



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Nickel-mediated N–N bond formation and N₂O liberation *via* nitrogen oxyanion reduction†

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The syntheses of (DIM)Ni(NO₃)₂ and (DIM)Ni(NO₂)₂, where DIM is a 1,4-diazadiene bidentate donor, are reported to enable testing of bis boryl reduced N-heterocycles for their ability to carry out stepwise deoxygenation of coordinated nitrate and nitrite, forming O(Bpin)₂. Single deoxygenation of (DIM)Ni(NO₂)₂ yields the tetrahedral complex (DIM)Ni(NO)(ONO), with a linear nitrosyl and κ¹-ONO. Further deoxygenation of (DIM)Ni(NO)(ONO) results in the formation of dimeric [(DIM)Ni(NO)]₂, where the dimer is linked through a Ni–Ni bond. The lost reduced nitrogen byproduct is shown to be N₂O, indicating N–N bond formation in the course of the reaction. Isotopic labelling studies establish that the N–N bond of N₂O is formed in a bimetallic Ni₂ intermediate and that the two nitrogen atoms of (DIM)Ni(NO)(ONO) become symmetry equivalent prior to N–N bond formation. The [(DIM)Ni(NO)]₂ dimer is susceptible to oxidation by AgX (X = NO₃[−], NO₂[−], and OTf[−]) as well as nitric oxide, the latter of which undergoes nitric oxide disproportionation to yield N₂O and (DIM)Ni(NO)(ONO). We show that the first step in the deoxygenation of (DIM)Ni(NO)(ONO) to liberate N₂O is outer sphere electron transfer, providing insight into the organic reductants employed for deoxygenation. Lastly, we show that at elevated temperatures, deoxygenation is accompanied by loss of DIM to form either pyrazine or bipyridine bridged polymers, with retention of a BpinO[−] bridging ligand.

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Introduction

Reduction of CO₂ is widely sought and has classically focused on deoxygenation to C₁ products,^{1–3} with recent advances moving towards multi-carbon and value-added products which preserve terrestrial use of that carbon resource.^{4–9} There is a nitrogen analogue of this challenge, accomplished biologically through denitrification, which requires an N–N bond formation upon reduction of NO_x[−] back to dinitrogen. It is widely recognized that the natural nitrogen cycle has been perturbed as an unintended consequence of inexpensive Haber–Bosch ammonia used to enhance food production.^{10–12} Soluble nitrate is abundant in runoff from agricultural fields, and accumulates in poorly flushed rivers, bays and seacoasts which ultimately leads to eutrophication and oxygen-poor “dead zones.”^{13–15}

The reduction of nitrogen oxidation states is biologically accomplished by a variety of enzymes^{16–23} and its synthetic conversion is essential to repurpose environmental pollutants

into value-added compounds that might be derived from nitrogen oxyanions. An attractive route to nitrogen oxyanion reduction is through deoxygenation,^{24–32} and the oxygen can be captured by protons, metal electrophiles, or main group elements known for their oxophilicity. The same is true for CO₂ reduction and both carbon and nitrogen reduction would be richer if they could form products with C–C and N–N bonds, respectively. Designing methods to encourage such element/element reductive coupling is a valuable pursuit.

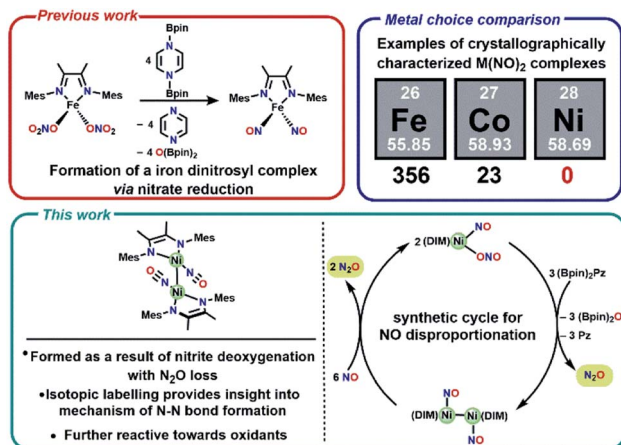
One attractive strategy to promote N–N bond formation is to design complexes which pre-organize two nitrogen oxyanions at a single metal center and explore the deoxygenation chemistry of these systems. There are several reported examples of N–N bond formation at a single metal complex,^{33–36} and most terminal *cis*-hyponitrite complexes are reported with group 10 metals.^{37–41} The reagent bis-pinacolylboryl pyrazine (Scheme 1) has been shown to deoxygenate two nitrates coordinated to iron to yield a dinitrosyl iron complex.³¹ This reagent is electron rich (8π electron ring) and carries two electrophilic boron atoms making it a potent reductant with attractive kinetic reactivity, including polar B–N bonds.

We chose a bidentate ancillary ligand with a steric profile designed to allow for multidentate binding of nitrogen oxyanions if needed. These 1,4-diazadienes (DIM, Scheme 1) ligands are established to be redox active and can assist in the stabilization of low valent nickel complexes.^{42–45} Herein we show

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† Electronic supplementary information (ESI) available. CCDC 2083107 (DIM)Ni(NO₃)₂, 2083108 (DIM)Ni(NO₂)₂, 2083111 (DIM)Ni(NO)(ONO), 2083110 [(DIM)Ni(NO)]₂, 2083109 (DIM)Ni(NO)(OTf), 2083112 (OBpin)₂Ni₂(NO)₂(Pz) and 2083113 (OBpin)₂Ni₂(NO)₂(Bpy). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d1sc02846d





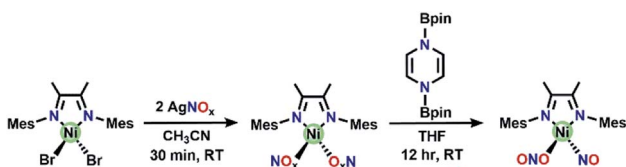
Scheme 1 Previously reported nitrate reduction and motivation for utilizing Ni compounds (top), and summary of new reactivity presented (bottom).

stepwise deoxygenation of oxanions coordinated to nickel, where the location of nickel late in the 3d series disfavors two coordinated nitrosyls in comparison to Fe or Co (Scheme 1), and instead deoxygenation leads to N-N bond formation and nitrous oxide liberation.

