Bioinformatics for Beginners, January 2025



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Learning Objectives

NIH HPC Systems

Additional training materials at hpc.nih.gov

• What is a high performance cluster (HPC)?

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Bioinformatics for Beginners: RNA-Seq

Course Description

Bioinformatics integrates biology, statistics, and computer science to develop and apply theory, methods, and tools for the collection, storage, and analysis of biological and related data. Some key application areas in bioinformatics include genomic and molecular analysis, drug discovery and development, medical diagnosis and treatment, agricultural biotechnology, and environmental monitoring. The National Cancer Institute (NCI) uses bioinformatics extensively in its research efforts to combat cancer, including research on the "origin, evolution, progression, and treatment of cancer" (https://www.cancer.gov/research/infrastructure/bioinformatics#:~:text=NCI's%20Role%20in%20Cancer%20Bioinformatics,-NCI%20Data%20Initiatives&As%20a%20federal%20agency%2C%20NCI,sharing%2C%20analysis%2C%20analy

This course was designed to teach the basic skills needed for bioinformatics, including working

This course was designed to teach the basic skills needed for bioinformatics, including working on the Unix command line. This course primarily focuses on RNA-Seq analysis. All steps of the RNA-Seq workflow, from raw data to differential expression and gene ontology analysis, are covered. However, many of the skills learned are foundational to most bioinformatics analyses and can be applied to the analysis of other types of next generation sequencing experiments.

Why learn bioinformatics?

Here are a few compelling reasons to explore the world of bioinformatics:

- Analyze your data: Empower yourself to delve into your own biological data, gaining valuable insights.
- Enhancing Scientific Skills: Broaden your scientific knowledge and skills by mastering bioinformatics tools and techniques. By understanding the principles involved with data collection and analysis, you'll be better equipped to design robust experiments and interpret their results effectively.
- Career Opportunities: Open doors to exciting career paths in the rapidly growing field of bioinformatics.
- Understand the Data Landscape: Gain a deeper appreciation for how others analyze biological data, fostering collaboration and critical thinking.

This course is divided into three modules.

Module 1: Unix and Biowulf

Lessons focus on developing command line skills, getting started and working on Biowulf (the NIH HPC cluster), and downloading data from NCBI.

- Lesson 1 What is Biowulf?
- Lesson 2 Navigating file systems with Unix
- Lesson 3 Useful Unix
- Lesson 4 Working on Biowulf
- Lesson 5 Downloading data from the SRA

Module 2: RNA-Seq Analysis

Lessons focus on RNA-Seq analysis including experimental design and best practices, quality control, trimming, alignment based methods, feature counts, differential expression analysis, and biological interpretation.

- Lesson 6 Introduction to RNA-Seq
- Lesson 7 Introduction to Next Generation Sequencing (NGS) Data and Quality Control
- Lesson 8 Cleaning and Preparing Next Generation Sequencing (NGS) Data for Downstream Analysis
- Lesson 9 Aligning Next Generation Sequencing (NGS) Data to Genome
- Lesson 10 Quantifying Gene Expression from RNA Sequencing Data
- Lesson 11 Visualizing Genomic Data: Preparing Files
- Lesson 12 Visualizing Genomic Data with the Integrative Genomics Viewer
- Lesson 13: Differential Expression Analysis: QC
- Lesson 14: Differential Expression Analysis for Bulk RNA Sequencing: The Actual Analysis

Module 3: Pathway Analysis

Lessons focus on gene ontology and pathway analysis.

- Lesson 15: Introduction to gene ontology and pathway analysis
- Lesson 16: Functional enrichment with DAVID
- Lesson 17: Pathway Analysis with Reactome

Course requirements:

Who can take this course?

There are no prerequisites to take this course. This course is open to NCI-CCR researchers interested in learning bioinformatics skills, especially those relevant to analyzing bulk RNA sequencing data.

How will we work through lesson content?

For the hands-on sessions, participants will use Biowulf student accounts. To sign up for a student account, click here (https://docs.google.com/spreadsheets/d/1Epjzdtg9NxtJ8nm8Nti-l0I5S-5FOCsASe4hE4GXmaA/edit?usp=sharing). Student accounts are only available to course registrants.

Lesson content and practice questions can be found in these pages.

Class documents are available at https://bioinformatics.ccr.cancer.gov/docs/bioinformatics-for-beginners-2025/ (https://bioinformatics.ccr.cancer.gov/docs/bioinformatics-for-beginners-2025/).

Class data

Below are the links for the class data in case participants would like to practice outside of and after this course series. There is no need to download these for this course as the instructors have made them available on Biowulf. If you do not have access to Biowulf, see the below instructions.

Module 1

You can find compressed Module 1 data here. Download the data and unzip.

unzip module 1.zip

Module 2

All Module 2 data were obtained from the Griffith lab RNA sequencing tutorial (https://rnabio.org/) and renamed for this course series.

Reference genome can be downloaded at https://rnabio.org/module-01-inputs/0001/02/01/Reference_Genomes/ (https://rnabio.org/module-01-inputs/0001/02/01/Reference_Genomes/)

Annotation can be downloaded at https://rnabio.org/module-01-inputs/0001/03/01/Annotations/ (https://rnabio.org/module-01-inputs/0001/03/01/Annotations/)

See https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/ (https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/) for instructions on downloading the HBR-UHR and hcc1395 data.

Contact Us

Email ncibtep@nih.gov if you have any comments, questions, or concerns.



Biowulf HPC Unix

Lesson 1: What is Biowulf?

To fully engage with the course material and complete the hands-on exercises, we'll be leveraging the powerful NIH HPC Biowulf system.

To make the most of this powerful tool, it's essential to grasp the fundamentals of working with HPC systems, specifically Biowulf. In this lesson and others in Module 1, we will delve into the key concepts and practical skills you'll need to navigate this environment effectively.

Learning Objectives

- 1. Understand the components of an HPC system. How does this compare to your local desktop?
- 2. Learn about Biowulf, the NIH HPC cluster.
- 3. Learn about the command line interface and resources for learning.

Additional training materials at hpc.nih.gov

Much of the content for this presentation is from hpc.nih.gov. For more information and more detailed training documentation, see hpc.nih.gov/training/ (https://hpc.nih.gov/training/).

Note

This lesson will be demo-based only. Hands-on lessons will not begin until Lesson 2.

What is a high performance cluster (HPC)?

A collection of standalone computers that are networked together. They will frequently have software installed that allow the coordinated running of other software across all of these computers. This allows these networked computers to work together to accomplish computing tasks faster. --- hpc-intro (Software carpentries) (https://carpentries-incubator.github.io/hpc-intro/files/jargon.html#p6)

NIH HPC Systems

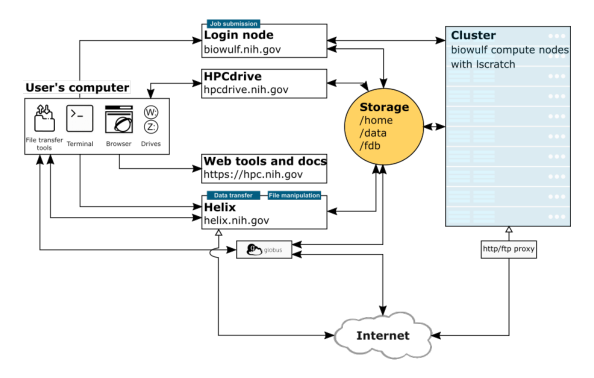


Image Credit: hpc.nih.gov/systems (https://hpc.nih.gov/systems/)

This diagram describes the NIH HPC Systems (Login node, Biowulf Cluster, HPC drive, and Helix), how the systems interact, and how you interact with the systems. Each "system" is described here (https://hpc.nih.gov/systems/). We will also discuss these further below.

When using an HPC

- We use a **command line interface** and a Secure shell protocol (SSH) to establish a remote connection to the login node / head node.
 - Most of us are likely used to a graphical user interface (GUI), which is a point-and-click interface, meaning we use a mouse to move around and click on various display icons. We visually interact with our compute environment. However, we do not generally interact with an HPC in this way. HPCs are remote resources that require connections using slow or intermittent interfaces (over WIFI and VPNs). Because of this, it is more practical to guide functionality over the command line using plain text. (https://carpentries-incubator.github.io/hpc-intro/11-connecting/index.html)
 - The cluster head node (login node) distributes compute tasks (the things we want to do outside of file manipulation and editing) using a scheduling system (e.g., SLURM). We use a scheduling system because the HPC is a shared resource with hundreds of worker nodes and thousands of processors.

Slurm stands for Simple Linux Utility for Resource Management.

What is Biowulf?

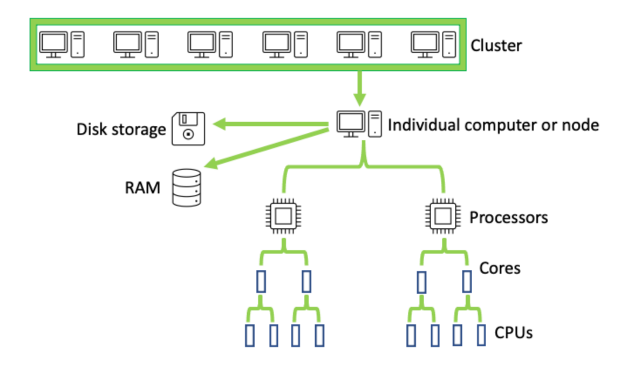
Biowulf is the National Institutes of Health's (NIH) high-performance computing (HPC) cluster. This massive Linux system boasts over 90,000 processors, allowing it to tackle enormous computational tasks by dividing jobs and running them simultaneously across multiple nodes. Biowulf comes pre-loaded with over 600 scientific software programs (https://hpc.nih.gov/apps/) and databases (https://hpc.nih.gov/apps/db.php) relevant to various fields like genomics, molecular biology, and bioinformatics. For security reasons, access to Biowulf is restricted to the NIH campus network or through a Virtual Private Network (VPN).

When should we use Biowulf?

You should use Biowulf when:

- Software is unavailable or difficult to install on your local computer and is available on Biowulf
- You are working with large amounts of data that can be parallelized to shorten computational time AND/OR
- You are performing computational tasks that are memory intensive.

Example of High Performance Computing Structure



Essentially Biowulf is a scaled up version of your local computer.

In Biowulf, many computers make up a cluster. Each individual computer or node has disk space for storage and random access memory (RAM) for running tasks. The individual computer is composed of processors, which are further divided into cores, and cores are divided into CPUs.

Info

Information on the NIH HPC architecture and hardware here (https://hpc.nih.gov/systems/hardware.html).

Getting an NIH HPC account

- If you do not already have a Biowulf account, you can obtain one by following the instructions here (https://hpc.nih.gov/docs/accounts.html).
- NIH HPC accounts are available to all NIH employees and contractors listed in the NIH Enterprise Directory.
- Obtaining an account requires PI approval and a nominal fee of \$40 per month.
- Accounts are renewed annually contigent upon PI approval.

Node types on the HPC

- Login node (head node)
 - Used for submitting resource intensive tasks as jobs
 - Editing and compiling code
 - File management and data transfers on a small scale
- Compute nodes (worker nodes)
 - For computational processes
 - Requires interaction with a job scheduling system (SLURM)
 - Batch jobs, sinteractive sessions
- Data transfer node (For Biowulf, this is Helix.)
 - more on this below

Info

sinteractive - work on biowulf compute nodes interactively; suitable for testing/debugging cpu-intensive code, Pre/post-processing of data, and using graphical applications.

sbatch - for submitting shell scripts via jobs, taking away any interactive component.

swarm - used for runnning embarassingly parallel code as independent jobs.

The Data transfer node: Helix

- Used for data transfers and file management on a large scale.
- 48 core system with 1.5 TB of main memory
- direct internet connection
- Helix should be used when
 - ∘ you are transferring >100 GB using scp
 - gzipping a directory containing >5K files, or > 50 GB
 - copying > 150 GB of data from one directory to another.
 - uploading or downloading data from the cloud.
- For more information on data transfers see hpc.nih.gov (https://hpc.nih.gov/docs/transfer.html).

Biowulf Data Storage

Summary of file storage options

	Location	Creation	Backups	Space	Available from
/home	network (NFS)	with Biowulf/Helix account	yes	16 GB default quota	B,C,H
/Iscratch (nodes)	local	created by user job	no	~850 GB shared	С
/scratch	network (NFS)	created by user	no	100 TB shared	В,Н
/data	network (GPFS/NFS)	with Biowulf/Helix account	no	100 GB default quota	В,С,Н

H = helix, B = biowulf login node, C = biowulf compute nodes

- You may request more space on /data, but this requires a legitimate justification.
- More information on data storage here (https://hpc.nih.gov/storage/).

Important

Data storage on the HPC system should not be for archival purposes.

Note

Though there aren't true back-ups of your data directories, there are snapshots with a view of your home and data directories at a specific point in time. You can learn more about snapshots (https://hpc.nih.gov/storage/backups.html) in the HPC documentation.

• To check disk space use:

checkquota - this shows the directories for which you have write access

OR

Look on the *user dashboard (https://hpcnihapps.cit.nih.gov/auth/dashboard) -> disk storage.

*Only works on VPN

Best practices file storage

BAD	GOOD
Submitting a swarm without knowing how much data it will generate	Run a single job, sum up the output and tmp files, and figure out if you have enough space before submitting the swarm
Directory with 1 million files	Directories with < 5,000 files
100 jobs all reading the same 50 GB file over and over from /data/\$USER/	Use /lscratch instead, copy the file there, and have each job access the file on local disk
100 jobs all writing and deleting large numbers of small temporary files	Use /Iscratch instead, have all tmp files written to local disk
Each collaborator having a copy of data on Biowulf	Ask for a shared area and keep shared files there to minimize duplication
Use Biowulf storage for archiving	Move unused or old data back to you local system

How do I create a directory in scratch?

mkdir /scratch/\$USER

Applications on Biowulf

- Bioinformatics applications and other programs are available on Biowulf via modules.
- View a list of available applications here (https://hpc.nih.gov/apps/).

Info

Loading software as environment modules allows us to better control our computational environment and easily use a large number of programs and even different versions of the same programs. Modules alter the user's environment variables such as the execution path.

The Command Line Interface (CLI)

What is Unix?

- Unix is a proprietary operating system like Windows or MacOS (Unix based).
- There are many Unix and Unix-like operating systems, including open source Linux and its multiple distributions.
- Biowulf nodes use a Unix-like (Linux) operating system (distributions RHEL8/Rocky8).
- Biowulf requires knowledge and use of the command line interface (shell) to direct computational functionality.
- To work on the command line we need to be able to issue Unix commands to tell the computer what we want it to do.

Tip

A basic foundation of Unix is advantageous for most scientists, as many bioinformatics open-source tools are available or accessible by command line on Unix-like systems.

Accessing your local terminal or command prompt

Mac OS

Type cmd + spacebar and search for "terminal". Once open, right click on the app logo in the dock. Select Options and Keep in Dock.

Windows 10 or greater

You can start an SSH session in your command prompt by executing ssh user@machine and you will be prompted to enter your password. ---Windows documentation (https://docs.microsoft.com/en-us/windows/terminal/tutorials/ssh? source=recommendations)

To find the Command Prompt, type cmd in the search box (lower left), then press Enter to open the highlighted Command Prompt shortcut.

Are you a Windows user and not affiliated with NIH?

If you are using a Windows operating system, Windows 10 or greater, you can use the Windows Subsystem for Linux (https://docs.microsoft.com/en-us/windows/wsl/) (WSL) for your computational needs.

The Windows Subsystem for Linux (WSL) is a feature of the Windows operating system that enables you to run a Linux file system, along with Linux command-line tools and GUI apps, directly on Windows, alongside your traditional Windows desktop and apps. --- docs.microsoft.com (https://docs.microsoft.com/en-us/windows/wsl/faq)

To install WSL, follow the instructions here (https://learn.microsoft.com/en-us/windows/wsl/install). There are multiple Linux distributions. We recommend new users install "Ubuntu".

Windows WSL is not available to NIH employees due to security policies.

How much Unix do we need to learn?

As with any language, the learning curve for Unix can be quite steep. However, to work on Biowulf you really need to understand the following:

- Navigating the File System: Understanding the hierarchical structure of directories, using the cd command to move between directories.
- File Paths: Learning how to specify the location of files using absolute and relative paths.
- Basic Unix Commands: Getting acquainted with common commands like 1s for listing files, mv for moving files, rm for removing files, mkdir for creating directories, cat for viewing file contents, and man for accessing command documentation.
- Getting help: Discovering how to find more information about Unix commands and their usage.
- Command Customization: Learning how to modify the behavior of Unix commands using flags or options, such as using 1s -1 to list files with detailed information compared to the basic 1s command.
- Redirecting Input and Output: Understanding standard input and output, and how to redirect the output of one command to the input of another using pipes (|) or redirection operators (>, <).

More on these in the next lesson.

Connecting to Biowulf

- To connect to Biowulf, we use a secure shell (SSH) protocol.
 - used to open an encrypted network connection between two machines, allowing you to send & receive text and data without having to worry about prying eyes. (https://carpentries-incubator.github.io/hpc-intro/11-connecting/index.html)
 - ∘man ssh

Establishing a remote connection

ssh username@biowulf.nih.gov

"username" = NIH/Biowulf login username.

Note

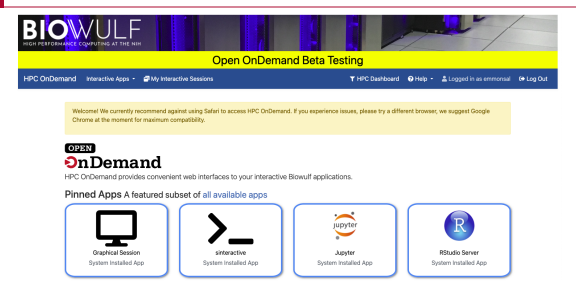
If this is your first time logging into Biowulf, you will see a warning statement with a yes/no choice. Type "yes".

Type in your password at the prompt. The cursor will not move as you type your password!

HPC OnDemand

Recently, the NIH HPC Team has provided on demand access to HPC resources via web browser through integration of Open OnDemand. This integration makes working with HPC resources less intimidating for new users, as they will not have to open a terminal and remotely connect via ssh. Instead, navigate to your web browser (Google Chrome is preferred) and connect to NIH HPC OnDemand using https://hpcondemand.nih.gov/).

HPC OnDemand (https://hpc.nih.gov/ondemand/index.html) provides an online dashboard for users to easily access command line interactive sessions, graphical linux desktop environments, and interactive applications including RStudio, MATLAB, IGV, iDEP, VS Code, and Juptyer Notebook.



HPC OnDemand Dashboard

SLURM commands

You will also need to know commands specific to the Biowulf job scheduling system:

- sbatch submit slurm job
- swarm submit a swarm of commands to cluster
- sinteractive allocate an interactive session
- sjobs show brief summary of queued and running jobs
- squeue display status of slurm batch job
- scancel delete slurm jobs

We will talk about many of these in more detail in later lessons.

How to load / unload a module

• To see a list of available software in modules use

```
module avail
module avail [appname|string|regex]
module -d
```

To load a module

```
module load appname/version
```

• To see loaded modules

module list

To unload modules

```
module unload appname
module purge #(unload all modules)
```

Note

You may also create and use your own modules.

Getting help on Biowulf: NIH HPC Documentation

The NIH HPC systems are well-documented at hpc.nih.gov (https://hpc.nih.gov/).

- User guides (https://hpc.nih.gov/docs/user_guides.html)
- Training documentation (https://hpc.nih.gov/training/)
- How To (https://hpc.nih.gov/docs/how_to.html) docs

Note

Existing safeguards make it nearly impossible for individual Biowulf users to irreparably mess up the system for others

WORST CASE SCENARIO - You are locked out of your account pending consultation with NIH HPC staff

Additional HPC help

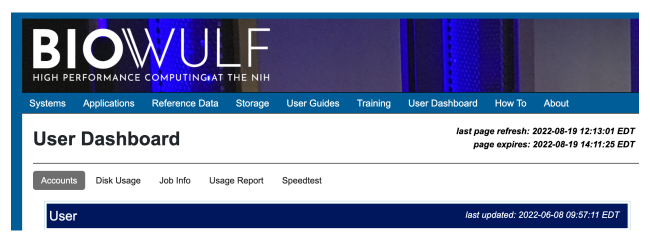
- Contact staff@hpc.nih.gov (mailto:staff@hpc.nih.gov)
 The HPC team welcomes questions and is happy to offer guidance to address your concerns.
- Monthly Zoom consult sessions

The HPC team offers monthly zoom consult sessions. "All problems and concerns are welcome, from scripting problems to node allocation, to strategies for a particular project, to anything that is affecting your use of the HPC systems. The Zoom details are emailed to all Biowulf users the week of the consult." (https://hpc.nih.gov/training/)

• Bioinformatics Training and Education Program

If you experience any difficulties or challenges, especially with different bioinformatics applications, please do not hesitate to email us at BTEP (mailto:ncibtep@nih.gov).

User Dashboard



- Can view disk usage and job info
- Request more disk space
- Evaluate job info for troubleshooting

Learning Unix: Classes / Courses

- Introduction to Biowulf (May Jun, 2023) (https://bioinformatics.ccr.cancer.gov/docs/biowulf-introduction-summer-2023/index.html)
- Introduction to Unix on Biowulf (Jan Feb, 2023) (https://bioinformatics.ccr.cancer.gov/docs/unix-on-biowulf-2023/index.html)
- Bioinformatics for Beginners: Module 1 Unix/Biowulf (https://bioinformatics.ccr.cancer.gov/docs/b4b/Module1_Unix_Biowulf/Lesson1/)

Additional Unix Resources:

- BashScripting_LinuxCommands from the NIH HPC team (https://hpc.nih.gov/training/handouts/BashScripting_LinuxCommands.pdf)
- Fosswire linux reference sheet (https://bioinformatics.ccr.cancer.gov/docs/b4b/ fosswire_reference.pdf)

Key points

- Biowulf is the high performance computing cluster at NIH.
- To work on Biowulf, you will need to use the command line interface, which requires some knowledge of unix commands.
- When you apply for a Biowulf account you will be issued two primary storage spaces:
 - 1. /home/\$User (16 GB)

2. /data/\$USER (100 GB).

- Hundreds of pre-installed bioinformatics programs are available through the module system.
- Computational tasks on Biowulf should be submitted as a job (sbatch, swarm) or through an interactive session (sinteractive).
- Do not run computational tasks on the login node.

Unix Command line

Lesson 2: Navigating file systems with Unix

Lesson 1 Review

- Biowulf is the high performance computing cluster at NIH.
- When you apply for a Biowulf account you will be issued two primary storage spaces: 1) /home/\$User and 2) /data/\$USER, with 16 GB and 100 GB of default disk space.
- Hundreds of pre-installed bioinformatics programs are available through the module system.
- Computational tasks on Biowulf should be submitted as a job (sbatch, swarm) or through an interactive session sinteractive.
- Connect to Biowulf using HPC OnDemand (https://hpc.nih.gov/ondemand/index.html) or ssh.
- Do not run computational tasks on the login node.

Lesson Objectives

In lesson 1, you learned about the NIH HPC cluster, Biowulf. Biowulf nodes use a Unix-like (Linux) operating system (distributions RHEL8/Rocky8), which requires knowledge and use of the command line interface (shell) to direct computational functionality. The purpose of today's lesson is to get you familiar with working on the command line. To this end, we will...

- Learn the basic structure of a unix command.
- Learn how to navigate your file system, including absolute vs relative directories.
- Learn unix commands related to navigating directories, creating and removing files or directories, and getting help.

A word about mistakes

YOU WILL MAKE MISTAKES...but, it is okay. We all make mistakes, and mistakes are how we learn. Remember, existing safeguards make it nearly impossible for individual Biowulf users to irreparably mess up the system for others. However, you can make your life difficult, for example, by misusing commands, ignoring existing tools, overwriting files, failing to redirect or output results, disregarding warnings, and not seeking help.

How can we overcome mistakes?

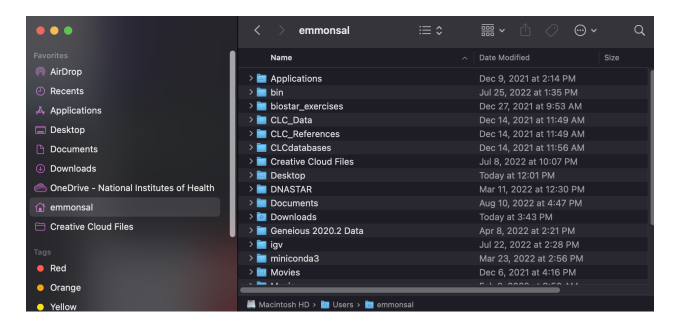
We practice. The more you use unix and bash scripting the better you will become.

You will need to learn how to troubleshoot error messages. Often this will involve googling the error in reference to the entered command. There are many forums that post help regarding specific errors (e.g., stack overflow, program repositories such as github).

File system

We manage files and directories through the operating system's file system. A directory is synonymous with a "folder", which is used to organize files, other directories, executables, etc.

On a Windows or Mac, we usually open and scroll through our directories and files using a GUI. For example, **Finder** is the default file management GUI from which we can access files or deploy programs on a macbook.



This same file system can be accessed and navigated via command line from the unix shell.

Some useful unix commands to navigate our file system and tell us some things about our files

- 1. pwd (print working directory)
- 2. 1s (list)
- 3. touch (creates an empty file)
- 4. nano (basic editor for creating small text files)
- 5. using the rm command to remove files. Be careful!
- 6. mkdir (make a directory) and rmdir (remove a directory, must be empty of all files)
- 7. cd (change directory), by itself will take you home, cd . . (will take you up one directory), cd /results dir/exp1 (go directly to this directory)
- 8. mv (for renaming files or moving files)

- 9. less (for viewing files, or more)
- 10. man (for viewing the man pages when you need help on a command)
- 11. cp (copy) for copying files

Getting Started

We have already seen some unix commands relevant to Biowulf. For example, we learned about ssh. The ssh command is used to securely log into a remote machine and execute commands on that machine. (https://www.tutorialspoint.com/common-ssh-commands-in-linux-with-

examples#:~:text=The%20ssh%20command%20is%20used,hostname%20of%20the%20remote%20machin

ssh is the command and username@biowulf.nih.gov is a command line argument, where username is the username that you wish to connect to on the remote system and biowulf.nih.gov is the hostname of the remote machine. For this lesson and the lessons that follow, we will use NIH HPC student accounts (https://hpc.nih.gov/nih/student.html) to connect to Biowulf.

More about student accounts

At the beginning of each class you must sign up for a student account. You can sign up for a student account using a Google spreadsheet, the link for which will be supplied at the beginning of each class via Webex. Click on the supplied link, and find an empty slot under the "Name" column. Type your name in the empty slot. The username under "Account Username" will now serve as your username for logging in to Biowulf.

This task will be repeated at the beginning of each lesson to allow students the option of flexible attendance.

Let's go ahead and get connected. Open a terminal and type the following:

ssh username@biowulf.nih.gov

username = NIH/Biowulf login username. Remember to use the student account username here.

Note

If this is your first time logging into Biowulf, you will see a warning statement with a yes/no choice. Type "yes".

Type in your password at the prompt. The cursor will not move as you type your password!

Success

We will connect to Biowulf at the beginning of each session.

We are now on the login node. Remember, you should not do work on the login node. However, you can do basic file management on the login node and edit and compile code. For now, we will stay on the login node.

Our Second Unix Command (1s)

Let's continue learning about the structure of linux commands using another common command, 1s. The 1s command "lists" the contents of the directory you are in. You may see files and other directories here.

1s

At this point, you are in your home directory, and so you will see whatever files and directories are located here. For example, if I had logged in to my Biowulf account, I would see the following:

```
Desktop R bin ncbi_error_report.txt
```

However, since this is a student account, I do not see anything, as I have not yet added any files.

How can you tell the difference between a file and a directory?

We can add some additional options (flags) to our command.

```
ls -lh
```

will show permissions and indicate directories (d). The -1h are flags. -1 refers to listing in long format, while -h provides human readable file sizes.

Or, many systems offset directories and files using colors (e.g., blue for directories). If you don't see colorized output, try the -G flag.

We can also label output by adding a marker to indicate files, directories, links, and executables using the -F flag.

ls -F

a terminal / = directory

a@=link

a * = executable

Anatomy of a command

Using 1s as an example, we can get an idea of the overall structure of a unix command.

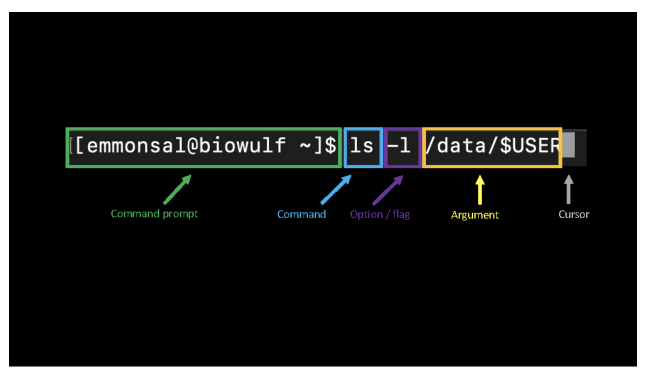


Image inspired by "Learn Enough Command Line to Be Dangerous"

The first thing we see is the command line prompt, usually \$ or %, which will vary by operating system. The prompt let's us know that the computer is waiting for a command. Next we see the actual command, in this case, 1s, telling the computer to list the files and directories. Most commands will have various options / flags that can be included to modify the command function. We can also supply an argument, which in the case of 1s is optional. For example, here we supplied an alternative directory from which we are interested in listing files and directories. We hit enter or return after each command, and when the command has finished running, the command prompt will reappear prompting us to enter more commands.

Where am I? (pwd)

pwd

pwd stands for "print working directory". When you run this, you should see something like this.

/home/username

where username is your name or student account. This is your home directory - where you start from when you open a terminal. This is an example of a "path". The path tells us the location of a file or directory.

Note

While Windows computers use a \ as a path separator, unix systems use a /.

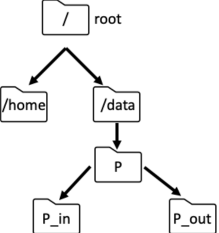
Therefore, the pwd command is very helpful for figuring out where you are in the directory structure. If you are getting "file not found" errors while trying to run something, it is a good idea to pwd and see if you are where you think you are. Type the pwd command and make a note of the directory you are in.

More on the home directory (~)

We see that we are in our home directory. But where is that exactly?

The file system on any computer is hierarchical. On a Unix system, the top level of the file system, or root directory, is denoted by /. All subdirectories on the file system branch from this root directory. See the below example.

File system hierarchy



Example of file system hierarchy structure.

In our example hierarchy, we have subdirectories /home and /data, and within data, we see additionally subdirectories, P_in and P_out. Only the first / denoted a directory (root). All other /s in the path serve as separating characters.

Absolute vs Relative directories

A file path that starts with the root or / is known as an absolute path. A path that does not start with the root directory is called a relative path. For example, in Unix, . is used to denote here in the present working directory and . . is used to denote one directory back. Thus, a path that starts with . or . . is a relative path. Going back to pwd. The output (/home/username) is an absolute file path. Absolute file paths will break scripts when collaborating because the likelihood that your file system matches another's is low.

We can use the tree command to get an idea of the structure of our home directory on Biowulf.

Getting around the Unix directory structure (cd)

How do we navigate this directory tree. We use cd, which means "change directory". Let's change directory to our data directory, which is the larger of the two allocations we are allotted on Biowulf.

```
cd /data/$USER # (1) change to your data directory
pwd #print working directory
ls #list the contents of /data/$USER
```

\$USER and other environment variables

~

\$USER is an example of an environment variable.

Environment variables contain user-specific or system-wide values that either reflect simple pieces of information (your username), or lists of useful locations on the file system. --- Griffith Lab (https://pmbio.org/module-01-setup/0001/03/01/Unix_Primer/)

We can display these variables using echo.

echo \$USER echo \$HOME

\$PATH is an important environment variable.

echo \$PATH

This results in a colon separated list of directories containing programs that you can run without specifying those directories each time you run the program.

You will likely need to add to your \$PATH at some point in the future.

To do this use:

export PATH=\$PATH:/path/to/folder

This change will not remain when you close the terminal. To permanently add a location to your path, add the above line to your bash shell configuration file, ~/.bashrc.

By itself, the cd command takes you home. Let's try that, and then do a pwd to see where we are.

cd pwd We are back in our home directory.

```
/home/username
```

How can we go back to the /data/\$USER directory? We need to give the "path" to that directory.

```
cd /data/$USER
pwd
ls
```

Check where you are with pwd and look at the contents of the directory with 1s. What do you see?

Home shortcut

We can also use ~ as a shortcut for our home directory.

ls ~

Once we create more files and directories, we can learn a bit more about the directory structure and absolute vs relative file paths.

Creating files (touch)

The touch command creates a file, but the file is empty, so it is not a command you will use very often, but good to know about.

```
touch file1.txt
touch file2.txt
ls
```

Now we see something like this.

```
file1.txt file2.txt
```

The nano editor is a text editor useful for small files.

```
nano file2.txt
```

Let's put something in this file.

```
Unix is an operating system, just like Windows or MacOS.
Linux is a Unix like operating system;
sometimes the names are used interchangeably.
```

Nano commands for saving your file and exiting nano:

- 1. control O write file (equivalent to save as)
- 2. File name to write: file2.txt (Hit return/enter on your keyboard to save the file with this name).
- 3. control X to Exit

This brings us to our next topic which is very important!

Avoid spaces in file names and directories.

There should not be spaces in Unix file names or directories. There are many strategies that can be used to avoid spaces in file names (https://builtin.com/articles/pascal-case-vs-camel-case#:~:text=Pascal%20case%20is%20a%20casing,capitalized%2C%20such%20as%3A%20camelCase%2. Though, consistency is key. One good method is using snake_case, in which words are separated by an _.

For example, we can use the underscore (_) where a space would go, like this, to name a directory for module 1.

```
Module_1
```

To use snake_case with file names, we may see something like this:

```
brain_rna.fastq
liver_rna.fastq
```

The first part of the file name provides info about the file, and the extension (.fastq) tells what kind of file it is. (Examples of file extensions are .csv, .txt, .fastq, .fasta and many more.)

More on file organization to come.

Understanding file extensions

It's important to understand file extensions, to know what kinds of data you are working with.

.txt are text files. These are likely but not always tab delimited.

.tsv are tab delimited files.

.csv are "comma-separated values" - good for importing into MS Excel spreadsheets

.tar.gz indicates a tarred and zipped file - so it is a compressed file

.fastq tells you that these are FASTQ files, containing sequence data and quality scores

.fasta indicates FASTA formatted sequence data, either protein or nucleotide

Removing files with rm

Warning

A Unix system will delete something when you ask to delete it and there is usually no way of getting it back. Be extremely careful when removing files and directories.

By adding the -i option, the system will ask if you're sure you want to delete. Generally speaking, when a file on a Unix system is deleted, it is gone.

rm -i file1.txt

will remove a file we created.

Creating directories (mkdir)

A couple things to note - this is a good time to give your directories meaningful names, which will help you keep things organized. Organization is key. I generally like to have a new directory per project, and within that directory, subdirectories separating raw data from analysis files. From there, each analysis would also get its own subdirectory. However, there are many ways to organize files and you should do whatever makes sense for your data and helps you (and others) stay organized.

Keep raw data raw

Always keep your raw data raw, and save outputs to new files. Do not overwrite raw data! Consider setting the permissions on these files to "read only". More on permissions later.

For now, let's create a directory called Module_1, where we can store Module 1 lesson content. To create a directory, we use mkdir.

mkdir Module_1

Removing directories (rmdir)

Directories must be completely empty of all files and other contents before you can delete them with rmdir. There are ways to "recursively" remove files and directories using the -r option of rm, for example (rm -r directory). This would remove all of the files and subdirectories in our hypothetical directory. Keep in mind that once these files are deleted they are gone for good. Be extremely careful with the -r option. As beginners it can be safer to navigate to the directory and remove content directly.

Let's take a quick second to apply some of the things we have learned and create more content to work with.

Navigate to the directory we just made (Module_1), and make another directory within it called directory to delete.

```
cd Module_1 #change directory
pwd #print working directory (wd)
mkdir directory_to_delete #make a new directory
ls #list the contents of wd
```

Now let's move to directory_to_delete and create a file, myseq.txt.

```
cd directory_to_delete #change directory
touch myseq.txt #Create a new file
```

Let's check the contents of our directory and see where we are located in our directory tree.

```
ls
pwd
```

To summarize what we have done:

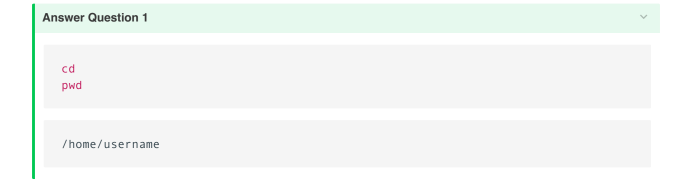
We've moved to the Module_1 directory, checked our directory with pwd, created a directory called directory_to_delete, and listed the contents of Module_1, so we can see the directory we just created. We then navigated to directory_to_delete using cd, created a file with touch, listed the contents with ls, and printed our working directory (pwd).

Let's move up one directory back to Module 1.

```
cd ..
```

Test Your Knowledge: Question 1

How could you move back to your home directory?



Test Your Knowledge: Question 2

If you changed to your home directory, how can you return to directory_to_delete?

```
Answer Question 2

We need to give the "path" to that directory.

cd /data/$USER/Module_1/directory_to_delete
pwd
ls
```

Getting back to removing directories (rmdir)

Now that we have created some directories, let's use rmdir to remove one.

