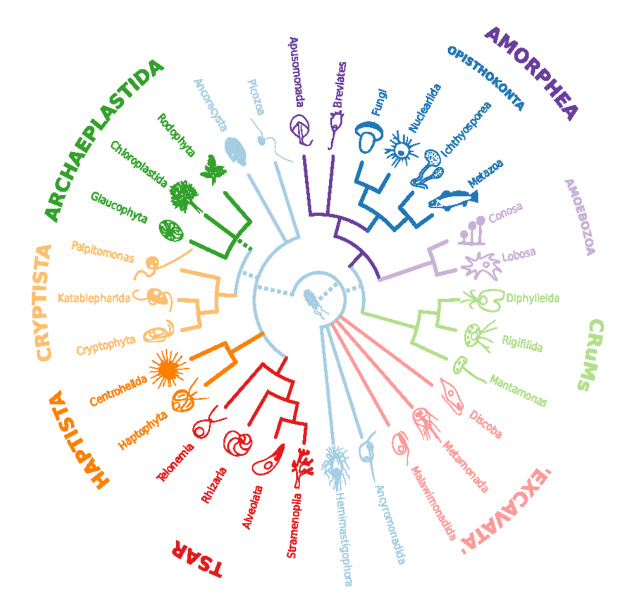
# Genome and chromosome evolution



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### Genome and chromosome evolution

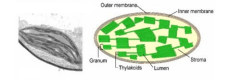
- genome size variation
- chromosome number variation
- variation in coding DNA amount
- variation in non-coding DNA

## Genome size variation

#### Polychaos dubium



...perhaps the largest known genome -670 billion base pairs (670 Gb) (~200-times larger than the human genome, 3.2 Gb; some authors suggest treating the value with caution)





organism	genome size (base pairs)	protein coding genes	number of chromosomes	
model organisms				
model bacteria E. coli	4.6 Mbp	4,300	1	
budding yeast S. cerevisiae	12 Mbp	6,600	16	
fission yeast S. pombe	13 Mbp	4,800	3	
amoeba D. discoideum	34 Mbp	13,000	6	
nematode C. elegans	100 Mbp	20,000	12 (2n)	
fruit fly D. melanogaster	140 Mbp	14,000	8 (2n)	
model plant A. thaliana	140 Mbp	27,000	10 (2n)	
moss P. patens	510 Mbp	28,000	27	
mouse M. musculus	2.8 Gbp	20,000	40 (2n)	
human H. sapiens	3.2 Gbp	21,000	46 (2n)	
viruses				
hepatitis D virus (smallest known animal RNA virus)	1.7 Kb	1	ssRNA	
HIV-1	9.7 kbp	9	2 ssRNA (2n)	
influenza A	14 kbp	11	8 ssRNA	
bacteriophage λ	49 kbp	66	1 dsDNA	
Pandoravirus salinus (largest known viral genome)	2.8 Mbp	2500	1 dsDNA	
organelles				
mitochondria - H. sapiens	16.8 kbp	13 (+22 tRNA +2 rRNA)	1	
mitochondria – S. cerevisiae	86 kbp	8	1	
chloroplast – A. thaliana	150 kbp	100	1	
bacteria				
C. ruddii (smallest genome of an endosymbiont bacteria)	160 kbp	182	1	
M. genitalium (smallest genome of a free living bacteria)	580 kbp	470	1	
H. pylori	1.7 Mbp	1,600	1	
Cyanobacteria S. elongatus	2.7 Mbp	3,000	1	
methicillin-resistant S. aureus (MRSA)	2.9 Mbp	2,700	1	
B. subtilis	4.3 Mbp	4,100	1	
S. cellulosum (largest known bacterial genome)	13 Mbp	9,400	1	
archaea				
Nanoarchaeum equitans (smallest parasitic archaeal genome)	490 kbp	550	1	
Thermoplasma acidophilum (flourishes in pH<1)	1.6 Mbp	1,500	1	
Methanocaldococcus (Methanococcus) jannaschii (from ocean bottom hydrothermal vents; pressure >200 atm)	1.7 Mbp	1,700	1	
Pyrococcus furiosus (optimal temp 100°C)	1.9 Mbp	2,000	1	
eukaryotes - multicellular				
pufferfish Fugu rubripes (smallest known vertebrate genome)	400 Mbp	19,000	22	
poplar P. trichocarpa (first tree genome sequenced)	500 Mbp	46,000	19	
corn Z. mays	2.3 Gbp	33,000	20 (2n)	
dog C. familiaris	2.4 Gbp	19,000	40	
chimpanzee P. troglodytes	3.3 Gbp	19,000	48 (2n)	
wheat T. aestivum (hexaploid)	16.8 Gbp	95,000	42 (2n=6x)	
marbled lungfish P. aethiopicus (largest known animal genome)	130 Gbp	unknown	34 (2n)	
herb plant Paris japonica (largest known genome)	150 Gbp	unknown	40 (2n)	





## **Eukaryotes: minimal chromosome numbers**



Myrmecia pilosula "Jack jumper ant", Australia; males (haploid) n = 1, females (diploid) 2n = 2



five angiosperm species
e.g., Haplopappus gracilis, Asteraceae,
n = 2

## **Eukaryotes: highest chromosome numbers**

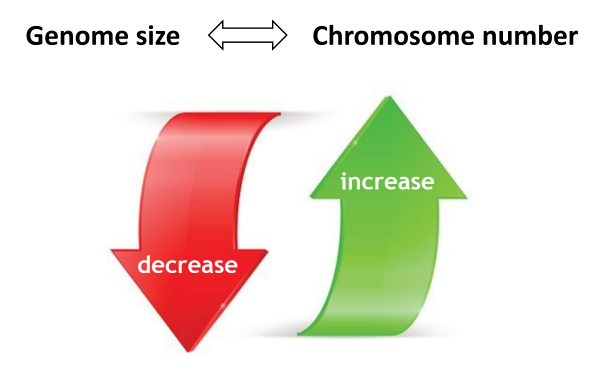


Polyommatus atlanticu<mark>s</mark> n = c. 220



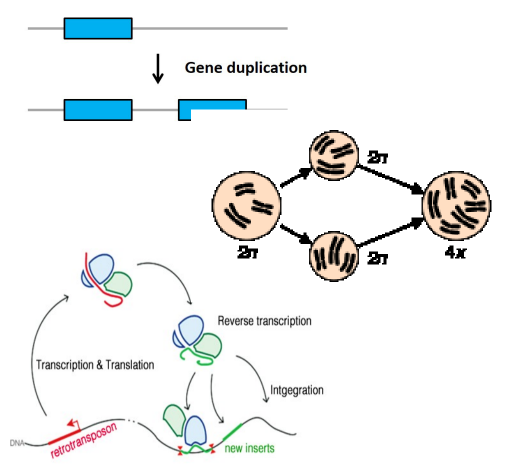
fern Ophioglossum reticulatum n = c. 530



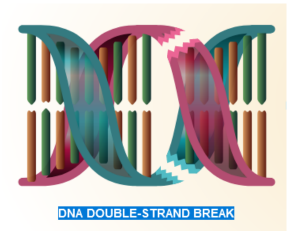


## Variation in genome size and chromosome number is driven by two principal processes

#### **DNA/genome duplication**



#### **DNA recombination**



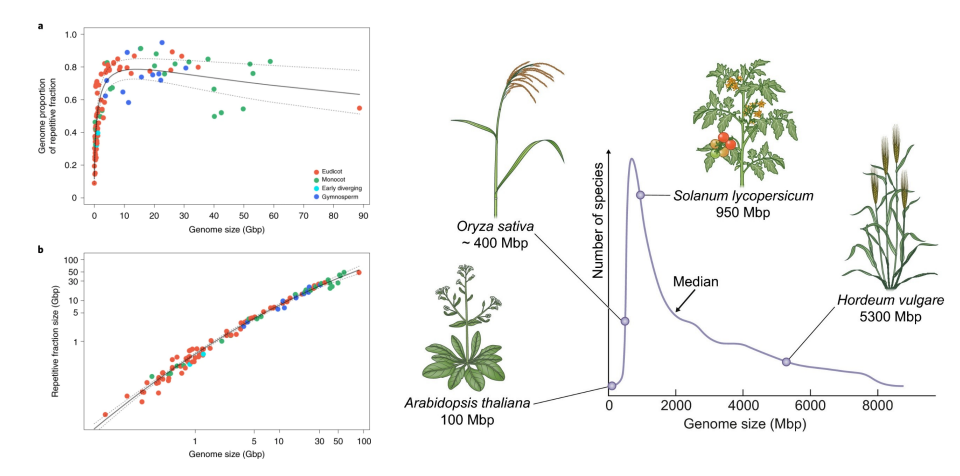
## $\int$

#### recombination

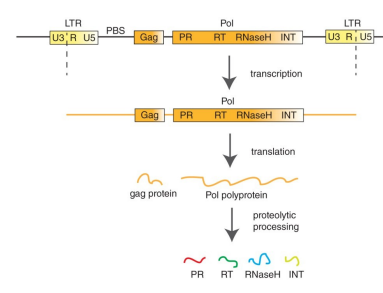
#### **Genome size increase**

- amplification of retrotransposons (and tandem repeats)
- gene and segmental duplications
- polyploidy

