

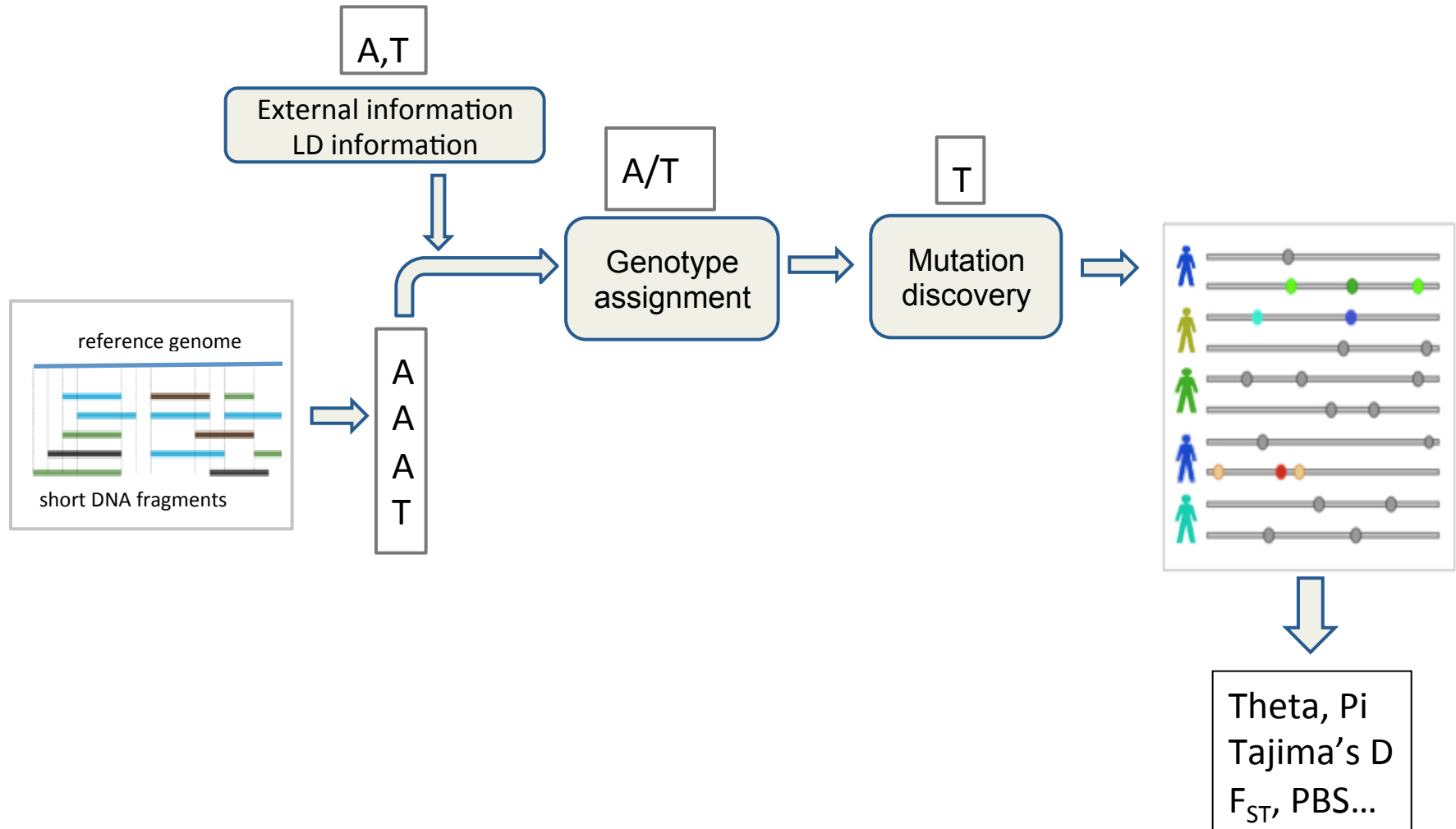
Detecting selection: from low-depth data

