UNIVERSITY of York

This is a repository copy of A Simple Route to Strong Carbon-13 NMR Signals Detectable for Several Minutes.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/117890/</u>

Version: Accepted Version

Article:

Roy, Soumya S. orcid.org/0000-0002-9193-9712, Norcott, Philip orcid.org/0000-0003-4082-2079, Rayner, Peter J. orcid.org/0000-0002-6577-4117 et al. (2 more authors) (2017) A Simple Route to Strong Carbon-13 NMR Signals Detectable for Several Minutes. Chemistry : A European Journal. pp. 10496-10500. ISSN 1521-3765

https://doi.org/10.1002/chem.201702767

Reuse Other licence.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/

Illustration of a simple route to generate strong carbon-13 NMR signals that are detectable for several minutes

Soumya S. Roy^a, Philip Norcott^a, Peter J. Rayner^a, Gary G. R. Green^b and Simon B. Duckett^a

Abstract: Nuclear Magnetic Resonance (NMR) and Magnetic Resonance Imaging (MRI) suffer from low sensitivity and limited nuclear spin memory lifetimes. While hyperpolarization techniques increase sensitivity there is also a desire to increase relaxation times to expand the range of applications addressable by these methods. Here we demonstrate a route to create hyperpolarized magnetization in ¹³C nuclear spin pairs that last much longer than normal lifetimes by storage in a singlet state. By combining molecular design and low-field storage with para-hydrogen derived hyperpolarization we achieve more than three-orders of signal amplification relative to equilibrium Zeeman polarization and an order of magnitude extension in state lifetime. These studies use a range of specifically synthesized pyridazine derivatives and dimethyl p-tolyl phenyl pyridazine is the most successful, achieving a lifetime of ca 190 s in low-field which leads to a ¹³C-signal that is visible for 10 minutes.

While carbon is one of the most abundant elements in nature, its nuclear magnetic resonance (NMR) active form carbon-13 is present at just a ~1.1% level which, when coupled with its low magnetogyric ratio, results in low detectability. Consequently, ¹³C magnetic resonance imaging (MRI) produces a negligible response when compared to proton measurement in the body which is facile due to high water content and high sensitivity. ¹³C detection does, however, benefit from potentially long relaxation times when compared to those of the proton.

A number of methods, commonly known as hyperpolarization, exist that can increase NMR sensitivity in nuclei such as ¹³C and are being used to overcome these issues.^[1-2] These approaches artificially increase the associated spin population differences between the energy levels that are probed. For example, Golman et al. reported a para-hydrogen (p-H₂) induced nuclear polarization (PHIP)^[3-4] study which achieved the rapid in vivo detection of a ¹³C-MRI response in 2001.^[5] Two years later, they described the results of a similar study using dissolution dynamic nuclear polarization (DNP),^[6] where a normally inaccessible response was seen in vivo. Bhattacharya et al. have since incorporated p-H₂ into sodium 1-¹³C acetylene dicarboxylate to facilitate the collection of an arterial ¹³C-MRI image of a rat brain.^[7] More recently, a DNP derived ¹³C-MRI response with chemical shift resolution has been shown to distinguish different metabolic flux between normal and tumor cells in humans.^[8-11] These studies illustrate

[a] Dr. S. S. Roy, Dr. P. Norcott, Dr. P. J. Rayner, Prof. Dr. S. B. Duckett Department of Chemistry, University of York Heslington, York, YO10 5DD, UK E-mail: simon.duckett@york.ac.uk

[b] Prof. Dr. G. G. R. Green York Neuroimaging Centre, The Biocentre, York Science Park Innovation Way, Heslington, York, YO10 5NY, UK

Supporting information for this article is given via a link at the end of the document.

the potential benefits to human health if such methods were to become widely accessible and hence establish the need for a rapid, low cost delivery method for long-lived ¹³C hyperpolarization.

In this article, we demonstrate that the goal of rapidly producing a long-lived ¹³C hyperpolarized response can be met by applying the signal amplification by reversible exchange (SABRE) effect.^[12-14] In SABRE, a catalyst reversibly binds *p*-H₂ and the substrate to transfer dormant spin order from p-H2 into the substrate via the scalar-coupling framework, as shown in Scheme 1. We use this approach here to hyperpolarize a series of coupled ¹³C spin-pairs in a range of pyridazine derivatives, a motif that exhibits pharmacological activity.^[15-16] Polarization is then stored in specially created singlet spin order to enable a response to be seen several minutes later. While a range of nicotinamide and pyridazine based substrates have been shown to deliver long-lived ¹H hyperpolarization, [17-18] and analogous ¹⁵N based singlets have been created by Warren and coworkers^[19] we believe the ¹³C responses reported here are significant due to the growing use of ¹³C-MRI for *in vivo* study.



