

Optimising NMR Spectroscopy through Method and Software Development



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Abstract

Solution-state nuclear magnetic resonance (NMR) spectroscopy is one of the most important analytical techniques in modern organic chemistry. A series of ‘core’ NMR techniques is widely used for the routine characterisation of organic molecules, ranging from 1D ^1H and ^{13}C experiments to more complex 2D experiments such as COSY, HSQC, HMBC, and more. Nonetheless, there remains much room for developments in NMR, especially considering the increased *complexity* and *quantity* of molecules which are being studied. This thesis describes multiple approaches towards improving the quality, as well as the speed, of NMR data acquisition.

In chapter 1, I first set out the quantum mechanical formalisms required for the analysis of NMR pulse sequences. The work which follows may be divided into three sections:

- *Improved techniques for pure shift NMR* (chapter 2). In a pure shift experiment, the effects of all homonuclear scalar couplings are suppressed, leading to highly-resolved spectra where each multiplet is collapsed to a singlet. A variety of theoretical and experimental approaches are used to search for pure shift methods which have both high sensitivity and low artefact intensity.
- *On-the-fly optimisation of NMR experiments* (chapter 3). Here, I couple NMR acquisition with derivative-free optimisation algorithms which seek to minimise a cost function measured on spectral data. The resulting software, called POISE, allows experiment parameters to be tailored specifically for each sample and spectrometer in a fully automated fashion. The use of POISE to improve spectral sensitivity and purity is demonstrated on a wide-ranging series of NMR experiments.
- *Accelerated 2D NMR data collection* (chapter 4). NOAH supersequences allow multiple 2D experiments ('modules') to be acquired in the time of one, through the careful manipulation of different magnetisation pools. This chapter contains work on pulse sequence development, including new NOAH-compatible modules, as well as improvements to various pre-existing modules. I also describe the GENESIS web site, which creates Bruker pulse programmes for arbitrary NOAH supersequences.

Declaration of authorship

This thesis consists entirely of my own work, except for:

- Figures 2.25a to 2.25c, which illustrate data acquired by Corentin Jacquemmoz and Jean-Nicolas Dumez at the University of Nantes; and
- § 3.5 on ESR-POISE, where all experimental data were acquired by, and a portion of the code written by, Jean-Baptiste Verstraete, David Goodwin, and William Myers, at the University of Oxford. In particular, the data used for Figure 3.32 were acquired by Jean-Baptiste Verstraete.

These contributions are also explicitly indicated within the text.

This work has not been submitted, either wholly or substantially, for any qualification at this or any other institution.

Acknowledgements

‘I cannot make speeches, Emma:’—he soon resumed; and in a tone of such sincere, decided, intelligible tenderness as was tolerably convincing.—‘If I loved you less, I might be able to talk about it more.’

— JANE AUSTEN, *Emma*

First of all, I would like to thank my supervisors, Tim Claridge and Mohammadali Foroozandeh. Throughout my studies, their guidance has been understated yet unerring; I have become a better scientist and person for it, and hope that this thesis bears witness to the quality of their counsel. Beyond that, I greatly appreciate being afforded the freedom to explore other things which did not directly feed back into my DPhil output. This includes the short period where I worked in the NMR team, as well as all the time I have spent on programming, which explains—but does not justify—many a week where no work was produced. Yet, because of this, I have gotten so much more out of these four years than just a DPhil, and can look towards my future career with excitement and optimism.

I am grateful to the Clarendon Fund, as well as the SBM CDT, for funding this work. They have made it possible for me to stay in Oxford—a place now very dear to me.

I am also deeply indebted to my industrial mentors, Peter Howe, Philip Sidebottom, Iain Swan, and Harry Mackenzie. They have always taken a keen interest in all aspects of my work, and I shall remember with much fondness the very kind invitation to the Jealott’s Hill Syngenta site. Here, I wish to also thank my professional collaborators in the world of magnetic resonance, Ēriks Kupče and Jean-Nicolas Dumez: it has been nothing short of a privilege to work with and learn from them.

Much credit is due to the people with whom I have been fortunate to work with in Oxford. In particular, I wish to thank the NMR staff: Caitlin, James, Maria, Nader, Nick, and Tina (and Tim, again); and various members of the NMR group past and present, including Ali, David, Fay, JB, Jos, Noelle, Sam, and Simon. By, at least occasionally, creating in me a desire to go into the office, they have tempered my natural indolence, which is no mean feat. It is my sincere hope that we may continue to stay in touch, now as friends rather than colleagues.

On a more personal level, learning the flute has been my greatest undertaking in Oxford outside of my work: one which has been sometimes stressful, but always joyful. For this, I must thank my teacher Jean, who has been kind to me well beyond her duty as an instructor, as well as her husband Robin.

I am also very grateful for the friends I have had during my time in Oxford, including (but not limited to) Alvin, Belinda, Jing Long, Liv, Steve, Jieyan, Sheryl, and Wearn Xin; and the rest of my SBM cohort, especially Henry and Kate. Many of you will already have moved on to other adventures, but I am so glad to have crossed paths with you at some point. Distinguished most in both constancy and intimacy, however, is Marie. I have been extremely fortunate to have enjoyed your company over the last few years: there are not many people who would understand me as you do, and on such a wide range of topics—thank you so very much.

Finally, I come to the people who have immeasurably shaped my life for the better. My parents have sacrificed much of their freedom to guarantee mine: the work I have done here is entirely due to theirs. I cannot even begin to repay their unconditional love, but I pray that through my life I may do them proud and thereby vindicate their choices. Last and dearest is my partner Bernie, who has been the source of every felicity during my time in Oxford. None of the achievements in this work can equal the profound joy which I derive from her constant presence and unfailing support.

Preface

Miss Dashwood at this point, turning her eyes on him with the most angry contempt, stopped him, by saying, ‘It is hardly worth while, Mr. Willoughby, for you to relate, or for me to listen any longer. Such a beginning as this cannot be followed by any thing.—Do not let me be pained by hearing any thing more on the subject.’

— JANE AUSTEN, *Sense and Sensibility*

Pulse sequence notation

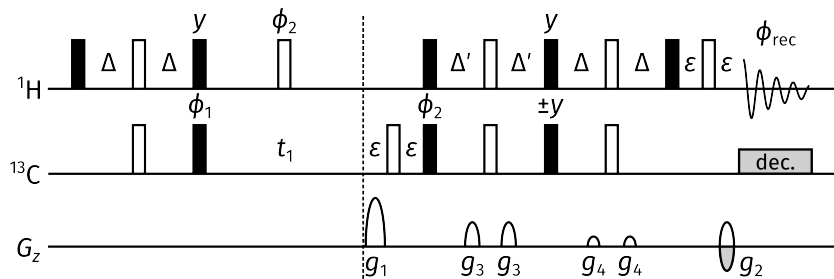


Figure 0.1: An example of a pulse sequence (the sensitivity-enhanced HSQC), used here to illustrate the notation used in this thesis.

NMR pulse sequences in this thesis are depicted in a reasonably conventional manner: fig. 0.1 shows an example of this. To avoid any ambiguity, I describe this notation here. Filled black rectangular bars indicate 90° pulses; empty bars indicate 180° pulses. Pulses with other flip angles are depicted using filled grey bars, typically with the Greek letter β above it representing the flip angle. Delays are variously represented by the letters Δ and τ ; exact values of these are given in the respective captions. Pulses without explicit phases are assumed to be applied along the (rotating-frame) $+x$ -axis; pulses with phases labelled as ϕ_i are typically phase-cycled, details of which are specified in the respective captions. Grey boxes labelled ‘dec.’ represent periods of heteronuclear decoupling.

In this thesis, I use the word *gradient* to refer to pulsed field gradients; this is more concise and reflects common usage. Where relevant, z -gradient amplitudes are given as percentages of

the maximum gradient amplitude, which is probe-dependent (see table 0.1). This maximum amplitude is unlikely to substantially affect the performance of any of the pulse sequences; consequently, in the text I quote gradient amplitudes only as percentages. Nevertheless, if necessary, absolute gradient amplitudes may still be determined for each individual dataset by referring to table 0.1. Gradients which are inverted for echo–antiecho selection are depicted as a pair of positive and negative gradients with different shading (e.g. g_2 above). ε is always used to denote the time required for a gradient, plus the subsequent recovery delay (200 μs throughout this work).

In practice, the implementation of a pulse sequence may differ in tiny ways: for example, delays may be modified to accommodate finite pulse widths and other technicalities. Furthermore, shaped pulses may be used in place of hard pulses in order to optimise the pulse sequence, e.g. by allowing more efficient refocusing: as a rule, adiabatic pulses are used for all ^{13}C 180° rotations.

For some experiments, particularly in the NOAH chapter (chapter 4), a product operator analysis is added above the pulse sequence. Typically, these show how a ^1H –X spin pair and an isolated ^1H spin evolve under during pulse sequence: these directly correspond to the so-called $^1\text{H}^X$ and $^1\text{H}^{IX}$ magnetisation pools (i.e. protons directly coupled and not directly coupled to ^{13}C , see § 4.1.2 for further explanation). One-letter terms m ($m \in \{x, y, z\}$) are shorthand for single-spin terms on proton, i.e. I_m . Two-letter terms mn are shorthand for two-spin terms on both the proton and heteronucleus, i.e. $2I_mS_n$. The effects of gradients are ignored in the product operator analysis: this can be justified because the coherence transfer pathways (CTPs) of interest are ultimately rephased, meaning that the gradients have no net effect (this is illustrated in greater detail in § 1.4.5). For simplicity, any modulation acquired during t_1 is also omitted in these analyses; it will be discussed in the text where relevant.

Software

All NMR data were processed using TopSpin 3 or 4. Quantum mechanical NMR simulations were done in Matlab R2021a or R2021b. This thesis was written using the L^AT_EX typesetting system: specifically, I used the LuaL^AT_EX engine.

Pulse sequences and other diagrams are drawn using the vector graphics programme Inkscape. Plots are generated using Python 3, using a number of packages, namely: `numpy`, `scipy`, `matplotlib`, `seaborn`, and `penguins`, which was written by me.

The L^AT_EX sources for this thesis, as well as all figures, can be accessed at GitHub: <https://github.com/yongrenjie/thesis>.

Samples used

The caption of every figure containing experimental data includes a ‘data code’ at the end, which specifies the spectrometer and sample used for the data, as well as the date it was acquired (in YYMMDD format). The spectrometers and samples used in this thesis are enumerated in tables 0.1 and 0.2 and fig. 0.2. Therefore, for example, the code 7Z-211225 would represent data acquired on the 700 MHz spectrometer, using the zolmitriptan sample, on Christmas Day 2021.

Peak assignments and other physical data for some of the samples are given in Appendix A.

Code	Internal name	Details
7	AV700	700 MHz ^1H resonance frequency 5 mm TCI $^1\text{H}/^{13}\text{C}/^{15}\text{N}$ inverse cryoprobe 53 G cm^{-1} maximum z -gradient amplitude AVANCE III console, TopSpin 3.6.2
6	AV600	600 MHz ^1H resonance frequency 5 mm Prodigy N ₂ broadband cryoprobe ($^1\text{H}/^{19}\text{F}$ outer coil) 66 G cm^{-1} maximum z -gradient amplitude AVANCE III HD console, TopSpin 3.6.2
5	AVX500	500 MHz ^1H resonance frequency 5 mm TBO $^1\text{H}/^{19}\text{F}/\text{X}$ broadband room-temperature probe 50 G cm^{-1} maximum z -gradient amplitude AVANCE III HD console, TopSpin 3.6.2
4	AVB400	400 MHz ^1H resonance frequency 5 mm broadband room-temperature SmartProbe 50 G cm^{-1} maximum z -gradient amplitude AVANCE NEO console, TopSpin 4.0.8

Table 0.1: Spectrometers used in this thesis. A more complete description may be accessed via the links in the ‘internal name’ column.

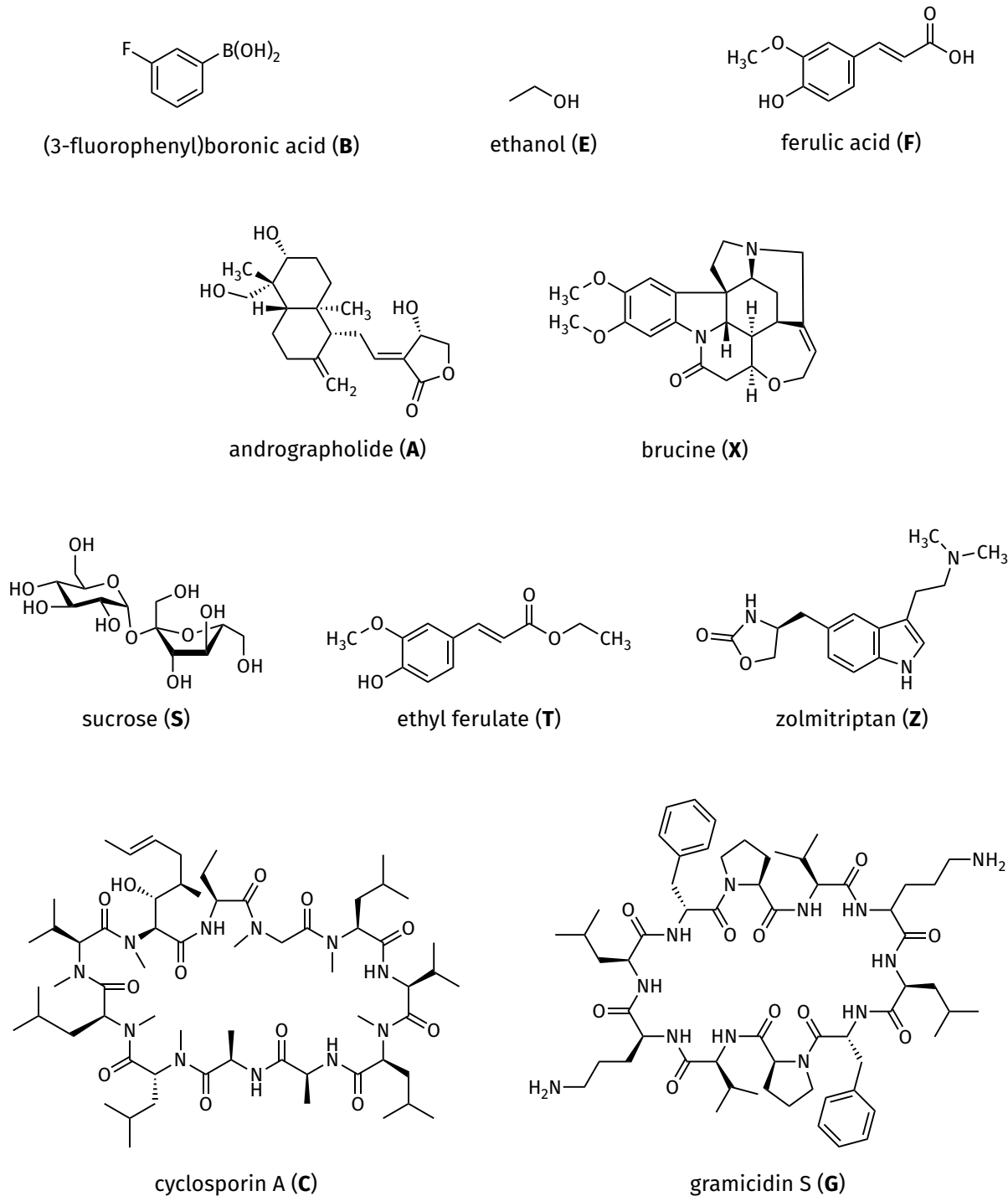


Figure 0.2: Chemical structures of samples used in this thesis. See table 0.2 for more information.

Code	Compound	Solvent	Concentration
A	Andrographolide	DMSO- <i>d</i> ₆	40 mM
B	(3-Fluorophenyl)boronic acid	DMSO- <i>d</i> ₆	120 mM
C	Cyclosporin A	C ₆ D ₆	50 mM
E	Ethanol	D ₂ O	1 M
F	Ferulic acid	DMSO- <i>d</i> ₆	50 mM
G	Gramicidin S	DMSO- <i>d</i> ₆	40 mM
P	Rodent urine	D ₂ O	-
S	Sucrose	90% H ₂ O / 10% D ₂ O	22 mM
T	Ethyl ferulate	DMSO- <i>d</i> ₆	200 mM
X	Brucine	CDCl ₃	50 mM
Z	Zolmitriptan	DMSO- <i>d</i> ₆	50 mM

Table 0.2: Samples used in this thesis. Note that concentrations are approximate and not necessarily constant, as samples were remade over time due to e.g. decomposition. However, it is reasonable to assume that the variation in concentration is below 10%. None of the results in this thesis depend on the exact concentrations of the samples used. See fig. 0.2 for chemical structures.

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List of acronyms

Many experiments in NMR spectroscopy are better known by their (occasionally very contrived) acronyms, and in my opinion, including the long name in the text often hinders rather than aids readability. I tabulate these and other abbreviations here for completeness.

2BOB	two-bond and one-bond correlation
2DJ	(2D) J-resolved
ADEQUATE	adequate sensitivity double-quantum spectroscopy
AP	antiphase
API	application programming interface
ASAP	acceleration by sharing adjacent polarisation
BANGO	β -TANGO (i.e. small flip angle)
BEBOP	broadband excitation by optimised pulses
BFGS	Broyden–Fletcher–Goldfarb–Shanno
BIBOP	broadband inversion by optimised pulses
BIG-BIRD	biselective independent gyrations BIRD
BIRD	bilinear rotation decoupling
BOBYQA	bound optimisation by quadratic approximation
BURBOP	broadband universal rotation by optimised pulses
CHORUS	chirped, ordered pulses for ultra-broadband spectroscopy
CLIP	clean in-phase
COSY	correlation spectroscopy
CPU	central processing unit
CRK	Cavanagh–Rance–Kay (seHSQC)
CTP	coherence transfer pathway
DEER	double electron–electron resonance
DFT	discrete Fourier transform
DIPSI	decoupling in the presence of scalar interactions
DMSO	dimethyl sulfoxide
DOSY	diffusion-ordered spectroscopy

dPSYCHE	discrete PSYCHE
DQF-COSY	double-quantum filtered COSY
DSE	double spin echo
DUMBO	decoupling using mind-boggling optimisation
EA	echo-antiecho
ECOSY	exclusive COSY
eDUMBO	experimental DUMBO
EPSI	echo-planar spectroscopic imaging
ES	excitation sculpting
ESCALADE	efficient spin control using analytical Lie algebraic derivatives
ESR	electron spin resonance
FE	function evaluation
FID	free induction decay
FT	Fourier transform
GEMSTONE	gradient-enhanced multiplet-selective targeted-observation NMR experiment
GENESIS	generation of supersequences <i>in silico</i>
GPU	graphics processing unit
H2OBC	heteronuclear two- and one-bond correlation
HMBC	heteronuclear multiple-bond correlation
HMQC	heteronuclear multiple-quantum correlation
HSQC	heteronuclear single-quantum correlation
iDOSY	internal DOSY
INADEQUATE	incredible natural-abundance double-quantum transfer experiment
INEPT	insensitive nuclei enhanced by polarisation transfer
IP	in-phase
IPAP	in-phase/antiphase
JRE	J-refocusing element
JRSE	J-refocused spin echo
JSON	JavaScript object notation
LP	linear prediction
LPJF	low-pass J-filter
MDS	multidirectional search
MLEV	Malcolm Levitt
NM	Nelder–Mead
NMR	nuclear magnetic resonance
NOAH	NMR by ordered acquisition using ^1H detection

NOE	nuclear Overhauser effect
NOESY	nuclear Overhauser effect spectroscopy
NUS	non-uniform sampling
PEP	preservation of equivalent pathways
POISE	parameter optimisation by iterative spectral evaluation
PSE	pure shift element
PSYCHE	pure shift yielded by chirp excitation
RF	radiofrequency
ROESY	rotating-frame Overhauser effect spectroscopy
rSNOB	refocusing selective excitation for biochemical applications
SAPPHIRE	sideband averaging by periodic phase incrementation of residual J evolution
seHSQC	sensitivity-enhanced HSQC
SERF	selective refocusing
SNR	signal-to-noise ratio
SW	spectral width / spectral window
TANGO	Testing for Adjacent Nuclei with a Gyration Operator
TIG-BIRD	triselective independent gyrations BIRD
TOCSY	total correlation spectroscopy
TPPI	time-proportional phase incrementation
TS	time-shared
TSE	triple spin echo
URL	uniform resource locator
URP	universal rotation pulse
ZF	zero filling
ZIP	<i>zz</i> isotope-selective pulse
ZQF	zero-quantum filter
ZS	Zanger-Sterk

Chapter 1

NMR theory

I have seen a great many lists of her drawing-up at various times of books that she meant to read regularly through—and very good lists they were—very well chosen, and very neatly arranged ... But I have done with expecting any course of steady reading from Emma. She will never submit to any thing requiring industry and patience.

— JANE AUSTEN, *Emma*

In the opening chapter of this thesis, I provide an overview of basic NMR theory, specifically, the dynamics of quantum systems containing one or more spin- $\frac{1}{2}$ particles. Starting from the Schrödinger equation, I progressively develop the rotating frame and density operator formalisms used in the analysis and simulation of simple NMR experiments. The important product operator formalism, used throughout this thesis, is exemplified through a selection of 1D and 2D experiments. Since 2D experiments form a very large part of this thesis, I also discuss a number of general principles in 2D NMR.

Note that this chapter is not intended to be an exhaustive account of magnetic resonance theory; for more complete coverage, the reader is directed to a suitable textbook.^{1–5}

1.1 Quantum mechanics

The most fundamental equation in (non-relativistic) quantum mechanics, which governs the time evolution of a quantum state $|\Psi(t)\rangle$ under a Hamiltonian H , is the time-dependent Schrödinger equation:

$$\frac{\partial |\Psi(t)\rangle}{\partial t} = -\frac{i}{\hbar} H |\Psi(t)\rangle \quad (1.1)$$

For a Hamiltonian which is constant over a period of time $t_1 \leq t \leq t_2$ (i.e. is *time-independent*), this can be integrated to yield an explicit solution:

$$|\Psi(t_2)\rangle = \exp \left[-\frac{iH(t_2 - t_1)}{\hbar} \right] |\Psi(t_1)\rangle \quad (1.2)$$

In NMR, it is conventional to use units of angular frequencies instead of energies, for example by replacing $H/\hbar \rightarrow H$; this will henceforth be assumed. The term $\exp[-iH(t_2 - t_1)]$ is called the *propagator* of the system and denoted $U(t_2, t_1)$; this is often further simplified to $U(\tau)$ where $\tau = t_2 - t_1$ is the duration of the evolution. For a Hamiltonian which varies with time but is piecewise constant, in that it can be broken up into several finite periods within which H is time-independent, the time evolution of the state is simply given by successive application of propagators:

$$|\Psi(t_n)\rangle = U(t_n, t_{n-1}) \cdots U(t_2, t_1) U(t_1, t_0) |\Psi(t_0)\rangle \quad (1.3)$$

where $t_n > t_{n-1} > \cdots > t_0$. The case where H continuously varies with time is more complicated, but we will not need to consider it in this thesis.

In NMR spectroscopy, we manipulate the *spin angular momentum* of atomic nuclei in order to obtain information about chemical structure and dynamics. The present work is restricted to nuclei with spin quantum number $I = 1/2$. These are two-level systems, where the eigenstates of I_z (denoted as $|\alpha\rangle$ and $|\beta\rangle$ for $m_I = +1/2$ and $-1/2$ respectively) are used as a standard basis, called the *Zeeman basis*. Primarily for mathematical convenience, the z -axis is conventionally chosen as the quantisation axis in textbook treatments of angular momentum. However, in the context of NMR, the z -axis bears even more significance as we define it to be the axis along which the static magnetic field is aligned. Since the matrix elements of an operator O are given by $O_{mn} = \langle m|O|n\rangle$, we can work out the matrix representations of the angular momentum operators in the Zeeman basis:

$$I_x = \frac{1}{2} \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}; \quad I_y = \frac{1}{2} \begin{pmatrix} 0 & -i \\ i & 0 \end{pmatrix}; \quad I_z = \frac{1}{2} \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix} \quad (1.4)$$

Their commutators are given by:

$$[I_i, I_j] = \sum_k i\varepsilon_{ijk} J_k, \quad (1.5)$$

where ε_{ijk} is the Levi-Civita symbol. We also define the following linear combinations:

$$\begin{aligned} I_+ &= I_x + iI_y = \begin{pmatrix} 0 & 1 \\ 0 & 0 \end{pmatrix}; & I_\alpha &= \frac{1}{2}E + I_z = \begin{pmatrix} 1 & 0 \\ 0 & 0 \end{pmatrix} \\ I_- &= I_x - iI_y = \begin{pmatrix} 0 & 0 \\ 1 & 0 \end{pmatrix}; & I_\beta &= \frac{1}{2}E - I_z = \begin{pmatrix} 0 & 0 \\ 0 & 1 \end{pmatrix} \end{aligned} \quad (1.6)$$

where E is the 2×2 identity matrix. The *coherence order* of an operator, denoted p , is defined by the Zeeman basis states it connects, i.e. the nonzero elements in its matrix form when expressed in this basis: an operator $O = |m_2\rangle\langle m_1|$ would represent $(m_2 - m_1)$ -order coherence, since $\langle m_2|O|m_1\rangle \neq 0$. Thus, in the above equations, $I_+ = |\alpha\rangle\langle\beta|$ represents a coherence order of +1; I_- a coherence order of -1; I_x and I_y are both a mixture of ± 1 -coherence; and the remainder have coherence order 0.

States (and operators) for composite systems are formally defined as tensor products of single-spin states (and operators).⁶ Operators on the same spin commute as per eq. (1.5), and operators on different spins fully commute. The Kronecker product allows these operators to be expressed in matrix form.⁵ For example, the operator $2I_xS_z$ can be represented as follows:^{*}

$$2I_xS_z = 2 \cdot \frac{1}{2} \cdot \frac{1}{2} \left[\begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix} \otimes \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix} \right] = \frac{1}{2} \begin{pmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \\ 1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 \end{pmatrix} \quad (1.7)$$

The Hamiltonians H for nuclear spin interactions, which will be encountered frequently in this chapter, are formed from such operators.³ In solution-state NMR, these interactions include:

$$H_{\text{cs}} = \sum_i \omega_{0,i} I_{iz} \quad (\text{chemical shift}) \quad (1.8)$$

$$H_J = \sum_{i>j} 2\pi J_{ij} (\mathbf{I}_i \cdot \mathbf{I}_j) \quad (\text{scalar coupling}) \quad (1.9)$$

$$H_{\text{pulse}} = \sum_i \omega_{i,x} I_{ix} + \sum_i \omega_{i,y} I_{iy} \quad (\text{radiofrequency pulses}) \quad (1.10)$$

$$H_{\text{grad}} = \sum_i \gamma_i Gz I_{iz} \quad (\text{pulsed field gradients on } z) \quad (1.11)$$

Pulsed field gradients (henceforth shortened to *gradients*) can in principle be applied along any axis, not just z , but this is dependent on hardware: all the work in this thesis was done on

*This representation is not unique; it is perfectly possible to reverse the order of the Kronecker product, and as long as this is consistently done, any physically measurable quantities calculated using this alternative will be the same.

z -gradient probes. In the above expressions:

- γ_i is the magnetogyric ratio of spin i ;
- $\omega_{0,i}$ refers to the Larmor, or precession, frequency of spin i (usually on the order of MHz). The Larmor frequency is defined as

$$\omega_{0,i} = -\gamma_i B_0, \quad (1.12)$$

where B_0 is the strength of the external (static) magnetic field;

- J_{ij} is the scalar coupling constant between spins i and j (expressed in units of Hz);
- ω_x and ω_y are amplitudes of radiofrequency (RF) pulses along the x - and y -axes, which are in general time-dependent, and are related to the so-called B_1 by a factor of γ_i .
- G is the amplitude of the gradient, typically in units of G/cm; and
- z is the position of the spin along the z -axis, typically in units of cm.

Finally, note that in the *weak coupling* regime where

$$\omega_{0,i} - \omega_{0,j} \gg J_{ij}, \quad (1.13)$$

the scalar coupling Hamiltonian may be simplified (the *secular approximation*^{*}) to

$$H_{J,\text{secular}} = \sum_{i>j} 2\pi J_{ij} I_{iz} I_{jz}. \quad (1.14)$$

This condition is always satisfied whenever spins i and j are different nuclides.

Throughout the course of an NMR experiment, RF pulses and gradients are turned on and off, and thus H_{pulse} and H_{grad} are time-dependent—although they will always satisfy the ‘piecewise constant’ criterion which allows us to use eq. (1.3). The ‘free precession’ (or simply ‘free’) Hamiltonian, H_{free} , refers to the Hamiltonian which is operative whenever no pulses or gradients are being applied:

$$H_{\text{free}} = H_{\text{cs}} + H_J. \quad (1.15)$$

*This result comes from the use of time-independent nondegenerate perturbation theory: it is based on the assumption that the eigenstates $\{|n\rangle\}$ of the main Hamiltonian H_0 are unchanged by the perturbation V (since the first-order correction varies as $\sum_m V_{mn}/(\omega_m - \omega_n) \ll 1$), and only the first-order correction to the energies $E_n^{(1)} = \langle n|V|n\rangle$ is retained. In this context, H_0 and V are respectively H_{cs} and H_J . When the condition eq. (1.13) does not hold, the nondegenerate treatment fails; see e.g. Sakurai.⁶

1.2 The rotating frame

The Hamiltonians described above refer to the ‘laboratory frame’ or the *Schrödinger picture*, where spins precess about the z -axis at their intrinsic frequencies and obey the equation of motion (1.1). However, this proves to often be unwieldy, in particular when analysing the effects of RF pulses. It is standard procedure to transform the frame of reference to a ‘rotating frame’, specifically, one which rotates about the z -axis at a defined rotation frequency ω_{rot} which is close to the Larmor frequencies ω_0 .

The rotating frame can be formalised using the *interaction picture* of quantum mechanics,⁶ which involves the separation of the Hamiltonian into two parts, with the first typically being completely time-independent:

$$H(t) = H_0 + H_1(t). \quad (1.16)$$

In this case, the static part H_0 simply corresponds to precession of the spins at a particular frequency:

$$H_0 = \sum_i \omega_{\text{rot},i} I_{iz}. \quad (1.17)$$

(Generally, each instance of the same nuclide (e.g. ^1H or ^{13}C) will share the same ω_{rot} , so the subscript i in $\omega_{\text{rot},i}$ is useful only for distinguishing different nuclear species.) This allows us to define H_1 as

$$\begin{aligned} H_1 &= H_{\text{J}} + H_{\text{pulse}} + H_{\text{grad}} + (H_{\text{cs}} - H_0) \\ &= H_{\text{J}} + H_{\text{pulse}} + H_{\text{grad}} + \sum_i \Omega_i I_{iz} \\ &= H_{\text{J}} + H_{\text{pulse}} + H_{\text{grad}} + H_{\text{offset}} \end{aligned} \quad (1.18)$$

where $\Omega_i = \omega_{0,i} - \omega_{\text{rot},i}$ is the *offset* of spin i . For reasons which will become clear later, the frequency ω_{rot} is chosen to be the centre of the spectral window for the given nuclide.

Having split up our Hamiltonian, we then define an *interaction-picture ket*:

$$|\Psi\rangle_I = \exp(iH_0t) |\Psi\rangle. \quad (1.19)$$

The time evolution of this ket is given by a transformation of the Schrödinger equation:

$$\begin{aligned}
 \frac{\partial |\Psi\rangle_I}{\partial t} &= iH_0 \exp(iH_0t) |\Psi\rangle + \exp(iH_0t) \frac{\partial |\Psi\rangle}{\partial t} \\
 &= iH_0 |\Psi\rangle_I + \exp(iH_0t)(-iH |\Psi\rangle) \\
 &= iH_0 |\Psi\rangle_I - i \exp(iH_0t)(H_0 + H_1) \exp(-iH_0t) |\Psi\rangle_I \\
 &= iH_0 |\Psi\rangle_I - iH_0 |\Psi\rangle_I - i \exp(iH_0t) H_1 \exp(-iH_0t) |\Psi\rangle_I \\
 &= -i \exp(iH_0t) H_1 \exp(-iH_0t) |\Psi\rangle_I \\
 &= -iH_{1,I} |\Psi\rangle_I,
 \end{aligned} \tag{1.20}$$

where

$$H_{1,I} = \exp(iH_0t) H_1 \exp(-iH_0t). \tag{1.21}$$

The underlying principle here is that the ‘interesting’ behaviour should be contained in H_1 , and instead of explicitly considering the time evolution under the ‘uninteresting’ H_0 , it is just used to transform H_1 into $H_{1,I}$.

We now turn our attention to how the various NMR Hamiltonians (eqs. (1.8) to (1.11)) are transformed in the interaction picture; that is to say, what the individual terms in the right-hand side of

$$\begin{aligned}
 H_{1,I} &= \exp(iH_0t) H_J \exp(-iH_0t) + \exp(iH_0t) H_{\text{pulse}} \exp(-iH_0t) \\
 &\quad + \exp(iH_0t) H_{\text{grad}} \exp(-iH_0t) + \exp(iH_0t) H_{\text{offset}} \exp(-iH_0t)
 \end{aligned} \tag{1.22}$$

are. We first note that H_0 (and hence $\exp(\pm iH_0t)$) is a function only of the I_{iz} operators; thus, any Hamiltonian which commutes with all I_{iz} 's will be untouched by this transformation. This is trivially true of H_{offset} and H_{grad} , which are themselves both functions of the I_{iz} 's. It can also be shown that H_J (in the homonuclear case) and $H_{J,\text{secular}}$ (heteronuclear case) fully commute with H_0 . So, for three of the four terms in eq. (1.22) we simply have the result that $\exp(iH_0t)H \exp(-iH_0t) = H$. This allows us to immediately write down the free precession Hamiltonian in the interaction picture:

$$H_{\text{free},I} = H_{\text{offset}} + H_J. \tag{1.23}$$

The fourth term, which does not commute with H_0 , is H_{pulse} . In the laboratory frame, *hard pulses* are applied as oscillating RF fields. Consider the case of a pulse acting on a single spin:

$$H_{\text{pulse}} = \omega_1 [\cos(\omega_{\text{rf}}t + \phi) I_x + \sin(\omega_{\text{rf}}t + \phi) I_y]. \tag{1.24}$$

Here, ω_1 represents the *amplitude* of the pulse, and ϕ the *phase*. This expression is similar to the expression in eq. (1.10), but here ω_1 and ϕ are both constants, with the time dependence explicitly specified using the *frequency* of the pulse, ω_{rf} . In the rotating frame, using that $H_0 = \omega_{\text{rot}} I_z$, we

then have the following interaction Hamiltonian:

$$H_{\text{pulse},I} = \omega_1 [\exp(i\omega_{\text{rot}}tI_z)I_x \cos(\omega_{\text{rf}}t + \phi) \exp(-i\omega_{\text{rot}}tI_z) + \exp(i\omega_{\text{rot}}tI_z)I_y \sin(\omega_{\text{rf}}t + \phi) \exp(-i\omega_{\text{rot}}tI_z)], \quad (1.25)$$

and using the formulae

$$\exp(i\theta I_z)I_x \exp(-i\theta I_z) = I_x \cos \theta - I_y \sin \theta \quad (1.26)$$

$$\exp(i\theta I_z)I_y \exp(-i\theta I_z) = I_y \cos \theta + I_x \sin \theta \quad (1.27)$$

(see Appendix A.2 of Levitt³ for a derivation), eq. (1.25) simplifies to

$$H_{\text{pulse},I} = \omega_1 [I_x \cos(\omega_{\text{rf}} - \omega_{\text{rot}} + \phi) + I_y \sin(\omega_{\text{rf}} - \omega_{\text{rot}} + \phi)]. \quad (1.28)$$

The frequency at which hard pulses are applied is termed the *transmitter frequency*, ω_{tx} . This is a parameter which can be controlled by the user, and is typically placed in the centre of the spectrum of the sample under study, in order to make the most use of its *bandwidth* (the region of frequencies over which the pulse is effective). For convenience, it is typical to then choose the rotating-frame frequency to be exactly the same frequency: $\omega_{\text{rot}} = \omega_{\text{tx}}$. This allows us to simplify the rotating-frame Hamiltonian to

$$H_{\text{pulse},I} = \omega_1 (I_x \cos \phi + I_y \sin \phi), \quad (1.29)$$

which is time-*independent*. Occasionally, I will also use the Cartesian components:

$$(c_x, c_y) = (\omega_1 \cos \phi, \omega_1 \sin \phi), \quad (1.30)$$

instead of the amplitude and phase, to describe the pulse.

Consider now the application of this pulse to an isolated spin for which $\omega_0 = \omega_{\text{rot}}$ and thus has an offset $\Omega = 0$. We have that $H_{\text{offset}} = H_J = H_{\text{grad}} = 0$, and the only active Hamiltonian is that of the pulse, which causes *nutation* of the spin magnetisation vector around the axis of the pulse; in this case, the pulse (or the spin) is said to be *on-resonance*.^{*} If a duration for the pulse τ_p is further specified, this also allows us to define a *flip angle* $\beta = \omega_1 \tau_p$. On the other hand, spins which are *off-resonance* ($\omega_0 - \omega_{\text{tx}} \neq 0$) evolve not only under the pulse Hamiltonian but also the offset; this leads to a different effective flip angle and axis of rotation. Off-resonance effects may be neglected when considering an idealised, infinitely hard pulse. However, this is of course not

^{*}Strictly speaking, the rotating frame is just a mathematical formalism, so the resonance condition does not necessitate $\Omega = 0$ or $\omega_{\text{rf}} = \omega_{\text{rot}} = \omega_0$. We only need that $\omega_{\text{rf}} = \omega_0$, or in other words, that the pulse is applied at the frequency of the spin—which may or may not be the same as the rotating-frame frequency. Practically, such a situation may arise in (for example) the application of selective pulses to a specific spin which is not at the centre of the spectrum.

possible on a spectrometer, and in practice off-resonance effects are noticeable even for hard pulses as short as several microseconds.

In general, RF pulses are more complicated than the simple case of the hard pulse shown here. For example, they may be constructed such that even in the rotating frame there is still a time dependence in the amplitudes and/or the phases; these are variously referred to as *shaped*, *amplitude-modulated*, or *frequency-modulated* pulses depending on the context. In principle, ω_1 and ϕ may both be continuous functions of time; however, for ease of construction and implementation, pulses are typically generated in a *piecewise* or discrete method using n points each of time δt , within which ω_1 and ϕ (or equivalently, c_x and c_y) are constant. The total length of the pulse is then simply $n(\delta t)$; δt is sometimes called the *timestep* of the pulse.

1.3 Density operators

NMR experiments are not executed on one single spin at a time; instead, the samples used typically contain on the order of 10^{20} spins. Furthermore, each of these spins may have its own wavefunction: it is generally impossible to force every spin to possess the same state. Since we are only interested in the *ensemble* behaviour such as expectation values, rather than the dynamics of each individual spin, we can use the *density operator* formalism instead of dealing with a composite wavefunction of many spins. The density operator, ρ , is defined (in the Schrödinger picture) as

$$\rho = \sum_j p_j |\psi_j\rangle \langle\psi_j| \quad (1.31)$$

where p_j is the probability that a spin is in the state $|\psi_j\rangle$ (and the $|\psi_j\rangle$'s are assumed to form a complete set of states).^{*} The use of ρ actually represents a loss of information, in that while eq. (1.31) gives us a straightforward recipe for constructing ρ from a given distribution of states $\{p_j, |\psi_j\rangle\}$, the reverse is not possible: given a known ρ , it is generally not possible to determine a unique distribution of states. This is not a problem, however, because ρ contains all the information required for the calculation of expectation values, in that for any operator A ,

$$\langle A \rangle = \sum_j \langle\psi_j|A\rho|\psi_j\rangle. \quad (1.32)$$

If A and ρ are expressed as matrices (through any choice of basis), then this is more easily expressed as the trace of the matrix product:

$$\langle A \rangle = \text{Tr}(A\rho). \quad (1.33)$$

^{*}This probability is a *classical* probability: that is, it is purely statistical in nature and should not be confused with the probability amplitudes associated with quantum superposition (i.e. $|c_j|^2$ in a single-spin wavefunction $\sum_j c_j |j\rangle$).

Other properties of the density operator are not discussed here, but can be found in virtually any textbook covering their use.^{6–8}

The time evolution of a Schrödinger-picture density operator is governed by the Liouville–von Neumann equation, which can be derived from eq. (1.1):

$$\begin{aligned}\frac{d\rho}{dt} &= \sum_j p_j \left(\frac{d|\psi_j\rangle}{dt} \langle\psi_j| + |\psi_j\rangle \frac{d\langle\psi_j|}{dt} \right) \\ &= \sum_j p_j (-iH|\psi_j\rangle\langle\psi_j| + |\psi_j\rangle i\langle\psi_j|H) \\ &= -iH \left(\sum_j p_j |\psi_j\rangle\langle\psi_j| \right) + i \left(\sum_j p_j |\psi_j\rangle\langle\psi_j| \right) H \\ &= -i[H, \rho].\end{aligned}\quad (1.34)$$

Note here that the weights p_j are time-independent, as the time evolution is contained entirely in the kets and bras.* For a time-independent H , this can be integrated to yield the solution:

$$\rho(t_2) = \exp(-iH\tau)\rho(t_1)\exp(iH\tau), \quad (1.35)$$

where $\tau = t_2 - t_1$.

In the interaction picture, the density operator is instead defined using interaction-picture states $\{|\psi_i\rangle_I\}$:

$$\begin{aligned}\rho_I &= \sum_j p_j |\psi_j\rangle_I\langle\psi_j|_I = \sum_j p_j \exp(iH_0t) |\psi_j\rangle\langle\psi_j| \exp(-iH_0t) \\ &= \exp(iH_0t) \left(\sum_j p_j |\psi_j\rangle\langle\psi_j| \right) \exp(-iH_0t) \\ &= \exp(iH_0t)\rho\exp(-iH_0t)\end{aligned}\quad (1.36)$$

(note the similarity to eq. (1.21)). Using a very similar proof as in eq. (1.34), it can be shown that ρ_I obeys a modified Liouville–von Neumann equation:

$$\frac{d\rho_I}{dt} = -i[H_{1,I}, \rho_I] \quad (1.37)$$

and analogously, for a time-independent $H_{1,I}$ we have that

$$\rho_I(t_2) = \exp(-iH_{1,I}\tau)\rho_I(t_1)\exp(iH_{1,I}\tau) = U\rho_I(t_1)U^\dagger, \quad (1.38)$$

*Strictly speaking, this only applies to a *closed* quantum system, which implies that effects such as relaxation are ignored (or at least, treated in only an empirical manner). The discussion of open quantum systems is beyond the scope of this work, but can be found elsewhere.^{9,10}

where $U = \exp(-iH_{1,I}\tau)$. Multiple propagators may be chained in a similar fashion to eq. (1.3). This result means that from a practical point of view, the effects of H_0 can be completely ignored when analysing or simulating NMR experiments using density operators.

Finally, a mention of the *equilibrium* or *thermal* density operator is in order. For a canonical ensemble, this is given by:

$$\rho_0 = \frac{\exp(-\beta\hbar H)}{\text{Tr}[\exp(-\beta\hbar H)]} \quad (1.39)$$

where $\beta = 1/(k_B T)$ and the Hamiltonian H is in units of angular momentum, as has been consistently used here. At equilibrium, no pulses or gradients are being applied, so the appropriate Hamiltonian is the free (Schrödinger-picture) Hamiltonian H_{free} (eq. (1.15)).^{*} Consider the case of a single spin: we have that $H_J = 0$, and hence $H_{\text{free}} = H_{\text{cs}} = \omega_0 I_z$. Thus,

$$\rho_0 = \frac{\exp(cI_z)}{\text{Tr}[\exp(cI_z)]} \approx \frac{E + cI_z}{\text{Tr}(E + cI_z)} = E + cI_z, \quad (1.40)$$

where $c = -\beta\hbar\omega_0 = -\hbar\omega_0/(k_B T)$ and E is the identity matrix; the approximation $\exp(cI_z) \approx E + cI_z$ is justified here as c is typically very small (on the order of 10^{-5}).

Throughout this thesis I consider only linear transformations of the form in eq. (1.38), which use unitary propagators of the form $U = \exp(-iH\tau)$:

$$U\rho_0 U^\dagger = U(E + cI_z)U^\dagger = UEU^\dagger + c(UI_zU^\dagger) = E + c(UI_zU^\dagger) \quad (1.41)$$

When describing NMR experiments, it is typical to simply ignore both the E term as well as the proportionality factor c , and focus only on the transformation of the I_z term. Thus, one may define a ‘simplified’ equilibrium density operator:[†]

$$\rho'_0 = I_z. \quad (1.42)$$

The E term is in fact truly inconsequential, as it cannot ever be transformed into detectable magnetisation. However, the constant c is still relevant: it is manifested in the magnitude of the NMR signal which is ultimately detected. It should be mentioned that ρ'_0 is not a true density operator: for example, $\text{Tr}(\rho'_0) = 0$ and not 1 as is required for a density operator. Nonetheless, all the physically interesting dynamics of the system such as expectation values are fully contained within ρ'_0 (at least up to the proportionality constant c).

^{*}The interaction-picture $H_{\text{free},I}$ would not be appropriate here, as its entire existence is merely a mathematical formalism. If that were not the case, it would imply that we can change the equilibrium state ρ_0 by simply *choosing* a different H_0 to factor out.

[†]This is similar to the ‘deviation’ density operator^{11,12} which measures how far a density operator deviates from the identity; but I have gone one step further in dropping the factor of c . Note that the alternative term ‘reduced density operator’ has a different meaning (it refers to the density operator of a subsystem, obtained by taking a partial trace over all other degrees of freedom).

1.4 Pulse sequences

It is impossible to provide a full overview of all, or even most, of the major NMR experiments in widespread use; the reader is directed to other books for this purpose.^{2,4,5,13,14} I seek only to (somewhat briefly) explain the general structure of one- and two-dimensional Fourier transform (FT) experiments, and in particular, how the formalisms developed in previous sections can be used to analyse and simulate such experiments.

1.4.1 1D pulse–acquire

Consider the simplest NMR experiment, a 1D ^1H pulse–acquire spectrum (§ 1.4.1). This consists of a 90° pulse, immediately followed by detection; for convenience, we will first consider the pulse as being applied along the $+y$ -axis, i.e. with a phase of $\phi = \pi/2$.



Figure 1.1: 1D ^1H pulse–acquire experiment.

To understand this, we begin with the thermal density operator $\rho'_0 = I_z$ (eq. (1.42)) and assume that there is only one spin in the sample, and that the pulse is applied on-resonance. The corresponding Hamiltonian during the pulse is simply $\omega_1 I_y$ (eq. (1.29)). If the duration of the pulse is τ_p , then the density operator immediately following the pulse is given by:

$$\rho = \exp(-i\omega_1 I_y \tau_p) I_z \exp(i\omega_1 I_y \tau_p) = \cos(\omega_1 \tau_p) I_z + \sin(\omega_1 \tau_p) I_x. \quad (1.43)$$

In this case, to obtain a 90° pulse, τ_p is specifically calibrated to ensure that $\omega_1 \tau_p = \pi/2$, which yields

$$\rho = I_x. \quad (1.44)$$

During the detection period, this term evolves under $H_{\text{free}} = H_{\text{cs}} = \omega_0 I_z$. (We use the Schrödinger-picture free Hamiltonian here because the measurement of the NMR signal takes place in the laboratory frame.) At a time t after detection has begun, the density operator is thus:

$$\rho(t) = \exp(-i\omega_0 I_z t) I_x \exp(i\omega_0 I_z t) = \cos(\omega_0 t) I_x + \sin(\omega_0 t) I_y. \quad (1.45)$$

The NMR signal derives from both x - and y -magnetisation (M_x and M_y), which are in turn proportional to I_x and I_y by a factor of γ . (If multiple spins are present, then each spin induces its own magnetisation: we would have that $M_x = \sum_i \gamma_i I_{ix}$, and likewise for M_y .) These are then

combined to form a complex signal (this process is known as *quadrature detection*):

$$s(t) = M_x(t) + iM_y(t) \quad (1.46)$$

$$\begin{aligned} &\propto \langle I_x(t) \rangle + i\langle I_y(t) \rangle \\ &= \text{Tr}[I_x\rho(t)] + i\text{Tr}[I_y\rho(t)] \\ &\propto \cos(\omega_0 t) + i \sin(\omega_0 t) \\ &= \exp(i\omega_0 t). \end{aligned} \quad (1.47)$$

Before the signal is digitised, the NMR spectrometer mixes this with a *reference* RF field oscillating at the transmitter frequency ω_{tx} . This results in downconversion of the detected frequencies by ω_{tx} , such that the actual digitised signal oscillates at the offset frequency Ω rather than ω_0 (recall we have chosen $\omega_{rot} = \omega_{tx}$, so $\omega_0 - \omega_{tx} = \omega_0 - \omega_{rot} = \Omega$). Therefore, instead of eq. (1.47), the signal we really see is:

$$s(t) \propto \exp(i\Omega t). \quad (1.48)$$

This result is the same as if we had pretended that during the detection period, ρ evolved under the *interaction-picture* free Hamiltonian $H_{free,I} = H_{offset}$; we will henceforth adopt this simplification, even though it is not physically accurate.

In practice, relaxation causes this signal to decay with time; this is frequently modelled as an exponential, in accordance with the Bloch equations:¹⁵

$$s(t) = \exp(i\Omega t) \exp(-t/T_2), \quad (1.49)$$

where T_2 is the transverse relaxation time constant.* The NMR signal is thus often called a *free induction decay* (FID). Fourier transformation of the FID then yields a spectrum with absorption- and dispersion-mode lineshapes in the real and imaginary parts respectively (fig. 1.2):

$$\begin{aligned} S(\omega) = F[s(t)] &= \frac{1}{\sqrt{2\pi}} \int_0^\infty s(t) \exp(-i\omega t) dt \\ &= \underbrace{\frac{k}{k^2 + (\omega - \Omega)^2}}_{A(\omega; \Omega)} + i \underbrace{\frac{\Omega - \omega}{k^2 + (\omega - \Omega)^2}}_{D(\omega; \Omega)}, \end{aligned} \quad (1.50)$$

where $k = 1/T_2$. The notation $A(\omega; \Omega)$ here means that the spectrum is a function of the frequency ω , but is parametrised by the peak offset Ω . Conventionally, only the real part of the spectrum is displayed, so it is desirable for the real part to contain the absorption-mode lineshape. This provides better resolution due to the narrower lineshape, and is also less affected by cancellation when multiple peaks overlap.

*Transverse (and longitudinal) relaxation are sometimes called spin–spin (and spin–lattice) relaxation, although the continued usage of these terms has been criticised.^{3,4,16}

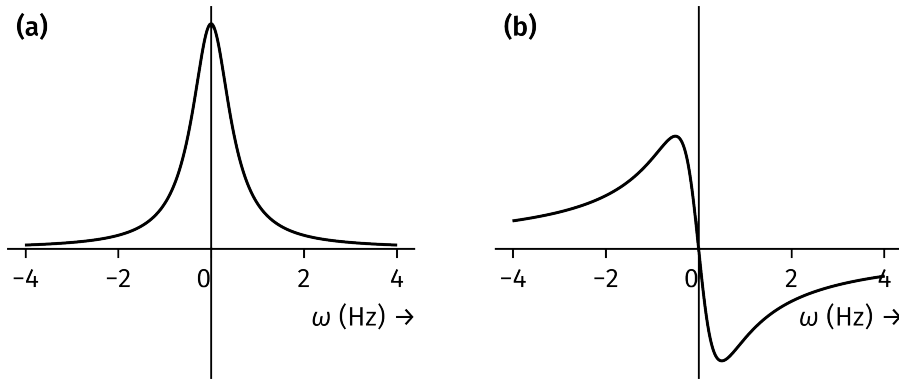


Figure 1.2: (a) Absorption-mode lineshape $A(\omega; \Omega = 0)$. (b) Dispersion-mode lineshape $D(\omega; \Omega = 0)$. Both lines have been plotted using $k = \pi \text{ rad s}^{-1}$.

Strictly speaking, the Lorentzian lineshapes above are only obtained when there is nonzero relaxation during the FID. For example, in the opposite limit where $k \rightarrow 0$, $A(\omega; \Omega)$ tends to a delta function $\delta(\omega = \Omega)$. However, for simplicity, in this thesis I will drop the relaxation term $\exp(-kt)$ unless absolutely necessary; I will simply pretend that a signal of the form $s(t) = \exp(i\Omega t)$ is directly Fourier transformed to give $A(\omega; \Omega) + iD(\omega; \Omega)$.

Consider now changing the initial pulse such that it is applied along the $+x$ -axis instead ($\phi = 0$). Repeating the above analysis, we find that the resulting signal will have a phase shift:

$$s'(t) = -i \exp(i\Omega t) \quad (1.51)$$

$$\Rightarrow S'(\omega) = F[s'(t)] = D(\omega; \Omega) - iA(\omega; \Omega). \quad (1.52)$$

If we were to take the real part of the spectrum here, then we would obtain the undesired dispersion-mode lineshape $D(\omega; \Omega)$. There are two ways of removing this phase shift. The first is to shift the *receiver phase* by ϕ_{rec} , which introduces an extra factor of $\exp(-i\phi_{\text{rec}})$ to the detected signal: we can thus choose $\phi_{\text{rec}} = 3\pi/2$ in order to cancel out the $-i$ term in $s'(t)$. Alternatively, the spectrum can be processed through *phase correction*, in which $S(\omega)$ is directly multiplied by a term $\exp(i\phi_{\text{corr}})$, where ϕ_{corr} is a linear function of the frequency ω :

$$\phi_{\text{corr}} = \phi_{\text{corr}}^{(0)} + \omega\phi_{\text{corr}}^{(1)}. \quad (1.53)$$

$\phi_{\text{corr}}^{(0)}$ and $\phi_{\text{corr}}^{(1)}$ are respectively termed the *zeroth-* and *first-order phase corrections*: in this idealised case, we can simply choose $(\phi_{\text{corr}}^{(0)}, \phi_{\text{corr}}^{(1)}) = (\pi/2, 0)$ to again remove the unwanted phase shift. More realistically, due to instrumental imperfections, both of these values will have to be nonzero in order to ensure that every peak in the spectrum has the correct phase, i.e. is displayed in absorption-mode.

An alternative framework for analysing pulse sequences is to use the ladder operators I_+ and I_- (eq. (1.6)). Using the original example with our initial pulse on $+y$, the density operator

immediately after the pulse is:

$$\rho = I_x = \frac{1}{2}(I_+ + I_-), \quad (1.54)$$

and during detection this evolves as:

$$\rho(t) = \cos(\Omega t)I_x + \sin(\Omega t)I_y = \frac{1}{2} [\exp(-i\Omega t)I_+ + \exp(i\Omega t)I_-]. \quad (1.55)$$

(Notice that the +1-coherence I_+ actually evolves at the negative frequency $-\Omega$.) To obtain the same signal as previously done in eq. (1.47), we ‘detect’ the I_- term:

$$s(t) \propto \text{Tr}[I_- \rho(t)] \propto \exp(i\Omega t), \quad (1.56)$$

which leads to the common assertion that *only quantum coherences of order -1 are detectable*. It is true that coherences with orders $p = 0, \pm 2, \pm 3, \dots$ can never be detected in an FID. However, it is worth pointing out that the ‘uniqueness’ of -1 -coherence is merely a result of how the x - and y -magnetisation are combined to form the complex signal (eq. (1.46)). We do not *physically* detect I_- : we detect I_x and I_y , and combine them to form a complex signal which is mathematically equal to detecting I_- . If we had instead chosen to combine them in a different way, such as $s(t) = M_x(t) - iM_y(t)$, this would give us $s(t) \propto \exp(-i\Omega t)$ —corresponding to ‘detection’ of +1-coherence—although this alternative does come with the drawback that frequencies must be reversed after Fourier transformation. In any case, we will stick to the established convention of detecting -1 -coherence here.

To end this section, it should be pointed out that the complex signal is not obtained as an infinitely-long, continuous function of time, as the treatment above implies. The complex-valued signal is digitised at an interval called the *dwell time*, τ_{dw} , and detection must be stopped after a finite period called the *acquisition time*, τ_{aq} . The transform being used is actually a discrete Fourier transform (DFT), which yields a periodic function $S(\omega)$; its period (in Hz) is given by $1/\tau_{\text{dw}}$.^{*} The NMR spectrum displayed to the user corresponds to one single period of $S(\omega)$, and thus the *spectral width* is also equal to $1/\tau_{\text{dw}}$.[†] In principle, the periodicity of the DFT means that signals which would ordinarily fall outside of the spectral width would appear at incorrect frequencies in the spectrum.¹⁷ On modern instrumentation, this is no longer the case for direct detection; peaks outside of the spectral width are removed using digital filters. However, *folding* or *aliasing* of peaks in the indirect dimension(s) of multidimensional NMR spectra still occurs.

^{*}The periodicity property of the DFT is equivalent to the Nyquist theorem, which is usually formulated as follows: the sampling rate required to correctly digitise a signal containing frequencies in the range $[0, f_{\text{max}}]$ is $1/(2f_{\text{max}})$. In the main text, it appears as if we have dropped the factor of 2 in the denominator; but in truth this statement of the Nyquist theorem is applicable to *real-valued* signals, and here we have a *complex-valued* signal $s(t)$, which effectively doubles the range of correctly sampled frequencies.

[†]Frustratingly, the DW parameter in Bruker’s TopSpin software is actually equal to $\tau_{\text{dw}}/2$. The reason is because this parameter corresponds to the interval between which *real* data is sampled, which is effectively twice as fast as complex-valued sampling.

The DFT $S(\omega)$ is also a discrete function itself, and its resolution is given by $1/\tau_{\text{aq}}$. It is possible to extend the effective acquisition time (and thus improve spectral resolution) without actually acquiring more data: this can be done either by *forward linear prediction* of the signal, or by simply adding zeros onto the end of the signal (*zero-filling*).

1.4.2 INEPT and product operators

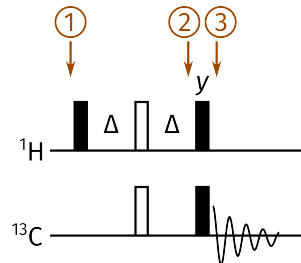


Figure 1.3: INEPT pulse sequence. The delay Δ is set to $1/(4 \cdot {}^1J_{\text{CH}})$.

Having tackled a simple single-spin case, we now move to the analysis of coupled spin systems and the development of the so-called ‘product operator formalism’.¹⁸ In particular, we look at the INEPT experiment,^{19,20} in which magnetisation is transferred from a nuclide with a high magnetogyric ratio to one with a low magnetogyric ratio through a scalar coupling: for example, from ${}^1\text{H}$ to ${}^{13}\text{C}$ using the one-bond coupling constant, ${}^1J_{\text{CH}}$ (fig. 1.3). Following tradition, the two nuclei are respectively labelled I and S .^{*} The Schrödinger-picture free Hamiltonian for a weakly coupled system (cf. eqs. (1.13) and (1.14)) is $H_{\text{free}} = \omega_{0,I}I_z + \omega_{0,S}S_z + 2\pi J_{IS}I_zS_z$. At the very beginning of the sequence (point ①), we formally have the equilibrium density operator

$$\rho_0 = \frac{\exp(-\beta\hbar H_{\text{free}})}{\text{Tr}[\exp(-\beta\hbar H_{\text{free}})]} \approx E - \beta\hbar(\omega_{0,I}I_z + \omega_{0,S}S_z + 2\pi J_{IS}I_zS_z), \quad (1.57)$$

using the same approximations as in eq. (1.40). The scalar coupling term can be safely neglected as $2\pi J_{IS}$ is several orders of magnitude smaller than the Larmor frequencies ω_0 . After removing the physically irrelevant E term and factoring out a constant of $\beta\hbar B_0$, we end up with:

$$\rho'_0 = \gamma_I I_z + \gamma_S S_z. \quad (1.58)$$

This represents equilibrium magnetisation (or *polarisation*) on both spins I and S , in proportion to their magnetogyric ratios. In general, an NMR experiment may manipulate—and ultimately detect—both of these terms. Since unitary evolution according to the Liouville–von Neumann equation is *linear*, in that $U(\rho_1 + \rho_2)U^\dagger = U\rho_1U^\dagger + U\rho_2U^\dagger$, we can treat these two terms separately:

^{*}This may seem insensible since I is the *sensitive* and S the *insensitive* nucleus, and indeed, in the original INEPT literature¹⁹ the meanings of I and S were swapped. However, this usage has not been universal,²¹ and in modern usage the identification of I as the sensitive nucleus seems to have prevailed.

we focus first on the spin- I polarisation, $\rho_I = \gamma_I I_z$. The first 90°_x ^1H pulse tips this magnetisation into the transverse plane (ignoring off-resonance effects):

$$\rho_I \rightarrow \exp[-i(\pi/2)I_x]\gamma_I I_z \exp[i(\pi/2)I_x] = -\gamma_I I_y. \quad (1.59)$$

In principle, we could continue in this manner through repeated application of the ‘sandwich’ formulae (eqs. (1.26) and (1.27), as well as an analogous version for the $I_z S_z$ term). For example, in the Δ delay which follows, we have that

$$\begin{aligned} \rho_I &\rightarrow -\gamma_I \exp(-iH_{\text{free},I}\Delta)I_y \exp(iH_{\text{free},I}\Delta) \\ &= -\gamma_I \exp(-iH_J\Delta) \exp(-iH_{\text{offset}}\Delta)I_y \exp(iH_{\text{offset}}\Delta) \exp(iH_J\Delta) \\ &= \dots \end{aligned} \quad (1.60)$$

When performing simulations of NMR experiments, such as those in later chapters, this is precisely what happens, with the slight difference that the Liouville–von Neumann equation (eq. (1.38)) is evaluated numerically rather than symbolically. Note that in going from the first to the second line, we can only ‘split up’ $H_{\text{free},I}$ into its constituent components H_{offset} and H_J because they commute.

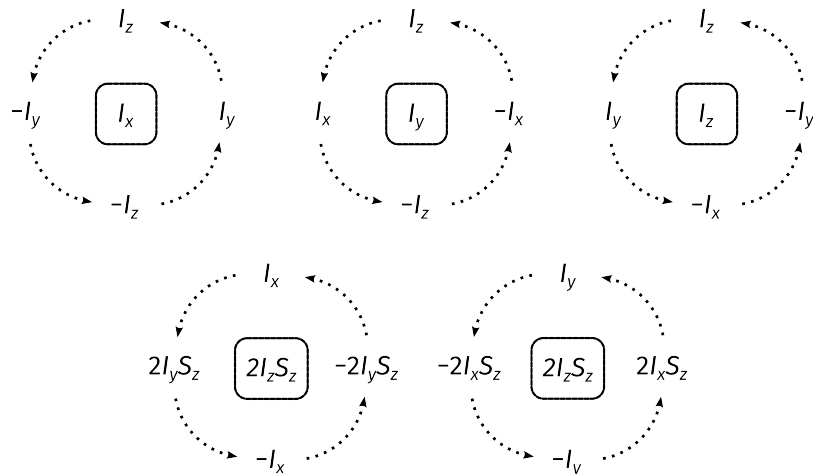


Figure 1.4: Simplified rules for the evolution of product operators under different common Hamiltonians (offset, weak/secular J-coupling, and pulses). These Hamiltonians often have the form ωM where M is some ‘base operator’, and are applied for a time τ . The boxed operators in the centre of each group refer to M ; the initial state is then ‘rotated’ about this by an angle of $\omega\tau$ to obtain the final state, or more formally, it is transformed into itself times $\cos(\omega\tau)$, plus the next term in the cycle times $\sin(\omega\tau)$. For example, a 90°_x pulse has the ‘base’ operator I_x and the angle $\omega\tau = \pi/2$; thus, the initial state I_z would be rotated to $I_z \cos(\pi/2) - I_y \sin(\pi/2) = -I_y$.

When analysing pulse sequences by hand, however, it is far more convenient to use a set of heuristics which summarise the effects of various pulse sequence elements. For example, fig. 1.4 summarises the evolution of a density operator under a single term of the Hamiltonian: as above,

since $[H_{\text{offset}}, H_J] = 0$, we only need to consider one term at a time. More high-level rules may be devised as well: for example, during the $\Delta-180^\circ_x(I, S)-\Delta$ spin echo which comes next, the J_{IS} interaction in $H_{\text{free},I}$ is allowed to evolve for a period of 2Δ , but the offset term is *refocused* and can be ignored. (The sign inversion caused by the 180° pulses must also be included.) As per fig. 1.4, this transforms the $-I_y$ term to $-2I_xS_z$ at point ②: the Hamiltonian is $\pi J_{IS}2I_zS_z$ for a total time of $2\Delta = 1/(2J_{IS})$, so the ‘angle’ rotated through is $\pi J_{IS}/(2J_{IS}) = \pi/2$. Immediately after this, the $90^\circ_x(I, S)$ pair of pulses rotates this magnetisation to $-2I_zS_y$ (point ③). These transformations are often denoted with simpler notation:

$$\gamma_I I_z \xrightarrow{90^\circ_x(I)} -\gamma_I I_y \xrightarrow{\Delta-180^\circ_x(I, S)-\Delta} -2\gamma_I I_x S_z \xrightarrow{90^\circ_y(I), 90^\circ_x(S)} -2\gamma_I I_z S_y \quad (1.61)$$

During the detection period, the term $-2\gamma_I I_z S_y$ evolves as

$$\begin{aligned} -2\gamma_I I_z S_y &\xrightarrow{H_{\text{free},I}} -2\gamma_I I_z S_y \cos(\Omega_S t) \cos(\pi J t) + \gamma_I S_x \cos(\Omega_S t) \sin(\pi J t) \\ &\quad - 2\gamma_I I_z S_x \sin(\Omega_S t) \cos(\pi J t) + \gamma_I S_y \sin(\Omega_S t) \sin(\pi J t), \end{aligned} \quad (1.62)$$

from which we extract the complex signal

$$s_I(t) = \langle S_x(t) \rangle + i\langle S_y(t) \rangle = \frac{\gamma_I}{2i} \{ \exp[i(\Omega_S + \pi J_{IS})t] - \exp[i(\Omega_S - \pi J_{IS})t] \}. \quad (1.63)$$

After Fourier transformation, the resulting spectrum has two peaks with opposite phase and have the frequencies $\Omega_S \pm \pi J_{IS}$; because of the factor of $1/(2i)$, the real part of the spectrum will contain dispersion-mode signals (fig. 1.5a). If desired, zeroth-order phase correction can be performed here, yielding instead a pair of absorption-mode signals still with opposite phases (fig. 1.5b). In either case, this is termed an *antiphase* doublet; the product operators which give rise to it ($2I_zS_x$ and $2I_zS_y$) are said to be antiphase with respect to spin I . Importantly, the amplitude of the signal scales as γ_I rather than γ_S ; since $\gamma_I > \gamma_S$, this represents a sensitivity enhancement compared to the direct excitation of S-magnetisation.

Of course, this is only half of the picture; we have not considered what happens to the other part of the magnetisation, namely $\rho_S = \gamma_S S_z$. Clearly, this is unaffected by the initial $90^\circ_x(I)$ pulse and the first Δ delay. The $180^\circ_x(S)$ pulse inverts it, and the final $90^\circ_x(S)$ pulse in fact transforms it into observable S-magnetisation:

$$\gamma_S S_z \xrightarrow{90^\circ_x(I)-\Delta} \gamma_S S_z \xrightarrow{180^\circ_x(I), 180^\circ_x(S)-\Delta} -\gamma_S S_z \xrightarrow{90^\circ_y(I), 90^\circ_x(S)} \gamma_S S_y \quad (1.64)$$

This term produces *in-phase* spin-S magnetisation during the detection period (where the two components of the doublet have the same phase):

$$s_S(t) = \frac{i\gamma_S}{2} \{ \exp[i(\Omega_S + \pi J_{IS})t] + \exp[i(\Omega_S - \pi J_{IS})t] \}, \quad (1.65)$$

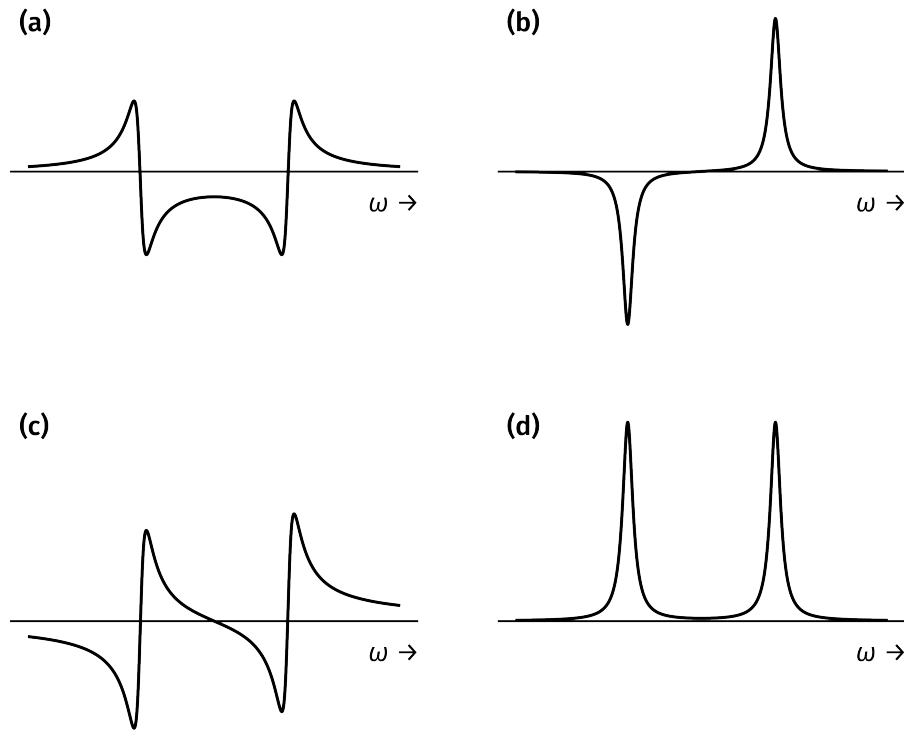


Figure 1.5: Peak shapes of a doublet. In all cases, the separation between the two peaks is $2\pi J_{IS}$. (a) Antiphase, dispersion-mode. (b) Antiphase, absorption-mode. (c) In-phase, dispersion-mode. (d) In-phase, absorption-mode.

Because of the factor of i , the real part of the spectrum will contain a dispersion-mode doublet (fig. 1.5c). The signal actually measured by the spectrometer is $s(t) = s_I(t) + s_S(t)$; and the spectrum is a weighted sum of in-phase and antiphase magnetisation. This leads to potentially unwanted phase distortions in the spectrum, which one would prefer to suppress.

This can be accomplished through the technique of *phase cycling*, where pulse and receiver phases are changed in concert and the resulting FIDs summed in order to select for a particular signal. In this case, the INEPT experiment is performed twice, once with the phases as given in fig. 1.3, and once where the initial $90^\circ_x(I)$ pulse is replaced with a $90^\circ_{-x}(I)$ pulse. The first of these gives us the same signals as above. However, inverting the initial I pulse leads to s_I acquiring a minus sign, because the initial I_z term is rotated to I_y instead of $-I_y$. On the other hand, the signal component s_S is unaffected by this pulse and thus does not experience a change of sign. The two FIDs we record are thus as follows:

$$s_1(t) = s_I(t) + s_S(t) \quad (1.66)$$

$$s_2(t) = -s_I(t) + s_S(t) \quad (1.67)$$

Simply taking the difference of these two FIDs yields a signal where the desired s_I has been accumulated and s_S has been cancelled out. In practice, instead of subtracting the two signals,

it is typical to shift the receiver phase ϕ_{rec} by 180° in the second experiment: this introduces a phase shift of $\exp(-i\pi) = -1$ to the signal, and the two signals can now be *added* together instead of subtracted to cancel out s_S . Since both ϕ_1 and ϕ_{rec} are 0 on the first experiment and π on the second experiment, we can express this as $\phi_x = \phi_{\text{rec}} = (0, \pi)$. This is more commonly denoted as $\phi_1 = \phi_{\text{rec}} = (x, -x)$, because the phases $(0, \pi)$ correspond to the $+x$ - and $-x$ -axes respectively.

The ‘simplified’ analysis of pulse sequences shown in eqs. (1.61) and (1.64) is often called ‘*product operator*’ analysis,¹⁸ because the underlying two-spin operators are products of single-spin Cartesian operators. Although this is often touted as being ‘simpler’ than full density operator calculations, it is really just a shorthand which masks the quantum mechanical theory developed in this chapter:

$$\underbrace{I_z \xrightarrow{90_x^\circ(I)} -I_y}_{\text{product operator}} \iff \underbrace{\exp(-iI_x\pi/2)I_z \exp(iI_x\pi/2) = -I_y}_{\text{density operator}} \quad (1.68)$$

Since the operators $\{E, I_x, I_y, I_z\}$ form a complete basis for a single-spin system, their products (i.e. product operators) likewise form a complete basis for multiple-spin systems, and so *any* density matrix for a multiple-spin system may be expressed as a linear combination of product operators. Thus, strictly speaking, the use of product operators therefore does not actually sacrifice any power in and of itself. However, the heuristics such as those in fig. 1.4 are limiting, in that the evolution under some Hamiltonians—for example, strong coupling $\mathbf{I} \cdot \mathbf{S}$, or pulses for off-resonance spins where H is a sum of I_x and I_z —cannot be neatly captured in such a pictorial form.

1.4.3 2D NMR: general principles

Much of this thesis is concerned with two-dimensional (2D) NMR experiments. Before considering an explicit example of a 2D experiment, we will first describe some general principles.^{22,23} 2D experiments contain one *indirect* and one *direct* time dimension, traditionally labelled t_1 and t_2 respectively. The t_1 period is a variable period which starts at 0^* and is incremented by a constant amount $\delta(t_1)$ on every iteration of the sequence. For each value of t_1 (or each t_1 *increment*) one complex signal $s'(t_2)$ is obtained. Putting this all together, the raw data thus take the form $s(t_1, t_2)$, which can be viewed as a 2D data matrix. Fourier transformation in both dimensions leads to a spectrum $S(\omega_1, \omega_2)$. In experimental contexts, the frequency dimensions are often referred to as F_1 and F_2 , but in this chapter I stick to the more mathematically consistent ω .

*Or as close to 0 as possible, considering that pulse elements during t_1 —such as the $180^\circ(I)$ pulse in fig. 1.6—require finite amounts of time. In some cases, it is possible to arrange spin echoes around t_1 such that the t_1 evolution on the first increment is refocused.

2D experiments generally comprise four components: *preparation*, *evolution*, *mixing*, and *detection*. The relevant product operators broadly follow this pattern:

$$\rho'_0 \xrightarrow{\text{preparation}} P \xrightarrow{\text{evolution}} P \cos(\Omega_P t_1) + P' \sin(\Omega_P t_1) \xrightarrow{\text{mixing}} Q \cos(\Omega_P t_1) + \dots \quad (1.69)$$

Here, P is some operator which, during the t_1 period, evolves into P' at a frequency of Ω_P (as per fig. 1.4). This leads to two terms which are *amplitude-modulated* in t_1 . The mixing process simply transforms the operator P to Q ; for now, we will assume that the sine-modulated term P' is turned into unobservable magnetisation.

During the detection period, the x - and y -magnetisation generated by the term Q is recorded. As discussed in § 1.4.1, Q must therefore be (or at least contain) -1 -quantum coherence on some spin, and thus evolves at the offset of that spin, Ω_Q . The complex signal $s(t_1, t_2)$ is therefore of the form $\cos(\Omega_P t_1) \exp(i\Omega_Q t_2)$. Note that, since the term P is not directly detected, it can have *any* coherence order; 2D NMR thus allows us to ‘detect’ frequencies of otherwise unobservable coherences.

We can equivalently express this signal as a sum of complex exponentials, which is easier to Fourier transform:

$$s(t_1, t_2) = \cos(\Omega_P t_1) \exp(i\Omega_Q t_2) = \frac{1}{2} [\exp(i\Omega_P t_1) \exp(i\Omega_Q t_2) + \exp(-i\Omega_P t_1) \exp(i\Omega_Q t_2)] \quad (1.70)$$

Fourier transformation of the first term reveals a peak centred at $(\omega_1, \omega_2) = (\Omega_P, \Omega_Q)$:

$$\begin{aligned} F[\exp(i\Omega_P t_1) \exp(i\Omega_Q t_2)] &= [A_1(\Omega_P) + iD_1(\Omega_P)][A_2(\Omega_Q) + iD_2(\Omega_Q)] \\ &= A_1(\Omega_P)A_2(\Omega_Q) - D_1(\Omega_P)D_2(\Omega_Q) \\ &\quad + i[A_1(\Omega_P)D_2(\Omega_Q) + D_1(\Omega_P)A_2(\Omega_Q)], \end{aligned} \quad (1.71)$$

where we have used the shorthand $A_i(\Omega)$ to denote what should properly be written as $A(\omega_i, \Omega)$. We should ideally like our 2D peaks to be absorption-mode in both dimensions, i.e. $S(\omega_1, \omega_2) = A_1(\Omega_P)A_2(\Omega_Q)$. Unfortunately, if we take the real part of this spectrum, we have an undesirable mixture of double absorption and double dispersion: this peak shape is called a *phase twist*.

This is not the only problem, however: if we perform the same Fourier transformation on the second term in eq. (1.70), we get *another* phase twist peak but this time centred at $(\omega_1, \omega_2) = (-\Omega_P, \Omega_Q)$. So, merely obtaining the complex signal $s(t) = \cos(\Omega_P t_1) \exp(i\Omega_Q t_1)$ is clearly not good enough:

- firstly, our spectra are not *pure phase* or *phase-sensitive*, in that the peaks are an inseparable mixture of absorptive and dispersive lineshapes;
- secondly, we have lost *quadrature detection* in the indirect dimension, in that we cannot

distinguish positive and negative offsets.

There are multiple different ways of solving these dual issues, which are extensively covered in NMR textbooks.^{1–4,14} Among these are the States method,²⁴ the time-proportional phase incrementation (TPPI) method,²⁵ and the echo–antiecho (EA) method. I will briefly cover the States and EA methods; the TPPI method can be shown to be essentially mathematically equivalent to the States method.^{26*}

In the States method, the cosine-modulated signal described above (and denoted $s_{\cos}(t_1, t_2)$ here) forms only one part of the signal. It is also necessary to acquire a *sine-modulated signal* of the form $s_{\sin}(t_1, t_2) = \sin(\Omega_P t_1) \exp(i\Omega_Q t_2)$. Once we have these two datasets, we can perform a Fourier transform along ω_2 first to get two intermediate signals:

$$s'_{\cos}(t_1, \omega_2) = \cos(\Omega_P t_1)[A_2(\Omega_Q) + iD_2(\Omega_Q)] \quad (1.72)$$

$$s'_{\sin}(t_1, \omega_2) = \sin(\Omega_P t_1)[A_2(\Omega_Q) + iD_2(\Omega_Q)]. \quad (1.73)$$

These idealised signals have the absorption-mode component A_2 as the real part in the ω_2 dimension. However, as before, an experimentally measured signal will typically require phase correction, which can be performed at this point by multiplication with a phase factor $\exp(i\phi_{\text{corr},2})$. After any phase correction, we discard the imaginary parts of these signals and use their real parts to construct another complex signal:

$$s''(t_1, \omega_2) = \Re\{s'_{\cos}(t_1, \omega_2)\} + i\Re\{s'_{\sin}(t_1, \omega_2)\} = \exp(i\Omega_P t_1)A_2(\Omega_Q). \quad (1.74)$$

Fourier transformation along ω_1 yields

$$S(\omega_1, \omega_2) = [A_1(\Omega_P) + iD_1(\Omega_P)]A_2(\Omega_Q), \quad (1.75)$$

which represents a peak only at the correct frequency (Ω_P, Ω_Q), and the real part of which is the desired double-absorption lineshape. Phase correction along ω_1 can now be carried out using a different phase factor $\exp(i\phi_{\text{corr},1})$. (In practice, the imaginary parts of s'_{\cos} and s'_{\sin} are not actually discarded: they are retained to allow phase correction in ω_2 to be performed at a later

*There is a slight difference in that the TPPI method pushes *axial peaks*—artefacts arising at $\omega_1 = 0$ —to the edge of the spectrum, whereas with the States method, these artefacts appear in the middle of the spectrum, potentially obscuring useful peaks. The reader is referred to the references cited in the main text for a discussion of this. In practice, both the States and EA methods can be easily adapted to incorporate this shifting of axial peaks by phase shifting the t_1 modulation as well as the receiver by 180° every time t_1 is incremented (in the former case it is creatively known as the *States–TPPI method*). Thus, I do not consider the TPPI method any further here.

stage if necessary.)^{*}

Naturally, the obvious question is how the sine-modulated signal $s_{\sin}(t_1, t_2)$ can be obtained. Returning to the product operators in eq. (1.69), we see that we can obtain this if we change the mixing period to transform $P' \rightarrow Q$ instead of $P \rightarrow Q$. This can usually be done by phase shifting one or more pulses after the t_1 period by 90° .[†] Alternatively (in fact, more commonly), we can modify the preparation period such that it produces an operator $-P'$ which rotates into P during t_1 . That way, after the t_1 evolution period we have a density operator of the form $-P' \cos(\omega_P t_1) + P \sin(\omega_P t_1)$; if we keep the mixing period the same then we will obtain the desired sine-modulated data. In contrast to before, this can be done by phase shifting one or more pulses before the t_1 period by -90° .

On the other hand, the EA method seeks instead to measure two signals which are *phase-modulated* in t_1 (instead of *amplitude-modulated* as in the States method):

$$s_{\text{echo}}(t_1, t_2) = \frac{1}{2} \exp(-i\Omega_P t_1) \exp(i\Omega_Q t_2) \quad (1.76)$$

$$s_{\text{antiecho}}(t_1, t_2) = \frac{1}{2} \exp(i\Omega_P t_1) \exp(i\Omega_Q t_2) \quad (1.77)$$

Once recorded, these signals can be added and subtracted to obtain the cosine- and sine-modulated signals of the States method; the standard States processing can then follow. The echo and antiecho signals can be obtained through the use of pulsed field gradients during the t_1 period, though a concrete example is deferred until the next section. The factor of $1/2$ in eqs. (1.76) and (1.77) requires an explanation: this arises because the ‘original’ signal is cosine- or sine-modulated, which can be thought of as a sum of two opposite phase modulations, i.e. $\cos(\Omega_P t_1) = [\exp(-i\Omega_P t_1) + \exp(i\Omega_P t_1)]/2$. The gradients effectively select for only one sense of

^{*}This treatment of the States method stems from the original paper²⁴ and is consistently reproduced in textbooks, although I have personally always found the steps rather contrived. I argue instead that a more coherent formulation can be given in terms of *quaternions*, a type of *hypercomplex number*, expressed as $a + ib + jc + kd$ where $a, b, c, d \in \mathbb{R}$, $i^2 = j^2 = k^2 = -1$, and $ij = -ji = k$. We can write that $s_{\cos} = \cos(\Omega_P t_1) \exp(j\Omega_Q t_2)$, and that $s_{\sin} = \sin(\Omega_P t_1) \exp(j\Omega_Q t_2)$; then, we can directly form a quaternionic 2D matrix $s(t_1, t_2) = s_{\cos} + is_{\sin} = \exp(i\Omega_P t_1) \exp(j\Omega_Q t_2)$. At this point, we can use a quaternion Fourier transform to *directly* obtain the data: $S(\omega_1, \omega_2) = (2\pi)^{-1} \iint \exp(-i\omega_1 t_1) s(t_1, t_2) \exp(-j\omega_2 t_2) dt_1 dt_2$, and phase correction in *both* dimensions can be carried out at will and simultaneously using $\exp(i\phi_{\text{corr},1})S(\omega_1, \omega_2)\exp(j\phi_{\text{corr},2})$, where the ϕ_{corr} ’s represent the phase corrections in the two respective frequency dimensions. (Note that quaternion multiplication is non-commutative, so the ordering of terms in these expressions matters.) Indeed, in the representation of 2D data as used by Bruker instruments, the four files `2rr`, `2ri`, `2ir`, and `2ii` essentially correspond to the four components (or *quadrants*) of a quaternionic $S(\omega_1, \omega_2)$. Of course, this is merely nice notation: none of the underlying science is changed. But the beauty of this is that we can consider the steps given in the main text to be an *implementation* of something more fundamental (the quaternion Fourier transform), a generalisation which naturally follows from the 1D case and immediately suggests its extension to the 3D case, rather than a *prescription* which—at least to a novice—seems to work only as if by magic. This treatment has been proposed before by Delsuc,²⁷ although it does not appear to have caught on much.

[†]Actually, if double-quantum coherence is sampled in t_1 , then the phase shift used must be halved, i.e. 45° ; and likewise for higher coherence orders. But we will not encounter such cases in this thesis.

the phase modulation and reject the other. Although this factor of $1/2$ can be cancelled out when combining the echo and antiecho datasets, in that:

$$s_{\cos,EA} = s_{\text{echo}} + s_{\text{antiecho}} = \cos(\Omega_P t_1) \exp(i\Omega_Q t_2) = s_{\cos} \quad (1.78)$$

(and likewise for the sine component), the process of adding up two separate datasets leads to a $\sqrt{2}$ increase in the noise level when compared to measuring s_{\cos} directly. Therefore, EA spectra have a $\sqrt{2}$ decrease in signal-to-noise ratio (SNR) compared to their States counterparts. This point is explored more fully in an article by Keeler and coworkers.²⁸ Despite this loss in SNR, the use of gradients typically leads to spectra with far better artefact suppression, completely obviating the need for long phase cycles. Consequently, a significant proportion of modern 2D experiments—especially heteronuclear experiments—use EA selection.

1.4.4 The States HSQC experiment

To illustrate the ideas developed in the previous section, we now turn our attention to two typical implementations of the 2D HSQC experiment, where both phase cycling as well as gradients are used to select for particular product operators. Figure 1.6 shows an HSQC experiment where quadrature detection in the indirect dimension is performed using the States method.

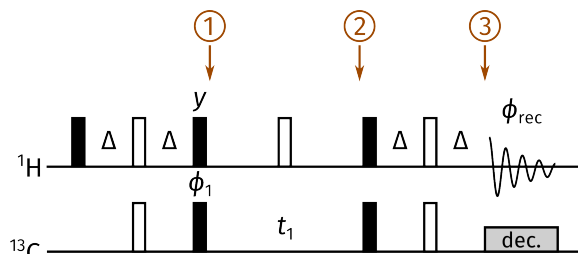


Figure 1.6: A typical HSQC pulse sequence utilising the States method for quadrature detection in ω_1 . The delay Δ is set to $1/(4 \cdot ^1J_{\text{CH}})$. To record the cosine-modulated dataset $s_{\cos}(t_1, t_2)$, we set $\phi_1 = \phi_{\text{rec}} = (x, -x)$. The sine-modulated dataset $s_{\sin}(t_1, t_2)$, on the other hand, is obtained using $\phi_1 = (-y, y)$ and $\phi_{\text{rec}} = (x, -x)$.

The HSQC experiment seeks to only detect protons directly bonded to ^{13}C ; the signals from all other protons must be suppressed. We first consider the simplest possible implementation of the HSQC, shown in fig. 1.6; as before, we will illustrate this with an *IS* spin pair. It can be shown that the equilibrium *S*-magnetisation cannot be transformed into observable *I*-magnetisation at the end of the sequence, so we will simply take $\rho'_0 = I_z$. Consider first the ‘basic’ case where $\phi_1 = x$. The *preparation period* is just the same INEPT element as analysed in § 1.4.2, so at point ① we already know that we have

$$\rho = -2I_zS_y. \quad (1.79)$$

During t_1 , the J-coupling is refocused, but the offset evolves to give:

$$\rho = 2I_z S_y \cos(\Omega_S t_1) - 2I_z S_x \sin(\Omega_S t_1), \quad (1.80)$$

at point ②, bearing in mind that the $180^\circ(I)$ pulse inverts the I_z component of both terms. Immediately after this, the reverse INEPT element (the ‘mixing’ segment) transfers the amplitude-modulated $2I_z S_y$ coherence I_x for detection. However, the $2I_z S_x$ term is turned into a mixture of unobservable zero- and double-quantum coherence, such that immediately before detection (point ③) we have:

$$\rho = I_x \cos(\Omega_S t_1) = \frac{1}{2}(I_+ + I_-) \cos(\Omega_S t_1). \quad (1.81)$$

Performing the usual quadrature detection in the direct t_2 dimension gives us the desired cosine-modulated signal of

$$s_{\cos}(t_1, t_2) = \frac{1}{2} \cos(\Omega_S t_1) \exp(i\Omega_I t_2), \quad (1.82)$$

In a similar way, it can be shown that setting $\phi_1 = -y$ yields the following after t_1 (point ②)

$$\rho = 2I_z S_x \cos(\Omega_S t_1) + 2I_z S_y \sin(\Omega_S t_1), \quad (1.83)$$

which after the reverse INEPT gives us the desired sine-modulated signal

$$s_{\sin}(t_1, t_2) = \frac{1}{2} \sin(\Omega_S t_1) \exp(i\Omega_I t_2). \quad (1.84)$$

These can be combined in the way described in § 1.4.3 to yield an HSQC spectrum with double absorption-mode lineshapes $A_1(\Omega_S)A_2(\Omega_I)$.

This alone is not sufficient to obtain a high-quality HSQC spectrum, however. Not every ^1H spin in a molecule will share a one-bond coupling to a ^{13}C spin: for example, in a natural-abundance sample, the HSQC experiment only detects around 1% of ^1H spins. Even small artefacts arising from the remaining 99% of proton magnetisation may have comparable intensity to the signals of interest. We therefore need to suppress any unwanted peaks which might arise from this ‘bulk’ magnetisation.

The simplest way to do this is to perform (at least) a two-step phase cycle, where the pulse phase ϕ_1 and receiver phase ϕ_{rec} are simultaneously inverted. This has no effect on the desired signal (it picks up two minus signs which cancel out), but will lead to cancellation of any signal arising from the bulk magnetisation, which does not evolve under H_J and therefore cannot be affected by the pulse on spin S. Note the similarity to the INEPT phase cycling in § 1.4.2, where we chose to cycle a pulse which the undesired pathway did not ‘experience’: this is not the only possible choice, but is one that is conceptually simple to understand. Naturally, this phase cycling must be carried out for both the cosine- and sine-modulated datasets. In practice, even longer phase

cycles are typically required to deal with experimental imperfections.

1.4.5 The echo–antiecho HSQC: gradients and coherence selection

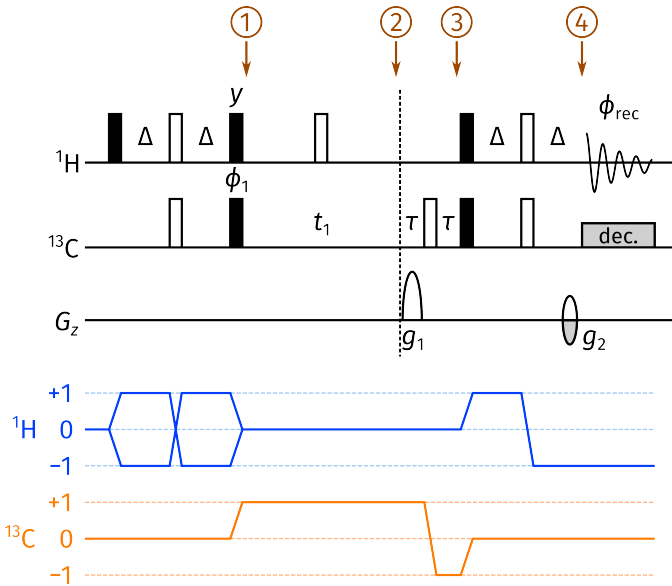


Figure 1.7: A typical EA HSQC pulse sequence. The delay Δ is set to $1/(4 \cdot ^1J_{\text{CH}})$. The gradient amplitudes are chosen such that $|g_1/g_2| = \gamma_{\text{H}}/\gamma_{\text{C}} \approx 4$. Specifically, the echo dataset is obtained by setting $(g_1, g_2) = (80\%, 20\%)$, and the antiecho dataset by setting $(g_1, g_2) = (80\%, -20\%)$, where gradient amplitudes are quoted as a percentage of the maximum gradient amplitude. Unlike in the States HSQC (fig. 1.6), phase cycling of ϕ_1 and ϕ_{rec} is no longer mandatory as the gradients g_1 and g_2 dephase unwanted magnetisation (although it can still be performed to attain *even* better artefact suppression). τ represents the duration of both gradients, and is usually on the order of 1 ms. The lines below the pulse sequence illustrate the coherence orders selected for during the echo experiment, which are collectively referred to as a coherence transfer pathway (CTP): for the antiecho experiment the ^{13}C CTP must be inverted.

The EA version of the HSQC experiment (fig. 1.7) is very similar to the States version discussed above. The only real difference is that two gradients are added: one immediately after t_1 , and one directly before detection. As we will see, the effect of this is to enforce a relationship between the coherence orders during the two gradients, or in other words, to select for a specific *coherence transfer pathway* (CTP), as illustrated by the lines beneath the pulse sequence in fig. 1.7.

Brute-force analysis

Recall that the Hamiltonian caused by a gradient of strength G is given by $H_{\text{grad}} = \sum_i \gamma_i G z I_{iz}$ (eq. (1.11)), where z is the z -position of the spin. In the case of our two-spin system, we can more explicitly write this as:

$$H_{\text{grad}} = \gamma_I G z I_z + \gamma_S G z S_z. \quad (1.85)$$

Points ① and ② are the same as in the States HSQC (fig. 1.6), so from eq. (1.80) we know that the density operator at point ② is:

$$\rho = 2I_zS_y \cos(\Omega_S t_1) - 2I_zS_x \sin(\Omega_S t_1). \quad (1.86)$$

In the next spin echo with the gradient g_1 , H_{offset} and H_J are refocused, so we can ignore their effects and focus only on H_{grad} . We assume here that the gradients are applied with duration τ .^{*} The I_z terms in ρ are unaffected by the gradient since they commute with H_{grad} ; however, the transverse S-magnetisation acquires a phase which depends on the z -position of the spin in the sample. Immediately after the gradient, we have:

$$\begin{aligned} \rho(z) &= 2I_zS_y \cos(\Omega_S t_1) \cos(\gamma_S g_1 z \tau) - 2I_zS_x \cos(\Omega_S t_1) \sin(\gamma_S g_1 z \tau) \\ &\quad - 2I_zS_x \sin(\Omega_S t_1) \cos(\gamma_S g_1 z \tau) - 2I_zS_y \sin(\Omega_S t_1) \sin(\gamma_S g_1 z \tau), \end{aligned} \quad (1.87)$$

where the notation $\rho(z)$ reminds us that this density matrix is spatially dependent. The $180^\circ_x(S)$ pulse then flips the S_y terms to give us, at point ③,

$$\begin{aligned} \rho(z) &= -2I_zS_y \cos(\Omega_S t_1) \cos(\gamma_S g_1 z \tau) - 2I_zS_x \cos(\Omega_S t_1) \sin(\gamma_S g_1 z \tau) \\ &\quad - 2I_zS_x \sin(\Omega_S t_1) \cos(\gamma_S g_1 z \tau) + 2I_zS_y \sin(\Omega_S t_1) \sin(\gamma_S g_1 z \tau). \end{aligned} \quad (1.88)$$

We know already from the States HSQC analysis that the mixing period (i.e. reverse INEPT) causes the transformation $2I_zS_y \rightarrow I_x$, and that the $2I_zS_x$ term is lost. So, if the gradient g_2 were absent, we would have the following terms at point ④:

$$\begin{aligned} \rho(z) &= -I_x \cos(\Omega_S t_1) \cos(\gamma_S g_1 z \tau) + I_x \sin(\Omega_S t_1) \sin(\gamma_S g_1 z \tau) \\ &= -I_x \cos(\Omega_S t_1 + \gamma_S g_1 z \tau). \end{aligned} \quad (1.89)$$

This is of course not the case. In principle, the last Δ delay should be split up into two parts: one of duration $(\Delta - \tau)$ where only $H_{\text{free},I}$ is active, and one of duration τ where the Hamiltonian $H_{\text{free},I} + H_{\text{grad}}$ operates. Thankfully, we know that H_{grad} commutes with $H_{\text{free},I}$, so we can in fact separate the relevant propagators:

$$\begin{aligned} &\exp[-i(H_{\text{free},I} + H_{\text{grad}})\tau] \exp[-iH_{\text{free},I}(\Delta - \tau)] \\ &= \exp(-iH_{\text{grad}}\tau) \exp(-iH_{\text{free},I}\tau) \exp[-iH_{\text{free},I}(\Delta - \tau)] \\ &= \exp(-iH_{\text{grad}}\tau) \exp(-iH_{\text{free},I}\Delta), \end{aligned} \quad (1.90)$$

where the first term is just the gradient on its own, and the second is just the delay without a gradient. So, we can directly apply $\exp(-iH_{\text{grad}}\tau)$ to the density operator in eq. (1.89) to

^{*}In practice, we also need to include a gradient recovery delay immediately after the gradient to allow for the dissipation of eddy currents, which causes the spin echo to be lengthened slightly. This is inconsequential to the analysis and will be ignored here.

get:^{*}

$$\rho(z) = -I_x \cos(\Omega_S t_1 + \gamma_S g_1 z \tau) \cos(\gamma_I g_2 z \tau) - I_y \cos(\Omega_S t_1 + \gamma_S g_1 z \tau) \sin(\gamma_I g_2 z \tau) \quad (1.91)$$

$$\begin{aligned} &= -\frac{1}{2} I_x \cos[\Omega_S t_1 + (\gamma_S g_1 + \gamma_I g_2) z \tau] - \frac{1}{2} I_x \cos[\Omega_S t_1 + (\gamma_S g_1 - \gamma_I g_2) z \tau] \\ &\quad - \frac{1}{2} I_y \sin[\Omega_S t_1 + (\gamma_S g_1 + \gamma_I g_2) z \tau] + \frac{1}{2} I_y \sin[\Omega_S t_1 + (\gamma_S g_1 - \gamma_I g_2) z \tau] \end{aligned} \quad (1.92)$$

(as a sanity check, it can be verified that this reduces to eq. (1.89) if $g_2 = 0$). The signal which we detect stems from the entire sample length, so we in fact need to perform an integration over z :

$$\rho = \frac{1}{L} \cdot \int_{-L/2}^{L/2} \rho(z) dz, \quad (1.93)$$

where the sample length is L and $z = 0$ is assumed to be the middle of the sample.

For the *echo* experiment, we choose the ratio $g_1/g_2 = \gamma_I/\gamma_S$; this means that $\gamma_S g_1 - \gamma_I g_2 = 0$. The second and fourth terms in eq. (1.92) thus *lose* their z -dependence; when integrated over z these terms are unchanged. On the other hand, the first and third terms are attenuated by a factor proportional to $\int_{-L/2}^{L/2} \cos[(\gamma_S g_1 + \gamma_I g_2) z \tau] dz$ (or the equivalent sine-modulated integrand). The term in square brackets here can be identified as the spatially dependent phase caused by the evolution under the gradient pulse; we say that the gradients *dephase* coherences. (In the case of the second and fourth terms, they also *rephase* coherences.) If the gradient strengths (g_1, g_2) and/or their durations τ are sufficiently large, the integral evaluates to a small number which is effectively zero, corresponding to complete dephasing. The result is that, just before detection, we have:

$$\rho_{\text{echo}} = -\frac{1}{2} I_x \cos(\Omega_S t_1) + \frac{1}{2} I_y \sin(\Omega_S t_1), \quad (1.94)$$

or, extracting the detectable I_- components,

$$\rho_{\text{echo}} = -\frac{1}{4} I_- \cos(\Omega_S t_1) + \frac{i}{4} I_- \sin(\Omega_S t_1) = -\frac{1}{4} I_- \exp(-i\Omega_S t_1). \quad (1.95)$$

Detection of this in t_2 gives us the echo signal

$$s_{\text{echo}}(t_1, t_2) = -\frac{1}{4} \exp(-i\Omega_S t_1) \exp(i\Omega_I t_2), \quad (1.96)$$

in accordance with eq. (1.76). Note that the prefactor here is $-1/4$, instead of $1/2$ in the States method (eqs. (1.82) and (1.84)): this accounts for the decreased SNR in the EA method as previously described. (The minus sign comes from the extra $180^\circ(S)$ pulse after t_1 , but is incon-

^{*}We have implicitly assumed here that the z -coordinate of the *IS* spin pair during the first gradient is the same as its z -coordinate during g_2 , or in other words, that there is no *diffusion* or *convection* between the two gradients. In general, this is not true, and these effects will lead to a loss of signal as the rephasing by the second gradient is not perfect.

sequential as it can be removed by phase correction.)

In a similar way, it can be shown that if we invert the sign of g_2 , we have that $g_1/g_2 = -\gamma_I/\gamma_S$. Now, the second and fourth terms in eq. (1.92) are dephased, and we get the antiecho spectrum from the first and third terms:

$$s_{\text{antiecho}}(t_1, t_2) = -\frac{1}{4} \exp(i\Omega_S t_1) \exp(i\Omega_I t_2). \quad (1.97)$$

The shift basis

In the above treatment, I have only used the ‘rules’ developed so far for Cartesian product operators to explain the effects of gradients. *This is clearly very tedious!* When gradients are involved, it proves simpler to use a different basis, specifically $\{E, I_z, I_+, I_-\}$: this is called the *shift basis*.^{*} The coherence orders of these, and their products, are easily read off from the terms involved: for example, $I_- S_-$ is double-quantum coherence with $p = -2$. The evolution of these terms under various Hamiltonians is summarised by Keeler,⁴ but two cases are particularly simple and important here:

1. 180°_x pulses invert I_z and interchange $I_+ \leftrightarrow I_-$;
2. An operator I_{ip} , on a spin i and with coherence order p , evolves under the Hamiltonian ωI_z for a time t to give:

$$I_{ip} \longrightarrow I_{ip} \exp(-ip\omega t) \quad (1.98)$$

We have seen examples of the latter before: compare, for example, eqs. (1.54) and (1.55). Being flexible in switching between the two bases can greatly simplify the mathematics involved, as terms which do not survive can be immediately dropped, and the simpler phase modulation $\exp(i\omega t)$ can be used instead of unwieldy $\cos(\omega t)$ and $\sin(\omega t)$ terms.

Consider now how a spatially dependent phase is imparted to a coherence as it proceeds through the HSQC sequence. We assume that between the two gradients the coherence I_{ip} is transferred to I_{jq} , i.e. q -order coherence on some other spin j :

$$I_{ip} \xrightarrow{g_1} I_{ip} \exp(-ip\gamma_i g_1 z\tau) \xrightarrow{\text{mixing}} I_{jq} \xrightarrow{g_2} I_{jq} \exp(-iq\gamma_j g_2 z\tau) \exp(-ip\gamma_i g_1 zt). \quad (1.99)$$

For the coherence to be rephased, we require that

$$p\gamma_i g_1 \tau + q\gamma_j g_2 \tau = 0, \quad (1.100)$$

^{*}The normalisation factors of each of these operators may be modified slightly for consistency, as in Cavanagh et al.,² but this is largely inconsequential for the purposes of this section.

and in the specific case of the HSQC, we know that spins i and j are respectively S and I , so

$$p\gamma_S g_1 + q\gamma_I g_2 = 0. \quad (1.101)$$

For the echo experiment, we choose $g_1/g_2 = \gamma_I/\gamma_S$, which means that eq. (1.101) is satisfied if and only if $p + q = 0$. Since I_q is only detectable if $q = -1$, this means that $p = +1$.

