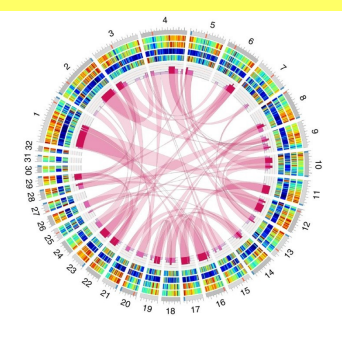
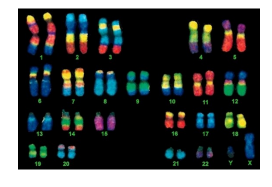
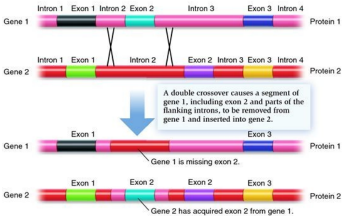
EVOLUTION OF GENETIC SYSTEMS

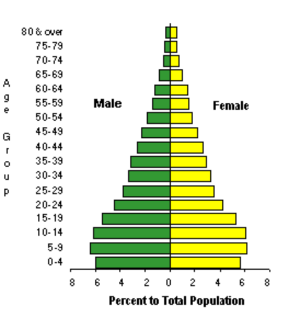












EVOLUTION OF THE GENOME

Genome size and C-value:

C-value = amount of DNA in haploid genome (pg, bp)

Prokaryotes:

 $6 \times 10^5 - 10^7$ bp (20-fold span)

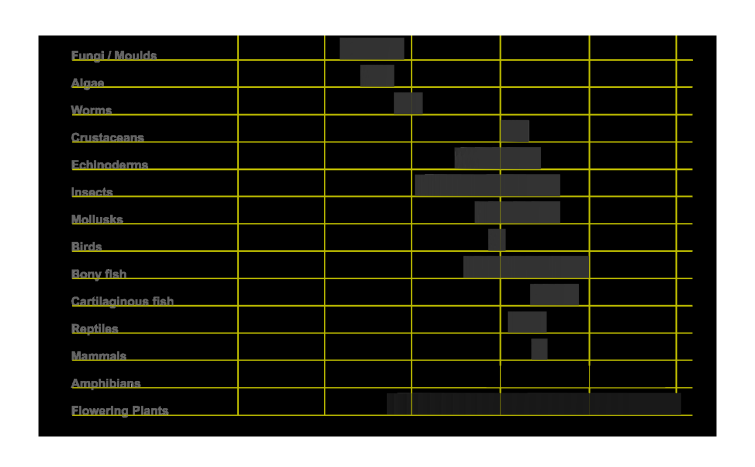
smalles: *Mycoplasma genitalium* 525 genes, the smallest fuctional artificially synthesized genome ≈ 473 genů (modified from *M. mycoides*)

largest: some G+ bacteria, cyanobacteria

Mycoplasma			in bp	
Gram positive bacteria				
Gram negative bacteria				

Eukaryotes:

 $8.8 \times 10^6 - 6.9 \times 10^{11}$ bp (80 000-fold span!)



no relation between genome size and organismal complexity or number of genes

large differences even in related organisms:

Paramecium caudatum (8 600 000 kb) × P. aurelia (190 000 kb)

human: ca. 6×10^9 bp (~ 6,5 pg DNA)

× Amoeba proteus: 2,9×10¹¹ bp Polychaos dubium (Amoeba dubia): 6,7×10¹¹ bp

⇒ C-value paradox (C-value enigma)

Table 4. C values from eukaryotic organisms ranked by genome size.

Species	C value (kb)				
Navicola pelliculosa (diatom)	35,000				
Drosophila melanogaster (fruitfly)	180,000				
Paramecium aurelia (ciliate)	190,000				
Gallus domesticus (chicken)	1,200,000				
Erysiphe cichoracearum (fungus)	1,500,000				
Cyprinus carpio (carp)	1,700,000				
Lampreta planeri (lamprey)	1,900,000				
Boa constrictor (snake)	2,100,000				
Parascaris equorum (roundworm)	2,500,000				
Carcarias obscurus (shark)	2,700,000				
Rattus norvegicus (rat)	2,900,000				
Xenopus laevis (toad)	3,100,000				
Homo sapiens (human)	3,400,000				
Nicotiana tabaccum (tobacco)	3,800,000				
Paramecium caudatum (ciliate)	8,600,000				
Schistocerca gregaria (locust)	9,300,000				
Allium cepa (onion)	18,000,000				
Coscinodiscus asteromphalus (diatom)	25,000,000				
Lilium formosanum (lily)	36,000,000				
Amphiuma means (newt)	84,000,000				
Pinus resinosa (pine)	68,000,000				
Protopterus aethiopicus (lungfish)	140,000,000				
Ophioglossum petiolatum (fern)	160,000,000				
Amoeba proteus (amoeba)	290,000,000				
Amoeba dubia (amoeba)	670,000,000 <				

Data from Cavalier-Smith (1985), Sparrow et al. (1972), and other references.

marbled lungfish: > 40× larger than human

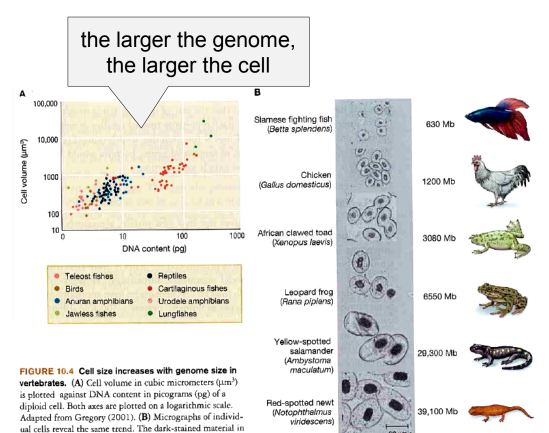
almost 200× larger than human

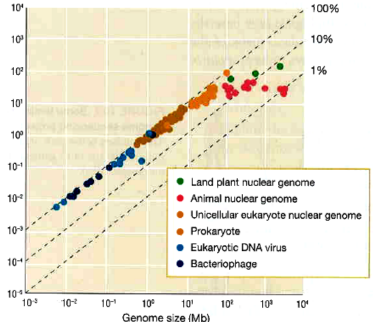
closely related species!

