# Analyzing RADSeq Data

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FISH 546: Bioinformatics

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 Can I perform those steps following along from the methods section of a paper?

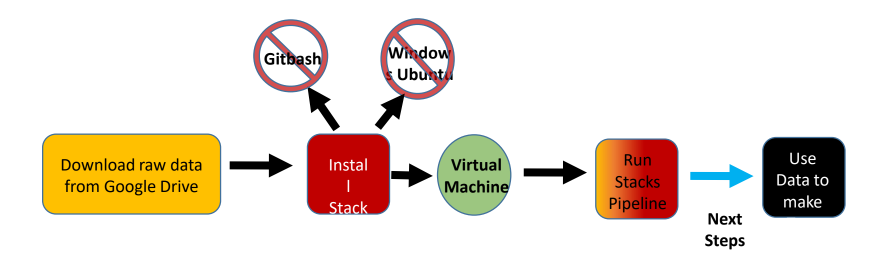
# Question:

What are the steps to go from raw RADSeq data to data that could be input into a tree?

 Can I perform those steps following along from the methods section of a paper?

Spoiler alert: NO

# RADSeq Workflow using Stacks v 2.2





Raw data (37 gb fq.gz)
Barcode file (.txt)

Demultiplex samples with 'process\_radtags'

Directory with 48 fq.gz files, 1 per specimen

Filter and cap samples by read number with my script

Directory with 44 fq.gz files. 1e5<n-reads<2e6

Run `ustacks` on each sample to find loci

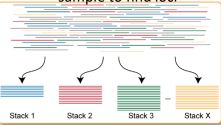


Figure http://catchenlab.life.illinois.edu/stacks/param\_tut.php

Directory with ustacks output (Name.tags.tsv.gz, Name.snps.tsv.gz, Name.alleles.tsv.gz

Run `cstacks` on all samples to build catalog of loci (de novo

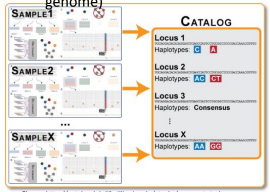


Figure http://catchenlab.life.illinois.edu/stacks/param\_tut.php

Directory with ustacks and cstacks output data (ustack files and catalog files)

Run 'sstacks' on all samples to align against the catalog

Directory with sstacks, ustacks and ccstacks data (name.matches.tsv.gz)

Directory with ustack, cstacks, and sstacks outputs

Run `tsv2bam` to transpose data to a bam alignment by loci instead of by specimen

Directory with tsv2bam output and ustacks, cstacks, and cstacks outputs (name.matches.bam)

Run `gstacks` to genotype

Directory with gstacks outputs?

Run `populations` to get population summary statistics such as F<sub>ST</sub> and output data into a fasta or phylip that can be used for analysis

Fasta or phylip files that can now be used to make a tree!!

Raw data (37 gb fq.gz) Barcode file (.txt)

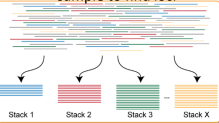
**Demultiplex samples** with 'process radtags'

Directory with 48 fq.gz files, 1 per specimen

Filter and cap samples by read number with my script

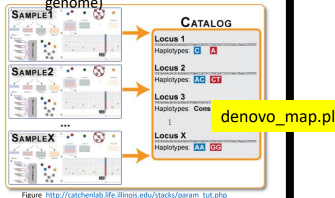
Directory with 44 fq.gz files. 1e5<n-reads<2e6

