

# Reads Alignment and Variant Calling

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# Genome Analysis

Alignment is different from Assembly:

- **Alignment** aims to find the best matches of a particular read to a reference genome
- **Assembly** finds the best overlaps among the reads to determine the most likely genome

# Mapping

Given the level of variability across individuals it is expected that an **high fraction of reads will not map.**

The complete **human pan-genome** will require additional 19-40 Mb of novel sequences. *Nature Biotechnology 28, 57–63 (2010)*

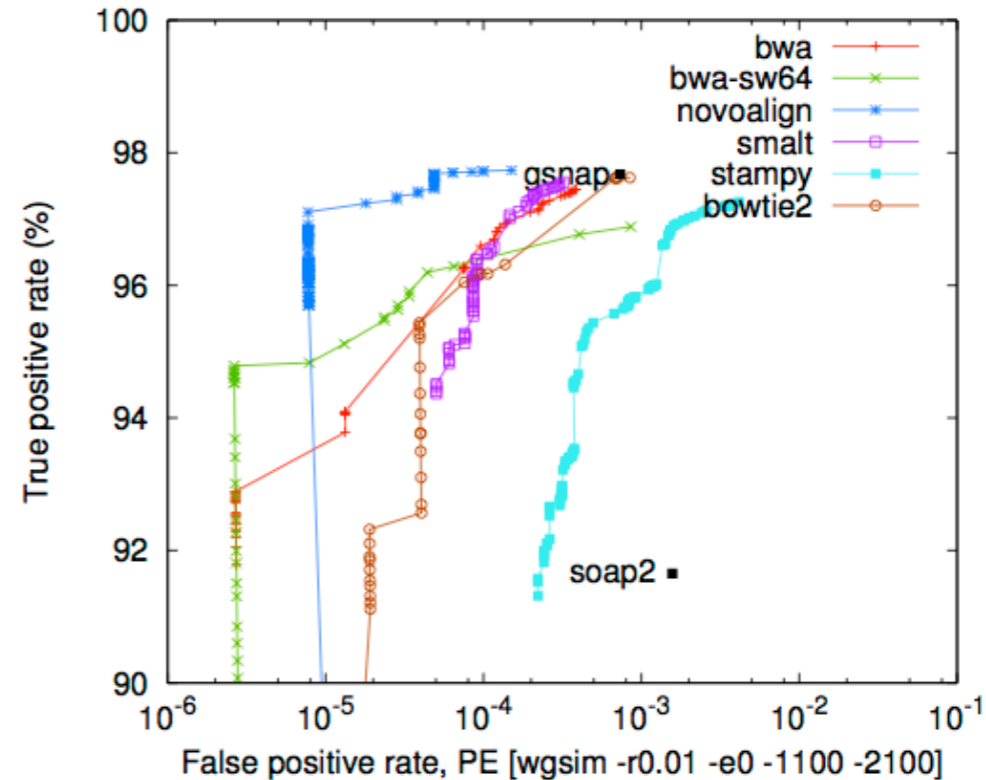
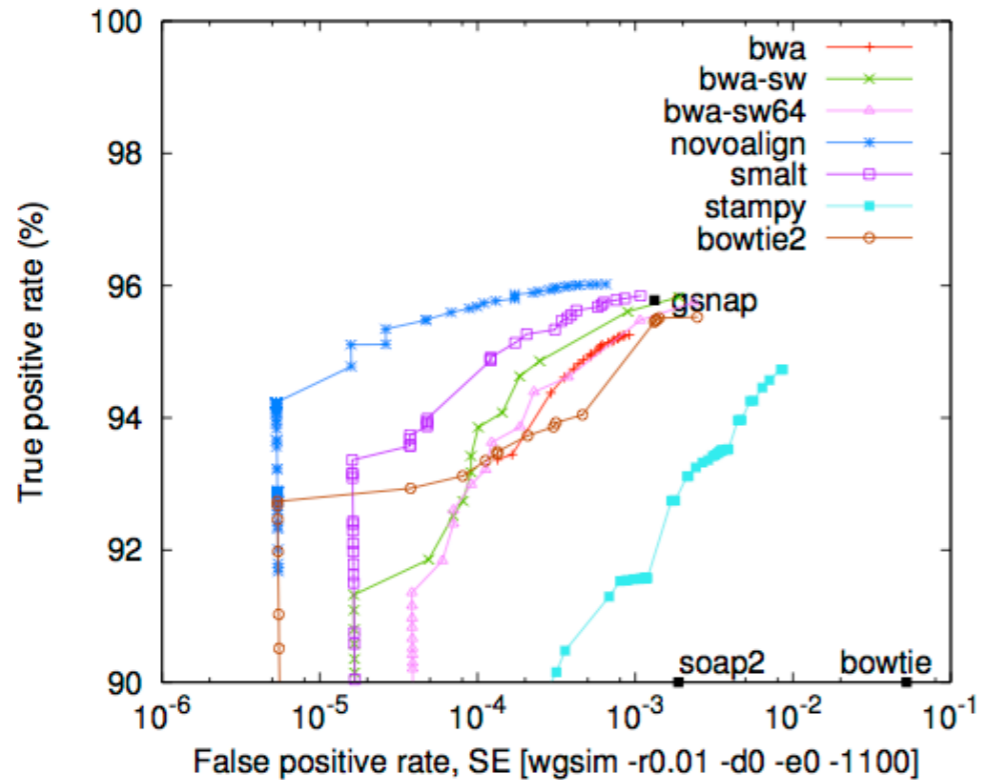
# The Alignment

The main limitation is the computational time.  
the best method should have a good balance between cpu and memory usage, speed and accuracy.

