

# Genetic Variation: What it is and how to summarize it

# Genetic Variation

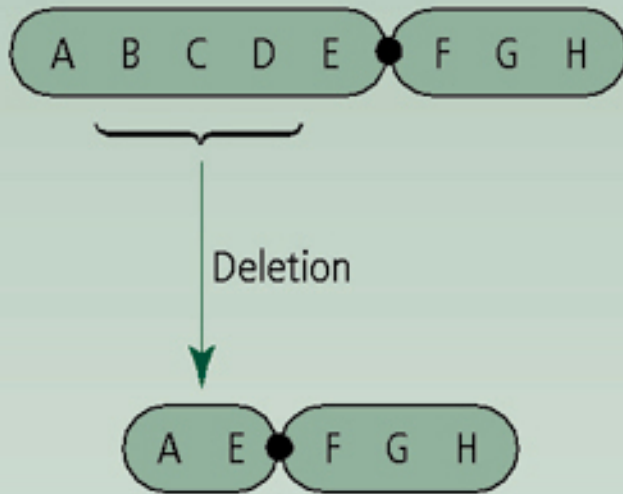
- Chromosomal variation
- Protein variation
- DNA variation

# Chromosomal Variation

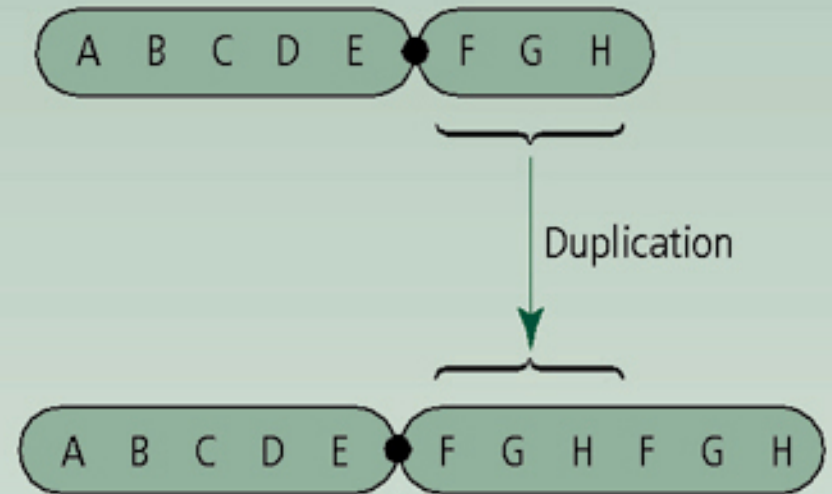
- Variation in chromosome number, gene number, gene order etc.

# Chromosomal rearrangements

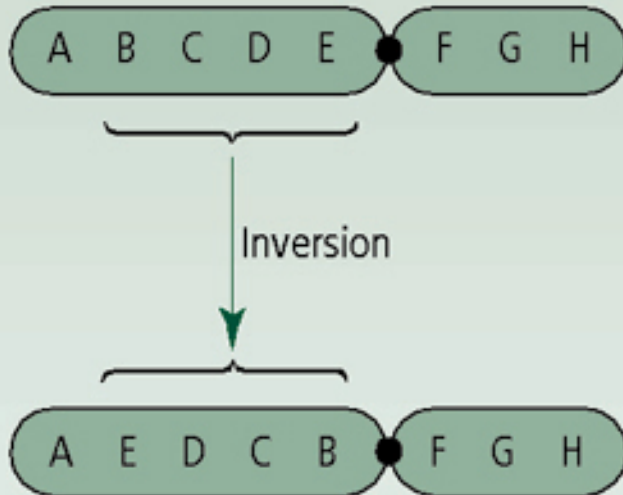
(a)



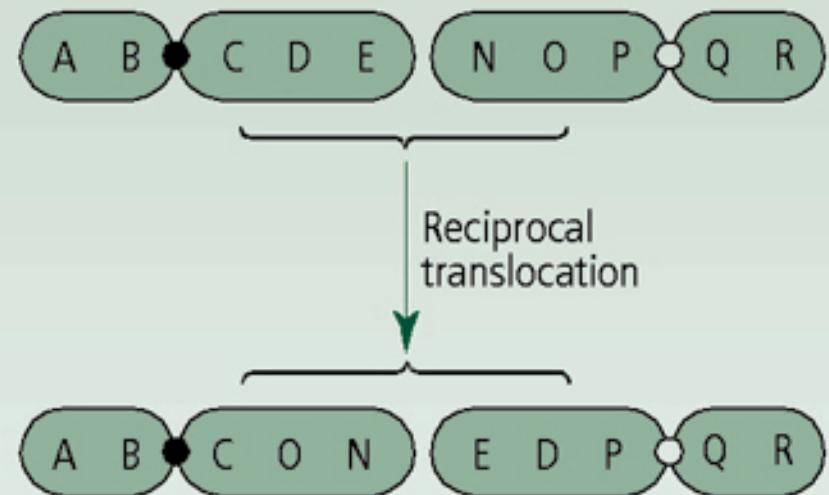
(b)



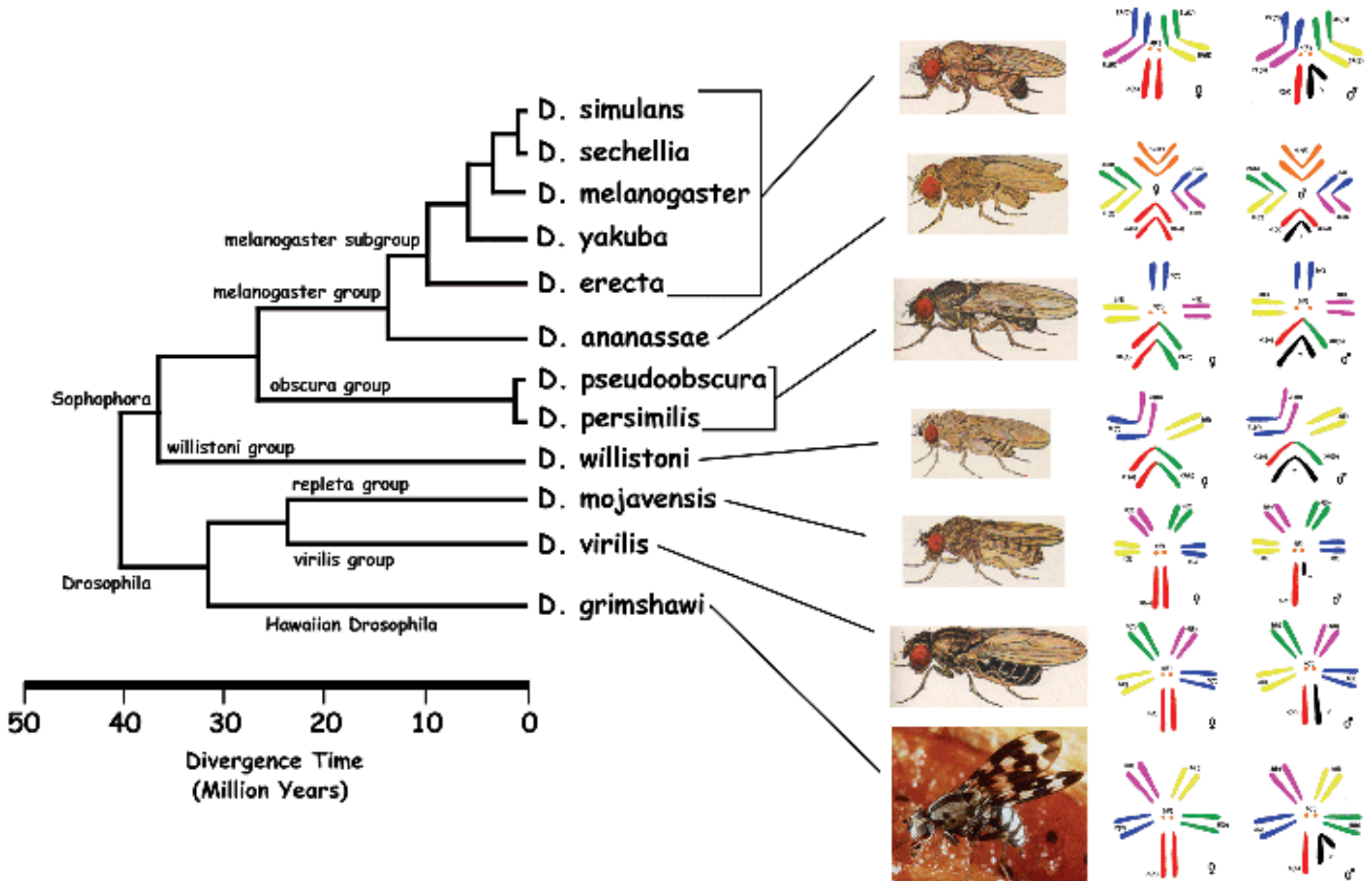
(c)



(d)



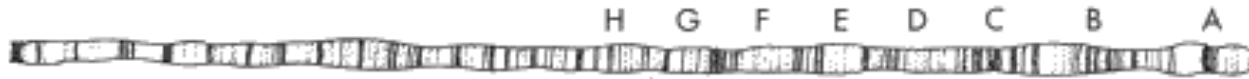
# Chromosomal Variation



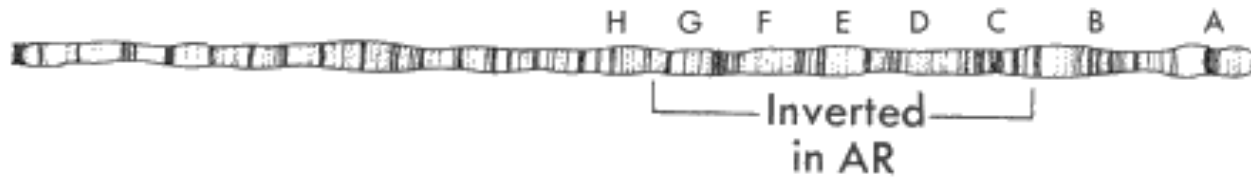
# Chromosomal Variation



# Chromosomal Variation

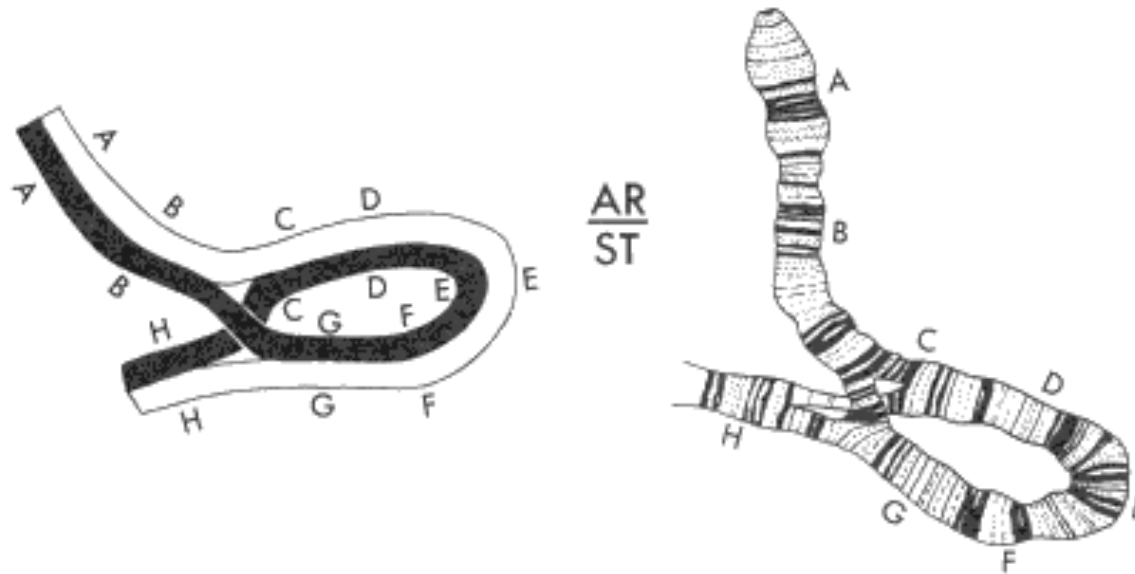
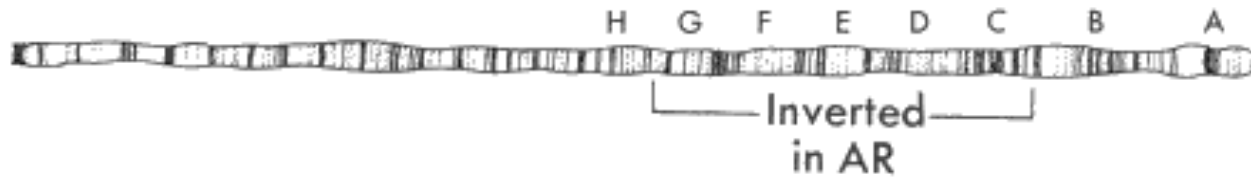


# Chromosomal Variation





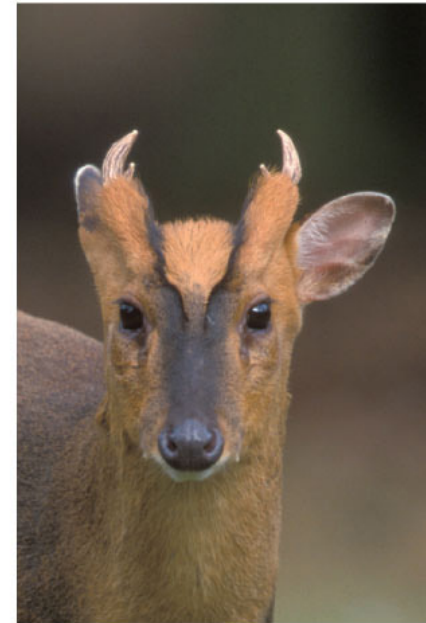
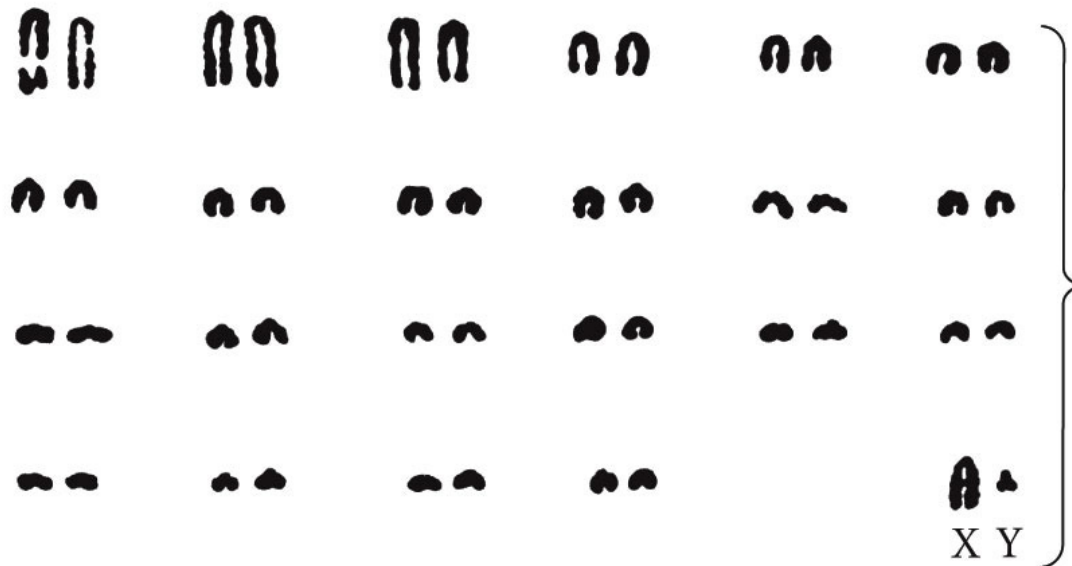
# Chromosomal Variation



From Dobzhansky and Sturtevant

# Chromosomal number changes

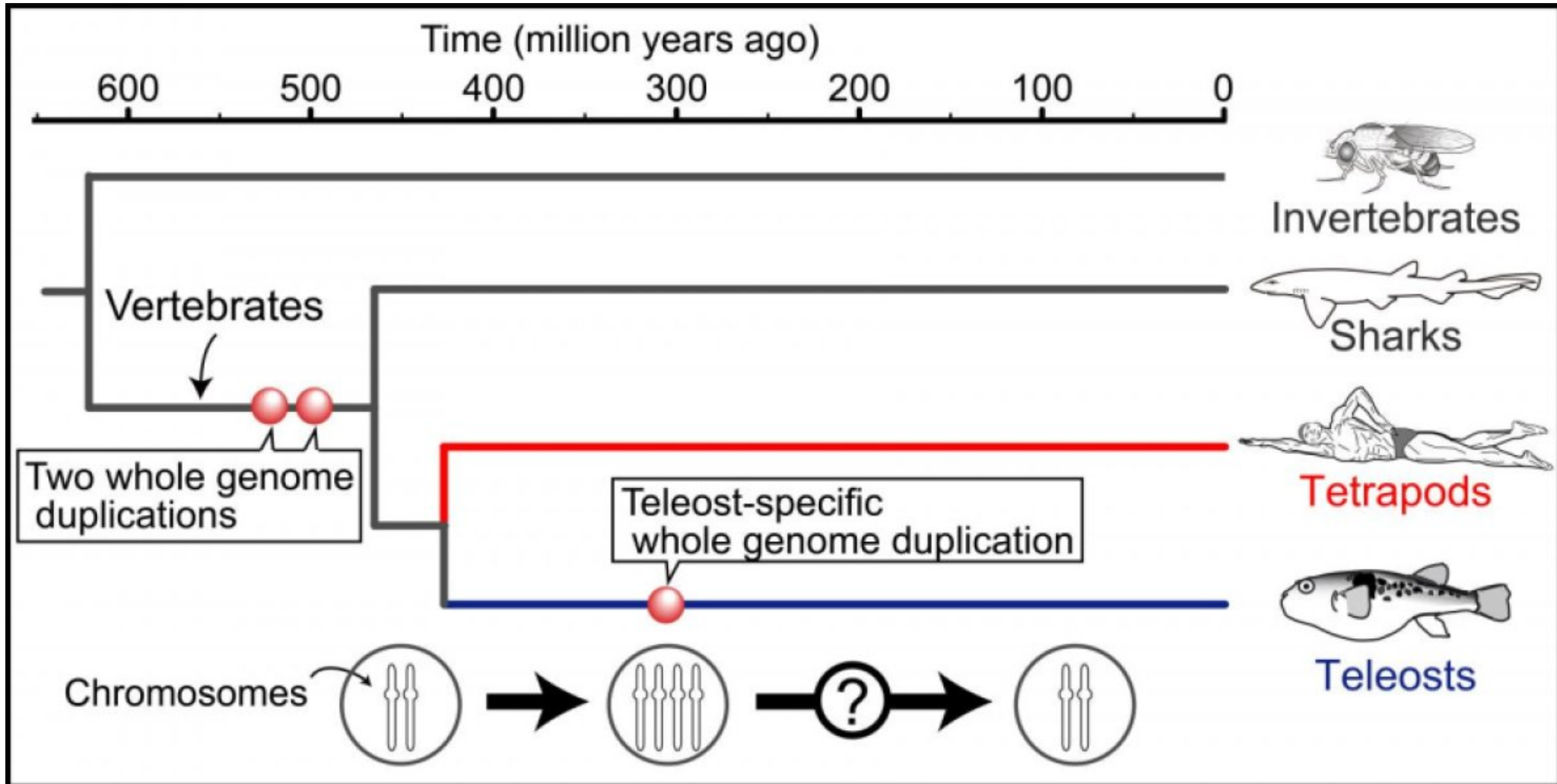
*Muntiacus reevesii* ( $2N = 46$ )



