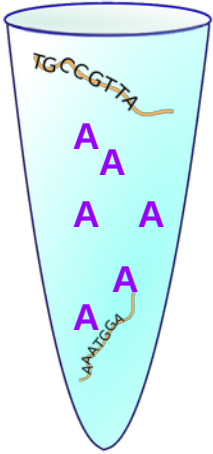


Genotype likelihoods, allele frequencies, and SNP calling from NGS data

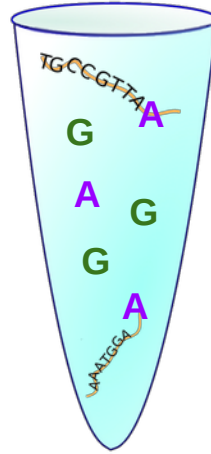
Tyler Linderoth
Physalia lcWGS course 2025

Why Probabilistic Methods?

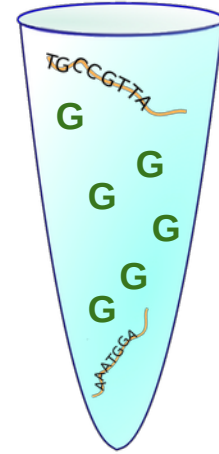
Why Probabilistic Methods?



The library for an individual homozygous for the **A** allele will consist only of **A**s.

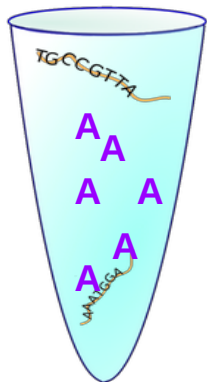


The library for a heterozygous individual at a site contains both **A**s and **G**s.



The library for an individual homozygous for the **G** allele will consist only of **G**s.

Why Probabilistic Methods?

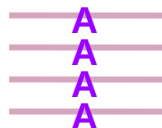


Sequence to average depth of 4x.

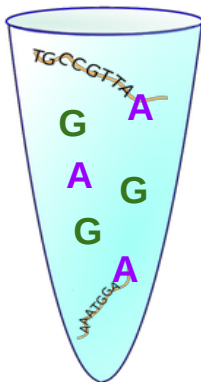
Depth \sim Poisson ($\lambda=4$)

$E[\text{depth}] = \lambda$

$\text{Var}[\text{depth}] = \lambda$



Expect 4 reads, all with A's at this ref position.



Sequencing (sampling) the two different alleles is just like flipping a coin.

A alleles \sim Binomial ($n \text{ reads}, p=0.5$)

$E[A \text{ depth}] = np = 0.5n$

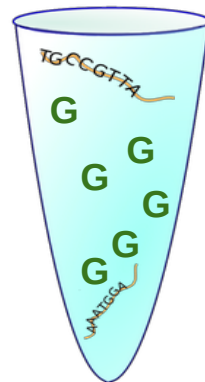
$\text{Var}[A \text{ depth}] = np(1-p) = 0.25n$



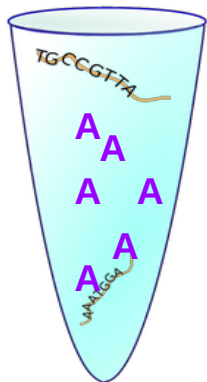
vs



Expect 2 A alleles and 2 G alleles



Why Probabilistic Methods?

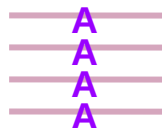


Sequence to average
depth of 4x.

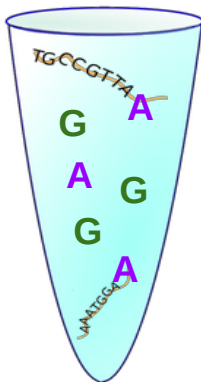
Depth \sim Poisson ($\lambda=4$)

$E[\text{depth}] = \lambda$

$\text{Var}[\text{depth}] = \lambda$



Expect 4 reads, all with
As at this ref position.



Sequencing (sampling) the two
different alleles is just like
flipping a coin.

A alleles \sim Binomial ($n \text{ reads}, p=0.5$)

$E[A \text{ depth}] = np = 0.5n$

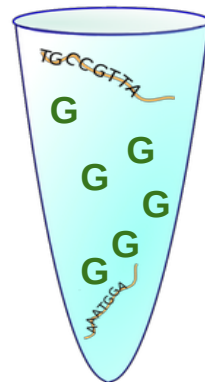
$\text{Var}[A \text{ depth}] = np(1-p) = 0.25n$



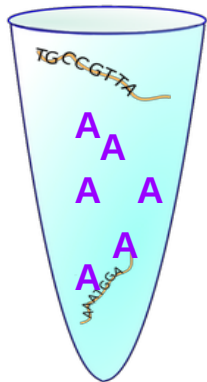
vs



Expect 2 A alleles
and 2 G alleles



Why Probabilistic Methods?



Sequence to average depth of 4x.

Depth \sim Poisson ($\lambda=4$)

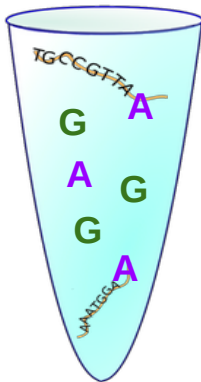
$E[\text{depth}] = \lambda$

$\text{Var}[\text{depth}] = \lambda$



A
T
A
A

Sequencing error.
Rates of $\sim 0.1\%$ for
some Illumina
platforms.



Sequencing (sampling) the two different alleles is just like flipping a coin.

