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# Article

# Real-Time High-Sensitivity Reaction Monitoring of Important Nitrogen-Cycle Synthons by <sup>15</sup>N Hyperpolarized Nuclear Magnetic Resonance

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exchange hyperpolarization of a range of <sup>15</sup>N-containing synthons can be used to enable studies of their reactivity by <sup>15</sup>N nuclear magnetic resonance (NO<sub>2</sub><sup>-</sup> (28% polarization), ND<sub>3</sub> (3%), PhCH<sub>2</sub>NH<sub>2</sub> (5%), NaN<sub>3</sub> (3%), and NO<sub>3</sub><sup>-</sup> (0.1%)). A range of iridium-based spin-polarization transfer catalysts are used, which for NO<sub>2</sub><sup>-</sup> work optimally as an amino-derived carbene-containing complex with a DMAP- $d_2$  coligand. We harness long <sup>15</sup>N spinorder lifetimes to probe in situ reactivity out to 3 ×  $T_1$ . In the case of NO<sub>2</sub><sup>-</sup> ( $T_1$  17.7 s at 9.4 T), we monitor PhNH<sub>2</sub> diazotization in acidic solution. The resulting diazonium salt (<sup>15</sup>N- $T_1$  38 s) forms within 30 s, and its subsequent reaction with NaN<sub>3</sub> leads to the detection of hyperpolarized PhN<sub>3</sub> ( $T_1$  192 s) in a second step via the formation



an identified cyclic pentazole intermediate. The role of PhN<sub>3</sub> and NaN<sub>3</sub> in copper-free click chemistry is exemplified for hyperpolarized triazole ( $T_1 < 10$  s) formation when they react with a strained alkyne. We also demonstrate simple routes to hyperpolarized N<sub>2</sub> in addition to showing how utilization of <sup>15</sup>N-polarized PhCH<sub>2</sub>NH<sub>2</sub> enables the probing of amidation, sulfonamidation, and imine formation. Hyperpolarized ND<sub>3</sub> is used to probe imine and ND<sub>4</sub><sup>+</sup> ( $T_1$  33.6 s) formation. Furthermore, for NO<sub>2</sub><sup>-</sup>, we also demonstrate how the <sup>15</sup>N-magnetic resonance imaging monitoring of biphasic catalysis confirms the successful preparation of an aqueous bolus of hyperpolarized <sup>15</sup>NO<sub>2</sub><sup>-</sup> in seconds with 8% polarization. Hence, we create a versatile tool to probe organic transformations that has significant relevance for the synthesis of future hyperpolarized pharmaceuticals.

# ■ INTRODUCTION

Positron emission tomography (PET) is a very sensitive technique that uses gamma cameras to image changes in metabolic processes, blood flow, and agent absorption in the body. It takes long-lived radionuclides generated using a cyclotron that are then embedded into a suitable receptor to create the radiopharmaceuticals that convey the diagnostic response. Unfortunately, this process can be complex and costly. Magnetic resonance imaging (MRI) is another powerful diagnostic method, but inherent low sensitivity means that routine clinical measurements probe highly abundant water.

Consequently, there has been a great deal of excitement in the clinical community with a method called hyperpolarization that gives MRI the sensitivity needed to visualize changes in metabolic flux by detecting biomolecules that encode disease. The most clinically developed method currently involves the use of dissolution dynamic nuclear polarization (d-DNP).<sup>1-3</sup> One alternative method to create hyperpolarization is *para*hydrogen  $(p-H_2)$ -induced polarization (PHIP), which despite being discovered in the 1980s is only now receiving worldwide attention. Three recent significant PHIP advances that utilize *p*-

 $H_2$  are signal amplification by reversible exchange (SABRE),<sup>4</sup> *p*- $H_2$ -induced polarization with side-arm hydrolysis,<sup>5</sup> and the rapid hyperpolarization and purification of the metabolite fumarate.<sup>6,7</sup> As *p*- $H_2$  can be prepared to a level of 50% purity by simply cooling  $H_2$  gas by liquid nitrogen,<sup>8</sup> one could imagine the widespread future use of this MR sensitization approach.

Here, we demonstrate how it is possible to turn PHIP into a versatile tool for the in situ synthesis of a family of long-lived and highly MR visible precursors containing <sup>15</sup>N, akin to the radionuclides of PET. These reactive intermediates are rapidly embedded into important molecular reporters to illustrate the creation of the hyperpharmaceutical. We achieve this by harnessing reactive species like nitrite ( $NO_2^-$ ), nitrosonium

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(NO<sup>+</sup>), and ammonia (NH<sub>3</sub>), representing simple building blocks which can be transformed into a wide range of hyperpolarized materials.

Our method takes their <sup>15</sup>N isotopologues and uses PHIP to first hyperpolarize them. Thus we create an imbalance in one of the <sup>15</sup>N's two possible nuclear spin orientations (+1/2 or -1/2) which can potentially be maintained for 10's of minutes if stored in an appropriate magnetic field.<sup>9–11</sup> We focus on establishing this concept by reference to nuclear magnetic resonance (NMR), a technique that is used by many scientific disciplines. NMR mainly detects <sup>1</sup>H responses, because the most commonly probed alternative nucleus, <sup>13</sup>C, is 6400 times harder to detect than <sup>1</sup>H. This is due to <sup>13</sup>C's 1% abundance and small gyromagnetic ratio ( $\gamma$ ). Consequently, the Zeeman splitting yielding the resonance frequency is four times smaller than <sup>1</sup>H, and a minute macroscopic nuclear magnetization occurs, which is detected by NMR.

Less utilized <sup>15</sup>N has a highly informative 1350 ppm chemical shift range and long  $T_{1,7}^{9}$  but as <sup>15</sup>N is only 0.36% abundant and has a  $\gamma$  10 times smaller than <sup>1</sup>H, it is 260,000 times harder to detect. Hence, high-concentration samples and extensive signal averaging are needed for NMR studies at natural abundance. Despite this limitation, nitrite ions have been probed by <sup>15</sup>N NMR in solution and the solid state<sup>12</sup> and used to study chemodenitrification in humic substances<sup>13</sup> and nitric oxide release from copper sites, so its utility is established.<sup>14</sup> Importantly, the <sup>15</sup>N isotope can be sourced cheaply in materials like <sup>15</sup>NH<sub>4</sub>Cl and Na<sup>15</sup>NO<sub>2</sub>, and this offers routes to synthesize other isotopically labeled compounds such as pharmaceuticals. Hyperpolarized Na<sup>15</sup>NO<sub>2</sub>, created by *d*-DNP, has been studied.<sup>15</sup>

The sensitivity gains provided by PHIP have already been used widely to aid the study of organic and inorganic chemicals, and it has made the detection of previously hidden intermediates possible.<sup>16–19</sup> Our aim here is to illustrate the hyperpharmaceutical concept while establishing that we can track chemical reactivity, complete the diagnostic fingerprinting of materials, and dramatically expand chemical diversity in the field of hyperpolarization.

We start with nitrite  $(NO_2^{-})$ , a reagent that is formed during the nitrification of ammonia by nitrosomas within the nitrogen cycle.<sup>20</sup> While mammals do not absorb nitrites directly, plants use it to form essential nitrogen-containing molecules such as amino acids and further aerobic oxidation of nitrite leads to nitrate. Nitrite is used as a food additive for cured meats<sup>21</sup> and approximately 7% of our ingested nitrite comes this source, while the remainder comes from the enterosalivary pathway.<sup>3,22</sup> While nitrites are noncarcinogenic, their ability to form nitrosamines can lead to toxicity<sup>23</sup> as examined by the research community and mainstream media.<sup>24,25</sup> The action of methmyogolbin production by nitrite is, however, beneficial in the treatment of cyanide poisoning and sodium nitrite remains as one of the primary antidotes for acute intoxication.<sup>26</sup>

The nitrite ion, usually as sodium nitrite, finds widespread use in the chemical industry, because of its oxidizing properties and role in organic transformations; common examples are the Sandmeyer reaction, which transforms aryl amines into aryl halides, and the diazotization reaction that is used en route to the formation of dyes and pigments.<sup>27</sup>

Nitrite is also an ambidentate ligand that can bind to metals via the N- or O-atoms to form nitro or nitrito complexes, respectively,<sup>28,29</sup> with Ni<sup>30–33</sup> and Pt<sup>34–36</sup> examples being the most prevalent. As the PHIP hyperpolarization method SABRE

works through reversible binding of the agent that is set to become hyperpolarized to a metal complex, we hypothesized that polarization of  $NO_2^-$  via such a route is possible.<sup>4,11,37,38</sup>