With this knowledge, we can work through the pulse sequence analysis with much greater facility. At the start of t_1 , we have the operator $-2I_z S_y = i(I_z S_+ - I_z S_-)$: of these two terms, the gradient combination selects for the $I_z S_+$ term during t_1 (for which $p = +1$), and rejects the $I_z S_-$ term ($p = -1$). This evolves during t_1 (and the $180^\circ(I)$ pulse) to give

$$-iI_z S_+ \exp(-i\Omega_S t_1). \quad (1.102)$$

The gradient echo after t_1 transforms this to

$$-iI_z S_- \exp(-i\Omega_S t_1) \exp(-i\gamma_S g_1 z\tau) = -iI_z (S_x - iS_y) \exp(-i\Omega_S t_1) \exp(-i\gamma_S g_1 z\tau). \quad (1.103)$$

Again, the mixing period only transforms $2I_z S_y \rightarrow I_x$, so we can discard the $I_z S_x$ term and get

$$-\frac{1}{2}I_x \exp(-i\Omega_S t_1) \exp(-i\gamma_S g_1 z\tau) = -\frac{1}{4}(I_+ + I_-) \exp(-i\Omega_S t_1) \exp(-i\gamma_S g_1 z\tau) \quad (1.104)$$

just before applying the g_2 gradient. The rest of the story is the same as before: the I_- component is rephased by g_2 (it picks up another $\exp(i\gamma_I g_2 z\tau)$ term, which cancels out with the phase from the first gradient), and is then detected to yield $s_{\text{echo}}(t_1, t_2)$ (eq. (1.96)). This *CTP selection* process is frequently depicted in the diagrammatic form of fig. 1.7, where solid lines indicate the coherence orders which are selected for at each stage of the pulse sequence. In this case, during the gradient g_1 , we have $+1$ -coherence on ^{13}C ; and during the gradient g_2 , we have -1 -coherence on ^1H .

Conversely, for the antiecho experiment, the gradient ratio $g_1/g_2 = -\gamma_I/\gamma_S$ necessitates instead that $p - q = 0$. This therefore selects for the $I_z S_-$ term during t_1 , and the remaining analysis can be carried out along the same lines. The relevant CTP diagram is similar to that in fig. 1.7, except that the ^{13}C component (orange line) is inverted.

Finally, we note here that any signal generated from the bulk magnetisation (protons not directly coupled to ^{13}C) cannot be rephased by these gradients. Since such magnetisation is always on spin I , this would require that

$$p\gamma_I g_1 + q\gamma_I g_2 = 0 \Leftrightarrow p\gamma_I + q\gamma_S = 0, \quad (1.105)$$

which cannot be satisfied for any sensible integer values of p and q except for $p = q = 0$, which is not observable during t_2 anyway. So, the CTP gradients effectively remove all unwanted signals arising from the bulk magnetisation: the cycling of ϕ_1 and ϕ_{rec} done in the States experiment is rendered unnecessary.

The points developed in this chapter are neatly demonstrated in fig. 1.8. The States–TPPI experiment used here is the same as the States experiment in fig. 1.6, except that on every t_1 increment ϕ_1 and ϕ_{rec} are inverted, causing the artefacts to be shifted to the edge of the spectrum. Figures 1.8a and 1.8b show the States–TPPI and EA HSQC spectra obtained with one scan per increment. Comparing the projections of these two spectra (fig. 1.8d), it can be seen that the SNR of the States–TPPI version is greater by a factor of $\sqrt{2}$. However, the spectral quality of the States–TPPI spectrum is far poorer: presentable data can only be obtained when using a two-step phase cycle, as was done in fig. 1.8c. In this case, although the EA spectrum sacrifices some SNR, its sensitivity is still perfectly usable: the improved spectral quality thus makes EA the preferred implementation for many heteronuclear experiments. For homonuclear experiments which do not have such stringent artefact suppression requirements, States–TPPI quadrature detection is still commonly used.

Coherence transfer pathways

To end this chapter, we generalise the CTP refocusing requirement introduced in eq. (1.100). The treatment here is similar to that of Mitschang et al.²⁹ In general, during a pulse sequence we may have n gradients in total, with amplitudes $g^{(1)}, g^{(2)}, \dots, g^{(n)}$. We assume that their durations $\tau^{(i)}$ are all the same, such that they can be factored out of the equation. We express the coherence during the i -th gradient as a product of single-spin coherences

$$M^{(i)} = \prod_j^{\text{spins}} I_j(p_j^{(i)}), \quad (1.106)$$

where $I_j(p_j^{(i)})$ represents $p_j^{(i)}$ -order coherence on spin j during gradient i . The spatially dependent phase imparted by the gradient $g^{(i)}$ is the sum of the phases acquired by each individual coherence:

$$\phi^{(i)} = -zg^{(i)} \sum_j p_j^{(i)} \gamma_j \quad (1.107)$$

For rephasing of a CTP, we require that $\sum_i \phi^{(i)} = 0$, or equivalently

$$\sum_i \left(g^{(i)} \sum_j p_j^{(i)} \gamma_j \right) = 0. \quad (1.108)$$

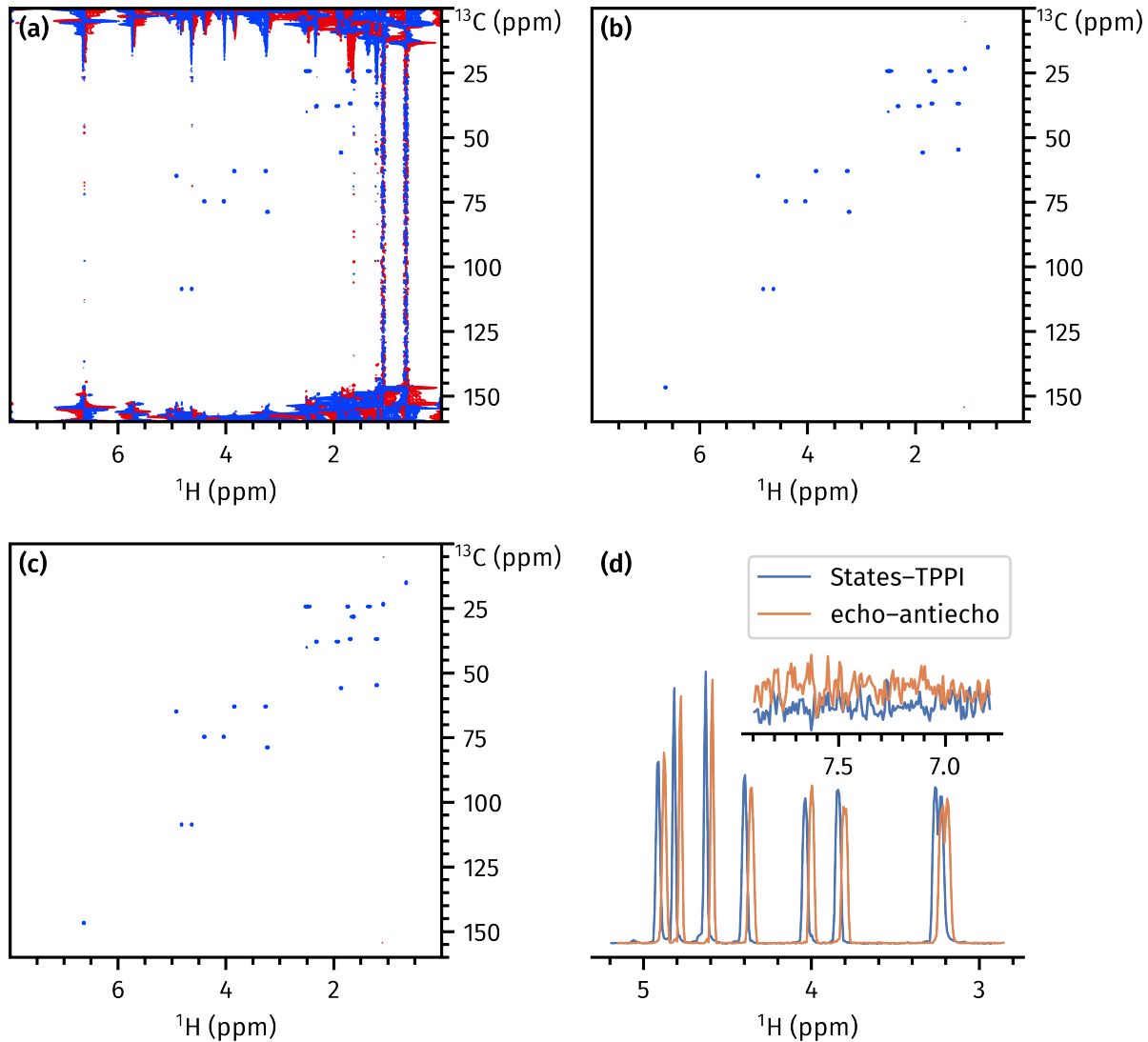


Figure 1.8: A comparison of HSQC data acquired using various quadrature detection schemes. (a) States-TPPI HSQC with one scan (i.e. no phase cycling) applied. (b) EA HSQC with one scan. (c) States-TPPI HSQC with two scans, using the phase cycling shown in fig. 1.6. The artefacts are essentially completely removed. (d) Projections of the States-TPPI and EA spectra in (a) and (b) onto the F_2 axis (to avoid picking up the artefacts, only the region between 60 and 110 ppm in F_1 was projected). The two projections are slightly horizontally offset for visual clarity. The signal intensity is the same, but the noise level in the EA spectrum is larger (by a factor of $\sqrt{2}$, as described in the text). Data code: 6A-220809.

We can now define a *weighted coherence order*³⁰ as

$$p^{(i)} = \sum_j p_j^{(i)} \gamma_j. \quad (1.109)$$

For example, the weighted coherence order for I_+S_- is simply $\gamma_I - \gamma_S$.^{*} This allows the rephasing condition to be very simply expressed as a scalar product:

$$\sum_i g^{(i)} p^{(i)} = \mathbf{g} \cdot \mathbf{p} = 0. \quad (1.110)$$

$\mathbf{g} \cdot \mathbf{p}$ can be considered to be an ‘extent of dephasing’: ideally, gradient amplitudes \mathbf{g} are chosen such that the desired CTP \mathbf{p} is rephased (eq. (1.110)), and undesired CTPs \mathbf{p}' are suppressed as much as possible in that $\mathbf{g} \cdot \mathbf{p}'$ is maximised.

1.5 References

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*Mitschang et al. define a *composite coherence order* as $\sum_j p_j^{(i)} \gamma_j / \gamma_I$, using some nuclide I as a reference. This follows an earlier paper³⁰ where I is explicitly chosen to be ^1H , and has the advantage that if the experiment is homonuclear (i.e. all the spins j are simply I), the composite coherence order is the same as the coherence order. Here, I prefer not to tie the analysis to a particular nuclide as the choice of I may, in principle, depend on the experiment under consideration. This different definition naturally induces a different choice of terminology.

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Chapter 2

Pure shift NMR

Then seating herself with a gentleman on each side, she ... discussed the possibility of improvements with much animation. Nothing was fixed on—but Henry Crawford was full of ideas and projects, and, generally speaking, whatever he proposed was immediately approved.

— JANE AUSTEN, *Mansfield Park*

Pure shift NMR refers to the technique of acquiring *broadband homodecoupled* NMR spectra, where all multiplet structure is suppressed and each chemical environment gives rise to a singlet.^{1,2} Here, ‘broadband’ means that the couplings are removed from the entire spectrum, and ‘homodecoupled’ refers to the removal of homonuclear couplings: unlike heteronuclear couplings, these cannot be removed by applying RF pulses during the acquisition period, as that will destroy the desired signal itself. Although pure shift techniques can be applied to any nuclide, ¹H spectra are of greatest interest because of the narrow chemical shift range of ¹H which often leads to peak overlap, as well as the abundance of ¹H–¹H couplings in typical organic molecules.

In the first two sections, I first cover the theory underpinning, and a brief history of, pure shift experiments. I then describe my work towards increasing the quality of pure shift experiments: this is measured both in terms of *sensitivity* as well as *purity*, i.e. the lack of spectral artefacts arising from imperfect decoupling. Each of the approaches described here is compared against the PSYCHE pure shift method, which is the current state of the art. Finally, I end with a section discussing the combination of pure shift diffusion spectroscopy—formally a pseudo-3D experiment—with the use of ultrafast NMR techniques to collapse the diffusion dimension. This last project was carried out in collaboration with Jean-Nicolas Dumez (University of Nantes).

The work in this chapter has not been submitted for publication.

2.1 Theoretical background

In chapter 1, I showed how density operators could be expressed in either of the two bases $\{E, I_x, I_y, I_z\}$ or $\{E, I_z, I_+, I_-\}$, depending on which was most mathematically expedient. To analyse pure shift NMR, it turns out to be most convenient to introduce a third basis, namely $\{I_\alpha, I_\beta, I_+, I_-\}$.³⁻⁵ The definitions of these terms are given in eq. (1.6): it is clear from there that, when expressed in the Zeeman basis, each matrix element of the density operator corresponds to one of these terms:

$$\rho = \begin{pmatrix} \rho_\alpha & \rho_+ \\ \rho_- & \rho_\beta \end{pmatrix} = \rho_\alpha I_\alpha + \rho_+ I_+ + \rho_- I_- + \rho_\beta I_\beta. \quad (2.1)$$

In systems containing multiple spins, the corresponding *single-element operators* are just the products of these operators. In this section, I use a system with two weakly coupled spins I_1 and I_2 to illustrate the ideas behind pure shift NMR. We have that $H_{\text{free},I} = \Omega_1 I_{1z} + \Omega_2 I_{2z} + 2\pi J I_{1z} I_{2z}$, which is diagonal in the Zeeman basis:

$$H_{\text{free},I} = \begin{pmatrix} \omega_{\alpha\alpha} & 0 & 0 & 0 \\ 0 & \omega_{\alpha\beta} & 0 & 0 \\ 0 & 0 & \omega_{\beta\alpha} & 0 \\ 0 & 0 & 0 & \omega_{\beta\beta} \end{pmatrix}. \quad (2.2)$$

Here, $\omega_{\lambda\mu} = \langle \lambda\mu | H_{\text{free},I} | \lambda\mu \rangle$ ($\lambda, \mu \in \{\alpha, \beta\}$) represents the precession frequency of the state $|\lambda\mu\rangle$. Given that $I_z |\alpha\rangle = (1/2) |\alpha\rangle$ and $I_z |\beta\rangle = -(1/2) |\beta\rangle$, these frequencies are relatively easy to work out:

$$\begin{aligned} \omega_{\alpha\alpha} &= \frac{1}{2}(\Omega_1 + \Omega_2 + \pi J), & \omega_{\alpha\beta} &= \frac{1}{2}(\Omega_1 - \Omega_2 - \pi J), \\ \omega_{\beta\alpha} &= \frac{1}{2}(-\Omega_1 + \Omega_2 - \pi J), & \omega_{\beta\beta} &= \frac{1}{2}(-\Omega_1 - \Omega_2 + \pi J). \end{aligned} \quad (2.3)$$

The corresponding propagator $U(\tau) = \exp(-iH_{\text{free},I}\tau)$ is then just:

$$U(\tau) = \begin{pmatrix} \exp(-i\omega_{\alpha\alpha}\tau) & 0 & 0 & 0 \\ 0 & \exp(-i\omega_{\alpha\beta}\tau) & 0 & 0 \\ 0 & 0 & \exp(-i\omega_{\beta\alpha}\tau) & 0 \\ 0 & 0 & 0 & \exp(-i\omega_{\beta\beta}\tau) \end{pmatrix}. \quad (2.4)$$

Consequently, the evolution of the single-element operators under $H_{\text{free},I}$ is extraordinarily simple to calculate in matrix form: for example, we have that

$$I_{1+}I_{2\alpha} = \begin{pmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \end{pmatrix}, \quad (2.5)$$

so $U(\tau)I_{1+}I_{2\alpha}U^\dagger(\tau)$ is

$$\begin{pmatrix} 0 & 0 & \exp(-i\omega_{\alpha\alpha}\tau)\exp(i\omega_{\beta\alpha}\tau) & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \end{pmatrix} = \exp[-i(\Omega_1 + \pi J)\tau]I_{1+}I_{2\alpha}. \quad (2.6)$$

Essentially, under $H_{\text{free},I}$, all of these operators acquire phase factors which depend on the difference between two of the frequencies $\omega_{\lambda\mu}$. I explicitly state the rules for the single-quantum operators on spin 1 here:^{*}

$$I_{1+}I_{2\alpha} \longrightarrow \exp[-i(\Omega_1 + \pi J)\tau]I_{1+}I_{2\alpha} \quad (2.7)$$

$$I_{1+}I_{2\beta} \longrightarrow \exp[-i(\Omega_1 - \pi J)\tau]I_{1+}I_{2\beta} \quad (2.8)$$

$$I_{1-}I_{2\alpha} \longrightarrow \exp[i(\Omega_1 + \pi J)\tau]I_{1-}I_{2\alpha} \quad (2.9)$$

$$I_{1-}I_{2\beta} \longrightarrow \exp[i(\Omega_1 - \pi J)\tau]I_{1-}I_{2\beta} \quad (2.10)$$

The rules for the corresponding operators on spin 2 can be easily obtained by permutation of labels. Notice that the evolution frequencies of the -1 -quantum operators (eqs. (2.9) and (2.10)) each correspond to one peak of the corresponding multiplet in an NMR spectrum: for example, $\Omega_1 + \pi J$ and $\Omega_1 - \pi J$ correspond to the two peaks of the spin-1 doublet.

Consider now a simple spin echo sequence: $90^\circ_x - \tau - 180^\circ_x - \tau - \text{detection}$. The initial excitation pulse acts on both spins 1 and 2, and thus generates a mixture of all eight possible single-quantum operators (the four above plus four more on spin 2). For simplicity, we consider only the $I_{1+}I_{2\alpha}$ term. This evolves in the first τ delay to give $\exp[-i(\Omega_1 + \pi J)\tau]I_{1+}I_{2\alpha}$. The 180° pulse *flips* both spins 1 and 2, in that it causes the transitions $I_+ \leftrightarrow I_-$ and $I_\alpha \leftrightarrow I_\beta$; consequently, we have that

$$\exp[-i(\Omega_1 + \pi J)\tau]I_{1+}I_{2\alpha} \longrightarrow \exp[-i(\Omega_1 + \pi J)\tau]I_{1-}I_{2\beta}. \quad (2.11)$$

^{*}Note that § 10.4.2 of Keeler's text³ has a sign error in these equations—the sign of the πJ evolution is flipped—though it proves to be inconsequential as the sign of J cannot be observed.

In the second delay, we get a second phase factor from the evolution of the $I_{1-}I_{2\beta}$ operator:

$$\begin{aligned}\exp[-i(\Omega_1 + \pi J)\tau]I_{1-}I_{2\beta} &\longrightarrow \exp[-i(\Omega_1 + \pi J)\tau] \exp[i(\Omega_1 - \pi J)\tau]I_{1-}I_{2\beta} \\ &= \exp(-2i\pi J\tau)I_{1-}I_{2\beta}.\end{aligned}\quad (2.12)$$

Detection of this gives us one of the two peaks of the spin-1 doublet, as described previously, but with a phase factor tacked on. The Ω_1 terms in the phase factor are cancelled out, which reflects the fact that the offset (or chemical shift) is refocused by the 180° pulse. However, the J -evolution is not refocused, which leads to characteristic phase distortions in the detected multiplets. The same is true of the seven other single-quantum operators.

In order to refocus the J -evolution as well as the chemical shift, we would need—instead of a 180° pulse—a pulse sequence element which simultaneously effects *all* of the following transitions on spin 1:

$$I_{1+}I_{2\alpha} \longrightarrow I_{1-}I_{2\alpha}; \quad I_{1+}I_{2\beta} \longrightarrow I_{1-}I_{2\beta}; \quad I_{1-}I_{2\alpha} \longrightarrow I_{1+}I_{2\alpha}; \quad I_{1-}I_{2\beta} \longrightarrow I_{1+}I_{2\beta}; \quad (2.13)$$

and on spin 2:

$$I_{1\alpha}I_{2+} \longrightarrow I_{1\alpha}I_{2-}; \quad I_{1\beta}I_{2+} \longrightarrow I_{1\beta}I_{2+}; \quad I_{1\alpha}I_{2-} \longrightarrow I_{1\alpha}I_{2+}; \quad I_{1\beta}I_{2-} \longrightarrow I_{1\beta}I_{2-}. \quad (2.14)$$

Such an element forms the basis of a pure shift technique, and I refer to it here as a *pure shift element* (PSE). The difficulty in designing a PSE is that *all* spins must be simultaneously decoupled from each other (and not just one spin). For example, if we only had to invert spin 1 and not spin 2 (i.e. only eq. (2.13) and not eq. (2.14)), this could be trivially accomplished by a selective 180° pulse on spin 1. However, this would not bring about the correct transitions for spin 2. Yet another complicating factor is that the spins will have offsets and couplings which are *a priori* not known; so the design of the PSE cannot use these parameters as inputs. These limitations mean that it is impossible to accomplish the above transitions *in full*; rather, a more realistic scenario involves

$$I_{1+}I_{2\alpha} \longrightarrow cI_{1-}I_{2\alpha} + \sum_i c'_i M_i \quad (2.15)$$

and likewise for the other operators. Here, the desired transition probability $c < 1$ directly correlates with the sensitivity of the PSE, and the M'_i 's are some other undesired operators which (if detectable) lead to artefacts if not suppressed.

In the above discussion, note that the role of the PSE is to invert the I_+ and I_- terms, and to leave the I_α and I_β terms untouched. The spins with I_+ and I_- terms are referred to as *active spins*, and the I_α and I_β spins as *passive spins*. Thus, for example, in the context of eq. (2.13), spin 1 is active and spin 2 is passive. The detected signal always arises from the active spins (unless coherence transfer between spins occurs, for example in the mixing period of a 2D pure shift

experiment).

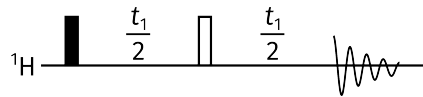


Figure 2.1: Pulse sequence for the J-resolved (2DJ) experiment.

Before moving on to the discussion of how PSEs are implemented in practice, I insert a slight digression about *J-resolved* (or *2DJ*) *spectroscopy*, which is very closely related to pure shift NMR. The basic 2DJ sequence involves a spin echo of duration t_1 , immediately followed by detection (fig. 2.1). If we only consider a single operator and reuse the analysis above, setting the spin echo delay τ to be $t_1/2$ instead, then we have that

$$I_{1+}I_{2\alpha} \xrightarrow{\frac{t_1}{2}} \xrightarrow{180^\circ} \xrightarrow{\frac{t_1}{2}} \exp(-i\pi J t_1) I_{1-}I_{2\beta}. \quad (2.16)$$

This yields a complex signal of the form

$$s(t_1, t_2) = \exp(-i\pi J t_1) \exp[i(\Omega_1 - \pi J)t_2], \quad (2.17)$$

which when Fourier transformed yields a phase twist lineshape at $(-\pi J, \Omega_1 - \pi J)$. The other component on spin 1 (starting from $I_{1+}I_{2\beta}$) likewise yields a phase twist at $(\pi J, \Omega_1 + \pi J)$. It has long been known that *shearing* this 2DJ spectrum by 45° (i.e. moving each data point (Ω_1, Ω_2) to $(\Omega_1, \Omega_2 - \Omega_1)$) generates a spectrum which only has chemical shift information in the ω_2 dimension. After magnitude-mode processing, projection of this spectrum onto the ω_2 axis, for example, would in principle yield a pure shift spectrum.* This is true, but in practice the phase twist lineshapes cause the resulting resolution to be very poor, which defeats the purpose of using a pure shift spectrum. To circumvent this issue, a number of special processing techniques have been proposed^{7–9} (see also references therein); but more ideally, we want a phase-sensitive 2DJ spectrum, which can be obtained by recording a pair of ‘echo’ and ‘antiecho’ signals:[†]

$$s_{\text{echo}}(t_1, t_2) = \exp(i\pi J t_1) \exp[i(\Omega_1 - \pi J)t_2] \quad (2.18)$$

$$s_{\text{antiecho}}(t_1, t_2) = \exp(-i\pi J t_1) \exp[i(\Omega_1 - \pi J)t_2] \quad (2.19)$$

These can be processed in the same way as described in § 1.4.3 to yield double absorption-mode lineshapes. The antiecho signal (eq. (2.19)) is of course the same as in eq. (2.17), but to obtain the echo signal in eq. (2.18), we require a different pulse sequence with a PSE inserted just prior

*Or equivalently, projection of the unsheared spectrum along a 45° axis.⁶

[†]The terms ‘echo’ and ‘antiecho’ refer to the relative senses of the coherences evolving during t_1 and t_2 : in the echo spectrum these have opposite signs, e.g. $I_z S_+$ and I_- in the HSQC, and in the antiecho spectrum they have the same sign. As pointed out by Pell and Keeler,¹⁰ this is not really appropriate for the 2DJ experiment since each half of t_1 has a coherence with a different sense, but we will stick to this nomenclature as the underlying concept is very similar to that of echo–antiecho processing.

to detection:

$$I_{1-}I_{2\alpha} \xrightarrow{t_1/2} \exp[i(\Omega_1 + \pi J)t_1/2]I_{1-}I_{2\alpha} \quad (2.20)$$

$$\xrightarrow{180^\circ} \exp[i(\Omega_1 + \pi J)t_1/2]I_{1+}I_{2\beta} \quad (2.21)$$

$$\xrightarrow{t_1/2} \exp(i\pi J t_1)I_{1+}I_{2\beta} \quad (2.22)$$

$$\xrightarrow{\text{PSE}} c \exp(i\pi J t_1)I_{1-}I_{2\beta} \quad (2.23)$$

$$\xrightarrow{t_2} c \exp(i\pi J t_1) \exp[i(\Omega_1 - \pi J)t_2]. \quad (2.24)$$

(Note that we began the analysis with a different operator here, $I_{1-}I_{2\alpha}$, in order to end up with the same $I_{1-}I_{2\beta}$ operator just before detection.) In order to apply echo–antiecho processing, the decrease in sensitivity by a factor of c must also be applied to the antiecho spectrum: this can be done by simply inserting the PSE immediately after the initial excitation pulse, which simply scales all operators down by c . Thus, we see that *exactly the same PSE* allows us to generate pure shift spectra as well as absorption-mode 2DJ spectra: this has been previously demonstrated using various PSEs.^{10,11}

In fact, the same formalism can be used to describe a family of small flip angle COSY experiments, including ECOSY^{5,12,13} and z -COSY;^{14–16} these are also closely related to pure shift NMR. In particular, the anti z -COSY experiment is a precursor to PSYCHE, and is analysed in § 2.2.3. However, a full discussion of these is beyond the scope of this thesis.

2.2 Pure shift in practice

In the previous section, I described the underlying theory used for analysing PSEs and showed how such an element could be used to record absorption-mode 2DJ spectra. From this, one can obtain a pure shift spectrum through shearing and projection. However, this is only an *indirect* route to a pure shift spectrum. In this section, we will tackle the main question of how pure shift experiments may be *directly* acquired using a PSE. Following this, I cover several examples of PSEs reported in the literature. This is not an exhaustive survey of pure shift methods: I only choose to cover a handful of PSEs which specifically accomplish the transformations listed in eqs. (2.13) and (2.14). Thus, for example, constant-time techniques (which are widely used to suppress ^{13}C – ^{13}C couplings in labelled biomolecules) are not mentioned.

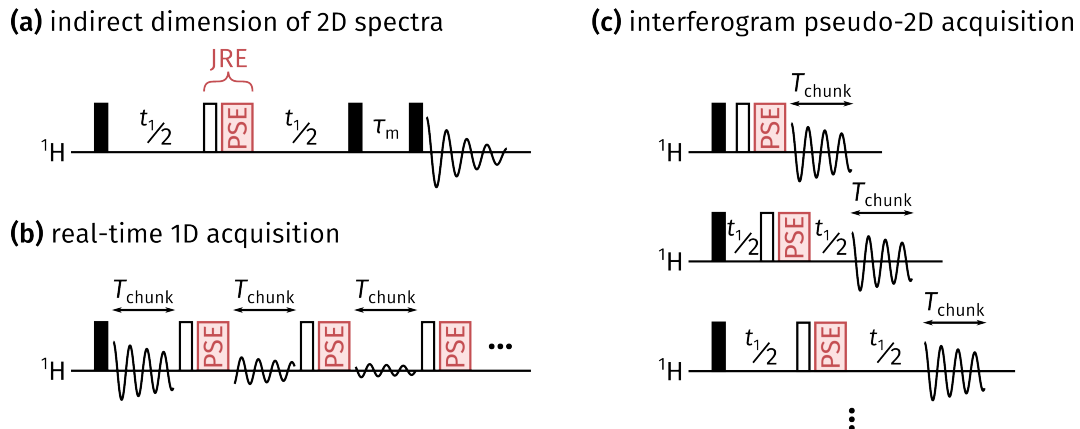


Figure 2.2: Possible acquisition modes for pure shift spectroscopy. The red box labelled ‘PSE’ indicates a generic pure shift element, which can be any of those described in the main text. In practice, gradients are also used to suppress unwanted coherence transfers; these are not shown here for simplicity. (a) Insertion of a J-refocusing element (JRE) in the centre of an indirect-dimension evolution period, which leads to a spectrum which is pure shift in F_1 . The $90^\circ-\tau_m-90^\circ$ block shown here corresponds to a NOESY experiment, but in principle any 2D experiment can be adapted in this fashion. (b) Real-time acquisition of a 1D pure shift spectrum in chunks of duration T_{chunk} . (c) Interferogram acquisition of a 1D pure shift spectrum, where t_1 is lengthened by T_{chunk} every increment.

2.2.1 Acquisition modes

Restating eq. (2.15), suppose we have a PSE which accomplishes the transformation

$$I_{1+} I_{2\alpha} \longrightarrow c I_{1-} I_{2\alpha} + \sum_i c'_i M_i \quad (2.25)$$

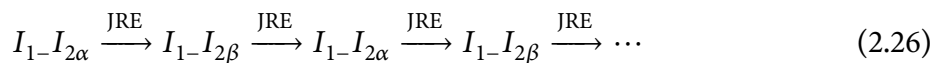
(and likewise for the other single-quantum operators, which are not shown here). The simplest method of using this is to insert it in the middle of a t_1 period of a 2D experiment. This is actually not entirely desirable, because the PSE causes *both* chemical shifts and J-couplings to be refocused; consequently, there will be *no* frequency modulation during t_1 at all! It is more sensible to combine the PSE with a hard 180° pulse, which refocuses only chemical shifts. Together, the effect is to refocus J-couplings and allow chemical shifts to evolve; this combination is thus called a *J-refocusing element*, or JRE (fig. 2.2a). We can equivalently say that the JRE flips all passive spins and leaves active spins untouched. This distinction between a JRE and a PSE is important and will be referred to several times in this chapter.

Implementing a JRE in the middle of a t_1 period is simple, requiring minimal modification of existing 2D experiments. Furthermore, the pure shift ‘character’ of the F_1 dimension may then be mapped to the F_2 dimension through indirect covariance processing.^{17–22} However, the increased resolution in the F_1 dimension provided by homodecoupling cannot really be reaped unless many t_1 increments are acquired. Furthermore, this does not help with acquiring a 1D pure shift

spectrum, where there is no indirect dimension.

This brings us to the second acquisition mode for pure shift data, called *real-time acquisition*.^{23–26} Here, JREs are inserted at regular intervals throughout an acquisition period, causing the chemical shift evolution to be effectively ‘suspended’ for the duration of the JRE, and the sense of J-evolution to be reversed (fig. 2.2b).^{*} This leads to a series of FID ‘chunks’ which must then be concatenated to form the desired FID. To prevent the J-coupling from evolving too much during a single chunk, the required spacing of the JREs, or equivalently the duration of each chunk T_{chunk} , must satisfy $T_{\text{chunk}} \ll 1/J$ (in practice, it is on the order of $1/(2J)$). Naturally, real-time acquisition still comes with the sensitivity penalty of c . However, it allows a pure shift spectrum to be acquired in effectively the same time as the original coupled spectrum; its ‘single-scan’ nature also allows, for example, the application of hyperpolarisation techniques which cannot be reproducibly repeated over multiple increments.^{27,28}

Unfortunately, it is not always possible to perform real-time acquisition. The reason is because the JRE is applied multiple times, and each time it is, it must select for the same active and passive spins in the same molecule as it did the last time. In other words, *for any given molecule in the sample*, it must enforce this CTP:



As will be described later, the BIRD and Zangerer–Sterk methods always select the same active spins in the same molecules, but the PSYCHE method does not. Therefore, in order to acquire pure shift PSYCHE spectra, we have to resort to the *interferogram method*, where each chunk is obtained as a separate increment of a 2D experiment (fig. 2.2c). The insertion of the JRE in the middle of the t_1 period means that when detection is started, it is ‘as if’ only the chemical shift has evolved for a period t_1 . On each increment, one chunk—again of duration $T_{\text{chunk}} \ll 1/J$ —is detected, and then t_1 is incremented by T_{chunk} so that the next chunk can be recorded. Finally, the chunks are stitched together to form the requisite FID.[†] Since the indirect dimension is not processed by a Fourier transform, this is sometimes called a *pseudo-2D* experiment (or *pseudo-3D* if this is applied to the direct dimension of a 2D experiment, and so on). In this case, the sensitivity drop c is incurred, but there is on top of that also a time penalty in that the experiment duration must be lengthened by n times in order to collect n chunks. n is typically

^{*}Relaxation during the JRE must also be taken into account for the real-time method: this causes each successive chunk to decay in intensity faster than usual, thereby leading to peak broadening, which can be an issue for very long JREs. In contrast, the interferogram method only applies one JRE per increment, so the relaxation losses in each chunk are just a constant.

[†]Not included in figs. 2.2b and 2.2c is the extra detail that scalar couplings are usually allowed to evolve for a period of $T_{\text{chunk}}/2$ at the start of the sequence by the addition of a spin echo: this amounts to a *prefocusing* of J-evolution, such that J-coupling is refocused in the middle of the chunk rather than the beginning.²⁹ This allows chunk sizes twice as large to be used, reducing the total duration of the experiment. This J-prefocusing can also be done in a more intelligent manner via the SAPPHIRE method,³⁰ which is discussed in more detail in § 4.3.6.

on the order of 16–32.

2.2.2 Pure shift elements

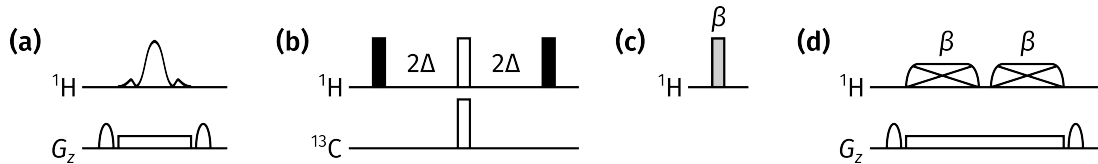


Figure 2.3: A selection of pure shift elements. (a) Zanger–Sterk PSE,³¹ involving the combination of a selective 180° refocusing pulse and a weak gradient. (b) BIRD PSE;^{32,33} the delay Δ is set to $1/(4 \cdot ^1J_{\text{CH}})$. (c) Time-reversal PSE,¹³ simply consisting of a hard pulse with variable flip angle β . Multiple spectra with different values of β must be co-added to suppress artefacts (though this suppression is not perfect, as discussed in the text). (d) PSYCHE PSE,³⁴ consisting of two saltire pulses^{35,36} with flip angle β , and a weak gradient.

Zanger–Sterk

We are finally now in a position to study individual PSEs and their mechanisms of action. We begin with the Zanger–Sterk (ZS, or ‘slice-selective’) PSE,³¹ in which a selective refocusing pulse and a weak gradient are simultaneously applied (fig. 2.3a). In practice, an rSNOB pulse³⁷ is often used as the refocusing pulse. The effect of the gradient is to make each spin in the sample have a spatially dependent offset; therefore, in each *slice* (or cross-section) of the sample, a different spin will fall within the specific bandwidth of the refocusing pulse. This spin is refocused by the PSE and therefore becomes the active spin *within that specific slice*; the bracketing pair of CTP gradients serve to destroy coherences on all the other spins which are not inverted. Each signal of the pure shift spectrum therefore derives from a specific slice of the sample; during direct detection, all slices simultaneously contribute to the signal, thus yielding a broadband pure shift spectrum.

The sensitivity of the ZS method tends to be low (the factor c tends to be on the order of 0.01 to 0.05), as each signal only comes from a narrow section of the sample. Nevertheless, it still finds wide usage in pure shift applications nowadays, especially because it is compatible with the real-time acquisition mode:²⁴ as long as the pulse and the weak gradient are the same each time, then the same active spins will always be chosen in the same slice (as long as diffusion effects are ignored). The PSE can also be customised through the bandwidth of the refocusing pulse: decreasing this improves the spatial differentiation between spins which have similar intrinsic offsets, yielding better decoupling quality, albeit at the cost of sensitivity. The ZS element can be easily—and has been—adapted for use in many experiments, including (but not limited to) absorption-mode 2DJ spectroscopy¹⁰ and selective refocusing (SERF) experiments for the measurement of ${}^nJ_{\text{HH}}$.^{38–41}

BIRD

Next up is the *bilinear rotation decoupling* (BIRD) pulse element (fig. 2.3b). BIRD is not spatially selective like the ZS method; instead, it is *isotope-selective* in that it acts as a 180°_y pulse on ^{13}C -bound protons, and does not affect ^{12}C -bound protons. Consequently, all ^{13}C -bound protons become the active spins in the context of pure shift NMR. The first report of the BIRD element,³² in 1982, was clearly ahead of its time: it reported the use of an interferogram-type approach to obtain 1D pure shift spectra. However, in the subsequent decades, this seemed to have been forgotten: BIRD found much more use as an isotope-selective rotation element in heteronuclear NMR,⁴² until its use as a pure shift element was ‘rediscovered’.^{33,43}

An immediate drawback of BIRD is that it does not decouple geminal (diastereotopic) CH_2 groups, as both protons would be either both active or both passive. The sensitivity penalty of BIRD is also relatively severe: the factor c derives from the natural abundance of ^{13}C , which is approximately 0.011. However, it is also compatible with real-time acquisition,²³ and has found particular success as a pure shift element in the F_2 dimension of HMQC and HSQC experiments.^{26,43–47} In this case, the use of BIRD leads to no loss of sensitivity as only ^{13}C -bound protons are detected in HSQC experiments anyway.* It should be noted that the BIRD element does not need to be combined with a hard 180° pulse to form a JRE: the inverse effect of only flipping the passive spins can be accomplished by simply changing the phase of both internal 180° pulses to $+y$. Using the nomenclature of Uhrín et al.,⁴² the PSE and JRE forms of the BIRD pulse can be labelled as $\text{BIRD}^{\text{d},\text{X}}$ and $\text{BIRD}^{\text{r},\text{X}}$ respectively.

Time-reversal

The *time-reversal* PSE (fig. 2.3c) is even simpler: it consists only of one hard pulse with flip angle β .¹³ The catch is that the experiment must be repeated with different values of β , and the results added up with specific weightings.⁵ This leads to cancellation of *some*, but not all, of the unwanted coherences: for example, on its own, it does not suppress COSY-type coherence transfer such as $I_{1-}I_{2\alpha} \rightarrow I_{1\alpha}I_{2+}$. This proved to be inconsequential in the original application of F_1 decoupling in a 2D NOESY;¹³ however, it is not acceptable in a 1D context as it leads to substantial artefacts. This will be discussed in more detail in § 2.5, so a fuller analysis of the time-reversal PSE is deferred until then.

PSYCHE

Finally, we come to the PSYCHE method, which is generally considered the current state of the art for pure shift NMR.³⁴ The corresponding PSE (fig. 2.3d) consists of two swept-frequency pulses

*In fact, the sensitivity is increased by the collapse of multiplet structure.

applied during a weak gradient: for example, a pair of chirps with opposite sweep directions can be used. The signal-to-artefact ratio of PSYCHE can be further improved by using two *saltire pulses*: these are superpositions of chirps which simultaneously sweep in both frequency directions.³⁵ The operation of this PSE is not easy to fully explain.³⁶ However, we can adopt the (not fully accurate, but still useful) instant-flip approximation^{48,49} —that the swept-frequency pulse acts as an instantaneous 180° rotation on each frequency it passes through. Using this, the PSYCHE element (or strictly, the PSYCHE JRE, including a hard 180° pulse) may be viewed as a spatially parallelised version of the anti *z*-COSY experiment,^{14,15} which we now describe.

2.2.3 PSYCHE in detail

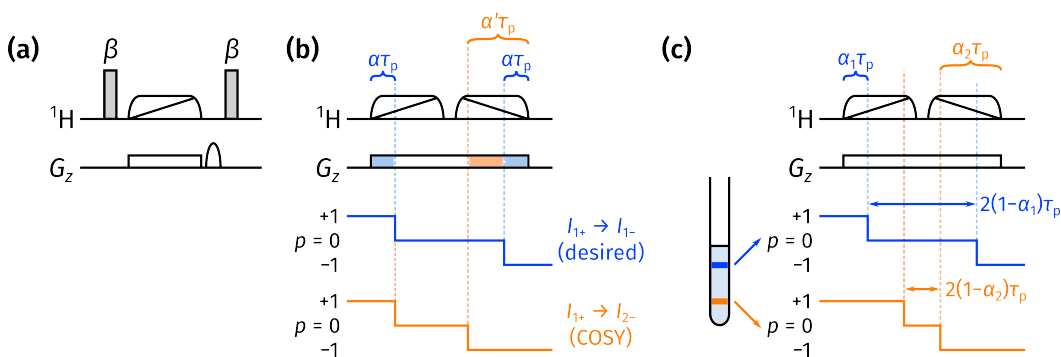


Figure 2.4: A closer look at the mechanism of the PSYCHE PSE. (a) The β -ZQF- β mixing period used in the original anti *z*-COSY experiment. This has a similar action to a JRE, but does not suppress COSY-type coherence transfers from spin *i* to spin *j*. (b) An illustration of how COSY-type artefacts are suppressed by the PSYCHE pulse element. The desired CTP which remains on spin 1 is rephased by the gradients, but the COSY CTP is dephased. (c) An illustration of how zero-quantum terms are suppressed by the PSYCHE element through spatial averaging: in each slice of the sample (highlighted in blue and orange), the zero-quantum terms are allowed to evolve for a different duration.

The anti *z*-COSY experiment utilises a β -zero-quantum filter (ZQF)- β mixing period (fig. 2.4a), where β is a small angle, typically 10° to 20°. The role of the ZQF⁵⁰ is to remove terms such as $I_{1-}I_{2+}$ between the two β pulses, retaining only population terms such as $I_{1\alpha}I_{2\alpha}$. The CTPs which this mixing period selects for ultimately give rise to peaks which lie perpendicular to the main diagonal of the spectrum. In an elegant paper, Pell et al.¹⁵ showed that by isolating the diagonal peaks of a 2D anti *z*-COSY experiment and taking the 45° projection of these, a pure shift spectrum could be obtained. Here, we will go one step further and consider the *direct* use of this element as a JRE: this will reveal some problems which are neatly taken care of by the PSYCHE experiment.

We first need to introduce how the basis operators $\{I_+, I_-, I_\alpha, I_\beta\}$ evolve under a hard pulse

(applied along the x -axis) with flip angle β :

$$I_{\pm} \xrightarrow{\beta I_x} c^2 I_{\pm} + s^2 I_{\mp} \pm \frac{iS}{2}(I_{\alpha} - I_{\beta}) \quad (2.27)$$

$$I_{\alpha} \xrightarrow{\beta I_x} c^2 I_{\alpha} + s^2 I_{\beta} + \frac{iS}{2}(I_{+} - I_{-}) \quad (2.28)$$

$$I_{\beta} \xrightarrow{\beta I_x} c^2 I_{\beta} + s^2 I_{\alpha} - \frac{iS}{2}(I_{+} - I_{-}) \quad (2.29)$$

where $S = \sin \beta$, $s = \sin(\beta/2)$, and $c = \cos(\beta/2)$. Using these formulae, we can show that for a two-spin system (see Pell et al.¹⁵ for the equivalent analysis on a three-spin system), the β -ZQF- β element converts the term $I_{1+}I_{2\alpha}$ to

$$I_{1+}I_{2\alpha} \longrightarrow \underbrace{\frac{1}{2}S^2c^4I_{1+}I_{2\beta}}_{\text{term 1}} + \underbrace{S^2c^2s^2I_{1+}I_{2\alpha}}_{\text{term 2}} - \underbrace{\frac{1}{4}S^2c^4I_{1\alpha}I_{2+}}_{\text{terms 3 and 4}} + \frac{1}{4}S^2c^4I_{1\beta}I_{2+} + \dots, \quad (2.30)$$

where other terms with different coherence orders have been neglected (on the basis that they can be easily suppressed with bracketing CTP gradients), and terms with higher orders in s have been discarded since $s = \sin(\beta/2) \ll 1$ for small β .

The first term $I_{1+}I_{2\beta}$, corresponding to the flipping of passive spins only, is the only term we want to see from a JRE. The second term, $I_{1+}I_{2\alpha}$, corresponds to the case where neither active nor passive spins have been flipped. In the original anti z -COSY work, these give rise to ‘off-diagonal’ peaks which are part of a multiplet on the diagonal, but when projected at 45° generate artefacts around the pure shift peak. In the context of pure shift NMR, these are called ‘recoupling artefacts’, as they arise from imperfect J -refocusing. Note that the ratio of recoupling artefacts to desired signal is proportional to $S^2c^2s^2/S^2c^4 = \tan^2(\beta/2)$: using a smaller value for β therefore leads to better signal-to-artefact ratios, but also lower overall sensitivity. The PSYCHE element is similar to the β -ZQF- β element in this regard: it does not suppress the recoupling artefacts, but instead relies on the user choosing a suitable value for β such that the artefact-to-signal ratio is tolerably small. If the sensitivity proves to be insufficient, the flip angle β may be increased instead: this leads to a larger artefact-to-signal ratio, but if the sample is not concentrated anyway, it may well be that the artefacts do not rise above the noise level.

The third and fourth terms, $I_{1\lambda}I_{2+}$ ($\lambda \in \{\alpha, \beta\}$), represent ‘COSY-type’ coherence transfer to a coupled spin. In the original anti z -COSY, these led to crosspeak multiplets at (Ω_1, Ω_2) , which could be removed by hand before taking the projection. However, in a pure shift sequence, the peaks arising from these terms cannot simply be removed in the same way. It is precisely this issue which precludes the β -ZQF- β element from being directly used as a JRE, and motivates the development of the PSYCHE PSE, which *does* suppress these coherence transfers.

To understand how this occurs, we invoke the instantaneous spin-flip assumption. Each coher-

ence I_{i+} is converted (or ‘flipped’) to a population term $I_{1\lambda}$ at a specific point $\alpha\tau_p$ after the start of the first chirp, and can be reconverted to a coherence on the same spin I_{i-} at a time $\alpha\tau_p$ before the end of the second chirp (the blue CTP in fig. 2.4b).^{*} Here, τ_p is the duration of the chirp, and $0 < \alpha < 1$. In this case, the coherence is perfectly rephased by the weak gradient applied during the chirp pulses, since the *time* it experiences these gradients for is the same on both sides of the chirps. Now, if the I_{i+} term is instead converted to a coherence on a different spin I_{j-} , it experiences the gradient for a total duration of $\alpha\tau_p$ after the start of the first chirp, and also $\alpha'\tau_p$ before the end of the second chirp (the orange CTP in fig. 2.4b). In general, since spins i and j have different offsets, we have that $\alpha \neq \alpha'$. Therefore, this CTP is dephased by the gradients, resulting in suppression of the COSY-type artefacts in the spectrum.

It remains to also consider how the PSYCHE element selects for the population terms between the two spin flips. Any terms with nonzero coherence order are of course simply dephased by the weak gradient. However, zero-quantum terms (in homonuclear systems) are not dephased by gradients, and to eliminate them in a single-scan manner, they must be spatially averaged, for example by a ZQF. It turns out that the PSYCHE element also results in a similar spatial averaging. Following on from the previous paragraph, the time *between* the spin flips (for the desired pathways, i.e. not COSY-type coherence transfer) is given by $2(1 - \alpha)\tau_p$. At the same time, the weak gradient induces a range of offsets across the sample, much like in the Zanger-Sterk experiment. Thus, the offset, and thus the value of α , for a given spin depends on which slice it is in; for example, fig. 2.4c uses values of α_1 and α_2 for two different slices (blue and orange). If zero-quantum terms are present between the spin flips, they evolve during this time and accrue a spatially-dependent phase: summation of these during FID acquisition leads to a cancellation of these terms. Only population terms such as $I_{1\alpha}I_{2\alpha}$ survive during this, as they do not precess during this time.

The sensitivity of PSYCHE is significantly better than for other methods: depending on the flip angle β chosen, c is typically on the order of 0.05–0.15 (see also fig. 2.5 for explicit simulations). Furthermore, it is generally more robust with respect to strong coupling compared to other pure shift methods (artefacts from strong coupling often arise due to unexpected coherence transfer,⁴ which is suppressed in a similar way to the COSY-type artefacts). These two factors alone have meant that PSYCHE has enjoyed substantial adoption since its introduction: a large number of 2D experiments utilising PSYCHE decoupling in either F_1 or F_2 have been developed,^{22,51–56} notably the PSYCHE-iDOSY diffusion experiment,⁵⁷ where the increased resolution provided by pure shift spectroscopy translates into increased resolution in the *diffusion* dimension as well. Like the ZS element before it, the PSYCHE element has also been used for the acquisition of absorption-mode 2DJ spectra.¹¹

^{*}Note the change in the sign of the coherence, which differs from the analysis of the anti *z*-COSY experiment. This arises because we are only considering the PSYCHE PSE on its own, *not* the JRE.

Despite this success, PSYCHE suffers from one significant drawback: it cannot be used in a real-time fashion. The PSYCHE PSE is often said to select active and passive spins in a ‘statistical’ manner: this is because of the c^2 and s^2 terms arising from the low-flip angle pulses. What this really means is that we do not care *exactly* which spins are active and which are passive, but that a certain proportion of the spins are active and passive. Repeated application of the PSE therefore does not select for the same active spins each time, which precludes its application to real-time acquisition.

Although the PSYCHE PSE may appear deceptively simple at first glance, the closer analysis given here (and elsewhere³⁵) clearly shows that its inner workings are anything but. Along with other ingenious experiments such as the ZQF,⁴ ultrafast NMR,^{58–60} and more recently GEMSTONE,⁶¹ PSYCHE is a prime example of how *spatiotemporal averaging* and pulsed field gradients can be used to great effect in modern NMR spectroscopy.⁶²

At the same time, PSYCHE itself is not *perfect*: it does not fully suppress recoupling artefacts, and can only be used in the interferogram mode. To improve on PSYCHE would therefore entail one of the following:

1. increasing the sensitivity (while maintaining purity);
2. increasing the purity (while maintaining sensitivity); or
3. developing a pure shift element which is compatible with real-time acquisition while giving comparable sensitivity and purity to PSYCHE.

The sections which follow describe my efforts towards objectives (1) and (2).

2.3 PSYCHE with a variable number of saltires

The first attempted method was to change the number of saltire pulses used in the PSYCHE PSE. As described in § 2.2.3, PSYCHE relies on spatiotemporal averaging to suppress unwanted artefacts: this crucially relies on the fact that the pulse(s) used in the PSE are symmetric. This can be accomplished with two opposing chirps, or two saltires, both of which are symmetric. However, a *single* saltire is also symmetric in itself: it is not hard to show that a single saltire can provide the requisite averaging. Likewise, the use of four saltires would also be valid.

Theoretical simulations show that the overall profile of signal and artefact versus flip angle varies with the number of saltires (fig. 2.5). Generally, using a larger number of saltires but with a smaller flip angle accomplishes a similar sensitivity level. This may be qualitatively rationalised as more saltires providing more possible CTPs which generate the desired signal (the same idea is generally invoked when discussing the difference between unidirectional chirps and saltires³⁵).

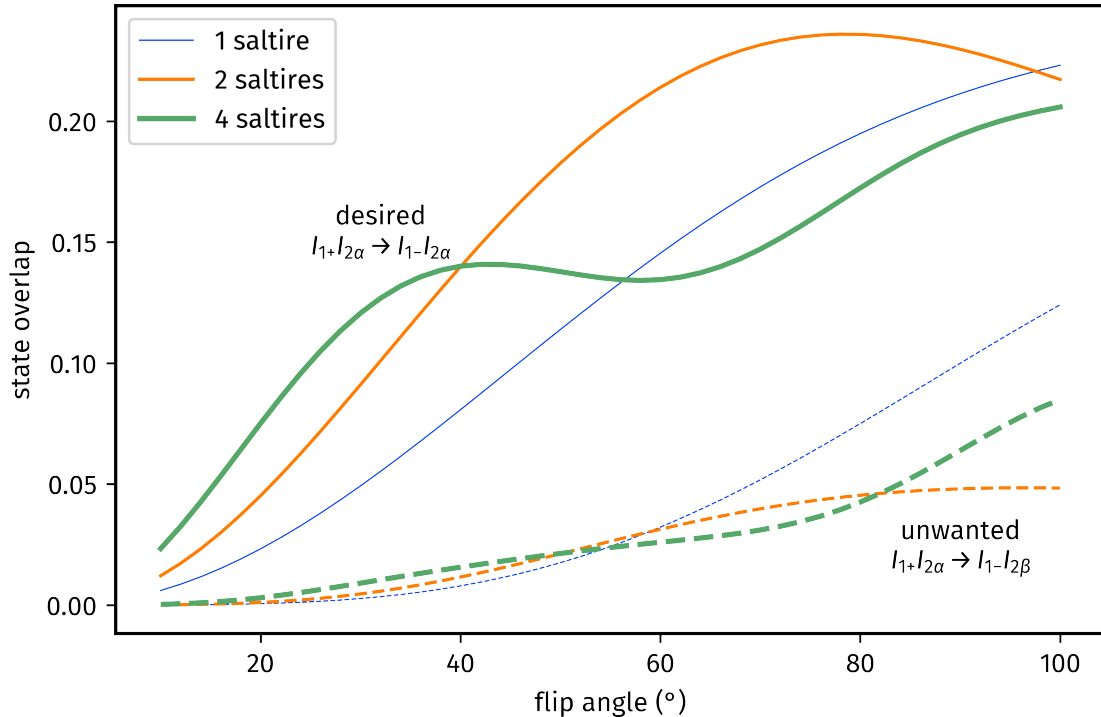


Figure 2.5: Simulated signal and artefact intensity for various PSYCHE-style PSEs as a function of the flip angle used. Calculations were performed on a two-spin system with a coupling of 7 Hz, and an offset difference of 1.5 kHz. The total PSE duration was 30 ms (so, for example, in the four-saltire PSYCHE, each saltire was 7.5 ms). Solid lines indicate the coefficients for the desired $I_{1+}I_{2\alpha} \rightarrow I_{1-}I_{2\alpha}$ pathway which contributes to the pure shift signal; dashed lines the coefficients for the undesired $I_{1+}I_{2\alpha} \rightarrow I_{1-}I_{2\beta}$ pathway, which gives rise to recoupling artefacts.

However, regardless of the number of saltires, the fundamental strategy of adjusting the flip angle to control the signal-to-artefact ratio remains valid, which naturally raises the question of whether specific waveforms and flip angles can be chosen in order to obtain the best decoupling quality.

Quadruple-saltire PSYCHE

The quadruple-saltire PSYCHE was first evaluated experimentally. In the first instance, I set the total duration of the PSE to 30 ms, meaning that each saltire was 7.5 ms long. In this experiment, the sensitivity was defined as the maximum height of the main peak in fig. 2.6a, and the artefact as the mean of the maximum heights of the two artefacts surrounding it. The plot in fig. 2.6b shows how these quantities vary as a function of the flip angle. Interestingly, for the artefacts, the profile observed is similar to that in the simulations: the double-saltire version performs better at low and very high flip angles, but in the middle there is a region where the quadruple-saltire version has lower artefact intensity. The signal intensities for both the double- and quadruple-saltire versions, however, seem to plateau off rather more quickly than the simulations suggest.

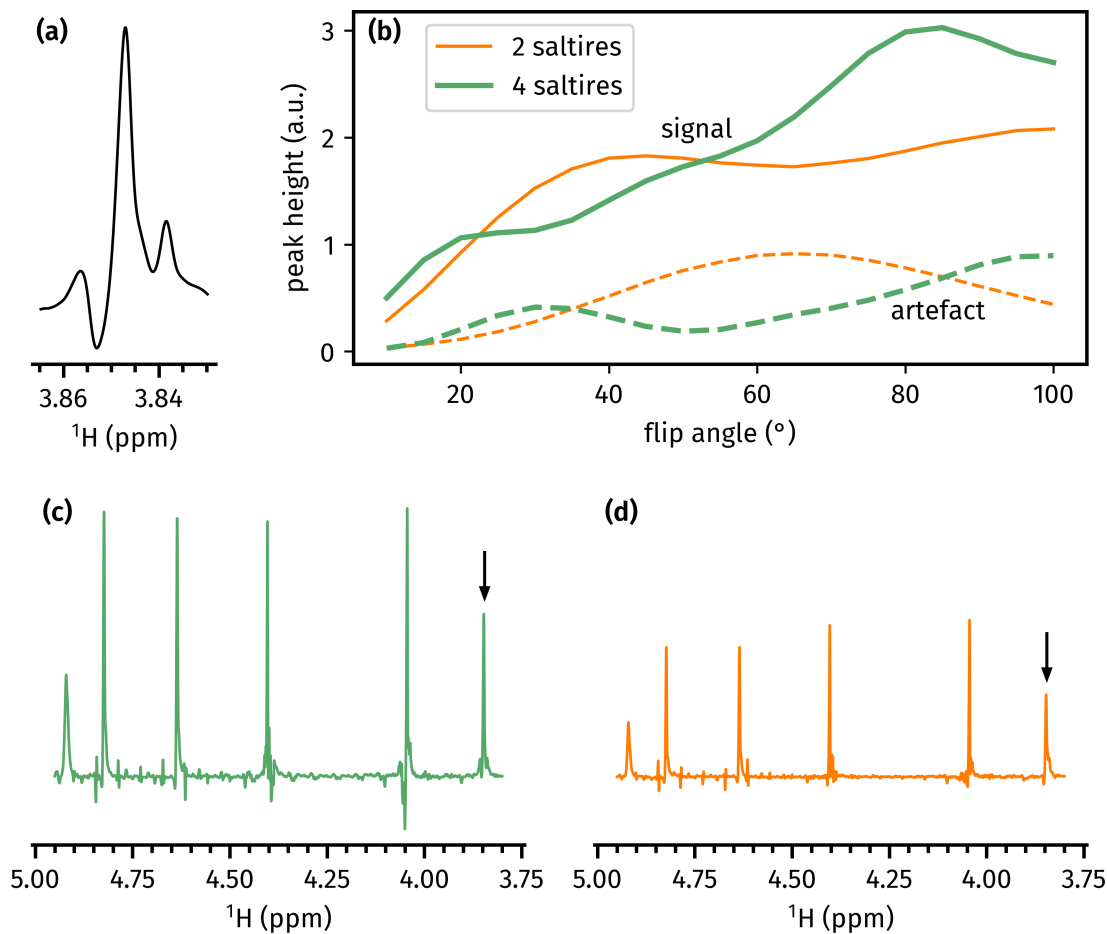


Figure 2.6: (a) The peak in the pure shift spectrum of andrographolide used for calculation of signal and artefact intensity. The recoupling artefacts flanking the main peak are clearly visible. (b) Peak heights of the desired signal (the central peak, solid lines) and artefacts (the mean of the two flanking peaks, dashed lines), as a function of flip angle. (c) Quadruple-saltire PSYCHE with $\beta = 55^\circ$. (d) The reference spectrum, a double-saltire PSYCHE with $\beta = 20^\circ$. The peak at 3.85 ppm used for the sensitivity and purity analysis is labelled with an arrow. *Data code:* 7A-201016.

To highlight one particular data point, fig. 2.6b suggests that the quadruple-saltire experiment with $\beta \approx 55^\circ$ has a similar artefact level to the double-saltire experiment with $\beta \approx 20^\circ$, but with substantially greater signal intensity. Insets from these two spectra are respectively shown in figs. 2.6c and 2.6d. This conclusion does indeed seem to be true for the specific peak used for this analysis, which is at the right edge of the spectral insets shown here. Overall, the quadruple-saltire 55° experiment does have better sensitivity than the double-saltire 20° experiment. However, the artefacts are not always suppressed as nicely as in the chosen reference peak: for example, the peak at 4.05 ppm is significantly less clean in the quadruple-saltire experiment. Any apparent advantage over the double-saltire experiment is therefore not very clear—at least from this data

alone.*

Single-saltire PSYCHE

Moving on to the single-saltire case, in addition to searching for a better signal and artefact profile as before, another possible motivation would be that the duration of the PSE can be decreased. This would minimise losses due to relaxation and diffusion during the PSE, which were not taken into account in fig. 2.5 (and the simulations there did not vary τ_p anyway). A single-saltire PSYCHE experiment was thus recorded with various combinations of flip angle ($\beta/\circ \in \{15, 20, 25, 28, 30\}$) and pulse duration ($\tau_p/\text{ms} \in \{15, 25, 30, 35, 45, 55\}$). In a similar way to the quadruple-saltire study, the sensitivity was defined as the maximum height of the main peak in fig. 2.7a,[†] and the artefact as the mean of the heights of the two artefacts surrounding it. (However, note that the sample used was different.) The purity, or signal-to-artefact ratio, is simply the former divided by the latter.

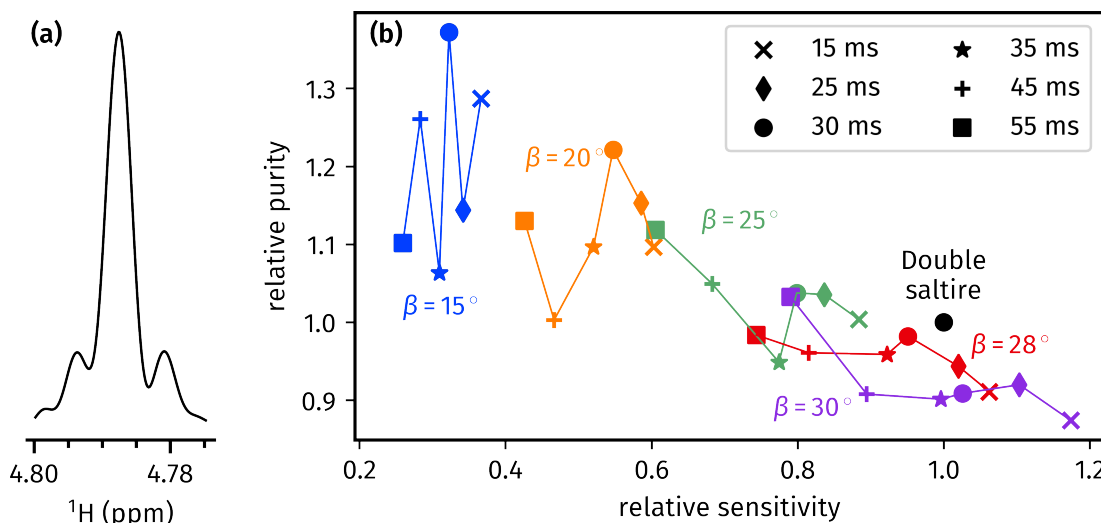


Figure 2.7: (a) The peak in the (reference, double-saltire) pure shift spectrum of cyclosporin used for calculation of signal intensity and signal-to-artefact ratio. (b) Plot showing the signal intensity and signal-to-artefact ratio obtained in various single-saltire PSYCHE experiments, normalised against the double saltire experiment with $\beta = 20^\circ$ and $\tau_p = 30$ ms. Each line represents a series of single-saltire experiments acquired with the same value of β ; τ_p generally decreases going left to right (i.e. a longer pulse means less signal). *Data code:* 5C-190617.

*In fact, I also performed some preliminary experiments where the four-saltire PSE was lengthened to 60 ms. The artefact behaviour in these spectra were better than in their 30 ms counterparts, which is to be expected since a longer PSE leads to better spatiotemporal averaging. However, I unfortunately did not compare these against a 60 ms double-saltire experiment, so these results are not included in this thesis. (It would be rather unfair to compare them against the 30 ms double-saltire.)

[†]As an alternative, the Bruker `sinocal` AU programme was also used to measure the sensitivity of the spectrum; it yielded extremely similar results, so is not shown here. I found the `sinocal` routine to be rather unrepeatable as the exact value calculated depends on random noise.

The results thus obtained are shown in fig. 2.7b. In this plot, both the sensitivities as well as the signal-to-artefact ratios are normalised with respect to the reference double-saltire experiment (black dot at $(1, 1)$, acquired using $\beta = 20^\circ$ and a total $\tau_p = 30$ ms, i.e. 15 ms per saltire). An ideal PSE would fall in the top-right corner of this plot. Clearly, as the flip angle increases, the sensitivity increases but at the cost of the purity: this is hardly unsurprising given that the double-saltire experiment has the same property. The effect of the pulse duration is less clear-cut: in general, a shorter PSE yields to increased signal, and very usable results were obtained even with a simple 15 ms single-saltire PSE. However, a shorter PSE also leads to poorer spatiotemporal suppression of artefacts and thus lower purity.

While there is no standout candidate which is *clearly* better than the double saltire (i.e. greater sensitivity as well as purity), the single-saltire pulse with $\beta = 28^\circ$ and $\tau_p = 30$ ms comes close in performance to the double-saltire experiment. (The use of $\beta = 28^\circ$ here is not coincidental: this flip angle for a single saltire was chosen to (approximately) match the sensitivity of the $\beta = 20^\circ$ double-saltire experiment.) Therefore, this does at least prove that a single saltire can be used as a PSE. However, both the single- and quadruple-saltire cases suffer from the classic dilemma of pure shift NMR: just as in the original PSYCHE, and in the Zangerer–Sterk method before it, there is a compromise between sensitivity and purity, and one can only be increased at the cost of the other. Arguably, then, there is not much value in changing the number of saltire pulses as simply varying the *flip angle* of the basic double-saltire already gives the experimentalist a way to balance these competing objectives.

2.4 Direct optimisation of PSYCHE waveform

The changes to the PSYCHE waveform discussed in the previous section—changing the number of saltire pulses to either 1 or 4—are fairly minor, in that they do not fundamentally alter the form of the PSE. Furthermore, there is an obvious issue in that the quality of the spectrum is not measured in a particularly rigorous manner. Ideally, we would like to have a mathematical way of measuring how well the pure shift sequence has worked. Such a metric would also enable a more automated optimisation process, where a programme is allowed to find the best PSE without any interference from human subjectivity.

In this section, I discuss more radical changes which depart from the tried-and-tested saltire pulse. I also show how two different *cost functions*—functions which determine how ‘bad’ a spectrum is—can be used to objectively evaluate PSEs. Although the work in this section did not quite yield any groundbreaking results, it provided substantial insights into the nature of pure shift optimisations, which were later used in the context of POISE (chapter 3).

2.4.1 Techniques for pure shift optimisations

Throughout this chapter (and more generally this thesis), various algorithms are used for *optimisation*: that is, to find the parameters $\mathbf{x} \in \mathbb{R}^n$ which minimise a *cost function* $f : \mathbb{R}^n \rightarrow \mathbb{R}$. These algorithms can loosely be categorised as either *derivative-based* or *derivative-free*: the former use extra information in the form of ∇f to help locate the optimum, whereas the latter do not, using only the value of $f(\mathbf{x})$. While derivative-based algorithms typically converge to an optimum more quickly, they are unsuitable for problems where the cost function f is noisy. In this section, the Nelder–Mead (NM) simplex algorithm⁶³ was used: it is a very popular derivative-free method, and although mathematical convergence is not guaranteed,⁶⁴ in practice such cases are rather unlikely to arise. In this section, the implementation of the NM algorithm in the Python `scipy` package was simply used as-is.

The cost function measuring the performance of a pure shift experiment can be measured in one of two ways: either *theoretically*, in that the pure shift experiment is simulated using the density operator formalism, or *experimentally*, in that the experiment is run on a spectrometer. Unfortunately, the simulation of PSYCHE-type PSEs, where a shaped pulse is applied together with a gradient, requires a large amount of time. The pulse itself already has $m \sim 10000$ points, but on top of that, the application of gradients also requires splitting up the sample into multiple slices ($n \sim 100$ to 1000) such that the evolution of ρ can be simulated in each slice and the results summed up. To make matters worse, $H_{\text{pulse},I}$ does not commute with H_{grad} , so a propagator

$$U(i, z) = \exp \left[-i(H_{\text{grad}}(z) + H_{\text{free},I} + c_x^{(i)} I_x + c_y^{(i)} I_y)(\delta t) \right] \quad (2.31)$$

must be calculated for each pulse point, in each slice of the sample, for a total of mn matrix exponentials. As a result, the simulation of PSYCHE spectra in the Spinach package⁶⁵ typically requires minutes to hours.*

This in fact makes it faster to experimentally acquire a pure shift spectrum and calculate a cost function based on that. Running an actual pure shift experiment, though, is suboptimal: firstly, the pseudo-2D interferogram method is slow, and secondly, there is no easy way to devise a cost function for the resulting spectrum without prior knowledge of where peaks and artefacts lie. Instead, we can use a simple 1D ‘J-refocused spin echo’ (JRSE) sequence, which has the form $90^\circ-\tau-\text{PSE}-\tau-\text{detect}$ (fig. 2.8b). An ideal PSE would lead to complete refocusing of both chemical shifts and J-couplings, and the signal detected after this would simply be the same as in a pulse–acquire experiment (fig. 2.8a).

*With highly optimised handwritten code, exploiting the symmetry of the PSYCHE element, the fastest simulation I could do took 16 seconds on a 20-core computer, for a simple 2-spin system. This number increases exponentially for larger spin systems. It is possible that GPU acceleration could result in substantial speedups, but I have not looked into this sufficiently. Anyway, experimental acquisition of the JRSE spectrum (to be described) takes only around 5 seconds.

Of course, there is a sensitivity penalty which reflects that of the PSE (there are also relaxation losses during the τ delays, but these are a constant, unaffected by the form of the PSE). On top of that, if the J-refocusing is not perfect, then the multiplets in the spin echo sequence acquire a degree of phase distortion: the delay τ has to be long enough to allow for this to evolve, but its exact value is otherwise largely insignificant. These distortions are just about visible in fig. 2.8b.

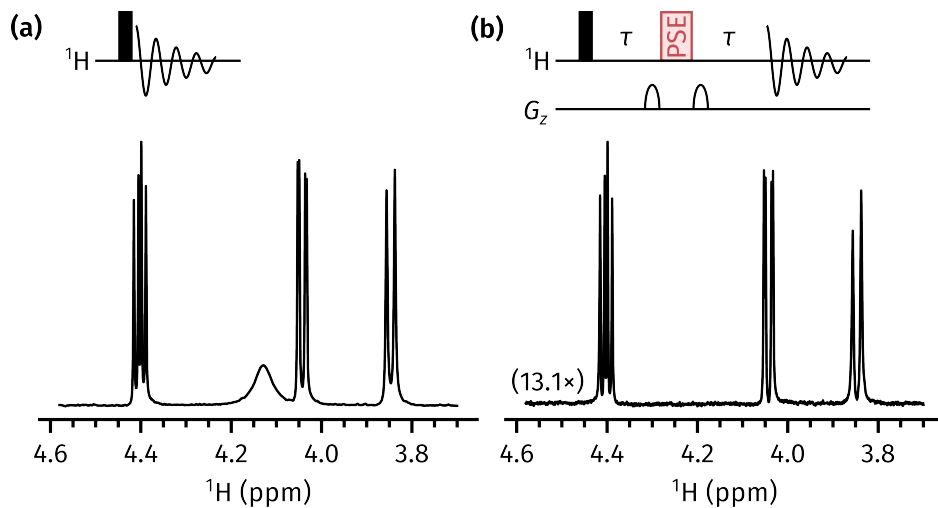


Figure 2.8: (a) Pulse–acquire experiment and the resulting spectrum. (b) J-refocused spin echo experiment and the resulting spectrum. The PSE used was the PSYCHE double saltire, with a flip angle of 25°; the delay τ was 11 ms. The OH peak at 4.1 ppm is lost, most likely due to chemical exchange. *Data code:* 6A-200816.

Two cost functions were designed and used in this section:

$$f_{\text{phase}} = \text{Var}_i \left[\arctan \left(\frac{S_{\text{re},i}}{|S_{\text{im},i}|} \right) \right] \quad (2.32)$$

$$f_{\text{diff}} = \left\| \frac{\mathbf{S}_{\text{re}}}{\|\mathbf{S}_{\text{re}}\|} - \frac{\mathbf{T}_{\text{re}}}{\|\mathbf{T}_{\text{re}}\|} \right\|, \quad (2.33)$$

where the JRSE and pulse–acquire 1D spectra are treated as complex-valued vectors \mathbf{S} and \mathbf{T} respectively (for ‘spectrum’ and ‘target’). \mathbf{S}_{re} and \mathbf{S}_{im} are the real and imaginary parts of the spectrum \mathbf{S} , and $S_{\text{re},i}$ is the i -th point of the real part of the spectrum. The operator Var_i represents the variance over all points in the spectrum (indexed by i), and $\|\mathbf{x}\|$ denotes the 2-norm of the vector \mathbf{x} , i.e. $\sqrt{\sum_i x_i^2}$. Python implementations of these are given in listing 2.1.*

These two cost functions were chosen as they exhibited desirable characteristics on synthetic data (fig. 2.9). In these simulations, the ‘target’ spectrum was chosen to simply be an in-phase

*The use of `np.arctan` (the arctangent), and *not* `np.arctan2` (the argument of a complex number), is intentional. The behaviour shown in fig. 2.9 isn’t reproduced with `arctan2`. Of course, this means the name ‘phase’ is a misnomer; it’s not really the phase of anything meaningful.

```

import numpy as np
# assume S and T are complex numpy arrays which have been read in
Sr = np.real(S); Si = np.imag(S); Tr = np.real(T)
f_phase = np.var(np.arctan(Sr / np.abs(Si)))
f_diff = np.linalg.norm((Sr / np.linalg.norm(Sr))
                        - (Tr / np.linalg.norm(Tr)))

```

Listing 2.1: Pure shift cost functions.

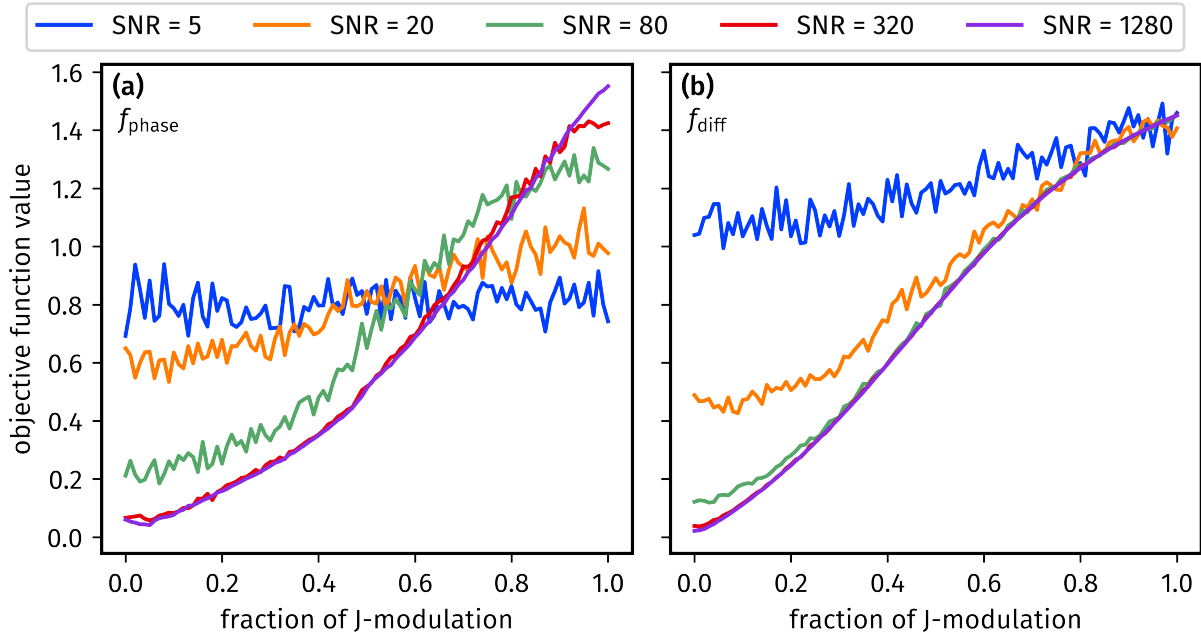


Figure 2.9: Behaviour of the two cost functions, f_{phase} and f_{diff} , on synthetic spectra with various SNRs. Zero phase distortion refers to an in-phase absorption-mode doublet, whereas complete phase distortion refers to an antiphase dispersion-mode doublet. (a) The f_{phase} cost function. (b) The f_{diff} cost function, measured against a spectrum with no phase distortion and an SNR of 500.

absorption-mode doublet with an SNR of 500. Synthetic data with increasing amounts of J-modulation (i.e. spectra ranging from in-phase absorption, to antiphase dispersion) were generated, and extra Gaussian noise added to mimic different SNRs. It can be seen that, for data which have little J-modulation (left edges of the plots), both f_{phase} (fig. 2.9a) and f_{diff} (fig. 2.9b) penalise lower SNRs. Furthermore, both of the cost functions penalise J-modulation, since they increase going from left to right. This penalty is stricter for high-SNR spectra, which is also desirable, since it is only in high-SNR spectra that the J-modulation becomes noticeable.

The cost function f_{diff} is easier to comprehend: it simply scales both the target and JRSE spectra down by their respective intensities, and compares each point to determine whether the peak shapes obtained are similar. Although this seems like it should be agnostic towards signal intensity,

this is only true for noiseless spectra. If a (genuine) JRSE spectrum has low SNR, $\|S_{re}\|$ will be small, and the noise will be scaled down less than for the target spectrum; this difference in the *noise* (rather than the signal) then contributes towards the cost function. On the other hand, a proper rationalisation of why the cost function f_{phase} works is unfortunately not within my capabilities! It was mostly developed by trial-and-error (based on the notion that phase distortions would have something to do with S_{re} and S_{im}), and I do not have a good explanation of why it works.

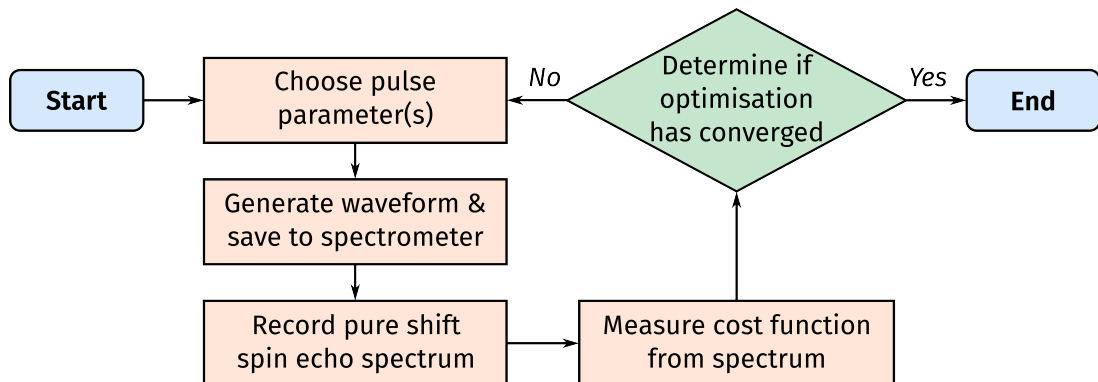


Figure 2.10: Flowchart illustrating the steps for optimisation of a pure shift spectrum.

The general optimisation procedure is conceptually simple and largely consists of the loop shown in fig. 2.10: this is essentially a specialised version of the POISE flowchart (fig. 3.1). The optimisation algorithm is responsible for determining convergence, as well as choosing the new parameters based on previously obtained information; the initial parameters must be supplied by the user.

In practice, it is a technical challenge to implement this loop on a spectrometer as the cost function calculation is performed either in Matlab or Python 3, both of which are not compatible with Bruker's TopSpin software. TopSpin instead provides Jython (Python 2.7) and C programming interfaces;^{*} the former is not compatible with Python 3 packages like numpy or scipy, and the latter is too low-level to be worth implementing numerical algorithms in.[†] Thus, we require a means of *communication* between the spectrometer and the optimisation control programme: this includes a signal from the controlling programme to trigger acquisition on the spectrometer, as well as a signal from the spectrometer that acquisition is done so that the cost function can be calculated. As it turns out, the code used for the optimisations in this section was a very rudimentary and fragile form of that eventually used in POISE (for example, the aforementioned

^{*}TopSpin 4.1.4 introduced a Python 3 interface which would have made much of this work simpler. Unfortunately, this was not available at the time of this work.

[†]Of course, heavily optimised code in low-level languages such as C and Fortran—or perhaps even Matlab—would run faster. However, speeding up the code has virtually no impact on the optimisation, since its rate is limited by spectrum acquisition. In this situation, it makes far more sense to save *developer time*.

signals were transmitted via the creation and deletion of files). I therefore defer the discussion of this issue to § 3.2.5, where the more robust POISE interface is explained in detail.

2.4.2 Flip angle optimisation

Having described the rest of the optimisation setup, it remains to choose exactly which parameters are subjected to optimisation. The simplest option is to only optimise one parameter, namely the flip angle of the (double-)saltire pulse. The flip angle dependence of PSYCHE spectra is well-understood, which crucially allows us to evaluate the cost functions outlined above and determine whether they are functioning correctly.

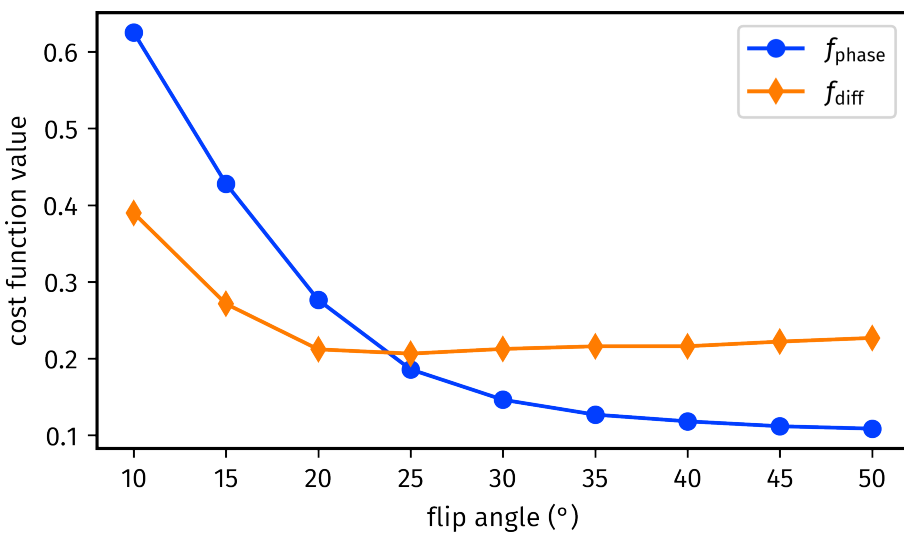


Figure 2.11: Behaviour of the two cost functions, f_{phase} and f_{diff} , on experimental J-refocused spin echo spectra. Data code: 5C-190809.

I first sought to measure how the cost functions described above varied with the flip angle. To this end, JRSE spectra using a *single* saltire as the PSE and various flip angles (from 10° to 50°) were acquired (fig. 2.11). Since both cost functions penalise both low sensitivity and low purity, we might expect that there is be an intermediate value where neither sensitivity nor purity were penalised too much: this would be the ‘optimum’ flip angle. In the event, it was found that only f_{diff} yielded a useful—albeit shallow—minimum at around 25° (recall that the PSE here is a single saltire, so this corresponds roughly to a 15°–20° double saltire). The f_{phase} cost function, on the other hand, was strictly decreasing within the range of flip angles tested: it is possible that there is an optimum at an even larger flip angle, but this would have been a scientifically unsound conclusion given that the sample was decently concentrated (50 mM).

This naturally raises the question of why f_{diff} yields an optimum which falls within what we would consider a ‘sensible’ region. It is tempting to believe that the form of f_{diff} (eq. (2.33)), which is at first glance quite intuitive, naturally leads to a better result. I argue here, however,

that this is mostly down to *coincidence*. This is difficult to explain quantitatively, but in a broader sense, we may imagine that the cost function separately penalises low sensitivity and low purity, i.e. it can be decomposed into something of the form

$$f = g(s) + \lambda h(p), \quad (2.34)$$

where s and p are respectively the sensitivity and purity, and g and h are some unknown functions which *decrease* with increasing s and p (since we want to minimise the cost function). This is clearly a simplification, because the plots in fig. 2.9 show that the effects of sensitivity and purity on the cost functions are not additive; however, it is sufficient to make the point here. The parameter λ represents the relative weighting of purity to sensitivity: if λ is large than the purity is more strongly emphasised, and vice versa if λ is small.

When we say that an optimum is ‘sensible’ or ‘sound’, this is with respect to the sensitivity/purity balance in the parent pseudo-2D homodecoupled spectrum. In other words, what we *really* seek is a cost function which measures the sensitivity and purity of that spectrum:

$$f' = g'(s') + \lambda' h'(p'), \quad (2.35)$$

where everything is marked with a prime symbol to indicate that it is with respect to the decoupled spectrum, and the parameter λ' is chosen to fit our judgement of the required balance, or in other words, yield an optimum of around 20°. Now, we may reasonably assume that s and s' are proportional, but p and p' are hardly the same thing: one is manifested in terms of artefact intensity and the other in terms of phase distortions. Furthermore, the λ provided to us by the cost function f may not necessarily be the same as our ideal choice of λ' ; let alone the forms of the functions g and h . The fact that eq. (2.34) *happens* to provide the same flip angle optimum as the idealised eq. (2.35) cannot truly be attributed to design!

Of course, just because a cost function works mostly by serendipity does not mean that it cannot be used. So, I ran several *actual* optimisations of the flip angle using the cost function f_{diff} , which reliably converged to optima between 20° and 25° regardless of the initial point chosen. Typically, around 10 spectrum acquisitions were required, corresponding to a time of around 2–4 minutes. This is not surprising in light of our knowledge of f_{diff} ; however, it provides us with a proof-of-principle that automated optimisation of NMR parameters is possible. Further optimisations of the PSYCHE flip angle are also discussed in § 3.4.8.

2.4.3 Waveform parameterisation and optimisation

Given that a working optimisation setup, including cost functions, had been developed, it was a logical step to then test it out on a more challenging problem: namely, how the waveform used in

the PSYCHE PSE could be modified. This goes beyond simply modifying the number of saltires, as was done in § 2.3. There is no real reason why the pulse *must* be an integer number of saltires: in principle it can have *any* shape, although being symmetric about the centre of the pulse would likely still be beneficial in terms of preserving the mechanism of spatiotemporal averaging.

Sum-of-chirp parameterisation

A naive attempt at optimising the pulse would simply involve modifying every pulse point in the double-saltire waveform used in the PSYCHE element. As described in § 1.2, each pulse point consists of a pair of x - and y -amplitudes (c_x, c_y); therefore, for a pulse with m points, we would have a parameter vector $\mathbf{x} \in \mathbb{R}^{2m}$. Unfortunately, for PSYCHE, m is on the order of 10000, and an optimisation with 20000 points is totally unfeasible.*

As a result of this, we must consider other ways of parameterising the waveform which use fewer degrees of freedom. Several approaches to this issue have surfaced in the literature, such as the use of Fourier series,^{37,70–73} Gaussian cascades,⁷⁴ or spline interpolation between a subset of pulse points.⁷⁵ In this instance, we can use the knowledge that the PSYCHE PSE is composed of saltire pulses to our advantage. Each saltire pulse is a linear combination of two chirps, defined by:

$$\phi(t) = \phi_0 + 2\pi f_0 t + \pi\tau_p(\Delta F) \left(\frac{t}{\tau_p} - \frac{1}{2} \right)^2 \quad (2.36)$$

$$c_x(t) = A \cos[\phi(t)] \quad (2.37)$$

$$c_y(t) = A \sin[\phi(t)] \quad (2.38)$$

for $t \in [0, \tau_p]$. Here, τ_p is the duration of the chirp, A is the amplitude of the chirp (which is time-independent), ϕ_0 the phase of the chirp, ΔF the bandwidth, and f_0 its frequency offset (i.e. where the centre of the bandwidth lies). The latter two parameters are assumed to be given in units of ordinary frequencies (Hz), not angular frequencies. (Note that eqs. (2.37) and (2.38), defining the x - and y -coefficients of the pulse, are just a restatement of eq. (1.30).)

Given these expressions, we see that there are five parameters of the chirp which can be modified: A , τ_p , ϕ_0 , ΔF , and f_0 . The two chirps which form one saltire pulse simultaneously sweep in opposite directions, which mean that ΔF for one chirp is the negative of the other; however, their parameters are otherwise equal. We may, however, also envision a case where the pulse is constructed from two chirps which are applied at a different point in time. This adds one more

*Although problems of this size have been tackled using optimal control theory,^{66–69} it is not really feasible to use it in cases where the pulse is applied *together* with a gradient, as is the case in PSYCHE. On top of that, the coupling networks and spin systems of interest are rather more complicated than in typical applications of optimal control.

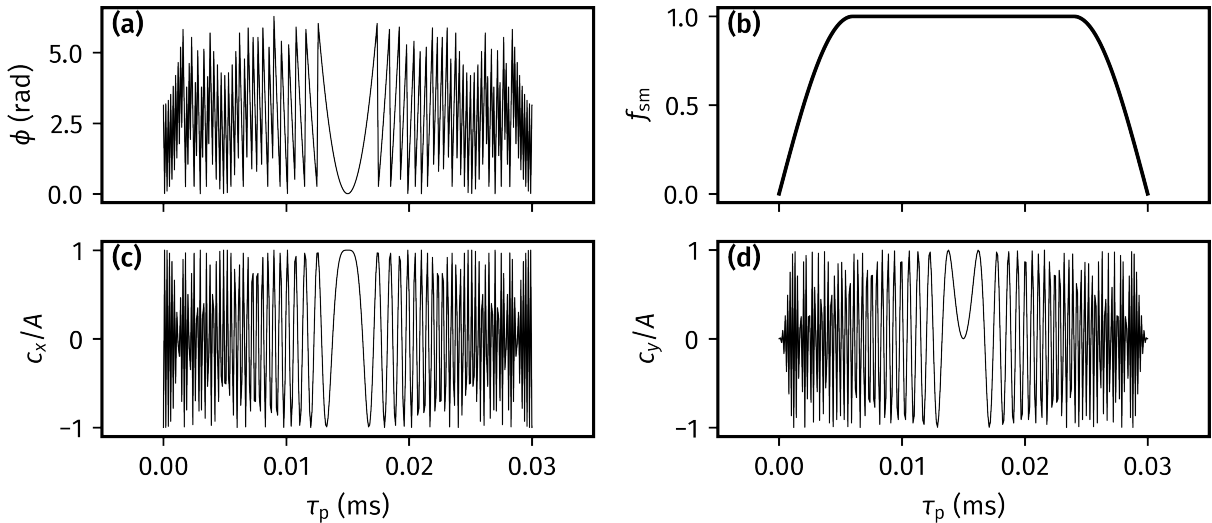


Figure 2.12: Plots of the quantities in eqs. (2.36) to (2.38) for a typical chirp pulse ($\tau_p = 30$ ms, $\Delta F = 10$ kHz). (a) $\phi(t)$ wrapped to the range $[0, 2\pi]$. (b) The quarter-sine smoothing profile which is later applied to the entire waveform (eq. (2.39)); here $s_{sm} = 0.2$. (c) $c_x(t)/A$ prior to smoothing. (d) $c_y(t)/A$ prior to smoothing. The amplitude A is simply a constant, which is related to the flip angle of the chirp.³⁵

parameter to each chirp, namely t_0 , the starting time of the pulse. Each chirp therefore sweeps from the frequency $f_0 - (\Delta F)/2$ at a time t_0 , to the frequency $f_0 + (\Delta F)/2$ at a time $t_0 + \tau_p$. In total, this gives us 12 parameters to optimise.^{36,76}

After constructing the sum of two chirps, ‘empty’ regions of no RF application ($c_x = c_y = 0$) at the beginning and the end were trimmed off. Since a sum of two chirps is not necessarily symmetric (with respect to reflection in time), the waveform was then reflected about its end, thus doubling the length of the pulse. The entire waveform was then multiplied by a *smoothing function* $f_{sm}(t)$ (fig. 2.12b), which prevents large jumps in RF amplitude at the beginning and end of the pulse. f_{sm} depends on a smoothing parameter s_{sm} , which is typically 0.1–0.2:

$$f_{sm}(t) = \begin{cases} \sin\left(\frac{\pi t'}{2s_{sm}}\right) & 0 \leq t' < s_{sm}; \\ 1 & s_{sm} \leq t' < 1 - s_{sm}; \\ \sin\left[\frac{\pi(1-t')}{2s_{sm}}\right] & s_{sm} \leq t' \leq 1, \end{cases} \quad (2.39)$$

where $t' = t/\tau_p$ (and here τ_p refers to the duration of the *entire* waveform, after reflection).

Optimisations

The initial point chosen was:

- Chirp 1: $\tau_p = 15$ ms; $\Delta F = -5$ kHz; $\phi_0 = 0$; $A = 36$ Hz; $t_0 = 0$; $f_0 = 5$ kHz;

- Chirp 2: $\tau_p = 15$ ms; $\Delta F = 5$ kHz; $\phi_0 = 0$; $A = 36$ Hz; $t_0 = 0$; $f_0 = -5$ kHz.

After the sum of these two pulses is reflected about its end, we obtain a single saltire with bandwidth 10 kHz, duration 30 ms, and an amplitude of 72 Hz, corresponding to a flip angle of approximately 32°.

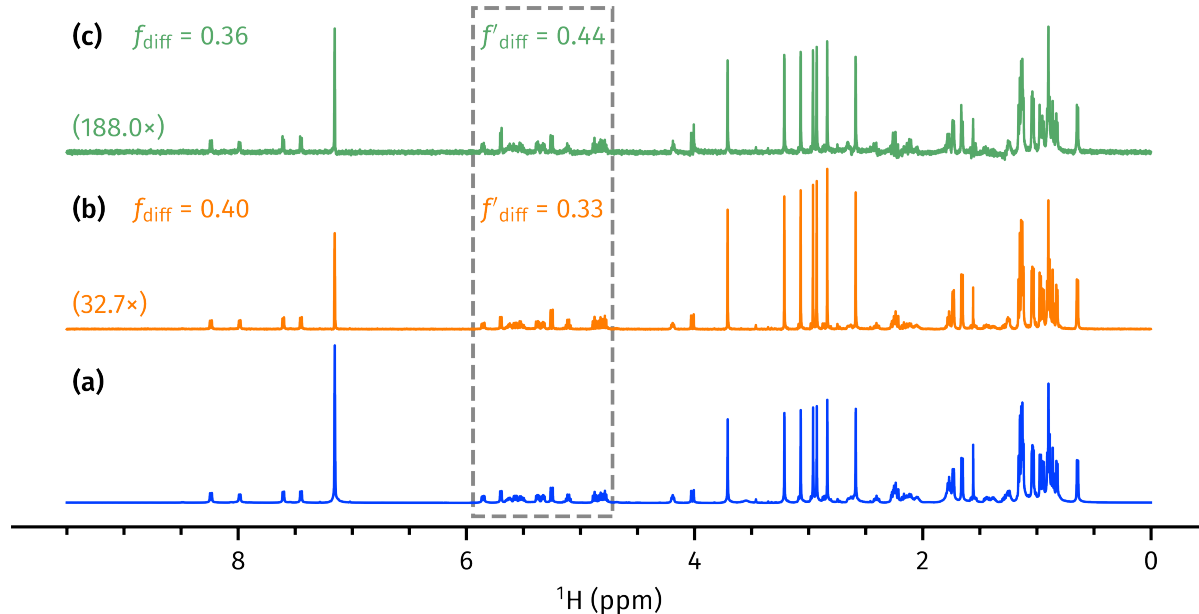


Figure 2.13: (a) Target spectrum (pulse–acquire). (b) JRSE spectrum obtained using the initial guess (a saltire). (c) JRSE spectrum obtained with a spurious optimum point. The grey dotted box shows the restricted region over which f_{diff} was subsequently applied to, yielding a formally different cost function f'_{diff} ; this yielded more sensible results where the cost function for the saltire pulse was smaller. *Data code:* 6C-190823.

Using this setup, several optimisations of the 12 parameters above were conducted using the f_{diff} cost function, which had performed well enough in the flip angle optimisations. However, it was quickly noticed that this led to spurious optima being located, such as the one in fig. 2.13c. Although the initial point (a saltire) yielded a much larger SNR (fig. 2.13b), the value of f_{diff} was still larger (i.e. worse) than for this false optimum. The reason for this is almost certainly because the saltire pulse distorts the relative intensity of the strong singlets in the spectrum, notably the C₆D₅H peak at 7.15 ppm, and the N-methyl groups between 2.5 and 3.8 ppm, whereas the false optimum does not (most likely by coincidence).

Of course, singlets are completely unimportant when devising a pure shift experiment. Unfortunately, singlets are also typically more intense than the rest of the spectrum, and thus contribute disproportionately to the cost function. A simple and effective way to circumvent this is to restrict the region of the spectrum being evaluated. In this case, I chose to use the cyclosporin H α region between 4.72 and 5.94 ppm (grey dotted box in fig. 2.13). This yields a formally different cost function, which I label as f'_{diff} . With this, much more logical behaviour was observed: in

particular, the saltire pulse performed better than the spurious optimum previously found.

While this new cost function could be successfully used to run optimisations, most of these unfortunately failed to find anything performing better than the original saltire pulse. On the rare occasion where something ‘better’ was found (as judged by the new cost function f'_{diff}), these ‘optimised’ pulses were fairly close to a saltire, and the corresponding decreases in the cost function extremely small—suggesting that the ‘better’ result may simply just have been due to noise in the cost function. Nevertheless, these new ‘optima’ *did* function as perfectly serviceable PSEs: for example, fig. 2.14 compares a triple spin echo (TSE) PSYCHE spectrum obtained with an ‘optimised’ pulse to one obtained with the single saltire pulse. There is virtually no difference. This is a meaningful result, as it demonstrates that f'_{diff} is actually an accurate metric to determine the quality of a PSE (it is noisy, but this is to be expected of an experimentally measured cost function). Unfortunately, although the pulse shape is shown in fig. 2.14b, the exact parameters which led to this pulse shape have been lost to time.

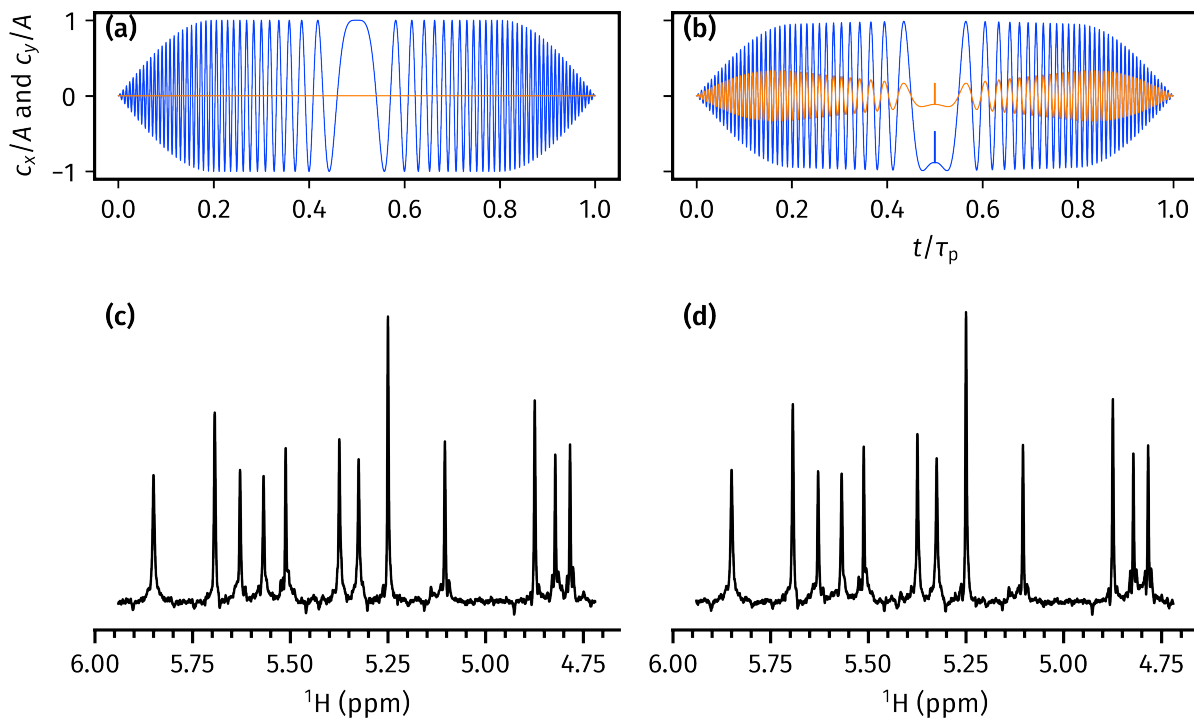


Figure 2.14: (a) x - and y -coefficients of the initial saltire pulse (as a fraction of the maximum amplitude A). (b) x - and y -coefficients of the ‘optimised’ pulse (as a fraction of the maximum amplitude A). (c) TSE-PSYCHE spectrum obtained with the initial guess (a saltire pulse). (d) TSE-PSYCHE spectrum obtained with the ‘optimised’ pulse, with virtually equivalent performance. Data code: 6C-190831.

One issue is that the optimisation algorithms used here can only perform a *local* optimisation, i.e. it may not necessarily locate a global optimum. We cannot rule out the possibility that *some* pulse within this parameter space may in fact outperform a saltire pulse. However, finding this optimum—which may lie very far away from the initial guess of a saltire pulse—is not

generally something which can be accomplished in a reasonable amount of time, even though 12 parameters is far more tractable than 20000. Furthermore, although the cost functions described here do work, they are generally quite ‘flat’ in that they do not discriminate very sharply between ‘good’ and ‘bad’ spectra. Combined with the fact that the cost function is noisy, this makes experimental optimisation of the waveform an uphill task. Nevertheless, much of the knowledge (and code) in this section was later used in the development of POISE.

In the next two sections in this chapter, I move on from PSYCHE and instead discuss completely different methods of obtaining pure shift spectra using only hard pulses in the PSE.

2.5 Time-reversal method

This section focuses on the time-reversal element briefly introduced in § 2.2.2: the aim is to determine whether a pure shift method based on this can have better performance than PSYCHE. Before that, I will first provide a more detailed theoretical analysis of it. The time-reversal element consists of just a single hard pulse with a flip angle β , and we first analyse its potential use as a JRE (instead of a PSE), using eqs. (2.27) to (2.29).

Recall that an ideal JRE should invert passive spins only, not active spins: thus, we seek a transformation of the form $I_{1+}I_{2\alpha} \rightarrow I_{1+}I_{2\beta}$. However, the hard pulse does this:

$$I_{1+}I_{2\alpha} \rightarrow \underbrace{c^2 s^2 I_{1+}I_{2\beta}}_{\text{term 1}} + \underbrace{c^4 I_{1+}I_{2\alpha}}_{\text{term 2}} - \underbrace{\frac{S^2}{4} I_{1\alpha}I_{2+}}_{\text{terms 3 and 4}} + \underbrace{\frac{S^2}{4} I_{1\beta}I_{2+}}, \quad (2.40)$$

where (as before) $S = \sin \beta$, $s = \sin(\beta/2)$, and $c = \cos(\beta/2)$. As in the analysis for PSYCHE (§ 2.2.3), term 1 here represents the desired signal; term 2 recoupling artefacts; and terms 3 and 4 COSY-type artefacts arising from coherence transfer. Terms with difference coherence orders have been neglected as they can be removed using CTP gradients.* Unlike in the PSYCHE analysis, however, we have not neglected any other terms of smaller order in s .

