

# FID-A: The FID Appliance

## Version 1.0β2

### User Manual

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## 1. Introduction

The FID Appliance (FID-A) is an open-source software package for simulation of MRS experiments, design and analysis of radiofrequency (RF) pulses, and processing of MRS data. The software is freely available for download ([www.github.com/CIC-methods/FID-A](http://www.github.com/CIC-methods/FID-A)).

The FID-A software package consists of four separate toolboxes:

- The **Simulation Toolbox** for simulation of in-vivo MRS experiments,
- The **RF-pulse Toolbox**, for designing and simulating radiofrequency pulse waveforms,
- The **Input-output Toolbox**, for reading and writing data between MATLAB and other useful data formats,
- and The **Processing toolbox**, for processing of in-vivo MRS data.

In addition to the toolboxes listed above, FID-A also comes with a library of **Example Run Scripts**, which provide examples of useful “pipelines” for NMR simulation and data processing. The example run scripts each make use of combinations of the various functions within the FID-A toolboxes. Finally, although the FID-A toolbox is primarily MATLAB command-line based, a few **graphical user interfaces** are also provided to assist with processing tasks in which visual feedback is required, such as manual phasing of spectra, and visual alignment of spectra or sub-spectra prior to subtraction.



The FID-A toolboxes are each described briefly below:

### **1.1. Simulation Toolbox**

The FID-A NMR simulation toolbox is based on an implementation of the density matrix formalism, where the evolution of the spin system in a given NMR experiment is described by successive evolutions of the density matrix by time-independent Hamiltonian operators.

The simulation toolbox contains built-in functions for simulating the basic components of an MRS pulse sequence, including excitation (`sim_excite.m`), delays (`sim_evolve.m`), rotation using ideal (`sim_rotate.m`) and shaped (`sim_shapedRF.m`) radiofrequency pulses, and signal readout (`sim_readout.m`). Using these basic pulse sequence elements, common in-vivo MRS pulse sequences are also implemented, including the FID (`sim_onepulse.m`), PRESS (`sim_press.m`), STEAM (`sim_steam.m`) and spin-echo (`sim_spinecho.m`) sequences.

Simulation is performed on a given spin-system by specifying the pulse timings and the chemical shifts and coupling constants of the spin-system of interest. A full set of common metabolite spin-system definitions is provided based on values previously published by Govindaraju et al (NMR Biomed 2000, 13:129-153). Spin system definitions can be found in `{FID-A_path}/simulationTools/metabolites/`, where `{FID-A_path}` is the path to the root folder of the FID-A software package. Each metabolite's spin system definition is contained within a structure with two fields: "shifts" and "J". Shifts is a vector of length N, which contains the chemical shift value (in ppm) for each of the N spins in the spin system. J is an N x N matrix specifying the coupling constant (in Hz) between each spin and every other spin in the system. For spin systems with more than 6 or 7 spins, the spin system is broken down into multiple independent (uncoupled) subgroupings with fewer spins. This is because the computation time increases more than linearly with increasing number of spins, and so

it is much more computationally efficient to simulate, say two groups of spins with 6 protons each rather than simulating one group of 12 protons. Finally, some spin systems have been reduced in size due to redundancy. For example, scyllo inositol consists of nine magnetically equivalent protons, and is therefore most easily simulated as a single proton. However, the user must then remember to scale the intensity of Scyllo Inositol Simulations by a factor of 9. Such an amplitude scaling can easily be done using one of the tools from the FID-A processing toolbox. The example below shows how one can easily generate and display a scyllo inositol spectrum using just four lines of code:

```
load Scyllo; %Load the spin system
scyllo = sim_onepulse(2048,2000,3,6,sysScyllo); %Simulate
scyllo_scaled = op_ampScale(scyllo,9); %Do amplitude scaling
op_plotspec({scyllo,scyllo_scaled}); %Plot
```

This example highlights one of the nice features of the FID-A software package; namely that the operations in the processing toolbox can be applied to both simulated data as well as real experimental data.

In FID-A simulations, excitation and refocusing RF pulses can be modeled as ideal (instantaneous) rotations, or fully shaped RF waveforms, depending on the user requirements. In the case of shaped RF pulses, phase cycling is performed to remove unwanted coherences. Furthermore, in the case of a shaped slice selective pulse, a single simulation corresponds to one point in space, which must be specified relative to the centre of the slice selective pulse. In order to simulate the result of a full in-vivo MRS experiment involving shaped localization pulses, it is necessary to run the simulation over many points in space (with phase cycling), and then combine the results. Function names in the simulation toolbox begin with the prefix "sim\_".

## **1.2. RF Pulse Toolbox**

The FID-A RF toolbox enables the creation of basic RF pulse waveforms, and Bloch simulation to determine the resulting excitation/refocusing/inversion profiles, as well as frequency shifting and resampling of rf waveforms. RF pulses are stored in MATLAB structure format, with fields corresponding to the RF waveform, the type of rf pulse (excitation, inversion, refocusing), the time-bandwidth product of the rf pulse, and the time-B1 product. The time-bandwidth product and time-b1 product of the rf pulse are calculated automatically when the rf pulse is initialized using the `io_loadRFwaveform.m` function. These values can then be used to calculate the required B1 amplitude to achieve a certain flip angle in bloch simulations (see `rf_blochSim.m`) or to calculate the required gradient strength for slice selection (see `run_simMegaPressShaped.m`). Function names in the RF-pulse toolbox begin with the prefix "rf\_".

## **1.3. Input-Output Toolbox**

The FID-A Input-Output toolbox contains "load" functions to accept MRS data in MRI vendor data formats (Siemens, Agilent, Philips, GE), and store them in MATLAB. It also contains "load" functions to accept MRS data from other data processing or analysis software packages (LCModel, jMRUI), and "write" functions to export MRS data back into those formats. Finally, the Input-Output toolbox also contains "load" functions to accept radiofrequency pulse waveforms in MRI vendor data formats (Siemens, Agilent), and "write" functions to export newly designed RF pulses back into vendor formats for use on the scanner. All of the functions in the Input-Output toolbox begin with the prefix "io\_".

## 1.4. Processing Toolbox

Once data has been loaded into MATLAB using the FID-A Input-Output toolbox, the data can then be operated on using any of the over 50 different processing operations, including (but not limited to) filtering (`op_filter.m`), zeropadding (`op_zeropad.m`), time domain truncation (`op_leftshift.m` and `op_zerotrim.m`), frequency domain truncation (`op_fregrange.m`), eddy current correction (`op_ecc.m`), removal motion corrupted averages (`op_rmbadaverages.m`), retrospective frequency and phase drift correction (`op_alignAverages.m` and `op_alignAverages_fd.m`, Near et al, Magn Reson Med 2014, DOI: 10.1002/mrm.25094), combination of multi-element RF coil data (`op_addrcvrs.m`), and zero- and first-order phase corrections (`op_addphase.m`).

These functions can be nested within one another. For example, to load a spectrum, combine the receivers, combine the averages and then filter the result, can be done using the following single line of code:

```
out= op_filter(op_averaging(op_addrcvrs(op_loadspec_twix('filename.dat'),1,'w'),5);
```

where the argument "5" represents a 5 Hz exponential filter, the argument "'w'" specifies that a weighted coil recombination should be performed, and the argument "1" specifies that the phases and amplitudes of the rf coil channels should be determined using the first point in the time domain.

Function names in the processing toolbox begin with the prefix "op\_", which stands for operator.

## 1.5. Example Run Scripts

The {FID-A\_dir}/exampleRunScripts directory contains a few examples of full MRS data processing pipelines. These include:

run\_specialproc.m, run\_pressproc.m, and run\_megapressproc.m. These “pipeline” operations begin with raw MRS data in Siemens .dat format and then process the data in a logical step-by-step fashion to generate fully processed data that is ready to be analyzed using one of the leading MRS analysis software packages (LCModel, jMRUI or Tarquin). Processing steps performed in these pipelines include combination of receive channels (op\_addrcvrs.m), removal of motion corrupted averages (op\_rmbadaverages.m), spectral registration of averages (op\_alignAverages.m), and combination of averages (op\_averaging.m).

In addition to processing pipelines, the {FID-A\_dir}/exampleRunScripts directory contains some examples of how to run complicated simulations involving pulse sequences with shaped rf pulses (run\_simMegaPressShaped.m, run\_simMegaPressShapedEdit.m, run\_simMegaPressShapedRefoc.m, run\_simMegaSpecialShaped.m, run\_simSpinEchoShaped.m). Finally, the {FID-A\_dir}/exampleRunScripts directory contains an example script showing how to generate a basic LCModel basis set (run\_simExampleBasisSet.m).

## **1.6. Data structure formatting**

The FID-A software package is implemented in MATLAB (Natick MA, USA). Within FID-A, simulated or experimental MRS datasets are stored in uniquely formatted data structures that encapsulate all of the data, (in both the time-domain and frequency-domain) as well as the relevant header information in the fields of the structure. By encapsulating data and header information in this way, processing operations have access to all relevant information and thus require as few input arguments as possible.

FID-A is unique among MRS processing software tools in that it enables the user to easily and efficiently manage MRS datasets with higher dimensionality (datasets with multiple averages, multiple coils, and multiple subspectra). When multiple averages, coil

channels, or subspectra of data are present, they are stored in separate dimensions of a data arrays, and indexed within the header. The FID-A processing operations are designed to automatically recognize the dimensionality of the data based on the header information and perform their operations accordingly. The fields of the MRS data structure are listed and briefly described below:

```

fids      - time domain MRS data.
specs     - frequency domain MRS data.
t         - vector of time values for plotting in the time domain [s]
ppm       - vector of frequency values for plotting in the frequency domain [ppm]
sz        - size of the fids and specs arrays
date      - date that the data was acquired or simulated
averages  - number of averages in the dataset (possibly altered by processing)
rawAverages - number of averages in the original dataset (not altered by
              processing).
subspecs  - number of subspectra (ISIS, edit on/off, etc) in the dataset
              (possibly altered by processing).
rawSubspecs - number of subspectra (ISIS, edit on/off, etc) in the original dataset
              (not altered by processing).
Bo        - magnetic field strength [Tesla]
txfrq     - Centre frequency [MHz];
linewidth - linewidth of data (only used for simulated data) [Hz]
n         - number of spectral points
dwelltime - dwell time of the data in the time domain [s] (dwelltime =
              1/spectralwidth)
sim       - type of simulation (ideal vs. shaped pulses), only used for simulated
              data.
te        - echo time of acquisition [ms], only used for simulated data
seq       - type of sequence used (only used for simulated data).
dims      - structure specifying which data dimensions are stored along which
              dimensions of the fids/specs arrays. Fields include:
    t      - time/frequency dimension (usually this is 1, the first
              dimension of the fids/specs array).
    coils  - for multiple receiver array, this is the dimension of the
              arrayed receiver data (can be 2, 3 or 4).
    averages - for multiple averages, this is the dimension of the averages
              (can be 2, 3 or 4).
    subSpecs - in the case of subtraction data (ISIS, MEGA-PRESS), this is

```

the dimension of the subSpectra (can be 2, 3 or 4).

flags - structure specifying what processing operations have already been done on the data. fields include:

- writtentostruct - Has the dataset been written to a structure (1 or 0)
- gotparams - Have the parameters been retrieved from the dataset (1 or 0)
- filtered - Has the dataset been filtered (1 or 0)
- zeropadded - Has the dataset been zeropadded (1 or 0)
- freqcorrected - Has the dataset been frequency corrected (1 or 0)
- phasescorrected - Has the dataset been phase corrected (1 or 0)
- averaged - Have the averages been combined (1 or 0)
- addedrcvrs - Have the rcvr channels been combined (1 or 0).
- Subtracted - Have the subspecs been subtracted (1 or 0)
- Writtentotext - Has the data been writte to text file (1 or 0)
- Downsampled - has the data been resampled to a different spectral resolution (1 or 0)
- avgNormalized - Has the data been amplitude scaled following combination of the averages (1 or 0)
- isISIS - Does the dataset contain ISIS subspectra (1 or 0)

RF pulses are also stored in uniquely formatted data structures. Each RF pulse structure stores the rf waveform as well as information about the rf pulse type (excitation, refocusing or inversion), the time-bandwidth product, and the time-B1 product. The fields of the RF pulse structure are listed and briefly described below:

- waveform - an n x 3 array, where n is the number of points in the rf pulse, the first column (:,1) contains the rf pulse phase, the second column (:,2) contains the absolute rf amplitude, and the third column (:,3) contains the duration of each time step (normally this is a vector of ones, but not necessarily).
- type - The type of RF pulse. Options are 'exc' for an excitation pulse, 'inv' for an inversion pulse, and 'ref' for a refocusing pulse.
- twl - The product of the duration of the rf pulse [s] and the w1max of the rf pulse [Hz].
- tbw - The product of the duration of the rf pulse [s] and the bandwidth of the rf pulse [Hz]

Below is a brief summary of all of the functions in the FID-A Software package. The same information can be obtained by typing 'help functionName" at the MATLAB command line:

## **2. Simulation Tools**

### **2.1. sim\_Hamiltonian.m**

**USAGE:**

```
[H,d] = sim_Hamiltonian(sys,Bfield);
```

**DESCRIPTION:**

Creates the nxn Hamiltonian matrix for a spin system which can then be used in other functions to simulate NMR experiments.

**INPUTS:**

```
sys          = spin system definition structure.  
Bfield       = magnetic field strength (Tesla).
```

### **2.2. sim\_evolve.m**

**USAGE:**

```
d_out = sim_evolve(d_in,H,t)
```

**DESCRIPTION:**

This function simulates free evolution of the spin system under the effects of chemical shift and scalar coupling.

**INPUTS:**

```
d_in        = input density matrix structure.  
H           = Hamiltonian operator structure.  
t           = duration of evolution (s)
```

### **2.3. sim\_excite.m**

**USAGE:**

```
d_out = sim_excite(H,axis,angle)
```

**DESCRIPTION:**

This function simulates the effect of an ideal (instantaneous) excitation pulse on the density matrix. Used in simulation tools.

**INPUTS:**

```
d_in        = input density matrix structure.
```



**H** = Hamiltonian operator structure.  
**axis** = Axis of rotation ('x' or 'y');  
**angle** = Flip angle of excitation (degrees). Optional. Default=90.  
 If angle is a scalar, then the same flip angle is applied to all spins in the spin system. If angle is a vector, then the elements of the vector specify the flip angles to apply to each spin in the system. In this case, the length of the vector must be the same as the number of spins in the spin system.

## 2.4. **sim\_excite\_arbPh.m**

### USAGE:

`d_out = sim_excite_arbPh(H,phase,angle)`

### DESCRIPTION:

This function simulates the effect of an ideal (instantaneous) excitation pulse on the density matrix. Used in simulation tools. The phase of the excitation pulse can be arbitrarily chosen. To achieve an arbitrary phase, this code executes a rotation about z by an angle of -phi (the rf pulse phase), then executes a rotation about x by an angle of "angle" (the flip angle of the excitation pulse), and then executes a rotation back about z by an angle of phi.

### INPUTS:

**d\_in** = input density matrix structure.  
**H** = Hamiltonian operator structure.  
**phase** = Phase of rotation in degrees (ie. 0='x', 90='y', etc);  
**angle** = Flip angle of excitation (degrees). Optional. Default=90. If angle is a scalar, then the same flip angle is applied to all spins in the spin system. If angle is a vector, then the elements of the vector specify the flip angles to apply to each spin in the system. In this case, the length of the vector must be the same as the number of spins in the spin system.

## 2.5. **sim\_gradSpoil.m**

### USAGE:

`d_out = sim_gradSpoil(d_in,H,gradVect,dur,posVect)`

### DESCRIPTION:

This function simulates the effect of a rectangular spoiler gradient with a given amplitude, direction and duration.

