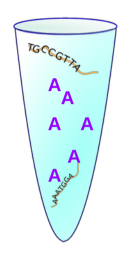
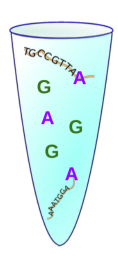
Genotype likelihoods, allele frequencies, and SNP calling from NGS data

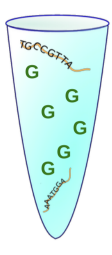
Tyler Linderoth
Physalia lcWGS course 2025



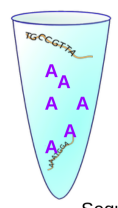
The library for an individual homozygous for the A allele will consist only of As.

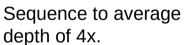


The library for a heterozygous individual at a site contains both As and **G**s.



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





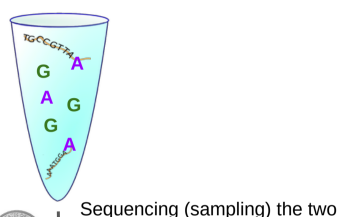
Depth ~ Poisson ($\lambda = 4$)

 $E[depth] = \lambda$

 $Var[depth] = \lambda$



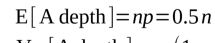
Expect 4 reads, all with As at this ref position.

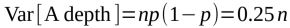




different alleles is just like flipping a coin.

A alleles \sim Binomial (nreads, p=0.5)



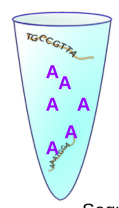


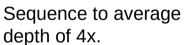




G G granded

Expect 2 A alleles and 2 G alleles





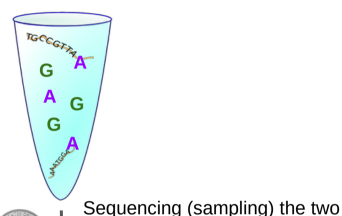
Depth \sim Poisson ($\lambda = 4$)

 $E[depth] = \lambda$

 $Var[depth] = \lambda$

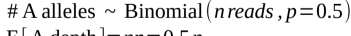
Expect 4 reads, all with

A As at this ref position.





different alleles is just like flipping a coin.



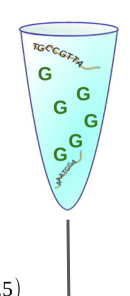
E[A depth] = np = 0.5 n

Var[A depth] = np(1-p) = 0.25 n

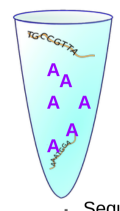




Expect 2 A alleles and 2 G alleles







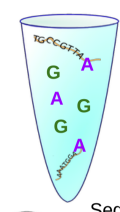
Sequence to average depth of 4x.

Depth ~ Poisson ($\lambda = 4$)

 $E[depth] = \lambda$

 $Var[depth] = \lambda$

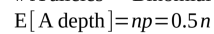
Sequencing error.
Rates of ~0.1% for some Illumina platforms.





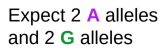
Sequencing (sampling) the two different alleles is just like flipping a coin.

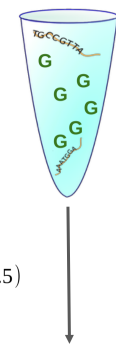
A alleles \sim Binomial (n reads, p = 0.5)