The most popular methods (BWA and Bowtie) are based on suffix array: a sorted array of all suffixes of a string.

BWA and other aligners such as Bowtie use an implementation of the Burrows–Wheeler transform algorithm which is a technique for data compression.

# Testing methods



Program	Version	Options	100k 100bp SE	100k 2x100bp PE (CPU sec)
bowtie2	2.0.0-beta4	-X 650; mapQ>1	78.1	154.0 (to be updated)
bwa	0.5.9-r26-dev	(default); mapQ>0	106.5	230.1
bwa-sw	0.5.9-r26-dev	(default); mapQ>0	237.4	502.0
bwa-sw64	0.6.0-r79-dev	(default); mapQ>0	139.4	286.5
gsnap	2011-10-16	(default); mapQ>3	98.9	538.9
novoalign	2.05.33	-k14 -s3 -i 500 50; mapQ>3	359.7	349.5
smallt	~2011-10-17	-k20 -s13 -i 650; mapQ>0	468.8	640.2

# Genome Indexing

The first step consist in the creation of an **indexed reference genome**.

```
> bwa index $db.fasta
```

two useful option:

-p database prefix

-a indexing algorithm

“bwtsw” for large genome (> 50,000,000 BP)

“is” for smaller genomes

Reference from different institutions

<ftp://gsapubftp-anonymous@ftp.broadinstitute.org/bundle/2.8/hg19/>

# Other Indexing

GATK requires **two specific index files** with extension fai and dict.

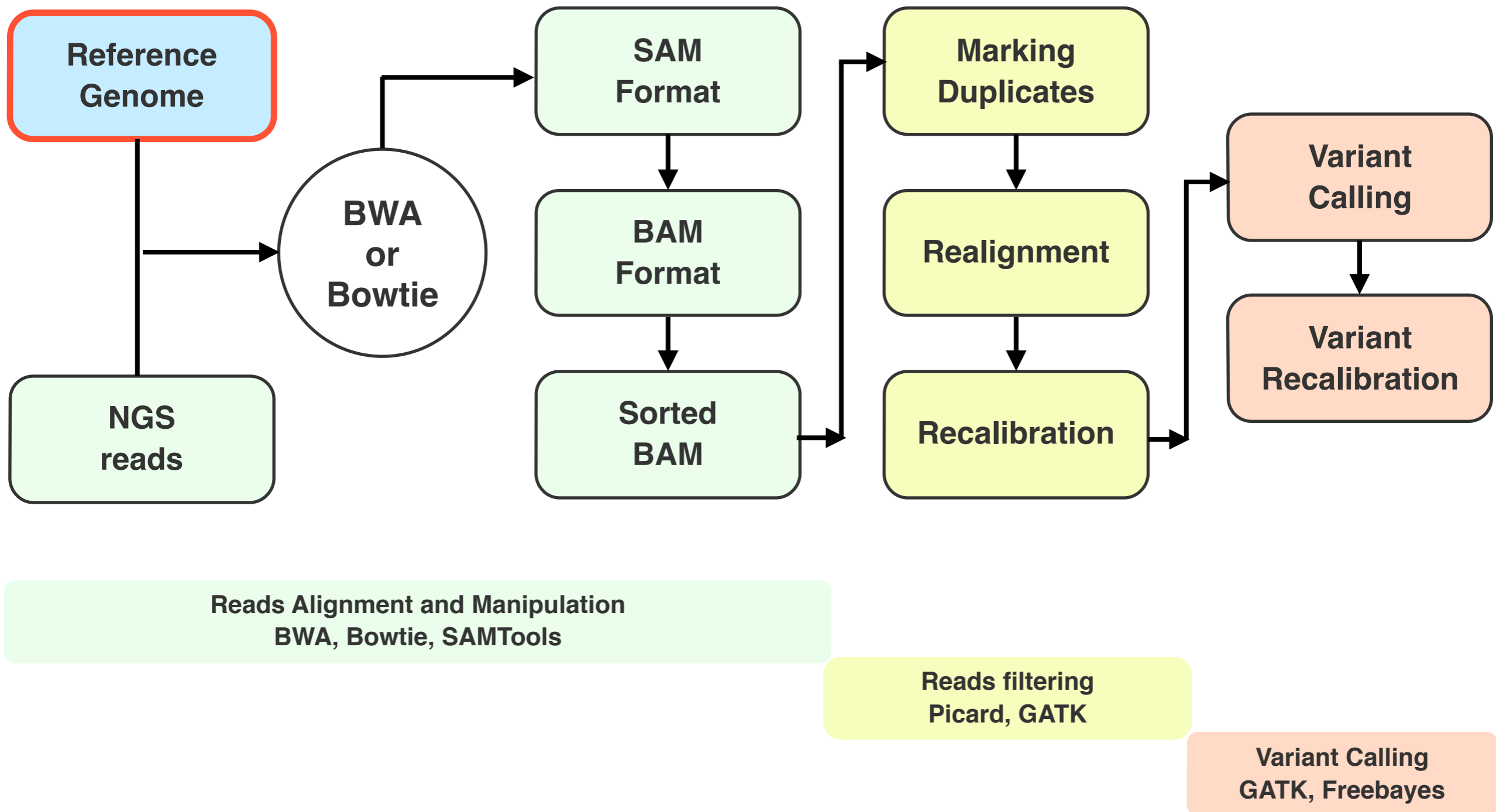
The fai file is generated by *samtools*

```
> samtools faidx $db.fasta
```

