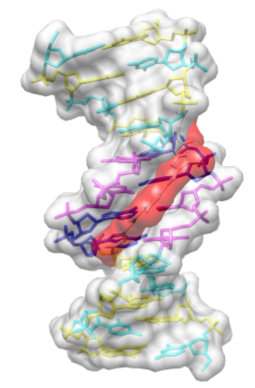
### **UCSF Chimera - Getting Started**

This tutorial provides an overview of basic features in Chimera. You can interact with Chimera using menus and/or commands. The basic features of Chimera are available either way, but not all command functions are available in menus or graphical interfaces, and not all menu or graphical interface functions are available in commands. Thus, it is useful to become familiar with both ways of interacting with Chimera.

The Working with menus and **Working with commands** sections are independent of each other and (for the most part) cover identical operations, accomplished in different ways. If you go through both sections, you can skip portions that cover issues you already understand. You can also go back and forth



DNA helix with bound netropsin between the sections to see the correspondence between menu and command

#### **Outline:**

operations.

- Working with menus Part 1
  - Getting started
    - Opening a structure
    - Side View
  - Using the mouse
    - Selection with the mouse
  - Selection/Action
  - Models and model status
- Working with menus -

#### Part 2

- Setup
- Representations
- Surfaces
- Front image how-to (menu)

**Typographical Conventions** 

-JF - <b>J</b>		
Item	Example	Description
Keyboard key	Ctrl	The control key
Mouse key	Btn1	Mouse button 1 (left button)
Menu action	File→Open	File Menu bar pulldown, followed by Open

#### • Working with commands - Part 1

- Getting started
  - Opening a structure
  - Side View
- Using the mouse
  - Selection with the mouse
- Command/Target
- Models and model status
- Working with commands Part 2
  - <u>Setup</u>
  - Representations
  - Surfaces
- Front image how-to (commands)

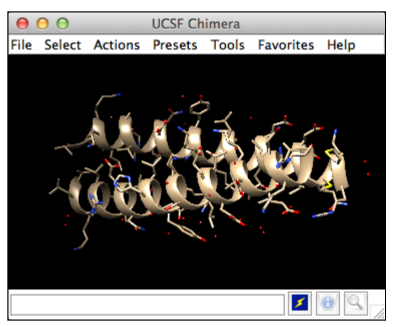
### Working with Menus, Part 1 - Manipulation, Selection, and Chains

#### ← Getting started

Start Chimera by clicking or doubleclicking the Chimera icon (depending on its location). Typically, this icon

location). Typically, this icon will be present on the desktop. The Chimera executable can also be run from its installation location.

A splash screen will appear, to be replaced in a few seconds by the main Chimera window containing either the graphics display or the **Rapid Access** list of recently used files (it doesn't matter which, since opening a structure will automatically



**UCSF Chimera with 1zik** 

switch the display to the graphics window). If you like, enlarge the window by clicking and dragging its lower right corner. The window can also be moved by clicking its top bar and dragging.

#### Opening a structure

Now open a structure. Choose **File→Fetch by ID** from the menu and enter **1zik** as the **ID** of the structure to fetch from the **PDB**. The structure will appear in the graphics window; it is a leucine zipper formed by two peptides.

The default initial display is ribbons. To also display atoms:

#### Actions→Atoms/Bonds→show

This shows all of the atoms and bonds in the structure, except that those in the peptide backbone are suppressed by the ribbon display. How to indicate specific parts of a structure for display, coloring, *etc.* is discussed <u>below</u>. Initially, heteroatoms (atoms other than carbon) are color-coded by element: oxygens red, nitrogens blue, *etc.* The carbons retain the model color, in this case tan.

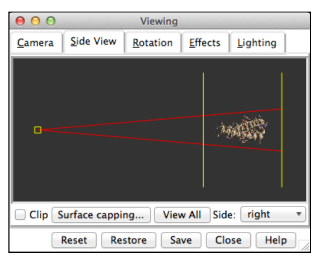
Hide ribbons to reveal the backbone atoms, then show ribbons again:

Actions→Ribbon→hide Actions→Ribbon→show

#### **Side View**

Show the **Side View** for interactive scaling and clipping (menu: **Tools→Viewing Controls→Side View**). By default, the **Side View** is also listed in the **Favorites** menu.

It shows a tiny version of the structure. Try moving the eye position (the small square) and the clipping planes (vertical lines) by clicking and dragging with the left mouse button. The **Side View** will renormalize itself after movements, so that the eye or clipping plane positions may appear to "bounce back," but your adjustments have been applied.



