

Analyzing RADSeq Data

Jennifer Gardner

FISH 546: Bioinformatics

Question:

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

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?

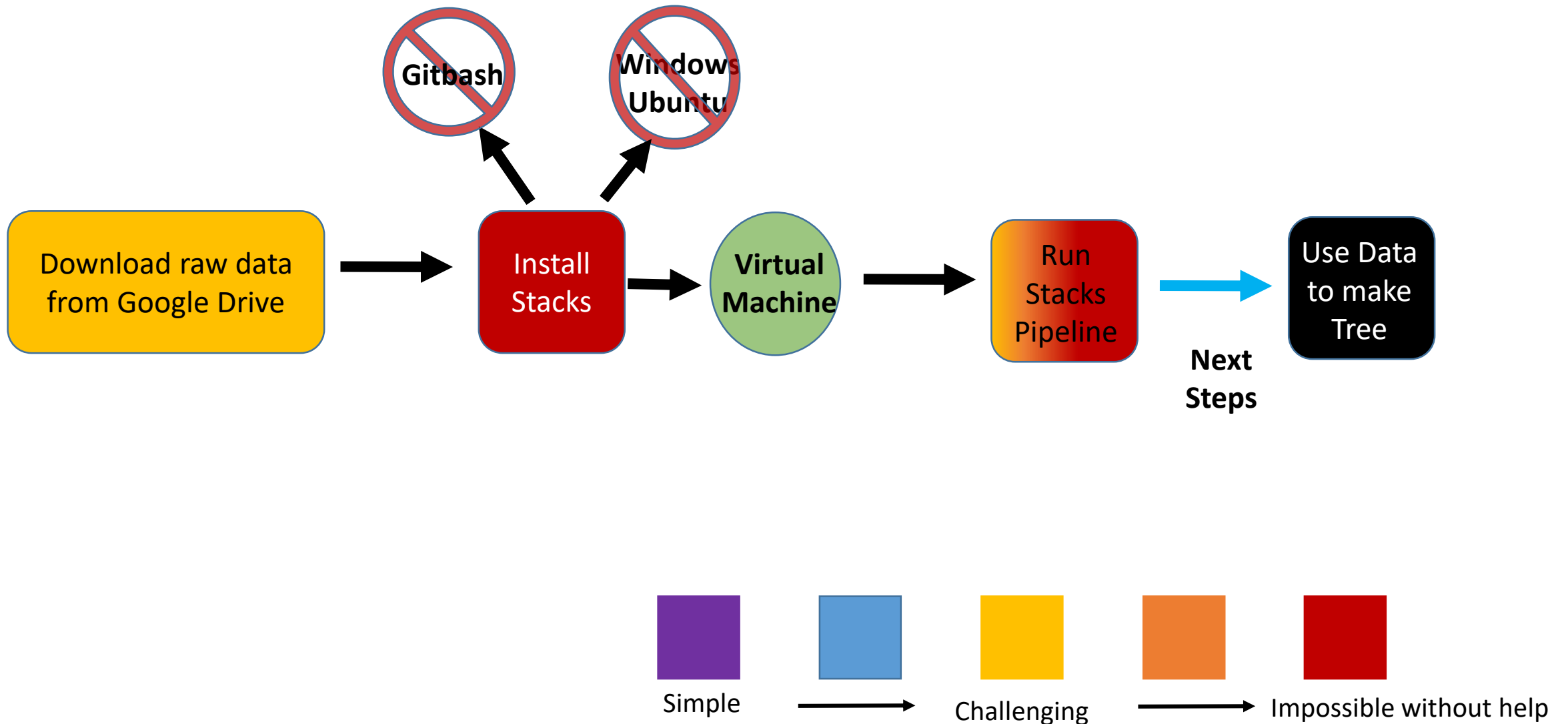
Question:

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.
 $1e5 < n\text{-reads} < 2e6$

Run `ustacks` on each
sample to find loci

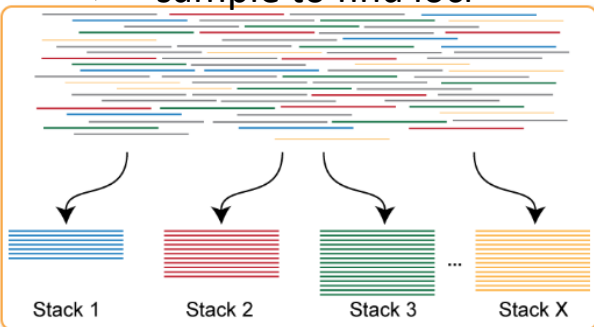


Figure http://catchenlab.life.illinois.edu/stacks/param_tut.php

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 genome)

