# An Introduction to the Command Line Interface

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This is a preliminary draft – please send comments to charlierahal@gmail.com

# 1 Introduction and History

The command line interface ('CLI') can often feel like a harsh, unforgiving environment, but it represents an indispensable way to interface with the underlying operating system of a computer. In comparison to graphical user interfaces ('GUI'), CLIs can require substantial effort in order to understand and acquire the skills to execute specific tasks, but conversely, they represent an almost infinite space of opportunity productive analysis which would otherwise remain largely impossible. If you are a Microsoft Windows user, you may have already encountered the command line through the 'Command Prompt' (cmd): the counterpart of COMMAND.COM in DOS and antiquated Windows 9x systems. If you are running Mac OS X or Linux, you are already operating a 'Unix-like' (hereafter '\*nix-like') system with access to command line utilities (through the 'terminal'), likely all neatly packaged behind an accessible graphical environment. To more concretely motivate our use of the command line in a general context, the following reasons are of specific importance:

• **Speed:** At the most primitive level, using a keyboard is faster than using a mouse.

For correspondence: Charles Rahal, Department of Sociology, University of Oxford, Manor Road Building, Manor Road, Oxford, OX1 3UQ. E-mail: charlie rahal at gmail dot com. Accompanying shell scripts for this chapter can be found at github.com/crahal, in addition to a set of slides to be delivered as a workshop. All text in typewriter font indicates a command or filepath.

- **Automation:** No matter how complex the command or series of commands, the CLI doubles as a scripting language which can be ran as a batch, or 'scheduled' to run at specific times in the future.
- Control: Commands are often more powerful and precise, giving more control over the operations to be performed, and this is particularly relevant with more complex tasks.
- **Consistency:** CLIs enable more 'replicability' either later by the original user or others, as all options and operations are invoked and documented in consistent form.
- **Resources:** A computer which relies only on the CLI takes a lot less of the computer's system resources than a GUI.

There are additional factors which convincingly motivate bio-informaticians and social scientists to integrate the CLI into their workflow. The reduced cost of genomic sequencing – due to the development of 'Next-Generation Sequencing' (NGS) machines – is rapidly generating enormous waves of data. Analysis of such large caches of genotype data typically requires hardware more powerful than a standard laptop or desktop computer, typically in the form of 'high performance computing' (HPC). These almost exclusively run Linux or Unix, and a large volume of genomic software is only compatible with Linux or Mac OS-X based systems (such as one of the two discussed in Section 6.3). That is not to say that you should leave behind your graphical environment entirely! Some tasks are without a doubt best suited to a GUI, such as word-processing. However, tasks such as data manipulation and file management – critical tasks in our domain – are more naturally suited to the command line.

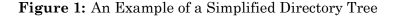
Before, we proceed, it might be useful to consider the history, and therefore the evolutionary distinction between Unix and other \*nix-like systems. Unix was created 1969 at Bell Laboratories – then a development division at AT&T – by Ken Thompson and Dennis Richie. Prior to this, all software was hardware dependent, written in assembler language. Unix was originally developed as a proprietary, commercial enterprise, with portability as a primary strength. Shortly after, researchers at the University of California, Berkley began developing Berkley Software Distribution (BSD) – based on the AT&T code base. This has since been rewritten to remove any copyright issues originating from the original Unix code. Commercial Unix derivatives which combined elements of Unix and BSD began appearing in the 1980s (such as AIX and Sun Solaris), but fragmentation began to cause problems. To address this issue, the Institute of Electrical and Electronics Engineers

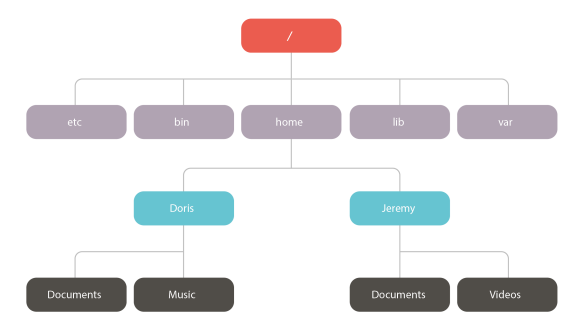
(IEEE) developed the Portable Operating System Interface for Computer Environments (POSIX). On September 17, 1991, Linus Torvalds first released the Linux operating system kernel. It shares much of the core concepts of Unix, but, because it contains no Unix code, it can be freely shared and replicated. Specifically, the term Linux refers to the kernel, and for this reason, members of the free and open-source community strongly advocate the term GNU/Linux when referring to the operating system in order to acknowledge and promote the fact that GNU was a longstanding project begun in 1984 to develop a free operating system, with Linux merely providing the 'missing piece'. This has since spun off into hundreds of 'distributions', each with specific functionality. Notable examples include the Android operating system for smartphones and Chrome OS which runs on Chromebooks. Variants also appear on multiple types of embedded devices such as televisions, video games consoles and smartwatches. It is also by far the most popular server operating system in use today. Therefore, we can summarize the classification of \*nix-like systems into one of the three main categories, a.) Unix: Commercial implementations based on the original AT&T code, b.) BSD: Open source system derived from the original Unix and c.) Linux: Open source clone of Unix, written from scratch and freely distributed.

