

MODULARIZACE VÝUKY EVOLUČNÍ A EKOLOGICKÉ BIOLOGIE CZ.1.07/2.2.00/15.0204



HYBRIDIZATION AND HYBRID ZONES

















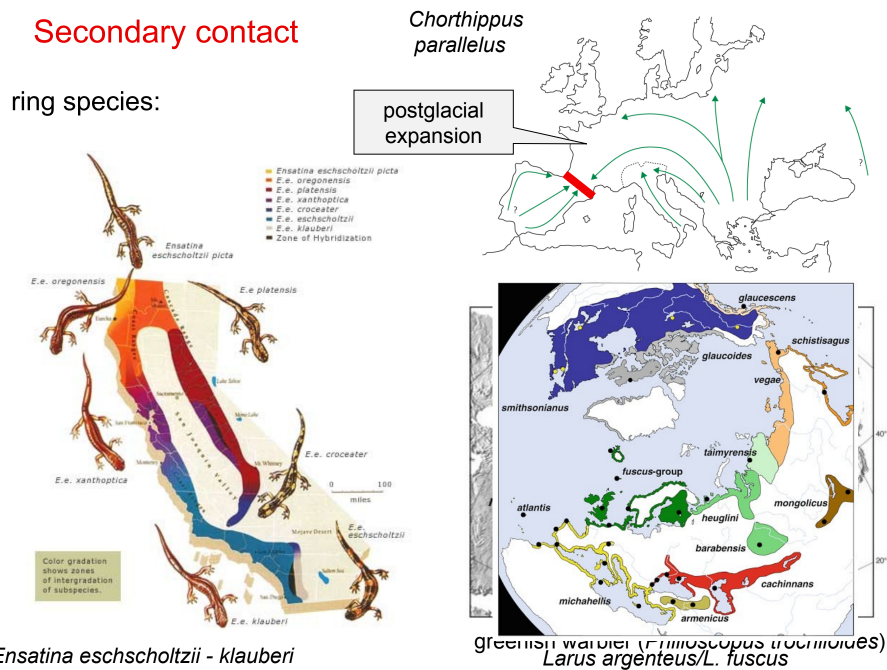












Ensatina eschscholtzii - klauberi

Hybridization:

25% species of vascular plants

10% species of animals

probably underestimation (only conspicuous species: ducks, birds of paradise, butterflies)

often result of environmental disturbance:

eg. "Darwin's finches" Geospiza fuliginosa, G. fortis and

G. scandens after El Niño event



Geospiza fuliginosa

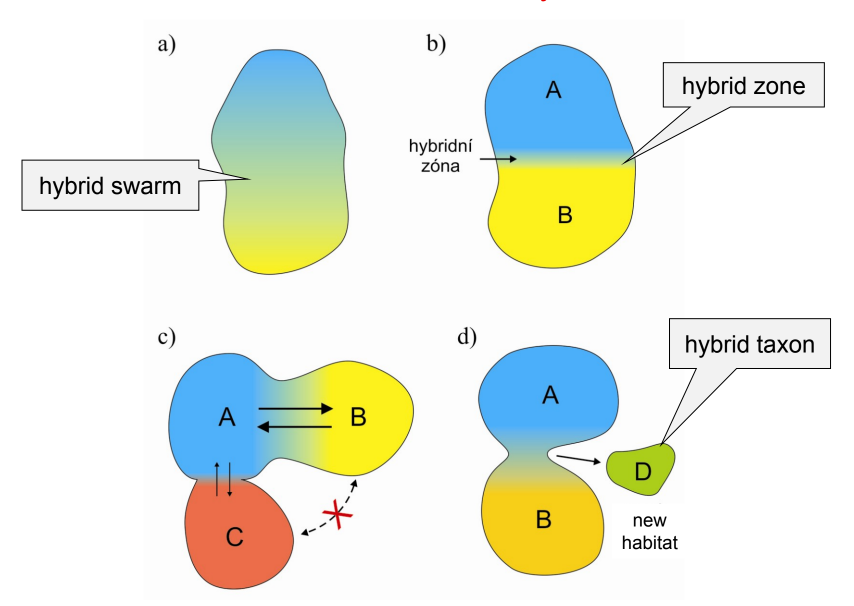


G. fortis



G. scandens

Possible outcomes of hybridization



Program NewHybrids:

A Model-Based Method for Identifying Species Hybrids Using Multilocus Genetic Data

E. C. Anderson¹ and E. A. Thompson

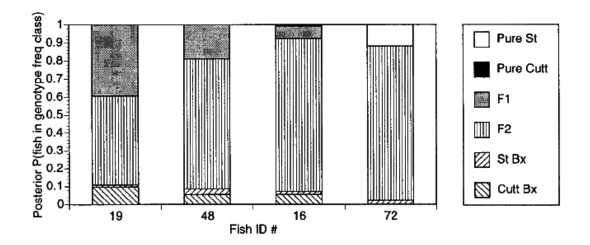
Department of Statistics, University of Washington, Seattle, Washington 98195

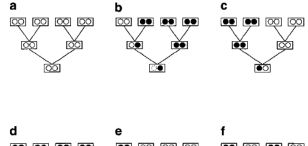
Manuscript received October 3, 2001

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ABSTRACT

We present a statistical method for identifying species hybrids using data on multiple, unlinked markers. The method does not require that allele frequencies be known in the parental species nor that separate, pure samples of the parental species be available. The method is suitable for both markers with fixed allelic differences between the species and markers without fixed differences. The probability model used is one in which parentals and various classes of hybrids (F₁'s, F₂'s, and various backcrosses) form a mixture from which the sample is drawn. Using the framework of Bayesian model-based clustering allows us to compute, by Markov chain Monte Carlo, the posterior probability that each individual belongs to each of the distinct hybrid classes. We demonstrate the method on allozyme data from two species of hybridizing trout, as well as on two simulated data sets.





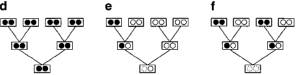
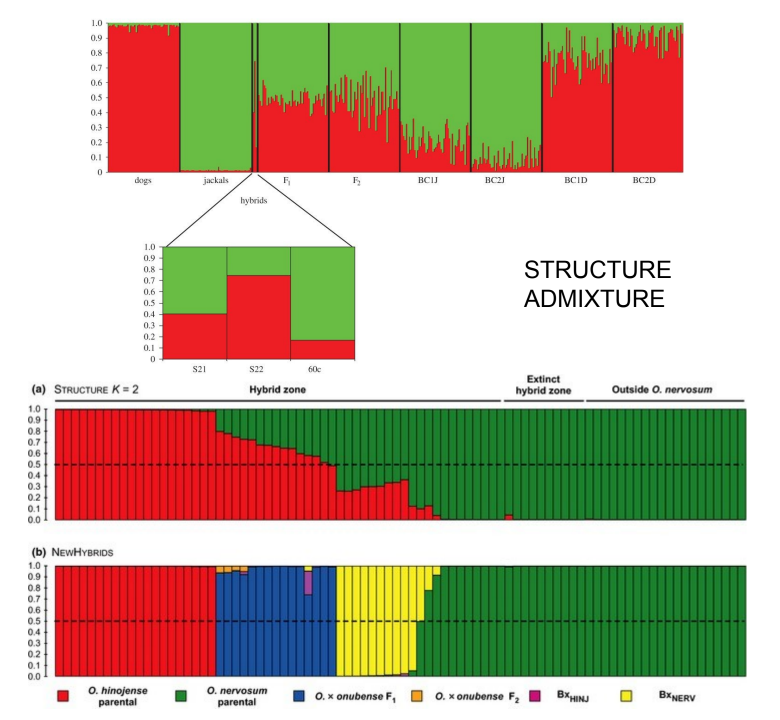


FIGURE 1.—Six arrangements of founders on a pedigree of n=2 generations. Each box represents a locus. The circles within each box represent the two genes possessed by the diploid organism at the locus. The founders are the individuals in the top row of each pedigree. Black gene copies are those originating from the species A population, and the white genes are from species B. Genes that are not determined to be either black or white by the pedigree and the founders in it are denoted by broken circles. The individual at the bottom of each pedigree belongs to a different hybrid class, determined by the arrangement of species among the founders. a–f represent six distinct genealogical classes. a–f also represent six distinct genealogical classes. There are, however, only five distinct gene frequency classes, the individuals at the bottoms of pedigrees c and f are both in the same gene frequency classe.



Hybrid zone (Barton a Hewitt 1985)

= area, where genetically different populations meet, mate and give rise at least some hybrid offspring

Hybrid zones may be classified as:

primary secondary

tension, mosaic, staggered, "mottled" ...

extrinsic selection (external environment) intrinsic selection (prezygotic or postzygotic barriers)

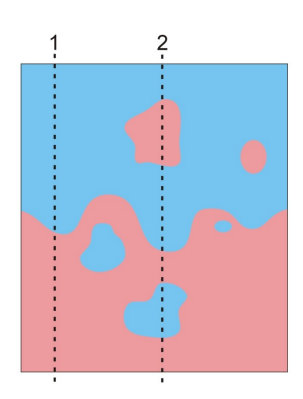
Mosaic hybrid zone:

influence of environment in fact a set of several hybrid zones

eg.: *Gryllus firmus* x *G. pennsylvanicus* (NE USA) sandy x clayish soils





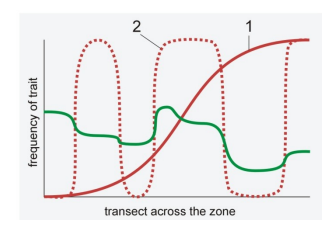


Iris fulva x I. brevicaulis:

I. fulva is limited to more forested sites



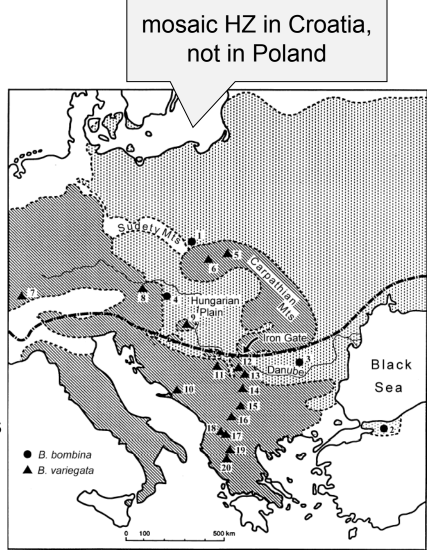




fire-bellied toad *B. bombina*:

lowlands
mostly in water
larger water surfaces
thiner skin
territorial
530 Hz
longer development

Bombina:





yellow-bellied toad *B. variegata*:

hills, highlands terrestrial mating in puddles thick skin nonterritorial 580 Hz shorter development

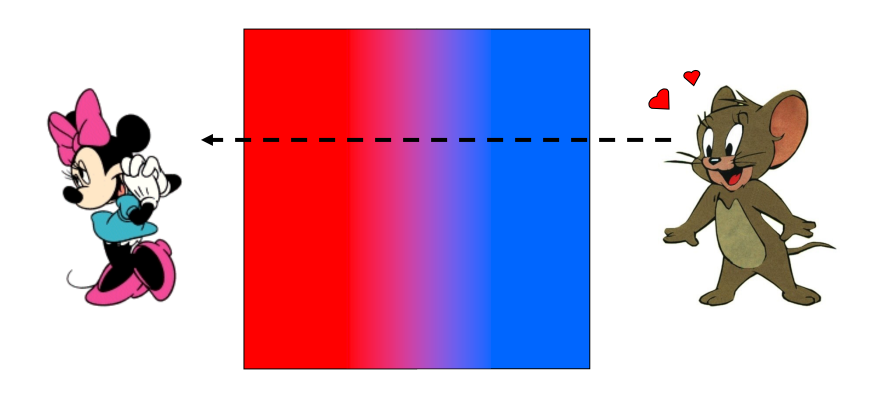


Most hybrid zones are tension zones.

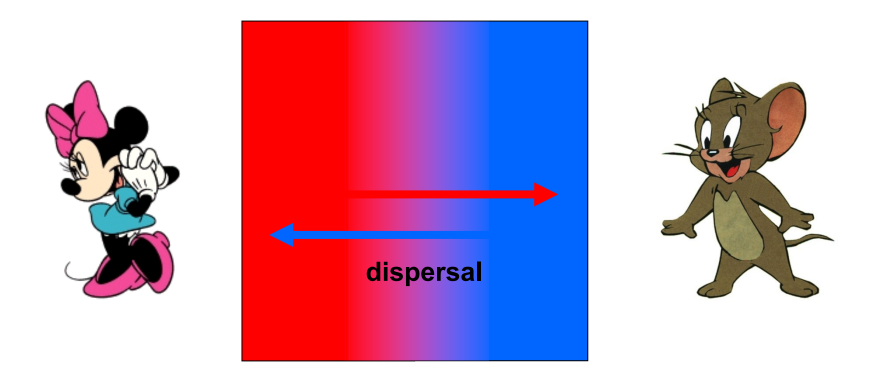
Nick Barton

... i.e., they are maintained by balance between dispersal and selection (Barton & Hewitt, 1985)

Tension zone is when...