### INPUTS:

**d\_in** = input density matrix structure.  
**H** = Hamiltonian operator structure.  
**gradVect** = Vector of spoiler gradient amplitudes [Gx Gy Gz] in G/cm.  
**posVect** = Position vector of spins of interest [x y z] in cm.  
**dur** = Duration of the gradient pulse in ms.

## 2.6. `sim_lcmrawbasis.m`

### USAGE:

```
[RF,out]=sim_lcmrawbasis(n,sw,Bfield,linewidth,metab,tau1,tau2,addref, makeraw,seq)
```

### DESCRIPTION:

Generate an LModel .RAW file to be used as an individual metabolite basis spectrum in an LModel basis set. The relevant characteristics of the acquisition can be specified (pulse sequence, number of points, spectral width, etc)

### INPUTS:

```
n          = number of points in fid/spectrum
sw         = desired spectral width in [Hz]
Bfield     = main magnetic field strength in [T]
linewidth  = linewidth in [Hz]
tau1       = first echo time in [s] (if seq='st', tau1 = TE)
tau2       = second echo time in [s]. (Used in Press, but not used in S      (If
            seq='st', tau2=TM).
addref     = add reference at 0ppm (for use in LModel makebasis) ['y' or 'n']
makeraw    = make output file for lmodel ['y' or 'n']
seq        = pulse sequence ['se' for Spin Echo or 'p' for Press]
metab      = one of the following choices
    'H2O'   = Water
    'Ala'   = Alanine
    'Asp'   = Aspartate
    'PCh'   = PhosphoCholine
    'Cr'    = Creatine
    'PCr'   = PhosphoCreatine
    'GABA'  = Gamma-aminobutyric acid (kaiser)
    'GABA3' = Gamma-aminobutyric acid (de Graaf)
    'Gln'   = Glutamine
    'Glu'   = Glutamate
    'GSH'   = Glutathione
    'Gly'   = Glycine
    'Ins'   = Myo-inositol
    'Lac'   = Lactate
    'NAA'   = N-acetyl aspartate
    'Scyllo' = Scyllo-inositol
    'Tau'   = Taurine
    'Asc'   = Ascorbate (Vitamin C)
    'bHB'   = beta-Hydroxybutyrate
    'bHG'   = beta-Hydroxyglutarate
    'Glc'   = Glucose
    'NAAG'  = N-acetyl aspartyl glutamate
    'GPC'   = Glycero-phosphocholine
    'PE'    = Phosphoryl ethanolamine
    'Ser'   = Serine
```

## 2.7. `sim_make2DSimPlot.m`

### USAGE:

```
sim_make2DSimPlot(in,ppmmin,ppmmax,plotDiff)
```

### DESCRIPTION:

This function takes the output of a spatially resolved simulation, and plots the array of spectra on a single figure. The input should be a cell array where each element of the cell array is a simulated spectrum from one spatial position in the spatially resolved simulation. Each element of the array is also in FID-A data structure format. By including the optional input argument ppmmin and ppmmax, only a the corresponding range of each spectrum will be plotted. If the

**INPUTS:**

in = input cell array of simulated spectra from a spatially resolved simulation  
 ppmmin = lower limit of ppm range to plot [ppm]  
 ppmmax = upper limit of ppm range to plot [ppm]

**2.8. sim\_megapress.m****USAGE:**

out=sim\_megapress(n,sw,Bfield,linewidth,sys,taus,refoc1Flip,refoc2Flip, editFlip)

**DESCRIPTION:**

Simulate the MEGA-PRESS sequence with instantaneous localization and editing pulses. Provides the ability to specify the flip angle of each refocusing pulse and editing pulse on each spin in the spin system.

**INPUTS:**

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 sys = spin system definition structure  
 taus = pulse sequence timing vector:  
   taus(1) = time in [ms] from 90 to 1st 180  
   taus(2) = time in [ms] from 1st 180 to 1st edit pulse  
   taus(3) = time in [ms] from 1st edit pulse to 2nd 180  
   taus(4) = time in [ms] from 2nd 180 to 2nd edit pulse  
   taus(5) = time in [ms] from 2nd edit pulse to ADC  
 refoc1Flip= array of refoc1 flip angles for each spin in system  
 refoc2Flip= array of refoc2 flip angles for each spin in system  
 editFlip = array of editing flip angles for each spin in system

**2.9. sim\_megapress\_shaped.m****USAGE:**

sim\_megapress\_shaped(n,sw,Bfield,linewidth,taus,sys,editPulse,editTp,editPh1, editPh2,refPulse,refTp,Gx,Gy,dx,dy,refPh1,refPh2)

**DESCRIPTION:**

This function simulates the MEGA-PRESS sequence with shaped localization pulses and shaped editing pulses. Enables choice of the timings of all of the rf pulses as well as the choice of the phase of both the editing pulse and the refocusing pulses. This allows phase cycling of the editing and refocusing pulses by repeating simulations with different editing pulse phases, which is necessary to remove phase artefacts from the editing pulses. For the editing pulses, an eight step phase cycling scheme is typically sufficient, where by the first editing pulse is cycled by 0 and 90 degrees, and the second editing pulse is cycled by 0,90,180, and 270 degrees, and all phase cycles should be added together to remove unwanted coherences. For the refocusing pulses, a four step phase cycling scheme is typically sufficient, where both refocusing pulses are phase cycled by 0 and 90 degrees, and the phase are combined in the following way:

signal = ([0 90] - [0 0]) + ([90 0] - [90 90]);

where, in [X Y], X is the phase of the first refocusing pulse and Y is the phase of

the second refocusing pulse

Note that this code only simulates one subspectrum at a time (edit-on or edit-off). The difference spectrum can be obtained by simulating one of each, and then subtracting.

INPUTS:

```
n          = number of points in fid/spectrum
sw         = desired spectral width in [Hz]
Bfield     = main magnetic field strength in [T]
linewidth  = linewidth in [Hz]
taus(1)    = time in [ms] from 90 to 1st 180
taus(2)    = time in [ms] from 1st 180 to 1st edit pulse
taus(3)    = time in [ms] from 1st edit pulse to 2nd 180
taus(4)    = time in [ms] from 2nd 180 to 2nd edit pulse
taus(5)    = time in [ms] from 2nd edit pulse to ADC
            FOR MEGA-PRESS on SIEMENS SYSTEM:
            taus=[4.545,12.7025,21.7975,12.7025,17.2526];
editPulse  = Editing pulse shape - [Nx3] array with [:,1]=phase, [:,2]=amplitude,
            [:,3]=duration
editTp     = duration of editing pulse in [ms];
sys        = Metabolite spin system definition structure;
editPh1    = the phase of the first editing pulse in [degrees];
editPh2    = the phase of the second editing pulse in [degrees];
refPulse   = Refocusing pulse shape - [Nx3] array with [:,1]=phase,
            [:,2]=amplitude, [:,3]=duration
refTp      = duration of refocusing pulse in [ms]
Gx         = gradient strength for first selective refocusing pulse [G/cm]
Gy         = gradient strength for second selective refocusing pulse [G/cm]
dx         = position offset in x-direction (corresponding to first refocusing
            pulse) [cm]
dy         = position offset in y-direction (corresponding to second refocusing
            pulse) [cm]
refPh1     = the phase of the first refocusing pulse in [degrees];
refPh2     = the phase of the second refocusing pulse in [degrees];
```

## 2.10. **sim\_megapress\_shapedEdit.m**

USAGE:

```
sim_megapress_shapedEdit(n,sw,Bfield,linewidth,taus,sys,editPulse,editTp,
                        editPh1,editPh2)
```

DESCRIPTION:

This function simulates the MEGA-PRESS sequence with instantaneous localization pulses and shaped editing pulses. Enables choice of the timings of all of the rf pulses as well as the choice of the phase of the editing pulse. This allows phase cycling of the editing pulses by repeating simulations with different editing pulse phases, which is necessary to remove phase artefacts from the editing pulses. For the editing pulses, an eight step phase cycling scheme is typically sufficient, where by the first editing pulse is cycled by 0 and 90 degrees, and the second editing pulse is cycled by 0,90,180, and 270 degrees, and all phase cycles should be added together to remove unwanted coherences.

Note that this code only simulates one subspectrum at a time (edit-on or edit-off). The difference spectrum can be obtained by simulating one of each, and then subtracting.

INPUTS:

```
n          = number of points in fid/spectrum
```

```

sw          = desired spectral width in [Hz]
Bfield      = main magnetic field strength in [T]
linewidth   = linewidth in [Hz]
taus(1)     = time in [ms] from 90 to 1st 180
taus(2)     = time in [ms] from 1st 180 to 1st edit pulse
taus(3)     = time in [ms] from 1st edit pulse to 2nd 180
taus(4)     = time in [ms] from 2nd 180 to 2nd edit pulse
taus(5)     = time in [ms] from 2nd edit pulse to ADC
              FOR MEGA-PRESS on SIEMENS SYSTEM:
              taus=[4.545,12.7025,21.7975,12.7025,17.2526];
editPulse   = Editing pulse shape - [Nx3] array with [:,1]=phase, [:,2]=amplitude,
              [:,3]=duration
editTp      = duration of editing pulse in [ms];
sys         = Metabolite spin system definition structure;
editPh1     = the phase of the first editing pulse in [degrees];
editPh2     = the phase of the second editing pulse in [degrees];

```

## 2.11. **sim\_megapress\_shapedRefoc.m**

### USAGE:

```

sim_megapress_shapedRefoc(n,sw,Bfield,linewidth,taus,sys,editFlip,refPulse,refTp,Gx
,Gy,dx,dy,refPh1,refPh2)

```

### DESCRIPTION:

This function simulates the MEGA-PRESS sequence with shaped localization pulses and instantaneous editing pulses. Enables choice of the timings of all of the rf pulses as well as the choice of the phase of both the editing pulse and the refocusing pulses. This allows phase cycling of the editing and refocusing pulses by repeating simulations with different editing pulse phases, which is necessary to remove phase artefacts from the editing pulses. For the editing pulses, an eight step phase cycling scheme is typically sufficient, where by the first editing pulse is cycled by 0 and 90 degrees, and the second editing pulse is cycled by 0,90,180, and 270 degrees, and all phase cycles should be added together to remove unwanted coherences. For the refocusing pulses, a four step phase cycling scheme is typically sufficient, where both refocusing pulses are phase cycled by 0 and 90 degrees, and the phase are combined in the following way:

$$\text{signal} = ([0 \ 90] - [0 \ 0]) + ([90 \ 0] - [90 \ 90]);$$

where, in [X Y], X is the phase of the first refocusing pulse and Y is the phase of the second refocusing pulse

Note that this code only simulates one subspectrum at a time (edit-on or edit-off). The difference spectrum can be obtained by simulating one of each, and then subtracting.

### INPUTS:

```

n          = number of points in fid/spectrum
sw         = desired spectral width in [Hz]
Bfield     = main magnetic field strength in [T]
linewidth  = linewidth in [Hz]
taus(1)    = time in [ms] from 90 to 1st 180
taus(2)    = time in [ms] from 1st 180 to 1st edit pulse
taus(3)    = time in [ms] from 1st edit pulse to 2nd 180
taus(4)    = time in [ms] from 2nd 180 to 2nd edit pulse
taus(5)    = time in [ms] from 2nd edit pulse to ADC
              FOR MEGA-PRESS on SIEMENS SYSTEM:
              taus=[4.545,12.7025,21.7975,12.7025,17.2526];
sys        = Metabolite spin system definition structure;

```

```

editFlip    = vector of editing flip angles in [degrees] at chemical shifts
              corresponding to 'shifts'.
refPulse    = Refocusing pulse shape - [Nx3] array with [:,1]=phase,
              [:,2]=amplitude, [:,3]=duration
refTp       = duration of refocusing pulse in [ms]
Gx          = gradient strength for first selective refocusing pulse [G/cm]
Gy          = gradient strength for second selective refocusing pulse [G/cm]
dx          = position offset in x-direction (corresponding to first refocusing
              pulse) [cm]
dy          = position offset in y-direction (corresponding to second refocusing
              pulse) [cm]
refPh1      = the phase of the first refocusing pulse in [degrees];
refPh2      = the phase of the second refocusing pulse in [degrees];

```

## 2.12. **sim\_megaspecial\_shaped.m**

### USAGE:

```

sim_megaspecial_shaped(n,sw,Bfield,linewidth,taus,sys,editPulse,editTp,
                      editPh1,editPh2,refPulse,refTp,Gx,dx,refPh)

```

### DESCRIPTION:

This function simulates the MEGA-SPECIAL sequence with a shaped localization pulse and shaped editing pulses. Enables choice of the timings of all of the rf pulses as well as the choice of the phase of both the editing pulses and the refocusing pulse. This allows phase cycling of the editing and refocusing pulses by repeating simulations with different editing pulse phases, which is necessary to remove phase artefacts from the editing pulses. For the editing pulses, an eight-step phase cycling scheme is typically sufficient, where by the first editing pulse is cycled by 0 and 90 degrees, and the second editing pulse is cycled by 0,90,180, and 270 degrees, and all phase cycles should be added together to remove unwanted coherences. For the refocusing pulse, a two step phase cycling scheme is typically sufficient, where the refocusing pulses is phase cycled by 0 and 90 degrees, and the two phase cycles are subtracted from each other.

Note that this code only simulates one subspectrum at a time (edit-on or edit-off). The difference spectrum can be obtained by simulating one of each, and then subtracting.

### INPUTS:

```

n           = number of points in fid/spectrum
sw          = desired spectral width in [Hz]
Bfield      = main magnetic field strength in [T]
linewidth   = linewidth in [Hz]
taus(1)     = time in [ms] from 90 to 1st edit pulse
taus(2)     = time in [ms] from 1st edit pulse to the 180
taus(3)     = time in [ms] from the 180 pulse to 2nd edit pulse
taus(4)     = time in [ms] from 2nd edit pulse to ADC
editPulse    = Editing pulse shape structure
editTp      = duration of editing pulse in [ms];
sys         = Metabolite spin system definition structure;
editPh1     = the phase of the first editing pulse in [degrees];
editPh2     = the phase of the second editing pulse in [degrees];
refPulse     = Refocusing pulse shape structure
refTp       = duration of refocusing pulse in [ms]
Gx          = gradient strength for selective refocusing pulse [G/cm]
dx          = position offset in x-direction (corresponding to refocusing pulse)
              [cm]
refPh       = the phase of the refocusing pulse in [degrees];

```

### 2.13. **sim\_onepulse.m**

**USAGE:**

```
out=sim_onepulse(n,sw,Bfield,linewidth,sys)
```

**DESCRIPTION:**

This function simulates a pulse-acquire experiment with an ideal (instantaneous) excitation pulse and an assumed lorentzian lineshape. The function calls the function 'sim\_Hamiltonian' which produces the free evolution Hamiltonian for the specified number of spins, J and shifts.

**INPUTS:**

```
n          = number of points in fid/spectrum
sw         = desired spectral width in [Hz]
Bfield     = main magnetic field strength in [T]
linewidth  = linewidth in [Hz]
sys        = spin system definition structure
```

### 2.14. **sim\_onepulse\_arbPh.m**

**USAGE:**

```
out=sim_onepulse_arbPh(n,sw,Bfield,linewidth,sys,ph)
```

**DESCRIPTION:**

This function simulates a pulse-acquire experiment with an ideal (instantaneous) excitation pulse and an assumed lorentzian lineshape. The function calls the function 'sim\_Hamiltonian' which produces the free evolution Hamiltonian for the specified number of spins, J and shifts. This function enables an excitation pulse with an arbitrary phase.

