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Direct and Indirect Hyperpolarisation of Aminesusing Parahydrogen

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Wissam Iali, ^a Peter J. Rayner, ^a Adel Alshehri, ^a A. Jonathan. Holmes, ^a Amy J. Ruddlesden ^a and Simon B. Duckett* ^a

Nuclear Magnetic Resonance (NMR) and Magnetic Resonance I magng (MRI) are two widely used techniques for the study of molecules and materiab. Hyperpolarisation methods, such as Signal Amplification By Reversible Exchange (SABRE), turn typically weak magnetic resonance responses into strong signab. In this article we detal how tis possible to hyperpolarise the ¹H, ¹³C and ¹⁵N nuclei of a range of amines. This involved showing how primary a mines form stable but labile complex es of the type [Ir (H) ₂(I Mes)(amine)₃] CI that allow *para* hydrogen to reby its latent polarisation into the amine. By optimising the temperature and *para* hydrogen pressure a 1000-fold per proton NH signal gain for deuterated benzylamine is a chieved at 9.4 T. Additionally, we show that sterically hindered and electron poor amines that bind poorly to iridium can be hyperpolarised by either empbying a co-ligand for complex stab lisation, or harnessing the fact that it is possible to exchange hyperpolarised protons between amines in a mixture, through the recently reported SABRE-RELAY method. These chemical refinements have significant potential to extend the classes of agent that can be hyperpolarised by readily accessible *par* phydrogen.

Introduction

Hyperpolarisation methods are used to overcome the inherent insensitivity of Nuclear Magnetic Resonance (NMR) spect roscopy and Magnetic Resonance I maging (MRI) where their use may lead to dramatic time and cost savings. One such hyperpolarisation method, Para Hydrogen Induced Polarisation (PHIP). produces the required non-Boltzmann nuclear spin distribution by the incorporation of para hydrogen (p-H $_2$), an example of a nuclear singlet, into a suitable substrate molecule. This effect was shown to yield an enhanced NMR signal in 1987^2 and has been the subject of intense investigation. $^{1, 36}$ A drawback of PHIP though, is the requirement for chemical change, caused by p-H2 addition to an unsaturated centre such as an alkene. However, recently a p-H₂ technique that does not change the chemical identity of the sensitised molecule, called Signal Amplification By Reversible Exchange (SABRE), was reported. 7,8 In this process, p-H₂ is not directly incorporated into the substrate. Instead, polarisation is transferred via the J-coupling network that exists within a metal complex that co-locates $p-H_2$ derived hydride ligands and a weakly bound substrate (ligand). 911 Ligand exchange with excess unbound substrate and $p-H_2$ enables the build-up of a pool of polarised substrate molecules in solutionina catalytic fashion as shown in Scheme 1. 12 The SABRE polarisation of ¹H nuclei typical vutilises a ⁴J_{HH} coupling

between the catalysts hydride and substrate ligand protons. Tessari et~al. have quantified these small spin-spin couplings to be $\approx 1.2~{\rm Hz.}^{13}$ Alternatively, stronger $^2J_{\rm HN}$ couplings have now been used to achieve 15 N polarisation transfer at micro-Tesla fields in a variant known as SABRE-SHEATH (SABRE-in shield enables alignment transfer to heteronuclei). $^{14,~15}$ Intramol ecular spin-spin coupling networks within the substrate subsequently enables transfer to remote spins which do not exhibit direct coupling to the hydride ligands. 16