C-value paradox:

the center of each cell is DNA.

large genomes include large amount of non-coding DNA





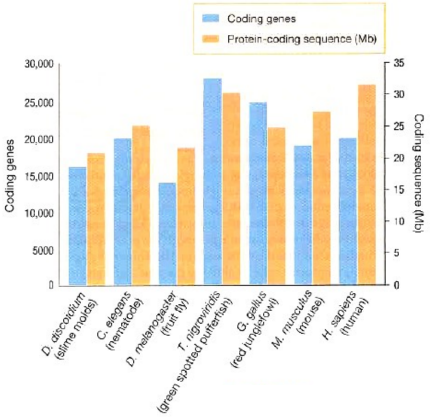
Coding DNA (Mb)

large genomes ⇒ large cells ⇒ influence on:

cell division speed
metabolism efficiency
rate of ions/proteins
exchange
body size

G-value paradox:

despite diversity of organismal complexity, metazoans tend to have similar numbers of protein-coding genes (G-value)



No dependency on the total number of genes but on complexity of gene regulation networks – organisms with similar number of genes may have very different patterns of gene regulation networks

					8	00		3	1	W W W W W W W W W W W W W W W W W W W	
whole genome sequence		0.0036 0.0054 Mb Mb	1.8 Mb	12.5 Mb		100 Mb	123 Mb	115 Mb	3100 Mb		
		1976	1977	$-n_{\parallel}$	1995	1996	1997	1998	1999	2000	2001
	Bacteriophage MS2	Bacteriophage		Haemophilus influenzae	Saccharomyces cerevisiae		Caenorhabditis elegans	Drosophila melanogaster	Arabidopsis thaliana	Homo sapiens sapiens (draft; completed 2003)	

How many coding genes are in the human genome?

before 2001 (draft version of the genome) estimates from 50 000 till > 140 000 (max. 212 278) genes

Int. Human Genome Sequencing Consortium (IHGSC) 2001: 30 000–40 000 protein coding genes

IHGSC 2004: 20 000-25 000 protein coding genes

Ensembl – May 2012: 21 065 coding genes

Ensembl – January 2013: 20 848 coding genes

Ensembl – February 2014: 20 805 coding genes

Ensembl – December 2014: 20 364 coding genes

Repetitive DNA:

- 1. Highly repetitive = satellite
- 2. Moderately repetitive = minisatellites, microsatellites
- 3. Transposable elements, retroelements (SINE, LINE)

Why does repetitive DNA exist?

Cavalier-Smith (1978): there must be some function

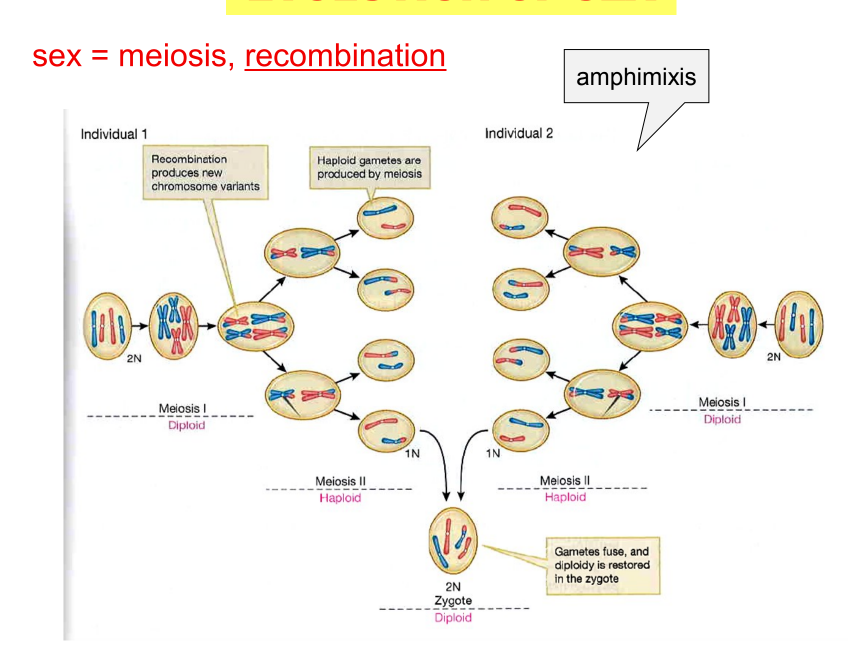
Doolittle and Sapienza, Orgel and Crick (1980): repetitive DNA is "selfish"

Susumu Ohno (1972): "junk DNA"



