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# Simultaneous nitrosylation and N-nitrosation of a Ni-thiolate model complex of Ni-containing SOD†

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Nitric oxide (NO) is used as a substrate analogue/spectroscopic probe of metal sites that bind and activate oxygen and its derivatives. To assess the interaction of superoxide with the Ni center in Ni-containing superoxide dismutase (NiSOD), we studied the reaction of NO<sup>+</sup> and NO with the model complex, Et<sub>4</sub>N [Ni(nmp)(SPh-*o*-NH<sub>2</sub>-*p*-CF<sub>3</sub>)] (**1**; nmp<sup>2-</sup> = dianion of *N*-(2-mercaptoethyl)picolinamide; <sup>-</sup>SPh-*o*-NH<sub>2</sub>-*p*-CF<sub>3</sub> = 2-amino-4-(trifluoromethyl)benzenethiolate) and its oxidized analogue **1**<sup>ox</sup>, respectively. The ultimate products of these reactions are the disulfide of <sup>-</sup>SPh-*o*-NH<sub>2</sub>-*p*-CF<sub>3</sub> and the *S,S*-bridged tetrameric complex [Ni<sub>4</sub>(nmp)<sub>4</sub>], a result of S-based redox activity. However, introduction of NO to **1** affords the green dimeric {NiNO}<sup>10</sup> complex (Et<sub>4</sub>N)<sub>2</sub>{Ni(k<sup>2</sup>-SPh-*o*-NNO-*p*-CF<sub>3</sub>(NO))<sub>2</sub>} (**2**) *via* NO-induced loss of nmp<sup>2-</sup> as the disulfide and N-nitrosation of the aromatic thiolate. Complex **2** was characterized by X-ray crystallography and several spectroscopies. These measurements are in-line with other tetrahedral complexes in the {NiNO}<sup>10</sup> classification. In contrast to the established stability of this metal-nitrosyl class, the Ni–NO bond of **2** is labile and release of NO from this unit was quantified by trapping the NO with a Co<sup>II</sup>-porphyrin (70–80% yield). In the process, the Ni ends up coordinated by two *o*-nitrosaminobenzenethiolato ligands to result in the structurally characterized *trans*-(Et<sub>4</sub>N)<sub>2</sub>[Ni(SPh-*o*-NNO-*p*-CF<sub>3</sub>)<sub>2</sub>] (**3**), likely by a disproportionation mechanism. The isolation and characterization of **2** and **3** suggest that: (i) the strongly donating thiolates dominate the electronic structure of Ni-nitrosyls that result in less covalent Ni–NO bonds, and (ii) superoxide undergoes disproportionation *via* an outer-sphere mechanism in NiSOD as complexes in the {NiNO}<sup>9/8</sup> state have yet to be isolated.

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## Introduction

Nitric oxide (NO) and its derivatives (termed reactive nitrogen species or RNS) play a vital role in a variety of mammalian (and in some cases bacterial) physiological and pathological processes.<sup>1–4</sup> Additionally, this gaseous free radical has applications in fundamental research, especially in bioinorganic chemistry, where it is utilized as a structural/spectroscopic probe of O<sub>2</sub> (and other reactive oxygen species, *e.g.*, O<sub>2</sub><sup>•-</sup> and H<sub>2</sub>O<sub>2</sub>) binding/activating metalloenzymes.<sup>5–9</sup> In general, this approach is employed because metal-nitrosyl (MNO) bonds are

highly covalent, and hence more stable, than metal-dioxygen (M–O<sub>2</sub>) adducts.<sup>10</sup> The use of NO as an O<sub>2</sub> analogue is based on similar electronic structures between these diatoms and their reduced derivatives.<sup>3</sup> For example, <sup>3</sup>NO<sup>-</sup> (termed the nitroxyl anion), the one electron reduced analogue of NO, is isoelectronic with O<sub>2</sub> with two unpaired π\* electrons in the HOMO. Additionally, NO, while not isoelectronic with O<sub>2</sub><sup>•-</sup>, has the same ground state electronic structure with a singly occupied π\* MO. Thus, NO interactions with the active sites of O<sub>2</sub>-activating/ROS-breakdown enzymes report coordination (inner-sphere substrate binding) and the extent of substrate bond activation from vibrational spectroscopic measurements of the N–O and M–NO stretching frequencies.

Since 2009, our lab has designed and constructed numerous low molecular weight models of the active site of Ni-containing superoxide dismutase (NiSOD).<sup>11–18</sup> NiSOD is an unprecedented SOD due to Ni<sup>III/II</sup>-coordination to cysteinato-S (CysS) and peptido-N donors (Chart 1), the former of which is susceptible to oxidative modification by the substrate (O<sub>2</sub><sup>•-</sup>) and products (O<sub>2</sub> and H<sub>2</sub>O<sub>2</sub>) of the SOD catalyzed reaction.<sup>19,20</sup> Few models employ ligands with the correct spatial disposition and

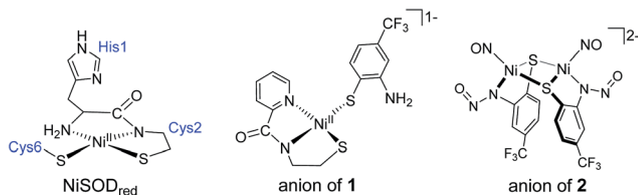
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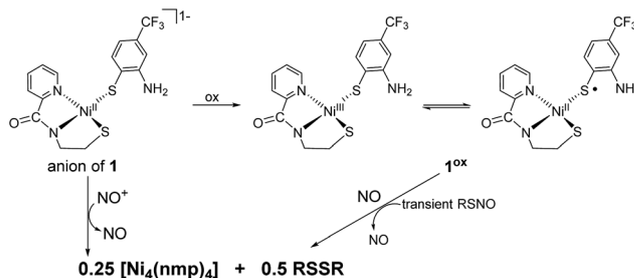
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**Chart 1** Structures of the active site of NiSOD<sub>red</sub> (left; His1 coordinates to Ni<sup>III</sup> in NiSOD<sub>ox</sub>), the anion of the NiSOD model complex Et<sub>4</sub>N[Ni(nmp)(SPh-*o*-NH<sub>2</sub>-*p*-CF<sub>3</sub>)] (1) (center; nmp<sup>2-</sup> = dianion of the N<sub>2</sub>S ligand *N*-(2-mercaptoethyl)picolinamide), and the anion of the {NiNO}<sup>10</sup> complex (Et<sub>4</sub>N)<sub>2</sub>{Ni(κ<sup>2</sup>-SPh-*o*-NNO-*p*-CF<sub>3</sub>)(NO)}<sub>2</sub> (2).