One of the most effective precatalysts for this process is [IrCl(COD)(I Mes)] (1) [where I Mes = 1, 3-bis (2,4,6-tri methyl phenyl)i midazol -2-ylidene, COD = cis,cis-1,5-cycl ooctadiene] which, after reaction with H_2 and an excess of substrate, typically forms $[Ir(H)_2(IMes)(substrate)_3]$ Cl in protic solvent ssuch as methanol. ¹⁷ Neutral active catalysts of the type $[Ir(H)_2(CI)(IMes)(substrate)_2]$ have also been reported to achieve similar results. ¹⁸ These metal based polarisation transfer catalysts have been shown to act on a range of substrate sthat contain multiple bonds to nitrogen, such as nicotinami de, ^{19,20} i soniazid, ^{21,22} metronidazole, ²³ pyr azole, ²⁴ imines, ²⁵ diazirines ²⁶ and ni triles, ²⁷ and I ead to pol arised ¹H, ¹³C, ¹⁵N, ¹⁹F, ²⁹Si, ³¹P, and ¹²⁹Sn nuclei that yield substantially enhanced NMR responses in just a few seconds. ^{19,283} In fact, ¹H polarisations of 50% have been reported, while for ¹⁵N, values of over 20% have been achieved.

While SABRE-induced polarisation can also be achieved using in-field rf. transfer methods, 3437 whose efficiency varies with pulse sequence, 3739 spontaneous polarisation transfer occurs readily at low-field and it is this method we employ here. Moreover, as predicted, 9 it has also been established that SABRE can be used to produce hyperpolarised singlet states 40 with long-lifetimes through transfer in ultra-low field,

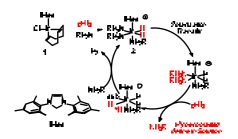
Electronic Supplementary Information (ESI) available: Full experimental procedures, characterisation data and example spectra. See DOI: 10.10.39/x0xx000000x

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or after the implementation of rf. transfer. ⁴¹⁻⁴⁶ Hence the diversity of applications found for this approach is growing and it clearly reflects not only a successful medium to test hyperpolar isation concepts but a potential route to transform the analytical potential of NMR. ⁴⁷⁻⁵⁰



Sche me 1: Route to SABRE hyperpolarisation of an amine, NH 2 R

In this article, we introduce a new class of substrate into the SABRE repertoire, the amine. This is achieved by the formation of iridium-amine complexes of type [Ir(H)₂(IMes)(RNH₂)₂]Cl (2. Scheme 1), whose kinetic behaviour is determined. Whilst the synthesis and use of iridium-amine complexes has been reported for catalytic transformations such as hydrogenation, ^{51:53} we use them here for polarisation transfer catalysis. We have recently shown a limited number of amines are amenable to SABRE.⁵⁴ Here, we start by detailing the hyperpolarisation of ammonia and benzylamine ($BnNH_2$) and its associated optimisation to achieve large NMR signal enhancements. We then show how hyperpolarisation can be achieved in a range of primary amines. Upon changing to steri call y bul ky primary ami nes, secondary amines or aromatic amines, we show that an active SABRE catalyst does not for m upon reaction with 1. However, we exemplify co-ligand and relayed polarisation transfer protocols to overcome this limitation and hence expand further the range of amines amenable to polarisation by $p-H_2$.

Results and Discussion

Direct ¹HHyperpolarisation of Ammonia and BnNH₂ by SABRE

Our objective was to investigate the efficiency of the SABRE polarization of amines and ammonia and to determine their ligand exchange dynamics. A 5 mM solution of ${\bf 1}$ in dry dichloromethane- d_2 containing an \approx 6-fold excess of NH $_3$ relative to ${\bf 1}$ at 298 K was therefore prepared. The aprotic solvent ensures that we maintain the necessary J-coupling network in $[Ir(H)_2(IMes)(NH_3)_3]CI$ ($({\bf 2-NH_3})$) during the study, as

