UCSF Chimera QUICK REFERENCE GUIDE June 2008

mask

match

mclip*

move

movie msc*

neon

open*

pdbrun

preset

read

reset

ribrepr

rlabel*

rmsd rock

roll

save

scale*

section select*

setattr*

shape

show*

sleep

solvate

source

split

set*

Commands

*reverse function ~command available

2 dlabala	aroute arbitrary text labels and place them in 2D	
2dlabels	create arbitrary text labels and place them in 2D	
ac addaa	enable accelerators (keyboard shortcuts) add an amino acid to a peptide C-terminus	
	1 1	
addcharge	assign partial charges to atoms	
addh	add hydrogens	
alias*	create an alias or list the existing aliases	
align	align two atoms or sets of atoms along the line of sight	
angle	measure a bond angle or torsion angle	
bond*	add/delete bonds	
bondcolor*	color bonds independently from atoms	
bonddisplay	control how bond display depends on atom display	
bondrepr	control bond style (wire, stick)	
bondzone*	make zoning tools use points along bonds	
brotation	make a bond rotatable	
cd	change the working directory	
center	center the view on specified atoms	
chain	chain specified atoms, undisplay the others	
chirality	report the R/S configuration of a chiral center	
clip*	move clipping planes	
close	close a model	
cofr*	report or change the center of rotation	
color*	color atoms/bonds, ribbons, labels, and surfaces	
colordef	define a new color	
conic	create a shadowed space-filling image	
сору	save an image (Chimera graphics or POV-Ray)	
crystalcontacts	identify clashes between PDB symmetry copies	
defattr	assign attribute values to atoms, residues, or models	
delete	delete atoms and bonds	
display*	display specified atoms	
distance*	measure the distance between two atoms	
echo	send text to the status line and Reply Log	
export	save the scene (x3d, vrml, pov-ray, renderman, obj)	
findclash*	identify clashes and/or contacts	
focus	adjust the view and center of rotation	
freeze	stop all motion	
getcrd	report untransformed coordinates	
hbonds*	(findhbond) identify possible hydrogen bonds	
help	display the manual page for a command	
hkcage	create a hexagon/pentagon mesh that covers an icosahedron	
intersurf	generate and display interface surfaces	
ksdssp	determine secondary structure from protein coordinates	
label*	display atom labels	
labelopt	control the information in atom labels	
linewidth	control the width of wire bonds	
load	restore a saved Chimera session	

longbond* show/hide pseudobonds representing missing segments extract volume data bounded by surfaces superimpose two models matrixcopy apply the transformation matrix of one model to another write the current transformation matrices to a file matrixget matrixset read and apply transformation matrices from a file control per-model clipping meshmol create a "molecule" from surface mesh for stick display minimize energy-minimize structures mmaker (matchmaker) align models in sequence, then in 3D modelcolor set color at the model level modeldisplay* set display at the model level molmap create a density map from atomic coordinates translate along the X, Y, or Z axis capture image frames and assemble them into a movie color multiscale surfaces to match atoms name and save the current selection namesel create a shadowed stick/tube image (not on Windows) objdisplay* display graphical objects read local files or fetch by ID send an annotated PDB file to the system shell perframe* specify an alias to be executed at each display frame apply a predefined combination of display settings push or pop images on the picture stack push,pop color residues, chains, or models over a range rainbow rangecolor color over a range according to attribute values execute a command file, updating display at the end represent control atom/bond style (wire, stick, bs, sphere) restore default or saved orientations ribbackbone* allow display of both ribbon and backbone atoms ribbon* display ribbon ribcolor* set ribbon color ribinside color*set a separate color for inside protein helix ribbons control ribbon style (flat, edged, rounded) ribscale control ribbon scaling (Chimera default, licorice) display residue labels evaluate the RMSD between specified sets of atoms rock about the X, Y or Z axis roll about the X, Y, or Z axis rotation make a bond rotatable save the current Chimera session savepos* save the current orientations scale the view move the clipping planes in parallel activate models for motion or select atoms set options (see Set/Unset Options) set an attribute to a specified value create a surface of a specified geometric shape display specified atoms, undisplay the others pause command processing add solvent using AmberTools execute a command file, updating display continually make chains of a molecule model separate submodels

start	start Chimera tools by name
stereo	switch amongst stereo options and mono viewing
stop	exit from Chimera
surface*	calculate and display molecular surfaces
surfcat	(msms cat) group atoms for surface calculations
surfcolor	set surface color source
surfrepr	(msms repr) control surface style (solid, mesh, dot)
surftransparency	* adjust molecular surface transparency
swapaa	mutate amino acids or swap rotamers
swapna	mutate nucleic acid residues
sym*	generate symmetry copies that update automatically
system	send a command to the system shell
thickness	move the clipping planes in opposite directions
topography	plot values in a volume data plane as surface heights
turn	rotate about the X, Y, or Z axis
vdw*	display van der Waals (VDW) surface
vdwdefine*	set VDW radii
vdwdensity	set VDW surface dot density
version	show copyright information and Chimera version
viewdock	start ViewDock and load docking results
volume	visualize volume data such as electron density
vop	edit volume data to create a new volume data set
wait	suspend command processing until motion has stopped
window	adjust the view to contain the specified atoms
windowsize	adjust the dimensions of the graphics window
write	save atomic coordinates (pdb, mol2)
writesel	write a list of the currently selected (or unselected) items

Set/Unset Toggle Options

autocolor	make each newly opened model a unique color	
independent	t make each model rotate about its own center of m instead of the combined center of mass	
	instead of the combined center of mass	
	Set/Unset Value Options	

bg_color	set background color; value can be any color name
dc_color	set depth cue color; value can be any color name

Miscellaneous Operations (Default Settings)

	······································
Action	Procedure
selection from screen	Ctrl-left mouse button
add/toggle selection	Shift-Ctrl-left mouse button
XY-rotation	left mouse button inside "spaceball"
Z-rotation	left mouse button outside "spaceball"
XY-translation	middle mouse button
Z-translation	Ctrl-middle mouse button
scaling	right mouse button or the Side View
preferences	Favorites Preferences
searching help	Help Search Documentation

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Spe	cific	ation	Sym	bols
- OPC	cinc	auton	- J J II	10010

Symbol

#

#.