In fact, there are a few examples of ionic species such as sodium pyruvate,<sup>39,40</sup> sodium acetate,<sup>41</sup> and naicin<sup>42</sup> that undergo SABRE. This method requires the creation of a scalar coupling network between the target agent and *p*-H<sub>2</sub>-derived protons in a catalyst.<sup>43–46</sup> Hence, an  $\eta^1$ -NO<sub>2</sub> (*N*-nitro) complex with a potentially large hydride-<sup>15</sup>N coupling would be preferred over  $\eta^1$ -ONO (*O*-nitrito) or  $\eta^2$ -O–N–O (*O*,*O*-bidentate) linkage isomers. Theoretical descriptions of SABRE are provided by Barskiy and others<sup>43,44,47</sup> and account for the magnetization transfer conditions needed to sensitize a range of agents.<sup>45</sup> Transfer is optimized at low magnetic fields, typically 6 mT for <sup>1</sup>H, or through *r.f.* excitation at high field.<sup>48</sup> When combined with suitable catalyst lifetimes, this has driven the efficient sensitization of <sup>1</sup>H, <sup>13</sup>C, <sup>15</sup>N, <sup>19</sup>F, <sup>31</sup>P, and <sup>29</sup>Si (etc.)<sup>39-41,49-59</sup> nuclei.

Here, we also evaluate the azide anion, an excellent nucleophile that readily forms organic azides such as the antiretroviral AZT, Avapro, Diova, and Tamiflu. This functionality can be readily reduced to create amines, and through the Curtius rearrangement carbamates. Copper-catalyzed azidealkyne cycloadditions or click reactions are also important. Consequently, azide represents an important precursor to agrochemicals, pharmaceuticals, and natural products so its successful hyperpolarization is also desirable.

Typically, when an iridium *N*-heterocyclic carbene (NHC) catalyst is used, products can be created whose NMR signal strengths are many orders of magnitude higher than those which would be obtained at thermal equilibrium. $^{60,61}$  Warren et al. in particular stand-out for their work on  $^{15}N^{68}$  in a refinement called SABRE-SHEATH,<sup>51,56</sup> and up to 79% <sup>15</sup>N polarization has recently been reported for a range of neutral Lewis bases.<sup>62</sup> Several alternative radio-frequency transfer strategies have also been exemplified, <sup>48,63,64</sup> and given one of the goals of SABRE is in vivo detection, water-soluble SABRE catalysts have been described,<sup>65,66</sup> with the in vitro MRI detection of an <sup>15</sup>N response already illustrated.<sup>66</sup> Tessari et al. have developed a number of analytical science applications for SABRE<sup>67</sup> and other catalyst types have been reported.<sup>68</sup> Furthermore, hyperpolarized long-lived singlet states, as pioneered by Levitt,69 have been created and detected after their formation. 50,70-74 Consequently, we might expect the benefits of such a simple approach to sensitize a range of <sup>15</sup>N-containing reagents to be substantial.

#### RESULTS AND DISCUSSION

**Demonstration That an Active SABRE Catalyst Forms** with Na<sup>15</sup>NO<sub>2</sub>. As indicated, for successful SABRE transfer to occur, the formation of a complex exhibiting spin-spin couplings between the bound substrate and p-H<sub>2</sub>-derived hydride nuclei is required. Classically, this involves the reaction of a precatalyst (most commonly [IrCl(COD)(IMes)] (1) (IMes = 1,3-bis(2,4,6-trimethylphenyl)imidazolylidene)), with an excess of the selected substrate under a H<sub>2</sub> atmosphere. Complexes of type [Ir(H)<sub>2</sub>(IMes)(sub)<sub>3</sub>]Cl, when the substrate is a neutral *N*-heterocycle such as pyridine, meet this requirement.<sup>4</sup> Consequently, our initial efforts targeted the synthesis of an active SABRE catalyst with bound NO<sub>2</sub><sup>-</sup> rather than pyridine.

When Na<sup>15</sup>NO<sub>2</sub> (1 equiv) was added to a solution of [IrCl(COD)(IMes)] (1, 5 mM) in methanol- $d_4$ , the complete

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**Figure 1.** (a) Reaction of [IrCl(COD)(IMes)] with Na<sup>15</sup>NO<sub>2</sub> in the presence of hydrogen and pyridine (inset: structure of the IMes and pyridine ligand). (b) <sup>15</sup>N NMR spectrum of Na<sup>15</sup>NO<sub>2</sub> after SABRE hyperpolarization under 3 bar *p*-H<sub>2</sub> (inset: thermally polarized <sup>15</sup>N NMR spectrum of a 5.0 M solution of <sup>15</sup>NH<sub>4</sub>Cl in D<sub>2</sub>O for comparison).

conversion to  $[Ir(^{15}NO_2)(COD)(IMes)]$  (2) at 298 K (Figure 1a) is observed. This change was readily evident as the <sup>15</sup>N signal for free Na<sup>15</sup>NO<sub>2</sub> at  $\delta_{\rm N}$  611.8 moved to  $\delta_{\rm N}$  490.7 for the bound  $NO_2^-$  at 255 K (see the Supporting Information). When 2 was then exposed to a 3 bar pressure of H<sub>2</sub> at 254 K, the oxidative addition of hydrogen took place to form  $[Ir(H)_2(^{15}NO_2)-$ (COD)(IMes) (3). This complex exhibits <sup>1</sup>H NMR for its hydride ligands at  $\delta_{\rm H}$  –18.77 (hydride *trans* to <sup>15</sup>NO<sub>2</sub><sup>-</sup>, <sup>2</sup>J<sub>HN</sub> = 23.1 Hz and  ${}^{2}J_{\rm HH}$  = 3.3 Hz) and  $\delta_{\rm H}$  –14.17 (hydride *trans* to COD,  ${}^{2}J_{HH}$  = 3.3 Hz). Additionally, a signal for bound  ${}^{15}NO_{2}^{-1}$ appears at  $\delta_N$  476.1. Hence, there is a strong hydride-<sup>15</sup>N coupling in this material that would be commensurate with SABRE. Subsequently, this sample was warmed to 298 K for 20 min. This led to the formation of multiple hydride-containing products, some of which display PHIP on exposure to p-H<sub>2</sub> (see the Supporting Information). Pleasingly, a hyperpolarized signal for free Na<sup>15</sup>NO<sub>2</sub> is observed at 298 K in the <sup>15</sup>N NMR spectrum after SABRE transfer at -5 mG; a field in the mG range will be needed for efficient SABRE transfer.<sup>51,55,56</sup> The resulting <sup>15</sup>N signal enhancement was 134-fold and confirms reversible binding of NO2<sup>-</sup>. Unfortunately, when this sample was left at room temperature for >2 h, SABRE activity was lost because of catalyst decomposition. Hence, we sought to create alternative catalysts that would not only improve the <sup>15</sup>N signal enhancement level but also be suitable for repeated measurement over long periods. Coligands have been used to achieve stability in conjunction with weakly binding substrate,<sup>40,41,73</sup> to reduce spin dilution<sup>42,75,76</sup> and to create hydride ligand chemical, rather than magnetic, inequivalence.<sup>77</sup> Hence, we chose to follow this path as it offers a multitude of benefits to SABRE.

Thus, a sample containing [IrCl(COD)(IMes)] (1, 5 mM), Na<sup>15</sup>NO<sub>2</sub> (5 equiv), and pyridine (3 equiv) was investigated by NMR spectroscopy. The initial formation of [Ir(<sup>15</sup>NO<sub>2</sub>)-(COD)(IMes)] (2) was indicated. Clearly, nitrite outcompetes pyridine for the [Ir(COD)(IMes)]<sup>+</sup> center. Subsequently, exposing this sample to 3 bar H<sub>2</sub> at 254 K led again to the formation of neutral 3. Characterization data are provided in the Supporting Information. We note that known [Ir(H)<sub>2</sub>(IMes)-( $\eta^1$ -COD)(pyridine)<sub>2</sub>]Cl (4<sub>A</sub>) forms alongside 3 in a 1:4 ratio. After warming the sample for 1 h at room temperature, further reaction to form two additional hydride-containing products takes place. Of these, [Ir(H)<sub>2</sub>(<sup>15</sup>NO<sub>2</sub>)(IMes)(pyridine)<sub>2</sub>] (5<sub>A</sub>), with characteristic hydride peaks at  $\delta_H$  –21.24 and –22.45, dominates. The former resonance exhibits a <sup>2</sup>J<sub>NH</sub> splitting of 29 Hz, and both show <sup>2</sup>J<sub>IHH</sub> couplings of –8 Hz.