What do you see when you try to remove this directory?

```
rmdir directory_to_delete
```

What should we do? We need to remove the contents of a directory before we can remove the directory. Here's one safe option.

```
cd directory_to_delete
ls
rm myseq.txt
ls
cd ..
ls
rmdir directory_to_delete
```

Moving and renaming files and directories, all with one command (mv)

The mv command is a handy way to rename files if you've created them with a typo or decide to use a more descriptive name. For example:

```
cd ..
mv file2.txt README.txt
ls
```

We can also move this file into a different location using the same command.

```
mv README.txt Module_1
cd Module_1
ls
```

Be careful when moving files, a mistake in the command can yield unexpected results. The - i interactive option will help keep you safe.

For example:

```
mkdir dir1
touch dir1/hello.txt
touch hello.txt
mv -i dir1/hello.txt hello.txt
```

Copying files (cp)

This is similar to mv but will create an actual copy of a file. You will need to specify what you are copying (the source) and where you want to make the copy (the target).

For example, let's copy a file from the BTEP teaching materials to Module_1.

```
cp /data/classes/BTEP/B4B_2025/Module_1/sample.fast* .
```

Remember, the . is a relative path shortcut denoting our current directory, so we are copying this into our current working directory.

* is a wildcard, matching zero or more characters including spaces. We are using this to copy two files that differ in the last letter of the file extension. More on these in Lesson 3.

We can also copy an entire directory using the recursive flag (cp -r). For example, let's copy the directory Practice_Sessions from the BTEP teaching materials to our current directory.

cp -r /data/classes/BTEP/B4B_2025/Module_1/Practice_Sessions .

Viewing file content

There are several ways to view files. We can use the less command to view the contents of a file like this.

```
less sample.fasta
```

You'll need to type q to get out of less and back to the command line. Before the less command was available, the more command was commonly used to look at file content. The less command has more options for scrolling through files, so it is now the preferred command.

Another command for reading files is cat, but this will print the contents in their entirety.

Help! (man)

Lastly, all Unix commands have a man or "manual" page that describes how to use them. If you need help remembering how to use the command 1s, you would type:

```
man ls
```

To exit man, again use q.

There are quite a few flags/options that we can use with the 1s command, and we can learn all about them on the man page. My favorite flags for 1s are -1 and -h. We will use flags often, and you won't get far in Unix without knowing about them. Try this:

```
cd
ls -lh
```

We have already seen these flags, but as a reminder...

- -h when used with the -1 option, use unit suffixes (Byte, Kilobyte, Megabyte, Gigabyte, Terabyte and Petabyte) in order to reduce the number of digits to three or less using base 2 for sizes.
- -I (The lowercase letter "ell".) List in long format. (See below). If the output is to a terminal, a total sum for all the file sizes is output on a line before the long listing.

Additional Resources

Software Carpentry: The Unix Shell (https://swcarpentry.github.io/shell-novice/01-intro/index.html)

Help Session

Practice navigating the file system and creating files. Instructions are here.

Lesson 2 Review

- Directories are folders that can store files, other directories, links, executables, etc.
- The file system is hierarchical with the root directory (/) at the top.
- Commands useful for navigating our file system and handling files include:
 - pwd = print working directory
 - ∘ 1s = list contents
 - cd = change directory
 - o mkdir, rmdir = make directory; remove directory
 - \circ rm = remove file
 - touch = create file
 - nano opens text editor
- absolute file paths include the entire location from the root of the file system
- relative file paths include the location from the working directory

Learning Objectives

In this lesson, we will continue to learn tips and concepts that facilitate working on the command line, including

- Flags and command options making programs do what they do
- Wildcards (e.g., *)
- Tab complete for less typing
- Accessing user history with the "up" and "down" arrows on the keyboard
- cat, head, and tail for reading files
- Working with file content (input, output, and append)
- Combining commands with the pipe (1).
- Finding information in files with grep
- Performing repetitive actions with Unix (e.g., for loop)
- File Permissions

For this lesson, you will need to connect to Biowulf.

Reminder: How to Connect to Biowulf

For this lesson and the lessons that follow, we will use NIH HPC student accounts (https://hpc.nih.gov/nih/student.html) to connect to Biowulf.

Open a terminal and type the following:

```
ssh username@biowulf.nih.gov
```

username = NIH/Biowulf login username. Remember to use the student account username here.

Type in your password at the prompt. The cursor will not move as you type your password!

Congrats! You are now connected to the login node. Your current working directory will be your home directory.

Getting Started

To begin this lesson, let's move to our data directory /data/\$USER.

```
cd /data/$USER
```

Let's also grab some files that we will need for this lesson from BTEP Teaching Materials.

```
cp -R /data/classes/BTEP/B4B_2025/Module_1 .
```

Did you forget what cp does? See Lesson 2.

Navigate to your Module 1 directory.

```
cd ./Module_1
```

Flags and command options - making programs do what they do

In Lesson 2, I introduced the idea of command options (flags). Command options allow us to change the behavior of a command. Command options, which ultimately allow you to modify command parameters, are extremely important to obtain expected results.

Let's return to 1s for an example. Compare the output from these commands:

```
ls
ls -S
ls -lh
```

1s -h (when used with -1 option, prints file sizes in a human readable format with the unit suffixes: Byte, Kilobyte, Megabyte, Gigabyte, Terabyte. This reduces the number of digits displayed.)

- 1s -1 (list in long format). The column output is as follows:
 - 1. directory / file type
 - 2. Content permissions
 - 3. Number of hard links to content
 - 4. Owner
 - 5. Group owner
 - 6. Content size (bytes)
 - 7. Last modified date / time
 - 8. File / directory name
- 1s S (sort from largest to smallest file size)

What do you see when combining the -h and -1 flags?

There are many flags you can use with 1s. How would we find out what they are?

```
man ls
```

Or to see a more user friendly display, google to the rescue. Google "man Is unix" and see what you get. Here's (https://shapeshed.com/unix-Is/) a useful, readable explanation of the "Is" command with examples.

Try combining some of the 1s flags.

```
ls -lhS
ls -alt
```

- -a (show hidden dot files .)
- t (sort by modification time with most recent listed first)

Flags and options add a layer of complexity to unix commands but are necessary to get the command or program to behave in the way you expect. For example, here is a command line for running "blastn" an NCBI/BLAST application. The Basic Local Alignment Search Tool (BLAST) finds regions of local similarity between sequences. (https://hpc.nih.gov/apps/Blast.html)

```
module load blast # Try this out by first loading the blast module
blastn -db /fdb/blastdb/nt -query seq1.fasta -out results.out
```

What's going on in this command line? First, the BLAST algorithm is specified, in this case it is blastn, then the -db flag is used to choose the database to search against (nt for nucleotide). The query flag specifies the input sequence, which is in FASTA format, and the -out flag specifies the name of the output file. See more about using Blast on NIH Biowulf here (https://hpc.nih.gov/apps/Blast.html).

Note

We are not running this today. This is just an example. If you want to run this, make sure you are not on the login node (Use sinteractive) and be sure to allocate enough memory. See the Biowulf Blast help docs (https://hpc.nih.gov/apps/Blast.html).

Use of wildcards

Wildcard characters are a handy tool when working at the command line. Wildcards "are symbols or special characters that represent other characters." (https://www.tecmint.com/use-wildcards-to-match-filenames-in-linux/) Want to list all your FASTA files?

Use *:

```
ls *.fasta
```

The * wildcard matches zero or more characters including spaces.

This example will work as long as all your FASTA files end in .fasta. But sometimes they don't (e.g., fas or fa). Read more about FASTA file extensions here (https://en.wikipedia.org/wiki/FASTA_format#FASTA_file).

For example, to find a file ending in . fa, you could use the following:

```
touch file.fa #First create a file ending in fa
ls *.fa
```

or you want all the the FASTA and FASTQ files.

```
ls *.fa*
```

In addition to the asterisk (*) character, you can use other wildcards on the unix command line, not limited to the following:

? - matches a single character

{} - used for multiple matches

[] - specify a range of characters or numbers. For example, [a-z] would specify all lower case letters and [0-9] would mean all digits.

To see some more practical examples of using wildcards, see this article (https://www.tecmint.com/use-wildcards-to-match-filenames-in-linux/) from tecmint.com and this (https://medium.com/@leedowthwaite/advanced-wildcard-patterns-most-people-dont-know-52f7fd608cb3) from the medium.com. This second article provides a nice discussion on how wildcards differ from regular expressions.

Using tab complete for less typing

Here's a good Unix trick to know - tab complete. Start typing the name of the file or directory you want, and hit the tab key. The system will auto-complete the name of the file or directory if the name is unique. It may only give a partial name if there is more than one file with a similar name in the directory. You can fill in the next part of the name and then try tab-complete again.

Let's create some files to test this.

```
touch file.txt
touch file.fasta
touch file.fastq
```

Start typing...

```
less f (hit tab)
```

What do you get? How does that differ from:

```
less file (hit tab)
```

The tab complete will save you lots of typing, and also help to figure out if you are where you think you are in the directory structure.

Access your history with the "up" and "down" arrows on your keyboard

Here's another Unix trick to make your life easier. Access previous commands with the up and down arrows on your keyboard. You can scroll backwards and forwards. This helps when you've got a typo or small mistake in your command lines that you can fix without retyping the whole thing.

The history command

You can also search, view, and retrieve recently using commands using history. See this guide (https://www.geeksforgeeks.org/history-command-in-linux-with-examples/).

Keyboard shortcuts

There are also a few handy keyboard shortcuts to make life on the command line easier. For example:

ctrl c to kill a running process

ctrl 1 to clear the screen

ctrl a skip to the beginning of a command

ctrl e skip to the end of a line

See more examples here (https://unix.stackexchange.com/questions/255707/what-are-the-keyboard-shortcuts-for-the-command-line).

cat, head, and tail

Who says Unix programmers don't have a sense of humor? Let me introduce cat, head, and tail. The cat command (short for "concatenate") is an extremely useful command for creating new files and viewing file content. You can use it to open files for reading input and writing output. Or you can use it to copy several files into a new file. Also you can "append" the contents of a file to the end of another file.

This command reads the content of sample.fasta and outputs to standard output (i.e., the screen). This is not helpful for very large files, as it moves to the end of the file quickly. Less is a better command option for reading large files.

```
cat sample.fasta
```

You can use cat to combine several files into one file, such as:

```
cat seq1.fasta seq2.fasta
```

Although, this again prints to standard output, the screen. To capture that output, we need to learn how to redirect output. (Coming up next!)

In the meantime, let's take a quick look at head and tail.

head - prints the first 10 lines of a specified file (by default)

```
head sample.fasta
```

tail - prints the last 10 lines of a specified file (by default)

```
tail sample.fasta
```

You can specify how many lines you would like to see (-n), or you can use the default value, which is 10.

```
head -n 20 sample.fasta
```

Want to be sure of those 20 lines? Let's use cat -n to show the file with numbered lines and compare.

```
cat -n sample.fasta
```

What if you didn't know what -n did? How could you find out more about "cat"?

```
man cat
```

Working with file content (input <, output >, and append >>)

By default, commands take input from the standard input (your keyboard) and send the results to standard output (your screen). If you want to redirect the output (results) of a command to file, you need to use output redirection. Similarly, we can also redirect input, for example, instead of coming from our keyboard, it can instead come from file. This is known as input redirection. Learn more here. (https://www.geeksforgeeks.org/input-output-redirection-in-linux/)

- < input redirection operator
- > output redirection operator
- >> append redirection operator

Want to put the output from cat, head, or tail into a new file?

```
head -n 20 seq1.fasta > smaller.fasta
```

Or we could put the last 20 lines into a file with tail.

```
tail -n 20 seq1.fasta > smaller2.fasta
```

What if we want the first 20 lines and the last 20 lines in one file, with the first at the top and the last at the bottom? Use append, >> to paste the second file to the bottom of the first file. Let's try it.

```
head -n 20 sample.fasta > smaller.fasta
tail -n 20 sample.fasta >> smaller.fasta
```

Keep in mind that if you input into the same file multiple times, you are overwriting the previous contents. For example, what is the final content of our file covid.fasta?

```
head -n 20 seq1.fasta > covid.fasta
head -n 20 seq2.fasta > covid.fasta
head -n 20 seq3.fasta > covid.fasta
```

How many lines are now in covid.fasta? How can you check?

Let's try wc, short for word count.

```
wc covid.fasta
```

wc is a very useful function. Without opening a file, we can find out how many lines, words and characters are in it. Line counts are extremely useful to assess your data output.

If we created a file where we were expecting there to be 1000 lines of output? The wc command provides a quick way to check.

What happened to all of our content? The final results are from "seq3.fasta" only. The other two results files have been overwritten.

So, how would you get all three files into covid.fasta? You'll need to use append.

```
cat seq1.fasta > covid.fasta
cat seq2.fasta >> covid.fasta
cat seq3.fasta >> covid.fasta
```

OR

```
cat seq1.fasta seq2.fasta seq3.fasta > covid2.fasta
```

How could you test to see if the file has the expected number of lines?

```
wc covid.fasta
wc covid2.fasta
```

To redirect input, we would use something like this:

```
less < covid.fasta
```

Yes, you could use less covid.fasta. However, these commands act differently. In less covid.fasta, the file is directly passed to the less. However, with less < covid.fasta the content of covid.fasta is passed to less.

Combining commands with pipe (|). Where the heck is pipe anyway?

The hardest thing about using pipe (|) is finding it on your keyboard!

The pipe symbol "|" (a.k.a., vertical bar) is way over on the right hand side of your keyboard, above the backslash \.

Pipe is used to take the output from one command, and use it as input for the next command.

For example,

```
head -n <mark>20</mark> sample.fasta | wc
```

Using what we've learned, all together now.

Let's say you've got a very large FASTA or FASTQ file, and you want to run an analysis on it. Before working on the whole file, it can be useful to set up a smaller test file instead.

Here's one way to do it.

```
cat sample.fasta | head -n 20 > output.fasta
```

This combines several things we have learned about. The cat command opens the file sample.fasta for writing. The pipe | command is used to take that output and run it through the head command where we only want to see the first 20 lines, and we want them output > into a file called "output.fasta".

Note

We could simply use head without cat, but we wanted to use the pipe for an example.

Let's compare the files. How are they different?

```
ls -lh
```

and

```
less sample.fasta
less output.fasta
```

Finding information in files with grep

The grep utility is used to search files looking for a pattern match. It is used like this.

```
grep pattern options filename
```

As our first example we will look for restriction enzyme (EcoRI) sites in a sequence file (eco.fasta). The file has four EcoRI sites, but two of them are across the end of the line (and won't be found).

```
cat eco.fasta
grep -n GAATTC eco.fasta
```

We can modify the eco.fasta file to remove the line breaks (\n) at the ends of the lines using tr.

```
grep -v ">" eco.fasta | tr -d "\n" | grep GAATTC
```

-v: This prints out all the lines that do not match the pattern, effectively removing the header line (>) of the file.

The unix tr (translate) command is used for translating or deleting characters.

Usage:

```
tr [option] set1 [set2]
```

-d: Deletes characters in the first set from the output

So this part of the command line is finding the line breaks \n and removing them.

And now we can see all four of the EcoRI sites.

What if we just wanted to count the occurrence of the EcoRI sites in the sequence?

```
grep -v ">" eco.fasta | tr -d "\n" | grep GAATTC -c
```

Perhaps "1" is not what you expected. Check out man grep.

The -c option prints only a count of the lines that match a pattern. Here we are counting the entire line as one.

If we wanted to see each of the EcoRI sites listed.

```
grep -v ">" eco.fasta | tr -d "\n" | grep GAATTC -o
```

And if we want to count the number of EcoRI sites.

```
grep -v ">" eco.fasta | tr -d "\n" | grep GAATTC -o | wc
```

Let's create a word file that we can input to grep. We can input multiple restriction enzyme sites and search for all of them.

```
nano wordfile.txt
```

Put in the words (GAATTC, TTTTT). Now we can use that file to find lines.

```
grep -v ">" eco.fasta | tr -d "\n" | grep -f wordfile.txt -o | wc
```

We can find a list of options to be used with grep using man:

```
man grep
```

Performing repetitive actions with Unix

We can create a "for" loop to do iterative actions in Unix.

For each commands all on one line or separate lines: (i can be any variable name). These steps can be saved as a file, thereby creating a simple Unix script.

What does this "for loop" do?

```
for i in *.fasta; do ls $i; done
```

What do these command lines do?

```
for i in *.fasta; do echo $i; grep ">" $i; done
```

This one pulls out all the header ">" lines in the fasta files.

```
for i in seq*.fasta; do echo $i; grep ">" $i; done
```

While this one pulls out the header lines from files named seq*.fasta.

Learn more about for loops here (https://www.geeksforgeeks.org/bash-scripting-for-loop/).

Permissions - when all else fails check the permissions

Permissions dictate who can access your files and directories, and what actions they can perform. If you are consistently getting error messages from your command line and you're sure you are typing it correctly, it's worthwhile checking the permissions. So what does it all mean? How do we read the permissions information?

As we have seen, 1s -1, provides information about file types, the owner of the file, and other permissions.

For example:

```
ls -l sample.fasta
```

Here is an overview of what these permissions mean:

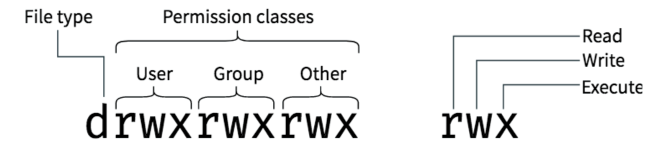


Image from booleanworld.com (https://www.booleanworld.com/introduction-linux-file-permissions/)

The first letter d indicates whether this is a directory or not - or some other special file type. The next 3 positions are the owner's/user's permissions. In this image, the owner can "read", "write" and "execute". So they can create files and directories here, read files here, and execute/run programs. The next 3 positions show the permissions for the "group". The last 3 positions shows permissions for everyone ("other").

You can modify permissions using chmod. Let's see this in action.

```
touch example.txt
chmod u-w example.txt
ls -l example.txt
```

Now, reassign write privilege to the user/owner

```
chmod u+w example.txt
ls -l example.txt
```

Help Session

Let's complete a Unix treasure hunt.

Lesson 4: Working on Biowulf

Lesson 3 Review

- Flags and command options
- Wildcards (*)
- Tab complete
- Accessing user history with the "up" and "down" arrows
- cat, head, and tail
- Working with file content (input, output, and append)
- Combining commands with the pipe (|)
- grep
- for loop
- File Permissions

Lesson Objectives

- Learn about the slurm system by working on Biowulf. Learn about batch jobs, swarms jobs, interactive sessions.
- Retrieve data from NCBI through a batch job.
- Learn how to troubleshoot failed jobs.

For this lesson, you will need to connect to Biowulf.

Working on Biowulf

Now that we are becoming more proficient at the command line, we can use these skills to do more than navigate our file system. We can actually begin working on Biowulf. Today's lesson will focus on submitting computational jobs on the Biowulf compute nodes.

Login using ssh

To get started, as always, we will need to log in to Biowulf. Make sure you are on VPN.

Open your Terminal if you are using a mac or the Command prompt if you are using a Windows machine.

ssh username@biowulf.nih.gov

username = NIH/Biowulf login username. Remember to use the student account username here.

Type in your password at the prompt. The cursor will not move as you type your password!

When you log in to Biowulf, you are automatically in your home directory (/home/\$USER). This directory is very small and not suitable for large data files or analysis.

Use the cd command to change to your data directory (/data).

```
cd /data/$USER
```

where \$USER is an environment variable holding your username.

If you do not yet have one, create a directory to work in for this lesson and move to that directory.

```
mkdir Module_1
cd Module_1
```

Note

When working on Biowulf, you can not use any computational tools on the "login node". Instead, you need to work on a node or nodes that are sufficient for what you are doing.

To run jobs on Biowulf, you must designate them as interactive, batch or swarm. Failure to do this may result in a temporary account lockout.

What kind of work can we do on the login node?

The login node can be used for:

- Submitting resource intensive tasks as jobs
- Editing and compiling code
- File management and data transfers on a small scale

Batch Jobs

Most jobs on Biowulf should be run as batch jobs using the "sbatch" command.

```
sbatch yourscript.sh
```

Where yourscript.sh is a shell script containing the job commands including input, output, cpus-per-task, and other steps. Batch scripts always start with #!/bin/bash or similar call. The sha-bang (#!) tells the computer what command interpreter to use, in this case the Bourneagain shell.

For example, to submit a job checking sequence quality using fastqc (MORE ON THIS LATER), you may create a script named fastqc.sh:

```
nano fastqc.sh
```

Inside the script, you may include something like this:

```
#!/bin/bash

module load fastqc
fastqc -o output_dir -f fastq seqfile1 seqfile2 ... seqfileN
```

where -o names the output directory

- f states the format of the input file(s)

and seqfile1 ... seqfileN are the names of the sequence files.

Note

fastqc is a available via Biowulf's module system, and so prior to running the command, the module had to be loaded.

For more information on running batch jobs on Biowulf, please see: https://hpc.nih.gov/docs/userguide.html (https://hpc.nih.gov/docs/userguide.html).

Multi-threaded jobs and sbatch options

In high-performance computing, multithreading lets a single program split itself into multiple "workers" that can run at the same time on different parts of the computer's brain (CPUs), speeding up complex tasks. Many bioinformatics programs use multi-threading.

For multi-threaded jobs, you will need to set --cpus-per-task. You can do this at the command line or from within your script.

Example at the command line:

```
sbatch --cpus-per-task=# yourscript.sh
```

In your script:

```
#!/bin/bash
#SBATCH --job-name qc
#SBATCH --mail-type BEGIN,END
#SBATCH --cpus-per-task #

module load fastqc
fastqc -o output_dir -t $SLURM_CPUS_PER_TASK -f fastq seqfile1 seqfi.
```

Within the script we can use **directives** denoted by **#SBATCH** to support command line arguments such as --cpus-per-task. If included within the script, you will not need to call these at the command line when submitting the job. You should also pass the environment variable, **\$SLURM_CPUS_PER_TASK** to the thread argument. Some other useful directives include --job-name, where you assign a name to the submitted job, and --mail-type, which you can use to direct **slurm** to send you an email when a job begins, ends, or both.

Tip

The jobscript should always be the last argument of sbatch.

To see more sbatch options, use

```
sbatch --help
```

Some slurm commands

Once you submit a job, you will need to interact with the slurm system to manage or view details about submitted jobs.

Here are some useful commands for this purpose:

Command	Purpose	Pros & Cons
squeue	Information about jobs	Lots of options, odd syntax
scancel	Delete job(s)	Only way to cancel your jobs!
sacct	Job accounting for completed jobs	Useful to get a list of jobs you submitted recently (default is 'today'), odd syntax
sjobs (Biowulf utility)	Info about running and pending jobs	Informative but slow

Courtesy of NIH HPC Team Training Documentation

Standard error and standard output

Output that you would expect to appear in the terminal (e.g., standard error and standard output) will not in batch mode. Rather, these will be written by default to slurm#####.out in the submitting directory, where ##### represents the job id. These can be redirected using --output=/path/to/dir/filename and --error=/path/to/dir/filename on the command line or as an #SBATCH directive.

Partitions

Your job may be in a waiting phase ("Pending" or "PD") depending on available resources. You can specify a particular node partition using --partition.

Use freen to see what's available.

Summary of partitions

Nodes available to all users				
norm	the default partition. Restricted to single-node jobs			
multinode	Intended to be used for large-scale parallel jobs. Single node jobs are not allowed. See here for detailed information.			
largemem	Large memory nodes. Reserved for jobs with memory requirements that cannot fit on the norm partition. Jobs in the largemem partition must request a memory allocation of at least 350GB.			
unlimited	Reserved for jobs that require more than the default 10-day walltime. Note that this is a small partition with a low CPUs-per-user limit. Only jobs that absolutely require more than 10 days runtime, that cannot be split into shorter subjobs, or that are a first-time run where the walltime is unknown, should be run on this partition.			
quick	For jobs < 4 hours long. These jobs are scheduled at higher priority. They may run on the dedicated quick partition nodes, or on the buy-in nodes when they are free.			
gpu	GPU nodes reserved for applications that are built for GPUs.			
visual	Small number of GPU nodes reserved for jobs that require hardware accelerated graphics for data visualization.			
	Buy-in nodes			
ccr*	for NCI CCR users			
forgo	for individual groups from NHLBI and NINDS			
persist	for NIMH users			

Student partition

The student accounts have their own partition for running jobs, --partition=student. This will need to be included on the command line or in the job script.

Walltime

The default walltime, or amount of time allocated to a job, is 2 hours on the norm partition. To change the walltime, use --time=d-hh:mm:ss.

Here are the walltimes by partition:

batchlim

	Walltime		
Partition	Default	Maximum	
norm	 02:00:00	10-00:00:00	
multinode	08:00:00	10-00:00:00	
+turbo		08:00:00	
interactive	08:00:00	1-12:00:00	
gpu	02:00:00	10-00:00:00	
unlimited	UNLIMITED	UNLIMITED	
largemem	02:00:00	10-00:00:00	
visual	08:00:00	1-12:00:00	
quick	02:00:00	04:00:00	
persist (NIMH)	UNLIMITED	UNLIMITED	
ccr (NCI_CCR)	04:00:00	10-00:00:00	
forgo (lab only)	1-00:00:00 	3-00:00:00	

You can change the walltime after submitting a job using

```
newwall --jobid <job_id> --time <new_time_spec>
```

Submit an actual job

Let's submit a batch job. We are going to download data from the Sequence Read Archive (SRA), a public repository of high throughput, short read sequencing data. We will discuss the SRA a bit more in detail in Lesson 5. For now, our goal is to simply download multiple fastq files associated with a specific BioProject on the SRA. We are interested in downloading RNAseq files associated with BioProject PRJNA578488, which "aimed to determine the genetic and molecular factors that dictate resistance to WNT-targeted therapy in intestinal tumor cells".

We will learn how to pull the files we are interested in directly from SRA at a later date. For now, we will use the run information stored in sra_files_PRJNA578488.txt.

Let's copy this file to our working directory and view using less.

```
cp /data/classes/BTEP/B4B_2025/Module_1/sra_files_PRJNA578488.txt .
less sra_files_PRJNA578488.txt
```

Now, let's build a script downloading a single run, SRR10314042, to a directory called /data/ \$USER/testscript.

```
mkdir testscript

cd testscript
```

Open the text editor nano and create a script named filedownload.sh.

```
nano filedownload.sh
```

Inside our script we will type

```
#!/bin/bash
#SBATCH --cpus-per-task=6
#SBATCH --gres=lscratch:10
#SBATCH --partition=student

#load module
module load sratoolkit

fasterq-dump -t /lscratch/$SLURM_JOB_ID SRR10314042
```

Notice our use of sbatch directives inside the script.

fasterq-dump assigns 6 threads by default, so we are specifying --cpus-per-task=6. This can be modified once we get an idea of the CPUs needed for the job. -t assigns the location to be used for temporary files. Here, we are using /lscratch/\$SLURM_JOB_ID, which is created upon job submission. In combination, we need to request local scratch space allocation, which we are setting to 10 GB (--gres=lscratch:10). The final directive, --partition=student is required because we are using student accounts for this lesson. If you are with the Center for Cancer Research and using your own account, you can use --partition=ccr.

Remember:

Default compute allocation = 1 physical core = 2 CPUs

Default Memory Per CPU = 2 GB. Therefore, default memory allocation = 4 GB

Now, let's run the script.

sbatch filedownload.sh

Let's check our job status.

```
squeue -u $USER
```

Once the job status changes from PD (pending) to R (running), let's check the job status.

```
sjobs -u $USER
```

Some other useful job monitoring commands include jobload, and jobhist. The latter is useful for when the job completes.

When the job completes, we should have a file called SRR10314042.fastq.

```
ls -lth
```

Now, what if we want to download all of the runs from sra_files_PRJNA578488.txt. We could use a for loop, GNU parallel, which acts similarly to a for loop (MORE ON THIS NEXT LESSON), or we can submit multiple fasterq-dump jobs in a job array, one subjob per run accession.

Note

There are instructions for running SRA-Toolkit on Biowulf here (https://hpc.nih.gov/apps/sratoolkit.html).

Swarm-ing on Biowulf

Swarm is for running a group of commands (job array) on Biowulf. swarm reads a list of command lines and automatically submits them to the system as sub jobs. "By default, swarm runs one command per core on a node, making optimum use of a node. Thus, a node with 16 cores will run 16 commands in parallel." (https://hpc.nih.gov/apps/swarm.html). To create a swarm file, you can use nano or another text editor and put all of your command lines in a file called file.swarm (file is just a placeholder). Then you will use the swarm command to execute.

For example,

```
$ swarm -f file.swarm
```

Note

By default, each subjob is allocated 1.5 gb of memory and 1 core (consisting of 2 cpus) --- hpc.nih.gov (https://hpc.nih.gov/apps/swarm.html)

Swarm creates two output files for each command line, one each for STDOUT (file.o) and STDERR (file.e). You can look into these files with the less command to see any important messages.

For example,

```
$ less swarm_jobid_subjobid.o
$ less swarm_jobid_subjobid.e
```

View the swarm options using

```
swarm --help
```

For more information on swarm-ing on Biowulf, please see: https://hpc.nih.gov/apps/swarm.html (https://hpc.nih.gov/apps/swarm.html)

Let's create a swarm job

To retrieve all the files at once, you can create a swarm job.

```
nano set.swarm
```

Copy the following and paste into the swarm file.

```
#SWARM --threads-per-process 6
#SWARM --gb-per-process 4
#SWARM --gres=lscratch:20
#SWARM --module sratoolkit
#SWARM --partition=student

fasterq-dump -v -t /lscratch/$SLURM_JOB_ID SR10314043 #Add error to state to the st
```

Here, based on our swarm directives, each subjob will request 6 cpus and 4 GB of RAM. We are also denoting local scratch space of 20 GB per each command.

There is advice for generating a swarm file using a for loop and echo in the swarm user guide (https://hpc.nih.gov/apps/swarm.html). If you can help it, do not type each of these commands.

Something like this could generate the lines above:

```
cat ../sra_files_PRJNA578488.txt | while read line; do
    echo 'fasterq-dump -v -t /lscratch/$SLURM_JOB_ID' $line >> script
done
```

You may have noticed that the while loop above does not produce the same lines. The order of the files is in reverse, and there is an extra line of code. fasterq-dump -v -t /lscratch/\$SLURM_JOB_ID SRR10314045 fasterq-dump -v -t /lscratch/\$SLURM_JOB_ID SRR10314044 fasterq-dump -v -t /lscratch/\$SLURM_JOB_ID SRR10314043 fasterq-dump -v -t /lscratch/\$SLURM_JOB_ID How can we fix the lines written to script.sh? Use the skills you have learned and a little help from google. Possible Solution One solution is to sort the script using sort and drop the first line of the file using tail -n +2. sort script.sh | tail -n +2 > script_mod.sh

Let's run our swarm file. Because we included our directives within the swarm file, the only option we need to include is - f for file.

```
swarm -f set.swarm
```

Running Interactive Jobs

Interactive nodes are suitable for testing/debugging cpu-intensive code, pre/post-processing of data, graphical application, and to GUI interface to application.

To start an interactive node, type sinteractive at the command line and press Enter/Return on your keyboard.

```
sinteractive
```

You will see something like this printed to your screen. You only need to use the sinteractive command once per session. If you try to start an interactive node on top of another interactive node, you will get a message asking why you want to start another node.

```
[username@biowulf ]$ sinteractive
salloc.exe: Pending job allocation 34516111
salloc.exe: job 34516111 queued and waiting for resources
salloc.exe: job 34516111 has been allocated resources
salloc.exe: Granted job allocation 34516111
salloc.exe: Waiting for resource configuration
salloc.exe: Nodes cn3317 are ready for job
srun: error: x11: no local DISPLAY defined, skipping
[username@cn3317 ]$
```

You can use many of the same options for sinteractive as you can with sbatch. (https://hpc.nih.gov/docs/userguide.html#int) The default sinteractive allocation is 1 core (2 CPUs) and 3 GB (1.5 GB per CPU) of memory and a walltime of 8 hours.

For example,

```
sinteractive --gres=lscratch:20 --cpus-per-task=6
module load sratoolkit
fasterq-dump -t /lscratch/$SLURM_JOBID SRR2048331 -0 /data/$USER/sra
```

To terminate / cancel the interactive session use

```
exit
```

Exiting from Biowulf

To disconnect the remote connection on Biowulf, use

```
exit
```

Transferring files to Biowulf

Before ending this lesson, I want to briefly discuss file transfers. At some point you will need to know how to get your data on Biowulf. The NIH HPC Team has fantastic documentation on this subject at https://hpc.nih.gov/docs/transfer.html (https://hpc.nih.gov/docs/transfer.html).

Remember, Helix (helix.nih.gov) is the interactive data transfer and file management node for the NIH HPC Systems. If you are interactively transferring files to and from NIH HPC directories, you should connect to Helix:

```
ssh username@helix.nih.gov
```

where username is your Biowulf username.

Recommended methods for file transfers

Large scale transfers

For large scale transfers and large files use, Globus. Find instructions here. (https://hpc.nih.gov/docs/globus/)

Small scale transfers

For small scale transfers, use scp to or from Helix or drag and drop files by mounting HPC Systems directories to your local machine.

For example, to copy Module 1 to your local machine, use:

```
scp -r username@helix.nih.gov:/data/username/Module_1 .
```

Note

This command and the next are issued from your local computer. The . in the above command means that you are copying Module_1 into your current working directory.

To copy from your local machine to Biowulf, use:

```
scp -r ./Module_1 username@helix.nih.gov:/data/username/Module_1
```

Make sure to substitute username with your Biowulf username.

To mount the HPC directories to your local computer, follow the instructions here. (https://hpc.nih.gov/docs/hpcdrive.html)

Help Session

Practice the skills learned in this lesson here.

So you think you know Biowulf?

Quiz yourself using the hpc.nih.gov biowulf-quiz (https://hpc.nih.gov/training/intro_biowulf/biowulf-quiz/).

Lesson 5: Downloading data from the SRA

Lesson 4 Review:

- The majority of computational tasks on Biowulf should be submitted as jobs: sbatch or swarm
- the SRA-toolkit can be used to retrieve data from the Sequence Read Archive

Learning Objectives:

- 1. Download data from the SRA with fastq-dump
 - split files into forward and reverse reads
 - o download part, not all, of the data
- 2. Compare fastq-dump to fasterq-dump
- 3. Introduce prefetch
- 4. Grab SRA run info and run accession information
- 5. Learn to automate data retrieval with the parallel command
- 6. Learn about alternative download options

Get Started

Connect to Helix

Helix is the dedicated data management and file transfer node.

ssh username@helix.nih.gov

username = NIH/Biowulf login username. Remember to use the student account username here.

Type in your password at the prompt. The cursor will not move as you type your password!

Congrats! You are now connected to Helix. Your current working directory will be your home directory.

Navigate to your Module_1 directory

```
cd /data/$USER/Module_1
```

If you do not have a Module_1 directory, use the following:

```
cp -R /data/classes/BTEP/B4B_2025/Module_1 /data/$USER
cd /data/$USER/Module_1
```

Create a lesson directory

Here, we will download data from the NCBI/SRA. Let's create a directory named sra_data to store the data we are using today.

```
mkdir sra_data
cd sra_data
```

A brief word about sequence formats

 FASTA – commonly used text format for downstream analysis such as sequence similarity searches

```
head ../sample.fasta
```

• FASTQ – output format from NGS technologies with quality scores

```
head ../sample.fastq
```

@SRR1553607.1 1 length=202

GTTAGCGTTGTTGATCGCGACGCAACAACTGGTAAAGAATCTGGAAGAAGGATATCAGTTCAAACGCT(
AGCGAGATGATGGATATTTTTGAACGACTCATCATCGTCACATCCCGTTCAACCTGATTGAGTCGTT(
AAAATATCCATCATCTCGCTTGAGCGTTTGAACTGATATCCTTCTTCCAGATTCTTTACCAG
+SRR1553607.1 1 length=202

@SRR1553607.2 2 length=202

GGTGTAAGCACAGTACTCGGCCCACATCGCCTTTGTGTTAATGAAGTTTGGGTATCAACTTTCATCCC(
ATCTTCCGTGGAAGGAGTATGTTCCGTCAATCCGTACATGTTAACCCTGGGCCTGTTACTTGGGCCTT(
CGGGTTCTAATACAAATTGACGGAACATACTCCTTCCACGGAAGATTGGGGATGAAAGTTGA
+SRR1553607.2 2 length=202

What is the SRA?

The SRA (Sequence Read Archive) at NCBI is a large, public database of DNA sequencing data. The repository holds "short reads" generated by high-throughput next-generation sequencing, usually less than 1,000 bp.

We will download data from the SRA using the command line package sratoolkit. We will also explore how you can download these data from NCBI via a web browser.

Helpful documentation for the SRA:

- 1. SRA handbook at NCBI (https://www.ncbi.nlm.nih.gov/books/NBK47528/)
- 2. SRA Toolkit Documentation (NCBI) (https://github.com/ncbi/sra-tools/wiki)

SRA terms to know

BioProject (Prefix: PRJNA, example: PRJNA257197 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA257197/)): provides a description of the research project associated with the sequencing study. It includes information such as the study's abstract, links to publications (if available), and links to project data, including experiments, PubMed, and GEO datasets.

BioSample (Prefix: SAMN): contains sample-specific details. It could include details such as disease information, sex, response to drug, race, and other relevant attributes.

Sample Accession (Prefix: SRS): A Sample is an object imported from BioSample, which contains metadata describing the physical sample upon which a sequencing experiment was performed.

Experiment (Prefix: SRX): an object that contains metadata describing the library preparation, the sequencing technique, instrument used, and other experiment-specific details.

