excellent in vivo HNO imaging function and highly selective HNO detection was reported, which has a different fluorophore (coumarin-based) but the same Cu^{II} coordination site compared to Cu^{II}[BOT1].²⁰ Its reaction with HNO also results in a Cu^I complex. These results indicate the importance of this Cu center in these excellent Cu-based HNO probes. Interestingly, the Cu coordination environment of four nitrogen-containing ligands including the nitrogen-containing

bridging ligand in these systems is essentially the same as that found in Cu,Zn-superoxide dismutase (SOD),^{22,23} which was found to play a significant role in the cellular interconversion between HNO and NO.^{1,4,12} However, the conversion mechanisms in these Cu-based systems, identities of important species in the conversion pathways, and key structural and electronic factors responsible for reactivities remain to be elucidated.

Previous quantum chemical investigations have provided useful information to understand structural, spectroscopic, and mechanistic properties of HNO involvement in metalloproteins and related models.^{24–32} Here, we present results from a quantum chemical investigation of the HNO/NO conversion mechanisms via Cu^{II}[BOT1]. This first mechanistic information of HNO/NO conversion via copper-containing systems shall facilitate the development of additional HNO probes and studies of HNO reactions with other copper systems,^{12,13,16,20,21} particularly Cu,Zn-SOD.^{12,22,23}

As shown in Figure 1, based on the experimental results,¹⁹ HNO/NO conversion via Cu^{II}[BOT1] starts after HNO replaces the chloride ligand in the first step ($\mathbf{R} \rightarrow \mathbf{I-1}$). Then, the bound HNO is converted to bound NO, which is followed by the release of NO from the Cu complex. The HNO conversion to NO involves both proton transfer and electron transfer, which is similar to the superoxide reduction step by Cu^{II},Zn-SOD.^{33,34} Cu^{II}[BOT1] also has four nitrogen-contain-

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Figure 1. HNO/NO conversion pathways via Cu^{II}[BOT1].

ing ligands around the redox center like Cu^{II} ,Zn-SOD, including a bridging ligand, triazole, which is known as an efficient proton-transfer mediator like the bridging histidine ligand in Cu^{II} ,Zn-SOD. However, because triazole can accept the proton via the terminal and middle nitrogen atoms in the triazole ring, respectively,^{35–39} called the imidazole-like pathway (I) and the pyrazole-like pathway (II), both pathways as illustrated in Figure 1 were investigated. This process involves transition states (TS/TS') between the HNO-bound Cu^{II} complex (I-1) and the NO-bound Cu^{II} complex (I-2/I-2') and is completed with the formation of corresponding products (P/P') and the concomitant release of NO.

In order to obtain the appropriate first HNO/NO conversion mechanistic information, we first performed a methodological investigation of a number of density functional theory (DFT) methods, including the frequently used hybrid method B3LYP⁴⁰ and another hybrid functional mPW1PW91⁴¹ as well as the more recently developed functionals M06⁴² and ω B97XD⁴³ recommended previously for reaction energy calculations,^{44–48} together with several basis sets of different sizes and relativistic effects for metal centers. On the basis of the detailed comparisons in the Supporting Information, the mPW1PW91 method with the LanL2DZ basis for Cu, the 6-311++G(2d,2p) basis for core atoms including HNO, all copper coordinated atoms, and all heavy atoms in the triazole ligand, plus the 6-31G(d) basis for the rest part of molecules, were used in subsequent studies. Solvent effects are included in all geometry optimizations and subsequent frequency and energy calculations via the PCM model (see the Supporting Information). Because the complete molecules including the exact fluorophores in the experimental systems were used here, the best method was then used to investigate the conformational effect of the fluorophore on the reaction pathways, which

was found to be basically negligible; see the Supporting Information. This is consistent with the fact that it is separated from the metal reaction center by the bridging ligand, as well as the experimental results that both $Cu^{II}[BOT1]$ and $Cu^{II}[COT1]$ with the same reaction site and only different by the fluorophore part are excellent HNO probes.^{19,20} Therefore, the relatively more stable fluorophore conformation was used in the following investigations, as shown by the optimized structures in Figure 2.

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Figure 2. Optimized structures of the studied species in HNO/NO conversion pathways. Atom color scheme: N, blue; O, red; C, cyan; H, gray; Cu, orange; B, pink; F, purple.

