Variant Calling

Goal: Learn how to use various tools to identify variants after re-sequencing

Input(s): Mo_FR13_IP{1-3}_accepted_hits.bam

Output(s): Mo_FR13_alignments_merged.bam

Moryzae_FR13_alignments.bcf Moryzae_FR13_alignments.vcf

Three of the RNA samples used for RNAseq analysis were generated from fungal strain FR13, which is expected to show several sequence differences relative to the 70-15 strain that was used to generate the reference genome. Therefore, we will use the RNAseq alignments to search for nucleotide differences between FR13 and the 70-15 reference strain.

7.1 Merge the FR13 alignment (.bam) files

Having confidence in variant calls requires one to have multiple reads aligned across the region where there is a genetic difference between the sample and the reference. When sequencing DNA, this is easily accomplished by ensuring that we have enough sequence data to provide several-fold coverage of the genome (≥ 20x). However, with RNAseq data, we have no control over coverage because this is determined by the expression level of each gene at the time and place where the RNA was extracted. Therefore, to maximize coverage across each gene for the purpose of our analysis, we will merge the alignments file for the three *in planta* RNA replicates.

- ☐ We will use the **samtools** merge tool. (Note that bam files must be sorted by coordinates before they are merged. We already sorted the files in the *maseq* lab, so we can use them directly here.)
 - samtools merge Mo_FR13_alignments_merged.bam
 - ../rnaseq/alignments/*FR13*bam

	Take a quick look at the .bam file. Is it in binary format? You can take a look at it by using samtools again:		
	• samtools view Mo_FR13_alignments_merged.bam		
	☐ Whoa, can you read the output? Quit the process (control-c) and re-run the command with into another program that will enable you to look at a few lines at a time.		
	Does the output still look like it contains sam alignment data? If so, proceed to the following.		
	We must provide mpileup with the reference genome that was used for alignment. To speed up its computations, mpileup uses an indexed version of the genome. This index is different to the one generated with bowtie2 , and we will use samtools to generate it.		
	• samtools faidx/blast/magnaporthe_oryzae_70-15_8_supercontigs.fasta		
	This will create an index file named magnaporthe_oryzae_70-15_8_supercontigs.fasta.fai.		
☐ Check to make sure the index was created. (Note that the file is created in the same or reference genome.)			
7.2 F	Perform the variant calling		
insertio informa	I use the bcftools mpileup utility to extract information on nucleotide variations (substitutions, ns, deletions) between our sequence sample and a reference genome. mpileup does this by using ation contained in the CIGAR (Concise Idiosyncratic Gapped Alignment Report) string and the (back track operations) string in the MD:Z: field of the .sam (or .bam)file.		
Usage:	ols mpileup [options] -f <reference_genome> <bam alignment(s)=""></bam></reference_genome>		
	Make sure you are using screen .		
	Run mpileup on the sorted file.		
	• bcftools mpileupthreads 2 -f/blast/magnaporthe_oryzae_70-15_8_supercontigs.fasta Mo_FR13_alignments_merged.bam		
	threads number of processors to use		
	-f [FILE] name of reference genome		
	ls, like samtools, outputs results to the screen, making the output kind of hard to read. However, at u can see that the program is doing something interesting.		
	Stop the process (using control-c) so you don't have to wait a LONG time for the output to finish printing to the screen.		

- □ Re-run the previous command but redirect the results to a file called *Mo_FR13_alignments.vcf*. (Note: the program will take several minutes to complete, during which time you will receive no progress updates—all program outputs are directed to the specified output file.)
 □ Inspect the *Mo_FR13_alignments.vcf* file. This summarizes variant statistics for every position in the reference genome for which there are aligned reads. However, it does contain variant calls.
 □ We will use **bcftools** to call the variants. **bcftools** is like **samtools** in that it sends results to the
 - bcftools call -v -c --ploidy 1 Mo_FR13_alignments.vcf > Mo_FR13_alignments_called.vcf
 - -v output potential variant sites only (i.e., skip monomorphic ones)
 - -c call variants using the original method implemented in samtools mpileup
 - **--ploidy** set to 1 because *Magnaporthe* is haploid, and we only expect one copy of each gene unless the sequence is repeated

7.3 Examine the resulting variant calls

screen so we need to re-direct the output to a file.

☐ Inspect the *Mo_FR13_alignments_called.vcf* file.

At the head of the file is some information on how to interpret the various fields. Below, each line provides information on a predicted variant at a specific nucleotide position within the chromosome. Here you should be able to recognize data that make sense.