rapid ^2H exchange results to form ND $_3$ in deuterated protic solvents. This complex yields a hydride signal at δ –23.8, alongside a broad response at δ 0.47 for free NH $_3$. The corresponding equatorial and axial NH $_3$ ligand ^1H NMR signals of **2-NH_3** appear at δ 2.19 and 2.88 respectively. 2D $^1\text{H}^{-15}\text{N}$ HMQC measurements were subsequently used to locate the corresponding 15 N signals for these ligands at $\delta_{a\text{xial}}$ –47.8 and δ_{equ} –35.5. Full characterisation data for **2-NH** $_3$ is available in the ESI. 54 EXSY met hooks were then used to probe NH $_3$ and H $_2$ loss in **2-NH** $_3$. At 298 K, the associated rate constant for NH $_3$ loss proved to be 1.64 s 3 while that of H $_2$ loss is 0.32 s 3 . For comparison, the dissociation rate for pyridine in [Ir (H) $_2$ (IMes)(py) $_3$]Cl is 13.2 s 4 and suggests a higher stability for **2-NH** $_3$ which agrees with the greater basicity of NH $_3$ relative to pyridine. 55

As $\mathbf{2\text{-}NH_3}$ undergoes both NH_3 and H_2 loss in solution, we sought to prove that it underwent SABRE catalysis. Thus, a 3 bar pres sure of p-H₂ was introduced at 298 K and polarisation transfer was conducted at 60 G. A ¹H NMR spectrum at 9.4 T was then recorded which showed a 154-fold signal enhancement per proton for the free NH₃ response while the corresponding equatorial ligand signal , at δ 2.19, showed a 77fold enhanced response (Figure 1). Hence 2-NH3 acts as a SABRE catalyst as it produces a hyperpolarised free ammonia response. In the presence of water, the observed signal enhancement of the protons in free NH3 decreased to 40-fold per proton, matching that now observed for the equatorially bound NH₃ ligand This drop is reflected in the signal at δ 1.88, for what is a H₂O response, exhibiting a 75-fold signal gain per proton due to concomitant proton exchange; the ratio of 2- $NH_3: H_2O: NH_3$ in this sample was 1:5:17.5. Under these conditions, the T_1 value for free NH_3 in the presence of the active SABRE catalyst was measured by inversion recovery to

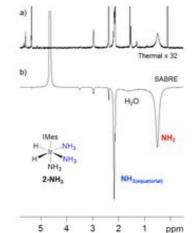


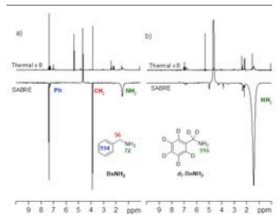
Figure 1: (a) Thet her maly po b rised 1 HNMR S ABRE s pect 1 u m(x 32 ve it cale xpansion) recorded of 2 -NH $_3$ (for med byre action of 1 with NH $_3$ and H $_2$)in dichloro metha re- 0 2 at 298 K. (b)Th ecorres ponding SABRE polarised 3 4.7 4 HNMR spectrum after transfer

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under $p-H_2$ at 60G. Thehy perpolarised responses off ree NH_3 and $I-NH_3$ (eq ua 0.66) of $2-NH_2$ and residual H_3O are indicated.

The SABRE-induced hyperpolarisation of benzylamine (BnNH₂) was also investigated. As ample containing 1 (5 mM) and $BnNH_2$ (10 eq.) in dichloromethane- d_2 solution was exposed to 3 bar of H2. The immediate formation of [Ir(H), (IMes)(BnNH,),] d (2-BnNH,) was observed. It gives a characteristic hydride resonance in the $^1\!H$ NMR spectrum at $\,\delta$ -23.97. Full characterisation data for this product is available in the ESI. Interestingly, the ¹H NMR spectrum of **2-BnNH**₂ showed that the $BnNH_2$ ligand that lies $\mbox{\it tr}$ ans to hydride, yields inequivalent responses for its NH, protons at δ 4.92 and 2.30, and CH₂ protons at δ 3.60 and 3.18. This is due to hindered rotation around the Ir-N bond which results in an up/down distinction for the resonances of the equatorial ligand. In contrast, the axial ligand yields single responses which are equival ent at δ 4.24 (NH $_{\!2})$ and δ 3.83 (CH $_{\!2})$ due to free rotation on the NMR timescale about the Ir-N bond. The corresponding EXSY-derived rate constant for equatorial BnNH₃loss from 2-**BnNH₂** was $3.33 \,\text{s}^{-1}$ while the rate of H₂ loss was $2.83 \,\text{s}^{-1}$ at 298 K. Hence the rate of BnNH₂ loss is higher than that of NH₃ loss in $2-NH_3$. This difference is due to NH_3 forming a st ronger Ir-N bond as reflected in their relative pK, values and suggests that it might perform better under SABRE that NH₃.