Side View showing 1zik

#### ← Using the mouse

Try manipulating the structure in the main window with the mouse. By default:

 the left mouse button controls rotation

#### **Default Mouse Button Assignments**

Mouse button	Modifier	Action
Btn1 (left button)		Rotation
Btn2 (middle button)		XY Translation
Btn3 (right button)		Scaling
Btn1	Ctrl	Picking (selection)
Btn1	Ctrl-Shift	Addition to (removal from) selection

- the middle mouse button controls XY translation (panning)
- the right mouse button performs scaling (zooming)

If you are using a touchpad or single-button mouse, modifier keys allow emulating the middle and right mouse buttons. These are **option** and **command** (♯) on **Mac** keyboards. Multitouch gestures on a Mac touchpad are enabled by default, but can be turned off in the preferences (menu: Favorites→Preferences, category Mouse). Continue moving and scaling the structure with the mouse in the graphics window and **Side View** as desired throughout the tutorial.

When the mouse focus is in the graphics window (you may need to click into it if you have been interacting with a different window), hovering the mouse cursor over an atom or bond without clicking any buttons will show identifying information in a pop-up "balloon." The balloon will disappear when the cursor is moved away. For an atom, the balloon information is of the form:

#### res-name res-num.chain atom-name

You can see from the balloons that this structure contains two peptide chains, A and B, and water (HOH residues), also with chain identifiers A and B.

#### **Selection with the mouse**

In combination with modifier keys, the mouse buttons have additional functions. By default, *picking* from the screen (a type of *selection*) is done by <a href="Ctrl-clicking">Ctrl-clicking</a> an atom or bond with the left mouse button, <a href="Btn1">Btn1</a>. You can also drag out a selection area with <a href="Ctrl-Btn1">Ctrl-Btn1</a> adds to or toggles an existing selection. The selection is outlined in green, and placing the mouse cursor over the green magnifying glass icon <a href="mailto:near the bottom right corner of the window reports what is selected in a pop-up "balloon."

The arrow keys can be used to broaden ( $\uparrow$ ), narrow ( $\downarrow$ ), or invert ( $\rightarrow$ ) a selection. The hierarchy for broadening and narrowing a selection could include (depending on the initial selection): atom/bond, residue, protein secondary structure element, bonded set of atoms, all atoms with the same chain ID, entire

model. When a selection is inverted, the selected atoms become deselected and *vice versa*.

Spend some time selecting various parts of the structure. An easy way to clear the selection (deselect everything) is to use <a href="Ctrl-Btn1">Ctrl-Btn1</a> in any blank space in the graphics window.

 $\leftarrow$ 

#### **Selection/Action**

In general, operations performed with the Chimera Actions menu apply to the current **selection**. Selections can be made in many ways, including with the Select menu or with the mouse (as described above). When nothing is selected, the Actions menu applies to everything.

The following will color all residues named LYS hot pink.

#### **Actions Menu Items**

	Actions Memoritans	
Menu Item	Description	
Atoms/Bonds	Controls the display and representation of atoms and bonds.	
Ribbon	Controls the display and representation of ribbons.	
Surface	Controls the display and representation of molecular surfaces.	
Color	Colors selected objects. Color target can be limited to object types in the <b>all options</b> dialog.	
Label	Labels selected atoms. The <b>residue</b> submenu labels residues containing the selected atoms.	
Focus	Focuses the view on the selected atom(s), zooming and translating if necessary.	
Set Pivot	Sets the center of rotation based on the selected atom(s) without adjusting the view.	
Inspect	Launches the Selection Inspector; same as clicking $lacktriangle$ .	
Write List	Writes a list of the currently selected objects to a parsable text file.	
Write PDB	Writes the coordinates of the currently selected atoms to a PDB file.	

#### Select→Residue→LYS Actions→Color→hot pink

The selection is highlighted in green, and the magnifying glass icon lear the bottom right corner of the window is also green, indicating that something is selected. Clearing the selection (deselecting) beforehand will color everything:

Select→Clear Selection Actions→Color→hot pink

**Select** menu choices also include chain ID, element, and many other categories of atoms and residues. (More complicated selections can be built up by intersecting some of these choices, as shown near the end of <a href="Part 2">Part 2</a>.)

