

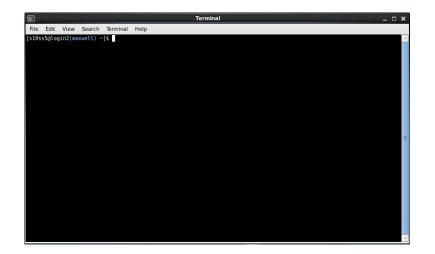
# Cloud Computing and Unix: An Introduction

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## What We're Going To Do

- Why Unix?
- Cloud Computing
- Connecting to AWS
- Introduction to Unix Commands



## Etiquette

- PowerPoint interspersed with Challenges
- Ask me questions
- Ask demonstrators
- Work together
- Cheat!



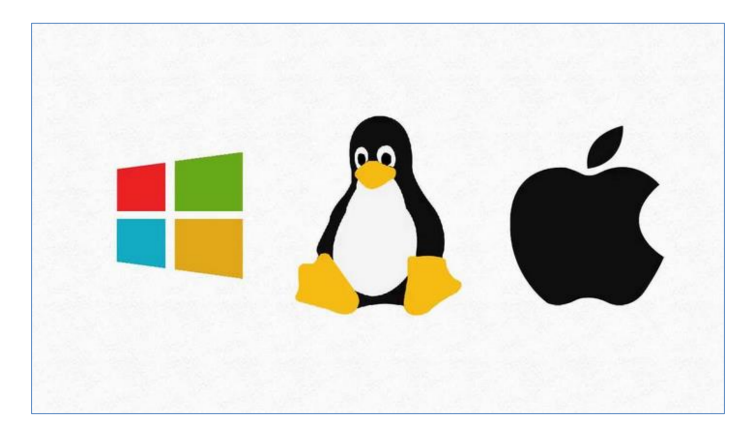
#### Linix/Linux Command Deference



Unix/Linux Command Ref	erence FOSSwire.com							
File Commands	System Info							
ls - directory listing	date - show the current date and time							
1s -al - formatted listing with hidden files	cal - show this month's calendar							
cd dir - change directory to dir	uptime - show current uptime							
cd - change to home	w - display who is online							
pwd - show current directory	whoami - who you are logged in as							
mkdir dir – create a directory dir	finger user - display information about user							
rm file - delete file	uname -a - show kernel information							
rm -r dir - delete directory dir	cat /proc/cpuinfo - cpu information							
rm -f file - force remove file	cat /proc/meminfo - memory information							
<pre>rm -rf dir - force remove directory dir * cp file1 file2 - copy file1 to file2</pre>	man command - show the manual for command df - show disk usage							
cp -r dir1 dir2 - copy dir1 to dir2; create dir2 if it								
doesn't exist	free - show memory and swap usage							
mv file1 file2 - rename or move file1 to file2	where is app - show possible locations of app							
if file2 is an existing directory, moves file1 into	which app - show which app will be run by default							
directory file2								
In -s file link - create symbolic link link to file	Compression							
touch file - create or update file	tar cf file.tar files - create a tar named							
<pre>cat &gt; file - places standard input into file</pre>	file.tar containing files							
more file - output the contents of file	tar xf file.tar - extract the files from file.tar							
head file - output the first 10 lines of file	tar czf file.tar.gz files - create a tar with Gzip compression							
tail file - output the last 10 lines of file	tar xzf file.tar.gz - extract a tar using Gzip							
<b>tail -f</b> <i>file</i> - output the contents of <i>file</i> as it grows, starting with the last 10 lines	tar cjf file.tar.bz2 - create a tar with Bzip2							
	compression							
Process Management	tar xjf file.tar.bz2 - extract a tar using Bzip2							
ps – display your currently active processes	gzip file - compresses file and renames it to							
top - display all running processes	file.gz							
kill pid - kill process id pid	gzip -d file.gz - decompresses file.gz back to							
killall proc - kill all processes named proc * bg - lists stopped or background jobs; resume a	file							
stopped job in the background	Network							
fg - brings the most recent job to foreground	ping host - ping host and output results							
<b>fg</b> <i>n</i> - brings job <i>n</i> to the foreground	whois domain - get whois information for domain							
File Permissions	dig domain - get DNS information for domain							
chmod octal file - change the permissions of file	dig -x host - reverse lookup host							
to <i>octal</i> , which can be found separately for user,	wget file - download file							
group, and world by adding:	wget -c file - continue a stopped download							
• 4 - read (r)	Installation							
• 2 - write (w)	Install from source:							
<ul> <li>1 - execute (x)</li> </ul>	./configure							
Examples:	make							
chmod 777 - read, write, execute for all	make install							
chmod 755 - rwx for owner, rx for group and world	dpkg -i pkg.deb - install a package (Debian)							
For more options, see man chmod.	<pre>rpm -Uvh pkg.rpm - install a package (RPM)</pre>							
SSH	Shortcuts							
<pre>ssh user@host - connect to host as user</pre>	Ctrl+C - halts the current command							
<pre>ssh -p port user@host - connect to host on port</pre>	<b>Ctrl+Z</b> - stops the current command, resume with							
port as user	fg in the foreground or bg in the background							
ssh-copy-id user@host - add your key to host for	Ctrl+D - log out of current session, similar to exit							
user to enable a keyed or passwordless login	Ctrl+W - erases one word in the current line							
Searching	Ctrl+U - erases the whole line							
grep pattern files - search for pattern in files	Ctrl+R - type to bring up a recent command							
grep -r pattern dir - search recursively for	!! - repeats the last command							
pattern in dir	exit - log out of current session							
<b>command</b>   grep pattern - search for pattern in the output of command								
locate file - find all instances of file	* use with extreme caution.							
inter inter an interaction of the								

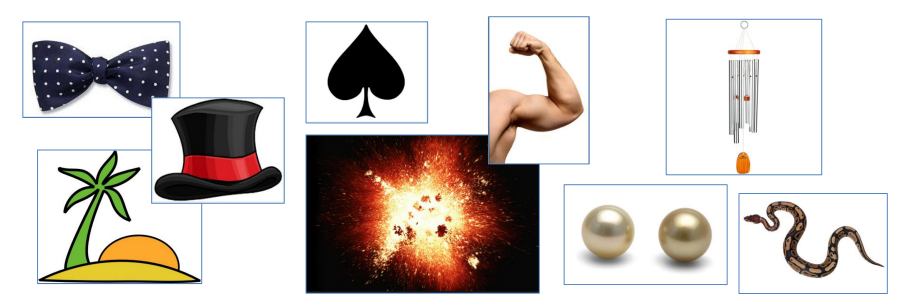
## What is Unix?

• Operating System



# Why Unix?

- Bioinformatics software designed to run on Unix platforms.
- Large amounts of data.
- Much faster than your Windows PC.



## How Can We Use Unix?

- Linux computers or servers.
- Compute clusters.
- The cloud.
  - What we're going to use this week

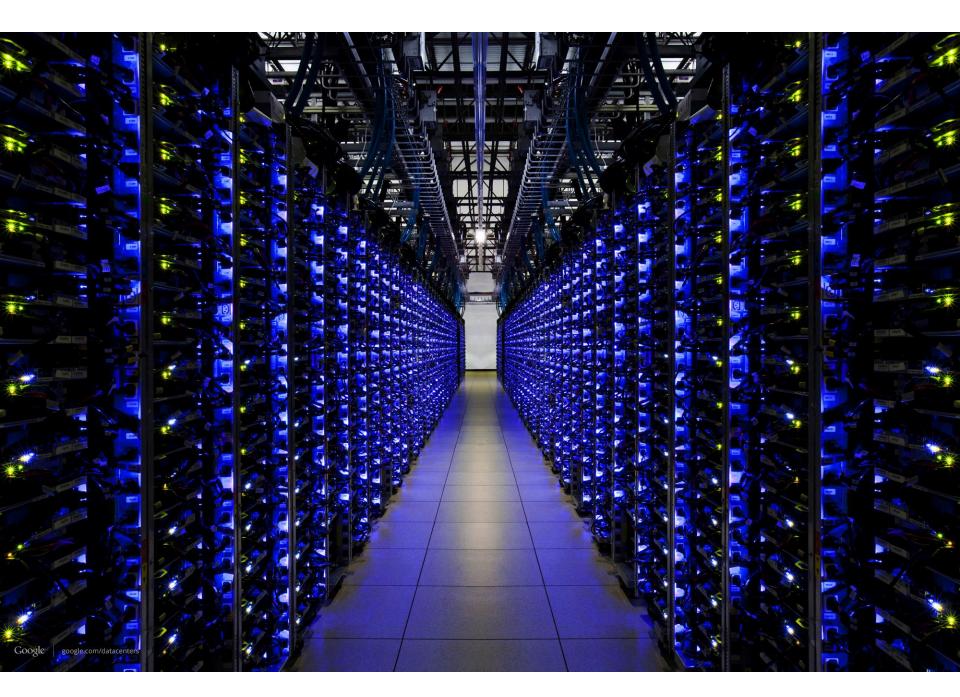


## So What is Cloud Computing?





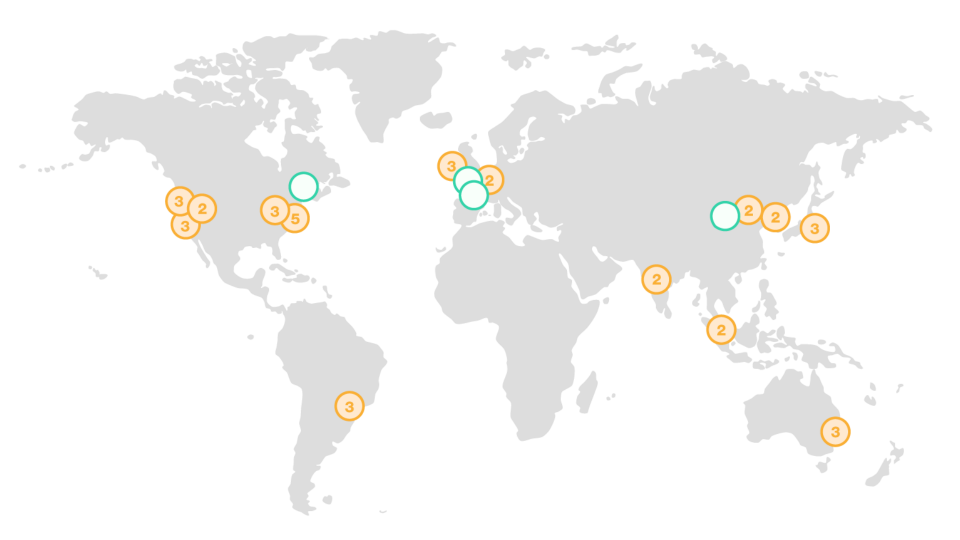




## **Cloud Computing Solutions**



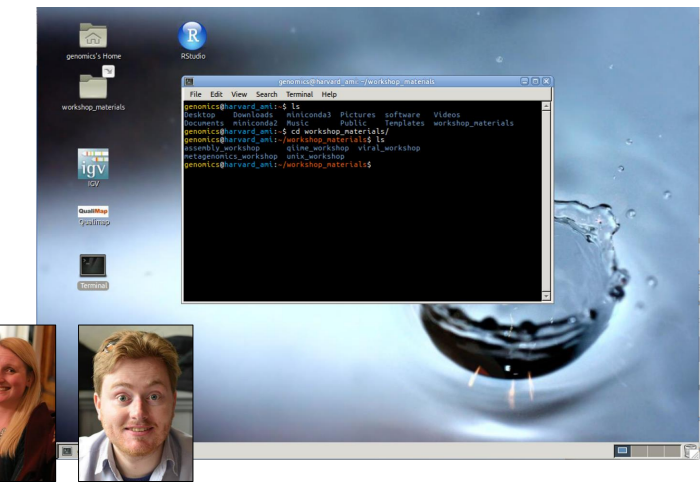
Google Compute Engine



#### AWS "Availability Zones" and Data Centres

### How it Works

#### AMI ("Amazon Machine Image") Base computer with all data and software



### How it Works











Own copy of the AMI = Instance (Virtual Machine or VM)

## Terminology

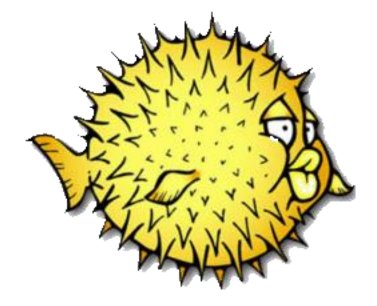
- Creating an instance buying a brand new computer with software already installed.
- Starting an instance *turning that computer on.*
- Stopping an instance *turning that computer off.*
- Terminating an instance setting that computer on fire and throwing it out of the window.

