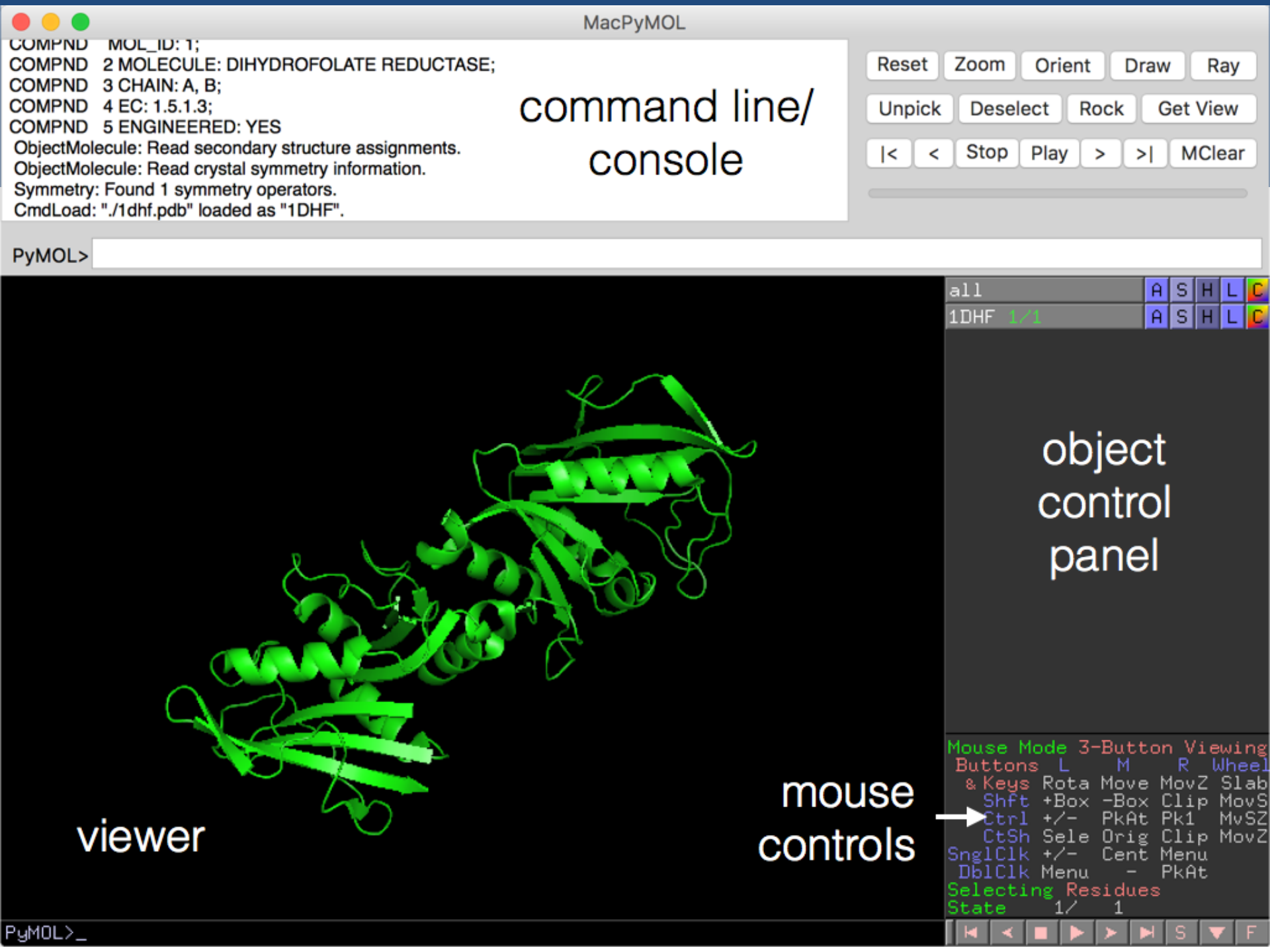


Protein structure and PyMOL scripting



Object control panel



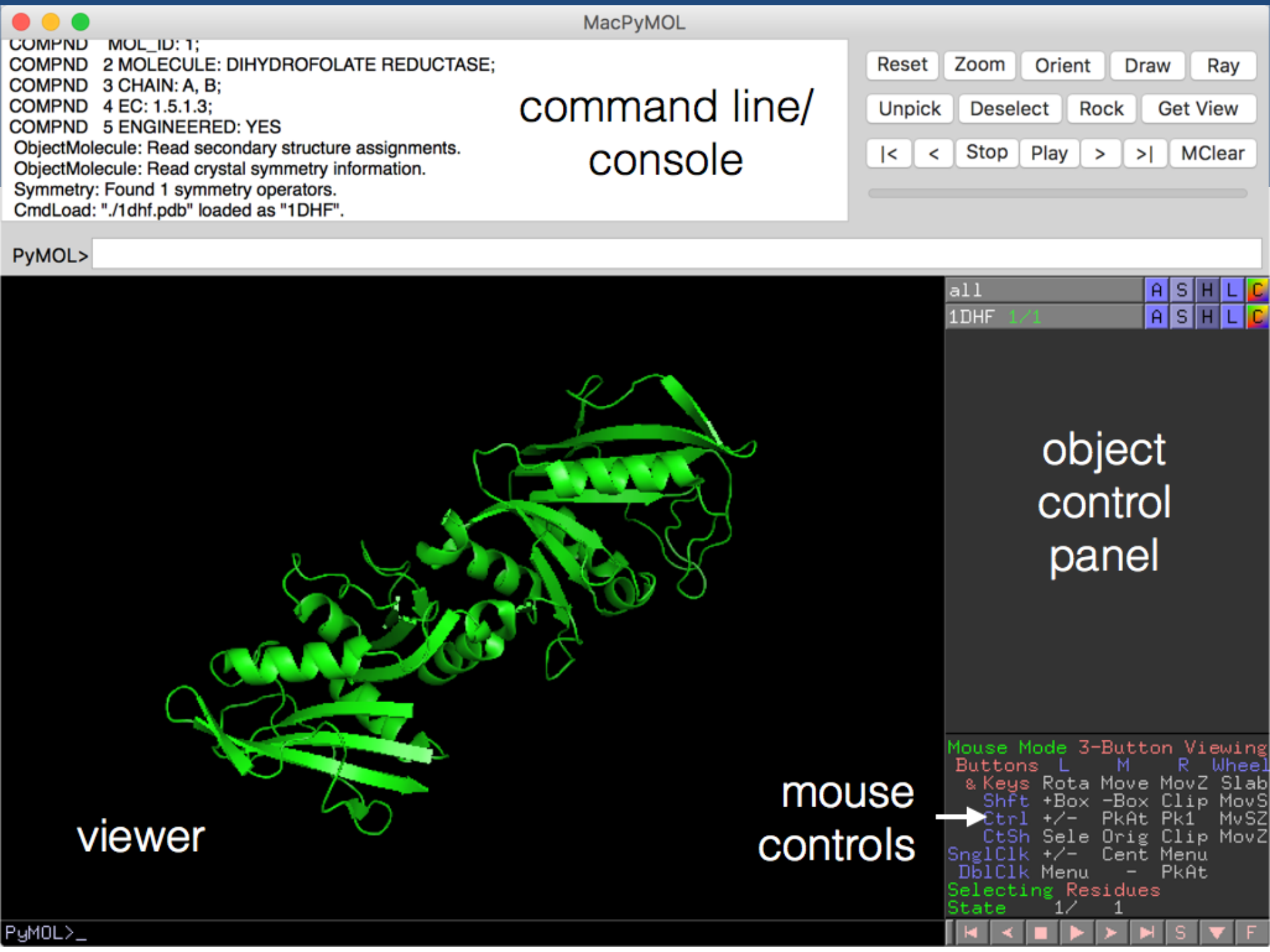
A (Actions): Rename, duplicate, remove

S (Show): Change the way things appear, e.g. change to stick or cartoon view.

H (Hide): Things that are shown using **S** accumulate, and don't automatically replace the last view. **H** is the opposite of **S** and hides unwanted representations.

L (Label): Label atoms, residues, etc.