**INPUTS:**

```
n          = number of points in fid/spectrum
sw         = desired spectral width in [Hz]
Bfield     = main magnetic field strength in [T]
linewidth  = linewidth in [Hz]
sys        = spin system definition structure
ph         = excitation pulse phase (in degrees)
```

### 2.15. **sim\_onepulse\_shaped.m**

**USAGE:**

```
out = sim_onepulse_shaped(n,sw,Bfield,linewidth,sys,RF,tp,PhCyc,dwdx,G)
```

**DESCRIPTION:**

This function simulates the effect of a frequency selective or slice selective excitation, followed immediately by the acquisition window. This is mainly an exercise to see if I can get slice selective excitation working.

Note that when simulating a frequency selective pulse, it is okay to specify only 8 arguments (no gradient needs to be specified). If the 9th argument, G, is specified and is non-zero, then a slice selective pulse is assumed.

**INPUTS:**

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 sys = spin system definition structure  
 RF = radiofrequency pulse array [N x 3]. Phase, Amplitude, Duration.  
 tp = RF pulse duration in [ms]  
 phCyc = Phase of excitation rf pulse in [degrees].  
 dwdx = if simulating a frequency selective pulse, this argument should be the frequency offset [Hz]. If simulating a slice selective pulse, this argument should be the position offset [cm].  
 G = gradient strength for slice-selective pulse [G/cm];

## 2.16. **sim\_press.m**

### USAGE:

```
out = sim_press(n,sw,Bfield,linewidth,sys,taul,tau2)
```

### DESCRIPTION:

This function simulates an ideal PRESS experiment with first echo time "taul" and a second echo time of "tau2". The function calls the function 'sim\_Hamiltonian.m' which produces the free evolution Hamiltonian and rotation Hamiltonians for the specified spin system.

### INPUTS:

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 sys = spin system definition structure  
 taul = Echo time in [s] of first press Spin Echo  
 tau2 = Echo time in [s] of second press Spin Echo

## 2.17. **sim\_press\_shaped.m**

### USAGE:

```
out =sim_press_shaped(n,sw,Bfield,linewidth,sys,taul,tau2,RF,tp,
                     dx,dy,Gx,Gy,phCyc1,phCyc2))
```

### DESCRIPTION:

This function simulates the PRESS experiment. The excitation is simulated as an instantaneous rotation, and the refocusing pulse is simulated as a shaped rotation.

This code enables the choice of the phase of the refocusing pulses. This enables phase cycling of the refocusing pulses by repeating simulations with different editing pulse phases, which is necessary to remove phase artefacts from the editing pulses. A four-step phase cycling scheme is typically sufficient, where both refocusing pulses are phase cycled by 0 and 90 degrees, and the phase are combined in the following way:

$$\text{signal} = ([0 \ 90] - [0 \ 0]) + ([90 \ 0] - [90 \ 90]);$$

where, in [X Y], X is the phase of the first refocusing pulse and Y is the phase of the second refocusing pulse

Finally, this code simulates the spectrum at a given point in space (x,y), given the values of the slice selection gradients (Gx, and Gy). The pulse waveform is



assumed to be the same for both refocusing pulses. In order to fully simulate the MEGA-PRESS experiment, you have to run this simulation many times at various points in space (x,y), and then add together the resulting spectra.

#### INPUTS:

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 J = matrix of coupling constants in [Hz] for spin system  
 shifts = vector of chemical shifts in [ppm] for spin system  
 tau1 = echo time 1 in [ms].  
 tau2 = echo time 2 in [ms].  
 RF = radiofrequency pulse array [N x 3]. Phase, Amplitude, Duration.  
 tp = RF pulse duration in [ms]  
 peakB1 = peak B1 amplitude in [kHz]  
 dx = position offset in x-direction (corresponding to first refocusing pulse) [cm]  
 dy = position offset in y-direction (corresponding to second refocusing pulse) [cm]  
 Gx = gradient strength for first selective refocusing pulse [G/cm]  
 Gy = gradient strength for second selective refocusing pulse [G/cm]  
 phCycl = initial phase of the first refocusing pulse in [degrees];  
 phCycl2 = initial phase of the second refocusing pulse in [degrees];

### 2.18. **sim\_readout.m**

#### USAGE:

d\_out = sim\_readout(d\_in,H,n,sw,linewidth,rcvPhase,shape)

#### DESCRIPTION:

This function simulates an ADC readout of the transverse magnetization during the free evolution of the spin system under the effects of chemical shift and scalar coupling.

#### INPUTS:

d\_in = input density matrix structure.  
 H = Hamiltonian operator structure.  
 n = number of readout points  
 sw = spectral width [Hz]  
 linewidth = full width at half maximum of spectral peaks [Hz]  
 rcvPhase = receiver phase [degrees]. Optional. Default = 0 (corresponds to x'-axis readout);  
 shape = line broadening function. Optional,  
           'L' = lorentzian (default)  
           'G' = gaussian

### 2.19. **sim\_rotate.m**

#### USAGE:

d\_out = sim\_rotate(d\_in,H,angle,axis)

#### DESCRIPTION:

This function simulates the effect of an ideal (instantaneous) rotation on the density matrix. Used in simulation tools.

**INPUTS:**

**d\_in** = input density matrix structure.  
**H** = Hamiltonian operator structure.  
**angle** = RF pulse flip angle (degrees). If this value is a scalar, then the same flip angle will be applied to all spins in the system. To apply a different flip angle to the different spins in the system, the angle variable can be a vector of flip angles with length equal to the H.nspins.  
**axis** = Axis of rotation ('x', 'y' or 'z'); (A z-rotation technically doesn't correspond to an rf pulse rotation, but it is included here anyway).

**2.20. sim\_rotate\_arbPh.m****USAGE:**

**d\_out** = sim\_rotate\_arbPh(**d\_in**,**H**,**angle**,**ph**)

**DESCRIPTION:**

This function simulates the effect of an ideal (instantaneous) rotation on the density matrix. Used in simulation tools. The phase of the rf pulse can be arbitrarily chosen. To achieve an arbitrary phase, this code executes a rotation about z by an angle of -phi (the rf pulse phase), then executes a rotation about x by an angle of "angle" (the flip angle of the pulse), and then executes a rotation back about z by an angle of phi.

**INPUTS:**

**d\_in** = input density matrix structure.  
**H** = Hamiltonian operator structure.  
**angle** = RF pulse flip angle (degrees). If this value is a scalar, then the same flip angle will be applied to all spins in the system. To apply a different flip angle to the different spins in the system, the angle variable can be a vector of flip angles with length equal to the H.nspins.  
**ph** = Phase of rotation (in degrees; ie. 0='x', 90='y');

**2.21. sim\_shapedRF.m****USAGE:**

**d\_out** = sim\_shapedRF(**d\_in**,**H**,**RFstruct**,**flipAngle**,**phase**,**dfdx**,**grad**)

**DESCRIPTION:**

This function simulates the effect of a shaped rf pulse on the density matrix. The temporal shape of the refocusing pulses is modeled as a series of N instantaneous rotations about the effective RF field, where N is the number of time points in the RF waveform. The instantaneous effective RF field can be an arbitrary vector, and can be represented in polar coordinates as B(Beff,alpha,zeta), where Beff is the magnitude field, alpha is the polar angle (the angle between the transverse plane and the effective B field), and zeta is the azimuthal angle, which is given by the phase of the RF). Rotation about the effective B-field is achieved by a composite rotation: Rotate about Y by -alpha, rotate about Z by -zeta, then rotate about X by  $2\pi\gamma\text{Beff}dt$ , then rotate back about Z by zeta and back about Y by alpha.

**INPUTS:**

**d\_in** = input density matrix structure.  
**H** = Hamiltonian operator structure.  
**RF** = Radiofrequency pulse. This can be the filename of a Siemens .pta file, or an RF pulse definition structure (obtained using rf\_init.m)

Tp = Pulse duration in [ms];  
 flipAngle = RF pulse flip angle [degrees].  
 phase = Phase of RF pulse [degrees]. Optional. Default = 0 (x'-axis. 90 degrees corresponds to +y' axis)  
 dfdx = if simulating a frequency selective pulse, this argument should be the frequency offset [Hz] (Optional. Default = 0 Hz). If simulating a slice selective pulse, this argument should be the position offset [cm].  
 grad = Gradient strength [G/cm]. Optional (for slice selective pulses only).

## 2.22. **sim\_spinecho.m**

### USAGE:

out = sim\_spinecho(n,sw,Bfield,linewidth,sys,tau)

### DESCRIPTION:

This function simulates a spin-echo experiment with instantaneous RF pulses.

### INPUTS:

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 sys = spin system definition structure  
 tau = echo time in [s]

## 2.23. **sim\_spinecho\_shaped.m**

### USAGE:

sim\_spinecho\_shaped(n,sw,Bfield,linewidth,sys,TE,RF,Tp,grad,pos,ph)

### DESCRIPTION:

This function simulates a localized spin-echo sequence with a shaped refocusing pulse. It enables choice of the echo-time as well as the choice of the phase refocusing pulse. This allows phase cycling of the refocusing pulses by repeating simulations with different pulse phases, which is necessary to remove unwanted coherences from outside the volume of interest. For the refocusing pulse, a two step phase cycling scheme is typically sufficient, where the refocusing pulse is phase cycled by 0 and 90 degrees the phase are combined by subtraction.

### INPUTS:

n = number of points in fid/spectrum  
 sw = desired spectral width in [Hz]  
 Bfield = main magnetic field strength in [T]  
 linewidth = linewidth in [Hz]  
 sys = Metabolite spin system definition structure;  
 TE = Echo time in [ms]  
 RF = Refocusing pulse structure (load using io\_loadRFwaveform);  
 Tp = duration of refocusing pulse in [ms]  
 grad = gradient strength for the selective refocusing pulse [G/cm]  
 pos = position offset in the direction corresponding to the refocusing pulse [cm]  
 ph = the phase of the refocusing pulse in [degrees];

## 2.24. **sim\_spoil.m**

### USAGE:

```
d_out = sim_spoil(d_in,H,angle)
```

### DESCRIPTION:

This function simulates the effect of a rotation about the z-axis.

### INPUTS:

```
d_in      = input density matrix structure.  
H         = Hamiltonian operator structure.  
angle     = Spoil angle (degrees).
```

## 2.25. **sim\_steam.m**

### USAGE:

```
out = sim_steam(n,sw,Bfield,linewidth,sys,te,tm)
```

### DESCRIPTION:

Simulate the STEAM sequence using ideal (instantaneous) RF pulses. To remove unwanted coherences, a 4 step phase cycle is automatically performed, with the first and third rf pulses being cycled by 0, 90 180, and 270 degrees. THIS CODE IS NOT TESTED. RESULTS MAY NOT BE ACCURATE!!

### INPUTS:

```
n          = number of points in fid/spectrum  
sw         = desired spectral width in [Hz]  
Bfield     = main magnetic field strength in [T]  
linewidth  = linewidth in [Hz]  
sys        = spin system definition structure  
te         = echo time in [s]  
tm         = mixing time in [s]
```

## 2.26. **sim\_steam\_gradSim.m**

### USAGE:

Script can be run by pressing "run".

### DESCRIPTION:

This function runs the `sim_steam_spoil` function multiple times with different spoiler gradient intensities. The result is a spoiled STEAM sequence.

### INPUTS:

Initialize the following variables and then click "run":

```
spinsys    = Spin system.  
TE         = Echo time [s].  
TM         = Mixing time [s].  
N          = Number of 'phase cycles'
```

### 3. RF Pulse Tools

#### 3.1. rf\_blochSim.m

USAGE:

```
[mv,sc]=rf_blochSim(RF,tp,fspan,f0,peakB1);
```

DESCRIPTION:

Perform a bloch simulation of an RF pulse. This code simply runs Martyn Klassen's excellent bloch equation simulator. For more information, see help file for bes.m. (~FID-A/rfPulseTools/mklassenTools/bes.m).

INPUTS:

```
RF          = RF pulse definition structure
tp          = pulse duration in [ms]
fspan       = Frequency span in [kHz] (optional)
f0          = Centre of frequency span [kHz] (optional)
peakB1      = Peak B1 amplitude in [kHz] (optional)
```

#### 3.2. rf\_dualBand.m

USAGE:

```
[rf,AMPINT]=rf_dualBand(tp,df,n,bw,ph,shft)
```

DESCRIPTION:

Creates an n-point dual banded gaussian inversion RF pulse with duration tp(ms). The first band will be at f=0Hz and the second band will be at df Hz. Bw is the bandwidth of the two selection bands in Hz.

INPUTS:

```
tp          = pulse duration in ms.
df          = frequency of 2nd gaussian band [Hz].
n           = number of points in rf waveform.
bw          = bandwidth of both selection bands [Hz].
ph          = phase of the second gaussian.
shft        = frequency shift applied to both bands.
```

#### 3.3. rf\_freqshift.m

USAGE:

```
RF_shift=rf_freqshift(RF,Tp,f);
```

DESCRIPTION:

Apply a frequency shift to an RF pulse.

INPUTS:

```
RF          = RF pulse definition structure.
Tp          = duration of the rf pulse in [ms].
F           = amount that you would like to frequency shift the rf pulse in [Hz].
```

### 3.4. rf\_gauss.m

USAGE:

```
[rf,AMPINT]=rf_gauss(tp,df,n,bw);
```

DESCRIPTION:

Create an n point gaussian rf waveform. Waveform can be converted in to siemens pta file using rf\_writepta.m or into varian/agilent rf file using io\_writeRF.m.

INPUTS:

tp           = duration of rf pulse in ms.  
df           = frequency of gaussian pulse in Hz. 0 frequency will correspond to the reference frequency of the rf transmitter.  
n            = number of points in the rf waveform.  
bw           = FWHM of the gaussian inversion profile in the frequency domain (Hz).

### 3.5. rf\_hs.m

USAGE:

```
[rf]=rf_hs(outfile,N,n,tbw,Tp,trunc,thk)
```

DESCRIPTION:

This function creates any desired HS pulse. N is the number of steps, n is the order of the HS pulse, tbw is the time bandwidth product of the pulse, Tp is the duration of the pulse and thk is the desired thickness of the pulse.

INPUTS:

outfile       = name of output rf file.  
N            = Number of points in RF waveform.  
n            = order of the HS pulse.  
tbw           = Time bandwidth product.  
Tp           = Duration of the RF pulse (ms).  
trunc        = Truncation of the amplitude modulation function.  
thk           = thickness of the slice selective pulse (optional).

### 3.6. rf\_resample.m

USAGE:

```
RF_out=rf_resample(RF_in,N);
```

DESCRIPTION:

Resample the input RF pulse into a new waveform with N discrete points.

INPUTS:

RF           = RF pulse definition structure  
N            = Number of points in new RF waveform

### 3.7. rf\_sinc.m

USAGE:

```
[rf,AMPINT]=rf_sinc(lobes,n,df);
```

DESCRIPTION:

Create an n point sinc rf waveform. Waveform can be converted in to siemens pta file using rf\_writepta.m or into varian/agilent rf file using io\_writeRF.m.

**INPUTS:**

lobes        = Number of lobes in the sinc pulse.  
n            = number of points in the rf waveform.  
type        = Type of pulse:  
             Refocusing = 'ref'  
             Inversion = 'inv'  
             Excitation = 'exc'

## **4. Input-Output Tools**

### **4.1. io\_loadRFwaveform.m**

**USAGE:**

```
[RF_struct]=io_loadRFwaveform(filename,type,f0);
```

**DESCRIPTION:**

Initialize an RF pulse structure to contain an RF Pulse waveform as well as its accompanying header information. This function finds the time-bandwidth product (tbw) and the time-wl product (twl) of the pulse, and stores this information in the header fields of the output RF structure.

**INPUTS:**

filename    = filename of RF pulse waveform text file. Can be in Siemens format (.pta) or Varian/Agilent format (.RF).  
type        = Excitation ('exc'), Refocusing ('ref') or Inversion ('inv')  
f0          = centre frequency of the rf pulse [Hz]. Optional. Default=0.

