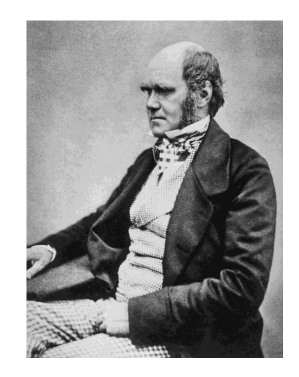
GENETIC AND PHENOTYPIC VARIATION



Evolution as a two-stage process:

- 1. variation among individuals in a population
- 2. changes in the proportion of variants from generation to generation

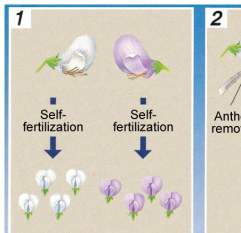


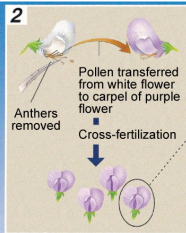


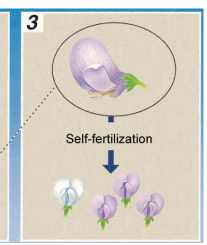
R.A. Fisher

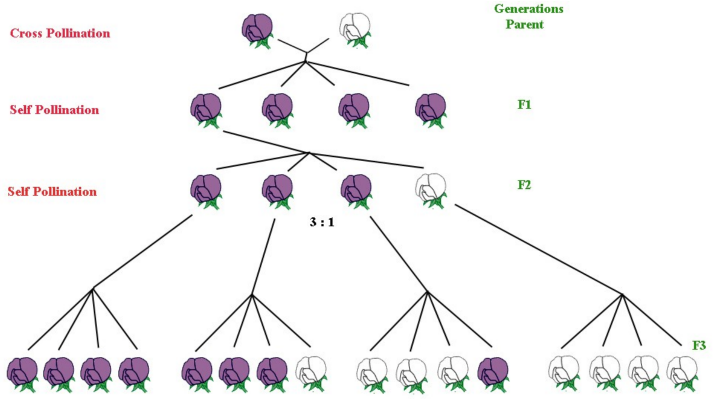
The increase in mean fitness due to natural selection is proportional to the additive genetic variance in fitness.



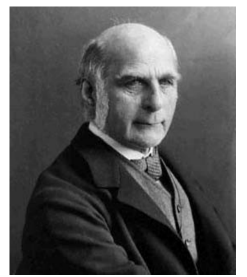




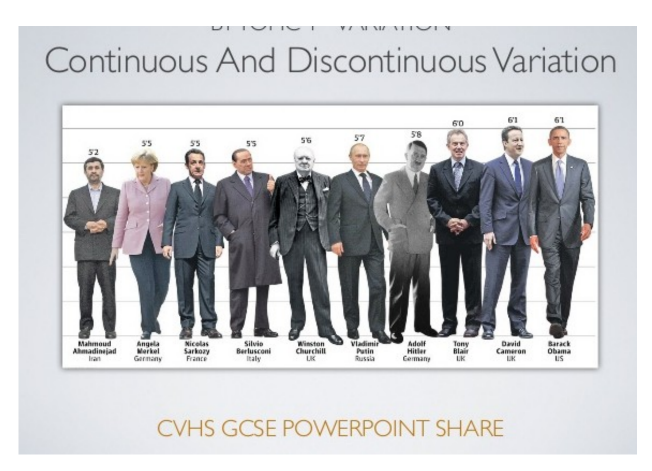




3:1 3:1



F. Galton



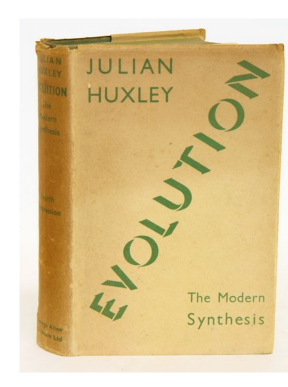
Biometricians: continual variation

many genes

often strong influence of environment

Sources of phenotypic variation:

differences in <u>genotype</u>
differences in <u>environmental conditions</u>
<u>maternal</u> influences (paternal influences)

