

Summary of the
analysis pipeline
until now

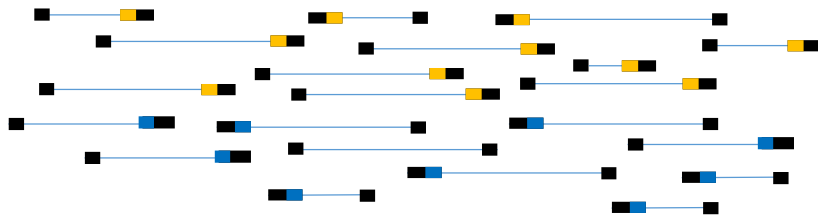
Whole-genome short-read sequencing

DNA

Random shearing



Illumina adapter ligation
incl. individual index



Size selection

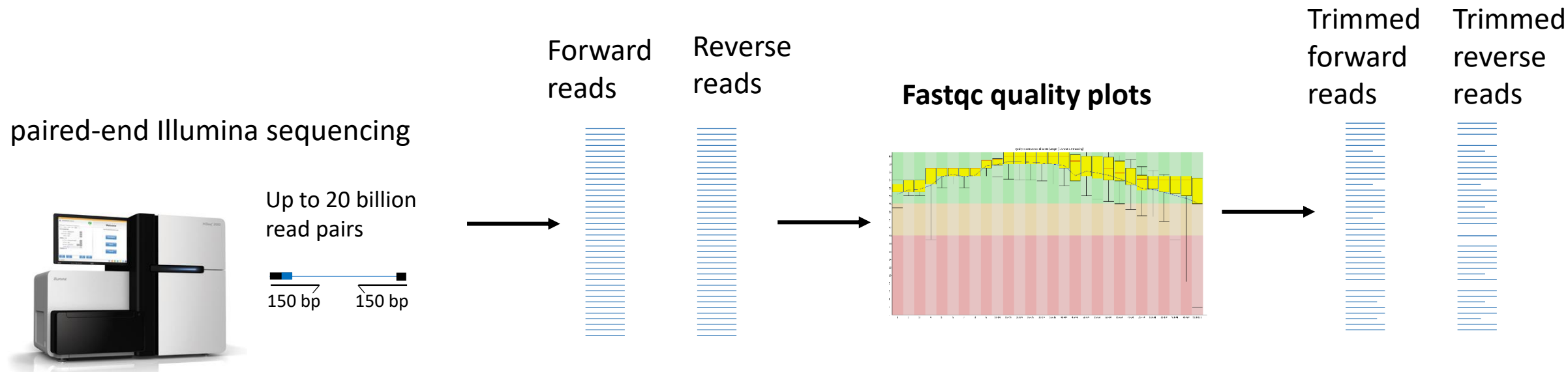


paired-end sequencing

Up to 20 billion read pairs



1. Quality check and trimming raw reads



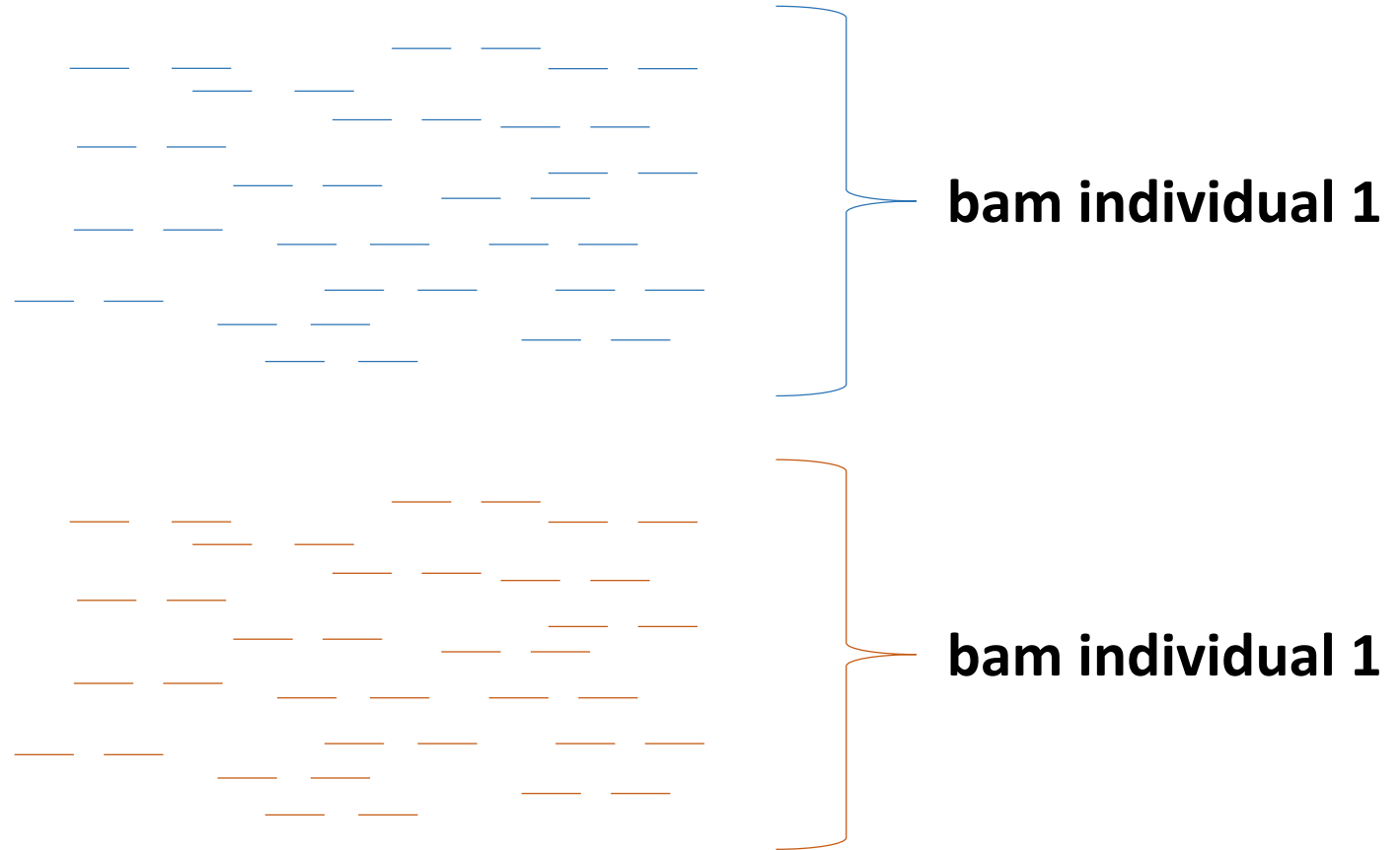
Fastq file of reads

```
@HWUSI-EAS611:34:6669YAAXX:1:1:5069:1159 1:N:0:  
TCGATAATACCGTTTTTTTCCGTTTGATGTTGATACCATT  
+  
IIHHIIHIIIIIIIIIIIIIIIIIIIIIIIIIIIIHIIIIHIIIII
```

