

1 A modular library for fast prototyping of solution-state nuclear 2 magnetic resonance experiments

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8 **Abstract.** We present a framework library (Modular Elements, ME) for the development of pulse sequences for Bruker
9 spectrometers. It implements a two-level abstraction approach—the lower level comprises basic functional elements of pulse
10 sequences and the higher one often-reused blocks comprising ~~multiple spin echoes~~ several evolution periods. The low-level
11 abstractions reduce code duplication between variants of experiments such as hard-pulse and selective variants of individual
12 NMR experiments. The high-level modules enable further reuse of pulse program code and aid in the construction of
13 complex experiments. We show the library’s functionality by presenting pulse programs that can be switched between
14 standard and TROSY variants, hard and shaped pulses and can seamlessly incorporate real-time homodecoupling—~~—~~.
15 Adaptability is further demonstrated in a configurable 4D NOESY program.

16 1 Introduction

17 NMR is an extraordinarily powerful and adaptable spectroscopic method, with just the solution-state variant being capable of
18 discerning the structure and dynamics of molecules ranging in size from simple organic compounds to large protein
19 complexes such as a proteasome (Sprangers and Kay, 2007). The variety of experimental objects and the great number of
20 parameters that can be measured has led to the proliferation of not only general experimental schemes—~~—~~ such as an ^1H , ^{15}N
21 HSQC (Bodenhausen and Ruben, 1980) or a HNCQ (Kay et al., 1990b; Ikura et al., 1990)), but also their variants and thus
22 the pulse sequences, implementing them as computer code. As an example, for the often-used HNCQ experiment, the non-
23 exhaustive list of meaningful implementation choices is: the experiment can use hard pulses or avoid saturating water using
24 selective pulses (Schanda et al., 2006); the final transfer element can be a simple spin-echo (Palmer et al., 1991), a set of
25 three echoes implementing a sensitivity-enhanced transfer or one of many TROSY variants (Salzmann et al., 1999b;
26 Nietlispach, 2005), with possible optimizations (Salzmann et al., 1999a; Schulte-Herbrüggen and Sørensen, 2000); radiation
27 damping can be suppressed with bipolar gradients (Sklenar, 1995). Even without implementing all ~~specialized~~ specialised
28 experiment variants, the standard library supplied with the TopSpin software (Bruker) contains over a thousand pulse
29 programs.

30 A common problem with pulse sequences, especially in biological NMR, is thus the requirement to code multiple variants of
31 a given sequence. If this is done in separate files (as in the TopSpin built-in library) it results in a lot of code repetition and if
32 made using conditional statements, it can substantially complicate the structure of the file, making trouble-shooting harder.
33 Similarly, many pulse sequences share large amounts of code, often with no or minimal changes. Because this repeated code
34 is scattered across different sequences and variants of experiments adding new variants (using different soft pulses, adding
35 homodecoupling) requires applying the same modification across a large part of the whole pulse sequence library, which is
36 tedious and error-prone. It is possible to implement such a library using standard systems programming language like C or
37 Python, but we decided to use the native programming language of the spectrometer system, since any user writing pulse
38 sequence needs to be familiar with it and requiring knowledge of separate programming language and its tooling would be
39 an unnecessary hurdle to adoption. Here we show that by abstracting certain functionality using the somewhat limited macro
40 and "define" functionality built into the TopSpin software, the above-described problems can still be avoided and the code
41 can be made more readable and easier to modify. Here, we present the Modular Elements (ME) library for Bruker
42 spectrometers. Although the library is specific to a particular hardware vendor, the modular approach it implements is more
43 general and can be implemented on other instruments. A previous implementation of a modular library for pulse program
44 implementation (NMR blocks) can be found in (Zawadzka-Kazimierczuk, 2012) for Varian/Agilent spectrometers, where
45 ~~spin-echos and transferevolution~~ periods such as INEPT or COS-INEPT where abstracted as C functions. Alternative
46 approaches to a modular library include domain specific pulse program generators, like GENESIS (Yong et al., 2022) for
47 NOAH supersequences. Specialized libraries combining custom pulse programs and various tools (Favier and Brutscher,
48 2019; Vallet et al., 2020; Lukavsky and Puglisi, 2001), are suitable for routine use, but have limited applicability in the
49 prototyping of new sequences.

50 **2 General approach to pulse sequence modularisation**

51 We categorise the library's functionality as low- and high-level. Low-level functionality encompasses the creation of
52 variables and functions (technically functional macros), abstracting the basic building blocks of pulse sequences like pulses,
53 gradients and delays. A pulse function can ~~evaluate to expand to~~ a 90 degree proton hard pulse or an HN selective excitation
54 pulse depending on global settings. A gradient function can ~~evaluate expand~~ to "no operation" in standard HSQC, or a
55 selection gradient ~~it in the g~~ ~~the gradient-selected~~ HSQC variant, with its corresponding delay variable containing a zero or
56 correspondingly non-zero length of time. Decoupling functions for protons and deuterium can likewise be enabled and
57 disabled depending on whether a TROSY variant is desired and if the sample is deuterated. This functionality simplifies the
58 writing of pulse sequences implementing multiple variants of a given NMR experiment and gives its user the ability to easily
59 test and compare the effectiveness of the variants for a given sample and the commonization of parameters across variants
60 enables faster optimisation.