A further minor product formed in this reaction proved to be  $Na[Ir(H)_2(^{15}NO_2)_2(IMes)(pyridine)]$  (6<sub>A</sub>). It yields hydride resonances at  $\delta_H$  -22.02 (<sup>2</sup> $J_{NH}$  = 29 Hz) and -23.01 with

mutual  ${}^{2}J_{\text{HH}}$  splittings of -7 Hz. Interestingly, no evidence for the formation of *tris* pyridine-containing  $[\text{Ir}(\text{H})_{2}(\text{IMes})-(\text{pyridine})_{3}]\text{Cl}$  was observed<sup>78</sup> and unlike the complexes formed in the absence of pyridine, pyridine-derived  $S_{A}$  and  $6_{A}$  proved stable when left at room temperature for >24 h. Given this stability, these species were suitable probes for rigorous assessment of their SABRE performance. When the ratio of Na<sup>15</sup>NO<sub>2</sub> to pyridine was set to 5:3, the ratio of  $S_{A}$  to  $6_{A}$  in solution proved to be 85:15. Further addition of Na<sup>15</sup>NO<sub>2</sub> (25 equiv), while maintaining the pyridine concentration, only moderately shifted the equilibrium between  $S_{A}$  and  $6_{A}$  to 80:20 thereby confirming that neutral  $[\text{Ir}(\text{H})_{2}({}^{15}\text{NO}_{2})(\text{IMes})-(\text{pyridine})_{2}]$  ( $S_{A}$ ) is the most thermodynamically stable of these products.

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SABRE Assessment of [Ir(H)<sub>2</sub>(<sup>15</sup>NO<sub>2</sub>)(IMes)(A)<sub>2</sub>] (5<sub>A</sub>) and  $Na[Ir(H)_2(^{15}NO_2)_2(IMes)(A)]$  (6<sub>A</sub>) Activity. In order for effective SABRE, the lifetime of the active catalyst must match with the propagating couplings and a level anticrossing condition should be met.<sup>43,44,46</sup> To assess the SABRE performance of  $5_A$  and  $6_{A\nu}$  a series of shake and drop measurements were undertaken using a mu-metal shield to attenuate the Earth's field by a factor of 300 to bring it into the range needed for efficient transfer. These measurements involved first exposing an NMR tube equipped with a J. Youngs Tap containing a solution of [IrCl(COD)(IMes)] (1, 5 mM),  $Na^{15}NO_2$  (5 equiv), and pyridine (4 equiv) in methanol- $d_4$  (0.6 mL) to H<sub>2</sub> (3 bar) for 1 h to form an 85:15 ratio of  $5_A$  to  $6_A$  in solution. Subsequently, the H<sub>2</sub> atmosphere was replaced with *p*- $H_2$  (3 bar), and the sample was shaken for 10 s inside the mumetal shield. After shaking, the sample was transferred into the 9.4 T detection field and an <sup>15</sup>N NMR spectrum was recorded immediately.

Spectral analysis revealed that the free <sup>15</sup>N signal of Na<sup>15</sup>NO<sub>2</sub> was now ~880-fold larger than that of the corresponding thermally polarized NMR spectrum, corresponding to a 0.29% <sup>15</sup>N polarization level (Figure 1b). SABRE transfer to the <sup>15</sup>N of unlabeled pyridine was also observed, and a 172-fold signal gain was quantified for its resonance at  $\delta_N$  301. <sup>15</sup>N NMR signals for coordinated NO<sub>2</sub><sup>-</sup> ligands were also readily visible at  $\delta_{\rm N}$  511.28  $(J_{\rm HN}$  = 29 Hz) for 5<sub>A</sub> and at  $\delta_{\rm N}$  509.7  $(J_{\rm HN}$  = 29 Hz) for the NO<sub>2</sub><sup>-</sup> in the equatorial position and at  $\delta_{\rm N}$  483.7 for the ligand in the axial position of  $6_A$ . Repeating the experiment after polarization transfer at 70 G and subsequently recording a <sup>1</sup>H NMR spectrum revealed that PHIP-enhanced hydride resonances for  $\mathbf{5}_{A}$  and  $\mathbf{6}_{A}$ . SABRE hyperpolarization was also quantified for the <sup>1</sup>H resonances of free pyridine as  $\sim$ 230, 60, and 150-fold for its ortho, meta, and para positions, respectively, after 60 G transfer. No evidence for a PHIP-enhanced hydride resonance for

 $[Ir(H)_2(IMes)(py)_3]Cl$  at  $\delta_H$  –22.7 was observed. We conclude therefore that <sup>15</sup>NO<sub>2</sub><sup>-</sup> sensitization is possible through the action of this coligand-supported catalyst.

Effect of Polarization Transfer Field on the Level of <sup>15</sup>N NMR Signal Gain in Na<sup>15</sup>NO<sub>2</sub>. To improve the levels of signal gain, a more precise polarization transfer field needs to be used. To investigate this effect, a sample containing [IrCl(COD)-(IMes) (1, 5 mM), Na<sup>15</sup>NO<sub>2</sub> (5 equiv), and pyridine (4 equiv) in methanol- $d_4$  (0.6 mL) was exposed to p-H<sub>2</sub> (3 bar), and polarization transfer fields from +10 to -10 mG were deployed; these were created by a solenoid located within an mu-metal shield. A profile of the resulting SABRE enhanced resonance for Na<sup>15</sup>NO<sub>2</sub> is presented in the Supporting Information. The highest signal enhancements were observed when the polarization transfer field was nominally +5 or -3 mG with gains of 1948 and 2054-fold, respectively. More precise probing of the polarization transfer field around these maxima revealed that improvement could be achieved using a -3.5 mG value. At this polarization transfer field, a 2329-fold signal gain was quantified through subsequent measurement at 9.4 T which corresponds to an <sup>15</sup>N polarization level of 0.77%.

Effect of the Coligand on SABRE Catalysis. To form  $5_A$  and  $6_A$ , <sup>15</sup>NO<sub>2</sub><sup>-</sup> must out-bind the stabilizing coligand pyridine. Consequently, we hypothesized that the SABRE processes will be sensitive to the identity of this coligand; such a behavior has been observed previously during the SABRE polarization of sodium pyruvate by sulfoxides,<sup>39</sup> and there are other examples.<sup>41,72,79</sup> Additionally, isotopic labeling of these coligands, which reduces the number of polarization-acceptor spins at the metal center, has proven beneficial while additionally attenuating the effect of relaxation.<sup>42,80,81</sup> A suitable range of coligands were therefore examined to test if it were possible to improve the measured polarization levels in free Na<sup>15</sup>NO<sub>2</sub>, as detailed in Figure 2.

In each case, samples containing [IrCl(COD)(IMes)],  $Na^{15}NO_2$  (25 equiv), and the coligand (A-H, 4 equiv) were prepared and then exposed to 3 bar H<sub>2</sub> at 298 K for 1 h to form the corresponding complexes 6 and 7. Subsequently, a sample was exposed to 3 bar p-H<sub>2</sub>, in a -3.5 mG field, prior to its rapid insertion into the 9.4 T detection field. The second coligand tested was <sup>15</sup>N labeled pyridine (A-<sup>15</sup>N), and this reduced the signal enhancement level for Na<sup>15</sup>NO<sub>2</sub> from 2329-fold to 2107fold. This is likely to reflect the increase in spin dilution associated by increasing the proportion of spin-1/2 nuclei that can accept polarization. The resonance for free <sup>15</sup>N-pyridine at  $\delta_{\rm N}$  301 now exhibits a signal gain of 1558-fold. In contrast, the use of pyridine- $d_5$  (A- $d_5$ ) improves the SABRE hyperpolarization for Na<sup>15</sup>NO<sub>2</sub> as the new enhancement level increases to 3007-fold. As expected, all the pyridine isotopologues yield analogous complexes, 5 to 6, in a common 85:15 ratio. Hence, catalyst speciation is constant and the ca. 30% improvement, compared to undeuterated A, likely reflects both slower relaxation in the active catalyst and reduced polarization.

