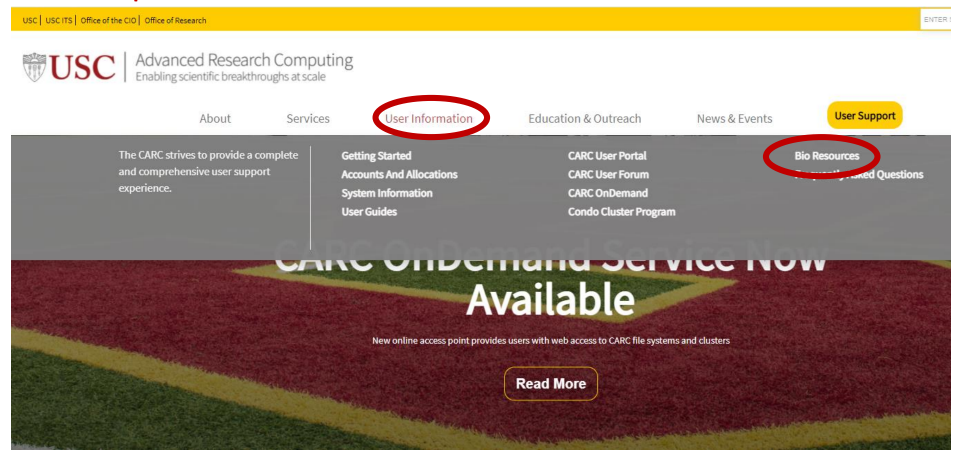
2021 Summer bootcamp Bio Resources at CARC

Center for Advanced Research Computing
USC, June 2021
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Bio Resources (updated quarterly)

https://carc.usc.edu/user-information/bio-resources



Bio Resources (updated quarterly)

https://carc.usc.edu/user-information/bio-resources

- **Genomes** reference sequences and annotations for commonly analyzed organisms
- **Genbank** collection of all public nucleotide sequences and their protein translations
- Genome Taxonomy Database (GTDB) an initiative to establish a standardized microbial taxonomy based on genome phylogeny
- <u>Pfam Database</u> large collection of protein families, each represented by multiple sequence alignments and hidden Markov models (HMMs)
- <u>TIGRFAMs</u> curated multiple sequence alignments, Hidden Markov Models (HMMs) for protein sequence classification
- <u>UniProt</u> UniProtKB (curated protein information), UniRef (closely related sequences),
 UniParc (all protein sequences, consisting only of unique identifiers and sequences)

Bio Resources (upcoming)

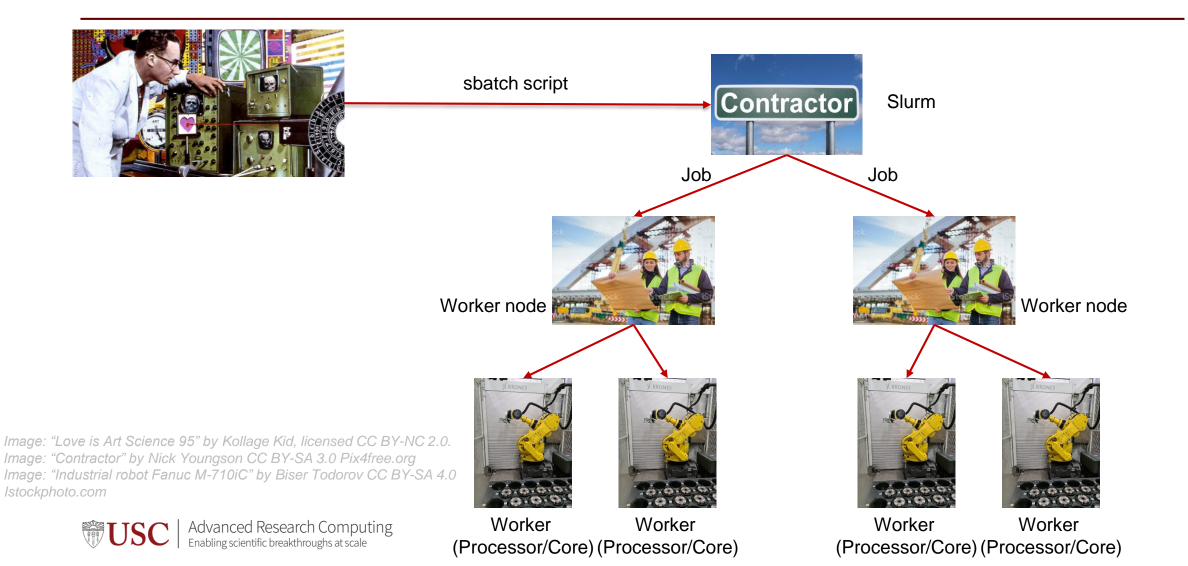
https://carc.usc.edu/user-information/bio-resources

