



RasMol v2.7.4.1

Quick Reference Card

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Mouse Buttons

Clicking on an atom identifies that atom in the command window. Moving the mouse whilst holding mouse buttons and/or control keys manipulates the molecule. The default bindings are described below.

Mac	Windows	Action
---	Left	Rotate X-Y (or Z)
Command	Right	Translate X-Y
Shift	Shift Left	Zoom
Shift-Cmnd	Shift Right	Rotate Z
Control	Control Left	Z-Clipping (Slab)

General Commands

load [format] <filename> Load a molecule (up to 5)

pdb	Brookhaven Protein Databank
mdl	Molecular Design Limited's Mol file
mol2	Tripos' Sybyl Mol2 file format
alchemy	Tripos' Alchemy file format
charmm	CHARMm format card file
xyz	MSC's XMOL XYZ file format
mopac	J.P. Stewart's MOPAC file format
cif	IUCr CIF or mmCIF file format

exit Exit from RasMol Script
quit Terminate pgm execution

help [topic [subtopic]] Display on-line help topic

select <expression> Update part of molecule

restrict <expression> Display only part of mol.

set bondmode [mode] Change bond selection

script <filename> Execute file of commands

zap Delete molecule

Bond Commands

bond <src> <dst> + Add a bond

bond <src> <dst> pick Pick bond for rotation

unbond <src> <dst> Remove a bond

Display Commands

wireframe [boolean] Display wireframe

wireframe <rad> Display stick bonds

set bondmode all Mark all atoms

set bondmode none Mark no atoms

set bondmode not bonded Mark non-bonded atoms

set bondmode not bonded Mark non-bonded atoms

spacefill [boolean] Display spacefill spheres
spacefill <value> Specify atom sphere radius

spacefill temperature
spacefill user

star ... Display stars for spheres

surface molecule <value>

surface solvent <value>

Display molecule surface

Display alpha backbone

Specify backbone radius

ribbons [boolean] Display solid ribbons

ribbons <value> Specify ribbon width

strands [boolean] Draw ribbon as strands

strands <value> Specify ribbon width

set strands <value> Number of ribbon strands

label [boolean] Draw default atom labels

label <string> Label with arbitrary text

set fontsize <value> [FSIPS]

Set label font height

set fontstroke<value> Set label stroke width

ssbonds [boolean] Display disulphide bonds

ssbonds <value> Specify ssbond radius

set ssbonds backbone SSBonds between alphas

set ssbonds sidechain SSBonds between sulphurs

hbonds [boolean] Display hydrogen bonds

hbonds <value> Specify hbond radius

set hbonds backbone HBonds between alphas

set hbonds sidechain HBonds donor/acceptor

dots [boolean] Display dot surface

dots <value> Specify dot density

set solvent [boolean] VDW or solvent surface

set radius <value> Specify probe sphere rad.

set axes [boolean] Display co-ordinate axes

set boundingbox [boolean] Display bounding box

set unitcell [boolean] Display crystal unit cell

set monitor [boolean] Show distance monitor labels

set backfade [boolean] Shade to any background color

set display selected Currently selected portion

set picking Series of 13 commands:
off | ident | distancel angle
| torsion | label | monitor | center
coord | bond | atom | group | chain

Language Commands

Bulgarian Bulgarian menus and messages

Chinese Chinese menus and messages

English English menus and messages

French French menus and messages

Italian Italian menus and messages

Japanese Japanese menus and messages

Spanish Spanish menus and messages

Colour Commands

colour [object] <colour> Colour representation

Objects:

atoms	bonds	backbone
ribbons	labels	hbonds
ssbonds	dots	axes
ribbons1	ribbons2	

Predefined Colours:

Black	Blue	BlueTint	Brown
Cyan	Gold	Grey	Green
GreenBlue	GreenTint	HotPink	Magenta
Orange	Pink	PinkTint	Purple
Red	RedOrange	SeaGreen	SkyBlue
Violet	White	Yellow	YellowTon

Atom Colour Schemes:

cpk, cpknew	amino	shapely
group	chain	structure
temperature	charge	user
alt	model	

colour hbonds type Colour hbonds by offset

colour dots potential Display potential surface

Manipulation Commands

depth[boolean] Enable/disable back-clipping

depth <value> Move back-clipping plane

molecule <number> Specify molecule to manipulate

rotate <axis> [-] <value> Rotate molecule

rotate bond Rotate bond

rotate molecule Rotate selected molecule

rotate all Rotate all molecules

translate <axis> [-] <value> Translate molecule

zoom [boolean] Scale molecule

zoom <value> Specify magnification

slab [boolean] Enable/disable slabbing

slab <value> Move Z-clipping plane

centre [expression] {centre|translate}

Set centre of rotation

reset Initial transformation

set stereo [boolean] Control L&R images

Scripted Commands

load [format] inline Load molecule from script

pause Suspend script execution

echo Display text on command line

refresh Redraw image

set write [boolean] Save & write in scripts

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Map Commands

map {<map_selector>} {map_subcommand} <parameters>
<map_selector>: all, new, next or <number>
manipulate maps for current molecule

map {<map_selector>} **generate** {dots|mesh|surface}
generate a map from the current atoms

map {<map_selector>} **level** {MEAN} <number>
select a contour level

map {<map_selector>} { **load** | **save** } <filename>
load a CBF or CCP4 map, save a CBF map

