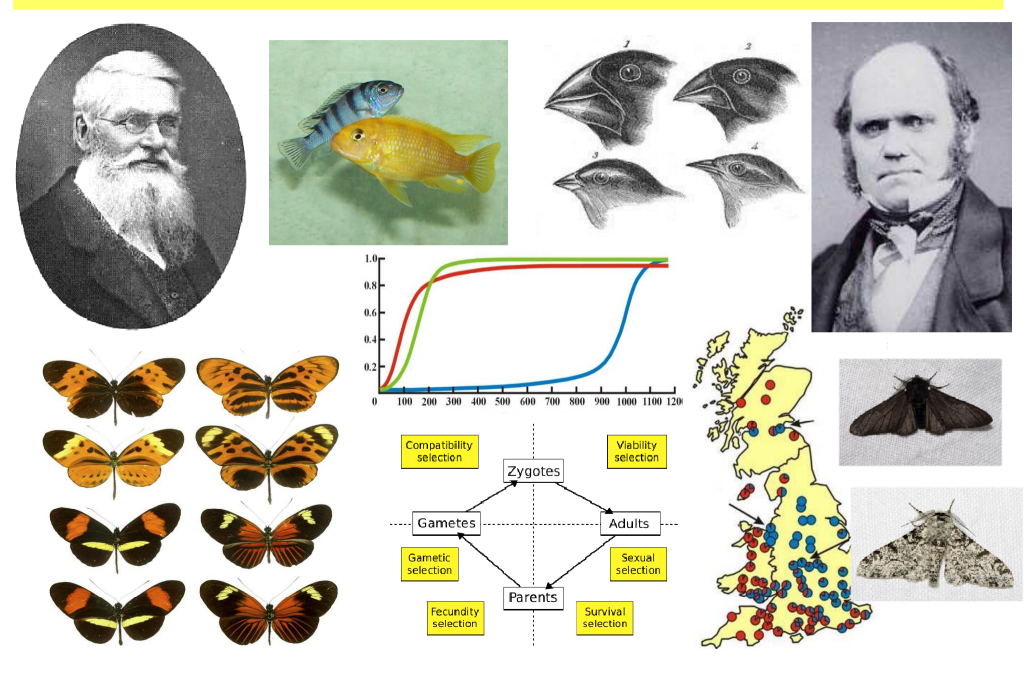
## **NATURAL SELECTION**



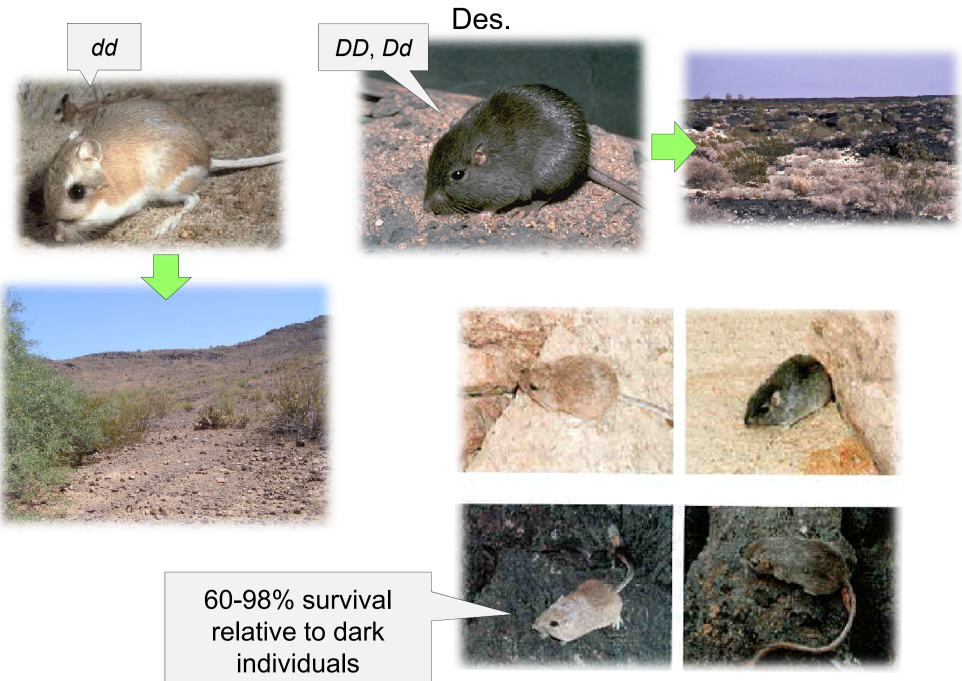
#### Evolution by natural selection:

All organisms produce more offspring than can survive and reproduce.

Individuals (genotypes) differ in heritable traits related to survival and reproduction.

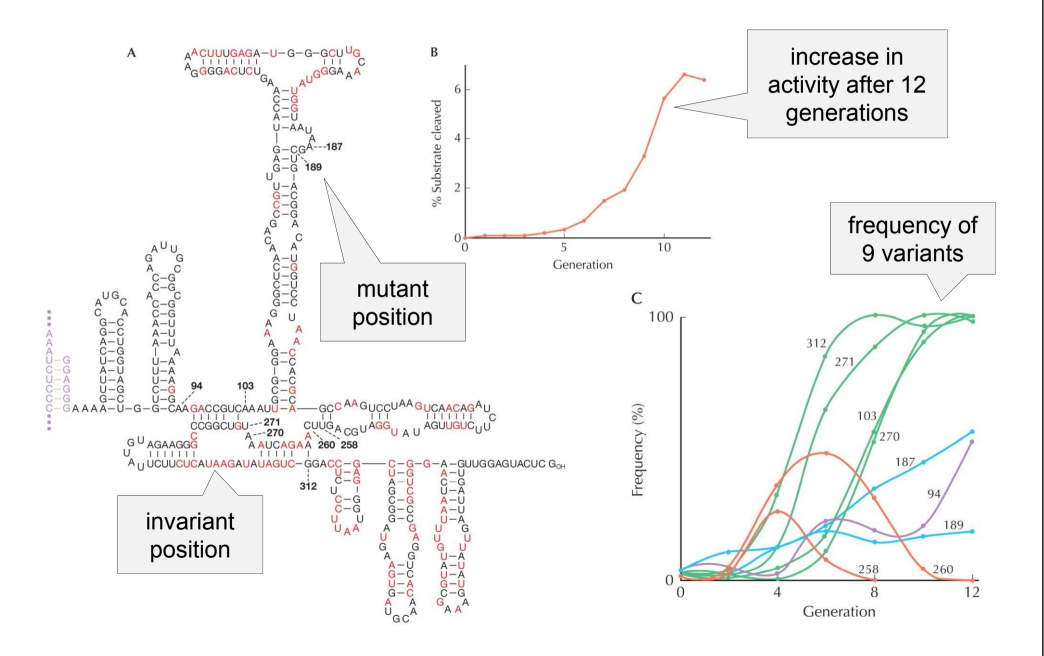
> The genotypes differ in their contribution to the next generation, ie. the most fit genotypes contribute more than the less fit ones.

#### Rock pocket mouse (Chaetodipus intermedius): Sonoran and Chihuahuan



#### Selection on the RNA level:

#### intron *Tetrahymena*: Ca<sup>+</sup> instead of Mg<sup>+</sup> (normal state)



## **REPRODUCTIVE FITNESS,** *w*

= average per capita lifetime contribution of individuals of a given genotype to the population after one or more generation

absolute number of the offspring = absolute fitness

discrete generations, stable population  $\rightarrow$  fitness  $\approx$  1 in asexual organisms,  $\approx$  2 in sexual organisms; even with a slight deviation the population goes either to extinction or to overpopulation

continuous time scale  $\rightarrow$  growth rate  $\approx 0$ 

in evolution relationships between genotypes in a population more important  $\rightarrow$  relative fitness

discrete time  $\rightarrow$  = <u>ratio</u> of absolute fitness; continuous time  $\rightarrow$  = <u>difference</u> between growth rates

usually relative fitness of the most fit genotype = 1 alternatively we may relate to the <u>mean population fitness</u> Components of fitness:

viability

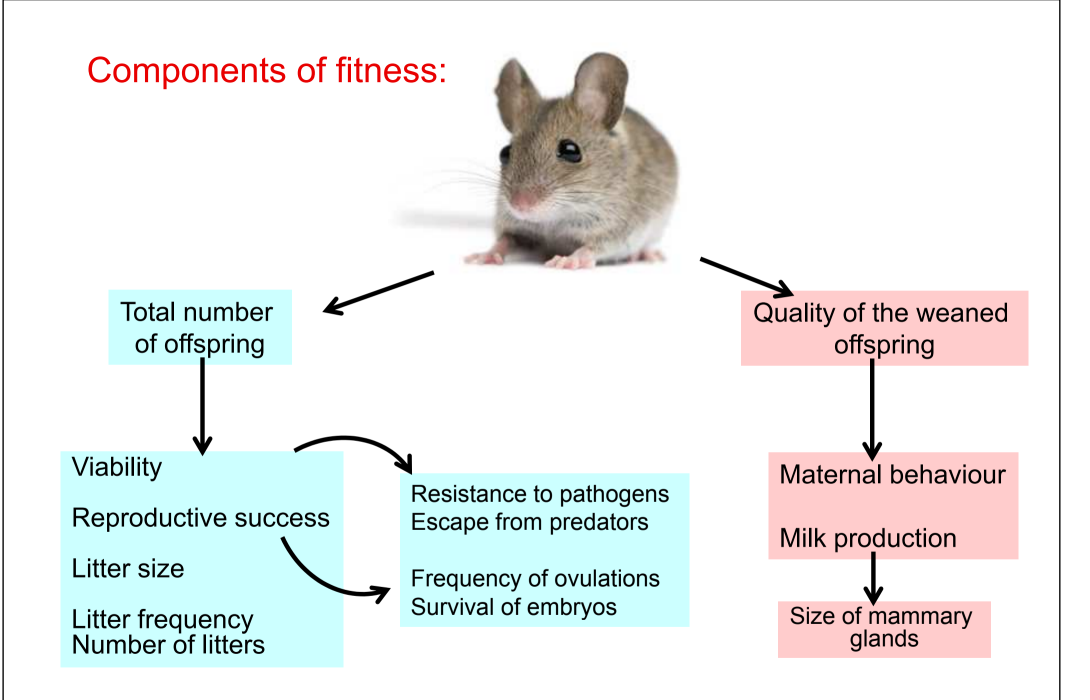
reproductive success

fertility/fecundity









zygotic selection:

viability reproductive success fertility/fecundity

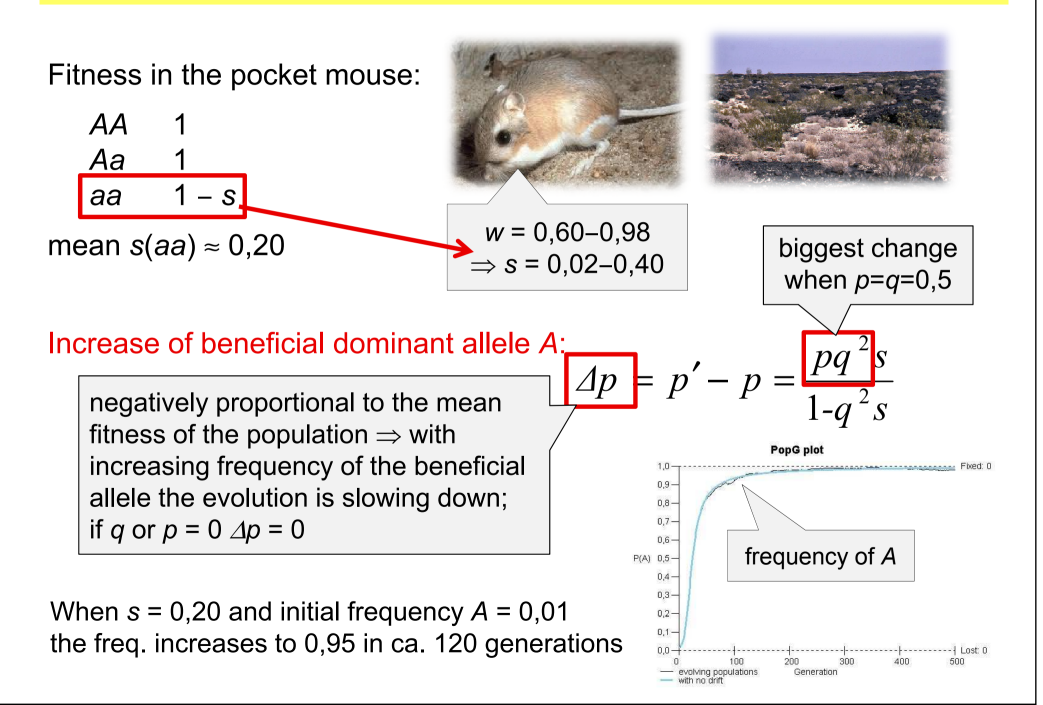
gametic selection:

gamete viability fertilisation success segregation distortion

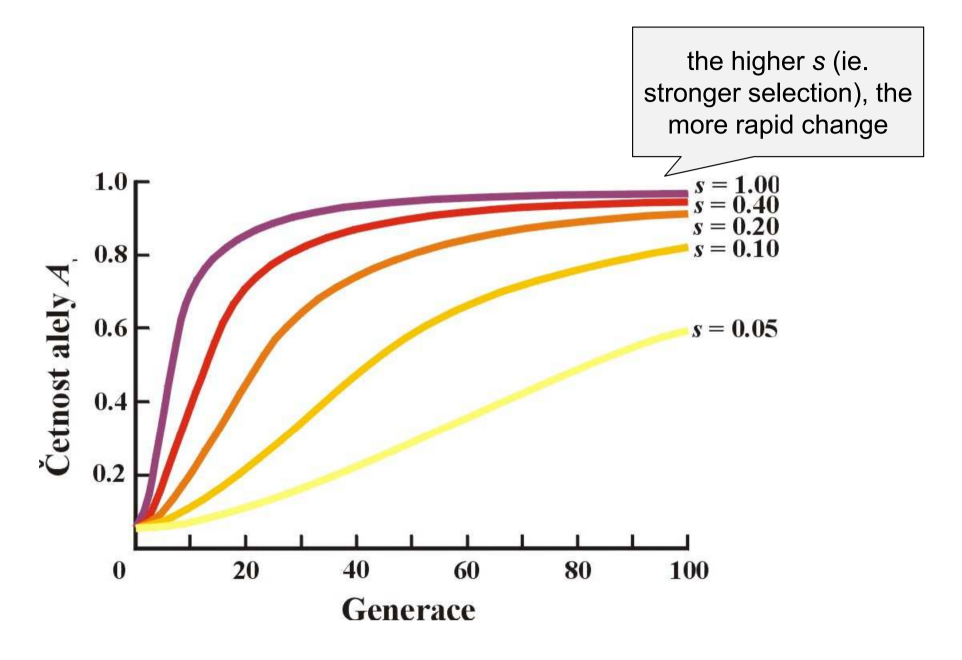


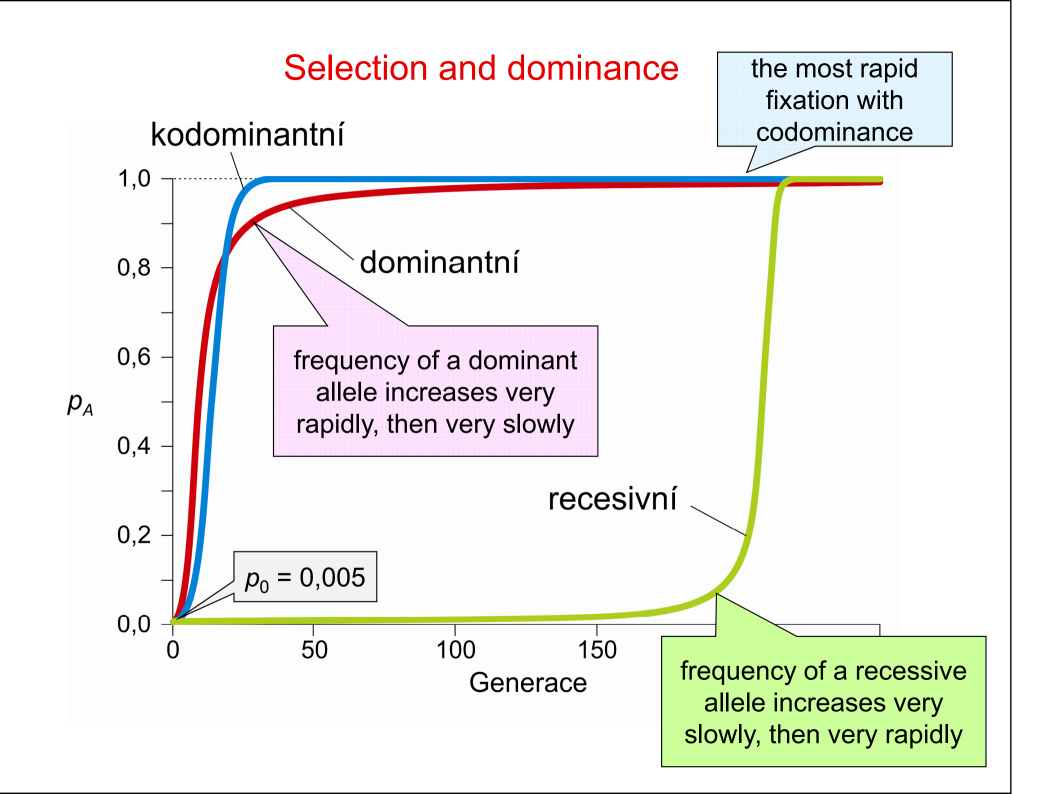


## Change of allele frequencies and selection coefficient, s

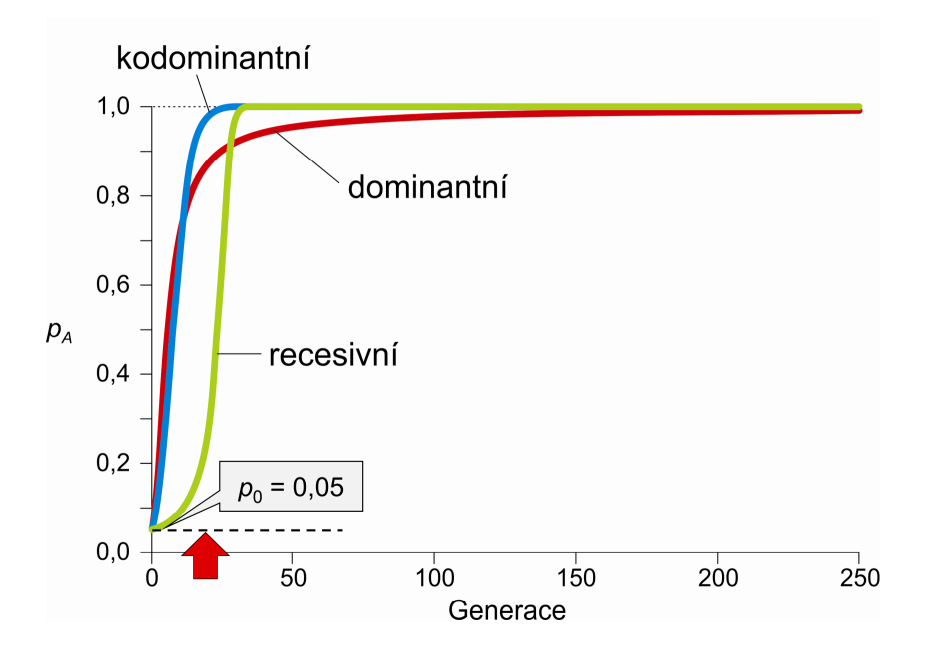


#### Increase of a advantageous dominant allele A:





#### Effect of the initial allele frequency:

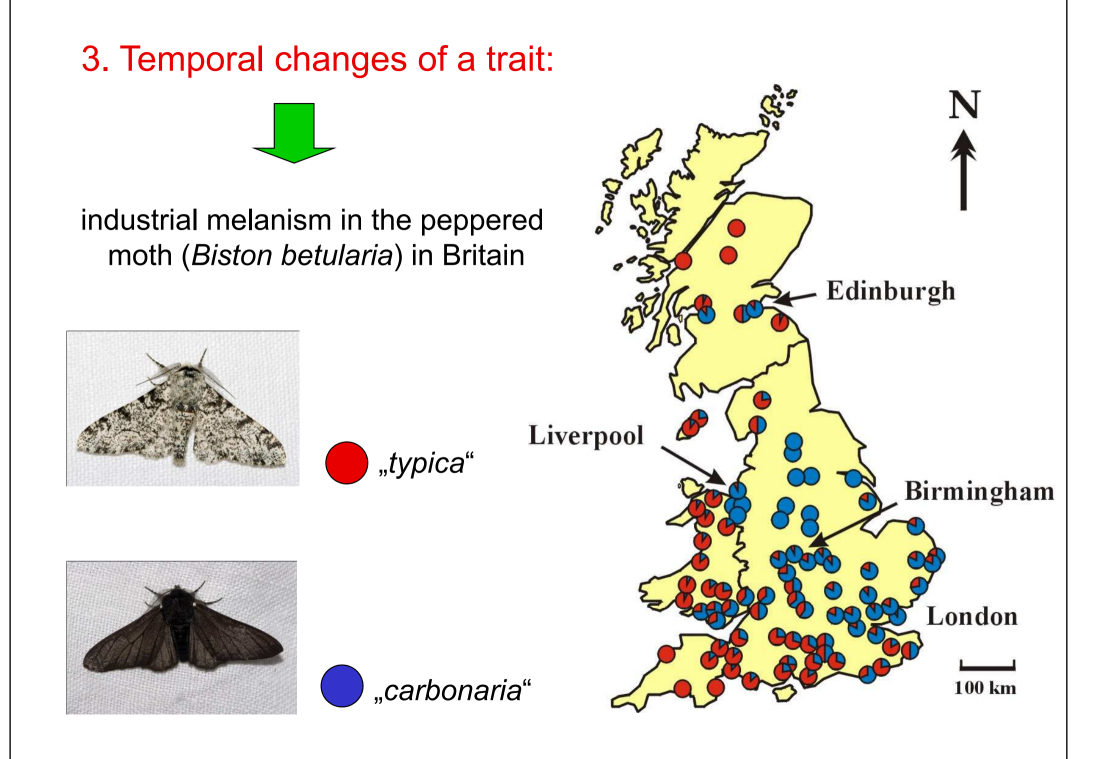


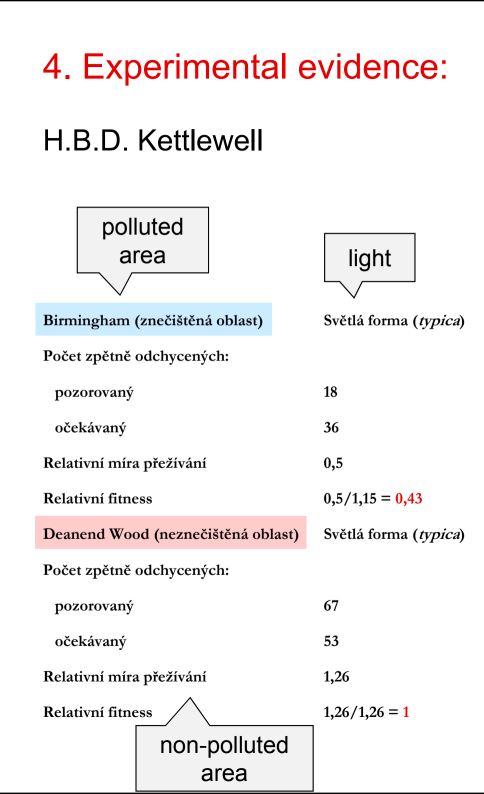
## **STUDY OF NATURAL SELECTION:**

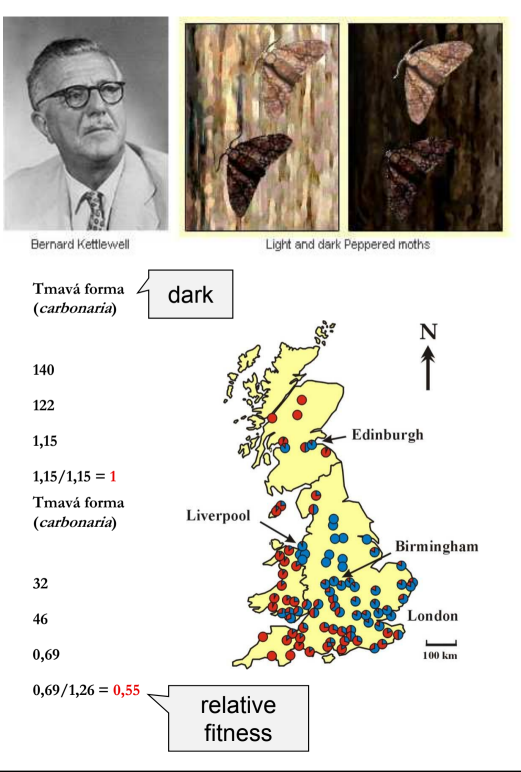
- 1. Deviations from expected genotype frequencies (HW)
- 2. Correlation of allele frequencies across populations

Adh<sup>F</sup> in D. melanogaster





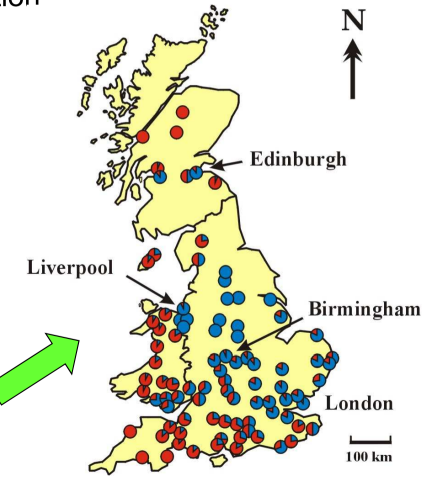




#### Objections and problems:

~90 % of predation by bats × these deaths are non-selective!

in some areas weak correlation between melanism and pollution  $\times$  active and passive migration



**Objections and problems:** 

errors in the experiment?

during the day, peppered moths stay on horizontal branches, not on trunks (different lichen species); in butterflies and birds different perception of UV

increase of melanism also in species not endangered by predation by insectivorous birds (pigeons, cats, some beetles)

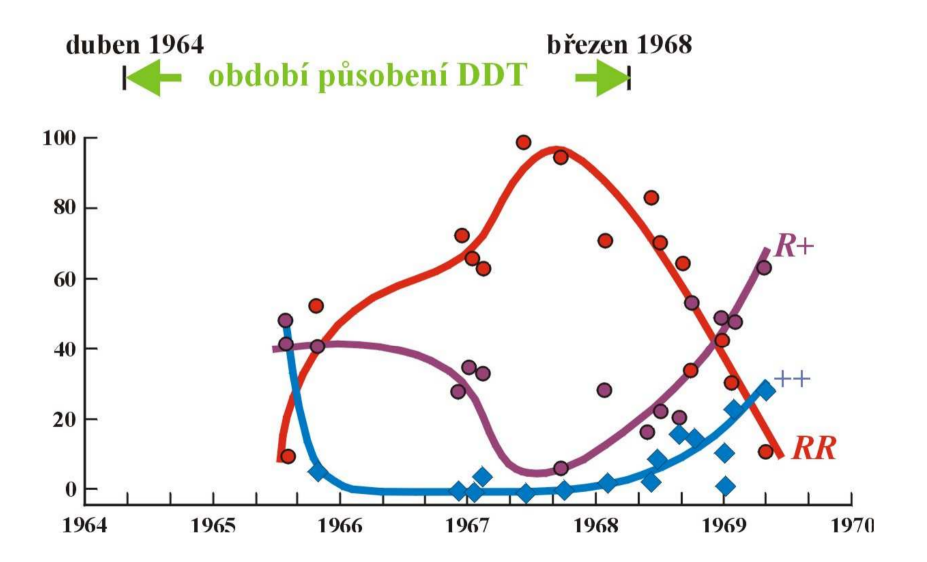
under laboratory conditions the *typica* viability by 30% lower than that of *carbonaria* 

better absorption of solar radiation in melanic forms? (eg. two-spot ladybird)

