

# Mechanisms of coherent re-arrangement for long-lived spin order

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This article is dedicated to Geoffrey Bodenhausen for his 70<sup>th</sup> birthday.

## ABSTRACT

Long-lived spin order-based approaches for magnetic resonance rely on the transition between two magnetic environments of different symmetries, one governed by the strong magnetic field of the spectrometer and the other where this magnetic field is inconsequential. Research on the excitation of magnetic-symmetry transitions in nuclear spins is a scientific endeavour that debuted in Southampton in the years 2000. We advanced in this field carrying the baggage of pre-established directions in NMR spectroscopy. Herein, we discuss the part of discoveries that may have been obscured at the time by our choice to mainly look at them through the experience of such directions. We emphasise methodological developments concerning the mechanisms of translation between coherences that generate long-lived spin order and outline zero-quantum rotations in the starting blocks of long-lived state populations and magnetisation transfers between hyperpolarised heteronuclei and protons, as well as spin dynamics leading to the emergence of long-lived coherences. These pulse sequences can seed further applications: for instance, we discuss the combined use of introduced coherence rotations with classical pulse blocks to obtain 2D correlations between protons and heteronuclei. We hope the pulse sequence building blocks discussed herein open further perspectives for magnetic resonance experiments with long-lived spin order.

## KEYWORDS

NMR methods / Long-lived states / Long-lived coherences

## 34 1. Introduction

35 This paper is an opportunity to present several magnetic resonance concepts free of particular  
36 application-specific introductions. This may allow to such concepts for what they are worth simply  
37 as magnetisation transfer mechanisms and comment on their potential usefulness in further  
38 experiments. We point out that all concepts presented herein were already addressed, albeit  
39 concisely, in references(Ahuja et al., 2010; Sarkar et al., 2007, 2010; P. R. Vasos et al., 2009) or  
40 in the supporting material of these papers.

41 Proposing that a presentation free of application-specific introductions may reveal magnetic  
42 resonance progress to the fullest implies that the drive for traditional discipline-oriented  
43 applications may have obscured part of the concepts in the original papers. A legitimate question  
44 is whether these articles would have been accepted by the journal editors without the applications  
45 in mind, or whether they would have been worth accepting. With hindsight, challenges by editors  
46 on our work in Lausanne and Paris concerned more often the applications than the soundness of  
47 methods developments. For instance, in the search for new singlet-based excitation sequences  
48 (Marina Carravetta et al., 2004) on the route of hyperpolarised magnetisation to long-lived spin  
49 states (LLS), we were never tormented by the question *‘is transport of hyperpolarisation really*  
50 *long-lived?’*(P. R. Vasos et al., 2009) (Pileio, 2020). However, *‘is LLS-based polarisation storage*  
51 *in peptides better than the mere longitudinal relaxation time constant of heteronuclei with which*  
52 *peptides are often isotopically enriched,  $T_1(^{15}N)$ ,  $T_1(^{13}C)$ ?’* was a harrowing question. Equally  
53 present was the doubt: *“are long-lived states, with their complicated excitation and sustaining*  
54 *mechanism, really a better way of measuring slow diffusion, slow exchange constants than*  
55 *heteronuclei(Ferrage et al., 2003) such as  $^{15}N$ ?”* or *‘are long-lived coherences (LLC’s) actually a*  
56 *good route to improved spectral resolution in NMR?’*.

57 When we dedicated the first of a series of papers(Sarkar et al., 2007) to Anatole Abragam along  
58 with a letter expressing our hopes that the discoveries may be useful for diffusion studies, he seized  
59 the essence of our work in his answer (mainly addressed to Geoffrey Bodenhausen): *‘nice to see*  
60 *a way of skillfully sending spins to sleep in their soft bed’, ‘envoyer les spins se reposer dans leur*  
61 *lit douillet’*. The remark, thus rhythmized by alliteration, was as concise as it was exact, since the  
62 singlet state we were searching for is magnetically inactive, i.e., the spins are *‘sleeping’*. This  
63 commentary alone may have replaced the introduction to our original paper.

64

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## 68 **2. Zero-quantum rotation in the starting block of long-lived states**

69 The structure of singlet-triplet population differences, or long-lived states operators,  $Q_{LLS}$ :

70

$$71 \quad Q_{LLS} = |S_0\rangle\langle S_0| - \frac{1}{3}(|T_{-1}\rangle\langle T_{-1}| + |T_0\rangle\langle T_0| + |T_1\rangle\langle T_1|) \quad (1)$$

72

73 was first discussed in formulae adapted to the zero-field magnetic structure for a two-spin system,

74 as first created (Marina Carravetta et al., 2004) in non-equivalent nuclei. While the preference of

75 writing highly-symmetrical long-lived states in spherical tensor operators is natural, we recurred

76 in Lausanne, however, to Cartesian operators (Sørensen et al., 1984) in the Liouvillian space:

77

$$78 \quad Q_{LLS} = -N_{LLS}(I_x S_x + I_y S_y + I_z S_z) \quad (2)$$

79

80 with  $N_{LLS} = 4/3$ .

81 The form of this operator allowed us to understand the structure of coherences prone to evolution:

82

$$83 \quad LLS = -\frac{4}{3}ZQ_x - \frac{2}{3}2I_z S_z \quad (3)$$

84

85 where  $ZQ_x = \frac{1}{2}(2I_x S_x + 2I_y S_y)$  is a zero-quantum coherence.