Since the desired and undesired terms have different coefficients, it is possible to *fully cancel out the recoupling artefacts* by recording (in this case) two different spectra with different values of β and performing an appropriate linear combination:

$$I_{1+}I_{2\alpha} \xrightarrow{\beta=0^\circ} I_{1+}I_{2\alpha} \quad (2.41)$$

$$I_{1+}I_{2\alpha} \xrightarrow{\beta=90^\circ} \frac{1}{4}I_{1+}I_{2\beta} + \frac{1}{4}I_{1+}I_{2\alpha} - \frac{1}{4}I_{1\alpha}I_{2+} + \frac{1}{4}I_{1\beta}I_{2+} \quad (2.42)$$

If we take eq. (2.42) minus 1/4 of eq. (2.41), the recoupling artefacts (arising from the $I_{1+}I_{2\alpha}$

*In the original work¹³ which predated the widespread use of field gradients, other terms were removed through phase cycling, which is essentially equivalent.

term) are fully removed. In general, for an N -spin system, there are several different ‘types’ of recoupling artefacts where different numbers of passive spins (between 1 and $N - 1$) are not inverted. Each of these pathways will have different coefficients, as each spin that is flipped contributes s^2 , whereas each spin that is not flipped contributes c^2 . Suppressing all of these requires the acquisition and summation of N spectra with different flip angles and appropriate weights.

Before we go on further, notice even in the two-spin system that the COSY-type artefacts are *not* suppressed! In the original work,¹³ this time-reversal element was used in the middle of the t_1 period in a NOESY experiment. The coherence transfer peaks were not deemed to be problematic in this context: they gave rise to artefacts which had F_1 frequencies of $(\Omega_1 + \Omega_2)/2$, but different phase properties to genuine crosspeaks, allowing them to be easily identified. Of course, these artefacts are not acceptable in an actual pure shift spectrum.

To remove these peaks, I adopted the strategy first reported by Thrippleton *et al.* for suppression of COSY-type transfer pathways in 2DJ spectra.⁴ In a 2DJ experiment, the central 180° pulse should in principle not cause coherence transfers between different spins; however, in strongly coupled systems this can happen. The solution was to bracket the 180° pulse, as well as half of the t_1 period, with a pair of opposing chirps and gradients. The same idea was also used in the TSE-PSYCHE experiment¹¹ to (further) suppress strong coupling responses in the parent PSYCHE experiment. This works because the unwanted CTPs have coherences on different spins during each of the two chirps; consequently, the coherences are inverted at different times by the chirp pulses, and are ultimately dephased by gradients. The resulting time-reversal experiment was thus simply the same as the parent TSE-PSYCHE experiment, except that the central PSYCHE element was replaced by a β hard pulse (fig. 2.15).

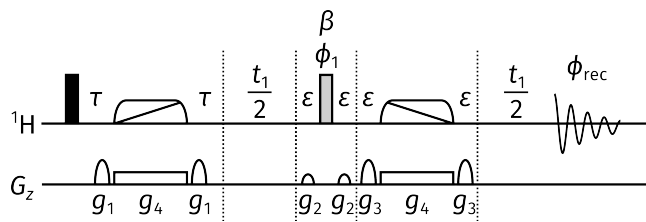


Figure 2.15: Time-reversal pure shift pulse sequence. The flip angle β is varied as described in eq. (2.43). Pulse phases are: $\phi_1 = (x, y, -x, -y)$; $\phi_{\text{rec}} = (x, -x, x, -x)$. The delay τ is set to $1/(4 \cdot T_{\text{chunk}})$, and allows for J-coupling to be refocused in the middle of the chunk. Gradient amplitudes are $(g_1, g_2, g_3, g_4) = (35\%, 49\%, 77\%, 1\%)$ (note that, in principle, g_4 should be calibrated according to the bandwidth of the chirp used).

In such a sequence, the JRE is in fact not only the β pulse itself but also the second chirp (which takes the place of a 180° pulse). So, the β pulse here fulfils the role of a PSE, not a JRE. To accomplish this for a system containing N mutually coupled spins, N different experiments must

be acquired using the following flip angles β_j and summed with the corresponding weights W_j ($j = 1, 2, \dots, N$):

$$\beta_j = \frac{j\pi}{N} \quad (2.43)$$

$$W_j = \frac{N}{8} \cdot \frac{(-1)^j}{\sin^2(\beta_j/2)}. \quad (2.44)$$

For the sake of completeness, the values of β and W for a JRE are given here as well:

$$\beta_k = \frac{k\pi}{N} \quad (2.45)$$

$$W_k = \frac{N}{8} \cdot \frac{(-1)^{k+N}}{\cos^2(\beta_j/2)} \quad (2.46)$$

for $k = 0, 1, \dots, N - 1$. The derivation of these expressions is discussed more thoroughly in a paper by Griesinger et al.⁵ In the context of this specific paper, note that the weights for the PSE correspond to that used for the ECOSY experiment, and the weights for the JRE correspond to that used for the complementary ECOSY experiment.

In practice, $N = 5$ is likely to cover most realistic spin systems. Explicitly evaluating eqs. (2.43) and (2.44) yields $\beta = \{36^\circ, 72^\circ, 108^\circ, 144^\circ, 180^\circ\}$ and $W = \{-6.545, 1.809, -0.955, 0.691, -0.625\}$. Figure 2.16 shows insets of the five subspectra acquired with the above values of β and scaled by their respective weights W . The weighted sum (i.e. the pure shift spectrum) was phased, and the resulting phase correction values were propagated back to the individual subspectra.

Although the experiment seems to work, in that the weighted sum *is* indeed a pure shift spectrum, the fact that it is obtained through summation of N different experiments brings some immediate drawbacks. Firstly, the minimum duration of the experiment is lengthened by a factor of N : this is essentially the same as an N -step phase cycle. However, and perhaps more importantly in the context of *pure shift* NMR, the artefacts surrounding the main peaks are not perfectly cancelled through the process of summation. As a result, random distortions are observed around the desired peaks in the pure shift spectrum: this is noticeable in the 4.04 ppm peak in fig. 2.16, and is even worse for more intense signals.

In terms of sensitivity, the time-reversal spectrum is not particularly exceptional either. Each of the five subspectra above were acquired with 2 scans; when compared against a typical TSE-PSYCHE experiment acquired with only 4 scans (i.e. 40% of the experiment duration), the TSE-PSYCHE experiment had comparable or perhaps even slightly better SNR (fig. 2.17). The likely reason for this is because in the time-reversal experiment, signal is actually being *cancelled out* through the process of summation, as is quite clearly shown in fig. 2.16. In principle, the sensitivity of the time-reversal experiment could be optimised by acquiring the more heavily weighted spectra with

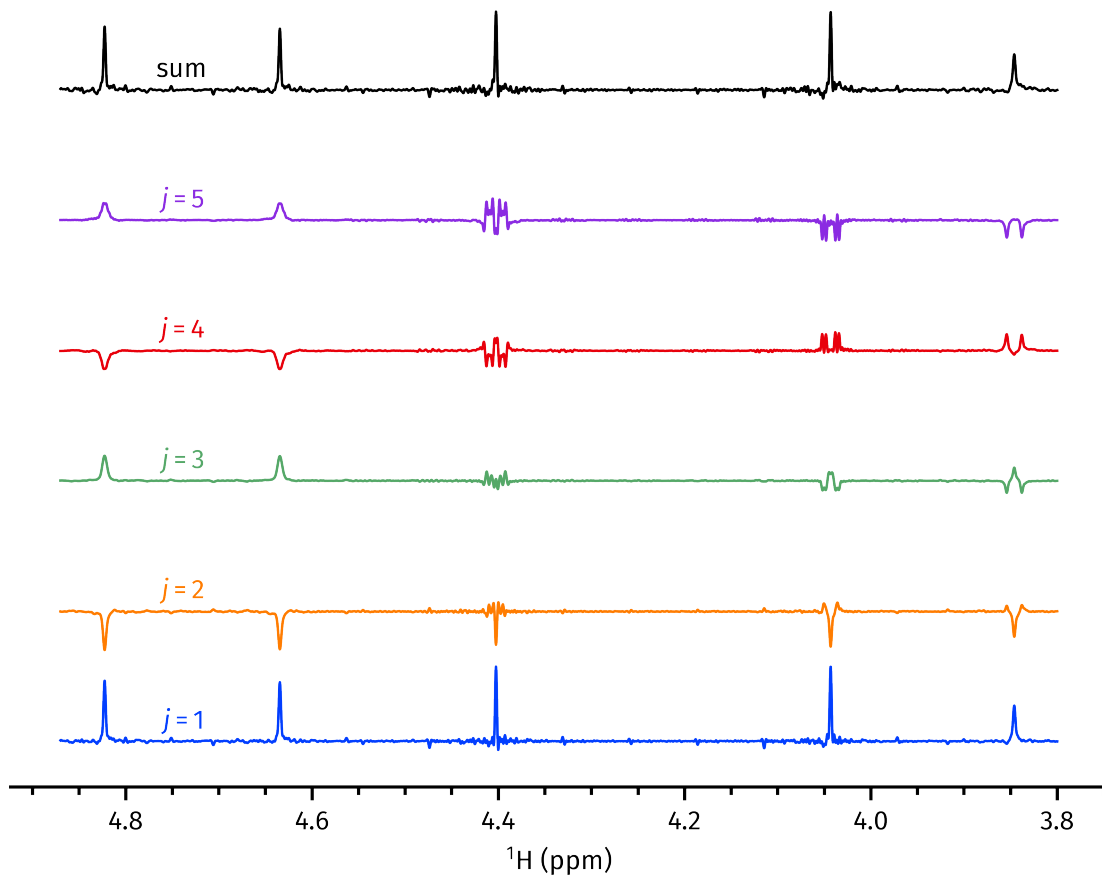


Figure 2.16: Insets of weighted time-reversal subspectra (with $j = 1$ through 5), as well as their sum (the pure shift spectrum). Data code: 7A-201020.

more scans. However, I consider this unlikely to make a substantial difference to the conclusions drawn here. The idea of re-optimising the weights to better suppress artefacts was also briefly considered. However, given that eq. (2.44) already yields *theoretically* complete suppression, it was deemed unlikely that anything substantially better could be obtained, considering that the artefacts arise from the summation process itself and are likely to appear regardless of what weights are chosen. Performing such an optimisation would also require a pure shift spectrum to have been acquired beforehand (for comparison), thus defeating the purpose of optimising the weights anyway.

Interestingly, fig. 2.16 suggests that the $j = 1$ spectrum *on its own* already provides as good a result as the summed pure shift spectrum. This is not surprising, as the use of a hard pulse as the PSE yields a conceptually very similar result to PSYCHE in that the signal-to-artefact ratio depends on $\tan^2(\beta/2)$ (of course, the COSY-type artefacts must still be suppressed through the TSE scheme). This suggests that even without summation of multiple subspectra, the TSE time-reversal pulse sequence in fig. 2.15 is a viable pure shift experiment—albeit one which does not have any significant advantage over PSYCHE.

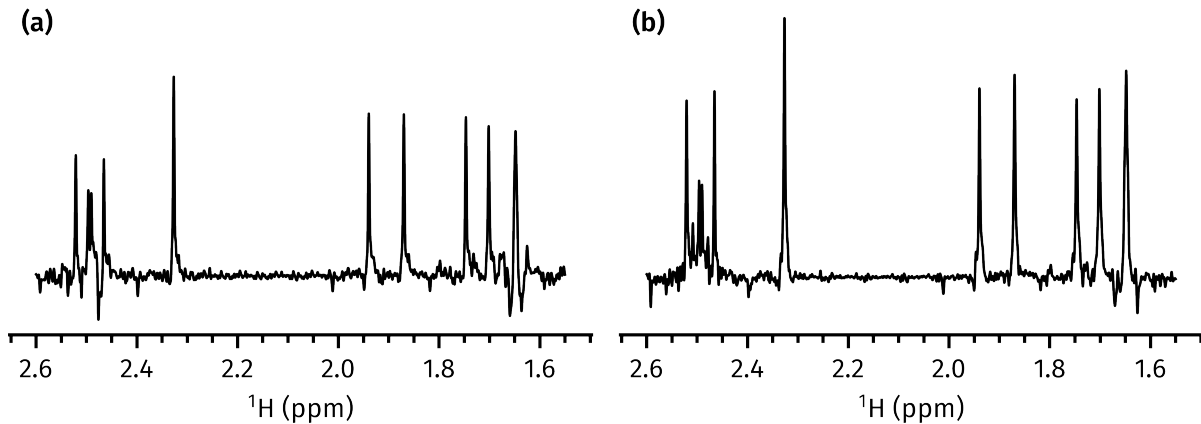


Figure 2.17: Comparison of time-reversal and TSE-PSYCHE sensitivity. (a) Time-reversal pure shift spectrum (the same as the sum in fig. 2.16) acquired with 2 scans for each subspectrum. (b) TSE-PSYCHE (double saltire, flip angle 15°) acquired with 4 scans. The spectra have been scaled so that their noise levels are similar: the signal intensity is comparable, or perhaps slightly better in the TSE-PSYCHE. *Data code: 7A-201020.*

2.6 ‘Discrete PSYCHE’

The last pure shift method in this chapter is completely original, and represents perhaps the most fruitful attempt so far at optimising pure shift experiments. It relies on what is essentially a ‘temporal discretisation’ of the PSYCHE waveform and gradient combination: instead of applying a shaped pulse and a gradient simultaneously, hard pulses and gradients are interleaved in the PSE (fig. 2.18).

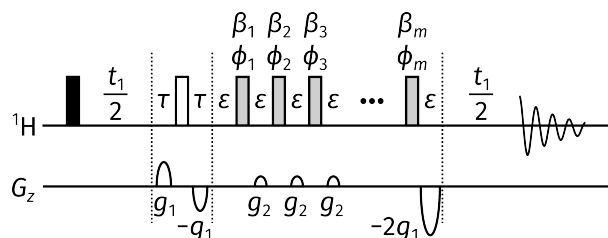


Figure 2.18: dPSYCHE pulse sequence. Gradient amplitudes are $(g_1, g_2) = (35\%, 41\%)$; the gradients in the PSE g_2 are applied with a duration of $500 \mu\text{s}$. The hard pulses in the PSE are applied with an RF amplitude of 18 kHz . The delay τ is set to $1/(4 \cdot T_{\text{chunk}})$, and allows for J-coupling to be refocused in the middle of the chunk.

For this reason I have dubbed this experiment the ‘discrete PSYCHE’, or dPSYCHE for short. There are two major reasons why this is more amenable towards optimisation than many of the previous experiments:

1. Pulses and gradients are no longer applied simultaneously, which makes simulation of the experiment *extremely* fast compared to the original PSYCHE. This opens up the possibility of entirely theoretical optimisations, as the noise can be completely eliminated from the

cost function.

2. There are much fewer ‘pulse points’ than in the original PSYCHE: effectively, the phase and flip angle of every hard pulse has to be optimised, leading to $2m$ parameters. Even for $m \sim 10$, this is quite tractable if the optimisation is noiseless.

One downside of this is that it is difficult, or perhaps even impossible, to explain how the PSE works.* For a symmetric PSE where $\beta_1 = \beta_m$ (and so on) it is probably possible to reuse an explanation based on PSYCHE-style CTP selection, but this is clearly inapplicable if the flip angles and phases are scrambled.

2.6.1 Speeding up dPSYCHE simulations

To begin, I first explain how the exact simulation of dPSYCHE experiments can be greatly accelerated through efficient propagator calculations. Although Spinach⁶⁵ is the leading simulation package for NMR, and covers an extremely impressive range of experiments, this generality also prevents it from providing optimal performance for any single experiment. As it turns out, handwritten, specialised Matlab code can outperform Spinach by orders of magnitude.

The NMR simulations developed here simply use the density operator formalism in Hilbert space, as outlined in § 1.3. The Zeeman basis is used, and non-unitary transformations such as relaxation are neglected. Propagation under the Liouville–von Neumann equation (eq. (1.38)) requires the calculation of matrix exponentials $\exp(-iH\tau)$. For an $N \times N$ matrix, the matrix exponential requires $O(N^3)$ time to calculate (and for a system containing p spins, we have $N = 2^p$); it is often this which is the bottleneck in NMR simulations. Minimising the number of matrix exponentials, and/or their computational cost, is the key to achieving speedups, as will be shown in the following text.[†]

Generally, the accurate simulation of pulsed field gradients requires the sample to be divided up into n discrete slices, each simulated with a different H_{grad} .[‡] Thus, a very naive implementation of the dPSYCHE experiment would require mn matrix exponentials, one per pulse per slice. The

*Of course, we could simulate it and say that it works because the maths says it does; but that isn’t very illuminating. Some of the optimisations done in this thesis are somewhat like a scaled-down version of machine learning, in that they produce better results at the cost of interpretability.

[†]Note that in my simulations, I simply used the built-in `expm` Matlab function, which implements the matrix exponential using a combination of the scaling-and-squaring method and Padé approximation.⁷⁷ This is in contrast to Spinach, which primarily uses a scaled-and-squared Taylor series (according to the `propagator.m` file, various other methods supposedly did not ‘live up to their marketing’). An in-depth discussion of matrix exponential methods is outside the scope of this thesis, but can be found in a classic paper by Moler and Van Loan.⁷⁸

[‡]In simple cases this can be avoided by simply removing all terms with the wrong coherence orders as we know they will be dephased (eq. (1.110)), but this is too naive an approach for pure shift techniques.

overall structure of the code would resemble listing 2.2.* Note that the term H_{free} , and the corresponding propagator U_{free} , technically refer to the interaction-picture free Hamiltonian, $H_{\text{free},I}$.

```
% loop over slices
for slce=1:n
    H_grad = I_z * G * z(slce);

    % loop over pulse points
    for pulse_point=1:m
        H_pulse = (c_x(pulse_point) * I_x) + (c_y(pulse_point) * I_y);

        % calculate propagators; m*n total matrix exponentials
        U_pulse = expm(-1j * (H_free + H_pulse) * t_pulse);
        U_grad = expm(-1j * (H_free + H_grad) * t_grad);

        rho = U_grad * U_pulse * rho * U_pulse' * U_grad';
    end
end
```

Listing 2.2: Rough structure of a naive dPSYCHE implementation. Note that I use the variable name `slce` as `slice` is an existing builtin Matlab function.

It is not difficult to come up with a more sensible approach which cuts this down by a factor of m : since the pulses are not applied together with the gradients, the pulse propagators U_{pulse} can be pre-calculated outside of the loop. Furthermore, all of the gradients within the PSE are the same, so U_{grad} can be moved out of the inner loop (listing 2.3).

Spinach, which is designed to be general, has no idea that these optimisations are possible, so is naturally rather slower. However, even this is relatively inefficient. It can be shown that the two components of the gradient propagator, $H_{\text{free},I}$ and H_{grad} , actually commute with one another (even in the strong coupling case). Thus, we can write:

$$\exp[-i(H_{\text{free},I} + H_{\text{grad}})t] = \exp(-iH_{\text{free},I}t) \exp(-iH_{\text{grad}}t) \quad (2.47)$$

(in general, for matrices A and B , $\exp(A + B) = \exp(A) \exp(B)$ if and only if $[A, B] = 0$). This on its own does not reduce the number of matrix exponentials required, but notice now that H_{grad} is a sum of I_{iz} terms and is therefore *diagonal* in the Zeeman basis. The exponential of a diagonal

*Strictly speaking, there is a slight inaccuracy in this code: the final gradient should have strength $-2G$ and not G , but that is a minor detail which I leave out for clarity in the code.

```
% precalculate pulse propagators; m total matrix exponentials
for pulse_point=1:m
    H_pulse = (c_x(pulse_point) * I_x) + (c_y(pulse_point) * I_y);
    U_pulse(m) = expm(-1j * (H_free + H_pulse) * t_pulse);
end

% loop over slices
for slce=1:n
    % calculate gradient propagators; n total matrix exponentials
    H_grad = I_z * G * z(slce);
    U_grad = expm(-1j * (H_free + H_grad) * t_grad);

    % loop over pulse points
    for point=1:m
        rho = U_pulse(m) * rho * U_pulse(m)';
        rho = U_grad * rho * U_grad';
    end
end
```

Listing 2.3: Rough structure of a slightly faster implementation of dPSYCHE.

matrix is almost trivial to calculate, as we only need to exponentiate the diagonal *elements*:

$$\exp \begin{pmatrix} \lambda_1 & 0 & \dots & 0 \\ 0 & \lambda_2 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & \lambda_n \end{pmatrix} = \begin{pmatrix} \exp(\lambda_1) & 0 & \dots & 0 \\ 0 & \exp(\lambda_2) & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & \exp(\lambda_n) \end{pmatrix}. \quad (2.48)$$

Instead of using the $O(N^3)$ `expm(M)` function, this can instead be done in $O(N)$ time using `diag(exp(diag(M)))` (the `diag` Matlab function converts a diagonal matrix to a vector of its diagonal entries, and vice versa). So, we now require only $m+1$ ‘true’ matrix exponentials. At this point, our matrix exponentials have almost been eliminated and the largest remaining bottleneck is almost certainly the matrix *multiplications* required for the propagation. We can cut these down by a factor of two simply by calculating the overall propagator

$$U_{\text{total}} = U_n \cdots U_2 U_1 \quad (2.49)$$

and then performing the propagation only at the very end:

$$\rho = U_{\text{total}} \rho_0 U_{\text{total}}^\dagger \quad (2.50)$$

instead of performing every individual step $\rho \rightarrow U_1 \rho_0 U_1^\dagger$. The final optimised code therefore resembles that in listing 2.4.

```
% precalculate propagator due to free evolution during gradient
% only 1 matrix exponential
U_free = expm(-1i * H_free * t_grad);

% precalculate pulse propagators; m total matrix exponentials
for pulse_point=1:m
    H_pulse = (c_x(pulse_point) * I_x) + (c_y(pulse_point) * I_y)
    U_pulse(m) = expm(-1j * (H_free + H_pulse) * t_pulse)
end

% loop over slices
for slce=1:n
    % initialise propagator for this slice
    U_slce = eye(2 ^ p);

    % calculate gradient propagators; no matrix exponentials required
    H_grad = I_z * G * z(slce);
    U_grad = U_free * diag(exp(diag(-1j * H_grad * t_grad)));

    % loop over pulse points
    for point=1:m
        U_slce = U_pulse(m) * U_slce;
        U_slce = U_grad * U_slce;
    end

    % perform propagation only at the end
    rho = U_slce * rho * U_slce';
end
```

Listing 2.4: Rough structure of a fast dPSYCHE implementation.

The performance of this handwritten code as compared to Spinach is summarised in table 2.1. In all cases, the spectra produced by the two methods were entirely equivalent. It should be noted that the handwritten code does not even utilise CPU parallelisation, whereas Spinach does. I investigated the possibility of parallelising the loop over slices (replacing the outer `for` in listing 2.4 with `parfor`): however, this in fact made the code *slower*, presumably due to overhead. This is a good thing: it means that `parfor` can be used in an external loop, e.g. for the parallel simulation of the dPSYCHE experiment on different spin systems.

Number of spins	Number of couplings	Execution time (s)	
		Spinach	Handwritten
1	0	3.33	0.35
2	0	4.59	0.32
	1	6.22	0.32
3	0	9.90	0.43
	1	12.95	0.45
	2	31.86	0.47
	3	35.01	0.48
4	0	30.59	1.02
	1	38.63	0.99
	2	44.61	1.04
	3	365.04	1.52
	4	446.03	1.57
	5	521.72	1.57
	6	588.31	1.69

Table 2.1: Comparison of wall-clock times for dPSYCHE simulations. The dPSYCHE sequence used contained 15 pulses, each applied with a flip angle of 15° and a phase of 0° . Spin systems were generated pseudo-randomly. All timings were measured on a 2019 MacBook Pro with a 2.6 GHz 6-core Intel i7 CPU.

2.6.2 Optimisations and experimental evaluation

Spectrum-comparison cost function

As previously discussed, the fact that the dPSYCHE experiment can be very quickly simulated opens up the potential for entirely computational optimisation of the pulse sequence. For any arbitrary spin system, it is trivial to remove all the couplings *in silico* and simulate a pulse–acquire spectrum: this gives us a theoretically perfect pure shift spectrum. The simulated dPSYCHE experiment (on a system with couplings) can then be compared against this. The entire process is repeated using s different spin systems, and the cost function is defined as:

$$f_{\text{diff},2} = \frac{1}{s} \sum_s \left\| \frac{\mathbf{S}_{\text{re}} - c\mathbf{T}_{\text{re}}}{c\|\mathbf{T}_{\text{re}}\|} \right\|^2 \quad (2.51)$$

where \mathbf{S} is the dPSYCHE spectrum, and \mathbf{T} is the target spectrum. The prefactor c will be discussed later; for now I treat it as 1. This cost function appears superficially very similar to f_{diff} (discussed in § 2.4.1), and is based on the same principle that we want the spectra \mathbf{S} and \mathbf{T} to match one another, but there are several points of note:

- The spectrum \mathbf{S}_{re} is not scaled down by its norm. This means that the sensitivity penalty

no longer comes from the difference in the noise (as was previously the case), but rather directly from the difference in peak intensity. Since simulated spectra are noiseless, the original f_{diff} would not work here.

- The division of the entire cost function by $\|T_{\text{re}}\|$ is not important if only one spin system is being simulated as it is simply a constant factor. However, if more than one spin system is being simulated, $\|T_{\text{re}}\|$ differs from system to system and this factor helps to essentially normalise the contributions from each spin system.
- The norm in the cost function here is squared. Again, this makes no difference to the optimum if only one spin system is being investigated, because x^2 is strictly increasing for $x > 0$.^{*} However, for multiple spin systems it makes sense to square the norm, as the largest deviations will be penalised more greatly: this means that a pure shift spectrum which works reasonably well across a wide range of spin systems will be prioritised over one which works perfectly well for some and fails badly for others.

I began by first checking how many t_1 increments (i.e. chunks) were required in the simulation to obtain reliable cost function values. If too few chunks are simulated, the resulting pure shift spectrum will have truncation artefacts, which are likely to mask artefacts from unwanted CTPs. The value of $f_{\text{diff},2}$ was thus tested with a wide variety of randomly chosen phases and angles, with the number of chunks set to 4, 8, and 16 (fig. 2.19).

The 16-chunk and 8-chunk $f_{\text{diff},2}$ (fig. 2.19a) do in fact line up quite well. Notably, as the inset shows, they both agree on the ‘best’ candidate (note that this is not necessarily anywhere near a *perfect* candidate, since these inputs were randomly generated). The 4-chunk $f_{\text{diff},2}$ also has the correct overall behaviour (fig. 2.19b). However, its ranking of the ‘best’ candidates is not very accurate: the 4-chunk cost function rates the red cross in the inset as the optimum, but that is only the 13th-best candidate when using the 16-chunk cost function. Ultimately, I decided to use the 4-chunk cost function for ‘quick and dirty’ optimisations, where only an approximate optimum was required. However, for anything requiring more accuracy, the 8-chunk cost function was used.

An optimisation was then carried out with a (rather arbitrarily chosen) setting of $m = 9$, i.e. nine hard pulses in the PSE. A total of $s = 20$ randomly generated spin systems were used, matching the number of CPU cores on the computer used for the optimisations: these were further subdivided into four two-spin systems, eight three-spin systems, and eight four-spin systems. The derivative-based BFGS algorithm was used to carry out the optimisation: this is a popular line search algorithm which uses an approximate Hessian to calculate the search direction at each iteration.^{79,80} No lower or upper bounds were placed on the flip angles or

^{*}It may affect the rate of convergence, but this is not something I tested.

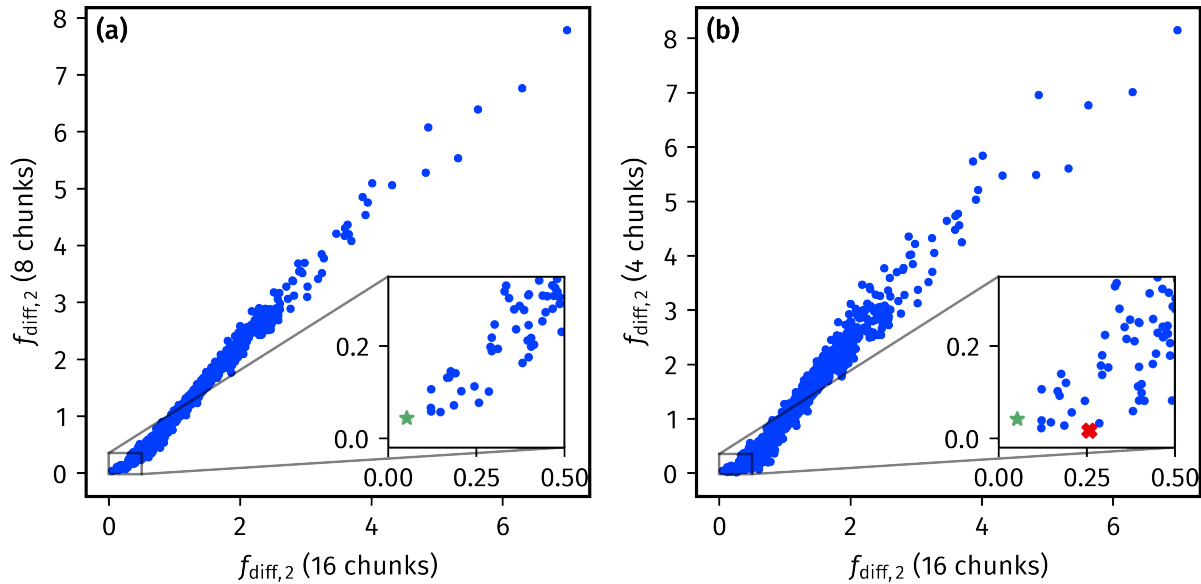


Figure 2.19: Comparison of $f_{\text{diff},2}$ values when simulated with different numbers of chunks. 16 chunks is assumed to be the ‘gold standard’. (a) Correlation between 16-chunk and 8-chunk cost functions. The ‘optimum’ identified by both cost functions is plotted as a green star in the inset. (b) Correlation between 16-chunk and 4-chunk cost functions. The ‘optima’ identified by the 16- and 4-chunk cost functions are respectively plotted as a green star and red cross in the inset.

phases: instead, phases can simply be wrapped to the region $[0, 2\pi)$, and in the simulations, the hard pulses were modelled as being instantaneous rotations, so their flip angles can also just be wrapped to $[0, 2\pi)$. (This would not be completely valid if realistic, finite pulses were used, since changing the flip angle would also change their duration.) This first optimisation yielded the following optimised parameters for the nine pulses:

$$\{\beta_i\} = \{118.6972^\circ, 29.8400^\circ, 107.4850^\circ, 190.4788^\circ, 138.4710^\circ, 144.5939^\circ, 18.9674^\circ, 73.8900^\circ, 130.6071^\circ\}; \quad (2.52)$$

$$\{\phi_i\} = \{144.5641^\circ, 173.3596^\circ, 38.9878^\circ, 146.3121^\circ, 127.7346^\circ, 7.5104^\circ, 36.9805^\circ, 110.2791^\circ, 182.3894^\circ\}. \quad (2.53)$$

To evaluate the quality of the decoupling on a real sample, the dPSYCHE experiment was performed experimentally, and the results compared against a PSYCHE experiment (fig. 2.20). Firstly, it is worth pointing out that a pure shift spectrum (if not a very good one) was obtained: this validates the form of the PSE and the optimisation approach used here, especially considering that the optimisation was not tailored towards this particular sample. The dPSYCHE experiment clearly has much greater sensitivity than the PSYCHE experiment; however, the decoupling quality is extremely poor. The reason for this is likely because the optimisation is prioritising sensitivity too highly over purity, and ultimately stems from the factor c in eq. (2.51) which

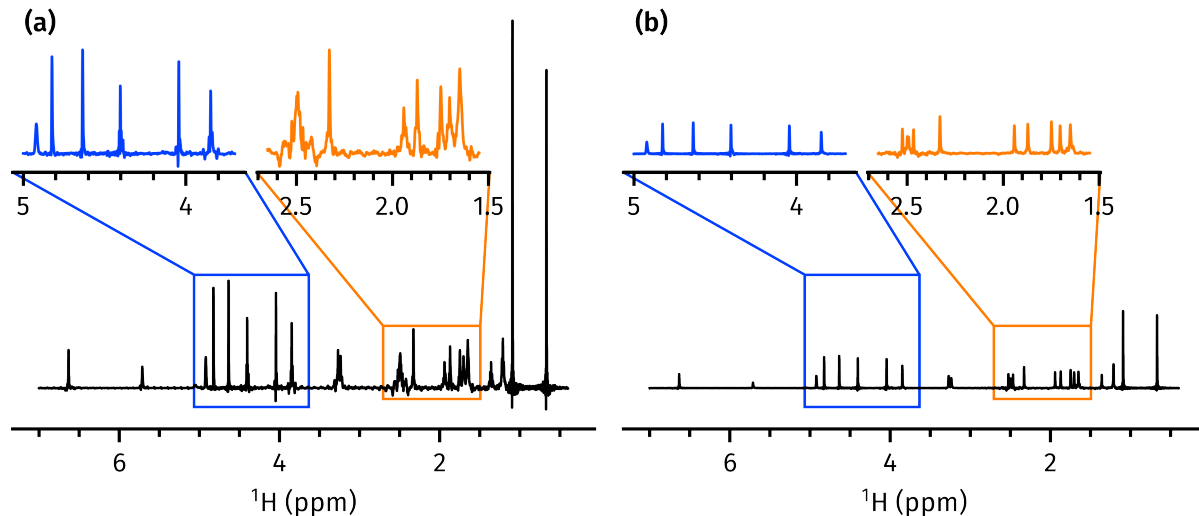


Figure 2.20: (a) dPSYCHE experiment acquired using the flip angles and phases in eqs. (2.52) and (2.53). (b) (Double-saltire) PSYCHE experiment acquired with $\beta = 20^\circ$. Spectra and insets with the same colour share the same scaling, allowing for direct sensitivity comparisons. Data code: 6A-211027.

we have ignored until now. Since $c = 1$, the optimisation is essentially guiding the dPSYCHE experiment towards having *exactly the same sensitivity as a pulse–acquire experiment*.

Although this might be a nice idea in principle, it is not physically sensible: no pure shift method has sensitivity which is even close to that of a pulse–acquire experiment. It makes more sense to scale down the target spectrum by introducing a factor of $c < 1$ into the cost function (eq. (2.51)); the parameter c therefore represents the ‘target sensitivity’. By changing the parameter c , we can control whether the optimisation emphasises sensitivity or purity more. (Note that this option was *not* available to us in the experimental JRSE-based optimisations in § 2.4.)

A series of new optimisations were therefore run, with c ranging from 0.2 to 1 (table 2.2). The 4-chunk cost function was used, and the maximum number of function evaluations capped at 5000. Each optimisation was run 10 times with a different starting seed, and the best of the 10 results chosen for further evaluation (table 2.2).

The resulting spectra (fig. 2.21) show that adjusting c has the desired effect: larger values lead to greater sensitivity and lower purity, and vice versa. In this spectrum of andrographolide, the 3.5 ppm to 5 ppm region (blue inset) is an ‘easier’ region to decouple: there are fewer couplings here and all spin systems are firmly within the weak coupling regime. The 1.5 ppm to 2.8 ppm region (orange inset) is ‘more difficult’ to decouple, especially the two peaks at 2.5 ppm, which are mutually strongly coupled. The dPSYCHE spectrum with $c = 0.4$ appeared to be a good compromise between purity and sensitivity: its sensitivity was higher than that of the original ($\beta = 20^\circ$) PSYCHE experiment. Furthermore, it provided good decoupling in the ‘easier’ deshielded region and somewhat acceptable performance in the shielded region, with the notable

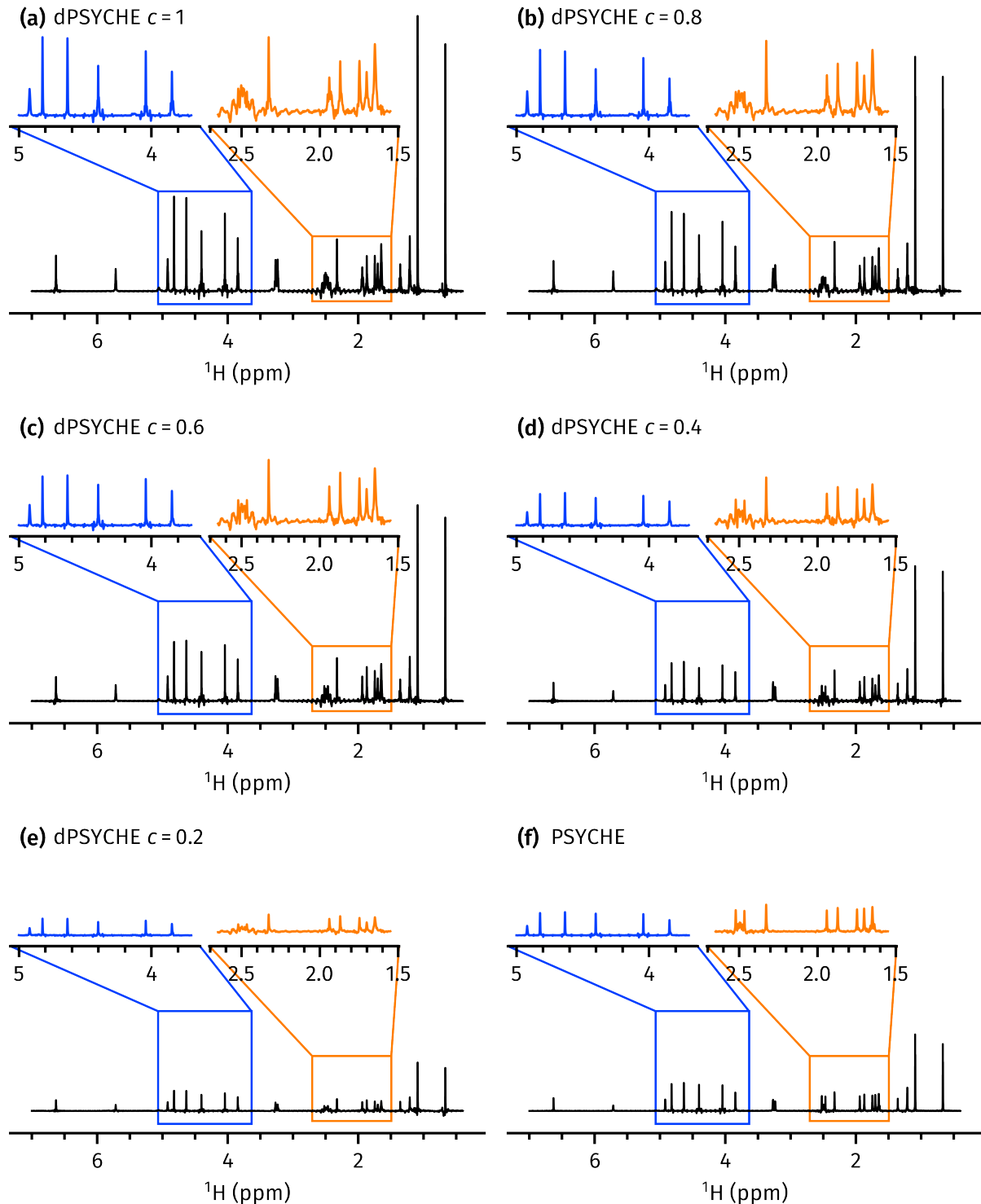


Figure 2.21: (a) dPSYCHE, $c = 1$. (b) dPSYCHE, $c = 0.8$. (c) dPSYCHE, $c = 0.6$. (d) dPSYCHE, $c = 0.4$. (e) dPSYCHE, $c = 0.2$. (f) Double-saltire PSYCHE with $\beta = 20^\circ$. Spectra and insets with the same colour share the same scaling, allowing for direct sensitivity comparisons. Data code: 6A-211231.

c	Flip angles (°)	Phases (°)	$f_{\text{diff},2}$
1	59.8813, 96.0748, 111.5862, 118.7285, 144.701, 176.9773, 29.2866, 40.0658, 63.8237	355.773, 81.741, 99.8752, 81.6675, 337.0056, 150.7004, 105.9666, 317.9979, 27.7897	0.0404
0.8	81.9461, 65.311, 90.3488, 106.6388, 73.3462, 196.7857, 51.791, 34.8188, 53.9506	329.8571, 60.3564, 137.3929, 68.5286, 340.74, 126.29, 64.4601, 145.4714, 27.3019	0.0321
0.6	53.1313, 82.2547, 88.6093, 161.9291, 83.462, 161.3548, 14.2976, 66.2199, 53.9709	347.2835, 59.5518, 68.072, 113.5358, 34.5537, 156.191, 268.7641, 302.7427, 16.29	0.0250
0.4	77.2998, 127.7274, 87.6663, 104.6371, 99.6474, 168.8171, 8.8569, 52.7865, 58.1408	342.2787, 47.7526, 76.4114, 97.3304, 14.2178, 163.9513, 94.2575, 280.0273, 16.0971	0.0271
0.2	120.6613, 107.6712, 84.0427, 133.5377, 82.1999, 223.122, 41.1617, 41.9398, 12.016	347.2051, 79.4503, 109.8796, 83.5017, 345.4284, 133.8355, 66.4118, 201.0041, 4.4419	0.0261
0.4*	92.4395, 133.2136, 38.9704, 34.9182, 56.0256, 57.4097, 31.5916, 140.1088, 128.7125	62.036, 16.411, 319.5634, 128.4443, 49.7357, 328.5103, 210.6498, 44.0123, 327.4517	0.0108

Table 2.2: Results of dPSYCHE optimisations for different sensitivities. Note that since the cost function depends on the value of c , the exact value of $f_{\text{diff},2}$ for these optimisations cannot be compared to one another; they are only presented here for completeness. The final entry, labelled with an asterisk, was run with no limit on the number of function evaluations (this is discussed further in the text).

exception of the strongly coupled peaks at 2.5 ppm.

I therefore ran a longer optimisation using the same starting point and with no limit on the number of function evaluations, which successfully lowered the cost function value by twofold (the last entry in table 2.2). The pure shift spectrum obtained using these optimised pulses is shown in fig. 2.22. The decoupling quality in both regions is comparable to that in the PSYCHE spectrum, again with the exception of the strongly coupled peaks at 2.5 ppm.

To make the dPSYCHE experiment more robust towards strong coupling, a TSE version of the sequence was also written and tested (fig. 2.23). This did indeed improve the decoupling at 2.5 ppm, as expected. However, although the TSE-dPSYCHE version still retains its *overall* sensitivity advantage over TSE-PSYCHE (especially evident in the deshielded region), this is not true of every peak: the two peaks at 1.65 and 1.70 ppm seem to have decreased intensities in the TSE-dPSYCHE experiment. The reason for this is not fully clear. One could say that the optimised pulses are unlikely to work equally well on *all* combinations of chemical shifts

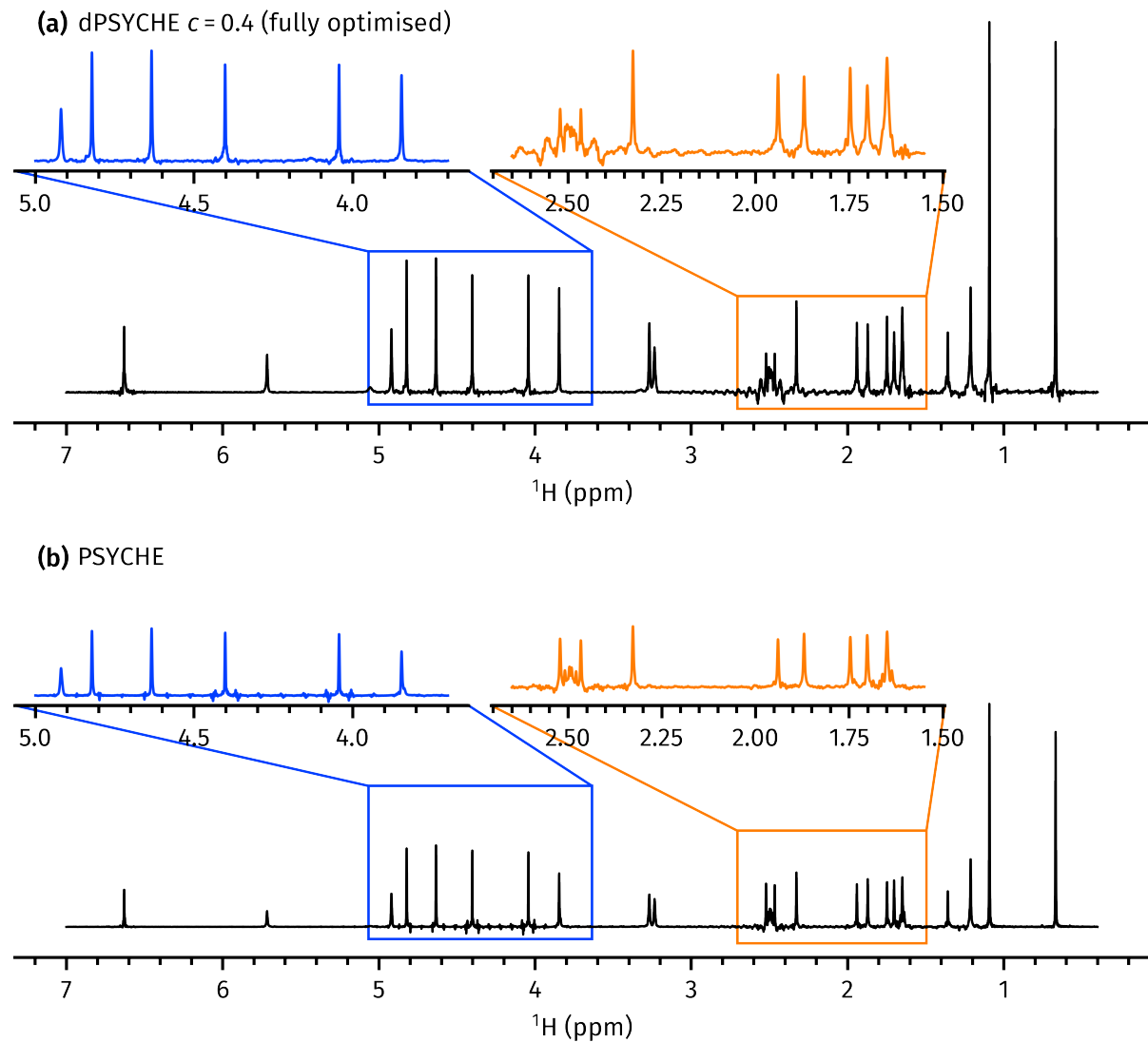


Figure 2.22: (a) Fully optimised dPSYCHE experiment with $c = 0.4$. (b) Double-saltire PSYCHE with $\beta = 20^\circ$. Spectra and insets with the same colour share the same scaling, allowing for direct sensitivity comparisons. Data code: 6A-211231.

and couplings, and these peaks may simply fall into one of the less effective regions; but in the (non-TSE) dPSYCHE experiment (fig. 2.22a), these distortions in intensity are not observed. One possible conclusion is that some of the intensity in the non-TSE experiment in fact stems from artefacts which are not fully suppressed: this may also explain the slight broadening at the base of the peaks in fig. 2.22a.

State-transfer cost function

The optimisations conducted so far have only used the simulated pure shift spectrum as an input to the cost function. However, since these optimisations are conducted in a purely theoretical manner, it is possible to directly calculate the effect of the PSE on each input state to determine

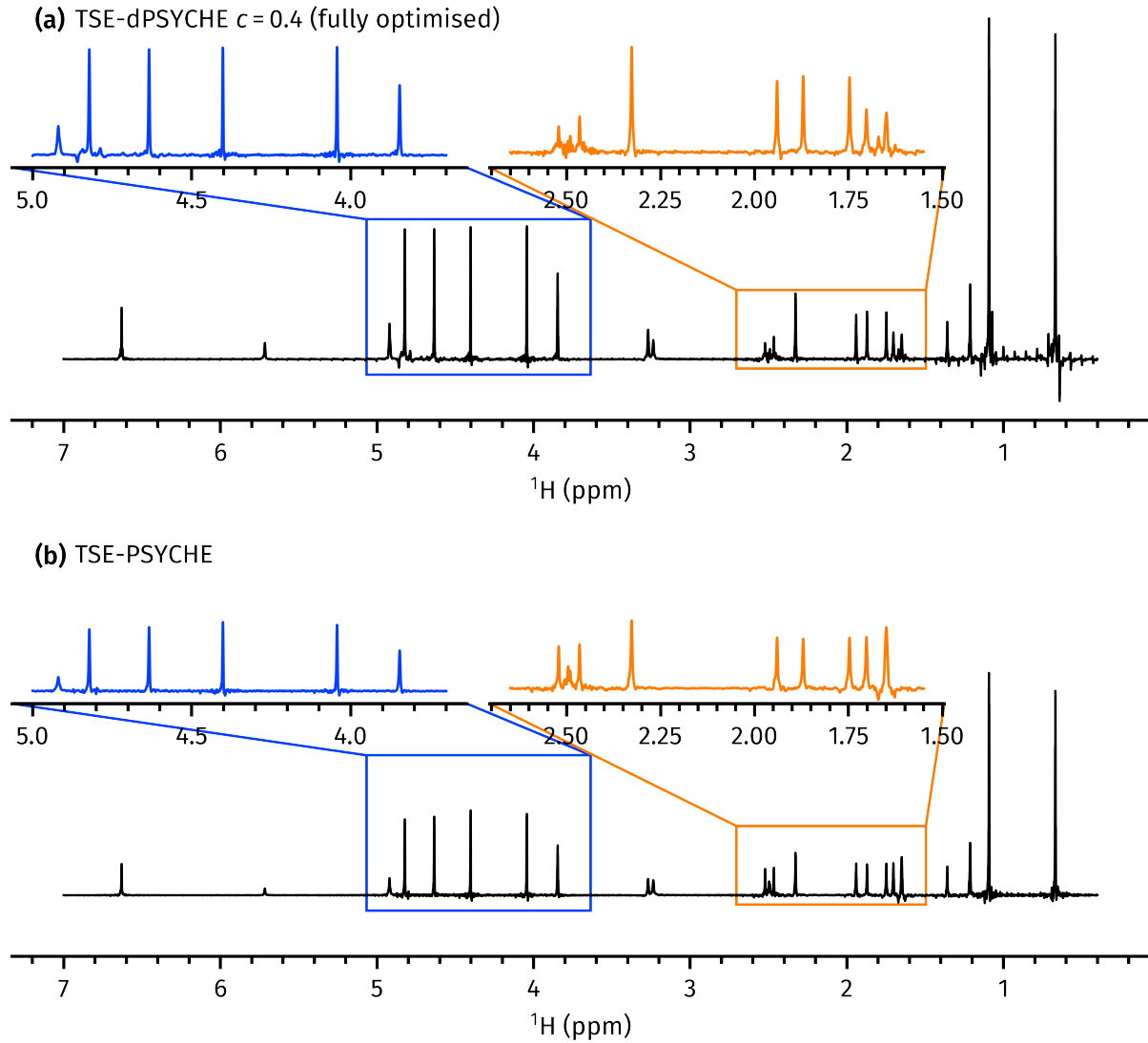


Figure 2.23: (a) Fully optimised TSE-dPSYCHE experiment with $c = 0.4$. (b) Double-saltire TSE-PSYCHE with $\beta = 20^\circ$. Spectra and insets with the same colour share the same scaling, allowing for direct sensitivity comparisons. Data code: 7A-220129.

its quality. The possible state transfers caused by the PSE can be categorised as follows:

1. desired (e.g. $I_{1+}I_{2\alpha} \rightarrow I_{1-}I_{2\alpha}$);
2. undesired, but can be dephased by CTP gradients (e.g. $I_{1+}I_{2\alpha} \rightarrow I_{1+}I_{2\alpha}$);
3. undesired, but can be suppressed by the TSE element (e.g. $I_{1+}I_{2\alpha} \rightarrow I_{1\alpha}I_{2-}$);
4. undesired, but survive even with the TSE element (e.g. $I_{1+}I_{2\alpha} \rightarrow I_{1-}I_{2\beta}$).

We want to maximise the degree of desired state transfer, item (1), which leads to the pure shift signal; and minimise the degree of unwanted state transfers (which lead to artefacts). For a TSE-dPSYCHE experiment, the only relevant artefacts are contained in item (4); without the TSE addition, we would additionally need to take item (3) into account. These state transfer

coefficients would also have to be summed (or averaged) over all possible single-quantum input states: for an n -spin system, there are $n \cdot 2^{n-1}$ possible input states (n possible spins which could be active, and 2^{n-1} permutations of passive spin states). The cost function would then be constructed as a linear combination of the relevant terms, where the weightings serve a similar function to the parameter c in the spectrum-comparison cost function $f_{\text{diff},2}$ (eq. (2.51)), in that they can be tweaked to control the balance between sensitivity and purity.

Entry	Description	Signal	All artefacts	TSE artefacts
1	dPSYCHE, $c = 1$	0.1544	0.1906	0.1126
2	dPSYCHE, $c = 0.8$	0.1417	0.1313	0.0707
3	dPSYCHE, $c = 0.6$	0.1226	0.1016	0.0359
4	dPSYCHE, $c = 0.4$	0.0869	0.0561	0.0171
5	dPSYCHE, $c = 0.2$	0.0443	0.0315	0.0087
6	dPSYCHE, $c = 0.4$ (long optimisation)	0.0915	0.0282	0.0119
7	PSYCHE, 20° double saltire	0.0420	0.0083	0.0040
7	PSYCHE, 34° double saltire	0.0918	0.0342	0.0249

Table 2.3: Evaluation of previous optima using the state-transfer cost functions. One PSYCHE experiment is included for comparison. ‘Signal’ refers to the coefficient for the desired state transfer; ‘all artefact’ the average coefficient for all undesired state transfers; and ‘TSE artefact’ the average coefficient for all undesired state transfers which are not suppressed by the TSE sequence. These are further averaged over all possible input states with coherence order +1. A three-spin system with couplings of 7 Hz and 10 Hz was used for this calculation.

Table 2.3 shows the values of each of these components for all of the optima obtained so far. Entries 1 through 5 show the effect of varying c : as expected, decreasing c leads to less signal and also fewer artefacts. These are consistent with the spectra shown in fig. 2.21. Unsurprisingly, the ‘long’ optimisation (entry 6) also performs better in all respects than the corresponding ‘short’ optimisation with $c = 0.4$ (entry 4). Finally, it is worth comparing the best dPSYCHE result (the $c = 0.4$ long optimisation, entry 6) with the standard PSYCHE method (entries 7 and 8). When PSYCHE is run with a ‘normal’ flip angle of 20° , the purity is far superior to any of the dPSYCHE experiments. However, the sensitivity is only around half that of the dPSYCHE. In order to accomplish the same sensitivity as the dPSYCHE, the PSYCHE flip angle must be increased to approximately 34° ; in this case, the artefact intensity is greater than that of the dPSYCHE experiment.

Unfortunately, even though all of the code for efficient (parallelised) computation of this cost function was written, I did not have the time to actually perform optimisations using this cost function and to experimentally validate the results. This is a slight disappointment to me, as the initial results shown in this section have been quite promising, and it is likely that the state transfer-based optimisations would have improved this even further. However, even from just the

results obtained so far, it is clear that the interleaved pulses and gradients used in the dPSYCHE PSE form a reasonable *ansatz* which can yield pure shift spectra which are comparable to the PSYCHE method.

2.7 Ultrafast PSYCHE-iDOSY

In the final section of this chapter, I turn to something entirely different: the combination of pure shift diffusion NMR with ultrafast acquisition. This work was done in collaboration with Jean-Nicolas Dumez (University of Nantes): the original project idea was first conceived and implemented there. Unfortunately, the results we obtained at Oxford were not promising enough to drive the project further, especially in light of time constraints. However, for the sake of completeness, I describe the overall concept as well as these results in this section.

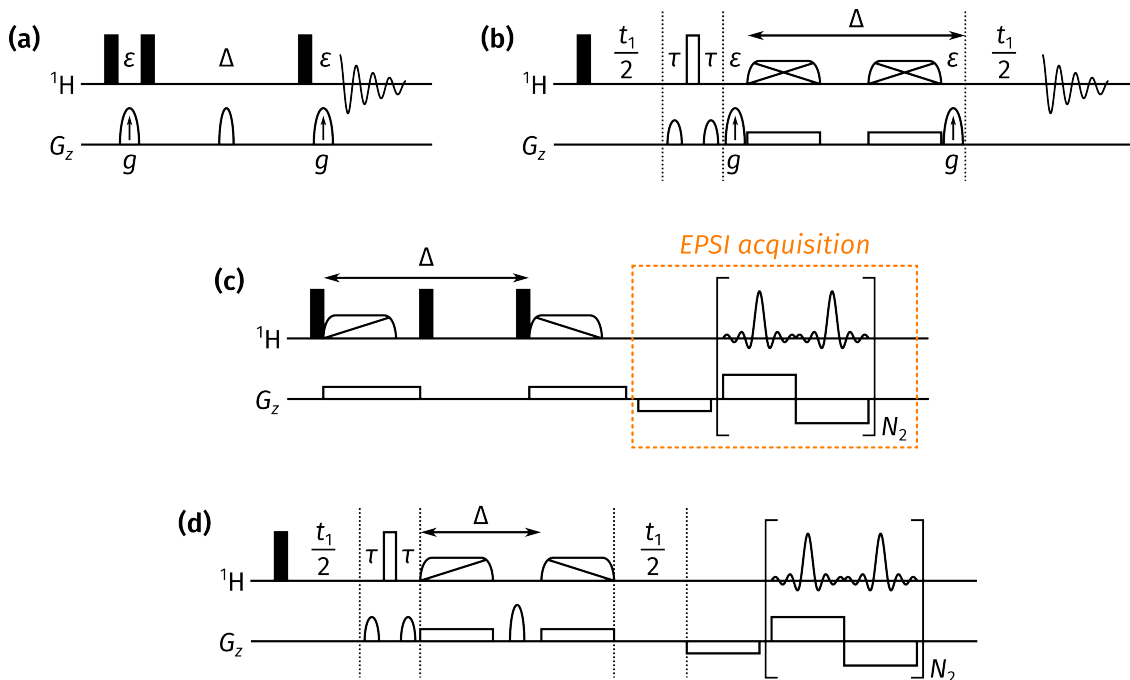


Figure 2.24: (a) Basic stimulated echo DOSY experiment. To acquire the diffusion dimension, the amplitudes of gradients with arrows inside are incremented. Δ indicates the diffusion delay. (b) PSYCHE-iDOSY experiment. τ is set to $1/(4 \cdot T_{\text{chunk}})$, as before. (c) Ultrafast DOSY experiment. The EPSI acquisition block is highlighted in orange, consisting of a prephasing gradient followed by detection in the presence of alternating gradients. (d) Ultrafast PSYCHE-iDOSY experiment. Note the use of unidirectional chirps rather than saltire pulses.

I begin with a brief overview of diffusion NMR;⁸¹ this topic will also be covered in some detail in § 3.4.10. Figure 2.24a shows a typical 2D stimulated echo DOSY experiment, where the indirect (diffusion) dimension involves the incrementation of gradient amplitudes rather than an evolution delay. For each gradient amplitude, one time-domain signal is recorded. Molecular

diffusion during a delay placed between a pair of gradients leads to attenuation of the signal, as the spatially-dependent phase imparted by the first gradient is not perfectly refocused by the second. This attenuation is described by the Stejskal–Tanner equation,^{82,83} which in its simplest form is:

$$I(G) = I(0) \exp [-(\gamma \delta G)^2 D \Delta']. \quad (2.54)$$

Here, $I(G)$ represents the signal intensity when measured with a gradient amplitude of G ; γ is the magnetogyric ratio, δ is the duration of the bracketing gradients, D the diffusion coefficient, and Δ' a ‘corrected’ diffusion delay. The exact correction required depends on the shapes of the gradients used. By measuring the signal intensity I for at least two different values of G , the diffusion coefficient D can be estimated.*

The fact that different molecules have different diffusion coefficients, and thus different attenuation profiles, can be used to separate mixtures of molecules through the basic DOSY experiment. However, when signals from different species overlap in the ^1H spectrum (as is often the case in mixtures), it is not possible to cleanly extract peak intensities, resulting in poor resolution in the diffusion dimension. There are several ways to solve this issue: for example, peaks can be resolved using another chemical shift dimension in a 3D experiment.^{84–88} A slightly less time-consuming option is to use pure shift NMR to improve the resolution. The first pure shift diffusion experiments used either the 45° projection of a 2DJ spectrum,⁸⁹ or the Zanger–Sterk PSE:^{29,90,91} in both cases, improvements in the resolution of diffusion coefficients were reported. The PSYCHE-iDOSY experiment⁵⁷ (fig. 2.24b) improved upon these, much in the same way that PSYCHE itself improved on existing pure shift methods. The ‘i’ refers to the fact that the diffusion delay is *internal* to the sequence, i.e. it is simply added in the middle of the PSYCHE element, which (using the instantaneous flip approximation) can itself be thought of as a spatially parallel stimulated echo. This avoids the need to add an entirely separate stimulated echo at the beginning or end of the sequence.

While the addition of a pure shift dimension is less expensive compared to a full chemical shift dimension in terms of time, the fact remains that the PSYCHE-iDOSY is a (pseudo)-3D experiment. One way to shorten this is to use spatial encoding techniques to collapse one dimension, specifically the diffusion dimension.⁹² Two separate steps are required for this: firstly, a gradient whose amplitude varies across the sample must be applied (the ‘encoding’ step), and then an imaging acquisition technique must be used to read out the signal in each slice of the sample (the ‘detection’ step). In the context of ‘standard’ 2D DOSY, this was first done by Loening et al.,⁹³ using the field generated by a z^2 shim coil for spatial encoding. The readout was then performed by simply acquiring the FID in the presence of a weak gradient, such that the peak

*As will be discussed in § 3.4.10, this should really be an *apparent* diffusion coefficient as there are various other factors which can affect the intensity, notably convection.

shapes directly reflect the distribution of signal intensities across the sample. Later, Thrippleton et al.⁹⁴ introduced the (by now familiar) chirp-gradient combination for spatial encoding: since spins in different slices are inverted at different times, the total gradient area applied in each slice is different. In later work by the groups of Frydman⁹⁵ and Dumez,^{96,97} the spatial encoding of the diffusion attenuation is still accomplished using the very same chirp-gradient combination. However, the detection step is accomplished using the echo-planar spectroscopic imaging (EPSI) acquisition technique,^{98,99} much like in ultrafast 2D NMR (fig. 2.24c).^{58–60,62,100–102}

Unsurprisingly, the aim in this section is to unite the PSYCHE-iDOSY and ultrafast DOSY techniques to form an ultrafast PSYCHE-iDOSY experiment. This would yield PSYCHE-iDOSY spectra in a much reduced time, at the cost of sensitivity. The resulting pulse sequence is shown in fig. 2.24d. The EPSI acquisition is used to collapse the diffusion dimension, not the PSYCHE chunking dimension.* One benefit of this is that only one PSYCHE chunk needs to be acquired at a time, circumventing the need for long EPSI detection periods (which can cause spectrometer damage). Note also that it is mandatory to use unidirectional chirps in the PSYCHE PSE: in the original PSYCHE-iDOSY, these pulses were used only for the purpose of homonuclear decoupling, and thus saltire pulses were acceptable (and indeed, superior). However, in the ultrafast version, the PSE is also responsible for creating the requisite spatial encoding of gradient areas. Using saltire pulses here would lead to an undesired ‘double’ encoding, yielding data which cannot be processed correctly.

The sequence itself was written and evaluated by Corentin Jacquemmoz and Jean-Nicolas Dumez at the University of Nantes, using a concentrated (1 M) sample of ethanol in D₂O. Their results are shown in figs. 2.25a to 2.25c. When I later tried to reproduce these results, I additionally used the POISE routine described in § 3.4.6 to optimise the amplitude ratio between the positive and negative gradients during the EPSI acquisition (without optimisation, the results obtained were even poorer); my results are shown in figs. 2.25d to 2.25f. The data processing used to generate these figures was written by me,[†] and broadly consists of the following steps: reshaping of the EPSI raw data, concatenation of PSYCHE chunks, and Fourier transformation in both dimensions.[‡] For simplicity, the data points acquired during the negative gradients is discarded.

In principle, the Fourier transformed (z, δ) plots should reveal a diffusion profile similar to that in Guduff et al.⁹⁶ This is the case for the Nantes data; in fact, it was possible to process this further

*The other option would require an ultrafast PSYCHE experiment, which—to date—has not been developed.

[†]The Nantes team have a more complete Matlab package for this, which goes several steps further than that shown here and includes features such as phase correction and fitting of the diffusion profiles to various modified Stejskal–Tanner equations. However, these were not necessary here as the project never got to that stage, and I preferred to use Python.

[‡]As discussed in Frydman et al.,⁶⁰ the k -domain in ultrafast NMR is directly proportional to the indirect-dimension frequency domain. So, to obtain the diffusion profile—which varies with space, i.e. z —a Fourier transform along this axis is required.

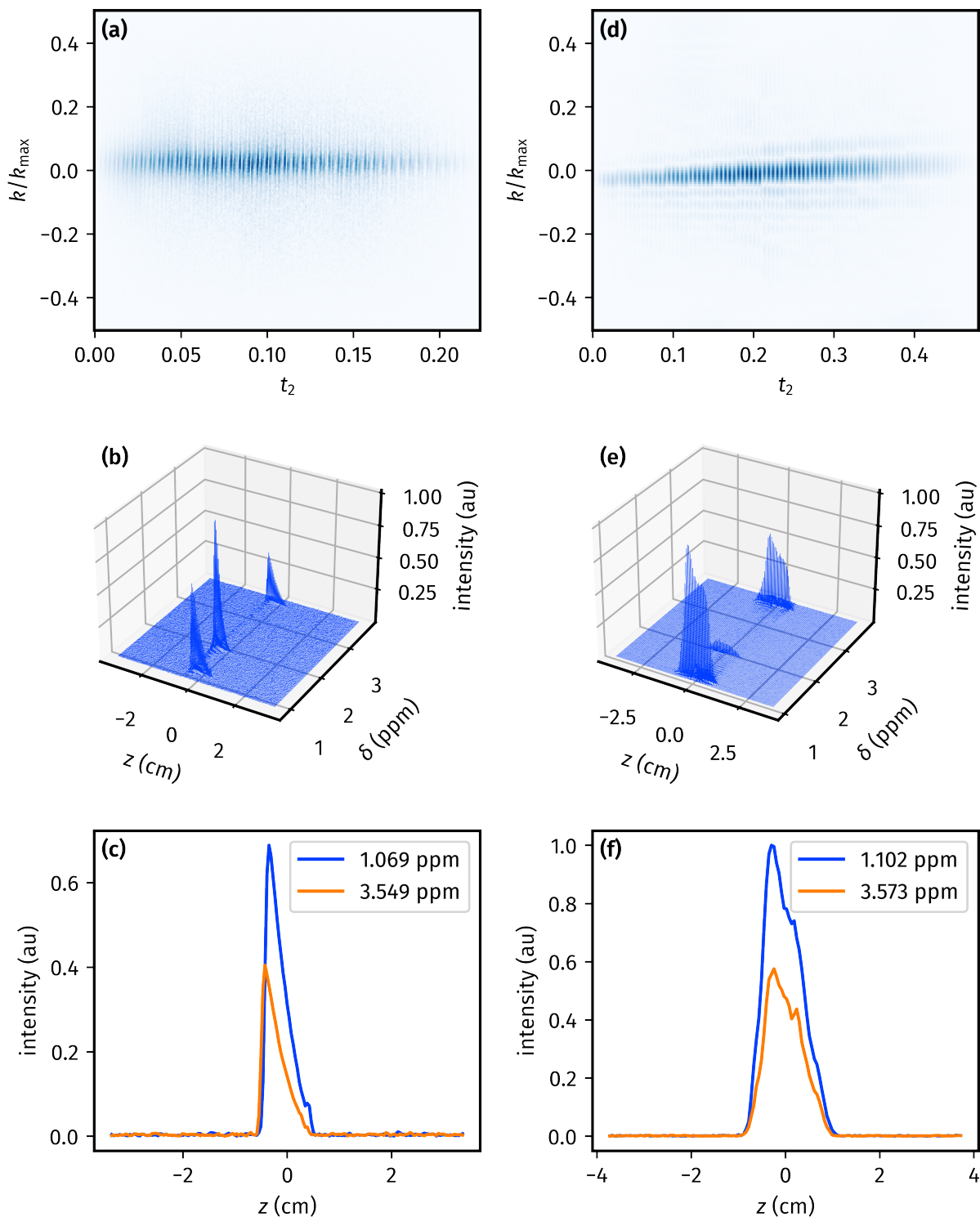


Figure 2.25: (a)–(c) EPSI PSYCHE-iDOSY data obtained by Corentin Jacquemmoz and Jean-Nicolas Dumez at the University of Nantes. The three plots respectively show the 2D raw (k, t_2) data (after PSYCHE processing has been carried out); the Fourier transformed (z, δ) data; and the individual diffusion profiles for each peak. (d)–(f) The equivalent data which I recorded in Oxford. The central peak corresponds to HDO; the Oxford sample used was drier and thus it is less intense in (e) compared to (b). Data code: 6E-210426.

to obtain a conventional 2D DOSY spectrum, where the diffusion coefficient for each component is extracted and plotted. For the Oxford data, although the general *form* of the curve is correct (in that there is a generally decreasing pattern), there are clearly ‘spikes’ in the (z, δ) diffusion profiles.

My current best hypothesis is that these spikes are artefacts from unwanted CTPs which are ordinarily suppressed through spatial averaging, but are instead being refocused by the gradients during the EPSI acquisition. This is supported by the fact that the Nantes data were acquired on a microimaging probe equipped with triaxial gradients, which allowed CTP gradients to be applied along the x - and y -axes (note that the PSYCHE gradient must still be applied along the z -axis). Naturally, any artefacts dephased by these cannot be refocused by the z -gradients in the EPSI block. On the other hand, the Oxford data were acquired on a Prodigy cryoprobe with only z -gradients.

The intensity of these ‘spike’ artefacts can, in fact, be controlled by varying other acquisition parameters. For example, increasing the EPSI gradient amplitude leads to increased artefacts (figs. 2.26a and 2.26b): this lends credence to the hypothesis above that the artefacts arise from refocused CTPs. Notice also, in fig. 2.26a, the appearance of signals at $k \neq 0$: this should not happen because in ultrafast experiments, the k -axis is proportional to the evolution frequency as a function of z (i.e. indirect-dimension frequencies in a typical 2D experiment). In a diffusion experiment, however, there is no frequency modulation in the indirect or spatial dimension, only amplitude modulation arising from diffusion attenuation; so, all signals should fall along $k = 0$. In principle, the EPSI gradient could instead be reduced, but this has the drawback of lowering the resolution in the diffusion dimension. Alternatively, when the PSYCHE gradient is increased, the artefacts seem to be reduced (figs. 2.26c and 2.26d). However, this cannot be varied too much as the PSYCHE gradients are also responsible for the diffusion encoding (the value of G in the Stejskal–Tanner equation is directly related to the PSYCHE gradient).

Another possible reason for the artefacts was a potentially poorer B_1 homogeneity on the Oxford instrument. However, a simple pulse–EPSI experiment (fig. 2.27) shows that the B_1 spatial profile, though not perfect, is relatively uniform.

Curiously, fig. 2.25d shows that in the Oxford data, there is a progressive ‘shifting’ along the k -axis in each PSYCHE chunk. (The POISE optimisation in § 3.4.6 was used to correct for the drifting *within* each chunk, but cannot be applied to the overall drift after the concatenation of chunks.) One possible explanation for this is perhaps lingering effects from gradients applied during the pulse sequence, which have more time to dissipate when t_1 is longer. This, however, does not have an appreciable impact on the data: it can be crudely corrected by circularly shifting the data along the k -axis by an appropriate amount, and Fourier transformation of the resulting data yielded effectively the same results.

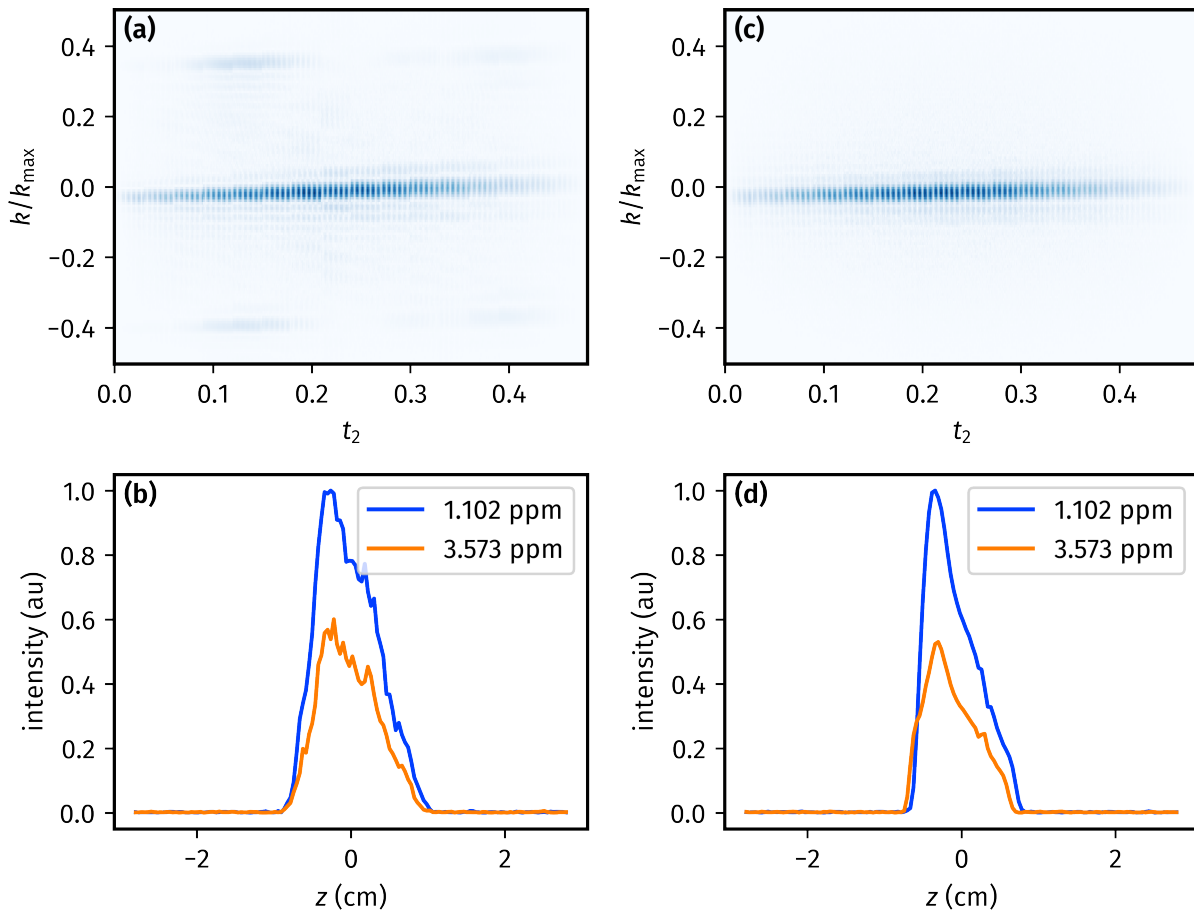


Figure 2.26: (a)–(b) The same as in figs. 2.25d and 2.25f, but with the EPSI acquisition gradient increased to 32% (from 24% in the original). *Data code:* 6E-210426. (c)–(d) The same as in figs. 2.25d and 2.25f, but with the PSYCHE gradient increased to 5% (from 3% in the original). *Data code:* 6E-210505.

All in all, it appears that the fundamental idea behind the ultrafast PSYCHE-iDOSY experiment is sound: the Nantes data form an excellent proof-of-principle. However, the implementation of the sequence almost certainly needs to be optimised in order for high-quality data to be extracted. An important subsequent step would then be to derive the appropriate form of the Stejskal–Tanner equation for extracting diffusion coefficients from the data. Unfortunately, I simply did not have the time to pursue this in any further detail.

2.8 Conclusion

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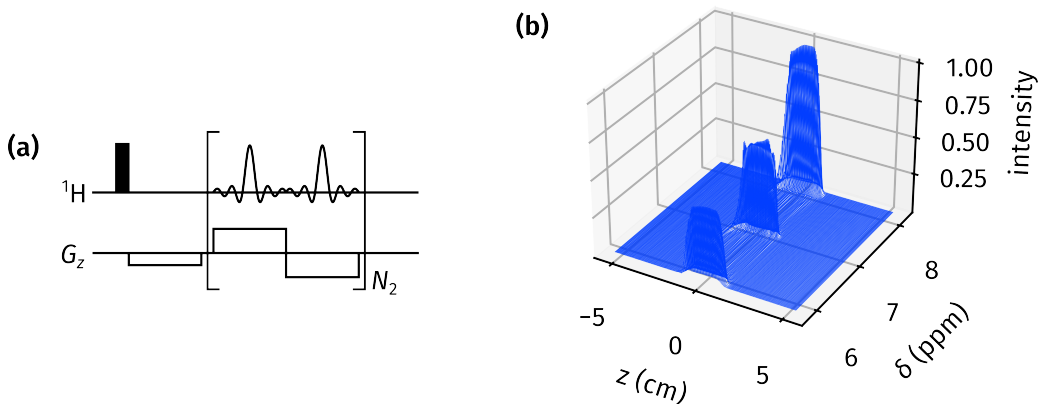


Figure 2.27: (a) Pulse-EPSI pulse sequence. The variation of the peak intensity along the z -axis directly corresponds to the B_1 spatial profile. (b) The resulting data after Fourier transformation in both dimensions ($k \leftrightarrow z$ and $t_2 \leftrightarrow \delta$). Data code: 6E-210426.

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Chapter 3

POISE

'And have you never known the pleasure and triumph of a lucky guess?—I pity you.—I thought you cleverer—for, depend upon it a lucky guess is never merely luck. There is always some talent in it.'

— JANE AUSTEN, *Emma*

This chapter describes the development of a piece of software for on-the-fly optimisation of NMR experimental parameters, titled POISE (*Parameter Optimisation by Iterative Spectral Evaluation*). POISE is primarily written in Python 3, and includes an interface to Bruker's TopSpin, allowing optimisations to be started directly from the TopSpin command line. Through this optimisation process, NMR acquisition parameters can be adjusted for individual spectrometers and samples, ultimately leading to improved spectral quality.

In this chapter, I first provide some details about the implementation of POISE. The bulk of the text which follows is devoted to a number of applications in solution-state NMR spectroscopy. At the end, the extension of the concept of on-the-fly optimisation to ESR spectroscopy is also briefly discussed: I contributed code for this, but the experimental ESR work and data analysis were carried out by Jean-Baptiste Verstraete and William Myers (University of Oxford).

The work in this chapter forms the subject of two publications:

- Yong, J. R. J.; Foroozandeh, M. On-the-Fly, Sample-Tailored Optimization of NMR Experiments. *Anal. Chem.* **2021**, 93, 10735–10739, DOI: [10.1021/acs.analchem.1c01767](https://doi.org/10.1021/acs.analchem.1c01767)
- Verstraete, J.-B.; Yong, J. R. J.; Goodwin, D. L.; Myers, W. K.; Foroozandeh, M. Towards fully optimised and automated ESR spectroscopy. *Chem. Commun.* **2022**, 58, 10715–10718, DOI: [10.1039/d2cc02742a](https://doi.org/10.1039/d2cc02742a)

3.1 Introduction

In the previous chapter, I covered various approaches to improving pure shift NMR through the use of optimisation. Although the optimisation code written there was highly specialised and only designed to work on pure shift applications, it was envisioned that this optimisation approach could be applied to essentially *any* NMR experiment which required parameter optimisation. This qualification is technically true of *every* experiment: even the simplest pulse–acquire experiment can be optimised for maximal SNR per unit time through the use of Ernst angle excitation. More complex examples, such as 2D experiments, typically have parameters which should be chosen to optimally match coupling constants (e.g. the INEPT delay, fig. 1.3) or relaxation rates (e.g. the NOE mixing time).

In routine NMR spectroscopy, this requirement for accurate parameters is often ‘solved’ through the use of compromise values, which typically fall in the middle of an expected range for typical molecules. For example, the INEPT delay may be optimised for a ${}^1J_{\text{CH}}$ value of 145 Hz, which is reasonably close to most actual ${}^1J_{\text{CH}}$ constants for sp^3 and sp^2 carbons. As an alternative, parameter values may be optimised ‘by hand’ prior to acquisition. Such a procedure is often recommended for more complicated experiments such as solvent suppression, diffusion NMR or saturation transfer difference experiments. However, compared to these, the use of experimental optimisation has several benefits. It is:

1. *sample-specific*, and as long as the default values are within the optimisation bounds, the optimisation will yield performance which is no worse than the defaults;
2. more *robust* towards unusual molecular structures, which have physical or chemical properties which fall outside of an expected range;
3. *instrument-specific*, so can compensate for spectrometer imperfections;
4. *automated*, so does not require an expert to adjust parameter values manually, or even any user intervention for that matter;
5. *objective*, in that the quality of a spectrum can (in principle) be mathematically measured through a cost function; and
6. *fast*, in that it uses an algorithm which is designed to achieve rapid decreases in the objective function: many ‘manual’ optimisations involve either trial-and-error or an exhaustive grid search (i.e. increasing a parameter value one step at a time), neither of which are efficient.

Despite these advantages, experimental optimisation of NMR parameters has seen only limited use. In fact, although there are several examples of such optimisations in laser,³ nuclear quadrupole resonance,^{4–6} and ESR⁷ spectroscopies, the only direct parallel in NMR which I

am aware of is that of the eDUMBO pulses for heteronuclear^{8,9} and homonuclear dipolar¹⁰ decoupling in solid-state magic-angle spinning experiments. In this work, the Emsley group used ‘direct spectral optimisation’ to determine the best coefficients for a pulse expressed as a Fourier series. The performance of these pulses was measured by a cost function which primarily took into account the intensity of the detected peaks: a larger intensity corresponds to better decoupling performance. Interestingly, the aim of using an experimental optimisation here was not to obtain sample-specific pulses (point (1)), but rather to account for the ‘spectrometer response’, i.e. instrumental non-idealities (point (3)). It was assumed that the compound used for the optimisation was a suitably representative choice, so that the optimisation result could simply be applied to other samples with no change.

The likely reason for the low popularity of experimental optimisations is *time*. Each *function evaluation* (FE), i.e. each measurement of the cost function, corresponds to the acquisition of an NMR spectrum which may take seconds to minutes. Thus, where possible, it is easier to run NMR optimisations in a *theoretical* manner, which can be much faster and also circumvents the effect of noise. Examples of such optimisations include the design of shaped pulses through optimal control theory,^{11–17} by simple parameterisation,^{18–23} or the optimisation of entire pulse sequences^{24–28} (the dPSYCHE experiment in § 2.6 falls under this category). In fact, even the aforementioned eDUMBO pulses were not *originally* designed as an experimental optimisation: they are actually an enhancement of the DUMBO decoupling schemes, which were optimised using numerical simulations.²⁹

In this chapter, I aim to provide a convincing argument that experimental optimisation is not necessarily slow. In particular, I show that it is often possible to devise optimisation routines which yield improved results in a matter of minutes. All the optimisations here are performed using a software package written by me, called POISE (Parameter Optimisation by Iterative Spectral Evaluation). POISE is open-source (<https://github.com/foroozandehgroup/nmrpoise>) and can be installed in a single step through `pip install nmrpoise`. Furthermore, it comes with extensive user documentation, both in the form of a text guide (<https://foroozandehgroup.github.io/nmrpoise>) as well as video (<https://www.youtube.com/watch?v=QTCeSCRZs4I>).

In contrast to previous work, which typically feature optimisations targeted at one specific application, I have endeavoured to make POISE as customisable and as broad as possible. Although it is just a single piece of software, this generality is what allows POISE to perform all the optimisations described in this chapter. Furthermore, it also means that other users can devise specific cost functions and optimisation procedures for their own use which can also be run using POISE. Thus, *POISE is more than just the applications shown later in this chapter*: it is really a platform which makes it possible to carry out arbitrary optimisations on an NMR spectrometer.

3.2 Technical overview

In this section, I first cover the general principles underlying, and the implementation of, POISE. The basic operation of POISE is summarised in the flowchart of fig. 3.1: this is a generalised version of the pure shift optimisations carried out in § 2.4. Almost all aspects of this can be customised by the user, as will be described.

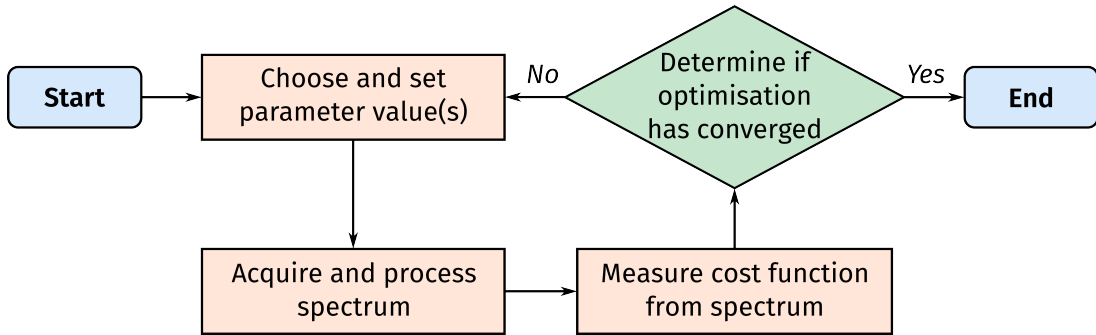


Figure 3.1: Flowchart depicting the main steps in a POISE optimisation.

I begin by drawing a distinction between an optimisation *routine*, as well as the *way* in which these routines can be run (fig. 3.2). Routines consist of a series of predefined variables, such as the parameter(s) to be optimised: however, these may be optimised in different ways, which is where the experiment used to perform the optimisation, as well as other options, come in. These ‘ingredients’ of an optimisation will be repeatedly referred to in the discussion of individual applications in § 3.4.

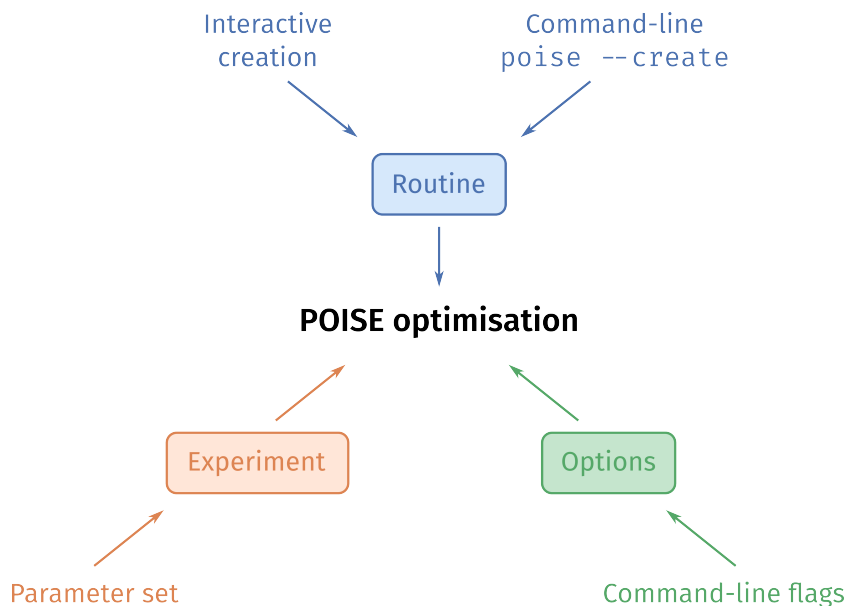


Figure 3.2: The various ‘ingredients’ of a POISE optimisation and how they can be set up.

3.2.1 Routines

An *optimisation routine*, as defined in POISE, consists of the following components:

1. *Name*

This is an identifier used to refer to the entire routine, which is arbitrary, but should ideally be descriptive.

2. *Parameters*

The parameters to be optimised. These are given as strings and directly correspond to TopSpin parameter names, for example, P1 for a pulse width.

3. *Initial guess* (one per parameter)

The point at which the optimisation is started. Naturally, this should represent the user's best guess as to where the optimum lies. It is generally sensible to choose unoptimised, 'default' values for these.

4. *Lower and upper bounds* (one each per parameter)

These may be chosen based on a range of values which are 'chemically sensible', or alternatively, instrumental specifications may sometimes restrict the range of parameter values which can be explored.

5. *Tolerances* (one per parameter)

The tolerance loosely corresponds to the level of accuracy required for the optimisation. Of course, setting the tolerance to be too large will simply yield an imprecise and meaningless result. However, there is no point in setting this to be too small (i.e. requesting an overly accurate optimum): this increases the time required for convergence, because the value of the cost function evaluated at two points too close together will differ only by noise. Furthermore, parameter values cannot be implemented with arbitrary precision on a spectrometer, and the available resolution sets a lower bound for the tolerance (see § 3.4.6 for a more concrete discussion of this). These conditions make it sound as if there is little room for error, but in practice the desired accuracy is often reasonably clear from the context, and it generally suffices to get the order of magnitude correct;

6. *AU programme*

The AU programme defined in an optimisation routine is used to acquire and process the spectrum. The user may leave this empty, in which case POISE automatically detects the dimensionality of the experiment and performs standard processing steps (Fourier transformation, window multiplication, phase correction, and baseline correction). This

is often sufficient for many applications. However, this does allow for almost infinite customisation of the actual spectral measurement: for example, the AU programme may call other scripts in TopSpin which create shaped pulses (as shown in § 3.4.8).

7. Cost function

The cost function measures the ‘badness’ of the spectrum thus recorded. As in the previous chapter, the optimisation algorithm seeks to minimise this value. POISE cost functions must be written in Python 3: this design decision is considered later in § 3.2.5. For ease of use, several cost functions which cover ‘typical’ optimisation scenarios, such as maximising or minimising signal intensity, come pre-installed with POISE. This means that users do not necessarily need to write their own cost function if they are not familiar with Python.

Collectively, a routine is therefore a description of *what is to be optimised*. POISE allows users to create new routines interactively through a series of dialog boxes. Alternatively, routines themselves can be created on-the-fly using the `poise --create` command: this is useful when some components are not known beforehand, such as if the optimum from a different optimisation is to be used as the initial point in a new one. However, this is limited to single-parameter routines.

After being created, routines are stored in the human-readable JSON format: they can therefore be modified using any text editor. Examples of these JSON files are presented in subsequent sections.

3.2.2 The experiment

Once the user has defined a (named) routine, it can then be run from the TopSpin command line using the command `poise ROUTINE_NAME`. However, the routine itself does not specify what experiment is to be run (i.e. the pulse programme), nor any of the other parameters in the experiment. These must be set by the user, and can most conveniently be stored in a TopSpin parameter set which can simply be loaded before starting the optimisation.

This flexibility means that the same *type* of optimisation may be applied to different pulse sequences: for example, an experiment to optimise the NOE mixing time (as described in § 3.4.4) can be run with different versions of the NOESY sequence depending on what is most appropriate, as long as the parameter being optimised always has the same identifier in TopSpin (D8 for the mixing time). Likewise, parameters such as the number of scans can be adjusted in order to run optimisations on samples with different concentrations, without having to redefine a new routine.

3.2.3 Optimisation options

Finally, POISE provides a few command-line options which control how the optimisation is carried out:

- the `--maxfev` option allows the user to control the maximum number of FEs, or in other words, the maximum number of experiments run. If the optimisation has not converged after acquiring this many spectra, the best result so far is simply returned. This indirectly allows a cap to be placed on the time spent on optimisation.
- the `--quiet` option silences all output from POISE. This is useful when, for example, a optimisation is to be run under automation. (The optima found are stored in the dataset itself after the optimisation ends, and can therefore always still be retrieved.)
- the `--separate` option allows each FE to be run in a new experiment number, so that the optimisation trajectory can be analysed after its conclusion.
- perhaps most importantly, the `--algorithm` option allows the user to choose one of three optimisation algorithms, which I now describe in more detail.

3.2.4 Optimisation algorithms

As was briefly mentioned in § 2.4.1, derivative-based optimisation algorithms cannot be used for experimental optimisations. To be more precise, when analytical gradients are not available, derivative-based algorithms calculate gradients using a finite difference approximation:

$$\nabla f(x) \approx \frac{f(x + \varepsilon) - f(x - \varepsilon)}{2\varepsilon}, \quad (3.1)$$

where ε is the step size used for the finite difference calculation. In Nocedal and Wright,³⁰ it is shown that the error in this finite difference gradient (as compared to the true gradient) has an upper bound of

$$\frac{\eta(x; \varepsilon)}{\varepsilon} + O(\varepsilon^2), \quad (3.2)$$

where $\eta(x; \varepsilon)$ is (roughly) the noise in the region $[x - \varepsilon, x + \varepsilon]$. The first term represents the error due to this noise, and if ε is small, this term is large; on the other hand, the second term represents the error due to the finite difference approximation, and if ε is large *this* term is large. This means that finite difference gradients, and any algorithms which use them, are entirely unreliable in the presence of noise. Instead, POISE provides a choice of three derivative-free optimisation algorithms: the Nelder–Mead (NM) method,³¹ the multidirectional search (MDS) method,^{32,33} and the Py-BOBYQA trust-region method.^{34,35}

Nelder–Mead

The NM method is a highly popular derivative-free optimisation algorithm, which maintains a set of points $\{y_1, y_2, \dots, y_{n+1}\}$ during the optimisation, n being the number of parameters being optimised (or the *dimension* of the optimisation problem). The convex hull of these points, Y , is the smallest possible set of points which contains all the y_k and is convex, in that

$$\forall x_1, x_2 \in Y, \forall \alpha \in [0, 1], \alpha x_1 + (1 - \alpha)x_2 \in Y, \quad (3.3)$$

and in this context is called a *simplex*. If this convex hull is nonempty—or equivalently, if the n vectors $y_k - y_1$ ($2 \leq k \leq n + 1$) are linearly independent—then the simplex is called *nonsingular*. (To provide an analogy for $n = 2$, the convex hull is the shape obtained by stretching a rubber band around three pins placed at the points y_1, y_2, y_3 . The simplex would be singular if the three points were collinear.)

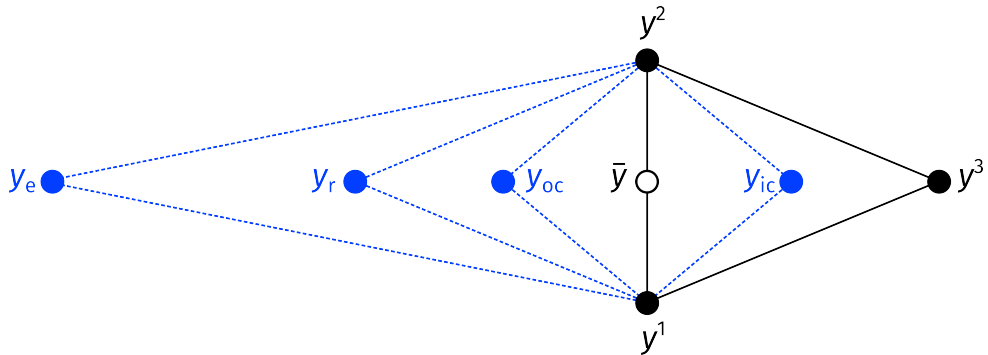


Figure 3.3: Diagram showing various points evaluated in one iteration of the Nelder–Mead algorithm (for an optimisation of two parameters). The solid black lines indicate the boundaries of the simplex, which is assumed to be ordered such that y_1 is the best point (has the lowest cost function value) and y_3 the worst. The current points in the simplex are shown as filled black circles, and the centroid \bar{y} as an empty black circle. The blue circles indicate the trial points which the algorithm attempts to replace y_3 with, and are further discussed in the text. Blue dashed lines indicate the simplex which would result if the corresponding trial point were accepted.

The NM algorithm is in fact quite intuitive to understand. The initial simplex is first constructed using the supplied initial point: POISE specifically uses the method of Spendley et al.³⁶ to construct a *regular simplex*. The optimisation itself begins by measuring the cost function f at every point of the simplex, and sorting the points in ascending order of cost function values (i.e. from best to worst), such that $f(y_1) \leq f(y_2) \leq \dots \leq f(y_{n+1})$. The centroid of the simplex is defined by the n best points:

$$\bar{y} = \sum_{i=1}^n y_i. \quad (3.4)$$

On each iteration of the NM algorithm, we attempt to replace the worst point y_{n+1} with a better

point (fig. 3.3). The search for the new point is performed in several steps: first, the worst point is *reflected* about the centroid of the simplex to obtain a new point y_r :

$$y_r = \bar{y} - (y_{n+1} - \bar{y}). \quad (3.5)$$

The value of the cost function is evaluated at this point, and is compared against the existing points in the simplex. If this reflected point falls in the middle of the pack, such that $f(y_1) \leq f(y_r) < f(y_n)$, this represents a ‘modest’ improvement in the cost function: we simply replace the worst point with this and continue to the next iteration.

On the other hand, if the reflected point is better than all the other points (i.e. $f(y_r) < f(y_1)$), then we ambitiously attempt to *expand* the simplex even further in that direction:

$$y_e = \bar{y} - 2(y_{n+1} - \bar{y}). \quad (3.6)$$

Of course, there is no guarantee that this is necessarily better than y_r ; therefore, we choose whichever point of y_r or y_e had a lower value of f , and replace the worst point with this and continue to the next iteration.

If the reflected point is an improvement on the worst point but is no better than the remaining points, in that $f(y_n) \leq f(y_r) < f(y_{n+1})$, then the algorithm performs an *outside contraction*, which resembles a half-hearted reflection:

$$y_{oc} = \bar{y} - (1/2)(y_{n+1} - \bar{y}). \quad (3.7)$$

Conversely, if the reflected point is even worse than the worst point, in that $f(y_{n+1}) \leq f(y_r)$, then this suggests that that search direction is very poor: we thus perform an *inside contraction*, which uses a point halfway between the worst point and the centroid:

$$y_{ic} = \bar{y} + (1/2)(y_{n+1} - \bar{y}). \quad (3.8)$$

If either of these contracted points are any better than y_r , then we replace the worst point in the simplex and continue to the next iteration; otherwise, we conclude that no search direction was good, and simply shrink the simplex towards the current best point by replacing each point y_k with $(y_k + y_1)/2$. In practice, these ‘last-resort’ shrink steps occur very rarely.

Finally, convergence is signalled when in every dimension of the optimisation, the width of the simplex is smaller than the chosen optimisation tolerance.* For a multiple-parameter optimisa-

*The implementation of the NM algorithm in the `scipy` library only accepts a single value for the ‘tolerance’, which is then used in all dimensions. This is designed to be used by scaling the parameters beforehand such that the tolerance in each dimension is equal (and in fact, POISE was later updated to do so). However, during initial development I chose instead to re-implement the NM algorithm with a convergence check which allowed for different tolerances to be specified in each dimension.

tion, this can potentially mean that extra accuracy is obtained in one of the parameters (because the simplex may have shrunk along that dimension more quickly). However, it does guarantee that *at least* the specified level of accuracy in every dimension is achieved.

Multidirectional search

In the preceding discussion, we noted that the simplex Y was nonsingular if the n vectors $y_k - y_1$ were linearly independent. Equivalently, the matrix M formed by concatenating these vectors

$$M = (y_2 - y_1, y_3 - y_1, \dots, y_{n+1} - y_1) \quad (3.9)$$

must be nonsingular, i.e. have a nonzero determinant. We can quantify how ‘close’ the simplex Y is to being singular, using the l^2 condition number of the matrix M , which in this context is usually referred to as the *simplex condition*:

$$\kappa(Y) = \|M\| \|M^{-1}\|, \quad (3.10)$$

where $\|M\|$ is the matrix norm induced by the Euclidean norm,

$$\|M\| = \max_{x \neq 0} \frac{\|Mx\|}{\|x\|}. \quad (3.11)$$

A singular simplex Y of course does not have a well-defined condition, since M^{-1} does not exist. However, the larger the condition of a simplex is, the closer it is to being singular. Very loosely speaking, a long and thin simplex has a large condition number, and would be singular if its width were to go to zero.

The simplex updates made in the process of the NM algorithm mean that the simplex condition changes throughout the course of the optimisation. This is good for achieving rapid decreases in the cost function, since the simplex shape adapts to the cost function being optimised. However, if the simplex condition gets too large, it is possible that the optimisation will stall at a nonstationary point, since the search directions of the simplex are severely limited. The MDS algorithm was proposed partially for the purpose of avoiding this ill-conditioning.*

The MDS method is also simplex-based, and uses similar reflection/expansion/contraction steps to the NM algorithm. However, instead of (e.g.) reflecting a single worst point y_{n+1} about the other points, it reflects *all* of the n worst points $\{y_2, y_3, \dots, y_{n+1}\}$ about the best point y_1 . This means that the shape of the simplex Y , and thus its condition $\kappa(Y)$, is always preserved: this

*The main reason was in fact to better exploit computer parallelism, but it was also noticed that the MDS method proved to be generally more robust than NM.

provides it with much better convergence properties, as was shown by Torczon et al.^{32,37*} The increased reliability of the MDS algorithm over the NM algorithm was demonstrated on a variety of example optimisation problems: even in the very simple case where the cost function was simply the norm of a vector,

$$f(y) = \|y\|, \quad (3.12)$$

it was shown that the NM algorithm stalled when the dimension of the problem, n , was sufficiently large. The value of n needed to precipitate this failure depended on the problem being solved, and generally ranged from 8 to 40. On the other hand, the MDS method proved to be robust under the same conditions, eventually converging to the optimum—although in the cases where NM *did* work, the MDS method generally required more FEs.

It was this improved robustness of the MDS algorithm which prompted Goodwin et al. to use it in their (experimental) optimisation of ESR pulse shapes,⁷ and for me to later include it in POISE. In the ESR work, the number of pulse points being optimised was 11 or 21, which fell into the regime where the MDS method would likely have better convergence properties than NM. However, in NMR, each experiment takes a substantially longer time than in ESR, and even optimisations with $n > 2$ become rather time-consuming due to the number of FEs required (the largest n explored in the present work is 4). Thus, it is unlikely that the NMR optimisations explored in POISE will lie in the regime where the NM algorithm is prone to failure. As will be shown later, I generally found that the NM and MDS methods were equally reliable in our optimisations, with NM generally being faster.

Py-BOBYQA

Unlike the NM and MDS algorithm, Py-BOBYQA is not simplex-based, but is a trust-region algorithm.^{34,35} The fundamental idea behind a (derivative-free) trust-region method is to sample the cost function at a set of points Y , and construct a model m through interpolation, which matches the cost function at these points:

$$\forall y \in Y, m(y) = f(y). \quad (3.13)$$

The model at iteration k is labelled m_k . Most trust region methods use a quadratic model, and Py-BOBYQA is no exception. This can be expressed as:

$$m_k(x_k + p) = c + g^T p + p^T G p, \quad (3.14)$$

*Specifically, it can be concluded that at least one of the search directions was bounded away from being orthogonal to the gradient; or in simpler (and less precise) terms, at least one of the search directions is close enough to a direction in which the cost function f decreases.

where G is a symmetric matrix and x_k is the centre of the model at iteration k (x_0 being the user-specified initial point). For this model to be fully determined, the set Y must therefore contain $(n + 1)(n + 2)/2$ points in total.

