

# Calling variants from low-coverage NGS data

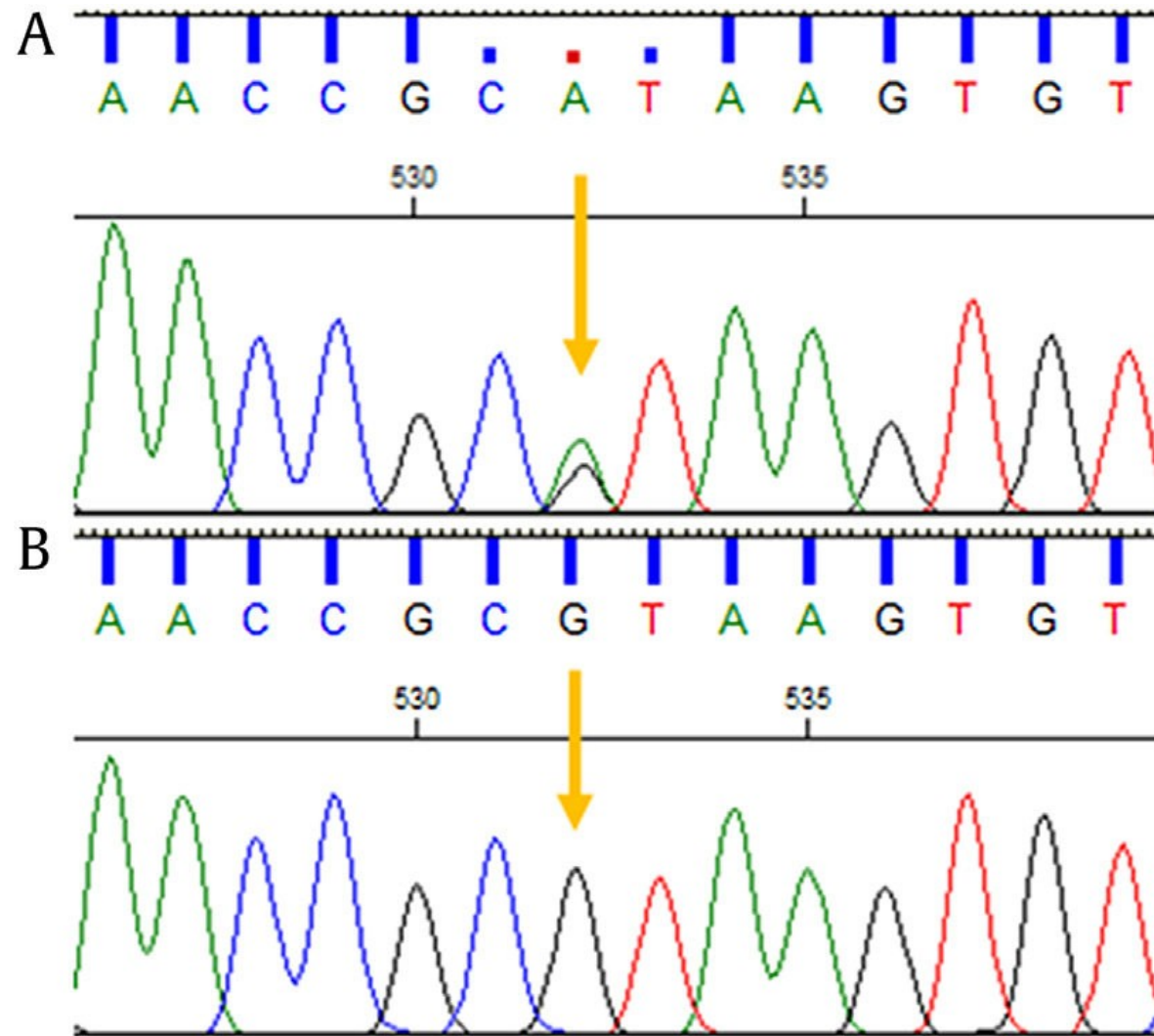
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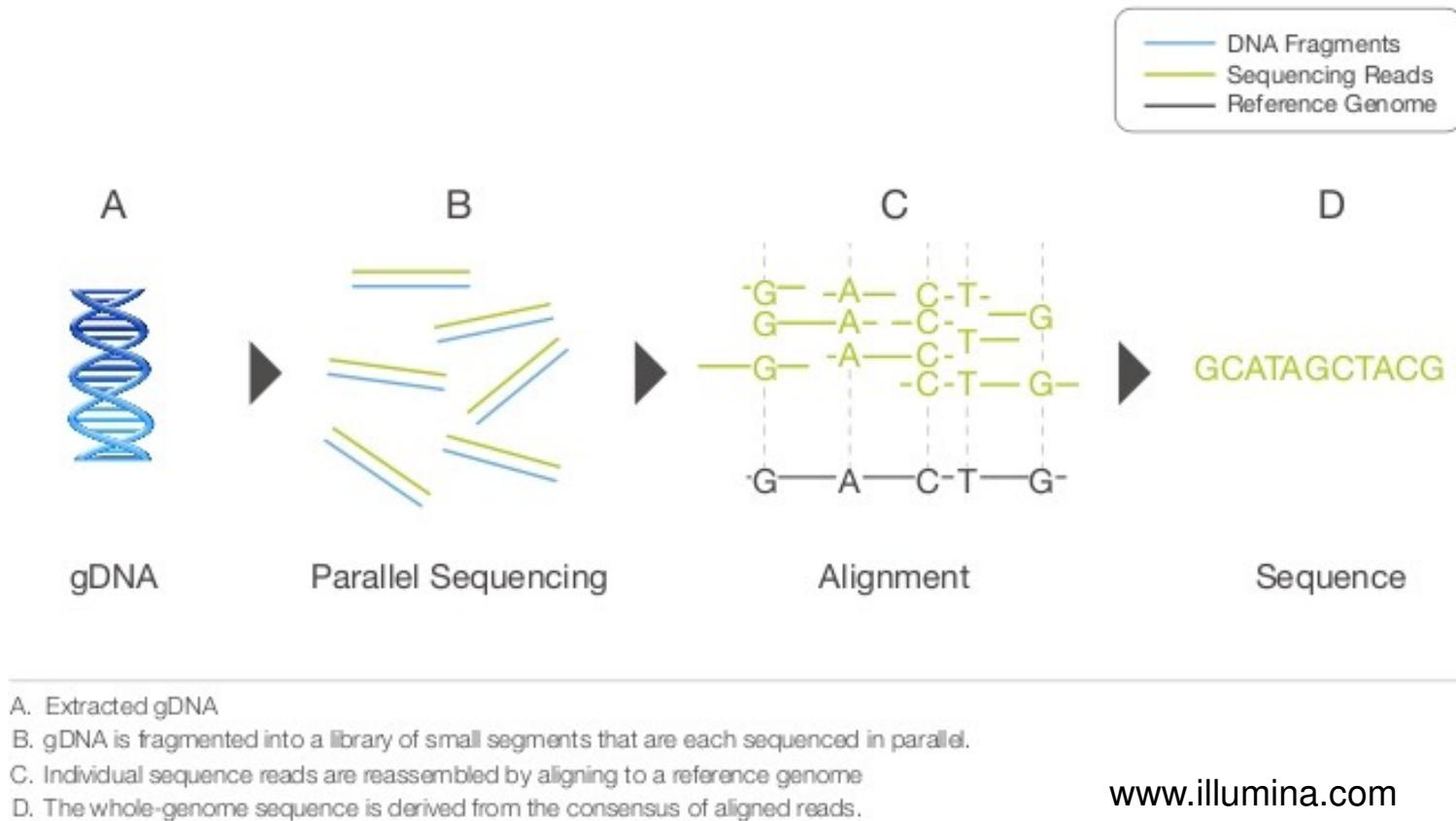