**Scheme 1** NO and NO<sup>+</sup> reactions of 1 and 1<sup>ox</sup>, respectively. R = 2-amino-4-(trifluoromethyl)benzenethiolate.

electronic nature of the unique N<sub>3</sub>S<sub>2</sub> donor set found in the active site.<sup>21–23</sup> Moreover, fewer report reversible electrochemical and/or spectroscopic evidence for the Ni<sup>III</sup> oxidation state due to redox associated with the coordinated thiolates. One model from our lab, namely Et<sub>4</sub>N[Ni(nmp)(SPh-*o*-NH<sub>2</sub>-*p*-CF<sub>3</sub>)] (1; nmp<sup>2-</sup> = dianion of the N<sub>2</sub>S ligand *N*-(2-mercaptoethyl)picolinamide; see Chart 1) displays a reversible redox-event at –0.43 V (vs. Fc/Fc<sup>+</sup> in DMF) that, based on EPR, UV-vis, MCD, and DFT computations, represents the electrochemical conversion from Ni<sup>II</sup> in 1 to a Ni<sup>II</sup>-thiyl ↔ Ni<sup>III</sup>-thiolate resonance species termed 1<sup>ox</sup>.<sup>16</sup> Because substrate binding to Ni in NiSOD has not been defined, although most reports favor an outer-sphere mechanism,<sup>15,24</sup> we were curious to use NO as an O<sub>2</sub><sup>•-</sup> probe to define potential intermediates that may be traversed in the NiSOD mechanism. We report here, for the first time, the reactions and product characterization of NO (and NO<sup>+</sup>) with 1 and the well-defined analogue of NiSOD<sub>ox</sub> (1<sup>ox</sup>). NO/NO<sup>+</sup> oxidize the aromatic thiolate ligand in 1<sup>ox</sup> and 1, respectively. However, introduction of NO to 1 affords the green dimeric {NiNO}<sup>10</sup> complex (Et<sub>4</sub>N)<sub>2</sub>{Ni(κ<sup>2</sup>-SPh-*o*-NNO-*p*-CF<sub>3</sub>)(NO)}<sub>2</sub> (2) *via* NO-induced loss of nmp<sup>2-</sup> as the disulfide and N-nitrosation of the aromatic thiolate (Chart 1). While 2 bears little resemblance to NiSOD, its formation indicates how reactive NiSOD models such as 1 are in the presence of redox-active diatoms and suggest similar paths for other biological Ni-thiolate sites. Additionally, 2 contains a labile Ni–NO bond, a new feature for the {NiNO}<sup>10</sup> formulation that appears to be controlled by the presence of the thiolate ligands. We describe the synthesis, spectroscopy, electronic structure, reactivity and mechanistic insight into the formation of the Ni-nitrosyl in this account.

## Results and discussion

In the anticipation of isolating a Ni-nitrosyl as an analogue of a potential Ni-superoxo/peroxo catalytic intermediate of NiSOD, we examined the reaction of 1 with NOBF<sub>4</sub> and *in situ* prepared 1<sup>ox</sup> with NO (Scheme 1). In theory, both reactions yield the same product. For example, nitrosonium (NO<sup>+</sup>; a strong oxidant, *E* = +0.56 V vs. Fc/Fc<sup>+</sup> in DMF<sup>25</sup>) will oxidize 1 to 1<sup>ox</sup> and form NO in the process. The newly generated 1<sup>ox</sup> (*S* = 1/2) then reacts with NO to form the Ni-nitrosyl, formally a {NiNO}<sup>8</sup> complex, assuming binding of NO and no other coordination sphere

changes, using the notation defined by Enemark and Feltham.<sup>26</sup> Likewise, NO will readily intercept paramagnetic 1<sup>ox</sup> to generate the same species. Mixing a DMF solution of 1 with NOBF<sub>4</sub> (1 : 1) resulted in instantaneous bleaching of the solution, consistent with oxidation of the RS<sup>-</sup> ligand to disulfide (RSSR), and the appearance of a dark-red precipitate that was spectroscopically identified to be the neutral *S,S*-bridged tetramer [Ni<sub>4</sub>(nmp)<sub>4</sub>] (Scheme 1).<sup>16</sup> This outcome is typical for all [Ni(nmp)(SR)]<sup>-</sup> complexes when treated with chemical oxidants, *i.e.*, *S*-oxidation of the coordinated monodentate thiolate to RSSR.<sup>15</sup> Incidentally, the same result was obtained when introducing NO(g) into a DMF solution of *in situ* generated 1<sup>ox</sup>. In this case, formation of the disulfide may traverse a fleeting, and yet to be characterized, RSNO intermediate that releases NO *via* homolytic cleavage of the RS–NO bond (Scheme 1).<sup>27</sup> Overall, a Ni-nitrosyl was not isolated. This result may not be too surprising considering that all known Ni-nitrosyls are in the {NiNO}<sup>10</sup> Enemark–Feltham (EF) classification,<sup>28</sup> although a {NiNO}<sup>9/8</sup> species is not entirely unrealistic in light of the strong donors present in 1 and in NiSOD, *i.e.*, peptido-*N* and alkyl-thiolato-*S*.

