# An Introduction to the UNIX Shell

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June 12, 2017

### 1 What is the shell?

The shell, or command line interface (CLI), is one of the most fundamental ways to interact with a computer. Rather than using a mouse to click on icons or windows, the shell relies entirely on typing commands. At a low level, the shell allows the user to access the hardware at the heart of the machine. Part of what makes the shell so powerful is its ability to run other commands, which in turn can accomplish complicated tasks or control things automatically. By the end of this tutorial, you will be able to navigate your system using the command line, as well as run other programs. You'll also be well on your way to understanding how to write your own programs for solving problems. Let's get started!

These notes are based on the lessons presented by the Software Carpentry website, http://swcarpentry.github.io/shell-novice/. There are many other valuable lessons there, beyond just the shell. Definitely check it out!

# 2 Navigating using the shell

An important first question when learning to use the shell is "How do I get around my filesystem?" In some sense, the command line gives you a text version of the directory structure that you might be used to navigating graphically, such as with Windows Explorer or macOS Finder.

### 2.1 UNIX Filesystem Structure

UNIX (and its descendents) use a hierarchical file system, which means that there is a root directory (denoted as /), with files and subdirectories that are located within that directory. Those subdirectories can in turn contain files and sub-subdirectories of their own, to help organize the contents of the filesystem. Each file and directory "knows" which directory it is located in, so that you can eventually move up the directory structure (sometimes called a "tree") and end up back at the root directory /.

One of the benefits of the hierarchical structure is that each file or directory has a unique name associated with it. Subdirectories are denoted by a / character between its name and

that of its parent directory. For instance, your "home" directory, which is where Ubuntu defaults to putting your personal files and downloaded files, is called "champ1", which is located in the home directory, which is located in the root directory /. The folder's location in the filesystem can be represented as:

#### /home/champ1

This directory is frequently used when writing commands, and so the tilde character ~ can be used to substitute for it.

### 2.2 Our first commands: 1s, cd, and pwd

#### 2.2.1 ls

Now that we know how the filesystem is laid out, we can begin using the command line to move around. Let's begin by opening a new terminal. You should see something that looks like this:

\$

This dollar sign is known as the "command prompt". This symbol means the shell is listening for your input, and is ready to run a command.

The first thing we want to do is to see what is in the current directory. Let's find out! The command for this is 1s (the "listing" of the directory's contents). To run the command, we type the characters 1s, followed by the Enter key. You should see something like this:

\$ ls
Desktop Documents Downloads Pictures Videos pic.png
\$

The first line (ls) is the command we entered, and the next line (Documents, Downloads, etc.) is the output of that command. On the following line is the command prompt again, meaning the shell has finished executing the command we asked it to run and is listening for the next command.

Commands will often have different options available for us to use, which are usually entered by entering a dash and a character after the command name. Let's look at 1s as an example. When listing directory contents, we can have the computer tell us which entries are directories with a trailing slash by using the -F option. Let's try it:

# \$ ls -F Desktop/ Documents/ Downloads/ Pictures/ Videos/ pic.png

Now we can see that every entry in our home directory is another directory, except for one. We will soon learn how to move between directories using another command.

As a side note, you can see many of a command's options by using the --help option. Alternatively, you can look at a command's manual page by running man <command>, e.g.,

man 1s. This will bring up a separate environment, where you use the space bar to move forward and b to move back. You can quit using the q key. The man page has more information than you ever cared to know about a specific command, so if you have specific questions, it's a great place to start.

We can also provide "arguments" to commands, which change how they behave. For the ls command, if we give it an argument, we will see a list of that directory's contents instead of the one we're currently in. Let's try it with the Desktop folder:

```
$ ls -F Desktop
data-shell/
```

We can now use ls to look at the contents of this sub-directory:

```
$ 1s -F Desktop/data-shell
creatures/ molecules/ notes.txt solar.pdf
data/ north-pacific-gyre/ pizza.cfg writing/
```

We can see that there are more directories and files in this directory. Now we're going to learn how to change our directory so we're in this one.