> Run 'ustacks' on each sample to find loci



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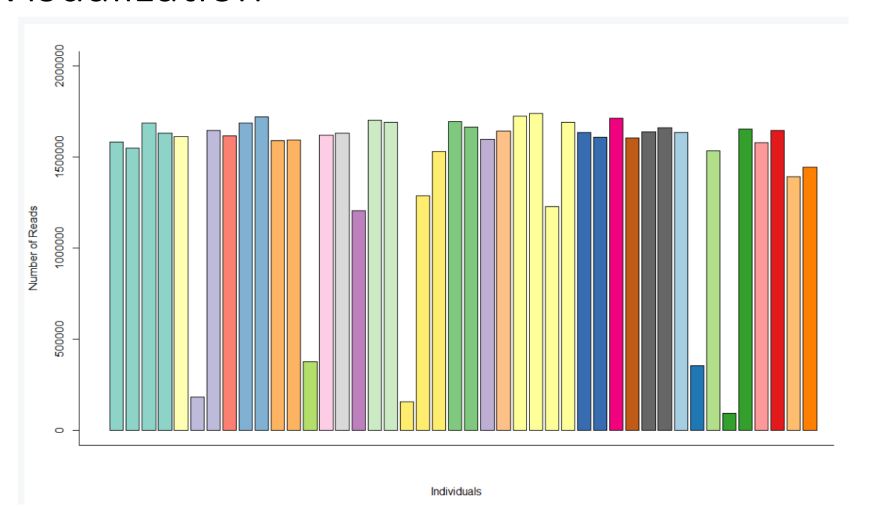
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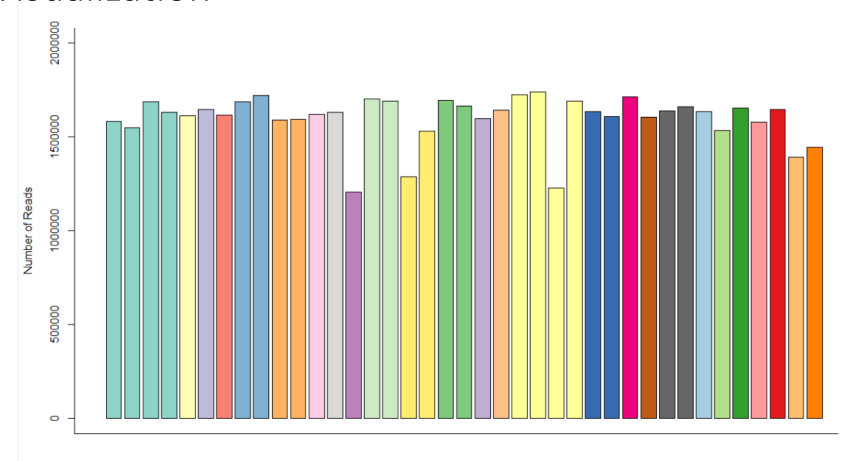
Run 'gstacks' to genotype

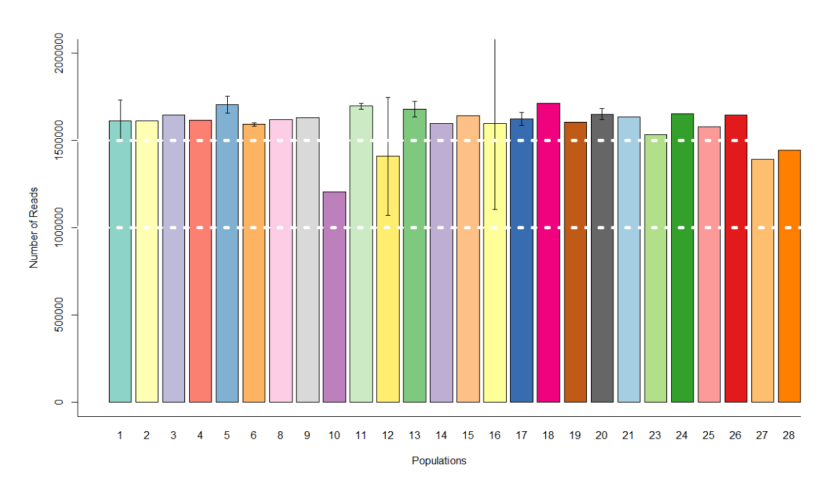
Directory with gstacks outputs?

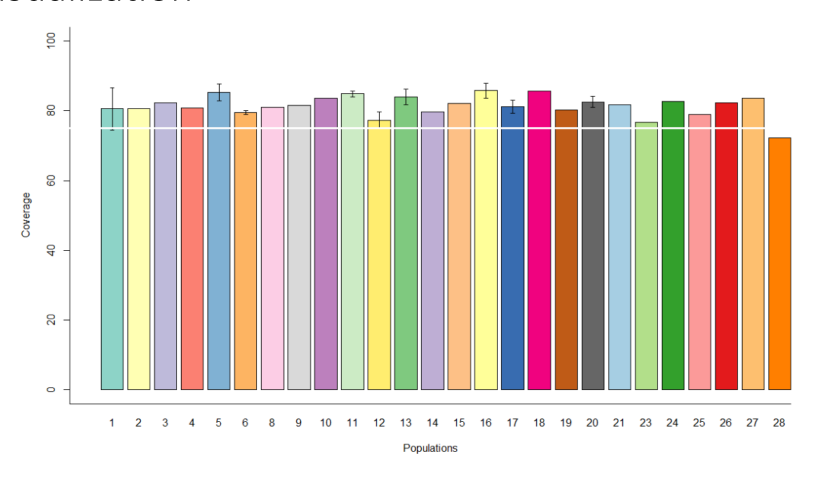
Run 'populations' to get population summary statistics such as F<sub>sT</sub> and output data into a fasta or phylip that can be used for analysis

Fasta or phylip files that can now be used to make a tree!!









# Next Steps RADSeq

- Analyze output from denovo\_map.pl
- Determine why it was failing outside the pipeline
- Take the data and make a tree
- Compare that tree to one generated in the paper I'm following
- Try it all again with Ipyrad and see if it gives similar trees

# **Bonus Question:**

Can I use the command line to automate downloading and renaming CT scanning files to save time and remove user error?

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Spoiler Alert! YES!!