*Muntiacus muntiacus* ( $2N = 8$ )

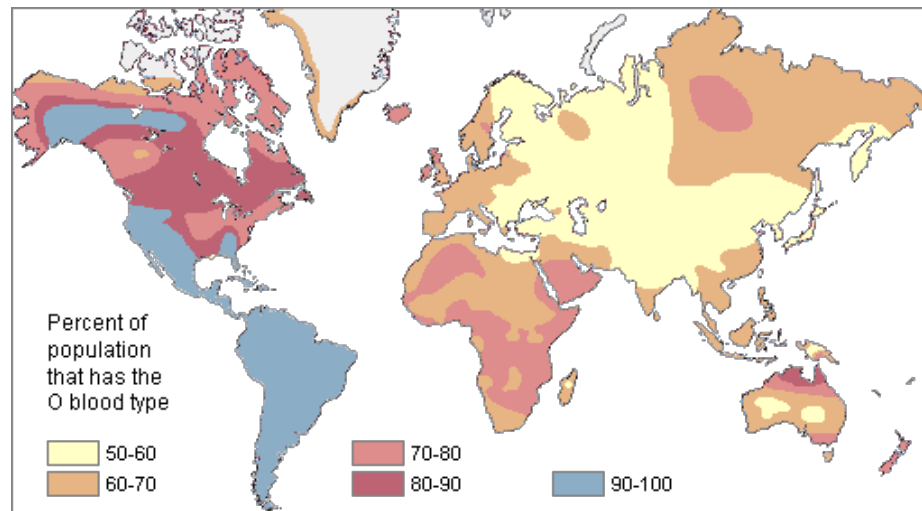
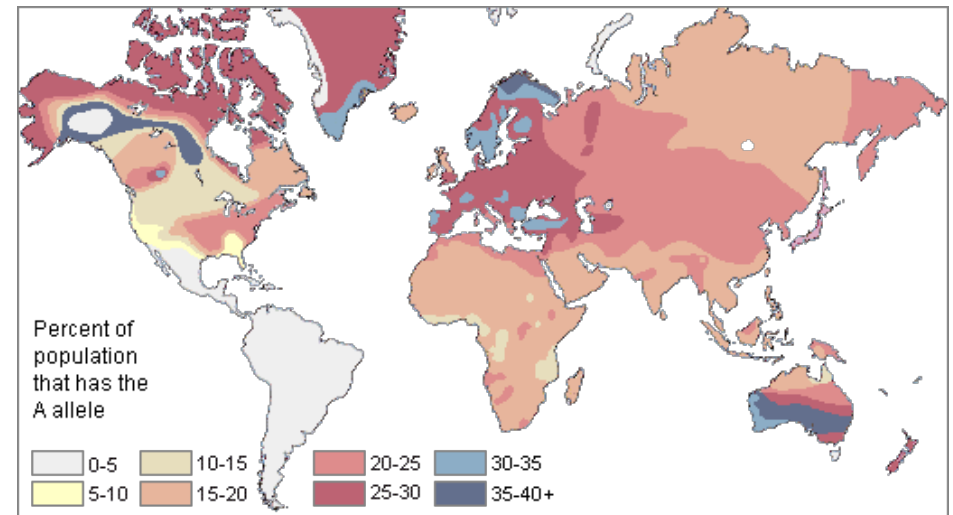
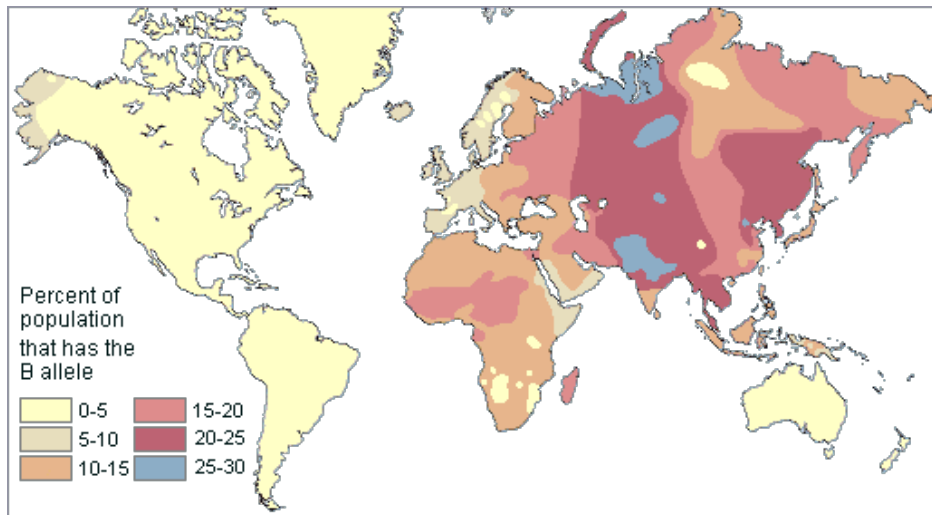


# Genome Duplication



# Protein variation

- ABO blood groups



# Protein variation

- Allozymes: variant forms of an enzyme encoded by different alleles at the same locus
- Variation revealed using electrophoresis

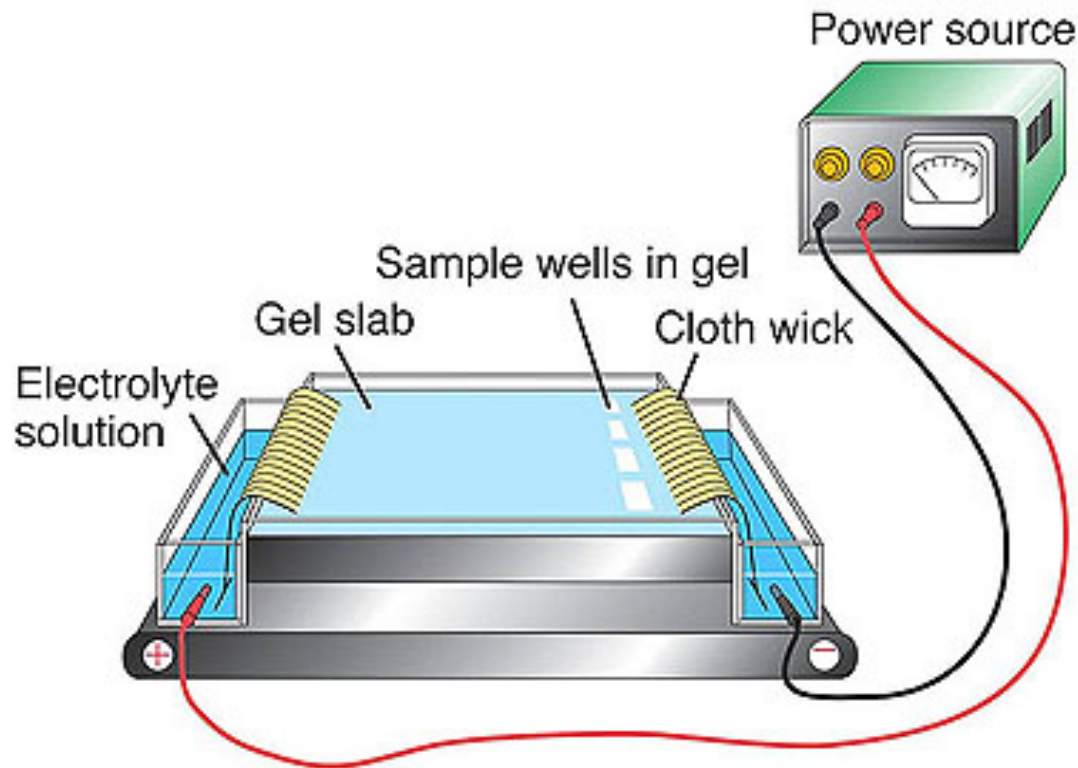
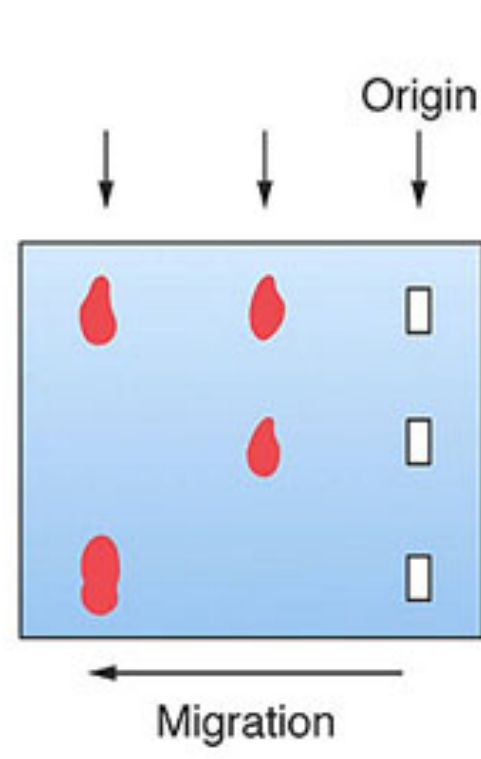


Figure 4: Schematic of devices used in protein electrophoresis

# Protein variation

- Allozymes: variant forms of an enzyme encoded by different alleles at the same locus
- Variation revealed using electrophoresis

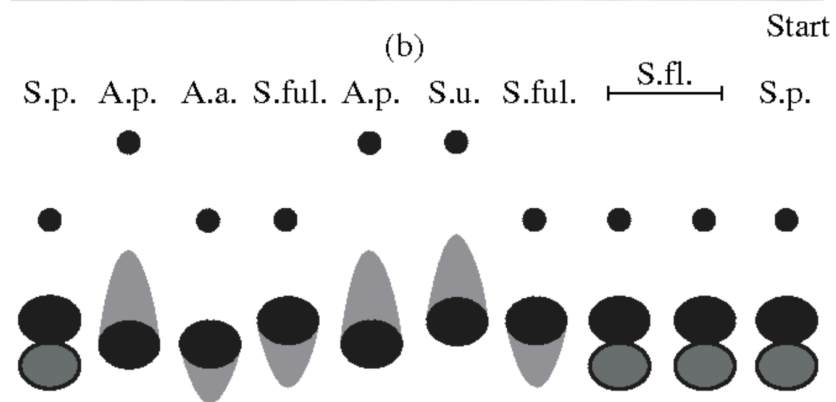
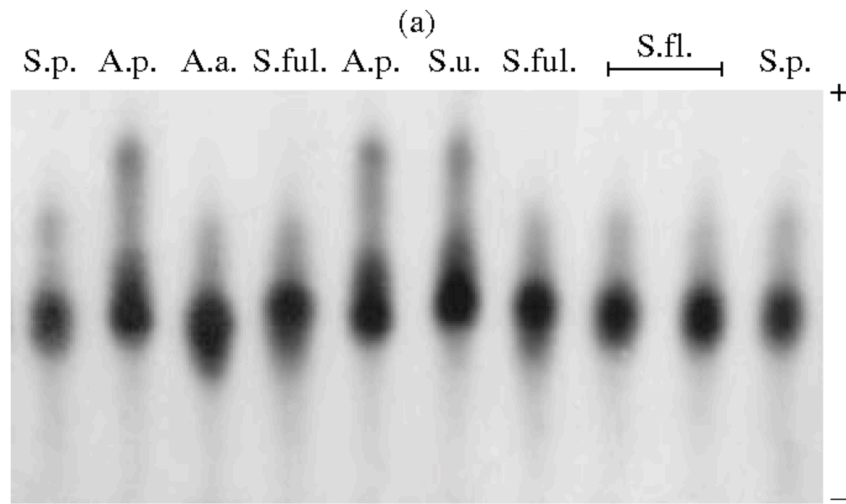


# How does it work?

- Nonsynonymous mutations can change enzyme's overall ionic charge
- Leads to differences in electrophoretic mobility

Amino Acid	3-Letter	1-Letter	Side chain polarity	Side chain charge (pH 7.4)
Alanine	Ala	A	nonpolar	neutral
Arginine	Arg	R	polar	positive
Asparagine	Asn	N	polar	neutral
Aspartic acid	Asp	D	polar	negative
Cysteine	Cys	C	nonpolar	neutral
Glutamic acid	Glu	E	polar	negative
Glutamine	Gln	Q	polar	neutral
Glycine	Gly	G	nonpolar	neutral
Histidine	His	H	polar	positive(10%), neutral(90%)
Isoleucine	Ile	I	nonpolar	neutral
Leucine	Leu	L	nonpolar	neutral
Lysine	Lys	K	polar	positive
Methionine	Met	M	nonpolar	neutral
Phenylalanine	Phe	F	nonpolar	neutral
Proline	Pro	P	nonpolar	neutral
Serine	Ser	S	polar	neutral
Threonine	Thr	T	polar	neutral
Tryptophan	Trp	W	nonpolar	neutral
Tyrosine	Tyr	Y	polar	neutral
Valine	Val	V	nonpolar	neutral

# Protein variation

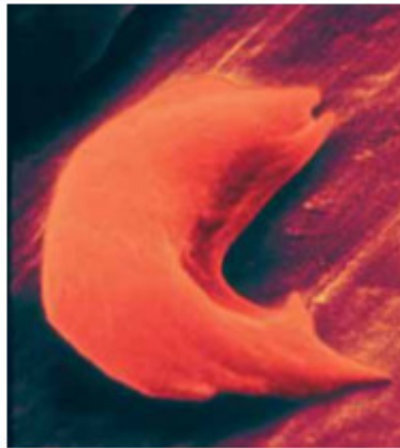


- HB-1
- HB-2
- HB-3
- HB-4

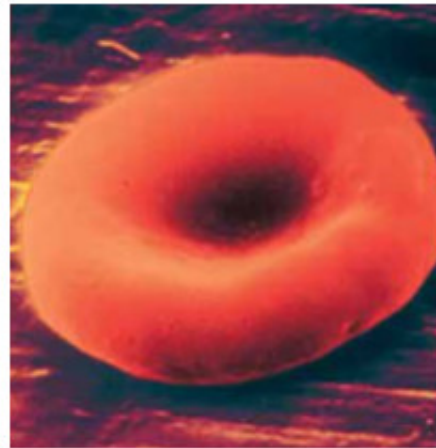


# Protein variation

- Amino acid variation
  - Alternative forms of proteins arising from variation in the amino acid sequence
    - ❖ Sickle-cell disease (HbS): ONE amino acid change in beta-globin chain of hemoglobin



Sickle-cell phenotype



Normal phenotype

# DNA variation

- RFLP: Restriction fragment length polymorphism
  - Created by mutation that changes a restriction site