#### 2.2.2 cd

How do we move between directories? We need a new command, called cd (for change directory). Let's move to the data-shell directory we saw before:

```
$ ls
Desktop Documents Downloads Pictures Videos pic.png
$ cd Desktop
$ cd data-shell
$ cd data
$ ls -F
amino-acids.txt elements/ pdb/ salmon.txt
animals.txt morse.txt planets.txt sunspot.txt
```

The cd command didn't produce any output, but if we run ls after the cd command, we see that we have several new entries.

#### 2.2.3 Aside: pwd

When we're moving around the file structure, it's easy to get lost and forget where you are. The pwd command will pring the working directory. Let's try it:

```
$ pwd
/home/champ1/Desktop/data-shell/data
```

We now know where we are, which is very helpful for when we've moved around and forgotten where exactly we're located. Now, back to cd.

#### 2.2.4 Back to cd

So now that we've learned how to move down in the directory hierarchy, let's find out how to move back up. One thing you might try is cd data-shell, the name of the parent directory. Let's see what happens:

```
$ cd data-shell
cd: no such file or directory: data-shell
$ pwd
/home/champ1/Desktop/data-shell/data
```

cd gave us an error message, and we didn't actually move anywhere! What happened? The command cd is only able to move to directories that are contained in the current one. Fortunately in UNIX, every folder "knows" what directory it's in. Let's see how:

```
$ cd ..
$ pwd
/home/champ1/Desktop/data-shell
```

.. means "this directory's parent directory". But wait! We said that cd can only move to directories contained in the current one! We can see that it actually is in the current directory, it's just "hidden". Let's try 1s with the -a option:

```
$ ls -F -a
./ creatures/ notes.txt
../ data/ pizza.cfg
.bash_profile molecules/ solar.pdf
Desktop/ north-pacific-gyre/ writing/
```

The . and . . entries are special, and are found in every directory. . means "the current directory", and we'll see some uses for it soon. We also see the . . entry, the parent directory. We also see a new file we didn't before, .bash\_profile. By default, ls doesn't show any files or directories that begin with a . character. These are usually configuration files the user does not usually have reason to modify, so they are hidden to prevent cluttered directories.

As a couple of last points on cd, let's see what happens if we run cd without any commands:

```
$ cd
$ pwd
/home/champ1
```

We were brougt back to our home directory. We can also use the ~ character to represent our home directory:

```
$ cd ~/Desktop/data-shell
$ ls -F
creatures/ molecules/ notes.txt solar.pdf
data/ north-pacific-gyre/ pizza.cfg writing/
```

We're back in the data-shell directory.

#### 2.2.5 Aside: Absolute vs. Relative Paths

When we want to specify the path to a file or directory in UNIX, there are two options: absolute paths and relative paths. Absolute paths contain the entire path, starting from the root directory /. These paths work no matter which directory we're located in, since they tell the filesystem how to get somewhere starting from the root directory.

Conversely, we've seen that we can reference a directory's parent by using the .. alias. Let's suppose that we're in ~/Documents. If we wanted to get to ~/Desktop, we could use the command:

\$ cd /home/champ1/Desktop

OR

\$ cd ../Desktop

In both cases, we'd end up in the same folder.

#### 2.2.6 Aside: tab-completion

Are you tired of typing out the entire name of a file or directory yet? Or making typos? (This is a recurring problem for me...) Enter tab-completion, which is one of the best features of the shell.

Tab-completion works by typing the first few letters of a command or argument, and then hitting the <TAB> key. If the shell knows what you're trying to say, it will automatically fill in the rest for you! If it doesn't know exactly what you want (like if there are several different things that match the first few letters you've typed), then hitting <TAB> a second time will print out to the screen all of the possible completions for your partial command or argument. So, if you type cd ~/Desk<TAB>, the shell will auto-complete to cd ~/Desktop. On the other hand, typing cd ~/D<TAB> will not auto-complete, since this could turn into ~/Desktop, ~/Documents, or ~/Downloads. Hitting <TAB> a second time will show you these options, so you can see for yourself why the shell isn't completing the command.