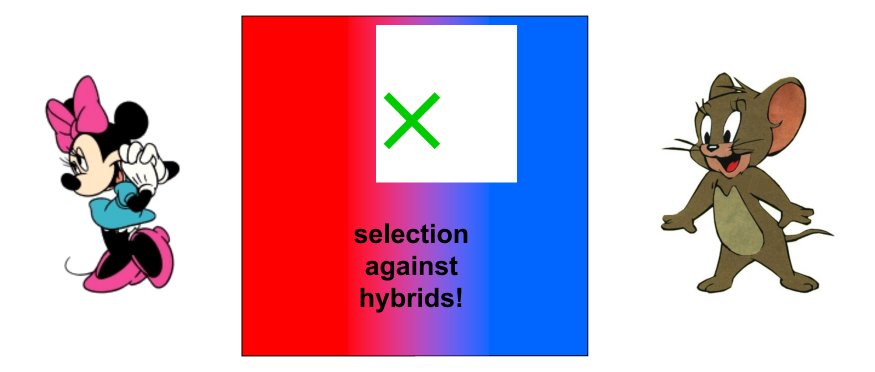


Tension zone is when...



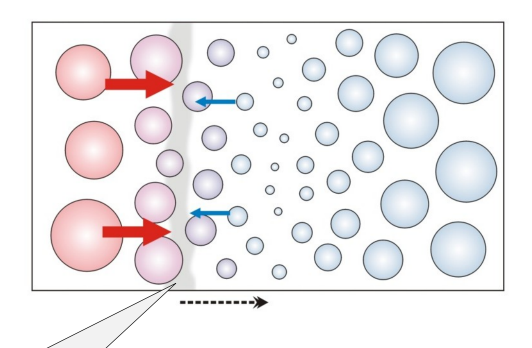
 \Rightarrow zone widening

Tension zone is when...



⇒ zone narrowing

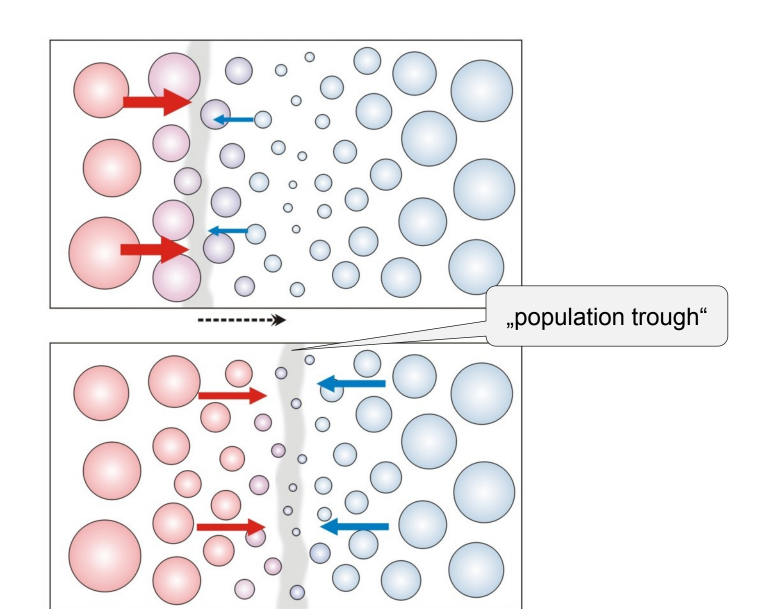
Tension zone is maintained by dynamic equilibrium between *dispersal* and *selection*

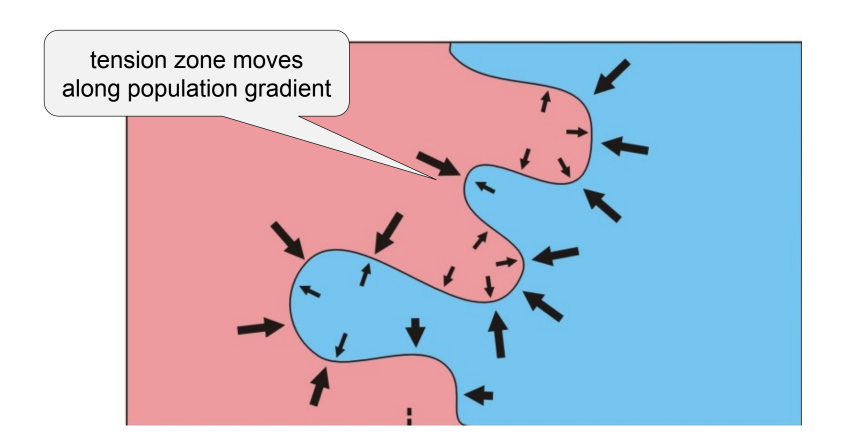


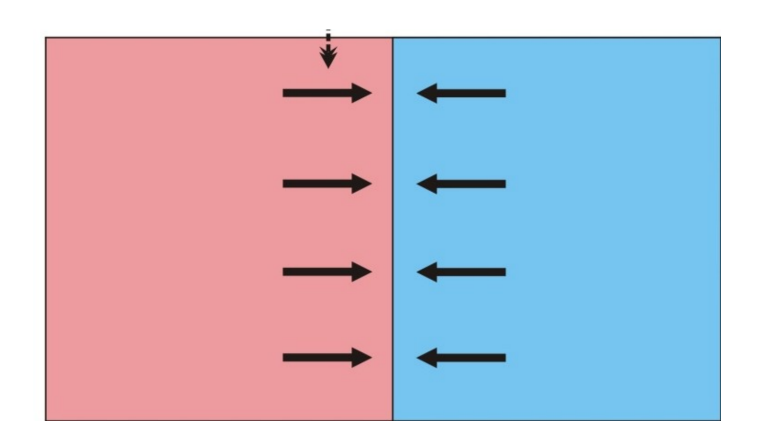
tension zone moves along the gradient of population density

Tension zone is independent of external conditions (*intrinsic selection*)

⇒ its movement ends at a geographical barrier or in the area
of the lowest population density (*"population/density trough*")





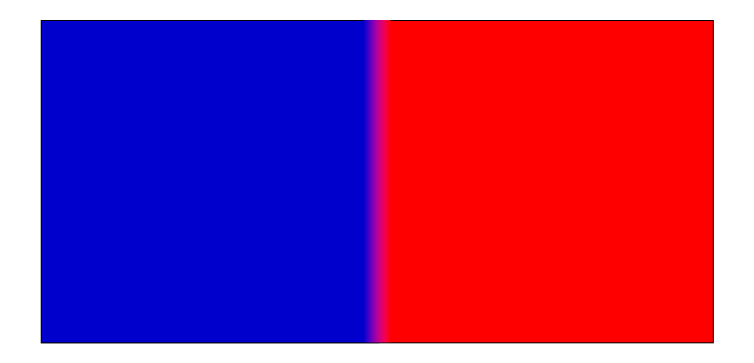


Theory of cline:

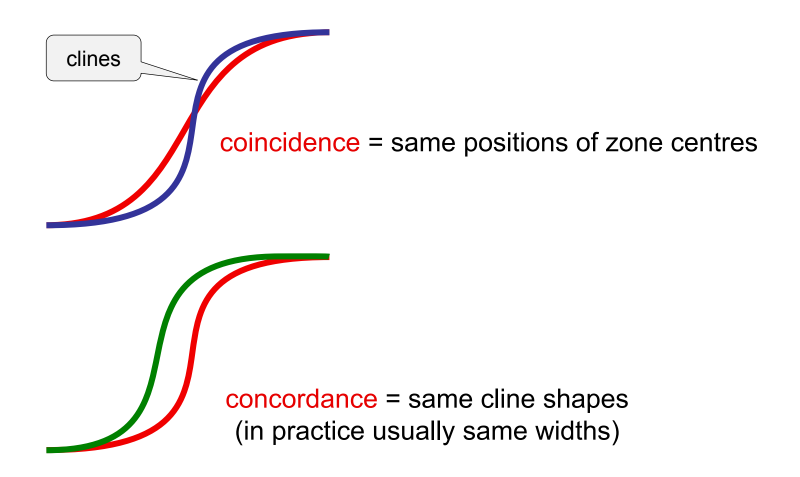
secondary contact:

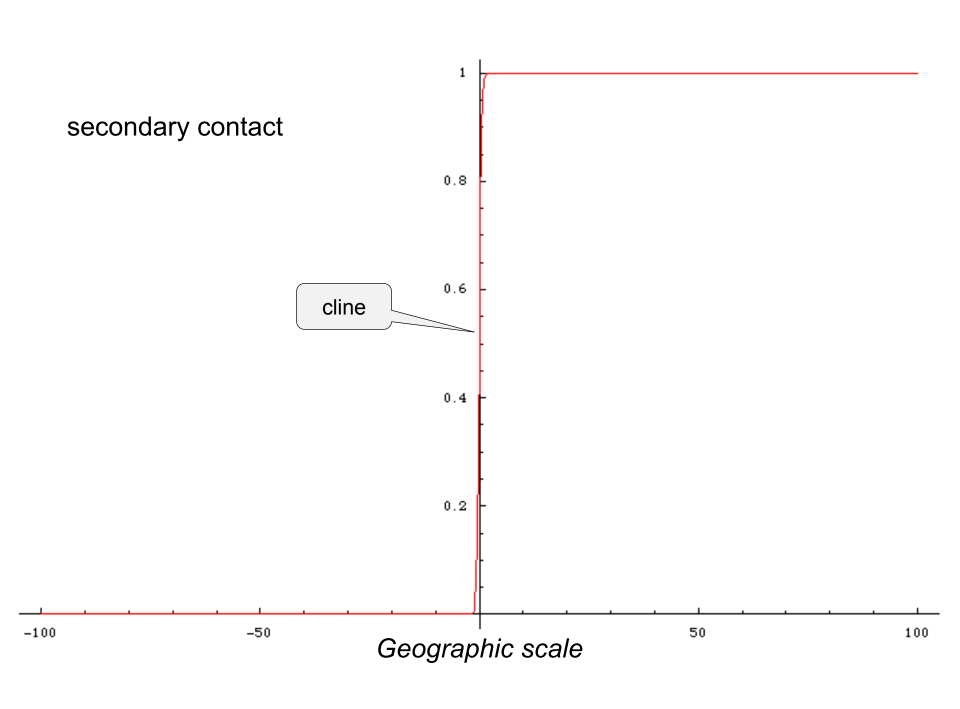


coincident and concordant clines



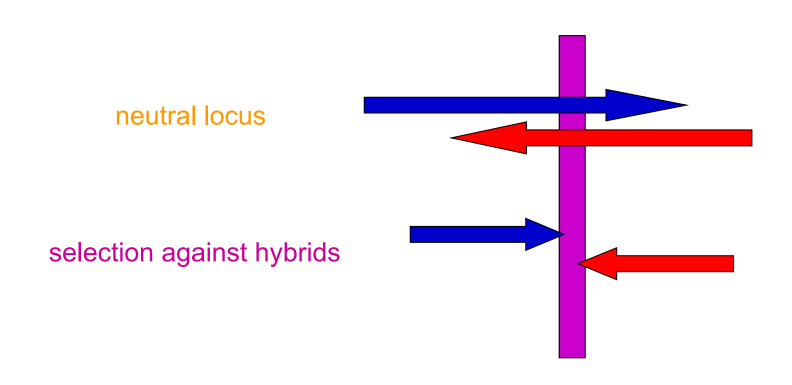
Cline = gradient of trait(s) (eg. allele frequency or mean of quantitative trait) across spatially continuous habitat

