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#### Introduction

Anthropogenic nitrous oxide (N2O) emissions have approximately 500 times the global warming potential of CO<sub>2</sub> per molecule<sup>1</sup> and represent the leading cause of ozone layer depletion.<sup>2</sup> Therefore, it is critical to understand the mechanisms by which both natural and synthetic catalysts convert N<sub>2</sub>O to benign compounds.<sup>3-5</sup> In nature, N<sub>2</sub>O is converted to N<sub>2</sub> according to eqn (1) as part of the nitrogen cycle.<sup>6,7</sup> This N<sub>2</sub>O reduction reaction (N2ORR) is catalyzed by nitrous oxide reductase (N2OR),8-10 a copper-dependent enzyme crucial to bacterial denitrification. The catalytic site of N<sub>2</sub>OR is a coppersulfide cluster known as Cuz,11 which has consistently been found to contain a  $[Cu_4(\mu_4-S)]^{n+}$  core in all its catalytically active forms.<sup>12-15</sup> Alternatively, heterogeneous catalysts are known to mediate the decomposition of N<sub>2</sub>O (deN<sub>2</sub>O) according to eqn (2). Although this disproportionation reaction is catalyzed most efficiently by Rh,16 explorations of earth-abundant alternatives have identified Cu-doped zeolites and CuO/CeO2 as viable catalyst materials.17-19

# Triazenide-supported [Cu<sub>4</sub>S] structural mimics of Cu<sub>Z</sub> that mediate N<sub>2</sub>O disproportionation rather than reduction<sup> $\dagger$ </sup>

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As part of the nitrogen cycle, environmental nitrous oxide (N<sub>2</sub>O) undergoes the N<sub>2</sub>O reduction reaction (N<sub>2</sub>ORR) catalyzed by nitrous oxide reductase, a metalloenzyme whose catalytic active site is a tetranuclear copper–sulfide cluster (Cu<sub>2</sub>). On the other hand, heterogeneous Cu catalysts on oxide supports are known to mediate decomposition of N<sub>2</sub>O (deN<sub>2</sub>O) by disproportionation. In this study, a Cu<sub>2</sub> model system supported by triazenide ligands is characterized by X-ray crystallography, NMR and EPR spectroscopies, and electronic structure calculations. Although the triazenide-ligated Cu<sub>4</sub>( $\mu_4$ -S) clusters are closely related to previous formamidinate derivatives, which differ only in replacement of a remote N atom for a CH group, divergent reactivity with N<sub>2</sub>O is observed. Whereas the formamidinate-ligated clusters were previously shown to mediate single-turnover N<sub>2</sub>ORR, the triazenide-ligated clusters are found to mediate deN<sub>2</sub>O, behavior that was previously unknown to natural or synthetic copper–sulfide clusters. The reaction pathway for deN<sub>2</sub>O by this model system, including previously unidentified transition state models for N<sub>2</sub>O activation in N–O cleavage and O–O coupling steps, are included. The divergent reactivity of these two related but subtly different systems point to key factors influencing behavior of Cu-based catalysts for N<sub>2</sub>ORR (*i.e.*, Cu<sub>2</sub>) and deN<sub>2</sub>O (*e.g.*, CuO/CeO<sub>2</sub>).

$$N_2O + 2H^+ + 2e^- \rightarrow N_2 + H_2O$$
 (1)

$$N_2 O \rightarrow N_2 + \frac{1}{2}O_2 \tag{2}$$

Our group has reported previous synthetic studies on structural<sup>20-22</sup> and functional<sup>23-25</sup> mimics of Cu<sub>z</sub> involving  $[Cu_4(\mu_4-S)]$  cores supported by bridging diphosphine or formamidinate ligands.<sup>11</sup> One of these complexes,  $[Cu_4(\mu_4-S)(NCN)_4]^-$  ( $[1^C]^-$ ), not only possesses an electronic structure similar to Cu<sub>z</sub> according to XAS analysis<sup>24</sup> but also was found to mediate a single turnover of N<sub>2</sub>ORR, producing N<sub>2</sub> and O<sup>2-</sup> along with  $[Cu_4(\mu_4-S)(NCN)_4]$  ( $1^C$ ) quantitatively (NCN = [MesN = CH-NMes]<sup>-</sup>, see Scheme 1).<sup>23</sup> In this reaction, it was proposed that one equivalent of  $[1^C]^-$  activates the N<sub>2</sub>O substrate (mimicking Cu<sub>z</sub> in N<sub>2</sub>OR) while another  $[1^C]^-$  equivalent acts as a sacrificial electron donor (mimicking the Cu<sub>A</sub> electron



Scheme 1 Divergent N2O reactivity of Cu4S model systems (Mes = 2,4,6-Me\_3C\_6H\_2).