## The Rules

- Only create one instance each.
- Stop your instance at the end of each day (unless you have software running).
- Name your instance (with YOUR name! No Bruce Waynes please)
- Only start or stop your own instance.
- Only terminate your own instance.

### **Connecting to Your Instance**



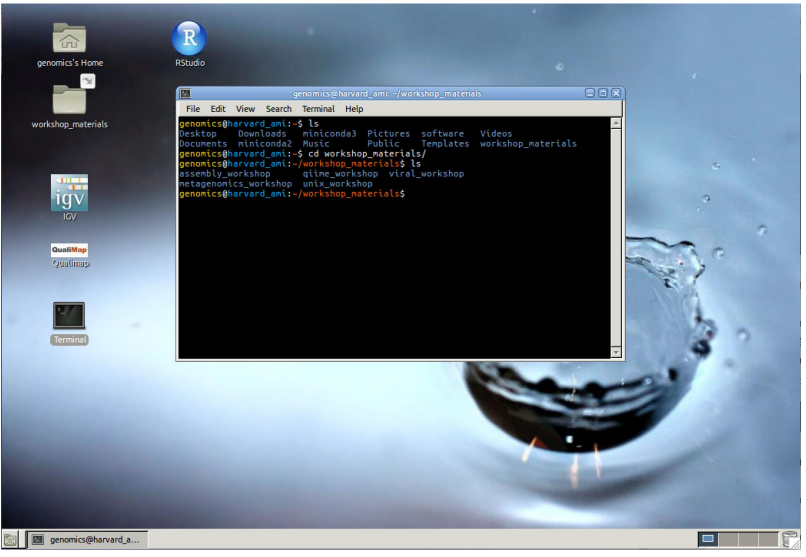


Remote Desktop Software e.g. X2Go Secure Shell – "SSH" e.g. SSH or PuTTY

## Now What?!

• You're each going to create, start and connect to your own instance.

#### \*\*\*INSERT LIVE DEMO\*\*\*

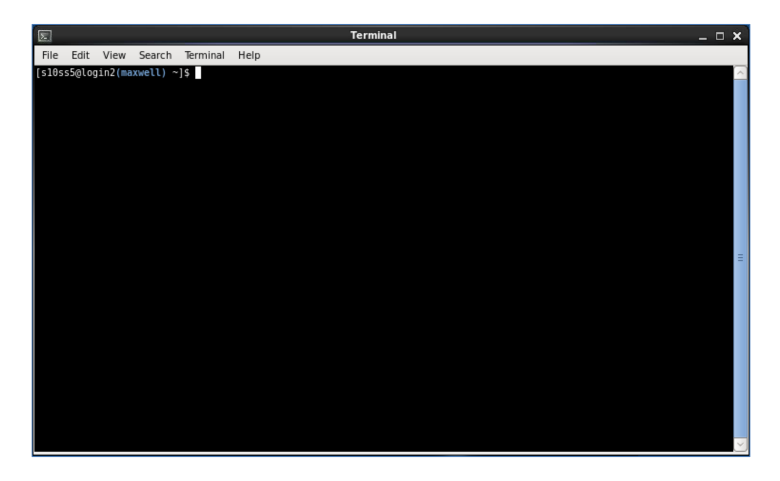


You're now connected to your instance and you're ready to learn some Unix!

#### Any Questions So Far?



## The Terminal Window



The Command Line, The Shell, The Prompt

Where you see this "\$" followed by text, I want you to type the text on your command line

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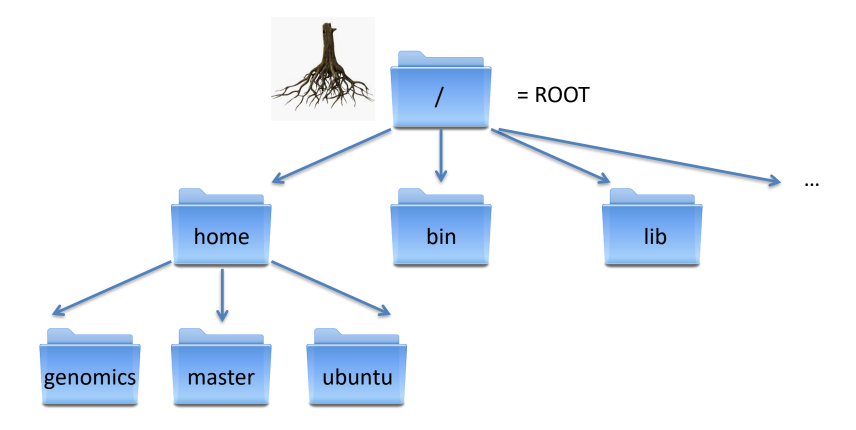
### Location is Important

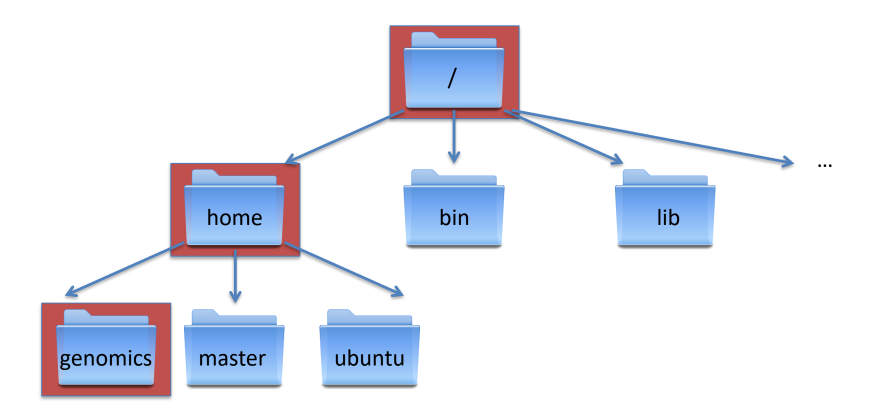
First Task – Where am I?



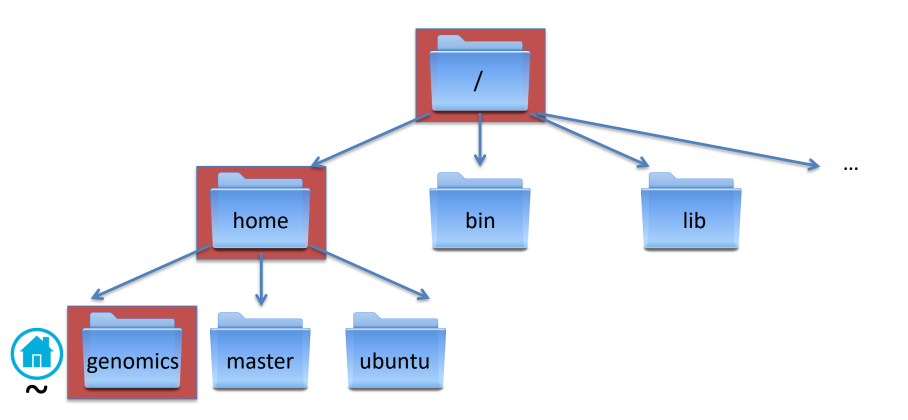
genomics@harvard\_ami:~\$ pwd
/home/genomics
genomics@harvard\_ami:~\$

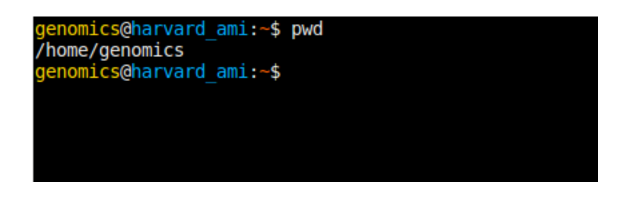
This is your "present working directory"





genomics@harvard\_ami:~\$ pwd
/home/genomics
genomics@harvard\_ami:~\$





This location is also known as your Home Directory

Tilde is shorthand for Home:

#### Now let's create some directories and files

Make a directory

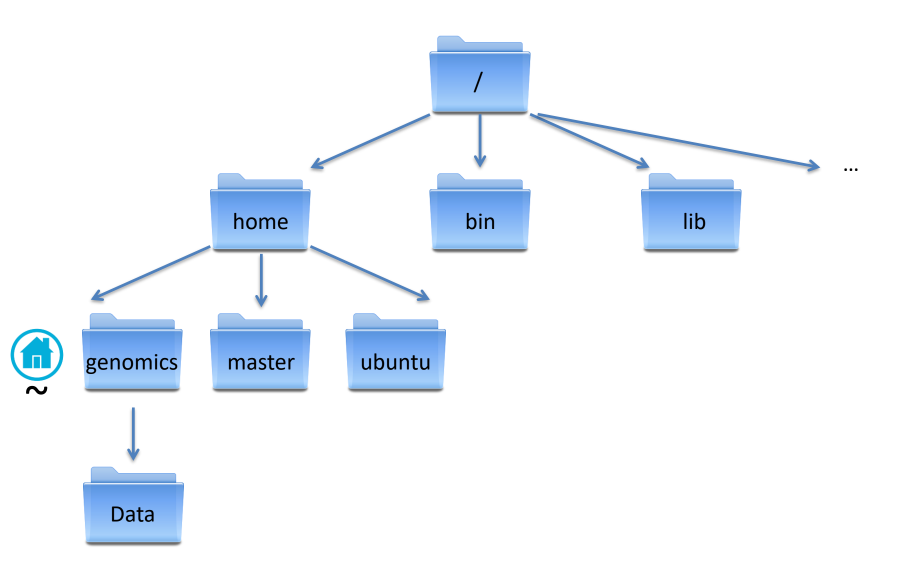


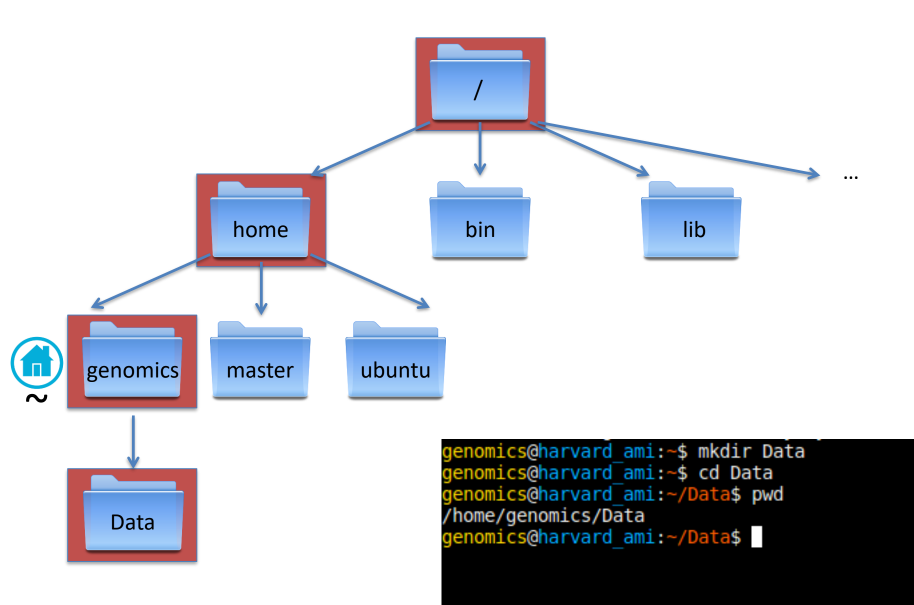
Change into this directory



Now what is your present working directory?