Results and discussion

Nickel nitrate and nitrite complexes

A precursor complex, $(DIM)NiBr_2$, is synthesized from $(DME)NiBr_2$ ($DME = \text{dimethoxyethane}$) and DIM in a 1 : 1 mole ratio in DCM and workup affords a red/brown solid. 1H NMR spectroscopy (Fig. S1[†]) of $(DIM)NiBr_2$ reveals a paramagnetic species with resonances ranging from -22 to 36 ppm and is fully consistent with a C_{2v} symmetric product. Salt metathesis reactions of $(DIM)NiBr_2$ with $AgNO_3$ or $AgNO_2$ in acetonitrile at room temperature quickly form the bis-nitrate and bis-nitrite complexes respectively with concurrent precipitation of $AgBr$ (Scheme 2). The bis-nitrate and bis-nitrite complexes are also paramagnetic and twofold symmetric as judged by NMR spectroscopy (Fig. S3 and S5[†]) and their molecular structures (Fig. 1) exhibit bidentate binding of both nitrate and nitrite creating an octahedral coordination environment about the nickel metal centers.



Scheme 2 Synthesis of $(DIM)Ni(NO_x)_2$ complexes, as well as the synthesis of $(DIM)Ni(NO)(ONO)$ from $(DIM)Ni(NO_2)_2$. The bis-nitrate and bis-nitrite complexes are O-bound and bidentate in the solid-state molecular structures (Fig. 1), we have drawn them as monodentate here for simplicity.

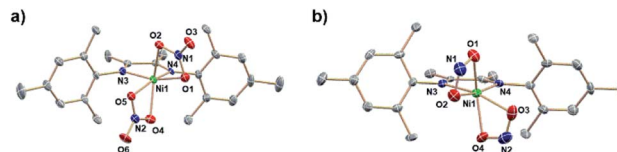


Fig. 1 (a) ORTEP representation of the molecular structure (50% probabilities) of the nonhydrogen atoms of (a) $(DIM)Ni(NO_2)_3$ and (b) $(DIM)Ni(NO_2)_2$ (right), showing selected atom labelling (unlabelled atoms are carbon). Selected structural parameters for $(DIM)Ni(NO_2)_3$ (Å): Ni1–O1, 2.110(4); Ni1–O2, 2.093(4); Ni1–O4, 2.050(4); Ni1–O5, 2.097(4); Ni1–N3, 2.009(5); Ni1–N4, 2.058(4); O1–N1, 1.265(6); O2–N1, 1.273(6); O3–N1, 1.230(6); O4–N2, 1.298(6); O5–N2, 1.271(6); O6–N2, 1.255(6); and $(DIM)Ni(NO_2)_2$ (Å): Ni1–O1, 2.094(6); Ni1–O2, 2.091(6); Ni1–O3, 2.099(6); Ni1–O4, 2.072(6); Ni1–N3, 2.035(7); Ni1–N4, 2.038(7); O1–N1, 1.287(9); O2–N1, 1.269(9); N2–O3, 1.257(10); N2–O4, 1.290(11).

Single deoxygenation of $(DIM)Ni(NO_2)_2$

Reaction of $(DIM)Ni(NO_2)_2$ with equimolar $(Bpin)_2Pz$ occurs to completion within 12 h at room temperature in THF with a color change from brown to green (Scheme 2). Analysis by 1H NMR spectroscopy shows complete consumption of the boryl reagent, formation of equimolar pyrazine and $(Bpin)_2O$ and a diamagnetic metal complex with the expected 4 : 6 : 6 : 12 intensity 1H NMR resonances of a single DIM-containing product. Workup by pentane wash to remove organic materials yields a green solid, and the molecular structure (Fig. 2) confirms the presence of the singly deoxygenated species $(DIM)Ni(NO)(ONO)$, assigned as $\{Ni(NO)\}^{10}$ in the Enemark–Feltham notation. The Ni1–N3–O1 angle is linear (175.5°), and the complex has a strong nitrosyl stretching frequency at 1785 cm^{-1} , similar to other $LNi(NO)(X)$ complexes.^{46–49} The nitrito linkage in $(DIM)Ni(NO)(ONO)$ contrasts the N-bound nitrite linkage in $(Me_2\text{-ortho-phenanthroline})Ni(NO)(NO_2)$.⁴⁹

Deoxygenation of $(DIM)Ni(NO)(ONO)$

The tetrahedral nickel(0) $(DIM)Ni(NO)(ONO)$ complex reacts with $(Bpin)_2Pz$ (1 : 1.5 mole ratio) at room temperature in THF. Monitoring the reaction by 1H NMR spectroscopy shows the

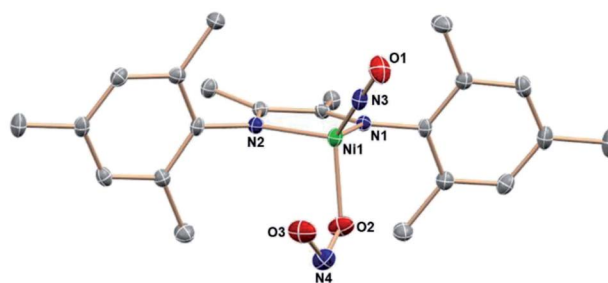


Fig. 2 ORTEP representation of the molecular structure (50% probabilities) of the nonhydrogen atoms of $(DIM)Ni(NO)(ONO)$ showing selected atom labelling (unlabelled atoms are carbon). Selected structural parameters: Ni1–N3, 1.637(2); Ni1–O2, 1.9952(19); Ni1–N2, 2.0075(19); Ni1–N1, 2.0204(19); O1–N3, 1.176(3); O2–N4, 1.280(3); O3–N4, 1.228(3); N4–O2–Ni1, 118.25(16); O1–N3–Ni1, 175.5(2).



formation of boryl ether and pyrazine, as well as the depletion of signals for (DIM)Ni(NO)(ONO). Crystalline precipitate forms during the course of the reaction. Executing this reaction in CD₃CN shows complete consumption of starting materials in less than four hours at room temperature. All DIM-containing product is insoluble based on the lack of any ligand resonances, but the pyrazine and boryl ether byproducts are detected by ¹H NMR spectroscopy (Fig. S19†) in the correct stoichiometric ratio in the colorless solution. This confirms that no free ligand is formed upon deoxygenation, nor any degradation products containing the DIM ligand. The reaction executed in CD₂Cl₂ shows the same result, including no detectable ¹H NMR signals of the product nickel complex. This solvent gives slightly larger crystals, which were used for the better of two single crystal structure determinations. The molecular structure (Fig. 3) as determined by SCXRD reveals the product to be [(DIM)Ni(NO)]₂, a dimer with a Ni–Ni bond.

