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1 Introduction

The goal of this manual is to describe the basic steps of working with the TopSpin software package to process NMR data on a data station. This should enable a relatively unexperienced user to install, start, display, process and analyze acquired data independently.

1.1 How to Use this Manual

Please note: The TopSpin icons provide access to many different graphically oriented functionalities. This functionality is also available through commands that one can type into the TopSpin command line or use in automation routines. When the mouse cursor is moved over an icon, further explanatory information about the function of the icon is shown as well as the corresponding command name. For better reading of our manuals, we therefore don’t explicitly list or mention these command names in the description of the icons.

2 Installation

In the following chapter, the software installation will be described only for data station. In case of an NMR Spectrometer, used for NMR data acquisition, please refer to the Installation Guide Manual delivered with TopSpin (under Help | Manuals (docs) | Installation Guides | TopSpin Installation Guide) or download it from the Bruker website under the following link: https://www.bruker.com/service/information-communication/user-manuals/nmr/installation.html (Please note that access to the download section is restricted to customers. Registration for a free customer account is provided directly on the Bruker website.).

There are two main ways to install the TopSpin Software on your computer:

Download from the internet:

If you have access to an internet connection the TopSpin up to date version can be directly downloaded from the Bruker website: https://www.bruker.com/service/support-upgrades/software-downloads.html

If you don’t have an account, please register for an account. If you already have an account please log in to access the download section containing all listed software products.

Using the DVD

Insert the TopSpin DVD and open the DVD drive with your usual system explorer.

Find and double click the install command (Windows: install.cmd, Linux: install, Mac OS: TopSpin<version>.pkg) and follow the instructions of the Installer.

The Installation Procedure of TopSpin on all operating systems is mainly the same. Nevertheless, there are some small differences that are explained in detail in the Installation Manual delivered with TopSpin (see above).

Please note: For all operating systems the administrator password is required when installing TopSpin.
3 First Start and License

Start TopSpin by double clicking the TopSpin desktop Icon ( ). If you already have a TopSpin license, TopSpin will open immediately. If not, a dialog will be opened which guides you through the process of obtaining and installing a TopSpin license (see figure below).

For further information on licenses, please refer to the CodeMeter License Management Manual on the Bruker website: https://www.bruker.com/fileadmin/user_upload/8-PDF-Docs/MagneticResonance/Service_NMR/Installation-Guides/software_license_agreement.pdf
4 Getting Started

Once the TopSpin license has been activated, TopSpin will start immediately:

4.1 TopSpin Interface - Overview

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4.2 First Start of TopSpin - Load a Dataset

Figure 4.1: TopSpin starting window.

At the starting point of TopSpin no dataset is selected. You first have to select it in the browser (see figure below). Load it in the active window by drag and drop.
Figure 4.2: Browser window with list of available datasets (spectra+parameters)

Click on the plus sign left of the dataset name you want to select. The figure below shows this for a 1D proton experiment (exam1d_1H). The type of experiment (here zg30) and the name of the samples (cyclosporin and cholesterylacetat) will be shown. You can navigate in this browser window and select the dataset you want to inspect.

You can also use the Search functionality to get a list of datasets containing the search text.

Figure 4.3: Browser & search window.

You can also load a dataset by drag & drop from an Explorer window (e.g. Windows Explorer, Konqueror, Finder etc.).

The figure below shows the TopSpin window with the selected dataset after drag & drop.
4.3 Basic Processing and Manipulations

The workflow button bar opens the spectrum processing tools.

![Workflow button bar for spectrum processing.](image)

The toolbar gives access to the most useful tools to manipulate the spectrum.

Please note: ![Offers access to further icons for spectrum manipulation functionalities.](image)
4.4 Basic Processing

Clicking on the Proc. Spectrum button in the workflow button bar (see Figure 4.5) performs all basic processing steps fully automatically. The steps can be configured through the proc1d button in the pulldown menu.

The screenshot below shows the configuration options for the basic 1D automatic processing.
4.5 Phase Correction

For further improvement of the phase of the spectrum, \( \uparrow \downarrow \text{Adjust Phase} \) gives access to the interactive phase correction. The dataset window will show icons specific for phasing.