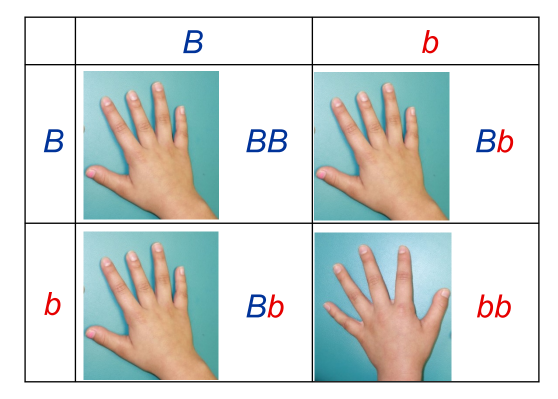


Paradox:

for evolutionary biologists important to study phenotypes for geneticists easier to directly study molecules



Reginald C. Punnett: brachydactyly

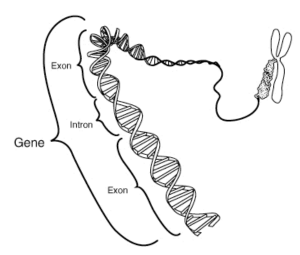




George Udny Yule

Why don t we observe the 3:1 ratio in *populations*?





gene ... till now difficult to define/delimit

locus ... here = gene or any other molecular trait

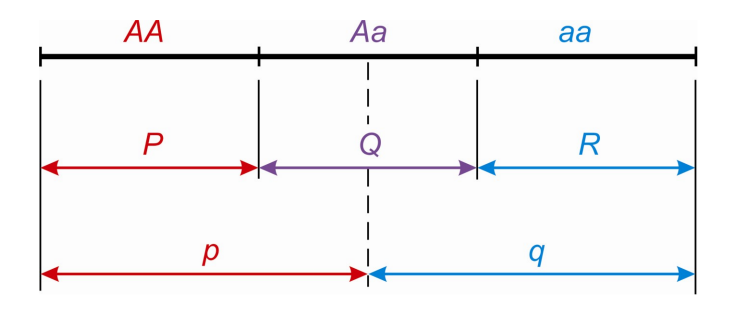
alleles = alternative forms of genes (now broader meaning – segment of DNA)

genome = set of all genes of an individual (nuclear, mitochondrial...)

genotype = set of alelles of one or more genes of an individual

haplotype (haploid genotype) = combination of alelles inherited together

Genotype and allele frequencies



Frequencies: genotype: $P(f_{AA})$, $Q(f_{Aa})$, $R(f_{aa})$

allele (gene): p(A), q(a)

$$P + Q + R = 1$$
$$p + q = 1$$

Evolution takes place in populations...

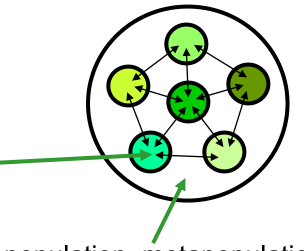
T. Dobzhansky, E. Mayr:

population as a shared gene pool

≈ set of shared alelles or gametes

local populations (subpopulations, demes)

Metapopulation structure Patchy Highly connected Classical Mainland-island Patch isolation Occupied habitat patches Vacant habitat patches Boundaries of populations Nonequilibrium Dispersal Highly Boundary of metapopulation isolated All small Small and large Patch size (Based on Harrison and Taylor 1997; Stith et al. 1996)



global population, metapopulation

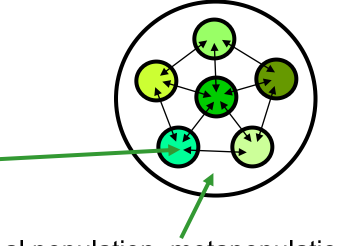
Evolution takes place in populations...

T. Dobzhansky, E. Mayr:

population as a shared gene pool

≈ set of shared alelles or gametes

local populations (subpopulations, demes)



global population, metapopulation

Local populations also share a system of mating

populations natural, experimental, agricultural, model

Model populations – Hardy-Weinberg population

Characteristics:

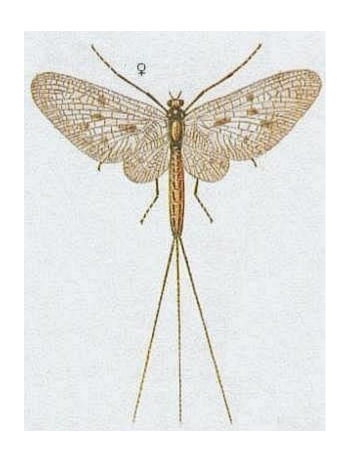
diploid

sexual reproduction

discrete generations

2 alleles, "fair" segregation 1:1

same frequencies of alleles in both sexes



Model populations – Hardy-Weinberg population

Characteristics:

random mating (panmixis)

non-random: assortative mating, inbreeding

very <u>large</u> (effectively infinite) <u>size</u>

no gene flow

no mutation

no selection

Why don t we observe the Mendelian ratios in nature?



