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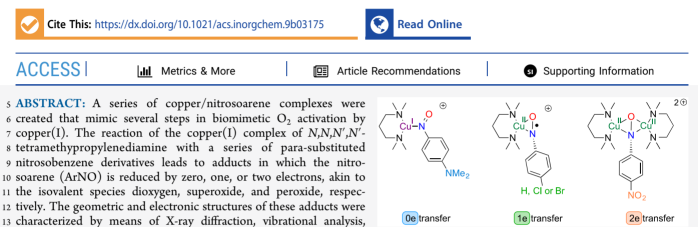
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Inorganic Chemistry

¹ Tuning Inner-Sphere Electron Transfer in a Series of Copper/ ² Nitrosoarene Adducts

3 Mohammad S. Askari, Farshid Effaty, Federica Gennarini, Maylis Orio, Nicolas Le Poul,* 4 and X. Ottenwaelder*



15 density functional theory (DFT) calculations. The bonding mode of the NO moiety depends on the oxidation state of the 16 ArNO moiety: κ N for ArNO, mononuclear η^2 -NO and dinuclear μ - η^2 : η^1 for ArNO^{•-}, and dinuclear μ - η^2 : η^2 for ArNO²⁻. ¹⁵N 17 isotopic labeling confirms the reduction state by measuring the NO stretching frequency (1392 cm⁻¹ for κ N-ArNO, 1226 cm⁻¹ for 18 η^2 -ArNO^{•-}, 1133 cm⁻¹ for dinuclear μ - η^2 : η^1 -ArNO^{•-}, and 875 cm⁻¹ for dinuclear μ - η^2 : η^2 for ArNO²⁻). The ¹⁵N NMR signal 19 disappears for the ArNO^{•-} species, establishing a unique diagnostic for the radical state. Electrochemical studies indicate reduction 20 waves that are consistent with one-electron reduction of the adducts and are compared with studies performed on Cu-O₂ analogues. 21 DFT calculations were undertaken to confirm our experimental findings, notably to establish the nature of the charge-transfer 22 transitions responsible for the intense green color of the complexes. In fine, this family of complexes is unique in that it walks 23 through three redox states of the ArNO moiety while keeping the metal and its supporting ligand the same. This work provides 24 snapshots of the reactivity of the toxic nitrosoarene molecules with the biologically relevant Cu(I) ion.

25 INTRODUCTION

26 The interaction of nitrosoarenes (ArNO) with metal centers 27 has drawn much attention because of its relevance to biological ²⁸ pathways¹⁻⁷ and catalytic C–N bond formation processes.^{8–12} 29 Chemists now have a good understanding of the geometric 30 structure of transition metal/nitrosoarene complexes.^{13,14} In 31 addition, ArNO species are redox-noninnocent ligands,^{15–17} 32 which portends a large landscape of electronic structures and 33 reactivity types upon interaction with redox-active metal ions. Because ArNO species are isovalent with O_2 , the reduction 34 $_{35}$ of ArNO by a transition metal is akin to the reduction of O_2 to $_{36}$ the superoxide ion (O2 $^{\bullet-}$, 1e reduction) or the peroxide ion $_{37}$ (O₂²⁻, 2e reduction). Therefore, metal/ArNO adducts are 38 often regarded as surrogates for metal/O₂ adducts. In 39 particular, and with relevance to the present paper, the 40 activation of O_2 by Cu(I) centers is paramount in the 41 biological world. This process fuels enzymes such as ⁴² dopamine- β -hydroxylase, tyrosinase, and particulate methane ⁴³ monooxygenase, to name but a few.^{18,19} This has inspired ⁴⁴ numerous biomimetic studies in which an electron-rich Cu(I)⁴⁵ species is reacted with O_2 .^{20–22} Without protection of the 46 protein backbone, however, the ensuing Cu/O_2 complexes are

14 ultraviolet-visible spectroscopy, NMR, electrochemistry, and

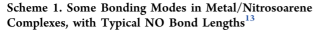
usually too oxidative to be stable above -60 °C. By contrast, ⁴⁷ Cu/ArNO adducts have been shown to have geometric and ⁴⁸ electronic structures very similar to those in Cu/O₂ adducts ⁴⁹ but were advantageously characterized at ambient temper- ⁵⁰ ature. ^{17,23–25} ⁵¹

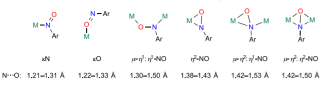
Owing to the asymmetric structure of ArNO in comparison $_{52}$ with that of O₂, the structural variety of metal/nitrosoarene $_{53}$ complexes exceeds that of metal/O₂ compounds. Some of the $_{54}$ main bonding modes of ArNO to metal ions are shown in $_{55}$ Scheme 1, 13,14 with the most common one being through the $_{56 \text{ s1}}$ N atom (κ N). The other bonding modes are thought to be $_{57}$ more prevalent when the ArNO moiety is reduced to the $_{58}$ mono- or dianion.

The NO bond length in metal/ArNO complexes depends on 60 the bonding mode, nature, and oxidation state of the metal and 61 the supporting ligands but alone is insufficient to characterize 62

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63 the degree of reduction of the ArNO moiety, as was already 64 shown with metal/O₂ adducts.²⁶ A few studies have scrutinized 65 the electronic structure on metal/ArNO complexes, partic-66 ularly the oxidation state of the ArNO moiety, by means of 67 techniques such as X-ray absorption spectroscopy or vibra-68 tional analysis with isotopic labeling (Scheme 2 for group 10 69 and 11 complexes). Their main conclusions are the following: 70 (i) In the majority of mononuclear κ N nitrosoarene 71 complexes, the NO bond length, 1.209–1.31 Å, shows little 72 or no elongation compared with that in free nitrosoar-73 enes,^{13,14,27–29} unless back-bonding from the metal becomes 74 significant.¹⁶ A radical character of the κ N-ArNO moiety, and 75 thus formally a 1.5 bond order, has been confirmed or inferred 76 in a few species (Scheme 2a).^{16,30,31}

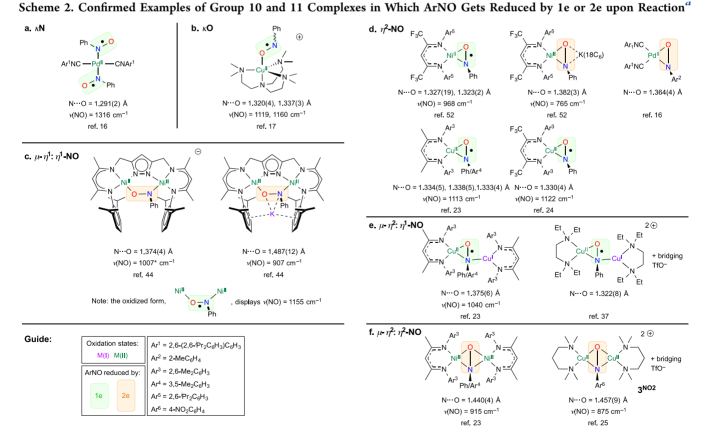
(ii) Most mononuclear κ O complexes with non-d¹⁰ r8 transition metals are structurally disordered, ^{13,14,32–36} and r9 conclusive statements about the extent of back-donation and 80 ArNO reduction cannot be made. By contrast, the non-81 disordered crystal structures of [(Me₆tren)Cu(κ O-PhNO)]X 82 (X = TfO⁻, SbF₆⁻; Scheme 2b) show significant NO bond 83 elongation. The radical character of the PhNO moiety (to an arylnitrosyl radical anion) was confirmed by magnetic ⁸⁴ measurements and vibrational and computational studies.^{17,37} ⁸⁵