A alleles \sim Binomial ($n \text{ reads}, p=0.5$)

$E[A \text{ depth}] = np = 0.5n$

$\text{Var}[A \text{ depth}] = np(1-p) = 0.25n$

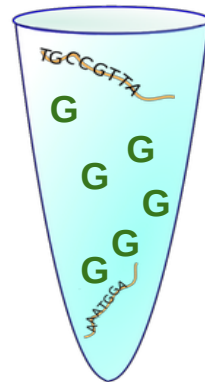


vs



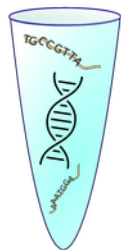
A
A
G
A

Expect 2 A alleles
and 2 G alleles



G
G
G
G

SAMtools mpileup representation of sequencing data for two individuals



Individual 1 data

6,	DEGEGG
6,	DEGEGG
6,	DEGEGG
6,	DEGEGG
6,	DEGEGG
6,	DEGEGG
6,	DEGEGG
7,^].	DEGEGGE
7,	DEGEGGG
8,.^],	DEGEGGGB
8,,	DEGEGGGB
8,,	DEGEGGGB
8,,	DEGEGGGB
9,.^],	DEGEGGGBE
9	.G...gG,,	DEGEGGGBG
9,,	DEGEGGGBG

Individual 2 data

9,	DABGIIIII
9,	DABGIIIII
9,	DABGIIIII
10,^].	DABGIIIIIE
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	DABGIIIIII
10,	D3BGIIIIII
10,	D3BGIIIIII

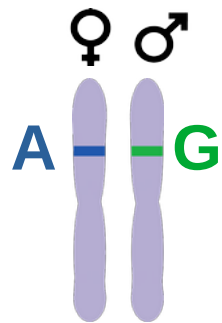
Each row is a different site in the reference genome

reads

base IDs

base qualities

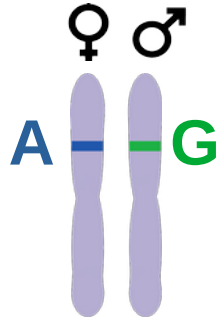
Example sequencing data for one individual at Chr1:472



<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	DEG

Maternally and paternally inherited chromosome 1 of a diploid individual.

Example sequencing data for one individual at Chr1:472



Maternally and paternally inherited chromosome 1 of a diploid individual.



<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	D EG



ASCII character “D” = decimal value 68

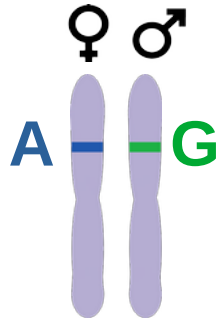


$68 - 33 = \text{base quality of } 35$



$$P(\text{error}) = \epsilon = 10^{\frac{-35}{10}} = 0.00032$$

Example sequencing data for one individual at Chr1:472



<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	DEG

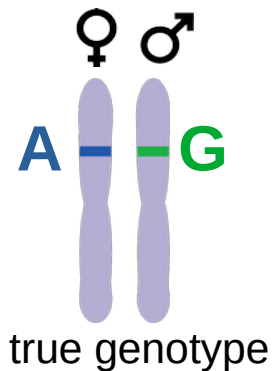
Maternally and paternally inherited chromosome 1 of a diploid individual.

This individual could have any of the following 10 genotypes (we can only see the sequencing data):

AA, AC, AG, AT, CG, CC, CT, GG, GT, TT

How do we figure out which genotype they are most likely to have based on the observed sequence data?

Genotype likelihoods



<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	DEG

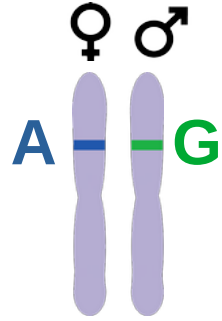
$P(\text{Data} | \text{Genotype} = bh) = L(\text{Genotype } bh) = \text{likelihood of genotype } bh$

$b, h \in \{A, C, G, T\}$

Possible genotypes: AA, AC, AG, AT, CG, CC, CT, GG, GT, TT

What is the likelihood that the individual's genotype is AC?

Possible genotypes: AA, **AC**, AG, AT, CG, CC, CT, GG, GT, TT



true genotype

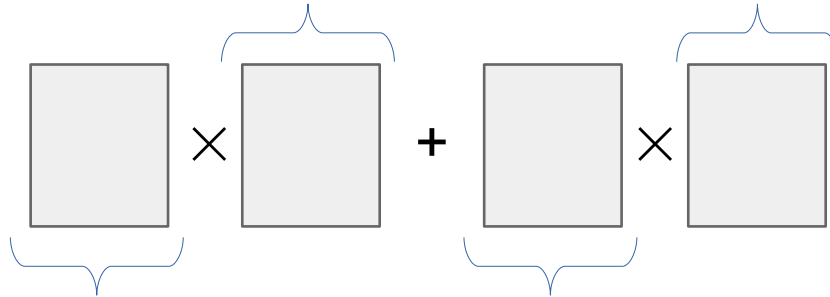


<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	DEG

$P(\text{observed } i\text{-th read} \mid \text{A allele})$

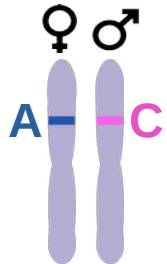
$P(\text{observed } i\text{-th read} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$



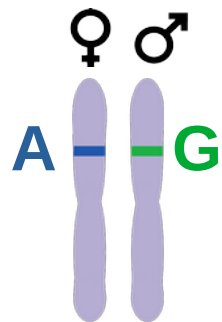
Probability of sampling
maternal chromosome
(A allele)

Probability of sampling
paternal chromosome
(C allele)



Assumed true genotype

What is the likelihood that the individual's genotype is AC?