61 | High-level functionality is implemented as modules that are included ~~whole~~ in the pulse sequences and can be classified as
62 | general modules and specific modules. General modules implement elements common to almost all pulse sequences. The
63 | functionally most significant ones are the preparation and acquisition modules. The preparation module gives the user the
64 | option to turn on functionality such as solvent presaturation or a combination of N/C pulses and pulse field gradients for
65 | spoiling of residual magnetisation on those nuclei. The acquisition module enables switching between standard or
66 | homodecoupled data acquisition. The specific modules are abstract blocks of pulse sequence elements that appear in many
67 | pulse sequences in an almost identical form. Two main types of specific modules are proximal and distal modules,
68 | abstracting the functionality of blocks including and following first excitation (distal) and right before acquisition
69 | (proximal). Despite a large variety of possible implementations, the proximal/distal fragments differentiate variants of a
70 | pulse sequence (for example a standard hard-pulse HNCO, selective/BEST-HNCO, hard pulse and BEST TROSY-HNCO
71 | (Solyom et al., 2013)) and the actual code is usually repeatable across different sequences. HNCO, HNCACO and HNCOCA
72 | (Yang and Kay, 1999) have very similar proximal and distal parts; HN(CA)CONH and HabCabCONH (Kazmierczuk et al.,
73 | 2010) have different proximal blocks, but the distal block is still very similar for all sequences listed. With the use of the
74 | low-level functionality described above a single proximal module can abstract the initial two transfer periods (first with
75 | transverse H magnetisation and second with transverse N/C magnetisation), with the choice of N/C nucleus and choice of
76 | evolved J coupling (CO in HNCO) made using define directives in the main pulse sequence. NOESY experiments are
77 | particularly susceptible to modularisation, with the NOE transfer period naturally splitting them into proximal and distal
78 | blocks. Standard 2D experiments of the HSQC, TROSY and HMQC type have thus been implemented as proximal modules,
79 | that can be used on their own as 2D experiments or included in a 3 or 4D NOESY (Kay et al., 1990a) with the chosen distal
80 | modules, which can themselves be modified 2D experiments or simpler blocks.

81 | **3 Library implementation**

82 | Description of implementation details and design choices requires a quick recapitulation of TopSpin pulse ~~programs~~
83 | ~~programs'~~ language specifics. TopSpin ~~allows~~ has two types of variables: user-adjustable numbered variables (*d1..d63* for
84 | delays, *cnst1..cnst63* for floating point constants, similarly for integer constants ("loopcounters") *IN*, pulse lengths *pN*, ...)
85 | and named variables (pulses, delays and loopcounters only, also lists of various kinds), which can only be manipulated
86 | within a pulse program. Some less-documented observations on the limitations of named variables are compiled in SI.
87 | TopSpin implements limited functionality for defining text-substitution macros ("--traditional" mode of the GNU C
88 | preprocessor `cpp` (Stallman and GCC Developer Community, 2012)), which can be used everywhere outside a "relation"
89 | (variable value calculations using a subset of C syntax), due to their implementation as text in quotes (treated as string
90 | literals by `cpp` and ignored for macro expansion), though this limitation can be overcome (see the file "notes on TopSpin.txt"
91 | in the ME library). The user can provide custom option choices to a pulse program using the ZGOPNTS variable to define
92 | appropriate macros.

93 **3.1.1 Low-level modularisation**

94 **3.1.1 Variables**

95 With no user-adjustable named variables, two approaches to making them consistent across different pulse programs are
96 possible - indirection through `a`-named variables or introducing a convention attaching constant meaning to numbered
97 variables. Due to the limited number and type of named variables, we predominantly use the latter option (with sets of
98 variables described in files such as `delays.incl`, `pulse.incl`, ...) with some focused use of indirection - for example, proximal
99 type modules use `timeHX` and `timeXY` for J coupling evolution times between the H, X and Y nuclei. [Default values for all](#)
100 [such variables can be set using the `me.set_parameters.py` TopSpin program.](#) For variables that don't ordinarily have
101 calculations performed on them (pulse phases `phN`, gradient programs `gpN`) ~~we~~ implemented full indirection, where the
102 user can use `phFree1` or `phFree3` without worrying as to which `phN` variables are used by other parts of a pulse program.

103 **3.1.2 Pulses**

104 The most important low-level abstractions are pulse functions. They are implemented using function-like macros of `cpp` and
105 have the general form of `nucleus_type(phase)`, where `nucleus` can be a general specifier like `H/C/N` or more specific like
106 `HN/HC/CA/CO` and `type` is classified based on the desired functionality, with the main ones being: excitation (for the
107 excitation of longitudinal magnetization), flipback (acting on transverse magnetization), refocussing, inversion (inverting
108 longitudinal magnetization), `excitation_UR` and `flipback_UR` (implementing universal rotations). The pulse macros will have
109 different replacement text based on global settings (usually `ZGOPTNS`). A proton pulse "`H_excitation(ph)`" will ~~evaluate~~
110 ~~to be replaced by~~ a hard pulse "`p1 ph p1`" by default, but with a "`-DH_SHAPED`" option will instead ~~evaluate to be replaced~~
111 ~~by~~ "`p54:sp54 ph`" for a selective soft pulse and the associated named variable `pH_excitation` will be set to have the same
112 value as `p1` or `p54`.