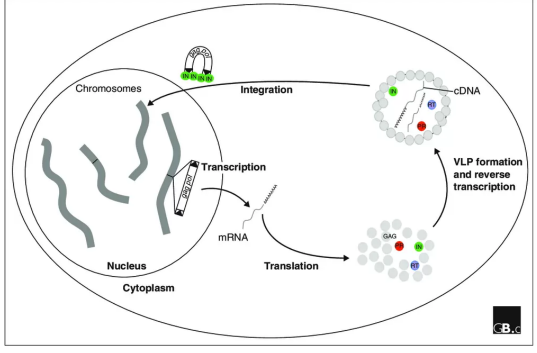
Genome size variation in angiosperms is driven by amplification (and elimination) of repetitive DNA



#### LTR (Long Terminal Repeat) retrotransposons (LTR-RTs)

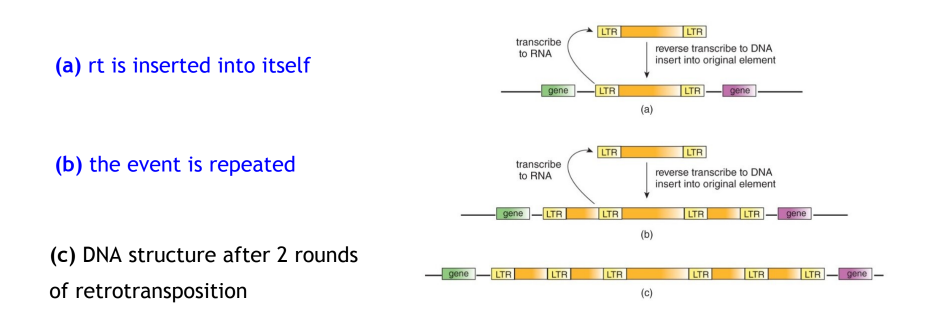


- Gag gene for the Gag protein
- **INT** integrase
- PBS primer binding site
- PR protease
- RT reverse transcriptase



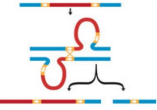
#### Genome size increase by retrotransposition

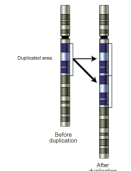
#### (nested retrotransposon insertion)



### Genome size increase by gene duplication

- replication slippage (errors in replication → gene duplication)
- ectopic recombination (between two direct repeats, typically TEs)

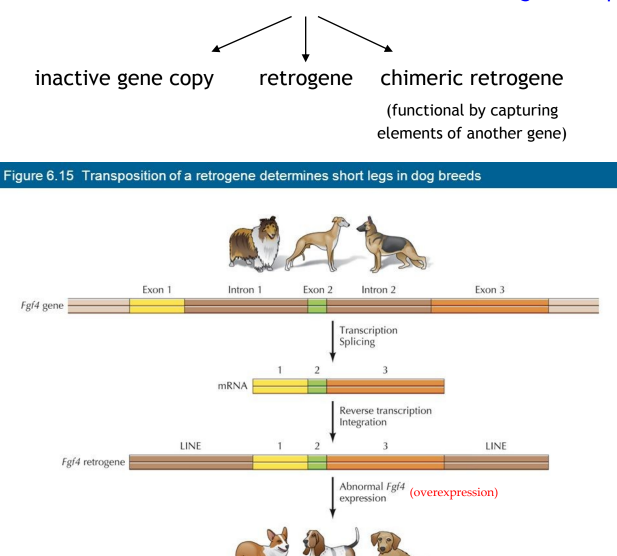




- unequal crossing-over in meiosis (due to missaligned chromosomes)
- via retrotransposition = retrogenes (cellular mRNA is transcribed into cDNA by reverse transcriptase of a retrotransposon or retrovirus; retrogene does not contain introns = lacking regulatory elements = pseudogene, but can evolve into a functional gene)

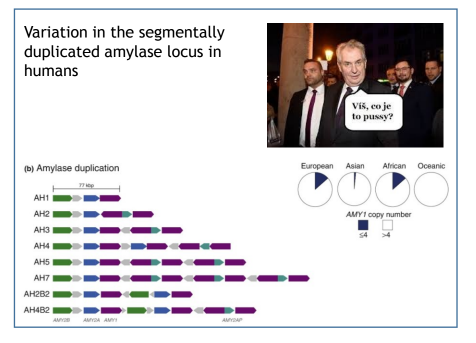
#### Retrogenes

• mRNA is reverse-transcribed into cDNA and inserted in a new genomic position



### **Segmental duplications**

- duplicated segment of chromosomal DNA (usually defined as > 1 kb in length)
- either tandem or interspersed organization, either intra-chromosomal or interchromosomal
- also known as low copy repeats
- human genome: 159 Mb gene-rich duplicated
   (5.5% of the genome) = c. Arabidopsis genome

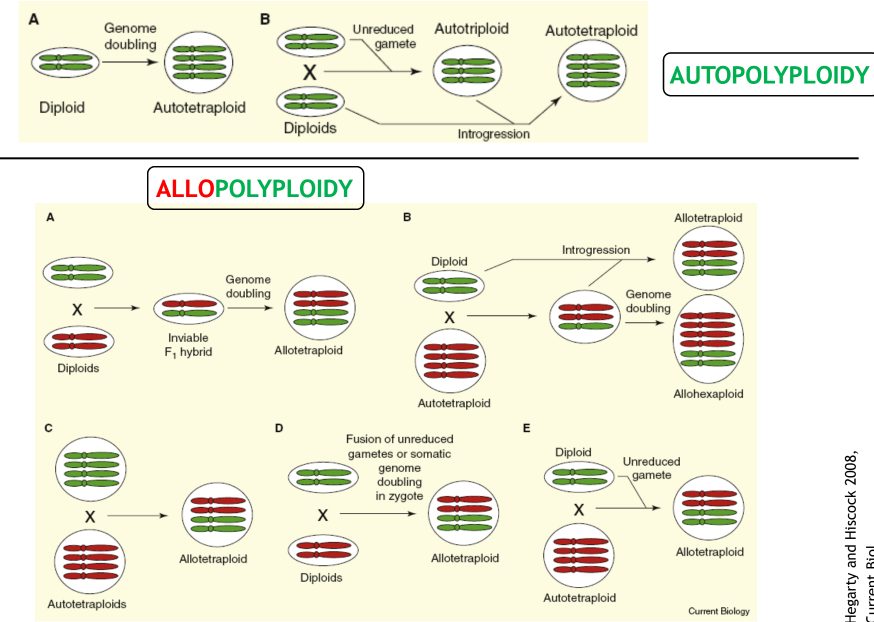


	Caenorhabditis elegans	Drosophila melanogaster	Human	Mouse	Rat	Chicken	Chimpanzee*
SDs of >1 kb	4.3%	1.2%	5.2%	2.7%	1.6%	2.7%	N.D.
SDs of >10 kb	0.7%	0.1%	4.5%	2.2%	1.5%	0.3%	N.D.
SDs of >20 kb	N.D.	N.D.	4.0%	1.7%	0.9%	0.0%	~4.8%
Genome size	97	123	2,866	2,506	2,566	1,040	2,866

 Table 1 | SD content of sequenced animal genomes

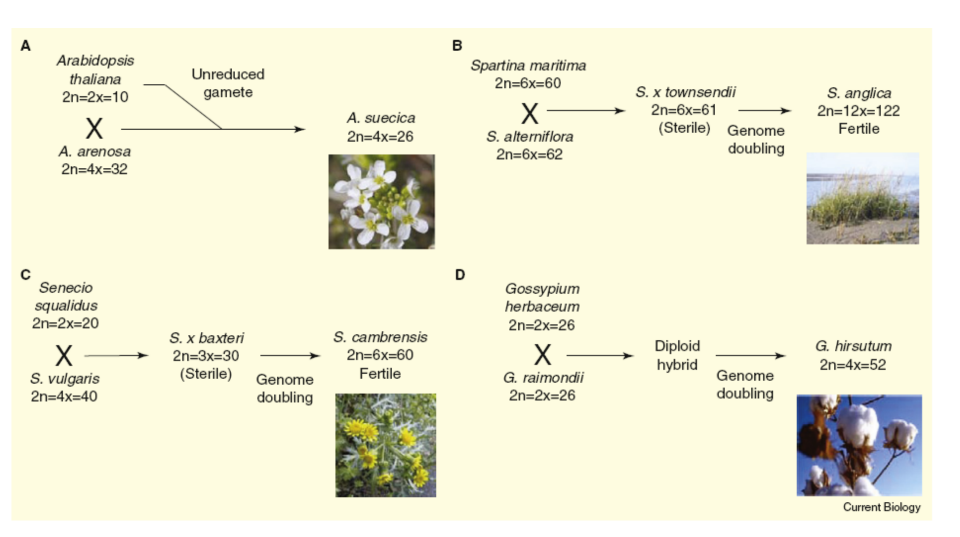
Data taken from REFS 2,7 for pairwise segmental duplications (SDs) with >90% identity. \*Given the fragmented nature of SDs in the draft chimpanzee genome, the duplication content can only be estimated indirectly on the basis of human duplication content, adjusting for detected differences in SD compared with chimpanzee whole-genome shotgun sequencing<sup>6</sup>. DNA not assigned to a chromosome was not included in these calculations. Consequently, in other genomes the estimate of recent duplication might rise as the quality of the sequence assembly improves. N.D., not determined.