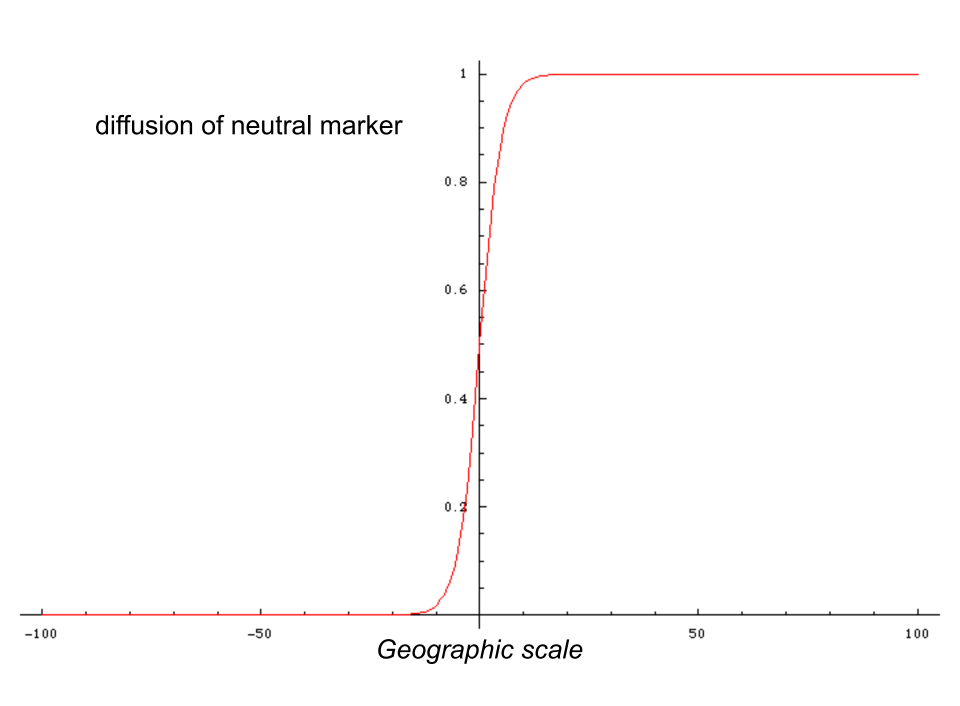


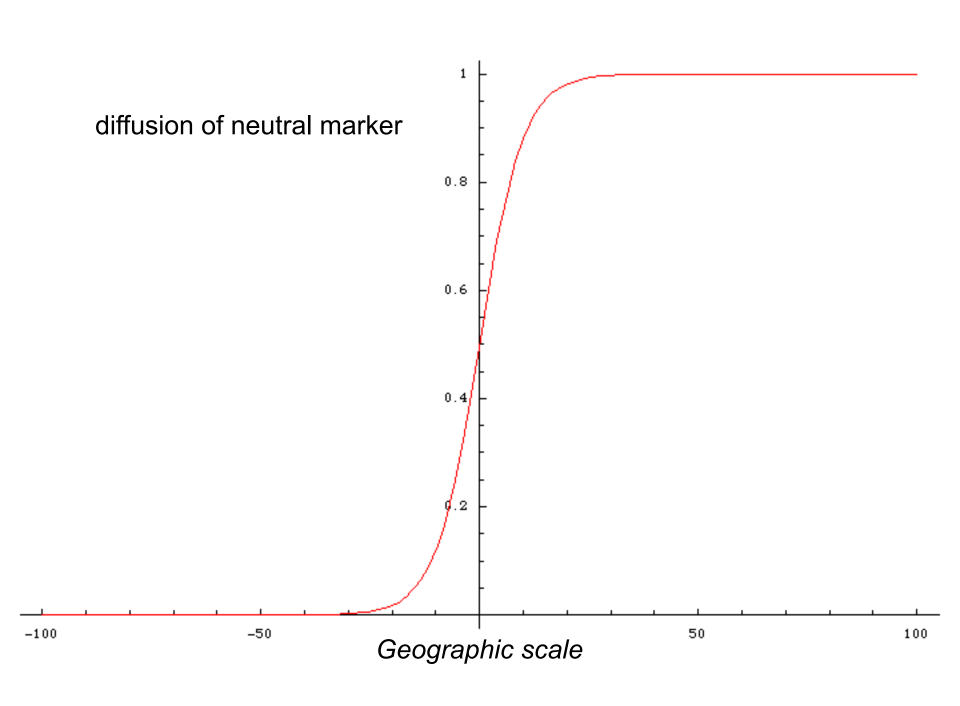


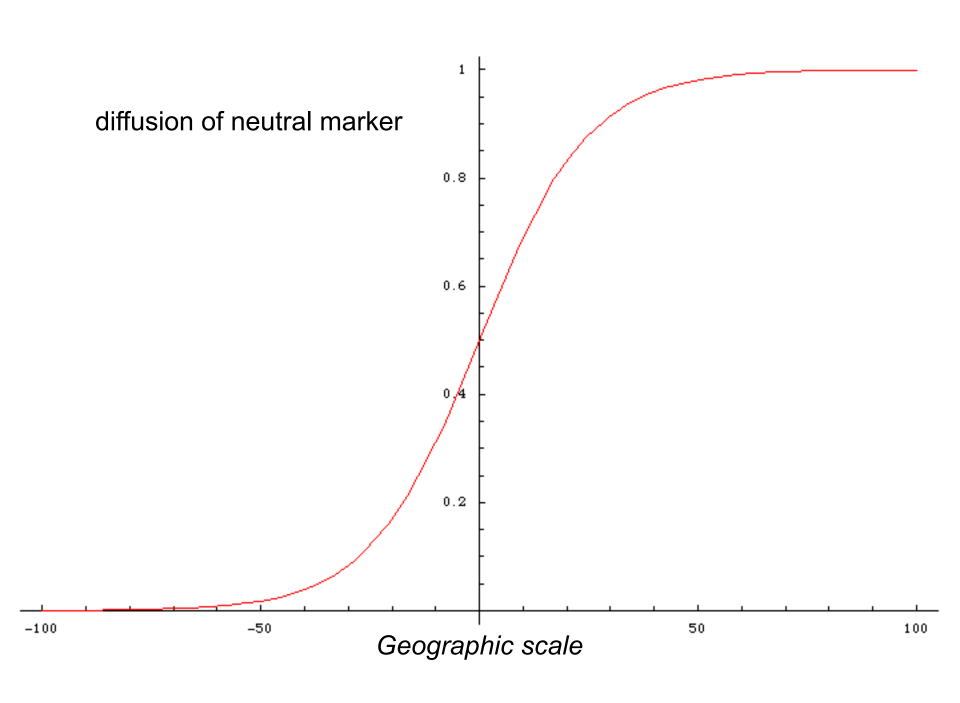
Theory of cline:

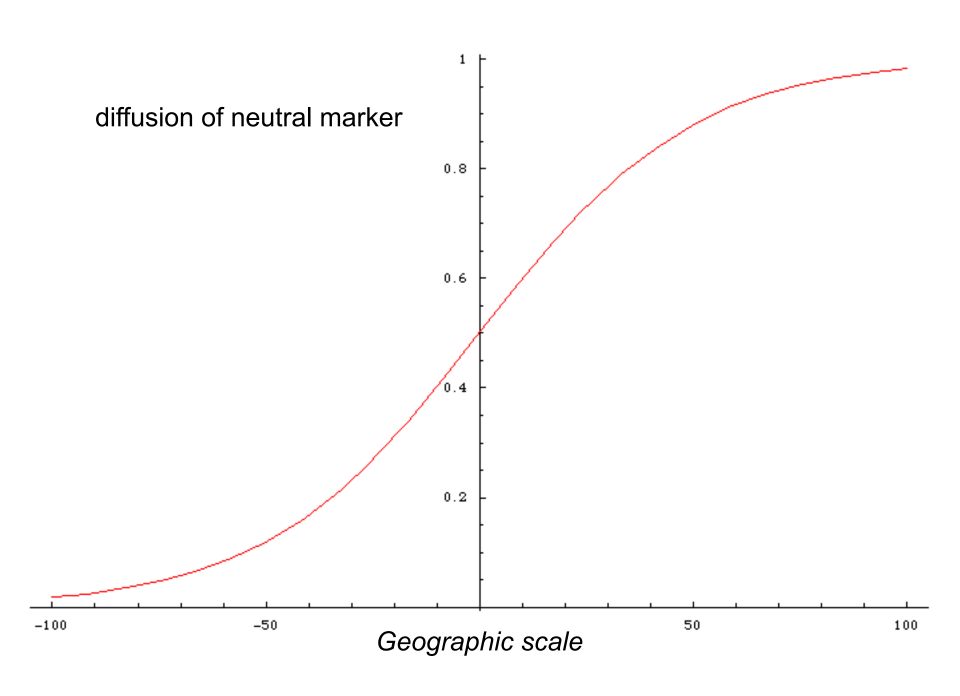
neutral vs. selected loci





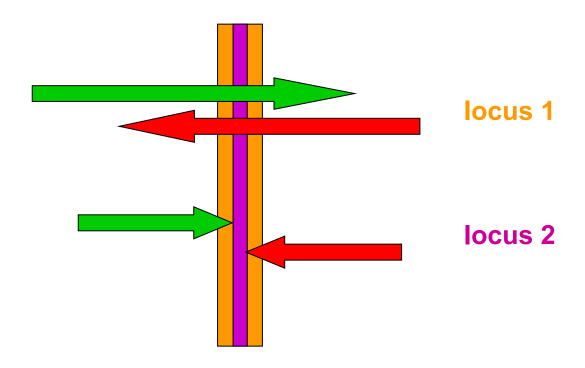




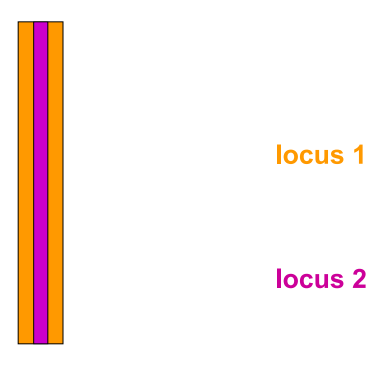


neutral vs. selected loci

with time, concordance is disappearing ...

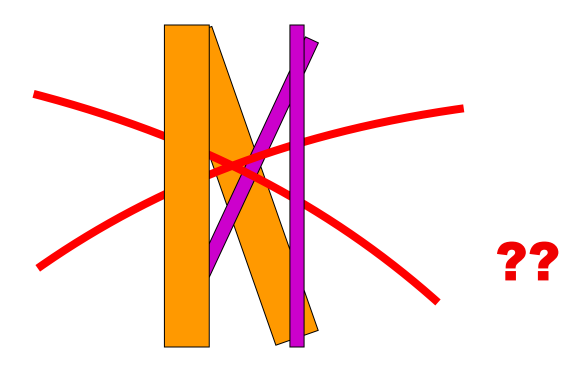


... but (in tension zone) selection pushes clines for individual loci to each other ⇒ maintains coincidence sometimes ...



... but clines still parallel

cline models (diffusion approximation etc.), linkage disequilibrium, evolutionary parameters



problem, how to analyse

Hybrid zone study

- Sampling along linear or 2D transect, geographic coordinates of localities
- Genetic (morphological, behavioural etc.) analysis
 problem of sample independence (F_{ST}, F_{IS} ... effective No. alleles)
- 3. Geographic clines
- 4. Estimation of dispersal, selection, and other parametres
- 5. Alternative approaches: monotonic clines2D analysis genomic clines concordance analysis

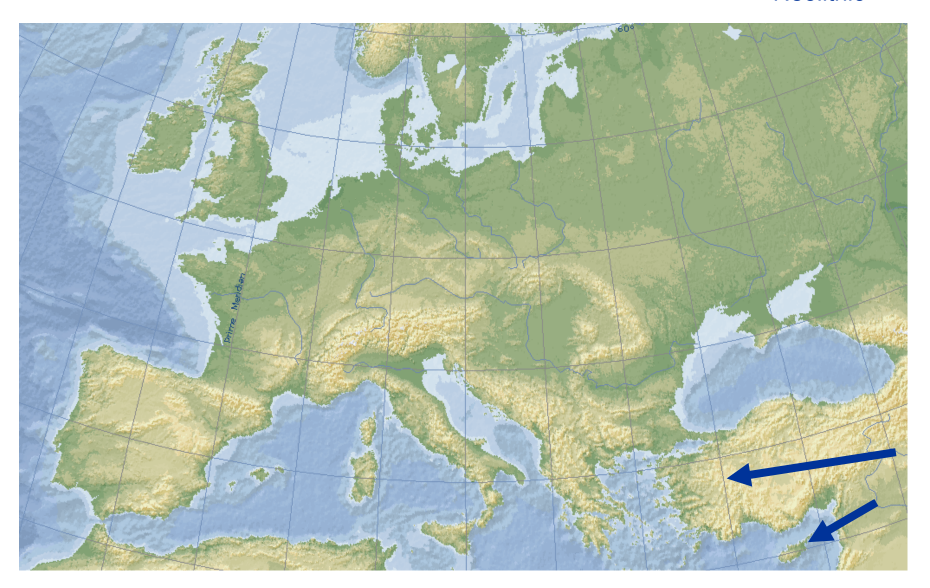
Case study: house mouse hybrid zone





Mouse colonization of Europe

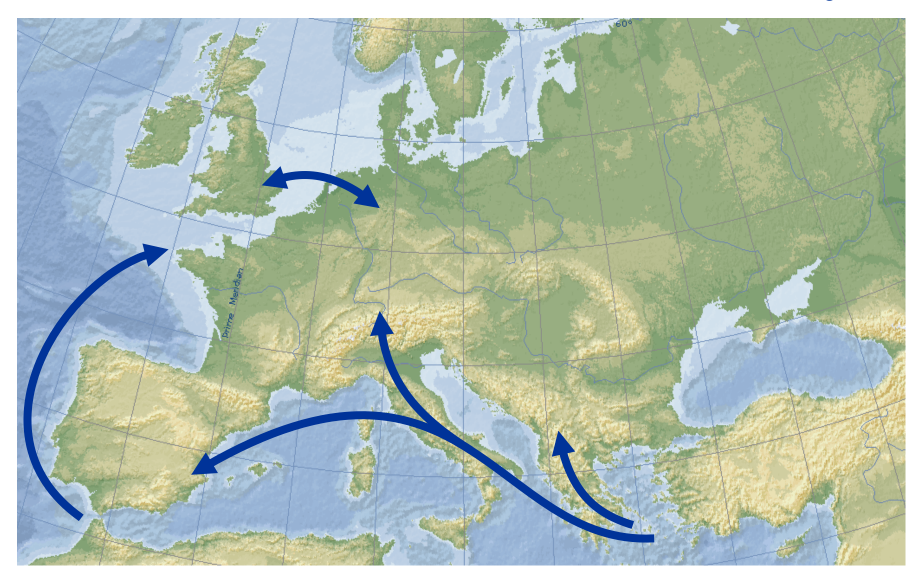
Neolithic



Cucchi et al. (2005)