Congrats! You now know how to move around to different directories. This an important first step on the road to mastering the shell.

### 3 Files and Directories

### 3.1 Making New Directories

It's helpful to know how to move around existing directories, but we also want to be able to make our own new ones. For this, we have the mkdir command, which is short for "make directory". Let's do that now:

```
$ cd ~/Desktop/data-shell
```

\$ 1s -F

```
molecules/
                                                           solar.pdf
creatures/
                                        notes.txt
data/
                   north-pacific-gyre/ pizza.cfg
                                                           writing/
$ mkdir thesis
$ 1s -F
                                     thesis/
creatures/
             north-pacific-gyre/
             notes.txt
                                     writing/
data/
Desktop/
             pizza.cfg
molecules/
             solar.pdf
```

We see that we have a new thesis directory. Great!

### 3.1.1 Naming Files and Directories, a.k.a., Spaces are Evil

When we're naming our files and directories, there are some important rules that we should follow:

- 1. Don't use spaces in names. Use \_ (underscore) or (dash) instead. Whitespace in directory names is frequently garbled by the shell unless you're careful, since it assumes a space means "move on to the next argument".
- 2. Don't begin names with -, since the shell interprets leading dashes as command arguments.
- 3. Try to use only letters, numbers, . (period), \_ (underscore), and (dash). We're going to learn soon that most punctuation marks have special meaning in the shell.
- 4. Don't use spaces in names.

### 3.2 Making New Files

Now that we've made a new directory, we want to put something in it. We're going to start a new text file. To do that, we're going to use a program called nano. Let's make a new file called draft.txt:

```
$ cd thesis
$ nano draft.txt
```

This opens up nano, which allows you to write plain text files. We're going to use it (or another text editor) to write our programs in later lessons, so it's good to get familiar with it. You can type the contents of the file into nano directly. You'll notice at the bottom of the screen, there are commands listed. The combination of  $^0$ , for instance, means to hold down the Ctrl key while pressing 0. Let's write some text, save it with  $^0$ , and exit with  $^X$ .

Let's look at our directory now:

```
$ ls
draft.txt
```

We didn't like that draft, so we're going to start over and try again. We get rid of the current file using the rm command:

```
$ rm draft.txt
$ ls
```

There's no output from ls anymore, because we've deleted the only file that was in this directory. Let's get rid of the directory too:

```
$ cd ..
$ rm thesis
rm: cannot remove 'thesis': Is a directory
```

The rm command by default cannot remove directories. There is a command rmdir to do that, or we can pass an option to rm:

```
$ rm -r thesis
```

This will get rid of the directory, and all of its contents. (The -r option instructs rm to recurse into all subdirectories.)

### 3.2.1 Aside: Always Have the Safety On

One very important aspect of rm is that once deleted, the files are gone forever. There is no "trash can" or "recycle bin" on the command line: one a file has been rm'd, there is no way to get it back. To get around this, we can use the rm -i command, which will interactively ask you about every file, and whether you really want to delete it. This gives you an extra layer of security, and keeps you from accidentally deleting all of your code you worked so hard to write.

# 3.3 cp and mv: Moving Things Around

The cp command is very straightforward: it copies files (or directories) from one place to another. The general syntax is cp existing\_file new\_file. The contents of existing\_file will be copied to new\_file as an exact copy. This gives us 2 versions of the file, and can be useful for making a backup copy of code or data files.

The mv command is also straightforward: it uses the same syntax as the cp command, but deletes the original copy. So, if you run mv existing\_file new\_file, you'll notice that there's only one file, new\_file, when you're done. mv can be used to move files from one directory to another, or to rename files and directories (e.g., using mv old\_name new\_name).