113 Pulse programs should account for the effective evolution time `a`-during pulse (which can be as much as 1 ms for longer
114 selective pulses) to give correctly phased spectra and optimal J coupling evolution times. This library only accounts for
115 linear phase slope using the modelling method described in (Lescop et al., 2010), that is treating a pulse as sequence (delay,
116 ideal pulse, delay), which accounts for the phase slope of many commonly used pulses and can be [explicitly](#) optimized for
117 ~~consciously~~-during pulse design (Gershenson et al., 2008; Asami et al., 2018). This phase slope is compensated for using
118 variables such as `eH_excitation`, which for the hard pulse above would be set to $\frac{2p1}{\pi}$. We assume that the flipback and
119 `flipback_UR` pulses act as if they were time-reversed excitation pulses and so the effective evolution time for a flipback
120 pulse acting on transverse magnetization is also `eH_excitation`. ~~For a~~ `H_excitation_UR` pulse of phase `x` will give an
121 effective time of `eH_excitation` for z magnetization, `eH_flipback` for y magnetization and `eH_excitation + eH_flipback` for x
122 magnetization. By compensating delays using the above mentioned variables, the whole sequence can be switched from a

123 hard pulse implementation to a shaped pulse version, whether to account for field inhomogeneity or perform band-selective
124 excitation.

125 3.1.3 Code blocks

126 There are many small blocks of code that can be included/excluded in a pulse program based on a sequence variant. To limit
127 the number of conditional statements in the main pulse program, many are defined as macros that ~~evaluate-will expand~~ to
128 pulse program code based on options, for example “H2O_FLIPBACK(ph2)” will ~~evaluate-be replaced to-by~~ “(11:sp1
129 ph2):f1” ~~orin~~ a pulse sequence with water flipback and ~~byte~~ whitespace if using selective pulses. Similarly
130 DECOUPLE_H_ON and DECOUPLE_H_OFF macros will turn on proton decoupling in a standard HNC0 experiment but
131 will have no effect in TROSY-HNC0.

132 3.2 High-level modularization

133 TopSpin pulse programs follow a defined sequential structure that complicates the implementation of high-level modules as
134 individual files and, in general, is:

- 135 1) configuration and compile-time calculations
- 136 2) a "zd" or "ze" statement
- 137 3) pulse program body (pulses and delays) and real-time calculations
- 138 4) signal acquisition block
- 139 5) loop statements for scans of a FID and points of a multidimensional experiment
- 140 6) phase program definitions

141 3.2.1 General modules

142 The general modules fit into this sequential structure as follows:

- 143 1a) configuration and compile-time calculations
- 144 1b) **init.incl**
- 145 1c) configuration and compile-time calculations continued
- 146 2) a "zd" or "ze" statement
- 147 3a) real-time calculations
- 148 3b) **start.incl**
- 149 3c) pulse program body (pulses and delays) and real-time calculations
- 150 ~~3d)~~ **end.incl**
- 151 5) loop statements for scans of a FID and points of a multidimensional experiment
- 152 6a) **phasescycles.incl**

153 6b) phase program definitions

154 The general modules have numerous conditional statements and imports evaluating the option provided in point 1) above
155 and using the built-in ZGOPTNS variable and interact with the specific modules (this is covered below). The init.incl
156 module provides the [libraries-library's](#) core functionality by defining macros for functions and variable descriptions.
157 start.incl executes the relaxation delay (with possible solvent presaturation) and optional operations, such as crushing
158 residual C or N magnetization (gradient pulse after an excitation pulse) or inverting N magnetization before the relaxation
159 delay in BEST-TROSY. For non-protein experiments an ASAP (Kupče and Freeman, 2007) period would be added here, but
160 the relevant code is experimental and provided in a commented-out form due to the method's potential to damage
161 probeheads. The end.incl module handles acquisition with the option for real-time homodecoupling - here provided with ¹³C-
162 GBIRD^{FX} (Garbow et al., 1982; Haller et al., 2022) and BASHD (Brüschweiler et al., 1988; Krishnamurthy, 1997) types.

