

Mapping and Variant Calling (June 2016)

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PhD course: High throughput sequencing of non-model organisms

Contents

1 Mapping with Bowtie2	1
1.1 The mapping commands	2
1.2 Running the commands in a script for posterity	4
2 Filter mappings	6
2.1 Removing duplicate reads	6
2.2 Re-alignment around indels	7
3 Visualizing alignments	7
3.1 Samtools tview: command-line viewer	7
3.2 IGV: viewer with a graphical user interface	9
4 BONUS: SNP calling with samtools mpileup and bcftools	10

Once you have assembled a draft genome, you can map your reads against it and identify potential read variants, like SNPs (Single Nucleotide Polymorphisms) and InDels (Insertions and Deletions). The library that you sequenced in this course is likely to represent a single diploid individual. Read variants, thus, occur at loci that differed between the mother and the father of this individual. Read variants become interesting when they are associated with certain environmental factors or phenotypes. This, however, requires sequencing several individuals of differing phenotypes or obtained from several environments.

1 Mapping with Bowtie2

Mapping next-generation sequencing data against genomes requires very high performance algorithms that must balance the accuracy of the mapping with the time taken. This is an interesting problem to work on and a rather large number of different applications have been developed. To map reads against the assembled draft genome, we will use [Bowtie2](#). Bowtie2 defaults to finding global alignments (no soft-clipping of terminal bases) and it allows for gaps in the alignment, thereby increasing mapping accuracy ([Schlotterer *et al.* \(2014\) *Nature Reviews Genetics*](#)).

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A number of aligners have been written to either compensate or make use of the specific properties of sequence data obtained using different technologies. For example the aligner [TMAP](#) was specifically designed to be used with Ion Torrent data which has problems with resolving homopolymer lengths accurately. Similarly there are specific aligners that handle SoliD color-space data which cannot be effectively aligned as nucleotide sequences. In this tutorial we will make use of Bowtie2 as it is one of the most commonly used aligners that balances speed, accuracy and memory requirements. My current favourite aligner is the [STAR aligner](#), which was designed to align RNA sequences to genomic locations; it is rather accurate and very fast, but requires far more memory to run than bowtie/bowtie2. An alternative aligner that is also widely used is the [BWA](#) aligner. You can find an unpublished comparison of mapping performance [here](#), a benchmarking of 9 mapping tools [here](#) and a more in-depth look at mapping RNA sequences to genomes [here](#).

1.1 The mapping commands

What you need for mapping are two sequence files, one containing the quality-trimmed reads (fastq format) and the other containing the draft genome (in the fasta format).

Create a new folder named **Mapping** (with `mkdir`) and copy the two sequence files into it (with `cp`). For example for the assembly made using IonTorrent data:

```
1 mkdir Mapping
2
3 cp GenomeAssembly/IonTorrentDeNovoAssembly_assembly/IonTorrentDeNovoAssembly_d_results/\
4 IonTorrentDeNovoAssembly_out.unpadded.fasta \
5 Mapping/
6
7 cp RAWREADS_trimmed_trimmed.fq \
8 Mapping/
9
10 cd Mapping
```

Here the `RAWREADS_trimmed_trimmed.fq` refer to the IonTorrent reads after quality trimming using TrimGalore. Here we are simply making copies of the various files in different locations for performing different analyses. This is simple to describe and understand, but is not a particularly good way to organise your files, both because it wastes disk space¹ and (more seriously) it makes it easy to end up with files with the same names, but with slightly different content².

In the Mapping folder, we first create an index for the reference genome using the following command (enter the folder using `cd` before calling this command) :

```
1 bowtie2-build -f IonTorrentDeNovoAssembly_out.unpadded.fasta GENOMENAME
```

You should change `GENOMENAME` to something that is suitable for your data.

¹In some modern file systems that make use of Copy-On-Write, there won't actually be any copying of the data unless one of the files is modified.

²How to handle lots of files created by various versions of data flows or pipelines is not a simple problem and there are many systems that have been developed to address such problems. This comes under the general heading of version control systems, and is outside the scope of this course. In general though, it is wise to follow rule number one of database design: 'never store a piece of information in more than one location'. I.e. don't copy stuff around like we are doing here.

