Practical Computing and Bioinformatics for Conservation and Evolutionary Genomics

To my son, without whom I should have finished this book two years earlier

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Preface

This is a collection of blurbs Eric started writing in the last year to help remind himself and others of some useful things to know for bioinformatics.

It was started during an extended government shutdown, when it seemed like writing a book might be in order. Fortunately, the shutdown ended eventually. It is not clear whether a book will ever come of it, but these notes shall be up on the web indefinitely, so feel free to use them!

0

Introduction

Nothing here yet.

Eric's Notes of what he might do

This is where I am going to just throw out ideas and start to organize them. My thought was that while I am actually doing bioinformatics, etc. in my normal day-to-day work I will analyze what I am doing and figure out all the different tools that I am using and organize that or pedagogy.

• Note: I am going to make a companion repository called mega-bioinf-pop-gen-examples that will house all of the data sets and things for exercises.

0.1 Table of topics

Man! There is going to be a lot to get through. My current idea is to meet three times a week. The basic gist of those three sessions will be like this:

- 1. Fundamental Tools / Environments: I am thinking 5 weeks on Unix, 1 Week on HPC, 6 Weeks on R/Rstudio, and 3 on Python from within Rstudio (so that students know enough to run python modules like moments.)
- 2. Theory and Background: Population-genetic and bioinformatic theory. Alignment and BW transforms, the coalescent, Fst, etc. Basically things that are needed to understand (to some degree) what various programs/analyses are doing under the hood.
- 3. Application and Practice: Getting the students to get their feet wet and their fingers dirty actually doing it. This time should be entirely practical, with students doing an exercise (in pairs or groups, possibly) with me (or someone else, maybe CH) overseeing.

Week	Fundamental	Theory and	Application and
	Tools	Background	Practice
1	Unix Intro: filesystem; absolute and relative paths, everything is a file; readable, writable, executable; PATH; .bashrc — hack everyone's to get the time and directory; TAB- completion; cd, ls (colored output), cat, head, less; stdout and stderr and file redirection of either with > and 2>; the; vs &. Using TextWrangler with edit and we need a PC	Data Formats: fasta, fastq, SAM, BAM, VCF, BCF	Command line drills

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Week	Fundamental Tools	Theory and Background	Application and Practice
2	Programs,	Fundamental ·	samtools, veftools,
	binaries,	programming	beftools. hands on,
	compiling, installing,	concepts; Scripts vs binaries (i.e. compiled	doing stuff with them, reading the man
	package	vs interpreted	~
	management;	languages);	pages, exercises.
	software	dependencies: headers	
	distribution;	and libraries;	
	GitHub and	Modularization;	
	sourceforge;	Essential algorithms;	
	admin	compression;	
	privileges and	compression,	
	sudo, and how		
	you probably		
	won't have that		
	on a cluster.		
3	Programming		
	on the shell:		
	variables and		
	variable		
	substitution;		
	Globbing and		
	path expansion;		
	variable		
	modifications;		
	loops;		
	conditionals;		
	sed, awk, and		
	regular		
_	expressions		
5	HPC: clusters;		
	nodes; cores;		
	threads. SGE		
	and/or SLURM;		
	qsub; qdel;		
	<pre>qacct; myjobs;</pre>		
	job arrays.		

Part I

Part I: Essential Computing Skills

1

Overview of Essential Computing Skills

What up with this? It appears that bookdown does not let you write things at the beginning of a Part without putting it under a chapter heading. Oh well.

Essential Unix/Linux Terminal Knowledge

Unix was developed at AT&T Bell Labs in the 1960s. Formally "UNIX" is a trademarked operating system, but when most people talk about "Unix" they are talking about the *shell*, which is the text-command-driven interface by which Unix users interact with the computer.

The Unix shell has been around, largely unchanged, for many decades because it is *awesome*. When you learn it, you aren't learning a fad, but, rather, a mode of interacting with your computer that has been time tested and will likely continue to be the lingua franca of large computer systems for many decades to come.

For bioinformatics, Unix is the tool of choice for a number of reasons: 1) complex analyses of data can be undertaken with a minimum of words; 2) Unix allows automation of tasks, especially ones that are repeated many times; 3) the standard set of Unix commands includes a number of tools for managing large files and for inspecting and manipulating text files; 4) multiple, successive analyses upon a single stream of data can be expressed and executed efficiently, typically without the need to write intermediate results to the disk; 5) Unix was developed when computers were extremely limited in terms of memory and speed. Accordingly, many Unix tools have been well optimized and are appropriate to the massive genomic data sets that can be taxing even for today's large, high performance computing systems; 6) virtually all state-of-the-art bioinformatic tools are tailored to run in a Unix environment; and finally, 7) essentially every high-performance computer cluster runs some variant of Unix, so if you are going to be using a cluster for your analyses (which is highly likely), then you have gotta know Unix!

2.1 Getting a bash shell on your system

A special part of the Unix operating system is the "shell." This is the system that interprets commands from the user. At times it behaves like an interpreted programming language, and it also has a number of features that help to minimize the amount of typing the user must do to complete any particular

tasks. There are a number of different "shells" that people use. We will focus on one called "bash," which stands for the "Bourne again shell." Many of the shells share a number of features.

Many common operating systems are built upon a Unix or upon Linux—an open-source flavor of Unix that is, in many scenarios, indistinguishable. Hereafter we will refer to both Unix and Linux as "Unix" systems). For example all Apple Macintosh computers are built on top of the Berkeley Standard Distribution of Unix and bash is the default shell. Many people these days use laptops that run a flavor of Linux like Ubuntu, Debian, or RedHat. Linux users should ensure that they are running the bash shell. This can be done by typing "bash" at the command line, or inserting that into their profile. To know what shell is currently running you can type:

echo \$0

at the Unix command line. If you are running bash the result should be

-bash

PCs running Microsoft Windows are something of the exception in the computer world, in that they are not running an operating system built on Unix. However, Windows 10 now allows for a Linux Subsystem to be run. For Windows, it is also possible to install a lightweight implementation of bash (like Git Bash). This is helpful for learning how to use Unix, but it should be noted that most bioinformatic tools are still difficult to install on Windows.

2.2 Navigating the Unix filesystem

Most computer users will be familiar with the idea of saving documents into "folders." These folders are typically navigated using a "point-and-click" interface like that of the Finder in Mac OS X or the File Explorer in a Windows system. When working in a Unix shell, such a point-and-click interface is typically not available, and the first hurdle that new Unix users must surmount is learning to quickly navigate in the Unix filesystem from a terminal prompt. So, we begin our foray into Unix and its command prompt with this essential skill.

When you start a Unix shell in a terminal window you get a *command prompt* that might look something like this:

my-laptop:~ me\$

or, perhaps something as simple as:

\$

or maybe something like:

/~/--%

We will adopt the convention in this book that, unless we are intentionally doing something fancier, the Unix command prompt is given by a percent sign, and this will be used when displaying text typed at a command prompt, followed by output from the command. For example

% pwd /Users/eriq

shows that I issued the Unix command pwd, which instructs the computer to print working directory, and the computer responded by printing /Users/eriq, which, on my Mac OS X system is my home directory. In Unix parlance, rather than speaking of "folders," we call them "directories;" however, the two are essentially the same thing. Every user on a Unix system has a home directory. It is the domain on a shared computer in which the user has privileges to create and delete files and do work. It is where most of your work will happen. When you are working in the Unix shell there is a notion of a current working directory—that is to say, a place within the hierarchy of directories where you are "currently working." This will become more concrete after we have encountered a few more concepts.

The specification /Users/eriq is what is known as an absolute path, as it provides the "address" of my home directory, eriq, on my laptop, starting from the root of the filesystem. Every Unix computer system has a root directory (you can think of it as the "top-most" directory in a hierarchy), and on every Unix system this root directory always has the special name, /. The address of a directory relative to the root is specified by starting with the root (/) and then naming each subsequent directory that you must go inside of in order to get to the destination, each separated by a /. For example, /Users/eriq tells us that we start at the root (/) and then we go into the Users directory (Users) and then, from there, into the eriq directory. Note that / is used to mean the root directory when at the beginning of an absolute path, but in the remainder of the path its meaning is different: it is used merely as a separator

between directories nested within one another. Figure 2.1 shows an example hierarchy of some of the directories that are found on the author's laptop.

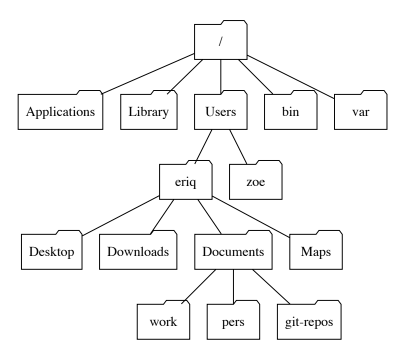


FIGURE 2.1: A partial view of the directories on the author's laptop.

From this perspective it is obvious that the directory eriq lives inside Users, and also that, for example, the absolute path of the directory git-repos would be /Users/eriq/Documents/git-repos.

Absolute paths give the precise location of a directory relative to the root of the filesystem, but it is not always convenient, nor appropriate, to work entirely with absolute paths. For one thing, directories that are deeply nested within many others can have long and unwieldy absolute path names that are hard to type and can be difficult to remember. Furthermore, as we will see later in this book, absolute paths are typically not reproducible from one computer's filesystem to another. Accordingly, it is more common to give the address of directories using relative paths. Relative paths work much like absolute paths; however, they do not start with a leading /, and hence they do not take as their starting point the root directory. Rather, their starting point is implicitly taken to be the current working directory. Thus, if the current

working directory is /Users/eriq, then the path Documents/pers is a relative path to the pers directory, as can again be seen in Figure 2.1.

The special relative path symbol .. means "the directory that is one level higher up in the hierarchy." So, if the current working directory were /Users/eriq/Documents/git-repos, then the path .. would mean /Users/eriq/Documents, the path ../work gives the directory /Users/eriq/Documents/work, and, by using two or more .. symbols separated by forward slashes, we can even go up multiple levels in the hierarchy: ../../zoe is a relative path for /Users/zoe, when the current working directory is /Users/eriq/Documents/git-repos.

When naming paths, another useful Unix shorthand is ~ (a tilde) which denotes the user's home directory. This is particularly useful since most of your time in a Unix filesystem will be spent in a directory within your home directory. Accordingly, ~/Documents/work is a quick shorthand for /Users/eriq/Documents/work. This is essential practice if you are working on a large shared computing resource in which the absolute path to your home directory might be changed by the system administrator when restructuring the filesystem.



A useful piece of terminology: in any path, the "final" directory name is called the *basename* of the path. Hence the basename of /Users/eriq/Documents/git-repos is git-repos. And the basename of ../../Users is Users.

2.2.1 Changing the working directory with cd

When you begin a Unix terminal session, the current working directory is set, by default, to your home directory. However, when you are doing bioinformatics or otherwise hacking on the command line, you will typically want to be "in another directory" (meaning you will want the current working directory set to some other directory). For this, Unix provides the cd command, which stands for change directory. The syntax is super simple:

cd path

where *path* is an absolute or a relative path. For example, to get to the git-repos directory from my home directory would require a simple command cd Documents/git-repos. Once there, I could change to my Desktop directory with cd ../../Desktop. Witness:

% pwd
/Users/eriq

```
% cd Documents/git-repos/
% pwd
/Users/eriq/Documents/git-repos
% cd ../../Desktop
% pwd
/Users/eriq/Desktop
```

Once you have used cd, the working directory of your current shell will remain the same no matter how many other commands you issue, until you invoke the cd command another time and change to a different directory.

If you give the cd command with no path specified, your working directory will be set to your home directory. This is super-handy if you have been exploring the levels of a Unix filesystem above your home directory and cannot remember how to get back to your home directory. Just remember that

% cd

will get you back home.

Another useful shortcut is to supply - (a hyphen) as the path to cd. This will change the working directory back to where you were before your last invocation of cd, and it will tell you which directory you have returned to. For example, if you start in /Users/eriq/Documents/git-repos and then cd to /bin, you can get back to git-repos with cd - like so:

```
% pwd
/Users/eriq/Documents/git-repos
% cd /bin/
% pwd
/bin
% cd -
/Users/eriq/Documents/git-repos
% pwd
/Users/eriq/Documents/git-repos
```

Note the output of cd - is the newly-returned-to current working directory.

2.2.2 Updating your command prompt

When you are buzzing around in your filesystem, it is often difficult to remember which directory you are in. You can always type pwd to figure that out,

but the bash shell also provides a way to print the current working directory within your command prompt.

For example, the command:

```
PS1='[\W]--% '
```

redefines the command prompt to be the basename of the current directory surrounded by brackets and followed by --%:

```
% pwd
/Users/eriq/Documents/git-repos
% PS1='[\W]--% '
[git-repos]--% cd ../
[Documents]--% cd ../
["]--% cd ../
[Users]--%
```

This can make it considerably easier to keep track of where you are in your file system.

We will discuss later how to invoke this change automatically in every terminal session when we talk about customizing environments in Section 2.5.

2.2.3 TAB-completion for paths

Let's be frank...typing path names in order to change from one directory to another can feel awfully tedious, especially when your every neuron is screaming, "Why can't I just have a friggin' Finder window to navigate in!" Do not despair. This is a normal reaction when you first start using Unix. Fortunately, Unix file-system navigation can be made much less painful (or even enjoyable) for you by becoming a master of *TAB-completion*. Imagine the Unix shell is watching your every keystroke and trying to guess what you are about to type. If you type the first part of a directory name after a command like cd and then hit the TAB key, the shell will respond with its best guess of how you want to complete what you are typing.

Take the file hierarchy of Figure 2.1, and imagine that we are in the root directory. At that point, if we type cd A, the shell will think "Ooh! I'll bet they want to change into the directory Applications because that is the only directory that starts with A. Sure enough, if you hit TAB, the shell adds to the command line so that cd A becomes cd Applications/ and the cursor is still waiting for further input at the end of the command. Boom! That was way easier (and more accurate) than typing all those letters after A.

Developing a lightning-fast TAB-completion trigger finger is, quite seriously, essential to surviving and thriving in Unix. Use your left pinky to hit TAB. Hone your skills. Make sure you can hit TAB with your eyes closed. TAB early and TAB often!

Once you can hit TAB instantly from within the middle of any phrase, you will also want to understand a few simple rules of TAB completion:

- 1. If you try TAB-completing a word on the command line that is not at the beginning of the command line (i.e., you are typing a word after a command like cd), then the shell tries to complete the word with a directory name or a file name.
- 2. The shell will only complete an *entire* directory or file name if the name *uniquely* matches the first part of the path that has been entered. In our example, there were no other directories than Applications in / that start with A, so the shell was certain that we must have been going for Applications.
- 3. If there is more than one directory or file name that matches what you have already typed, then, the first time you hit TAB, nothing happens, but the *second* time you hit TAB, the shell will print a list of names that match what you have written so far. For example, in our Figure 2.1 example, hitting TAB after typing cd ~/D does nothing. But the second time we hit TAB we get a list of matching names:

```
% cd ~/D
Desktop/ Documents/ Downloads/
```

So, if we are heading to Documents we can see that adding oc to our command line, to create cd Doc would be sufficient to allow the shell to uniquely and correctly guess where we are heading. cd Doc will TAB-complete into cd Documents/

4. If there are multiple directory or file names that match the current command line, and they share more letters than those currently on the command line, TAB-completion will complete the name to the end of the shared portion of the name. An example helps: let's say I have the following two directories with hideously long names in my Downloads folder:

```
WIFL.rep_indiv_est.mixture_collection.count.gr8-results
WIFL.rep_indiv_est.mixture_collection.count-results
```

Then, TAB completing on ~/Downloads/WIFL.rep will partially complete so that the prompt and command look like:

```
% cd ~/Downloads/WIFL.rep_indiv_est.mixture_collection.count
```

and hitting TAB twice gives:

```
% cd ~/Downloads/WIFL.rep_indiv_est.mixture_collection.count
WIFL.rep_indiv_est.mixture_collection.count-results
WIFL.rep_indiv_est.mixture_collection.count.gr8-results
```

At this point, adding - and TAB completing will give the first of those directories.