### **4.2. io\_loadjmrui.m**

**USAGE:**

```
out=io_loadjmrui(filename);
```

**DESCRIPTION:**

Load a jMRUI text file into matlab structure format.

**INPUTS:**

filename    = filename of the jMRUI txt file.

### **4.3. io\_loadlcmdetail.m**

**USAGE:**

```
[metabs,corrMatrix]=io_loadlcmdetail(filename);
```

**DESCRIPTION:**

This function loads in the "detailed output" of LCModel and returns the matrix of metabolite correlation coefficients.

**INPUTS:**

filename    = Filename of the lcmodel detailed output file.

#### 4.4. io\_loadspec\_GE.m

**USAGE:**

```
[out,out_w]=io_loadspec_GE(filename,sw,Larmor,subspects,te,tr);
```

**DESCRIPTION:**

Reads in GE P file (.dat file) using code adapted from GEREad.m, provided as part of the Gannet software package by Richard Edden (gabamrs.blogspot.com).

op\_loadspec\_GE outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox. NOTE: Since the Gannet code is geared towards edited GABA MRS data, this code may not be general enough to handle all types of MRS data. Suggestions are most welcome.

**INPUTS:**

```
filename    = filename of GE P file to be loaded.  
sw          = spectral width (Hz)  
Larmor      = Larmor frequency (Hz/ppm, ie. 127 for 3T)  
subspects   = number of subspectra in the data (from spectral editing, ISIS, etc.)  
te          = Echo time (ms). Optional. Default is [].  
tr          = Repetition time (ms). Optional. Default is [].
```

#### 4.5. io\_loadspec\_IMA.m

**USAGE:**

```
out=io_loadspec_IMA(filename,Bo,spectralwidth,te,tr);
```

**DESCRIPTION:**

Loads a siemens .IMA file into matlab structure format.

**INPUTS:**

```
filename      = Filename of Siemens .IMA file to load.  
Bo            = Field strength (Tesla).  
spectralwidth = spectral width of the input spectrum (Hz).  
te            = Echo time (ms). Optional. Default is [].  
tr            = Repetition time (ms). Optional. Default is [].
```

#### 4.6. io\_loadspec\_irBruk.m

**USAGE:**

```
out=io_loadspec_irBruk(filename);
```

**DESCRIPTION:**

Reads in Bruker MRS data (1i and 1r files). Generally with this data, the averages and coil channels have already been combined.

op\_loadspec\_brukIR outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox.

**INPUTS:**



inDir = Path to the scan directory that contains the 'pdata' folder.

#### 4.7. io\_loadspec\_data.m

**USAGE:**

```
[out,out_w]=io_loadspec_data(filename,sw,Larmor,subspecs,te,tr);
```

**DESCRIPTION:**

Reads in Philips MRS data (.data and .list files) using code adapted from PhilipsRead\_data.m, provided as part of the Gannet software package by Richard Edden (gabamrs.blogspot.com).

op\_loadspec\_data outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox. NOTE: Since the Gannet code is geared towards edited GABA MRS data, this code may not be general enough to handle all types of MRS data. Suggestions are most welcome.

**INPUTS:**

filename = filename of Philips .data file to be loaded.  
sw = spectral width (Hz)  
Larmor = Larmor frequency (Hz/ppm, ie. 127 for 3T)  
subspecs = number of subspectra in the data (from spectral editing, ISIS, etc.)  
te = Echo time (ms). Optional. Default is [].  
tr = Repetition time (ms). Optional. Default is [].

#### 4.8. io\_loadspec\_sdat.m

**USAGE:**

```
out=io_loadspec_sdat(filename,subspecs);
```

**DESCRIPTION:**

Reads in Philips MRS data (.spar and .sdat files) using code adapted from PhilipsRead.m, provided as part of the Gannet software package by Richard Edden (gabamrs.blogspot.com).

op\_loadspec\_sdat outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox. NOTE: Since the Gannet code is geared towards edited GABA MRS data, this code may not be general enough to handle all types of MRS data. Suggestions are most welcome. ALSO: This code is not currently smart enough to parse out all of the relevant information from the header file, such as the number of subspectra. So for now, these details must be passed to the function as input arguments. Help implementing these improvements are most welcome!!

**INPUTS:**

filename = filename of GE P file to be loaded.  
subspecs = number of subspectra in the data (from spectral editing, ISIS, etc.)

#### **4.9. io\_loadspec\_twix.m**

**USAGE:**

```
out=io_loadspec_twix(filename);
```

**DESCRIPTION:**

Reads in siemens twix raw data (.dat file) using the mapVBVD.m and twix\_map\_obj.m functions from Philipp Ehses (philipp.ehses@tuebingen.mpg.de).

op\_loadspec\_twix outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox.

**INPUTS:**

filename = filename of Siemens twix data to load.

#### **4.10. io\_loadspec\_varian.m**

**USAGE:**

```
out=io_loadspec_varian(filename);
```

**DESCRIPTION:**

Reads in varian .fid data using the readfid.m and readprocpar.m functions from Martyn Klassen (mklassen@robarts.ca).

io\_loadspec\_varian outputs the data in structure format, with fields corresponding to time scale, fids, frequency scale, spectra, and header fields containing information about the acquisition. The resulting matlab structure can be operated on by the other functions in this MRS toolbox.

**INPUTS:**

filename = filename of Varian .fid data to load.

#### **4.11. io\_readRF.m**

**USAGE:**

```
[rf,info]=io_readRF(filename)
```

**DESCRIPTION:**

Read a Varian/Agilent .RF file into matlab. The resulting RF matrix will have 3 columns specifying magnitude, phase, and duration. This function simply calls Martyn Klassen's readrfvnmr.m function.

**INPUTS:**

filename = filename of the .RF file to read in.

#### **4.12. io\_readjmrui.m**

**USAGE:**

```
out=io_readjmrui(filename);
```

**DESCRIPTION:**

Reads jMRUI .txt format into the FID-A data structure format in MATLAB.

INPUTS:  
filename = filename of jMRUI .txt file.

#### 4.13. io\_readlcmcoord.m

USAGE:  
out = io\_readlcmcoord(filename,metab)

DESCRIPTION:  
Reads a LCModel .coord file and extracts the desired part.

INPUTS:  
filename = filename of the LCModel .coord file.  
part = Which metabolite fit to extract from the .coord file - The abbreviated metabolite name should be given (ie. 'Cr','PCr','Glu','GABA',etc.)

#### 4.14. io\_readlcmcoord\_getBackground.m

USAGE:  
out = io\_readlcmcoord\_getBackground(filename,part)

DESCRIPTION:  
Reads a LCModel .coord file and extracts the desired part.

INPUTS:  
filename = filename of the LCModel .coord file.  
part = Which part of the .coord file to extract - 'bg' extracts the LCModel baseline signal, 'sp' extracts the spectrum, and 'fit' extracts the fit.

#### 4.15. io\_readlcmraw.m

USAGE:  
out=io\_readlcmraw(filename,type);

DESCRIPTION:  
Reads LCModel raw data format into the FID-A data structure format in MATLAB.

INPUTS:  
filename = filename of LCModel raw file.  
type = type of LCModel raw file:  
    'rda' - .raw file generated from Siemens RDA file  
    'dat' - .raw file generated by FID-A using io\_writelcm.  
    'sim' - .raw file generated from FID-A simulated data.  
    'raw' - not sure about this one.

#### 4.16. io\_readlcmraw\_basis.m

USAGE:  
out=io\_readlcmraw\_basis(filename);

**DESCRIPTION:**

Reads entire LCModel .basis file into multiple FID-A data structures in MATLAB.

**INPUTS:**

filename = filename of LCModel .basis file.

#### **4.17. io\_readlcmraw\_dotraw.m**

**USAGE:**

out=io\_readlcmraw\_dotraw(filename);

**DESCRIPTION:**

Reads LCModel .RAW model spectrum file into the FID-A data structure format in MATLAB.

**INPUTS:**

filename = filename of LCModel raw file.

#### **4.18. io\_readlcmtab.m**

**USAGE:**

out = io\_readlcmtab(filename)

**DESCRIPTION:**

Reads a LCModel .table output file and stores the metabolite concentrations into a matlab structure array.

**INPUTS:**

filename = filename of the LCModel .table file.

#### **4.19. io\_readpta.m**

**USAGE:**

[rf,info]=io\_readpta(filename)

**DESCRIPTION:**

Read a Siemens .pta file into matlab. The resulting RF matrix will have 2 columns specifying magnitude and phase.

**INPUTS:**

filename = filename of the .pta file to read in.

#### **4.20. io\_writeRF.m**

**USAGE:**

RF=io\_writeRF(rf,outfile);

**DESCRIPTION:**

Write a matlab RF pulse structure (containing 3 x N waveform array field with rf.waveform(1,:)= phase, rf.waveform(2,:)=amplitude, and

rf.waveform(3,:)=timestep), to a varian/agilent format .RF file.

**INPUTS:**

rf           = matlab RF pulse.  
outfile      = name of the output .RF file to be written.

#### **4.21.       io\_readRFBruk.m**

**USAGE:**

rf=io\_readRFBruk(filename);

**DESCRIPTION:**

Read a Bruker RF pulse file into matlab. The resulting RF matrix will have 2 columns specifying magnitude and phase.

**INPUTS:**

filename     = filename of the .pta file to read in.

#### **4.22.       io\_writejmrui.m**

**USAGE:**

RF=io\_writejmrui(in,outfile);

**DESCRIPTION:**

Takes MRS data in matlab structure format and writes it to a text file that can be read by jMRUI.

**INPUTS:**

in           = input data in matlab structure format.  
outfile      = Desired filename of output text file.

#### **4.23.       io\_writelcm.m**

**USAGE:**

RF=io\_writelcm(in,outfile,te);

**DESCRIPTION:**

Takes MRS data in matlab structure format and writes it to a text file that can be read by LCModel.

**INPUTS:**

in           = input data in matlab structure format.  
outfile      = Desired filename of output text file.  
te           = Echo time of acquisition (in ms).

#### **4.24.       io\_writelcmraw.m**

**USAGE:**

RF=io\_writelcmraw(data\_struct,outfile,metab);

**DESCRIPTION:**

Take a simulated metabolite basis spectrum in matlab structure format, and output it into LCModel RAW format to be used in an LCModel basis spectrum.

**INPUTS:**

data\_struct = simulated metabolite basis spectrum in matlab structure format.  
 outfile = name of the output .RAW file.  
 metab = Abbreviated name of the metabolite (ie. 'Cr', 'Glu', etc.)

**4.25. io\_writepta.m****USAGE:**

RF=io\_writepta(rf,outfile);

**DESCRIPTION:**

Write a matlab RF pulse structure (containing N x 3 waveform array field with rf.waveform(:,1)= phase, rf.waveform(:,2)=amplitude, and rf.waveform(:,3)=timestep), to a siemens format .pta file.

**INPUTS:**

rf = matlab RF pulse.  
 outfile = name of the output .pta file to be written.

**5. Processing Tools****5.1. addphase.m****USAGE:**

PhasedSpecs=addphase(specs,AddedPhase);

**DESCRIPTION:**

Add equal amount of complex phase to each point of a vector. This function operates on a vector (fid or spectrum), not on a FID-A data structure. For a phase shifting function that operates on a FID-A data structure, see 'op\_addphase.m'.

**INPUTS:**

specs = Input vector.  
 AddedPhase = Amount of phase (degrees) to add.

**5.2. addphase1.m****USAGE:**

PhasedSpecs=addphase1(specs,ppm,timeShift,ppm0,B0);

**DESCRIPTION:**

Add first order phase to a spectrum (added phase is linearly dependent on frequency). This function operates on a vector (fid or spectrum), not on a FID-A data structure. For a phase shifting function that operates on a FID-A data structure, see 'op\_addphase.m'.

**INPUTS:**

specs = input vector

ppm = frequency scale (ppm) corresponding to the specs vector  
timeShift = This defines the amount of 1st order phase shift by specifying the equivalent horizontal shift (in seconds) in the time domain.  
ppm0 = The frequency "origin" (ppm) of the 1st order phase shift. (this frequency will undergo 0 phase shift).  
B0 = The main magnetic field strength (needed since ppm depends on B0)

### 5.3. op\_ISIScombine.m

**USAGE:**

```
out=op_ISIScombine(in,addInd);
```

**DESCRIPTION:**

Combine dimensions corresponding to ISIS on/off acquisitions to produce fully localized MRS volumes. Mostly used for MEGA-SPECIAL data to combine the ISIS subspectra but not the Edit-on/edit-off subspectra.

**INPUTS:**

in = input data in matlab structure format.  
addInd = (optional) If add==1, then row indices [1 2] and [3 4] will be added. Otherwise, row indices [1 2] and [3 4] will be subtracted.

### 5.4. op\_addNoise.m

**USAGE:**

```
[out,noise]=op_addNoise(in,sdnoise,noise);
```

**DESCRIPTION:**

Add noise to a spectrum (useful for generating simulated data). Normally distributed random noise is added to both the real and imaginary parts of the data. Real and imaginary noise parts are uncorrelated.

**INPUTS:**

in = Input data in matlab structure format.  
sdnoise = Standard deviation of the random noise to be added in the time domain.  
noise = (optional) Specific noise kernel to be added (if specified, sdnoise variable is ignored).

### 5.5. op\_addScans.m

**USAGE:**

```
out=op_addScans(in1,in2,subtract);
```

**DESCRIPTION:**

Add or subtract two scans together.

**INPUTS:**

in1 = First spectrum to add (in matlab structure format)  
in2 = Second spectrum to add (also in matlab structure format).  
subtract = (optional). Add or subtract? (0 = add, 1=subtract). Default=0;

## 5.6. op\_addphase.m

### USAGE:

```
out=op_addphase(in,ph0,ph1,ppm0,suppressPlot);
```

### DESCRIPTION:

add zero and first order phase to a spectrum.

### INPUTS:

```
in          = input spectrum in matlab structure format
ph0         = zero order phase to add (degrees)
ph1         = 1st order phase to add (in seconds);
ppm0        = (optional) frequency reference point. Default = 4.65;
suppressPlot = (optional) Boolean to suppress plots. Default = 0;
```

## 5.7. op\_addphaseSubspec.m

### USAGE:

```
out=op_addphaseSubspec(in,ph);
```

### DESCRIPTION:

Add zero order phase to one of the subspectra in a dataset. For example, the edit-on spectrum of a mega-press acquisition.

### INPUTS:

```
in  = Input spectrum in matlab structure format.
ph  = Phase (in degrees) to add to the second subspectrum.
```

## 5.8. op\_addrcvrs.m

### USAGE:

```
[out,fids_presum,specs_presum,ph,sig]=op_addrcvrs(in,point,mode,coilcombos);
```

### DESCRIPTION:

Perform weighted coil recombination for MRS data acquired with a receiver coil array.

### INPUTS:

```
in          = input spectrum in matlab structure format.
point       = point of fid to use for phase estimation.
mode        = '-w' performs amplitude weighting of channels based on the maximum
              signal of each coil channel.
              '-h' performs amplitude weighting of channels based on the maximum
              signal of each coil channel divided by the square of the noise in
              each coil channel (as described by Hall et al. Neuroimage 2014).
coilcombos  = (optional) The predetermined coil phases and amplitudes as
              generated by the op_getcoilcombos.m function. If this argument is
              provided, the 'point', and 'mode', arguments will be ignored.
```

## 5.9. op\_alignAllScans.m

### USAGE:

```
[out,ph,frq]=op_alignAllScans(in,tmax,ref,mode);
```



**DESCRIPTION:**

Use spectral registration to align many separate scans.