R. C. Punnett



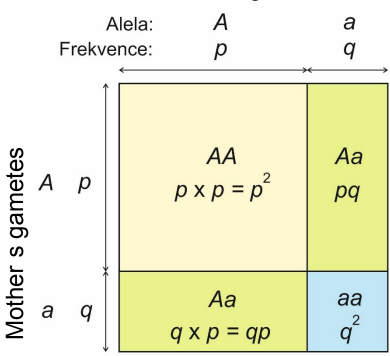




Godfrey Harold Hardy

HARDY-WEINBERG PRINCIPLE

Father s gametes





Godfrey Harold Hardy (1877-1947)

Genotype frequencies in zygotes:

$$f'_{AA} = p^{2}$$

$$f'_{Aa} = pq + qp = 2pq$$

$$f'_{aa} = q^{2}$$

$$p^{2} + 2pq + q^{2} = 1$$

Wilhelm Weinberg (1862-1937)



HARDY-WEINBERG PRINCIPLE

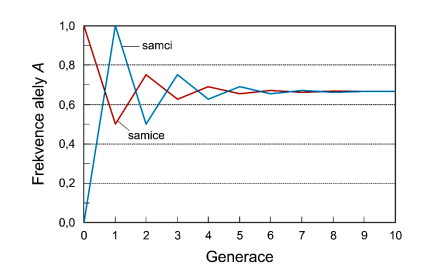
- 1. Alelle frequencies stable across generations
 - = Hardy-Weinberg equilibrium (HWE)
- 2. HWE achieved within a single generation of random mating

Generalization:

X-linked genes:

females: $p^2 + 2pq + q^2$

males: p + q

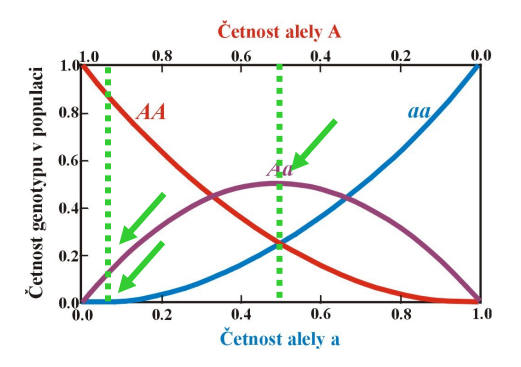


more than 2 alleles:

3 alleles: $p^2 + q^2 + r^2 + 2pq + 2pr + 2qr$

in general $p_i^2 + 2p_{ii}$

Frekvencies of rare alleles



heterozygotes most frequent when p = q = 0.5

 f_{Aa} decreases with 2pq f_{aa} decreases with $q^2 \Rightarrow f_{Aa}/f_{aa}$ increases \rightarrow rare allele "hidden" for selection in heterozygous state

Possible causes of HWE violation:

Methodic causes:

null alleles, allelic dropout

Violation of some of the assumptions of the H-W population:

Heterozygote deficiency:

selection against heterozygotes nonrandom mating (inbreeding, assortative mating) structured populations (different allele frequencies, cf. Wahlund effect)

Heterozygote excess:

selection in favour of heterozygotes nonrandom mating (outbreeding, negative assortative mating) migration mutation

GENETIC VARIATION IN POPULATIONS

Methods of the study of genetic variation:

protein electrophoresis analysis of restriction fragments (Southern blotting, RFLP, DNA fingerprinting) PCR, sequencing, NGS, microsatellites ... Polymorfism and polytypy

Polymorfism:

```
proportion of polymorphic loci (P)
```

sample size usually finite ⇒

limit 5% (
$$P_{0.05}$$
) or 1% ($P_{0.01}$)

number of alleles per locus (A; allele diversity, allele richness)

mean observed heterozygosity (H_o)

mean expected heterozygosity (H_e) = gene diversity

nucleotide polymorphism (θ)

nucleotide diversity (π)

GENETIC VARIATION IN NATURAL POPULATIONS

Issue of the extent of variation in natural populations:



T.H. Morgan, H. Muller: "classical" model limited variability







A. Sturtevant, T. Dobzhansky: "equilibrium" model variation widespread

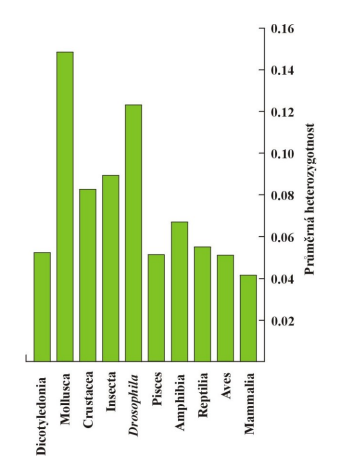




GENETIC VARIATION IN NATURAL POPULATIONS

1966: Harry Harris – humans; Richard Lewontin, John Hubby – D. pseudoobscura

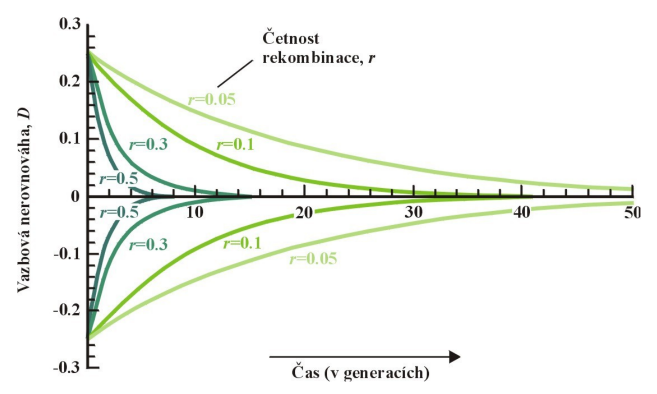
Taxon	Počet zkoumaných druhů	Podíl lokusů polymorfních	Průměrná heterozygotnost	
Bezobratlí				
mořští plži	5	0.175	0.083	
suchozemští plži	5	0.437	0.150	
ostatní mořští bezobratlí	9	0.587	0.147	
haplodiploidní blanokřídlí	6	0.243	0.062	
Drosophila	43	0.431	0.140	
ostatní hmyz	23	0.329	0.074	
bezobratlí celkem	93	0.397	0.112	
Obratlovci				
ryby	51	0.152	0.051	
obojživelníci	13	0.269	0.079	
plazi	17	0.219	0.047	
ptáci	7	0.150	0.047	
hlodavci	26	0.202	0.054	
savci	46	0.147	0.036	
obratlovci celkem	135	0.173	0.049	
Rostliny celkem	473	0.505		



microsatellites, minisatellites → high mutation rate, high variability question to what extent protein electrophoresis representative?

VARIATION AT MORE LOCI

proximity of loci = linkage valid H-W assumptions \Rightarrow formation of linkage equilibrium this proces can be slow \Rightarrow linkage disequilibrium (LD) coefficient of LD: D relation of D to recombination r:



Causes of linkage disequilibrium:

absence of recombination (eg. inversion)

LD needn t exist only between loci on the same chromosome!

nonrandom mating

selection

recent mutation

sample is a mixture of 2 species with different allele frequencies

recent merging of 2 populations

random genetic drift

INBREEDING

= mating between relatives

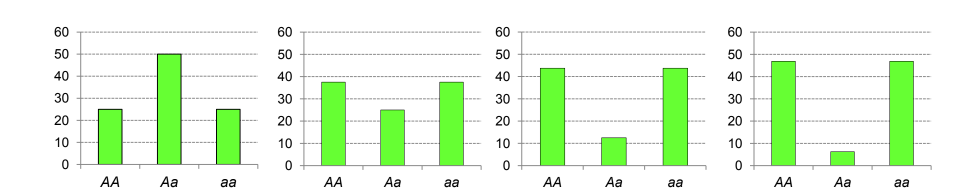
eg. repeated autogamy (self-fertilization, self-pollination):

initial generation (HWE): 1/4 AA, 2/4 Aa, 1/4 aa

1. gen. of selfing: 3/8 AA, 2/8 Aa, 3/8 aa

2. gen. of selfing: 7/16 AA, 2/16 Aa, 7/16 aa

3. gen. of selfing: 15/16 AA, 2/32 Aa, 15/16 aa



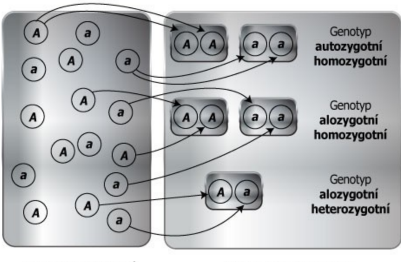
Ovules (produce female

INBREEDING COEFFICIENTS

1. Pedigree inbreeding, *F*:

= probability of autozygosity

Vztah alel v genotypech nové populace vzhledem k původní



Alely v populaci předků, které nejsou IBD

Genotypy v nové populaci

autozygosity:

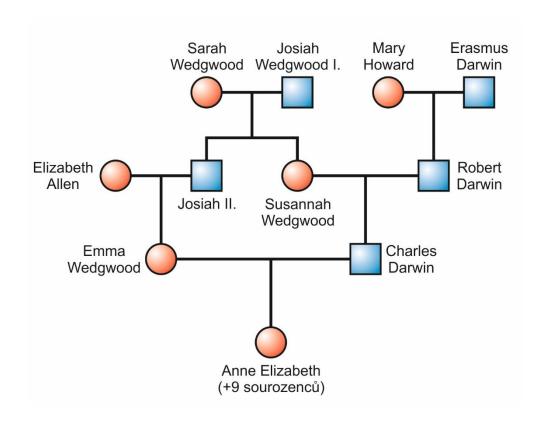
alleles identical by descent (IBD), always homozygous

allozygosity:

either heterozygote or homozygote (alleles identical by state, IBS)

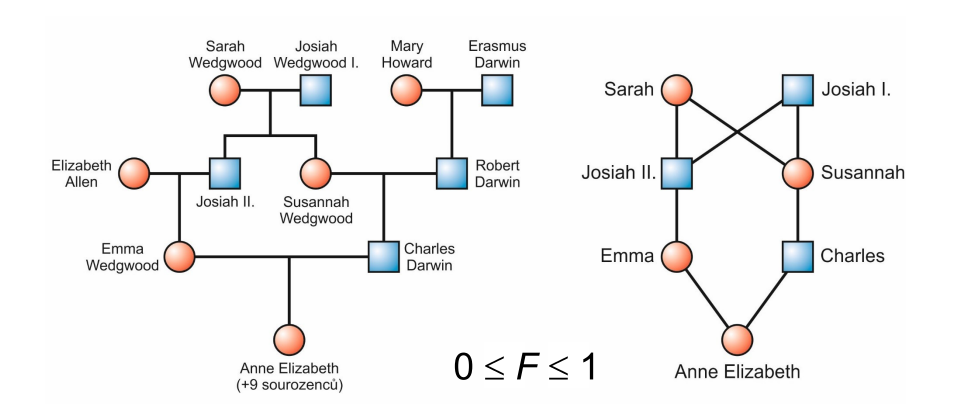
Inbred population = pop. in which the probability of autozygosity due to inbreeding > in panmictic population

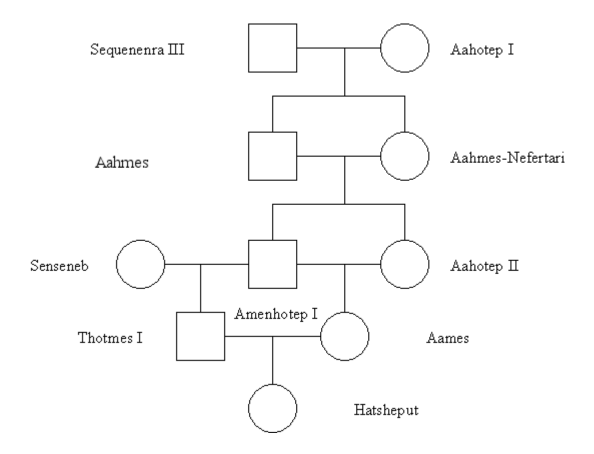
F = probability that an individual inherited both alleles at a locus from the same ancestor (both alleles are IBD)



Inbred population = pop. in which the probability of autozygosity due to inbreeding > in panmictic population

F = probability that an individual inherited both alleles at a locus from the same ancestor (both alleles are IBD)





a`	Amenhoten	I. and Aahote	n II	25%
a	Amemoteb	i. and Aanole	β 11.	ZJ /0

b) Aames 37.5%

c) Hatsheput 25%

d) Remaining in the pedigree are not inbred, ie F = 0

2. System-of-mating inbreeding, F_{1S} :

= deviation from HWE

$$F_{IS} = (H_e - H_o)/H_e$$
 $-1 \le F_{IS} \le +1$

$$-1 \le F_{1S} \le +1$$

 H_0 = observed

 $H_{\rm e}$ = expected heterozygosity



F and F_{IS} don t measure the same thing!