[(DIM)Ni(NO)]₂ contains a crystallographic inversion center at the Ni–Ni midpoint and the Ni–Ni bond length (2.6103(10) Å) is short compared to other Ni–Ni bonds with no bridging ligands.^{50–53} There are no other examples of two Ni–NO units linked solely by a Ni–Ni bond. The Ni lies only 0.16 Å out of the three-nitrogen atom plane, and the Ni'/N1 distance is longer by 1.24 Å, indicating that the NO is not bridging. Analysis of the bond lengths in the DIM backbone (N–C–C–N) (Table S1†) reveals a long-short-long pattern (*vs.* the comparison compounds in Table S1†) consistent with increased electron density in the di-imine moiety. The C–C bond in particular is quite short (1.436 Å) compared to the other (DIM)Ni complexes, indicative of either a singly or doubly reduced ligand. The monomer (DIM)Ni(NO) is an odd-electron (radical) species which dimerizes to the crystallographically observed product. The Δ*G*^o for dissociation of the dimer into two (DIM)Ni(NO) radical monomers is calculated to be 14.7 kcal mol^{−1}, similar to reported values for dissociation of an iron carbonyl dimer.⁵⁴ The MALDI-TOF spectrum (Fig. S18†) of [(DIM)Ni(NO)]₂ has an *m/z* at 816.3, consistent with the intact dimer. The *trans* rotational conformation around the nickel–nickel bond is sterically favored. This conformation has the stabilizing effect of

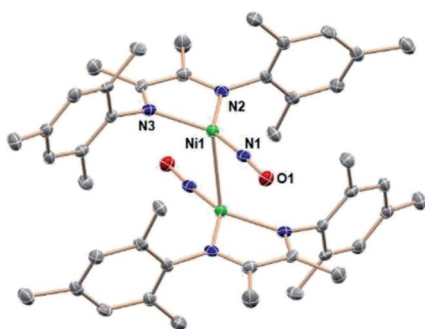


Fig. 3 ORTEP representation of the molecular structure (50% probabilities) of the nonhydrogen atoms of centrosymmetric [(DIM)Ni(NO)]₂, showing selected atom labelling (unlabelled atoms are carbon). Selected structural parameters: Ni1–N3, 1.960(4) Å; Ni1–N2, 1.893(4); Ni1–N1, 1.647(3); O1–N1, 1.193(4); O1–N1–Ni1 172.5(3)^o.

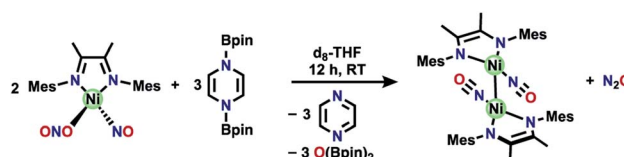
opposing the dipoles in each (DIM)Ni(NO) monomer and benefits from opposed NO^{δ+} to DIM^{δ−} attraction between monomer units. Because the dimeric product is insoluble in a wide range of solvents of polarity up to acetonitrile and dichloromethane, a ¹H NMR spectrum could not be obtained. The solid state IR spectrum (Fig. S15†) shows a strong NO absorption at 1664 cm^{−1}, a low value consistent with greater back donation to the nitrosyl than in (DIM)Ni(NO)(ONO). This can be attributed to the increased donor ability of the reduced DIM ligand as well as better orbital overlap.

The reaction of (DIM)Ni(NO)(ONO) with equimolar (Bpin)₂Pz shows complete consumption of borylated reagent along with incomplete consumption of nickel starting material. In order to establish the correct stoichiometry, (DIM)Ni(NO)(ONO) was reacted with excess (3 : 1 mole ratio) (Bpin)₂Pz in the presence of naphthalene as an internal standard. From this experiment, pyrazine is produced in a 1 : 1.5 mole ratio with respect to the (DIM)Ni(NO)(ONO) starting material, indicating a 2 : 3 stoichiometry of nickel to borylating reagent. Labelling studies are employed here for characterization, and later for mechanistic insight. The nitrosyl stretch shifts to 1625 cm^{−1} (Fig. S16†) for isotopically labelled [(DIM)Ni(¹⁵NO)]₂, which is consistent with the expected reduced mass calculation. The reduced nitrogen containing byproduct upon deoxygenation of (DIM)Ni(NO)(ONO) was confirmed to be N₂O *via* vacuum transfer of the headspace volatiles to a gas IR cell, which clearly show the P and R bands of N₂O at 2224 cm^{−1} (Fig. S30†). All of this allows us to write a balanced reaction for the formation of [(DIM)Ni(NO)]₂ with concomitant loss of N₂O (Scheme 3).

Deoxygenation of isotopically labelled (DIM)Ni(¹⁵NO)(O¹⁵NO) results in the formation of (¹⁵N)₂O, evidenced by a band at 2158 cm^{−1} *via* gas IR (Fig. S30†). The deoxygenating agent employed in Scheme 3 can be replaced by the reduced 4,4'-bipyridine analogue, (Bpin)₂Bpy, which reacts completely within 2 hours. The faster deoxygenation using (Bpin)₂Bpy *vs.* (Bpin)₂Pz is a consequence of reduced bipyridine being a more potent reductant than reduced pyrazine.⁵⁵

XPS of nickel complexes

XPS provides useful resolution of the chemical environment in these four molecules (see Fig. S44†). The nitrogen 1s region resolves nitrate, nitrite, nitrosyl and all the DIM nitrogen atoms, with progressively lower NO_x binding energy along that series. The oxygen 1s intensity decreases progressively from nitrate to nitrite to nitrosyl when integrated with respect to the nitrogen. Compound (DIM)Ni(NO)(ONO) shows two oxygen 1s binding energies with relative intensities of 1 : 2, indicating resolution



Scheme 3 Synthesis of [(DIM)Ni(NO)]₂ from (DIM)Ni(NO)(ONO) and (Bpin)₂Pz.