true genotype

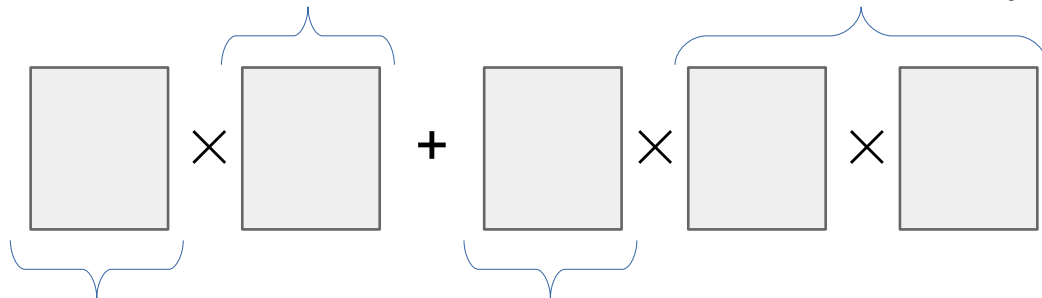


<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	A AG	DEG

$P(\text{observed read A} \mid \text{A allele})$

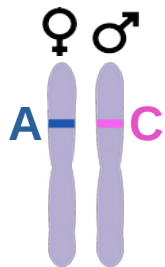
$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$



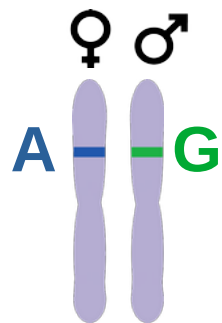
**Probability of
sampling maternal
chromosome (A allele)**

**Probability of sampling
paternal chromosome
(C allele)**

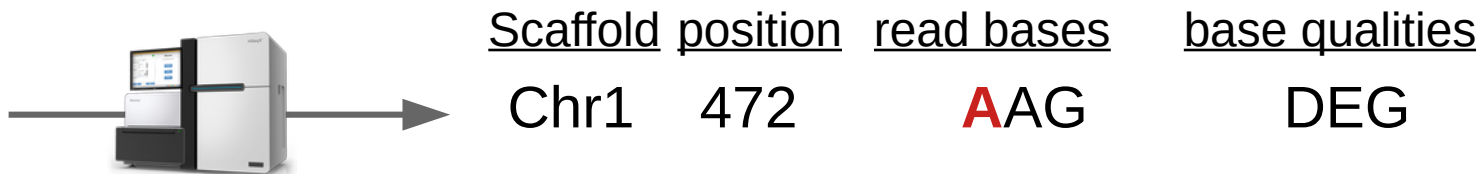


Assumed true genotype

What is the likelihood that the individual's genotype is AC?



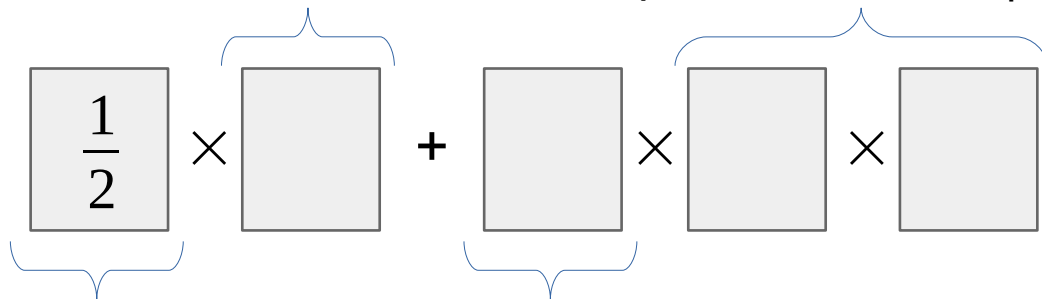
true genotype



$P(\text{observed read A} \mid \text{A allele})$

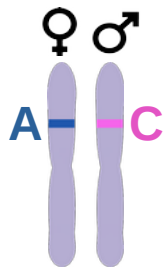
$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$



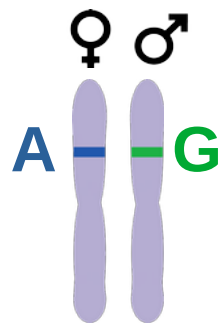
Probability of sampling
maternal chromosome
(A allele)

Probability of sampling
paternal chromosome
(C allele)



Assumed true genotype

What is the likelihood that the individual's genotype is AC?



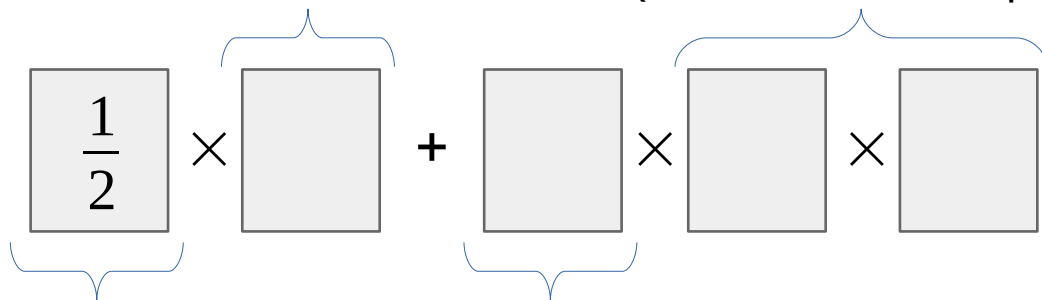
true genotype



P(observed read A | A allele)

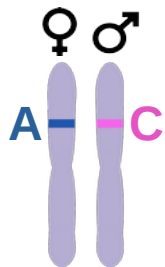
P(observed read A | C allele)

$$P(\text{Data} | \text{Genotype} = \text{AC}) =$$



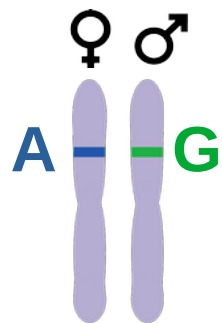
Probability of sampling
maternal chromosome
(A allele)

Probability of sampling
paternal chromosome
(C allele)



Assumed true genotype

What is the likelihood that the individual's genotype is AC?



true genotype



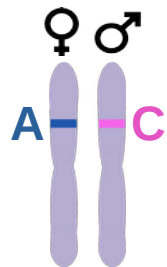
<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	A AG	DEG