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:. @

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Function

residue

chain ID

range

atom name

model number

residue name

submodel number

alternate location ID

name separator

whole wildcard

partial wildcard

@/display

Usage		"on"
# model (integer) #. submodel (integer)	@/drawMode=mode	<i>mode</i> can be 0 (dot, as in wireframe), 1 (sphere, as in CPK), 2 (endcap, as in stick), or
: residue (name or number)		3 (ball, as in ball-and-stick)
:: residue	<pre>@/element=atno</pre>	atomic number
:. chain	<pre>@/idatmType=type</pre>	Chimera atom type
@atom	@/label	whether the atom is labeled
@.alt_loc	@/label=label	text of the atom label
specifies a range of models,	@/labelColor=labcolor	color of the atom label
submodels, or residues	@/name=name	atom name
separates models or residues, ranges of models or residues, or	@/occupancy=occupancy	crystallographic occupancy
names of atoms	@/radius=radius	current VDW radius
matches whole atom or residue	@/serialNumber=n	serial number in the input file
names, <i>e.g.</i> ,:*@ CA specifies the alpha carbons of all residues	@/surfaceCategory=category	surface calculation category (main, ligand, <i>etc</i> .)
matches partial atom or residue names, <i>e.g.</i> , @C= specifies all atoms with names beginning with C	@/surfaceDisplay	per-atom surface display bit (can be true for buried atoms with no surface)
used for atom and residue names only, <i>e.g.</i> , :G?? selects all residues with three-letter names		sidue Attributes
beginning with G		vent-accessible surface area

Usage	Description
:/areaSAS=sasa	solvent-accessible surface area
:/areaSES=sesa	solvent-excluded surface area
:/isHelix	whether the residue is in an alpha helix
:/isHet	whether the residue is in PDB HETATM records (or the mmCIF equivalent)
:/isStrand or :/isSheet	whether the residue is in a beta strand
:/isTurn	whether the residue is assigned to a turn in the input file
:/kdHydrophobicity=value	Kyte-Doolittle amino acid hydrophobicity
:/ribbonColor=ribcolor	color of the residue's ribbon segment
:/ribbonDisplay	per-residue ribbon display bit (can be true for residues such as water that cannot be shown with ribbon)
:/type=resname	residue name

whether the atomic display bit is

.. ..

Selected Molecule Model Attributes

Usage	Description
#/ballScale=factor	ball size relative to VDW radius
#/color=color	model-level color assignment
#/display	model display bit
#/explicitHydrogens	whether the model has hydrogen atoms
#/lineWidth=width	linewidth of wire representation

Specification Examples

#	
- all	models
#0	110
	del 0
	5-83,90-98 idues 45-83 and 90-98 in model 3
•	ine and arginine residues
	14@ca ha carbons in residues 12 and 14
	14@ca atoms in residue 12 and the alpha carbon in residue 14
	@ca,c,n,o
	ptide backbone atoms in chain A
	B,.D idue 50 in chain B and all residues in chain D
- res	15,26-28.a,45.b idues 12-15 in all chains (except het/water), 26-28 in chain A, 45 in chain B
# 0.1 - sut	-3,5 pmodels 1-3 of model 0 and all of model 5
	-3,.5 pmodels 1-3 of model 0 and submodel 5 of all models
liga - ang	nd y/all residues automatically classified as ligand
	tent.S sulfur atoms
	// !label and color!=green and color!=red ms named CA which are not labeled, and are not green or red
	olor=yellow or color=blue and label ons that are yellow and atoms that are both blue and labeled
	/ isHelix baragine residues in alpha helices
	sp,glu & #0 z<10
	partate and glutamate residues in model 1 within 10 angstroms
- sol	ent & Ng+ z<3 solvent & N3+ z<3 vent residues within 3 angstroms of guanidinium nitrogens or hybridized, formally positive nitrogens
	factor>50 & ~ solvent & ~ ions ms with B-factor values over 50, excluding solvent and ions

UCSF Chimera was developed by the Computer Graphics Laboratory at the University of California, San Francisco, under support of NIH grant P41-RR01081. The software is copyrighted and licensed by the Regents of the University of California.

?	single-char wildcard	used for atom and residue names only, <i>e.g.</i> , :G?? selects all residues with three-letter names beginning with G
;	command separator	separates multiple commands on a single line
Z<	zone specifier	<pre>z<zone all<br="" or="" specifies="" zr<zone="">residues within zone angstroms, za<zone all="" atoms<br="" specifies="">(rather than entire residues) within that distance. Using > instead of < gives the complement.</zone></zone></pre>
&	intersection	intersection of specified sets
1	union	union of specified sets
~	negation	negation of specified set (when space-delimited)

Selected Atom Attributes

Usage	Description
@/altLoc=altloc	alternate location ID
@/areaSAS=sasa	solvent-accessible surface area
@/areaSES=sesa	solvent-excluded surface area
@/bfactor=bfactor	B-factor
@/color=color	atom-level color assignment
<pre>@/defaultRadius=rad</pre>	default VDW radius