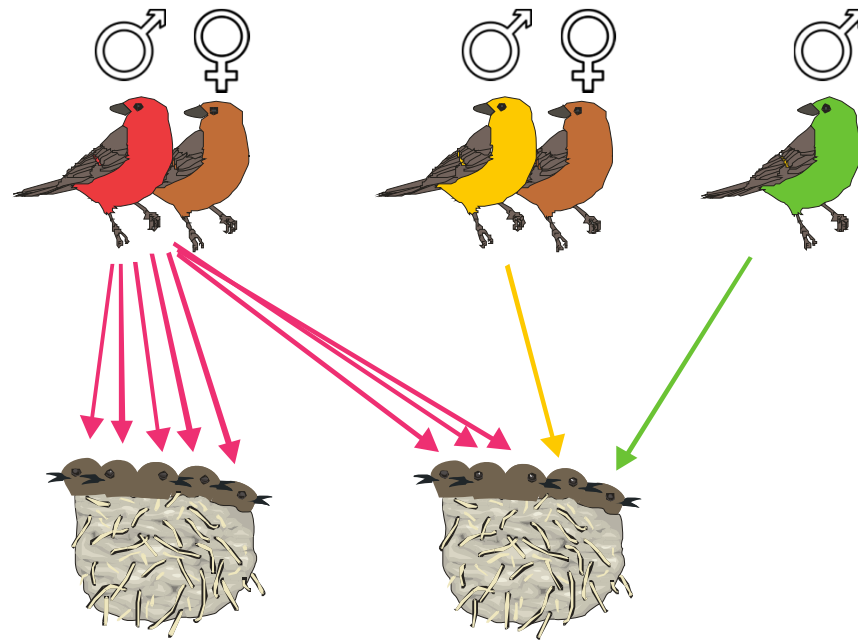


# Molekulární identifikace

## Druh, jedinec, pohlaví



# Identifikace druhů

DNA barcoding

# Definice druhu

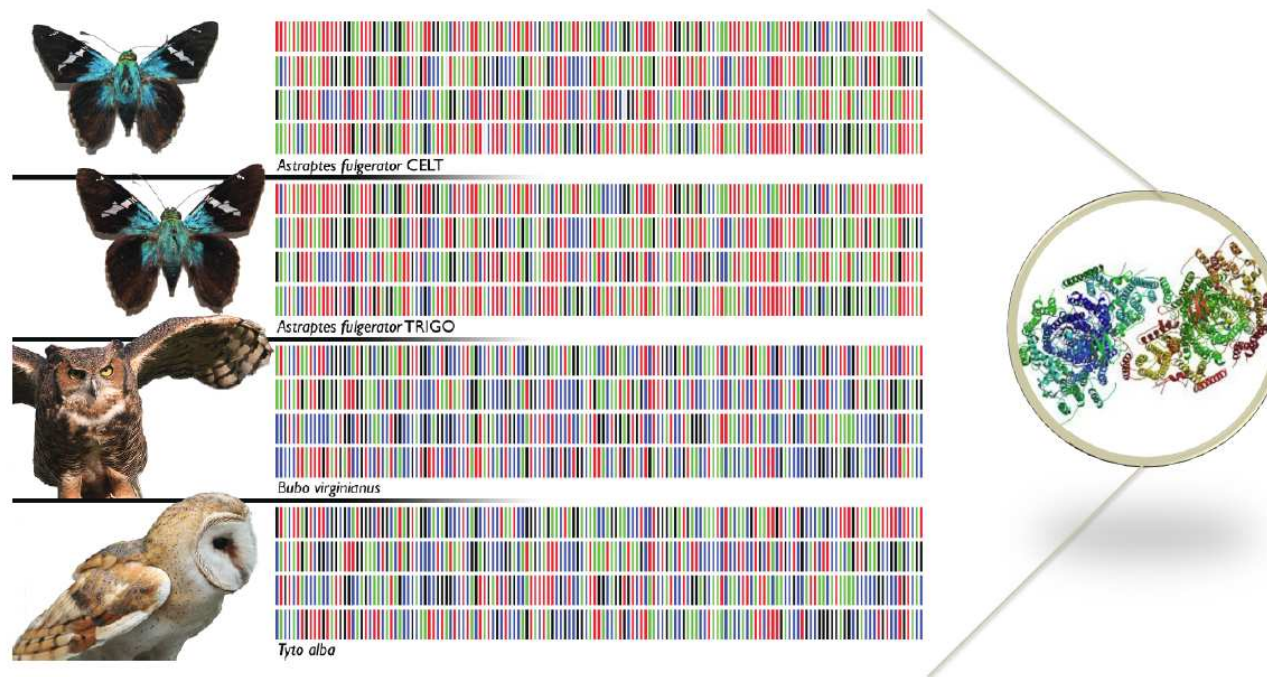
- Koncept **biologického druhu** (Mayr, 1942) – RIM = post- nebo prezygotické bariéry toku genů (ne vždy tak jednoduché – hybridní zóny, alopatická speciace, asexuální druhy atd.)
- Další koncepty (např. typologický = morfologický, genetický, fylogenetický aj.)

## Proč je vůbec potřeba druhu určovat (geneticky)?

- Ochranná biologie – je nutno rozhodnout o taxonomické jednotce (druhu), která vyžaduje pozornost
- Forezní genetika, vývojová stadia bez determinačních znaků, identifikace kořisti v trávicím traktu predátorů, atd.