- Biogeotraces set of metagenomes, collected under the auspices of the bioGEOTRACES component of the international GEOTRACES program
- TaraOceans marine microbial metagenomes sampled across space and time
- Variant Effect Predictor cache VEP can use a variety of annotation sources to retrieve the transcript models used to predict consequence types. Cache contains all transcript models, regulatory features and variant data for a species and allows for an offline use of VEP

Some terms

- Head Node The system that controls the cluster
- Worker (Compute) Node Systems that perform the computations in a cluster
- Login Node System that users log into to use a cluster
- Scheduler Software that controls when jobs are run and the node they are run on
- Shell A program that users employ to type commands
- Script A file that contains a series of commands that are executed
- Job A chunk of work that has been submitted to the cluster

How does it work?



What partition should I use?

- debug small, short or test jobs that take less than 30m; short queue
- main (default) most jobs; up to 48h runtime (2 days)
- epyc-64 larger multithreaded jobs; up to 48h (2 days)
- largemem jobs that require huge amount of memory (up to 1TB); up to 168h (7 days)
- oneweek long running jobs; up to 168h (7 days)

Log into CARC

- Open the terminal:
 - Mac: Applications>Utilities>Terminal or open Spotlight and start typing "terminal"
 - Windows: Start menu>cmd (or use PuTTY or Cygwin)
 - Linux: System tools>Terminal or Accessories>Terminal or search for Terminal
- Type ssh <u>ttrojan@discovery1.usc.edu</u>
- Enter your password
- Choose an option in Duo-2FA, and confirm your access
- Answer "No" when asked to save your password
- If successful, your prompt should look something like:
 [ttrojan@discovery1 ~]



Review of Important Commands: squeue

Shows the status of jobs running in the queues

```
[osinski@discovery1 ~]$ squeue | head
          4679566
                                                                  8 d23-[13,15-16], e21-14, e22-[08-09,12], e23-01
                        main discover
                                      sunwool R
                                                     2:19:33
          4680126
                                                                  8 d23-[13-16],e22-[05-06,08-09]
                       main discover
                                      sunwool R
                                                       39:11
          4678655
                       main job.slur liukuang R
                                                    11:09:20
                                                                  1 d14 - 08
                       main 1086-7B
                                                    4:18:00
          4679445
                                        asareh R
                                                                  1 d11-46
          4679444
                       main 1086-7B asareh R
                                                    4:19:31
                                                                  1 d05-40
```

You can also show the status of just your jobs

```
[osinski@discovery1 ~]$ squeue -u ttrojan
            JOBID PARTITION
                                NAME
                                         USER ST
                                                      TIME
                                                           NODES NODELIST (REASON)
          3678639
                  epyc-64
                              test 1 ttrojan PD
                                                      0:00
                                                                4 (Resources)
          3678721 epyc-64
                              test 2 ttrojan PD
                                                      0:00
                                                                4 (Priority)
          3675759
                    epyc-64
                              test 3 ttrojan R 1-01:48:12
                                                                2 b22-[29-30]
```

CD is complete, R is running, PD is waiting to run



Review of Important Commands: sinfo

Show the properties of queues and nodes

```
[osinski@discovery1 ~]$ sinfo
PARTITION AVAIL TIMELIMIT
                           NODES
                                   STATE NODELIST
                                    idle a02-26,e05-[42,76,78,80],e22-13
debug
                     30:00
             up
epyc-64
             up 2-00:00:00
                               32 alloc b22-[01-32]
main*
                               16 alloc$ d11-[02-04],e16-[01-13]
             up 2-00:00:00
             up 2-00:00:00
                               11 maint e16-[14-24]
main*
             up 2-00:00:00
                                4 drain* d17-[03,30],d23-[11-12]
main*
                                6 down* d17-[39-44]
main*
             up 2-00:00:00
main*
             up 2-00:00:00
                                1 drain d06-23
             up 7-00:00:00
                                     mix = 01 - [52, 76], e02 - [70 - 71]
oneweek
                                9 alloc e01-[46,60],e02-[40-46]
oneweek
             up 7-00:00:00
             up 7-00:00:00
                                  idle e01-[48,62,64],e02-[48-69,72-80]
oneweek
largemem
             up 7-00:00:00
                                     mix a16-[02,04]
largemem
             up 7-00:00:00
                                1 alloc a16-03
```