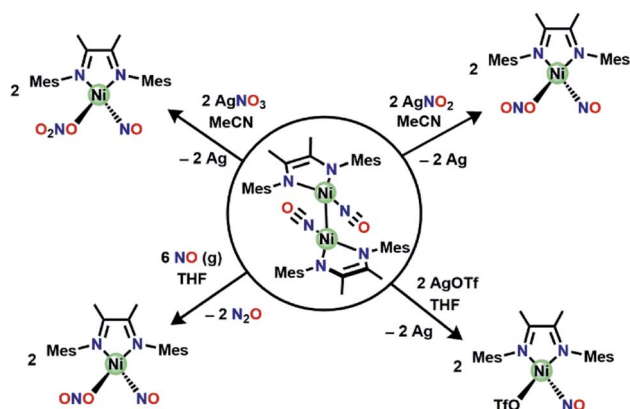


of two of the three types of oxygens. The oxygen 1s binding energy decreases progressively from (DIM)Ni(NO₃)₂ to (DIM)Ni(NO₂)₂ to [(DIM)Ni(NO)]₂, consistent with a lower nitrogen oxidation state being less electron withdrawing towards the oxygen. Although linear nitrosyl and nitrite are both formally trivalent nitrogen, the lower binding energy of oxygen by 0.57 eV in [(DIM)Ni(NO)]₂ compared to (DIM)Ni(NO₂)₂ can be attributed to the increased backdonation into the nitrosyl of the dimer, consistent with the observed low ν_{NO} stretching frequency. The nitrogen 1s spectrum of [(DIM)Ni(NO)]₂ resolves the nitrosyl from imine in the intensity ratio of 1 : 2. The Ni 2p spectrum shows a lower metal binding energy for [(DIM)Ni(NO)]₂ compared to the bis-nitrate and bis-nitrite complexes, consistent with it having a lower metal oxidation state.

Redox reactivity of [(DIM)Ni(NO)]₂

Despite the insolubility of [(DIM)Ni(NO)]₂, it reacts readily (Scheme 4) with oxidants to cleave the Ni–Ni bond. The reaction between a slurry of [(DIM)Ni(NO)]₂ with AgNO₂ or AgNO₃ results in homogeneity of the solution and a color change to green in both cases. Isolation of the green solid from the reaction with AgNO₂ reveals the product to be the previously characterized (DIM)Ni(NO)(ONO) by ¹H and ¹³C NMR spectroscopy along with the diagnostic NO stretch at 1785 cm⁻¹. The corresponding reaction of the dimer and AgNO₃ results in the formation of the {Ni(NO)}¹⁰ complex (DIM)Ni(NO)(ONO₂), indicating that the one electron oxidation product of [(DIM)Ni(NO)]₂ is susceptible to nucleophilic attack even by weak nucleophiles such as nitrate. It is important here to control the Ag : Ni stoichiometry because both (DIM)Ni(NO)(ONO) and (DIM)Ni(NO)(ONO₂) are subject to further oxidation by silver salts, forming the bis-nitrite or bis-nitrate complexes respectively. The oxidations of the mixed oxanyon complexes result in liberation of an equivalent of nitric oxide, which was detected *via* gas phase IR (Fig. S34†). There is precedence for nitric oxide liberation upon one electron oxidation of [Ni(NO)(bipy)][PF₆] with either AgPF₆ or [NO][PF₆] as the oxidant.⁴⁸

Even an oxidant with a more weakly nucleophilic triflate anion succeeds in this oxidative process. The reaction of [(DIM)Ni(NO)]₂ with AgOTf in THF results in a color change to green



Scheme 4 Summary of oxidative reactivity with [(DIM)Ni(NO)]₂.

and homogeneity of the solution. The ¹⁹F NMR spectrum (Fig. S21†) is indicative of one diamagnetic product, and the ¹H NMR (Fig. S21†) spectrum indicates twofold symmetry. The solid state nitrosyl stretch appears at 1780 cm⁻¹ (Fig. S22†) which is shifted to lower energy by 100 cm⁻¹ compared to authentic 3-coordinate (bpy)Ni(NO)⁺ (bpy = bipyridine).⁴⁹ The shift in ν_{NO} can be attributed to coordination of THF solvent to give the 4-coordinate species [(DIM)Ni(NO)(THF)]⁺. When the reaction of [(DIM)Ni(NO)]₂ and AgOTf is repeated in the non-coordinating solvent DCM, the ν_{NO} is higher by 31 cm⁻¹ in the solid state (Fig. S23†) and 45 cm⁻¹ in a DCM solution (Fig. S24†). If the same material is redissolved in THF, the IR spectrum shows two nitrosyl stretches suggesting an equilibrium of THF binding to the nickel center. Crystals were grown *via* vapor diffusion of pentane into a concentrated THF solution and the solid state molecular structure (Fig. 4) shows the {Ni(NO)}¹⁰ complex (DIM)Ni(NO)(OTf), with triflate coordinated to nickel. However, based on spectroscopic data, the OTf is in equilibrium when dissolved in coordinating solvents.

With this weak nucleophile, the tetrahedron is distorted to flatten the (DIM)NiN(O) plane: the angle from the DIM N–C=C–N midpoint to the Ni center to N₁ is 135.6°. Regardless of the abundance of THF as crystallization solvent, this neutral molecule is apparently the least soluble equilibrium participant.

