KnowItAll® Informatics Training

NMR Processing Tools

Process NMR

How to Import and Process 1D NMR Files

Purpose

This exercise demonstrates how to use the KnowltAll Informatics System's ProcessIt NMR application to import and process 1D NMR files.

Objectives

This exercise will teach you:

- > How to use ProcessIt NMR tools to improve spectral appearance and correct experimental artifacts
- How to create and use processing macros

Background

You can use the ProcessIt application to import raw data files from major NMR instrument vendors and processed data formats. The application can then be used to process the files and is useful for improving spectral appearance and correcting experimental artifacts.

Training Files Used in This Lesson

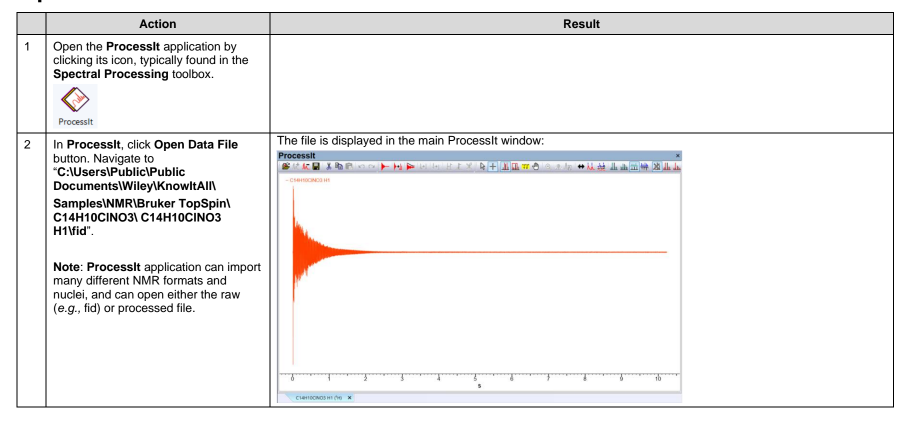
C:\Users\Public\Public
 Documents\Wiley\KnowItAll\
 Samples\NMR\Bruker TopSpin\
 C14H10CINO3\ C14H10CINO3 H1\fid