$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

$$\underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of sampling paternal chromosome (C allele)}} + \underbrace{\phantom{\frac{1}{2}}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}}$$

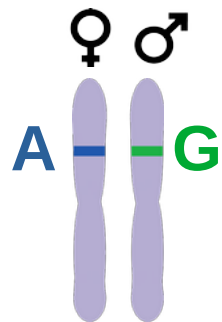


Assumed true genotype

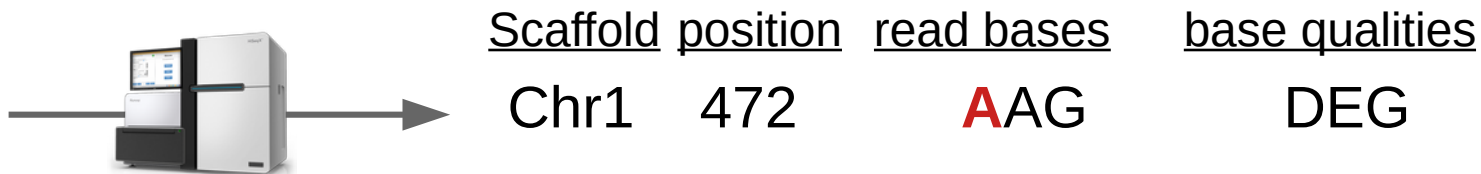
Probability of sampling maternal chromosome (A allele)

Probability of sampling paternal chromosome (C allele)

What is the likelihood that the individual's genotype is AC?



true genotype

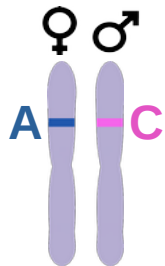


$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

$$\underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of sampling paternal chromosome (C allele)}} + \underbrace{\phantom{\frac{1}{2}}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}}$$

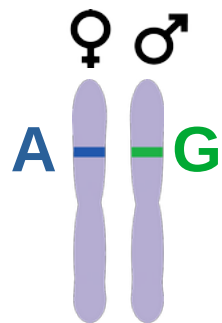


Assumed true genotype

Probability of sampling maternal chromosome (A allele)

Probability of sampling paternal chromosome (C allele)

What is the likelihood that the individual's genotype is AC?



true genotype



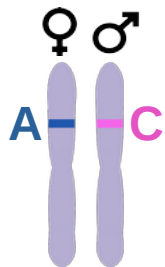
<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	A AG	DEG

$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

$$\underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of sampling paternal chromosome (C allele)}} + \underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}}$$

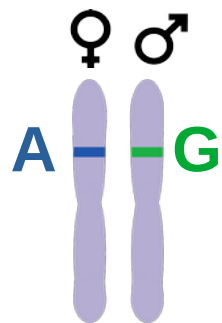


Assumed true genotype

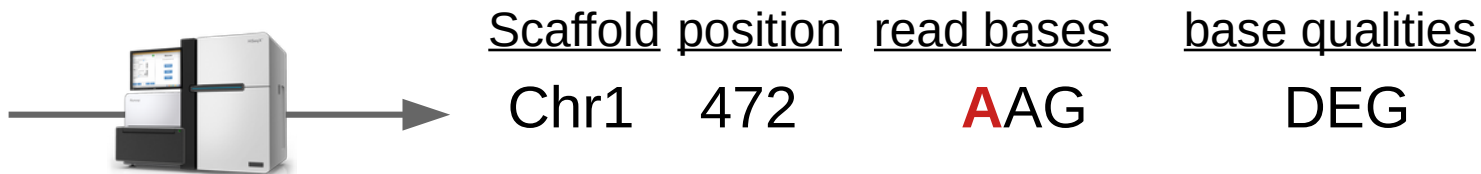
Probability of sampling
maternal chromosome
(A allele)

Probability of sampling
paternal chromosome
(C allele)

What is the likelihood that the individual's genotype is AC?



true genotype

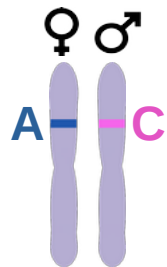


$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

$$\underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of sampling paternal chromosome (C allele)}} + \underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{}_{\text{Probability of sampling paternal chromosome (C allele)}}$$

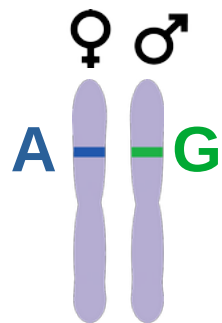


Assumed true genotype

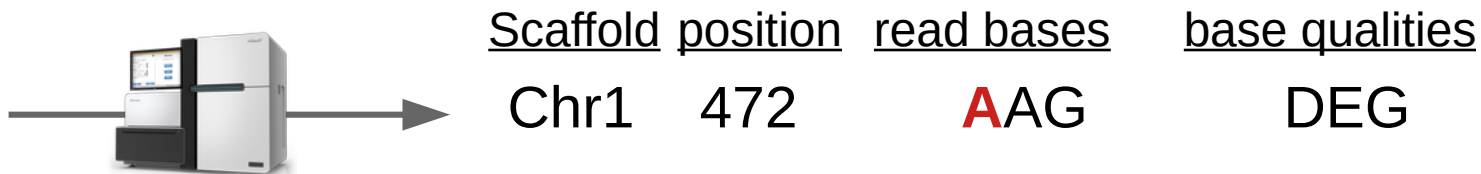
Probability of sampling maternal chromosome (A allele)

Probability of sampling paternal chromosome (C allele)

What is the likelihood that the individual's genotype is AC?



true genotype

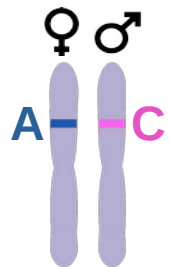


$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

$$= \underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of sampling maternal chromosome (A allele)}} + \underbrace{\frac{1}{2}}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{\epsilon_1}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{\frac{1}{3}}_{\text{Probability of sampling paternal chromosome (C allele)}}$$