(iii) Dinuclear μ - η^{1} : η^{1} (end-on) complexes present varying 86 degrees of ArNO reduction: by 0e (NO = 1.257-1.32 s7 Å),^{28,29,38-41} 1e (1.33-1.35 Å),⁴² and 2e (1.37-1.49 Å; 88 Scheme 2c).^{43,44}

(iv) In η^2 -NO complexes, the NO bond length (1.323–90 1.432 Å)^{16,23,31,45–52} is significantly longer than that in free 91 nitrosoarenes. 1e reduction of the ArNO moiety has been 92 confirmed in Cu and Ni complexes (Scheme 2d).^{23,52} Further 93 reduction of the Ni complex led to a doubly reduced PhNO^{2–}94 moiety.⁵² 2e reduction of the ArNO moiety was also confirmed 95 in a square-planar Pd(II) species upon reaction of a Pd⁰ species 96 with TolNO.¹⁶ 97

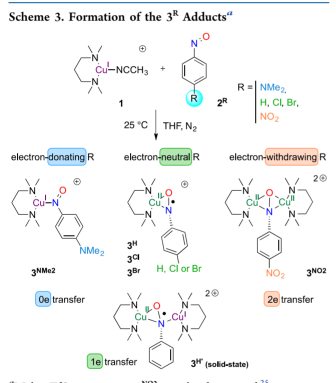
(v) Alongside several ArNO²⁻ examples (N–O = 1.40–1.53 98 Å),^{31,53–58} dinuclear μ - η^2 : η^1 complexes have been found in the 99 solid-state structures of Cu complexes with shorter NO bond 100 lengths (1.322–1.375 Å).^{23,37} Typically, the 1e-reduced 101 ArNO^{•-} moiety binds η^2 to a Cu(II) center and η^1 to a 102 Cu(I) center (Scheme 2e). These species are thought to be in 103 equilibrium with the mononuclear form [Cu^{II}(η^2 -ArNO^{•-})] in 104 solution.^{23,37}

(vi) Dinuclear μ - η^2 : η^2 complexes are quite rare, and only a 106 few examples with Rh,⁵⁷ Zr,⁵⁹ Hf,⁵⁹ Ni,²³ and Cu²⁵ are 107 reported in the literature. With an NO bond length in the 108 range of single bonds (1.422–1.500 Å), these complexes 109 possess a doubly reduced ArNO^{2–} moiety. In the case of the 110 Cu complex (Scheme 2f), this 2e reduction was made possible 111 by using a very electron-poor nitrosoarene bearing a *p*-NO₂ 112 substituent. More on this complex will be discussed below. 113



^aAsterisks indicate calculated values.

114 To summarize, 1e reduction of the ArNO moiety is usually 115 indicated by NO bond lengths in the range 1.29-1.37 Å and 116 NO stretching frequencies in the range 1000-1300 cm⁻ 117 (Scheme 2). Reduction by 2e is revealed by NO bond lengths 118 of 1.36-1.46 Å and NO stretching frequencies below 950 119 cm⁻¹. When no reduction occurs, the NO bonds are short 120 [1.261(4) and 1.268(4) Å for free PhNO] and the NO 121 stretching frequency is high (1506 cm⁻¹ for free PhNO), 122 although these values can be modified significantly when back-123 bonding is present.¹⁶ Last, 4e reduction of PhNO, with 124 complete NO bond cleavage, is possible with very electron-rich 125 metal complexes such as cobalt(I) β -diketiminate species.⁶⁰ Noting that these examples comprise different supporting 126 127 ligands and metal ions, we aimed at providing a systematic 128 study of the degree of inner-sphere ArNO reduction by using a 129 single Cu(I) precursor. Thus, in the present study, we report 130 on adducts 3^{R} (R = NMe₂, H, Cl, Br, NO₂) formed upon 131 intermolecular reaction of para-substituted nitrosobenzenes 2^{R} 132 with the Cu(I) complex of $N_1N_1N_1'$, N'-tetramethyl-1,3-133 propanediamine (TMPD), 1 (Scheme 3), for which analogous 134 Cu/O₂ chemistry is known.^{61–}



^{*a*}With a TfO⁻ counterion. 3^{NO2} was already reported.²⁵

135 **RESULTS AND DISCUSSION**

Synthesis and Crystallography. The slow addition of a 137 $[(MeCN)_4Cu](TfO)$ (MeCN = acetonitrile) solution to a 138 solution of TMPD and nitrosoarene, in a 1:1:1 ratio in 139 tetrahydrofuran (THF) at 25 °C (1:1:2 for R = NO₂), results 140 in the formation of deeply colored complexes that remain 141 stable under inert conditions. Crystallization of the complexes 142 by the slow diffusion of pentane into the reaction mixtures at 143 -30 °C afforded crystals suitable for X-ray diffraction analysis. 144 Several binding motifs consisting of mono- and dinuclear 145 complexes are obtained depending on the para substituent of 146 the nitrosoarene (Figure 1 and Table S1).

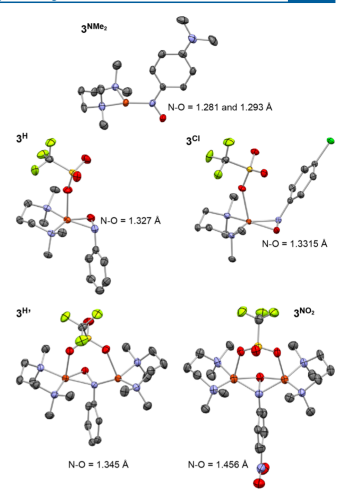


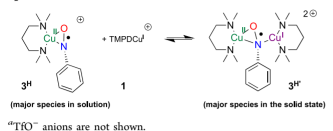
Figure 1. ORTEP at 50% probability of 3^{NMe_2} (one of two independent molecules), $3^{\rm H}$, $3^{\rm H\prime}$, $3^{\rm Cl}$, and $3^{\rm NO_2}$, with relevant N–O bond lengths. Uncoordinated TfO⁻ anions $(3^{NMe_2} \text{ and } 3^{\rm NO_2})$ and H atoms were omitted for clarity.

The structure of 3^{NMe_2} is that of a copper(I) arylnitroso 147 complex, i.e., mere κ N coordination of 1^{NMe_2} onto the [TMPD- 148 Cu^I]⁺ complex **1**. The N–O bond lengths in the two 149 independent molecules, 1.281 and 1.293 Å, are typical for 150 N=O double bonds. The trigonal-planar ligand field of Cu is 151 consistent with a Cu(I) oxidation state. In the solid state, the 152 species dimerizes via two weak Cu···O interactions (2.242 and 153 2.278 Å) between two crystallographically related 3^{NMe_2} 154 cations. This fact, coupled with the back-bonding of Cu into 155 the nitroso π^* orbital, could explain the slight elongation of the 156 N–O bond compared with a true double bond.^{16,37} 157

Two types of crystals were obtained in the same 158 crystallization pot with R = H. The minor component, of 159 green color, is the mononuclear [TMPDCu^{II}(η^2 -PhNO^{•-})- 160 (TfO)] species (3^H). This complex displays an η^2 -NO 161 coordination with an elongated N–O bond of 1.327 Å, 162 consistent with a 1.5 bond order.^{23,52} Cu sits in a square- 163 pyramidal environment with a TfO⁻ anion as a weak axial 164 ligand (Cu···O = 2.345 Å). The brown major component, 3^H, 165 of the formula [TMPDCu^{II}(μ - η^2 : η^1 -PhNO^{•-})(μ -TfO)- 166 Cu^ITMPD](TfO) also displays an elongated N–O bond 167 1.345 Å, consistent with a 1.5 bond order. One of the Cu 168 centers is bonded to both N and O of the PhNO moiety (Cu– 169 N = 2.019 Å; Cu–O = 1.853 Å), while the other is only 170 bonded to the N atom of PhNO (Cu–N = 1.904 Å; Cu···O = 171

¹⁷² 2.828 Å). Species $3^{H'}$ is therefore well described as formed ¹⁷³ from the association of the minor green component 3^{H} with ¹⁷⁴ one molecule of 1 (Scheme 4). Such architecture and

Scheme 4. Mononuclear/Dinuclear Equilibrium in 3^H Species^a



175 association were already described in the literature.^{23,37} 176 Because the mononuclear complex prevails in solution (the 177 solution is green, and a Job titration confirms a 1:1 178 stoichiometry; Figure S14), formation of the dinuclear 179 compound is an artifact of crystallization.