The dict file is generated by *picard*

```
>java -jar CreateSequenceDictionary.jar \
REFERENCE=$db.fasta OUTPUT=$db.dict
```

# Variant Calling Steps



# Alignments of the reads

The alignment is generated using **bwa aln**

```
> bwa aln -t [opts] $db $file.fastq
```

two useful options:

- f output file
- t number of threads

If you are analyzing **pair-end sequencing** you need to repeat the alignment for both *fastq* files.

# Generate the SAM file

The SAM file is generated using **bwa samse** (single-end) or **sampe** (pair-end)

```
> bwa samse $db $file.sai $file.fastq  
> bwa sampe $db $file1.sai $file2.sai $file1.fastq $file2.fastq
```

two useful option:

**-f output file**

**-r group\_info**

for example

```
@RG\tID:.. .\tLB:.. .\tSM:.. .\tPL:ILLUMINA
```

# SAM to BAM

To save space and make all the process faster we can convert the **SAM file** to **BAM** using *samtools*

```
> samtools view -bS $file.sam > $file.bam
```

useful option:

- b output BAM
- S input SAM
- T reference genome if header is missing

# Samtools functions

*Samtools* can be used to perform several tasks:

Sorting BAM file

```
> samtools sort $file.bam $file.sorted
```

Create an index

```
> samtools index $file.bam $file.bai
```

Filtering unmapped reads

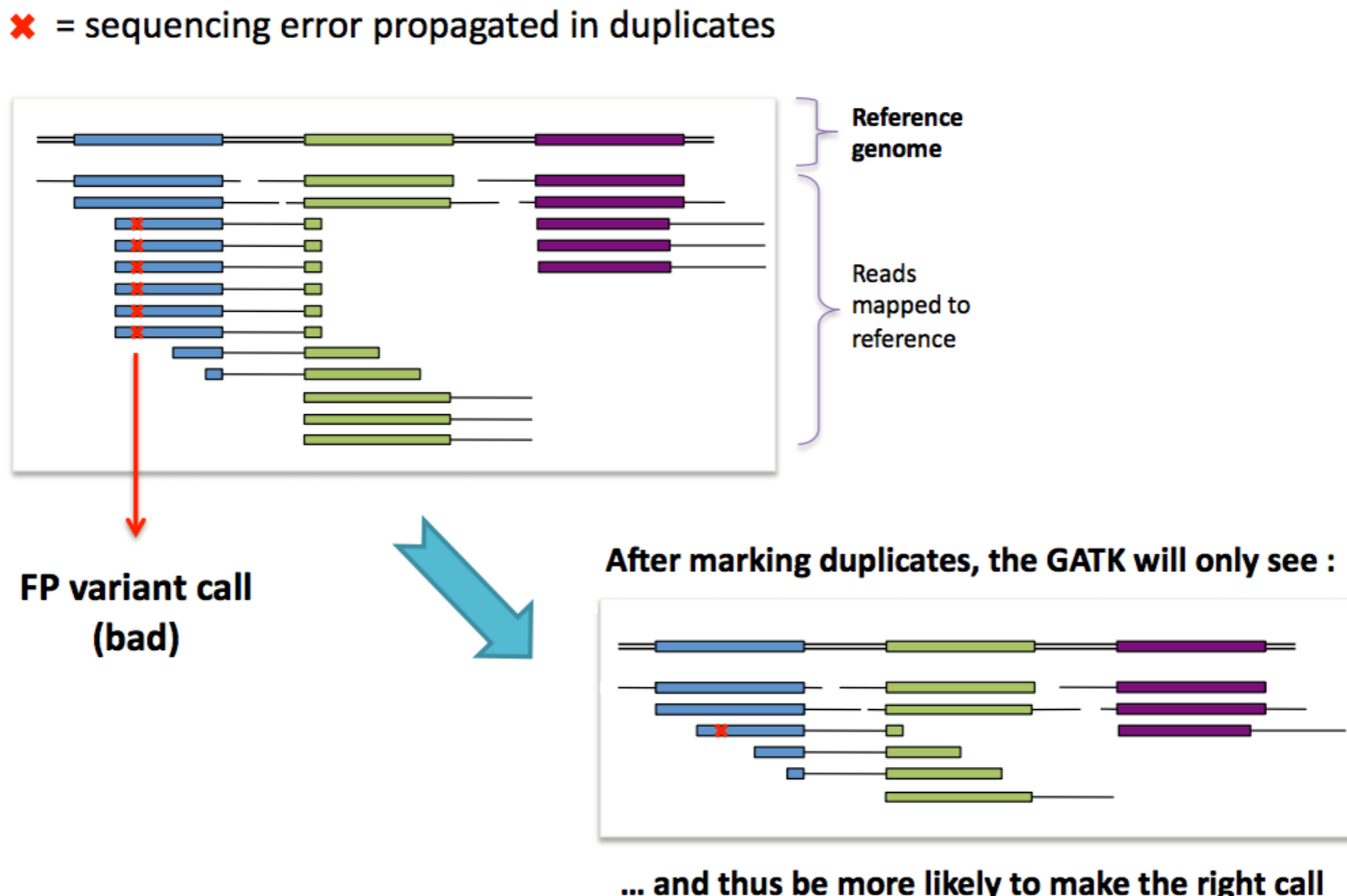
```
> samtools view -h -F 4 $file.bam $file.mapped.bam
```