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transfer site in N<sub>2</sub>OR). Based on computational modeling, µ-1,3-N<sub>2</sub>O binding across a Cu-S edge of the cluster enabled by sulfide redox non-innocence was proposed,24 although details of the N–O bond breaking pathway (e.g., the transition state structure) were not elucidated. This manuscript details the synthesis, characterization, and N<sub>2</sub>O reactivity of an analogous  $[Cu_4(\mu_4$ -S)(NNN)]<sup>*n*-</sup> system (NNN = [MesN=N-NMes]<sup>-</sup>; n = 0, 1; see Scheme 1). Unlike its formamidinate analogue  $[1^{C}]^{-}$ , the triazenide derivative  $[1^N]^-$  reacts with N<sub>2</sub>O in a 1 : 1 stoichiometry upon reductive activation, mediating a single turnover of deN2O rather than N2ORR. Computational analysis of the deN2O process includes transition state models for N-O cleavage and O-O coupling steps that were not identified in previous studies. The divergent selectivity of N<sub>2</sub>ORR for  $[1^{C}]^{-}$  vs. deN<sub>2</sub>O for  $[1^{N}]^{-}$ points to key factors influencing behavior of Cu-based catalysts for N<sub>2</sub>ORR (*i.e.*, Cu<sub>Z</sub>) and deN<sub>2</sub>O (*e.g.*, CuO/CeO<sub>2</sub>).

#### Results and discussion

Synthesis and characterization of the  $1^N$  system somewhat parallel those of the  $1^C$  system<sup>21-24</sup> but will be detailed here to facilitate important comparisons between complexes that are summarized in Table 1.

Addition of a toluene solution of elemental sulfur to a bright yellow THF solution of dicopper(1) precursor, Cu<sub>2</sub>(NNN)<sub>2</sub>, immediately produced an inky blue solution from which  $[Cu_4(\mu_4-S)(NNN)_4]$  (1<sup>N</sup>) began to spontaneously crystallize. Complex 1<sup>N</sup> is highly crystalline, showing only sparing solubility in THF, CH<sub>2</sub>Cl<sub>2</sub>, and CHCl<sub>3</sub> and no measurable solubility in CH<sub>3</sub>CN, Et<sub>2</sub>O, and toluene. Notably,  $1^{N}$  is indefinitely stable on a benchtop in open air. The <sup>1</sup>H NMR of  $1^{N}$  in CDCl<sub>3</sub> (Fig. S1<sup>†</sup>) is consistent with two inequivalent NNN environments (syn and anti to the  $\mu_4$ -S ligand, respectively), each with restricted N-C bond rotation. A similar interpretation of the <sup>1</sup>H NMR spectrum of  $\mathbf{1}^{\mathbf{C}}$  was attributed to intramolecular  $\pi$ -stacking interactions between neighboring mesityl groups evident by X-ray crystallography.<sup>23</sup> The UV-Vis-NIR spectrum of  $\mathbf{1}^{N}$  in CH<sub>2</sub>Cl<sub>2</sub> (Fig. 1a) is dominated by an intense charge transfer transition at  $\lambda_{max} =$ 602 nm ( $\varepsilon = 1.2 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ ) with a shoulder at  $\lambda_{\text{max}} \approx$ 768 nm. Another feature at higher energy ( $\lambda_{max} = 369$  nm) is

likely ligand-based, as it appears in all complexes examined, including previously reported Cu<sub>2</sub>(NNN)<sub>2</sub>.<sup>26</sup> For comparison, the analogous charge transfer transitions for **1**<sup>C</sup>, which is purple-colored, were observed at 561 ( $\epsilon = 1.4 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ ) and ~470 nm (shoulder).<sup>22</sup>

Complex  $1^{N}$  crystallized with cubic symmetry in the  $R\bar{4}3n$ space group, with the molecule residing on a crystallographic special position. The asymmetric unit (Fig. S3<sup>†</sup>) contains a single NNN ligand, a single Cu atom disordered over two positions, and a single S atom. Applying crystallographic symmetry elements reveals a cluster with four well-ordered NNN ligands related by a molecular  $S_4$  axis, along with a disordered inorganic core with eight distinct Cu positions and two S positions (Fig. S3<sup>†</sup>). To assist with interpretation, a computational model of 1<sup>N</sup> was built and optimized by DFT. (Here and throughout the study, computational models use methyl groups in place of mesityls.) The resulting structure (Fig. 1b) features a rectangle-based pyramidal Cu<sub>4</sub>S core with Cu-Cu distances of 2.45 and 2.85 Å and Cu-S distances of 2.24 Å. The crystallographic model was then analyzed by grouping together four Cu atoms with one S position and the other four Cu atoms with the other S position to match the computational model as closely as possible. This analysis indicated a rectangular pyramidal Cu<sub>4</sub>S core disordered over two positions, with experimental Cu-Cu distances of 2.405(3) and 2.965(3) Å and Cu-S distances of 2.176(2) Å (Fig. 1a). Grouping together the Cu and S atoms in alternative ways provided geometries dissimilar to the computational model, e.g., with unreasonably long Cu-S bonds. Additionally, this interpretation of the crystallographic data provides a model that is isostructural to 1<sup>C</sup>, which was reported to have Cu–Cu distances of 2.4226(6) and 3.0353(6) Å within its rectangle-based pyramidal Cu<sub>4</sub>S core.<sup>22</sup>