2000–2012 Michael Majerus: ~5000 moths, corroboration of the hypothesis

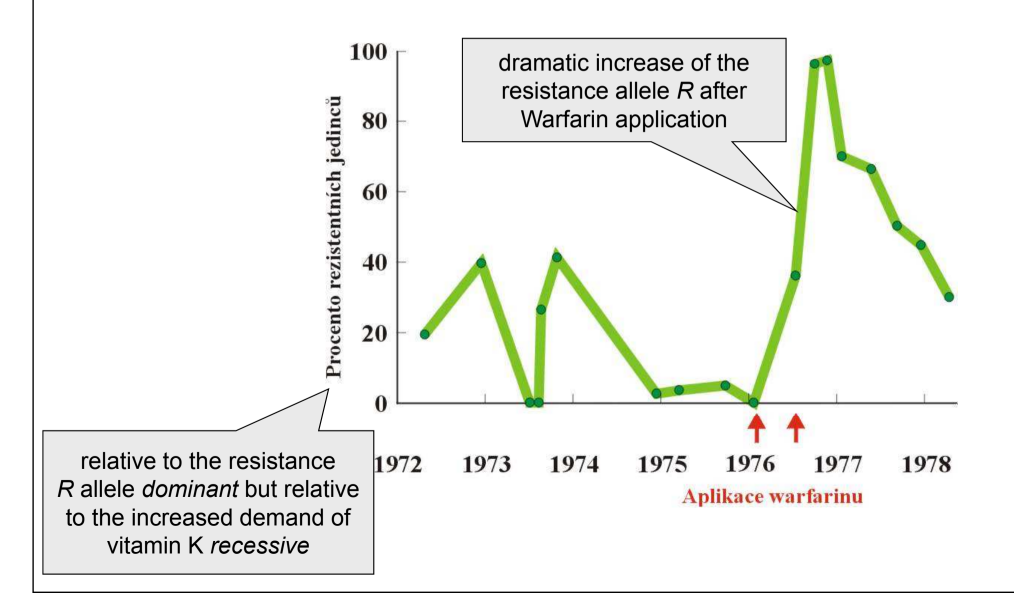
## 5. Resistance

eg.: DDT resistance in mosquitos (Aedes, Anopheles):

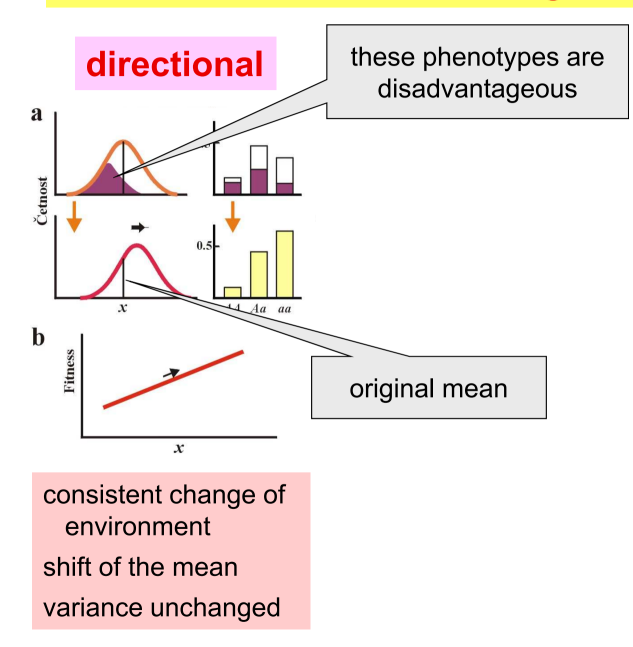


eg.: Warfarin resistance in rats:

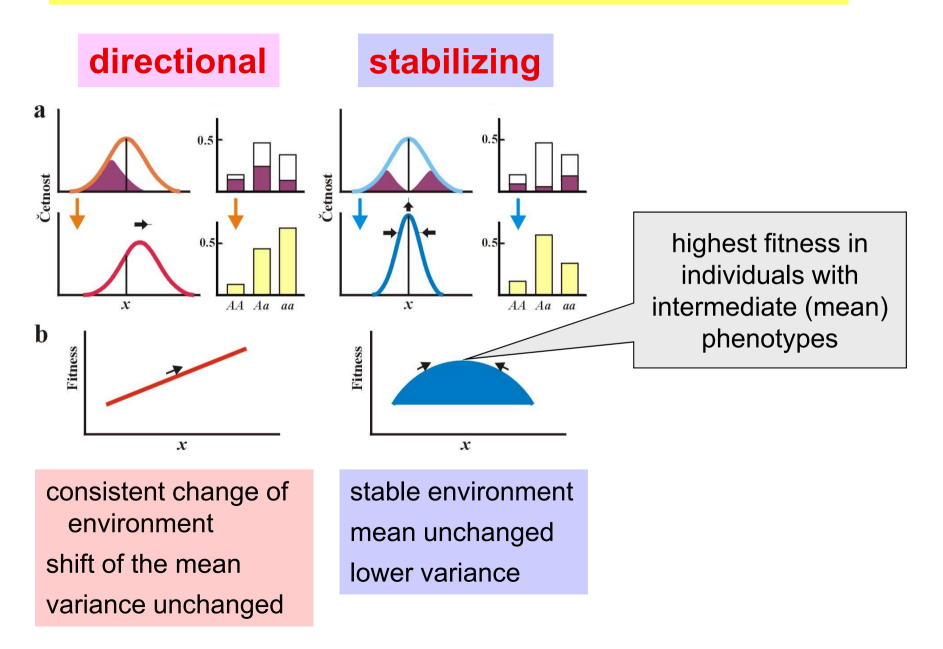
Warfarin = blood anticoagulant, inhibiting the enzyme responsible for the recovery of vitamin K (coagulation cofactor)



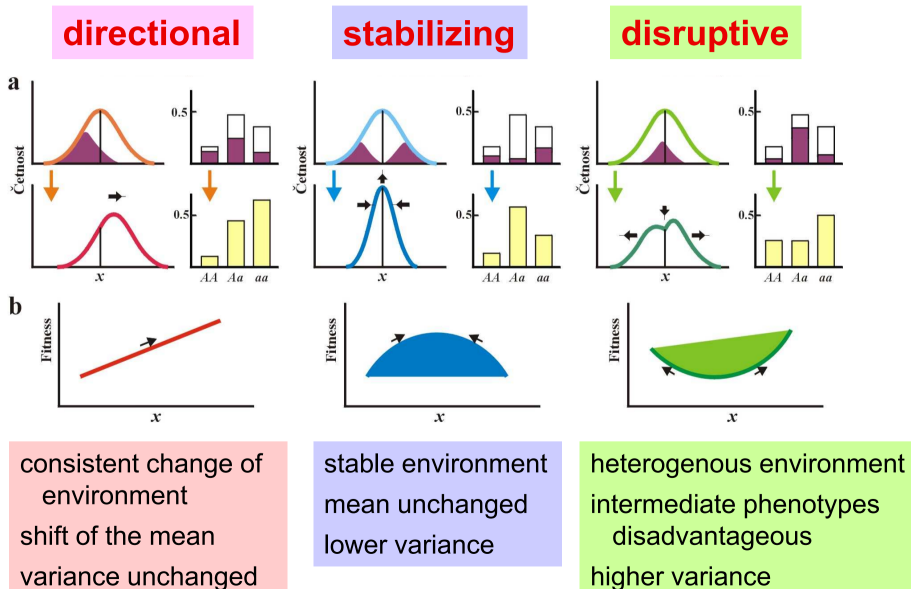
# Relationship between phenotype and fitness: basic selection regimes



## Relationship between phenotype and fitness: basic selection regimes

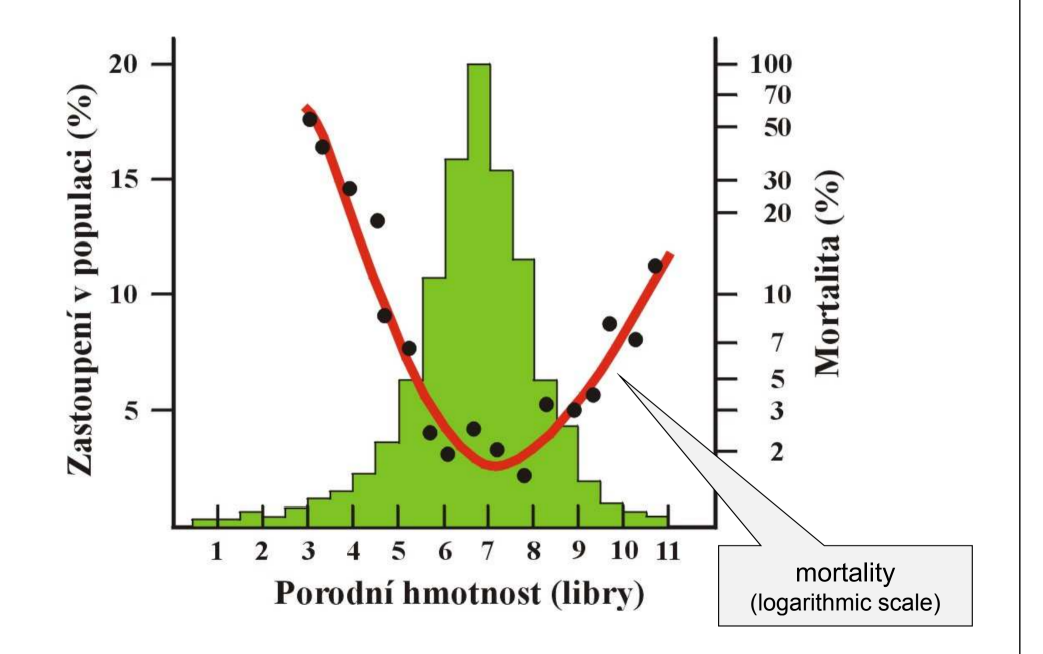


## **Relationship between phenotype and fitness:** basic selection regimes



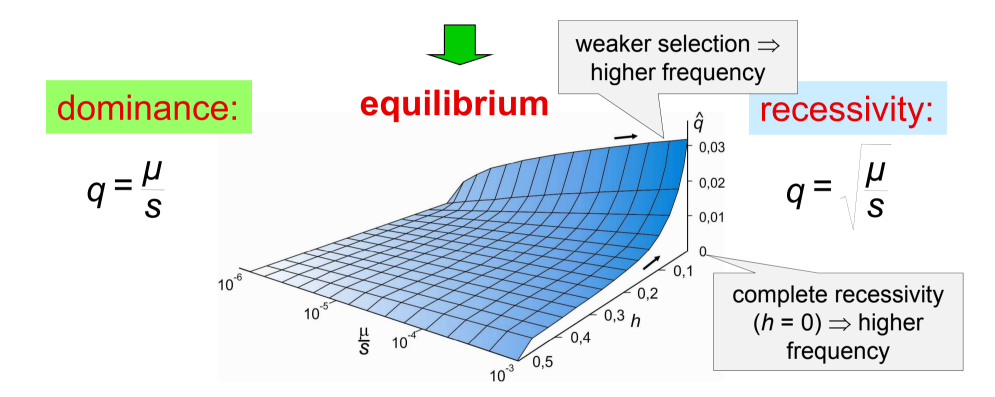
variance unchanged

### stabilizing selection – birth weight in humans



## Equilibrium between selection and mutation

recurrent emergence of a deleterious mutation  $\times$  elimination by selection