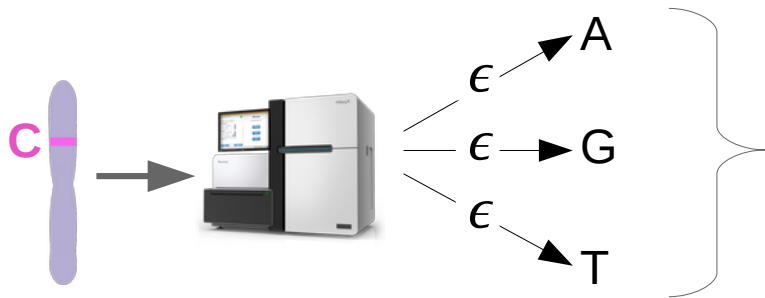


Assumed true genotype

Probability of sampling maternal chromosome (A allele)

Probability of sampling paternal chromosome (C allele)

What is the likelihood that the individuals genotype is AC?



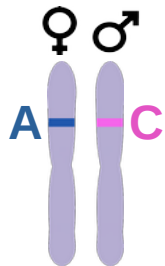
Assuming equal probability of changing to any of the possible 3 erroneous bases.

$P(\text{observed read A} \mid \text{A allele})$

$P(\text{observed read A} \mid \text{C allele})$

$$P(\text{Data} \mid \text{Genotype} = \text{AC}) =$$

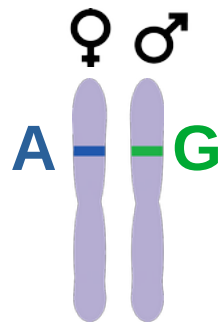
$$\underbrace{\frac{1}{2}}_{\text{Probability of sampling maternal chromosome (A allele)}} \times \underbrace{1 - \epsilon_1}_{\text{Probability of observing A given A allele}} + \underbrace{\frac{1}{2}}_{\text{Probability of sampling paternal chromosome (C allele)}} \times \underbrace{\epsilon_1}_{\text{Probability of observing A given C allele}} \times \underbrace{\frac{1}{3}}_{\text{Probability of observing A given C allele}}$$



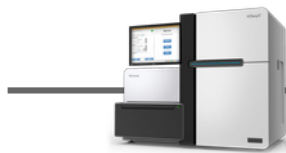
Assumed true genotype

Probability of sampling maternal chromosome (A allele)

Probability of sampling paternal chromosome (C allele)

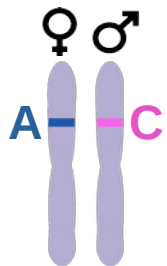


true genotype

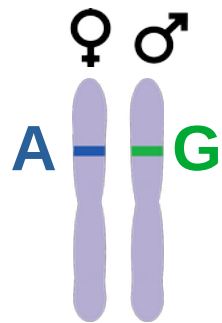


<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AAG	DEG

$$P(\text{Data} | \text{Genotype} = AC) = \left[\left[\frac{1}{2} \times (1 - \epsilon_1) + \frac{1}{2} \times \epsilon_1 \times \frac{1}{3} \right] \times \left[\frac{1}{2} \times (1 - \epsilon_2) + \frac{1}{2} \times \epsilon_2 \times \frac{1}{3} \right] \right]$$



Assumed true genotype



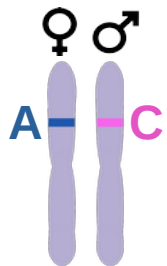
true genotype



<u>Scaffold</u>	<u>position</u>	<u>read bases</u>	<u>base qualities</u>
Chr1	472	AA G	DEG

$P(\text{Data} | \text{Genotype} = AC) =$

$$\left[\left[\frac{1}{2} \times (1 - \epsilon_1) + \frac{1}{2} \times \epsilon_1 \times \frac{1}{3} \right] \times \left[\frac{1}{2} \times (1 - \epsilon_2) + \frac{1}{2} \times \epsilon_2 \times \frac{1}{3} \right] \times \left[\frac{1}{2} \times \epsilon_3 \times \frac{1}{3} + \frac{1}{2} \times \epsilon_3 \times \frac{1}{3} \right] \right]$$



Assumed true genotype

General genotype likelihood expression

$$P(\text{Data} \mid \text{Genotype} = bh) = \prod_{i=1}^{r \text{ reads}} \left(\frac{L_b^{(i)}}{2} + \frac{L_h^{(i)}}{2} \right)$$

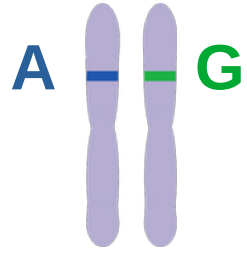
$$b, h \in \{A, C, G, T\}$$

$$L_b^{(i)} = P(\text{observed read} = x_i \mid \text{assumed true allele} = b)$$

$$L_h^{(i)} = P(\text{observed read} = x_i \mid \text{assumed true allele} = h)$$

$$\left. \begin{array}{l} L_b^{(i)} \\ L_h^{(i)} \end{array} \right\} \begin{array}{l} \frac{\epsilon_i}{3} \text{ if } b, h \neq x_i \\ 1 - \epsilon_i \text{ if } b, h = x_i \end{array}$$