Scheme 1 Schematic depiction of the SABRE hyperpolarization technique. IMes = 1,3-bis(2,4,6-trimethylphenyl)imidazol-2-ylidene.

The term singlet $(|S_0\rangle = (|\alpha\beta\rangle - |\beta\alpha\rangle)/\sqrt{2})$ that is used here represents the spin-zero magnetic alignment of a coupled spin-1/2 system whose conversion into the associated triplet states $(|T_0\rangle = (|\alpha\beta\rangle + |\beta\alpha\rangle)/\sqrt{2}$; $|T_1\rangle = |\alpha\alpha\rangle$; $|T_{-1}\rangle = |\beta\beta\rangle$) is symmetry-forbidden. Consequently, any population difference that can be created between these singlet and triplet forms is expected to relax more slowly than the usual time constant T_1 .^[20] The symmetry properties that make such states long-lived also make them challenging to generate and probe.[20-21] Levitt and co-workers have demonstrated a number of strategies to do this in a range of chemically inequivalent spin systems^[22-26] and have achieved a lifetime of over 1 hour in an optimized chemical system at low-field.^[27] However, when a substantial chemical shift difference exists between these spin-pairs, the application of a spin-lock, or sample-shuttling to low field, is necessary to extend state lifetime.^[22, 28-29] This effect has recently been illustrated by monitoring the effect of solvent dependent chemical shift changes.[17-18] Warren and co-workers have

COMMUNICATION

reported a parallel approach that exploits magnetic inequivalence to create related singlet states.^[21, 30-33] Hence while SABRE has been shown to create hyperpolarized ¹H and ¹⁵N derived singlets there is a need to expand these methods to ¹³C given the success of DNP.^[8-11] However, ¹³C-SABRE itself has though currently seen limited application^[34] and reported efficiency gains are relatively low. We develop here a molecular design strategy for use with SABRE and radio frequency (*rt*) excitation to achieve greater than 2% net ¹³C polarization in a long-lived form.

In this study, we employ magnetic and chemical inequivalence effects through the synthesis of specific substrates where their carbon-4 and carbon-5 sites are ¹³C enriched as detailed in Scheme 2 (full synthetic strategy and characterization data are available in the supporting information, Section S1-3). The *Type-1* form agents exhibit chemically equivalent but magnetic inequivalent ¹³C spin-pairs ($\Delta \delta = 0$) and have local C_2 symmetry. The *Type-2a* form is constructed such that R¹ and R² are chemically different and a small chemical shift difference results between the ¹³C spin-pair ($\Delta \delta \neq 0$). Chemical inequivalence is also derived by remote substitution, at R² and R³, in the *Type-2b* agents of Scheme 2. Whilst these synthetic strategies allow access to two distinct classes of molecular system, our results illustrate that both are equally viable.



Scheme 2 The molecular systems studied here are of *Type-1*, which reflect a chemically equivalent but magnetically distinct ¹³C spin-pair (\bullet), or *Type-2a* and *Type-2b*, which reflect chemically inequivalent ¹³C spin-pairs ($R^1 \neq R^2 \neq R^3$).

In order to explore the singlet states of these systems their NMR properties must first be analyzed. The Type-1 substrate, 1, of Table 1 reflects an AA'XX' type spin system (Figure 1a) and produces the ¹³C NMR spectrum shown in Figure 1b. This trace illustrates the effect of magnetic inequivalence, but does not immediately yield the individual carbon-proton couplings $(^{2}J_{CH})$ and ${}^{3}J_{CH}$) necessary to create a singlet state by the method of Warren^[21] as the peak-to-peak separations reflect the mean value of the ¹³C-¹H *J*-couplings (5.25 Hz = $[^{2}J_{CH} + {}^{3}J_{CH}]/2$). By employing a *J*-synchronized experiment,^[23, 30, 32] it is possible to show the difference in these J-couplings is 3.1 Hz (see Section S5 in the supporting information). We harness this difference in coupling (ΔJ_{CH}) to populate the singlet state via rf pulse sequencing as detailed in Figure 1c. Table 1 details the chemical structures of Type-1 agents 1-3 that are examined here. A value of zero for ΔJ_{CH} means that it is not possible to induce interconversion between the singlet and triplets forms via rf pulsing (e.g. agent 3, see Section S5).[20]