NOTE! Directory names (and file names for the matter) can not contain spaces. Underscores are often used instead if you want to separate words.





#### Now let's create some directories and files

Make an empty file



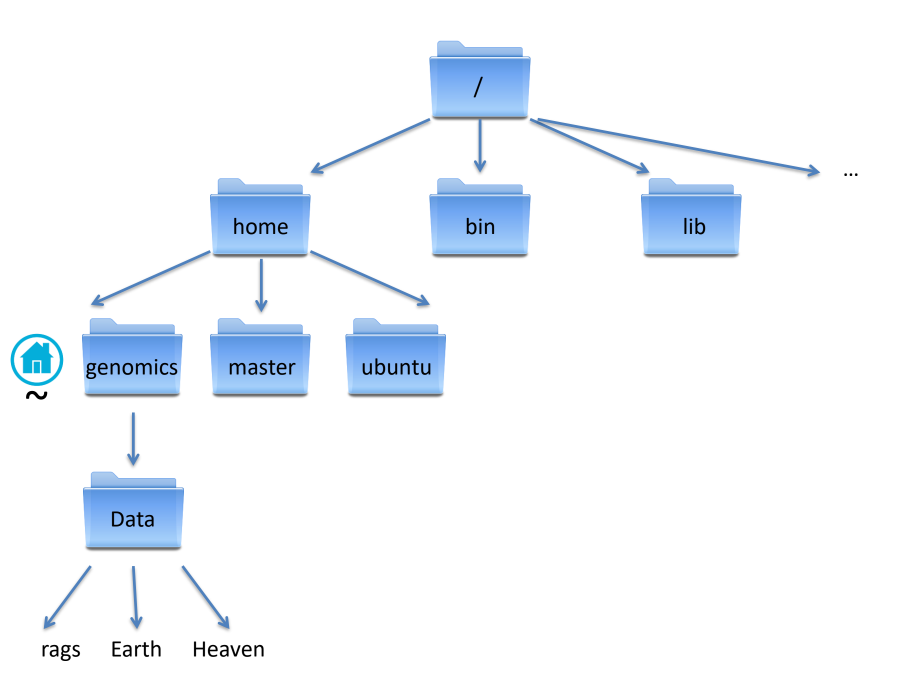
And another two

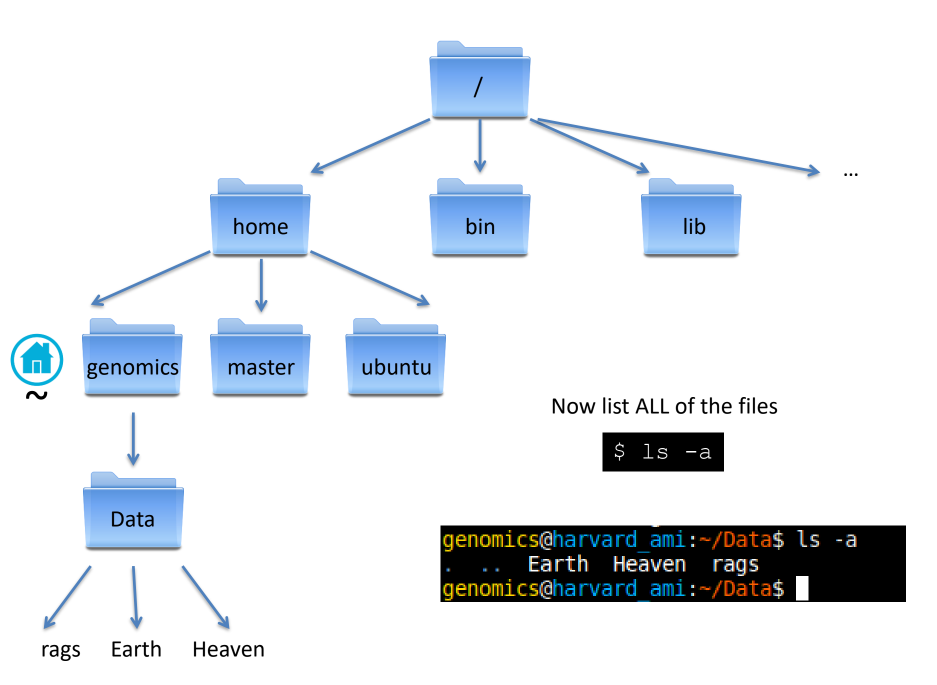
\$ touch Earth Heaven

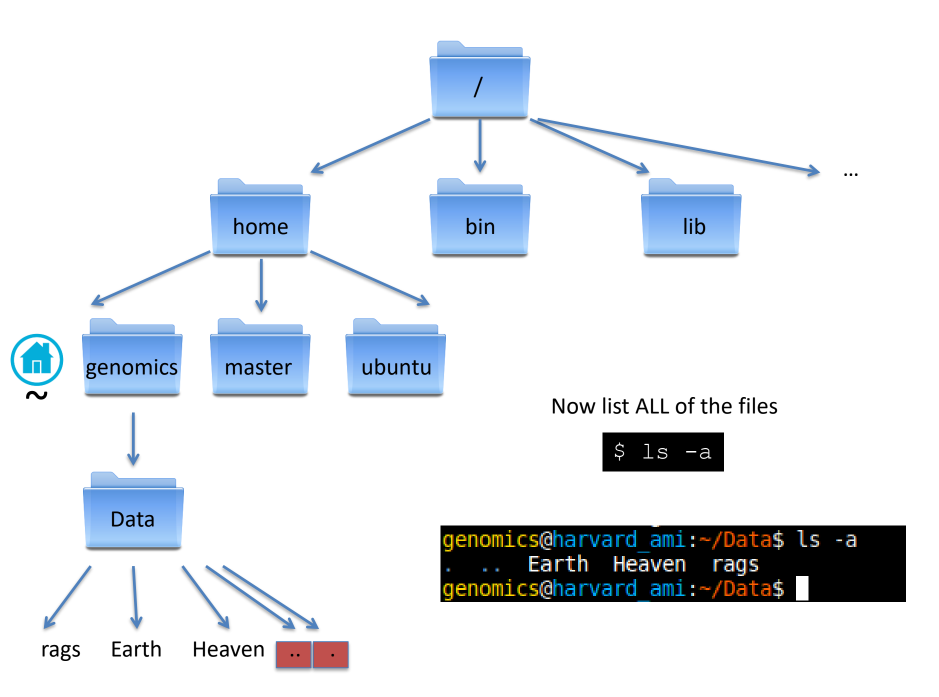
Now let's list the contents of the current directory (Data)

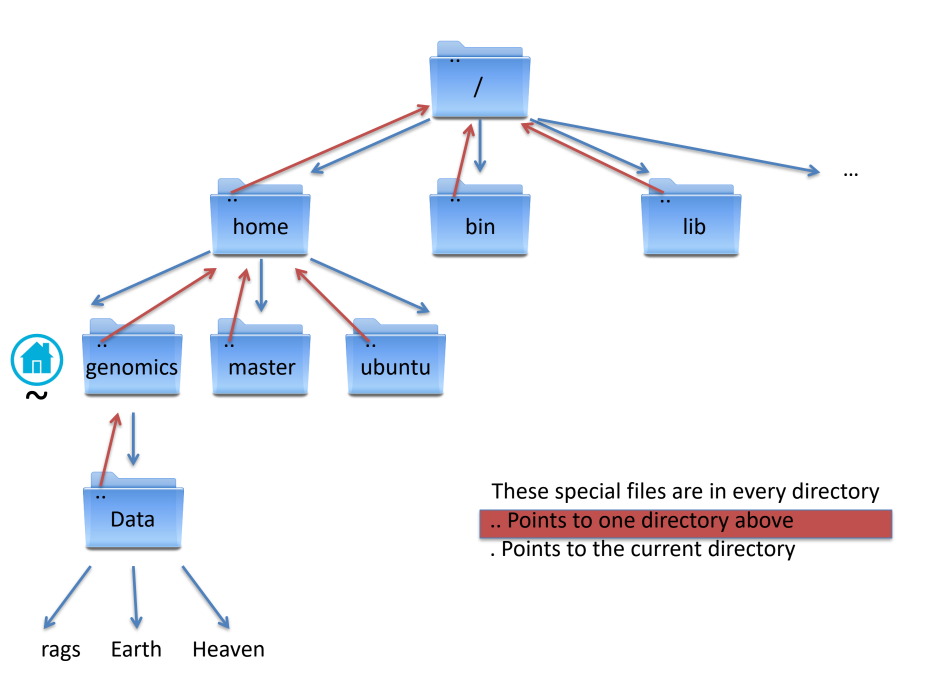


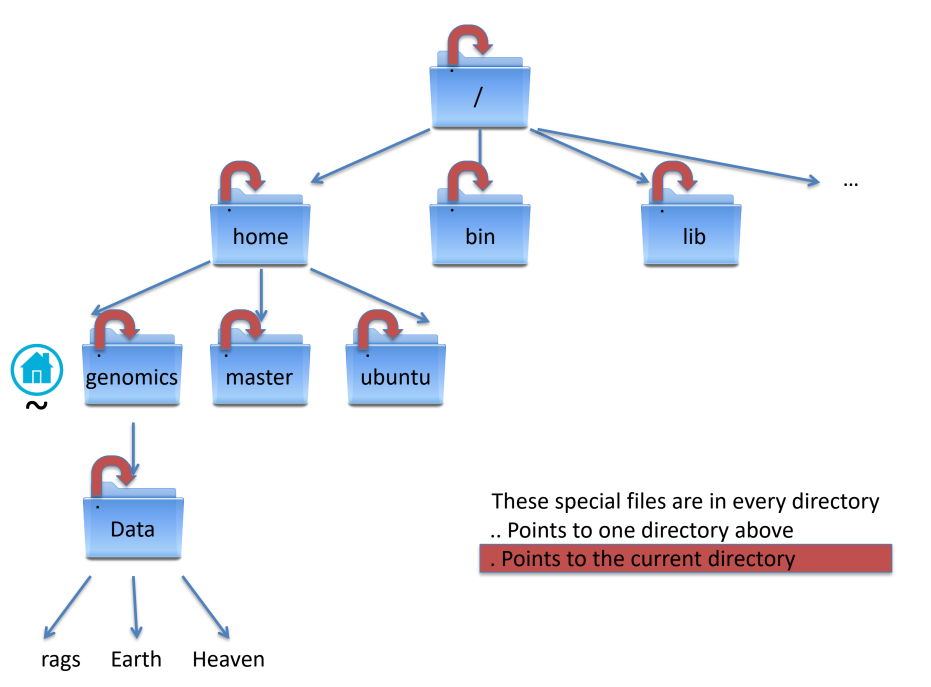
genomics@harvard\_ami:~/Data\$ touch rags
genomics@harvard\_ami:~/Data\$ touch Earth Heaven
genomics@harvard\_ami:~/Data\$ ls
Earth Heaven rags
genomics@harvard\_ami:~/Data\$