Unfortunately, the header does not provide much information on the overall structure of the .vcf file. The main fields in the tab-delimited section are as follows:

Column 1: Chromosome number

Column 2: Nucleotide position

Column 3: SNP ID (if previously characterized and named)

Column 4: Nucleotide in reference genome

Column 5: Alternate allele(s) identified in sequence reads

Column 6: Quality of SNP call

Column 7: Filtering information (":" = no filter; "Low Qual"; or "PASS")

Column 8: SNP information (see list of INFO fields in file header)

Columns 9 & 10: SNP formats (see the list of FORMAT fields in file header)

A complete description of the VCF format is in the VCFv4.1.pdf file on the Canvas site.

Column 10 in the Mo_FR13_alignments_called.vef file contains information that informs us about the likelihood that a given SNP (or INDEL) is valid. First, to have confidence in the call, we want to have several reads that support it, and because we are working with a haploid fungus, ideally we want all reads that overlap a given site to have the alternate (variant) allele. Unfortunately, variant callers are not very smart when operating in haploid mode (they were mostly designed for human variant calling). As a result, they often call variants at sites that are clearly heterozygous for the reference/alternate alleles. Heterozygosity in a haploid organism indicates that there are two or more copies of the site being interrogated but, more importantly, it tells us that the SNP is likely to be false. There are many instances of heterozygosity for variant calls in the Mo_70-15_FR13.vef file, so we need to filter the file to remove these suspect calls.

	5 -	file, so we need to filter the file to remove these suspect calls.	
	10. (Note: this is a w-i-d-e co-supporting ref allele, reverse s	ew pages of the vcf file and pay attention to the DP4= field in column lumn!) There are four values that follow DP4=: forward strand reads strand reads supporting ref allele, forward strand reads supporting the supporting the alt allele. For a true SNP, there should be no reads at	
	of low quality calls where the values for one such suspect re	umber of SNP calls (don't count INDELs). Next, count the number re are some reads supporting the reference allele, position, and DP4=ecord. ; Low quality (suspect) SNPs:	
confide	ent that these variant calls aren	were called were supported by very few reads, and so we can't be it due to sequencing errors. Therefore, we will filter the SNP calls to ted by at least 10 reads, with no reads containing the ref allele.	
	*	is purpose, but why waste our time trying to remember the necessary is we can easily do this with grep ?	
	• grep DP=[1-9][0-9 > Mo_FR13_alignme] Mo_FR13_alignments_called.vcf grep DP4=0,0, nts_filtered.vcf	
	grep DP=[1-9][0-9]	look for "DP=" followed by any number between 1 and 9, followed by any number between 0 and 9. (This is how we make sure that DP= is followed by a number that is equal to, or greater than, 10.)	
	ne the resulting filtered file cor o answer the following questio	ntaining "high quality" variant calls, and use the suggested tools with ns:	
	How many SNPs were identified on Chromosome 8.1? (grep)		
	How many INDELs were found on Chromosome 8.1? (grep)		
	Are there any indels that are greater than one nucleotide in length? (awk {print length()}, sort uniq)" yes; no		
	W/I	f acres on (DD-) for a CND in the filtered detect) (arrive and cont	
Ц	uniq)	f coverage (DP=) for a SNP in the filtered dataset? (awk, sed, sort,	

Appendix

The variant call format

This is a standard text file format used in **bioinformatics** for storing information about **genetic variants** (like SNPs, insertions, deletions, etc.) identified in DNA sequencing data. It has a header section (lines starting with ##) which documents the mpileup and variant call commands that were used. This also provides information about the reference genome used, as well as the various data "tags" that describe the data. This is followed by the data section which lists the sites on the reference where variants have been found as well as various statistics that support each call. These statistics are denoted with shorthand tags, short descriptions of which can be found in the header.

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The various data columns are as follows:

CHROM	Chromosome (e.g., chr1)
POS	Position on the chromosome

ID Variant ID (e.g., official rsID in dbSNP)

REF Reference base(s)
ALT Alternate base(s)

QUAL Quality score for the variant (PHRED scale)
FILTER Filter status (e.g., PASS or reason for filtering out)

INFO Additional information (e.g., read depth: DP=100; BQBZ=base quality bias) FORMAT Format of the genotype fields (e.g., GT:PL:AD = genotype:phred

likelihood:allelic depths)

sample1, sample2... Genotypes for each sample (0=ref; 1=alt; 0/1 =heterozygous). For our

analysis we only have one genotype column but when there are multiple

samples, there will be one genotype fields per sample.