## **Polyploidy (whole-genome duplication)**



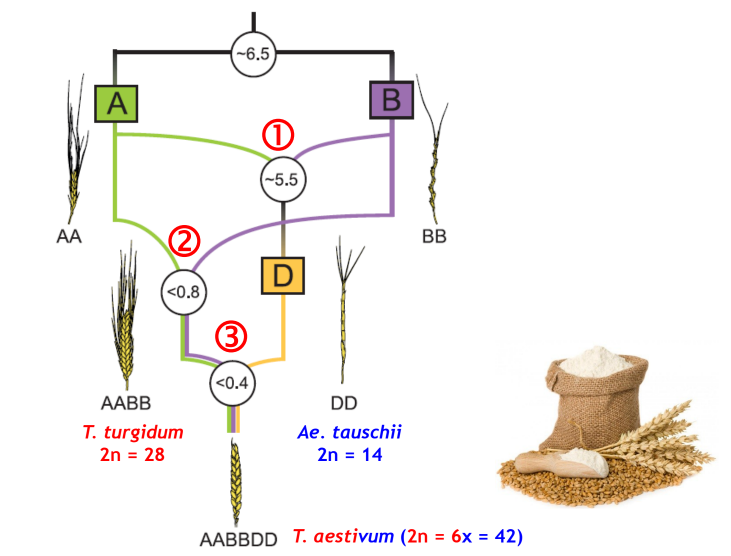
**Current Biol** 

### **Examples of allopolyploid speciation**



Phylogenomic history of bread wheat (Triticum aestivum; AABBDD).

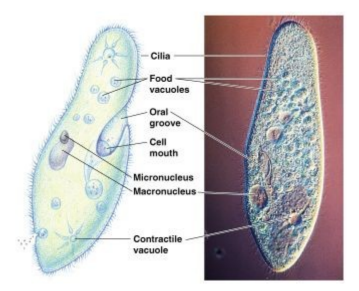
Three rounds of hybridization/polyploidy.

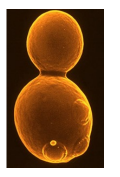


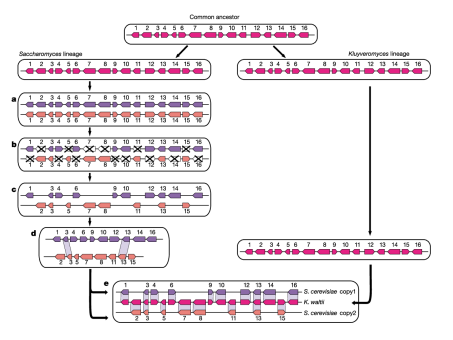
Marcussen et al. (2014), Science

#### Whole-genome duplications in protozoa

- the unicellular eukaryote *Paramecium tetraurelia*
- most of 40,000 genes arose through at least 3 successive whole-genome duplications
- most recent duplication most likely caused an explosion of speciation events that gave rise to the *P. aurelia* complex (15 sibling species)
- some genes have been lost, some retained
- many retained (duplicated) genes do not generate functional innovations but are important because of the gene dosage effect







#### Whole-genome duplications in yeast

genome comparison between two yeast
 species, Saccharomyces cerevisiae (n = 16) and
 Kluyveromyces waltii (n = 8)

- each region of *K. waltii* corresponding to two regions of *S. cerevisiae*
- the S. cerevisiae genome underwent a WGD after the two yeast species diverged

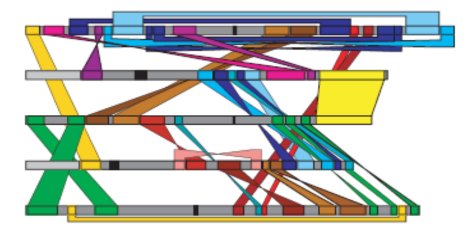
• in nearly every case (95%), accelerated evolution was confined to only one of the two paralogues (= one of the paralogues retained an ancestral function, the other was free to evolve more rapidly and acquired a derived function)



First evidence of a WGD in plants

# Analysis of the genome sequence of the flowering plant *Arabidopsis thaliana*

The Arabidopsis Genome Initiative\* AGI (2000)

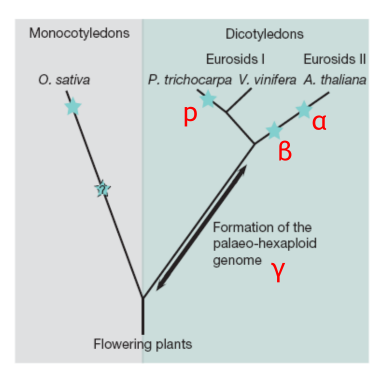


What does the duplication in the Arabidopsis genome tell us about the ancestry of the species? As the majority of the Arabidopsis genome is represented in duplicated (but not triplicated) segments, it appears most likely that Arabidopsis, like maize, had a tetraploid ancestor ...The diploid genetics of Arabidopsis and the extensive divergence of the duplicated segments have masked its evolutionary history.

# The grapevine genome sequence suggests ancestral hexaploidization in major angiosperm phyla

The French-Italian Public Consortium for Grapevine Genome Characterization\*

Nature 449, 2007

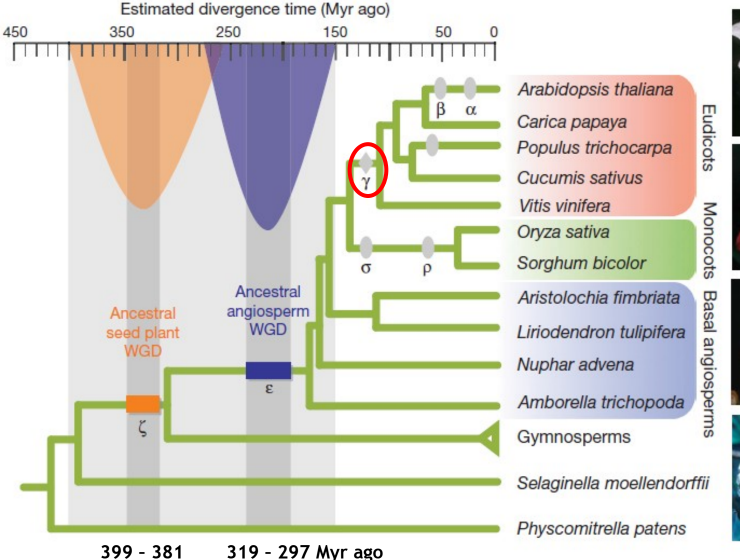


The formation of the palaeo-hexaploid ancestral genome occurred after divergence from monocots and before the radiation of the Eurosids. Star = a WGD (tetraploidization) event. The **γ triplication** may have been an ancient **auto-hexaploidy** formed from fusions of three identical genomes, or **allo-hexaploidy** formed from fusions of three somewhat diverged genomes.

Tang et al. 2008, Genome Res



### WGD events in seed plants and angiosperms



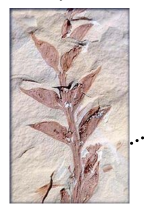


Jiao et al. (2011) Nature; Clark and Donoghue (2017) Proc R Soc

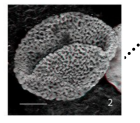
### Charles Darwin's abominable mystery solved (?)