Barcoding is a standardized approach to identifying plants and animals by minimal sequences of DNA, called DNA barcodes

**DNA Barcode:** A short DNA sequence, from a uniform locality on the genome, used for identifying species



# Biological identifications through DNA barcodes

Paul D. N. Hebert\*, Alina Cywinska, Shelley L. Ball  
and Jeremy R. deWaard

Department of Zoology, University of Guelph, Guelph, Ontario N1G 2W1, Canada

CBOL in 2005

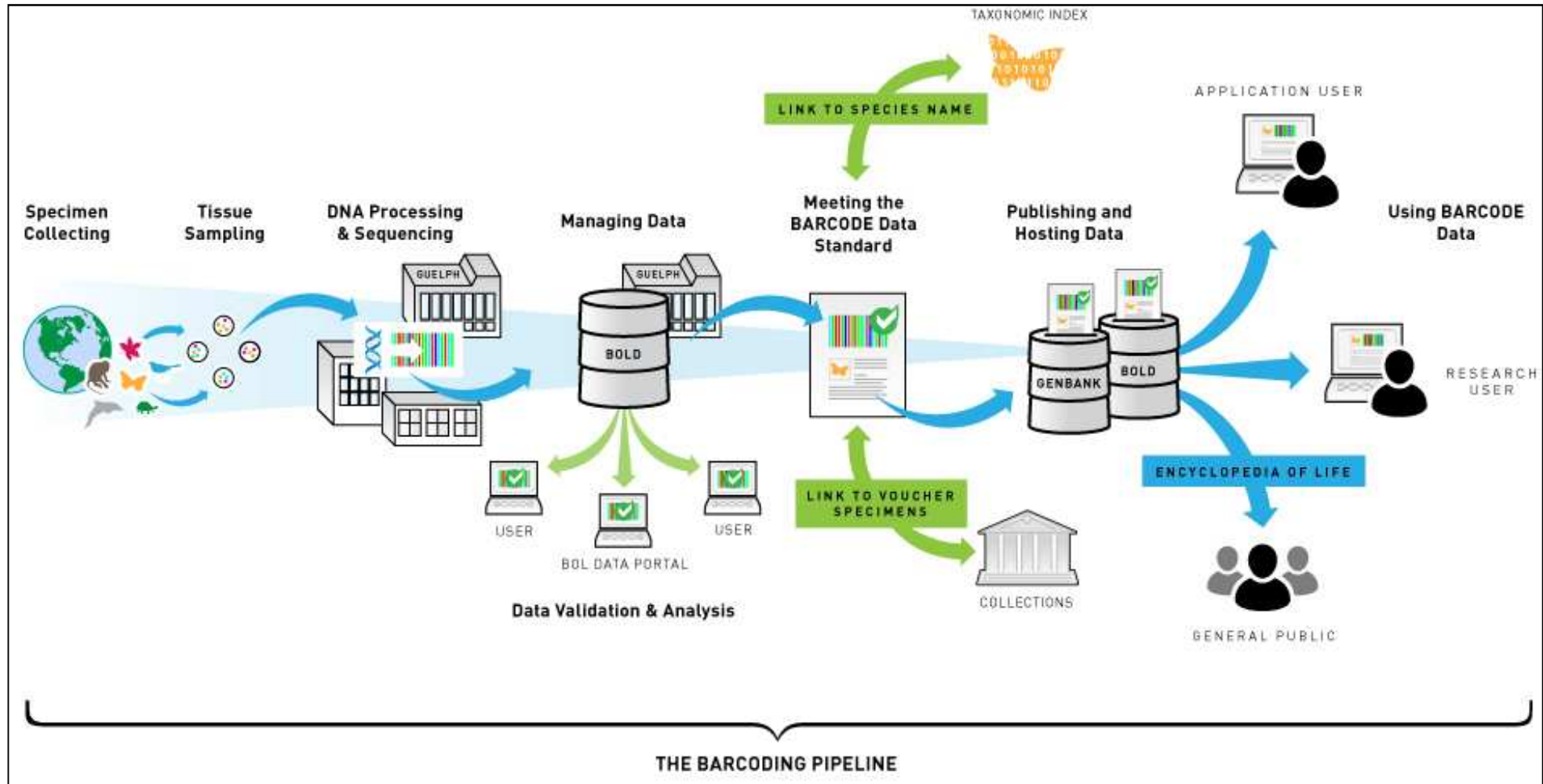
international  
BARCODE  
OF LIFE



iBOL 2010-2015

500 000 species  
barcoded in 2015

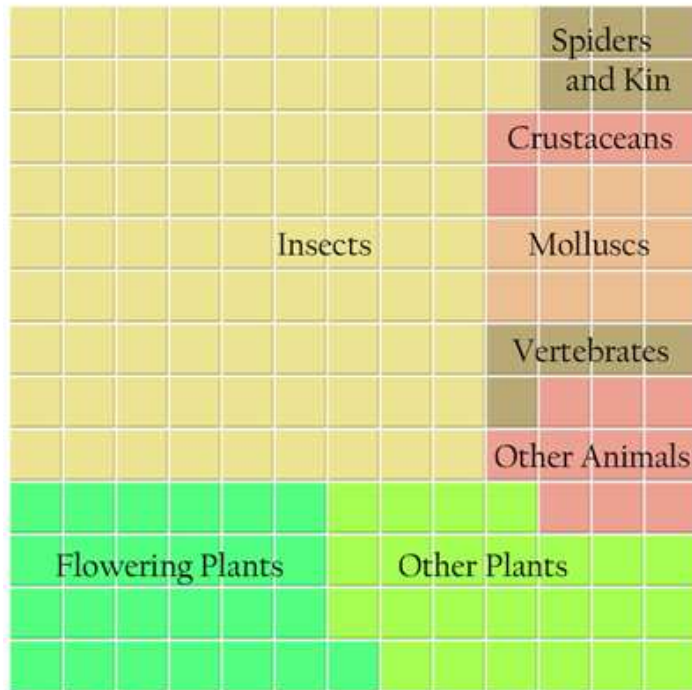
first idea in 2003





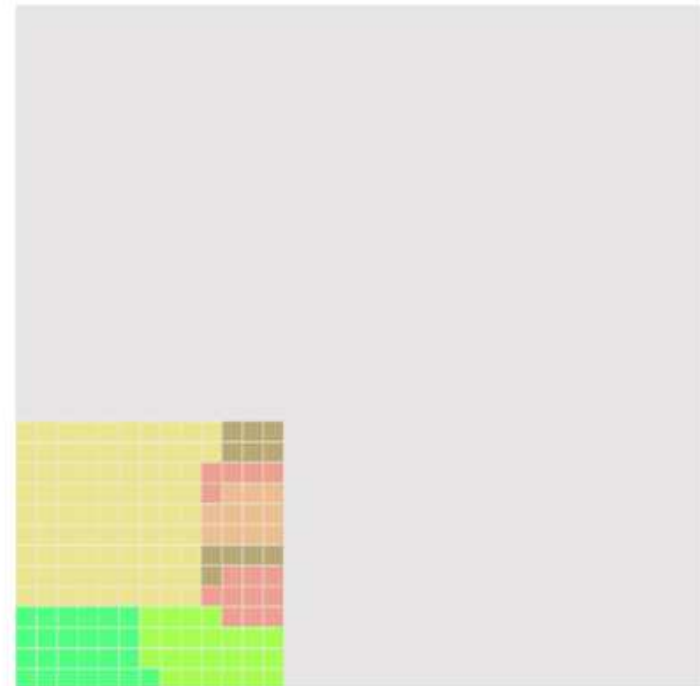
# Why barcode animal and plant species?

Known Biodiversity  
*Approximately 1.7 million named species of plants and animals.*



