Calling genetic variants with **breseq**

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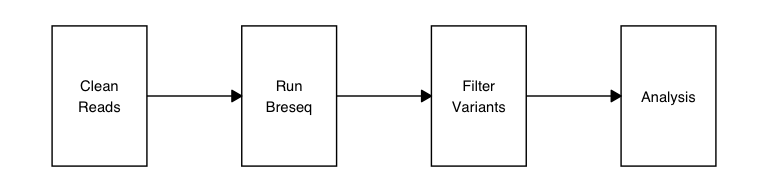
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# Outline

This protocol describes a general workflow for calling genetic variant using **breseq**, which is developed by the Barrick Lab. I like **breseq** because it is quite conservative, gives lots of intuitive output - as well as the traditional VCF file - and has really good support.

The general workflow is as follows.



# Resources

There are many resources that go into more depth than this walkthrough does here. You should go and read them! Especially if you are new to *breseq* and want to understand how it works and its output.

* breseq [documentation](https://github.com/barricklab/breseq/wiki).

# Pipeline

## Where to run this?

If you have only a couple of files and you are not running it in polymorphism mode, you can probably run this on your local computer. However, if you have a lot of files with a lot of sequencing data each (e.g. you want to call variants from an experimental evolution experiment) then I would recommend trying to run this on either the RStudio Servers (which I have done before) or a HPC at Exeter (which I should have done before).

## Install miniforge and breseq

This can be one of the most laborious, and possibly anger inducing, steps. Bioinformatics software can be notoriously difficult to install. However, there is a code chunk that can helpfully help with this. It first installs [miniforge](https://github.com/conda-forge/miniforge) that installs **conda** and **mamba** package managers. It then sets a one time configuration for which repositories to use when installing packages, prioritising **conda-forge** and **bioconda** based on [current recommendations](https://bioconda.github.io/#usage).

Firstly we install **tmux** which allows us to run multiple terminal sessions at once, and is particularly useful for running long-running commands. A cheatsheet for tmux is [here](https://tmuxcheatsheet.com/). The think I found most awkward was the Ctrl-b, then d for detach, but is easier when you realise Ctrl-b is the prefix action to all **tmux** commands.

We then create a new environment (which is a bit like a virtual machine) for all of the tools in the pipeline (actually just breseq and fastp). This is useful to avoid conflicts between different projects. You can learn more about conda environments [here](https://docs.conda.io/projects/conda/en/latest/user-guide/tasks/manage-environments.html).

# install new mamba/conda installation if needed  
curl -L -O "https://github.com/conda-forge/miniforge/releases/latest/download/Miniforge3-$(uname)-$(uname -m).sh"  
  
bash Miniforge3-$(uname)-$(uname -m).sh  
  
# update conda if needed  
conda update conda  
  
# set up channel configuration based on current best practice  
conda config --add channels bioconda  
conda config --add channels conda-forge  
conda config --set channel\_priority strict  
  
# install tmux  
conda install -c conda-forge tmux  
  
# create conda environment for breseq  
conda create -n breseq\_env -c bioconda breseq fastp seqkit

## Filter reads

Next we need to do some additional filtering of the short reads. Sequencing services do some standard filtering, but we will do our own just to make sure.

Our filtering removes: - Adapters - Median Q score, all bases > 30 - Expected read length > 95% of expected read length (e.g. for 2x 150bp reads, this would be 143bp, for 2x 300bp reads, this would be 285bp) - We check the average read length of a single file using seqkit stats to see what the expected read length is.

# activate conda environment  
conda activate breseq\_env  
  
# set working directory to where the short reads are  
wd=short\_reads  
  
# make fastp reports folder  
mkdir -p "$wd/fastp\_reports"  
  
# look at one file to look for read length  
file=$wd/307504\_NalR2\_1\_trimmed.fastq.gz  
seqkit stats $file  
  
# run fastp on all the short reads  
for file in $wd/\*1\_trimmed.fastq.gz; do  
  
 fwd=$file  
 # replace 1\_trimmed.fastq.gz with 2\_trimmed.fastq.gz to get the reverse read  
 rev=${fwd%1\_trimmed.fastq.gz}2\_trimmed.fastq.gz  
  
 #echo $fwd  
 #echo $rev  
  
 # run fastp on the file  
 fastp -i $fwd -I $rev -o "$wd/trimmed/$(basename $fwd)" -O "$wd/trimmed/$(basename $rev)" -w 4 --detect\_adapter\_for\_pe -l 237 -q 30 -j "$wd/fastp\_reports/$(basename ${fwd%1\_trimmed.fastq.gz}fastp.json)" -h "$wd/fastp\_reports/$(basename ${fwd%1\_trimmed.fastq.gz}fastp.html)"  
done

You can then look at the output in **fastp\_reports**. Or run **seqkit stats** on the filtered files.