For 3^{Cl} , only green crystals of [TMPDCu^{II}(η^2 - $2^{\text{Cl}\bullet-}$)(TfO)] 181 were formed. The molecular structure is very similar to that of 182 3^{H} , with a 1.5 N–O bond order (1.3315 Å), except that the 183 TfO⁻ anion and the aromatic ring are on the same side of the 184 CuNO plane.

The complex with the most electron-poor ArNO moiety, 185 The complex with the most electron-poor ArNO moiety, 186 3^{NO_2} , was characterized in a previous communication.²⁵ It is a 187 dinuclear species of the formula [TMPDCu^{II}(μ - η^2 : η^2 -188 PhNO²⁻)(μ -TfO)Cu^{II}TMPD](TfO), where 2^{NO_2} is reduced 189 by 2e (N-O = 1.456 Å) and both Cu centers are in the 2+ 190 oxidation state.

Overall, the crystallographic study concludes on an increased degree of electron transfer from 1 to 2^{R} inasmuch as the *p*-R substituent is made more electron-poor: 0e in 3^{NMe_2} , 1e in 3^{H} / $3^{H'}$ and 3^{Cl} , and 2e in 3^{NO_2} . The lability of Cu complements the self-assembly process by allowing TfO⁻ or extra Cu(I) coordination when necessary.

IR Properties. Vibrational analysis by IR spectroscopy was sound the network of the network o

²⁰⁷ Comparing the IR properties of the organic precursors 2^{R} is ²⁰⁸ tentative because they have different structures in the solid ²⁰⁹ state: monomeric for 2^{NMe_2} , syn dimeric for 2^{H} , and anti ²¹⁰ dimeric for 2^{Br} (Tables 1 and S2 and Figures S1–S6). Still, the ²¹¹ correlation between the experimental and calculated spectra is ²¹² excellent, providing confidence that the calculations can enable ²¹³ us to locate the NO stretch accurately in the complexes.

t1

²¹⁴ Drastic changes in the NO stretching frequency are seen in ²¹⁵ 3^{R} complexes, consistent with NO bond weakening upon ²¹⁶ electron transfer (Tables 1 and S3 and Figures S7–S12). While ²¹⁷ the symmetry of the complexes is different and some ²¹⁸ complexes have multiple vibrational modes involving the NO ²¹⁹ stretch, the NO stretching energy decreases from 1315/1392 ²²⁰ cm⁻¹ for 3^{NMe_2} to 1226 cm⁻¹ for 3^{Br} to 875 cm⁻¹ for $3^{NO_2.25}$ ²²¹ This trend, supported by DFT calculations, is fully consistent

Table 1. NO Stretching Frequencies^a

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species	$ u \ (\Delta)/{ m cm}^{-1}$	species	$ u \ (\Delta)/\mathrm{cm}^{-1}$
2 ^{NMe₂}	1365 (12), 1340 (19)	3 ^{NMe₂}	1392 (14), 1315 (6)
2 ^{H b} 2 ^{Br d}	1388 (27) 1286 (4), 1256 (24)	3 ^{H/c} 3 ^{Br}	1162 (10), 1133 (23) 1226 (6)
$2^{NO_2 d}$	1238 (20)	3 ^{NO₂}	875 (15)

^{*a*}Measured at 25 °C on species labeled with ¹⁴N and ¹⁵N on the NO moiety. Full data are given in the Supporting Information. ^{*b*}Syn ArN(O)N(O)Ar dimer. ^{*c*}Contains a small amount of mononuclear species 3^{H} . ^{*d*}Anti ArN(O)N(O)Ar dimer.

with reduction of the bond order upon inner-sphere electron 222 transfer from the Cu center(s). For mixed-valent dinuclear 223 species $3^{H'}$, the NO stretch is lowered from 3^{Br} by about 70–224 100 cm⁻¹, consistent with the electron density being 225 delocalized onto the additional Cu(I) center. 226

NMR Properties. In CDCl₃, CD₂Cl₂, or acetone- d_6 ²²⁷ solutions, all 3^{R} species display diamagnetic ¹H and ¹³C ²²⁸ NMR spectra (Figures S37–S47). For 3^{H} , 3^{Cl} , and 3^{Br} , this ²²⁹ indicates a singlet ground state, as was observed for similar η^2 -²³⁰ ArNO complexes.^{23,24,37} By analogy with structurally similar ²³¹ η^2 -superoxocopper(II) species, this ground-state singlet is ²³² expected to be highly delocalized.⁶⁴ This situation also ²³³ contrasts with the end-on topology, where end-on ²³⁴ superoxocopper(II) complexes have a S = 1 ground ²³⁵ state, ^{65–68} as do Cu^{II}(κ O-ArNO^{•–}) complexes when Cu– ²³⁶ O–N–C_{Ar} is coplanar.¹⁷

The ¹⁵N NMR spectra of the ¹⁵N-labeled 3^{R} species are 238 most informative on the degree of electron transfer (Figure 2 239 f2t2

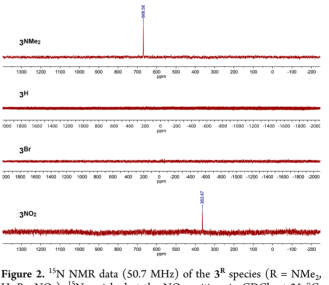


Figure 2. ¹⁵N NMR data (50.7 MHz) of the 3^{R} species (R = NMe₂, H, Br, NO₂), ¹⁵N-enriched at the NO position, in CDCl₃ at 25 °C. Species 3^{H} was made in situ by mixing equimolar amounts of TMPD, [Cu(MeCN)₄](TfO), and 2^{H} .

and Table 2). For comparison, the ¹⁵N NMR spectra of the ²⁴⁰ t2 ¹⁵N-labeled 2^{R} species reveal a logical downfield shift of the ²⁴¹ signal inasmuch as the R substituent becomes more electron-²⁴² poor. Cu(I) coordination on 2^{NMe_2} to form 3^{NMe_2} leads to an ²⁴³ upfield shift of the signal by 119 ppm, consistent with the ²⁴⁴ presence of a partial charge transfer from Cu(I) to the ArNO ²⁴⁵ moiety. On the other end of the series, the formation of 3^{NO_2} ²⁴⁶ leads to a dramatic upfield shift of the signal by 550 ppm, ²⁴⁷ consistent with the ArNO moiety being doubly reduced ²⁴⁸

Table 2. ¹⁵N NMR Data^a

	species	$\delta(^{15}\mathrm{N})/\mathrm{ppm}$	species	$\delta(^{15}\mathrm{N})/\mathrm{ppm}$
	2 ^{NMe2}	787.58	3 ^{NMe2}	668.58
	2 ^H	885.83	3 ^H	not observed
	2^{Br}	878.67	3 ^{Br}	not observed
	2 ^{NO2}	913.23	3 ^{NO2}	363.67
4	^a Measured in	CDCL at 25 °C on	a 500 MHz ii	$nstrument: \nu(^{15}N) =$

"Measured in CDCl₃ at 25 °C on a 500 MHz instrument; ν (¹⁵N) = 50.7 MHz.