KnowltAll Applications Used

ProcessIt



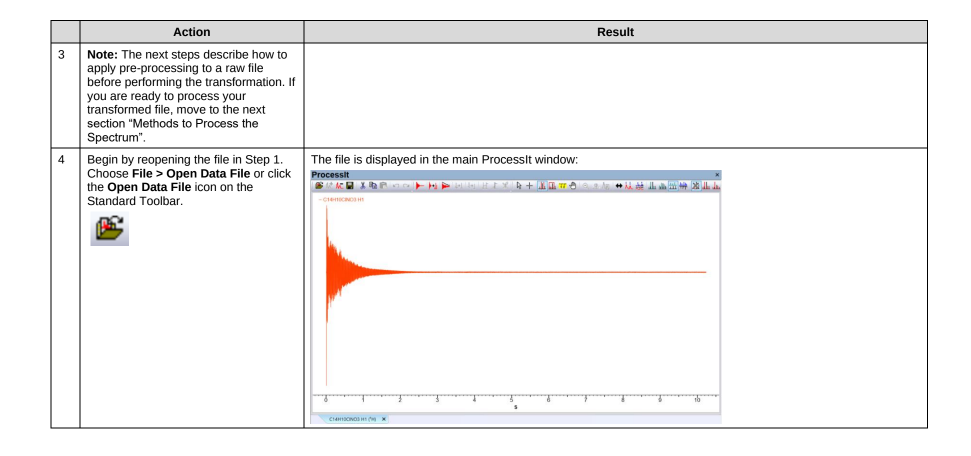
Open a raw NMR file in ProcessIt



Methods to Transform the File

Action Result Fourier Transform: If you wish to The Fourier Transform dialog appears with a preview of the transformed proton spectrum behind it: apply a Fourier Transform (FT) without additional preprocessing, begin here (otherwise move to step 4). Choose Process > Transforms > Fourier Transform Fourier Transform, or click the Fourier Transform icon on the Default O Complex Process Toolbar. Close Replace C14H10CINO3 H1 (*H) X Click **Replace** on the **Fourier** The transformed proton spectrum is displayed: Transform dialog to accept the previewed transformation using the ● はんに回 ※ 鳴 戸 め ○ □ **Default** algorithm. **Note:** Four different algorithms are available in the Fourier Transform dialog: Default, Complex, Real and Bruker. 3e+08-C14H10CINO3 H1 (¹H) X





The Zero Filling dialog box opens:

Choose Process > Zero Fill.

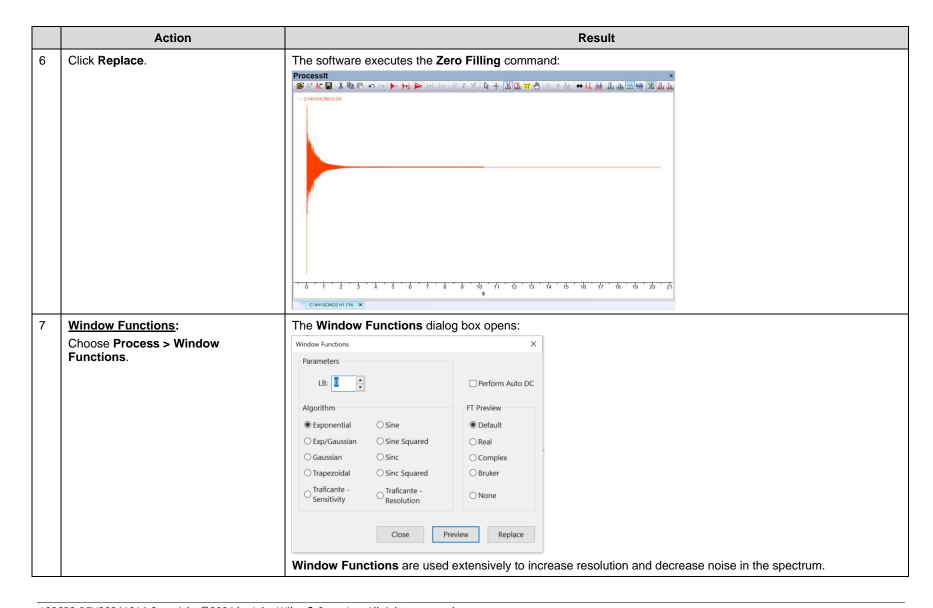
The Zero Filling dialog box opens:

Zero Filling

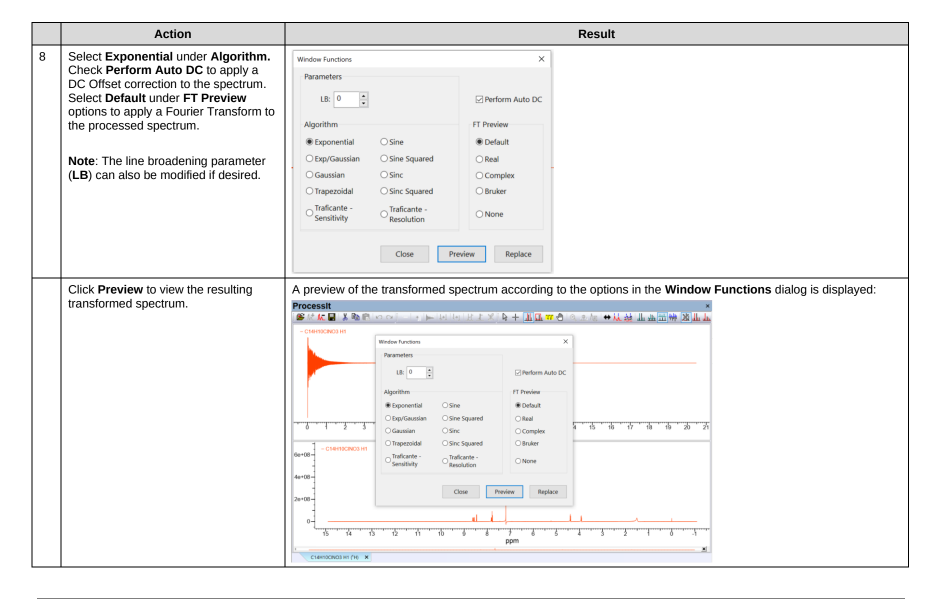
Original Point Count: (Real + Imaginary) 131072