Review of Important Commands: sbatch

- Submit a job to the cluster
- --partition partition you want to summit to
 - Default is "main"
- --nodes Number of nodes
 - Default is 1
- --ntasks Number of CPUs per node
 - Default is 1
- Many more options
 - https://slurm.schedmd.com/sbatch

Review of Important Commands: module

- Loads the necessary environment for a program
- module avail
 - Shows all modules available, or all the software installed
- module load
 - Load the environment for a program
- module list
 - Shows modules loaded
- module unload
 - Removes a loaded module
- module purge
 - Removes all loaded modules

Pay attention to module messages

```
[ttrojan@discovery1 ~]$
[ttrojan@discovery1 ~]$ module purge
[ttrojan@discovery1 ~]$ module load gcc/9.2.0
[ttrojan@discovery1 ~]$ module load bowtie2
[ttrojan@discovery1 ~]$ module load bedtools2
[ttrojan@discovery1 ~]$ module load fastqc
[ttrojan@discovery1 ~]$ module load blast-plus
```

Review: Transferring Files

- SFTP
 - Cyberduck (OSX, Windows) https://cyberduck.io
 - WinSCP (Windows) https://winscp.net/eng/index.php
 - FileZilla (OSX, Windows, Linux) https://filezilla-project.org
- Globus Online (Best way to get data from CARC)
 - go to https://www.globus.org in your browser and click Log In
 - Search for University of Southern California in the box that says "Use your existing organizational login"
- Command line tools rsync or scp

More info:

https://carc.usc.edu/user-information/user-guides/data-management/transfer-overview

Resources

- CARC home page
 - https://carc.usc.edu
- Bio Resources at CARC
 - https://carc.usc.edu/user-information/bio-resources
- CARC User Forum
 - https://hpc-discourse.usc.edu/categories
- SLURM tutorials
 - https://slurm.schedmd.com/tutorials.html
- SLURM quick reference
 - https://slurm.schedmd.com/pdfs/summary.pdf



Resources

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 - https://carc.usc.edu/user-information/bio-resources
- CARC User Forum ← the most value for the community!
 - https://hpc-discourse.usc.edu/categories
- SLURM tutorials
 - https://slurm.schedmd.com/tutorials.html
- SLURM quick reference
 - https://slurm.schedmd.com/pdfs/summary.pdf



Review: Interactive Jobs

When you need to provide unpredictable input

```
[ttrojan@discovery1 ~]$ hostname
discovery1.usc.edu
[ttrojan@discovery1 ~]$ srun -p debug --pty bash
[ttrojan@a02-26 ~]$ hostname
a02-26.hpc.usc.edu
[ttrojan@a02-26 ~]$ exit
exit
[ttrojan@discovery1 ~]$ hostname
discovery1.usc.edu
[ttrojan@discovery1 ~]$
```

Review: Bash Scripts

- Bash scripts are a series of commands that can be grouped together within files to accomplish a series of tasks
- This allows you to run one command instead of several successive commands

Exercise:

- Start an interactive job to the debug queue
- This program sleeps for 10 seconds and then prints out "Hello World"
- Make this file, give it execute permissions, and run

```
#!/bin/bash
# This program: sleeps for 10 seconds, then prints "Hello World"
sleep 10
echo "Hello World"
```



Bash Variables

```
cd raw-seq
i=1
ls -l yeast_${i}_50K.fastq
i=2
ls -l yeast_${i}_50K.fastq
```

Lets get going

- Detailed policies and directions
 - https://carc.usc.edu/user-information/getting-started
- Do not install software yourself, contact us
 - https://carc.usc.edu/education-and-outreach/office-hours (Tue, 2:30-5:00)
 - Submit a ticket! (<u>https://carc.usc.edu/user-support/</u>)
 - When we install software, it is available to everyone
- Program running slow? Submit a ticket!
- Don't know what resources to use? Submit a ticket!
- Any other questions? Submit a ticket or visit our forum

Prepare to Run Jobs

Copy example data to your home directory

```
[ttrojan@discovery1 ~]$
[ttrojan@discovery1 ~]$ git clone https://github.com/uschpc/workshop-bioresources-bootcamp-2021.git
[ttrojan@discovery1 ~]$ cd workshop-bioresources-bootcamp-2021
[ttrojan@discovery1 ~]$ ls
```

