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 C_CON_IPAP.nan

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We use this protocol and it's working

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Disclaimer

This protocol is part of the knowledge base content of NAN: The Network for Advanced NMR (<https://usnan.nmrhub.org/>)

Specific filenames, paths, and parameter values apply to spectrometers in the NMR facility of the Complex Carbohydrate Research Center (CCRC) at the University of Georgia.

Abstract

This protocol describes running a 2D carbon detected CON_IPAP pulse sequence, which provides correlations between CO(i) carbons and N(i-1) amide nitrogens.

It yields two 2D sub-spectra, so use 'splitcomb ipap 2' to process in Topspin.

^{15}N and ^{13}C labeling are required.

It uses the Topspin library pulseprogram 'c_con_iasq'.

Attachments



[biotop.pdf](#)

2.5MB



[NUS_poisson_gap_file...](#)

12KB



Guidelines

NUS sampling is usually not required for 2D experiments, since time savings are small, unless running multiple 2D experiments. If using NUS keep sampling amount ~30-50%.

For processing in TopSpin use 'splitcomb ipap 2' command to extract individual sub-spectra.

Before start

A sample must be inserted in the magnet either locally by the user after training, or by facility staff if running remotely.

This protocol requires a sample that is locked, tuned/matched on 1H, 13C and 15N channels, and shimmed. At a minimum, 1H 90° pulse width and offset O1 should be calibrated and a 1D proton spectrum with water suppression has been collected according to the protocol **PRESAT_bio.nan**. Prior acquisition of a **2D 15N HSQC** and **3D HNCO** are also recommended, according to protocols **HSQC_15N.nan** and **HNCO_3D.nan**.

General aspects of the use of the **bioTop** module are described in more detail in the protocols listed below.

It is recommended to calibrate 1H carrier offset, 1H H2O selective flip-back pulse, as well as 1H, 13C, and 15N 90° pulse widths using the "Optimization" tab of BioTop. Alternatively, 1H 90° pulse width and offset can be calibrated using other methods, such as pulsecal or calibo1p1. Additional parameters, like 15N and 13C offsets and spectral width can be either optimized or manually entered in the "Optimization" tab of BioTop. Note that since BioTop optimizations are saved in the dataset folder, all experiments should be created under the same dataset name when using BioTop for acquisition setup.

Refer to protocols

1. Acquisition Setup Workflow, Solution NMR Structural Biology
2. PRESAT_bio.nan
3. HSQC_15N.nan
4. HNCO_3D.nan
5. Biotop-Calibration and Acquisition setup



Create C_CON_IPAP experiment file

- 1 Start with existing dataset containing 1D proton or 15N-HSQC data for this sample.
- 1.1 Open create dataset window with **edc** command or through menu.
Edit text in Title box.
- 1.2 To select starting parameter set, check 'Read parameterset' box, and click Select.
For standard NAN parameter sets, change the Source directory at upper right corner of the window:
Source = /opt/NAN_SB/par
Click 'Select' to bring up list of parameter sets.
Select C_CON_IPAP_xxx.par, where xxx=900,800 or 600.
Click OK at bottom of window to create the new EXPNO directory.
- 1.3 If not done, tune Nitrogen and Carbon channels:
-on the Acquire menu, click Tune (or type atma on command line).

Calibrate pulses and adjust parameters

2

Note

Loading the C_CON_IPAP_xxx.par parameter set enters the default parameters into the experiment directory. While they represent a good starting point, they may not be fully optimal or accurate for your particular sample or spectrometer hardware. The probe- and solvent-specific parameters, specifically the ^1H 90° pulse length, and possibly the ^{13}C and ^{15}N 90° pulse lengths, along with other dependent pulse widths and powers may need to be updated.

There are two ways of automatically updating experimental parameters:

- 1) Use **getprosol** command, which typically only updates proton pulse widths and power levels. It is most useful for running routine experiments using the default parameters.
- 2) Or use **bioTop** module that organizes calibrated and defined parameters for a dataset.

2.1 Using getprosol:

Use the calibrated proton P1 value obtained from the proton experiment (protocol PRESAT_bio.nan) and note the standard power level attenuation in dB for P1 (PLW1); otherwise type calibo1p1 and wait till finished.

Then execute the getprosol command:

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

e.g. getprosol 1H 9.9 -13.14.

Where for example, the calibrated P1=9.9 at power level -13.14 dB attenuation

This also loads 15N and 13C pulse widths and power levels from the 'edprosol' table, and are assumed to be sufficiently accurate.

Skip to step 2.3.**2.2 Use bioTop**

If you previously performed parameter calibrations using the "Optimizations" tab of the BioTop GUI, or entered parameters manually in the "Optimizations" tab, you can **type btprep** at the command line.

See protocol 'BioTop : calibration and acquisition parameters' and attached Bruker manual 'biotop.pdf' for details.

Inspect Parameters

- 2.3** Examine parameters by typing '**eda**', or select the 'Acqpars' tab and get the 'eda' window by clicking on the '**A**' icon. This view shows the two dimensions, F2 (13C) and F1 (15N) in columns.

Parameters to check:

2 TD: Number of ¹³CO time domain real points (~512-2048, preferably 2^N, keep **2 AQ** at ~50-200 ms)

1 TD: Number of 15N time domain real points, 1 AQ at ~30-60 ms (TopSpin 4.x) or ~60-120 ms (TopSpin 3.x). In TopSpin 3.x inner IPAP loop counts toward the TD points.

2 SW: ¹³CO spectral width (~20 ppm, defined in **BioTop**)

1 SW: 15N amide spectral width (~46 ppm, defined in BioTop)

O1P: ¹³CO offset (~173 ppm, defined in **BioTop**)

O2P: ¹H offset (~4.7 ppm, defined in **BioTop**)

O3P: 15N amide offset (~115-120 ppm, defined in bioTop)

- 2.4** Then examine the parameters in Acqpars 'ased' mode (click 'pulse' icon), or **type 'ased'**;

Most of the default parameters are suitable, however it's useful to compare values in the fields against those proposed by the original parameter file and the pulseprogram.

D1: recycle delay (~1 s for protonated samples, ~2-3 s for perdeuterated samples)



DS: 32-128 'dummy' scans that are not recorded ; allows system to reach steady state equilibration.

NS: multiple of 4; increase for increased signal to noise (S/N increases as $\sqrt{NS}$)

channel f1:

CNST21 - ^{13}C O offset (~173 ppm, same as **O1P**, defined in **BioTop**)

CNST22 - ^{13}C A offset (~48 ppm, defined in **BioTop**)

channel f2:

CPDPRG2 - ^1H decoupling (garp)

channel f3:

P21 - 15N 90° high power pulse (calibrated with BioTop)

CPDPRG3 - ^{15}N decoupling (garp, garp4)

Acquire and Process Data

- 3 Type '**rga**' or click on 'Gain' in Topspin Acquire menu to execute automatic gain adjustment.
Type '**expt**' to calculate the expected run time
Type '**zg**' or click on 'Run' in Topspin Acquire menu to begin acquisition.
- 3.1 You can always check the first FID by typing 'efp' to execute an exponential multiplied Fourier transform. It will ask for a FID #, choose the default #1. You can evaluate the 1D spectrum for amide proton signal to noise and water suppression.
- 3.2 2D planes with NUS or uniformly sampled can be processed with Topspin: **xfb**

Protocol references

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'nusPGSv8' written by Scott Anthony Robson 2013