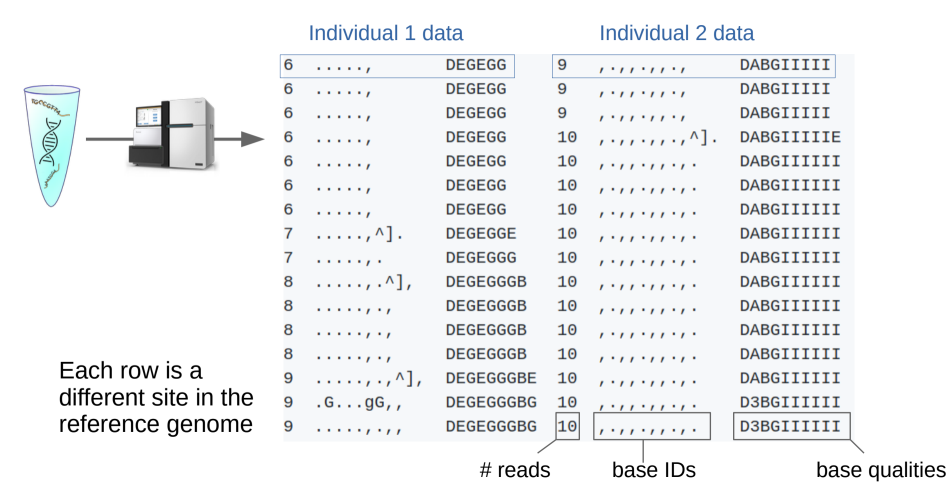
Var[A depth] = np(1-p) = 0.25 n



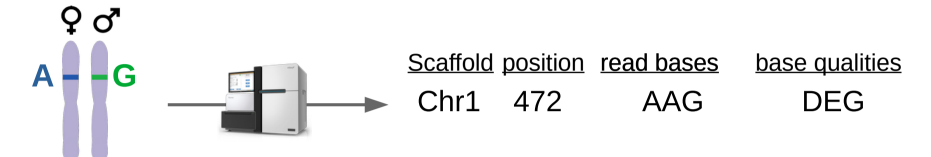




SAMtools mpileup representation of sequencing data for two individuals



Example sequencing data for one individual at Chr1:472

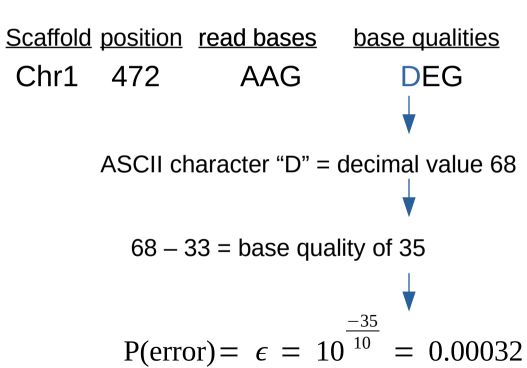


Maternally and paternally inherited chromosome 1 of a diploid individual.

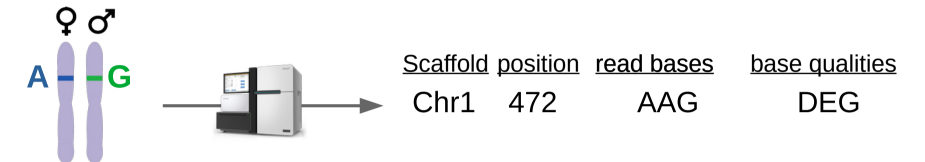
Example sequencing data for one individual at Chr1:472



Maternally and paternally inherited chromosome 1 of a diploid individual.



Example sequencing data for one individual at Chr1:472



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



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

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

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

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



Scaffold position read bases

(C allele)

AAG

472

base qualities DFG

P(observed *i*-th read | C allele)

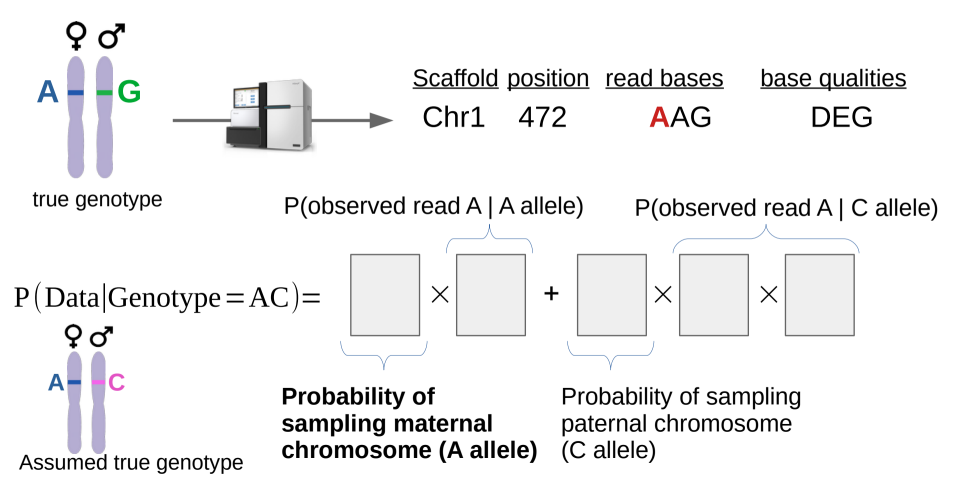
true genotype P(observed *i*-th read | A allele)