The algorithm further maintains a *trust region radius* Δ_k at each iteration, which is a measure of how reliable the model is. The initial trust region radius, Δ_0 , can be arbitrarily chosen: in the case of POISE, I elected to set Δ_0 to be 10 times the desired tolerance. The model m_k is then used to calculate the next step s_k , which is obtained by minimising m_k over all points within a radius of Δ_k from the centre x_k (the *trust region subproblem*):

$$s_k = \arg \min_{\|s\| \leq \Delta_k} m_k(x_k + s). \quad (3.15)$$

Since m_k itself is noiseless, this can be solved using almost any algorithm: Py-BOBYQA uses a conjugate gradient method. The cost function is then evaluated at the trial point $x_k + s_k$, and compared against the value predicted by the model. If the ratio of ‘actual improvement’ to ‘predicted improvement’ is large enough, i.e.

$$r_k = \frac{f(x_k) - f(x_k + s_k)}{m_k(x_k) - m_k(x_k + s_k)} \geq \eta \quad (3.16)$$

for some threshold value η , then the step s_k is accepted and x_{k+1} is set to $x_k + s_k$, replacing the worst point in Y . Additionally, the trust region radius Δ_k may be increased so that the next step(s) can be more ambitious. Conversely, if $r_k < \eta$, then there are one of two possibilities: either the model is poorly conditioned (in that the points in Y are very unevenly distributed), in which case one of the points is replaced and the model recalculated; or the model is sufficiently well-conditioned, in which case the step is rejected, and Δ_k is decreased.

Py-BOBYQA goes beyond a standard derivative-free trust-region algorithm in further limiting the rate at which the radius Δ_k can change (amongst others). Specifically, Py-BOBYQA also maintains a lower bound on the trust region radius ρ_k , and on unsuccessful iterations Δ_k is not allowed to decrease further than ρ_k . This prevents Δ_k from decreasing too quickly until the algorithm is certain that Y is sufficiently well-conditioned.³⁸ Another core feature of Py-BOBYQA is the implementation of multiple restarts, which endows it with greater robustness towards noise and also allows it to escape local minima.^{35,39} However, the multiple-restarts feature in Py-BOBYQA was disabled in POISE as this often led to overly long optimisations.*

Crucially, Py-BOBYQA differs from the simplex-based methods in that *it cares about the actual value of the cost function*. In the NM and MDS methods, only the relative ordering of the points

*Most mathematics papers on optimisation have no qualms in using hundreds or even thousands of FEs, and it is this context in which Py-BOBYQA outperforms other algorithms. Unfortunately for me, POISE works in an *extremely* restrictive regime where even 50 FEs would be considered very expensive.

in the simplex matters; it makes no difference to the algorithm whether the worst point has a cost function value of 10 or 1000. However, in Py-BOBYQA, the value of f is used in constructing the model, and thus directly influences the optimisation trajectory. Although this is beneficial in cases where the underlying cost function is relatively well-behaved,^{*} and is reflected in faster convergence rates for such problems, it can be problematic for some cost functions. Py-BOBYQA is set as the default optimiser in POISE, but the user is strongly recommended to try the NM method as a first step when troubleshooting optimisations.

3.2.5 Implementation details

In this section, I discuss some behind-the-scenes details about how POISE is implemented and several design choices. This information is relevant for anybody looking to improve or otherwise modify the POISE codebase.

Firstly, POISE is written in Python 3, and since TopSpin does not have a built-in Python 3 interface,[†] this means that POISE is not entirely self-contained within TopSpin: in particular, an external installation of Python 3 is required, which may be a slight inconvenience. This choice was necessary because it would have been too time-consuming to implement numerical optimisation algorithms using the existing C or Python 2 application programming interfaces (APIs) in TopSpin: notably, the Python 2 API uses the Jython implementation of the language, which is incompatible with numpy. However, a benefit of this is that since the ‘cost’ of installing Python 3 is already paid, we can also allow users to define their own cost functions using libraries such as numpy and scipy. Without these, performing any kind of non-trivial data processing is significantly more awkward.

Like most Python 3 packages, POISE is available on the Python Package Index (PyPI), so can be installed using a single command: `pip install nmrpoise`. As usual, POISE is first installed to the Python `site-packages` directory. If the `nmrpoise` package is imported from a Python 3 script, then this code is read. This may be required on occasion, as the `nmrpoise` package provides a few functions to analyse optimisation logs created by POISE. We might refer to this code as the ‘library’ component of `nmrpoise`.

This, however, is irrelevant for actually *running* optimisations. When POISE is installed, on top of the default installation to `site-packages`, it automatically searches for TopSpin installations in either `C:\` (Windows), or `/opt/` (Unix/Linux). (If necessary, a non-standard TopSpin installation

^{*}I think this means cases where the cost function is well described by a quadratic model. Of course, because of Taylor’s theorem, every non-noisy cost function can be locally described by a quadratic model within a sufficiently small region. However, for meaningful progress to be made with noisy cost functions, the model must be built over a large enough region such that noise becomes less relevant.

[†]Version 4.1.4 of TopSpin now comes with a Python 3 API; however, this was introduced too late for the work in this chapter.

location can be specified using the `$TS` environment variable.) The installation then creates:

- a *frontend script* at `$TS/exp/stan/nmr/py/user/poise.py`, which allows POISE to be invoked by simply typing `poise` in the TopSpin command line and is responsible for controlling data acquisition; as well as
- a *backend directory* at `$TS/exp/stan/nmr/py/user/poise_backend`, within which all of the POISE data and logic is stored. For example, routines can be found in the `routines` subdirectory, and cost functions in the `costfunctions.py` and `costfunctions_user.py` files.

All optimisations are run using the code *only* in the backend directory, and not anything in Python's `site-packages` folder. This is because the frontend script must know how to launch the backend (i.e. where to find the files), and it is simply easiest to predefined this location.*

```
try:
    # Launch backend
    backend = subprocess.Popen([p_python3, "-u", p_backend],
                               stdin=subprocess.PIPE,
                               stdout=subprocess.PIPE)

    # Pass information from frontend to backend
    for item in [args.algorithm, routine_id, p_spectrum, args.maxfev]:
        print >>backend.stdin, item
    backend.stdin.flush()

    while True:
        # Receive information from backend
        line = backend.stdout.readline()
```

Listing 3.1: Excerpt from the POISE frontend script, illustrating the two-way communication between frontend and backend.

Having files in two different places does mean that some form of communication between the two must be established. In POISE, this is accomplished through the use of anonymous pipes, one for each direction of communication (listing 3.1). In this way, the backend can signal to the frontend what values of parameters should be evaluated; the frontend can then begin data acquisition, and signal to the backend when this is complete so that the cost function can be calculated. Although this setup works perfectly fine when left to run untouched, a frustrating number of ‘tricks’ are required to keep these synchronised if either the frontend or the backend

*In fact, it is possible to dynamically determine the `site-packages` installation location at runtime, meaning that the entire backend does not need to be copied to TopSpin directories. However, that would mean the cost functions would be buried inside the `site-packages` directory, which can be difficult to find.

are terminated unexpectedly, or if acquisition is prematurely stopped by the user (which usually suggests that they wish to stop the optimisation). This includes the backend creating a file with its process ID every time it is called and deleting it upon exit (listing 3.2), meaning that the frontend can locate any stray backend processes which were not appropriately terminated.

```
from contextlib import contextmanager

@contextmanager
def pidfile():
    # Create a file with the PID
    pid = os.getpid()
    pid_fname = Path(__file__).parent / f".pid{pid}"
    pid_fname.touch()
    # Run the code in the 'with' block
    try:
        yield
    # Delete the file after the 'with' block is exited
    finally:
        if pid_fname.exists():
            pid_fname.unlink()

if __name__ == "__main__":
    with pidfile() as _:
        main()
```

Listing 3.2: Simplified excerpt from POISE backend script, showing a context manager used to keep track of backend process IDs. The context manager ensures that when the script is started, a file with the process ID is created; and when the script exits, this file is deleted. The ‘main()’ function carries out the actual optimisation.

Finally, there is one other quirk of TopSpin surrounding data acquisition: it is possible to start the acquisition from a Python script (such as the frontend `poise.py` script), but it is not possible to block execution of the Python script while acquisition is running. Thus, it is not possible to trigger acquisition and wait until it is done before sending a signal to the backend.* The workaround is to call an AU programme containing acquisition commands, which (somehow) blocks the Python script. Of course, this also means that the AU programme can be extended to include other commands, which is yet another way in which optimisations can be customised. During optimisations, the frontend must also be careful not to overwrite other experiments: this can easily happen if, for example, a user opens a new dataset in TopSpin just before acquisition is started. To ensure that this is the case, the frontend *always* brings the optimisation dataset to the foreground immediately before the acquisition AU programme is executed. This has a slight

*The TopSpin Python documentation claims that this *can* be accomplished using, for example `XCMD("zg", wait=WAIT_TILL_DONE)`. However, none of the suggestions in the documentation worked as intended.

drawback in that it can be difficult to view other spectra in TopSpin while an optimisation is proceeding. (Unfortunately, this is out of my control: TopSpin does not provide any documented way of running an acquisition AU programme on a background dataset.)

3.3 What POISE is not

Before moving on to cover applications of POISE, I want to make a note about several limitations of the approach chosen.

Firstly, *POISE is not specialised*. Generality is a strength in that POISE can be applied to a diverse range of NMR experiments. However, it can also be a weakness, because the capabilities of POISE must be restricted such that it can be adapted to different scenarios. POISE *always* follows the framework in fig. 3.1: in particular, it simply seeks to find the optimum \mathbf{x}^* , defined by

$$\arg \min_{\mathbf{x}} f(\mathbf{x}). \quad (3.17)$$

Although this conforms to the mathematical notion of an optimisation, there are many other ways of searching for ideal NMR parameters, such as fitting data to a model or directly reading parameters off a spectrum.

This rigidity in the underlying logic means that it is very conceivable that in specific instances, specialised optimisation routines which use customised strategies for data acquisition and analysis *can* outperform POISE in terms of speed and/or accuracy. For example, we see this in § 3.4.1: the TopSpin `pulsecal` routine for pulse width calibration can be much faster than POISE, because it only needs to perform one experiment to obtain an answer. I particularly want to distinguish POISE from other types of ‘optimisations’ reported in the literature, which typically *accumulate* data points until a given threshold is reached (in terms of, say, SNR or confidence in a parameter). Examples of such procedures can be found in the contexts of relaxation measurements^{40,41} and undersampling in multidimensional NMR.^{42–44}

Secondly, *POISE is not a global optimiser*. The optimisation algorithms provided within POISE are not designed to go beyond the first local minima found (except for Py-BOBYQA, but as described in § 3.2.4, I disabled the multiple restarts option responsible for this). In challenging optimisation cases where multiple local minima exist, it is not generally possible to predict which local minimum the algorithm will converge to. What *can* be guaranteed is that if the initial point is not already an optimum, then the optimisation will always provide a decrease in the cost function: in other words, it will always lead to an improvement in the spectrum (insofar as the cost function accurately represents the quality of the spectrum).

Finally, *POISE is not a panacea*. It should be noted that there is always an inherent tradeoff against

the time required for the optimisation itself. For example, it makes little sense to spend several minutes optimising the sensitivity of a pulse–acquire experiment: the time could simply be used to improve the SNR by collecting more scans. There is also the critical—though undeniably subjective—question of whether the optimisation is *necessary*: even if an optimisation can be run in a relatively short time, are the results really significantly better than a ‘compromise’ value in a default parameter set?* I do not profess to have a definitive answer to this, and I leave the reader to form their own conclusions in the specific contexts where they may consider using POISE. In any case, a ‘meaningful’ optimisation is likely to either be fast, and/or solve a problem which cannot simply be tackled through signal averaging in the same amount of time. It is my hope that this is true of (most of) the examples shown.

3.4 Applications

In this section, I cover a number of scenarios in which POISE can be used. These are generally ordered from simple to complex, and progressively show how the features in POISE can be used to customise optimisation procedures.

All POISE optimisations run in this chapter were performed five times to check for potential reproducibility issues. Due to noise in the cost function, these optimisations are not deterministic, and the optima obtained (as well as the time required to locate them) typically span a range. Where possible, this range is quoted in all the results shown in this chapter.

3.4.1 Pulse width calibration

The first of these applications is the calibration of a 90° ^1H pulse, which is applicable to virtually every NMR experiment. Essentially, we seek to determine τ_p for which

$$\tau_p \omega_1 = \frac{\pi}{2}, \quad (3.18)$$

where the RF amplitude ω_1 is not known *a priori* (it is only indirectly controlled via the power level). This pulse width is conventionally specified as the P1 parameter in TopSpin.

Optimisation setup

In theory, performing a pulse–acquire spectrum with a perfect 180° or 360° pulse would yield no detectable (i.e. transverse) magnetisation, i.e. a *null*. Generally, the 360° null is preferred as it

*Of course, a similar argument can be applied to *many* scientific advances. To use an example from the next chapter, is it really necessary to acquire NOAH spectra when one can just acquire the standalone 2D experiments? I have seen arguments on both sides—some people simply do not need the speedups provided and do not want to spend the time to set up or troubleshoot new experiments.

minimises effects due to radiation damping, and also allows a smaller recovery delay to be used. We can use POISE to search for this by acquiring the spectrum, performing a magnitude-mode calculation, and using the intensity of the resulting spectrum as a cost function:

$$f_{\text{minabsint}} = \sum_i |S_i|, \quad (3.19)$$

where (reusing notation from § 2.4.1) \mathbf{S} is the spectrum under consideration represented as a complex-valued vector, and the i -th point of the spectrum is $S_i = \sqrt{S_{\text{re},i}^2 + S_{\text{im},i}^2}$. The label `minabsint` refers to how this cost function drives the optimisation to *minimise* the *absolute intensity* of the spectrum. To show how easily this can be coded in Python 3, an implementation of this is shown in listing 3.3 (for all other cost functions in this chapter, the reader is directed to the POISE source code for their implementations). The `get1d_...` helper functions are provided by POISE.

```
def minabsint():
    r = get1d_real()
    i = get1d_imag()
    mag = np.abs(r + 1j * i)
    return np.sum(mag)
```

Listing 3.3: The implementation of the `minabsint` cost function in POISE.

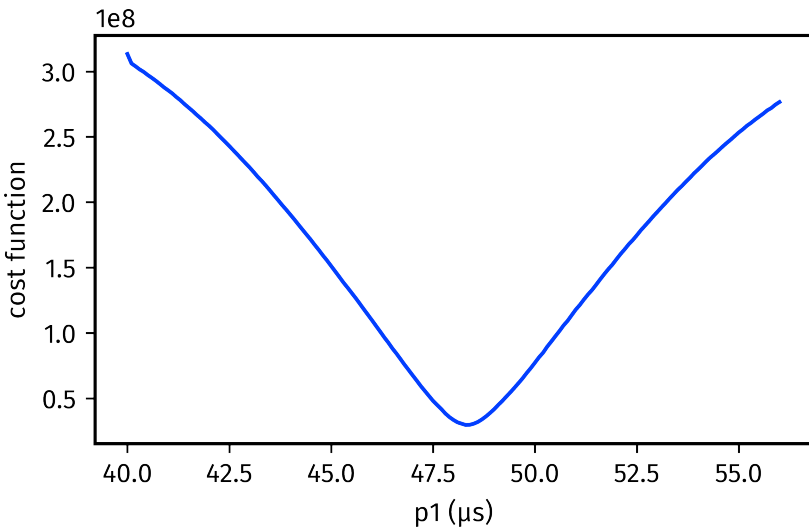


Figure 3.4: Reference grid search, showing how the `minabsint` cost function varies with the pulse width $P1$. Data code: 6F-200826.

To check whether this cost function was sensible, I manually acquired a series of spectra with increasing pulse widths and calculated $f_{\text{minabsint}}$ for all of these (fig. 3.4). In this chapter, I will refer to this process as a *reference grid search*. It should be noted that reference grid searches are

a time-consuming procedure, and an end-user of POISE generally does *not* need to do this (it would defeat the purpose of the optimisation). I only perform one here to provide some insight into the nature of the optimisation. In any case, the reference grid search makes it clear that there is a well-defined minimum, located in this case at 48.3 μs ; if the POISE optimisation process is accurate, it should converge to this point.

I chose the initial point to be four times the `prosol` value for P1: this represents our ‘best guess’ and is derived from prior calibration of the pulse width on a standard sample. The tolerance is set to 0.2 μs , which corresponds to an accuracy of 0.05 μs for the 90° pulse width itself. The lower and upper bounds are set to be 8 μs away from the initial point, representing a ‘sensible’ region within which we expect the null to lie (this may need to be adjusted for samples with high ionic strength, but for typical organic samples this is more than enough). A standard `poise_1d` acquisition AU programme is used, which simply acquires the spectrum and performs Fourier transformation, phase correction, and baseline correction. The routine in JSON format is shown in the caption of table 3.1. Finally, in order to reduce the time taken for the optimisation, several tricks are used: each FE is run using no dummy scans and only one scan, the acquisition time is set to just 1.1 s (high resolution is not required for a reliable cost function value), and the recovery delay D1 is set to 0. In practice, there is an extra gap of around 5 s between successive FEs due to spectrometer initialisation, so an extra recovery delay is not needed.

The competition

The performance of POISE can be compared against two ‘competitors’ in this area. The traditional method of determining the 90° pulse width is to measure a pulse width array (colloquially, ‘to array the pulse width’).⁴⁵ This entails measuring a series of pulse–acquire spectra, over which τ_p is evenly incremented: in optimisation parlance this would be called a *grid search*.^{*} This leads to a sinusoidal pattern in the peak intensities, from which the 360° null can be directly read off. An example of this is shown in fig. 3.5, where the 360° null at $\tau_p \approx 48 \mu\text{s}$ is visible (it is never a *perfect* null because of off-resonance effects and ω_1 , i.e. B_1 , inhomogeneity).

TopSpin provides a built-in mechanism for measuring a pulse width array using the `popt` command. The pulse–acquire spectrum is measured for pulse widths between a lower and upper bound, and the user specifies a spectral region of interest which `popt` uses to determine the null in spectrum intensity.[†] While this usually yields highly accurate results, the acquisition of so

^{*}This differs from the *reference grid search* above only in spirit (hence the similar names). The process is exactly the same, but the grid search is used to actually locate the optimum, whereas in the reference grid search the aim is purely to verify whether the faster optimisation algorithms converge to the correct point.

[†]In fact, `popt` uses the notion of a cost function as well, in that it determines the point where the cost function is minimised. In this case, I set it to use the `MAGMIN` cost function, which seeks to minimise the intensity of the magnitude-mode spectrum; this is essentially identical to the `minabsint` cost function which was used for the POISE optimisations except that it only applies to the spectral region of interest.

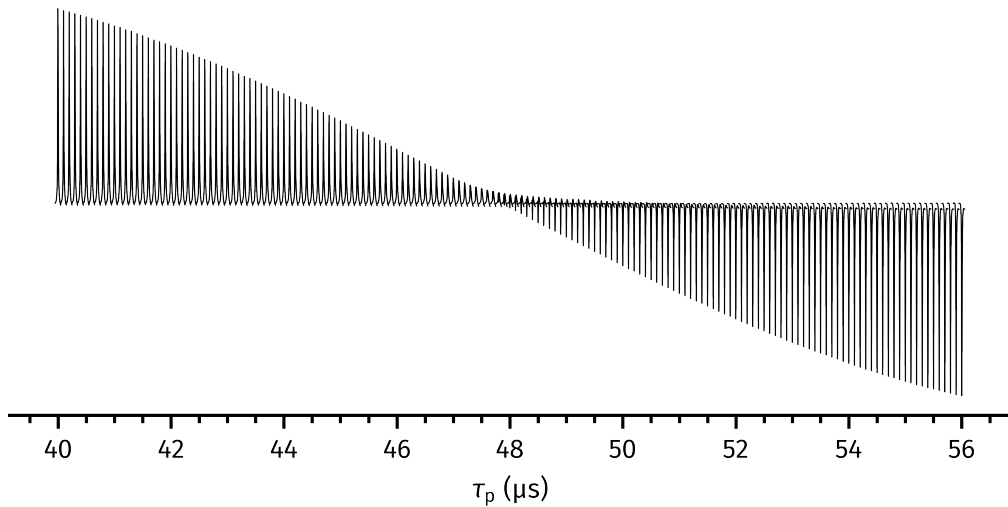


Figure 3.5: An example of a pulse width array, where the variation of the water peak is monitored with changes in pulse width (the sample used in DMSO- d_6 , but is slightly wet). These spectra were acquired manually; the TopSpin `popt` command would yield essentially identical results. *Data code:* 6F-200826.

many spectra is relatively time-consuming and arguably unnecessary if the only purpose is to determine the null.

A more rapid method for pulse calibration is the nutation experiment of Wu and Otting,⁴⁶ which allows the 90° pulse width to be determined in a single-scan experiment. In this experiment, an RF pulse with a given power level, corresponding to an (unknown) amplitude of ω'_1 , is applied during acquisition.* Assuming that the pulse is applied along the x -axis, this leads to the following product operators during the acquisition period:

$$I_z \xrightarrow{\omega'_1 t_2} I_z \cos(\omega'_1 t_2) - I_y \sin(\omega'_1 t_2) \quad (3.20)$$

and an FID of

$$s(t_2) = -\sin(\omega'_1 t_2) = -\frac{1}{2i}(\exp(i\omega'_1 t_2) - \exp(-i\omega'_1 t_2)), \quad (3.21)$$

which when Fourier transformed yields an antiphase doublet where the two peaks are separated by the frequency $2\omega'_1$. Measuring the separation between the two peaks directly yields the unknown amplitude ω'_1 , from which $\tau'_p = \pi/(2\omega'_1)$ can be calculated. Typically, the RF amplitude ω'_1 is rather smaller than the amplitude ω_1 which we would like to apply the hard pulse at, and thus $\tau'_p > \tau_p$. However, this can be adjusted for using the power levels applied (which are known to the user).

Although the nutation experiment can be performed extremely quickly using the TopSpin `pulsecal` command, often only requiring a few seconds, the pulse widths calculated are generally

*To be precise, it is applied for a proportion d of the dwell time between acquisition of successive points in the FID; d is called the *duty cycle* and must be accounted for when calculating the pulse width using this method.

slightly shorter compared to the value obtained from a 360° null. This is a known effect which arises because the separation is calculated from the top of the peaks, which correspond to the most homogeneous region of the B_1 profile.⁴⁶ On the other hand, the 360° null (as measured through `popt` or `POISE`, for example) measures a signal which is averaged over the entire B_1 profile.

Optimisation results

Entry	Method	Optimum found (μs)	FEs	Time taken (s)
1	<code>popt</code>	48.40	41	299
2	<code>pulsecal</code>	46.64	–	37
3	<code>POISE (NM)</code>	48.38	10	76–79
4	<code>POISE (MDS)</code>	48.38	10	77–80
5	<code>POISE (BOBYQA)</code>	48.29–48.41	6–7	46–54

Table 3.1: Comparison of methods for 360° pulse width determination. `popt` grid searches were run between values of $40\ \mu\text{s}$ and $56\ \mu\text{s}$, with a linear increment of $0.4\ \mu\text{s}$ (which, through interpolation, provides a precision of approximately $0.2\ \mu\text{s}$ in the result, matching the tolerance used for `POISE`). `pulsecal` was run as normal and the reported pulse width multiplied by 4 to obtain the 360° pulse width for comparison. `POISE` optimisations were run according to the routine in table 3.1. *Data code:* 6F-200826.

Compared to these two existing methods, we expect `POISE` to be faster than the `popt` grid search, and also more accurate than the nutation experiment in `pulsecal`. This is borne out in practice (table 3.1). `popt` yields an optimum of $48.4\ \mu\text{s}$, which is closely matched by `POISE`. However, across all five optimisation runs performed for each algorithm, `POISE` locates this optimum using far fewer FEs because its algorithms are more efficient than a simple grid search. While the `pulsecal` routine is even faster than `POISE`,^{*} it underestimates the 90° pulse width by about 4%. In this particular case, `POISE` is the only option which strikes a useful balance between speed and accuracy. These results also provide the first evidence that Py-BOBYQA is generally faster than the simplex-based methods: this observation is consistently reproduced in many of the other optimisations in this chapter.

Different initial points

One question we might reasonably ask is how robust `POISE` is towards poor initial guesses. In the case of the pulse width calibration, the answer is *very* robust. Tables 3.2 and 3.3 summarise the results obtained with an initial guess of $43\ \mu\text{s}$ and $53\ \mu\text{s}$ respectively. There is slightly decreased

^{*}It could be yet faster if it was instructed to not optimise the receiver gain prior to performing the nutation experiment.

performance in that a few more FEs are required for convergence, but the accuracy of the result is unchanged.

Entry	Method	Optimum found (μs)	FEs	Time taken (s)
1	POISE (NM)	48.38	14	109–114
2	POISE (MDS)	48.38	14	108–112
3	POISE (BOBYQA)	48.27–48.33	9	70

Table 3.2: Pulse width optimisations with an initial point of 43 μs . The POISE routine is the same as in table 3.1, except with "init": [43.0]. Data code: 6F-200826.

Entry	Method	Optimum found (μs)	FEs	Time taken (s)
1	POISE (NM)	48.38	14	110–114
2	POISE (MDS)	48.25–48.38	16	123–126
3	POISE (BOBYQA)	48.26–48.33	9	69–70

Table 3.3: Pulse width optimisations with an initial point of 53 μs . The POISE routine is the same as in table 3.1, except with "init": [53.0]. Data code: 6F-200826.

It is tempting to use this example to draw the conclusion that the initial point does not matter in POISE optimisations. However, this is only really true for a simple optimisation like this. Looking again at the reference grid search in fig. 3.4, it is clear that there is no other possible minimum that the optimiser could converge to. Furthermore, the noise in the cost function is almost indiscernible. These represent the *ideal* conditions for an experimental optimisation to work, and it is not surprising that extremely good performance is obtained with POISE. Some of the subsequent examples include more difficult or more noisy cost functions. We will see that POISE does indeed have *some* tolerance towards poor initial points, even in the presence of noise (after all, this is the entire purpose of using derivative-free algorithms). However, for very challenging optimisations (especially ones with many parameters) it is very likely that the optimisation will ultimately converge to a local minimum close to the initial point.

3.4.2 Ernst angle optimisation

Often, in a simple 1D pulse–acquire spectrum it is not hugely important to know the exact 90° pulse width: instead, it is more valuable to optimise the sensitivity per unit time of the spectrum. Before launching straight into how this may be obtained through optimisation, it is instructive to first consider which parameters are worth optimising. For a pulse–acquire experiment (fig. 3.6), the repetition time τ_r is the sum of the acquisition time AQ plus the recovery delay D1; the flip angle θ is controlled via the pulse width P1. We assume that the experiment has been repeated enough times to reach a *steady state*, that is, the amount of z -magnetisation prior to the excitation

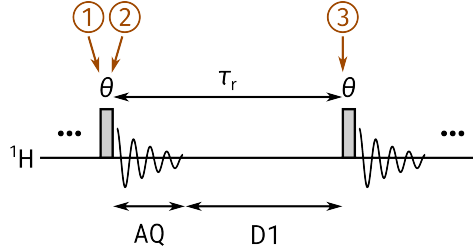


Figure 3.6: Steady-state pulse–acquire experiment. The excitation flip angle is θ , and the repetition time between experiments is τ_r .

pulse (point ①) is a constant, $M_{z,ss}$. Application of the excitation pulse leads to a signal scaling as $M_{z,ss} \sin \theta$, and residual (unexcited) longitudinal magnetisation of $M_{z,ss} \cos \theta$ at point ②. After the repetition time τ_r (point ③), it can be shown using the Bloch equations⁴⁷ that the z -magnetisation recovers to

$$M_{z,0}(1 - c) + cM_{z,ss} \cos \theta, \quad (3.22)$$

where $c = \exp(-\tau_r/T_1)$ and $M_{z,0}$ is the initial, equilibrium z -magnetisation (before the experiment begins). Since the experiment has reached a steady state, points ① and ③ are equivalent: thus, we have that

$$M_{z,0}(1 - c) + cM_{z,ss} \cos \theta = M_{z,ss}, \quad (3.23)$$

which can be rearranged to give

$$\frac{M_{z,ss}}{M_{z,0}} = \frac{1 - c}{1 - c \cos \theta}. \quad (3.24)$$

The signal amplitude s therefore scales as

$$s = \frac{(1 - c)}{1 - c \cos \theta} \cdot \sin \theta, \quad (3.25)$$

and is maximised when $ds/d\theta = 0$, the solution of which is the celebrated *Ernst angle*:⁴⁸

$$\theta_E = \arccos c = \arccos \left[\exp \left(-\frac{\tau_r}{T_1} \right) \right]. \quad (3.26)$$

In general, T_1 and hence θ_E varies across the different spins in a given sample, so some degree of compromise is required in order to maximise sensitivity for all peaks.

To begin with, we may consider fixing τ_r and optimising P1 to locate the Ernst angle (or to be precise, the pulse width which corresponds to the Ernst angle). This is by no means an illogical proposition. However, we can go one step further, because τ_r itself is comprised of two parameters, and the sensitivity *per unit time* may be affected by varying τ_r . Since the signal scales as $1/\tau_r$ (a shorter τ_r means more repetitions per unit time) but the noise scales only as $\sqrt{1/\tau_r}$,

the sensitivity per unit time is

$$S = \frac{(1 - c) \sin \theta}{(1 - c \cos \theta) \sqrt{\tau_r}}. \quad (3.27)$$

Assuming that θ is always set to the respective Ernst angle for different τ_r , it can be shown that the best sensitivity per unit time is attained when $\tau_r \rightarrow 0$.^{49,50} Of course, this limit is not physically possible: τ_r comprises the acquisition time which must be nonzero. However, it does imply that AQ should be kept as short as possible, and D1 set to zero, as shown in fig. 3.7.

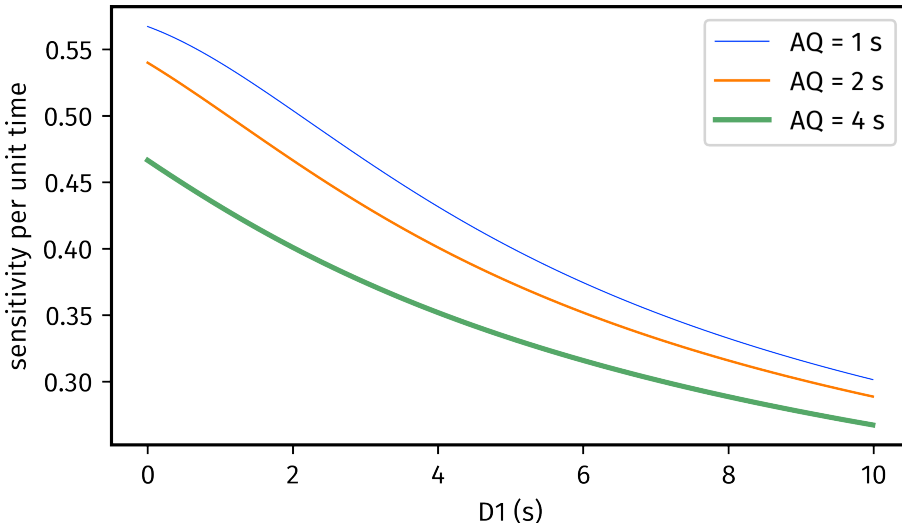


Figure 3.7: Simulated sensitivity per unit time as a function of AQ and D1 (as given by eq. (3.27)), assuming that an Ernst angle excitation pulse is used. T_1 was set to 1.5 s.

Optimisation setup

Knowing this, we can then set up a meaningful optimisation routine. We seek to optimise the pulse width P1 such that the intensity of the real part of the spectrum is maximised (corresponding to a `maxrealint` cost function). In practice, I took the extra step of calibrating the 90° pulse width (as per the previous section) and modifying the pulse programme such that the flip angle could be specified as the parameter `CNST20`: this is not generally necessary and is only useful for evaluating the results, as will be shown later. AQ and D1 were set to be 1.2 s and 0 respectively, in accordance with the theory outlined above. The number of scans (NS) can be set to 1, but unlike in the pulse width calibration (§ 3.4.1), we must use enough dummy scans (DS) to ensure that a steady-state signal intensity is recorded: in practice I set DS=4. This means that each FE, and thus the overall optimisation, requires a slightly longer time than the pulse width calibrations previously shown.

Optimisation results

The Ernst angle optimisation was run with two different spectral regions of interest: firstly, on all the aromatic and olefinic peaks in the sample of ferulic acid (table 3.4), and secondly, on only a single peak at 6.79 ppm (table 3.5). (This spectral region can be selected using the TopSpin dp1 command, which stores the bounds using the parameters F1P and F2P: all built-in cost functions respect these two parameters.) Generally, optimisations could be completed in under two minutes. The optima found for these two optimisations are different: this is because the former searches for a compromise Ernst angle which balances T_1 of all peaks within the region, and the latter optimises only for one T_1 .

Entry	Algorithm	Optimum found (°)	FEs	Time taken (s)
1	NM	67.5–73.1	9–13	91–132
2	MDS	67.5–73.1	9	90–92
3	BOBYQA	70.1–70.7	7	70–71

Table 3.4: Ernst angle optimisation, performed on all aromatic and olefinic peaks in ferulic acid (between 6 and 8 ppm). The POISE routine used here is: `{"name": "ernst", "pars": ["cnst20"], "lb": [10.0], "ub": [90.0], "init": [30.0], "tol": [3.0], "cf": "maxrealint", "au": "poise_1d"}`. The mean of all five theoretical Ernst angles is 65.2°. Data code: 5F-210619.

Entry	Algorithm	Optimum found (°)	FEs	Time taken (s)
1	NM	60.0–67.5	9–11	91–111
2	MDS	65.6–67.5	11	110–111
3	BOBYQA	60.0–65.2	6–7	59–71

Table 3.5: Ernst angle optimisations on the peak at 6.79 ppm in ferulic acid. The POISE routine is the same as in table 3.4, but the spectral region under optimisation was set to be 6.71–6.87 ppm. The theoretical optimum, as given in table 3.6, is 61.6°. Data code: 5F-210619.

To determine the accuracy of the optima found, rather than performing a reference grid search as in § 3.4.1, I measured T_1 of each of these peaks using a typical gradient-enhanced inversion-recovery experiment. From this, the theoretical Ernst angles for each peak could be calculated (table 3.6): they range from 60° to 73°. The first optimisation, which should yield a weighted average of the five Ernst angles, appears at first glance to be biased towards the upper end of this range. However, this can be rationalised by the fact that a flip angle larger than θ_E is less detrimental to sensitivity compared to one that is smaller (this can be seen by plotting eq. (3.25)). On the other hand, the second optimisation yields an accurate value for the relevant peak (61.6°, peak number 4 in table 3.6).

Peak	^1H chemical shift (ppm)	T_1 (s)	θ_E ($^\circ$)	$T_1 \ln 2$ (s)
1	7.49	1.750	59.8	1.213
2	7.27	0.977	73.0	0.677
3	7.08	1.279	67.0	0.887
4	6.79	1.615	61.6	1.119
5	6.36	1.415	64.6	0.981

Table 3.6: T_1 and corresponding Ernst angles for each peak in ferulic acid, calculated for a repetition time of 1.20 s (peak assignments can be found in table A.3). The values of $T_1 \ln 2$, measured via a full pseudo-2D inversion–recovery experiment, are also provided here in anticipation of the optimisations in § 3.4.3. Data code: 5F-210619.

It is worth considering, though, whether this optimisation is truly worth it. ^1H pulse–acquire spectra already have a very high intrinsic sensitivity, and in the two minutes taken to optimise the flip angle, one could easily just acquire (around) 64 more scans, at which point knowledge of the Ernst angle would cease to be useful. It *may* be more useful for nuclei which have lower sensitivity, such as ^{13}C : I did not evaluate this possibility. However, it must be borne in mind that a low-sensitivity experiment will also require more scans per FE, which leads to a corresponding increase in the optimisation time.

In my estimation, a more useful application of this optimisation routine would be to use it to determine an average T_1 value for a group of peaks. This could then be used to inform the choice of recovery delay for multidimensional experiments^{51,52} or quantitative NMR experiments.^{53,54} It is, however, possible to more directly obtain T_1 values from an inversion–recovery experiment, which I describe next.

3.4.3 Inversion–recovery

Optimisation setup

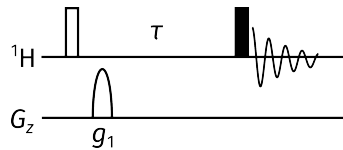


Figure 3.8: Inversion–recovery pulse sequence.

If what we really want to measure is T_1 for a particular peak (or a set of peaks), a more direct way is to perform an inversion–recovery experiment (fig. 3.8). This can either be recorded in a 2D form where the τ delay is incremented and the resulting intensities fit to an exponential curve, or in an iterative fashion by searching for the null in spectral intensity, which occurs at $\tau = T_1 \ln 2$: this latter option is particularly suited to optimisation. The delay τ was specified

as the parameter D27, and the `zerorealint` cost function used here is just the absolute value of the integral over the region of interest: an ideal null would have a cost function value of 0. Table 3.6 provides the theoretical values of $T_1 \ln 2$, which were calculated using the more accurate 2D fitting procedure.

A slight drawback of using this method, compared to the Ernst angle optimisations in the previous section, is that the recovery delay must be sufficiently long to allow for complete relaxation between FEs: in this case, I used a D1 of 5 s. This is compensated for by the fact that dummy scans are no longer needed to reach a steady-state; I therefore set DS=0 and NS=1.

Optimisation results

Just as in the Ernst angle optimisations, two different types of optimisations were performed: one on the entire 6 ppm to 8 ppm region, and one on just a single peak (note that here, the chosen peak is at 7.08 ppm, or peak 3 in table 3.6). The results are shown in tables 3.7 and 3.8. The optimisation over the entire aromatic region correctly converges to an ‘averaged’ value of $T_1 \ln 2$ over the five peaks (the mean value for the five peaks is 0.975 s). The second optimisation performed only on a single peak is no less accurate, yielding an optimum which closely matches the theoretical value for the peak in question (0.887 s).

Entry	Algorithm	Optimum found (s)	FEs	Time taken (s)
1	NM	0.938–0.969	14–16	204–235
2	MDS	0.956–0.975	16	233–235
3	BOBYQA	0.953–0.971	9–11	130–160

Table 3.7: Inversion–recovery optimisations on all aromatic and olefinic peaks in ferulic acid (6 ppm to 8 ppm). The POISE routine used here is: `{"name": "invrec", "pars": [{"d27": 0.35}, {"ub": 1.75}, {"init": 0.6}, {"tol": 0.01}, {"cf": "zerorealint"}, {"au": "poise_1d"}]`. The mean of $T_1 \ln 2$ values for all five peaks is 0.975 s. Data code: 5F-210619.

Entry	Algorithm	Optimum found (s)	FEs	Time taken (s)
1	NM	0.863–0.875	14	202–205
2	MDS	0.863–0.869	14	203–204
3	BOBYQA	0.862–0.873	9–10	128–145

Table 3.8: Inversion–recovery optimisations on the peak at 7.08 ppm in ferulic acid. The POISE routine is the same as in table 3.4, but the spectral region under optimisation was set to be 7.02–7.15 ppm. The theoretical optimum (from table 3.6) is 0.887 s. Data code: 5F-210619.

The only downside of these optimisations would then be the time required, which is on the order of 2–4 minutes. Although this is less time than required for a full pseudo-2D inversion–recovery

experiment, POISE has the drawback that an optimisation can only be run on one peak at a time. Thus, if the aim is to determine T_1 for all peaks, then the 2D experiment may well end up being faster. On top of that, there are many other ways to measure T_1 which are faster than a full 2D inversion–recovery experiment and almost certainly also faster than POISE.^{55–59} However, no explicit comparisons were performed in this work.

3.4.4 NOE mixing time

In all of the optimisations done previously, the cost functions used are relatively simple, simply seeking to maximise or minimise some intensity. In fact, the Bruker popt interface does come with a number of cost functions itself, which can be used for grid search-based optimisations: so, in principle, all the previous examples could have been done with popt (albeit with a much longer time). However, POISE goes beyond this in that it allows users to define their own cost functions. In this section, we exploit this customisability to devise a more complicated cost function for optimising mixing times in NOE experiments.

Optimisation setup

The ideal NOE mixing time for a given compound depends on the rates of various relaxation processes: too short a mixing time does not allow for sufficient buildup of the NOE, but too long a mixing time leads to loss of signal through relaxation. In this section, I do not deal with this theoretically: the optimisation process is merely used to find the empirically best value (for the sample under study).

In the chosen sample, 3-fluorophenylboronic acid, there are four crosspeaks of interest in the 2D NOESY spectrum. I first performed a reference grid search to determine where the crosspeak intensities were maximised (fig. 3.9). Generally, a broad minimum between 2.5 and 4 s is observed: any result within this range should be considered as ‘correct’. The relatively large value for the optimal mixing time reflects the small size of the molecule used in this study (it has a molar mass of 139.92 g mol^{-1}), which corresponds to rapid tumbling and long T_1 values.* Although the broad minimum may at first glance seem imprecise, it merely reflects the underlying physical characteristics of the sample under study: an optimisation process cannot ‘discover’ extra precision where there is none to be found. We also see here the first example of a cost function where the noise is significant: this provides a good test of the derivative-free algorithms used in POISE.

*There is a complicating factor in that the use of such long mixing times also leads to a noticeable increase in the experiment duration. As such, it may be more prudent to consider the sensitivity *per unit time*. In the optimisations which follow, I have neglected this issue: the 2× increase in sensitivity demonstrated below would only be cancelled out by a 4× increase in time, which is not the case here. However, to be more rigorous, this factor could be accounted for by modifying the form of the cost function.

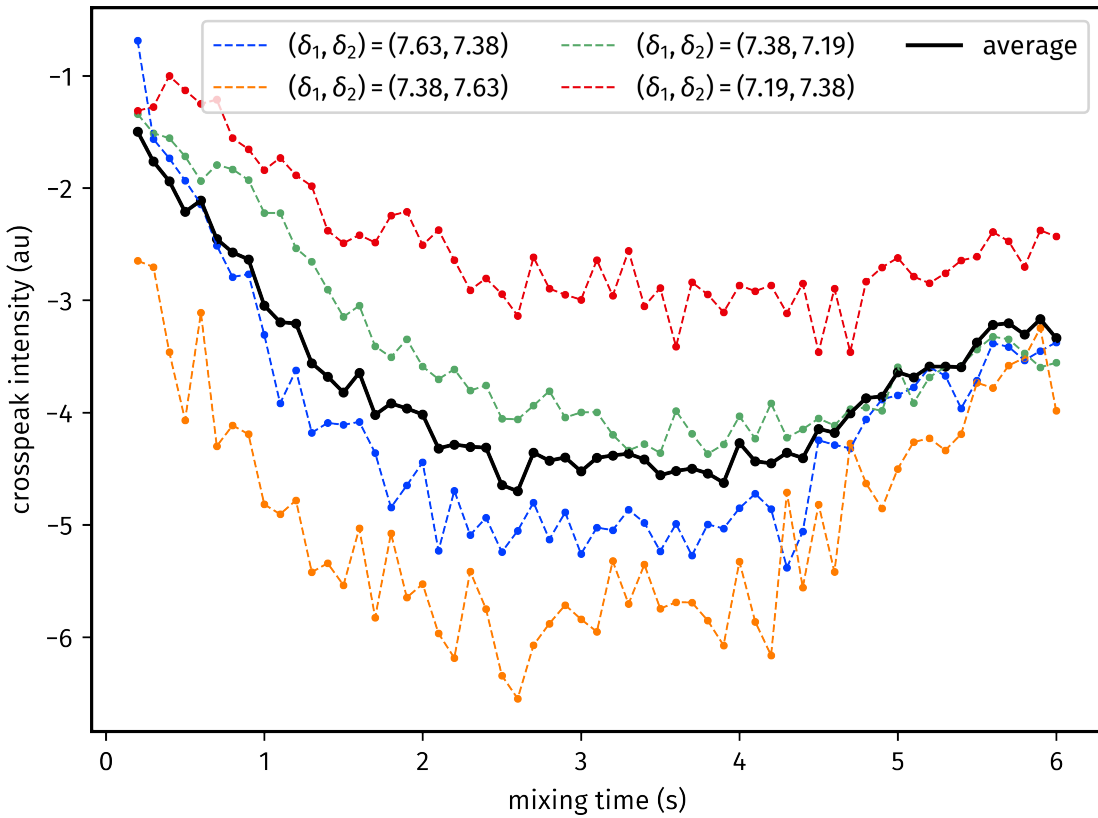


Figure 3.9: Reference grid search of 2D NOE crosspeak intensities. The 2D NOESY spectra themselves are shown in fig. 3.11. The individual crosspeak intensities are shown in dashed lines; the solid black line is the average of these four, which represents the quantity we seek to optimise. Data code: 7B-200725.

While the 2D reference grid search offers first-hand insight into what our target optimum should be, it is unwise to run an optimisation using the full 2D experiment, simply because of the time required for each FE. A more sensible method is to use a selective 1D NOESY experiment. Although this is much faster, it does come with two caveats:

1. The frequency for selective irradiation must be first chosen, likely after acquisition of a 1D ^1H spectrum. Thus, the optimisation does require some *a priori* knowledge of the system being studied.
2. The peaks observed in the 1D NOESY *must* be sufficiently representative of those in the full 2D NOESY.

As for the cost function, it must be able to pick out only the peaks from the 1D NOESY spectrum which correspond to genuine NOE transfer, rejecting (for example) the selectively irradiated peak as well as other artefacts. The cost function used here (`noe_1d`) does this by reading the frequency used for the selective irradiation from the SPOFFS2 parameter, excising a region of ca. 100 Hz around this irradiation frequency, and integrating the remainder of the spectrum.

In order to account for the fact that the desired peaks may be either positive or negative, the absolute value of the integral is taken, and the negative of this is used as the cost function (since we seek to maximise the intensity). Note that if a different 1D NOESY pulse programme is used with different parameter definitions, then the cost function must be adjusted accordingly.

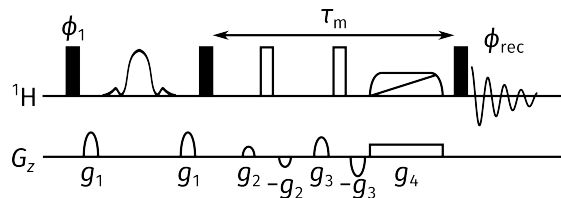


Figure 3.10: Selective 1D NOESY pulse sequence used for POISE optimisations. Phase cycling was performed using $\phi_1 = \phi_{\text{rec}} = (x, -x)$. Gradient amplitudes were set as follows: ($g_1, g_2, g_3, g_4 = 17\%, 40\%, 25\%, 10\%$).

In the event, I used a modified 1D NOESY pulse programme (fig. 3.10), with two extra inversion pulses inserted during the mixing time: these minimise artefacts arising from relaxation during the mixing time, which can be especially problematic for long mixing times of several seconds. (In principle, these artefacts have equal positive and negative components and thus should not contribute to the cost function except in terms of noise; however, it is always a good idea to minimise the noise in the cost function as much as possible.) The mixing time is represented by the parameter D8. The initial value for the optimisation was set to 0.5 s, which is a reasonable compromise value for most ‘small’ organic molecules.

Optimisation results

The results of this optimisation are shown in table 3.9. The peak at 7.38 ppm was used for selective irradiation. In all cases, the optimisations converged to the correct region within 2.5 minutes (for BOBYQA) or 5 minutes (for the simplex-based algorithms). The resulting 2D NOESY spectra, with the initial and optimised mixing times of 0.5 s and 3.5 s respectively, are shown in fig. 3.11, where the improvement in crosspeak sensitivity is clearly visible.

Entry	Algorithm	Optimum found (s)	FEs	Time taken (s)
1	NM	3.25–3.88	16–18	268–312
2	MDS	3.63–3.75	16–18	269–305
3	BOBYQA	3.38–3.80	6–10	88–162

Table 3.9: NOE mixing time optimisations on a sample of 3-fluorophenylboronic acid. The POISE routine used here is: `{"name": "1dnoe", "pars": ["d8"], "lb": [0.2], "ub": [6.0], "init": [0.5], "tol": [0.1], "cf": "noe_1d", "au": "poise_1d"}`. Data code: 7B-200721.

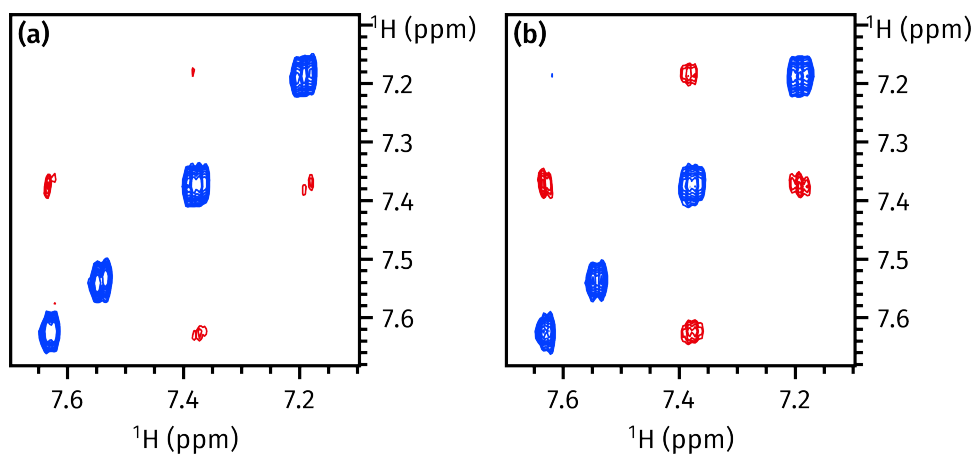


Figure 3.11: 2D NOESY spectra of 3-fluorophenylboronic acid, obtained (a) before and (b) after optimising the mixing time on the 1D NOESY sequence in fig. 3.10. Both spectra are plotted with the same contour levels. *Data code:* 7B-200725.

It should be mentioned here that each FE was run using one dummy scan and two scans. This was only made possible by the high SNR afforded by a concentrated sample (120 mM), as well as a cryogenic probe. For more dilute samples where SNR is insufficient, the POISE optimisation will require more scans per FE, and consequently will take longer. However, it can be argued that the *benefit* reaped from the optimisation is also larger, since the final (optimised) 2D NOESY would also take a correspondingly longer time.

A different sample

To more clearly illustrate the *sample-specific* nature of POISE optimisations, I also performed the same optimisation on a different sample: in this case, the decapeptide gramicidin S. For this rather larger compound, we would expect the optimisation to converge to a shorter mixing time; this is indeed observed in practice (table 3.10). The peak at 4.76 ppm was used for selective irradiation: this is the H^α proton of the ornithine residue. In this case, the sample is rather less concentrated, and I used DS=2 and NS=4; this is still a relatively small number of scans, but was sufficient to yield accurate results.

Entry	Algorithm	Optimum found (s)	FEs	Time taken (s)
1	NM	0.63–0.78	9–14	253–384
2	MDS	0.44–0.75	9–11	254–309
3	BOBYQA	0.58–0.77	5–8	136–223

Table 3.10: NOE mixing time optimisations on a sample of gramicidin S. The POISE routine used here is identical to before. *Data code:* 7G-210815.

3.4.5 ASAP-HSQC excitation delay

The next example of a POISE optimisation is the ASAP-HSQC experiment:^{60,61} here, I use it to again illustrate the new possibilities which custom cost functions enable. In the ASAP-HSQC experiment (fig. 3.12), an HSQC spectrum is recorded using only ^{13}C -bound proton magnetisation, and the ^{12}C -bound ('bulk') proton magnetisation is returned to the equilibrium $+z$ axis (i.e. the I_z state). In this way, instead of having a conventional recovery delay, the $^{13}\text{C}-^1\text{H}$ magnetisation can be directly replenished using isotropic mixing, which causes transfer of z -magnetisation from bulk protons. The elision of the recovery delay from the sequence thus leads to significantly shorter experiment durations: in one of the more extreme examples, an HSQC spectrum could be recorded within 7 seconds. This separation of different 'magnetisation pools' is conceptually very similar to that in NOAH experiments (§ 4.1.2), and in anticipation of that, I use the notation $^1\text{H}^{\text{C}}$ and $^1\text{H}^{\text{IC}}$ to represent magnetisation belonging to protons coupled and not coupled to ^{13}C respectively.

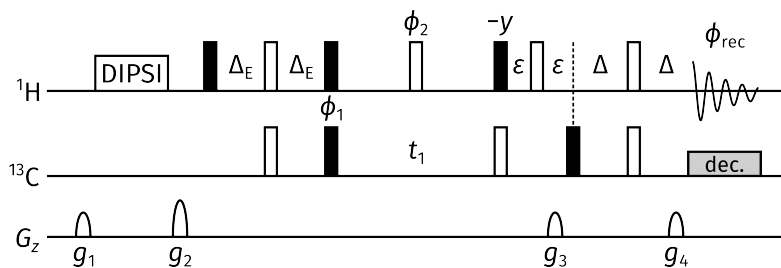


Figure 3.12: ASAP-HSQC pulse sequence. Phase cycling is performed using $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x)$, and $\phi_{\text{rec}} = (x, -x, x, -x)$. Gradient amplitudes are as follows: $g_1 = 33\%$, $g_2 = 43\%$, and for echo-antiecho selection, $(g_3, g_4) = (59.9\%, 80\%)$ and $(63.9\%, 80\%)$ respectively. The delay Δ is set to $1/(4 \cdot ^1J_{\text{CH}})$; the delay Δ_E was optimised as described in the text. The BIBOP, BEBOP, and BURBOP optimal control pulses^{12,14,15} were used for ^1H 180° pulses and all ^{13}C pulses; for more details, refer to the original paper by Luy and coworkers.⁶⁰

By modifying the length of the delay Δ_E in the initial INEPT block, it is possible to adjust the proportion of the $^1\text{H}^{\text{C}}$ magnetisation excited during the sequence: the remainder is stored along z . The delay Δ_E can be related to an *effective flip angle*, θ_{eff} , through

$$\theta_{\text{eff}} = 2\pi J \Delta_E. \quad (3.28)$$

where J is here short for $^1J_{\text{CH}}$. This partial excitation means that the signal is decreased by a factor of $\sin \theta_{\text{eff}}$, but means that on the next scan or increment, there is a larger pool of $^1\text{H}^{\text{C}}$ magnetisation to start from. The combination of these factors means that there is an optimum Δ_E which yields the greatest steady-state signal. Equivalently, we can define an 'effective J-coupling'

J_{eff} , for which the INEPT delay is optimised:

$$\Delta_E = \frac{1}{4J_{\text{eff}}} \iff J_{\text{eff}} = \frac{\pi J}{2\theta_{\text{eff}}}, \quad (3.29)$$

and we can search for the ideal J_{eff} to maximise the sensitivity of the ASAP-HSQC spectrum.

Notice that this is entirely analogous to the Ernst angle previously discussed in § 3.4.2. However, here, the magnetisation is being recovered partly through isotropic mixing and partly through relaxation during the FID. This makes an analytic description of the ASAP-HSQC experiment rather more complicated (although it has been studied before⁶²). Directly optimising the excitation delay Δ_E with POISE, which circumvents the theory entirely, is therefore an attractive option for practical day-to-day use.

Optimisation setup

Given the analysis above, it is extraordinarily easy to set up the optimisation: the parameter being optimised is J_{eff} , and the initial point chosen is a ‘compromise’ $^1J_{\text{CH}}$ value of 150 Hz. Although generally it is not a good idea to use a 2D experiment as part of the FE, in this specific case it is acceptable because the ASAP-HSQC has such a short running time. In this case, we simply take the projection of the 2D ASAP-HSQC onto the F_2 axis, integrate it, and take the negative of that to obtain the cost function.

A reference grid search was performed so that I could later verify the optimisation results (fig. 3.13). The exact point where the crosspeak intensities are maximised is not obvious: generally, the signal increases up until $J_{\text{eff}} \approx 230$ Hz, after which it plateaus off. In a similar fashion to the NOE optimisations (§ 3.4.4), we may therefore consider any value above this to be ‘correct’.

To speed up the optimisation, optimisations were run using a ‘low-quality’ ASAP-HSQC spectrum with only 32 t_1 increments. A recovery delay of 0.1 s was used.

Optimisation results

Entry	Algorithm	Optimum found (Hz)	FEs	Time taken (s)
1	NM	250.0–256.3	8–9	169–195
2	MDS	237.5–243.8	8	171–179
3	BOBYQA	229.8–245.6	4–7	114–157

Table 3.11: ASAP-HSQC INEPT delay optimisations. The POISE routine used was:

```
{"name": "asaphsqc", "pars": ["cnst3"], "lb": [120.0], "ub": [280.0], "init": [150.0], "tol": [10.0], "cf": "asaphsqc", "au": "poise_2d"}. Data code: 7B-200722.
```

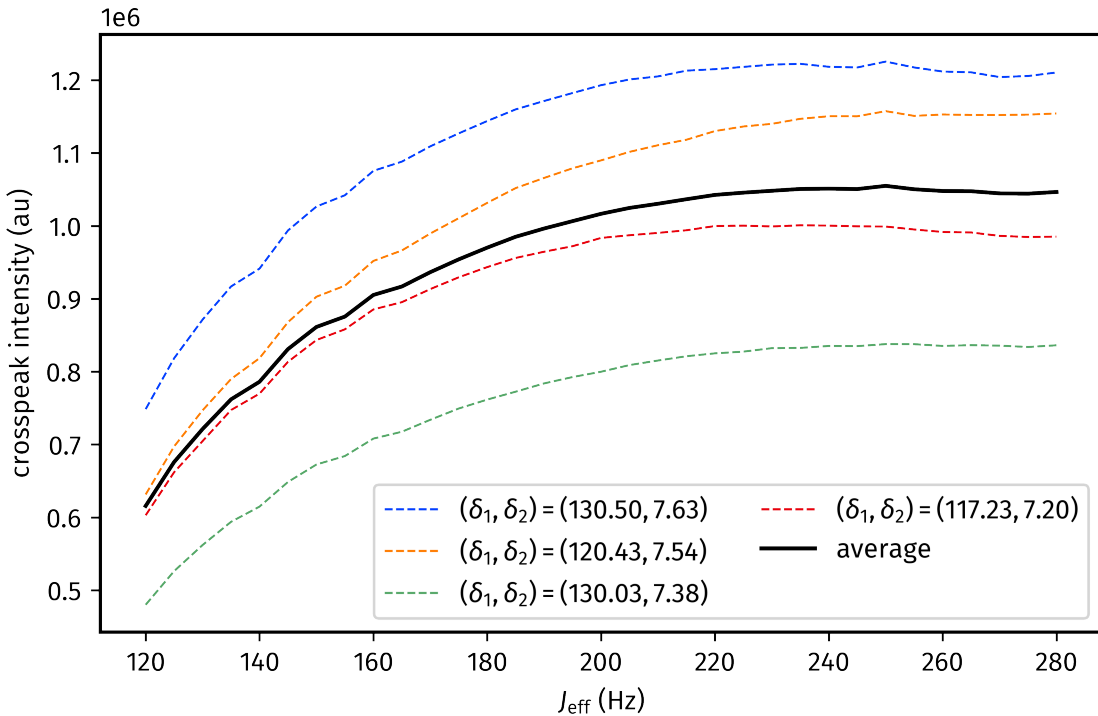


Figure 3.13: Reference grid search showing how the ASAP-HSQC signal intensity for the 3-fluorophenylboronic acid sample varies with J_{eff} . Data code: 7B-200722.

The results from the POISE optimisations are collated in table 3.11. As can be seen, even though the FE involves acquisition of a 2D spectrum, the optimisation itself completes in just 2–3 minutes, correctly converging to J_{eff} values in the region of around 240 Hz. When re-evaluated on a ‘full’ ASAP-HSQC experiment with 64 t_1 increments (fig. 3.14a),^{*} the optimisation yielded an 18–27% improvement in the sensitivity of the ASAP-HSQC spectrum, as shown in fig. 3.14b.

In fact, this specific example is not particularly impressive in itself. The ASAP-HSQC experiment shown in fig. 3.14a takes only around 45 seconds to acquire, so simply repeating it two or three more times would yield a similar or even larger increase in SNR. However, this argument may easily be swung in the opposite direction if more t_1 increments were to be used, such as 256 for a more ‘typical’ ^{13}C spectral window, or even more if extremely high F_1 resolution is desired (the original paper⁶⁰ provides an example of a three-hour ASAP-HSQC experiment where 16384 t_1 increments were recorded to differentiate two peaks separated by 3 Hz in F_1).

3.4.6 Ultrafast NMR

The final example of a single-parameter POISE optimisation is also the most complicated: it pertains to the EPSI acquisition technique, which was previously introduced in § 2.7. EPSI

^{*}The ^{13}C resonances of interest in this compound fall within a very small window, so relatively few t_1 increments are needed.

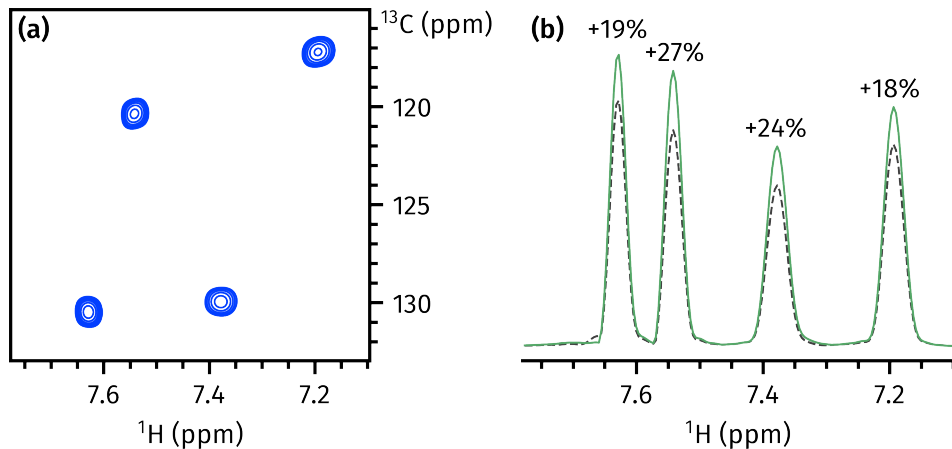


Figure 3.14: (a) ASAP-HSQC spectrum of the 3-fluorophenylboronic acid sample. (b) Projections of the ASAP-HSQC spectra before (grey dotted line, using $J_{\text{eff}} = 150 \text{ Hz}$) and after (green dotted line, using $J_{\text{eff}} = 245 \text{ Hz}$) optimisation of the INEPT delay. Sensitivity increases for each peak are indicated. *Data code:* 7B-200722.

acquisition allows signals from different slices of the sample to be simultaneously detected in a single FID, and (in liquid-state NMR) has most famously been used in the context of *ultrafast* 2D experiments,^{63–69} where the t_1 evolution is spatially encoded and read out using the EPSI technique.

During an EPSI acquisition period, alternating gradients of equal magnitude but opposite sign are applied, as shown in figs. 3.15a and 3.15b. Each data point collected therefore depends not only on t_2 (the acquisition time domain) but also on k , a wavenumber which measures the extent of gradient dephasing caused by the acquisition gradients:

$$k = \int_0^{t_2} \gamma G(t) \, dt. \quad (3.30)$$

This value of k increases during positive gradients and decreases during negative gradients, leading to a zigzag pattern in k -space, all while t_2 is still increasing (fig. 3.15c). The *prephasing gradient* immediately before acquisition has the same duration as the EPSI gradients, but has half the amplitude, meaning that k begins from a negative value, specifically, $-k_{\max}/2$, where k_{\max} is the change in k caused by one complete positive EPSI gradient. The indirect-dimension frequencies F_1 can be directly read off from the k -domain as F_1 is directly proportional to k : this can also be understood as the EPSI gradients unwinding a phase-modulated ‘helix’ created by chemical shift evolution during the t_1 period, as has been beautifully illustrated in previous reviews.^{65,66}

This rapid alternating of gradients is very demanding on spectrometers, and it is often the case that the positive and negative EPSI gradients—although *nominally* specified with the same amplitude—are not perfectly balanced. This causes a ‘drift’ in the k -domain over time, as illus-

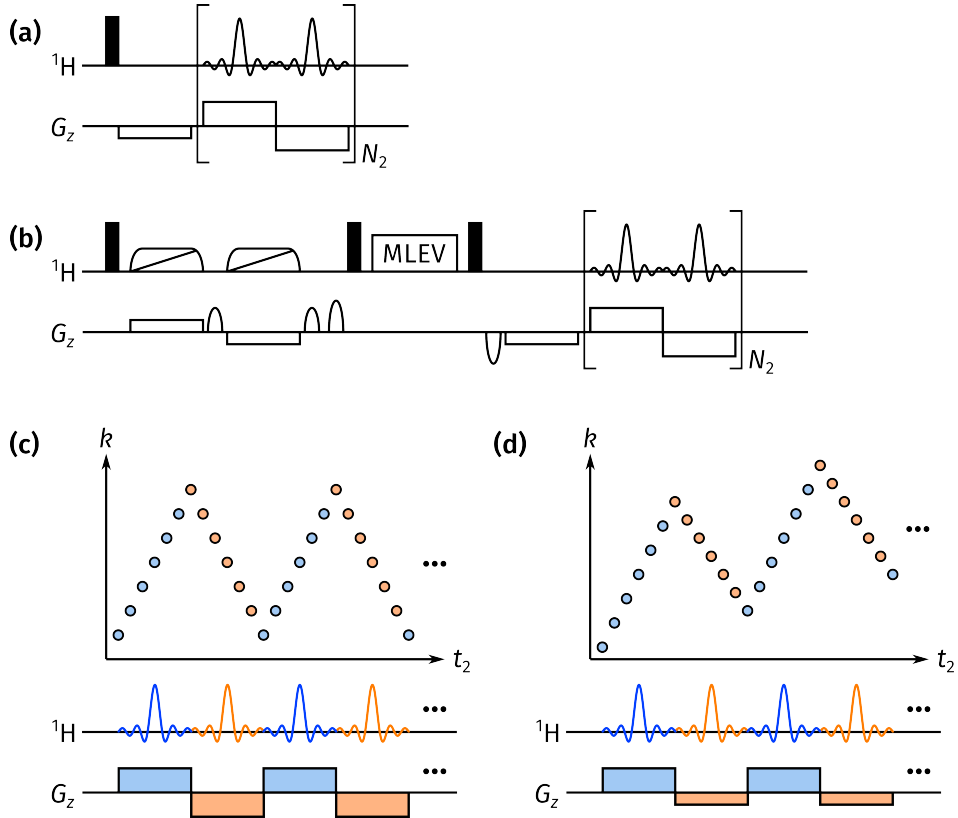


Figure 3.15: Pulse sequences used for EPSI optimisation, and depiction of the (k, t_2) data matrices thus obtained. (a) Pulse-EPSI experiment. (b) Ultrafast TOCSY experiment; gradients were set up according to the protocol of Giraudeau et al.⁶⁸ (c) Illustration of the (k, t_2) data matrix obtained during an EPSI acquisition period. Blue dots indicate data acquired during a positive gradient, and orange dots data acquired during the negative gradient. (d) The (k, t_2) data matrix obtained when the positive and negative EPSI gradients (blue and orange in the pulse sequences) are not balanced. In this case, the negative gradient is slightly weaker, leading to a ‘drift’ towards larger values of k .

trated in fig. 3.15d. In this section, POISE is used to perform an *instrument-specific* optimisation in order to correct for this effect.

Optimisation setup

To measure the drift in k -space, it is easiest to use a pulse sequence for which there is no spatial encoding. The pulse-EPSI experiment (fig. 3.15a) is perfectly suited for this. In the pulse programme, I multiplied the amplitudes of the negative gradients by a factor α , represented as CNST16 in TopSpin: the objective of POISE is therefore to determine the optimum value for this factor which minimises the drift. Calculating this drift requires fairly substantial processing, which is most easily done inside the cost function using the numpy library. The `epsi_gradient_drift` cost function therefore consists of the following steps:

1. reading in the 1D FID and reshaping it into a 2D (k, t_2) matrix;

2. discarding data obtained during negative gradients;
3. determining, for each point in t_2 , the value of k for which the maximum signal is found;
4. performing linear regression on these points and obtaining the slope of k against t_2 , the absolute value of which directly represents the extent of k -space drifting.

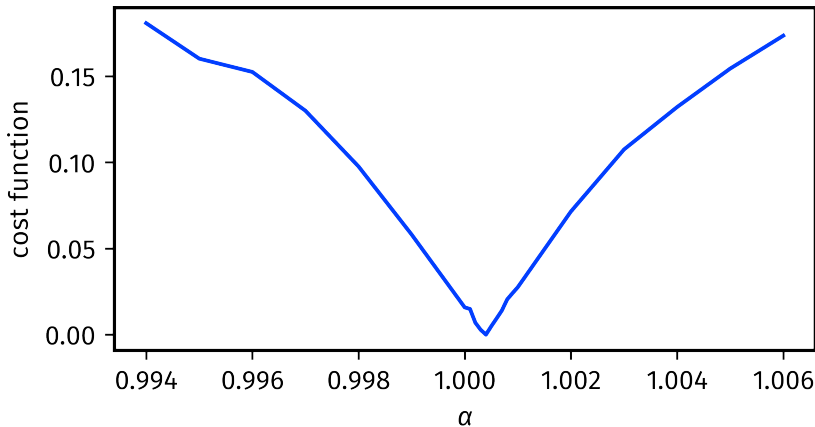


Figure 3.16: Reference grid search for EPSI optimisation. Data code: 6F-210316.

Since the cost function constructed here was relatively complicated, I carried out a reference grid search to confirm that it was working as intended (fig. 3.16). This yielded an expected optimum of $\alpha = 1.0004$. (The sample used here was ferulic acid, but in principle this should not matter as the drift is due to the spectrometer and not the sample.)

To save time in the optimisation itself, I reused the strategy of setting DS=0 and NS=1. As before, since each FE is separated by approximately five seconds of processing and spectrometer initialisation, this means that there is no need for another recovery delay; thus, D1 was set to 0.5 s. However, note that such a short recovery delay should *not* be used if DS is larger than 0 or NS larger than 1, as EPSI acquisitions should not be carried out in rapid succession. Furthermore, AQ should be kept under 100 ms (I used 91.8 ms here).

Optimisation results

In the event, all three optimisation algorithms successfully find the the optimal value of α in under a minute (table 3.12). This is *extremely* fast and the resulting value of α can be used for any subsequent ultrafast experiment, including the ultrafast PSYCHE-iDOSY experiment described in § 2.7.

However, it is worth clarifying one subtlety of POISE here. In the table, the optima are quoted to a rather greater precision than one might expect at first: however, the actual precision of the optimisation result is not given by the number of significant figures,^{*} but rather a combination

^{*}The entire concept of ‘significant figures’ is anyway a very crude method of expressing uncertainty.

Entry	Algorithm	Optimum found	FEs	Time taken (s)
1	NM	1.000375	10	49–52
2	MDS	1.000375	10	49–50
3	BOBYQA	1.000383–1.000388	7	34–35

Table 3.12: Results of EPSI gradient imbalance optimisations. Note that in practice, all of these optimisations actually converge to $\alpha = 1.0004$; the extra decimal place is not used by the spectrometer (see text for further discussion). The POISE routine used here is: `{"name": "epsi", "pars": ["cnst16"], "lb": [0.99], "ub": [1.01], "init": [1.0], "tol": [0.0001], "cf": "epsi_gradient_drift", "au": "poise_1d"}`. Data code: 6F-210316.

of two factors:

- first, the *optimisation tolerance*, which is purely determined by the optimisation routine. Thus, in table 3.12, the optimum of 1.000375 actually means 1.000375 ± 0.0001 (and not 1.000375 ± 0.0000005 , as ‘significant figures’ would imply); and
- second, the available *spectrometer resolution*, which represents how precisely the spectrometer can implement a given parameter value. In this case, the available resolution is 0.0001, which means that even if we input the ‘optimum’ of $\alpha = 1.000375$, the experiment is executed with $\alpha = 1.0004$ (see also fig. 3.17). The optimisation algorithm has no knowledge of this, so it still reports a value of 1.000375. **When reporting optima from POISE, users should be wary of any rounding performed by the spectrometer.**

This in fact further underlines the importance of choosing a sensible optimisation tolerance: it should be *at least* as large as the instrument resolution (larger values can be used when an extremely accurate result is not needed).

Returning to the current optimisation, the (k, t_2) data matrices obtained from the excitation–EPSI pulse sequence are shown in figs. 3.17a and 3.17b. In the former, where the default value of $\alpha = 1$ is used, there is a noticeable drift in k -space. This is removed in the optimised experiment with $\alpha = 1.0004$. The optimised value of α can then be transferred to other ultrafast experiment, such as the ultrafast TOCSY (fig. 3.15b). The improvements thus obtained are shown in figs. 3.17c and 3.17d: in the optimised spectrum, substantial improvements in lineshapes are obtained, even allowing some couplings to be properly resolved in the indirect dimension. In principle, this k -drift could be compensated for by shearing of the (k, t_2) data matrix prior to Fourier transformation. However, similar improvements in lineshapes could not be accomplished using any existing processing scripts, including those in the protocol of Giraudeau et al.;⁶⁸ I did not attempt to manually shear the data myself either.

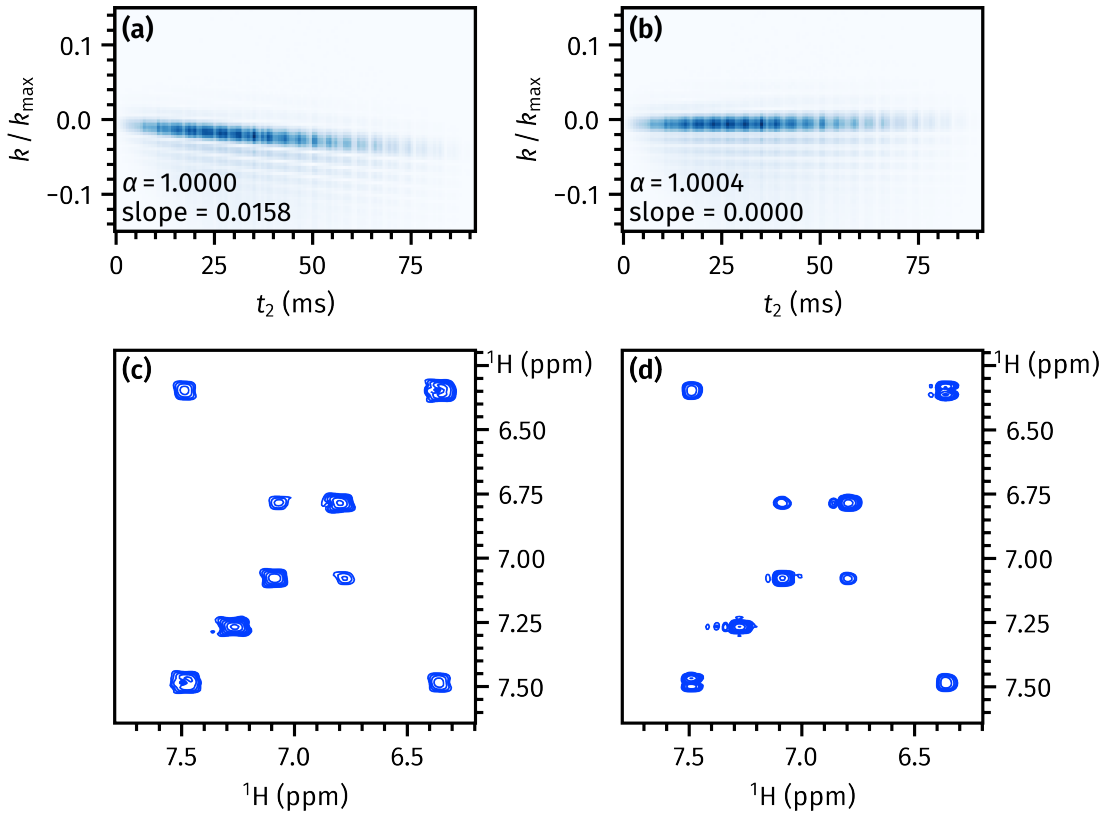


Figure 3.17: (a,b) Magnitude-mode (k, t_2) data matrices obtained from the pulse-EPSI experiment, before and after optimisation of α respectively. (c,d) The corresponding 2D ultrafast TOCSY spectra, recorded with the unoptimised and optimised value of α respectively. The MLEV mixing period used was 10 ms.

3.4.7 HMBC low-pass J-filter

In this and the remaining examples, we look at optimisations involving *multiple parameters*, one of the key strengths of POISE. Grid searches are extremely ineffective for this purpose, as the number of FEs required scales exponentially with the number of parameters; human trial-and-error can also be very difficult as parameters may be tightly *coupled*, meaning that the optimum cannot simply be found by optimising one parameter at a time. In contrast, the optimisation algorithms used in POISE make no assumption about the relationship between the two parameters. Furthermore, since only local minima are sought, the time required for convergence scales approximately linearly with the number of parameters, at least in the regimes which I have tested.*

In the simplest example of a multiple-parameter optimisation, I use POISE to search for the best delays in the low-pass J-filter (LPJF) of a HMBC experiment (fig. 3.18a). Specifically, a second-order LPJF is used, which has the form $\tau_1 - 90^\circ(^{13}\text{C}) - \tau_2 - 90^\circ(^{13}\text{C})$. The role of the LPJF

*I am not actually aware of any *theoretical* results which show how the number of function evaluations scales with the number of parameters in an optimisation; I suspect the answer depends on the exact problem being solved.

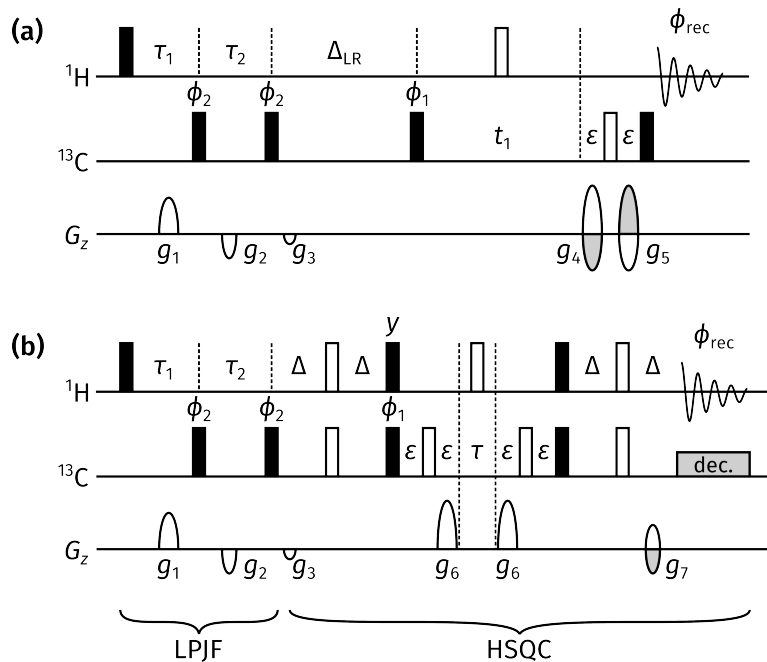


Figure 3.18: (a) Standard 2D HMBC pulse sequence with a second-order LPJF. (b) 1D LPJF-HSQC pulse sequence used in the POISE optimisation. In practice, only the first increment of this experiment was recorded. Delays are set as follows: $\Delta = 1/(4 \cdot ^1J_{CH})$; $\Delta_{LR} = 1/(2 \cdot ^nJ_{CH})$; τ is as short as possible (several microseconds); and τ_1 and τ_2 are as described in the text. Phase cycling is performed with $\phi_1 = \phi_{rec} = (x, -x)$, and $\phi_2 = (x, x, -x, -x)$. Gradient amplitudes are: $(g_1, g_2, g_3) = (15\%, -10\%, -5\%)$ for the LPJF, $(g_4, g_5) = (80\%, -47.85\%)$ (or vice versa) for EA selection in the HMBC experiment,⁷⁰ and $(g_6, g_7) = (70\%, \pm 35.21\%)$ for EA selection in the LPJF-HSQC experiment.

is to destroy $^1\text{H}^{\text{C}}$ magnetisation at the start of the HMBC, which would otherwise generate one-bond artefacts in the resulting spectrum. Assuming that the $^1J_{\text{CH}}$ values span a range of J_{\min} to J_{\max} , Sørensen and coworkers have provided ‘theoretical’ best values for the delays:^{71,72}

$$\tau_1 = \frac{1}{2} [J_{\min} + 0.146(J_{\max} - J_{\min})]^{-1}; \quad (3.31)$$

$$\tau_2 = \frac{1}{2} [J_{\max} - 0.146(J_{\max} - J_{\min})]^{-1}. \quad (3.32)$$

In practice, the TopSpin standard library HMBC sequence (`hmbcetgp12nd`) uses delays of

$$\tau_1 = \frac{1}{2J_{\min}}; \quad \tau_2 = \frac{1}{2J_{\max}}. \quad (3.33)$$

In either case, though, these delays neglect the actual *distribution* of $^1J_{\text{CH}}$: if many of the couplings are clustered around one particular value, then adjusting the delays to better suppress these values will yield better results. (Since $^1J_{\text{CH}}$ roughly correlates with the hybridisation of the ^{13}C atom,⁷³ this can easily happen, for example, if the molecule is predominantly aliphatic or predominantly aromatic.)

Optimisation setup

In order for the optimisation routine to minimise the artefacts, it must first have a cost function which can measure the extent of these artefacts. Given a full 2D HMBC spectrum, it is very difficult to discern which of the peaks are indeed artefacts (this is exactly why it is worthwhile to suppress them). However, by using a specially designed LPJF–HSQC experiment (fig. 3.18b), the artefacts can be isolated without any of the HMBC signals appearing: this works because any $^1\text{H}^{\text{C}}$ magnetisation which is *not* destroyed by the LPJF will be sampled in the HSQC segment which follows. Although the LPJF–HSQC can in principle be recorded as a full 2D experiment, to save time, I chose to only record the first increment as a 1D experiment. ^{13}C decoupling during acquisition is not mandatory, but the boost in SNR obtained through decoupling shortens the time needed for optimisation, and also reduces noise in the cost function.

We could directly perform an optimisation on the two delays τ_1 and τ_2 . However, for simplicity, I chose to reuse the definitions of τ_1 and τ_2 in the Bruker library sequence (eq. (3.33)) and instead optimise the parameters $J'_{\min} = 1/(2\tau_1)$ and $J'_{\max} = 1/(2\tau_2)$: these are represented in the pulse sequence as CNST25 and CNST26 respectively.* Both parameters were constrained to lie between 100 and 200 Hz, with a tolerance of 3 Hz. To match the magnitude-mode processing typically used for 2D HMBC experiments, the acquisition AU programme was modified to also process the LPJF–HSQC spectrum in magnitude mode. The integral of the entire spectrum can therefore be used as the desired cost function for the POISE optimisation. More specifically, I chose to use a sum of squares:

$$f_{\text{sos}} = \sum_i S_i^2, \quad (3.34)$$

which has the effect of penalising intense artefacts more strongly.

Optimisation results

In previous sections, I have quoted ranges for the optima found in an effort to show that the optimisation results were reproducible. In multiple-parameter optimisations, it is no longer appropriate to report separate ranges for both parameters, since the parameters are in general *interdependent*, in that the spread of values obtained for the first parameter depends on the value obtained for the second, and vice versa. I therefore only provide ranges for the number of FEs, as well as the time taken. As for optima found, I will quote only the *best* of these, as measured by the reduction in the cost function. These optima should therefore not be viewed as a complete summary of the five different runs, but rather as an *example* of the improvement which may be obtained through POISE.

* Prime symbols are used here because J_{\min} and J_{\max} are *physical constants* which depend on the sample being studied, whereas J'_{\min} and J'_{\max} are *parameters* which can be chosen to have any value.

Entry	Algorithm	Best optimum found			Aggregated results	
		J'_{\min}	J'_{\max}	$f_{\text{sos}}/10^7$	FEs	Time taken (s)
1	NM	143.07	185.71	8.65	20–24	358–430
2	MDS	141.73	185.82	8.36	18	321–323
3	BOBYQA	133.38	155.03	6.42	14–17	251–304

Table 3.13: Results of HMBC POISE optimisations. For each optimisation algorithm, the best optimum found is reported along with the value of the cost function f_{sos} (a lower value represents a better optimum). The number of FEs, as well as the time taken, is still reported as a range over the five optimisation runs performed per algorithm. The POISE routine used here is: `{"name": "lpjf", "pars": ["cnst25", "cnst26"], "lb": [100.0, 100.0], "ub": [200.0, 200.0], "init": [120.0, 180.0], "tol": [3.0, 3.0], "cf": "zerorealint_squared", "au": "poise_1d_mc"}`. Data code: 7Z-210814.

Interestingly, BOBYQA finds an optimum which is different from—and substantially better than—the other two algorithms. As mentioned previously, it is not generally possible to predict which optimum an algorithm will converge to, and this is complicated even further by the presence of noise, which imparts some stochastic character to the optimisation. Nevertheless, a closer inspection of the other optimisation runs actually shows that the better performance of BOBYQA on this problem is reproducible: the optima found from each of the five rounds, as well as a reference grid search, are plotted in fig. 3.19. It appears that the simplex-based methods get stuck in a suboptimal region, whereas BOBYQA is usually able to move leftwards and downwards into a more productive region. However, the exact reasons for this remain unclear. Also, BOBYQA does not always outperform NM and MDS in the other optimisations in this chapter (this is possibly a manifestation of the ‘no free lunch theorem’,⁷⁴ which states that when averaged over all problems, no optimisation algorithm strictly outperforms another).

In any case, when the best of these optimised values (i.e. entry 3 in table 3.13) were used to run a standard 2D HMBC, a significant reduction in one-bond artefacts was observed. It should be emphasised that these optimised values were obtained in a matter of several minutes, which is far shorter than a typical HMBC experiment requires. The performance of the optimised second-order LPJF (fig. 3.20b) was in fact almost comparable to that of a third-order LPJF (fig. 3.20c). However, using a third-order LPJF risks suppressing some of the desired HMBC signal as well, so if all else is equal, the optimised second-order LPJF should be preferred. In principle, the third-order LPJF itself can also be subjected to optimisation: this possibility was not explored in the present work, but would be easy to set up using the methods outlined in this section.

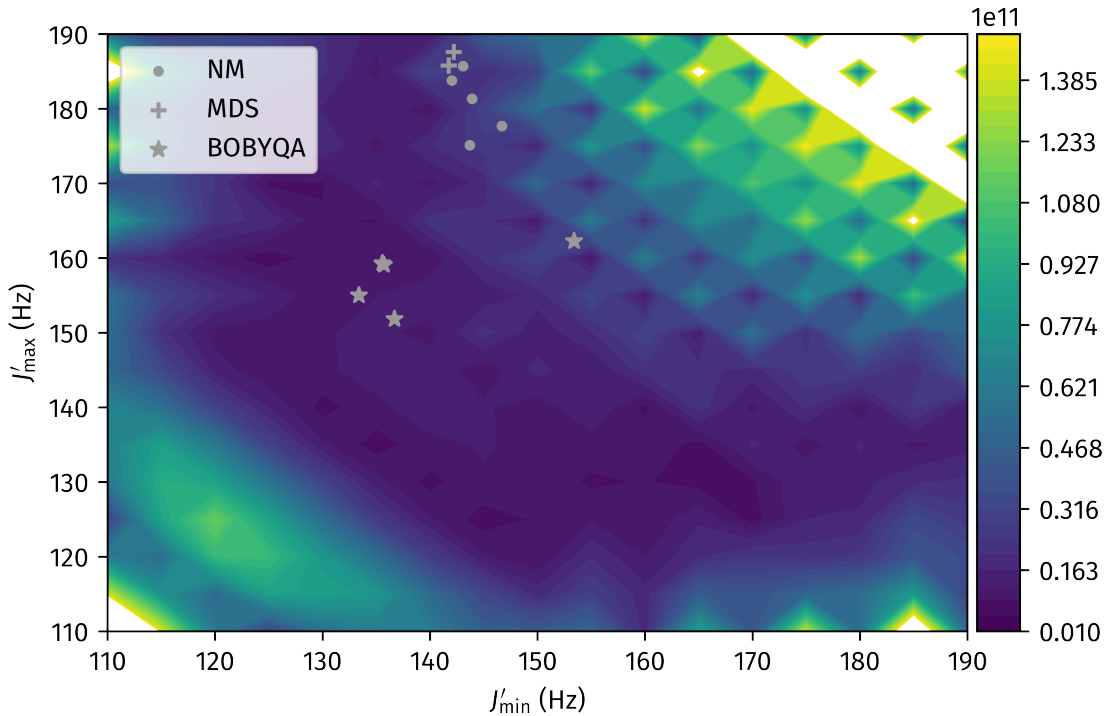


Figure 3.19: Reference grid search for HMBC optimisation. Darker regions correspond to lower cost function values, i.e. better spectra. The results of the 15 optimisations (5 per algorithm) are plotted on the same axes. Note that the cost functions here are rather larger than in table 3.13: this is solely because a different receiver gain was used. *Data code: 7Z-210814.*

Application to NOAH HMBC

In NOAH supersequences, the HMBC experiment (or ‘module’) is typically modified by replacing the initial 90° excitation pulse with a zz -filter (fig. 3.21).^{75,76} This causes the $^1\text{H}^{13}\text{C}$ magnetisation used in the HMBC to be excited, but leaves $^1\text{H}^{13}\text{C}$ magnetisation along the $+z$ axis so that it can be preserved for later modules in a supersequence. In an ideal situation, this means that the one-bond artefacts in the HMBC should in fact be suppressed even without an LPJF. However, significant one-bond artefacts are in fact still observed in NOAH HMBC spectra. I therefore sought to apply the same optimisation routine to the zz -HMBC module to see whether this could be improved.

Unfortunately, the results here were far less impactful: generally, the optimisation converged to values very close to the initial starting point, and in some cases the behaviour of the optimised spectrum was even (very slightly) worse than the unoptimised spectrum. (The deterioration is very marginal, and likely arises only due to noise in the cost function during the optimisation.) This strongly suggests that the artefacts are not due to imperfect LPJF suppression, but instead stem from the zz -filter component of the sequence. This was indeed verified experimentally: the addition of a ^{13}C 90° pulse at the end of the zz -filter led to much better one-bond artefact

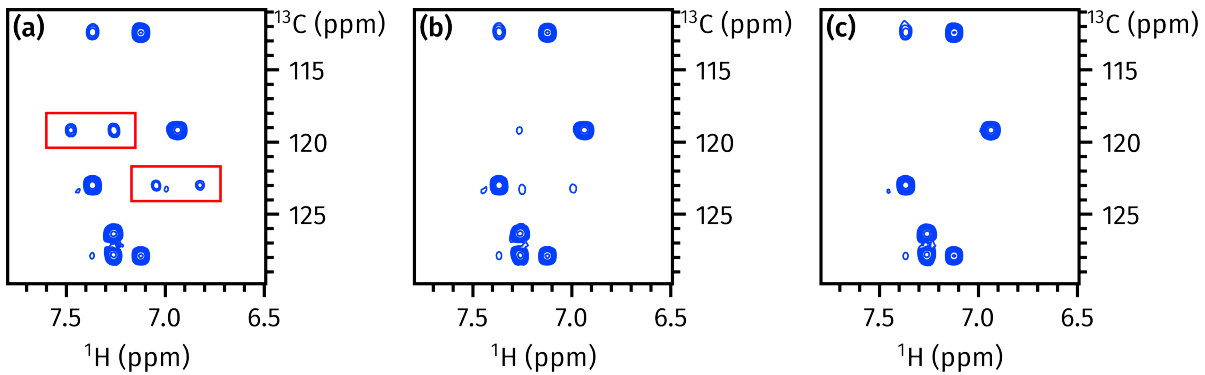


Figure 3.20: (a) HMBC spectrum with ‘compromise’ values of 120 and 180 Hz for J'_{\min} and J'_{\max} respectively in the second-order LPJF. The one-bond artefacts are highlighted in red boxes. (b) HMBC spectrum with optimised second-order LPJF delays. (c) HMBC spectrum with third-order LPJF. Data code: 7Z-210814.

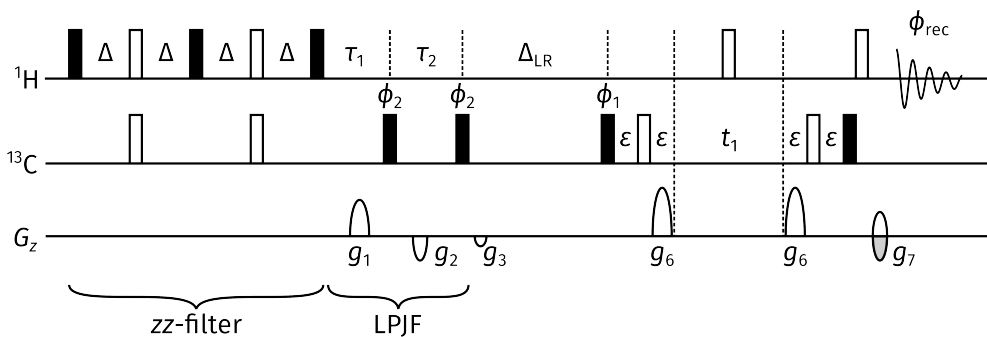


Figure 3.21: NOAH HMBC module.

suppression (fig. 3.22c; this strategy is further explained in § 4.3.7).

3.4.8 PSYCHE pure shift NMR

In § 2.4, I described the motivation behind, and early attempts towards, the optimisation of PSYCHE pure shift spectra. The content in this section is similar, except that it was performed within the framework of POISE.

Optimisation setup

In this section, the standard double-saltire PSYCHE pure shift element was used (fig. 3.23a).^{77,78} As described in § 2.4.1, the PSYCHE PSE can be described using six parameters; in this section, we investigate only three of these, namely the amplitude (i.e. flip angle β), bandwidth ΔF , and duration τ_p . In addition to this, the amplitude of the weak gradient during the PSE g_3 was also chosen as a fourth parameter to vary. As before, the quality of the PSE is evaluated using a JRSE experiment (fig. 3.23b), which is then compared against a pulse–acquire spectrum: the cost function used is f_{diff} .

Entry	Algorithm	Best optimum found			Aggregated results	
		J'_{\min}	J'_{\max}	$f_{\text{sos}}/10^9$	FEs	Time taken (s)
1	NM	117.05	188.59	5.37	25–29	451–523
2	MDS	125.21	193.16	5.35	14–17	249–310
3	BOBYQA	119.49	182.74	5.78	9–16	164–293

Table 3.14: Results of POISE optimisations on the LPJF in the NOAH HMBC module. All optimisation details are the same as in table 3.13, except that the LPJF-HSQC pulse sequence used for the optimisation was modified accordingly to include a *zz*-filter. Data code: 7Z-210814.

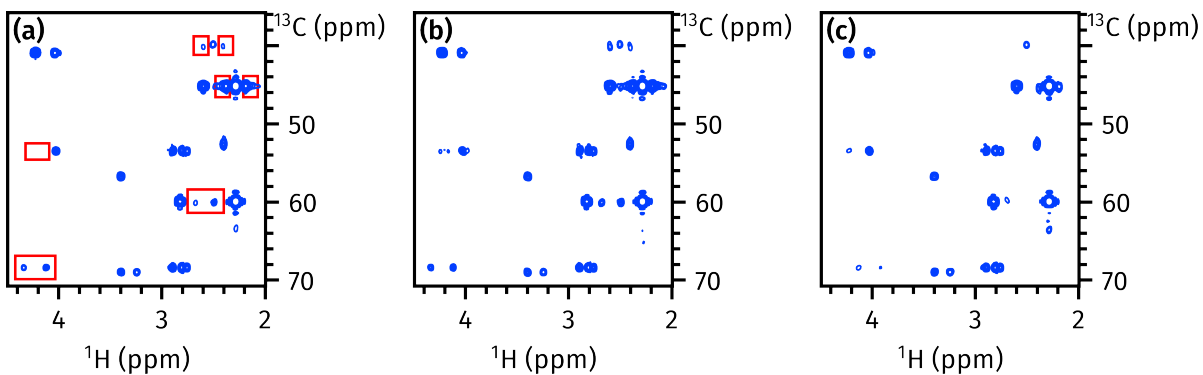


Figure 3.22: (a) NOAH HMBC spectrum before POISE optimisation, using the default settings of 120 and 180 Hz. One-bond artefacts are highlighted in red boxes. (b) NOAH HMBC spectrum after POISE ‘optimisation’ (using $J'_{\min} = 125.2$ Hz, $J'_{\max} = 193.2$ Hz, as per entry 2 of table 3.14). (c) NOAH HMBC spectrum with no POISE optimisation, but instead adding a 90° ^{13}C pulse at the end of the *zz*-filter (see also § 4.3.7). Data code: 7Z-210814.

In the JRSE pulse sequence used for optimisation, the four parameters (flip angle, bandwidth, duration, and gradient amplitude) are respectively CNST20, CNST21, P40, and GPZ10. There is, however, a slight complication: whenever the bandwidth or duration is changed, the entire pulse must be re-created, because the *x*- and *y*-coefficients depend on these parameters. Therefore, a (TopSpin) Python script was written to generate the double saltire pulse using the parameters above, and the POISE AU programme was modified to call this Python script before acquiring the spectrum. Although the overall setup may perhaps be slightly confusing,^{*} I contend that this is an excellent demonstration of the customisability that POISE provides.

One slightly odd behaviour noted with the PSYCHE optimisations was that at least one dummy scan was required for the optimisation to be robust: if no dummy scans were used, the *very first* FE in an optimisation run would yield a rather unreliable result, although subsequent FEs were unaffected. It is not obvious why this is the case, as FEs are already separated by a delay of

^{*}We have here a Python script (the POISE frontend) calling an AU programme (for acquisition) which calls a Python script (to make the double saltire).

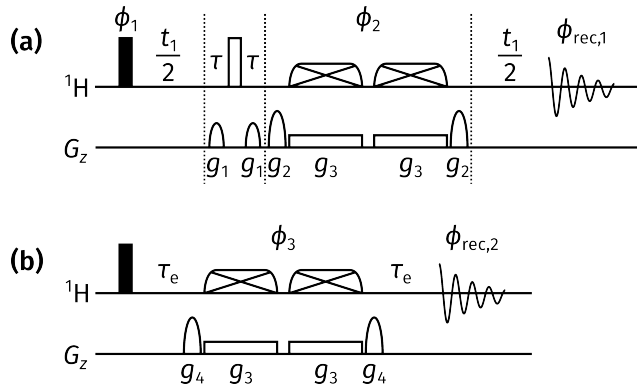


Figure 3.23: (a) Pseudo-2D PSYCHE pure shift experiment. (b) J-resolved spin echo experiment (see also § 2.4.1) using the PSYCHE PSE. Phase cycling is performed using: $\phi_1 = (x, -x)$, $\phi_2 = (x, x, y, y)$, $\phi_{rec,1} = (x, -x, -x, x)$ for the full PSYCHE experiment, and $\phi_3 = (x, y, -x, -y)$, $\phi_{rec,2} = (x, -x, x, -x)$ for the JRSE experiment. CTP gradient amplitudes are $(g_1, g_2, g_4) = (35\%, 77\%, 50\%)$ (though the exact values are likely immaterial); the PSYCHE gradient g_3 was subjected to optimisation. The delay τ is set to $1/(4 \cdot T_{\text{chunk}})$ for the PSYCHE experiment, and τ_e is 16 ms.

several seconds, which is on the order of $5T_1$: thus, each FE should be starting from (almost) full equilibrium magnetisation. Nevertheless, to avoid this issue, all optimisations in this section were run using DS=1 and NS=2.

The sample used was andrographolide: this is the same sample as was used in the dPSYCHE section (§ 2.6), and is fairly useful for pure shift studies as it contains quite diverse coupling patterns. The spectral region was limited to 1.15 ppm and above to exclude strong singlets from the methyl groups in andrographolide, which are irrelevant to pure shift NMR but disproportionately influence the value of the cost function (this issue was also discussed in § 2.4.3).

Optimisation results

I ran several different optimisations of increasing complexity:

- one-parameter: flip angle only
- two-parameter: flip angle and gradient amplitude
- three-parameter: flip angle, gradient amplitude, and bandwidth
- four-parameter: flip angle, gradient amplitude, bandwidth, and duration

Table 3.15 summarises the results obtained from all optimisations, whereas tables 3.16 to 3.19 show more detailed results for each individual set of optimisations, including a breakdown by algorithm. We can see from these results that—perhaps unsurprisingly—optimising more parameters leads to greater improvements in the cost function, albeit at the cost of more FEs and more time.

Description	Aggregated results from all runs			Parameters from best optimum			
	f_{diff}	FEs	Time (s)	β ($^{\circ}$)	g_3 (%)	ΔF (kHz)	τ_p (ms)
Initial point	0.353	–	–	(25.0)	(2.00)	(10.0)	(30.0)
1 parameter	0.340–0.343	5–10	84–168	17.5	(2.00)	(10.0)	(30.0)
2 parameters	0.325–0.338	12–33	205–565	16.5	1.00	(10.0)	(30.0)
3 parameters	0.320–0.328	18–77	315–1344	17.4	1.73	15.9	(30.0)
4 parameters	0.316–0.333	39–85	705–1504	13.9	1.45	15.9	36.0

Table 3.15: Summary of all optimisations performed on the PSYCHE PSE. Unlike before, aggregated results are not grouped by optimisation algorithm; these ranges are therefore collected from a total of 15 runs (5 per algorithm). Parameters in parentheses were not optimised and were simply carried over from the initial point. A detailed breakdown of the results by algorithm, as well as the POISE routines used, are described in tables 3.16 to 3.19. Data code: 6A-200822.

Entry	Algorithm	Optimum found ($^{\circ}$)	FEs	Time (s)
1	NM	15.0–20.0	8–10	135–168
2	MDS	17.5–20.0	8	133–136
3	BOBYQA	18.4–19.9	5	84–85

Table 3.16: PSYCHE one-parameter (flip angle) optimisations. The POISE routine used was: `{"name": "psyche1", "pars": ["cnst20"], "lb": [10.0], "ub": [35.0], "init": [25.0], "tol": [2.0], "cf": "specdiff", "au": "poise_1d"}`. Data code: 6A-200822.

Entry	Algorithm	Best optimum found			Aggregated results	
		β ($^{\circ}$)	g_3 (%)	f_{diff}	FEs	Time (s)
1	NM	16.48	1.00	0.3253	18–24	307–410
2	MDS	18.47	0.82	0.3261	25–33	424–565
3	BOBYQA	15.77	0.96	0.3279	12–16	205–271

Table 3.17: PSYCHE two-parameter (flip angle and gradient amplitude) optimisations. The POISE routine used was: `{"name": "psyche2", "pars": ["cnst20", "gpz10"], "lb": [10.0, 0.2], "ub": [35.0, 5.0], "init": [25.0, 2.0], "tol": [2.0, 0.2], "cf": "specdiff", "au": "poise_1d"}`. Data code: 6A-200822.

Entry	Algorithm	Best optimum found				Aggregated results	
		β (°)	g_3 (%)	ΔF (kHz)	f_{diff}	FEs	Time (s)
1	NM	17.40	1.73	15.93	0.3196	33–43	576–770
2	MDS	16.75	1.71	16.33	0.3215	46–77	804–1344
3	BOBYQA	16.48	1.65	16.71	0.3200	18–31	315–540

Table 3.18: PSYCHE three-parameter (flip angle, gradient amplitude, and bandwidth) optimisations. The POISE routine used was: `{"name": "psyche3", "pars": ["cnst20", "gpz10", "cnst21"], "lb": [10.0, 0.2, 1000.0], "ub": [35.0, 5.0, 20000.0], "init": [25.0, 2.0, 10000.0], "tol": [2.0, 0.2, 500.0], "cf": "specdiff", "au": "poise_psych"}`. Data code: 6A-200822.