We also prepared agents **4-8** which reflect a series of *Type-2* molecular systems. Their spin system is illustrated in Figure 1d, while Figure 1e shows the ¹³C NMR spectrum of agent **5** in methanol-*d*₄. In this case, the partially resolved 1.05 Hz ($\Delta\delta^2\nu^2/2J_{CC}$) splitting signifies that a strongly coupled ¹³C spin-pair results when R¹ and R² are deuterated phenyl and *para*-tolyl groups respectively.



Figure 1 a) Spin topology of the *Type-1* agent **1**, showing the *J*-couplings that exist between the ¹H and ¹³C nuclei, where R^1 = deuterated phenyl group; (b) corresponding ¹³C NMR spectrum of agent **1** in methanol-*d*₄; (c) M2S-S2M pulse sequence used here; (d) spin topology of *Type-2* substrate **5** and corresponding ¹³C NMR spectrum in methanol-*d*₄(e).

The pulse sequence that is used to create and examine the lifetime of the singlet state in these *Type-1* and *Type-2* molecules consists of three parts as detailed in Figure 1c. Part I converts longitudinal magnetization into singlet order (M2S), part II preserves this singlet order and part III converts it back into a visible form. The first and last steps are realized experimentally by a train of n 180° pulses that are separated by delay (τ) which is a molecule specific parameter. For the *Type-1* system, **1** where J_{CC} >> J_{HH} , the following equations provide τ and n.^[30, 32]

$$\tau = \frac{1}{(2\sqrt{(J_{CC} + J_{HH})^2 + (\Delta J_{CH})^2})};$$

$$n_1 = \frac{\pi}{(2\tan^{-1}[\Delta J_{CH}/(J_{CC} + J_{HH})])}; n_2 \simeq n_1/2$$

In contrast, in the case of the *Type-2* spin systems (agents **4-8**) these parameters come from the equations shown below.

$$\tau = \frac{1}{(4\sqrt{J^2 + \Delta\delta^2\nu^2})}; n_1 = \frac{\pi}{(2\tan^{-1}[\Delta\delta, \nu/J])}; n_2 \simeq n_1/2$$

Section S7 in the supporting information details these values for **1-8**. The resulting singlet states were then stored either in high field or in low field (after sample transfer). For **1**, the singlet state lifetimes (T_S) were measured to be 75 ± 5.5 s and 115 ± 12 s at high and low field respectively. We therefore see an ~10-fold increase over the 9.4 T T_1 relaxation time of 9.7 s. The effect of a spin-lock during high-field storage proved to be minimal, increasing the T_S by only ~10%. In the case of agent **5**

COMMUNICATION

we achieved a $T_{\rm S}$ of 90 ± 3 s in high field which increases to 165 ± 18 s in low-field. Table 1 summarizes these values for agents 1-8 and confirms that this strategy allows the creation of long-lived singlet states in these molecules. ²H-labeled 7 contained the optimal molecular environment of the series, delivering a low-field $T_{\rm S}$ of 186 ± 18 s.

Table 1. ¹³C (**O**) SABRE signal enhancement (ε) over the corresponding thermal measurement at 9.4 T after transfer at the indicated field (G), net polarization (P, %) and T_1 and T_S lifetimes (s) of substrate **1-8** in high field (HF: 9.4 T) and low field (LF: ~10 mT). The *J*-coupling between the ¹³C spin-pair was found to be ~58.5 \pm 2.0 Hz in all cases. The ΔJ_{CH} values for Type-1 substrates, and the chemical shift difference ($\Delta \nu$) for Type-2 substrates are noted.