Representation of genotype likelihoods in ANGSD



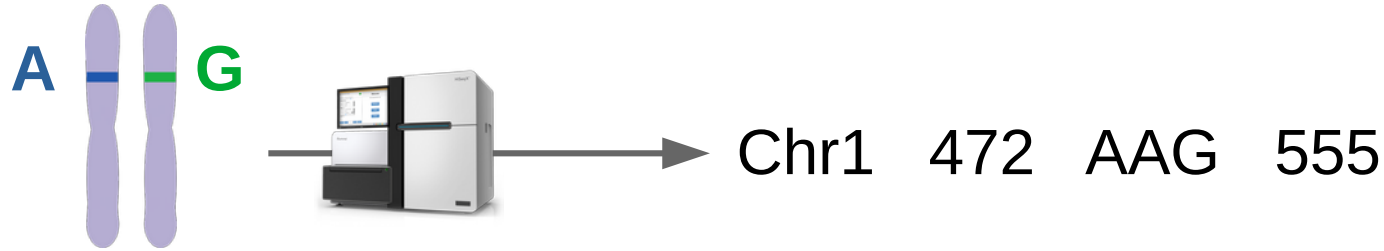
Chr1 472 AAG 555

ASCII character "5" = decimal value 53

$53 - 33 = \text{base quality of } 20$

$$\epsilon = 10^{\frac{-20}{10}} = 0.01$$

Representation of genotype likelihoods in ANGSD

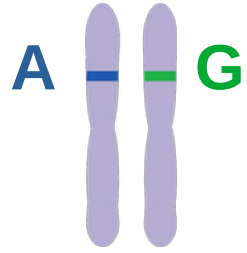


Instead of a single genotype, AG, we have a distribution over all possible genotypes:

Genotype	AA	AC	AG	AT	CC	CG	CT	GG	GT	TT
Log ₁₀ likelihood	-2.49	-3.08	-0.91	-3.08	-7.43	-5.26	-7.43	-4.96	-5.26	-7.43

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Representation of genotype likelihoods in ANGSD



Chr1 472 AAG 555

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Log ₁₀ likelihood	-2.49	-3.08	-0.91	-3.08	-7.43	-5.26	-7.43	-4.96	-5.26	-7.43



Maximum likelihood estimate of the genotype

$$\epsilon = 10^{\frac{-20}{10}} = 0.01$$

Exercise. Calculate genotype likelihoods with ANGSD.

Estimating allele frequencies

When genotypes are known, allele frequencies can be calculated by simply counting alleles.

Minor
allele
frequency
(MAF)

	Ind1	Ind2	Ind3	Ind4	Ind5	Ind6	Ind7	Ind8	Ind9	Ind10
Site 1	0	0	1	0	0	0	1	1	0	0
Site 2	0	2	0	0	1	0	0	0	1	0

= 0.3

= 0.4

Genotype notation

0 = zero minor alleles

1 = one minor alleles

2 = two minor alleles

Estimating allele frequencies

When genotypes are known, allele frequencies can be calculated by simply counting alleles.

	Ind1	Ind2	Ind3	Ind4	Ind5	Ind6	Ind7	Ind8	Ind9	Ind10	Minor allele frequency (MAF)
Site 1	0	0	1	0	0	0	1	1	0	0	= 0.3
Site 2	0	2	0	0	1	0	0	0	1	0	= 0.4

But how do you estimate allele frequencies when you have a distribution of genotype likelihoods?

Maximum likelihood estimation of allele frequencies

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$

D_i = sequencing data for individual i

f = population minor allele frequency

Genotype notation

0 = zero minor alleles

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Maximum likelihood estimation of allele frequencies

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} \underbrace{P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)}_{\text{This is the likelihood of genotype } g \text{ for individual } i \text{ calculated as shown previously.}}$$

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D_i = sequencing data for individual i

f = population minor allele frequency

Genotype notation

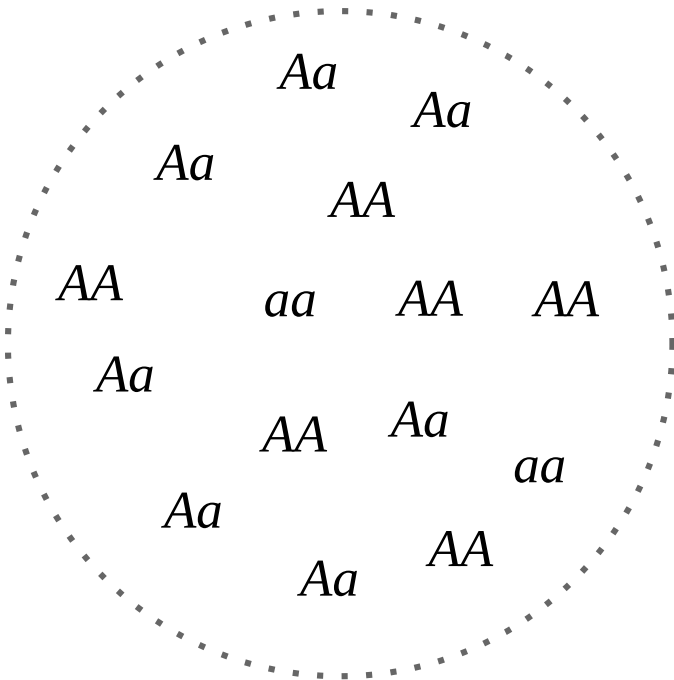
0 = zero minor alleles

1 = one minor alleles

2 = two minor alleles

Genotype frequencies under Hardy-Weinberg Equilibrium (HWE)

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$

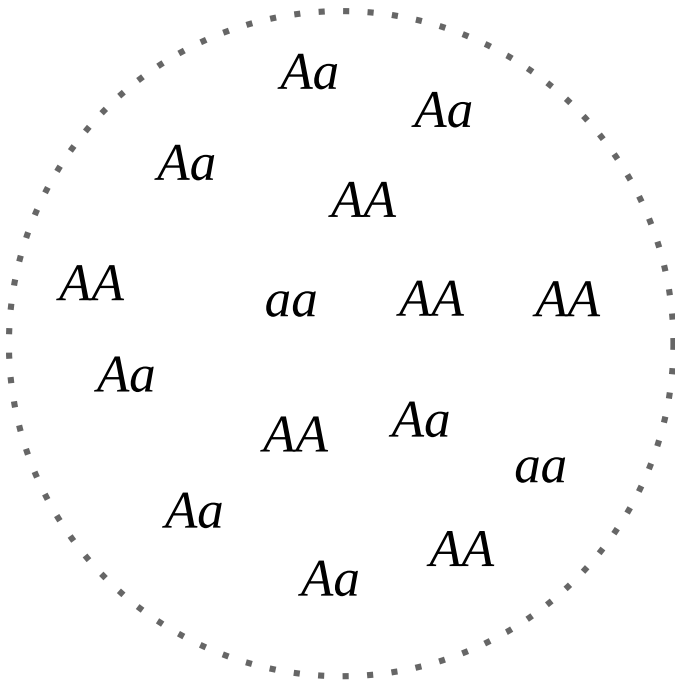


An “infinitely” large population of sexually reproducing diploid organisms segregating for alleles A and a , and for which