## 2 An Overview of the Architecture

At the core of every \*nix-like operating system is the **kernel**, which provides a layer between the user applications and the hardware of the computer. For example, the kernel facilitates the printing of output to a command line (or visualising graphics in a GUI) on a monitor, and writes files to the disk drive. It directs all of the actions of an operating system to work in harmony. Most \*nix-like systems share a similar directory structure. The most common directories which we need to briefly itemize are listed below, with a basic summary visualization seen in Figure 1:

- /: otherwise known as 'root', this is the top of the file system hierarchy.
- /bin: Binaries and other executable programs.
- /etc: System configuration files.
- /home: Home directories.
- /opt: Optional/third party software.
- /tmp: Temporary space, typically cleared upon every reboot.





- · /usr: User related programs.
- /var: Variable data, such as log files.

User generated files are typically stored in the /home directory, and all files stored outside of this directory are only editable by root users. This protects core system processes either from malicious actors or accidental removal by the root user themselves. A variety of different types of file systems are supported, namely ext2-4, JFS, HFS+, UFS, VxFS and ZFS. All have a directory known as /dev which contains device files pertaining to hardware installed on the system. These devices can be previewed by typing 1s -1 /dev/ into the command line — with commands and options which we will come to explain later in this chapter.

This brings us to the **shell** – the command line interpreter which accepts the command line inputs and performs the required tasks. At this stage, it is critical to distinguish between the two types of shell prompts: \$ is the normal shell user, and # represents the root user. These types of user have significant implications with respect to what can and cannot be accessed - where the root typically has unrestricted access (with the potential for accidental damage!). Finally, it is important to be aware that while there are many types of shells available (such as sh, chs, tsch and ksh), the Bourne Again Shell (bash) is by far the most popular shell in use today (where scripts for different shells are typically not

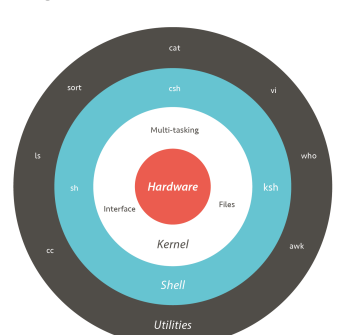


Figure 2: An Overview of the Architecture

portable). An overview of architecture can be seen in Figure 2.

We must also mention 'case sensitivity' where ExampleFile is not the same as examplefile and the fact that there is no recycle bin (as per Windows): when you delete something from the CLI in \*nix-like systems, the files are irrecoverable. One final thing to realize about \*nix-like systems is that the whole philosophy revolves around a small set of building blocks and programs which we can then 'pipe' together with sophisticated pattern matching and programming to create bigger things. For that reason, we start in Section 3 with a set of fundamental CLI commands. Just like mathematics, learning the CLI requires practice. If you have access to a Mac or Linux, you are likely acutely aware of the value of the terminal to your system, and you can follow along without issue. If you are using Windows, one option is to create a new partition and 'duel-boot' Linux alongside it, or 'live-boot' from a USB directly. Another way to get access to a terminal (if you are not on a machine which natively has one) is to run Linux (or another open source Unix variant) within a virtual machine - an emulation of a computer system ran inside your default operating system. One free tool of note is VirtualBox. A final option, which may only be accessible to some readers, is to access a remote machine through their workplace or university (through the ssh command). Without further ado, please allow me to welcome you to the command line:

#### Code Example 1: A Warm Welcome

```
user@system:~$ echo ''Welcome to the Command Line!''
Welcome to the Command Line!
```

# 3 Fundamental CLI Commands

All of the commands in the following section are summarized in the glossary at the end of this chapter. The first command which we saw in the previous section was echo: one of the most commonly used built-in commands which writes its arguments to standard output (the display screen by default). Another important echo to try is echo \$SHELL, which will print which shell we are using by default. The man command shows the manual for a given page, including information on options and usage. This is the default resource for getting help on specific commands. For example, the man echo command will display the manual for the echo command.