As a control, we also explored the reaction of Ni<sup>II</sup> complex 1 with NO. In general, NO does not react with square-planar [Ni(nmp)(SR)]<sup>-</sup> (R = simple aryl or alkyl groups) complexes due to their diamagnetic nature. However, when R contains a potentially bidentate chelate, as in 1, a different course takes place. For instance, exposing a DMF solution of 1 with NO(g) for 30 s resulted in a gradual change of the solution from dark-red to green over several minutes. Workup of this reaction indicated a Ni-nitrosyl based on the strong double-humped peak in the N–O stretching ( $\nu_{\text{NO}}$ ) region of the IR spectrum (*vide infra*). Subsequent crystallization of the bulk material from MeCN/Et<sub>2</sub>O at –20 °C resulted in green crystals of a dinuclear thiolate-bridged {NiNO}<sup>10</sup> complex (Et<sub>4</sub>N)<sub>2</sub>{Ni(κ<sup>2</sup>-SPh-*o*-NNO-*p*-CF<sub>3</sub>)(NO)}<sub>2</sub> (2) as depicted in Fig. 1 with selected metric parameters listed in Table 1. The Ni centers in 2 are distorted tetrahedral ( $\tau_4 = 0.73$  (ref. 29)) resulting from N<sub>2</sub>S<sub>2</sub> coordination of the thiolato-*S*/deprotonated amine-*N* of the *S*-bridged *o*-nitrosaminobenzenethiolate and a terminal nitrosyl. To our knowledge, complex 2 represents the first example of a structurally characterized first-row metal complex with both a coordinated nitrosyl and amine-*N*-bound nitrosamine. In accord with other tetrahedral {NiNO}<sup>10</sup> complexes,<sup>28</sup> the Ni–N(O)



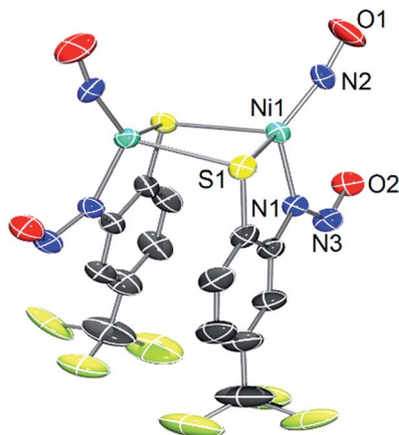


Fig. 1 X-ray structure of the anionic portion of **2** with the atom labeling scheme (50% thermal probability). H atoms and  $\text{Et}_4\text{N}^+$  counterions are omitted for clarity.

Table 1 Selected bond distances (Å) and bond angles (deg) from the X-ray crystal structure of **2**, compared with the DFT-optimized model **2**\*

	X-ray structure <b>2</b>	DFT (BP86/def2-TZVPP) optimized structure <b>2</b> *
Ni1–S1	2.3169(7)	2.294
Ni1–S1'	2.3555(6)	2.344
Ni1–N2	1.659(7)	1.648
Ni1–N1	1.971(2)	1.974
N2–O1	1.182(8)	1.191
N1–N3	1.299(3)	1.324
N3–O2	1.269(3)	1.254
S1–Ni–S1'	96.66(2)	90.05
S1–Ni–N1	86.21(6)	86.67
S1'–Ni–N1	99.10(6)	98.40
S1–Ni–N1	123.8(5)	125.92
N2–Ni–S1'	109.8(6)	113.43
N1–Ni–N2	133.6(5)	132.15
Ni1–N2–O1	167.8(12)	171.42
N1–N3–O2	114.2(2)	115.80
$\tau_4$	0.73	0.72

distance is short (1.659 Å), the N–O bond (1.182 Å) is intermediate between free  $\text{NO}^+$  (1.15 Å) and  $^1\text{HNO}$  (1.21 Å),<sup>10</sup> and the Ni–N–O bond angle is close to linear albeit slightly bent (167.8°) (see Table 1). Complex **2** is analogous to the limited number of four-coordinate/S-bound Ni-nitrosyls,<sup>30–34</sup> fewer of which contain Ni–S<sub>thiolate</sub> bonds<sup>31,33</sup> which display Ni–N(O) (1.663–1.683 Å), N–O (1.131–1.173 Å), and Ni–N–O (156.6–173.9°) distances/angles in similar ranges. Even neutral/cationic P-<sup>35–40</sup> and N-bound<sup>34,41–43</sup>  $\text{L}_3\text{Ni–NO}/\text{L}_2\text{XNi–NO}$  complexes exhibit similar metric parameters. The coordinated nitrosamine is bent (N–N–O: 115.4°), *i.e.*,  $\text{sp}^2$ -hybridized nitroso-N, with N–N and N–O distances of 1.299 and 1.269 Å, respectively. These values suggest a small degree of delocalization in the R–N–N–O unit. However, the structure is more biased towards the nitrosamino R–N<sup>–</sup>–N=O *versus* diazoate R–N=N–O<sup>–</sup> resonance form. To compare, the structure of *syn*-methanediazoate (N–N: 1.246 Å,

N–O: 1.306 Å) reflects the true double bond character in an authentic R–N=N–O unit.<sup>44</sup> These values are somewhat comparable to other N-bound nitrosamine complexes,<sup>45,46</sup> especially  $[\text{CpNi}(\text{PPh}_3)(\text{ON}_2\text{Ph-}p\text{-NO}_2)]$  (**1**)<sup>47</sup> (N–N: 1.327 Å, N–O: 1.249 Å, N–N–O: 113.1°). Structures of coordinated nitroso-N-metal complexes (*vs.* amine-N as in **2**) also afford similar structural parameters in the R<sub>2</sub>NO.<sup>48</sup> In contrast, O-bound nitrosamine complexes appear to favor more of a resonance delocalized structure as the N–N (1.275–1.288 Å) and N–O (1.251–1.275 Å) distances in a series of  $[\text{Fe}^{\text{III}}(\text{P})(\text{ONNR}_2)_2]^+$  (P = porphyrin) complexes are nearly identical and result in a single <sup>15</sup>N-sensitive peak in the IR due to overlapping  $\nu_{\text{NN}}/\nu_{\text{NO}}$  modes.<sup>49–51</sup>