Run (Prefix: SRR): A Run is an object that contains the actual sequencing data for a particular sequencing experiment. Experiments may have multiple Runs. -- Decoding SRA Accessions: A Primer on Navigating NCBI's Sequence Read Archive (https://gauravsh.com/decoding_sra_accessions/

#:~:text=SRA%20Experiment%20Accession%20(SRX%23)%3A,physical%20sample%20imported%20

Using fastq-dump

fastq-dump and fasterq-dump are modules in the SRA Toolkit and can be used to download FASTQ-formatted data. Both download the data in SRA format and convert it to FASTQ format.

If working on a high performance computing system such as Biowulf, either submit a script to cluster, use an interactive session, or connect to helix. Make sure to do module load sratoolkit prior to issuing either the fastq-dump or fasterq-dump commands. Alternatively, install the SRA Tool kit on local computer (see https://github.com/ncbi/sra-tools/wiki/01.-Downloading-SRA-Toolkit (https://github.com/ncbi/sra-tools/wiki/01.-Downloading-SRA-Toolkit)).

module load sratoolkit

fastq-dump SRR1553607

creates the file:

SRR1553607.fastq

Check the file to make sure it is in fastq format. How would you do this? What is FASTQ format?

What are "spots"?

Spots are a legacy term referring to locations on the flow cell for Illumina sequencers. All of the bases for a single location constitute the spot including technical reads (e.g., adapters, primers, barcodes, etc.) and biological reads (forward, reverse). In general, you can think of a spot as you do a read. For more information on spots, see the linked discussion (https://

www.biostars.org/p/12047/) on Biostars. When downloading "spots", always split the spots into the original files using:

```
--split-files
```

For paired-end reads, we need to separate the data into two different files like this:

```
fastq-dump --split-files SRR1553607
```

which creates

```
SRR1553607_1.fastq
SRR1553607_2.fastq
```

Split 3

There is an additional option --split-3 that will split the reads into forward and reverse files and a third file with unmatched reads. Since many bioinformatic programs require matching paired end reads, **this is the preferred option. **

fastq-dump first downloads the data in SRA format, then converts it to FASTQ. If we want to work with a subset of the data, for example the first 10,000 reads, we can use -X:

```
fastq-dump --split-3 -X 10000 SRR1553607 --outdir SRR1553607_subset
```

We have additionally included --outdir to save our sub-setted data to a different directory, so that we do not overwrite our previous downloads.

To see other fastq-dump options, use

```
fastq-dump --help
```

Other options of interest:

- --gzip or bzip2 allows you to compress the reads
- --skip-technical allows you to only download biological reads

To generate an XML report on the data that shows us the "spot" or read count, and the count of bases "base_count", including the size of the data file:

```
sra-stat --xml --quick SRR1553607
```

which yields this data in XML format:

```
<Run accession="SRR1553607" spot_count="203445" base_count="41095890"</pre>
  <Member member name="A" spot count="203445" base count="41095890" |</pre>
 <Size value="25452196" units="bytes"/>
  <AlignInfo>
 </AlignInfo>
 <QualityCount>
    <Quality value="2" count="3523771"/>
   <Quality value="5" count="40006"/>
   <Quality value="6" count="30160"/>
   <Quality value="7" count="79961"/>
    <Quality value="8" count="80987"/>
   <Quality value="9" count="21904"/>
   <Quality value="10" count="63700"/>
   <Quality value="11" count="23695"/>
   <Quality value="12" count="42479"/>
   <Quality value="13" count="24910"/>
   <Quality value="14" count="22036"/>
   <Quality value="15" count="135747"/>
   <Quality value="16" count="56174"/>
   <Quality value="17" count="81514"/>
   <Quality value="18" count="123393"/>
   <Quality value="19" count="79256"/>
   <Quality value="20" count="269807"/>
   <Quality value="21" count="68295"/>
   <Quality value="22" count="125196"/>
   <Quality value="23" count="267733"/>
   <Quality value="24" count="287688"/>
   <Quality value="25" count="296721"/>
   <Quality value="26" count="338062"/>
   <Quality value="27" count="507890"/>
   <Quality value="28" count="226423"/>
   <Quality value="29" count="569612"/>
   <Quality value="30" count="813014"/>
   <Quality value="31" count="1309505"/>
   <Quality value="32" count="812951"/>
   <Quality value="33" count="2398417"/>
   <Quality value="34" count="2324453"/>
    <Quality value="35" count="10399039"/>
   <Quality value="36" count="1250313"/>
   <Quality value="37" count="2681177"/>
   <Quality value="38" count="1210457"/>
    <Quality value="39" count="2744683"/>
   <Quality value="40" count="2268539"/>
   <Quality value="41" count="5496222"/>
```

```
<pr
```

where we can see the spot count, the base count, and the number of bases corresponding to quality scores.

```
spot_count="203445" base_count="41095890"
```

Side bar: (XML) is Extensible Markup Language, a document format that is both human and machine-readable. XML aims for: "simplicity, generality and usability across the Internet".

```
Using seqkit stat

This tool cannot be used on Helix.

We can get similar information from our downloaded files using a program called seqkit (https://bioinf.shenwei.me/seqkit/). If you get a chance, grab an interactive session on Biowulf and check out seqkit.

module load seqkit

First, let's see what options are available?

seqkit --help

We can see that seqkit stats provides "simple statistics of FASTA/Q files". Let's try it out.

seqkit stat SRR1553607_1.fastq
```

fasterq-dump

We have already seen fasterq-dump in action (See Lesson 4). fasterq-dump is faster than fastq-dump because it uses multi-threading (default --threads 6). --split-3 and --

skip-technical are defaults with fasterq-dump, so you do not need to worry about specifying how you want the files to be split.

However, you can not grab a subset of the file like you can with fastq-dump nor can you compress the file during download, so fasterq-dump is not necessarily a replacement for fastq-dump.

Using prefetch

Both fastq-dump and fasterq-dump are faster when following prefetch (https://github.com/ncbi/sra-tools/wiki/08.-prefetch-and-fasterq-dump), and fasterq-dump paired with prefetch is the fastest way to pull the files from the SRA. There are alternative options, which we will mention later.

prefetch will first download all of the necessary files to your working directory. Runs are downloaded in the SRA format (compressed).

```
mkdir prefetch
cd prefetch
prefetch SRR1553607
ls -l
```

fasterq-dump will then convert the files to the fastq format.

```
fasterq-dump -p -t /scratch/$USER SRR1553607
```

What does -p do? How can we find out?

Note

We are not on a computational node, so we will use scratch instead of lscratch.

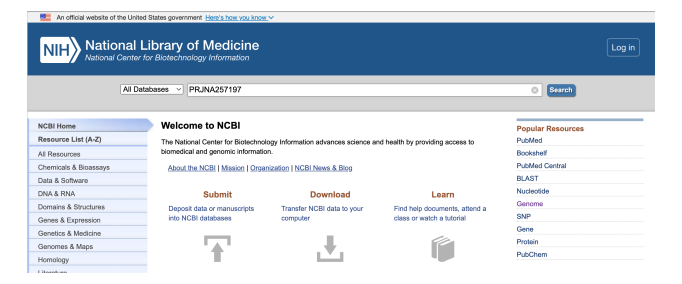
Navigating NCBI SRA

Where can we get an accession list or run information for a particular project?

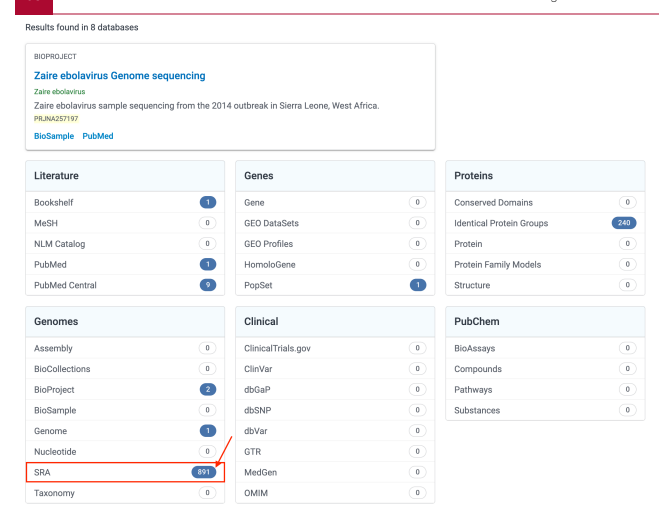
Navigating the NCBI website can be challenging. From a user perspective, nothing is as straight forward as you would expect. For the sequence read archive (SRA), there are fortunately some options. There are the convuluted e-utilites, which can grab run information without navigating to the NCBI website, or there is the SRA Run Selector (https://www.ncbi.nlm.nih.gov/Traces/study/). The Run Selector is nice because you can filter the results

for a given BioProject and obtain only the accessions that interest you in a semi user-friendly format. We can search directly from the SRA Run Selector, or simply go to the main NCBI website and begin our search from there. Let's see the latter example, as this is likely the primary way you will attempt to find data in the future.

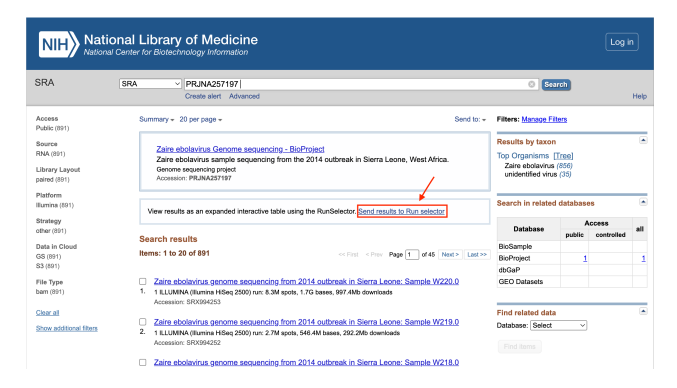
Step 1: Start from the NCBI homepage (https://www.ncbi.nlm.nih.gov/). Type the BioProject ID (PRJNA257197) into the search field.



Step 2: In the search results, select the entries next to the SRA.



Step 3: From the SRA results, select "Send results to Run Selector".



Step 4: From there, you can download the metadata or accession list to your local computer. You could also download the fastq files directly to local.

Copy and paste the accession list into a file using nano. Save to runaccessions.txt. Now use head to grab the first few results.

First, let's create a new directory to work in.

```
mkdir run_selector
cd run_selector
nano runaccessions.txt
```

```
head -n 3 runaccessions.txt > runids.txt
```

E-utilities

Using e-utilities to programmatically grab the run info

esearch and efetch, can be used to query and pull information from Entrez (https://www.ncbi.nlm.nih.gov/Web/Search/entrezfs.html). They aren't the easiest to understand or use, but you can use them to get the same run info that we grabbed using the Run Selector with the following one liner:

```
module load edirect
esearch -db sra -query PRJNA257197 | efetch -format runinfo > runinfo.csv
```

Here we provide the BioProject ID to esearch, which queries the SRA. That info can be piped to efetch, which will pull down the run info in comma separated format, which we can save to file using >.

Then we can use a combo of cat, cut, grep, and head to grab only the accession info we are interested in:

```
cat runinfo.csv | cut -f 1 -d ',' | grep SRR | head > runids.txt
```

Here, we opened the file with cat, piped it to the cut command, piped it to grep for only the accession IDs (skipping the column header), and get only the first few results.

So what is cut? It is a unix command, that is used to cut out selected portions of the file (man cut for more info). The -f option specifies which field to cut (the first field), and -d tells the program which delimiter to use. We know this is a comma-separate value file, so we do -d ', ' to specify the comma.

Using the "parallel" tool for automating downloads

Now that we have accession numbers to work with, let's use parallel and fastq-dump to download the data.

GNU parallel is a shell tool for executing jobs in parallel using one or more computers. (https://www.gnu.org/software/parallel/man.html) It can serve as a nice replacement of the for loop; though, its behavior is different. Check out this post from the Biostars forum, Tool: Gnu Parallel -

Parallelize Serial Command Line Programs Without Changing Them (https://www.biostars.org/p/63816/), and this from OMGenomics Lab, Using GNU parallel painlessly (https://omgenomics.com/blog/parallel).

Let's see how this works.

```
mkdir parallel
cd parallel
cat ../runids.txt | parallel -j 1 fastq-dump -X 10000 --split-files
```

This gives us six files, two files for each run and 3 runs. It takes the place of the multiple fastq-dump commands we would need to use.

What's up with the curly braces {}?

The curly braces are default behavior for parallel. This is the default replacement string; when empty, it appends inputs. So the following gets the same result.

```
cat ../runids.txt | parallel fastq-dump -X 10000 --split-files
```

```
More on parallel

The real power of parallel comes when using something between the brackets such as {.}, {/}, and {//}, like this...

nano filelist

Include the following:

/dir/subdir/file.txt
/other/list.csv
/raw/seqs.fastq

Save the file -> Ctrl 0, Yes, Ctrl X.

cat filelist | parallel echo {} "without extension" {.}

/dir/subdir/file.txt without extension /dir/subdir/file
/other/list.csv without extension /other/list
/raw/seqs.fastq without extension /raw/seqs

(This replaces the extension.)

cat filelist | parallel echo {} "without path" {/}
```

/dir/subdir/file.txt without path file.txt /other/list.csv without path list.csv /raw/seqs.fastq without path seqs.fastq

(This removes the path.)

```
cat filelist | parallel echo {} "with root" {//}
```

/dir/subdir/file.txt with root /dir/subdir /other/list.csv with root /other /raw/seqs.fastq with root /raw

(This keeps only the path.)

Note

parallel will default to 1 concurrent job per available CPU. On a shared system like helix with 48 CPUs, where users do run fast(er)q-dump, that can cause an overload. Even when submitting a job on Biowulf, you may not want to run as many concurrent jobs as there are allocated CPUs (e.g. if you ran fasterq-dump with 2 threads and you have 12 CPUs allocated --jobs would be 6). Therefore, it is good practice to always specify the -j flag, which assigns the number of jobs for the parallel command.

Using swarm

We can also take advantage of swarm for running jobs in parallel, as we did in the previous lesson.

Note

This should be run from Biowulf, not Helix.

```
cd ..
mkdir swarm
cd swarm
```

We can generate a script using:

```
cat ../runids.txt | while read file; do
    echo prefetch $file >> download.sh
    echo 'fasterq-dump -t /lscratch/$SLURM_JOB_ID' $file >> download
done
```

Edit to include #SBATCH directives.

```
nano download.sh
```

At the top of the file add the following:

```
#SWARM --threads-per-process 6

#SWARM --gb-per-process 4

#SWARM --gres=lscratch:10

#SWARM --module sratoolkit

#SWARM --partition=student
```

Submit the script.

```
swarm -f download.sh
```

European Nucleotide Archive (ENA)

Everything in the SRA is also in the ENA (See PRJNA257197 on ENA (https://www.ebi.ac.uk/ena/browser/view/PRJNA257197?show=reads)). Files in the ENA share the same naming convention as the SRA but are stored directly as gzipped fastq files and bam files. You can easily use wget or curl to pull files directly from the ENA without using the sratoolkit. Check out the ENA training modules (https://ena-docs.readthedocs.io/en/latest/retrieval/file-download.html#) for more information on downloading data from the ENA, including examples.

```
Using curl and wget
```

The curl command is used to retrieve data from web sites. A similar command is wget. The Unix system you are working with may have either curl or wget installed. To see which is active on your system, just type the command at the command line.

```
wget
curl
```

Let's see how we can use wget to grab a couple of files from the ENA.

We use wget followed by the path to the file:

```
wget -nc ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR155/006/SRR1553426/SRR1553426_2.fast wget -nc ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR155/006/SRR1553426/SRR1553426_1.fast
```

The -nc option refers to "no clobber", which will not overwrite existing files.

To see additional options, use

man wget

These files are compressed. To decompress a file use gunzip.

gunzip SRR1553426_2.fastq.gz

In the case of .zip you can use unzip, and with tar.gz, you can use tar -xvf. When in doubt, use google.

Note

Many programs accept compressed fastq files, so rarely will you need to decompress a fastq file.

There is also a fantastic web service called SRA Explorer (https://sra-explorer.info/#) that can be used to easily generate code for file downloads from the SRA, if you are looking to download a maximum of 500 samples. Use caution, the SRA-explorer is not associated with the ENA or NCBI, and program support / updates cannot be guaranteed.

Help Session

Practice the skills learned in this lesson here.

Module 2 - Introduction to RNA Sequencing

Lesson 7: Introduction to Next Generation Sequencing (NGS) Data and Quality Control

Lesson 6 Review

Lesson 6 introduced the basics of RNA sequencing, including experimental considerations and basic ideas behind data analysis. In lessons 7 through 14 participants will get hands-on experience with RNA sequencing analysis.

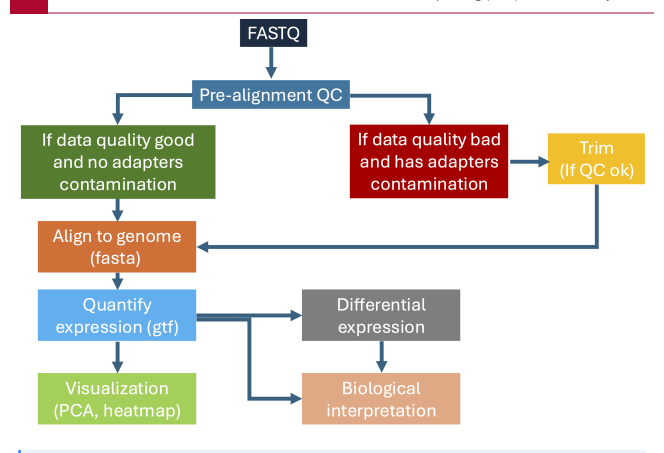
Learning Objectives

After this class, participants will be able to:

- Describe the format in which NGS data is stored.
- Know why sequence quality score is important and how this is calculated.
- Run quality assessment on NGS data and interpret results.

Basic RNA Sequencing Analysis Workflow

The diagram below shows a basic workflow for RNA sequencing analysis, which starts with FASTQ files as input. Quality check is performed on the FASTQ files ensure that sequencing quality is good and that there are no contaminations such as those arising from adapter read through (https://knowledge.illumina.com/software/general/software-general-reference_material-list/000002905). If there are no issues arising from quality check, perform alignment to determine where in the genome each sequence comes from. If issue(s) arise from quality check, then perform trimming (either to remove low quality reads or contamination such as adapters) and perform QC to ensure the trimmed sequences are good prior to alignment. Quantification of gene expression can be performed on the aligned data. From the expression quantification, the researcher can construct visualizations (ie. PCA and heatmap) as well as perform differential expression analysis and biological interpretation.



Note

A reference genome in the form of a fasta (or .fa) file is needed for the alignment step while a gtf file describing location of genomic features (ie. genes, transcripts, exons, and introns) is needed for the quantification step.

Example Data

The data used are located in /data/classes/BTEP/hcc1395_b4b on Biowulf. This data was obtained from rnabio.org (https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/) and essentially a transcriptomic profiling study comparing between tumor and normal HCC1395 breast cancer cells. There a 3 normal and 3 tumor samples for each condition.

Sign onto Biowulf

Before following along in these exercises, please sign onto Biowulf using the ssh construct below where "user" is the class participant's assigned Biowulf student account ID.

ssh user@biowulf.nih.gov

Request an Interactive Session

Request an interactive session to perform compute intensive tasks on Biowulf by issuing the sinteractive command construct below where:

- --mem: enables the user to request the desired amount of RAM (https://en.wikipedia.org/wiki/Random-access_memory) for analysis (12gb requested in this example).
- --gres: enables the user to request generic compute resources such as local temporary storage space or lscratch (5gb requested in this example).

```
sinteractive --mem=12gb --gres=lscratch:5
```

Copy the Data to User's /data folder

The data is located in the hcc1395_b4b folder of /data/classes/BTEP. Copy this to the participants' own /data directory in Biowulf prior to proceeding using the cp command with the -r option for copying folders. The directory path where the data resides is /data/classes/BTEP/hcc1395_b4b and the destination to which it needs to copied is /data/user/ and name the copied folder hcc1395_b4b. Be sure to change "user" to the participant's assigned Biowulf student ID.

```
cp -r /data/classes/BTEP/hcc1395_b4b /data/user/hcc1395_b4b
```

Use the 1s command to list the content in the participant's Biowulf /data folder to make sure the copy was successful.

1s

hcc1395_b4b

FASTQ Files for hcc1395 b4b

Change into the hcc1395 b4b folder in the participant's Biowulf /data directory.

```
cd hcc1395_b4b
```

Next, use 1s with the -1 option to list the contents in the reads directory one item per line.

```
ls -1 reads
```

The reads folder contains the FASTQ or .fq files for the hcc1395 data. There are 12 of these, two for each sample as this study generated paired end sequences.

```
hcc1395_normal_rep1_R1.fq
hcc1395_normal_rep2_R1.fq
hcc1395_normal_rep2_R2.fq
hcc1395_normal_rep3_R2.fq
hcc1395_normal_rep3_R2.fq
hcc1395_tumor_rep1_R1.fq
hcc1395_tumor_rep1_R2.fq
hcc1395_tumor_rep2_R2.fq
hcc1395_tumor_rep2_R1.fq
hcc1395_tumor_rep2_R2.fq
hcc1395_tumor_rep3_R2.fq
hcc1395_tumor_rep3_R2.fq
```

Read 1 Read 2

In paired end sequencing, a template or unknown nucleotide fragment is sequenced from both ends, thus generating two FASTQ files per sample. A gap between the two reads is usually present in this sequencing protocol.

Image source: https://www.ebi.ac.uk/training/online/courses/functional-genomics-ii-common-technologies-and-data-analysis-methods/rna-sequencing/performing-a-rna-seq-experiment/design-considerations/ (https://www.ebi.ac.uk/training/online/courses/functional-genomics-ii-common-technologies-and-data-analysis-methods/rna-sequencing/performing-a-rna-seq-experiment/design-considerations/).

Overview of a FASTQ File

Each FASTQ file is composed of many sequences. The tool seqkit and its stats function can be used to get statistics the hcc1395 FASTQ files.

Change in the reads folder for this.

cd reads

Load seqkit.

```
module load seqkit
```

```
[+] Loading seqkit 2.7.0
```

Run seqkit stats for all of the hcc1395 FASTQ files in one go by using * before the .fq extention. * acts as a wildcard.

```
seqkit stats *.fq
```

Query of the stats for the FASTQ files generates the results below, which informs of things such as the number of sequences (or reads) in a FASTQ file. For the hcc1395_normal_rep1 biological replicates, both files in the pair (R1 and R2) have 331,958 sequences. The average length of the sequences in these files is 151 bases. Pairs in paired end sequencing should have the same number of sequences because the reads were generated from the same template.

file	format	type	num_seqs	sum_len	min_le
hcc1395_normal_rep1_R1.fq	FASTQ	DNA	331,958	50,125,658	1!
hcc1395_normal_rep1_R2.fq	FASTQ	DNA	331,958	50,125,658	1!
hcc1395_normal_rep2_R1.fq	FASTQ	DNA	331,958	50,125,658	1!
hcc1395_normal_rep2_R2.fq	FASTQ	DNA	331,958	50,125,658	1!
hcc1395_normal_rep3_R1.fq	FASTQ	DNA	331,956	50,125,356	1!
hcc1395_normal_rep3_R2.fq	FASTQ	DNA	331,956	50,125,356	1!
hcc1395_tumor_rep1_R1.fq	FASTQ	DNA	390,607	58,981,657	1!
hcc1395_tumor_rep1_R2.fq	FASTQ	DNA	390,607	58,981,657	1!
hcc1395_tumor_rep2_R1.fq	FASTQ	DNA	390,607	58,981,657	1!
hcc1395_tumor_rep2_R2.fq	FASTQ	DNA	390,607	58,981,657	1!
hcc1395_tumor_rep3_R1.fq	FASTQ	DNA	390,607	58,981,657	1!
hcc1395_tumor_rep3_R2.fq	FASTQ	DNA	390,607	58,981,657	1!

Recall from the seqkit stat results shown above that each FASTQ file contains many sequencing reads. The record for each sequencing read is composed of four lines and these include the following.

• The first line is the sequencing read metadata (ie. instrument used, location on the flow cell where the read was derived, and whether the read is the first or second of a pair in paired end sequencing) - in this example,

- "/1" at the end of the metadata line in the reads from hcc1395_normal_rep1_R1.fq denotes the first read of a pair
- "/2" at the end of the metadata line in the reads from hcc1395_normal_rep1_R2.fq denotes the second read of a pair
- The second line is the biological sequence
- The third line is a "+"
- The fourth line contains the quality score of each of the bases in the sequencing read.

 The quality score reflects the likelihood that a base at a particular location along the read is wrong.

Below, head -4 is used to show the first sequencing reads of hcc1395_normal_rep1_R1.fq and hcc1395_normal_rep1_R2.fq. -4 in the head command tells it to only retrieve the first 4 lines in a file.

```
head -4 hcc1395_normal_rep1_R1.fq
```

@K00193:38:H3MYFBBXX:4:1101:10003:44458/1
TTCCTTATGAAACAGGAAGAGTCCCTGGGCCCAGGCCTGGCCCACGGTTGTCAAGGCACATCATTGCC/+

```
head -4 hcc1395_normal_rep1_R2.fq
```

```
@K00193:38:H3MYFBBXX:4:1101:10003:44458/2
ATGGAACTCCCTACTGACCTCTGAGAACTGGAAACGAGTTTGTACAGAAGTCAGAACTTTGGGTTGGG/+
```

Click here (http://www.drive5.com/usearch/manual/quality_score.html) and here (https://training.galaxyproject.org/training-material/topics/sequence-analysis/tutorials/quality-control/slides-plain.html) to learn more about the quality scores. The image below shows a table that translates the characters you see in the quality score lines to a numeric score.

ASCII_BASE=33 Illumina, Ion Torrent, PacBio and Sanger											
Q	P_error	ASCII	Q	P_error	ASCII	Q	P_error	ASCII	Q	P_error	ASCII
0	1.00000	33 !	11	0.07943	44 ,	22	0.00631	55 7	33	0.00050	66 B
1	0.79433	34 "	12	0.06310	45 -	23	0.00501	56 8	34	0.00040	67 C
2	0.63096	35 #	13	0.05012	46 .	24	0.00398	57 9	35	0.00032	68 D
3	0.50119	36 \$	14	0.03981	47 /	25	0.00316	58 :	36	0.00025	69 E
4	0.39811	37 %	15	0.03162	48 0	26	0.00251	59 ;	37	0.00020	70 F
5	0.31623	38 €	16	0.02512	49 1	27	0.00200	60 <	38	0.00016	71 G
6	0.25119	39 '	17	0.01995	50 2	28	0.00158	61 =	39	0.00013	72 H
7	0.19953	40 (18	0.01585	51 3	29	0.00126	62 >	40	0.00010	73 I
8	0.15849	41)	19	0.01259	52 4	30	0.00100	63 ?	41	0.00008	74 J
9	0.12589	42 *	20	0.01000	53 5	31	0.00079	64 @	42	0.00006	75 K
10	0.10000	43 +	21	0.00794	54 6	32	0.00063	65 A			

Note

 $\qquad $\qquad = -10\times \log 10(error\ probability)$

Quality Assessment on the hcc1395 FASTQ Files

Go back to the hcc1395 b4b folder.

```
cd /data/users/hcc1395_b4b
```

Make a folder called pre_alignment_qc_raw.

```
mkdir pre_alignment_qc_raw
```

Load fastqc, which is a tool used to perform quality check on Illumina sequencing data.

```
module load fastqc
```

```
[+] Loading singularity 4.1.5 on cn0047
[+] Loading fastqc 0.12.1
```

Change into the pre alignment qc raw folder.

```
cd pre_alignment_qc_raw
```

Next, run fastqc using the following command construct where:

• ../reads/*.fq tells fastqc to look one directory back ../ and then in the reads folder and perform QC on all of the .fq files. * is used as a wildcard.

• -o is an option in fastqc that allows for specification of output folder. In this case, the output folder should be pre_alignment_qc_raw. Since the current directory is pre_alignment_qc_raw, the '.' can be used to denote here, in the current folder.

```
fastqc ../reads/*.fq -o .
```

List the contents for pre_alignment_qc_raw after QC completes. FASTQC generates a html report and provides the results in a zip file for each file. The html report can be viewed on a local web browser.

ls

```
hcc1395_normal_rep1_R1_fastqc.html
                                    hcc1395_tumor_rep1_R1_fastqc.htm
hcc1395 normal rep1 R1 fastqc.zip
                                    hcc1395 tumor rep1 R1 fastqc.zip
hcc1395_normal_rep1_R2_fastqc.html
                                    hcc1395_tumor_rep1_R2_fastqc.htm
hcc1395 normal rep1 R2 fastqc.zip
                                    hcc1395 tumor rep1 R2 fastqc.zip
hcc1395 normal rep2 R1 fastqc.html
                                    hcc1395 tumor rep2 R1 fastqc.htm
hcc1395_normal_rep2_R1_fastqc.zip
                                    hcc1395_tumor_rep2_R1_fastqc.zip
hcc1395 normal rep2 R2 fastqc.html
                                    hcc1395 tumor rep2 R2 fastqc.htm
hcc1395_normal_rep2_R2_fastqc.zip
                                    hcc1395_tumor_rep2_R2_fastqc.zip
hcc1395 normal rep3 R1 fastqc.html
                                    hcc1395 tumor rep3 R1 fastqc.htm
                                    hcc1395_tumor_rep3_R1_fastqc.zip
hcc1395_normal_rep3_R1_fastqc.zip
hcc1395_normal_rep3_R2_fastqc.html
                                    hcc1395_tumor_rep3_R2_fastqc.htm
hcc1395_normal_rep3_R2_fastqc.zip
                                    hcc1395_tumor_rep3_R2_fastqc.zip
```

Copy to hcc1395_normal_rep1_R1_fastqc.html to local computer to learn how to interpret a FASTQC report. To do this open another terminal (Mac) or command prompt (Windows) and change into the local Downloads folder.

```
cd Downloads
```

In the scp construct below, change user to the participant's assigned Biowulf student ID and can be broken down as follows.

- user@helix.nih.gov: connects the user to helix, which is the HPC partition used for interactive data transfer.
- : separates the helix login part from the path of the file or folder to transfer, which is / data/user/hcc1395_b4b/pre_alignment_qc_raw/ hcc1395_tumor_rep1_R1_fastqc.html.
- .: specifies for scp to download to the current directory, which should be the local Downloads folder.

Enter the user's Biowulf password when prompted and the download should commence.

scp user@helix.nih.gov:/data/user/hcc1395_b4b/pre_alignment_qc_raw/hc

Interpreting FASTQC Report

Report Navigator and Summary

The first item is the navigation panel that enables scientists to quickly link to different portions of the FASTQC report. The QC modules that passed are indicated by a green circle with a check. Those with warnings are denoted by a yellow circles with a "!". Failed modules are indicated by a red circle with "x".

Next there is a summary statistics table, which includes the following information.

- Name of the FASTQ file in which the report was generated for (hcc1395_tumor_rep1_R1.fq in this case)
- The number of sequences in the file.
- Number of sequences flagged with poor quality.
- The sequence length (ie. how many bases are in each sequencing read of the FASTQ file).

PastQC Report

Summary





Per tile sequence quality

Per sequence quality scores

Per base sequence content

Per sequence GC content

Per base N content

Sequence Length Distribution

Sequence Duplication Levels

Overrepresented sequences

Adapter Content



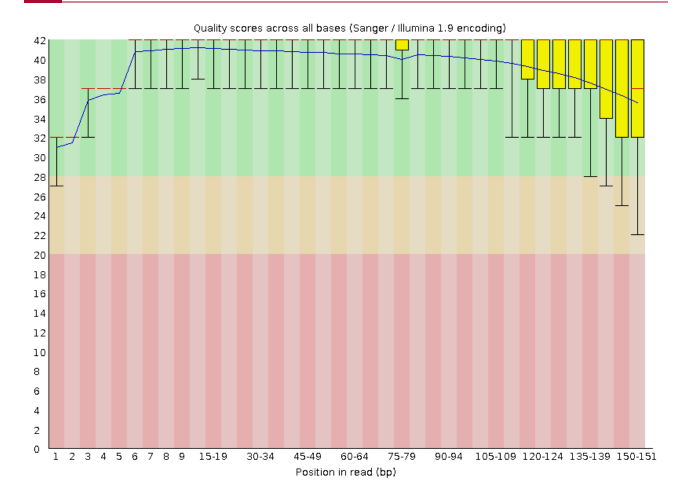
Measure	Value			
Filename	hcc1395_tumor_rep1_R1.fq			
File type	Conventional base calls			
Encoding	Sanger / Illumina 1.9			
Total Sequences	390607			
Total Bases	58.9 Mbp			
Sequences flagged as poor quality	0			
Sequence length	151			
%GC	53			

Per Base Sequence Quality

Next, there is the "Per base sequence quality" plot.

This chart plots the error likelihood at each base position averaged over all sequences in a FASTQ file.

- On the vertical axis are the quality scores that are in row 4 of the sequencing reads in a FASTQ file. These quality scores represent error probabilities, where:
 - 10 corresponds to 10% error (1/10),
 - 20 corresponds to 1% error (1/100),
 - 30 corresponds to 0.1% error (1/1000) and
 - 40 corresponds to one error every 10,000 measurements (1/10,000) that is an error rate of 0.01%
- The three colored bands (green, yellow, red) illustrate the typical labels assigned to these measure:
 - reliable (28-40, green)
 - less reliable (20-28, yellow)
 - error prone (1-20, red)
- The yellow boxes contain 50% of the data, the whiskers indicate the 75% outliers.
- The red line inside the yellow boxes is the median quality score for that base.
- The blue line is the average quality score at a particular base.



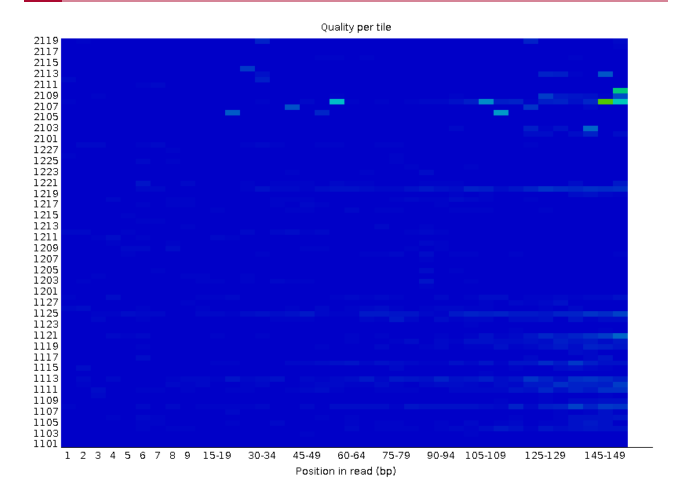
Per Tile Sequence Quality

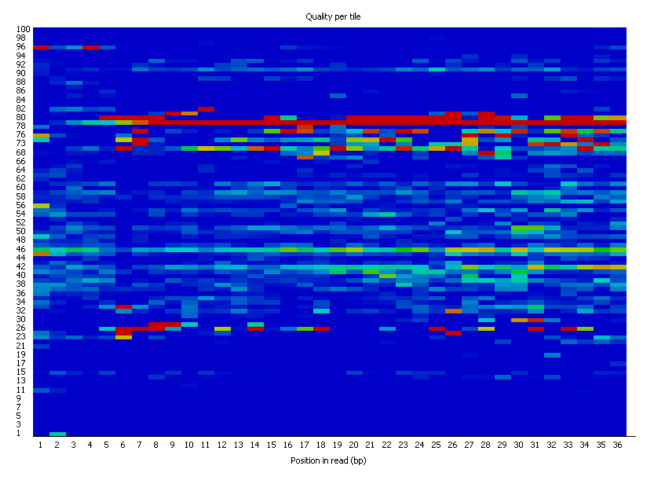
Next, there is the "Per tile sequence quality" plot. This graph informs whether something is wrong with a tile in the flow cell. Along the horizontal axis are the base positions. The vertical axis represents the tile number in which the read came from.

This plot compares the average quality score of a tile to the average quality score of all tiles at a particular base position. -- https://sequencing.qcfail.com/articles/position-specific-failures-of-flowcells/ (https://sequencing.qcfail.com/articles/position-specific-failures-of-flowcells/)

The interpretation for this plot is as follows

- Colder colors indicate that the average quality score at a tile is at or above the average quality score of all tiles at a particular base position. So a plot that is entirely blue is good.
- Hotter colors indicate that the average quality score at a tile is below the average quality score of all tiles at a particular base position. So a plot with red indicates a part of the flow cell has problems.

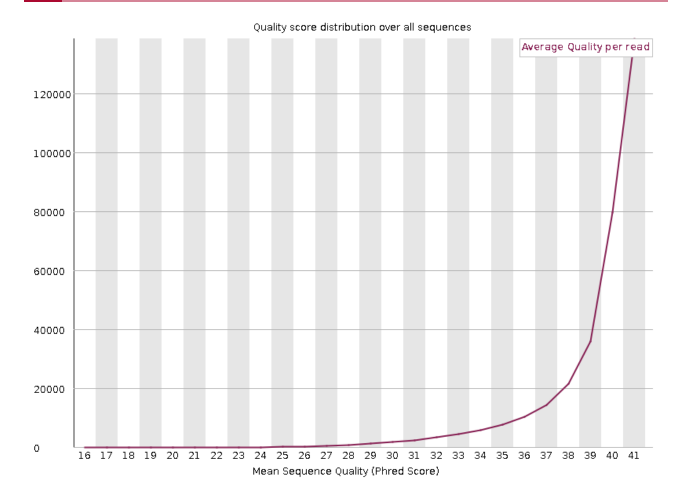




Bad per tile quality. Source: https://www.bioinformatics.babraham.ac.uk/projects/fastqc/Help/3%20Analysis%20Modules/12%20Per%20Tile%20Sequence%20Quality.html (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/Help/3%20Analysis%20Modules/12%20Per%20Tile%20Sequence%20Quality.html)

Per Sequence Quality Scores

Next, the distribution of the sequence quality scores is shown in the "Per sequence quality scores" plot. This graph can reveal whether the quality of a subset of sequences is poor. Here the sequences have good quality, where most reads have quality that clusters at around 37 or above (~0.02% error likelihood).