1 square = 10,000 species

Estimated Biodiversity  
*10 million species*



# Crisis of biodiversity and classical taxonomy



DNA barcoding is  
important part of  
„integrative taxonomy“

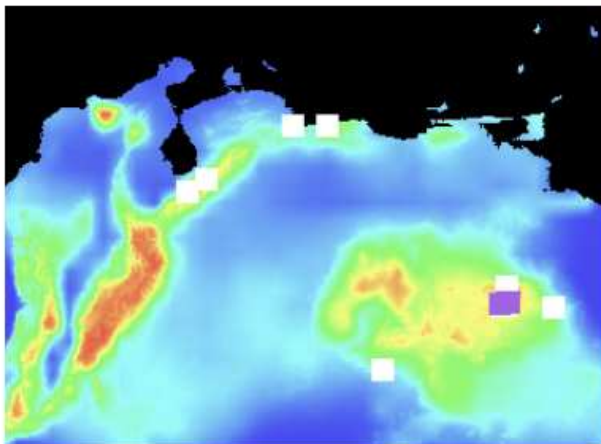


# Integrative taxonomy

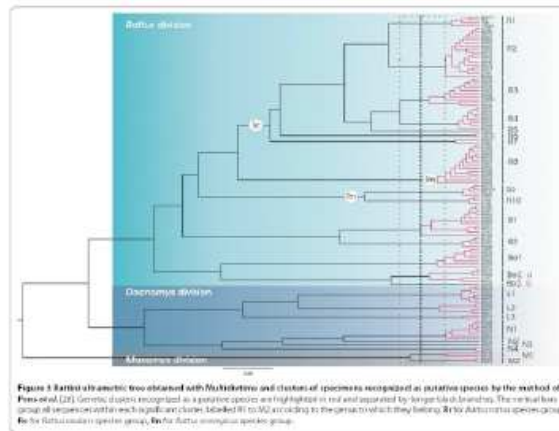
Integr. Taxonomy



Ecology



Genetics

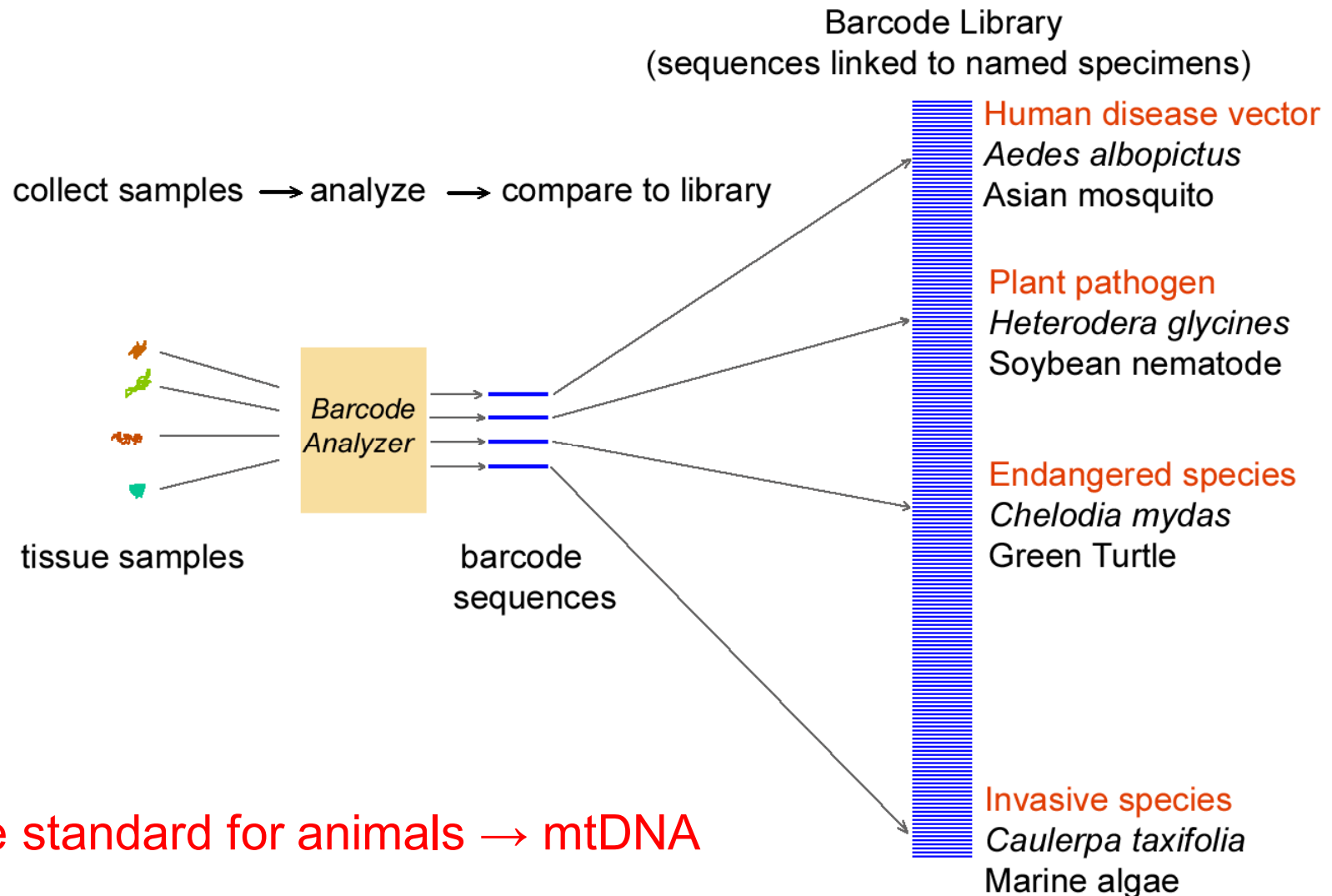


Behavioural Biol.



