Analysis V3 shortcuts

Mouse events

Context menu

Pan

Zoom into area

Zoom into area

Zoom in/out around cursor

Zoom in/out for selected axis

Select new

Add/Remove one to/from Selection

Select in area and add

Pick one peak at position

Pick peaks in area Move single peak

Move module

Move whole window

Pop out module (manage as windows)

Make new object View/Edit object Right click

Left drag

Shift right drag

Middle drag

Scroll wheel in display

Scroll wheel in axis active area

Left click

Ctrl/Cmd left click

Ctrl/Cmd left drag

Ctrl/Cmd shift left click

Ctrl/Cmd shift left drag

Ctrl/Cmd middle drag

Drag&drop purple bar

Alt Left drag

Double click on Purple bar

Sidebar Sidebar

Keyboard shortcuts

	Shortcut	Related
Graphics:		
Delete selected	Del (Fn+Backspace)	
Toggle Grid	GS	CH, TA, PC
Toggle Crosshairs	СН	GS, TA, PC
Toggle Horizontal Trace	TH	TV, TU, TD
Toggle Vertical Trace	TV	TH, TU, TD
Increase Trace Scale	TU	TH, TV, TD
Decrease Trace Scale	TD	TH, TV, TU
Create Mark at cursor position	MK	MC
Clear All Marks	MC	MK
Navigate to Peak position	FP	FN
Navigate to NmrResidue position	FN	FP
Copy PeakList	CL	СР
Copy Peaks	СР	CL
Clear Current Selected Peaks	CZ	
Refit Selected Peaks	RP	SE
Snap to Extremum Current Selected Peaks	SE	RP
Select All Peaks	Ctrl/Cmd+A	
Set Phasing Pivot to cursor position	PV	PC, PT, TR
Add Integral in 1D plot	I1	
Add Phasing Trace	PT	PC, PV, TR
Remove Phasing Trace	TR	PC, PV, PT
Switch Mouse Mode	MM	
Zoom in to Spectrum	ZI or +	ZO, ZS, ZR

Zoom out of Spectrum	ZO or –	ZI, ZS, ZR
Toggle Share Y Axis Across Display	TA	GS, CH, PC
Cycle Peak Labelling	PL	PS
Cycle Peak Symbols	PS	PL
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	
Preferences	Ctrl/Cmd+,	
Quit	Ctrl/Cmd+Q	
Spectrum:		
Load Spectrum	LS	SS
Show Spectrum Groups	SS	LS
Set Experiment Types	ET	
Pick 1D Peaks	P1	PP
Pick ND Peaks	PP	P1
Copy PeakList	CL	CP
Copy Peaks	СР	CL
Make Projection	PJ	
Print to File	Ctrl/Cmd+P	
Molecules :		
Reference Chemical Shifts	RC	
Acces		
Assign:	CNI	
Set up NmrResidues	SN	
Pick and assign	PA	
Backbone assignment	BB	
Peak Assigner	AA	
Assignment Inspector	Al	
Residue information	RI	
View		
View:	СТ	
NmrAtom Chemical Shift Table	NT	
NmrResidue Table		
Peak Table	LT	
Integral Table	IT pr	
Restraint Table	RT	
Structure Table	ST	
Sequence Graph	SG	
Atom Assigner	AS	
Chemical Shift Mapping	CM	

Notes Table	NO	
Python Console	Space-Space	
For current strip		
Show/Hide Toolbar	TB	
Show/Hide Phasing Console	PC	PV, PT, TR
Move to Next Spectrum (and Hide Current)	TabTab	
Move to Previous Spectrum (and Hide Current)	TabQ	
Toggle all Spectra on Current Display	TabA	
Store Zoom	ZS	ZI, ZO, ZR
Restore Zoom	ZR	ZI, ZO, ZS
Flip X-Y Axis	XY	XZ, YZ
Flip X-Z Axis	XZ	XY, YZ
Flip Y-Z Axis	YZ	XY, XZ
Macro :		
Define User Shortcuts	DU	
Reserved for user macros	Q0, Q1,Q9	
	1110 1114 1110	

DU
Q0, Q1,Q9
W0, W1,W9
Y0, Y1,Y9