This was examined by $p-H_2$ -based polarisation transfer at 60 G which resulted in hyperpolarised free BnNH $_2$ in solution. The signal enhancements were quantified to be 72- (NH $_2$), 56-(CH $_2$) and 194-fold (Ph) per proton as shown in Figure 2a. However, by using d_7 -BnNH $_2$ instead we were able to focus the SABRE polarisation into the two amino protons alone and this led to an improved signal enhancement of 916-fold per proton (Figure 2b).



Figur e 2: (a) ¹HNMR spectra of BnNH2, thermally polarised, to p, and hyper polarise d, bot tom. (b) ¹HNMR spectra for d > BnNH2, thermally polarised, top, and hyperpolarised, bot tom.

In or der to investigate the T_1 contribution to this effect we determined values for BnNH $_2$ and d_7 -BnNH $_2$ at 9.4 T. BnNH $_2$

proved to have effective T_1 values of 1.1 s (NH $_2$) and 4.7 s (CH $_2$) respectively while it s 2 H-labelled variant exhibited a similar 1.1 s T_1 value for the amino group in the presence of the active catalyst. Hence, the improved NH signal gain seen with d_7 -BnNH $_2$ is due to a reduction in spin dilution which leads to more efficient SABRE transfer. The relaxation rates for BnNH $_2$ and d_7 -BnNH $_2$ are both slower in the absence of the active SABRE catalyst in agreement with earlier reports that the catalyst plays a role in reducing relaxation times due to reversible binding. Consequently, BnNH $_2$ now shows T_1 values of 9.0 s (NH $_2$) and 11.0 s (CH $_2$), whereas d_7 -BnNH $_2$ has a T_1 value of 10.1 s for its NH $_2$ group.

Effect of Catalyst to Substrate Ratio on SABRE Polarisation

Previous studies have shown that the SABRE effect is dependent upon the catalyst to substrate ratio as a consequence of ki netic and relaxation effects. Therefore, we studied the effect of changing the ratio of BnNH₂ relative to 1 from 4-fold to 20-fold in a series of further experiments, under taking the associated SABRE transfer studies at 60 G and 298 K. It was found that similar total polarisation levels result within experimental error during these experiments (see ESI). Hence, we conclude that the observed signal enhancements under these conditions are essentially independent of ligand excess which suggests that slow exchange and fast relaxation within the catalyst restrict the maximum polarisation level.

Effect of p-H₂ Pressure on SABRE Polarisation of BnNH₂

As SABRE derives its polarisation from p-H₂, it could be the limiting reagent in this catalytic process and therefore affect the observed substrate polarisation level. ¹⁹ Up until this point, we have been utilising 3 bar pressure of p-H₂ which reflects an ca. 6-fold excess when compared to the 50 mM substrate present in a 5 mm NMRt ube. A sample containing 1 (5 mM), $BnNH_2$ (50 mM, 10 eq.) in dichloromethane- d_2 solution was therefore prepared and exposed to between 2 and 4 bar of p-H.. The resulting signal gains, after polarisation transfer at 60 G, are shown in Figure S14 (see ESI) and a strong dependence on p-H₂ pressure is seen. This is consistent with the fact that H₂ exchange takes place after ligand dissociation and the remaining equatorially bound BnNH₂ ligand will experience a higher level of lat ent p-H₂ polarisation (see Scheme 1). When d_7 -BnNH₂ is examined with 4 bar of p-H₂, the NH signal gain increases to 1079-fold per proton from the 916-fold signal gain achieved with 3 bar.