Matteo Fumagalli

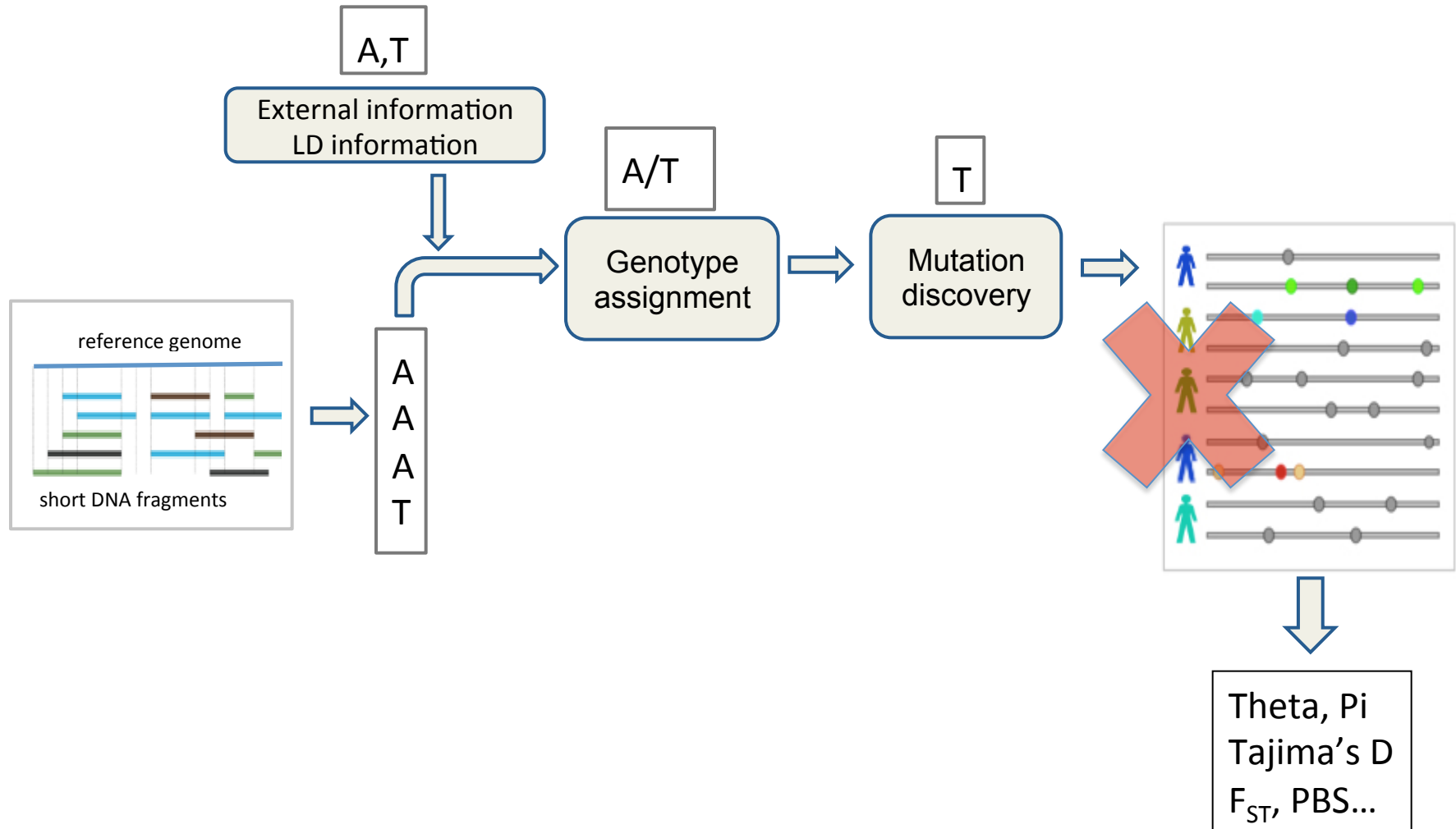
Outline

- Brief introduction to natural selection
- Modes of selection
- Inferring selection at the intra-species level
 - Genetic differentiation
 - Haplotype variation
 - Model-based approaches
 - Testing for significance
- Inferring selection at the inter-species level
- Detecting selection from low-depth sequencing data
- Brief notes on optimal experimental design

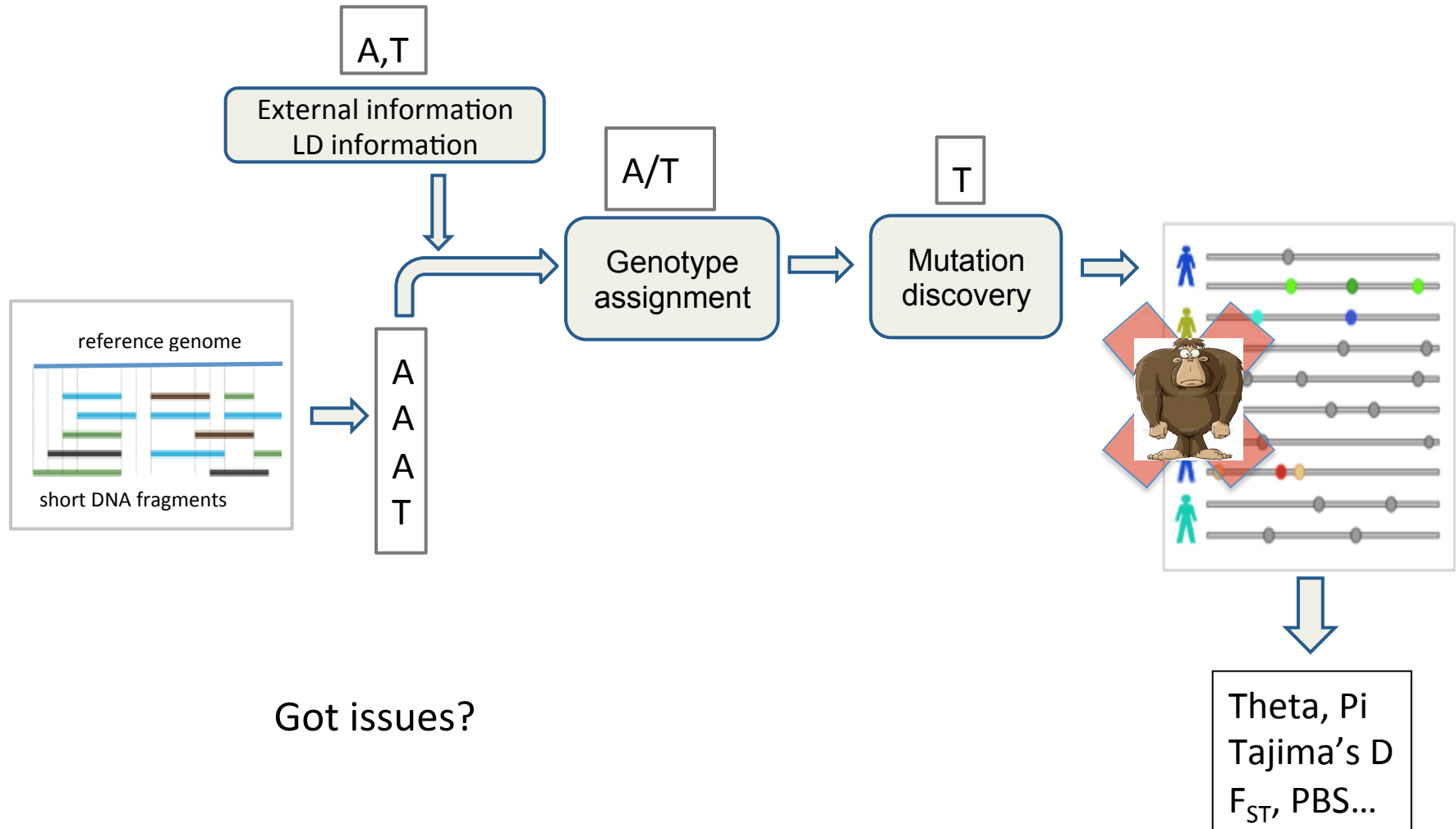
Next Generation Sequencing data processing