# Code Example 2: man

The whatis command gives a short summary description of the specific command, and command inputs can be stacked together. Each manual page has a short description available within it - and whatis searches the manual page names and displays the manual page descriptions of any name matched. For example:

# Code Example 3: whatis

Is is a Linux shell command (with some very subtle differences to dir) that lists directory contents of files and directories. There are a number of important examples however. Is / lists the root directory. Is .. lists the parent directory. Is ~ lists the users home directory. The lists can also be sorted, accept wildcards, and pipe outputs to file (see below). It can show hidden files, show file size, and has the option for a 'long' format.

#### Code Example 4: ls

```
user@system:~$ ls
```

2 Desktop Documents Downloads Music Pictures Public Videos

However, this isn't altogether much use without knowing *where* we are in the file tree. In \*nix-like (and some other operating systems), the pwd command (which stands for print working directory) writes the full pathname of the current working directory to the standard output.

#### Code Example 5: pwd

```
user@system:~$ pwd
```

2 /home/user

We can also change between the directories which we are presently in using the cd command (also known as chdir: change directory). If we carry on the example from above, we can then move from our user folder (/home/user/) to the Documents folder (/home/user/Documents). Note the change in the command prompt to indicate the change of location (on some systems):

#### Code Example 6: cd

```
user@system:~$ cd Documents
user@system:~/Documents$ pwd
/home/user/Documents
user@system:~$ cd ..
user@system:~$ pwd
/home/user/
```

We should make a new directory (or folder - a container for other files) for the purposes of following along with these examples. This is done with the mkdir command:

# Code Example 7: mkdir

```
user@system:~$ mkdir nix
user@system:~$ ls
nixTextbookChapter
user@system:~$ cd nix
user@system:~/nix$ pwd
/home/user/nix
```

This brings us to absolute and relative paths. From the documents folder above, we can navigate to the new folder (nix) through its relative path: cd nix (when we are already in /home/user/, or through the absolute path with cd /home/user/nix. There are three useful shortcuts in pathing to be aware of: ~ is the shortcut for your home directory, . is the reference to your current directory (i.e. cd ./nix) and .., which is a reference to the parent directory. There are multiple ways of identifying the same file path in the directory tree.

The touch command is a standard program for Unix/Linux operating systems which is used to create, change and modify timestamps of a file.

# Code Example 8: touch

```
user@system:~/nix$ touch testfile
user@system:~/nix$ ls
testfile
```

Despite the lack of detail surrounding it, the file testfile might seem unfamiliar to you for one obvious reason: the lack of an extension. That is because we are now operating within an extensionless environment, where a file extension is normally a small set of characters after a full stop which denote which type of file it is (such as file.exe for executables, .txt for text files, and so on). In operating systems such as Windows, these are critical, as the system uses this information in order to determine how to run the file. In \*nix-like systems, the extension is ignored, and the file type is determined automatically (and the command file can tell us additional information, if required).

Now that we've discussed how to create directories and files, we should also talk about the daunting rm command, which removes specified *files* (by default, it does not remove directories). To remove the file we just created above, we can use the command rm testfile. There are two key options with the rm command to be aware of. -i (which is the short hand, where the long hand is --interactive=always) creates a prompt before every removal (and -I or --interactive=always creates a prompt before removing more than three files, or when removing recursively). This is the default behavior within some distributions (through bash aliases: i.e. alias rm='rm -i'). If the -r (--recursive) option is specified, the command will remove directories and their contents recursively. For example, we could *recursively* delete the contents of the directory which we created above.

#### Code Example 9: rm

```
user@system:~/nix$ cd ..
user@system:~/$ rm -ri nix
rm: remove directory 'nix'?
```

This not only introduces us to command [OPTION]s (of the form command [OPTION]... FILE...), but it also shows how we can stack options together, where rm -ri is the equivalent to rm -r -i. Note, that command line options are not universal between commands, and implementation across different operating systems may vary. Note how our use of cd .. took us up one level of the directory tree. We should also note the existence of rmdir, the negative equivalent to mkdir, which removes *empty* directories. Now that we have learnt how to create files and directories, it's only natural that we learn how to move and copy them using one of the directory shortcuts we learnt above (..).

#### Code Example 10: mv and cp

```
user@system:~/nix$ cp testfile ..
user@system:~/nix$ rm ../testfile
user@system:~/nix$ mv testfile ..
```

An important thing to mention here is that we need not actually remove testfile after copying it and before moving it, as the mv command would simply overwrite it.

