





# Plant of the day!

- Pebble plants, *Lithops*, dwarf xerophytes
- Aizoaceae
- South African
- Plants consist of one or more pairs of bulbous leaves – almost no stem
- Leaf markings appear to help plant match its background and be less vulnerable to herbivory



*Lithops lesliei*

# Genomics of Adaptation



# Questions

- What are the genetic changes that underlie adaptation?
- What are the population genetic or genomic signatures of adaptation?
- How do non-adaptive processes affect tests of selection?

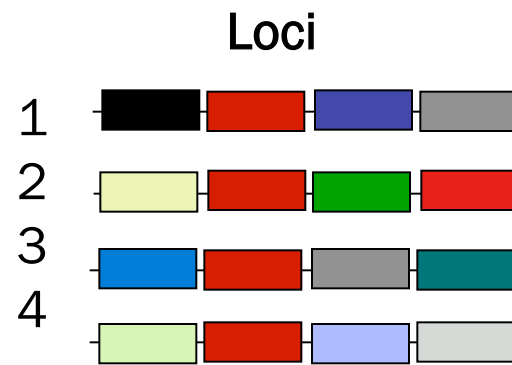
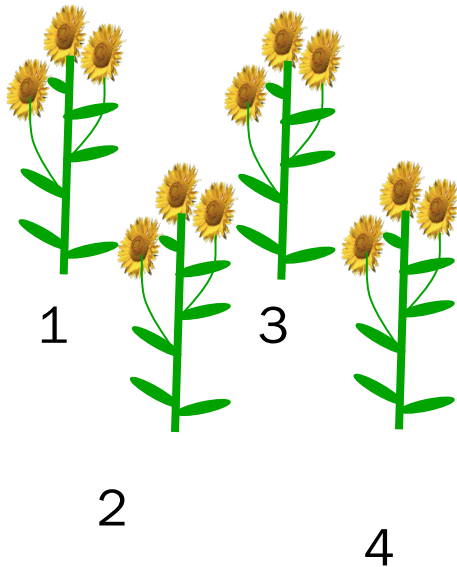
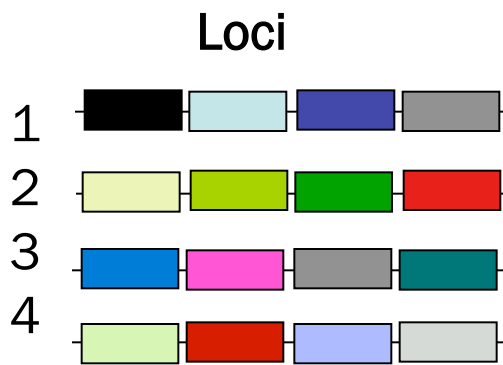
# Goals

- Understand some top down and bottom up approaches used to identify genes responsible for adaptation
- Explain patterns of sequence variation expected with directional and balancing selection
- Understand the principles of population genetic tests of selection

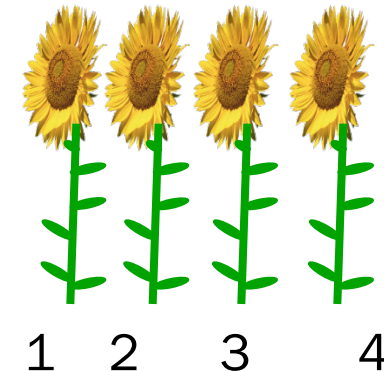
# The genetic basis of adaptation

- Phenotype to genotype (Top down)
  - Identify important trait then find loci associated with it
  - QTL, association mapping, bulk segregant analysis
- Genotype to phenotype (Bottom up)
  - Identify loci under selection, then find trait associated with loci
  - Population genetics

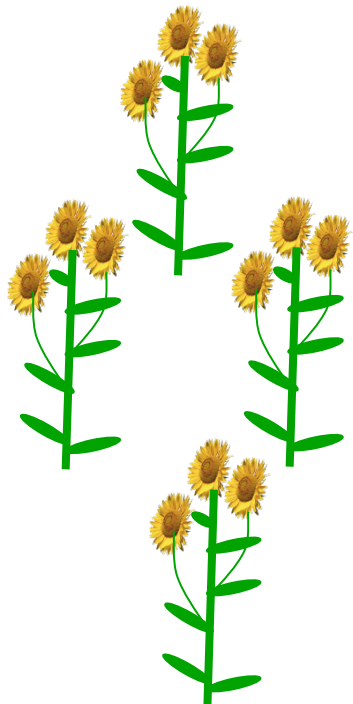
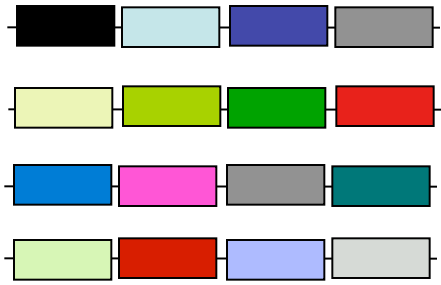
Which locus is likely involved in the change in floral phenotype?



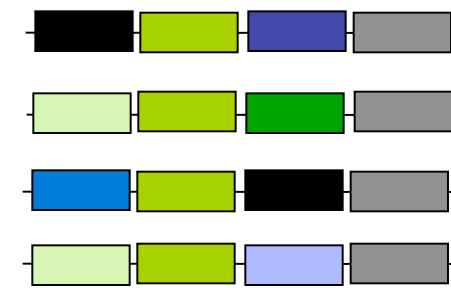
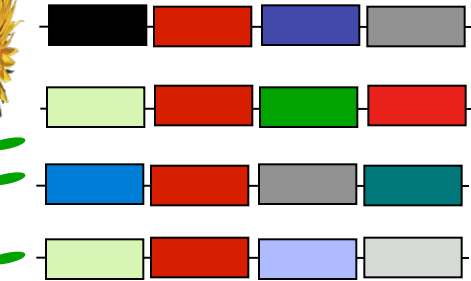
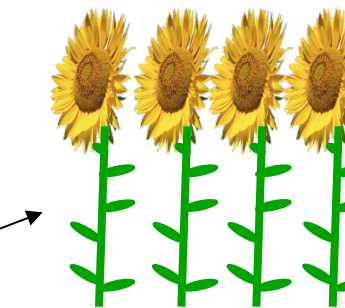
Selective sweep



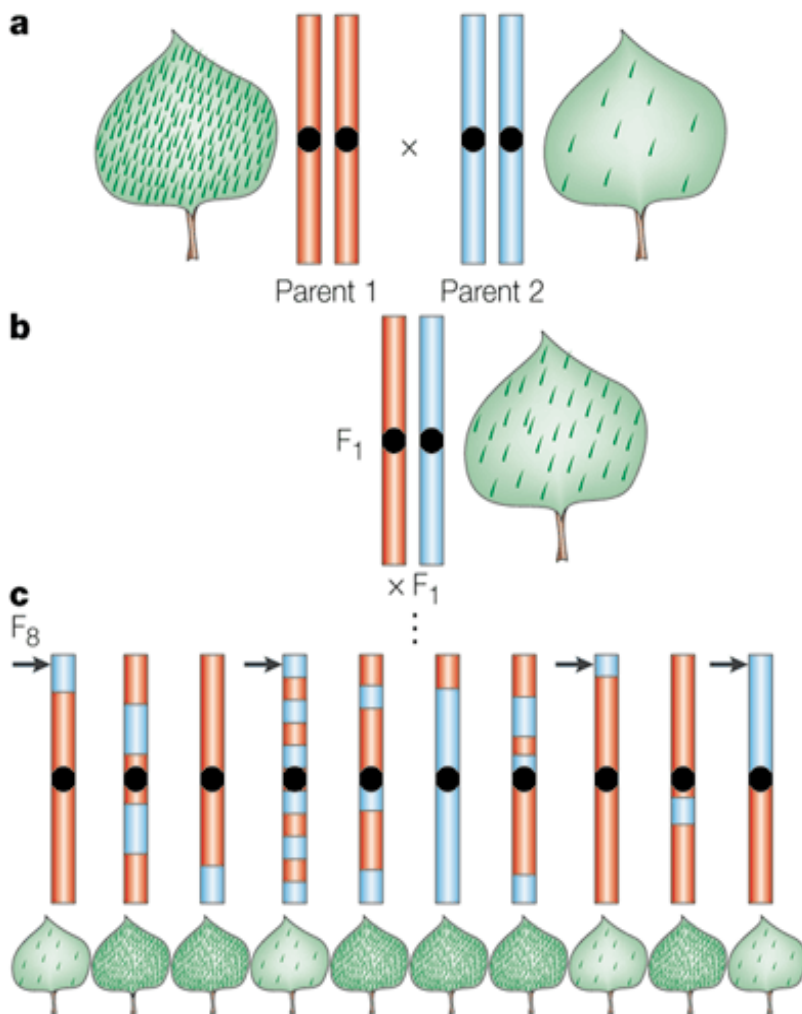
# Which locus is likely involved in the divergence in floral phenotype?



divergence

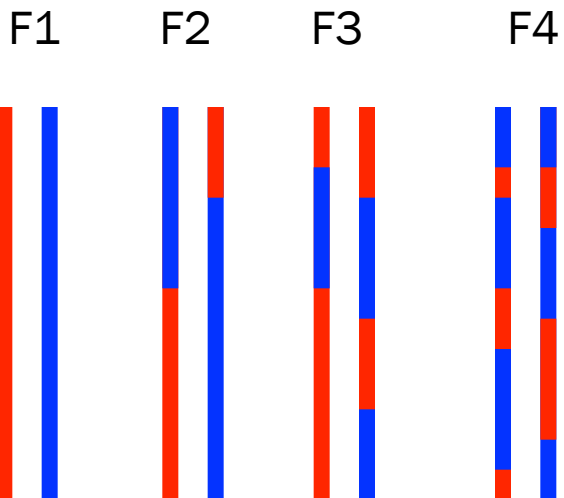


# Quantitative trait loci (QTL)



- Genomic regions associated with trait variation
- Loci detected may differ across individuals/environments
- Statistical issues (sample size, genes of small effect, epistasis)
- Can be large regions of a chromosome (further mapping in region needed)
- Can't perform in all species