To further modulate the coligand, other pyridyl derivatives having different steric and electronic properties were examined. Recently reported 2,6-lutidine (**B**)<sup>82,83</sup> was chosen as its *ortho* methyl groups hinder binding to the metal center, a change which might promote ligand loss. When **B** is employed with Na<sup>15</sup>NO<sub>2</sub>, an increase in the SABRE polarization level is indeed observed when compared to pyridine. Interestingly, the ratio of **5**<sub>B</sub> to **6**<sub>B</sub> has changed to 95:5. However, slow activation means that **3** (c.f. Figure 1a) is still visible after 1 h at room temperature; at this stage, it exists in a 1:1 ratio with **5**<sub>B</sub>. Unfortunately, when



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A-15N A-d5

**Figure 2.** Graphical representation of the effect the coligand, **A-H**, has on the resulting SABRE polarization efficiency for Na<sup>15</sup>NO<sub>2</sub> based on the precatalyst [IrCl(COD)(IMes)], Na<sup>15</sup>NO<sub>2</sub> (25 equiv), and coligand (4 equiv) in methanol- $d_4$  at 298 K. The <sup>15</sup>N NMR signal enhancements occur after polarization transfer from 3 bar *p*-H<sub>2</sub> in a -3.5 mG field.

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this sample is left under a 3 bar atmosphere of  $H_2$  for a longer time to allow full activation, sample degradation and the formation of multiple hydride-containing complexes are noted. Hence, **B** fails to provide the stability needed.

The use of electron-deficient methyl  $4,6-d_2$ -nicotinate (C), which has been shown to exhibit <sup>1</sup>H polarization levels of ca. 60% itself under SABRE,<sup>42,60</sup> as a coligand was found to decrease the signal enhancement of Na<sup>15</sup>NO<sub>2</sub> to 1894. The formation of mono-C-substituted  $6_C$  is favored by this change as the ratio of  $S_C$  to  $6_C$  became 1:2. In contrast, electron-rich dimethylamino pyridine (D, DMAP) forms  $S_D$  in a 17:1 ratio with  $6_D$ . Consequently, electron-donating coligands favor the formation of a *bis*-co-ligand substituted complex. Additionally, a significantly improved 10,313-fold <sup>15</sup>N signal enhancement is now observed for Na<sup>15</sup>NO<sub>2</sub> which corresponds to the creation of a 3.4% <sup>15</sup>N polarization level.

Nonheterocyclic ligands can also be utilized for SABRE. As such, amine ligands have been shown to be able to form stable SABRE catalysts and are effective agents for SABRE-relay polarization transfer.<sup>80,81,84–86</sup> When utilized as a coligand for the hyperpolarization of Na<sup>15</sup>NO<sub>2</sub>, benzylamine- $d_7$  (E- $d_7$ ) led to a <sup>15</sup>N signal gain of 2070-fold. The two hydride-containing complexes  $S_E$  and  $6_E$  were formed under these conditions in a ca. 1:1 ratio with hydride resonances at  $\delta_H$  –22.10 and –23.40 and  $\delta_H$  –22.36 and –22.72, respectively. When aniline (F) was used as the coligand, a 3322-fold signal gain for Na<sup>15</sup>NO<sub>2</sub> was quantified. In this sample,  $S_F$  now dominates.

Similarly, sulfoxides have proven to be efficacious for the hyperpolarization of sodium pyruvate and weakly coordinating

substrates.<sup>39,40,79,87</sup> The coligand DMSO- $d_6$  (G) gave a 6270fold signal enhancement for Na<sup>15</sup>NO<sub>2</sub>. Interestingly, while Na[Ir(H)<sub>2</sub>(<sup>15</sup>NO<sub>2</sub>)<sub>2</sub>(IMes)(DMSO- $d_6$ )] (**6**<sub>G</sub>) is now dominant, a second isomer 7<sub>G</sub>, where the two <sup>15</sup>NO<sub>2</sub> ligands lie *cis* to one another and *trans* to hydride is observed. This complex gives rise to a single hydride resonance at  $\delta_{\rm H}$  –22.32 where  $J_{\rm NH$ *cis* $} + J_{\rm NH$ *trans* $}$ is 27.6 Hz. The <sup>15</sup>NO<sub>2</sub> resonance of 7<sub>G</sub> appears at  $\delta_{\rm H}$  502. Isomer **5**<sub>G</sub> is also detected, but now as a minor species, with the ratio of **5**<sub>G</sub>:**6**<sub>G</sub>:**7**<sub>G</sub> in solution being ~1:9:5. Characterization data for these complexes are provided in the Supporting Information. Finally, acetonitrile<sup>88</sup> gave a 2029-fold <sup>15</sup>N signal gain for NO<sub>2</sub><sup>-</sup>. For acetonitrile, the neutral complex [Ir(H)<sub>2</sub>(<sup>15</sup>NO<sub>2</sub>)(IMes)-(acetonitrile)<sub>2</sub>] (**5**<sub>H</sub>), with hydride resonances at  $\delta_{\rm H}$  –22.66 (<sup>2</sup> $J_{\rm NH}$  = 26.7 Hz, <sup>2</sup> $J_{\rm HH}$  = –7 Hz) and –21.77 (<sup>2</sup> $J_{\rm HH}$  = –7 Hz), was the only complex observed. Clearly substantial coligand effects occur, with **D** proving optimal.

Identifying the Optimum DMAP (D):Na<sup>15</sup>NO<sub>2</sub> Ratio. Interestingly, this ligand yielded the highest concentration of isomer 5. We postulated that the concentration of  $S_D$  in solution could be further manipulated by changing the number of equivalents of D in relation to [IrCl(COD)(IMes)] (1) and Na<sup>15</sup>NO<sub>2</sub>. Therefore, a series of samples were prepared with between 3 and 20 equiv of D relative to 1. After activation, they were exposed to 3 bar *p*-H<sub>2</sub> while located in a -3.5 mG polarization transfer field. The resulting signal enhancements at 9.4 T are shown in the Supporting Information.

When three equivalents of D (with respect to iridium) are utilized, a 9086-fold signal enhancement is observed with the corresponding  $\mathbf{5}_{D}$ : $\mathbf{6}_{D}$  ratio being 8:1. Increasing the concentration of DMAP to four equivalents improved the signal gain seen at 9.4 T to 11,019-fold. The ratio of complex  $5_{D}:6_{D}$  also increased to 17:1. Further incremental increases in DMAP concentration, to 6, 8, and 10 equiv, gave signal enhancements of 12,036, 12,079, and 11,888-fold, respectively. The ratio  $5_{\rm D}$ : $6_{\rm D}$ was now 24:1 in all three samples. At higher loadings of D, the formation of  $[Ir(H)_2(IMes)(D)_3]Cl$  is observed, as a single hydride resonance at  $\delta_{\rm H}$  –23.00. Clearly, this catalyst does not transfer hyperpolarization to Na<sup>15</sup>NO<sub>2</sub> and hinders the overall <sup>15</sup>N signal gain because of consumption of *p*-H<sub>2</sub>. Therefore, we conclude that a sensible DMAP level lies between 6 and 10 equivalents with respect to iridium. This creates SABRE beneficial  $5_{\rm D}$  as the dominant species in solution.

**Synthesis and Utilization of DMAP-** $d_2$  **for SABRE.** As stated, deuteration of ligands within the active catalyst can be beneficial.<sup>42,70,76,77,88</sup> We postulated that deuteration of the *ortho* protons in **D**, to give DMAP- $d_2$ , may lead to further improvements in <sup>15</sup>N polarization. Thus, DMAP- $d_2$  was synthesized via H/D exchange from DMAP in D<sub>2</sub>O under microwave irradiation as reported in the literature.<sup>89</sup> Examination of a sample containing [IrCl(COD)(IMes)] (5 mM), DMAP- $d_2$  (6 equiv), and Na<sup>15</sup>NO<sub>2</sub> (25 equiv) in methanol- $d_4$  and exposure to 3 bar *p*-H<sub>2</sub> at a polarization transfer at -3.5 mG led to a signal gain of 13,811 after investigation at 9.4 T (4.56% <sup>15</sup>N polarization level). The corresponding value with <sup>1</sup>H-DMAP was 12,036, and hence introducing the <sup>2</sup>H is beneficial to SABRE.