Now, to align the trimmed reads against the genome, use the following command:

```
1 nohup bowtie2 -p 1 \  
2 -q \  
3 --phred33 \  
4 --sensitive \  
5 --no-unal \  
6 --al-gz FILE_Aligned.fq.gz \  
7 --un-gz FILE_Unaligned.fq.gz \  
8 --met-file MetricsFile.txt \  
9 -x GENOMENAME \  
10 -U RAWREADS_trimmed_trimmed.fq \  
11 -S MAPPEDREADS.sam >Logfile.log &
```

Here an overview of the meaning of the used options:

-p 1 Causes Bowtie 2 to use a single thread. Depending on the number of users and libraries we will probably increase this.

-q Informs the program that the reads to map are saved in fastq files.

--phred33 Sets the quality encoding of the fastq files to "Phred+33".

--sensitive sets several options at once regarding the seeding and other adjustments.

--no-unal Suppress SAM records for reads that do not align.

--al-gz Write unpaired reads that align at least once to to the specified file.

--un-gz Write unpaired reads that failed to align to the specified file.

--met-file Write bowtie2 metrics to Metricsfile.txt.

-x Specifies the name of the genome.

-U Specify the unpaired reads to align (can contain a comma-separated list of several fq files).

-S Specify the sam file to which the alignment shall be saved.

You can't set the exact number of mismatches in the seed, but you can adjust the mismatch penalty.

The program should run no longer than 10-20 mins. The resulting output file will be in the SAM format. For a detailed description of this format, see [here](#).

To map the Illumina data we follow a similar procedure; however, we need to modify the call to **bowtie2** as the Illumina data is paired ended. To find out how we can do this, we can run **bowtie2** without any arguments or specifying the **--help** option. This will print out the usage information. Knowing how to read usage information is one of the most important things you can do as you'll then be able to run most applications without relying on others. If you do this, you'll see something like this:

```

1 bowtie2 --help
2 Bowtie 2 version 2.1.0 by Ben Langmead (langmea@cs.jhu.edu, www.cs.jhu.edu/~langmea)
3 Usage:
4   bowtie2 [options]* -x <bt2-idx> {-1 <m1> -2 <m2> | -U <r>} [-S <sam>]
5
6   <bt2-idx>  Index filename prefix (minus trailing .X.bt2).
7             NOTE: Bowtie 1 and Bowtie 2 indexes are not compatible.
8   <m1>       Files with #1 mates, paired with files in <m2>.
9             Could be gzip'ed (extension: .gz) or bzip2'ed (extension: .bz2).
10  <m2>       Files with #2 mates, paired with files in <m1>.
11             Could be gzip'ed (extension: .gz) or bzip2'ed (extension: .bz2).
12  <r>        Files with unpaired reads.
13             Could be gzip'ed (extension: .gz) or bzip2'ed (extension: .bz2).
14  <sam>      File for SAM output (default: stdout)
15
16  <m1>, <m2>, <r> can be comma-separated lists (no whitespace) and can be
17  specified many times.  E.g. '-U file1.fq,file2.fq -U file3.fq'.
18
19 Options (defaults in parentheses):
20
21 Input:
22   -q          query input files are FASTQ .fq/.fastq (default)
23   --qseq      query input files are in Illumina's qseq format
24   .... more options

```

Let us consider the top lines first. This is the basic usage information that tells you the arguments you need to specify and their order.

```

1 Usage:
2   bowtie2 [options]* -x <bt2-idx> {-1 <m1> -2 <m2> | -U <r>} [-S <sam>]

```

Things contained in square brackets [stuff in square brackets] denote optional arguments. So, the above (bowtie2 [options] ...) indicates that optional options (specified with - or --) should be specified before other arguments. After these options (of which there may be none) you should specify the value of the -x option. Looking down, you can see that <bt2-idx>, is a placeholder for the name of the index that you built using bowtie2 in the previous section. If you have assembled a genome from the Illumina data on its own this will be a different index file based on a different assembly sequence, so we will need to change this value.

The next section of the usage line is contained in squiggly brackets (usually referred to as braces) indicating that you have a choice of two or more alternatives. These alternatives are separated by the pipe (|) character which in computing languages is usually taken to mean 'or'. So the section {-1 <m1> -2 <m2> | -U <r>} reads as 'either specify the values of -1 and -2 or the value of -U. Looking at the explanation further down, you can see that <m1> and <m2> refer to mate or paired sequences, whereas <r> refers to unpaired reads. The last section simply specifies to which file we wish to write the output; if you don't specify this, it will simply be written to the terminal (i.e. STDOUT). This is useful, because we can then pipe the data to other applications in a single command.