Region  
Hovedstaden

# Sanger Sequencing

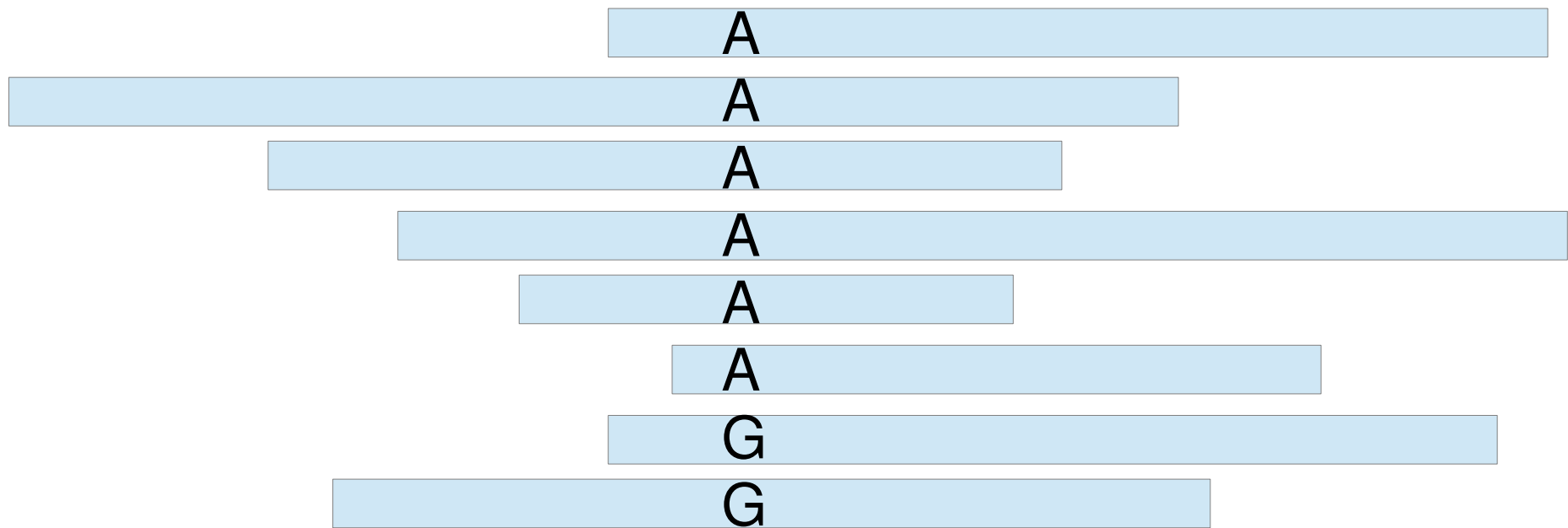
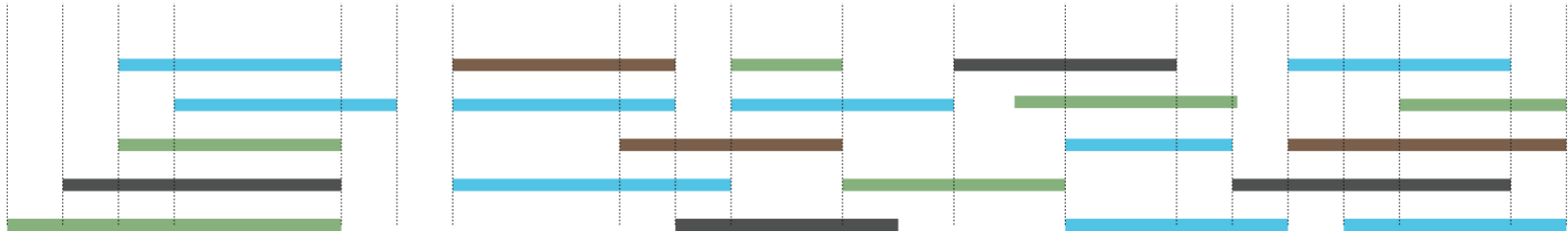


# Next Generation Sequencing (NGS)

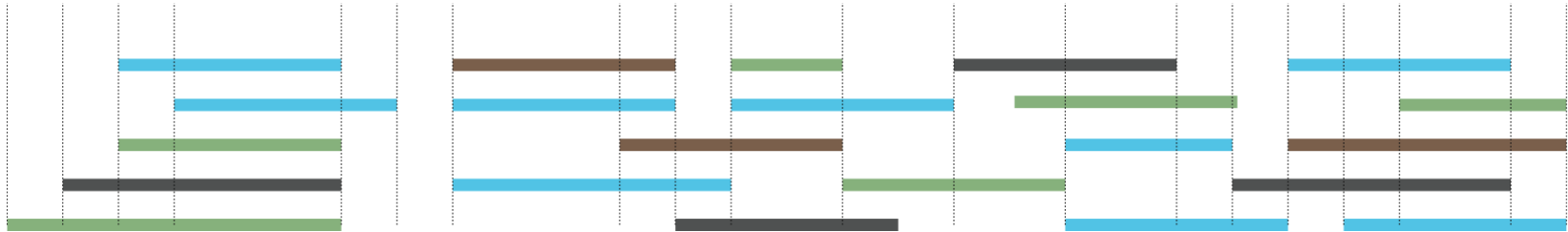


- However NGS has drawbacks:
  - High error rates
  - Shorter reads

# Sequencing depth / coverage

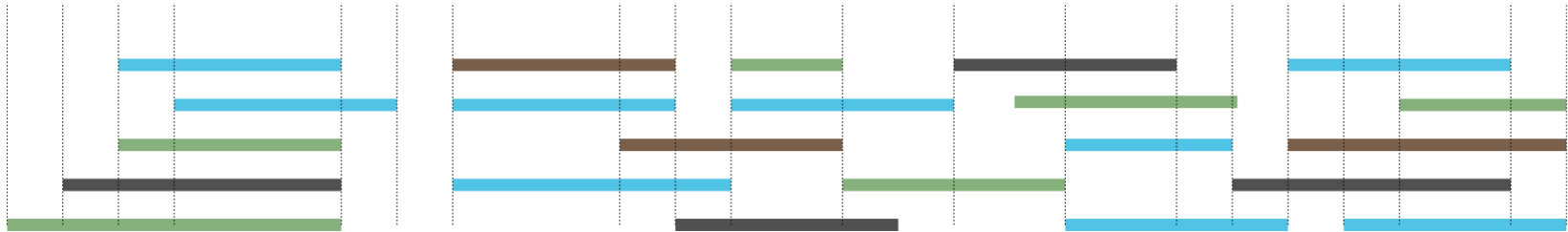


# Is the site variable in the sample?



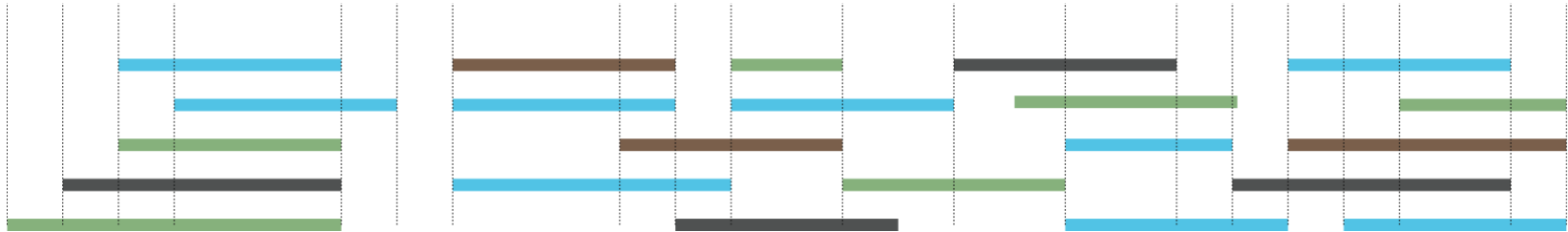
A  
A  
A  
A  
A  
A  
G  
G

Is the site still variable in the sample?



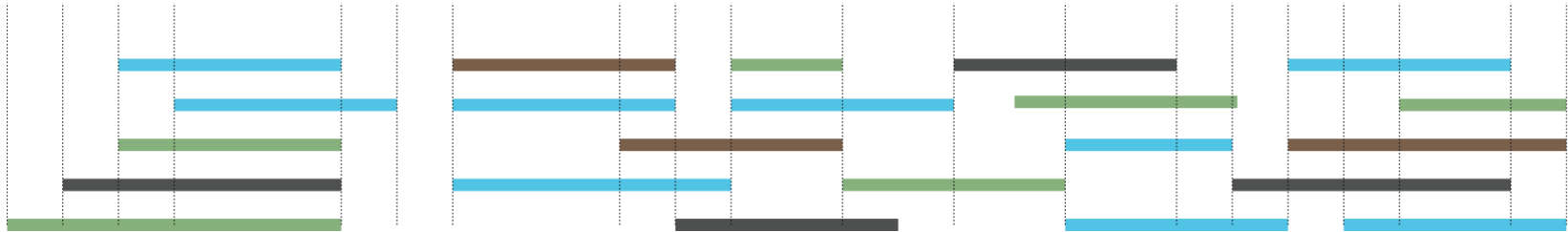
A	A
A	A
A	A
A	A
A	A
A	A
G	G
G	

Is the site still variable in the sample/individual?