Effect of NHC Identity on the Efficiency of  $Na^{15}NO_2$ Polarization. Aiming to improve the polarization outcome still further, a study of the effect of the NHC ligand was completed in conjunction with DMAP- $d_2$ . Previously, we have shown how manipulation of the steric and electronic properties of this ancillary ligand can result in improved <sup>1</sup>H, <sup>13</sup>C, and <sup>15</sup>N signal enhancements because of changes in the rate of ligand



**Figure 3.** Effect of NHC on the <sup>15</sup>N NMR signal gain for Na<sup>15</sup>NO<sub>2</sub> with precatalyst [IrCl(COD)(NHC)] (5 mM), DMAP- $d_2$  (6 equiv), and Na<sup>15</sup>NO<sub>2</sub> (25 equiv) in methanol- $d_4$  after polarization transfer at -3.5 mG under 3 bar *p*-H<sub>2</sub>.

exchange.<sup>60</sup> We sequentially increased the steric bulk of the NHC (quantified by the magnitude of %BurV<sup>90,91</sup>) to drive ligand exchange (Figure 3). On moving from the IMes ligand (%BurV = 31.2) to SIMes (32.7), we saw a 14,628-fold signal gain which is a slight improvement from the 13,811-fold signal gain previously observed for IMes. IPr (33.6) and SIPr (35.7) both also led to increased signal enhancements of 15,799 and 17,149-fold. However, the best result was obtained for IPent as a 20,337-fold signal gain which has the highest %BurV of 43.4.

Next our focus turned to the electronic properties of the NHC ligand (Figure 3). As expected, electron-deficient IMes<sup>Cl</sup>, which has chloro substituents on the imidazole ring, reduced the signal enhancement to just 5471. Introducing methyl groups on the imidazole ring showed a minimal effect when compared to IMes (13,336-fold vs 13,811-fold, respectively). However, introduction of a single  $-NMe_2$  group increased the signal enhancement level to 16,427-fold at 9.4 T.

To combine the steric and electronic effects, we utilized the ligand IPr<sup>NMe2</sup>, which has previously proven to be effective for Buchwald–Hartwig amination catalysis,<sup>92,93</sup> to form the precatalyst [IrCl(COD)(IPr<sup>NMe2</sup>)]. This catalyst system gave the highest <sup>15</sup>N signal enhancement for free Na<sup>15</sup>NO<sub>2</sub> seen, 21,967-fold at 9.4 T which is equivalent to 7.2% polarization.

Effect of Na<sup>15</sup>NO<sub>2</sub> Concentration on Signal Enhancement. A series of samples were prepared which contained varying excesses of Na<sup>15</sup>NO<sub>2</sub> relative to 5 mM of [IrCl(COD)-(IPr<sup>NMe2</sup>)] and a constant 30 mM of DMAP- $d_2$ . The corresponding hyperpolarization results are presented in the Supporting Information. Reducing the substrate excess to 10 equiv with respect to iridium increased the <sup>15</sup>N signal enhancement to 36,629-fold, but for just 4 equiv a value of



**Figure 4.** Establishing SABRE hyperpolarization allows the reactivity of <sup>15</sup>N-containing synthons to be assessed. (a) Multistep reaction from Na<sup>15</sup>NO<sub>2</sub> that tracks diazotization with aniline-<sup>15</sup>N and the subsequent formation of two isotopomers of phenyl azide after reaction with  $1^{-15}$ N-NaN<sub>3</sub> in a process that proceeds through a cyclic intermediate; (b) copper-free click reaction of PhN<sub>3</sub> formed as in part (a); (c) formation of N<sub>2</sub>(g); (d) ND<sub>3</sub> quaternization with DCl(aq) and imine formation with benzaldehyde; (e) amidation, sulfonamidation, and imine formation of benzylamine-<sup>15</sup>N; (f) copper-free click reaction of  $1^{-15}$ N-NaN<sub>3</sub>.

62,470-fold was found which is equivalent to 20.6% <sup>15</sup>N polarization. Conversely, increasing the substrate loading to 50 equiv reduced the signal gain to 10,382 (3.17%), albeit with an improved signal to noise ratio. When the concentration of [IrCl(COD)(IPr<sup>NMe2</sup>)] is reduced to 0.25 mM, while maintaining the 1:4 molar ratio with Na<sup>15</sup>NO<sub>2</sub>, the <sup>15</sup>N polarization level increases to 28.42%. This phenomenon is likely the result of increasing the excess of *p*-H<sub>2</sub>, the limiting reagent, relative to this substrate.

**Detection of Unlabeled NaNO**<sub>2</sub>. Given the high signal gains obtained for Na<sup>15</sup>NO<sub>2</sub>, we tested a sample where the <sup>15</sup>N label was present at natural abundance. This sample contained 20 mM of NaNO<sub>2</sub> and was examined with [IrCl(COD)-(IPr<sup>NMe2</sup>)] (5 mM) and DMAP- $d_2$  (6 equiv,). An <sup>15</sup>N signal was easily seen whose signal enhancement was 115,592-fold at 9.4 T; this corresponds to a 38.1% polarization level.

Effect of 15-Crown-5 and Alternative Solvents. Unfortunately, the ionic nature of NaNO2 acts to limit its solubility in the range of organic solvents that are typically employed for SABRE catalysis; it has a moderate solubility in methanol; however, it is sparingly soluble in other primary alcohols and insoluble in most apolar solvents. In an attempt to increase methanol- $d_4$  solubility, the macrocycle 15-crown-5 was added, which has a high chelating affinity for  $Na^{+,94,95}$  SABRE transfer was therefore undertaken on a sample containing [IrCl(COD)(IMes)] (5 mM), Na<sup>15</sup>NO<sub>2</sub> (25 equiv), DMAP (6 equiv), and 15-crown-5 (25 equiv) in methanol- $d_4$ . This led to an <sup>15</sup>N signal enhancement of 12,044-fold at 9.4 T for <sup>15</sup>NO<sub>2</sub><sup>-</sup> which corresponds to 3.97% and reflects a 10% improvement over the analogous measurement with no 15-crown-5. Interestingly, the ratio of  $5_D$  to  $6_D$  in solution was 99:1, as opposed to 91:9 seen in the absence of 15-crown-5. For the optimized catalyst and coligand  $([IrCl(COD)(IPr^{NMe2})]$ (5 mM), Na<sup>15</sup>NO<sub>2</sub> (25 equiv), and DMAP- $d_2$  (6 equiv)), the effect of 15-crown-5 proved to be less pronounced, with the <sup>15</sup>N signal gain improving from 21,967-fold to just 23,114-fold. Hence, solvent effects on this catalysis are substantial and likely to change ligand exchange rates in addition to catalyst speciation.

When an NMR sample containing [IrCl(COD)(IMes)] (5 mM),  $Na^{15}NO_2$  (25 equiv), and D (6 equiv) was prepared in dichloromethane- $d_2$  (0.6 mL), the impact of insolubility of  $Na^{15}NO_2$  was immediately evident. After sonication for 30 min, the sample was exposed to  $H_2$  (3 bar). Investigation by NMR spectroscopy revealed just  $[Ir(H)_2Cl(IMes)(DMAP)_2]$ . However, when an analogous sample was prepared containing 15crown-5 in a 1:1 ratio with Na<sup>15</sup>NO<sub>2</sub>, a different hydridecontaining complex formed. Its hydride resonances appear at  $\delta_{\rm H}$ -22.66 (<sup>2</sup> $J_{\rm HN}$  = 27.5 Hz and <sup>2</sup> $J_{\rm HH}$  = -7 Hz) and  $\delta$  -23.00 (<sup>2</sup> $J_{\rm HH}$ = -7 Hz) and match those of  $5_{D}$ . After SABRE transfer in a -3.5mG field, a 3586-fold signal enhancement was observed at 9.4 T for the free  ${}^{15}NO_2^{-1}$  resonance at  $\delta_N$  618. Warming this sample to 308 K prior to polarization transfer significantly improved the signal gain to 7248-fold and indicates that slow ligand exchange limits the polarization level attained. However, warming further to 323 K yielded no further increase. Using the electron-rich and sterically encumbered precatalyst [IrCl(COD)(IPr<sup>NMe2</sup>)] also yielded improved polarization transfer as an 8149-fold signal gain is seen at 9.4 T. Warming this sample, however, had no benefit. Hence, we have demonstrated how significant polarization levels for  ${}^{15}NO_2^{-}$  result in dichloromethane- $d_2$  if 15crown-5 is present.