Select→Chain→B
Actions→Color→cyan
Actions→Ribbon→hide
Select→Structure→solvent
Actions→Atoms/Bonds→hide
Select→Chemistry→element→N
Actions→Atoms/Bonds→sphere
Select→Clear Selection
Actions→Atoms/Bonds→stick

One way to select specific residues or ranges of residues is in the Sequence tool

(menu Favorites→Sequence, show sequence for chain A). When the sequence window has mouse focus, placing the cursor over a residue symbol in the sequence shows information for the corresponding structure residue at the bottom of the window. Click-drag a box within the sequence window to select one or more residues (as opposed to simply clicking within the light yellow box, which will select the entire helix), then hide their atoms:

#### Actions→Atoms/Bonds→hide

Quit from the sequence window and display all protein atoms again:

Select→Structure→protein Actions→Atoms/Bonds→show

Coloring can be limited to only certain representations, such as atoms only (not ribbons, surfaces, *etc.*):

Select→Residue→GLU Actions→Color→all options

In the resulting Color Actions dialog:

- 1. choose to Show all colors (lower right)
- 2. change the Coloring applies to (target) setting to atoms/bonds
- 3. click to choose any color

Notice that only the atoms/bonds of the selected residues and not their ribbon segments change color. Clear the selection (main menu Select→Clear Selection), then in the Color Actions dialog:

- 1. change the coloring target back to all of the above
- 2. click to choose tan (colors are listed alphabetically in the all-colors section)
- 3. click **Close** to dismiss the dialog

Restore heteroatom color-coding:

#### Actions→Color→by heteroatom

Coloring by heteroatom is useful for showing functional groups, yet keeping different models distinguishable by their different carbon colors.

Try picking two atoms in different residues (Ctrl-click the first, Shift-Ctrl-click the second). Show residue labels for the atoms you have selected:

#### Actions→Label→residue→name + specifier

(Actions→Label→name would show the atom names instead.) These 3D labels move along with structures and are mainly for interactive use. For figures and movies, 2D Labels are recommended instead.

Promote the selection to the entire residues with the keyboard up arrow  $\uparrow$  *or* the following:

Select→Broaden

Show only the selected atoms:

#### Actions→Atoms/Bonds→show only

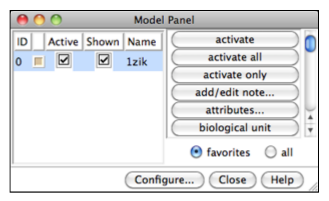
Clear the selection by **Ctrl**-clicking in empty space, as if picking "nothing." Turn off residue labels, hide ribbon, display all atoms, and color by element:

Actions→Label→residue→off Actions→Ribbon→hide Actions→Atoms/Bonds→show Actions→Color→by element

The **by element** coloring is the same as **by heteroatom** except it also color-codes carbons (gray).

#### ← Models and model status

Generally, each file of coordinates opened in Chimera becomes a *model* with an associated model ID number. Models are assigned successive numbers starting with 0. The *Model* Panel lists the current models and enables many operations upon them. Open this tool with Tools→General Controls→Model Panel.



**Chimera Model Panel** 

The columns on the left side of the Model Panel show:

- model ID number
- · model color
- A(ctive) whether activated for motion
- **S**(hown) whether display-enabled
- model name

Try toggling the checkboxes to see what happens; a model that is not activated for motion cannot be rotated or translated interactively. In the list of functions on the right side of the **Model Panel** (*not* the button at the bottom), click **close** to remove model **1zik**. Use the **Close** button at the bottom to close the **Model Panel**.

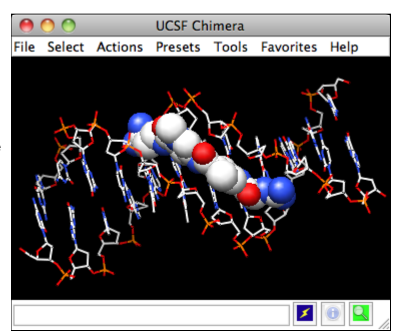
Go on to Part 2 below, or exit from Chimera with File→Quit.

# Working with Menus, Part 2 - Molecular Representations and Surfaces

#### ← Setup

With Chimera started as described at the beginning of Part 1, open a different structure. Choose File→Fetch by ID from the menu and enter 1d86 as the ID of the structure to fetch from the PDB. The structure contains the molecule netropsin bound to double-helical DNA, initially shown with ribbons and stylized representations of the nucleic acid sugars and bases.

Rotate, translate, and scale the structure as needed to get a better look (see <u>Using the</u> <u>mouse</u> to review how this is done). Continue moving and



done). Continue moving and Chimera showing netropsin as spheres scaling the structure as desired throughout the tutorial.