Effect of Temperature on SABRE Polarisation of BnNH $_{\mathrm{2}}$

The temperature at which SABRE is conducted is also known to affect the efficiency of the polar isation transfer due to changes in the lifetime of the SABRE-active catalyst. We found here that cooling a dichloromethane- d_2 solution containing 1, BnNH_2 and 3 bar $p\text{-H}_2$ to 288 K results in a reduction in the level of signal enhancement when compared to 298 K data (Figure S15, ESI). Conversel y308 K gave an improved response with the overall polarisation level increasing by $^\sim$ 40 %. This fits with the observed rate constant for BnNH_2 dissociation increasing to 9.85 s $^{-1}$ from the 3.33 s $^{-1}$ value at 298 K. We

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therefore conclude the retained polarisation level in BnNH₂ is improved by the faster rate of substrate dissociation and shorter catalyst lifetime. For NH₃, a 251-fold ¹H signal gain per NH proton is observed at 308 K when compared to the 154fold value at 298 K. This is consistent with the increase in the NH_3 dissociation rate constant to 10.42 s^{4} at 308 K when compared to 1.64 s⁻¹ at 298 K.

SABRE Transfer to ¹³C and ¹⁵N

SABRE-induced hyperpolarisation of ¹³C was also observed for BnNH_a. Whilst polarisation transfer into the ortho phenyl carbon was readily observed using a standard ^{13}C acquisition sequence after polarisation transfer 60 G under 4 bar p-H₂, the other $^{13}\mathrm{C}$ resonances had poor signal-to-noise ratios. We overcame this by using a $^{1}\mathrm{H}$ - $^{13}\mathrm{C}$ refocussed INEPT experiment that gave rise to a spectrum showing all 5 carbon envir onments after polarisation transfer at 60 G. We utilised long-range J-H-C-couplings to transfer this polarisation. 13C signal gains of up to 65-fold were achieved using this method (Figure 3a). We further note that there is a very strong 13 C si gnal polari sati on transfer field dependence on the BnNH₃ intensities which is consistent with earlier reports on

When ${\rm Bn^{15}NH_2}$ is used instead of BnNH2, the detection of a hyperpolarised $^{15}{\rm N}$ response is readily evident as shown in Figur e 3b. The ¹⁵N signal gain for the free material in solution proved to be ~880-fold after polarisation transfer at 60 G and 308 K. The equatorially bound 15 N resonance at δ -5.59, is 4 times larger than the free amine signal. As the ratio of free ami ne to equatorially bound Bn 15NH 2 in solution is actually 7: 2, the rate of Bn¹⁵NH, loss must be relatively slow, even at 308 K. Under this 60 G condition, polarisation transfer is likely to occur via the ${}^{3}J_{HH}$ coupling between the Bn $^{15}NH_{2}$ and the hydride ligands. To investigate the effect of using a $^2J_{\rm HN}$ coupling we repeated this measurement after polarization transfer within a μ -metal shield (ca. 350-fold shielding). Under these SABRE-SHEATH type conditions, 14, 15 an ~800-fold 15 Nsignal gain was observed and further optimisation may therefore be needed to maximise this response. The corresponding ¹H signal gains with this ¹⁵N labelled material after transfer at 60 G were now 33- (NH2), 34- (CH2) and 52fold (Ph). These comparet o the analogous values of 72-, 56and 192-fold respectively with Bn 14NH 2. Interestingly, the 1H polari sati on levels therefor e decrease with 15N addition and we propose that this is an example of spin dilution.