Assessment of Na<sup>15</sup>NO<sub>2</sub> Relaxation Rates. DNP hyperpolarized Na<sup>15</sup>NO<sub>2</sub> is reported to have a  $T_1$  of 14.8 s in D<sub>2</sub>O at 5.8 T.<sup>15</sup> We used a low-tip angle approach to assess the  $T_1$  of this SABRE-polarized product at 9.4 T. It was found to be comparable at 16.45 s in the presence of a SABRE catalyst. This value was also determined using an automated hyperpolarization device under reversible flow,<sup>96</sup> after first conducting the SABRE process at -3.5 mG, prior to turning off the  $p-H_2$ supply and holding the sample in a defined magnetic field for a period of time, prior to transfer to 9.4 T to acquire a spectrum. Repeating this process for a number of time points enables the effective low field  $T_1$  value to be calculated. This analysis was undertaken on samples that were stored in the mu-metal shield (ca. 300-fold shielding) or at -3.5 mT. The new  $T_1$  values were 14.9 and 11.2 s, respectively (see the Supporting Information). These values suggest that there will be sufficient time to use the hyperpolarized <sup>15</sup>NO<sub>2</sub><sup>-</sup> resulting from SABRE synthetically to create other hyperpolarized products as  $3 \times T_1$  is available before a signal vanishes. Interestingly, as the  $T_1$  values for <sup>15</sup>N nuclei can dramatically be extended when they are located in an appropriate magnetic field, accessing reaction times of many minutes may be possible.<sup>55,97,98</sup> We are currently exploring this, but here we show how rapid reactions can be evaluated through <sup>15</sup>N NMR at high field is detailed in the following sections.

Conversion of Hyperpolarized Na<sup>15</sup>NO<sub>2</sub> to a Diazonium via NO<sup>+</sup>. The first reaction we consider is the important Sandmeyer reaction that rapidly converts arylamines into arylhalides via a diazonium salt intermediate.<sup>99</sup> Since the first reported example in 1884,<sup>100</sup> it has become a mainstay of organic chemistry and many related reactions have been discovered.<sup>101</sup> Classically, it utilizes either stoichiometric or catalytic amounts of a copper halide, although a number of metal free variants are known.<sup>102–104</sup> The formation of the diazonium salt intermediate proceeds via nitrous acid addition, which is formed in situ from the reaction of NaNO<sub>2</sub> and a strong acid. We sought to follow a diazotization reaction by <sup>15</sup>N hyperpolarized NMR spectroscopy. To do this, we first created a solution of hyperpolarized Na<sup>15</sup>NO<sub>2</sub> using the previously optimized conditions ([IrCl(COD)(IPr<sup>NMe2</sup>)] (5 mM), DMAP (30 mM), Na<sup>15</sup>NO<sub>2</sub> (125 mM) in methanol- $d_4$  (0.6 mL)). A solution of aniline (150 mM) and conc. HCl (100  $\mu$ L) in methanol- $d_4$  (100  $\mu$ L) was then added, and the resulting NMR tube was immediately transferred into the spectrometer and investigated using a  $T_1$ -corrected variable flip angle pulse sequence. It took between 3 and 5 s to start this series of measurements, and the resulting hyperpolarized signals were indicative of nitrous acid (H<sup>15</sup>NO<sub>2</sub>,  $\delta_N$  563) forming phenyl diazonium chloride ( $\delta_N$  314) and ortho-<sup>15</sup>N<sub>2</sub> ( $\delta_N$  308). Their identity was confirmed by their independent synthesis and comparison to literature data.<sup>105</sup> Over the course of 30 s, the response for H<sup>15</sup>NO<sub>2</sub> vanished.

When this process was repeated with aniline-<sup>15</sup>N, the reaction monitoring step revealed the detection of hyperpolarized responses for both of the <sup>15</sup>N centers in the diazo product at  $\delta_N$  315.0 and 232.6 in agreement with the literature (Figure 4a).<sup>106</sup> The hyperpolarization of both of the <sup>15</sup>N sites happens even though aniline itself was not hyperpolarized. Consequently, efficient polarization transfer between them takes place during their time as a coupled spin pair at low field. As a control, we exposed a sample of phenyl diazonium chloride and the catalyst to *p*-H<sub>2</sub> at -3.5 mG and noted no hyperpolarized <sup>15</sup>N resonance result. Consequently, all the hyperpolarized signals seen during Figure S12 of the Supporting Information details how the hyperpolarization level of unlabeled  $NaNO_2$  is sufficient to allow the detection of phenyl diazonium chloride without the need for isotopic labeling.

**Reactions of Hyperpolarized** <sup>15</sup>N<sub>2</sub>-Phenyl Diazonium Chloride. Phenyl diazonium chloride proved to have hyperpolarized  $T_1$  values for <sup>15</sup>N<sub>1</sub> and <sup>15</sup>N<sub>2</sub> of 29.4 and 39.2 s, respectively, at 9.4 T. Additionally, it proved to be relatively stable under these conditions as only limited decomposition to hyperpolarized <sup>15</sup>N<sub>2(g)</sub> ( $\delta_N$  308) was seen. This meant that we could explore the reactivity of this diazonium salt in situ. It is known that such salts liberate N<sub>2</sub> under photochemical or transition-metal-catalyzed processes.<sup>107,108</sup> Under our hyperpolarized regime, addition of CuI saw its rapid conversion into N<sub>2</sub> and consequently a strong signal was seen at  $\delta_N$  308.

A similar hyperpolarized diazonium salt solution was prepared and then treated with NaN3 to examine the formation of phenyl azide. Rapid monitoring enabled the collection of a hyperpolarized <sup>15</sup>N NMR spectrum with strong resonances at  $\delta_{N}$ 242.2 and 90.1 that share a common  ${}^{2}J_{NN}$  of 13.8 Hz because of this species. According to the literature, this reaction could proceed via a cyclic and/or acyclic intermediate, species which would deliver five and three distinct <sup>15</sup>N signals, respectively.  $^{109-111}$  Interestingly, we detect transient signals at  $\delta_{\rm N}$ 356.8 and 298.2 (both with  ${}^{2}J_{NN} = 16.7$  Hz) for the site connected to the  $C_6H_5$  ring which we assign to this product. Upon repeating this study with 1-15N NaN<sub>3</sub>, these two signals gain further complexity and appear alongside one other at  $\delta_N$ 387.3 (d, 17 Hz). These additional features are reflective of the two possible isotopologues that can result from  ${}^{15}N_1 - N_3^{-1}$ addition to form a cyclic intermediate, which place a Ph-<sup>15</sup>N next to two chemically equivalent <sup>15</sup>N groups (a triplet at  $\delta_N$ 298.6 of 17 Hz is seen for it alongside a doublet of 17 Hz at  $\delta_{
m N}$ 356.9) or one (a doublet at  $\delta_N$  298.6 of 17 Hz is now seen) alongside a further triplet at 356.9 of 17 Hz and a doublet at  $\delta_{
m N}$ 387.3 (d 17 Hz) for the next and more remote center (Figure 4a) <sup>15</sup>N of the N<sub>5</sub> ring. Hence, all three unique signals for this cyclic intermediate have been detected. We note that its conversion into phenyl azide  $(Ph-{}^{15}N={}^{15}N^+={}^{15}N^-$  and  $Ph-^{15}N=^{15}N^{+}=N^{-}$ ) proceeds rapidly at 298 K, and the signals for this product also appear,  $\delta_N$  90.3, 242.5, and 232, with apparent  $T_1$  values of 56, 192, and 101 s at 9.4 T, all respectively.

These long  $T_1$  values enable the creation of strongly hyperpolarized phenyl azide. When the reactive alkyne, (1R,8S,9s)-bicyclo[6.1.0]non-4-yn-9-ylmethanol,<sup>112</sup> is added to this hyperpolarized sample in a third synthetic step, further reaction to form the corresponding triazole occurs. Despite the corresponding <sup>15</sup>N signal's  $T_1$  in this product proving to be <10 s, its formation is readily indicated in the associated hyperpolarized <sup>15</sup>N NMR measurements through three signals at  $\delta_N$ 335.4, 255.3, and 252.7 that can be linked through mutual <sup>2</sup> $J_{NN}$ couplings of 12.8 Hz (Figure 4b). As copper-free click chemistry is used widely for bioconjugation with nuclei acids, we expect such measurements to help in the optimization of pharmaceutical preparations and/or in vivo detection.<sup>113,114</sup>

These data have clearly illustrated the successful examination of a multistep reaction as it proved possible to simultaneously see <sup>15</sup>N signals for the phenyl diazonium salt, the pentazole intermediate and phenyl azide (Figure 4a) or the pentazole intermediate, phenyl azide, and the triazole (Figure 4b). We plan

to develop methods to extract precise kinetic data for these changes in the future.