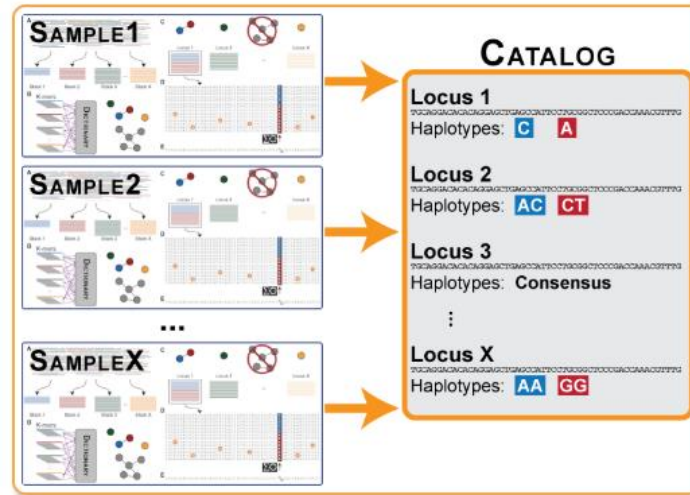


Figure http://catchenlab.life.illinois.edu/stacks/param_tut.php

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
cstacks 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

STUCK HERE

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

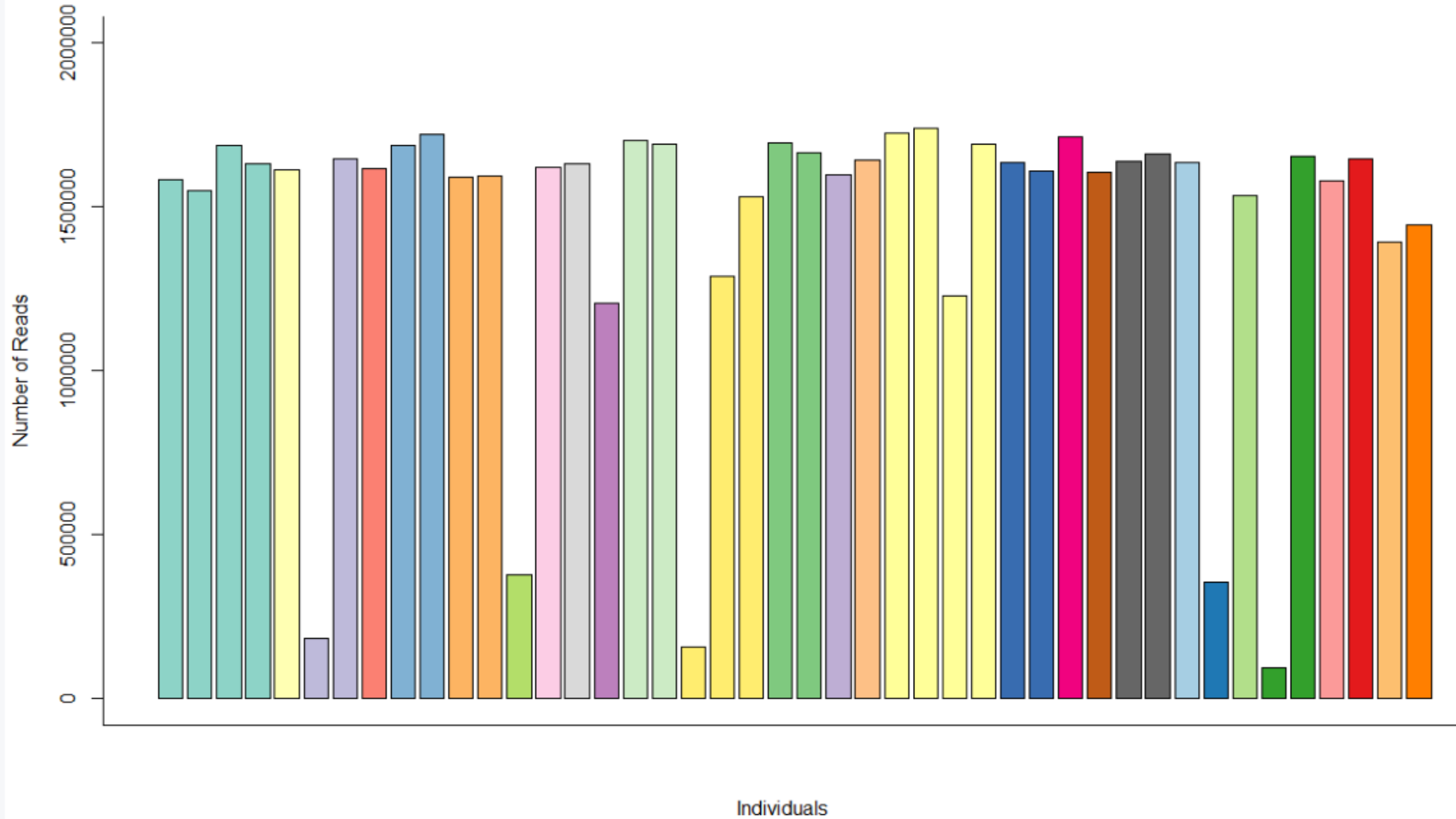
Run `gstacks` to do something?