The cyclic voltammogram of  $\mathbf{1}^{\mathbf{N}}$  in  ${}^{n}\text{Bu}_{4}\text{NPF}_{6}/\text{THF}$  (Fig. S6†) showed a reversible reduction at -0.85 V vs. FeCp<sub>2</sub><sup>+/0</sup> to the  $[\mathbf{1}^{\mathbf{N}}]^{-}$  state, and an irreversible reduction at lower potentials to the  $[\mathbf{1}^{\mathbf{N}}]^{2-}$  state that is unstable on this timescale. The redox behavior mirrors that of  $\mathbf{1}^{\mathbf{C}}$ , which possesses a reversible reduction at lower potentials.<sup>22</sup> Thus, the substitution of four formamidinate

Table 1         Data comparison of triazenide- and formamidinate-ligated Cu <sub>4</sub> S clusters				
Property	$Cu_4S(NNN)_4$ (1 <sup>N</sup> )	$[\mathrm{Cu}_4\mathrm{S}(\mathrm{NNN})_4]^-([\mathbf{1^N}]^-)$	$\operatorname{Cu}_4 S(\operatorname{NCN})_4 (1^{\mathbb{C}})^a$	$[\mathrm{Cu}_4\mathrm{S}(\mathrm{NCN})_4]^- ([1^\mathbf{C}]^-)^b$
$\lambda_{\rm max}$ (nm)	602, 768 (sh)	620, 934	470 (sh), 561	566
$\varepsilon (M^{-1} cm^{-1})$	$1.2  imes 10^4$	$4.1  imes 10^3, 3.3  imes 10^3$	$1.4  imes 10^4$	$8.6 imes10^3$
d (Cu···Cu) (Å)	2.405(3)	2.5853(16)	2.4226(6)	2.502(1)
	2.965(3)	2.6031(16)	3.0353(6)	2.809(1)
	$2.405(3)^{c}$	$2.5853(16)^c$	$2.4226(6)^{c}$	2.532(1)
	$2.965(3)^{c}$	$2.6031(16)^{c}$	$3.0353(6)^{c}$	2.831(2)
$E^{\circ}$ (V vs. [FeCp <sub>2</sub> ] <sup>+/0</sup> )	-0.85	irreversible	-1.28	irreversible
g tensors	n/a	2.143, 2.066, 2.005 (rhombic)	n/a	2.043, 2.090 (axial)
$A(^{63,65}Cu)$ tensors (MHz)	n/a	≤122, 90, 100 (from <i>H</i> -strain)	n/a	15, 100
Redox-active MO <sup>d</sup>	S 3p, 25%	S 3p, 26%	S 3p, 21%	S 3p, 20%
	Cu 3d. 11% each	Cu 3d. 12% each	Cu 3d, 13% each	Cu 3d, 14% each

<sup>*a*</sup> From Johnson *et al.*<sup>22</sup> <sup>*b*</sup> From Johnson *et al.*<sup>23</sup> and Rathnayaka *et al.*<sup>24</sup> <sup>*c*</sup> Generated by crystallography symmetry. <sup>*d*</sup> Mulliken populations for the LUMO and SOMO, respectively.



Fig. 1 (a) Experimental and (b) computational data: UV-Vis-NIR of neutral (blue,  $CH_2Cl_2$ ) and anionic (red, THF) forms of  $1^N$  (left) and molecular structures of the neutral redox state of  $1^N$  (right). The X-ray crystal structure is depicted as 50%-probability ellipsoids for non-C, H atoms and as wireframe for C atoms; H atoms are omitted; and only one component of the disordered  $Cu_4S$  core is shown. Computations were done at the B3LYP/def2TZVPP level of DFT using the CPCM implicit solvation model ( $CH_2Cl_2$ ); computed UV-Vis-NIR spectra were generated using excitations calculated by TD-DFT and assuming 0.333 eV half-width at half height.

ligands for four triazenides shifts the redox potentials positively by approximately 0.4 V.