Create the FastQC Job Script

 Use a text editor to create a file name samplefastqc.sh that contains what follows:

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH --partition debug
#SBATCH --chdir /home1/ttrojan/bio-bootcamp
#SBATCH --account=renney_710
module purge
module load gcc/9.2.0
module load fastqc
echo "Example FastQC start"
sleep 20
fastqc -o results/fastqc-rawseq raw-seq/yeast_1_50K.fastq
echo "Example FastQC end"
```

Run the FastQC Job Script

Submit the job

```
[ttrojan@discovery1 ~]$ sbatch fastqc1.job
Submitted batch job 33723
```

Check the status of the job

```
[osinski@discovery1 ~]$ squeue -u osinski

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

33723 dbeug fastqc.s osinski R 0:02 1 a02-26
```



Check Output File for Errors

Check Output File for Errors

```
[ttrojan@discovery1 ~]$ cat slurm-33723.out
Started analysis of yeast 1 50K.fastq
Approx 5% complete for yeast 1 50K.fastq
Approx 10% complete for yeast 1 50K.fastq
Approx 15% complete for yeast 1 50K.fastq
Approx 20% complete for yeast 1 50K.fastq
Approx 25% complete for yeast 1 50K.fastq
Approx 30% complete for yeast 1 50K.fastq
Approx 35% complete for yeast 1 50K.fastq
Approx 40% complete for yeast 1 50K.fastq
Approx 45% complete for yeast 1 50K.fastq
Approx 50% complete for yeast 1 50K.fastq
Approx 55% complete for yeast 1 50K.fastq
Approx 60% complete for yeast 1 50K.fastq
Approx 65% complete for yeast 1 50K.fastq
Approx 70% complete for yeast 1 50K.fastq
Approx 75% complete for yeast 1 50K.fastq
Approx 80% complete for yeast 1 50K.fastq
Approx 85% complete for yeast 1 50K.fastq
Approx 90% complete for yeast 1 50K.fastq
Approx 95% complete for yeast 1 50K.fastq
Approx 100% complete for yeast 1 50K.fastq
Analysis complete for yeast 1 50K.fastq
```

Important Things to Note

- Job length
 - If over 24 hours, can this be split up, can threads be increased?
- Many small files
 - To be avoided!
 - Group into larger files
- Data
 - Save space by removing temp files
 - Archive data as soon as reasonable
 - Let us know if you are adding several TB of data
 - Use /scratch or /scratch2 whenever possible for temporary files

Important Things to Note

- Make sure you are not on the login node when you launch an application
 - You can check the system you are on by typing hostname
- Make sure you reserve as many processors as you need
 - A mismatch here can increase your runtime or wait time
- Make sure you reserve as much RAM as needed
 - Overestimating increases wait time, underestimating crashes
- Know which resources work the best
 - Sometimes using a debug or epyc-64 is better

Important Things to Note

No HIPAA Data is allowed on the cluster!

(we are working on that part)

SLURM Environment Variables

- \$SLURM JOB ID The job number
 - Assigned automatically by SLURM
- \$SLURM_JOB_NAME The name of the job
 - Similar to the script name
 - Or you can specify one with -J
- \$SLURM_NTASK Number of reserved Processors
 - Assigned automatically by SLURM
 - Value is the multiple of the --ntasks and --nodes values

Multi-Processor Jobs

- The program must support it!
- Our default nodes have mostly 20 cores. Some programs loose efficiency after 8 or 16 processors.
- Wait time and run time adds up if not properly submitted
- Try "program --help" or "man program"
- Use \$SLURM NTASKS

Create the BLAST Job Script

Replace swissprot with the path to the v5 of swissprot db obtained from

https://carc.usc.edu/user-information/bio-resources/genbank

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 10
#SBATCH --partition debug
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
#SBATCH -time 00:05:00
#SBATCH --account=reneny_710
module purge
module load gcc/9.2.0
module load blast-plus
echo "Start BLAST Job"
blastp -db swissprot -query blast/query.txt -out results/blast/results.txt -num_threads
$SLURM_NTASKS
echo "Finish BLAST Job"
```

Run the BLAST Job Script

Submit the job

```
[ttrojan@discovery1 ~]$ sbatch blast1.job
Submitted batch job 4773117
```

Check the status of the job

```
[ttrojan@discovery1 ~]$ squeue -u ttrojan

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

4773117 Main blast1.j ttrojan R 0:02 1 a02-d11
```