# run seqkit on filtered files  
seqkit stats $wd/trimmed/\*.fastq.gz

| File Name | Format | Type | Num Seqs | Sum Len | Min Len | Avg Len | Max Len |
| --- | --- | --- | --- | --- | --- | --- | --- |
| 307504\_NalR2\_1\_trimmed.fastq.gz | FASTQ | DNA | 650,074 | 163,155,244 | 237 | 251 | 251 |
| 307504\_NalR2\_2\_trimmed.fastq.gz | FASTQ | DNA | 650,074 | 163,100,866 | 237 | 250.9 | 251 |
| 307505\_NalR3\_1\_trimmed.fastq.gz | FASTQ | DNA | 444,294 | 111,504,398 | 237 | 251 | 251 |
| 307505\_NalR3\_2\_trimmed.fastq.gz | FASTQ | DNA | 444,294 | 111,465,524 | 237 | 250.9 | 251 |
| 307506\_NalR4\_1\_trimmed.fastq.gz | FASTQ | DNA | 656,664 | 164,809,449 | 237 | 251 | 251 |
| 307506\_NalR4\_2\_trimmed.fastq.gz | FASTQ | DNA | 656,664 | 164,741,485 | 237 | 250.9 | 251 |
| 307507\_NalR5\_1\_trimmed.fastq.gz | FASTQ | DNA | 655,940 | 164,628,344 | 237 | 251 | 251 |
| 307507\_NalR5\_2\_trimmed.fastq.gz | FASTQ | DNA | 655,940 | 164,573,108 | 237 | 250.9 | 251 |

## Run breseq

Now we can run **breseq**. The command runs a for loop through all of the different samples, and runs **breseq** against a reference. This command is written as a task which allows us to run it in parallel. An example of running it as a simple for loop in bash is also provided.

This example is just mapping some reads back to the same reference genome, and is done on clonal sequencing data, so we are not expecting polymorphisms.

Things you will want to watch out for are: - If your reference genome is not a single contig, you might want to use the --contig-reference option. - If you have population level data, you will want to use the -p option to run in polymorphism mode. - This code assumes your forward and reverse reads are in the same folder, and their identifier is **1\_trimmed.fastq.gz** and **2\_trimmed.fastq.gz**. If this is not the case for your forward and reverse reads, you will need to change this! - **breseq** likes references in .gbk format. - **-j** is the number of threads. Set this to 8 or so if running on the RStudio server, might need to run it lower if on your local machine.

# set working directory  
wd=BASEFOLDER\_OF\_CHOICE  
  
# set reference genome  
ref=$wd/reference/reference.gbk  
  
# set output folder  
output\_folder=$wd/breseq  
# make folder if it does not exist  
mkdir -p $output\_folder  
  
# write up the breseq command in a task  
task(){  
 # assign fwd and rev files  
 file\_fwd="$1"  
  
 # assign fwd and rev files  
 file\_fwd="$1"  
 file\_rev=${file%\_1\_trimmed.fastq.gz}\_2\_trimmed.fastq.gz  
  
 #stub  
 stub=$(basename ${file%\_1\_trimmed.fastq.gz})  
  
 echo $stub  
  
 mkdir -p $wd/breseq/$stub  
  
 # run breseq - there are two contigs so run them in -c contig mode  
 breseq $file\_fwd $file\_rev -j 4 -o $stub -r $ref  
}  
  
# run one instance of this   
# lets do 6  
N=1  
  
# try and run it on 6 instances at once  
(  
for file in $trimmed\_files/\*val\_1.fq.gz; do   
 ((i=i%N)); ((i++==0)) && wait  
 task "$file" &   
done  
)