**INPUTS:**

```

in          = cell array of inputs (spectra all to be registered)
tmax        = Maximum time (s) in time domain to use for registration.
ref         = Align to what? (optional)
              'f' - Align to the first input? (default)
              'a' - Align to average of inputs.
mode        = (optional)'f' - Frequency align only
              'p' - Phase align only
              'fp or pf' - Frequency and phase align (default)

```

**5.10. op\_alignAllScans\_fd.m****USAGE:**

```
[out,ph,frq]=op_alignAllScans_fd(in,fmin,fmax,tmax,ref,mode);
```

**DESCRIPTION:**

Use spectral registration to align many separate scans. In this version only a limited frequency range is used for fitting;

**INPUTS:**

```

in          = cell array of inputs (spectra all to be registered)
fmin        = Minimum frequency for spectral alignment (ppm).
fmax        = Maximum frequency for spectral alignment (ppm).
tmax        = Maximum time (s) in time domain to use for registration.
ref         = Align to what? (optional)
              'f' - Align to the first input? (default)
              'a' - Align to average of inputs.
mode        = (optional)'f' - Frequency align only
              'p' - Phase align only
              'fp or pf' - Frequency and phase align (default)

```

**5.11. op\_alignAverages.m****USAGE:**

```
[out,fs,phs]=op_alignAverages(in,tmax,avg,initPars);
```

**DESCRIPTION:**

Perform spectral registration in the time domain to correct frequency and phase drifts. As described in Near et al. Frequency and phase drift correction of magnetic resonance spectroscopy data by spectral registration in the time domain. Magn Reson Med. 2014 Jan 16. doi: 10.1002/mrm.25094. [Epub ahead of print]

**INPUTS:**

```

in          = Input data structure.
tmax        = Maximum time (s) in time domain to use for alignment.
avg         = Align averages to the average of the averages? (y or n)
initPars    = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].
              Default=[0,0];

```

### 5.12. **op\_alignAverages\_fd.m**

**USAGE:**

```
[out,fs,phs]=op_alignAverages_fd(in,minppm,maxppm,tmax,avg,initPars);
```

**DESCRIPTION:**

Perform time-domain spectral registration using a limited range of frequencies to correct frequency and phase drifts. As described in Near et al. Frequency and phase drift correction of magnetic resonance spectroscopy data by spectral registration in the time domain. Magn Reson Med. 2014 Jan 16. doi: 10.1002/mrm.25094. [Epub ahead of print]

**INPUTS:**

```
in           = Input data structure.
Minppm       = Minimum of frequency range (ppm).
Maxppm       = Maximum of frequency range (ppm).
tmax         = Maximum time (s) in time domain to use for alignment.
avg          = Align averages to the average of the averages? (y or n)
initPars     = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].
               Default=[0,0];
```

### 5.13. **op\_alignISIS.m**

**USAGE:**

```
[out,fs,phs]=op_alignISIS(in,tmax,initPars);
```

**DESCRIPTION:**

Apply spectral registration to align ISIS subspectra prior to subtraction. This is intended to be used prior to averaging, so that the alignment can be performed independently for each average.

**INPUTS:**

```
in           = Input data structure.
tmax         = Maximum time (s) in time domain to use for alignment.
initPars     = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].
               Default=[0,0];
```

### 5.14. **op\_alignMPSubspecs.m**

**USAGE:**

```
[out,fs,phs]=op_alignMPSubspecs(in,initPars);
```

**DESCRIPTION:**

Apply spectral registration to align MEGA-PRESS subspectra prior to subtraction. This function is designed to minimize subtraction artefacts from choline, and residual water. This is intended to be used after averaging.

**INPUTS:**

```
in           = Input data structure.
initPars     = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].
               Default=[0,0];
```

### 5.15. **op\_alignMPSubspecs\_fd.m**

**USAGE:**

```
[out,fs,phs]=op_alignMPSubspecs_fd(in,initPars);
```

**DESCRIPTION:**

Apply spectral registration to align MEGA-PRESS subspectra prior to subtraction. This function is designed to minimize subtraction artefacts from choline, and residual water. This is intended to be used after averaging.

**INPUTS:**

```
in          = Input data structure.  
initPars    = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].  
              Default=[0,0];
```

### 5.16. **op\_alignScans.m**

**USAGE:**

```
[out,ph,frq]=op_alignScans(in,in1,tmax,mode);
```

**DESCRIPTION:**

Use spectral registration to align two separate scans (align in to in1);

**INPUTS:**

```
in          = input (spectrum to be registered)  
in1         = base (spectrum that the input is to be registered to).  
tmax        = Maximum time (s) in time domain to use for registration.  
mode        = (optional)'f' - Frequency align only  
              'p' - Phase align only  
              'fp or pf' - Frequency and phase align (default)
```

### 5.17. **op\_alignScans\_fd.m**

**USAGE:**

```
[out,ph,frq]=op_alignScans_fd(in,in1,fmin,fmax,tmax,mode);
```

**DESCRIPTION:**

Use spectral registration on a limited frequency range to align two separate MRS datasets (align in to in1);

**INPUTS:**

```
in          = input (spectrum to be registered to the base)  
in1         = base (spectrum that the input is to be registered to).  
fmin        = Minimum frequency for spectral alignment (ppm).  
fmax        = Maximum frequency for spectral alignment (ppm).  
tmax        = Maximum time (s) in time domain to use for registration.  
mode        = (optional)'f' - Frequency align only  
              'p' - Phase align only  
              'fp or pf' - Frequency and phase align (default)
```

### 5.18. **op\_alignrcvrs.m**

**USAGE:**

```
[out,ph,sig]=op_alignrcvrs(in,point,mode,coilcombos);
```

**DESCRIPTION:**

Phase align the receiver channels without combining them.

**INPUTS:**

in = input spectrum in matlab structure format.  
point = Index of point in time domain to use for phase refernece.  
Coilcombos = (optional) The predetermined coil phases and amplitudes as generated by the op\_getcoilcombos.m function. If this argument is provided, the 'point', and 'mode', arguments will be ignored.

### 5.19. **op\_ampScale.m**

**USAGE:**

```
out=op_ampScale(in,A);
```

**DESCRIPTION:**

Scale the amplitude of a spectrum by factor A.

**INPUTS:**

in = input data in matlab structure format  
A = Amplitude scaling factor.

### 5.20. **op\_arsos.m**

**USAGE:**

```
out=op_arsos(in);
```

**DESCRIPTION:**

Perform all rank statistic order filter (see Slotboom J et al, Meas Sci Technol. 20 (2009)). This effectively rank-orders the data along the averages dimension. This can be done in either the time domain (default) or the frequency domain.

**INPUTS:**

in = input data in matlab structure format.  
Domain = time domain ('t', (default)) or frequency domain ('f').

### 5.21. **op\_averaging.m**

**USAGE:**

```
out=op_averaging(in);
```

**DESCRIPTION:**

Combine the averages in a scan by adding the averages together and then dividing by the number of averages.

**INPUTS:**

in = input data in matlab structure format.

## 5.22. **op\_combinesubspecs.m**

### USAGE:

```
out=op_combinesubspecs(in,mode);
```

### DESCRIPTION:

Combine the subspectra in an acquisition either by addition or subtraction.

### INPUTS:

```
in      = input data in matlab structure format.  
mode    = -"diff" adds the subspectra together (this is counter intuitive, but  
          the reason is that many "difference editing" sequences use phase  
          cycling of the readout ADC to achieve "subtraction by addition".  
          -"summ" performs a subtraction of the subspectra.
```

## 5.23. **op\_complexConj.m**

### USAGE:

```
out=op_complexConj(in)
```

### DESCRIPTION:

Take the complex conjugate of the data;

### INPUTS:

```
in      = Input data in matlab structure format.
```

## 5.24. **op\_concatAverages.m**

### USAGE:

```
out=op_concatAverages(in1,in2);
```

### DESCRIPTION:

Concatenate two scans along the averages dimension. Two scans with 50 averages each will now look like a single scan with 100 averages.

### INPUTS:

```
in1     = first input in matlab structure format.  
in2     = second input in matlab structure format.
```

## 5.25. **op\_concatSubspecs.m**

### USAGE:

```
out=op_concatSubspecs(in1,in2);
```

### DESCRIPTION:

Concatenate two scans along the subspecs dimension. Two scans with 50 averages each will now look like a single scan with 100 averages.

### INPUTS:

```
in1     = first input in matlab structure format.  
in2     = second input in matlab structure format.
```

## 5.26. **op\_creFit.m**

### USAGE:

```
parsFit=op_creFit(in,ph0,ph1);
```

### DESCRIPTION:

Perform a Lorentzian lineshape fit to the creatine resonance in a brain proton MRS dataset.

### INPUTS:

in = input data in matlab stucture format.  
ph0 = zero order phase to add to the input data.  
ph1 = 1st order phase to add to the input data.

## 5.27. **op\_dccorr.m**

### USAGE:

```
out=op_dccorr(in,mode,var);
```

### DESCRIPTION:

Do a DC Correction on the data. This method is a frequency domain operation.

### INPUTS:

in = input data in matlab structure format.  
Mode = Point('p') or Value('v'). In point mode, the DC offset is calculated automatically at a specific point in the spectrum. In value mode, the user has to provide the value of the desired DC offset.  
var = If mode is 'p', then 'var' is the index of the spectral point that you wish to use to calculate the DC offset. If mode is 'v', then 'var' is the value of the dc offset correction that you wish to employ.

## 5.28. **op\_downsamp.m**

### USAGE:

```
out=op_downsamp(in,dsFactor);
```

DESCRIPTION: Change the time domain sampling rate of a spectrum by a factor of 'dsFactor'. Nearest neighbour interpolation is performed by default.

### INPUTS:

in = input data in matlab structure format.  
dsFactor = factor by which to divide the sampling rate of the fid.

## 5.29. **op\_ecc.m**

### USAGE:

```
[out,outw]=op_ecc(in,inw);
```

### DESCRIPTION:

Perform an eddy current correction by estimating any non-linearity in the phase of the water unsuppressed data in the time domain and applying the appropriate correction to both the water suppressed and water unsuppressed data.

**INPUTS:**

in = water suppressed input data in matlab structure format.  
in2 = water unsuppressed input data in matlab structure format.

**5.30. op\_fddccorr.m**

**USAGE:**

out=op\_fddccorr(in,npts);

**DESCRIPTION:**

Correct and DC offset in the frequency domain. This is equivalent to a vertical shift in the frequency domain. The required vertical shift is calculated by taking the average of the first and last "NPTS" points in the frequency domain. This requires that those points are in the noise floor.

**INPUTS:**

in = input data in matlab structure format.  
npts = number of points at both edges of the frequency domain that will be used for estimation of the DC offset of the spectrum.

**5.31. op\_filter.m**

**USAGE:**

out=op\_filter(in,lb);

**DESCRIPTION:**

Perform line broadening by multiplying the time domain signal by an exponential decay function.

**INPUTS:**

in = input data in matlab structure format.  
lb = line broadening factor in Hz.

**5.32. op\_freqAlignAverages.m**

**USAGE:**

[out,fs]=op\_freqAlignAverages(in,tmax,avg,initPars);

**DESCRIPTION:**

Perform spectral registration in the time domain using only frequency adjustment (no phase adjustment).

**INPUTS:**

in = Input data structure.  
tmax = Maximum time (s) in time domain to use for alignment.  
avg = Align averages to the average of the averages? (y or n)  
initPars = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].  
Default=[0,0];

### 5.33. **op\_freqAlignAverages\_fd.m**

**USAGE:**

```
[out,fs,phs]=op_freqAlignAverages_fd(in,minppm,maxppm,tmax,avg,initPars);
```

**DESCRIPTION:**

Perform time-domain spectral registration using a limited range of frequencies and using only frequency adjustment (no phase adjustment).

**INPUTS:**

in = Input data structure.  
minppm = Minimum of frequency range (ppm).  
maxppm = Maximum of frequency range (ppm).  
tmax = Maximum time (s) in time domain to use for alignment.  
avg = Align averages to the average of the averages? (y or n)  
initPars = (Optional) Initial fit parameters [freq(Hz), phase(degrees)].  
Default=[0,0];

### 5.34. **op\_freqrange.m**

**USAGE:**

```
out=op_freqrange(in,ppmmin,ppmmax);
```

**DESCRIPTION:**

Output only a specified frequency range of the input spectrum.

**INPUTS:**

in = input data in matlab structure format.  
ppmmin = minimum extent of frequency range in ppm.  
ppmmax = maximum extent of frequency range in ppm.

### 5.35. **op\_freqshift.m**

**USAGE:**

```
out=op_freqshift(in,f);
```

**DESCRIPTION:**

Apply a frequency shift to the input spectrum by 'f' Hz.

**INPUTS:**

in = input data in matlab structure format.  
f = frequency shift to apply (in Hz).

### 5.36. **op\_freqshiftSubspec.m**

**USAGE:**

```
out=op_freqshiftSubspec(in,f);
```

**DESCRIPTION:**

Apply a frequency shift to only one of the sub-spectra in a dataset. This is used to minimize subtraction artefacts from MEGA\_PRESS data, for instance.

**INPUTS:**

in = input data in matlab structure format.



f = frequency shift to apply to subspectrum (in Hz).

### 5.37. op\_gaussianPeak.m

**USAGE:**

out=op\_gaussianPeak(n,sw,Bo,lw,ppm0,amp);

**DESCRIPTION:**

Generate a noiseless spectrum containing a single gaussian peak with desired parameters (frequency, amplitude, linewidth, etc.).

**INPUTS:**

n = Number of points in spectrum.  
sw = spectral width of spectrum (Hz).  
Bo = Magnetic field strength (Tesla).  
lw = Linewidth of gaussian peak (Hz).  
ppm0 = Frequency of gaussian peak (ppm).  
amp = Amplitude of gaussian peak.

### 5.38. op\_getLW.m

**USAGE:**

[FWHM]=op\_getLW(in,zpfactor,Refppmmin,Refppmmax);

**DESCRIPTION:**

Estimates the linewidth of a reference peak in the spectrum. Two methods are used to estimate the linewidth: 1. FWHM is measured by simply taking the full width at half max of the reference peak. 2. The FWHM is measured by fitting the reference peak to a lorentzian lineshape and determine the FWHM of the best fit. The output FWHM is given by the average of these two measures.

**INPUTS:**

in = input spectrum in structure format.  
zpfactor = zero-padding factor (used for method 1.)  
Refppmmin = Min of frequency range (ppm) in which to search for reference peak.  
Refppmmax = Max of frequency range to (ppm) in which search for reference peak

### 5.39. op\_getSNR.m

**USAGE:**

[SNR]=op\_getSNR(in,NAAppmmin,NAAppmmax,noiseppmmin,noiseppmmax);

**DESCRIPTION:**

Find the SNR of the NAA peak in a spectrum.

**INPUTS:**

in = input data in matlab structure format  
NAAppmmin = min of frequency range in which to search for NAA peak.  
NAAppmmax = max of frequency range in which to search for NAA peak.  
noiseppmmin = min of frequency range in which to measure noise.  
noiseppmmax = max of frequency range in which to measure noise.

#### 5.40. **op\_getcoilcombos.m**

**USAGE:**

```
coilcombos=op_getcoilcombos(file_or_struct,point);
```

**DESCRIPTION:**

This function finds the relative coil phases and amplitudes. Coil phases are found by determining the phase and amplitude of the pointth point in the time domain.

**INPUTS:**

file\_or\_struct = this function will accept either a string filename or the name of a structure. If the input is a string, the program will read in the data corresponding to that filename. If the input is a structure, it will operate on that structure.

point = The index of the datapoint in the fid that is used for determination of signal intensity and phase.

#### 5.41. **op\_getcoilcombos\_specReg.m**

**USAGE:**

```
coilcombos=op_getcoilcombos_specReg(file_or_struct,tmin,tmax,point);
```

**DESCRIPTION:**

This function finds the relative coil phases and amplitudes. Coil phases are found by fitting in the time domain. (In the original op\_getcoilcombos, the phases were determined simply by observation of the pointth point in the time domain). The "Base" receiver channel is determined as the one with the highest signal (all other coil channels will be registered to that channel).