Directory with gstacks outputs?

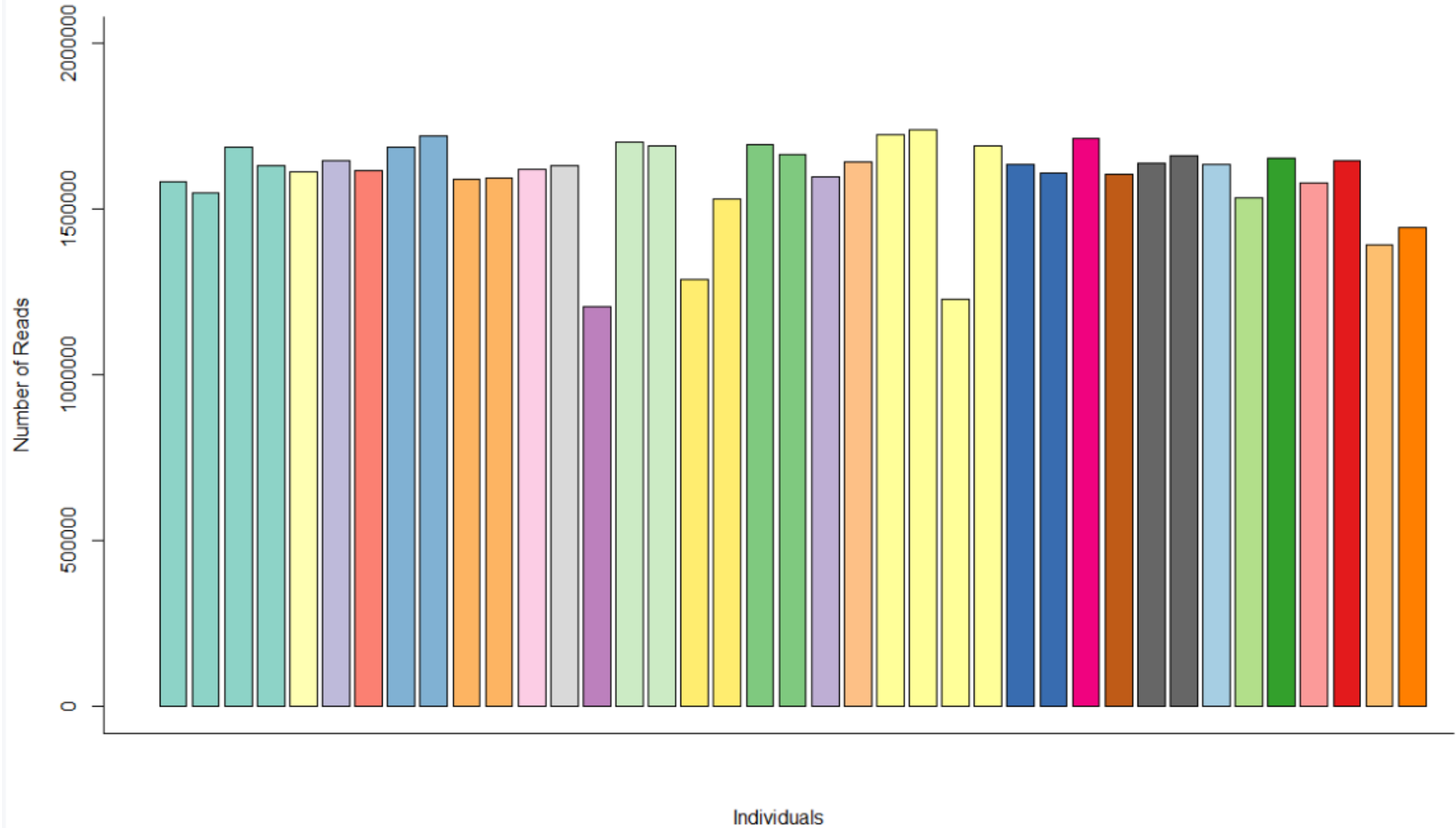
Run `populations` to get population
summary statistics such as F_{ST} and output
data into a fasta or phylip that can be used
for analysis