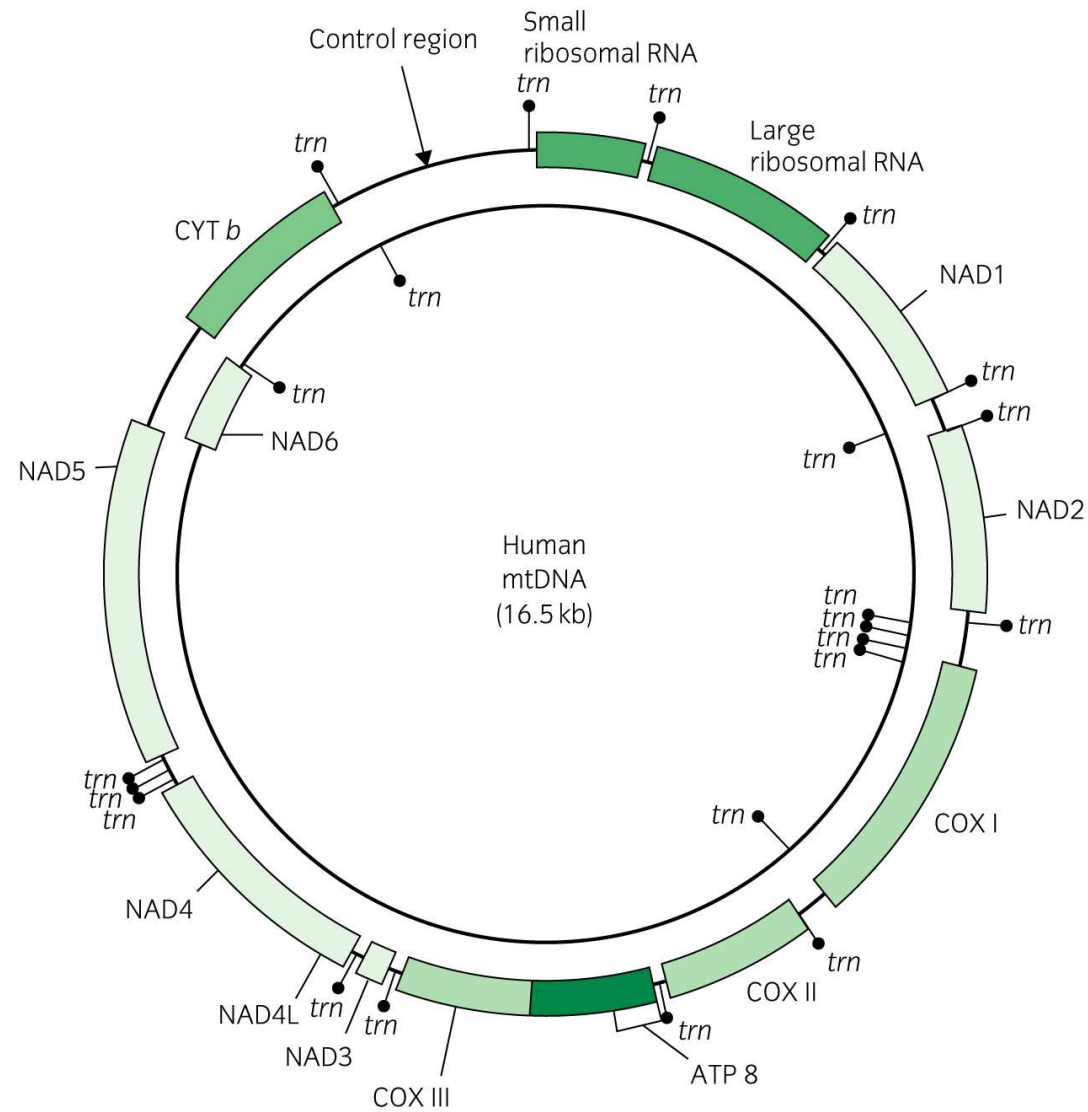


# What are the benefits of standardization?

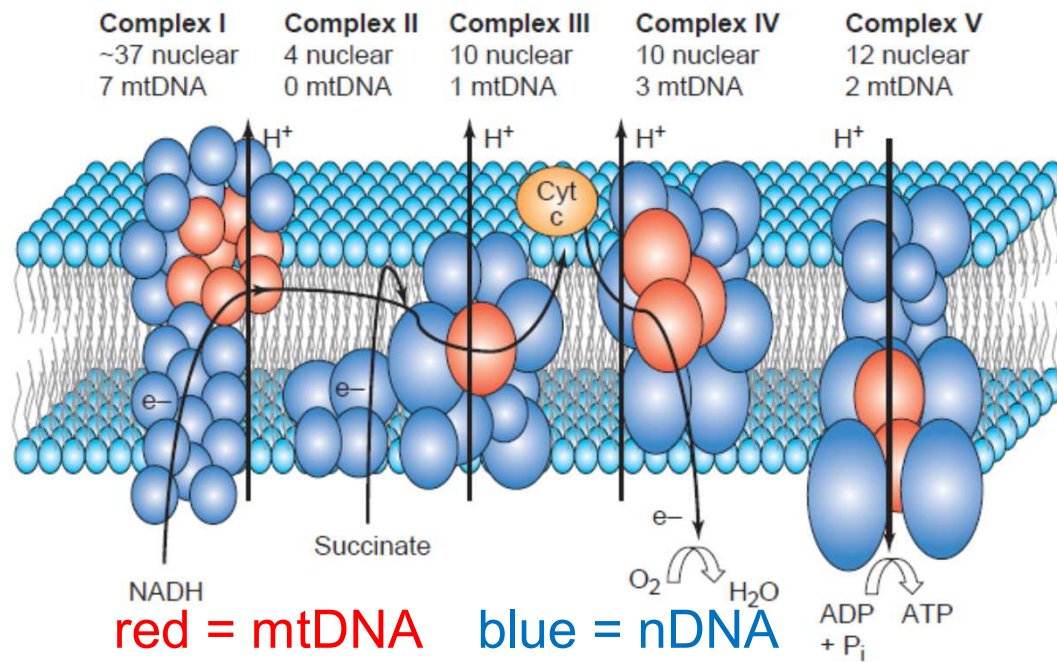