"junk" ≠ "garbage" ⇒ in future it may gain some function

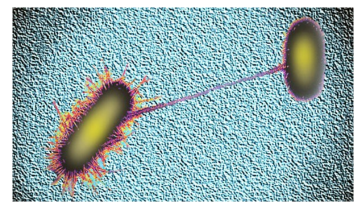
EVOLUTION OF SEX

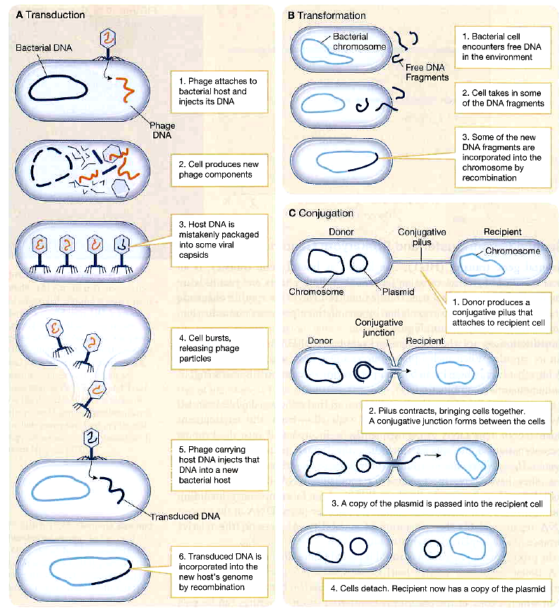


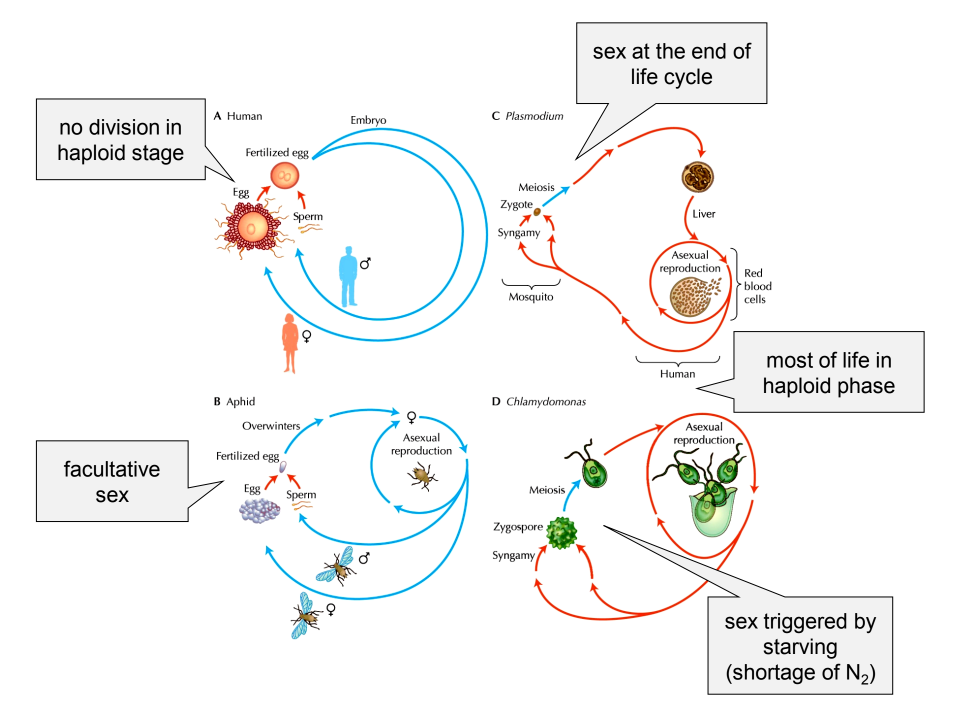
"sex" in Prokaryotes:

conjugation transformation transduction

conjugation in *E.coli*:

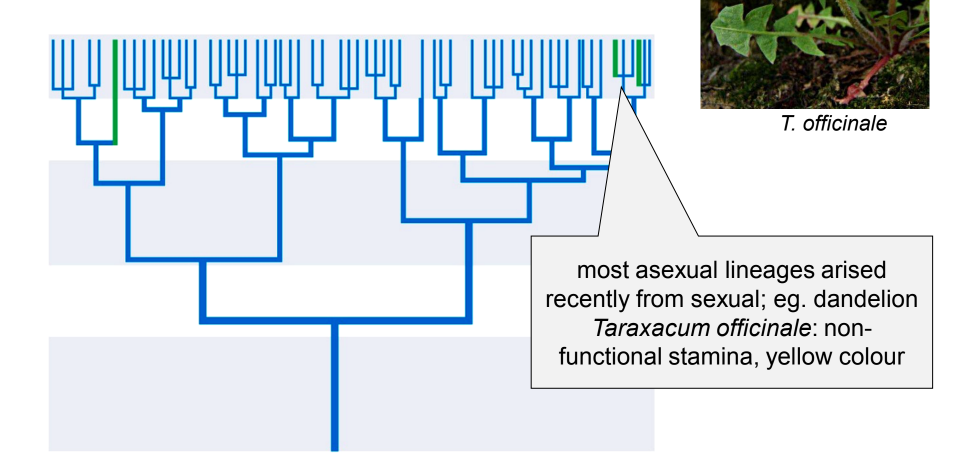






phylogenetic position of asexual taxa:

mostly recent lineages taxa scattered



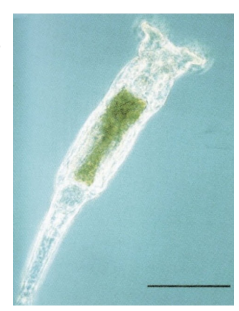
exceptions:

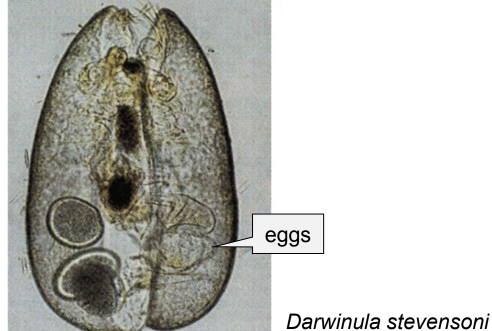
Bdelloidea rotifers: fossils in amber 35-40 MY existency ~100 MY