249 (ArNO^{2–}) and therefore quite electron-rich. Interestingly, no 250 ¹⁵N signal was observed for 3^{H} and 3^{Br} under the same 251 recording conditions or with a wider acquisition window. This 252 behavior is consistent with the radical character of the ArNO^{•–} 253 moiety in these species. A small amount of triplet character 254 admixture in the ground-state singlet at room temperature 255 could lead to a paramagnetic shift of the ¹⁵N NMR resonance 256 outside the acquisition window (Fermi contact at ¹⁵N).⁶⁹ 257 Hence, the lack of a signal in a standard acquisition window 258 can be used as a local diagnostic of radical character on N. 259 Overall, the NMR data confirm, in solution, the assignments 260 that were made in the solid state.

²⁶¹ **Ultraviolet–Visible (UV–Vis) Absorption Properties.** ²⁶² The electronic structure of the complexes was probed by UV– ²⁶³ vis absorption spectroscopy (Figure 3 and Table 3).

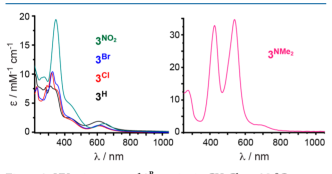


Figure 3. UV-vis spectra of 3^R species in CH₂Cl₂ at 25 °C.

Table 3. UV-Vis Data of 3^R Complexes^a

species		$\lambda_{\max} (\varepsilon)^{b}$		
3 ^{NMe₂}	426 (32.8)		538 (34.7)	690 (2.3)
3 ^H	313 (8.0), 338 (7.4)	440 (sh)		609 (1.9)
3 ^{CI}	319 (10.2), 350 (7.3)	440 (sh)		614 (1.1)
3 ^{Br}	325 (10.4), 350 (8.2)	440 (sh)		614 (1.2)
3 ^{NO₂}	345 (19.4)	440 (sh)		644 (1.5)
^{<i>a</i>} Measured in CH ₂ Cl ₂ at 25 °C. ^{<i>b</i>} λ_{max} /nm (ϵ /mM ⁻¹ cm ⁻¹).				

264 Complexes 3^{H} , 3^{Cl} , and 3^{Br} display sensibly the same UV– 265 vis spectrum, with an intense band around 330 nm and a less 266 intense feature around 610 nm. Compound 3^{NO^2} exhibits the 267 same spectral shape, but the 345 nm band is twice as intense. 268 The spectrum for complex 3^{NMe^2} is very different from the 269 other four spectra. It shows two very intense bands at 426 and 270 538 nm, while the weaker feature is red-shifted to 690 nm. 271 These absorptions will be analyzed in the next section.

272 **DFT Calculations.** DFT calculations have been undertaken 273 on the 3^{R} complexes to gain insight into the nature of the 274 species observed experimentally and to correlate their 275 electronic structures to the experimental data. The structures 276 of the 3^{R} species were subjected to geometry optimization, and their electronic properties were investigated. A good agreement 277 is found upon a comparison of the molecular geometries with 278 the X-ray crystallographic data (Figure S13). The calculated 279 NO bond lengths are 1.243 Å for 3^{NMe_2} , 1.287–1.288 Å for 3^H , 280 3^{Cl} , and 3^{Br} , 1.313 Å for $3^{H'}$, and 1.405 Å for 3^{NO_2} . While these 281 values are all underestimated (up to 0.05 Å), they lie within the 282 typical error range of DFT and provide a fair trend along the 283 series, being thus informative on the redox state of the ArNO 284 moiety. The DFT-optimized structures are very close to those 285 observed experimentally, with root-mean-square deviations of 286 0.543, 0.427, 0.407, 0.419, 0.444, and 0.338 from the crystal 287 molecular structures of 3^{NMe_2} , 3^H , 3^{Cl} , 3^{Br} , $3^{H'}$, and 3^{NO_2} , 288 respectively.

Time-dependent DFT (TD-DFT) calculations were per- 290 formed on the 3^R complexes, and the predicted spectroscopic 291 data provide calculated spectra that compare well with the 292 experimental observations (Tables S4-S8 and Figures S15- 293 \$20). Our computations support that the UV-vis spectra of 294 3^H, 3^{Cl}, 3^{Br}, and 3^{NO₂} are similar and dominated by two main 295 absorption bands of different intensities, while that of 3^{NMe2} 296 displays two intense electronic transitions. For the latter, the 297 band at 538 nm is assigned to a metal-to-ligand charge transfer 298 (MLCT), and the band at 426 nm is attributed to a ligand-to- 299 ligand charge transfer (LLCT). For both transitions, the 300 acceptor states mainly involve the nitroso moiety (Figure S16). 301 The electronic transitions for 3^{NO_2} were already analyzed.²⁵ 302 The 345 and 644 nm bands correspond to MLCT transitions 303 involving the μ - η^2 : η^2 -NO²⁻ moiety, in a very similar manner to 304 the transitions in the $(\mu$ - η^2 : η^2 -O₂²⁻)Cu^{II}₂ cores that mimic the 305 active sites of oxytyrosinase and oxyhemocyanin.²² For 3^H, 3^{Cl}, 306 and 3^{Br}, the absorptions near 320 nm are due to a combination 307 of MLCT and LLCT, with the acceptor state involving the NO 308 moiety, while the transitions in the visible around 610 nm 309 display a mixed character with similar contributions from the 310 metal and the nitroso moiety in both donor and acceptor states 311 (Figures S17–S19). Our TD-DFT calculations thus adequately 312 reproduce the energy of the key features of the experimental 313 spectra, which further support the geometries and electronic 314 properties of the 3^R complexes. Vibrational analysis also 315 confirmed the experimental observations (vide supra). $_{316}$ Consequently, 3^{NMe_2} can be described as a Cu(I) complex, $_{317}$ while 3^{H} , 3^{Cl} , and 3^{Br} are Cu^{II} -(ArNO^{•-}) species. The dimer ³¹⁸ $3^{H'}$ is assigned to Cu^{II} - Cu^{I} -(ArNO^{•-}) species, while 3^{NO_2} was ³¹⁹ previously shown to be a Cu(II)-Cu(II) complex with a ³²⁰ ArNO²⁻ moiety (2e-reduced ArNO). 321

Electrochemical Studies. Because this work aims at 322 tuning the redox properties by simple substitution, we studied 323 the electrochemical behavior of both precursors 2^{R} and 324 complexes 3^{R} for the whole series of R substituents (NMe₂, 325 H, Cl, Br, and NO₂). The goal was to correlate the 326 electrochemical properties with the reactivity (0e, 1e, or 2e 327 transfer) observed upon reaction with the [(TMPD)Cu¹]⁺ 328 complex 1 and the H-atom-transfer (HAT) reactivity of the 3^{R} 329 complexes (see below). The data were also compared to 330 existing records for analogous ArNO and O₂ complexes. Cyclic 331 voltammetry (CV) studies were performed at a glassy-carbon 332 working electrode in dry CH₂Cl₂ with 0.1 M NBu₄OTf as the 333 supporting electrolyte. In what follows, all potentials are 334 referenced to the Fc^{+/0} couple. 335

Substituted Nitrosoarenes, 2^{R} . CV studies of the free 336 nitrosoarenes, 2^{R} , led to the summary in Scheme 5. 2^{H} was first 337 s5 studied for a comparison with the literature.⁷⁰⁻⁷² When 338 scanned negatively, it displays two reversible responses at $E_{1/2}^{A}$ 339

f3t3



 $_{340} = -1.40$ V (process A) and $E_{1/2}^{B} = -1.86$ V versus Fc^{+/0} $_{341}$ (process B) (Figure S21 and Table 4). An irreversible

Table 4. Electrochemical	Data of the	e 2 ^R Nitrosoarenes ^a
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species	$E_{1/2}^{\mathrm{A}}(2^{\mathrm{R}})$	$E_{1/2}^{\mathrm{B}}(2^{\mathrm{R}})$	$\sigma_{ m para}^{\ \ b}$
2^{NMe_2}	$-1.69 (120)^{c}$	d	-0.83
2 ^H	-1.40 (90)	-1.86 (100)	0
2 ^{CI}	-1.32 (110)	-1.79 (90)	0.227
2^{Br}	-1.30 (90)	-1.75 (90)	0.232
2^{NO_2}	-0.93 (90)	$-1.33 (100)^{e}$	0.78

^{*a*}In CH₂Cl₂ containing 0.1 M NBu₄OTf at 25 °C (glassy-carbon working electrode); scan rate $v = 0.1 \text{ V s}^{-1}$, E/V versus $Fc^{+/0} (\Delta E_p/\text{mV})$. ^{*b*} σ_{para} Hammett parameters. ^{*c*}Determined at $v = 0.5 \text{ V s}^{-1}$. ^{*d*}Not determined. ^{*e*}An intermediate wave at -1.17 V was observed at faster scan rates.