#### 3.1 Other Basic Utilities

Before we move on to more advanced commands, there are a few further commands to briefly discuss. We can clear the terminal screen and we can also display all environmental variables with env. find and locate search for files and directories (with the latter being faster - performing on a database of indexed filenames). date displays or sets (with the option -s) the system time, and cal displays the calender. history and specifically -history [NUM] reports the last [NUM] commands to be executed in the session. With some specific shells, the up and down arrow keys can be used to traverse the history manually. We can exit from the current terminal session, logout as a specific user, or shutdown the machine entirely.

# 4 Slightly More Advanced Concepts

#### 4.1 Editors at the Command Line

In the last section we created files (with touch), but they were blank. Now is an important time to introduce the use of text editors at the command line through the use of programs such as vi, pico, nano and emacs. These programs are intended as plain text editors (as per Notepad on Windows) as opposed to a word processing suites (such as Word). They do, however, have a lot more power than Notepad (or Textedit on Mac). To use vi on a file, type in vi testfile to create a file called testfile. If the named file exists, then the first page of the file will be displayed in the command line, and if it doesn't, an empty file will be created and appear on the screen for editing. It is important to note, that unlike many GUI based editors, the mouse does not move the cursor. Unlike PC editors, you cannot modify text by highlighting it with a mouse. Despite these restrictions and several others, there are an extremely wide range of keyboard shortcuts which can be used to enhance productivity. Lets enter vi to create a file which we will use for some examples later on: a

list of all the fruits we can think of (possibly with some help from Wikipedia). We'll save it in the nix directory and is call it allfruits.

# Code Example 11: vi allfruits

```
1 Apple
2 Apricot
3 Avocado
4 .
5 .
6 "allfruits" 90 lines, 846 characters
```

# 4.2 I/O Redirection

# 4.2.1 Standard Input and Output

Now that we have a firm handle on the basic commands at our disposal, it is necessary to discuss a couple of slightly more advanced concepts before proceeding further. One of the most important facilities is the ability to redirect the input and output ('I/O') of commands, and even link commands together. As discussed above, most command line programs output their results to the 'standard output' (STDOUT): which, by default, is the display. We can redirect standard output to specific files using the '>' character, and append to a file using '> >'.

## Code Example 12: >

```
user@system:~/nix$ ls > filelist.txt
```

While the above relates to 'standard output', commands can also accept input from 'standard input', (STDIN) which, by default, is the keyboard. To redirect from standard input from a file, we use the '<' character in conjunction with the sort command which we will come to shortly:

#### Code Example 13: <

```
user@system:~/nix$ sort < filelist.txt</pre>
```

There is also a third stream which will go unexamined ('standard error' or STDERR) which is used for error messages – and also defaults to the terminal.

#### 4.2.2 Filters

This brings us to a class of programs called *filters*, which are extremely useful for I/O Redirection. Rather than give examples of each of them individually, we provide a list of the most commonly used filters (some of which will feature later):

- sort: Sorts standard input then outputs the sorted result on standard output. By default it will sort alphabetically (on the first column), but there are an extremely large number of options available to modify the sorting mechanism.
- uniq: Given a sorted stream of data from standard input, it removes duplicate lines of data (i.e., it makes sure that every line is unique).
- grep: Examines each line of data it receives from standard input and outputs every line that contains a specified pattern of characters.
- fmt: Reads text from standard input, then outputs formatted text on standard output.
- head: Outputs the first few lines of its input. Useful for getting the header of a file. By default it will print the first 10 lines.
- tail: Outputs the last few lines of input. Useful for things like getting the most recent entries from a log file, and is the opposite of head also defaults to (the last) 10 lines.
- cat: Concatenates files and displays their contents, or just displays contents if given one input.
- cut: Divides a file into several columns.
- more: Displays output of a file one page at a time: scrolled using spacebar.
- less: Displays output of a file one page at a time: scrolled up or down using arrow keys.

Two other filters to mention briefly are n1, which prints the line number before data, and wc which prints the count of lines, words and characters.

#### **4.2.3** The Pipe

The final I/O redirection tool to discuss is the pipe (|), which allows you to connect multiple commands together. The standard output of one command is fed into the standard input of another, enabling you to chain together individual commands to create something really powerful. The example below *pipes* the output from 1s into the head command which takes the input 1 to show us the first 1 line of the 1s output: we've chained the two commands together. We could optionally then redirect the standard output away from the command line into a file which contained the same output with a command such as: 1s | head -1 > myfirstpipe.txt.

