Operating Procedure for the MERCURY 400 FT-NMR

I. Logging on spectrometer/inserting sample
When users are not employing the SMS sample changer, the spectrometer should remain logged in to the “organic” account. There are various “operators” in this account (who do not have passwords) and these operator sessions should be “exited” but not logged out of completely (after use, get to the gray screen – with large Varian logo, not the Shiley Science Center).

If the screen displays the Shiley Science Center, you should log in as “organic” using the correct password. If the screen displays the large Varian logo, log in as the appropriate operator.

Operators: organic (owner of account) bolender malachow
plnu (Pt. Loma Nazarene College) ddehaan debbiet
grossmnt (Grossmont College) tdwyer tshields
miramar (Miramar College) piovine
exp (Experimental Interface) leigh

At this point, a Windows screen appears and on the bottom panel is an icon (single click) this to launch vnmrj.

In the bottom, center window, under the Start tab click the first menu item entitled Study, then click on Eject. Remove the standard and place the sample (use depth gauge to set proper depth) on the air stack. Return to the console and click the Insert button.

II. Setting up the experiment(s)
  a. choose the experiment(s) you wish from the protocol list on the left, e.g. Proton, Carbon, Dept, DQCSOY, etc. This creates a “new sample” line in the automation window (below the protocol window).
  
b. in the bottom, center panel under the Start tab, click the first selection entitled Study. In this window, you must type in the sample name (NO SPACES) and select the solvent from the pulldown menu. Also in this panel, make sure the boxes on the right for “Find z0” and “Gradient shim” are “checked”. Here is where you would also check “Plot all data” if you want plots to be initiated automatically after acquisition.
  
c. If you wish to change any parameter for the experiment(s) prior to collecting data, you must do it now. To change any parameters, double click on the time field for the experiment in the protocol window (the field turns yellow with italic letters). To change acquisition parameters, click on the Acquire tab and then either the Default or Acquisition menu item and change the appropriate parameter. To change a processing parameter, click on the Process tab and then the Process menu item.

For example, some parameters you may wish to check and/or change:
  - Number of scans
  - Block size (the smallest increment of acquisitions after which you can view spectrum as it collects)
  - Line broadening (typically 0.5 Hz for proton, 2 Hz for carbon)
  - Spectral width

  d. When the parameters are set correctly, click on Submit.

While the experiment is running …
The “new sample” line changes to the sample name that you gave it in step II.b. If you fail to give a sample name, it will simply be named “s_#” where # is 01, 02, etc. Below that is a line
the states the type of experiment (proton, carbon, etc), followed by a third line that begins with
the length of time the experiment will take followed by the type of experiment. This third line
turns yellow during the lock/shim procedure (automated), then turns blue when the data is being
collected, then turns back to yellow when the experiment is finished (and the time field turns into
an icon of a lock). Also, at the bottom of the window, there is an acquisition status bar that turns
blue when the experiment is in progress and turns green (with the word Idle) when the
experiment is finished.

If you wish to abort an acquisition, click on the red Stop button next to the Process tab.

III. Viewing/adjusting/plotting the spectrum
Most spectral adjustments are made using the column of icons to the left of the spectral window
that will appear as you process the data.

a. view the entire spectrum: drag and drop the line beneath the name of the file in the
protocol window (the line begins with an icon of a lock). This will process the spectrum
with the parameters listed in the Process tab and display it in the spectral window. For the
time being, you can hide the integrals (for proton spectra) by clicking on the integration
icon (top, green integral - click it twice to turn them off). To display the chemical shift
scale under the spectrum, click on the scale icon (looks like a white ruler). You can
adjust the vertical scale of the spectrum using the middle mouse "wheel" (click) or set to
maximize the tallest peak by selecting the "Autoscale" button in the Process tab under the
Default or Display menu item.

b. phasing the spectrum: To autophase, click the Process tab, then the Proces menu item,
then click “Autophase Full”. To phase the spectrum manually, click on the phasing icon
that looks like a red clock 🕒. Then click and drag the left mouse button to phase the
downfield region; click and drag the right mouse button to phase the upfield region.
When finished phasing, click the cursor icon (very top icon with one or two vertical red
dlines).

c. expanding regions: use the left and right mouse buttons in the spectral display window to
place the red cursor lines on the downfield (left mouse button) and upfield (right mouse
button), then click on the icon that looks like a magnifying glass, 🕵️.

d. setting the reference peak: having chosen the solvent in II.c., the reference should be
automatically set. If an adjustment is needed, you can do this in the Process tab under the
Display menu item.

e. displaying peak frequencies: adjust the threshold for selecting peaks. Click on the icon
with the yellow horizontal lines on it 📉; click and drag the left mouse button to
position the yellow bar lower than the smallest peak you wish to be "seen". Click on icon
again to hide the bar. To display the frequencies on the screen, click on the "Peak
frequencies" button in the Process tab under the Display menu item.

IV. Integration
a. Start by selecting "Clear Integral Resets" in the Process tab under the Cursors/Integration
menu item. You can toggle on/off the integrals using the icon 📊.
b. Define the integration regions: click on the integral resets icon. Next, place the cursor arrow on the downfield side of the peak and click the left mouse button. Move to the upfield side of the peak and click the left mouse button again. Repeat for each peak of interest.

c. Correct the integral baselines: Click on the integral phasing icon and use the left/right mouse buttons to drag the phase and slope into adjustment (make the integral baselines parallel with the spectrum baseline).

V. Plotting the spectrum
In the Process tab, under the Plot menu item you can select what you wish to plot. In general, click the following: "Plot spectrum", "Plot Spectrum Scale", "Normalized Integrals" (for proton spectra). If you wish the peak frequencies to be marked above each peak, select the box "On peaks" under "Plot Peak Frequencies". To generate the plot, click on “Plot Page”.

VI. Shutdown
Remove sample by clicking on Eject in the Start tab under the Study menu item. Replace the standard (any sealed sample in CDCl₃) into the sample holder and replace the sample on the stack. Click on Insert. Exit vnmrj under the Utilities pulldown.