The last example shows just how much typing TAB completion can save you. So, don't be shy about hitting that TAB key. When navigating your filesystem (or writing longer command lines that require paths of files) you should consider hitting TAB after every 1 or 2 letters. In routine work on the command line, probably somewhere around 25% or more of my keystrokes are TABs. Furthermore, a TAB is never going to execute a command, and it typically won't complete to a path that you don't want (unless you got the first part of its name wrong), so there isn't any risk to hitting TAB all the time.

2.2.4 Listing the contents of a directory with 1s

So far we have been focusing mostly on directories. However, directories themselves are not particularly interesting—they are merely containers. It is the *files* inside of directories that we typically work on. The command 1s lists the contents—typically files or other directories—within a directory.

Invoking the ls command without any other arguments (without anything after it) returns the contents of the current working directory. In our example, if we are in /Users then we get:

```
% ls eriq zoe
```

By default, 1s gives output in several columns of text, with the directory contents sorted lexicographically. For example, the following is output from the 1s command in a directory on a remote Unix machine:

% ls map-sliced-fastqs-etc.sh

bam-slices play

bwa-run-list.txt REDOS-map-sliced-fastqs-etc.sh

fastq-file-prefixes.txt sliced

fqslice-22.error slice-fastqs.sh
fqslice-22.log slicer-lines.txt

map-etc.sh Slicer-Logs-summary.txt

The first line shows the command prompt and the command: % ls, and the remainder is the output of the command.

Invoked without any further arguments, the ls command simply lists the contents of the current working directory. However, you can also direct ls to list the contents of another directory by simply adding the path (absolute or relative) of that directory on the command line. For example, continuing with the example in Figure 2.1, when we are in the home directory (eriq) we can see the directories/files contained within Documents like so:

```
[~]--% ls Documents
git-repos/ pers/ work/
```

If you give paths to more than one directory as arguments to ls, then the contents of each directory are listed after a heading line that gives the directory's path (as given as an argument to ls), followed by a colon. For example:

You might also note in the above example, that some of the paths listed within each of the two directories are followed by a slash, /. This ls customization denotes that they are directories themselves. Much like your command prompt, ls can be customized in ways that make its output more informative. We will return to that in Section 2.5.

If you pass the path of a file to ls, and that file exists in your filesystem, then ls will respond by printing the file's path:

```
% ls Documents/git-repos/lowergranite_0.0.1.tar.gz
Documents/git-repos/lowergranite_0.0.1.tar.gz
```

If the file does not exist you get an error message to that effect:

```
% ls Documents/try-this-name
ls: Documents/try-this-name: No such file or directory
```

The multi-column, default output of 1s is useful when you want to scan the contents of a directory, and quickly see as many files as possible in the fewest lines of output. However, this output format is not well structured. For example, you don't know how many columns are going to be used in the default output of 1s (that depends on the length of the filenames and the width of your terminal), and it offers little information beyond the names of the files.

You can tell the ls command to provide more information, by using it with the -l option. Appropriately, with the -l option, the ls command will return output in *long* format:

```
2019-02-08 21:09 /osu-chinook/--% ls -l
total 108
drwxr-xr-x 2 eriq kruegg 4096 Feb 7 08:26 bam
drwxr-xr-x 14 eriq kruegg 4096 Feb 8 15:56 bam-slices
-rw-r--r- 1 eriq kruegg 17114 Feb 7 20:16 bwa-run-list.txt
-rw-r--r-- 1 eriq kruegg
                           824 Feb 6 14:14 fastq-file-prefixes.txt
                                   7 20:14 fqslice-22.error
-rw-r--r-- 1 eriq kruegg
                             0 Feb
                             0 Feb 7 20:14 fqslice-22.log
-rw-r--r-- 1 eriq kruegg
-rwxr--r-- 1 eriq kruegg 1012 Feb
                                   7 07:59 map-etc.sh
-rwxr--r-- 1 eriq kruegg 1138 Feb
                                   7 20:56 map-sliced-fastqs-etc.sh
drwxr-xr-x 3 eriq kruegg 4096 Feb
                                   7 13:01 play
                                   8 15:08 REDOS-map-sliced-fastqs-etc.sh
-rwxr--r-- 1 eriq kruegg 1157 Feb
drwxr-xr-x 14 eriq kruegg 4096 Feb
                                   8 15:49 sliced
-rwxr--r-- 1 eriq kruegg
                           826 Feb
                                   7 20:09 slice-fastqs.sh
-rw-r--r-- 1 eriq kruegg
                         1729 Feb
                                   7 16:11 slicer-lines.txt
```

Each row contains information about only a single file. The first column indicates what kind of file each entry is, and also tells us which users have permission to do certain things with the file (more on this in a few sections). The third and fourth columns show that the owner of each file is eriq, who is a user in the group called kruegg. After that is the size of the file (in bytes) and the date and time it was last modified.

There are a few options to 1s that are particularly useful. One is -a, which causes 1s to include in its listing all files, even *hidden* ones. In a Unix file

system, any file whose name starts with a . is considered a hidden file. Commonly, such files are configuration files or other files used by programs that you typically don't interact with directly. (We will see an example of this when we start working with git for version control, Section 8.2.) The -d option for 1s is also' quite handy. Recall that when you provide the name of a directory as an argument to 1s, the default behavior is to list the contents of the directory. This can be troublesome when you are listing the contents of a subdirectory: 1s ~/Documents/git-repos/* lists the contents (which can be substantial) of each of the directories in my directory, but I might only want to know the name of each of those directories, rather than their full contents. 1s -d ~/Documents/git-repos will do that for you. Finally, the -R option to 1s will cause the operating system to drill down, recursively into all the subdirectories of the one you supplied to the command, and list their contents, as well.

2.2.5 Globbing

If you have ever had to move a large number of files of a certain type from one folder to another in a Finder window, you know that individually clicking and selecting each one and then dragging them can be a tedious task (not to mention the disaster that ensues if you slip on your mouse and end up dropping all the files some place you did not intend). Unix provides a wonderful system called *filename expansion* or "globbing" for quickly providing the names of a large number of files and paths which let's you operate on multiple files quickly and efficiently. In short, globbing allows for wildcard matching in path names. This means that you can specify multiple files that have names that share a common part, but differ in other parts.

The most widely used (and the most permissive) wildcard is the asterisk, *. It matches anything in a file name. So, for example:

- *.vcf will expand to any files in the current directory with the suffix .vcf.
- D*s will expand to any files that start with an uppercase D and end with an $\ensuremath{\mathtt{g}}$
- *output-*.txt will expand to any files that include the phrase output-somewhere in their name and also end with .txt.
- * will expand to all files in the current working directory.
- /usr/local/*/*.sh will expand to any files ending in .sh that reside within any directory that is within the /usr/local directory.



Actually, there is some arcana here: Names of files or directories that start with a dot (a period) will not expand unless the dot is included explicitly. Files with names starting with a dot are "hidden" files in Unix. You

also will not see them in the results of ls, unless you use the -a option: ls -a.

After the asterisk, the next most commonly-used wildcard is the question mark,? The question mark denotes any single character in a file name. For example. If you had a series of files that looked like AA-file.txt, AB-file.txt, ..., AZ-file.txt. You could get get all those by using A?-file.txt. This would not expand to, for example, AAZ-file.txt, if that were in the directory.

You can be more specific in globbing by putting things within [and]. For example: A[A-D]* would pick out any files starting with, AA, AB, AC, or AD. Or you could have said A[a-d]* which would get any files starting with Aa, Ab, Ac, or Ad. And you can also do it with numbers: [0-9]. You can also negate the contents of the [], with ^. Thus, 100_[^ABC]* picks out all files that start with 100 followed by anything that is *not* and A, B, or a C.

Finally, you can be really specific about replacements in file names by iterating over different possibilities with a comma-separated list within curly braces. For example, img.{png,jpg,svg} will iterate over the values in curly braces and expand to img.png img.jpg img.svg. Interestingly, with curly braces, this forms all those file names whether they exist or not. So, unlike * it isn't really matching available file names.

The last thing to note about all of these globbing constructs is that they are not intimately associated with the ls command. Rather, they simply provide expansions on the command line, and the the ls command is listing all those files. For example, try echo *.txt.

2.2.6 What makes a good file-name?

If the foregoing discussion suggests to you that it might not be good to use an actual *, ?, [, or { in names that you give to files and directories on your Unix system, then congratulations on your intuition! Although you can use such characters in your filenames, they have to be preceded by a backslash, and it gets to be a huge hassle. So don't use them in your file names. Additionally, characters such as #, |, and : do not play well for file names. Don't use them!

Another pet peeve of mine (and anyone who uses Unix) are file names that have spaces in them. In Windows and on a Mac it is easy to create file names that have spaces in them. In fact, the standard Windows system comes with such space-containing directory names as My Documents or My Pictures. Yikes! Please don't ever do that in your Unix life! One can deal with spaces in file names, but there is really no reason to include spaces in your file names, and having spaces in file names will typically break a good many scripts. Rather

than a space, use an underscore, _, or a dash, -. You've gotta admit that, not only does My-Documents work better, but it actually looks better too!

However, should you have to deal with files having spaces in their name, you can address them by either backslash escaping the spaces, or putting the whole file name in quotation marks (single or double quotation marks will work). If you have a file called dumb file name.jpg, you can address it on the command line as either of the following three:

```
dumb\ file\ name.jpg
"dumb file name.jpg"
'dumb file name.jpg'
```

To make your life easier, however, the bottom line is that you should name your files on a Unix system using only upper- and lowercase letters (Unix file systems are case-sensitive), numerals, and the following three punctuation characters: ., -, and _. Though you can use other punctuation characters, they often require special treatment, and it is better to avoid them altogether.

2.3 The anatomy of a Unix command

Nearly every Unix command that you might invoke follows a certain pattern. First comes the command itself. This is the word that tells the system the name of the command that you are actually trying to do. After that, often, you will provide a series of *options* that will modify the behavior of the command (for example, as we have seen, -1 is an option to the 1s command). Finally, you might then provide some *arguments* to the functions. These are typically paths to files or directories that you would like the command to operate on. So, in short, a typical Unix command invocation will look like this:

command options arguments

Of course, there are exceptions. For example, when invoking Java-based programs from your shell, arguments might be supplied in ways that make them look like options, etc. But, for the most part, the above is a useful way of thinking about Unix commands.

Sometimes, especially when using samtools or bcftools, the command part of the command line might including a command and a subcommand, like samtools view or bcftools query. This means that the operating system is calling the program samtools (for example), and then samtools interprets the next token (view) to know that it needs to run the view routine, and interpret all following options in that context.

We will now break down each element in "command options arguments".

2.3.1 The command

When you type a command at the Unix prompt, whether it is a command like ls or one like samtools (Section 14.2.1), the Unix system has to search around the filesystem for a file that matches the command name and which provides the actual instructions (the computer code, if you will) for what the command will actually do. It cannot be stressed enough how important it is to understand where and how the bash shell searches for these command files. Understanding this well, and knowing how to add directories that the shell searches for executable commands will alleviate a lot of frustration that often arises with Unix.

In brief, all Unix shells (and the bash shell specifically) maintain what is called an *environment variable* called PATH that is a colon-separated list of pathnames where the shell searches for commands. You can print the PATH variable using the **echo** command:

echo \$PATH

On a freshly installed system without many customizations the PATH might look like:

/usr/bin:/bin:/usr/sbin:/sbin

which is telling us that, when bash is searching for a command, it searches for a file of the same name as the command first in the directory /usr/bin. If it finds it there, then it uses the contents of that file to invoke the command. If it doesn't find it there, then it next searches for the file in directory /bin. If it's not there, it searches in /usr/sbin, and finally in /sbin. If it does not find the command in any of those directories then it returns the error command not found.

When you install programs on your own computer system, quite often the installer will modify a system file that specifies the PATH variable upon startup. Thus after installing some programs that use the command line on a Mac system, the "default" PATH might look like:

/usr/local/bin:/usr/bin:/usr/sbin:/sbin:/opt/X11/bin:/Library/TeX/texbin

2.3.2 The options

Sometimes these are called flags, and they provide a convenient way of telling a Unix command how to operate. We have already seen a few of them, like the -a, -1 or -d options to 1s.

Most, but not all, Unix tools follow the convention that options specified by a single letter follow a single dash, while those specified by multiple letters follow two dashes. Thus, the tar command takes the single character options -x, -v, and -f, but also takes an option named like --check-links. Some utilities also have two different names—a single-letter name and a long name—for many options. For example, the bcftools view program uses either -a or --trim-alt-alleles to invoke the option that trims alternate alleles not seen in a given subset of individuals. Other tools, like BEAGLE, are perfectly have options that are named with multiple letters following just a single dash.

Sometimes options take parameter values, like bcftools view -g het. In that case, het is a parameter value. Sometimes the parameter values are added to the option with an equals-sign.

With some unix utilities' single-letter options can be bunged together following a single dash, like, tar -xvf being synonymous with tar -x -v -f. This is not universal, and it is not recommended to expect it.

Holy Cow! This is not terribly standardized, and probably won't make sense till you really get in there and starting playing around in Unix...

2.3.3 Arguments

These are often file names, or other things that are not preceded by an option flag. For example, in the ls command:

ls -lrt dir3

-lrt is giving ls the options -l, -r, and -t and dir3 is the *argument*—the name of the directory whose contents you should list.

2.3.4 Getting information about Unix commands

Zheesh! The above looks like a horrible mish-mash. How do we find out how to use/invoke different commands and programs in Unix? Well, most programs are documented, and you have to learn how to read the documentation.

If a utility is properly installed, you should be able to find its manual page with the man command. For example, man ls or man tar. These "man-pages",

as the results are called, have a fairly uniform format. They start with a summary of what the utility does, then then show how it is invoked and what the possible options are by showing a skeleton in the form:

"command options arguments"

and usually square brackets are put around things that are not required. This format can get quite ugly and hard to parse for an old human brain, like mine, but stick with it.

If you don't have a man-page for a program, you might try invoking the program with the --help option, or maybe with no option at all.

2.4 Handling, Manipulating, and Viewing files and streams

In Unix, there are two main types of files: regular files which are things like text files, figures, etc.—Anything that holds data of some sort. And then there are "special" files, which include directories which you've already seen, and symbolic links which we will talk about later.

2.4.1 Creating new directories

You can make a new directory with:

```
mkdir path
```

where path is a path specification (either absolute or relative). Note that if you want to make a directory within a subdirectory that does currently not exist, for example:

```
mkdir new-dir/under-new-dir
```

when new-dir does not already exist, then you have to either create new-dir first, like:

```
mkdir new-dir
mdkir new-dir/under-new-dir
```

or you have to use the -p option of mkdir, which creates all necessary parent directories as well, like:

mkdir -p new-dir/under-new-dir

If there is already a file (regular or directory) with the same path specifiation as a directory you are trying to create, you will get an error from mkdir.