Suitable standard for animals → mtDNA

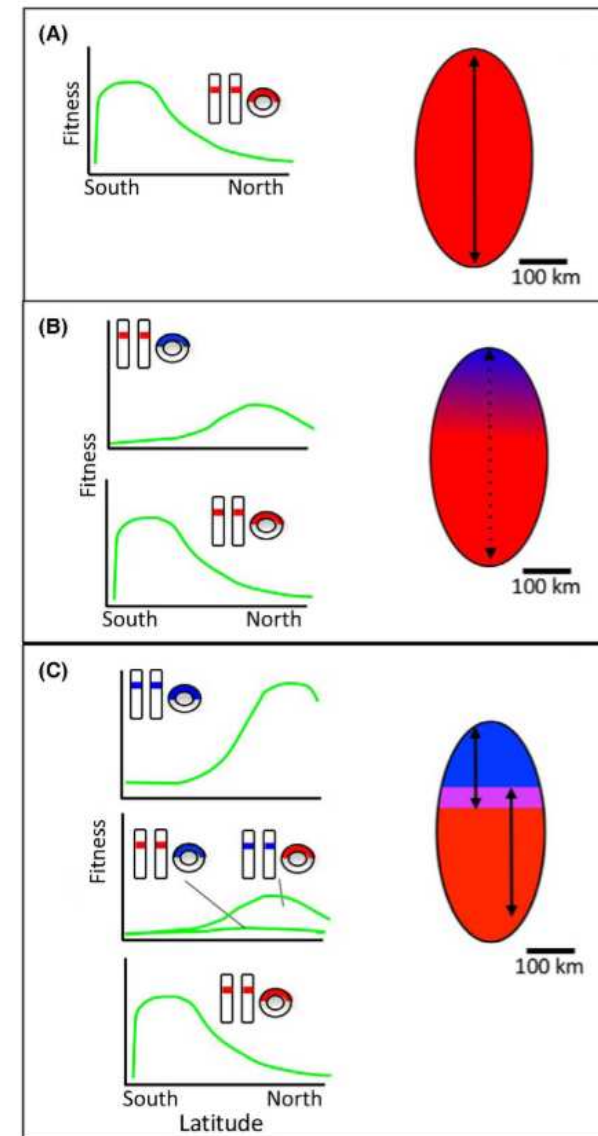
# Why barcode animals with mitochondrial DNA?



