Analysis V3beta2 shortcuts

Mouse events

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

Drag&drop purple bar

Move whole window Alt Left drag

Pop out module (manage as windows)

Double click on Purple bar

Make new object

View/Edit object

Double click <New> in Sidebar

Double click object in Sidebar

Keyboard shortcuts

Shortcut

Graphics:

Delete selected Del (Fn+Backspace)

Toggle Grid GS **Toggle Crosshairs** CH TH **Toggle Horizontal Trace** TV **Toggle Vertical Trace** TU **Increase Trace Scale** TD **Decrease Trace Scale** MK Create Mark at cursor position MC **Clear All Marks** FP Navigate to Peak position Navigate to NmrResidue position FN CL Copy PeakList **Copy Peaks** CP

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	TR
Switch Mouse Mode	MM
Zoom in to Spectrum	ZI
Zoom out of Spectrum	ZO
Toggle Share Y Axis Across Display	TA

Project:

New project Ctrl/Cmd+N
Open project Ctrl/Cmd+O

Load Data LD

Save Ctrl/Cmd+S

Save as SA
Import Nef File IN
Export Nef File EX

Undo Ctrl/Cmd+Z
Redo Ctrl/Cmd+Y
Quit Ctrl/Cmd+Q

Spectrum:

LS **Load Spectrum** SS **Show Spectrum Groups** ET **Set Experiment Types P1** Pick 1D Peaks PP **Pick ND Peaks** CL **Copy PeakList** CP **Copy Peaks** Make Projection ΡJ PR **Print to File**

Molecules:

Reference Chemical Shifts RC

Assign:

Set up NmrResidues

Pick and assign

PA

Backbone assignment

Peak Assigner

AA

Assignment Inspector

Residue information

SN

PA

AB

BB

RI

View:

New Blank Display ND
NmrAtom Chemical Shift Table CT
NmrResidue Table NT

Peak Table	LT
Integral Table	IT
Restraint Table	RT
Structure Table	ST
Sequence Graph	SG
Atom Selector	AS
Chemical Shift Mapping	CM
Show Sequence	SQ
Notes Table	NO
Python Console	PY
For current strip	
Show/Hide Toolbar	TB
Show/Hide Phasing Console	PC
Move to Next Spectrum (and Hide Current)	MN
Move to Previous Spectrum (and Hide Current)	MP
Reset Zoom	RΖ
Flip X-Y Axis	XY
Flip X-Z Axis	ΧZ
Flip Y-Z Axis	ΥZ

Macro :

Define User Shortcuts	DU
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Reserved for user macros Q0, Q1, ...Q9
Reserved for user macros W0, W1, ...W9
Reserved for user macros Y0, Y1, ...Y9