# Quantitative trait loci (QTL)



-Precision limited by density of markers and number of recombination events

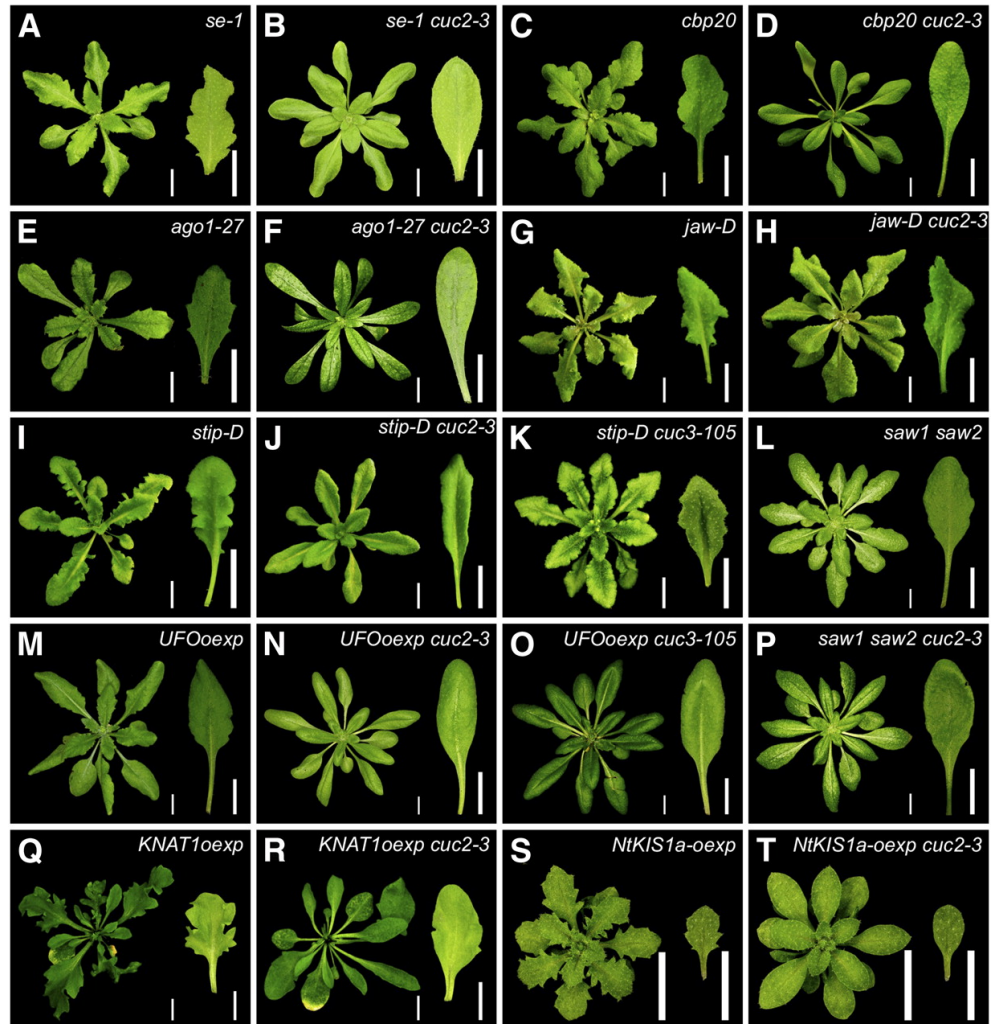
-Recombination events limited by the number of individuals and their degree of recombination between the parental genomes (i.e. F2, F3, etc)

Parental genomes are more finely recombined with each generation of consecutive intercrosses.

# Association mapping

Associations between markers (SNPs) and phenotypes in natural populations

- Different populations of *Arabidopsis* have different leaf shape.
- Look through whole genome to find SNPs associated with leaf shape.



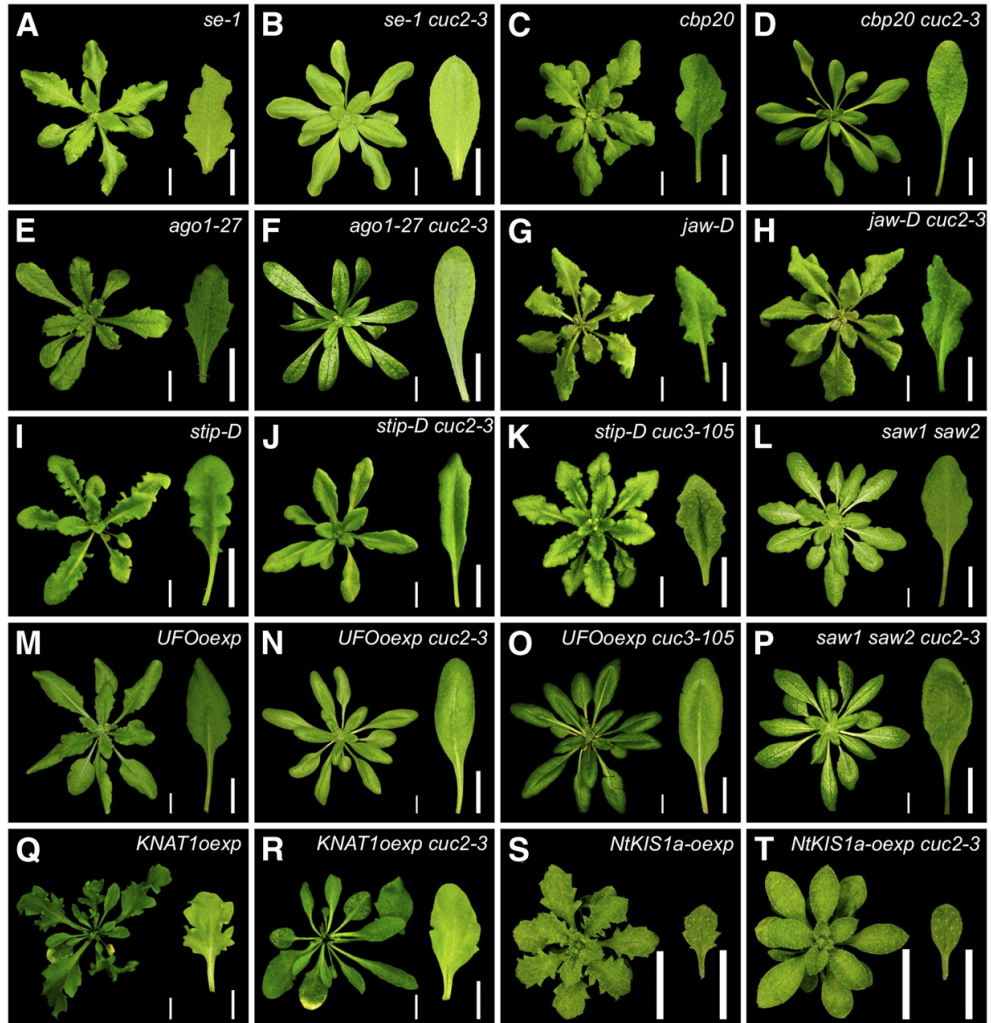
# Association mapping

Pros:

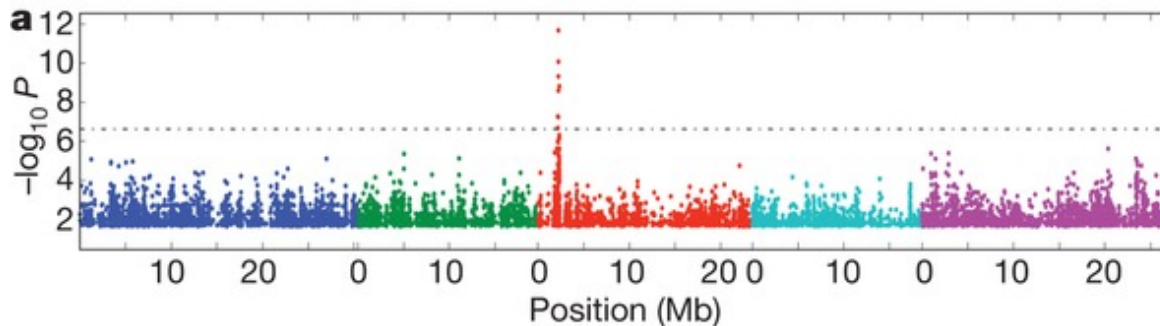
- Much higher resolution
- No need for crosses

Cons:

- Population structure may lead to spurious associations
- Need many many markers, more than QTL mapping.