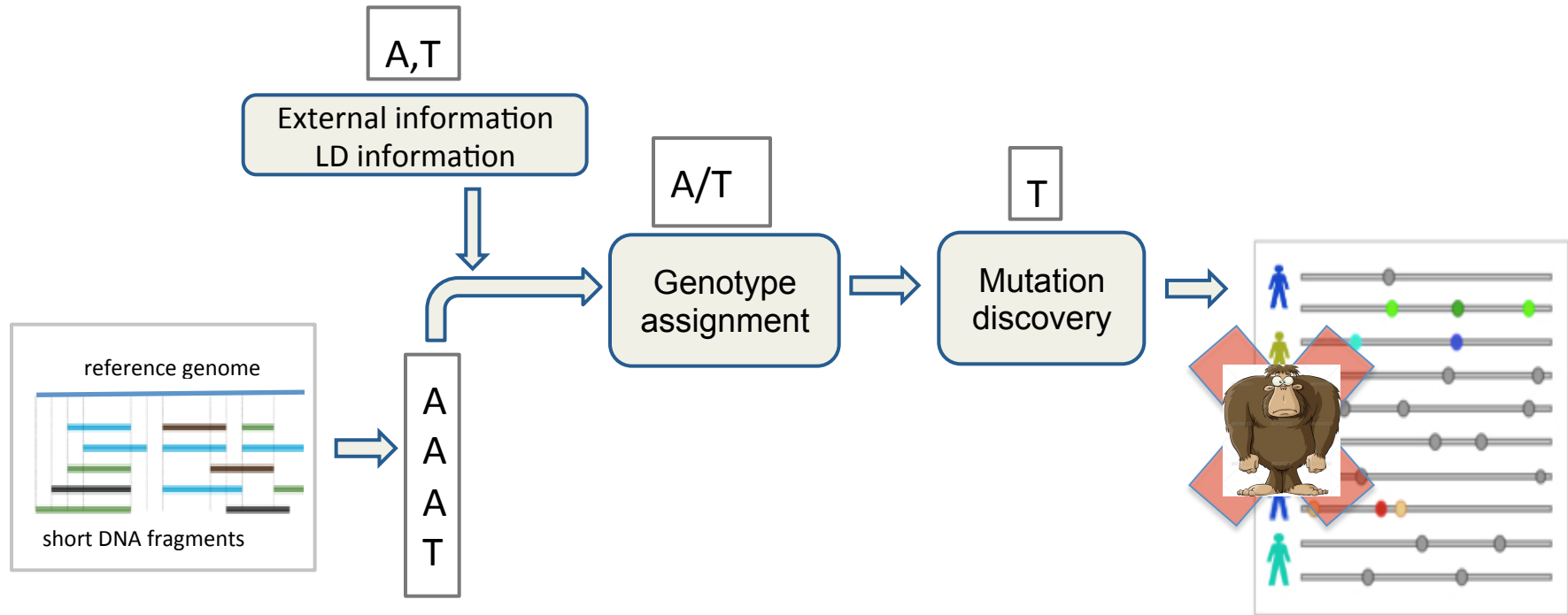
Next Generation Sequencing data processing



Next Generation Sequencing data processing in the non-model world



Next Generation Sequencing data processing in the non-model world

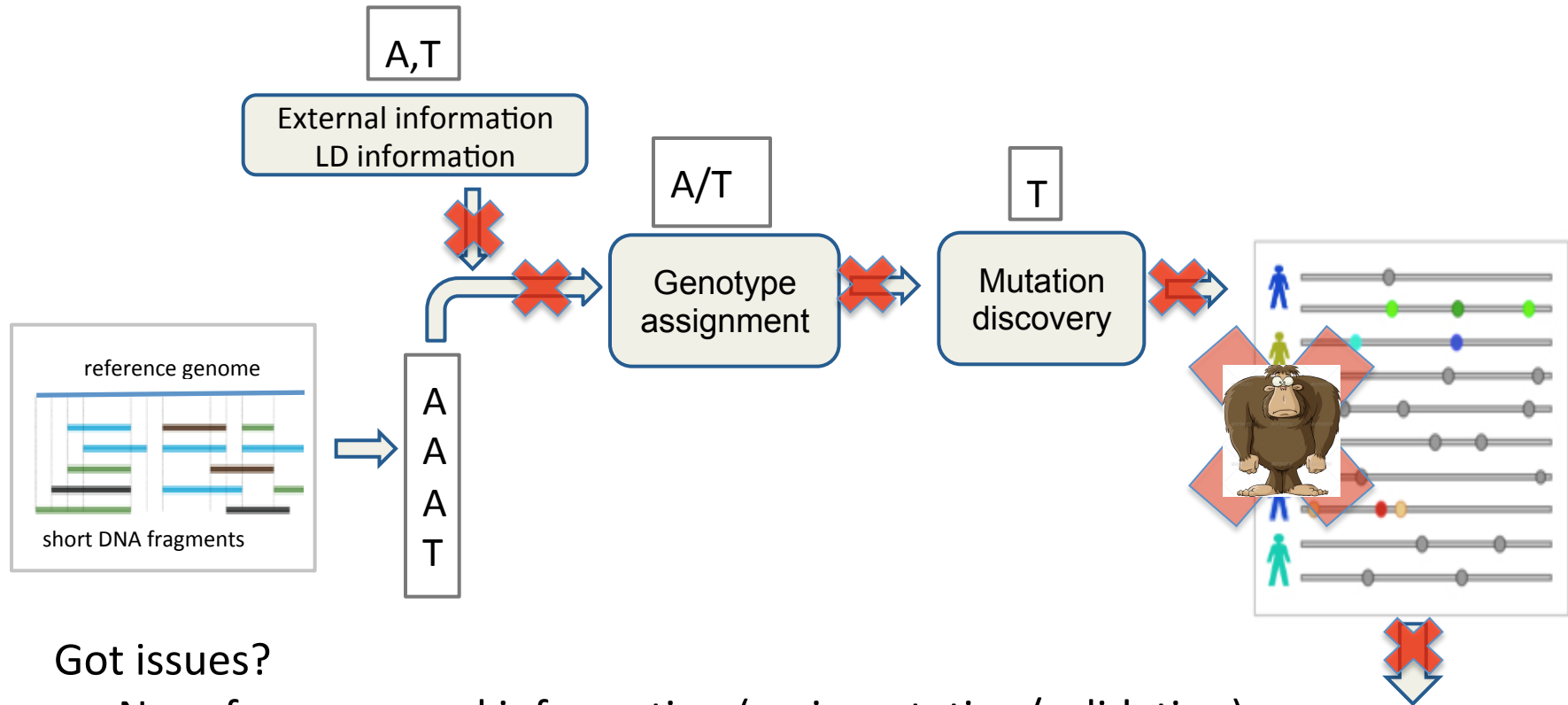


Got issues?