Chemical reduction of 1<sup>N</sup> was carried out with either CoCp<sub>2</sub>  $(E^{\circ'} = -1.33 \text{ V vs. FeCp}_2^{+/0})$  or K[FeCp(CO)<sub>2</sub>]  $(E^{\circ'} = -1.8 \text{ V vs.})$  $FeCp_2^{+/0}$  in THF or  $CH_2Cl_2$ .<sup>27</sup> In each case, a subtle color change from inky blue to a duller blue was observed. While the  $[1^N]$ [CoCp<sub>2</sub>] salt was too insoluble for solution analysis, use of Kryptofix-222 to form [1<sup>N</sup>][K(Krypt<sub>222</sub>)] provided material suitable for characterization. As expected, only resonances for the cationic portion were observed by <sup>1</sup>H NMR (Fig. S7<sup>†</sup>) due to the S = 1/2 ground state of the  $[1^N]^-$  anion. The  $[1^N][K(Krypt_{222})]$  salt is air-sensitive in solution, gradually converting to neutral  $1^{N}$ when left on the benchtop. The UV-Vis-NIR spectrum of  $[1^N]$ [K(Krypt<sub>222</sub>)] in THF (Fig. 1a) has three main features, all of which have lower molar extinction coefficients than for corresponding peaks for 1<sup>N</sup>. Charge transfer transitions at  $\lambda_{max} =$ 382 nm and  $\lambda_{max}$  = 620 nm ( $\epsilon$  = 4.1  $\times$  10<sup>3</sup> M<sup>-1</sup> cm<sup>-1</sup>) are minimally shifted in wavelength compared to 1<sup>N</sup>. However, the feature for  $\mathbf{1}^{\mathbf{N}}$  at  $\lambda_{\max} = 768$  nm shifts significantly to  $\lambda_{\max} =$ 934 nm ( $\epsilon = 3.3 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ) for  $[\mathbf{1}^{\bar{\mathbf{N}}}]^-$ . For comparison, purple-colored  $[1^{C}]^{-}$  was reported to have a charge transfer band at  $\lambda_{\text{max}} = 566 \text{ nm}$  ( $\varepsilon = 8.6 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ), and no NIR data was reported in the previous study.<sup>23</sup> Generally, both 1<sup>N</sup> and  $[1^N]^-$  have their optical transitions shifted to longer wavelengths by 40–50 nm compared to  $\mathbf{1}^{\mathbf{C}}$  and  $[\mathbf{1}^{\mathbf{C}}]^{-}$ .

Crystallization of [1<sup>N</sup>][K(Krypt<sub>222</sub>)] invariably gave thin plates with very weak X-ray diffraction, and X-ray quality crystals of other salts of  $[1^N]^-$  (e.g.,  $[1^N]$ [CoCp<sub>2</sub>],  $[1^N]$ [K(18-crown-6)]) were not obtained after repeated attempts. Compound [1<sup>N</sup>] [K(Krypt<sub>222</sub>)] crystallizes with orthorhombic symmetry in the C222<sub>1</sub> space group, with both cation and anion portions residing on crystallographic special positions (Fig. S9<sup>†</sup>). The  $[K(Krypt_{222})]^+$  unit exhibited severe disorder. Furthermore, within the  $[1^N]^-$  unit, the S atom was disordered over two positions and the NNN ligands exhibited significant disorder, too. Fortunately, the four Cu positions were well ordered. The crystallographic data indicates that, upon reduction from 1<sup>N</sup> to  $[1^N]^-$ , the Cu<sub>4</sub>S core converts from a rectangular towards a square-based pyramid shape, with experimental Cu-Cu distances of 2.5853(16) and 2.6031(16) Å. A DFT model of  $[1^N]^-$ (Fig. S10<sup>†</sup>) was found to have computed Cu-Cu distances of 2.54 and 2.74 Å, which are also shifted from rectangular towards square shaped compared to the DFT model of  $1^{N}$  (vide supra). The change from rectangular to square shape in the Cu<sub>4</sub> base was also observed previously upon reduction of  $\mathbf{1}^{C}$  to  $[\mathbf{1}^{C}]^{-}$ , the latter of which has experimental Cu-Cu distances of 2.502(1) and 2.809(1) Å.23

Having optimized computational models for  $1^{N}$  and  $[1^{N}]^{-}$ , next we conducted TD-DFT calculations to better understand the electronic transitions evident by UV-Vis-NIR. As shown in Fig. 1b, the salient features of the experimental UV-Vis-NIR

spectra are well captured in the computed spectra, especially when considering that the computational models used a truncated [MeN=N-NMe]<sup>-</sup> ligand in place of the [MesN=N-NMes]<sup>-</sup> ligand used experimentally. For  $[1^N]^-$ , excitations calculated at 639 nm (experimental: 620 nm) and 961 nm (experimental: 934 nm) are both charge transfers into the LUMO (MO145 $\beta$ ), which is a highly delocalized MO with 26% S 3p character, 48% Cu 3d character (12% per Cu), and significant contribution from the two NNN ligands syn to the  $\mu_4$ -S atom (Fig. 2a). The transition calculated at 639 nm involves excitation of an electron from MO144 $\beta$ , which has 74% Cu 3d character. Thus, this transition can be viewed as a combination of  $Cu \rightarrow S$  and  $Cu \rightarrow NNN$  MLCT. The transition calculated at 961 nm involves excitation of an electron from MO141 $\beta$ , which is exclusively based on the four NNN ligands (<5% Cu 3d, <5% S 3p). Thus, this transition can be viewed as NNN  $\rightarrow$  [Cu<sub>4</sub>S] CT. The corresponding transitions for neutral 1<sup>N</sup> are qualitatively similar but less readily interpreted because of some admixture of states via configuration interactions at each excitation wavelength (Fig. S4<sup>†</sup>).