In addition, the Adjust Phase pulldown menu gives access to further phasing operations.
4.6 Baseline Correction

For further improvement of the spectrum baseline, leads to the interactive baseline correction. The dataset window will show icons specific for baseline correction.

In addition, the Baseline pulldown menu gives access to further baseline correction operations.

4.7 Axis Calibration

gives access to following pulldown menu with axis calibration options:

4.8 Advanced Processing Options

gives access to further options through following pulldown menu:
5 Analyse the Spectrum

To enter the Analyse Process, click on the Analyse button, the following workflow appears:

![Workflow button bar for spectrum analysis.](image)

*Figure 5.1: Workflow button bar for spectrum analysis.*

5.1 Peak Picking

...gives access to the interactive peak picking. The dataset window shows specific icons for peak picking:

![Icons for peak picking.](image)

In addition, the Pick Peaks pulldown menu gives access to further peak picking operations and options:
5.2 Spectrum Integration

The **integrate** button gives access to the integration pulldown menu:

![Integration menu](image)

*Figure 5.3: Integration pulldown menu and Integrate Regions File menu (command `intrng`).*

In addition, more options can be obtained through the command `int`: 

*Figure 5.2: Peak picking pulldown menu and options (command `pp`).*
5.3 Further Analysis Options

Further analysis options can be obtained through the corresponding workflow:

- Define integral regions manually
- Auto-find regions, integrate & display result
- Integrate existing regions (file ‘intmg’) & display result
- List peaks and integrals (using regions file ‘intmg’) within the displayed region
- List peaks and integrals (using regions file ‘intmg’) of the entire spectrum
- Integrate a list of spectra

Required parameters

- Integration sensitivity factor ABSL (0..100) = 3
- Minimum separation between independent integral regions AZFW [ppm] = 0.1
- Integral region extension factor AZFE [ppm] = 0.1
- Integral sensitivity factor with reference to the largest integral ISEN (>0) = 128
- Degree of polynomial ABSG (0..5) = 5
- Left spectral range limit F1P [ppm] = 9.99032020668847
- Right spectral range limit F2P [ppm] = -0.99583438312942
- Scale 1D integrals relative to a reference dataset INTSCL (-1, 0, >0) = 1
- Automatic baseline correction of integrals (if regions auto-detected!) INTBC = no

Figure 5.4: Further options for integration (command int).

Please note: For operations specific tutorials are available via the pull-down menu.
6 Printing the Spectrum

Printing the spectrum can be done in two different ways:

6.1 Print Screen Directly

Click on the printer icon , located on the upper right part of the TopSpin window, allows direct printing of the spectrum. Details can be obtained as usual by dragging the mouse on the icon. Clicking the button gives access to the print window:

![Figure 6.1: Print window.](image-url)
6.2 Print Spectrum Using Layout (Plot)

To use this TopSpin feature, click the button \[ \text{PLOT} \], or use the \text{plot} command in the TopSpin command line. The following TopSpin window appears:

![TopSpin plot window](image)

Figure 6.2: Part of TopSpin plot window (with layout).

The left part of the plot window shows the names of the selected layout and printer. They can be changed with the arrow button, as well as other plotting view parameters, like limits or display. New NMR elements (title, parameters, 2D spectrum, etc.) can be inserted.

The right part presents the spectrum like it will be plotted. Changes on the left part will be shown in real time in the layout on the right.

7 Further Documentation

For detailed information please refer to the TopSpin Manual section under Manuals (docs) or visit the Bruker website under https://www.bruker.com/service/information-communication/user-manuals/nmr.html. (Please note that a customer account is needed.)
8  Contact

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Bruker BioSpin Hotlines
Contact our Bruker BioSpin service centers.
Bruker BioSpin provides dedicated hotlines and service centers, so that our specialists can respond as quickly as possible to all your service requests, applications questions, software or technical needs.
Please select the service center or hotline you wish to contact from our list available at:
https://www.bruker.com/service/information-communication/helpdesk.html