A	A	A
A	A	A
A	A	A
A	A	A
A	A	A
A	A	A
G	G	
G		

# Is the site still variable in the sample/individual?



A	A	A
A	A	A
A	A	A
A	A	A
A	A	A
A	A	A
G	G	
G		

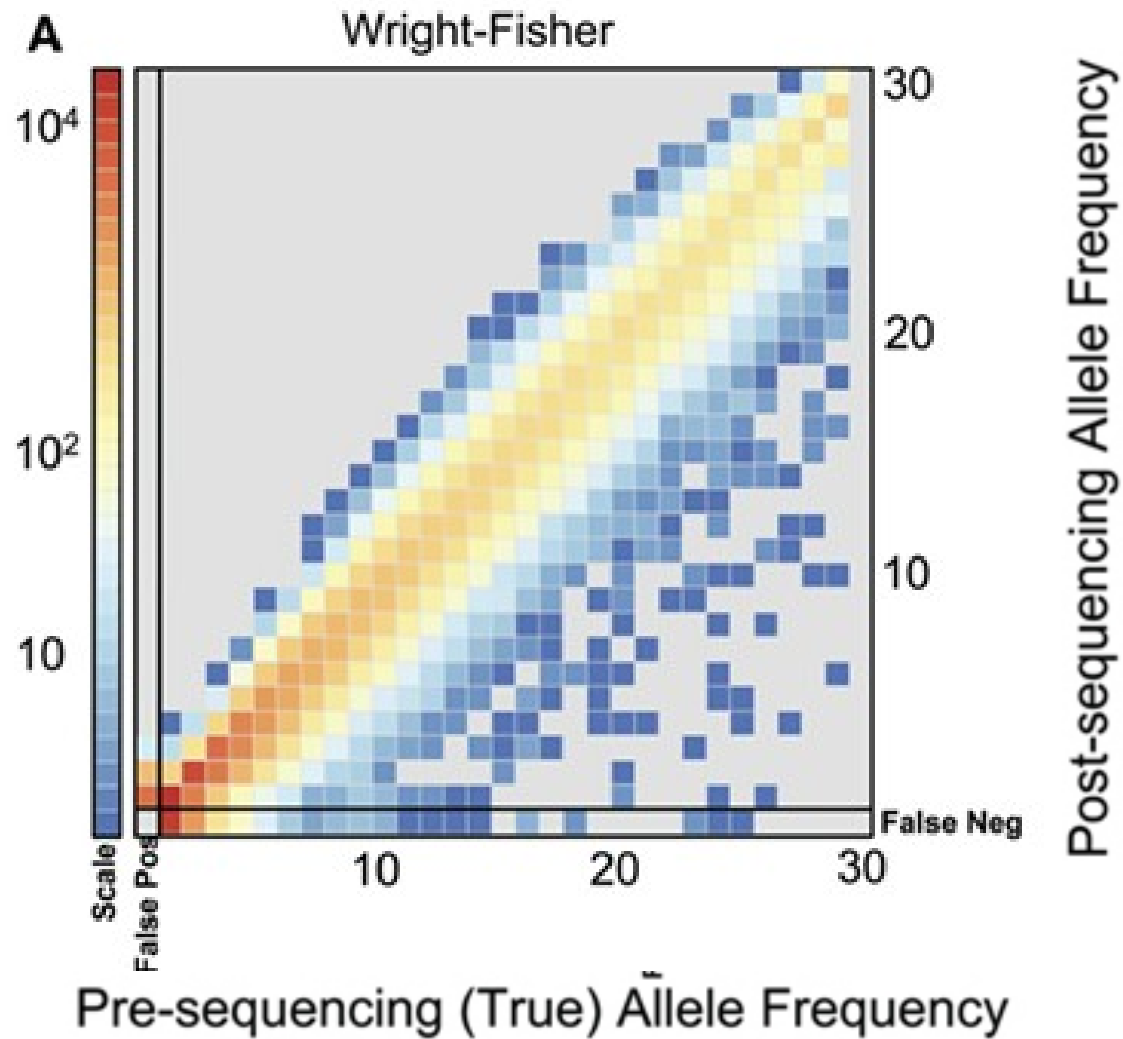
Common errors introduced here:

**SNP calling:** identification of variable sites.

**Genotype calling:** determination of the genotype for each site for each individual.



# Bias in allele frequencies



Crawford and Lazzaro 2012

# Possible solutions



# Possible solutions



More sequencing depth?

More samples?

# It depends...

- Fixed budget
  - Balance between sample size and coverage (uncertainty)
  - Depends on objective
    - Reference genome (high coverage)
    - Rare variants (large sample sizes at high coverage)
    - Population genetics (large sample sizes)
  - How low can we go?
- How to deal with uncertainty?
  - Stricter filtering → Loss of data
  - Probabilistic framework (genotype likelihoods)
    - Increased analytical power
    - Associated measure of statistical uncertainty
    - Incorporation of **prior** information

# Objective

- 1) What are **genotype likelihoods** (GL)?
- 2) How to do **SNP calling** from GL?
- 3) How to do **genotype calling** from GL?
- 4) What is the **error** in population genetic inferences using naïve strategies for **SNP and genotype calling**?
- 5) What is the optimal **sequencing design** for population genetics purposes?

# Objective

1) What are **genotype likelihoods (GL)**?

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# Genotype likelihoods

Probability of observing the read data, given a particular **genotype**

$$p(X|G=bh) = \frac{1}{2^r} \prod_{i=1}^r (L_b^{(i)} + L_h^{(i)})$$



Likelihood of observing allele  $b$  at read  $i$

# Genotype likelihoods – an example

Where can we get the error rate from?

-A-  
-A-  
-C-  
-T-

$$P(X|AA) = \left(\frac{L_A^{(1)}}{2} + \frac{L_A^{(1)}}{2}\right) * \left(\frac{L_A^{(2)}}{2} + \frac{L_A^{(2)}}{2}\right) * \left(\frac{L_A^{(3)}}{2} + \frac{L_A^{(3)}}{2}\right) * \left(\frac{L_A^{(4)}}{2} + \frac{L_A^{(4)}}{2}\right)$$

$$L_A^{(1)} = L_A^{(2)} = 1 - \epsilon \quad L_A^{(3)} = L_A^{(4)} = \frac{\epsilon}{3} \quad (1 - \epsilon) + \left(\frac{\epsilon}{3}\right) + \left(\frac{\epsilon}{3}\right) + \left(\frac{\epsilon}{3}\right) = 1$$

$$P(X|AC) = \left(\frac{L_A^{(1)}}{2} + \frac{L_C^{(1)}}{2}\right) * \left(\frac{L_A^{(2)}}{2} + \frac{L_C^{(2)}}{2}\right) * \left(\frac{L_A^{(3)}}{2} + \frac{L_C^{(3)}}{2}\right) * \left(\frac{L_A^{(4)}}{2} + \frac{L_C^{(4)}}{2}\right)$$

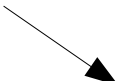
$$L_A^{(1)} = L_A^{(2)} = L_C^{(3)} = 1 - \epsilon \quad L_C^{(1)} = L_C^{(2)} = L_A^{(3)} = L_A^{(4)} = L_C^{(4)} = \frac{\epsilon}{3}$$



# Posterior probabilities of genotypes

**Prior** is derived assuming **HWE** from the estimated Minor Allele Frequency.

Genotype likelihood


$$P(G_s^{(i)} | X_s^{(i)}) = \frac{P(X_s^{(i)} | G_s^{(i)}) P(G_s^{(i)})}{\sum_{G=0}^2 P(X_s^{(i)} | G_s^{(i)}) P(G_s^{(i)})}$$

The term  $P(G_s^{(i)})$  in the numerator is circled in red.

$$P(A | B) = \frac{P(B | A)P(A)}{P(B)}$$

Nielsen et al 2012

- Model organisms

- Reference genome
- SNP databases
- Patterns of LD
- Known allele or genotype frequencies
- ...

- Non-model organisms

- Expected genotype frequencies under a model (e.g. HWE)
  - Works for most case, if population follows HWE
  - Exceptions:
    - Inbreeding (e.g. self-pollination plants)
    - Asexual reproduction

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2) How to do **SNP calling** from GL?

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5) What is the optimal sequencing design for population genetics purposes?

# Objective

Sample	True genotype	Reads allele A	Read allele G
1	AA	7	0
2	AA	25	1
3	AG	5	3
4	AG	4	4
5	GG	0	2
6	GG	0	4
Total		41	14

**What is the true frequency?**

**What is the estimated frequency?**

**What is the problem with that estimate?**

# Estimating Allele Frequencies - ML

$$P(D|f) = \prod_{i=1}^N \sum_{g \in \{0,1,2\}} P(D|G = g)P(G = g|f)$$

- Likelihood function
- What is?
  - $P(D | G) = P(X | G)$
  - $P(G = g | f)$
- Estimate  $f$ , by optimizing the likelihood function through (e.g.) EM
  - $f = 0.46$

- ANGSD uses the minor allele frequency (MAF) to call SNPs
  - Naive:
    - $f > t$  (e.g.,  $t = 1/2N$ )
  - Likelihood Ratio Test (LRT), comparing the goodness of fit (chi2) between:
    - Null model:  $f = 0$
    - Alternative model:  $f \neq 0$

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## Calling genotypes – 10 GL

Genotype	Likelihood (log10)
AA	-2.49
AC	-3.38
AG	-1.22
AT	-3.38
CC	-9.91
CG	-7.74
CT	-9.91
GG	-7.44
GT	-7.74
TT	-9.91

What is the  
genotype?



