

VERSION 2

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# OPEN ACCESS



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# noesypr1d\_metab.nan V.2 Version 1 is forked from noesypr1d.nan

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### **ABSTRACT**

This is a protocol for running the Bruker pulse program "noesypr1d".

#### **GUIDELINES**

This protocol intends to provide concise instructions to carry out the experiment. For more comprehensive information, see Bruker's documentation including "Basic NMR Experiments" by clicking ? → Manuals (docs) on the menu bar on TopSpin.

**Protocol status:** Working

We use this protocol and it's working

#### **BEFORE START INSTRUCTIONS**

This protocol assumes your sample is loaded, locked, tuned, and shimmed in the magnet.

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# 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

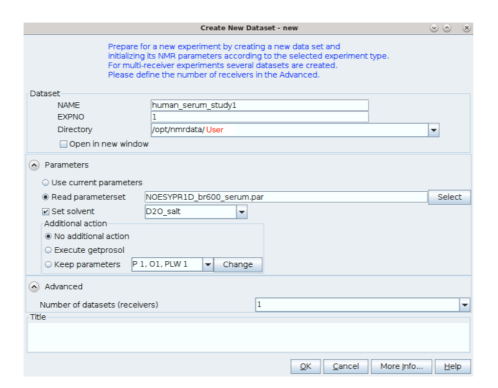
You can also use the **new** command in the command line to do this step.

## 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

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

NOESYPR1D\_br600\_serum.par: Parameter set optimized for serum and plasma samples.

NOESYPR1D\_br600\_urine.par: Parameter set optimized for urine samples.

#### Note

Parameter set names in the list vary between spectrometers (e.g., NOESYPR1D\_br800\_serum.par).

#### Click

### OK



# 1.5 Click

OK

# **Acquire a spectrum**

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

STEP CASE

Use default

7 steps

parameters:

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

3

# 3.1 Click on

Acquire  $\rightarrow$  Prosol  $\rightarrow$  Single scan pulse calibration (pulsecal) to calibrate the 90° proton pulse (i.e., P1).



A new window will appear to show the calibrated P1.

#### Note

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

3.2 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)

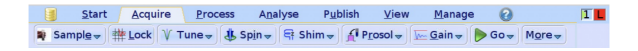
#### Note

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

### 3.3 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.

# 3.4 Click

#### Go

in the menu bar to acquire a spectrum.

### Note

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

# 3.5 Click on

# **Process** → **Proc. Spectrum**

in the menu bar to execute an automated processing macro.



3.6 If you want to modify parameters to improve your spectrum, and move to a step case "MODIFY PAR".