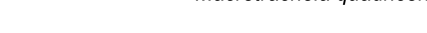
ostracods:

asexual ~100 MY × recently males found











Disadvantages of sexual reproduction

time and energy necessary for finding a partner (finding itself may be a problem), further effort before copulation

increased risk of predation or parasitation, transmission of venereal diseases

susceptibility to extinction at low N_e

lower capability of colonization

complex meiotic molecular machinery meiosis: 10-100 h × mitosis: 15 min – 4 h

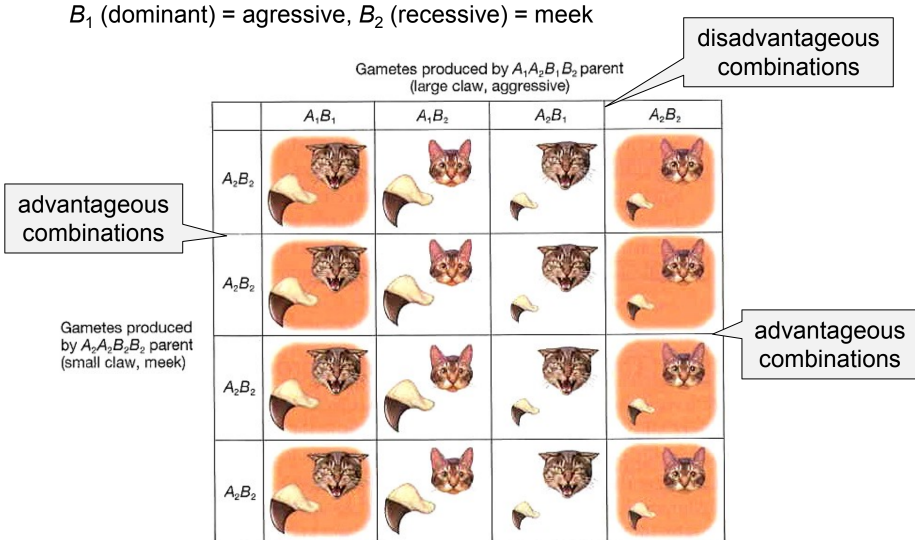
impact of sexual selection on males → reduction of population fitness eg. Soay sheep (St. Kilda): males die during the first winter × females and castrated males several years

Disadvantages of sexual reproduction:

break-up of advantageous allele combinations by recombination

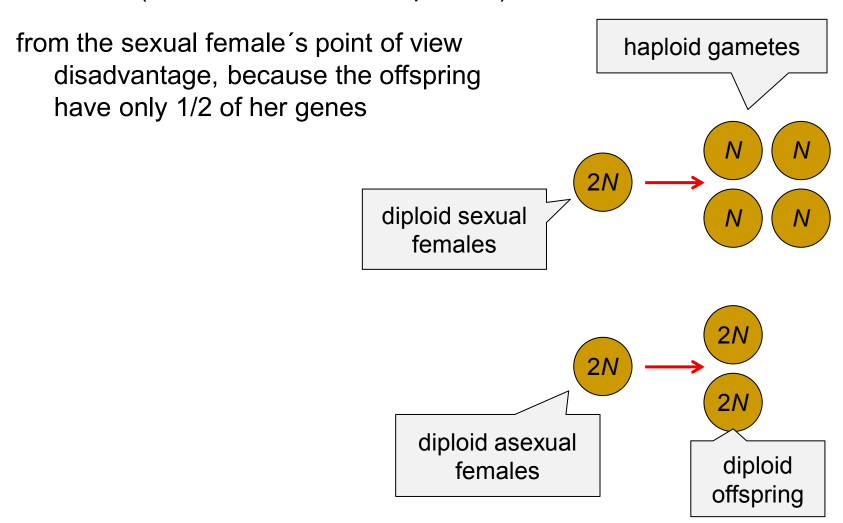
Eg.: A_1 (dominant) = large claws, A_2 (recessive) = small claws





Disadvantages of sexual reproduction:

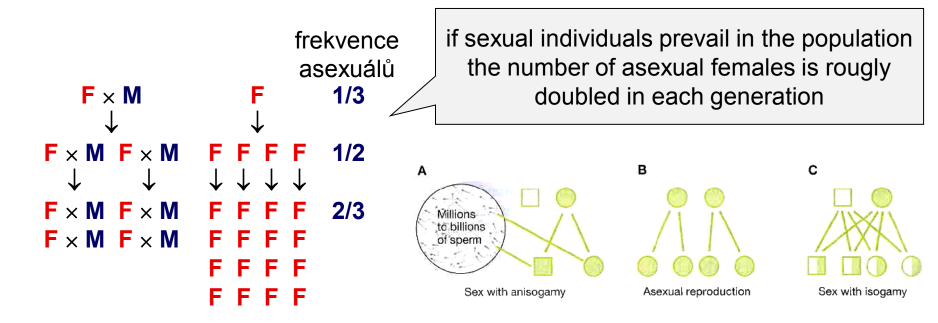
action of selfish elements (conflict of genes) → reduction of population fitness (B chromosomes, transposons)



J. Maynard Smith: What is the fate of sexual and asexual population?

assumptions: way of reproduction has no effect on

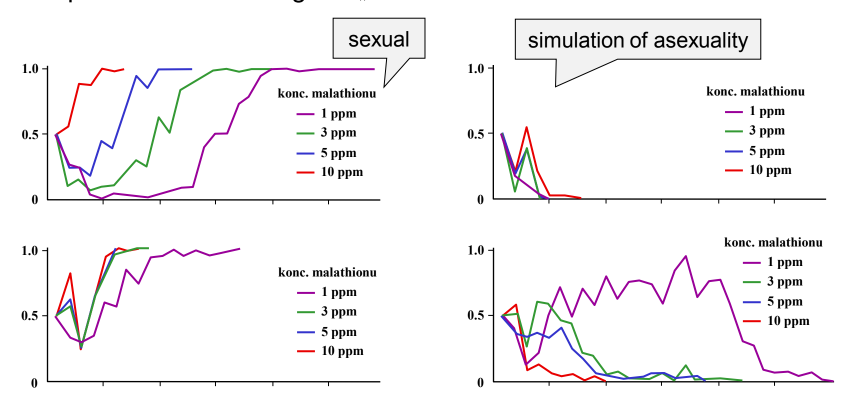
- 1. number of descendants (eg. when males take care of offspring)
- 2. probability of offspring survival