Proposed Point Count: (Real + Imaginary) 262144

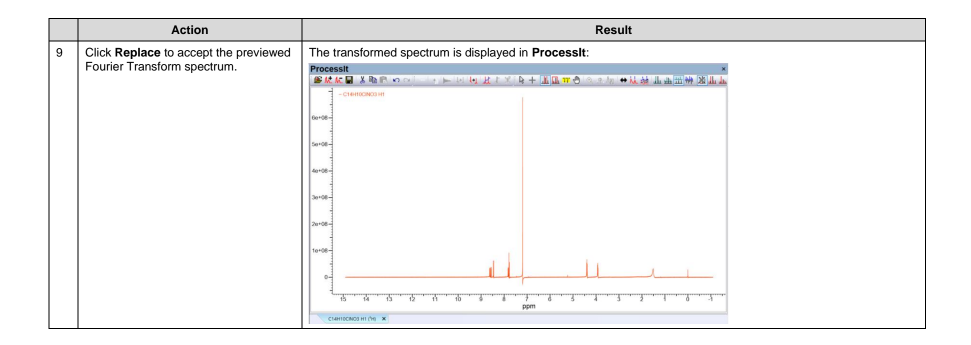
The software proposes a new number of points for the spectrum based on the original (experimental) count.



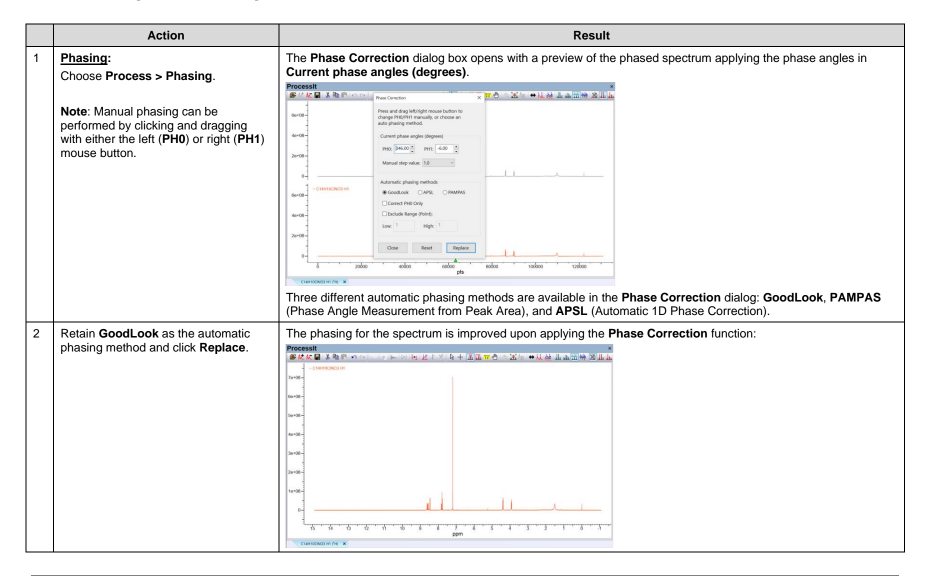




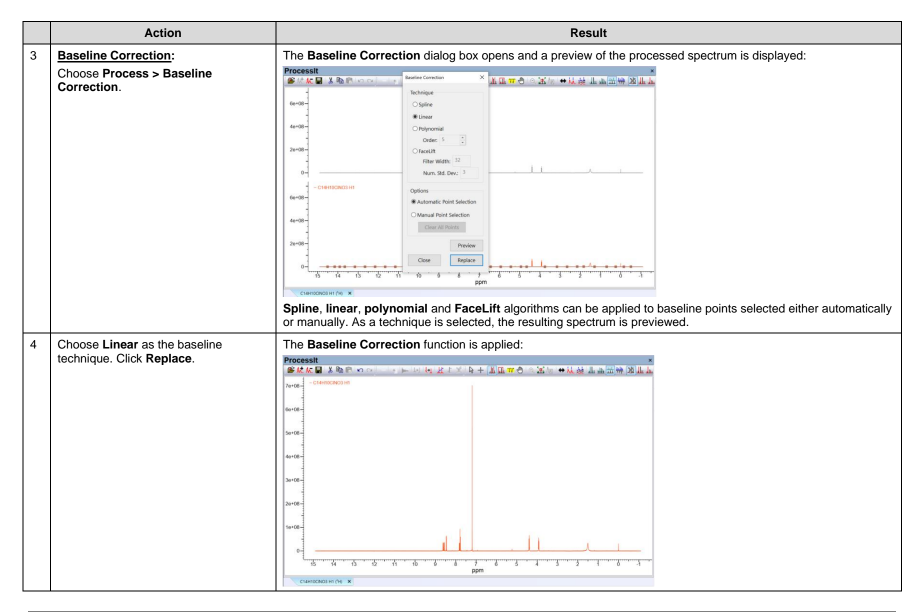




Methods to process the spectrum

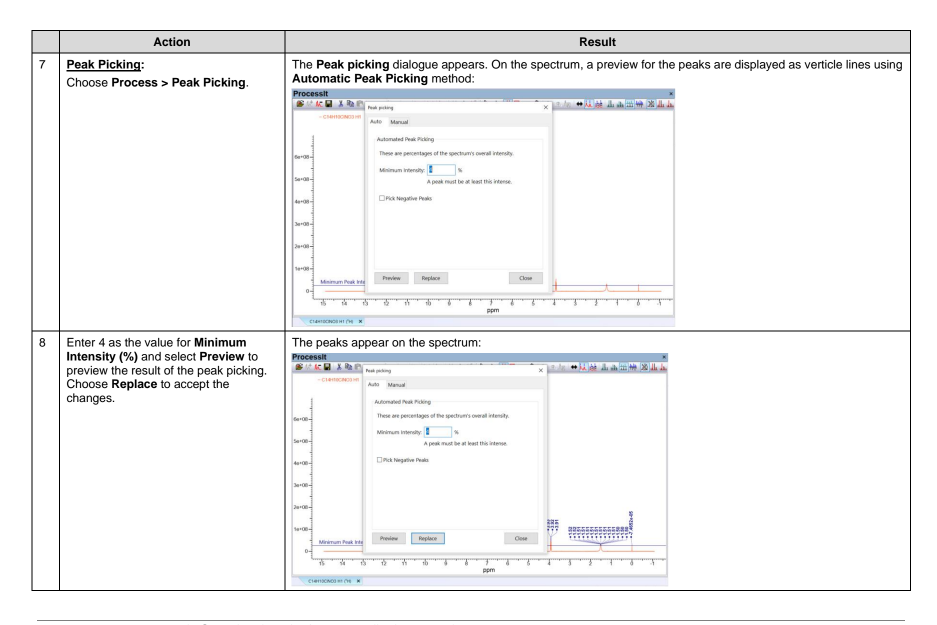




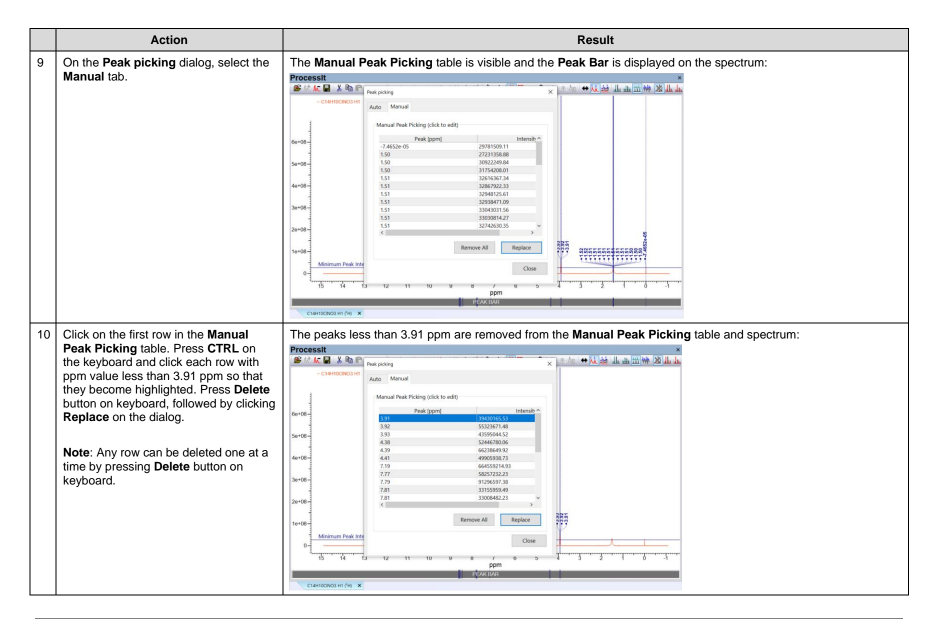








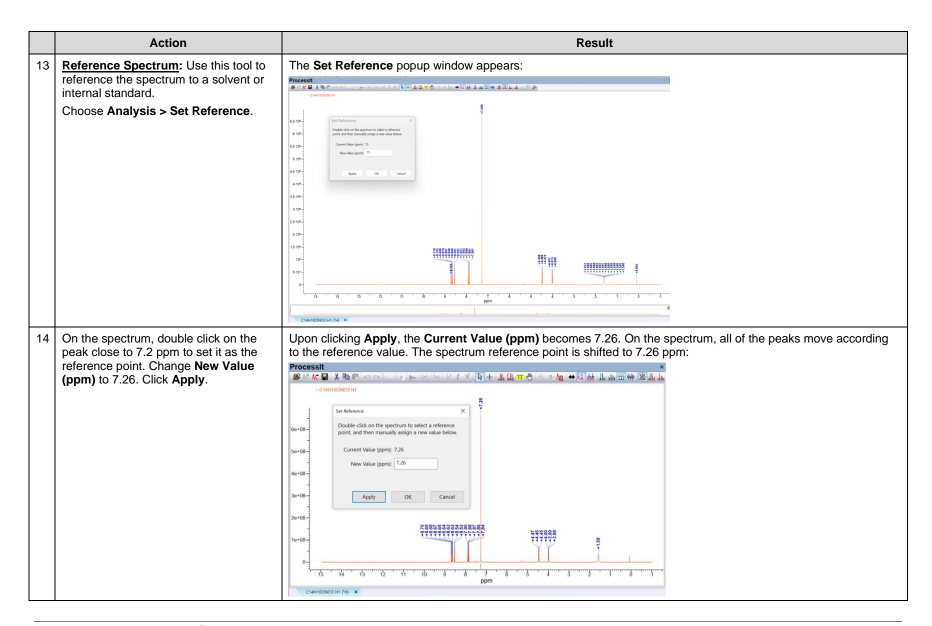