163 3.2.2 Specific modules

164 In contrast to the general modules, specific modules implement a specific form of proximal or distal block and serve to
165 localize the relevant code in a single file. The biggest hurdle to writing self-contained modules for TopSpin is the sequential
166 pulse program structure necessitating the separation of related code segments in the post-preprocessing file. To mitigate this
167 problem, each module is entirely enclosed in a conditional statement with alternative conditions (an if...elif...else structure)
168 and including the file once will only insert a selected part of the module into a file. Since the 4 general modules already
169 perform the sequential separation of code, each of them sets the appropriate conditions (defines a macro) and imports the
170 distal_2D.incl and proximal_2D.incl which themselves import the selected specific modules at each of the 4 positions in the
171 pulse program. Thus, the initialization phase statements (variable declarations, some calculations, macro definitions) are
172 included in init.incl, runtime calculations of both types of modules are included through start.incl, together with the main
173 body (pulses and delay statements) of the distal. Similarly, the main body of the proximal module is included through the
174 end.incl before the latter's acquisition portion. Phase cycles of both modules are inserted into a pulse program file through
175 phasecycles.incl with some basic logic, allowing for coordinating the cycles between them if two modules are used.

176 For triple-resonance experiments (in the implementation limited to amide protons, but should be possible to extend to
177 aliphatic/aromatic groups) the proximal module hx.incl and the distal module hx.incl provide the ability to compartmentalize
178 the relatively standard blocks for both out-and-back and straight through type experiments and a more detail description in
179 the context of a HNCO experiment is provided below. Although sub-optimal in some circumstances the library provides ~~a~~
180 default 2 step phase cycles for each of the modules, leaving the implementation of 8 step and longer cycles for the central
181 part of the program. [A more detailed description of individual modules is provided in library documentation. In the
182 supplemental we provide a detailed step-by-step description of the proximal HSQC module and the way it is used in the 2D
183 experiment pulse program.](#)

184

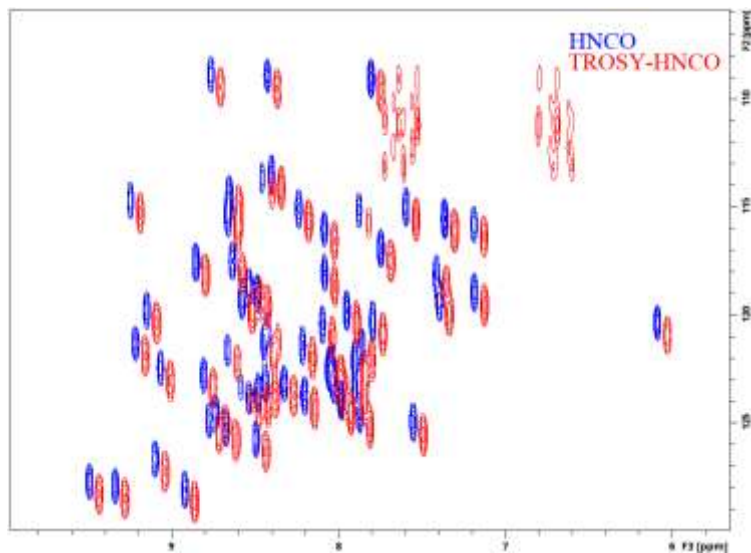
185 A specific module separate from the proximal-distal type can also be based on the same structure and either manually
186 included in the pulse program after each general module or in a specific module itself - se.incl is module implementing the
187 sensitivity-enhanced COS-INEPT and TROSY transfers and is imported in both the hsqc_se.incl and hx.incl modules.

188 4 Application examples

189 4.1 HNCO

```
pruscl relations=<me>
# include <Avance.incl>
# include <Grad.incl>
# define DIMS 3
/*Select options for distal and proximal blocks:*/
# define MH
# define HX
# define DISTAL_M
# define DISTAL_Y_CO
# define DISTAL_A_CA
# define PROXIMAL_MH
# define PROXIMAL_Y_CO
# define PROXIMAL_A_CA
; Variable definitions for the distal (H->N) and proximal (N->H) blocks:
# include <PE/includes/init.incl>
i ze
; Relaxation and distal block Hz -> NHz -> COHz:
# include <PE/includes/start.incl>
; 200Hz CO evolution (T1):
(CO_excitation(phFree1)):fCO
T1*0.5
(center (CA_CO_inversion(ph0)):fCA (N_inversion(ph0)):fN)
T1*0.5
(CO_refocussing(ph0)):fCO
(CA_CO_inversion(ph0)):fCA ; BSP compensation.
(CO_T1ipback(ph0)):fCO
GRAB(gpFree1)
; Proximal block COHz -> NHz-> H and acquisition:
# include <PE/includes/end.incl>
F1PH(calph(phFree1,+90), caldel(T1, +in1))
exit
# include <PE/includes/phasecycles.incl>
phFree1 = 0 0 0 0 2 2 2 2
; Receiver phases:
phRec = PROXIMAL_PHS1 + DISTAL_PHS1 + phFree1
;gpzFree1: gradient after CO echo: 21%.
;gpnanFree1: SMSQ10.100
```