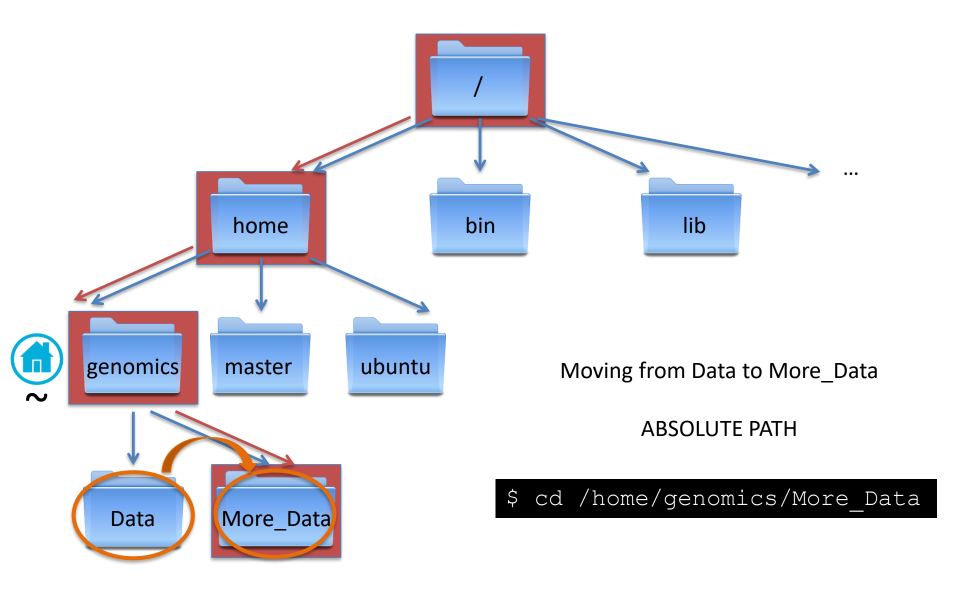




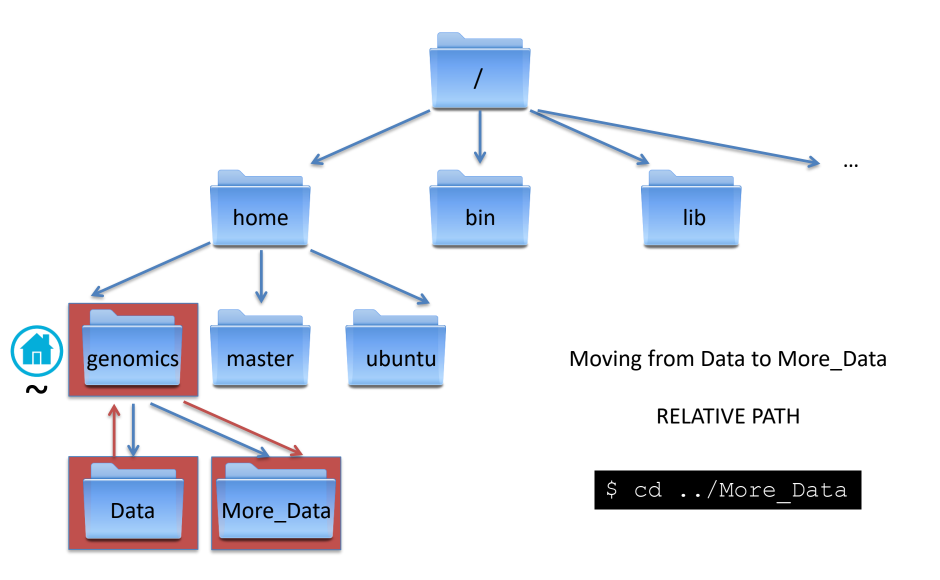


# ABSOLUTE AND RELATIVE PATHS: GETTING FROM ONE PLACE TO ANOTHER

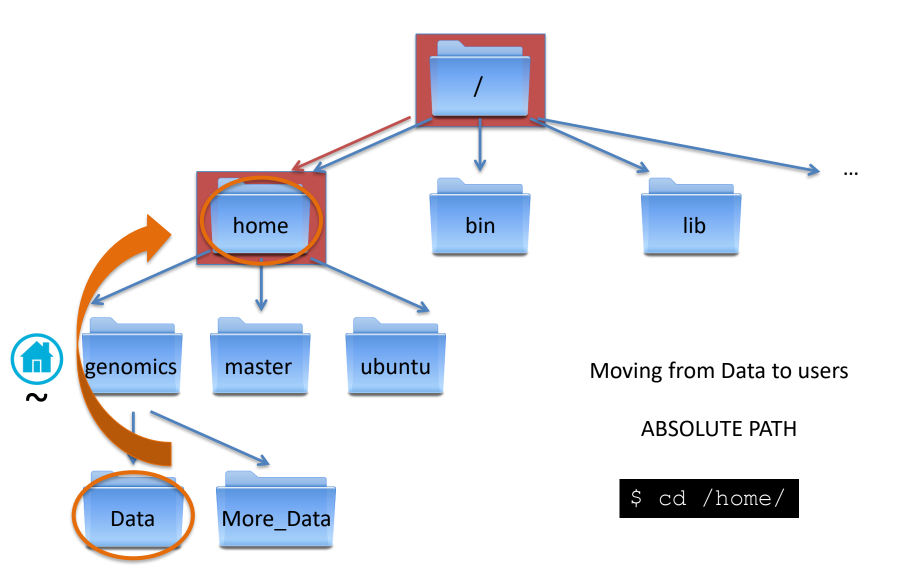
## **Absolute and Relative Paths**



# Absolute and Relative Paths



## Absolute and Relative Paths



### **Absolute and Relative Paths** • • • home bin lib genomics Moving from Data to users master ubuntu **RELATIVE PATH** \$ cd ../. Data More\_Data

# Let's put this to practice

Where am I right now? (Should be the Data directory)



Change to the directory above



Let's list the contents of the Data directory



# Let's put this to practice

Where am I right now? (Should be the Data directory)



Change to the directory above



Let's list the contents of the Data directory



#### CHALLENGE 1!

- 1. Move into the Data directory and list the contents of your home directory
- 2. In Data, make a new directory and move into this location
- 3. From this new directory, move into your home directory IN ONE COMMAND and check your location

### If You're Typing, You're Doing Something Wrong!

Tab complete is a nice trick to save you typing paths

For this examples we are going to list everything in directory /var/run

Start by typing:



Followed by tab twice quickly

genomics@harvar	d ami:~\$ ls /				
bin/	lib/	root/	usr/		
boot/	lib64/	run/	var/		
dev/	lost+found/	sbin/	vmlinuz		
etc/	media/	snap/	vmlinuz.old		
home/	mnt/	srv/			
initrd.img	opt/	sys/			
initrd.img.old	proc/	tmp/			
<pre>genomics@harvard_ami:~\$ ls /</pre>					

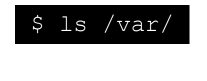
This shows the contents of the root directory

### If You're Typing, You're Doing Something Wrong!

Now type:



Followed by tab once. The path to the /var/ directory has filled in.



Now type: ls /var/r

Followed by tab once. The path to the /var/run/ directory has filled in.

\$



Tab complete will fill in paths, save you time in typing and prevent typos!

### If You're Typing, You're Doing Something Wrong!

Two more tricks for less typing!

\* Represents a special character For example:

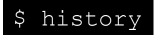
\$ ls /home/genomics/\*.txt

Will list everything in my home directory ending .txt

The up arrow can be used to re-run commands

Press your up arrow and see

If you want all of these commands listed, simply type



### Any Questions So Far?



# **Binary programs**

These are all programs installed on the Unix machine.

#### They can be found in /bin

#### \$ ls /bin

genomics@harvard am	i:~\$ ls /bin/				
bash	chmod	hciconfig	mv	pwd	systemd-tty-ask-password-agent
btrfs	chown	hostname	nano	rbash	tailf
btrfs-calc-size	chvt	ip	nc	readlink	tar
btrfsck	ср	journalctl	nc.openbsd	red	tempfile
btrfs-convert	cpio	kbd mode	netcat	rm	touch
btrfs-debug-tree	dash	kill	netstat	rmdir	true
btrfs-find-root	date	kmod	networkctl	rnano	udevadm
btrfs-image	dd	less	nisdomainname	run-parts	ulockmgr server
<pre>btrfs-map-logical</pre>	df	lessecho	ntfs-3g	sed	umount
btrfs-select-super	dir	lessfile	ntfs-3g.probe	setfacl	uname
btrfs-show-super	dmesg	lesskey	ntfs-3g.secaudit	setfont	uncompress
btrfstune	dnsdomainname	lesspipe	ntfs-3g.usermap	setupcon	unicode start
btrfs-zero-log	domainname	ln	ntfscat	sh	vdir
bunzip2	dumpkeys	loadkeys	ntfscluster	sh.distrib	vmmouse_detect
busybox	echo	login	ntfscmp	sleep	wdctl
bzcat	ed	loginctl	ntfsfallocate	SS	which
bzcmp	egrep	lowntfs-3g	ntfsfix	static-sh	whiptail
bzdiff	false	ls	ntfsinfo	<u>st</u> ty	ypdomainname
bzegrep	fgconsole	lsblk	ntfsls	su	zcat
bzexe	fgrep	lsmod	ntfsmove	sync	zcmp
bzfgrep	findmnt	mkdir	ntfstruncate	systemctl	zdiff
bzgrep	fsck.btrfs	mkfs.btrfs	ntfswipe	systemd	zegrep
bzip2	fuser	mknod	open	systemd-ask-password	zfgrep
bzip2recover	fusermount	mktemp	openvt	systemd-escape	zforce
bzless	getfacl	more	pidof	systemd-hwdb	zgrep
bzmore	grep	mount	ping	systemd-inhibit	zless
cat	gunzip	mountpoint	ping6	systemd-machine-id-setup	zmore
chacl	gzexe	mt	plymouth	systemd-notify	znew
chgrp	gzip	mt-gnu	ps	systemd-tmpfiles	

These include pwd, mkdir, ls ...

