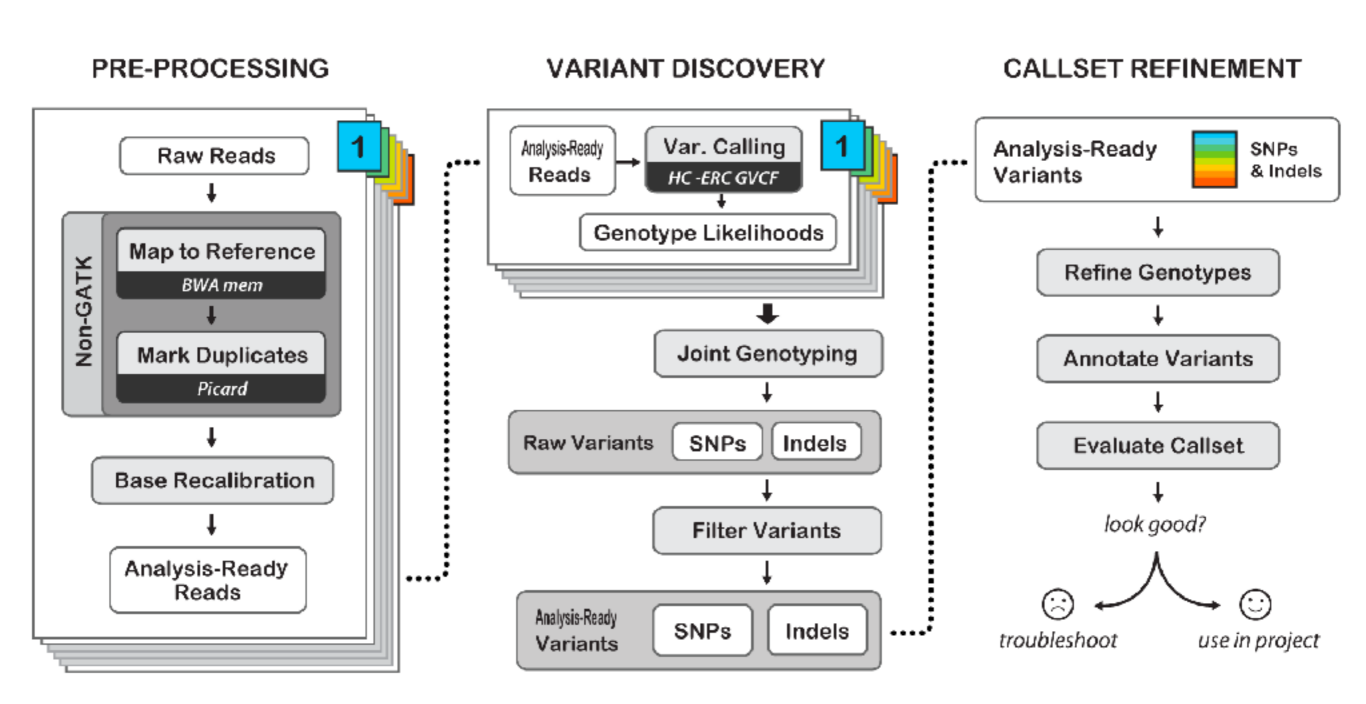
Topic 7: Variant calling using GATK

Learning Goals

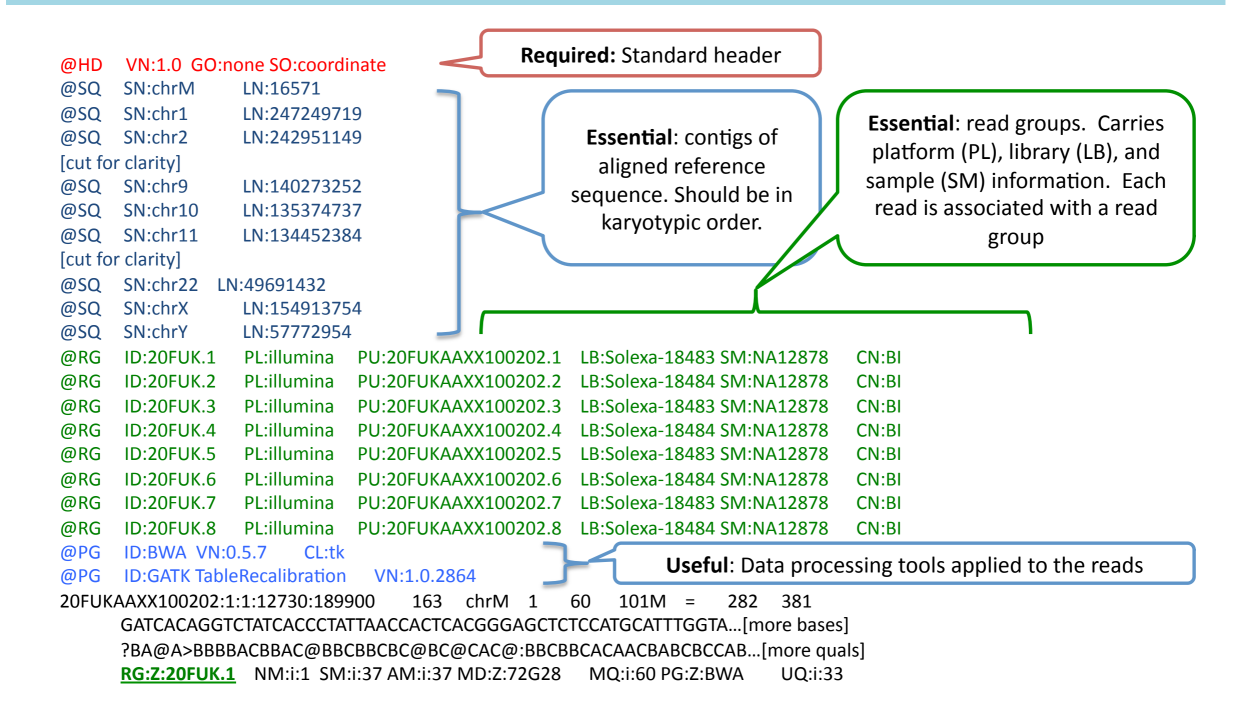
- Define the steps involved in SNP calling and what they are doing.
- Understand the reason for haplotype based SNP calling.
- Understand the recalibration approach to variant filtering.
- Define the N+1 problem in genotyping.

GATK Best Practises for Variant Discovery in DNAseq



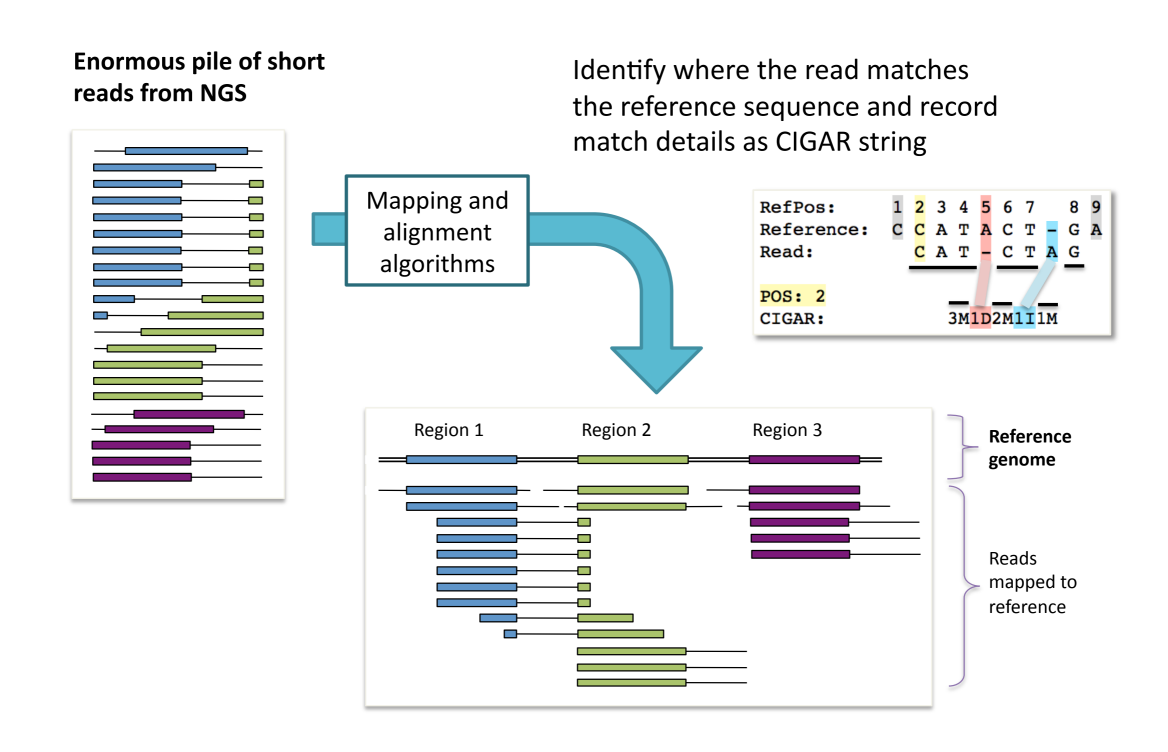
Best Practices for Germline SNPs and Indels in Whole Genomes and Exomes - June 2016