191 Fig. 1. Pulse program code for the implementation of the HNCO experiment.

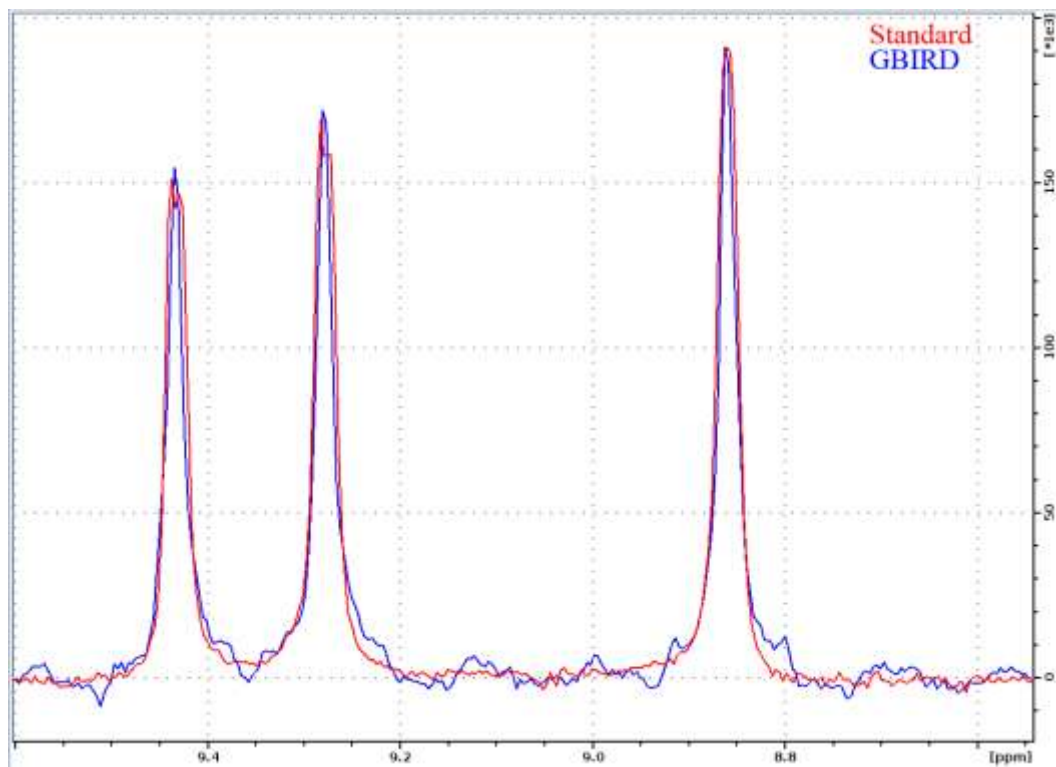


192
 193 **Fig. 2. Experimental demonstration of the implementations of the HNC(O) and TROSY-HNC(O) experiments for ubiquitin 8 (kDa) at**
 194 **25 °C. Spectra were recorded as ^1H - ^{15}N planes with maximum evolution times of 85.2 ms (^1H) and 9.87 ms (^{15}N) and processed**
 195 **using cosine squared window functions.**

196 | -HNC(O) is one of the simplest triple-resonance experiments and thus a good candidate to demonstrate the strengths and
 197 | limitations of the presented approach to library building. We present its ME NMR implementation in Fig. 1. We use a
 198 | custom prosol file (used mostly for automatic precalculation of pulse parameters) to free up a number of variables. Evolution
 199 | delays and increments are defined explicitly due to the proximal module's numbered variables (here $td2$ and $in2$) being
 200 | dimensionality-dependant. The block of defines specifies options for ME library - specifying the proximal (xh.incl) and
 201 | distal (hx.incl) modules and the couplings to be evolved (Y is $^2J_{\text{NCO}}$) and decoupled (A is $^2J_{\text{NCA}}$). After importing the first two
 202 | general modules, which includes the distal modules, two spin-echo evolution periods, the carbonyl echo is implemented
 203 | using the library's low-level functionality. Since ~~the~~ channels and pulses aren't selected explicitly explicitly, the sequence
 204 | this block will function with split CA and CO channels (with the right spectrometer configurations and "CACO_SPLIT"
 205 | defined in ZGOPTNS) or using a single carbon channel and frequency-offset pulses. The rest of the pulse program includes
 206 | the end.incl module (with the two proximal echoes and acquisition) and standard configuration of gradients and phasecycles.
 207 | To demonstrate the libraries functionality in Fig 2. we present 2D spectra (recorded as HN(CO) experiments) of a standard
 208 | variant of the experiment (no ZGOPTNs) and a TROSY-HNC(O) (adding the TROSY define to ZGOPTNS) selecting only the
 209 | H_{β} and N_{β} component (the lower right component using standard display convention). It's possible to choose a ^{13}C -GBIRD^{r,x}
 210 | appending the "ACQ_BIRD_C" option to ZGOPTNS, with an example of line narrowing demonstrated in Fig. 3.