Nitric oxide disproportionation

With the knowledge that [(DIM)Ni(NO)]₂ reacts rapidly with silver oxidants, we were interested in its reactivity with NO gas. A suspension of [(DIM)Ni(NO)]₂ in THF was exposed to excess NO and there was an immediate color change to a homogeneous brown/green solution. Workup of the solution indicates that the nickel-containing product of this reaction is (DIM)Ni(NO)(ONO), which is established by a variety of spectroscopic techniques along with an independent unit cell determination of the molecular structure *via* SCXRD. Vacuum transfer of the volatiles generated during the reaction into a gas IR cell show

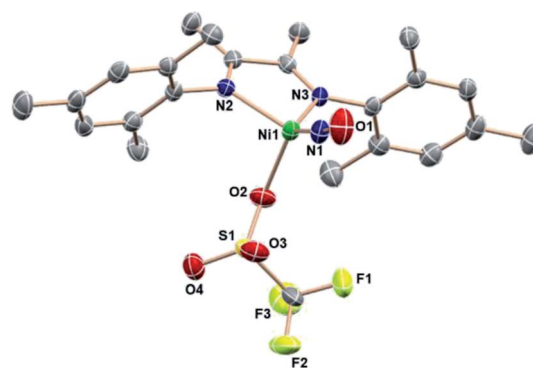


Fig. 4 ORTEP representation of the molecular structure (50% probabilities) of the nonhydrogen atoms of (DIM)Ni(NO)(OTf), showing selected atom labeling (unlabelled atoms are carbon). Selected structural parameters (Å, °): Ni1–O2, 2.059(2); Ni1–N3, 1.994(2); Ni1–N2, 2.010(2); Ni1–N1, 1.630(2); O1–N1–Ni1, 173.1(3); N1–Ni1–O2, 124.22(11).





Scheme 5 Metal-mediated nitric oxide disproportionation.

the formation of N_2O (Fig. S31†). This reaction therefore accomplishes disproportionation of nitric oxide to N_2O and NO_2 , but the latter undergoes one electron reduction by nickel to yield the coordinated nitrito product (Scheme 4).

As evidenced by Scheme 5, two low valent nickel centers of $[(\text{DIM})\text{Ni}(\text{NO})]_2$ mediate the disproportionation of 6 moles of NO resulting in the formation of two $\text{Ni}(\text{ONO})$ complexes and two moles of N_2O , a reaction which has precedence with first row transition metals.^{49,56–63} The reaction of $[(\text{DIM})\text{Ni}(\text{NO})]_2$ with NO gas can be accomplished in the absence of solvent. Addition of excess nitric oxide to a solid sample of $[(\text{DIM})\text{Ni}(\text{NO})]_2$ over the duration of 15 min, followed by extraction into THF indicates the formation of $(\text{DIM})\text{Ni}(\text{NO})(\text{ONO})$ *via* IR and ^1H NMR spectroscopy. Attempts to detect an intermediate in this disproportionation reaction by solid state IR were investigated by exposing a solid sample of $[(\text{DIM})\text{Ni}(\text{NO})]_2$ to 1–4 equivalents of nitric oxide. This resulted in up to 15% growth of $(\text{DIM})\text{Ni}(\text{NO})(\text{ONO})$ but no new intermediate terminal nitrosyl and no growth of other bands between 1000 and 1600 cm^{-1} (Fig. S37†). The fact that this reaction occurs cleanly in the solid state, where diffusive separation of intermediate nickel monomers is physically inhibited, makes it plausible that an intermediate is two identical $(\text{DIM})\text{Ni}(\text{NO})_2$.

Nitrogen isotope label studies

Preparation of isotopomers. We sought an isotopic crossover experiment with deoxygenation of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ in a 1 : 1 mole ratio, where any detected $^{14}\text{N}^{15}\text{NO}$ or $^{15}\text{N}^{14}\text{NO}$ would indicate that the N–N bond formation occurs within a bimolecular Ni_2 species. We isotopically labelled $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and found that it exhibits a broad ^{15}N NMR chemical shift at 573.8 ppm that shows a T^{-1} dependence of chemical shift which is characteristic of paramagnetic complexes (Fig. S7†). Furthermore, the nitrite ligand stretches

shift from 1295 cm^{-1} and 1210 cm^{-1} to 1276 cm^{-1} and 1180 cm^{-1} in the ^{15}N isotopomer (Fig. S9†). Doubly labelled $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ can be synthesized by reductive bor-ylation of the labelled $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ starting material. Isotopically labelled $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ shows a diagnostic shift for both NO and nitrite IR signals (Fig. S13†). Curiously, ^{15}N NMR of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ in both THF and MeCN at room temperature shows only one signal at 3.6 ppm (Fig. S11†) despite chemically inequivalent nitrogen atoms. However, room temperature ^{15}N NMR in DCM (Fig. S11†) shows the same signal at $\sim 3.6\text{ ppm}$ along with a broad signal at 206 ppm . Cooling this solution to $-42\text{ }^\circ\text{C}$ results in two sharp signals at -19.1 and 216.3 ppm in a 1 : 1 intensity ratio (Fig. 5). To confirm the assignment of the two signals, the isotopomers $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ were synthesized *via* the reaction of the appropriately labelled $[(\text{DIM})\text{Ni}(\text{NO})]_2$ dimer with either AgNO_2 or $\text{Ag}^{15}\text{NO}_2$ (see ESI† for isotopic details). At $-42\text{ }^\circ\text{C}$ in DCM, ^{15}N NMR of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ shows one sharp signal at 216.3 ppm , and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ shows one sharp signal at -19.1 ppm (Fig. 5). The specific isotopomers confirm that the original unseen ^{15}N NMR signal of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ in THF and MeCN corresponds to the nitrito ligand, consistent with a dynamic process that exchanges the nitrito ligand between nickel complexes. This dynamic process is confirmed *via* mass spectrometry with a 1 : 1 THF solution of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$. The ESI+ spectrum is consistent with isotopic scrambling which shows a statistical mixture of the original doubly labelled and unlabelled compounds, together with equal amounts of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ (Fig. S38†). The presence of the mixed labelled isotopomers is consistent with a dynamic process of intermolecular nitrite exchange between metal centers, and this dynamic process is supported by the line broadening ^{15}N NMR observations above.