2.4.2 Fundamental file-handling commands

For the day-to-day business of moving, copying, or removing files in the file system, the three main Unix commands are:

- mv for moving files and directories
- cp for copying files and directories
- rm for removing files and directories

These obviously do different things, but their syntax is somewhat similar.

2.4.2.1 mv

mv can be invoked with just two arguments like:

mv this there

which moves the file (or directory) from the path this to the path there.

- If this is a regular file (i.e. not a directory), and:
 - there is a directory, this gets moved inside of there.
 - there is a regular file that exists, then there will get overwritten, becoming a regular file that holds the contents of this.
 - there does not exist, it will be created as regular file whose contents are identical to those of this.
- If this is a directory and:
 - there does not exist in the filesystem, the directory there will be made and its contents will be the (former) contents of this
 - if there already exists, and is a directory, then the directory this will be moved inside of the directory there (i.e. it will become there/this).
 - if there already exists, but is not a directory, then nothing will change
 in the filesystem, but an an error will be reported. In all cases, whatever
 used to exist at path this will no longer be found there.

And mv can be invoked with multiple arguments, in which case the last one must be a directory that already exists that receives all the earlier arguments inside it. So, if you already have a directory named dest_dir then you can move a lot of things into it like:

```
mv file1 file2 dir1 dir2 dest dir
```

You can also write that as as

```
mv file1 file2 dir1 dir2 dest_dir/
```

which makes its meaning a little more clear, but there is no requirement that the final argument have a trailing /.

Note, if any files in dest_dir have the same name as the files you are moving into dest_dir they will get overwritten.

So, you have gotta be careful not to overwrite stuff that you don't want to overwrite.

2.4.2.2 cp

This works much the same way as mv with two different flavors:

```
cp this there
```

and

```
cp file1 file2 dest_dir
# or
cp file1 file2 dest_dir/
```

The result is very much like that of mv, but instead of moving the file from one place to another (an operation that can actually be done without moving the data within the file to a different place on the hard drive), the cp command actually makes a full copy of files. Note that, if the files are large, this can take a long time.

2.4.2.3 rm

Finally we get to the very spooky rm command, which is short for "remove." If you say "rm myfile.txt" the OS will remove that file from your hard drive's directory. The data that were in the file might live on for some time on your hard drive—in other words, by default, rm does not wipe the file off your hard drive, but simply "forgets" where to look for that file. And the space that file took up on your hard drive is no longer reserved, and could easily be overwritten the next time you write something to disk. (Nonetheless, if you do rm a file, you should never expect to be able to get it back). So, be very careful

about using rm. It takes an -r option for recursively removing directories and all of their contents.

When used in conjunction with globbing, rm can be very useful. For example, if you wanted to remove all the files in a directory with a .jpg extension, you would do rm *.jpg from within that directory. However, it's a disaster to accidentally remove a number of files you might not have wanted to. So, especially as you are getting familiar with Unix, it is worth it to experiment with your globbing using ls first, to see what the results are, and only when you are convinced that you won't remove any files you really want should you end up using rm to remove those files.

2.4.3 "Viewing" Files

In a typical GUI-based environment, when you interact with files on your computer, you typically open the files with some application. For example, you open Word files with Microsoft Word. When working on the Unix shell, that same paradigm does not really exist. Rather, (apart from a few cases like the text editors, nano, vim and emacs, instead of opening a file and letting the user interact with it the shell is much happier just streaming the contents of the file to the terminal.

The most basic of such commands is the cat command, which catenates the contents of a file into a very special data stream called stdout, which is short for "standard output." If you don't provide any other instruction, data that gets streamed to stdout just shoots by on your terminal screen. If the file is very large, it might do this for a long time. If the file is a text file then the data in it can be written out in letters that are recognizable. If it is a binary file then there is no good way to represent the contents as text letters, and your screen will be filled with all sorts of crazy looking characters.

It is generally best not to cat very large files, especially binary ones. If you do and you need to stop the command from continuing to spew stuff across your screen, you can type cntrl-c which is the universal Unix command for "kill the current process happening on the shell." Usually that will stop it.



A note regarding terminals: On a Mac, the Terminal app is quite fast at spewing text across the screen. Megabytes of text or binary gibberish can flash by in seconds flat. This is not the case with the terminal window within RStudio, which can by abysmally slow, and usually doesn't store many lines of output.

Sometimes you want to just look at the top of a file. The head command shows you the first 10 lines of a file. That is valuable. The less command

shows a file one screenful at a time. You can hit the space bar to see the next screenful, and you can hit q to quit viewing the file.

Try navigating to a file and using cat, head, and less on it.

One particularly cool thing about cat is that if you say

```
cat file1 file2
```

it will catenate the contents of both files, in order, to stdout.

Now, one Big Important Unix fact is that many programs written to run in the Unix shell behave in the same way regarding their output: they write their output to *stdout*. We have already seen this with 1s: its output just gets written to the screen, which is where *stdout* goes by default.

2.4.4 Redirecting standard output: > and >>

Unix starts to get really fun when you realize that you can "redirect" the contents of stdout from any command (or group of commands...see the next chapter!) to a file. To do that, you merely follow the command (and all its options and arguments) with > path where path is the path specifying the file into which you wish to redirect stdout.

Witness, try this:

```
# echo three lines of text to a file in the /tmp directory
echo "bing
bong
bong" > /tmp/file1

# echo three more lines of text to another file
echo "foo
bar
baz" > /tmp/file2

# now view the contents of the first file
cat /tmp/file1

# and the second file:
cat /tmp/file2
```

It is important to realize that when you redirect output into a file with >, any contents that previously existed in that file will be deleted (wiped out!). So be

careful about redirecting. Don't accidentally redirect output into a file that has valuable data in it.

The >> redirection operator does not delete the destination file before it redirects output into it. Rather, >> file means "append *stdout* to the contents that already exist in file." This can be very useful sometimes.

2.4.5 stdin, < and |

Not only do most Unix-based programs deliver output to standard output, but most utilities can also receive input from a file stream called *stdin* which is short for "standard input."

If you have data in a file that you want to send into standard input for a utility, you can use the < like this:

command < file

But, since most Unix utilities also let you specify the file as an argument, this is not used very much.

However, what is used all the time in Unix, and it is one of the things that makes it super fun, is the pipe, I, which says, "take *stdout* coming out of the command on the left and redirect it into *stdin* going into the command on the right of the pipe.

For example, if I wanted to count the number of files and directories stored in my git-repos directory, I could do

```
% ls -dl Documents/git-repos/* | wc 174 1566 14657
```

which pipes the output of ls -dl (one line per file) into the *stdin* for the wc command, which counts the number of lines, words, and letters sent to its standard input. So, the output tells me that there are 174 files and directories in my directory Documents/git-repos.

Note that pipes and redirects can be combined in sequence over multiple operations or commands. This is what gives rise to the terminology of making "Unix pipelines:" the data are like streams of water coming into or out of different commands, and the pipes hook up all those streams into a pipeline.

2.4.6 stderr

While output from Unix commands is often written to *stdout*, if anything goes wrong with a program, then messages about that get written to a different stream called *stderr*, which, you guessed it! is short for "standard error". By default, both *stdout* and *stderr* get written to the terminal, which is why it can be hard for beginners to think of them as separate streams.

But, indeed, they are. Redirecting stdout with >, that does **not** redirect stderr.

For example. See what happens when we ask ls to list a file that does not exist:

```
[~]--% ls file-not-here.txt
ls: file-not-here.txt: No such file or directory
```

The error message comes back to the screen. If you redirect the output it still comes back to the screen!

```
[~]--% ls file-not-here.txt > out.txt
ls: file-not-here.txt: No such file or directory
```

If you want to redirect *stderr*, then you need to specify which stream it is. On all Unix systems, *stderr* is stream #2, so the 2> syntax can be used:

```
[~]--% ls file-not-here.txt 2> out.txt
```

Then there is no output of *stderr* to the terminal, and when you cat the output file, you see that it went there!

```
[~]--% cat out.txt
ls: file-not-here.txt: No such file or directory
```

Doing bioinformatics, you will find that there will be failures of various programs. It is essential when you write bioinformatic pipelines to redirect *stderr* to a file so that you can go back, after the fact, to sleuth out why the failure occurred. Additionally, some bioinformatic programs write things like progress messages to *stderr* so it is important to know how to redirect those as well.

2.4.7 Symbolic links

Besides regular files and directories, a third type of file in Unix is called a *symbolic link*. It is a special type of file whose contents are just an absolute or

a relative path to another file. You can think of symbolic links as "shortcuts" to different locations in your file system. There are many useful applications of symbolic links.

Symbolic links are made using the ln command with the -s option. For example, if I did this in my home directory:

```
[~]--% ln -s /Users/eriq/Documents/git-repos/srsStuff srs
```

then srs becomes a file whose full listing (from ls -l srs) looks like:

```
lrwxrwxr-x 1 eriq staff 40B Jan 9 19:24 srs@ -> /Users/eriq/Documents/git-repos/srsS
```

2.4.8 File Permissions

Unix systems often host many different users. Some users might belong to the same research group, and might like to be able to read the files (and/or use the programs) that their colleagues have in their accounts.

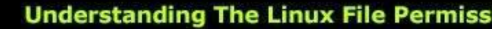
The Unix file system uses a system of permissions that gives rights to various classes of users to read, write, or execute files. The permissions associated with a file can be viewed using ls -1. They are captured in the first column which might look something like -rwxr-xr-x. When you first start looking at these, they can be distressingly difficult to visually parse. But you will get better at it! Let's start breaking it down now.

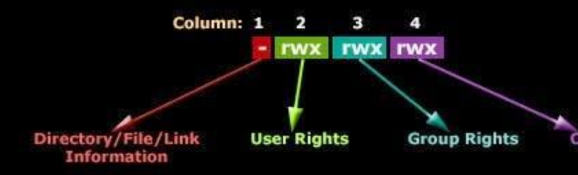
The file description string, in a standard Unix setting, consists of 10 characters.

- The first tells what kind of file it is: = regular file, d = directory, 1 = symbolic link.
- The next group of three characters denote whether the owner/user of the file has permission to either read, write, or execute the file.
- The following two groups of three characters are the same thing for users within the users group, and for all other users, respectively.

Here is a figure from the web¹ that we can talk about:

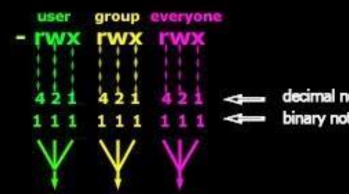
 $^{^{1}} https://unix.stackexchange.com/questions/183994/understanding-unix-permissions-and-file-types$





While the first column defines a directory, file or link, the next 3 column the permissions for the User, Group and Others (everyone else) group

Linux Permissions Made Easy



decimal notification: add each number to obtain the value (4 + 2 + 1 = 7)

binary notification: convert it to decimal then you should have the value (r-x = 101 b

Permissions can be changed with the chmod command. We will talk in class about how to use it with the octal representation of permissions.

2.4.9 Editing text files at the terminal

Sometimes you might need to edit text files at the command line.

The easiest text editor to use on the command line is nano. Try typing nano a-file.txt and type a few things. It is pretty self explanatory.

2.5 Customizing your Environment

Previously we saw how to modify your command prompt to tell you what the current working directory is (remember PS1='[\W]--%'). The limitation of giving that command on the command line is that if you logout and then log back in again, or open a new Terminal window, you will have to reissue that command in order to achieve the desired look of your command prompt. Quite often a Unix user would like to make a number of customization to the look, feel, and behavior of their Unix shell. The bash shell allows these customization to be specified in two different files that are read by the system so as to invoke the customization. The two files are hidden files in the home directory: ~/.bashrc and ~/.bash_profile. They are used by the Unix system in two slightly different contexts, but for most purposes, you, the user, will not need or even want to distinguish between the different contexts. Managing two separate files of customization is unnecessary and requires duplication of your efforts, and can lead to inconsistent and confusing results, so here is what we will do:

- 1. Keep all of our customization in ~/.bashrc.
- 2. Insert commands in ~/.bash_profile that say, "Hey computer! If you are looking for customization in here, don't bother, just get them straight out of ~/.bashrc.

We take care of #2, by creating the file ~/.bash_profile to have the following lines in it:

```
if [ -f ~/.bashrc ]; then
    source ~/.bashrc
fi
```

Taking care of #1 is now just a matter of writing commands into ~/.bashrc. In the following are some recommended customization.

2.5.1 Appearances matter

Some customization just change the way your shell looks or what type of output is given from different commands. Here are some lines to add to your ~/.bashrc along with some discussion of each.

```
export PS1='[\W]--% '
```

This gives a tidier and more informative command prompt. The export command before it tells the system to pass the value of this *environment variable*, PS1, along to any other shells that get spawned by the current one.

```
alias ls='ls -GFh'
```

This makes it so that each time you invoke the ls command, you do so with the options -G, -F, and -h. To find out on your own what those options do, you can type man ls at the command line and read the output, but briefly: -G causes directories and different file types to be printed in different colors, -F causes a / to be printed after directory names, and other characters to be printed at the end of the names of different file types, and -h causes file sizes to be printed in an easily human-readable form when using the -1 option.

2.5.2 Where are my programs/commands at?!

We saw in Section 2.3.1 that bash searches the directories listed in the PATH variable to find commands and executables. You can modify the PATH variable to include directories where you have installed different programs. In doing so, you want to make sure that you don't lose any of the other directories in PATH, so there is a certain way to go about redefining PATH. If you want to add the path /a-new/program/directory to your PATH variable you do it like this:

PATH=\$PATH:/a-new/program/directory

2.6 A Few More Important Keystrokes

If a command "gets stuck" or is running longer than it should be, you can usually kill/quit it by doing cntrl-c.

Once you have given a command, it gets stored in your bash history. You can use the up-arrow key to cycle backward through different commands in your history. This is particularly useful if you are building up complex pipelines on the command line piece by piece, looking at the output of each to make sure it is correct. Rather than re-typing what you did for the last command line, you just up-arrow it.

Once you have done an up-arrow or two, you can cycle back down through your history with a down-arrow.

Finally, you can search through your bash history by typing cntrl-r and then typing the word/command you are looking for. For example, if, 100 command lines back, you used a command that involved the program awk, you can search for that by typing cntrl-r and then typing awk.

One last big thing to note: the # is considered a *comment character* in bash. This means that any text following a # (unless it is backslash-escaped or inside quotation marks), until the next line ending, will be ignored by the shell.

2.7 A short list of additional useful commands.

Everyone should be familiar with the following commands, and the options that follow them on each line below. One might even think of scanning the manual page for each of these:

- echo
- cat
- head, -n, -c
- tail, -n
- less
- sort, -n -b -k
- paste
- cut, -d
- tar, -cvf, -xvf
- gzip, -c
- du, -h -C,

- W(
- date
- uniq
- chmod, u+x, ug+x
- grep

2.8 Two important computing concepts

2.8.1 Compression

Most file storage types (like text files) are a bit wasteful in terms of file space: every character in a text file takes the same number of bytes to store, whether it is a character that is used a lot, like s or e, or whether it is a character that is seldom seen in many text files, like ^. Compression is the art of creating a code for different types of data that uses fewer bits to encode "letters" (or "chunks" of data) that occur frequently and it reserves codewords of more bits to encode less frequently occurring chunks in the data. The result is that the total file size is smaller than the uncompressed version. However, in order to read it, the file must be decompressed.