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211

212 Fig. 3. 1D slices (for $N = 128.5$ ppm) through ^1H - ^{15}N planes recorded for a TROSY-HNCO with standard acquisition and
 213 TROSY-HNCO with ^{13}C -GBIRD $^{\text{r-s}}$ demonstrating the effectiveness of the homodecoupling and the resultant line narrowing. Both
 214 spectra were acquired for ubiquitin 8 (kDa) at 25 °C with maximum evolution times of 340.7 ms (^1H) and 9.87 ms (^{15}N) and
 215 processed using a cosine squared window function in the N dimension and sine 2 squared shifted by $\frac{\pi\pi}{2}$ in the H dimension. The
 216 GBIRD spectrum was shifted right by 4 Hz (shift was possibly induced by sample heating) and scaled up to match the amplitude of
 217 the standard TROSY-HNCO. For the GBIRD spectrum, 18 chunks were acquired with a 11.96 ms inter-chunk delay, 3.5 ms $^2J_{\text{HC}}$
 218 evolution time and using a 120 μs BIP-720-100-10 (Smith et al., 2001) pulse for ^{13}C inversion. Linewidths at half height are (from
 219 left to right) 19.6 Hz, 19.5 Hz and 19.1 Hz for the standard spectrum and 13.2 Hz, 13.2 Hz and 13.7 Hz for the homodecouple
 220 spectrum (TopSpin peakw function).

221

222

```

prosol relations=<me>

# include <Avance.incl>
# include <Grad.incl>

# define NOESY
# define DIMS 4

; Variable definitions and calculations for the proximal and distal 2Ds:
# include <ME/includes/init.incl>

define delay mixTime
;d10: NOESY mixing time [40-400 ms]
"mixTime = d10 - pGRAD - dGRAD" ; Corrected for gradient.

1 ze

; Distal 2D:
# include <ME/includes/start.incl>

; NOESY mixing:
# ifdef MIX_LOCKED
(
  refalign (mixTime):fH
  lalign (1m 4u BLKGRAD):fH
  ralign (2m UNBLKGRAD):fH
)
# else
  mixTime
# endif

  GRAD(gpNOESY)

; Proximal 2D and acquisition:
# include <ME/includes/end.incl>
exit

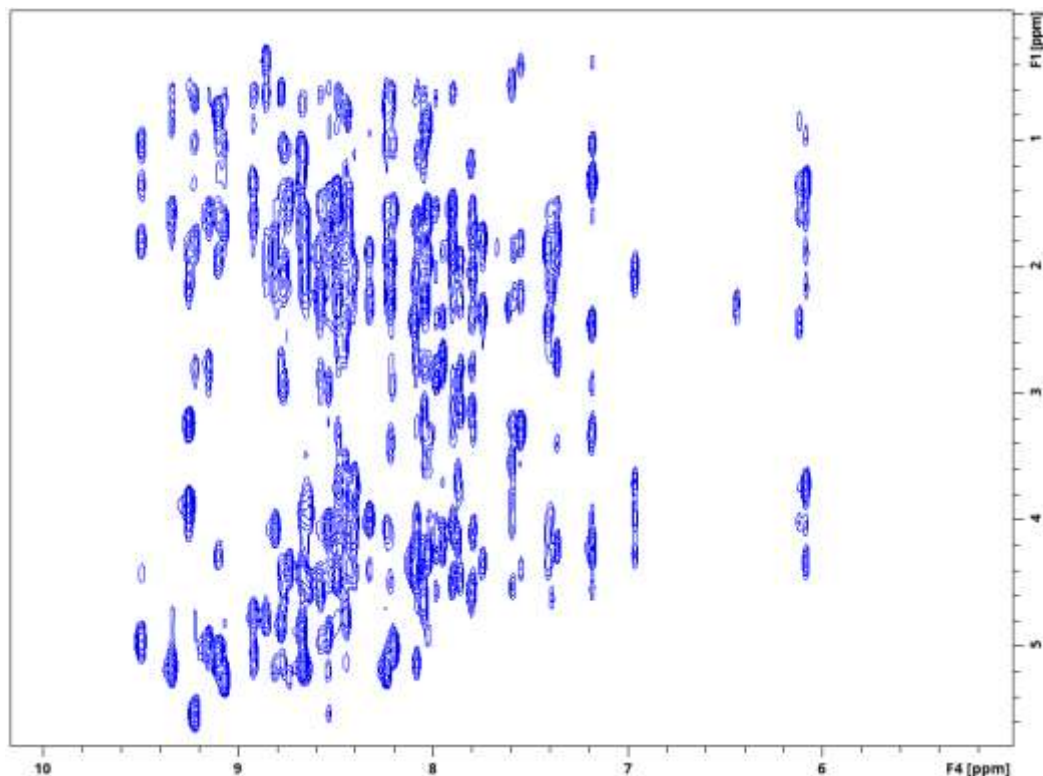
# include <ME/includes/phasecycles.incl>

; Receiver phase:
phRec = PROXIMAL_PH31 + DISTAL_PH31

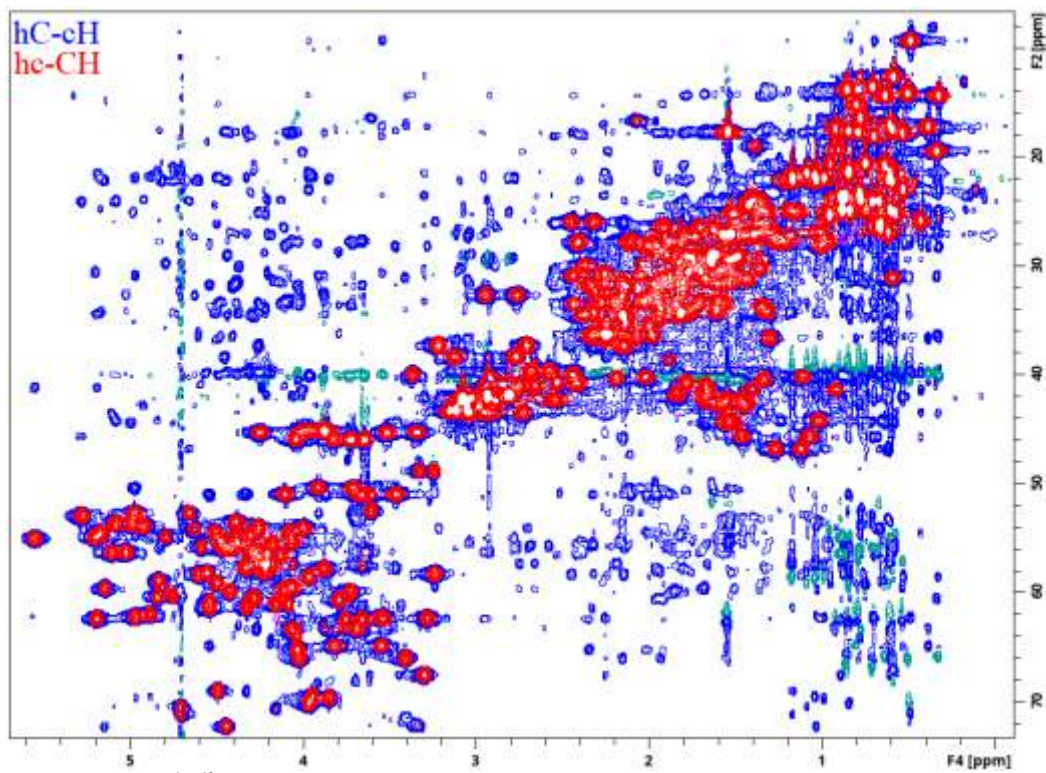
;gpzNOESY: gradient after NOESY: -7%.
;gpnamNOESY: SMSQ10.100