Expanding the Substrate Range

In order to test the generality of amine polarisation via SABRE. we prepared a series of samples containing 1 (5 mM) and 10 eq. of the substrates shown in Figure 4 in dichloromethane-d2 solution. These substrates include a number of primary amines and each is successfully hyperpolarised after transfer at 60 G upon reaction with 1 and p-H, In fact. SABRE polarisation of phenylethylamine (PEA) and phenylpropylamine (PPA) results strong signal enhancements and transfer is found to proceed across the corresponding C_2 and C_3 carbon chains into their phenyl rings. For PEA we found that the NH2

gain is actually increased to 108-fold per proton compared to the 72-fold BnNH, value, and that the CH, CH, bridge gave 50fold (NCH₂) and 45-fold (CH₂) enhancement sper proton. The 5proton containing phenyl group gave a 92-fold gain per proton. Spin-isolation of the phenyl group, by introducing an ether linkage, as in phenoxyethylamine (POEA) resulted in signal enhancements of 99- (NH₂), 47- (NCH₂), 147- (CH₂O) and as expected. just 8-fold (Ph) per proton for our test sample. We therefore conclude that polarisation transfer across the oxygenlinker is inefficient at 60 G and a stronger aliphatic proton response results. The amines isobut vlamine, all vlamine and tryptamine were also studied as shown in Figure 4. In all cases, the formation of [Ir(H), (IMes)(amine),]CI was indicated (see ESI) and polarisation transfer results.

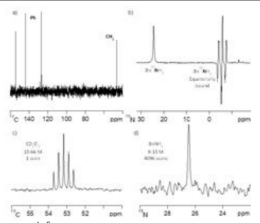
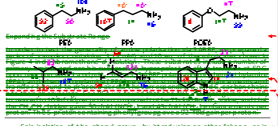


Figure 3. (a) 1 H- 18 Cre focuss ed INEPTNMR spectrum of hyperpolarised B nNH₂ (35 mM) achieved via 2-Bn NH₂(5 m M)u nder SABREin dichlor om ethane -d₂ solution aft er tra nsfer at 60 Ga rd 308K; (b) 15NNMRs pec trum o f 15N lab elled Bn 15NH2 (35 m M) aft er SABRE trans fer via 2-8 rN H ₂(5 m M) at 60G and 3 08 K which gives rise t o hyper polarised resonances for free (δ 24.42) and equatorially bound(δ -5.59) substrate: (c) Single scant her maly polarised 13C NMR spectrum in CD₂C_b (15.66 M) and (d) 4 096 scan thermaly polar ised ¹⁵NNMR spectrum of BnNH ₂(9.15 M).



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Figure 4: Amine substrates polarised by SABRE using precatalyst 1 in dichlor ome thane $-d_2$ so Lition. Per proton signal gains are given for the indicated $^1\mathrm{H}$ sites (*average ac ross two sites due to peak overlap) observed at 9.4 T. Corresponding $^1\mathrm{H}$ NMR spectra for thermally pobrised and SABRE polarised experiments a regiven in the ESI.

When secondary amines, such as dibenzylamine, were examined, no evidence for the formation of an active SABRE catalyst was observed. A similar result was observed for sterically hindered primary amines, such as isopropylamine and aromatic amines, such as aniline. Sterically demanding substrates, such as 2, 6-lutidine, have been previously shown to be unable to be polarised using SABRE. SA full list of the amines probed in this study is available in the ESI. We therefore postulate that sterically demanding or electron deficient amines fail to activate and form the necessary [Ir(H)₂(IMes)(amine)₃]CI SABRE catalyst.

This problem could be overcome for aniline by the addition of the co-ligand 1-methyl 1, 2,3-t riaz ole (mtz) or CH₃CN. For the corresponding sample containing 1 (5m M), aniline (10 eq.) and mtz (3 eq.) in dichloromethane- d_2 we achieved signal enhancements of 51-fold for the NH₂ group and 17-fold for the phenyl group, per proton. These signal gains are summarised in Figure 5. When CH₃CN (8 eq.) is used instead of mtz, the polarisation levels increase to 306- (NH₂) and 193-fold (Ph) per proton. The active complex in this SABRE process was characterised as [Ir(H)₂(IMes)(aniline)₂(CH₃CN)] Cl and yields a distinctive hydride resonance at δ –24.78 (see ESI). Utilis at ion of such a co-substrate strategy was however unsuccessful for the secondary amines as detailed in the ESI.