86

87 Under such a configuration, the system is immune to the scalar-coupling evolution and also to

88 chemical shift evolution, provided the chemical shift difference between the two spins is eclipsed

89 by ample radio-frequency radiation or by cycling the main field (Cavanagh et al., 1995).

90

91 Equation (1) proved to be one the most useful formulae in developing the general theory of long-

92 lived states by pointing out the very nature of their extended lifetime, the population imbalance

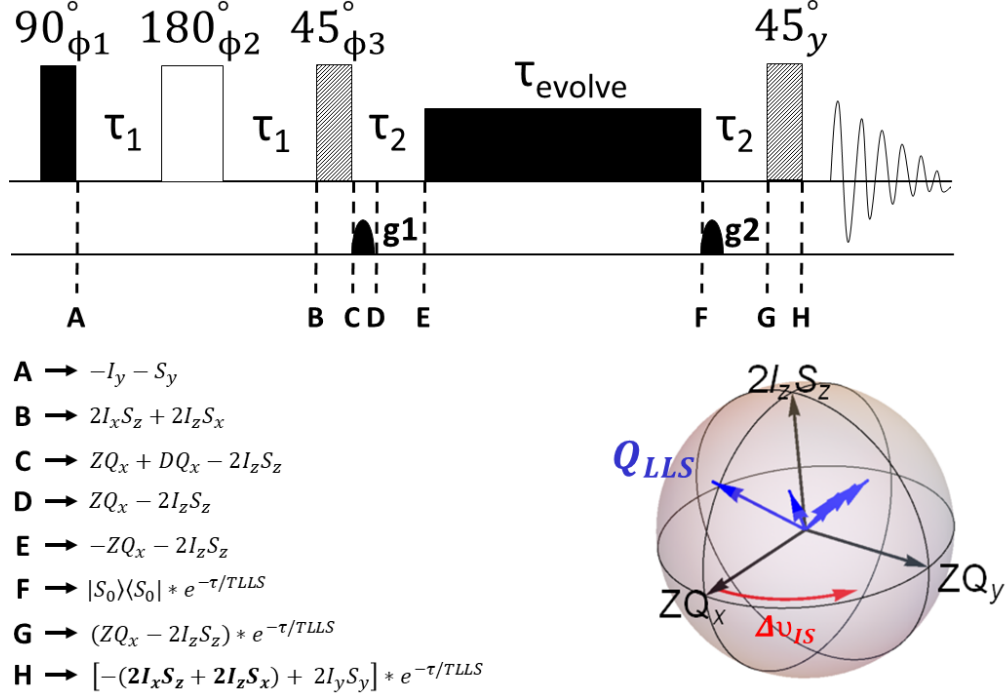
93 between states or manifolds of different symmetries with respect to spin permutations (Stevanato,

94 2015, 2020) which cannot be interconverted by relaxation mechanisms with certain symmetries.

95 The novelty of Equation (3) was that it strongly connected the singlet-states explorations to  
96 research in the I. Solomon - consecrated (I,S) homonuclear and heteronuclear magnetisation  
97 transfer(Solomon, 1955).

98 The first method of excitation for long-lived populations, developed by Levitt and  
99 collaborators(Marina Carravetta & Levitt, 2004), worked for a pair of spins  $I,S$  provided carefully-  
100 chosen delays dependent on the chemical shifts,  $\nu_I$  and  $\nu_S$ , were used, making a sweep through  
101 frequencies necessary to excite different pairs of coupled spins (I,S), (R,K)... in different  
102 experiments, just like 1D magnetic resonance spectroscopy necessitated a sweep of the main field  
103 through resonance conditions for different chemical environments before the introduction of  
104 Fourier transform(Ernst & Anderson, 1966; *Richard R. Ernst – Nobel Lecture.*, n.d.). The  
105 chemical-shift dependency of long-lived states rendered impossible any 2D investigations of  
106 phenomena involving two or more spin pairs or several chemical environments of spin pairs with  
107 encoded LLS, such as exchange or interaction dynamics, in the same experiment.

108  
109 The first concept introduced in the Lausanne paper(Sarkar et al., 2007) was the broadband  
110 excitation of singlet states. The topic may have deserved, in retrospect, a paper on its own. Our  
111 way towards broadband LLS excitation passed through zero-quantum coherences, as explained  
112 below. The first attempts to excite  $Q_{LLS}$  in Lausanne (Figure 1) posed challenges regarding the  
113 evolution and relative orientation of zero-quantum coherences ( $ZQ_x, ZQ_y$ ) and ZZ-magnetization  
114 ( $2I_zS_z$ ).



115

116 Figure 1. Pulse sequence adapted from reference(Sarkar et al., 2007) showing the evolution of the density  
 117 operator at different stages with an emphasis on its three projections ( $ZQ_x, ZQ_y, 2I_z S_z$ )between time points  
 118 C and E (figure generated with SpinDynamica(Bengs & Levitt, 2018)). Here,  $\tau_1 = 1/(4 \cdot J_{IS})$  and  $\tau_2 =$   
 119  $1/(2 \cdot \Delta\nu_{IS})$  where  $J_{IS}$  is the scalar coupling constant (Hz) between spins I and S,  $\Delta\nu_{IS} = \nu_I - \nu_S$  is the  
 120 chemical shift difference (Hz) between the two spins. The phase cycling is  $\phi_1 = (x, -x)$ ,  $\phi_2 = x$ ,  $\phi_3 =$   
 121  $2(y), 2(-y)$  and  $\phi_{rec} = (x, -x, -x, x)$ .