Insight into the N–N bond formation. The nitrito scrambling in a 1 : 1 mixture of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ precludes any mechanistic insight obtained from a mixture of these two complexes. Instead, we focused on the individual isotopomers $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ to obtain mechanistic information about the

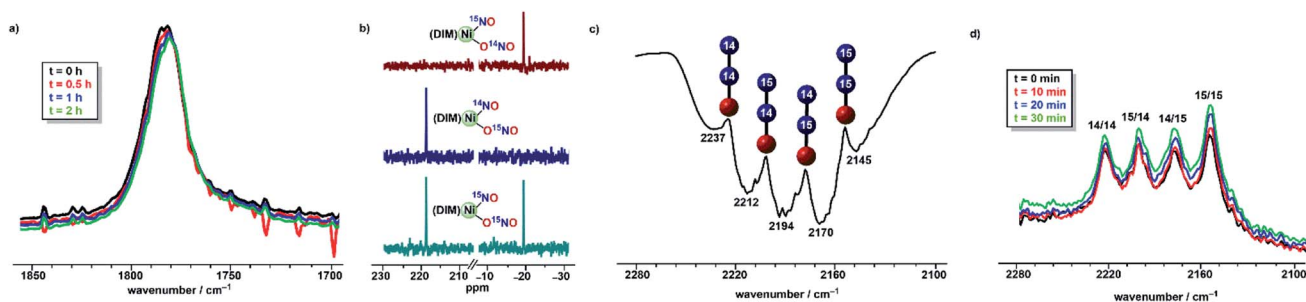


Fig. 5 (a) Solution IR of the nitrosyl stretch in $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ monitored over time (b) ^{15}N NMR of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ (bottom), $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{14}\text{NO})$ (middle), and $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ (top) all in CD_2Cl_2 at $-42\text{ }^\circ\text{C}$ (c) gas phase IR spectrum of volatiles from the reaction of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and 3 equivalents of $(\text{Bpin})_2\text{Pz}$ and (d) THF solution IR spectrum of the reaction of $(\text{DIM})\text{Ni}(\text{NO})(\text{O}^{15}\text{NO})$ and 3 equivalents of $(\text{Bpin})_2\text{Bpy}$.



formation of N_2O . Both $(DIM)Ni(^{14}NO)(O^{15}NO)$ and $(DIM)Ni(^{15}NO)(O^{14}NO)$ are isotopically pure as observed by the nitrosyl stretching frequencies and ^{15}N NMR (Fig. 5), and monitoring a solution of either isotopomer in THF shows no isotopic scrambling as judged by ^{15}N NMR and IR spectroscopy (Fig. 5) among the two NO_x sites during the time scale relevant to the reactivity with $(Bpin)_2Pz$. Focusing on one isotopomer, the reaction between $(DIM)Ni(^{14}NO)(O^{15}NO)$ and $(Bpin)_2Pz$ in a 2 : 3 mole ratio provides insight into the mechanism based on the nature of the N_2O produced. Vacuum transfer of the product volatiles to a gas IR cell reveals the presence of all four isotopomers, $(^{15}N)_2O$, $(^{14}N)_2O$, $^{14}N^{15}NO$ and $^{15}N^{14}NO$ (Fig. 5), indicating the N–N bond forming event occurs in a dinickel species. Moreover, the detection of both $^{14}N^{15}NO$ and $^{15}N^{14}NO$ demands an intermediate where both nitrogen atoms are chemically equivalent, which makes $(DIM)Ni(NO)_2$ a plausible precursor.

Isotopomer $(DIM)Ni(^{14}NO)(O^{15}NO)$ was then reacted with $(Bpin)_2Bpy$ which begins to react in the time of mixing. The solution infrared spectrum (Fig. 5) recorded within five minutes of combining the reagents showed all four isotopomers of nitrous oxide. This excludes any mechanism where nitrous oxide comes exclusively from $(DIM)Ni(NO)(ONO)$ nitrite or exclusively from its nitrosyl nitrogen. The presence of ^{15}N in both $N\alpha$ and $N\beta$ of nitrous oxide requires a transition state or intermediate where the two nitrogen atoms are symmetry-equivalent and thus equally likely to become $N\alpha$ or $N\beta$ in a subsequent N–N bond forming step. In addition, seeing both unlabelled and doubly ^{15}N labelled N_2O in this experiment indicates that the N–N bond forming event occurs after formation of a dinickel intermediate because N–N bond formation at a single nickel center would give only singly labelled N_2O . Confirming the above, the solid state IR spectrum of $[(DIM)Ni(NO)]_2$ formed from $(DIM)Ni(^{14}NO)(O^{15}NO)$ and 3 equivalents of $(Bpin)_2Pz$ shows a diagnostic nitrosyl stretches for all three expected isotopomers, $[(DIM)Ni(^{14}NO)]_2$, $[(DIM)Ni(^{15}NO)]_2$, and $(DIM)_2Ni_2(^{14}NO)(^{15}NO)$ (Fig. S17†), consistent with computationally predicted nitrosyl stretches for each isotopomer.

Evidence for outer sphere electron transfer. Closely monitoring the reaction by solution IR (Fig. 6) in THF shows that $(DIM)Ni(NO)(ONO)$ is immediately converted to a new species with a ν_{NO} at 1765 cm^{-1} , distinct from starting material ($\nu_{NO} = 1785\text{ cm}^{-1}$). The lower stretching frequency suggests a more electron rich complex, and the fast reaction suggests electron transfer from $(Bpin)_2Bpy$ to $(DIM)Ni(NO)(ONO)$ to form the anion $[(DIM)Ni(NO)(ONO)]^-$ as the identity of this first intermediate. To test this hypothesis, addition of equimolar decamethylcobaltocene (Cp^*_2Co) to $(DIM)Ni(NO)(ONO)$ results in an immediate color change to dark green and solution IR after stirring for 10 minutes indicates the formation of the same $[(DIM)Ni(NO)(ONO)]^-$ anion as judged by the growth of ν_{NO} at 1765 cm^{-1} (Fig. S40†). To further support this assignment, EPR spectroscopy of a solution of $(DIM)Ni(NO)(ONO)$ after addition of Cp^*_2Co yields an EPR spectrum indicative of $[(DIM)Ni(NO)(ONO)]^-$ where the electron resides in the DIM ligand backbone and couples to both nitrogen atoms and the six $-CH_3$,

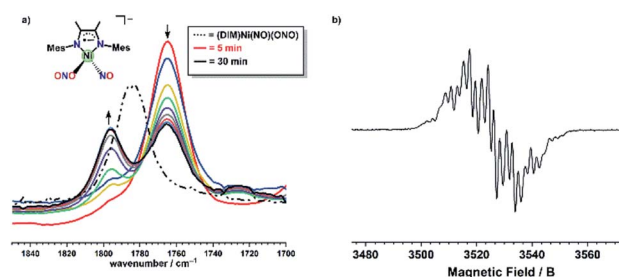


Fig. 6 (a) Solution IR of $(DIM)Ni(NO)(ONO)$ and $(Bpin)_2Bpy$ in a 1 : 1 mole ratio showing complete conversion to the reduced $[(DIM)Ni(NO)(ONO)]^-$ species by the first spectrum collected (red trace), as well as the increase of the new nitrosyl stretch (solid black trace) after 30 minutes. The dotted black trace corresponds to the nitrosyl stretch of unreduced $(DIM)Ni(NO)(ONO)$. (b) X-band EPR spectrum of $[(DIM)Ni(NO)(ONO)]^-$ produced from OSET using $(Bpin)_2Bpy$.