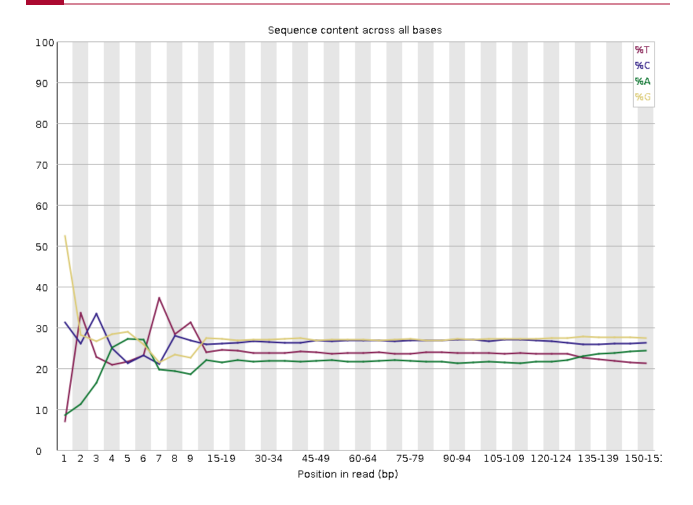


Per Base Sequence Content

The next figure shows "Per base sequence content", which is essentially the sequence make up along the bases of reads in the FASTQ file. If a library is random, then the percent composition of each nucleotide base (A,T,C,G) should be the same (~25%).

This module fails for hcc1395_tumor_rep1_R1.fq because the difference between the percentage of A and the percentage of T is larger than 20 at a particular location **OR** the difference between the percentage of C and the percentage of G is larger than 20 at a particular location -- Babraham bioinformatics Per base sequence content (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/Help/3%20Analysis%20Modules/4%20Per%20Base%20Sequence%20Content.html).

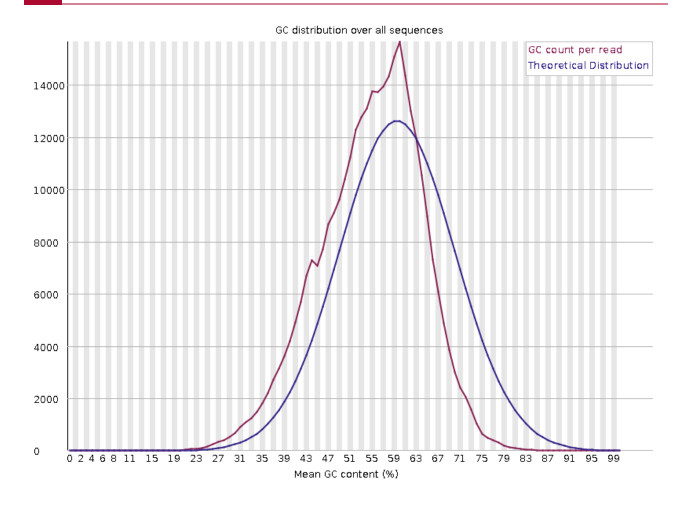
In hcc1395_tumor_rep1_R1.fq, it looks like the difference between the percent composition of A and T at base position 2 is causing the failure. Unfortunately, this type of uneveness in base distribution at the beginning of a read is observed in RNA sequencing due to priming with random hexamers during the library preparation stage.



Per Sequence GC Content

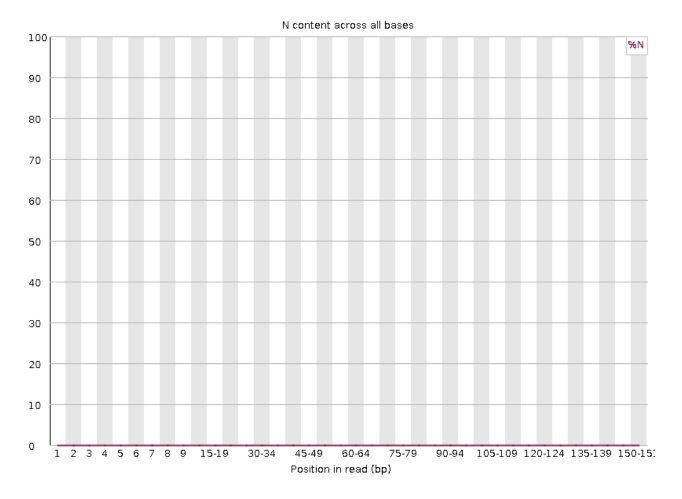
Next, the GC content across each sequence compared to a normal distribution in what is shown in the "Per sequence GC content" plot. The GC content in hcc1395_tumor_rep1_R1.fq is off from the normal theoretical distribution.

The peak of this theoretical distribution is an estimate of the GC content of the underlying genome. Deviation from of the GC content from the theoretical distribution could be caused by contamination or sequencing bias. -- Babraham bioinformatics Per base sequence content (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/Help/3%20Analysis%20Modules/5%20Per%20Sequence%20GC%20Content.html).

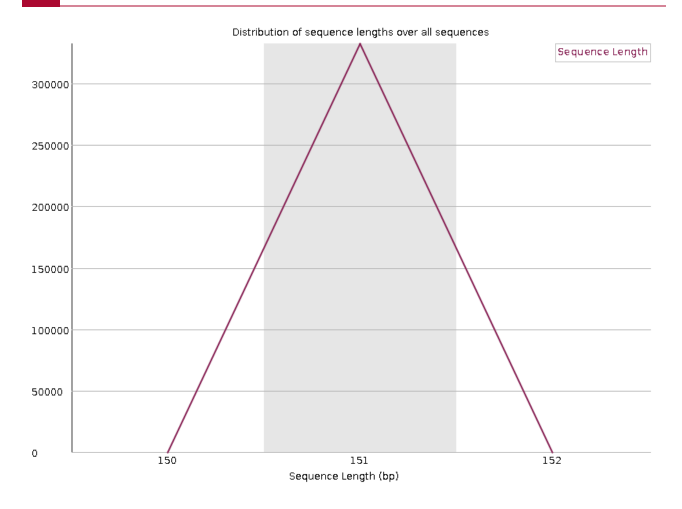


Per Base N Content

Whether there are any unknown bases in the sequencing reads is shown in next plot, which is the "Per base n content".



The sequence length distribution of hcc1395_tumor_rep1_R1.fq in what is known as the "Sequence Length Distribution" plot. All sequences have 151 bases as trimming has not been performed.

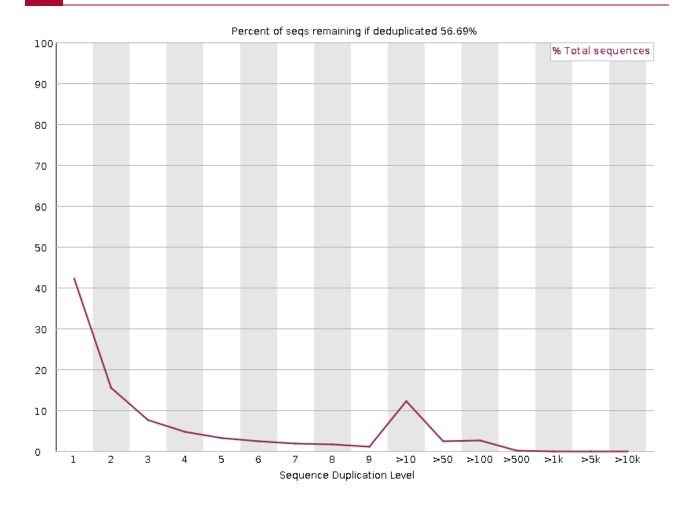


Sequence Duplication Level

The plot below shows the sequence duplication levels. High levels of duplication may indicate an enrichment bias such as over-amplification in the PCR step. However, in RNA sequencing, duplicate sequences could be biologically meaningful as these could be derived from multiple copies of the same transcript.

See here (https://dnatech.ucdavis.edu/faqs/should-i-remove-pcr-duplicates-from-my-rna-seq-

data#:~:text=The%20short%20and%20generalized%20answer,are%20analyzed%20without%20duplic for advice regarding handling duplicates in RNA sequencing.



Overrepresented Sequences

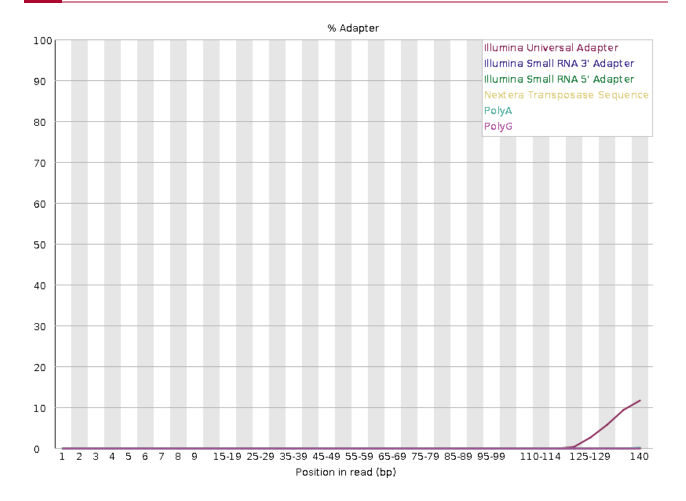
The overrepresented sequences graph is shown next. Presence of overrepresented sequences may indicate enrichment artifact or the sequencing library is not very diverse. On the other hand, overrepresented sequences could have biological significance.



Sequence	Count	Percentage	Possible Source
${\tt CTGGAGTCTTGGAAGCTTGACTACCCTACGTTCTCCTACAAATGGACCTT}$	1544	0.39528221460444896	No Hit
${\tt CTACGTTCTCCTACAAATGGACCTTGAGAGCTTGTTTGGAGGTTCTAGCA}$	654	0.1674317152534389	No Hit
${\tt GGGAGCAGAGGCAGCTGCCTTTATTGGGGGCCATGGGCTGGTTTTCA}$	597	0.1528390428230933	No Hit
${\tt GTGGAAAATAACTTTTATTGAGACCCCACCAACTGCAAAATCTGTTCCTG}$	594	0.15207100743202245	No Hit

Adapter Contamination

If there is adapter contamination in the sequences, then this will be shown in the next plot.



Conclusions from hcc1395_normal_rep1_R1.fastq Quality Assessment

In general, FASTQC shows that the file hccl395_normal_repl_R1.fastq have good quality sequences (less than 1% error likelihood). The concern is the presence of Illuminal universal adapters.

Merging Multiple FASTQC Reports into One

Reviewing multiple FASTQC reports is cumbersome but there is an application called MultiQC that allows scientists to combine many FASTQC reports into one.

To use MultiQC on Biowulf (or other HPC systems), first load it using the following.

```
module load multiqc
```

The --filename option allows users to enter a base name (ie. file name without an extension) for the output MultiQC report. Here, the base name pre_alignment_qc_raw will be used.

```
multiqc --filename pre_alignment_qc_raw .
```

ls

MultiQC generates a folder that contains the QC data should the researcher choose to interrogate these programmatically. It provides a report in the html format as well, which can be downloaded to local computer and viewed in a web browser.

```
pre_alignment_qc_raw_data
pre_alignment_qc_raw.html
```

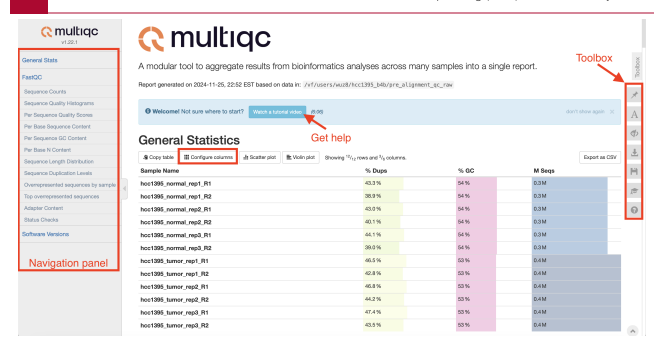
Take a look at the MultiQC html report by copying it to the Downloads folder on local computer using scp.

```
scp user8@helix.nih.gov:/data/user/hcc1395_b4b/pre_alignment_qc_raw,
```

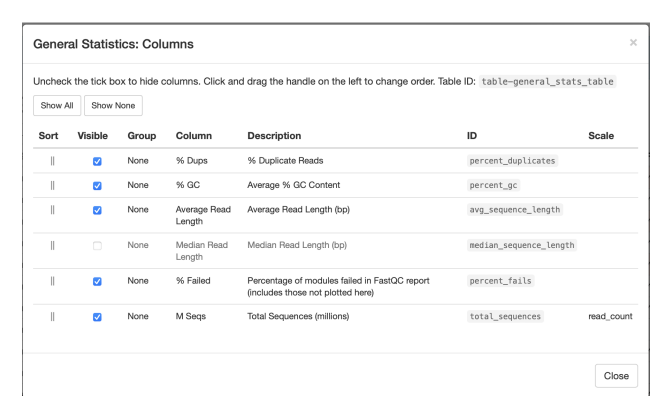
MultiQC Navigation Panel and Summary Statisics

Upon opening the MultiQC report, the user will be greeted with a navigation panel (similar to that found in the individual FASTQC reports) that allows users to quickly move to different sections of the report. A link to get help for using the MultiQC report is available as well. To the right of the report page, there is a tool box that allows the researcher to do things like highlighting different samples in different colors for better visualization, rename samples, and export each of the individual QC plots as an image for inclusion in presentations and/or publications. MultiQC reports are interactive.

The General Statistics table shows some basic statistics about the samples, including percentage of duplicate reads, GC content, and number of sequences (reported as million of sequences in the M Seqs column).



Click on the Configure Columns tab to choose which columns to include in the General Statistics table and the plot button to visualize the data in graphical format.



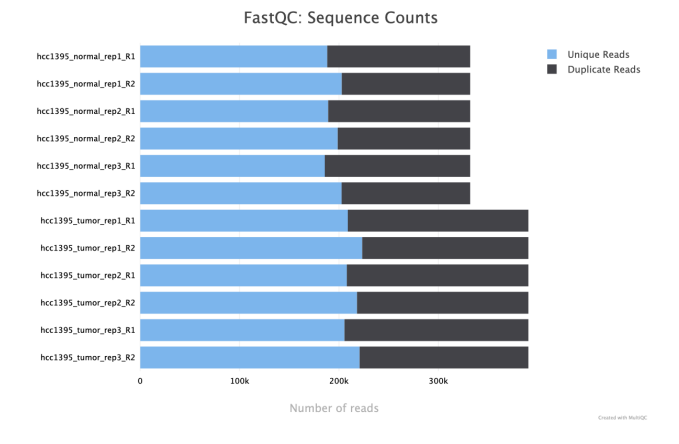
Altering the way the total number of sequences is reported

The number of sequences could be reported as the actual number rather than in millions. Users can alter this by issuing the `--config` option in `multiqc` and specifying a custom report configuration `yaml` file. The content for this configuration file is shown below and can be explained as follows. * `read_count_multiplier`: set to 1 to report the actual sequence number. * `read_count_prefix`: set to "" to remove the units at the end of the number of sequences. The default is "M" for milion. * Under `table_columns_name` change the `total_sequences` column heading to "# Sequences" or any name that reflects the fact the actual number of sequences is being reported.

```
read_count_multiplier: 1
read_count_prefix: ""
table_columns_name:
   FastQC:
     total_sequences: "# Sequences"
```

MultiQC Sequence Count

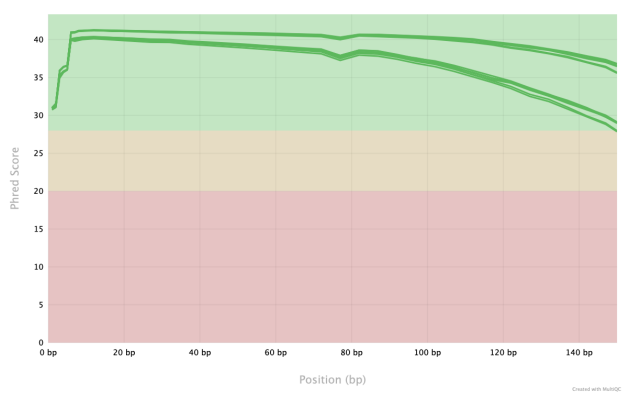
The Sequence Count plot shows the break down of unique and duplicate reads for each FASTQ file. Again, duplication suggests some sort of enrichment bias or in the case of RNA sequencing, it could be that the duplicates are biologically meaningful due to multiple copies of the same trancript being expressed.



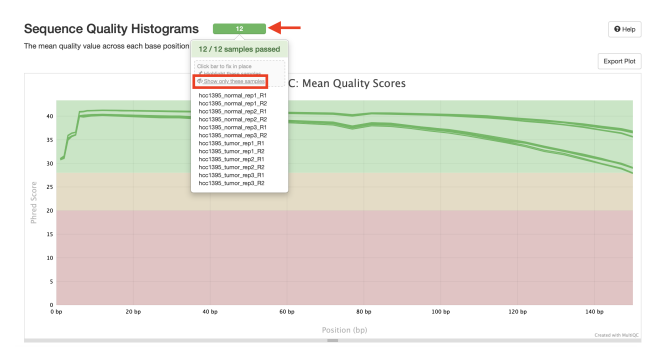
MultiQC Mean Quality Score

The average quality score of the sequencing reads in FASTQ files along each base position is shown in the figure below. Regarding the boxes at the top of the QC plots, green means QC passed while orange and red indicate warning and failed, respectively.

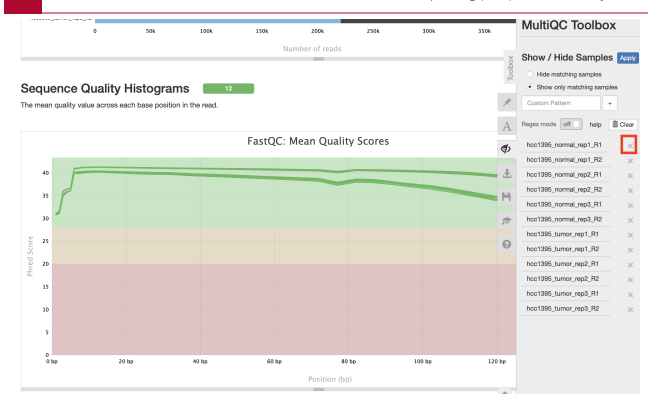




Click on the green rectangle labeled with the number of files to filter out samples to see in the plot.

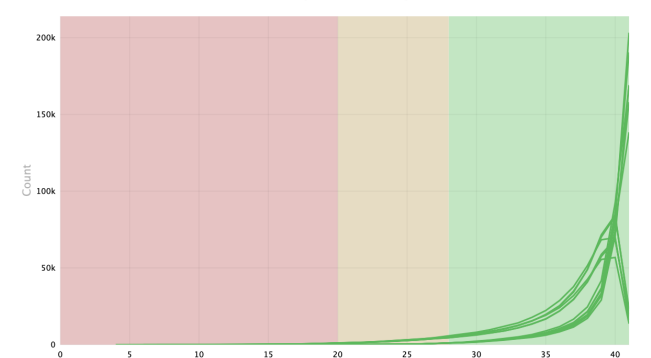


Upon clicking on the green rectangled labeled with the number of files, a tool box will appear and users can click on the "x" next to each file to deselect and remove it from the visualization.



MultiQC Quality Score Distribution

The quality score distributions of each of the FASTQ files are plotted in the next figure.



Mean Sequence Quality (Phred Score)

FastQC: Per Sequence Quality Scores

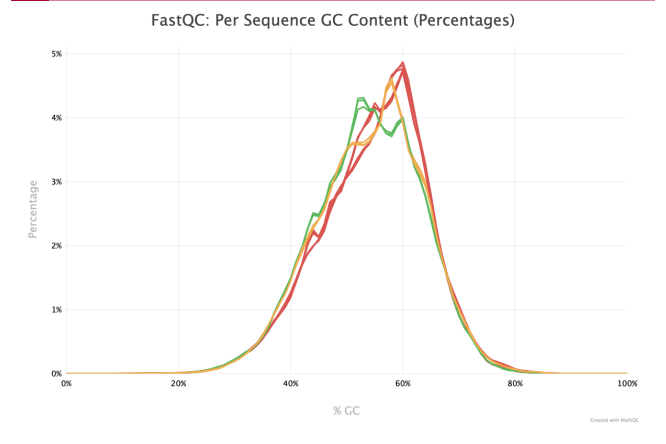
MultiQC Per Base Sequence Content

An interactive heatmap of the percent composition of each nucleotide base (A,T,C,G) along the bases (horizontal axis) for each of the FASTQ files (vertical axis) is presented next. Hover over a tile to see the corresponding numbers for a given sample. Click on any row in this heatmap to get the base composition plot for just that sample.



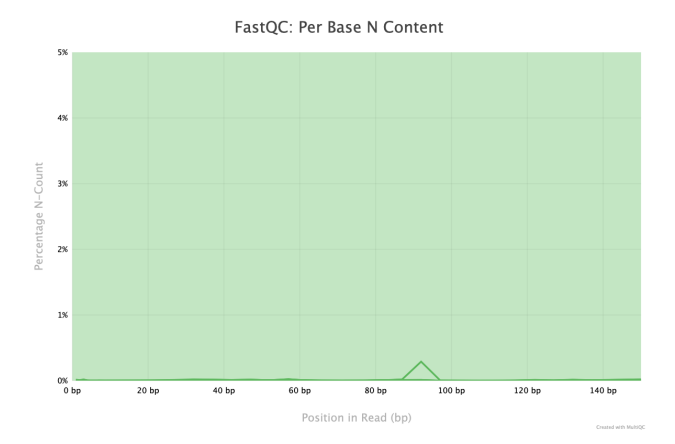
MultiQC GC Distribution

The GC distribution for each of the FASTQ files is shown in figure below.



MultiQC Per Base N Content

The percentage of unknown bases (or N) per FASTQ file is available as the next figure.



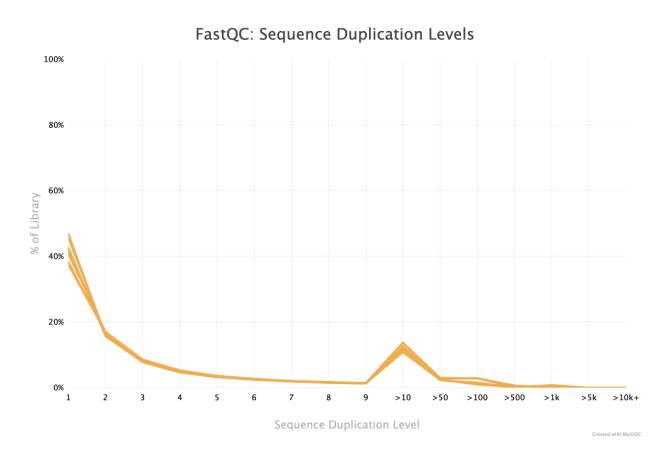
MultiQC Sequence Length Distribution

As shown below, all FASTQ files for the hcc13995 study have sequences that contain 151 bases.



MultiQC Duplication Level

The percentage of sequences with the duplication level shown on the horizontal axis.

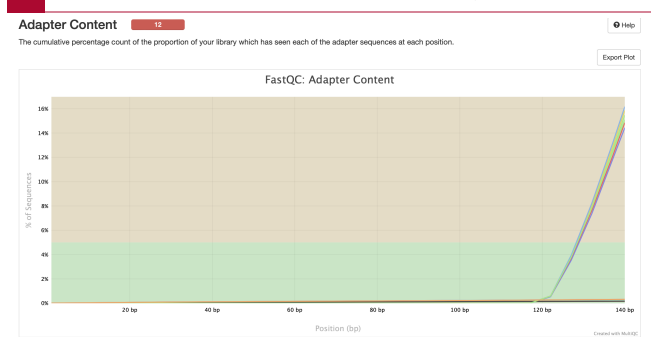


MultiQC Adapter Contamination

The sequences in the hcc1395 data have adapter contamination. Mousing over the area where the adpater content starts to increase suggests Illumina Universal Adapter.

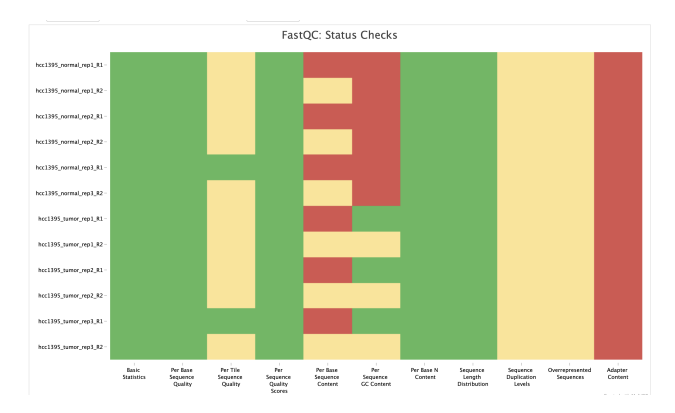
Note

FASTQC will issue a warning if any sequence is present in more than 5% of all reads and failure if any sequence is present in more than 10% of all reads. -- FASTQC manual (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/Help/3%20Analysis%20Modules/10%20Adapter%20Content.html)



MultiQC Status Heatmap

Finally, a heatmap showing the per file QC modules that have passed, warn, or failed is provided.



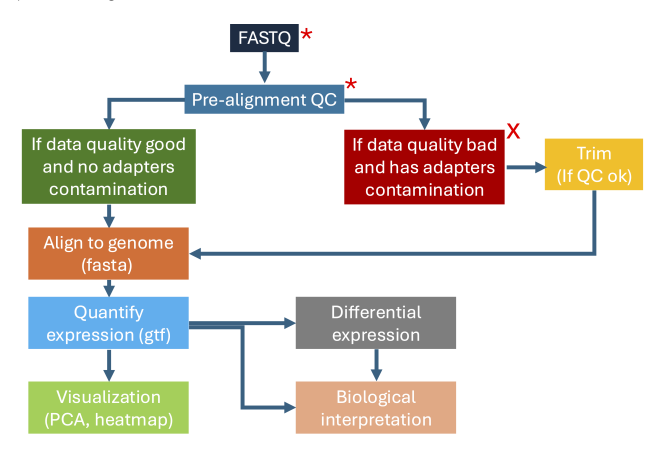
View FASTQC and MultiQC HTML Reports

Untrimmed FASTQ Files

- hcc1395_normal_rep1_R1
- hcc1395_normal_rep1_R2
- hcc1395_normal_rep2_R1
- hcc1395_normal_rep2_R2
- hcc1395_normal_rep3_R1
- hcc1395_normal_rep3_R2
- hcc1395_tumor_rep1_R1
- hcc1395_tumor_rep1_R2
- hcc1395_tumor_rep2_R1
- hcc1395_tumor_rep2_R2
- hcc1395_tumor_rep3_R1
- hcc1395_tumor_rep3_R2
- MultiQC

Lesson 7 Review

Lesson 7 introduced the FASTQ file, which is the format used to store Next Generation Sequencing (NGS) data. Participants also learned about assessing quality of the sequences in FASTQ files using the tool FASTQC. This step is essential as it will inform whether sequencing is of high quality and if there are contamination such as adapters in the sequences. QC results revealed that while sequencing data quality is good, there are adapters in the sequences. Adapters will interfere with the mapping stage where algorithms are used to determine where in the genome the sequences came from. Thus, the next step is to trim away the adapters and perform QC again on the trimmed data to make sure the contamination has been removed.



Learning objectives

At the end of this session, participants will be able to use the tool Trimmomatic (http://www.usadellab.org/cms/?page=trimmomatic) to remove adapters from the NGS data.

Sign onto Biowulf and Request an Interactive Session

Before getting started, sign onto Biowulf and request an interactive session. In the ssh command below, replace user with the participant's assigned Biowulf student ID.

ssh user@biowulf.nih.gov

Change into /data/user/hcc1395_b4b.

cd /data/user/hcc1395_b4b

Next, request an interactive session with 12 gb of RAM and 10 gb of local temporary storage. The option --cpus-per-task is used to request 6 CPUs on Biowulf in the sinteractive command below in addition to the 12 gb of memory and 10 gb of local temporary storage.

sinteractive --cpus-per-task 6 --mem=12gb --gres=lscratch:10

Adapter Trimming with Trimmomatic

Load Trimmomatics

Trimmomatic is a tool that can remove both low quality sequence and adapters. The first step to using Trimmomatic on Biowulf is to load it.

module load trimmomatic

Tip

Other tools used for trimming include bbduk (https://jgi.doe.gov/data-and-tools/software-tools/bbtools/bb-tools-user-guide/bbduk-guide/) and Cutadapt (https://cutadapt.readthedocs.io/en/stable/). Both are capable of quality and adapter trimming.

Make a New Folder to Stored the Trimmed Reads

mkdir trimmed_reads

Stay in the /data/user/hcc1395 b4b folder for these exercises.

Run Trimmomatic

To run remove adapters for all FASTQ files in one go, the parallel command will be introduced. This command enables the analyst to run multiple tasks in parallel such as trimming of high throughput sequencing data. The command construct is broken down below.

- cat is used to print the hcc1395 samples ids stored in the file hcc1395_sample_ids.txt. Rather than printing to terminal, | is used to send the output from cat to parallel.
 - The Trimmomatic construct is enclosed in double quotes within the parallel command. The components are as follows.
 - j: enables users to specify how many jobs to run in parallel (6 in this case since there are 6 samples).
 - java -jar \$TRIMMOJAR: when running on Trimmomatics on Biowulf, start with this. It essentially runs the Trimmomatic Java archive located in the folder pointed to by environmental variable \$TRIMMOJAR.
 - PE: this next piece tells Trimmomatic to expect paired end sequencing.
 - phred33: this indicates the quality score encoding, which is used by modern Illumina sequencers.
 - The input FASTQ files are specified next. Because the input FASTQ files are in the reads folder, this must be included in the path. {} in the file path acts as a place holder to store the sample ids sent by cat. All users have to do is to add the _R1 and _R2 part.
 - The outputs are specified next. These will be written to the trimmed_reads folder. Trimmed versions will be labeled with "_trimmed". The files labeled "unpaired" will store sequences where only one of the pair met the trimming threshold.
 - The adapter file (see references/illumina_multiplex.fa) is specified after the ILLUMINACLIP argument followed by some parameters to help Trimmomatic decide whether a portion of a sequence matches an adapter and continue with trimming. See the Trimmomatic manual to learn more about options and parameters (http://www.usadellab.org/cms/uploads/supplementary/Trimmomatic/TrimmomaticManual_V0.32.pdf)
 - The MINLEN argument allows users to specify a sequence length threshold after trimming. If the length of the sequence post trimming is shorter than this number than it will be discarded. The threshhold is set to 25 bases here. Short sequences may also interfere with alignment as they could be aligned to multiple spots in a genome.

cat hcc1395_sample_ids.txt | parallel -j 6 "java -jar \$TRIMMOJAR PE

Note

When running the parallel command, users will see the message below regarding citation and donation to the creators. To turn off the citation, use parallel --citation.

Academic tradition requires you to cite works you base your article on. If you use programs that use GNU Parallel to process data for an article in a scientific publication, please cite:

Tange, O. (2024, December 22). GNU Parallel 20241222 ('Bashar'). Zenodo. https://doi.org/10.5281/zenodo. 14550073

This helps funding further development; AND IT WON'T COST YOU A CENT. If you pay 10000 EUR you should feel free to use GNU Parallel without citing.

More about funding GNU Parallel and the citation notice: https://www.gnu.org/software/parallel/parallel_design.html#citation-notice

To silence this citation notice: run 'parallel --citation' once.

Come on: You have run parallel 166 times. Isn't it about time you run 'parallel --citation' once to silence the citation notice?

Run QC on Trimmed FASTQ Files

Make a directory pre_alignment_qc_trimmed to store the QC results for the adapter trimmed FASTQ files.

```
mkdir pre_alignment_qc_trimmed
```

Then load FASTQC and MultiQC.

```
module load fastqc
```

```
module load multiqc
```

In the fastqc construct below, specify the path to the trimmed FASTQ files, which are located in the folder trimmed_reads. Then use *trimmed*.fq to get FASTQC to check all of the trimmed FASTQ files (* is used as a wild card). Write the results into the pre_alignment_qc_trimmed folder using the -o option.

```
fastqc trimmed_reads/*trimmed*.fq -o pre_alignment_qc_trimmed/
```

Next, combine the FASTQC reports for the trimmed data into one using MultiQC. In the multiqc command below, specify the path the FASTQC reports (pre alignment qc trimmed) and then use --filename option to assign a base name to

the report (ie. hcc1395_multiqc, which will write a the report to the file hcc1395_multiqc.html).

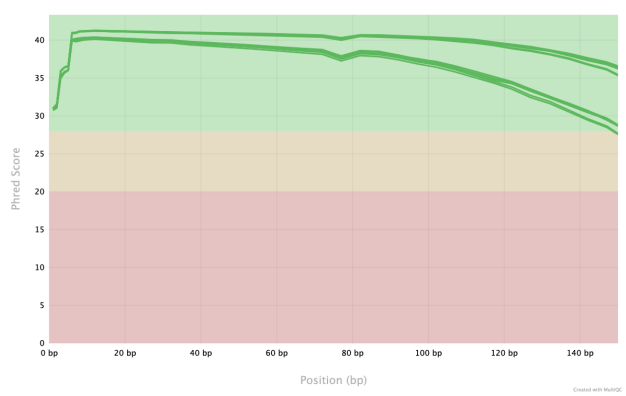
```
multiqc pre_alignment_qc_trimmed/ --filename hcc1395_multiqc
```

Then use scp to copy hcc1395 multiqc.html to local Downloads folder.

```
scp user@helix.nih.gov:/data/user/hcc1395_b4b/hcc1395_multiqc.html .
```

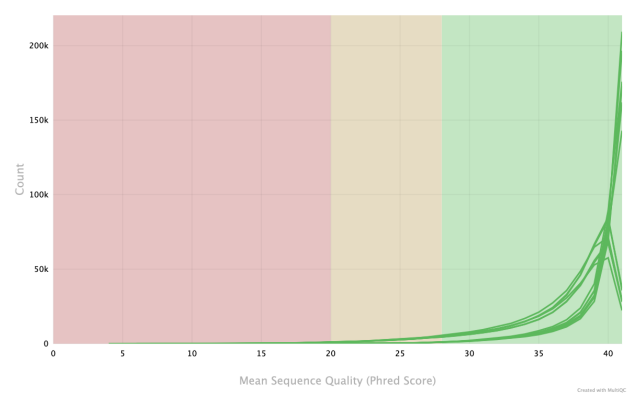
Adapter Trimming Conclusion

Adapter trimming did not influence the quality scores of the FASTQ files but sequences were cleaned of adapter contamination.

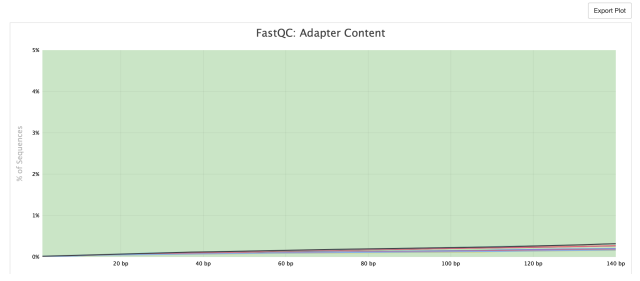


FastQC: Mean Quality Scores









View FASTQC and MultiQC HTML Reports

Adapter Trimmed FASTQ Files

- hcc1395_normal_rep1_R1
- hcc1395_normal_rep1_R2
- hcc1395_normal_rep2_R1

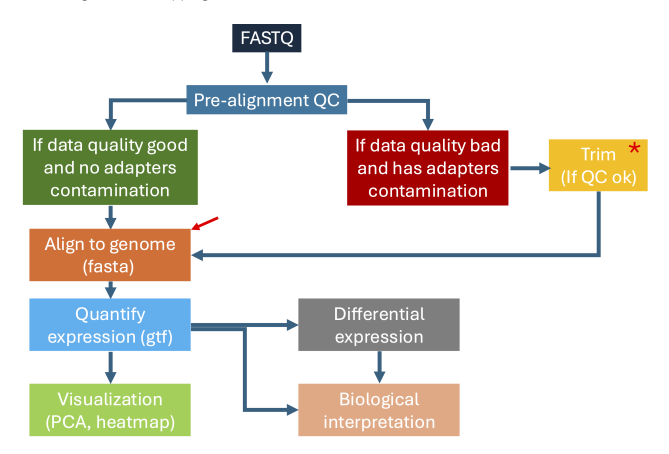
126

- hcc1395_normal_rep2_R2
- hcc1395_normal_rep3_R1
- hcc1395_normal_rep3_R2
- hcc1395_tumor_rep1_R1
- hcc1395_tumor_rep1_R2
- hcc1395_tumor_rep2_R1
- hcc1395_tumor_rep2_R2
- hcc1395_tumor_rep3_R1
- hcc1395_tumor_rep3_R2
- MultiQC

Lesson 9: Aligning FASTQ Sequences to Reference Genome

Lesson 8 Review

In Lesson 8, participants were introduced to the tool Trimmomatic, which can be used to remove low quality reads and adapter contamination from sequencing data. For the hcc1395 dataset, only adapters had to be removed. Subsequent QC of the trimmed data revealed that adapter contamination was removed. Thus, the next step will determine where in the genome each of the sequences in the trimmed FASTQ files for hcc1395 came from in a process known as the alignment or mapping.