## Calling genotypes – 3 GL

Genotype	Likelihood
AA	-5.73
AG	-2.80
GG	-17.12

**What is the genotype?**

# Calling genotypes – GL ratio

$$\log_{10} \frac{L_{G(1)}}{L_{G(2)}} > t$$

i.e.  $t = 1$  meaning that the most likely genotype is 10 times more likely than the second most likely one

**Pros and Cons?**

**Genotype Quality?**

**Missing data?**

# Calling genotypes – Posterior Probabilities (PP)

AAAG (A,G alleles)

$\varepsilon = 0.01$

Genotype	Likelihood (log)	Prior	Posterior
AA	-5.73	1/3	0.05
AG	-2.80	1/3	0.95
GG	-17.12	1/3	0

# Calling genotypes – PP (reference prior)

AAAG (A,G alleles)

$\varepsilon = 0.01$

A is reference  $\rightarrow P(AA) > P(AG) > P(GG)$

Genotype	Likelihood (log)	Prior	Posterior
AA	-5.73	0.80	0.22
AG	-2.80	0.15	0.78
GG	-17.12	0.05	0

# Calling genotypes – PP (HWE prior)

AAAG (A,G alleles)

$\varepsilon = 0.01$

$f(a) = 0.7$  (from a reference panel)

$P(AA) = ?$ ;  $P(AG) = ?$ ;  $P(GG) = ?$

Genotype	Likelihood (log)	Prior	Posterior
AA	-5.73	0.49	0.06
AG	-2.80	0.42	0.94
GG	-17.12	0.09	0

**Can we assume HWE?**

# Calling genotypes – PP (HWE prior)

AAAG (A,G alleles)

$\varepsilon = 0.01$

$f(a) = 0.7$  (from the data itself)

$P(AA) = ?$ ;  $P(AG) = ?$ ;  $P(GG) = ?$

Genotype	Likelihood (log)	Prior	Posterior
AA	-5.73	0.49	0.06
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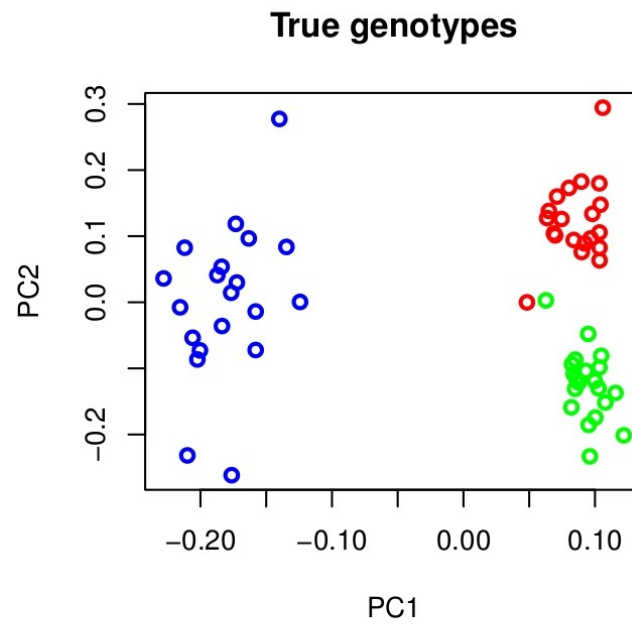
**Can we estimate  
frequencies  
accurately?**

**Can we assume HWE?**

# Objective

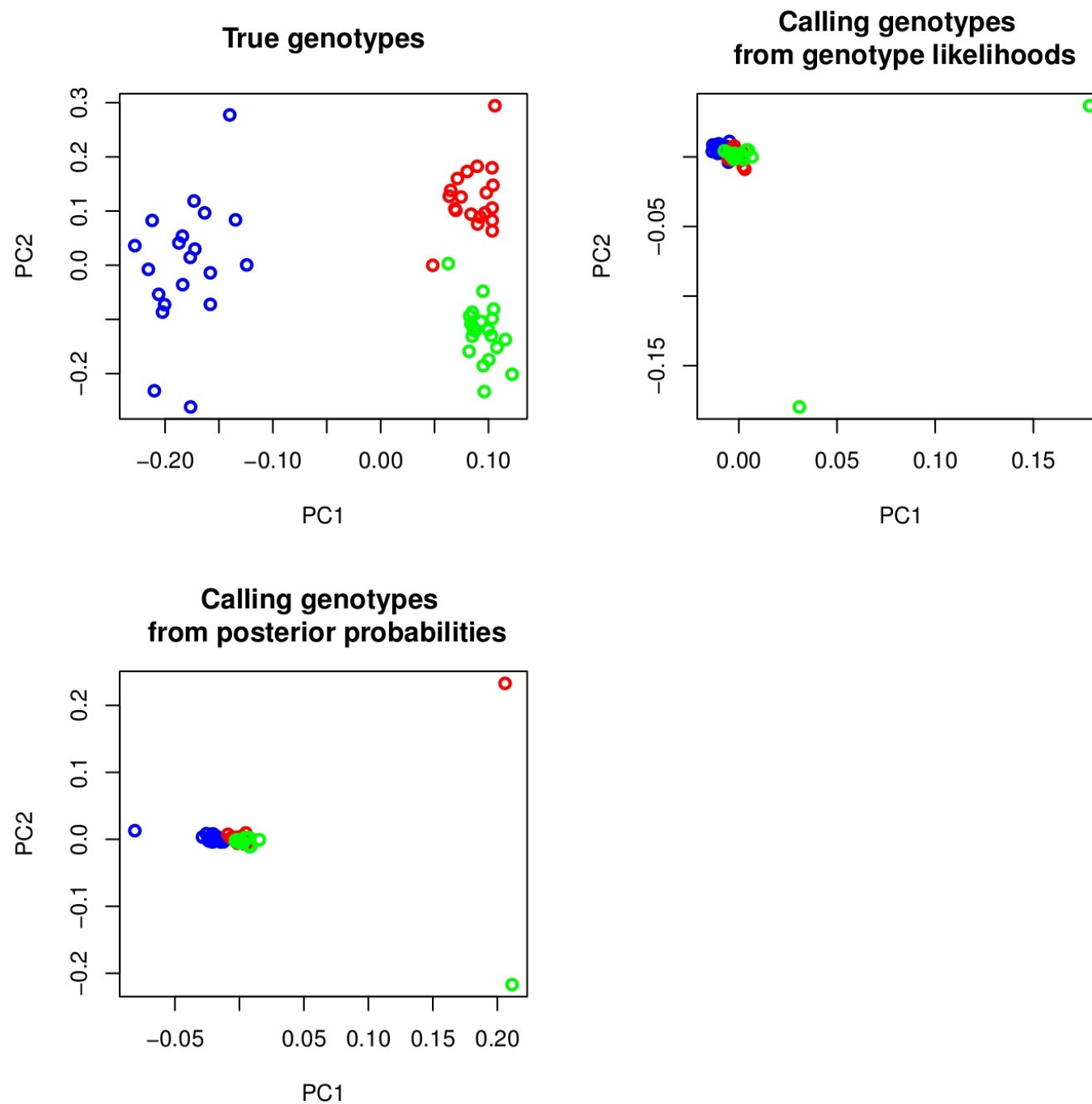
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# Population structure - PCA