Entry	Algorithm	Best optimum found				Aggregated results	
		β (°)	g_3 (%)	ΔF (kHz)	τ_p (ms)	f_{diff}	FEs
1	NM	13.87	1.45	15.93	36.00	0.3163	40–47
2	MDS	19.15	1.06	11.19	36.52	0.3185	57–85
3	BOBYQA	17.59	1.20	12.76	43.06	0.3178	39–62
							733–845
							1006–1504
							705–1130

Table 3.19: PSYCHE four-parameter (flip angle, gradient amplitude, bandwidth, and pulse duration) optimisations. The POISE routine used was: `{"name": "psyche4", "pars": ["cnst20", "gpz10", "cnst21", "p40"], "lb": [10.0, 0.2, 1000.0, 5000.0], "ub": [35.0, 5.0, 20000.0, 75000.0], "init": [25.0, 2.0, 10000.0, 30000.0], "tol": [2.0, 0.2, 500.0, 2000.0], "cf": "specdiff", "au": "poise_psych"}`. Data code: 6A-200822.

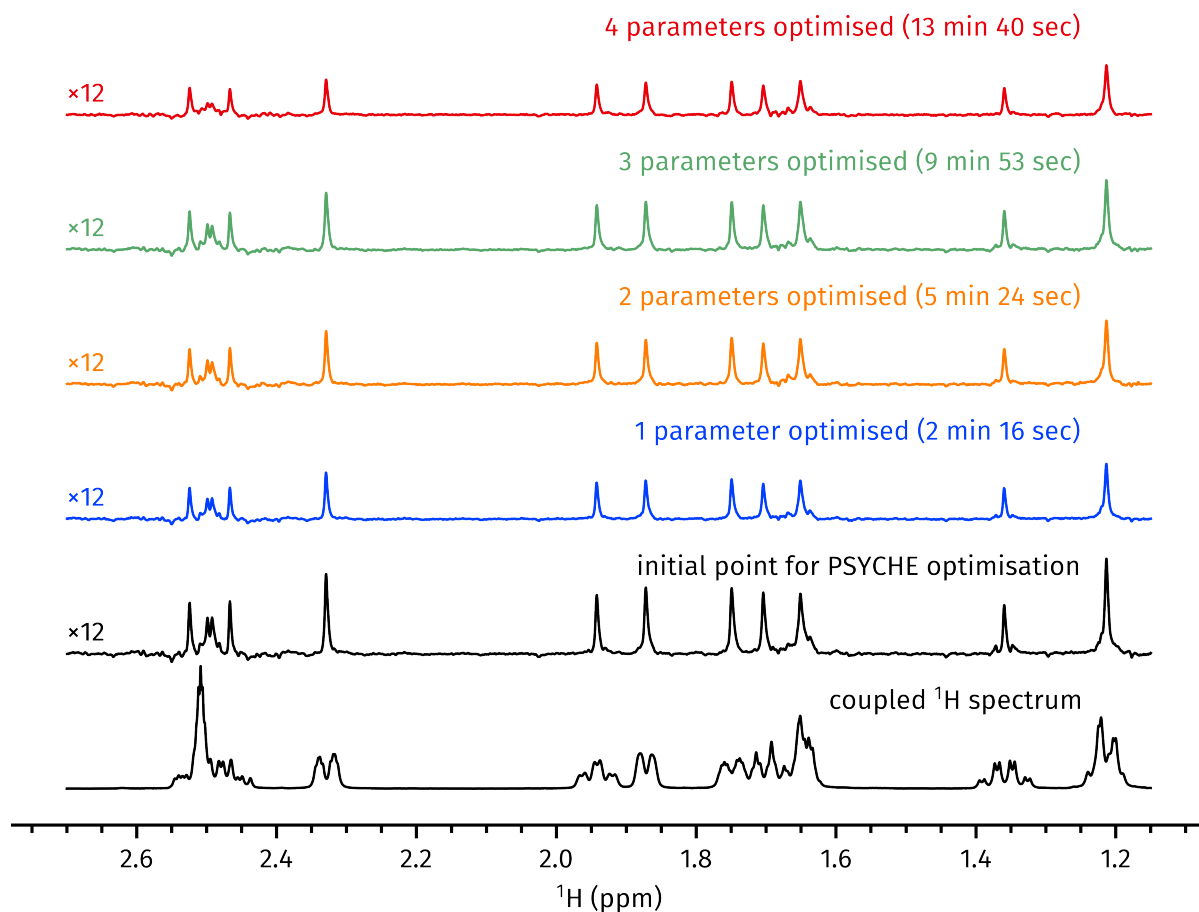


Figure 3.24: Insets of PSYCHE spectra obtained using the best optimum from each set of optimisations in table 3.15; the time required to obtain the optimised parameters is also indicated for each spectrum. The original coupled ^1H spectrum is shown as a reference. Data code: 6A-200823.

An important question is whether these reductions in the cost function (measured on a JRSE experiment) do actually translate into improved pure shift spectra. The pure shift spectra, obtained with the optimised parameters in table 3.15, are shown in fig. 3.24. Of particular interest are the two strongly coupled protons at 2.5 ppm: in the pure shift spectrum, the two peaks on either side are genuine, but strong coupling artefacts appear in the middle of these peaks. On top of that, the decoupling performance for the peaks at 1.36 ppm and 1.65 ppm are also worth noting.

Broadly speaking, all the optimised spectra provide better performance than the initial point in terms of decoupling quality. In particular, the four-parameter optimisation successfully suppresses some of the strong coupling artefacts: this may be explained by the fact that the pulse duration was increased from 30 ms to 36 ms, which generally results in better spatiotemporal

averaging, as was noted in § 2.3.* Generally, this optimisation *does* come at a cost in sensitivity, which is particularly noticeable for the four-parameter optimised result. However, since the cost function f_{diff} also penalises sensitivity losses, it ensures that any drops in sensitivity are not excessive.

Given these improvements, one might wonder why it would ever be worth optimising fewer than four parameters. The first obvious drawback is the time required: in fact, it is probably more economical to use the TSE-PSYCHE experiment (which also has improved performance in the presence of strong coupling).† Furthermore, f_{diff} is a rather ‘flat’ cost function and optimisations using it are very susceptible to noise: this factor is specific to pure shift optimisations, and was previously mentioned in § 2.4.3. A closer inspection of the POISE logs for the three- and four-parameter optimisations (which are not provided here, but can be accessed in the raw data for the POISE paper, available at <https://doi.org/10.5281/zenodo.4698423>) reveals that the optimisations are perhaps not as consistent as one would hope. Although *generally* the optimisation leads to similar results in that (for example) β is often decreased and τ_p increased, the extents of these changes are not quite uniform: for example, τ_p is optimised to anywhere between 36 ms and 55 ms, which suggests that there are *many* local minima and which of these the optimisation converges to depends on the noise in the cost function.‡

Putting all of these factors together, it is (in my opinion) only really worthwhile to optimise two parameters at a time: either the flip angle plus gradient amplitude, or the flip angle plus the duration, appear to be sensible choices. That said, this *does* demonstrate that optimising multiple parameters at once—ordinarily a very challenging task for a human—does not actually require prohibitively long times when performed using a suitable algorithm.

3.4.9 Water suppression

The suppression of solvent signals is one of the most important techniques in modern NMR, especially in cases where solvent deuteration is not a viable strategy (e.g. when exchangeable protons must be observed).^{79–81} Here, we specifically seek to optimise the 1D NOESY experiment with presaturation⁸² (fig. 3.25): although this sequence is popular, its performance can depend on at least four parameters, namely the transmitter offset, presaturation power, the NOE mixing time, and the presaturation duration or recovery delay. In this section, I use the symbols v_{tx} , k_{ps} ,

*This does raise the question of whether a two-parameter optimisation of just the flip angle and duration could yield similar results, but in a shorter time. I think it is possible, but I did not test this, though, as I simply did not have the appetite to try optimising *every* possible combination of parameters.

†Another thing I did not (thoroughly) investigate was to optimise the TSE-PSYCHE experiment. This could, in principle, be done using a TSE form of the JRSE experiment: 90°–chirp– τ –PSE– τ –chirp–detect. However, this has decreased sensitivity over the original JRSE experiment, which means even longer optimisations.

‡I still think, however, that it is misleading to quote these ranges in tables 3.16 to 3.19, because this gives the impression that there are no correlations between the optimised parameters.

τ_m , and τ_r to respectively refer to these four parameters: the corresponding TopSpin parameters are O1, CNST20, D8, and D1.

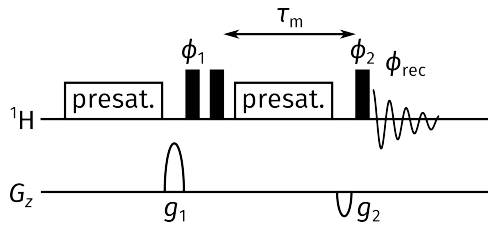


Figure 3.25: 1D NOESY pulse sequence used for water suppression. Presaturation of the water resonance is applied during the recovery delay as well as the NOE mixing time. Phase cycling is performed with $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x, y, y, -y, -y)$, and $\phi_{\text{rec}} = (x, -x, -x, x, y, -y, -y, y)$. Gradient amplitudes are $(g_1, g_2) = (50\%, -10\%)$. Boxes labelled with ‘presat.’ indicates periods of weak presaturation.

Optimisation setup

In a similar style to the PSYCHE optimisations (§ 3.4.8), a series of optimisations with increasing numbers of parameters were run. However, in this case, the cost function is far easier to construct and far less noisy: we can simply integrate the water peak, which was defined to be the region of the spectrum between 4.65 and 4.75 ppm, and use that (or its absolute value) as the cost function. In practice, because the phase of the water peak can vary rather unpredictably, I preferred to reuse the sum-of-squares cost function f_{sos} (eq. (3.34)). Squaring each point of the spectrum not only accounts for the possible sign changes from out-of-phase peaks, but also more strongly penalises intense absorption-mode water peaks.*

One drawback of the water suppression optimisation is that the phase cycle of the 1D NOESY is integral to its performance: thus, several scans must be used for a reliable cost function to be obtained, making for relatively long optimisations. In this case, I used DS=2 and NS=4.

For these optimisations, I used a sample of rodent urine in D_2O , which was kindly provided by Abi Yates and Fay Probert (both University of Oxford).

Optimisation results

As before, I first provide a ‘summary’ table comparing the optimisations with different numbers of parameters (table 3.20). From this, we see a similar story to before, in that optimising more parameters takes more time but leads to larger decreases in the cost function. Individual details of each set of optimisations are given in tables 3.21 to 3.24.

*Dispersion-mode peaks are possibly more problematic because of their long ‘tails’. In hindsight, I could have considered trying the `minabsint` cost function instead (eq. (3.19)). However, in practice this did not seem to matter very much: all of these cost functions do ultimately share the same goal, i.e. to reduce the intensity of the water peak.

Description	Aggregated results from all runs			Parameters from best optimum			
	$f_{\text{sos}}/10^{18}$	FEs	Time (s)	ν_{tx} (Hz)	k_{ps} (Hz)	τ_m (s)	τ_r (s)
Initial point	14.7	–	–	(1880.61)	(50.0)	(0.100)	(2.00)
1 parameter	1.85–2.49	6–12	259–520	1880.41	(50.0)	(0.100)	(2.00)
2 parameters	1.21–9.68	9–21	390–911	1880.20	51.94	(0.100)	(2.00)
3 parameters	1.24–4.20	19–26	825–1128	1879.90	47.79	0.118	(2.00)
4 parameters	0.165–1.65	25–53	1143–2314	1881.10	53.28	0.150	3.00

Table 3.20: Summary of optimisations on 1D NOESY / presaturation pulse sequence for water suppression. Parameters in parentheses were not optimised and were simply carried over from the initial point. A detailed breakdown of the results by algorithm, as well as the POISE routines used, are described in tables 3.21 to 3.24. Data code: 4P-210620.

Entry	Algorithm	Optimum found (Hz)	FEs	Time taken (s)
1	NM	1880.24–1880.49	10–12	436–519
2	MDS	1880.24–1880.36	12	518–520
3	BOBYQA	1880.34–1880.47	6–7	259–303

Table 3.21: Water suppression one-parameter (transmitter offset) optimisations. The POISE routine used was: `{"name": "solvsupp1", "pars": ["o1"], "lb": [1870.61], "ub": [1890.61], "init": [1880.61], "tol": [0.2], "cf": "zerorealint_squared", "au": "poise_1d_noapk"}`. Data code: 4P-210620.

The corresponding spectra post-optimisation are shown in fig. 3.26. Clearly, the optimisations succeed in reducing the size of the water peak, especially the four-parameter optimisation. However, perhaps equally importantly, the doublet at 4.56 ppm is unaffected by the optimisation. Since the cost function does not actually check for the retention of peaks outside the target window, it is in principle possible that the optimisation will converge to parameters which achieve excellent water suppression but also cause nearby peaks to be lost. This possibility is averted here by setting a conservative upper bound of 55 Hz on the presaturation power k_{ps} : this avoids inadvertently saturating nearby resonances.

In fact, it may be justifiable in this case to run a long, four-parameter optimisation (which takes around 30 minutes), especially if many samples are to be run using the same optimised water suppression parameters. However, for more routine usage, it is probably preferable to limit the number of parameters being optimised to two.

Optimisation on a sucrose sample

The water suppression optimisations shown above were in fact first developed and evaluated on a sample of sucrose in 90% H₂O/10% D₂O. This sample is less ‘interesting’ in that there are no resonances close to the water peak, and I also did not perform replicates of each optimisation run (this data never made it into the POISE paper). Nevertheless, I still believe it is still worth adding

Entry	Algorithm	Best optimum found			Aggregated results	
		ν_{tx} (Hz)	k_{ps} (Hz)	$f_{sos}/10^{18}$	FEs	Time (s)
1	NM	1880.20	51.94	1.209	17–21	737–911
2	MDS	1880.85	50.81	2.600	15	649–652
3	BOBYQA	1881.12	50.03	2.887	9–14	390–608

Table 3.22: Water suppression two-parameter (transmitter offset and presaturation power) optimisations. The POISE routine used was: `{"name": "solvsupp2", "pars": ["o1", "cnst20"], "lb": [1870.61, 10.0], "ub": [1890.61, 55.0], "init": [1880.61, 50.0], "tol": [0.2, 2.5], "cf": "zerorealint_squared", "au": "poise_1d_noapk"}`. Data code: 4P-210620.

Entry	Algorithm	Best optimum found				Aggregated results	
		ν_{tx} (Hz)	k_{ps} (Hz)	τ_m (ms)	$f_{sos}/10^{18}$	FEs	Time (s)
1	NM	1879.90	47.79	117.7	1.238	23–26	1002–1128
2	MDS	1882.79	52.95	49.91	1.843	24–25	1034–1080
3	BOBYQA	1881.85	52.14	134.7	1.738	19–26	825–1133

Table 3.23: Water suppression three-parameter (transmitter offset, presaturation power, and mixing time) optimisations. The POISE routine used was: `{"name": "solvsupp3", "pars": ["o1", "cnst20", "d8"], "lb": [1870.61, 10.0, 0.010], "ub": [1890.61, 55.0, 0.150], "init": [1880.61, 50.0, 0.100], "tol": [0.2, 2.5, 0.010], "cf": "zerorealint_squared", "au": "poise_1d_noapk"}`. Data code: 4P-210620.

Entry	Algorithm	Best optimum found					Aggregated results	
		ν_{tx} (Hz)	k_{ps} (Hz)	τ_m (ms)	τ_r (s)	$f_{sos}/10^{18}$	FEs	Time (s)
1	NM	1880.72	49.73	110.6	1.972	1.008	29–40	1250–1726
2	MDS	1882.65	53.41	69.46	2.137	1.199	34–53	1487–2314
3	BOBYQA	1881.10	53.28	150.0	3.000	0.1653	25–40	1143–1843

Table 3.24: Water suppression four-parameter (transmitter offset, presaturation power, mixing time, and presaturation duration) optimisations. The POISE routine used was: `{"name": "solvsupp4", "pars": ["o1", "cnst20", "d8", "d1"], "lb": [1870.61, 10.0, 0.010, 1.0], "ub": [1890.61, 55.0, 0.150, 3.0], "init": [1880.61, 50.0, 0.100, 2.0], "tol": [0.2, 2.5, 0.010, 0.1], "cf": "zerorealint_squared", "au": "poise_1d_noapk"}`. Data code: 4P-210620.

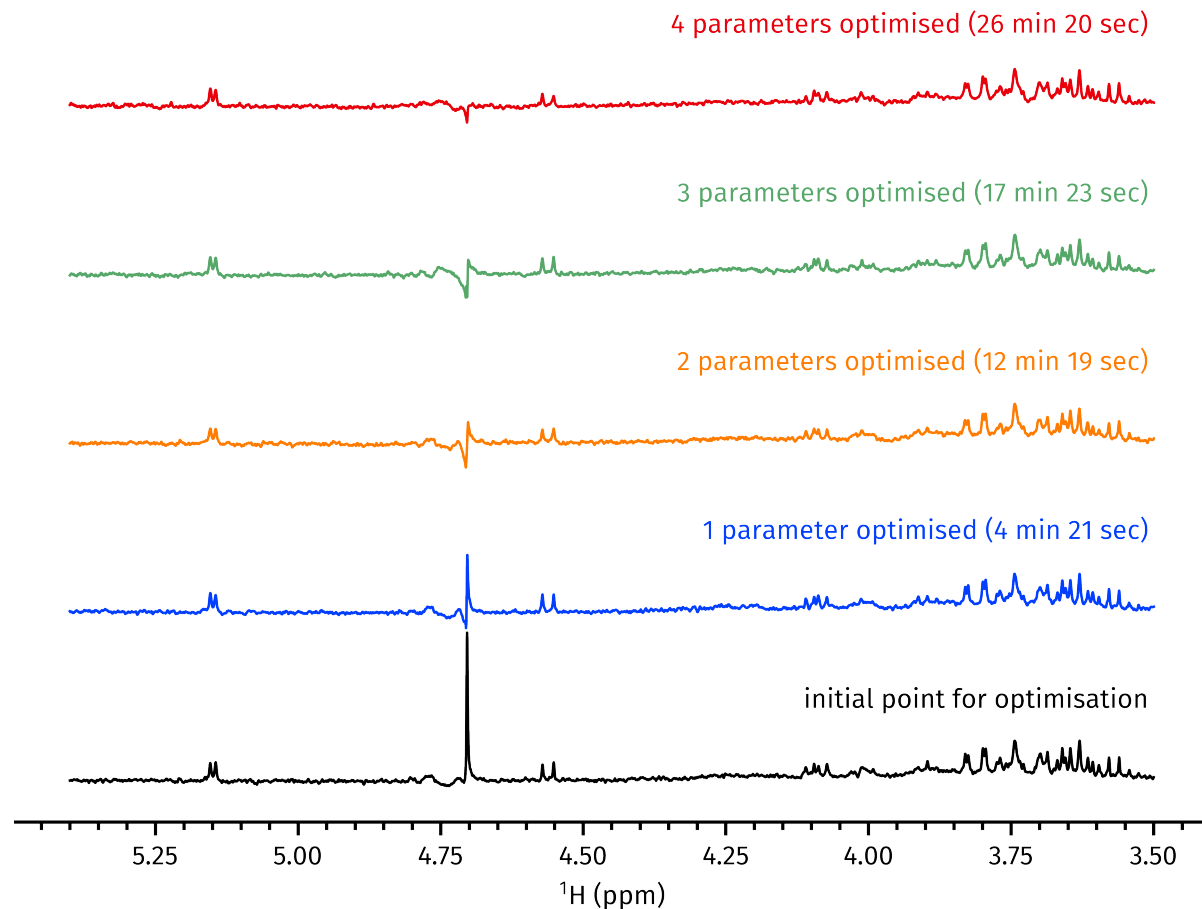


Figure 3.26: Insets of 1D NOESY spectra acquired using optimised parameters from each set of optimisations in table 3.15; the time required to obtain the optimised parameters is also indicated for each spectrum. The sample used was rodent urine in D_2O . Note that, for the three- and four-parameter optimisations, the best optimum from the 15 runs (i.e. the optima listed in table 3.20) was used to acquire the spectra shown here. However, this was not the case for the two-parameter optimisation: I (inexplicably) used a different optimum. This was probably an oversight on my part. Nevertheless, it does not affect any of the conclusions drawn here. *Data code:* 4P-210620.

the results here, as they demonstrate that the optimisation procedure can be easily applied to different samples.

In this case, the one-parameter optimisation seemed to have no significant effect on the result. However, the two- through four-parameter optimisations all yielded significant improvements in the water suppression (fig. 3.27). The parameters used for each of these spectra are listed in table 3.25.

3.4.10 Diffusion NMR

As the final example of a POISE optimisation, I demonstrate its application to diffusion-ordered spectroscopy (DOSY) experiments.⁸³ DOSY experiments measure molecular diffusion during a

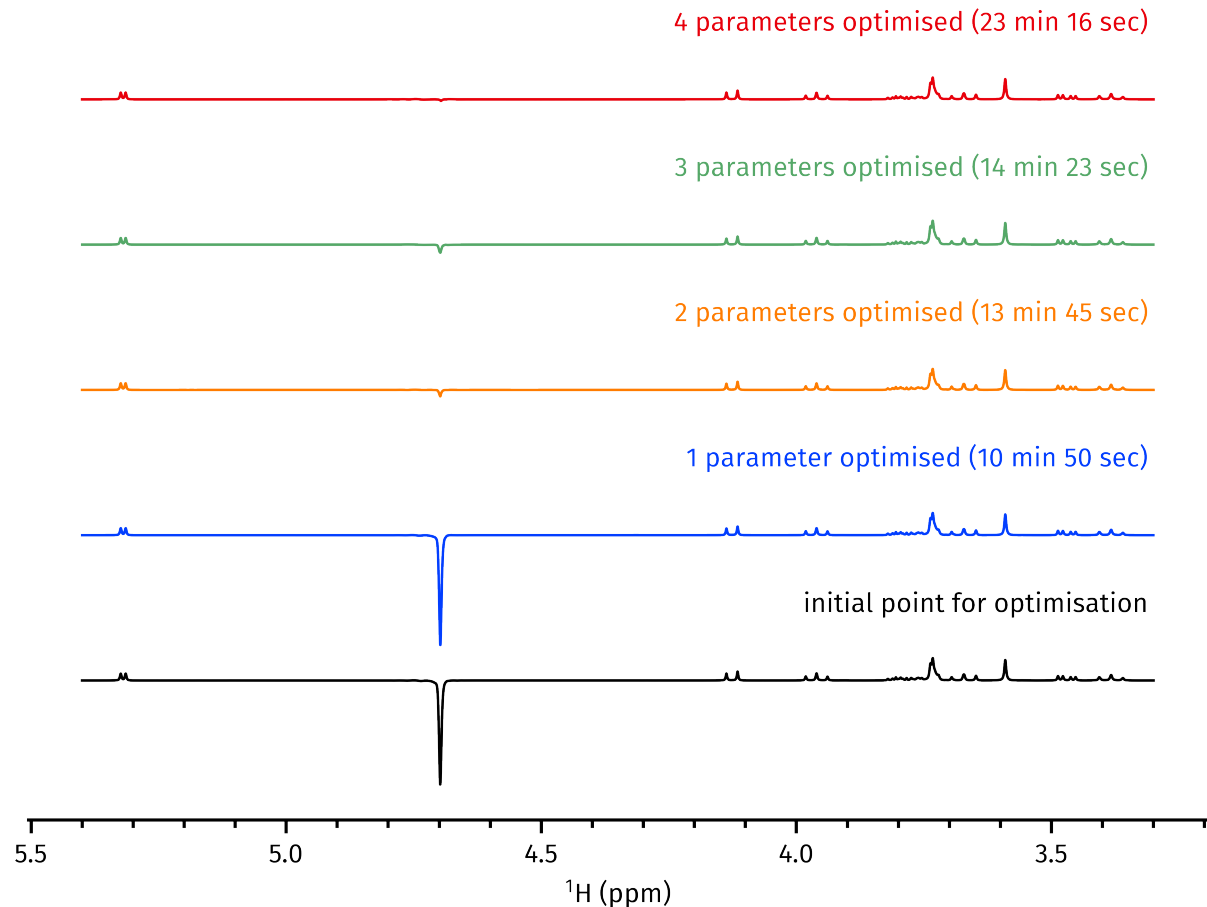


Figure 3.27: Insets of 1D NOESY spectra acquired after optimisation of 1, 2, 3, or 4 parameters (the exact parameters being optimised are the same as in fig. 3.26). The sample used was sucrose in 90% H₂O/10% D₂O. The time required to obtain the optimised parameters is also indicated for each spectrum. The parameters used for each of these spectra are in table 3.25. Data code: 4S-210617.

Description	$f_{\text{sos}}/10^{10}$	FEs	Time (s)	ν_{tx} (Hz)	k_{ps} (Hz)	τ_m (s)	τ_r (s)
Initial point	7.74	–	–	(1880.61)	(50.0)	(0.100)	(2.00)
1 parameter	7.25	15	650	1882.11	(50.0)	(0.100)	(2.00)
2 parameters	3.65	19	825	1881.50	75.58	(0.100)	(2.00)
3 parameters	5.71	20	863	1881.40	78.48	0.045	(2.00)
4 parameters	4.62	29	1396	1881.09	58.54	0.107	2.91

Table 3.25: Optimised water suppression parameters for a sample of sucrose in 90% H₂O/10% D₂O. Parameters in parentheses were not optimised and were simply carried over from the initial point. These correspond to the spectra in fig. 3.27. Data code: 4S-210617.

delay period, Δ , which is placed between two gradients, such as in the stimulated echo experiment (fig. 3.28a), or the bipolar pulse pair version (fig. 3.28b) which uses opposing gradients to refocus the lock signal during the sequence. In these sequences, the first ‘encoding’ gradient (or set thereof) imparts a spatially-dependent phase, which—in the absence of diffusion—is perfectly

refocused by the second ‘decoding’ gradient (or set thereof). However, if diffusion is present, this refocusing is not complete, leading to signal attenuation which is described by the Stejskal–Tanner equation:

$$I(G) = I(0) \exp [-(\gamma \delta G)^2 D \Delta']. \quad (3.35)$$

Here, G is the amplitude of the encoding and decoding gradients, δ the duration of the gradients, and Δ' is a corrected diffusion delay.

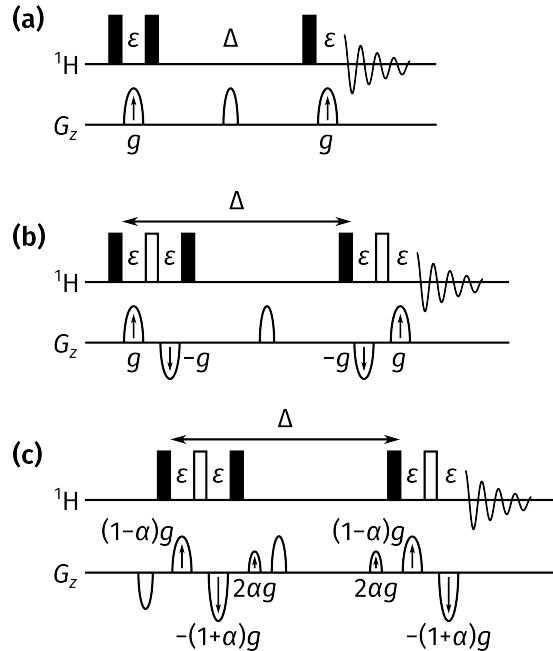


Figure 3.28: (a) Stimulated echo DOSY pulse sequence. The diffusion-encoding and -decoding gradients, whose amplitudes are incremented, are marked with arrows. (b) Stimulated echo bipolar pulse pair DOSY pulse sequence. (c) Oneshot DOSY pulse sequence.

In a pseudo-2D DOSY experiment, each increment is recorded with a different gradient amplitude G . The value of D can then be extracted through various means, most simply an exponential fitting of the measured intensities $I(G)$ for single species (or distinct species which do not overlap). In order to minimise the uncertainty in the calculated diffusion coefficients, the parameters Δ and δ , as well as the range of gradient amplitudes used (and possibly also their distribution), should be chosen in order to yield a diffusion profile of the form in fig. 3.29b. Generally, the last increment (acquired using the maximum gradient strength, G_{\max}) should have an attenuation of 5–35%* when compared against the first increment (acquired with the minimum gradient amplitude, G_{\min} , which may be 0). Traditionally, this is done ‘by hand’ by acquiring individual increments of the pseudo-2D diffusion experiment, checking the attenuation, and adjusting the parameters accordingly.^{83,84}

* Recommendations tend to vary.

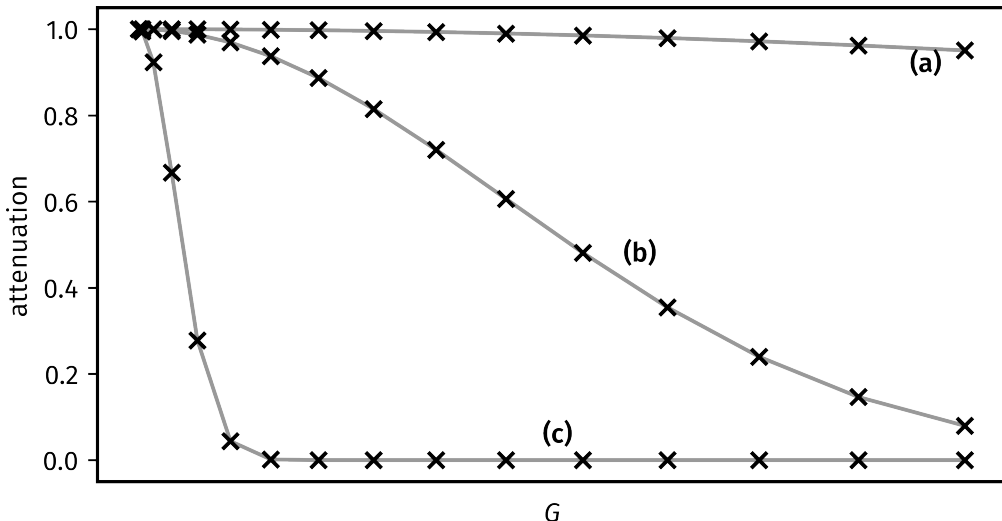


Figure 3.29: Simulated diffusion profiles for species with different diffusion rates. (a) A slowly diffusing species: insufficient attenuation is observed with the given parameters, meaning that generally Δ or δ need to be increased. (b) An ‘ideal’ diffusion profile where attenuation is neither too little or too much. (c) A rapidly diffusing species, for which complete attenuation is already accomplished using low gradient amplitudes. This generally suggests that Δ or δ need to be decreased, or a new gradient ramp calculated.

This task could of course be delegated to a computer programme such as POISE. We could simply tackle it head-on by optimising all of Δ , δ , and G_{\max} at once (assuming that G_{\min} is fixed); however, this is actually rather inefficient. Firstly, we previously saw that multiple-parameter optimisations took a longer time than the corresponding single-parameter optimisations. However, on top of that, it is often not even necessary to change all three parameters: the Stejskal–Tanner equation shows that the diffusion attenuation is controlled only by the overall product $\delta^2 G_{\max}^2 \Delta'$. Thus, at least to a first approximation, the individual values of Δ , δ , and G_{\max} do not actually matter. It makes sense to fix two of these and optimise one parameter, only changing the other two if necessary.

It should be mentioned here that obtaining *accurate* values of D requires careful consideration of various factors such as the exact DOSY sequence and gradient shapes used,⁸⁵ peak overlap,^{86–90} and convection.^{91,92} This may influence the type of processing which is chosen for extraction of diffusion coefficients. However, these issues are not our concern here: the role of POISE is only to find a good set of acquisition parameters such that the resulting data are amenable towards further processing.

Optimisation setup

The overall strategy for this ‘optimisation’ can be briefly summarised in a flowchart (fig. 3.30). Note that this is *not* a classic optimisation which falls nicely into the POISE workflow (fig. 3.1):

thus, the entire procedure cannot simply be summarised into one POISE routine. However, it does contain two optimisation subproblems: one to adjust Δ if necessary, and one to find G_{\max} . We can therefore create a wrapper script for the overall procedure (here called `dosy_opt.py`, which invokes POISE to solve the two individual subproblems. This approach is made possible by the command-line interface of POISE, which allows for the creation and execution of optimisation routines. We also exploit the fact that during the optimisation, POISE stores the value of the cost function as the `TI` parameter in TopSpin. This fact is often irrelevant as the value of the cost function is not often of great interest, but proves to be handy in this particular instance as it allows POISE to pass information back to the wrapper script. Previously, we explored situations where the functionality of POISE could be extended from *within*, e.g. by invoking different scripts in its acquisition AU programme. In this case, we are going in the other direction, allowing POISE to be incorporated in other scripts.

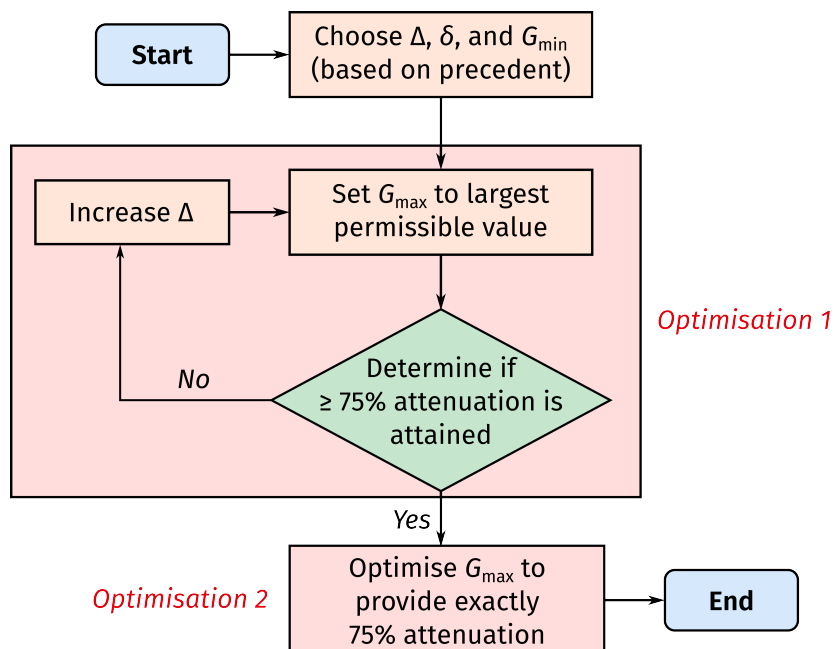


Figure 3.30: Flowchart for DOSY parameter optimisation. In fact, this consists of two subproblems which can be individually solved using POISE.

The DOSY optimisation procedure uses a 1D pulse programme throughout: this should essentially be a single increment of the desired pseudo-2D DOSY experiment.* In this work, I specifically used the Oneshot DOSY sequence⁹³ (fig. 3.28c): this sequence has the advantage of not requiring onerously long phase cycles, which would otherwise lead to very long FEs and optimisations.[†] In principle, though, the optimisation can be run with any DOSY variant, in-

*1D versions of most DOSY variants are available in the TopSpin standard library, but can also be easily obtained by disabling the gradient incrementation in the pseudo-2D versions.

[†]However, using *too few* scans leads to inaccuracies in the diffusion coefficients.⁹³ I chose to use 4 scans per FE here.

cluding convection-compensated sequences:^{94–96} the `dosy_opt.py` wrapper script is sufficiently general that only minor modifications are needed to use it with other DOSY sequences.

The first optimisation subproblem is the determination of an appropriate diffusion delay Δ . Since larger diffusion delays lead to a greater loss of signal through T_1 relaxation, as well as more convection,⁹¹ Δ should always be kept as short as possible.* Since the procedure in fig. 3.30 only *increases* Δ , the wrapper script should always be started with the smallest value of Δ which is sensible for the sample under study: in the present case, I used an initial value of 50 ms. The next step is to determine whether using the full range of available gradient amplitudes leads to sufficient attenuation (which I defined to be 75%), relative to the first increment of the DOSY acquired using G_{\min} . This involves setting G_{\max} to the largest permissible value (for the Oneshot sequence, this is 80%), then calculating a cost function $f_{\text{dosy,aux}}$:

$$f_{\text{dosy,aux}} = \frac{\sum_i S_i(G_{\max})}{\sum_i S_i(G_{\min})} - 0.25. \quad (3.36)$$

In the fraction, the numerator represents an integral over some region of the spectrum acquired with G_{\max} , and the denominator an integral over the same region for the spectrum acquired with G_{\min} . If the attenuation is sufficient, then $f_{\text{dosy,aux}}$ will be negative.

In order for the wrapper script to monitor this cost function, POISE is instructed to perform an ‘optimisation’ with only one function evaluation, using the `--maxfev 1` flag: essentially, POISE is being used here purely as a way of evaluating the cost function $f_{\text{dosy,aux}}$.† At each point, POISE stores the value of $f_{\text{dosy,aux}}$ in the `TI` parameter, and returns control to the wrapper script so that it can check this value. If the value of $f_{\text{dosy,aux}}$ is negative, then the wrapper script moves on to the next subproblem. However, if it is still positive, then there is no way the desired attenuation can be accomplished using the gradient strengths available. Thus, Δ is increased by a fixed amount (50 ms in this case), and the sub-optimisation step repeated.‡

Once a suitable value of Δ has been found, the script then searches for the correct value of G_{\max} in order to provide *exactly* the ‘correct’ amount of attenuation (in this case, 75%). This is done through a conventional POISE optimisation, where the cost function is

$$f_{\text{dosy}} = |f_{\text{dosy,aux}}| = \left| \frac{\sum_i S_i(G_{\max})}{\sum_i S_i(G_{\min})} - 0.25 \right|. \quad (3.37)$$

*In hindsight, it would likely have been better to increase δ instead: this would also have a larger effect as the attenuation depends quadratically on δ but only linearly on Δ . Care must be taken to not increase δ beyond what the spectrometer can tolerate, though.

†Of course, this cost function evaluation could just be done in the wrapper script itself. In fact, it would likely be simpler to do so. But there is one clear benefit of doing it in this rather circuitous route: POISE allows the user to write cost functions in Python 3.

‡If sample spinning is used, then this incrementation of Δ needs to be chosen carefully, as the diffusion delay should always correspond to an integral number of rotations.

The absolute value here makes sure that the optimisation converges to the value of G_{\max} which gives *precisely* a 75% attenuation, no more, and no less. At this point, the wrapper script reports the appropriate values of Δ and G_{\max} , and then exits; these values can then be used to set up a DOSY experiment (e.g. using the dosy TopSpin AU programme).

Before presenting some results, it is worth contemplating what would happen if the sample under study contained a mixture of different molecules, as would often be the case in DOSY experiments. Assuming that the integrals $\sum_i S_i$ are taken over the entire spectrum, the parameters thus obtained would then be ‘compromise values’ which are weighted more heavily towards the major component(s) in the sample. This spectral region being evaluated can be changed by the user (via the dpl command), but that relies on some knowledge of the sample being studied. Thus, in this case, our best hope is really that the ‘compromise’ parameters obtained are *good enough* for all the components in the mixture. Obviously, this can hardly be guaranteed, and the larger the difference in the diffusion coefficients of the constituent species, the more prone this approach is to failing.

In my defence, any script which seeks to optimise DOSY parameters must account for the presence of mixtures in some way, for example by identifying rapidly- and slowly-diffusing species. This is therefore not truly an issue with POISE *per se*, but rather with the approach taken in the wrapper script. (For example, if the wrapper script can identify distinct resonances which have different attenuation profiles, then POISE could be used to perform individual optimisations on each of these.) Nevertheless, for this reason, I refrain from claiming that the work in this section constitutes a *full* method for automating DOSY.* I am more comfortable labelling this as a demonstration of how POISE can be integrated into larger procedures—for example, in the setup of single-component DOSY experiments.

Optimisation results

The DOSY optimisation procedure described above was tested on the sample of andrographolide (in DMSO- d_6). Despite this being a single-component sample, it does at least have some mildly interesting behaviour, in that the exchangeable OH protons have a larger apparent diffusion coefficient compared to the CH protons.

As described previously, the procedure was initialised with a value of $\Delta = 50$ ms. The gradient duration δ , which is not modified, was set to 2 ms. The first sub-optimisation always increased the value of the diffusion delay by one step to 100 ms. Since this part only uses one function evaluation in POISE, the optimisation algorithm used does not actually matter. However, the algorithm does matter for the second sub-optimisation (which searches for the best value of G_{\max}).

*Doing such an endeavour proper justice would anyway have required rather more time in my DPhil and more space in this thesis.

This part was therefore performed five times per algorithm, in line with previous optimisations in this chapter (table 3.26).

Entry	Algorithm	Optimum found (%)	FEs	Time taken (s)
1	NM	75.00	9	195–197
2	MDS	75.00	9	196–197
3	BOBYQA	75.00–75.60	8–9	175–197

Table 3.26: Results of maximum gradient amplitude sub-optimisations for DOSY parameter setup. The POISE routine used here is: `{"name": "dosy", "pars": ["gpz1"], "lb": [20.0], "ub": [80.0], "init": [50.0], "tol": [2.0], "cf": "dosy", "au": "poise_1d"}`. Data code: 6A-200823.

In this case, all three algorithms converged to the same result of 75%. The full Oneshot DOSY spectrum was then acquired using these parameters of $\Delta = 100$ ms, $\delta = 2$ ms, and $G_{\max} = 75\%$ (nominally 49.3 G cm^{-1} on the spectrometer used here, see table 0.1). The overall time taken was approximately five minutes: two for the first sub-optimisation to find Δ , and three for the second sub-optimisation (as shown in table 3.26). This figure may of course be larger depending on the number of scans used (here I used DS=2 and NS=4), as well as the number of iterations required in the first sub-optimisation.

The resulting attenuation profiles for one CH and one OH peak are shown in fig. 3.31. The apparently more slowly-diffusing CH peak is attenuated to 28% of its original intensity, whereas the exchanging OH peak is attenuated to 20%. This is certainly a case where the ‘different components’ have diffusion coefficients which are not too dissimilar, allowing reasonable results to be obtained. Gaussian fitting using the modified Stejskal–Tanner equation

$$I(G) = I(0) \exp \left\{ -(\gamma \delta G)^2 D \left[\Delta - \frac{\delta(5 - 3\alpha^2)}{16} - \frac{\tau(1 - \alpha^2)}{2} \right] \right\} \quad (3.38)$$

yielded *apparent* diffusion coefficients of $(7.411 \pm 0.013) \cdot 10^{-9} \text{ m}^2 \text{ s}^{-1}$ for the CH peak and $(9.369 \pm 0.049) \cdot 10^{-9} \text{ m}^2 \text{ s}^{-1}$ for the OH peak. (The equation was obtained from the pulse programme provided on the University of Manchester website; here τ represents the time between the midpoints of each gradient within the bipolar pulse pair, and α the gradient imbalance factor as shown in fig. 3.28c.)

Simultaneous optimisation of Δ and G_{\max}

It is possible—albeit inefficient, as previously mentioned—to dispense with the wrapper script entirely and perform a single optimisation of both Δ and G_{\max} . In order to ensure that Δ is not

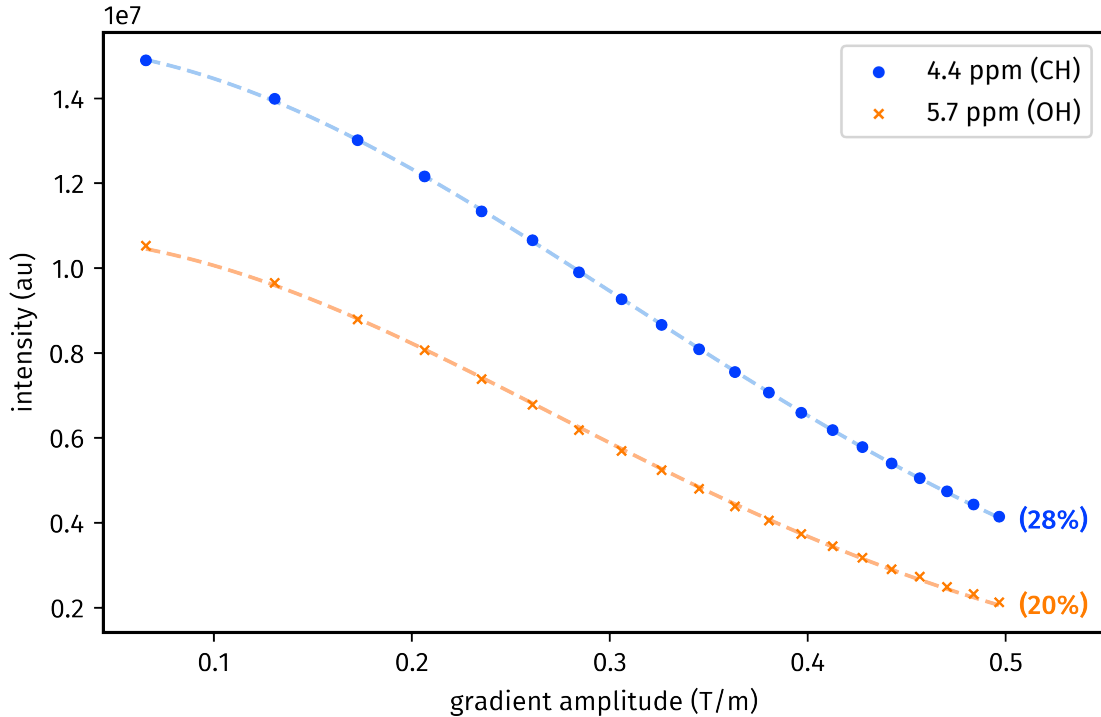


Figure 3.31: Diffusion profiles obtained using a Oneshot experiment for a CH peak (blue, circles) and an OH peak (orange, crosses) in andrographolide. The dashed lines represent the Gaussian curves obtained through non-linear least-squares fitting (`scipy.optimize.curve_fit`). Data code: 6A-200823.

increased too much, we must add a penalty term to the cost function:

$$f_{\text{dosy},2p} = f_{\text{dosy}} + (\Delta/s) = \left| \frac{\sum_i S_i(G_{\max})}{\sum_i S_i(G_{\min})} - 0.25 \right| + (\Delta/s) \quad (3.39)$$

(where ‘2p’ stands for two-parameter). One significant drawback of this is that the reference spectrum $S(G_{\min})$ must be reacquired every time Δ is changed: so, each FE requires twice as long as before. For example, using the NM algorithm, the optimisation converged to $\Delta = 91.6$ ms and $G_{\max} = 79.3\%$. Although this result is quite similar to what we had obtained before, the optimisation took over 26 minutes, over five times longer than previously. I therefore opted not to perform the full 15 optimisations.

Curiously, BOBYQA did not work as well with this cost function: in the two times I tested it, BOBYQA converged to an ‘optimum’ of $\Delta = 225$ ms and $G_{\max} = 50\%$. This did yield the expected 75% attenuation in the spectrum, but Δ is clearly rather longer than we would like it to be. Furthermore, because of the penalty term, the value of the cost function at this point (0.229) was far larger than the corresponding optimum found using the NM algorithm (0.095). It is likely that some of the trust-region optimisation parameters must be tweaked to make this work properly, but I did not spend any further time investigating this.

3.5 POISE for ESR

In the final section of this chapter, I briefly discuss how the concept of on-the-fly optimisation may be extended to pulsed ESR spectroscopy as well. Much of the underlying code from NMR-POISE was recycled for this, including the implementation of the three optimisation algorithms (NM, MDS, and BOBYQA). I added only one small difference in ESR-POISE, namely, a way to control the initial size of the simplex (for NM and MDS) or the trust region radius (for BOBYQA): in NMR-POISE, this is fixed at ten times the desired optimisation tolerance. The overall optimisation framework (fig. 3.1), as well as the concept of the optimisation routine (§ 3.2.1), are retained: this preserves the generality of POISE, which is probably its greatest strength.

Naturally, TopSpin-specific sections of the code had to be rewritten to instead be compatible with Bruker's Xepr software. This in fact proved to be easier than anticipated: most of the TopSpin code could be completely deleted, because Xepr provides an API which can be accessed from external programmes such as Python 3. Thus, ESR-POISE is in fact completely written in Python 3, and there is no need for an artificial separation between ‘frontend’ and ‘backend’ (which also neatly circumvents most of the issues discussed in § 3.2.5). The Xepr-facing code was written by Jean-Baptiste Verstraete (University of Oxford) and David Goodwin (University of Southampton, formerly Oxford).

A number of applications in ESR-POISE were explored. All ESR experimental work was done by Jean-Baptiste Verstraete and William Myers (Oxford); thus, I only provide very brief descriptions here. The examples include:

- optimisation of the signal phase in a simple spin echo experiment, by maximising the intensity of the detected echo;
- similarly, calibration of 90° and 180° pulse widths and powers in a spin echo;
- optimisation of an inversion pulse used on the pump channel in a DEER experiment,⁹⁷ in order to increase the modulation depth of the resulting DEER traces (which reveal dipolar couplings between electrons);
- calibration of shaped pulse amplitudes for CHORUS broadband excitation,^{98,99}
- compensation for resonator distortions when applying shaped pulses, again demonstrated using the CHORUS sequence.

An example of the results obtained via the latter two optimisations is shown in fig. 3.32. In fig. 3.32a, the CHORUS broadband excitation experiment was run without any optimisations: the resulting spectrum (blue, solid) is compared against that obtained via a standard field sweep

(grey, dashed). Clearly, there are substantial mismatches. Figure 3.32b shows the improved performance of CHORUS after a simple optimisation of the shaped pulse amplitudes, using the spectral intensity (or the negative thereof, since we seek to maximise it) as the cost function: the POISE optimisation therefore allows for a much more accurate spectrum to be obtained.

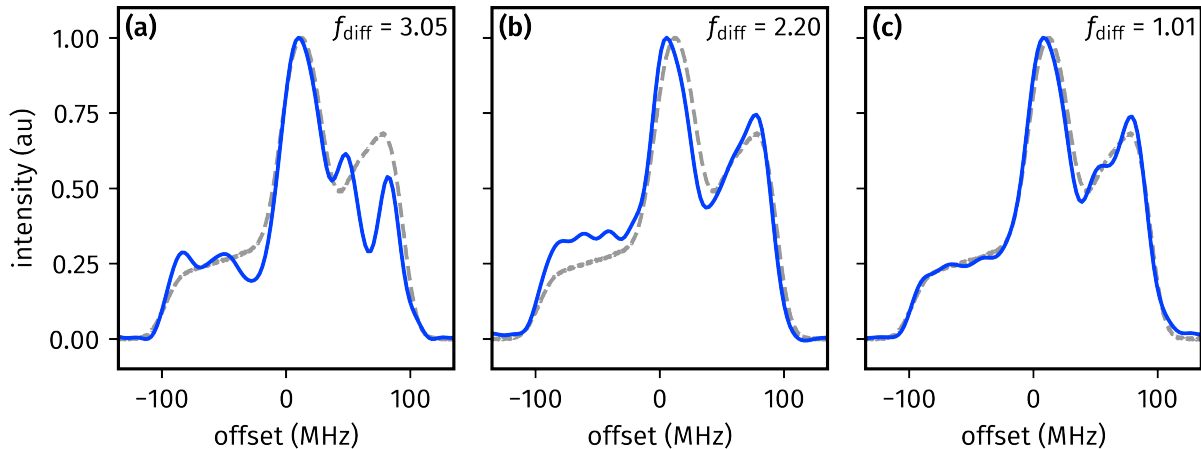


Figure 3.32: Comparison between CHORUS spectrum (blue, solid line) and field sweep profile (grey, dashed line): (a) without any optimisation of CHORUS, (b) after optimisation of the CHORUS amplitudes, and (c) using a compensated CHORUS pulse obtained through determination of the resonator transfer function. The value of f_{diff} is indicated on each plot. The sample used was a bisnitroxide; further experimental details are provided in the paper.² The data used for this figure were acquired by Jean-Baptiste Verstraete.

This can be further improved, however, by compensating for distortions in the pulse shape caused by the resonator. Typically, this is a time-consuming process, requiring the determination of a transfer function: in this case, however, it can be performed in a fully automated fashion using POISE. The coefficients in the transfer function are used as optimisation parameters, and are used within each function evaluation to back-calculate a compensated pulse shape: the cost function used is the difference between the compensated CHORUS spectrum and the field sweep (f_{diff} , as defined in eq. (2.33)). At the point where the cost function is minimised, the transfer function parameters will have been accurately determined. The final compensated CHORUS spectrum is shown in fig. 3.32c.

3.6 Conclusion

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Chapter 4

NOAH

The power of doing any thing with quickness is always much prized by the possessor, and often without any attention to the imperfection of the performance.

— JANE AUSTEN, *Pride and Prejudice*

This final chapter describes my work on NOAH (NMR by Ordered Acquisition using ^1H detection) *supersequences*, pulse sequences which record multiple 2D datasets ('*modules*') in the time required for one. This is an attractive NMR technique for several reasons: the time savings are clearly a key factor, but the flexibility of being able to combine almost any set of modules also makes NOAH supersequences applicable to a variety of contexts.

I begin by introducing the concepts underlying NOAH supersequences, as well as a general discussion of the time savings (and sensitivity per unit time benefits) thus realised. I then describe the GENESIS (GENERation of Supersequences In Silico) website, written by me, which allows users to generate Bruker pulse programmes for almost every imaginable NOAH supersequence. After this, my work on various aspects of the actual sequences themselves is described, with a special focus on newly developed and/or improved modules. Finally, the design of 'parallel' supersequences, which comprise multiple interleaved supersequences, is discussed.

This work was done in close collaboration with Ēriks Kupče (Bruker UK). However, all results and analysis shown in this chapter are mine. The work in this chapter forms the subject of several publications:

- Yong, J. R. J.; Hansen, A. L.; Kupče, Ē.; Claridge, T. D. W. Increasing sensitivity and versatility in NMR supersequences with new HSQC-based modules. *J. Magn. Reson.* 2021, 329, 107027, DOI: [10.1016/j.jmr.2021.107027](https://doi.org/10.1016/j.jmr.2021.107027)
- Kupče, Ē.; Yong, J. R. J.; Widmalm, G.; Claridge, T. D. W. Parallel NMR Supersequences:

Ten Spectra in a Single Measurement. *JACS Au* **2021**, *1*, 1892–1897, DOI: [10.1021/jacsa.1c00423](https://doi.org/10.1021/jacsa.1c00423)

- Yong, J. R. J.; Kupče, Ě.; Claridge, T. D. W. Modular Pulse Program Generation for NMR Supersequences. *Anal. Chem.* **2022**, *94*, 2271–2278, DOI: [10.1021/acs.analchem.1c04964](https://doi.org/10.1021/acs.analchem.1c04964)
- Yong, J. R. J.; Kupče, Ě.; Claridge, T. D. W. Uniting Low- and High-Sensitivity Experiments through Generalised NMR Supersequences. **2022**, manuscript in preparation

The material in the introductory sections also closely follow two reviews which I have contributed to:

- Kupče, Ě.; Frydman, L.; Webb, A. G.; Yong, J. R. J.; Claridge, T. D. W. Parallel nuclear magnetic resonance spectroscopy. *Nat. Rev. Methods Primers* **2021**, *1*, No. 27, DOI: [10.1038/s43586-021-00024-3](https://doi.org/10.1038/s43586-021-00024-3)
- Yong, J. R. J.; Kupče, Ě.; Claridge, T. D. W. In *Fast 2D solution-state NMR: concepts and applications*, Giraudeau, P., Dumez, J.-N., Eds., forthcoming, 2022

4.1 Introduction

The characterisation of small molecules and biomolecules by NMR spectroscopy relies on a suite of standard 2D NMR experiments, which seek to detect heteronuclear scalar couplings (e.g. HSQC and HMBC), homonuclear scalar couplings (e.g. COSY and TOCSY), or through-space interactions (e.g. NOESY and ROESY). Although such 2D experiments provide far superior resolution and information content compared to 1D spectra, they also require substantially longer experimental durations, as the indirect dimension must be constructed through the acquisition of many t_1 increments. This problem is further exacerbated by the fact that structural elucidation or verification often requires combined information from multiple different 2D experiments.

The acceleration of 2D NMR has thus proven to be a popular area of research. We may broadly categorise these existing techniques into two classes.* First come the methods which seek to directly speed up the acquisition of *individual* 2D spectra: these include non-uniform sampling (NUS),^{7–10} fast pulsing (i.e. shortening of recovery delays),^{11–14} ultrafast NMR,^{5,15–19} Hadamard encoding,^{20,21} and spectral aliasing.^{22–25}

On the other hand, *multiple-FID experiments* aim to collect two or more 2D spectra in the time required for one. The corresponding FIDs may either be detected at the same time (as in time-shared NMR^{26–28} and multiple-receiver NMR^{29–31}), or *sequentially*: this is the category which NOAH supersequences^{5,32,33} fall under. NOAH supersequences are furthermore set apart from other sequential-acquisition experiments^{34–39} by their ‘modular’ nature: they can be constructed from building blocks, or ‘modules’, which (usually) contain one FID each. Typically, each of these building blocks corresponds to a specific 2D experiment, as will be shown below. It should be noted that there are several other multiple-FID experiments which, while not explicitly advertised as ‘NOAH experiments’, are built from ‘modules’ and are conceptually identical;^{40–43} I do not discuss these here.

The time savings provided by NOAH experiments stems from the elision of *recovery delays*—the time required for spins to relax to their equilibrium state (ρ_0 in eq. (1.39)), such that the next transient or t_1 increment can be recorded. These recovery delays are responsible for most of the experimental duration of a typical 2D NMR experiment. In a NOAH supersequence, this is accomplished by designing each module to draw on a different *magnetisation pool*: this refers to a source of magnetisation belonging to a specific isotopologue. For example, ^{13}C -bound protons and ^{12}C -bound protons are two different magnetisation pools. This allows modules to be directly concatenated, without the addition of extra recovery delays between them; only one overall recovery delay is required for the entire supersequence, where *all* magnetisation pools

*These are by no means mutually exclusive: many of the techniques here can be combined to provide even greater efficiency.

simultaneously recover.

In this section, I first provide a formal overview of how the time savings for NOAH experiments may be quantified and analysed. This includes an analysis of the conditions under which multiple-FID techniques such as NOAH are most valuable. Following this, I turn to the design of NOAH pulse sequences, using specific case studies to illustrate the general principles underlying these experiments.

4.1.1 Time savings and sensitivity analyses

Time savings in the sampling-limited regime

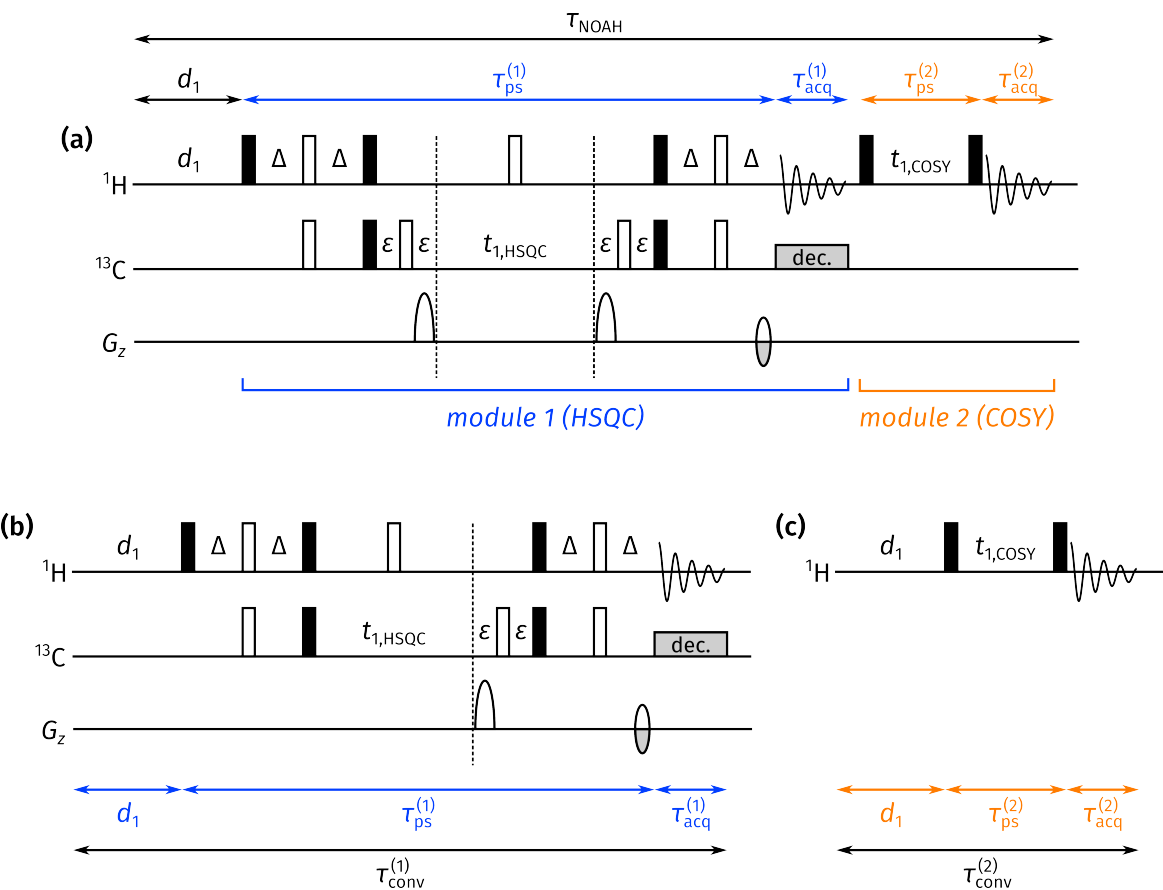


Figure 4.1: (a) NOAH-2 SC supersequence, comprising HSQC and COSY modules (see table 4.1 for an explanation of the single-letter module codes used). (b) ‘Conventional’ echo-antiecho HSQC (the same as in fig. 1.7). (c) ‘Conventional’ COSY. The timings referred to in the text are highlighted for all three experiments; d_1 for each experiment is assumed to be the same. Note that the lengths are not to scale: d_1 is typically far longer than τ_{ps} and τ_{acq} .

The duration of an NMR experiment, τ_{exp} , can be expressed as a sum of its parts:

$$\tau_{\text{exp}} = \tau_{\text{ps}} + \tau_{\text{acq}} + d_1, \quad (4.1)$$

where τ_{ps} is the time required for the pulse sequence itself (typically several milliseconds), τ_{acq} is the acquisition time (several hundred milliseconds), and d_1 is the recovery delay (one or more seconds). These timings are illustrated in fig. 4.1 for a NOAH supersequence formed from two modules (HSQC and COSY). The *time-saving factor* ρ_t for a NOAH supersequence, as compared to a series of conventional standalone experiments, may be expressed as:

$$\rho_t = \frac{\sum_i \tau_{\text{conv}}^{(i)}}{\tau_{\text{NOAH}}} = \frac{\sum_i (\tau_{\text{ps}}^{(i)} + \tau_{\text{acq}}^{(i)} + d_1^{(i)})}{d_1 + \sum_i (\tau_{\text{ps}}^{(i)} + \tau_{\text{acq}}^{(i)})}, \quad (4.2)$$

where τ_{NOAH} is the duration of the NOAH experiment, τ_{conv} is the duration of a conventional experiment, and the superscript (i) represents the i -th module or conventional experiment being acquired. The sum runs from $i = 1$ to N , where N is the number of modules. If we assume that $d_1^{(i)} = d_1$ is the same for all N conventional experiments and the supersequence, then in the limit where

$$d_1 \gg \sum_i \tau_{\text{ps}}^{(i)} + \tau_{\text{acq}}^{(i)}, \quad (4.3)$$

we have that $\rho_t \rightarrow Nd_1/d_1 = N$, i.e. an N -fold time saving.

This analysis makes plenty of assumptions, and is not entirely valid in practice. For example, *each* τ_{acq} is often around 5–10% of d_1 , so is not entirely negligible, especially when N is larger. Furthermore, some modules require longer τ_{ps} : most notable is the NOESY module, which contains a mixing time of several hundred milliseconds. (HMBC, TOCSY, and ROESY modules are also lesser offenders.)

These factors serve to reduce ρ_t from its idealised value of N ; generally, this deviation is larger as N increases, because eq. (4.3) becomes less and less valid. Despite this, it remains true that the time savings are approximately proportional to N , as can be appreciated from table 4.2.

For relatively concentrated samples, where sensitivity is not an issue, these time savings are the sole relevant point. In this *sampling-limited regime*, the minimum 2D experiment duration is dictated purely by the number of t_1 increments needed to obtain sufficient resolution in the indirect dimension, as well as the minimum phase cycle required for artefact suppression.* NOAH supersequences are identical to conventional experiments in both these aspects, but come with the added benefit of time savings. The development of modern NMR instrumentation (such as high-field magnets and cryogenic probes) plays an important role in extending the sampling-limited regime to ever lower concentrations.

*With modern gradient-enhanced experiments, the minimum phase cycle may not even be a ‘cycle’, as one scan per increment may suffice; see also fig. 1.8.

Sensitivity comparisons

However, the opposite *sensitivity-limited regime* is still very commonly encountered. This can happen, for example, with naturally insensitive experiments (e.g. ADEQUATE), low-field benchtop NMR, or most simply, dilute samples.* In such cases, the benefits of NOAH supersequences must be much more carefully considered: in particular, it becomes mandatory to compare the SNRs of the NOAH modules and conventional experiments. To do so, we define for each module an *SNR factor A*, which is the SNR of the NOAH module divided by the SNR of a conventional experiment, acquired with the same parameters.[†] The SNR factor of the i -th module in a supersequence may be denoted as $A^{(i)}$.

In general, we have that $A \leq 1$, because (as will be explained in § 4.1.2) NOAH modules frequently contain small modifications with respect to conventional experiments. For sensitivity-limited samples, we must balance these losses against the time savings obtained, by defining a *gain in sensitivity per unit time*, ε as

$$\varepsilon = A \sqrt{\rho_t}. \quad (4.4)$$

Here, the square root accounts for the fact that SNR scales only as the square root of the number of scans, or equivalently, the number of times the experiment can be repeated in a given time. If $\varepsilon^{(i)} > 1$, as is frequently the case, this means that the NOAH supersequence provides greater sensitivity per unit time in the i -th module compared to a standalone experiment. Equivalently, performing a NOAH experiment allows data of sufficient sensitivity to be obtained in less time.

Time savings in the sensitivity-limited regime

An important—yet often ignored—caveat with sensitivity-limited samples is that the time savings obtained are not *truly* on the order of N as advertised above. This is because each module in a NOAH supersequence is run with the same number of scans. In contrast, with conventional experiments, it is common to run more insensitive experiments with a larger number of scans but to use fewer for sensitive experiments. In this case, the *effective* time savings provided by

*If the SNR factor $A^{(i)}$ as discussed below is *very small*, then it is possible that even concentrated samples may be shifted into the sensitivity-limited regime. This is never really the case in practice, though, as will be shown in § 4.1.3.

[†]The relative SNR will likely vary from peak to peak in the spectrum, and A should in theory be quoted either as an average over all peaks, or as a range. This is what I have done in this thesis. Furthermore, the exact values calculated for A will depend on the sample used for the comparison. Any values given here should therefore be assumed to be valid only for similar samples. Of course, my hope is that the samples chosen are reasonably representative of ‘typical’ organic molecules.

NOAH experiments are smaller:

$$\rho_{t,\text{eff}} = \frac{\sum_i \tau_{\text{conv}}^{(i)}}{\tau_{\text{NOAH}}} = \frac{\sum_i S^{(i)}(\tau_{\text{ps}}^{(i)} + \tau_{\text{acq}}^{(i)} + d_1^{(i)})}{Sd_1 + S \sum_i (\tau_{\text{ps}}^{(i)} + \tau_{\text{acq}}^{(i)})}, \quad (4.5)$$

where each standalone experiment is acquired with $S^{(i)}$ scans, and the NOAH experiment with S scans. We could also analogously define $\varepsilon_{\text{eff}} = A \sqrt{\rho_{t,\text{eff}}}$.

Typically, S is simply the largest of the $S^{(i)}$, as this ensures sufficient data quality for the most insensitive module in the supersequence. This means that generally, $\rho_{t,\text{eff}} < \rho_t$. In such a situation, it is probably more appropriate to describe a NOAH supersequence as ‘measuring the most insensitive module and getting the others for free’. Indeed, in the extreme case where $S = S^{(i)} \gg S^{(j \neq i)}$, then ‘the other’ modules require almost no time to measure (relative to the least sensitive module), and $\rho_{t,\text{eff}}$ tends towards 1, meaning that even the time-saving utility of NOAH vanishes. A corollary of this is that NOAH supersequences are generally best constructed from modules which have similar intrinsic sensitivities and hence similar $S^{(i)}$ ’s.

Returning to the sampling-limited regime for a short while, we generally have that $S^{(i)} = S$ for all i (i.e. all conventional experiments are acquired with the same number of scans). In this case, eq. (4.5) simply reduces to eq. (4.2).

As the reader will no doubt appreciate by now, the comparison of NOAH and conventional spectra is fraught with subtleties. In fact, it is very much possible to construct yet more edge cases which challenge the framework used in this analysis. For example, one may not want to acquire all the individual spectra ‘conventionally’: for example, NUS may be used for an HSQC experiment but not for others; or d_1 may be varied for different experiments. These will have an impact on both the durations of the experiments, as well as their sensitivities. To make any meaningful comparisons or generalisations, it is therefore necessary to restrict the discussion to values of ρ_t , A , and ε , which can be objectively calculated. These should, however, be read with the understanding that other factors may, depending on the context, lead to *some*—but never a *complete*—decrease in the utility of NOAH experiments.

4.1.2 Magnetisation pools

Having dealt with this relatively dry material, I now turn to exactly how NOAH supersequences are constructed. Ordinarily, if the recovery delay is removed from an NMR experiment, its sensitivity will be greatly reduced because insufficient magnetisation will have recovered between repetitions; or in other words, $A^{(i)}$ will be very small. Such experiments would only really be useful if sensitivity was greatly abundant.

As briefly mentioned earlier, the key to avoiding this in NOAH supersequences is to make sure

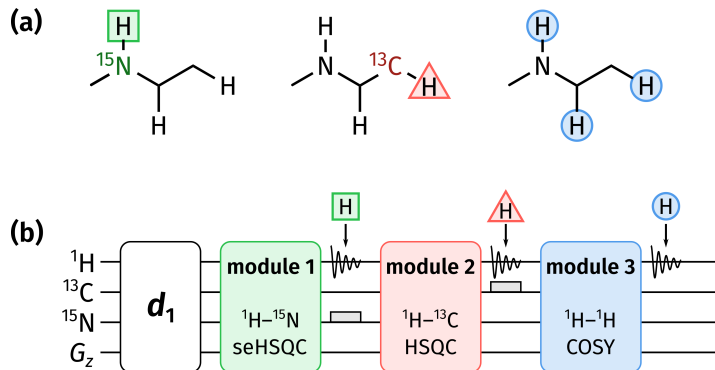


Figure 4.2: (a) An illustration of the different magnetisation pools used in a typical NOAH supersequence: the three pools here (¹⁵N-bound protons, ¹³C-bound protons, and all others) would be referred to as ¹H^N, ¹H^C, and ¹H^{IX} respectively. (b) A NOAH supersequence consisting of three modules, which each consume one of the above magnetisation pools.

that *each module samples a different source of magnetisation*. These are referred to as *magnetisation pools*, and arise because of the different isotopologues present in natural-abundance samples (fig. 4.2a). For example, an HSQC module can be designed to only sample magnetisation of protons directly bonded to the 1.1%-natural abundance ¹³C, and leave all other proton magnetisation untouched. Immediately following this, the remainder of the proton magnetisation can then be used to record (say) a COSY module, without needing a separate recovery delay. Figure 4.2b shows a more elaborate example, which includes both ¹⁵N and ¹³C HSQC experiments.

Using the notation of Orts and Gossert,⁴⁴ the magnetisation of ¹³C-bound protons is denoted as ¹H^C, and that of ¹⁵N-bound protons is ¹H^N. The magnetisation of protons *not* bonded to ¹³C is denoted as ¹H^{!C} (and likewise ¹H^{!N}). Protons not directly bonded to *any* NMR-active heteronucleus are labelled ¹H^{IX}, and are often referred to as ‘bulk’ magnetisation, since the majority of protons in natural-abundance samples fall into this category.

Since standard 2D experiments typically seek to *destroy*—instead of *preserve*—unused magnetisation, NOAH modules often require some modifications compared to standard experiments. For example, compared to the echo-antiecho HSQC (discussed in § 1.4.5), the NOAH HSQC module³² adds an extra CTP gradient. This ensures that the bulk magnetisation is refocused after t_1 , and ultimately returned to the $+z$ equilibrium state (fig. 4.3a). (This is largely identical to the ‘symmetrised’ ASAP-HSQC experiment.⁴⁵) Sometimes, the modifications required are more extensive, such as in the HMBC module. In order for this module to preserve ¹H^C magnetisation (e.g. for a later HSQC module), the initial 90° excitation pulse must be replaced with a *zz*-filter (fig. 4.3b). This performs an *isotope-selective rotation* in that ¹H^C magnetisation is stored along the z -axis, but ¹H^{!C} magnetisation is excited and subsequently detected.

In general, sequences which are thus modified have lower sensitivities (i.e. $A < 1$) than the ‘original’ sequences from which they were derived. This is partly because of imperfect manipu-

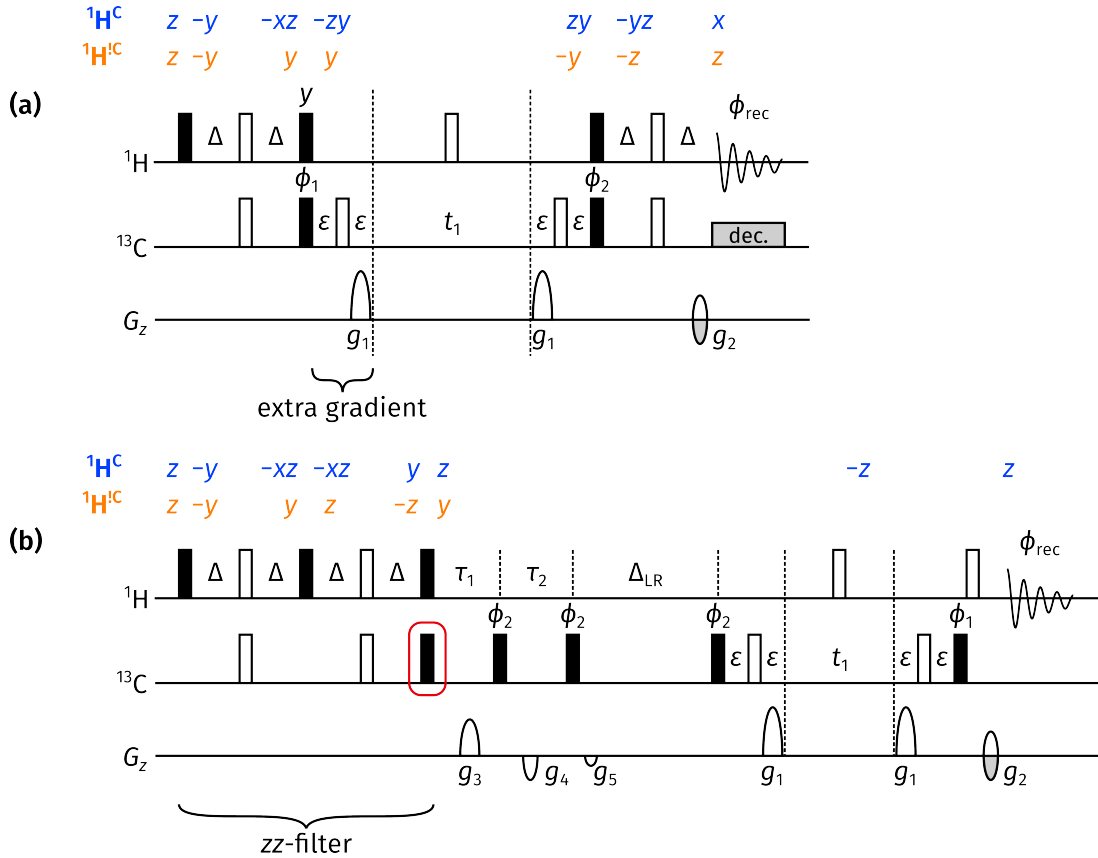


Figure 4.3: (a) NOAH HSQC module. (b) NOAH HMBC module. The 90° pulse highlighted in red is described in § 4.3.7. Delays are set as: $\Delta = 1/(4 \cdot {}^1\text{J}_{\text{CH}})$; $\Delta_{\text{LR}} = 1/(2 \cdot {}^3\text{J}_{\text{CH}})$; $\tau_1 = 1/(2 \cdot {}^1\text{J}_{\text{CH,max}})$; $\tau_2 = 1/(2 \cdot {}^1\text{J}_{\text{CH,min}})$ (see also § 3.4.7 for the LPJF). Phase cycling is performed with $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x)$, and $\phi_{\text{rec}} = (x, -x, -x, x)$. Gradient amplitudes are $(g_1, g_2, g_3, g_4, g_5) = (80\%, \pm 40.2\%, 15\%, -10\%, -5\%)$. Product operator analysis is provided above both modules for both the ${}^1\text{H}^{\text{C}}$ and ${}^1\text{H}^{\text{IC}}$ magnetisation pools; the notation for this is explained in the *Preface*.

lation of magnetisation by the extra pulse sequence elements, and also increased losses due to relaxation during these extended sequences. In contrast, modules placed towards the *end* of a supersequence do not need to be modified, as they do not need to preserve any magnetisation. This includes virtually all homonuclear modules, which are allowed to simply consume any remaining magnetisation. Although this makes their implementation very straightforward, in general these modules will *also* suffer some losses in sensitivity, because the preceding modules do not perfectly retain all magnetisation.

It is therefore fairly rare for a NOAH module to have $A = 1$: for this to happen, the module must be placed first in the supersequence *and* not have undergone any modifications relative to

the standalone experiment.* Such cases are very rare, and it is thus necessary to accept some decreases in A , which are often fairly small (on the order of 10–20%). For sensitive (typically homonuclear) modules, such sensitivity losses are often perfectly tolerable, as even with this sensitivity penalty they are still more intense than the other (heteronuclear) modules in the supersequence. However, for insensitive modules, it is imperative to make sure that A is not decreased by too much.

4.1.3 Case studies

Using all of the concepts described in the previous sections, we now look at a few ‘typical’ supersequences to understand their construction. Before beginning, a quick note about the nomenclature of NOAH supersequences is warranted. Supersequences are labelled by the number of modules N , plus a series of single-letter codes corresponding to the identity and ordering of the modules involved (table 4.1). Occasionally, superscripts or subscripts are used to qualify the modules involved.[†] Thus, a NOAH supersequence containing three modules—say ^{15}N HMQC, ^{13}C HSQC, and CLIP-COSY—would be referred to as a NOAH-3 M_NSC^c. Table 4.2 provides values of ρ_t and A for the example supersequences used in this section; these values which will be rationalised in the text which follows.

$^1\text{H}-^{15}\text{N}$ modules		$^1\text{H}-^{13}\text{C}$ modules		$^1\text{H}-^1\text{H}$ modules	
Module	Code	Module	Code	Module	Code
HMQC	M _N	HSQC	S	COSY	C
HSQC	S _N	seHSQC	S ⁺	CLIP-COSY	C ^c
seHSQC	S _N ⁺	HSQC-TOCSY	S ^T	DQF-COSY	C
HMBC	B _N	HSQC-COSY	S ^C	TOCSY	T
		2BOB	O	NOESY	N
		HMBC	B	ROESY	R
		ADEQUATE	A	PSYCHE	P
				TSE-PSYCHE	P ^T
				PSYCHE 2DJ	J

Table 4.1: A (non-exhaustive) list of single-letter module codes for NOAH modules. In the literature, the ^{15}N HMQC module has been referred to simply by ‘M’, since the HSQC module is preferred for $^1\text{H}-^{13}\text{C}$ correlations. In this thesis, I include the subscript N throughout to avoid any ambiguity.

*Of course, this also depends on exactly *what* standalone experiment the NOAH supersequence is being compared against. Sometimes, in the literature, the NOAH experiment has been compared against its constituent modules acquired in a standalone fashion; in this case, the first module will always have $A = 1$, since by definition it is not modified. This tells us how much we gain through the act of concatenating modules, but is less meaningful in the ‘real world’ where one is interested in how useful NOAH is relative to ‘typical’ optimised 2D experiments.

[†]With the increasing number of modules, and the variety of modern NMR experiments which could be incorporated into NOAH supersequences, keeping these abbreviations short yet meaningful has been a challenge.

Entry	Sequence	τ_{NOAH}	ρ_t	A				
				HMBC	seHSQC	HSQC	COSY	TOCSY
1	SC	15 min 0 s	1.87			0.97	0.90	
2	SCT	16 min 25 s	2.60			1.01	0.99	0.79
3	BS	15 min 40 s	1.82	0.93		0.87		
4	SB	15 min 35 s	1.83	0.99		0.96		
5	BSCT	17 min 48 s	3.22	0.95		0.90	0.36	0.28
6	BS_{N}^+ SCT	18 min 57 s	3.74	0.95	0.71	0.66	0.38	0.30
7	S_{N}^+ BSCT	18 min 56 s	3.75	0.76	0.79	0.74	0.33	0.26

Table 4.2: Sensitivity and time-saving analyses of several typical NOAH supersequences. All experiments were acquired with 2 scans per increment, 256 t_1 increments, an acquisition time of 67 ms, and a recovery delay of 1.5 s. The HMBC module used here includes the extra ^{13}C 90° pulse described later in § 4.3.7: this has no significant impact on the SNR, and is only mentioned as a technicality. The ^{15}N seHSQC module is that described in § 4.3.3. The CT module here was run with States indirect-dimension quadrature detection, and the individual C module (in entry 1) with echo-antiecho. The following Bruker library sequences were used as the ‘conventional’ experiments: `hmbcetgpl2nd`, `hsqcetf3gpsi2`, `hsqcetgpss.2`, `cosygpqf`, and `dipsi2gpphzs`. *Data code:* 7Z-220224.

NOAH-2 SC: HSQC + COSY

We begin with perhaps the simplest example of a NOAH supersequence, one containing the HSQC and COSY modules: this is labelled as a NOAH-2 SC experiment (entry 1, table 4.2). Using the implementation shown in fig. 4.3a, the HSQC module only samples $^1\text{H}^{\text{C}}$ magnetisation, and leaves $^1\text{H}^{\text{!C}}$ magnetisation along the $+z$ -axis. This modification to the HSQC module is very small, so its sensitivity is practically unaffected as compared to a ‘standard’ HSQC ($A = 0.97$). Furthermore, the COSY module retains *most* of its sensitivity ($A = 0.90$). The small loss here is because the HSQC module does not *perfectly* preserve the $^1\text{H}^{\text{!C}}$ magnetisation: for example, evolution of homonuclear couplings as well as relaxation occur during the HSQC pulse sequence, both of which were ignored in the product operator analysis of fig. 4.3a.

Furthermore, the value of the time-saving factor, $\rho_t = 1.87$, is very close to the theoretical limit of $N = 2$. This reflects the fact that the pulse sequence itself, τ_{ps} , is fairly short for both the HSQC and COSY modules; the deviation therefore chiefly arises from the acquisition time, τ_{acq} . In all respects, this is therefore an example of an ‘ideal’ NOAH supersequence, where the combination of two modules provides almost 2× time savings without compromising on sensitivity.

It is worth pointing out that the order of the modules cannot be reversed: the COSY module cannot be (easily) modified to preserve $^1\text{H}^{\text{C}}$ magnetisation. In a hypothetical NOAH-2 CS supersequence, the later HSQC module would only be able to use magnetisation recovered during the COSY FID, leading to a substantial decrease in sensitivity.

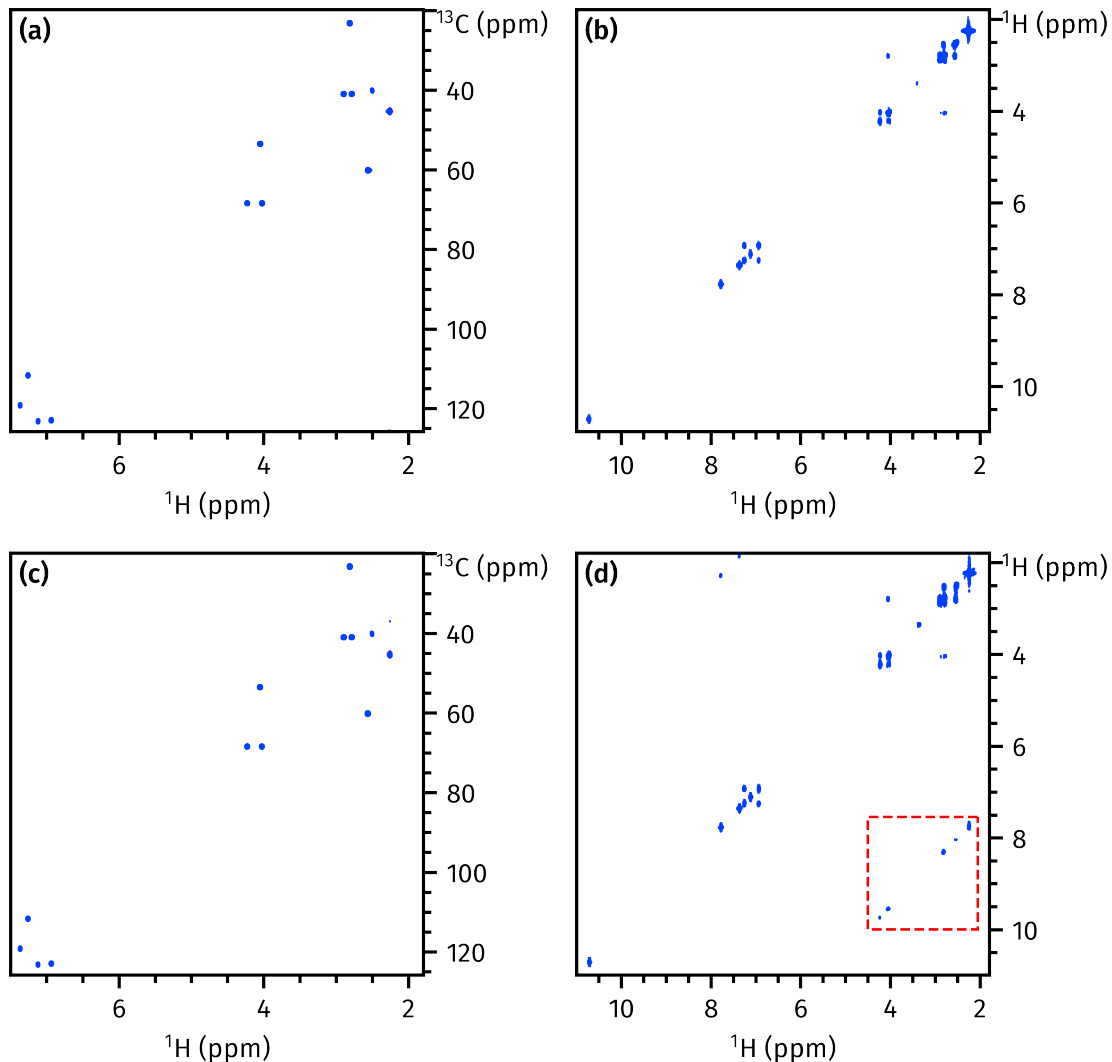


Figure 4.4: (a) HSQC from NOAH-2 SC supersequence. (b) COSY from NOAH-2 SC supersequence. (c) Standalone HSQC. (d) Standalone COSY; off-diagonal artefacts are highlighted in the red box. *Data code:* 7Z-220224.

A final point to consider would be whether the NOAH data has comparable spectral quality in terms of (for example) artefacts. In this case, the answer is yes: the NOAH HSQC spectrum is virtually identical to the standalone (figs. 4.4a and 4.4c; both spectra have low-level artefacts of different kinds, which do not seriously impede the interpretation and are not shown). On the other hand, the NOAH COSY spectrum seems to actually *improve* on the standalone COSY, in that it better suppresses off-diagonal artefacts (figs. 4.4b and 4.4d). These artefacts likely arise in the standalone COSY because of accidental refocusing of magnetisation which has not completely relaxed between successive t_1 increments.⁴⁶ In contrast, the NOAH COSY module has an extra set of HSQC gradients between every repetition of the COSY, so accidental refocusing is less likely. (Similar artefacts have previously been noted in the DQF-COSY experiment,^{47,48} and have also shown to be attenuated in the corresponding NOAH module.⁴⁹) That said, such

improvements are not always guaranteed. It is possible for specific artefacts to arise exclusively in NOAH experiments; some of these will be discussed in the following sections.

NOAH-3 SCT: HSQC + COSY + TOCSY

Evidently, the fact that the HSQC preserves almost all $^1\text{H}^{1\text{C}}$ magnetisation means that *any* homonuclear module can be placed after it. This means that supersequences such as SC^c, SC, and ST can be constructed in a very similar way. It would be very useful if we could go beyond this and place *multiple* homonuclear modules in a single supersequence.

However, since every homonuclear module tends to consume all remaining bulk magnetisation, creating combinations of homonuclear modules which do not ‘compete’ for this bulk magnetisation is a difficult task. The only notable exceptions to this are ‘COSY+X’ combinations, where X can be NOESY, ROESY, or TOCSY: instead of concatenating the COSY and X modules, the COSY pulse sequence can instead be nested *within* the X module, as was first demonstrated with X = NOESY.^{34,35}

Here, we use the COSY/TOCSY combination³⁷ as an example (fig. 4.5). The $90^\circ-t_1-90^\circ$ COSY pulse sequence leads to a mixture of both transverse and longitudinal magnetisation which is frequency-labelled in t_1 . The transverse component is sampled during the COSY acquisition period, whereas the longitudinal component (usually discarded in a COSY experiment) is subjected to isotropic mixing and subsequently sampled as part of the TOCSY. Unlike the HSQC and COSY in the previous example, the COSY and TOCSY in fig. 4.5c cannot be clearly separated into different sections of the pulse sequence: they in fact share a common t_1 period. Nevertheless, in the context of NOAH, it is common to still refer to these as separate modules.

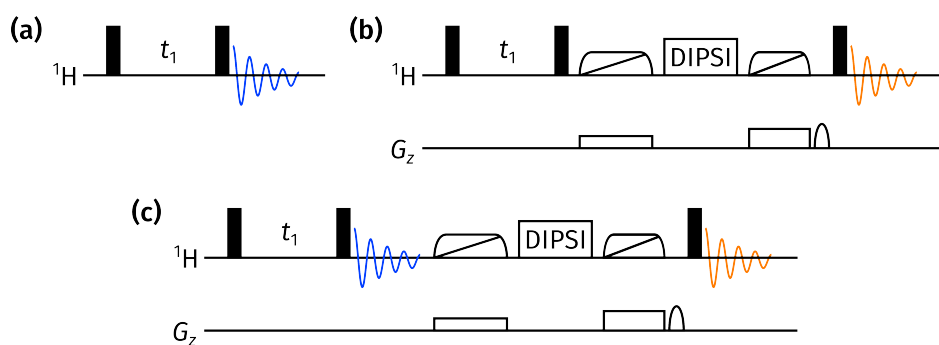


Figure 4.5: (a) COSY module. (b) TOCSY module; zero-quantum suppression is employed before and after the isotropic mixing period. (c) Combined COSY/TOCSY module, where the COSY FID is acquired immediately before the TOCSY mixing.

As shown in entry 2 of table 4.2, this nesting of the COSY module does not materially affect the TOCSY sensitivity. A small loss of approximately 20% is observed, which is partly due to the imperfect magnetisation preservation by the HSQC, and perhaps also due to relaxation during

the COSY acquisition period. As for the time-saving factor, a slightly larger deviation ($\rho_t = 2.60$) is observed from the ideal value of 3. This reflects the addition of an extra acquisition period and the TOCSY mixing period, which make the assumption in eq. (4.3) less valid.

NOAH-2 BS: HMBC + HSQC

As mentioned previously, the HMBC module shown in fig. 4.3b is designed to retain $^1\text{H}^{\text{C}}$ magnetisation through the addition of the zz -filter. This can be used in a subsequent HSQC module in a NOAH-2 BS supersequence. Entry 3 of table 4.2 shows that the addition of the zz -filter to the HMBC causes a relatively small 7% decrease in sensitivity; on the other hand, the HSQC loses 13% of its sensitivity because of incomplete magnetisation preservation. These sensitivity losses, however, are reasonably acceptable considering the time savings provided by this supersequence ($\rho_t = 1.82$).

Generally, it has been recommended that less sensitive modules are placed earlier in the supersequence so that they can access a larger proportion of the equilibrium magnetisation. The NOAH-2 BS does not, however, actually obey this principle. Since the HMBC is the less sensitive of the two modules, this rule of thumb suggests that the BS supersequence should be better than the alternative SB supersequence. In fact, the opposite is true, as entry 4 of table 4.2 shows. This can be understood as follows: the HSQC module experiences a boost in sensitivity because it is placed first in the supersequence, and no longer needs to rely on the $^1\text{H}^{\text{C}}$ magnetisation preserved by the HMBC; and the HMBC benefits because the zz -filter modification is no longer needed.* This may just be the exception that proves the rule, but in my view, this provides some justification for considering the ordering of modules on a case-by-case basis.

NOAH-4 BSCT: HMBC + HSQC + COSY + TOCSY

We now move on to a longer supersequence containing four modules, with a correspondingly larger ρ_t value of 3.22. The sensitivity of the HSQC module is practically the same as in the NOAH-2 BS supersequence just described: however, the COSY and TOCSY modules expose one weakness of the HMBC module which has so far been overlooked. In principle, the HMBC module should only excite magnetisation of protons which have long-range couplings to ^{13}C (which we could, for example, denote as $^1\text{H}^{\text{C(LR)}}$). This magnetisation pool is distinct from both the directly coupled protons ($^1\text{H}^{\text{C}}$), as well as protons which are not coupled to any ^{13}C at all ($^1\text{H}^{\text{!C}}$). Unfortunately, this is not the case: it is not actually possible to separate the $^1\text{H}^{\text{C(LR)}}$ and $^1\text{H}^{\text{!C}}$ magnetisation pools. The HMBC excites both of these magnetisation sources, dephases the latter using CTP gradients, and detects the signal arising from the former.

*In fact, the final 180° pulse in the HMBC module could also be removed: this is likely to give a further boost in SNR, as discussed in § 4.3.7. However, this was not done here.

This means that the COSY and TOCSY modules, which rely on $^1\text{H}^{1\text{C}}$ magnetisation, will have substantially lower sensitivities. The signal detected in these two modules derives only from polarisation that has recovered during the previous two acquisition periods, as shown in entry 5 of table 4.2 (the values of A for the COSY and TOCSY are 0.36 and 0.28 respectively). That said, this is in fact not likely to be an issue *even* for sensitivity-limited samples. Because the intrinsic sensitivity of the HMBC is orders of magnitude lower than that of the COSY and TOCSY, the COSY and TOCSY spectra still have greater intensities than the HMBC, even after these large losses in sensitivity. Thus, as long as the entire supersequence is acquired with enough scans such that the HMBC SNR is sufficient, the COSY and TOCSY will *also* have acceptable SNR. This is proven by the spectra shown in fig. 4.6.

A rather more insidious problem is that different signals relax at different rates: thus, the COSY and TOCSY spectra (or indeed, any homonuclear module) will have uneven intensities and are frequently asymmetric. This can be seen in the COSY spectrum, where a pair of asymmetric crosspeaks are highlighted. Adding a period of isotropic mixing before the COSY module⁵⁰ can ameliorate this to some extent (although the spectra in fig. 4.6 were not acquired with this).

NOAH-5 BS_N⁺SCT: HMBC + ^{15}N seHSQC + HSQC + COSY + TOCSY

As the final example, we add a further magnetisation pool to the mix, namely protons directly coupled to ^{15}N (i.e. $^1\text{H}^{\text{N}}$). As of the time of writing, the implementation of multiple-FID experiments on Bruker spectrometers limits N to a maximum of 5, so a supersequence such as this NOAH-5 BS_N⁺SCT is the current limit. (However, there is no *scientific* argument forbidding $N > 5$, and it is likely that in future versions of TopSpin this restriction will be lifted.)

The values of A for each module are given in entry 6 of table 4.2. If the HMBC module is placed at the beginning of the supersequence, then in order to preserve *both* $^1\text{H}^{\text{N}}$ and $^1\text{H}^{\text{C}}$ magnetisation, the zz -filter must be extended to include ^{15}N pulses.⁵¹ As before, this modification leads to a slight drop in sensitivity for the HMBC (5%).

After this, the ^{15}N seHSQC and ^{13}C HSQC modules both suffer drops in sensitivity. For the ^{15}N seHSQC, this is partly because of imperfect preservation of $^1\text{H}^{\text{N}}$ magnetisation by the HMBC, but also stems from the addition of the zz isotope-selective pulse (ZIP) element to the seHSQC pulse sequence; this is described further in § 4.3.3. On the other hand, for the ^{13}C HSQC, the sensitivity loss stems purely from imperfect retention of $^1\text{H}^{\text{C}}$ magnetisation by the HMBC. Finally, because the HMBC dephases $^1\text{H}^{\text{IX}}$ magnetisation, the COSY and TOCSY at the end have lower sensitivities: however, as discussed above, this is not an issue in practice.

It is also possible to move the ^{15}N seHSQC module to the front: this gives it a slightly greater sensitivity, at the cost of the HMBC (entry 7, table 4.2). In general, these two modules tend to

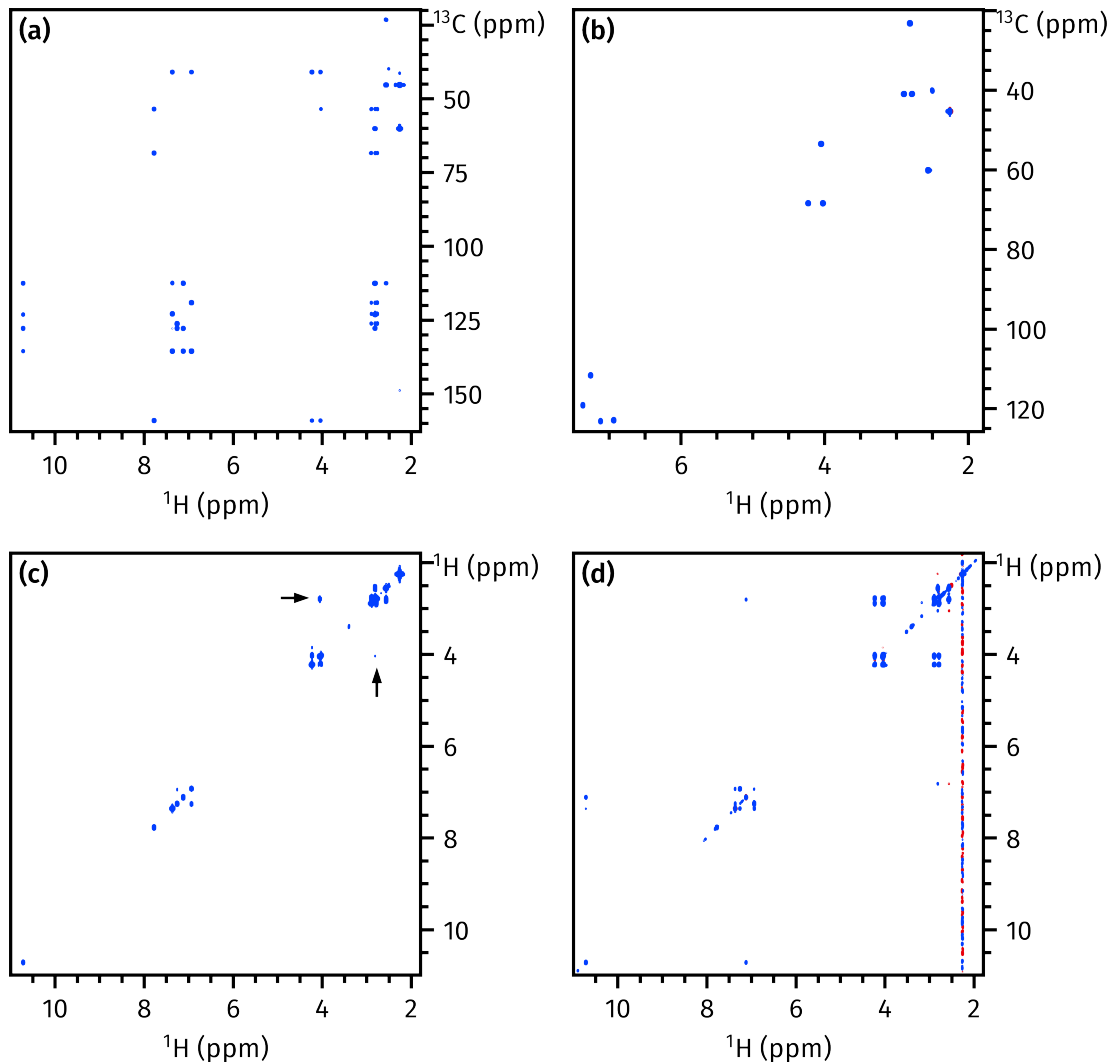


Figure 4.6: Spectra obtained from a NOAH-4 BSCT supersequence. (a) HMBC. (b) HSQC. (c) COSY; a pair of crosspeaks with asymmetric intensities is highlighted with black arrows. (d) TOCSY (60 ms DIPSI-2 mixing). Despite the COSY and TOCSY having only around 30% of their sensitivity compared to standalone experiments, the intensity of the spectra obtained is still perfectly acceptable (the contour levels chosen are 1–2 orders of magnitude larger than for the HMBC). Data code: 7Z-220224.

have comparable sensitivity, and which of these two arrangements is better depends on which module the sensitivity needs to be optimised for.

Finally, the value of ρ_t given here of 3.74 represents an effective upper limit on the time-saving factor. Although ρ_t increases with N , the extent to which it deviates from the ideal value of N also increases: it is very difficult to obtain $\rho_t > 4$, even with five modules in the supersequence. Of course, it is possible to increase ρ_t further by lengthening the recovery delay d_1 used for the experiments: for example, if d_1 is increased to 2 s from its present value of 1.5 s, ρ_t increases to 3.94. Obviously, this can only be pushed so far before it becomes meaningless.

4.2 GENESIS: automated pulse programme creation

In this section, I discuss the development of the GENESIS (GENEration of Supersequences In Silico) website (fig. 4.7).^{*} As its name suggests, GENESIS is a tool which programmatically generates (Bruker) pulse programmes for arbitrary NOAH supersequences. The website also provides extensive instructions on acquiring and processing NOAH data. The present version of GENESIS is available at <https://nmr-genesis.co.uk>; the source code can also be obtained from <https://github.com/yongrenjie/genesis>.

The screenshot shows the 'Module selector' interface of the GENESIS website. It features five columns of modules:

- 13C HMBC**: Contains 'None' (selected) and 'HMBC'.
- 15N-1H**: Contains 'None' (selected), 'HMQC', and 'seHSQC'.
- 13C-1H #1**: Contains 'None' (selected), 'HSQC', 'F2 J-HSQC', 'HSQC-COSY', and 'HSQC-TOCSY'.
- 13C-1H #2**: Contains 'None' (selected), 'HSQC', 'F2 J-HSQC', 'HSQC-COSY', 'HSQC-TOCSY', 'HMQC', 'seHSQC', 'F2 J-seHSQC', and 'seHSQC-TOCSY'.
- 1H-1H**: Contains 'None' (selected) and a grid of other modules: COSY, COSY-ROESY St, COSY (QF), COSY-NOESY EA, CLIP-COSY, COSY-NOESY St, DQF-COSY, COSY-TOCSY EA, TOCSY, COSY-TOCSY St, NOESY, 2DJ (QF), ROESY, PSYCHE 2DJ, ROESY (ad.), 1D PSYCHE, COSY-ROESY EA, and 1D TSE-PSYCHE.

Below the modules are several buttons: 'Reset all', 'Download pulse programme', 'Read the FAQ', and 'Download processing scripts'. A 'Developer mode' toggle switch is located in the top right corner.