# Mitonuclear coevolution as the genesis of speciation (?)



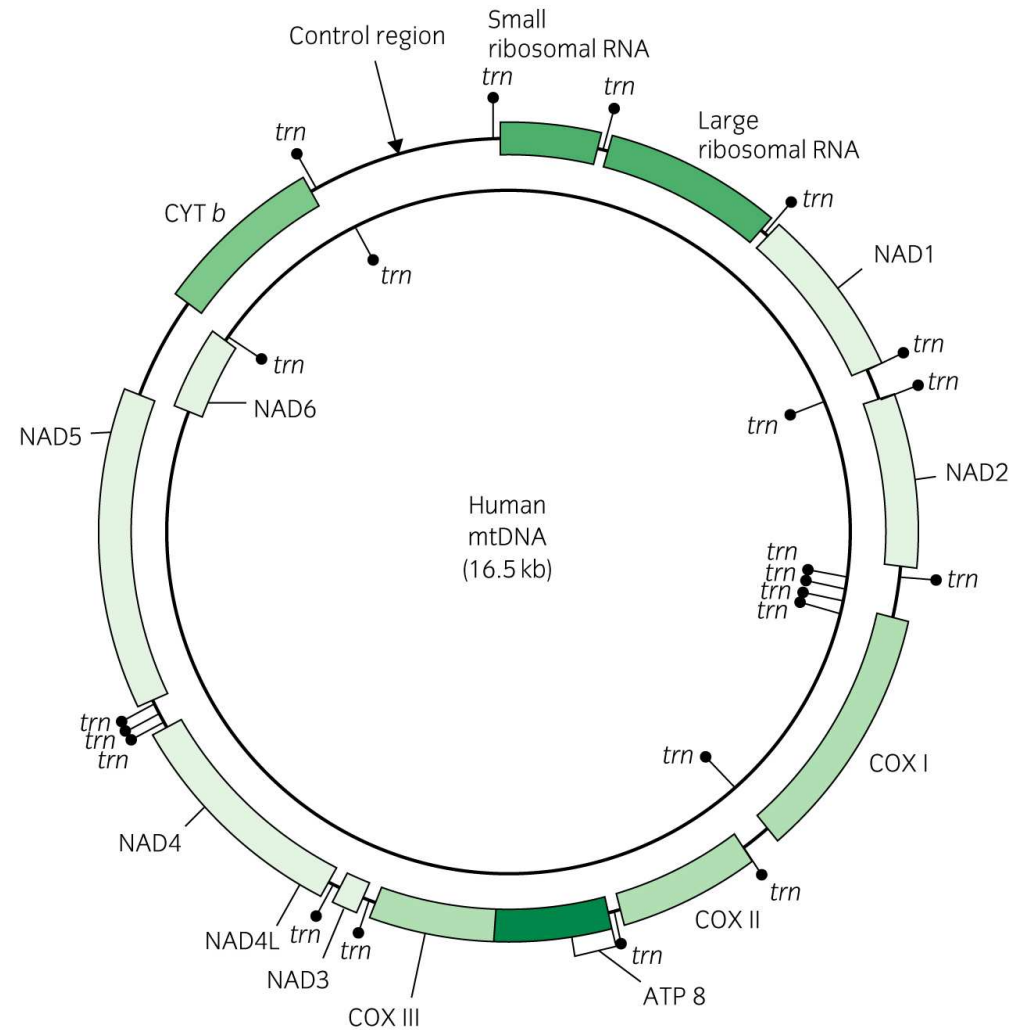
Rand et al. 2004



Hill 2016

# Why barcode animals with mitochondrial DNA?

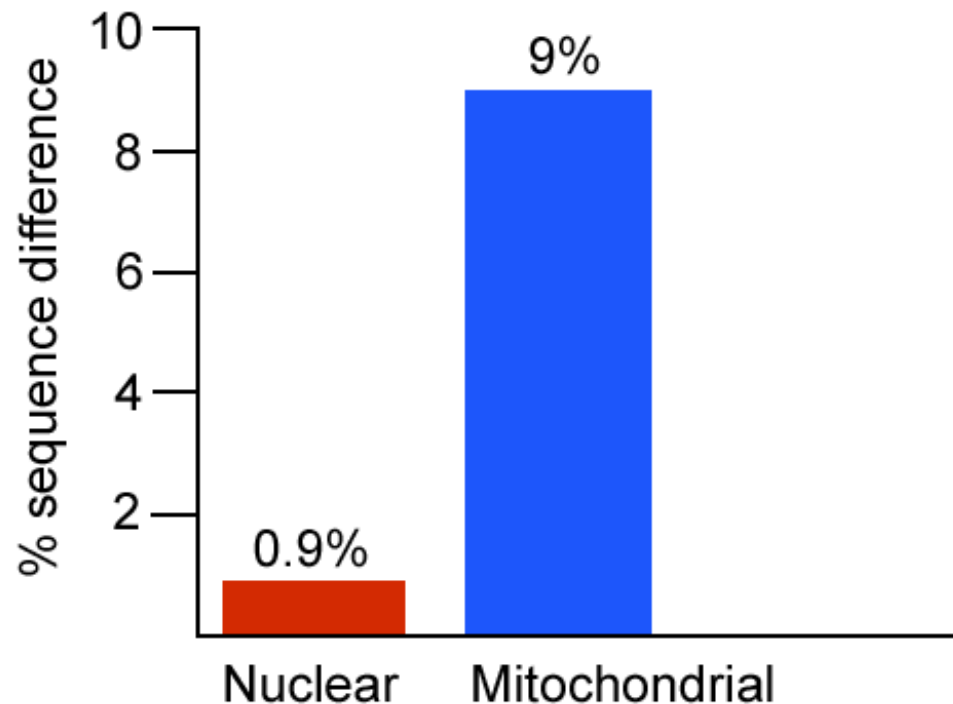
Four properties make mitochondrial genomes especially suitable for identifying species





1. **Greater differences among species**, on average 5- to 10-fold higher in mitochondrial than in nuclear genes. Thus shorter segments distinguish among species, and because shorter, less expensively.

Average sequence differences in nuclear and mitochondrial DNA between human and chimp



- **2. Copy number** There are 100-10,000 more copies of mitochondrial than nuclear DNA per cell, making recovery, especially from small or partially degraded samples, easier and cheaper.
- **3. Relatively few differences within species** in most cases. Small intraspecific and large interspecific differences signal distinct genetic boundaries between most species, enabling precise identification with a barcode.
- **4. Introns, which are non-coding regions interspersed between coding regions of a gene, are absent from mitochondrial DNA** of most animal species, making amplification straightforward. Nuclear genes are often interrupted by introns, making amplification difficult or unpredictable.

