

# **Vespa – DataSim User Manual and Reference**

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# Overview of the Vespa Package

Vespa stands for “Versatile Simulation, Pulses and Analysis”. The Vespa package is an integrated, open-source, platform for magnetic resonance spectroscopy (MRS) research and data analysis. Applications in the Vespa package include:

- Vespa-Simulation – software for spectral simulation (using the GAMMA library)
- Vespa-Pulse – software for RF pulse design
- Vespa-Analysis – spectral data processing and analysis
- Vespa-DataSim – used to create ‘fake’ MRS data sets from Simulation results

The Vespa project addresses software limitations seen across the MRS field, including: non-standard data access, closed source multiple language software that complicates algorithm extension and comparison, lack of integration between programs for sharing prior information, and incomplete or missing documentation and educational content.

## Installation

Vespa is a Python package. It requires that a Python environment be installed with certain required additional modules (all easily obtained) and then Vespa can be automatically installed from the PyPI web site. For historical reasons, the package on PyPI is called Vespa-Suite. As of version 1.0.0, Vespa only runs under Python 3.

Full instructions for installation can be accessed online at:

<https://scion.duhs.duke.edu/vespa/project/wiki/HowToInstallVespa>

Vespa is an actively developed project that has frequent releases. Once you have it installed, it is easy to upgrade as described here:

<https://scion.duhs.duke.edu/vespa/project/wiki/Upgrading>

## Other Online Resources

The Vespa project and each of its applications have Trac Wiki sites with extensive information about how to use, and develop new functionality for, each application. These can be accessed through the main portal site at

<https://scion.duhs.duke.edu/vespa/>

# Introduction to Vespa-DataSim

Vespa-DataSim is application written in the Python programming language that allows users to create simulated MR data sets. Vespa-DataSim allows users to:

- 1) Select from existing Vespa-Simulation Experiments in the Vespa database.
- 2) Specify a single set of Experiment loop parameters (if more than one) on which to base the datasim's metabolite basis functions.
- 3) Specify the spectral resolution as the bandwidth and number of points in the FIDs.
- 4) Scale individual metabolite areas.
- 5) Create a Voigt lineshape envelope that has a specific  $T_a$  ( $T_2$ ) value for each metabolite and a global  $T_b$  ( $T_2^*$ ) value for the entire datasim.
- 6) Add unlimited Gaussian spectral peaks with independent area, ppm, linewidth and phase to create simulated macromolecular signals.
- 7) Add unlimited Gaussian spectral peaks with independent area, ppm, linewidth and phase to create simulated metabolite baseline signals.
- 8) SNR can be set using an independent reference line (not included in spectrum) to maintain RMS noise levels even as metabolite line shape changes.
- 9) Output to Vespa-Analysis compatible data format or a simple text/binary header/data format.
- 10) Output a single simulated spectrum or an array of spectra, containing the same signal model but different noise, for use in Monte Carlo evaluations.

**What is a DataSim?** A 'DataSim' consists of a simulated MR spectroscopy (MRS) data set that has been created using prior information from a Vespa-Simulation Experiment. The combination of "simulated data set" has been shortened to "datasim" in this manual. A datasim can only contain metabolites from a single set of loop parameters from the Experiment. On output, the datasim contains header information about all signals, processing and noise that were performed to create the data.

**What is the DataSim Notebook?** This is the main window of the DataSim application. It contains one or more tabs, each of which contains the data and processing for an entire datasim. Multiple datasims can be created/loaded as tabs in the DataSim notebook.

DataSims are created in four steps, organized as four sub-tabs in each DataSim tab. These sub-tabs are 'Spectral Settings', 'Metabolite Signals', 'Macromolecular Signals', and 'Baseline Signals'. Upon output, a full provenance for sub-tab parameters is created as part of the DataSim XML output data format. A variety of graphical and text-based methods are available for saving results, as well.

The following chapters run through the operation of the Vespa-DataSim program both in general and widget by widget.

In this manual, command line instructions will appear in a fixed-width font on individual lines, for example:

```
~/Vespa-DataSim/ % ls
```

Specific file and directory names will appear in a fixed-width font within the main text.

Online Resources:

The Vespa project and each of its applications have Trac Wiki sites with extensive information about how to use, and develop new functionality for, each application. These can be accessed through the main portal site at