# Every binary program has a manual

To view the manual page, type man followed by the name of the program

\$ man <PROGRAMME>

Open the manual page for Is

Scroll through (enter) and find the options for: long listing format, human-readable sizes and sort by modification time

Exit the manual page (type q) and give these Is options a go in your Data directory



# Every binary program has a manual

To view the manual page, type man followed by the name of the program

\$ man <PROGRAMME>

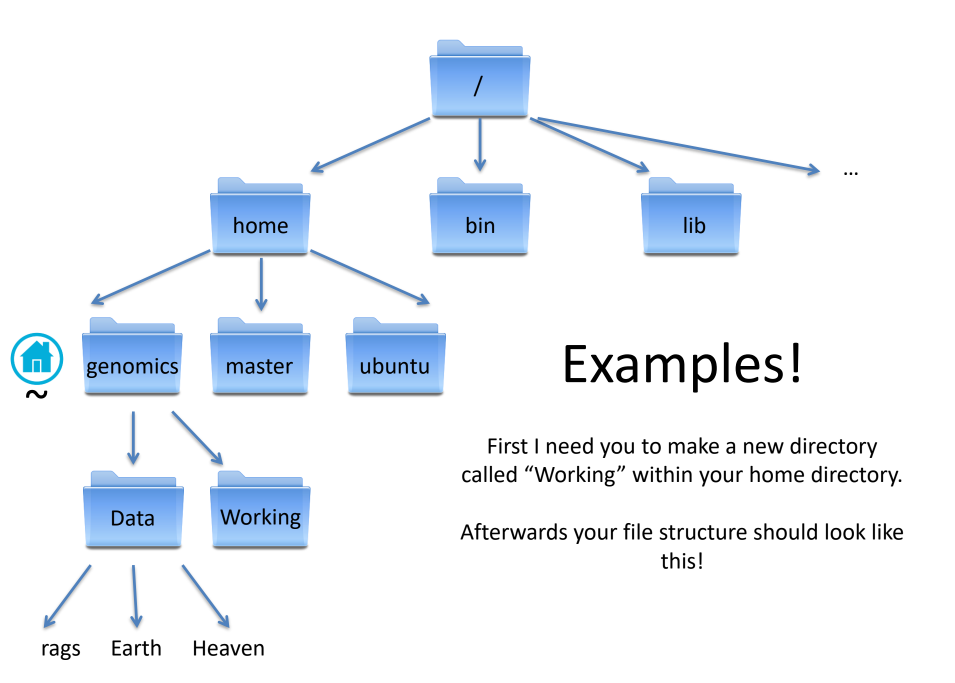
#### Open the manual page for Is

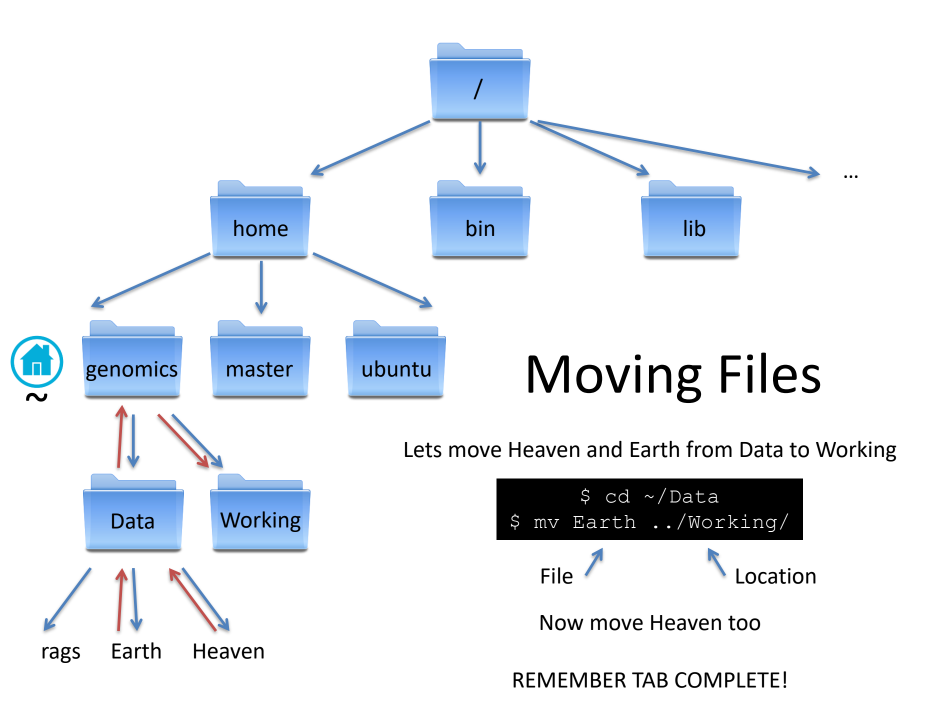
\$ man ls

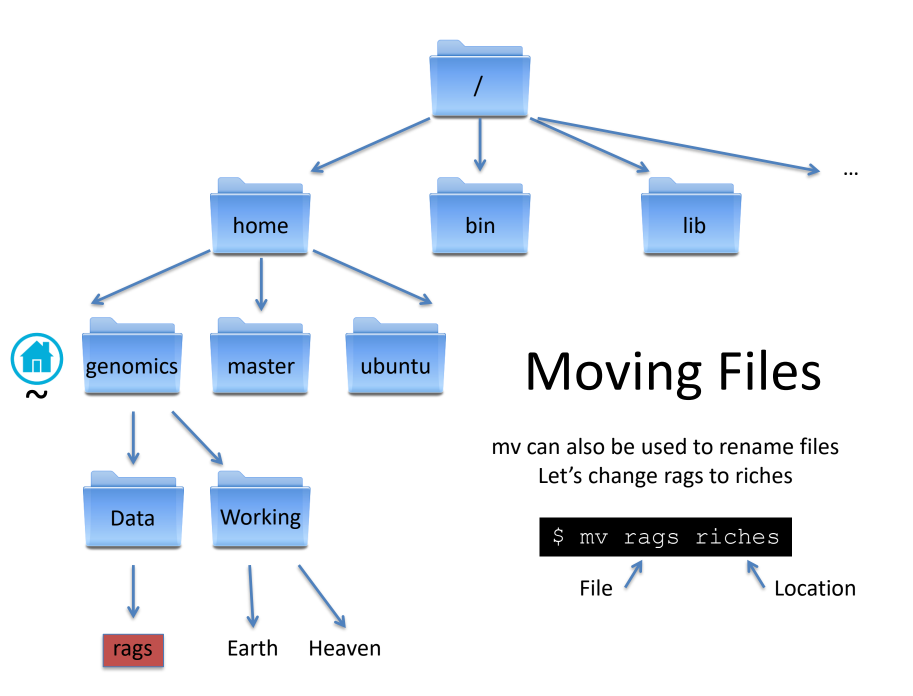
Scroll through (enter) and find the options for: long listing format (-l), human-readable Sizes (-h) and sort by modification time (-t)

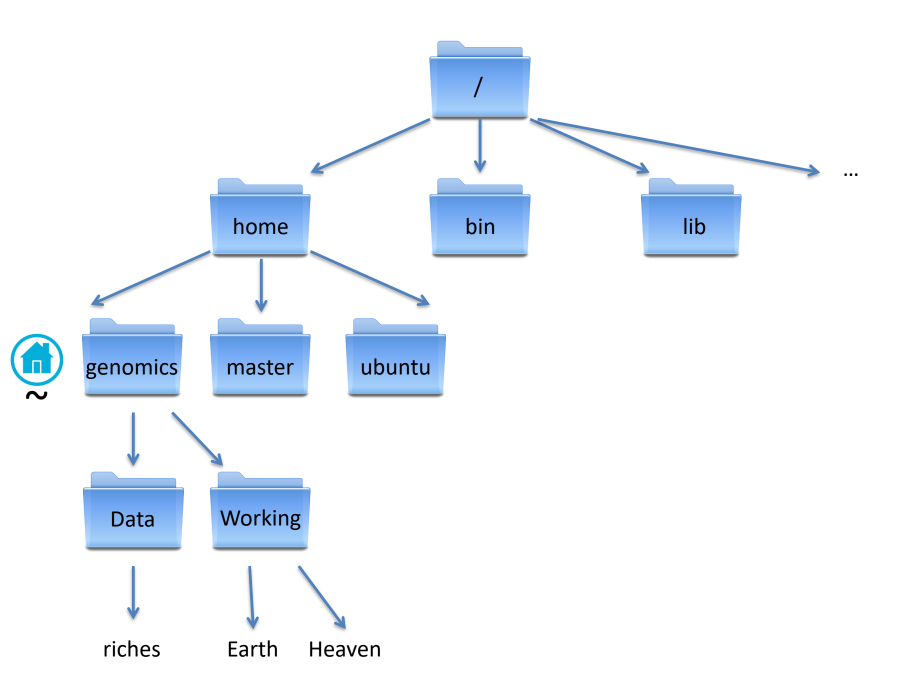
Exit the manual page (type q) and give these Is options a go in your Data directory