In bioinformatics, many of the files you deal with will be compressed, because that can save many terabytes of disk space. Most often, files will be compressed using the gzip utility, and they can be uncompressed with the gunzip command. Sometimes you might want to just look at the fist part of a compressed file. If the file is compressed with gzip, you can decompress to stdout by using gzcat and then pipe it to head, for example.

A central form of compression in bioinformatics is called bgzip compression which compresses files into a series of blocks of the same size, situated in such a way that it is possible to *index* the contents of the file so that certain parts of the file can be accessed without decompressing the whole thing. We will encounter indexed compressed files a lot when we start dealing with BAM and vcf.gz files.

2.8.2 Hashing

The final topic we will cover here is the topic of hashing, an in particular the idea of "fingerprinting" files on ones computer. This process is central to how the git version control system works, and it is well worth knowing about.

Any file on your computer can be thought of as a series of bits, 0's and 1's, as,

fundamentally, that is what the file is. A *hashing algorithm* is an algorithm that maps a series of bits (or arbitrary length) to a short sequence of bits. The SHA1 hashing algorithm maps arbitrary sequences of bits to a sequence of 160 bits.

There are $2^{160} \approx 1.46 \times 10^{48}$ possible bit sequences of length 160. That is a vast number. If your hashing algorithm is well randomized, so that bit sequences are hashed into 160 bits in a roughly uniform distribution, then it is exceedingly unlikely that any two bit sequences (i.e. files on your filesystem) will have the same hash ("fingerprint") unless they are perfectly identical. As hashing algorithms are often quite fast to compute, this provides an exceptionally good way to verify that two files are identical.

The SHA1 algorithm is implemented with the shasum command. In the following, as a demonstration, I store the recursive listing of my git-repos directory into a file and I hash it. Then I add just a single line ending to the end of the file, and hash that, to note that the two hashes are not at all similar even though the two files differ by only one character:

```
[~]-% ls -R Documents/git-repos/* > /tmp/gr-list.txt
[~]-% # how many lines is that?
[~]-% wc /tmp/gr-list.txt
93096 88177 2310967 /tmp/gr-list.txt
[~]-% shasum /tmp/gr-list.txt
1396f2fec4eebdee079830e1eff9e3a64ba5588c /tmp/gr-list.txt
[~]-% # now add a line ending to the end
[~]-% (cat /tmp/gr-list.txt; echo) > /tmp/gr-list2.txt
[~]-% # hash both and compare
[~]-% shasum /tmp/gr-list.txt /tmp/gr-list2.txt
1396f2fec4eebdee079830e1eff9e3a64ba5588c /tmp/gr-list.txt
23bff8776ff86e5ebbe39e11cc2f5e31c286ae91 /tmp/gr-list2.txt
[~]-% # whoa! cool.
```

2.9 Unix: Quick Study Guide

This is just a table with quick topics/commands/words in it. You should understand each and be able to tell a friend a lot about each one. Cite it as 2.1

TABLE 2.1: Terms/ideas/etc. to know forward and backward

1 1.	- 1 14 41-	1. 4: 41.
bash	absolute path	relative path
	and a second power	Pooli

TABLE 2.1: Terms/ideas/etc. to know forward and backward

/ at beginning of path	/ between directories	home directory
~	current working directory	pwd
cd		
cd -	basename	PS1
TAB-completion	ls(-a, -d, -R)	globbing
*	?	[0-9]
[a-z]	[^CDcd]	${png,jpg,pdf}$
echo	man command	mkdir
mv	ср	rm
cat	head	less
stdout	stdin	stderr
>	<	
ln -s	symbolic link	PATH
-rw-rr	.bashrc	$.$ bash_profile
sort, -n -b -k	paste	cut, -d
tar, -cvf, -xvf	gzip	$\mathtt{du}, \mathtt{-h} \mathtt{-C},$
WC	date	uniq
cntrl-c	cntrl-r	#
$\operatorname{up-arrow/down-arrow}$	chmod, ug+x, 664	grep

Shell programming

Discuss the programming interface, and also maybe discuss & and ; and how to put things into scripts.

In here, let's also talk about compression with gzip (and using $stuff \mid gzip -c > this.gz$ to gzip and send to stdout.)

3.1 Advanced repitition

I want to get constructs like $\{1..20\}$ and $\{csv,pdf,jpg\}$ in here too.

3.2 Variables

3.3 looping

3.4 Further reading

An excellent chapter on the development of Unix (Raymond, 2003)

3.5 reading files line by line

This is handy. Note the line can be broken into a shell array:

```
cat bwa-run-list.txt straggler-bwa-run-list.txt | while read -r line; do
 A=($line);
 file=${A[1]};
 num=${A[2]};
 du -h bam-slices/$file/${num}-sorted.bam;
done
2.3G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
2.2G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
2.2G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
2.3G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
2.2G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
2.2G
        bam-slices/chinook_Battle.Creek.Sacramento.River_Schluter_GBC_001_CH1-2011_Male/00
```

Note that this is not how you want to rip through files, typically, because it is slow and awk is a much better bet. But, if you want to do a system call for each line, it ends up being a decent way forward.

3.6 Difference between double and single quotes

This becomes important when writing awk scripts and using beftools expressions.

Sed, awk, and regular expressions

In the course of doing bioinformatics, you will be dealing with myriad different text files. As we noted in the previous chapters, Unix, with its file I/O model, piping capabilities, and numerous utilities, is well-suited to handling large text files. Two utilities found on every Unix installation—awk and sed—merit special attention in this context. awk is a lightweight scripting language that lets you write succinct programs to operate line-by-by line on the contents of text files. It is particularly useful for handling text files that have columns of data separated by white spaces or tabs. sed on the other hand, is particularly useful for automating "find-and-replace" operations on text files. Each of them is optimized to handle large files without storing a lot of information in memory, so they can be useful for quick operations on large bioinformatic data sets. Neither is a fully-featured programming language that you would want to write large, complex programs in; however they do share many of the useful text-manipulation capabilities of such languages, such as Perl and Python. Additionally, awk and sed are deployed in a consistent fashion across most Unix operating systems, and they don't require much time to learn to use effectively for common text-processing tasks. As a consequence awk and sed are a useful addition to the bioinformatician's tookbox.

Both awk and sed rely heavily on regular expressions to describe patterns in text upon which some operation should be performed. You can think of regular expressions as providing a succinct language for performing very advanced "find" operations in a text file.

High Performance Computing (HPC) Environments

Hey Eric! You might consider breaking this into two separate chatpers: 1 = working on remote computers and 2 = high-performance computing. The first could include all the stuff about scp, globus, rclone, and google drive.

This is going to be a chapter on using High Performance Computing clusters.

There is a lot to convey here about using queues and things.

I know SGE pretty well at this point, but others might use slurm.

Here is a good page with a comparison: https://confluence.csiro.au/display/SC/Reference+Guide%3A+Migrating+from+SGE+to+SLURM

And here is a good primer on SGE stuff: https://confluence.si.edu/display/HPC/Monitoring+your+Jobs

I guess I'll have to see what the CSU students have access to.

5.1 Accessing remote computers

Start off with some stuff about ssh and scp.

Maybe have a section on public and private keys so you don't have to put your password in every time (I would like to get better with that, as well).

Here is the deal on that. On a mac, you can use

```
ssh-keygen -t rsa -b 4096
```

and be sure to give it a password, so that your private key is not unencrypted. For, if it is, then there is a chance (I do believe) that if someone were to obtain that file, they could gain access to all the computers you are authorized on.

Note that a lot of tutorials on the web have you generating a private key without any encryption. That is lame.

Then copy the .ssh/id_rsa.pub key to the .ssh/authorized_keys file on the server (creating it if it needs to be there). Then ssh to the server and your Mac will pop up a window asking for a password or will prompt on the command line for one (note that the password stays local!). You put in the password that you used when you created the private key. That can be saved in the Mac keychain (on old version of OSX you might get asked if you want to save it in the Mac keychain). Voila! Now you have access to the server with no need to type a password in there.

Note, since Sierra, you need to add this to a .ssh/config file to get the password stored in the keychain:

Host * AddKeysToAgent yes UseKeychain yes

And, it turns out that you also need to make sure the permissions on that file are set appropriately:

```
chmod 600 ~/.ssh/config
```

It is actually pretty darn simple...

Note, to have access from another computer to the server, you probably just create a keypair for that computer, and add the public key to the authorized_keys.

Then add something like this to your .bashrc:

```
# for quick ssh to hoffman
alias hoffy='ssh eriq@hoffman2.idre.ucla.edu'
# to be used in scp. like scp file $hoff:~/
hoff=eriq@hoffman2.idre.ucla.edu
```

Note that this should only be done on a private computer account, not on a shared account on a computer. Otherwise, everyone using that account will have access.

Also, an interesting thing to investigate might be SSHF/FUSE which just might let one mount the HPC filesystem on a local directory, so you can work with files there using RStudio, etc, get them all debugged, and then run them.

Check out some information about that here: https://www.digitalocean.com/community/tutorials/how-to-use-sshfs-to-mount-remote-file-systems-over-ssh.

I went to https://osxfuse.github.io/ and I downloaded SSHFS 2.5.0 and

FUSE for Mac OS X. Then I installed them, doing a simple default install for each.

Then, you make a mountpoint, which you can do in your home directory if you want:

```
mkdir ~/hoffman # this is the mountpoint
sshfs -o allow_other,defer_permissions,IdentityFile=~/.ssh/id_rsa
# close it out:
umount ~/hoffman/
```

Note that the absolute path to my home folder there seemed to be important.

Also, I can't get to /u/nobackup/kruegg/eriq from here by going through the symlink in my home directory on hoffman, since that is essentially a different volume. But I should be able to do this:

```
sshfs -o allow_other,defer_permissions,IdentityFile=~/.ssh/id_rsa eriq@hoffman2.idre.ucla.
```

Yep! that works. And I can open all those files from within RStudio. Cool.

Note, I will have to have a separate mount point for \$SCRATCH as well.

Shoot! That works really seamlessly!

I can also add to my .bashrc:

```
alias hoffuse='sshfs -o allow_other,defer_permissions,IdentityFile=~/.ssh/id_rsa eriq@hoff
```

Now, the bad news: you can open an Rstudio project from that remote, mounted volume, but it doesn't seem to really work. It is incredibly slow, and then fails to quit properly, etc.

After force-quitting it, I am unable to unmount the hoffuse volume. What a mess. OK, I probably won't try (running the rstudio project over FUSE). But it might be useful still for editing small things.

5.2 Transferring files to remote computers

5.2.1 scp

5.2.2 Globus

5.2.3 Interfacing with "The Cloud"

Increasingly, data scientists and tech companies alike are keeping their data "in the cloud." This means that they pay a large tech firm like Amazon, Dropbox, or Google to store their data for them in a place that can be accessed via the internet. There are many advantages to this model. For one thing, the company that serves the data often will create multiple copies of the data for backup and redundancy: a fire in a single data center is not a calamity because the data are also stored elsewhere, and can often be accessed seamlessly from those other locations with no apparent disruption of service. For another, companies that are in the business of storing and serving data to multiple clients have data centers that are well-networked, so that getting data onto and off of their storage systems can be done very quickly over the internet by an end-user with a good internet connection.

Five years ago, the idea of storing next generation sequencing data might have sounded a little crazy—it always seemed a laborious task getting the data off of the remote server at the sequencing center, so why not just keep the data inhouse once you have it? To be sure, keeping a copy of your data inhouse still can make sense for long-term data archiving needs, but, today, cloud storage for your sequencing data can make a lot of sense. A few reasons are:

- 1. Transferring your data from the cloud to the remote HPC system that you use to process the data can be very fast.
- 2. As above, your data can be redundantly backed up.
- 3. If your institution (university, agency, etc.) has an agreement with a cloud storage service that provides you with unlimited storage and free network access, then storing your sequencing data in the cloud will cost considerably less than buying a dedicated large system of hard drives for data backup. (One must wonder if service agreements might not be at risk of renegotiation if many researchers start using their unlimited institutional cloud storage space to store and/or archive their next generation sequencing data sets. My own agency's contract with Google runs through 2021...but I have to think that these services are making plenty of money, even if a handful of researchers store big sequence data in the cloud. Nonetheless, you

- should be careful not to put multiple copies of data sets, or intermediate files that are easily regenerated, up in the cloud.)
- 4. If you are a PI with many lab members wishing to access the same data set, or even if you are just a regular Joe/Joanna researcher but you wish to share your data, it is possible to effect that using your cloud service's sharing settings. We will discuss how to do this with Google Drive.

There are clearly advantages to using the cloud, but one small hurdle remains. Most of the time, working in an HPC environment, we are using Unix, which provides a consistent set of tools for interfacing with other computers using SSH-based protocols (like scp for copying files from one remote computer to another). Unfortunately, many common cloud storage services do not offer an SSH based interface. Rather, they typically process requests from clients using an HTTPS protocol. This protocol, which effectively runs the world-wide web, is a natural choice for cloud services that most people will access using a web browser; however, Unix does not traditionally come with a utility or command to easily process the types of HTTPS transactions needed to network with cloud storage. Furthermore, there must be some security when it comes to accessing your cloud-based storage—you don't want everyone to be able to access your files, so your cloud service needs to have some way of authenticating people (you and your labmates for example) that are authorized to access your data.

These problems have been overcome by a utility called rclone, the product of a comprehensive open-source software project that brings the functionality of the rsync utility (a common Unix tool used to synchronize and mirror file systems) to cloud-based storage. (Note: rclone has nothing to do with the R programming language, despite its name that looks like an R package.) Currently rclone provides a consistent interface for accessing files from over 35 different cloud storage providers, including Box, Dropbox, Google Drive, and Microsoft OneDrive. Binaries for rclone can be downloaded for your desktop machine from https://rclone.org/downloads/. We will talk about how to install it on your HPC system later.

Once rclone is installed and in your PATH, you invoke it in your terminal with the command rclone. Before we get into the details of the various rclone subcommands, it will be helpful to take a glance at the information rclone records when it configures itself to talk to your cloud service. To do so, it creates a file called ~/.config/rclone/rclone.conf, where it stores information about all the different connections to cloud services you have set up. For example, that file on my system looks like this:

[gdrive-rclone] type = drive scope = drive root_folder_id = 1I2EDV465N5732Tx1FFAiLWOqZRJcAzUd
token = {"access_token":"bs43.94cUF0e6SjjkofZ","token_type":"Bearer","refresh_token":"1/Mr
client_id = 2934793-oldk97lhld88dlkh301hd.apps.googleusercontent.com
client_secret = MMq3jdsjdjgKTGH4rNV_y-NbbG

In this configuration:

- gdrive-rclone is the name by which rclone refers to this cloud storage location
- root_folder_id is the ID of the Google Drive folder that can be thought of as the root directory of gdrive-rclone. This ID is not the simple name of that directory on your Google Drive, rather it is the unique name given by Google Drive to that directory. You can see it by navigating in your browser to the directory you want and finding it after the last slash in the URL. For example, in the above case, the URL is: https://drive.google.com/drive/u/1/folders/1I2EDV465N5732Tx1FFAiLWOqZRJcAzUd
- client_id and client_secret are like a username and a shared secret that
 rclone uses to authenticate the user to Google Drive as who they say they
 are.
- token are the credentials used by rclone to make requests of Google Drive on the basis of the user.

Note: the above does not include my real credentials, as then anyone could use them to access my Google Drive!

To set up your own configuration file to use Google Drive, you will use the rclone config command, but before you do that, you will want to wrangle a client_id from Google. Follow the directions at https://rclone.org/drive/#making-your-own-client-id. Things are a little different from in their step by step, but you can muddle through to get to a screen with a client_ID and a client secret that you can copy onto your clipboard.