BAM headers: an essential part of a BAM file



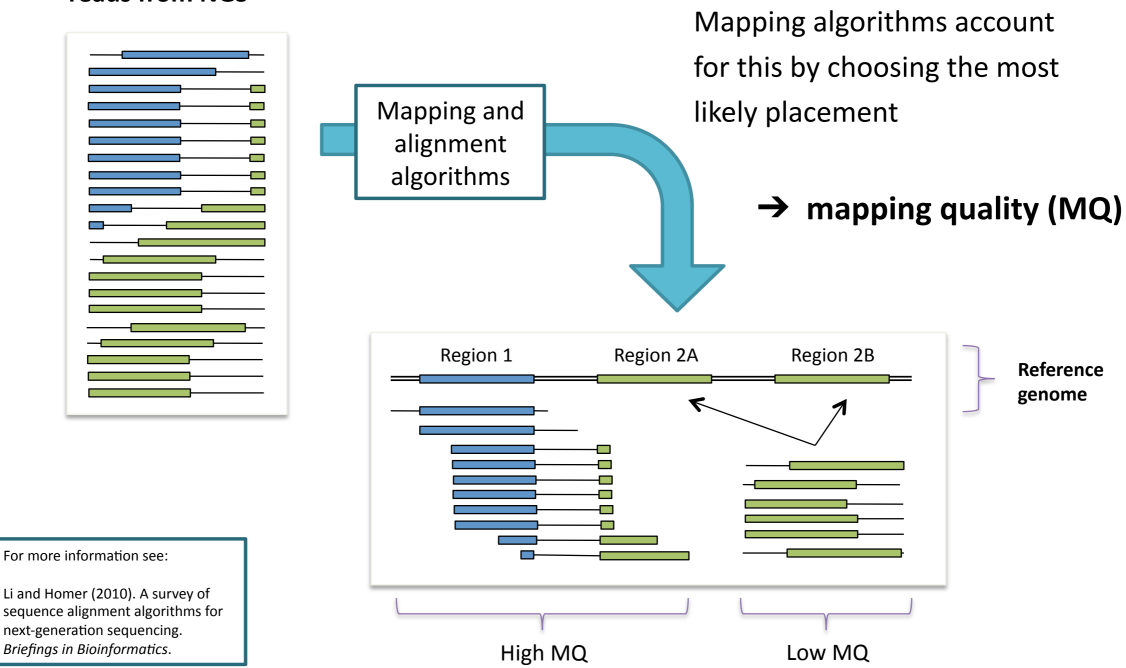
Official specification in http://samtools.sourceforge.net/SAM1.pdf

Mapping short reads to a reference is simple in principle



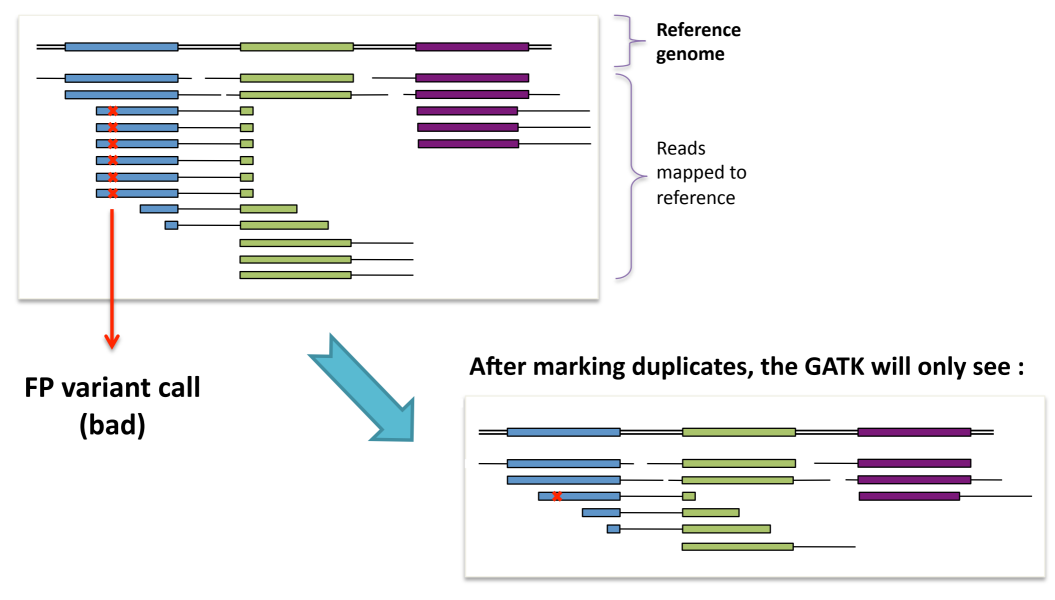
But mapping is actually very hard because of mismatches (true mutations or sequencing errors), duplicated regions etc.