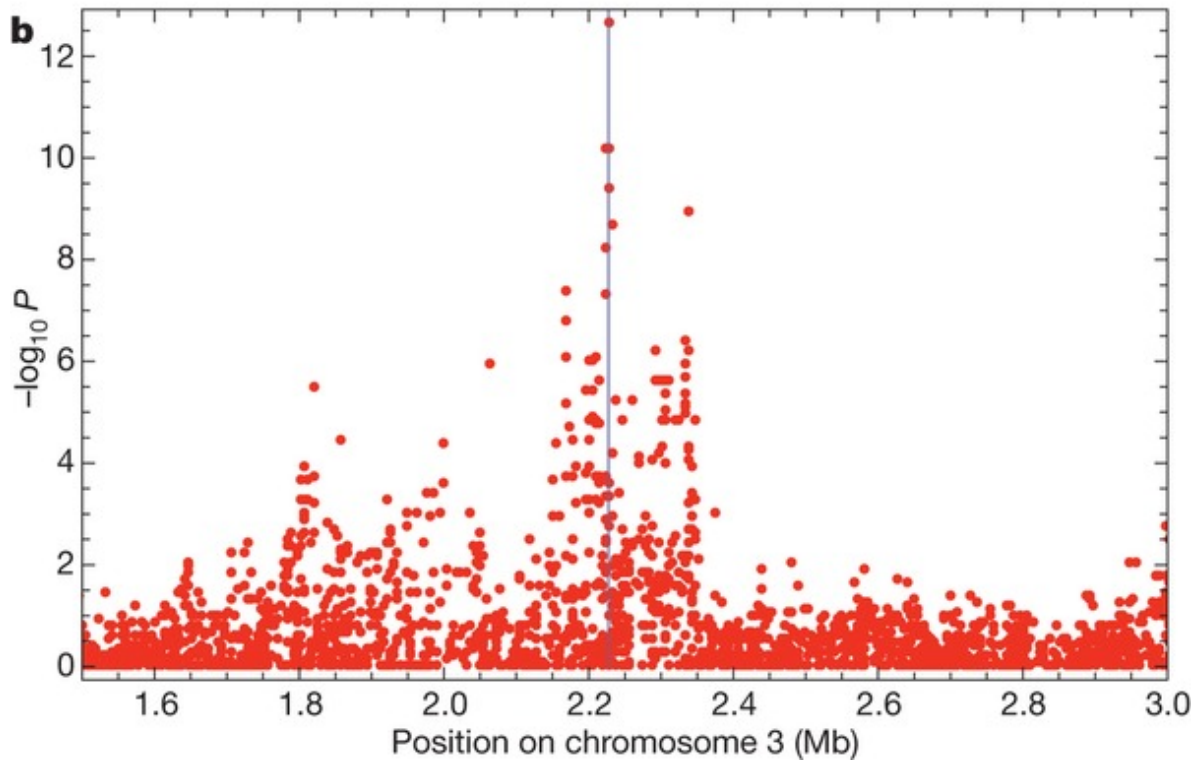


# Example association map



Response to the avirulence gene *AvrRpm1*

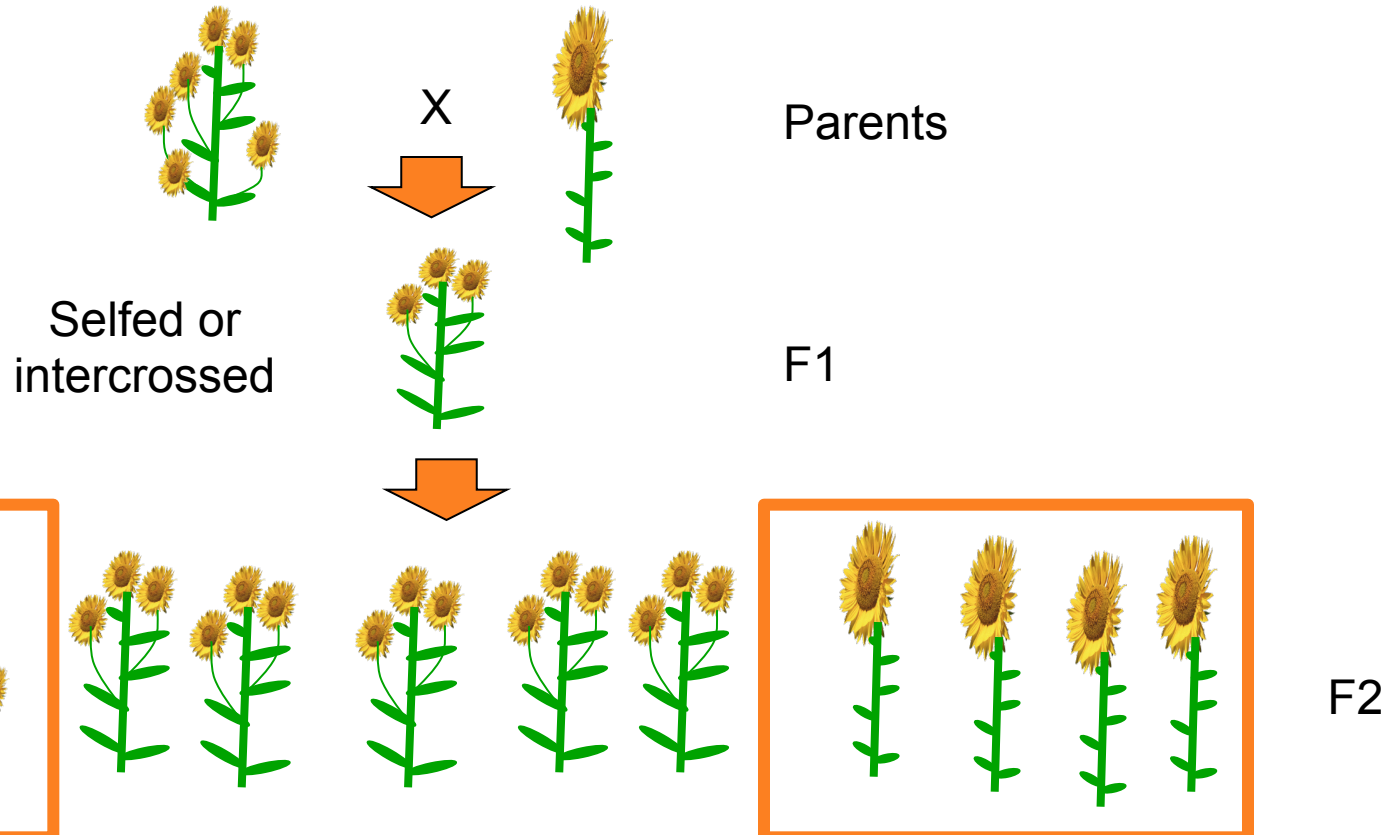
- A very simple trait



Used 95 inbred lines and 250,000 SNPs

# Bulk Segregant Analysis

- Cross two plants divergent phenotype, then self or intercross to make an F<sub>2</sub> population
- Select F<sub>2</sub> individuals with extreme phenotypes for the trait
- Genotype both pools for many markers
- Look for genes where different alleles are enriched in each pool



# The genetic basis of adaptation

- Phenotype to genotype (Top down)
  - Identify important trait then find loci associated with it
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- Genotype to phenotype (Bottom up)
  - Identify loci under selection, then find trait associated with loci
  - Population genetics

# Detecting natural selection

- The Neutral theory suggests that most molecular changes are neutral and are caused by random genetic drift
- This is used as a null hypothesis and deviations from neutral expectations are evidence of selection
- Important to consider how non-selective processes like population structure and linkage affect the statistics

# The effect of selection on the genome

## Directional selection

- Best allele(s) sweep to fixation
- Loss of variation
- Change in frequency distribution of polymorphisms
- Increase in linkage disequilibrium around the site

# The effect of selection on the genome

## Directional selection

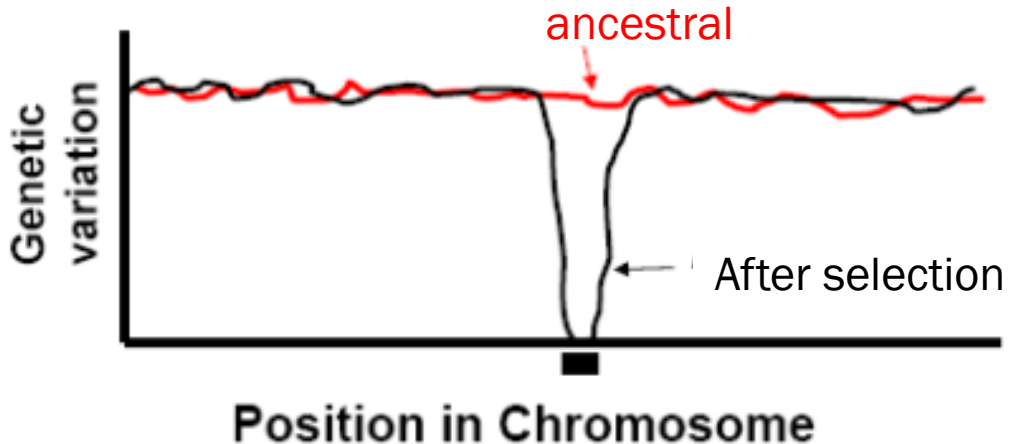
- Best allele(s) sweep to fixation
- Loss of variation
- Change in frequency distribution of polymorphisms
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## Balancing selection

- Maintains variation that otherwise would be lost to drift
- Heterozygote advantage, frequency dependent selection, fluctuating selection, (divergent selection)

# Directional selection

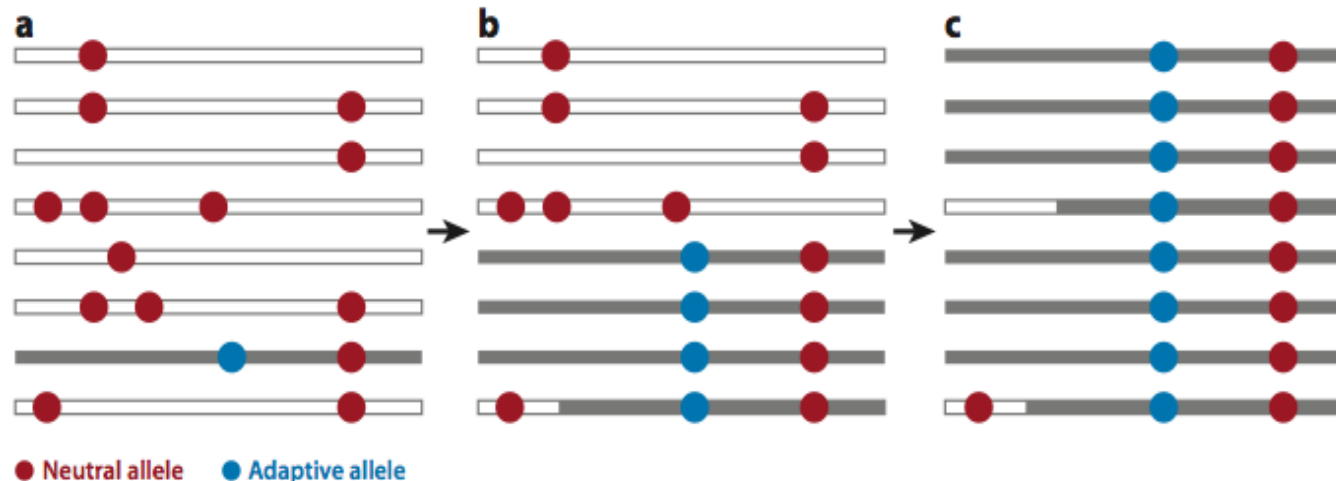
- A beneficial allele arises
- Variants with this allele rapidly spread through the species
- Genetic diversity is reduced around this adaptive locus



# Chance of detecting natural selection

Depends on:

- Time
- Strength of selection
- Recombination, mutation
- Initial frequency



Selective sweep

# Methods for detecting selection

- A. MacDonald-Kreitman Type Tests
- B. Site Frequency Spectrum Approaches
- C. Linkage Disequilibrium (LD) and Haplotype Structure
- D. Population Differentiation: Lewontin-Krakauer Methods

These tests can be applied to single genes, or across the whole genome.