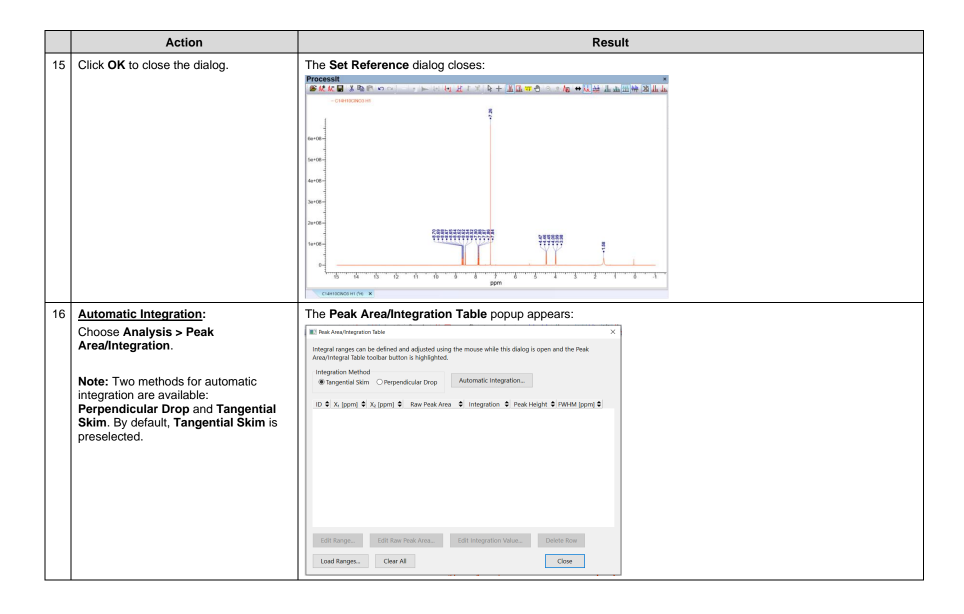




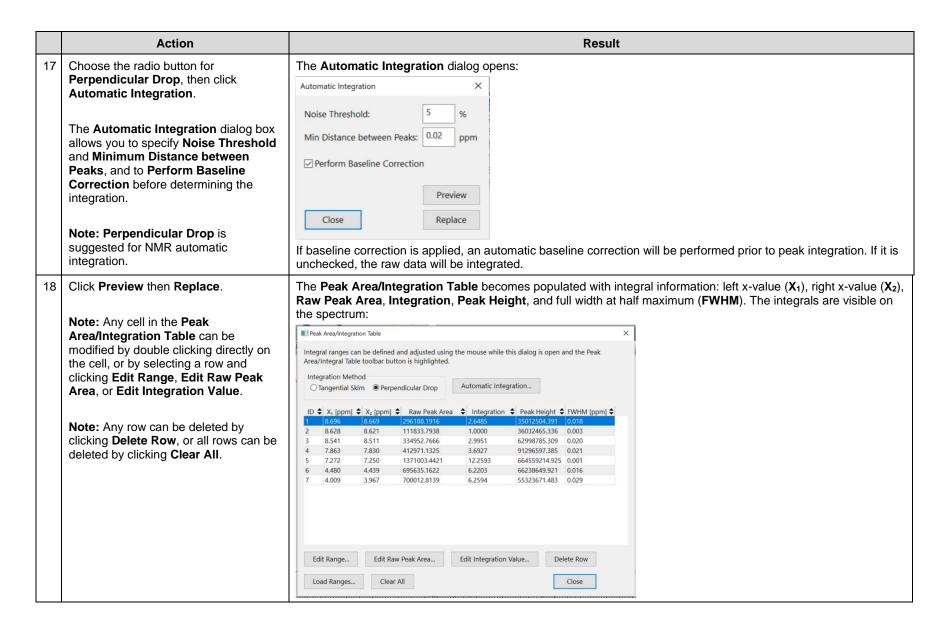




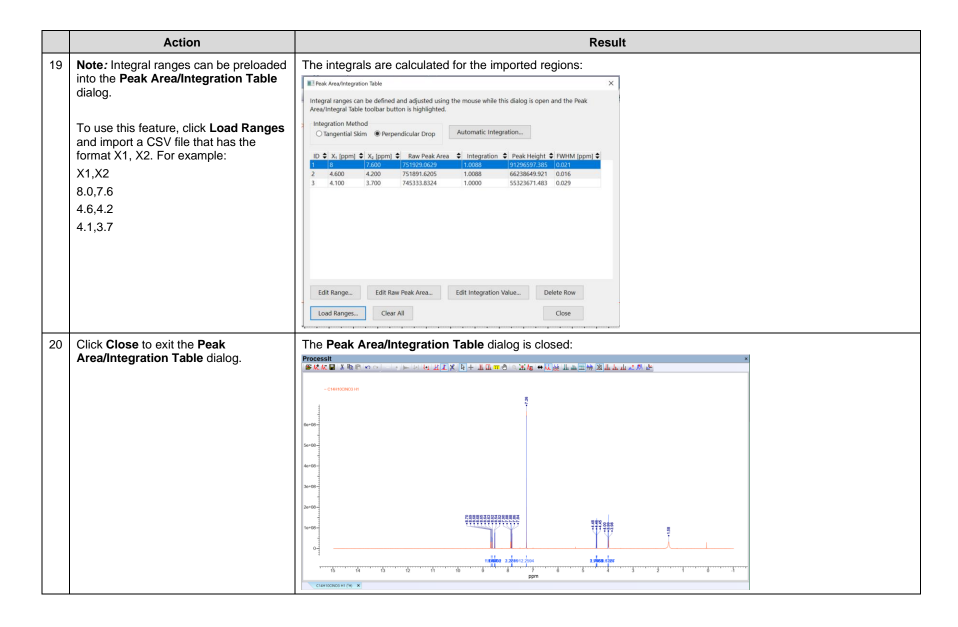


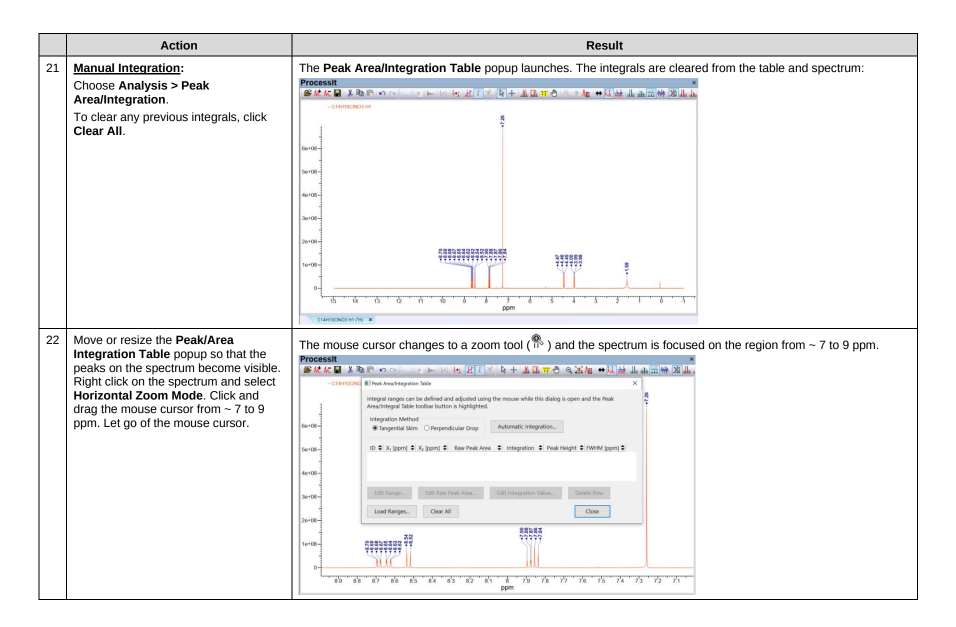








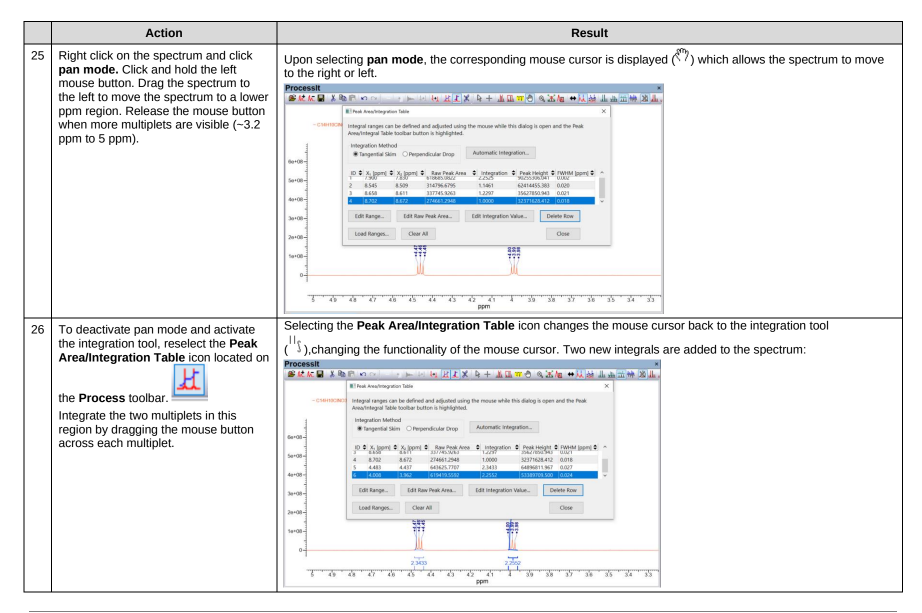






	Action	Result	
23	Select the Peak Area/Integration Table icon located on the Process toolbar to reactivate the integration tool.	Clicking the Peak Area/Integration Table icon changes the mouse cursor from the zoom tool () to the integration tool ():	