Complex **2** was characterized by a variety of spectroscopic methods. The solid-state IR spectrum (KBr matrix) of **2** exhibits two closely spaced, but well-resolved,  $\nu_{\text{NO}}$  at 1759 and 1743  $\text{cm}^{-1}$  (1724, 1708  $\text{cm}^{-1}$  for **2**-<sup>15</sup>NO;  $\Delta\nu_{\text{NO}}$ : 35  $\text{cm}^{-1}$ ; see Fig. 2). These values fall in the range of known tetrahedral, neutral, and anionic  $\{\text{NiNO}\}^{10}$  complexes.<sup>28</sup> Because **2** is of  $C_2$  symmetry (*cis* NO, *syn* bridging thiolates), two IR-active N–O vibrational modes are expected. The other feasible isomer of **2** would be of  $C_i$  symmetry (*trans* NO, *anti* bridging thiolates) and would display one IR-active N–O stretch. Indeed, the IR spectrum of **2** in DMSO exhibits one  $\nu_{\text{NO}}$  at 1784  $\text{cm}^{-1}$  suggesting possible *cis/trans*-NO conversion in solution (or an averaged  $\nu_{\text{NO}}$  value due to rapid tumbling) or thiolate-bridge splitting to yield a four-

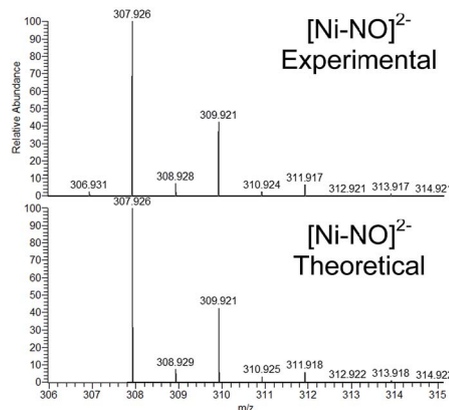
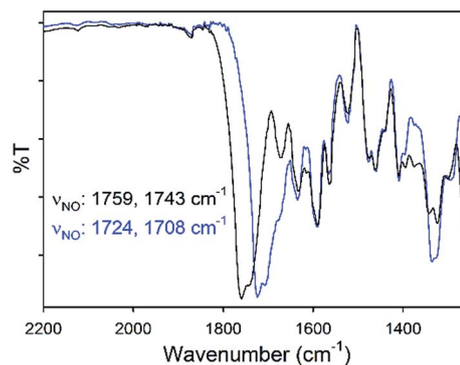


Fig. 2 (Top) Solid-state IR of the  $\nu_{\text{NO}}$  region for **2** (black) and **2**-<sup>15</sup>NO (blue) in a KBr matrix. (Bottom) High resolution ESI-MS(–) of **2** with the theoretical isotopic distribution.



coordinate mononuclear  $\{\text{NiNO}\}^{10}$  with DMSO as the fourth ligand, *i.e.*,  $[\text{Ni}(\kappa^2\text{-SPh-}o\text{-NNO-}p\text{-CF}_3)(\text{DMSO})(\text{NO})]^-$ . The  $^1\text{H}$  NMR spectrum of **2** in  $\text{CD}_3\text{CN}$  (Fig. S6†) or  $\text{DMSO-}d_6$  (not shown) are similar and thus do not distinguish any of the proposed structures. Comparable IR spectral changes in the opposite direction are observed for the one other thiolate-supported anionic dinuclear  $\{\text{NiNO}\}^{10}$ ,  $(\text{Et}_4\text{N})_2[\text{Ni}_2(\text{NO})_2(\mu\text{-SPh})_2(\text{SPh})_2]$  (**II**), with *trans* NO ligands ( $\nu_{\text{NO}}$ : 1709  $\text{cm}^{-1}$  in KBr; 1751, 1721  $\text{cm}^{-1}$  in THF).<sup>33</sup> A similar situation is described for a pyrazolate-bridged anionic dinuclear  $\{\text{NiNO}\}^{10}$  complex.<sup>52</sup> IR peaks arising from the nitrosamine were not as obvious due to multiple overlapping peaks in the region (Fig. S5†). However,  $^{15}\text{N}$ -sensitive peaks in the IR of **2** at 1342 and 1258  $\text{cm}^{-1}$  (1326, 1249  $\text{cm}^{-1}$  in **2- $^{15}\text{NO}$** ) are assigned as  $\nu_{\text{NO}}$  and  $\nu_{\text{NN}}$ , respectively. In comparison, a series of secondary nitrosamines display  $\nu_{\text{NO}}$ : 1428–1463  $\text{cm}^{-1}$  and  $\nu_{\text{NN}}$ : 1035–1154  $\text{cm}^{-1}$  in  $\text{CCl}_4$ .<sup>53</sup> Therefore, a significant degree of delocalization occurs in the RNNO unit of **2** to cause the corresponding downshift in  $\nu_{\text{NO}}$ /upshift in  $\nu_{\text{NN}}$ . While no paramagnetically shifted resonances are observed in the  $^1\text{H}$  NMR ( $\text{CD}_3\text{CN}$ ) of **2**, several species are indicated in freshly prepared solutions (Fig. S6†) that are likely caused by the lability of the Ni–NO bond and presence of  $\text{nmpS}_2$  (*vide infra*). The  $^{15}\text{N}$  NMR spectrum of **2- $^{15}\text{NO}$**  confirms multiple solution speciation with four major peaks in the range for nitrosamines and linearly coordinated NO ( $\delta$ : 40–190 ppm in  $\text{CD}_3\text{CN}$ , *vs.*  $\text{CH}_3\text{NO}_2$ , Fig. S7†).<sup>54–57</sup> Moreover, the  $^1\text{H}$  NMR of thiolate-bridged dinuclear complex **II** displays broadened aryl-H resonances caused by rapid exchange of  $\text{PhS}^-$  ligands because of disproportionation to the mononuclear  $(\text{Et}_4\text{N})_2[\text{Ni}(\text{NO})(\text{SPh})_3]$  (**III**) and an uncharacterized  $[\text{Ni}(\text{NO})(\text{SPh})]$  species.<sup>33</sup> However, high-resolution electrospray ionization mass spectrometry (HR-ESI-MS; negative mode) displays one dominant compound with the formula and isotopic distribution consistent with the dianionic portion of **2** ( $m/z$ : 307.926,  $z = 2$ , Fig. 2, S8 and S9†) and **2- $^{15}\text{NO}$**  ( $m/z$ : 309.920,  $z = 2$ ; Fig. S10 and S11†), although this measurement does not discriminate against *cis* and *trans* NO conformers. Another minor peak in the HR-ESI-MS(–) is centered at  $m/z$ : 248.960 ( $z = 2$ ; Fig. S8†) that suggests a new  $[\text{Ni}(\text{N}_2\text{S}_2)]^{2-}$  species through loss of the Ni-coordinated NO and one Ni (*vide infra*).