# A. MacDonald-Krietman type tests

- **Synonymous substitutions:**
  - Mutations that do not cause amino acid change (usually 3rd position)  
“silent substitutions”
- **Nonsynonymous substitutions:**
  - Mutations that cause amino acid change (1st, 2nd position)  
“replacement substitutions”

First base	Second base				Third base
	U	C	A	G	
U	UUU Phenylalanine UUC Phenylalanine UUA Leucine UUG Leucine	UCU Serine UCC Serine UCA Serine UCG Serine	UAU Tyrosine UAC Tyrosine UAA Stop UAG Stop	UGU Cysteine UGC Cysteine UGA Stop UGG Tryptophan	U C A G
	CUU Leucine CUC Leucine CUA Leucine CUG Leucine	CCU Proline CCC Proline CCA Proline CCG Proline	CAU Histidine CAC Histidine CAA Glutamine CAG Glutamine	CGU Arginine CGC Arginine CGA Arginine CGG Arginine	U C A G
	AUU Isoleucine AUC Isoleucine AUA Isoleucine AUG Start (Methionine)	ACU Threonine ACC Threonine ACA Threonine ACG Threonine	AAU Asparagine AAC Asparagine AAA Lysine AAG Lysine	AGU Serine AGC Serine AGA Arginine AGG Arginine	U C A G
	GUU Valine GUC Valine GUA Valine GUG Valine	GCU Alanine GCC Alanine GCA Alanine GCG Alanine	GAU Aspartic Acid GAC Aspartic Acid GAA Glutamic Acid GAG Glutamic Acid	GGU Glycine GGC Glycine GGA Glycine GGG Glycine	U C A G
Codon      Amino acid					

# A. MacDonald-Krietman type tests

First base	Second base				Third base
	U	C	A	G	
U	UUU Phenylalanine UUC Phenylalanine UUA Leucine UUG Leucine	UCU Serine UCC Serine <b>UCA</b> Serine UCG Serine	UAU Tyrosine UAC Tyrosine UAA Stop UAG Stop	UGU Cysteine UGC Cysteine UGA Stop UGG Tryptophan	U C A G
	CUU Leucine CUC Leucine CUA Leucine CUG Leucine	CCU Proline CCC Proline CCA Proline CCG Proline	CAU Histidine CAC Histidine CAA Glutamine CAG Glutamine	CGU Arginine CGC Arginine CGA Arginine CGG Arginine	
	AUU Isoleucine AUC Isoleucine AUA Isoleucine AUG Start (Methionine)	ACU Threonine ACC Threonine ACA Threonine ACG Threonine	AAU Asparagine AAC Asparagine AAA Lysine AAG Lysine	AGU Serine AGC Serine AGA Arginine AGG Arginine	
	<b>GUU</b> Valine GUC Valine GUA Valine GUG Valine	GCU Alanine GCC Alanine <b>GCA</b> Alanine GCG Alanine	GAU Aspartic Acid GAC Aspartic Acid GAA Glutamic Acid GAG Glutamic Acid	GGU Glycine <b>GGC</b> Glycine GGA Glycine GGG Glycine	
Codon      Amino acid					
GCU - Alanine					
GCC - Alanine	GUU - Valine	Synonymous	Nonsynonymous		

# A. MacDonald-Krietman type tests

## $K_a/K_s$ Test

Nonsynonymous substitutions

Synonymous substitutions

$\frac{K_a}{K_s}$

- Uses coding sequence (sequence that codes proteins)
- Controls for max possible rate of each type of substitution
- $K_s$  doesn't change protein so is "neutral" and is used as baseline rate
- Important to remember that both types of mutations occur at the same rate, it is fixation rate that varies.

# A. MacDonald-Krietman type tests

## $K_a/K_s$ Test

Nonsynonymous substitutions

$\frac{K_a}{K_s}$

Synonymous substitutions

- $K_a/K_s = 1$  --- Neutral drift. Protein changes aren't being selected for or against.
- $K_a/K_s > 1$  --- Positive selection. Protein changes are being selected for
- $K_a/K_s < 1$  --- Purifying selection. Protein changes are being selected against.

# A. MacDonald-Krietman type tests

## $K_a/K_s$ Test

Nonsynonymous substitutions

Synonymous substitutions

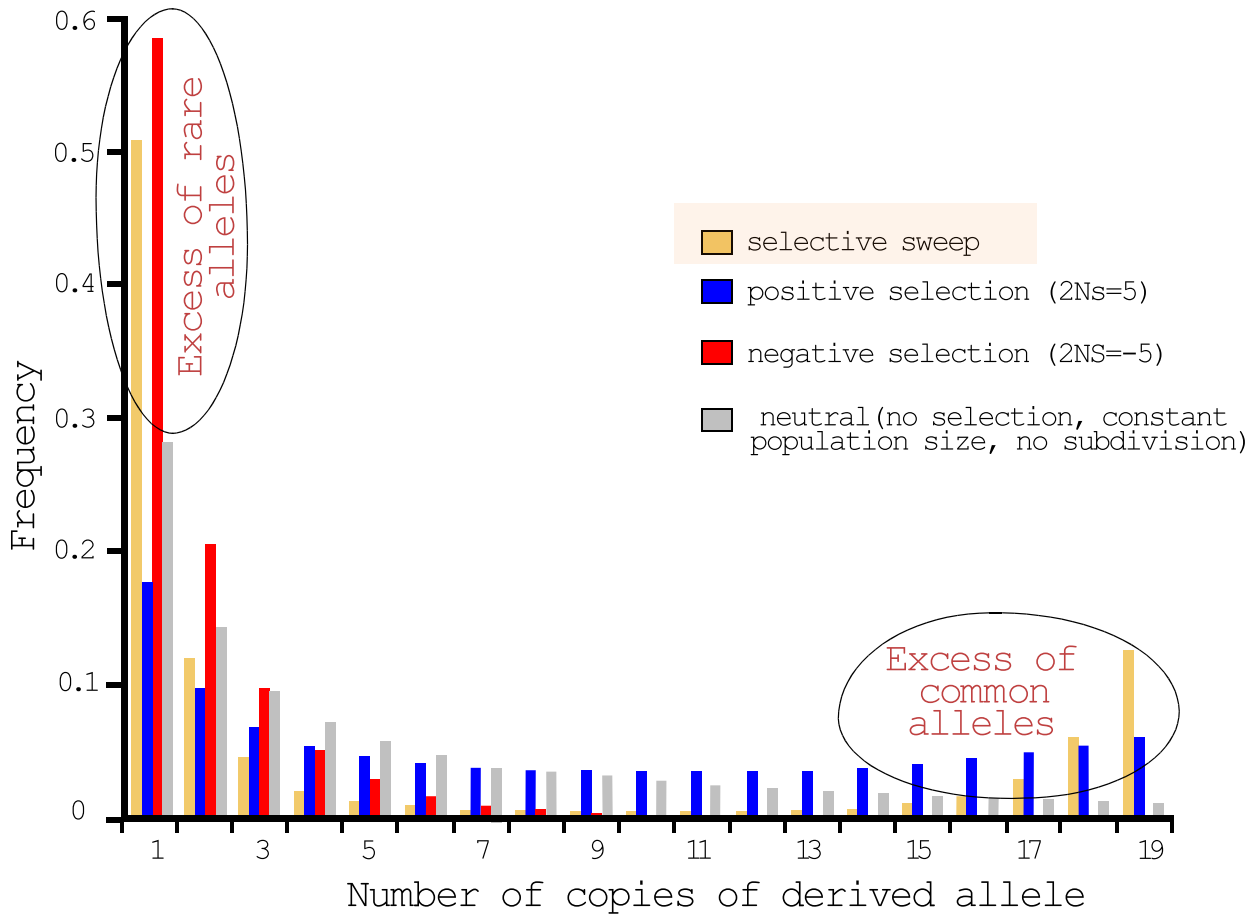
$$\frac{K_a}{K_s}$$

- Can be done with single sequences per species/group (don't need population genetics data)
- Can pinpoint where selection occurred on a phylogeny
- Proteins very rarely have  $K_a/K_s > 1$  for their entire sequence, often only small pieces or single codons are under selection
  - Proteins with  $K_a/K_s > 1$  are often under diversifying selection, e.g. immune or self-incompatibility genes