Check BLAST Job Stats with sacct

sacct can get stats for a job after its completed

https://slurm.schedmd.com/sacct.html

[ttrojan@discovery1 ~] \$ sacct -j 4773117 --format=JobID, State, Elapsed, NCPUS, MaxRSS

[ttrojan@disco	overy1 ~]\$	sacct -j 4773	117forma	at=JobID,Stat	e,Elapsed,NCPUS,MaxRS
JobID	State	Elapsed	NCPUS	MaxRSS	
1773117	COMPLETED	00:00:09	10		
1773117.bat+	COMPLETED	00:00:09	10	1228K	
4773117.ext+	COMPLETED	00:00:09	10	832K	

GPU Jobs – Example

Use gpu partition

Reserve gpus with --gres parameter

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH --gres=gpu:p100:1
#SBATCH --mail-user=ttrojan@usc.edu
#SBATCH --mail-type=ALL
#SBATCH --chdir /home1/ttrojan
#SBATCH -accountrenney_710
module load gcc/8.3.0
module load cuda/10.0.130
module load motioncor2
```

Job Arrays

- A way to run the same commands on many (hundreds, thousands) of datasets/samples.
- A variable called \$SLURM_ARRAY_TASK_ID is used to determine the element of the array being run.
- #SBATCH --array=1-1000
- \$SLURM ARRAY TASK ID becomes 1 in first job, 2 in second job, etc...



Without Job Arrays – Numbered Files

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH --partition main
#SBATCH --time 00:05:00
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
#SBATCH --account=renney 710
module purge
module load gcc/9.2.0
module load fastqc
echo "Starting FastQC job"
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 1 50K.fastq
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 2 50K.fastq
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 3 50K.fastq
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 4 50K.fastq
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 5 50K.fastq
fastqc -o results/fastqc-rawseq-ordered raw-seq-ordered/yeast 6 50K.fastq
echo "Finish FastQC job"
```

Job Arrays – Numbered Files

 Here is an example SLURM script for a job array. Save as fastqc_numbered_array.job

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH --partition main
#SBATCH --time 00:05:00
#SBATCH --array=1-6
#SBATCH --array=1-6
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
#SBATCH --account=renney_710
module purge
module load gcc/9.2.0
module load fastqc
echo "Starting FastQC job"
sleep 20
fastqc -o results/fastqc-rawseq-ordered-arr raw-seq-rdered/yeast_${SLURM_ARRAY_TASK_ID}_50K.fastq
echo "Finish FastQC job"
```

View Job Array

squeue -u uscnetid

```
[ttrojan@disocvery1 bio-bootcamp]$ squeue -u ttrojan
                                               NODES NODELIST (REASON)
JOBID PARTITION
                   NAME
                            USER ST
                                          TIME
1152 main
          bash ttrojan R
                                2:17:32
                                             1 d05-40
1153 main
             bash ttrojan R
                                2:17:12
                                             1 d05-40
1207 1 main numbered ttrojan R
                                               1 d05-41
                                     0:02
1207 2 main numbered ttrojan R
                                     0:02
                                               1 d05-40
1207 3 main numbered ttrojan R
                                               1 d05-42
                                     0:02
1207 4 main numbered ttrojan R
                                     0:02
                                               1 d05-45
1207 5 main numbered ttrojan R
                                     0:02
                                               1 d05-44
1207 6 main numbered ttrojan R
                                               1 d05-44
                                     0:02
```

Job Arrays – Unnumbered Files

- Start by creating a list of all of the unnumbered filenames
- Then create slurm array script for fastqc jobs that have unnumbered filenames

```
#!/bin/bash
#SBATCH --nodes 1
#SBATCH --ntasks 1
#SBATCH --partition main
#SBATCH --time 00:05:00
#SBATCH --array=1-6
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
#SBATCH --account=renney 710
module purge
module load gcc/9.2.0
module load fastgc echo "Starting FastQC job"
sleep 20
ls raw-seq/ > unnumbered-filenames.txt
line=$(sed -n -e "$SLURM ARRAY TASK ID p" unnumbered-filenames.txt)
fastqc -o results/fastqc-rawseq-unordered raw-seq/${line}
echo "Finish FastQC job"
```

Job Dependencies

- Instructions on how jobs relate to other jobs
- Useful for if you want to run a series of jobs that depend on the output from other jobs
- Examples:

```
-d depend=afterok:jobid

Starts after jobid has finished without errors.
```

-d depend=afterok:jobid, before:jobid2

Starts after jobid is finished, but not until jobid2 has started.