<http://scion.duhs.duke.edu/vespa/>

# Using DataSim – A User Manual

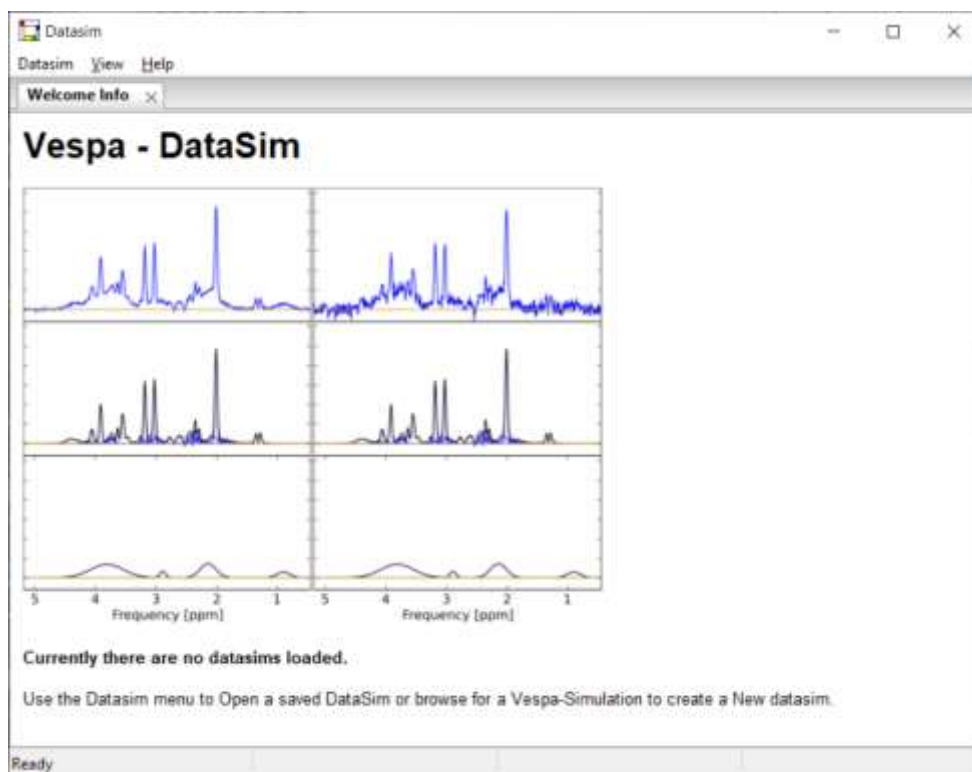
*This section assumes Vespa-DataSim has been downloaded and installed. See the Vespa Installation guide on the Vespa main project wiki for details on how to install the software and package dependencies. <http://scion.duhs.duke.edu/vespa>.*

In the following, screenshots are based on running DataSim on the Windows OS, but aside from starting the program, the basic commands are the same on all platforms.

## 1. Overview – How to launch Vespa-DataSim

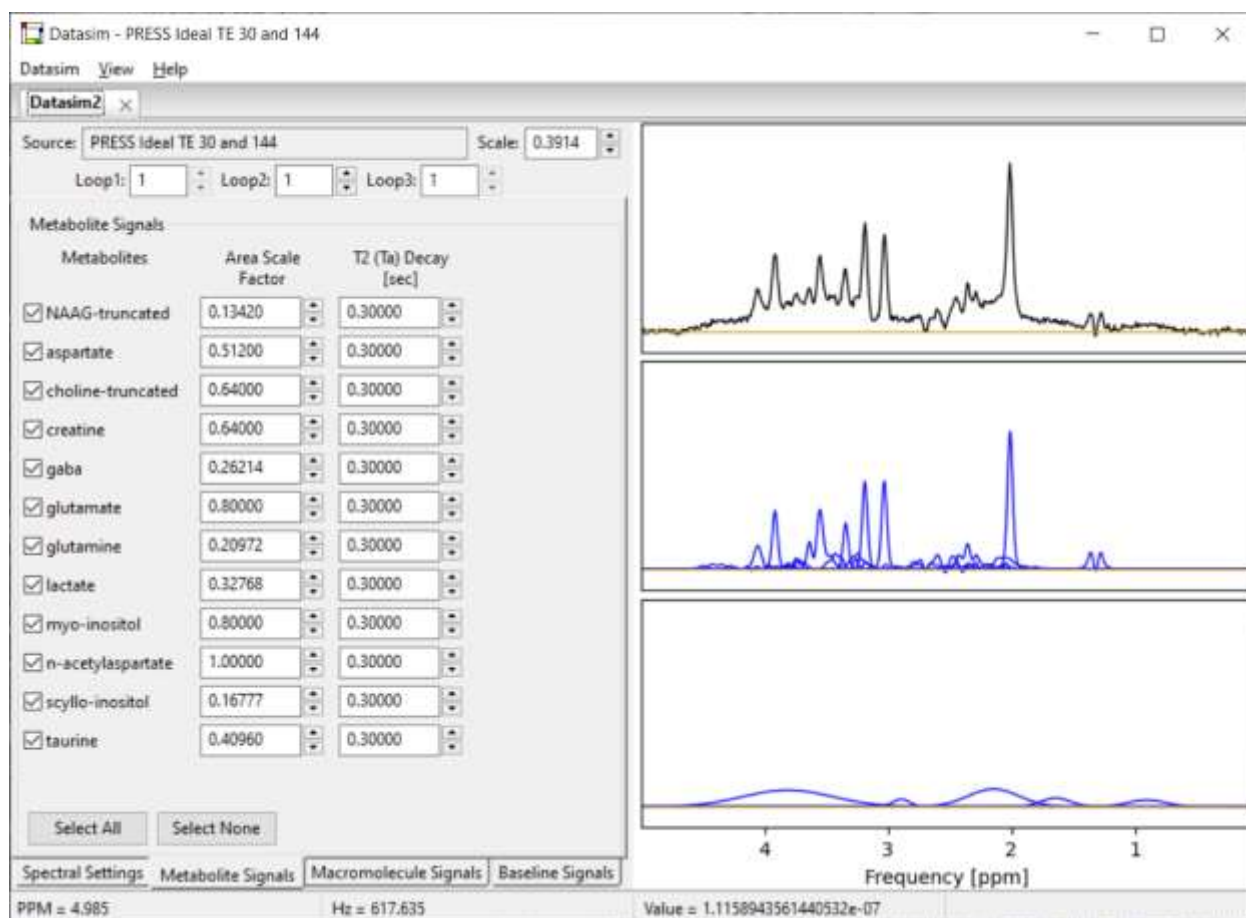
Double click on the DataSim icon that the installer created on your Desktop.

Shown below is the Vespa-DataSim main window as it appears on first opening. No actual DataSim windows are open, only the 'Welcome' banner is displayed.



Use the **DataSim → Open DataSim** menu to open saved datasims into tabs, or the **DataSim → New DataSim from Experiment** menu to create a new datasim.

