\left.|  | UCSF Chimera Quick Reference Guide |
| :--- | :--- |
|  | April 2013 |$\right]$|  | Commands (*reverse function command available) |
| :--- | :--- |

lighting
linewidth
longbond*
mask
match
matchmaker
matrixcopy
matrixget
matrixset
mclip*
mсору
measure
meshmol
minimize
modelcolor
modeldisplay*
molmap
morph
move
movie
$m s c^{*}$
namesel*
nucleotides*
objdisplay*
open*
pause
perframe*
play
preset
rainbow
rangecolor
read
represent
reset
resrenumber
ribbackbone*
ribbon*
ribclass
ribinsidecolor*
ribrepr
ribscale
ribspline
rlabel*
rmsd
rock
roll
rotation*
runscript
save
savepos*
scale*
scene*
scolor
section
adjust lighting and shininess
control the width of wire bonds
show/hide pseudobonds representing missing segments extract volume data bounded by surfaces
perform least-squares fitting of specified atoms
(mmaker) align models in sequence, then in 3D apply the transformation of one model to another write the current transformation matrices to a file read and apply transformation matrices from a file control per-model clipping
copy settings from one molecule model to another perform calculations on structures, surfaces, maps create a "molecule" to show surface mesh as sticks energy-minimize structures set color at the model level set display at the model level
create a density map from atomic coordinates morph (interpolate) between different structures translate models
capture image frames and assemble them into a movie color multiscale surfaces to match atoms
save and name the current selection
create special nucleotide representations
display graphical objects
read local files or fetch by ID
pause script execution until the user presses a key specify commands to be executed at each display frame script various complex motions
apply a predefined combination of display settings color residues, chains, or models over a range color over a range according to attribute values execute a command file, updating display at the end control atom/bond style (wire, stick, bs, sphere) restore default or saved orientations
reassign residue numbers
allow display of both ribbon and backbone atoms display ribbon
set ribbon residue class
set a separate color for inside protein helix ribbons control ribbon style (flat, edged, rounded) control ribbon scaling (Chimera default, licorice) control ribbon path (B-spline or cardinal spline) display residue labels
evaluate the RMSD between specified sets of atoms rock (rotate back and forth)
roll (rotate continuously)
make a bond rotatable
run Python script with command-line arguments
save the current Chimera session
save model positions
scale the view
save/restore scenes (positions, styles, colors, labels, etc.) color surfaces by volume data or geometry move global clipping planes in parallel
segment select* set* setattr*
shape
show*
sleep
solvate
sop
split
start stereo* stop surface* surfcat surfrepr swapaa swapna sym* system thickness tile*
topography transparency* turn
$v d w^{*}$
$v d w d e f i n e *$ $v d w d e n s i t y$ version viewdock volume vop
vseries
wait
window windoworigin windowsize* write writesel zonesel
act on segmentation models select atoms, (de)activate models for motion set visual effects, individual model rotation set an attribute to a specified value create a surface of a specified geometric shape display specified atoms, undisplay the others pause script execution for a specified time
add solvent using AmberTools adjust capping, edit surface models partition a molecule model into separate submodels start Chimera tools by name switch amongst stereo options and mono viewing exit from Chimera
calculate and display molecular surfaces (msms cat) group atoms for surface calculations (msms repr) control surface style (solid, mesh, dot) mutate amino acids or swap rotamers mutate nucleic acid residues generate symmetry-related copies of a structure send a command to the system shell move global clipping planes in opposite directions arrange models in a plane plot values in a volume data plane as surface heights make atoms/bonds, ribbons, and surfaces transparent rotate models
display van der Waals (VDW) dot surface set VDW radii set VDW surface dot density show copyright information and Chimera version start ViewDock and load docking results display volume data such as electron density edit volume data
display an ordered sequence of volume data sets suspend command processing until motion has stopped adjust the view to contain the specified atoms set graphics window location adjust the dimensions of the graphics window save atomic coordinates (pdb, mol2) write a list of the currently selected (or unselected) items select atoms/surfs within cutoff of specified atoms/surfs