```
-d depend:afterok:jobid
-d depend:afterok:jobid2
```

Starts after both jobid and jobid2 have finished.

-d depend=afterokarray:jobid

Starts after the job array jobid has finished without errors.

Job Dependencies

- Why would you do this:
 - Mostly for job pipelines, a series of programs that depend on each other's output that are all submitted at once.

Example:

Step 1:

```
[ttrojan@discovery1 ~]$ sbatch preprocessing-step.sh Submitted batch job 18866
```

Step 2:

```
[ttrojan@discovery1 ~]$ sbatch -d after:18866 job-array-step.sh Submitted batch job 18870
```

Step 3:

```
[ttrojan@discovery1 ~]$ sbatch -d afterok:18870 postprocessing-step.sh Submitted batch job 18867
```

What is Wrong

What is Wrong

The module is not loaded

What is Wrong II

```
#!/bin/bash
# ------SLURM Parameters-----
#SBATCH --partition main
#SBATCH --ntasks 20
#SBATCH --mem=10g
#SBATCH --nodes 1
# ------Load Modules------
module purge
module load gcc/9.2.0
module load blast-plus
# ------Commands------
blastn -query fasta.file -db database_name -outfmt 6 \
-num_alignments 1 -num_descriptions 1 -out output_file
```

What is Wrong II

Number of processors and no working directory

```
#!/bin/bash
# -------SLURM Parameters------
#SBATCH --partition main
#SBATCH --ntasks 20
#SBATCH --mem=10g
#SBATCH --nodes 1
#SBATCH --chdir /homel/ttrojan/workshop-bioresources-bootcamp-2021
# --------Load Modules-------
module purge
module load gcc/9.2.0
module load blast-plus
# -------Commands-------
blastn -query fasta.file -db database_name -outfmt 6 num_alignments 1 \ -num_descriptions 1 -out output_file _-num_threads 20
```

What is Wrong II

- Number of processors and no working directory
- Better to use \$SLURM_NTASKS

```
#!/bin/bash
# -------SLURM Parameters-----
#SBATCH --partition main
#SBATCH --ntasks 20
#SBATCH --mem=10g
#SBATCH --nodes 1
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
# ------Load Modules-------
module purge
module load gcc/9.2.0
module load blast-plus
# -------Commands------
blastn -query fasta.file -db database_name -outfmt 6 num_alignments 1 \ -num_descriptions 1 -out
output_file _-num_threads $SLURM_NTASKS
```

What is Wrong III

```
#!/bin/bash
# ------SLURM Parameters------
#SBATCH --partition main
#SBATCH --ntasks 1
#SBATCH --mem=200g
#SBATCH --nodes 1
#SBATCH --chdir /home1/ttrojan/workshop-bioresources-bootcamp-2021
# ------Load Modules-------
module purge
module load gcc/8.3.0
module load R
# -------Commands-------
Rscript /home1/ttrojan/R_example.R
```

What is Wrong III

Wrong partition/mem requirements too high

What is Wrong IV

What is Wrong IV

GPU resources not specified

What is Wrong V

What is Wrong V

- No bash shebang line, #!/bin/bash
- Can use long names for SBATCH parameters

Genome mapping and tools: Read mapping

- Aim: to find coordinates of reads in the reference genome.
- Challenges:
 - Millions of short sequences
 - Sequences are often paired
 - Errors are not randomly distributed
- Most popular programs are <u>bow</u>tie and <u>bwa</u> (both use <u>Burrows-Wheeler</u> Transform algorithm). Two-step approach:
 - Create an index for the reference genome (one time for one genome).
 - Map reads to the reference genome using this index

Genome mapping and tools – overview I

- FastQC
 - FastQC is a quality control application for high throughput sequence data
 - Checks the quality of their sequence data
 - Generates an HTML report

http://www.bioinformatics.babraham.ac.uk/projects/fastqc/

Genome mapping and tools – overview II

- bowtie
 - The first version of bowtie [Langmead et al. 2009] is optimal for:
 - short reads (under 50 bp)
 - reads without indels (insertions/deletions)
- bowtie2
 - The second version of bowtie2 [Langmead & Salzberg 2012] is optimal for:
 - long reads (more than 50 bp)
 - reads with indels
 - various alignment options
- Each version has its own index file format (bowtie-build / bowtie2-build tools).
- A popular RNA-seq analysis toolset (tophat, cufflinks) is based on bowtie / bowtie2

http://bowtie-bio.sourceforge.net

Genome mapping and tools – overview III

- bwa
 - bwa backtrack [Li, Durbin 2009]:
 - for short reads (< 100bp)
 - bwa bwasw [Li, Durbin 2010]:
 - for long reads (70bp 1Mbp)
 - short indels
 - bwa mem [Li 2013]:
 - for long reads (70bp 1Mbp)
 - faster and more efficient

Genome mapping and tools – overview IV

- samtools package A set of utilities for processing SAM/BAM files
- samtools view
 - convert a bam file into a sam file samtools view sample.bam > sample.sam
 - Convert a sam file into a bam file samtools view -bS sample.sam > sample.bam
 - Extract all the reads aligned to the range specified. An index of the input file is required