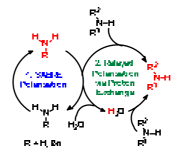
Figur e 5: 1 HNMR signalga 'n s perp rotonobs er ved f ort heindicated aniline res orances when hyp erpolarised by SABR Eint he presence of the described co-lgandat 9.4 T .

Indi rect Hyperpol ari sati on of Am ines by SABRE-REL AY

As expected, substrate binding to the metal centre is needed for polarisation transfer to occur. We hypothes ised that these amines might also be hyperpolarised indirectly. In this scenario, hyperpolarisation of a primary amine or ammonia is achieved and subsequent proton exchange, which may be

mediated by residual water, allows for a polarised proton to be shuttled into the non-SABRE-active amine. Subsequent intrasubstrate polarisation transfer then relays the signal gain more widely in this agent.

In order to test this hypothesis, a series of samples containing 1 (5 mM), target amine (10 eq.) and NH₃ (3-5 eq.) were prepared in dichlor omethane- d_2 solution. **2-NH**₃ for med in all cases as confirmed by the presence of a hydride resonance in the corresponding 1H NMR spectra at δ -23.8. Polarisation transfer was then conducted at 60 G, and the resulting signal gains that were observed at 9.4 T are presented in Figure 6.



Scheme 2: SABRE-RELAY polar sation of amines. (1) SABRE pobrisation of an inter med brytransferagert, in this casea primaryamine or ammonia. (2) Pobrisation is then rebyed into the target amine via proton exchange either directly or via residual water present in the sample.

For is opropylamine (PrNH₂), the SABRE-RELAY polarised NH₂ signal showed a 220-fold signal gain while 27- and 150-fold enhancements were seen for the CH and CH3 resonances respectively. This reflects a breakthrough as PrNH2 was unable to be directly polarised by SABRE due to its steric bulk preventing adequate binding. Dibenz vlamine (Bn₂NH) was also successfully polar ised using this method, and yi elds ¹H signal gains of 274-(NH), 200-(CH₂) and 395-fold (Ph) per proton. Additionally, a ¹³C spectrum can be acquired in a single scan on these materials after polarisation transfer at 60 G such that a 475-fold signal gain for the CH₂ resonance is observed. Full NMR spectra are available in the ESI. Furthermore, the aromati camine, ani li ne, now exhi bits a 150-fold NH2 proton signal enhancement and a 9-fold signal gain for the phenyl ring under analogous conditions. We note that these signal gains are lower than than those seen when CH₃CN is used as a coligand to achieve direct SABRE transfer as detailed in Figure 5. We suggest that this difference in behaviour arises because a 60 G polarisation transfer field is no-longer optimal for intramolecular polarisation transfer after proton exchange. This is clearly is not the case for transfer via directly bound aniline and the complexes scalar coupling network which is in fact commonly maximised for $^{1}\mathrm{H}$ transfer at 60 G.

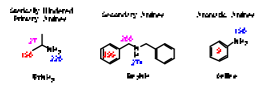
From these results we can conclude that the SABRE-RELAY effect is able to polarise sterically hindered primary amines,

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secondary amines and aromatic amines that are not themselves accessible to SABRE. Thus, the scope of amine polarisation is vastly increased.



