



# RasMol v2.7.2

## 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        |
| 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> [<rad>]** Display stick bonds  
**set bondmode all** Mark all atoms  
**set bondmode none** Mark no 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

**backbone [boolean]** Display alpha backbone  
**backbone <value>** 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> [FS|PS]** 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]**

**set display selected** Shade to any background color  
Currently selected portion

**set picking** Series of ten commands:  
off | ident | distance  
angle | torsion | label  
monitor | center | coord | bond

### 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>Black</b>     | <b>Blue</b>      | <b>BlueTint</b> | <b>Brown</b>      |
| <b>Cyan</b>      | <b>Gold</b>      | <b>Grey</b>     | <b>Green</b>      |
| <b>GreenBlue</b> | <b>GreenTint</b> | <b>HotPink</b>  | <b>Magenta</b>    |
| <b>Orange</b>    | <b>Pink</b>      | <b>PinkTint</b> | <b>Purple</b>     |
| <b>Red</b>       | <b>RedOrange</b> | <b>SeaGreen</b> | <b>SkyBlue</b>    |
| <b>Violet</b>    | <b>White</b>     | <b>Yellow</b>   | <b>YellowTont</b> |

**Atom Colour Schemes:**

|                    |               |                  |
|--------------------|---------------|------------------|
| <b>cpk</b>         | <b>amino</b>  | <b>shapely</b>   |
| <b>group</b>       | <b>chain</b>  | <b>structure</b> |
| <b>temperature</b> | <b>charge</b> | <b>user</b>      |
| <b>alt</b>         | <b>model</b>  |                  |

**colour hbonds type** Colour hbonds by offset  
**colour dots potential** Display potential surface

### Manipulation Commands

**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]** 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

## Atom Expressions

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

## Predefined Sets

|                   |                 |                    |                  |
|-------------------|-----------------|--------------------|------------------|
| <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

## Rendering Commands

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

## Export Commands

|  |                         |
|--|-------------------------|
| <b>write [format] &lt;filename&gt;</b> | Output image file       |
| <b>gif</b>                             | CompuServe GIF format   |
| <b>iris</b>                            | IRIS RGB                |
| <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         |
| <b>sun, sunrle</b>                     | Sun Rasterfile          |

**set vectps <boolean>** Enable cartoon outlines

|   |                           |
|---|---------------------------|
| <b>write script &lt;filename&gt;</b>    | Generate RasMol script    |
| <b>write povray &lt;filename&gt;</b>    | Generate POVray data      |
| <b>write vrml&lt;filename&gt;</b>       | Generate VRMLdata         |
| <b>write molscript &lt;filename&gt;</b> | Output MolScript script   |
| <b>write kinemage &lt;filename&gt;</b>  | Output Kinemage file      |
| <b>save &lt;filename&gt;</b>            | Save selected atoms       |
| <b>set kinemage &lt;boolean&gt;</b>     | Set Mage file detail      |
| <b>set transparent&lt;boolean&gt;</b>   | Allow transparent GIFs    |
| <b>write phipsi&lt;filename&gt;</b>     | Generate phi-psi data     |
| <b>write RDF&lt;filename&gt;</b>        | Ramachandran plot data    |
| <b>write RPP&lt;filename&gt;</b>        | Ramachandran printer plot |

## Misc. Commands

|                          |                              |
|--------------------------|------------------------------|
| <b>structure</b>         | DSSP secondry structure      |
| <b>connect [boolean]</b> | Recalculate connectivity     |
| <b>renumber</b>          | Sequentially number chains   |
| <b>show information</b>  | Display molecule statistics  |
| <b>show phipsi</b>       | Display trosion angles       |
| <b>show RPP</b>          | Ramachandran printer plot    |
| <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 |

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

### Amino Acid Colours

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

### Secondary Structure Colours

|             |                  |                      |
|-------------|------------------|----------------------|
| Alpha Helix | <b>magenta</b>   | <b>[240,0,128]</b>   |
| Beta Sheet  | <b>yellow</b>    | <b>[255,255,0]</b>   |
| Turns       | <b>pale blue</b> | <b>[96,128,255]</b>  |
| Other       | <b>white</b>     | <b>[255,255,255]</b> |

### Hydrogen Bond Type Colours

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