A *preset* is a predefined combination of display settings. Use the "all atoms" preset, which will show the DNA as wire and netropsin as spheres:

#### Presets→Interactive 2 (all atoms)

Color carbons white, then undisplay the water:

Select→Chemistry→element→C Actions→Color→white Select→Structure→solvent Actions→Atoms/Bonds→hide

Remember that hiding atoms does not deselect them; they remain selected, as indicated by the green magnifying glass icon near the bottom right of the window, until the selection is cleared or replaced with a new selection.

Residue names can be identified by looking in the **Select→Residue** menu or by hovering the cursor over an atom or bond to see information in a pop-up "balloon." Color the different nucleotides different colors, for example:

#### **Select**→**Residue**→**DA**

#### Actions→Color→blue

Analogously, color DC residues cyan, DG residues yellow, and DT residues magenta. Clear the selection with **Select\rightarrowClear Selection** or  $\boxed{\textbf{Ctrl}}$ -click in empty space.

#### ← Representations

Next, try some different display styles, or representations.

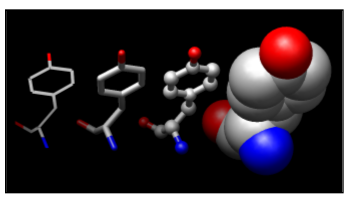
Actions→Atoms/Bonds→sphere Select→Chain→A

Actions→Atoms/Bonds→ball & stick Select→Clear Selection Actions→Atoms/Bonds→stick

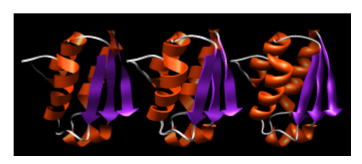
Showing ribbon automatically hides the mainchain (backbone) atoms.

Actions→Ribbon→show Actions→Ribbon→edged Actions→Ribbon→rounded

DNA can be shown with special nucleotide objects. We will show "lollipops," boxes, and a ladder.



Atoms/Bonds: wire, stick, ball & stick, and sphere



Ribbon: flat, edged, and rounded

#### Actions→Atoms/Bonds→nucleotide objects→settings

In the resulting **Nucleotides** dialog:

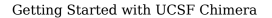
- 1. set Show side (sugar/base) as to tube/slab
- 2. set **Show base orientation** to **false**
- 3. click **Slab Style** tab, set slab style to **skinny**
- 4. click **Slab Options** tab, set **Slab object** to **ellipsoid**
- 5. click **Apply**; these are the "lollipops"

Nucleotide settings can be applied to just the selected residues (not necessarily all of the DNA). One way to select specific residues is in the **Sequence** tool:

#### **Favorites→Sequence**

Show the sequence of chain A and select one or a few residues in the sequence window with the mouse; this selects the corresponding part of the structure. Quit from the sequence window. In the **Nucleotides** dialog (also under **Tools→Depiction** in the menu):

- 1. set **Show base orientation** to **true**
- 2. set **Slab object** to **box**
- 3. click **Apply**; base orientations are shown with "bumps"



https://www.rbvi.ucsf.edu/Outreach/Tutorials/Get...

Clear the selection (Select→Clear Selection), then use Nucleotides to show the DNA as a ladder:

- set Show side (sugar/base) as to ladder
- 2. in the **Ladder Options**, set **Rung** radius to **0.3** Å
- 3. click **OK** (which will also dismiss the dialog)

To return to more general display styles, turn off the nucleotide objects:

Ribbons and nucleotide ladder

Actions→Atoms/Bonds→nucleotide objects→off

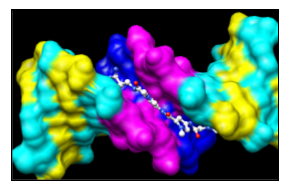
Hide the ribbons and show everything as ball-and-stick:

Actions→Ribbon→hide Actions→Atoms/Bonds→ball & stick

#### Surfaces

Finally, have some fun with molecular surfaces. There are built-in categories within structures such as main and ligand; when nothing is selected, Actions→Surface→show displays the surface of main.

Actions→Surface→show Actions→Surface→hide Select→Structure→ligand Actions→Surface→show Actions→Surface→mesh



Molecular surface (main)

Surface color can be specified separately from the colors of the underlying atoms. The ligand surface is tan and white because the original model color (tan) is used for surfaces of atoms not explicitly recolored by the user, and above, only the carbon atoms were changed to white. With the ligand still selected, choose Actions→Color→all options... to open the Color Actions dialog. In that dialog:

- 1. change the Coloring applies to (target) setting to surfaces
- 2. click red
- 3. click Close (which will automatically reset the coloring target back to all of the above)

Clear the selection, change back to a solid surface, and then undisplay the surface.