Figure 4.7: The front page of the GENESIS website (<https://nmr-genesis.co.uk>), as of 10 September 2022.

4.2.1 Motivation

The preceding discussion in § 4.1 makes clear how supersequences may be constructed from a large variety of modules. Furthermore, modules which consume (and preserve) the same magnetisation pools may be directly swapped out for one another. Thus, for example, the NOAH-2 SC supersequence can in fact be generalised to any $^1\text{H}^{\text{C}}$ module plus any $^1\text{H}^{!C}$ module. Very broadly speaking, we may define a generic supersequence as having any or all of the following:

*Although placing this section first may at first glance appear to be nonchronological, in that the GENESIS paper³ was published later than much of the other work in this chapter, an early version of the GENESIS tool was in fact created much earlier (by July 2020).

- an HMBC module, which actually uses $^1\text{H}^{1\text{X}}$ magnetisation but can be placed at the front as discussed in the NOAH-3 BSC example above;
- a $^1\text{H}^{\text{N}}$ module;
- one or more $^1\text{H}^{\text{C}}$ modules (it is possible to partition the $^1\text{H}^{\text{C}}$ magnetisation pool between two modules, as will be discussed in § 4.3.4);
- finally, a $^1\text{H}^{1\text{X}}$ module (or a COSY+X combination) which consumes all remaining bulk magnetisation.

In the first NOAH paper in 2017,³² a total of 285 ‘viable’ supersequences were already listed. If we further take into account some of the new modules which were developed over the course of my DPhil (§ 4.3), the generic formula above provides for over 4000 viable supersequences.* (‘Non-viable’ sequences would be those which have unwanted sensitivity losses, such as the wrongly-ordered NOAH-2 CS supersequence.)

In spite of this diversity, *only around 45 pulse programmes* had been made available prior to the GENESIS website (these were distributed either via the supplementary information of NOAH papers, or the Bruker User Library). The reason for this gap between theory and practice is simple: traditionally, pulse programmes must be written by hand, which is a laborious and fairly error-prone process made worse by the sheer length of NOAH experiments. Doing this for thousands of supersequences is clearly impractical. Furthermore, each time a new module is developed, or an old module is improved, updating every relevant supersequence would in itself be a mammoth task.

To bridge this gap, I turned towards the *programmatic* generation of pulse programmes.[†] This not only allows users to create their own customised supersequences, but also provides an easy way for pulse sequence updates to be rapidly disseminated to the NMR community, as new pulse programmes can be accessed immediately upon the deployment of updated source code. Furthermore, the website can serve as a ‘one-stop’ shop where—after downloading pulse programmes—users may download associated NOAH processing scripts and also access instructions on how to run NOAH experiments. This information did already exist, but was scattered across several different websites and/or supplementary information documents, and would have been needlessly confusing to a new user (not even taking into account the different versions of scripts available in different publications).

*This is also ignoring the ‘parallel’ supersequences, which are discussed in § 4.5. The support for parallel supersequences in GENESIS is not complete: integrating these fully would require substantial changes to the user interface, which I have not had time to do.

[†]This is actually a bit of a lie: GENESIS was initially created *for my own convenience*. Throughout this chapter, I have had to perform many comparisons of different supersequences, and this tool spared me from having to write everything by hand (and—more often than not—subsequently discover mistakes which invalidated the results). Of course, it soon became apparent that it could find much wider use.

4.2.2 Implementation details

I will now describe a few features of GENESIS pulse programmes, as well as how their generation is implemented in code. GENESIS is written in TypeScript; during website deployment, this is compiled to JavaScript, which can then be directly executed in the client's web browser. No server-side code is required, meaning that the GENESIS web page is actually a static site; it is currently hosted using GitHub Pages.

Overall structure

The structure of a pulse programme can loosely be separated into three parts, which are shown in listing 4.1:

1. the *preamble*, which consists of everything up until the beginning of the actual pulse sequence (signified by the `ze` command). This includes header comments as well as definitions of parameters, such as delays and pulse widths;
2. the *main section*, which contains the actual pulse sequence;
3. the *epilogue*, which contains phase cycle information as well as footer comments describing each parameter. Instructions for generating shaped pulses using Bruker's WaveMaker software are also included here, as well as instructions for automatic processing (§ 4.2.3), and comments indicating how the pulse programme was generated (for reproducibility purposes).

The algorithm used for pulse programme construction can be broken up into similar sections. The construction of the preamble and main section is largely accomplished through the collation of module-specific information, the most important of which are:

- descriptive information about the modules themselves, which go into the header comments;
- parameter definitions, which are collated to form the preamble. Duplicates must be removed here to avoid errors; and
- the pulse programmes themselves, which are directly concatenated to form the main section.

These, as well as other smaller bits of information (e.g. relevant citations, appropriate processing scripts), are stored within `NOAHModule` objects (an example is provided in listing 4.2). Each distinct module corresponds to one such object. Therefore, if one wants to add a new module to GENESIS, most of the work can be completed by simply defining a new `NOAHModule` object: no changes to the algorithm itself are needed.

```

/* PREAMBLE */
; gns_noah2-SCqf
; 13C HSQC
; 1H magnitude-mode COSY
"d4      = 0.25s/cnst2" ; 13C INEPT
"in0     = inf1/2"       ; 13C HSQC increment
"in11    = 2*dw"         ; COSY increment
; ...
define delay DC_HSQCa
"DC_HSQCa   = d4-p14/2"
"10        = td1/2"       ; TD1/NBL

/* MAIN SECTION */
1 ze
4 d1 st0
; MODULE 1 - HSQC
(p1 ph0):f1
; ...
goscnph30 cpd2:f2
50u do:f2
2m st
; MODULE 2 - COSY
(p1 ph12):f1
; ...
go=2 ph26
1m iu1      ; TD1/NBL counter
1m igrad EA ; echo-antiecho gradients
1m id11     ; COSY t1
30m wr #0 if #0 zd
if "l1 % 2 == 0" {
  1m id0      ; HSQC t1
}
lo to 4 times 10
exit

/* EPILOGUE */
ph0=0
;gpnam4: SMSQ10.100
;gpz4: 70% (13C CTP)
;cpd2:wvm:wudec: cawurst_d-20(220 ppm, 1.4 ms; L2H)
;d4: 1/4J(CH)
;auprog: noah_hsqc:noah_cosy QF
;module identifiers: C_HSQC H_COSY_QF
;pulse programme created by genesis-v2.2.2, https://nmr-genesis.co.uk
;Sun Sep 11 2022 16:04:54 GMT+0800 (Malaysia Time)

```

Listing 4.1: Abridged GENESIS pulse programme for the NOAH-2 SC supersequence shown in fig. 4.1a.

```

let shortDescription = `; 13C HSQC`

let preamble =
  "d4      = 0.25s/cnst2"           ; 13C INEPT
; ...
"D[ID]a  = d4-p14/2"
` // [ID] is later replaced with the module code

let pulprog =
  `; 13C-1H HSQC

  ; INEPT
  (p1 ph0):f1
  ; ...
  goscnp ph30 cpd2:f2
  50u do:f2
`


const mod = new NOAHModule(
  "C_HSQC",           // internal module code
  "c13",              // module category
  "S",                // single-letter code, Table 4.1
  [],                 // relevant citations (if any)
  "noah_hsqc",        // AU programme for processing
  shortDescription,   // short description
  [AF_EDIT],          // available acquisition flags
  preamble,           // preamble text
  pulprog,            // pulse programme text
  1,                  // number of FIDs
  false               // flag for 'parallel' modules
);
export default mod;

```

Listing 4.2: An excerpt of the NOAHModule object for the HSQC module shown in fig. 4.3a (internal code C_HSQC).

To put the epilogue together, the pulse programme constructed so far is scanned for pulse phases, shaped pulses, and all other parameters. Using predefined lookup tables, GENESIS then creates pulse phase definitions, WaveMaker directives (where appropriate), and comments containing textual descriptions of each parameter. These comments are not merely cosmetic; they are also displayed in the `ased` screen when setting up an experiment. Finally, instructions for automatic processing of the NOAH data (to be explained in § 4.2.3) are added to the bottom, together with a timestamp and the version number of GENESIS for reproducibility purposes.

Phase/delay incrementation and looping

Since NOAH experiments are 2D experiments, there is one additional complication: the pulse programme must contain appropriate looping statements, together with pulse phase and delay incrementation, in order to correctly generate the indirect dimension. In many existing NOAH pulse programmes, looping in 2D experiments was written using the equivalent of nested for loops (listing 4.3, *left*). Although this suffices for the vast majority of supersequences, whenever any parameter must be incremented in a different manner (e.g. in parallel supersequences where multiple supersequences are interleaved, or when using the PSYCHE module which has a different number of t_1 increments), the nested loop structure must be modified, which is not easy to reason about. I therefore opted to change the structure to use only one loop, and to control the phase and delay incrementation using modular arithmetic (listing 4.3, *right*). The outcome is entirely equivalent, but this strategy allows for other cases to be implemented simply by adding another check on the loop counter L1.

<pre>"10 = td1/4" 1 ze 3 1m 4 d1 st0 ; ... (pulse sequence goes here) ; in inner loop 1m igrad EA ; HSQC gradients 1m id11 ; COSY t1 30m wr #0 if #0 zd lo to 4 times 2 ; in outer loop 1m ip5*2 ; HSQC 13C 90 1m ip30*2 ; HSQC receiver 1m id0 ; HSQC t1 lo to 3 times 10 end</pre>	<pre>"10 = td1/2" "11 = 0" 1 ze 4 d1 st0 ; ... (pulse sequence goes here) ; on every pass 1m iu1 ; loop counter 1m igrad EA ; HSQC gradients 1m id11 ; COSY t1 30m wr #0 if #0 zd ; on every second pass if "11 % 2 == 0" { 1m ip5*2 ; HSQC 13C 90 1m ip30*2 ; HSQC receiver 1m id0 ; HSQC t1 } lo to 4 times 10 end</pre>
---	---

Listing 4.3: Implementation of phase/delay incrementation and looping in previous NOAH sequences (*left*, using nested loops) and in GENESIS (*right*, using modular arithmetic).

Parameter standardisation

Each NOAH module contains a number of parameters, including pulse widths, delays, gradient amplitudes, and shaped pulse waveforms. Since different modules are stored separately (as different `NOAHModule` objects), directly concatenating their pulse programmes may lead to conflicting parameter definitions if care is not taken. GENESIS avoids this by maintaining a global table of parameter definitions which are applicable to all modules: when new modules are added, they must be checked against this to ensure that there are no inconsistencies.

In general, where possible, these parameters are chosen to be consistent with pulse programmes in the Bruker standard library: thus, for example, `P1` is the ${}^1\text{H}$ 90° pulse width, and `CNST2` is the ${}^1J_{\text{CH}}$ value used for calculating INEPT (and other) delays. This makes it easy to read in parameters either from the *prosol* relation tables in TopSpin, or from other existing parameter sets. Some delays are module-specific and do not need to be reused, and in standard library sequences, are often labelled as `DELTA1`, `DELTA2`, and so on. To avoid conflicting definitions and also to improve readability, I renamed these such that they include the name of the module: thus, in a ${}^{13}\text{C}$ HSQC these may be labelled `DC_HSQC_1`. Here, `C_HSQC` is the name associated with the `NOAHModule` object.

If combined with some caution when adding new modules, these measures ensure that there will be no parameter clashes between the modules *within a given supersequence*: we may view this as a *local uniqueness* of parameters. However, the impact of this design choice is even more far-reaching: since parameters are stored globally, they will always have the same value in *all possible supersequences* (or in other words, the parameters are *globally unique*). This makes it exceptionally easy to set up multiple different supersequences in TopSpin using GENESIS pulse programmes, as virtually all of the parameter values may simply be copied from a previous NOAH dataset.

One potential issue with this strategy is that TopSpin provides only a finite number of named pulse widths (for example). Thus, there are only so many different parameters which can be stored in a global table before running into inevitable conflicts. A workaround would be to sacrifice the global uniqueness of each parameter, and only have it be unique within a given supersequence. Fortunately, this situation has not (yet) surfaced.*

*The number of *pulse phases* in GENESIS is in fact dangerously close to the maximum number of 32. However, the global uniqueness criterion is not really important for pulse phases, because—unlike, say, delays—pulse phases are hardcoded in the pulse programme, and cannot be copied from one dataset to another. So, if necessary, I could dispense with the global uniqueness for pulse phases only, at the cost of some increased code complexity. I did briefly contemplate this possibility, but since I am at the end of my DPhil and am unlikely to add any new modules soon, this will likely remain a hypothetical.

Parameter descriptions

At the epilogue of the pulse programme, extra comments are added for every parameter present in the pulse programme. Most of these are purely textual in nature, and appear in the TopSpin based parameter setup screen; this naturally helps to make the pulse programmes as easy to use as possible. However, some of these comments have special meanings: gradient amplitudes and shapes, for example, are specified in a way which allows them to be automatically populated using the gppp Python script packaged with TopSpin. Furthermore, WaveMaker directives are also specified for some shaped pulses, allowing them to be created in an on-the-fly manner using the wvm command. This means that the user (generally) need not separately download and install a set of shaped pulses.

Module choice

One final issue which must be overcome is the fact that some NOAH modules may be implemented differently depending on the supersequence which it is being used in. The HMBC module described in § 4.1.3 is one such example: the form of the *zz*-filter depends on whether the HMBC module is required to retain ${}^1\text{H}^{\text{C}}$ and/or ${}^1\text{H}^{\text{N}}$ magnetisation for subsequent modules. Thus, in the NOAH-2 BS supersequence the *zz*-HMBC must be used, but in the NOAH-2 SB supersequence the ‘original’ HMBC without a *zz*-filter is preferable. Since these have different pulse programmes, each of these ‘versions’ of the HMBC are described by *separate* NOAHModule objects (C_HMBC_CF and C_HMBC_NOF respectively).

However, it is unlikely that the majority of users would want to manually configure the supersequence in such detail by selecting NOAHModule objects. Thus, the GENESIS web interface actually hides these different versions from the user, only showing one button labelled ‘HMBC’. Under the hood, the logic above is used to deduce the correct version of the HMBC based on what other modules the user has chosen (fig. 4.8).

Should the user indeed want to control the exact module used, the website also offers a ‘developer mode’ switch: turning this on allows the user to directly choose the desired module. (Enabling developer mode also reveals a handful of extra modules which were used exclusively for my DPhil and are not generally of interest.)

Tests

One risk inherent in any NMR experiment is the possibility of causing spectrometer damage when executing malformed pulse programmes. To minimise the chances of this, each version of GENESIS is checked against a series of automatic tests before being released. The most important tests ensure that every module containing decoupling statements also turns off decoupling

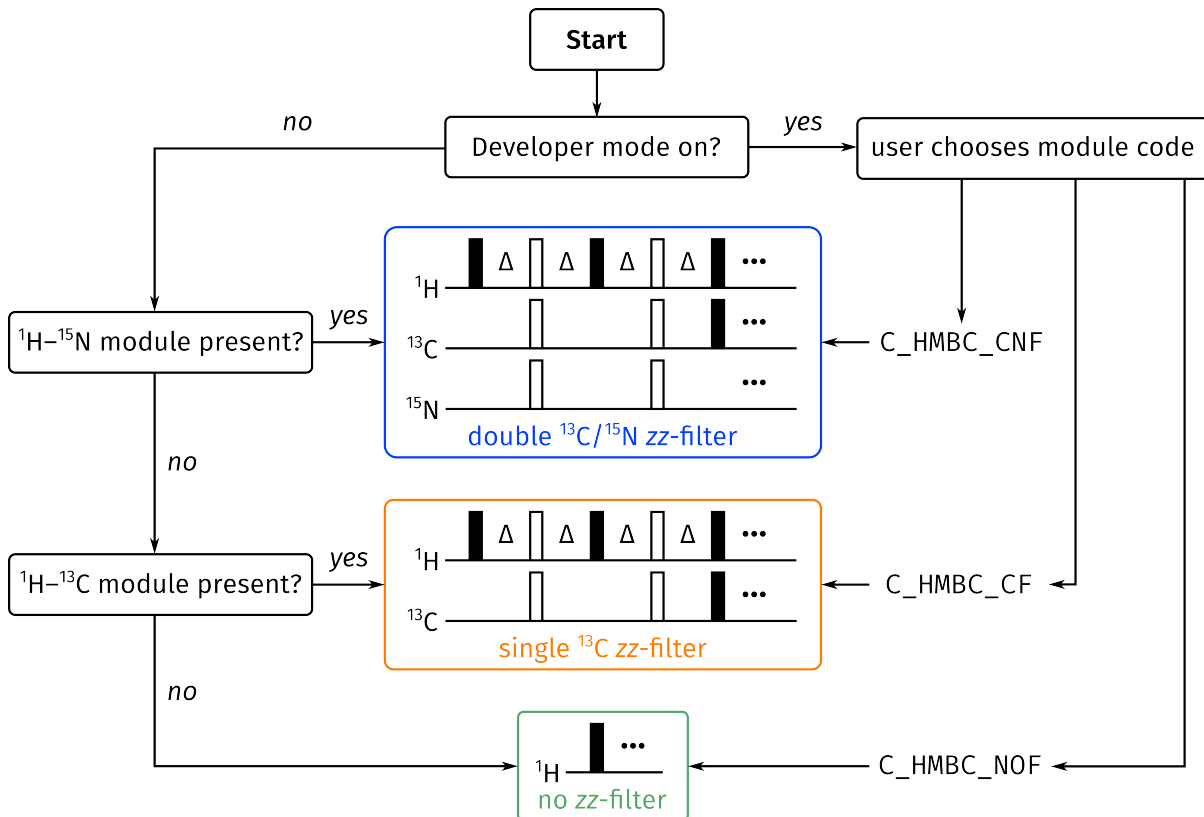


Figure 4.8: Flowchart showing how the correct ‘version’ of the HMBC module is determined when constructing a supersequence using GENESIS. When developer mode is on, the user directly chooses module codes corresponding to each version of the HMBC. When it is off, the appropriate version is chosen based on which other modules are present in the supersequence.

immediately after acquisition. On top of this, there are also a series of regression tests where (some representative) supersequences are checked against the same ones produced by previous versions of GENESIS: any differences in these are automatically flagged for review.

None of this can actually *stop* wrong pulse programmes from being written. For example, if a module with an incorrect pulse programme is created, then any supersequence containing that module will not be fully functional. There is no adequate way to check for this, as GENESIS cannot verify the correctness of an NMR experiment. However, the inclusion of the tests described above minimises the chances of creating pulse programmes which are outright dangerous; any mistakes will at worst lead to the wrong (or no) data being obtained. It should be mentioned that this is not a *new* problem which GENESIS introduces: errors in pulse programmes can arise equally easily (if not even more easily) when they are handwritten.

Reproducibility

The final section of a GENESIS pulse programme consists of comments intended solely for reproducibility purposes. These comments indicate exactly which NOAHModule objects were used to create the pulse programme, as well as a GENESIS *version number*. Since GENESIS has been updated with some regularity during my DPhil, each release is assigned a version number, which broadly follows the principles of *semantic versioning*. Old versions of GENESIS may be accessed by navigating to the specific URL <https://nmr-genesis.co.uk/X/Y/Z> for version X.Y.Z. Then, after enabling developer mode, the specific list of NOAHModule objects may be input in order to create the corresponding pulse programme. Thus, these two pieces of information together allow all GENESIS pulse programmes to be exactly regenerated whenever necessary: this is important in ensuring that the data thus acquired are reproducible.

How smart is GENESIS?

Having written several pages of text about the *features* present in GENESIS, it is tempting to think that it is ‘intelligent’ in its design of pulse programmes. At various points in time, extensions to the concept have even been proposed: in GENESIS, the building blocks used for pulse programme construction are NOAH modules, but one could envision breaking up these into the smallest possible units of pulses, delays, gradients, and so on, and using something like GENESIS to construct *arbitrary* pulse programmes. This was even speculatively mentioned in the GENESIS paper.³

However, in truth, GENESIS is a long way from being able to do such things. In particular, it does not have any actual understanding of the Bruker pulse programming syntax: *almost* all of the pulse programme instructions are hardcoded as strings.* Thus, the creation of pulse programmes is not *truly* being done from the ground up: instead of ‘combining pulse sequence elements’, a more accurate description is that the *text* corresponding to different pulse sequence elements is being concatenated. There is actually a substantial amount of brute force involved, and this is also why it is not possible for the automated tests to really check the correctness of the pulse programmes.

Of course, it is hardly a trivial task to write something more sophisticated: I would need to construct abstract representations of each pulse sequence element (pulses, delays, etc.), and functions which could translate these into actual pulse programme commands. I feel that I would need rather more knowledge in computer science, especially a course on compilers. However, this was something I would have very much liked to do had I had more time! I will also leave open the possibility of—although I refrain from *committing to*—expanding the GENESIS code

*There are several exceptions where some parameters (for example, loop counters in DIPSI mixing) are dynamically generated, mainly to avoid clashes between different parts of the supersequence.

in this direction as a future personal project.

4.2.3 Processing improvements

Finally, I touch on a few small improvements in processing NOAH data. These are not strictly part of the GENESIS website, but having a unified source for pulse programmes and processing scripts does make it substantially easier to propagate these developments to NMR users.

AU processing scripts

In the processing of NOAH experiments, the first step is to split up the FIDs of different modules into different files: this is performed by the `splitx_au` AU programme. However, after that is done, `splitx_au` also calls on auxiliary AU programmes to process each of the resulting datasets. Each module is associated with a specific auxiliary AU programme: for example, HSQC data are processed with `noah_hsqc`, COSY data with `noah_cosy`, and so on.

Previously, these auxiliary AU programmes had to be specified manually using the `USERP1` through `USERP5` processing parameters, which apply to the first through fifth modules in a supersequence respectively. Although this was a perfectly serviceable setup, I realised that it was a source of annoyance when running multiple different supersequences, as modules would often be processed incorrectly if these processing parameters were inadvertently copied from an old dataset. Even when just running a single supersequence, this would nonetheless represent an extra cognitive load, especially for users who are not familiar with NOAH experiments.

Since the pulse programme specifies which modules are run, and the ‘correct’ processing parameters also depend on the modules, it stands to reason that the processing parameters can be encoded in the pulse programme itself. Specifically, the auxiliary AU programmes are part of the `NOAHModule` objects (see listing 4.2), which makes this information accessible to the GENESIS algorithm. Using this, GENESIS inserts a line near the bottom of every pulse programme, specifying which AU programmes are to be used for each of the modules present. I furthermore modified the `splitx_au` AU programme to parse the pulse programme for this extra information. The overall effect is that the `USERP1` through `USERP5` parameters no longer need to be specified, as long as a GENESIS pulse programme and a recent version of `splitx_au` are used. However, if the parameters *are* specified, they will override the ‘default’ processing instructions found in the pulse programme. This maintains backwards compatibility with, for example, parameter sets which have been previously saved; it also allows advanced users to customise the processing, if so desired.

Non-uniform sampling

Another small improvement relates to the implementation of non-uniform sampling (NUS) in NOAH supersequences. Because the indirect dimension in NOAH pulse programmes is generated through explicit looping, instead of the `mc` macro used in most 2D experiments, the standard NUS implementation in TopSpin cannot be used. Instead, one must explicitly define a list of t_1 increments to sample (stored as the `VCLIST` parameter). This is true regardless of which looping implementation in listing 4.3 is used.

On top of this, the original NUS implementation⁴⁹ used a Python script (`noah_nus.py`) to modify the NOAH pulse programme such that it only sampled t_1 values from this list. This, however, necessitated storing two copies of the same pulse programme, one with NUS and one without.

There is no way to circumvent the requirement for the `VCLIST` parameter, but in GENESIS pulse programmes, I opted to implement NUS as an acquisition flag which could be toggled. This means that there is no need to store two versions of the same pulse programme, and also makes it easier to seamlessly switch between non-uniform and uniform sampling as desired.

Unfortunately, this change is slightly more problematic than the `splitx_au` change: this new NUS implementation is not backwards-compatible, so requires a new NUS setup script, which I have called `noah_nus2.py`. This script cannot be used with old pulse programmes, and the old NUS script cannot be used with GENESIS pulse programmes. Nevertheless, it does represent a real improvement over the previous implementation: the GENESIS website also helps to make the changeover as smooth as possible.*

4.3 Discussion of individual modules

The GENESIS website makes it easy to implement new modules and update old ones: simply changing the underlying `NOAHModule` objects is sufficient to propagate changes to all supersequences generated using those modules. In this section, I discuss a number of new pulse sequence developments made in the course of my DPhil; all of these have been successfully implemented in GENESIS and are available to download.

With regards to sensitivity analyses, since this chapter discusses the design of *individual modules*, I have chosen to focus almost entirely on comparing the SNRs of the modules. This is related to the SNR factor A (although my basis for comparison is not always the standalone experiments, so the definition is not totally the same). I ignore the quantities ρ_t and ε_t , as these depend on

*That said, there were still some bugs in the new NUS implementation right up until August 2022, so this change was not as smooth-sailing as I had hoped for.

the *supersequence* within which the module is used, and can be trivially improved by adding more modules to the supersequence under consideration. These numbers reflect the utility of the NOAH *technique*, but are not relevant to the individual modules from which they are constructed.

4.3.1 ^{13}C sensitivity-enhanced HSQC

The first new module is the sensitivity-enhanced HSQC (seHSQC) experiment, which provides up to $2\times$ increased SNR over a standard (echo–antiecho) HSQC.* In the original version of the seHSQC (fig. 4.9a), developed by Cavanagh, Rance, and Kay ('CRK'),^{53,54} this is accomplished through the so-called *preservation of equivalent pathways* (PEP) technique,⁵⁵ which I now provide a product operator analysis of.

Just like in a standard HSQC, the seHSQC experiment begins with an INEPT block and t_1 evolution. At the end of t_1 , there are two terms which are cosine- and sine-modulated with respect to Ω_S . In the standard HSQC, only the $2I_zS_y$ term is returned into observable ^1H magnetisation by the reverse INEPT block (see § 1.4.4). In contrast, the PEP block transfers *both* terms back to ^1H and subsequently detects both. Specifically, in the echo experiment it accomplishes the transfer $2I_zS_y \rightarrow I_y$ and $2I_zS_x \rightarrow I_x$, meaning that the density operator at the end of t_1

$$-2I_zS_y \cos(\Omega_S t_1) - 2I_zS_x \sin(\Omega_S t_1) \quad (4.6)$$

is transformed into

$$-I_y \cos(\Omega_S t_1) - I_x \sin(\Omega_S t_1) \quad (4.7)$$

just prior to acquisition. During the FID, only the -1 -coherence component of this is detected:

$$\frac{1}{2i}I_- \cos(\Omega_S t_1) - \frac{1}{2}I_- \sin(\Omega_S t_1) = \frac{1}{2i}I_- \exp(-i\Omega_S t_1), \quad (4.8)$$

yielding a signal of

$$\frac{1}{2i} \exp(-i\Omega_S t_1) \exp(i\Omega_I t_2), \quad (4.9)$$

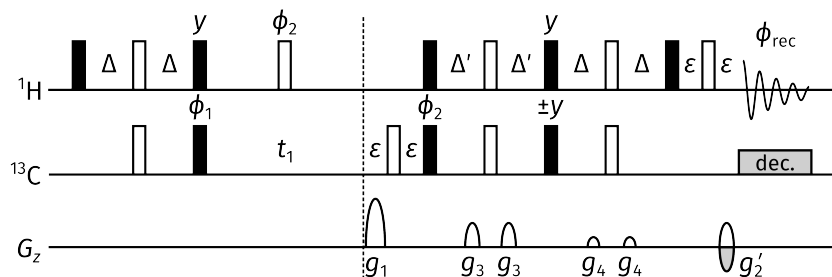
which has a $2\times$ larger amplitude than the original EA HSQC (eq. (1.96)). The antiecho experiment can be similarly analysed. Since the seHSQC has the same noise level as in an EA HSQC, this also corresponds to a $2\times$ increase in SNR.

It should, however, be noted that the PEP transfer is only fully attained for *IS* spin pairs if the delay Δ' is set to $1/(4 \cdot ^1J_{\text{CH}})$. For $I_2\text{S}$ or $I_3\text{S}$ spin systems, no gain in sensitivity is accomplished

*Since a States HSQC has $\sqrt{2}$ times the SNR of an EA HSQC (as shown in fig. 1.8), this also means that the seHSQC has a $\sqrt{2}$ SNR improvement over a States HSQC. The literature can be somewhat confusing on this point: sometimes the gain in signal is even conflated with the gain in SNR. The clearest exposition I have found is that provided by Kontaxis et al.⁵²

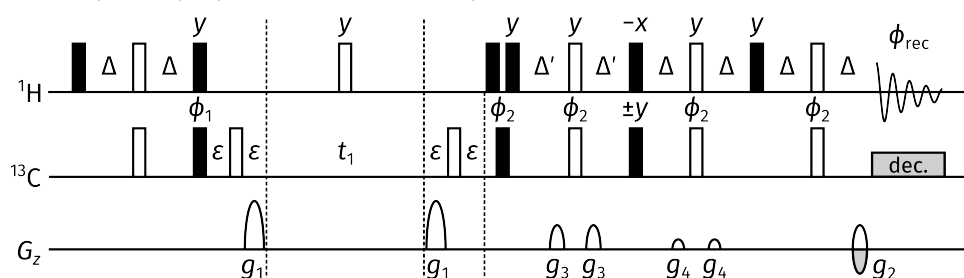
(a) CRK seHSQC

$\cos^1 H^C$	$z -y$	$-xz -zy$	$-zy yz$	$-x z$	$-z y$	$-y$
$\sin^1 H^C$	$z -y$	$-xz -zy$	$-zx yx$	$-yx \pm yz$	$\mp x \mp x$	$\mp x$
$^1H^{iC}$	$z -y$	$y y$	<i>...dephased</i>			



(b) NOAH seHSQC1

$\cos^1 H^C$	$z -y$	$-xz -zy$	zy	yz	$x x$	$-yz -yz$	x
$\sin^1 H^C$	$z -y$	$-xz -zy$	$-zx$	$-yx$	$-yx \pm zz$	$\pm zz \pm xz$	$\mp y$
$^1 H^{!C}$	$z -v$	$v v$	v	x	$-x -x$	$x -z$	z



(c) NOAH seHSOC2

$\cos^1 H^C$	$z -y$	$-xz$	$-xz$	$y y$	xz	zy	$-zy$	yz	$-x z$	$-z y$	$-y$
$\sin^1 H^C$	$z -y$	$-xz$	$-xz$	$y y$	xz	zy	$-zx$	yx	$-yx \pm yz$	$\mp x \mp x$	$\mp x$
$^1 H^{!C}$	$z -y$	y	z	$-z x$	$x -z$		$z -y$	$y y$	$-y -z$	$-z z$	z

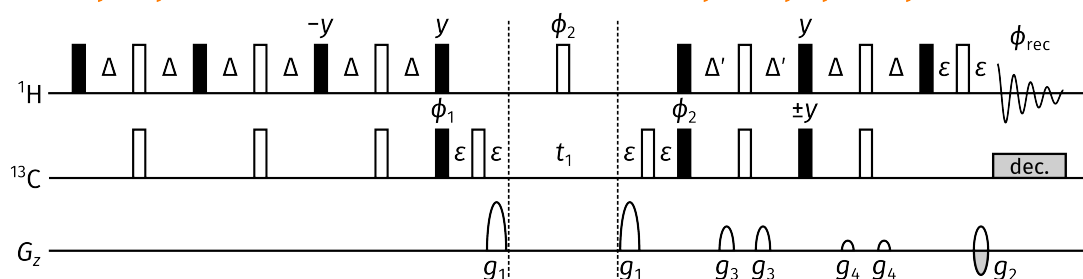


Figure 4.9: Sensitivity-enhanced HSQC sequences discussed in this section, along with product operator analysis. This analysis is provided only for the first step of the phase cycle, and assumes an *IS* spin pair with $\Delta' = 1/(4 \cdot ^1J_{\text{CH}})$. ‘cos $^1\text{H}^{\text{C}}$ ’ refers to the component of the $^1\text{H}^{\text{C}}$ magnetisation which is cosine-modulated during t_1 . (a) Cavanagh–Rance–Kay seHSQC. (b) NOAH seHSQC1 module. (c) NOAH seHSQC2 module. Phase cycling is performed with $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x)$, and $\phi_{\text{rec}} = (x, -x, -x, x)$. The delay Δ is set to $1/(4 \cdot ^1J_{\text{CH}})$; see the text for a discussion of Δ' . A value of 145 Hz was used for $^1J_{\text{CH}}$. The pulses marked with a phase of $\pm y$ are applied with a phase of y in the echo experiment and $-y$ in the antiecho. Gradient amplitudes are $(g_1, g_2, g'_2, g_3, g_4) = (70\%, \pm 35.2\%, \pm 17.6\%, 11\%, -5\%)$.

with this setting of $\Delta' = 1/(4 \cdot {}^1J_{\text{CH}})$. It is more common, therefore, to shorten Δ' to $1/(8 \cdot {}^1J_{\text{CH}})$: this sacrifices some transfer efficiency for *IS* systems, but allows for some sensitivity enhancement for *I₂S* and *I₃S* systems (table 4.3).

Spin system	Theoretical sensitivity enhancement		
	$\Delta' = 1/(4 \cdot {}^1J_{\text{CH}})$	$\Delta' = 1/(8 \cdot {}^1J_{\text{CH}})$	$\Delta' = 1/(12 \cdot {}^1J_{\text{CH}})$
<i>IS</i>	2	1.71	1.5
<i>I₂S</i>	1	1.41	1.37
<i>I₃S</i>	1	1.21	1.25

Table 4.3: Theoretical sensitivity enhancements for *IS*, *I₂S*, and *I₃S* spin systems in the seHSQC, as a function of the delay Δ' . The values are taken from Schleucher et al.⁵⁶

NOAH seHSQC versions

Just like the original EA HSQC (fig. 1.7), the CRK seHSQC must be modified in order to be compatible with NOAH supersequences. In particular, the CTP gradients in the CRK seHSQC dephase the bulk ${}^1\text{H}^{1\text{C}}$ magnetisation pool: we would very much like it to return that to $+z$ instead, so that it can be sampled in later homonuclear modules (or, indeed, a HMBC module). This experiment is more tricky to adapt than the HSQC, because there are *three* different magnetisation components to juggle: the cosine-modulated ${}^1\text{H}^{\text{C}}$, sine-modulated ${}^1\text{H}^{\text{C}}$, and ${}^1\text{H}^{1\text{C}}$. Nevertheless, there are at least two ways of doing so; these two modified modules are labelled seHSQC1 and seHSQC2 respectively.

The seHSQC1 module (fig. 4.9b) was developed by me.* It retains the same general structure of the CRK seHSQC up until t_1 , but immediately after t_1 a composite ${}^1\text{H}$ pulse is used to effect the transformations

$$I_z \rightarrow I_y; \quad I_y \rightarrow I_x, \quad (4.10)$$

where the former is required for the ${}^1\text{H}^{\text{C}}$ pool and the latter for ${}^1\text{H}^{1\text{C}}$. This has the effect of storing the sine-modulated term as $2I_z S_z$ magnetisation for one spin echo (as opposed to the CRK seHSQC, which stores the cosine-modulated term as I_z). At the beginning of the final spin echo, this is transformed into antiphase magnetisation of the form $2I_x S_z$; thus, this spin echo must be lengthened to a total duration of 2Δ in order to fully refocus ${}^1J_{\text{CH}}$. The final modification involves the addition of an extra gradient immediately before t_1 : this ensures that the bulk ${}^1\text{H}^{1\text{C}}$ magnetisation is not dephased, just like in the NOAH HSQC module (fig. 4.3a).

The seHSQC2 module (fig. 4.9c), on the other hand, was first developed by Hansen et al.⁵⁷ In this pulse sequence, the initial ${}^1\text{H}$ 90° excitation pulse is replaced with a *zz* isotope-selective pulse

*Through a great deal of trial and error, and certainly *not* intelligent design.

(ZIP) element, which is very similar to the zz -filter but has different pulse phases. The effect of this on the $^1\text{H}^{13}\text{C}$ magnetisation pool amounts to a 90°_x pulse, and thus the same signals are ultimately detected from this magnetisation pool (save for a trivial 180° phase shift). However, on $^1\text{H}^{13}\text{C}$ magnetisation, the ZIP element acts as a 90°_y pulse; it turns out that this modification alone is sufficient to return the bulk $^1\text{H}^{13}\text{C}$ magnetisation to $+z$ at the end of the sequence. The ZIP element therefore represents an *isotope-specific rotation* on protons, where the rotation axis depends on whether the proton is directly coupled to ^{13}C or not.

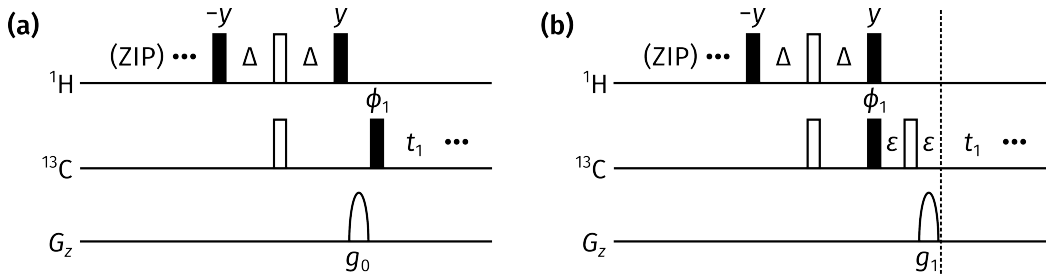


Figure 4.10: Comparison of CTP gradient schemes in seHSQC2 module. (a) As reported in Hansen et al.⁵⁷ g_0 is a purge gradient with arbitrary amplitude; the amplitude of the final CTP gradient must be halved. (b) The version used in this work (corresponding to fig. 4.9c).

In my work, I made one change to the experiment, namely the addition of an extra gradient echo prior to t_1 (thus leading to a symmetric gradient scheme similar to that in seHSQC1). Instead of this, the original paper had in fact inserted a purge gradient between the ^1H and ^{13}C 90° pulses just after the INEPT spin echo (fig. 4.10). The scheme I used leads to similar results, but has one advantage in that it allows the amplitude of the final CTP gradient (g_2 in fig. 4.9) to be twice as large: this is particularly relevant to ^{15}N experiments, as will be discussed in § 4.3.2.

The primary consideration when evaluating the different seHSQC versions in fig. 4.9 is naturally sensitivity. Figure 4.11 compares the sensitivities of the various possible NOAH-2 S^+C^c supersequence, against a NOAH-2 SC^c experiment. As can be seen, all three seHSQC implementations yield improvements in the sensitivities of CH peaks: the original CRK seHSQC has the largest effect here, followed by seHSQC2 and seHSQC1. For CH_2 and CH_3 peaks, the CRK seHSQC and seHSQC2 perform better than the standard HSQC, but the relative sensitivities in seHSQC1 have dipped slightly below 1. This is not entirely surprising in light of the discussion in § 4.1.1: the modified seHSQC1 and seHSQC2 sequences are expected to have lower sensitivities than the original experiment from which they are derived. In this case, it appears that the modifications made to the seHSQC1 sequence cause sensitivity losses which outweigh the benefits of using a sensitivity-enhanced experiment.

In the context of a NOAH supersequence, it is also important to consider how well each module preserves $^1\text{H}^{13}\text{C}$ magnetisation for subsequent homonuclear module(s). In fig. 4.11, the CLIP-

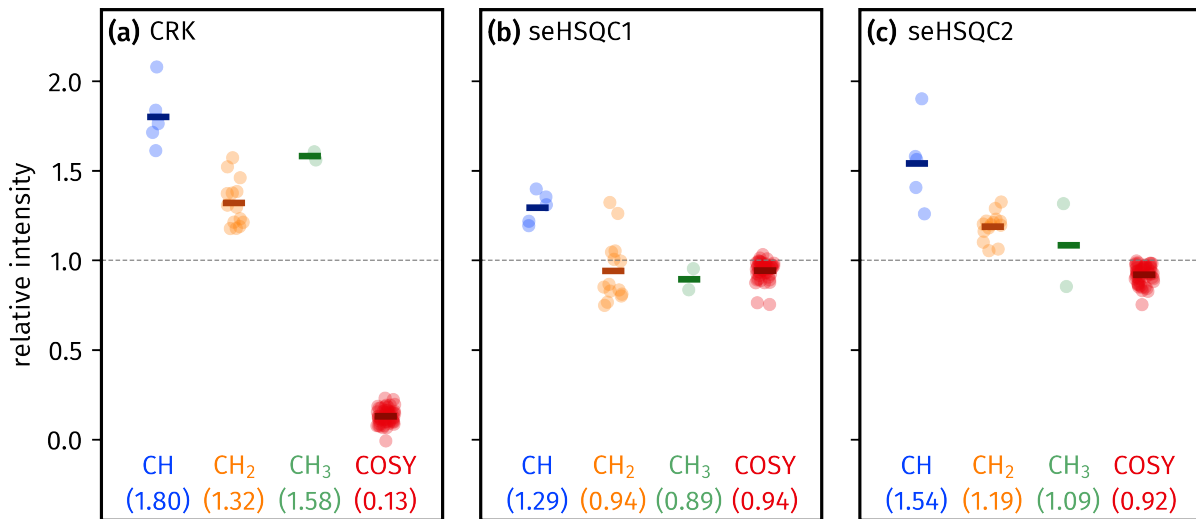


Figure 4.11: Sensitivities of seHSQC and CLIP-COSY modules in the NOAH-2 S^+C^c supersequences (run using $\Delta' = 1/(8 \cdot {}^1J_{CH})$). Intensities are reported relative to the corresponding peaks in a NOAH-2 SC^c supersequence; HSQC peaks are further broken down by their multiplicity. Each dot represents one peak; the solid lines as well as the numbers at the bottom of each plot represent averages over all peaks. (a) Using the CRK seHSQC. (b) Using the seHSQC1 module. (c) Using the seHSQC2 module. Data code: 7A-201115.

COSY module is used as the ‘reporter’ module, but the conclusions drawn here are applicable to *any* homonuclear module (or the HMBC). The CRK seHSQC does poorly in this respect, as it dephases the bulk magnetisation: this leads to almost an order-of-magnitude sensitivity loss in the CLIP-COSY. Both the other NOAH seHSQC modules, however, perform almost identically to the original HSQC module, as expected from the product operator analysis in fig. 4.9.

From fig. 4.11, we can draw some very simple conclusions: if the seHSQC is to be used at (or near) the *beginning* of a NOAH supersequence, where it needs to preserve ${}^1H^{1C}$ magnetisation, then the seHSQC2 module is the best choice. On the other hand, if the seHSQC module is placed at the *end* (say in a NOAH-2 BS⁺ supersequence), then the CRK seHSQC is optimal. This logic is used by GENESIS to choose the best seHSQC implementation based on other modules in the requested supersequence.

COSY-type artefacts

One (minor) way in which the seHSQC1 module outperforms seHSQC2 is in the suppression of COSY-type artefacts in the seHSQC spectrum.⁵⁸ These arise due to evolution of ${}^nJ_{HH}$ during the second-last spin echo of the CRK seHSQC. Specifically, if we assume the presence of a separate 1H spin K which is coupled to spin I , the sine-modulated $2I_yS_z$ term can evolve under both ${}^1J_{IS}$ and ${}^nJ_{IK}$ into $2I_yK_z$, and the last 1H 90° pulse transfers this coherence to the spin K . This leads to peaks correlating spin S with spin K . Since the seHSQC2 module shares the same product

operator analysis as the CRK seHSQC, these artefacts are also visible in seHSQC2 spectra.

However, for the seHSQC1 module, the corresponding term giving rise to these artefacts would be the cosine-modulated I_x term (at the beginning of the second-last spin echo). During this spin echo, this can again evolve under both J-couplings into $4I_yK_zS_z$, and the final 90° pulse would transform this to $-4I_zK_yS_z$. However, crucially, this term is antiphase with respect to the heteronucleus S: therefore, when decoupling is applied during acquisition, this term should not be observed. This analysis can be verified experimentally by inspection of the seHSQC spectra thus obtained. The COSY-type artefacts, labelled with red boxes in the seHSQC2 spectrum (fig. 4.12b), are largely absent in the seHSQC1 spectrum (fig. 4.12a). Some artefacts still remain, which perhaps arise due to pulse imperfections (or perhaps more complicated spin systems than the three-spin system considered here; I did not analyse this in any further detail).

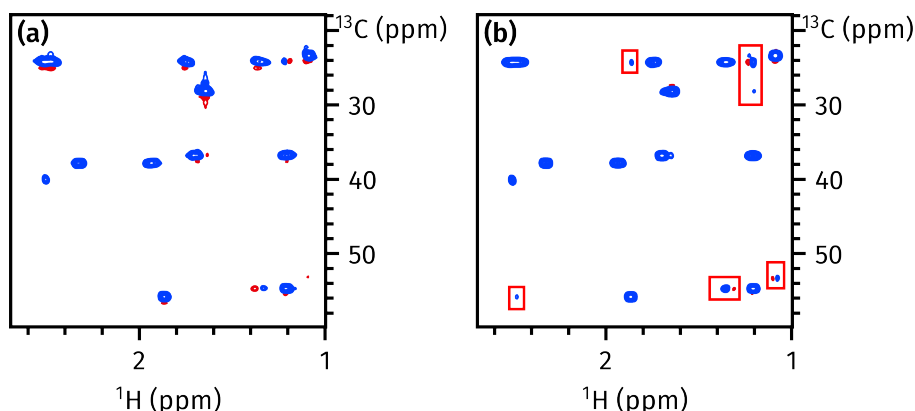


Figure 4.12: Comparison of COSY-type artefacts in NOAH seHSQC modules. (a) seHSQC1. (b) seHSQC2. The artefacts are highlighted in red boxes. Data code: 7A-201115.

Wing artefacts

One point worth considering is whether the extra gradient which I placed in the seHSQC2 module (fig. 4.10b) is truly needed. In the HSQC and seHSQC1 modules, the bulk magnetisation is placed along the transverse axis during t_1 ; therefore, if a pair of symmetric gradients are not used, this magnetisation will be dephased (as in the CRK seHSQC). However, in the seHSQC2 module, the bulk magnetisation is longitudinal, meaning that this gradient is not actually required for $^1\text{H}^{13}\text{C}$ preservation.

This gradient can indeed be removed in the seHSQC2 module without compromising the preservation of $^1\text{H}^{13}\text{C}$ magnetisation. However, the omission of this gradient leads to artefacts in the CLIP-COSY spectrum, which I term ‘wing artefacts’ because of the way they flank peaks in the CLIP-COSY (fig. 4.13). These artefacts can most clearly be seen in the diagonal peaks of methyl groups, but are present across the entire spectrum.

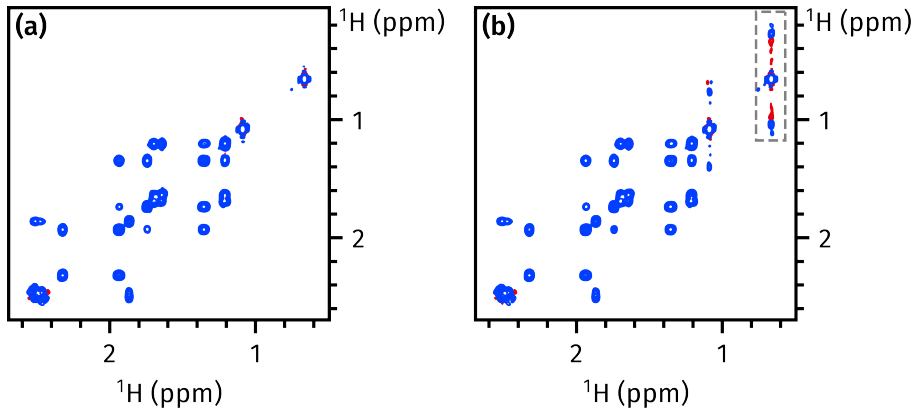


Figure 4.13: Wing artefacts in CLIP-COSY spectra, taken from NOAH-2 S^+C^c supersequences. (a) Using the seHSQC2 module as shown in fig. 4.9c. (b) Using a modified seHSQC2 module where the gradient echo just before t_1 is removed. The wing artefacts flanking the two rightmost diagonal peaks can be clearly seen; the boxed artefacts are studied further in fig. 4.14. Data code: 7A-201115.

A clue as to their origin is provided by the characteristic frequencies of the artefacts:

$$\left(\Omega_1, \Omega_2 = \Omega_I \pm \frac{\Omega_I \cdot [t_{1,\text{HSQC}}/2]}{t_{1,\text{COSY}}}, \Omega_I \right) \quad (4.11)$$

where $t_{1,X}$ refers to the value of t_1 in experiment X (it does not matter which increment is used, because the ratio of t_1 in the fraction above is constant). In accordance with this, when the seHSQC indirect-dimension spectral width is reduced, the artefacts are displaced outwards (fig. 4.14e). This suggests that these artefacts arise from $^1\text{H}^{13}\text{C}$ magnetisation which is frequency-labelled in two different t_1 periods: specifically, it evolves once during *half* of the seHSQC t_1 period, and then again during the CLIP-COSY t_1 period. In fact, it is necessary to apply CTP gradients on both sides of t_1 in order to suppress evolution of this stray magnetisation during both halves: if either or both of the gradients are removed, the artefacts are present (figs. 4.14a to 4.14d). Alternatively, the placement of a purge gradient before the first ^{13}C 90° pulse accomplishes a similar result, as in the original seHSQC2 implementation (fig. 4.10a).

These wing artefacts are unique to fast-pulsing experiments such as NOAH supersequences: in a typical 2D experiment, these are less likely to arise because of the long(er) recovery delays between data acquisition periods. In the sections which follow, we will see these wing artefacts surface repeatedly. Fortunately, it proves to be relatively easy to suppress them through the judicious use of gradients.

Multiplicity editing

Multiplicity editing is reasonably easy to include in all of the seHSQC sequences above. It suffices to add a spin echo of total duration $4\Delta = 1/J_{\text{CH}}$ just after t_1 , while also making sure to change

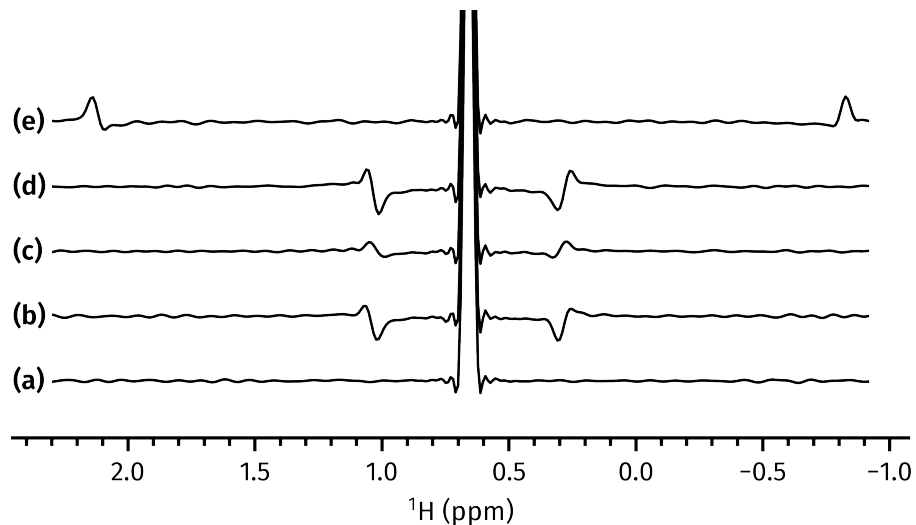
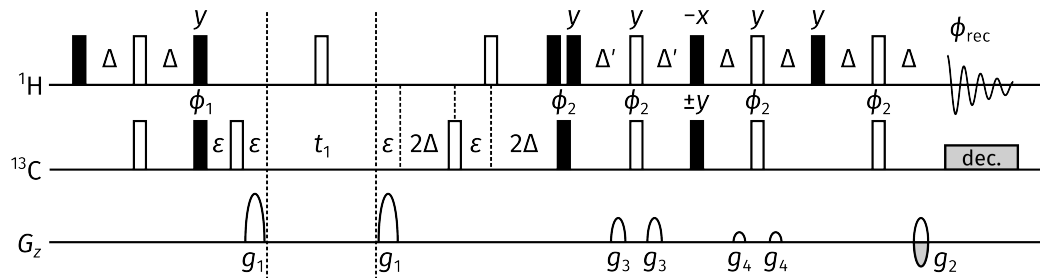


Figure 4.14: A closer look at wing artefacts in CLIP-COSY spectra, taken from NOAH-2 S⁺C^c supersequences with various modifications made to the seHSQC2 module. The spectra being plotted are 1D slices through $F_2 = 0.67$ ppm of the 2D CLIP-COSY spectrum; this corresponds to the boxed region in fig. 4.13b. (a) Using the default seHSQC2 module, with one gradient before and one after t_1 . (b) The same as (a), but with the gradient before t_1 turned off (the spin echo is still present, but the gradient intensity is set to 0). (c) The same as (a), but with the gradient after t_1 turned off. (d) The same as (a), but with both gradients turned off. (e) The same as (b), but with the seHSQC spectral width reduced by a factor of 4. *Data code: 7A-201115.*

some pulse phases to account for the extra ^1H 180° pulse (fig. 4.15).

(a) NOAH seHSQC1, edited



(b) NOAH seHSQC2, edited

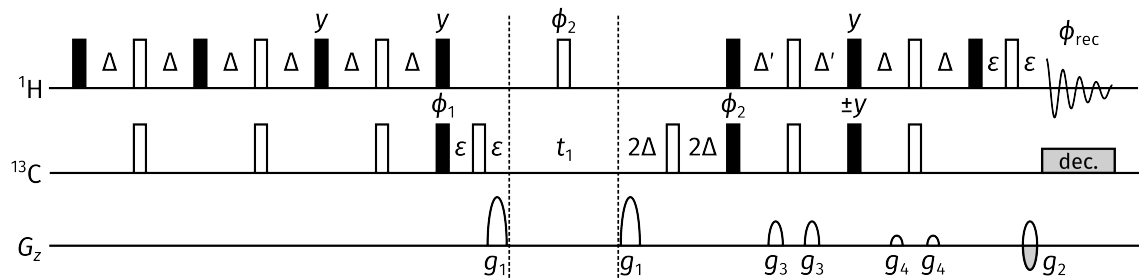


Figure 4.15: Multiplicity-edited NOAH seHSQC modules. (a) Edited seHSQC1. (b) Edited seHSQC2.

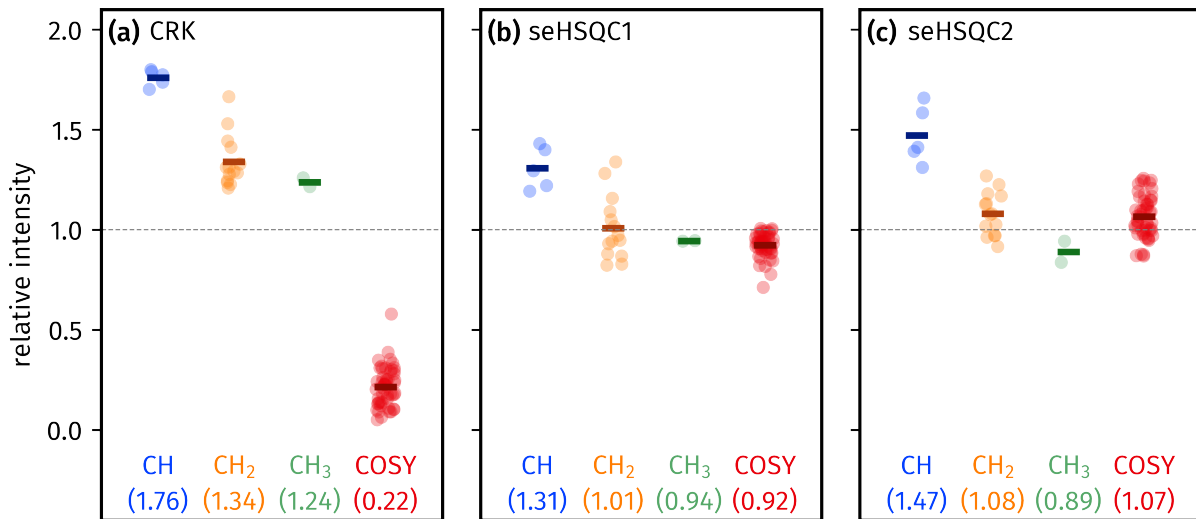


Figure 4.16: Sensitivity comparisons for multiplicity-edited seHSQC in NOAH-2 S^+C^c supersequences. The delay Δ' was set to $1/(8 \cdot {}^1J_{CH})$. (a) Using the edited CRK seHSQC. (b) Using the edited seHSQC1 module. (c) Using the edited seHSQC2 module. *Data code: 7A-201115.*

The inclusion of editing does not make a substantial difference in the sensitivity comparisons (fig. 4.16). However, it is interesting to note that the edited seHSQC2 in fact performs *better* than the edited HSQC in terms of preserving bulk magnetisation, as evidenced by the COSY intensities in fig. 4.16c which are greater than unity. This can be explained by the fact that in the editing period of the HSQC experiment (and seHSQC1), the bulk magnetisation is placed in the transverse plane. The evolution of homonuclear couplings will thus lead to a small loss in the amount of ${}^1H^{}C$ magnetisation preserved. In the seHSQC2 sequence, the bulk magnetisation is longitudinal during the editing period, so does not evolve under J_{HH} .

Choice of Δ'

As discussed previously, there are several possible values for the delay Δ' . I also specifically investigated the possibility of setting $\Delta' = 1/(4 \cdot {}^1J_{CH})$; the corresponding sensitivity comparisons are shown in fig. 4.17.

As can be seen, the sensitivity boosts obtained for CH groups are higher than in the corresponding spectra with $\Delta' = 1/(8 \cdot {}^1J_{CH})$ (fig. 4.11). However, there is also a small sensitivity *loss* for CH₂ and CH₃ groups as compared to the original HSQC: this is consistent with previous studies.⁵⁶ For CH₃ groups, which typically have large intensities in HSQC-type spectra, this is unlikely to be of any consequence; however, for diastereotopic CH₂ groups this sensitivity loss may not be desirable.

It is interesting to note that the performance of the seHSQC1 module is much closer to that of the seHSQC2 here. The fact that seHSQC1 performs worse with the reduced value of $\Delta' = 1/(8 \cdot {}^1J_{CH})$

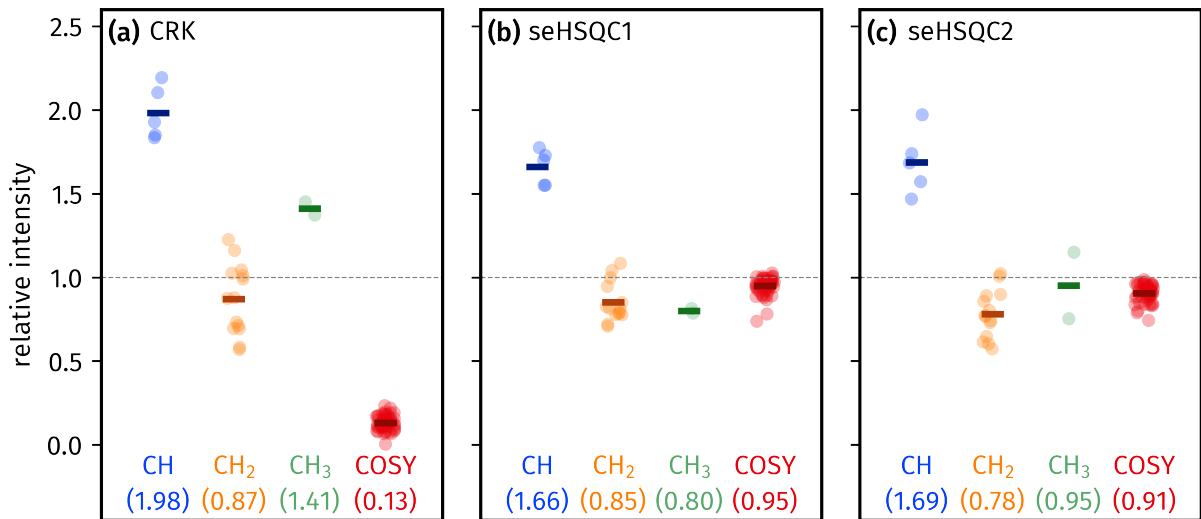


Figure 4.17: Sensitivity comparisons with Δ' set to $1/(4 \cdot {}^1J_{\text{CH}})$. Multiplicity editing was not used. (a) Using the CRK seHSQC. (b) Using the seHSQC1 module. (c) Using the seHSQC2 module. Data code: 7A-201115.

suggests that there are some inefficiencies in this section of the pulse sequence; however, there is no immediate explanation for this in the product operators (the relevant terms in the seHSQC1 are entirely similar to that in the CRK, save for a minus sign).

The conclusions drawn are entirely similar when multiplicity editing is enabled (fig. 4.18), so will not be further discussed.

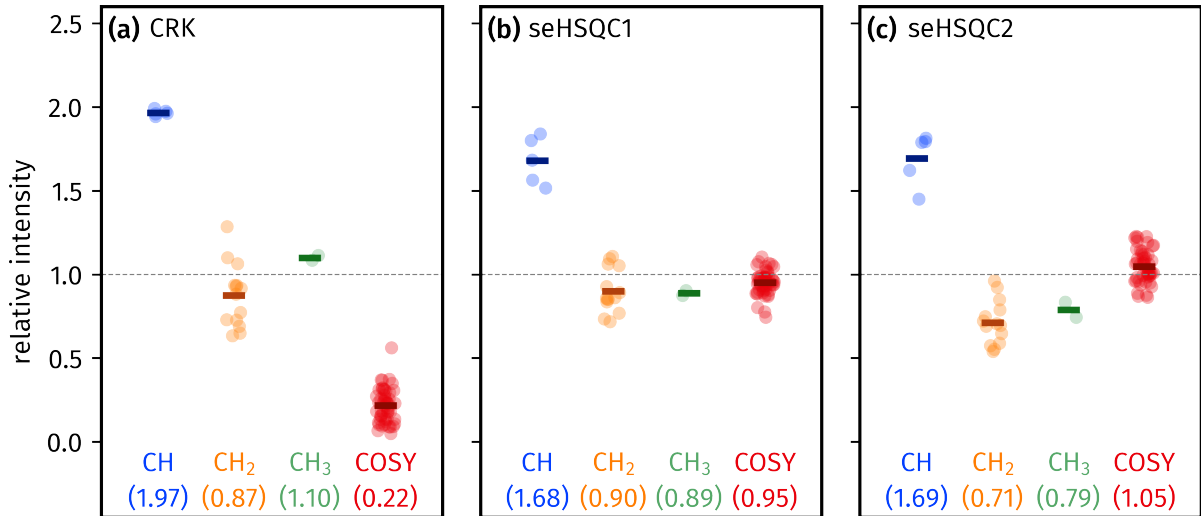


Figure 4.18: Sensitivity comparisons with Δ' set to $1/(4 \cdot {}^1J_{\text{CH}})$ and with multiplicity editing. (a) Using the edited CRK seHSQC. (b) Using the edited seHSQC1 module. (c) Using the edited seHSQC2 module. Data code: 7A-201115.

Optimal control for seHSQC1

One unresolved question is why the seHSQC1 module has a lower sensitivity than the seHSQC2, despite being shorter and containing fewer 180° pulses. There is also nothing in the product operator analysis to explain why this should be the case. One remaining possibility is the presence of some non-ideality in the pulses themselves, and the composite ^1H pulse is an obvious candidate for investigation.*

In order to investigate the extent to which this was responsible, I sought to use optimal control theory to develop a *universal rotation pulse* (URP) which could replace this composite pulse. Such a pulse must accomplish the transformations shown in eq. (4.10), namely $I_z \rightarrow I_y$ and $I_y \rightarrow I_x$.[†] This optimisation was performed using an interior-point algorithm (the default in Matlab `fmincon`) and the ESCALADE method⁶⁰ for the calculation of analytic derivatives. Pulse fidelity was calculated as an average over 51 ^1H spins, evenly distributed across a frequency range of 11 kHz (corresponding to 15.7 ppm at 700 MHz); the duration of the pulse was fixed at 200 μs , and 200 pulse points were used. The maximum B_1 allowed was 15 kHz; the optimisation was run with a 10% variation in B_1 to minimise the effects of inhomogeneity.

The ^{13}C 90° pulse which is simultaneously applied during the pulse sequence was not optimised together with this. Since the ^1H URP has a longer duration than this, the sequence must be modified slightly (fig. 4.19). In particular, the delays α and β must be chosen in order to satisfy the relations $2\varepsilon = \alpha + \beta$ (to ensure $^1\text{H}^{13}\text{C}$ magnetisation is refocused), and $\alpha = \beta + \tau$ (to ensure that for $^1\text{H}^{13}\text{C}$ magnetisation, the ^{13}C chemical shift evolves for a total duration of t_1).

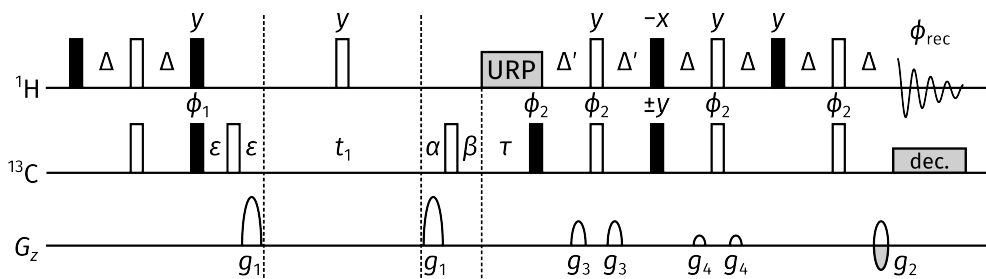


Figure 4.19: seHSQC1 pulse sequence using a ^1H URP in place of the double- 90° composite pulse. τ is the difference in duration between the URP and the ^{13}C 90° hard pulse. Delays are set as: $\alpha = \varepsilon + \tau/2$; $\beta = \varepsilon - \tau/2$.

This optimisation process yielded a pulse with 99.99% fidelity: the (theoretical) performance of

*The concept of a *composite pulse*⁵⁹ is usually associated with greater efficiency / uniformity, but here I have used the term to loosely refer to a consecutive series of pulses.

[†]The effect of this pulse on x -magnetisation is not important for the seHSQC1, but is in fact fully determined by these two constraints: $UI_xU^\dagger = -iU(I_yI_z - I_zI_y)U^\dagger = -i(UI_yU^\dagger UI_zU^\dagger - UI_zU^\dagger UI_yU^\dagger) = -i(I_xI_y - I_yI_x) = I_z$. This reveals a geometric interpretation of this pulse element: it is actually a 120° rotation about the vector $(1, 1, 1)$, which interchanges the three Cartesian axes; it is closely related to the C_3 symmetry operation in the octahedral point group.

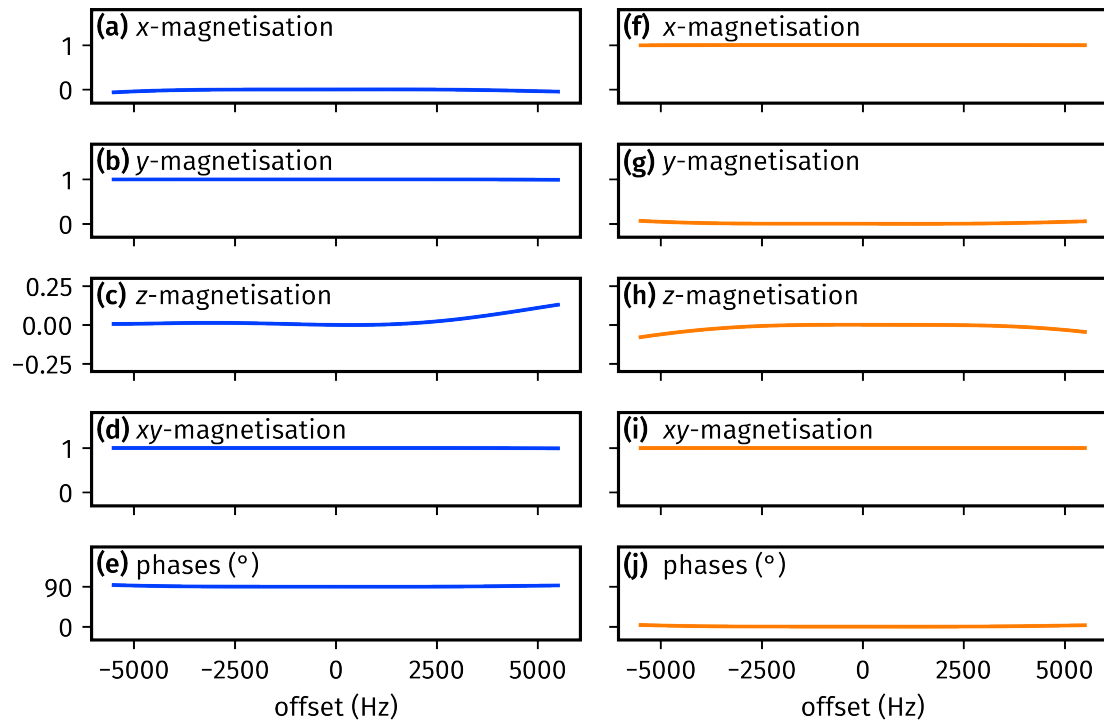


Figure 4.20: Simulated performance of URP used for seHSQC1 module on z - and y -magnetisation. The pulse fidelity was 99.99%. (a)–(e) Using z -magnetisation as input: the plots respectively show the amount of x -magnetisation, y -magnetisation, z -magnetisation, transverse magnetisation ($M_{xy} = \sqrt{M_x^2 + M_y^2}$), and the phase of the transverse magnetisation generated, as a function of offset frequency. (f)–(j) The same, but using y -magnetisation as input.

this pulse on z - and y -magnetisation is shown in fig. 4.20. However, when tested in the actual seHSQC1 experiment, this failed to yield any substantial difference compared to the original double 90° pulse (figs. 4.21a and 4.21b). Importantly, the performance still falls below that of the seHSQC2 module (fig. 4.21c). The reason for the poorer sensitivity therefore likely lies elsewhere.

BIG-BIRD versus ZIP

The final point in this section pertains to the implementation of the seHSQC2 module. As it stands, the isotope-specific rotation element placed at the start of the module is the ZIP element: its role is to effect 90° rotations with different phases on the ${}^1\text{H}^{\text{C}}$ and ${}^1\text{H}^{!C}$ magnetisation pools. However, such pulse sequence elements have been known for a long time: these include TANGO,⁶¹ BANGO,⁶² BIRD,^{63–65} BIG-BIRD,⁶⁶ and TIG-BIRD.⁶⁷ In particular, the BIG-BIRD element, which allows for independent excitation of ${}^1\text{H}^{\text{C}}$ and ${}^1\text{H}^{!C}$ magnetisation with arbitrary flip angles and phases, can be designed to accomplish the same overall effect as the ZIP element.

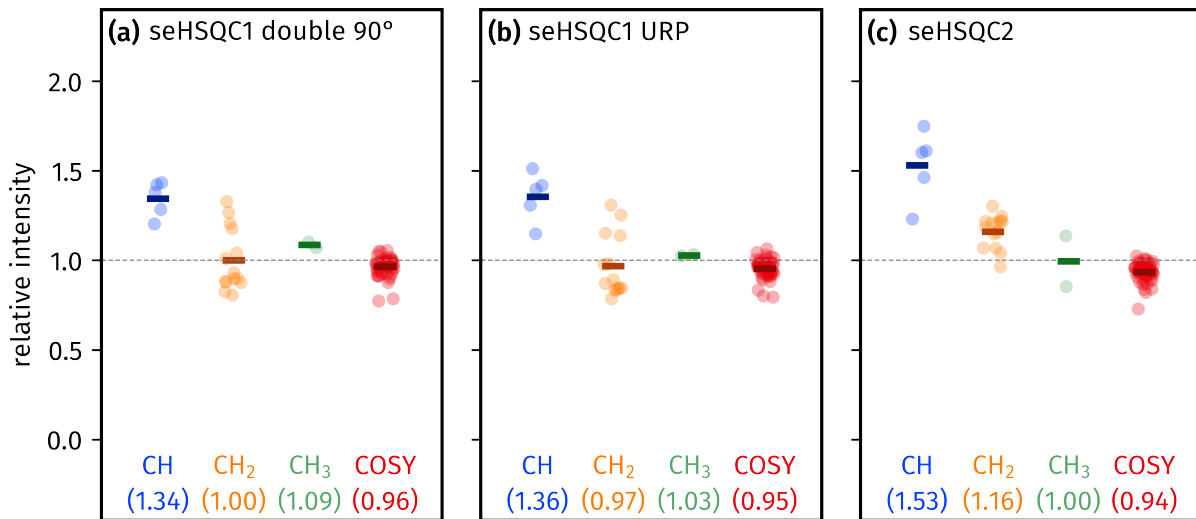


Figure 4.21: Sensitivity comparisons using the seHSQC1 URP. Δ' was set to $1/(8 \cdot ^1J_{\text{CH}})$; no multiplicity editing was used. (a) Original seHSQC1 module with double ^1H 90° pulse. (Note that different datasets were used for this figure and fig. 4.11, so the numbers are very slightly different.) (b) seHSQC1 using the optimised URP shown in fig. 4.20. (c) seHSQC2 module for comparison. *Data code:* 7A-220110.

For the seHSQC2 without multiplicity editing, the relevant BIG-BIRD sequence is

$$45^\circ_{45^\circ}({}^1\text{H}) - 2\Delta - 180^\circ({}^1\text{H}, {}^{13}\text{C}) - 2\Delta - 45^\circ_{225^\circ}({}^1\text{H}), \quad (4.12)$$

where β_ϕ represents a hard pulse with flip angle β and phase ϕ . For the edited seHSQC2, the phases must be altered slightly:

$$45^\circ_{315^\circ}({}^1\text{H}) - 2\Delta - 180^\circ({}^1\text{H}, {}^{13}\text{C}) - 2\Delta - 45^\circ_{135^\circ}({}^1\text{H}). \quad (4.13)$$

Figure 4.22 provides sensitivity comparisons between the BIG-BIRD and ZIP versions of the seHSQC2. The fact that similar numbers were achieved with both elements indicates that the BIG-BIRD element is (largely) effecting the desired rotations on the different magnetisation pools. However, in all respects, the performance of the BIG-BIRD sequence was not as good as the ZIP version: both the seHSQC sensitivity itself, as well as the sensitivity of the later CLIP-COSY, were slightly decreased when the BIG-BIRD element was used. This is true regardless of whether multiplicity editing was used.

4.3.2 ${}^{15}\text{N}$ HMQC

In § 4.3.3, I will discuss how the sensitivity-enhanced HSQC modules developed above may be adapted into ${}^1\text{H}-{}^{15}\text{N}$ experiments. However, before that, I make a slight detour to cover the ${}^{15}\text{N}$ HMQC experiment, which (up until my DPhil) was the experiment of choice for detecting one-bond ${}^1\text{H}-{}^{15}\text{N}$ correlations.

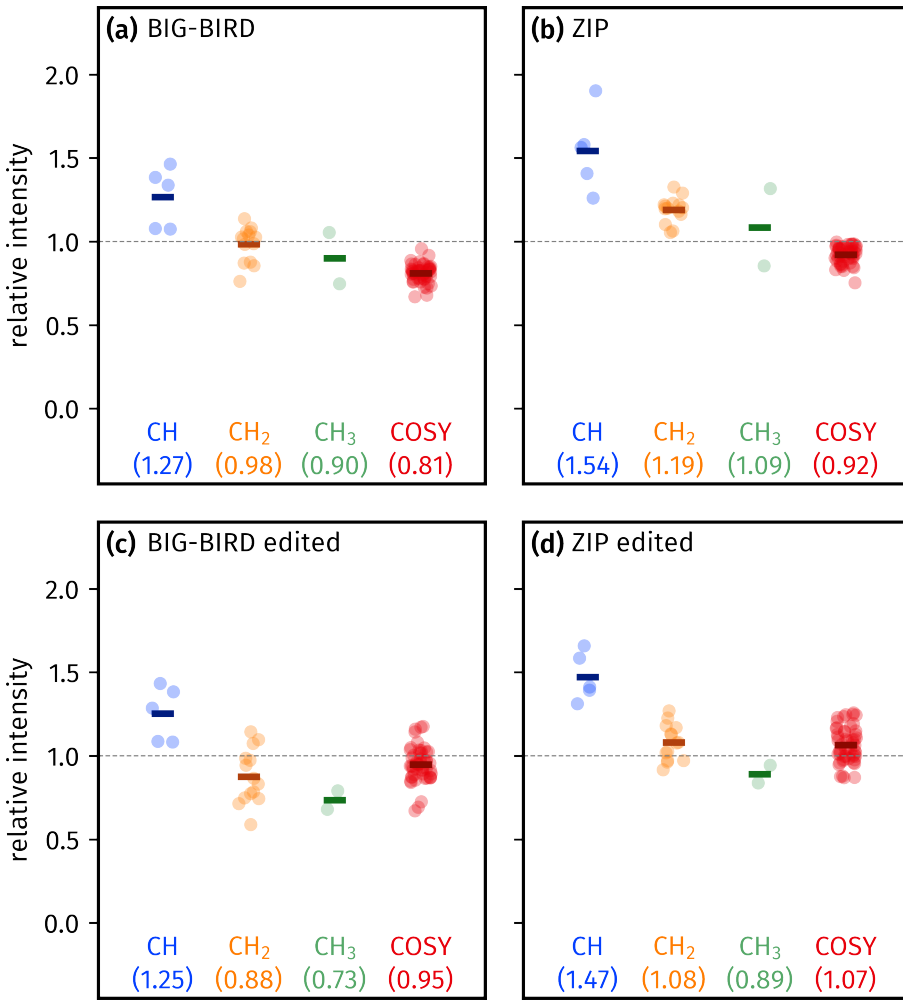


Figure 4.22: Sensitivity comparisons of BIG-BIRD and ZIP pulse elements in seHSQC2 module. The reference dataset being compared against is still the NOAH-2 SC^c (but with editing in (c) and (d)). The delay Δ' was set to $1/(8 \cdot {}^1J_{\text{CH}})$. (a) Unedited seHSQC2 using BIG-BIRD. (b) Unedited seHSQC2 using ZIP. (c) Edited seHSQC2 using BIG-BIRD. (d) Edited seHSQC2 using ZIP. Data code: 7A-201115.

CTP gradient scheme

The HMQC module is based on the ASAP-HMQC reported by Kupče and Freeman¹³. In the original report, the ASAP-HMQC used a symmetric gradient scheme similar to that in the seHSQC modules previously described (fig. 4.23a). However, in the NOAH HMQC module,³² this was expanded into a four-gradient scheme where bipolar gradient pairs were placed before and after t_1 (fig. 4.23b). This was likely implemented in order to allow the final gradient, g_2 , to have as large an amplitude as possible. In heteronuclear experiments, this final gradient is particularly important for dephasing bulk magnetisation which is transverse just prior to detection (due to pulse imperfections or relaxation). If this gradient is too weak, this unwanted magnetisation will be incompletely dephased, leading to artefacts in the resulting spectrum.

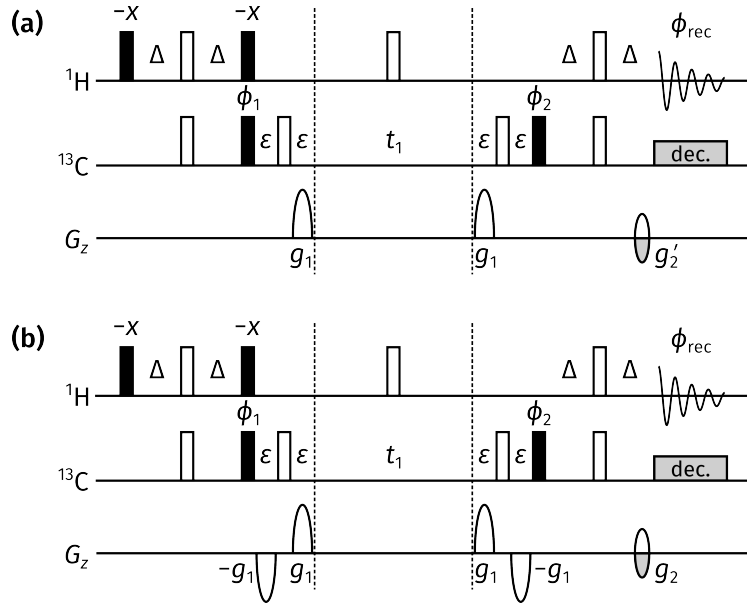


Figure 4.23: (a) With two encoding gradients around t_1 . (b) With four encoding gradients around t_1 . Phase cycling is performed with $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x)$, and $\phi_{\text{rec}} = (x, -x, -x, x)$. The delay Δ is set to $1/(4 \cdot ^1J_{\text{NH}})$. Gradient amplitudes are: $g_1 = 80\%$; $g_2 = \pm 32.4\%$; $g'_2 = g_2/2$.

Strategies to maximise the amplitude of this gradient are particularly crucial in ¹H–¹⁵N experiments (as compared to ¹H–¹³C experiments) for two reasons. Firstly, the natural abundance of ¹⁵N (0.36%) is even smaller than ¹³C (1.1%), which translates into a weaker signal: better artefact suppression must be achieved in order for the signal to not be obscured. Secondly, the gyromagnetic ratio of ¹⁵N is also smaller: since $g_2/g_1 \propto |\gamma_N/\gamma_H|$, g_2 will naturally be smaller if no modifications are made to the pulse sequence.

In this respect, the four-gradient scheme in fig. 4.23b is superior to the two-gradient scheme, because the gradient g_2 will have an amplitude of $4\gamma_N g_1/\gamma_H$. However, when used in a NOAH supersequence, this leads to wing artefacts in downstream modules, since bulk ¹H–¹⁵N magnetisation effectively does not experience any coherence order selection during t_1 . This motivates a return to the two-gradient scheme of fig. 4.23a, but to compensate for the fact that the decoding gradient g'_2 has half of the amplitude of g_2 , all CTP gradients were instead *lengthened* from their usual duration of 1 ms to 2.5 ms. This ensures that any stray transverse bulk magnetisation at the end of the HMQC module is effectively dephased by the final gradient.

The HMQC spectra thus obtained are shown in fig. 4.24. These, and all other ¹H–¹⁵N spectra in this chapter, were acquired assuming a $^1J_{\text{NH}}$ value of 90 Hz. The first column shows the spectra obtained with the original four-gradient scheme: although the artefacts at 2.2 ppm are reasonably well-suppressed in the HMQC module (fig. 4.24b), the CLIP-COSY spectrum clearly has a set of wing artefacts (fig. 4.24c). (Note that the wing artefacts occur at different frequencies compared

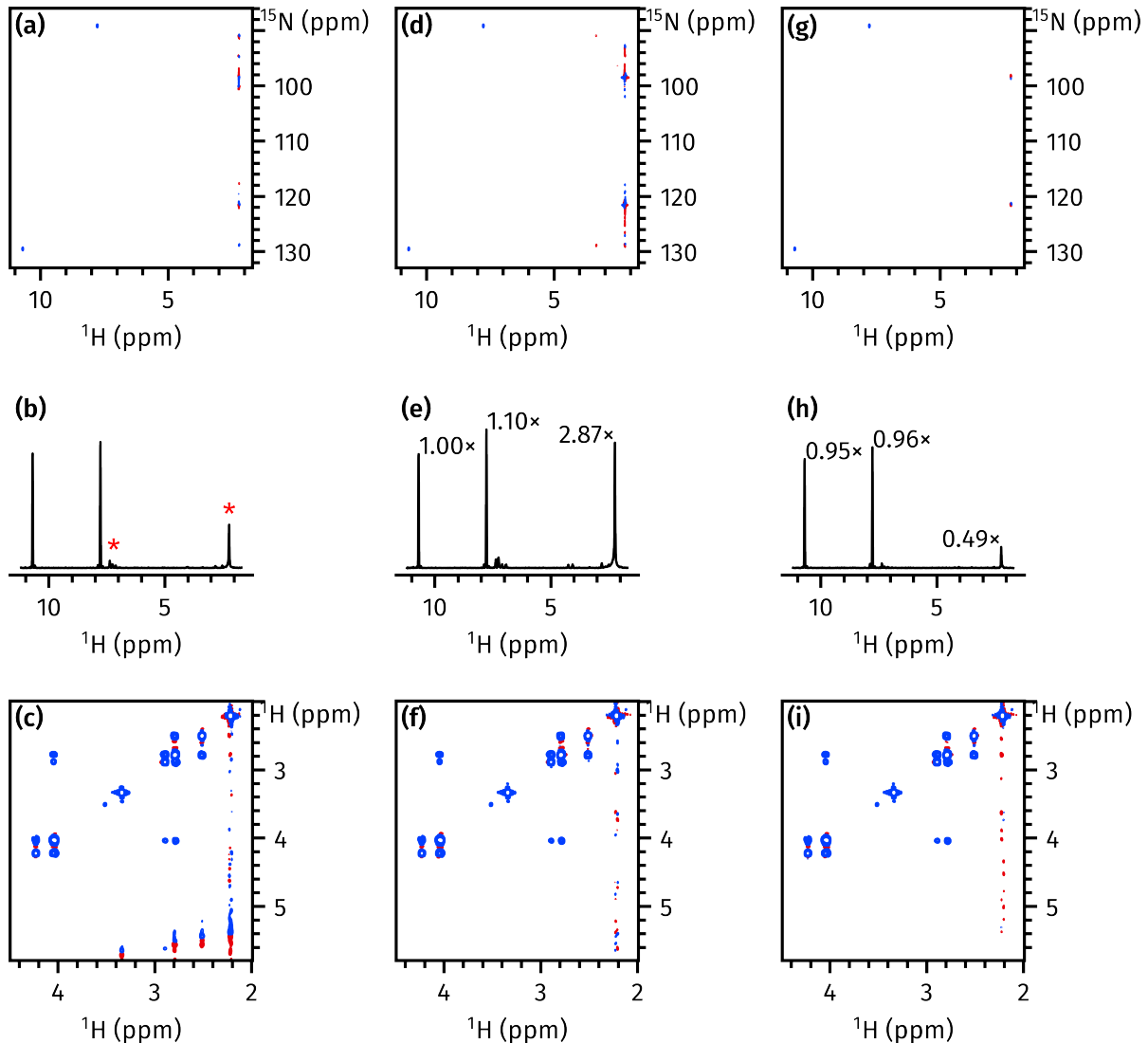


Figure 4.24: Comparison of HMQC and CLIP-COSY spectra obtained from NOAH-3 $M_N^+ S^+ C^$ supersequences, acquired using different HMQC gradient schemes. In the first row, the HMQC spectrum itself is shown. In the second row, the positive projection of the HMQC spectrum onto the F_2 axis is shown; the numbers indicate peak intensities with respect to the reference dataset (the left column). The asterisks indicate artefacts arising from bulk magnetisation which is not sufficiently dephased by the final gradient. In the third row, (an inset of) the CLIP-COSY spectrum is shown. (a)–(c) Using the four-gradient scheme of fig. 4.23b, with 1 ms gradients. (d)–(f) Using the two-gradient scheme of fig. 4.23a, with 1 ms gradients. (g)–(i) Using the two-gradient scheme of fig. 4.23a, with 2.5 ms gradients. Data code: 7Z-200926.

to the ^{13}C seHSQC case, because the t_1 increment in the ^{15}N HMQC module is different.)

The second column shows what happens when the two-gradient scheme is adopted without changing the gradient duration. Although the CLIP-COSY wing artefacts disappear (fig. 4.24f), the HMQC artefacts are over twice as intense (fig. 4.24e), and (in this case) have comparable intensity to the desired peaks.

By using the two-gradient scheme and increasing the gradient duration to 2.5 ms, we obtain the best of both worlds: the HMQC artefacts are well-suppressed (fig. 4.24h, in fact even better than in the original spectrum), and the CLIP-COSY is free of wing artefacts (fig. 4.24i). The only drawback is a slight loss in signal intensity in the HMQC, which arises due to diffusion and relaxation during the longer pulse sequence. However, this decrease is only on the order of 5% (for this example), which is a totally acceptable price to pay in return for the improved spectral quality.

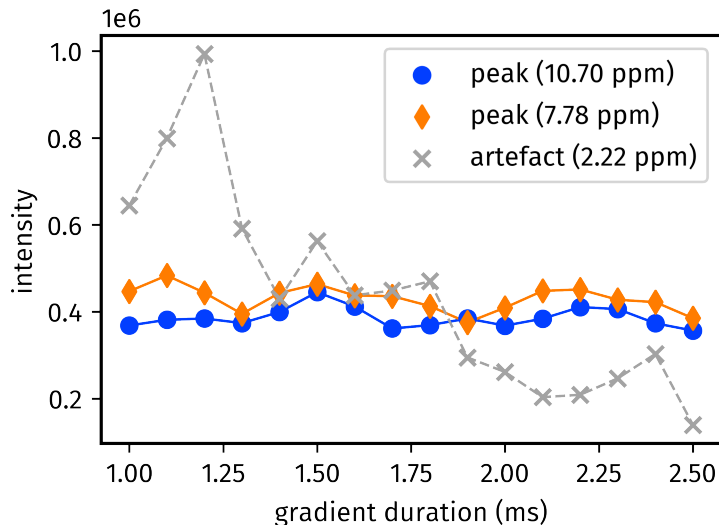


Figure 4.25: Variation of HMQC signal and artefact intensity with CTP gradient duration, as measured by (absolute) peak heights in the positive F_2 projection. Data code: 7Z-200926.

The signal loss does suggest that we should check whether such a long gradient is truly needed. I therefore ran the two-gradient HMQC experiment with a series of gradient durations from 1 ms to 2.5 ms; the signal and artefact intensities in each of these experiments are plotted in fig. 4.25. Generally, there is little variation in the intensities of the two desired peaks. The artefact intensity is more erratic, which possibly reflects the fact that gradient dephasing varies sinusoidally with the gradient duration τ due to the spatial integral

$$\frac{1}{L} \int_{-L/2}^{L/2} \exp(i\gamma G z \tau) dz = \frac{\sin(\gamma GL\tau/2)}{\gamma GL\tau/2}, \quad (4.14)$$

where L is the sample length and G the gradient amplitude (see also eq. (1.93)). Nevertheless,

increased gradient durations clearly lead to better artefact suppression, which (at least in these datasets) is greatest with 2.5 ms gradients. This value of 2.5 ms was therefore fixed on as a suitable choice for ^{15}N experiments, including the seHSQC which is discussed in § 4.3.3.

k- and SW-scaling

$^1\text{H}-^{15}\text{N}$ spectra of small molecules are often relatively sparse in the indirect dimension, and do not typically need to be acquired with the same resolution as a $^1\text{H}-^{13}\text{C}$ experiment. However, if these two different classes experiments were to be combined together within a supersequence, it would not ordinarily be possible to toggle the resolution of each module individually.

One way of changing this would be to decrease the frequency with which the $^{15}\text{N} t_1$ duration is incremented in the pulse programme. This practice was first proposed by Parella et al. in the context of time-shared NMR,^{27,68} and in this thesis is called ‘*k*-scaling’. This means that each t_1 increment in the ^{15}N experiment is acquired several times before t_1 is incremented, which effectively leads to k times fewer t_1 increments, but with each increment having k times the number of scans (after the data have been combined). An alternative to this would be to increase the spectral width of the ^{15}N module (which is encoded as the CNST40 parameter in GENESIS): this leads to very similar results, and is referred to as ‘SW-scaling’ here.

Figure 4.26 illustrates all of these possibilities in detail, focusing only on the t_1 sampling schedule of the scaled ^{15}N experiments. In a ‘typical’ NOAH supersequence (fig. 4.26a), 256 t_1 increments would be recorded (corresponding to the eight blue dots); for the gramicidin sample used here, the largest value of t_1 (denoted as AQ_1) is around 60 ms. The default TopSpin processing routine for 2D NMR applies forward linear prediction (LP)^{69–72} in the indirect dimension, such as to double the number of data points: the data obtained through LP are indicated by orange dots. This leads to an *effective* AQ_1 which is twice as large, denoted as $\text{AQ}_{1,\text{eff}}$. Finally, zero filling (ZF) is applied to increase the digital resolution to a given value, as represented by the green dots.

In the *k*-scaled spectrum, both AQ_1 (and thus $\text{AQ}_{1,\text{eff}}$) are halved, in return for a doubling of the number of scans (fig. 4.26b). The resolution in the indirect dimension is proportional to $\text{AQ}_{1,\text{eff}}$; thus, this also corresponds to a 2× decrease in resolution. However, this can be counteracted through the use of extra linear prediction (fig. 4.26c) to extend $\text{AQ}_{1,\text{eff}}$ back to its original value. The SW-scaled spectra (figs. 4.26d and 4.26e) are almost entirely equivalent to the *k*-scaled spectra in terms of their effect on AQ_1 and $\text{AQ}_{1,\text{eff}}$, except that the sampling schedule is *very* slightly different.

Figure 4.27 shows the effects of performing *k*- or SW-scaling on the ^{15}N HMQC module, without any extra linear prediction beyond the default. In this case, these scaling procedures alone can in

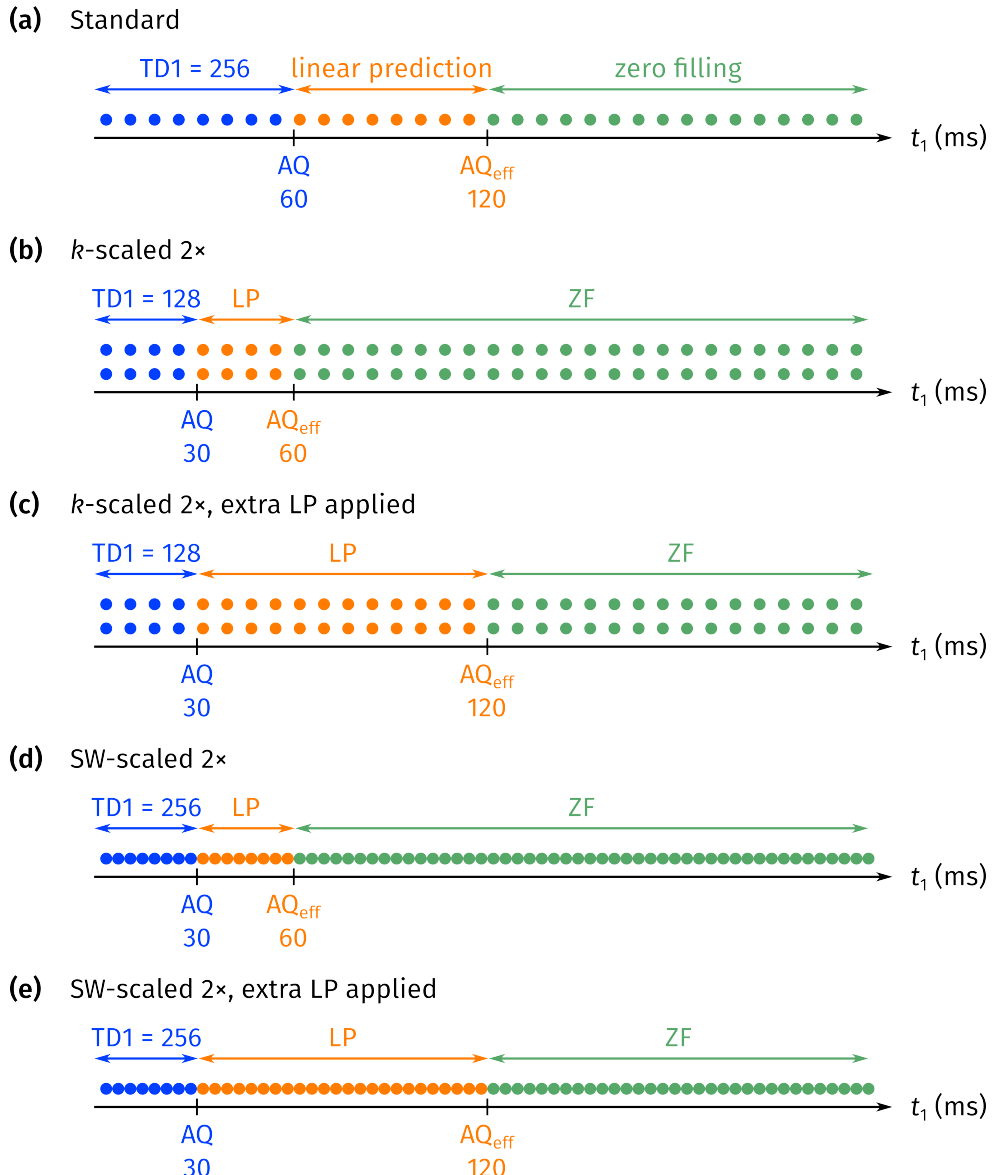


Figure 4.26: Pictorial representation of k - and SW-scaling in NOAH ^{15}N modules. Each blue dot represents $32 t_1$ increments physically acquired as part of the experiment; the experimental time is proportional to the number of blue dots, and is constant across all of these experiments. Orange dots represent $32 t_1$ increments obtained through forward linear prediction, and green dots represent zeroes. (a) The default experiment. (b)–(c) k -scaled experiment, without and with extra linear processing to restore the original value of $\text{AQ}_{1,\text{eff}}$. (d)–(e) SW-scaled experiment, without and with extra linear processing to restore the original value of $\text{AQ}_{1,\text{eff}}$.

fact lead to significant gains in SNR (as measured by peak heights), with up to $2.6\times$ improvements being observed at the extreme of 8-fold k - or SW-scaling. This likely arises because the J_{HH} modulation in the indirect dimension (visible in the standard spectrum, fig. 4.27b) is no longer being resolved: the largest gains are attained for the peak at 123 ppm, where this modulation is especially prominent.

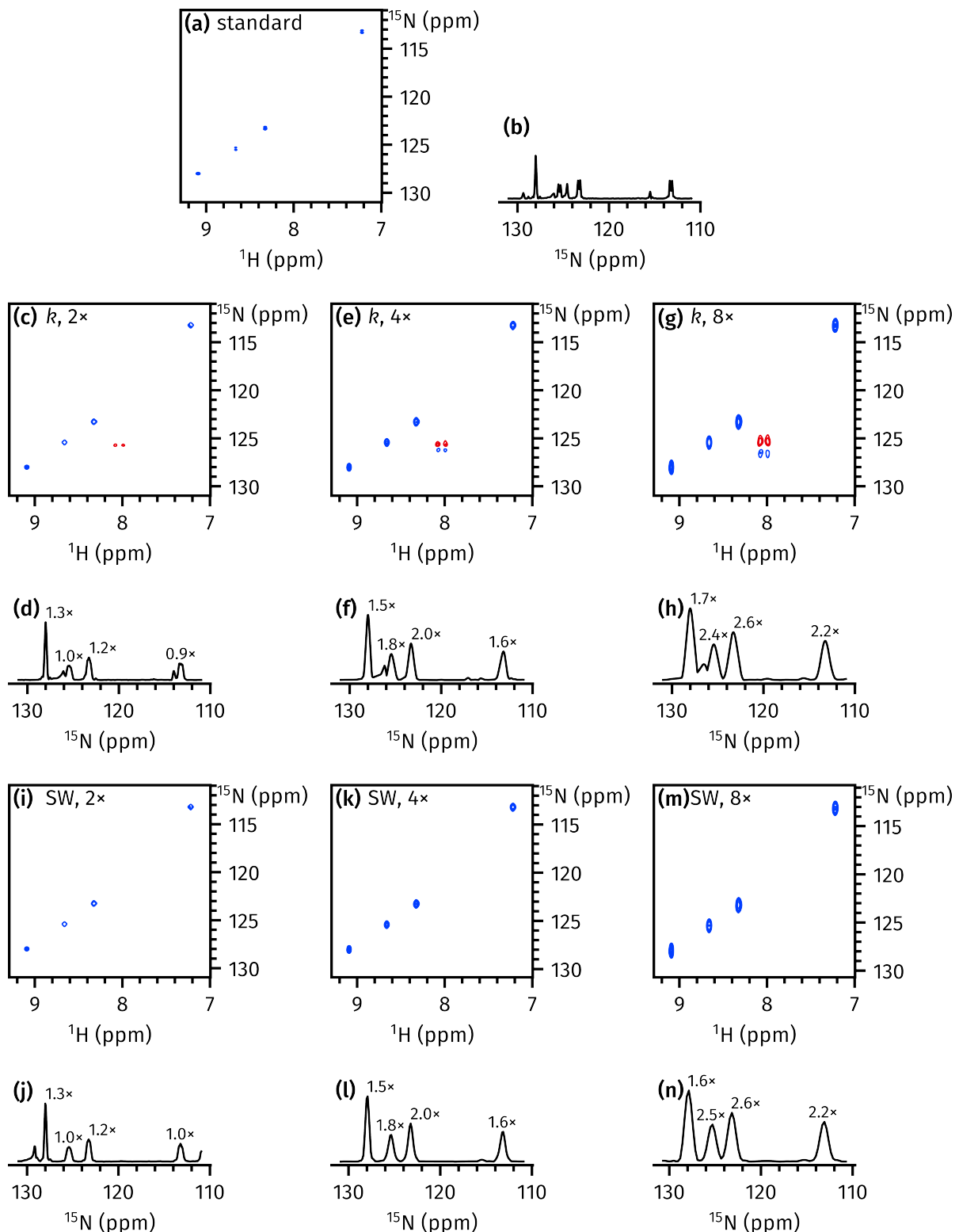


Figure 4.27: Effects of k - and SW-scaling on NOAH HMQC spectrum (taken from NOAH-3 $\text{M}_\text{N}\text{S}^+\text{C}^\text{c}$ supersequences). Each HMQC spectrum is shown together with a positive projection onto the F_1 axis. The relative SNR of each peak, with respect to the standard spectrum, is indicated on each of the other projections. (a)-(b) Standard spectrum. (c)-(h) k -scaled spectra. (i)-(n) SW-scaled spectra. Data code: 7G-210310.

The effect of scaling *and* adding extra linear prediction is shown in fig. 4.28. In general, this combination of scaling plus LP leads to improvements in spectral SNR of up to 6 \times . However, this is accompanied by distortions in the F_1 multiplet structure, especially for the k -scaled spectra (figs. 4.28c to 4.28h).

However, caution should be exercised when interpreting these results. Although this improvement in SNR is genuine, it is not necessarily the case that this represents a true improvement in *detection sensitivity*: in other words, processing techniques such as LP do not always allow for better discrimination between signal and noise.^{73,74*} Further tests would need to be done on more dilute samples[†] to ascertain the benefits of the scaling-plus-LP routine on spectra with lower intrinsic SNRs.

4.3.3 ^{15}N sensitivity-enhanced HSQC

The implementation of the ^{15}N seHSQC modules is entirely identical to the ^{13}C versions previously discussed (§ 4.3.1), except for the following points:

- CTP gradient amplitudes were set using $g_1 = 80\%$ and $g_2 = \mp 16.2\%$ (note the sign change because of the negative magnetogyric ratio of ^{15}N);
- CTP gradient durations were lengthened to 2.5 ms (as discussed for the ^{15}N HMQC) in order to suppress artefacts in the seHSQC itself;
- the delays Δ and Δ' were both set to $1/(4 \cdot ^1J_{\text{NH}})$ to maximise sensitivity for NH groups (as before, the value of $^1J_{\text{NH}}$ was set to 90 Hz);
- ^{13}C adiabatic pulses were replaced with hard pulses on ^{15}N .

Multiplicity editing was not implemented or used in the ^{15}N experiments; there is generally little need for this.