- Mating is random.
- Generations are nonoverlapping.
- Allele frequencies are the same in males and females.
- No migration, mutation, or selection.

Genotype frequencies under Hardy-Weinberg Equilibrium (HWE)

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$



Offspring are formed through independent draws of gametes from this population, expressed as $(f_A + f_a)^2$, which expanded yields:

$$f_{AA} = f_A^2 = (1 - f_a)^2 = P(\text{Genotype} = AA | f_a)$$

$$f_{Aa} = 2f_A f_a = 2(1 - f_a)f_a = P(\text{Genotype} = 1 | f_a)$$

$$f_{aa} = f_a^2 = P(\text{Genotype} = 2 | f_a)$$

Maximum likelihood estimation of allele frequencies

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \underbrace{\sum_{g \in \{0,1,2\}}}_{\text{Summing over the possible genotypes accounts for genotyping uncertainty.}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$

Summing over the possible genotypes accounts for genotyping uncertainty.

Marginal probabilities are used to account for uncertainty in various quantities associated with low coverage sequencing. In general, for random variables X and Y

$$P(X=x) = \sum_y P(X=x|Y=y) P(Y=y)$$

Maximum likelihood estimation of allele frequencies

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$

The value of f that maximizes the likelihood function above yields a **maximum likelihood estimate of f** :

$$\hat{f} = \operatorname{argmax}_f P(\text{Data}|f)$$

Using the ML allele frequency estimate to identify polymorphic sites

$$P(\text{Data}|f) = \prod_{i=1}^{n \text{ individuals}} \sum_{g \in \{0,1,2\}} P(D_i | \text{Genotype}_i = g) P(\text{Genotype}_i = g | f)$$

$$\hat{f} = \operatorname{argmax}_f P(\text{Data}|f) \quad \leftarrow \text{maximum likelihood estimate of } f$$

Probability of the sequencing data
when $f = 0$, i.e., the site is
monomorphic (null case).


$$\lambda = -2 \ln \left(\frac{P(\text{Data}|f_0)}{P(\text{Data}|\hat{f})} \right) = -2 [\ln(\underbrace{P(\text{Data}|f_0))}_{\text{monomorphic null case}}) - \ln(P(\text{Data}|\hat{f}))]$$

$$\lambda \sim \chi^2(1 \text{ degree of freedom}) \longrightarrow$$

Call SNPs at a given level of
statistical confidence.

Exercise. Estimate allele frequencies with ANGSD.

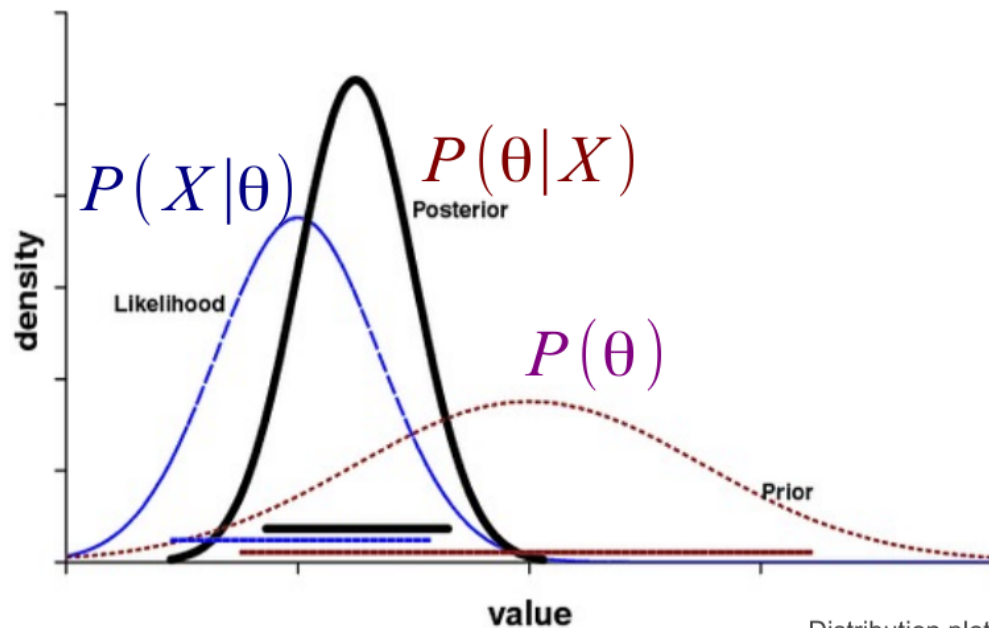
Can we use other information from our data to further increase our genotyping accuracy?