GCCG**C**ATTCTA  
CGGC**G**TAAGAT

GCCG**A**ATTCTA  
CGGC**T**TAAGAT

# DNA variation

- RFLP: Restriction fragment length polymorphism

– Created by mutations that alter the restriction site



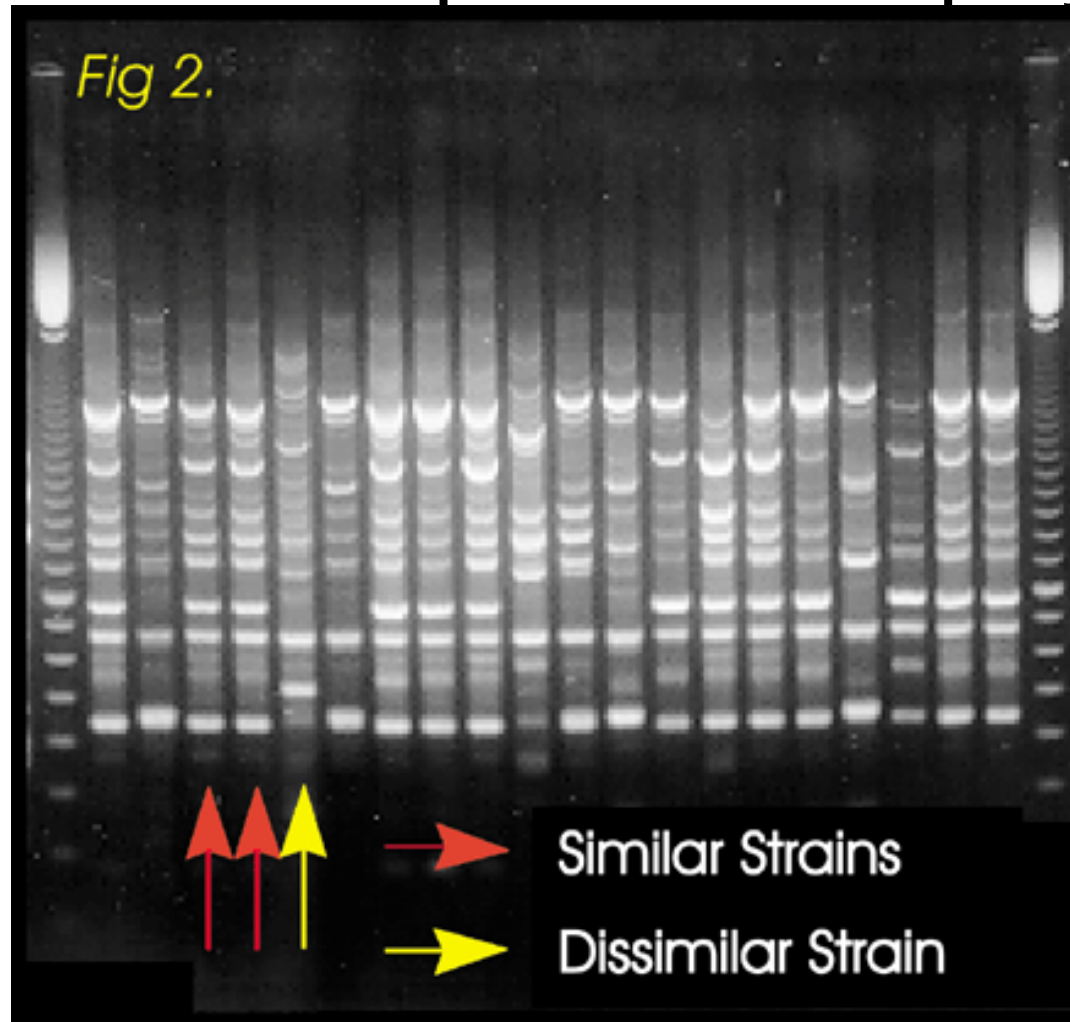
# DNA variation

- RFLP
- RAPD: Random amplification of polymorphic DNA
  - Like PCR, but segments are amplified randomly
  - Employs several arbitrary, short primers
  - Need no knowledge of underlying sequence
  - Variation in RAPD profile comes from variation in primer binding sites across individuals

# DNA variation

- RFLP
- RAPD: Random amplification of polymorphic DNA

Strains of  
Lactobacillus  
from 18  
types of  
Cheddar  
Cheese

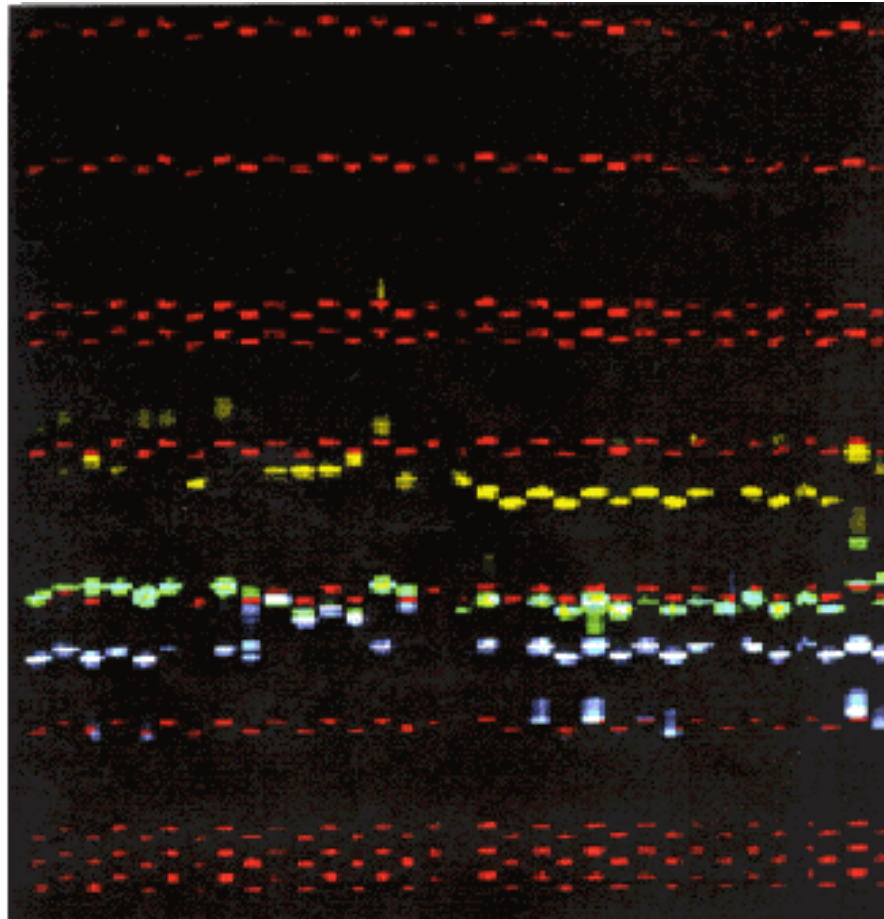






# DNA variation

- RFLP
- RAPD
- Microsatellites/minisatellites/VNTRs/SSRs:



Leishmania  
(Viannia) isolates

From London School of  
Hygiene and Tropical  
Medicine



# DNA variation

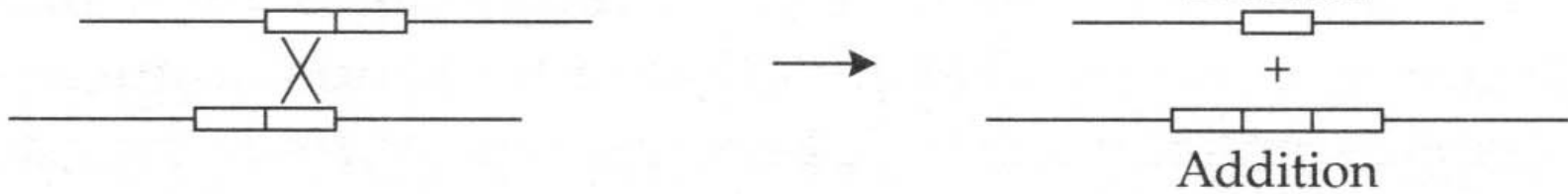
- RFLP
- RAPD
- Microsatellites/minisatellites/VNTRs/SSRs
- Insertion/Deletion: Gain or loss of DNA segment

# Small Scale Insertion & Deletions (InDels)

(a)



(b)



# DNA variation

- RFLP
- RAPD
- Microsatellites/minisatellites/VNTRs/SSRs
- Insertion/Deletion
- Single Nucleotide Polymorphism: Differences at a single nucleotide



# Discrete Genetic Variation

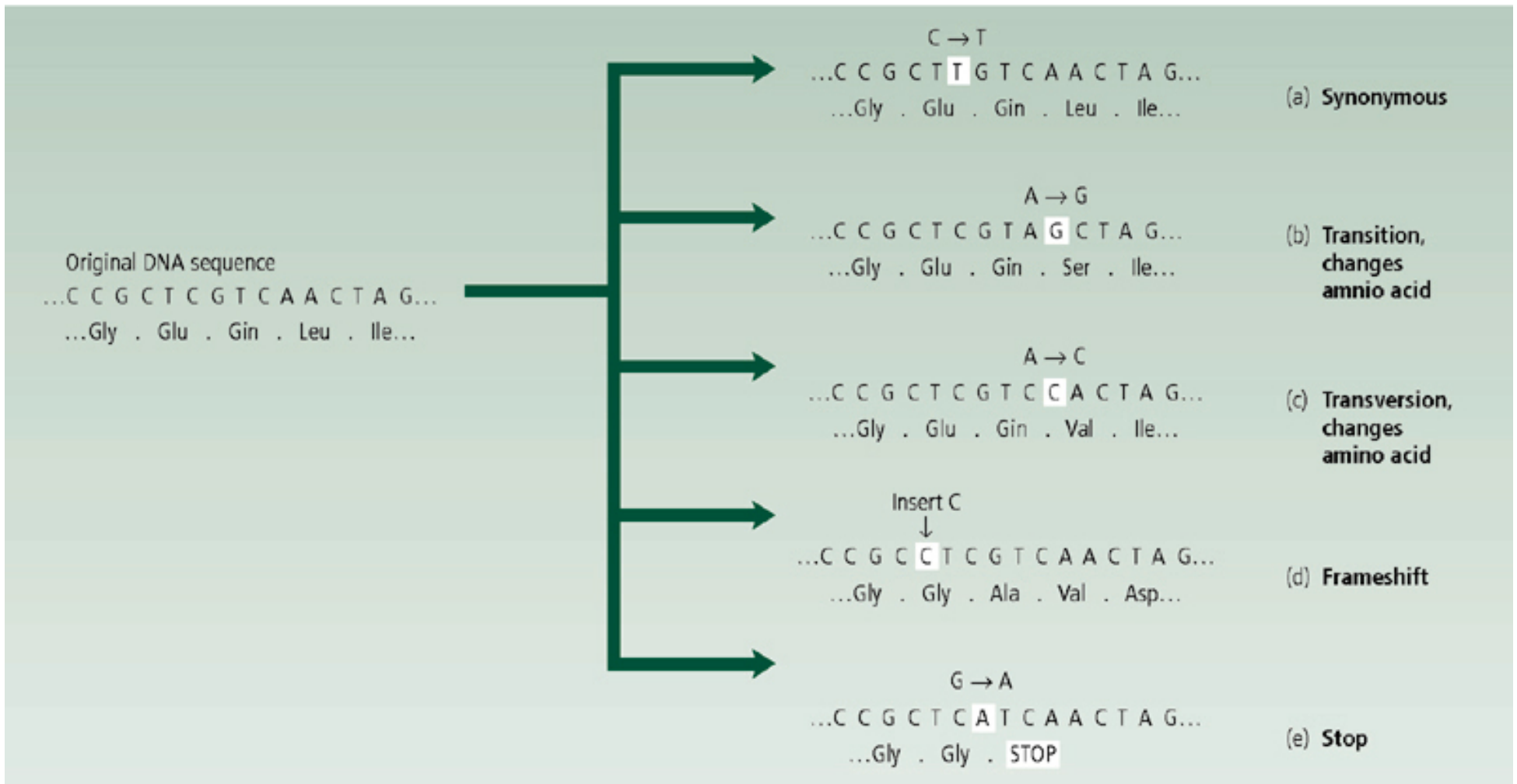
- Chromosomal variation
  - Inversions, chromosomes fusions/fissions
- Protein variation
  - Immunological, allozymes, amino acid variation
- DNA variation
  - RFLP, RAPD, VNTR, Indel, SNP

Where does genetic variation come from?

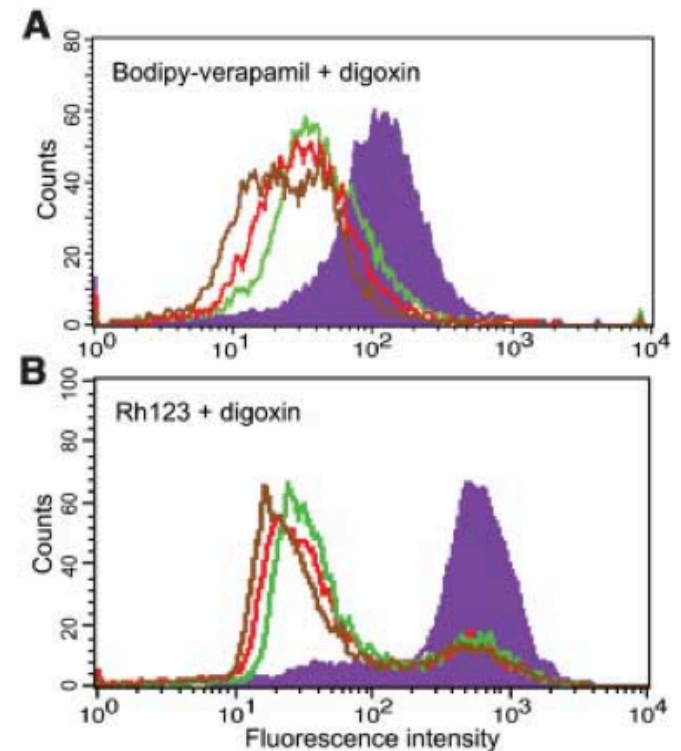
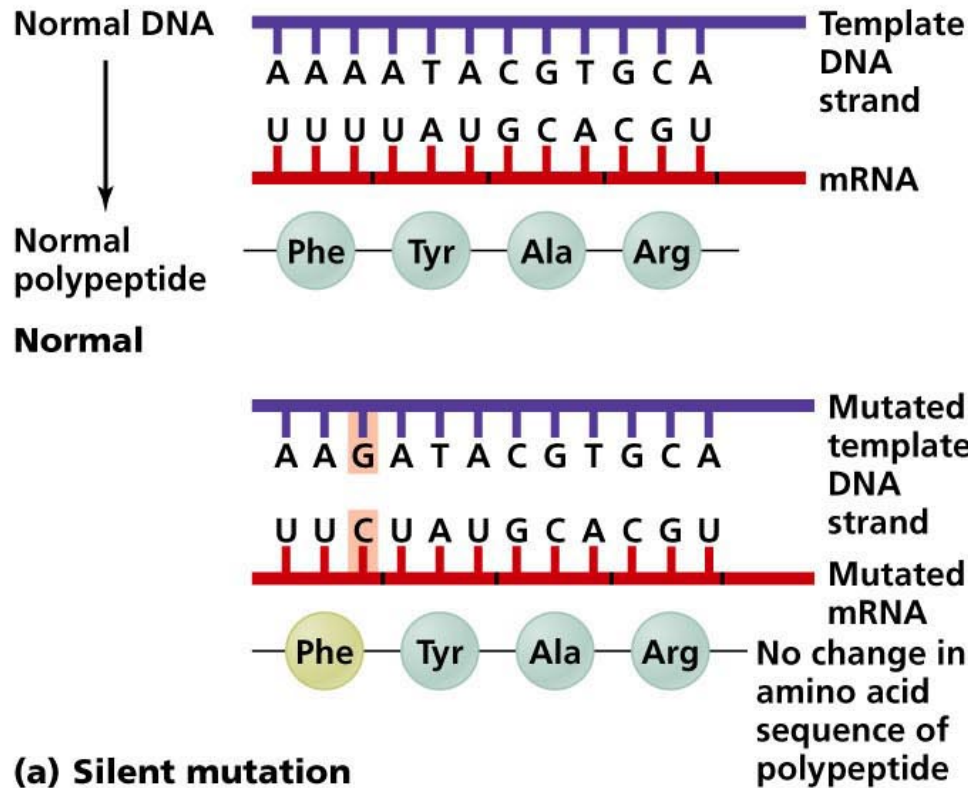
# Mutation is the substrate of evolution

- All (genetic) polymorphisms originate with mutation
- Point mutation (one base for another)

# Point Mutations



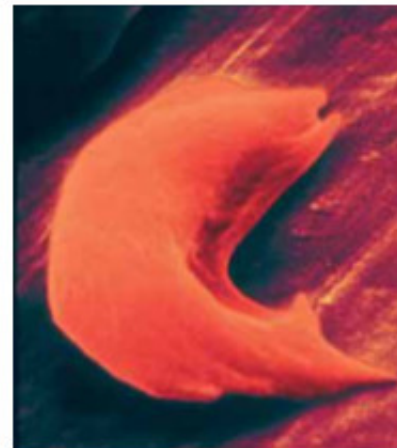
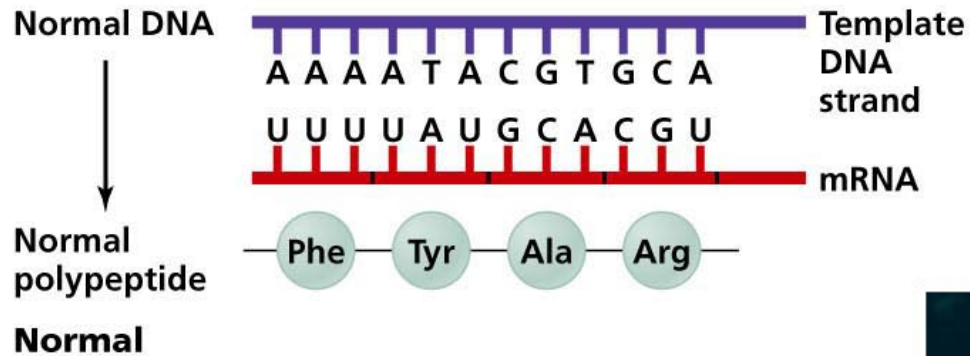
# Mutation is the substrate of evolution



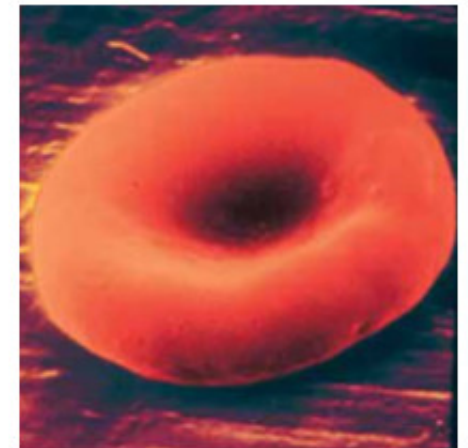
**Fig. 4.** Drug transport function of wild-type and two MDR1 haplotypes. The drug efflux of vaccinia infected/transfected HeLa cells was determined by FACS analysis (14). Cells were transfected with pTM1 (control; purple), MDR1, (wild-type P-gp; green), C1236T-G2677T-C3435T (red), and C1236T-G2677T-C3435A (brown). (A) 0.5  $\mu$ M bodipy-FL-verapamil in the presence of 500  $\mu$ M digoxin; (B) 0.5  $\mu$ M Rh123 in the presence of 150  $\mu$ M digoxin.