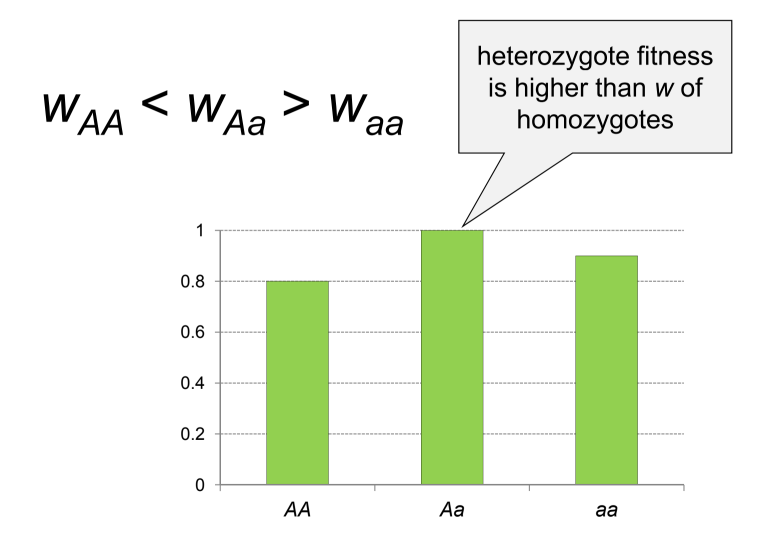
#### Muller-Haldane principle:

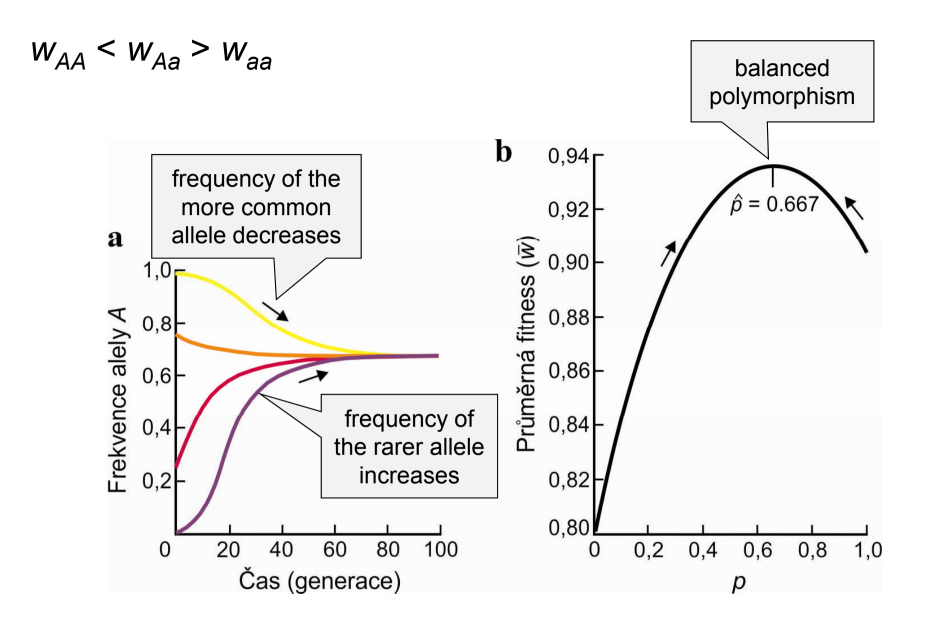
Regardless of dominance/recessivity of a deleterious mutation, its impact on decreasing fitness is independent of the level of its harmfulness.

#### **Equilibrium between selection and gene flow** repeated "influx" of a deleterious allele $\times$ elimination by selection 1. $m > s \Rightarrow$ allele fixation 2. $m < s \Rightarrow$ allele elimination $w_{12}$ intermediary equilibrium 3. $m=s \Rightarrow$ polymorphism 1.0 a m=0Rovnovážná genováč četnosť (q) v každém dému 5. 0 5. (q) v každém dému m=1Fitness, w W m=1m=0 $W_{12}$ higher b m=0m=1Fitness, w m=1m=0východ západ západ východ

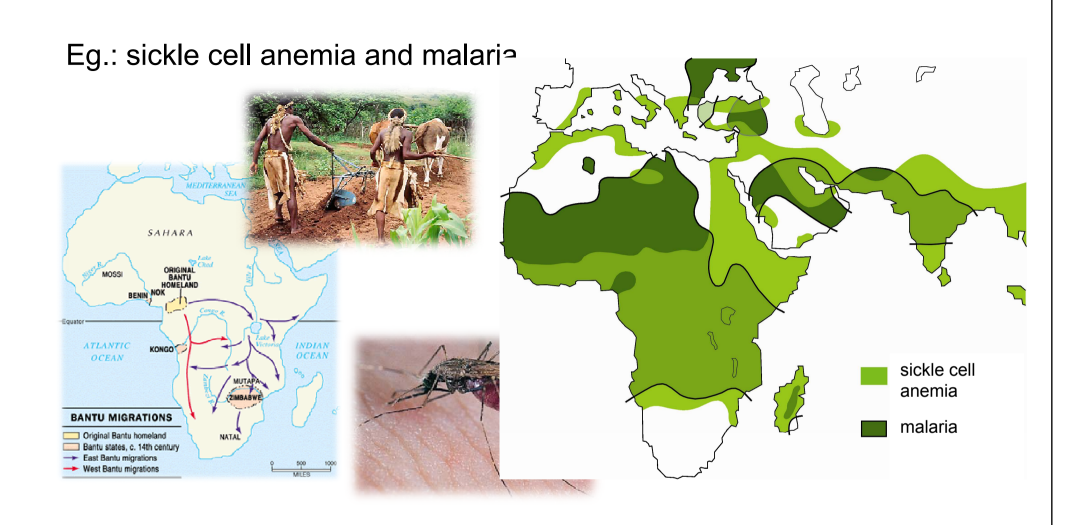


#### 1. Selective advantage of heterozygotes = overdominance





Selection maintains balanced polymorphism



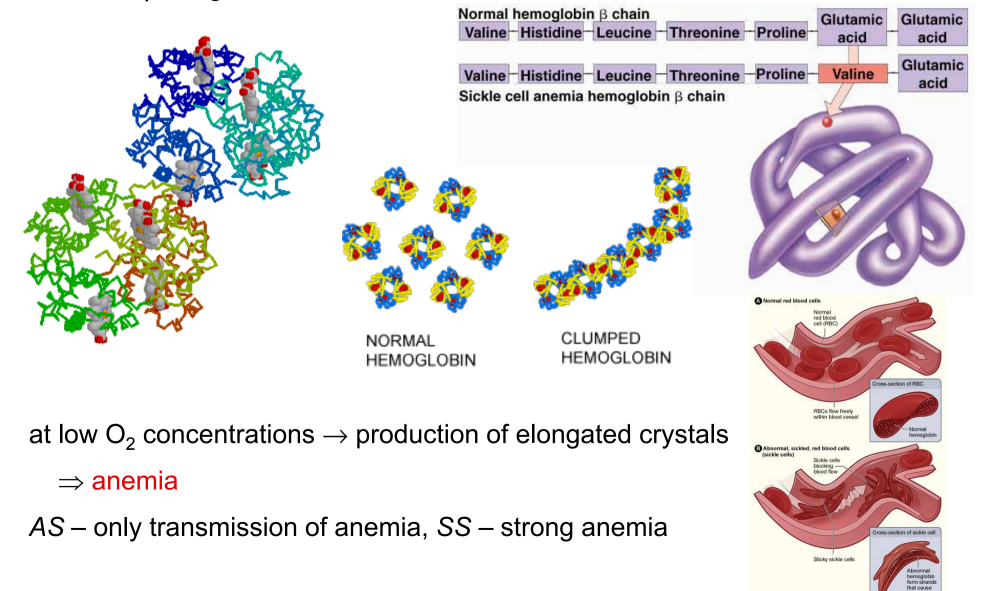
~ 2000 years ago expansion of Bantu peoples

burning off savannas and forests, increase of population density → suitable environment for *Anopheles* mosquitos (*A. gambiae*), the host of *Plasmodium falciparum* 

 $\Rightarrow$  malaria

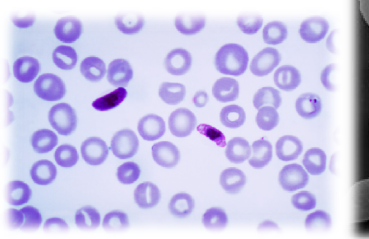
## Sickle cell anemia and malaria:

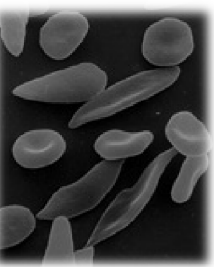
sickle cell anemia: S allele: substitution of 1 AA at 6th position in 6th codon of the  $\beta$ -Hb gene:



sickle-cell red blood cell invaded by *Plasmodium* is rapidly breaking  $\Rightarrow$  the parasite cannot reproduce and multiply  $\Rightarrow$  resistance