Once you have done that, then run rclone config and follow the prompts. A typical session of rclone config for Google Drive access is given here¹. Don't choose to do the advanced setup; however do use "auto config," which will bounce up a web page and let you authenticate rclone to your Google account.

5.2.3.1 Basic Maneuvers

The syntax for use is:

```
rclone [options] subcommand parameter1 [parameter 2...]
```

The "subcommand" part tells rclone what you want to do, like copy or sync,

¹https://rclone.org/drive/

and the "parameter" part of the above syntax is typically a path specification to a directory or a file. In using rclone to access the cloud there is not a root directory, like / in Unix. Instead, each remote cloud access point is treated as the root directory, and you refer to it by the name of the configuration followed by a colon. In our example, gdrive-rclone: is the root, and we don't need to add a / after it to start a path with it. Thus gdrive-rclone:this_dir/that_dir is a valid path for rclone to a location on my Google Drive.

Very often when moving, copying, or syncing files, the parameters consist of:

source-directory destination-directory

One very important point is that, unlike the Unix commands cp and mv, rclone likes to operate on directories, not on multiple named files.

A few key subcommands:

• ls, lsd, and lsl are like ls, ls -d and ls -l

```
rclone lsd gdrive-rclone:
rclone lsd gdrive-rclone:NOFU
```

• copy: copy the *contents* of a source *directory* to a destination *directory*. One super cool thing about this is that rclone won't re-copy files that are already on the destination and which are identical to those in the source directory.

```
rclone copy bams gdrive-rclone: NOFU/bams
```

Note that the destination directory will be created if it does not already exist. - sync: make the contents of the destination directory look just like the contents of the source directory. WARNING This will delete files in the destination directory that do not appear in the source directory.

A few key options:

- --dry-run: don't actually copy, sync, or move anything. Just tell me what you would have done.
- -P, -v, -vv: give me progress information, verbose output, or super-verbose output, respectively.
- --tpslimit 10: don't make any more than 10 transactions a second with Google Drive (should always be used when transferring files)
- ---fast-list: combine multiple transactions together. Should always be used with Google Drive, especially when handling lots of files.
- --drive-shared-with-me: make the "root" directory a directory that shows all of the Google Drive folders that people have shared with you. This is key for accessing folders that have been shared with you.

For example, try something like:

```
rclone --drive-shared-with-me lsd gdrive-rclone:
```

5.2.3.2 filtering: Be particular about the files you transfer

rclone works a little differently than the Unix utility cp. In particular, rclone is not set up very well to copy individual files. While there is a an rclone command known as copyto that will allow you copy a single file, you cannot (apparently) specify multiple, individual files that you wish to copy.

In other words, you can't do:

```
rclone copyto this_file.txt that_file.txt another_file.bam gdrive-rclone:dest_dir
```

In general, you will be better off using rclone to copy the *contents* of a directory to the inside of the destination directory. However, there are options in rclone that can keep you from being totally indiscriminate about the files you transfer. In other words, you can *filter* the files that get transferred. You can read about that at https://rclone.org/filtering/.

For a quick example, imagine that you have a directory called Data on you Google Drive that contains both VCF and BAM files. You want to get only the VCF files (ending with .vcf.gz, say) onto the current working directory on your cluster. Then something like this works:

```
rclone copy --include *.vcf.gz gdrive-rclone:Data ./
```

Note that, if you are issuing this command on a Unix system in a directory where the pattern *.vcf.gz will expand (by globbing) to multiple files, you will get an error. In that case, wrap the pattern in a pair of single quotes to keep the shell from expanding it, like this:

```
rclone copy --include '*.vcf.gz' gdrive-rclone:Data ./
```

5.2.3.3 Feel free to make lots of configurations

You might want to configure a remote for each directory-specific project. You can do that by just editing the configuration file. For example, if I had a directory deep within my Google Drive, inside a chain of folders that looked like, say, Projects/NGS/Species/Salmon/Chinook/CentralValley/WinterRun

where I was keeping all my data on a project concerning winterrun Chinook salmon, then it would be quite inconvenient to type Projects/NGS/Species/Salmon/Chinook/CentralValley/WinterRun every time I wanted to copy or sync something within that directory. Instead, I could add the following lines to my configuration file, essentially copying the existing configuration and then modifying the configuration name and the root_folder_id to be the Google Drive identifier for the folder Projects/NGS/Species/Salmon/Chinook/CentralValley/WinterRun (which one can find by navigating to that folder in a web browser and pulling the ID from the end of the URL.) The updated configuration could look like:

```
[gdrive-winter-run]
type = drive
scope = drive
root_folder_id = 1MjOrclmP1udhxOTvLWDHFBVET1dF6CIn
token = {"access_token":"bs43.94cUF0e6SjjkofZ","token_type":"Bearer","refresh_token":"1/Mr
client_id = 2934793-oldk97lhld88dlkh301hd.apps.googleusercontent.com
client_secret = MMq3jdsjdjgKTGH4rNV_y-NbbG
```

As long as the directory is still within the same Google Drive account, you can re-use all the authorization information, and just change the [name] part and the root_folder_id. Now this:

```
rclone copy src_dir gdrive-winter-run:
```

puts items into Projects/NGS/Species/Salmon/Chinook/CentralValley/WinterRun on the Google Drive without having to type that God-awful long path name.

5.2.3.4 Installing relone on a remote machine without sudo access

The instructions on the website require root access. You don't have to have root access to install relone locally in your home directory somewhere. Copy the download link from https://rclone.org/downloads/ for the type of operating system your remote machine uses (most likely Linux if it is a cluster). Then transfer that with wget, unzip it and put the binary in your PATH. It will look something like this:

```
wget https://downloads.rclone.org/rclone-current-linux-amd64.zip
unzip rclone-current-linux-amd64.zip
cp rclone-current-linux-amd64/rclone ~/bin
```

You won't get manual pages on your system, but you can always find the docs on the web.

5.2.3.5 Setting up configurations on the remote machine...

Is as easy as copying your config file to where it should go, which is easy to find using the command:

rclone config file

5.2.3.6 Encrypting your config file

This makes sense, and it easy: with rclone config encryption is one of the options. When it is encrypted, use rclone config show to see what it looks like in clear text.

5.2.3.7 Some other usage tips

Following an email exchange with Ren, I should mention how to do an md5 checksum on the remote server to make sure that everything is correctly there.

5.2.4 Getting files from a sequencing center

Very often sequencing centers will post all the data from a single run of a machine at a secured (or unsecured) http address. You will need to download those files to operate on them on your cluster or local machine. However some of the files available on the server will likely belong to other researchers and you don't want to waste time downloading them.

Let's take an example. Suppose you are sent an email from the sequencing center that says something like:

Your samples are AW_F1 (female) and AW_M1 (male). You should be able to access the data from this link provided by YCGA: http://sysg1.cs.yale.edu: 3010/5ln09bs3zfa8L0hESfsYfq3Dc/061719/

You can easily access this web address using rclone. You could set up a new remote in your rclone config to point to http://sysgl.cs.yale.edu, but, since you will only be using this once, to get your data, it makes more sense

to just specify the remote on the command line. This can be done by passing rclone the URL address via the --http-url option, and then, after that, telling it what protocol to use by adding :http: to the command. Here is what you would use to list the directories available at the sequencing center URL:

```
# here is the command
% rclone lsd --http-url http://sysg1.cs.yale.edu:3010/5ln09bs3zfa8L0hESfsYfq3Dc/061719/ :h
# and here is the output
-1 1969-12-31 16:00:00 -1 sjg73_fqs
-1 1969-12-31 16:00:00 -1 sjg73_supernova_fqs
```

Aha! There are two directories that might hold our sequencing data. I wonder what is in those directories? The rclone tree command is the perfect way to drill down into those directories and look at their contents:

```
\% \ \ \text{rclone tree --http-url http://sysg1.cs.yale.edu:} 3010/5ln09bs3zfa8L0hESfsYfq3Dc/061719/: 3010/5ln09bs3zfa8L0hESfsYfq3Dc/06170000000000000000000000000000000
          sjg73_fqs
                         AW_F1
                                       AW_F1_S2_L001_I1_001.fastq.gz
                                       AW_F1_S2_L001_R1_001.fastq.gz
                                       {\tt AW\_F1\_S2\_L001\_R2\_001.fastq.gz}
                         AW_M1
                                       AW_M1_S3_L001_I1_001.fastq.gz
                                       AW_M1_S3_L001_R1_001.fastq.gz
                                       AW_M1_S3_L001_R2_001.fastq.gz
                        ESP A1
                                         ESP_A1_S1_L001_I1_001.fastq.gz
                                         ESP_A1_S1_L001_R1_001.fastq.gz
                                         ESP_A1_S1_L001_R2_001.fastq.gz
          sjg73_supernova_fqs
                           AW_F1
                                         AW_F1_S2_L001_I1_001.fastq.gz
                                         AW_F1_S2_L001_R1_001.fastq.gz
                                         AW_F1_S2_L001_R2_001.fastq.gz
                           AW_M1
                                         AW_M1_S3_L001_I1_001.fastq.gz
                                         AW_M1_S3_L001_R1_001.fastq.gz
                                         AW_M1_S3_L001_R2_001.fastq.gz
                          ESP_A1
                                           ESP_A1_S1_L001_I1_001.fastq.gz
                                           ESP_A1_S1_L001_R1_001.fastq.gz
```

```
ESP_A1_S1_L001_R2_001.fastq.gz
8 directories, 18 files
```

Whoa! That is pretty cool!. From this output we see that there are subdirectories named AW_F1 and AW_M1 that hold the files that we want. And, of course, the ESP_A1 samples must belong to someone else. It would be great if we could just download the files we wanted, excluding the ones in the ESP_A1 directories. It turns out that there is! rclone has an --exclude option to exclude paths that match certain patterns (see Section 5.2.3.2, above). We can experiment by giving rclone copy the --dry-run command to see which files will be transferred. If we don't do any filtering, we see this when we try to dry-run copy the directories to our local directory Alewife/fastqs:

```
% rclone copy --dry-run --http-url http://sysg1.cs.yale.edu:3010/5ln09bs3zfa8L0hESfsYfq3Dc
2019/07/11 10:33:43 NOTICE: sjg73_fqs/ESP_A1/ESP_A1_S1_L001_I1_001.fastq.gz: Not copying a
2019/07/11 10:33:43 NOTICE: sjg73_fqs/ESP_A1/ESP_A1_S1_L001_R1_001.fastq.gz: Not copying a
2019/07/11 10:33:43 NOTICE: sjg73_fqs/ESP_A1/ESP_A1_S1_L001_R2_001.fastq.gz: Not copying a
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_I1_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_R2_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_I1_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_R1_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_R2_001.fastq.gz: Not c
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/ESP_A1/ESP_A1_S1_L001_I1_001.fastq.gz: Not
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/ESP_A1/ESP_A1_S1_L001_R1_001.fastq.gz: Not
2019/07/11 10:33:43 NOTICE: sjg73_supernova_fqs/ESP_A1/ESP_A1_S1_L001_R2_001.fastq.gz: Not
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_I1_001.fastq.gz: Not copying as
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_R1_001.fastq.gz: Not copying as
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_R2_001.fastq.gz: Not copying as
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_I1_001.fastq.gz: Not copying as
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz: Not copying as
2019/07/11 10:33:43 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_R2_001.fastq.gz: Not copying as
```

Since we do not want to copy the ESP_A1 files we see if we can exclude them:

```
% rclone copy --exclude */ESP_A1/* --dry-run --http-url http://sysg1.cs.yale.edu:3010/5lnC2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_I1_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_R2_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_F1/AW_F1_S2_L001_R1_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_I1_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_R2_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz: Not copying as 2019/07/11 10:37:22 NOTICE: sjg73_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz: Not copying as
```

```
2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_I1_001.fastq.gz: Not of 2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_R2_001.fastq.gz: Not of 2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_F1/AW_F1_S2_L001_R1_001.fastq.gz: Not of 2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_I1_001.fastq.gz: Not of 2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz: Not of 2019/07/11 10:37:22 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_R2_001.fastq.gz: Not of 2019/07/11 NOTICE: sjg73_supernova_fqs/AW_M1/AW_M1_S3_L001_R1_001.fastq.gz:
```

Booyah! That gets us just what we want. So, then we remove the --dry-run option, and maybe add -v -P to give us verbose output and progress information, and copy all of our files:

```
% rclone copy --exclude */ESP_A1/* -v -P --http-url http://sysg1.cs.yale.edu:3010/5ln09bs
```

5.3 Activating/Installing software

5.3.1 Modules

This is if your sys admin has made it easy.

5.3.2 Miniconda

This is how one will probably want to do it

5.3.2.1 Testing this on Summit

I just want to quickly try this:

```
# get on compile nodes
ssh scompile

# I checked modules and found no samtools, bcftools, etc.

# Now install miniconda
mkdir conda_install
```

```
cd conda_install/
wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
chmod u+x Miniconda3-latest-Linux-x86_64.sh
./Miniconda3-latest-Linux-x86_64.sh

# then you follow the prompts and agree to the license and the
# default install location.

## NOTE: Might want to install to a different location if there
## are serious caps on hard disk usage in home directory..

source ~/.bashrc

# after that I have it!
(base) [eriq@colostate.edu@shas0136 conda_install]$
```

Now, let's see if we can get samtools. Just google "samtools conda" to get an idea of how to do it:

```
conda install -c bioconda samtools
```

AFter that, it just works! Cool!

All right! That is amazing. Next steps:

1. Tell students about establishing different environments.

2. Learn about how to make a minimal environment for a project and how to record that and be able to distribute/propagate it.

Along those lines, I want to see if, when I install samtools into a new environment, it re-downloads it or not...

```
conda create --name quick-test
conda activate quick-test

# i also made an environment in a specified directory (you
# could put these within a project directory)
conda create --prefix ./test-envs-dir
conda activate ./test-envs-dir

# now, let's install bcftools there
conda install -c bioconda bcftools

# note that it doesn't re-download the dependencies, as far as I can tell.
conda activate base
bcftools # not found in environment base

conda activate ./test-envs-dir
bcftools # it is found in this environment. Cool.

conda activate quick-test
bcftools # it ain't here
```

Now, after that, bcftools is in ./test-envs-dir/bcftools

bcftools

So, what if we install it into another environment. Does it symlink it?

```
conda activate base
conda install -c bioconda bcftools
bcftools

# whoa! That errored out!
bcftools: error while loading shared libraries: libcrypto.so.1.0.0: cannot open shared obj
# that is a serious problem.

# Can I get it in my other environment?
conda activate quick-test
conda install -c bioconda bcftools
```

```
# that totally works, but it still doesn't in base...
# so, what if we add samtools to our other environments?
# that works fine.
```

But, samtools/bcftools dependency issues are a known problem: https://github.com/sunbeam-labs/sunbeam/issues/181 Basically they rely on different versions of some ssh libs.