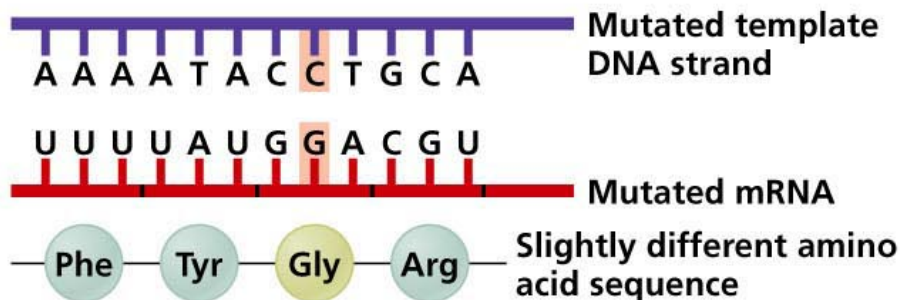
# Mutation is the substrate of evolution



Sickle-cell phenotype

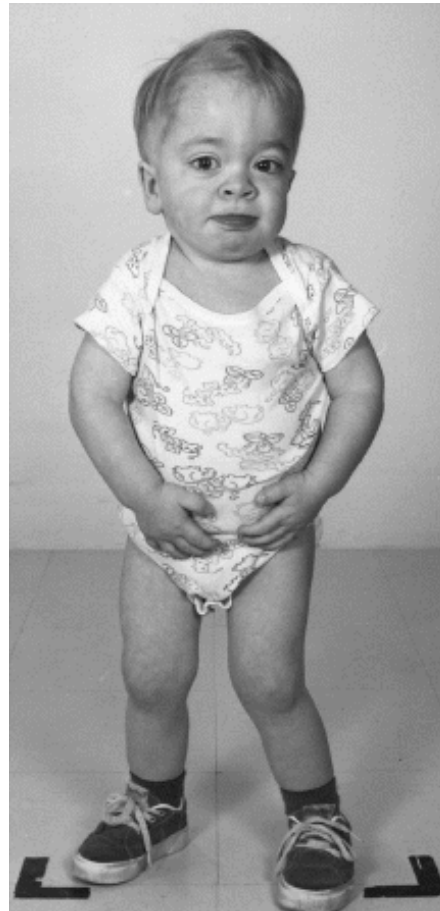
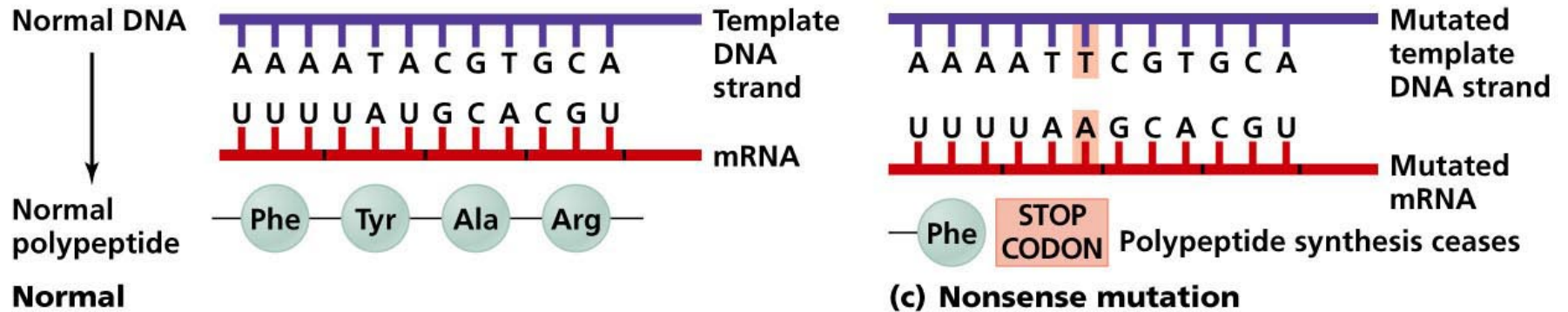


Normal phenotype



(b) Missense mutation

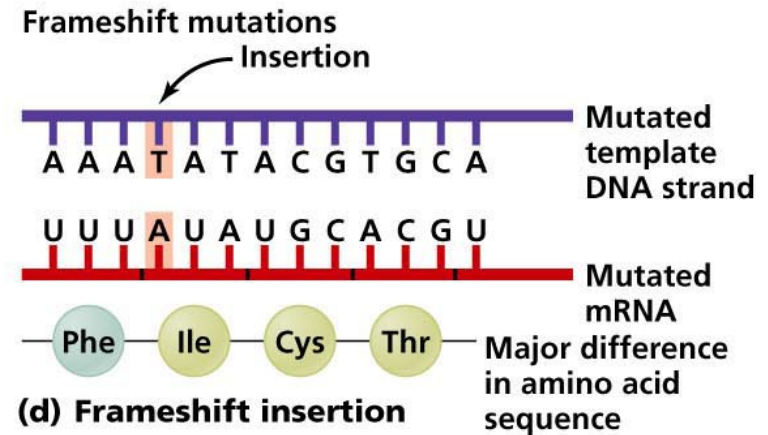
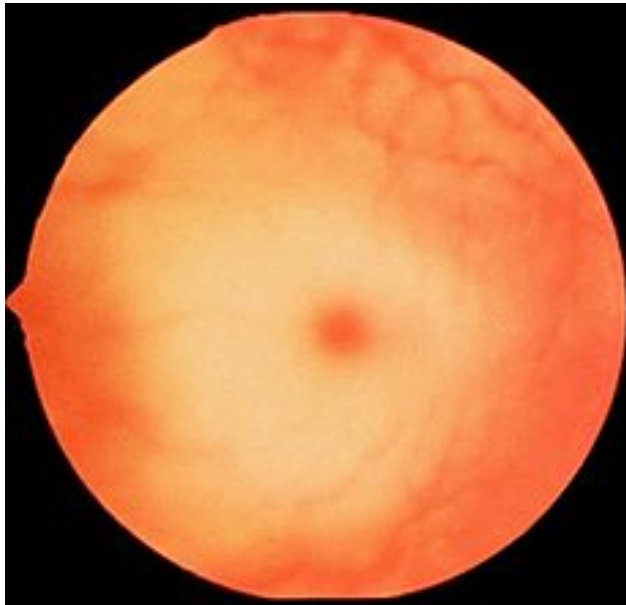
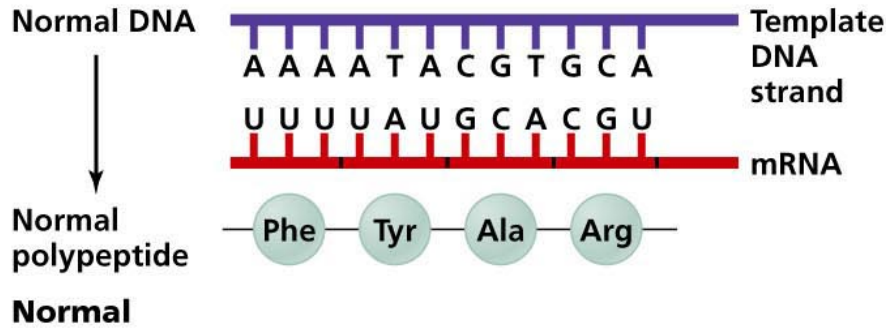
# Mutation is the substrate of evolution



# Mutation is the substrate of evolution

- All polymorphisms originate with mutation
- Point mutation (one base for another)
- Insertion (addition of DNA)

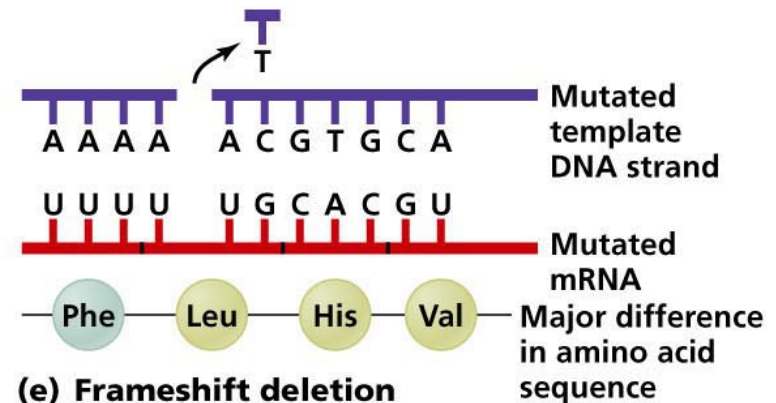
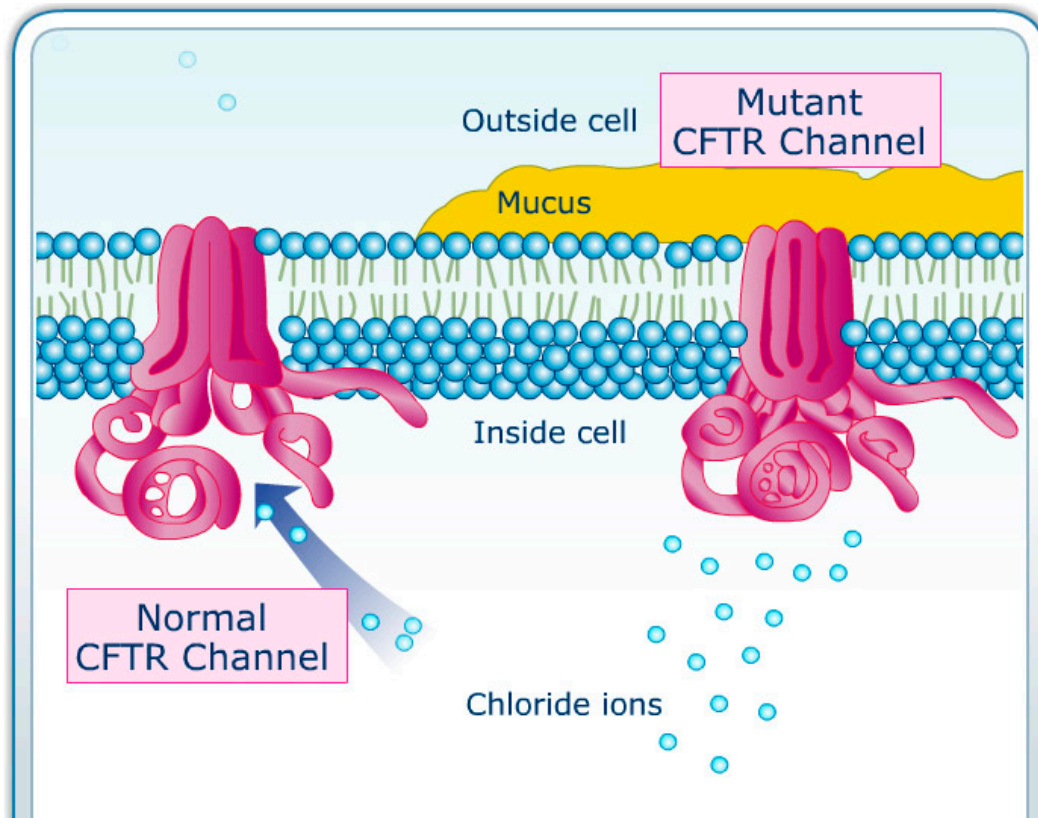
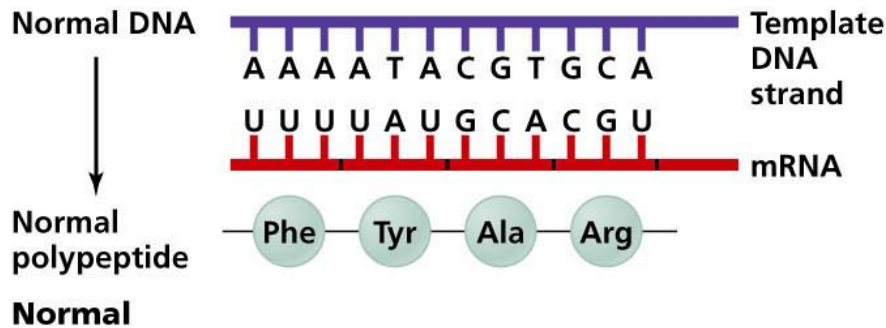
# Mutation is the substrate of evolution



# Mutation is the substrate of evolution

- All polymorphisms originate with mutation
- Point mutation (one base for another)
- Insertion (addition of DNA)
- Deletion (loss of DNA)

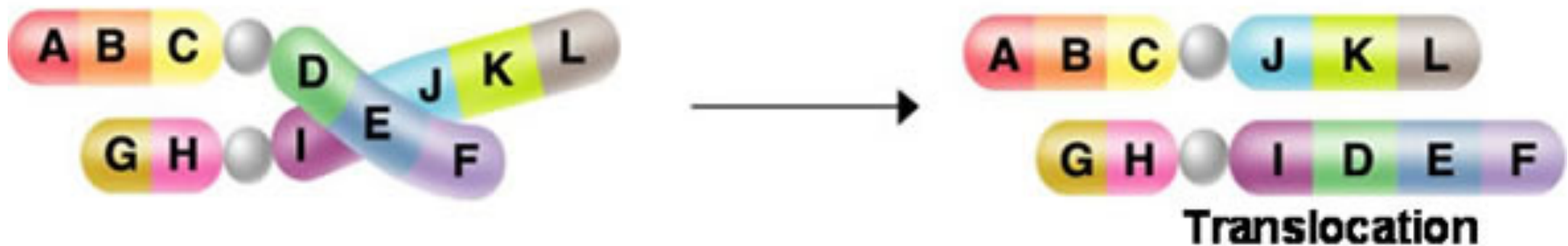
# Mutation is the substrate of evolution



# Mutation is the substrate of evolution

- All polymorphisms originate with mutation
- Point mutation (one base for another)
- Insertion (addition of DNA)
- Deletion (loss of DNA)
- Chromosomal mutations