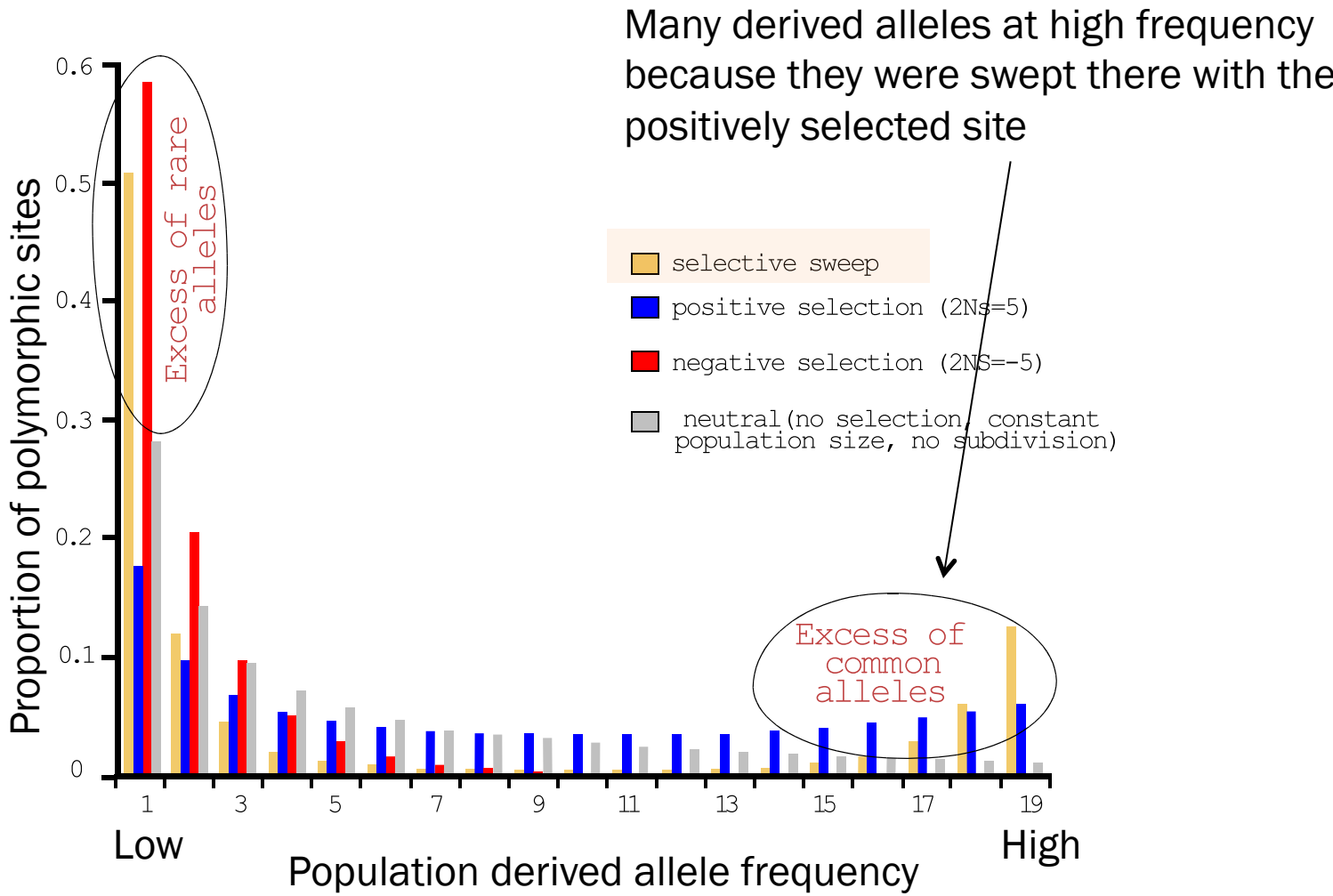
# B. Site Frequency Spectrum

- Selection affects the distribution of alleles within populations
- Method examines site frequency spectrum and compares to neutral expectations
- Could be applied to a single locus. Now used often for genomic scans for selective sweeps

# B. Site Frequency Spectrum

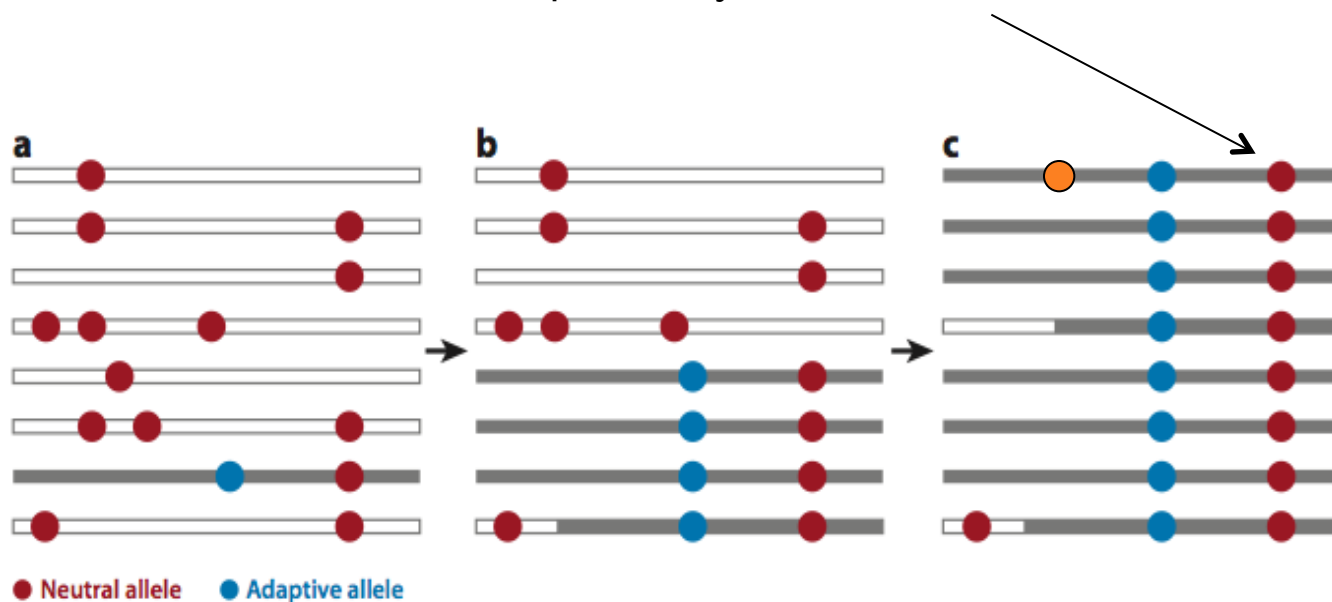


# B. Site Frequency Spectrum

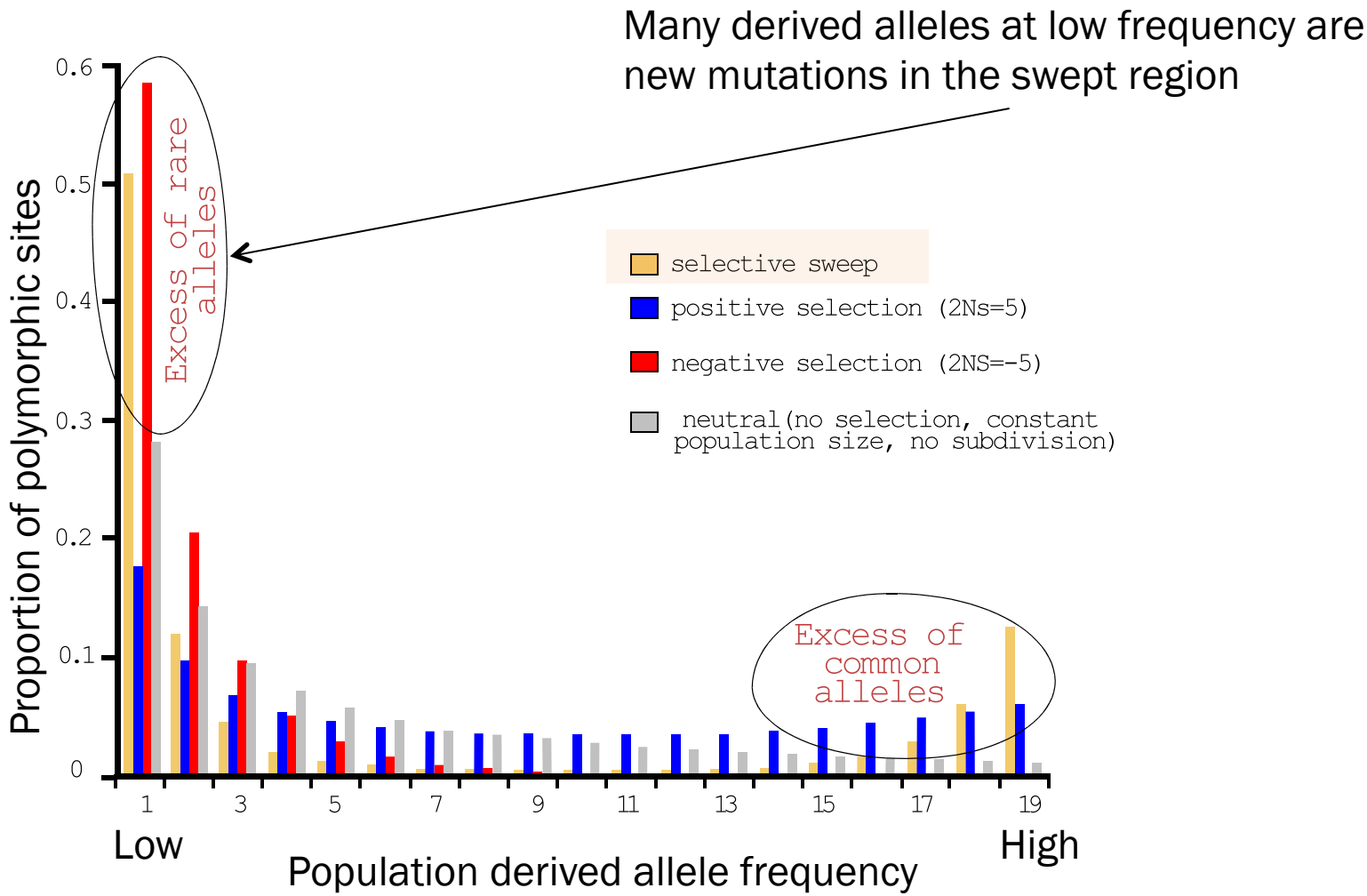


## B. Site Frequency Spectrum

Many derived alleles at high frequency because they were swept there with the positively selected site

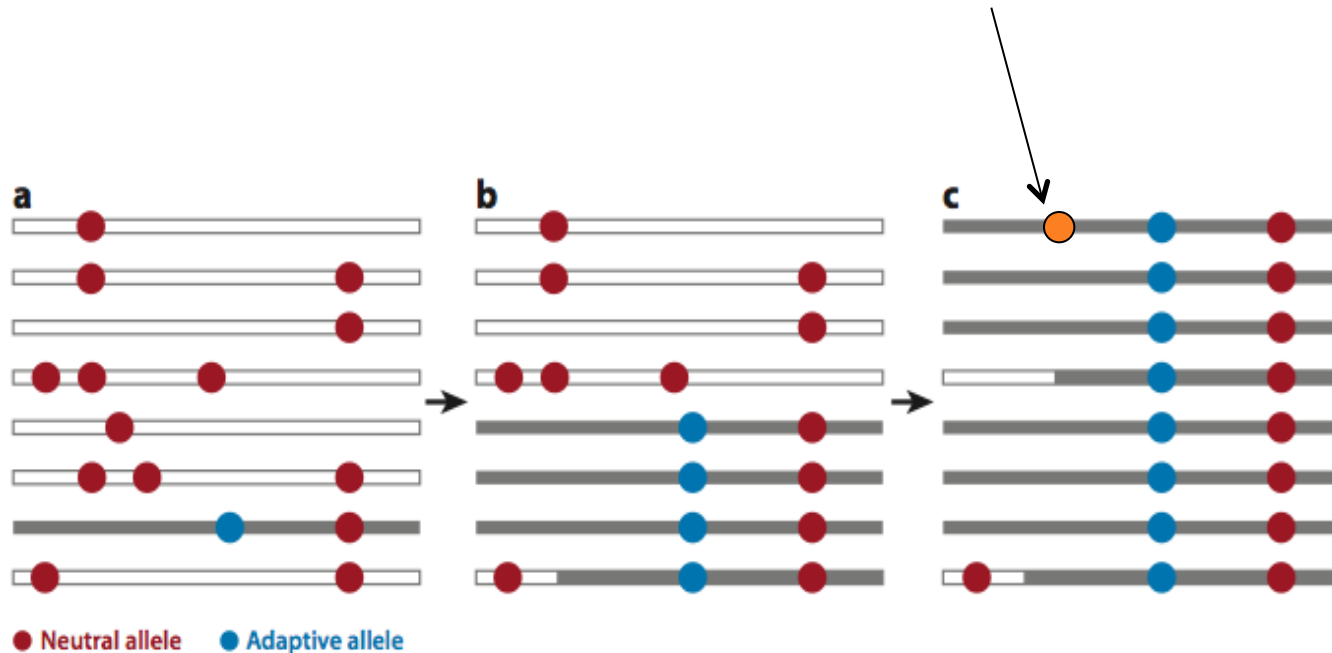


# B. Site Frequency Spectrum



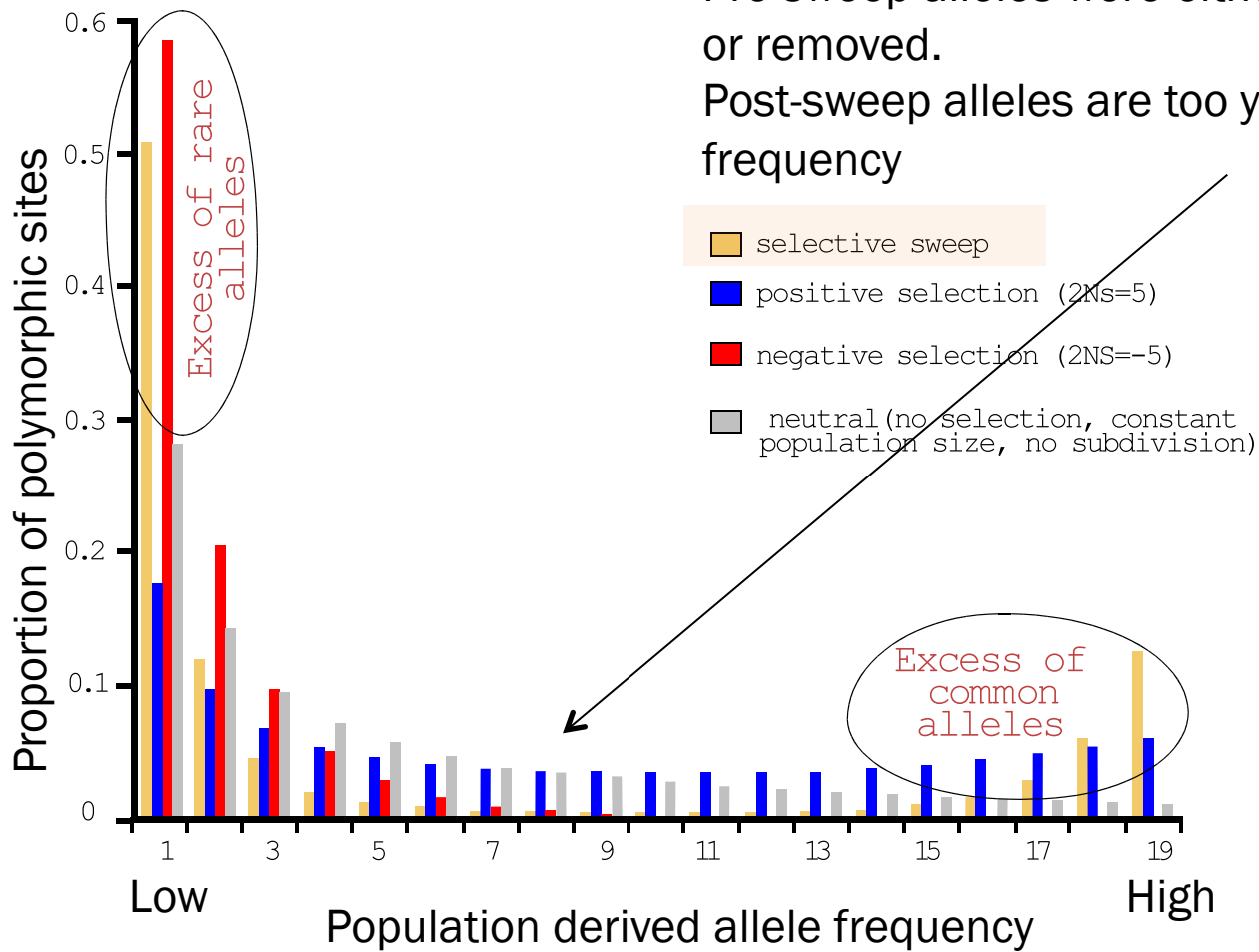
# B. Site Frequency Spectrum

Many derived alleles at low frequency are new mutations in the swept region



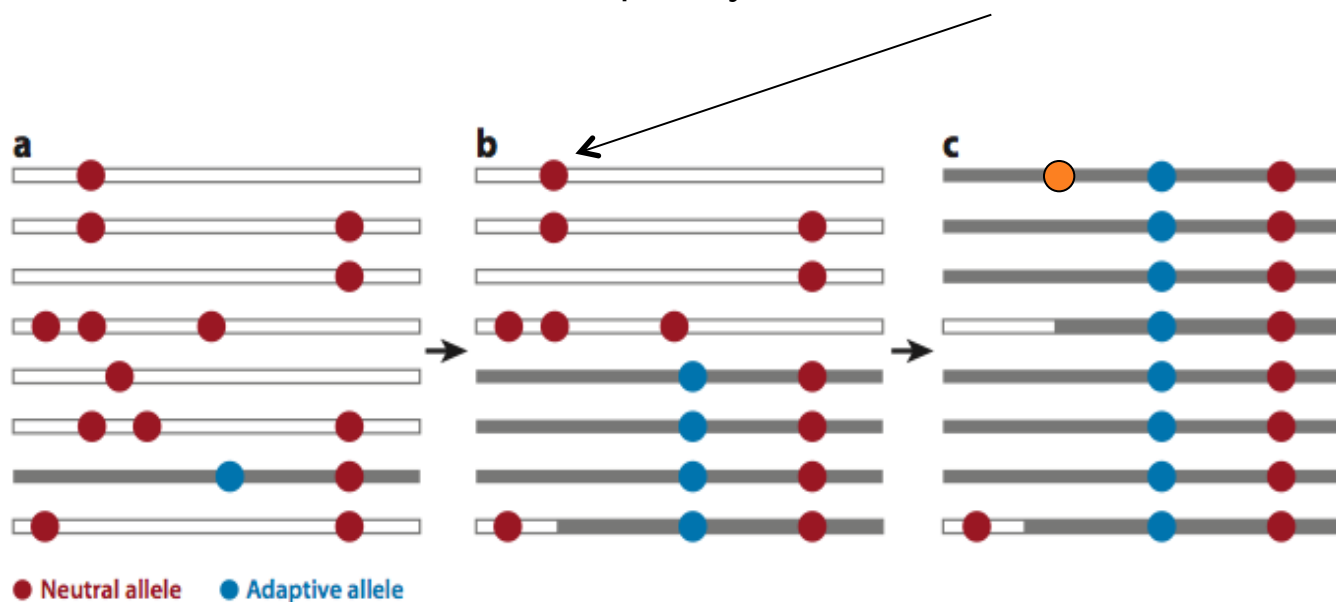
# B. Site Frequency Spectrum

Few medium frequency derived alleles.  
Pre-sweep alleles were either swept to high frequency or removed.  
Post-sweep alleles are too young to reach medium frequency