- No reference panel information (no imputation/validation)
- No reference sequence (lower mappability?)
- No HWE assumption (inbred)
- Hyper/Hypovariability or polyploidy or huge genome
- No money (?)
- ...

Theta, Pi
Tajima's D
F_{ST}, PBS...

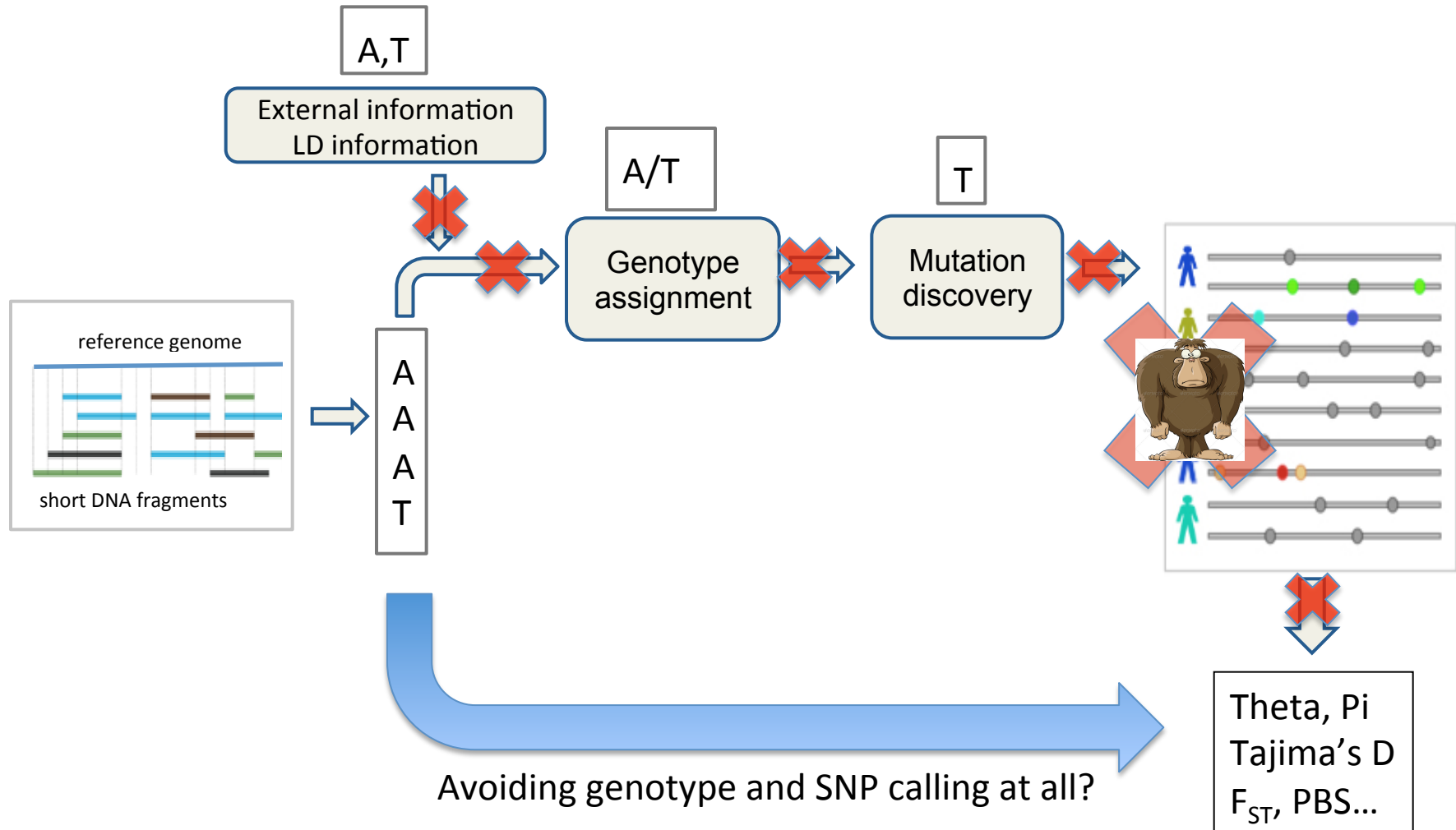
Next Generation Sequencing data processing in the non-model world



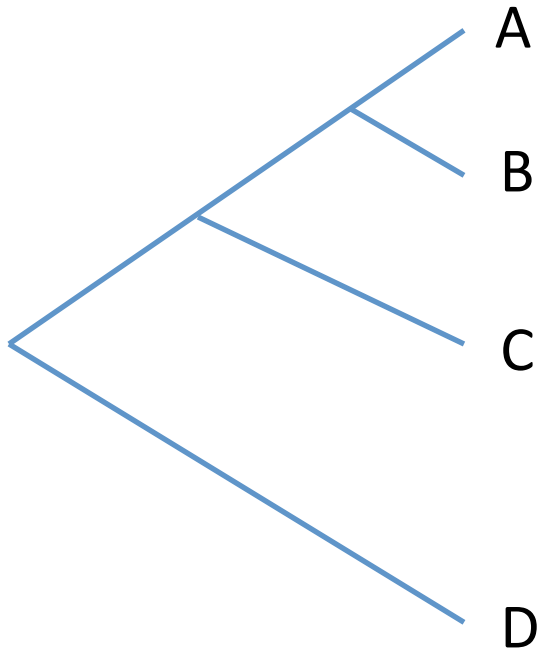
Got issues?