The X-band EPR spectrum of  $[1^N][K(Krypt_{222})]$  at 77 K in a frozen CH<sub>3</sub>CN/CH<sub>2</sub>Cl<sub>2</sub> glass is shown in Fig. 2b. Fine structure was not well resolved in the first-derivative spectrum but was emphasized by plotting the second derivative (Fig. 2c). Surprisingly, hyperfine splitting due to the four <sup>63,65</sup>Cu nuclei ( $I_{Cu} = 3/2$ ) was not evident. Instead, the spectra were best simulated with hyperfine splitting from two equivalent <sup>14</sup>N nuclei. The simulation also required inclusion of significant "*H*strain", *i.e.*, unresolved hyperfine coupling that can be assigned

to a combination of the four <sup>63,65</sup>Cu centers and remaining <sup>14</sup>N nuclei. This pattern can be rationalized by analyzing the spin density from the DFT model of  $[1^N]^-$ , which is plotted in Fig. 2d. Of the total 1e<sup>-</sup> spin, only 0.113e<sup>-</sup> resides on each Cu center  $(0.452e^{-}$  total), meaning that most of the unpaired spin is ligand-centered. The largest contributor is the  $\mu_4$ -S atom, which carries 0.300e<sup>-</sup>. Of the remaining 0.248e<sup>-</sup> of NNN-centered spin, nearly all resides on the two NNN ligands syn to the  $\mu_4$ -S ligand, with effectively none on the two anti-NNN ligands. Thus, the  $A(^{14}N)$  coupling observed by EPR spectroscopy can be assigned to the central N-atoms of the two syn-NNN ligands. The small and apparently unresolved  $A(^{63,65}Cu)$  coupling is consistent with the high degree of covalency (*i.e.*,  $\mu_4$ -S and NNN redox non-innocence) apparent from calculations. Although the EPR spectroscopy of  $[1^{C}]^{-}$  differs in that it is an axial signal dominated by  $A(^{63,65}Cu)$  coupling,<sup>23</sup> nonetheless the redox-active MOs for both the  $1^{N}$  and  $1^{C}$  systems are similar to each other (Table 1).24

No reaction was observed when  $[1^N][K(Krypt_{222})]$  was exposed to N<sub>2</sub>O (1 atm) in CH<sub>2</sub>Cl<sub>2</sub>. This behavior contrasts that of  $[1^C]$ [K(18-crown-6)], which converts quantitatively to  $1^C$  under the same conditions.<sup>23</sup> In the latter case, it was proposed that reversible N<sub>2</sub>O binding to  $[1^C][K(18-crown-6)]$  to transiently form  $[1^C \cdot N_2O][K(18-crown-6)]$  shifts the reduction potential such that a second equivalent of  $[1^C][K(18-crown-6)]$  can act as an electron donor, producing  $[1^C \cdot N_2O][K(18-crown-6)]_2$  and  $1^C$ (*i.e.*, N<sub>2</sub>O-induced disproportionation of the copper cluster). The shift of the  $1^N/[1^N]^-$  potential to more positive values



Fig. 2 (a) Charge transfer transitions (0.03 isosurfaces) for  $[1^N]^-$  calculated by TD-DFT at the B3LYP/def2TZVPP level using the CPCM implicit solvation model (CH<sub>2</sub>Cl<sub>2</sub>). Experimental and simulated X-band EPR spectroscopy for  $[1^N]$ [K(Krypt<sub>222</sub>)] in frozen CH<sub>3</sub>CN/CH<sub>2</sub>Cl<sub>2</sub>(1 : 1) glass at 77 K: (b) 1st derivative and (c) 2nd derivative spectra. (d) Spin density (0.001 isosurface) for  $[1^N]^-$  calculated by DFT. EPR simulation parameters: g = 2.143, 2.066, 2.005; A = 193, 68, 98 MHz for 2 equivalent <sup>14</sup>N nuclei; *H*-strain = 122, 90, 100 MHz; Gaussian line broadening = 0.43.

apparently prevents such N<sub>2</sub>O-induced disproportionation from occurring spontaneously for the triazenide derivative. Therefore, the reaction was repeated with a stronger sacrificial electron donor,  $CoCp_2$ , to reduce a transiently-formed  $[1^N \cdot N_2O]$ [K(Krypt<sub>222</sub>)]. A 1:1 mixture of  $[1^N]$ [K(Krypt<sub>222</sub>)] and CoCp<sub>2</sub> in CH2Cl2 solution was observed to darken slightly upon exposure to N<sub>2</sub>O (1 atm). Over 24 h, the major species in solution remained  $[1^N]^-$  according to UV-Vis-NIR spectroscopy. However, analysis of the reaction mixture by <sup>1</sup>H NMR spectroscopy indicated small conversion ( $\sim 10\%$ ) of CoCp<sub>2</sub> to  $[CoCp_2]^+$  along with formation of a new, diamagnetic product (2) containing the NNN ligand. The same product 2 was generated when exposing a 1 : 2 mixture of  $1^{N}$  and CoCp<sub>2</sub> to N<sub>2</sub>O (1 atm) or when replacing N2O with excess Me3NO under N2. Compound 2 could be extracted into toluene or Et<sub>2</sub>O, which served to separate it from  $[CoCp_2]^+$ ,  $[1^N]^-$ , and/or  $1^N$ . Washing 2 with pentane served to separate it from CoCp<sub>2</sub> (and excess Me<sub>3</sub>NO where relevant, see Fig. S12<sup>†</sup>). At this point, compound 2 was found to have a brown color, with a UV-Vis-NIR spectrum in CH<sub>2</sub>Cl<sub>2</sub> (Fig. 3a) showing a single distinct feature at  $\lambda_{max} = 420$  nm but lacking any welldefined charge transfer transitions in the 600-1000 nm region as observed for  $1^{N}$  and  $[1^{N}]^{-}$ . From these collected observations, it can be concluded that (1) the reaction involves O-atom transfer from N<sub>2</sub>O since Me<sub>3</sub>NO can replace it, (2) the resulting product 2 is diamagnetic and neutral in charge based on NMR characterization and solubility properties, and (3)