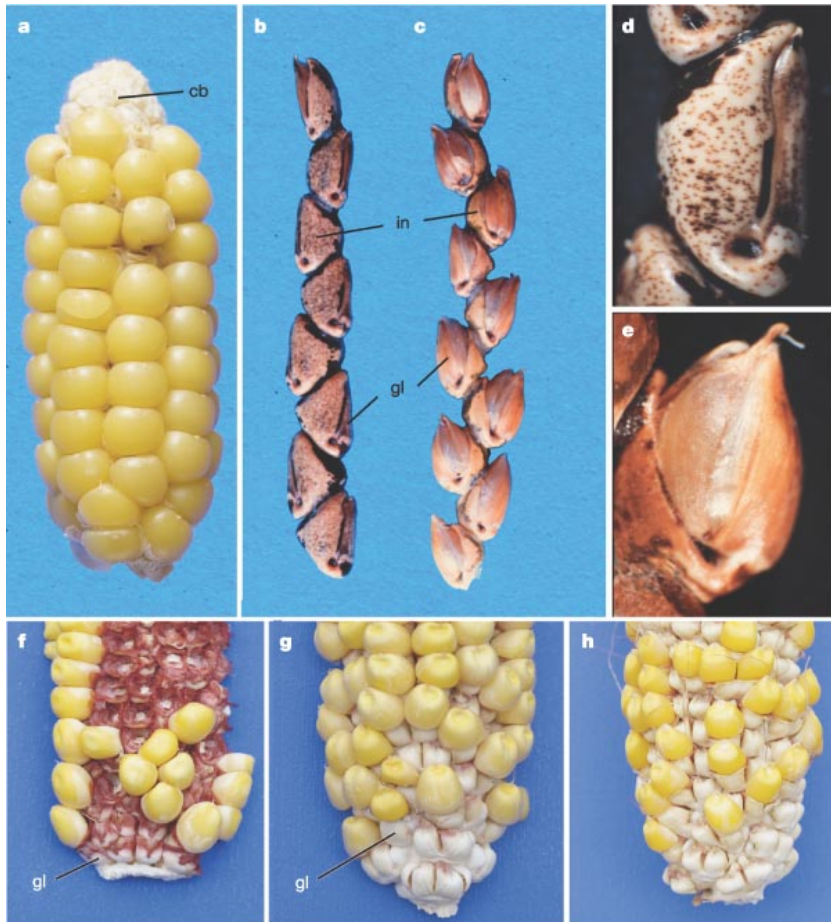


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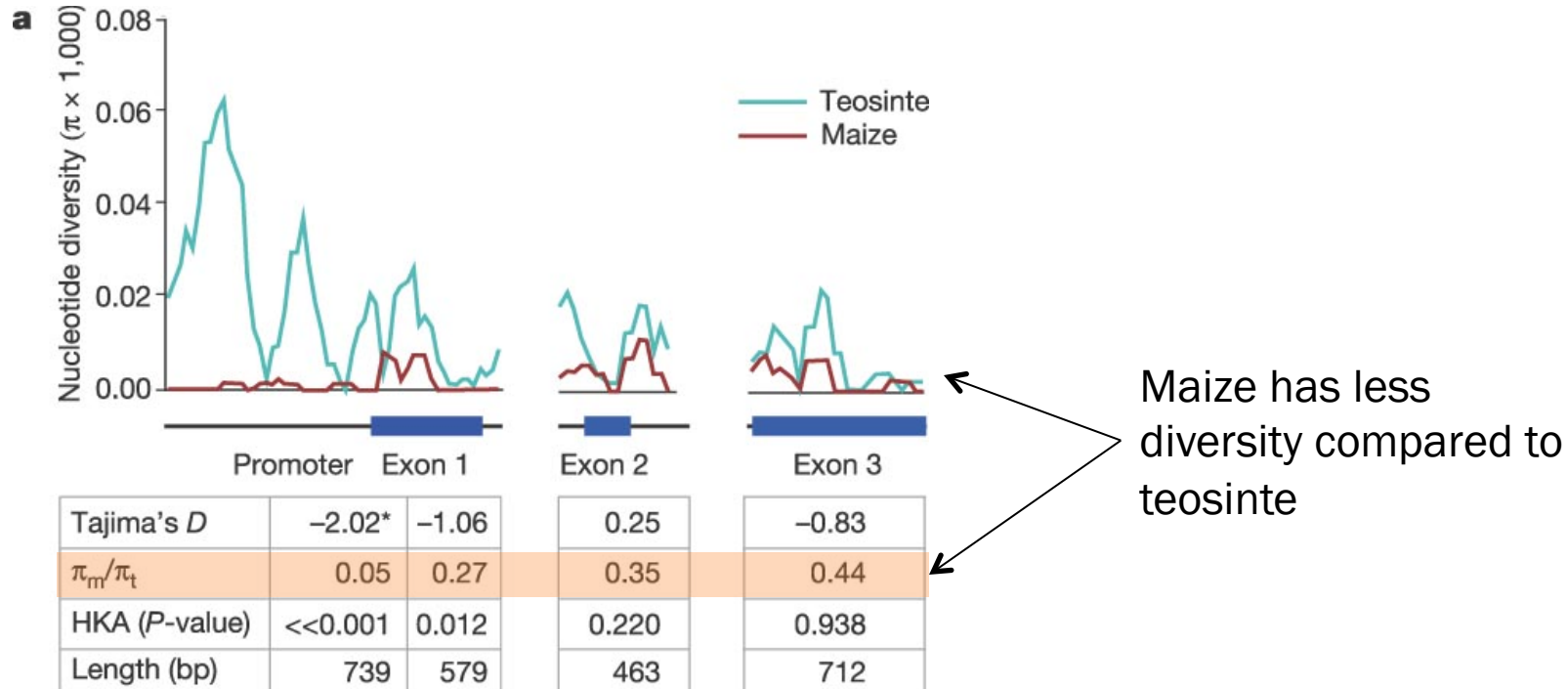
# Maize cupulate fruitcase genetics



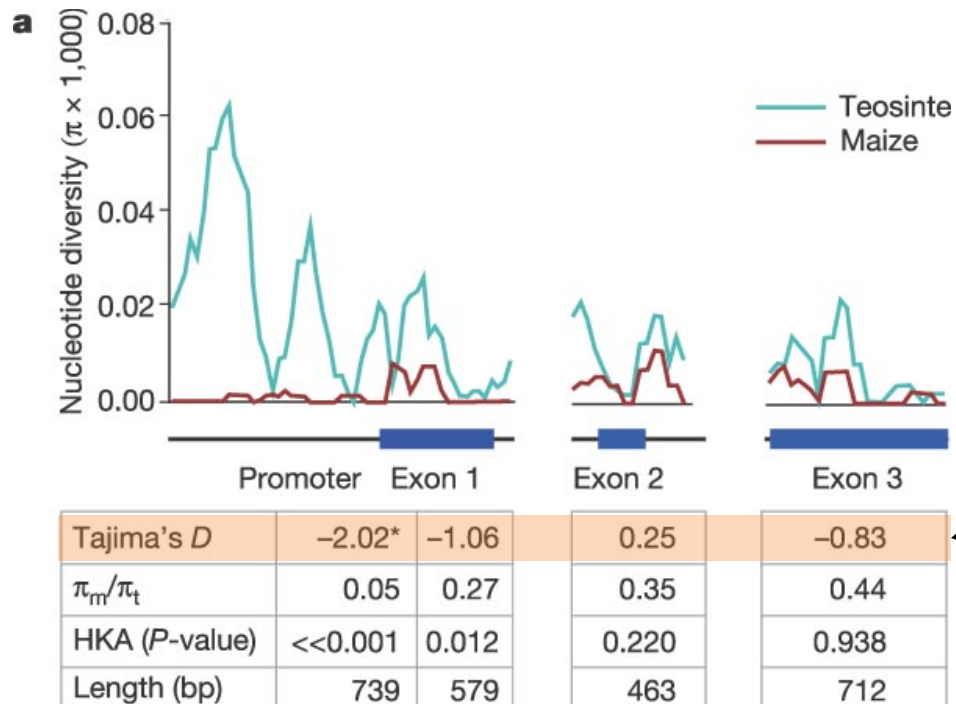
Wildtype teosinte hard fruitcase

Teosinte with maize *tga1* gene

# Maize cupulate fruitcase genetics

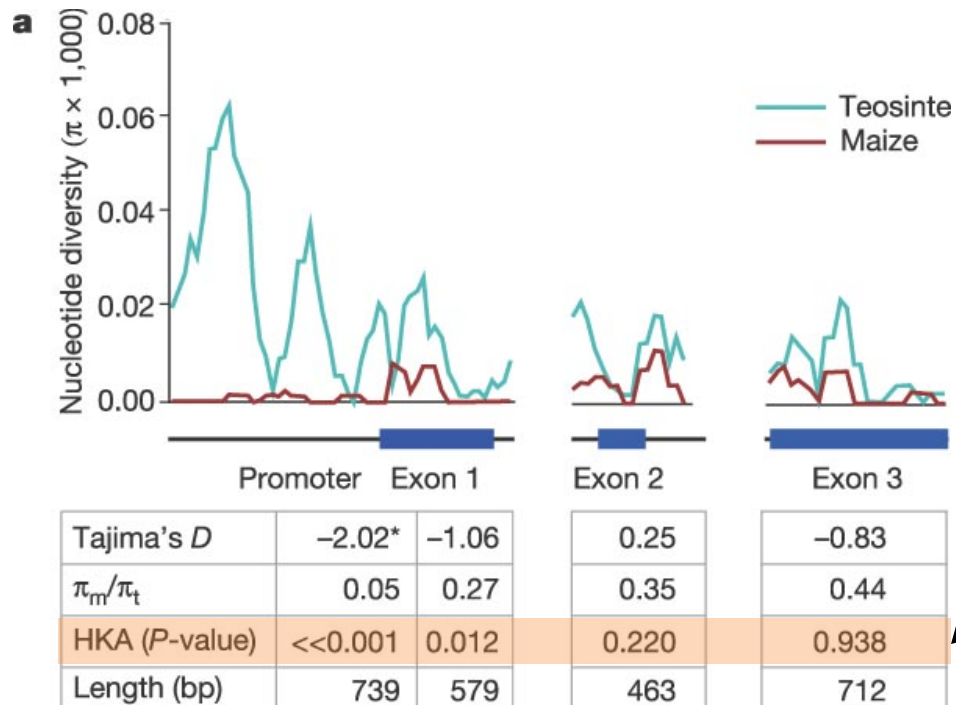


# Maize cupulate fruitcase genetics



Tajima's D looks at site frequency spectrum. Negative values suggests many rare polymorphisms, which occurs during positive selection.