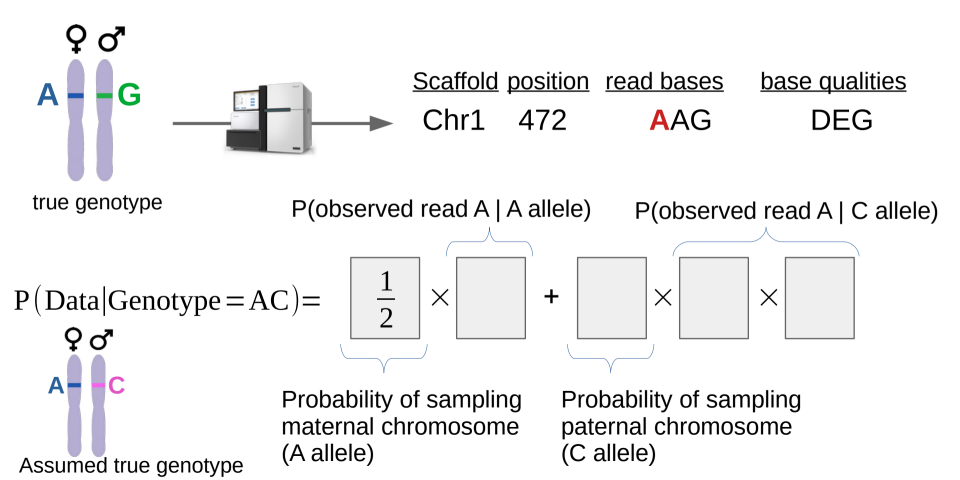
P(Data|Genotype = AC) =X

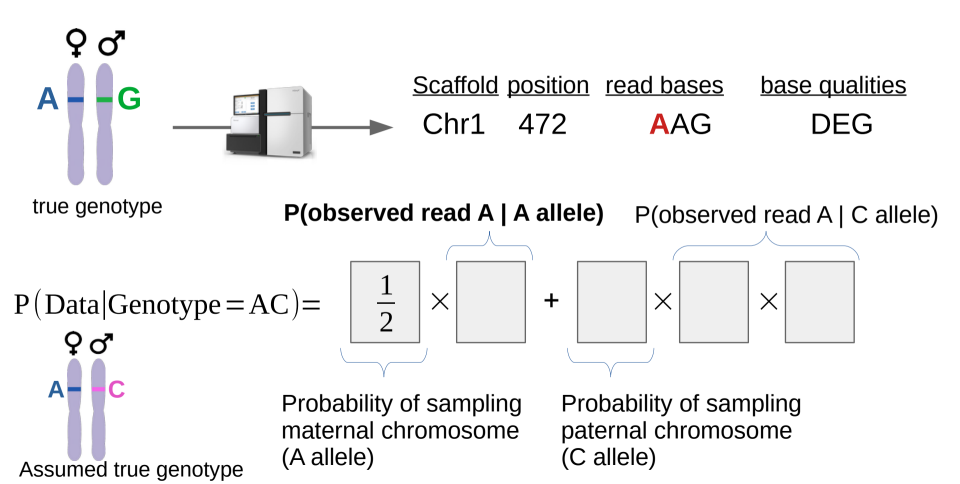
Assumed true genotype

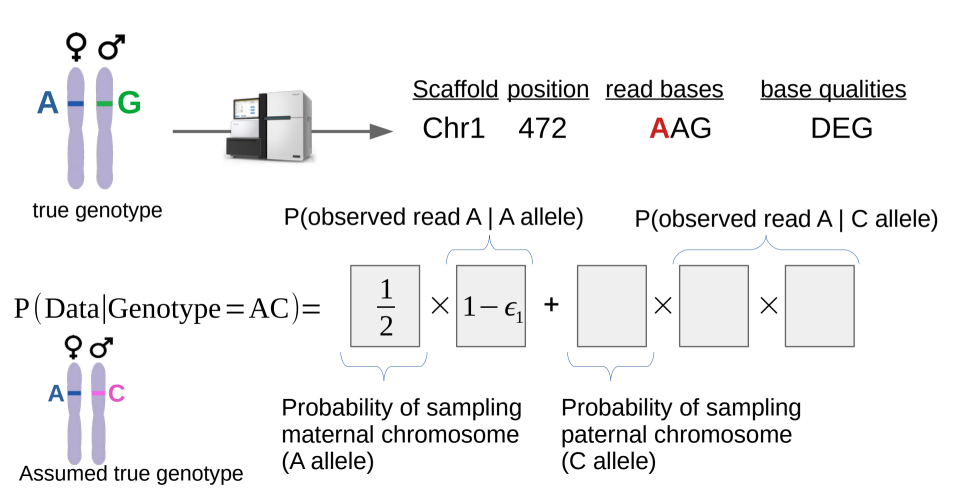