**INPUTS:**

file\_or\_struct = this function will accept either a string filename or the name of a structure. If the input is a string, the program will read in the data corresponding to that filename. If the input is a structure, it will operate on that structure.

tmin = The earliest timepoint in the fid to be used for spectral registration.

tmax = The latest timepoint in the fid to be used for spectral registration.

point = The index of the datapoint in the fid that is used for determination of signal intensity.

#### 5.42. **op\_leftshift.m**

**USAGE:**

```
out=op_leftshift(in,ls);
```

**DESCRIPTION:**

Remove leading datapoints from the fid to get rid of 1st order phase errors.

**INPUTS:**

in = input data in matlab structure format.

ls = number of points to remove from the beginning of the fid.

### 5.43. **op\_lorentz.m**

**USAGE:**

```
y=op_lorentz(pars,ppm);
```

**DESCRIPTION:**

Generate a parametrized lorentzian peak. This function is fed into fitting tools to enable fitting of peaks using lorentzian lineshapes.

**INPUTS:**

pars = The parameters of the Lorentzian function. This is a five element vector consisting of the following fields:  
[ Amplitude,  
FWHM, (In Hz)  
Centre Freq, (In ppm)  
baseline offset, (in Amplitude units)  
Phase shift]; (in degrees)  
ppm = frequency axis vector (in ppm);

### 5.44. **op\_lorentz\_linbas.m**

**USAGE:**

```
y=op_lorentz_linbas(pars,ppm);
```

**DESCRIPTION:**

Generate a parametrized lorentzian peak with a linearly sloping baseline. This function is fed into fitting tools to enable fitting of peaks using lorentzian lineshapes.

**INPUTS:**

pars = The parameters of the Lorentzian function. This is a six element vector consisting of the following fields:  
[ Amplitude,  
FWHM, (In Hz)  
Centre Freq, (In ppm)  
baseline slope, (in Amplitude units per ppm)  
baseline offset, (in Amplitude units)  
Phase shift]; (in degrees)  
ppm = frequency axis vector (in ppm);

### 5.45. **op\_lorentzianPeak.m**

**USAGE:**

```
out=op_lorentzianPeak(n,sw,Bo,lw,ppm0,amp);
```

**DESCRIPTION:**

Generate a noiseless spectrum containing a single lorentzian peak with desired parameters (frequency, amplitude, linewidth, etc.).

**INPUTS:**

n = Number of points in spectrum.  
sw = spectral width of spectrum (Hz).  
Bo = Magnetic field strength (Tesla).  
lw = Linewidth of gaussian peak (Hz).  
ppm0 = Frequency of gaussian peak (ppm).  
amp = Amplitude of gaussian peak.

#### 5.46. **op\_makeFreqDrift.m**

**USAGE:**

```
[out,fDrift]=op_makeFreqDrift(in,totalDrift,noise);
```

**DESCRIPTION:**

Add frequency drift to a dataset containing multiple averages. This is generally used to generate simulated datasets with phase drift.

**INPUTS:**

in = input data in matlab structure format.  
totalDrift = total amount of frequency drift (in Hz) to add over the whole scan.  
If totalDrift is a scalar, then a constant slope of drift will be added. If totalDrift is a vector with length equal to the number of averages in the input data, then this vector specifies the drift applied to each average.  
noise = the standard deviation of noise to add to the frequency drift function.

#### 5.47. **op\_makePhaseDrift.m**

**USAGE:**

```
[out,phDrift]=op_makePhaseDrift(in,totalDrift,noise);
```

**DESCRIPTION:**

Add phase drift to a dataset containing multiple averages. This is generally used to generate simulated datasets with phase drift.

**INPUTS:**

in = input data in matlab structure format.  
totalDrift = total amount of phase drift (in degrees) to add over the whole scan.  
If totalDrift is a scalar, then a constant slope of drift will be added. If totalDrift is a vector with length equal to the number of averages in the input data, then this vector specifies the drift applied to each average.  
noise = the standard deviation of noise to add to the phase drift function.

#### 5.48. **op\_median.m**

**USAGE:**

```
out=op_median(in);
```

**DESCRIPTION:**

Combine the averages in a scan by calculating the median of all averages, as described in Slotboom J et al, Meas Sci Technol 20 (2009).

**INPUTS:**

in = input data in matlab structure format.

### 5.49. **op\_phaseAlignAverages.m**

**USAGE:**

```
[out,phs]=op_phaseAlignAverages(in,Npts,avg,weighting)
```

**DESCRIPTION:**

Perform time-domain spectral registration using only phase adjustment (no frequency adjustment). This is rarely used.

**INPUTS:**

```
in           = Input data structure.
Npts         = Number of points in time domain to use for alignment.
avg          = Align averages to the average of the averages ('y'), or the first
              average in the series ('n');
weighting    = (Optional) Apply less weight to the later points of the fid?
```

### 5.50. **op\_phaseAlignAverages\_fd.m**

**USAGE:**

```
[out,phs]=op_phaseAlignAverages_fd(in,minppm,maxppm,Npts,avg,weighting)
```

**DESCRIPTION:**

Perform time-domain spectral registration using a limited range of frequencies and using only phase adjustment (no frequency adjustment). This is rarely used.

**INPUTS:**

```
in           = Input data structure.
minppm       = Minimum of frequency range (ppm).
maxppm       = Maximum of frequency range (ppm).
Npts         = Number of points in time domain to use for alignment.
avg          = Align averages to the average of the averages ('y'), or the first
              average in the series ('n');
weighting    = (Optional) Apply less weight to the later points of the fid?
```

### 5.51. **op\_plotfid.m**

**USAGE:**

```
out=op_plotfid(in,tmax,xlab,ylab,tit);
```

**DESCRIPTION:**

Plot the spectrum in the time domain.

**INPUTS:**

```
in           = input data in matlab structure format. This argument can be a MATLAB
              structure (formatted according to the FID-A structure format), or it can
              be a cell array, where the elements of each cell are FID-A structures.
              In the latter case, each spectrum in the structure will be plotted, with
              the option of a vertical offset.
tmax         = upper limit of time scale to plot in seconds (optional. Default =
              max(in.t)).
xlab         = Label for the x-axis (optional. Default = 'Time (sec)');
ylab         = label for the y-axis (optional. Default = 'FID Amplitude (arb units)');
tit          = label for the title of the plot (optional. Default = '');
```

## 5.52. **op\_plotspec.m**

### USAGE:

```
out=op_plotspec(in,ppmmin,ppmmax,xlab,ylab,tit);
```

### DESCRIPTION:

Plot the MR spectrum in the frequency domain.

### INPUTS:

```
in      = input data in matlab structure format. This argument can be a MATLAB
         structure (formatted according to the FID-A structure format), or it can
         be a cell array, where the elements of each cell are FID-A structures.
         In the latter case, each spectrum in the structure will be plotted, with
         the option of a vertical offset.
ppmmin  = lower limit of ppm scale to plot (optional. Default = 0.2 ppm).
ppmmax  = upper limit of ppm scale to plot (optional. Default = 5.2 ppm).
xlab    = Label for the x-axis (optional. Default = 'Frequency (ppm)');
ylab    = label for the y-axis (optional. Default = '');
tit     = label for the title of the plot (optional. Default = '');
```

## 5.53. **op\_ppmref.m**

### USAGE:

```
out=op_ppmref(in,ppmmin,ppmmax,ppmrefval);
```

### DESCRIPTION:

Search for the peak located between ppmmin and ppmmax, and then give that peak a new ppm reference value.

### INPUTS:

```
in      = input data in matlab structure format.
ppmmin  = minimum of ppm search range.
ppmmax  = maximum of ppm search range.
ppmrefval = new reference ppm value.
```

## 5.54. **op\_relyTest.m**

### USAGE:

```
out = op_relyTest(in);
```

### DESCRIPTION:

Perform reliability testing on a dataset with multiple averages, as described in Slotboom J et al, Meas Sci Technol 20 (2009). This involves calculating the skewness (3rd standard moment) and kurtosis (4th standard moment) of any dataset along the "averages" dimension. The output structure contains skewness and kurtosis vectors (as a function of t), as well as the mean of the absolute values skewness and kurtosis for all timepoints where the average FID SNR is greater than 2.

### INPUTS:

```
in      = input data in matlab structure format.
```

### **5.55.        `op_rmNworstaverages.m`**

**USAGE:**

```
[out,metric,badAverages]=op_rmNworstaverages(in,n);
```

**DESCRIPTION:**

Removes motion corrupted averages from a dataset containing multiple averages. The N most badly motion corrupted averages are discarded.

**INPUTS:**

```
in            = input data in matlab structure format
n            = number of bad averages to remove
```

### **5.56.        `op_rmbadaverages.m`**

**USAGE:**

```
[out,metric,badAverages]=op_rmbadaverages(in,nsd,domain);
```

**DESCRIPTION:**

Removes motion corrupted averages from a dataset containing multiple averages. Bad averages are identified by calculating a 'likeness' metric for each average. This is done by subtracting each average from the average of the averages, and then calculating the root mean squared of this difference spectrum. Averages whose likeness metrics are greater than 'nsd' above the mean are discarded.

**INPUTS:**

```
in            = input data in matlab structure format
nsd           = number of standard deviations to use a rejection threshold
domain        = domain in which to perform calculations ('t' or 'f')
```

### **5.57.        `op_rmworstaverage.m`**

**USAGE:**

```
[out,metric,badAverages]=op_rmworstaverage(in,n);
```

**DESCRIPTION:**

Removes motion corrupted averages from a dataset containing multiple averages. The most badly motion corrupted average is discarded.

**INPUTS:**

```
in            = input data in matlab structure format
```

### **5.58.        `op_subtractScans.m`**

**USAGE:**

```
out=op_subtractScans(in1,in2);
```

**DESCRIPTION:**

Subtract input 2 from input 1;

**INPUTS:**

```
in1           = 1st input data in matlab structure format.
in2           = 2nd input data in matlab structure format.
```

### **5.59.        op\_takeaverages.m**

**USAGE:**

```
out=op_takeaverages(in,index);
```

**DESCRIPTION:**

Extract the averages with indices corresponding to the 'index' input array.

**INPUTS:**

in        = input data in matlab structure format.

index    = vector indicating the indices of the averages you would like to extract.

### **5.60.        op\_takesubspec.m**

**USAGE:**

```
out=op_takesubspec(in,index);
```

**DESCRIPTION:**

Extract the subspectra with indices corresponding to the 'index' input array.

**INPUTS:**

in        = input data in matlab structure format.

index    = vector indicating the indices of the subspectra you would like to extract.

### **5.61.        op\_timerange.m**

**USAGE:**

```
out=op_timerange(in,ppmmin,ppmmax);
```

**DESCRIPTION:**

Output only a specified frequency range of the input spectrum.

**INPUTS:**

in        = input data in matlab structure format.

ppmmin    = minimum extent of frequency range in ppm.

ppmmax    = maximum extent of frequency range in ppm.

### **5.62.        op\_unfilter.m**

**USAGE:**

```
out=op_unfilter(in,lb);
```

**DESCRIPTION:**

Multiply the fid by an inverted exponential decay function to undo the effects of filtering.

**INPUTS:**

in        = input data in matlab structure format.

lb        = line narrowing factor (Hz).



### 5.63. `op_zeropad.m`

**USAGE:**

```
out=op_zeropad(in,zpFactor);
```

**DESCRIPTION:**

Apply zeropadding (a.k.a. zero-filling) to MRS data.

**INPUTS:**

```
in          = input data in matlab structure format.
zpFactor    = the factor by which the number of points in the fid will be increased.
              ie. if zpFactor =2, then the number of zeros added to the end of the
              fid will be equal to the number of points in the original spectrum.
```

### 5.64. `op_zerotrim.m`

**USAGE:**

```
out=op_zerotrim(in,numPointsToTrim);
```

**DESCRIPTION:**

Remove zeros (or even non-zero data points) from the end of the fid.

**INPUTS:**

```
in          = input data in matlab structure format.
numPointsToTrim = The number of points to trim from the end of the fid.
```

## 6. Example Run Scripts

The FID-A software package contains a library of example run scripts which are located in the {FID-A\_DIR}/exampleRunScripts directory. These are intended to provide the user with examples of useful “pipelines” for NMR simulation and data processing. In the case of NMR simulation pipelines, the provided example run scripts involve spatially resolved simulations for experiments with shaped localization pulses, or examples of how to generate a simple LCModel basis set. In the case of MRS data processing, the provided example run scripts demonstrate all of the necessary steps for a full processing pipeline, from importing the raw data into matlab, combining RF coil channels, removing motion corrupted averages, spectral registration to remove frequency and phase drift, signal averaging and phase correction, and exporting the final processed data into a format that is readable by one of the leading analysis software packages. Some of these example processing scripts may be

useable in their existing form for routine data processing of MRS data. Others may need to be modified to suit the user's specific needs. All of the example run scripts make use of the various functions in within the FID-A toolboxes. The available example run scripts are described below:

### 6.1. run\_getLWandSNR.m

**USAGE:**

```
[ FWHM,SNR ] = run_getLWandSNR(in);
```

**DESCRIPTION:**

Calculate the linewidth and SNR of a spectrum. This function calculates the Linewidth by measuring the FWHM of the unsuppressed water peak. SNR is calculated by measuring the height of the NAA peak and comparing this to the standard deviation of the noise in a signal-free region of the spectrum. SNR is measured four separate times using four different noise regions and the average of those four measurements is reported.