Select properly paired reads

```
> samtools view -h -f 0X0002 $file.bam $file.paired.bam
```

# Optical Duplication

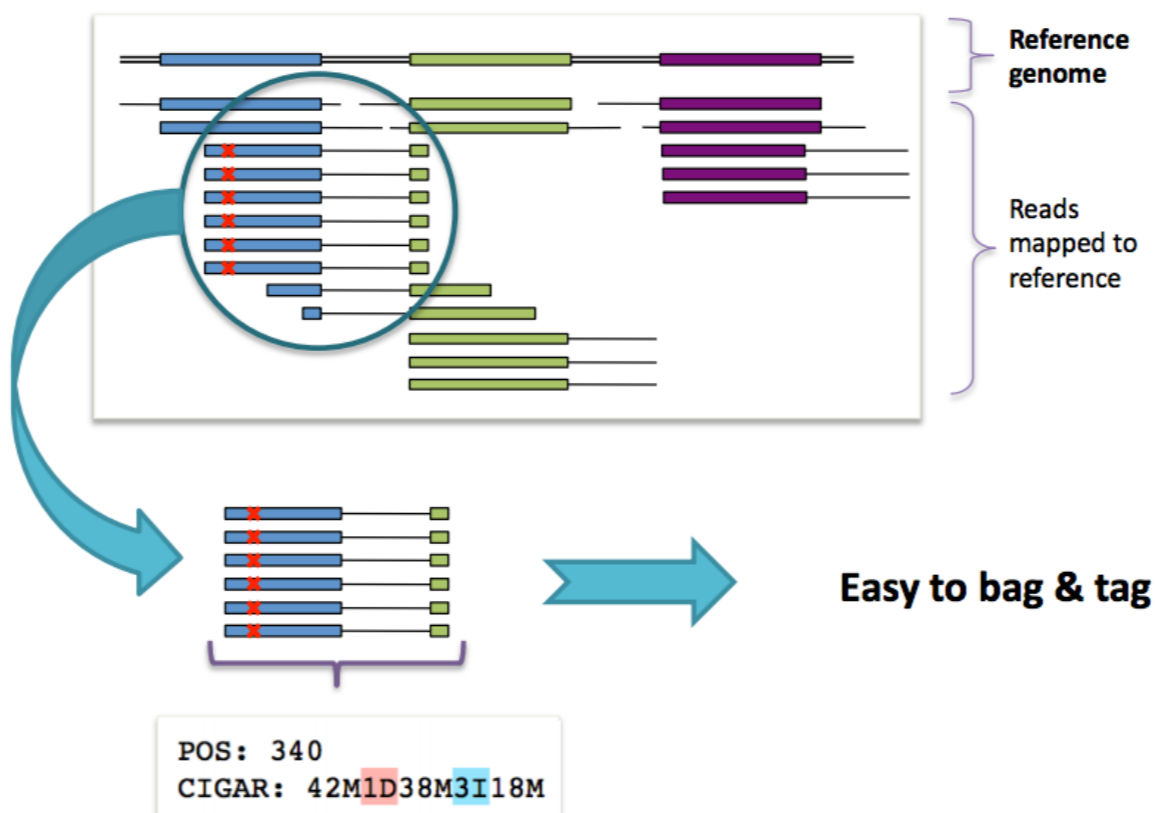
Optical duplicates are due to a read being read twice. The number of the duplicates depends on the depth of the sequencing, the library and sequencing technology.