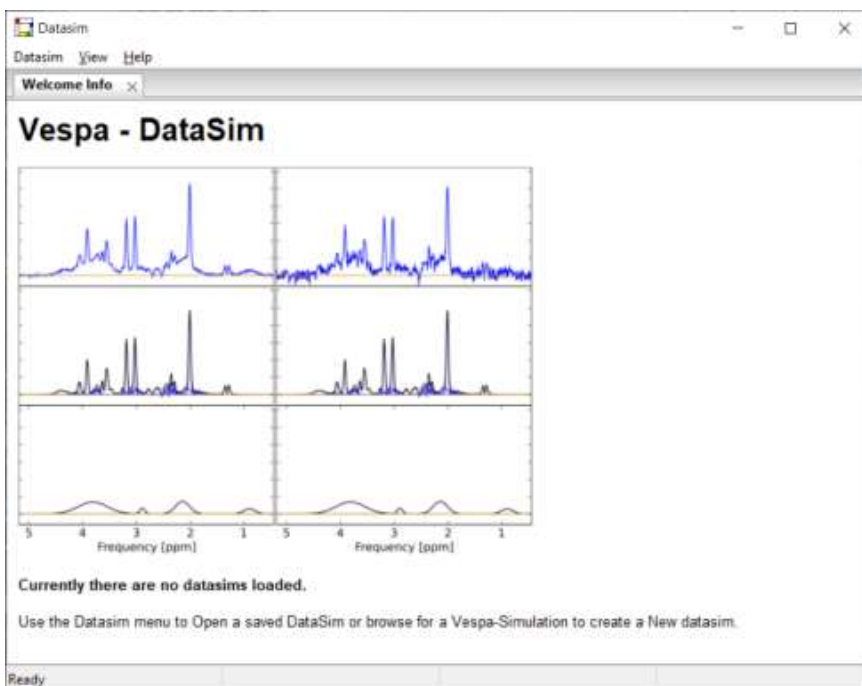
Shown below is a screen shot of a Vespa-DataSim session showing data from a PRESS 30ms simulation. The functionality of all sub-tabs will be described further in the following sections.



## 2. The DataSim Main Window

This is a view of the main Vespa-DataSim user interface window. It is the first window that appears when you run the program. It contains the datasim notebook, a menu bar and status bar. The datasim notebook can be populated with one or more datasim tabs, each of which contains input settings and simulated data from one datasim. As described above, a datasim is comprised of spectral settings, metabolite signals and baseline signals, each of which has its own sub-tab in the respective datasim tab. Sub-tabs are organized along the bottom edge, while datasim tabs are organized along the top edge.

The datasim Notebook is initially populated with a welcome text window, but no datasims are loaded. From the DataSim menu bar you can 1) open a datasim that has previously been processed by DataSim and then saved into the DataSim VIFF XML format, or 2) create a new datasim. In either case a tab will appear for each datasim that is opened or imported. The View menu items set the plotting options for whichever sub-tab is active. The Help menu provides links to useful resources.



The status bar provides information about where the cursor is located within the various plots in the interface throughout the program. During plot zooms or region selections, it also provides useful information about the cursor start and end points and the distance between. Finally, it also reports short messages that reflect current processing while events are running.

### On the Menu Bar

#### **DataSim→Open**

Opens an existing VIFF datasim XML file into a new datasim tab in the datasim Notebook. The state of the datasim as it was saved, including all sub-tab settings and simulated data, are restored as the datasim is opened into its tab.

#### **DataSim →New**

Allows the user to create a new datasim in a new Tab.

#### **DataSim →Save**

Saves the state of the datasim as it currently exists, including all sub-tab settings and data, into a VIFF (Vespa Interchange File Format) XML file.

*Note that this does NOT save out a simulated data set, but just a file that DataSim can read back in to recreate the current state shown in the program. In other words, it saves the DataSim program stuff, not the simulated data stuff. Use Export menu items (below) to save out simulated data.*

#### **DataSim →Save As**

Same as Save, but allows the user to change the file name into which the datasim is saved.

#### **DataSim → Close**

Closes the active tab.

#### **DataSim**



→ <b>Export Spectrum &lt;various&gt;</b>	Writes the simulated data from the datasim out to various formats. For formats that allow it, provenance for the simulated data is included. Otherwise, Provenance is saved to a separate text file in the same directory.
<b>DataSim</b>	
→ <b>Export Monte Carlo &lt;various&gt;</b>	Similar to Export Spectrum, but an array of spectral data with the same signal model, but different added noise is saved out to various formats.
<b>DataSim →Exit</b>	Closes the application window.
<b>View→&lt;various&gt;</b>	Changes plot options in the plots on each sub-tab of the active datasim tab, including: display a zero line, turn x-axis on/off or choose units, select the data type (real, imag, magn) displayed, and various output options for all plot windows and experiment in text format.
<b>Help→User Manual</b>	Launches the user manual (from vespa/docs) into a PDF file reader.
<b>Help→DataSim/Vespa Online Help</b>	Online wiki for the DataSim application and Vespa project
<b>Help→About</b>	Giving credit where credit is due.

## 2.1 The DataSim Notebook

The datasim notebook offers a lot of flexibility. Multiple tabs can be opened up inside the window. They can be moved around, arranged and “docked” as the user desires by left-click and dragging the desired tab to a new location inside the notebook boundaries. In this manner, the tabs can be positioned side-by-side, top-to-bottom or stacked. They can also be arranged in any mixture of these positions. There is only the one Notebook in the DataSim application, but it can display multiple simulated MRS data sets by loading them into DataSim tabs.