**INPUTS:**

in = input data in matlab structure **format**

### 6.2. run\_make2DSimPlot.m

**USAGE:**

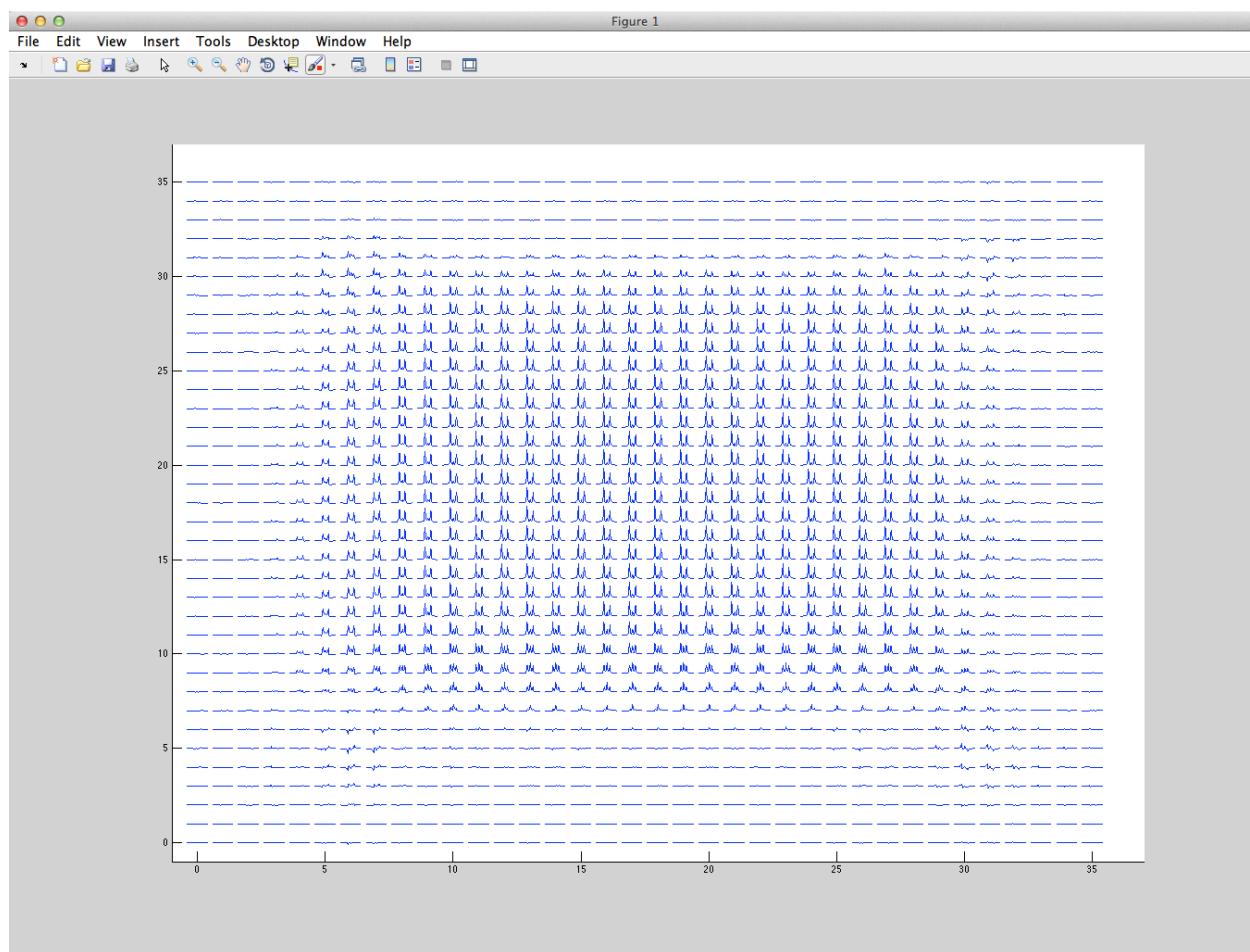
```
run_make2DSimPlot(in,ppmmin,ppmmax,plotDiff)
```

**DESCRIPTION:**

This function takes the output of a spatially resolved simulation, and plots the array of spectra on a single figure. The input should be a cell array where the grid of elements of the cell array are simulated spectra from a corresponding grid of spatial positions in the spatially resolved simulation. Each element of the cell array is also in FID-A data struture format. By including the optional input arguement ppmmin and ppmmax, only a the corresponding range of each spectrum will be plotted. An example of a plot generated by run\_make2DSimPlot is shown below in Figure 1.

**INPUTS:**

in = input cell array of simulated spectra from a spatially resolved simulation  
ppmmin = lower limit of ppm range to plot [ppm]  
ppmmax = upper limit of ppm range to plot [ppm]



**Figure 1:** The output of `run_make2DSimPlot`, which shows simulated, spatially resolved MEGA-PRESS difference spectra of the GABA spin system. This simulation was generated using `run_simMegaPressShaped`, and it incorporates the shaped editing and refocusing pulses. Only the 3ppm GABA resonance is shown.

### 6.3. `run_megapressproc.m`

#### USAGE:

```
[out1_diff,out1_sum,outw]=run_megapressproc(filestring,coilcombos,
                                           avgAlignDomain,aligns);
```

#### DESCRIPTION:

Processing script for Siemens MEGA-PRESS MRS data in .dat format (twix raw data). Includes combination of receiver channels, removal of bad averages, frequency drift correction, manual alignment of edit-on and edit-off spectra, and leftshifting.

#### INPUTS:

<code>filestring</code>	= String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '_w/'];
<code>coilcombos</code>	= (Optional). A structure obtained by running the <code>op_getcoilcombos</code> function. This allows the user to specify the coil phases and amplitudes as an input, rather calculating these from the input data by default.

avgAlignDomain = (Optional) Perform the spectral registration (drift correction) using the full spectrum ('t'), or only a limited frequency range ('f'). Default is 'f'.

alignSS = (Optional)  
 0 - Do not align the edit-on and edit-off subspectra.  
 2 - Perform manual alignment of edit-on and edit-off subspectra.

**OUTPUTS:**

out1\_diff = Fully processed difference spectrum.  
 out1\_sum = Fully processed sum spectrum.  
 outw = Fully processed water unsuppressed spectrum.

## 6.4. run\_megapressproc\_auto.m

**USAGE:**

```
[out1_diff,out1_sum,out1,outw,coilcombos]=run_megapressproc_auto(filestring,coilcombos,avgAlignDomain,alignSS);
```

**DESCRIPTION:**

Fully automated processing script for Siemens MEGA-PRESS MRS data in .dat format (twix raw data). Includes combination of receiver channels, removal of bad averages, frequency drift correction, manual alignment of edit-on and edit-off spectra, and leftshifting. This pipeline requires no user interaction. This function automatically generates an html report to describe the results of each processing step.

**INPUTS:**

Filestring = String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '\_w/'];

coilcombos = (Optional). A structure obtained by running the op\_getcoilcombos function. This allows the user to specify the coil phases and amplitudes as an input, rather than calculating these from the input data by default.

avgAlignDomain = (Optional) Perform the spectral registration (drift correction) using the full spectrum ('t'), or only a limited frequency range ('f'). Default is 'f'.

alignSS = (Optional)  
 0 - Do not align the edit-on and edit-off subspectra (default).  
 2 - Perform manual alignment of edit-on and edit-off subspectra.

**OUTPUTS:**

out1\_diff = Fully processed difference spectrum.  
 out1\_sum = Fully processed sum spectrum.  
 outw = Fully processed water unsuppressed spectrum.

## 6.5. run\_pressproc.m

**USAGE:**

```
[out,out_w,out_noproc,out_w_noproc]=run_pressproc(filestring,aaDomain,tmaxin,iterin);
```

**DESCRIPTION:**

Processing script for Siemens PRESS MRS data in .dat format (twix raw data). Includes combination of receiver channels, removal of bad averages, frequency drift correction, and leftshifting.

**INPUTS:**

filestring: String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '\_w/'];

aaDomain: (Optional) Perform the spectral registration (drift correction) using the full spectrum ('t'), or only a limited frequency range ('f'). Default is 'f'.

tmaxin: (Optional). Duration (in sec.) of the time domain signal used in the spectral registration (drift correction). Default is 0.2 sec.

iterin: (Optional). Maximum number of allowed iterations for the spectral registration to converge. Default is 20.

**OUTPUTS:**

out: Fully processed, water suppressed output spectrum.

out\_w: Fully processed, water unsuppressed output spectrum.

out\_noproc: Water suppressed output spectrum without pre-processing (No bad-averages removal, no frequency drift correction).

out\_w\_noproc: Water unsuppressed output spectrum without pre-processing.

## **6.6. run\_simExampleBasisSet.m**

**USAGE:**

This script is run simply by editing the input parameters and then clicking "Run".

**DESCRIPTION:**

Script to generate simulated basis spectra for all metabolites of interest in the human brain. The script will generate an LCModel format .RAW file for each metabolite basis spectrum, which can then be passed into LCModel's "makebasis" function to generate a complete LCModel basis set.

**INPUTS:**

To run this script, edit the following parameters as desired and then click run:

lb = linewidth (Hz)

np = Spectral points

sw = Spectral width (Hz)

Bo = Magnetic Field Strength (Tesla)

te1 = First PRESS echo time, or SPECIAL echo time (s)

te2 = Second PRESS echo time (if applicable) (s).

seq = Pulse sequence ('se'= SPECIAL, 'p'=press, 'st'=steam);

ref = Add reference peak at 0ppm (used in LCModel, y or n);

## **6.7. run\_simMegaPressShaped.m**

**USAGE:**

This script is run simply by editing the input parameters and then clicking "Run".

**DESCRIPTION:**

This script simulates a MEGA-PRESS experiment with fully shaped editing and refocusing pulses. Phase cycling of both the editing and refocusing pulses is performed. Furthermore, simulations are run at various locations in space to account for the within-voxel spatial variation of the GABA signal. Summation across phase cycles and spatial positions is performed. As a result of the phase

cycling and spatially resolved simulations, this code takes a long time to run. Therefore, the MATLAB parallel computing toolbox (parfor loop) was used to accelerate the simulations. Acceleration is currently performed in the direction of the slice selective pulse along the x-direction, but this can be changed. Up to a factor of 12 acceleration can be achieved using this approach. To enable the use of the MATLAB parallel computing toolbox, initialize the multiple worked nodes using "matlabpool size X" where "X" is the number of available processing nodes. If the parallel processing toolbox is not available, then replace the "parfor" loop with a "for" loop.

#### INPUTS:

To run this script, edit the parameters below as desired and then click "run":

```

refocWaveform    = name of refocusing pulse waveform.
editWaveform     = name of editing pulse waveform.
editOnFreq       = frequency of edit on pulse[ppm]
editOffFreq      = frequency of edit off pulse[ppm]
refTp            = duration of refocusing pulses[ms]
editTp           = duration of editing pulses[ms]
Bfield           = Magnetic field strength in [T]
Npts             = number of spectral points
sw               = spectral width [Hz]
Bfield           = magnetic field strength [Tesla]
lw               = linewidth of the output spectrum [Hz]
thkX             = slice thickness of x refocusing pulse [cm]
thkY             = slice thickness of y refocusing pulse [cm]
x                = vector of X positions to simulate [cm]
y                = vector of y positions to simulate [cm]
taus             = vector of pulse sequence timings [ms]
spinSys          = spin system to simulate
editPhCyc1       = vector of phase cycling steps for 1st editing pulse [degrees]
editPhCyc2       = vector of phase cycling steps for 2nd editing pulse [degrees]
refPhCyc1        = vector of phase cycling steps for 1st refocusing pulse
                  [degrees]
refPhCyc2        = vector of phase cycling steps for 2nd refocusing pulse
                  [degrees]

```

## 6.8. run\_simMegaPressShapedEdit.m

#### USAGE:

This script is run simply by editing the input parameters and then clicking "Run".

#### DESCRIPTION:

This script simulates a MEGA-PRESS experiment with fully shaped editing pulses. Phase cycling of editing pulses is performed, and summation across phase cycles is performed.

#### INPUTS:

To run this script, edit the parameters below as desired and then click "run":

```

editWaveform     = name of editing pulse waveform.
editOnFreq       = frequency of edit on pulse[ppm]
editOffFreq      = frequency of edit off pulse[ppm]
editTp           = duration of editing pulses[ms]
Npts             = number of spectral points
sw               = spectral width [Hz]
Bfield           = magnetic field strength [Tesla]
lw               = linewidth of the output spectrum [Hz]
taus             = vector of pulse sequence timings [ms]

```

```

spinSys           = spin system to simulate
editPhCyc1        = vector of phase cycling steps for 1st editing pulse
                  [degrees]
editPhCyc2        = vector of phase cycling steps for 2nd editing pulse
                  [degrees]

```

### 6.9. run\_simMegaPressShapedRefoc.m

#### USAGE:

This script is run simply by editing the input parameters and then clicking "Run".

#### DESCRIPTION:

This script simulates a MEGA-PRESS experiment with fully shaped refocusing pulses. Phase cycling of the refocusing pulses is performed and simulations are run at various locations in space to account for the within-voxel spatial variation of the GABA signal. Summation across spatial positions is performed. As a result of the phase cycling and spatially resolved simulations, this code takes a long time to run. Therefore, the MATLAB parallel computing toolbox (parfor loop) is used to accelerate the simulations. Acceleration is currently performed in the direction of the slice selective pulse along the x-direction, but this can be changed. Up to a factor of 12 acceleration can be achieved using this approach. To enable the use of the MATLAB parallel computing toolbox, initialize the multiple worked nodes using "matlabpool size X", where "X" is the number of available processing nodes. If the parallel processing toolbox is not available, then replace the "parfor" loop with a "for" loop.

#### INPUTS:

To run this script, edit the parameters below as desired and then click "run":

```

refocWaveform      = name of refocusing pulse waveform.
refTp              = duration of refocusing pulses[ms]
Npts               = number of spectral points
sw                 = spectral width [Hz]
Bfield             = magnetic field strength [Tesla]
lw                 = linewidth of the output spectrum [Hz]
thkX               = slice thickness of x refocusing pulse [cm]
thkY               = slice thickness of y refocusing pulse [cm]
x                  = vector of X positions to simulate [cm]
y                  = vector of y positions to simulate [cm]
taus               = vector of pulse sequence timings [ms]
spinSys            = spin system to simulate
editFlipON         = vector of edit-ON pulse flip angles for each spin in spin
                  system.
editFlipOFF        = vector of edit-OFF pulse flip angles for each spin in spin
                  system.
refPhCyc1          = vector of phase cycling steps for 1st refocusing pulse
                  [degrees]
refPhCyc2          = vector of phase cycling steps for 2nd refocusing pulse
                  [degrees]

```

### 6.10. run\_simMegaSpecialShaped.m

#### USAGE:

This script is run simply by editing the input parameters and then clicking "Run".

#### DESCRIPTION:

This script simulates a MEGA-SPECIAL experiment with fully shaped editing and refocusing pulses. Phase cycling of both the editing and refocusing pulses is

performed. Furthermore, simulations are run at various locations in space to account for the within-voxel spatial variation of the GABA signal. Summation across phase cycles and spatial positions is performed. As a result of the phase cycling and spatially resolved simulations, this code takes a long time to run. Therefore, the MATLAB parallel computing toolbox (parfor loop) was used to accelerate the simulations. Acceleration is performed in the direction of the slice selective pulse (along the x-direction). Up to a factor of 12 acceleration can be achieved using this approach. To enable the use of the MATLAB parallel computing toolbox, initialize the multiple worked nodes using "matlabpool size X" where "X" is the number of available processing nodes. If the parallel processing toolbox is not available, then replace the "parfor" loop with a "for" loop.

#### INPUTS:

To run this script, edit the parameters below as desired and then click "run":

```

refocWaveform    = name of refocusing pulse waveform.
editWaveform     = name of editing pulse waveform.
editOnFreq       = frequency of edit on pulse[ppm]
editOffFreq      = frequency of edit off pulse[ppm]
refTp            = duration of refocusing pulses[ms]
editTp           = duration of editing pulses[ms]
Bfield           = Magnetic field strength in [T]
Npts             = number of spectral points
sw               = spectral width [Hz]
Bfield           = magnetic field strength [Tesla]
lw               = linewidth of the output spectrum [Hz]
thkX             = slice thickness of x refocusing pulse [cm]
x                = vector of X positions to simulate [cm]
taus             = vector of pulse sequence timings [ms]
spinSys          = spin system to simulate
editPhCyc1       = vector of phase cycling steps for 1st editing pulse [degrees]
editPhCyc2       = vector of phase cycling steps for 2nd editing pulse [degrees]
refPhCyc         = vector of phase cycling steps for 1st refocusing pulse
                  [degrees]

```

### 6.11. run\_simSpinEchoShaped.m

#### USAGE:

This script is run simply by editing the input parameters and then clicking "Run".

#### DESCRIPTION:

This script simulates a localized spin-echo experiment with a fully shaped refocusing pulse. Phase cycling of the refocusing pulse is performed. Furthermore, simulations are run at various locations in space (1-D) to account for the within-voxel spatial variation of the metabolite signal due to J-evolution. Summation across phase cycles and spatial positions is performed. As a result of the phase cycling and spatially resolved simulations, this code takes a long time to run. Therefore, the MATLAB parallel computing toolbox (parfor loop) was used to accelerate the simulations. Acceleration is performed in the direction of the slice selective refocusing pulse. Up to a factor of 12 acceleration can be achieved using this approach. To enable the use of the MATLAB parallel computing toolbox, initialize the multiple worked nodes using "matlabpool size X", where "X" is the number of available processing nodes. If the parallel processing toolbox is not available, then replace the "parfor" loop with a "for" loop.