Miscellaneous Operations (Default Settings)

| selection from screen | Ctrl-left mouse button |
| :--- | :--- |
| add/toggle selection | Shift-Ctrl-left mouse button |
| rotation | left mouse button |
| XY-translation | middle mouse button |
| scaling | right mouse button or Side View |
| preferences | Favorites... Preferences... |
| searching help | Help... Search Documentation... |
| reporting a problem <br> mailing list | Help... Report a Bug... <br> chimera-users@cgl.ucsf.edu |


|  | Specification Symbols |  |
| :--- | :--- | :--- |
| Symbol | Function | Usage |
| \# | model number | \# model (integer) |
| \#. | submodel number | \#. submodel (integer) |
| : | residue | : residue (name or number) |
| :: | residue name | :: residue |
| :. | chain ID | :. chain |
| @ | atom name | @atom |
| @. | alternate location ID | @. alt_loc |
| - | range | specifies a range of models, <br> submodels, or residues |
|  |  | separates models or residues, |
|  |  | ranges of models or residues, or |
| names of atoms |  |  |

## @/element=atno

@/idatmType=type
@/label
@/label=label
@/labelColor=labcolor
@/name=name
@/occupancy=occupancy
@/radius=radius
@/serialNumber=n
@/surfaceCategory=category
@/surfaceDisplay
surface) surface) surface)

## Selected Residue Attributes

| Usage | Description |
| :--- | :--- |
| :/areaSAS=sasa | solvent-accessible surface area |
| :/areaSES=sesa | solvent-excluded surface area |
| :/isHet | residues in PDB HETATM <br> records (or the mmCIF <br> equivalent) |
| :/isHelix | amino acid residues in helices |
| :/isStrand or :/isSheet | amino acid residues in strands |
| :/kdHydrophobicity=value | Kyte-Doolittle amino acid <br> hydrophobicity |
| :/phi=angle | protein/peptide backbone phi <br> angle |
| :/psi=angle | protein/peptide backbone psi <br> angle |
| :/ssId=N | secondary structure element <br> identifier (1 for first helix and <br> first strand, etc.) |
| :/uniprotIndex=N | residue number in corresponding <br> UniProt sequence, if any |

## Selected Molecule Model Attributes

| Usage | Description |
| :--- | :--- |
| \#/ballScale=factor | ball radius relative to VDW radius |
| \#/color=color | model-level color assignment |
| \#/display | model display bit |
| \#/lineWidth=width | linewidth of wire representation |
| \#/numAtoms=N | total number of atoms |
| \#/numResidues=M | total number of residues |
| \#/stickScale=factor | stick radius relative to bond radius |

\#

- all models
\#0
- model 0
\#3:45-83,90-98
- residues 45-83 and 90-98 in model 3
:lys,arg
- lysine and arginine residues
:12,14@ca
- alpha carbons in residues 12 and 14
:12:14@ca
- all atoms in residue 12 and the alpha carbon in residue 14
:.A@ca,c,n,o
- peptide backbone atoms in chain A
:50.B,.D
- residue 50 in chain B and all residues in chain D
:12-15,26-28.a,45.b
- residues 12-15 in all chains (except het/water), 26 - 28 in chain A, and 45 in chain B
\#0.1-3,5
- submodels 1-3 of model 0 and all of model 5


## \#0.1-3,. 5

- submodels 1-3 of model 0 and submodel 5 of all models
ligand
- any/all residues automatically classified as ligand
$\mathbf{S} \mid \mathbf{F e}$
- all sulfur and iron atoms
@ca/!label and color!=green and color!=red
- atoms named CA which are not labeled, and are not green or red
@/bfactor>=20 and bfactor<=40
- atoms with B-factor values ranging from 20 to 40


## :asn \& helix

- asparagine residues in helices
\#1:asp,glu \& \#0 z<10
- aspartate and glutamate residues in model 1 within 10 angstroms of model 0
solvent \& Ng+ z<3 | solvent \& N3+ z<3
- solvent residues within 3 angstroms of guanidinium nitrogens or
$s p 3$-hybridized, formally positive nitrogens
@/bfactor>50 \& ~ solvent \& ~ions
- atoms with B-factor values over 50, excluding solvent and ions

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