⇒ twofold cost of sex, ie. 50% selective disadvantage of sex (not for isogamy! → so rather cost of males)

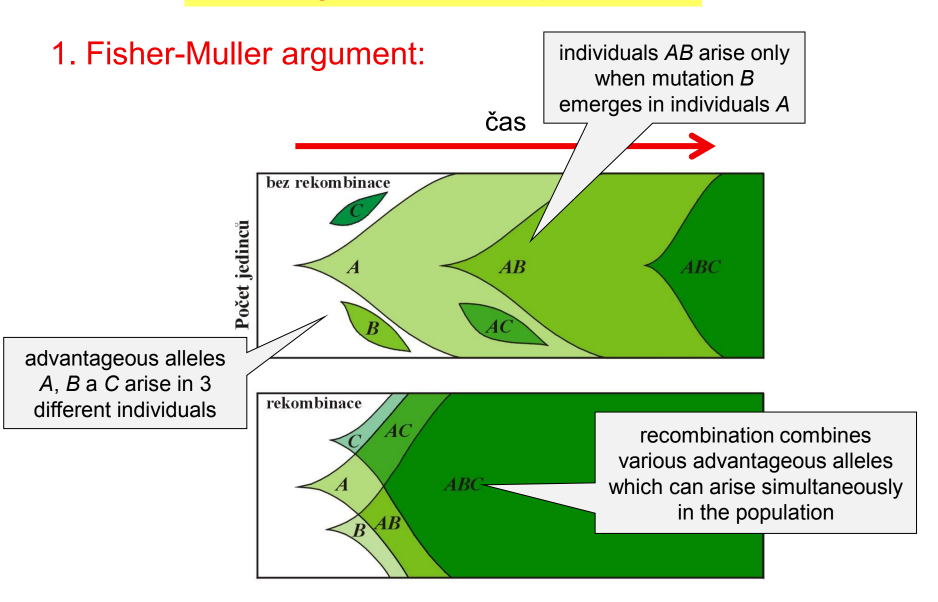
ad 2) effect of environment

experiment with *Tribolium castaneum*: competition, insecticide, reproductive advantage of "asexuals"



at first prevalence of asexuals, eventually fixation of sexuals faster at higher insecticide concentrations offspring of sexuals have higher fitness ⇒ assumption 2 is not valid

Advantages of sexual reproduction



Effects of recombination:

- 1 locus → max. 2 variants of gametes (heterozygote)
- 2 loci \rightarrow 4 variants: gametes $AB/ab \rightarrow ab$, aB, Ab, AB
- 10 loci \rightarrow 2¹⁰ = 1024 different gametes and 2ⁿ⁻¹(2ⁿ+1) = 524 800 diploid genotypes

for population genetics *the only* consequence of sex is linkage equilibrium

– when it is reached sex loses sense

every model explaining advantage of sex must include a mechanism which eliminates some gene combinations (LD arises), and explain why genes causing LD are favoured by selection

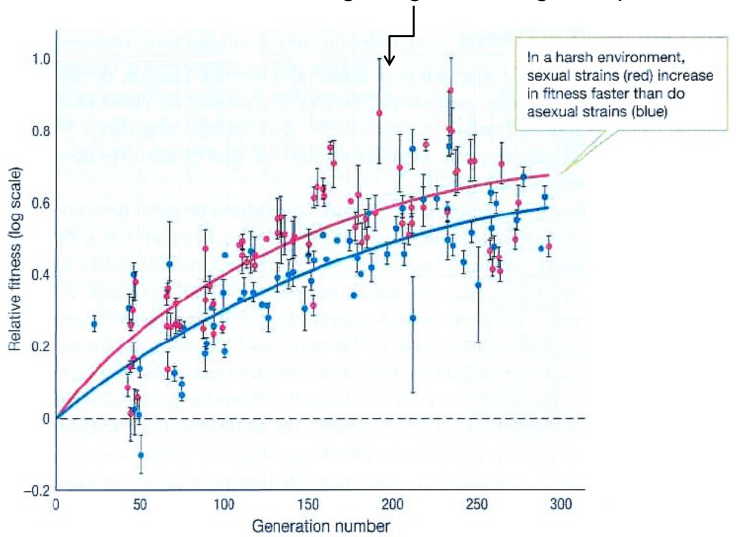
Sexual reproduction increases variation and hence rate of evolution but this advantage mostly in long-term perspective, asexuality in the short-term more advantageous

Eg.: yeast Saccharomyces cerevisiae

favourable environment: abundance of glucose, optimal temperature

→ no difference

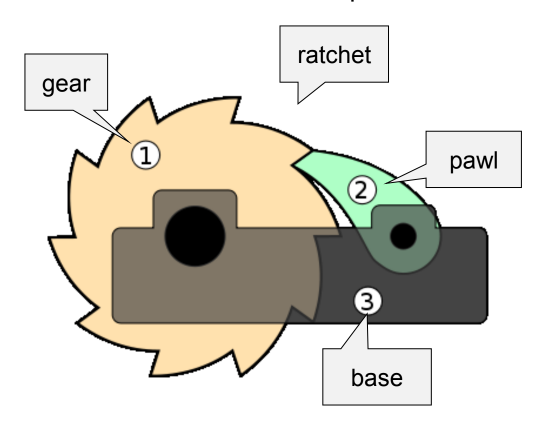
unfavourable environment: shortage of glucose, high temperature