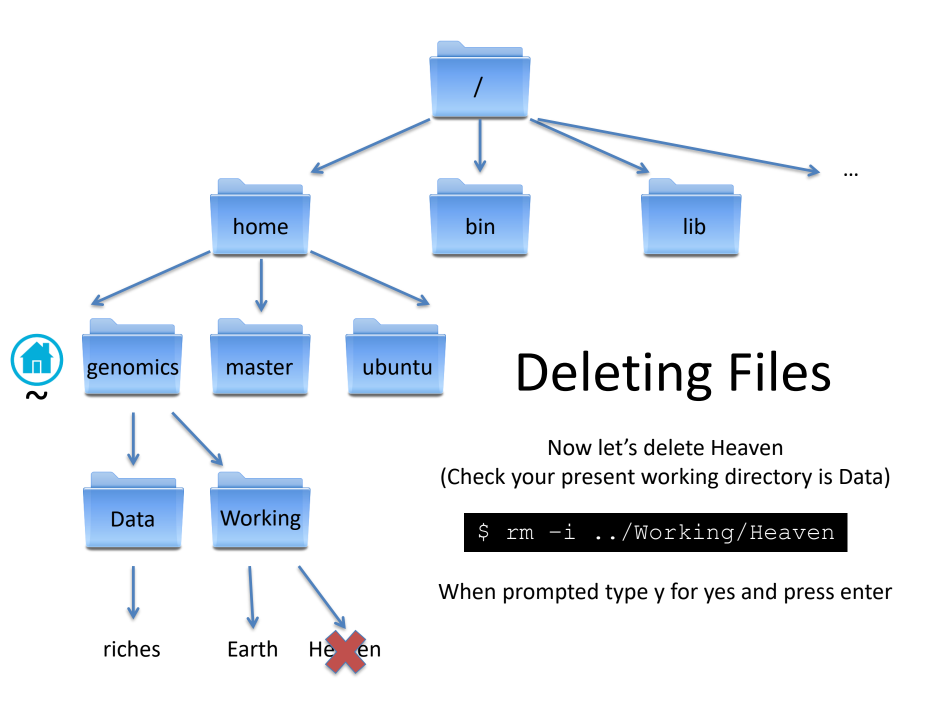


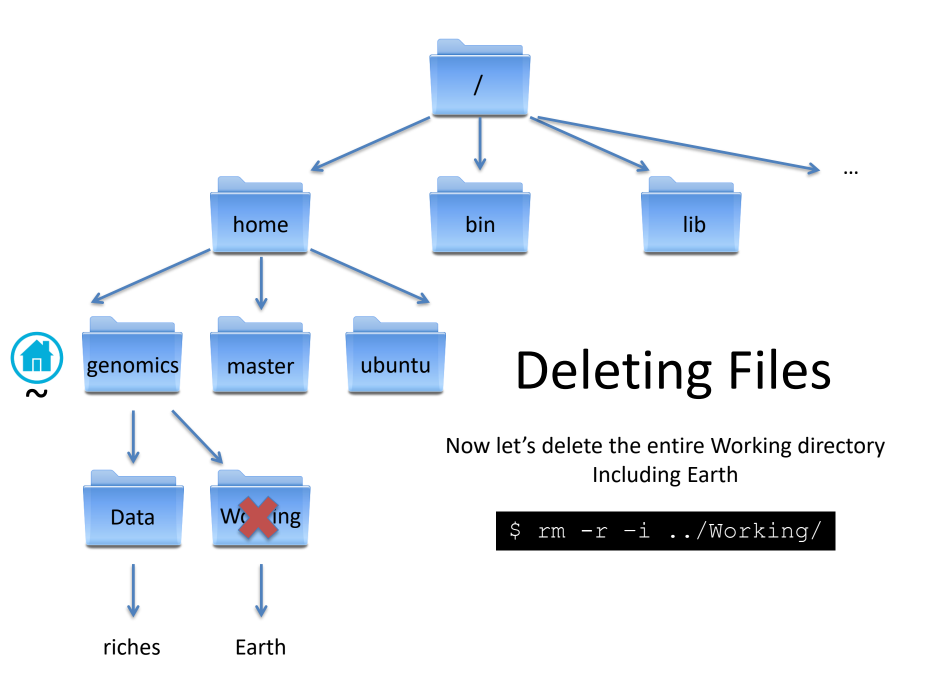


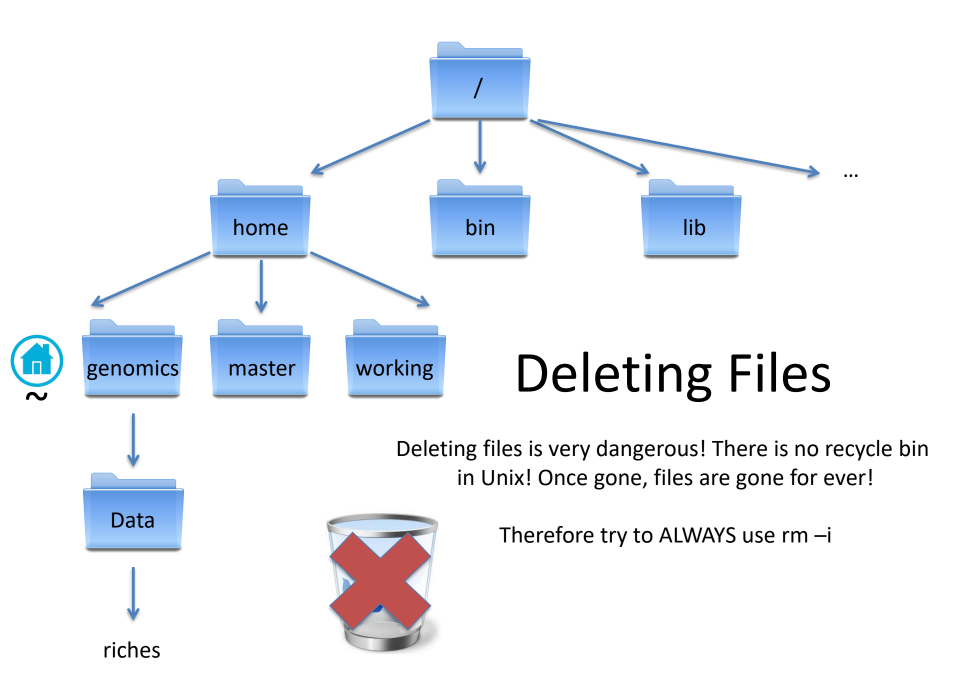


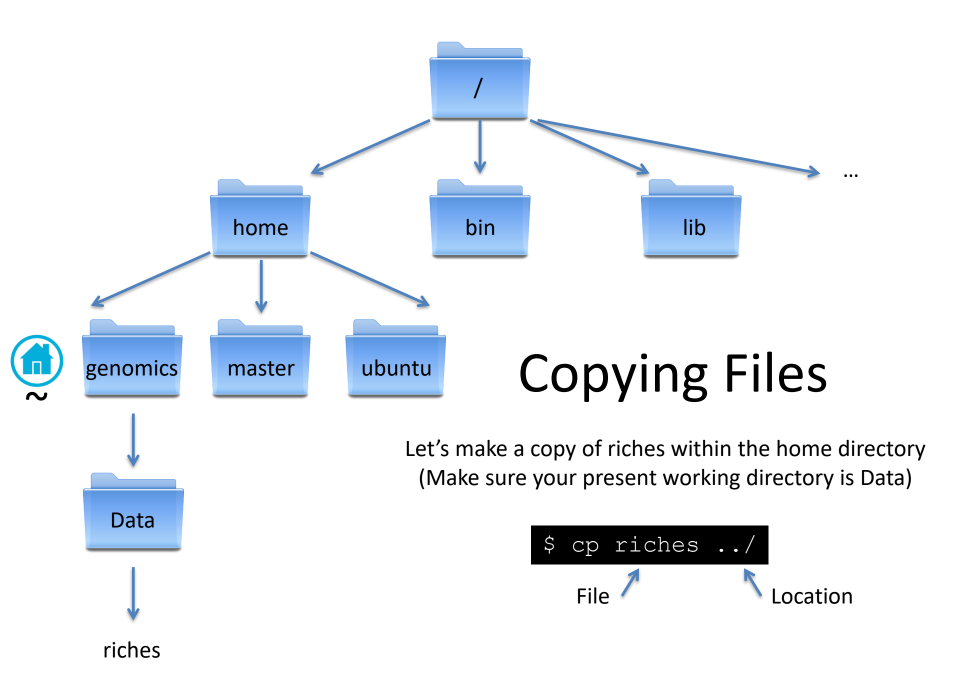


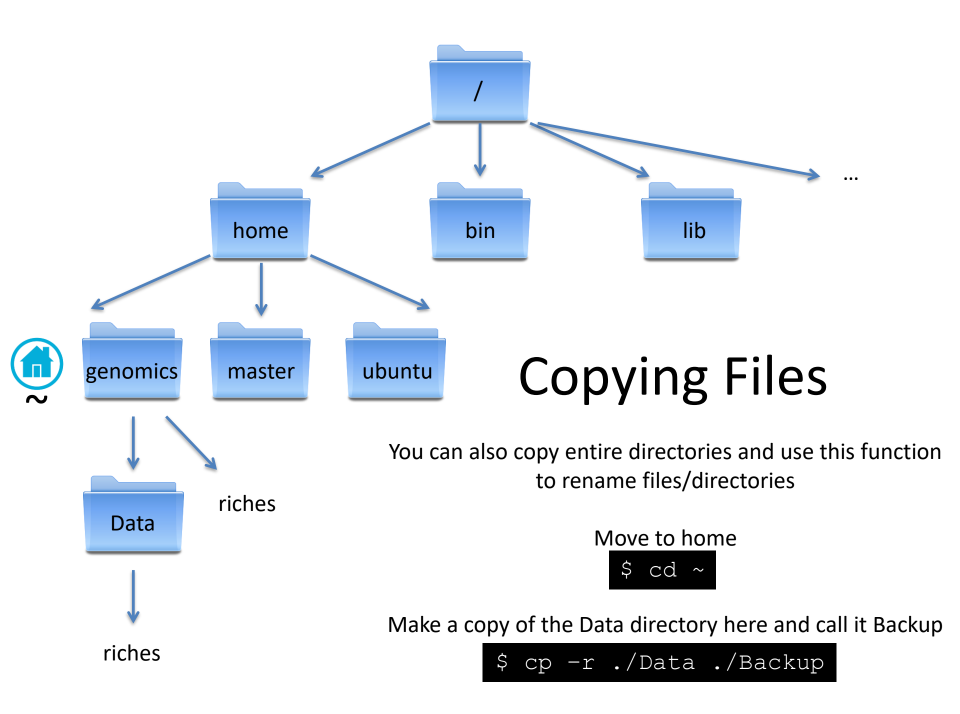


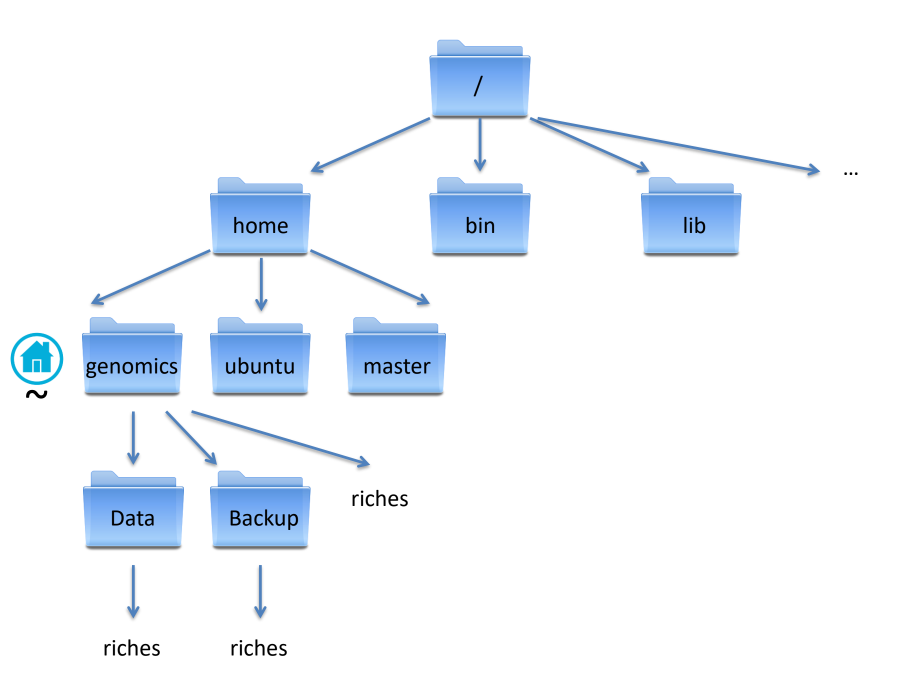












# **Typical File Sizes**



One Sequencing Sample on the Illumina NextSeq 3,000,000 reads = 1 Gb

But typically you will sequence more than one sample You may have different patients, different locations, replicates etc...

The size of the sequencing data file can easily become 100s of Gb

(or even bigger depending on the sequencer used)

# Archived/Compressed Files

Commonly, people will compress large files so that they are easier to store or share Here's an example: sequences.tar.gz

> .tar – means that it is a tape archive .gz – means that it is gzipped

These can be used alone or in combination

To uncompress A Tar Archive \$ tar -xvf <filename> (x = extract, v = verbose, f = all files)

A Gzipped file \$ gunzip <filename> A Gzipped Tar archive

5 tar -xzvf <filename>

### Any Questions So Far?



# Challenge 2!

1. Change to the unix\_workshop directory at the following path:

\$ cd ~/workshop\_materials/unix\_workshop
You should find a compressed directory:
 Sequences.tar

2. Make a copy of this file in a Backup directory

3. Un archive the directory

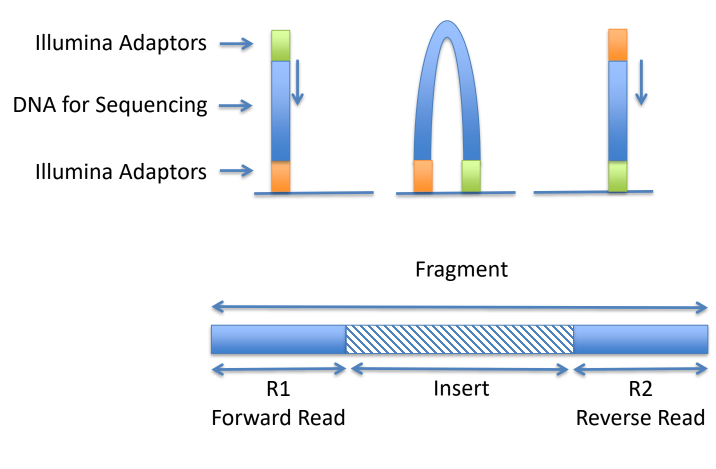
4. Unzip the read files

4. Rename the unarchived files – sequence\_1.fq and sequence\_2.fq

5. Delete the original .tar file

tar gunzip cp mv rm –i mkdir cd

### Paired Reads



An example: 300 bp paired end reads with a 700 bp fragment size R1 = 300 bp, R2 = 300 bp, Insert = 100bp