```

226 | -The modular nature of the library is exemplified by the 4D NOESY pulse program in Fig. 4. Apart ~~of~~ from the basic
227 | structure described above in the case of HNCO it only contains a mixing period joining the proximal and distal module, with
228 | the evolved heteronuclei and experiment types selected by the user using ZGOPTNS. A HC,NH-HMQC-NOESY-HSQC
229 | with sensitivity enhancement in the last dimension (Fig. 5.) can be changed to a HC,CH-HMQC-NOESY-HSQC (Fig. 6.)
230 | pulse program by changing the “PROXIMAL_N” option to “PROXIMAL_C” and adding the gradient selection option
231 | (“GS”, which isn’t a default for non-sensitivity-enhanced HSQC.
232

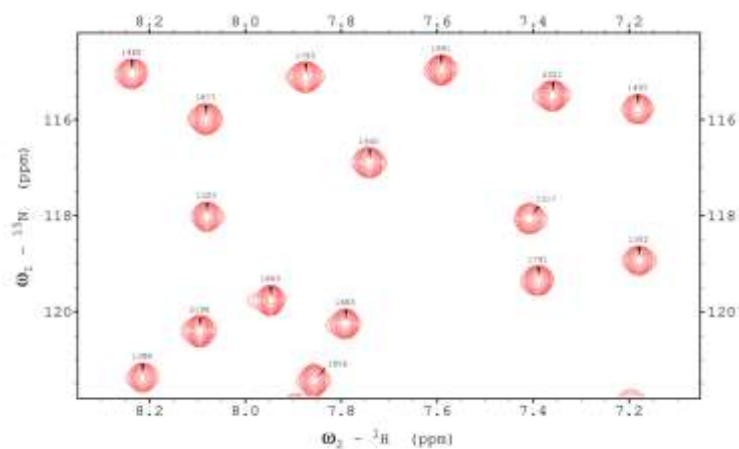


233 |
234 | **Fig. 5.** ^1H - ^1H planes recorded using a 4D HC,NH-HMQC-NOESY-HSQC experiment for ubiquitin 8 (kDa) at 25 °C. Spectra were
235 | recorded with maximum evolution times of 85,2 ms (^1H direct dimension) and 6.99 ms (^1H indirect dimension) and processed using
236 | cosine squared window functions.



