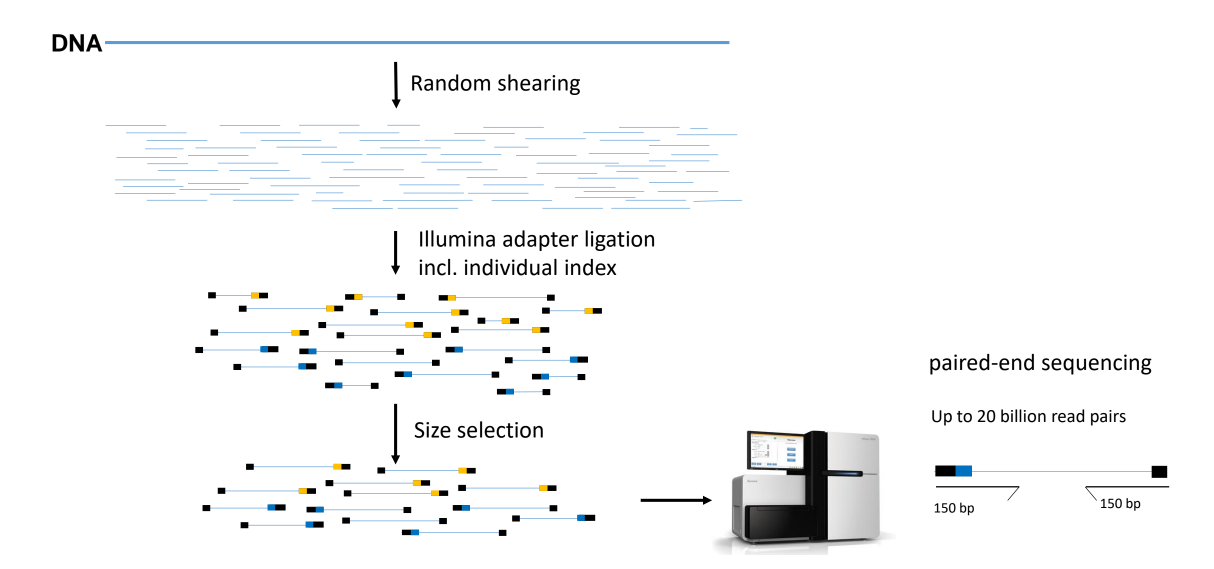
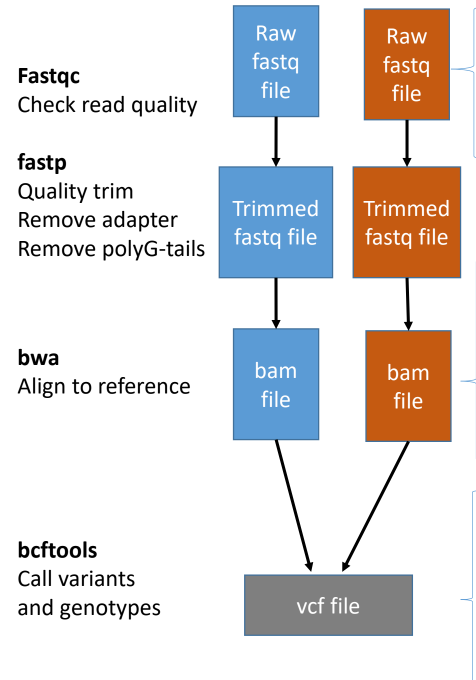
Summary of the analysis pipeline until now