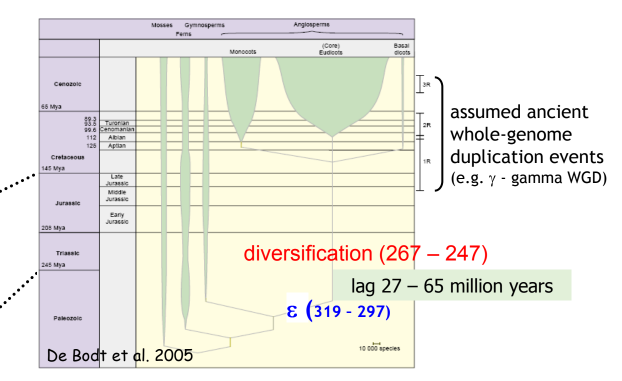
Archaefructus liaoningensis (140 million year old fossil)



Afropollis (245 million year old angiosperm pollen)

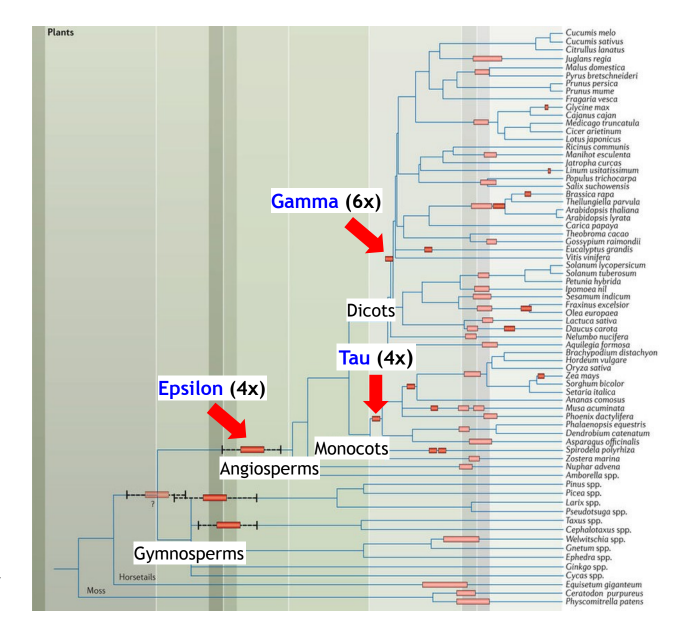


"The rapid development as far as we can judge of all the higher plants within recent geological times is an abominable mystery." (Charles Darwin in a letter to Sir Joseph Hooker, 1879)



Theres is evidence of ancient polyploidy throughout the major angiosperm lineages. It means that a genome-scale duplication event probably occurred PRIOR to the rapid diversification of flowering plants

## Multiple whole-genome duplications in evolution of land plants



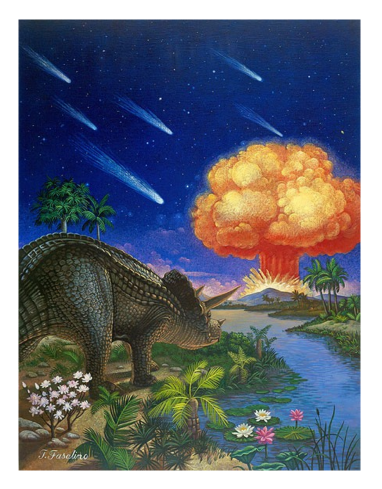
Van de Peer (2017) Nat Rev Genet (modified)

## Plants with double genomes might have had a better chance to survive the Cretaceous–Tertiary extinction event

Jeffrey A. Fawcett<sup>a,b,1</sup>, Steven Maere<sup>a,b,1</sup>, and Yves Van de Peer<sup>a,b,2</sup>

PNAS 106 (2009)

<sup>a</sup>Department of Plant Systems Biology, Flanders Institute for Biotechnology, 9052 Gent, Belgium; and <sup>b</sup>Department of Plant Biotechnology and Genetics, Ghent University, 9052 Gent, Belgium



Could WGD event(s) help plants to survive the mass extinction (one or more catastrophic events such as a massive asteroid impact) at the Cretaceous-Tertiary boundary ?

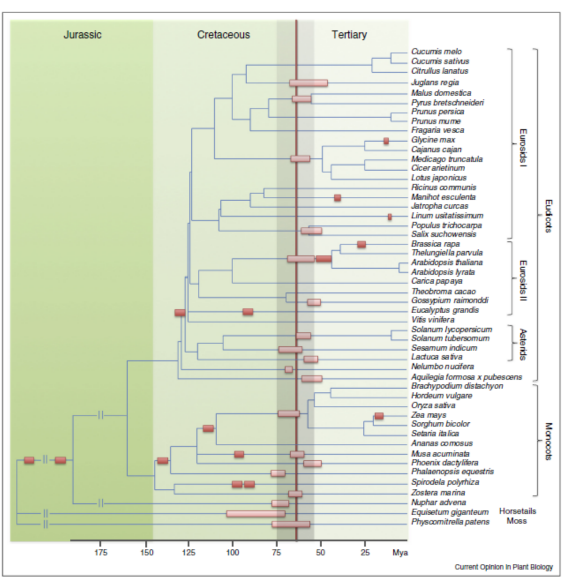


## Possible establishment of polyploid plants following the K/Pg mass extinction (66 million y. ago)

 WGDs clustered around the Cretaceous-Tertiary (KT) boundary

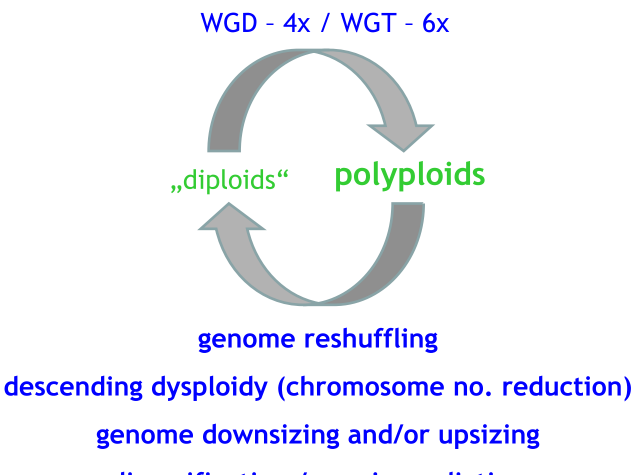
the KT extinction event the most recent mass extinction (one or more catastrophic events such as a massive asteroid impact and/or increased volcanic activity)

the KT extinction event extinction of 60% of plant species, as well as a majority of animals, including dinosaurs



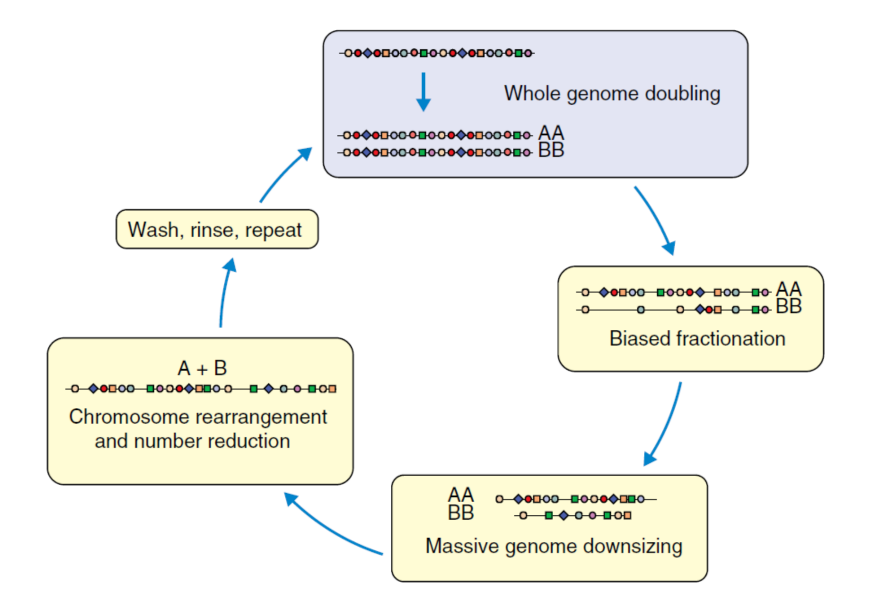
Lohaus and Van de Peer (2016) Curr Opin Pl Biol