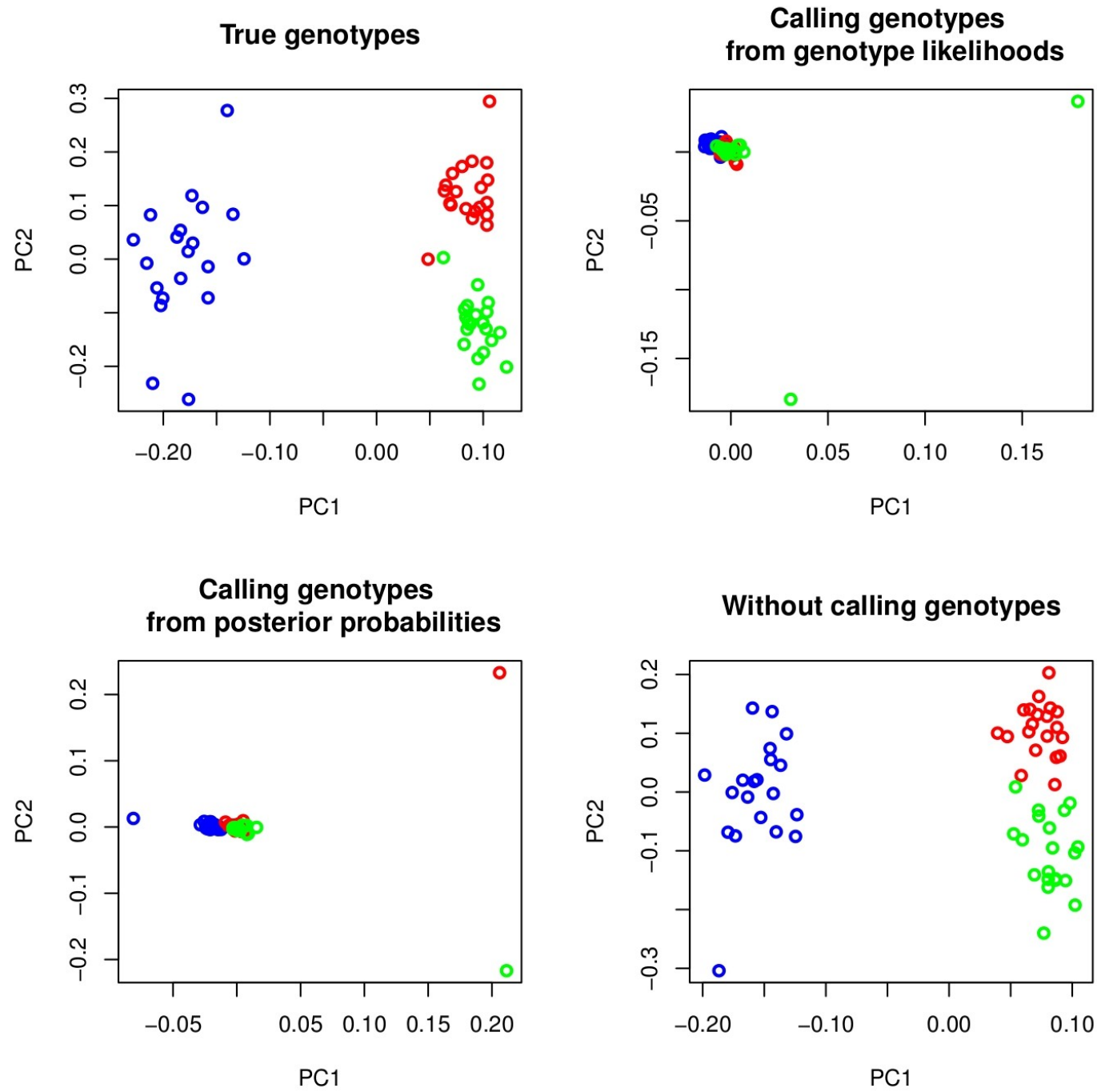




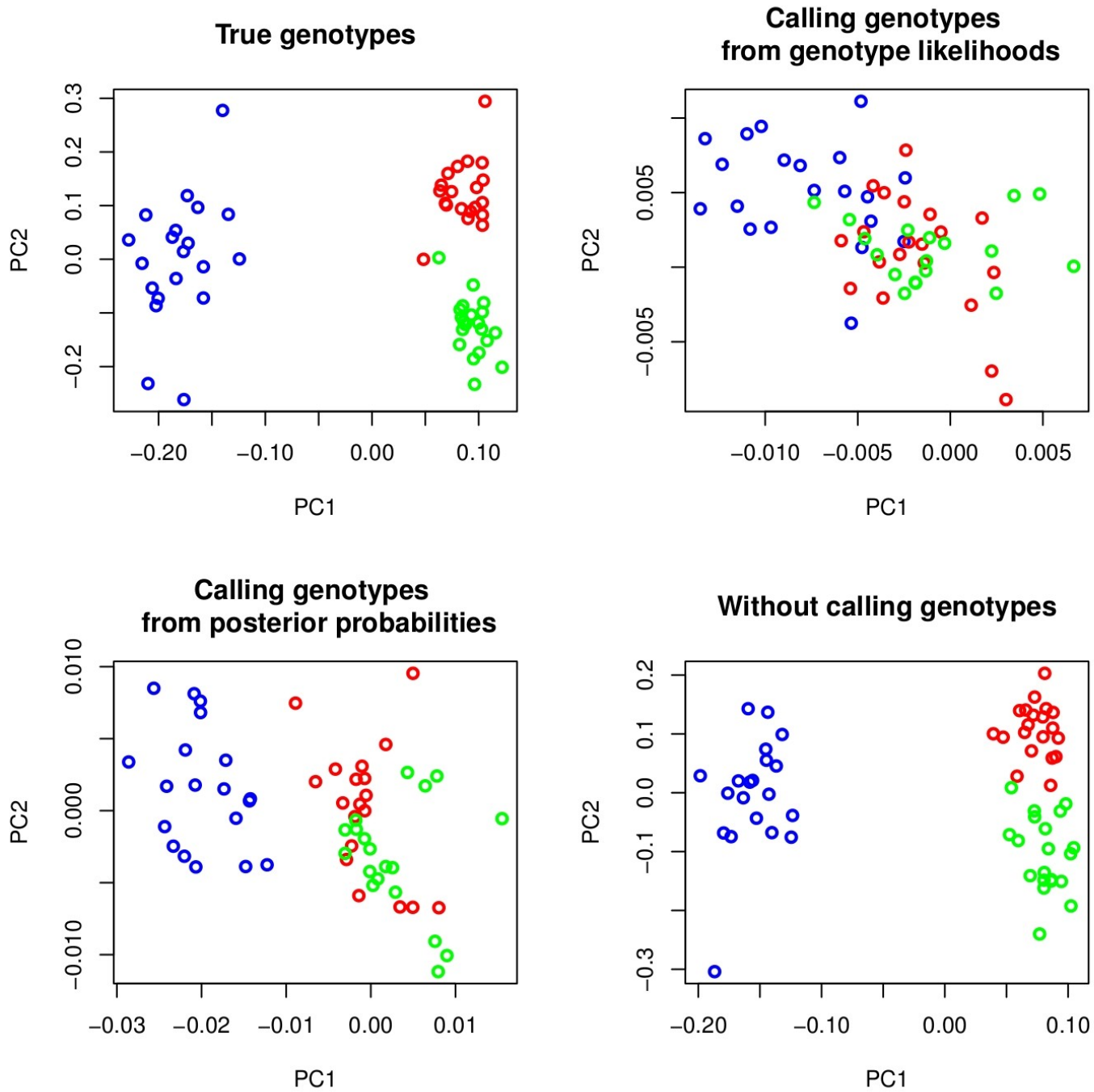
# Population structure - PCA



# Population structure - PCA



# Population structure – PCA (no outliers)



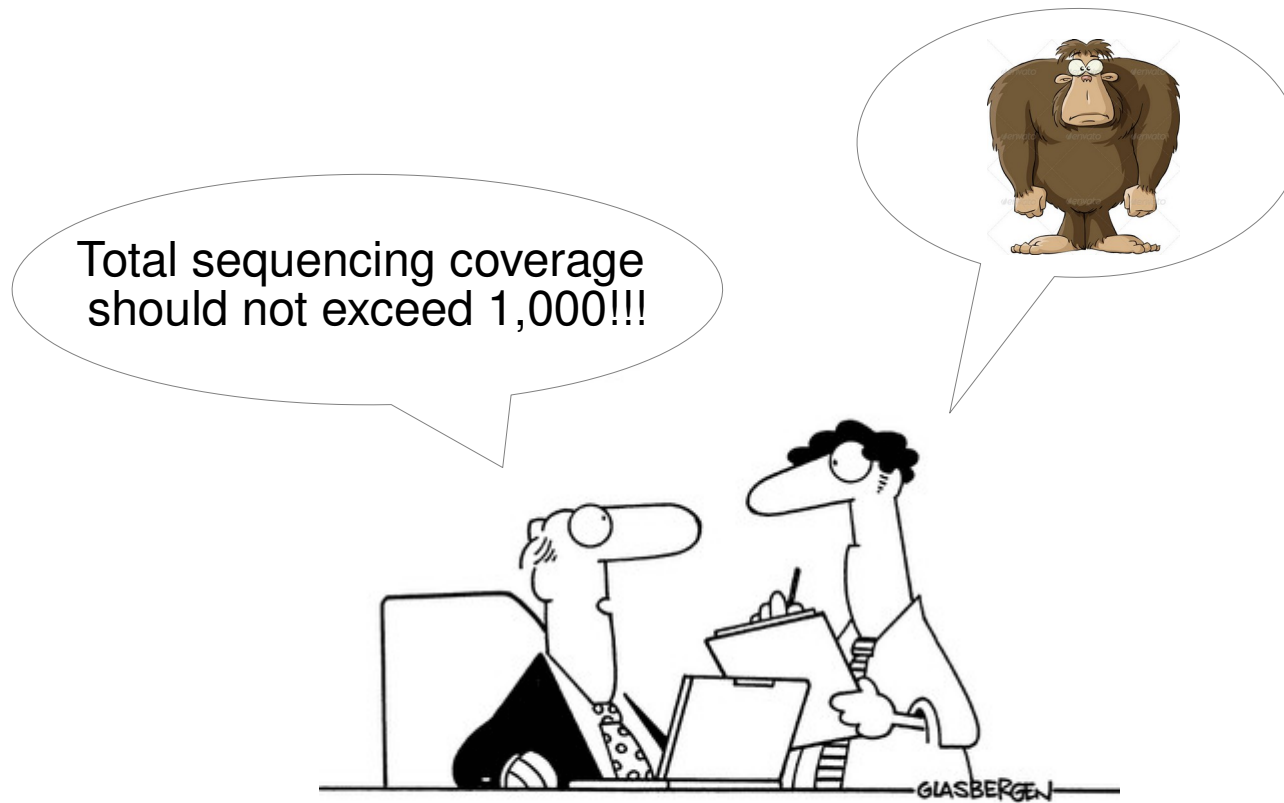
# Objective

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# Discovery of a “new” species/population

Population is comprised of **1,000 individuals**.