Enormous pile of short reads from NGS



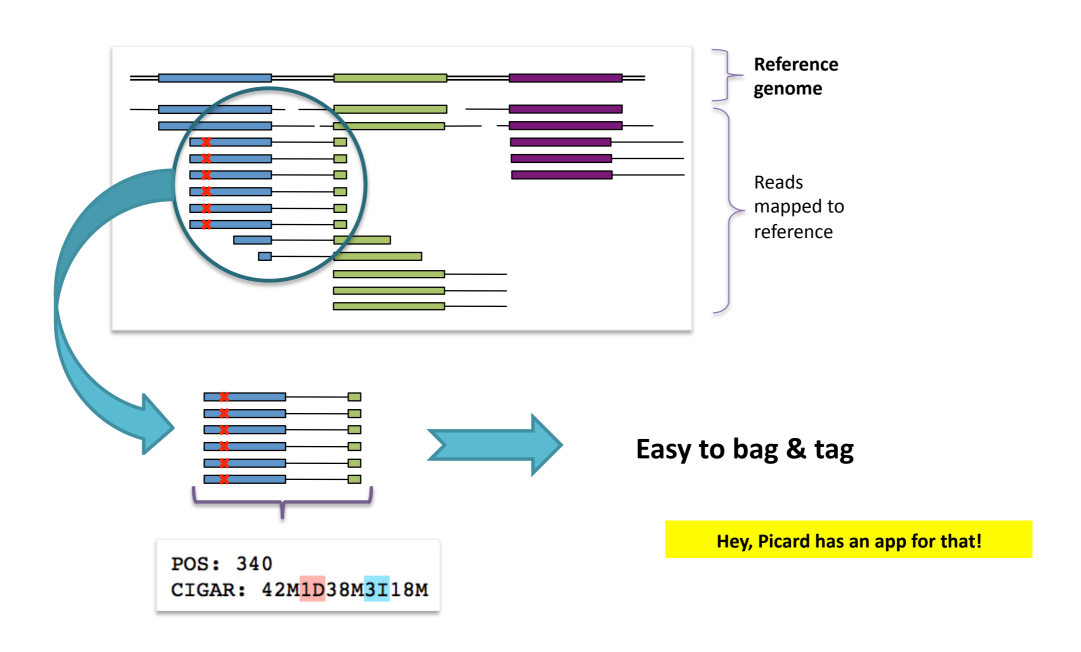
The reason why duplicates are bad

× = sequencing error propagated in duplicates



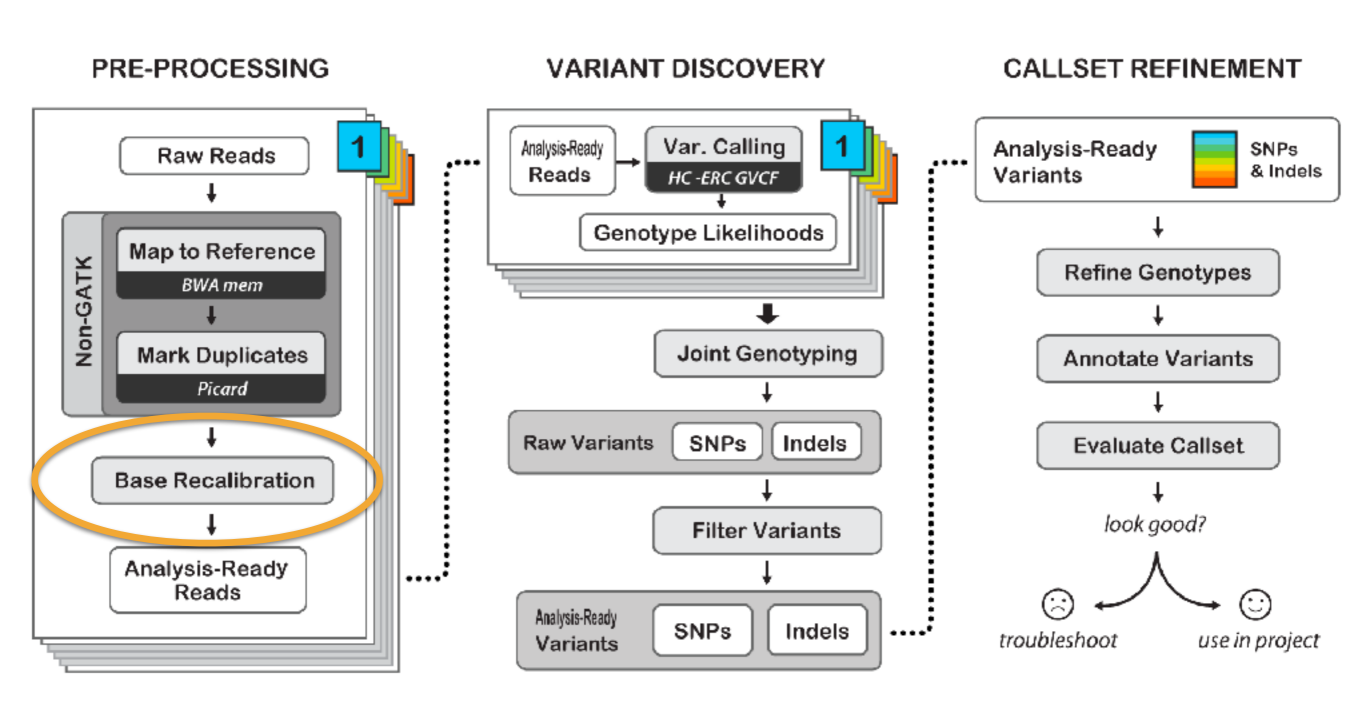
... and thus be more likely to make the right call

Easy to identify: duplicate reads have the same starting position and same CIGAR string



Why wouldn't we do this for GBS?

Base recalibration

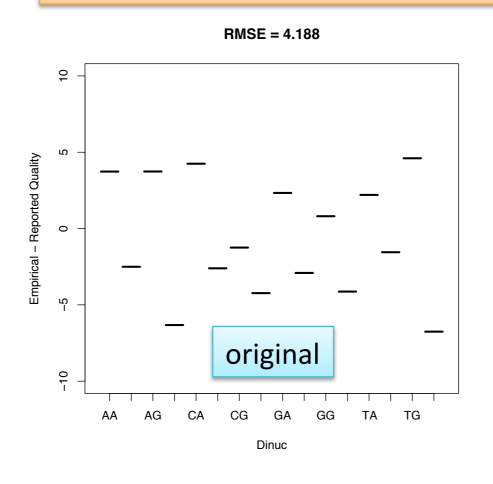


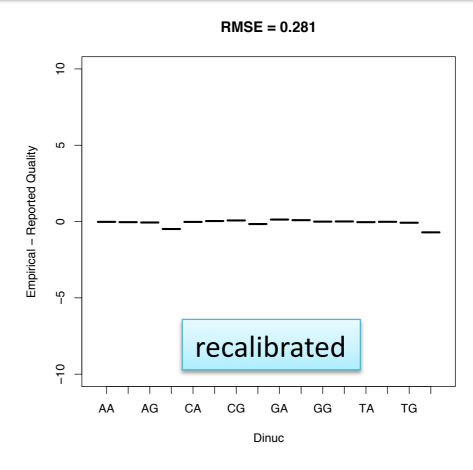
Best Practices for Germline SNPs and Indels in Whole Genomes and Exomes - June 2016

Quality scores issued by sequencers are **inaccurate** and **biased**

- Quality scores are critical for all downstream analysis
- Systematic biases are a major contributor to bad calls

Example of bias: qualities reported depending on nucleotide context





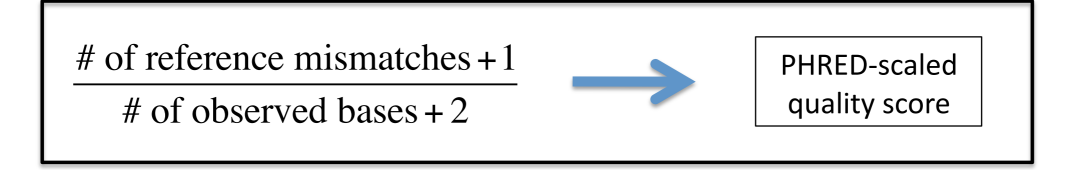
BQSR identifies patterns in how errors correlate with base features

- Empowered by looking at entire lane of data
- Analyze covariation among several features of a base, e.g.:
 - Reported quality score
 - Position within the read (machine cycle)
 - Preceding and current nucleotide (sequencing chemistry effect)
- Based on the patterns identified:
 - Apply corrections to recalibrate the quality scores of all reads in the BAM file.

How covariates are analyzed to identify patterns

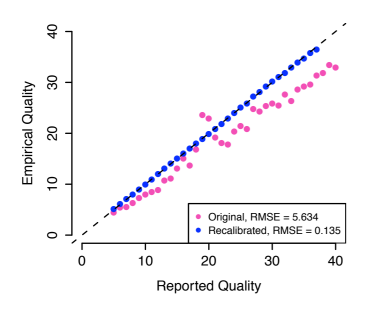
- Any sequence mismatch = error except known variants!
- Keep track of number of observations and number of errors as a function of various error covariates

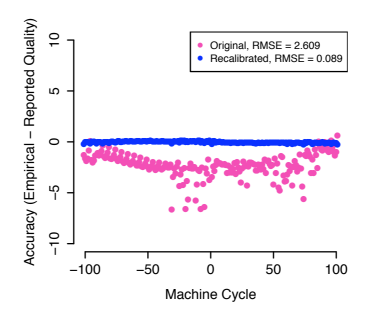
(lane, original quality score, machine cycle, and sequencing context)

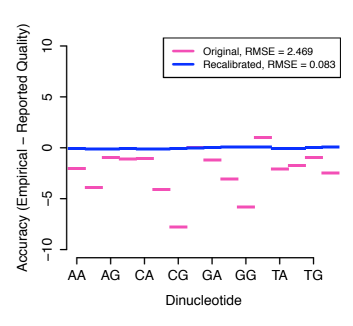


Did the recalibration work properly?