Learning objectives

After this lesson, participants will be able to:

- Describe a reference genome.
- Understand why a splice aware aligner is needed in RNA sequencing.
- Perform alignment using the tool HISAT2 (http://daehwankimlab.github.io/hisat2/).

Sign onto Biowulf

Prior to getting started, sign onto Biowulf.

```
ssh user@biowulf.nih.gov
```

Next, change into the /data/user/hcc1395_b4b folder.

```
cd /data/user/hcc1395_b4b
```

Finally, request an interactive session with the following resources. The option --cpus-per-task is used to request 6 CPUs on Biowulf in the sinteractive command below in addition to the 12 gb of memory and 10 gb of local temporary storage.

```
sinteractive --cpus-per-task 6 --mem=12gb --gres=lscratch:10
```

The Reference Genome

The reference genome is a completely assembled sequence (ie. the order in which the nucleotides are arranged is known). For high throughput sequencing, the known sequence help scientists find out where in the genome each of the reads came from. The reference genome in a way acts like a template that researchers can follow to reconstruct the genome of the unknown. In other words, think of the reference genome as a picture of the completed puzzle that helps guide the assembly of the actual puzzle, by allowing the overlap of pieces to see where they fit in the completed version. The file 22.fa contains the reference genome for human chromosome 22.

Tip

Biowulf staff has downloaded and made the genome as well as annotations available on the cluster (see https://hpc.nih.gov/refdb/index.php)) so users generally will not need to download anything on their own.

Change into the references folder in /data/user/hcc1395_b4b for this example. Since /data/user/hcc1395 b4b should be the current working directory, just do the following.

cd references

Use the head command to view the first 10 lines of 22.fa.

head 22.fa

A FASTA (.fa) file starts with a header line that contains a ">" followed by the name or some annotation of the sequence (ie. chr22). The first 10 lines indicate unknown sequences (denoted by N's).

>chr22

Note

The header line in a FASTA file can provide more information, depending on how the sequence was curated. As an example, in the human ADSL transcript nucleotide sequence (https://www.ncbi.nlm.nih.gov/nuccore/NM_001363840.3?report=fasta), the header line provides the accession number or sequence ID (NM_001363840.3), species in which the sequence was derived (Homo sapiens), name of the gene (ADSL) where the transcript originated, and that this is a mRNA sequence.

How many lines does 22.fa contain? To find out, use the wc command with -1 option. The -1 options tells wc to count the number of lines in a file.

wc -1 22.fa

The output of wc -1 is the line number count followed by the name of the input file (ie. 22.fa). The result below indicates that 22.fa had 846976 lines.

846976 22.fa

But are there actual sequences in 22.fa (ie. A, T, C, or, G)? To find out, use the grep command with the following options for pattern searching.

- -n: instructs grep to retrieve the line number of the file where the pattern is found.
- - E: treats the pattern as an extended regular expression (https://www.google.com/search?

client=safari&rls=en&q=what+is+an+extended+regular+expression&ie=UTF-8&oe=UTF-8).

- Within the single quotes is the search pattern A, T, C, or G where each is separated by "|".
- The grep output is sent via | (pipe) to the head command to print the first 10 lines of the results.

```
grep -n -E 'A|T|C|G' 22.fa | head
```

The results of the above grep command shows that the first line with actual nucleotide sequences is 175168.

How big is the 22.fa reference genome (ie. how many bases)? To answer this, use the tool segkit and it's stats feature.

Why is the size of the genome important?

Prior to the sequencing experiment, the size of the genome will help us determine the number of reads needed to achieve a certain coverage (https://www.illumina.com/documents/products/technotes/technote_coverage_calculation.pdf (https://www.illumina.com/documents/products/technotes/technote_coverage_calculation.pdf)).

After the experiment, we could use the size of the genome along with other information to decide the computing resources (ie. time and memory) needed for analysis. Chromosome 22 is the second smallest in humans (https://medlineplus.gov/genetics/chromosome/22/), so it would be faster and require less compute power to align to this than the entire human genome, thus this dataset was chosen for the class.

```
module load seqkit
```

seqkit stat 22.fa

```
file format type num_seqs sum_len min_len avg_len 22.fa FASTA DNA 1 50,818,468 50,818,468 50,818,468 50
```

Go back up one folder to /data/user/hcc1395_b4b.

```
cd ..
```

Aligning hcc1395 Sequencing Data to Reference Genome

The tool HISAT2 will be used to align the trimmed hcc1395 FASTQ files to the human chromosome 22 reference. RNA sequencing analyses require the use of splice aware aligners in order to map sequences that span across exons.



A sequencing read (red fragments) aligning two exons (e1 and e2). Modified from: https://training.galaxyproject.org/training-material/topics/transcriptomics/tutorials/rb-rnaseq/tutorial.html (https://training.galaxyproject.org/training-material/topics/transcriptomics/tutorials/rb-rnaseq/tutorial.html)

Build an Index

Users will need to build an index of the reference genome prior to aligning using HISAT2. This step essentially breaks down the reference into manageable pieces so that the alignment process is made more efficient.

Load HISAT2.

```
module load hisat2

This is HISAT2 v 2.2.1, built for NCBI NGS 3.0.0
[+] Loading samtools 1.21 ...
```

Stay in /data/user/hcc1395 b4b for this exercise.

To build the index for HISAT2, use the hisat2-build command where the arguments are:

- Path to the reference genome FASTA (or .fa) file: the reference genome file in this case is 22.fa and it is located in the references folder so the path starting from /data/user/hcc1395_b4b is references/22.fa.
- The folder to write the indices: in this case, write the indices to the reference folder. Thus, the path starting from /data/user/hcc1395_b4b is references/22. Note that an extension is not needed as HISAT2 will append.ht to the index files.

```
hisat2-build references/22.fa references/22
```

After indexing of 22.fa has been completed, use 1s with the -1 option to view the contents of the reference folder one item per line and confirm that 8 index files with .ht extension were generated in this step.

ls -1 references/

```
22.1.ht2
22.2.ht2
22.3.ht2
22.4.ht2
22.5.ht2
22.6.ht2
22.7.ht2
22.8.ht2
```

Alignment

Make a new folder hcc1395 hisat2 to store the HISAT2 alignment results.

```
mkdir hcc1395_hisat2
```

Stay in /data/user/hcc1395 b4b for this exercise.

To perform alignment for all samples at once, the parallel command will be used. Below is the explanation of the components.

• cat hcc1395_sample_ids.txt: print the hcc1395 sample ids and | will send the output to parallel rather than printing to screen.

- The HISAT2 alignment command is enclosed within double quotes within parallel. The alignment starts with the command hisat2 and arguments and options below are include:
- - j: enables users to specify how many jobs to run in parallel (6 in this case since there are 6 samples).
- -x: prompts user to enter the path to the reference genome indice (ie. references/22, note that the ht2 extension is not needed)
- -1: prompts for the first read in the pair for paired end sequencing.
 - The data is in the trimmed reads folder.
 - {} is used as a place holder for the sample ids sent by cat.
- -2: prompts for the second read in the pair for paired end sequencing.
- -S: prompts the output SAM file path. The alignment output will be stored in the folder hcc1395 hisat2.
- --summary-file: prompts for the text (txt) file that summarizes alignment statistics. The summary will be stored in the folder hcc1395 hisat2.

```
cat hcc1395_sample_ids.txt | parallel -j 6 "hisat2 -x references/22 ·
```

After alignment completes, list the contents of hcc1395_hisat2 to see what outputs were generated.

```
ls -1 hcc1395_hisat2
```

The folder hcc1395_hisat2 contains the aligned data in .sam format and alignment summary text files (ie. those labeled with summary.txt).

```
hcc1395_normal_rep1_hisat2_summary.txt
hcc1395_normal_rep1.sam
hcc1395_normal_rep2_hisat2_summary.txt
hcc1395_normal_rep2.sam
hcc1395_normal_rep3_hisat2_summary.txt
hcc1395_normal_rep3.sam
hcc1395_tumor_rep1_hisat2_summary.txt
hcc1395_tumor_rep1.sam
hcc1395_tumor_rep2_hisat2_summary.txt
hcc1395_tumor_rep2.sam
hcc1395_tumor_rep3_sam
hcc1395_tumor_rep3_sam
```

Change into the hcc1395 hisat2 directory.

```
cd hcc1395 hisat2
```

Use cat to take a look at the alignment summary for hcc1395_normal_rep1.

```
cat hcc1395_normal_rep1_hisat2_summary.txt
```

```
331945 reads; of these:

331945 (100.00%) were paired; of these:

43449 (13.09%) aligned concordantly 0 times

283794 (85.49%) aligned concordantly exactly 1 time

4702 (1.42%) aligned concordantly >1 times

----

43449 pairs aligned concordantly 0 times; of these:

10336 (23.79%) aligned discordantly 1 time

----

33113 pairs aligned 0 times concordantly or discordantly; of these

66226 mates make up the pairs; of these:

35020 (52.88%) aligned 0 times

30758 (46.44%) aligned exactly 1 time

448 (0.68%) aligned >1 times

94.73% overall alignment rate
```

Users will see the alignment statistics shown above and reports are generated for all samples. The first line of the HISAT2 alignment statistics says of 331945 reads (100.00%) were paired. Recall from FASTQC that read 1 and read 2 FASTQ files for hcc1395_normal_rep1 each has 331945 reads after trimming. So the first line in the HISAT2 alignment statistics is saying that out of all the reads from read 1 and read 2 FASTQ files of hcc1395_normal_rep1, there are 331945 pairs. This means there are 331945x2 or 663890 reads for the hcc1395_normal_rep1 sample.

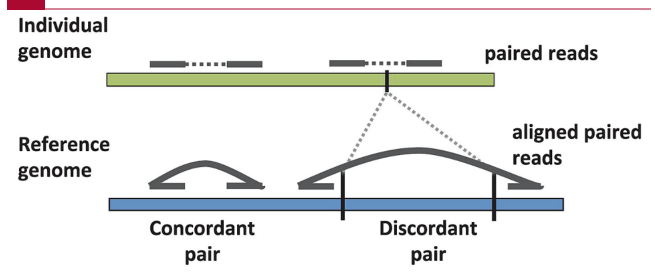


Measure	Value
Filename	hcc1395_normal_rep1_trimmed_R1.fq
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	331945
Total Bases	49.2 Mbp
Sequences flagged as poor quality	0
Sequence length	34-151
%GC	54

Basic Statistics

Measure	Value
Filename	hcc1395_normal_rep1_trimmed_R2.fq
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	331945
Total Bases	49.2 Mbp
Sequences flagged as poor quality	0
Sequence length	25-151
%GC	54

Following the first line of the HISAT2 alignment statistics, users will see some terminologies like concordant or discordant mapped reads. See the figure below for a visual explanation.



Source: Benjamin J. Raphael, Chapter 6: Structural Variation and Medical Genomics, PLOS Computational Biology, December 27, 2012 (https://journals.plos.org/ploscompbiol/article? id=10.1371/journal.pcbi.1002821)

- A concordant read pair is defined as those that align in an expected manner where the reads are oriented towards one another and the distance between the outer edges is within expected ranges.
- A discordant read pair is defined as those that do not align as expected such as where the distance between the outer edges is smaller or larger than expected.

For the hcc1395_normal_rep1 sample, there are:

- 283794 pairs (567588 reads) that aligned concordantly once
- 4702 pairs (9404 reads) that aligned concordantly more than once these are mapped to multiple parts of the genome and likely cause by duplicated or similar regions in the genome
- 10336 paris (20672 reads) that aligned discordantly once
- 30758 reads that did align (neither concordantly or discordantly) once
- 448 reads that aligned (neither concordantly or discordantly) more than once

Sum up the number of reads that mapped in the above break down and divide by the total number of reads in the two FASTQ files for the hcc1395_normal_rep1 sample to get an overall alignment rate of 94.73%.

SAM File

Aligned sequencing data are stored in the form of SAM files (https://www.google.com/url? sa=t&source=web&rct=j&opi=89978449&url=https://samtools.github.io/hts-specs/
SAMv1.pdf&ved=2ahUKEwiS2sSX8ImKAxWuF1kFHbE1NSYQFnoECBQQAQ&usg=AOvVaw0MUgu2LimUqu
The SAM file is human readable so users can use cat, less, and head to view its content in the terminal.

SAM Header

SAM files always start off with metadata information and these lines start with @. SAMTOOLS (http://www.htslib.org) can be used to view and manipulate SAM files.

module load samtools

To view the header for hcc1395_normal_rep1.sam, use the following samtools command construct, where:

- head: this argument is used to view SAM file header.
- hcc1395_normal_rep1.sam: SAM file in which the header should be viewed.

samtools head hcc1395_normal_rep1.sam

- @HD includes
- SAM file format information (in this case version 1.0 as indicated by VN:1.0)
- Whether the alignment file has been sorted (in this case no as indicated by SO:unsorted)
- @SQ provides reference information
- SN denotes reference sequence name, which is chr22
- LN is the reference length in bases, which is 50818468
- @PG provides information about the program that was used to generate the alignment
- ID is the program record identifier
- PN is the program name (HISAT2 and version 2.2.1 was used as indicated next to VN)
- CL is the command line call used to generate the alignment

After the metadata information, each SAM file contains 11 fields.

SAM File Information

```
samtools view hcc1395_normal_rep1.sam | head -1 | column -t | less -:
```

```
K00193:38:H3MYFBBXX:4:1101:10003:44458 99 chr22 31282436 60 151
```

- 1. QNAME or query template name, which is essentially the sequencing read that was mapped onto a reference genome. Use the grep to search for any template in the SAM file and users will retrieve a sequence in the corresponding FASTQ file.
- 2. The second column is a FLAG that provides detail on mapping. See https://broadinstitute.github.io/picard/explain-flags.html (https://broadinstitute.github.io/picard/

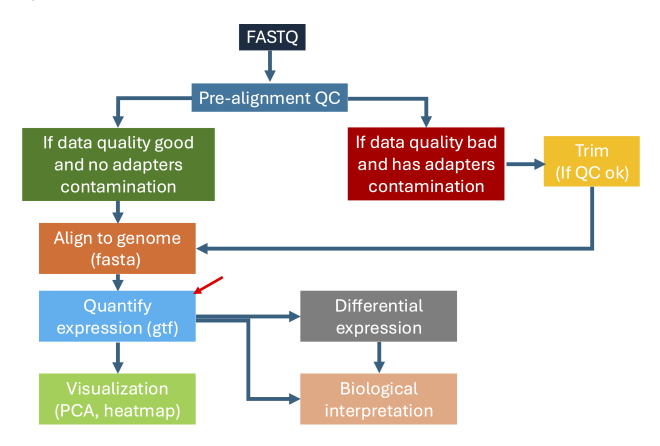
explain-flags.html) to learn about these FLAGS. For instance, the first alignment in the hcc1395_normal_rep1 SAM file has a FLAG of 99, which indicates that:

- 1. The read is properly paired.
- 2. Read is mapped in proper pair.
- 3. The mate is in the reverse strand.
- 4. The read is the first in a pair.
- 3. Column three contains the name of our reference genome (ie. chr22)
- 4. Column four provides the left most genomic coordinate where the sequencing read maps.
- 5. The mapping quality (MAPQ) is provided in the fifth column (the higher the number, the less likely that the mapping is due to error); a value of 255 in this column means that the mapping quality is not available.
- 6. Column six presents the CIGAR string, which informs about the match/mismatch, insertion/deletion, etc. of the alignment.
- 7. Column seven is the reference sequence name of the primary alignment of the NEXT read in the template. A "=" will appear if this name is the same as the current (which is expected for paired reads)
- 8. The alignment position of the next read in the template is provided in column eight. When this is not available, the value is set to 0.
- 9. Column nine provides the template length (TLEN), which should reflect the DNA fragment that was size selected for during library preparation.
- 10. The tenth column is just the sequencing read (some are written as the reverse complement so be cautious. The FLAGS in column two will tell whether the sequence is reverse complemented).
- 11. The eleventh column is the Phred quality scores of the sequencing read.
- 12. For definition of optional fields, see https://samtools.github.io/hts-specs/.

Lesson 10: Quantifying Gene Expression

Lesson 9 Review

In Lesson 9, participants learned to align the hcc1395 sequencing data to reference genome in order to determine where in the genome each of the sequences came from. The next step is to quantify at each gene, how many sequences mapped to it, which can be used to approximate expression.



Learning objectives

After this lesson, participants will be able to:

- Convert SAM files to machine readable BAM.
- Quantify gene expression from alignment results.

Sign onto Biowulf

Before getting started, sign onto Biowulf. Remember to replace user with the participant's assigned Biowulf student ID.

```
ssh user@biowulf.nih.gov
```

Then, change into the /data/user/hcc1395_b4b folder.

```
cd /data/user/hcc1395_b4b
```

Next, request an interactive session with 6 CPUs, 12 gb of RAM or memory and 10 gb of local temporary storage.

```
sinteractive --cpus-per-task 6 --mem=12gb --gres=lscratch:10
```

Converting SAM to BAM

SAM files are human readable. Many bioinformatics programs take the binary version of SAM files known as BAM for input. Below, SAMTOOLS will be used to convert the hcc1395 SAM alignment outputs to BAM.

```
module load samtools
```

Change into the hcc1395_hisat2 folder from /data/user/hcc1395_b4b and list the contents.

```
cd hcc1395_hisat2
```

```
ls
```

The following SAM alignment files should be present.

```
hcc1395_normal_rep1.sam
hcc1395_normal_rep2.sam
hcc1395_normal_rep3.sam
hcc1395_tumor_rep1.sam
hcc1395_tumor_rep2.sam
hcc1395_tumor_rep3.sam
```

The command construct below can be used to convert the SAM files to BAM and it can be explained as follows:

- cat is used to print the hcc1395 sample IDs to the terminal. The sample IDs are stored in a text file called hcc1395_sample_ids.txt one folder up, so it's path can be specified as ../hcc1395_sample_ids.txt. The output from this cat command is sent via pipe (|) to parallel. The samtools command used to convert SAM to BAM is enclosed in parentheses. Within samtools:
- sort will sort the alignment output by coordinate (see http://www.htslib.org/doc/samtools-sort.html) for more information).
- -o: prompts for the output file name, which has the .bam extension and prepended by the sample ID (specified using {} as a place holder) sent by cat.
- { } . sam is the input SAM file. Again, { } is used as a place holder for the sample IDs sent by cat.

```
cat ../hcc1395_sample_ids.txt | parallel -j 6 "samtools sort -o {}.ba
```

The next step is to index the BAM files. Again, cat and parallel will be used to performing the indexing step for all BAM files at once. The index subcommand of samtools is used for indexing. The option -b states create an index with extension .bai. The input BAM file follows the -b option and the output .bai file name is specified next.

```
cat ../hcc1395_sample_ids.txt | parallel -j 6 "samtools index -b {}.
```

Quantifying Gene Expression

Go back to /data/user/hcc1395 b4b.

```
cd /data/user/hcc1395_b4b
```

Create a directory called hcc1395_featurecounts.

```
mkdir hcc1395_featurecounts
```

Go back into /data/user/hcc1395_b4b/hcc1395_hisat2 for this exercise.

The module featureCounts (http://bioinformatics.oxfordjournals.org/cgi/reprint/btt656? ijkey=ZzPz96t2lqzAH6F&keytype=ref) from the subread (https://subread.sourceforge.net/) package will be used to quantify gene expression. So before getting started do the following.

module load subread

In the featureCounts command below:

- -p: specifies the presence of paired end data.
- --countReadPairs: specifies to count both reads in a pair.
- -a: prompt for path the gtf annotation file, which is locate one directory up from the current directory hcc1395_hisat2 (specified as . . /) and in the references folder with file name 22.gtf, so the path would be . . / references / 22.gtf.
- -g: indicates to report expression by gene name as reported in the gtf file.
- -o: prompts for path to write the output, which is one directory up from the current of hcc1395_hisat2 and in hcc1395_featurecounts with assigned file name hcc1395_gene_expression.txt. So the path for the output is ../ hcc1395_featurecounts/hcc1395_gene_expression.txt.
- *.bam: tells featureCounts to count all of the bam alignment output files in hcc1395_hisat2.

```
featureCounts -p --countReadPairs -a ../references/22.gtf -g gene_nar
```

After featureCounts has completed, change into the /data/user/hcc1395_b4b/hcc1395_featurecounts folder, which resides one folder up in /data/user/hcc1395 b4b.

```
cd ../hcc1395_featurecounts
```

head -1 reveals that the first line in the featureCounts gene expression table is the program and command line call used to perform the quantification. This can be removed in order to generate a gene by sample expression matrix.

```
head -1 hcc1395_gene_expression.txt
```

```
# Program:featureCounts v2.0.6; Command:"featureCounts" "-p" "--count
```

To remove the header in the gene expression table hcc1395_gene_expression.txt, use sed where the option 1d indicates to delete the first line in the file (ie. the header information). Then > will be use to write the output to hcc1395_gene_expression_no_header.txt.

```
sed '1d' hcc1395_gene_expression.txt > hcc1395_gene_expression_no_hea
```

Use head -1 to view the first line in hcc1395_gene_expression_no_header.txt. The expression table contains the Geneid (gene names) in the first column, followed by chromosome coordinates and then on column for each sample with the estimated expression.

```
head -1 hcc1395_gene_expression_no_header.txt
```

```
Geneid Chr Start End Strand Length hcc1395_normal_re
U2 chr22 10736171 10736283 - 113 0 (
```

The chromosomal information in columns 2 thru 6 can be removed using the cut command below where -f prompts for the columns to extract (ie. gene IDs in column 1 and sample expression estimates in columns 7 thru 12) with the construct being -f1,7-12. The file to extract these columns from (ie. hcc1395_gene_expression_no_header.txt) is specified next. Then | is used to send the output of cut to tr which will swap out the tab ('\t') that separate the columns with comma essentially turning the gene expression table in to a comma separate value (CSV) file saved under the name hcc1395_gene_expression.csv using >.

```
cut -f1,7-12 hcc1395_gene_expression_no_header.txt | tr '\t' ',' > hc
```

The column and head commands can be used to view the first two lines of hcc1395_gene_expression.csv in a way that the columns are nicely aligned. In the column command -t indicates to create a table while -s prompts for specification of column separator (ie. comma for CSV files). The file in which the content is to be displayed is hcc1395_gene_expression.csv. The output of column is sent to head with the -2 option to print the first two lines. The output indicates that this table has the Geneid (gene names) in the first column and the subsequent columns contain the expression estimates for each gene for each sample.

```
column -t -s ',' hcc1395_gene_expression.csv | head -2
```

```
Geneid hcc1395_normal_rep1.bam hcc1395_normal_rep2.bam
U2 0 0 0
```

Lesson 11: Visualizing Genomic Data: Part 1

Quick Review

The hcc1395 sequencing data were aligned to reference genome in Lesson 9. This enables the researcher to determine where in the genome each sequence from the Next Generation Sequencing experiment came from.

Learning objectives

Following this lesson, participants will know how to prepare sequencing alignment results for visual inspection using the Integrative Genomics Viewer (https://igv.org).

Signing onto Biowulf

Before getting started, sign onto Biowulf.

```
ssh user@biowulf.nih.gov
```

Next, change into the /data/user/hcc1395 b4b directory.

```
cd /data/user/hcc1395_b4b
```

Finally, request an interactive session with 6 CPUs, 12 gb of RAM or memory and 10 gb of local temporary storage.

```
sinteractive --cpus-per-task 6 --mem=12gb --gres=lscratch:10
```

Preparing Alignment Output for Visualization Using IGV

This exercise will create bigWig files from the hcc1395 BAM alignment output. bigWig files have pre-computed sequencing depth information and will enable researchers to see areas in the genome where there are sequences mapped to it and subsequently zoom in on these regions to further interrogate.

Step 1: Convert BAM to bedGraph

The first step for generating bigWig files is to convert the BAM alignment results to bedGraph (with extension bg) files that contains sequencing depth along genomic regions (ie. how many sequences mapped to a genomic region).

Definitions

BED file: this is also known as Browser Extensible Format and contains three columns, which are chromosome, start coordinate and end coordinate -- https://genome.ucsc.edu/FAQ/FAQformat.html#format1 (https://genome.ucsc.edu/FAQ/FAQformat.html#format1)

bedGraph: this has the same first three columns as the BED file but also has a fourth column that could be anything such as sequencing coverage -- https://genome.ucsc.edu/goldenPath/help/bedgraph.html (https://genome.ucsc.edu/goldenPath/help/bedgraph.html)

Example BED file is shown below -- source: https://bedtools.readthedocs.io/en/latest/content/tools/genomecov.html (https://bedtools.readthedocs.io/en/latest/content/tools/genomecov.html)

```
Chromosome Start End
chr1 10 20
chr1 20 30
chr2 0 500
```

Example bedGraph is shown below -- source: https://bedtools.readthedocs.io/en/latest/content/tools/genomecov.html (https://bedtools.readthedocs.io/en/latest/content/tools/genomecov.html)

```
Chromosome Start
                   End
                           Depth
           554304 554309
chr1
                           5
chr1
           554309 554313
chr1
           554313 554314
                          1
           554315 554316
chr1
           554316 554317
                           5
chr1
chr1
           554317 554318
                          1
chr1
           554318 554319
                           2
chr1
           554319 554321 6
chr1
           554321 554323 1
                          7
chr1
           554323 554334
```

The application bedtools (https://bedtools.readthedocs.io/en/latest/index.html) will be used to convert the hcc1395 BAM alignment outputs to bedGraph format. bedtools can be used for a range of tasks including compiling information on genomic intervals. The genomecov function

of bedtools, which calculates depth over a genomic range with the following options will be used:

- - i bam: prompts users to provide the name of the BAM input file(s).
- -split: when applied to RNA sequencing data, does not count reads that map to introns (see this post on why we get reads that map to introns in RNA sequencing (https://www.biostars.org/p/42890/)).
- -bg: reports sequencing depth along a genomic interval rather than at each nucleotide position (Figure 1 shows the ways that bedtools can report coverage information where some options report coverage along an interval, some report at each base position. bedtools also enables the fine tuning of coverage along splice junctions are reported using the split option).

To use bedtools on Biowulf, do:

```
module load bedtools
```

To generate bedGraph file for all of the hcc1395 BAM alignments in one go, the command construct below can be used.

- cat: used to print the modified sample IDs from the hcc1395 study to the terminal (recall that these new sample IDs not have "hcc1395" at the beginning). The output is sent to parallel using | or pipe.
- -j 6 in the parallel command tells it to perform 6 tasks simultaneously.
- The bedtools genomecov construct is enclosed in double quotes with parallel:
- The input BAM files are located in the folder hcc1395_hisat2 and {} is used as a place holder for the sample ID sent by cat. Thus, the path to the BAM files can be written as hcc1395_hisat2/{}.bam.
- > will write the output to the hcc1395_hisat2 folder into a file with sample name sent by cat and .bg extension. Thus, the path to the output is hcc1395_hisat2/{}.bg.

```
cat hcc1395_sample_ids.txt | parallel -j 6 "bedtools genomecov -ibam
```

Change into the hcc1395_hisat2 folder after the bedgraph files have been made and use the 1s command to ensure that that .bg files are there.

```
cd hcc1395_hisat2
```

Next, use head to view the first 10 lines of hcc1395_normal_rep1.bg.

```
head hcc1395_normal_rep1.bg
```

The last column in the .bg file is the sequencing depth.

```
chr22
       10564456
                   10564607
                               1
chr22
       10617976
                               1
                   10618127
chr22
       10684437
                   10684486
                               1
chr22
       10684486
                   10684588
                               2
       10684588
chr22
                   10684637
                              1
                               2
chr22
       10712579
                   10712719
chr22
       10712719
                   10712728
                               1
       10839950
                               2
chr22
                   10840096
chr22
       10865165
                   10865298
                               6
chr22
       10940293
                   10940443
                               1
```

Change back to the /data/user/hcc1395_b4b.

```
cd /data/user/hcc1395_b4b
```

Step 2: Create an Index for the Genome

To view genomic alignment output using IGV, an index for the reference needs to be created. samtools can be used to create the index. Recall that reference genome is located in the references folder and saved as 22.fa.

```
module load samtools
samtools faidx references/22.fa
ls -1 references
```

```
22.1.ht2
22.2.ht2
22.3.ht2
22.4.ht2
22.5.ht2
22.6.ht2
22.7.ht2
22.8.ht2
22.fa
22.fa.fai
```

22.gtf illumina_multiplex.fa

Step 3: Creating bigWig Files

The final step is to create the bigWig files using the application bedGraphToBigWig from UCSC tools (https://hpc.nih.gov/apps/Genome_Browser.html).

First, load ucsc tools into the Biowulf working environment.

module load ucsc

All of the hcc1395 .bg files can be converted to bigWig (.bw extension) at once using the command construct below. The first argument in bedGraphToBigWig is the path to the .bg files, which reside in the folder hcc1395_hisat2 (thus the path in the command is hcc1395_hisat2/{}.bg where {} is a place holder for the sample ID sent by cat). The second argument is the path to the reference index (ie. 22.fa.fai residing in the folder references, thus the path as constructed from the hcc1395_b4b folder is references/22.fa.fai). The final argument is the path to the bigWig file outputs, which will be written to the folder hcc1395_hisat2 and will have .bw extension. The for the output from the hcc1395_b4b folder will be hcc1395_hisat2/{}.bw.

cat hcc1395_sample_ids.txt | parallel -j 6 "bedGraphToBigWig hcc1395_

List the contents of hcc1395_hisat2 to confirm that the .bw files are written.

Lesson 12: Visualizing Genomic Data: Part 2

Lesson 11 Reviews

Participants learned how to prepare files (ie. bigWig) for visualizing genomic alignment results in Lesson 12.

Learning Objectives

After this lesson, participants will:

- Have a high level understanding of the Integrative Genomics Viewer (IGV) (https://igv.org) used for visualizing genomic data.
- Know the difference between visualizing genomic information from a bigWig file and BAM file.
- Understand the difference in output obtained from a splice aware versus a non-splice aware sequence aligner.

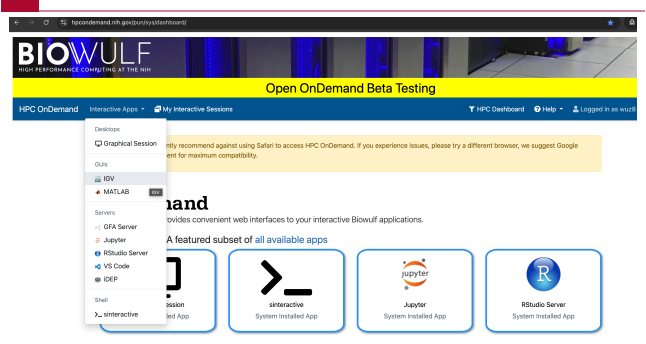
This class is demo only and not hands-on!

Launch IGV From HPC OnDemand

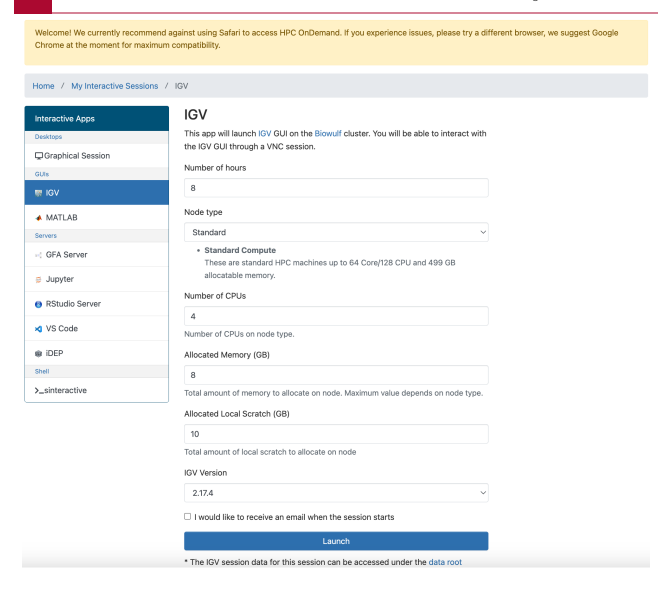
Note

Users sign onto HPC OnDemand using NIH credentials.

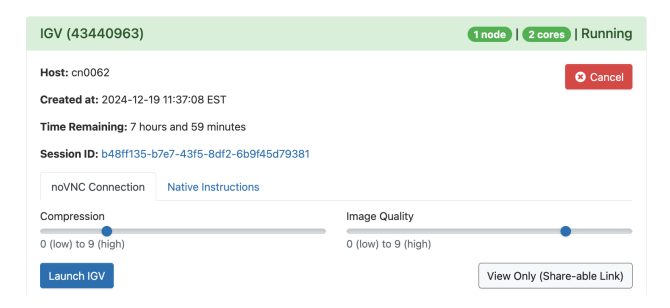
Launch the IGV session on NIH HPC OnDemand (https://hpcondemand.nih.gov/pun/sys/dashboard/) by clicking on "Interactive Apps" and then choosing "IGV".



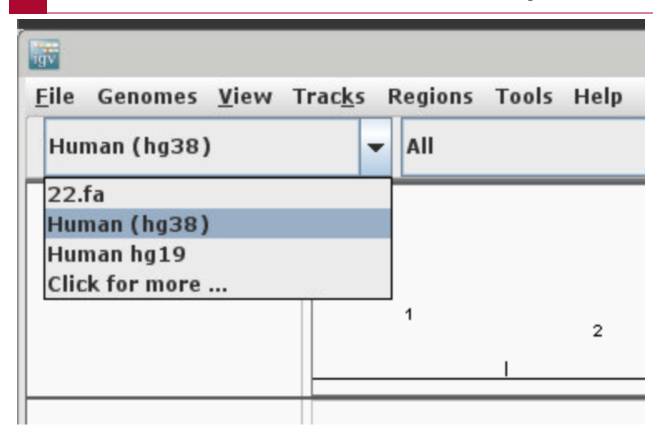
In the subsequent page, users will be able to select compute resources for the IGV session. Starting an application on HPC OnDemand will consume one of the two interactive sessions. Click on the "Launch" button when ready.



Once the IGV session's compute resources have been allocated, click on "Launch IGV" to get started.

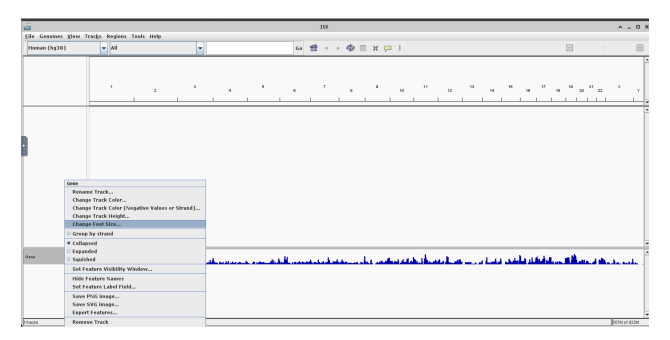


Select "Human (hg38)" as the reference in the genome selection drop down menu.



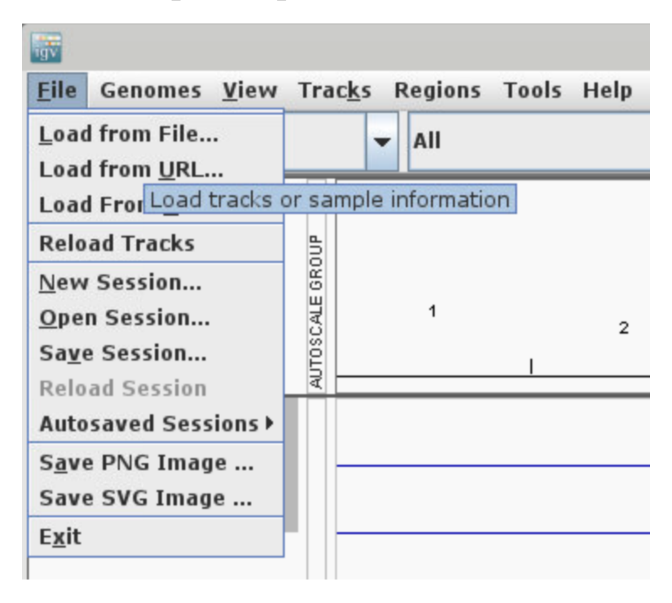
A track showing the genes for hg38 will appears. Right click on this track to see the configuration options, which include:

- Track color and font size.
- How densely the data should be displayed on the track (ie. collapsed, expanded, or squished).

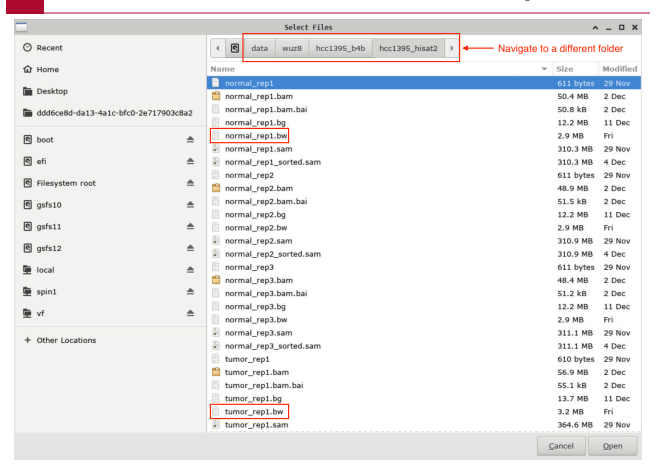


Viewing Coverage with bigWig Files

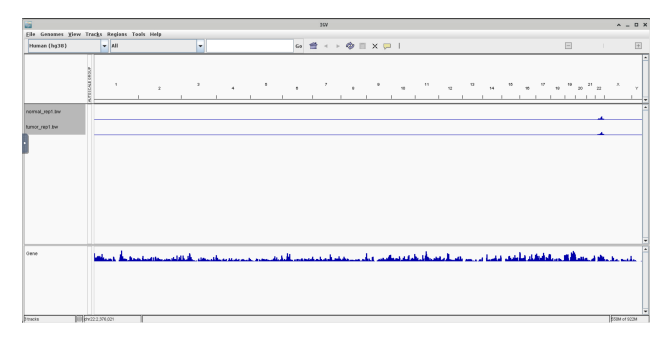
To load genomic data tracks, select "File" in the IGV menu bar. User can load from file, URL, or server. In this case, "Load from File" will be used to select alignment bigWig files from the /data/user/hcc1395_b4b/hcc1395_hisat2 folder.