As seen from Table 1, for the imidazole-like pathway I, the relative electronic energies with respect to I-1 (ΔE 's) show that after the uphill TS, the NO-bound I-2 state is actually more stable than the HNO-bound I-1 state. The NO binding with the Cu center in I-2 is also favorable as the release of NO consumes 7.11 kcal/mol of electronic energy. The inclusion of the quantum zero-point energy contribution (ΔE_{ZPE}) reduces

Table 1. Relative Ener	rgies of Key Sp	ecies in HNO/NO
Conversion Pathways	(unit: kcal/mol	l)

pathway	state	ΔE	ΔE_{ZPE}	ΔH	ΔG
Ι	I-1	0.00	0.00	0.00	0.00
	TS	12.55	10.11	9.49	12.12
	I-2	-10.03	-9.27	-9.19	-9.14
	$\mathbf{P} + \mathbf{NO}$	-2.92	-3.23	-1.68	-12.80
II	I-1	0.00	0.00	0.00	0.00
	TS'	19.98	17.05	16.55	18.88
	I-2′	-1.47	-1.68	-1.33	-2.07
	$\mathbf{P}' + \mathbf{NO}$	5.50	4.23	4.60	-5.36

the reaction barrier $(I-1 \rightarrow TS)$ by 2.44 kcal/mol. This barrier is further reduced at the enthalpy level (ΔH). However, after the inclusion of entropic contributions, the Gibbs free energy reaction barrier (ΔG^{\ddagger}) goes up by 2.63 kcal/mol. However, the formation of P in the NO release step becomes thermodynamically favorable due to a negative Gibbs free energy change from I-2 to P + NO. Because the enthalpy change in this step is positive, this thermodynamically favorable step is entropydriven. The qualitative energetic trends were found to be the same for the pyrazole-like pathway II; see Table 1. However, the imidazole-like pathway I was found to be the dominant mechanism compared to the pyrazole-like pathway II here due to both kinetically lower ΔG^{\ddagger} by 6.76 kcal/mol and thermodynamically lower overall reaction ΔG by 7.44 kcal/ mol. The preference of the imidazole-like pathway is consistent with previous mechanistic studies of triazole systems in other reactions.^{35,37–39} For this preferred imidazole-like pathway, the small reaction barrier (12.12 kcal/mol) coupled with a favorable overall Gibbs free energy change (-12.80 kcal/mol) is in good accord with the experimental observation that Cu^{II}[BOT1] is an excellent HNO detection system with facile reactivity.¹⁹ These results also suggest that for Cu,Zn-SOD with basically the same four nitrogen-containing ligands including an imidazole-like histidine bridging ligand, the HNO conversion to NO reaction may proceed in a similar imidazole-like mechanism as described here. The HNO to NO conversion via Cu,Zn-SOD may also be driven by both the more favorable enthalpy of the NO-bound system than the HNO-bound system (similar to I-2 versus I-1) and the favorable entropy change of NO release as in $I-2 \rightarrow P + NO$. Interestingly, the HNO/NO conversion barrier of 12.12 kcal/mol for Cu^{II}[BOT1] is also close to the experimental HNO/NO conversion barrier in Cu^{II},Zn-SOD (10.55 kcal/mol) calculated from the experimental rate constant of $1.0 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$ from the Eyring–Polanyi equation.¹⁵ The relatively more favorable ΔG^{\ddagger} in Cu,Zn-SOD could be due to the protein environment effect, which is under investigation in our lab now.

In I-1, the middle N1 atom (see Figure 2 for atom numbering) involved in the pyrazole-like pathway is actually in a better position to form hydrogen bonding with the H in HNO than the terminal N3 atom involved in the imidazole-like pathway; $\angle N1\cdots$ H-N of 109.7° is larger than $\angle N3\cdots$ H-N of 84.5°, and the N1…H distance is ~0.02 Å shorter than that of N3…H. However, the above data show that the imidazole-like pathway is more favorable. To help understand this result, a molecular orbital analysis of the HNO/NO conversion precursor I-1 was performed. As illustrated in Figure 3, in the highest occupied molecular orbital (HOMO), N3 is the only atom in the triazole ring with dominant contribution, while the contribution from the middle N1 is negligible. In addition, the lobe of N3's p orbital that is close to H of HNO was also found to be in-phase with this hydrogen atom in the HOMO (see



Figure 3. Isosurface representations of the (A) HOMO and (B) HOMO-3 in I-1. Contour values = ± 0.02 au.