## 2.2 DataSim Tabs

The datasim notebook can be populated with one or more datasim tabs, each of which contains the spectral settings, metabolite signals, macromolecule signals and baseline signals of one datasim. Tabs for datasims are arranged along the top of the notebook and can be grabbed (left-click and drag) and moved to different locations inside the notebook as you like. Tabs can be closed using the X box on the tab or with a middle-click on the tab itself. When a tab is closed, the datasim is removed from memory, but can be restored to its current state at a future time - assuming it was saved to DataSim VIFF format.

Each datasim tab has four sub-tabs that represent the spectral settings, metabolite signals, macromolecule signals and baseline signals in the data. Each processing sub-tab is described in more detail in the following section.

DataSims are only saved to file when specifically requested by the user. On selecting **DataSim → Save**, the current state of the datasim, ie. all settings and results in all tabs, is saved into a file in the Vespa Interchange File Format, or VIFF. This file can be updated when desired by the user by again hitting Save, or a new filename can be used to save different states in different files by using **DataSim → Save As**. When a VIFF file is opened in DataSim, all tabs and results are restored to the state they were in upon save.

Each processing sub-tab displays name of the Simulation Experiment that provides the basis functions for the metabolites, the indices of the specific Experiment set of loop parameters that is being used, and the y-scale of the plot in the sub-tab.

The View menu on the main menu bar can be used to modify the display of the plots in the active sub-tab. The state of plot options in each sub-tab is maintained in each sub-tab as the user switches between them. The following lists the functions on the View menu item:

## **On the Menu Bar**

**View (this menu affects the plots in the currently active DataSim tab)**

→ZeroLine→Show	toggle zero line off/on
→ZeroLine→Top/Middle/Bottom	display the zero line in the top 10% region, middle or bottom 10% region of the canvas as it is drawn on the screen
→Xaxis →Show	display the x-axis or not
→Xaxis→PPM/Hz	x-axis value in PPM or Hz
→Data Type	select Real, Imaginary, or Magnitude spectral data to display
→Plot Views→Final/All Three	select whether only the final simulated spectrum is displayed, or the final spectrum, the metabolite signals and , macromolecule signals + baseline signals are each displayed in their own plot.
→Output Experiment Text	displays the selected Experiment object in text format in a separate text editor, similarly to what can be displayed in the Vespa-Simulation application.
→Output→View→<various>	writes the entire plot to file as either PNG, SVG, EPS or PDF format

## **2.3 Mouse Events in Plots**

Most processing sub-tabs have plots in their right hand panels. These plots may contain one or more axes which may change dynamically. Typically, the DataSim plot will show either one spectrum which is the sum of metabolite signals + macromolecule signals + baseline signals + noise, or three plots which display the summed spectrum, the metabolite spectrum and the baseline spectrum, respectively.

You can control a number of functions by using your mouse interactively within the plot area of most sub-tabs. Vespa-DataSim is best used with a ‘two-button’ mouse that has a scroll wheel, but can also work fine with a ‘two-button’ mouse, as most mouse-driven features for the scroll wheel also have a corresponding widget that can be clicked on or typed in to cause the same effect. The following describes the typical actions that can be effected using the mouse in a plot window. Any variations from this will be noted in the following sub-tab sections.

The mouse can be used to set the X-axis and cursor values in sub-tab plots. When there are three plots, the same X-axis or cursors are set on all three. The left mouse button sets the X-axis zoom range. Click and hold the left mouse button in the window and a vertical cursor will appear. Drag the mouse either left or right and a second vertical cursor will appear. PPM value changes will be reflected in the status bar. Release the mouse and the plot will be redisplayed for the axis span selected. This zoom span will display its range in a pale yellow that disappears when the left mouse is released. Click in place with the left button and the plot will zoom out to its max x- and y-axis settings.

In a similar fashion, two vertical cursors can be set inside the plot window. Click and drag, then release, to set the two cursors anywhere in the window. This cursor xpan will display as a light gray span. Click in place with the right mouse button and the xursor span will be turned off.

The cursor values are used to determine the “area under the peak” values that are displayed in the status bar. While performing a right-click and drag to create a cursor span the status bar will also display the start/end location of the span and the delta Hz and delta PPM size of the span.

The roller bar can be used to increment/decrement the Y-axis scale value. A maximum value for the Y-axis scale is determined the first time a datasim is loaded and displayed. That max value is the value displayed in the scale widget (top right in the datasim) and used when you zoom all the way out. As you roll the ball up/down (or you click on the SpinCtrl widget next to the scale field) the scale value changes and the plot is updated. (Note. It may be necessary to actually click in

the plot window to move the focus of the scroll wheel into the plot, before the scroll wheel events will be applied to the Scale value.)