# Mutation is the substrate of evolution

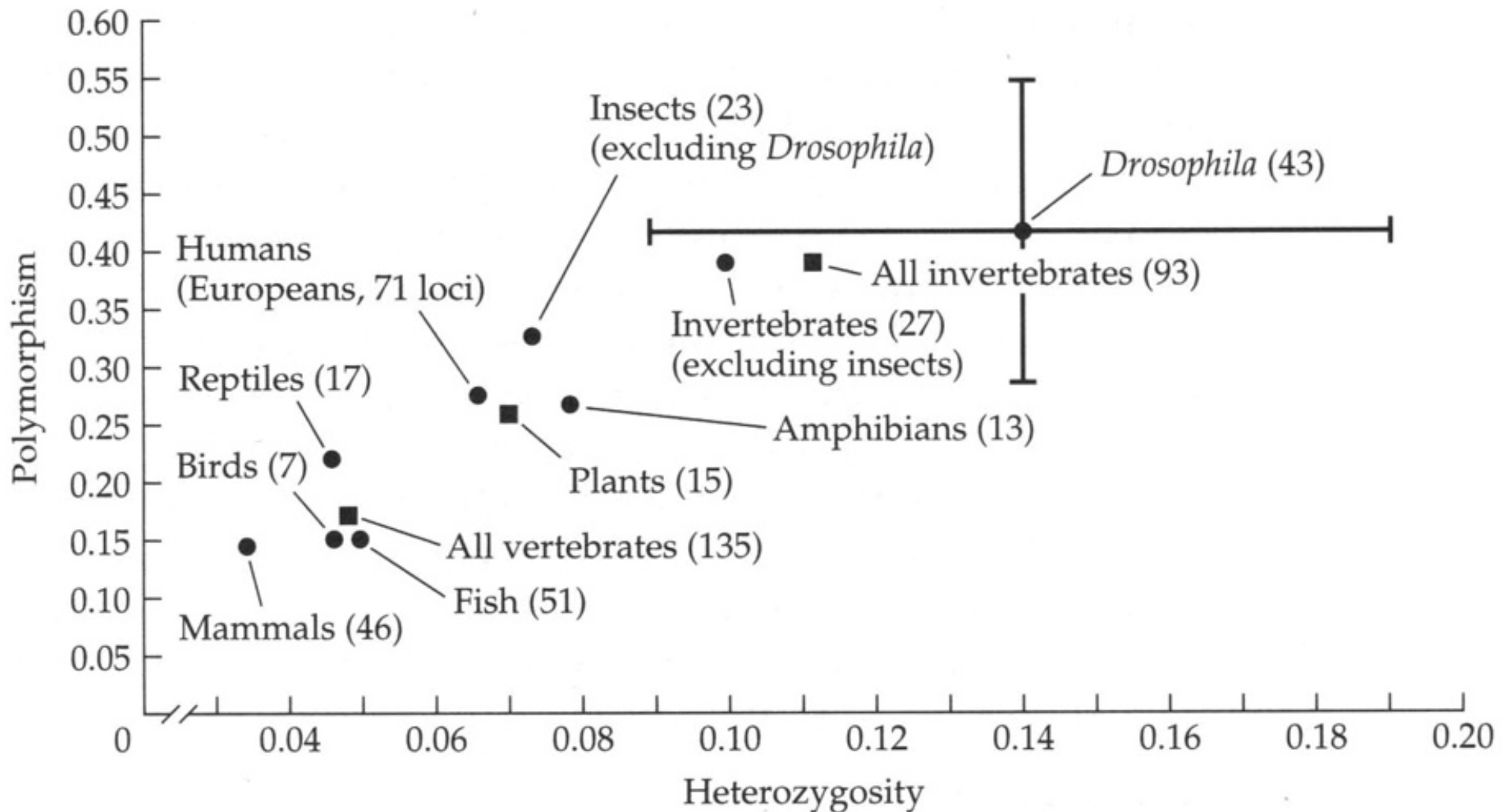




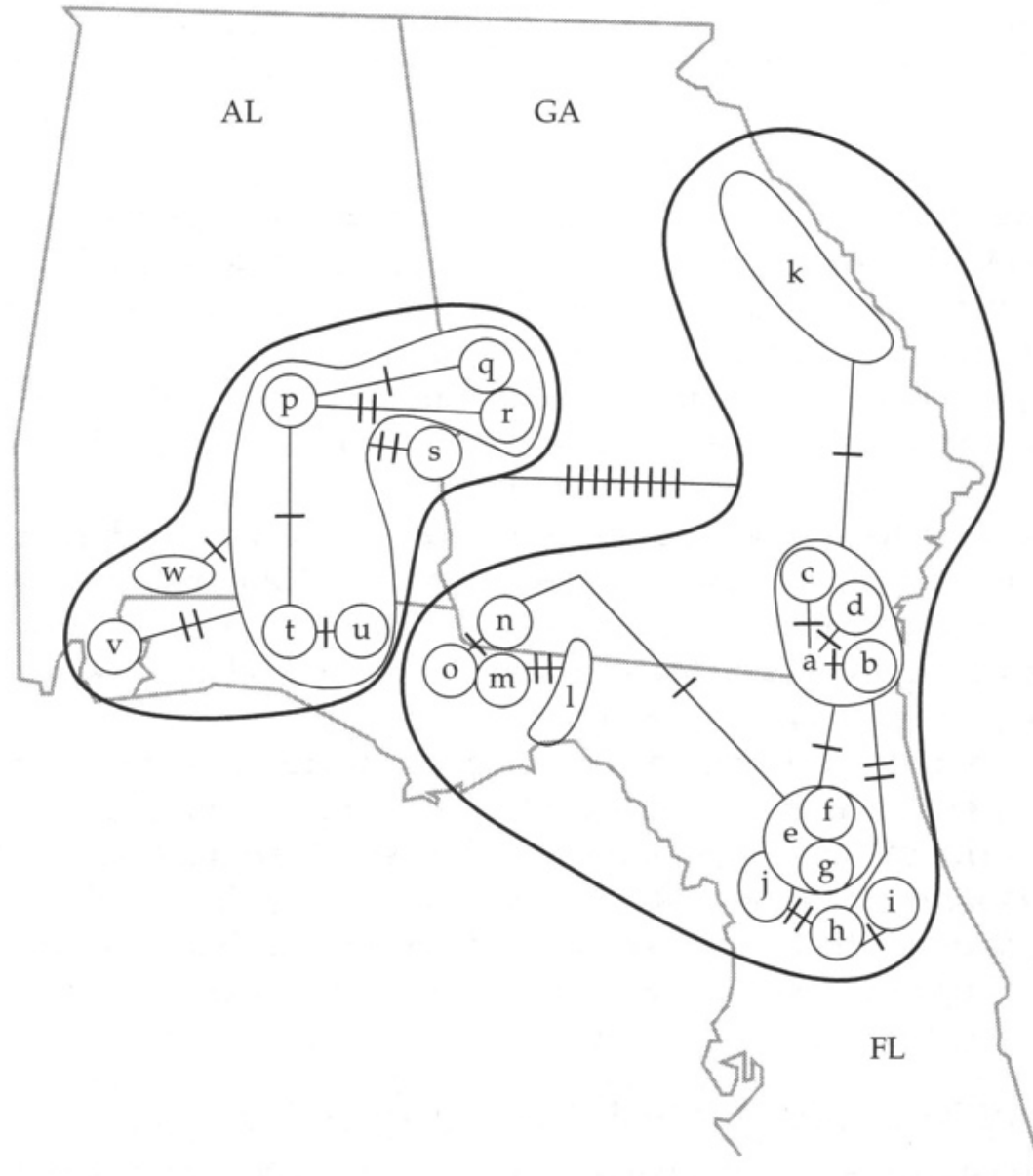
# Genetic Variation

- All genetic variation originates with mutation
  - Mutation is the substrate of evolution
- All levels of organization from single base pairs to entire genomes
- Understanding genetic variation has deep implications

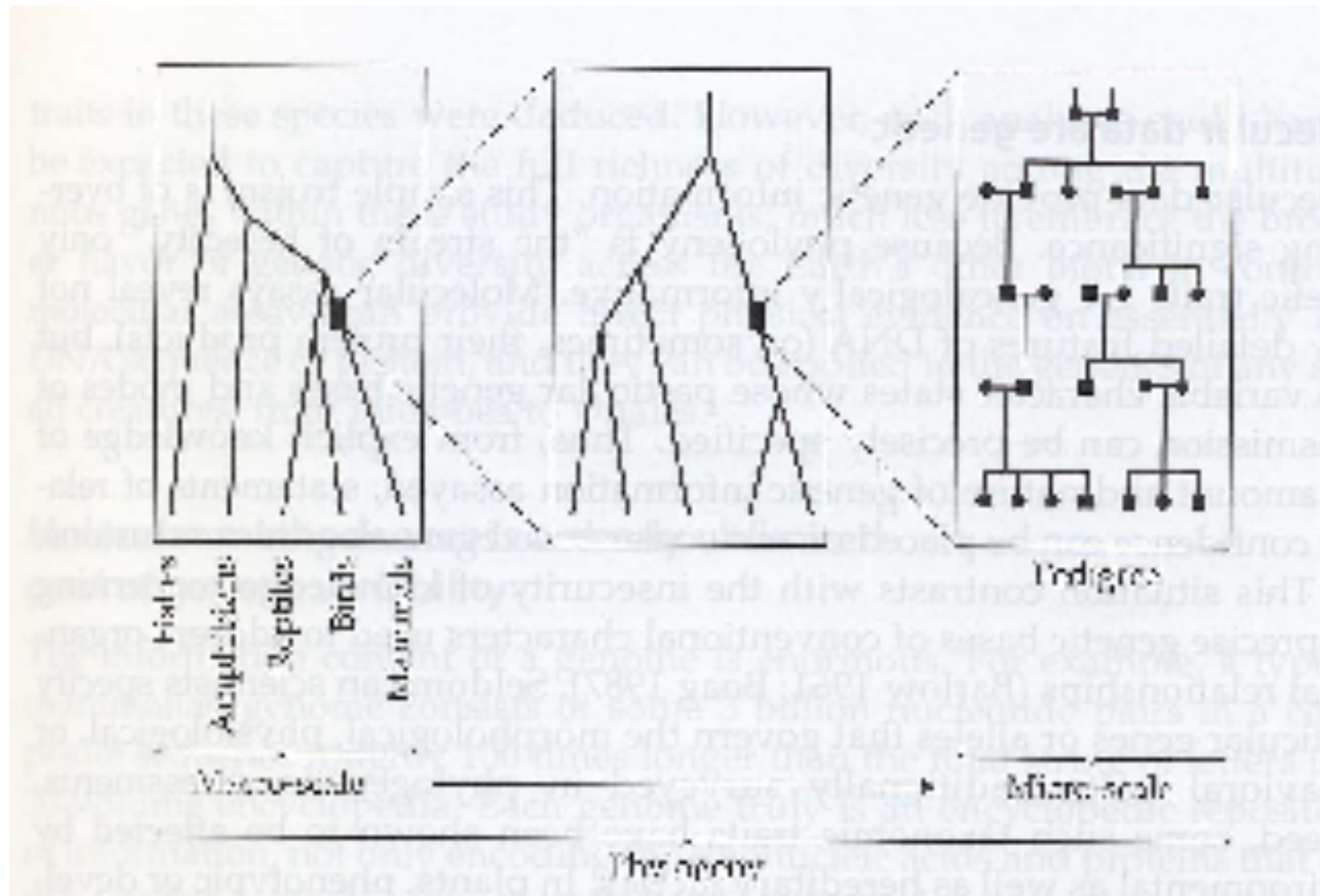
# Genetic variation varies greatly among organisms

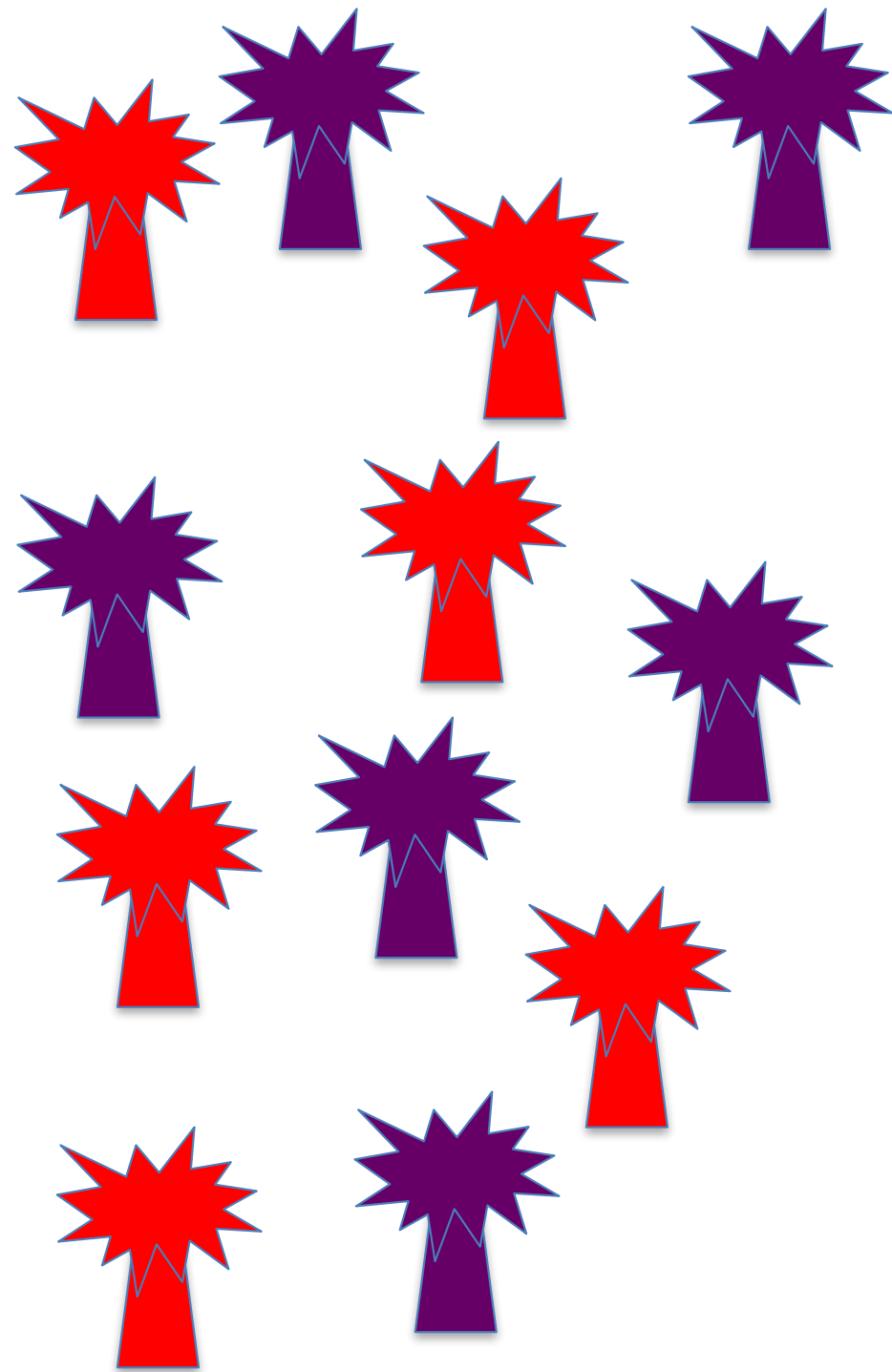
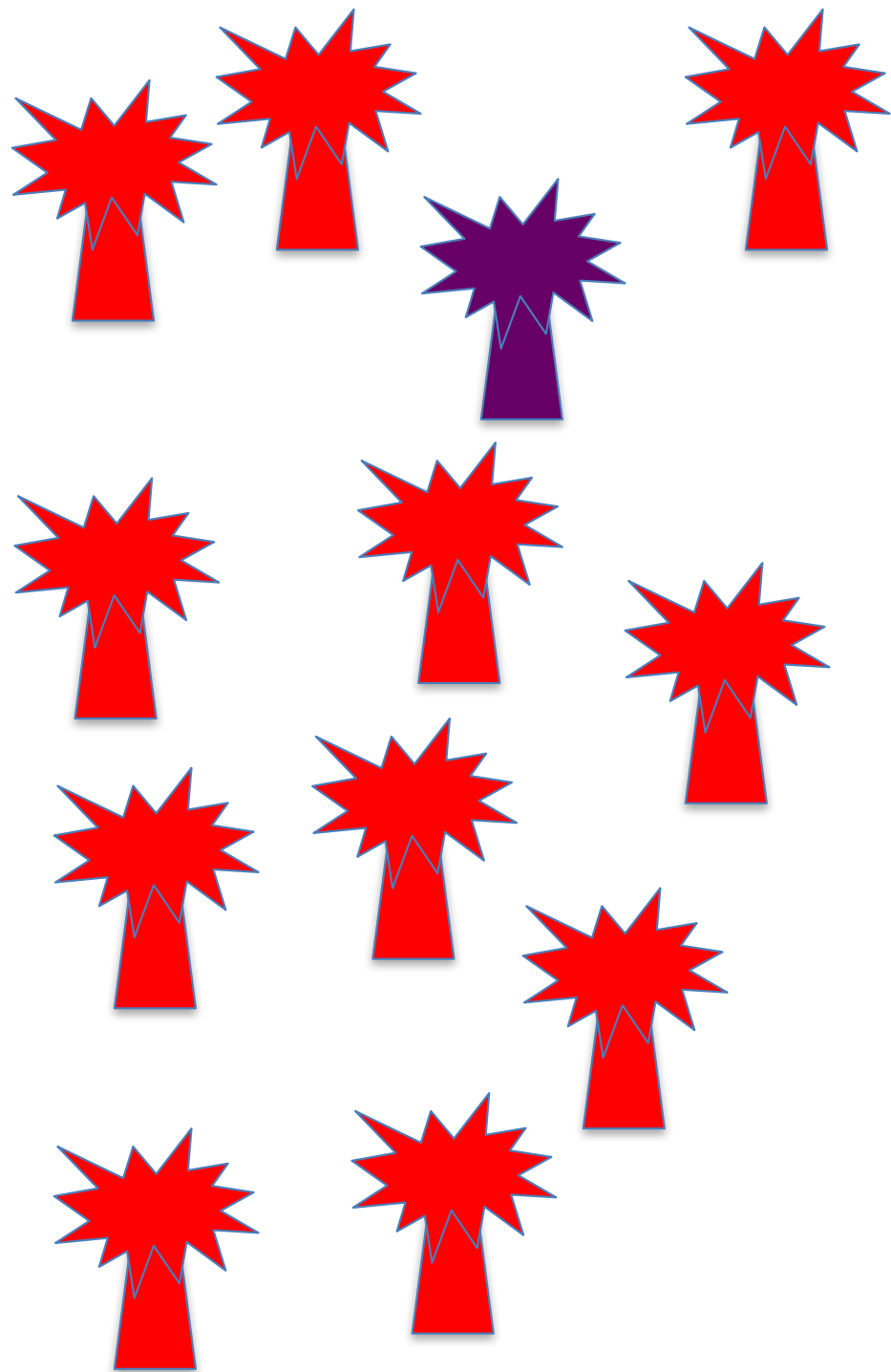


# Genetic variation is distributed geographically



# Genetic variation is nested phylogenetically





# Summarizing molecular data

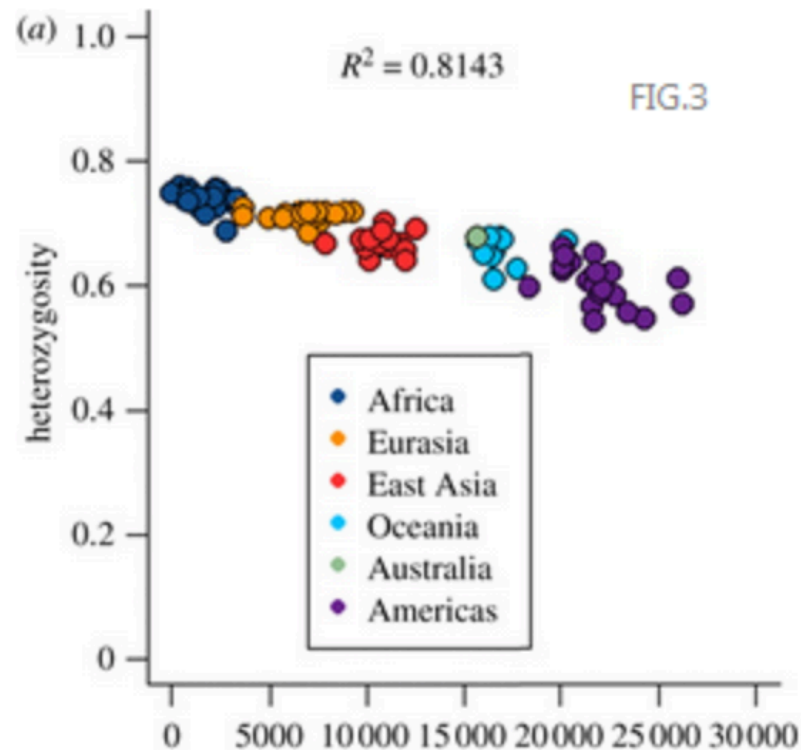
```
#1 CTGCCACCTTTTGTGGGTCTTAGTCCGCAGTGCACTTGTGCCGCCGAGGGGAATGTGGTGCGTTTCCATTGTCCGGATG
#2 .....C.....T.....C.....
#3 .....
#4 .A.....C.....T.....C.....
#5 .....G.....A.....
#6 .....
#7 .....C.....C.....
#8 .....T.....A.....
#9 .....A.....
#10 .....A.....
#11 .....C.....C.....C.....
#12 .....C.....C.....C.....
```

# Key terms

- Gene
- Locus
- Allele
- Genotype
- Haplotype
- Homozygote
- Heterozygote
- Polymorphism

# Heterozygosity

- Expected probability that an individual will be heterozygous at a locus
- Expected proportion of heterozygotes in a population at a locus