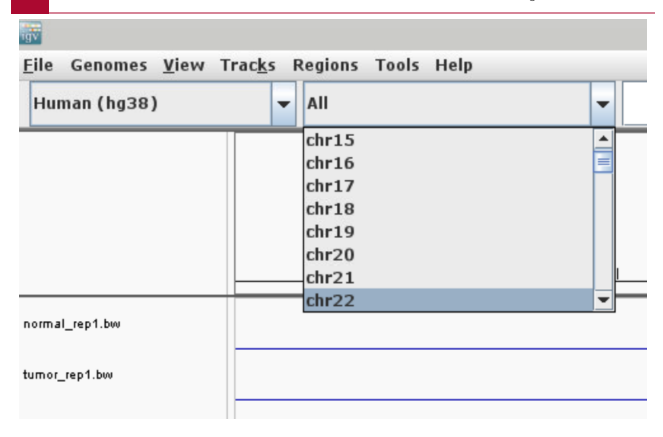


In the file explorer, choose the bigWig files normal_rep1.bw and tumor_rep1.bw (select multiple files by holding down the control key). Users can navigate to a different folder in the / data/users folder using the navigation tool at the top of the file selection window.

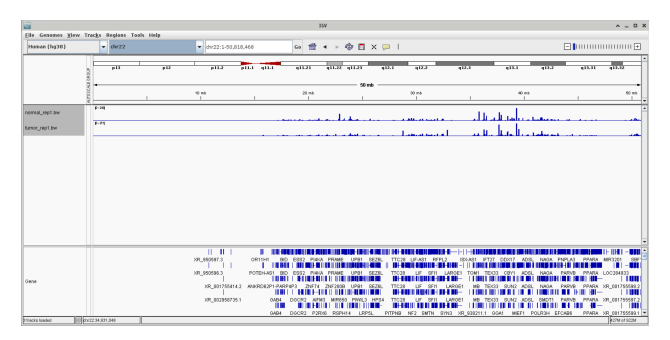


The bigWig files show pre-computed alignment coverage and it is clear that the only location where sequences have aligned is chromosome 22 as indicated by the peaks. Either click on the "22" above the peak or select from the chromosome selection drop down menu.

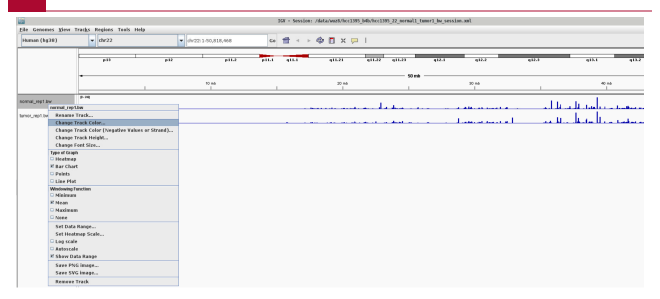




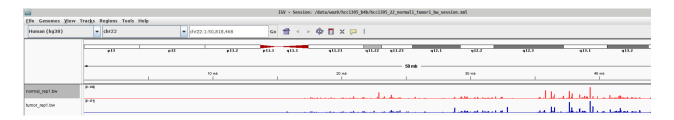
After filtering to only chromosome 22, the coverage data along with the genes on this chromosome are apparent.



Next, right click on the track labeled "normal_rep1.bw" and change the color to help distinguish is from the tumor sample (ie. tumor_rep1.bw).



Change the "normal_rep1.bw" track to red.

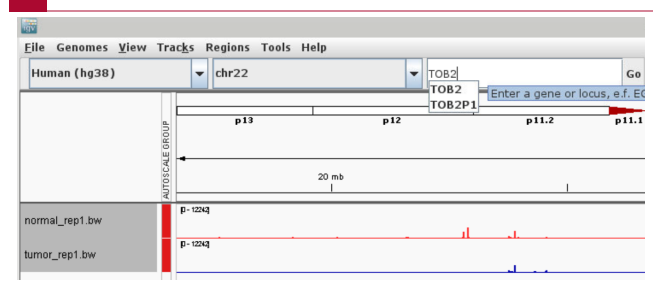


Next, select both the normal and tumor bigWig tracks, right click and select "none" for the Windowing function and Group auto scale to put the data ranges shown on the tracks on the same scale.

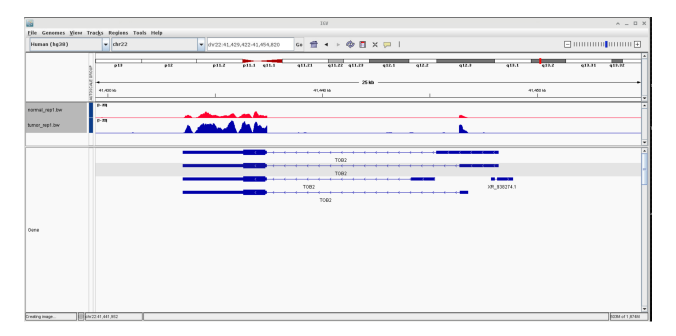
Note

Regarding the Windowing function: "When the view is zoomed out, each pixel on the screen may represent a genomic region that encompasses multiple numeric values in the data. The windowing function specifies which of the multiple values to display. To set the function, select one of the options in the Windowing Function section of the track right-click pop-up menu. The available options will depend on the file type, but most include: Minimum, Mean, Maximum, and None. By default, the function is set to Mean. The None option will display all the values, rather than combining them into one value, which can be useful for tracks displayed as points". -- IGV (https://igv.org/doc/desktop/#UserGuide/tracks/quantitative_data/#windowing-function)

Regarding group autoscale: "When comparing several sets of tracks, it is helpful to scale them on the same axis using the "Group Autoscale" option." -- https://eclipsebio.com/eblogs/how-to-use-igv-1/ (https://eclipsebio.com/eblogs/how-to-use-igv-1/)

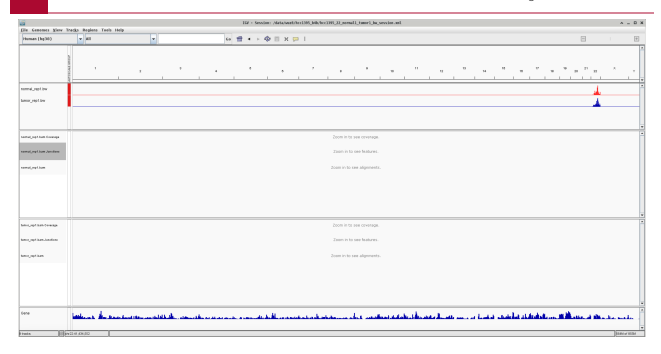


Search for the gene TOB2 (https://www.genecards.org/cgi-bin/carddisp.pl?gene=TOB2). From this IGV view, it appears that TOB2 is expressed higher in the "tumor_rep1" as compared to the "normal_rep1" sample due to more reads aligning to TOB2 in "tumor_rep1". In the expanded view of the gene track, transcript isoforms are shown. From the IGV image below, which one of the TOB2 transcripts is likely expressed?



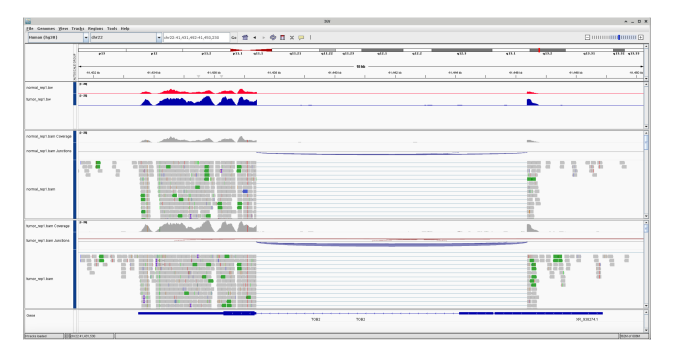
Viewing BAM files in IGV

For this exercise, click on the chromosome selection drop down and choose "All". Then load normal_rep1.bam and tumor_rep1.bam to the tracks. Unlike the bigWig files, which shows pre-calculated coverage in IGV as soon as they are loaded, BAM files requires users to zoom in to a specific location in order to see the coverage and alignment information.

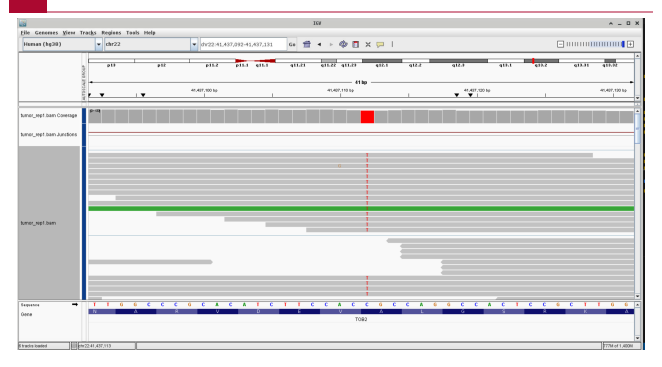


Upon zooming into TOB2, users will see that the BAM files contains more information than the bigWig file. These include:

- Coverage information (also shown in the bigWigs).
- The actual alignments.
- Splice junctions (note that the parts of sequencing reads that span across exons are connected by solid lines in the alignment track).

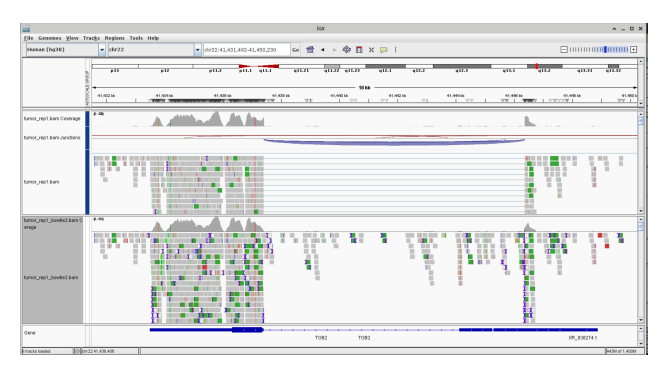


Zoom in a bit and a potential single nucleotide variant is apparent. Where the sequence contains a T, the reference contains a C. Users can also view insertions/deletions in IGV when looking at BAM files.



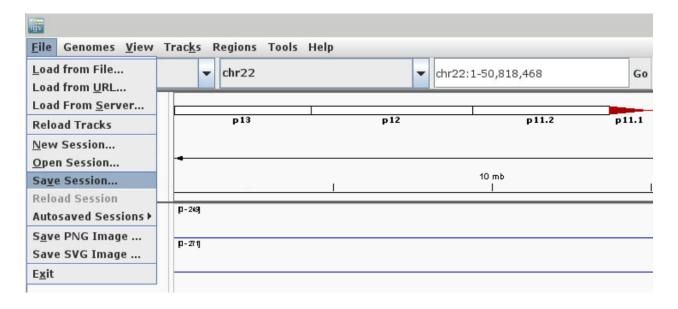
Difference Between HISAT2 and BOWTIE2 Alignment Results

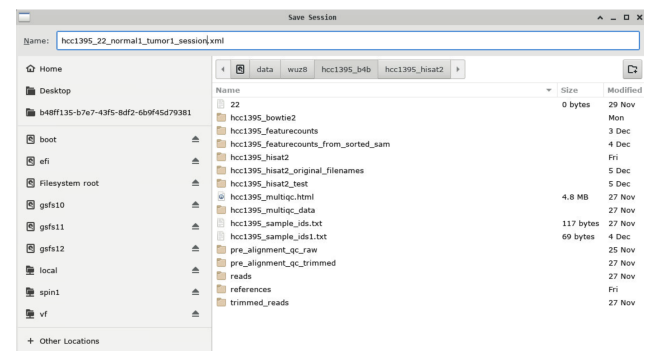
Right click on the normal_rep1.bw and tumor_rep1.bw tracks and remove them. Also remove the normal_rep1.bam track. Then, load the tumor_rep1_bowtie2.bam file into IGV. Note that the coverage and alignment information are available in the BOWTIE2 BAM outputs. However, because BOWTIE2 is not splice aware, the junction track is missing. Further, reads that map across exons are not connected by a line when the hcc1395 data is aligned to genome using BOWTIE2.



Saving IGV Session

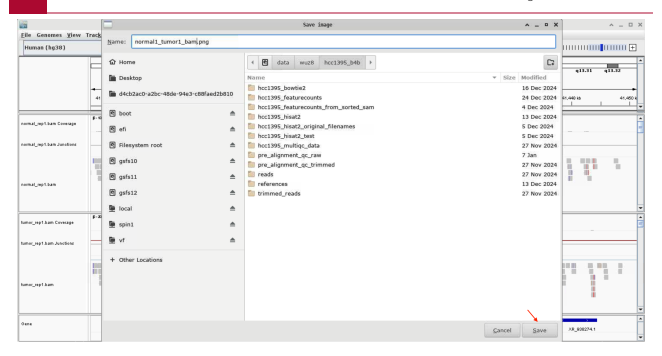
Users can save the IGV session and open it at a later time point to resume work. To do this, select File from the menu and then "Save Session". Subsequently, select the folder to save the session and provide a file name. The session is saved as a .xml file.



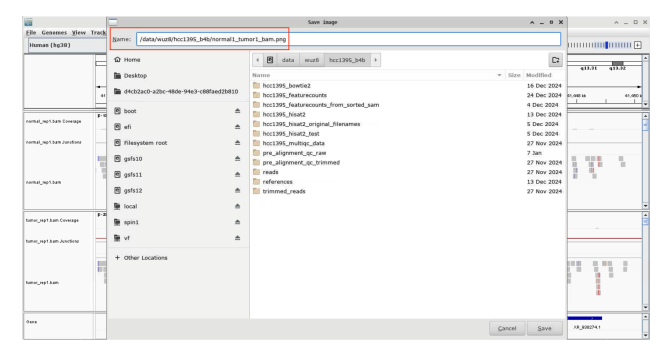


Saving IGV Image

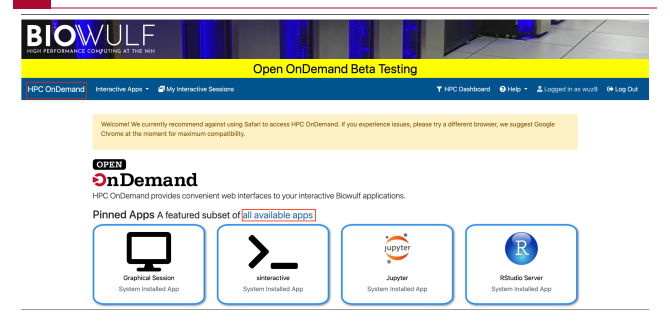
To save an IGV view as a snapshot image, goto File and Select Save PNG Image or Save SVG Image. PNG will be used here. In the dialogue box, enter the file name for the images (in this case it is normal1 tumor1 bam.png).



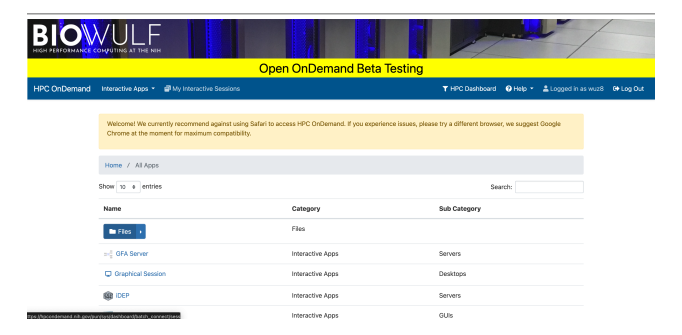
If not in the directory in which the images should be saved, just start typing the destination directory path and enter the desired file name in the address bar.



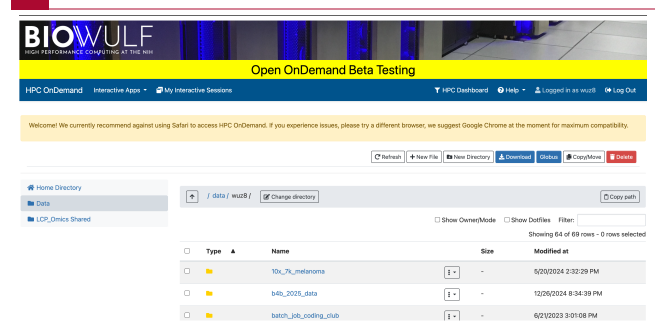
After the image has been saved, click on "HPC OnDemand" tab in the HPC OnDemand page. Then select "all available applications."



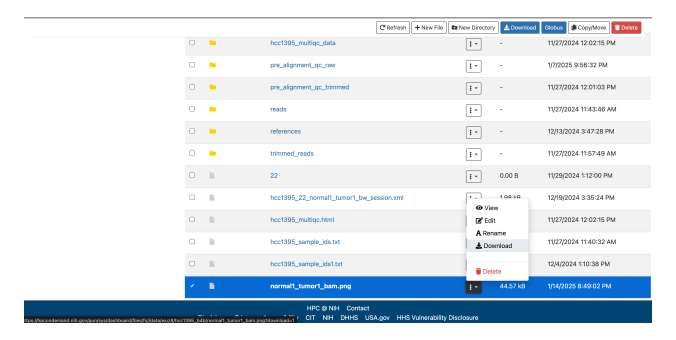
Click on "File" in the next screen.



Then choose the Data directory.



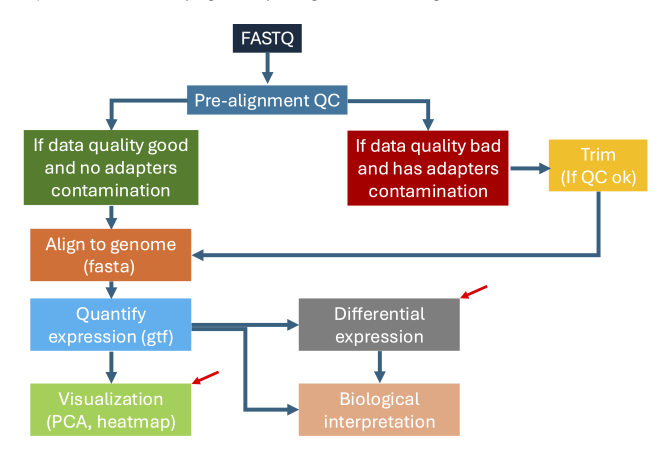
Scroll to the folder containing the IGV image, and select to download from the corresponding drop down menu.



Lesson 13: Differential Expression Analysis for Bulk RNA Sequencing: QC

Quick Review

Previously, in this Introduction to Bulk RNA Sequencing Analysis course series, participants learned to align data to reference to genome and estimate gene expression. The next 2 classes will address differential expression (DE) analysis where scientists will determine genes whose expression are statistically significantly changed between biological conditions.



Learning Objectives

After this class, participants will be able to:

- Describe rationale for filtering low expression genes and filtering these out from the gene expression estimates table.
- Perform quality control measures such as Principal Components Analysis on expression estimates and provide rationale for each QC measure.

This class will not introduce the R programming language but participants will run R scripts from the command line.

Sign onto Biowulf

Before getting started using the participant's assigned Biowulf student account ID.

```
ssh user@biowulf.nih.gov
```

Next, change into the /data/user/hcc1395 b4b folder.

```
cd /data/user/hcc1395_b4b
```

Request an interactive session with 12 gb of RAM, and 10 gb of local temporary storage.

```
sinteractive --mem=12gb --gres=lscratch:10
```

Load R

module load R

Background on R Scripts

The R scripts used for this class are in the folder b4b_scripts. Stay in /data/user/hcc1395_b4b and list the contents of b4b_scripts to learn what is available. These scripts were generated by BTEP and inspired by the Biostars Handbook's (https://www.biostarhandbook.com/).

```
ls b4b_scripts
```

There are 3 R scripts.

- deg. R will calculate differential gene expression using DESeq2.
- filter_expression.R will filter low expressing genes from the expression estimates.
- quality check. R enables QC of raw expression estimates.

```
deg.R filter_expression.R quality_check.R
```

Warning

These R scripts are for class and demonstration purposes. Do not take these and blindly use to analyze data as modification will be needed to meet the challenges of each study.

This session will use filter_expression.R and quality_check.R. To run R scripts the terminal, users will start with the RScript command follow by the script name and then arguments.

Stay in /data/user/hcc1395_b4b and create a folder called hcc1395_deg to store the differential expression analysis results.

mkdir hcc1395_deg

Note

This course will not teach participants how to script in R.

Filter Low Expressing Genes

It's common practice to filter low expression genes as these may represent noise. In these exercises, the criteria below will be used to determine which genes will be removed from the expression estimates stored in the folder hcc1395_featurecounts as file hcc1395 gene expression.csv.

- If a gene has 0 expression across all samples, then it is removed.
- If a 2 out of 3 samples in a treatment group has a gene expression of 0, then it is removed (ie. if a gene for a particular condition has 2 sample with 0 expression and 1 sample with expression of 10, then it is removed).

Filtering criteria is up to the researcher. Other methods include filtering based on the variance of expression of a gene across samples.

Issuing one of the following will pull the arguments needed for filter expression. R.

Rscript filter_expression.R

or

Rscript filter_expression.R -h

or

```
Rscript filter_expression.R --help
```

Regardless of method, users will see the following tips on how to run the R scripts, in this case filter_expression.R.

The first line is a description of the script followed by usage. The required arguments are listed as well.

```
This R script filters low expressing genes from RNA sequencing data.

Usage: Rscript filter_expression.R arguments

Use Rscript filter_expression.R -h or Rscript filter_expression.R --!

The following arguments are required.

Argument 1: File name of expression results in CSV format.

*The first column should contain gene identifiers followed by color Argument 2: The integer number of samples in each group that needs to Argument 3: File name of a phenotypes table in CSV format. This file

*Column - this contains the sample column headings from the expressample - these are the shortened sample names and will used to retain the sample are the biological conditions to which the sample argument 4: Study name! This will be prepended to the outputs.

Argument 5: Output directory! Outputs will be written here.
```

Stay in the /data/user/hcc1395 b4b folder for the exercises below.

To run filter_expression.R from the /data/user/hcc1395_b4b folder, use the following command construct where:

- b4b_scripts/filter_expression.R is the path for the filtered_expression.R script since it is being run from /data/user/hcc1395 b4b.
- hcc1395_featurecounts/hcc1395_gene_expression.csv is the path for the gene expression table as referenced from /data/user/hcc1395_b4b. The expression table is stored in the folder hcc1395_featurecounts and file hcc1395_gene_expression.csv
- hcc1395_phenotypes.csv is the path to the phenotypes table as referenced from / data/user/hcc1395_b4b.
- The study name will be hcc1395 and this will be prepended to all output files.
- The output folder is hcc1395 deg as referenced from /data/user/hcc1395 b4b.

Rscript b4b_scripts/filter_expression.R hcc1395_featurecounts/hcc139!

List the contents of hcc1395_deg from /data/user/hcc1395_b4b to check that a filterd expression CSV file is present.

```
ls hcc1395_deg
```

```
hcc1395_gene_expression_filtered.csv
```

Use wc -1 to compare number of lines in the unfiltered expression table versus the filtered.

```
wc -l hcc1395_featurecounts/hcc1395_gene_expression.csv
```

1336

```
wc -l hcc1395_deg/hcc1395_gene_expression_filtered.csv
```

593

The unfiltered expression table has 1336 lines while the filtered table has 593 lines. Thus, 743 genes were removed as a result of filtering based on the criteria set. The command expr can be used to perform arithmetic operations in Unix command line and the construct below subtracts 593 (number of lines in the filtered expression table) from the number lines in the unfiltered expression table (1336).

```
expr 1336 - 593
```

Quality Check

Below are the quality checks that will be performed on the filtered gene expression table.

• Principal Components Analysis (PCA): This transforms high dimensional data such as those derived from RNA sequencing so that researchers can see how study variables cluster together. The result of PCA is that the original data is projected onto a set of perpendicular axes where each axis accounts for a percentage of variance in the data. To learn the math behind PCA, see https://www.iro.umontreal.ca/~pift6080/H09/documents/papers/pca_tutorial.pdf). Given that this example dataset has normal and tumor samples, it is expected the samples under each condition would separate along the principal

- component axis that accounts for the most variance in the dataset. Thus, indicating it is the biology that differentiates normal and tumor samples.
- Box and density plots enable scientists to visualize the distribution of expression data and check for outliers. Ideally, the expression distribution for all samples should be roughly the same for differential expression analysis (see h3abionet (https://h3abionet.github.io/H3ABionet-SOPs/RNA-
 - Seq-4-1.html#:~:text=Whatever%20the%20shape%20of%20the,and%20extreme%20group/batch%20effects.)). These two visuals are also useful to determine whether a specific normalization method worked well.
- Distance plot shows the distance between samples. It is expected that samples within condition will be closer to each other (distance wise) than those samples from other conditions.

To run quality_check.R, use the construct below. Remember, to run a R script from the command line, start with the command Rscript, followed by:

- Script name, which is quality_check.R. However, because quality_check.R is in the folder b4b_script, it is referenced using b4b_script/quality_check.R from the current folder of /data/user/hcc1395_b4b.
- The name of expression table. Here, the filtered expression table
 hcc1395_gene_expression_filtered.csv in hcc1395_deg will be used. As
 referenced from /data/user/hcc1395_b4b, the path would be hcc1395_deg/
 hcc1395_gene_expression_filtered.csv.
- The study name (ie. hcc1395), which will be prepended to all output files.
- Directory in which to write the output (ie. hcc1395 deg).

```
Rscript b4b_scripts/quality_check.R hcc1395_deg/hcc1395_gene_express
```

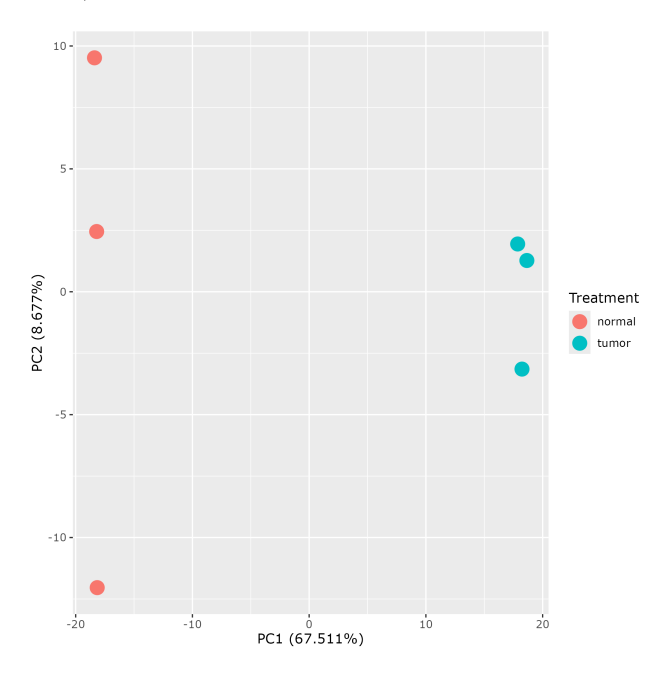
```
ls hcc1395 deg
```

```
hcc1395_box.png hcc1395_distance.png hcc1395_mehcc1395_density.png hcc1395_gene_expression_filtered.csv hcc1395_pc
```

Principal Components Results

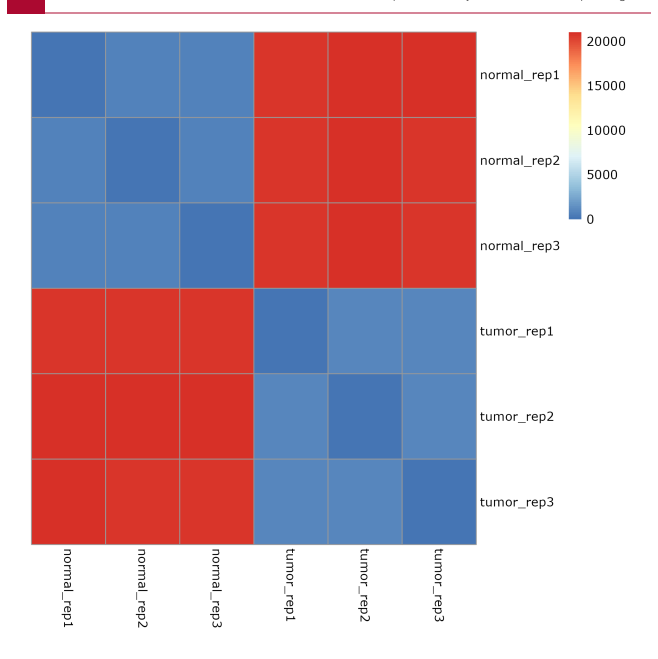
The Principal Components plot below graphs the sample along the PC1 and PC2 axes, which account for the highest and second highest variances in the data, respectively. PC1 explains 67.5% of variance while PC2 explains 8.7%. It is clear that the normal and tumor samples separate along PC1, which indicates that it is the biology that is differentiating these samples between these groups. Within group samples separation along PC2 could be due to differences

between samples of same group or maybe batch effect (although batch information is not available).



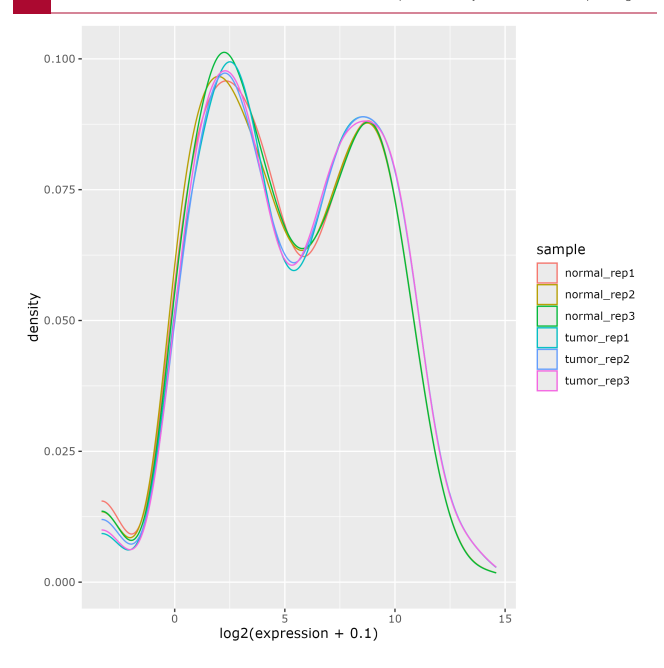
Distance Plot Results

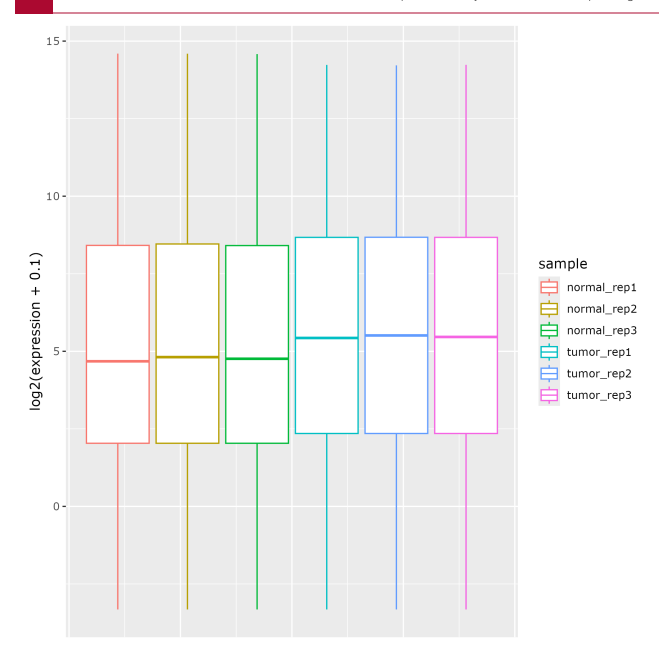
The distance plot is also a good tool for determining if the samples within groups are clustered to together. The legend scale indicates the distance and in this plot (the smaller the number the smaller the distance). It is expected that samples within the same group be closer to each other as compared to samples from a different biological condition. This QC step passes for the filtered gene expression table.



Expression Distribution

Even without normalization, the filtered expression distribution are very similar between the samples with the median and 25th to 75th percentiles higher in the tumor samples.





Lesson 14: Differential Expression Analysis for Bulk RNA Sequencing: The Actual Analysis

Lesson 13 Review

In the previous class, participants filtered gene expression results for the hcc1395 data to remove those genes whose expression is 0 across all samples and also those do not have enough replicates in a biological condition with greater than 0 expression. Subsequent QC of the filtered gene expression showed that the normal samples clustered together while tumor samples clustered with each other (PCA and distance plot), which indicates that biology is driving the underlying differences between the samples in this study.

Learning Objectives

After this lesson, participants will be able to:

- Describe on a high level how RNA sequencing data is modeled.
- Understand the importance of normalization in differential expression analysis.
- Interpret differential expression analysis results.

Sign onto Biowulf

Before getting started using the participant's assigned Biowulf student account ID.

```
ssh user@biowulf.nih.gov
```

Next, change into the /data/user/hcc1395_b4b folder.

```
cd /data/user/hcc1395_b4b
```

Request an interactive session with 12 gb of RAM, and 10 gb of local temporary storage.

```
sinteractive --mem=12gb --gres=lscratch:10
```

Load R

module load R

Run deg. R

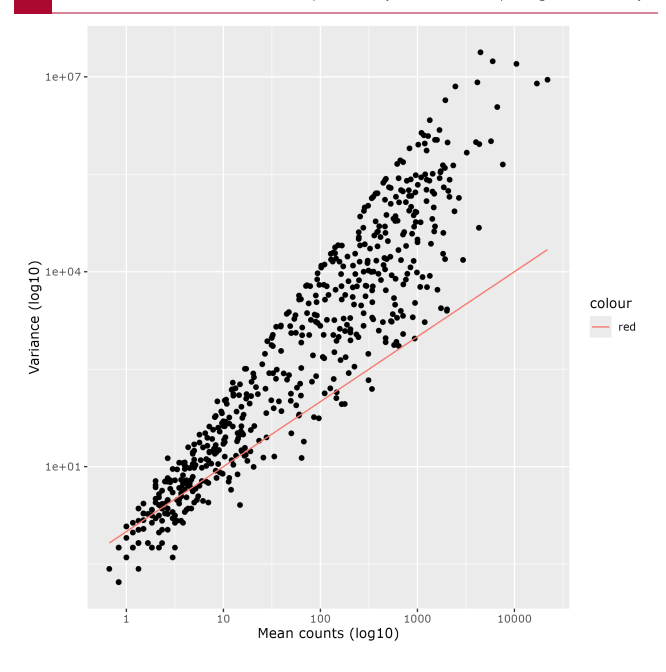
The R script for differential expression analysis, deg.R is stored in the b4b_scripts folder. It will generate differential expression results (using DESeq2), normalized expression, as well as visualizations.

A Bit of Background on the Negative Binomial Distribution

DESeq2 and another popular differential expression analysis package edgeR model expression data generated from RNA sequencing using the negative binomial distribution (https://en.wikipedia.org/wiki/Negative_binomial_distribution).

"We assume that the number of reads in sample j that are assigned to gene i can be modeled by a negative binomial (NB) distribution" -- DESeq2 paper (https://genomebiology.biomedcentral.com/articles/10.1186/gb-2010-11-10-r106) "We assume the data can be summarized into a table of counts, with rows corresponding to genes (or tags or exons or transcripts) and columns to samples. For RNA-seq experiments, these may be counts at the exon, transcript or genelevel. We model the data as negative binomial (NB) distributed" -- edgeR paper (https://academic.oup.com/bioinformatics/article/26/1/139/182458?login=true)

The rationale for modeling RNA sequencing expression data using negative binomial is that the assumption that mean equals variance under the Poisson distribution (https://en.wikipedia.org/wiki/Poisson_distribution) is not valid as variance is greater than the mean (see mean expression versus variance plot for the hcc1395 data below).



Note

The negative binomial distribution accommodates the unequal mean-variance relationship by modeling the expression data such that a both a mean and a variance parameter has to be estimated. It can be thought of that differential expression packages such as DESeq2 creates a model of and performs statistical testing for expected true value of gene expression. Statistical testing is not done directly on the input data.

Normalization

Normalization in RNA sequencing is important as this should place expression data for all samples under the same distribution. This process also removes technical variations due to sequencing depth (ie. more sequences generated from one sample as compared to another), RNA composition (ie. normal and tumor sample may differ in their composition of RNA), gene length, etc. For comparison between conditions, sequencing depth and RNA composition

should be accounted for. See here (https://hbctraining.github.io/DGE_workshop/lessons/02_DGE_count_normalization.html) for a summary of normalization methods and when it is appropriate to use them. DESeq2 uses median ratio normalization (see DESeq2 paper (https://genomebiology.biomedcentral.com/articles/10.1186/gb-2010-11-10-r106)) which normalizes for sequencing depth and RNA composition.