So reading the usage line (also known as the synopsis) we can design our command line. If our paired reads are in files RAWREADS_fw_trimmed_trimmed.fq and RAWREADS_rv_trimmed_trimmed.fq, and the index for our assembly genome is in GENOMENAME.X.bt2, the command without any of the optional options would be:

```

1 bowtie2 -x GENOMENAME -1 RAWREADS_fw_trimmed_trimmed.fq \
2 -2 RAWREADS_rv_trimmed_trimmed.fq -S MAPPED.sam

```

Here we haven't specified any of the options we used for the IonTorrent data above and the

program will simply use the default options. To see what the default options are you should read the rest of the help section that is printed out when you run `bowtie2` without any arguments. You can probably use most of the options as we used above, though you should not assume this.

Given that the Illumina data is paired end sequence data you should pay special attention to the Paired-end section of the help text. In particular consider the values of `-I` and `-X` and whether the default options are reasonable for your libraries.

1.2 Running the commands in a script for posterity

As was emphasised in the section on Unix tools for bioinformatics, you really shouldn't type these commands directly into a terminal window. It's too easy to make a mistake when you have to specify many options, and you will not have a record of the command that you actually used. Instead we will write the commands into a text file and ask the shell (in this case `bash`³) to run the commands non-interactively. In the simplest case you just make one file for each command, and run these separately. However, it is much better to embed the full process into a single script as all the information will be in a single place.

Here what we have done is:

- made a directory for our mapping (`mkdir`)
- copied the data files to that directory (`cp`)
- entered the directory (`cd`)
- run `bowtie2` to make an index
- run `bowtie2` to map the sequences

We can put all of those commands into a single shell script, or we can make the directories manually and only include the more complicated commands in the script. Which is better depends a little bit on the situation; if you have lots of different sequence files that you wish to map in different ways then you might want to put all the directory commands into the script; ideally doing this in an automated way using loops and assembling the directory names automatically. However, here I would suggest the simple option of manually making the directories and having simpler script files to avoid using more complex shell scripting.

Hence once you have created the appropriate directory and copied the sequence files (as above) you can write (eg: `nano pgm_map.sh`) a script (to map IonTorrent data) that looks a bit like:

³bash stands for Bourne Again Shell, and is a bit of a joke on the fact that Bash is an extension or enhancement of the Bourne shell. These days it's probably the most common shell used, but as always there are people who consider it an abomination.

```

1  #!/bin/bash
2
3  ## here you can define some variables that specify the names of
4  ## input and output files
5
6  RAWREADS=breiflabb_pgm
7  GENOMENAME=breiflabb_pgm
8  FILE="$GENOMENAME"_bt2
9
10 ## note that when you use the variables you have to put a $
11 ## sign in front of them
12 ## and if you want to concatenate to words you need to
13 ## to quote them
14
15 ## first build the index:
16 bowtie2-build -f IonTorrentDeNovoAssembly_out.unpadded.fasta $GENOMENAME
17
18 ## then use that to map the sequences:
19 bowtie2 -p 1 -q -phred33 --sensitive --no-unal \
20 --al-gz "$FILE"_Aligned.fq.gz --un-gz "$FILE"_Unaligned.fq.gz \
21 --met-file MetricsFile.txt \
22 -x $GENOMENAME -U $RAWREADS_trimmed_trimmed.fq \
23 -S "$GENOMENAME"_bt2_mapped.sam" > bt2_log.log
24
25 ## here you can put some comments to indicate what the different
26 ## options indicate and why you have chosen the specific options.

```

To run this script (pgm_map.sh) you can either :

```

1  bash pgm_map.sh

```

2 Filter mappings

To remove unmapped reads, reads below a mapping quality of 20, and reads that were not aligned uniquely (reads that were mapped to >1 places in the genome), use the python script [Bowtie2Filtering.py](#):

```

1  Bowtie2Filtering.py -mq -u -a -s MAPPEDREADS.sam

```

Your filtered reads will be saved in **MAPPEDREADSfiltered.sam**

Alternatively, you can use [samtools](#) to filter out reads with a mapping quality <20:

```

1  samtools view -Sh -q 20 -o MAPPEDREADS_QualityAbove20.sam MAPPEDREADS.sam

```

Options:

-S Input is in the sam format

-h Include the samfile header in the output

-q Skip alignments with a mapping quality below 20

Note that it is usually possible to limit the alignments reported by the mapping program by adjusting the options; for at least some programs you can instruct the program to only report unique matches and so it might seem unnecessary to perform post-filtering steps like these.