protons (Fig. S39†). The EPR spectrum of $(DIM)Ni(NO)(ONO)$ after addition of equimolar $(Bpin)_2Bpy$ (Fig. 6) is identical to the EPR obtained using Cp^*_2Co which confirms that outer sphere electron transfer from $(Bpin)_2Bpy$ to $(DIM)Ni(NO)(ONO)$ is the first mechanistic step in reductive borylation.

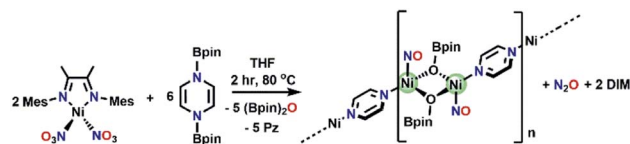
The EPR spectrum obtained after a one electron reduction of $(DIM)Ni(^{15}NO)(O^{15}NO)$ is identical to the EPR spectrum of doubly ^{14}N complex, indicating that the nitrosyl and nitrito nitrogen atoms are not involved in the coupling observed. The DIM ligand-based reduction is corroborated by density functional theory (DFT) electronic structure analysis of $[(DIM)Ni(NO)(ONO)]^-$ where the spin density and corresponding orbital diagram are consistent with a DIM ligand-based SOMO (Fig. S48†). Within 30 minutes of the reaction, there is a clear appearance of a second intermediate with ν_{NO} at 1795 cm^{-1} (Fig. 6). This intermediate is a result of downstream borylation, as this nitrosyl stretch at 1795 cm^{-1} does not appear following electron transfer from Cp^*_2Co .

Candidate intermediates by DFT calculation. After gaining experimental insight into the mechanism of the formation of $[(DIM)Ni(NO)]_2$, we employed density functional theory at the B3LYP-D3BJ/6-311G(d,p) level of theory to evaluate candidate intermediates after single deoxygenation of $(DIM)Ni(NO)(ONO)$, which initiates the reaction. For the intermediate with the chemical formula $(DIM)Ni(N_2O_2)$, we optimized both a nickel dinitrosyl (singlet and triplet) and a nickel hyponitrite (only singlet) (Fig. S47†). The optimized $S = 0$ nickel dinitrosyl was found to be 37 kcal mol^{-1} higher in energy than the nickel hyponitrite and has one linear and one bent nitrosyl ligand. The $S = 1$ dinitrosyl was 7.3 kcal mol^{-1} lower in energy than the singlet, with two bent nitrosyl ligands ($Ni-N-O = 139^\circ$) and pseudo C_2 symmetry. Based on the corresponding orbitals and the spin density plot of $S = 1$ $(DIM)Ni(NO)_2$, there is significant unpaired electron density on the nitrosyl ligands (Fig. S49†). Despite both dinitrosyl species being higher in energy than the calculated hyponitrite, experimental evidence suggests N–N bond formation does not happen at one nickel center. Therefore, we suspect the barrier for intramolecular hyponitrite formation from the $(DIM)Ni(NO)_2$ intermediate is large. With



the high energy of the dinitrosyl species, it is unlikely that (DIM)Ni(NO)₂ would reach a concentration high enough for a bimolecular reaction between two dinitrosyl complexes, and (DIM)Ni(NO)₂ was never observed spectroscopically. Although a transient dinitrosyl species is not a pre-requisite for N–N bond formation, one potential mechanistic pathway that coincides with experimental data is that the N–N bond is formed between transient (DIM)Ni(NO)₂ and (DIM)Ni(NO)(ONO). Within this mechanistic pathway, we postulate that after the bimolecular collision, a series of two more deoxygenation events results in the formation of the [(DIM)Ni(NO)]₂ dimer with liberation of N₂O. While no intermediate bridging hyponitrite species was observed spectroscopically in the reaction, there is literature precedence for N₂O formation from bridging hyponitrite complexes.^{36,64}

Analogous nitrate deoxygenation. Contrasting to the reactivity of (DIM)Ni(NO₂)₂, the analogous (DIM)Ni(NO₃)₂ is unreactive towards (Bpin)₂Pz at room temperature. However, reaction of (DIM)Ni(NO₃)₂ with (Bpin)₂Pz in d₈-THF (mole ratio 1 : 4) at 80 °C for two hours results in a color change from brown to blue. Monitoring by ¹H NMR spectroscopy (Fig. S28†) reveals complete consumption of (Bpin)₂Pz and the paramagnetic (DIM)Ni(NO₃)₂ complex, as well as the growth of O(Bpin)₂ and pyrazine. Upon cooling the NMR sample to room temperature, X-ray quality crystals are present in the J-Young tube. Single crystal XRD of these crystals show that the molecular structure is polymeric (Fig. 7), with a repeat unit Ni₂(NO)₂(OBpin)₂(Pz). Bond lengths within this structure are consistent with unreduced (neutral) pyrazine and the linear Ni–N–O is best described as tetrahedral Ni(0). Each OBpin ligand derived from deoxygenation of nitrate bridges two nickel centers creating a Ni₂(OBpin)₂ diamond core. We postulate that a zero valent nickel nitrosyl complex is a thermodynamic sink and having excess pyrazine in solution facilitates the crystal growth of this polymer. The OBpin ligand shows that borylation using (Bpin)₂Pz can be stepwise and it is not limited to concerted double boryl transfer to one oxygen. The stepwise borylation is supported by the ability of the borylated N-heterocycles to engage in outer sphere electron transfer. There is precedence for OBpin as a ligand including OBpin derived from deoxygenation of CO₂.^{65–70} The solid state IR spectrum (Fig. S29†) of



Scheme 6 Synthesis of pyrazine linked polymer facilitated by N₂O loss.