# Picard

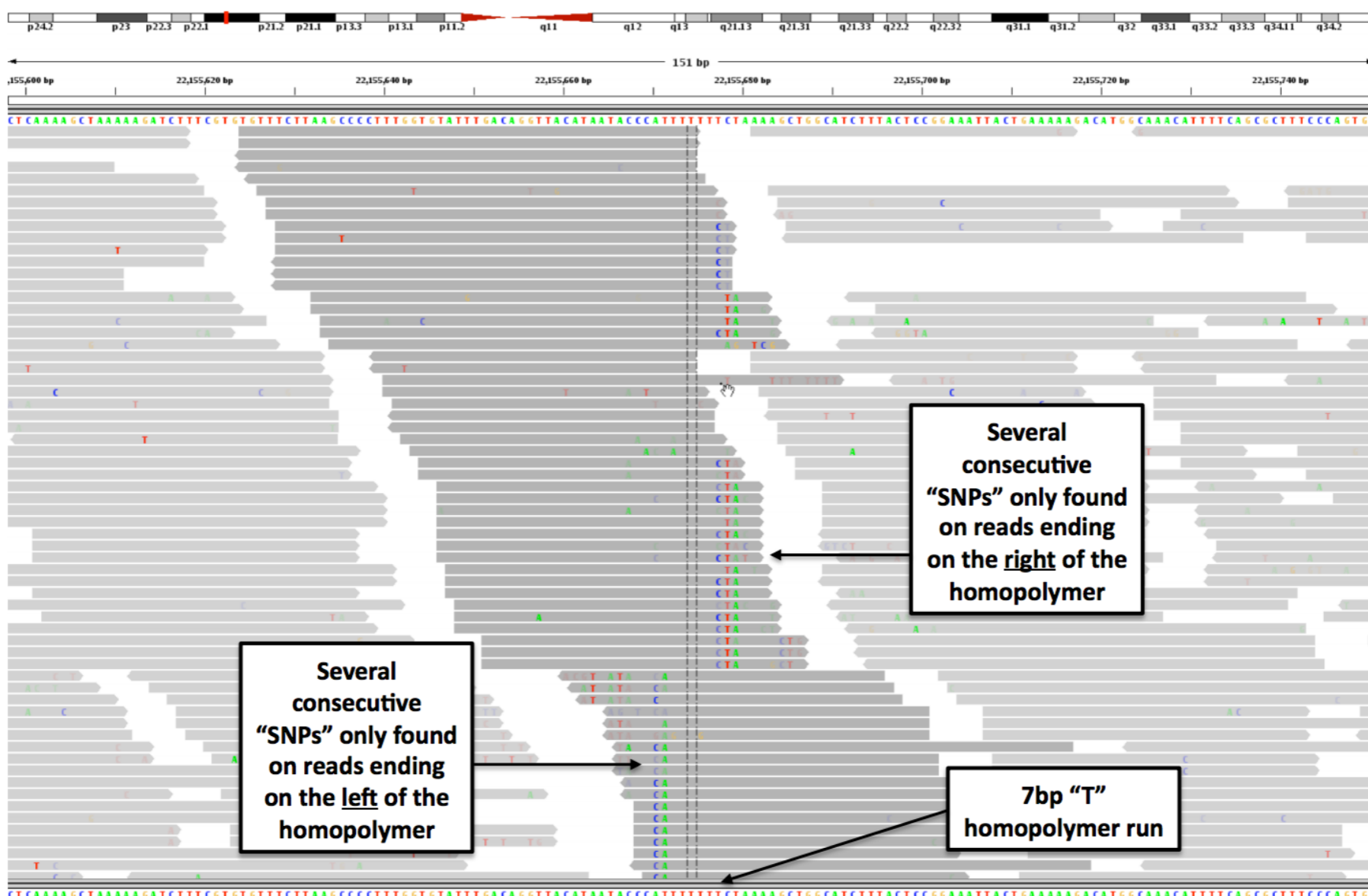
Duplication can be detect comparing the CIGAR.  
Picard is used to mark the duplicated reads.

```
# Mark duplicate  
  
java -Xmx4g -Djava.io.tmpdir=/tmp -jar MarkDuplicates.jar \  
INPUT=$file.bam OUTPUT=$file.marked.bam METRICS_FILE=metrics \  
CREATE_INDEX=true VALIDATION_STRINGENCY=LENIENT
```



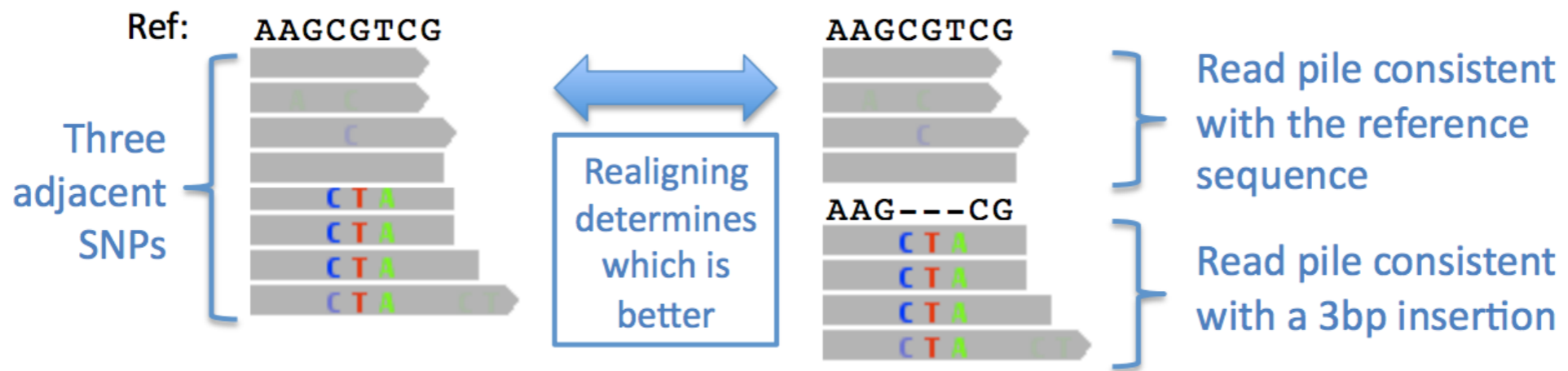
# Indel realignment

The mapping of **indels** especially in regions near to the ends can be seen as **mismatches**. Consecutive variants close to the ends are suspicious.