Select→Clear Selection

Actions→Surface→solid Actions→Surface→hide

As an example of a more complicated selection process, show the surface of the adenine and thymine deoxynucleotides in chain B only:

- 1. change the selection mode: Select→Selection Mode→append
- 2. Select→Residue→DA
- 3. Select→Residue→DT
- 4. change the selection mode: Select→Selection Mode→intersect
- 5. Select→Chain→B
- 6. Actions→Surface→show

To prepare for any subsequent operations, restore the selection mode and clear the selection:

```
Select→Selection Mode→replace
Select→Clear Selection (or Ctrl-click in empty space)
```

The command-line (Tools→General Controls→Command Line) equivalent is much more concise, but requires some knowledge of the atom specification syntax:

```
Command: surf:da.b,dt.b
```

Sometimes it is helpful to make a surface transparent:

```
Actions→Surface→transparency→50%
```

Choose File→Quit from the menu to terminate the Chimera session.

DNA helix with bound netropsin

#### ← Front image how-to (menu)

How to recreate the image at the front of the tutorial using the menu (see <a href="commands">commands</a>):

- 1. Choose File→Fetch by ID and fetch PDB entry 1d86
- 2. Use the all atoms preset:
  - Presets→Interactive 2 (all atoms)
- 3. Set the display style to stick:
  - o Actions→Atoms/Bonds→stick
- 4. Delete the waters:
  - Select→Structure→solvent
  - o Actions→Atoms/Bonds→delete
- 5. Color the residues:
  - Select→Residue→DA
  - o Actions→Color→blue
  - Select→Residue→DC
  - Actions→Color→cyan
  - Select→Residue→DG
  - Actions→Color→yellow
  - Select→Residue→DT
  - Actions→Color→magenta
  - Select→Residue→NT
  - Actions→Color→white
- 6. Broaden the selection to the whole chain and then to the whole model (both **ligand** and **main**), show surfaces, make them transparent:
  - Select→Broaden
  - Select→Broaden
  - Actions→Surface→Show
  - Actions→Surface→transparency→40%
- 7. Set coloring to surfaces only, make them light gray:
  - choose Actions→Color→all options... to show the Color Actions dialog, and in that dialog:
    - change the Coloring applies to (target) setting to surfaces
    - click light gray (keep the dialog open)
- 8. Select just netropsin again, make just its surface red:
  - Select→Residue→NT
  - in the Color Actions dialog:
    - click red (keep the dialog open)
  - Select→Clear Selection
- 9. Set coloring to background only, make it white:
  - in the Color Actions dialog:
    - change the coloring target to background
    - click white
    - click Close (which will automatically reset the coloring target back to all of the above)

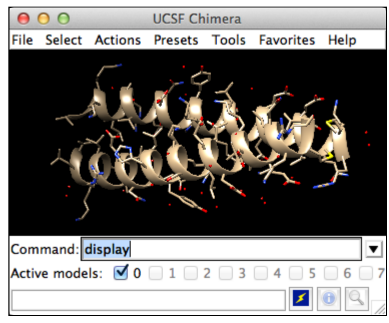
- 10. Adjust the view as desired
- 11. Save the image:
  - o File→Save Image

### Working with Commands, Part 1 - Manipulation, Selection, and Chains

#### ← Getting started

Start Chimera by clicking or doubleclicking the Chimera icon (depending on its location). Typically, this icon will be present on the desktop. The Chimera executable can also be run from its installation location.

A splash screen will appear, to be replaced in a few seconds by the main Chimera window containing either the graphics display or the **Rapid Access** list of recently used files (it doesn't matter which, since opening a structure will automatically



**Chimera with Command Line** 

switch the display to the graphics window). If you like, enlarge the window by clicking and dragging its lower right corner. The window can also be moved by clicking its top bar and dragging.

Show the Command Line with Tools→General Controls→Command Line. By default, the Command Line is also listed in the Favorites menu.

#### Opening a structure

Now open a structure. To fetch the structure of entry **1zik** from the Protein Data Bank (PDB), use the command:

Command: open 1zik

The structure will appear in the graphics window; it is a leucine zipper formed by two peptides.