#### INPUTS:

To run this script, edit the parameters below as desired and then click "run":

```

RFWaveform       = name of refocusing pulse waveform.
Tp               = duration of refocusing pulses[ms]
Bfield           = Magnetic field strength in [T]

```



Npts = number of spectral points  
 sw = spectral width [Hz]  
 lw = linewidth of the output spectrum [Hz]  
 thk = slice thickness of refocusing pulse [cm]  
 pos = vector of positions to simulate [cm]  
 TE = Echo-Time [ms]  
 spinSys = spin system to simulate  
 PhCyc = vector of phase cycling steps for refocusing pulse [degrees]

## 6.12. run\_specialproc.m

### USAGE:

```
[out,out_w,out_noproc,out_w_noproc]=run_specialproc(filestring,aaDomain,
tmaxin,iterin);
```

### DESCRIPTION:

Processing script for Siemens SPECIAL MRS data in .dat format (twix raw data). Includes combination of receiver channels, removal of bad averages, frequency drift correction, and leftshifting.

### INPUTS:

filestring: String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '\_w/'];  
 aaDomain: (Optional) Perform the spectral registration (drift correction) using the full spectrum ('t'), or only a limited frequency range ('f'). Default is 'f'.  
 tmaxin: (Optional). Duration (in sec.) of the time domain signal used in the spectral registration (drift correction). Default is 0.2 sec.  
 iterin: (Optional). Maximum number of allowed iterations for the spectral registration to converge. Default is 20.

### OUTPUTS:

out: Fully processed, water suppressed output spectrum.  
 out\_w: Fully processed, water unsuppressed output spectrum.  
 out\_noproc: Water suppressed output spectrum without pre-processing (No bad-averages removal, no frequency drift correction).  
 out\_w\_noproc: Water unsuppressed output spectrum without pre-processing.

## 6.13. run\_specialproc\_fmrs.m

### USAGE:

```
[out_stimOFF,out_stimON,out_w]=run_specialproc_fmrs(filestring,blockDesign,
leadingAveragesToRmv);
```

### DESCRIPTION:

Processing script for functional MRS data acquired using the SPECIAL MRS sequence. This script accepts data in Siemens .dat format (twix raw data). Processing steps include combination of receiver channels, removal of bad averages, frequency drift correction, and leftshifting. Also includes prior knowledge of the stimulation paradigm (given by the 'blockDesign' vector) and returns the averaged stimulus OFF and stimulus ON spectra.

### INPUTS:

filestring: String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '\_w/'];

**blockDesign:** This is a vector of positive and negative even integers that make up the ON/OFF block design. Each integer represents the number of sequential averages in a block. Positive integers refer to ON blocks, and negative integers refer to OFF blocks. For example, for a block design consisting of 30 OFF averages followed by 20 ON averages followed by 10 OFF averages, the blockDesign vector would be: [-30 20 -10];

**leadingAvgToRmv:** The number of averages to omit from the beginning of each block. This is done to account for a lag in the neurochemical response to stimulus. Must be an even integer. (optional. Default=0);

**OUTPUTS:**

**out\_stimOFF:** Fully processed water suppressed spectrum from the sum of the stimulus OFF periods.

**out\_stimON:** Fully processed water suppressed spectrum from the sum of the stimulus ON periods.

**out\_w:** Fully processed, water unsuppressed output spectrum.

#### **6.14. run\_specialproc\_fmrs\_slidingWindow.m**

**USAGE:**  
[out1,out\_w]=run\_specialproc\_fmrs\_slidingWindow(filestring>windowSize);

**DESCRIPTION:**  
Processing script for functional MRS data acquired using the SPECIAL MRS sequence. This script accepts data in Siemens .dat format (twix raw data). Processing steps include combination of receiver channels, removal of bad averages, frequency drift correction, and leftshifting. This code generates a 'sliding window timecourse' of MR spectra by combining the averages within a small window given by the windowSize argument, and then sliding the window by 1 average and combining again. Each summed window is output as an LCModel text file to be analyzed in LCModel. As a result, this function generates many text output files.

**INPUTS:**

**filestring:** String variable for the name of the directory containing the water suppressed .dat file. Water unsuppressed .dat file should be contained in [filestring '\_w/'];

**windowSize:** This is an integer that specifies the number of averages that are stored within the sliding window. It is recommended to choose a window size that is divisible by the number of phase cycles so that the window does not contain any partial phase cycles.

**OUTPUTS:**

**out1:** The first sliding window spectrum.

**out\_w:** Fully processed, water unsuppressed output spectrum.

## **7. Graphical User Interfaces (GUIs)**

Although the FID-A software package is primarily a MATLAB command-line software application, it does contain a three GUI based tools to enable visual manipulation of MR spectra. The three GUI based tools

are called DiffTool, SpecTool and subSpecTool, and they can be found in the {FID-A\_DIR}/processingTools/SpecTool/ Directory. A brief description of each tool is given below:

### 7.1. DiffTool

The DiffTool Gui is used to visualize at the difference between two spectra, while interactively controlling the relative frequency, phase ( $0^{\text{th}}$  and  $1^{\text{st}}$  order), amplitude, and DC offset between them. This tool is useful to enable subtraction of two spectra whilst minimizing subtraction errors due to frequency and phase shifts, etc. The DiffTool GUI is called in the following way:

```
DiffTool(in1,in2,ppmmin,ppmmax);
```

Where in1 and in2 are the two spectra to be subtracted, and ppmmin and ppmmax are the minimum and maximum frequencies over which to display the result. The DiffTool GUI is shown in Figure 2, below.

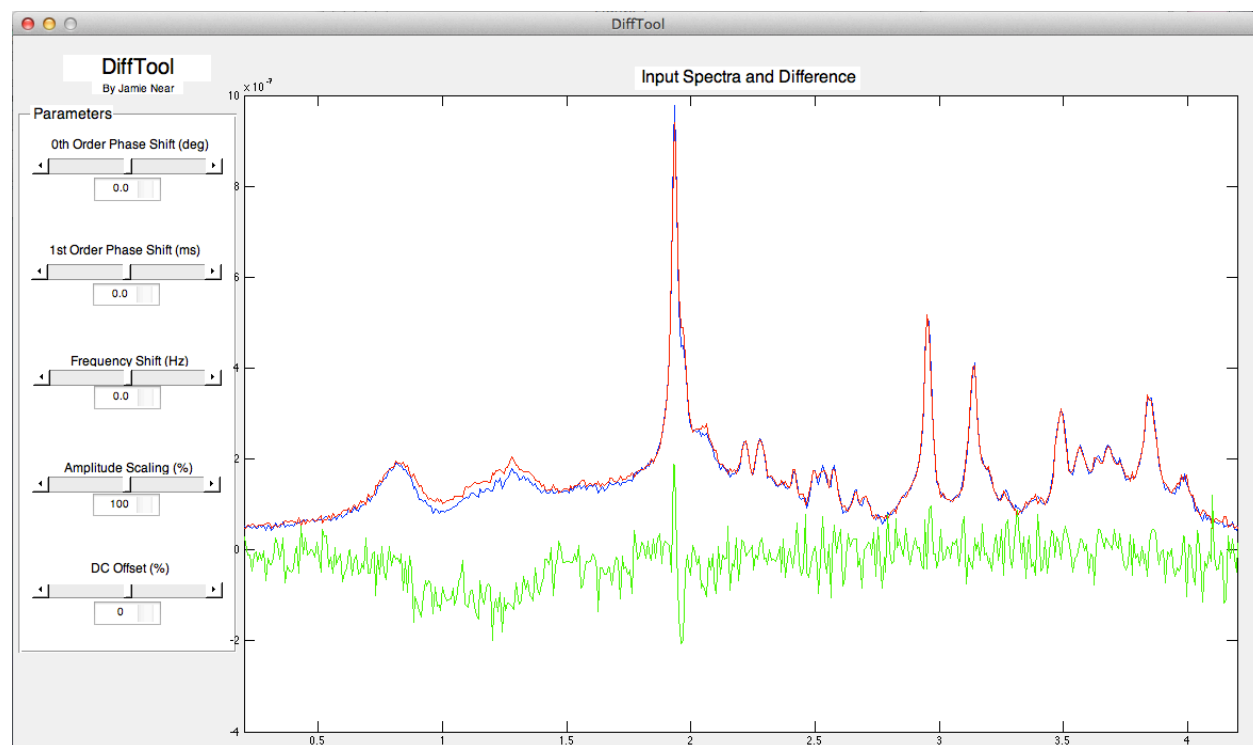


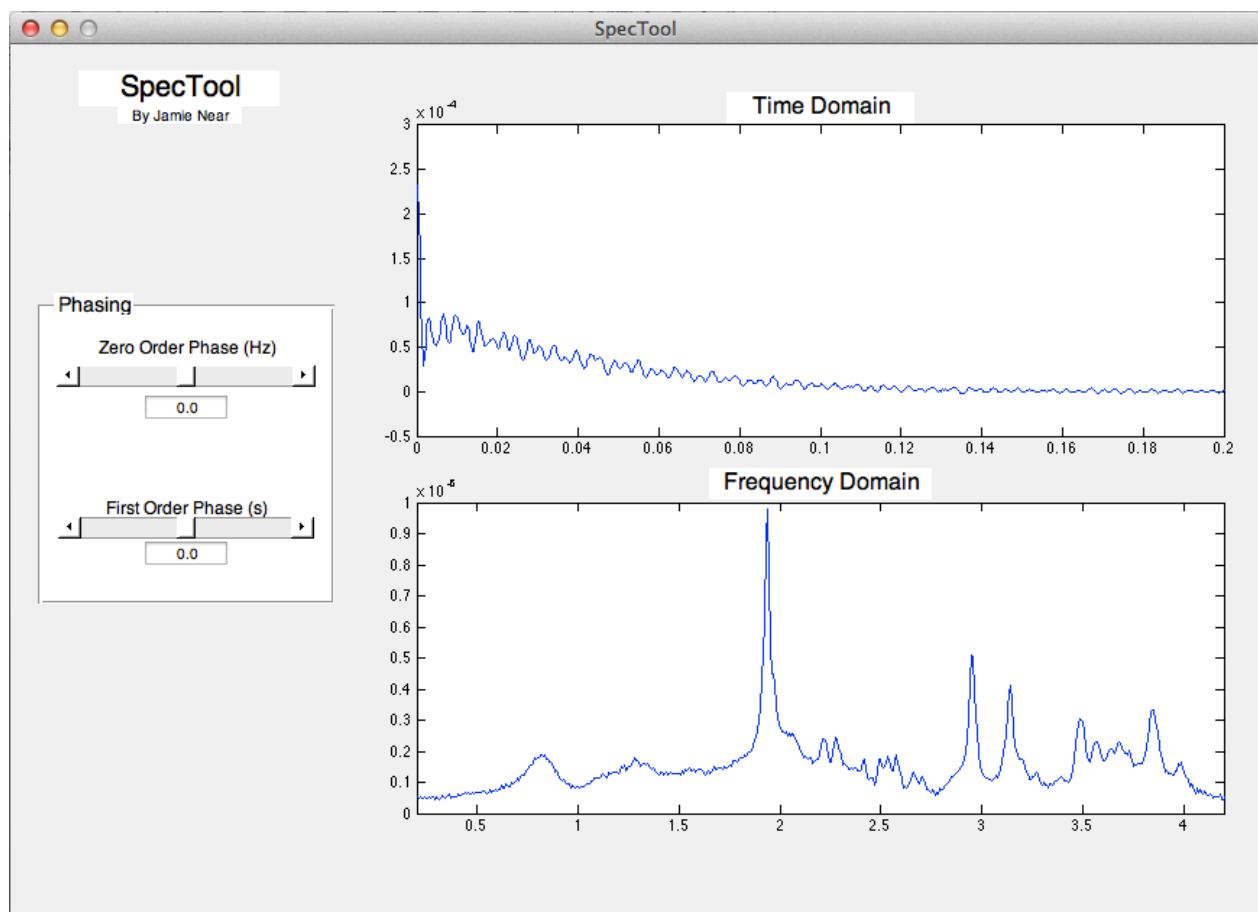
Figure 2: The DiffTool GUI.

## 7.2. SpecTool

The SpecTool GUI enables adjustment of the 0<sup>th</sup> and 1<sup>st</sup> order phase of the spectrum, while providing real-time visual feedback of both the time-domain and frequency-domain signals. The SpecTool GUI is called in the following way:

```
SpecTool(in,tmax,ppmmin,ppmmax);
```

Where `in` is the input spectrum, `tmax` is the maximum time over which to display the result in the time domain, and `ppmmin` and `ppmmax` are the minimum and maximum frequencies over which to display the result in the frequency domain. The SpecTool GUI is shown in Figure 3 below.



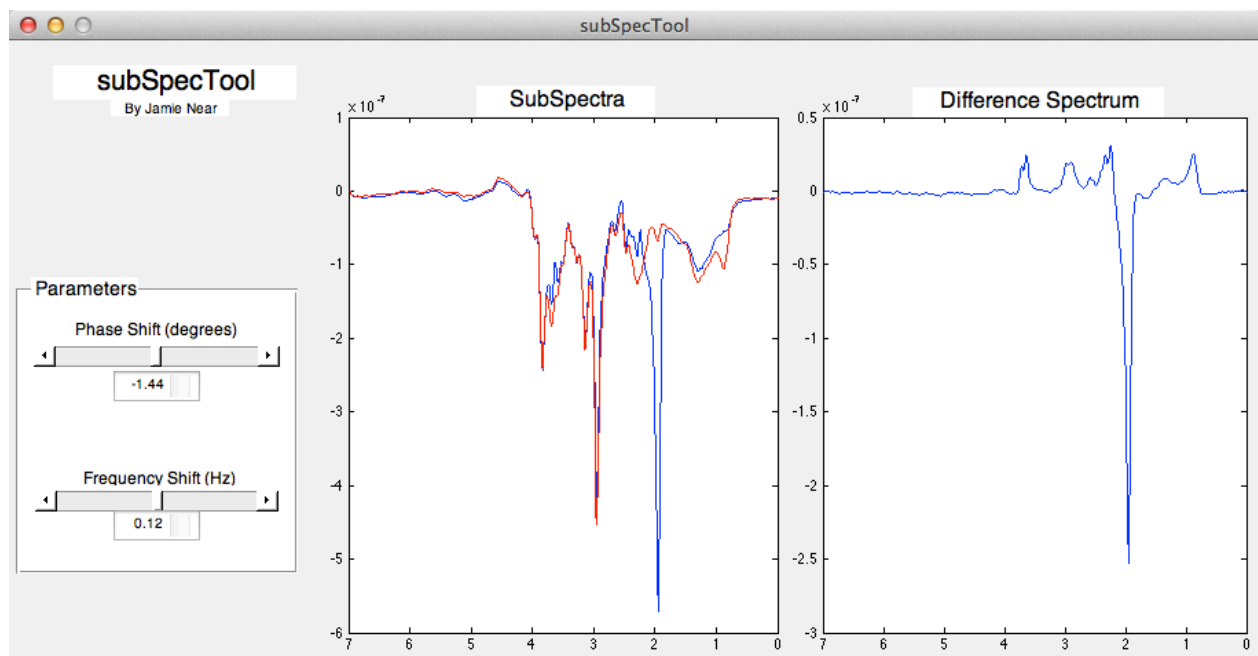
**Figure 3:** The SpecTool GUI.

### 7.3. subSpecTool

The subSpecTool GUI is used to visualize at the difference between two subspectra of an edited MRS scan, while interactively controlling the relative frequency and phase (0th order only), between them. This tool is useful to enable subtraction of two subspectra whilst minimizing subtraction errors due to frequency and phase shifts, etc. The subSpecTool GUI is called in the following way:

```
subSpecTool(in,ppmmin,ppmmax);
```

Where in is the input spectrum, and ppmmin and ppmmax are the minimum and maximum frequencies over which to display the result in the frequency domain. The subSpecTool GUI is shown in Figure 4 below.



**Figure 4:** The subSpecTool GUI.