

# Analysis V3<sub>beta2</sub> shortcuts

## Mouse events

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Context menu	Right click
Pan	Left drag
Zoom into area	Shift right drag
Zoom into area	Middle drag
Zoom in/out around cursor	Scroll wheel in display
Zoom in/out for selected axis	Scroll wheel in axis active area
Select new	Left click
Add/Remove one to/from Selection	Ctrl/Cmd left click
Select in area and add	Ctrl/Cmd left drag
Pick one peak at position	Ctrl/Cmd shift left click
Pick peaks in area	Ctrl/Cmd shift left drag
Move single peak	Ctrl/Cmd middle drag
Move module	Drag&drop purple bar
Move whole window	Alt Left drag
Pop out module (manage as windows)	Double click on Purple bar
Make new object	Double click <New> in Sidebar
View/Edit object	Double click object in Sidebar

## Keyboard shortcuts

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### *Graphics :*

	Shortcut
Delete selected	Del (Fn+Backspace)
Toggle Grid	GS
Toggle Crosshairs	CH
Toggle Horizontal Trace	TH
Toggle Vertical Trace	TV
Increase Trace Scale	TU
Decrease Trace Scale	TD
Create Mark at cursor position	MK
Clear All Marks	MC
Navigate to Peak position	FP
Navigate to NmrResidue position	FN
Copy PeakList	CL
Copy Peaks	CP
Clear Current Selected Peaks	CZ
Refit Selected Peaks	RP
Snap to Extremum Current Selected Peaks	SE
Select All Peaks	Ctrl/Cmd+A
Toggle Phasing Pivot	PV
Add Integral in 1D plot	I1
Add Phasing Trace	PT

**Remove Phasing Trace**  
**Switch Mouse Mode**

**TR**  
**MM**

*Project :*

**New project**  
**Open project**  
**Load Data**  
**Save**  
**Save as**  
**Import Nef File**  
**Export Nef File**  
**Undo**  
**Redo**  
**Quit**

**Ctrl/Cmd+N**  
**Ctrl/Cmd+O**  
**LD**  
**Ctrl/Cmd+S**  
**SA**  
**IN**  
**EX**  
**Ctrl/Cmd+Z**  
**Ctrl/Cmd+Y**  
**Ctrl/Cmd+Q**

*Spectrum :*

**Load Spectrum**  
**Show Spectrum Groups**  
**Set Experiment Types**  
**Pick 1D Peaks**  
**Pick ND Peaks**  
**Copy PeakList**  
**Copy Peaks**  
***Make Projection***  
**Print to File**

**LS**  
**SS**  
**ET**  
**P1**  
**PP**  
**CL**  
**CP**  
**PJ**  
**PR**

*Molecules :*

**Reference Chemical Shifts**

**RC**

*Assign :*

**Set up NmrResidues**  
**Pick and assign**  
**Backbone assignment**  
**Peak Assigner**  
**Assignment Inspector**  
**Residue information**

**SN**  
**PA**  
**BB**  
**AA**  
**AI**  
**RI**

*View :*

**New Blank Display**  
**NmrAtom Chemical Shift Table**  
**NmrResidue Table**  
**Peak Table**  
**Integral Table**  
**Restraint Table**  
**Structure Table**  
**Sequence Graph**  
**Atom Selector**

**ND**  
**CT**  
**NT**  
**LT**  
**IT**  
**RT**  
**ST**  
**SG**  
**AS**

<b>Chemical Shift Mapping</b>	<b>CM</b>
<b>Show Sequence</b>	<b>SQ</b>
<b>Notes Table</b>	<b>NO</b>
<b>Python Console</b>	<b>PY</b>
<i>For current strip</i>	
<b>Show/Hide Toolbar</b>	<b>TB</b>
<b>Show/Hide Phasing Console</b>	<b>PC</b>
<b>Move to Next Spectrum (and Hide Current)</b>	<b>MN</b>
<b>Move to Previous Spectrum (and Hide Current)</b>	<b>MP</b>
<b>Reset Zoom</b>	<b>RZ</b>
<b>Flip X-Y Axis</b>	<b>XY</b>
<b>Flip X-Z Axis</b>	<b>XZ</b>
<b>Flip Y-Z Axis</b>	<b>YZ</b>
 <i>Macro :</i>	
<b>Define User Shortcuts</b>	<b>DU</b>
<b>Reserved for user macros</b>	<b>Q0, Q1, ...Q9</b>
<b>Reserved for user macros</b>	<b>W0, W1, ...W9</b>
<b>Reserved for user macros</b>	<b>Y0, Y1, ...Y9</b>