Let's use this in practice to get one of the data directories off of the desktop folder, and into our Documents folder:

```
$ cd ~/Desktop/data-shell
$ mv north-pacific-gyre ~/Documents
```

We also could have given a relative path, mv north-pacific-gyre ../../Documents. Note that mv will not produce output unless the -v option is given to it. This has moved the data from where it was to a new location. All of the files contained in this directory are still inside of it, just at a new point in the filesystem.

# 4 Pipes and Filters

One of the most powerful features of UNIX is that you can very easily use the output of one command as the input for another. This allows us to chain together commands to do powerful tasks with just a few keystrokes. Let's see how these behave.

#### 4.1 Wildcards

We're going to use a new command, wc, which will give us a word count of a file that we pass as an argument. Let's go to the molecules folder:

```
$ cd ~/Desktop/data-shell/data/molecules
$ ls
cubane.pdb ethane.pdb methane.pdb
octane.pdb pentane.pdb propane.pdb
```

#### 4.1.1 The \* wildcard

We have six files here. Rather than run the wc command on each individually, we're going to use what's known as a "wildcard" character. The \* symbol tells the shell to match zero or more characters. For instance, let's use it to run wc on all files that end in .pdb:

```
$ wc *.pdb
20  156 1158 cubane.pdb
12  84  622 ethane.pdb
9  57  422 methane.pdb
30  246 1828 octane.pdb
21  165 1226 pentane.pdb
15  111  825 propane.pdb
107  819 6081 total
```

By reading the man page of wc, we would see that the three columns are the number of lines, words, and characters contained in each file. The total of all files inspected is the bottom row.

#### 4.1.2 The? wildcard

There is another commonly used wildcard, the ? character, which is similar to the \* character. Instead of matching any number of characters, the ? character matches exactly one character.

So if we ran wc \*ethane.pdb, we'd get word counts for both ethane.pdb and methane.pdb. On the other hand, if we ran wc ?ethane.pdb, we would only get methane.pdb, since the ? character must match at least one character.

### 4.2 Redirecting Output

Let's return to the wc command. The option -1 will give us just the number of lines in a file (the first column of the full wc command):

```
$ wc -1 *.pdb
20 cubane.pdb
12 ethane.pdb
9 methane.pdb
30 octane.pdb
21 pentane.pdb
15 propane.pdb
107 total
```

This output is manageable, since there are only six files. What if we had thousands? We can redirect the output of the wc command to a file, so that we can read it more easily. To do this, we use the > (greater than) symbol:

```
$ wc -l *.pdb > lengths.txt
```

Note that there's no output from this command, since it was sent to lengths.txt. Let's look at our directory contents again:

```
$ ls
cubane.pdb ethane.pdb lengths.txt methane.pdb
octane.pdb pentane.pdb propane.pdb
```

We have a new file, lengths.txt, that has shown up. Now let's take a look at what it contains.

#### 4.2.1 cat and less: Seeing File Contents

To see the contents of a file, we can use cat or less. cat will concatenate the file to the screen, and prints the whole file. less will show you less of the file—only one screen's worth—and is much easier for navigating large files. Like man pages, you can move through the less screen by using space bar to go forward a page, b to go back, and q to quit. Since we're looking at just a small file, we're going to use cat.

In general, though, less is more!

Okay, now let's look at the contents of lengths.txt:

```
$ cat lengths.txt
20  cubane.pdb
12  ethane.pdb
9  methane.pdb
30  octane.pdb
21  pentane.pdb
15  propane.pdb
107  total
```

Notice that this is the same as the output from wc -1 \*.pdb from before, without the redirect.

### 4.3 Sorting Output

Oftentimes, we want to be able to sort our output. Let's find out how to sort our lengths.txt file to figure out the longest file:

```
$ sort -n lengths.txt
9 methane.pdb
12 ethane.pdb
15 propane.pdb
20 cubane.pdb
21 pentane.pdb
30 octane.pdb
107 total
```

We can see that methane.pdb is the shortest file, and octane.pdb is the longest one. We can make a new file that has the lengths sorted. We also want to use the head command, to see just the top line of the resulting file:

```
$ sort -n lengths.txt > sorted-lengths.txt
$ head -n 1 sorted-lengths.txt
9 methane.pdb
```

Giving the head command the option -n 1 means that we see only the first line of the file.