Solutions of **2**, especially in donor solvents such as MeCN or DMF, gradually lose their green color to give red-brown solutions more reminiscent of square-planar  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  complexes.<sup>15,58</sup> Even freshly prepared  $\text{CD}_3\text{CN}$  solutions of **2** exhibit multiple peaks in the  $^1\text{H}/^{15}\text{N}$  NMR, and ESI-MS shows a new species with a Ni isotope pattern at  $m/z \sim 249$  (*vide supra*). This change is enhanced when vacuum is applied and FTIR spectra of these reaction mixtures lack any  $\nu_{\text{NO}}$  suggesting the loss of coordinated NO from **2** to generate a new Ni species. Slow diffusion of  $\text{Et}_2\text{O}$  into MeCN solutions of **2** that have been left standing for several weeks result in crystals (10–20% isolated yield from crystallization) of a square-planar ( $\tau_4 = 0.12$ )  $\text{Ni}^{\text{II}}$  compound where two N,S-chelating *o*-nitrosaminobenzenethiolato ligands bind to Ni in a *trans* configuration, *viz.* *trans*-( $\text{Et}_4\text{N}$ )<sub>2</sub>[Ni(SPh-*o*-NNO-*p*-CF<sub>3</sub>)<sub>2</sub>] (**3**) (Fig. 3). The bond lengths (Ni–S: 2.2072 Å, Ni–N: 1.896 Å) and angles (Table S3†) are similar to other planar  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  complexes that

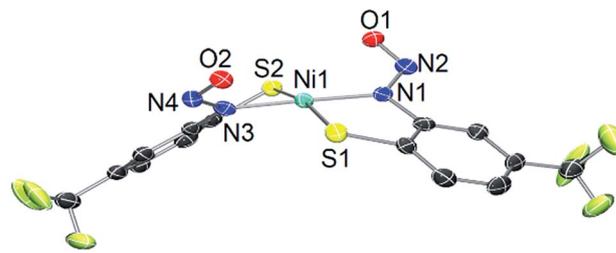


Fig. 3 X-ray structure of the dianion of **3** with the atom-labeling scheme (50% thermal probability). One of two crystallographically distinct molecules shown. H atoms,  $\text{Et}_4\text{N}^+$  counterions, and solvent of crystallization ( $\text{Et}_2\text{O}$ ) are omitted for clarity.

contain  $\kappa^2\text{-N,S-}o$ -aminobenzenethiolate ligands.<sup>59–61</sup> The Ni–N distance in **3** is shorter than the typical Ni– $\text{N}_{\text{amine}}$  bond and reflects the enhanced donor strength of the deprotonated nitrosamino-N, which is comparable to, although weaker than, a Ni– $\text{N}_{\text{carboxamido}}$  ( $\sim 1.86$  Å).<sup>15,58</sup> No evidence for a coordinated ligand radical is evident from the X-ray structure (*i.e.*, short C–S, C–N distances of the coordinated *o*-aminobenzenethiolate<sup>62</sup>) and confirm the N,S-ligand is a closed-shell dianion. The R–N–N–O linkage in **3** (avg. of two crystallographically distinct molecules, N–N: 1.309 Å, N–O: 1.264 Å; avg. N–N–O: 115.2°) is unremarkable from **2**.  $^1\text{H}$  and  $^{15}\text{N}$  NMR ( $\text{RN}^{15}\text{NO}$ ,  $\delta$ : 194 ppm *vs.*  $\text{CH}_3\text{NO}_2$ ) of crystals of **3** are consistent with the X-ray structure and analogous to other nitrosamines (Fig. S13 and S14†).<sup>55,57</sup> As expected, the IR of **3** lacks the intense  $\nu_{\text{NO}}$  from the NiNO of **2** (although IR and ESI-MS show that some **2** remains even in crystals of **3**, Fig. S12†) and the  $\nu_{\text{NO}}$  and  $\nu_{\text{NN}}$  of the R–N–N–O unit is similar. HR-ESI-MS(–) confirm this formulation with peaks corresponding to  $[\text{M}-2\text{Et}_4\text{N}]^{2-}$  ( $m/z$ : 248.960,  $z = 2$ , for **3**;  $m/z$ : 249.957,  $z = 2$ , for **3- $^{15}\text{NO}$** ) as the prominent peak (Fig. S15–S18†).