342 oxidation peak is also detected at 0.63 V on the backscan 343 after reduction at -1.90 V (Figure S22). Variation of the scan 344 rate ν induces a significant modification of the redox behavior 345 (Figure S23), which is typical of two successive electron-346 transfer processes coupled to a chemical reaction, namely, an 347 ECE mechanism (E = electrochemical and C = chemical; 348 Scheme 5). In agreement with previous electrochemical 349 studies,^{70,71} process A corresponds to the monoelectronic 350 reduction of 2^H (Scheme 5), while process B corresponds to 1e 351 reduction of the azoxybenzene formed in situ by reaction of 352 the radical anion with residual water. This dimerization 353 hypothesis is supported by the ratio of the cathodic peak 354 currents, $i_{pc}^{B}/i_{pc}^{A} \approx 0.5$ (assuming similar diffusion coefficients). 355 In addition, the value of $E_{1/2}^{B}$ is in good agreement with the 356 standard potential values for the azoxy species in organic 357 solvents.^{70–72}

The processes described in Scheme 5 occur for 2^{R} with different para substituents (R = NMe₂, H, Cl, Br, NO₂) but at different redox potentials (Figure S21 and Table 4). Under the same experimental conditions, 2^{H} , 2^{Cl} , and 2^{Br} display almost at ca. $E_{1/2}^{A} = -1.3$ V and a second one at ca. $E_{1/2}^{B} = -1.8$ V. For second one at ca. $E_{1/2}^{A} = -1.8$ V. For second one at ca. $E_{1/2}^{A} = -1.8$ V. For second one at ca. $E_{1/2}^{B} = -1.8$ V.

Thus, a span of +760 mV is observed for $E_{1/2}^{A}$ upon NMe₂/ 370 NO₂ substitution, consistent with the electron-donating/ 371 withdrawing properties of the substituents. Fittingly, plots of 372 $E_{1/2}^{A}$ versus the σ_{para} Hammett parameter follow a linear trend, 373 indicating that the value of the redox potential is mainly 374 controlled by electronic effects (Figure S29).

³⁷⁵ [*Cu*(*TMPD*)(*ArNO*)](*OTf*) Complexes 3^{R} . CV studies of the ³⁷⁶ 3^{R} complexes (R = NMe₂, H, Br, NO₂) were performed under ³⁷⁷ the same experimental conditions as those for 2^{R} ligands ³⁷⁸ (Figure 4 and Table 5). Adding 2^{R} to a solution of 1 under CV ³⁷⁹ monitoring led to the same conclusions as those below (Figure ³⁸⁰ S25). All 3^{R} complexes display a first irreversible reduction

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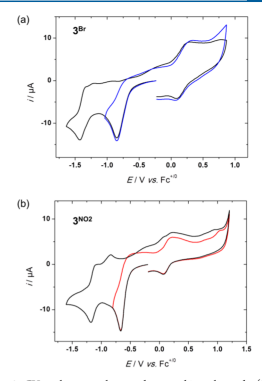


Figure 4. CV cycles at a glassy-carbon working electrode (E/V vs Fc^{+/0}; $\nu = 0.1 \text{ V s}^{-1}$) of 3^{R} (1.0 mM) in dry CH₂Cl₂ and 0.1 M NBu₄OTf: (a) R = Br; (b) R = NO₂.

Table 5. Electrochemical Data of 3^{R} Complexes^{*a*}

	$E_{\rm pc}^{\rm C}(3^{\rm R})$	$E_{1/2}^{\mathrm{D}}(3^{\mathbf{R}})$	$E_{1/2}^{\mathrm{E}}(3^{\mathrm{R}})$	$\sigma_{ m para}^{\ \ b}$
3 ^{NMe₂}	-1.05	-1.75	-0.36 (140)	-0.83
3 ^H	-0.92	-1.45^{c}	0.28	0.00
3 ^{Br}	-0.82	-1.22 (180)	0.08 (130)	0.23
3 ^{NO₂}	-0.72	-1.17 (145)	0.11 (120)	0.78

^{*a*}In CH₂Cl₂ containing 0.1 M NBu₄OTf at 25 °C; scan rate $\nu = 0.1$ V s⁻¹. *E*/V versus Fc^{+/0} ($\Delta E_p/mV$). ^{*b*} σ_{para} Hammett parameters. ^{*c*}Irreversible cathodic peak.

peak E_{pc}^{C} (process C) extending from -1.05 V for 3^{NMe_2} to $_{381}^{-0.72}$ V for 3^{NO_2} (Figure 4 and Table 5). As was the case with $_{382}^{-0.72}$ V for 3^{NO_2} (Figure 4 and Table 5). As was the case with $_{382}^{-0.72}$ v for 3^{RO_2} (Figure 4 and Table 5). As was the case with $_{382}^{-0.72}$ process A for the 2^{R} ligands, the redox potential is mainly $_{383}^{-0.72}$ controlled by electronic effects, which is confirmed by the $_{384}^{-0.72}$ linear variation of E_{pc}^{C} versus σ_{para} Hammett parameters (Figure $_{385}^{-0.72}$ S29). Whatever the nature of R, the first system remains $_{386}^{-0.72}$ irreversible at moderate scan rates ($\nu < 10 \text{ V s}^{-1}$; Figures 4 and $_{387}^{-0.72}$ S26 and S27a). This is indicative that a fast chemical reaction $_{388}^{-0.72}$ occurs upon electrochemical reduction. This EC mechanism $_{389}^{-0.72}$ versus log ν ($_{33}$ mV decade⁻¹; Figure S27b) and the constancy $_{391}^{-0.72}$ of the normalized current intensity ($t_{pc}^{C}\nu^{-1/2}$) with ν (inset $_{392}^{-0.72}$ Figure S27a), hence excluding an ECE process.

CV scanning until -1.8 V leads to the appearance of a 394 second system (process D) at $E^{\rm D}$ (-1.75 V $< E^{\rm D} < -1.21$ V), 395 which is quasi-reversible or irreversible, depending on R 396 (Figures 4 and S31). Increasing the scan rate induces a 397 decrease of the relative peak currents $i_{\rm pc}^{\rm C}$ and $i_{\rm pc}^{\rm D}$ (Figure S27c,d 398 for $3^{\rm NO_2}$), without modification of the peak potential values. 399 Altogether, this data set confirms that the chemical species that 400 is reduced reversibly through a simple electron transfer at $E^{\rm D}$ 401 originates from the first electrochemical reduction of the $3^{\rm R}$ 402 complex. As shown in Table 5, the potential value at $E^{\rm D}$ is 403

404 highly dependent on the substituting group R, meaning that 405 the chemical species or complex contains the ArNO moiety. 406 Possibly, reduction of the complex induces breaking of the 407 Cu–ArNO bond, liberating 2^{R} and explaining the similarity of 408 the E^{D} and E^{A} values. Such a hypothesis is verified in all cases 409 except for 3^{NO_2} (Figures S28 and S30).