#### **Polyploidization – Diploidization cycle**

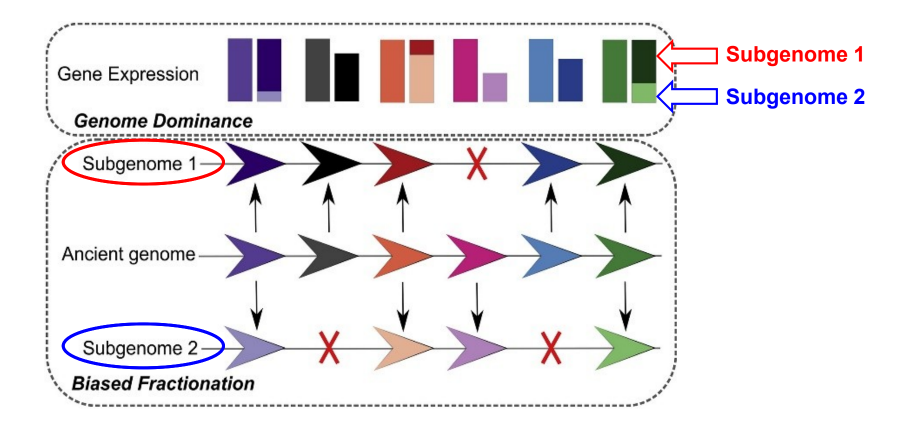


diversification / species radiation

#### Whole-genome duplication and diploidization



#### Genome diploidization: biased fractionation and (sub)genome dominance

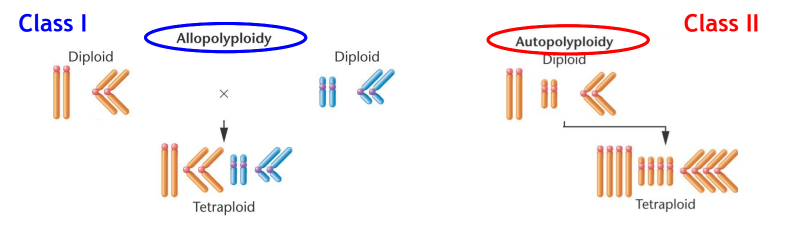


## Biased (sub)genome fractionation and dominance can be explained by the mode of polyploidization

Garsmeur et al. (2013) Mol Biol Evol

Species	WGD Class	Substitution Rate (Ks)	Bias Ratio between	Fractionation Pattern	Genome Dominance	Expression Data from
	Class	Race (RS)	Duplicate			
			Regions			
Medicago	Т	0.87	1.23	Biased	No data	
Sorghum	1	0.95	1.24	Biased (Schnable et al. 2012)	Yes	Dugas et al. (2011)
Arabidopsis	1	0.76	1.17	Biased (Thomas et al. 2006)	Yes	Gan et al. (2011)
Brassica	1	0.34	1.47	Biased (Wang et al. 2011)	Yes (Cheng et al. 2012)	
Maize	1	0.17	1.46	Biased (Woodhouse et al. 2010)	Yes (Schnable et al. 2011)	
Poplar	П	0.23	1.05	Unbiased	No data	
Soybean	Ш	0.15	1.03	Unbiased	No	Schmidt et al. (2011)
Banana	Ш	0.39	1.06	Unbiased	No	D'Hont et al. (2012) and supplementary table S4, Supplementary Material onlin

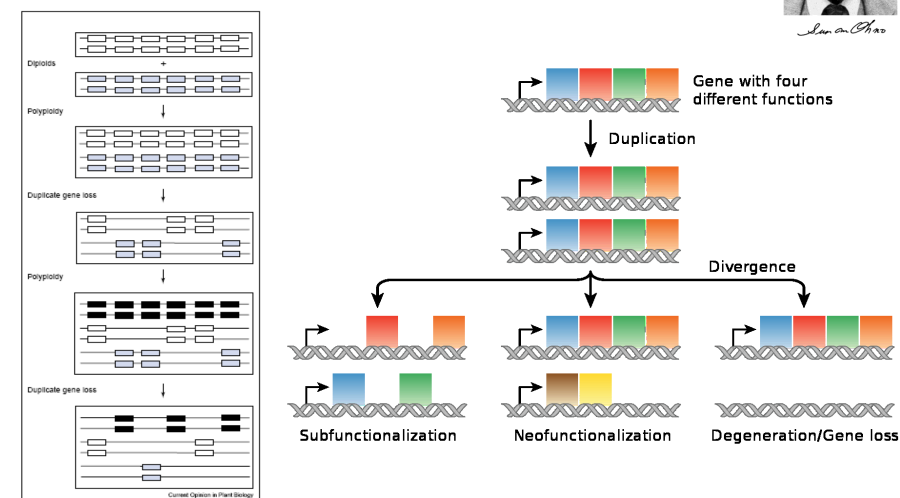
Table 2. Fractionation Pattern and Genome Dominance in Eight Species.



#### The fate of duplicated genes

#### Genome evolution through cyclic

#### WGD and diploidization



Adams and Wendel (2005)

Genome size decrease (downsizing)

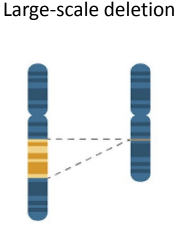
- recombination
- chromosome rearrangements

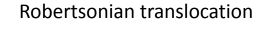
### Genome size decrease (downsizing)

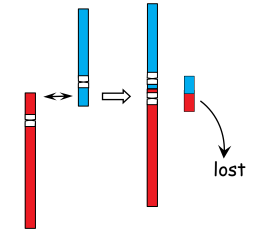
Recombinational deletions after double-strand breaks (DSBs) - DSB repair

- unequal homologous recombination including unequal crossing-over
- illegitimate recombination (non-homologous end joining, NHEJ)

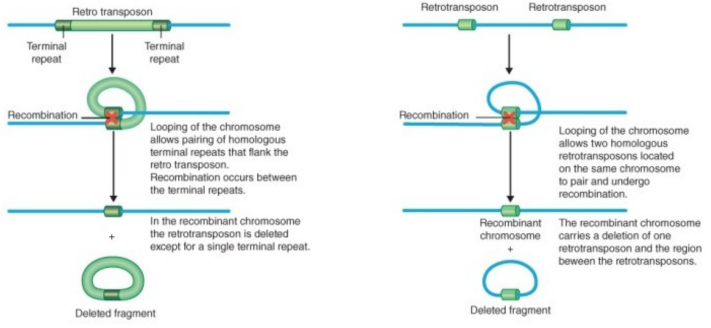
#### Chromosome rearrangements (...in principle again DSBs and recombination)







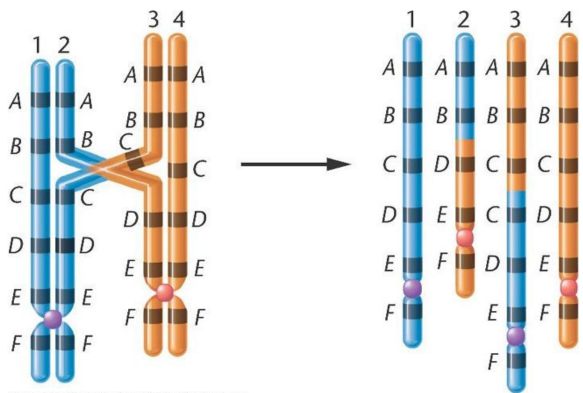
## Genome size decrease by unequal homologous recombination between two LTRs or between two LTRretrotransposons





~70% of retrotransposon sequences in the *A*. *thaliana* genome are no longer autonomous: solo LTRs = probably the consequence of unequal homologous recombination = inactive, truncated elements cannot contribute to genome expansion

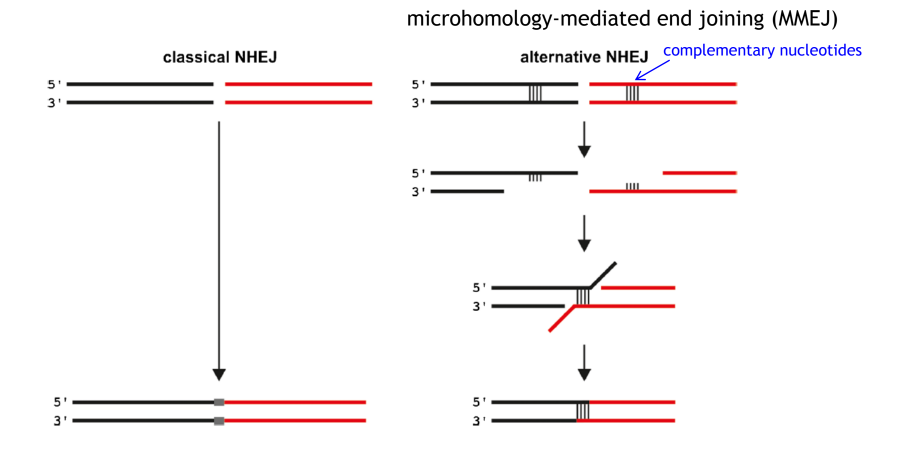
## **Deletion through unequal crossing-over**



This tetrad is mispaired at meiotic synapsis.

The result, after crossing over, is two unequal chromosomes: one with a duplication (3) and one with a deletion (2).