Ni₂(NO)₂(OBpin)₂(Pz) shows a strong NO stretch at 1810 cm⁻¹. We propose that loss of DIM occurs because its bite angle (~81°) is not ideal for the tetrahedral needs of nickel in competition with pyrazine available as another suitable donor. Free DIM is observed by ¹H NMR spectroscopy (Fig. S28†). Reacting (DIM)Ni(NO₃)₂ with 4 moles of (Bpin)₂Bpy as the deoxygenating agent at 80 °C proceeds analogously, forming Ni₂(NO)₂(OBpin)₂(Bpy) (Fig. 7). The τ₄ values for the nickel in the pyrazine and bipyridine linked polymers vary slightly: 0.73 and 0.67, respectively. The smaller τ₄ value for the bipyridine derivative is accompanied by a smaller nitrosyl angle, with Ni1–N1–O1 being 156.84°, compared to 176.06° for the pyrazine-bridged analogue.

Much like in the synthesis of [(DIM)Ni(NO)]₂, these polymers are deficient in nitrogen with respect to the bis-nitrate starting material. After reacting (DIM)Ni(NO₃)₂ with 4 moles of (Bpin)₂Pz in THF at 80 °C for 2 hours, vacuum transfer of the headspace gas into a gas IR cell shows the characteristic doublet for N₂O at 2224 cm⁻¹ (Fig. S41†), indicating N–N bond formation. With N₂O confirmed, as well as (Bpin)₂O, pyrazine, and free DIM, a balanced reaction for this transformation is presented in Scheme 6. The formation of an N–N bond from the reaction of (DIM)Ni(NO)(ONO) with (Bpin)₂Pz mirrors the N–N bond formation observed in Scheme 3; however, DIM now is replaced by bridging OBpin and bridging pyrazine or bipyridine ligands. From this result, we hypothesize that (DIM)Ni(NO)(ONO) is formed in the synthesis of polymeric Ni₂(NO)₂(OBpin)₂(Pz), and DIM is lost as a result of heating after the nitrite/nitrosyl complex is formed. This is supported by the fact that (DIM)Ni(NO)(ONO) is thermally unstable, and heating in THF for 16 hours results in a color change from brown/green to blue. Monitoring this decomposition by ¹H NMR shows the formation of many new DIM containing species, including free ligand, as judged by numerous new singlets in the aromatic region corresponding to the mesityl meta-H (Fig. S43†). Furthermore, these same polymeric structures can be obtained from the reactivity of (DIM)Ni(NO₂)₂ with either (Bpin)₂Pz or (Bpin)₂Bpy in a 1 : 2 mole ratio at 80 °C, once again showing elevated temperatures induce DIM loss and polymer formation.

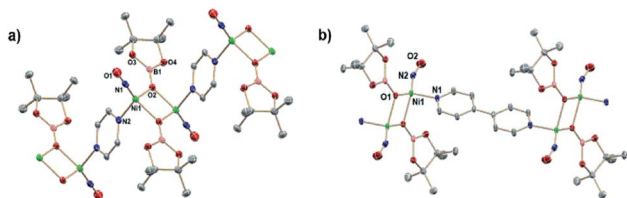


Fig. 7 ORTEP representation of the molecular structures (50% probabilities) of the nonhydrogen atoms of (a) Ni₂(NO)₂(OBpin)₂(Pz) and (b) Ni₂(NO)₂(OBpin)₂(Bpy), showing selected atom labelling (unlabelled atoms are carbon). Selected structural parameters for Ni₂(NO)₂(OBpin)₂(Pz) (Å): Ni1–N1, 1.634(3); Ni1–N2, 2.040(2); Ni1–O2, 1.988(2); N1–O1, 1.173(3); O1–N1–Ni1, 176.1(2); and Ni₂(NO)₂(OBpin)₂(Bpy) (Å): Ni1–N1, 1.9914(17); Ni1–N2, 1.6386(19); Ni1–O1, 1.9960(16); N2–O2, 1.181(3); O2–N2–Ni1, 156.83(18).

Conclusions

We have shown that reductive borylation on both (DIM)Ni(NO₂)₂ and (DIM)Ni(NO₃)₂ at elevated temperatures leads to loss of DIM and nitrous oxide formation, followed by the formation of either pyrazine or 4,4'-bipyridine bridged polymers. With milder conditions, single deoxygenation of (DIM)Ni(NO₂)₂ is facile to yield the mixed oxyanion compound (DIM)



Ni(NO)(ONO). The (DIM)Ni(NO)(ONO) complex undergoes further deoxygenation to form the dimeric [(DIM)Ni(NO)]₂, while also liberating nitrous oxide. Isotopic labelling studies indicate that the N–N bond formation event happens bimolecularly, with a C₂ symmetric intermediate. The dimer is susceptible to oxidation with AgX (X = NO₃[−], NO₂[−], or OTf[−]) to form (DIM)Ni(NO)(X) compounds, and is competent to facilitate nitric oxide disproportionation. While mechanistically complex, we have experimental evidence that the first step in the reaction to form [(DIM)Ni(NO)]₂ is outer sphere electron transfer from the electron rich (Bpin)₂Bpy reductant.

Data availability

The data supporting the findings of this study are available within the paper and its ESI files. Crystallographic data for the crystal structures has been deposited at the CCDC under 2083107 for (DIM)Ni(NO₃)₂, 2083108 for (DIM)Ni(NO₂)₂, 2083111 for (DIM)Ni(NO)(ONO), 2083110 for [(DIM)Ni(NO)]₂, 2083109 for (DIM)Ni(NO)(OTf), 2083112 for (OBpin)₂Ni₂(NO)₂(Pz), and 2083113 for (OBpin)₂Ni₂(NO)₂(Bpy), and can be obtained from <https://ccdc.cam.ac.uk/structures/>.

Author contributions

D. M. B. designed and performed the experiments and computations, interpreted the results and assisted with the manuscript. A. C. C. helped synthesize and characterize the nitrate/nitrite starting materials and the polymers, and assisted with the manuscript. M. P. and V. C. collected and refined the X-ray data. X. G. assisted with the nitrogen NMR experiments. K. G. C. helped design experiments, interpret results, and assisted with the manuscript writing.

Conflicts of interest

There are no conflicts to declare.

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