The default initial display is ribbons. To also display atoms:

Command: **display** 

This shows all of the atoms and bonds in the structure, except that those in the peptide backbone are suppressed by the ribbon display. How to indicate specific parts of a structure for display, coloring, *etc.* is discussed <u>below</u>. Initially, heteroatoms (atoms other than carbon) are color-coded by element: oxygens red, nitrogens blue, *etc.* The carbons retain the model color, in this case tan.

Hide ribbons to reveal the backbone atoms, then show ribbons again:

Command: ~ribbon
Command: ribbon

Many commands have "~" versions that perform the opposite function.

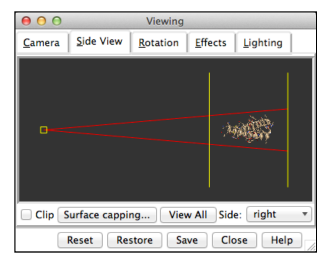
#### **Side View**

Show the **Side View** for interactive scaling and clipping:

Command: start Side View

By default, the **Side View** can also be started from the **Favorites** menu.

It shows a tiny version of the structure. Try moving the eye position (the small square) and the clipping planes (vertical lines) by clicking and dragging with the left mouse button. The **Side View** will renormalize itself after movements, so



Side View showing 1zik

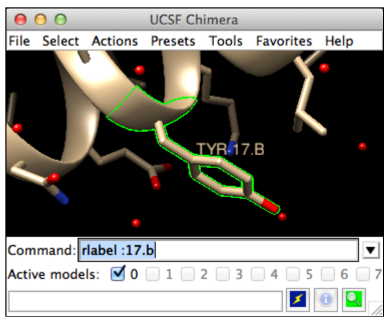
that the eye or clipping plane positions may appear to "bounce back," but your adjustments have been applied.

#### ← Using the mouse

Try manipulating the structure in the main window with the mouse. By default:

- the left mouse button Btn1 controls rotation
- the middle mouse button
   Btn2 controls XY
   translation (panning)
- the right mouse button
   Btn3 performs scaling
   (zooming)

If you are using a touchpad or single-button mouse, modifier keys allow emulating the middle and right mouse buttons. These are **option** and **command** (**X**) on **Mac** 



1zik with tyrosine 17 (B chain) selected

keyboards. Multitouch gestures on a Mac touchpad are enabled by default, but can be turned off in the preferences (menu: Favorites→Preferences, category Mouse). Continue moving and scaling the structure with the mouse in the graphics window and Side View as desired throughout the tutorial.

When the mouse focus is in the graphics window (you may need to click into it if you have been interacting with a different window), hovering the mouse cursor over an atom or bond without clicking any buttons will show identifying information in a pop-up "balloon." The balloon will disappear when the cursor is moved away. For an atom, the balloon information is of the form:

#### res-name res-num.chain atom-name

You can see from the balloons that this structure contains two peptide chains, A and B, and water (HOH residues), also with chain identifiers A and B.

#### Selection with the mouse

In combination with modifier keys, the mouse buttons have additional functions. By default, *picking* from the screen (a type of *selection*) is done by <a href="Ctrl">Ctrl</a>-clicking an atom or bond with the left mouse button, <a href="Btrl">Btrl</a>. Additionally pressing <a href="Shift">Shift</a> adds to or toggles an existing selection. The selection is outlined in green, and placing the mouse cursor over the green magnifying glass icon <a href="magnifying">near the bottom right corner of the window reports what is selected in a pop-up "balloon."</a>

The arrow keys can be used to broaden ( $\uparrow$ ), narrow ( $\downarrow$ ), or invert ( $\rightarrow$ ) a selection. The hierarchy for broadening and narrowing a selection could include

(depending on the initial selection): atom/bond, residue, protein secondary structure element, bonded set of atoms, all atoms with the same chain ID, entire model. When a selection is inverted, the selected atoms become deselected and *vice versa*.

Spend some time selecting various parts of the structure. An easy way to clear the selection (deselect everything) is to use <a href="Ctrl-Btn1">Ctrl-Btn1</a> in any blank space in the graphics window.

### ← Command/Target

A Chimera command may include *arguments* and a *target* (or *atom specification*). For example, in the following color command,

Command: color hot pink: lys

hot pink is an argument that specifies a color name, and :lys specifies that the target is all residues named LYS.