# Focus to date

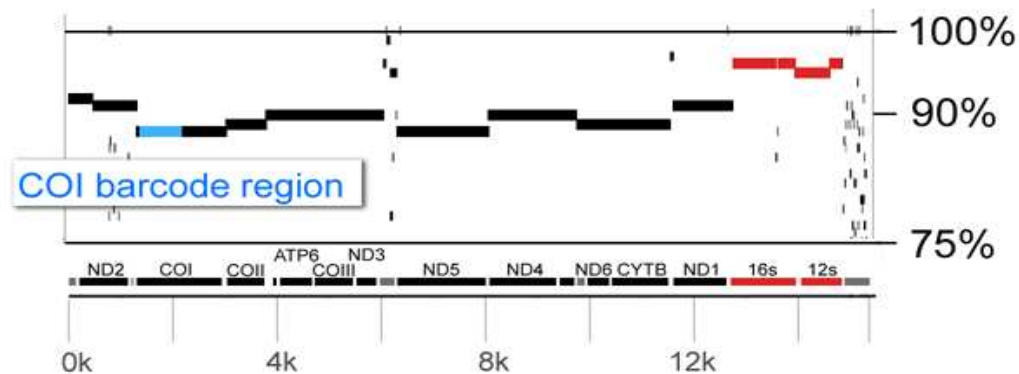
- **For animals**, a 658 base-pair fragment of the mitochondrial gene, **cytochrome oxidase subunit I** (mtCOI) – consensus for iBOL consortium; 18S rDNA – Nematoda etc.
- **For plants**, mitochondrial genes do not differ sufficiently to distinguish among closely related species. Promising markers are genes on cpDNA: matK and rbcL
- **For bacteria**, a 16S-rDNA emerges as very useful marker (especially when using next-generation sequencing)
- For particular taxonomic groups, also other barcodes are widely used, e.g. cytochrome *b* for mammals

# Cytochrome c oxidase I (COI or CoxI) contains differences representative of those in other protein-coding genes

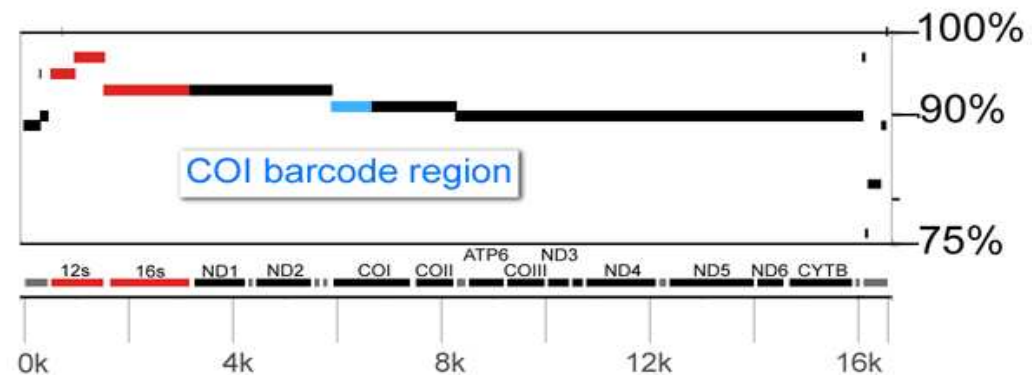
Possible gains in accuracy or cost using a different protein-coding gene would likely be small.