122

123 After the first  $45^\circ$  pulse, at point (C) in Figure 1, the density operator takes the following  
 124 expression:

125

$$126 \rho_{pre-LLS}^C = 2I_x S_x - 2I_z S_z = ZQ_x + DQ_x - 2I_z S_z \quad (4)$$

127

128 At first, we expected to induce the presence of  $Q_{LLS}$  at point (C) in this sequence, due to the  
 129 presence of projections on  $Q_{LLS}$  by both longitudinal two-spin order and  $2I_x S_x$ . However, we  
 130 realized these two contributions exactly annihilate one another, leaving us at a loss on how to  
 131 excite singlets in a broadband manner. We could have anticipated the mutual cancellation by  
 132 expressing the operator at point (C) in the singlet-triplet basis, relevant upon application of a  
 133 ‘sustaining’ radio-frequency (rf) field:

134  $\rho_{pre-LLS}^C = -\frac{1}{2}(|T_{-1}\rangle\langle T_{-1}| + |T_1\rangle\langle T_1| - 2|T_0\rangle\langle T_0| - |T_{-1}\rangle\langle T_1| - |T_1\rangle\langle T_{-1}|)$  (5)

135 indicating there was no singlet order to be found there.

136

137 By applying the pulsed field gradient  $g_I$ , the double-quantum term dissipates, and the density  
138 operator becomes:

139

140  $\rho_{pre-LLS}^D = (I_x S_x + I_y S_y) - 2I_z S_z = ZQ_x - 2I_z S_z$  (6)

141

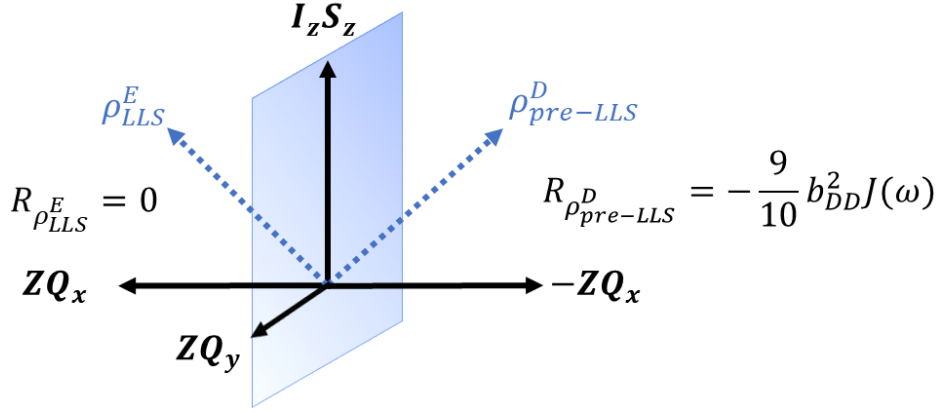
142 which possesses all the component of the long-lived state (Equation 1), but displays an opposite  
143 orientation of zero-quantum and ZZ components with equal projections on  $Q_{LLS}$ . Therefore, these  
144 projections cancel each other. To better understand this apparent conundrum, we can write the  
145 operator in the basis of singlet-triplet operators:

146

147  $\rho_{pre-LLS}^D = \frac{1}{2}(|T_0\rangle\langle T_0| - |S_0\rangle\langle S_0|) - \frac{1}{2}(|T_{-1}\rangle\langle T_{-1}| + |T_1\rangle\langle T_1| - |T_0\rangle\langle T_0| - |S_0\rangle\langle S_0|) =$   
148  $\frac{1}{2}(2|T_0\rangle\langle T_0| - |T_{-1}\rangle\langle T_{-1}| - |T_1\rangle\langle T_1|)$  (7)

149

150 The next step was figuring out how to interchange the singlet and central triplet populations in  
151 order to get the expression for  $LLS$ , a task which is not immediately obvious in this form. To do  
152 that, a reversion in the cartesian product basis proved fruitful:  $|T_0\rangle\langle T_0| - |S_0\rangle\langle S_0| = I_x S_x +$   
153  $I_y S_y = ZQ_x$ . After several weeks of calculations, a group seminar was dedicated to the otherwise  
154 well-known evolution of  $ZQ_x$  under a scalar coupling interaction (Cavanagh et al., 1995). The new  
155 aspect was that the rotation axis was this time also apparent in the density operator expression, so  
156 effectively one of the constituents of spin order was rotating around the other, thus changing the  
157 relative sign of the constituent product operators  $ZQ_x$  and  $2I_z S_z$  and yielding the sought-after  $LLS$   
158 (Figure 2). The atmosphere in the magnetic resonance laboratory in Lausanne should be credited  
159 for a substantial contribution to the birth of these concepts. However, as a side note, the physical  
160 exercises of magnetisation succeeded to captivate more attention on paperback than in the coffee-  
161 table setting around a group-meeting whiteboard.



162  
 163 Figure 2. Position of coherences prior to and after zero-quantum rotation by 180 degrees around  
 164 longitudinal two-spin order. The relaxation rate constants corresponding to the density operator given by  
 165 the two linear combinations of  $ZQ_x$  and  $I_z S_z$ , taking into account only the dipolar relaxation mechanism for  
 166 the pair of two spins, with a dipolar coupling constant  $b_{DD}$ ;  $J(\omega)$  is the spectral density function.