2. Alignment to the reference genome with bwa

reference

bwa



3. Variant and genotype calling with bcftools

reference

T

bcftools

T

T

T

T

C

C

C

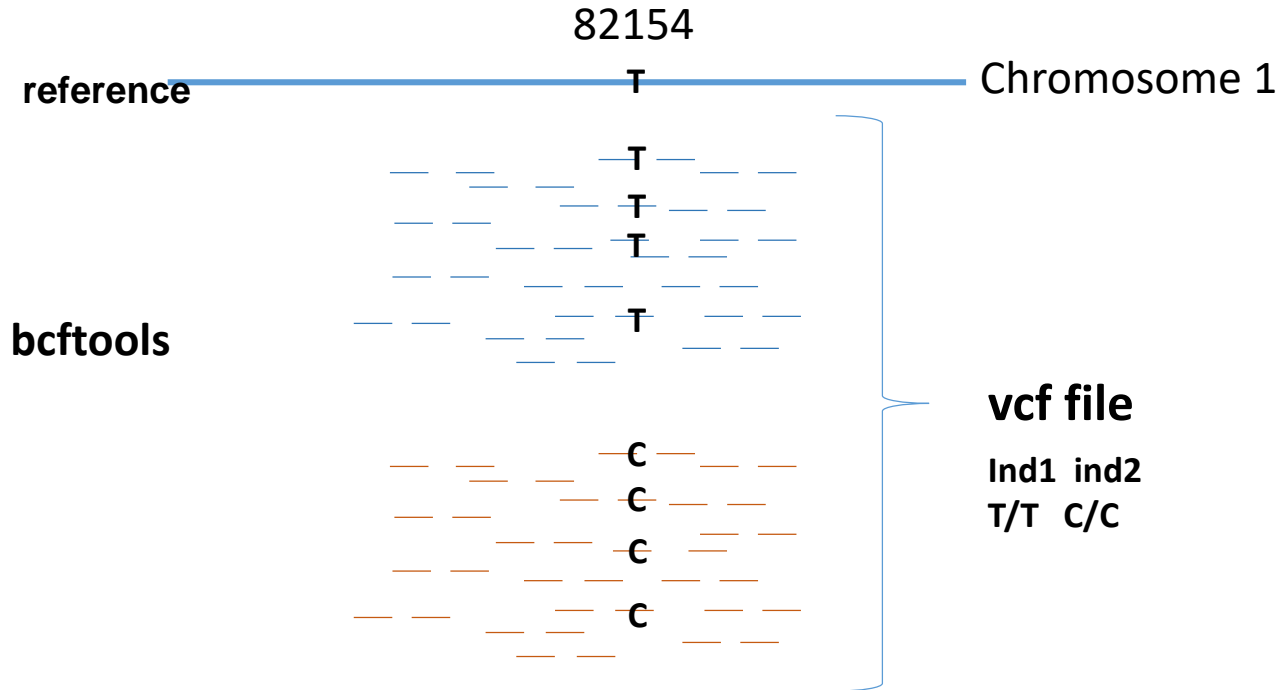
C

vcf file

Ind1 ind2

T/T C/C

3. Variant and genotype calling with bcftools



vcf file: Genotypes for each individual at genomic sites

```
##FORMAT=<ID=GT,Number=1,Type=String,Description="Genotype"  
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT GEN  
chr1 82154 . T C . GT 0/0 1/1 0/0 1/1 0/0 1/1  
chr1 752566 . T . . GT 0/0 0/0 0/0 0/0 0/0 0/0  
chr1 752721 . T C . GT 1/1 1/1 1/1 1/1 1/1 1/1  
chr1 752721 . A . . GT ./.
```