compound 2 likely lacks the  $\mu_4\mbox{-}S$  atom that would produce low-energy CT electronic transitions.

Unfortunately, all reaction conditions explored consistently provided only low conversion (<15%) to 2, with the remaining mass balance being predominantly  $[1^N]^-$ . Furthermore, repeated attempts at obtaining X-ray quality crystals of 2 failed. The identity of 2 was made clear by analyzing it using EI-HRMS. The parent ion mass and isotope distribution (Fig. 3a) indicate the empirical formula,  $Cu_4(O_2)(NNN)_4$ . To investigate this possibility further, the structure of 2 assuming that formula was modeled computationally. Two energy minima were located: one with  $\mu_2:\eta^2$  binding of an O<sub>2</sub> unit to one Cu from the periphery of the Cu<sub>4</sub> core, and another with  $\mu_3$ : $\eta^2$ , $\eta^1$ , $\eta^1$  binding of the O<sub>2</sub> unit to three Cu atoms in the capping position typically occupied by sulfide (Fig. S13<sup>†</sup>). The former isomer (Fig. 3b) was calculated to be lower in Gibbs free energy by -13.6 kcal mol<sup>-1</sup>. The predicted UV-Vis-NIR spectrum (Fig. 3b) of this structure reproduces key features of the experimental spectrum, with a prominent absorbance at  $\lambda_{max} = 448$  nm (experimental: 420 nm) that tails into the lower energy region of the spectrum without well-defined absorbance features in the 600–1000 nm region. The local structure within 2 of the  $Cu(O_2)$ unit in a nitrogen-rich environment resembles 1:1 Cu:O2 species that have been long studied since seminal works of Kitajima, Tolman, and others.<sup>28,29</sup> Notably, the presence of an absorbance at ~400 nm is characteristic of such compounds



Fig. 3 Characterization of complex 2: (a) experimental UV-Vis-NIR (left,  $CH_2Cl_2$ ) and high-resolution mass spectrometry (right, positive EI mode) data; (b) computed UV-Vis-NIR spectrum (left) and molecular structure (right). Computations were done at the B3LYP/def2TZVPP level of DFT using the CPCM implicit solvation model ( $CH_2Cl_2$ ); computed UV-Vis-NIR spectra were generated using excitations calculated by TD-DFT and assuming 0.45 eV half-width at half height.

and is typically assigned to the  $\pi \rightarrow \pi^*$  transition of the bound  $O_2$ .<sup>30</sup> The calculated O–O distance of 1.37 Å in 2 and the short calculated Cu–O distances of 1.85 and 1.86 Å are all indicative of a highly reduced  $O_2$  unit.<sup>31,32</sup> Curiously, the <sup>1</sup>H NMR spectrum for 2 shown only one set of NNN peaks, rather than the three sets expected based on the calculated structure. This is likely indicative of fluxionality due to the  $O_2$  ligand rapidly hopping from Cu site to Cu site.

Based on these observations, a plausible mechanism for a hypothetical deN<sub>2</sub>O reaction for is presented in Scheme 2. Reversible binding of N<sub>2</sub>O to  $[1^N]^-$  would generate a small equilibrium concentration of  $[1^N \cdot N_2O]^-$ . Despite CoCp<sub>2</sub> not being a strong enough reductant to spontaneously reduce  $[1^N]^-$ , reduction of  $[1^N \cdot N_2O]^-$  could be spontaneous due to the  $\pi$ accepting nature of N<sub>2</sub>O raising the reduction potential of the Cu<sub>4</sub>S core. The resulting  $[1^N \cdot N_2O]^{2-}$  intermediate would, then, be the active catalyst in the "fully reduced" 4Cu<sup>I</sup> redox state.<sup>33</sup> Based on previous computational modeling with  $[1^C \cdot N_2O]^{-,24}$ a  $\mu$ -1,3-N<sub>2</sub>O binding mode spanning a Cu–S cluster edge is proposed. Low conversion to 2 observed experimentally is likely due to inefficiency of this reductive activation process, *i.e.*, most of the dissolved complex remains unactivated as  $[1^N]^-$ .