167  
 168 The evolution of the  $ZQ_x$  during free precession is:

169  
 170 
$$ZQ_x \xrightarrow{(2\pi\nu_I I_z + 2\pi\nu_S S_z)\tau} ZQ_x \cos(2\pi\Delta\nu_{IS}\tau) + ZQ_y \sin(2\pi\Delta\nu_{IS}\tau) \quad (8)$$

171  
 172 where  $\nu_I$  is the Larmor frequency of spin  $I$ ,  $\nu_S$  is the Larmor frequency of spin  $S$ ,  $\Delta\nu_{IS} = \nu_I - \nu_S$  is  
 173 the chemical shift difference (Hz) between the two spins and  $J_{IS}$  is the scalar coupling constant  
 174 (Hz) between spins  $I$  and  $S$ . Thus, after an evolution period  $\tau_2 = 1/(2\Delta\nu)$ , the  $ZQ_x$  will change  
 175 sign such that  $\frac{1}{2}(|T_0\rangle\langle T_0| - |S_0\rangle\langle S_0|) \xrightarrow{(2\pi\nu_I I_z + 2\pi\nu_S S_z)\tau} \frac{1}{2}(|S_0\rangle\langle S_0| - |T_0\rangle\langle T_0|)$ . During the evolution  
 176 represented in Figure 2, the right-hand side in the first quadrant of the coherence clock representing  
 177  $\rho_{pre-LLS}^D$  is transformed into its symmetric image with respect to the  $(2I_z S_z, ZQ_y)$  plane, in the  
 178 fourth quadrant. The initial  $\rho_{pre-LLS}^D$  and its plane-symmetric image  $\rho_{LLS}^E$  have relaxation rate  
 179 constants governed by different symmetry rules. The density operator evolved into:

180  
 181 
$$\rho_{LLS}^E = |S_0\rangle\langle S_0| - \frac{1}{2}(|T_{-1}\rangle\langle T_{-1}| + |T_1\rangle\langle T_1|) \quad (9)$$

182  
 183 A note can be made on the time dependence of the relaxation rate constant for the case of free  
 184 precession of the LLS. Given the evolution of zero-quantum coherences under the chemical shift

185 difference, the density operator, starting from  $ZQ_x + I_z S_z$ , will have an oscillatory evolution  
 186 between the two  $ZQ_x$  and  $ZQ_y$  as:

187

$$188 \quad ZQ_x + I_z S_z \xrightarrow{(2\pi\nu_I I_z + 2\pi\nu_S S_z)\tau} ZQ_x \cos(2\pi\Delta\nu_{IS}\tau) + ZQ_y \sin(2\pi\Delta\nu_{IS}\tau) + I_z S_z \quad (10)$$

189

190 Considering a relaxation superoperator only for the dipolar interaction between two coupled spins:

191

$$192 \quad \hat{F} = -\frac{6}{5} b_{DD}^2 \sum_{m=-2}^2 (-1)^m J(m * \omega) [T_{2,m} [T_{2,-m-}]] \quad (11)$$

193

194 where  $b_{DD}$  is the dipolar coupling constant,  $J(\omega)$  is the spectral density and  $T_{2,m}$  are the spherical  
 195 tensor spin operators of rank 2, the computed relaxation rate constant for the density operator  
 196  $ZQ_x \cos(2\pi\Delta\nu_{IS}\tau) + ZQ_y \sin(2\pi\Delta\nu_{IS}\tau) + I_z S_z$

197 is

$$198 \quad R(\tau) = -\frac{6}{5} b_{DD}^2 ((1 + \cos(2\pi\Delta\nu_{IS}\tau)) \cdot J(0) + 3 \cdot J(\omega)) \cdot (\sin(\frac{2\pi\Delta\nu_{IS}\tau}{2}))^2 \quad (12)$$

199

200 Thus, only for  $\tau = \frac{2n}{2\pi\Delta\nu_{IS}}$  in the starting blocks the relaxation rate constant of the obtained state is  
 201 optimally low.

202

203 Juggling with operators in order to drive the spin system in its ‘*soft bed*’ we realized we should  
 204 always look at Nature from various perspectives. We learned that if only one of the longitudinal  
 205 two-spin order and zero-quantum components could be selected at time point (C) (Figure 1),  $Q_{LLS}$   
 206 would have been present already. For instance, ZZ-magnetization alone projects on the long-lived  
 207 state, given that  $2I_z S_z = \frac{1}{2} (|T_{-1}\rangle\langle T_{-1}| + |T_1\rangle\langle T_1| - |T_0\rangle\langle T_0| - |S_0\rangle\langle S_0|)$ . Thus, during the  
 208 sustaining period, only the singlet population will survive for a period much longer than  
 209 longitudinal magnetization. In order to do so, we employed a Thrippleton-Keeler(Thrippleton &  
 210 Keeler, 2003) filter to wipe out the troubling zero and double quantum coherences and obtained a  
 211 singlet state with an amplitude two-times lower than using both zero-quantum and ZZ-  
 212 magnetization. Other groups employed the so-called “pseudo singlet order”(Pileio, 2017) which is  
 213 just  $ZQ_x = \frac{1}{2} (|T_0\rangle\langle T_0| - |S_0\rangle\langle S_0|)$  as the source for long-lived state obtaining similar results.