410 Exhaustive electrolyses at E_{pc}^{C} and coulometric measurements 411 confirm that system C is a 1e process per mole of 3^{R} . For 412 example, electrochemical reduction of 3^{NMe_2} leads to its 413 disappearance, while a new wave appears at $E_{1/2}^{D}$ (Figure 414 S31), together with a significant color change of the solution 415 (purple to orange). A new system also appears in oxidation at 416 -0.2 , +0.45, and +0.65 V, suggesting a release of TMPD 417 (Figure S25a).

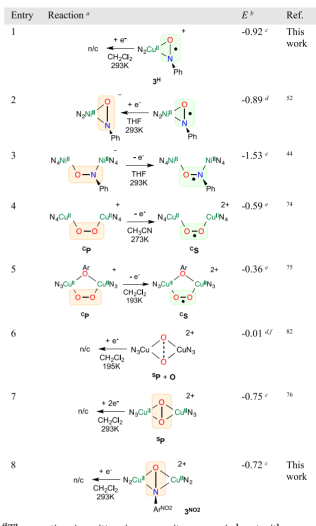
⁴¹⁸ In a general manner, 1e reduction of the 3^{R} complexes is ⁴¹⁹ accompanied by fast chemical processes that lead to partial ⁴²⁰ decomplexation and release of the TMPD ligand and/or 2^{R} . ⁴²¹ The transient electron-reduced species may thus be implicated ⁴²² in several reactions: radical dimerization and simple decoordi-⁴²³ nation, which themselves seem dependent on R.

On the oxidation side, a quasi-reversible system (process E) to the oxidation side, a quasi-reversible system (process E) to the system (process E)

The reduction data obtained for the 3^R complexes can be 430 431 compared with the few redox processes reported for $Ni_n/$ 432 ArNO and Cu_n/O_2 analogues (Table 6). The side-on 433 arylnitrosyl 3^H species (entry 1) gets reduced at a potential 434 similar to that of Warren's side-on arylnitrosylnickel(II) 435 complex.⁵² When ArNO binds in a 1,2-fashion (end-on) 436 between two Ni(II) centers, the potential for ArNO^{2-/•-} 437 conversion is decreased by ca. 650 mV (entry 3).⁴⁴ Comparing 438 ArNO with O₂ complexes would be interesting, but so far there 439 is no reported redox data for monocopper superoxo species 440 that would be similar in structure to 3^{H} . The exception is the 441 recent work by Reinaud et al., which showed by spectroelec-442 trochemistry that an in situ generated calix[6]amino-tren end-443 on superoxo complex could not be reduced above -0.90 V 444 versus Fc at -60 °C (113 K) in acetone.⁷³ On the other hand, 445 a few dicopper peroxo and superoxo species have been well 446 characterized by electrochemistry with the help of low-447 temperature approaches (entries 4-7). Here, the irreversible 448 1e reduction of 3^{R} is comparable to the monoelectronic and 449 reversible electron-exchange reactions detected for the end-on 450 superoxo/peroxo pyrazolate- and xylO-based complexes 451 (entries 4 and 5).^{74,75} Interestingly, the reduction potential $_{452}$ of 3^{NO_2} (entry 8) is close to that of Kodera's side-on 453 peroxodicopper(II) species (entry 7), although the latter is a 454 2e process.⁷⁶ Overall, using such comparisons to make 455 educated assignments of the electrochemical processes remains 456 tentative given the large variety of ligands, charge, nuclearity, 457 and bonding topology of the reported complexes. While data 458 for μ -hydroxodicopper complexes that are reminiscent of $Cu_n/$ 459 O₂ species is readily available, 77-82 comparisons with the 3^{R} 460 complexes would be even more tentative.

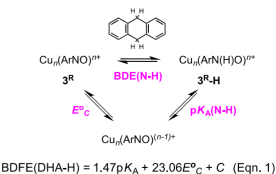
HAT Reactivity. We evaluated the reactivity of 3^{Cl} , 3^{Br} , and 462 3^{NO_2} for HAT reactivity (Scheme 6). In previous work, Warren 463 et al. reported a Ni^{II}-(η^2 -ArNO^{•-}) that converts into the 464 related Ni^{II}-(η^2 -ArN(H)O) complex (protonated hydroxyl-465 amine) upon reaction with 9,10-dihydroanthracene (DHA).⁵² 466 Conversely, Meyer et al. reported a dinuclear Ni^{II}₂-(μ - η^1 : η^1 - pubs.acs.org/IC

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^{*a*}The reaction is written in a way it was carried out either as an oxidation or as a reduction. Abbreviations: n/c, compound not characterized; ^CP, cis-end-on peroxo; ^CS, cis-end-on peroxo; ^SP, side-on peroxo; **O**, bis(μ -oxo). ^{*b*}Potential versus Fc^{+/0}. ^{*c*}Irreversible cathodic peak. ^{*d*}Reversible reduction. ^{*c*}Reversible oxidation. ^{*f*}Converted from the value versus saturated calomel electrode (SCE) using $E_{1/2}(Fc^{+/0}) = 560$ mV versus SCE in these conditions.

Scheme 6. Thermodynamic Analysis of HAT Reactivity



ArN(H)O) species that released its H atom to a phenoxyl 467 radical to form the related Ni^{II}_{2} -(μ - η^{1} : η^{1} -ArNO^{•-}) species and 468 evaluated a bond dissociation energy BDE(N–H) of around 469

s6

470 62–65 kcal mol⁻¹).⁴⁴ Following these examples, we reacted to 471 3^{Cl} , 3^{Br} , and 3^{NO_2} with DHA (bond dissociation free energy 472 BDFE = 76.0 mol⁻¹)⁸³ under UV–vis monitoring. A significant 473 decrease of the 3^{R} spectrum was observed upon the addition of 474 40 mol equiv of DHA in THF at 45 °C, which was corrected 475 for self-decomposition of the 3^{R} complexes at this temperature. 476 By analogy with the above examples, we presume that the 477 reaction yielded complexes of ArN(H)O, labeled 3^{R} -H 478 hereafter (Scheme 6), but their instability prevented further 479 analysis of the reaction and its mechanism.

⁴⁸⁰ The electron-withdrawing NO₂ group induces a faster ⁴⁸¹ oxidation of DHA, consistent with 3^{NO_2} being a stronger ⁴⁸² oxidant (higher $E_{1/2}^{C}$) than 3^{Br} and 3^{Cl} . The initial rates of ⁴⁸³ reaction depend on the R substituent: 0.029, 0.021, and 0.051 ⁴⁸⁴ \pm 0.005 mM min⁻¹ for 3^{Cl} , 3^{Br} , and 3^{NO_2} , respectively. Using ⁴⁸⁵ eq 1 (Scheme 6)⁸³ with a temperature correction, the value of ⁴⁸⁶ $E_{1/2}^{C}$ for 3^{R} taken as E_{pc}^{C} and using C = 66 kcal mol⁻¹ in THF,⁸⁴ ⁴⁸⁷ the pK_A value of the N–H bond in 3^{R} -H is evaluated around ⁴⁸⁸ 18 and 19.5 for the NO₂ and Br adducts, respectively, in order ⁴⁸⁹ to perform HAT from DHA. Similarly, 2^{NO_2} reacts, but slowly, ⁴⁹⁰ with 1,2-diphenylhydrazine (BDFE = 67.1 kcal mol⁻¹) at 25 ⁴⁹¹ °C. This brings the pK_A value to around 13, but this reaction is ⁴⁹² complicated by the byproduct azobenzene, which can interact ⁴⁹³ with Cu(I) and dissociate 2^{NO_2} . Further studies with different ⁴⁹⁴ substrates are necessary to decipher how nitrosarene ⁴⁹⁵ complexes perform this reaction, i.e., in a concerted or ⁴⁹⁶ sequential manner.^{85,86}