**Conversion of Hyperpolarized Na<sup>15</sup>NO<sub>2</sub> to <sup>15</sup>N<sub>2</sub> through Reaction with <sup>15</sup>NH<sub>4</sub>Cl/HCl.** N<sub>2</sub> gas spontaneously forms from the diazonium salts of primary amines. Consequently, as <sup>15</sup>NH<sub>4</sub>Cl is readily available we monitored its reaction with Na<sup>15</sup>NO<sub>2</sub> and saw strong signals for <sup>15</sup>N<sub>2</sub> in solution (see Figure 4c). We have therefore detailed two facile approaches to hyperpolarized <sup>15</sup>N<sub>2</sub>, we are currently exploring as routes to potentially important *p*-N<sub>2</sub>.<sup>54,74</sup>

Utilization of <sup>15</sup>N Hyperpolarized Azide, Amines, and Ammonia as Probes of Reactivity. The SABRE hyperpolarization of NH3 and amines, such as benzylamine, and their use in SABRE-relay have been extensively reported.<sup>80,81,84-86</sup> Additionally, ammonia and amines have been used as a coligand that leads to improved SABRE catalysis.<sup>62,115</sup> The <sup>15</sup>N polarization of benzylamine-<sup>15</sup>N (E-<sup>15</sup>N) is reported to be ca. 800-fold at 9.4 T.<sup>84</sup> This involves the action of  $[Ir(H)_2(IMes) (E^{-15}N)_3$  Cl in dichloromethane- $d_2$  solution. We restudied this process to improve the SABRE outcome and thereby provide access to a further functional group to demonstrate hyperpolarized reactivity screening. Using the same conditions as previously reported ([IrCl(COD)(IMes)] (5 mM) and benzylamine- ${}^{15}N$  (E- ${}^{15}N$ , 7 equiv)), we determined that optimal SABRE transfer occurs at -4 mG. At this transfer field, a 7751fold signal enhancement was achieved at 9.4 T. As the rate of benzylamine dissociation from  $[Ir(H)_2(IMes)(E-^{15}N)_3]Cl$  in dichloromethane- $d_2$  is slow,<sup>45,85</sup> we found that warming the sample to 308 K further improved the enhancement level to 11,211-fold which corresponds to 3.7% <sup>15</sup>N polarization; it has 14 s  $T_1$  at 9.4 T in the absence of the catalyst and 12.8 s when it is present. We predict that further optimizations could improve this value; however, the resulting signal strengths are sufficient to explore its reactivity. We exemplify now the utilization of hyperpolarized E-<sup>15</sup>N as a synthon for amidation, sulfonamidation, and imine formation. This resulted in the <sup>15</sup>N detection of the products shown in Figure 4e. Their identity was verified by independent synthesis as described in the Supporting Information or by comparison to literature data. In particular, the addition of trifluoroacetic anhydride to hyperpolarized E-<sup>15</sup>N led to the formation and detection of N-benzyltrifluoroacetamide-15N in the resulting 15N NMR spectrum through a signal at  $\delta_N$  116.4. Similarly, triflic anhydride reacted to yield the analogous sulfonamide with a resonance at  $\delta_{\rm N}$  88.6. Finally, addition of benzaldehyde to E-<sup>15</sup>N produced the imine condensation product as evident from a peak at  $\delta_{\rm N}$  327.7.

Ammonia is also widely used in synthetic chemistry and we sought to test whether its reactivity could be probed while hyperpolarized. As gaseous  $^{15}NH_3$  was expensive and difficult to handle, we used an alternative ammonia source.

This involved taking a 1:1 mixture of <sup>15</sup>NH<sub>4</sub>Cl/KO<sup>t</sup>Bu and adding it to 1 in the presence of H<sub>2</sub> with the result that  $[Ir(H)_2(IMes)(^{15}ND_3)_3]$ Cl forms in methanol-d<sub>4</sub>. After polarization transfer at -4 mG, a 3268-fold <sup>15</sup>N signal gain was quantified for the free <sup>15</sup>ND<sub>3</sub> signal. However, over the course of ca. 1 h, the signal enhancement diminishes when the SABRE process is repeated. In contrast, the use of <sup>15</sup>NH<sub>4</sub>OH (available as a 14 molar solution in H<sub>2</sub>O) yielded the same active catalyst, but the sample was now stable for >24 h. The <sup>15</sup>N signal enhancement is also slightly improved to 3765-fold. Changing the NHC ligand proved to have a modest effect on SABRE efficacy (see the Supporting Information), and warming the sample derived from 1 to 308 K improved the signal gain to



**Figure 5.** Establishing the imaging capability of SABRE hyperpolarized <sup>15</sup>N  $O_2^-$  in biocompatible solvents. (a) Setup for hyperpolarized imaging at 7 T of a reaction cell used for the biphasic preparation of <sup>15</sup>N $O_2^-$ ; (b) SABRE hyperpolarized <sup>15</sup>N NMR spectrum of Na<sup>15</sup>N $O_3$  created with  $[Ir(COD)(IMes)(pyridine)]BF_4$ , DMSO- $d_6$  (2 equiv), Na<sup>15</sup>N $O_3$  (25 equiv), and 3 bar *p*-H<sub>2</sub> at 9.4 T; (c) <sup>1</sup>H NMR spectra of the hydride region establish the conversion of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> via the detection of  $Ir(H)(^{15}NO_2)(IMes)(py)_2]$ ; (d) high-resolution three-dimensional axial acquisition using spiral (out) encoding of 6 cm<sup>3</sup> over a 64 × 64 × 8 matrix with a 6 s acquisition time; (e) hybrid MRS(I) data collected using an EPSI pulse sequence and a 64 × 64 matrix (512 spectral points).

4521-fold. However, dramatic improvements are observed with a coligand. While the coligands DMSO- $d_6$ , CD<sub>3</sub>CN, NO<sub>2</sub><sup>-</sup>, and DMAP (see the Supporting Information) were explored, pyridine- $d_5$  proved to give the highest signal gain of 15,145-fold (5.0% polarization). The <sup>15</sup>N  $T_1$  value for <sup>15</sup>ND<sub>3</sub> at 9.4 T proved to be 37 s so there is again a wide time window over which a reaction can be examined. Protonation of <sup>15</sup>ND<sub>3</sub> with DCl in D<sub>2</sub>O led to the detection of hyperpolarized <sup>15</sup>ND<sub>4</sub><sup>+</sup> as a signal  $\delta_N$  15.93 with resolved <sup>15</sup>N-D scalar couplings,  $J_{ND}$ , of 10.8 Hz and a hyperpolarized  $T_1$  of 33.6 s (Figure 4d).

The SABRE hyperpolarization of 1-<sup>15</sup>N NaN<sub>3</sub> itself using the coligand strategy with DMAP and 1 also proved successful. The reaction was found to proceed to form [Ir- $(H)_2(DMAP)_2(IMes)(^{15}N=N=N)$ ] which exhibits hydride signals at  $\delta_H - 23.1 (^2J_{HH} = 8 \text{ Hz})$  and  $\delta_H - 25.0 (^2J_{HH} = 8 \text{ Hz})$  alongside [Ir(H)<sub>2</sub>(DMAP)<sub>3</sub>(IMes)]Cl ( $\delta_H - 22.8$ ). SABRE transfer at -3.5 mG yielded 3.2% hyperpolarization of the N<sub>3</sub><sup>-</sup> signal at  $\delta_N$  95.7. A hyperpolarized solution of NaN<sub>3</sub> was then reacted directly with (1*R*,8*S*,9*s*)-bicyclo[6.1.0]non-4-yn-9-ylmethanol to form the triazole. Under these conditions, a single hyperpolarized <sup>15</sup>N response for the product was visible at  $\delta_N$  321.3 as expected and is shown in Figure 4f.