F is the <u>individual</u> measure, F_{IS} is the <u>group</u> measure

Př.: hutterites (anabaptists) of the Great Plains in USA and Canada:

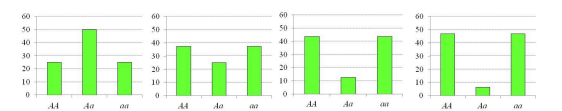
in spite of respecting the incest taboo this is one of the most inbred human groups known (F = 0.0255)

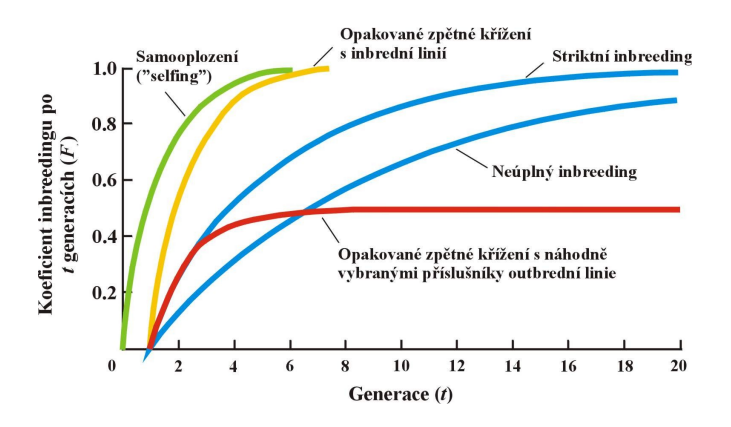
caused by a small number of founders (Protestants from Tyrol and Carinthia, 16th century)

Genetic effects of inbreeding:

inbreeding changes genotype frequencies (increase of homozygote freq.)

× <u>allele freqs. don t change</u> affects all loci





Phenotypic effects of inbreeding:

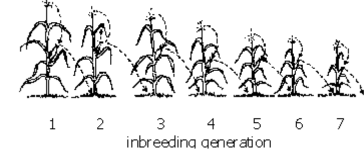
inbreeding depresion

diseases, reduced fertility and/or viability





Leavenworthia alabamica



BUT! Not always must inbreeding be deleterious (eg. many species of embryophyte (land) plants are self-fertilising). Moreover, the inbreeding effects can differ within a single species depending on environment.

Inbreeding depression in humans:

the Amish: haemophilia B, anemia, myotonic dystrophy, Ellis-van Creveld syndrome (dwarfness, polydactyly), defects in nail development, dental

defects







Vadoma tribe, Zimbabwe (tzv. "Ostrich people"): ectrodactyly

Mormons of Hilldale (Utah) and Colorado City (Arizona)

Amazonia Indians aristocratic dynasties



Human inbreeding depression:

Charles II of Spain:

unnaturally big head, deformed mandible, weak body, difficulties with walking and other defects, mental and psychical defects, impotence, sterility



Francis II:

in some children mental retardation, hydrocephaly, seizures,

some unable of living without assistence





Maria Theresa

hybrid vigour (heterosis)



Francis I of Lorraine

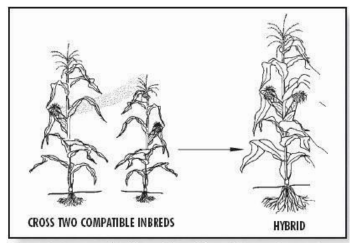


Figure 6. Cross pollination of two inbreds to produce a vigorous hybrid.

ASSORTATIVE MATING

= higher probability of mating between individuals with the same phenotype

can be caused by active mating preference but another causes can exist as well

- eg.: phytophagous insects individuals living at different host species can mature in different times ⇒ more frequent mating between individuals of the same phenotype (confinement to the host) without active mating preference
- ⇒ this is only a positive <u>phenotypic correlation</u> assortative mating causes deficit of heterozygotes assortative mating causes <u>linkage disequilibrium (LD)</u>

Differences between inbreeding and assortative mating:

affects only locus (loci) connected with preferred phnotype inbreeding affects all loci

ass. mating is <u>a powerful evolutionary force</u> (strong LD at more loci) × inbreeding only strenghtens existing LD, and only in the case of selfing, in other cases recombination "more succesful" → reduction of LD

NEGATIVE ASSORTATIVE (DISASSORTATIVE) MATING

= preference of mates with different phenotypes
 results in <u>intermediary allele frequencies</u>, <u>reduces LD</u>
 eg. preference of males with different MHC (mouse, man)

