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tocsy_metab.nan

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DOI

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We use this protocol and it's working

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Abstract

This is a protocol for running the Bruker pulse program "dipsi2qpphpr" for metabolomics samples.

Guidelines

This protocol intends to provide concise instructions to carry out the experiment. For more comprehensive information, see Bruker's documentation "Basic NMR Experiments" by clicking ? → **Manuals (docs)** on the menu bar on TopSpin. See also "Pulse Program Catalogue. 1D/2D" for the details about the pulse program used in this protocol.

Troubleshooting



Before start

This protocol assumes:

- Your sample is loaded, locked, tuned, and shimmed in the magnet
- The calibrated 90° pulse value for proton (i.e., P1) for the sample has been collected



Create a new dataset

1

1.1 On the menu bar on TopSpin, click on

Start → **Create Dataset**



(This protocol uses TopSpin 3.6.4, and the interface may look different on other TopSpin versions.)

Note

You can also use the **new** command in the command line to do this step.

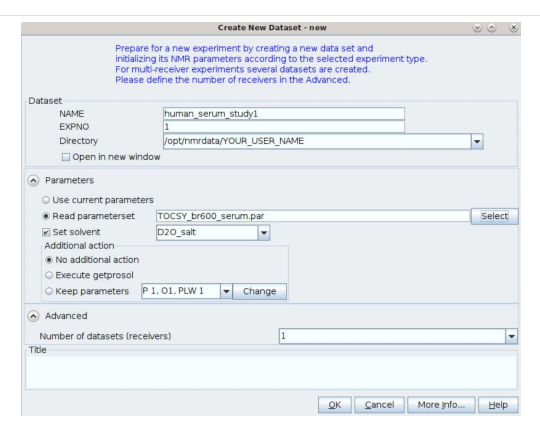
1.2 Enter

- NAME: Name of a set of datasets (e.g., human_serum_study1). Use a single string.
- **EXPNO**: Dataset number. Use a positive integer.

Select

Directory: Your directory.





Note Your new dataset will be stored in Directory/NAME/EXPNO

1.3 Select

Read parameterset

Click the button

Select

1.4 A new window opens. On the right top bar, select

Source = /opt/NAN_METAB/par



<u>File Options Help</u> Source			ource = /opt/NAN_METAB/	= /opt/NAN_METAB/par	
Find file names tocsy*	Exclude:	Clear			
Class = 🕎 Dim = 🕎 Show Recommended					
Type = SubType = SubTypeB =	Reset Filters				
TOCSY_br600_serum.par TOCSY_br600_urine.par TOCSY_NUS_br600_urine TOCSY_NUS_HighRes_br6					
				<u>O</u> K <u>C</u> lose	

In the list, select the one you want to use:

For serum and plasma samples:

■ **TOCSY_br600_serum.par**: Parameter set optimized for serum samples.

For urine samples:

- **TOCSY_br600_urine.par**: Parameter set optimized for urine samples.
- TOCSY_NUS_br600_urine.par: Parameter set using an acquisition mode "nonuniform sampling (NUS)". Higher resolution on the indirect dimension

Note

Parameter set names in the list vary between spectrometers (e.g., TOCSY_br800_serum.par).

Click

OK

1.5 Click

OK

2 Go to the "USE DEFAULT" tab below to proceed with the default optimized parameters.

STEP CASE

Use default parameters: 6 steps

This step case uses the default optimized parameters to acquire a spectrum.

3



3.1 Load the calibrated P1 using the following command in the command line.



getprosol 1H [calibrated P1 value] [power level for P1]

(e.g., getprosol 1H 10.01 -7.45)

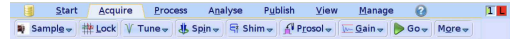
Note

[power level for P1] varies between spectrometers. Never use a wrong [power level for P1].

3.2 Click on

Acquire → **Gain**

in the menu bar to automatically set the receiver gain.



Note

You can also use the **rga** command in the command line.

3.3 Click

in the menu bar to acquire a spectrum.

Note

You can also use the **zg** command in the command line.

3.4 After the run, click on

$\textbf{Process} \rightarrow \textbf{Proc. Spectrum}$

in the menu bar to execute an automated processing macro.





3.5 If you want to modify parameters to improve your spectrum, go to step 2 and move to the step case "MODIFY PAR".