Mouse colonization of Europe

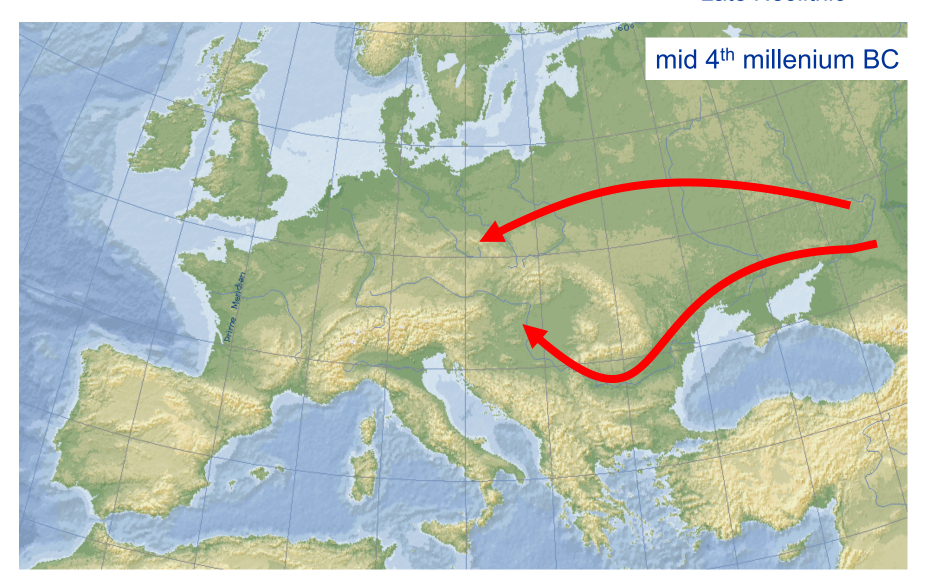
Bronze and Iron Age



Cucchi et al. (2005)

Mouse colonization of Europe

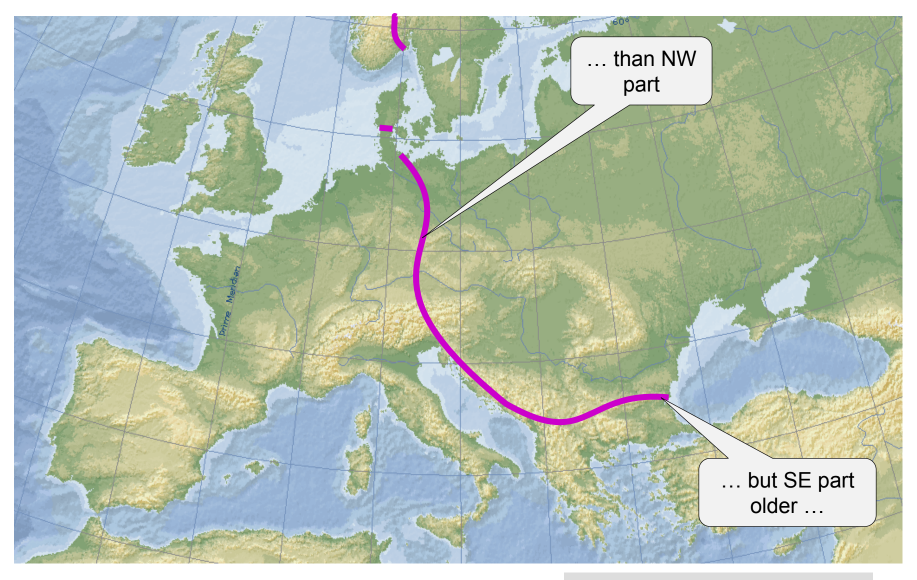
Late Neolitnic



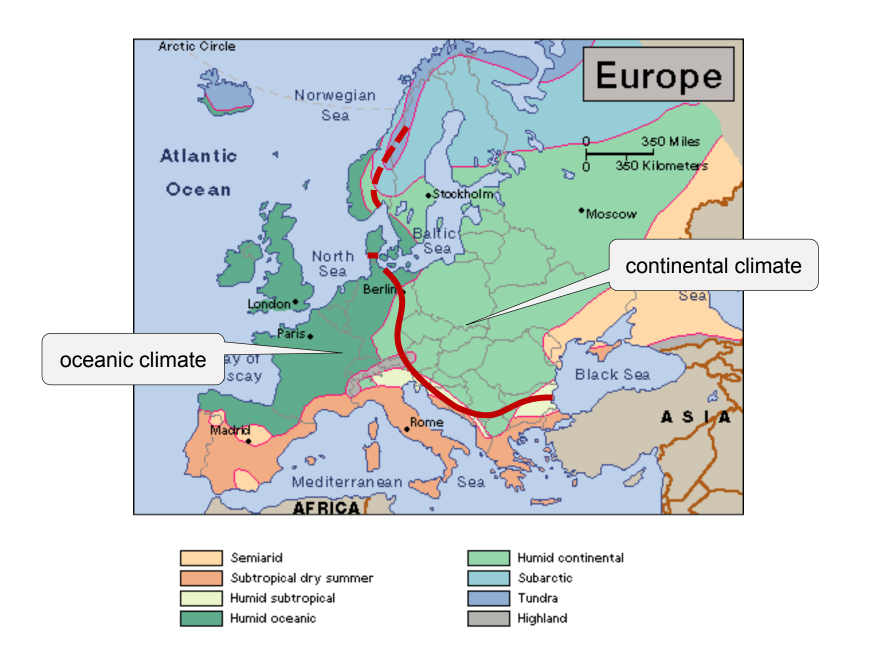
Cucchi et al. (2011)

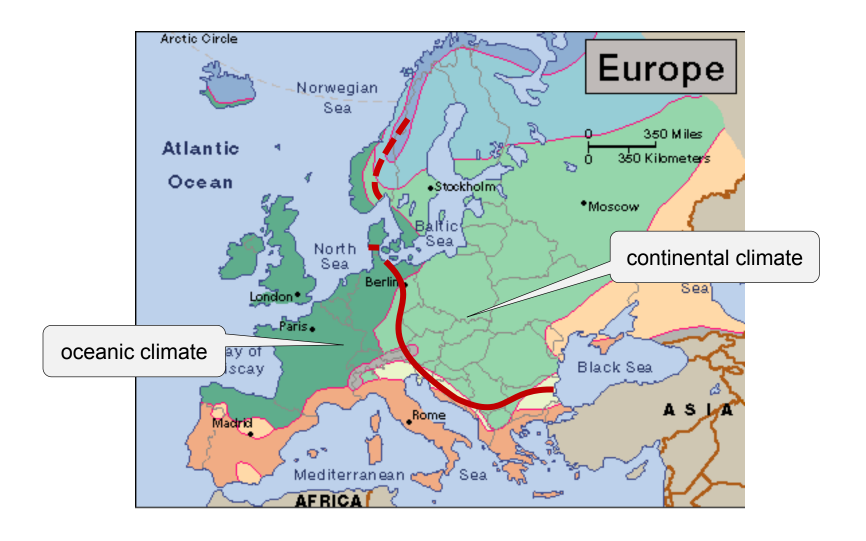
Hybrid zone in Europe musculus time of origin unknown... domesticus

Hybrid zone in Europe



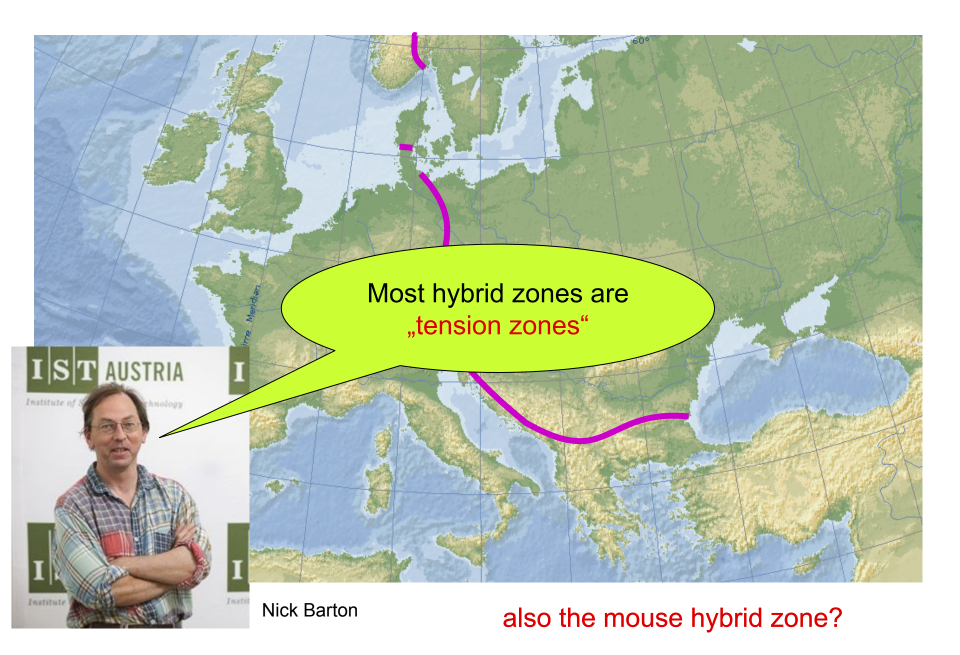
Co tuto zónu ovlivňuje?

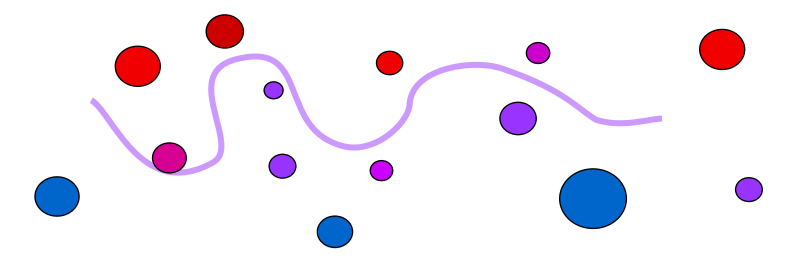




climatic factors don't determine

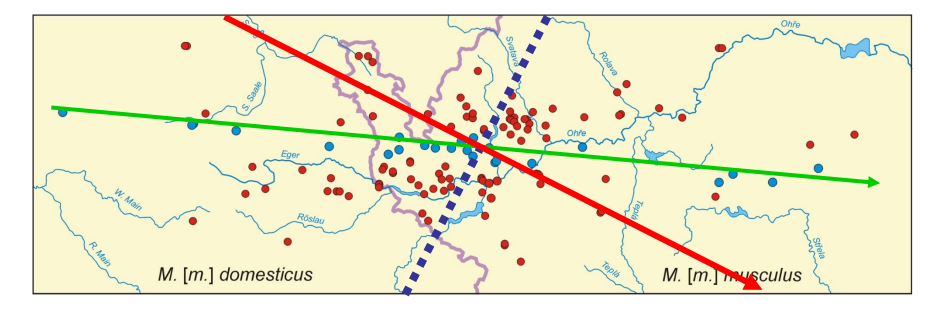
Hybrid zone in Europe

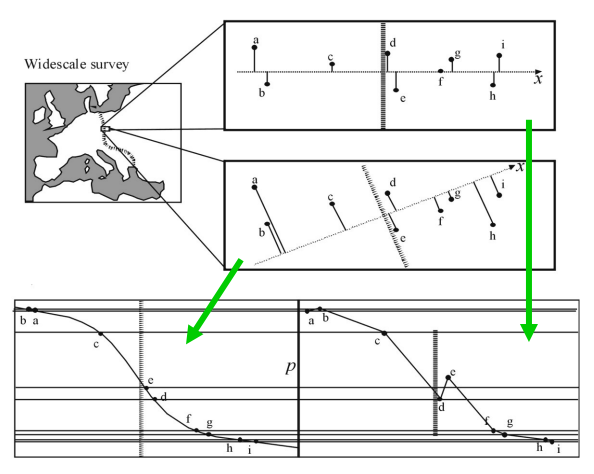




hybrid zone course may be complex....