# Maize cupulate fruitcase genetics



HKA asks if there is more divergence between species than would be expected by the amount of polymorphism in the species

# C. Linkage Disequilibrium (LD)

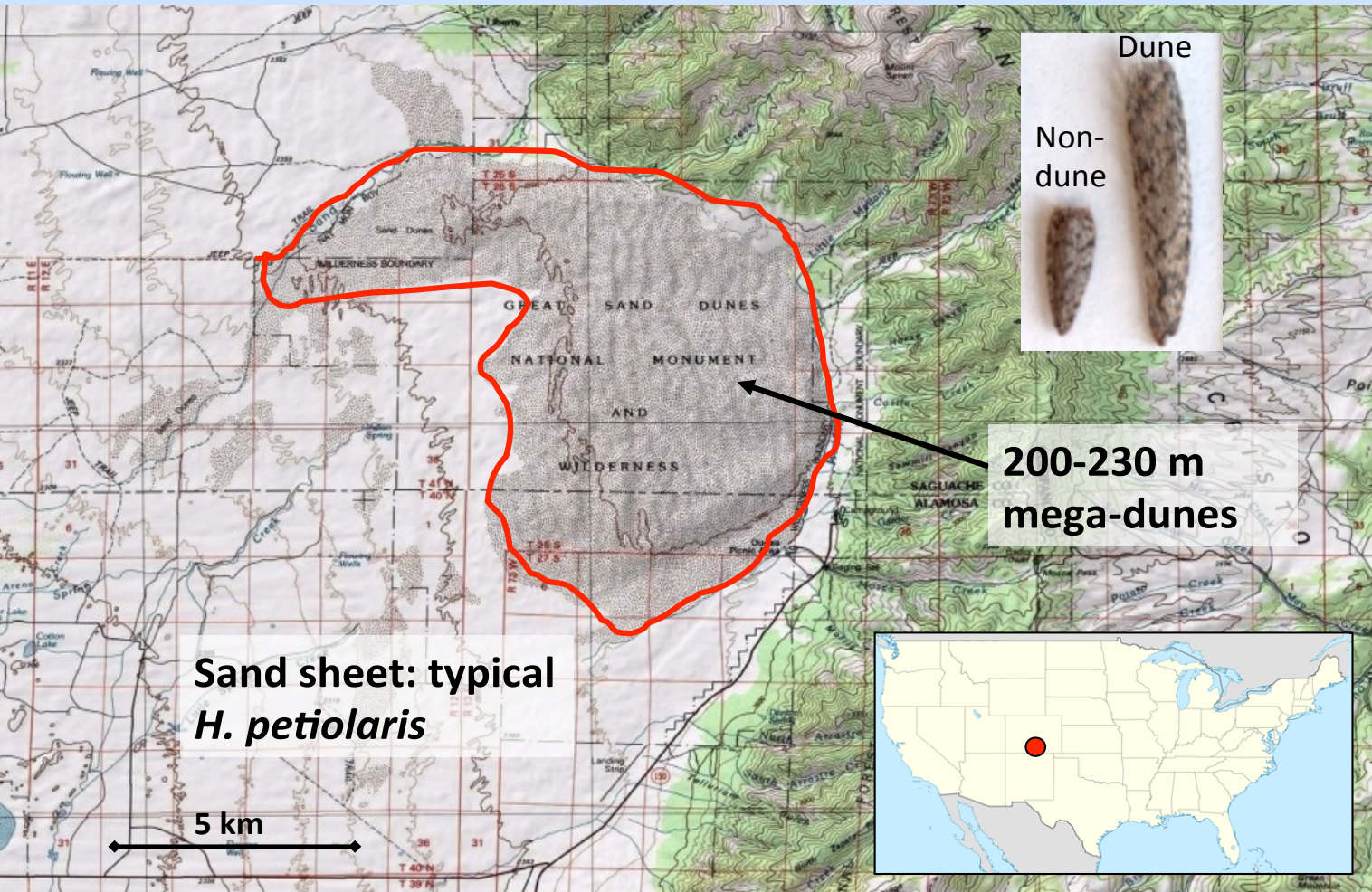
- The nonrandom association of alleles from different loci
- Levels of linkage disequilibrium will increase during selective sweeps
  - As a new mutation rises in frequency, it will drag along linked sites
  - This haplotype block will have high LD until recombination breaks it up over time

# D. Population Differentiation: Lewontin-Krakauer Methods

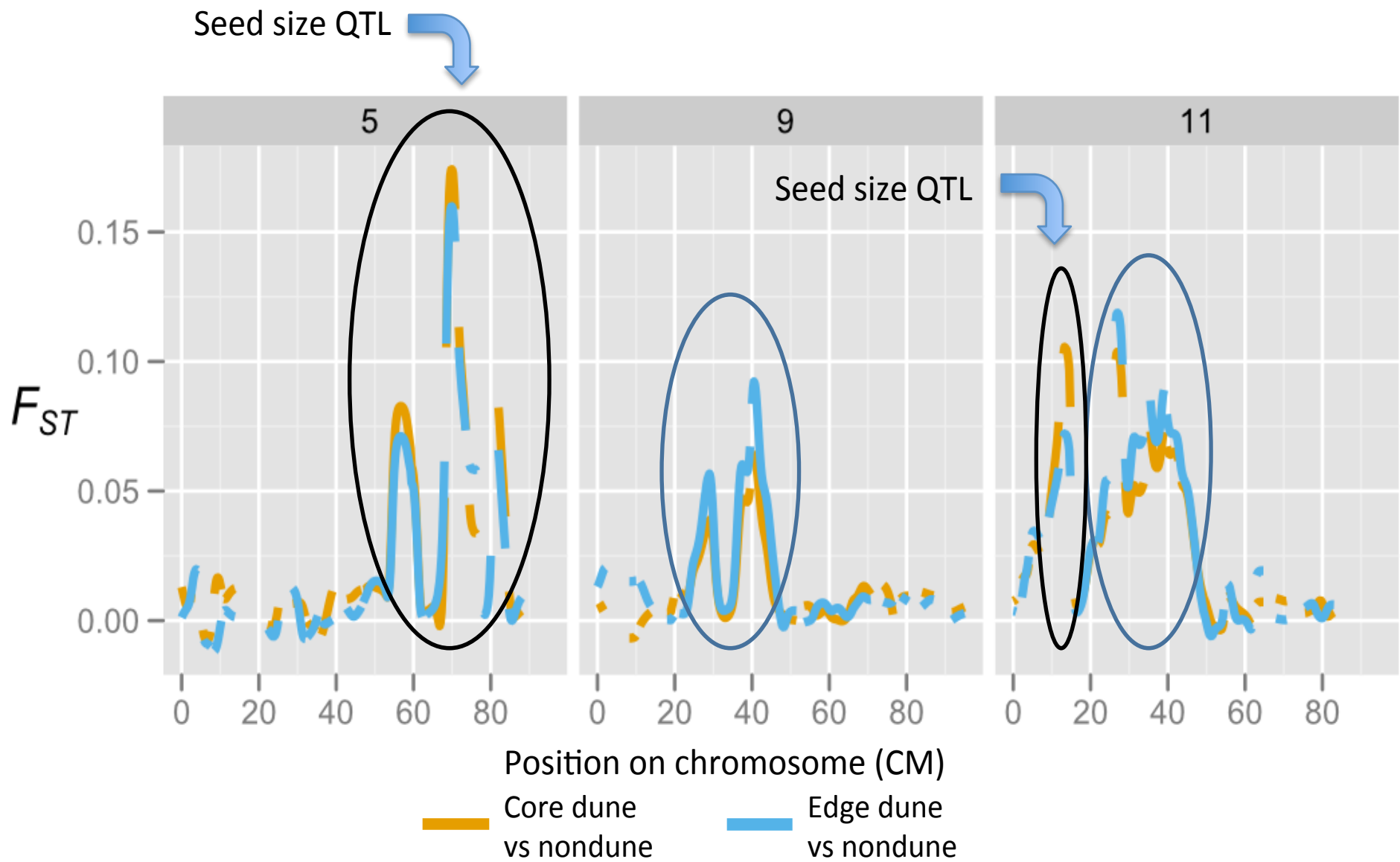
- Selection will often increase the degree of genetic distance between populations
- Compute pairwise genetic distances (e.g.,  $F_{ST}$ ) for many loci between populations
- When a locus shows extraordinary levels of genetic distance relative to other loci, this “outlier” locus is a candidate for positive selection

# Example of Genome Scan Approach in Sunflowers

Study System: Dune Sunflowers from Great Sand Dunes NP, CO



# Four regions of differentiation



# Unanswered questions

- What are the genes that underlie adaptation?
- Is it many genes or a few?
- How repeatable is the genetics of adaptation?
- Do adaptive mutations occur in coding or regulatory regions?
- What is the effect size adaptive alleles?
- What role does linkage play in adaptation?
- De novo mutations or standing genetic variation?