Microsatellite heterozygosity vs. geographical distance



# Allelic diversity

- Number of alleles per locus

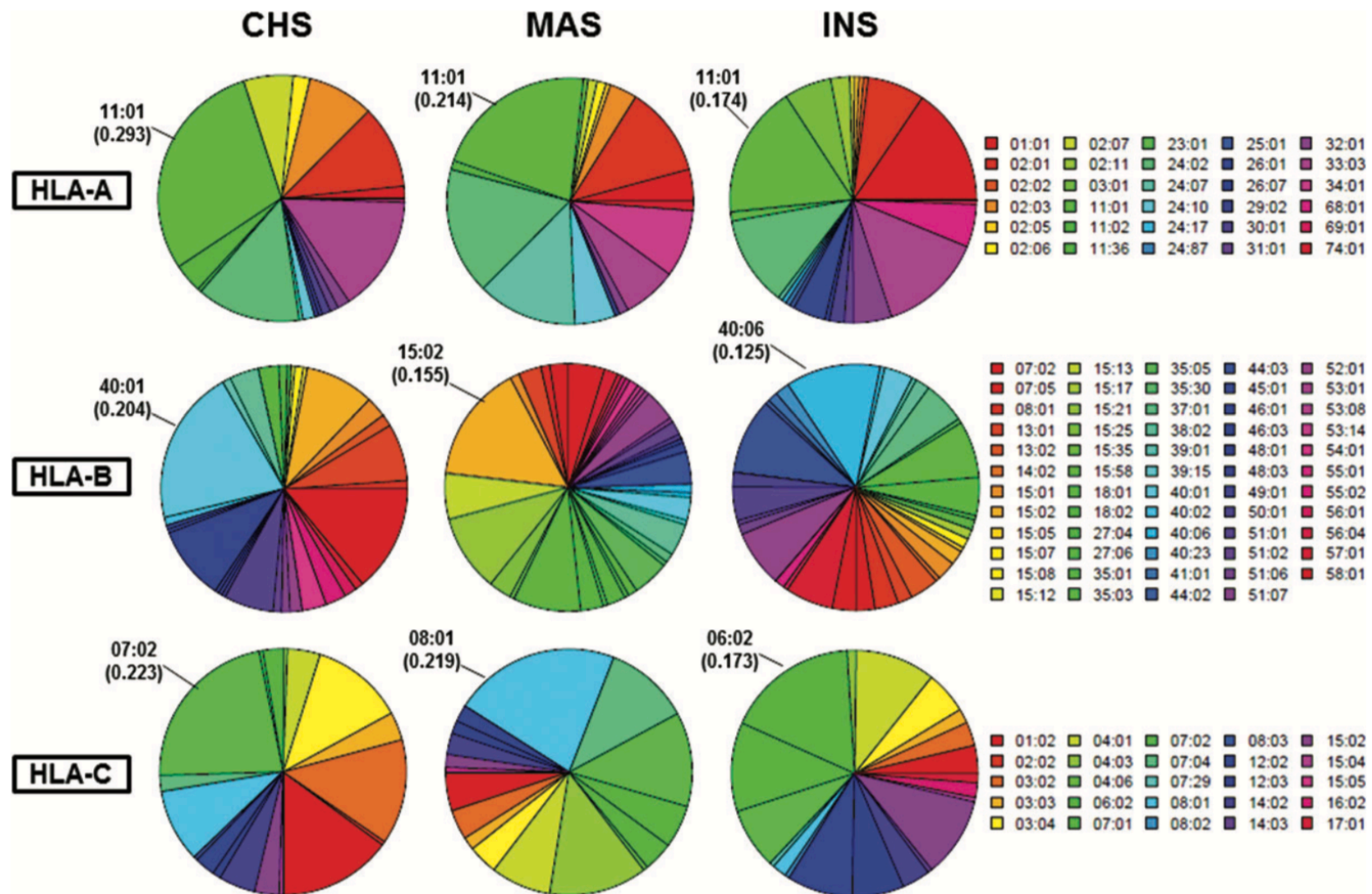
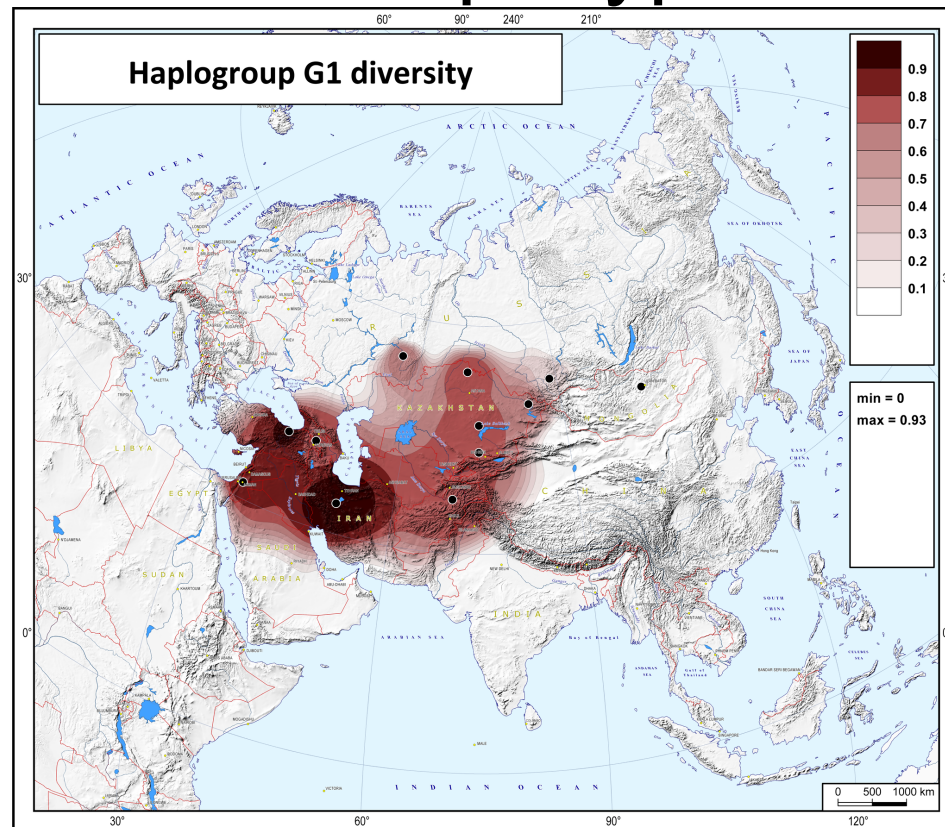


Figure 1. Allelic diversity and distribution in HLA Class I genes. Pie charts illustrating the allelic diversity of the three genes in HLA Class I in the three populations.

# Haplotype indices

- Haplotype number
- Unique ('private') haplotypes
- Haplotype diversity: probability that two randomly chosen haplotypes are different



# Nucleotide-level Indices

- $\Theta_w$

- Watterson's estimator

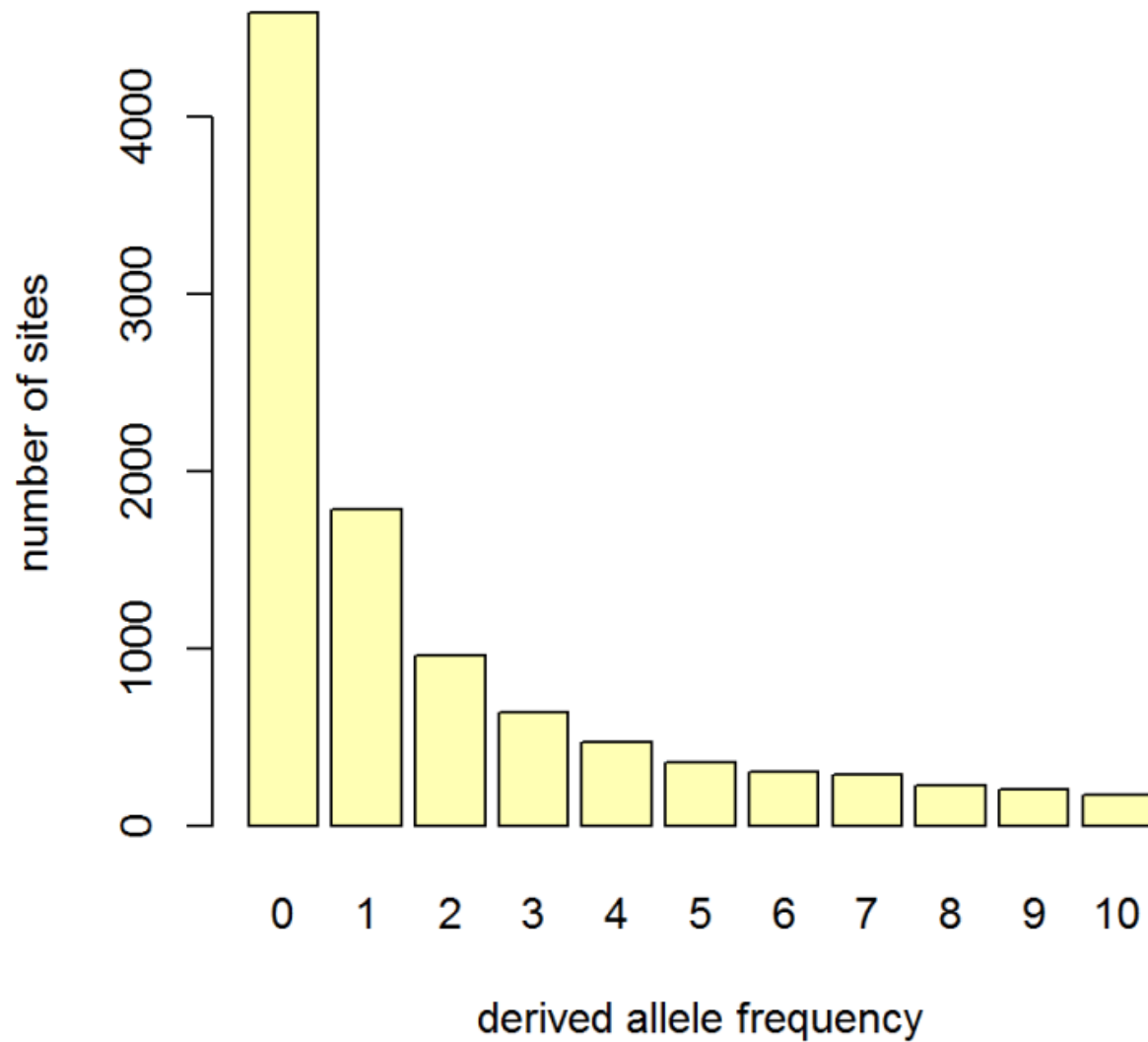
- $$= \frac{S}{\sum_{i=1}^{n-1} \frac{1}{i}}$$

- $\pi$

- Average pairwise difference between alleles

# Site Frequency Spectrum

5 diploid individuals  
10,000



# Genetic Variation: Software

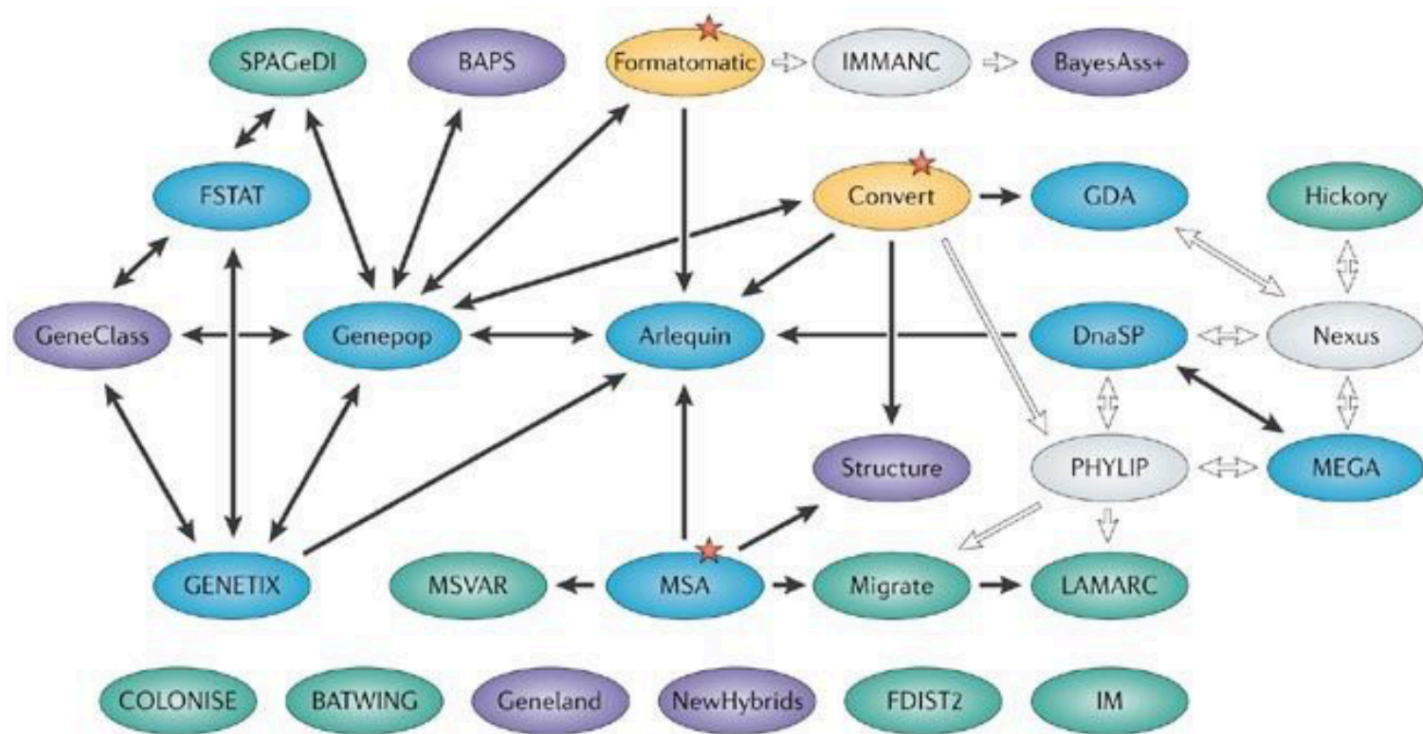
- DnaSP (DNA sequence polymorphism)
- DNA sequence
- Basic population genetic analyses
- Estimate genetic diversity
- Within/Between populations
- LD, recombination, gene flow, gene conversion
- Neutrality tests

# Genetic Variation: Software

- Arlequin
- MEGA
- PHYLIP
- ...
- Excoffier and Heckel 2006: Computer programs for population genetic data analysis: a survival guide

# Genetic Variation: Software

From: Computer programs for population genetics data analysis: a survival guide



# Why do we care about genetic variation?

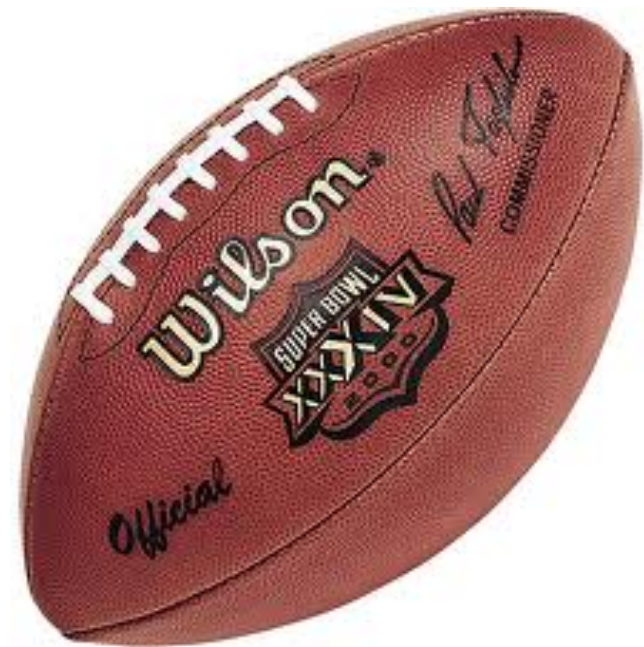
- Genetic variation underlies phenotypic differences among individuals
  - Including disease risk and responses to drugs and environmental factors





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  - Including disease risk and responses to drugs and environmental factors
- Individual identification



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  - Including disease risk and responses to drugs and environmental factors
- Individual identification
- Manage resources



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- Genetic variation underlies phenotypic differences among individuals
  - Including disease risk and responses to drugs and environmental factors
- Individual identification
- Manage resources
- Public Health

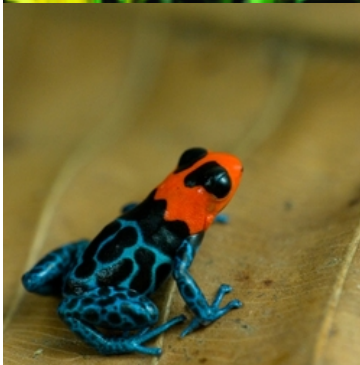


# Why do we care about genetic variation?

- Genetic variation underlies phenotypic differences among individuals
  - Including disease risk and responses to drugs and environmental factors
- Individual identification
- Manage resources
- Public Health
- Improve plant and animal food products
- Understand genetic basis of disease and other complex phenotypes including behaviours
- Insights into evolutionary history, ancestry

# Conservation genetics

- Use of genetic theories and techniques to study the loss and restoration of genetic diversity
- Derived from ecological, evolutionary, and quantitative genetics
- Focuses on small populations