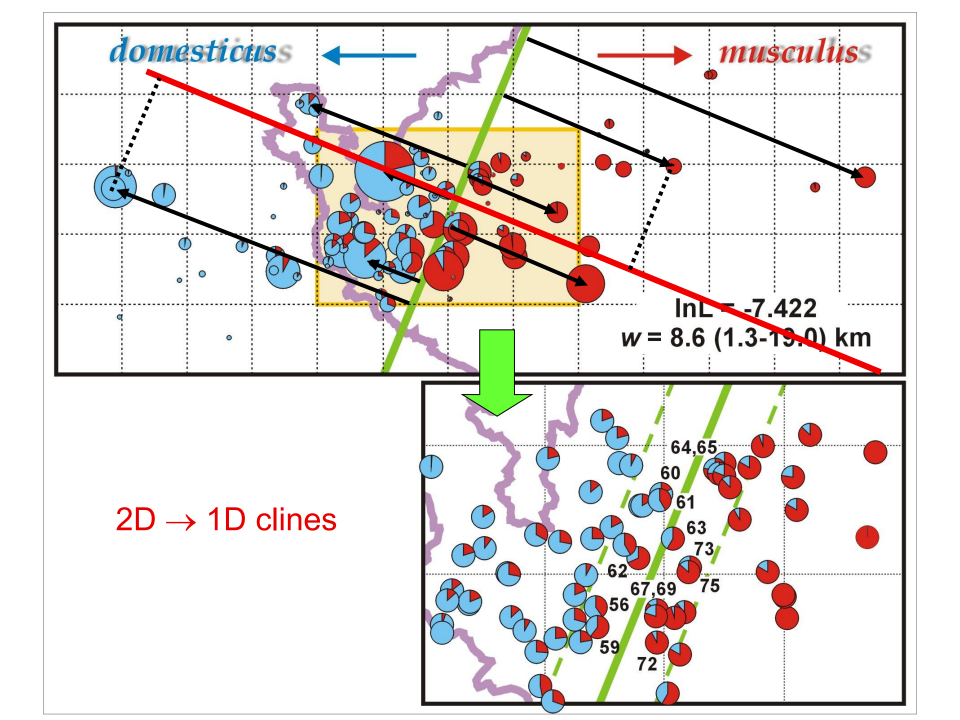
... moreover, usually we don't know a priori, or we extrapolate from global direction



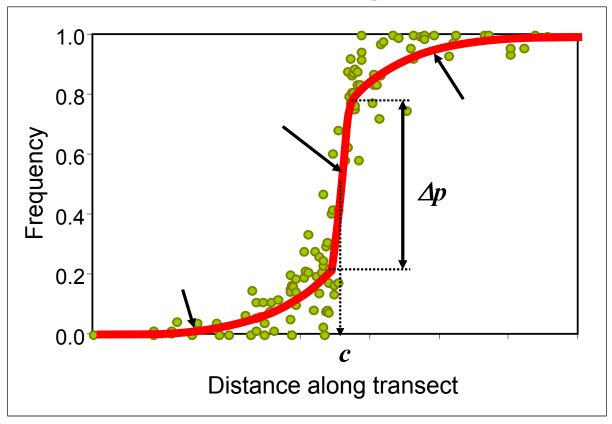


Real local cline

Cline interpolated from widescale survey



Multiple genes:



"stepped" model (symetrical, asymmetrical)

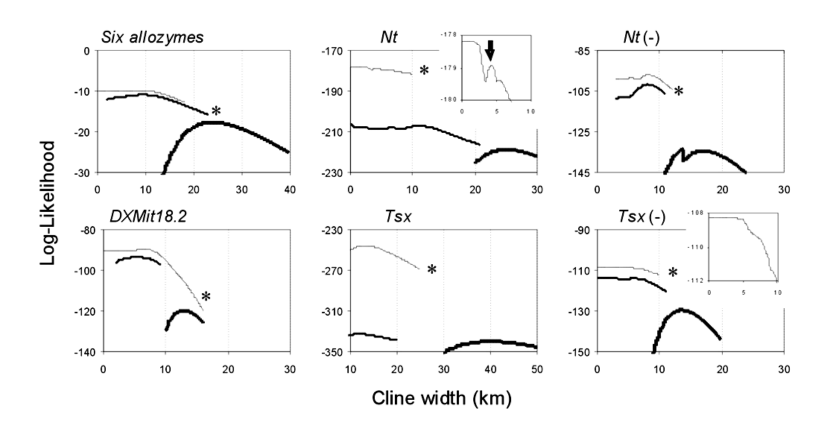
linkage disequilibrium resulting from influx of parental allele combinations ⇒ synergistic effect: strenghtening of selection in zone centre ⇒ central step × introgression tails reflect selection at individual loci

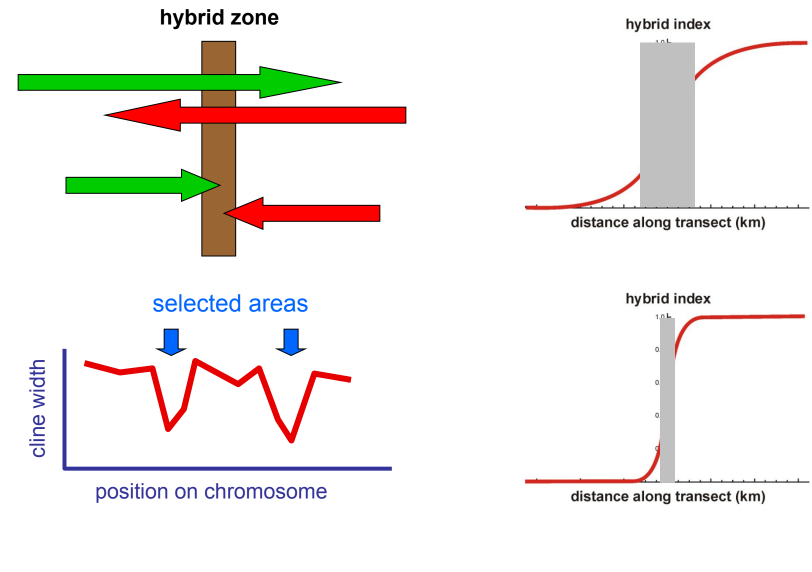
We can estimate some other key evolutionary parametres from LD and cline parametres:

| dispersal: | |
|---------------------------------|--|
| effective selection: | |
| selection on marker loci: | |
| selection on selected loci: | |
| fitness of hybrids: | |
| number of loci under selection: | |

model comparison: LRT (they are nested); d.f. = difference in number of parametres

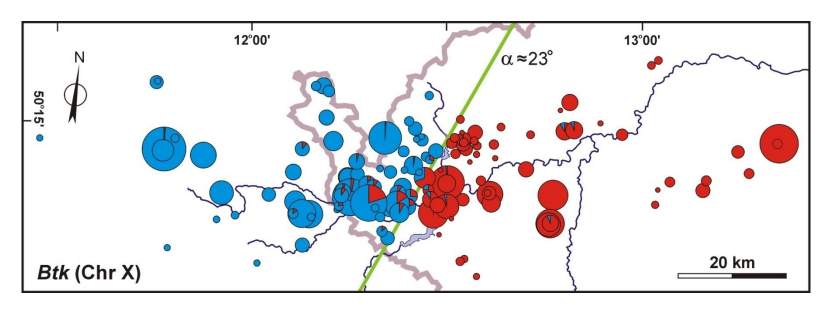
likelihood profiles:

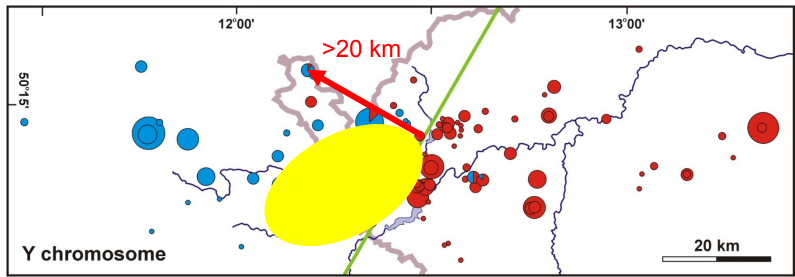




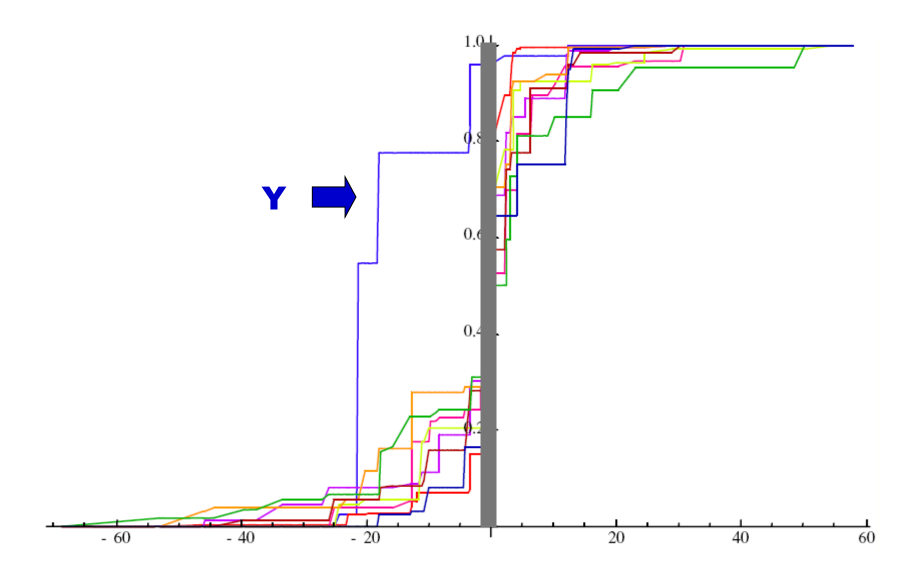


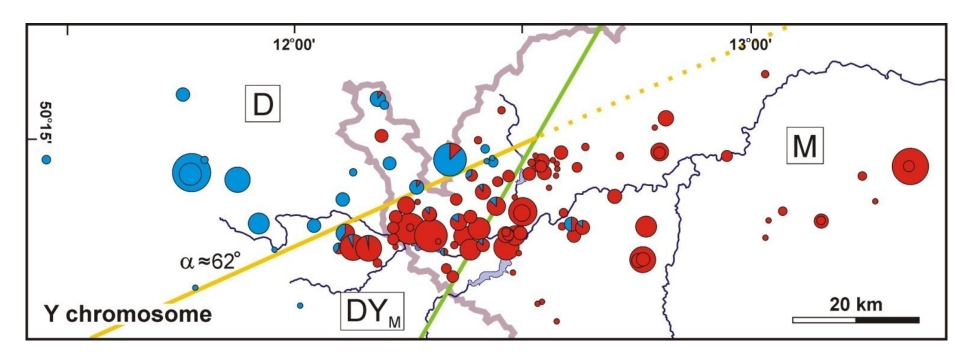
Problems – Y chromosome

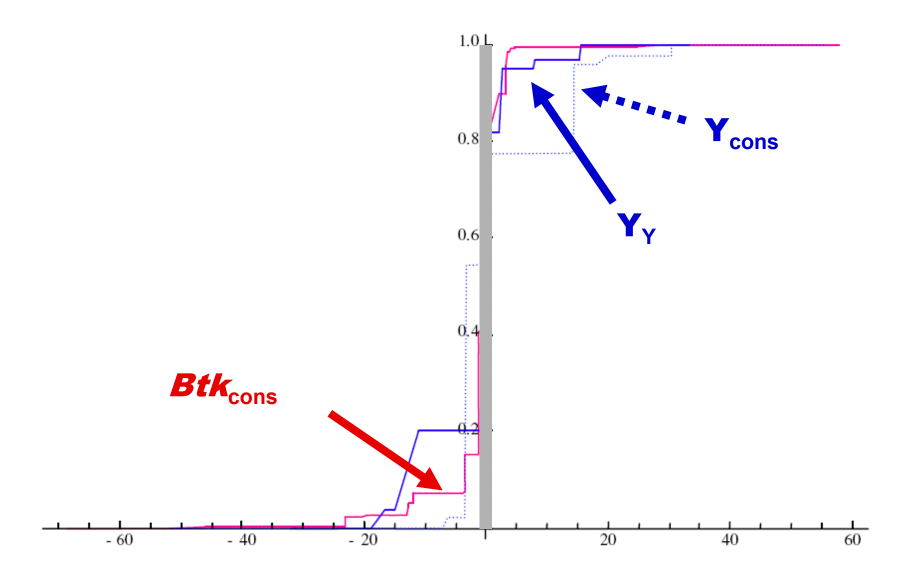


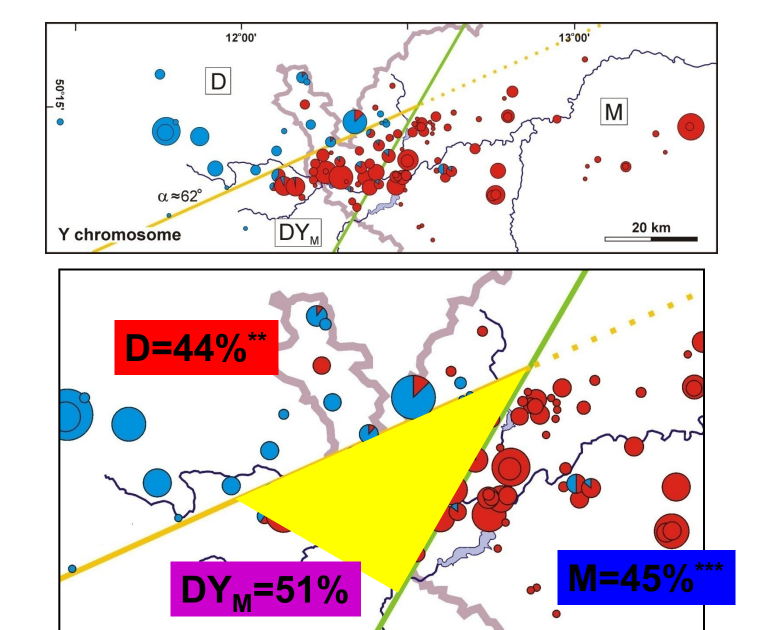


Monotonic clines – consensus orientation







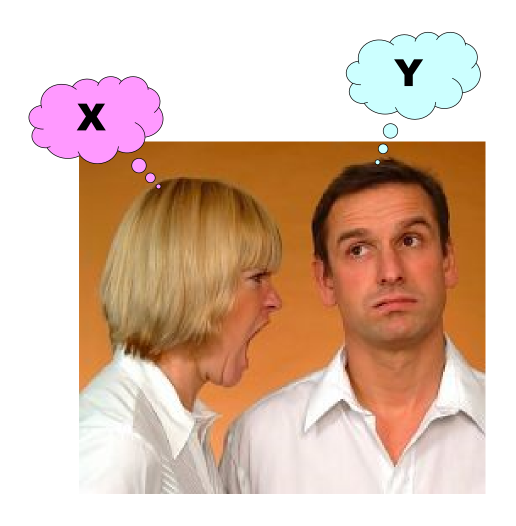


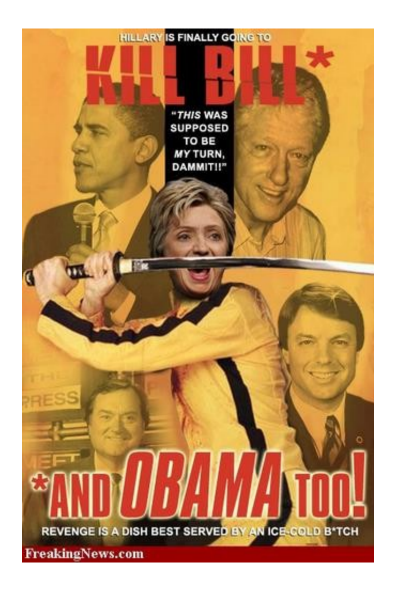
Weird behaviour of the Y in the hybrid zone – summary:

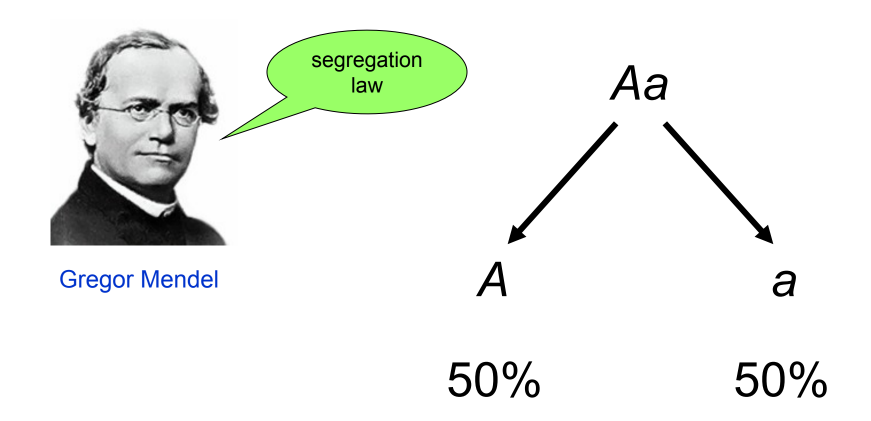
- 1. *musculus* Y <u>more successful</u> than *domesticus* Y <u>on</u> <u>its own genetic background</u>
- 2. <u>higer proportion of males</u> relative to other areas

Either coincidence, or ...

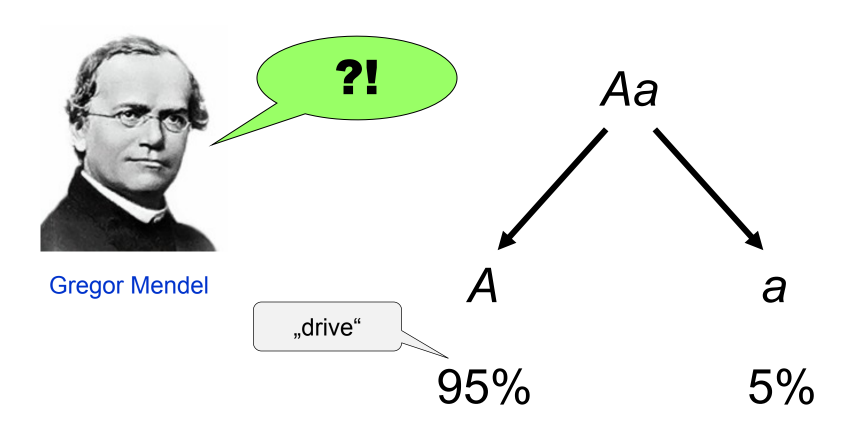
... or genetic conflict between X and Y and probably some autosomal genes as well





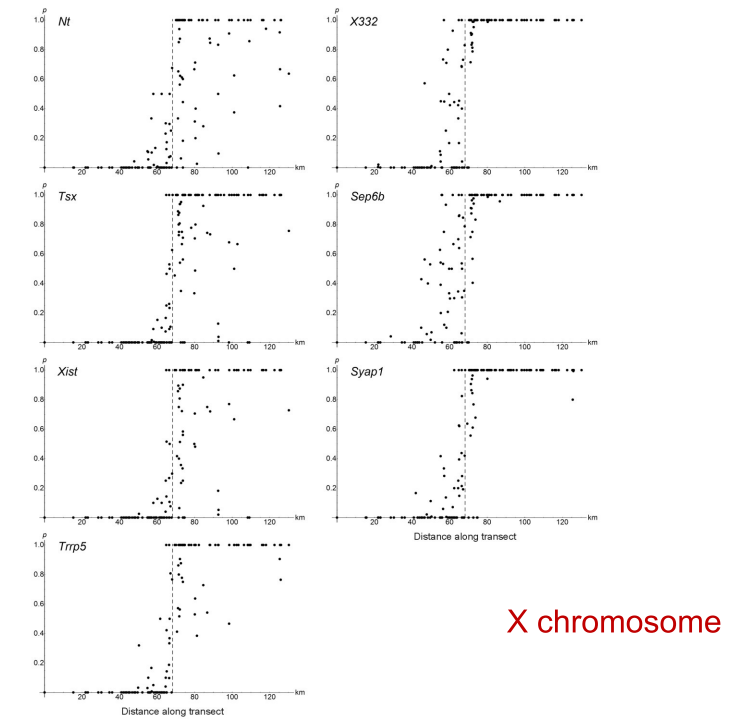


Intragenomic conflict results in higher proportion of a genomic element in the next generation

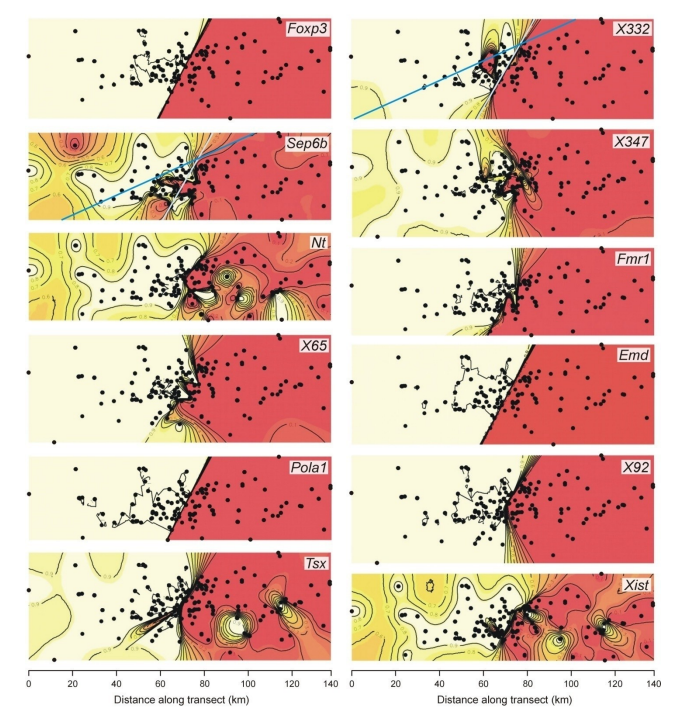


vychýlení segregačího (transmisního) poměru

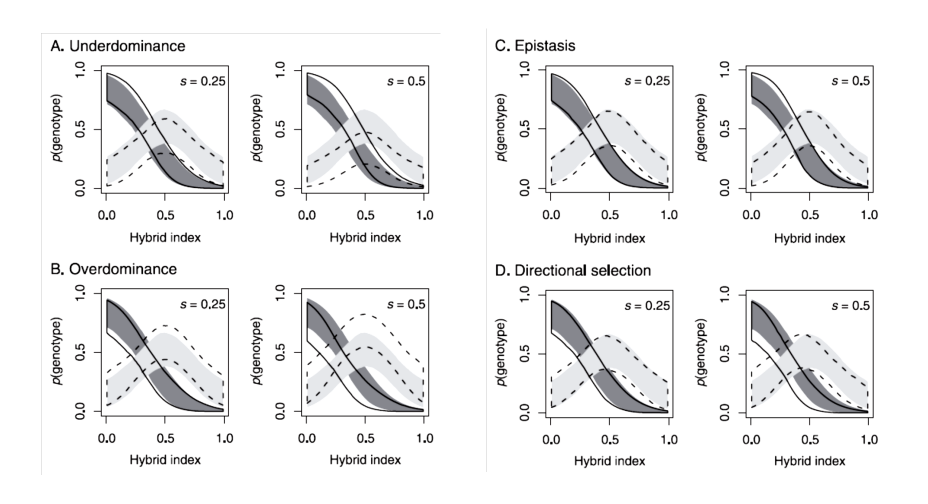
- = segregation distortion (SD)
- = transmission ratio distortion (TRD)



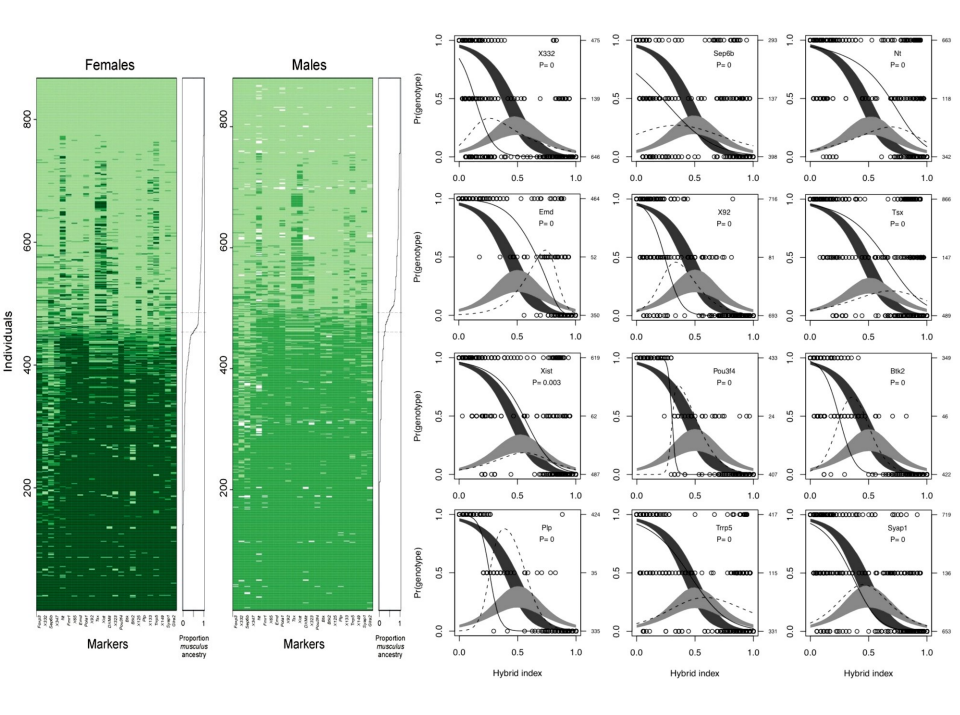
Chr. X - 2D analysis Geneland



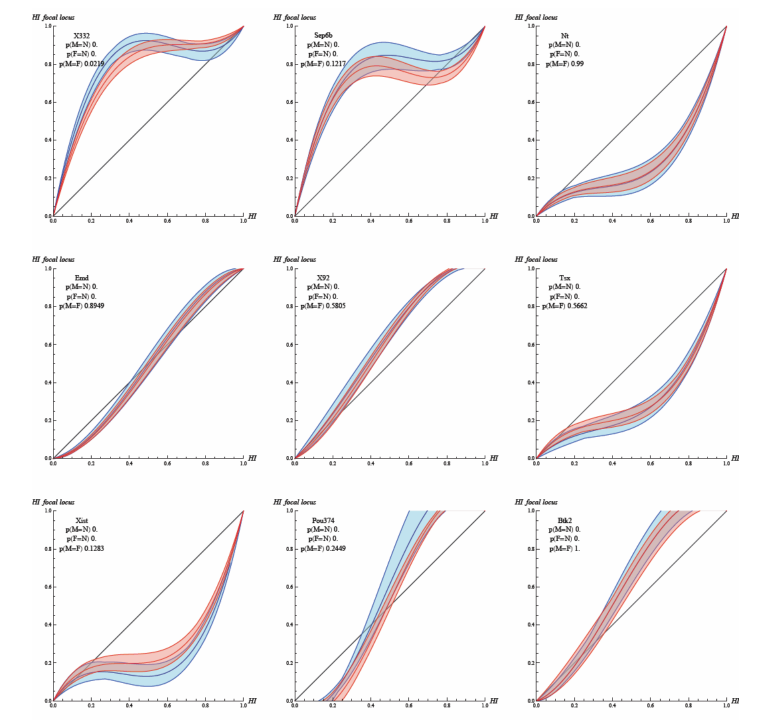
'Genomic clines'