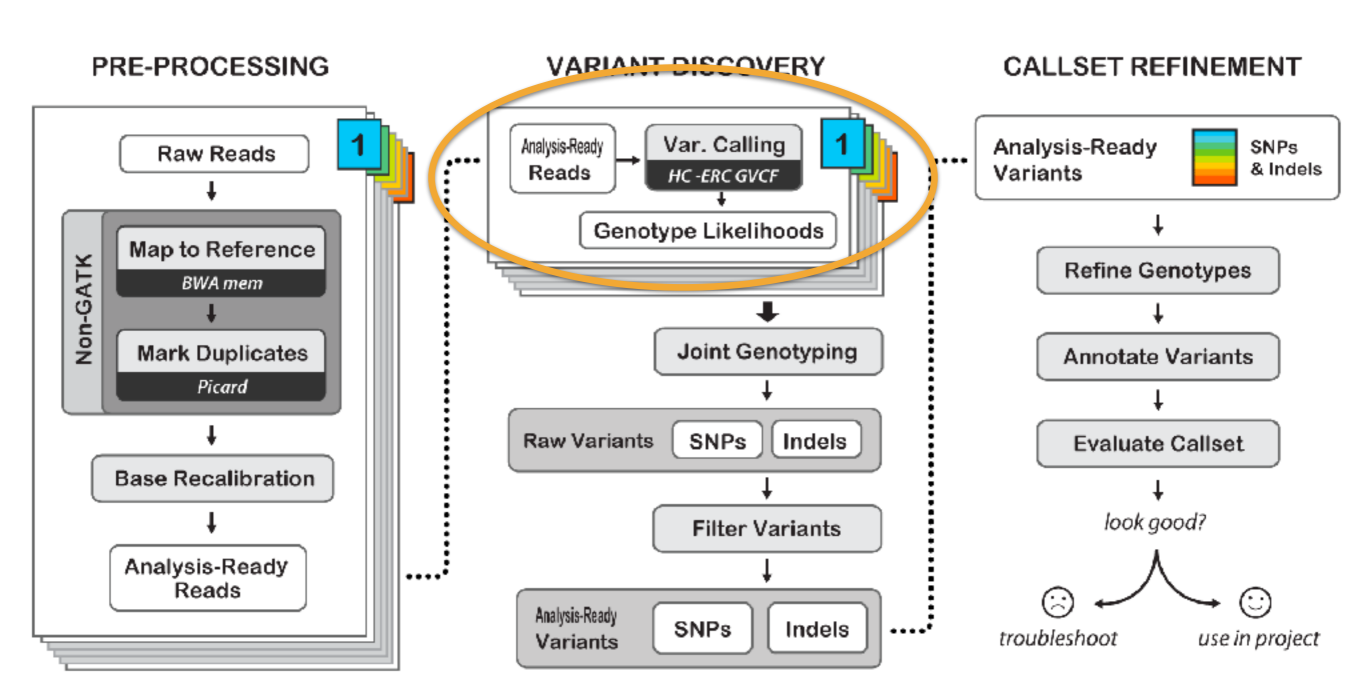
Post-recalibration quality scores should fit the empirically-derived quality scores very well; no obvious systematic biases should remain





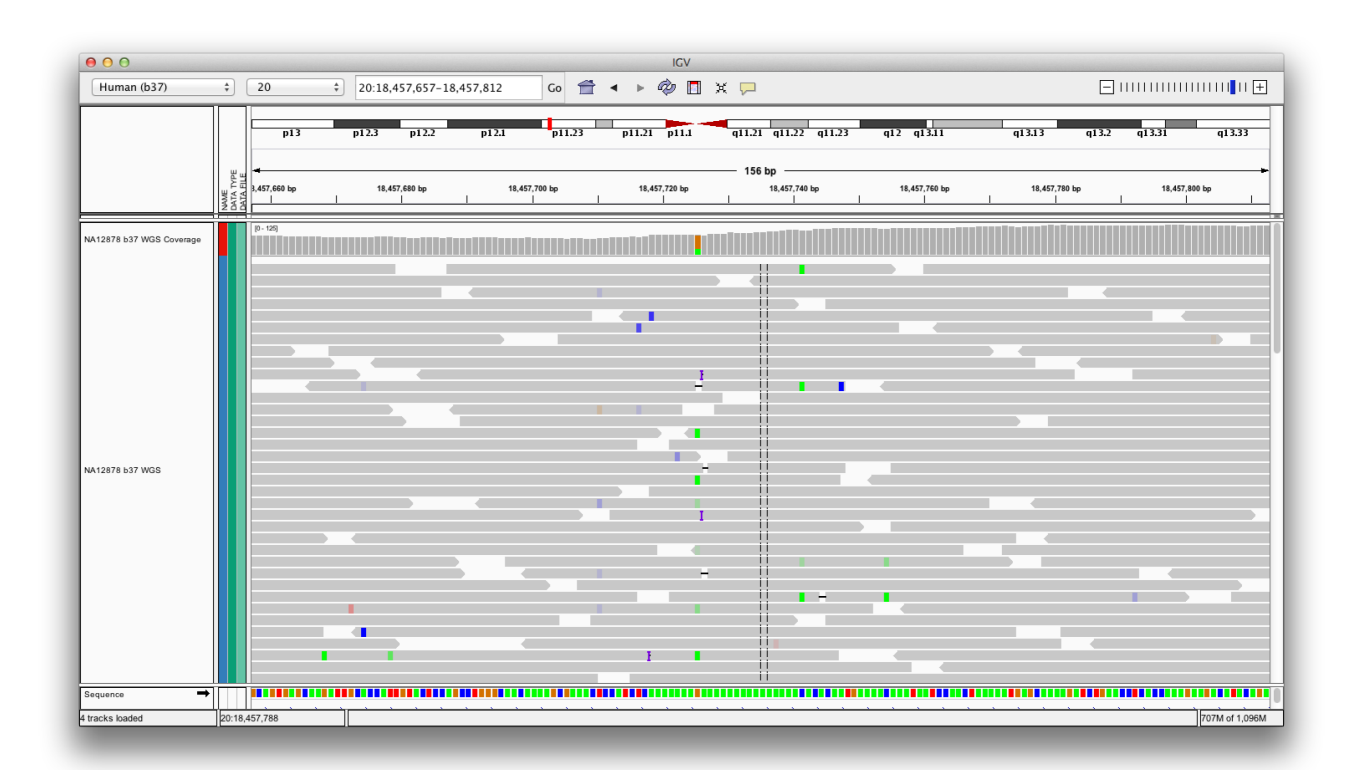


Base recalibration



Best Practices for Germline SNPs and Indels in Whole Genomes and Exomes - June 2016

Real mutations are hidden in the noise



Summed up in GATK terms

Prior of the Likelihood of the genotype
$$Pr\{G|D\} = \frac{\Pr\{G\}\Pr\{D|G\}}{\sum_{i}\Pr\{G_{i}\}\Pr\{D|G_{i}\}}, \ [\text{Bayes' rule}] \qquad \text{Diploid assumption}$$

$$\Pr\{D|G\} = \prod_{j} \left(\frac{\Pr\{D_{j}|H_{1}\}}{2} + \frac{\Pr\{D_{j}|H_{2}\}}{2}\right) \text{ where } G = H_{1}H_{2}$$

$$\Pr\{D|H\} \text{ is the haploid likelihood function}$$

Variant callers in GATK

UnifiedGenotyper

Call SNPs and indels separately by considering each variant locus independently

- Accepts any ploidy
- Pooled calling

HaplotypeCaller

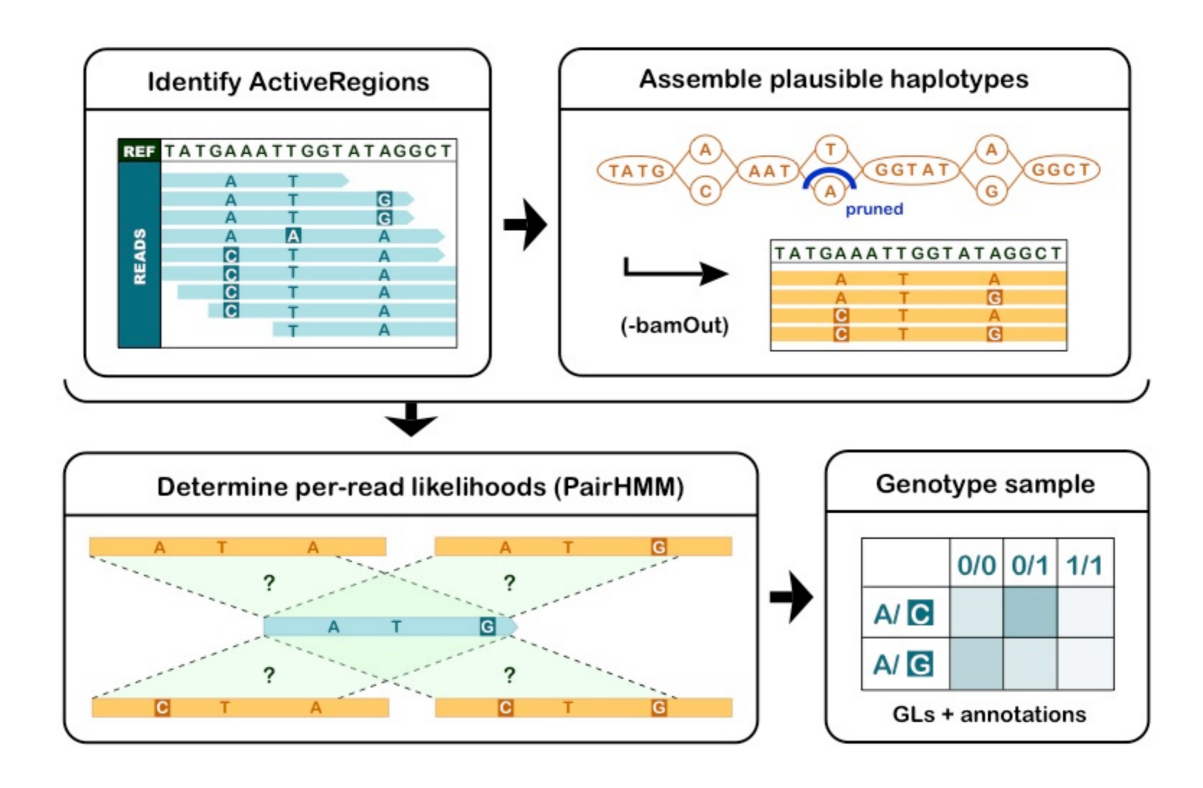
Call SNPs, indels, and some SVs simultaneously by doing local re-assembly and considering haplotypes