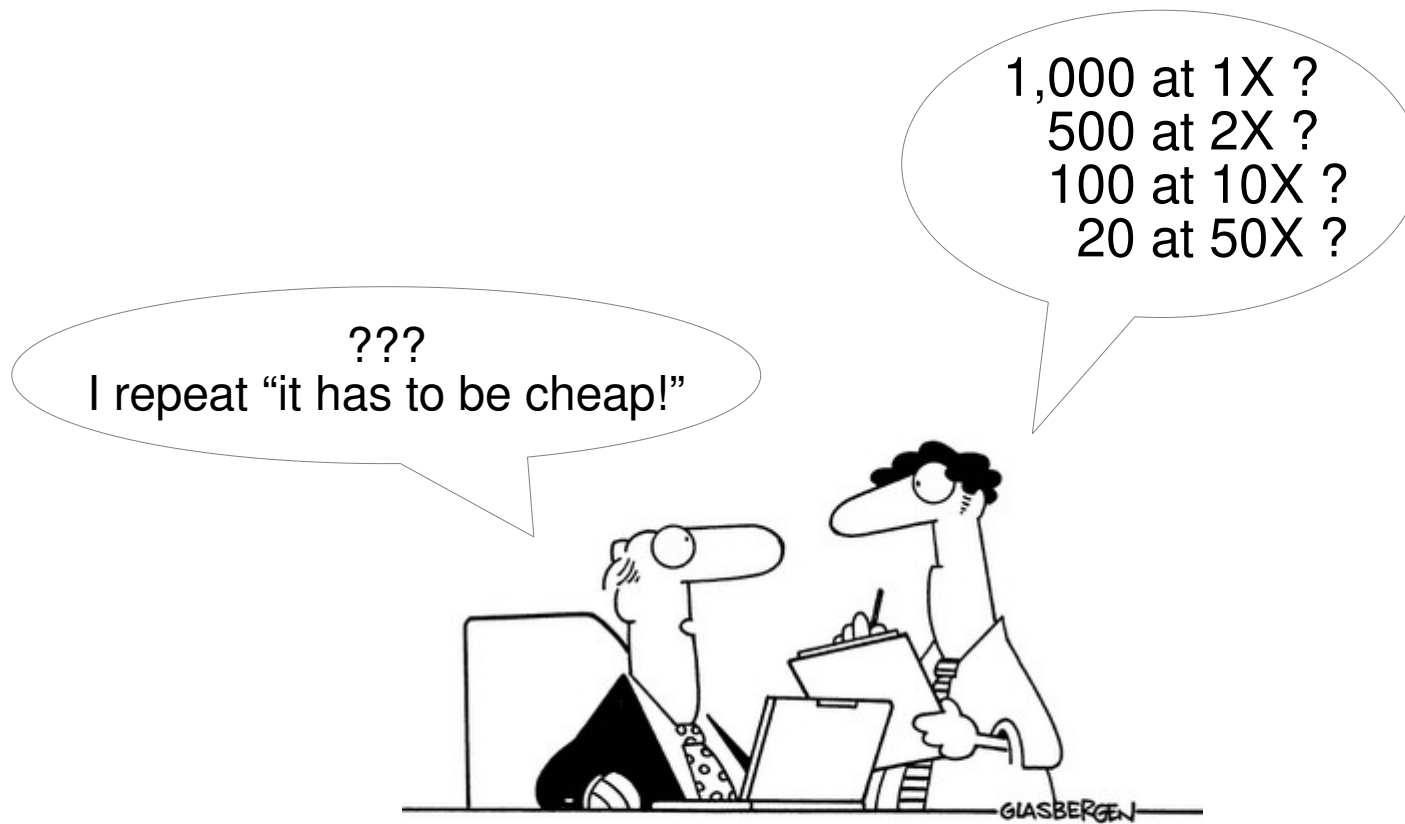
Genome is **100,000 bp** long.



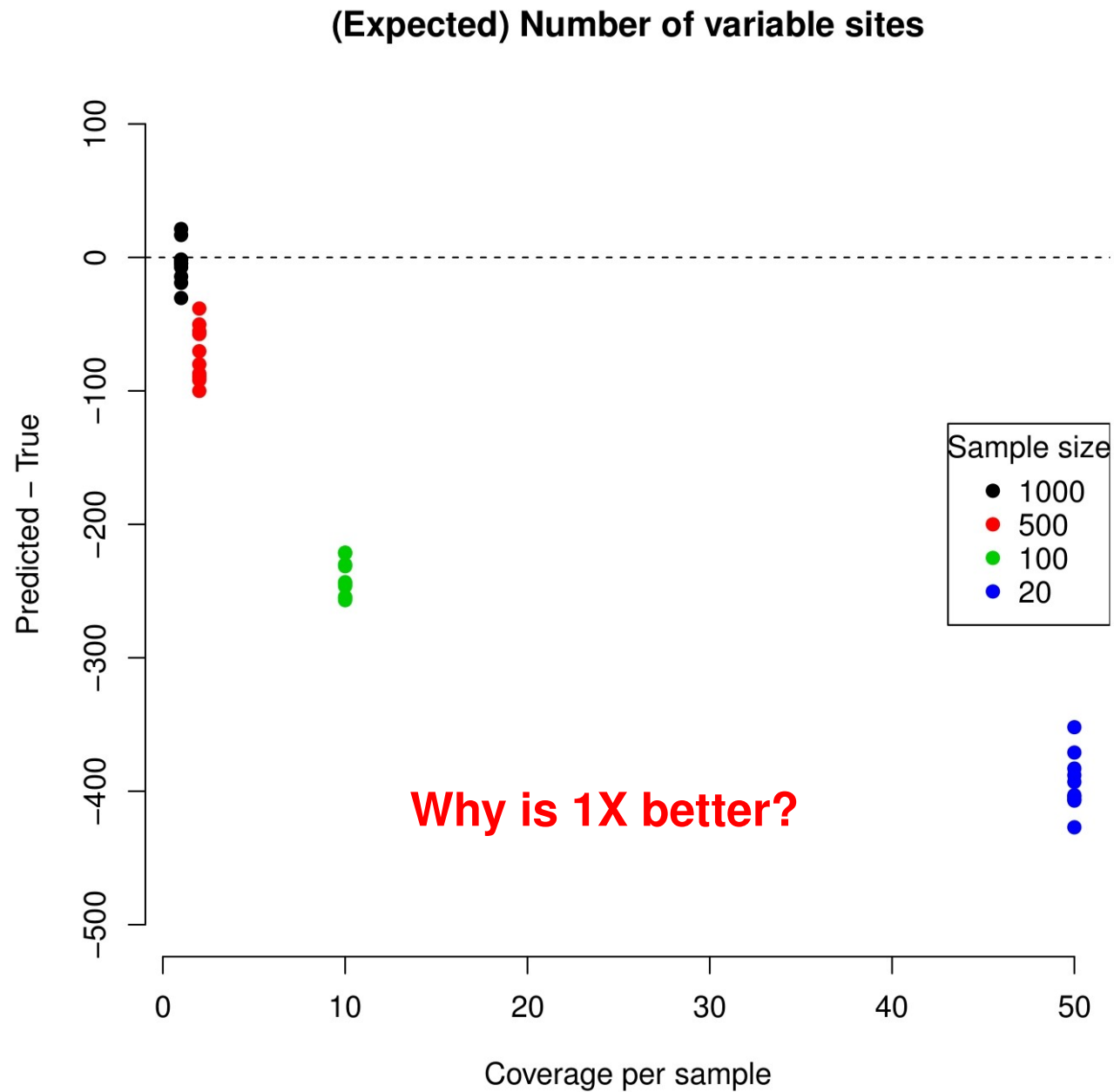
# Planning the experiment

Population is comprised of **1,000 individuals**.