However, given that the mapping process takes far more time than the filtering process it often makes sense to map using permissive criteria and then to filter these depending on the questions being asked.

2.1 Removing duplicate reads

After quality-trimming, we counted the fraction of duplicate reads. Duplicate reads have the same start and end coordinates and map to the same region. Duplicates result from primer or PCR bias towards these reads. As they can skew genotype estimates, they should be removed before SNP calling.

To remove duplicates, we will use 'MarkDuplicates' from the [Picard command line tools](#). An alternative tool is [samtools rmdup](#), which considers single-end reads to be duplicates when their mapping locations are the same - even if the base composition differs between the reads.

First, we need to convert our sam file to a bam file (a binary, compressed version of a sam file that is not human-readable) and sort the reads by the leftmost mapping coordinates.

```
1 samtools view -bSh MAPPEDREADS.sam > MAPPEDREADS.bam
2 samtools sort MAPPEDREADS.bam MAPPEDREADS_sorted
```

Meaning of the options:

-b output in bam format

-S input in sam format

-h include the header in the output

Then, you can use the java script 'MarkDuplicates.jar' from Picard tools to remove the duplicates from the sorted bam file:

```
1 picard-tools MarkDuplicates \
2 INPUT=MAPPEDREADS_sorted.bam \
3 OUTPUT=MAPPEDREADS_dedup.bam \
4 METRICS_FILE=MAPPED_metricsfile \
5 ASSUME_SORTED=true \
6 VALIDATION_STRINGENCY=SILENT \
7 REMOVE_DUPLICATES=true
```

Duplication metrics will be written to the `MAPPED_metricsfile`. We again very strongly recommend that you put these commands into a shell file and run that rather than to run directly from the command line.

2.2 Re-alignment around indels

Reads that are spanning InDels are often misaligned and can result in false SNPs (see [Schlotterer et al. \(2014\) Nature Reviews Genetics](#)). These reads should be removed or re-aligned. We have not enough time to re-align the reads in this course but the required steps (using [GATK](#)) are described in detail here: <http://sfg.stanford.edu/SFG.pdf>.

3 Visualizing alignments

3.1 Samtools tview: command-line viewer

The command line tool samtools tview allows you to view your alignments directly in the command line window. What you need is the reference genome (fasta file) and the sorted and deduplicated alignment file (bam file). First, you need to index the bam file before using **samtools tview**:

```
1 samtools index MAPPEDREADS_dedup.bam
2
3 samtools tview MAPPEDREADS_dedup.bam \
4 IonTorrentDeNovoAssembly_out.unpacked.fasta
```

Fig. 1 shows a screenshot of tview. When you hit the **?** on your keyboard, you will see the range of options to navigate through the alignment. You can change the contig that you are looking at by hitting **g** and then enter in the Goto-window the name of the contig, like **IonTorrentDeNovoAssembly_c3**. You can exit the alignment viewer by hitting **q**.