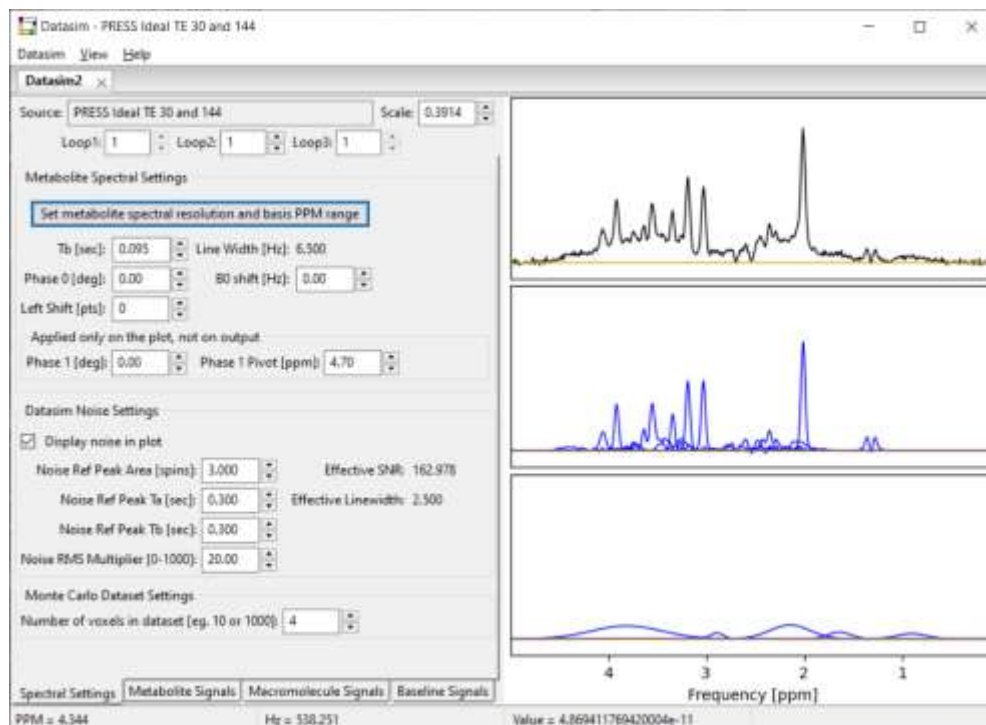
Roller balls can typically be used also as a 'middle button' but pushing down on it without rolling it up or down. DataSim plots do not make use of events from this middle button.

Click and release the left mouse button in place and the plot will zoom out to its max setting. Click and release the right mouse button in place and the cursor span will be turned off

### 3. Spectral Settings Sub-tab

#### 3.1 General

Each datasim tab has four sub-tabs called Spectral Settings, Metabolite Signals, Macromolecule Signals and Baseline Signals. The Spectral Settings tab is shown below.



#### Theoretical Details

The formula below describes how the simulated data signals are created in the time domain as a sum of metabolite  $M(t)$ , baseline  $B(t)$  and noise  $N(t)$  functions, and subsequently transformed into the spectral domain using the FFT for display in the DataSim plot panel.

The metabolite signal starts with a set of basis functions generated from the a priori Simulation Experiment information. The set of all a priori resonance lines for each metabolite are indexed over  $n$  from 1 to  $N_m$ . These sinusoids are created at the digital/spectral resolution set by the user. Only the metabolites selected by the user are included. Metabolite basis functions are modified by the user specified area ( $A_m$ ) and  $T_2$  decay ( $T_a$ ) and global frequency shift ( $\omega_0$ ), phase 0 ( $\phi_0$ ) and  $T_2^*$  decay ( $T_b$ ) terms.

If the 'Left Shift' parameter is not zero, then after the time domain FID has been calculated, it is truncated by  $N$  integer points (as set by the spectral resolution widget) before being Fourier transformed. The value  $N$  is taken from the Left Shift widget setting.

$$s(\omega) = FFT[M(t) + B(t) + N(t)]$$

$$N(t) = \text{numpy.random.randn}(\text{dims}[0])$$

$$B(t) = \sum_{b=1}^{Bas} A_b e^{-i[(\omega_b + \omega_0)t + \varphi_0]} e^{-\left[(t/T_b)^2\right]}$$

a priori relative area, frequency and phase of all resonances for each metabolite

$$M(t) = \sum_{m=1}^{Met} \sum_{n=1}^{N_m} A_m A_n e^{-i[(\omega_0 + \omega_n^m)t + \varphi_n^m + \varphi_0]} e^{-\left[\frac{t}{T_a^m} + (t/T_b)^2\right]}$$

$\uparrow$   
 metabolite area  
(concentration)  
of each metabolite

$\uparrow$   
 global  
frequency  
shift

$\uparrow$   
 global  
zero order  
phase

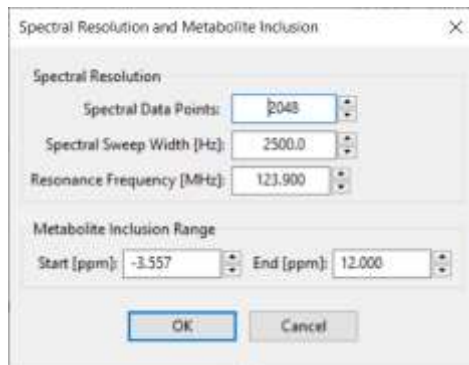
$\uparrow$   
metabolite  
specific  $T_2$   
decay

$\uparrow$   
 global  $T_2^*$   
decay

### 3.2 Metabolite Spectral Settings Section

The Metabolite Spectral Settings section contains controls that affect the spectral resolution, SNR and global spectral parameters for the simulated data. The data in the plot on the right will update interactively as you change parameters.

Spectral resolution and the PPM range of metabolite basis peaks is set by using the **Set metabolite spectral resolution and basis PPM range** button. This button will launch a dialog (shown below):



on which, you can set the **Frequency, Spectral Points, Spectral Width** and metabolite inclusion range in PPM using spin control widget. **Note** – changes to any of these widgets will result in a recalculation of all basis functions from the selected Experiment, but not until you hit the OK button. Depending on the number of steps in all Experiment parameter loops, this recalculation can take an appreciable amount of time. However, it does speed up display of different basis sets when the Loop indices change.

The **Phase 0** and **B0 shift** widget spin controls set these global values for the simulated data.