214  
215 Though broadband excitation of singlet states would have deserved publication as a discovery in  
216 its own right, we were cautious to avoid publication of our research in slices of ‘salami  
217 science’(Sweedler, 2019). However, this progress proved relevant for the advancement of long-  
218 lived state order(Bengs et al., 2020; Pileio, 2017, 2020; Teleanu et al., 2021) and was more  
219 challenging to obtain than the 2D spectroscopy application for the study of singlet-state-based  
220 exchange we describe in the same paper(Sarkar et al., 2007) (SS-EXSY). In the tradition of finding  
221 low-key names for sequences such as ‘INEPT’(Morris & Freeman, 1979) or  
222 ‘INADEQUATE’(Bax et al., 1980), we could have named the zero-quantum rotation block of the  
223 pulse sequence in Figure 1 a ‘(ZZ-)ZEROTATION’.

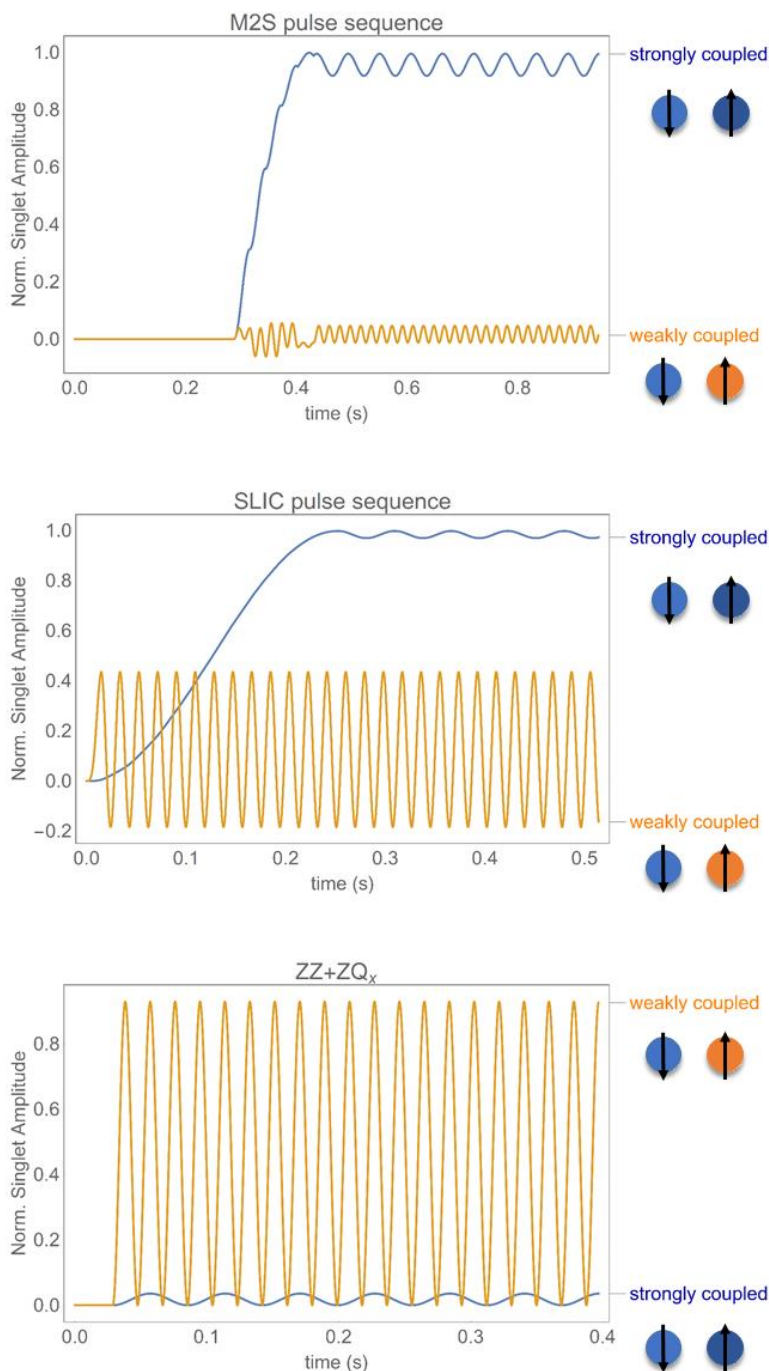
224  
225 As singlet-based applications ambition to store magnetisation for ever longer time periods, the  
226 most adapted systems to this purpose, quasi-equivalent spin pairs with  $J$ -couplings far  
227 overweighing the differences between the chemical shifts of the components, became increasingly  
228 studied. Spin dynamics that shift the magnetisation of the two spins differentially to create the  
229 singlet state are particularly challenging in such systems.

230 Pulse sequences of particular interest include the Magnetization-to-Singlet(Tayler & Levitt,  
231 2011a) and Spin-Lock Induced Crossover(DeVience et al., 2013). These methods are suited for  
232 strongly-coupled spins where the scalar coupling is larger than the chemical shift difference, while  
233 the pulse sequence described in Figure 2, which we identify herein as “ZZ+ZQ<sub>x</sub>” (Sarkar et al.,  
234 2007), performs better in terms of long-lived states excitation in the weakly-coupled regime.  
235 Several attempts to efficiently excite singlet order on broader domains of coupling regimes have  
236 been recently devised(Bengs et al., 2020; Mamone et al., 2020). Figure 3 depicts numerical  
237 simulations performed with SpinDynamica(Bengs & Levitt, 2018) for singlet population  
238 excitation using the aforementioned pulse sequences for both weakly and strongly coupled regime  
239 outlining the difference in excitation efficiency.