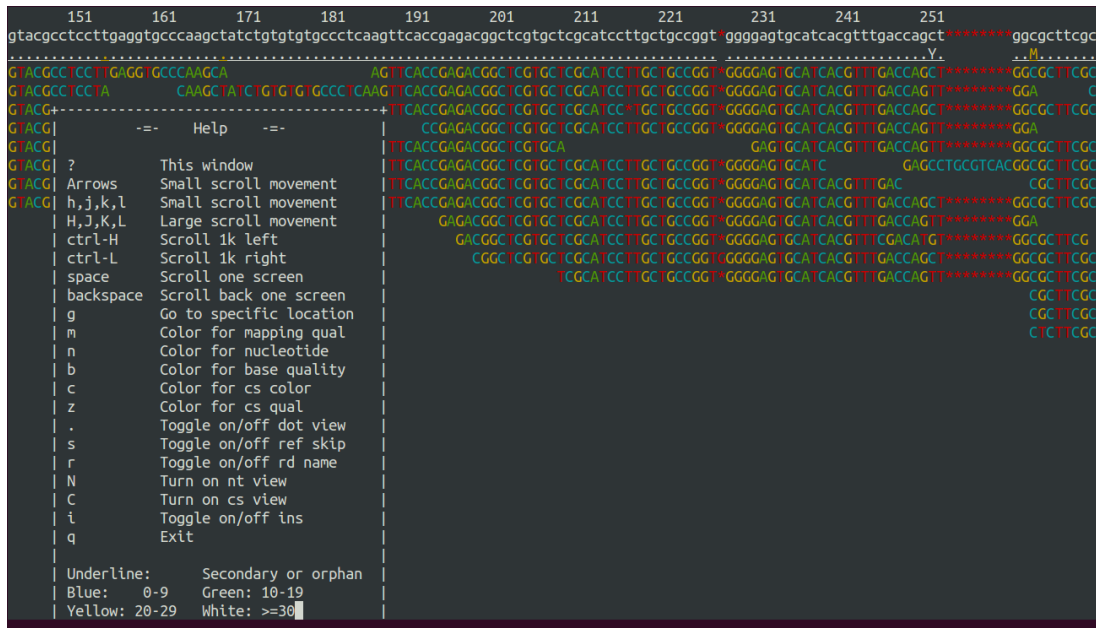


Figure 1: Screenshot of tview

3.2 IGV: viewer with a graphical user interface

I bet that many of you prefer to look at the alignment in a graphical user interface. A decent free alignment viewer is [igv](#), the Integrative Genomics Viewer (see Fig. 2 for a screenshot). Once you have registered, you can launch the program with Java Web Start. We can't promise that this works well in the course, since everything that relies on a graphical user interface can be quite slow when using a remote connection. Thus, you might want to download the required files (deduplicated SAM file and reference genome) and try out igv on your private computer. The interface is pretty much self-explanatory. To look at the alignment, you first need to load a genome and then add the mapped, sorted and indexed bam file.



Figure 2: Screenshot of igv with reads aligned to a reference and colored mismatches

4 BONUS: SNP calling with samtools mpileup and bcftools

Given sequences aligned to a reference it seems that it should be trivial to identify sequence variants. Surely any mismatches between the reference (in this case our assembly) and reads is evidence for the presence of a sequence variant. However, if the probability of observing a sequencing error is larger than the frequency of sequence variants within the population (an individual can be considered as a population of two haploid genomes) then most sequence mismatches will be caused by sequencing errors. This is usually the case (and overwhelmingly so) when looking at individuals from within a single species and in order to identify a position as a sequence variant we need to have more than one read diverging from the reference. How many reads are required depends on the total number of reads, the qualities of those reads and the expected variant frequency. If we are sequencing populations, then we also have to consider the rarity of a given allele; the rarer the allele one wishes to discover the larger the sequencing coverage required. This has led to the development of a rather large number of variant detection algorithms and programs (see [table 3](#) of [Schlotterer *et al*](#) for a list), and the difficulty of balancing computation times, sensitivity and accuracy makes it likely that more methods and or implementations will be written.

Here we will use the `samtools mpileup` in conjunction with `bcftools`. Computationally these are some of the simplest ways to detect variants and are widely used. For more in depth analyses we would recommend that you consider using other tool sets that have the potential to provide more accurate variant detection at the cost of more processing time.

The tool `samtools mpileup` defaults to creating a pileup file, which summarizes aligned base calls in text format (See [here](#) for an overview of its options, and [here](#) for a detailed characterization of a pileup file <http://samtools.sourceforge.net/pileup.shtml>). If you call `samtools mpileup` with the `-u` or `-g` option the output format is a vcf or bcf (compressed binary version of vcf) file; vcf stands for 'variant call format'. Its format specifications are described [here](#) and summarized in [Fig. 3](#).