Whole-genome short-read sequencing





Fastq: raw reads with sequencing quality information

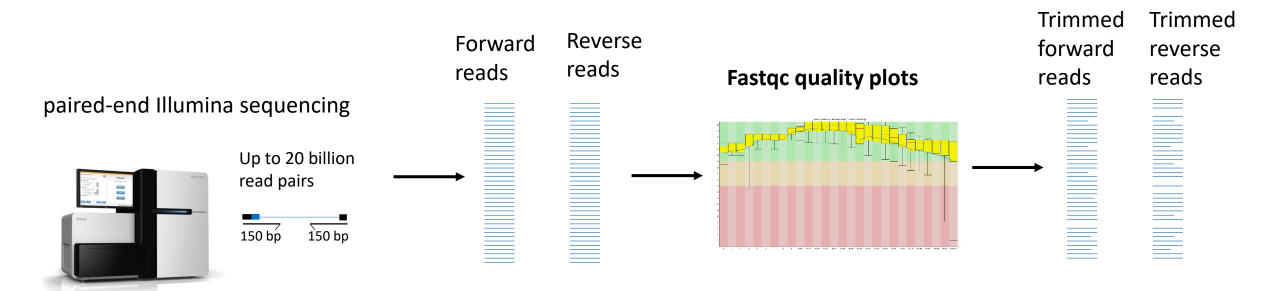
sam file: reads mapped to the reference genome -> binary version: bam file

HWI-ST1145:74:C101DACXX:7:1114:2759:41961 193953 50 100M TGCTGGATCATCTGGTTAGTGGCTTCTGACTCAGAGGACCTTCGTCCCCTGGGGCAGTGGACCTTCCAGTGATTCCCCTGACATAAGGGGCATGGACGA DCDDDDEDDDDDDDDDDDDDCCCDDDCDDDDEEC>DFFFEJJJJJIGJJJJIHGBHHGJIJJJJJGJJJIJJJJJHHJJJJJJHHHHHFFFFFCCC XM:i:3 X0:i:0 XG:i:0 MD:Z:60G16T18T3 NM:i:3 NH:i:1 HWI-ST1145:74:C101DACXX:7:1204:14760:4030 270877 50 100M DDDDDDDDDDDDDDDDDDDDDDDDDEEEEEEEFFFEFFEGHHHHFGDJJHJJJJJJJIIIIIGGFJJJHIIIJJJJJJJJGHHFAHGFHJHFGGHFFFDD@BB AS:i:-11 XM:i:2 X0:i:0 XG:i:0 MD:Z:0A85G13 NM:i:2 NH:i:1 HWI-ST1145:74:C101DACXX:7:1210:11167:8699 271218 50 50M4700N50M GTGGCTCTTCCACAGGAATGTTGAGGATGACATCCATGTCTGGGGTGCACTTGGGTCTCCGAAGCAGAACATCCTCAAATATGACCTCTCG

vcf file: Genotypes for each individual at genomic sites

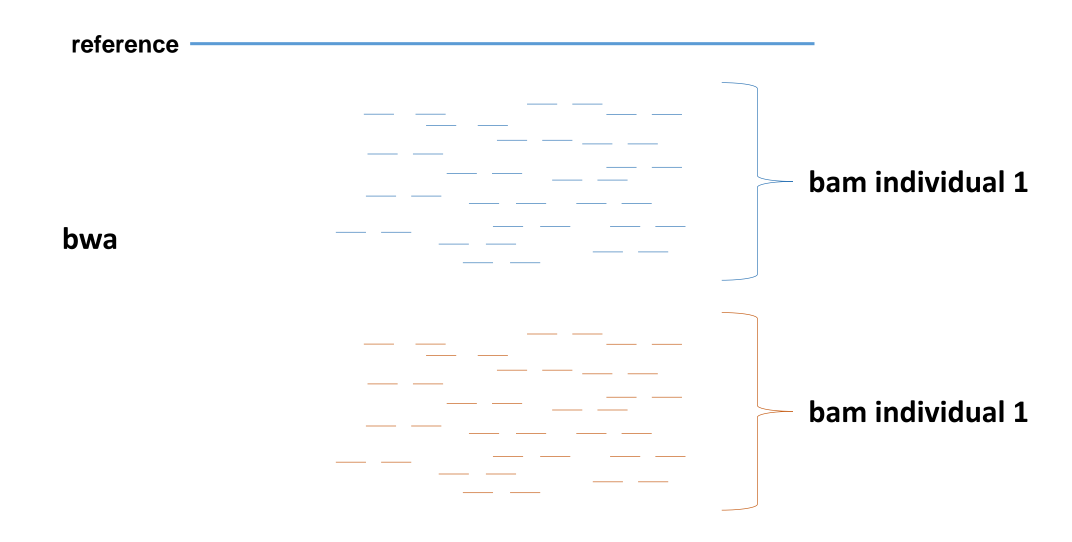
##FORMAT=<ID=GT, Number=1, Type=String, Description="Genotype" POS ID REF ALT QUAL FILTER #CHROM INF₀ FORMAT 82154 chr1 752566 chr1 GT chr1 752721 GT chr1 752721 0/0 0/0 0/0 0/0 0/0