```
samtools view -h -b sample_sorted.bam "chr1:10-13" > tiny_sorted.bam
```

samtools sort

samtools sort unsorted_in.bam sorted_out

samtools index

samtools index sorted.bam (creates an index file, sorted.bam.bai)

http://samtools.sourceforge.net



Genome mapping and tools – overview IV

- samtools package A set of utilities for processing SAM/BAM files
- samtools view

- -b: output BAM -S: read SAM
- convert a bam file into a sam file samtools view sample.bam > sample.sam
- Convert a sam file into a bam file samtools view -bS sample.sam > sample.bam
- Extract all the reads aligned to the range specified. An index of the input file is required •

```
samtools view -h -b sample_sorted.bam "chr1:10-13" > tiny_sorted.bam
```

samtools sort

```
samtools sort unsorted in.bam sorted out
```

samtools index

```
samtools index sorted.bam (creates an index file, sorted.bam.bai)

<a href="http://samtools.sourceforge.net">http://samtools.sourceforge.net</a>
```



Genome mapping and tools – overview IV

samtools flagstat – report basic statistics

```
samtools flagstat sample.bam
```

An example of output:

```
4198456 + 0 in total (QC-passed reads + QC-failed reads)
0 + 0 duplicates
4022089 + 0 mapped (95.80%:-nan%)
4198456 + 0 paired in sequencing
2099228 + 0 read1
2099228 + 0 read2
3796446 + 0 properly paired (90.42%:-nan%)
4013692 + 0 with itself and mate mapped
8397 + 0 singletons (0.20%:-nan%)
167574 + 0 with mate mapped to a different chr
72008 + 0 with mate mapped to a different chr (mapQ>=5)
```

samtools faidx – index a FASTA file

samtools faidx ref.fasta (creates an index file ref.fasta.fai)

samtools merge – merge several BAM files into one

samtools merge out.bam in1.bam in2.bam



Genome mapping and tools – overview V

BedTools package

- bamtobed Convert BAM alignments to BED (& other) formats
- bamtofastq Convert BAM records to FASTQ records
- bedtobam Convert intervals to BAM records
- closest Find the closest, potentially non-overlapping interval
- complement Extract intervals _not_ represented by an interval file
- coverage Compute the coverage over defined intervals
- genomecov Compute the coverage over an entire genome
- getfasta Use intervals to extract sequences from a FASTA file
- intersect Find overlapping intervals in various ways
- shuffle Randomly redistribute intervals in a genome
- sort Order the intervals in a file

Genome mapping and tools – overview V

bedtools intersect

• Report the intervals that represent overlaps between your two files:

bedtools intersect -a cpg.bed -b exons.bed

Report the original feature in each file:

bedtools intersect -a cpg.bed -b exons.bed -wa -wb

How many base pairs of overlap were there?

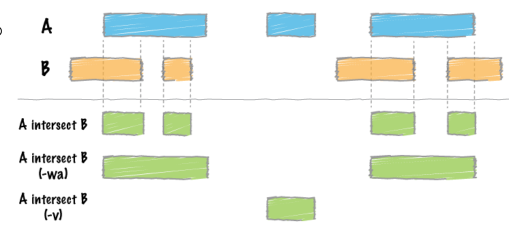
bedtools intersect -a cpg.bed -b exons.bed -wo

Counting the number of overlapping features:

bedtools intersect -a cpg.bed -b exons.bed -c

Find features that DO NOT overlap:

bedtools intersect -a cpg.bed -b exons.bed -v



Exercise

 There are paired reads of some DNA sequencing experiment of the human sample:

```
bio-bootcamp/R1.fastq.gz
bio-bootcamp/R2.fastq.gz
```

- You will study some particular region of the human genome
- Map reads to the human reference genome (version hg19 find path on our Bio Resources)
- Extract reads that map to your region only
- Upload the reads to UCSC genome browser as a custom track
- count the number of insertions and deletions in SAM file

How To: Mapping

load bowtie2 program:

```
module purge
module load gcc/9.2.0
module load bowtie2
```

- Copy sequence of a chromosome your region is located at as a FASTA file
 - find the path on our website in Homo sapiens > UCSC > hg19 > Chromosomes (7th column)
 - https://carc.usc.edu/user-information/bio-resources/reference-genomes
 - Add chr*.fa at the end of the path
 - cp path above /home1/ttrojan/bio-bootcamp/results/read-mapping
- Map reads to this chromosome using bowtie2 with the standard parameters

Don't forget to make an index (bowtie-build2) of the chromosome before mapping!

- You will get a SAM file as an output, convert to BAM (samtools view)
- Count the number of insertions and deletions in SAM file (use cut for field 6, and grep)

How To: Extracting reads

load BedTools:

```
module purge
module load gcc/9.2.0
module load bedtools2
```

• Create a tab-delimited BED file with the coordinates of your region:

```
chr21 1000000 2000000
```

- Convert SAM file with mapped reads to BAM file using samtools view
- Use bedtools intersect to extract the reads from the BAM file

You'll need a BED file to upload the result to UCSC genome browser, so figure out how to make bedtools intersect to produce an output in BED format.

How To: UCSC custom track

Upload the BED file to UCSC genome browser

'Add custom track' button \rightarrow Choose file \rightarrow Submit

Thanks to the whole CARC team:

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Ryan Sim

Carolina Jin

Reference material:

- HPCBio (Holmes J., Clark L., Drnevicch J., Valizadegan N.)
- CNRG (Davidson D., Leigh J.)
- Skoltech (Khrameeva E.)

Mapping exercise: Answers

```
#!/bin/bash
#SBATCH --partition main
#SBATCH --nodes 1
#SBATCH --ntasks 20
#SBATCH --time 01:00:00
#SBATCH --chdir /homel/osinski/bio-bootcamp
#SBATCH --account renney 710
#SBATCH --mem 4g
module purge
module load gcc/9.2.0
module load bowtie2
module load samtools
module load bedtools2
mkdir results/read-mapping
cp data/R*.gz results/read-mapping
gunzip results/read-mapping/R1.fastq.gz
gunzip results/read-mapping/R2.fastg.gz
cp -v /project/biodb/genomes/Homo sapiens/UCSC/hg19/Sequence/Chromosomes/chr21.fa results/read-mapping/
bowtie2-build --threads $SLURM NTASKS results/read-mapping/chr21.fa results/read-mapping/chr21index
bowtie2 --threads $SLURM NTASKS -x results/read-mapping/chr21index -q results/read-mapping/R1.fastq > results/read-mapping/R1.sam
bowtie2 --threads $SLURM NTASKS -x results/read-mapping/chr21index -q results/read-mapping/R2.fastq > results/read-mapping/R2.sam
samtools view results/read-mapping/R1.bam | cut -f 6 | grep -c 'D' > results/read-mapping/R1.no of deletions.txt
samtools view results/read-mapping/R1.bam | cut -f 6 | grep -c 'I' > results/read-mapping/R1.no of insertions.txt
samtools view results/read-mapping/R2.bam | cut -f 6 | grep -c 'D' > results/read-mapping/R2.no of deletions.txt
samtools view results/read-mapping/R2.bam | cut -f 6 | grep -c 'I' > results/read-mapping/R2.no of insertions.txt
samtools view -bS results/read-mapping/R1.sam > results/read-mapping/R1.bam
samtools view -bS results/read-mapping/R2.sam > results/read-mapping/R2.bam
samtools sort results/read-mapping/R1.bam > results/read-mapping/R1 sorted.bam
samtools sort results/read-mapping/R2.bam > results/read-mapping/R2 sorted.bam
samtools view -h -b results/read-mapping/R1 sorted.bam "chr21:10000000-20000000" > results/read-mapping/R1 sorted region.bam
samtools view -h -b results/read-mapping/R2 sorted.bam "chr21:10000000-20000000" > results/read-mapping/R2 sorted region.bam
bamToBed -i results/read-mapping/R1 sorted region.bam > results/read-mapping/R1 sorted region.bed
bamToBed -i results/read-mapping/R2 sorted region.bam > results/read-mapping/R2 sorted region.bed
bedtools intersect -a results/read-mapping/R1 sorted region.bed -b results/read-mapping/R2 sorted region.bed > results/read-mapping/reads.bed
```