Probability of sampling maternal chromosome (A allele)

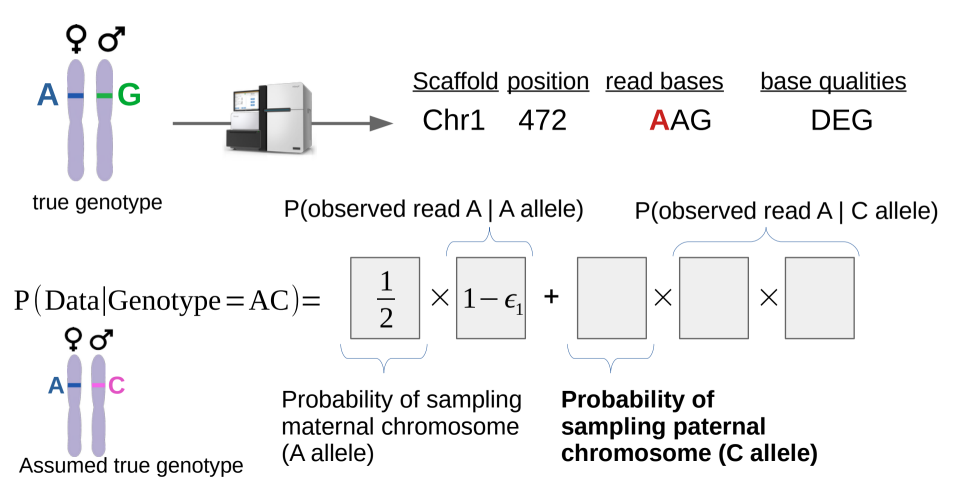
Probability of sampling paternal chromosome