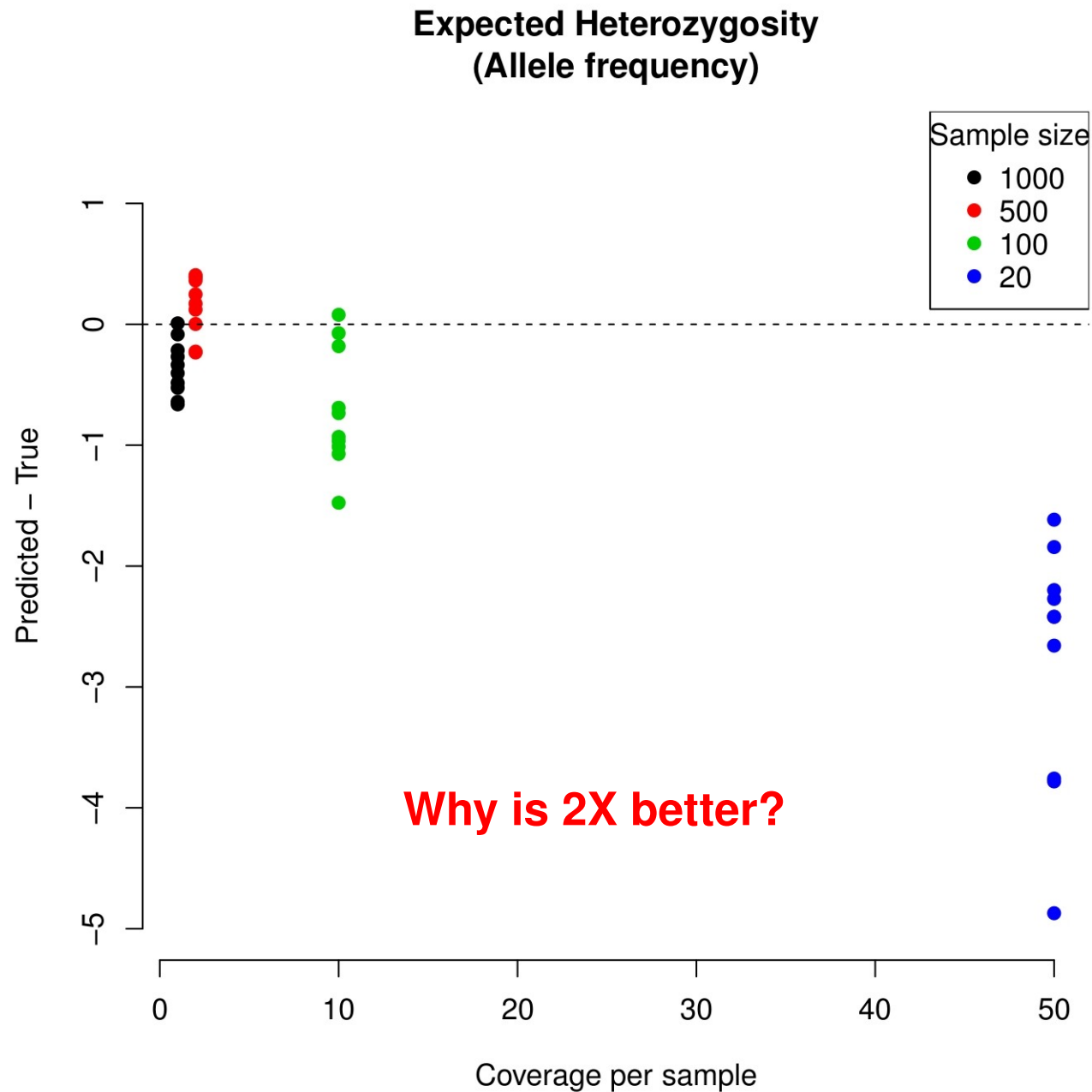
Genome is **100,000 bp** long.



# How many polymorphic sites?



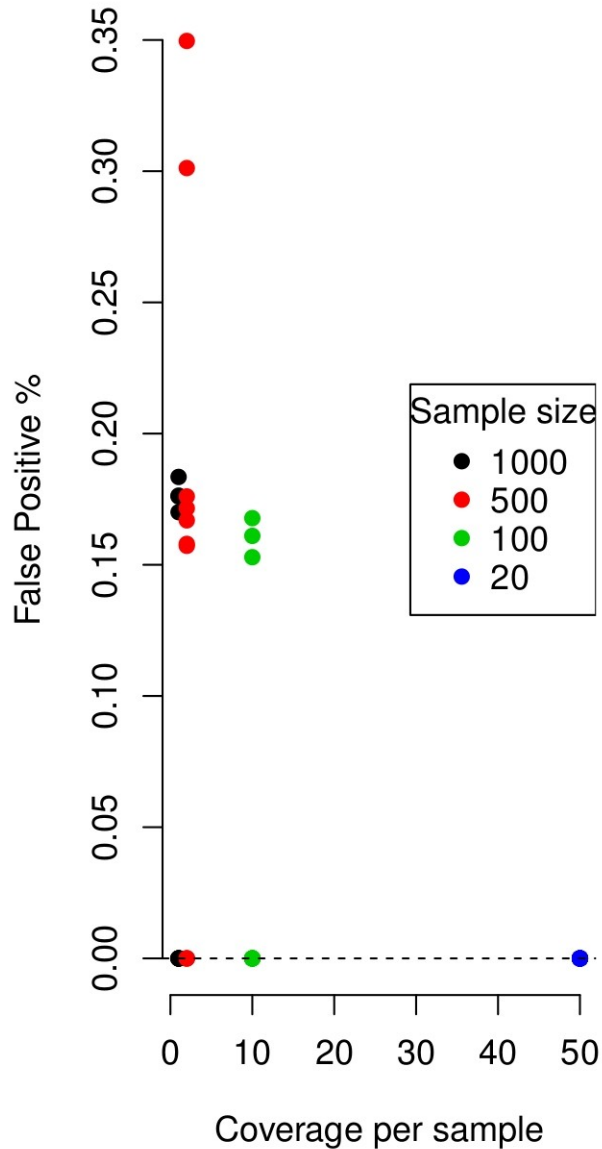
# How about the allele frequencies?





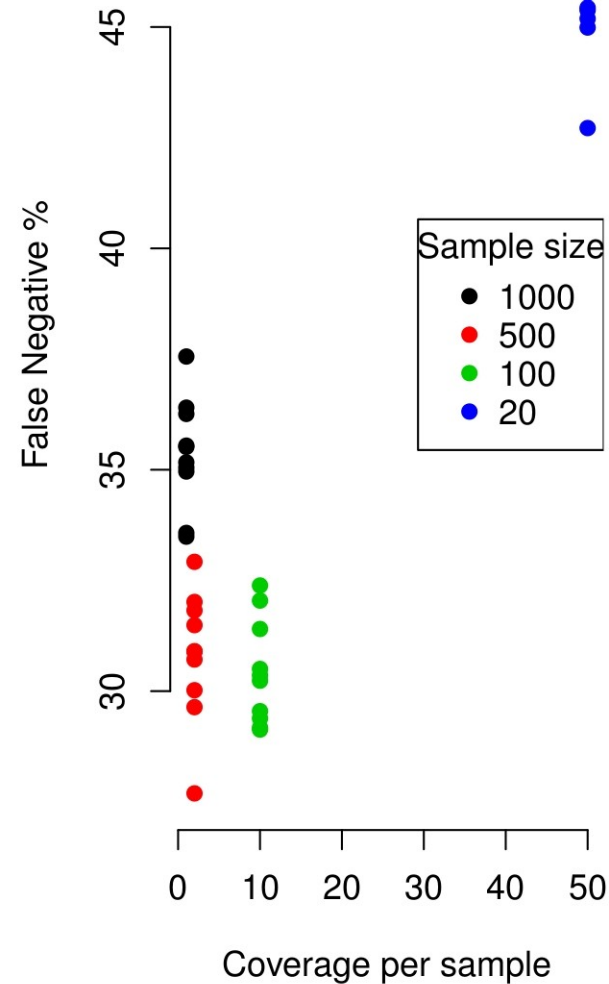
# Do you get the right SNPs?

