BARE BONES GUIDE TO 2D-COSY
on the VXRs-300 only

Note: On the Inova-500 and Inova-400, you should use gradient (gCOSY).

This guide is written assuming proficiency in basic operation of the Varian NMR instrument. You should be experienced in performing basic 1-dimensional NMR experiments before attempting to perform 2D experiments on your own. Please ask for help the first time you perform this, to minimize your frustration (if for no other reason).

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Conventions in this manual:

**Boldface** text indicates commands to be typed at the computer

<angle brackets> are used to designate a key to be pressed (i.e. <Ret> for Return/Enter)

[square brackets] designate an icon/button in the VNMR menu to be clicked

**Mouse Conventions:** click, by default, refers to the Left Mouse Button.
- LMB will be used to designate the Left Mouse Button
- MMB will be used to designate the Middle Mouse Button
- RMB will be used to designate the Right Mouse Button

Sometimes you will need to hold, rather than click the mouse button. This means that you should press and hold the button down throughout the operation.

Note: All **commands** in boldface assume that you press <Return> afterwards.

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1) Make sure you are working in Exp:1 or Exp:2 (jexp1, or jexp2).
2) Acquire a normal $^1$H NMR spectrum, setting NT=4.
3) Zoom-In on the region containing peaks of interest (Note: if you zoom-in on a region with peaks outside that region, you will have folded peaks in your spectrum...this can be OK, or it can be a problem; depending upon where the folded peaks land).
4) Type movesw, and this will set the spectral width and offset to match your selected window.
5) Type ga to acquire a new spectrum, then phase it.
6) Save this spectrum in your directory, as a 1-D trace for plotting beside your 2D contour plot.
7) Type mp(1,100) jexp100 (assuming you are working in Exp: 1, substitute "2" if you are starting in Exp:2). You should now be in Exp:100
8) Type cosy to load the COSY parameters.
9) Set NT (# transients):
   a) Usually nt=4 works fine (and takes the minimum amount of time).
   b) Use nt=8 for more dilute samples (will take 2x as long).
10) Set the number of increments in the $t_1$ dimension:
    a) Type ni?, and note the current value for ni.
    b) If the value of ni is < 128, type ni=128.
       i) Note: if you have time, ni=256 will give a better (higher-resolution) spectrum.
11) Type time to see how long the experiment will take.
12) Type go to start the experiment.
    a) After ~32 or more transients, you can type wft2d to see if your spectrum is coming along nicely. Otherwise, you can wait until it is done if you wish.
13) Type svf, followed by your filename, to save the data for processing.
14) Type jexp1, eject your sample, and re-insert the reference. Be sure to turn the spinner back on in [Acqi], and lock/shim on the reference before you sign-out.
Processing and Plotting the 2D-COSY spectrum:

1. Select your printer/plotter and paper size first.
2. Type `jexp1` to join Exp:1, and load the 1D-1H spectrum from step 8 on the previous page.
3. Type `jexp2` (or any other valid Exp:#), and load the 2D-COSY spectrum
4. Type `setLP1` to activate forward linear prediction (improves resolution and quality).
   a. Type `fn?<Ret>`, and note the value for fn. If greater than 2048, the processing will take a very long time.
   b. If fn>2048, type: `fn=2k<Ret>` then `fn1=fn <Ret>`.
5. Type `sinebell` to set the window-function parameters.
6. Type `wft2d` to process the data.
7. Type `foldt` to symmetrize the spectrum (recommended). This removes artifacts, but can reduce the information content of the spectrum in some cases.
8. If the contour plot is not centered, type `center` followed by `dconi`.
9. **NOTE**: the command `dconi` is analogous to “ds” for 1D spectra. If you lose your 2D image, and you want to view it, type `dconi` to re-display the 2D spectrum. You don’t need to re-type `wft2d` unless you want to try different processing parameters. In fact, re-processing every time will waste a lot of your time!

10. Adjusting the contour plot:
    a). Click the MMB on any spot in the 2d-display, to scale the spectrum so that spot just touches the floor of the contour plot. Clicking MMB on a taller peak will reduce the vs to make smaller peaks disappear.
    b). Alternatively, you can click MMB on the colored scale (on the right) to reduce the minimum level displayed. I recommend using (a), rather than (b).
    c). You can fine-tune the peaks displayed peaks by clicking on [vs+20%] or [vs-20%].
11. Plotting: there are lots of ways to do this…this is the recommended way.
    a). Plotting with your high-resolution 1D spectrum (in Exp: 1) displayed along the sides. Problem: If your 1D spectrum has very tall peaks, the size of your 2D plot will be reduced in size (sometimes, very small).
        1). Type `plcosy(15,1.5,1)` . This means: draw 15 contours, separated by 1.5 level units, and the 1D spectrum that is in Exp: 1 on the sides.
           a) Note: if your 1D spectrum is very small due to intense singlets, you can multiply the 1D trace by a factor of “n” by typing: `plcosy(15,1.5,1,n)` (i.e. plcosy(15,1.5,1,5) will multiply the 1D trace intensity by 5x).
        2). If using BIG (11”x17”) paper, substitute “plcosybig” for “plcosy”
    b). If you want to print to a PostScript file instead of to the printer, do the following:
        1). Substitute “plcosy_nopg” for “plcosy” above
        2). After executing `plcosy_nopg(15,1.5,1,n)`, type `page(‘filename.ps’)` and your plot will be saved in your directory as a PostScript file under the name “filename.ps”. Of course, use your own filename instead of “filename”.

-R.Shoemaker