# 4.4 Pipes

Wouldn't it be convenient if there were a way to avoid making a temporary file? We can accomplish this by using the pipe character | bewteen two commands to tell the shell "the output of the first command should become the input for the second command." We can get the same output result as above, but without having to make the sorted-lengths.txt file:

```
$ rm sorted-lengths.txt
$ sort -n lengths.txt | head -n 1
9 methane.pdb
```

If we were to ls in the directory, we would see that we didn't create a sorted-lengths.txt file this time. We can even chain multiple commands together, so we wouldn't even need a lengths.txt file!

```
$ wc -l *.pdb | sort -n | head -n 1
9 methane.pdb
```

Ta-da! No temporary files!

This paradigm is what makes UNIX so powerful and so successful: make a tool, that does one thing *really* well, and then use pipes to redirect output between tools, which act as filters to massage the original output into our desired format.

### 4.5 Putting it all together

Now, let's use what we learned on a small scale to look at (a facsimile of) real data. We're gong to go to the north-pacific-gyre folder:

```
$ cd ~/Documents/north-pacific-gyre

$ wc -1 *.txt | sort -n | head -n 5

240 NENEO2018B.txt

300 NENE01729A.txt

300 NENE01729B.txt

300 NENE01736A.txt

300 NENEO1751A.txt
```

As we saw above, this will give us the number of lines of all files that end in .txt, sort them, and then give us the top 5 entries (in this case, the 5 shortest files). We can see that one file is significantly shorter than the rest of them.

Let's also take a look at the 5 longest ones. This is done with the tail -n 5 command:

```
$ wc -1 *.txt | sort -n | tail -n 5
300 NENE02040B.txt
300 NENE02040Z.txt
300 NENE02043A.txt
300 NENE02043B.txt
5040 total
```

We've gotten some glimpses into why the command line is so powerful, and we've only used a few commands!

# 5 Loops

Loops are one of the most basic constructs in all of programming. Essentially, a loop says "I want to repeat the same set of actions many times, with only small changes each time." As control structures, they are clear to both humans and machines, and are very efficient at automating large tasks. Automation is one of the major benefits of programming in the first place, so we're going to take our first step toward how to do it in the shell.

### 5.1 For Loops

A "for loop" is a construct where we perform some task *for* a certain number of pre-defined iterations. To see this in action, let's go to another directory. Our first task is to make a backup of each of these files. Let's use cp for this:

```
$ cd ~/Desktop/data-shell/data/creatures
$ cp *.dat original-*.dat
cp: target 'original-*.dat' is not a directory
```

What happened? When we give cp multiple arguments, it expects the final one to be a directory to copy all of the files to. In this case, the first wildcard is expanded, so the full command reads cp basilisk.dat unicorn.dat original-\*.dat. Since there is no directory with that name, cp gives us an error.

We can use a loop to accomplish this for us. Enter the following commands:

```
$ for filename in *.dat
> do
> cp $filename original-$filename
> done
```

Note that the prompt has changed from the dollar sign to the "greater than" sign. This means that the shell is waiting for us to finish the for loop, and will change back once the loop has been completed.

Now we have achieved the original intent: we've made a backup of each of our files. Implicitly, in this construction, we have told the shell that it should do the same thing to two different files: for each file in our list, we want to make a copy, with original-prepended to the filename.

