



# RasMol v2.5

## 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.

Left Button	Rotate X-Y
Right Button	Translate X-Y
Shift Left Button	Zoom
Shift Right Button	Rotate Z
Control Left Button	Z-Clipping (Slab)

### General Commands

<b>load</b> [format] <filename>	Load a molecule
<b>pdb</b>	Brookhaven Protein Databank
<b>mdl</b>	Molecular Design Limited's Mol file
<b>mol2</b>	Tripos' Sybyl Mol2 file format
<b>alchemy</b>	Tripos' Alchemy file format
<b>charmm</b>	CHARMm format card file
<b>xyz</b>	MSC's XMOL XYZ file format
<b>exit</b>	Exit from RasMol
<b>quit</b>	
<b>help</b> [topic [subtopic]]	Display on-line help topic
<b>select</b> <expression>	Update part of molecule
<b>restrict</b> <expression>	Display only part of mol.
<b>set bondmode</b> [mode]	Change bond selection
<b>script</b> <filename>	Execute file of commands
<b>zap</b>	Delete molecule

### Display Commands

<b>wireframe</b> [boolean]	Display wireframe
<b>wireframe</b> <value>	Display stick bonds
<b>spacefill</b> [boolean]	Display spacefill spheres
<b>spacefill</b> <value>	Specify atom sphere radius
<b>spacefill temperature</b>	
<b>spacefill user</b>	

<b>backbone</b> [boolean]	Display alpha backbone
<b>backbone</b> <value>	Specify backbone radius
<b>ribbons</b> [boolean]	Display solid ribbons
<b>ribbons</b> <value>	Specify ribbon width
<b>strands</b> [boolean]	Draw ribbon as strands
<b>strands</b> <value>	Specify ribbon width
<b>set strands</b> <value>	Number of ribbon strands
<b>label</b> [boolean]	Draw default atom labels
<b>label</b> <string>	Label with arbitrary text
<b>set fontsize</b> <value>	Set label font height
<b>ssbonds</b> [boolean]	Display disulphide bonds
<b>ssbonds</b> <value>	Specify ssbond radius
<b>set ssbonds backbone</b>	SSBonds between alphas
<b>set ssbonds sidechain</b>	SSBonds between sulphurs
<b>hbonds</b> [boolean]	Display hydrogen bonds
<b>hbonds</b> <value>	Specify hbond radius
<b>set hbonds backbone</b>	HBonds between alphas
<b>set hbonds sidechain</b>	HBonds donor/acceptor
<b>dots</b> [boolean]	Display dot surface
<b>dots</b> <value>	Specify dot density
<b>set solvent</b> [boolean]	VDW or solvent surface
<b>set radius</b> <value>	Specify probe sphere rad.
<b>set axes</b> [boolean]	Display co-ordinate axes
<b>set boundingbox</b> [boolean]	Display bounding box
<b>set unitcell</b> [boolean]	Display crystal unit cell

### Colour Commands

**colour** [object] <colour> Colour representation

#### Objects:

<b>atoms</b>	<b>bonds</b>	<b>backbone</b>
<b>ribbons</b>	<b>labels</b>	<b>hbonds</b>
<b>ssbonds</b>	<b>dots</b>	<b>axes</b>
<b>ribbons1</b>	<b>ribbons2</b>	

#### Predefined Colours:

<b>blue</b>	<b>black</b>	<b>cyan</b>	<b>green</b>
<b>greenblue</b>	<b>magenta</b>	<b>orange</b>	<b>purple</b>
<b>red</b>	<b>redorange</b>	<b>violet</b>	<b>white</b>
<b>yellow</b>			

#### Atom Colour Schemes:

<b>cpk</b>	<b>amino</b>	<b>shapely</b>
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<b>group</b>	<b>chain</b>	<b>structure</b>
<b>temperature</b>	<b>charge</b>	<b>user</b>
<b>colour hbonds type</b>		Colour hbonds by offset
<b>colour dots potential</b>		Display potential surface

### Manipulation Commands

<b>rotate</b> <axis> [-] <value>	Rotate molecule
<b>translate</b> <axis> [-] <value>	Translate molecule
<b>zoom</b> [boolean]	Scale molecule
<b>zoom</b> <value>	Specify magnification
<b>slab</b> [boolean]	Enable/disable slabbing
<b>slab</b> <value>	Move Z-clipping plane
<b>centre</b> [expression]	Set centre of rotation
<b>reset</b>	Initial transformation

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## Atom Expressions

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<b>Predefined Sets:</b>	alpha hydrophobic
<b>Residue Ranges:</b>	3,16,12 9-20
<b>Boolean Operators:</b>	backbone and not alpha ligand or 196-199
<b>Primitive Expressions:</b>	cys, glu, arg, as? ser70a, **p, glu24:1 hem*p.fe, *.sg
<b>Comparison Operators:</b>	atomno=4,atomno=6 temperature>=900
<b>Within Expressions:</b>	within(8.0,ligand)

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## Predefined Sets

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

**define** <identifier> <expression> User-defined sets

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## Rendering Commands

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<b>background</b> <colour>	Set background colour
<b>set ambient</b> [value]	Depth-cueing/lighting
<b>set shadows</b> [boolean]	Enable/disable shadows
<b>set specular</b> [boolean]	Enable atom highlights
<b>set specpower</b> [value]	Control atom 'shininess'

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## Export Commands

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<b>write</b> [format] <filename>	Output image file
<b>gif</b>	CompuServe GIF format
<b>ps, epsf</b>	Encapsulated PostScript
<b>monops</b>	Monochrome PostScript
<b>vectps</b>	'Cartoon' PostScript
<b>bmp</b>	Microsoft Bitmap format
<b>pict</b>	Apple 'PICT' file
<b>ppm</b>	Portable Pixmap

**sun, sunrle** Sun Rasterfile

<b>set vectps</b> <boolean>	Enable	cartoon outlines
<b>write script</b> <filename>	Generate	RasMol script
<b>write molscript</b> <filename>	Output	MolScript script
<b>write kinemage</b> <filename>	Output	Kinemage file
<b>set kinemage</b> <boolean>	Set	Mage file detail

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## Misc. Commands

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<b>structure</b>	DSSP secondry structure		
<b>connect</b> [boolean]	Recalculate connectivity		
<b>renumber</b>	Sequentially	number chains	
<b>show information</b>	Display molecule statistics		
<b>show sequence</b>	Display molecule sequence		
<b>show symmetry</b>	Display	crystal	space group
<b>set mouse rasmol</b>	Default mouse bindings		
<b>set mouse quanta</b>	Polygen's Quanta bindings		
<b>set mouse insight</b>	Biosym's	Insight	II bindings

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## Command Line Editing

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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

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## Colour Schemes

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### CPK Atom Colours

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

### 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]

### 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]