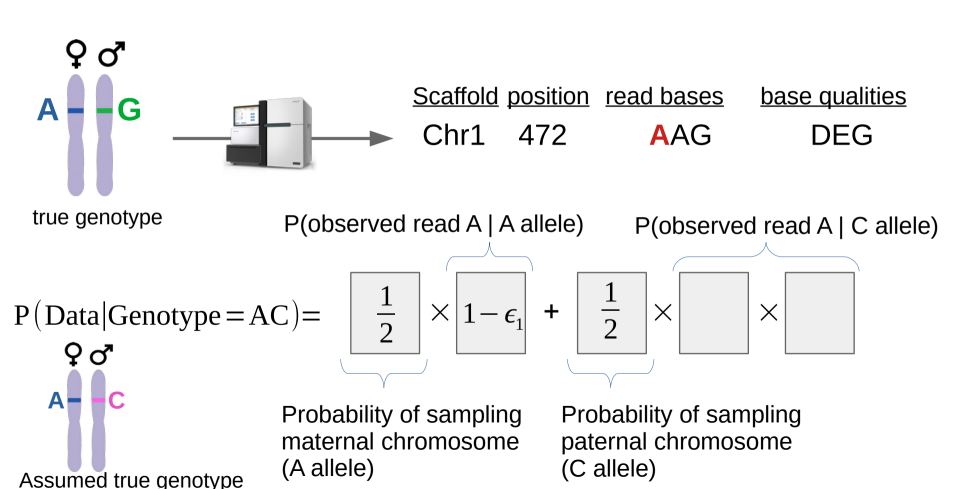


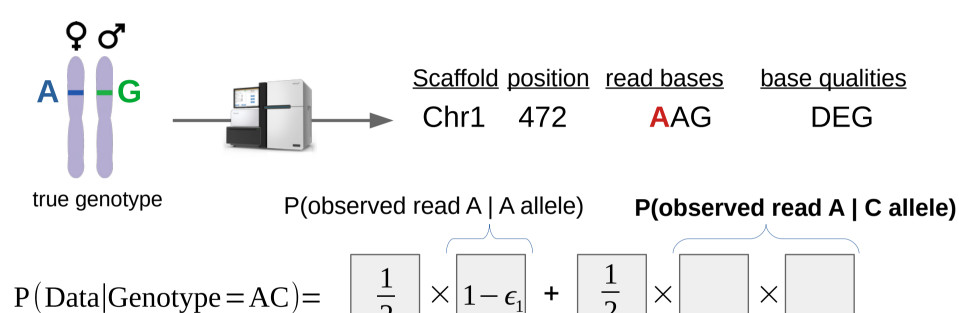






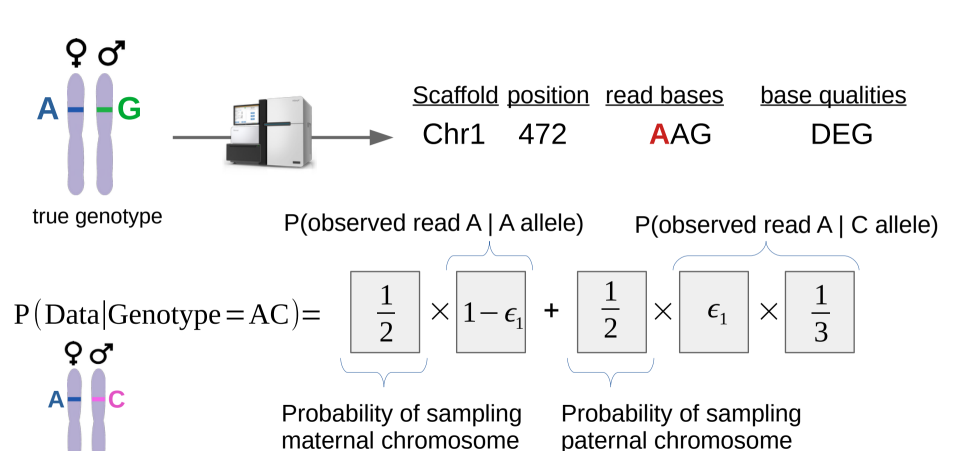






(A allele)

Assumed true genotype



(C allele)

P(observed read A | A allele)

Probability of sampling

maternal chromosome

(A allele)

Assumed true genotype



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

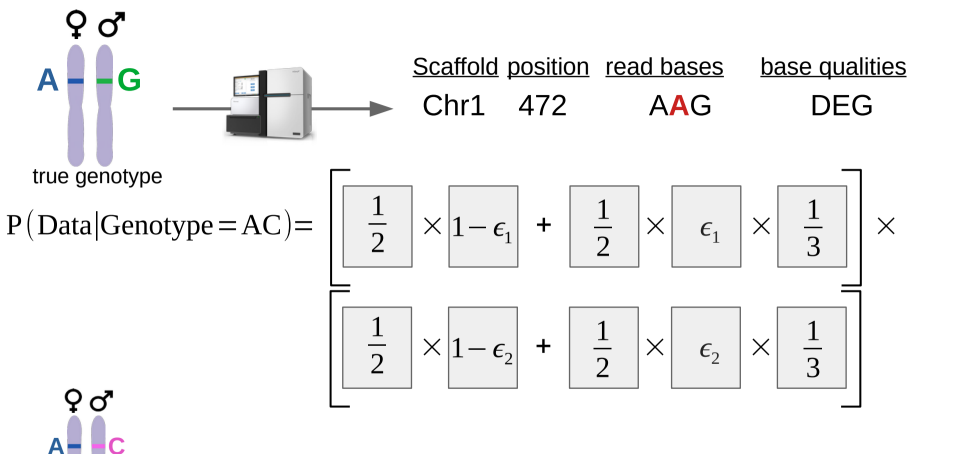
Probability of sampling

paternal chromosome

(C allele)

P(observed read A | C allele)