Sensitivity analysis

The seHSQC is expected to provide sensitivity gains over the HMQC for two reasons: first, the seHSQC peaks are not split by $^nJ_{\text{HH}}$ in the indirect dimension, and secondly, the PEP transfer element leads to (in theory) a doubling in sensitivity for NH groups. To quantify these gains, I ran several NOAH-3 XS $^+ \text{C}^c$ experiments, where the first module was a ^{15}N HSQC, seHSQC1, or seHSQC2 module (fig. 4.29). These were then compared against the basic NOAH-3 M $_{\text{N}}\text{S}^+ \text{C}^c$ supersequence. Including the ^{15}N HSQC module in this analysis allows us to determine how

*In recent years, this issue has been investigated more thoroughly in the context of NUS.⁷⁵

[†]Or, perhaps, less sensitive instruments—although the 700 MHz cryoprobe used here was the only option for these supersequences.

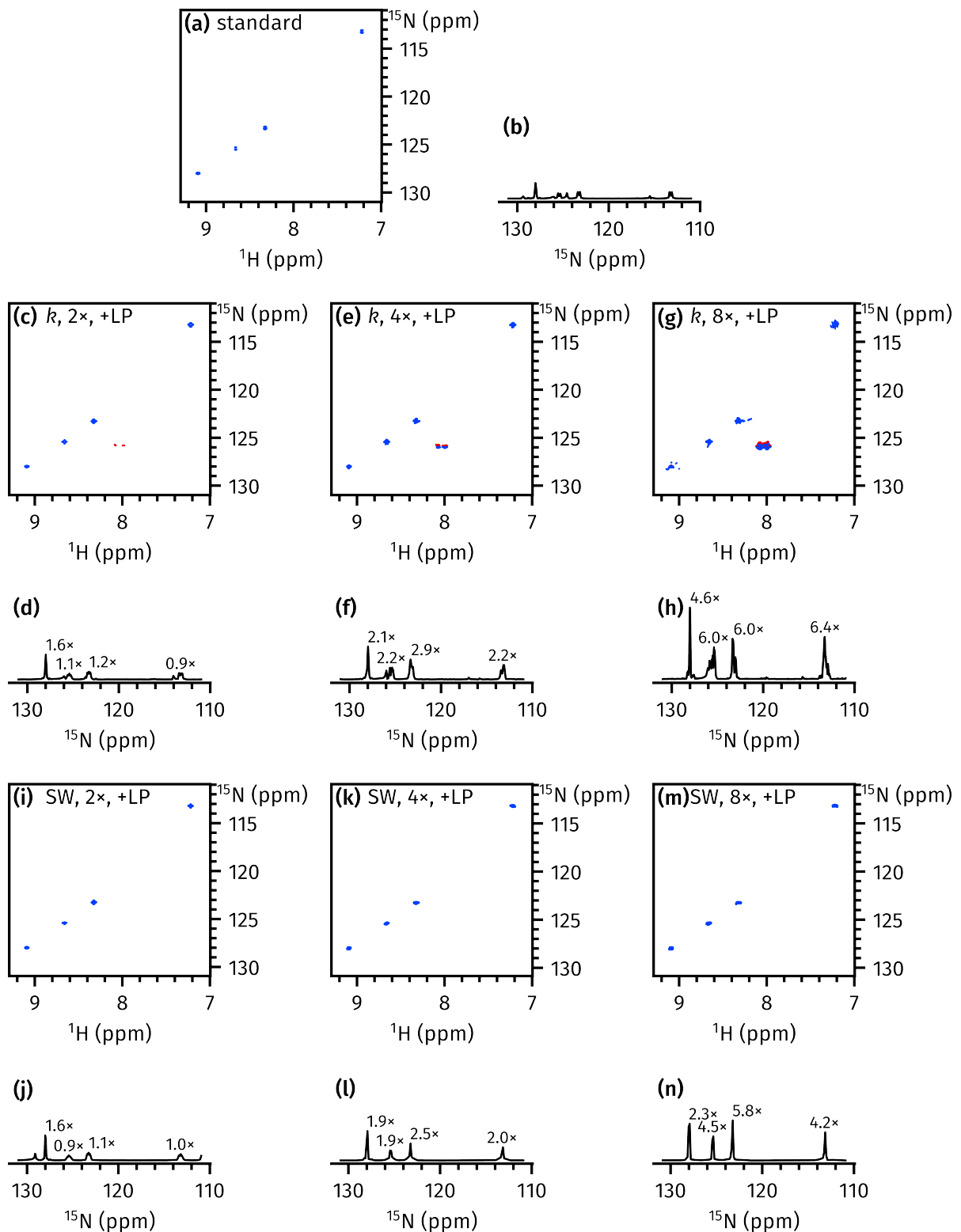


Figure 4.28: The same as in fig. 4.27, but with extra linear prediction applied to all scaled spectra to bring $AQ_{1,\text{eff}}$ up to its original value in the standard spectrum. Linear prediction of k times more points leads to a \sqrt{k} increase in noise; to account for this, all spectra are plotted with the same noise level. (a)–(b) Standard spectrum. (c)–(h) k -scaled spectra. (i)–(n) SW-scaled spectra. Data code: 7G-210310.

much of the sensitivity gain is due to multiplet collapse, and how much is due to the sensitivity enhancement block. Note that the peak at 8.0 ppm is folded and, for simplicity, is omitted from this analysis.

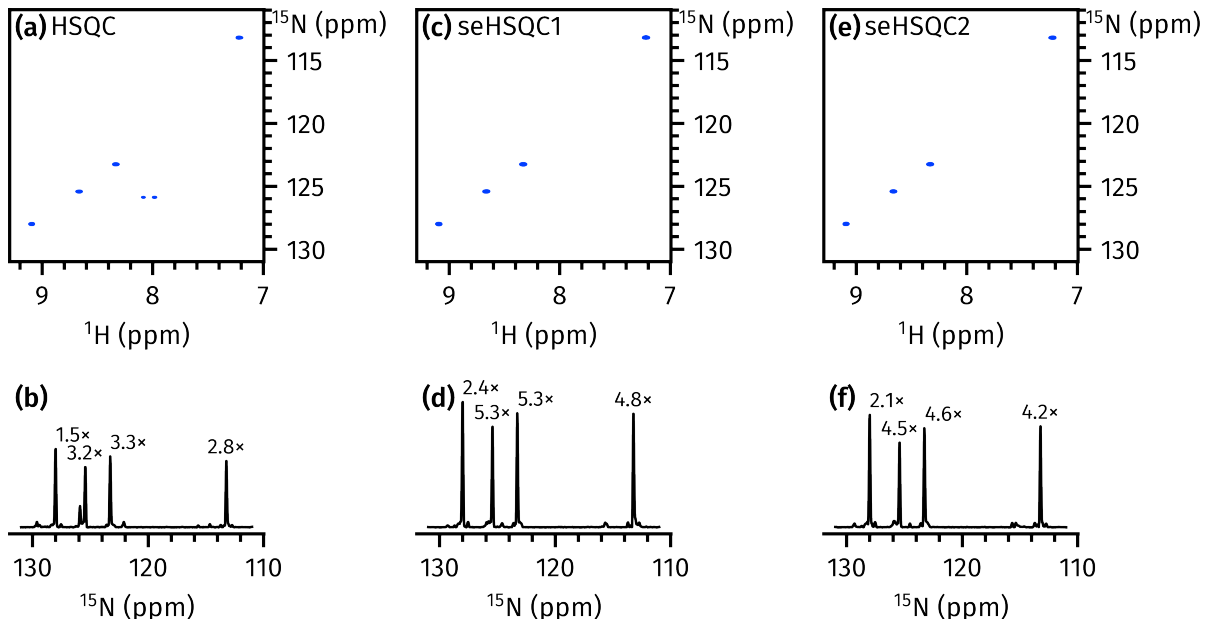


Figure 4.29: Comparison of NOAH ^1H - ^{15}N modules; the 2D spectra and their positive projections onto the F_1 axis are shown. Numbers above each peak in the projections indicate sensitivity increases compared to the HMQC module (the HMQC spectra themselves can be seen in, e.g., figs. 4.27a and 4.27b). (a)-(b) HSQC module. (c)-(d) seHSQC1 module. (e)-(f) seHSQC2 module. Data code: 7G-201017.

As can be seen, the (non-enhanced) HSQC alone allows for substantial sensitivity increases over the HMQC, with over 3 \times gains being accomplished simply due to multiplet collapse. The seHSQC experiments push this even further: the PEP transfer block provides an approximate 1.5 \times sensitivity improvement for all peaks, which is on top of the boost already provided by the HSQC. In contrast to the ^{13}C case, the ^{15}N seHSQC1 module outperforms the seHSQC2 in terms of raw sensitivity, though (as before) there is no clear explanation for this subtle difference.

However, this apparent advantage of the seHSQC1 vanishes when looking at the ^{13}C seHSQC module which comes later in the supersequence. When preceded by the ^{15}N HSQC or seHSQC1 modules, this module suffers from severely decreased sensitivity, compared to when the seHSQC2 module is used (fig. 4.30). As the spectra and their F_1 projections show, this reduction in sensitivity actually stems from broadening in the indirect dimension. This in turn arises because $^1\text{H}^{15}\text{N}$ magnetisation (which includes the $^1\text{H}^{13}\text{C}$ component sampled in this module—by definition, protons coupled to ^{13}C are not coupled to ^{15}N) precesses in the transverse plane during the t_1 period of the ^{15}N HSQC and seHSQC1. Evolution of ${}^nJ_{\text{HH}}$, as well as T_2 relaxation, lead to a t_1 -dependent signal loss which (after Fourier transformation) is manifested as line broadening and significantly reduced peak heights (figs. 4.30c and 4.30f).

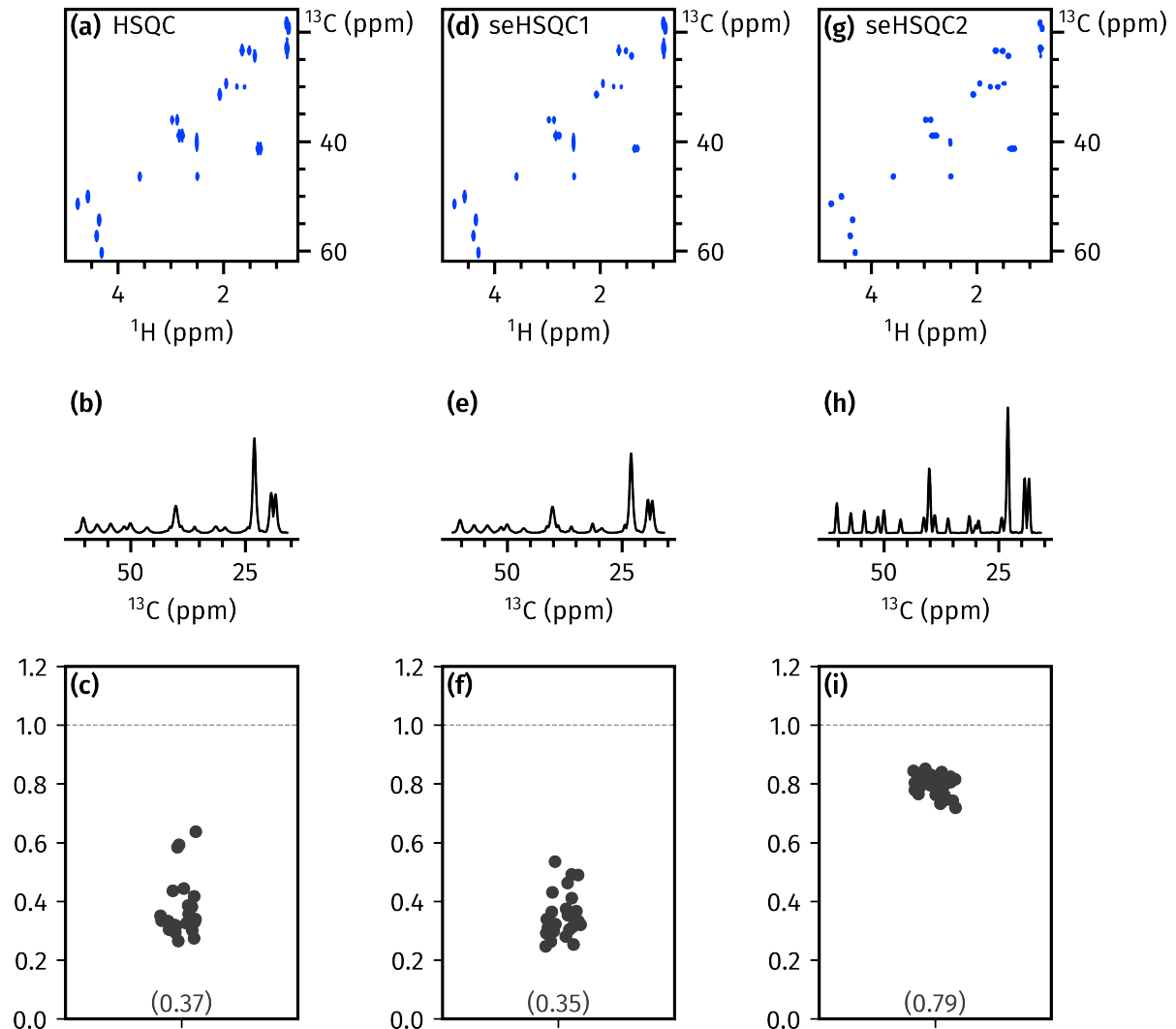


Figure 4.30: Comparison of the ^{13}C seHSQC module from NOAH-3 $\text{XS}^+\text{C}^{\text{c}}$ supersequences, where X is a $^1\text{H}-^{15}\text{N}$ module. The 2D spectra, their positive projections onto the F_1 axis, as well as the relative intensities of all peaks (as compared to the corresponding module in the NOAH-3 $\text{M}_\text{N} \text{S}^+\text{C}^{\text{c}}$ supersequence), are shown. (a)–(c) $\text{X} = ^{15}\text{N}$ HSQC. (d)–(f) $\text{X} = ^{15}\text{N}$ seHSQC1. (g)–(i) $\text{X} = ^{15}\text{N}$ seHSQC2. Data code: 7G-201017.

While the ^{15}N seHSQC2 module also does not preserve $^1\text{H}^{15}\text{N}$ magnetisation as well as the HMQC module (with an approximately 20% loss, see fig. 4.30i), this is not anywhere as drastic as with the other modules. Furthermore, these losses are constant across all t_1 increments, so there is no line broadening observed in the modules which follow. This small sensitivity loss in later modules is also a reasonable price to pay for the improvement in the ^{15}N module itself: typically, it is the ^{15}N module which has the smallest intrinsic sensitivity, so it is desirable to maximise this, even at the cost of the more sensitive modules.

It should be noted that the ^{13}C seHSQC1 (and HSQC) modules also do place $^1\text{H}^{13}\text{C}$ magnetisation in the transverse plane during t_1 . However, they do not cause (appreciable) broadening in

downstream modules, as evidenced by the almost complete preservation of CLIP-COSY peak heights in fig. 4.11. This is because indirect-dimension spectral widths for ^{13}C experiments are often much wider, and the acquisition times (AQ_1 , as defined in § 4.3.2) consequently shorter. For example, in the NOAH-3 XS^+C^c experiments shown here, the values of AQ_1 for the ^{15}N and ^{13}C experiments are respectively 60.1 ms and 5.38 ms. Thus, the effect of magnetisation losses during t_1 is not so keenly felt.

CTP gradient duration

To verify that the 2.5 ms CTP gradients were necessary in the ^{15}N seHSQC as well, I repeated the NOAH-3 $\text{S}_\text{N}^+\text{S}^+\text{C}^c$ experiment with increasing gradient durations. Only the seHSQC2 module was studied here, as the HSQC and seHSQC1 modules are not viable for use in NOAH supersequences. It should be noted that the durations of *only* the CTP gradients g_1 and g_2 (in fig. 4.9c) were modified; g_3 and g_4 retain their original durations of 1 ms. The results, shown in fig. 4.31, are qualitatively very similar to that obtained with the HMQC (in fig. 4.25): increasing the gradient duration does not lead to substantial differences in signal intensity, but does provide much improved artefact suppression. In this case, equivalent results can be obtained using gradient durations of 2.2 ms to 2.3 ms; however, 2.5 ms still appears to be a reliable ‘default’ value which can be used without having to perform further optimisations.*

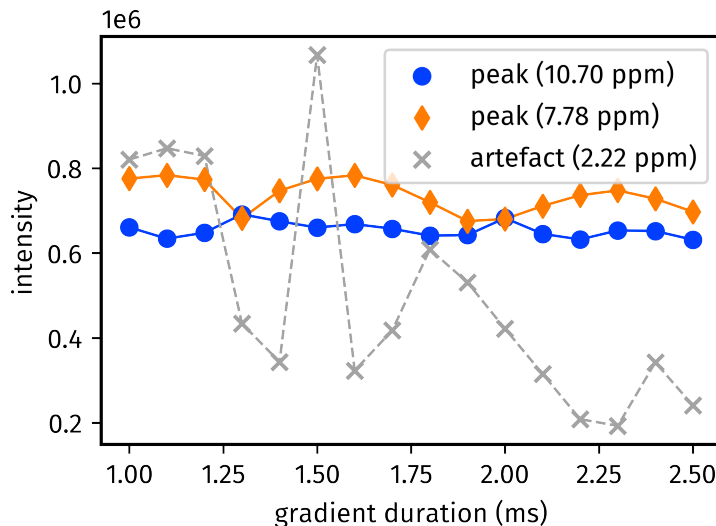


Figure 4.31: Variation of seHSQC2 signal and artefact intensity with CTP gradient duration, as measured by (absolute) peak heights in the positive F_2 projection. Data code: 7Z-200926.

*I am aware that this statement somewhat undermines my arguments for the use of POISE optimisations in chapter 3.

k- and SW-scaling

Finally, the effects of *k*- and SW-scaling were also investigated for the seHSQC2 module. The effects of just performing scaling are very minimal (fig. 4.32): unlike the HMQC module where lowering the resolution avoids losses in peak height due to multiplet structure, there are no such gains for the seHSQC module. The use of indirect-dimension linear prediction, however, is rather more successful for the seHSQC module (fig. 4.33): again, this likely stems from the lack of multiplet structure in F_1 . As before, the application of LP to the SW-scaled spectra yields better peak shapes than the *k*-scaled spectra (compare, e.g., figs. 4.33g and 4.33m). The same caveats as to detection sensitivity are also applicable in this instance.

4.3.4 Dual HSQC and HSQC-TOCSY

Having completed our survey of ^{15}N modules, we now return to ^{13}C modules. In particular, I was interested in how *two* (or more) modules drawing on $^1\text{H}^{\text{C}}$ magnetisation could be combined in the same supersequence. All of the ^{13}C modules described so far fully consume $^1\text{H}^{\text{C}}$ magnetisation: if we were to crudely concatenate two such modules, the second of these modules will have greatly reduced sensitivity. This is acceptable only if the second module has a far greater intrinsic sensitivity. More ideally, we should like to have a way to *partition* the magnetisation pool between the two different modules as needed: this would allow us to balance their sensitivities.

This can be accomplished by modifying the first module to only excite a portion of $^1\text{H}^{\text{C}}$ magnetisation, storing the remainder along $+z$. The NOAH HSQC module—which is derived from the ASAP-HSQC experiment,^{11,45} previously described in § 3.4.5—does in fact allow for this, through a modification to the INEPT delay. I therefore begin by describing SSX-type supersequences, where the first HSQC only partially excites $^1\text{H}^{\text{C}}$ magnetisation, and the second consumes the remainder. Here, X represents a homonuclear module, which in this section is the CLIP-COSY; however, the conclusions drawn here are applicable to any homonuclear module.

Supersequences with two HSQC modules

In order to accomplish partial $^1\text{H}^{\text{C}}$ excitation in the NOAH HSQC module (fig. 4.34), the INEPT delay Δ_E should be modified from its usual value of $1/(4J)$ (where J is short for $^1J_{\text{CH}}$). After the $90^\circ(I)-\Delta_E-180^\circ(I, S)-\Delta_E$ INEPT block, the relevant product operators are

$$\cos(2\pi J \Delta_E) I_y - \sin(2\pi J \Delta_E) 2I_x S_z. \quad (4.15)$$

In a ‘normal’ INEPT block, the choice of $\Delta_E = 1/(4J)$ makes the cosine term vanish, leaving us with only the term $-2I_x S_z$. Since this term is subsequently transferred to spin S and labelled in t_1 , this choice of $\Delta_E = 1/(4J)$ corresponds to *complete* excitation of $^1\text{H}^{\text{C}}$ magnetisation.

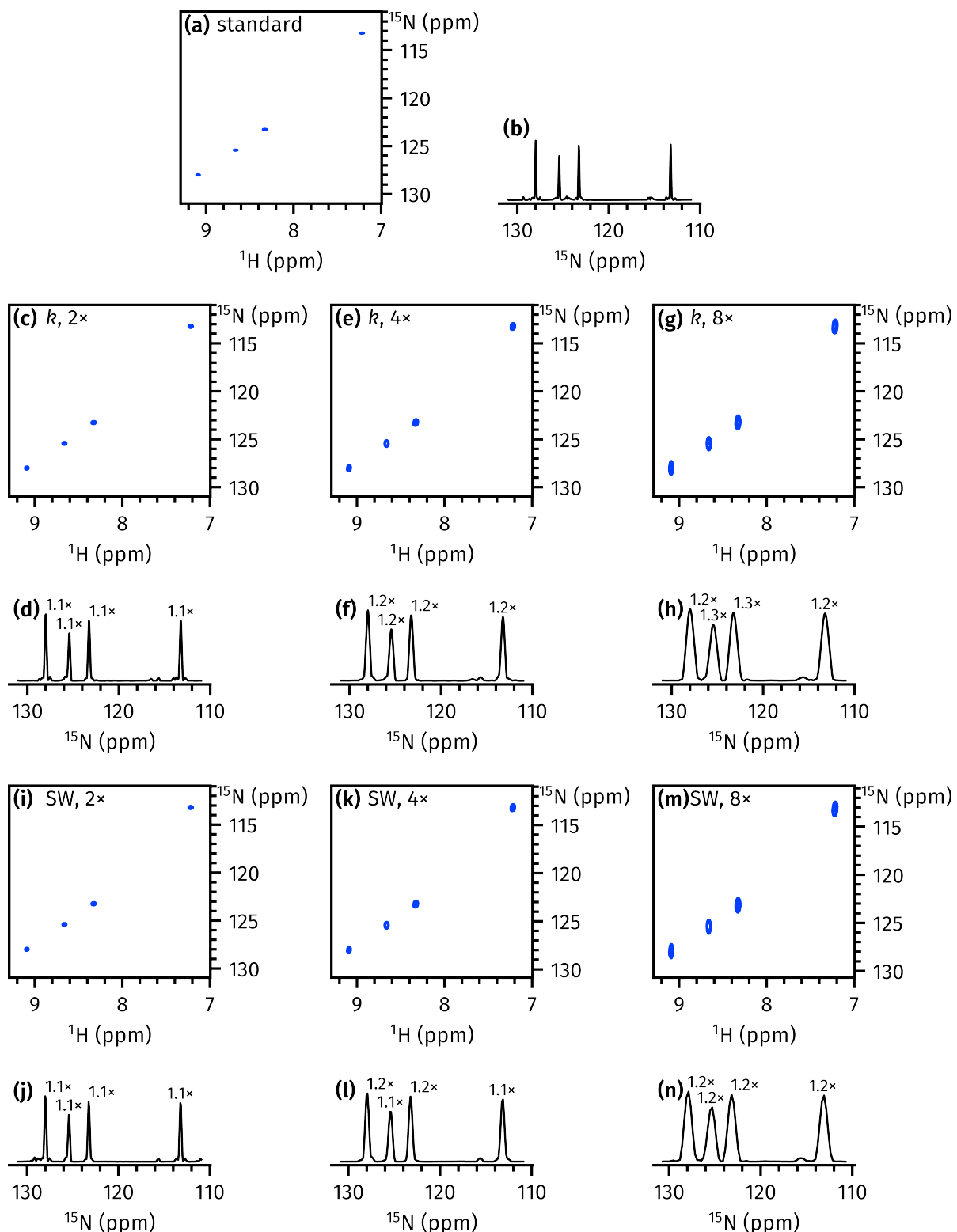


Figure 4.32: Effects of k - and SW-scaling on NOAH seHSQC2 spectrum (taken from NOAH-3 $\text{S}_\text{N}^+\text{S}^+\text{C}^\text{c}$ supersequences). Each seHSQC spectrum is shown together with a positive projection onto the F_1 axis. The relative SNR of each peak, with respect to the standard spectrum, is indicated on each of the other projections. (a)-(b) Standard spectrum. (c)-(h) k -scaled spectra. (i)-(n) SW-scaled spectra. Data code: 7G-210310.

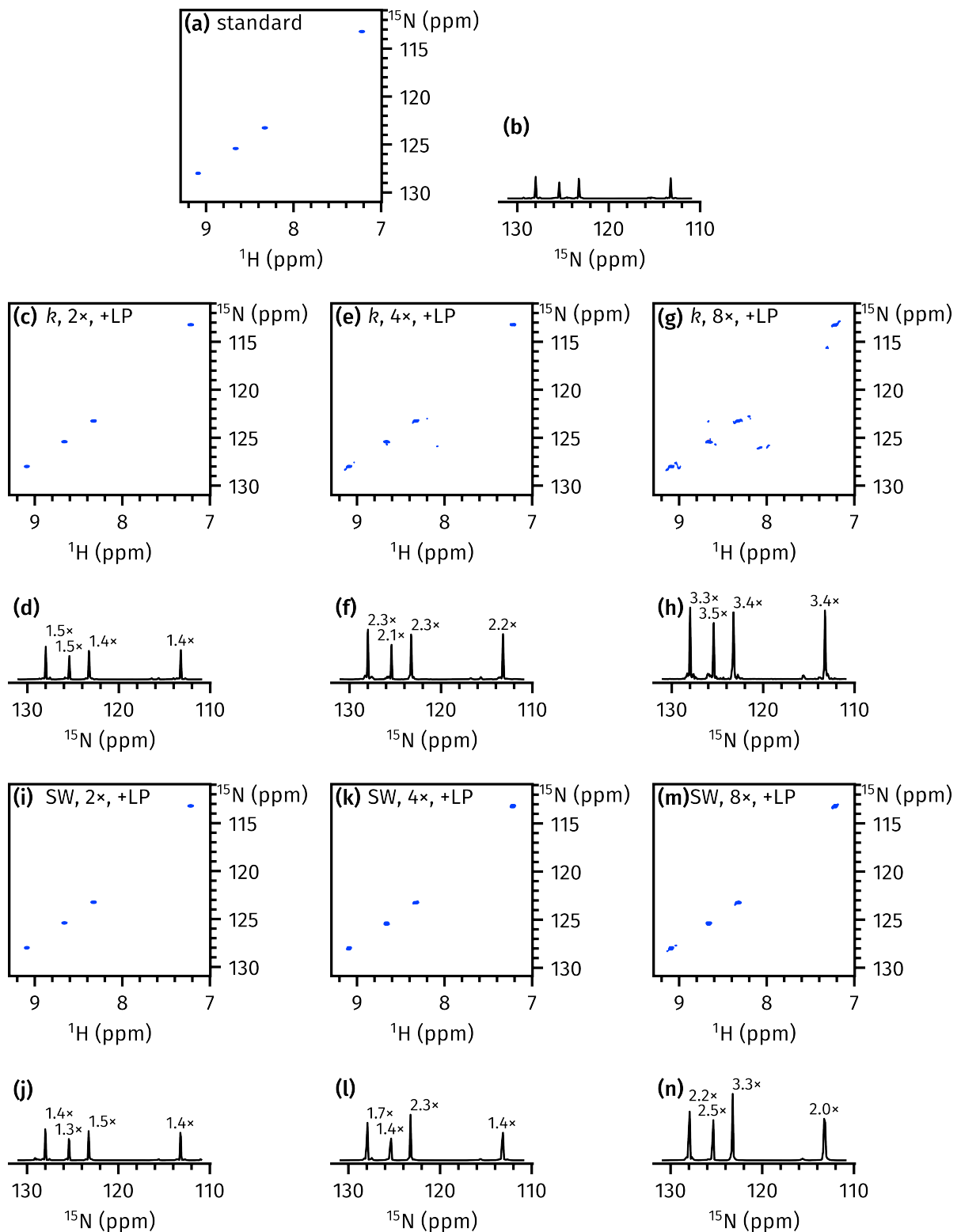


Figure 4.33: The same as in fig. 4.32, but with extra linear prediction applied to all scaled spectra to bring $\text{AQ}_{1,\text{eff}}$ up to its original value in the standard spectrum. Linear prediction of k times more points leads to a \sqrt{k} increase in noise; to account for this, all spectra are plotted with the same noise level. (a)–(b) Standard spectrum. (c)–(h) k -scaled spectra. (i)–(n) SW-scaled spectra. Data code: 7G-210310.

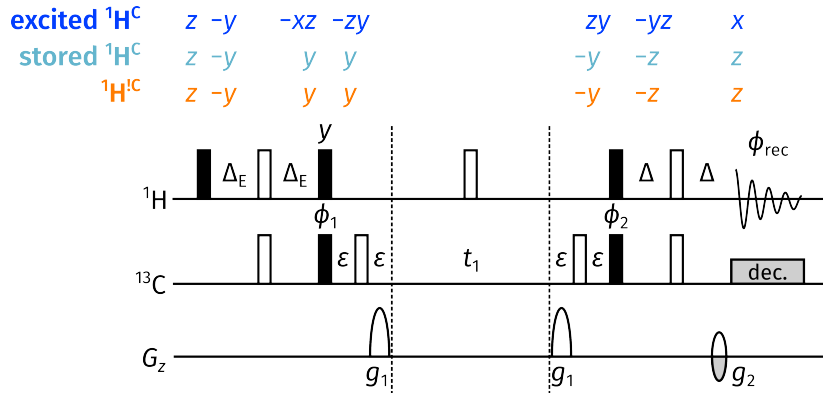


Figure 4.34: NOAH HSQC module with partial excitation, where the delay Δ_E is changed from the usual value of $1/(4J)$. As shown by the product operator analysis, proportion f of $^1\text{H}^C$ magnetisation is stored along $+z$ at the end of the sequence and can be sampled again in a later module. Gradient amplitudes are $(g_1, g_2) = (75\%, \pm 18.9\%)$; all other symbols have the same meaning as in fig. 4.9. (The different CTP gradient amplitudes are chosen to avoid accidental refocusing between different modules.)

However, if we choose $\Delta_E < 1/(4J)$, then the first I_y term—which corresponds to ‘stored’ or unexcited $^1\text{H}^C$ magnetisation—is eventually returned to I_z , as shown by the product operator analysis in fig. 4.34.* To be precise, in order to excite a fraction f of $^1\text{H}^C$ magnetisation (and store the remaining $(1 - f)$ for the next module), we require that

$$\Delta_E = \frac{2\Delta \arcsin f}{\pi}, \quad (4.16)$$

where Δ is the usual value of $1/(4J)$. The spectra obtained from a NOAH-3 SSC^C supersequence, using this partial $^1\text{H}^C$ excitation in the first HSQC module (with $f = 0.8$), are shown in fig. 4.35.

In fig. 4.36a, the intensities of these spectra are compared against the HSQC and CLIP-COSY in a NOAH-2 SC^C supersequence. As expected, the first HSQC retains (around) 80% of its ‘base’ sensitivity, corresponding to the fraction of excited magnetisation. The second HSQC spectrum *nominally* only has access to the remaining 20% of magnetisation: however, due to recovery of $^1\text{H}^C$ magnetisation during the FID of the first HSQC, this number is boosted to 65%. The CLIP-COSY sensitivity is (very slightly) decreased compared to a NOAH-2 SC^C supersequence: this is because of cumulative losses from the two HSQC modules.

Figures 4.36b to 4.36d detail several possible modifications to this basic scheme. In the first of these (fig. 4.36b), we further increase the sensitivity of the second HSQC module by converting

*In the context of the ASAP experiments (both HMQC and HSQC), this is termed *Ernst angle* excitation because the unexcited magnetisation is stored for later repetitions of the *same* experiment, which mirrors the original Ernst angle for the 1D pulse–acquire experiment. However, for NOAH supersequences, the magnetisation is stored for other modules, so it is more correct to refer to this as a *partial* excitation.

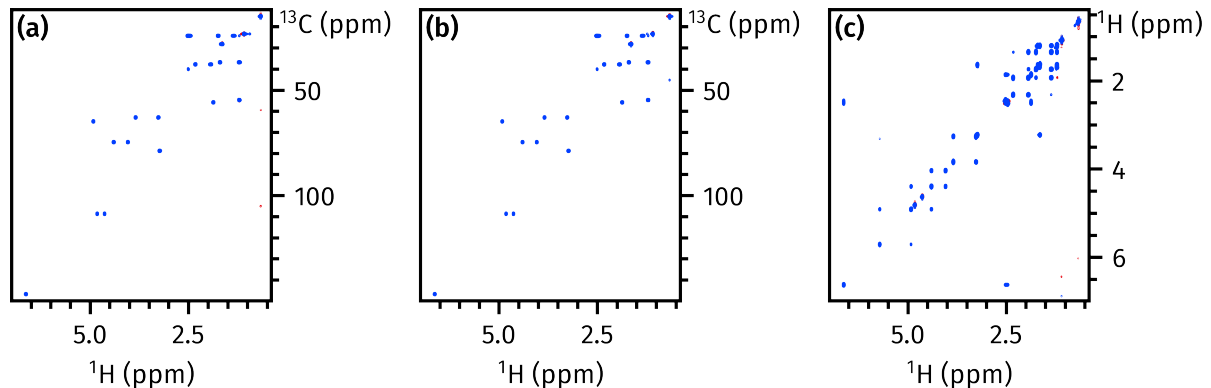


Figure 4.35: Spectra from a NOAH-3 SSC^c supersequence, where the INEPT delay of the first HSQC module was modified to only excite a fraction $f = 0.8$ of ${}^1\text{H}^{\text{C}}$ magnetisation. (a) First HSQC. (b) Second HSQC. (c) CLIP-COSY. Data code: 7A-201010.

it into a seHSQC2 module.* The CLIP-COSY sensitivity is slightly decreased due to poorer ${}^1\text{H}^{\text{C}}$ preservation, which is consistent with § 4.3.1. Note, however, that it is not possible to change the *first* HSQC into a seHSQC experiment: none of the seHSQC versions discussed in § 4.3.1 preserve unused ${}^1\text{H}^{\text{C}}$ magnetisation in the same way as the HSQC module above.

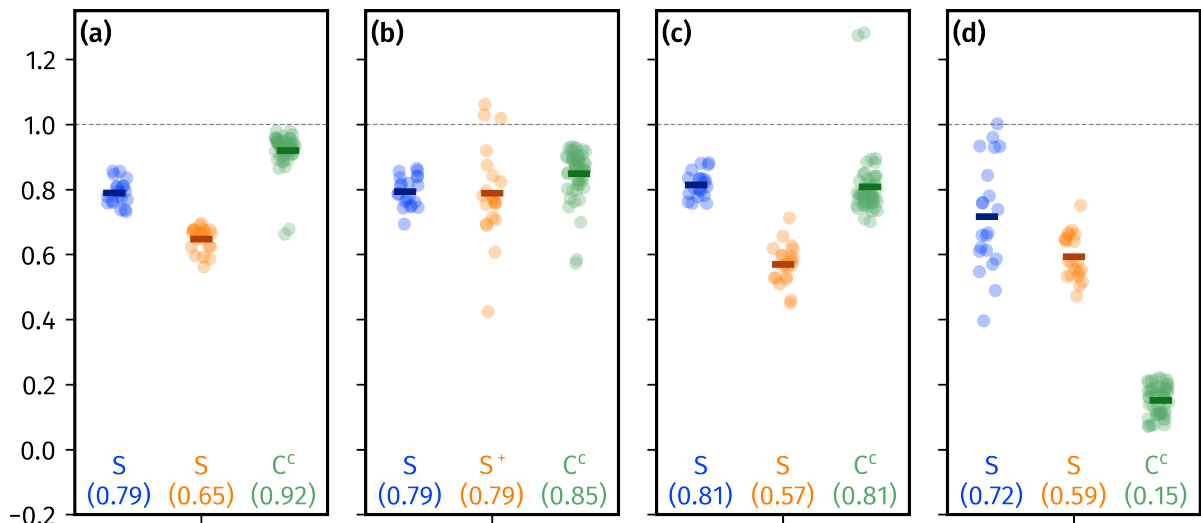


Figure 4.36: Comparisons of (se)HSQC and CLIP-COSY sensitivities of NOAH-3 SSC^c and SS^+C^c supersequences. The three groups of dots in each subplots refer to the three respective modules in the supersequence. Peak intensities are normalised against the HSQC and CLIP-COSY experiments in a NOAH-2 SC^c supersequence. (a) NOAH-3 SSC^c with $f = 0.8$ (the same spectra as shown in fig. 4.35). (b) NOAH-3 SS^+C^c with $f = 0.8$. (c) NOAH-3 SSC^c with $f = 0.8$, plus 35 ms DIPSI-2 mixing after the first HSQC module. (d) A NOAH-3 SSC^c supersequence, but using the split-seHSQC implementation of Nolis et al.³⁸ (as opposed to the ASAP-HSQC module) for the double HSQC. Data code: 7A-201010.

Alternatively, it is also possible to include a period of isotropic mixing between the two HSQC

*The sensitivity increases are not uniform because they depend on multiplicity.

modules: here, the DIPSI-2 sequence⁷⁶ was chosen. Since the $^1\text{H}^{\text{C}}$ magnetisation pool has been (partially) depleted, and the $^1\text{H}^{\text{C}}$ magnetisation pool is (almost) full, the application of DIPSI mixing should in theory lead to transfer of polarisation from the $^1\text{H}^{\text{C}}$ pool to ^1H . However, when tested, this was not found to have a beneficial impact: in fact, small *decreases* in sensitivity were observed for both the second HSQC module and the final CLIP-COSY (fig. 4.36c).

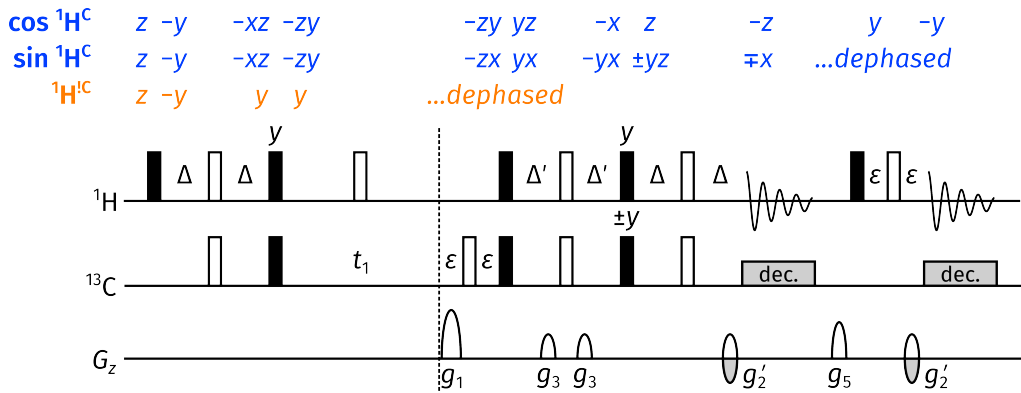


Figure 4.37: Split-seHSQC experiment used for the collection of two HSQC spectra.³⁸ All symbols have the same meaning as in fig. 4.9a; g_5 is a purge gradient with arbitrary amplitude.

Lastly, it should also be noted that the acquisition of two HSQC spectra in one experiment has also been accomplished in a different way by Parella and coworkers.^{38,39} This relies on ‘splitting’ up the CRK seHSQC experiment (fig. 4.9a), such that the cosine- and sine-modulated components are separately detected.* This can be easily accomplished by inserting a detection period before the final ^1H 90° pulse (fig. 4.37), where the cosine-modulated component is stored as longitudinal magnetisation and the sine-modulated component is transverse (and in-phase).

However, this implementation does not fare as well in the context of the NOAH-3 SSC^c supersequences investigated here. The sensitivity of the three modules when using this is shown in fig. 4.36d: the two HSQC spectra have comparable, but slightly poorer, sensitivity than the $f = 0.8$ partial excitation scheme used in fig. 4.36a. There is also a slightly larger spread in sensitivity, which can be attributed to the fact that the seHSQC does not boost intensities uniformly. Furthermore, since this experiment (like the parent CRK seHSQC) dephases $^1\text{H}^{\text{C}}$ magnetisation, the CLIP-COSY which follows suffers from a large drop in intensity. Prepending the experiment with the ZIP element to make a ‘split seHSQC2’ does not help, because the bulk magnetisation is in the transverse plane during the first FID. Finally, the split-seHSQC implementation does not easily allow for features such as multiplicity editing to be independently applied to one of the two signals.[†]

* In my paper,¹ this was referred to as an ‘MFA’ (multiple-FID acquisition) implementation. I refrain from using this term here because it is ambiguous—after all, NOAH experiments are also multiple-FID experiments.

[†] It should at least be *possible* to apply editing to one signal using a modification similar to that done by Nolis et al.³⁹ However, that is not as simple as the ASAP implementation, where each module is separate.

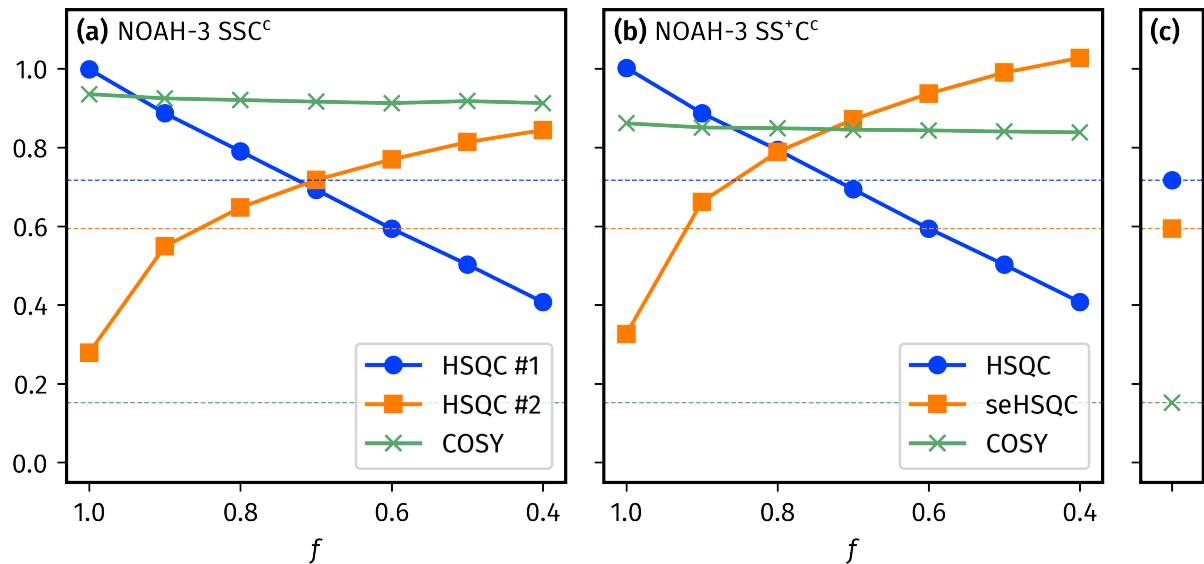


Figure 4.38: Sensitivity comparisons for NOAH-3 SSC^c and NOAH-3 SS^+C^c supersequences. Intensities are compared against the HSQC and CLIP-COSY modules in a NOAH-2 SC^c experiment, and averaged over all peaks. (a) NOAH-3 SSC^c supersequences acquired with different values of f . (b) NOAH-3 SS^+C^c supersequences acquired with different values of f . (c) Using the split-seHSQC implementation. Data code: 7A-201010.

Figure 4.38 summarises all of these factors, and further shows how sensitivity varies as a function of the parameter f . From figs. 4.38a and 4.38b, it is clear that the sensitivity of the first HSQC is directly proportional to f . The sensitivity of the second HSQC module in fig. 4.38a stems from both the unexcited $(1 - f)$ proportion of ${}^1\text{H}^{\text{C}}$ magnetisation, plus any that has recovered during the first FID: this latter component decreases as f increases. The CLIP-COSY intensities generally do not vary with f . In fig. 4.38b, the second HSQC module is replaced with the seHSQC2: this allows f to be increased in order to balance the sensitivities of the two HSQC spectra. Finally, for completeness, the split-seHSQC implementation is shown in fig. 4.38c; the numbers here are the same as in fig. 4.36d. It is clear that there are several values of f which allow the NOAH/ASAP-type implementation to outperform the split-seHSQC experiment in sensitivity.

On its own, the acquisition of two HSQC spectra—as has so far been shown—is not particularly groundbreaking. However, it is possible to differentiate the two HSQC signals and thereby extract more information. For example, one spectrum may be run without decoupling in order to measure one-bond coupling constants;^{38,77} multiplicity editing may be implemented;⁴⁵ or the indirect-dimension spectral width of one of the HSQC spectra can be changed in order to make use of spectral aliasing techniques.^{25,39} Most interesting of all, though, would be to modify the pulse sequence itself to obtain different *types* of ${}^1\text{H}-{}^{13}\text{C}$ spectra: for example, the addition of an isotropic mixing block to the first HSQC yields an HSQC-TOCSY + HSQC combination,³⁸ which I now discuss.

HSQC-TOCSY

The addition of DIPSI mixing inside the partial-excitation HSQC module leads to an HSQC-TOCSY module (fig. 4.39a), which is (conceptually) identical to the ASAP-HSQC-TOCSY experiment also previously reported by Luy and coworkers.⁷⁸ Note that the ^1H 90° pulse immediately after t_1 has a different phase from in the HSQC module: this ensures that $^1\text{H}^{13}\text{C}$ (and unused $^1\text{H}^1\text{H}$) magnetisation is returned to $+z$ after accounting for the extra 180° pulse at the end.

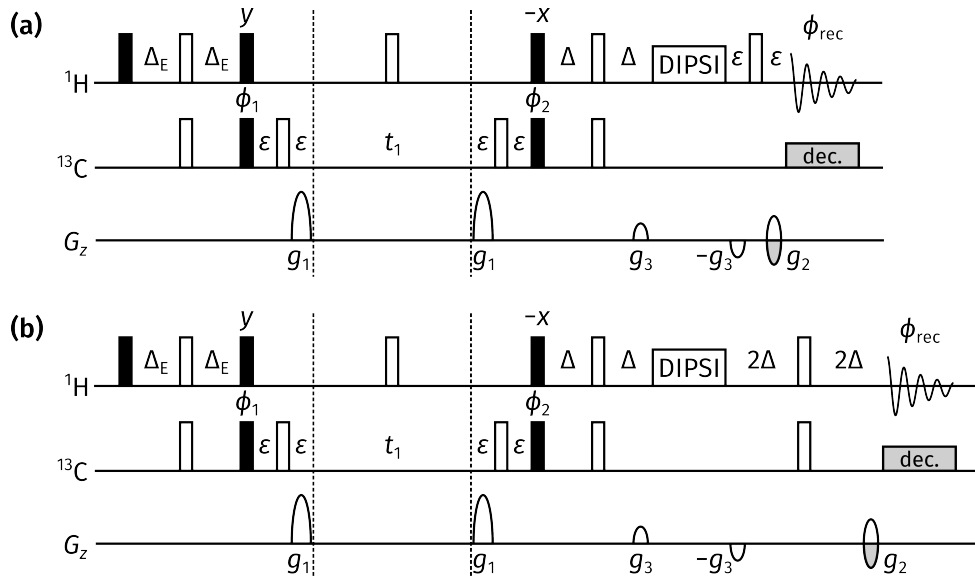


Figure 4.39: NOAH HSQC-TOCSY module: (a) without inversion of HSQC signals ('direct' responses); and (b) with inversion of HSQC signals.

The HSQC-TOCSY module may optionally be implemented with inversion of 'direct' HSQC-type responses, which correlate ^{13}C nuclei with directly bonded protons. This is to be contrasted with the 'indirect' responses, which correlate ^{13}C nuclei with other remote protons in the same spin system, and represent magnetisation which has been transferred during the isotropic mixing period. This inversion may be accomplished by expanding the final gradient echo to a duration of $4\Delta = 1/J$ (fig. 4.39b); ^{13}C -bound protons will evolve under $^1J_{\text{CH}}$ and acquire an additional 180° phase shift compared to remote protons.*

The use of such editing in the HSQC-TOCSY is perhaps slightly risky, because it may lead to unwanted peak cancellation through accidental overlap. Furthermore, if used together with a second HSQC module (as discussed here), this information does not need to be encoded in the HSQC-TOCSY. Thus, it is not enabled by default; the user must explicitly turn it on through an acquisition flag. The concept of inverting remote peaks is more useful in the HSQC-COSY, which will be discussed in § 4.3.5.

*Of course, the remote protons may also happen to be bonded to ^{13}C , but the chance of this is only roughly 1%.

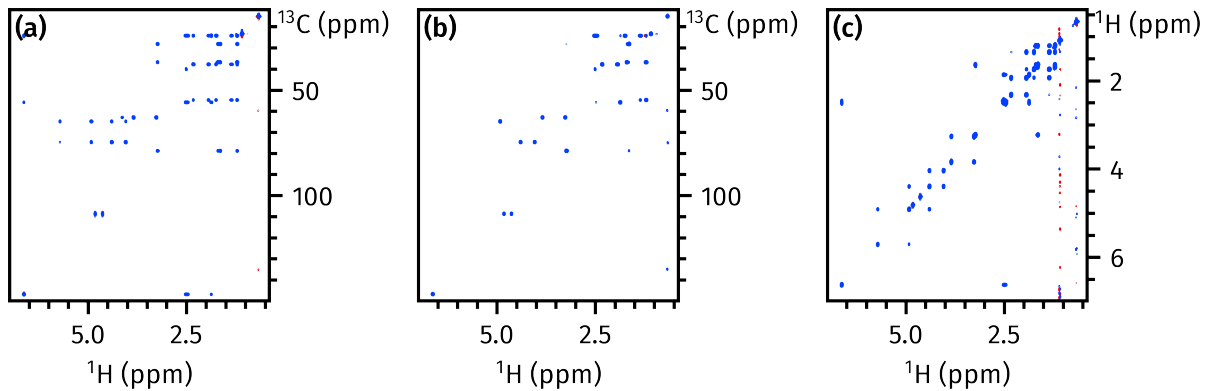


Figure 4.40: Spectra from a NOAH-3 $S^T S^+ C^c$ supersequence. (a) HSQC-TOCSY ($f = 0.9$). (b) seHSQC2. (c) CLIP-COSY. Data code: 7A-210126.

With this module in hand, it is possible to implement either NOAH-3 $S^T SC^c$ or NOAH-3 $SS^T C^c$ supersequences which yield both HSQC and HSQC-TOCSY data. In both cases, the first of the two ^{13}C modules can be adjusted to only partially excite $^1\text{H}^C$ magnetisation. However, the former is preferred as the HSQC-TOCSY has a lower intrinsic sensitivity compared to the HSQC: thus, it is better to place this first in the supersequence, and to also use a larger value of f in order to balance the relative sensitivities. Figure 4.40 shows an example of the spectra obtained with a setting of $f = 0.9$.

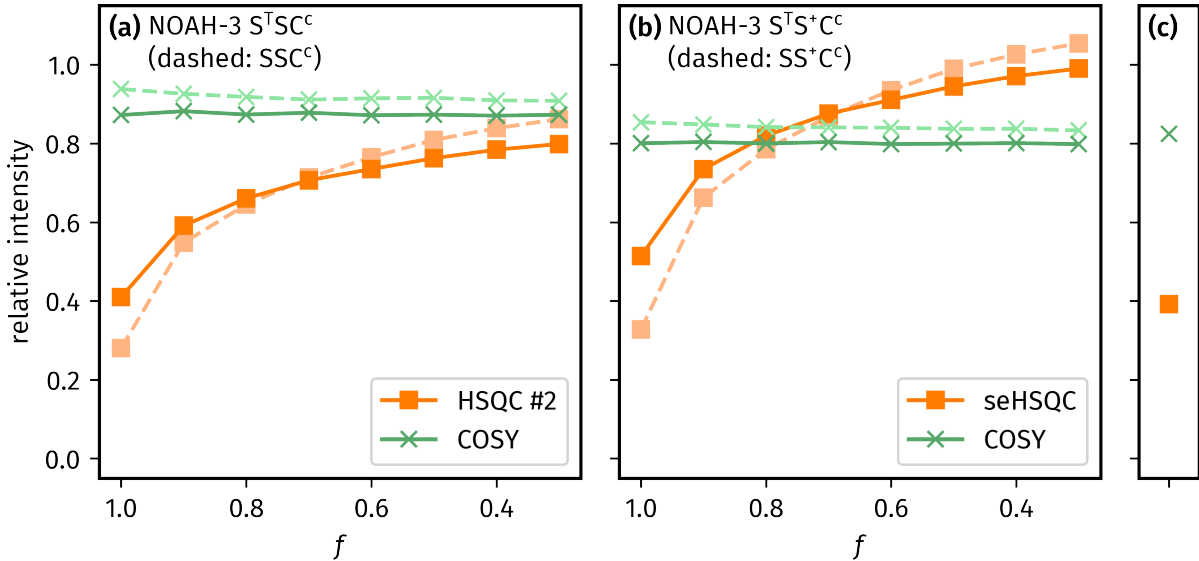


Figure 4.41: Sensitivity comparisons for NOAH-3 SSC^c and NOAH-3 $S^T SC^c$ supersequences. The intensities of the latter two modules are compared against the HSQC and CLIP-COSY modules in a NOAH-2 SC^c experiment, and averaged over all peaks. (a) NOAH-3 $S^T SC^c$ supersequences acquired with different values of f ; the dashed lines are the corresponding intensities in the NOAH-3 SSC^c supersequence. (b) The same, but with the seHSQC2 as the second module instead. (c) The corresponding intensities of the second and third modules in a seHSQC-TOCSY + HSQC + CLIP-COSY supersequence: this is to be compared with the first data point ($f = 1$) in (a). Data code: 7A-210126.

Finally, I performed a series of experiments to establish whether changing the first module from an HSQC (i.e. SSC^c) to an HSQC-TOCSY (i.e. NOAH-3 $S^T\text{SC}^c$) would affect the sensitivities of the two latter modules. The results are shown in figs. 4.41a and 4.41b, where the solid lines represent the supersequences starting with the HSQC-TOCSY, and the dashed lines those starting with the HSQC. At large values of f (corresponding to depletion of the ${}^1\text{H}^C$ pool), the subsequent ${}^{13}\text{C}$ module experiences sensitivity *gains* when the HSQC-TOCSY is used. However, this changes at smaller values of f , where a drop in intensity is instead observed. For the CLIP-COSY module, the HSQC-TOCSY leads to slightly lower sensitivities across the board. All of these changes are, however, extremely minor: the supersequences shown here still work entirely as expected.

Figure 4.41c shows one final alternative which has not yet been considered: the use of the seHSQC-TOCSY⁵⁷ as the first module. This experiment is similar to the seHSQC2 module, but with the addition of isotropic mixing. Although this fully maximises the intensity of the HSQC-TOCSY component (not shown in fig. 4.41), and also preserves ${}^1\text{H}^C$ magnetisation for the CLIP-COSY (as evidenced by the relatively high CLIP-COSY intensity in fig. 4.41c), it cannot retain any portion of ${}^1\text{H}^C$ magnetisation. Thus, any later ${}^{13}\text{C}$ module which draws on the same magnetisation pool will only be able to use any signal which has recovered during the FID, just like in an experiment run with $f = 1$. In this case, the second module is a plain HSQC (not seHSQC): thus, its sensitivity is similar to the leftmost point in fig. 4.41a.

4.3.5 HSQC-COSY

One downside of the HSQC-TOCSY experiment discussed in the previous section is the (largely) indiscriminate transfer of magnetisation effected by the DIPSI mixing. It is difficult to determine how many transfer steps give rise to each peak, unless several HSQC-TOCSY spectra with different mixing times are recorded, which is a time-consuming process.*

For unambiguous spectral assignment, it may be preferable to have a mixing process which only effects coherence transfer across a single coupling: this can be described as an HSQC-COSY experiment. Similar experiments, including the H2BC,^{79,80} 2BOB/H2OBC,⁸¹ and HMQC-COSY⁸² have previously been proposed: the 2BOB experiment in particular has previously been incorporated in NOAH supersequences.⁵¹ These often use constant-time techniques in order to remove ${}^nJ_{\text{HH}}$ splittings and minimise linewidths in the indirect dimension; however, the drawback of this approach is that peak *amplitudes* are modulated by ${}^nJ_{\text{HH}}$. Furthermore, it is not generally possible to obtain absorption-mode lineshapes for all peaks in the spectrum: typically the ‘direct’ responses are in-phase absorption, and ‘indirect’ responses antiphase dispersion. (Here, the

*Of course, as per the previous section, this could be done in a NOAH supersequence where each HSQC-TOCSY spectrum is assigned a fraction of the ${}^1\text{H}^C$ magnetisation. However, one must be wary of stretching the same magnetisation pool too thinly between many different modules: the resulting decrease in sensitivity may lead to peaks being lost.

terms in-phase and antiphase are used with respect to ${}^nJ_{\text{HH}}$; specifically, for the indirect peaks, this refers to the ‘active’ coupling over which the coherence was transferred).

HSQC-CLIP-COSY

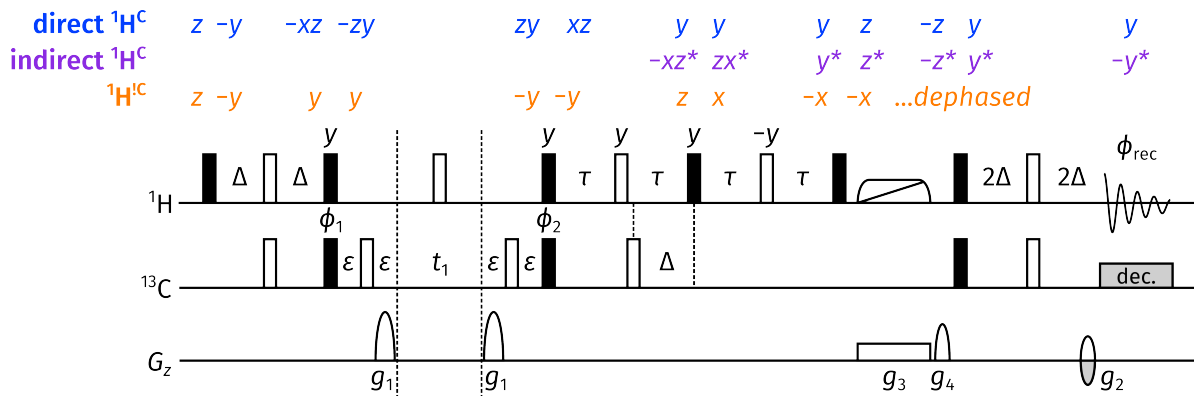


Figure 4.42: HSQC-CLIP-COSY experiment with product operator analysis for the HSQC-COSY signal derived from the ${}^1\text{H}^{13}\text{C}$ magnetisation, as well as the bulk ${}^1\text{H}^{13}\text{C}$ magnetisation. Both the ‘direct’, HSQC-type peaks, as well as the ‘indirect’ responses arising from coherence transfer in the perfect echo block, are analysed. The shorthand notation for product operators is expanded here to deal with a three-spin IKS system, where I and S are a mutually bonded ${}^1\text{H}-{}^{13}\text{C}$ pair as usual, and K is a ‘remote’ proton coupled to I . Terms with asterisks are on spin K ; thus, for example, zx^* refers to an $I_z K_x$ term. The delay τ is chosen to be $1/(4 \cdot \sum {}^nJ_{\text{HH}})$; typically, this sum of couplings is set as 30 Hz, leading to a value of $\tau = 8.33$ ms. The ZQF gradient g_3 should be calibrated as per Thrippleton et al.;⁸³ g_4 is a purge gradient with arbitrary amplitude. All other symbols have the same meaning as in fig. 4.9.

These problems are circumvented by the HSQC-CLIP-COSY experiment,^{84,85} where the basic HSQC experiment is combined with clean in-phase (CLIP) coherence transfer using a perfect echo^{86–88} and ZQF⁸³ (fig. 4.42). The use of a HSQC-type experiment means that ${}^nJ_{\text{HH}}$ does not evolve during t_1 , and the CLIP transfer ensures that all peaks have in-phase absorption lineshapes. Much like the HSQC-TOCSY experiment before it, this experiment may be implemented with direct/indirect response editing: it is this version of the experiment which is shown in fig. 4.42.* If this editing is not desired, then the final $2\Delta-180^\circ(I, S)-2\Delta$ spin echo can simply be reduced to a minimal gradient echo, i.e. $\epsilon-180^\circ(I)-\epsilon$.

Unfortunately, the HSQC-CLIP-COSY experiment does not preserve ${}^1\text{H}^{13}\text{C}$ magnetisation; thus, it cannot be directly used in a NOAH supersequence without sacrificing the sensitivity of terminal homonuclear modules. In particular, the ZQF used in the HSQC-CLIP-COSY experiment

*Note that this editing is orthogonal to *multiplicity editing*, which labels both direct and indirect responses with a sign that depends on the multiplicity of the ${}^{13}\text{C}$ nucleus detected in t_1 . The addition of multiplicity editing leads to highly confusing spectra, so is ignored here—although the option to do so *is* provided in the GENESIS pulse programmes if the user so desires.

dephases all magnetisation that is not along z . If we wished to retain ${}^1\text{H}{}^{13}\text{C}$ magnetisation, it would therefore have to (somehow) be placed along z during this period, and would have to be differentiated from the HSQC-COSY signal *after* the ZQF, which is not possible.* On top of that, the experiment also cannot be modified to provide partial excitation of ${}^1\text{H}{}^{13}\text{C}$ magnetisation, which limits the ways in which it can be combined with other ${}^{13}\text{C}$ modules.

Double spin echo HSQC-COSY

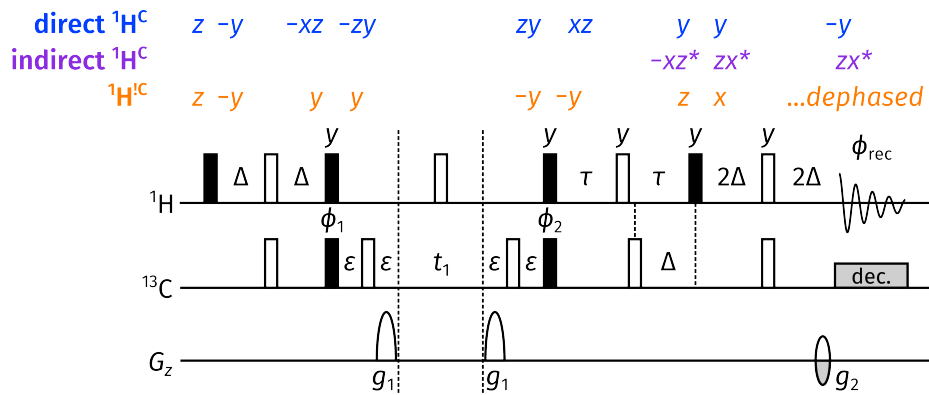


Figure 4.43: ‘Double spin echo’ HSQC-COSY experiment; all symbols have the same meaning as in fig. 4.42.

Before tackling these problems directly, I discuss a simpler version of the HSQC-COSY experiment, which uses a simple spin echo for ${}^2J_{\text{HH}}$ evolution instead of the perfect echo of the CLIP version. Together with the final 4Δ spin echo, this forms a ‘double spin echo’ (DSE) version of the HSQC-COSY experiment (fig. 4.43); the ‘double’ refers only to the mixing section after t_1 . The removal of the CLIP coherence transfer element leads to mixed lineshapes in this experiment, where the direct responses are (mostly) in-phase absorption, and the indirect responses (mostly) antiphase dispersion.[†]

Like the CLIP version, this DSE experiment does not preserve bulk magnetisation; it is also incompatible with partial ${}^1\text{H}{}^{13}\text{C}$ excitation. However, it does provide more raw sensitivity than the CLIP version: in the CLIP version, any antiphase signal at the end of the perfect echo is destroyed by the ZQF. All of this available signal is sampled in the DSE version, but this comes with a cost of not having pure absorption lineshapes.

It should be mentioned that prepending this DSE HSQC-COSY with the ZIP element would in fact return the bulk ${}^1\text{H}{}^{13}\text{C}$ magnetisation to $+z$ at the end of the sequence. Unfortunately, I did

*To be precise, although the ${}^1J_{\text{CH}}$ Hamiltonian could be used to separate bulk magnetisation and the HSQC-type ‘direct’ responses, the ‘indirect’ responses cannot be differentiated.

[†]The qualifier *mostly* is required because evolution of ${}^2J_{\text{HH}}$ during the final spin echo leads to mixtures of in-phase absorption and antiphase dispersion. This evolution is generally small, though, because 2Δ is smaller than τ .

not manage to test this experimentally, as my focus was on the development of the *triple* spin echo HSQC-COSY below.

Triple spin echo HSQC-COSY

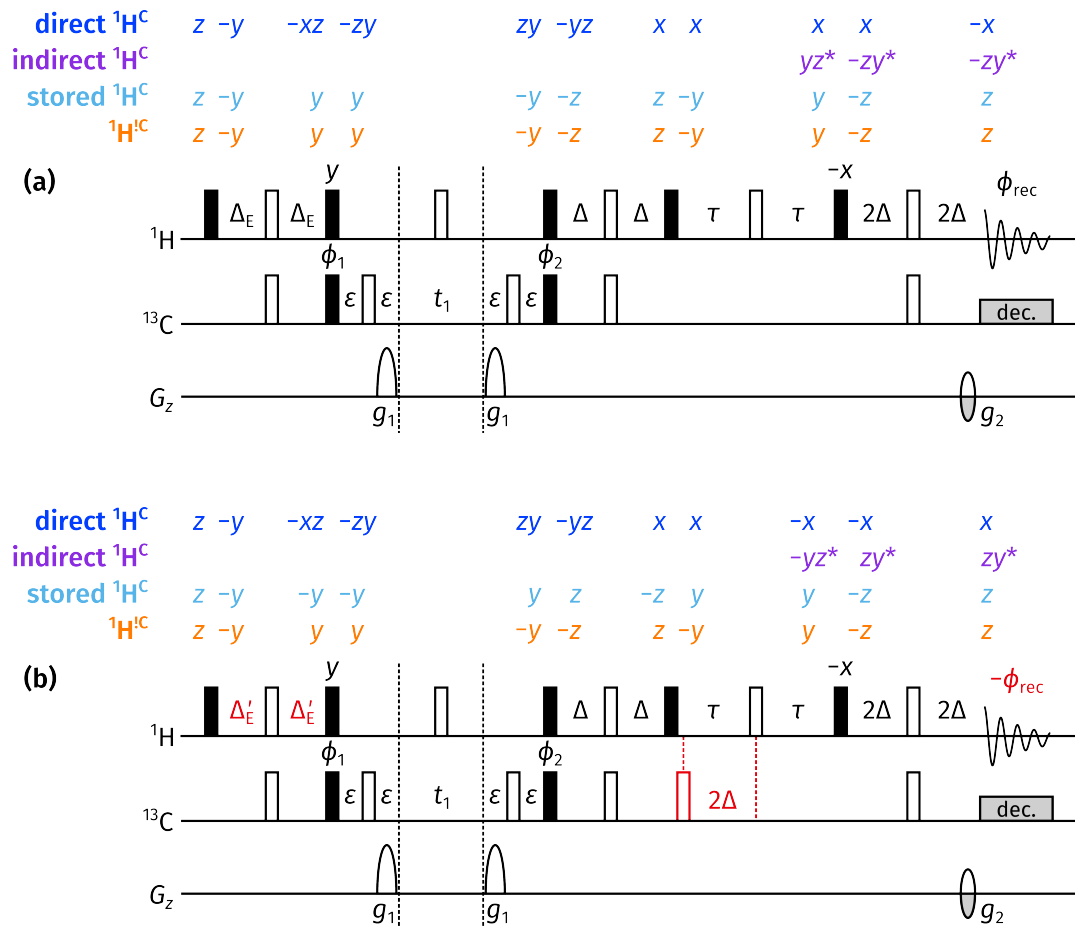


Figure 4.44: ‘Triple spin echo’ (TSE) HSQC-COSY experiment. (a) The first part of the experiment; this can be used on its own, but leads to spurious ‘relayed’ peaks arising via coherence transfer over two scalar couplings. (b) The second part of the experiment; co-adding this dataset with the first part leads to suppression of the relayed peaks. All symbols have the same meaning as in fig. 4.42.

In the DSE HSQC-COSY, the first of the two spin echoes after t_1 serves a dual purpose: $^1J_{\text{CH}}$ is allowed to evolve for a duration of $\tau - (\tau - \Delta) + \Delta = 2\Delta$ (thus generating peaks which are in-phase with respect to $^1J_{\text{CH}}$), and $^nJ_{\text{HH}}$ evolves for the total duration of 2τ (allowing coherence transfer to remote spins). The triple spin echo (TSE) HSQC-COSY is derived from the DSE version by separating the first spin echo into two distinct parts: one for $^1J_{\text{CH}}$ refocusing, and one for $^nJ_{\text{HH}}$ evolution. As shown by the product operator analysis in fig. 4.44a, this not only preserves the bulk $^1\text{H}^{\text{C}}$ magnetisation, but is also compatible with partial $^1\text{H}^{\text{C}}$ excitation.

The problem with the TSE version is the spectral quality: consider an even larger spin system

of the form S–I–K–L, where $\{I, K, L\}$ are ^1H and S is ^{13}C . At the end of t_1 , single-quantum magnetisation on spin I is present. Any evolution of J_{IK} in the first spin echo (of duration 2Δ) leads to terms of the form $2J_yK_z$, and can then be transferred onto spin K by the subsequent 90°_x pulse. This magnetisation can then further evolve under J_{KL} during the 2τ spin echo, and then be transferred a *second* time by the 90°_x pulse to spin L. Since the intensity of this transfer pathway is proportional to $\sin(2\pi J_{IK}\Delta)$, these ‘relay’ artefacts are especially prominent for large J_{IK} .

In order to suppress these, the experiment must be run a second time but with an additional 180° ^{13}C pulse inserted during the second spin echo (fig. 4.44b).^{*} Any magnetisation still on spin I will therefore evolve under $^1J_{\text{CH}}$ for a total period of 4Δ , leading to net inversion. However, magnetisation which has already been transferred to spin K at this point will not be inverted: it is this magnetisation which is responsible for the relay artefacts. Therefore, by subtracting the two datasets (or equivalently, inverting the receiver phase and adding the two datasets), the desired peaks will be added up and the artefacts cancelled through subtraction.

The insertion of this 180° pulse also causes any unexcited $^1\text{H}^{\text{C}}$ magnetisation to be inverted. In order to ensure that this is returned to $+z$ at the end of the sequence (and not $-z$), the initial INEPT delay must be *lengthened* instead of shortened, as per:

$$\Delta'_E = \frac{2\Delta(\pi - \arcsin f)}{\pi}, \quad (4.17)$$

where, as before, f is the proportion of magnetisation to be excited.

Supersequences

The spectral quality, and sensitivity, of all of these modules is captured in figs. 4.45 and 4.46. The CLIP version (figs. 4.45a and 4.46a) has the best lineshapes, but its sensitivity is slightly lower. The DSE version (figs. 4.45b and 4.46b) provides greater sensitivity, but at the cost of impure lineshapes.

For the TSE version, we first look at the importance of the relay artefact suppression procedure described above. The extra relay artefacts are clearly visible in fig. 4.45c, acquired using the ‘basic’ sequence in fig. 4.44a only. (As previously mentioned, these arise from large $^nJ_{\text{HH}}$ which evolve during the 2Δ spin echo; for this specific compound, the offending couplings are 3J between two axial protons in a six-membered ring, and 2J in diastereotopic methylenes). The presence of these peaks largely defeats the purpose of using a HSQC-COSY experiment; in fact, this unoptimised TSE HSQC-COSY is qualitatively very similar to a HSQC-TOCSY acquired with

^{*}A very similar strategy is used in the Bruker seHSQC pulse sequence hsqcedetgpsisp2.4 to suppress the COSY-type artefacts. However, to the best of my knowledge, this sequence has not been published anywhere.

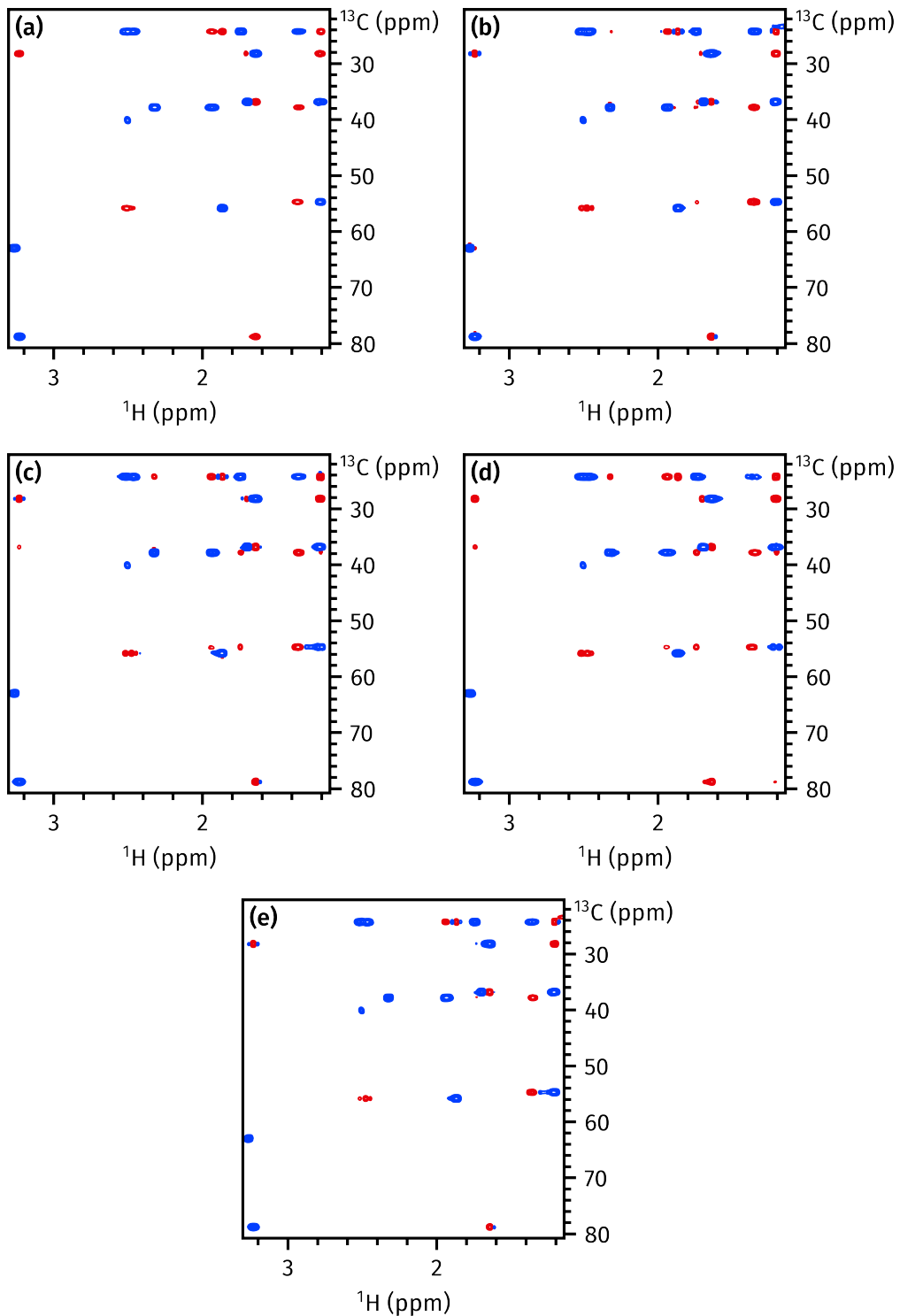


Figure 4.45: HSQC-COSY and HSQC-TOCSY spectra, taken from (respectively) NOAH- $^3\text{SC}^{\text{c}}$ and NOAH- $^3\text{STC}^{\text{c}}$ experiments. (a) HSQC-CLIP-COSY. (b) DSE HSQC-COSY. (c) TSE HSQC-COSY without suppression of relay artefacts (described in the text). (d) HSQC-TOCSY with 10 ms mixing time. (e) TSE HSQC-COSY with suppression of relay artefacts. Data code: 7A-210303.

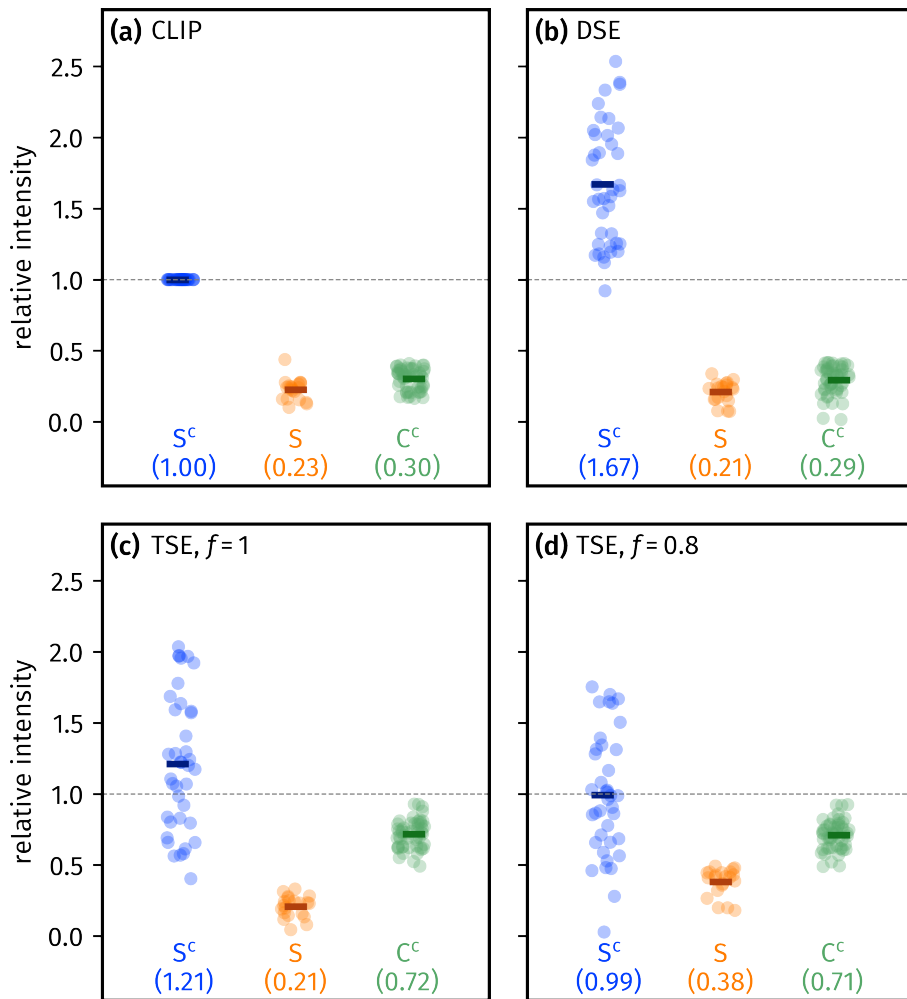


Figure 4.46: Sensitivity comparisons for all three modules in NOAH-3 S^CSC^c supersequences. Peak intensities are relative to the HSQC-CLIP-COSY module itself (the leftmost column in (a)), as well as HSQC and CLIP-COSY spectra from a NOAH-2 SC^c experiment. Numbers in parentheses indicate averages over all peaks. (a) HSQC-CLIP-COSY. (b) DSE HSQC-COSY. (c) TSE HSQC-COSY, acquired with $f = 1$. (d) TSE HSQC-COSY, acquired with $f = 0.8$. Data code: 7A-210723.

a short mixing time of 10 ms (fig. 4.45d).^{*} However, these artefacts can be efficiently removed using the suppression technique described in the text above; the result in fig. 4.45e is qualitatively similar to the two other HSQC-COSY experiments. Using this suppression technique, the sensitivity of the TSE HSQC-COSY (when acquired with $f = 1$) falls between that of the CLIP and DSE versions.

Figure 4.46 also shows the relative sensitivities of the later modules in NOAH-3 S^CSC^c supersequences. In all of the first three cases (CLIP, DSE, and TSE HSQC-COSY with $f = 1$, figs. 4.46a to 4.46c), the HSQC sensitivity is low (ca. 20%) because no $^1H^C$ magnetisation is retained for it

*The HSQC-TOCSY even gives better peak shapes, since the DIPSI mixing transfers in-phase magnetisation to in-phase magnetisation.

to use. Thus, the signal derives only from $^1\text{H}^{\text{C}}$ magnetisation which has recovered during the HSQC-COSY FID. However, when the TSE version is used, partial $^1\text{H}^{\text{C}}$ excitation can be used to control this sensitivity: for example, when $f = 0.8$ (fig. 4.46d), the HSQC-COSY sensitivity is decreased (on average equalling that of the HSQC-CLIP-COSY), but the sensitivity of the HSQC module is almost doubled. Generally, choosing a value of $f < 1$ allows for the HSQC-COSY and HSQC sensitivities to be better balanced.

Finally, the CLIP-COSY module suffers when the HSQC-CLIP-COSY or the DSE HSQC-COSY are used, because both of these dephase $^1\text{H}^{\text{C}}$ magnetisation; however, the TSE version successfully preserves around 70% of this magnetisation for it, regardless of the value of f . (This value is slightly lower than the approximately 90% magnetisation preserved by the HSQC module, because the bulk magnetisation is placed in the transverse plane during the 2τ spin echo (fig. 4.44a) and experiences losses due to ${}^nJ_{\text{HH}}$ evolution.)

HSQC-COSY in context

In NOAH-3 $\text{S}^{\text{C}}\text{SC}^{\text{c}}$ -type supersequences, using the TSE HSQC-COSY module here appears to be a sensible option as it is capable of preserving some $^1\text{H}^{\text{C}}$ magnetisation for the HSQC, as well as all $^1\text{H}^{\text{C}}$ magnetisation for the CLIP-COSY. However, this may not necessarily be so important in the context of a larger supersequence—particularly one which begins with the HMBC module, which *already* dephases $^1\text{H}^{\text{C}}$ magnetisation (meaning that there is not much of it to preserve).

Figure 4.47 provides the same sensitivity comparisons as in fig. 4.46, but in the context of a NOAH-4 $\text{BS}^{\text{C}}\text{SC}^{\text{c}}$ supersequence instead. The HSQC-COSY and HSQC modules largely follow the same pattern as before, but with an approximate 10% loss across the board: this reflects the imperfect preservation of $^1\text{H}^{\text{C}}$ magnetisation by the zz -HMBC module. The CLIP-COSY module, however, has a substantially lower sensitivity regardless of which HSQC-COSY module is chosen. When the CLIP or DSE HSQC-COSY modules are used, the CLIP-COSY retains only roughly 30% of its original intensity: this is the same as in fig. 4.46. With the TSE HSQC-COSY, this is boosted to around 40% because there is one extra FID in which the $^1\text{H}^{\text{C}}$ polarisation can be recovered. However, the use of the HMBC module at the beginning effectively puts a cap on the amount of signal available to this module.

For virtually all homonuclear modules (including the CLIP-COSY), this small difference in sensitivity will not make a real difference in the interpretability of the spectrum. This is especially so considering that the HMBC module—which has a far lower sensitivity—is also present in the supersequence: a CLIP-COSY with 30% of its original sensitivity is still more intense than the HMBC experiment. This argument was used in justifying the NOAH-3 BSC^{c} experiment, and logically, should be equally applicable to the NOAH-4 $\text{BS}^{\text{C}}\text{SC}^{\text{c}}$ experiment. In this case, the

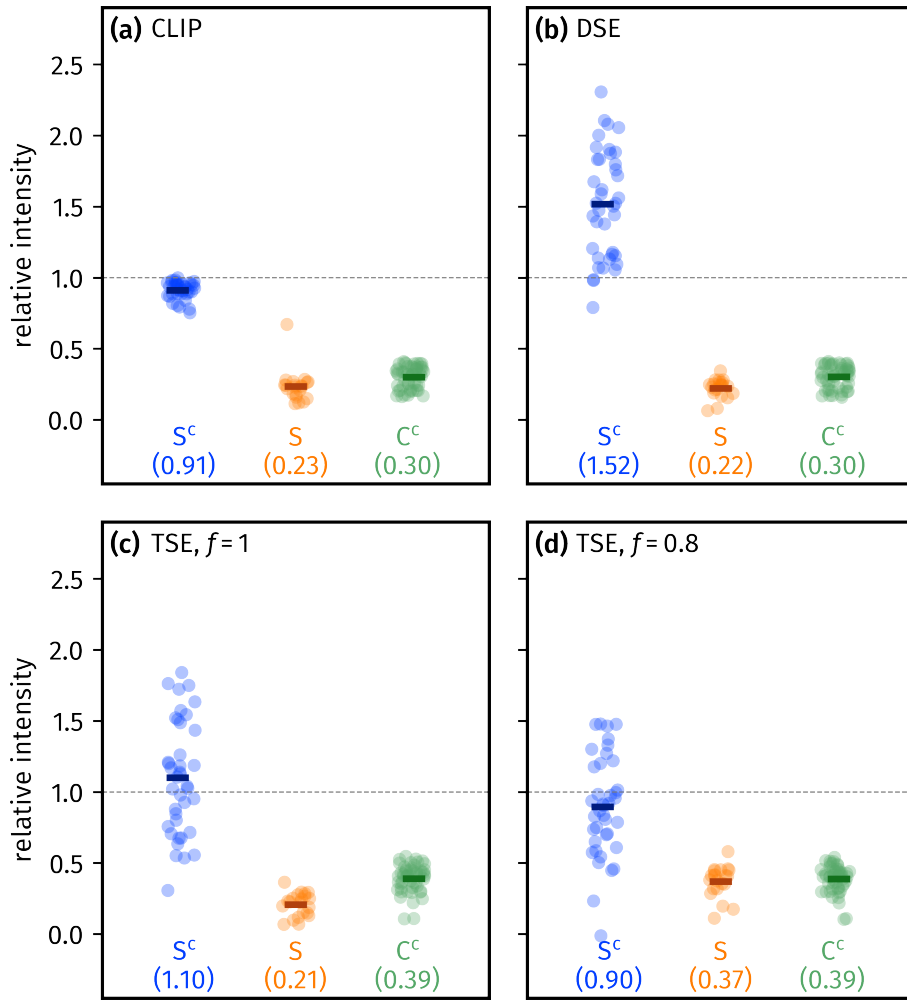


Figure 4.47: Sensitivity comparisons for the three last modules in NOAH-4 $\text{BS}^{\text{C}}\text{SC}^{\text{c}}$ supersequences. Peak intensities are relative to the HSQC-CLIP-COSY module from a NOAH-3 $\text{S}^{\text{C}}\text{SC}^{\text{c}}$ supersequence, and HSQC and CLIP-COSY spectra from a NOAH-2 SC^{c} experiment (i.e. the same reference spectra as in fig. 4.46). (a) HSQC-CLIP-COSY. (b) DSE HSQC-COSY. (c) TSE HSQC-COSY, acquired with $f = 1$. (d) TSE HSQC-COSY, acquired with $f = 0.8$. Data code: 7A-210723.

only compelling reason to use the TSE HSQC-COSY would be to preserve a portion of ${}^1\text{H}^{\text{C}}$ magnetisation for a later ${}^{13}\text{C}$ module. Thus, in this context, the decision of which HSQC-COSY module to use is slightly more nuanced: the cleaner lineshapes provided by the CLIP version, or the sensitivity of the DSE version, may be more preferable.

4.3.6 2DJ and PSYCHE

So far, I have exclusively used the CLIP-COSY as the final homonuclear module in NOAH supersequences. In general, since homonuclear modules are placed at the end of supersequences, there is rarely any need to adapt them for NOAH supersequences, as they do not need to preserve any magnetisation. One exception to this is the family of 2DJ and pseudo-2D pure shift

experiments (here typified by PSYCHE), where the spectral width in the indirect dimension is extremely small, on the order of 50 Hz. In such cases, the number of t_1 increments required is far smaller than for a typical 2D experiment. Since—by default—each module in a NOAH supersequence is acquired with the same number of increments, directly concatenating such modules to the end of a supersequence would therefore prove suboptimal.

However, as described previously in §§ 4.3.2 and 4.3.3, it is possible to reduce the number of t_1 increments for a particular module, and in exchange, increase the number of scans recorded for that module. In the context of ^{15}N modules, this was a ‘special’ procedure referred to as k -scaling; however, for these homonuclear modules, it is natural and necessary. One difference in the implementation is that, instead of specifying a value k by which TD1 is scaled down and NS scaled up, the user is allowed to directly specify the number of t_1 increments as an integer (CNST37 in TopSpin). This value must be a divisor of the *normal* number of t_1 increments for all other modules (or equivalently, in the context of ^{15}N modules, k must be an integer). The ‘indirect-dimension spectral width’ is specified as CNST38: this quantity is more properly referred to as the reciprocal of the chunk size, i.e. $1/T_{\text{chunk}}$.

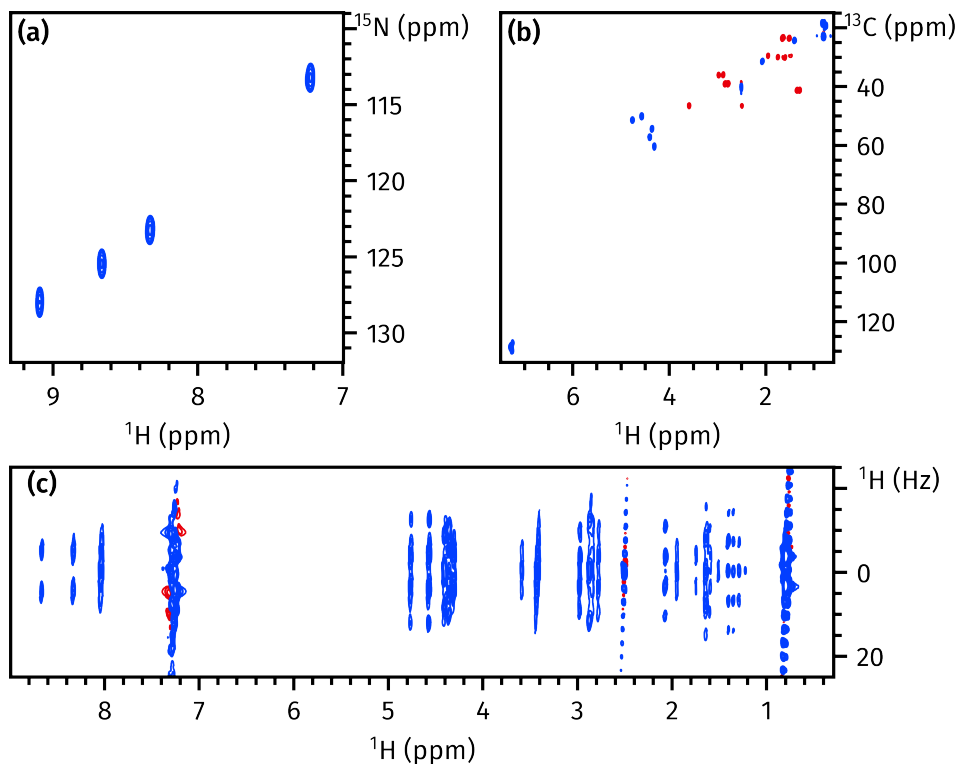


Figure 4.48: Spectra obtained from a NOAH-3 $\text{S}_\text{N}^+\text{S}^+\text{J}$ supersequence. (a) ^{15}N seHSQC (256 t_1 increments and 2 scans per increment). (b) ^{13}C seHSQC. (c) PSYCHE 2DJ (32 t_1 increments and 8 scans per increment). Data code: 7G-201028.

The modules thus implemented include the standard magnitude-mode 2DJ experiment, the PSYCHE 2DJ experiment,⁸⁹ the original 1D PSYCHE,⁹⁰ and the 1D TSE-PSYCHE.⁸⁹ An example

of the data thus obtained (with the PSYCHE 2DJ) is shown in fig. 4.48. One particular advantage of including PSYCHE-type modules in NOAH supersequences is that sensitivity is not likely to be at a premium: this is partly because of the increased number of scans, but also partly because other 2D experiments have comparably low sensitivity (meaning that in the time needed to acquire an HSQC, for example, the PSYCHE experiment will also have sufficient sensitivity). This allows the user to choose a relatively small flip angle (ca. 10°) for the PSYCHE saltire pulses in order to minimise artefacts from imperfect decoupling (see also § 2.2.3).

On top of this, for the 1D (TSE-)PSYCHE sequences, the extra transients can be used to perform SAPPHIRE averaging.⁹¹ Ordinarily, the PSYCHE pulse sequence seeks to refocus J-couplings in the middle of each chunk (of duration T_{chunk}). This is accomplished through the *prefocusing* of J-couplings in a spin echo of total duration $T_{\text{chunk}}/2$.⁹² In the SAPPHIRE procedure, each chunk of the pure shift interferogram is instead collected several times, all while varying the exact point in time where J-coupling is perfectly refocused. Summation of these data leads to the suppression of artefacts which arise due to the periodic J-modulation in the interferogram. This averaging is somewhat analogous to a phase cycle, and performing an 8-step SAPPHIRE averaging procedure (for example) would often require the experiment to be lengthened beyond the duration which is truly necessary. However, in the context of NOAH, since each chunk is acquired at least k times, a k -step SAPPHIRE procedure can be performed ‘for free’. In the GENESIS pulse programmes, this feature is enabled by default, although it can be turned off by selecting the original modules using developer mode.

A final and more prosaic implementation detail is that in the 1D pure shift modules, the chunk size T_{chunk} is automatically rounded to the nearest even multiple of the dwell time τ_{dw} (DW in TopSpin). This ensures that each chunk consists of an integral number of complex data points. Although this can be set by the user manually, it is very easy to forget, especially for someone not fully acquainted with the experiment; the results can be quite different, as illustrated in fig. 4.49. When an n -step SAPPHIRE averaging is used, the requirement is even stricter: T_{chunk} must be a multiple of $2n\tau_{\text{dw}}$. This is also encoded in the pulse programmes.

4.3.7 HMBC

The HMBC module is one which in fact does not fit perfectly into the NOAH principle of only exciting magnetisation which is needed. As described in § 4.1.3, the HMBC module should only require magnetisation of protons which have long-range couplings to ^{13}C ; however, it ends up exciting *all* $^1\text{H}^{13}\text{C}$ magnetisation. This leads to sharply reduced, and also unbalanced, intensities of homonuclear modules which come later in the supersequences.

I made some early (and brief) attempts at devising a pulse sequence which sought to discriminate

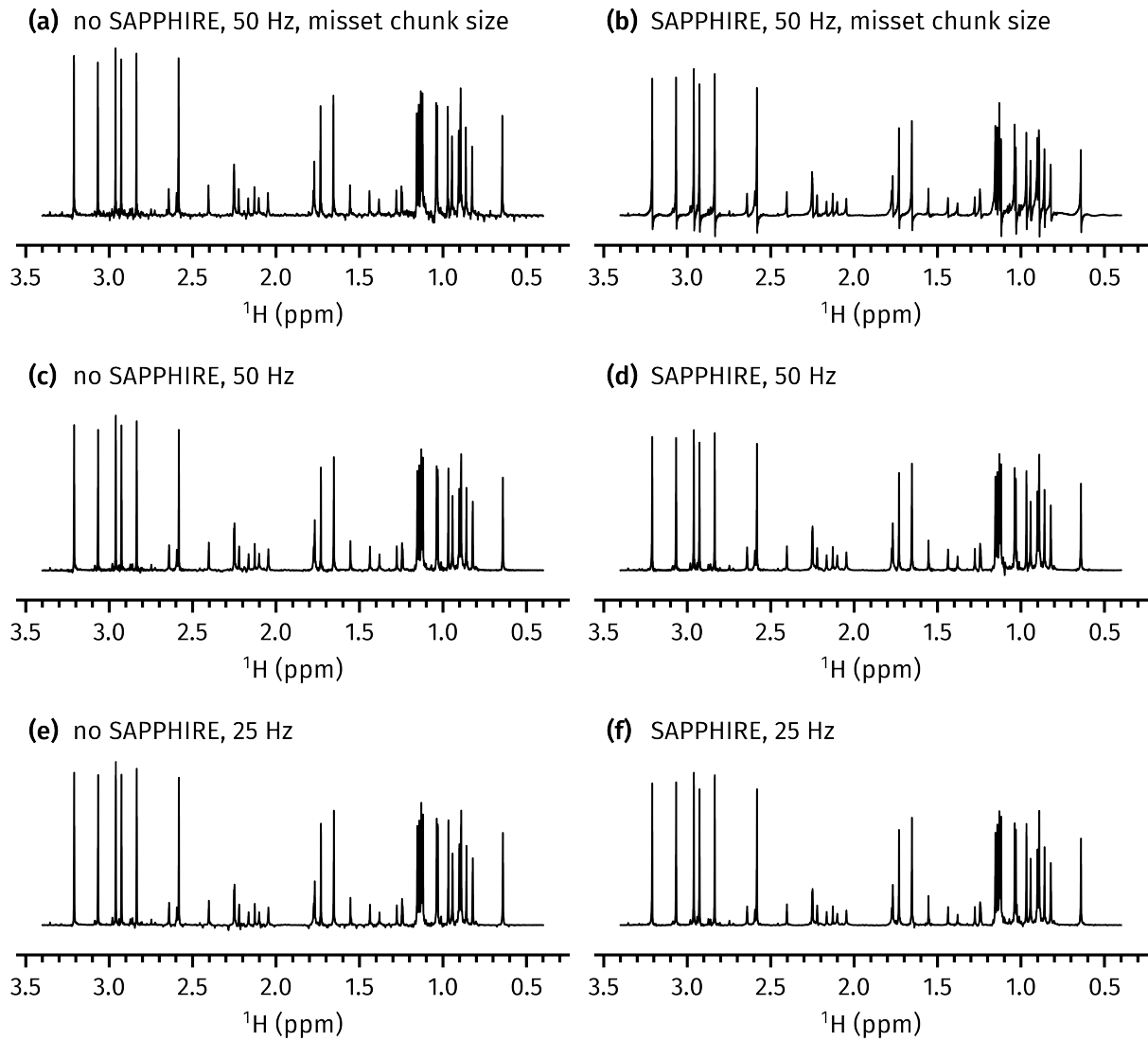


Figure 4.49: A series of 1D PSYCHE spectra obtained from the NOAH-3 $\text{S}_\text{N}^+\text{S}^+\text{P}$ super-sequence (saltire flip angle of 15°). (a) 1D PSYCHE spectrum acquired without SAPPHIRE averaging, and an incorrect chunk size which is not an integral number of complex data points. The ‘indirect-dimension spectral width’ (more precisely, $1/T_{\text{chunk}}$) is 50 Hz. (b) 1D PSYCHE spectrum with 8-step SAPPHIRE averaging and an incorrect chunk size; this manifests as phase errors which cannot be corrected. (c)–(d) The same as (a) and (b), but with the chunk size automatically corrected in the pulse programme. (e)–(f) The same as (c) and (d), but with double the chunk size. This leads to a more striking difference between the data acquired without and with SAPPHIRE, as discussed in the original paper.⁹¹ Data code: 7C-211123.

these two magnetisation components using a perfect echo.⁸⁷ However, this was quickly abandoned as it proved very difficult to *also* retain ${}^1\text{H}^{\text{C}}$ magnetisation. I do not claim here that it is impossible to come up with a pulse sequence which does this, but it is certainly not easy, and ultimately I turned my focus to improving (rather than replacing) the HMBC module.

Suppression of one-bond artefacts

One of the issues with the NOAH HMBC module was that there were an unusual amount of one-bond artefacts, which arise from ${}^1\text{H}^{\text{C}}$ magnetisation that is allowed to evolve during the pulse sequence. Generally, HMBC experiments seek to suppress this through the use of a low-pass J-filter (LPJF, see also § 3.4.7). The NOAH HMBC module *additionally* contains a zz -filter, which stores ${}^1\text{H}^{\text{C}}$ magnetisation along $+z$ before the LPJF. Thus, in theory, one-bond artefacts should be suppressed in the NOAH HMBC to an even greater extent.

However, this expectation is not borne out: in some cases, the NOAH HMBC has *more intense* one-bond artefacts when compared against a standard HMBC experiment (figs. 4.50a and 4.50c). Even performing a POISE optimisation of the LPJF delays, as described in § 3.4.7, did not lead to any reduction in artefacts. I hypothesised instead that these artefacts arose from imperfect manipulation of ${}^1\text{H}^{\text{C}}$ magnetisation by the zz -filter. In particular, any *antiphase* magnetisation (of the form $I_x S_z$ or $I_y S_z$) generated after the zz -filter would be reconverted into in-phase I_x or I_y terms, which would not be destroyed by the LPJF.* (The LPJF works based on the assumption that it needs to destroy in-phase magnetisation, which is true if the excitation element is just a ${}^1\text{H}$ 90° pulse, but is not necessarily applicable in the NOAH HMBC.)

Such antiphase terms can, however, be easily removed by the addition of a ${}^{13}\text{C}$ 90° pulse, which transforms them into a mixture of double- and zero-quantum terms: these are either unobservable, or can be efficiently dephased by CTP gradients. This technique is used in the CLIP-HSQC family of experiments,^{77,85} as well as the LPJF itself. In this case, we simply need to add an additional ${}^{13}\text{C}$ 90° pulse between the zz -filter and the LPJF: this pulse is highlighted in fig. 4.3b. This results in a striking reduction of the one-bond artefacts, as shown in fig. 4.50b. The suppression accomplished with the addition of this 90° pulse is superior to that in the standard HMBC (fig. 4.50c), and is comparable to a standard HMBC with a third-order LPJF (fig. 4.50d). Of course, the NOAH module—which by default uses a second-order LPJF—can also be ‘upgraded’ to use a third-order LPJF. In GENESIS pulse programmes, this can be done using the -DLP3 acquisition flag (although the results were not evaluated here).

Gradient selection schemes

Another point which was investigated (but bore less fruit) was the gradient scheme used for CTP selection. The NOAH module, as shown in fig. 4.3b, uses a ‘symmetric’ scheme where two gradients of equal amplitude surround the t_1 period: this encoding is later decoded by a third gradient just prior to acquisition. However, other choices exist: for example, the Bruker standard library HMBC (derived from the work of Cicero et al.⁹³) uses only two gradients which have

*It would be nice to back this up with simulations, but I did not have the time to run these.