# Human threats

- Poaching



# Human threats

- Poaching
- Habitat loss, degradation, fragmentation

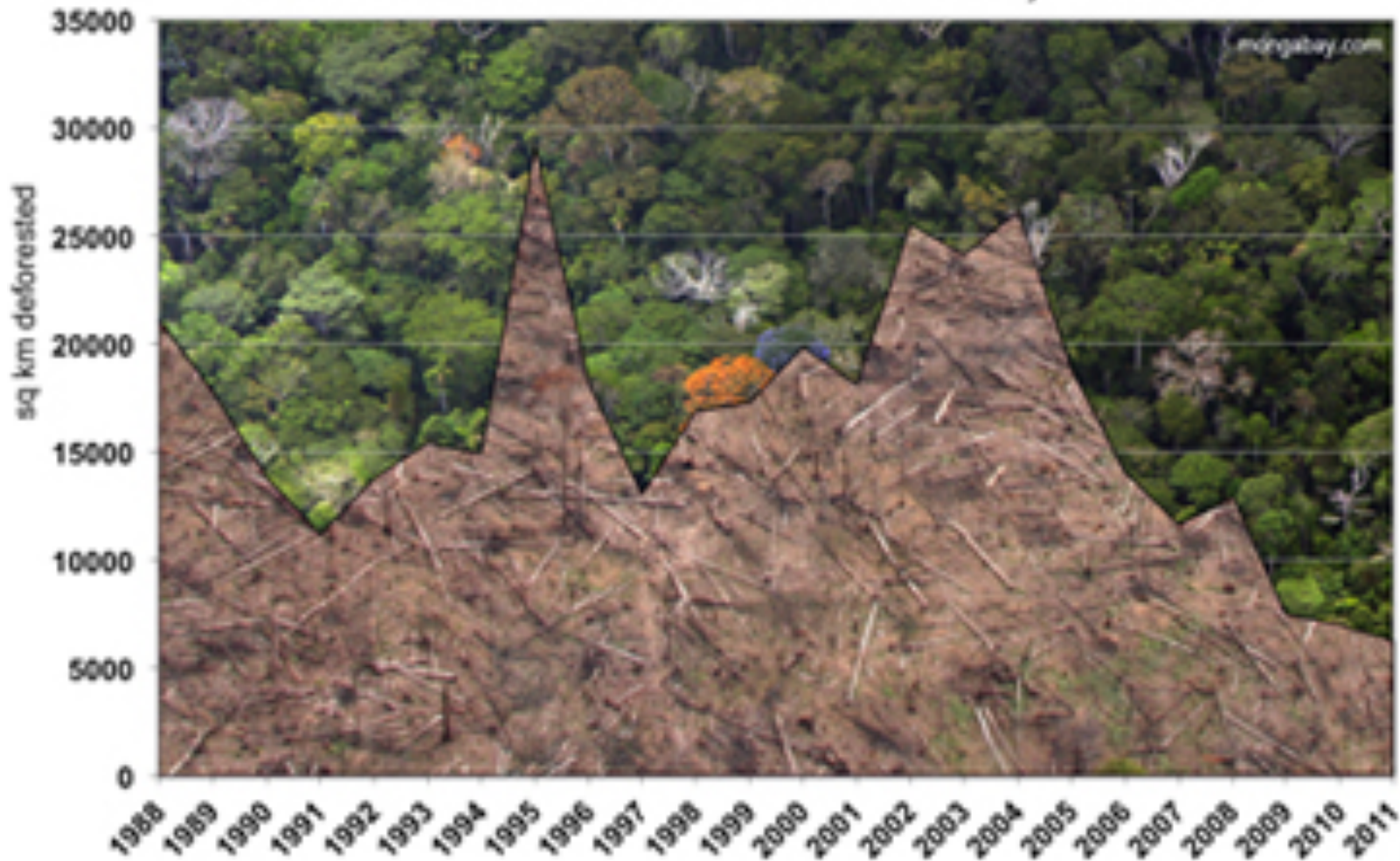




# Human threats

- Poaching
- Habitat loss, degradation, fragmentation

Deforestation in the Brazilian Amazon, 1988-2011



# Human threats

- Poaching
- Habitat loss, degradation, fragmentation



# Human threats

- Poaching
- Habitat loss, degradation, fragmentation
- Invasion



## Human threats

- Poaching
- Habitat loss, degradation, fragmentation
- Invasion
- Pollution



## Human threats

- Poaching
- Habitat loss, degradation, fragmentation
- Invasion
- Pollution
- Climate change



# Why is conservation important?

- Sixth mass extinction
- Aesthetic value
- Bioresources
- Ecosystem services
- Ethical, moral considerations

# How can genetics help?

- Minimize inbreeding and loss of genetic variation
- Identify populations of concern
- Resolve population structure
- Resolve taxonomic uncertainty
- Define management units
- Detect hybridization
- Detect and define invasive species
- Estimate population size and sex ratio
- Establish parentage
- Understand population connectivity
- Aid in management
- Enhance reproductive capacity of organisms
- How do genetic factors affect extinction risk?
- How can we best genetically rescue populations?
- What can we use genetics to identify hotspots?
- Applied genetic detective work
- ...

# Conservation genetics

- Management strategies
  - Captive Breeding design



# Captive breeding

- Establish populations in secure *ex situ* locations
- Educate/engage public
  - Fundraising
- Provide educational opportunities
- Provide animals for reintroduction
- Maintain demographically stable, genetically variable populations with little inbreeding
- 1150 zoos and aquaria worldwide house 1.2 million animals
  - 5-10% used for endangered species

# Captive breeding

**Table 1.** Types of contributions that molecular markers can make to captive breeding programs

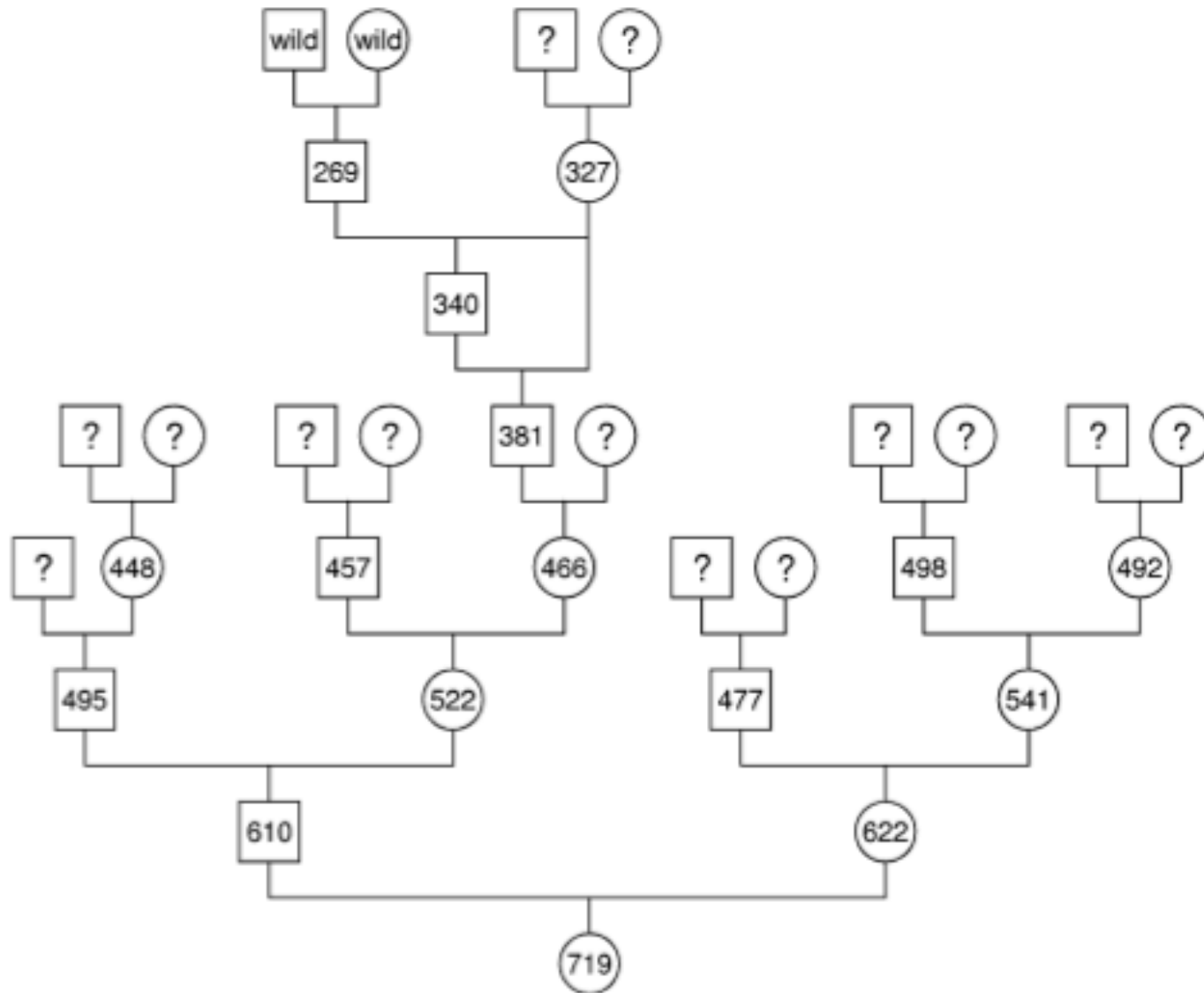
Contribution to breeding program	Species	Citation
Assessment of founder relationships	California condor ( <i>Gymnogyps californianus</i> )	Geyer et al. (1993)
	Guam rail ( <i>Rallus owstoni</i> )	Haig et al. (1994)
	Micronesian kingfishers ( <i>Halcyon cinnamomina</i> )	Haig et al. (1995)
Pedigree reconstruction	Bearded vulture ( <i>Gypaetus barbatus</i> )	Gautschi et al. (2003)
	Lion-tailed macaques ( <i>Macaca silenus</i> )	Morin and Ryder (1991)
Subspecies identification	Przewalski's horse ( <i>Equus ferus przewalskii</i> )	Bowling et al. (2003)
	Lion ( <i>Panthera leo</i> )	O'Brien et al. (1987)
Identification of geographic origin	Chimpanzee ( <i>Pan troglodytes</i> )	Ely et al. (2005)
	Galapagos tortoise ( <i>Geochelone nigra</i> )	Russello et al. (2007)
Quantification of wild genetic diversity captured	Baird's tapir ( <i>Tapirus bairdii</i> )	Norton and Ashley (2004)
	Iberian wolf ( <i>Canis lupus signatus</i> )	Ramirez et al. (2006)
Identification of genetically valuable individuals	Whooping crane ( <i>Grus americana</i> )	Jones et al. (2002)
	St Vincent parrot ( <i>Amazona guildingii</i> )	Russello and Amato (2004)
Assessment of hybridization	Lesser white-fronted goose ( <i>Anser erythropus</i> )	Ruokonen et al. (2007)

# Parma wallaby

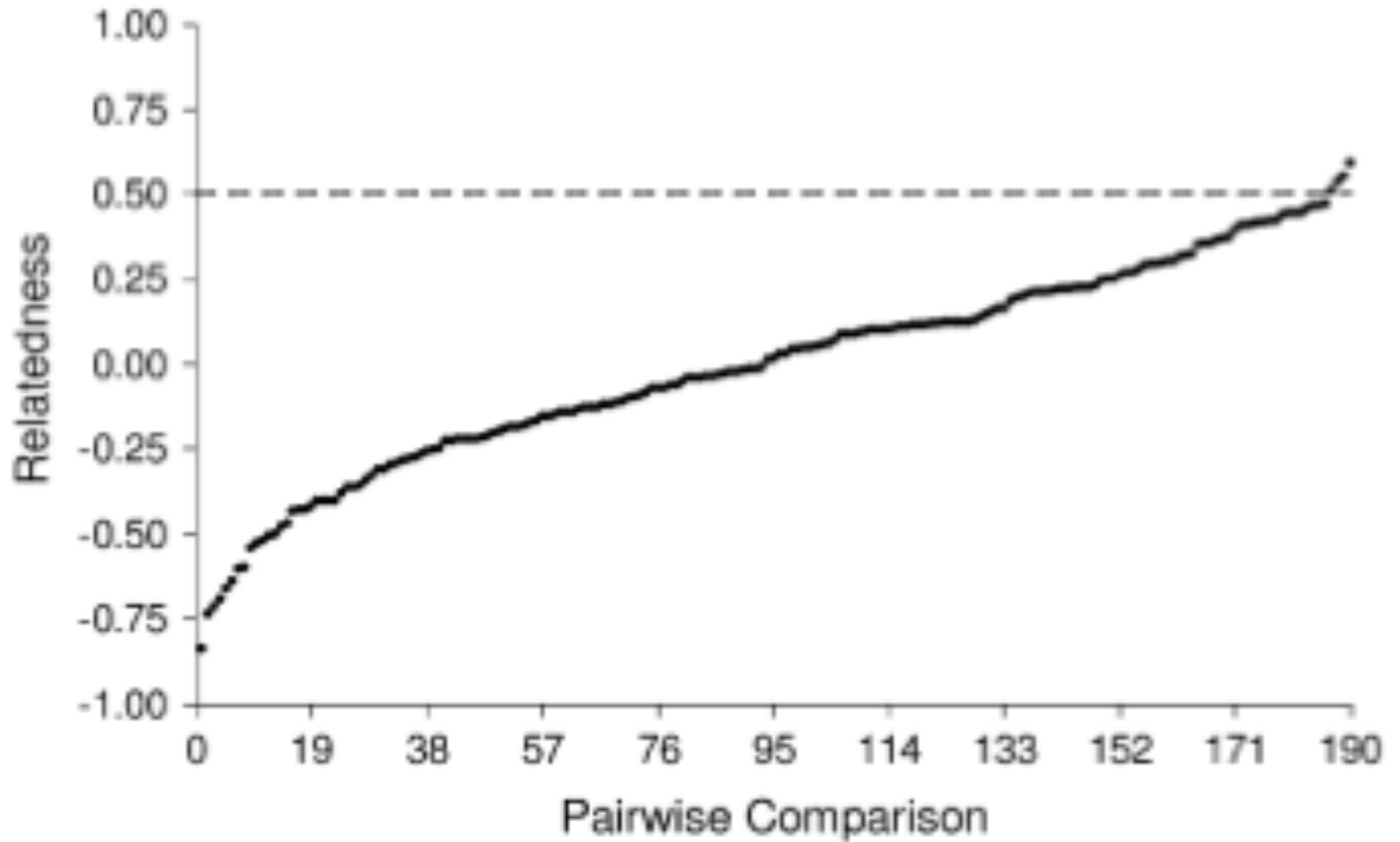
- Marsupial native to Australia
- Threatened by habitat degradation and low census size
- Captive population founded in 1960s
  - Managed by pedigree
  - Breeding recommendations based on kinship
  - 50% of individuals have unknown ancestry



# Parma wallaby



# Parma wallaby



# Hereditary defects

**TABLE 1. Possible, presumed, or proven hereditary defects observed in wild animals bred in captivity**

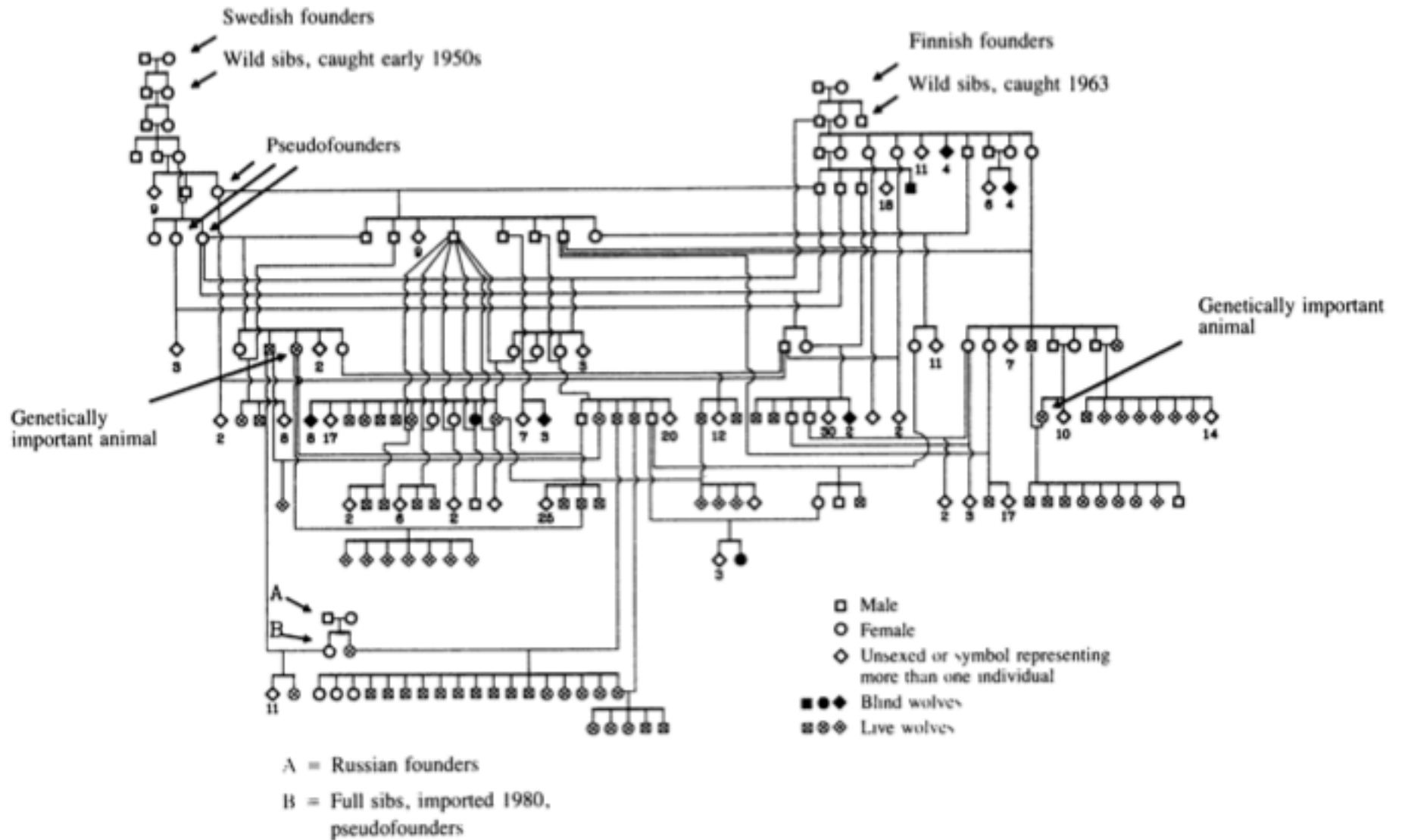
Species/taxonomic group	Defect	Heredity/mode of inheritance	Source
Arctic fox <i>Alopex lagopus</i>	Genu flexum (abnormal hind leg)	Hereditary	Krukowski & Klos, 1982
	Gingival hyperplasia	Hereditary/probably autosomal recessive with incomplete penetrance	Dyrendahl & Henricson, 1959
Asian small-clawed otter ( <i>Aonyx cinerea</i> )	Urolithiasis	Possibly hereditary	Karesh, 1983; Calle, 1985
Axis deer <i>Axis axis</i>	Ventricular septal defect	Possibly hereditary	Gendron, 1975
Bison Wood bison <i>Bison bison athabasca</i> Plains bison <i>Bison bison bison</i>	Arthrogryposis and degenerative joint disease	Possibly hereditary	Schuh & Haigh, 1990
Black-and-white ruffed lemur ( <i>Varecia variegata variegata</i> )	Pectus excavatum Diaphragm hernia Cleft palate Scoliosis Exencephaly Hydrocephaly	Autosomal recessive Presumed hereditary	Benirschke et al., 1985
Brown bear <i>Ursus arctos</i>	Albinism	Single autosomal recessive	Laikre et al., 1996; Andrén et al., 1997; this paper
Carnivores	Various dental diseases	Possibly hereditary	Fagan, 1980
Dear mouse <i>Peromyscus maniculatus</i>	Spherocytosis Post juvenile nude, hair deficiency	Autosomal recessive Autosomal recessive	Huestis et al., 1956 Clark, 1934
Dingo <i>Canis dingo</i>	Hip joint dysplasia	Presumed hereditary	Christoph et al., 1969
Douc langur <i>Pygathrix nemaus</i>	Pectus excavatum	Presumed hereditary	Sedgwick, 1981