Note that installing beftools first things work. But what if we make another environment and install samtools first again?

```
conda create --name samtools-first
conda activate samtools-first
conda install -c bioconda samtools # this didn't download anything new
conda install -c bioconda bcftools

# BOOM! THIS CREATES A FAIL. SO, YOU GOTTA INSTALL BCFTOOLS
# FIRST. FAR OUT.
```

5.3.3 Exporting environments

Looks like we should be able to do this. Let's do the one that works:

```
(quick-test) [~]--% conda env export > quick-test-env.yml
(quick-test) [~]--% cat quick-test-env.yml
name: quick-test
channels:
  - bioconda
  - defaults
dependencies:
 - _libgcc_mutex=0.1=main
  - bcftools=1.9=ha228f0b_4
  - bzip2=1.0.8=h7b6447c_0
  - ca-certificates=2019.5.15=1
  - curl=7.65.3=hbc83047_0
  - htslib=1.9=ha228f0b_7
  - krb5=1.16.1=h173b8e3_7
  - libcurl=7.65.3=h20c2e04 0
  - libdeflate=1.0=h14c3975_1
```

5.4 Boneyard 57

```
- libedit=3.1.20181209=hc058e9b 0
 - libgcc-ng=9.1.0=hdf63c60_0
  - libssh2=1.8.2=h1ba5d50 0
 - libstdcxx-ng=9.1.0=hdf63c60_0
 - ncurses=6.1=he6710b0_1
 - openssl=1.1.1c=h7b6447c_1
 - samtools=1.9=h10a08f8_12
 - tk=8.6.8=hbc83047_0
 - xz=5.2.4=h14c3975_4
  - zlib=1.2.11=h7b6447c_3
prefix: /home/eriq@colostate.edu/miniconda3/envs/quick-test
# OK, that is cool. Now, if we wanted to email that to someone,
# we could, and then they could do this:
conda env create --name dupie-quick -f quick-test-env.yml
# that environment then has samtools and bcftools
# note that it probably would try to name it quick-test if we didn't
# pass in the name there...
```

5.4 Boneyard

This is a variant of rsync that let's you sync stuff up to google drive. It might be a better solution than rcp for getting stuff onto and off of google drive. Here is a link: https://rclone.org/. I need to evaluate it. It might also be a good way to backup some of my workstuff on my laptop to Google Drive (and maybe also for other people to create replicas and have a decent backup if they have unlimited Google Drive storage).

```
I got this working. It is important to set your own OAuth client ID: https://forum.rclone.org/t/very-slow-sync-to-google-drive/6903
```

After that I did like this:

```
rclone sync -vv --tpslimit 10 --fast-list 0tsh_v1.0_genomic.fna gdrive-rclone:spoogee-spowhich did 2 Gb of fasta into the spoogee-spoogee directory pretty quickly.
```

But, with something that has lots of files, it took longer:

```
# this is only about 100 Mb but took a long time
rclone copy -P --tpslimit 10 --fast-list rubias gdrive-rclone:rubias
```

However, once that is done, you can sync it and it finds that parts that have changed pretty quickly.

it appears to do that by file modification times:

```
2019-04-19 23:21 /Otsh_v1.0/--% rclone sync -vv --tpslimit 10 --fast-list Otsh_v1.0_genomi
2019/04/19 23:21:36 DEBUG: rclone: Version "v1.47.0" starting with parameters ["rclone" "
2019/04/19 23:21:36 DEBUG : Using config file from "/Users/eriq/.config/rclone/rclone.conf
2019/04/19 23:21:36 INFO : Starting HTTP transaction limiter: max 10 transactions/s with
2019/04/19 23:21:37 DEBUG: GCF_002872995.1_Otsh_v1.0_genomic.gff.gz: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.dict: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.amb: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.ann: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.bwt: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.fai: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.pac: Excluded
2019/04/19 23:21:37 DEBUG : Otsh_v1.0_genomic.fna.sa: Excluded
2019/04/19 23:21:37 INFO : Google drive root 'spoogee-spoogee': Waiting for checks to fir
2019/04/19 23:21:37 DEBUG: Otsh_v1.0_genomic.fna: Size and modification time the same (di
2019/04/19 23:21:37 DEBUG: Otsh_v1.0_genomic.fna: Unchanged skipping
2019/04/19 23:21:37 INFO : Google drive root 'spoogee-spoogee': Waiting for transfers to
2019/04/19 23:21:37 INFO : Waiting for deletions to finish
2019/04/19 23:21:37 INFO
Transferred:
                         0 / 0 Bytes, -, 0 Bytes/s, ETA -
                        0
Errors:
Checks:
                        1 / 1, 100%
Transferred:
                        0 / 0, -
                     1.3s
Elapsed time:
2019/04/19 23:21:37 DEBUG : 5 go routines active
2019/04/19 23:21:37 DEBUG : rclone: Version "v1.47.0" finishing with parameters ["rclone"
```

So, for moving big files around that might be a good way forward. I will have to do a test with some big files.

And I need to test it with team drives so that multiple individuals can pull stuff off the Bird Genoscape drive for example.

It would be nice to have safeguards so people don't trash stuff accidentally....

5.4 Boneyard 59

5.4.0.1 rclone on Hoffman

Their default install script expects sudo access to put it in /usr/local but I don't on hoffman, obviously, so I just downloaded the the install script and edited the section for Linux to look like this at the relevant part

```
case $0S in
  'linux')
  #binary
  cp rclone ~/bin/rclone.new
  chmod 755 ~/bin/rclone.new
  #chown root:root /usr/bin/rclone.new
  mv ~/bin/rclone.new ~/bin/rclone
  #manuals
  #mkdir -p /usr/local/share/man/man1
  #cp rclone.1 /usr/local/share/man/man1/
  #mandb
  ;;
```

I don't get man pages, but I get it in ~/bin no problem.

To set up the configuration, check where it belongs:

```
% rclone config file
Configuration file doesn't exist, but rclone will use this path:
/u/home/e/eriq/.config/rclone/rclone.conf
```

And then I just put my config file from my laptop on there. I just pasted the stuff in whilst emacsing it. Holy cow! That is super easy.

Note that the config file is where you can also set default options like tpslimit and fast-list I think.

So, the OAuth stuff is all stored in that config file. And if you can set it up on one machine you can go put it on any others that you want. That is awesome.

When it was done, I tested it:

```
% rclone sync -vv --drive-shared-with-me gdrive-rclone:BaselinePaper BaselinePaper_here 2019/04/29 14:49:24 DEBUG: rclone: Version "v1.47.0" starting with parameters ["rclone" "2019/04/29 14:49:24 DEBUG: Using config file from "/u/home/e/eriq/.config/rclone/rclone.co 2019/04/29 14:49:25 INFO: Local file system at /u/home/e/eriq/BaselinePaper_here: Waiting 2019/04/29 14:49:25 INFO: Local file system at /u/home/e/eriq/BaselinePaper_here: Waiting 2019/04/29 14:49:26 DEBUG: Local file system at /u/home/e/eriq/BaselinePaper_here: File to 2019/04/29 14:49:26 DEBUG: BaselinePaper_Body.docx: Failed to pre-allocate: operation not
```

```
2019/04/29 14:49:26 INFO : BaselinePaper_Body.docx: Copied (new)
2019/04/29 14:49:26 DEBUG: BaselinePaper_Body.docx: Updating size of doc after download t
2019/04/29 14:49:26 INFO : BaselinePaper_Body.docx: Copied (Rcat, new)
2019/04/29 14:49:27 DEBUG : Local file system at /u/home/e/eriq/BaselinePaper_here: File t
2019/04/29 14:49:27 DEBUG: ResponseToReviewers_eca.docx: Failed to pre-allocate: operation
2019/04/29 14:49:27 INFO : ResponseToReviewers_eca.docx: Copied (new)
2019/04/29 14:49:27 DEBUG: ResponseToReviewers_eca.docx: Updating size of doc after downl
2019/04/29 14:49:27 INFO : ResponseToReviewers_eca.docx: Copied (Rcat, new)
2019/04/29 14:49:27 INFO : Waiting for deletions to finish
2019/04/29 14:49:27 INFO
                  193.543k / 193.543 kBytes, 100%, 79.377 kBytes/s, ETA Os
Transferred:
Errors:
Checks:
                        0 / 0, -
Transferred:
                        4 / 4, 100%
Elapsed time:
                     2.4s
2019/04/29 14:49:27 DEBUG : 6 go routines active
2019/04/29 14:49:27 DEBUG: rclone: Version "v1.47.0" finishing with parameters ["rclone"
```

That was fast and super solid.

5.4.0.2 Encrypt the config file

You can use rclone config edit to set a password for the config file. Then it encrypts that so no one is able to run wild if they just get that file. You have to provide your password to do any of the rclone commands. If you want to see the config file use rclone config show. You could always copy that elsewhere, and then re-encrypt it.

Here is some nice stuff for summarizing all the information from the different runs from the chinook-wgs project:

```
qacct -o eriq -b 09271925 -j ml | tidy-qacct
```

Explain scratch space and how clusters are configured with respect to storage, etc.

Strategies—break names up with consistent characters:

- dashes within population names
- underscores for different groups of chromosomes
- periods for catenating pairs of pops

etc. Basically, it just makes it much easier to split things up when the time comes.

5.5 The Queue (SLURM/SGE/UGE)

5.6 Modules package

5.7 Compiling programs without admin privileges

Inevitably you will want to use a piece of software that is not available as a module or is not otherwise installed on they system.

Typically these software programs have a frightful web of dependencies.

Unix/Linux distros typically maintain all these dependencies as libraries or packages that can be installed using a rpm or yum. However, the simple "plugand-play" approach to using these programs requires have administrator privileges so that the software can be installed in one of the (typically protected) paths in the root (like /usr/bin).

But, you can use these programs to install packages into your home directory. Once you have done that, you need to let your system know where to look for these packages when it needs them (i.e., when running a program or *linking* to it whilst compiling up a program that uses it as a dependency.

Hoffman2 runs CentOS. Turns out that CentOS uses yum as a package manager.

Let's see if we can install llvm using yum.

```
yum search all llvm # <- this got me to devtoolset-7-all.x86_64: Package shipping all avoid # a little web searching made it look like llvm-toolset-7-5.0.1-4.el7.x86_64.rpm or devtoo # might be what we want. The first is a dependency of the second...

mkdir ~/centos
```

Was using instructions at https://stackoverflow.com/questions/36651091/how-to-install-packages-in-linux-centos-without-root-user-with-automatic-depen

Couldn't get yum downloader to download any packages. The whole thing looked like it was going to be a mess, so I thought I would try with miniconda.

I installed miniconda (python 2.7 version) into /u/nobackup/kruegg/eriq/programs/miniconda/ and then did this:

```
# probably could have listed them all at once, but wanted to watch them go
# one at a time...
conda install numpy
conda install scipy
conda install pandas
conda install numba

# those all ran great.

conda install pysnptools

# that one didn't find a match, but I found on the web that I should try:
conda install -c bioconda pysnptools

# that worked!
```

Also we want to touch briefly on LD_PATH (linking failures—and note that libraries are often named libxxx.a) and CPATH (for failure to find xxxx.h), etc.

5.8 Job arrays

Definitely mention the eval keyword in bash for when you want to print command lines with redirects.

Show the routine for it, and develop a good approach to efficiently orchestrating redos. If you know the taskIDs of the ones that failed then it is pretty easy to write an awk script that picks out the commands and puts them in a new file. Actually, it is probably better to just cycle over the numbers and use the -t option to launch each. Then there is now changing the job-ids file.

In fact, I am starting to think that the -t option is better than putting it into the file.

Question: if you give something on the command line, does that override the directive in the header of the file? If so, then you don't even need to change the file. Note that using the qsub command line options instead of the directives really opens up a lot of possibilities for writing useful scripts that are flexible.

Also use short names for the jobs and have a system for naming the redos (append numbers so you know which round it is, too) possibly base the name on the ways things failed the first time. Like, fsttf1 = "Fst run for things"

that failed due to time limits, 1". Or structure things so that redos can just be done by invoking it with -t and the jobid.

5.9 Writing stdout and stderr to files

This is always good to do. Note that stdbuf is super useful here so that things don't get buffered super long. (PCAngsd doesn't seem to write antyhing till the end...)

5.10 Breaking stuff down

It is probably worth talking about how problems can be broken down into smaller ones. Maybe give an example, and then say that we will be talking about this for every step of the way in bioinformatic pipelines.

One thing to note—sometimes processes go awry for one reason or another. When things are in smaller chunks it is not such a huge investment to re-run it. (Unlike stuff that runs for two weeks before you realize that it ain't working right).

Part II

Part II: Reproducible Research Strategies

$Introduction\ to\ Reproducible\ Research$

```
(Pritchard et al., 2000)
```

And let's again try to pitch it in there: (Pritchard et al., 2000). Let's try chucking it in without parens: Pritchard et al. (2000)

Rstudio and Project-centered Organization

Somewhere talk about here::here(): https://github.com/jennybc/here_here

7.1 Organizing big projects

By "big" I mean something like the chinook project, or your typical thing this is a chapter in a dissertation or a paper.

I think it is useful for number things in order on a three-digit system, and at the top of each make directories outputs and intermediates, like this:

```
dir.create(file.path(c("outputs", "intermediates"), "203"), recursive = TRUE, showWarnings
```

I had previously used two variables output_dir and interm_dir to specify these in each notebook, but now I think it would be better to just hardwire those, for a few reasons:

- Sometimes you are working on two notebooks at once in the same environment and you don't want to get confused about where things should get written
- You can't use those variables in shell blocks of code, where you will just have to write the paths out anyway.
- Hard-wiring the paths forces you to think about the fact that once you establish the name for something, you should not change it, ever.
- Hard-wiring the paths makes it easy to identify access to those different files. In particular you can write an R script that rips through all the lines of code in the Rmds (and R files) in your project and records all the instances of writing and reading of files from the outputs and intermediates directories. If you do this, you can make a pretty cool dependency graph so that you can visualize what you need to keep to clean things up for a final reproducible project. Note: I should write a little R package that can analyze such dependencies in a project. Unless there is already something like that. (Note

that these are not package dependencies, but, rather, internal project dependencies. Note that if one is consistent with using readr functions it would be pretty easy to find all those instances of <code>read_*</code> and <code>write_*</code> and that makes it clear why standardized syntax like that is so useful. Hey! Notice that this type of analysis would be made simple if we just focused on dependencies between different Rmds. That is probably the level we want to keep it at as well. Ideally you can make a graph of all files that are output from one Rmd and read into another. That would be a fun graph to make of the Chinook project.

- Note. You should keep 900-999 as 100 slots for Rnotebooks for the final reproducible project to go with a publication. So, you can pare down all the previous notebooks and things.
- Hey! Sometimes you are going to want to write or read files that have been auto-produced. For example, if you are cycling over chromosomes, you might have output files that start something like: outputs/302/chromo_output_. So, when generating those names, make sure that the full prefix is in there, and has a trailing underscore. Then you can still find it with a regex search, and also recognize it as a signifying a class of output files.

Version control

- 8.1 Why use version control?
- 8.2 How git works
- 8.3 git workflow patterns
- 8.4 using git with Rstudio
- 8.5 git on the command line

A fast, furious overview of the tidyverse

Basically want to highlight why it can be so useful for bioinformatic data (and also some of the limitations with really large data sets).

(But, once you have whittled bams and vcfs down to things like GWAS results and tables of theta values, they dplyr is totally up for the job.)

A really key concept here is going to be the relational data model (e.g. tidy data) and how it is so much better for handling data.

A superpowerful example of this is provided by tidytree¹ which allows one to convert from phylo objects to a tidy object: Shoot! that is so much easier to look at! This is a great example of how a single approach to manipulating data works so well for other things that have traditionally not been manipulated that way (and as a consequence have been completely opaque for a long time to most people.)

Cool. I should definitely have a chapter on tidy trees and ggtree.

What I really want to stress is that the syntax of the tidyverse is such that it makes programming a relaxing and enjoying experience.