 $\rightarrow$  heterozygote advantage

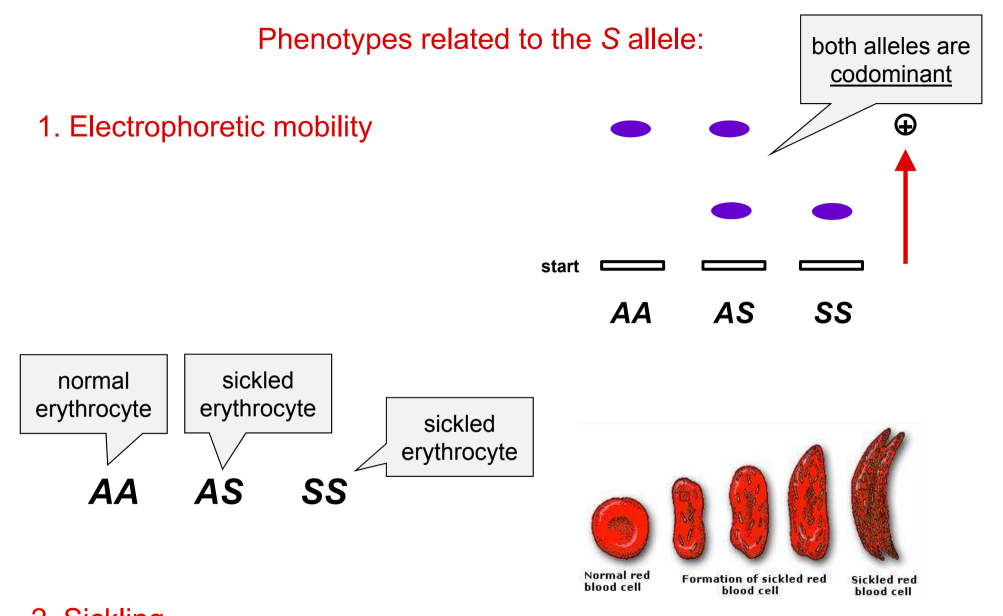




#### Relative fitness of genotypes related to sickle cell anemia:

Table 11.1. Phenotypic Attributes and Relative Fitnesses (Viabilities) of Six Genotypes Formed by *A*, *S*, and *C* Alleles at  $\beta$ -*Hb* Locus in Humans in Wet, Tropical Africa

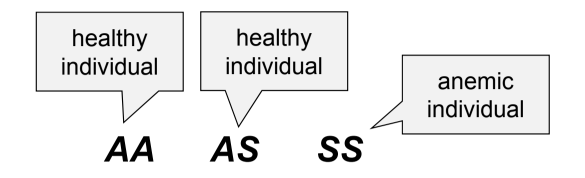
Genotype	Phenotypic Attributes	Fitness in Nonmalarial Environment	Fitness in Malarial Environment
AA	Malarial susceptibility	1.00	0.89
AS	Malarial resistance	1.00	1.00
SS	Hemolytic anemia	0.20	0.20



#### 2. Sickling

sickling in SS and AS individuals  $\Rightarrow$  with respect to deformation S dominant

#### Phenotypes related to the S allele:



#### 3. Anemia

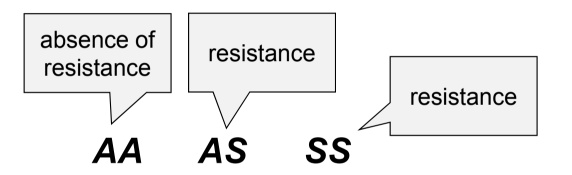
in SS individuals longer chains  $\Rightarrow$  stronger deformation of red blood cells  $\Rightarrow$  more fatal impacts on the organism: erythrocyte rupture (anemia), clogging of capillaries etc.

clinical syndromes only in  $SS \Rightarrow S$  allele <u>recessive</u>

Phenotypes related to the S allele:

4. Resistance to malaria

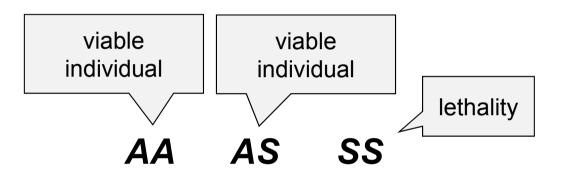
with respect to resistance the S allele dominant



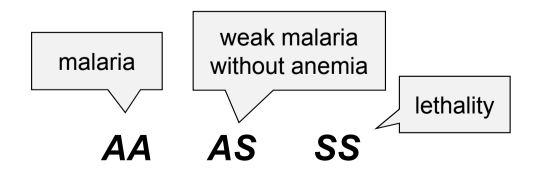
Phenotypes related to the S allele:

5. Phenotype of health (viability)

nonmalarial environment: S recessive



malarial environment: SS – strong anemia; AA – malaria; AS – no anemia, weak malaria  $\Rightarrow$  S is <u>overdominant</u>



Genotype	Phenotypic Attributes	Fitness in Nonmalarial Environment	Fitness in Malarial Environment
AA	Malarial susceptibility	1.00	0.89
AS	Malarial resistance	1.00	1.00
SS	Hemolytic anemia	0.20	0.20
AC	Malarial susceptibility	1.00	0.89
SC	Hemolytic anemia	0.71	0.70
CC	Malarial resistance	1.00	1.31

## Table 11.1. Phenotypic Attributes and Relative Fitnesses (Viabilities) of Six Genotypes Formed by *A*, *S*, and *C* Alleles at $\beta$ -*Hb* Locus in Humans in Wet, Tropical Africa

*Note:* The fitness of the *AS* heterozygote is set to 1. The malarial fitnesses are estimated from data given in Cavalli-Sforza and Bodmer (1971).

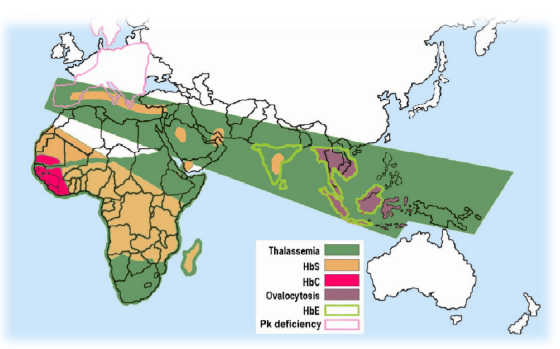
Emergence of *C* allele in the *AS* polymorphism region:

possible genotypes:  $w_{AC} = 0,89$ ;  $w_{SC} = 0,70$ 

 $w_{AS}$  = 1,00  $\Rightarrow$  selection acts against beneficial allele!

Although C higly beneficial, selection will decrease its frequency until it is completely removed!! Resistance against malaria can be mediated through other mechanisms:

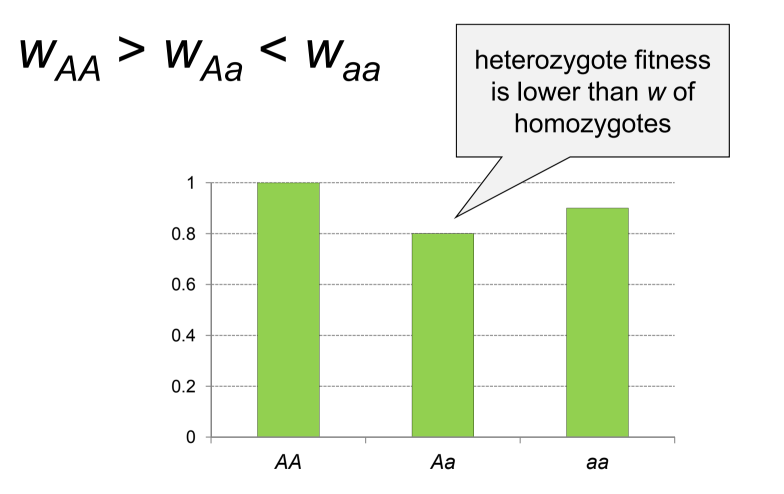
hemoglobin E (JV Asie)  $\alpha$ - a  $\beta$ -thalassemia G6PD<sup>\*)</sup> deficiency Pk<sup>\*\*)</sup> deficiency etc. etc.



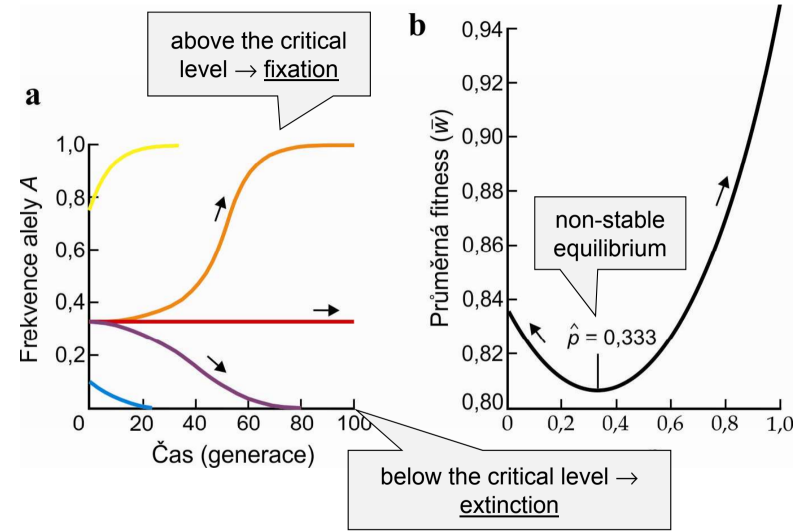
\*) glucose-6-phosphate dehydrogenase\*\*) pyruvate kinase

However, selection in favour of heterozygotes is not widespread in nature

Alternative equilibrium: selection against heterozygotes (underdominance)







Selection results in fixation of one of the alleles (and extinction of the other)