When we're building a loop, sometimes it's helpful to print the command you *actually* wrote, and see if it's the same as the one that you *wanted* to write. To do this, we use the echo command:

```
$ for filename in *.dat
> do
> echo "cp $filename original-$filename"
> done
cp basilisk.dat original-basilisk.dat
cp unicorn.dat original-unicorn.dat
```

Now we can see that the commands do what we expect them to do, and we can remove the echo and quotes to actually run the command. We've already done this, so we won't do it again. Just know that echoing the body of the loop is helpful for debugging potential issues.

# 6 Scripts

We've finally arrived at the teleological conclusion of the shell: scripts to do the business of repetitive tasks for us. You might have noticed that writing for loops by hand is repetitive and error-prone. It's also difficult to change a few small things and run the same commands again. We will now learn about writing scripts.

To start with, we're going to use nano again. Let's write a script that will take lines 11-15 of the octane.pdb files. Let's go back to the molecules folder.

```
$ cd ~/Desktop/data-shell/data/molecules
```

\$ nano middle.sh

This makes a new file called middle.sh. In nano, let's write the following line:

We can save with ^O and exit with ^X. Let's run it now, using the bash command:

\$ bash	middl	.e.sh						
MOTA	9	H	1	-4.502	0.681	0.785	1.00	0.00
MOTA	10	H	1	-5.254	-0.243	-0.537	1.00	0.00
MOTA	11	H	1	-4.357	1.252	-0.895	1.00	0.00
MOTA	12	Н	1	-3.009	-0.741	-1.467	1.00	0.00
MOTA	13	Η	1	-3.172	-1.337	0.206	1.00	0.00

Excellent! Now we can make changes to the shell script, and rerun it easily. It's not that flexible though: we can only run this on octance.pdb, and it gives us lines 11-15. Let's make some modifications to make it more flexible.

The flexibility can be achieved by using "positional arguments" which are represented by numbers. When we run a command, the variable \$1 represents the first argument passed in, \$2 is the second one, and so on. Let's open the file with nano again, and change the line to read:

```
head -n 15 "$1" | tail -n 5
```

We've protected the first argument by placing it in double quotes, which will allow for filenames with evil spaces to be passed in without generating an error. Now let's run middle.sh again, with a different file:

\$ bash	middl	e.sh	pentane.pdb					
MOTA	9	H	1	1.324	0.350	-1.332	1.00	0.00
MOTA	10	H	1	1.271	1.378	0.122	1.00	0.00
MOTA	11	Η	1	-0.074	-0.384	1.288	1.00	0.00
MOTA	12	H	1	-0.048	-1.362	-0.205	1.00	0.00
MOTA	13	Н	1	-1.183	0.500	-1.412	1.00	0.00

Great! We've added more flexibility. Let's make it even more flexible, and also allow the user to specify the numbers that get passed into head and tail:

```
head -n "$2" "$1" | tail -n "$3"
```

We can recover the same behavior as above using a new command:

\$ bash	middl	e.sh	pentane.pdb 1	15 5				
ATOM	9	H	1	1.324	0.350	-1.332	1.00	0.00
ATOM	10	H	1	1.271	1.378	0.122	1.00	0.00
ATOM	11	H	1	-0.074	-0.384	1.288	1.00	0.00
ATOM	12	H	1	-0.048	-1.362	-0.205	1.00	0.00
ATOM	13	H	1	-1.183	0.500	-1.412	1.00	0.00

At the same time, we can look at lines 16-20 instead:

```
$ bash middle.sh pentane.pdb 20 5
                                    -1.259
                                                                      0.00
MOTA
          14
              Η
                            1
                                              1.420
                                                       0.112
                                                               1.00
                            1
                                    -2.608
                                             -0.407
MOTA
          15
              Η
                                                       1.130
                                                               1.00
                                                                      0.00
MOTA
                            1
                                    -2.540
                                             -1.303
                                                      -0.404
          16
              Η
                                                               1.00
                                                                      0.00
MOTA
                            1
                                    -3.393
                                              0.254
                                                      -0.321
          17
              Η
                                                               1.00
                                                                      0.00
TER
          18
                            1
```

As we're going to learn, it's very helpful to document what was on your mind while you were writing the script, to help other programmers (or yourself!) to more easily understand what a program is doing. To do this, we'll add a few lines to the top of the script that start with the # character, which causes the shell to ignore the contents of that line. We'll open nano again, and run it:

```
# Select lines from the middle of a file.
# Usage: bash middle.sh filename end_line num_lines
head -n "$2" "$1" | tail -n "$3"
```

Great! We've successfully written our first shell script! You are now ready to go out and conquer the world by automating all of your tasks and running complicated code with a few keystrokes!