¹ https://cran.r-project.org/web/packages/tidytree/vignettes/tidytree.html

Authoring reproducibly with Rmarkdown

10.1 Notebooks

Here is a pro-tip. First, number your notebooks and have outputs and intermediates directories associated with them. And second, always save the R object that is a ggplot in the outputs so that if you want to tweak it without re-generating all the underlying data, you can do that easily.

10.2 References

Science, as an enterprise, has proceeded with each new generation of researchers building upon the discoveries and achievements of the previous. Scientific publication honors this tradition with the stringent requirement of diligent citation of previous work. Not only that, but it is incumbent upon every researcher to identify all current work by others that is related to their own, and discuss its similarities and differences. As recently as the early 90s, literature searches involved using an annual index *printed on paper!* And, if you found a relevant paper you had to locate it in a bound volume in the library stacks and copy it page by page at a Xerox machine (or send an undergraduate intern to do that...)

Today, of course, the Internet, search services like Google Scholar, and even Twitter, have made it far easier to identify related work and to keep abreast of the latest developments in your field. But, this profusion of new literature leads to new challenges with managing all this information in a way that makes it easy for you to access, read, and cite papers while writing your own publications. There are many reference management systems available today, to help you with this task. Some of these are proprietary and paid products like EndNote. Many institutions (like Colorado State University) have licenses that provide their students a no-cost way to obtain EndNote, but the license

will not extend to updates to the program, once the student has graduated. (CHECK THIS!!!).

An alternative citation manager is Zotero. It is an open source project that has been funded not by publishing companies (like its non-open-source competitors, Mendeley and ReadCube) but by the non-profit Corporation for Digital Scholarship. As an open-source project, outside contributors have been enabled to develop workflows for integrating Zotero with reproducible research authoring modalities like Rmarkdown, including RStudio integration that lets you drop citations from Zotero directly into your Rmarkdown document where they will be cited and included in the references list in the format of just about any journal you might want to choose from. Accordingly, I will describe how to use Zotero as a citation manager while writing Rmarkdown documents.

Install Zotero and be sure to install the connector for Chrome, Firefox, or Safari

10.2.1 Zotero and Rmarkdown

Zotero has to be customized slightly to integrate with Rmarkdown and Rstudio, you must install the R package citr, and you should make some configurations:

- First, you have to get a Zotero add-on that can translate your Zotero library into a different format called BibTeX format (which is used with the TeX typesetting engine and the LaTeX document preparation system). Do this by following the directions at https://retorque.re/zotero-better-bibtex/installation/.
- 2. When you restart Zotero, you can choose all the default configurations in the BetterBibTeX start-up wizard.
- 3. Then, configure the BetterBibTex preferences by going to the Zotero preferences, choosing BetterBibTex, and then selecting "Export" button. That yields a page that gives you a place to omit certain reference fields. You life will be easier if you omit fields that are often long, or which are not needed for citation. I recommend filling that field with:

abstract,copyright,file,pmid

And, you probably should restart Zotero after doing that.

4. Install the R package citr. It is on CRAN, but it is probably best

to first try the latest development version. Install it from within R using:

```
devtools::install_github("crsh/citr")
```

For more information about this package check out https://github.com/crsh/citr.

- 5. Once that package is installed. Quit and re-open RStudio. Now, if you go to the "Addins" menu (right under the name panel at the top of the RStudio window) you will see the option to "Insert citations." Choosing that brings up a dialog box. You can choose the Zotero libraries to connect to. It might take a while to load your Zotero library if it is large. Once it is loaded though, you just start typing the name of the author or part of an article title, and boom! if that article is in your library it appears as an option. If you select it, you get a markdown citation in your text.
- 6. To avoid having to go to the "Addins" menu, you can set a key-board shortcut for "Insert citations" by choosing the "Code" section of RStudio's preferences and, under the "Editing" tab, clicking the "Modify Keyboard Shortcuts" command, searching for "Insert citations" and then selecting the keyboard shortcut area of the row and keying in which keys you would like to give you the shortcut (for example, Shift-CMD-I).

After those steps, you are set up to draw from your Zotero library or libraries to insert citations into your R markdown document.

Pretty cool, but there are some things that are sort of painful—namely the Title vs. Sentence casing. Fortunately, citr just adds things to your references.bib, it doesn't re-overwrote references.bib each time, so you can edit references.bib to put titles in sentence case. Probably want to export without braces protecting capitals. Then it should all work. See this discussion¹. Just be sure to version control references.bib and commit it often. Though, you might want to go back and edit stuff in your Zotero library.

(Barson et al., 2015)

 $^{^{1} \}texttt{https://forums.zotero.org/discussion/61715/prevent-extra-braces-in-bibtex-export}$

10.3 Bookdown

Whoa! Bookdown has figured out how to do references to sections and tables and things in a reasonable way that just isn't there for the vanilla Rmarkdown. But you can use the bookdown syntax for a non-book too. Just put something like this in the YAML:

```
output:
  bookdown::word_document2: default
  bookdown::pdf_document2:
    number_sections: yes
  bookdown::html_document2:
    df_print: paged
```

10.4 Google Docs

This ain't reproducible research, but I really like the integration with Zotero. Perhaps I need a chapter which is separate from this chapter that is about disseminating results and submitting stuff, etc.

Using python

Many people live and breathe python. Most conservation geneticists and most biologists in general are probably more familiar and comfortable with R. Nonetheless, there are some pieces of software that are available in python and it behooves us to know at least enough to crack those open and start using them.

Happily, the makers of RStudio have made it very easy to use python within the RStudio IDE. This makes the transition considerably easier.

Topics:

- conda and installing and setting up python environments. (What are they under the hood?)
- the reticulate package
- step-by-step instructions.
- An example: 2Dsfs and the moments package.

Part III

Part III: Bioinformatic Analyses

$Overview\ of\ Bioinformatic\ Analyses$

This is going to be sequence based stuff and getting to variants.

Part IV will be "bioinformaticky" analyses that you might encounter once you have your variants.

I think now that I will insert QC relevant to each step in that section.

Hey Eric! Find a place to put a section about bedtools in there. It is pretty cool what you can do with it, even on VCF files, like coverage of a VCF file in windows to get a sliding window of variant density.

Next Generation Sequencing Data Acquisition

13.1 DNA Stuff

We are primarily going to be interested in representing DNA molecules in text format as series of nucleotide bases (A, C, G, T). But, in order to do this in a consistent fashion that lets us correctly and precisely compare one sequence of DNA to another we need to know how DNA "works" and we require some conventions.

Talk about the fact that it is double-stranded, and hence there is a reverse complement.

Genome references and forward vs reverse strand stuff.

13.2 Sequencing platforms

Basically talk a little about how the technologies work.

Go into paired-end vs single end reads, etc.

Strand direction. There is a good overview on a blog at: http://www.cureffi.org/2012/12/19/forward-and-reverse-reads-in-paired-end-sequencing/, to start getting one's head around it. It ends up being super important if you are looking for inversion breakpoints, etc (Corbett-Detig et al., 2012).

13.3 Library Prep Protocols

Gotta mention here about how barcodes work.

How you prepare your libraries dictates what type of data you get.

- 13.3.1 WGS
- $13.3.2 \quad \text{RAD-Seq methods}$
- 13.3.3 Amplicon Sequencing
- 13.3.4 Capture arrays, RAPTURE, etc.

Bioinformatic file formats and associated tools

Go over these, and for each, pay special attention to compressed and indexed forms and explain why that is so important. And I think that I should probably talk about the programs that are available for manipulating each of these.

14.1 Sequences

FASTA, FASTQ

14.2 Alignments

SAM/BAM. Talk about them being sorted or not. Note that the headers between these and VCF are similar.

Have a discussion here about naming positions within the genome and talk about 0-based vs 1-based coordinates.

14.2.1 Samtools

14.3 Variants

In terms of the format/standard, is going to be well worth explaining the early part of section 5 of the standard so that people know how insertions and deletions are coded. I hadn't really digested that until just today. Basically, the position in the VCF file correponds to the first character in either the

REF or the ALT field.

When you remember that, it all falls into place.

VCF. I've mostly used vcftools until now, but I've gotta admit that the interface is awful with all the –recode BS. Also, it is viciously slow. So, let's just skip it all together and learn how to use bcftools. One nice thing about bcftools is that it works a whole lot like samtools, syntactically.

Note that for a lot of the commands you need to have an indexed vcf.gz.

14.4 Segments

BED

14.5 Conversion/Extractions between different formats

 vcflib's vcf2fasta takes a phased VCF file and a fasta file and spits out sequence.

Genome Assembly

This is going to be a pretty light coverage of how it works. Maybe I could get Rachael Bay or CH to write it.

Alignment of sequence data to a reference genome

A light treatment of how bwa works. I think I will focus solely on bwa, unless someone can convince me that there are cases where something like bowtie works better.

16.1 Preprocess?

Will have a bit about sequence pre-processing (with WGS data it already comes demultiplexed, so maybe we can hold off on this until we get to RAD data). No, we need to talk about trimming and maybe slicing. Perhaps put that in a separate chapter of "preliminaries"

16.2 Read Groups

Gotta talk about this and make it relevant to conservation. i.e. maybe one individual was sampled with blood and also with tissue and each of those were included in four different library preps, etc. Give an example that makes it clear how it works.

16.3 Merging BAM files

There is a lot of discussion on biostars about how samtools does not reconstruct the \mathtt{QRG} dictionary. But I think that this must be a problem with an

older version. The newer version works just fine. That said, Picard's Merge-SamFiles seems to be just about as fast (in fact, faster. For a comparably sized file it took 25 minutes, and gives informative output telling you where it is at). And samtools merge is at well over 30 and only about 3/4 of the way through. Ultimately it took 37 minutes. Might have been on a slower machine...

However, if you have sliced your fastqs and mapped each separately, then samtools let's you not alter duplicate read group IDs, and so you can merge those all together faithfully, as I did in the impute project. Cool.

16.4 Divide and Conquer Strategies

At the end of each of these chapters, I think I will have a special section talking about ways that things can be divided up so that you can do it quickly, or at least, within time limits on your cluster.

Variant calling with GATK

Standard stuff here.

Big focus on parallelizing.

Bioinformatics for RAD seq data with and without a reference genome

We've gotta get our hands dirty with RAD and STACKS.

For some applications (like massive salamander genomes) this is the only way forward, at the moment. Can be useful, but is also fraught with peril.

Discuss all the problems, and strategies for dealing with them.

Note that stacks2 is a lot better than things were before. And, you can use a reference genome with them.

Amanda Stahlke gave a nice practical example at ConGen. Found all the materials on box through the link that Brian Hand gave me (in my email.)

That will be worth running through closely and carefully. Note that stacks2's denovo_map.pl script now takes all the tsv output and puts it into a series of bams.

Processing amplicon sequencing data

Super high read depths can cause problems for some pipelines.

I am going to mostly focus on short amplicons that are less than the number of sequencing cycles, and how we create microhaps out of those, and the great methods we have for visualizing and curating those.

Genome Annotation

I don't intend this to be a treatise on how to actually annotate a genome. Presumably, that is a task that involves feeding a genome and a lot of mRNA transcripts into a pipeline that then makes gene models, etc. I guess I could talk a little about that process, 'cuz it would be fun to learn more about it.

However, I will be more interested in understanding what annotation data look like (i.e. in a GFF file) and how to associate it with SNP data (i.e. using snpEff).

The GFF format is a distinctly hierarchical format, but it is still tabular, it is not in XML, thank god! 'cuz it is much easier to parse in tabular format.

You can fiddle it with bedtools.

Here is an idea for a fun thing for me to do: Take a big chunk of chinook GFF (and maybe a few other species), and then figure out who the parents are of each of the rows, and then make a graph (with dot) showing all the different links (i.e. gene -> mRNA -> exon -> CDS) etc, and count up the number of occurrences of each, in order to get a sense of what sorts of hierarchies a typical GFF file contains.

Whole genome alignment strategies

Basically want to talk about situations

21.1 Mapping of scaffolds to a closely related genome

I basically want to get my head fully around how SatsumaSynteny works.

After that, we might as well talk about how to get in and modify a VCF file to reflect the new positions and such. (It seems we could even add something to the INFO field that listed its position in the old scaffold system. awk + vcftools sort seems like it might be the hot ticket.)

21.2 Obtaining Ancestral States from an Outgroup Genome

For many analyses it is helpful (or even necessary) to have a guess at the ancestral state of each DNA base in a sequence. These ancestral states are often guessed to be the state of a closely related (but outgroup) species. The idea there is that it is rare for the same nucleotide to experience a substitution (or mutation) in each species, so the base carried by the outgroup is assumed to be the ancestral sequence.

So, that is pretty straightforward conceptually, but there is plenty of hardship along the way to do this. There are two main problems:

1. Aligning the outgroup genome (as a query) to the target genome. This typically produces a multiple alignment format (MAF) file. So, we have to understand that file format. (read about it here¹, on the

¹http://genome.ucsc.edu/FAQ/FAQformat#format5

UCSC genome browser site.) A decent program to do this alignment exercise appears to be LASTZ²

2. Then, you might have to convert the MAF file to a fasta file to feed into something like ANGSD. It seems that Dent Earl has some tools that might to do this hhttps://github.com/dentearl/mafTools³. Also, the ANGSD github site has a maf2fasta⁴ program, though no documentation to speak of. Or you might just go ahead and write an awk script to do it. Galaxy has a website that will do it: http://mendel.gene.cwru.edu:8080/tool_runner?tool_id=MAF_To_Fasta1, and there is an alignment too called mugsy that has a perl script associated with it that will do it: ftp://188.44.46.157/mugsy_x86-64-v1r2.3/maf2fasta.pl Note that the fasta file for ancestral sequence used by ANGSD just seems to have Ns in the places that don't have alignments.

It will be good to introduce people to those "dotplots" that show alignments.

Definitely some discussion of seeding and gap extensions, etc. The LASTZ web page has a really nice explanation of these things.

The main take home from my explorations here is that there is no way to just toss two genomes into the blender with default setting and expect that you are going to get something reasonable out of that. There is a lot of experimentation, it seems to me, and you really need to know what all the options are (this might be true of just about everything in NGS analysis, but in many cases people just use the defaults....)

21.2.1 Using LASTZ to align coho to the chinook genome

First, compile it:

```
# in: /Users/eriq/Documents/others_code/lastz-1.04.00
make
make install
# then I linked those (in bin) to myaliases
```

Refreshingly, this has almost no dependencies, and the compilation is super easy.

 $^{^2} http://www.bx.psu.edu/miller_lab/dist/README.lastz-1.02.00/README.lastz-1.02.00a.html$

³https://github.com/dentearl/mafTools

⁴https://github.com/ANGSD/maf2fasta

Now, let's find the coho chromomsome that corresponds to omy28 on NCBI. We can get this with curl:

```
# in: /tmp
curl ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/002/021/735/GCA_002021735.1_0kis_V1/GCA_00
gunzip coho-28.fna.gz
```

Then, let's also pull that chromosome out of the chinook genome we have:

```
# in / tmp samtools faidx \sim Documents/UnsyncedData/Otsh_v1.0/Otsh_v1.0_genomic.fna NC_037124.1 > chi
```

Cool, now we should be able to run that:

```
time lastz chinook-28.fna coho-28.fna --notransition --step=20 --nogapped --ambiguous=iupa
real 0m14.449s
user 0m14.198s
sys 0m0.193s
```

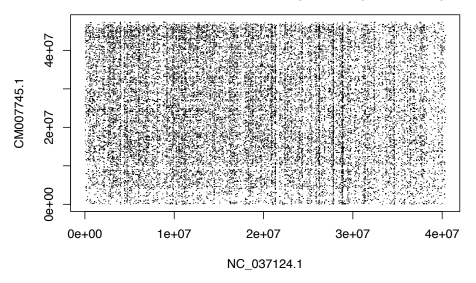
OK, that is ridiculously fast. How about we make a file that we can plot in R?

```
time lastz chinook-28.fna coho-28.fna --notransition --step=20 --nogapped --ambiguous=iupa
```

I copied that to inputs so we can plot it:

```
dots <- readr::read_tsv("inputs/chin28_vs_coho28.rdp.gz")

## Parsed with column specification:
## cols(
## NC_037124.1 = col_double(),
## CM007745.1 = col_double()
## )</pre>
plot(dots,type="l")
```



OK, clearly what we have there is just a bunch of repetitive bits. I think we must not have the same chromosomes in the two species.