C (Color): Change the color of atoms and groups



command line/
console

object
control
panel

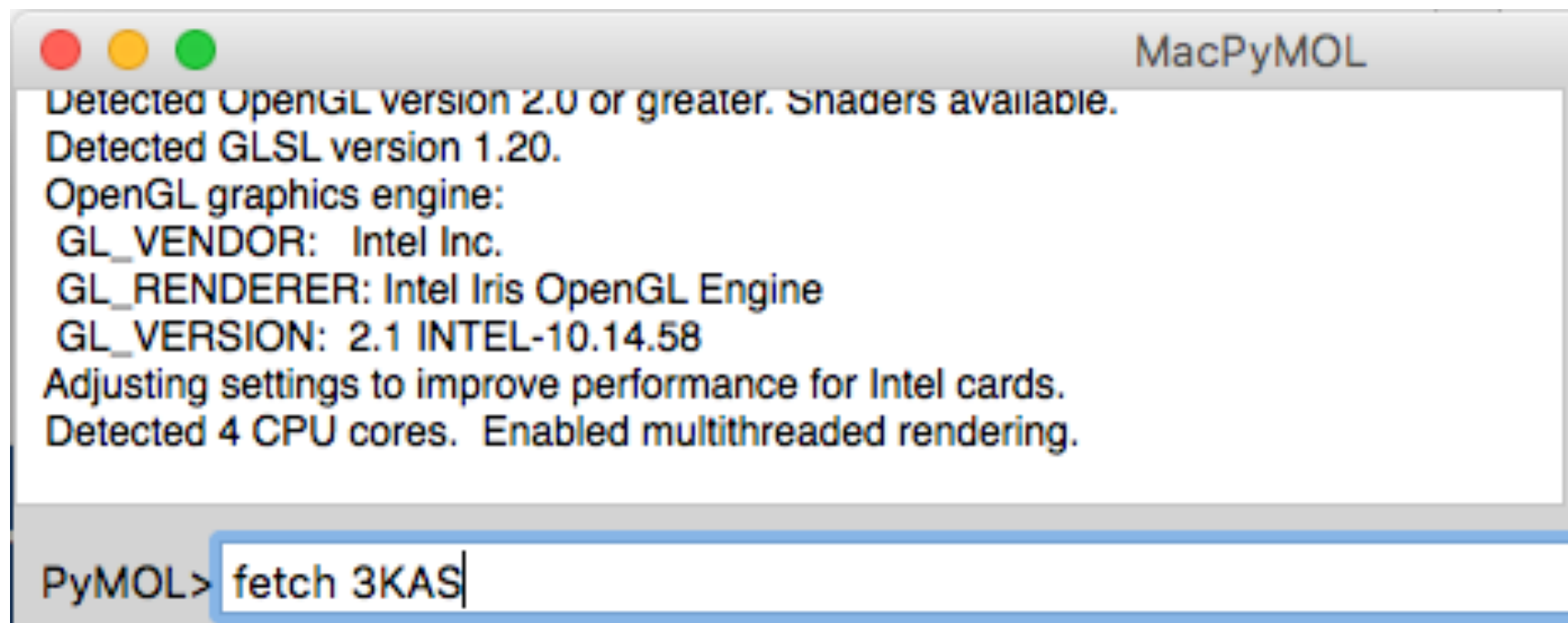
viewer

mouse
controls

Anything you can do with a menu, you can do with a command

`command arg1, arg2, arg3`

`>fetch 3KAS`



Anything you can do with a menu, you can do with a command

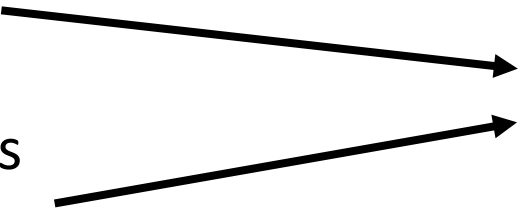
`select name, selector`

select chain A

`>select chain_a, chain A`

select all tyrosines

`>select all_tyr, resn TYR`



all	A	S	H	L	C
12AS 1/1	A	S	H	L	C
(chain_a)	A	S	H	L	C
(all_tyr)	A	S	H	L	C

More on selector syntax

http://pymolwiki.org/index.php/Property_Selectors

Anything you can do with a menu, you can do with a command

hide everything

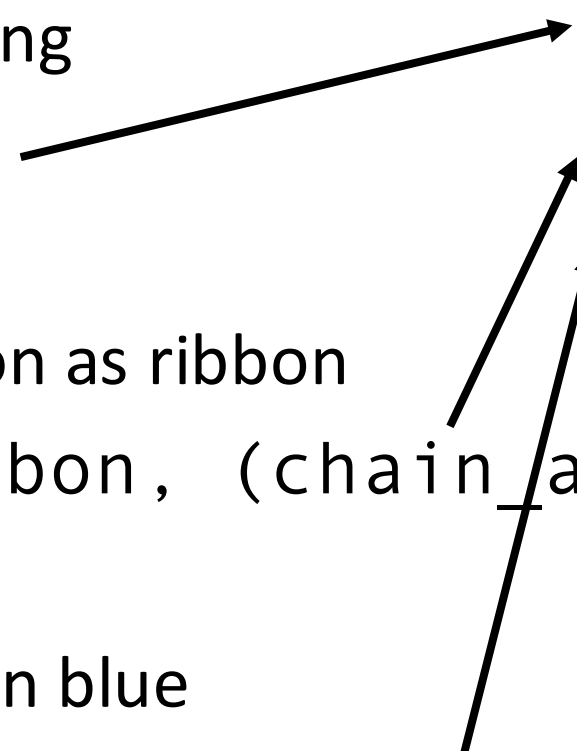
>hide all

show selection as ribbon

>show ribbon, (chain_a)

color selection blue

>color blue, (all_tyr)



all	A	S	H	L	C
12AS 1/1	A	S	H	L	C
(chain_a)	A	S	H	L	C
(all_tyr)	A	S	H	L	C

PyMOL exercises

1. Download & open structure 3KAS:
`fetch 3KAS`
 - Display in various forms (cartoon, stick, spheres, ...)
 - Color different chains
2. Download & open structure 1DLW
 - Display as cartoon
 - Show heme as sticks

Every PyMOL command has a python counterpart

PyMOL: >select chain_a, chain A

Python: `cmd.select("chain_a", "chain A")`

Need help? (PyMOL command line)

>help select

PYMOL API

`cmd.select(string name, string selection)`

You can navigate through your file system with the PyMOL console

Show the current working directory:

```
>pwd
```

(print working directory)