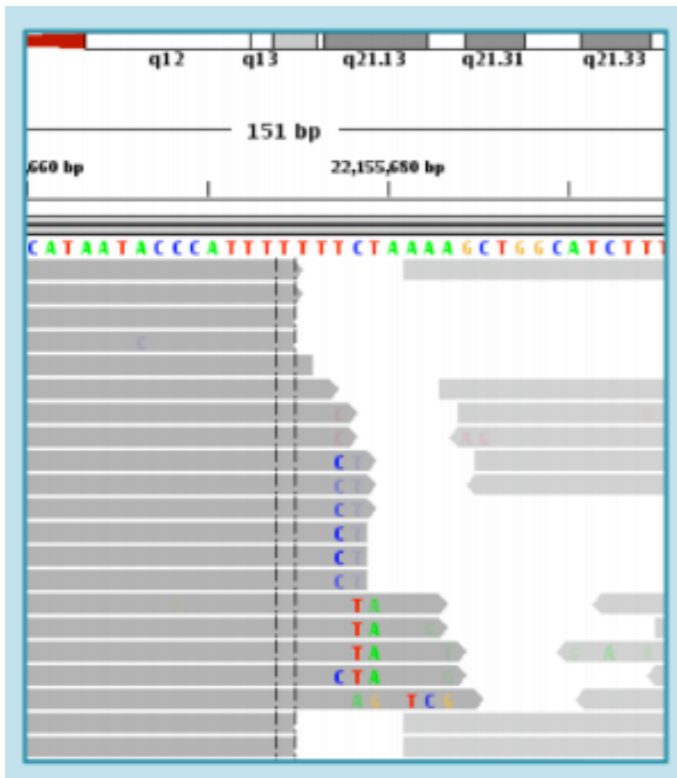


# Better Alignment

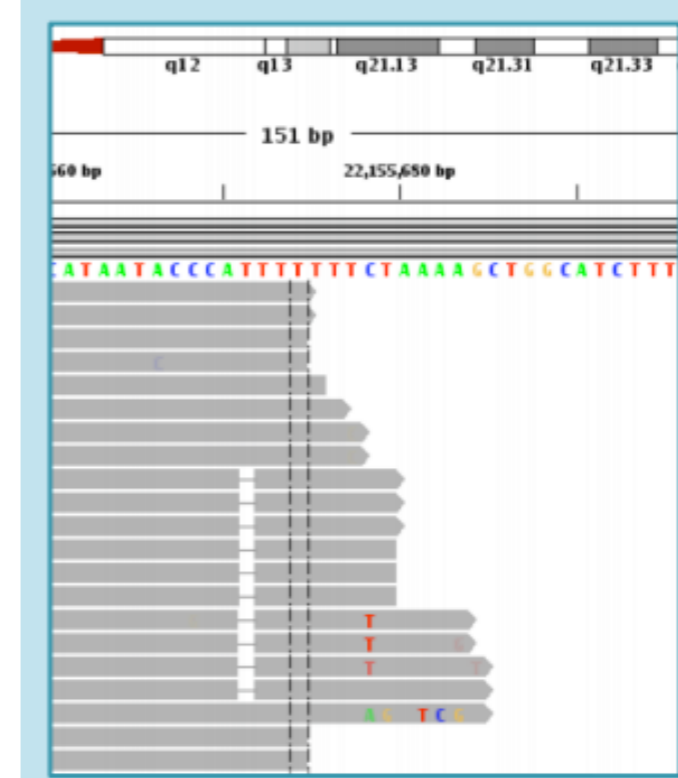
The realignment around the indels improve the quality of the alignment