- No reference panel information (no imputation/validation)
- No reference sequence (lower mappability?)
- No HWE assumption (inbred)
- Hyper/Hypovariability or polyploidy or huge genome
- No money (?)
- **Your inferences will be wrong!**

Next Generation Sequencing data processing in the non-model world

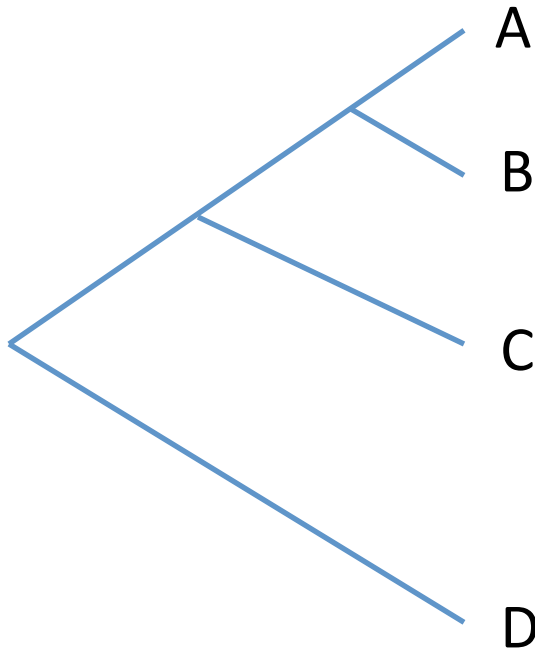


Genetic distances



Genotype 1	Genotype 2	Distance
aa	aa	0
aa	aA	1
aa	AA	2
aA	aa	1
aA	aA	0
aA	AA	2
...

Genetic distances



Genotypes are {aa, aA, AA} as {0, 1, 2}

For individuals i and j and N sites:

$$d(i, j) = -\log \left(1 - \frac{1}{N} \sum_{s=1}^N \frac{|g(i, s) - g(j, s)|}{2} \right)$$

genotype of i at site s

e.g. $G(i=A, s=1)=0$ and $G(j=B, s=1)=1$ then $d(i, j)=1$

Genetic distances from known genotypes

Genotypes are {aa, aA, AA} as {0, 1, 2}

For individuals i and j and N sites:

$$d(i,j) = -\log \left(1 - \frac{1}{N} \sum_{s=1}^N \frac{|g(i,s) - g(j,s)|}{2} \right)$$

$$d(i,j) = 1 * 1.00 = 1.00/2$$

B

A

	0	1	2
0	0	1	0
1	0	0	0
2	0	0	0

Expected value

- The expected value of a discrete random variable is the probability-weighted average of all possible values
- Average value if you perform the same experiment many times

Genetic distances from (un)known genotypes

Genotypes are {aa, aA, AA} as {0, 1, 2}

For individuals i and j and N sites:

$$d(i, j) = -\log \left(1 - \frac{1}{N} \sum_{s=1}^N \frac{|g(i, s) - g(j, s)|}{2} \right)$$

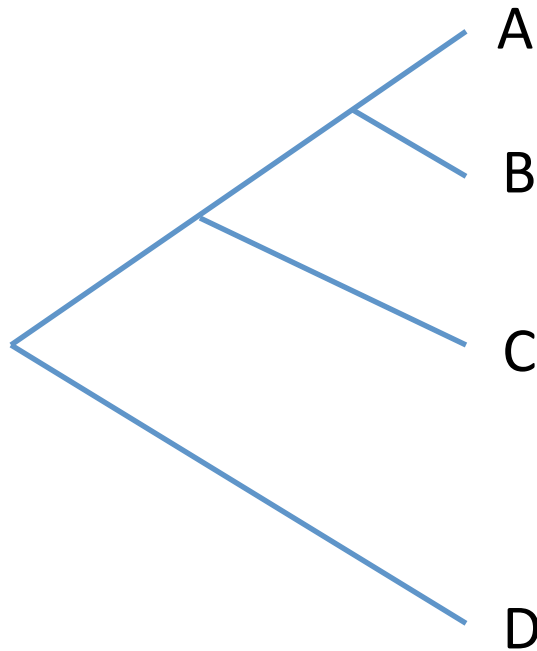
$$E[d(i, j)] = 0 \cdot 0.30 + 1 \cdot 0.50 + 2 \cdot 0.10 + 1 \cdot 0.10 + \dots = 0.80/2$$

B

A

	0	1	2
0	0.30	0.50	0.10
1	0.10	0	0
2	0	0	0

Genetic distances from unknown genotypes



Genotypes are {aa, aA, AA} as {0, 1, 2}

For individuals i and j and N sites:

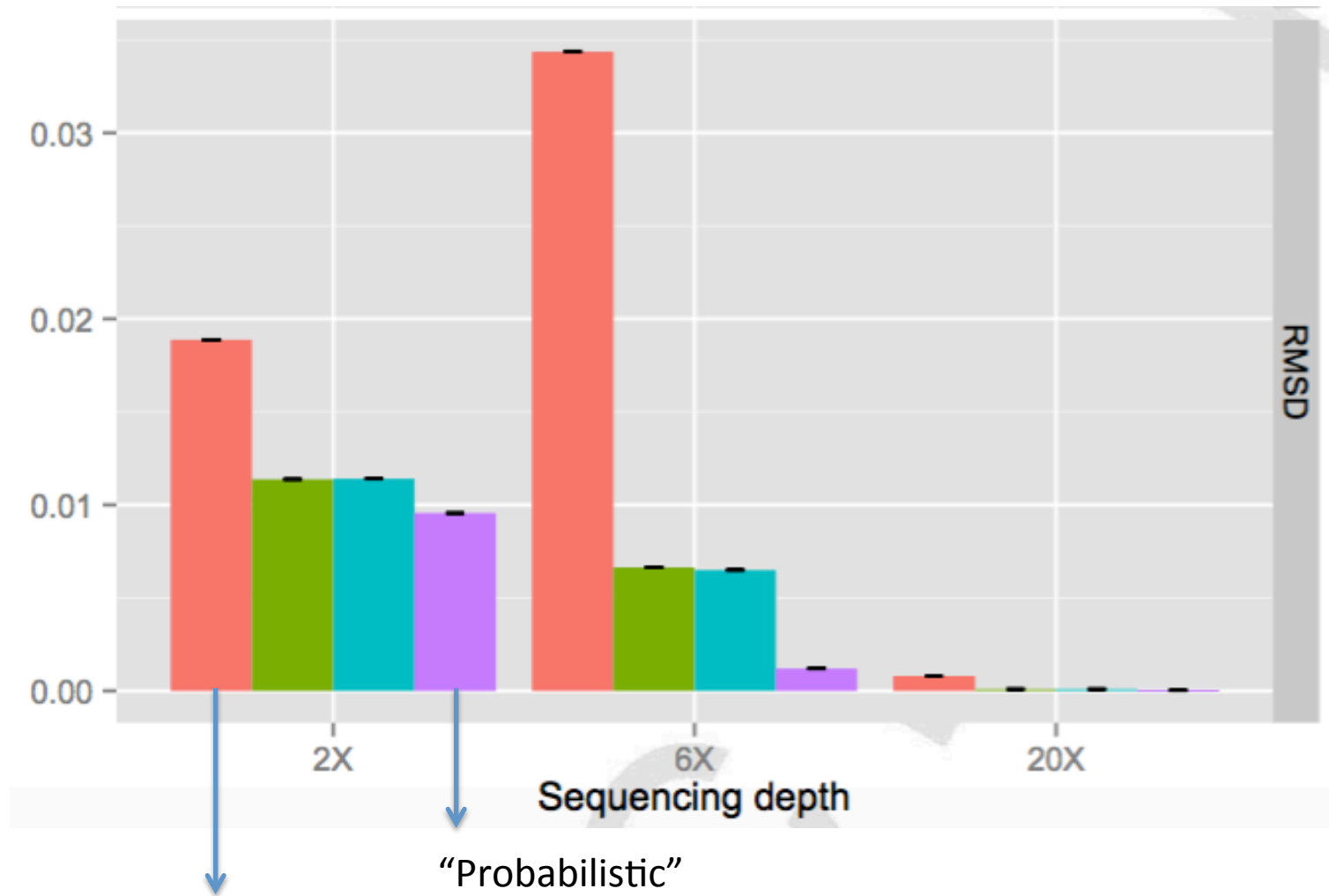
$$d(i, j) = -\log \left(1 - \frac{1}{N} \sum_{s=1}^N \frac{|g(i, s) - g(j, s)|}{2} \right)$$

Iterate across all possible genotypes

Genotypes probability

$$d(i, j) = -\log \left(1 - \frac{1}{N} \sum_{s=1}^N \sum_{g(i, s)=0}^2 \sum_{g(j, s)=0}^2 \frac{|g(i, s) - g(j, s)|}{2} * P(g(i, s), g(j, s)) \right)$$

Genetic distances from unknown genotypes



Genotype calling (no prior)

Vieira et al. BJLS 2016

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

If unfolded, $2k+1$ entries

p_0	p_1	p_2	p_3	...	p_{2k}
-------	-------	-------	-------	-----	----------

If folded, $k+1$ entries

p_0	p_1	p_2	...	p_k
-------	-------	-------	-----	-------

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

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p_0	p_1	p_2	p_3	...	p_{2k}
-------	-------	-------	-------	-----	----------

e.g. A is ancestral, G is derived (alternate)

AA AA AG AA AG AA AA AA AA

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

If unfolded, $2k+1$ entries



e.g. A is ancestral, G is derived (alternate)

AA AA AG AA AG AA AA AA AA

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

If unfolded, $2k+1$ entries

$p_0=0$	$p_1=0$	$p_2=1$	$p_3=0$...	$p_{2k}=0$
---------	---------	---------	---------	-----	------------



e.g. A is ancestral, G is derived (alternate)

AA AA AG AA AG AA AA AA AA

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

If unfolded, $2k+1$ entries

p_0	p_1	p_2	p_3	...	p_{2k}
-------	-------	-------	-------	-----	----------



e.g. A is ancestral, G is derived (alternate)

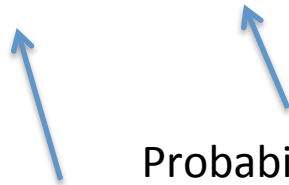
If genotypes are unknown? Counting is not possible?

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

If unfolded, $2k+1$ entries

p_0	p_1	p_2	p_3	...	p_{2k}
-------	-------	-------	-------	-----	----------



Probability of observing 1 copy

Probability of observing 0 copies

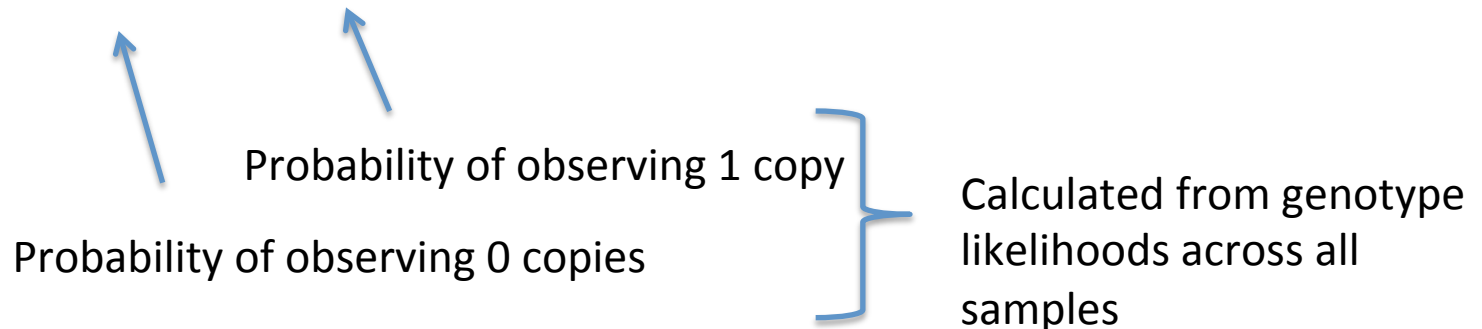
e.g. A is ancestral, G is derived (alternate)

Sample allele frequency

- *With k diploid individuals, how many possible sample allele frequencies can I observe?*