So, let's put LG28 in coho against the whole chinook genome. Note the use of the bracketed "multiple" in there to let it know that there are multiple sequences in there that should get catenated together.

```
time lastz ~/Documents/UnsyncedData/Otsh_v1.0/Otsh_v1.0_genomic.fna[multiple] coho-28.fna
FAILURE: in load_fasta_sequence for /Users/eriq/Documents/UnsyncedData/Otsh_v1.0/Otsh_v1.0
```

No love there. But that chinook genome has a lot of short scaffolds in there too, I think.

Maybe we could just try LG1. Nope. How about we toss every coho LG against LG1 from chinook...

```
# let's get the first 10 linkage groups from coho:
for i in {1..10}; do curl ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/002/021/735/GCA_00202
# now lets try aligning those to the chinook
for i in {1..10}; do time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --nog
```

Nothing looked good until I got to coho LG10:

```
dots <- readr::read_tsv("inputs/chinook1_vs_coho10.rdp.gz")
plot(dots,type="1")</pre>
```

There is clearly a big section that aligns there. But, we clearly are going to need to clean up all the repetive crap, etc on these alignments.

21.2.2 Try on the chinook chromosomes

So, it crapped out on the full Chinook fasta. Note that I could modify the code (or compile it with a -D): check this out in sequences.h:

```
// Sequence lengths are normally assumed to be small enough to fit into a // 31-bit integer. This gives a maximum length of about 2.1 billion bp, which // is half the length of a (hypothetical) monoploid human genome. The // programmer can override this at compile time by defining max_sequence_index // as 32 or 63.
```

But, for now, I think I will just go for the assembled chromosomes only:

```
# just get the well assembled chromosomes (about 1.7 Gb)
# in /tmp
samtools faidx ~/Documents/UnsyncedData/Otsh_v1.0/Otsh_v1.0_genomic.fna $(awk '/^NC/ {print
# then try tossing coho 1 against that:
time lastz chinook_nc_chroms.fna[multiple] coho-1.fna --notransition --step=20 --nogapped
# that took about 7 minutes
real 6m33.411s
user 6m28.391s
sys 0m4.121s
```

Here is what the figure looks like. It is clear that the bulk of Chromosome 1 in coho aligns for long distances to two different chromosomes in Chinoook (likely paralogs?). This is complex!

21.2.3 Explore the other parameters more

I am going to use the chinook chromosome 1 and coho 10 to explore things that will clean up the results a little bit.

```
# in /tmp
i=10; time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --nogapped --ambigu
```

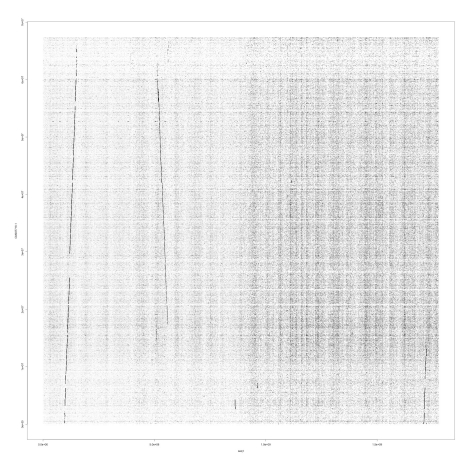


FIGURE 21.1: Coho Chromo 1 on catenated chinook chromos

```
real 0m29.055s
user 0m28.497s
sys 0m0.413s
```

That is quick enough to explore a few things. Note that we are already doing gap-free extension, because "By default seeds are extended to HSPs using x-drop extension, with entropy adjustment." However, by default, chaining is not done, and that is the key step in which a path is found through the high-scoring pairs (HSPs). That doesn't take much (any) extra time and totally cleans things up.

```
i=10; time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --nogapped --ambiguo
real 0m28.704s
```

So, that is greatly improved:

```
dots <- readr::read_tsv("inputs/chain-chinook1_vs_coho10.rdp.gz")
plot(dots,type="1")</pre>
```

So, it would be worth seeing if chaining also cleans up the multi-chrom alignment:

```
time lastz chinook_nc_chroms.fna[multiple] coho-1.fna --notransition --step=20 --nogapped real 6m42.835s user 6m35.042s sys 0m6.207s
```

```
dots <- readr::read_tsv("inputs/chin_nc_vs_coho1-chain.rdp.gz")
plot(dots,type="l")</pre>
```

OK, that shows that it will find crappy chains if given the chance. But if you zoom in on that stuff you see that some of the spots are pretty short, and some are super robust. So, we will want some further filtering to make this work. So, we need to check out the "back-end filtering." that is possible. Back-end filtering does not happen by default.

Let's say that we want 70 Kb alignment blocks. That is .001 of the total input sequence.

```
time lastz chinook_nc_chroms.fna[multiple] coho-1.fna --notransition --step=20 --nogapped
```

That took 6.5 minutes again. But, it also produced no output whatsoever. We probably want to filter on identity first anyway. Because that takes so long, maybe we could do it with our single chromosome first.

```
i=10; time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --nogapped --ambiguo
```

```
dots <- readr::read_tsv("/tmp/chinook1_vs_coho10-ident95.rdp")
plot(dots,type="l")</pre>
```

That keeps things very clean, but the alignment blocks are all pretty short (like 50 to 300 bp long). So perhaps we need to do gapped extension here to make these things better. This turns out to take a good deal longer.

```
i=10; time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --gapped --ambiguous
real 3m15.575s
user 3m14.048s
sys 0m0.936s
```

```
dots <- readr::read_tsv("/tmp/chinook1_vs_coho10-ident95-gapped.rdp")
plot(dots,type="1")</pre>
```

That is pretty clean and slick.

Now, this has got me to thinking that maybe I can do this on a chromosome by chromosome basis.

Check what 97% identity looks like:

```
i=10; \ time \ lastz \ chinook-1.fna \ coho-\$\{i\}.fna \ --notransition \ --step=20 \ --gapped \ --ambiguous \ --a
```

```
dots <- readr::read_tsv("/tmp/chinook1_vs_coho10-ident97-gapped.rdp")
plot(dots,type="1")</pre>
```

That looks to have a few more holes in it.

Final test. Let's see what happens when we chain it on a chromosome that doesn't have any homology:

Hey! That is cool. When there are no HSPs to chain, this doesn't take very long i=1; time lastz chinook-1.fna coho-\${i}.fna --notransition --step=20 --gapped --ambiguous=

```
dots <- readr::read_tsv("/tmp/chinook1_vs_coho1-gapped.rdp")
plot(dots,type="1")</pre>
```

OK, it finds something nice and crappy there.

What about if we require 95% identity?

```
dots <- readr::read_tsv("/tmp/chinook1_vs_coho1-gapped-ident95.rdp")
plot(dots,type="1")</pre>
```

That leaves us with very little.

Let's also try interpolation at the end to see how that does. Note that here we also produce the rdotplot at the same time as the maf.

```
i=10; time lastz chinook-1.fna coho-${i}.fna --notransition --step=20 --gapped --ambiguous
real 4m25.625s
user 4m22.957s
sys 0m1.478s
```

That took an extra minute, but was not so bad.

```
dots <- read_tsv("/tmp/chinook1_vs_coho10-ident95-gapped-inner1000.rdp")
plot(dots,type="l")</pre>
```

21.2.3.1 Repeat Masking the Coho genome

Turns out that NCBI site has the repeat masker output in GCF_002021735.1_Okis_V1_rm.out.gz. I save that to a shorter name. Now I will make a bed file of the repeat regions. Then I use bedtools maskfasta to softmask that fasta file.

```
# in: /Users/eriq/Documents/UnsyncedData/Okis_v1
gzcat Okis_V1_rm.out.gz | awk 'NR>3 {printf("%s\t%s\t", $5, $6, $7)}' > repeat-regions
bedtools maskfasta -fi Okis_V1.fna -bed repeat-regions.bed -fo Okis_V1-soft-masked.fna -
```

That works great. But it turns out that the coho genome is already softmasked.

But, it is good to now that I can use a repeat mask output file to toss repeat sites if I want to for ANGSD analyses, etc.

21.2.3.2 multiz maf2fasta

So, it looks like you can use single_cov2 from multiz to retain only a single alignment block covering each area. Then you can maf2fasta that and send it off in fasta format. Line2 holds the reference (target) sequence, but it has dashes added where the query has stuff not appearing in the target. So, what you have to do is run through that sequence and drop all the positions in the query that correspond to dashes in the target. That will get us what we want.

But maybe I can just use megablast like Christensen and company. They have some of their scripts, but it is not clear to me that it will be easy to get that back to a fasta for later analysis in ANGSD.

Not only that, but then there is the whole paralog issue. I am exploring that a little bit right now. It looks like when you crank the identity requirement up, the paralogs get pretty spotty so they can be easily recognized. For example setting the identity at 99.5 makes it clear which is the paralog:

time lastz chinook_nc_chroms.fna[multiple] coho-1.fna --notransition --step=20 --nogapped # takes about 6 minutes

And the figure is here.

So, I think this is going to be a decent workflow:

- 1. run lastz on each coho linkage group against each chinook chromosome separately. Do this at identity=92 and identity=95 and identity=99.9. For each run, produce a .maf file and at the same time a .rdotplot output.
- Combine all the .rdotplot files together into something that can be faceted over chromosomes and make a massive facet grid showing the results for all chromosomes. Do this at different identity levels so that the paralogs can be distinguished.
- 3. Visually look up each columns of those plots and determine which coho chromosomes carry homologous material for each chinook chromosome. For each such chromosome pair, run single_cov2 on them (maybe on the ident=92 version).
- 4. Then merge those MAFs. Probably can just cat them together, but there might be some sorting that needs to be done on them.
- 5. Run maf2fasta on those merged mafs to get a fasta for each chinook chromosome.

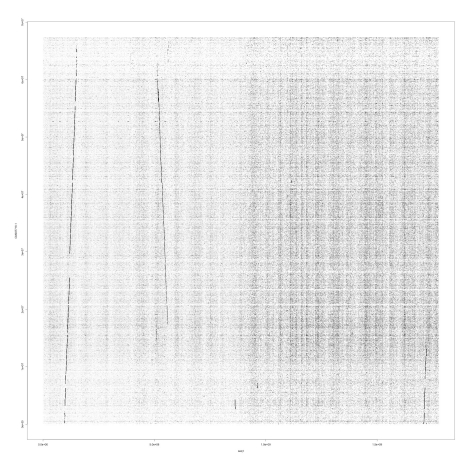


FIGURE 21.2: Coho Chromo 1 on catenated chinook chromos. Ident=99.5

6. Write a C-program that uses uthash to efficiently read the fasta for each chinook chromosome and then write out a version in which the positions that are dashes in the chinook reference are removed from both the chinook reference and the aligned coho sequence.

_Actually, one can just pump each sequence out to a separate file in which each site occupies one line. Then paste those and do the comparison...

```
2018-10-18 11:15 /tmp/--% time (awk 'NR==2' splud | fold -w1 > spp1.text) real 0m23.244s user 0m22.537s
```

```
# then you can use awk easily like this:
paste spp1.text spp2.text | awk 'BEGIN {SUBSEP = " "} {n[$1,$2]++} END {for(i in n) print
```

- 7. The coho sequence thus obtained will have dashes anywhere there isn't coho aligned to the chinook. So, first, for each chromosome I can count the number of dashes, which will tell me the fraction of sites on the chinook genome that were aligned (sort of—there is an issue with N's in the coho genome.) Then those dashes can be converted to N's.
- 8. It would be good to count the number of sites that are not N's in chinook that are also not Ns in coho, to know how much of it we have aligned.

Note, the last thing that really remains here is making sure that I can run two or more different query sequences against one chinook genome and then process that out correctly into a fasta.

Note that Figure 1 in christensen actually gives me a lot to go on in terms of which chromosomes in coho to map against which ones in chinook.

Part IV

Part IV: Analysis of Big Variant Data

Bioinformatic analysis on variant data

Standard analyses like computing Fst and linkage disequilibrium, etc., from data, typically in a VCF file.

Basically, we want to get comfortable with plink 2.0, bedtools, vcftools, etc.

The key in all of this is to motivate every single thing we do here in terms of an application in conservation genomics. That is going to be key.

This Part IV will be about standard bioinformatic tools for doing things with big variant data.

- Filtering
- Imputation
- LD, HWD, FST
- Etc.

I will probably have a chapter on unix tools.

Maybe another on R/Bioconductor tools.

Gotta have a chapter about "Look at your data!" and Whoa! and diagnostics using radiator.

Part V

Part V: Population Genomics

Topics in pop gen

This is just a bunch of ideas. But basically, I want to have some topics here that everyone should know about. Slanted toward things that are relevant for inference or simulation.

23.1 Coalescent

Gotta have a lecture on the coalescent. It would be nice to try to motivate all the topics from this backward in time perspective.

Get far enough to discuss π and the expected site frequency spectrum.

23.2 Measures of genetic diversity and such

It would really be good for me to write a chapter / give a few lectures on different measures like d_{xy} , $F_{\rm ST}$.

From Ash's paper: However, population genomic analyses (outlined below) use FST only, as dxy was highly correlated to nucleotide diversity () (for early stage diverging populations the correlation between dxy and π is > 0.91, Pearson correlation). As such variation in dxy across the genome reflects variation in diversity, not differentiation (Riesch et al., 2017).

Tajima's D and such. The influence of selection on such measures.

23.3 Demographic inference with $\partial a \partial i$ and moments

23.4 Balls in Boxes

Would be worthwhile to have a review of all these sorts of variants of population assignment, structure, admixture, etc.

Population structure and PCAs.

finestructure and fineRADstructure.

Might want to insert Bradburd et al. (2018).

Might also want to discuss Pickrell and Pritchard (2012).

Also: Pritchard et al. (2000)

What if we go and try to put the same one in? Like Pritch 2000 again: (Pritchard et al., 2000)

23.5 Some landscape genetics

After talking with Amanda about her dissertation I realized it would be good to talk about some landscape genetics stuff. For sure I want to talk about EEMS and maybe CircuitScape, just so I know well what is going on with the latter.

23.6 Relationship Inference

Maybe do a lecture on this...

23.7 Tests for Selection

A look at a selection of the methods that are out there. FST outliers, *Bayescan*, *Lositan*, *PCAdapt*, and friends. It would be good to get a nice succinct explanation/understanding of all of these.

23.8 Multivariate Associations, GEA, etc.

It really is time for me to wrap my head around this stuff.

23.9 Estimating heritability in the wild

Another from Amanda. It would be good to do some light Quant Genet so that we all understand how we might be able to use NGS data to estimate heritability in wild populations.

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