# Looking at File Contents

head	tail	more	less	cat
Shows the top lines of a file	Shows the bottom lines of a file	Shows the file one full screen at a time	Shows the file one full screen at a time	Shows an entire file all at once
-n specifies the number of lines (default 10)	-n specifies the number of lines (default 10)	Enter to scroll one line Space to scroll a page q to quit	Enter to scroll one line Space to scroll a page q to quit / to search	Ctrl + C to stop

Use these command line programs to look at the sequence files

# Let's put this to use

genomics@harvard\_ami:~/workshop\_materials/unix\_workshop\$ cd Sequences
genomics@harvard\_ami:~/workshop\_materials/unix\_workshop/Sequences\$ head sequence\_1.fq
@E.-371320/1
GCTGGTCAGCCAGGATAAAACCACCACTGACCCGATGGCGGTTGTTGACTGGATCAACATGTTTGCACTGGCAGTGAACGAAGAGAAACGCTGCTGG
CGGTCGCGGGGGGACTAAAAACCACCACTGACCGGTGCGGGGATTATCCCGGCAGTTCTGGCGTACTACGACAAGTTTATCCGCGAAGTGAACGCTGCTGA
CGGTCGCTGGTGACTGCGCGCGACTAACGGTGCGTGCGGGGATTATCCCGGCAGTTCTGGCGTACTACGACAAGTTTATCCGCGAAGTGAACGCTAA
CTCACTGGCTCGTTACCTGCTGGTAGCCAGCGCCATTGGTACTCTTTATAAGATGAAC
+
figgeright for the formation for the formation of t

HHHHFGIBEGHHHHHBFH=HHHFEEFGGEGHGDEBFG?FIFEG:8EBEEDG:GEEBGGGGGGGGGGGGGE?FECGFFCGFDFGFGEFEE??GG GCGE\*FG?:6E/FGCCEEHC:FGF-:G?6GCGGAAGGG6G)EGEC:GGFE'G;GC?G8

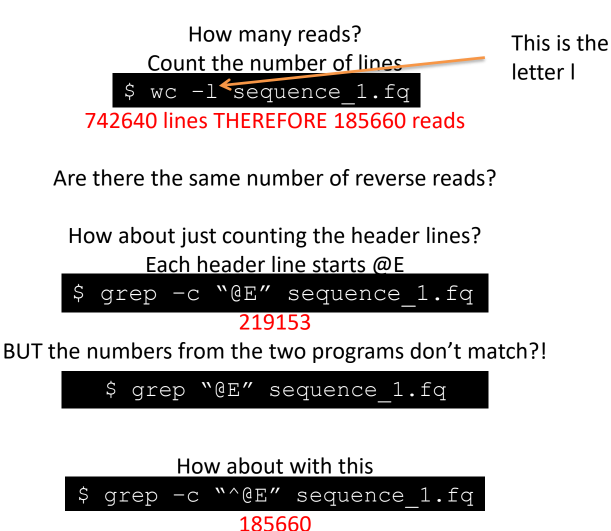
#### Fastq File Format:

Header		
Sequence		
Second Header (often +)		
Phred Quality Score		

Lot's of analysis software like paired reads to be in the same order

Use head to check that the top three headers are in the same order in sequence\_1.fq and sequence\_2.fq

## Sequencing Stats



^ matches this pattern at the start of the line – this is an example of a regular expression

### Any Questions So Far?



### AND NOW FOR A BRIEF SEGWAY INTO SCRIPTS...

## **Shell Scripts**

Imagine you have a complicated command to run. Take this as an example:

ref\_map.pl -o ./stacks\_gsnap/ -T 4 -O ./popmap -B middleton2\_ radtags -b 1 -s ./aligned\_gsnap/s13\_an\_01.bam -s ./aligned\_ gsnap/s13\_an\_02.bam -s ./aligned\_gsnap/s13\_an\_03.bam -s ./aligned\_gsnap/s13\_an\_04.bam -s ./aligned\_gsnap/s13\_an\_05.bam -s ./aligned\_gsnap/s13\_an\_06.bam -s ./aligned\_gsnap/s13\_an\_07 .bam -s ./aligned\_gsnap/s13\_an\_08.bam -s ./aligned\_gsnap/s13\_ fw\_01.bam -s ./aligned\_gsnap/s13\_fw\_02.bam -s ./aligned\_gsnap/ s13\_fw\_03.bam -s ./aligned\_gsnap/s13\_fw\_04.bam -s ./aligned\_ gsnap/s13\_fw\_05.bam -s ./aligned\_gsnap/s13\_fw\_06.bam -s ./aligned\_gsnap/s13\_fw\_07.bam -s ./aligned\_gsnap/s13\_fw\_08.bam

> But what if you make a mistake? Or want to run this command 10 times? You have to type it out every time 🔅

# **Shell Scripts**

Instead we can put this command inside a script.

Then it can easily be edited and ran multiple times

To understand shell scripts, we're going to look at a few topics:

- Shell scripting languages
  - Text editors
  - How to write a script
    - How to run a script

Scripts make a great record of what you've done, when and with what. You should also aim to keep a computational biology lab book.

# What is a Shell Script?

A computer program designed to be run by the Unix shell, the command line interpreter.

There are various types of shell scripts. These are scripting languages.

Today we are going to look at bash

First, let's run a simple bash command:

\$ echo Hello World

\$ echo Hello World Hello World

Try using echo with a different phrase

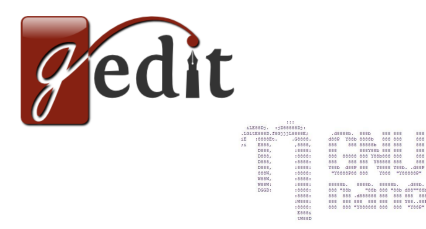
### **Text Editors**

These are pieces of software which can be used to write your script. Think of them as Unix versions of Notepad.

Some have an interactive user interface – E.G. gedit

Some work from within the command line – E.G. nano, vim, emacs

Today we are going to work with nano but have a play around with the others when you have a chance. Emacs and vim are notoriously difficult to use for the first time, so look up a cheat sheet.

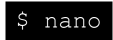






## Your First Script

#### Let's start by opening nano



N			Tutoria	I		_	o x
File Edit View							
GNU nano 2.	9.9	Ne	w Buffer			Modified	^
							=
^G Get Help	^0 Write	0ut ^R Read			^K Cut Text	^C Cur Pos	
^X Exit	^J Justi	.fy <mark>^W</mark> Wher	e Is 🔷	V Next Page	^U UnCut Text	^⊤ To Spell	$\sim$

## Key Nano Commands

**Ctrl + O** – This saves the file. You will be asked for a file name.

Type the name and press enter.



**Ctrl + X** – This exits nano. If the file is unsaved, you will be asked at this point if you'd like to save it.



#### **Your First Script**

Σ							T	lutorial									_ 0	×
File	Edit	View	Search	Terminal	Help													
GN	IU nai	no 2.	.0.9			Ne	w Buffe	r								Modif	ied	$\sim$
#!/b	in/b	ash																
echo	) "Th:	is is	s my fi	irst bas	sh scrip	)t."												
																		Ξ
^G (	iet H	eln	~0	Write(	)u+	^R Read	File	^Y	Prev I	Page	^K	Cut Te	avt	^C	Cur I	Pos		
	xit	erp	^3			^W When			Next I			UnCut			To S			$\overline{}$

#!/bin/bash tells the computer that this script is in the language bash. It always needs to go at the top of any bash script.

### Your First Script

E.		Tutorial		_ 0 X
File Edit View Search	Terminal Help			
GNU nano 2.0.9	New Bu	ffer		Modified
#!/bin/bash				
echo "This is my fi	rst bash script."			
				_
				=
File Name to Write:	firstscript sh			
^G Get Help	^T To Files	M-M Mac Format	M-P Prepend	
^C Cancel	M-D DOS Format	M-A Append	M-B Backup File	$\sim$

Then use Ctrl + O to save and give the file the name firstscript.sh.

Then use Ctrl + X to exit.

## Now Run Your Script

Simply Type:



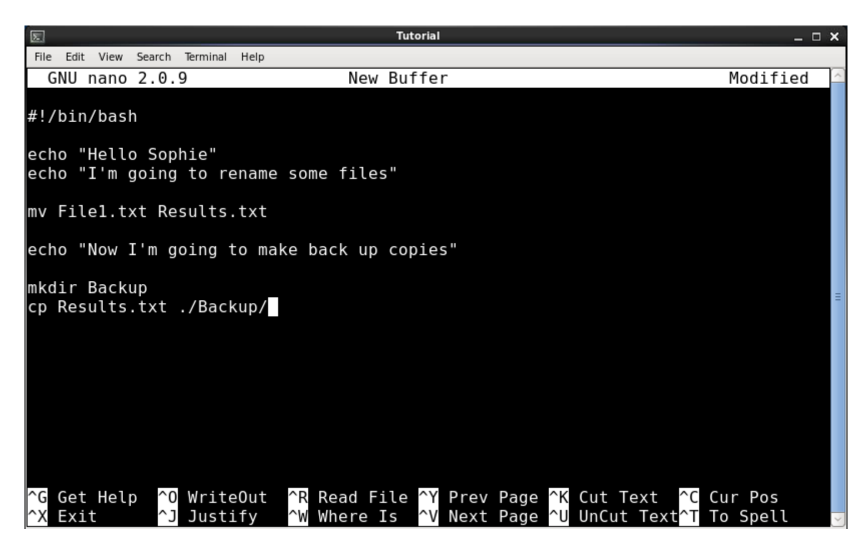
Reopen the same script:

\$ nano firstscript.sh

Change the phrase, save it and run the script again

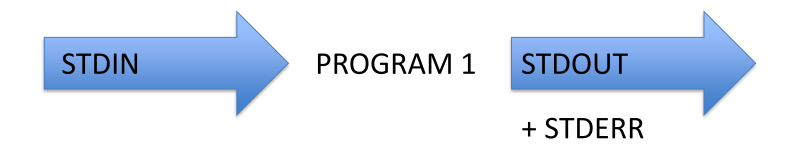
## **Bash Scripts**

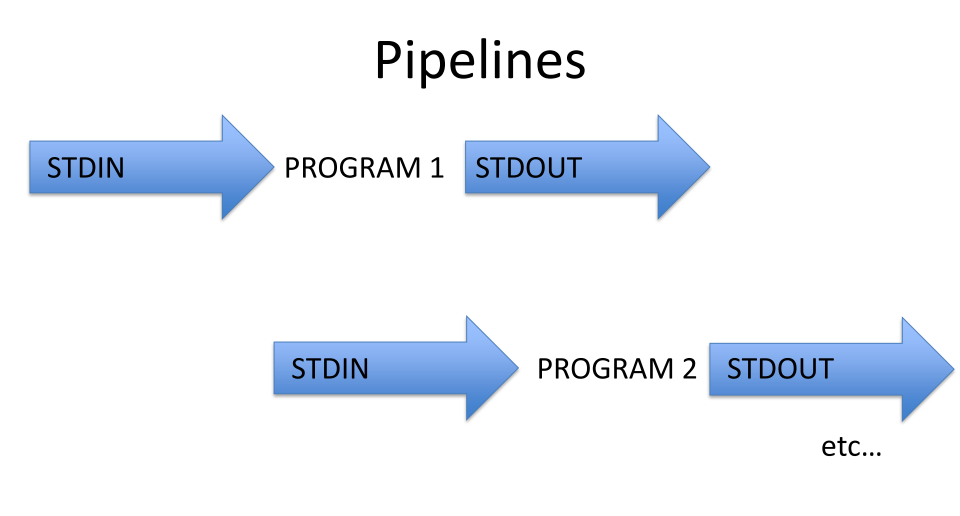
Bash scripts can be used to run binary programs like cd, mv, cp etc...