## Two main pathways of non-homologous end joining (NHEJ)



## DNA lost

(but some DNA can be inserted - filler DNA)

## NHEJ in plant somatic cells

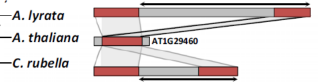
- NHEJ seems to be the main mode of DSB repair in higher eukaryotes
- NHEJ might lead, in some cases, to genomic changes (deletions, insertions or various kinds of genomic rearrangements)
- genomic alterations in meristematic cells can be transferred to the offspring
- alternative NHEJ can mediate genome size loss

Arabidopsis vs. tobacco (genome size larger in tobacco)

- tobacco: almost every second deletion event is accompanied by the insertion of filler sequence
- Arabidopsis: no insertions
- overall length of the deletions is about one-third shorter in tobacco than in Arabidopsis

>>> inverse correlation between genome size and the medium length of deletions
>>??? species-specific differences in DSB repair pathways can contribute to the evolution of eukaryotic genome size ???

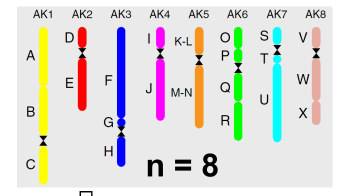
- *A. thaliana* (157 Mb) has lost  $\mathbf{6}$ × more introns than *Arabidopsis lyrata* (210 Mb) since the divergence of the two species but gained very few introns

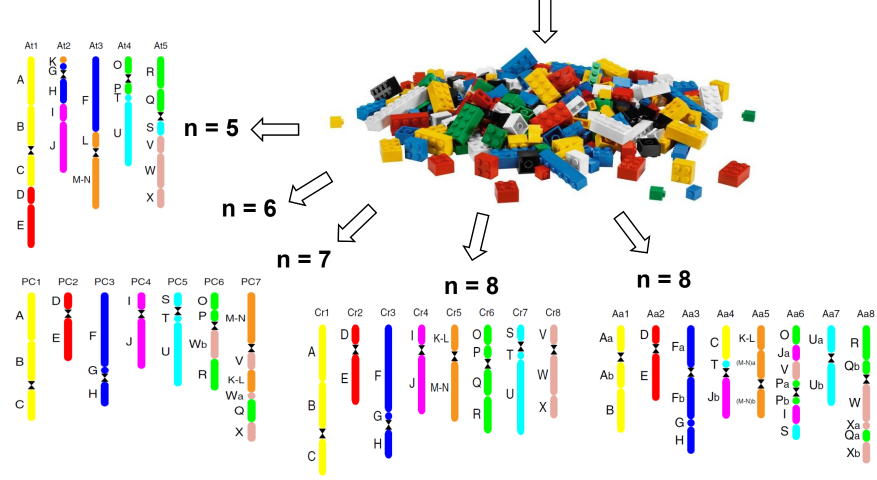


1C = 157 Mt

1C = 4.5 Gb

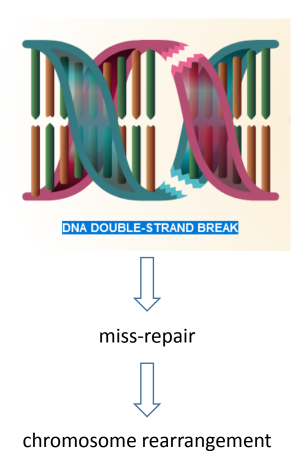
## Chromosome number variation: chromosome rearrangements





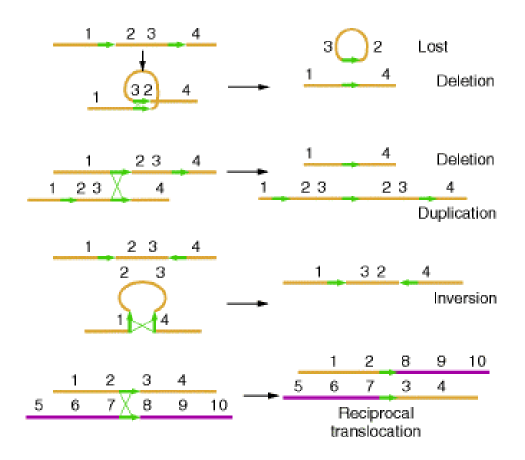
## Chromosome rearrangements results from double-strand breaks and their miss-repair

DSB



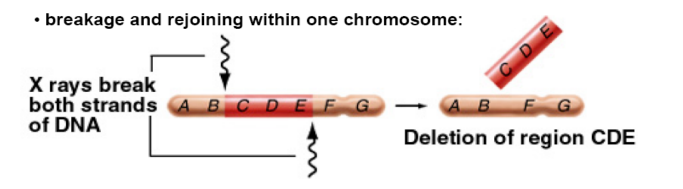
#### **Chromosome rearrangements – the role of repeats**

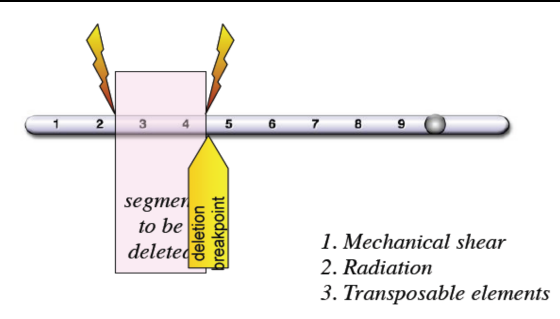
In organisms with repetitive DNA, homologous repetitive segments **within one chromosome** or **on different chromosomes** can act as sites of DSBs and their missrepair, i.e. non-allelic homologous recombination.



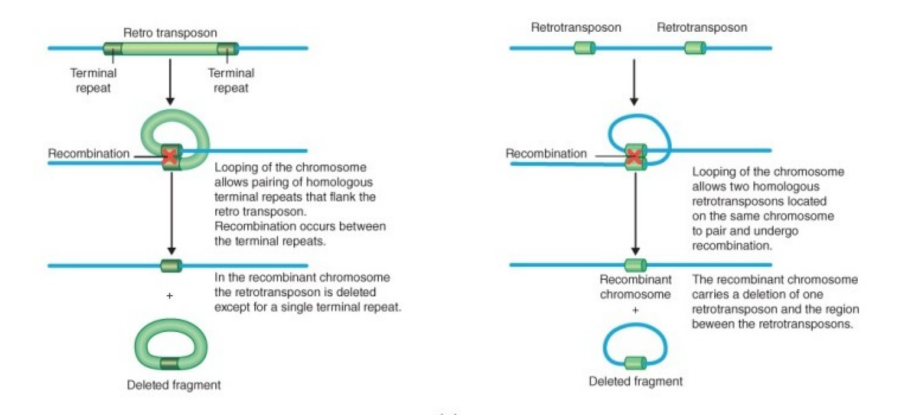
#### Deletion formation by breakage and rejoining

- = deficiencies = losses of chromosome segments
- can occur terminally or internally, e.g. caused by...



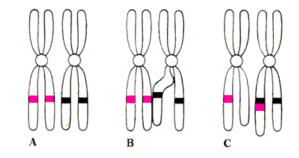


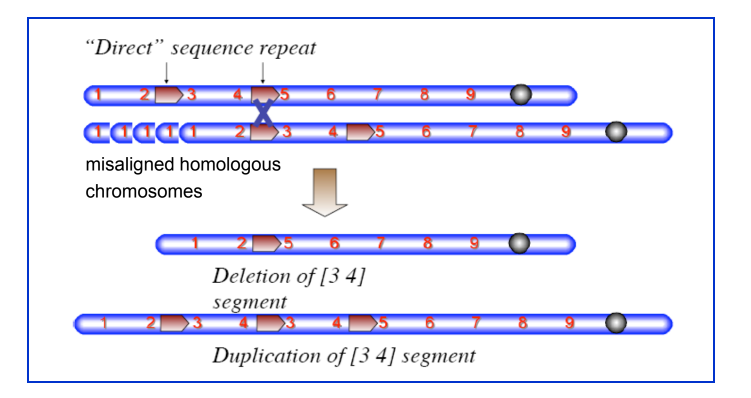
### Deletion formation by intra-chromosomal (unequal) recombination



#### Deletion (and duplication) formation by unequal cross-over

Sometimes during meiosis two chromatids from homologous chromosomes (A) are misaligned during a cross-over event (B) as a result, one chromatid gained a duplicated region and the another lost a deleted region (C). The duplication as well as the deletion are inherited by resulting gametes.

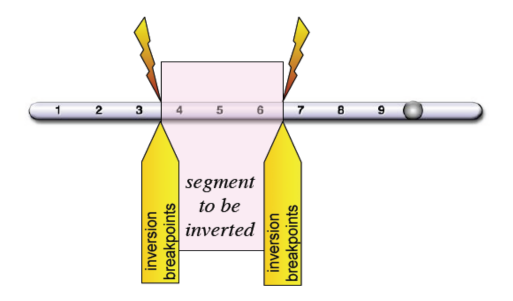


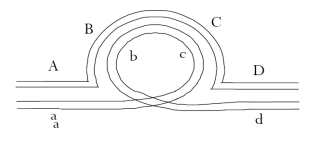


#### Inversions

Inversions as balanced rearrangements are generally viable and show no particular abnormalities at the phenotypic level. Many inversions can be made homozygous.