2. Elimination of deleterious mutations I. Muller's ratchet:

The only way how to escape from deleterious mutations either

back mutation, or mutation which invalidate effect of the previous mutation



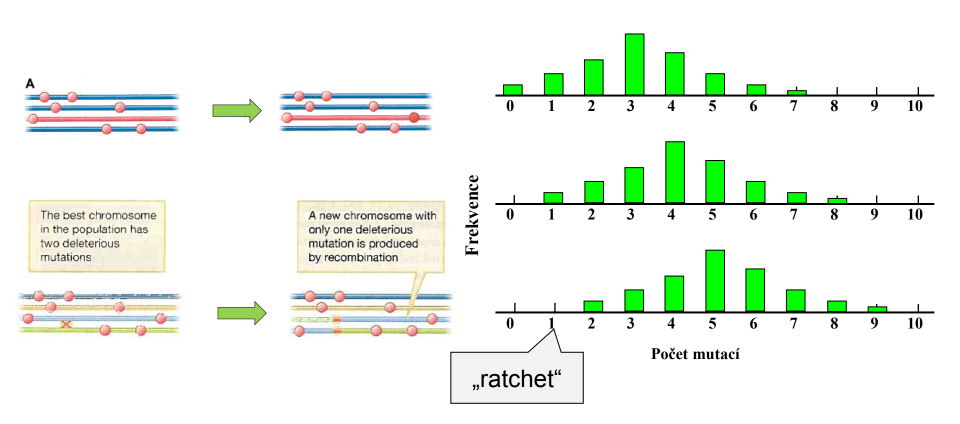
accumulation of deleterious mutations

small population \Rightarrow role of drift (stochastic process)

with sex chance to avoid "ratchet"

with increase of genotype frequencies without deleterious mutations spread of genes responsible for sex

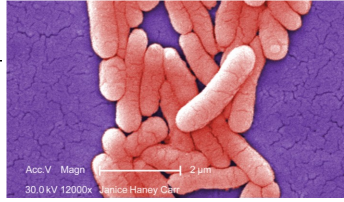
best when mutations are only slightly deleterious



Andersson and Hughes (1996) - Salmonella typhimurium

444 experimental cultures, each from 1 individual → growth overnight repetition ⇒ repeated drift, total of 1700 generations
 comparison with a free-living strain

→ 5 cultures (1%) with significantly reduced fitness, none with higher



Lambert and Moran (1998) – comparison of fitness of bacteria living within insect cells with free-living species

9 species of bacteria living only in insect cells
each species had its free-living relative counterpart
thermal stability of rRNA genes
did endosymbionts accumulated deleterious mutations?

→ in all cases rRNA of endosymbionts by 15 - 25% less stable

3. Elimination of deleterious mutations II. Kondrashov's model:

Alexey S. Kondrashov (1988)

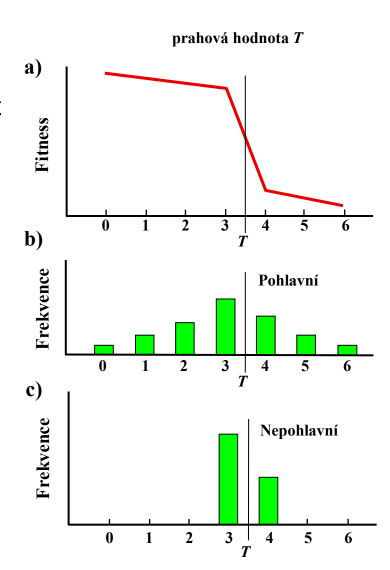
assumption that deleterious mutations act synergically → epistasis

"truncation selection" (deterministic process)

since in sexuals proportion of deleterious mutations exceeding *T* value is higher than in asexuals, elimination of these mutations is faster in the former (recombination combines them)

question if frequencies of deleterious mutations are sufficiently high (at least 1/generation/genome)

model proven in E. coli and S. cerevisiae



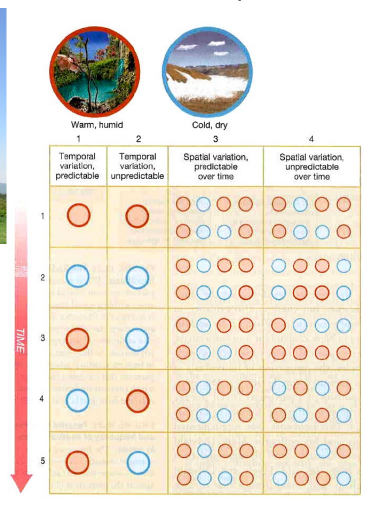
4. Unpredictable environment – lotery model, elm-oyster model

biotope divided into local sites to which descendants randomly "distributed"