240  
241 **3. Heteronuclei or proton long-lived states for conserving hyperpolarisation**  
242 In order to maximise the magnetisation lifetime, heteronuclear longitudinal spin-order (mainly on  
243 carbon-13) can be excited and used during evolution periods in both room-temperature(Bermel,  
244 Bertini, Duma, et al., 2005; Richter et al., 2010) and hyperpolarised NMR. We strived to also

245 preserve hyperpolarization on a pair of hydrogens entwined in a long-lived state(P. R. Vasos et al.,  
246 2009). Since the invention of dissolution-Dynamic Nuclear Polarisation (dissolution-  
247 DNP)(Ardenkjær-Larsen et al., 2003; Balzan et al., 2016; Comment et al., 2008) and its  
248 development in Lausanne by the team of A. Comment, S. Jannin, and J. van der Klink in the  
249 Functional Imaging Laboratory at EPFL(Ardenkjær-Larsen et al., 2003; Balzan et al., 2016;  
250 Comment et al., 2008), the topic was associated with our research due to its conjunction with long-  
251 lived spin order(Ardenkjær-Larsen et al., 2003; Balzan et al., 2016; Comment et al., 2008).

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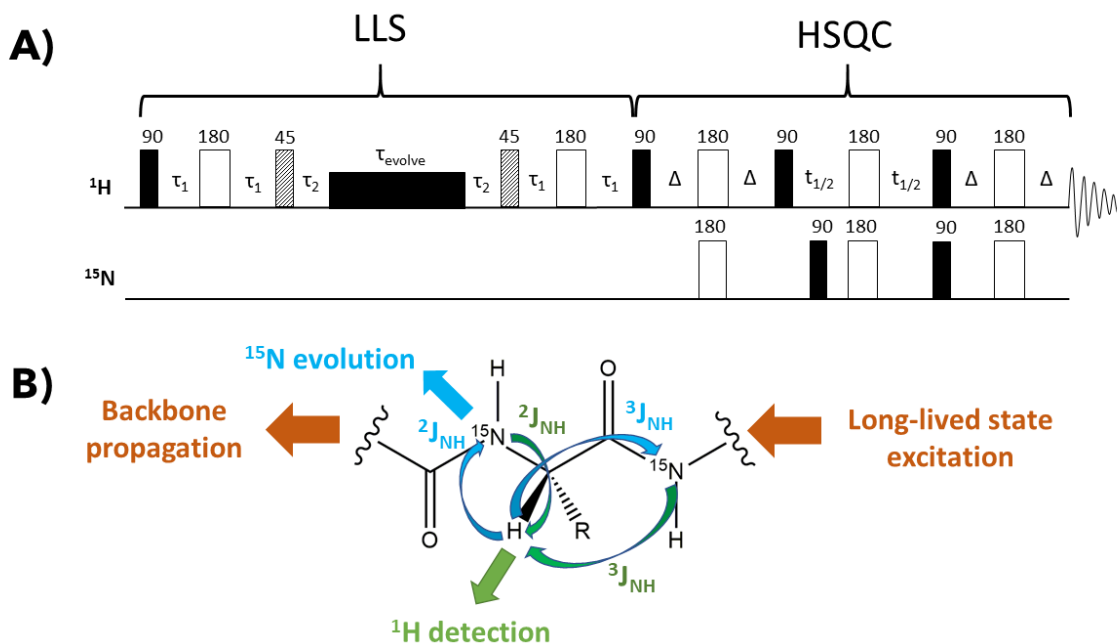
254 Figure 3. Numerical simulations using SpinDynamica outlining the efficiency of singlet polarisation for  
 255 M2S, SLIC and ZZ+ZQ<sub>x</sub> pulse sequences in a two-spin system (by projecting the density operator during  
 256 the pulse sequence onto the singlet population). The weakly-coupled system is described by  $\{\Delta\nu_{IS} =$   
 257  $50 \text{ Hz}; J_{IS} = 17.4 \text{ Hz}\}$ , while the strongly coupled system features  $\{\Delta\nu_{IS} = 2.8 \text{ Hz}; J_{IS} = 17.4 \text{ Hz}\}$ . For  
 258 each scenario, only the coherent evolution was considered in simulations (no relaxation dampening of the  
 259 amplitude of source operators or LLS is taken into account). The excitation period is followed by a free-  
 260 precession evolution after the maximum amplitude for the singlet population was reached.