**RealignerTargetCreator**



**IndelRealigner**



# Realignment

After marking the duplicated reads GATK the alignment is recalculated to improve the match to the indels.

```
# Local realignment

java -Xmx4g -jar GenomeAnalysisTK.jar -T RealignerTargetCreator \
-R $db.fa -o $file.bam.list -I $file.marked.bam

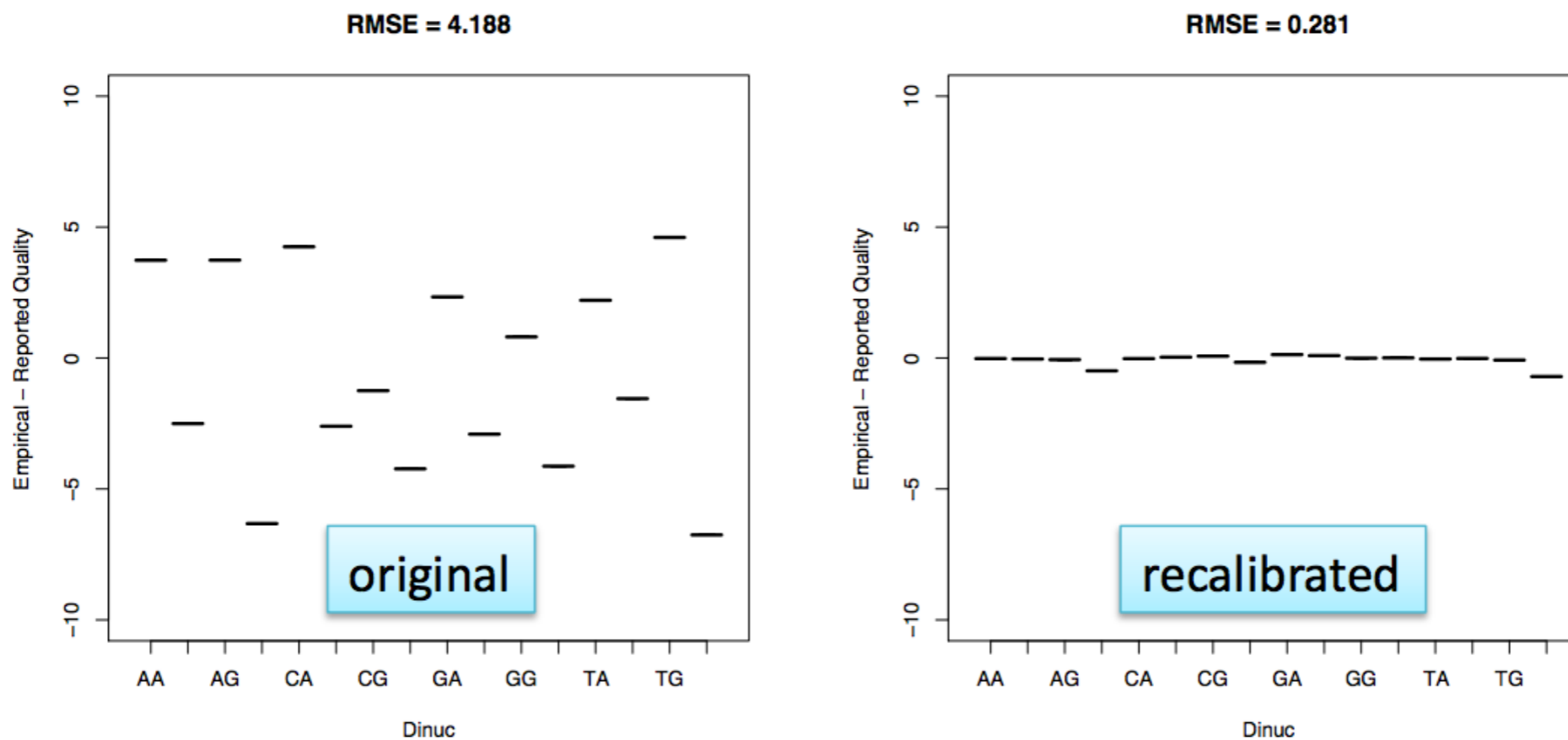
java -Xmx4g -Djava.io.tmpdir=/tmp -jar GenomeAnalysisTK.jar \
-I $file.marked.bam -R $genome -T IndelRealigner \
-targetIntervals $file.bam.list -o $file.marked.realigned.bam

# Picard realignment

java -Djava.io.tmpdir=/tmp/flx-auswerter -jar FixMateInformation.jar \
INPUT=$file.marked.realigned.bam OUTPUT=$file.marked.realigned.fixed.bam \
SO=coordinate VALIDATION_STRINGENCY=LENIENT CREATE_INDEX=true
```

# Quality Score

The quality score are critical for the downstream analysis. This score depends on the nucleotide context.



# The recalibration step

After removing the duplicated reads scores need to be recalibrated using GATK.  
The **recalibration** is detected calculating the **covariation among nucleotide** features.

```
# Recalibration

java -Xmx4g -jar GenomeAnalysisTK.jar -T BaseRecalibrator \
-R $db.fa -I $file.marked.realigned.fixed.bam -knownSites $dbsnp -o \
$output.recal_data.csv

java -jar GenomeAnalysisTK.jar -T PrintReads \
-R $db.fa -I $file.marked.realigned.fixed.bam \
-BQSR $file.recal_data.csv -o $file.marked.realigned.recal.bam
```

# The Variant Calling

There are two possible options: **UnifiedGenotyper** and **HaplotypeCaller**

```
# Variant calling

java -Xmx4g -jar GenomeAnalysisTK.jar -glm BOTH -R $db.fa \
-T UnifiedGenotyper -I $file.marked.realigned.fixed.recal.bam \
-D $dbsnp -o $file.vcf -metrics snps.metrics \
-stand_call_conf 50.0 -stand_emit_conf 10.0 -dcov 1000 -A AlleleBalance
```

or

```
java -Xmx4g -jar GenomeAnalysisTK.jar -T HaplotypeCaller -R $db.fa \
-I $file.marked.realigned.fixed.recal.bam --emitRefConfidence GVCF \
-variant_index_type LINEAR --variant_index_parameter 128000 --dbsnp $dbsnp \
-o $output.recalibrated.vcf
```

**UnifiedGenotyper** is faster and **HaplotypeCaller** is more accurate on the detection of indels.

# Improve Variant Calling

The final step consist in **recalibration and filtering**. The letter are based on previously known mutation events.