To confirm that NO(g) is released from **2** (forming **3** among other products), solutions of **2** were mixed with the NO(g) trap  $[\text{Co}(\text{T}(\text{OME})\text{PP})]$  ( $\text{T}(\text{OME})\text{PP} = 5,10,15,20\text{-tetrakis}(4\text{-methoxyphenyl})\text{-}21\text{H},23\text{H}\text{-porphine}$ ).<sup>63</sup> For example, mixing **2** and the  $\text{Co}^{\text{II}}\text{-P}$  (1 : 2) in  $\text{CH}_2\text{Cl}_2$  at RT for 24 h resulted in the  $\{\text{CoNO}\}^8$  complex  $[\text{Co}(\text{T}(\text{OME})\text{PP})(\text{NO})]$  in  $\sim 70\%$  avg. yield as quantified by  $^1\text{H}$  NMR ( $\text{CD}_2\text{Cl}_2$ ) and further verified by IR spectroscopy using **2- $^{15}\text{NO}$**  (Fig. S21–S23†). Notably, the reaction mixture becomes red over the course of the reaction. Workup of this solution after separating the Co–P compounds (MeOH-insoluble) reveals the presence of **3** (MeOH-soluble) *via*  $^1\text{H}$  NMR to confirm the fate of the  $\{\text{NiNO}\}^{10}$  complex **2**. To eliminate bimolecular NO-transfer *via* a putative  $\text{Co}\cdots\text{NO}\cdots\text{Ni}$  intermediate, NO(g) release was further verified by vial-to-vial trapping reactions wherein a  $\text{CH}_2\text{Cl}_2$  solution of the  $\text{Co}^{\text{II}}\text{-P}$  was separated from an MeCN solution of **2** ( $\text{Co}^{\text{II}}\text{-P}$  in excess, see the ESI†). Carrying out this reaction confirmed that NO(g) is indeed released from **2** (or **2- $^{15}\text{NO}$** ) to generate the  $\{\text{CoNO}\}^8$  porphyrin complex (80% avg. yield) as shown by  $^1\text{H}$  NMR and IR measurements (Fig. S24†). In contrast, no reaction takes place between THF solutions of **2** with  $[\text{Fe}(\text{TPP})\text{Cl}]$  (1 : 2; TPP = 5,10,15,20-tetraphenylporphyrin), a common HNO (or  $\text{NO}^-$ ) trap.<sup>64</sup> Although  $\{\text{NiNO}\}^{10}$  has not been characterized as



a particularly labile EF notation, we note that the majority of these complexes are cationic or neutral without coordinated thiolate ligands.<sup>28</sup> Indeed, the thiolate-ligated  $\{\text{NiNO}\}^{10}$  complex **III** photochemically releases NO to  $[\text{Co}(\text{TPP})]$  in MeCN suggesting some lability in the Ni–NO bond. Furthermore, the RN–NO bond is quite stable (as noted by formation of **3**) and the energetically stabilized MOs that contribute to the electronic structure of **2** and **3** where HOMO–3 represents a bonding MO with primary contributions from  $\sigma$ -NR and  $\sigma$ -NO orbitals (Fig. S25†).

Density functional theory (DFT) computations have provided a deeper understanding of the electronic structure of a variety of metal nitrosyls,<sup>65,66</sup> and we have employed them here for **2** and **3** at the OLYP/def2-TZVPP level of theory. Pure functionals such as BP86 and OLYP were used for geometry optimization and single point energy calculations, respectively, as these functionals have been established to deliver better matches with experimental geometries in MNO systems.<sup>67–69</sup> Geometry optimization of **2** was performed with coordinates from the crystal structure to yield DFT-optimized complex **2\*** (Fig. 4, Tables 1, S5 and S7 in the ESI†). Structurally, **2\*** replicates the metrics of **2** well, suggesting the computational model is reasonable. While the distances in **2\*** are within  $\pm 0.025$  Å of experimental values, the bond angles (especially S–Ni–S:  $-6.6^\circ$ , and Ni–N–O:  $+3.6^\circ$  from **2**) are slightly beyond the allowable tolerances for satisfactory DFT performance in small molecules (*i.e.*, distances  $\pm 0.03$  Å; angles  $\pm 1^\circ$ ).<sup>70</sup> However, these rules may be broken to some degree because of the enhanced complexity arising from the covalent MNO unit in **2**. The computations also reasonably match the two closely spaced N–O stretching frequencies for the symmetric and asymmetric  $\nu_{\text{NO}}$  in the IR at 1730 and 1708  $\text{cm}^{-1}$ , respectively. The  $\sim 30$   $\text{cm}^{-1}$  downshift from **2** is likely due to a slight overestimation of Ni–NO bond covalency arising from Ni–d $\pi$  backbonding. Previous calculations on three-<sup>71</sup> and four-coordinate<sup>43,72</sup>  $\{\text{NiNO}\}^{10}$  complexes support a  $\text{Ni}^{\text{II}}\text{-}^3\text{NO}^-$  ( $S_{\text{tot}} = 0$ , antiferromagnetically coupled) oxidation state assignment. This is comparable to high-spin nonheme  $\{\text{FeNO}\}^7$  systems that are classified as  $\text{Fe}^{\text{III}}\text{-}^3\text{NO}^-$  ( $S_{\text{tot}} = 3/2$ ).<sup>66,66,73</sup> In the Fe case,  $^3\text{NO}^-$  serves as a strong  $\pi$ -donor to afford a highly covalent Fe–NO bond.<sup>74</sup> The strength of this interaction originates from the effective nuclear charge on the metal, which is controlled by the