### Code Example 14: |

```
user@system:~/nix$ ls | head -1
allfruits
```

For even more power, it's possible to combine pipes and redirection.

# 4.3 Wildcards and Regular Expressions

Another slightly more advanced concept worth consideration is the 'wildcard'. Wildcards are a set of tools that allow you to create a pattern which defines a specific set of files or directories. There are two simple types of wildcards: 1.) \* represents zero or more characters, 2.) ? represents a single character. Lets create a set of files for this example called fileonea, fileoneb, filetwoa, filetwob.

# Code Example 15: ? and \*

```
user@system:~/nix$ touch fileonea fileoneb filetwoa filetwob
user@system:~/nix$ ls file*a
fileonea filetwoa
user@system:~/nix$ ls filetwo?
filetwoa filetwob
```

A regular expression (or 'regex') includes such functionality, but is a much more powerful pattern matcher providing the ability to restrict the specific type of characters which are searched for. This allows a more flexible specification overall. There are a number of books dedicated exclusively to regular expressions, and a detailed examination of the subject is far beyond the scope our endeavor. Below we can list a small number of regular expression operators and then provide two quick examples to combine this functionality with the powerful grep command which wasn't fully explored above.

- . : Match any character.
- ^: Match the empty string at the beginning of a line or string.
- \$: Match the empty string at the end of a line.
- A: Match an uppercase i.e. A.
- · a: Match a lowercase a.
- \d : Match any single digit.
- \D : Match any single non-digit character.
- \w: Match any single alphanumeric character.
- [A-E]: Match any of uppercase A, B, C, D, or E (most commonly seen as [A-Z]).
- [^A-E]: Match any character except uppercase A, B, C, D, or E.
- (abc|xyz): Match a sequence of at least one of abc or xyz.

Remember from above: the grep utility filters input, looking for matches. In the simplest use, grep prints those lines that contain text that matches a pattern. In our example, we pass it the -E option, which allows it to interpret the pattern as an extended regular expression.

#### Code Example 16: grep -E

```
user@system:~/Documents$ grep -E [z] allfruits
```

- 3 user@system:~/Documents\$ grep -E ^'(Red|Blue)' allfruits
- 4 Blueberry
- 5 Redcurrent

<sup>2</sup> Yuzu

The first example searches for any fruit which contains the lowercase character z ([z]) in the list of fruits, where the only fruit meeting this criteria is the yuzu - a citrus fruit which looks like a small grapefruit originating from China and Tibet. The second grep command looks for instances at the start of the line (^) which contain either 'Red' or 'Blue'. Further to these tools, awk is another powerful utility for text processing and pattern matching which can be used to perform batch processing functions (such as renaming multiple files). However, covering it fully is far beyond the scope of this chapter.

# 4.4 Users, Groups and Permissions

Permissions specify what a user can and cannot do - perhaps, for example, you want to lock your files in such a way that other people cannot change them, or perhaps you want to secure system files from damage. Permissions are split into three distinct categories which govern the ability to read (r), write (w) and execute (x - reserved for programs, scripts and directories). For every file, we need to define each of these permissions for three sets of potential users: the owner of the file, the group which owns the file, and everyone else. There are typically only two people who can manage the permissions of a given file or directory: the owner and the root user. Groups are used to simplify the management of system security, where users can be a member of one or more groups (and at least one by default). To view the permissions associated with an individual file, we can use the -1 option on the 1s command:

# Code Example 17: ls -l

```
user@system:~/nix$ ls -l allfruits
-rw-rw-r-- 1 user user 846 May 29 14:57 allfruits
```

How can we interpret this? The first character to the left determines whether it is a file (-) or a directory (d). Following this, we have information on the permissions for the owner, where r, w and x are described as above. If the permissions are missing, a - represents the omission of a permission. For example, with regards to allfruits as above, the owner of the file has read and write but not execute permissions. The same is true for the group of the owner, but everyone else only has read permission. To change the permissions associated with a file or a directory, we need to use the chmod command, where, again, the permission arguements are made up of three components: 1.) Who are we changing the permission for?

The user (u), others (o), the group (g), or all (a)? 2.) Are we giving (+) or taking permission away (-)?, 3.) Which of the three permissions are we setting - r, w or x? Lets look at an example whereby we remove the read permissions from the group of the owner and others:

#### Code Example 18: chmod

```
user@system:~/nix$ chmod og-r allfruits
user@system:~/nix$ ls -l allfruits
rw----- 1 user user 846 May 29 15:09 allfruits
```

There is also a 'shorthand' method which can achieve the same objectives (e.g. chmod 751 allfruits). There exists the same series of permissions for directories, albeit with slightly different meanings. In the directory context (d), r means you have the ability to read what's in the directory, w gives the permission to write new files and directories into it, and x gives you the permission to enter into it. Importantly, permissions are not inherited from the parent directory - and the recursive option (-R) for chmod can be especially useful. Other important commands include chown which changes the owner of a file or directory, and chgrp which changes the group associated with a file or directory. The groups command prints which groups a user is in.