- More accurate (esp. indels)
- Reference confidence model
- Replaces UG

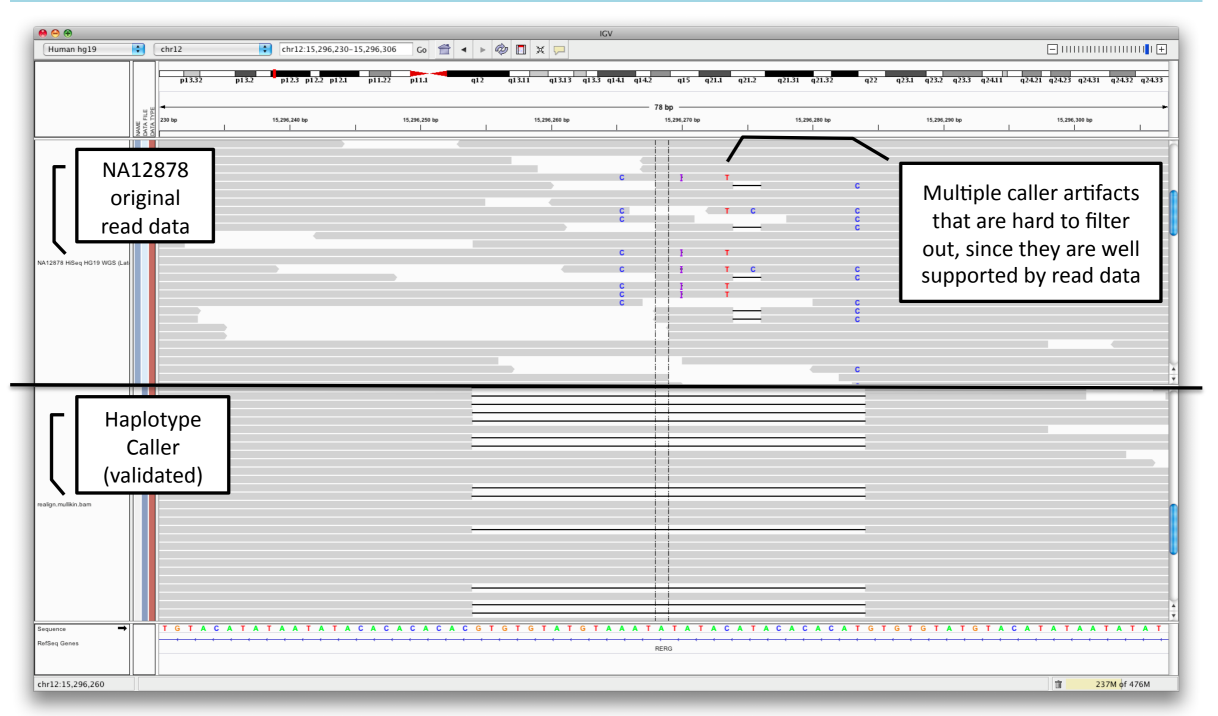
HaplotypeCaller method overview

- Call SNPs, indels, and some SVs simultaneously by doing local re-assembly and considering haplotypes
 - Determine if a region has potential variation
 - Make deBruijn assembly graph of the region
 - Paths in the graph = potential haplotypes to evaluate
 - Calculate data likelihoods given the haplotypes (PairHMM)
 - Identify variants on most likely haplotypes
 - Compute allele frequency distribution to determine most likely allele count, and emit a variant call if appropriate
 - If emitting a variant, assign genotype to each sample

HC method illustrated

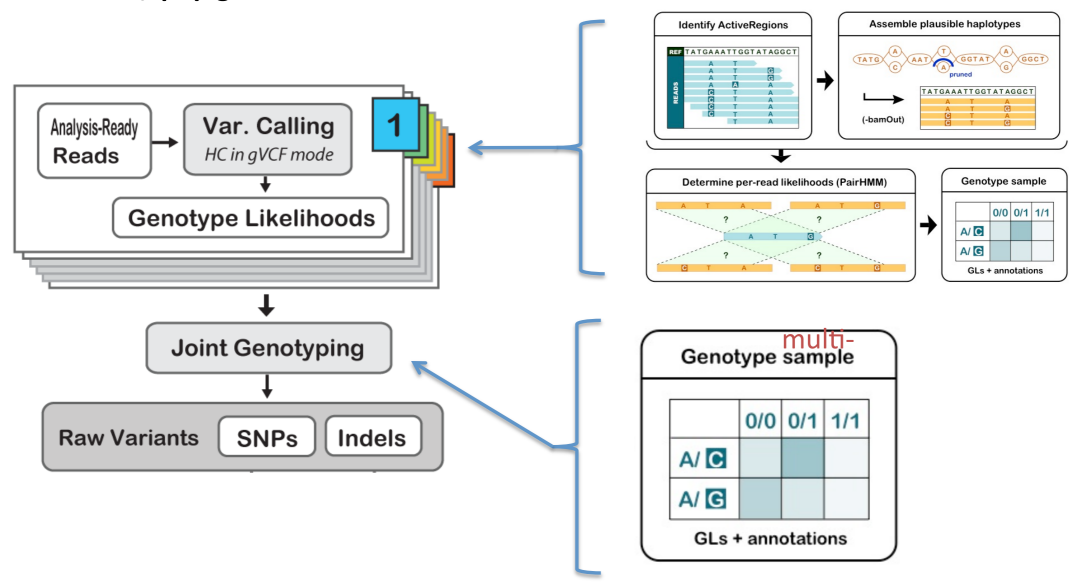


Artifactual SNPs and small indels caused by large indel can be recovered by assembly

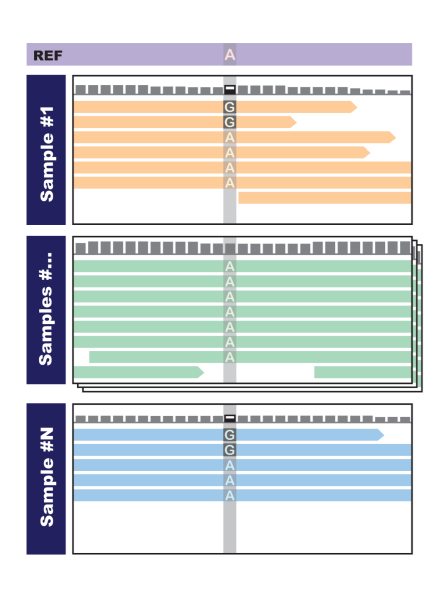


Joint Genotyping

Add a joint analysis step to take advantage of cohort / pop genetics data



Joint discovery empowers discovery at difficult sites

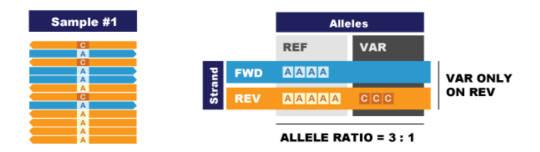


 If we analyze Sample #1 or Sample #N alone we are not confident that the variant is real