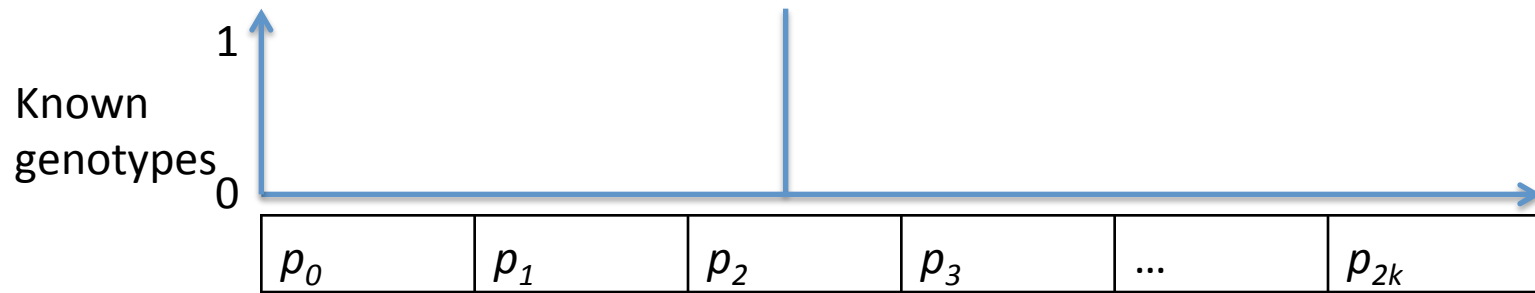
If unfolded, $2k+1$ entries

$p_0=0.05$	$p_1=0.15$	$p_2=0.70$	$p_3=0.10$...	p_{2k}
------------	------------	------------	------------	-----	----------

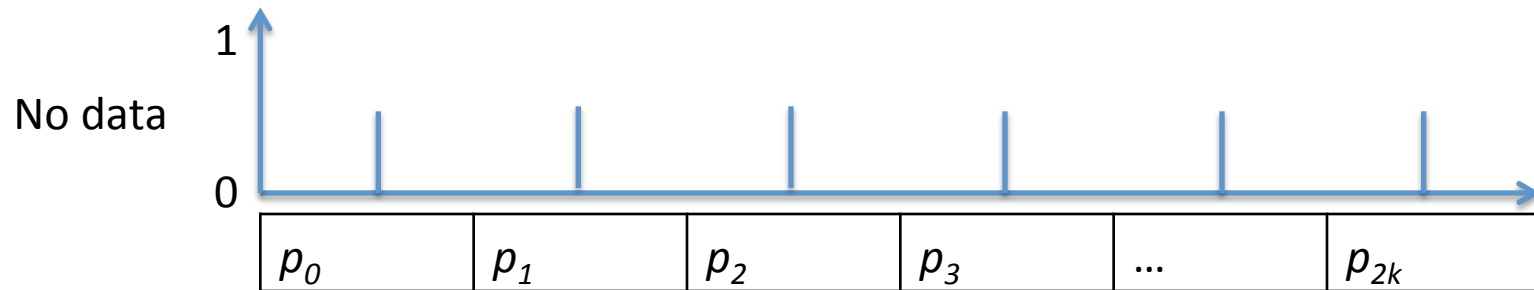
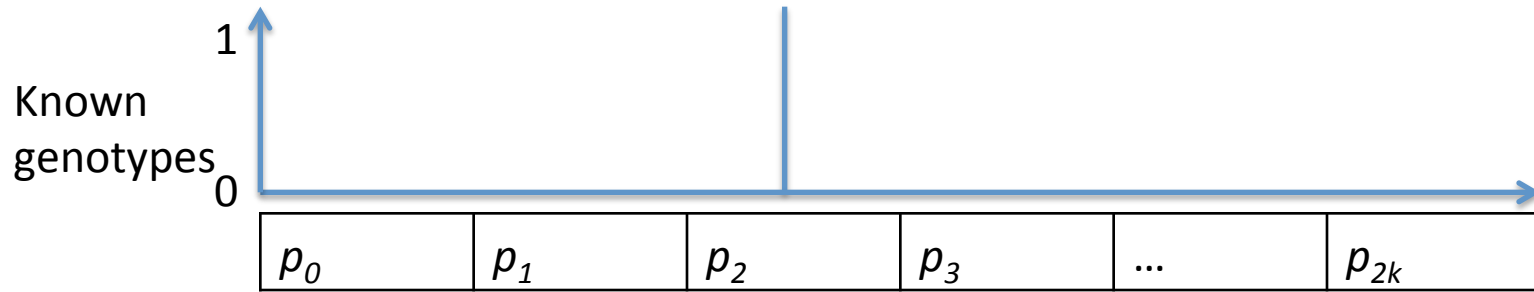


e.g. A is ancestral, G is derived (alternate)