497 CONCLUSIONS

498 In summary, placing a synthetic handle at the para position of 499 nitrosoarenes enables control over the degree of electron 500 transfer from Cu(I) complexes, from 0e with electron-donating 501 substituents to 1e with electron-neutral substituents and 2e 502 with electron-poor substituents. As the Cu/ArNO adducts are 503 undergoing self-assembly, the geometric preferences of the Cu 504 center will prevail.³⁷ Thus, Cu(I) will be found in trigonal 505 geometries, with κ N-ArNO coordination, whereas a square-506 pyramidal Cu(II) will force η^2 -ArNO^{•-/2-} coordination. One $_{\rm 507}$ of the novel features of this work is the use of ^{15}N NMR as a 508 direct, local probe for the redox level of the ArNO moiety. 509 Thus, the absence of a ¹⁵N NMR signal coincides with the 510 radical state. A side effect of the self-assembly is, however, the 511 relative instability of the adducts upon external electron-512 transfer events. Notwithstanding, this series of complexes s13 provides structural snapshots of the isovalent Cu/O_2 514 chemistry, without the complication of thermal sensitivity of $_{515}$ Cu/O₂ species. It also enables redox studies to be performed, s16 although much remains to be done before a proper $ArNO/O_2$ 517 redox benchmark can be established. This series also highlights 518 the variety of intermediates that could occur during Cu-519 catalyzed ArNO transformations and suggests that, perhaps, 520 bond-forming events from ArNO precursors may proceed via 521 radical states.

522 **EXPERIMENTAL SECTION**

523 **Materials.** Chemicals were obtained from Sigma-Aldrich and Alfa 524 Aesar, except acetanilide-¹⁵N, which was purchased from Cambridge 525 Isotope Laboratories. Air-sensitive samples were handled under an 526 inert atmosphere (N₂) in a dry nitrogen glovebox (O₂ < 0.1 ppm; 527 H₂O < 0.1 ppm) or using standard Schlenk techniques. Solvents were 528 dried by standard procedures, degassed, and stored over 4 Å 529 molecular sieves in the glovebox. $N_iN_iN'_iN'$ -Tetramethyl-1,3-530 propanediamine (TMPD) was distilled over CaH₂ under nitrogen and stored in the glovebox. The copper salt [(MeCN)₄Cu](TfO) was 531 prepared by adapting the Kubas procedure using TfOH.⁸⁷ 4-532 Dimethylaminonitrosobenzene (2^{NMe_2}) ,^{88,89} 4-chloronitrosobenzene 533 (2^{Cl}) ,⁹⁰ 4-bromosonitrobenzene (2^{Br}) ,⁹⁰ and 4-nitrosonitrobenzene 534 $(2^{\text{NO}_2})^{91}$ were prepared via literature procedures. Isotopically labeled 535 ¹⁵N-4-nitrosonitrobenzene and [(TMPDCu)₂(μ -TfO)(μ - η^2 : η^2 -p-536 NO₂-C₆H₄NO)](TfO) were prepared following the procedure 537 reported earlier.²⁵ 4-Bromoaniline-¹⁵N was prepared from ¹⁵N-538 acetamide as reported in the Supporting Information.¹⁵N derivatives 539 of 2^H, 2^{Cl}, and 2^{Br} were prepared similarly to the ¹⁴N samples (see the 540 Supporting Information).⁹⁰

Characterization Methods. NMR spectra were recorded on a 542 Varian Innova-500 MHz instrument. All spectra were recorded in 543 CDCl₃ unless otherwise noted. ¹H and ¹³C NMR spectra were 544 referenced to internal tetramethylsilane. For 3^{R} species, the signal for 545 the TfO⁻ anion is not reported; it is observed at 119.5 ppm in 546 concentrated samples. ¹⁵N NMR spectra were referenced to external 547 formamide in dimethyl sulfoxide. IR spectra were recorded on a 548 Nicolet iS5 (Thermo Scientific) attenuated-total-reflectance instrument. UV–vis spectra were recorded on an Agilent 8453 550 spectrophotometer or a B&W Tek i-Trometer. Elemental analysis 551 was performed by the Laboratoire d'Analyze Élémentaire de 552 l'Université de Montréal. The presence of F atoms in the samples 553 interfered with the normal integration peak for H atoms. The value 554 for H is not necessarily trustworthy. 555

X-ray Crystallography. Crystallographic analysis was performed 556 on a Bruker APEX-DUO diffractometer. The frames were integrated 557 with the Bruker *SAINT* software package using a narrow-frame 558 algorithm. Data were corrected for absorption effects using the 559 multiscan method (*SADABS* or *TWINABS*). The structures were 560 solved by direct methods and refined using the *APEX3* software 561 package. All non-H atoms were refined with anisotropic thermal 562 parameters. H atoms were generated in idealized positions, riding on 563 the carrier atoms with isotropic thermal parameters. 564

Electrochemistry. Room temperature electrochemical studies of 565 the nitrosoarene ligands and their copper complexes were performed 566 in a glovebox (Jacomex; $O_2 < 1$ ppm and $H_2O < 1$ ppm) with a home- 567 designed three-electrode cell (WE, glassy carbon or platinum; RE, 568 platinum wire in a Fc⁺/Fc solution; CE, platinum or graphite rod). 569 Ferrocene was added at the end of the experiments to determine the 570 redox potential values. The potential of the cell was controlled by an 571 AUTOLAB PGSTAT 100 (Metrohm) potentiostat monitored by the 572 NOVA 1.11 software. Dichloromethane (Acros) was distilled over 573 CaH₂ under an inert atmosphere and stored in a glovebox. The 574 supporting salt NBu₄PF₆ was synthesized from NBu₄OH (Acros) and 575 HPF₆ (Aldrich). It was then purified, dried under vacuum for 48 h at 576 100 °C, and then kept under argon in the glovebox. NBu₄OTf 577 (Aldrich, 99%) was stored as received in the glovebox. Electrolytic 578 solutions were prepared in the glovebox and dried for a few days 579 under molecular sieves (3 Å) to remove traces of water before use. 580

Computational Details. All theoretical calculations were 581 performed with the ORCA program package.⁹² Full geometry 582 optimizations were carried out for all complexes using the generalized 583 gradient approximation functional BP86⁹³⁻⁹⁵ in combination with the 584 TZV/P⁹⁶ basis set for all atoms and by taking advantage of the 585 resolution of the identity (RI) approximation in the Split-RI-J 586 variant⁹⁷ with the appropriate Coulomb fitting sets.⁹⁸ Increased 587 integration grids (Grid4 in the ORCA convention) and tight self- 588 consistent-field convergence criteria were used. IR spectra were 589 obtained from numerical frequency calculations performed on DFT- 590 optimized structures. Isotope shift effects $({}^{14}N/{}^{15}N)$ were taken into 591 account using the orca vib utility program, and vibrational normal 592 modes were visualized with $Chemcraft^{99}$ software. Solvent effects were 593 accounted for according to the experimental conditions. For that 594 purpose, we used the CH_2Cl_2 (e = 9.08) solvent within the framework 595 of the conductor-like screening (COSMO) dielectric continuum 596 approach.¹⁰⁰ Single-point optical properties were predicted from 597 additional single-point calculations using the same functional/basis set 598 as that employed previously. Electronic transition energies and dipole 599 moments for all models were calculated using $\mathrm{TD}\text{-}\mathrm{DFT}^{101-103}$ within 600