basicity of the supporting ligands.<sup>75</sup> Thus, electron rich supporting ligands attenuate the  $\pi$ -basicity of  $^3\text{NO}^-$  to result in diminished M–NO bond covalency. This property has been established in the  $\{\text{FeNO}\}^7$  case, but not yet for  $\{\text{NiNO}\}^{10}$ . Indeed, examination of the frontier MOs of **2\*** show that, much like other  $\{\text{NiNO}\}^{10}$  systems with Tp ligands<sup>43,72</sup> (Tp = tris(pyrazolyl)borate), the LUMO is a  $\pi^*$  MO primarily comprised of antibonding interactions between Ni–d $\pi$  and NO– $\pi^*$  orbitals (Fig. S25†). On the other hand, the HOMO (Fig. 4) and HOMO–1 have little contribution from NO, but large contributions from Ni–d $\sigma$  (38.0%) and S–p $\sigma$  (19.3%) orbitals of the Ni( $\mu$ -SR)<sub>2</sub>Ni core. The HOMO is antibonding in nature and suggests a thermally unstable structure. As expected from analogous  $\{\text{FeNO}\}^7$  systems, based on the increased donor strength of the anionic nitrosamine-N/thiolate-S supporting ligands in **2\***, the covalency in the Ni–N–O unit is less than in TpNi–NO complexes and rationalizes the observed lability of the Ni–NO bond and the Ni( $\mu$ -SR)<sub>2</sub>Ni core in **2**.

DFT computations on **3\*** were performed in the same fashion as for **2\***. Geometry optimized **3\*** is square-planar ( $\tau_4 = 0.09$ ) with metric parameters on-par with the X-ray structure of **3** and within the error of the DFT method (Table S9†). Unlike **2\***, the  $\pi^*$  HOMO of **3\*** is comprised primarily of Ni(d $\pi$ )/S(p $\pi$ ) contributions (Fig. S26†), typical of planar  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  complexes with strong-field ligands and suggests a highly covalent Ni–SR bond.<sup>15</sup>

The formation of **2** likely follows a mechanistic path analogous to those observed in the reductive nitrosylation of Cu-amine systems, where one-equiv. of NO reacts with  $\text{Cu}^{\text{II}}\text{-NR}_2$  complexes to yield  $\text{R}_2\text{N-NO}$  and deligated  $\text{Cu}^{\text{I}}$ .<sup>76,77</sup> The difference here is that the nitrosated ligand remains coordinated and the resulting paramagnetic Ni binds NO radical. Our working model is depicted in Scheme 2. Complex

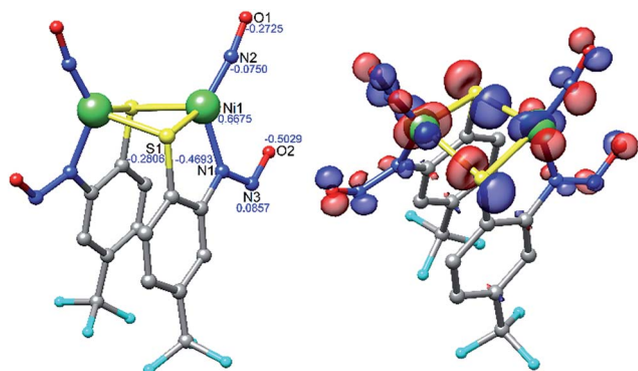
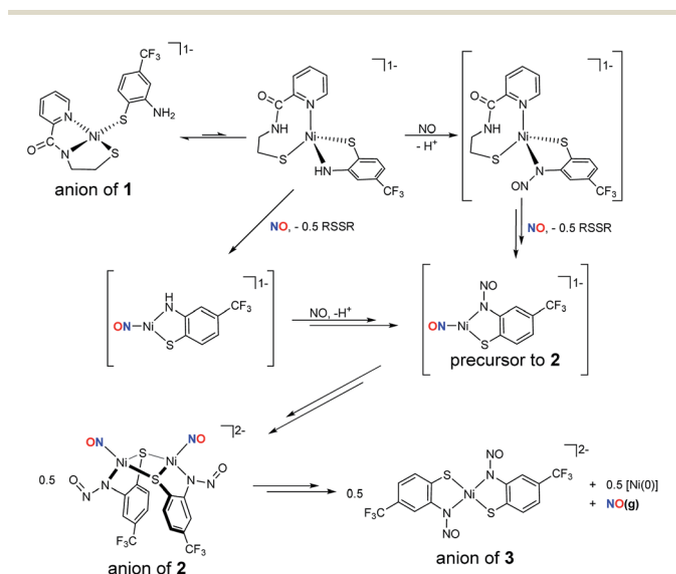


Fig. 4 DFT (OLYP/def2-TZVPP) optimized structure of **2\*** (left) with natural population analysis charges in blue and HOMO (right).



Scheme 2 Working model for the formation of  $\{\text{NiNO}\}^{10}$  complex **2** and  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  complex **3** starting from  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  monomer **1**. RSSR: disulfide of  $\text{nmp}^{2-}$ , *i.e.*, *N*-(2-mercaptoethyl)picolinamide. Intermediates represented in brackets have not been spectroscopically identified. The py-N of RSSR is a possible H<sup>+</sup> receptor.