Α	Substrate structure	Enhancement	Lifetime (s)	$\varDelta J_{CH}^{\dagger}$
g		(ε) , transfer		or Av@
n		polarization		диш 9.4 Т
t		level P (%)		(Hz)
		ε: 2500 ± 300	$T_1: 9.7 \pm 0.3$	3.1 ±
		@30 G	$T_{S(HF)}:75\pm5.5$	0.2
		P ≈ 2.0	$T_{S(LF)}$: 115 ± 12	
2		ε: 1600 ± 280	$T_1: 12.4 \pm 0.9$	²J _{CD} ∼
2		@150 G	T _{S(HF/LF)} : -	0.4
		P ≈ 1.3		
•		ε: 600 ± 50	T ₁ : 16.0 ± 1.5	0
3		@20 mG	$T_{S(HF/LF)}:No$	
		P≈0.5	acess	
4		ε: 1600 ± 300 @150 G	$T_1: 10.2 \pm 0.6$	11.0 ±
			$T_{S(HF)}:22\pm3.0$	0.1
		P ≈ 1.3	$T_{S(LF)}: 28 \pm 6.5$	
5		ε: 550 ± 50 @5 mG	T₁: 15.5 ± 1.2	10.4 ±
J		@3 mG	$T_{S(HF)}$: 90 ± 3.0	0.1
		P ≈ 0.45	T _{S(LF)} : 165 ± 18	
6		ε: 350 ± 40	$T_1: 10.4 \pm 0.3$	14.5 ±
		@10 mG	T _{S(HF)} : 115±5.5	0.4
		P≈0.35	T _{S(LF)} : 148±20	
		ε: 600 ± 50	T ₁ : 15.2 ± 0.3	4.4 ±
7		@1 mG	T _{S(HF)} : 145±6.0	0.3
		P≈0.50	T _{S(LF)} : 186 ± 18	
		ε: 800 ± 150	T ₁ : 7.5 ± 0.5	78.8 ±
8		@10 mG	T _{S(HF)} : < 5	0.5
		P≈0.65	$T_{S(LF)}: 45 \pm 6.0$	

For **2**, the ¹³C-²H couplings are too small to exploit the M2S sequence to prepare the singlet. For **4**, the singlet state lifetime proved low due to the ¹³C-deuterium coupling which provides a route to scalar relaxation.^[35] In **8**, the chemical shift difference between the ¹³C pairs is similar to the *J*-coupling constant in high field and a low lifetime results but in low field this extends to 45 s. In contrast, agents **5**, **6** and **7** operate well in both low and high field, exhibiting lifetimes in excess of 150 s in low field.

A series of SABRE experiments were then undertaken to see if it was possible to create hyperpolarized longitudinal spin

order within the ¹³C manifold of agents 1-8 (Table 1). This involved taking methanol-d₄ solutions that contained 20 mM of the substrate, and 5 mM of the IMes catalyst. p-H₂ gas was bubbled through the solution for 20 s in low field and the sample transferred into the NMR spectrometer for further analysis. Figure 2 highlights the results of this process, with the level of ¹³C polarization reaching ~2% as compared to the corresponding thermal polarization of only 0.0008 % at 9.4 T in the case of agent 1 after relayed transfer from ¹H-¹³C at 30 G (see Section S6). No H/D-exchange is observable on the timescale of the SABRE experiment. The relayed transfer process was then examined as a function of the magnetic field experienced by the sample, and three maxima were observed, at ~10 mG (using µ-metal shield), ~30 G, and ~100 G. Simulation revealed the ~10 mG maxima is associated with direct hydride-carbon spin-spin transfer via the ${}^{4}J_{H}{}^{13}{}_{C}$ and ${}^{5}J_{H}{}^{1}{}_{C}$ ¹³_C couplings in the catalyst. The remaining maxima appear to result from relayed transfer via the agents ¹H response (see Section S4).

When agent 2 is examined the ²H labels should prevent the relayed response operating and restrict transfer to the ~10 mG field range. Under these conditions a strong ^{13}C signal is seen. However, upon moving from 10 G to 150 G, ^{13}C and ^{1}H SABRE enhanced signals are observed in the ¹H and ¹³C frequency ranges. These results reveal readily detectable contributions from the ²H-¹H isotopolog, which is present at 1%, through the observation of a 13 C response that contains a $J_{H^{-13}C}^{1}$ splitting of 5.4 Hz. This reflects one of the challenges faced when work with hyperpolarization in so far as low-concentration species can be readily detected. Agents 3 and 5-8 also require direct polarization transfer as there is no suitable relayed transfer pathway and again work well between 1 mG and 20 mG. These ¹³C hyperpolarization data are summarized in Table 1 (and Section S5). Polarization levels approaching 2% are readily achieved, which would be expected to increase further through catalyst optimization.[36] We then transferred the resulting 13Chyperpolarization into singlet order via the methods described earlier. The efficiency of singlet conversion in all successful cases was found to be in the range of 50-80 %.