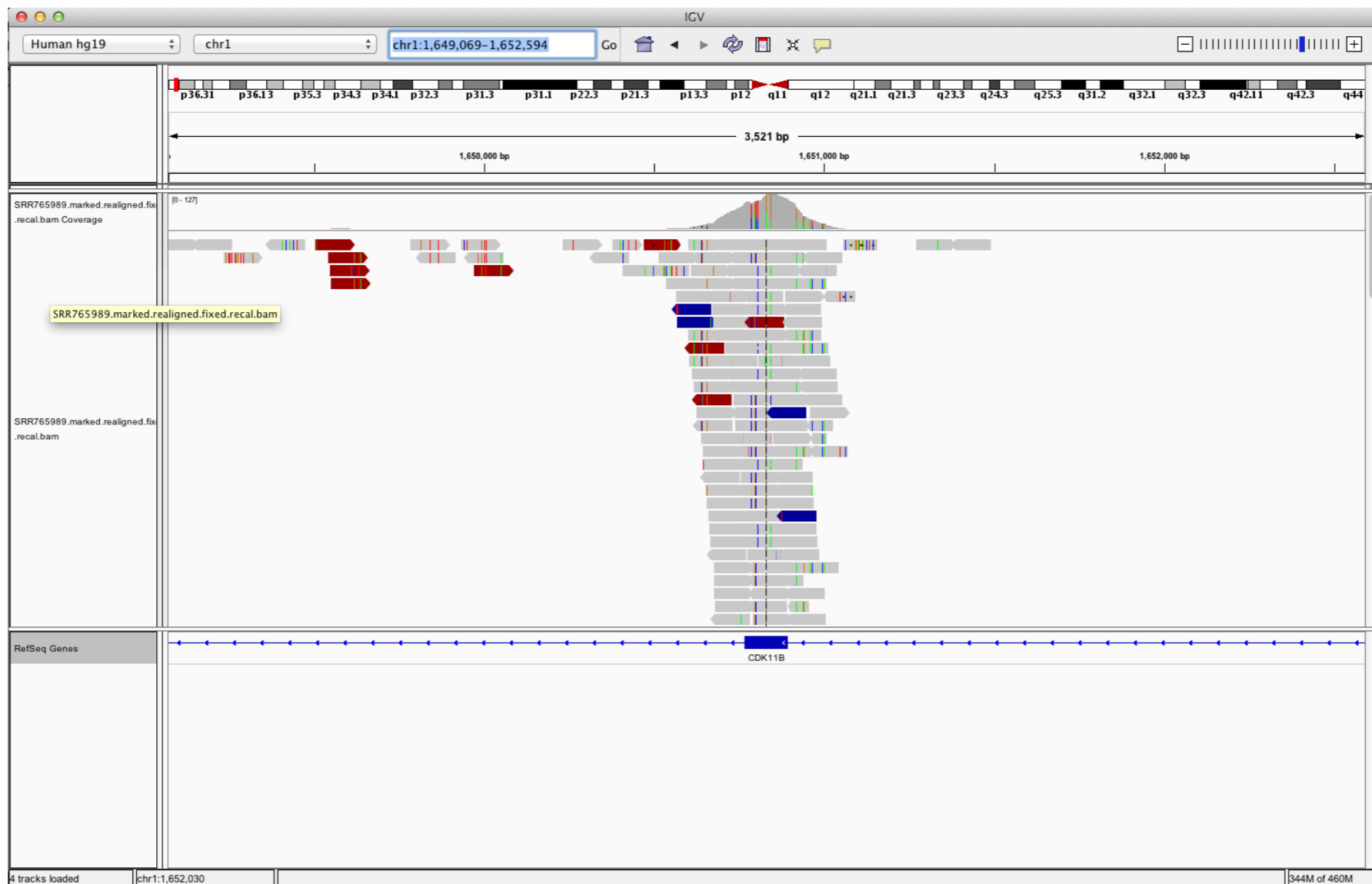
```
# Variant Quality Score Recalibration
java -jar GenomeAnalysisTK.jar -T VariantRecalibrator \
-R $db.fa -input $file.vcf \
-resource: {dbsnp, 1000Genomes, Haplotype } \
-an QD -an MQ -an HaplotypeScore {...} \
-mode SNP -recalFile $file.snps.recal \
-tranchesFile $file.recalibrated.tranches

java -jar GenomeAnalysisTK.jar -T ApplyRecalibration \
-R $db.fa -input $file.vcf -mode SNP \
-recalFile $file.snps.recal -tranchesFile raw.SNPs.tranches \
-o $file.recalibrated.vcf -ts_filter_level 99.0

# Variant Filtering
java -Xmx4g -jar GenomeAnalysisTK.jar -R $db.fa \
-T VariantFiltration -V $file.recalibrated.vcf \
-o $file.recalibrated.filtered.vcf --clusterWindowSize 10 \
--filterExpression "some filter --filterName \"filter_name"
```

# Visualization

Broad Institute have developed the **Integrative Genome Viewer (IGV)** for the **visualization of genomic data** from different sources of data.



[http://cmb.path.uab.edu/training/docs/CB2-201-2015/IGV\\_2.3.40.zip](http://cmb.path.uab.edu/training/docs/CB2-201-2015/IGV_2.3.40.zip)

# For more details

## Samtools

<http://www.htslib.org/doc/>

## GATK Guide

<https://www.broadinstitute.org/gatk/guide/>

## Best practices for variant calling with GATK

<http://www.broadinstitute.org/partnerships/education/broade/best-practices-variant-calling-gatk>

## IGV

<http://www.broadinstitute.org/igv/>