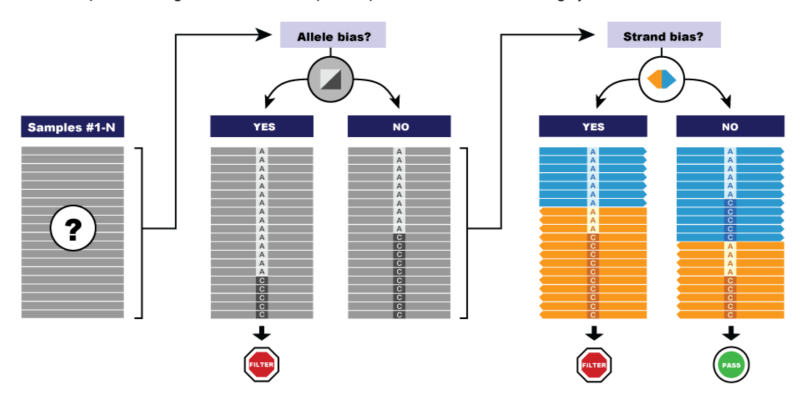
 If we see both samples then we are more confident that there is real variation at this site in the cohort

Joint discovery helps resolve bias issues

A. Single sample showing strand and allelic biases

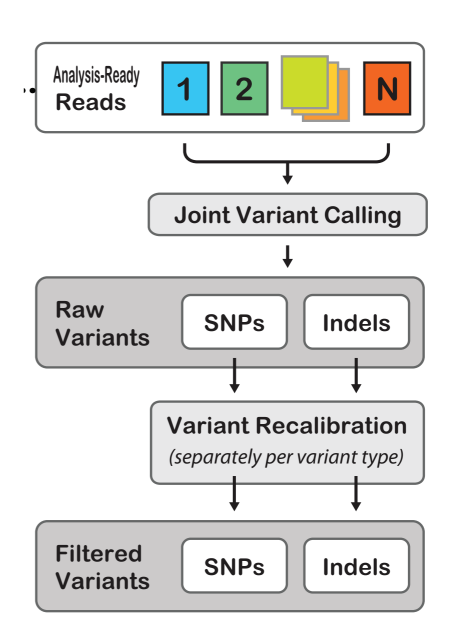


B. Decision process using evidence from multiple samples to filter out sites showing systematic biases



Classic approach to multi-sample variant discovery

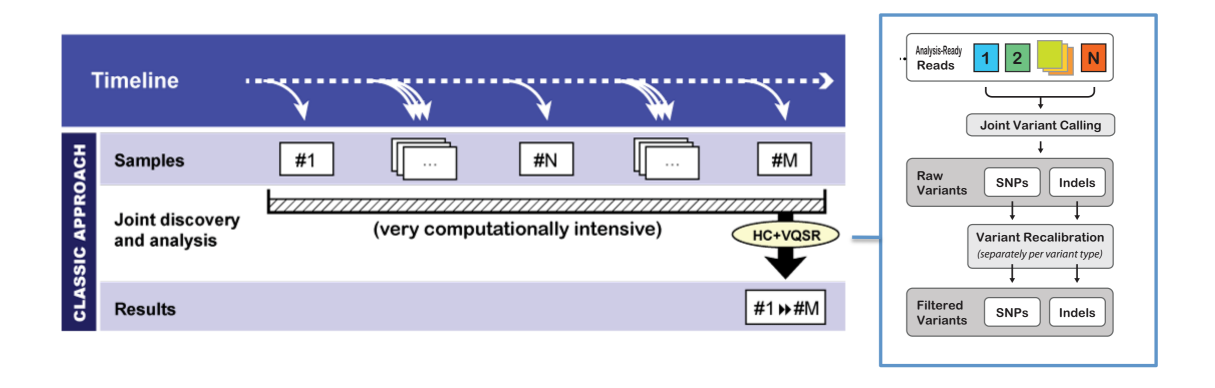
- Call variants jointly on all sample data
 - Scales badly -> limitations in amount of data that can be processed
 - Slow with UnifiedGenotyper (per-locus calculations)
 - Impossibly slow with
 HaplotypeCaller
 (so much extra work!)



But we want to use HaplotypeCaller because it is so much better!

Problems with the "all together" approach

- Computing costs
- The "N+1 problem"



##fileformat ##ALT ##ALT ##FILTER ##FILTER ##FORMAT ##INFO ##contig ##INFO ##reference ##contig **HEADER** ##reference #record headers **RECORDS** variant site record variant site record variant site record * Some tools may output an all-sites VCF that looks like what you can get using HC with -ERC BP_RESOLUTION but they do not provide an accurate estimate of

Regular* VCF

reference confidence.

HaplotypeCaller gVCF

-ERC GVCF

-ERC BP_RESOLUTION

##fileformat ##ALT ##FORMAT ##GVCFBlock ##INFO

##fileformat ##FILTER ##FORMAT ##contig ##reference

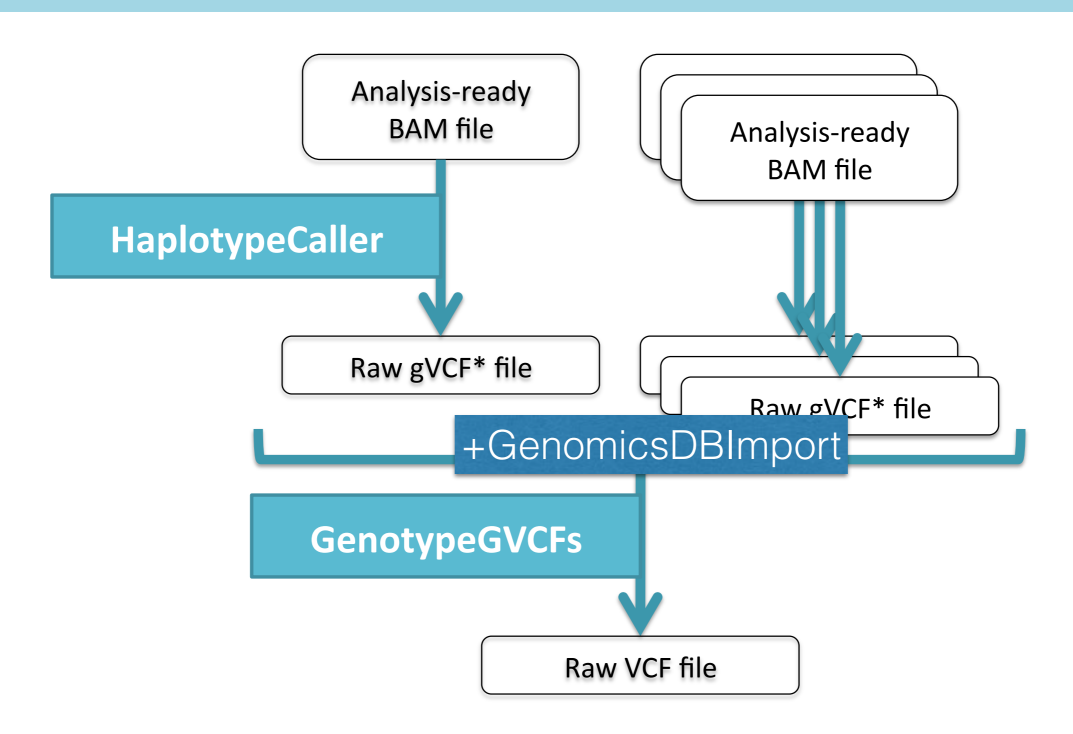
#record headers

- non-variant block record
- variant site record
 - non-variant block record
- variant site record
 - non-variant block record
- variant site record
 - non-variant block record

#record headers

- non-variant site record
- variant site record
- non-variant site record
- non-variant site record
- non-variant site record
- variant site record
- non-variant site record
- non-variant site record
- variant site record
- non-variant site record
- non-variant site record
- non-variant site record

Variant calling + joint genotyping workflow



Variant annotations provide key information to identify and remove artifacts!