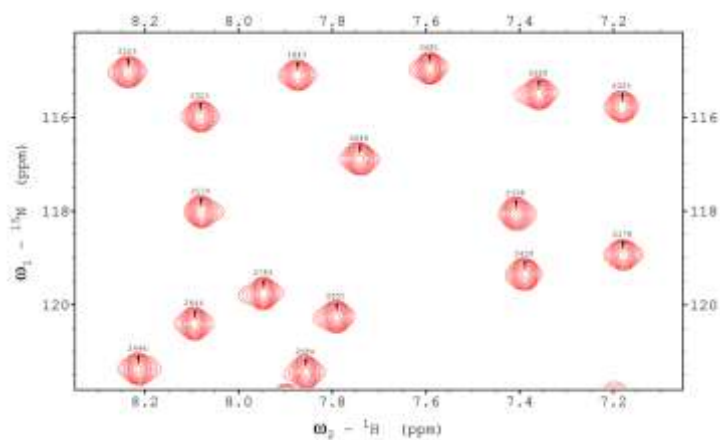
237
 238 Fig. 6. Two different ^1H - ^{13}C 2D planes recorded using a ME implementation of a 4D HC,CH-HMQC-NOESY-HSQC experiment
 239 for ubiquitin 8 (kDa) at 25 °C. Spectra were recorded with maximum evolution times of 85.2 ms (^1H direct dimension) and 7.96 ms
 240 (both ^{13}C dimensions) and processed using cosine squared window functions.

241 4.3 ^1H - ^{15}N correlation – shaped pulses



242

243 Fig. 7. ^1H , ^{15}N TROSY spectrum recorded using a ME implementation with hard pulses and water flipback for ubiquitin 8 (kDa) at
244 25 °C. The spectrum was recorded with maximum evolution times of 85,2 ms (^1H) and 39,5 ms (^{15}N) and processed using cosine
245 squared window functions.



246

247 Fig. 8. ~~Fig. 7.~~ ^1H , ^{15}N TROSY spectrum recorded using a ME implementation with shaped pulses (E400B and RE-BURP) for
248 ubiquitin 8 (kDa) at 25 °C. The spectrum was recorded with maximum evolution times of 85,2 ms (^1H) and 39,5 ms (^{15}N) and
249 processed using cosine squared window functions.

250

251 Since BEST-type experiments utilizing shaped pulses can bring improved sensitivity ~~even~~-especially at higher scan
252 repetition rates (Schanda et al., 2006) ~~†~~ we demonstrate the library's inherent ability to automatically adapt to the substantial
253 chemical shift and coupling evolution during the 90-degree universal rotation E400B (Veshtort and Griffin, 2004) (using a
254 time-reversed version of the original pulse for excitation) pulses with the length of 1073.1 μ s (equivalent to an ideal pulse
255 followed by a 611.7 μ s delay) and refocussing pulse RE-BURP (Geen and Freeman, 1991) ~~-~~with length of 1108.8 μ s
256 (modelled as an ideal refocussing pulse flanked by 554 μ s delays) in Fig. 7. and 8. With a relaxation delay of 0.65 s all peaks
257 in the selected region are over 20% stronger in the shaped pulse version. ~~-~~Full datasets for a number of different relaxation
258 delays are provided, as in the data availability section.

259 **5 Materials & methods**

260 For all experiments we used a 2 mM ^{13}C , ^{15}N -double labelled human ubiquitin (ASLA Biotech) in a 5 mm Shigemi NMR
261 microtube. All spectra were acquired using a Bruker Avance IIIHD 800 MHz spectrometer with a 5 mm TCI z-gradient
262 cryo-probe. Pulse lengths for 90 degree hard pulses were 10.47 μ s for ^1H , 12.3 μ s for ^{13}C and 33.22 μ s for ^{15}N . Full
263 acquisition and processing parameters are provided in the dataset linked below in the Data availability section. Acquisition
264 and library testing was, performed using the TopSpin 3.6.5 Service Pack 2 software (Bruker). Data processing and plotting
265 (aside from Fig. 7. and 8.) was carried out in TopSpin. Figures 7 and 8 were prepared using the NMRFAM-SPARKY
266 software (Goddard and Kneller, 2004; Lee et al., 2015).

267 **6 Conclusions**

268 We have described a framework library implementing a two-level approach to pulse program modularization and
269 demonstrated its utility. We hope it can be used by others either directly for the streamlining of pulse program code or as an
270 inspiration for similar frameworks. Although the usefulness of the modularization approach is most obvious for the case of
271 protein experiments presented here, it should extend to nucleic acids and, to a more limited extent, small molecules. In the
272 latter case, the ability to modularize preparation period operations (presaturation, ASAP), WATERGATE (Piotto et al.,
273 1992; Sklenar et al., 1993) type solvent suppression and real-time acquisition should be particularly useful.

274 **Code availability**

275 The initial version of the ME library [code and documentation](https://doi.org/10.5281/zenodo.1057868) is available online at:
276 <https://doi.org/10.5281/zenodo.1057868> <https://doi.org/10.5281/zenodo.10841749>. Current library version is available
277 [online at https://github.com/nmr-cnbc/MEnmr_pubcode](https://github.com/nmr-cnbc/MEnmr_pubcode) or from the authors upon request.

278 **Data availability**

279 All data used in the preparation of this article is available online at: <https://doi.org/10.5281/zenodo.10578330>.

280 **Author contributions**

281 MG and WK designed the general workflow of the ME library. MG wrote the library code and performed the experiments.

282 MG wrote the manuscript with input from WK.

283

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286 **Competing interests**

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

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