If no target is specified, the command acts on all applicable items. For example, the following makes everything hot pink:

Command: color hot pink

A basic specification may contain residue names, residue numbers, chain identifiers, and/or **Atom Specification Symbols** 

Symbol	Function	Usage
#	model number	# model (integer)
:	residue	: residue (name or number)
:.	chain ID	:.chain
@	atom name	@atom
*	whole wildcard	matches whole atom or residue names, e.g., :*@CA specifies the $\alpha$ -carbons of all residues
=	partial wildcard	matches partial atom or residue names, e.g., @C= specifies all atoms with names beginning with C
?	single-character wildcard	used for atom and residue names only, e.g., :G?? specifies all residues with three-letter names beginning with G
z<	zone specifier	z <zone or="" specifies<br="" zr<zone="">all residues within zone angstroms of the indicated atoms, and za<zone specifies<br="">all atoms (rather than entire residues) within zone angstroms of the indicated atoms. Using &gt; instead of &lt; gives the complement.</zone></zone>
&	intersection	intersection of specified sets
I	union	union of specified sets
~	negation	negation of specified set (when space-delimited)

atom names (see the table of symbols, right). Note also that commands can be truncated to unique strings.

Command: color yellow:20-22
Command: col gray:20-22.a
Command: ~disp:hoh.a

Command: ~disp inoling
Command: col cyan :.b
Command: ~ribbon :.b

Command: ~disp:5-10.a,15-20.b

Command: represent sphere @cd2

Command: rep stick

Command: col red: leu.b@ca

The Chimera Quick Reference Guide lists all of the commands and gives some examples of atom specification. It can be accessed by choosing Help→Tutorials from the Chimera menu and clicking the "Chimera Quick Reference Guide" link.

Many other types of specifications can be used, including element symbols and built-in classifications such as **solvent**:

Command: ~disp solvent
Command: color blue S
Command: disp protein

The command **help** can be used to show the manual page for any command:

Command: help color

As explained in the manual page, the **color** command also allows coloring only certain representations. For example, "**,a**" in the following means atoms only (not ribbons, surfaces, *etc.*):

Command: col gold,a:glu,lys

Restore the original coloring:

Command: col tan
Command: col byhet

Coloring by heteroatom is useful for showing functional groups, yet keeping different models distinguishable by their different carbon colors.

Try picking two atoms in different residues (Ctrl-click the first, Shift-Ctrl-click the second). Unlike the Actions menu, commands do not automatically act on the current selection. However, the current selection can be specified as the target of a command with the word selected, sel, or picked. Show residue labels for the atoms you have selected:

Command: rlabel sel

(The **label** command shows atom information instead.) The 3D labels from **rlabel** and **label** move along with structures and are mainly for interactive use. For figures and movies, **2D Labels** are recommended instead.

The following command can be used to promote the selection to the entire residues:

Command: select up

(The keyboard up arrow 🗈 also broadens a selection, but you may need to click

into the graphics window first to use that approach.) Show only the selected atoms:

Command: show sel

Clear the selection by **Ctrl**-clicking in empty space, as if picking "nothing." Turn off residue labels, hide ribbon, display all atoms, and color by element:

Command: ~rlab
Command: ~ribbon
Command: disp

Command: col byelement

Coloring **byelement** is the same as **byhet** except it also color-codes carbons (gray).

#### ← Models and model status

Generally, each file of coordinates opened in Chimera becomes a **model** with an associated model ID number. Models are assigned successive numbers starting with 0.

The Active models line right under the Command Line shows which models are activated for motion. Unchecking the box for 0 makes it impossible to rotate or translate model 0 interactively. Checking the box again restores the movable state. A similar row of checkboxes for toggling model display can be shown using the preferences (menu: Favorites→Preferences, category Command Line).

Remove the model:

Command: close 0

Go on to Part 2 below, **OR** exit from Chimera with the following command:

Command: **stop** 

## Working with Commands, Part 2 - Molecular Representations and Surfaces

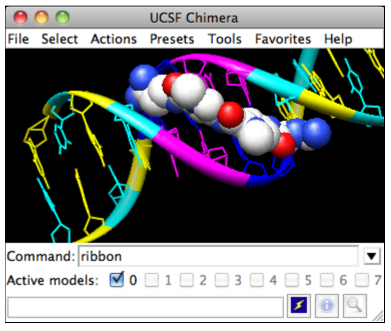
#### ← Setup

With Chimera started and the **Command Line** opened as described at the beginning of **Part 1**, fetch the structure of entry **1d86** from the Protein Data Bank (PDB):

Command: open 1d86

The structure contains the molecule netropsin bound to double-helical DNA, initially shown with ribbons and stylized representations of the nucleic acid sugars and bases.

Rotate, translate, and scale the structure as needed to get a better look (see <u>Using the</u> mouse to review how this is



**Chimera with Command Line** 

done). Continue moving and scaling the structure as desired throughout the tutorial.