Figure 2. ¹³C NMR spectra of **1** after (a) SABRE at a mixing field 5 mG and corresponding thermally equilibrated signal of 1000 transients. (c) Similar SABRE studies of **7** at a mixing field of 1mG and (d) its thermal equilibrium spectra acquired by 600 transients.

Figure 3 shows the decay of the resulting hyperpolarized ¹³C singlet derived signals for agents **1**, **5-8** as a function of their storage time ($T_{\rm S}$) in low field. The ¹³C lifetimes proved to be directly comparable to those measured without hyperpolarization and signals can be readily observed for several minutes after

creation when stored in a low-field region. In the case of 7, hyperpolarized signals were detectable for well over 10 mins.



Figure 3. Hyperpolarized ¹³C singlet state decay (log10 scale) as a function of low-field storage time (τ_{delav}) for agents 1, 5-8. Results are summarized in Table 1

In summary, we have demonstrated that a series of novel agents can be prepared which contain two adjacent ¹³C labels in addition to two nitrogen based lone pairs which make them suitable for SABRE. Despite the weak J-coupling that exists between the hydride ligands and the targeted ¹³C sites we achieve a hyperpolarized response at the 2% level. This hyperpolarization has then been efficiently converted into singlet spin order within the two ¹³C labels by rf excitation with a lowfield relaxation time of ~190 s being the result for deuterated dimethyl p-tolyl phenyl pyridazine. This process has been exemplified for both magnetic and chemical inequivalence. Our method provides a fast and low-cost technique to create ¹³C hyperpolarization in a reversible fashion with very little waste. Because of the simplicity of this approach we envisage that this strategy will be adopted more widely to hyperpolarize related tracers. We are currently seeking to improve on the purity of these states in order to test the in vivo detection of these agents.

Acknowledgements

We thank the Wellcome Trust (092506 and 098335) for funding. We are grateful to discussions with Prof. H. Perry.

Keywords: NMR spectroscopy • hyperpolarization • long-lived singlet states • para-hydrogen • structure elucidation

- J. H. Lee, Y. Okuno, S. Cavagnero, J. Magn. Reson. 2014, 241, [1] 18-31.
- J.-H. Ardenkjaer-Larsen, et al., Angew. Chem. Int. Ed. 2015, 54, [2] 9162-9185
- [3] J. Natterer, J. Bargon, Prog. Nucl. Mag. Res. Sp. 1997, 31, 293-315
- C. R. Bowers, D. P. Weitekamp, Phys. Rev. Lett. 1986, 57, 2645-[4] 2648
- K. Golman, O. Axelsson, H. Johannesson, S. Mansson, C. [5] Olofsson, J. S. Petersson, Magn. Reson. Med. 2001, 46, 1-5.
- K. Golman, J. H. Ardenaer-Larsen, J. S. Petersson, S. Mansson, I. Leunbach, *Proc. Natl. Acad. Sci. U.S.A.* 2003, 100, 10435-10439. [6]
- P. Bhattacharya, E. Y. Chekmenev, W. H. Perman, K. C. Harris, A. P. Lin, V. A. Norton, C. T. Tan, B. D. Ross, D. P. Weitekamp, J. [7] Magn. Reson. 2007, 186, 150-155.
- S. J. Nelson, et al., Science Translational Medicine 2013, 5. [8]