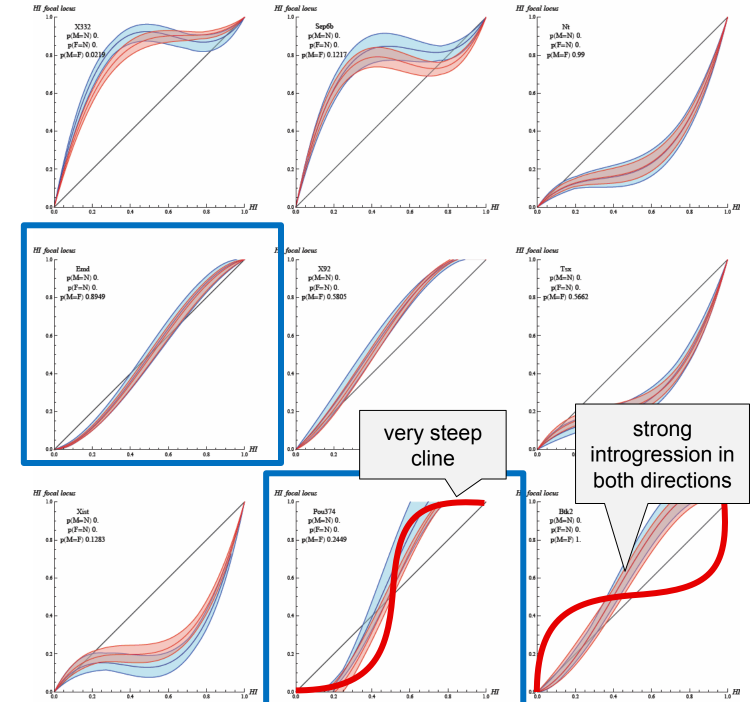
Z. Gompert & A. Buerkle



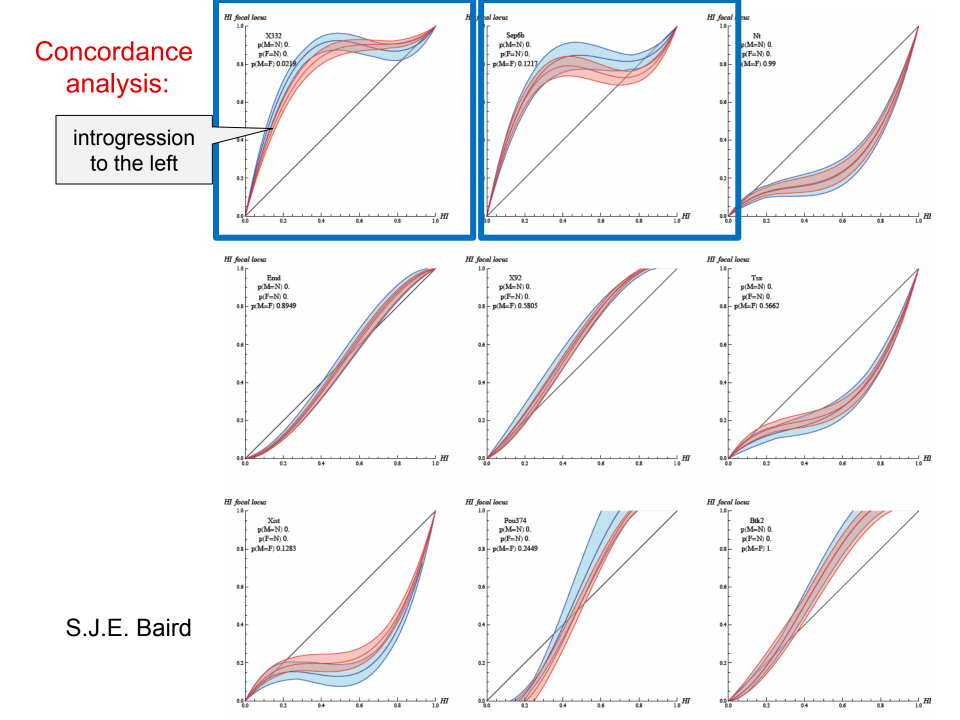
Concordance analysis:



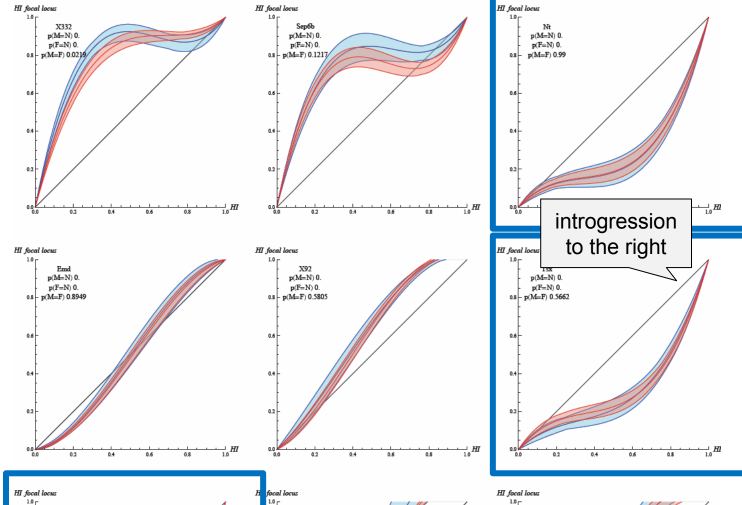
S.J.E. Baird

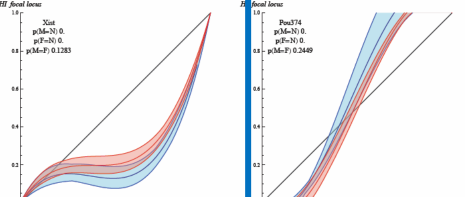


S.J.E. Baird

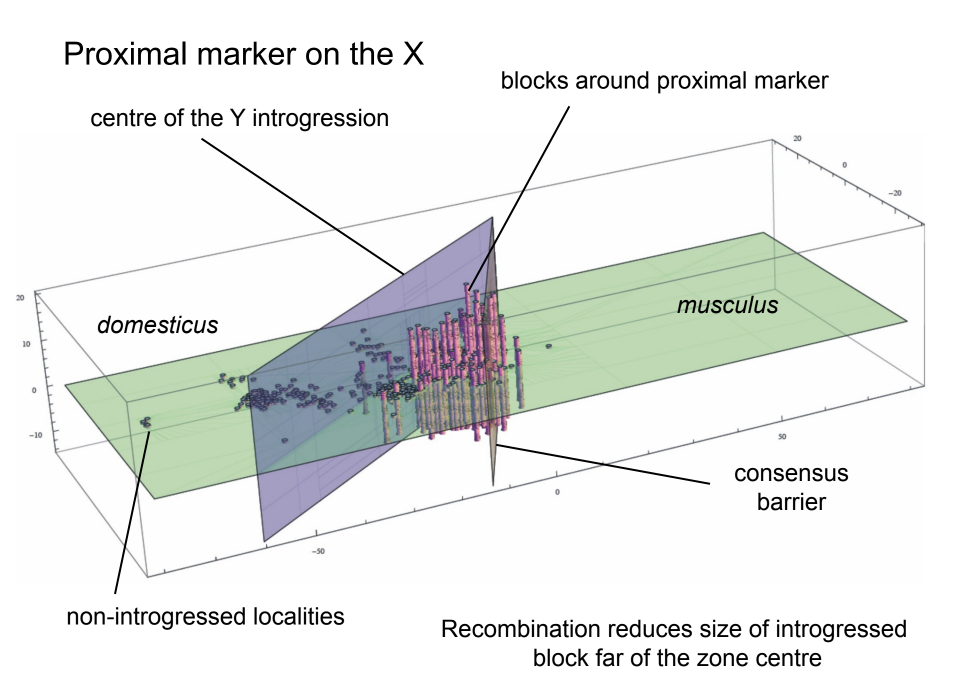


Concordance analysis:

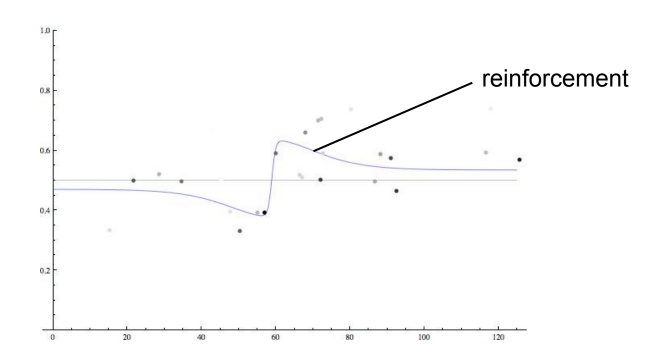




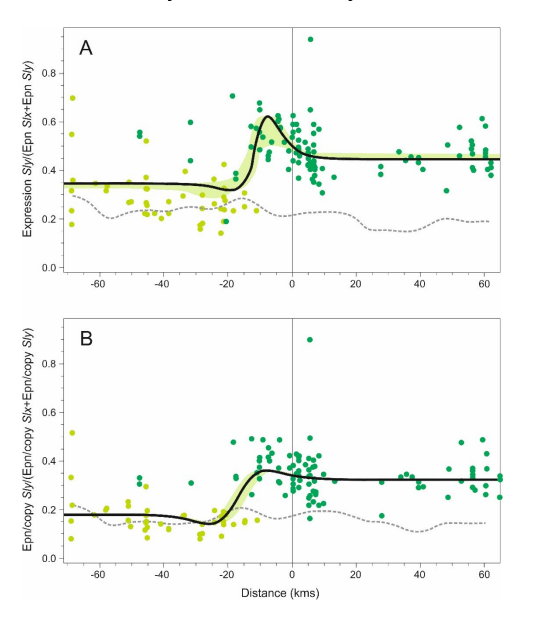
S.J.E. Baird



Using cline model for analysis of reinforcement – odour preference in the mouse hybrid zone



Using cline model for analysis of gene expression in the mouse hybrid zone – asymmetric model

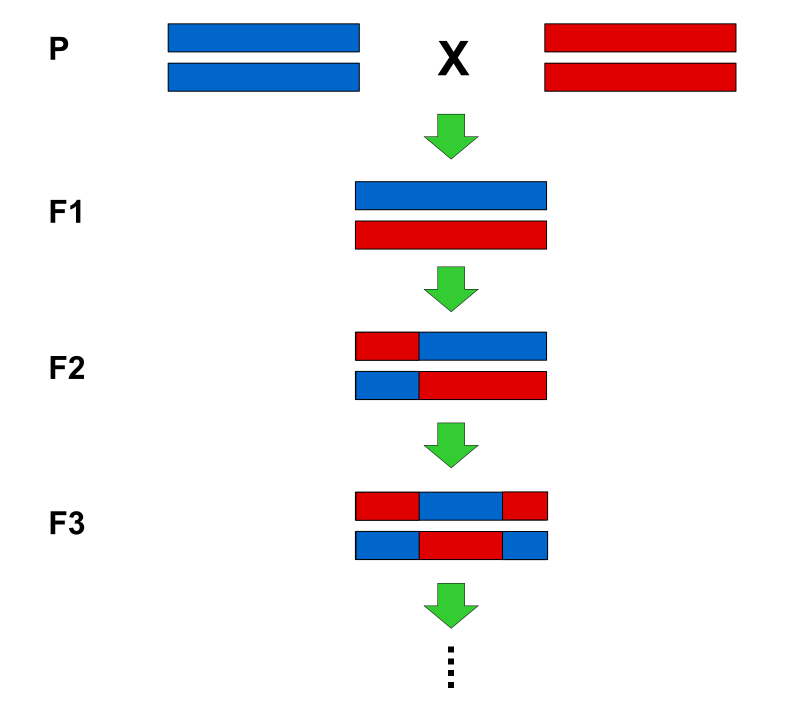




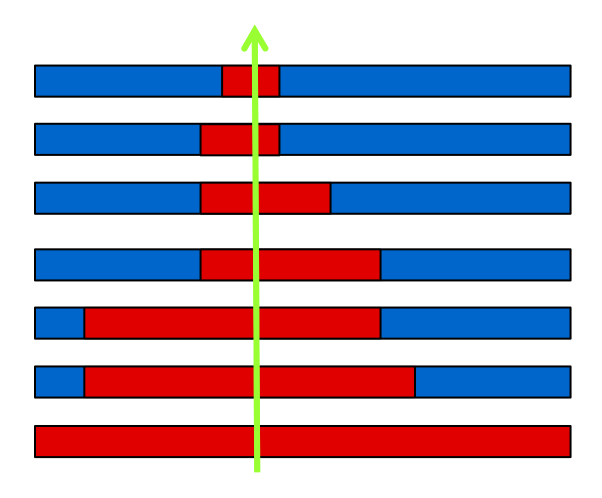
Neanderthal DNA specialist Svante Pääbo examines the femur, found near Ust'-Ishim in western Siberia. Photogi

Blocks of Neanderthal DNA found in modern humans can act like a biological clock, because they are fragmented more and more with each generation since interbreeding happened. The blocks of Neanderthal DNA in the Siberian man were on average three times longer than those seen in people alive today. Working backwards, the scientists calculate that Neanderthals contributed to the man's genetic ancestry somewhere between 7,000 and 13,000 years before he lived.