On the active cycle, loss of  $N_2$  from  $[1^{N} \cdot N_2 O]^{2-}$  would generate  $[1^N \cdot O]^{2-}$ , which is proposed to feature  $\mu_2 \cdot O^{2-}$  binding based on previous modeling for  $[1^C \cdot O]^{2-}$ .<sup>23</sup> For N<sub>2</sub>ORR chemistry associated with the  $1^C$  system, dissociation of  $O^{2-}$  from  $[1^C \cdot O]^{2-}$  generates  $1^C$ , which can hypothetically undergo  $2e^$ reduction under N<sub>2</sub>O to regenerate  $[1^C \cdot N_2 O]^{2-}$ . In the  $1^N$ 



Scheme 2 Plausible mechanisms for N<sub>2</sub>ORR and deN<sub>2</sub>O.

system, N<sub>2</sub>ORR chemistry is not observed. Instead, a second N<sub>2</sub>O molecule presumably intercepts  $[1^{N} \cdot O]^{2^-}$  to generate  $[1^{N} \cdot O_2]^{2^-}$  with loss of N<sub>2</sub>. To complete the hypothetical deN<sub>2</sub>O cycle, O<sub>2</sub> loss would form  $[1^{N}]^{2^-}$ , which would regenerate  $[1^{N} \cdot N_2O]^{2^-}$  upon N<sub>2</sub>O coordination. On the other hand, if S<sup>2-</sup> dissociation outcompetes O<sub>2</sub> dissociation, then off-cycle species 2 is formed. (Other pathways to 2 cannot be ruled out, *vide infra.*) Experimentally, examination of the headspace gas above the N<sub>2</sub>O reaction revealed only trace quantities of N<sub>2</sub> and O<sub>2</sub>, consistent with the off-cycle decomposition to 2 outcompeting on-cycle N<sub>2</sub>O-for-O<sub>2</sub> substitution.

To gain further insight, the deN<sub>2</sub>O cycle was modeled computationally. The reaction energy profile is shown in Fig. 4, which features models for the four key intermediates as well as the two transition states associated with N-O cleavage of N<sub>2</sub>O. The computed intermediate  $[1^{N} \cdot N_2 O]^{2-}$  (A) is isostructural to  $[1^{C}\!\cdot\!N_{2}O]^{2-}$  calculated previously,^4 with  $\mu\text{-}1,3\text{-}N_{2}O$  binding across a Cu-S edge of the cluster and significant N<sub>2</sub>O activation according to the bent O-N-N angle of 126°. Loss of N<sub>2</sub> from A to form  $[\mathbf{1}^{\mathbf{N}} \cdot \mathbf{O}]^{2-}$  (**B**) is exergonic by  $\Delta G = -28.2 \text{ kcal mol}^{-1}$  and proceeds *via* transition state **TS1** with a barrier height of  $\Delta G^{\ddagger}$  = 12.9 kcal mol<sup>-1</sup>. The structure of **TS1** differs from that of reactant **A** in that the  $N_2O$  ligand occupies a  $\mu$ -1,1-O binding mode spanning a Cu-Cu cluster edge in TS1. This binding mode has been previously proposed for a synthetic model complex<sup>34</sup> and for N<sub>2</sub>O activation by Cu-ZSM-5 <sup>35</sup> but differs from that proposed for Cu<sub>z</sub> itself.<sup>12</sup> At **TS1**, the N<sub>2</sub>O molecule has undergone further activation as evidenced by the elongated N-O bond distance (1.57 Å) and contracted N-N bond distance (1.14 Å) relative to those in A (1.297 and 1.233 Å, respectively). The Cu-Cu edge of the cluster with the N<sub>2</sub>O bound is short (2.54 Å) to accommodate the bridging ligand and is compensated by a long Cu-Cu distance of 3.35 Å along the edge opposite the N<sub>2</sub>O ligand. The structure of B following N2 loss features a µ-oxo ligand with a Cu–O–Cu angle of 87° and a Cu–Cu distance (2.56 Å) similar to that in TS1.

Oxygenation of **B** by N<sub>2</sub>O to form  $[1^{N} \cdot O_2]^{2-}$  (C) was calculated to be exergonic by  $\Delta G = -9.6$  kcal mol<sup>-1</sup> and proceeds *via* transition state **TS2**. The structure of **TS2** features the N<sub>2</sub>O substrate approaching the Cu–O–Cu unit with an O···O distance of 1.84 Å. The N<sub>2</sub>O unit is significantly activated in **TS2**, with a bent O–N–N angle of 143°, a long O–N distance of 1.50 Å, and a short N–N distance of 1.13 Å. Unlike compound 2 formed from S<sup>2–</sup> loss, the structure of **C** features a  $\mu_2$ : $\eta^1$ -O<sub>2</sub> ligand. The calculated O–O distance of 1.35 Å in C is indicative of significant O<sub>2</sub> reduction,<sup>31</sup> though the Cu–O distances in C (2.02 and 2.06 Å) are longer than those calculated for **2**.