The **Tb** spin control is used to set effects of simulated  $T_2^*$  line broadening. An estimate of the resulting Line Width (in Hz) is displayed in the field below. This estimate is based on an assumed  $T_a$  ( $T_2$ ) value of 0.3 seconds because each metabolite can have its own  $T_2$  value, thus we use this value as a reasonable average.

### 3.3 Dataset Noise Settings Section

The Dataset Noise Settings section contains parameters that affect the amount of noise that is added to each simulated spectrum. The data in the plot on the right will update interactively as you change parameters. Note, the noise displayed is updated every time a parameter is changed, thus low SNR plots will change appreciably each time a parameter changes.

#### Background on SNR calculation

Arrays of random noise are created using the `numpy.random.randn()` method which creates random floats sampled from a univariate “normal” (Gaussian) distribution of mean 0 and variance 1. Two numpy ndarrays of the proper length are combined to create a complex set of random noise samples.

The normalized noise is scaled to create a particular SNR setting based on the traditional “peak height divided by RMS noise value” method. In DataSim, the RMS noise is taken directly from the `randn()` method, but the peak height is measured from a user defined ‘reference peak’ that is set up on the Spectral Settings sub-tab. The reference peak is not plotted as part of the final spectrum. Its definition is used to create a temporary peak at high spectral resolution whose peak height for a given line and sweep width can be measured. This is the peak height used to calculate the SNR value displayed in the **Effective SNR** field.

The reason we use the reference method is that it allows us to keep the same noise scaling factor as metabolite line width changes. Thus we can create datasims that have the same noise level and the same metabolite areas but at different peak line widths. It also allows users to use a simple definition of SNR in spectra that do not have simple (singlet) lines against which to set an SNR value.

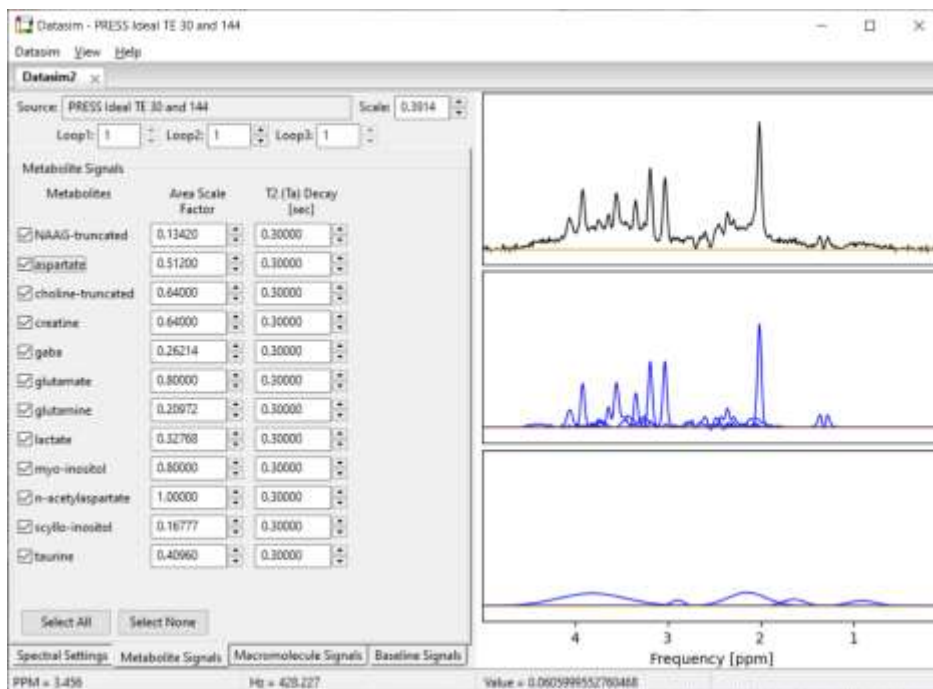
The default reference line is a singlet peak with 3 spins (aka.  $-\text{CH}_3$  group). The user sets the **Reference Peak  $T_a$**  and  **$T_b$**  values to achieve the desired **Effective Linewidth** value and then scales to the desired SNR level using the **Noise RMS Multiplier** spin control.

Finally, the user can select to turn the noise in the spectrum on/off using the **Display noise in the spectrum** check box. This is a convenience function only.

## 4. Metabolite Signals Sub-tab

### 4.1 General

Each datasim tab has four sub-tabs called Spectral Settings, Metabolite Signals, Macromolecule Signals and Baseline Signals. The Spectral Settings tab is shown below.