The findings, published in the journal Nature, suggest that humans and Neanderthals had reproductive sex around 50,000 to 60,000 years ago...



Hybridization makes a cascade of blocks



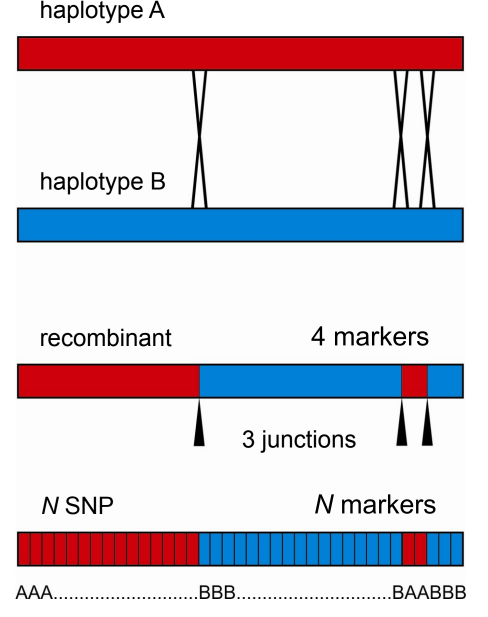
advancing introgression into 'blue' genome

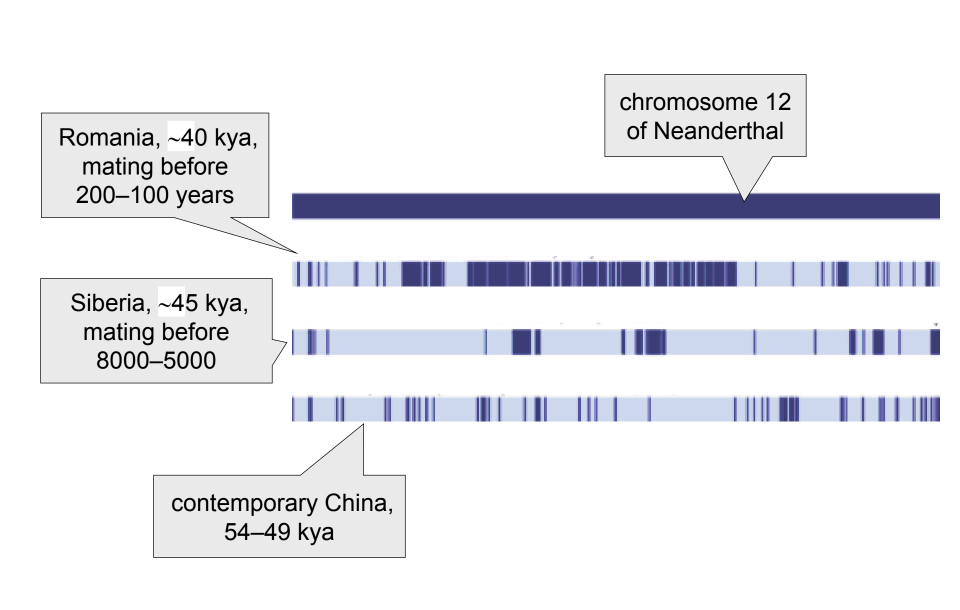


R. A. Fisher

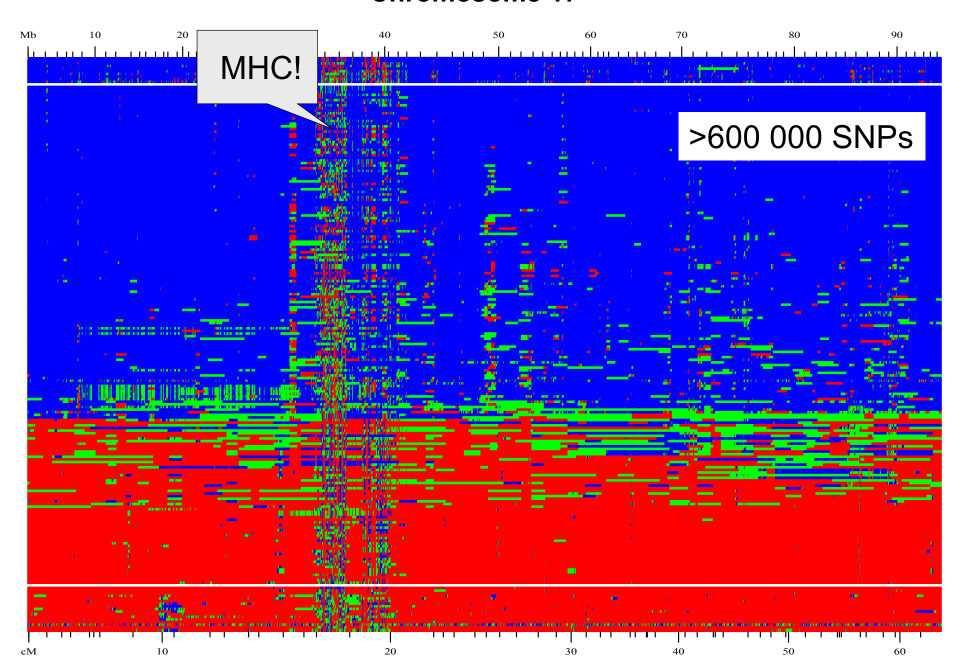
Recombination brings together DNA of different origin and makes *junctions* (*breakpoints*)

they divide genome into *blocks* (*chunks*, *tracts*, *segments*)

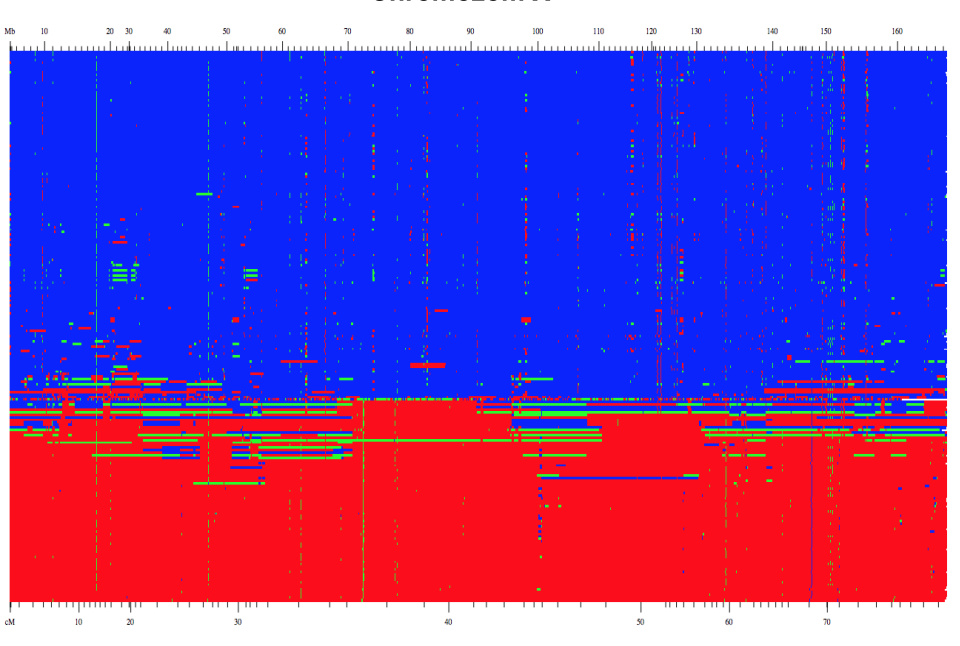




Chromosome 17



Chromozom X



Why hybrid zones? Reproductive barriers and speciation!

Dobzhansky-Muller model

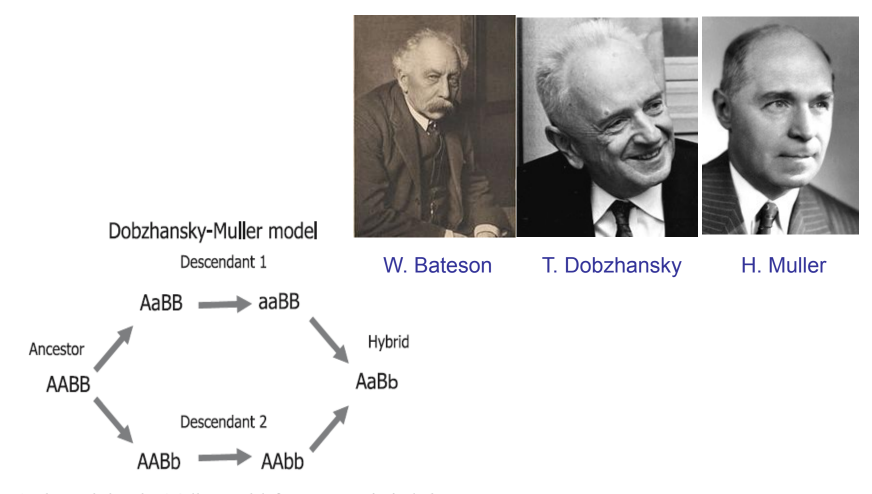


Fig. 1 The Dobzhansky-Muller model for postzygotic isolation

"Arms races" and secondary contact



genetic conflict: "classical" scenario



incompatible!



subpop. 1



arms race in ancestral population

continuing arms race

subpop. 2

secondary contact



MAD = mutually assured destruction

"speciation genes"

genetic conflict: alternative scenario



arms race in ancestral population



subpop. 1

continuing arms race

subpop. 2



Never bring a knife to a gunfight!

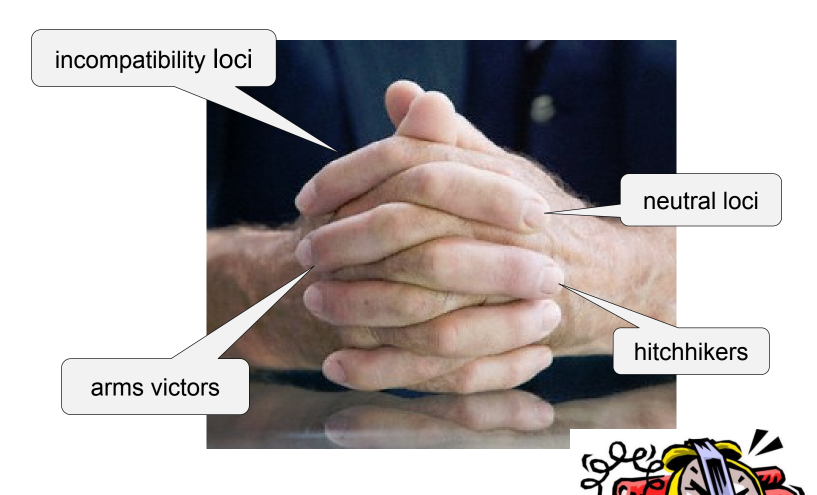


secondary contact

the winner thrives on "naive" genetic background

"antispeciation genes"

Why we don't see this more often?



Ticking time-bomb...

Cytonuclear disequilibria

- = non/random associations of nuclear and cytoplasmic (mitochondrial) alleles
- 3 × 2 table

| | nuclear genotype: | | | |
|--------|-------------------|-----------------------|-------|-------|
| mtDNA: | AA | Aa | aa | total |
| М | U_1 | V ₁ | W_1 | x |
| m | $u_2^{}$ | V_2 | W_2 | У |
| total | u | V | W | 1 |

No hybridization

| | nuclear genotype: | | |
|--------|-------------------|----|-----|
| mtDNA: | AA | Aa | aa |
| М | +++ | 0 | 0 |
| m | 0 | 0 | +++ |

Random mating, hybrid swarm

| | nuclear genotype: | | |
|--------|-------------------|---------|---------|
| mtDNA: | AA | Aa | aa |
| М | obs=exp | obs=exp | obs=exp |
| m | obs=exp | obs=exp | obs=exp |

Hybridization without apparent introgression, crossing independent of sex

| | nuclear genotype: | | |
|--------|-------------------|---------|----|
| mtDNA: | AA | Aa | aa |
| М | ++ | obs=exp | 0 |
| m | 0 | obs=exp | ++ |

Hybridization without apparent introgression, crossing depends on sex

| | nuclear genotype: | | |
|--------|-------------------|----|----|
| mtDNA: | AA | Aa | aa |
| М | ++ | ++ | 0 |
| m | 0 | | ++ |

Hybrids mate more often with less discriminating species

| | nuclear genotype: | | |
|--------|-------------------|----|----|
| mtDNA: | AA | Aa | aa |
| М | obs=exp | ++ | |
| m | obs=exp | | ++ |

Potential introgression, crossing dependent on sex

| | nuclear genotype: | | |
|--------|-------------------|----|----|
| mtDNA: | AA | Aa | aa |
| М | ++ | ++ | |
| m | 0 | 0 | ++ |

Symmetrical introgression

| | nuclear genotype: | | |
|--------|-------------------|---------|----|
| mtDNA: | AA | Aa | aa |
| М | ++ | obs=exp | |
| m | | obs=exp | ++ |