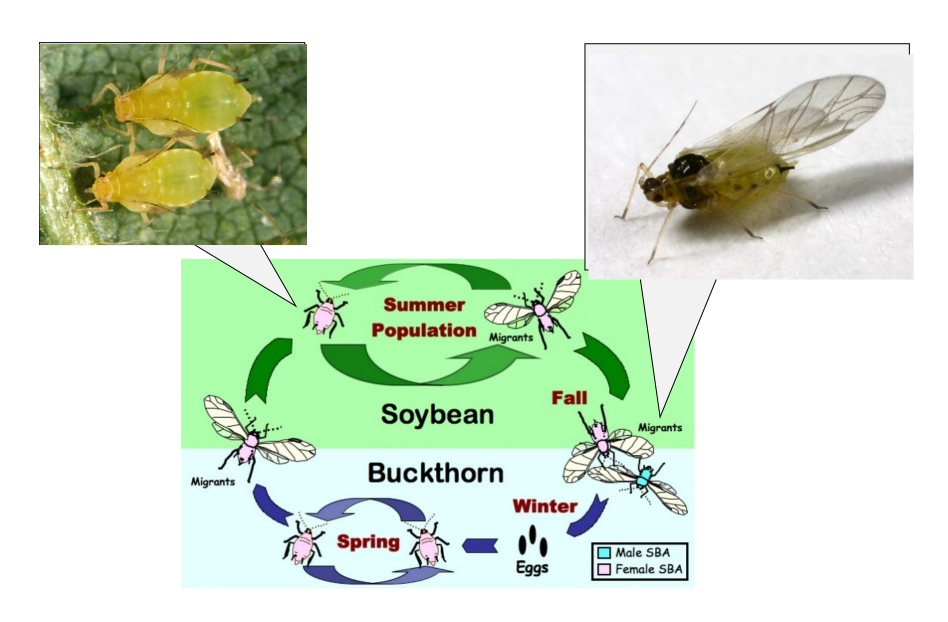
→ only best adapted ones survive, parents cannot know a priori which

of them will do

analogy with purchase of a lottery ticket

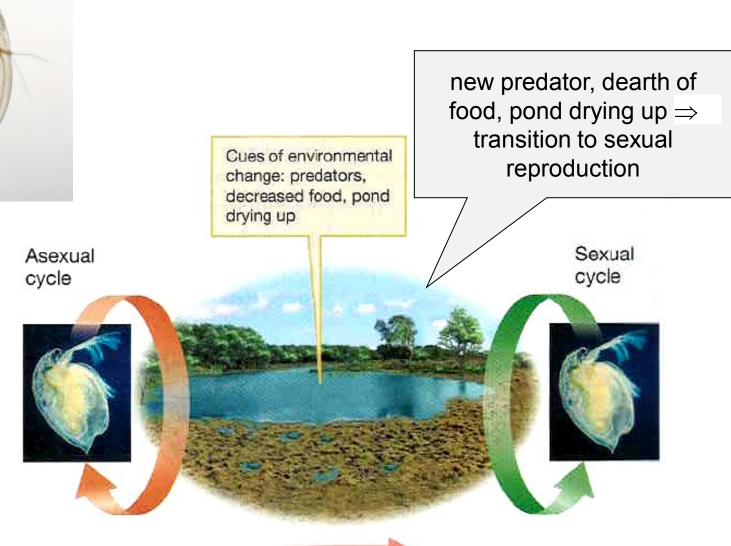


Eg. aphids:





Eg. Daphnia:





Unpredictable environment – elbow room model

assumption that in heterogenous and homogenous biotopes genotypes can differ in usage of limited sources

competition among siblings → more descendants of sexual parents can coexist at the same site because competition of asexual offspring is more intense

Problem: models 4 and 5 are valid only for organisms with high fertility

Fluctuation of environment:

itself does not maintain sex → <u>fluctuation of epistasis</u> necessary

eg. 2 loci: alternatio of association cold-wet and warm-dry ↔ cold-dry and warm-wet

this model can work eg. in parasite-host interaction

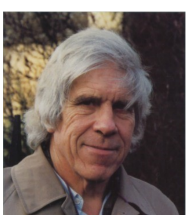
5. Red Queen hypothesis

William D. Hamilton

based on the Red Queen hypothesis (Leigh Van Valen)



"The Red Queen has to run faster and faster in order to keep still where she is. That is exactly what you all are doing!"