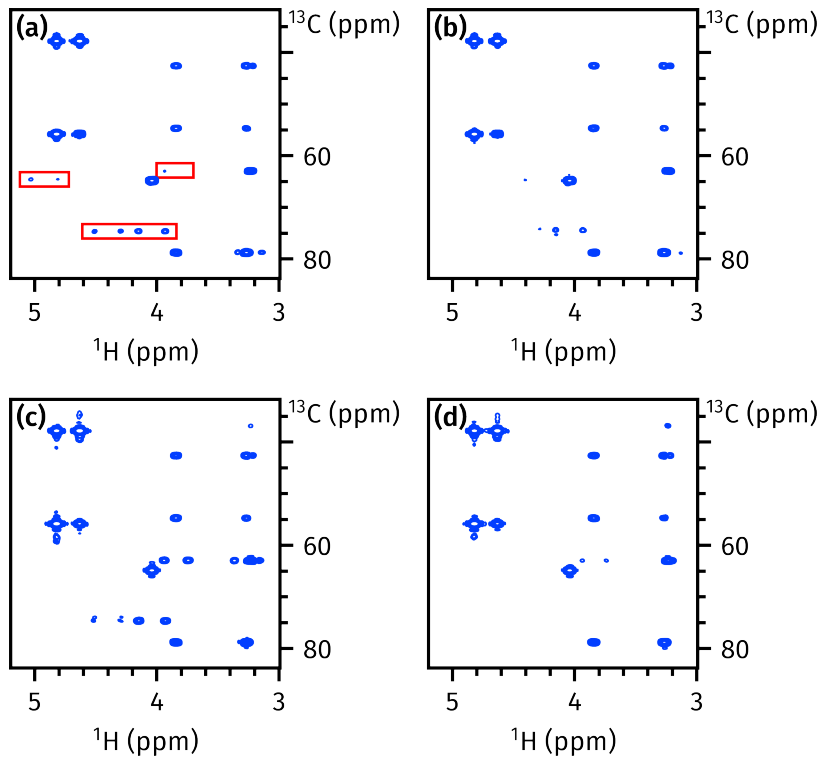


Figure 4.50: (a) NOAH zz -HMBC module without the additional 90° pulse. One-bond artefacts are highlighted in red. (b) NOAH zz -HMBC with the 90° pulse. (c) Standard library HMBC with a second-order LPJF. (d) Standard library HMBC with a third-order LPJF. Data code: 7A-210916.

unequal amplitudes. This gradient scheme cannot be directly used in a NOAH HMBC module, though. This is because the zz -filter element places $^1\text{H}^{\text{C}}$ magnetisation along the $+z$ axis just before the HMBC J-evolution delay (see fig. 4.3b). This magnetisation later experiences the ^1H 180° pulse in the middle of t_1 , which means that an extra 180° pulse must be added at the very end of the sequence to finally return it to $+z$. It is necessary to ensure that there is at least one gradient placed after this 180° pulse to ensure proper CTP selection (in the ‘symmetric’ scheme of fig. 4.3b, this is fulfilled by the gradient g_2).

Using this knowledge, it is possible to construct several ‘asymmetric’ gradient schemes:

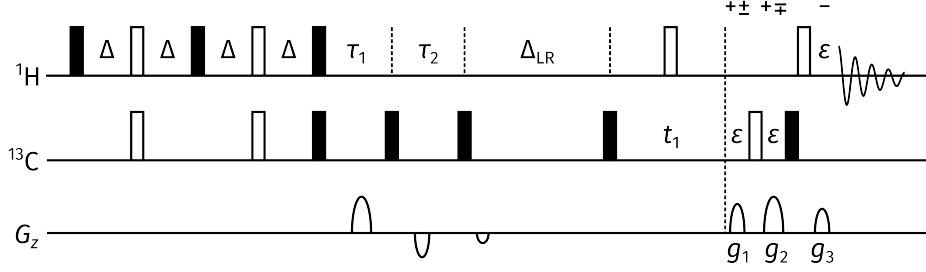
- ‘Scheme A’ (fig. 4.51a) is modified from the Bruker standard library to include a 180° pulse and gradient at the end. The presence of an additional gradient means that there is a free parameter, here denoted as α , which can be used to control the relative amplitudes of these three CTP gradients. The gradient amplitudes are chosen as follows:

$$\text{echo:} \quad g_1 = gc_1 \quad g_2 = g \quad g_3 = gc_2 \quad (4.18)$$

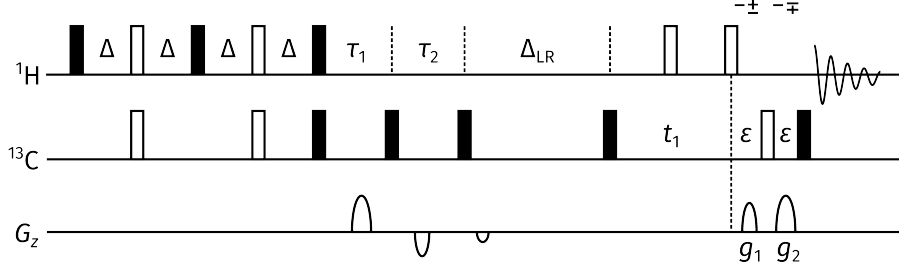
$$\text{antiecho:} \quad g_1 = g \quad g_2 = gc_1 \quad g_3 = gc_2 \quad (4.19)$$

where $c_1 = -\alpha(\gamma_{\text{H}} - \gamma_{\text{C}})/(\gamma_{\text{H}} + \gamma_{\text{C}})$ and $c_2 = (1 - \alpha)(\gamma_{\text{H}} - \gamma_{\text{C}})/\gamma_{\text{H}}$. In principle g is also a

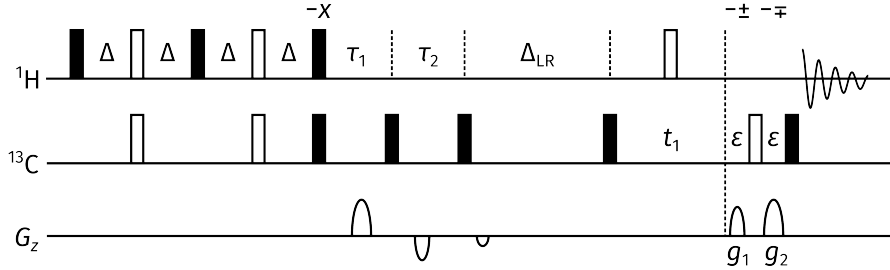
(a) Asymmetric scheme A



(b) Asymmetric scheme B



(c) Asymmetric scheme C



(d) Symmetric scheme (original)

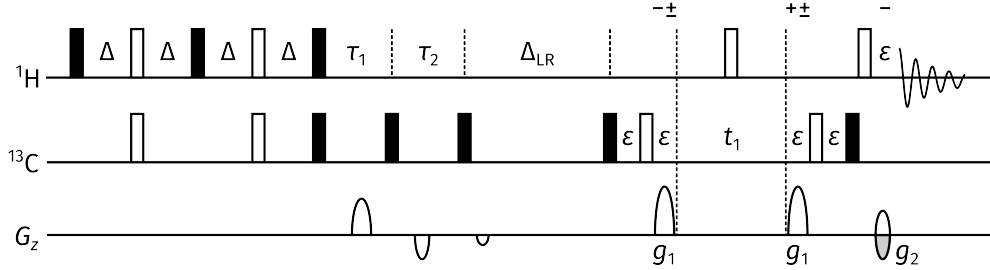


Figure 4.51: Alternative CTP gradient schemes investigated for the NOAH HMBC module. The coherences selected for during each gradient are indicated above each gradient, using the same notation for product operators as described in the *Preface*: the ‘upper’ term (e.g. + in ±) refers to the echo experiment, and the ‘lower’ term to the antiecho experiment. So, for example, +± refers to selection of I_+S_+ during the echo experiment and I_+S_- during the antiecho experiment. Gradient amplitudes are as described in the text. (a) ‘Asymmetric scheme A’, modified from the standard library sequence to include an additional 180° pulse and gradient. (b) ‘Asymmetric scheme B’, where the 180° pulse is shifted forward to the end of t_1 . (c) ‘Asymmetric scheme C’, which modifies the zz-filter instead of using an extra 180° pulse. (d) The original ‘symmetric’ scheme (the same as in fig. 4.3b), placed here for convenience.

free parameter; for maximum suppression of artefacts I chose a relatively large value of 80%.

2. In ‘Scheme B’ (fig. 4.51b), the 180° pulse is shifted to immediately after t_1 , before any of the CTP gradients have been applied. This means that there is no need for a third gradient, and the CTP gradient amplitudes can be directly taken from the standard library sequence:

$$\text{echo:} \quad g_1 = g \quad g_2 = gc \quad (4.20)$$

$$\text{antiecho:} \quad g_1 = gc \quad g_2 = g \quad (4.21)$$

where $c = -(\gamma_H - \gamma_C)/(\gamma_H + \gamma_C)$ and $g = 80\%$.

3. ‘Scheme C’ (fig. 4.51c) simply does not add a 180° pulse, but instead modifies the phases of the zz -filter in order to place $^1\text{H}^{\text{C}}$ magnetisation along the $-z$ axis during the HMBC J-evolution delay. Here, the gradient amplitudes are the same as those in the standard library sequence as well as in scheme B.

It is of interest to note two limiting cases of scheme A. When $\alpha = (\gamma_H + \gamma_C)/(\gamma_H - \gamma_C) \approx 1.67$, we obtain the ratio $g_1 : g_2 : g_3 = 1 : -1 : \pm 2\gamma_C/\gamma_H$, which mimics the original ‘symmetric’ scheme (fig. 4.51d); and when $\alpha = 1$, we have that $g_3 = 0$, i.e. a return to the two-gradient tactic of schemes B and C. In the tests which follow, I ran scheme A with $\alpha = 1.67$, $\alpha = 0.6$, and $\alpha = 0.3$.

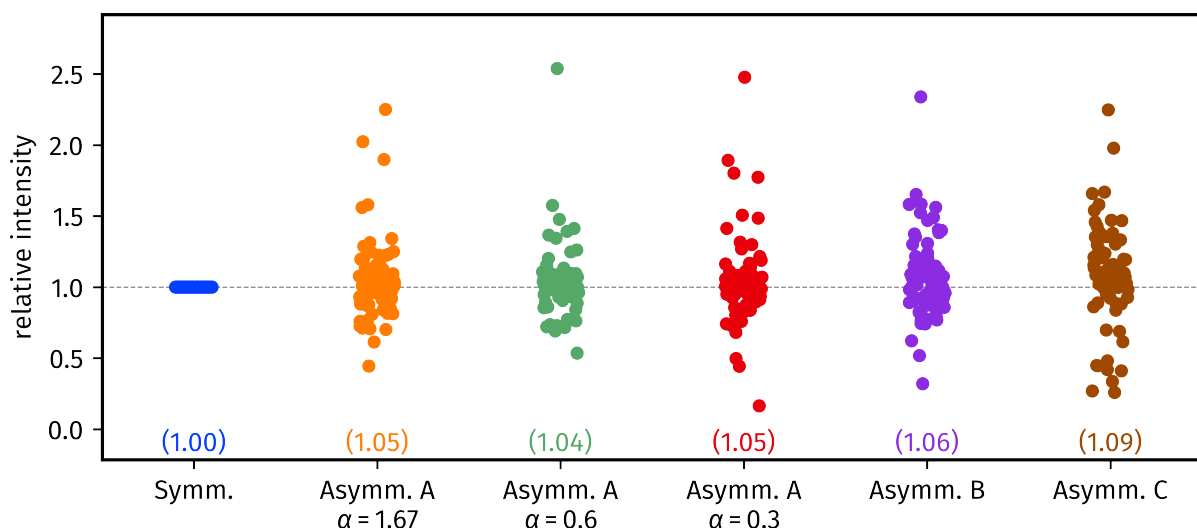


Figure 4.52: Sensitivities of various asymmetric HMBC gradient schemes, as compared to the symmetric scheme in fig. 4.3b. Each dot indicates one crosspeak in the HMBC spectrum; the numbers in parentheses are the average over all peaks. Data code: 7A-211226.

All of the different HMBC versions above, plus the original ‘symmetric’ scheme in fig. 4.3b, were tested in the context of a NOAH-2 BS supersequence using the andrographolide sample (fig. 4.52).

As can be seen, there is not much at all which separates the different versions (outliers with $> 2\times$ ‘sensitivity improvements’ can be attributed to different J-modulation in the multiplet). The most sensitive of these is asymmetric scheme C, which may be explained by the fact that it has one fewer 180° pulse: however, this comes with an immediate drawback. Since scheme C places the $^1\text{H}^C$ magnetisation along $-z$ during the LPJF as well as the J-evolution delay Δ_{LR} (a total of ca. 70 ms), relaxation losses during this period lead to poorer retention of $^1\text{H}^C$ magnetisation for later modules, as shown by the decreased HSQC sensitivities in fig. 4.53. In contrast, all the other gradient schemes retain $^1\text{H}^C$ magnetisation equally well.

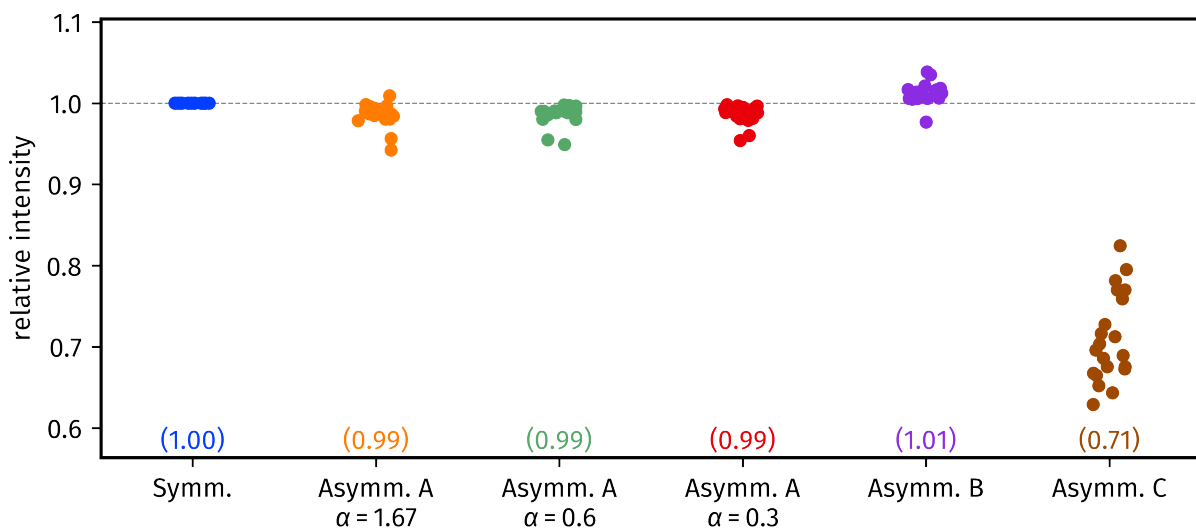


Figure 4.53: Sensitivities of the HSQC module in a NOAH-2 BS supersequence, where the HMBC module is implemented using the gradient schemes of fig. 4.52. Data code: 7A-211226.

The final point worth studying is the quality of the HMBC spectrum itself. To do this, we need to look at the actual spectra (fig. 4.54). For the most part, the spectra are all the same; however, there is a notable set of artefacts present in fig. 4.54b (scheme A with $\alpha = 1.67$) as well as fig. 4.54e (scheme B), highlighted in red boxes. These artefacts occur at the frequencies

$$(\Omega_1, \Omega_2) = \left(\Omega_S \pm \frac{\Omega_I}{2}, \Omega_I \right), \quad (4.22)$$

and are in fact ‘wing’ artefacts similar to that observed in other modules (§§ 4.3.1 to 4.3.3). In this case, they arise due to imperfect refocusing of the ^1H chemical shift during t_1 : specifically, whenever $I_z S_\pm$ terms are present during the second half of t_1 . Extra evidence for the origin of these artefacts comes from the observation that when the 180° pulse in the middle of t_1 is phase cycled, the artefacts are removed. In a standard HMBC, these terms would not be detected in the final FID; however, in this case, the addition of an extra 180° pulse after t_1 provides an opportunity for these to be converted back into observable spin- I magnetisation (through off-resonance effects or miscalibration).

The poorer performance may therefore be understood as follows: when scheme A is acquired with $\alpha = 1.67$, the gradients g_1 and g_2 have equal and opposite amplitudes, and so do not enforce any coherence selection on spin I during the second half of t_1 . Likewise, scheme B contains no gradients during the second half of t_1 .

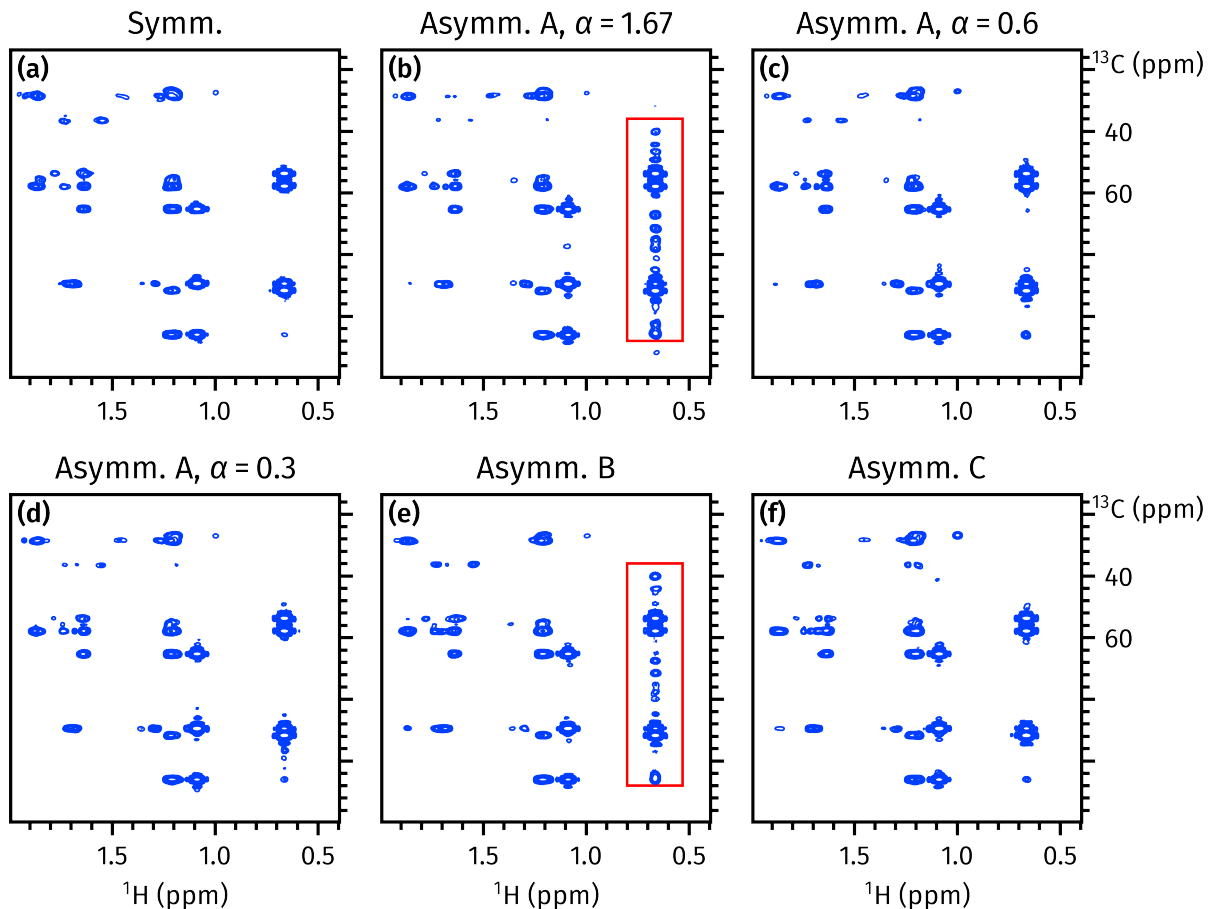


Figure 4.54: HMBC spectra acquired with the gradient schemes of fig. 4.52. Extra ‘wing’ artefacts present in two of the spectra (asymmetric scheme A with $\alpha = 1.67$, (b), and asymmetric scheme B, (e)) are highlighted in red boxes. Data code: 7A-211226.

The characteristics of these gradient schemes are summarised in table 4.4. As can be seen, the ‘best’ schemes are either the original symmetric scheme, or asymmetric scheme A with $\alpha \neq 1.67$. However, there is not much difference between these: it is not clear whether the improvement in sensitivity is reproducible across a wide range of samples, and in any case, the gains are extremely marginal.

Other artefacts

It has been established that the HMBC module (and supersequences containing it) are not fully ideal in terms of magnetisation preservation. However, there are also some other curious phenomena which have not been fully described in the literature. One of these is the presence

Gradient scheme	HMBC sensitivity	HSQC sensitivity	Wing artefacts
Symmetric	1	1	No
Asymmetric A, $\alpha = 1.67$	1.05	0.99	Yes
Asymmetric A, $\alpha = 0.6$	1.04	0.99	No
Asymmetric A, $\alpha = 0.3$	1.05	0.99	No
Asymmetric B	1.06	1.01	Yes
Asymmetric C	1.09	0.71	No

Table 4.4: Comparison of HMBC gradient schemes discussed in this section: the data are a summary of figs. 4.52 to 4.54. Data code: 7A-211226.

of *inverted peaks* in the homonuclear X module(s) in a NOAH-3 BSX supersequence: this is illustrated in fig. 4.55a with the CLIP-COSY module ($X = C^c$). It is not clear why this occurs, because the HMBC module (and the gradients which follow) should dephase all $^1H^{1C}$ magnetisation. Although this leads to reduced sensitivity in the homonuclear module, in that the signal derives from polarisation which has recovered during the preceding FIDs, it is not clear why this polarisation should be *negative*. One clue lies in the fact that these peaks are very sensitive to the 1H 90° pulse width: simply changing this by 0.5 μs is sufficient to restore the correct signal sign (fig. 4.55b).

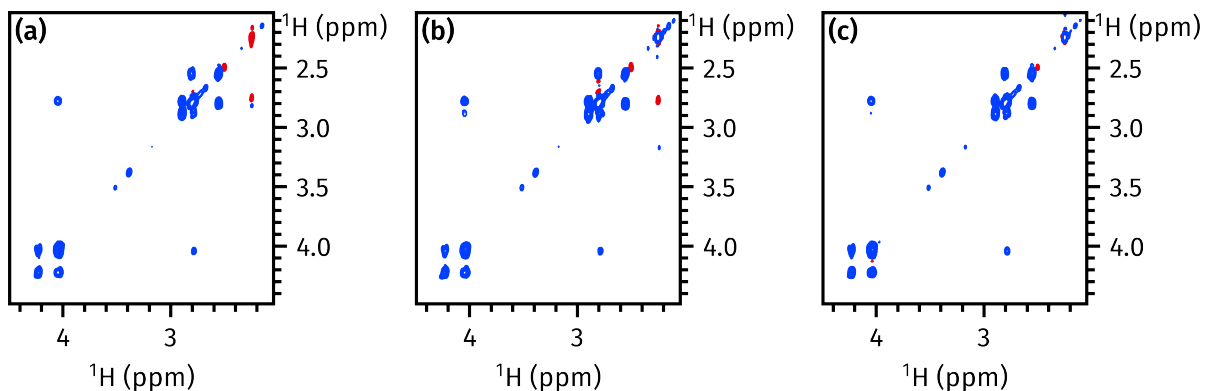


Figure 4.55: (a) CLIP-COSY from a NOAH-3 BSC^c supersequence, acquired with a 1H 90° pulse width of 11.28 μs (this value was obtained using the POISE calibration described in § 3.4.1). An inverted peak is visible at 2.25 ppm. (b) The same, but acquired using a 90° pulse width of 11.78 μs . (c) CLIP-COSY from a NOAH-4 BSSC^c supersequence. The 90° pulse width was 11.28 μs , the same as in (a). Data code: 7Z-220214.

The modules placed between the HMBC and the homonuclear module also play an important role. When *two* HSQC modules are used, i.e. a NOAH-4 BSSC^c supersequence (using $f = 0.7$ as described in § 4.3.4—although this is unlikely to matter), the negative peaks are no longer observed (fig. 4.55c). In fact, having *no modules* between the HMBC and the homonuclear module is also (at least sometimes) acceptable: a separate set of data shows that the inverted peaks in an NOAH-3 BS⁺C^c experiment are not seen in a NOAH-3 S⁺BC^c supersequence

(figs. 4.56a and 4.56b). The use of isotropic mixing (implemented as a sequence of adiabatic pulses,⁹⁴ and referred to as ‘ASAP’ mixing) just before the homonuclear module does not remedy this (figs. 4.56c and 4.56d). Unfortunately, a good explanation for these artefacts has remained elusive.

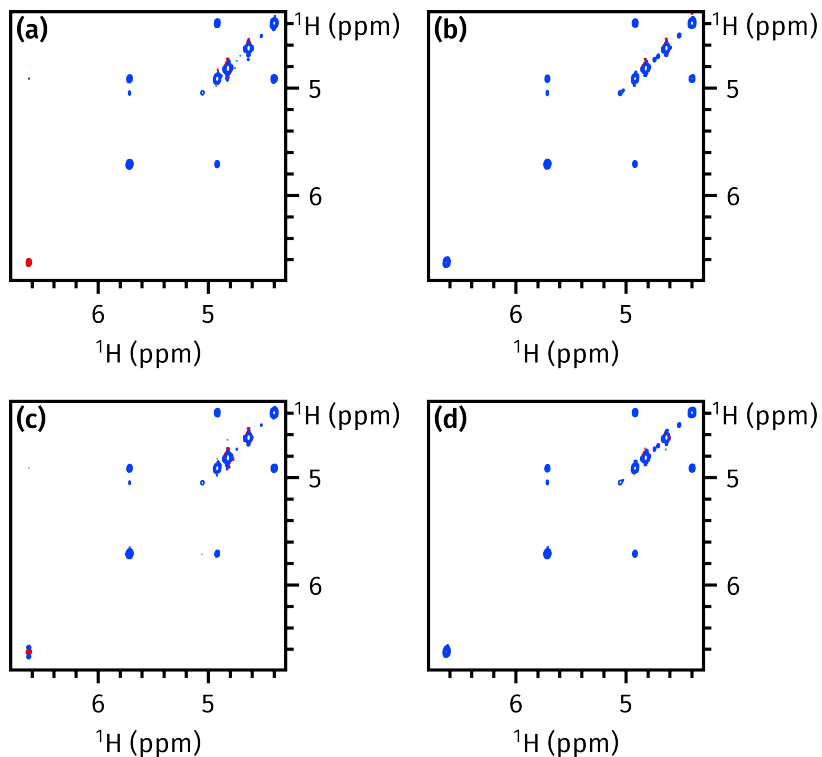


Figure 4.56: (a) CLIP-COSY from a NOAH-3 BS⁺C^c supersequence. An inverted diagonal peak can be seen at 6.6 ppm. (b) From a NOAH-3 S⁺BC^c supersequence. (c)–(d) The same as (a) and (b), but with 40 ms ASAP mixing placed just before the CLIP-COSY module. *Data code:* 7A-211227.

¹⁵N HMBC module

The entirety of this section has—until now—been devoted to the ¹³C HMBC module. However, the techniques used in constructing this, including the implementation of the *zz*-filter, are equally applicable to a ¹⁵N HMBC. For simplicity, the NOAH ¹⁵N HMBC module uses a first-order LPJF (since directly bonded NH pairs are less common); this may be omitted if desired. To minimise the number of pulses, a simple magnitude-mode version of the HMBC is used (fig. 4.57). The implementation of this module within supersequences is discussed in greater detail within the context of *generalised supersequences*, in § 4.5.

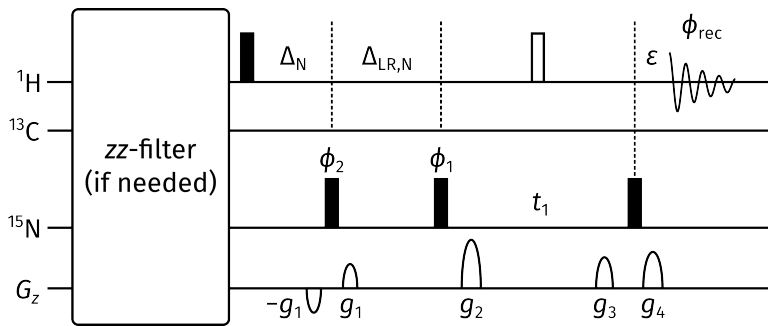


Figure 4.57: NOAH ^{15}N HMBC module. The zz -filter can be implemented as necessary in the same way as for the ^{13}C HMBC module (a final ^1H 180° pulse may also be required, but is not shown here). Delays are set as follows: $\Delta_N = 1/(2 \cdot ^1J_{\text{NH}})$; $\Delta_{\text{LR},N} = 1/(2 \cdot ^nJ_{\text{NH}})$. Phase cycling is performed using $\phi_1 = \phi_{\text{rec}} = (x, -x)$ and $\phi_2 = (x, x, -x, -x)$. Gradient amplitudes are $(g_1, g_2, g_3, g_4) = (5\%, 70\%, 30\%, 50.1\%)$.

4.3.8 ADEQUATE

The final ‘new’ module to be described in this section is the 1,1-ADEQUATE experiment,⁹⁵ which provides two-bond C–H correlations through a combination of $^1J_{\text{CH}}$ and $^1J_{\text{CC}}$. Because of the requirement for two adjacent ^{13}C nuclei, the sensitivity of the ADEQUATE experiment is considerably lower than almost all other NOAH modules considered so far. As established in § 4.1.1, sensitivity imbalances between different modules lead to less ideal supersequences with smaller effective time savings $\rho_{t,\text{eff}}$. Thus, the ADEQUATE module has to date not been implemented in NOAH supersequences.*

In this section, I only consider the design of the NOAH ADEQUATE module itself, which turns out to be extraordinarily simple. The ^1H pulses in the original ADEQUATE experiment mirror that of the CRK seHSQC almost perfectly, save for one extra 180° pulse. In particular, the INEPT excitation and the PEP transfer element are the same. Thus, the ADEQUATE module can be modified to preserve $^1\text{H}^{13}\text{C}$ magnetisation using exactly the same strategy in seHSQC2: namely, replacing the first ^1H excitation pulse with the ZIP element (fig. 4.58). In order to compensate for the extra 180° pulse (which arises in the constant-time period), the pulse phases in the ZIP element are modified accordingly: thus, the third ^1H 90° pulse has a phase of y (instead of $-y$ in seHSQC2).

An example of a NOAH-2 AB supersequence is shown in fig. 4.59. This is not the best setting to use the ADEQUATE module in, because the ^{13}C HMBC module has roughly 100 times the sensitivity of the ADEQUATE; however, it demonstrates that the ADEQUATE module does

*There is, in fact, one exception to my claim.⁹⁶ Here, the ADEQUATE and seHSQC modules were not modified to preserve any magnetisation for later modules. Since the ADEQUATE itself has such low sensitivity, it is likely that the sensitivity losses in the later modules were tolerable; however, no sensitivity comparisons against standalone experiments were provided.

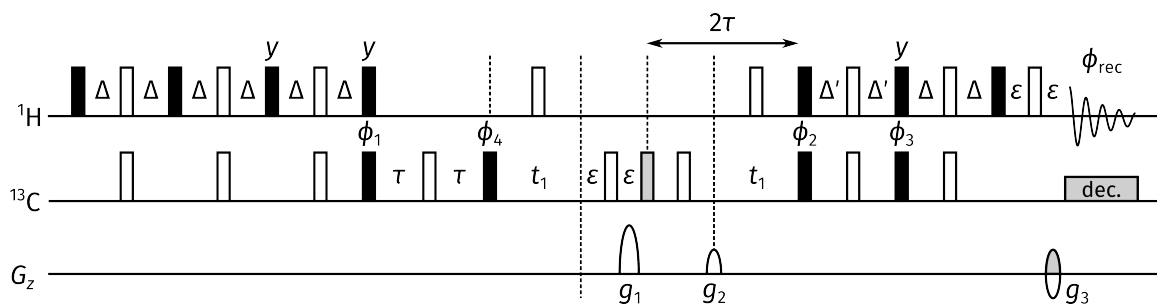


Figure 4.58: NOAH 1,1-ADEQUATE module. The grey filled bar is a 120° pulse, optimised for ^{13}C double-quantum to single-quantum coherence transfer.⁹⁷ Delays are set as follows: $\Delta = 1/(4 \cdot ^1\text{J}_{\text{CH}})$, $\Delta' = 1/(8 \cdot ^1\text{J}_{\text{CH}})$, and $\tau = 1/(4 \cdot ^1\text{J}_{\text{CC}})$. Phase cycling is performed using $\phi_1 = (x, -x)$, $\phi_2 = (x, x, -x, -x)$, $\phi_3 = (y, y, -y, -y)$, $\phi_4 = (x, x, x, x, -x, -x, -x, -x)$, and $\phi_{\text{rec}} = (x, -x, -x, x, -x, x, x, -x)$. Gradient amplitudes are $(g_1, g_2, g_3) = (78.5\%, 77.6\%, \mp 59\%)$. Echo-antiecho selection is achieved by inverting the sign of g_3 as well as the pulse phase ϕ_3 .

indeed work as intended. More constructive uses of the ADEQUATE module are deferred to § 4.5: in there, I discuss combinations with the (lower-sensitivity) ^{15}N HMBC module, and show how other modules may be added to form so-called ‘generalised’ supersequences.

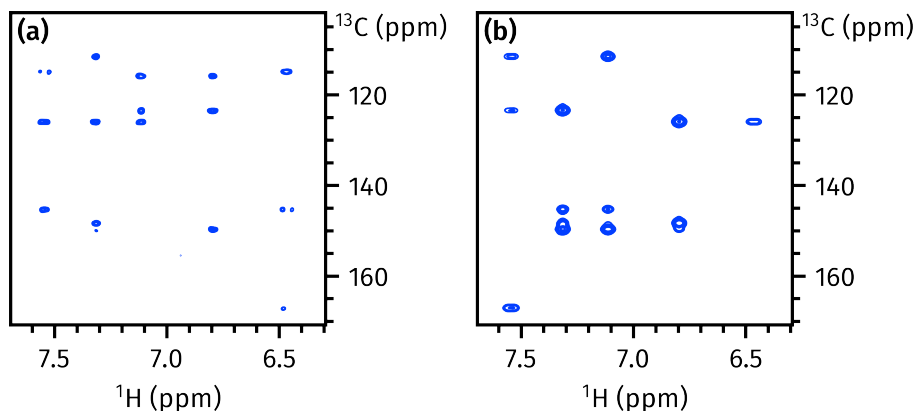


Figure 4.59: Spectra from a NOAH-2 AB supersequence. (a) 1,1-ADEQUATE module. (a) HMBC module. Data code: 7T-211231.

4.4 Solvent suppression in NOAH

Solvent suppression is one of the most important aspects of modern NMR, and has been incorporated into a large number of experiments. It is not a particularly difficult task to implement some basic solvent suppression techniques in NOAH supersequences, as will be described in this section. All suppression techniques shown here can be enabled via acquisition flags (the TopSpin ZGOPTNS parameter); this means that there is no need to switch pulse sequences.

4.4.1 Presaturation

A simple first step is to implement simple presaturation of the solvent resonance during the recovery delay of the supersequence. This often provides excellent suppression in the first module in a supersequence, and to a lesser extent, later modules (as the solvent magnetisation recovers during acquisition periods).

Presaturation is also included during long delays, in particular mixing times in NOESY experiments. The use of presaturation during the HMBC J-evolution delay was also tested, but was found to be unnecessary, especially since the HMBC module is typically placed first in supersequences.

4.4.2 Intrinsic suppression

A number of NOAH modules in fact come with an *intrinsic* form of solvent suppression, in that they return $^1\text{H}^{\text{IX}}$ magnetisation (including that of solvents) to the $+z$ axis at the end of the module. This is generally true of many of the HSQC-based NOAH modules, which seek to only sample $^1\text{H}^{\text{X}}$ magnetisation pools ($\text{X} = \text{C}$ or N). Thus, these modules already have far better solvent suppression properties compared to standard library sequences: there is no need for further modification.

For these sequences, it is possible to go one step further and to see how the solvent suppression varies with transmitter offset. This is less relevant if the solvent is just water (where it suffices to put the water peak on resonance), but may be important for samples in protonated solvents or mixtures thereof. I tested this by recording the first increment of the NOAH HSQC and seHSQC modules on an aqueous sample of sucrose, and integrating the water peak. The results (fig. 4.60) indicate that the HSQC module provides the best suppression across a range of offsets; of the two seHSQC modules, seHSQC1 provides more uniform suppression, whereas seHSQC2 has some regions ('spikes') where suppression is poorer than average.

It is not surprising that the more complicated seHSQC sequences have slightly poorer suppression: there are substantially more pulses involved which lead to cumulative errors, especially for peaks which are further away from resonance. However, it is not entirely clear why seHSQC2 has such large spikes. Some further investigation showed that the poorer suppression partly arises from imperfect cancellation of the water signal by phase cycling. This is, however, not a full explanation; it merely replaces one mystery with another. Nevertheless, it should be reiterated that all of these sequences still provide excellent water suppression. The standard library seHSQC sequences, for example, do not even come close to this level of suppression: although these do dephase water magnetisation prior to acquisition, radiation damping during the FID still leads to a very large water signal.

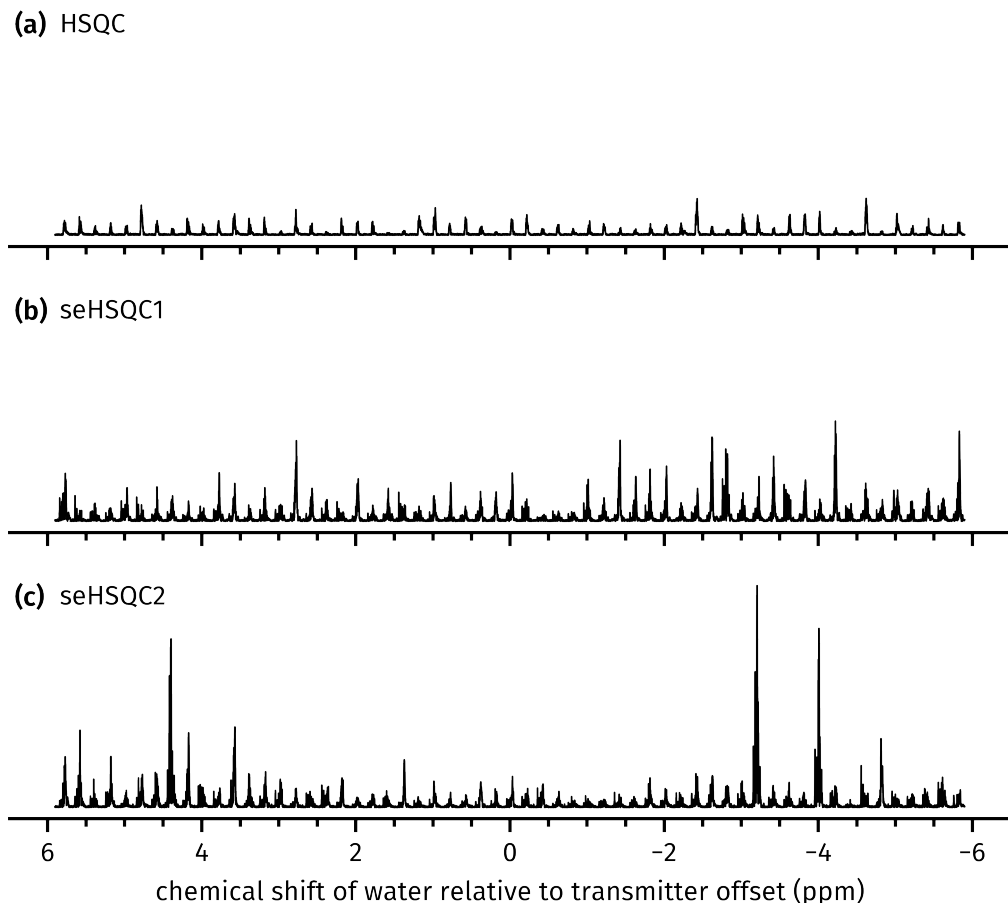


Figure 4.60: Residual water peaks observed in the first increment of NOAH HSQC and seHSQC modules, as a function of transmitter offset. The *y*-axes of the three subplots are the same. (a) HSQC module. (b) seHSQC1 module. (c) seHSQC2 module. *Data code:* 7S-201014.

4.4.3 Excitation sculpting

Since solvent suppression in HMBC and HSQC-type modules can be adequately accomplished through presaturation and intrinsic suppression, it remains to consider the suppression in homonuclear modules. These modules invariably occur at the end of supersequences, and thus almost any form of solvent suppression can be used: there is no need to consider how magnetisation needs to be preserved for other modules.

In practice, I chose to implement excitation sculpting⁹⁸ (ES) just prior to acquisition: the refocusing element chosen was a combination of a selective 180° sinc pulse and a hard 180° pulse. This worked perfectly for almost all of the homonuclear modules used in NOAH supersequences, with only a few adjustments needed, such as for the PSYCHE modules where it was necessary to ensure that chemical shifts and J-couplings evolved for the correct amount of time.

The case of the ‘double’ COSY + X homonuclear modules, however, proved to be more intricate.

This is illustrated here using $X = \text{TOCSY}$, but the considerations below are equally applicable to $X = \text{NOESY}$ or ROESY . It is tempting to simply place ES blocks prior to both FIDs: in the COSY module, this dephases transverse solvent magnetisation as desired, and (in principle) should leave longitudinal magnetisation untouched, meaning that the TOCSY module which later consumes this should be unaffected.

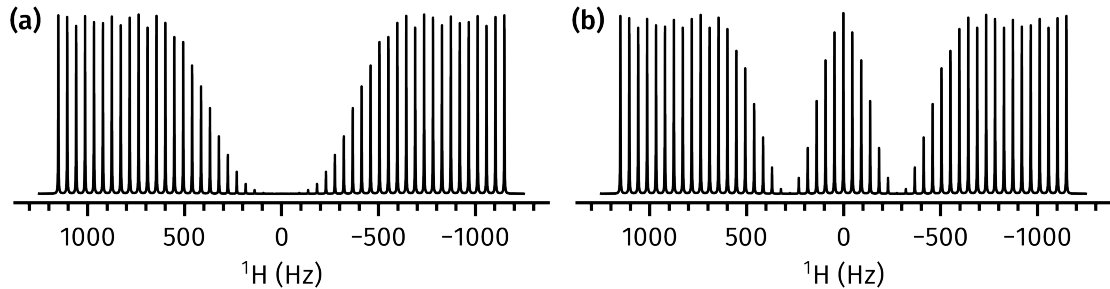


Figure 4.61: Simulations showing the proportion of magnetisation retained by the excitation sculpting block (using a 2 ms sinc pulse). (a) Retention of transverse magnetisation: this was obtained by simulating the spectrum of a 90° -ES-detect pulse sequence. (b) Retention of longitudinal magnetisation: this was obtained using an ES- 90° -detect pulse sequence.

However, this is not totally true. *Inside* the bandwidth of the selective 180° pulse, any longitudinal magnetisation experiences an 720° rotation; and *outside* of the bandwidth, it experiences a 360° rotation. However, *between* these two extremes, there is a crossover point where longitudinal magnetisation is sent into the transverse plane and subsequently dephased by gradients. This is reflected in simulations of the ES element, specifically the nulls in fig. 4.61b: peaks which fall within this range will be lost in the TOCSY module. This is visible in the spectra of fig. 4.62: although the water suppression obtained using this ‘double ES’ scheme is better, several peaks in the TOCSY spectrum have disappeared, because they fall precisely into these nulls. It proves better to omit the ES in the COSY module: an adequate degree of water suppression in the COSY can still be attained thanks to the use of presaturation.

Of course, it may well be that *no* peaks fall within this null, and thus the better solvent suppression may be obtained at no cost. This was the case when the experiments in fig. 4.62 were reacquired on a 700 MHz spectrometer; or when the sinc pulse was lengthened to 4 ms. However, such a coincidence cannot always be relied on.

4.5 Parallel and generalised NOAH supersequences

To conclude this chapter, I discuss how *multiple* NOAH supersequences may be run in parallel in order to obtain yet more data from a single experiment. As of the time of writing, current software limitations in TopSpin limit the NBL parameter (the number of memory blocks) to 5. This makes

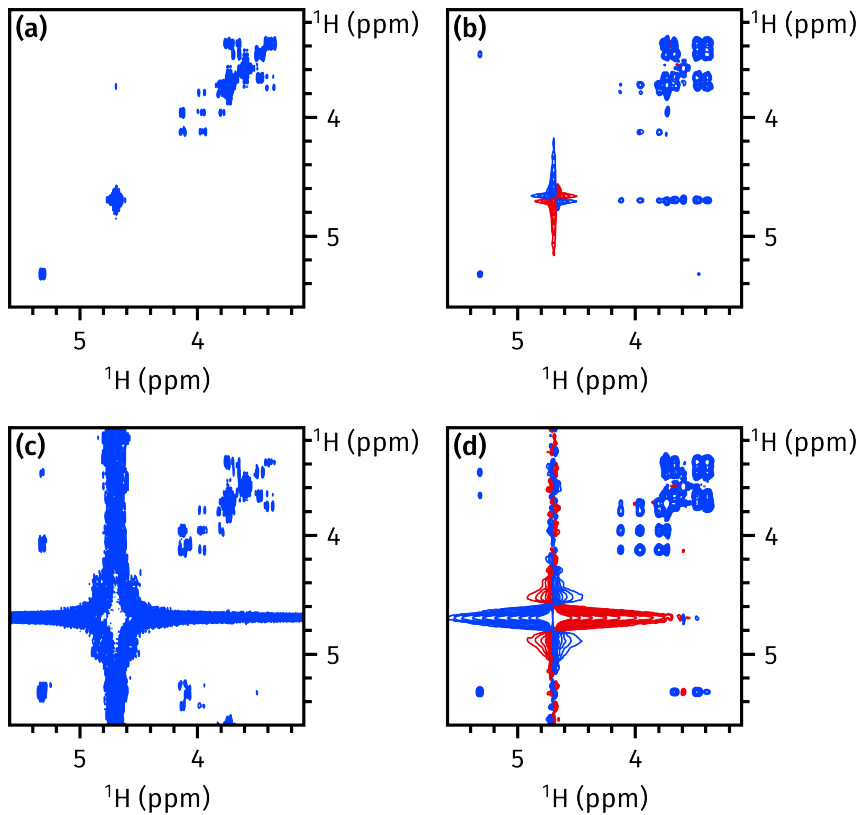


Figure 4.62: (a)–(b) COSY and TOCSY spectra obtained from a NOAH-3 SCT supersequence where excitation sculpting was placed before both the COSY and TOCSY FIDs. Although the water suppression is better, some peaks in the TOCSY module are lost. (c)–(d) The same, except that excitation sculpting was applied only in the TOCSY module. For all spectra shown here, presaturation of the water resonance was applied during the recovery delay. *Data code:* 4S-211105.

it presently impossible to linearly extend a supersequence beyond five modules. However, it is possible to *interleave* different supersequences, such that one t_1 increment of supersequence A is acquired, followed by one t_1 increment of supersequence B, before the value of t_1 is increased for both supersequences. In this text, I refer to this as a ‘vertical’ stacking of modules (as opposed to the traditional NOAH concept, which focuses on ‘horizontal’ concatenation of modules).

4.5.1 Parallel NOAH supersequences

This concept is more clearly illustrated in fig. 4.63, which contains a more explicit depiction of how t_1 is incremented for each module. In fig. 4.63a, a ‘traditional’ NOAH-2 ST supersequence is shown: here, t_1 is incremented for both the HSQC and TOCSY module at the same time, and on each t_1 increment, the sequence of modules being acquired is always the same. In fig. 4.63b, this is different: the second module is alternated between a TOCSY and a NOESY. If the total experimental duration (which is largely proportional to the number of recovery delays, d_1) is to be kept the same as before, this means that the TOCSY and NOESY modules will be recorded

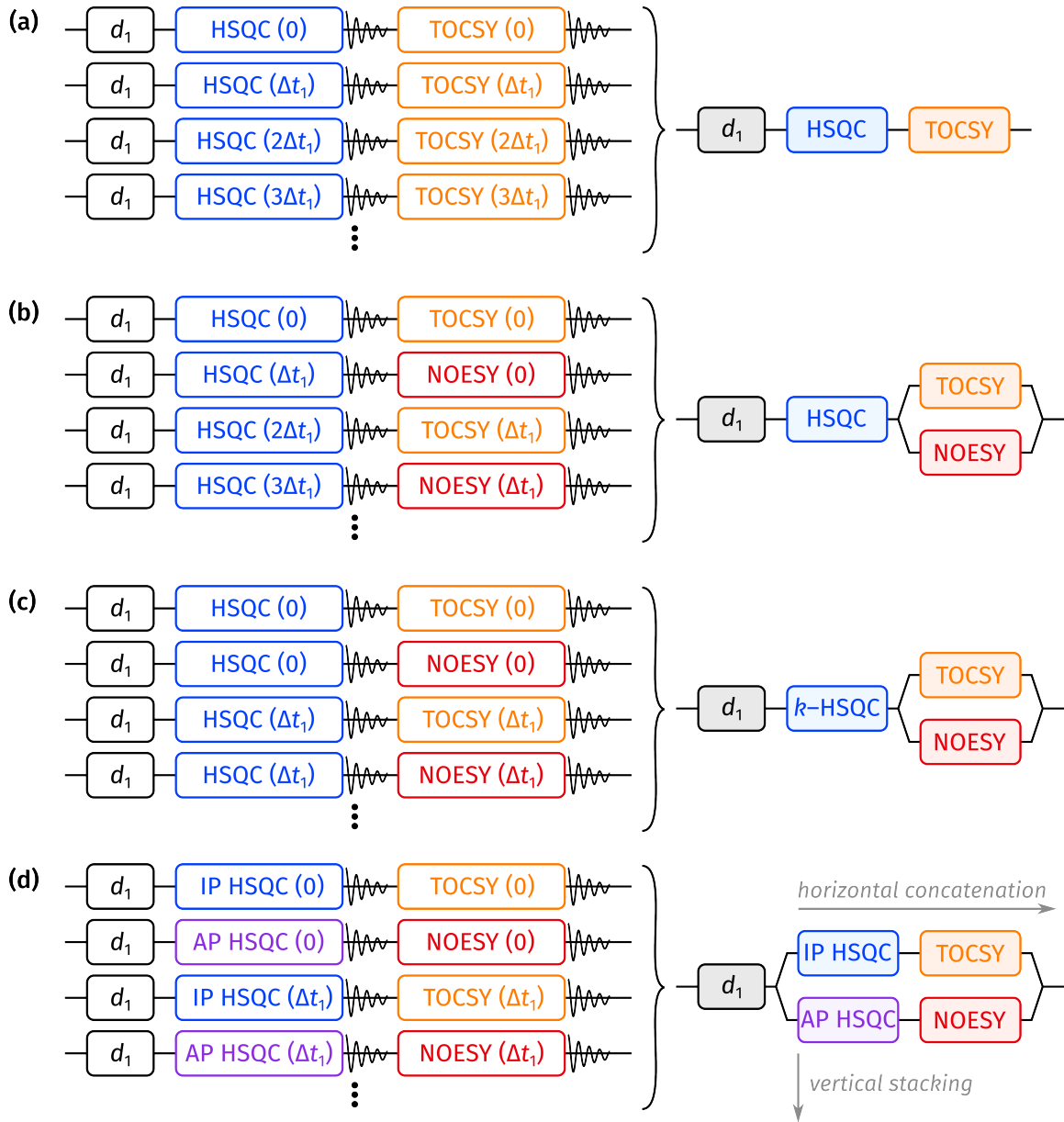


Figure 4.63: An overview of ‘parallel’ NOAH supersequences. The left side of each diagram explicitly spells out how t_1 is incremented for each module: the value in parentheses indicates the current value of t_1 , which begins at 0 and is incremented by Δt_1 each time. The right side is a ‘condensed’ depiction of the entire supersequence. (a) A ‘standard’ NOAH supersequence, where t_1 for every module is incremented at the same time. (b) The first example of a ‘parallel’ supersequence: the HSQC module is acquired as normal, but the TOCSY and NOESY modules are interleaved in the second slot. (c) The same as in (b), but the HSQC module is subjected to k -scaling (see also § 4.3.2). (d) A fully parallel supersequence, where both the first and second module slots are varied systematically. This amounts to the interleaved acquisition of two different ‘standard’ supersequences.

with half the number of t_1 increments each. Although this loss of resolution may be considered a drawback, this scheme does allow us greater *flexibility* in the design of NOAH supersequences: it is not ideal to acquire both a TOCSY and a NOESY using a traditional ‘linear’ supersequence, since both of these modules depend on $^1\text{H}^{\text{IX}}$ magnetisation.

Figure 4.63c shows how the F_1 resolutions for all three modules can be equalised, by performing the equivalent of k -scaling on the HSQC module (described in § 4.3.2). This entails reducing the number of t_1 increments by a factor of 2 (in this case), and acquiring each increment twice instead, which corresponds to a doubling of the number of scans after the data have been appropriately combined. This generally has very little impact on the spectra (as was previously shown), but provides a convenient lead into the final example of fig. 4.63d. Here, instead of acquiring each increment of the HSQC module twice, the time is used to acquire two different modules, an in-phase (IP) and antiphase (AP) HSQC (both run without ^{13}C decoupling). Through appropriate linear combination of the data, it is possible to isolate the two peaks of the $^1\text{H}-^{13}\text{C}$ doublets and measure $^1J_{\text{CH}}$ values: this is known as in-phase/antiphase (IPAP) processing.^{77,99–101} Much like the TOCSY and NOESY combination, it is not possible to acquire IP and AP HSQC spectra in a single, linear supersequence, not even with the partial $^1\text{H}^{\text{C}}$ excitation technique described in § 4.3.4. For the IPAP processing to work, each doublet has to have the same intensity in the IP and AP spectra. A careful choice of f in a linear supersequence may lead to the same *average* intensity, but the distribution of intensities amongst the peaks will not be the same.

The reader will no doubt notice at this point that fig. 4.63d corresponds simply to the interleaved acquisition of two different supersequences (one is IP-HSQC + TOCSY, and the other AP-HSQC + NOESY), which could just as well be acquired separately. Furthermore, both supersequences are acquired with only half of the original F_1 resolution (assuming the same experimental time as fig. 4.63a), which means that there are *no real time savings*. This is indeed correct: fundamentally, ‘vertical’ stacking does not increase the number of FIDs recorded per recovery delay, so does not improve the time efficiency (or ρ_t). The main benefits of this process are, in my opinion:

- the *flexibility* to construct supersequences that combine previously incompatible modules (such as in the examples above); and
- a way of *adjusting for relative sensitivities of different modules*. For example, the arrangement in fig. 4.63c amounts to the acquisition of the HSQC module for $2n$ scans, and the TOCSY and NOESY modules for n scans each (where n is some positive integer). In this way, less intrinsically sensitive modules can be assigned a larger number of scans, such that all modules in the supersequence have (relatively) equalised intensities.

In principle, all of the above may still be accomplished in a roundabout manner by acquiring multiple separate supersequences and then combining the results. This is most obvious in

fig. 4.63d. However, the current implementation of parallel supersequences provides *convenience* for the user, as all experiments may be simultaneously acquired and processed. This factor should not be overlooked, especially considering that all the different constructions (figs. 4.63b to 4.63d) must be processed in a slightly different manner: it is easier to do this from within a single experiment, rather than attempting to combine FIDs from separate datasets after the fact. There is also another argument for acquiring increments in an interleaved manner: this helps to minimise drifts in the spectra which arise from environmental instabilities such as temperature fluctuations.

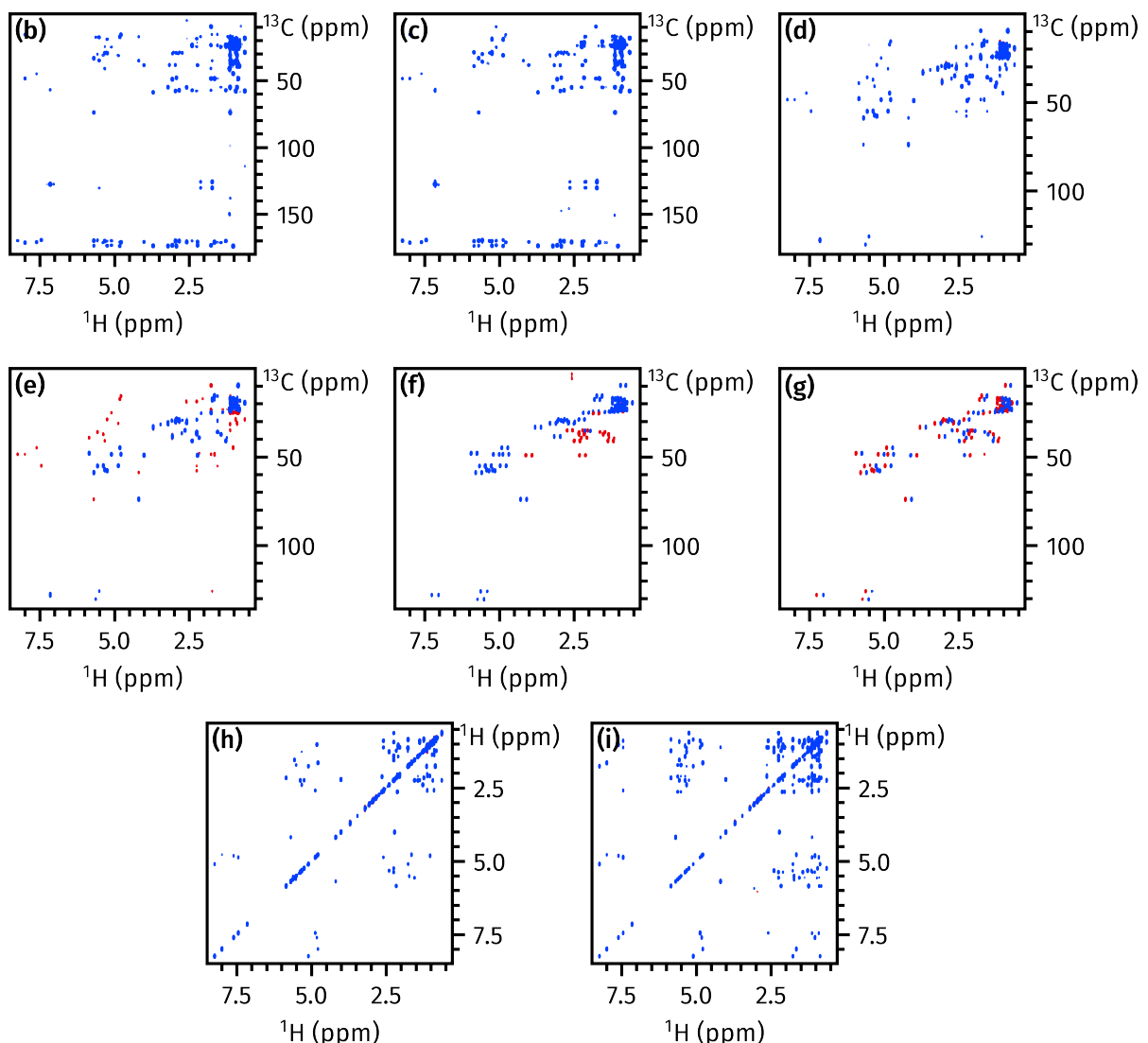


Figure 4.64: (a) Condensed representation of the supersequence, which may be interpreted in the same way as fig. 4.63d. (b) HMBC, optimised for ${}^nJ_{CH} = 5$ Hz. (c) HMBC, optimised for ${}^nJ_{CH} = 10$ Hz. (d) IP HSQC-CLIP-COSY. (e) AP HSQC-CLIP-COSY. (g) IP multiplicity-edited seHSQC. (g) AP multiplicity-edited seHSQC. (h) CLIP-COSY. (i) TOCSY (60 ms mixing time).

Figure 4.64a shows one example of a more complex parallel supersequence containing 8 modules.

In this supersequence, the HMBC module is recorded twice with the delay Δ_{LR} optimised for two different values of ${}^nJ_{CH}$ (figs. 4.64b and 4.64c): these spectra furnish a (slightly) different set of correlations, which allow for more complete structure determination. The HSQC-CLIP-COSY is acquired in an IPAP fashion, where direct/indirect editing is used in one experiment (fig. 4.64e) and not in the other (fig. 4.64d). Likewise, the F_2 -coupled seHSQC experiment is also performed in an IPAP manner, where this time the IPAP components are the two peaks in each doublet. Multiplicity editing is additionally used here in both IP and AP seHSQC spectra. Finally, we have two interleaved homonuclear modules, the CLIP-COSY and TOCSY (figs. 4.64h and 4.64i), which provide complementary information about spin networks. In fig. 4.64, the heteronuclear spectra are shown before IPAP processing (in other words, these are the IP and AP spectra). The four IPAP-processed spectra (IP – AP and IP + AP) from this supersequence are shown in fig. 4.65.

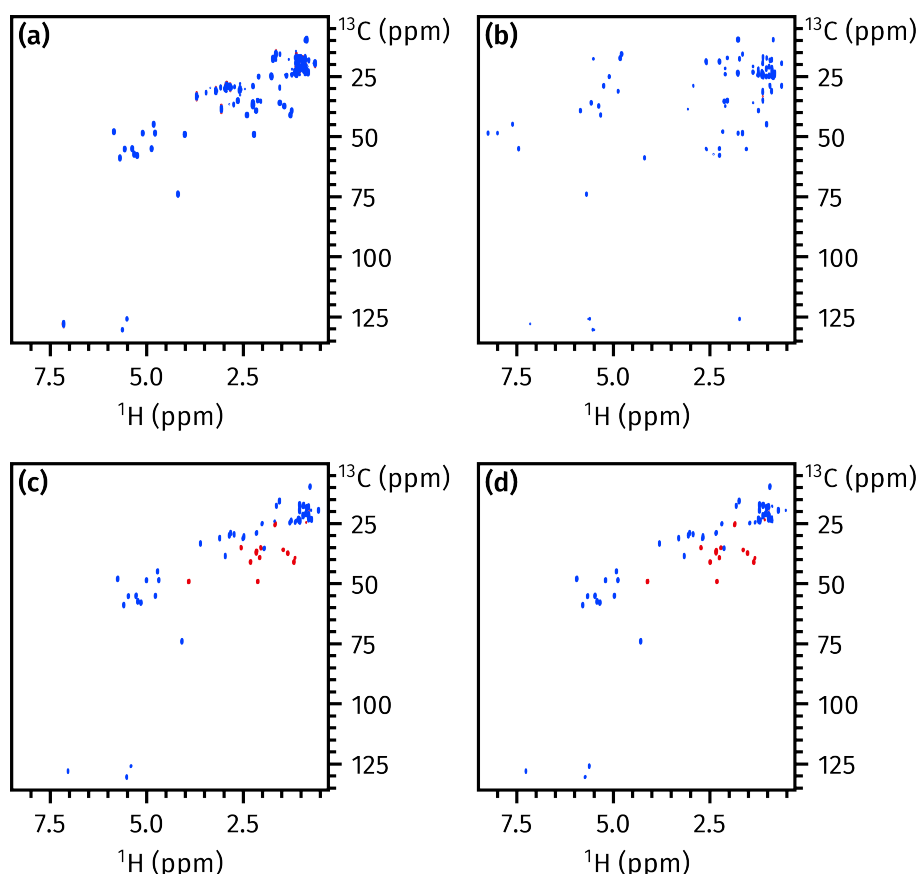


Figure 4.65: (a)–(b) IPAP-processed HSQC-COSY spectra (from figs. 4.64d and 4.64e), which separate the ‘direct’ and ‘indirect’ HSQC-COSY responses described in § 4.3.5. (c)–(d) IPAP-processed seHSQC spectra (from figs. 4.64f and 4.64g), each of which contain one peak of each 1H – ${}^{13}C$ doublet.

Time-shared NMR

In my published work,² what I refer to as IPAP here is frequently labelled as a TS, or *time-shared* experiment. In TS NMR, pulse sequences are specially designed in order to record two (or more) different signals at once, such as a ¹³C HSQC and a ¹⁵N HSQC: this has found use in both biomolecular^{102–107} and small-molecule NMR.^{26,27,108,109} Often, the design of TS experiments involves the joint evolution of both heteronuclear chemical shifts in a shared t_1 period: hence the term ‘time-shared’.

It should be noted that TS experiments use *simultaneous* acquisition, as opposed to the *sequential* acquisition of NOAH experiments. In other words, the two or more signals recorded in a TS experiment are part of the same FID, meaning that the resulting spectra will contain peaks from both signals. If these can be easily disambiguated (e.g. by the differing chemical shifts of ¹³C- or ¹⁵N-bound protons), then the TS experiment allows more data to be collected in the same amount of time as a non-TS experiment. However, if these signals must be separated, then the TS experiment must be repeated: once where the two signals have the same phase, and once where they have opposite phases. The underlying signals can then be disentangled through linear combination;^{102,110} this process leads to an improvement in sensitivity, which can be re-converted into time savings by using fewer scans for the TS experiment (if this does not compromise spectral quality too much).

Mathematically, this is exactly the same process as IPAP processing. In my view, what separates the two is the fact that the IPAP technique seeks to disentangle components which are *innate* to the pulse sequence under consideration and cannot usually be isolated—such as doublets in a F_2 -coupled HSQC—whereas TS experiments must be *specially crafted* to record these multiple components. Equivalently, the time savings (or sensitivity improvement) from a TS experiment stem from the fact that it is being compared against a pair of experiments which each yield only one of the two signals detected in the TS sequence. To justly label an experiment as being TS, appropriate choices must be made for these ‘base’ spectra.

In this case, the IPAP label clearly applies to the pair of F_2 -coupled seHSQC modules in figs. 4.64f and 4.64g. However, for the HSQC-COSY modules in figs. 4.64d and 4.64e, the situation is less clear. The two ‘signals’ which we should consider are the ‘direct’ and ‘indirect’ peaks; however, it is not entirely obvious which experiments can be used for these. The ‘direct’ peaks correspond to a (¹³C-decoupled) HSQC experiment.* However, for the ‘indirect’ peaks, there is no real equivalent spectrum to be compared against: the H2BC experiment^{79,80} yields a similar set of

* It should be noted that the intensities of the direct peaks in an HSQC-COSY spectrum are lower than in the corresponding HSQC spectrum, because the direct peaks in an HSQC-COSY stem only from magnetisation which does *not* evolve under $^2J_{\text{HH}}$ during the mixing period. However, TS experiments do generally have lower sensitivity than the non-TS counterparts, so this is not really an issue.

peaks, but in this experiment, the ‘direct’ peaks have in fact been intentionally suppressed through the application of a LPJF. When the LPJF is removed (as in the 2BOB/H2OBC experiments⁸¹), a similar information content to the HSQC-COSY is obtained. So, it is not entirely accurate to write that the HSQC-COSY combines a HSQC plus H2BC spectrum. Consequently, I personally prefer to view it as an IPAP experiment.

Of course, the analysis above is only applicable to the modules shown in fig. 4.64. None of this precludes the application of *true* time-shared experiments to NOAH supersequences: it is completely possible to implement, for example, a $^{13}\text{C}/^{15}\text{N}$ TS experiment within the context of a parallel supersequence. However, many TS experiments are already designed to detect signals arising from different magnetisation pools (e.g. $^1\text{H}^{\text{C}}$ and $^1\text{H}^{\text{N}}$). This is already something which can be accomplished in a cleaner manner using (linear) NOAH supersequences.

4.5.2 Generalised supersequences

From the discussion above, it is evident that there is no need to stop at *two* interleaved modules: it is possible to stack an arbitrary number of modules in a vertical manner. Furthermore, different modules may be acquired a different number of times. Thus, in fig. 4.63b, two supersequences were interleaved, and the TOCSY and NOESY modules were acquired once each in each t_1 increment. However, if M different supersequences are being interleaved, then a set of two different modules may be acquired m_1 and m_2 times, as long as $m_1 + m_2 = M$. For example, a supersequence with $m_1 = 1$ and $m_2 = 3$ is shown in fig. 4.66. Generally, it is desirable to assign larger m_i values to less sensitive modules, as the results can be summed up to yield improved signal-to-noise.

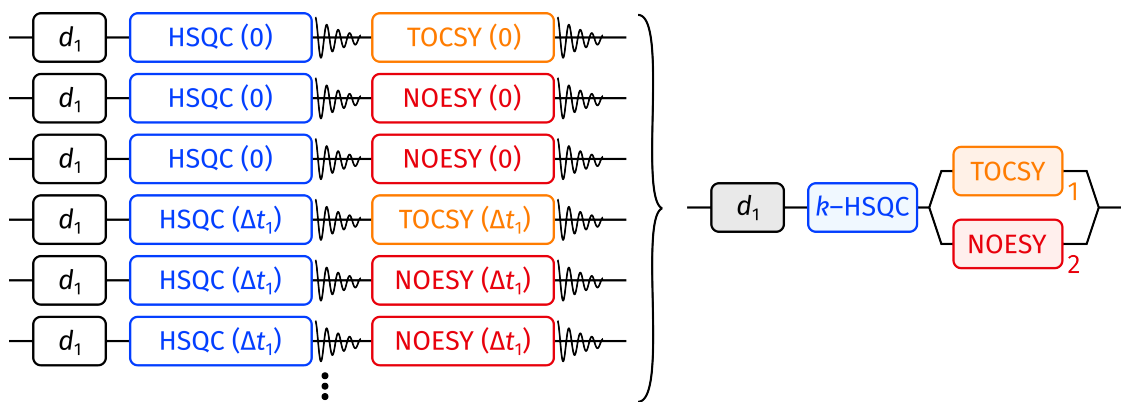


Figure 4.66: A basic example of a ‘generalised’ NOAH supersequence, where an arbitrary number of supersequences (here three) can be vertically ‘stacked’. For each value of t_1 , the TOCSY is acquired once, the NOESY twice, and the HSQC three times, as indicated by the subscripts in the condensed representation on the right.

Pulse programmes

Unfortunately, GENESIS is not (at present) advanced enough to create pulse programmes for generalised supersequences. Thus, the pulse programmes for this section have been written by hand. The overall structure of such a pulse programme is shown in listing 4.4. The parameter CNST51 corresponds to the number of supersequences being interleaved, i.e., M in the discussion above. The values of m_1 and m_2 are encoded as CNST52 and CNST53; the latter of these can be automatically calculated.

This structure provides a clear blueprint for how GENESIS can be adapted to produce generalised supersequences. The only thing which GENESIS needs to detect is how many modules are being interleaved, which determines the number of m_i 's. It should be noted that the ‘parallel’ supersequence of fig. 4.63c can be obtained by setting $M = 2$ and $m_1 = m_2 = 1$, and standard linear supersequences simply have $M = m_1 = 1$. Thus, if generalised supersequences were to be implemented in GENESIS, all other types of supersequences can be obtained ‘for free’.

A small technical detail in listing 4.4 is that each `go=2` statement loops back to the label 2, such that the experiment is repeated `NS` times. This means that in the left side of fig. 4.66, each row is already acquired a total of `NS` times, in addition to any repetition which is explicitly shown. Thus, after summation of the explicitly repeated increments, the HSQC will in fact end up with a total of $3 \times \text{NS}$ scans. However, the *phase cycle* is only moved forward within the `NS` loop; each time `NS` repetitions are completed, the phase cycle is reset to the beginning. This means that even though each FID in the HSQC is acquired $3 \times \text{NS}$ times, only an `NS`-step phase cycle is used. Thus, in order to obtain spectra of the highest possible quality, any common divisors of M and the m_i 's should be factorised out and placed into `NS` instead.

Examples

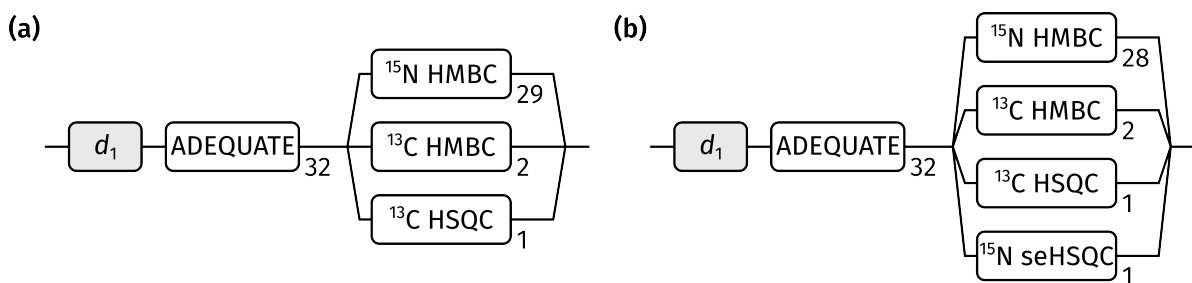


Figure 4.67: Two examples of generalised NOAH supersequences. (a) A NOAH-4 AB_N BS supersequence. (b) A NOAH-5 AB_N BSS_N⁺ supersequence.

Since generalised supersequences allow the effective number of scans to be individually customised for each module, they offer an avenue for modules with (very) different intrinsic sensitivities to be combined. On the very-low-sensitivity end are spectra such as ADEQUATE and ¹⁵N

```

"10      = td1/2"                      ; TD1/NBL
"cnst53 = cnst51 - cnst52"    ; automatically calculate m_2 = M - m_1
; ...

1 ze
2 30m
4 50u UNBLKGRAD
d1 st0

; HSQC goes here
; ...
goscnp ph30 cpd2:f2
50u do:f2

; check which module to run
if "13 % cnst51 < cnst52"
{
; TOCSY goes here
; ...
go=2 ph31
}
else
{
; NOESY goes here
; ...
go=2 ph31
}

; move to next 'row' in diagram
1m iu3
30m wr #0 if #0 zd

; check if M 'rows' have passed
if "13 % cnst51 == 0"
{
1m iu1
1m igrad EA   ; HSQC echo-antiecho gradients
; ...
}

lo to 4 times 10
end

```

Listing 4.4: The overall structure of the pulse programme for the generalised supersequence in fig. 4.66.

HMBC; compared to these, the HSQC experiments may be considered high-sensitivity. It should be reiterated that the time savings to be obtained through this combination (ρ_t) are still on the order of 2, because only two modules are being horizontally concatenated at a time.

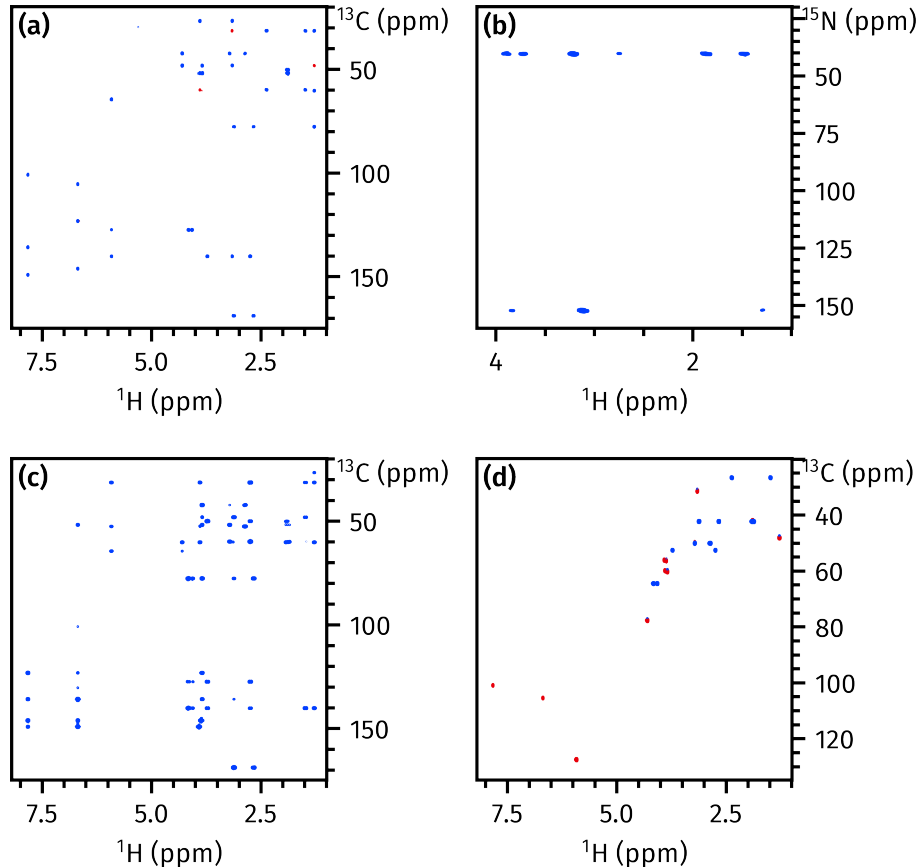


Figure 4.68: Spectra from the NOAH-4 AB_NBS generalised supersequence (fig. 4.67a). (a) ADEQUATE. (b) ¹⁵N HMBC, optimised for an $^nJ_{\text{NH}}$ value of 8 Hz. (c) ¹³C HMBC, optimised for an $^nJ_{\text{CH}}$ value of 8 Hz. (d) ¹³C HSQC. Data code: 7X-220604.

Figure 4.67 shows some examples of such experiments, where the relative numbers of scans are chosen with the aim of balancing the sensitivities of each module. The ADEQUATE module used here is the ZIP-ADEQUATE experiment shown in fig. 4.58; this consumes ¹H^C magnetisation and preserves the ¹H^{I,C} magnetisation required by the ¹⁵N HMBC, ¹³C HMBC, and ¹⁵N seHSQC. However, the ¹³C HSQC module draws on the same magnetisation pool. Therefore, in order to maximise the intensities in this experiment, a period of isotropic mixing is applied only before the HSQC module in order to effect polarisation transfer from ¹H^{I,C} spins. The spectra thus obtained are shown in figs. 4.68 and 4.69.

Another additional implementation detail, related to the earlier point about phase cycling, is that between each of the 28 or 29 repetitions of the ADEQUATE/¹⁵N HMBC increments, the ¹⁵N HMBC signal (specifically, the first ¹⁵N 90° pulse) and receiver phases are additionally inverted. This can be accomplished by adding the corresponding extra phase incrementation instructions

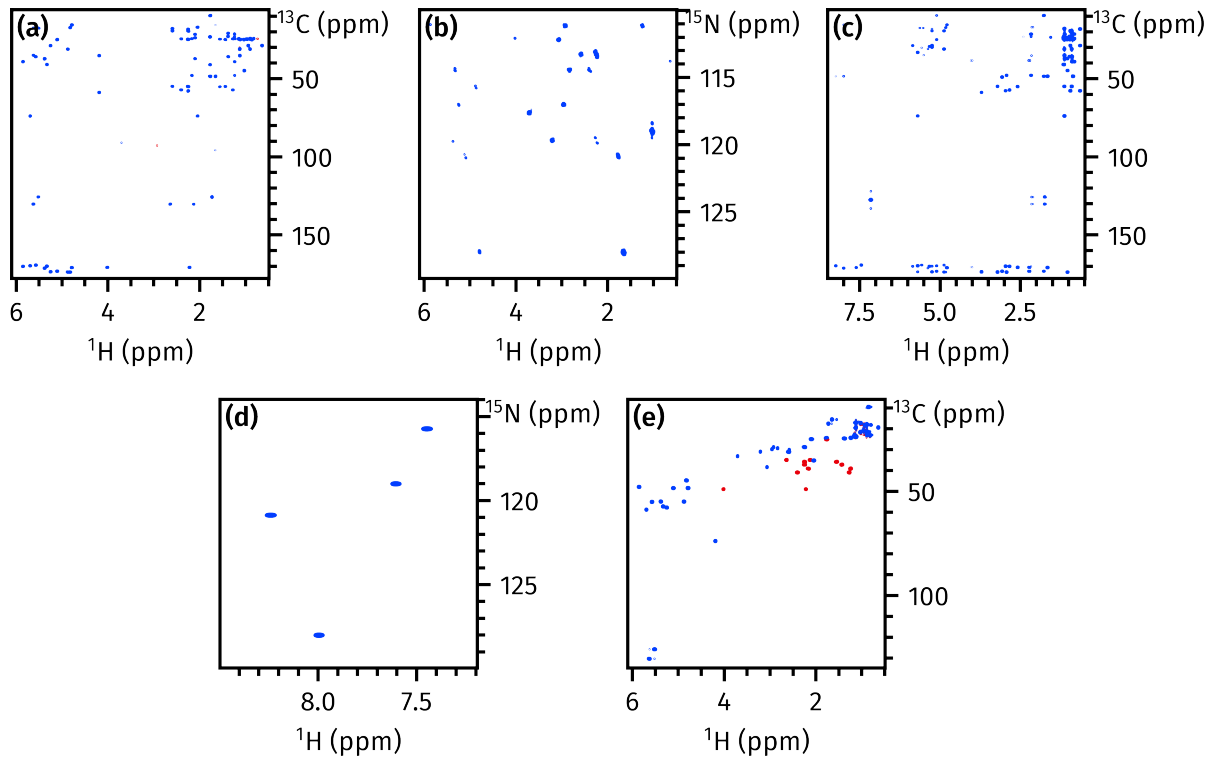


Figure 4.69: Spectra from the NOAH-5 AB_NBSS_N⁺ generalised supersequence (fig. 4.67b). (a) ADEQUATE. (b) ^{15}N HMBC, optimised for an $^nJ_{\text{NH}}$ value of 4 Hz. (c) ^{13}C HMBC, optimised for an $^nJ_{\text{CH}}$ value of 8 Hz. (d) ^{13}C HSQC. (e) ^{15}N seHSQC. Data code: 7C-220722.

just underneath the $1\text{m} \text{ iu3}$ line in listing 4.4, and can be thought of as a form of ‘manual’ phase cycling, which is independent of (and complementary to) the usual phase cycle specified in the pulse programme. The reason for doing this is to suppress artefacts which appear at $F_1 \approx 0$ in the ^{15}N HMBC, which stem from the ADEQUATE module (when the ^{15}N HMBC is run on its own, these artefacts do not appear). The exact origin of these artefacts is still unclear,* but their phases are constant; thus, inverting the ^{15}N HMBC receiver phase leads to their cancellation. It should be noted that the issue of phase cycling is not unique to generalised supersequences: indeed, such phase cycling would also be necessary in a standard linear NOAH-2 AB_N supersequence. However, because the degree of built-in phase cycling in a generalised supersequence is lower than the actual number of transients recorded for each module, some of it must be explicitly specified in this way.

*Their frequencies in F_1 are not exactly zero, so there is *some* kind of modulation in the t_1 period of the ADEQUATE module, but it is not entirely clear where this is happening. My current best guess is that the offending magnetisation evolves during the constant-time period of the ADEQUATE, where the double-quantum frequencies are reconverted into single-quantum frequencies.

Covariance analysis

The generalised supersequences shown in figs. 4.67 to 4.69 yield very thorough information about heteronuclear connectivity, including one-bond and long-range ^1H - ^{13}C and ^1H - ^{15}N couplings. These spectra can be used as inputs for indirect covariance processing^{111–113} in order to yield ‘unnatural’ (but very informative) double heteronuclear correlation spectra, in which each peak represents a correlation between two dilute heteronuclei (^{13}C or ^{15}N). Directly recording such spectra through an NMR experiment would require substantially more time, because of the extremely low probability of finding two heteronuclei in the same molecule.

Some possible covariance spectra obtained from the data in figs. 4.68 and 4.69 are shown in fig. 4.70. All covariance calculations were performed in Python, according to the formulae:

$$\mathbf{C} = \mathbf{S}_1 \mathbf{S}_2^T \quad (4.23)$$

for unsymmetrical indirect covariance (where \mathbf{S}_1 and \mathbf{S}_2 are the input spectra and \mathbf{C} the covariance spectrum), or

$$\mathbf{M} = \left[\begin{pmatrix} \mathbf{S}_1 \\ \mathbf{S}_2 \end{pmatrix} (\mathbf{S}_1 \quad \mathbf{S}_2) \right]^\lambda \quad (4.24)$$

for generalised indirect covariance; the covariance spectrum \mathbf{C} can then be obtained as an off-diagonal block of the matrix \mathbf{M} . λ is a parameter which is chosen in order to balance peak intensity and artefact intensity:¹¹² smaller values of λ lead to better suppression of false positives, but lower signal intensity (and also cause distortions in peak shapes).

In fig. 4.70a, the brucine ^{15}N HMBC and ^{13}C HSQC spectra (figs. 4.68b and 4.68d) are processed using unsymmetrical indirect covariance (eq. (4.23)) to yield a ^{13}C - ^{15}N correlation spectrum which contains both one- and long-range correlations.^{114,115} The sign of the peaks indicates the multiplicity of the ^{13}C involved, and stems from the use of multiplicity editing in the ^{13}C HSQC spectrum. Unfortunately, one of the weaknesses of covariance spectra is on show here: there are several artefacts in this spectrum (marked by asterisks), caused by overlap in the ^1H dimension of both input spectra.

The other three spectra (figs. 4.70b to 4.70d) combine the cyclosporin ^{13}C HSQC and AD-EQUATE spectra (figs. 4.69a and 4.69d) to generate a ^{13}C - ^{13}C one-bond correlation spectrum, using generalised indirect covariance (eq. (4.24)) with $\lambda = 0.5$. This is essentially an INAD-EQUATE spectrum, but with single-quantum frequencies in both dimensions.^{116,117} Such a correlation spectrum should in principle be symmetric about the main diagonal. However, since the HSQC only detects protonated carbons, and the ADEQUATE only detects correlations involving at least one protonated carbon, correlations between two quaternary carbons do not appear at all, and correlations between protonated and quaternary carbons will appear only once

in the spectrum (instead of twice). This does not pose a problem in the case of cyclosporin, though, as the only quaternary carbons are those of the carbonyl (CO) group, which will still display correlations to the adjacent C α carbons.

These C α -CO correlations are shown in fig. 4.70b. There are 11 peaks here, one for each amino acid residue in cyclosporin: again, the sign indicates the C α multiplicity (all are CH, except for the CH₂ of the sarcosine residue). Figures 4.70c and 4.70d shows the C α -C β correlations. Figure 4.70c shows the original spectrum, but in fig. 4.70d, the spectrum has been further subjected to a sign-preserving symmetrisation procedure, where the intensity at each point $p(\Omega_1, \Omega_2)$ is replaced by

$$p(\Omega_1, \Omega_2) \rightarrow \text{sgn}[p(\Omega_1, \Omega_2)] \cdot \min \{|p(\Omega_1, \Omega_2)|, |p(\Omega_2, \Omega_1)|\}. \quad (4.25)$$

Here, sgn x refers to the sign of x , or equivalently $x/|x|$ (for $x \neq 0$). Since all C α and C β carbons in cyclosporin are protonated, this region of the spectrum is expected to be symmetric;^{*} the symmetrisation process therefore helps to remove some of the artefacts which tend to plague the interpretation of covariance spectra.

4.6 Conclusion

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^{*}The peak intensities $p(\Omega_1, \Omega_2)$ and $p(\Omega_2, \Omega_1)$ may not be exactly equal, but both peaks should at least be present.

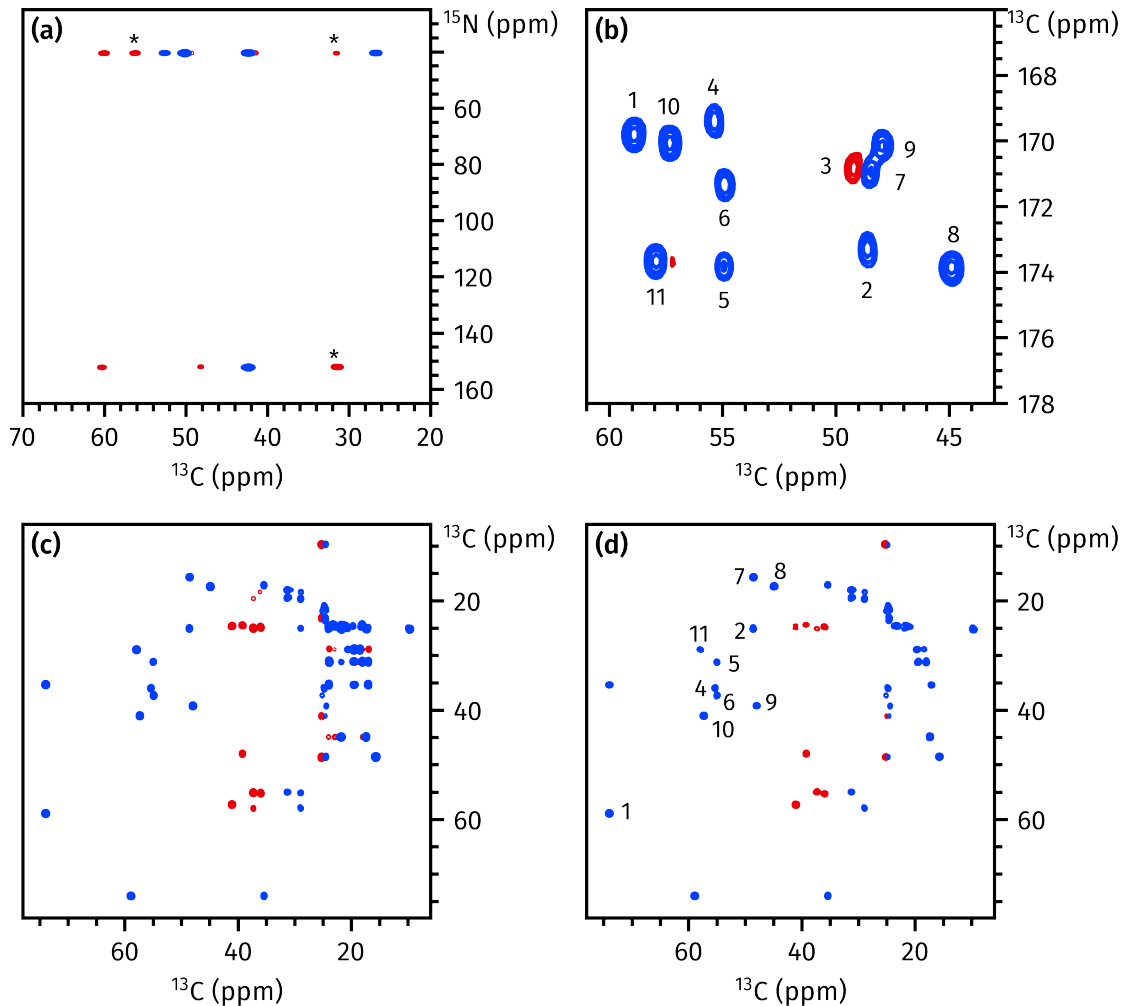


Figure 4.70: Examples of covariance spectra obtained from generalised NOAH supersequences. (a) ^{13}C - ^{15}N correlation spectrum for brucine. Asterisks indicate artefacts arising from peak overlap. (b) Inset of the ^{13}C - ^{13}C one-bond correlation spectrum for cyclosporin, obtained by processing the ADEQUATE and ^{13}C HSQC spectra in figs. 4.69a and 4.69d using generalised indirect covariance ($\lambda = 0.5$). The region shown here contains correlations between CO and C α carbons. (c)-(d) The same ^{13}C - ^{13}C correlation spectrum as in (b), but this time showing correlations in the alkyl region. The spectrum in (c) is untouched, but the spectrum in (d) has been further subject to a sign-preserving symmetrisation process. The CO-C α and C α -C β correlations for all residues are labelled in (b) and (d), following the residue numbering in fig. A.2.

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Appendix A

Peak assignments for some samples

‘It is over! it is over!’ she repeated to herself again, and again, in nervous gratitude.
‘The worst is over!’

— JANE AUSTEN, *Persuasion*

In this appendix, I provide assignments of ^1H and ^{13}C chemical shifts for some of the more frequently-used samples in this thesis, occasionally with some additional data. I have made no attempt to distinguish the shifts of diastereotopic protons or methyl groups.

A.1 Andrographolide

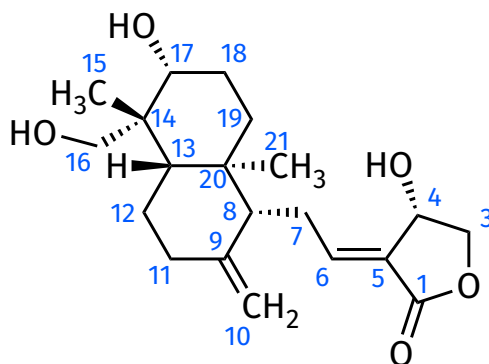


Figure A.1: Structure of andrographolide.

Label in fig. A.1	Label in Claridge (2016)	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)
1			170.42
3	g, i	4.40, 4.04	74.80
4	d	4.92	64.99
4-OH	b	5.71	
5			129.46
6	a	6.63	146.76
7	m, n	2.52, 2.47	24.43*
8	q	1.87	55.96
9			148.07
10	e, f	4.82, 4.63	108.71
11	o, p	2.33, 1.94	37.98
12	r, v	1.75, 1.36	24.42*
13	w	1.21	54.84
14			42.75
15	y	1.09	23.54
16	j, k	3.85, 3.27	63.11
16-OH	h	4.13	
17	l	3.24	78.91
17-OH	c	5.05	
18	t, u	1.65, 1.65	28.36
19	s, x	1.70, 1.21	36.99
20			39.06
21	z	0.67	15.22

Table A.1: Peak assignments for andrographolide; these can also be found in Figure 9.26 (page 340) of: Claridge, T. D. W., *High-Resolution NMR Techniques in Organic Chemistry*, 3rd ed.; Elsevier: Amsterdam, 2016. Asterisks indicate ^{13}C chemical shifts which could not be disambiguated.

A.2 Cyclosporin

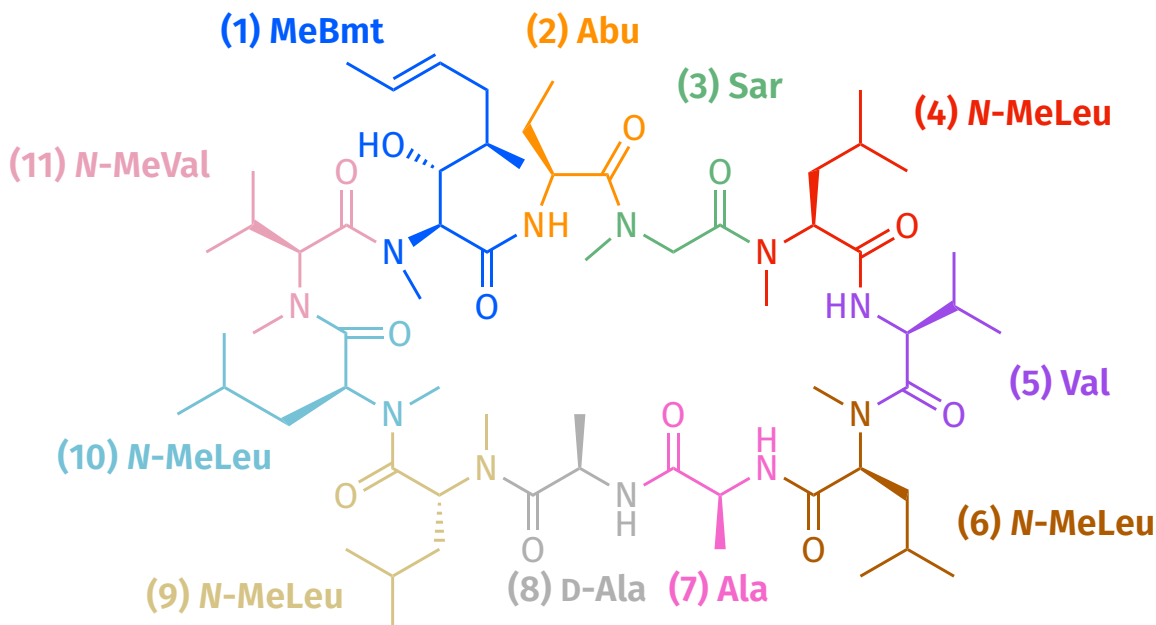


Figure A.2: Structure of cyclosporin.

Residue	Group	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$\delta(^{15}\text{N})$ (ppm)
1 MeBmt	NCH ₃	3.70	33.5	117.7
	α -CH	5.69	59.0	
	β -CH	4.19	74.1	
	CO		169.9	
2 Abu	NH	8.24		120.9
	α -CH	5.10	48.7	
	β -CH ₂	1.76	25.3	
	CO		173.2	
3 Sar	NCH ₃	3.07	38.6	112.2
	α -CH ₂	4.01	49.2	
	CO		170.9	
4 N-MeLeu	NCH ₃	2.58	30.4	113.3
	α -CH	5.57	55.3	
	β -CH ₂	1.35	36.1	
	CO		169.4	
5 Val	NH	7.45		115.7
	α -CH	4.87	55.2	
	β -CH	2.60	31.3	
	CO		173.8	

Table A.2: Peak assignments for cyclosporin.

Table A.2: (continued)

Residue	Group	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$\delta(^{15}\text{N})$ (ppm)
6 N-MeLeu	NCH ₃	3.21	31.2	119.7
	α -CH	5.37	55.1	
	β -CH ₂	2.25, 1.44	37.4	
	CO		171.4	
7 Ala	NH	8.00		128.0
	α -CH	4.78	48.7	
	β -CH ₃	1.65	15.8	
	CO		171.0	
8 D-Ala	NH	7.61		119.0
	α -CH	4.82	45.0	
	β -CH ₃	1.03	17.6	
	CO		173.9	
9 N-MeLeu	NCH ₃	2.92	29.1	111.2
	α -CH	5.85	48.0	
	β -CH ₂	2.16, 1.25	39.3	
	CO		170.2	
10 N-MeLeu	NCH ₃	2.84	29.6	114.4
	α -CH	5.32	57.5	
	β -CH ₂	2.40, 1.28	41.2	
	CO		170.1	
11 N-MeVal	NCH ₃	2.96	30.1	117.7
	α -CH	5.25	58.0	
	β -CH	2.25	29.1	
	CO		173.8	

Table A.2: Peak assignments for cyclosporin.

A.3 Ferulic acid

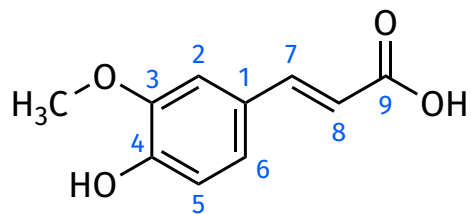


Figure A.3: Structure of ferulic acid.

Label	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$^1J_{\text{CH}}$ (Hz)	$T_1(^1\text{H})$ at 600 MHz (s)
1		126.25		
2	7.28	111.62	157.6	1.15
3		148.38		
3-OCH ₃	3.82	56.16	147.8	1.08
4		149.54		
5	6.79	115.98	159.5	1.59
6	7.08	123.27	158.6	2.00
7	7.49	144.97	154.3	1.96
8	6.36	116.08	160.6	1.75
9		168.45		

Table A.3: Peak assignments and other physical data for ferulic acid.

A.4 Gramicidin

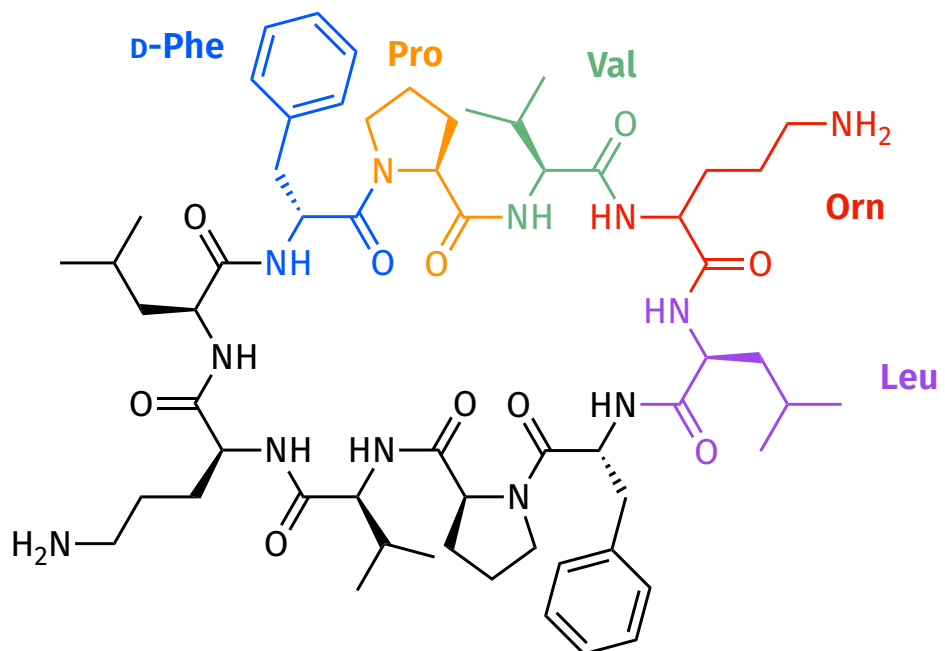


Figure A.4: Structure of gramicidin.

Residue	Group	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$\delta(^{15}\text{N})$ (ppm)
Leu	NH	8.33		123.3
	α -CH	4.57	50.09	
	β -CH ₂	1.35, 1.29	41.38	
	γ -CH	1.41	24.45	
	δ -CH ₃	1.41	23.20, 23.02	
Orn	NH	8.66		125.4
	α -CH	4.76	51.43	
	β -CH ₂	1.75, 1.60	30.12	
	γ -CH ₂	1.65	23.52	
	δ -CH ₂	2.84, 2.78	39.02	
	ε -NH ₂	8.04		36.0
Val	NH	7.22		113.2
	α -CH	4.41	57.31	
	β -CH	2.07	31.49	
	γ -CH ₃	0.80, 0.77	19.44, 18.50	
Pro	α -CH	4.31	60.36	
	β -CH ₂	1.95, 1.48	29.50	
	γ -CH ₂	1.52	23.58	
	δ -CH ₂	3.59, 2.50	46.48	
D-Phe	NH	9.09		128.0
	α -CH	4.36	54.38	
	β -CH ₂	3.59, 2.50	46.48	
	ipso-C		136.77	
	ortho-CH	7.26	129.80	
	meta-CH	7.29	128.71	
	para-CH	7.25	127.34	

Table A.4: Peak assignments for gramicidin.

A.5 Brucine

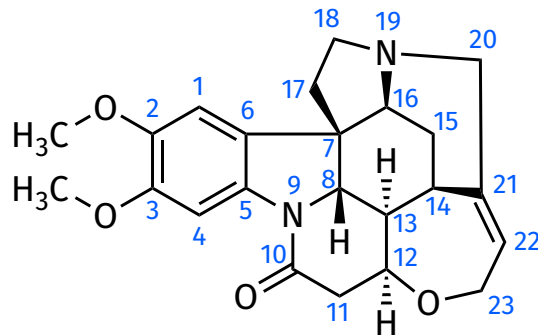


Figure A.5: Structure of brucine.

Label	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$\delta(^{15}\text{N})$ (ppm)
1	6.69	105.52	
2		146.24	
2-OCH ₃	3.87	56.46	
3		149.25	
3-OCH ₃	3.92	56.21	
4	7.83	101.00	
5		135.98	
6		123.33	
7		51.94	
8	3.85	60.35	
9			152.38
10		168.94	
11	3.12, 2.67	42.39	
12	4.30	77.79	
13	1.28	48.26	
14	3.16	31.55	
15	2.37, 1.49	26.79	
16	3.90	60.00	
17	1.89	42.41	
18	3.22, 2.87	50.21	
19			40.55
20	3.73, 2.75	52.71	
21		140.32	
22	5.92	127.54	
23	4.16, 4.07	64.60	

Table A.5: Peak assignments for brucine.

A.6 Zolmitriptan

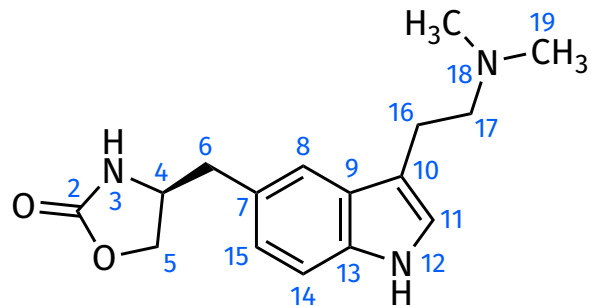


Figure A.6: Structure of zolmitriptan.

Label	$\delta(^1\text{H})$ (ppm)	$\delta(^{13}\text{C})$ (ppm)	$\delta(^{15}\text{N})$ (ppm)	$^1J_{\text{CH}}$ (Hz)
2		159.16		
3	7.77		89.2	
4	4.05	53.64		146.7
5	4.23, 4.03	68.54		153.0, 151.7
6	2.90, 2.79	41.11		127.7, 127.1
7		126.32		
8	7.37	119.24		155.4
9		127.99		
10		112.92		
11	7.12	123.18		179.7
12	10.71		129.6	
13		135.64		
14	7.26	111.71		158.9
15	6.93	122.97		155.7
16	2.81	23.55		125.8
17	2.53	60.46		132.2
18			106.6	
19	2.26	45.65		132.5

Table A.6: Peak assignments and other physical data for zolmitriptan.