# 4.5 Archiving

Archiving in our context is equivilent to the .zip extension in Windows. The standard way to do this is to use the *tape archive* – tar. It creates and manipulates archive files – extracting from *tar*, *pax*, *jar*, etc, with the -x option, and creating them with the -c option. One reason why you might want to use tape archives is if you want to transfer files, but if this is your motivation, you usually want to compress the files also. The typical \*nix ways to compress archives are to either use gzip (Lempel-Ziv coding) or bzip2 (Burrows-Wheeler block sorting) compression. Lets see some simple examples, first making a second file from our allfruits list to tar them together before zipping them.

#### Code Example 19: tar, gzip and bzip2

```
user@system:~/nix$ head -5 allfruits > top5fruits
user@system:~/nix$ ls -la
3 -rw----- 1 user user 846 May 30 13:32 allfruits
4 -rw-rw-r-- 1 user user
                           38 May 30 13:33 first5fruits
5 user@system:~/nix$ tar -cvf fruits.tar allfruits first5fruits
6 user@system:~/nix$ gzip fruits.tar
7 user@system:~/nix$ tar -cvf fruits.tar allfruits first5fruits
8 user@system:~/nix$ bzip2 fruits.tar
g user@system:~nix$ bzip2 ls -1
10 -rw----- 1 user user 10240 May 30 14:07 allfruits
11 -rw-rw-r-- 1 user user
                           38 May 30 13:33 first5fruits
12 -rw-rw-r-- 1 user user
                           214 May 30 14:13 fruitlists.tar.bz2
                           224 May 30 14:13 fruitlists.tar.gz
13 -rw-rw-r-- 1 user user
```

We more commonly create the archive and compression in one step with options like -cz, -xz (for gzip) and -cj, -xj for (bzip2).

# 5 Bash Scripting

Bash scripting represents a powerful tool which allows you to perform complex, and often repetitive tasks with minimal effort. We are essentially writing a series of commands (as above) which tell the Bash shell what to do next. A bash script is nothing more than a plain text file which contains commands which could also be taken out be ran on the command line. It is a convention to give bash scripts an extension of .sh. Lets create our first script (called myfirstscript.sh and describe how it works:

```
#!/bin/bash
2 echo Hello There! Welcome to Bash Scripting!
```

The first line is called the 'shebang'. When a script with a shebang is ran, the program loader parses the rest of the script's initial line (after #!) as an interpretter directive the path to the program that should be used to run the rest of the lines in the text file. The second line is the echo command which we saw above, and prints the message to the STDOUT. We can execute the script in two ways to obtain the same result:

# Code Example 20: ./ and bash

```
user@system:~/nix$ ./myfirstscript.sh
Hello There! Welcome to Bash Scripting!
user@system:~/nix$ bash myfirstscript.sh
Hello There! Welcome to Bash Scripting!
```

It is important to check the file permissions (especially: x!) before trying to execute a .sh script. Now that we've learned to execute the scripts, we can begin to learn about variables: temporary methods of storing information. There are two key actions: setting - which requires no \$, and reading, which does. Lets see an example of how we can pass a variable to the script (called variables.sh), by defining a simple echo script as above which accepts \$1 and \$2 as inputs:

```
#!/bin/bash
2 echo Hello $1! Message sent from $2!
```

We can then execute it as before:

#### Code Example 21: variables

```
user@system:~/nix$ ./variables.sh Reader Author
Hello Reader! Message sent from Author!
```

There are also a number of special variables which we need to mention:

- \$0: The filename of the current script.
- n: The arguments with which a script was invoked.
- \$#: The number of arguments supplied.
- \$? : The exit status of the last command.
- \$\$: Process number of the current shell.