# 7 ssh and scp: How to Work on a Supercomputer

As you're going to see in the course of this bootcamp, most of the work that we do is too much for a single laptop or desktop computer to handle. Many times, we log into a different computer, typically a supercomputer, to help us with our most computationally challenging tasks. This involves getting an account on the machine, which has a username and password. Then, logging in is as simple as running a few commands on the command line.

### 7.1 ssh: Logging In

The most common tool for accessing your account on a different machine is **ssh**, which is a **secure shell**. In addition to your username and password, you'll need to know the remote computer's *hostname*, which is how you'll tell your machine which computer to connect to. The general structure for a command is:

#### \$ ssh <user>@<hostname>

At this point, the remote machine will ask for your password.

Let's suppose that your username is champ1, and the computer we'd like to connect to is the Folio computer at UPenn's campus. For this machine, the hostname is folio2.sas.upenn.edu. So, the ssh command is:

```
$ ssh champ1@folio2.sas.upenn.edu champ1@folio2.sas.upenn.edu's password:
```

At this point, you would enter your password. Note that the cursor does not move, to keep people looking over your shoulder from knowing how long your password is. Once you've entered it, the machine will grant you access, and give you a new command prompt. This time, you're running a shell remotely on the new machine. Exciting! From here, you can access data that's on the other machine, as well as run programs. We'll talk more about how to do these things in future lessons.

### 7.2 scp: Bringing Data Home

Let's assume that you've written a great script, do\_cool\_science.sh, and generated new data or a plot that you want to have on your personal machine. ssh is only a way to execute commands remotely, not a way to move files. Fortunately, there's a command scp to securely cp files over the network. It uses the same security as ssh to make sure other people can't eavesdrop on your connection, and is very simple for moving files to or from the remote machine.

From your machine, you can either "push" files to the remote computer, or "pull" them from the remote machine to yours. Unless you've set up your personal computer's connection in a special way, it's much harder to push and pull data from the remote machine to your personal one. So you should always run the scp command from your personal machine.

The typical scp command for pushing to the remote machine is:

\$ scp <local\_data> <user>@<hostname>:<remote\_destination\_folder>

For instance, let's say that I have a file on my machine called awesome\_data.txt, and on the remote machine I want to copy it to ~/Documents/data\_backup. The command I would run is:

\$ scp awesome\_data.txt champ1@folio2.sas.upenn.edu:~/Documents/data\_backup champ1@folio2.sas.upenn.edu's password:

After you enter your password, scp will begin transferring the file. A copy is going to end up on folio2.sas.upenn.edu, in the ~/Documents/data\_backup folder, just like we wanted. scp will show you the status of the data transfer while it's happening, with information like how much of the file has been transferred so far, what the current speed is, and how long the transfer has been going on for. Once it's done, it will display average values for the whole transfer.

For pulling data from the super computer to your local machine, we use a similar syntax:

\$ scp champ1@folio2.sas.upenn.edu:~/Documents/data\_backup/old\_data.txt .

Once again, it will ask for our password. Note that we've used the shortcut . to represent the current directory. This means that a copy of the file old\_data.txt will end up in the directory we're currently in.

Congratulations! You've taken your first steps toward mastering using the shell, one of the most powerful modern tools for science research and discovery. You're well on your way to becoming a data scientist, and gaining new insight on problems you wouldn't have otherwise. Don't be afraid to ask for help if you have problems, and the Internet is your friend if you have any questions!