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

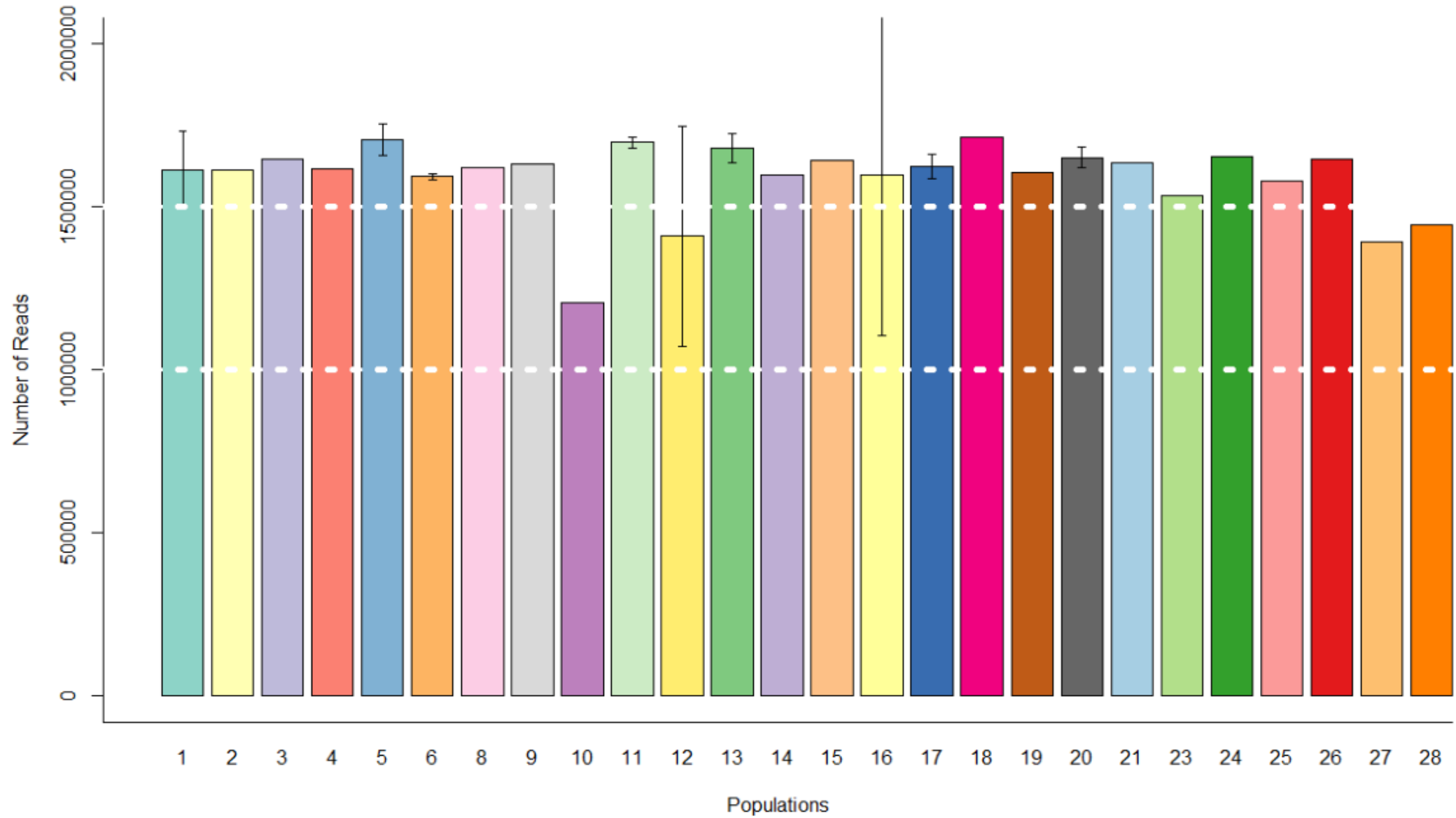
Visualization



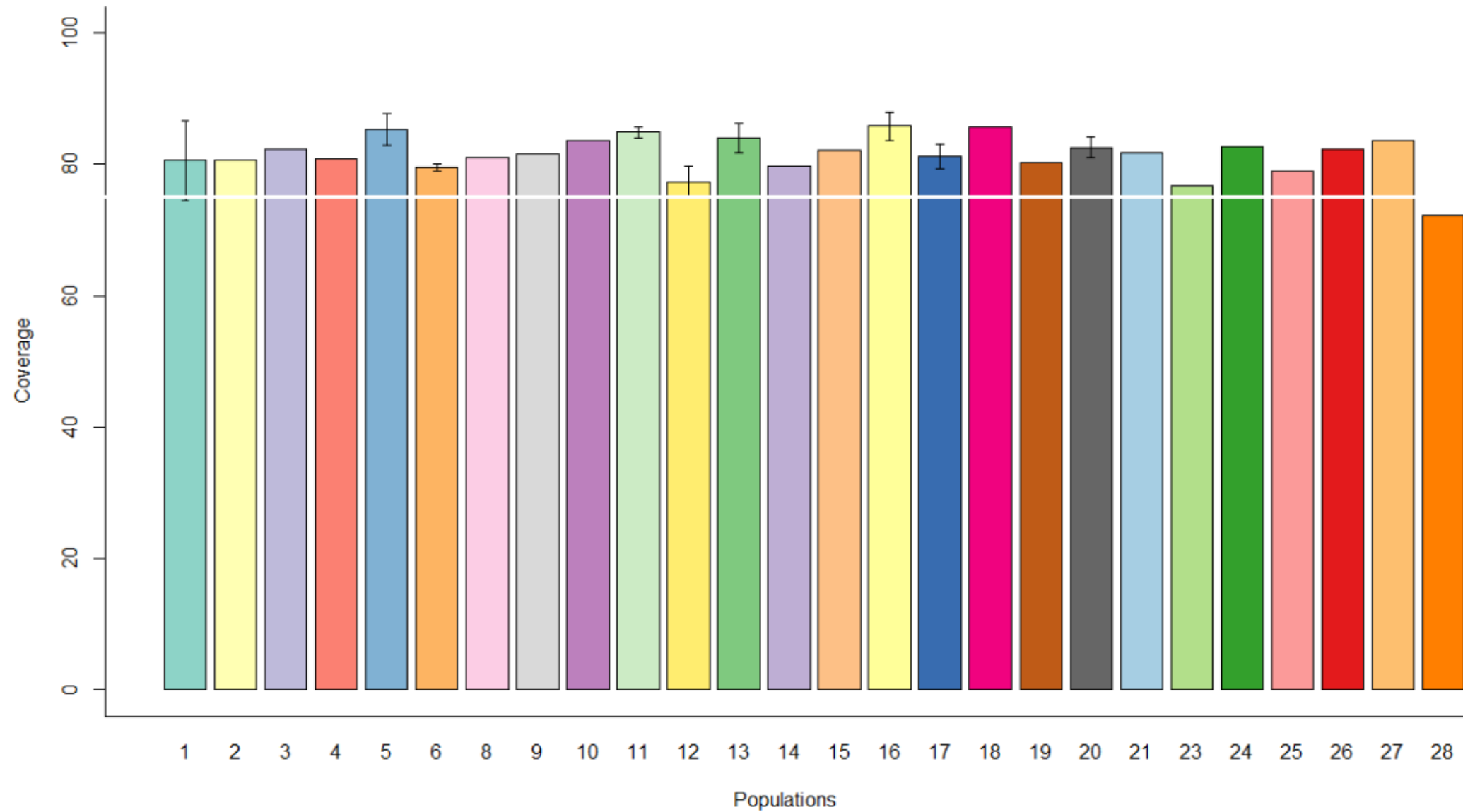
Visualization



Visualization



Visualization



Next Steps RADSeq

- Finish running 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 lpyrad 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?

Bonus Question:

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

Spoiler Alert! YES!!