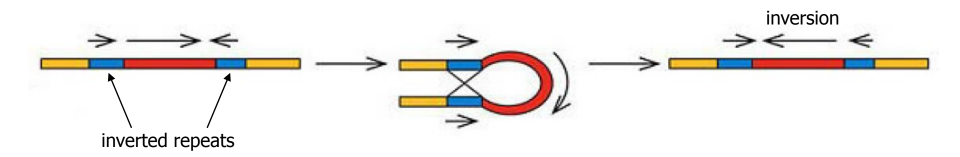
**Inversion heterozygote** - cells that contain one normal haploid chromosome set plus one set carrying the inversion. Microscopic observation of meioses in inversion heterozygotes reveals an **inversion loop**.



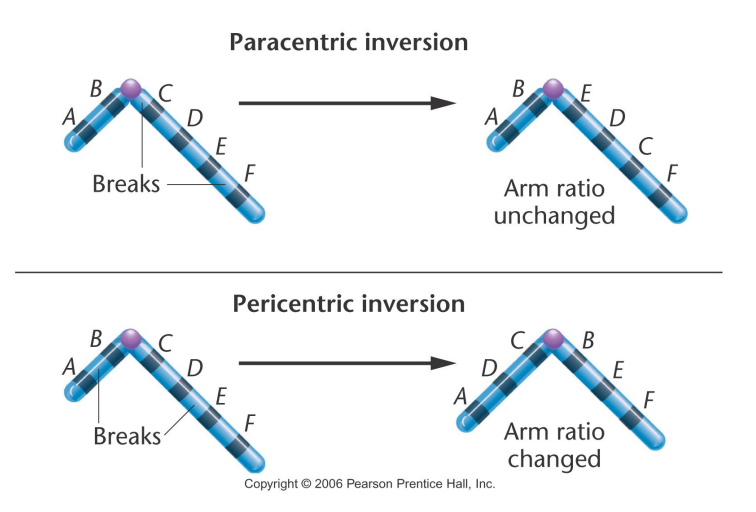


meiotic inversion loop

#### Inversion formation by intra-chromosomal recombination



#### Two types of inversions

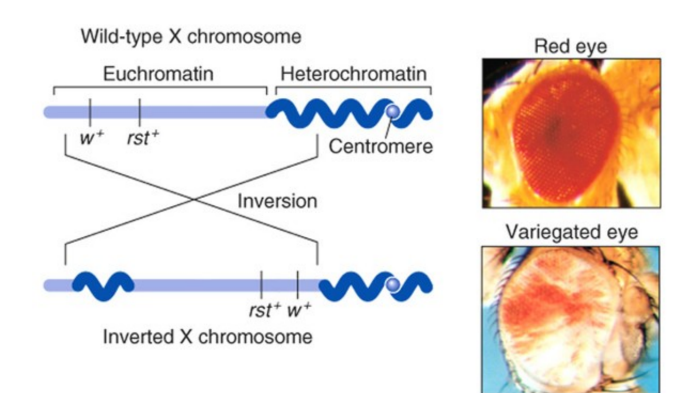


mechanism of inversion formation: breakage and rejoining

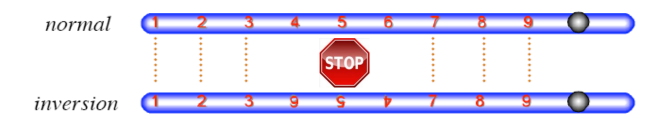
### Inversions and recombination: evolutionary significance

Can be "adaptive" when it stabilizes/disrupt a superior combination of alleles on a chromosome (examples seen in *Drosophila*)

## **Position-effect variegation**

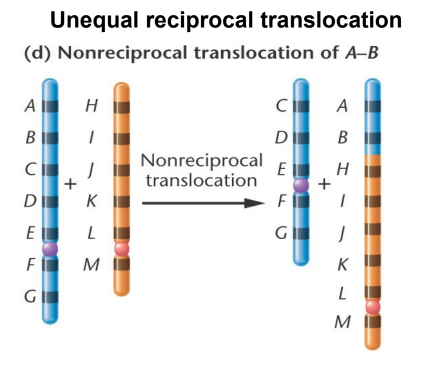


#### **Inversions may suppress recombination**

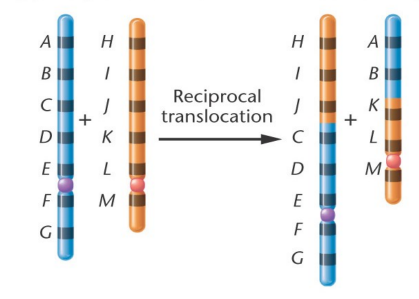


Chromosome rearrangements (typically inversions) may reduce gene flow by suppressing recombination. Inversions allow genes located in these regions to differentiate, in contrast to genes in freely recombining collinear regions.

### **Reciprocal translocations**



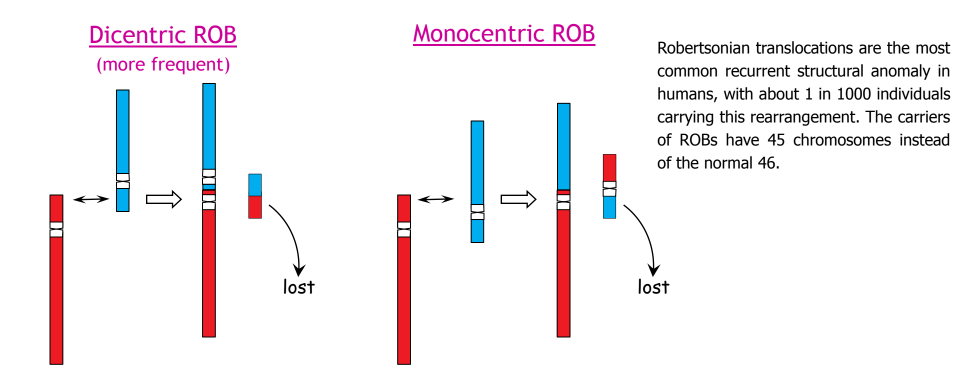
attachment of chromosome fragment to a non-homologous chromosome (leading to deletions and duplications in the progeny) (e) Reciprocal translocation of A-B and H-I-J



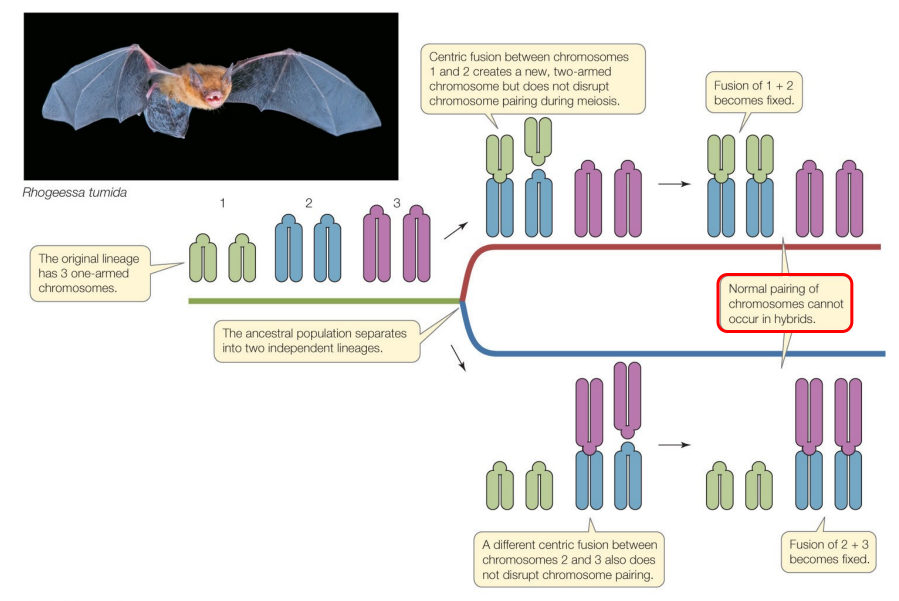
exchange of chromosome fragments between non-homologous chromosomes