**1** is likely in resonance with a distorted tetrahedral species which places the anilido-N in the coordination sphere. This proposal is supported by the presence of low intensity peaks in the  $^1\text{H}$  NMR spectrum of **1** and may explain the difficulty in crystallizing this complex.<sup>16</sup> On the other hand, X-ray absorption spectroscopic (XAS) characterization of **1**, not reported previously, suggests a four-coordinate planar  $\text{Ni}^{\text{II}}$  center (XANES analysis, see Fig. S3†) with two O/N- and S-ligands at 1.90 Å and 2.17 Å (EXAFS, Fig. S3, Table S4†), respectively. Thus, **1** is structurally analogous to other  $[\text{Ni}(\text{nmp})(\text{SR})]^-$  complexes at least in the solid-state. Introduction of  $\text{NO}(\text{g})$  can then result in either: (i) reduction of  $\text{Ni}^{\text{II}}$  to  $\text{Ni}^{\text{I}}$  and formation of  $\text{NO}^+$  that nitrosates the coordinated amine, or (ii) nitrosylation of Ni to yield  $\{\text{NiNO}\}^{10}$  with the electron originating from the coordinated thiolate of  $\text{nmp}^{2-}$  to result in the disulfide. Our results do not differentiate either of these transformations, but the disulfide of  $\text{nmp}^{2-}$  (*i.e.*,  $\text{nmpS}_2$   $^1\text{H}$  NMR and IR of the reaction mixture, see Fig. S19 and S20†) is spectroscopically observed in the reaction mixture and checked against independently synthesized  $\text{nmpS}_2$ . Thus, the fate of one proton and one electron is reasonably confirmed. At this point these intermediates can react with another equiv. of  $\text{NO}$  to yield the three-coordinate precursor to **2**. Compound **3** forms through either disproportionation (shown in Scheme 2) to yield a  $\text{Ni}^0$  species or ligand rearrangement *via* the loss of a  $\text{Ni}^{\text{I}}\text{-NO}$  fragment (not shown). In ligand rearrangement, the products would be a  $\text{Ni}^{\text{I}}\text{-N}_2\text{S}_2$  precursor to **3** (**3-PC**), an  $\text{L-Ni}^{\text{I}}\text{-NO}$  species ( $\text{L}$  = solvent), and free  $\text{NO}$ . Ultimately this  $\text{Ni}^{\text{I}}$  intermediate oxidizes **3-PC** to generate  $\text{Ni}^{\text{II}}$  complex **3** and an  $\text{L-Ni}^0\text{-NO}$  complex that would presumably release  $\text{NO}(\text{g})$  as evidenced by the  $\text{NO}(\text{g})$  trap experiments (*vide supra*). While the reaction mechanism for the conversion of **2**-to-**3** is likely more complex, similar chemistry has been proposed for N-heterocyclic carbene (NHC) Ni-nitrosyls.<sup>28,78</sup> The details of this mechanism are still under investigation.

## Conclusions

In conclusion, NiSOD model complex **1** reacts with  $\text{NO}(\text{g})$  in the  $\text{Ni}^{\text{II}}$  state to form the metastable  $\{\text{NiNO}\}^{10}$  dimeric complex **2** *via* loss of the  $\text{nmp}^{2-}$  ligand as the disulfide and N-nitrosation of the *o*-aminobenzenethiolate ligand. Reaction of  $\text{NO}$  with  $\mathbf{1}^{\text{ox}}$ , or  $\text{NO}^+$  with **1**, only yields the *S,S*-bridged tetrameric compound  $[\text{Ni}_4(\text{nmp})_4]$  through oxidation of the aromatic thiolate ligand. While any reaction with  $\text{NO}$  ( $S = 1/2$ ) is generally unexpected for square-planar ( $S = 0$ )  $\text{Ni}^{\text{II}}$  complexes, this Ni-nitrosyl likely forms due to an equilibrium mixture of **1** and a tetrahedral ( $S = 1$ ) or five-coordinate derivative (Scheme 2). Even if  $\text{NO}$  were to result in an  $\text{nmp}$ -bound Ni- $\text{NO}$  complex, the resulting  $\{\text{NiNO}\}^9$  (reaction of **1** with  $\text{NO}$ ) or  $\{\text{NiNO}\}^8$  (reaction of  $\mathbf{1}^{\text{ox}}$  with  $\text{NO}$ ) oxidation levels have yet to be defined and support an outer-sphere superoxide interaction in NiSOD. Although these EF notations have yet to be accessed, one would propose that NiSOD mimetics, especially with strong-field carboxamido-N and alkyl-thiolato-S donors, would surely stabilize such an electron poor species. Furthermore, the properties of complexes such as **2** extend to biology, where analogous S-bridged

mononitrosyl species, *i.e.*, Fe-S clusters and tetrahedral  $(\text{RS})_3\text{Fe-NO}$  complexes are proposed as intermediates in the repair of NO-damaged clusters.<sup>79–81</sup> Complex **2** is stable in the solid-state but breaks down slowly in solution causing rupture of the  $\text{Ni}(\mu\text{-SR})_2\text{Ni}$  core and release of  $\text{NO}$  that was trapped in near quantitative yield with a  $\text{Co}^{\text{II}}$ -porphyrin receptor. The resulting  $\text{Ni}^{\text{II}}\text{-N}_2\text{S}_2$  complex **3** (coordination of two *o*-aminobenzenethiolate in *trans* configuration) was isolated and structurally/spectroscopically characterized as the ultimate Ni breakdown product with the nitrosamine unit still intact. This release may take place through a disproportionation mechanism (or through ligand rearrangement), as has been proposed in other Ni-nitrosyls, to a yet ill-defined  $\text{Ni}^0$  complex (see Scheme 2).<sup>28,78</sup> Hence, thiolate-supported  $\{\text{NiNO}\}^{10}$  cores are reactive. While nitrosamines have been utilized as sources of  $\text{NO}$ , the  $\text{RN-NO}$  homolytic bond dissociation energy (BDE) is high (87.7 kcal mol<sup>-1</sup> (ref. 82)) compared to more traditional small molecule sources of  $\text{NO}$  such as nitrosothiols ( $\text{RSNO}$ ) that have  $\text{RS-NO}$  BDEs between 20–32 kcal mol<sup>-1</sup>.<sup>83,84</sup> Overall, the electronic structures of  $\{\text{NiNO}\}^{10}$  complexes are modulated by the supporting ligands. Indeed, the majority are stable entities; however, a small number are reactive and result in release of  $\text{NO}$  (thiolate-supported/anionic complexes **2** and **III**) or generate other reactive intermediates of environmental significance such as hyponitrite ( $\text{N}_2\text{O}_2^{2-}$ ) in five-coordinate  $\{\text{NiNO}\}^{10}$  species<sup>85</sup> (highly reduced  $\text{NO}$ , with a severely bent Ni-N-O angle = 130°).

## Conflicts of interest

There are no conflicts to declare.

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