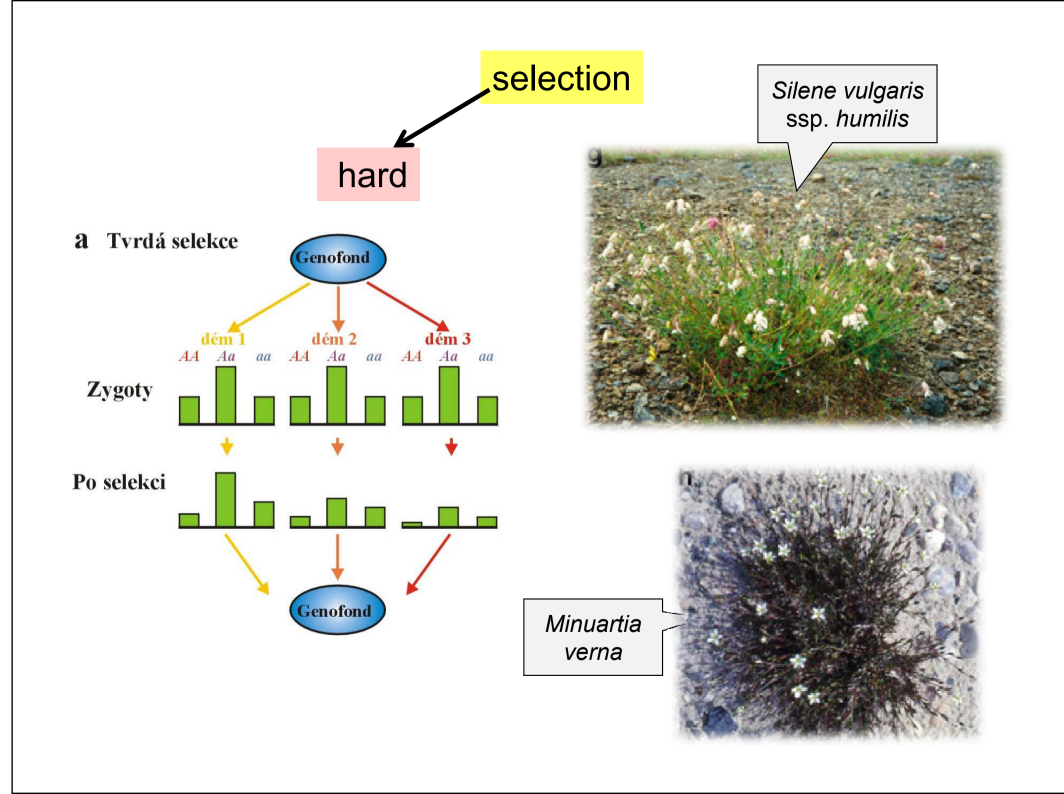
2. Selection in heterogeneous environment

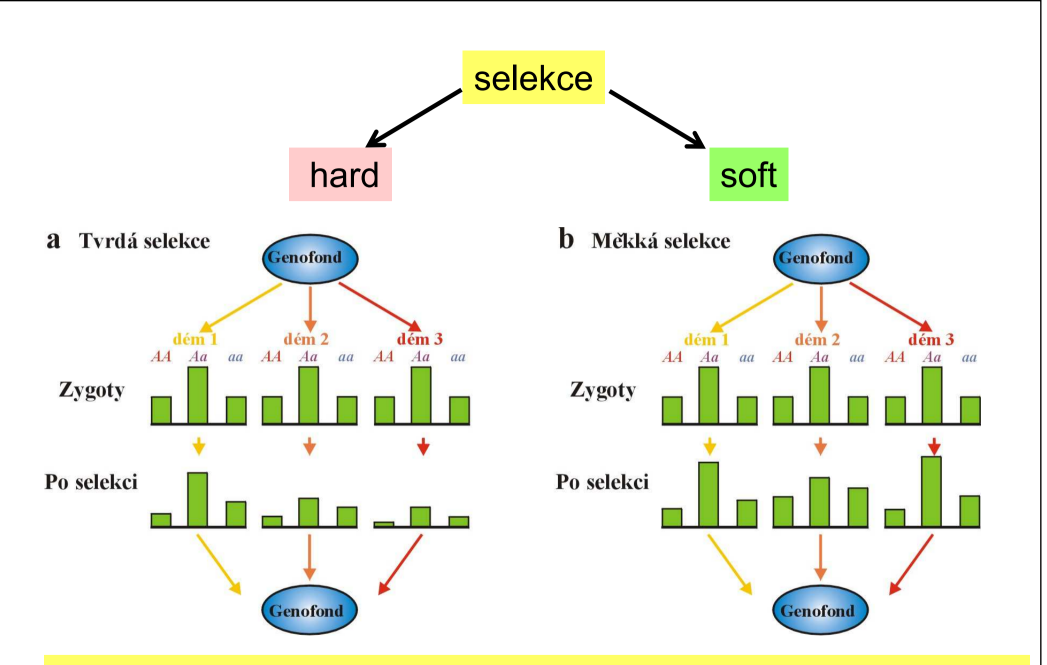
environmental variation: spatial

temporal

coarse-grained: single environment throughout lifetime fine-grained: environmental heterogeneity throughout lifetime

selection: soft hard



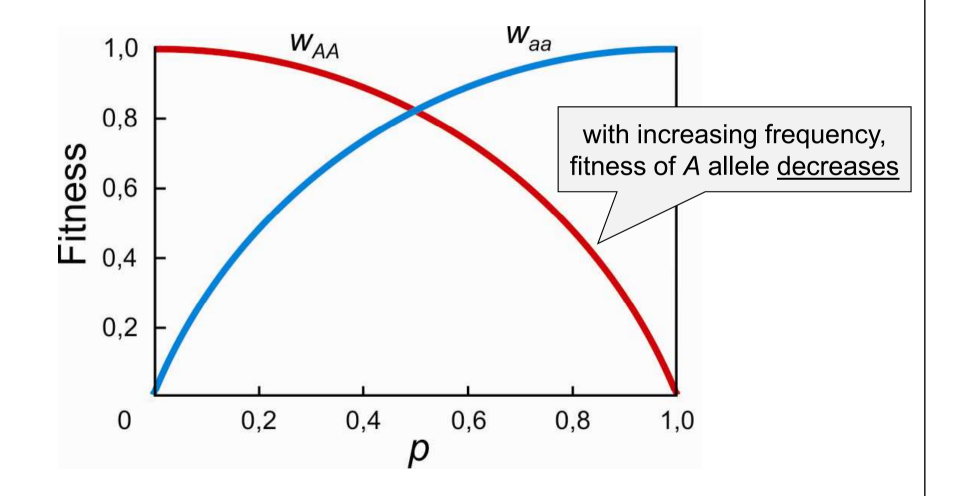


Coarse-grained environment and soft selection will maintain polymophism in the population with higher probability than fine-grained environment and hard selection.

### 3. Antagonistic selection

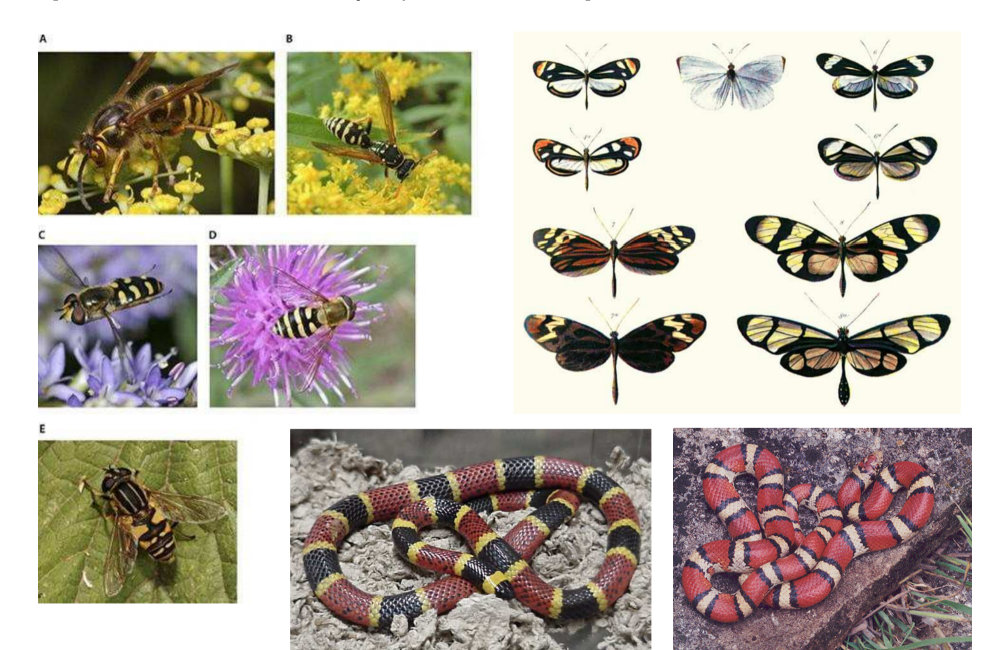
different sexes different ontogenetic stages gametic  $\times$  zygotic phase

## 4. Frequency-dependent selection I. Negative



#### Eg.: Batesian mimicry

[in this case it is rather density-dependent selection]