- K. Golman, R. in't Zandt, M. Lerche, R. Pehrson, J. H. Ardenkjaer-Larsen, Cancer Res. 2006, 66, 10855-10860.
- K. Golman, R. in't Zandt, M. Thaning, Proc. Natl. Acad. Sci. U.S.A. [10] 2006, 103, 11270-11275.
- [11] S. E. Day, M. I. Kettunen, F. A. Gallagher, D.-E. Hu, M. Lerche, J. Wolber, K. Golman, J. H. Ardenkjaer-Larsen, K. M. Brindle, Nat. Med. 2007, 13, 1382-1387.
- R. W. Adams, J. A. Aguilar, K. D. Atkinson, M. J. Cowley, P. I. P. [12] Elliott, S. B. Duckett, G. G. R. Green, I. G. Khazal, J. Lopez-Serrano, D. C. Williamson, *Science* **2009**, *323*, 1708-1711.
- [13] N. Eshuis, B. J. A. van Weerdenburg, M. C. Feiters, F. P. J. T. Rutjes, S. S. Wijmenga, M. Tessari, *Angew. Chem. Int. Ed.* **2015**, 54, 1481-1484.
- A. N. Pravdivtsev, A. V. Yurkovskaya, H.-M. Vieth, K. L. Ivanov, J. Phys. Chem. B 2015, 119, 13619-13629. [14]
- G. Heinisch, H. Frank, Prog Med Chem 1990, 27, 1-49.
 M. Asif, Curr. Med. Chem. 2012, 19, 2984-2991. [15]
- [16]

[9]

[34]

[35]

[36]

- [17] S. S. Roy, P. J. Rayner, P. Norcott, G. G. R. Green, S. B. Duckett, Phys. Chem. Chem. Phys. 2016, 18, 24905-24911
- [18] S. S. Roy, P. Norcott, P. J. Rayner, G. G. Green, S. B. Duckett, Angew. Chem. Int. Ed. 2016, 55, 15642-15645. [19]
 - T. Theis, et al., Sci. Adv. 2016, 2, e1501438.
- M. H. Levitt, in *Annu. Rev. Phys. Chem., Vol. 63* (Eds.: M. A. Johnson, T. J. Martinez), **2012**, pp. 89-105. W. S. Warren, E. Jenista, R. T. Branca, X. Chen, *Science* **2009**, [20]
- [21] 323. 1711-1714.
- M. Carravetta, M. H. Levitt, J. Am. Chem. Soc. 2004, 126, 6228-[22] 6229.
- M. C. D. Tayler, M. H. Levitt, Phys. Chem. Chem. Phys. 2011, 13, [23] 5556-5560.
- M. C. D. Tayler, I. Marco-Rius, M. I. Kettunen, K. M. Brindle, M. H. Levitt, G. Pileio, *J. Am. Chem. Soc.* 2012, *134*, 7668-7671. [24]
- G. Pileio, J. T. Hill-Cousins, S. Mitchell, I. Kuprov, L. J. Brown, R. [25] C. D. Brown, M. H. Levitt, J. Am. Chem. Soc. 2012, 134, 17494-17497.
- [26] G. Stevanato, S. S. Roy, J. Hill-Cousins, I. Kuprov, L. J. Brown, R. C. D. Brown, G. Pileio, M. H. Levitt, *Phys. Chem. Chem. Phys.* 2015, *17*, 5913-5922.
- G. Stevanato, J. T. Hill-Cousins, P. Hakansson, S. S. Roy, L. J. [27] Brown, R. C. D. Brown, G. Pileio, M. H. Levitt, Angew. Chem. Int. Ed. 2015, 54, 3740-3743.
- [28] G. Pileio, M. Carravetta, M. H. Levitt, Proc. Natl. Acad. Sci. U.S.A. 2010, 107, 17135-17139.
- [29] Y. Zhang, P. C. Soon, A. Jerschow, J. W. Canary, Angew. Chem. Int. Ed. 2014, 53, 3396-3399.
- Y. Feng, T. Theis, T.-L. Wu, K. Claytor, W. S. Warren, *Journal of Chemical Physics* 2014, 141. [30]
- K. Claytor, T. Theis, Y. Feng, W. Warren, J. Magn. Reson. 2014, [31] 239.81-86
- [32] Y. Feng, R. M. Davis, W. S. Warren, Nature Physics 2012, 8, 831-837 [33]
 - J. F. P. Colell, et al., Journal of Physical Chemistry C 2017, 121, 6626-6634
 - J.-B. Hoevener, et al., Anal. Chem. 2014, 86, 1767-1774.

 - G. Pileio, Prog. Nucl. Mag. Res. Sp. 2010, 56, 217-231.
 P. J. Rayner, M. J. Burns, A. M. Olaru, P. Norcott, M. Fekete, G. G. R. Green, L. A. R. Highton, R. E. Mewis, S. B. Duckett, Proc. Natl. Acad. Sci. U.S.A. 2017, 114, E3188-E3194.

COMMUNICATION