1. Quality check and trimming raw reads

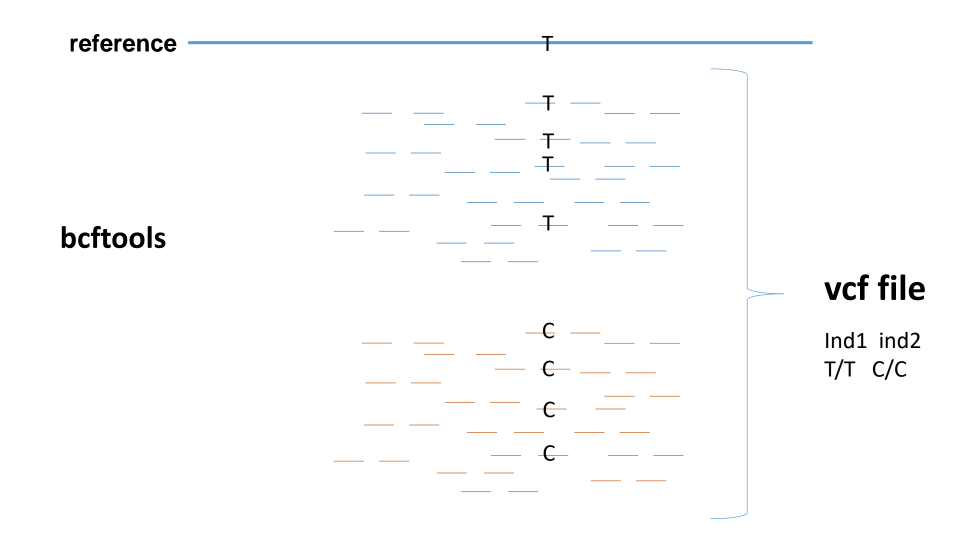


Fastq file of reads

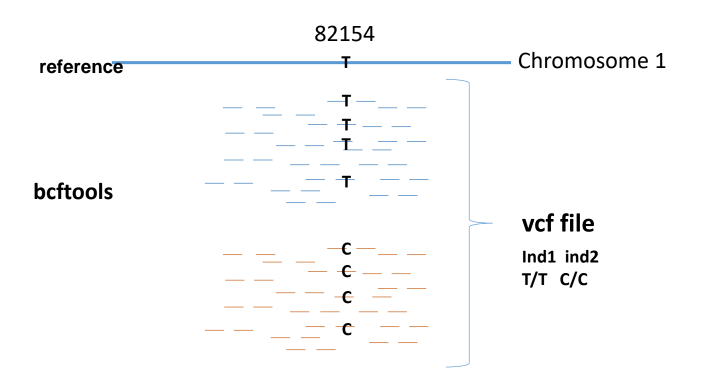
2. Alignment to the reference genome with bwa



3. Variant and genotype calling with bcftools



3. Variant and genotype calling with bcftools



vcf file: Genotypes for each individual at genomic sites

##FORMAT= <id=gt, <="" description="Genotype" number="1," th="" type="String,"></id=gt,>												
#CHROM	POS ID	REF	ALT	QUAL		FILTE	R I	NF0	F	ORMA	ΓG	EN
chr1	82154	•	Т	C	•	GT	0/0	1/1	0/0	1/1	0/0	1/1
chr1	752566	•	Т	•		GT	0/0	0/0	0/0	0/0	0/0	0/0
chr1	752721	•	Т	C		GT	1/1	1/1	1/1	1/1	1/1	1/1
chr1	752721	•	Α			GT	./.	0/0	0/0	0/0	0/0	0/0