The first step for calling SNPs from your aligned and deduplicated reads is:

```
1 samtools mpileup -g \  
2 -f \  
3 IonTorrentDeNovoAssembly_out.unpadded.fasta \  
4 -q 20 \  
5 -Q 20 \  
6 -t DP \  
7 -t SP \  
8 MAPPEDREADS_dedup.bam > MAPPEDREADS_dedup.bcf
```

The chosen options are described on this [page](#). By setting the `-t SP` and `-t DP` tags, `samtools mpileup` provides:

`-t SP` per-sample Phred-scaled strand bias P-value

`-t DP` per sample read depth

To call SNPs from the bcf file, we use `bcftools`:

```
1 bcftools call -vm -V indels MAPPEDREADS_dedup.bcf > MAPPEDREADS_variants.vcf
```

Options:

-v Output variant sites only

-V indels Skip indels

-m model for multiallelic and rare-variant calling

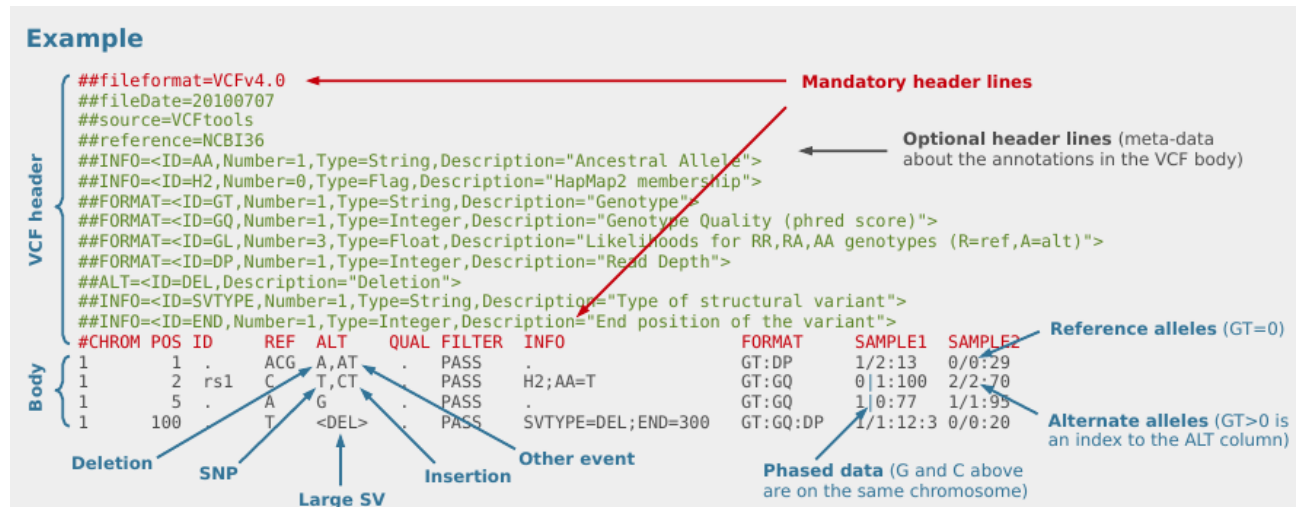


Figure 3: VCF file overview from Petr Danecek

To count how many SNPs were found, use the following command:

```
1 grep -v -c '^#' MAPPEDREADS_variants.vcf
```

The option -v in combination with ^# excludes all header lines that start with (^) the #-sign. With the -c option, grep counts the lines instead of writing them out.

To filter out SNPs that are low quality or covered by low depth, we can use the vcfutils.pl varFilter that comes with samtools:

```
1 vcfutils.pl varFilter -d 5 -w 3 -Q 20 MAPPEDREADS_variants.vcf > MAPPEDREADS_variants_filtered.vcf
```

Options used:

-d 5 minimum read depth of 5

-w 3 SNP within 3 bp around a gap to be filtered. This may be an alternative solution to re-alignment around indels

-Q 20 minimum mapping quality of 20

Another useful option can be:

-1 0.0001 min P-value for strand bias (given the PV4-tag in the vcf file). We obtained the PV4-tag by setting the -t SP tag in samtools mpileup. This option filters out the SNPs that have a strong strand-bias: SNPs that are supported by one strand and not the other.

Count how many SNPs are left after filtering

```
1 grep -v -c '^#' MAPPEDREADS_variants_filtered.vcf
```

The SNPs can be visualized with IGV. For this, we first need to compress and index the vcf files:

```
1 bgzip -c \  
2 MAPPEDREADS_variants_filtered.vcf \  
3 > MAPPEDREADS_variants_filtered.vcf.gz  
4  
5 tabix \  
6 -p vcf \  
7 MAPPEDREADS_variants_filtered.vcf.gz
```

Open IGV and load the indexed bam file and the indexed vcf file. Emacs 24.5.1 (Org mode 8.3beta)