W.D. Hamilton



L. Van Valen

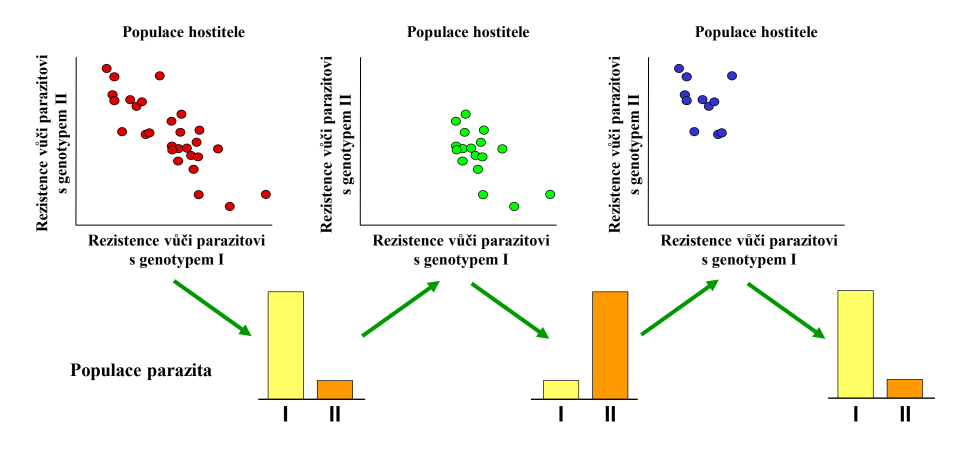
fluctuation of epistasis

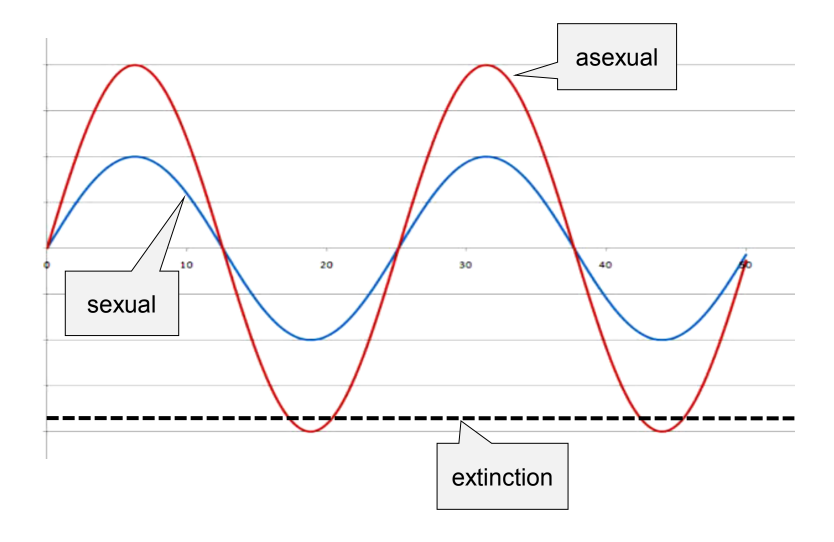
fitness and gene frequencies cycles

coevolution of parasite and host ⇒ arms races

multilocus "gene-for-gene" relation

oscillation of gene frequencies higher in asexual individuals





model assumption: in heterogonous organisms (changing of sexual and asexual reproduction) and organisms with facultative sexuality sexual reproduction more frequent in case of increased parasitation

Curtis Lively (1992): freshwater gastropod *Potamopyrgus antipodarum*

New Zealand lakes and rivers both sexual and asexual females



Lake Alexandria, South Island, New Zealand

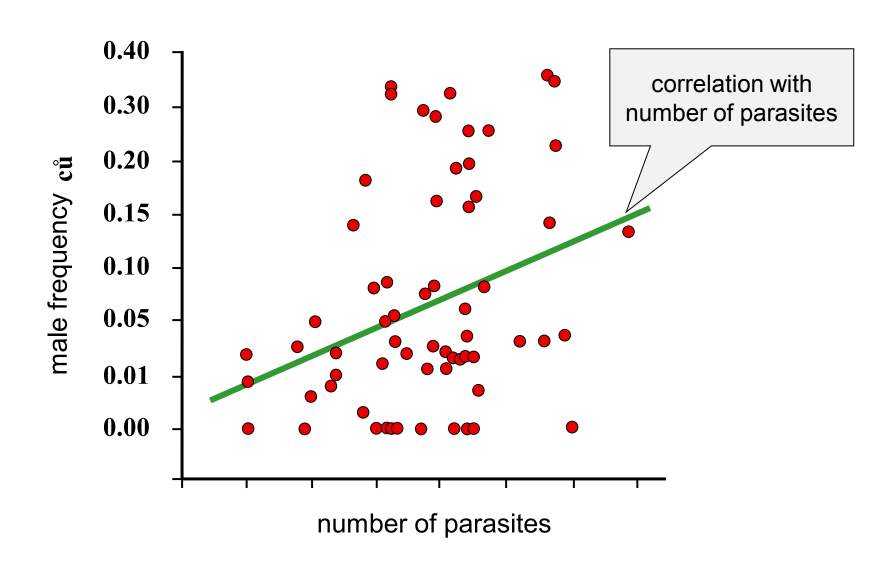


Potamopyrgus antipodarum

>12 parasitic trematode species (host castration \Rightarrow strong selection) 66 lakes

number of males as indicator of sexual reproduction

Lively et al. (1992):



EVOLUTION OF SEX RATIO

sex ratio often 1:1 \rightarrow why to waste for males?

R. A. Fisher (1930)

frequency-dependent selection

condition for validity of Fisher's argument:

- 1. random mating
- 2. same costs of both sexes

ad 1) Local mating competition:

mites Adactylidium, Pyemotes ventricosus, Acarophenax tribolii parasitoid wasps (eg. Nasonia vitripennis)



Nasonia vitripennis

Melittobia digitata

Neobellieria bullata (Pupa)

(C) Jorge M. González

Nasonia vitripennis



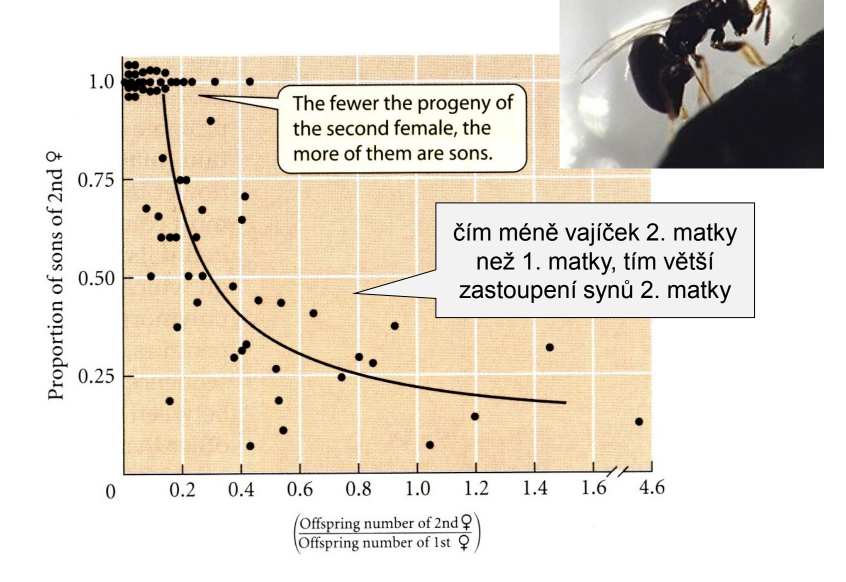
Pyemotes ventricosus



Acarophenax tribolii

theoretical prediction: with increasing number of egg laying females

percentage of sons increases



ad 2) Trivers-Willard hypothesis:

Robert L. Trivers, Dan Willard

investment in sex ensuring higher fitness in next generation

dominant mother → investment in sons and vice versa

sex ratio bias or unequal parental investment

Eg.: deers









D. Willard

sons of dominant mothers have higher fitness