To learn about the math behind median ratio normalization view the Stat Quest on DESeq2 normalization (https://youtu.be/UFB993xufUU?si=rY2f_iP1BHE9p7Mu).

Note

Underneath the hood, given the raw expression table, DESeq2 will model the mean and variance of expression for each gene as well as perform normalization and calculate differential expression.

Perform Differential Expression Analysis

The script deg.R in b4b_script will be used perform differential expression analysis on the hcc1395 data. This script will use DESeq2 and takes the following as input.

- Filtered gene expression table, which is located in the folder hcc1395_deg and stored in the file hcc1395_gene_expression_filtered.csv so the path constructed from the current working directory of hcc1395_b4b is hcc1395_deg/hcc1395_gene_expression_filtered.csv.
- The phenotype information, which is located in the current working directory of hcc1395 b4b and stored in file hcc1395 phenotypes.csv.
- Experimental factor of interest for the differential expression analysis. Here, it is Treatment and the goal is to find whether there are genes whose expressions differ between normal and tumor samples.
- Study name: all output will have this prepended (in this case, the study name is hcc1395).
- Output directory for storing the results (ie. hcc1395_deg).

```
Rscript b4b_scripts/deg.R hcc1395_deg/hcc1395_gene_expression_filter
```

As DESeq2 is running, the following messages will appear and informs the user that it is performing the normalization, estimating gene expression variance, fitting the expression data to a generalized linear model and performing statistical testing.

estimating size factors estimating dispersions gene-wise dispersion estimates mean-dispersion relationship

```
final dispersion estimates fitting model and testing
```

Interpreting Differential Expression Analysis Results

Change into hcc1395_deg from hcc1395_b4b.

```
cd hcc1395_deg
```

List the contents. Use the -1 option to view directory contents 1 item per line.

```
ls -1
```

The files generated from deg.R are:

- hcc1395_deg.csv: contains the differential expression analysis results for each gene. This table is used to generate the following plots:
 - ∘ hcc1395_volcano.png
- hcc1395_filter_deg.csv: contains the results for genes whose
 - log2 Fold Change is either greater than or equal to 4.5 with adjusted p-value less than 0.01.
 - log2 Fold Change is either less than or equal to -4.5 with adjusted p-value less than 0.01.
- Normalized gene expression are stored in hcc1395_normalized_counts.csv and is used to generate quality control plots post-normalization. The plots include:
 - ∘ hcc1395_normalized_pca.png
 - hcc1395 normalized distance.png
 - ∘ hcc1395_normalized_box.png
 - hcc1395 normalized density.png
- Normalized expression for genes appearing in the filtered differential analysis results (ie. hcc1395_filtered_normalized_expression.csv), which is used to generate the expression heatmap (ie. hcc1395_expression_heatmap.png).

```
hcc1395_box.png
hcc1395_deg.csv
hcc1395_density.png
hcc1395_distance.png
hcc1395_expression_heatmap.png
hcc1395_filter_deg.csv
hcc1395_filtered_normalized_expression.csv
hcc1395_gene_expression_filtered.csv
```

```
hcc1395_mean_variance.png
hcc1395_normalized_box.png
hcc1395_normalized_counts.csv
hcc1395_normalized_density.png
hcc1395_normalized_distance.png
hcc1395_normalized_pca.png
hcc1395_pca.png
hcc1395_volcano.png
```

Take a look at differential expression analysis results stored in hcc1395_deg.csv using column, where:

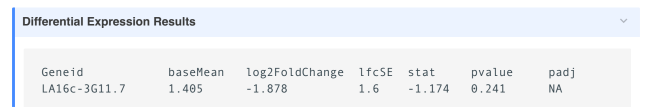
- -t tells the column command create a table.
- -s prompts for the field separator (comma in this case).
- | is used to send the output of column to head and to view the first 20 lines include the -20 option in head.

```
column -t -s ',' hcc1395_deg.csv | head -20
```

Resize Terminal or Command Prompt window if content from the differential expression analysis table does not fit the entire screen.s

The columns in hcc1395_deg.csv can be explained as follows.

- Geneid: gene identifier.
- baseMean: mean expression across all samples for the corresponding gene.
- log2FoldChange: this is a metric for gene expression change between conditions. In the results below, this is calculated by taking the ratio between mean expression of the gene from the tumor samples and that for the normal samples and then taking the log2 of this ratio. A log2 fold change of 1 indicates an expression ratio between tumor and normal of 2, while a log2 fold change of -1 indicates an expression ratio between tumor and normal of 1/2.
- IfcSE: the standard error of the log2 fold change.
- pvalue: the uncorrected p-value of the likelihood of observing the effect of the size expression change (or larger) by chance alone. This p-value is not corrected for multiple comparisons.
- padj: Multiple comparison adjusted p-value.
- normalMean: average expression for a gene in the normal samples.
- tumorMean: average expression for a gene in the tumor samples.

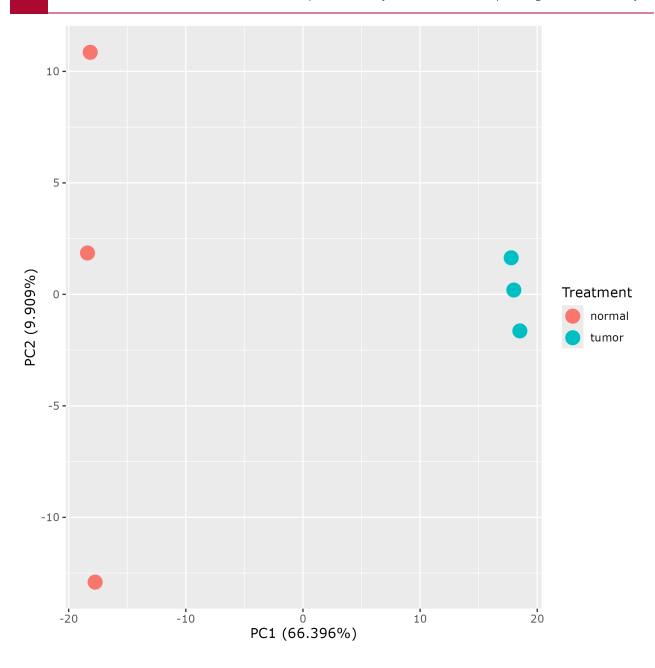


DUXAP8	556.167	8.763	0.548	15.985	1.63e-57	1.1e-56
LL22NC03-N64E9.1	65.516	6.891	0.849	8.118	4.74e-16	1.5e-15
KCNMB3P1	6.339	-1.173	0.723	-1.623	0.105	0.15
XKR3	2.034	-0.778	1.26	-0.618	0.537	0.636
AC006548.28	2.586	0.453	1.047	0.432	0.666	0.746
CECR7	85.621	0.976	0.193	5.046	4.5e-07	1.05e-06
AC006946.16	28.073	1.322	0.335	3.945	7.99e-05	0.00016
AC006946.17	2.14	-0.062	1.128	-0.055	0.956	0.967
IL17RA	281.737	-0.694	0.103	-6.728	1.72e-11	4.79e-11
CECR6	2.253	0.569	1.155	0.493	0.622	0.711
AC006946.15	1.415	1.518	1.534	0.989	0.322	NA
CECR5	401.063	-1.79	0.09	-19.786	3.9e-87	4.09e-86
CECR2	3.109	0.173	0.935	0.185	0.854	0.89
SLC25A18	4.281	-0.062	0.806	-0.077	0.939	0.958
ATP6V1E1	1012.665	-0.723	0.054	-13.426	4.26e-41	2.3e-40
BCL2L13	1133.492	-0.493	0.05	-9.784	1.32e-22	5.16e-22
BID	713.738	-2.16	0.072	-29.844	1.04e-195	2.84e-194
LINC00528	38.191	-5.972	0.757	-7.887	3.09e-15	9.58e-15

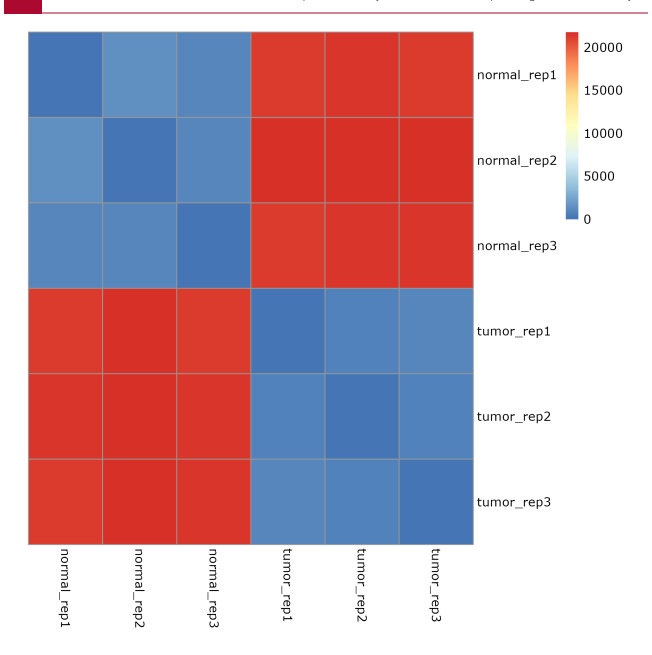
QC on Normalized Expression

It is a good idea to run QC on the normalized expression results to ensure that this step did not negatively alter the clustering of the samples and distribution of the gene expression.

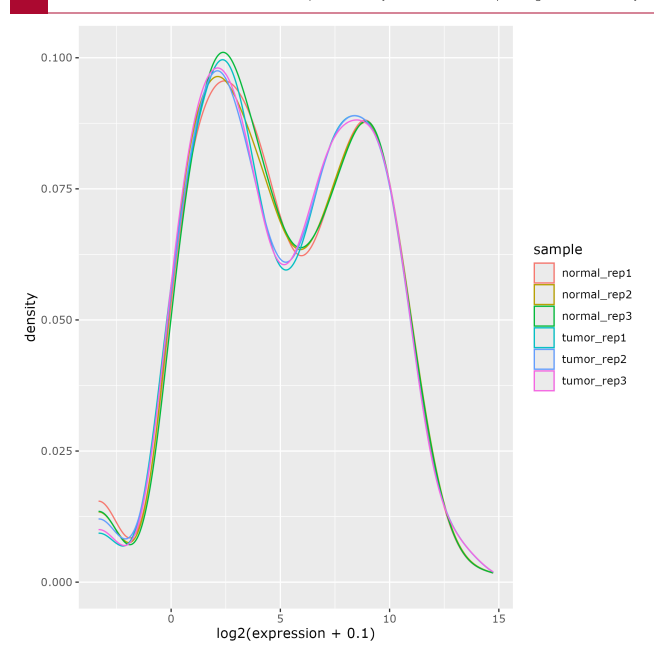
Normalization of the expression data brought the tumor samples closer together on the PCA plot.



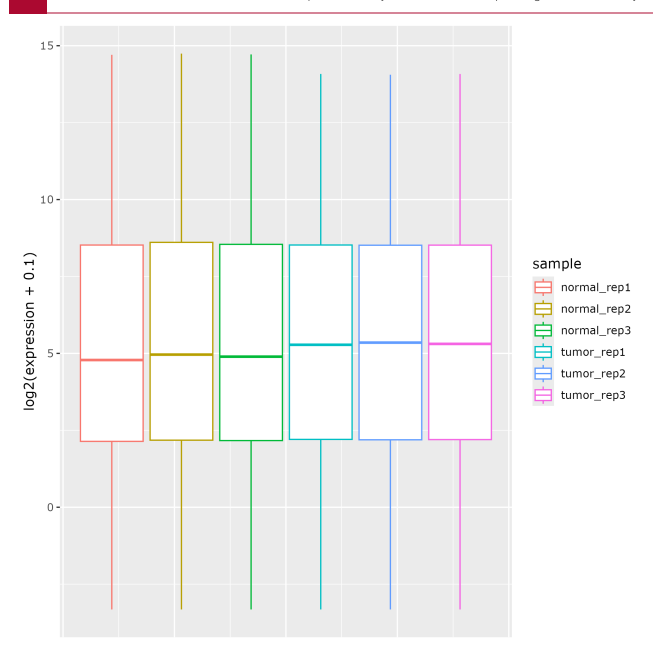
Normal samples were still closer together distance-wise as compared to the tumor samples according to the distance matrix.



Normalizing the filtered expression made the extremes (low and high expression) of the density plot overlap.



The 25th to 75th percentiles as well as median of gene expression of all samples are now more similar to each other after normalization.

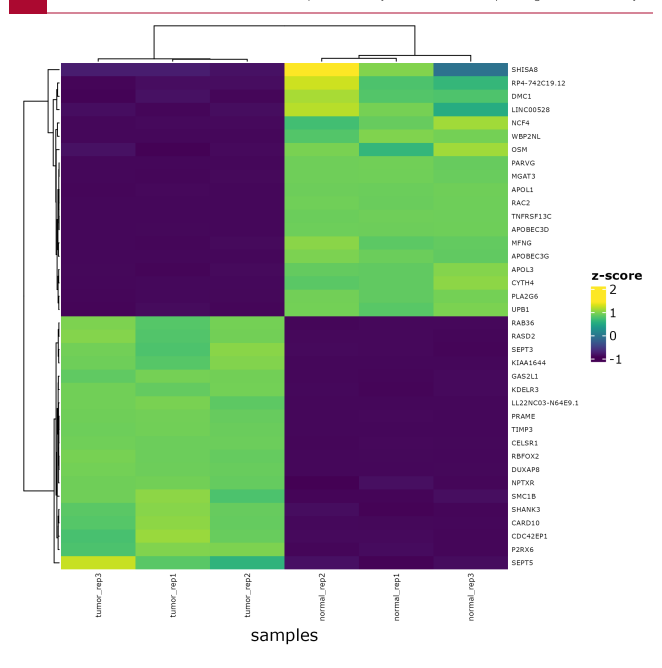


Visualizations

Expression Heatmap

A heatmap depicts numerical data on a color scale and is often used to visualize gene expression. The plot below is constructed using the normalized expression table that contains a subset of genes that were determined to be significantly changed between the normal and tumor samples according to the criteria below:

- log2 Fold Change is either greater than or equal to 2 with adjusted p-value less than 0.01
- log2 Fold Change is either less than or equal to 2 with adjusted p-value less than 0.01



Note

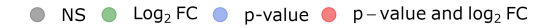
The log2 fold change and p-value cutoff for statistically significant gene expression change is determined by the researcher.

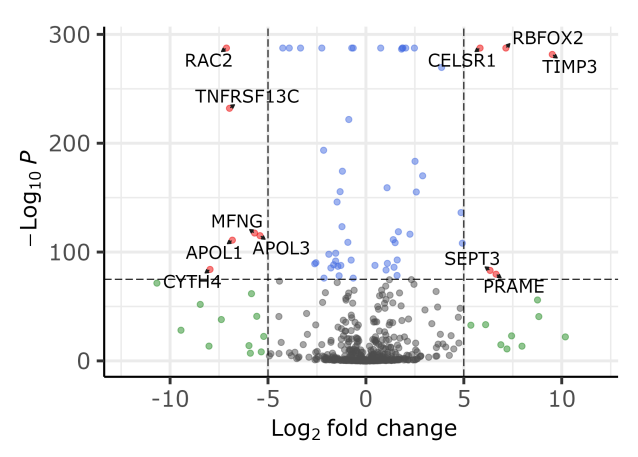
Volcano Plot

A volcano plot helps scientists visualize change and statistical confidence of change. In RNA sequencing, this is log2 Fold Change plotted on the horizontal axis and -1og10 of the p-value on the vertical axis (adjusted p-value was used here).

Volcano plot

EnhancedVolcano





total = 592 variables

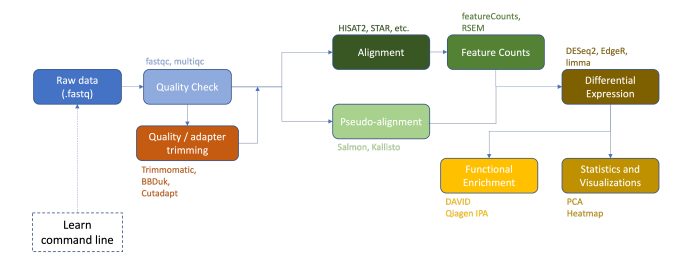
Module 3 - Pathway Analysis

Gene ontology and pathway analysis

Objectives

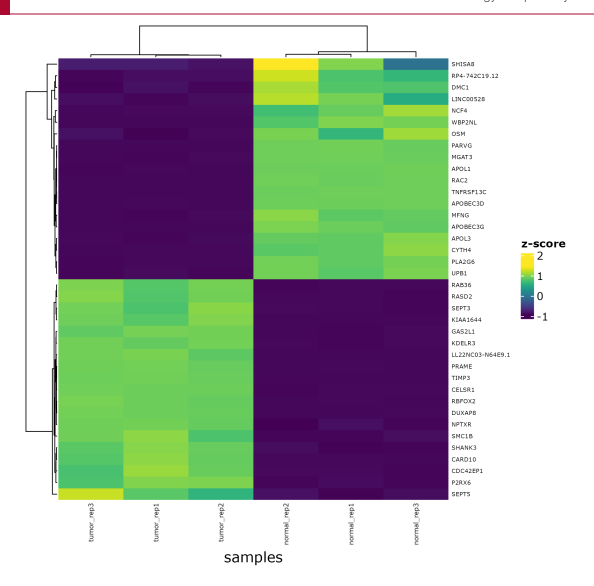
- 1. Determine potential next steps following differential expression analysis.
- 2. Tour geneontology.org and understand the three main ontologies.
- 3. Learn about different methods and tools related to functional enrichment and pathway analysis.
- 4. Get familiar with databases commonly used by popular functional enrichment tools.

Where have we been and where are we going?



Thus far we have:

- 1. Downloaded raw RNA-Seq data (.fastq files).
- 2. Examined raw data quality using fastqc and multiqc.
- 3. Performed adapter and quality trimming using Trimmomatic.
- 4. Aligned the raw sequences to a reference genome (human chromosome 22 from the GRCh38 version of the human reference genome) using HISAT2.
- 5. Viewed and compared alignments using IGV.
- 6. Generated a gene count matrix using featureCounts.
- 7. Performed differential expression analysis (DESeq2).
- 8. Generated a heatmap of differentially expressed genes.



Heatmap of differentially expressed genes (Normal vs Tumor).

You now have a potentially large list of differentially expressed genes. Now what? If you are like most biologists, you are interested in understanding these genes within their biological context.

To do that, we can examine gene ontology and perform some type of functional enrichment analysis or pathway analysis.

These types of analyses exploit the use of gene sets, and not all gene sets represent a pathway. Gene sets are collections of genes "formed on the basis of shared biological or functional properties as defined by a reference knowledge base. Knowledge bases are database collections of molecular knowledge which may include molecular interactions, regulation, molecular product(s) and even phenotype associations" Mathur et al. 2018 (https://biodatamining.biomedcentral.com/articles/10.1186/s13040-018-0166-8).

Whereas, a pathway is not a simple list of genes but rather includes an interaction component usually related to a specific mechanism, process, etc.

What is gene ontology?

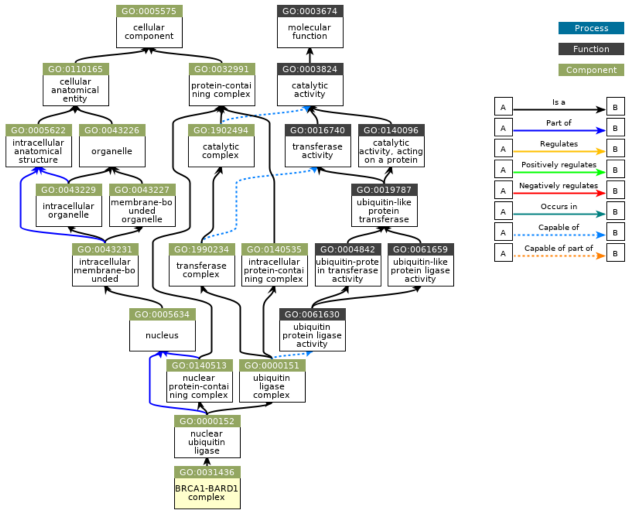
Many of the tools used to understand functional enrichment will use sets of GO terms, examining GO enrichment. What do we mean by GO?

The Gene Ontology (GO) provides a framework and set of concepts for describing the functions of gene products from all organisms. --- https://www.ebi.ac.uk/ols/ontologies/go (https://www.ebi.ac.uk/ols/ontologies/go).

This is manually curated by team members of the GO consortium.

There are two parts to the gene ontology: (Check out https://www.youtube.com/watch?v=6Am2VMbyTm4 (https://www.youtube.com/watch?v=6Am2VMbyTm4) for a more detailed overview)

- 1. the ontology (the GO terms and their hierarchical relationship) form a directed, acyclic graph structure (nodes = GO terms, edges = relationships)
- 2. the annotations (the annotated genes linked to various GO terms)



QuickGO - https://www.ebi.ac.uk/QuickGO

Image from https://www.ebi.ac.uk/QuickGO/GTerm?id=GO:0031436 (https://www.ebi.ac.uk/QuickGO/GTerm?id=GO:0031436) .

What is a GO term?

- GO terms provide information about a gene product
- GO terms as a vocabulary are species agnostic, but there are species constraints
- ontology and annotations are updated regularly
- computer readable suitable for bioinformatics

GO integrates information about gene product function in the context of three domains:

- 1. molecular function (F) "the molecular activities of individual gene products" (e.g., kinase)
- 2. cellular component (C) "where the gene products are active" (e.g., mitochondria)
- 3. *biological process (P) "the pathways and larger processes to which that gene product's activity contributes " (e.g., transport)

What is GO Slim? Reducing Semantic Similarity

There is a lot of redundancy in pathway analysis results, and there are different tools that can be used for reducing this redundancy (e.g., DAVID's Functional Annotation Clustering, REVIGO, GOSemSim). Alternatively, users can focus on sets of terms at different levels in the GO hierarchy, which range from broad to more specific. There are also GO Slim terms (https://www.ebi.ac.uk/training/online/courses/goa-and-quickgo-quick-tour/what-is-quickgo/what-can-i-do-with-a-go-slim/), which are simplified subsets from GO. GO slim terms are useful for providing a high level summary of functions.

Checkout geneontology.org to learn more about GO.

Approaches to gene set analysis / pathway analysis?

Functional enrichment and pathway analysis have broad and varying definitions. For our purposes, there are three general approaches: 1. Over-Representation Analysis (ORA), 2. Functional Class Scoring (FCS), and 3. Pathway Topology (PT) (Khatri et al. 2012 (https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi.1002375#pcbi-1002375-t001)).

Examining genes in a set allows us to:

- 1. increase the statistical power in our analysis
- 2. ease interpretation
- 3. predict new roles for genes
- 4. better integrate data from different methods

^{*}Commonly used for pathway enrichment analysis

Over-representation analysis (ORA)

statistically evaluates the fraction of genes in a particular pathway found among the set of genes showing changes in expression --- Khatri et al. 2012 (https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi. 1002375#pcbi-1002375-t001)

From this, ORA determines which pathways are over or under represented by asking "are there more annotations in the gene list than expected?"

Things to know:

- Tests based on hypergeometric, fisher's exact, chi-square, or binomial distribution; these determine the probability that the number of genes in our gene list and found in a given gene set are observed by chance.
- Assumes independence between genes.
- Requires an appropriate background gene set for comparison.
 - This could be:
 - all genes in the organism of interest
 - all protein coding genes
 - only the genes that are measured (microarray)
 - only the genes that are expressed (RNA-Seq) (https:// genomebiology.biomedcentral.com/articles/10.1186/s13059-015-0761-7)
 - genes in a gene set collection
- Prioritizes a subset of genes using an arbitrary, user determined threshold.
- Doesn't require the data, just the gene identifiers.
- Example tools include DAVID and Qiagen IPA.

To understand the statistics behind ORA, see this guide (https://www.pathwaycommons.org/guide/primers/statistics/fishers_exact_test/) from Pathway Commons and this video (https://www.youtube.com/watch?

v=H1cUs6pql9s&list=PLRGSLMDpRxCrTybN8g8GjNk39lmzJ2UTF&index=2) from Biostatsquid.

Functional Class Scoring (FCS)

Includes 'gene set scoring' methods such as GSEA, which first compute DE scores for all genes measured, and subsequently compute gene set scores by aggregating the scores of contained genes. --- Geistlinger et al. 2021 (https://academic.oup.com/bib/article/22/1/545/5722384?login=true)

These methods are more sensitive than ORA methods, but are more challenging to implement.

GSEA

• Ignores gene position and role

- Does not pre-select genes; considers all gene expression in the form of a ranked list. You must include data with gene identifiers for ranking.
 - Ranking by magnitude of change in gene expression between conditions and pvalue.
 - Top = upregulated + significant
 - Bottom = down-regulated + significant
 - Middle = non-significant
 - Determines where genes from a gene set fall in the ranking (i.e., at the top of the list or the bottom of the list).
 - Creates a running sum statistic
 - uses a permutational approach to determine significance (e.g., Kolmogorov-Smirnov)
- Broad Institute software but also available using web-based tools, R (See fgsea for a modified approach), and Qlucore (proprietary).
- also considered a strategy encompassing a range of methods
 - self-contained methods vs competitive methods

Check out this video for a nice overview of GSEA (https://www.youtube.com/watch? v=egO7Lt92gDY).

What is MSigDB (https://www.gsea-msigdb.org/gsea/msigdb/index.jsp) and how does it relate to GSEA?

The Molecular Signatures Database (MSigDB) is a curated resource of thousands of gene sets by the Broad Institute. These sets were curated for use with GSEA software but are used with other tools as well.

- Includes both human and mouse collections.
- There are 34,837 gene sets in the Human Molecular Signatures Database (MSigDB) (not all gene sets are related to pathways).
 - Includes 9 larger, themed collections

Human Collections hallmark gene sets are coherently ontology gene sets consist of genes expressed signatures derived by aggregating annotated by the same ontology term. many MSigDB gene sets to represent welldefined biological states or processes. positional gene sets corresponding to oncogenic signature gene sets defined human chromosome cytogenetic bands. directly from microarray gene expression data from cancer gene perturbations. **curated gene sets** from online pathway immunologic signature gene databases, publications in PubMed, and sets represent cell states and perturbations knowledge of domain experts. within the immune system. regulatory target gene sets based on cell type signature gene sets curated gene target predictions for microRNA seed from cluster markers identified in single-cell sequencing studies of human tissue. sequences and predicted transcription factor binding sites. computational gene sets defined by mining large collections of cancer-oriented expression data.

- Highlighted examples:
 - C5 the gene ontology (GO) collection
 - C2 curated gene sets from publications and pathway databases (e.g., KEGG and REACTOME).
 - Hallmark collection a summary list of well-defined gene sets with decreased redundancy. A good choice for many studies.
 - C7 great for immunological research.

Pathway Topology

ORA and FCS discard a large amount of information. These methods use gene sets, and even if the gene sets represent specific pathways, structural information such as gene product interactions, positions of genes, and types of genes is completely ignored. Pathway topology methods seek to rectify this problem.

PT methods are mostly considered network based.

Some examples:

Impact analysis (iPathwayGuide)

constructs a mathematical model that captures the entire topology of the pathway and uses it to calculate a perturbation for each gene. Then, these gene perturbations are combined into a total perturbation for the entire pathway and a p-value is calculated by comparing the observed value with what is expected by

chance. (https://advaitabio.com/ipathwayguide/more-accurate-pathway-rankings-using-impact-analysis-instead-of-enrichment/)

Other tools include Pathway-Express, SPIA, NetGSA, TPEA, etc. (See Nguyen et al. 2019 (https://genomebiology.biomedcentral.com/articles/10.1186/s13059-019-1790-4) for a review of PT methods.)

These methods have shown to produce more accurate results when the user wishes to understand the types and directions of gene interactions and underlying mechanisms. However, such methods often "require experimental evidence for pathway structures and genegene interactions, which is largely unavailable for many organisms." (https://www.sciencedirect.com/science/article/pii/S0168952523000185?via%3Dihub#bb0055)

What tools are available?

This is neither a comprehensive list of tools nor an endorsement of certain tools, but rather a list of semi-popular tools with different approaches.

Note

There are a ton of tools out there. Be aware of the background methods used and the quality of results returned. Also, check to see when the tool was last updated.

Gene set analysis tools

Tool	Author	Year	Citations ¹	Availability	Gene sets	$Methods^2$
WEBGESTALT	Zhang et al. [73]	2005	1423	Web server	GO, KEGG, +20 more	ORA, GSEA
GOSTATS	Falcon and Gentleman [74]	2007	1437	R package	GO	ORA
G:PROFILER	Reimand et al. [75]	2007	534	Web server	GO, KEGG, +7 more	ORA
GENETRAIL	Backes et al. [76]	2007	360	Web server	GO, KEGG, +28 more	ORA, GSEA
DAVID	Huang et al. [8]	2009	19 569	Web server	GO, KEGG, +38 more	ORA
GORILLA	Eden <i>et al</i> . [77]	2009	1881	Web server	GO	ORA
TOPPGENE	Chen <i>et al</i> . [78]	2009	1200	Web server	GO, KEGG, +45 more	ORA
CLUSTER-PROFILER	Yu <i>et al</i> . [10]	2012	1305	R package	GO, KEGG, +8 more	ORA, GSEA
PANTHER	Mi <i>et al</i> . [79]	2013	1405	Web server	GO, +2 more	ORA, GSEA
ENRICHR	Chen <i>et al</i> . [9]	2013	1246	Web server	GO, KEGG, +33 more	ORA

Google Scholar, July 2019.

Table from Geistlinger et al. 2020 (https://academic.oup.com/bib/article/22/1/545/5722384)

²Detailed summary of implemented methods in Supplementary Methods S1.2.

Other and related tools

- Gene Set Enrichment Analysis (GSEA) (http://software.broadinstitute.org/gsea)
- EnrichmentMap (https://apps.cytoscape.org/apps/enrichmentmap)
- REVIGO (http://revigo.irb.hr/) (reducing and visualizing gene ontology)
- Pathview (https://pathview.uncc.edu/)
- iPathwayGuide (https://advaitabio.com/ipathwayguide/) (proprietary)
- Qiagen IPA (https://btep.ccr.cancer.gov/docs/resources-for-bioinformatics/software/ipa/) (proprietary, CCR license)
- Qlucore (https://btep.ccr.cancer.gov/docs/resources-for-bioinformatics/software/qlucore/)
 (proprietary, CCR license)
- GeneMANIA (https://apps.cytoscape.org/apps/genemania)
- CellNetAnalyzer (http://www2.mpi-magdeburg.mpg.de/projects/cna/cna.html)
- PARADIGM (http://paradigm.five3genomics.com)

Other databases

There are many databases devoted to relating genes and gene products to pathways, processes, and other phenomenon. Again, the following is not meant to be a comprehensive list.

Kyoto Encyclopedia of Genes and Genomes (KEGG) (https://www.genome.jp/kegg/kegg1a.html)

- Curated database
- Biological pathways
- Molecular interaction networks
- Includes very nice pathway maps metabolic pathways, disease pathways, drug-target interactions, etc.
- System-level integration
- Restricted licenses

Reactome (https://reactome.org/)

The Reactome Knowledgebase systematically links human proteins to their molecular functions, providing a resource that functions both as an archive of biological processes and as a tool for discovering novel functional relationships in data such as gene expression studies or catalogs of somatic mutations in tumor cells. --- Jassal et al. 2019 (https://academic.oup.com/nar/article/48/D1/D498/5613674)

- Curated database including metabolism, signaling, and other biological processes
- Human specific
- Also includes disease super-pathways
- Several built-in pathway analysis tools

Pathway Commons (https://www.pathwaycommons.org/)

- A meta-database of pathways from other pathway databases
- Standardized format

PANTHER (http://www.pantherdb.org/pathway/)

The PANTHER (Protein ANalysis Through Evolutionary Relationships) Classification System was designed to classify proteins (and their genes) in order to facilitate high-throughput analysis. The core of PANTHER is a comprehensive, annotated "library" of gene family phylogenetic trees. --- pantherdb.org/about.jsp (http://pantherdb.org/about.jsp)

• Especially useful when considering evolutionary relationships

WikiPathways (https://www.wikipathways.org/index.php/WikiPathways)

- Community driven meta-database of pathways
- A great source of less well-established pathways

NDEx (https://home.ndexbio.org/index/)

The NDEx Project provides an open-source framework where scientists and organizations can store, share, manipulate, and publish biological network knowledge. - ndexbio.org (https://home.ndexbio.org/about-ndex/)

HumanCyc (https://humancyc.org/) (See BioCyc)

HumanCyc provides an encyclopedic reference on human metabolic pathways, the human genome, and human metabolites. --- humancyc.org (https://humancyc.org/)

DisGeNET (https://disgenet.com/)

A knowledgebase of genes and associated diseases.

Pathguide (http://pathguide.org/?organisms=all&availability=all&standards=all&order=alphabetic&DBID=none)

- Looking for a specific database? Pathguide contains a resource list of pathways searchable by organism and resource type.
- Most recent update was 2017

Importance of Gene IDs

To use various tools for functional analysis, you will need a list of annotated genes. Gene annotations come in a variety of flavors and not all flavors are compatible with every tool. For example, Gene Ontology (GO) is associated with Entrez, Ensemble, and offical gene symbols (assigned by the HUGO Gene Nomenclature Committee (HGNC)).

Note: Genome builds will have differences in the names and coordinates of genomic features, which will impact gene ID conversions. See this tutorial (https://github.com/hbctraining/Training-modules/blob/master/DGE-functional-analysis/lessons/Genomic_annotations.md) from the Harvard Chan Bioinformatics Core.

Some tools to help with annotation / conversion:

- g:Convert (https://biit.cs.ut.ee/gprofiler/convert)
- Ensembl Biomart
- AnnotationHub (https://bioconductor.org/packages/release/bioc/html/AnnotationHub.html)
- DAVID Gene ID Conversion (https://davidbioinformatics.nih.gov/conversion.jsp)

Other Considerations

- Describe what method(s) you use for functional enrichment or pathway analysis clearly in any resulting publications. This should include parameters used and software versions.
- For ORA, select appropriate background genes.
- Use p-values corrected for multiple comparisons. If more than one gene set is tested, the p-values should be corrected to minimize false positive.
- Many of the classic methods developed within the above approaches were designed specifically for transcriptomic data. Other types of -omic data (e.g., proteomics, metabolomics, scRNA-seq, GWAS) differ intrinsically from transcriptomic data, and these differences should be considered when selecting a method. For methods/tools specific to proteomics, GWAS, epigenomics, and multi-omics, see Zhao and Rhee 2023 (https://www.sciencedirect.com/science/article/pii/S0168952523000185?via%3Dihub#bb0055).

Resources:

- ClusterProfiler tutorial (https://bioinformatics.ccr.cancer.gov/docs/btep-coding-club/ CC2023/FunctionalEnrich_clusterProfiler/)
- Functional enrichment and comparison with R (https://alexslemonade.github.io/refinebio-examples/03-rnaseq/pathway-analysis_rnaseq_01_ora.html).
- ClusterProfiler, pathview, and good introductory information (https://github.com/ hbctraining/Training-modules/blob/master/DGE-functional-analysis/lessons/ functional_analysis_2019.md)
- Article on the impact of the evolving GO (https://www.nature.com/articles/ s41598-018-23395-2)
- Ten Years of Pathway Analysis: Current Approaches and Outstanding Challenges, PLOS Computation Biology, 2012 (https://journals.plos.org/ploscompbiol/article?id=10.1371/ journal.pcbi.1002375)
- Pathway enrichment analysis and visualization of omics data using g:Profiler, GSEA,
 Cytoscape and EnrichmentMap (https://www.nature.com/articles/s41596-018-0103-9)
- Toward a gold standard for benchmarking gene set enrichment analysis, Briefings in Bioinformatics, 2021 (https://academic.oup.com/bib/article/22/1/545/5722384)

• Introductory lectures on functional enrichment and R from DIY Transcriptomics (https://diytranscriptomics.com/project/lecture-10)

Help Sessions

201 Lesson 2 Practice

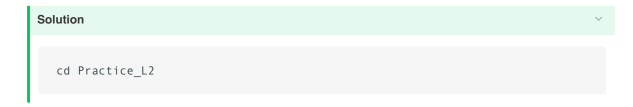
Lesson 2 Practice

The instructions that follow were designed to test the skills you learned in Lesson 2. Thus, the primary focus will be navigating directories and manipulating files.

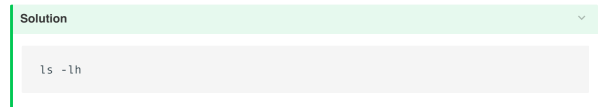
- 1. Let's navigate our files using the command line. Begin in your data directory.
- Copy the Practice_L2 directory (/data/classes/BTEP/B4B_2025/Module_1/ Practice_Sessions/Practice_L2) and all of its contents to your class working directory.

```
cd /data/$USER/Module_1
  cp -R /data/classes/BTEP/B4B_2025/Module_1/Practice_Sessions/Practice_L2 .
```

3. Change directories to Practice_L2/.

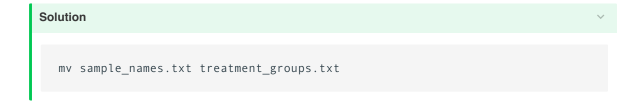


4. List the contents of Practice_L2. If there are files, when were they last modified and what is the file size?



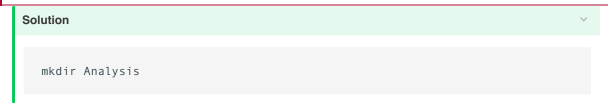
The -1h flag will allow you to obtain a list of directory content in long format, which provides information including the date the file was last modified and the file size.

5. Rename sample_names.txt to treatment_groups.txt.

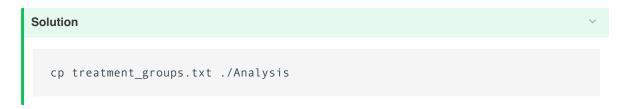


6. Within Practice_L2, create a new directory called Analysis.

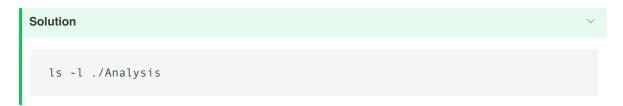
202 Lesson 2 Practice



7. Copy treatment_groups.txt to the newly created Analysis directory.



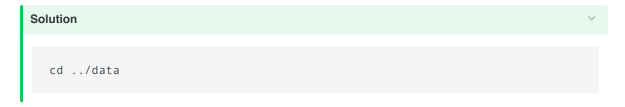
8. List the contents of the Analysis directory.



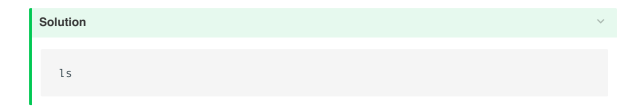
9. Change directories to Analysis. What is the path to Analysis?



10. Change to the data directory within the lesson practice directory (Practice_L2).



11. List the contents of data.



12. View A_R1. fastq.

203 Lesson 2 Practice

```
Solution 

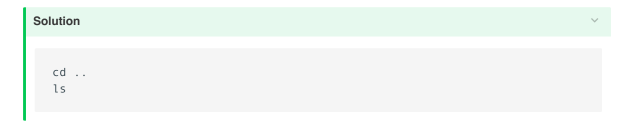
less A_R1.fastq
q
```

- 13. Copy and paste the first line of A_R1.fastq into a new file called Line1.txt using keyboard shortcuts and nano.
- 14. Move Line1.txt to Practice_L2 using a relative file path.

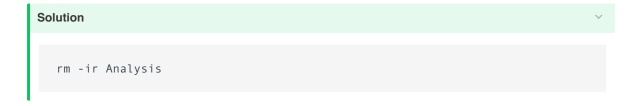
```
Solution v

mv Line1.txt ..
```

15. Change to the Practice_L2 directory and list the content of the directory.



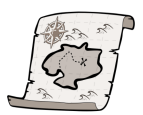
16. Remove the Analysis directory.



Lesson 3 Practice

For today's practice, we are going to embark on a Unix treasure hunt created by the **Sanders** Lab (https://sanderslab.github.io/code/) at the University of California San Francisco. Note: the treasure hunt materials can be obtained directly from the Sanders lab code repository linked above.

UNIX treasure hunt tutorial



This perl script will install a series of directories and clues that teaches basic UNIX command line skills including cd, ls, grep, less, head, tail, and nano. Run the perl script from the command line on a UNIX based machine (e.g. Mac or Linux) using the command: perl treasureHunt_v2.pl. Then use ls to find the first clue. A PDF of command line commands is also available to download.

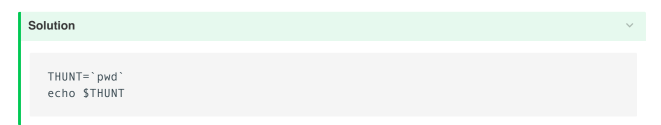
- 🔗 Source
- 🖪 Manual

To begin create a directory called treasure_hunt in your class working directory and run the perl script from /data/classes/BTEP/B4B_2025/Module_1/Practice_Sessions/Practice_L4 in the treasure_hunt directory.

```
mkdir treasure_hunt
cd treasure_hunt
perl /data/classes/BTEP/B4B_2025/Module_1/Practice_Sessions/Practice_L4/treasureHur
ls -l
```

Read the first clue and begin.

Recommendation: Create an environment variable to store the path to the treasure hunt directory to facilitate movement through the directory.

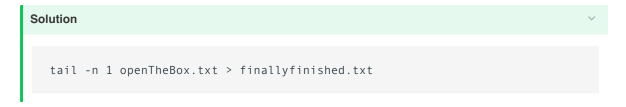


When you have found the treasure, answer or do the following:

1. How many words are in the last line of the file containing the teasure?

```
Solution 
tail -n 1 openTheBox.txt | wc -w
```

2. Save the last line of openTheBox.txt to a new file called finallyfinished.txt without copying and pasting.



3. Now append the first line of openTheBox.txt to the same file that you just saved the last line.



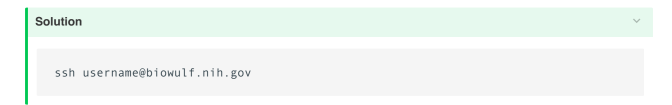
Congratulations! You have found the treasure and have gained some useful unix practice throughout your hunt.

Lesson 4 Practice

The following can be used to practice skills learned in Lesson 4.

Login to Biowulf

If you are already logged in, exit the remote connection and reconnect. Remember, you must be on the NIH network.



Navigate to your working directory.

```
cd /data/$USER/Module_1/testscript
```

Let's run fastqc (https://hpc.nih.gov/apps/fastqc.html), a quality control program, on the files we downloaded from the SRA.

Using sbatch

Start a new script, named fastqc.sh in the same directory in which you downloaded data from Lesson (/data/\$USER/testscript).

The command you will include in the script is as follows:

```
mkdir fastqc
fastqc -o ./fastqc/ -t 4 *.fastq
```

This command will output the fastqc results to a directory named fastqc inside the current working directory. It will also run using four threads and will run on all fastq files in your working directory.

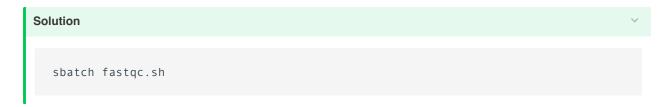
You need edit this script in order to submit as a job on Biowulf. What is missing?

207 Lesson 4 Practice

```
#!/bin/bash
#SBATCH --cpus-per-task=4

module load fastqc
mkdir fastqc
fastqc -o ./fastqc/ -t 4 *.fastq
```

Submit the job.



How can we check on our job? What is the job's status? How much memory is it using?

```
Solution

squeue -u $USER
sjobs -u $USER
```

How can we cancel this job?

```
Solution

scancel job-id

where job-id is the id of the job. Check the output of squeue -u $USER if you are unsure what the job id is.
```

Using sinteractive

What if we want to run this same task interactively? How can we access an interactive session to run the same code above?

```
Solution 

sinteractive --cpus-per-task=4
```

Additional practice materials from hpc.nih.gov

- Submit a job using sbatch. (https://hpc.nih.gov/training/intro_biowulf/hands-on-sbatch.html)
- Submit a multi-threaded job. (https://hpc.nih.gov/training/intro_biowulf/hands-on-multi.html)
- Sumbit a swarm job. (https://hpc.nih.gov/training/intro_biowulf/hands-on-swarm.html)

Lesson 5 Practice

The following was designed to practice skills learned in lesson 5.

Find the data

Here (https://journals.asm.org/doi/full/10.1128/mSystems.00065-20?rfr_dat=cr_pub++0pubmed&url_ver=Z39.88-2003&rfr_id=ori%3Arid%3Acrossref.org) is a paper examining the relationship between the oral microbiome and nasopharyngeal carcinoma. Where can you find the associated data?

Notice that the data was originally submitted to the ENA.

Download the data

Make a directory called Lesson5 practice and change directories.

```
Mkdir Lesson5_practice
cd Lesson5_practice
```

Navigate to the NCBI website (https://www.ncbi.nlm.nih.gov/) and grab the accession information for the associated BioProject.

1. Download the first 10 samples using fastq-dump with parallel.

```
You can download the accession list directly from the SRA Run Selector.

From the command line:

esearch -db sra -query PRJEB37445 | efetch -format runinfo | cut -f 1 -d ',' | sort cat accessions.txt | parallel -j 1 fastq-dump --split-3 {}

Or the same command with prefetch...

cat accessions.txt | parallel -j 1 'prefetch {} | fastq-dump --split-3' {}
```

- 1. Download five samples with the aid of the sra-explorer.
- 2. Navigate to the ENA. How might you go about downloading the data?

Unix Review Practice Session

Author: Stephan Sanders, PhD (UCSF)

In this exercise, we are going to embark on a Unix treasure hunt created by the Sanders Lab (https://sanderslab.github.io/code/) at the University of California San Francisco. Note: the treasure hunt materials can be obtained directly from the Sanders lab code repository linked above.

UNIX treasure hunt tutorial



This perl script will install a series of directories and clues that teaches basic UNIX command line skills including cd, ls, grep, less, head, tail, and nano. Run the perl script from the command line on a UNIX based machine (e.g. Mac or Linux) using the command: perl treasureHunt_v2.pl. Then use ls to find the first clue. A PDF of command line commands is also available to download.

- **Ø** Source
- 🖪 Manual

Note to start at the /data/username folder for this exercise (replace username with the student account ID). To begin create a directory called treasure_hunt in your data directory (ie. /data/username) and change into it. Next, run the treasureHunt_v2.pl perl script in / data/classes/BTEP.

```
mkdir treasure_hunt
cd treasure_hunt
perl /data/classes/BTEP/treasureHunt_v2.pl
ls -l
```

Read the first clue and begin.

Recommendation: Create an environment variable to store the path to the treasure hunt directory to facilitate movement through the directory.



THUNT=`pwd` echo \$THUNT

When you have found the treasure, answer or do the following:

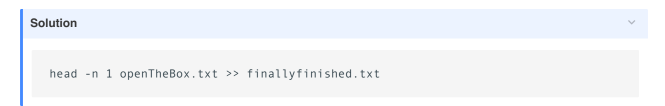
1. How many words are in the last line of the file containing the teasure?



1. Save the last line to a new file called finallyfinished.txt without copying and pasting.



1. Now append the first line to the same file that you just saved the last line.



Congratulations! You have found the treasure and have gained some useful unix practice throughout your hunt.

Lesson 7 Practice

This session will allow users to practice assessing FASTQ quality using data from the HBR-UHR study (see https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/ (https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/)).

First step is to make sure that the participant is signed onto Biowulf. If not then do the following to sign on. Remember to replace user with the participant's own Biowulf sign on ID.



Then change into the data directory.



The practice data is stored in the folder hbr_uhr_b4b in the folder /data/classes/BTEP. Copy it to the data directory.

```
Solution

cp -r /data/classes/BTEP/hbr_uhr_b4b .
```

Change into the hbr_uhr_b4b folder in the participant's data directory.

```
cd hbr_uhr_b4b
```

Request an interactive session with 12 gb of RAM or memory and 10 gb of local temporary storage space.

```
Solution 

sinteractive --mem=12gb --gres=lscratch:10
```

214 Lesson 7 Practice

What are the contents in hbr_uhr_b4b?

```
Solution
   ls -1
   drwxr-x---. 2 wuz8 wuz8 4096 Dec 20 15:41 reads
   drwxr-x---. 2 wuz8 wuz8 4096 Dec 20 14:37 references
There are two folders, 'reads' and 'references'.
   ls -1 reads
The 'reads' folder contains the FASTQ files. There are 12 of these, two for each sample.
  HBR Rep1 R1.fq
  HBR_Rep1_R2.fq
  HBR_Rep2_R1.fq
  HBR_Rep2_R2.fq
  HBR_Rep3_R1.fq
  HBR_Rep3_R2.fq
   {\tt UHR\_Rep1\_R1.fq}
  UHR_Rep1_R2.fq
   UHR_Rep2_R1.fq
  UHR_Rep2_R2.fq
   UHR_Rep3_R1.fq
   UHR_Rep3_R2.fq
   ls references
The references folder contains the chromosome 22 reference (fa or FASTA) and annotation (gtf) files.
  22.fa 22.gtf
There is also a file, 'hbr_uhr_samples.txt' that contains the HBR-UHR sample IDs.
  cat hbr_uhr_samples.txt
  HBR_Rep1
  HBR_Rep2
  HBR_Rep3
   UHR Rep1
   UHR_Rep2
   UHR_Rep3
```

Make a directory called hbr_uhr_b4b_raw_qc inside /data/user/hbr_uhr_b4b.



Load FASTQC.



Run FASTQC for raw FASTQ files and save the results in hbr_uhr_b4b_raw_qc. Stay in /data/user/hbr_uhr_b4b for this.

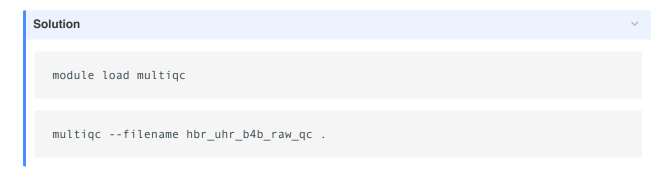
```
fastqc reads/*.fq -o hbr_uhr_b4b_raw_qc
```

Change into hbr_uhr_b4b_raw_qc.

```
Solution 

cd hbr_uhr_b4b_raw_qc
```

Load and run MultiQC to combine all of the FASTQC reports in hbr_uhr_b4b_raw_qc. Name the MultiQC results with prefix hbr_uhr_b4b_raw_qc.



Copy the hbr uhr b4b raw qc.html MultiQC report to local Downloads to view the report.



216 Lesson 7 Practice

Open a new Terminal (Mac) or Command Prompt (Windows). Then change into the local `Downloads` folder.

cd Downloads

scp_user@helix.nih.gov:/data/user/hbr_uhr_b4b/hbr_uhr_b4b_raw_qc/hbr_uhr_b4b_raw_qc

Based on the MultiQC report for the raw FASTQ files, what is the next step.

Solution

Align the data to reference genome since the quality of the data is good and there does not appear to be adapter contamination.

Lesson 8 Practice

Participants will practice trimming using trimmomatic in this help session but with data downloaded from the SRA instead as the HBR-UHR FASTQC report indicated that there are no adapter contamination.

Before getting started, be sure to sign onto Biowulf.



Then change into the participant's data directory.



Create a new folder called practice_trim and change into it.

```
Solution

mkdir practice_trim

cd practice_trim
```

Request an interactive session with 12 gb of RAM (or memory) and 10 gb of local temporary storage.



Use the SRA Tool kit to obtain the first 10,000 sequences for SRR1553606 (https://www.ncbi.nlm.nih.gov/sra/?term=SRR1553606). These are paired end sequences.



```
module load sratoolkit

fastq-dump --split-files -X 10000 SRR1553606

Read 10000 spots for SRR1553606
Written 10000 spots for SRR1553606

Stay in `practice_trim` and list its contents.

ls

SRR1553606_1.fastq SRR1553606_2.fastq
```

QC the two FASTQ files for SRR1553606 (ie. SRR1553606_1.fastq and SRR1553606_2.fastq).



Copy the HTML FASTQC report to local computer <code>Downloads</code> folder to view the results. Is there adapter contamination?



Try trimming away the adapter sequence below from the FASTQ files and rerun FASTQC.

>nextera CTGTCTCTTATACACATCTCCGAGCCCACGAGAC

Hint: copy the above sequence and save it to the file nextera.fa.



Trim away the sequence in nextera.fa using Trimmomatic.

- For quality trimming, use a SLIDINGWINDOW of size 4 bases and average score in the window of 30.
- For adapter trimming, set the adapter strigencies
- seed mismatches: 2
- palindrome clip threshold: 30
- simple clip threshold: 5
- Retain on those trimmed sequences whose length is greater than 50 bases

```
module load trimmomatic

java -jar $TRIMMOJAR PE SRR1553606_1.fastq SRR1553606_2.fastq SRR1553606_trimmed_1.
```

QC the two paired and trimmed FASTQ files.



Copy the two HTML reports for the trimmed FASTQ files to local **Downloads** folder to review the results.

```
Solution

scp user@helix.nih.gov:/data/user/practice_trim/SRR1553606_trimmed_1_fastqc.html .
```

220

scp user@helix.nih.gov:/data/user/practice_trim/SRR1553606_trimmed_2_fastqc.html .

Did trimming remove the adapters and improve the data quality?



Lesson 9 Practice

Since the HBR-UHR data did not need trimming, this practice session will have participants align this data to the chromosome 22 human reference.

Sign onto Biowulf and change into the /data/user/hbr_uhr_b4b folder.



Request an interactive session with 12 gb RAM (or memory) and 10 gb of local temporary storage.

```
Solution

sinteractive --mem=12gb --gres=lscratch:10
```

Create a new folder in /data/user/hbr uhr b4b called hbr uhr hisat2.



Load HISAT2.



Stay in the folder hbr_uhr_b4b for this exercise.

Build HISAT2 indices for the chromosome 22 reference genome. The file for the chromosome 22 genome is located in the folder references and the file name is 22.fa. Give the indices a

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base name (ie. file name without the extension) of 22. List the contents of the references folder to make sure that the build succeeded (ie. the .ht2 files are present).

```
Solution 

hisat2-build references/22.fa references/22
```

After building the indices, align the HBR-UHR FASTQ files to genome. Use the parallel command for this and store the alignment output in the folder hbr uhr hisat2.

```
cat hbr_uhr_samples.txt | parallel "hisat2 -x references/22 -1 reads/{}_R1.fq -2 re
```

Look at the overall alignment rates in the HISAT2 alignment summary files. Why are they low?

```
cat hbr_uhr_hisat2/*summary.txt | grep "overall"

52.75% overall alignment rate
52.51% overall alignment rate
52.68% overall alignment rate
47.91% overall alignment rate
51.26% overall alignment rate
46.94% overall alignment rate
```

"In addition, a spike-in control was used. Specifically we added an aliquot of the ERCC ExFold RNA Spike-In Control Mixes to each sample." -- (Griffith lab RNA Bio, https://rnabio.org/module-01-inputs/0001/05/01/RNAseq_Data/) This dataset has ERCC spike-ins as a quality control measure so not all sequences will map to the human chromosome 22 genome.

Create sorted BAM files and BAM indices for the alignment results.

```
Solution

Create BAM from SAM.

cat hbr_uhr_samples.txt | parallel "samtools sort -o hbr_uhr_hisat2/{}.bam hbr_uhr_
Index BAM.

cat hbr_uhr_samples.txt | parallel "samtools index -b hbr_uhr_hisat2/{}.bam hbr_uhr_
```

Lesson 10 Practice

In this session, participants will practice generating a gene expression matrix for the HBR-UHR data.

Sign onto Biowulf and change into the /data/user/hbr_uhr_b4b folder.



Create a new folder called hbr_uhr_expression.



Request an interactive session with 12 gb of RAM and 10 gb of local temporary storage space.

```
Solution 

sinteractive --mem=12gb --gres=lscratch:10
```

Generate a gene expression table using featureCounts for all of the samples in one go. Change into /data/user/hbr uhr b4b/hbr uhr hisat2 for this.

```
module load subread

featureCounts -p --countReadPairs -a ../references/22.gtf -g gene_name -o ../hbr_ur
```

Change back to the hbr_uhr_expression folder after the gene express matrix has been generated.

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```
Solution 

cd ../hbr_uhr_expression
```

Convert the gene expression matrix hbr_uhr_gene_expression.txt to a CSV file without a header line containing featureCounts information as well as with only the following columns:

- Gene name
- Columns for the expression of each sample

Do this using | (or pipe) to avoid writing intermediate files.

```
sed '1d' hbr_uhr_gene_expression.txt | cut -f1,7-12 | tr '\t' ',' > hbr_uhr_gene_e>

Make sure the result is correct.

column -t -s ',' hbr_uhr_gene_expression.csv | less -S
```

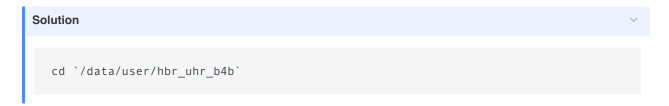
Lesson 11 Practice

In this session, participants will practice making bigWig files from the HBR-UHR alignment results.

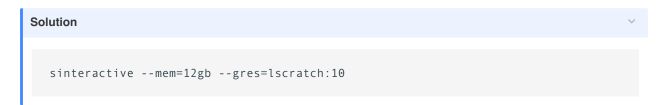
As always, be sure to be signed on to Biowulf for this set of exercises.



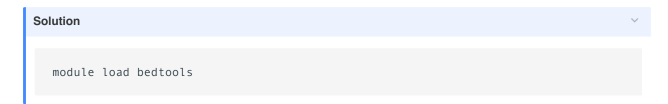
Change into the /data/user/hbr_uhr_b4b folder.



If not done already, request an interactive session with 12gb of RAM and 10gb of local temporary storage space.

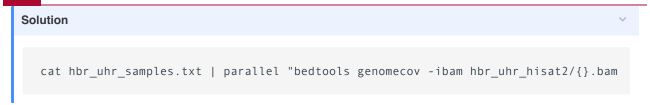


Load bedtools.



Create bedGraph files from the HBR-UHR alignment results. Stay in /data/user/hbr_uhr_b4b and recall that these are RNA sequencing data. Write the bedGraph files to the folder hbr_uhr_hisat2.

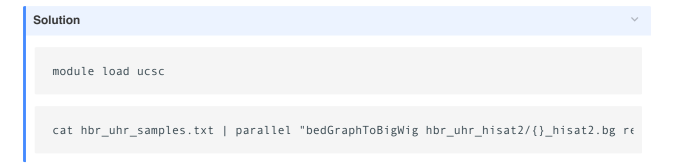
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Next, index the chromosome 22 reference genome stored in the folder references with file name 22.fa. Which module is needed?



Create bigWig files from the HBR-UHR bedGraph files. Which module needs to be loaded to do this? Stay in /data/user/hbr_uhr_b4b for this exercise.



Lesson 12 Practice

Participants can work on these practice questions on their own time as a Biowulf account or local installation of IGV is needed.

Open the bigWig files for HBR_Rep1 (HBR_Rep1_hisat2.bw) and UHR_Rep1 (UHR_Rep1_hisat2.bw). Search of the gene TEF. Does it appear to be more highly expressed in the HBR_Rep1 sample? Use "Human (hg38)" as reference.

Solution

Open the HBR_Rep1_hisat2.bw and UHR_Rep1_hisat2.bw `bigWig` files. Zoom to chromosome 22. Select both `bigWig` tracks, right click and choose "Group Autoscale". Search for TEF. From the IGV view, it does appear that TEF is more highly expressed in HBR_Rep1.

How many transcript isoforms are there for TEF.

Solution

Right click on the Gene track and select expand. There are 5 transcript isoforms associated with TEF, although 3 of these are predicted as the accession starts with "XM" and "XR" (see https://www.ncbi.nlm.nih.gov/books/NBK50679/).

Which of the TEF isoforms is likely to be expressed?

Solution

Load the BAM files for these two samples. It appears that from the splice junctions that NM_003216.4 is expressed. Although in the HBR_Rep1 sample, reads are mapping to the first exon of NM_001145398.2.

Zoom to chromosome 22, position 41,387,655. Notice anything in the HBR_Rep1 sample?

Solution

It looks like there is a heterozygous SNP, where the reference is C but some reads are showing T.

Lesson 13 Practice

In this practice session, participants will filter and perform some quality checks on the HBR-UHR gene expression data.

Before getting started, make sure to be connected to Biowulf and an interactive session with 12 gb of memory and 10 gb of local temporary storage is created.

Change into the /data/user/hbr uhr b4b folder.



Create directory called hbr_uhr_deg to store the differential expression analysis outputs.



What is folder is the HBR-UHR gene expression data table stored and what is the file name?

Solution The gene expression table is stored in the folder `hbr_uhr_expression/`. The name of the gene expression table is `hbr_uhr_gene_expression.csv`. To reference this from `hbr_uhr_b4b` use `hbr_uhr_expression/hbr_uhr_gene_expression.csv`.

Load R.



The scripts are the in the folder b4b_script

Filter low expressing genes out of hbr_uhr_gene_expression.csv. Set the minimum number of samples per group that have greater than 0 expression to be 2. Assign hbr_uhr to the study name and write the output to hbr_uhr_deg.



Rscript b4b_scripts/filter_expression.R hbr_uhr_expression/hbr_uhr_gene_expression.

How many genes are in the filtered expression table?

```
Wc -l hbr_uhr_deg/hbr_uhr_gene_expression_filtered.csv

Since the filter gene expression CSV file has 562 lines then this means 561 genes are left after filtering.
```

Run QC on the filtered expression data. Assign hbr_uhr as the study name and write the output to the folder hbr_uhr_deg.

```
Solution

Rscript b4b_scripts/quality_check.R hbr_uhr_deg/hbr_uhr_gene_expression_filtered.cs
```

After running QC, download the images to the Downloads folder of personal computer to view the results. To do this, open and a new terminal (mac) or command prompt window (Windows 10 or above) and change into the local Downloads folder.

```
cd Downloads
```

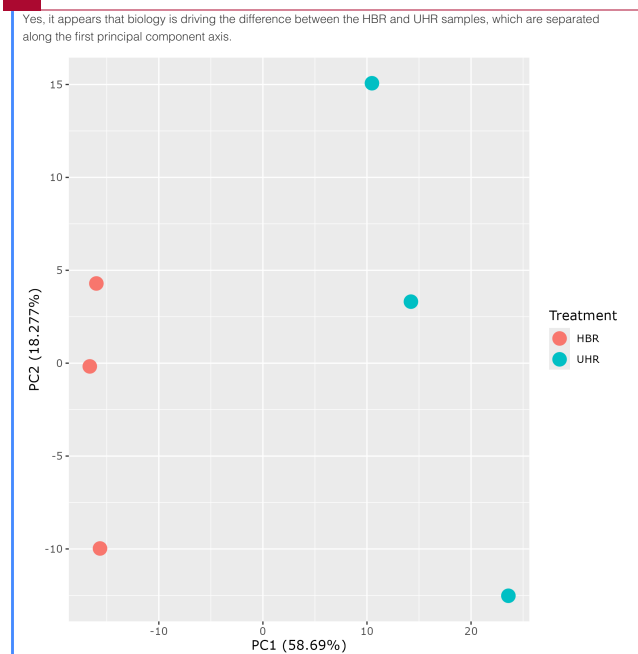
Then use the scp command construct below to download. Remember to replace user with the participants assigned Biowulf student ID.

```
scp -r user@helix.nih.gov:/data/user/hbr_uhr_b4b/hbr_uhr_deg .
```

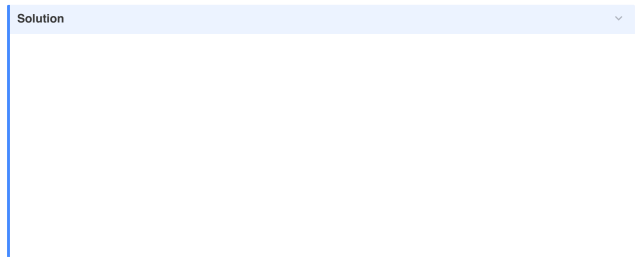
Use the Mac Finder or Windows Explorer to navigate to hbr_uhr_deg in the local Downloads directory to begin exploring the results.

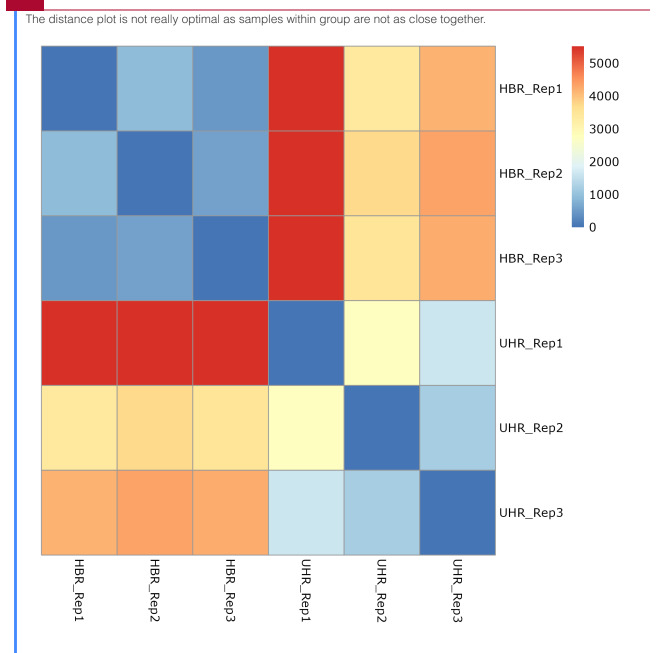
Does it look like the samples are separated by biology?





What is the distance plot informing of?

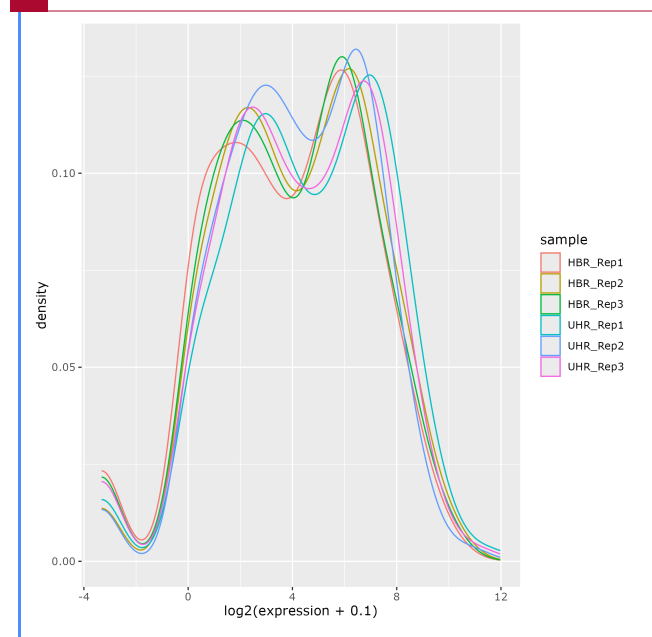




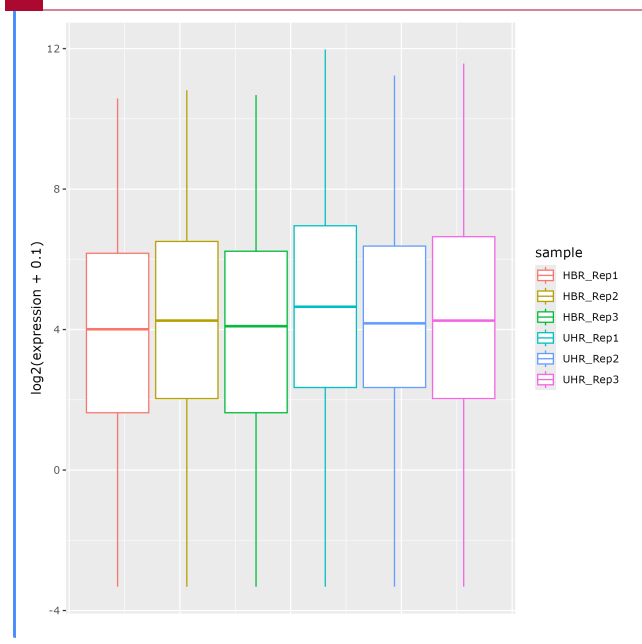
How does the expression distribution among samples look?

Solution

From the density and box plots, the expression distribution are not equal among samples. Hopefully normalization improves this.



Lesson 13 Practice



Lesson 14 Practice

In this practice session, participants will run the actually differential gene expression analysis on the HBR-UHR gene expression data.

Before getting started, be sure to be connected to Biowulf and have an interactive session with 12 gb of memory and 10 gb of local temporary storage space ready. Remember to replace user with the participant's assigned Biowulf student account ID.

Next, change into the /data/user/hbr_uhr_b4b folder.



Load R.

Next, change into the /data/user/hbr_uhr_b4b folder.



Run the deg2.R script in the folder b4b_scripts to generate differential expression analysis results. Use hbr uhr for the study name and write the output to hbr uhr deg.

```
Solution 

Rscript b4b_scripts/deg.R hbr_uhr_deg/hbr_uhr_gene_expression_filtered.csv hbr_uhr_
```

After running the differential expression analysis, copy the hbr_uhr_deg folder to local downloads folder to review the images.

To do this, open and a new terminal (mac) or command prompt window (Windows 10 or above) and change into the local Downloads folder.

cd Downloads

Then use the scp command construct below to download. Remember to replace user with the participants assigned Biowulf student ID.

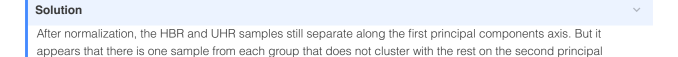
```
scp -r user@helix.nih.gov:/data/user/hbr_uhr_b4b/hbr_uhr_deg .
```

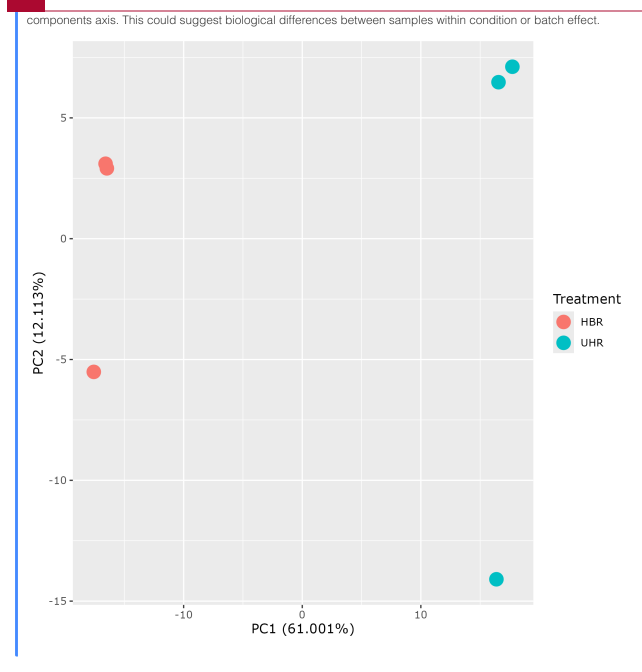
How many genes matched the criteria of log2FoldChange>=4.5 and adjusted p-value <=0.01 or log2FoldChange<=-4.5 and adjusted p-value <=0.01?



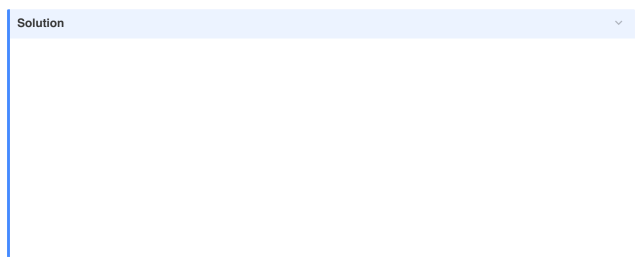
There are 23 (ie. 24 lines in hbr_uhr_filter_deg.csv, so minus 1 line for the column headers to get 23 genes) genes that matched the specified log2FoldChange and adjust p-value threshhold.

What is the PCA obtained from normalized expression table suggesting?

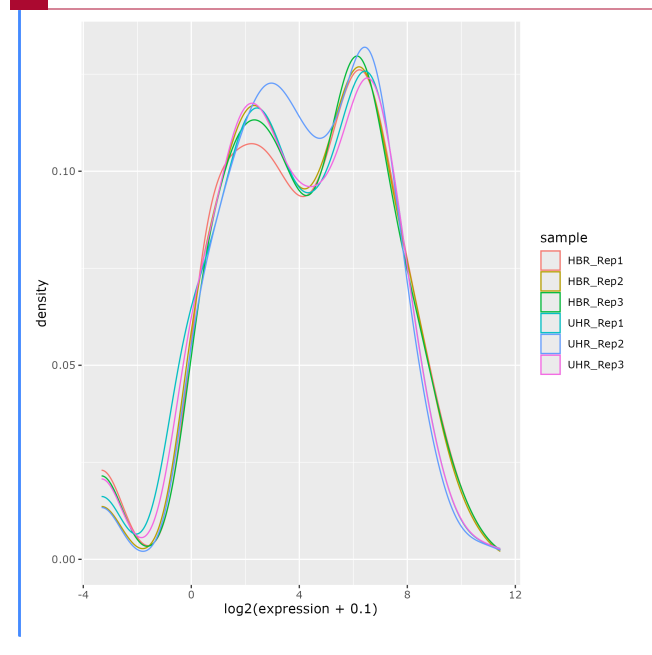




Did normalization help in terms of pushing the gene expression distribution for each sample towards the same.

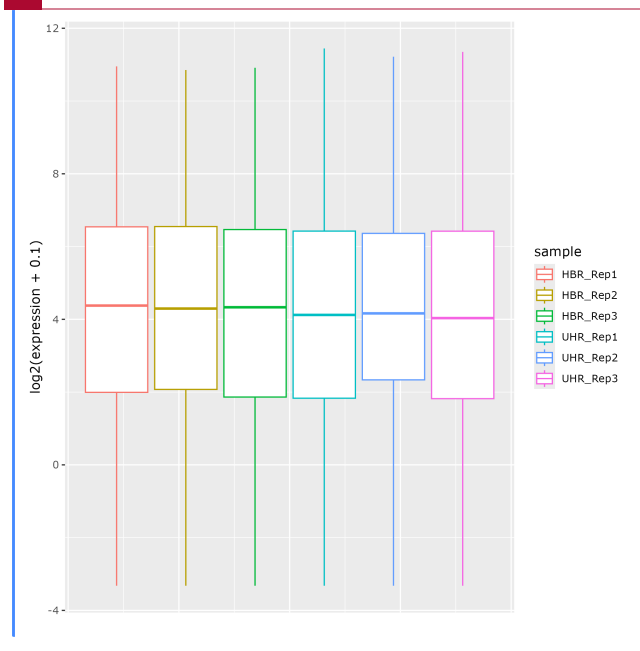


Lesson 14 Practice





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Take a look at the rest of the visualizations (ie. the distance plot, expression heatmap, and volcano plot) to see what can be concluded.