#### Robertsonian translocations - ROBs (centric "fusions")

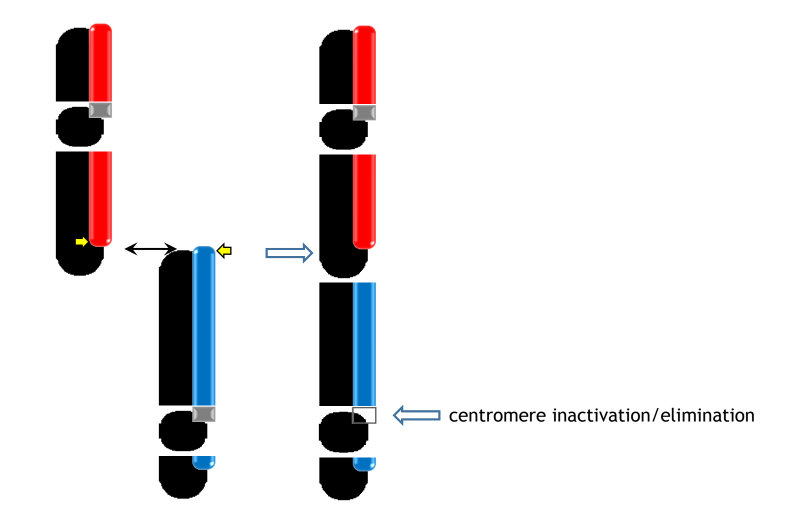
- type of a reciprocal translocation between two acrocentric/telocentric chromosomes
- also called whole-arm translocations or centric-fusion translocations
- named after the American insect geneticist W. R. B. Robertson, who first described a Robertsonian translocation in grasshoppers in 1916
- evolutionary significance >>> <u>chromosome number reduction</u> (from 2 acrocentric chromosomes one metacentric chromosome)



#### Speciation by Robertsonia translocations ("centric fusions")



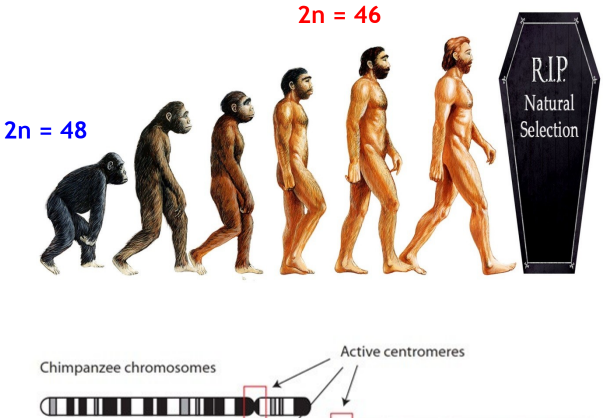
#### End-to-end chromosome translocations ("chromosome fusions")

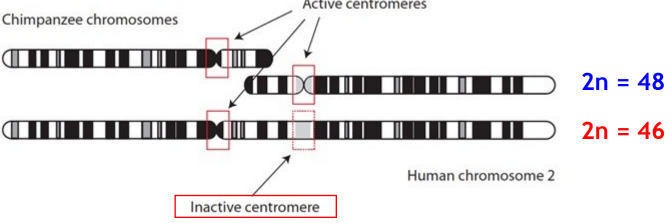


In principle unequal reciprocal translocation with breakpoints in (sub)telomeric regions.

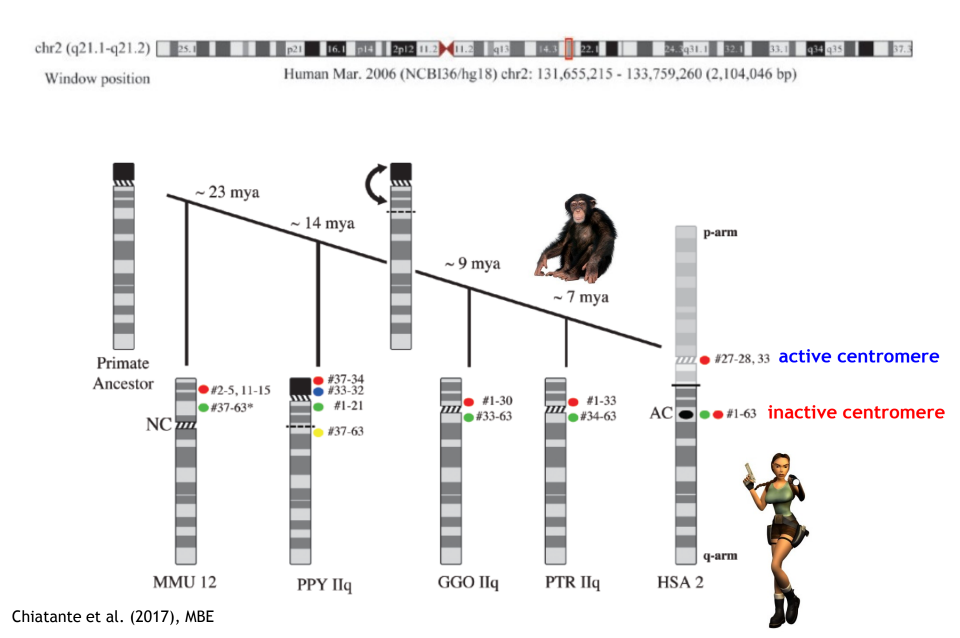
The second translocation product is minute and eliminated.

Chromosome "fusion" – the origin of the human (dicentric) chromosome 2

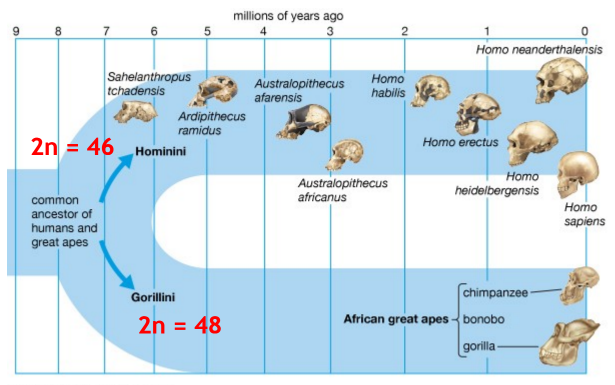




#### Chromosome "fusion" – the origin of the human (dicentric) chromosome 2



## Did the origin of "fusion" chromosome 2 contributed to reproductive isolation of hominid species from great apes?



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- different no. of chromosomes  $\rightarrow$  reproductive isolation
- loss of gene(s)  $\rightarrow$  adaptive advantage
- gene linkage? changed regulation of gene expression?

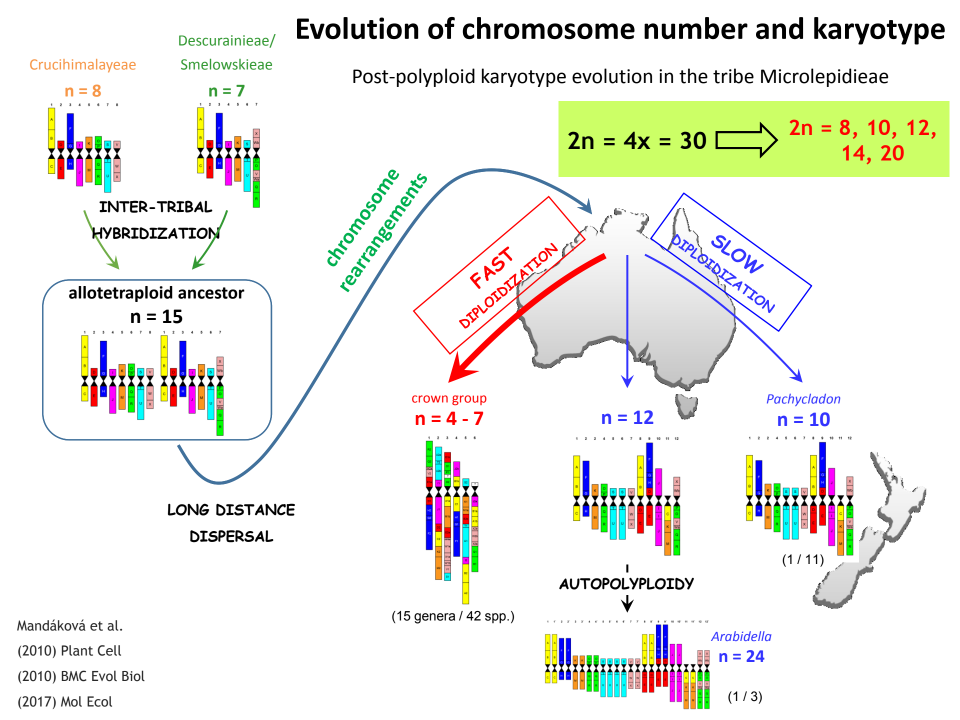
## **Evolution of chromosome number and karyotype**

Allopolyploid origin and diploidization in the tribe Microlepidieae (Brassicaceae)

- Australia: 15 genera, 47 species
- New Zealand: *Pachycladon*, 11 species
- chromosome number variation (from n = 4 to n = 24)







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