261 The preservation of hyperpolarised magnetisation obtained by dissolution-DNP in long-lived  
262 states raised fewer challenges than the comparison of the LLS with heteronuclear lifetimes in terms  
263 of performance as polarisation batteries. The hyperpolarised magnetisation in samples stemming  
264 from a polarizer such as the one developed in Lausanne and the similar one installed as the first  
265 dissolution-DNP system in France(Ardenkjær-Larsen et al., 2003; Balzan et al., 2016; Comment  
266 et al., 2008) followed intently the rf pulses in our high-resolution magnets. Inhomogeneities due  
267 to fast dissolution and injection could be tamed to run pulse sequences for LLS excitation and  
268 decoding and observe the signal. To us, the main issue remained that the benefits of a procedure  
269 consisting of  $Q_{LLS}$  excitation on protons compared to simply preserving hyperpolarised  
270 magnetisation in heteronuclei had to be carefully considered. We scrutinised this issue in terms of  
271 magnetisation lifetimes in the given conditions (room temperature, molecular size) as well as in  
272 the case of extreme molecular sizes or crowding of the environment and came to the conclusion  
273 that proton-based LLS were valuable for storing magnetisation even when the molecules contained  
274 isotope-enriched heteronuclei like  $^{15}\text{N}$  or  $^{13}\text{C}$ . In this analysis, we were inspired by similar  
275 comparisons between proton and heteronuclear magnetisation carried out for relaxation rates of  
276 heteronuclei and protons in large or paramagnetic proteins(Bermel, Bertini, Felli, et al., 2005).  
277 Journal editors understood the interest of the research topic and the papers we sent for publication  
278 encountered no uphill sisyphian battles (Molinié & Bodenhausen, 2013; *The Myth of Sisyphus* -  
279 *Wikiwand*, n.d.). The only significant delays in publication we incurred were for a paper submitted  
280 directly to a specialized journal that discussed results obtained using our freshly-installed DNP  
281 system at the time (Balzan et al., 2017).

282 Potential applications of symmetry-adapted states as magnetisation reservoirs for various 2D  
283 experiments led us to investigate the polarization transfer from long-lived state of protons towards  
284 heteronuclei across proteins' backbones (Figure 4)(L. Fernandes et al., 2013; Laetitia Fernandes  
285 et al., 2013). The particular relaxation rates of LLS reflect conformational exchange and act as  
286 probes for unravelling protein's inner dynamics, while 2D correlations (e.g.,  $^1\text{H}$ - $^{15}\text{N}$ ) help  
287 disentangle complex spectra, which is acutely needed in the analysis of unfolded or intrinsically  
288 disordered proteins. The field of 2D heteronuclear experiments using long-lived spin order (Figure  
289 4) is likely to develop further. As the field advances, long-lived states-based explorations of inter-  
290 molecular(Bornet et al., 2011; Stavarache et al., 2017) and intra-molecular(Ahuja et al., 2007;

291 Tayler et al., 2010) interactions become important, and so becomes spectral resolution for the study  
 292 of systems of increasing complexity.

293  
 294



295  
 296 Figure 4. A) Pulse sequence designed to generate a two-dimensional correlation spectrum between  
 297  $^1\text{H}$  and  $^{15}\text{N}$  via scalar coupling of the type  $^2J_{\text{NH}}$  and  $^3J_{\text{NH}}$  starting from a proton long-lived state. B)  
 298 Schematic representation of polarization transfer along protein's backbone which generates a two-  
 299 dimensional correlation spectrum via the  $^2J_{\text{NH}}$  and  $^3J_{\text{NH}}$  coupling constants. Protons are excited via  
 300 the first part of the pulse sequence from (A) into a long-lived state spin order. Figure adapted from  
 301 reference(Laetitia Fernandes et al., 2013).

302

#### 303 4. Long-lived coherences, eppur' si muove

304 Aware of the potential of long magnetisation lifetimes for line-narrowing in NMR, we strived to  
 305 obtain some type of magnetisation akin to LLS to rotate. Long lifetimes of magnetisation had been  
 306 traded for spectral resolution, e.g., for  $^1\text{H}$ - $^{15}\text{N}$  pairs in cross-correlated relaxation experiments and  
 307 for  $^{15}\text{N}$  heteronuclei for narrowing spectroscopic lines(Goldman, 1984; Pervushin et al., 1997; Paul  
 308 R. Vasos et al., 2006).

309 Our search for ‘moving’ long-lived configurations first involved complicated coherences in  
 310 alanine, serine, and other molecules with up to five coupled spins(Ahuja et al., 2009). When we  
 311 finally resorted to our favorite paired Gly aliphatic protons of AlaGly, the  $I_x$ - $S_x$  configuration was  
 312 deduced from the diagonalized Liouvillian(M. Carravetta & Levitt, 2005). The next hurdle  
 313 appeared in fitting the exotic long-lived states with products of oscillating and decaying functions  
 314 and translating them to signals in 2D spectra via Fourier transform-adapted spectroscopy (prior  
 315 experience non-conventional heteronuclear 2D experiments(Bertini et al., 2004; P.R. Vasos et al.,  
 316 2005) helped at this point). The simple scheme involving a  $180^\circ$  pulse for creating a  $Q_{LLC}$   
 317 observable in the indirect dimension of a 2D experiment, where:

318

$$319 \quad Q_{LLC} = (I_x - S_x) \cos(2\pi J_{IS}\tau) + (2I_y S_z - 2I_z S_y) \sin(2\pi J_{IS}\tau) \quad (13)$$

320

321 was further refined by different approaches (Figure S1), of which the most advanced itches  
 322 coherences with spins pointing in opposite ways in molecules with almost-equivalent  
 323 nuclei(Sheberstov et al., 2019).

324

325 When we sought for long-lived coherences in high fields, we were not aware yet that the  
 326 contemporary work on extremely low-frequency oscillations(Pileio et al., 2009) in low magnetic  
 327 fields developed at Southampton involved, practically, the same operators.