### 4.2 Metabolite Signals Parameters

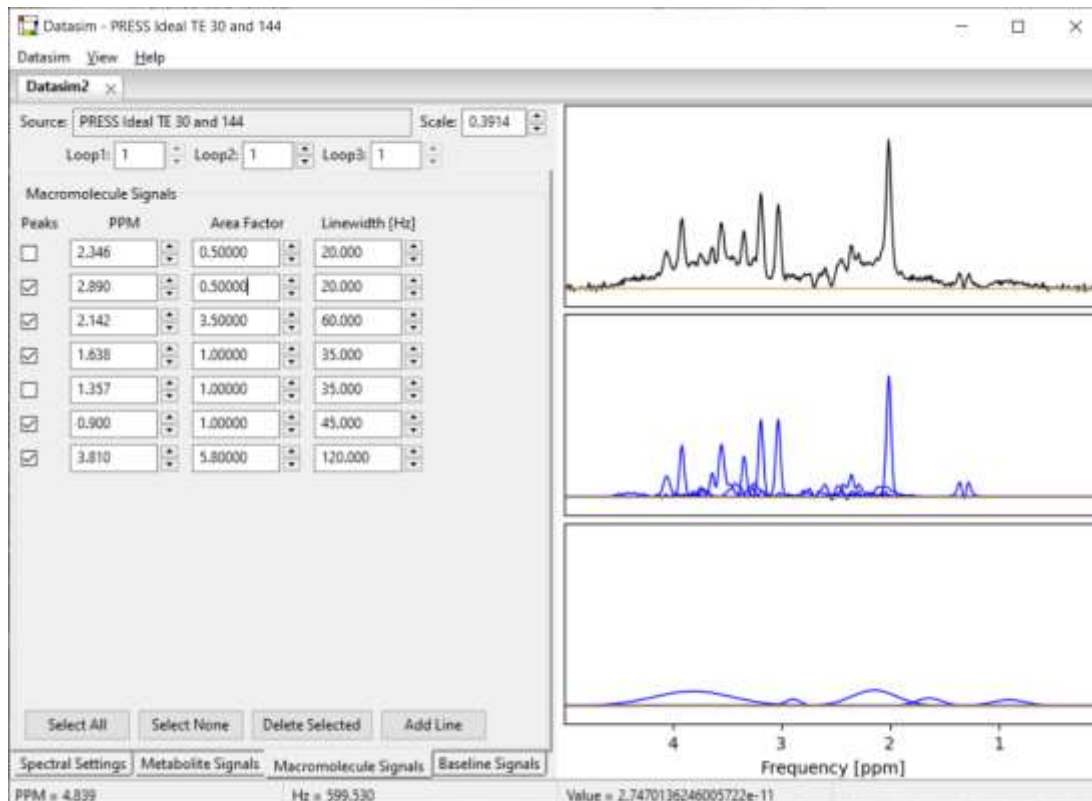
The metabolites included in the spectrum can be selected in two ways. First, as shown in the previous section, a metabolite inclusion range (in PPM) can be set using the 'spectral resolution' dialog. Depending on the spectral lines in each metabolite, the metabolite list is repopulated with metabolites that have lines within this range. Second, the user can explicitly select a metabolite for inclusion by clicking on the check box at the start of a line in the metabolite list. Metabolite areas and an intrinsic  $T_2$  decay value can be set in each line for each metabolite.

The plot to the right displays the spectrum you are designing. The top plot shows the summed metabolite + baseline + noise signals. The middle plot shows the summed metabolite signals (black) overlaid on the individual metabolite signals (blue). The bottom plot shows the summed macromolecule + baseline signals (black) overlaid on the individual baseline signals (blue). You can zoom in/out of this plot the same as described in Section 2.3. You will likely need to zoom in to clearly see the lines you are creating.

## 5. Macromolecule Signals Sub-tab

### 5.1 General

Each datasim tab has four sub-tabs called Spectral Settings, Metabolite Signals, Macromolecule Signals and Baseline Signals. The Macromolecule Signals tab is shown below.



### 5.2 Macromolecule Signals Controls

You can add/delete as many lines to the Macromolecule signals list as you like using the **Add Line**, **Delete Selected**, **Select All** and **Select None**. Only the lines that have a check in the box will be included in the calculated macromolecule signal. For each line, you can specify the PPM value for the peak center, the area of the peak and the Gaussian line width.

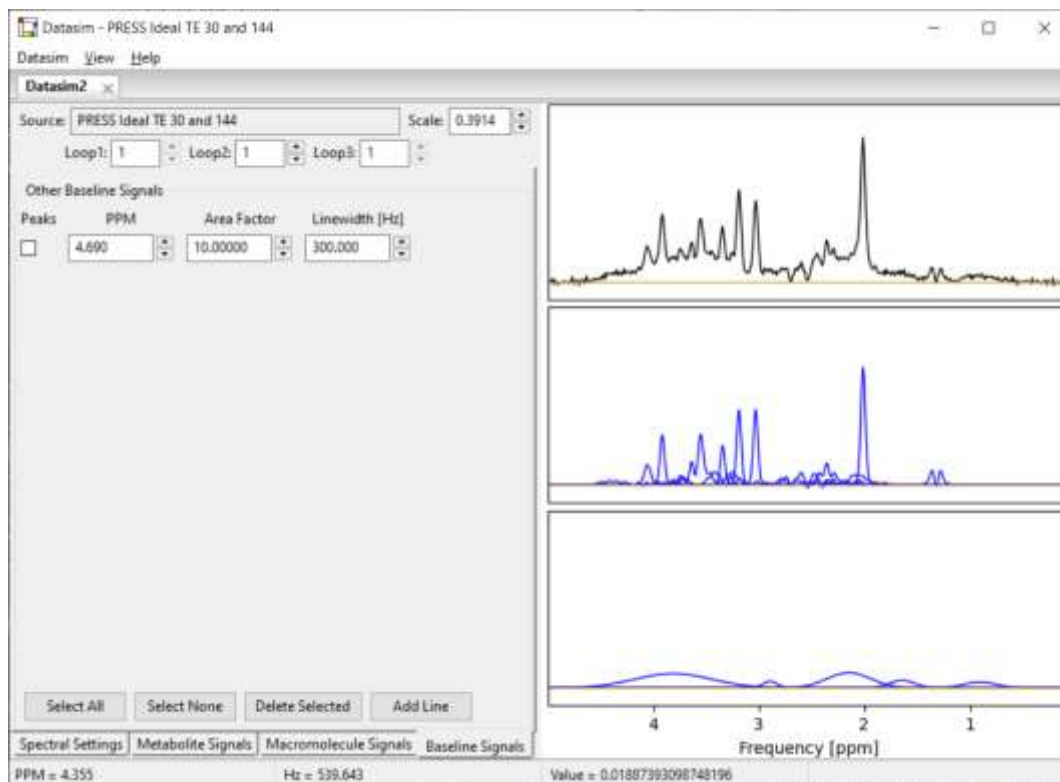
The plot to the right displays the spectrum you are designing. The top plot shows the summed metabolite + baseline + noise signals. The middle plot shows the summed metabolite signals (black) overlaid on the individual metabolite signals (blue). The bottom plot shows the summed macromolecule + baseline signals (black) overlaid on the individual macromolecule and baseline signals (blue). You can zoom in/out of this plot the same as described in Section 2.3. You will likely need to zoom in to clearly see the lines you are creating.