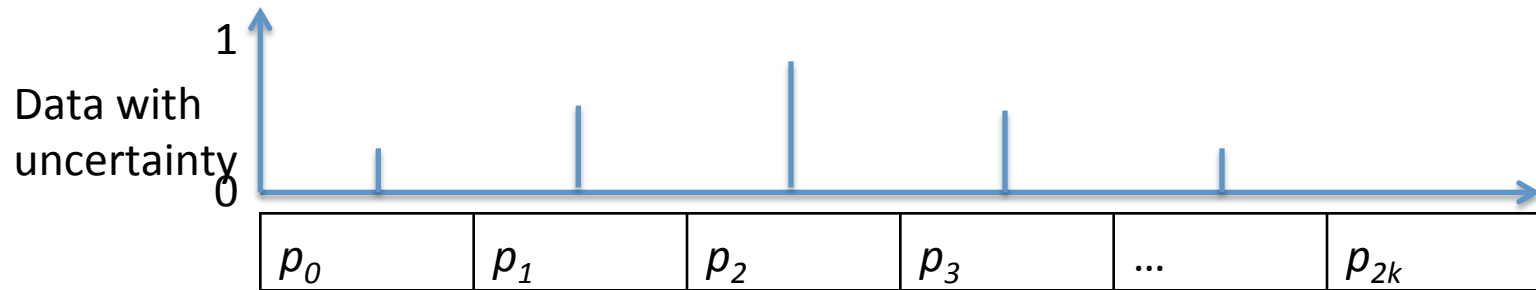
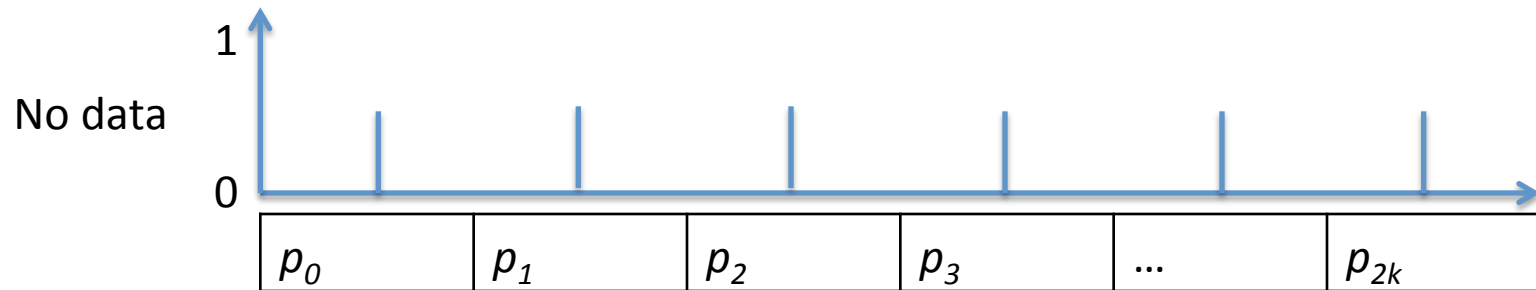
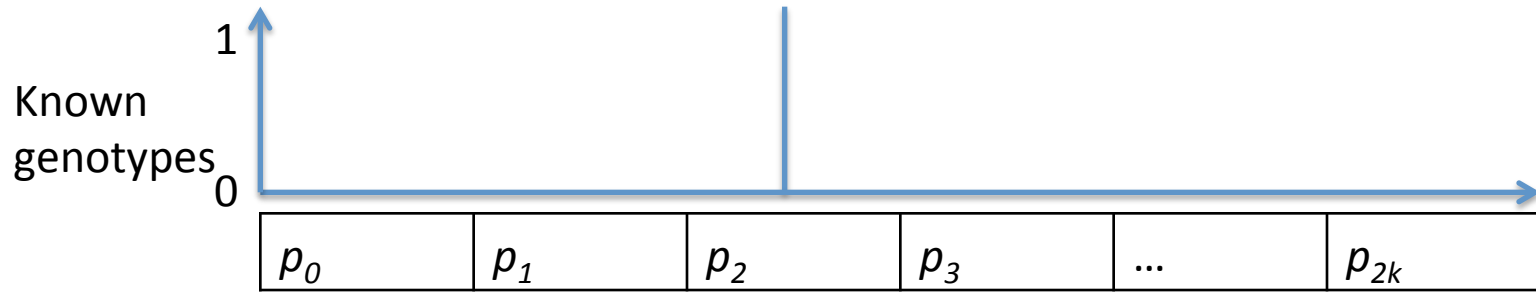
Sample allele frequency probabilities



Sample allele frequency probabilities



Sample allele frequency probabilities



Sample allele frequency posterior probabilities

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

- Estimating allele frequency

$$\hat{f} =$$

Sample allele frequency posterior probabilities

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

- Estimating allele frequency

$$\hat{f} = \sum_{i=0}^{2k} \binom{2k}{i} p(S = i)$$

Sample allele frequency posterior probabilities

With 6 chromosomes (3 diploids)

$p_0=0.10$	$p_1=0.15$	$p_2=0.50$	$p_3=0.15$	$p_4=0.05$	$p_5=0.05$	$p_6=0.00$
------------	------------	------------	------------	------------	------------	------------

- SNP calling

$$p_{\text{var}} = ?$$

$$p_{\text{var}} > t$$

with t being 0.95, 0.99, 0.999 and so on.

Sample allele frequency posterior probabilities

$p_0=0.10$	$p_1=0.15$	$p_2=0.50$	$p_3=0.15$	$p_4=0.05$	$p_5=0.05$	$p_6=0.00$
------------	------------	------------	------------	------------	------------	------------

- SNP calling

$$p_{\text{var}} = 1 - p(S = 0) - p(S = 2k) = 0.90$$

$$p_{\text{var}} > t$$

with t being 0.95, 0.99, 0.999 and so on.

Nr of segregating sites

Site 1

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Site 2

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Site 3

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

...

Site M

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Nr of segregating sites

Site 1

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Site 2

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Site 3

$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
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...

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$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
------------	------------	------------	------------	-----	-------------

Nr of segregating sites

Site 1	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 2	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 3	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
...						
Site M	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$

$$E[S] = \sum_{m=1}^M p_{\text{var}}^{(m)} = \sum_{m=1}^M (1 - p(S_m = 0) - p(S_m = 2k))$$

Nucleotide diversity

Site 1	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 2	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 3	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
...						
Site M	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$

$$D = 2f(1-f)$$

$$E[D] =$$

Nucleotide diversity

Site 1	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 2	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
Site 3	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$
...						
Site M	$p(S_m=0)$	$p(S_m=1)$	$p(S_m=2)$	$p(S_m=3)$...	$p(S_m=2k)$

$$E[D] = \sum_{m=1}^M \sum_{j=0}^{2k} 2 \binom{i}{2k} \binom{2k-i}{2k} p(S_m = i)$$

Applications



...

- Model and non-model species
- Plants
- Ancient genomes
- ...

Software

Such advanced methods have been implemented in several software and utilities, such as:

- **ANGSD** (<http://popgen.dk/ANGSD>)
- **ngsTools** (<https://github.com/mfumagalli/ngsTools>)
- <http://jnpopgen.org/software/>

A Hierarchical Bayesian Model for Next-Generation Population Genomics

Zachariah Gompert¹ and C. Alex Buerkle

Genetics, 2011

which we will explore during the practical session.

Summary

- SNP calling should be performed including information from all samples (and inbreeding coefficient estimates, if relevant)
- Probabilistic methods for estimation of allele frequencies and statistics should be preferred (especially for mean sequencing depth $< 20X$)