SNP Calling



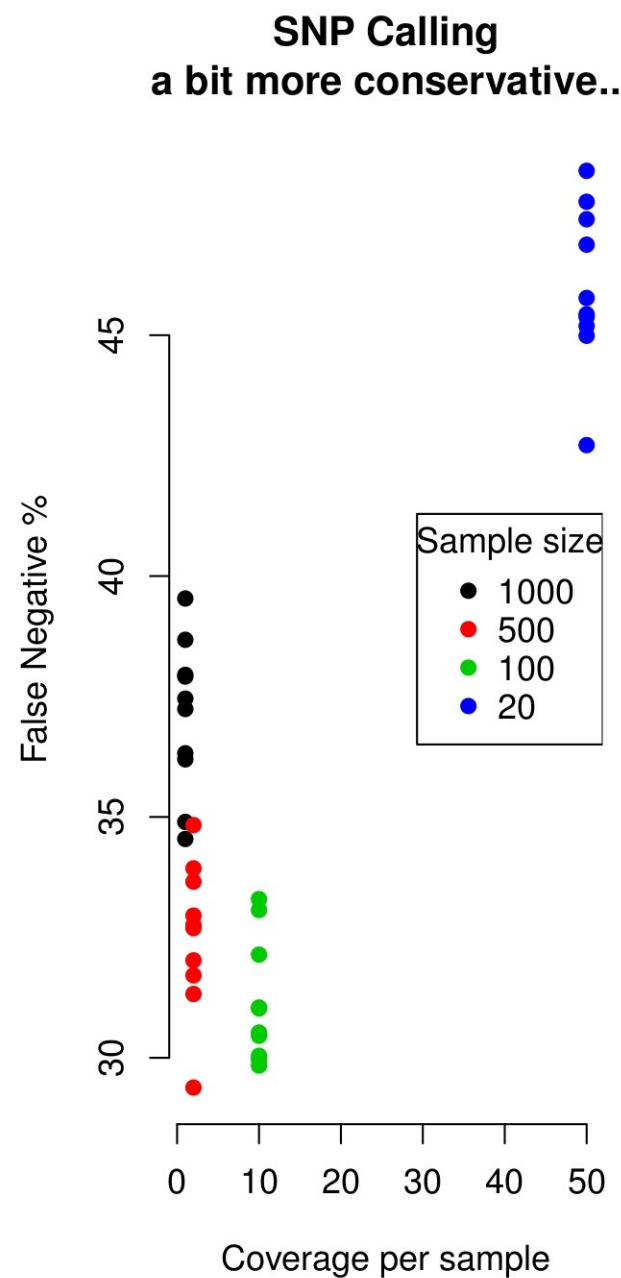
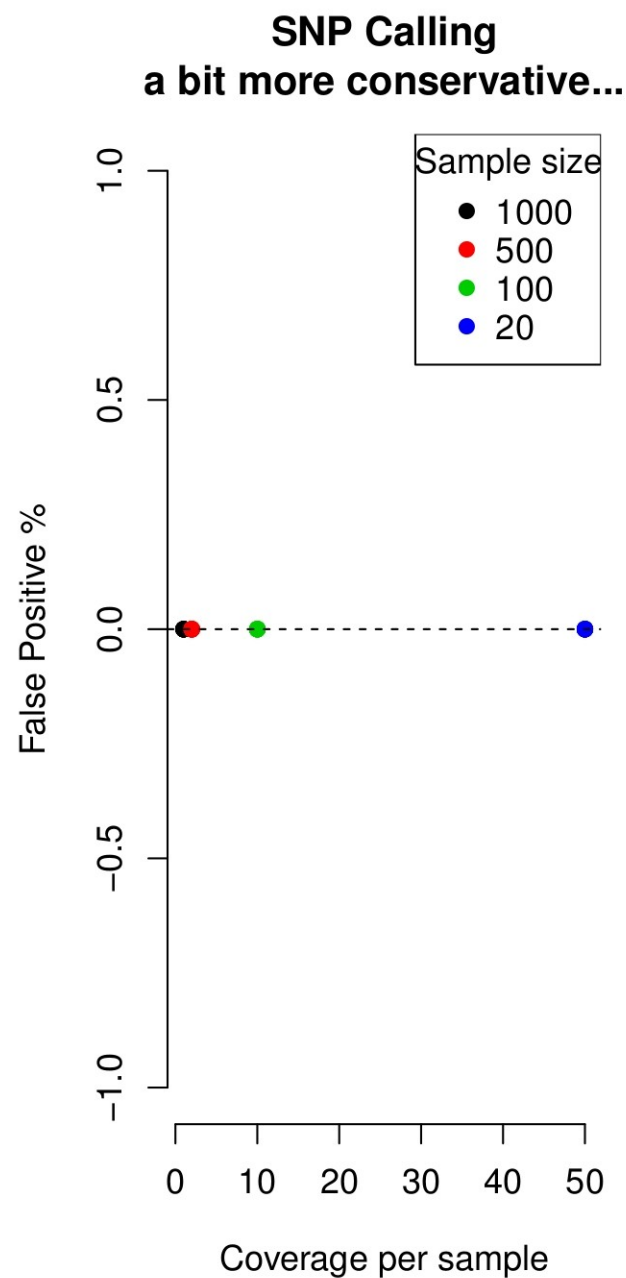
Why is FP rate high at low coverage?

SNP Calling

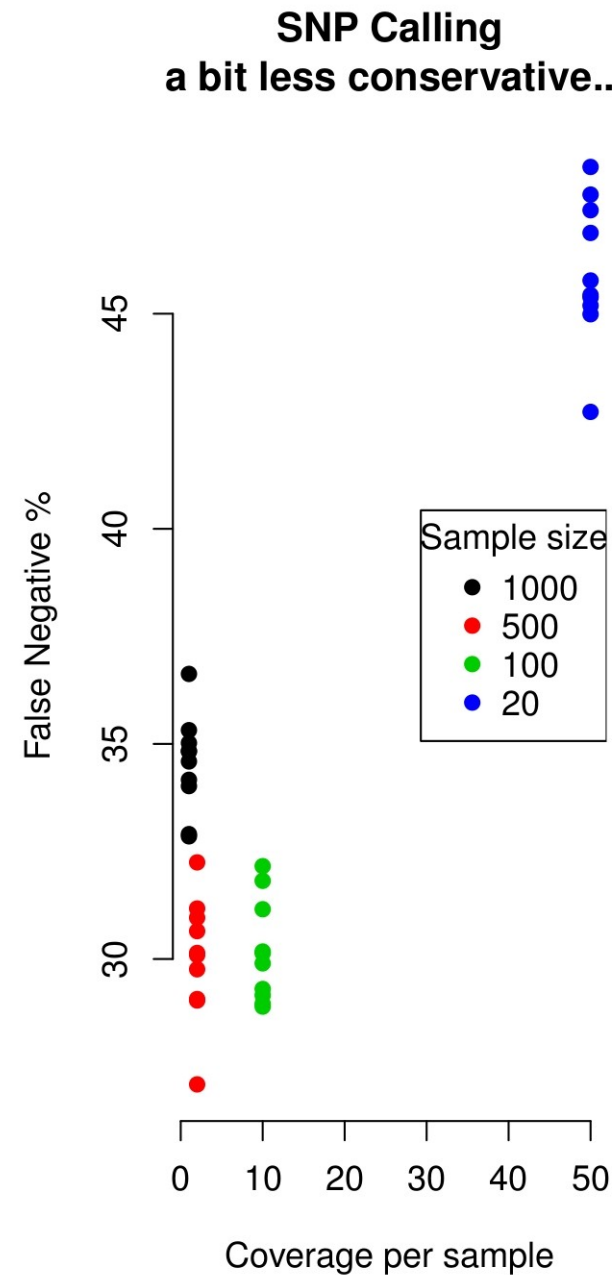
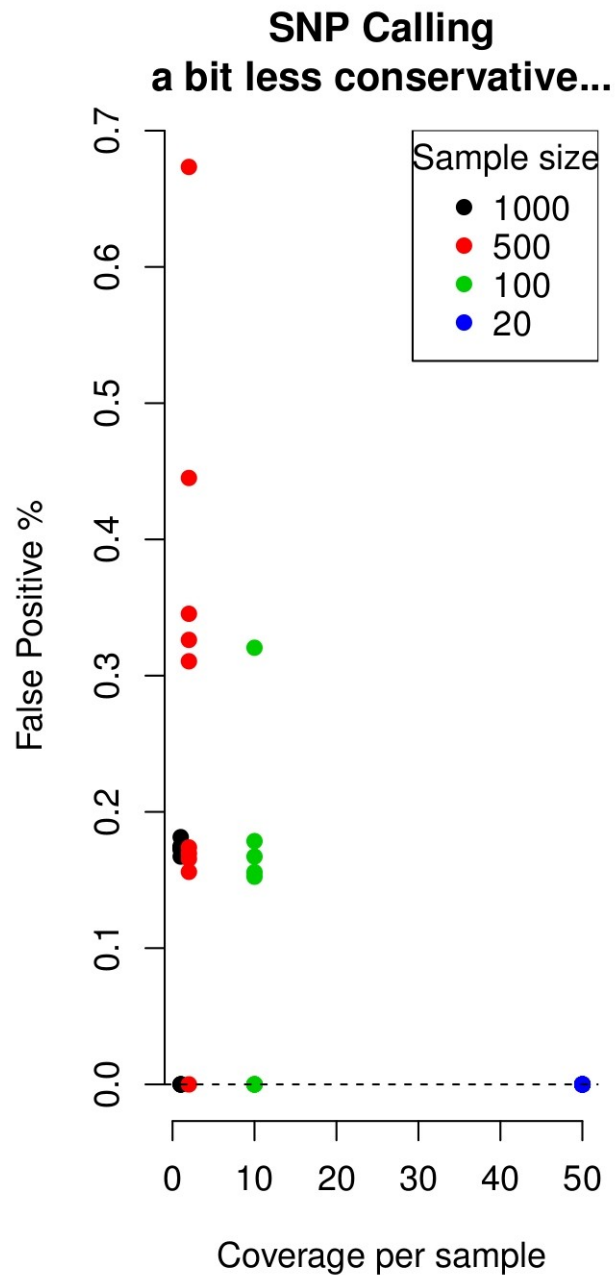


Why is FN rate high at high coverage?

# Do you get the right SNPs (more strict)?



# Do you get the right SNPs (less strict)?



# Conclusions

It is important to take **statistical uncertainty** into account, specially for low coverage samples.

The methods presented provide **tools** for investigating population genetic variation for multiple populations on a large scale.

The great improvement in accuracy for low coverage data can be explained by the fact that we **do not call SNPs or genotypes**.

# Acknowledgments



Rasmus Nielsen



Thorfinn Korneliussen  
Anders Albrechtsen



Matteo Fumagalli

Software available at:

<http://popgen.dk/software/angsd.html>

<https://github.com/fgvieira>

<https://github.com/mfumagalli/ngsTools>

# Performance of PCA