# Grey wolf

- Largest member of Canidae
- Ancestor of domestic dog
- Once abundant over Eurasia, North Africa, North America
- Current range restricted
- Threatened by habitat loss

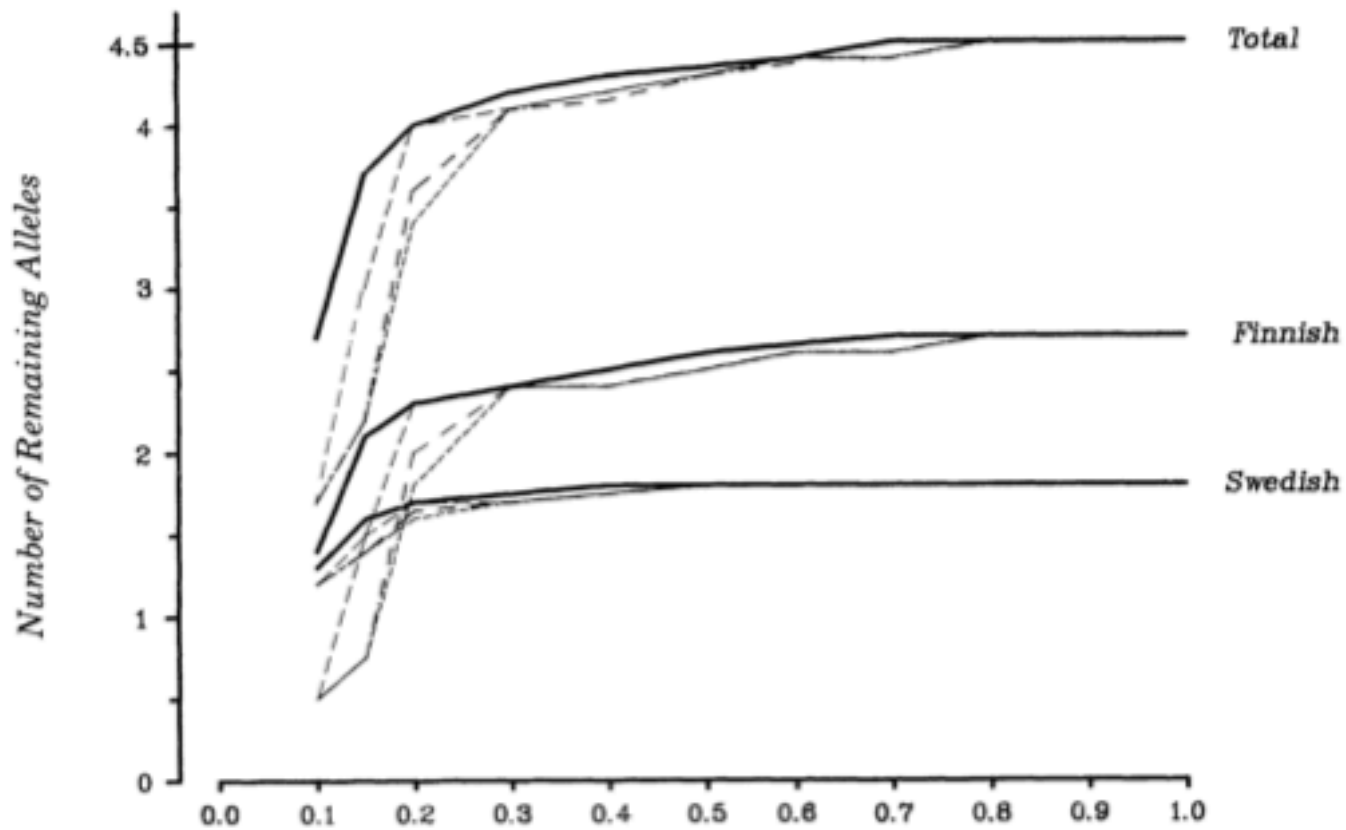


# Grey wolf





# Grey wolf



<i>Minimum Carrier Probability for Removal</i>	<i>Number of Animals Remaining</i>	$F_T$	$F_F$	$F_S$
1.0 (no removals)	55	0.34	0.17	0.17
0.6	49	0.35	0.17	0.18
0.5	43	0.36	0.18	0.18
0.4	40	0.36	0.18	0.18
0.3	36	0.37	0.19	0.18
0.2	29	0.38	0.22	0.16
0.1	2	0.34	0.16	0.18

# Black footed ferrets



# Black footed ferrets



	<i>n</i>	$H_O (\pm SD)$	$H_{dem}$	A	$N_c$	$N_{e-dem}$	$N_{e-gen}$ (95% $CI^a$ )
Agate-Sage Creek	207	$0.39 \pm 0.12$	0.43	2.0	81	75	10.6 (2.2-18.9)
Heck Table	47	$0.39 \pm 0.16$	0.41	2.0	17	14	2.0 (0.0-3.6)
Combined population, 5 loci <sup>b</sup>	254	$0.44 \pm 0.04$	0.43	2.0			

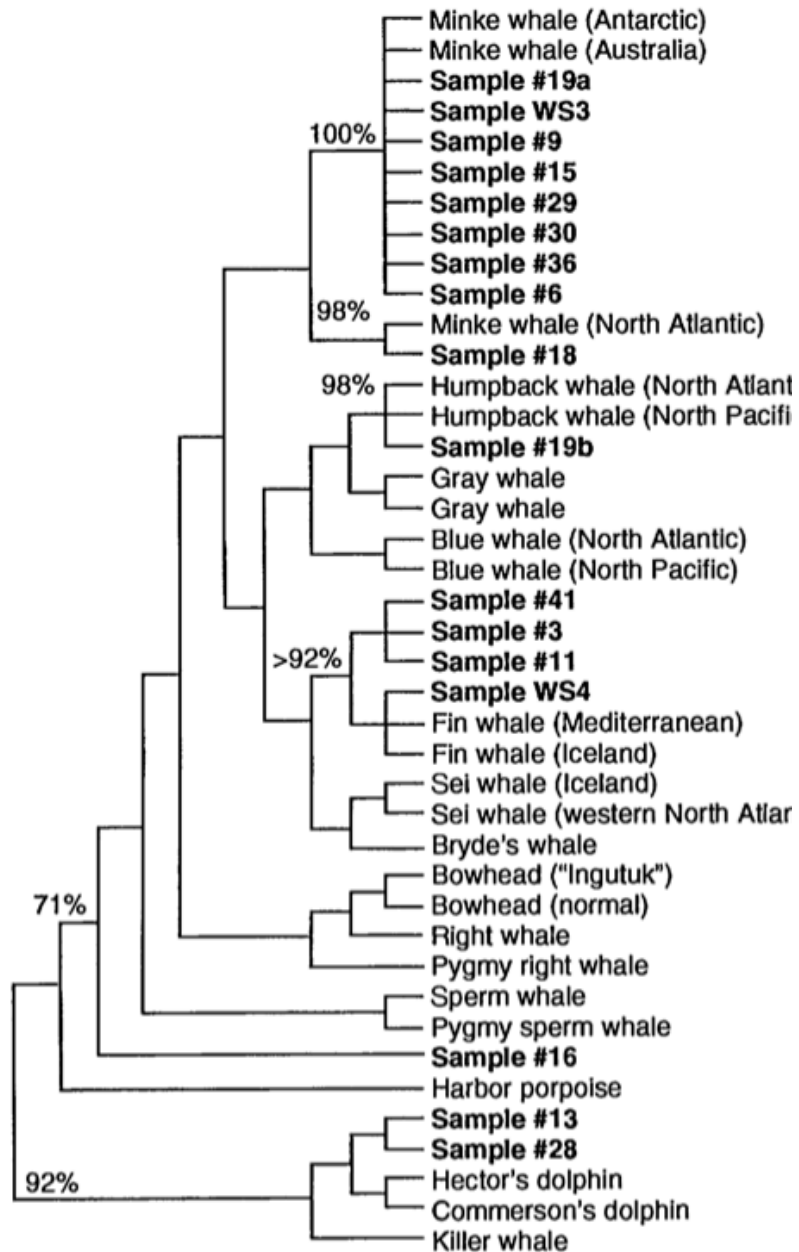
# Conservation genetics

- Management strategies
  - Captive Breeding design
  - Forensics

# Whaling



# Whaling

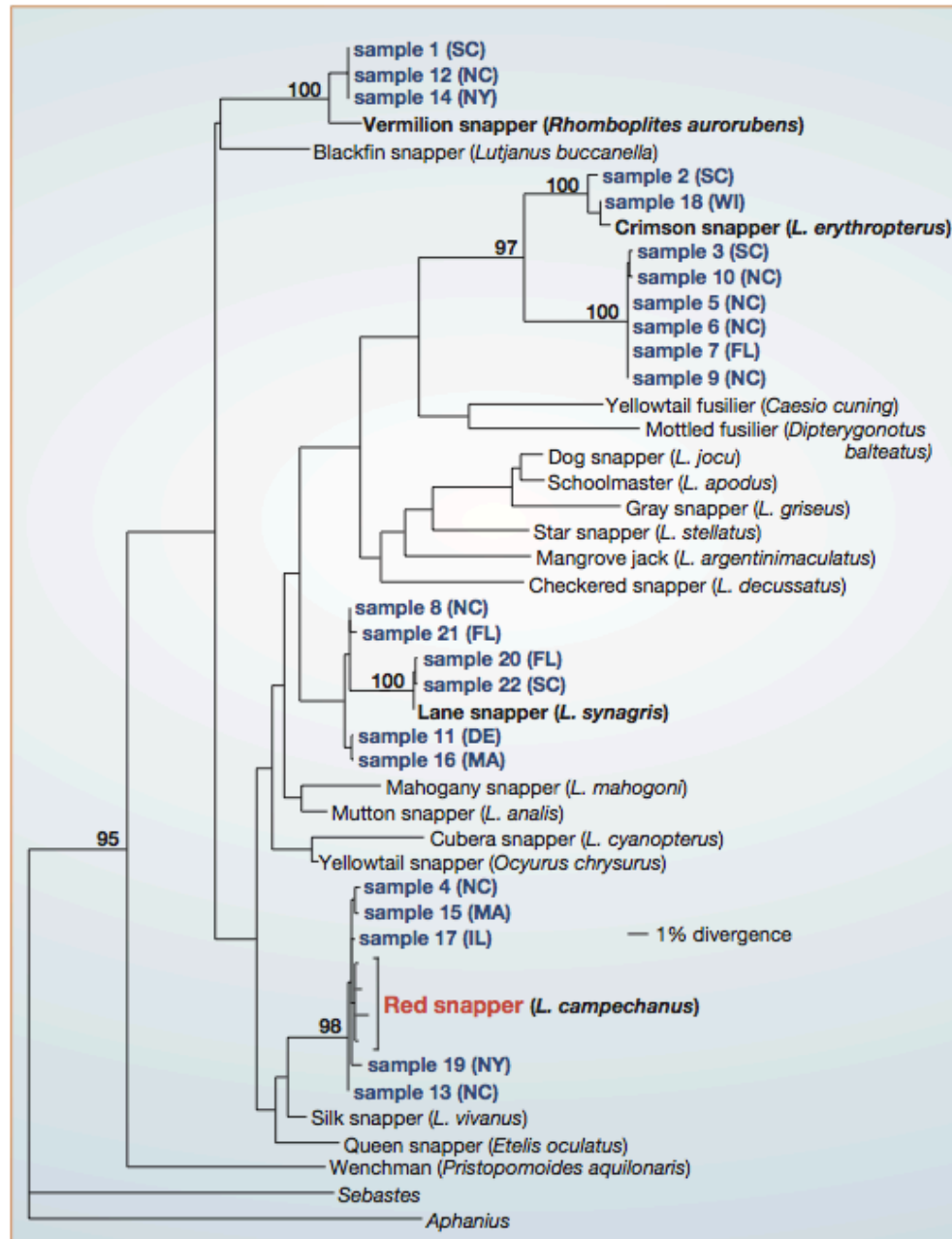


# Red Snapper

- Commercially and recreationally-caught
- Most commonly caught snapper in US
- Restrictions to maintain populations
  - 51% commercial
  - 49% recreational

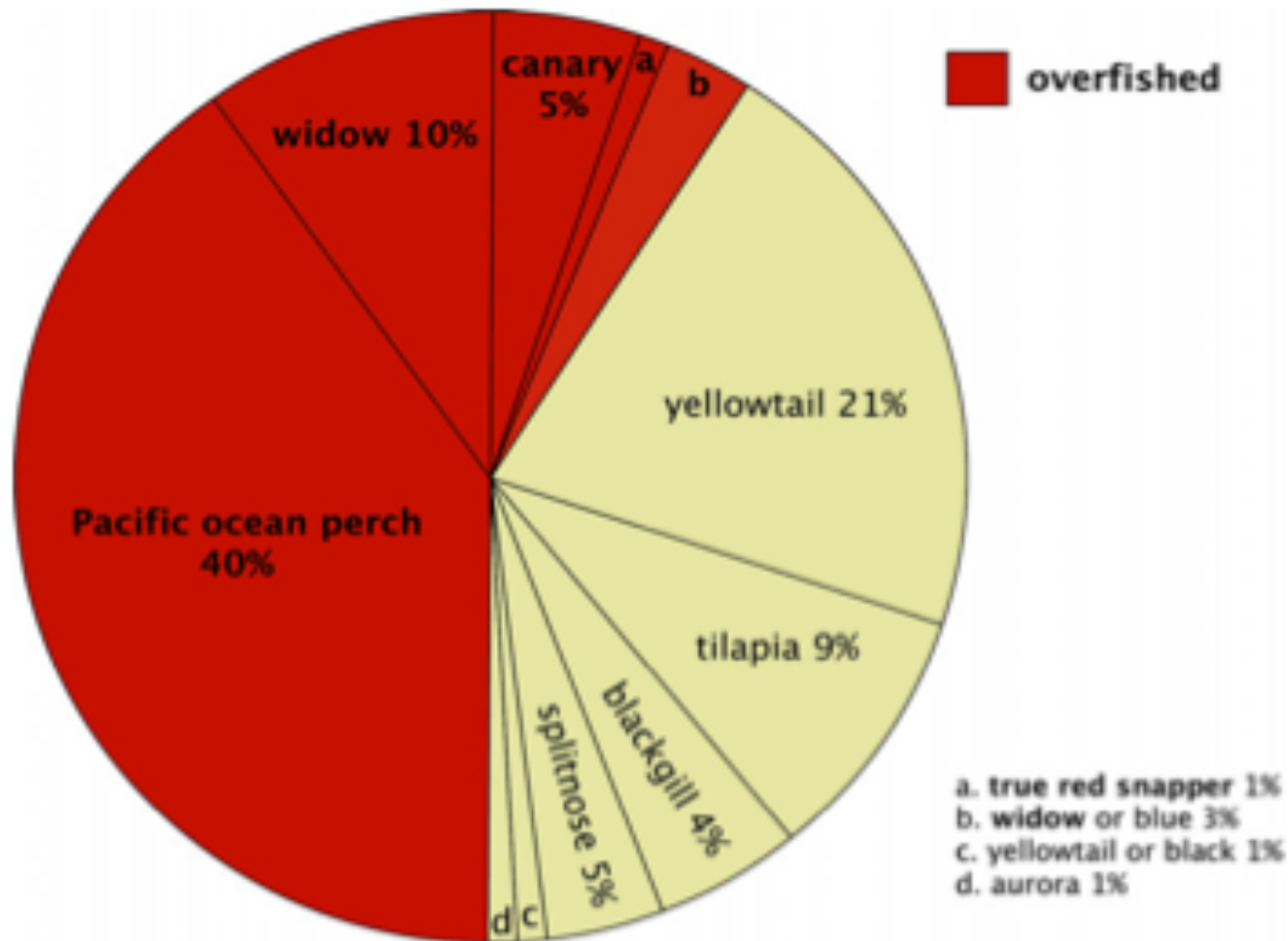


# Red Snapper





# Pacific Red Snapper



**Fig. 2 – Percent of Pacific red snapper samples ( $n = 77$ ) that have been declared overfished within the past three years by the National Marine Fisheries Service.**

# Conservation genetics

- Management strategies
  - Captive Breeding design
  - Forensics

# Future of conservation genetics

- Open questions:
  - Genetic variation and fitness
  - Mechanism connecting genetic variation and fitness
  - Genotype by environment interaction
  - Intermediate phenotypes
- New technologies
  - DNA sequencing