We can also set our own variables such as ourvariable=value. Remembering this, and the fact that if we want variables with more complex values however we need quotations, lets try a second example (called settingvariables.sh):

```
1 #!/bin/bash
2 recipient=''Person One''
3 sender=''Person Two''
4 echo Hello $recipient. Sent from: $sender
```

Which is then executed to give:

#### Code Example 22: setting variables

```
user@system:~/nix$ ./settingvariables.sh
Hello Person One! Message sent from Person Two!
```

We can also substitute the output of a command or program and save it to the value of a variable and export it elsewhere. We can also design shell scripts (readvariable.sh)to ask the user for input using the read command. For example:

```
#!/bin/bash
2 echo Hey, Buddy! What is your favorite color?
3 read favoritecolor
4 echo Wow! $favoritecolor is my favorite color too!
```

And then execute it as:

#### Code Example 23: bash reading variables

```
user@system:~/nix$ ./readvariable.sh
Hey, Buddy! What is your favorite color?
Blue
Wow! Blue is my favorite color too!
```

We can also do simple arithmetic operations using the let and expr commands. We can utilize the standard opreators of +, -, \\*, / which represent add, subtract, multiple and divide, as well as var++ for 'increase by one' and var- for 'decrease by one'. Finally, we can calculate the modulus with %. In particular, let stores the variable, but expr prints it out. Lets try out each of these features, feeding variables into our script:

```
1 #!/bin/bash
2 let "a = $1 + $2"
3 echo adding $1 and $2 together gives $a
4 echo multiplying $1 and $2 together gives $(expr $1 \* $2)
```

And then execute it as:

# Code Example 24: bash arithmetic

```
user@system:~/nix$ ./arithmetic.sh 10 5
2 adding 10 and 5 together gives 15
3 multiplying 10 and 5 together gives 50
```

We can also embed if statements, just like in other languages:

```
if [ condition ]
then
< execute commands >
fi
```

with everything between if and fi beign executed. There is no need to indent, and we can also further augment the script with nested statements, else, elif; and so on. We can also utilize various types of loops; while, until and for which we can continue and break out of as expected. We can also define custom functions, although this is a more advanced topic to be left for another discussion.

# 6 Sociogenomics Specifics

Now that we have a fuller understanding of the command line, we move on to some specific examples related to genomics. In particular, we'll outline three practical applications: some basic commands used on genomics data, executing scripts in languages most useful for sociogenomics (R and Python), and then executing genomics related programs to undertake a common task (calculating polygenic scores).

# 6.1 Apples

We're first going to take genome and annotations files related to apple (apple.genome and apple.gene) in order to answer a list of practical questions. The genome file is a Multi-FASTA format, as shown below:

#### Code Example 25: more apple.genome

```
user@system:~/nix$ more apple.genome
```

9 . . .

Using the commands which we learnt above, we can calculate the number of chromosomes in the genome using the fact that each chromosome begins with a header line which includes the distinctive character '>'. Therefore, we can use a combination of grep, | and wc to calculate the number of chromosomes:

# Code Example 26: grep apple.genome

```
user@system:~/nix/$ grep '>' apple.genome | wc -1.
2 3
```

We can also answer a series of descriptive questions based on the apple.genes file.

<sup>2 &</sup>gt;chr1

<sup>3</sup> ATTTTTTTCCTTTTGTTTTTGCATACAAGTCAATTTCATTCGAA

<sup>4</sup> GTTTGGTTGGATCGAAAAGTCGGGGTTCAATTTTTTGATCCGAC

<sup>5</sup> CACAATCAAGCTTCAACCTATGAAGATTGTCAATTTGGCCAAGG

<sup>6</sup> AAAGAGTTAGGCGACCTGTTTTGACATTTGAAAAGGAAATATG

<sup>7</sup> GTTGGCCTGGAAGAAATATTCAAGTTTCTATGCCA AATATAAA

<sup>8</sup> AATTGATTCAGAAGGTGATTCGTCGAATAAATGCTTAGATGATGA

These include things like 'How many genes are there?', 'How many transcript variants are there?' and 'How many genes have a single splice variant?'. As always, we can first have a glance at the file to check that it is the format which we expect:

Code Example 27: head apple.genes

```
user@system:~/nix$ head -3 apple.genes
2 MDP0000303933
                   MDP0000303933
                                      chr1
                                                   4276
                                                            5447
3 MDP0000223353
                   MDP0000223353
                                                   77339
                                                             79628
                                     chr1
                                              +
4 MDP0000322928
                   MDP0000322928
                                      chr1
                                                   103533
                                                              103686
```

And then undertake the analysis:

Code Example 28: grep and cut apple.genes

```
user@system:~/nix$ cut -f1 apple.genes | sort -u | wc -l
5453
user@system:~/nix$ cut -f2 apple.genes | sort -u | wc -l
5456
user@system:~/nix$ cut -f1 apple.genes | uniq -c | grep ' 1 ' | wc -l
5450
```