VCF record for an A/G SNP at 22:49582364

22 49582364 AC=3; AF=0.50; AN=6; DP=87; MLEAC=3; MLEAF=0.50; MQ=71.31; MQ0=22; QD=2.29; SB=-31.76	INFO field	•	A G 19		198.	08.96 .	
		AC	No. chrom alt allele	osomes carrying	MLEAF	Max likelihood AF 🔽 🔽	
		AN	Total no. of chromosomes MC			RMS MAPQ of all reads	
		AF	Allele frequency			No. of MAPQ 0 reads at locus	
		DP	Depth of coverage QD QUAL score over depth			QUAL score over depth	
		MLEAC	Max likelihood AC		SB	Estimated strand bias score	
GT:DP:GQ	0/1:12:99.00 0/1:11:89.43 0					0/1:28:37.78	

VCF Files store variant information

```
##fileformat=VCFv4.1
##reference=1000GenomesPilot-NCBI36
##INFO=<ID=DP, Number=1, Type=Integer, Description="Total Depth">
##INFO=<ID=AF, Number=A, Type=Float, Description="Allele Frequency">
##INFO=\langleID=DB, Number=0, Type=Flag, Description="dbSNP membership, build 129^{\dagger}\rangle
##FILTER=<ID=s50, Description="Less than 50% of samples have data">
                                                                               Header
##FORMAT=<ID=GT, Number=1, Type=String, Description="Genotype">
##FORMAT=<ID=GQ, Number=1, Type=Integer, Description="Genotype Quality">
##FORMAT=<ID=DP, Number=1, Type=Integer, Description="Read Depth">
#CHROM POS
               ΙD
                          REF ALT
                                     OUAL FILTER INFO
                NA00001
                                NA00002
                                                NA00003
    FORMAT
       14370 rs6054257 G A
20
                                                      DP=14; AF=0.5; DB
                                               PASS
    GT:GQ:DP 0|0:48:1 1|0:48:8 1/1:43:5
                                                      DP=10;AF=0.333,0.667;DB
2.0
       1110696 rs6040355 A
                            G,T
                                       67
                                               PASS
    GT:GQ:DP 1|2:21:6 2|1:2:0
                                 2/2:35:4
                                                                              Variant
       1230237 .
                                               PASS
20
                                                      DP=13
                                                                              records
    GT:GQ:DP 0|0:54:7 0|0:48:4 0/0:61:2
20
       1234567 microsat1 GTCT
                                G,GTACT 50
                                               PASS
                                                      DP=9
                0/1:35:4
    GT:GQ:DP
                                0/2:17:2
                                                1/1:40:3
```

Official specification in

www.1000genomes.org/wiki/Analysis/Variant Call Format/vcf-variant-call-format-version-41

Variant Filtering

- By default, GATK is very permissive. It will output false positive sites!
- Two ways of filtering:
 - Hard filters
 - Variant recalibration

Hard Filters

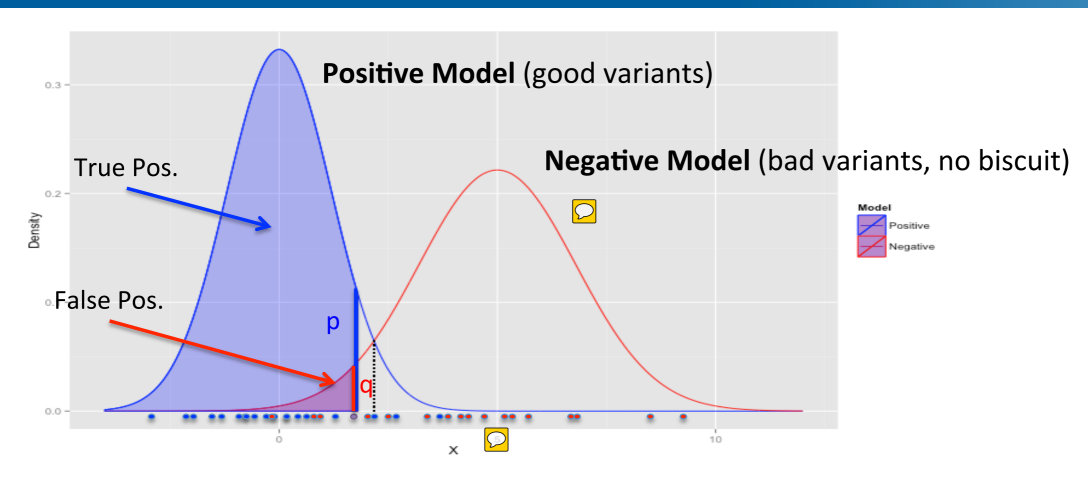
- User defined thresholds for each site. Hard to know where to make cut offs?
 - Mapping quality high enough
 - Depth above a minimum but not too high
 - Minor allele frequency above a minimum
 - Heterozygosity not too high

How variant recalibration works

Train on high-confidence known sites to determine the probability that other sites are true or false

- Assume annotations tend to form Gaussian clusters
- Build a "Gaussian mixture model" from annotations of **known variants** in our dataset
- Score **all variants** by where their annotations lie relative to these clusters
- Filter base on sensitivity to truth set

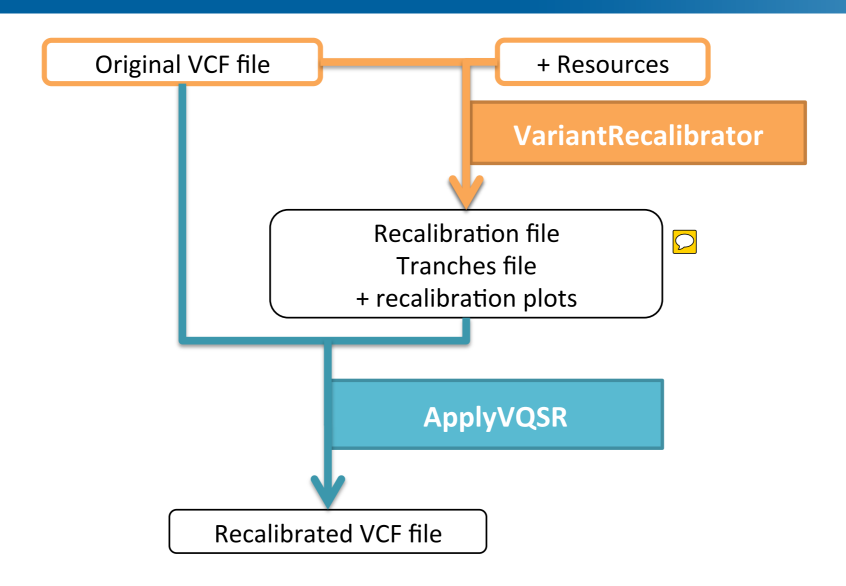
Actually two models: positive and negative



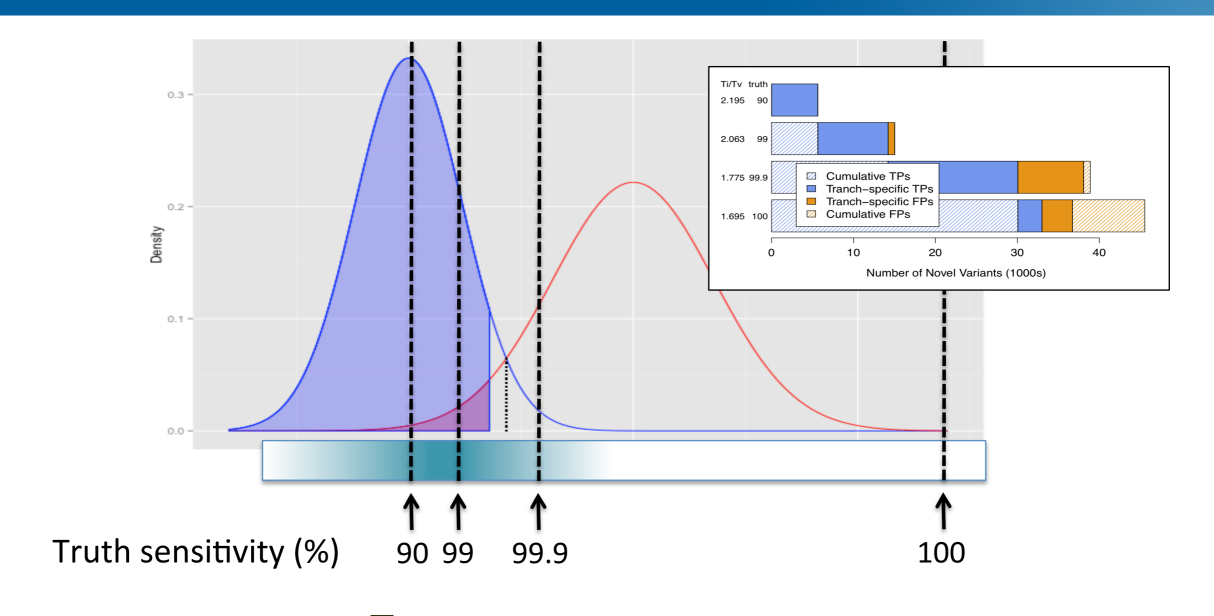
VQSLOD(x) = Log(p(x)/q(x))

Done for each annotation X then integrated into single overall VQSLOD

Step 1: VariantRecalibrator



Tranches: slices of sensitivity threshold values



Lower tranche = More stringent filtering

Where to get truth set?