Loss of  $O_2$  from **C** to form  $[\mathbf{1}^{\mathbf{N}}]^{2-}$  (**D**) was calculated to be exergonic by  $\Delta G = -29.8$  kcal mol<sup>-1</sup>. Unlike intermediates **B** and **C** and transition states **TS1** and **TS2** that all involve bridging *O*-ligands along one cluster edge, the structure of **D** features a nearly square-shaped tetracopper core, with all calculated Cu– Cu distances between 2.66 and 2.78 Å. To complete the hypothetical catalytic deN<sub>2</sub>O cycle, coordination of N<sub>2</sub>O to **D** to regenerate **A** was calculated to be endergonic by  $\Delta G =$ +25.2 kcal mol<sup>-1</sup>. Thus, although the net deN<sub>2</sub>O reaction is calculated to be favorable ( $\Delta G_{rxn} = -42.4$  kcal mol<sup>-1</sup>), the rate-



Fig. 4 Reaction intermediates and transition states for deN<sub>2</sub>O calculated by DFT, with Gibbs free energy values given in units of kcal mol<sup>-1</sup>. Computations were done at the B3LYP/def2TZVPP level of DFT using the CPCM implicit solvation model (CH<sub>2</sub>Cl<sub>2</sub>) except for **TS2** (see ESI†).

determining N<sub>2</sub>O binding step prevents efficient catalysis under the experimental conditions examined, allowing for off-cycle conversion of C to inactive 2. At this time, the path from C to 2 is ambiguous: it could involve simple  $S^{2-}$  dissociation from C to form 2 directly but could also involve exergonic O<sub>2</sub> dissociation from C to form D followed by O<sub>2</sub>-promoted conversion of D to 2.

Oxygenation of **B** to form **C** presumably involves nucleophilic addition of the N<sub>2</sub>O oxygen to the electrophilic  $\mu$ -oxo ligand of **B** (see **TS2**). In accord with that hypothesis, analysis of the frontier molecular orbitals of **B** indicated that the LUMO has significant (24%) oxygen 2p character (Fig. 5). One reason that the **1**<sup>N</sup> system is more likely than **1**<sup>C</sup> to do deN<sub>2</sub>O chemistry might be



Fig. 5 Lowest unoccupied molecular orbital (0.03 isosurface) calculated for  $[1^N \cdot O]^{2-}$  (B) by DFT at the B3LYP/def2TZVPP level using the CPCM implicit solvation model (CH<sub>2</sub>Cl<sub>2</sub>).

the stabilization of this LUMO due to the electronegative nature of the NNN<sup>-</sup> ligands relative to less electronegative NCN<sup>-</sup>, thus favoring O-O coupling by N<sub>2</sub>O nucleophilic addition in the NNN<sup>-</sup> case. A second contributing reason could be that the relatively electron-withdrawing NNN<sup>-</sup> ligands make the tetracopper core more Lewis acidic than for the NCN<sup>-</sup> analogues, thus causing O<sup>2-</sup> loss required for N<sub>2</sub>ORR to be less facile for **B** and instead shunting the **1**<sup>N</sup> system into the deN<sub>2</sub>O pathway.

#### Conclusions

A redox pair of  $Cu_4(\mu_4$ -S) clusters supported by triazenide ligands was synthesized and thoroughly characterized. While much of the characterization data mirrored those of the related formamidinate-supported system studied previously,21-24 divergent reactivity behavior with N2O was observed. While the formamidinate-ligated system was found to promote singleturnover N<sub>2</sub>ORR akin to the biological Cu<sub>Z</sub> cluster it mimics, the triazenide system was found to promote (inefficient) deN<sub>2</sub>O reactivity that is a typical hallmark of heterogeneous catalysts like CuO/CeO<sub>2</sub>. Computational modeling of the deN<sub>2</sub>O pathway allowed for comparisons to be made with previously studied N<sub>2</sub>ORR chemistry, enabling deeper understanding of factors influencing selectivity. A simple working model is as follows. The formamidinate system is relatively electron-rich, providing a "soft" tetracopper core that behaves like Cuz (that features a soft sulfide ligand) by readily dissociating  $O^{2-}$  upon  $N_2O$ deoxygenation to favor N2ORR. On the other hand, the triazenide system is relatively electron-poor, providing a "hard" tetracopper core that behaves like CuO/CeO<sub>2</sub> (that features hard oxide ligation) by forming a tightly-bound and electrophilic  $\mu$ -oxo intermediate upon  $N_2O$  deoxygenation to favor deN\_2O. These insights stand to inform future  $N_2O$ -fixing catalyst designs.

### Data availability

Spectral data is reproduced in ESI,<sup>†</sup> and raw data are available from the author upon request. Computational output coordinates have been uploaded as ESI.<sup>†</sup> X-ray crystallography data is available upon request from the CCDC under deposition numbers 2299540 and 2299541.

### Author contributions

N. P. M. carried out experiments, conducted computational modeling, and wrote the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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