List all files and directories in the current working directory:

```
>ls
```

You can navigate through your file system with the PyMOL console

Change directories

```
>cd directory_name
```

Navigate to a directory called downloads

```
>cd Downloads
```

Navigate to a parent directory

```
>cd ..
```

PyMOL Exercises

- Use the PyMOL console to navigate to your Downloads folder
- Use the PyMOL console to navigate to your Desktop

An aside about text editors

- Text editors edit plain or raw text (no bold, italic, underline, etc.)
- Microsoft Word is **not** a text editor
- Windows: Notepad
- Mac OSX: TextEdit (Format > Make plain text)
- Cross-platform: Atom
 - www.atom.io

A simple python script to view a protein structure

- Open a new text file
- Add the contents:

```
cmd.fetch("3KAS")
```
- Save as "myscript.py" (or any name you want)
 - Text files can have any extension, but for python scripts we use ".py" by convention
- Using the PyMOL console, navigate to the directory where you saved your script
- In the PyMOL console:

```
>run myscript.py
```

Every PyMOL command has a python counterpart

PyMOL: >select chain_a, chain A

Python: `cmd.select("chain_a", "chain A")`

Need help? (PyMOL command line)

>help select

PYMOL API

`cmd.select(string name, string selection)`

PyMOL exercises

Write a python script to do the following:

1. Download & open structure 3KAS:
`fetch 3KAS`
 - Display in various forms (cartoon, stick, spheres, ...)
 - Color different chains
2. Download & open structure 1DLW
 - Display as cartoon
 - Show heme as sticks