## 6. Baseline Signals Sub-tab

### 6.1 General

Each datasim tab has four sub-tabs called Spectral Settings, Metabolite Signals, Macromolecule Signals and Baseline Signals. The Baseline Signals tab is shown below.



### 6.2 Baseline Signals Controls

You can add/delete as many lines to the Baseline signals list as you like using the **Add Line**, **Delete Selected**, **Select All** and **Select None**. Only the lines that have a check in the box will be included in the baseline signal. For each line, you can specify the PPM value for the peak center, the area of the peak and the Gaussian line width.

The plot to the right displays the spectrum you are designing. The top plot shows the summed metabolite + baseline + noise signals. The middle plot shows the summed metabolite signals (black) overlaid on the individual metabolite signals (blue). The bottom plot shows the summed macromolecule + baseline signals (black) overlaid on the individual macromolecule and baseline signals (blue). You can zoom in/out of this plot the same as described in Section 2.3. You will likely need to zoom in to clearly see the lines you are creating.

## 7. Results Output

### 7.1 Plot results to image file formats

The plots displayed in all sub-tabs which contain View panels can all be saved to file in PNG (portable network graphic), PDF (portable document file) or EPS (encapsulated postscript) formats to save the results as an image. The Vespa-DataSim **View** menu lists commands that only apply to the active DataSim tab and selected processing sub-tab. Select the **View→OutputPlots→** option and further select either **View to PNG**, **View to EPS** or **View to PDF** item. The user will be prompted to pick an output filename to which will be appended the appropriate suffix.

### 7.2 Plot results to vector graphics formats

The plots displayed in all sub-tabs which contain View panels can all be saved to file in SVG (scalable vector graphics) or EPS (encapsulated postscript) formats to save the results as a vector graphics file that can be decomposed into various parts. This is particularly desirable when creating graphics in PowerPoint or other drawing programs. At the time of writing this, only the EPS files were readable into PowerPoint.

The Vespa-DataSim **View** menu lists commands that only apply to the active DataSim tab and selected processing sub-tab. Select the **View→Output→** option and further select either **Plot to SVG**, or **Plot to EPS** item. The user will be prompted to pick an output filename to which will be appended the appropriate suffix.

### 7.3 Simulated Data Output to File

The simulated data can be output to file in a number of formats. In all cases you will be prompted to select a director/filename for the output.

Use the **DataSim→Export Spectrum** and **DataSim→Export Monte Carlo** menu items to select if you want to output just the single spectrum shown on the plot, or if you want DataSim to create an output file with multiple spectra consisting of copies of the pure metabolite + macromolecule signals + baseline signals with different noise signals. Use the **Voxels in Dataset** spin control on the **Spectral Settings** sub-tab to indicate how many voxels to create in the Monte Carlo output file. Note, in a Monte Carlo output file the data will be stored as a two dimension data set. with **Spectral points** and **Voxels in Dataset** spin control values as the dimensions.

#### 7.3.1 VIFF – Vespa Interchange File Format

Both single spectrum and Monte Carlo datasims can be output to **VIFF** (Vespa Interchange File Format) **Raw Data** format. In both cases, the data output is time domain (k-space) data. The **VIFF Raw Data** format is importable into Vespa-Analysis and has the added benefit of being able to store a full provenance of the creation of the simulated data in its XML hierarchy.

Five files are output with provenance stored in each one's header. The user selects a base filename (fbase) and then extension strings are added to fbase to indicate the data in each file:

1. fbase\_metabolites.xml - time domain summed metabolite signals, no noise
2. fbase\_macromolecules.xml - time domain summed macromolecule signals, no noise
3. fbase\_baseline.xml - time domain summed baseline signals, no noise

- 4. fbase\_noise.xml                - the time domain added noise in the summed data
- 5. fbase\_summed.xml            - the sum of the above three files, final spectrum

### 7.3.2                **Siemens \*.RDA Format**

This is a simple MRS data export format supported on Siemens MR scanners. It consists of a text header with single “parameter = value” pairs on each line. At the end of the text header, the double complex floating point data is appended as a byte encoded string. See Siemens (non-existent) documentation for more details. However, this format has the benefit in that it can be read by a variety of MRS processing programs including LCModel. Vespa-Analysis can also read this format.

Only a single spectrum result can be saved to the **Siemens \*.RDA** file format. Six files are output, five data and one provenance. The user selects a base filename (fbase) and then extension strings are added to fbase to indicate which data is in the file:

- 1. fbase\_metabolites.rda    - time domain summed metabolite signals, no noise
- 2. fbase\_macromolecules.rda - time domain summed macromolecule signals, no noise
- 3. fbase\_baseline.rda        - time domain summed baseline signals, no noise
- 4. fbase\_noise.rda            - the time domain added noise in the summed data
- 5. fbase\_summed.rda          - the sum of the above three files, final spectrum
- 6. fbase\_provenance.rda    - text file, info about Vespa-DataSim setting used to create the above files. This is created because Siemens RDA has limited ability to store provenance information within its header.