

# Quick Start Guide

# ACD/1D NMR Processor: Basic Training

Version 12

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# Introduction

The following document outlines how to utilize ACD/1D NMR Processor for processing, assigning, and reporting a <sup>11</sup>

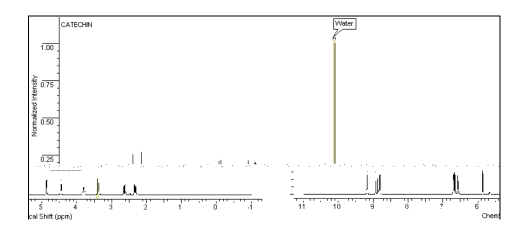


# **Fast Fourier Transform, Baseline, and Phase Corrections**

Shortcut Zero Filling FID Shift LinearPred WFunctions Fourier Tr. | Apodization | Manual Offset

### To automatically Fourier transform, baseline correct, and phase correct

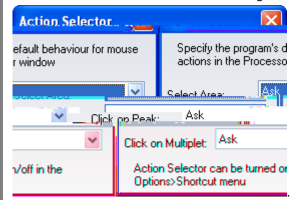
On the Operations toolbar, click Shortcut





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Note Ensure the Action Selector dialog box settings are as follows:



Peak Picking, Integration, and Multiplet Creation



# Attaching a Chemical Structure to a Spectrum

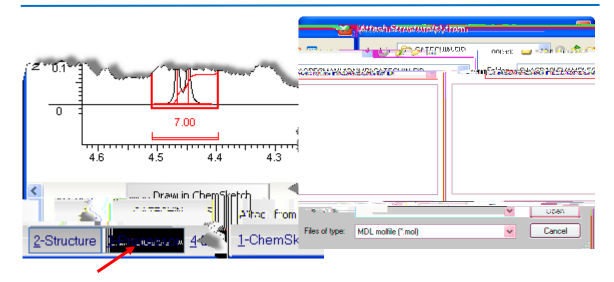
Once attached to a spectrum, the structure information is imbedded directly in the ACD/Labs spectral data format making it less likely that the identity information will be misplaced or lost.

#### To attach a chemical structure to a spectrum

- 1. On the bottom Switching bar, rest the cursor on 2-Structure to show the button menu.
- From the button menu, choose **Draw in ChemSketch** to use the built-in structure drawing package.

-OR-

From the button menu, choose **Attach from File** to open the Attach Structure(s) from dialog box.



Chemical structures associated with a spectrum can be automatically included in publication quality reports. The attached chemical structure is used for facilitating spectral assignments and verification.

#### To clear a chemical structure from a spectrum

• On the **Edit** menu, point to **Clear**, and then click **Structure**.

#### Practice Task:

Attach the structure of catechin to the spectrum.

**Hint!** Instead of drawing the chemical structure for catechin and other well known compounds by hand, check the ACD/Dictionary first. Search for compounds by name by clicking the **Dictionary** button on the right-side vertical toolbar. Found structures can be pasted directly in the workspace.

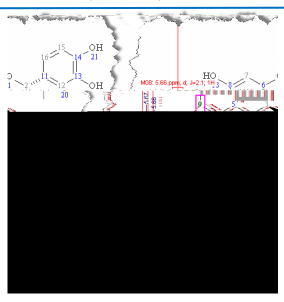


# **Peak Assignment**

#### To assign peaks to corresponding proton (or carbon) in the structure

- 1. On the Operation toolbar, click Assignment to enter Assignment mode.
- 2. On the Assignment toolbar, click By Multiplet
- 3. Click an atom in the structure, and then click a multiplet in the spectrum to create the assignment.
  - -OR-

Click a multiplet in the spectrum, and then click the atom to assign it.



If you select the atom first, a red line will go from the cursor to this atom. As well, a vertical bar will indicate your position on the spectrum and the nearest multiplet near this point will get highlighted. Similar functionality exists in assigning from multiplet to atom.

Note To abort an assignment, click an empty background area.

Practice Task:

Assign a multiplet in the spectrum to an atom.



# **Preparing a Report**

#### To preview a report in the ChemSketch editor

• On the **Edit** menu of ACD/1D NMR Processor, point to **Create Report**, and click **Standard**.

From ChemSketch you can print the report as shown, save it in ChemSketch format, or produce a PDF version.

Note Larger reports automatically create a multipage report. On the bottom bar, use the controls to navigate between the pages of the report. .

To paste segments of a ChemSketch report to





# Conclusion

The process described above is a very basic overview of the main workflow in ACD/1D NMR Processor. Many more processing options exist that may be valuable in your process.

More in-depth instructions are available in the *ACD/1D NMR Processor Reference Manual*. The User's Guide can be accessed in Adobe PDF format from the Processor window (Help>Documents>1D NMR>Reference Manual).

A detailed User's Guide for the ACD/ChemSketch structure editor is also available. To access the ChemSketch user guide, you must be in the ChemSketch window (Help>Documents>Guides).

Whenever prompted, you are encouraged to watch the technical movies that are included in the software. These movies cover important material that is not included in this guide.



# 1D NMR Processor Quick Start Summary Sheet

# **Importing Raw Data**

# To import spectral data

On the main toolbar, click





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Preparing a Report