**Producing Hyperpolarized NO**<sub>2</sub><sup>-</sup> in Water. For biological applications, it is desirable to produce hyperpolarized NO<sub>2</sub><sup>-</sup> in water. Unfortunately,  $S_A$  did not form when the analogous reaction was conducted in this solvent. Preforming  $S_D$ in methanol- $d_4$  prior to removing the solvent and replacing it with D<sub>2</sub>O was also unsuccessful. One further way to achieve an aqueous bolus is to use a biphasic<sup>116</sup> approach with dichloromethane- $d_2$ , which could benefit from the fact that the catalyst is not present in the aqueous layer.<sup>65,117</sup> A sample containing [IrCl(COD)(IMes)] (5 mM), Na<sup>15</sup>NO<sub>2</sub> (25 equiv) and DMAP (6 equiv) and 15-crown-5 (25 equiv) in dichloromethane- $d_2$ (0.3 mL) was prepared and exposed to H<sub>2</sub> (3 bar) to form the active catalyst. D<sub>2</sub>O (0.3 mL) was then added under a nitrogen atmosphere. After SABRE transfer at -3.5 mG and phase separation, two hyperpolarized signals were seen in the corresponding <sup>15</sup>N NMR spectrum for Na<sup>15</sup>NO<sub>2</sub> at  $\delta_N$  618 and  $\delta_{\rm N}$  609. These peaks had relative intensities of 1:70 and were assigned to  $Na^{15}NO_2$  dissolved in the dichloromethane- $d_2$  and D<sub>2</sub>O phases respectively by comparison with data from independent solutions. Assuming that all of the Na<sup>15</sup>NO<sub>2</sub> was present, the D<sub>2</sub>O layer results in a <sup>15</sup>N signal gain of 4667-fold. This will be an underestimate of the actual signal gain. As 15crown-5 can also play a role as a phase-transfer catalyst,<sup>118</sup> a further 25 equiv was added to the sample and this proved to increase the signal enhancement level to 13,794-fold. Further additions of 15-crown-5 did not improve on this; however, warming the sample to 308 K resulted in a 26,327-fold signal gain at 9.4 T. This is equivalent to an 8.69% polarization level in aqueous Na<sup>15</sup>NO<sub>2</sub>. Hence, we have created a simple route to hyperpolarized Na<sup>15</sup>NO<sub>2</sub> in biocompatible water. The level of signal gain compares favorably with the <1% reported with DNP.<sup>15</sup> To confirm the phase distribution of Na<sup>15</sup>NO<sub>2</sub>, a series of <sup>15</sup>N MRI images were recorded on a 10 mm-diameter sample tube as detailed in the Supporting Information. The data in Figure 5 details these results which confirm both separation and the fact that the associated signal strengths are sufficient to allow for high-sensitivity <sup>15</sup>N imaging of  $NO_2^{-1}$ .

SABRE Hyperpolarization of Na<sup>15</sup>NO<sub>3</sub>. In contrast to nitrite, nitrate is usually noncoordinating; however, there are examples of it functioning as a weak monodentate or bidentate ligand.<sup>29,119–123</sup> To further explore SABRE's use as a tool to polarize materials featuring in the nitrogen cycle, we explored the SABRE hyperpolarization of Na<sup>15</sup>NO<sub>3</sub>. As expected, in the absence of a coligand no active SABRE catalyst formed in the reaction between [IrCl(COD)(IMes)] and Na<sup>15</sup>NO<sub>3</sub> under a  $H_2$  atmosphere (3 bar) in methanol- $d_4$ . We screened a number of coligands (DMAP, 2-picoline, DMSO-*d*<sub>6</sub>, DPSO, or CD<sub>3</sub>CN) and saw no evidence for the <sup>15</sup>N polarization of Na<sup>15</sup>NO<sub>3</sub>. In each case, the dominant hydride-containing species in solution was  $[Ir(H)_2(IMes)(coligand)_3]Cl$  or  $[IrCl(H)_2(IMes) (coligand)_2$ ]. However, when the ionic precatalyst [Ir(COD)-(IMes)(pyridine)]BF<sub>4</sub> was used with DMSO- $d_6$  (2 equiv), a 547-fold signal enhancement for the <sup>15</sup>NO<sub>3</sub><sup>-</sup> signal at 9.4 T

(0.18% <sup>15</sup>N polarization) is observed (Figure 5b). No direct evidence for an  $NO_3^-$  containing complex could be found, and therefore, polarization transfer must occur through a very low concentration species.

Unexpected Reduction of Sodium Nitrate. During the course of these investigations, a hyperpolarized <sup>15</sup>N NMR signal at  $\delta_{\rm N}$  511.28 ( $J_{\rm HN}$  = 28.5 Hz) appears over the course of 0.5 h when pyridine alone is used as a coligand. This matches the equatorial NO<sub>2</sub><sup>-</sup> resonance previously observed for [Ir- $(H)_2({}^{15}NO_2)(IMes)(pyridine)_2]$  (5<sub>A</sub>). Relatively strong polarized signals for free pyridine and the pyridine ligand trans to hydride in  $[Ir(H)_2(IMes)(py)_3]Cl$  ( $\delta_N$  299.6 and 255.7, respectively) were also observed in these NMR spectra. As expected, the corresponding <sup>1</sup>H NMR spectrum is dominated by the hydride signal of  $[Ir(H)_2(IMes)(py)_3]Cl$  which appears at  $\delta_{\rm H}$  –22.7, although a weakly PHIP-enhanced signal for 5<sub>A</sub> is visible in this spectrum at  $\delta_{\rm H}$  –21.49 (the peak at  $\delta_{\rm H}$  –22.71 cannot be observed because of overlap). No evidence for  $6_A$  was observed in either the <sup>1</sup>H or <sup>15</sup>N NMR spectra which indicates that  $\mathbf{5}_{A}$  is likely to be the kinetic product of this reaction. After waiting for a further 1 h, refreshing the sample with p-H<sub>2</sub>, and repeating the SABRE process, a polarized signal for free Na<sup>15</sup>NO<sub>2</sub> ( $\delta_{\rm N}$  611.9) could also be detected and the observed signal for  $5_A$  increased in size. We therefore suspect that the reducing environment of this medium converts nitrate to nitrite in a metal-catalyzed reduction. To further probe this reduction, a sample containing [IrCl(COD)(IMes)] (20 mM), pyridine (3 equiv) and  $Na^{15}NO_3$  (25 equiv) was exposed to 3 bar of H<sub>2</sub> at 298 K for 24 h and the growth of the hydride ligand resonance for  $\mathbf{5}_{A}$  at  $\delta_{H}$  –21.49 was monitored by thermally polarized <sup>1</sup>H NMR spectroscopy over the course of 24 h. The resulting integral data for this peak could be fitted to an exponential growth curve (see the Supporting Information). After 24 h and refreshing the  $H_2$  atmosphere, further conversion to  $5_A$  could again be seen which indicates that H<sub>2</sub> is needed to drive this reaction. While the electrochemical reduction of nitrate is widely known<sup>124,125</sup> and limited examples of heterogeneous hydrogenative reduction of nitrate are also reported,  $^{120-128}$  to the best of our knowledge the molecular reduction of nitrate using transition-metal catalysis has not received significant attention. Optimization of the phenomenon reported here may therefore provide a useful alternative.

# CONCLUSIONS

In this work, we have demonstrated how the <sup>15</sup>N hyperpolarization of a range of important <sup>15</sup>N-synthons, including some which feature in the important nitrogen cycle  $(NO_2^{-}(28\%))$ polarization),  $NH_3$  (3%),  $PhCH_2NH_2$  (5%),  $NaN_3$  (3%), and  $NO_3^{-}$  (0.1%)), is possible. When monitored by <sup>15</sup>N NMR, all these species yield strong signals that can be detected readily. The in-field,  $T_1$  values of NO<sub>2</sub><sup>-</sup> (17 s), ND<sub>3</sub> (36 s), PhCH<sub>2</sub>NH<sub>2</sub> (12 s), and NaN<sub>3</sub> (50 s) mean that sufficient time exists to monitor their reactivity through hyperpolarized product responses. This has been demonstrated for the formation of phenyl diazonium, phenyl azide, a triazole, an amide, a sulfonamide, and two imines (Figure 4). In the case of phenyl azide formation, a pentazole intermediate was detected whose cyclic, rather than acyclic, formulation has been confirmed. Studies of the unlabeled formation of phenyl diazonium are also detailed.

Hence, these results demonstrate how SABRE reflects a versatile tool capable of tracking the preparation of a range of nitrogen rich products. We expect the future application of this

approach to aid in achieving the optimized the synthesis of many materials, including important pharmaceuticals.

Furthermore, we demonstrate a biphasic method using a 15crown-5 as a phase-transfer agent that yields >8% aqueous NO<sub>2</sub><sup>-</sup> polarization. We expect this route to help SABRE deliver biocompatible products in the future as we expect it to produce large amounts of such hyperpolarized reagents in seconds. The recent report of 79% <sup>15</sup>N-derived SABRE hyperpolarization<sup>62</sup> suggests that with further optimization a currently unrivaled low-cost approach to rapidly deliver <sup>15</sup>N NMR sensitivity will therefore be obtained. Hence, the pioneering work of Bowers and Weitekamp again continues to expand beyond its original horizons.<sup>129</sup>

#### ASSOCIATED CONTENT

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#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c02619.

Full experimental methods, hyperpolarized NMR spectra, and characterization data (PDF)

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## ABBREVIATIONS

SABRE signal amplification by reversible exchange

p-H2para-hydrogenPHIPparahydrogen induced polarizationd-DNPdissolution dynamic nuclear polarizationPETpositron emission tomographyDMAPdimethylamino pyridine

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