P(Data|Genotype=AC)=
$$\frac{1}{2} \times 1 - \epsilon_1 + \frac{1}{2} \times \epsilon_1 \times \frac{1}{3}$$



Assumed true genotype



AAG DEG

base qualities

true genotype
$$P(Data|Genotype = AC) = \begin{bmatrix} \frac{1}{2} \\ \end{bmatrix}$$

Assumed true genotype

Scaffold position read bases

General genotype likelihood expression

$$r$$
r

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

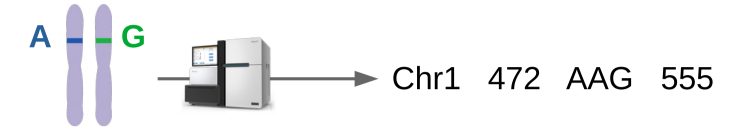
P(Data | Genotype=
$$bh$$
)= $\prod_{i=1}^{r \text{ reads}} \left(\frac{L_b^{(i)}}{2} + \frac{L_h^{(i)}}{2}\right)$

 $\begin{vmatrix} L_b^{(i)} = P(\text{observed read} = x_i | \text{assumed true allele} = b) \\ L_h^{(i)} = P(\text{observed read} = x_i | \text{assumed true allele} = h) \end{vmatrix} \begin{vmatrix} \frac{\epsilon_i}{3} & \text{if } b, h \neq x_i \\ 1 - \epsilon_i & \text{if } b, h = x_i \end{vmatrix}$

Representation of genotype likelihoods in ANGSD



Representation of genotype likelihoods in ANGSD

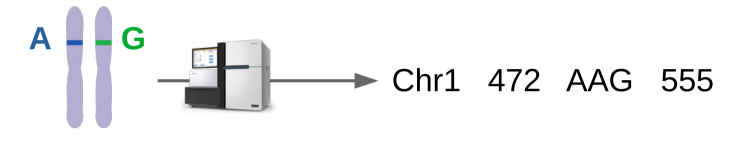


Instead of a single genotype, AG, we have a distribution over all possible genotypes:

Genotype	AA	AC	AG	AT	СС	CG	СТ	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

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

Representation of genotype likelihoods in ANGSD



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

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

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

Exercise. Calculate genotype likelihoods with ANGSD.

Estimating allele frequencies

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

S		Ind2		Ind4	Ind5	Ind6	Ind7	Ind8	Ind9	Ind10		equency (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

Minor

allele

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		equency (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

Minor

allele

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

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

$$D_i = \text{ sequencing data for individual } i$$

$$f = \text{ population minor allele frequency}$$

$$\frac{\text{Genotype notation}}{0 = \text{zero minor alleles}}$$

$$1 = \text{one minor alleles}$$

$$2 = \text{two minor alleles}$$

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

This the likelihood of genotype *g* for individual *i* calculated as shown previously.

$$D_i$$
= sequencing data for individual i
 f = population minor allele frequency

2 = two minor alleles

Genotype notation

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

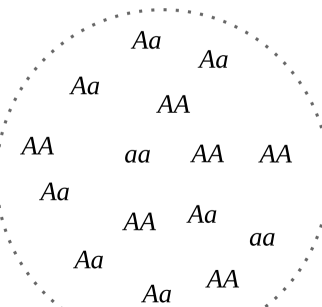
This the likelihood of genotype q for individual *i* calculated as shown previously.

Hardy-Weinberg frequency of genotypes.

D_i = sequencing data for individual i
f = population minor allele frequency

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)

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

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

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

expressed as
$$(f_A + f_a)^2$$
, which expanded yields:
 $f_{AA} = f_A^2 = (1 - f_a)^2 = P(Genotype = AA|f_a)$
 $f_{Aa} = 2f_A f_a = 2(1 - f_a)f_a = P(Genotype = 1|f_a)$

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

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

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:

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

Using the ML allele frequency estimate to identify polymorphic sites

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

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

maximum likelihood estimate of f

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

$$\lambda = -2\ln\left(\frac{P(Data|f_0)}{P(Data|\hat{f})}\right) = -2\left[\ln\left(\frac{P(Data|f_0)}{P(Data|\hat{f})}\right) - \ln\left(\frac{P(Data|\hat{f})}{P(Data|\hat{f})}\right)\right]$$

$$\lambda \sim \chi^2(1 \, \text{degree of freedom})$$
 — Call SNPs at a given level of statistical confidence.

