¹³C 1D Spectrum Guide

Step	Function or Dialog Box	<keystroke>/[Select]/<data entry=""></data></keystroke>	Comment
			See Sample Preparation Guide. Position sample
1	Sample		spinner using the depth gauge, place in probe.
2	Enter PNMR program.	<alt+tab></alt+tab>	(If necessary.)
3	Select ¹³ C observe.	H1>nu c13 <enter></enter>	Required only if the prompt is not C13>.
3b	Optional: Shim sample.	C13>shim <enter></enter>	Follow on-screen directions. Shims are optimized. Dilute sample RD=5; Conc. sample RD=2
4	Acquire ¹ H spectrum.	C13>zgh <enter></enter>	Obtain ¹ H spectrum using default conditions
_	Enter NUTS and	<alt+tab></alt+tab>	Trim phase as required. Use cursor to determine
5	process data.	>a2	TMS peak position in ppm, including sign.
6	Return to PNMR and enter TMS peak position.	<alt+tab> C13>fo<enter> value<enter> 0<enter></enter></enter></enter></alt+tab>	Enter the current position (in ppm) of the TMS peak to the first dialog box and 0 (zero) to the second dialog box. Repeat to confirm.
бb	Optional: Confirm field offset and save ¹ H spectrum for border of HETCOR plot.	C13>zgh <enter> <alt+tab>,>a2 >sa filename</alt+tab></enter>	Acquire proton spectrum. Switch to NUTS and process with a2 link. Enter filename for ¹ H spectrum, for example border_h1. <alt+tab> to return to PNMR</alt+tab>
7	Verify parameters.		Verify that parameters make sense; for neat samples NS=12, for 1M samples NS=60, for concentrations <1M use the BAPR program.
8	Acquire data.	<pre>C13>zg<enter> then filename<enter> or <enter> for default</enter></enter></enter></pre>	Enter file name if desired but it is usually better to use the default (pnmrfid) unless intending to save the data long term.
9	Enter NUTS.	<alt+tab></alt+tab>	
10	Process data.	<pre><ctrl+f3> then [filename][Open] to select a file or [Open] for default</ctrl+f3></pre>	Process using aii_C13.mac that references TMS and sets display range from 220 to -10 ppm. Does an automatic peak pick.
11	Enter line broadening.	value <enter></enter>	LB=0.5 Hz is a typical value.
	Optional: Pick peaks manually.	> dp <enter></enter>	The cursor becomes a crosshair with a DP label. <c> clears all peak picks; <k> removes a single peak pick at the cursor location. Add peaks by aligning cursor on a peak and clicking the left MB. <t> to write peak list to the table.</t></k></c>
12	Plot Data.	>pl	
	Optional: Save ¹³ C spectrum for border of HETCOR plot.	<pre><ctrl+b></ctrl+b></pre>	Remove peak labels and table displays. Enter filename for ¹³ C spectrum, for example borderc13.
	Optional: Add scans	C13>Go <enter></enter>	In case the signal to noise is too low, the acquisition can be extended.
_	Number of added scans	scans <enter></enter>	For neat samples ns-12, for 1M samples ns=60, for concentration <1M use the BAPR program.
		filename <enter></enter>	The program will ask for a filename when finished. Use the same name as step 8.

NOTE: For weak samples use Block Averaging with Peak Registration (BAPR)