24	Click and drag the mouse button across a multiplet. Release the mouse button at the end of the multiplet. Repeat for each multiplet in visible in the displayed region. The multiplets are: (1) 8.67 – 8.70 ppm, (2) 8.62 – 8.65 ppm, (3) 8.51 – 8.54, (4) 7.83 – 7.91 ppm. Note: Dragging functionality is bidirectional (left to right, or right to left). Note: Do not integrate the peak at 7.26 ppm which represents the ¹ H-NMR solvent signal.	The integration value for each multiplet is added as a new row in the Peak Area/Integration Table. The integral also display on the spectrum. Processit Peak Area/Integration Table Peak Area/Integration Table	







	Action	Result	
27	Note: On the spectrum, any integral can be resized. To resize an integral, click on the edge of the integral region and drag its edge to left or right along the x-axis.	When the mouse cursor is positioned on the edge of the incursor () to display an arrow pointing right () or left (dragging the integral egg to the right:). The example integral below becomes wider upon
		Before dragging right integral edge:	After dragging integral edge to the right:
28	Note: On the spectrum, any integral can be shifted. To relocate an integral, click inside of the integral region and drag it to the left or right on the x-axis.	the integral center to the right: Before dragging from integral center:	egion, the cursor changes from the integral cursor () to example integral below moves by clicking and dragging After dragging to the right from integral center:
29	Click Close on the Peak Area/Integration Table when the integration is complete. To zoom back out, right click and click View Entire Spectrum.	The entire spectrum is displayed with integrals: Processit CHENDONOMIN CONTINUE OF THE PROPERTY OF THE PROP	8.54 8.53 8.52 8.51 8.5 X



Apply a macro