601 the Tamm-Dancoff approximation.^{104,105} To increase the computa-602 tional efficiency, the RI approximation¹⁰⁶ was used to calculate the 603 Coulomb term. At least 40 excited states were calculated in each case, 604 and difference transition density plots were generated for each 605 transition. For each transition, difference density plots were generated 606 using the ORCA plot utility program and visualized with the 607 Chemcraft program. The same procedure was also employed to generate and visualize spin-density plots as well as molecular orbitals. 608 Synthetic Procedures. General Procedure for the Synthesis of 609 $_{610}$ 3^R Complexes (R = NMe₂, H, Cl, Br).²⁵ To a stirring solution of 611 TMPD (0.28 mmol, 1.1 equiv) and the corresponding nitrosobenzene 612 2^R (0.27 mmol, 1.05 equiv) in 5 mL of THF was added dropwise at 613 25 °C a solution of [(MeCN)₄Cu](TfO) (0.26 mmol, 1 equiv) in 2 614 mL of THF. The solution was stirred for 15 min and then cooled to 615 -30 °C. Dropwise addition of the solution to 15 mL of swirling 616 pentane previously cooled to -30 °C resulted in the precipitation of a 617 solid. The solid was isolated, washed with diethyl ether and pentane, 618 and dried in vacuo (yields typically 70-85%). Crystals suitable for X-619 ray structure determination were grown through the slow layered 620 diffusion of pentane into a concentrated solution of the complex in THF at -30 °C. 621

 $[(TMPDCu)(TfO)(\kappa N-p-NMe_2-C_6H_4NO)](TfO) \quad (\boldsymbol{3^{NMe_2}}). \text{ Yield: dark}$ 622 623 purple solid. ¹H NMR (500 MHz, CDCl₃): $\delta_{\rm ppm}$ 1.76 (m, 2H), 2.51 624 (s, 12H), 2.84 (m, 4H), 3.22 (s, 6H), 6.81 (br, 2H), 9.09 (very br, 625 2H). ¹³C NMR (125 MHz, CDCl₃): δ_{ppm} 22.89, 40.80, 48.69, 61.75, 626 112.2, 122.05, 156.26, 158.23. ¹⁵N NMR (50.7 MHz, CDCl₃): δ_{ppm} 627 668.58 (NO moiety). Anal. Calcd for C₁₆H₂₈CuF₃N₄O₄S: C, 38.98; 628 H, 5.72; N, 11.36; S, 6.50. Found: C, 37.86; H, 5.82; N, 11.13; S, 629 6.61

 $[(TMPDCu)_2(\mu-TfO)(\mu-\eta^2:\eta^1-PhNO)](TfO)$ (3^H). Yield: brown solid. 630 ₆₃₁ ¹H NMR (500 MHz, CDCl₃): δ_{ppm} 1.72 (br, 4H), 2.55 (br, 24H), 632 2.69 (br, 8H), 7.49 (t, 2H), 7.67 (t, 1H), 8.09 (d, 2H). ¹³C NMR 633 (125 MHz, CDCl₃): $\delta_{\rm ppm}$ 22.36, 48.59, 60.59, 120.97 (d upon 15 N 634 labeling, $J({}^{13}C-{}^{15}N) = 3$ Hz), 130.94 (d upon ${}^{15}N$ labeling, $_{635} J(^{13}C - ^{15}N) = 2 Hz)$, 131.78, 160.93. Anal. Calcd for 636 C₂₂H₄₁Cu₂F₆N₅O₇S₂: C, 33.33; H, 5.21; N, 8.83; S, 8.09. Found: 637 C, 31.28; H, 5.35; N, 8.37; S, 8.19 (precision is lacking because this 638 compound contains a minor quantity of 3^{H} in the solid state).

 $[(TMPDCu)(TfO)(\eta^2-PhNO)](TfO)$ (3^H). Yield: prepared in situ 639 640 (green solution). ¹H NMR (500 MHz, CDCl₃): $\delta_{\rm ppm}$ 1.69 (br, 2H), 641 2.47 (12H), 2.63 (4H), 7.43 (t, 2H), 7.63 (t, 1H), 7.97 (d, 2H). ¹³C 642 NMR (125 MHz, CDCl₃): $\delta_{\rm ppm}$ 20.37, 46.59, 59.24, 119.34 (d upon ¹⁵N labeling, $J(^{13}C-^{15}N) = 5$ Hz), 127.97 (d upon ¹⁵N labeling, 643 $_{644} J(^{13}C-^{15}N) = 2 Hz$, 130.78, 160.67 (d upon ¹⁵N labeling, 645 $J({}^{13}C-{}^{15}N) = 6$ Hz). ¹⁵N NMR (50.7 MHz, CDCl₃): not observed. $[(TMPDCu)(TfO)(\eta^2 - p - ClC_6H_4NO)](TfO)$ (**3**^{Cl}). Yield: green solid. ¹H 646 647 NMR (500 MHz, acetone- d_6): $\delta_{\rm ppm}$ 2.42 (br, 2H), 3.06 (s, 12H), 3.45 648 (br, 4H), 7.77 (d, 2H), 7.97 (d, 2H). ¹³C NMR (125 MHz, acetone-649 d₆): δ_{ppm} 20.17, 43.19, 55.50, 122.16, 126.10, 130.01, 166.47. Anal. 650 Calcd for C₁₄H₂₂ClCuF₃N₃O₄S: C, 34.71; H, 4.58; N, 8.67; S, 6.62. 651 Found: C, 34.27; H, 4.49; N, 8.16; S, 6.38.

 $[(TMPDCu)(TfO)(\eta^2-p-Br-C_6H_4NO)](TfO)$ (3^{Br}). Yield: green solid. 652 ₆₅₃ ¹H NMR (500 MHz, CDCl₃): $\delta_{\rm ppm}$ 1.74 (br, 2H), 2.65 (br, 12H), 654 2.78 (br, 4H), 7.54 (d, 2H), 7.91 (d, 1H). ¹³C NMR (125 MHz, 655 CDCl₃): δ_{ppm} 22.28, 48.52, 60.29, 122.81 (d upon ¹⁵N labeling, $(556 J(^{13}C-^{15}N)^{17} = 5 Hz)$, 126.01, 133.55 (d upon ¹⁵N labeling, $_{657} J(^{13}C-^{15}N) = 2.5 Hz)$, 160.59 (d upon ¹⁵N labeling, $J(^{13}C-^{15}N) =$ 658 5 Hz). ¹⁵N NMR (50.7 MHz, CDCl₃): not observed. Anal. Calcd for 659 C14H22BrCuF3N3O4S: C, 31.80; H, 4.19; N, 7.95; S, 6.06. Found: C, 660 31.29; H, 4.49; N, 7.81; S, 6.37

X-ray data for 3^{NMe_2} , 3^H , $3^{H\prime}$, and 3^{CI} are available as CCDC 661 662 1959040–1959043, respectively. Note that 3^{NO2} is CCDC 1029423.

ASSOCIATED CONTENT 663

664 Supporting Information

665 The Supporting Information is available free of charge at 666 https://pubs.acs.org/doi/10.1021/acs.inorgchem.9b03175.

Experimental supplements, including a crystallography 667 table, a Job plot, ¹⁵N labeling, IR data, electrochemistry 668 supplements, DFT data, and NMR spectra (PDF) 669

Accession Codes

CCDC 1959040-1959043 contain the supplementary crys- 671 tallographic data for this paper. These data can be obtained 672 free of charge via www.ccdc.cam.ac.uk/data request/cif. or by 673 emailing data request@ccdc.cam.ac.uk, or by contacting The 674 Cambridge Crystallographic Data Centre, 12 Union Road, 675 Cambridge CB2 1EZ, UK; fax: +44 1223 336033. 676

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