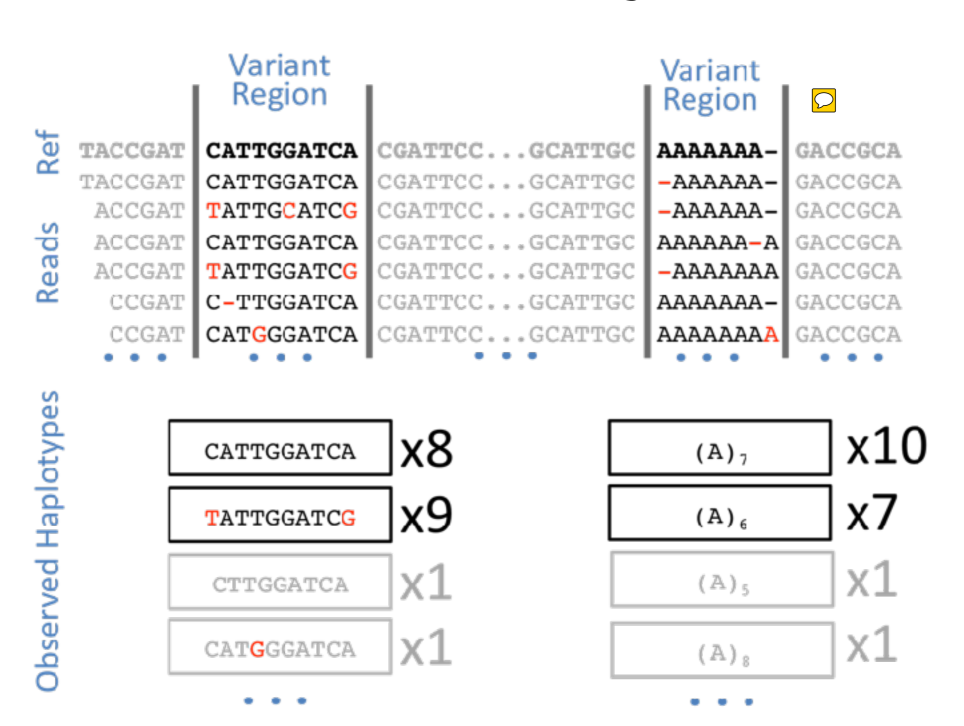
- Reference sets (e.g. 1000 genomes)
- Take your dataset, call SNPs, hard filter heavily, then use those as a truth set for recalibrating the unfiltered dataset.

Alternative Filtering

 Call SNPs using multiple programs and look for variants called in multiple programs, or combine the information in each.

• e.g. BAYSIC

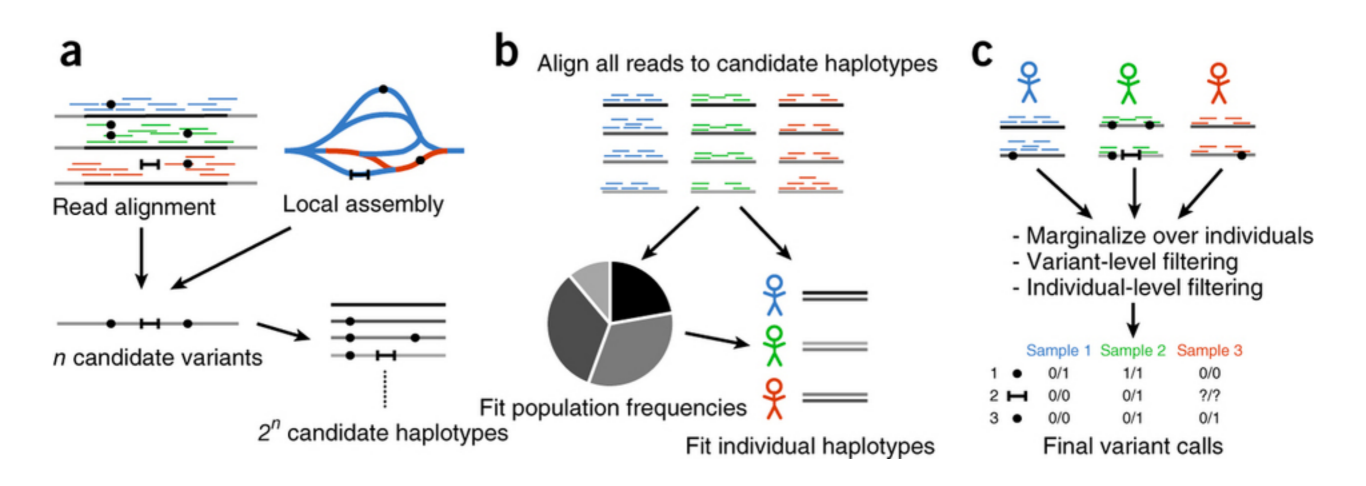
FreeBayes



FreeBayes

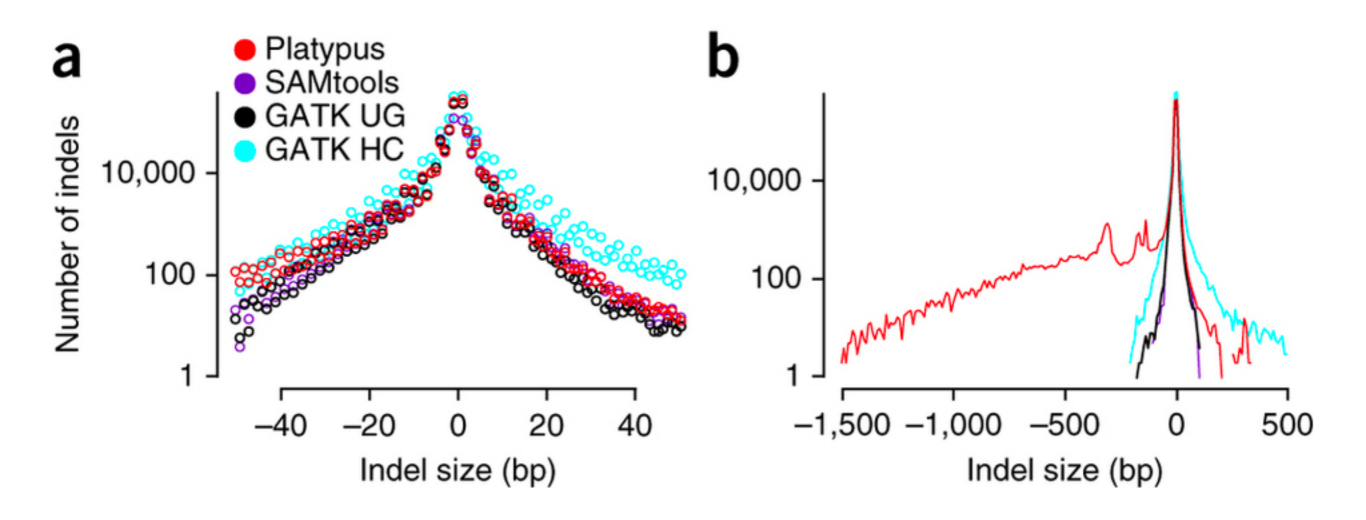
- Free and open source.
- Uses literal sequences of bases and haplotypes to call SNPs so is less affected by local alignment issues.
- Does not have "gvcf" n+1 method, although complicated work around exists.
- Generally faster than GATK, although RAM intensive.

Platypus



Includes local assembly, better for large indels

Platypus



Includes local assembly, better for large indels

ANGSD •

- Calls SNPs based on reads per site, no realignment.
- Outputs genotype likelihoods.
- Links with algorithms that use likelihood.
- Questionable with high diversity systems.