Figur e 6: 1 HNMR signalga in s observed per proton f ort heindicated aminere sonance s when hyp erpolar ised by SA BRE-R ELAY us n g $\,\textbf{2-NH}_{\textbf{3}}\,\text{at}\,9\,.4\,\text{T}.$

Conclusions

In summary, we have shown here how SABRE can be used to hyperpolarise a series of primary amines. This substrate extension opens up the SABRE approach to operate with a much wider range of analytes than was previously thought possible, as we extend beyond the original aromatic Nheterocycles, imines and nitriles. Activity is achieved by the formation of a series of complexes of the form [Ir(H)₂(IMes)(amine)₃]Cl. Relaxation studies, in conjunction with ligand dissociation rate measurements were used to demonstrate that the high relative stability of these complexes acts to limit the degree of SABRE signal gain. This hypothesis is consistent with the fact that increasing the p-H₂ pressure or reaction temperature leads to improved signal gains. Therefore, significant catalyst optimisation will be important if very high levels of hyperpolaris ation are to be achieved by this route in the future.

Nonetheless, in the case of BnNH₂, ¹H NMR signal enhancement values of ~100-fold per NH proton were achieved for benzylamine using [IrCl(COD)(IMes)]. Consequently, when d_7 -benzylamine was used, the resulting focusing of the hyperpolarisation into the NH, resonance resulted in a 900-fold signal enhancement per proton at 9.4 T with a p-H₂ pressure of 3 bar. This value reduced to 33-fold for Bn¹⁵NH₃ after transfer at 60 G. Hence, we predict that further improvements can be made through a more detailed study of the effect of isotopic labelling. We have also demonstrated transfer to ¹³C and ¹⁵N with diagnostic NMR spectra being collected at a 35 mM concentration in a single 1. _____S. B. Duckett and R. E. Mewis, Accounts of Chemical scan. We predict that application of high-field SABRE transfer techniques, 3437, 39 such as the LIGHT-SABRE 38 approach, might subsequently enable this process to work inside the magnet. but not e that a rigorous study of the effect the polarisation transfer field plays on the resulting signal enhancement levels is justified.

In the course of these studies we found that sterically hindered primary amines, secondary amines and aromatic amines were unable to form an active SABRE catalyst of the $type \quad [\,Ir(\,H)_{\,2}\!(I\,Mes)\,(a\,mi\,ne)_{\,3}]\,CI. \quad Thi\,s \quad meant \quad that \quad di\,rect$ polari sati on transfer vi a such a complex was not possible. We found for aniline that the addition of a co-ligand such as CH₃CN overcame this problem via the formation

[Ir (H)₂(IMes)(ani line)₂(CH₂CN)]Cl s uch that si gnal enhancements of up to 306-fold per NH proton could be achieved.

An indirect route was described to over come this limitation more generally, such that hindered primary amines, secondary amines and ar omaticamines can be hyperpolarised by SABRE-Now, a SABRE-hyperpolarised intermediary, such as ammonia, is able to readily transfer polarisation into agents such as isopropylamine, benzylamine and aniline via either direct proton exchange or mediated by residual water present in the sample. This approach expands the range of amines that can be hyperpolarised without changing their chemical identity through interactions with p-H₃.

Given the increase in signal intensity that is observed for the amines in this study, we are now working towards their use as agents for mechanistic study ⁵⁸⁶⁴ in transfer hydrogenation, ^{65,66} hydroamination, ^{67,68} and vitally important N₂ fixation reactions. ⁶⁹⁻⁷¹ Additionally, since phenylethylamine is a naturally occurring monoamine based alkaloid that acts as a promoter of cat echolamine (dopamine and norepinephrine) release in plants and animals we expect these observations to be of wide interest. 72, 73 Furthermore, the SABRE-RELAY method⁵⁴ has recently been shown to offer a route to hyperpolarise an even larger range of hydrogen transfer acceptors using OH functional groups. Optimisation of the intermediaries NH polarisation level reflects a key part to optimi sati on of this technique and hence these results will be of interest to any potential developer.

Conflicts of interest

The authors declare no conflicts of interest.

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