6,	DEGEGG	9,	DABGIIIIII	5	>AB/A	6	FGG	
6,	DEGEGG	9,	DABGIIIIII	5	>ABDA	6	3GGGGG	
6,	DEGEGG	9,	DABGIIIIII	5	>ABDA	6	3GGGBG	
6,	DEGEGG	10,^].	DABGIIIIIE	5	>AB/A	6	3GGGBG	
6,	DEGEGG	10,	DABGIIIIII	5	>AB/A	6	3GGGBG	
6,	DEGEGG	10,	DABGIIIIII	5	>ABDA	6	BGGGBG	
6,	DEGEGG	10,	DABGIIIIII	5	>ABDA	6	C,...	5G/BGB	
7,^].	DEGEGGE	10,	DABGIIIIII	5	>ABDA	6	5GIBGB	
7,	DEGEGGG	10,	DABGIIIIII	5	>ABDA	6	5GIBGB	
8,^],	DEGEGGGB	10,	DABGIIIIII	5	>ABDA	6	5GIBGB	
8,	DEGEGGGB	10,	DABGIIIIII	5	>AB/A	6	DGIBGB	
8,	DEGEGGGB	10,	DABGIIIIII	5	>ABAA	6	DGIBGB	
8,	DEGEGGGB	10,	DABGIIIIII	5	>/BAA	6	DGIBGB	
9,^],	DEGEGGGBE	10,	DABGIIIIII	5	>BBAA	6	DGIBGB	
9	.G...gG,,	DEGEGGGBG	10,	D3BGIIIIII	5	...,C	>BBAA	6	DGIBGB	
9,,,	DEGEGGGBG	10,	D3BGIIIIII	5	>BBAA	6	/GIBGB	

Wouldn't it be awesome if you knew what the frequency of C was in the population.

Bayesian Inference

$$P(\theta|X) = \frac{P(X|\theta)P(\theta)}{P(X)} = \frac{P(X|\theta)P(\theta)}{\sum_{\theta} P(X|\theta)P(\theta)}$$



X = Data

θ = Parameter

Genotype posterior probabilities to improve genotyping accuracy

Using Bayes' Theorem, the posterior probability of genotype g is

$$P(\text{Genotype} = g | \text{Data}) = \frac{P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}{\sum_{g \in \{0,1,2\}} P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}$$

Genotype posterior probabilities to improve genotyping accuracy

Using Bayes' Theorem, the posterior probability of genotype g is

Likelihood of genotype g
calculated as shown
previously.



$$P(\text{Genotype} = g | \text{Data}) = \frac{P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}{\sum_{g \in \{0,1,2\}} P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}$$

Genotype posterior probabilities to improve genotyping accuracy

Using Bayes' Theorem, the posterior probability of genotype g is

Likelihood of genotype g
calculated as shown
previously.

Given an estimate of the
population minor allele
frequency, f , under HWE

$$P(\text{Genotype}=0|f) = (1-f)^2$$

$$P(\text{Genotype}=1|f) = 2f(1-f)$$

$$P(\text{Genotype}=2|f) = f^2$$

$$P(\text{Genotype}=g|\text{Data}) = \frac{P(\text{Data}|\text{Genotype}=g) P(\text{Genotype}=g)}{\sum_{g \in \{0,1,2\}} P(\text{Data}|\text{Genotype}=g) P(\text{Genotype}=g)}$$

Genotype posterior probabilities to improve genotyping accuracy

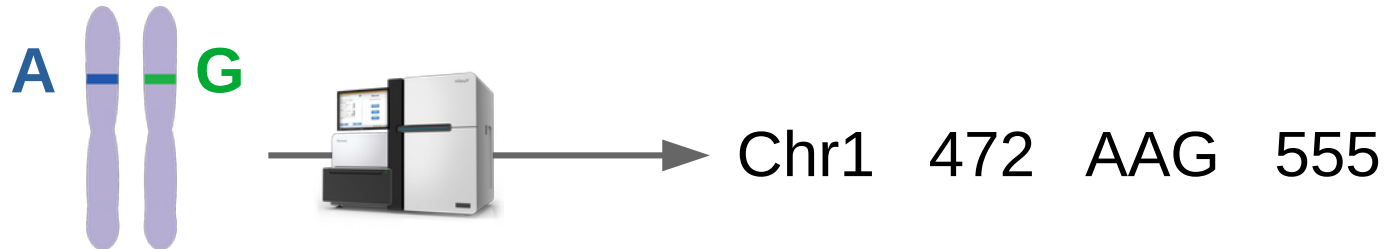
Using Bayes' Theorem, the posterior probability of genotype g is

Likelihood of genotype g
calculated as shown
previously.

Note: factors like inbreeding can
easily be incorporated into the
genotype posterior probabilities
by conditioning on the allele
frequency and inbreeding
coefficient.

$$P(\text{Genotype} = g | \text{Data}) = \frac{P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}{\sum_{g \in \{0,1,2\}} P(\text{Data} | \text{Genotype} = g) P(\text{Genotype} = g)}$$

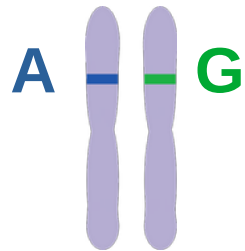
Example genotype posterior probability distribution



Assume we estimate $f(A) = 0.7$, $f(G) = 0.3$

Genotype	Log ₁₀ likelihood	Prior	Posterior probability
AA	-2.49	$P(\text{Genotype} = AA) = 0.7^2 = 0.49$	0.03
AG	-0.91	$P(\text{Genotype} = AG) = 2 \times 0.7 \times 0.3 = 0.42$	0.97
GG	-4.96	$P(\text{Genotype} = GG) = 0.3^2 = 0.09$	0.00

Example genotype posterior probability distribution



Chr1 472 AAG 555

Assume we estimate $f(A) = 0.7$, $f(G) = 0.3$

We could call most probable genotype, AG, and have an associated degree of confidence (prob = 0.97).

Genotype	Log_{10} likelihood	Prior	Posterior probability
AA	-2.49	$P(\text{Genotype} = AA) = 0.7^2 = 0.49$	0.03
AG	-0.91	$P(\text{Genotype} = AG) = 2 \times 0.7 \times 0.3 = 0.42$	0.97
GG	-4.96	$P(\text{Genotype} = GG) = 0.3^2 = 0.09$	0.00

Exercise. Calculate genotype posterior probabilities and call genotypes with ANGSD.