#### PIPELINES (TIME DEPENDING)

## Pipelines





## **Pipelines** PROGRAM 1 **STDIN** PIPE | PROGRAM 2 STDOUT etc...

### Let's put this to practice: Building Pipelines

Count the number of files and folders in your home directory

Let's build the first part of the pipeline, listing the files:

Number 1 =

\$ ls -1 /home/genomics/

PIPE this into wc –l to count the number of lines: (i.e. the number of files and folders)

Letter l

#### Let's put this to practice: Building Pipelines

How many base pairs in first sequence?

Firstly let's get the top two lines of the sequence file:

\$ head -n 2 sequence\_1.fq

Now let's PIPE this into tail to get just the sequence line

Finally PIPE this into word count of characters to count the base pairs

Is the first reverse read the same length?

Within the Unix Workshop directory you should find a file called scientists.txt

\$ cd ~/workshop materials/unix\_workshop/

Take a look at the contents

\$ more scientists.txt

First	Last	DOB
Charles	Darwin	12 February 1809
Marie	Curie	07 November 1867
Stephen	Hawking	08 January 1942
Rosalind	Franklin	25 July 1920
Isaac	Newton	04 January 1643
Richard	Dawkins	26 March 1941

#### \$ cat scientists.txt | cut -f 1

First
Charles
Marie
Stephen
Rosalind
Isaac
Richard

Now take a look at the original file

\$ more scientists.txt

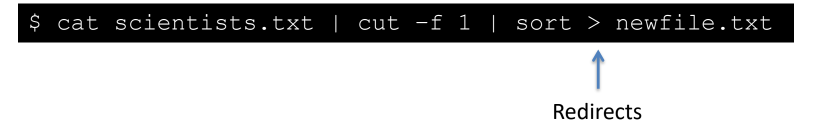
#### \$ cat scientists.txt | cut -f 1,3

First	DOB
Charles	12 February 1809
Marie	07 November 1867
Stephen	08 January 1942
Rosalind	25 July 1920
Issac	04 January 1643
Richard	26 March 1941

#### \$ cat scientists.txt | cut -f 1 | sort

Charles	
First	
Isaac	
Marie	
Richard	
Rosalind	
Stephen	

What if you wanted to keep the sorted list?



#### CHALLENGE 3!

Looking at the Saccharomyces cerevisiae gff3 file

GFF = general feature format

This is a file which lists all of the genome features, their coordinates, and info about them (genes, tRNAs, exons etc...)

##aff-version 3	
##sequence-region I 1 230218	
#sequence-region II 1 813184	
##sequence-region III 1 316620	
##sequence-region III 1 1531933	
#sequence-region IX 1 439888	
##sequence-region Mito 185779	
##sequence-region V1 55/874	
##sequence-region VII270161	
##sequence-region vi I 2 /0004 ##sequence-region VI I 1 090940	
##sequence-region VII 1 1000000	
##sequence-region X 1 745751	
##sequence_region XI 1 666816	
##sequence-region XII 1 1078177	
##sequence-region XIII 1 924431	
##sequence-region XIV 1 784333	
##sequence_region VXV 1 1891291	
##sequence-region XVI 1 948066	
#lgenome-build SGD R64-1-1	
#lgenome-version R64-1-1	
#!genome_date 2011-09	Screen Shot
#!genome-build-accession_SCA_000146045.2	
#Igenebuild-last-updated 2011-12	2016-0813.34
I SGD chromosome 1 230218 ID=chromosome:I;Alias=BK006935.2	The second
##	
I ensembl gene 335 649 . + . ID=gene:YAL069W;biotype=protein_coding;description=Dubious open reading frame%3B unlikely to encode a functional protein%2C based	on available experimenta
l and comparative sequence data [Source:SGDX3BAcc:S000002143];gene_id=YAL069W;logic_name=sgd	
I ensembl transcript 335 649 . + . ID=transcript:YAL069W;Parent=gene:YAL069W;Name=YAL069W;biotype=protein_coding;transcript_id=YAL069W	Screen Shot
I ensembl exon 335 649 . + . Parent=transcript:YAL069W;Name=YAL069W.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL069W.1;rank=1	2016-0813.30
I ensembl CDS 335 649 . + 0 ID=CDS:YAL069W;Parent=transcript:YAL069W;protein_id=YAL069W	2010-0010.00
I ensembl gene 538 792 . + . ID=gene:YAL068W-A;biotype=protein_coding;description=Dubious open reading frame%3B unlikely to encode a functional protein%2C bas	
tal and comparative sequence data%3B identified by gene-trapping%2C microarray-based expression analysis%2C and genome-wide homology searching [Source:SGD%3BAcc:S000028594];gene_id=YAL068W-A;1c	gic_name=sgd
I ensembl transcript 538 792 . + . ID=transcript:YAL068W-A;Parent=gene:YAL068W-A;Name=YAL068W-A;biotype=protein_coding;transcript_id=YAL068W-A	
I ensembl exon 538 792 . + . Parent=transcript:YAL0668W-A.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL068W-A.1;rank=1	
I ensembl CDS 538 792 . + 0 ID=CDS:YAL0668W-A;Parent=transcript:YAL0668W-A;protein_id=YAL0668W-A	
***	
I ensembl gene 1807 2169 . – . ID=gene:YAL068C;Name=PAU8;biotype=protein_coding;description=Protein of unknown function%3B member of the seripauperin multigene	family encoded mainly in
subtelomeric regions [Source:SGDX3BAcc:S000002142];gene_id=YAL068C;logic_name=sgd	
I ensembl transcript 1807 2169 . – . ID=transcript:YAL068C;Parent=gene:YAL068C;Name=PAU8;biotype=protein_coding;transcript_id=YAL068C	
I ensemble xono 1887 2169 Parent=transcript:X4L066C,Name=X4L066C,Liconstitutive=1ensembl_end.phase=8jexem_id=Y4L068C.1;rank=1	
I ensembl exon 1887 2169 - Parent=transcript:YAL068C;Name=YAL068C.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL068C.1;rank=1	
I ensembl exon 1887 2169 . – . Parent=transcript:YAL068C;Namé=YAL068C.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL068C.1;rank=1 I ensembl CDS 1887 2169 . – 0 ID=CDS:YAL068C;Parent=transcript:YAL068C;protein_id=YAL068C	54.700 inc.

#### CHALLENGE 3!

##aff-version 3	
##sequence-region I 1 230218	
##sequence-region II 1 813184	
#sequence-region III 1 316620	
#sequence-region IV 1 1531933	
##sequence-region IX 1 439688	
#sequence-region Mito 1 65779	
##sequence_region Y 1 576874	
##sequence-region VI 1 270161	
##sequence_region VII 1 1090940	
##sequence-region VIII 1 562643	
##sequence_region X 1 745751	
##sequence_region XI 1 666816	
##sequence-region XII 1 1078177	
##sequence_region XIII 1 924431	
##sequence-region XIV 1 784333	
##sequence-region XV 1 1891291	
##sequence_region XVI 1 948866	
#lgenome-build SGD R64-1-1	
#lgenome-version R64-1-1	
#Igenome-date 2011-89	
#lgenome-build-accession GCA_000146045.2	Screen Shot
#Jgenebuild-last-updated 2011-12	2016-0813.34
I SGD chromosome 1 230218 ID=chromosome:I;Alias=8K006935.2	And in case of the local division of the loc
***	
I ensembl gene 335 649 . + . ID=gene:YAL069W;biotype=protein_coding;description=Dubious open reading frame%38 unlikely to encode a functional protein%2C based on avail	lable experimenta
l and comparative sequence data [Source:SGDX3BAcc:S000082143];gene_id=YAL069W;logic_name=sgd	
I ensembl transcript 335 649 + ID=transcript:YAL069W;Parent=gene:YAL069W;biotype=protein_coding;transcript_id=YAL069W	Screen Shot
I ensembl exon 335 649 . + . Parent=transcript:YAL869W;Name=YAL869W.1;constitutive=1;ensembl_end_phase=8;exon_id=YAL869W.1;rank=1	2016-0813.3(
I ensembl CDS 335 649 . + 0 ID=CDS:YAL069W;Parent=transcript:YAL069W;protein_id=YAL069W	2010-0015.50
I ensembl gene 538 792 . + . ID=gene:YAL068W-A;biotype=protein_coding;description=Dubious open reading frame%3B unlikely to encode a functional protein%2C based on ava	
tal and comparative sequence dataX3B identified by gene-trappingX2C microarray-based expression analysisX2C and genome-wide homology searching [Source:SGDX3BAcc:S000028594];gene_id=YAL066W-A;logic_name=	=sgd
I ensembl transcript 538 792 . + . ID=transcript:YAL068W-A;Parent=gene:YAL068W-A;biotype=protein_coding;transcript_id=YAL068W-A	
I ensembl exon 538 792 . + . Parent=transcript:YAL068W-A;Name=YAL068W-A.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL068W-A.1;rank=1	
I ensembl CDS 538 792 . + 0 ID=CDS:YAL068W-A;Parent=transcript:YAL068W-A;protein_id=YAL068W-A	
###	
I ensembl gene 1807 2169 ID=gene:YAL068C;Name=PAU8;biotype=protein_coding;description=Protein of unknown function%3B member of the seripauperin multigene family en	ncoded mainly in
subtelomeric regions [Source:SGDX3BAcc:S000002142];gene_id=YAL068C;logic_name=sgd	
I ensembl transcript 1887 2169 ID=transcript:YAL068C;Parent=gene:YAL068C;Name=PAU8;biotype=protein_coding;transcript_id=YAL068C	
I ensembl exon 1807 2169 Parent=transcript:YAL068C;Name=YAL068C.1;constitutive=1;ensembl_end_phase=0;ensembl_phase=0;exon_id=YAL068C.1;rank=1	
I ensembl CDS 1807 2169 . – 0 ID=CDS:YAL068C;Parent=transcript:YAL068C;protein_id=YAL068C	
More(8%)	/20 ing

Lines that start # are comments – just run information

Column 1 = Chromosome

Column 3 = Feature/Type e.g. gene, chromosome, exon

Column 4 = Start Location

Column 5 = Stop Location

## CHALLENGE 3!

1. In the Unix workshop directory you should find a gff3 file.

\$ cd ~/Unix\_Workshop/Challenge5

Saccharomyces cerevisiae.R64-1-1.85.gff3.gz

2. Unzip the file.

3. How many feature entries are there?

- 4. List and count all the different types of features
  - 5. Which chromosome is the longest?

# Hints!

- Use head to work with 10 lines whilst testing what your pipe does!
  - This is a tab delimited file with a column layout.
  - Google "gff3 format" to find out what each of the columns are.
    - Remember that cat opens an entire file at once.
- There are a number of info lines at the start which begin with a hash. Look into grep with invert matches to skip these.
  - Cut can be used to isolate certain columns. You'll want the field option.
    - The programs sort and uniq may be helpful.
      - Sort must be used before uniq.
        - Uniq has a counting option.
    - Sort uses the key option to sort by a column.

more	gunzip	head	uniq
ср	mv	grep	WC
rm —i	mkdir	cut	
cd	cat	sort	man

#### Any Questions So Far?