Figure 3) by the same green colors, which may help the formation of the new N3–H bond in I-2 after proton transfer. In contrast, the first occupied orbital with a reasonably large contribution from the middle N1 atom, the proton acceptor in the pyrazole-like pathway, is the HOMO–3, which is of lower energy than the HOMO and thus harder to be accessed. Therefore, these results suggest that the preference of the imidazole-like pathway is due to relatively easier access of N3 than N1 as the proton acceptor.

Further analysis of the electronic and geometric results (full data in Tables S7–S9 (Supporting Information) and selected data in Table 2) shows that the proton transfer and electron

Table 2. Key Geometric and Electronic Properties in HNO/NO Conversion Pathways

pathway	state	$R_{\rm N5-H}$ (Å)	$R_{\rm Cu-N3}$ (Å)	$\rho_{\alpha\beta}^{ \mathrm{Cu}}(\mathrm{e})$	$\rho_{\alpha\beta}^{\rm NO}$ (e)
Ι	I-1	1.043	2.085	0.549	0.099
	TS	1.191	3.161	0.430	0.258
	I-2	2.238	3.561	0.024	0.946
	P + NO		3.540	0.000	1.000
II	I-1	1.043	2.085	0.549	0.099
	TS'	1.346	2.022	0.339	0.436
	I-2′	3.716	2.411	-0.122	0.758
	$\mathbf{P}' + \mathbf{NO}$		2.144	0.000	1.000

transfer are coupled during HNO/NO conversion pathways I and II, similar to the reduction step of Cu^{II},Zn-SOD by superoxide.^{22,23} For instance, for the preferred imidazole-like pathway I, the N5-H bond in HNO is elongated by ~0.2 Å from I-1 to TS and is further elongated by \sim 1.0 Å at I-2. These results indicate that the proton transfer is partially done in TS. From I-1 to I-2, as shown in Table S7 (Supporting Information), the most significant geometric change in the Cu coordination environment occurs with the Cu-N3 bond, associated with the proton-acceptor site in the bridging triazole ligand. As shown in Table 2, this bond is already broken in TS, and its change is basically done at I-2 because the R_{Cu-N3} difference between I-2 and P is only 0.021 Å. This kind of bridging ligand change is also similar to the reduction step of Cu^{II},Zn-SOD by superoxide.^{22,23} As seen from Table 2, the NO molety's spin density ($\rho_{\alpha\beta}^{NO}$) increases from 0.099 e in I-1 to 0.258 e in TS and subsequently changes to 0.946 e in I-2, which is basically the same as 1.000 e observed as released NO in the experimental products.¹⁹ Accordingly, copper's spin density $(\rho_{\alpha\beta}^{Cu})$ decreases by 22% from I-1 to TS and then further reduces by 74% to be 0.024 e at I-2, which indicates that the metal center is essentially Cu^I, as in P observed experimentally.¹⁹ These data show that similar to the proton-transfer process, the electron transfer from the NO moiety of HNO to the Cu center is also partially done in TS and mostly finished at I-2, indicating that it is coupled with the proton transfer. Additional calculations of initial setups with either proton transfer preceding electron transfer or vice versa (i.e., they are not coupled), however, all end up with the same results as those described above, which suggests that the proton-transfer and electron-transfer processes in HNO/NO conversion have a strong tendency to be coupled. In addition, these results indicate that during HNO conversion to NO, Cu^{II} and triazole are the electron and proton acceptors, respectively. Therefore, the following approaches may help build metal-based HNO probes with enhanced reactivities: (1) a metal center with stronger oxidizing power to facilitate electron transfer; (2) the

proton-accepting ligand with higher proton affinity and/or with electron-rich substituents to facilitate proton transfer; (3) the other metal ligand(s) in the system with electron-deficient substituents to facilitate electron transfer to the metal center. Other efforts such as using the ligand with strong fluorescence response and strong metal binding capability may also help.

In summary, this work provides the first structural, energetic, and electronic profiles of HNO/NO conversion pathways in the metal-based systems. Results not only are consistent with experimental observations but also provide numerous structural and mechanistic details unknown before. Results also suggest a HNO/NO conversion mechanism for Cu,Zn-SOD, which has the same coordination sphere and a similar bridging ligand with a similar HNO/NO conversion barrier. In addition, key structural and electronic factors responsible for observed reactivities were revealed for the first time, and specific guidelines for future design of metal-based HNO probes with enhanced reactivities were reported. These first time HNO/NO mechanistic results shall facilitate development of additional direct HNO probes and investigations of HNO/NO conversions via other metal complexes and metalloproteins.

ASSOCIATED CONTENT

S Supporting Information

Computational details and optimized coordinates (Tables S1–S24) are available. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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