A *preset* is a predefined combination of display settings. Use the "all atoms" preset, which will show the DNA as wire and netropsin as spheres:

Command: preset apply int 2

Color carbons white, then undisplay the water:

Command: color white C
Command: ~disp solvent

Residue names can be identified by looking in the **Select→Residue** menu or by hovering the cursor over an atom or bond to see information in a pop-up "balloon." Color the different nucleotides different colors, specifying them by residue name:

Command: color blue :da
Command: color cyan :dc
Command: color yellow :dg

Command: color magenta :dt

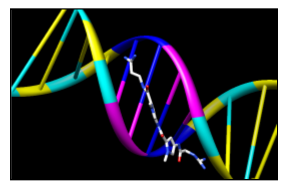
#### ← Representations

Next, try some different display styles, or representations.

Command: represent sphere

Command: repr bs :.a
Command: rep stick

Notice that commands but not necessarily their keyword arguments can be truncated to unique strings. For example, the command **represent** can be shortened to **repr** or **rep** but not **re** (because other commands also



Ribbons and nucleotide ladder

start with **re**), whereas its keywords **stick**, **sphere**, *etc*. cannot be truncated. If the truncation is not unique, one of the corresponding commands will be executed, but it may not be the one intended.

Showing ribbon automatically hides the mainchain (backbone) atoms.

Command: ribbon

Command: ribrep edged Command: ribr rounded

DNA can be shown with special nucleotide objects. We will show "lollipops," boxes with orientation bumps, and then a ladder. You can copy and paste into the **Command Line**. The command-line contents can be edited, and past commands can be accessed using the up and down arrow keys or **Ctrl-p** (previous) and **Ctrl-n** (next).

Command:

nuc side tube/slab shape ellipsoid orient false style skinny Command:

nuc side tube/slab shape box orient true style skinny :8-10.a

Command: nuc side ladder radius 0.3

To return to more general display styles, turn off the nucleotide objects:

Command: ~nuc

Hide the ribbons and show everything as ball-and-stick:

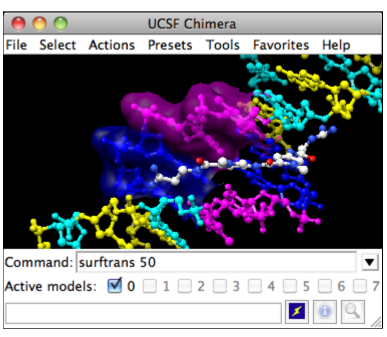
Command: ~ribbon Command: rep bs

#### ← Surfaces

Finally, have some fun with molecular surfaces. There are built-in categories within structures such as **main** and **ligand**; when nothing is specified, **surface** shows the surface of **main**.

Command: surface
Command: ~surf
Command: surf ligand
-OR- (equivalent)
Command: surf:nt

Surface color can be specified separately from the colors of the underlying atoms. The ligand surface is tan and white because the original model color (tan) is used for surfaces



Chimera showing a transparent surface

of atoms not explicitly recolored by the user, and above, only the carbon atoms were changed to white. Show the ligand surface as red mesh:

Command: surfrep mesh
Command: color red,s ligand
Command: surfrep solid

Parts of a surface can be shown:

Command: ~surf

Command: surf:da,dt

Command: ~surf

Command: surf:da.b,dt.b

Sometimes it is helpful to make a surface transparent:

Command: transp 50,s

When finished, exit from Chimera:

Command: stop now

#### ← Front image how-to (commands)

How to recreate the image at the front of the tutorial using commands (see menu approach):

- 1. Fetch 1d86:
  - Command: open 1d86
- 2. Use the all atoms preset:
  - Command: preset apply int 2
- 3. Set the display style to stick:
  - Command: repr stick
- 4. Delete the waters:
  - Command: del solvent
- 5. Color the residues:
  - Command: color blue :da
  - Command: color cyan :dc
  - Command: color yellow :dg
  - Command: color magenta :dt
  - o Command: color white :nt
- 6. Show surfaces for the whole model (both **ligand** and **main**), make them transparent:
  - Command: surf #0
  - Command: surftrans 40
- 7. Color the **main** (DNA) surface light gray and the **ligand** (netropsin) surface red:
  - Command: color light gray,s main
  - Command: color red,s ligand
- 8. Change the background color to white:
  - Command: set bg color white
- 9. Adjust the view as desired
- 10. Save the image:
  - Command: copy png file ~/Desktop/myfile.png

DNA helix with bound netropsin