# 6.2 Running Scripts from the Command Line

The majority of analytics undertaken with large datasets in social science now typically involve the use of HPCs which often run a \*nix and often do not have a GUI. This necessitates the submission of scripts to the HPCs through the command line. In our experience, the most prevalent languages for genetic data science and sociology are R, Python and Stata. Given the restrictive licensing associated with the third of these software platforms, we will outline how to execute scripts using the the first two. This is as simple as creating your .R and .py files (or using rsync or scp to transfer them from your local machine to the HPC where applicable) in your directory, and executing them. Lets create an example file called helloworld (remembering that we are in an extensionless environment), which simply has the single line: print(''Hello World! How are you?''). Assuming that you have Python 3 and not Python 2 installed, we can then execute the same files with both languages as follows to receive almost identical output:

#### **Code Example 29:** Python and R

```
user@system:~/nix$ python helloworld
Hello World! How are you?
user@system:~/nix$ Rscript helloworld
[1] ''Hello World! How are you?''
```

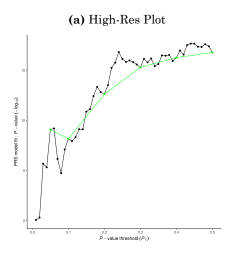
# 6.3 Genomic Software: Examples with Python and R

As part of our final example, we'll utilize two closely related pieces of genomics software directly from the command line. The first is PRSice (pronounced 'precise'): a software package for calculating, applying, evaluating and plotting the results of polygenic risk scores. It is written in R and includes wrappers for bash data management scripts and PLINK2. To further motivate our use of \*nix-like systems, PRSice is a perfect example of a software which is only compatible with Unix/Linux/Mac operating systems. Further information is available at http://PRSice.info. To begin, download and upzip the software to a local directory (i.e. nix). The '\' command shows how we can break across lines:

# Code Example 30: PRSice

```
user@system:~/nix$ R -q --file=./PRSice_v1.25.R --args base \
TOY_BASE_GWAS.assoc \
target TOY_TARGET_DATA \
slower 0 \
supper 0.5 \
sinc 0.01 \
covary F \
clump.snps F \
plink ./plink_1.9_linux_160914 \
figname EXAMPLE_1
```

Figure 3: PRSice Outputs with the Toy Dataset



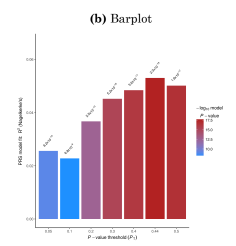


Figure 3 shows the two different outputs from this command. The second command line software which we will examine is LDpred: a Python based software package that adjusts GWAS summary statistics for the effects of linkage disequilibrium (LD). In order to calculate polygenic scores from this software package, we need to execute three commands, and one way to do this simultaneously is via a shell script:

#### Code Example 31: A Script for LDPred

```
echo ''Now Running: coord_genotypes''
python ldpred/coord_genotypes.py \
--gf=test_data/LDpred_data_p0.001_test_0 \
--ssf=test_data/LDpred_data_p0.001_ss_0.txt --N=100 --out=bogus.hdf5
ceho ''Now Running LDpred.py''
python ldpred/LDpred.py --coord=bogus.hdf5 --ld_radius=1 \
--PS=0.3 --N=100 --out=ldpredoutput
ceho ''Running the validate.py''
python ldpred/validate.py \
--vgf=test_data/LDpred_cc_data_p0.001_train_0 \
--rf=ldpredoutput --out=pgsoutputs
```

# 7 Conclusion

While we have introduced the command line, we have only begun to breach the surface of what it makes possible. There are a whole range of more advanced topics from both within a \*nix-like operating system perspective (such as networking commands or process control and scheduling) as well as those concerning genomics and bioinformatics (such as SAMTools which provides a range of tools for manipulating read alignments or PLINK, a free, open-source whole genome association analysis toolset). For this reason, we provide a reading list below to allow you to further explore the opportunities which the command line makes available.

# 8 Further Reading

Chadwick, R. (2017), 'Linux Tutorial', http://ryanstutorials.net/.

Florea, L. (2017), 'Command Line Tools for Genomic Data Science', Coursera Module: https://www.coursera.org/learn/genomic-tools/home/welcome.

Kerrisk, M. (2010), 'The Linux Programming Interface: A Linux and UNIX System Programming Handbook', No Starch Press, 1st Edition, ISBN: 1593272200.

Marsh, N. (2010), 'Introduction to the Command Line', Second Edition, Second Revision, ISBN: 1450588301.

Sobell, M. (1994), 'A Practical Guide to the UNIX System Paperback', Pearson, Third Edition, ISBN: 0805375651.