328

329 Just as in the case of LLS for hyperpolarisation safekeeping was compared to heteronuclear  
 330 storage, once LLC’s were developed comparisons with zero-quantum coherences came to mind.  
 331 For the case of two  $J$ -coupled, non-equivalent spins,  $I$  and  $S$ , the evolution of  $\rho_1(0) = I_x - S_x$  and  
 332  $\rho_{ZQx}(0) = I_x S_x + I_y S_y$  during free-precession (without any radio-frequency ‘sustaining’ applied)  
 333 is given by:

334

$$335 \quad \rho_{ZQx}(\tau) = (I_x S_x + I_y S_y) \cos(2\pi \Delta\nu_{IS}\tau) + (I_y S_x - I_x S_y) \sin(2\pi \Delta\nu_{IS}\tau) = ZQx \cos(2\pi \Delta\nu_{IS}\tau) +$$

$$336 \quad ZQy \sin(2\pi \Delta\nu_{IS}\tau) \quad (15)$$

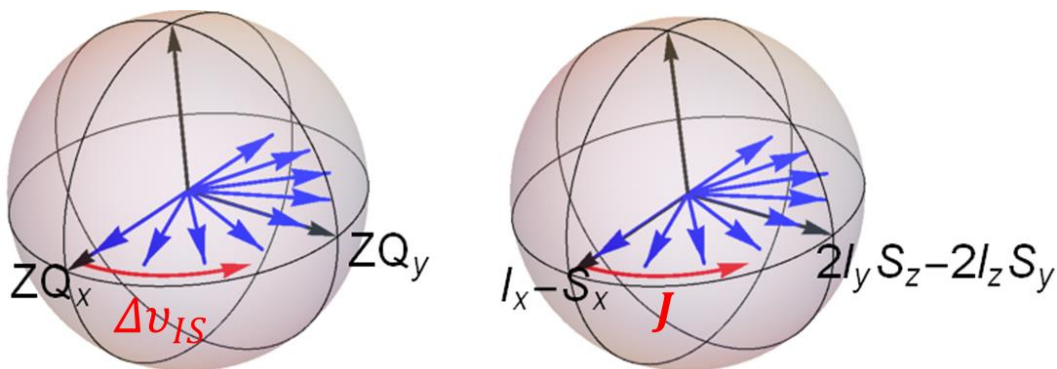
337

338 while in presence of rf fields with the carrier placed at the middle of their offsets ( $v_1 = -\Delta v_{IS}/2$   
 339 and  $v_2 = \Delta v_{IS}/2$ ) and an amplitude  $v_1 \gg \Delta v_{IS}$ , the evolutions for the LLC and  $ZQ_x$  are:

340  
 341 
$$\rho_{LLC}(\tau) = (I_x - S_x) \cos(2\pi J_{IS}\tau) + (I_y S_z - I_z S_y) \sin(2\pi J_{IS}\tau) \quad (16)$$

342  
 343 
$$\rho_{ZQx}(\tau) = I_x S_x + I_y S_y * \cos(2\pi v_1 \tau)^2 + I_z S_z * \sin(2\pi v_1 \tau)^2 + (I_y S_z +$$
  
 344 
$$+ I_z S_y) \sin(2\pi v_1 \tau) \cos(2\pi v_1 \tau) \quad (17)$$

345



346  
 347 Figure 5. Evolution of zero-quantum coherences during free precession with an oscillating frequency of  
 348  $\Delta v_{IS}$  and evolution of long-lived coherences during sustaining period with an oscillating frequency equal  
 349 to the scalar coupling constant  $J_{IS}$ .

350

351 In the absence of sustaining rf fields, differences of single-quantum transverse coherences, the  
 352 source of LLC's, evolve under the chemical shift difference and the scalar coupling, while  $ZQ$ 's  
 353 are immune to the latter (Cavanagh et al., 1995). In the second scenario, LLC's evolve only under  
 354 the effect of  $J$ -coupling (Sarkar et al., 2010), oscillating between in-phase ( $I_x - S_x$ ) and anti-phase  
 355 ( $2I_y S_z - 2I_z S_y$ ), with the coherence order equal to 1. Thus, the scalar coupling evolution sets  
 356 LLC's aside from  $ZQ$ 's (Figure 5).

357 Broadband excitation of LLC's in molecules with broadly different  $J$ -couplings and chemical  
 358 shifts is still a challenge, despite the progress. We explored part of the territory by exciting with a  
 359 series of selective 180 degrees pulses (Sarkar et al., 2011) and by sustaining with various pulse  
 360 trains (Sadet et al., 2014), but we can safely say that LLC's benefit from their simple and  
 361 parametric-free excitation scheme which consists of a selective  $\pi$  pulse and non-selective  $\pi/2$  hard  
 362 pulse followed by spin-lock.

363

364

## 365 **Conclusions**

366 We present from today's perspective several challenging aspects in the introduction of coherent  
367 dynamics designed to render spin order resilient to Chronos' decrees. The focus is placed on zero-  
368 quantum inversion with respect to longitudinal two-spin order in homonuclear spin systems.  
369 Numerical simulations outlining the efficiency of different pulse sequences to create long-lived  
370 states in different coupling regimes are discussed. The presented methods are delineated in a  
371 manner designed to render them useful as building blocks in further applications.

372

## 373 **Author contribution**

374 FT and PV conceived the paper, FT performed the numerical simulations, FT and PV wrote the  
375 paper.

376

## 377 **Competing interests**

378 The authors declare that they have no conflict of interest.

379

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387

## 388 **Code availability**

389 The Mathematica notebook used for the simulations is provided as supplementary information.

390

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