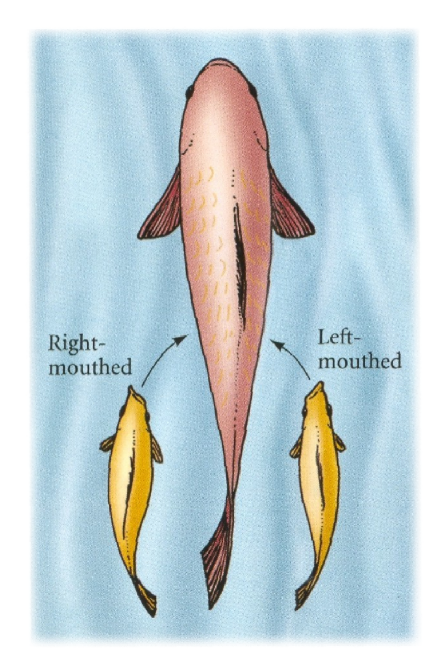


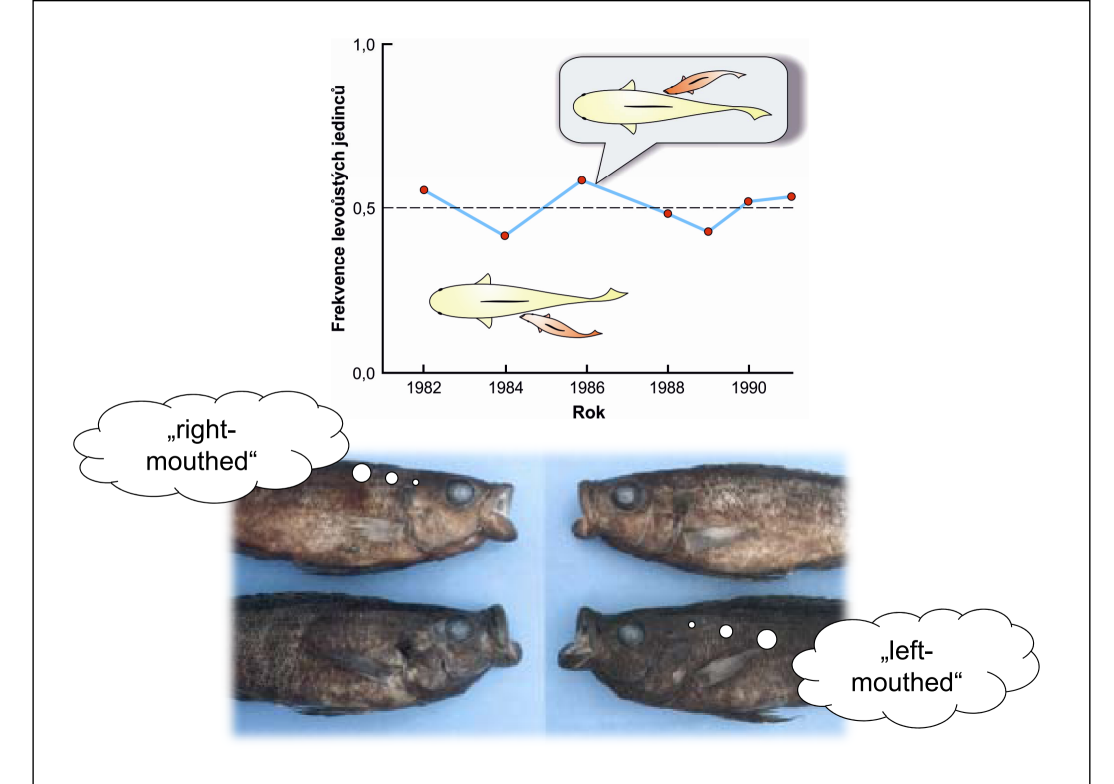
#### Eg.: cichlid Perissodus microlepis (Tanganyika)



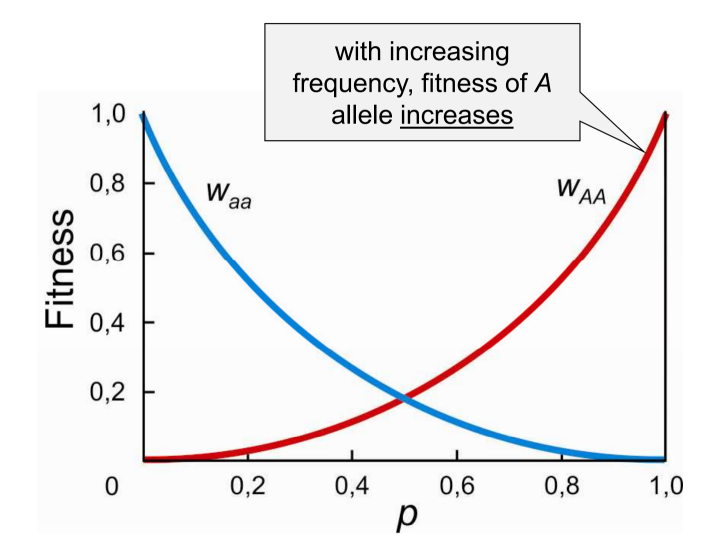


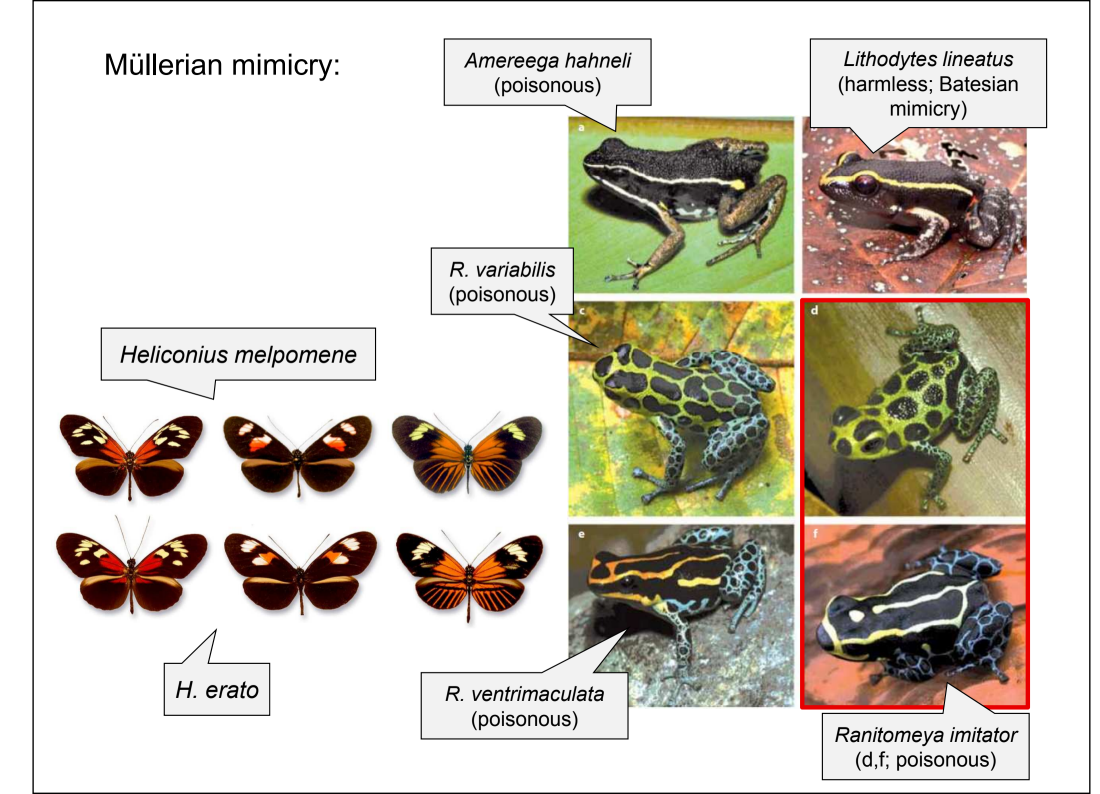
anit uter Street

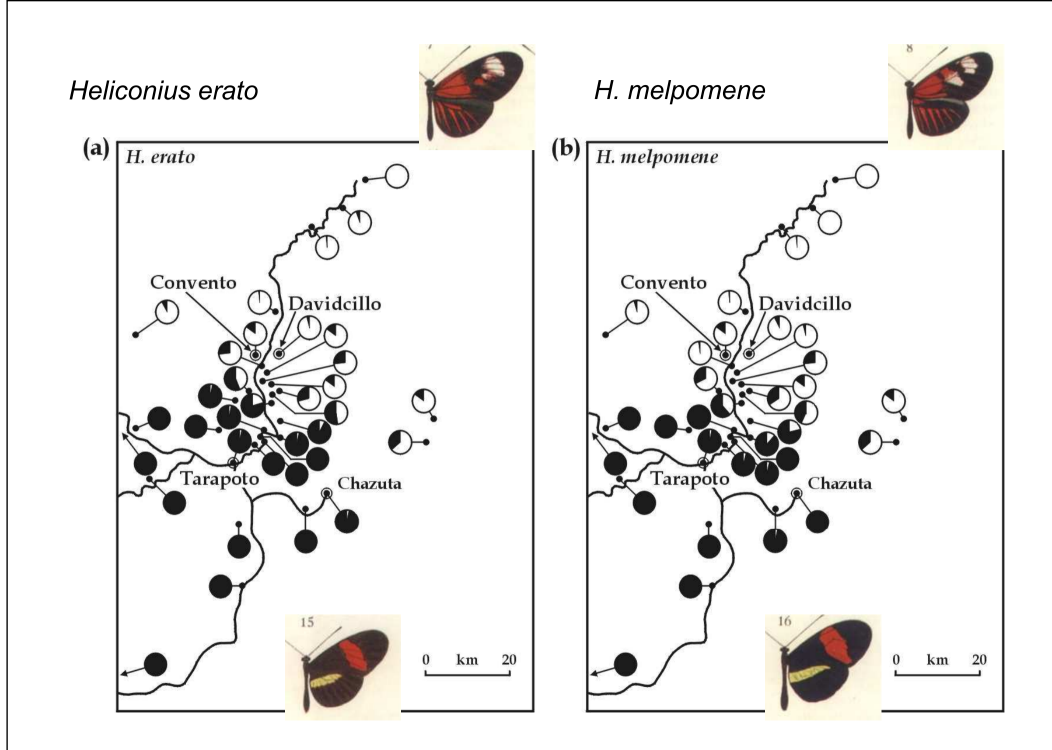




# 4. Frequency-dependent selection II. Positive







#### Balancing selection at the molecular level:

