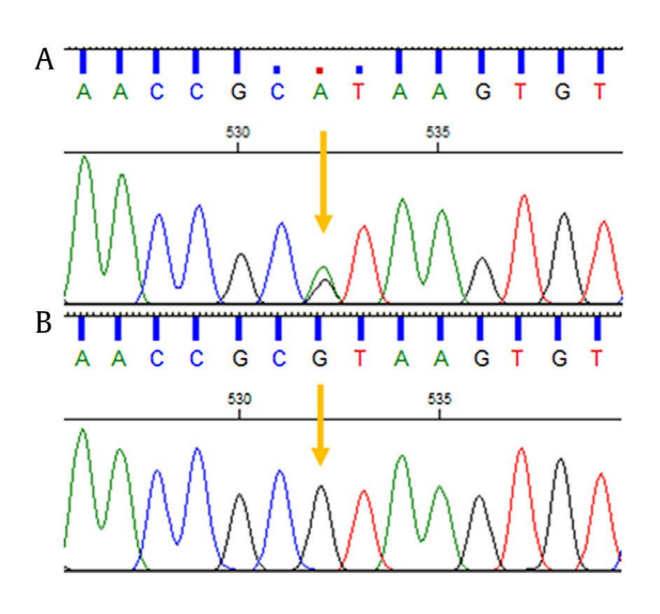
Calling variants from low-coverage NGS data

Filipe G. Vieira
Center for Genomic Medicine
Copenhagen University Hospital, Rigshospitalet filipe.garrett.vieira@regionh.dk

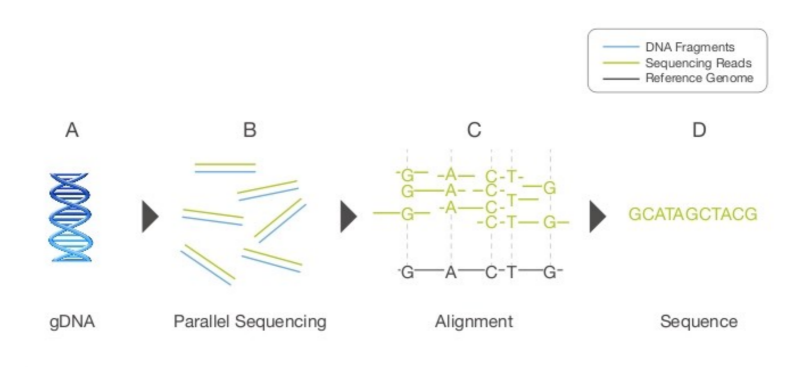




Sanger Sequencing



Next Generation Sequencing (NGS)



A. Extracted gDNA

www.illumina.com

However NGS has drawbacks:

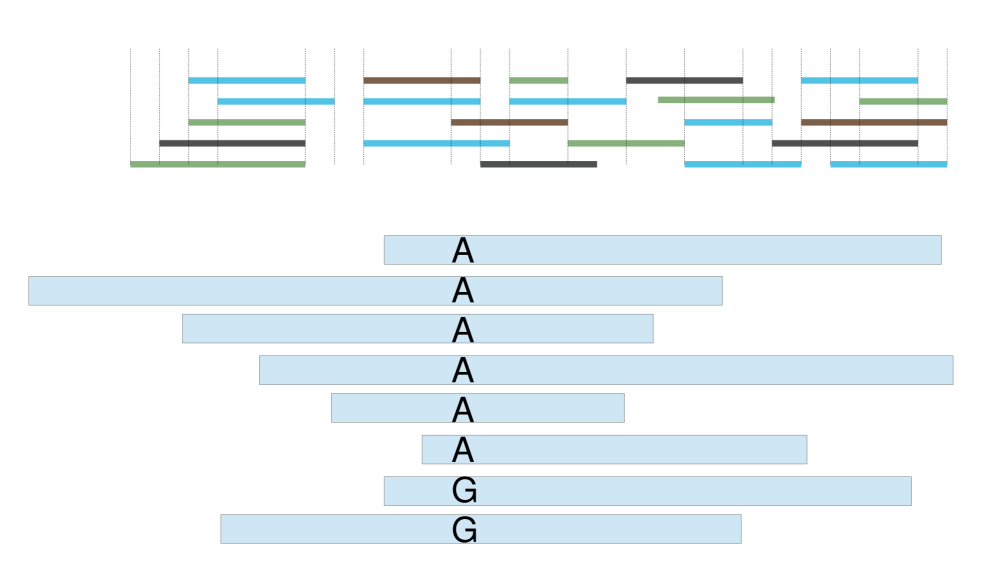
- High error rates
- Shorter reads

B. gDNA is fragmented into a library of small segments that are each sequenced in parallel.

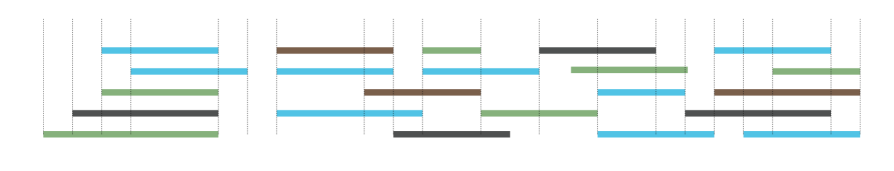
C. Individual sequence reads are reassembled by aligning to a reference genome

D. The whole-genome sequence is derived from the consensus of aligned reads.

Sequencing depth / coverage

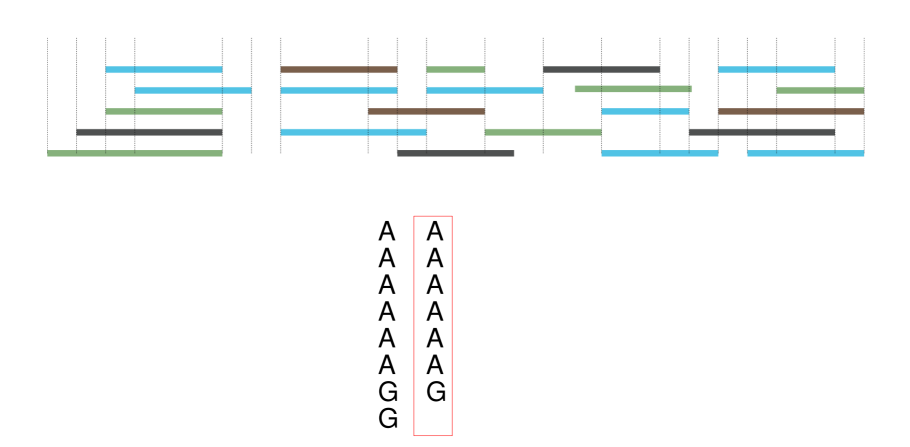


Is the site variable in the sample?

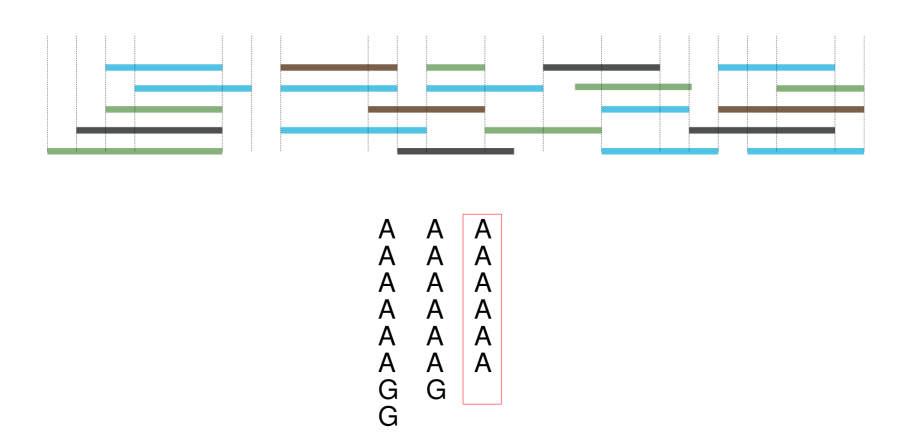


AAAAAGG

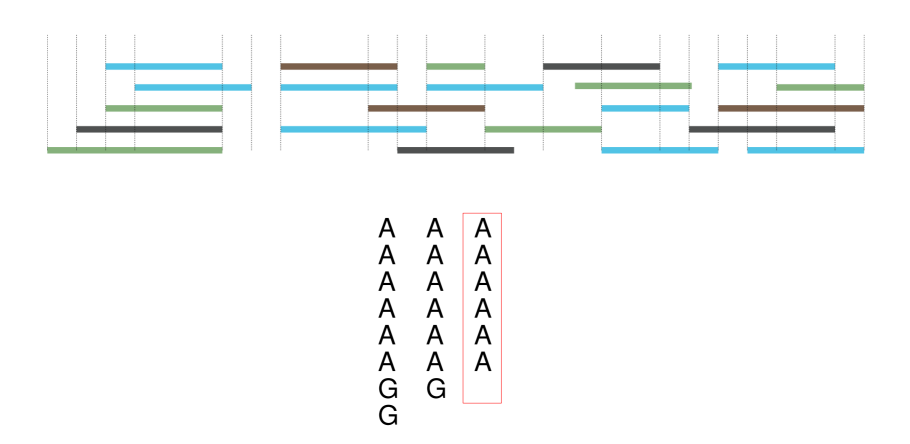
Is the site <u>still</u> variable in the sample?



Is the site still variable in the sample/individual?



Is the site still variable in the sample/individual?

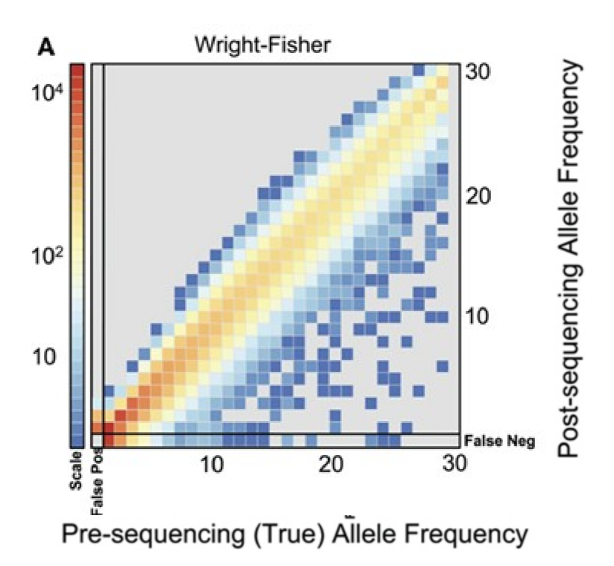


Common errors introduced here:

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 <u>sample size</u> and <u>coverage</u> (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?

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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?

$$\begin{split} P(X|AA) &= (\frac{L_A^{(1)}}{2} + \frac{L_A^{(1)}}{2}) * (\frac{L_A^{(2)}}{2} + \frac{L_A^{(2)}}{2}) * (\frac{L_A^{(3)}}{2} + \frac{L_A^{(3)}}{2}) * (\frac{L_A^{(4)}}{2} + \frac{L_A^{(4)}}{2}) \\ L_A^{(1)} &= L_A^{(2)} = 1 - \epsilon \qquad L_A^{(3)} = L_A^{(4)} = \frac{\epsilon}{3} \qquad (1 - \epsilon) + (\frac{\epsilon}{3}) + (\frac{\epsilon}{3}) + (\frac{\epsilon}{3}) + (\frac{\epsilon}{3}) = 1 \\ P(X|AC) &= (\frac{L_A^{(1)}}{2} + \frac{L_C^{(1)}}{2}) * (\frac{L_A^{(2)}}{2} + \frac{L_C^{(2)}}{2}) * (\frac{L_A^{(3)}}{2} + \frac{L_C^{(3)}}{2}) * (\frac{L_A^{(4)}}{2} + \frac{L_C^{(4)}}{2}) \\ L_A^{(1)} &= L_A^{(2)} = L_C^{(3)} = 1 - \epsilon \qquad L_C^{(1)} = L_C^{(2)} = L_A^{(3)} = L_A^{(4)} = L_C^{(4)} = \frac{\epsilon}{3} \end{split}$$

Posterior probabilities of genotypes

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

Genotype likelihood
$$P(G_{s}^{(i)}|X_{s}^{(i)}) = \frac{P(X_{s}^{(i)}|G_{s}^{(i)})P(G_{s}^{(i)})}{\sum_{G=0}^{2} P(X_{s}^{(i)}|G_{s}^{(i)})P(G_{s}^{(i)})}$$

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

Nielsen et al 2012

Priors

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-polinatign plans)
 - Asexual reproduction

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

$$- f = 0.46$$

Priors

- ANGSD uses the minor allele frequency (MAF) to call SNPs
 - Naive:
 - t > t (e.g., t = 1/2N)
 - Likelihood Ratio Test (LRT), comparing the goodness of fit (chi2) between:
 - Null model: *f* = 0
 - Alternative model: *f* <> 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	What is the genotype?
GT	-7.74	
TT	-9.91	

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 assume HWE?

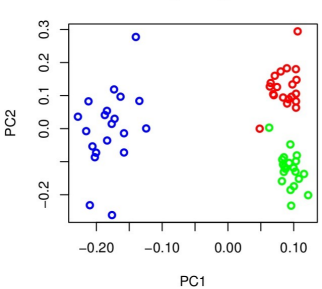
Can we estimate frequencies accurately?

Objective

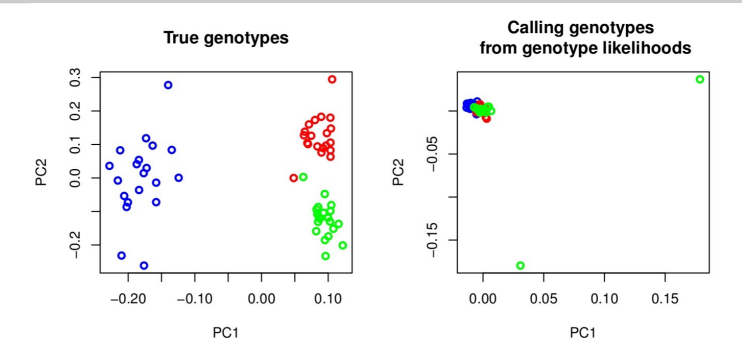
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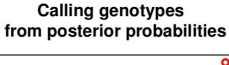
Population structure - PCA

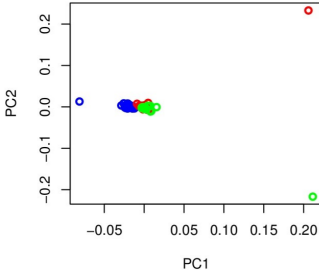




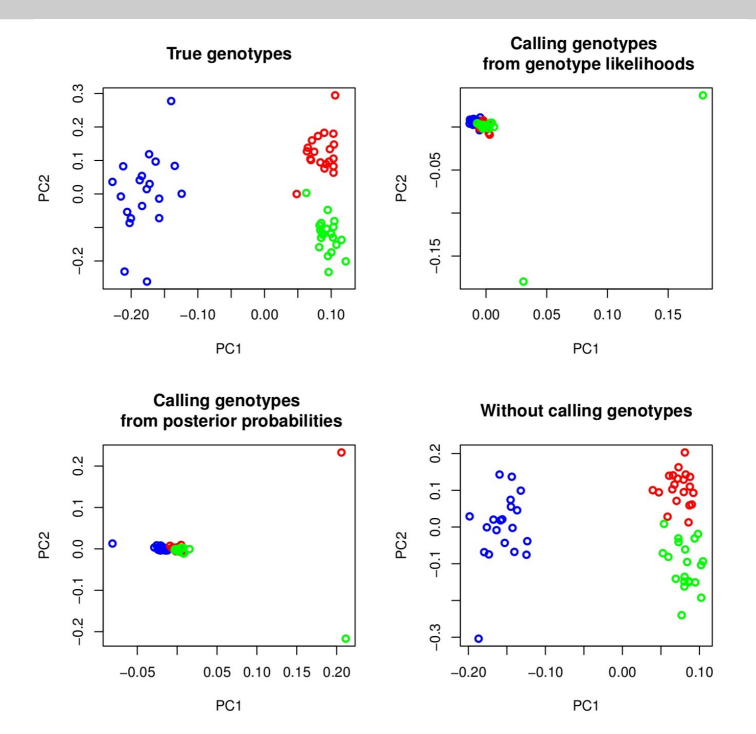
Population structure - PCA



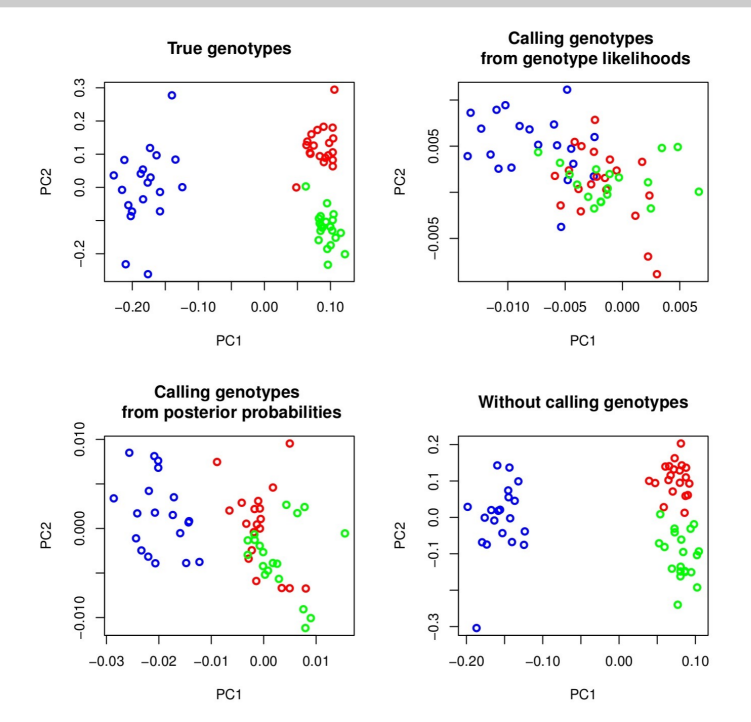




Population structure - PCA



Population structure – PCA (no outliers)



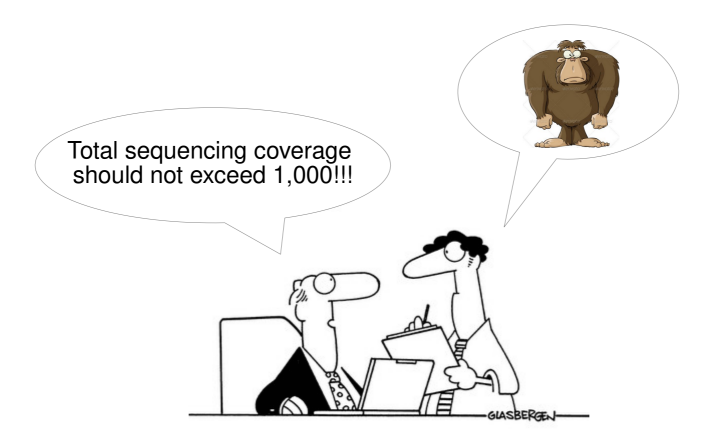
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Discovery of a "new" species/population

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

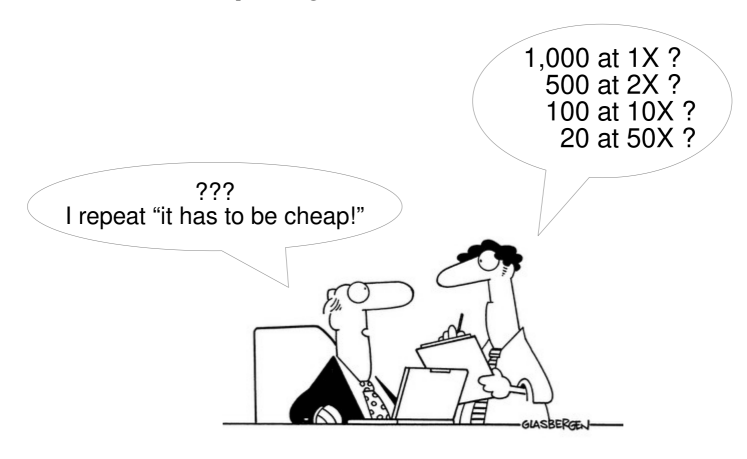
Genome is 100,000 bp long.



Planning the experiment

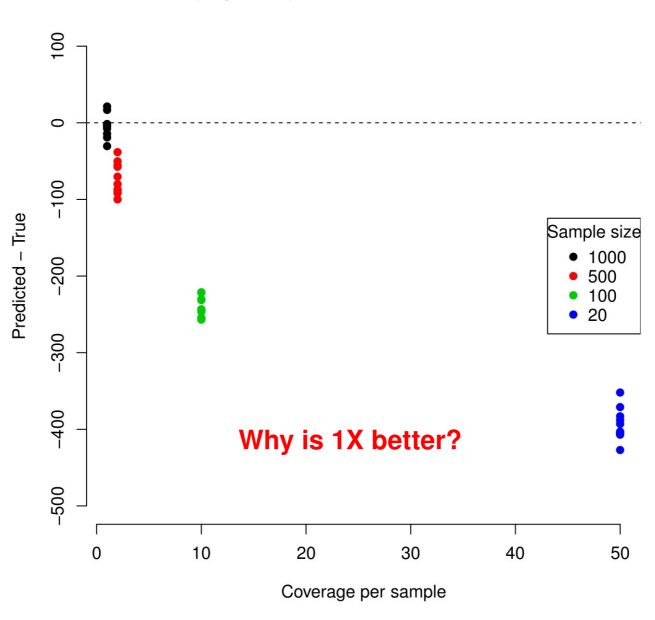
Population is comprised of **1,000 individuals**.

Genome is 100,000 bp long.

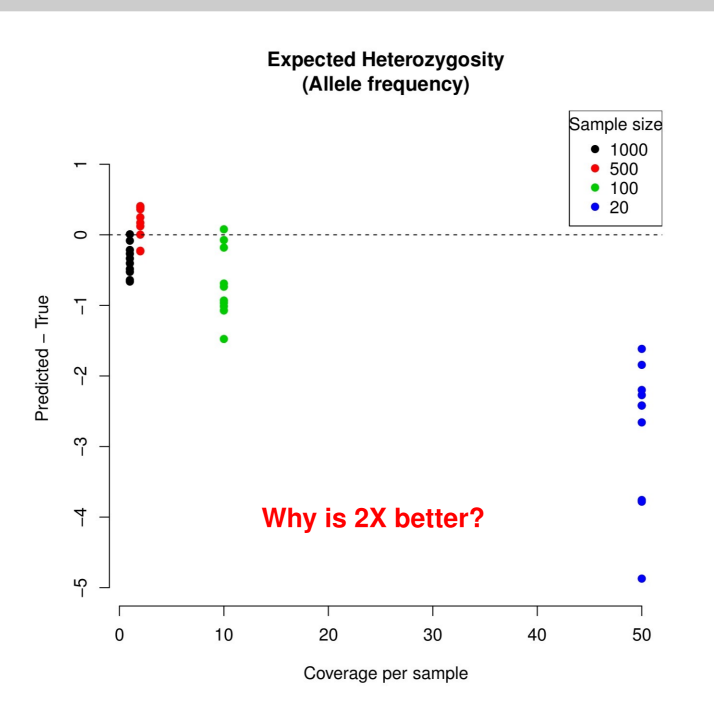


How many polymorphic sites?

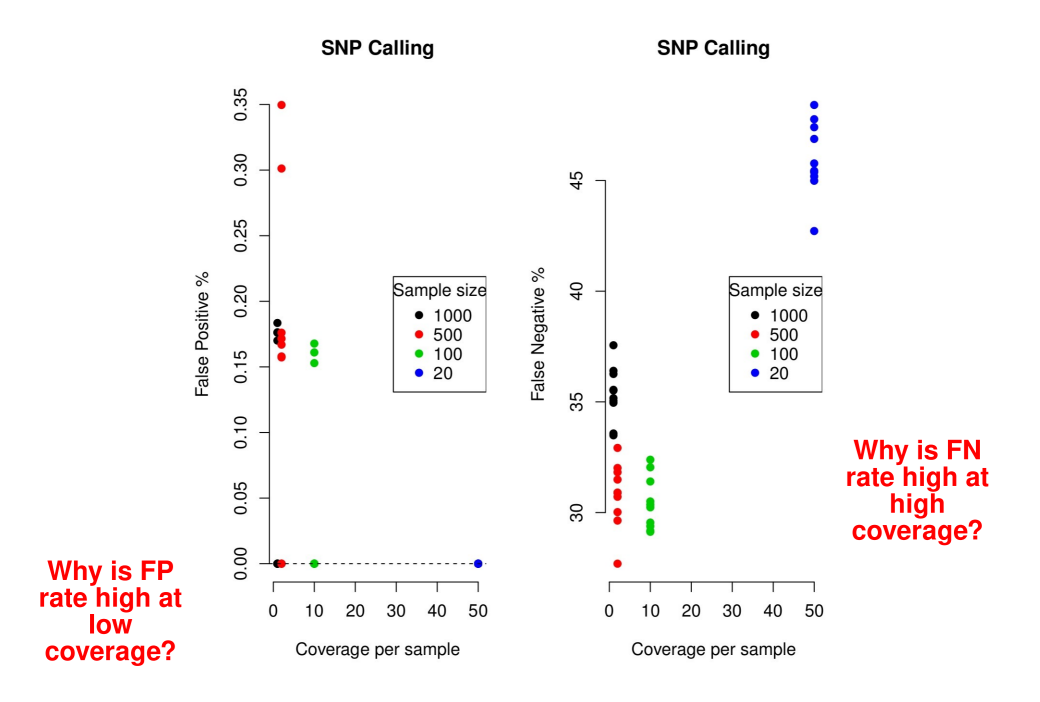




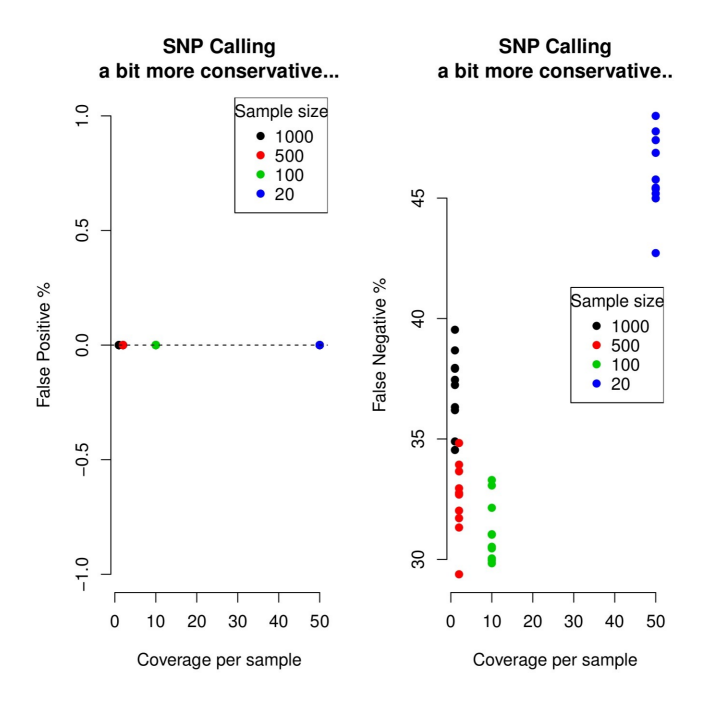
How about the allele frequencies?



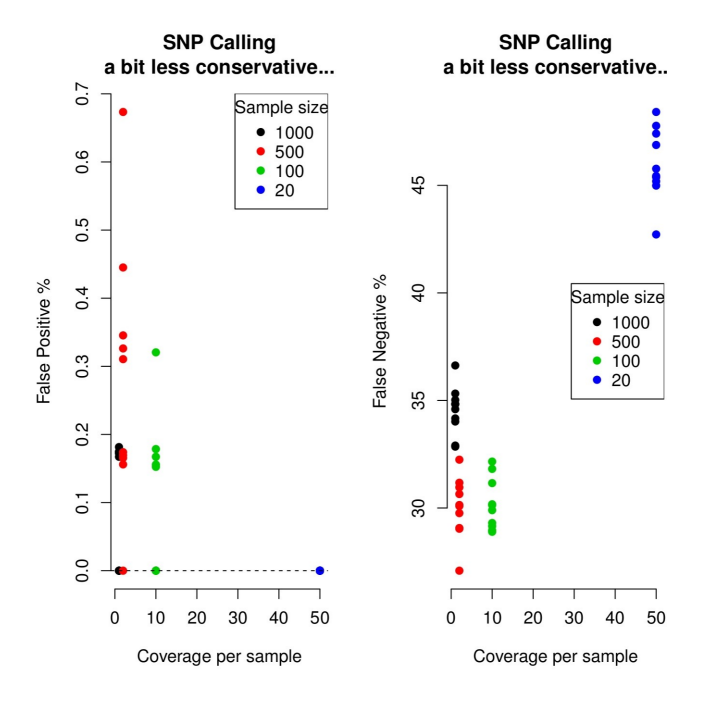
Do you get the right SNPs?



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