map {<map_selector>} **mask** { **selected** | <number> | **none** }
generate a mask from selected atoms or
an existing map by number

map {<map_selector>} { **resolution** | **spacing** | **spread** } <number>
set spacing and/or spread, spacing=resolution
spread = 2/3 resolution

map {<map_selector>} {**restrict** | **select**} **select** maps
if restrict disable display of the others

map {<map_selector>} **show**
show information about selected maps

map {<map_selector>} **zap**
erase selected maps

Atom Expressions

Predefined Sets: alpha
hydrophobic

Residue Ranges: 3,16,12
9-20

Boolean Operators: backbone and not alpha
ligand or 196-199

Primitive Expressions: cys, glu, arg, as?
ser70a, **p, glu24:1
hem*p.fe, *.sg

Comparison Operators: atomno=4,atomno=6
temperature>=900

Within Expressions: within(8.0,ligand)

Predefined Sets

at	acidic	acyclic	aliphatic
alpha	amino	aromatic	backbone
basic	bonded	buried	cg
charged	cyclic	cystine	helix
hetero	hydrogen	hydrophobic	ions
large	ligand	medium	neutral
nucleic	polar	protein	purine
pyrimidine	selected	sheet	sidechain
small	solvent	surface	turn

define <identifier> <expression>

User-defined sets

Rendering Commands

background <colour> Set background colour

set ambient [value] Depth-cueing/lighting

set shadows [boolean] Enable/disable shadows

set specular [boolean] Enable atom highlights

set specpower [value] Control atom 'shininess'

set shadepower [value] Control atom 'contrast'

Export Commands

write [format] <filename> Output image file

gif CompuServe GIF format

iris IRIS RGB

ps, epsf Encapsulated PostScript

monops Monochrome PostScript

vectps 'Cartoon' PostScript

bmp Microsoft Bitmap format

pict Apple 'PICT' file

ppm Portable Pixmap

sun, sunrle Sun Rasterfile

set vectps <boolean> Enable cartoon outlines

write script <filename> Generate RasMol script

write povray <filename> Generate POVray data

write vrml<filename> Generate VRMLdata

write molscript <filename>
Output MolScript script

write kinemage <filename>
Output Kinemage file

save <filename> Save selected atoms

set kinemage <boolean> Set Mage file detail

set transparent<boolean> Allow transparent GIFs

write phipsi<filename> Generate phi-psi data

write RDF<filename> Ramachandran plot data

write RPP<filename> Ramachandran printer plot

Misc. Commands

structure DSSP secondary structure

connect [boolean] Recalculate connectivity

renumber Sequentially number chains

show information Display molecule statistics

show phipsi Display torsion angles

show RPP Ramachandran printer plot

show sequence Display molecule sequence

show symmetry Display crystal space group

set mouse rasmol Default mouse bindings

set mouse quanta Polygen's Quanta bindings

set mouse insight Biosym's Insight II bindings

set cisangle CIS angle cutoff

Command Line Editing

In addition to the cursor keys, the following 'emacs' control keys may be used to edit the command line.

Ctrl-H / Ctrl-D Delete previous/next character

Ctrl-B / Ctrl-F Move backward/forward a character

Ctrl-A / Ctrl-E Move to beginning/end of line

Ctrl-P / Ctrl-N Display previous/next history

Colour Schemes

CPK Atom Colours

Carbon	light grey	[200,200,200]
Oxygen	red	[240,0,0]
Nitrogen	sky blue	[143,143,255]
Hydrogen	white	[255,255,255]
Sulphur	yellow	[255,200,50]
Phosphorous	orange	[255,165,0]
Chlorine	green	[0,255,0]
Bromine, Zinc	brown	[165,42,42]
Calcium	dark grey	[128,128,144]
Unknown	deep pink	[255,20,147]

CPKnew Atom Colours

Carbon	light grey	[211,211,211]
Oxygen	red	[255,0,0]
Nitrogen	sky blue	[135,206,235]
Hydrogen	white	[255,255,255]
Sulphur	yellow	[255,255,0]
Phosphorous	orange	[255,170,0]
Chlorine	green	[0,255,0]
Bromine, Zinc	brown	[128,40,40]
Calcium	dark grey	[105,105,105]
Unknown	deep pink	[250,22,145]

Amino Acid Colours

ASP, GLU	bright red	[230,10,10]
CYS, MET	yellow	[230,230,0]
LYS, ARG	blue	[20,90,255]
SER, THR	orange	[250,150,0]
PHE, TYR	mid blue	[50,50,170]
ASN, GLN	cyan	[0,220,220]
GLY	light grey	[235,235,235]
LEU, VAL, ILE	green	[15,130,15]
ALA	dark grey	[200,200,200]
TRP	pink	[180,90,180]
HIS	pale blue	[130,130,210]
PRO	flesh	[220,150,130]
others	tan	[190,160,110]

Secondary Structure Colours

Alpha Helix	magenta	[240,0,128]
Beta Sheet	yellow	[255,255,0]
Turns	pale blue	[96,128,255]
Other	white	[255,255,255]

Hydrogen Bond Type Colours

Offset +2	white	[255,255,255]
Offset +3	magenta	[255,0,255]
Offset +4	red	[255,0,0]
Offset +5	orange	[255,165,0]
Offset -3	cyan	[0,255,255]
Offset -4	green	[0,255,0]
default	yellow	[255,255,0]