Exercise. Estimate allele frequencies with ANGSD.

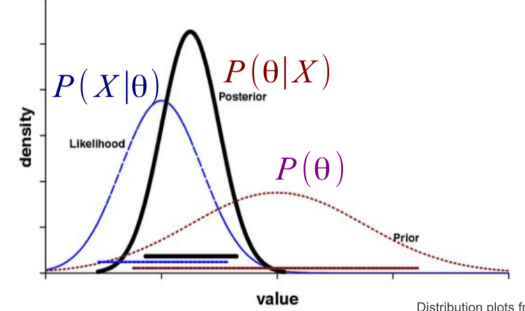
Can we use other information from our data to further increase our genotyping accuracy?

6	,	DEGEGG	9	/ · / / · / / · /	DABGIIIII	5	,,.,.	>AB/A	6	.,,	FGC 🗀	
6	,	DEGEGG	9	, . , , . , , . ,	DABGIIIII	5	,,.,.	>ABDA	6	.,,	3GGDGD	
6	,	DEGEGG	9	, . , , . , , . ,	DABGIIIII	5	,,.,.	>ABDA	6	.,,	3GGBGB	
6	,	DEGEGG	10	,.,,.,,^].	DABGIIIIE	5	,,.,.	>AB/A	6	.,,	3GGBGB	
6	,	DEGEGG	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>AB/A	6	.,,	3GGBGB	
6	,	DEGEGG	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABDA	6	.,,	BGGBGB	
6	,	DEGEGG	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABDA	6	C,,	5G/BGB]
7	,^].	DEGEGGE	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABDA	6	.,,	5GIBGB	
7	, .	DEGEGGG	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABDA	6	.,,	5GIBGB	
8	,.^],	DEGEGGGB	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABDA	6	.,,	5GIBGB	
8	, . ,	DEGEGGGB	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>AB/A	6	.,,	DGIBGB	
8	, . ,	DEGEGGGB	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>ABAA	6	.,,	DGIBGB	
8	,	DEGEGGGB	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>/BAA	6	.,,	DGIBGB	
9	,.,^],	DEGEGGGBE	10	, . , , . , , . , .	DABGIIIII	5	,,.,.	>BBAA	6	.,,	DGIBGB	
9	.GgG,,	DEGEGGGBG	10	, . , , . , , . , .	D3BGIIIII	5	,,.,C	>BBAA	6	.,,	DGIBGB	
9		DEGEGGGBG	10		D3BGIIIII	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 $\theta = Parameter$

Distribution plots from Bink 2008

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

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

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

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

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

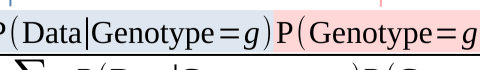
Likelihood of genotype *q* calculated as shown previously.

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

$$P(Genotype=0|f)=(1-f)^2$$

$$P(Genotype=1|f)=2f(1-f)$$

$$P(Genotype=2|f)=f^2$$



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

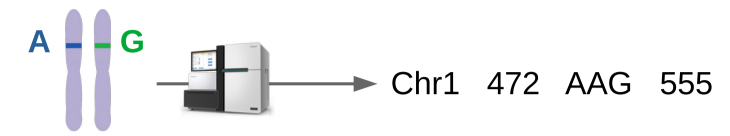
Using Bayes' Theorem, the posterior probability of genotype g is

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.

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

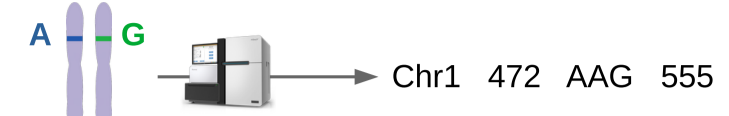
Example genotype posterior probability distribution



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

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

Example genotype posterior probability distribution



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	$P(Genotype = AA) = 0.7^2 = 0.49$	0.03		
AG	-0.91	$P(Genotype = AG) = 2 \times 0.7 \times 0.3 = 0.42$	0.97		
GG	-4.96	$P(Genotype = GG) = 0.3^2 = 0.09$	0.00		

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