Percent identity plot (PIP) analysis of complete mitochondrial genomes

*Anopheles gambiae*  
vs.  
*A. quadrimaculatus*



*Homo sapiens*  
vs.  
*Pan troglodytes*

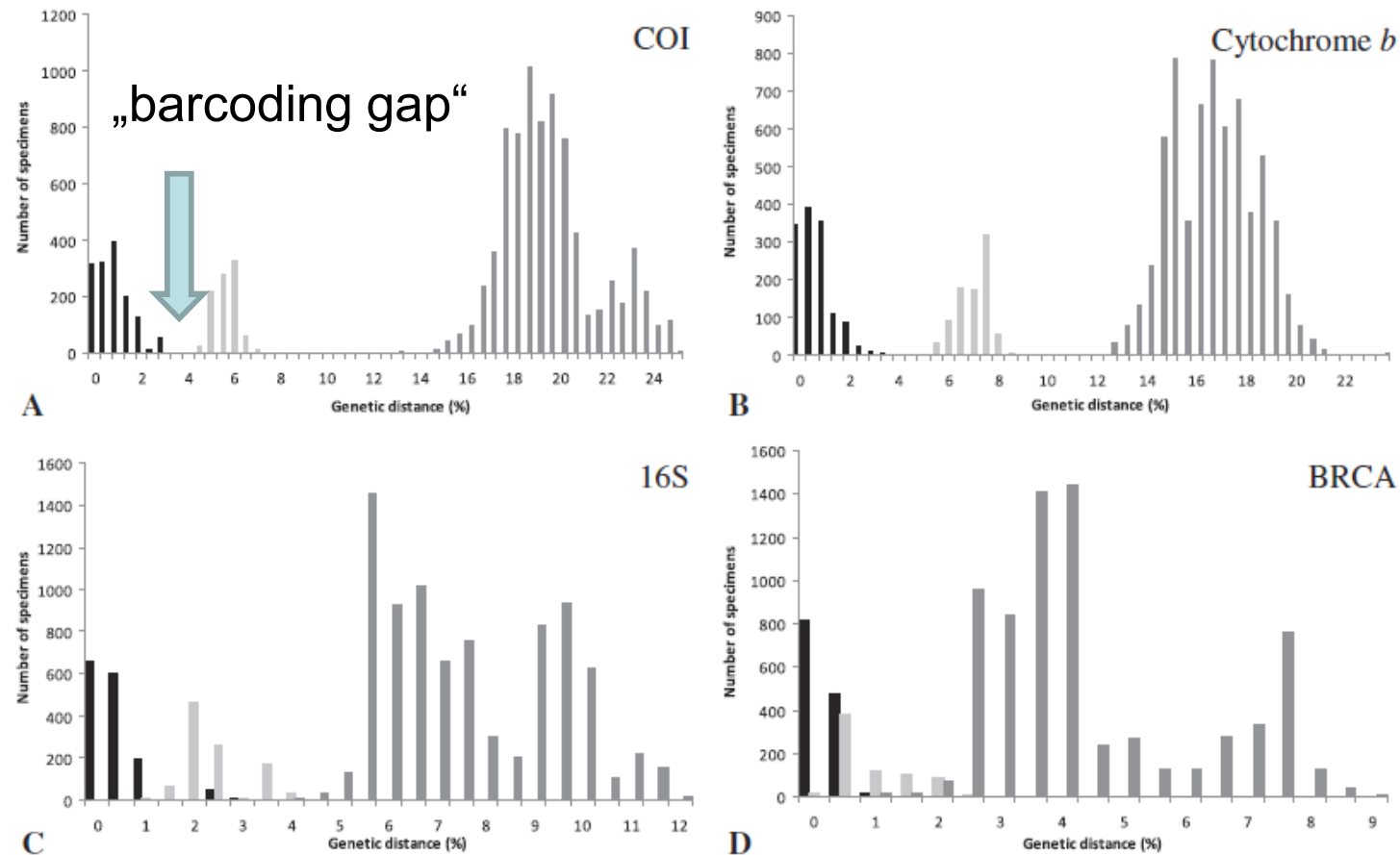


— protein coding gene  
— ribosomal RNA gene



# Barcoding, molecular taxonomy, and exploration of the diversity of shrews (Soricomorpha: Soricidae) on Mount Nimba (Guinea)

FRANÇOIS JACQUET<sup>1\*</sup>, VIOLAINE NICOLAS<sup>1</sup>, CELINE BONILLO<sup>2</sup>,  
CORINNE CRUAUD<sup>3</sup> and CHRISTIANE DENYS<sup>1</sup>



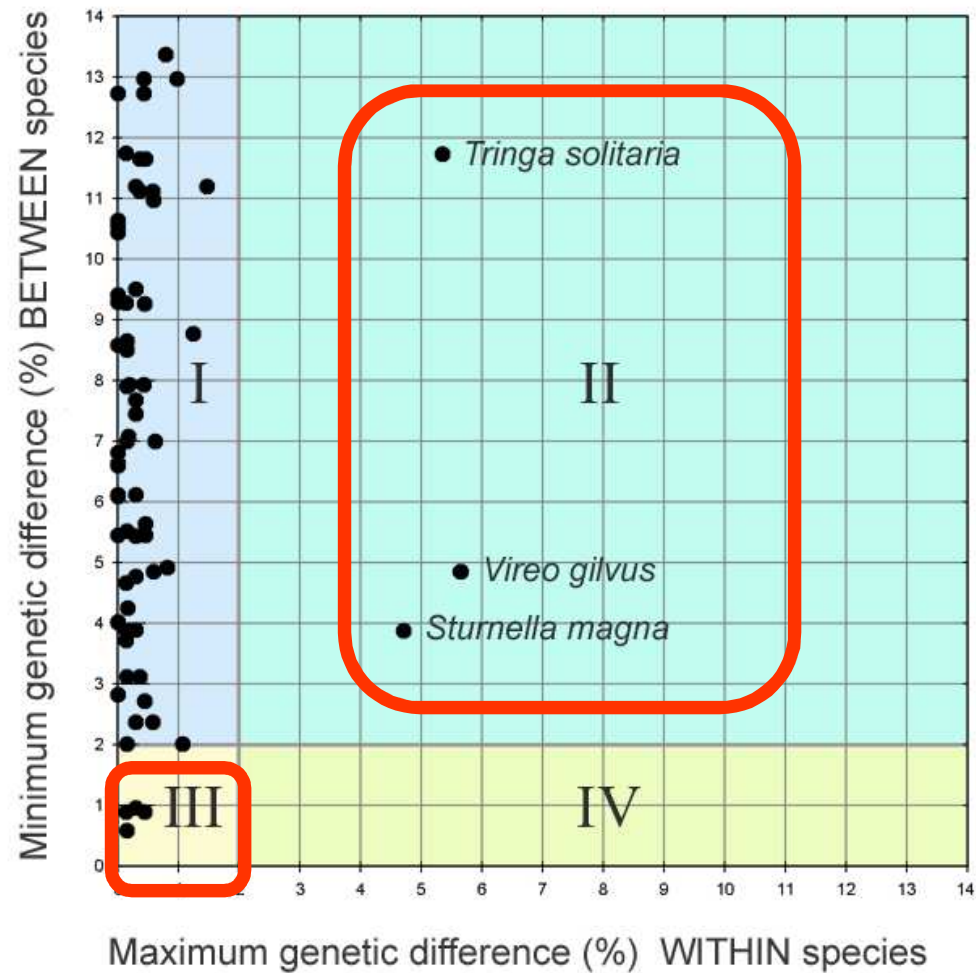
**Figure 3.** Histograms of divergence percentage values between all sequences. Black, intraspecific distances; light grey, interspecific distances amongst clade C; dark grey, interspecific distances excluding clade C. All sequences were corrected using the Kimura two-parameter substitution model. Sequences corresponding to nuclear copies of cytochrome *b* were excluded from the analysis. BRCA, breast cancer gene; COI, cytochrome oxidase I.

## What do barcode differences among and within animal species studied so far suggest?

- barcodes identify most animal species unambiguously
- approximately 2-5% of recognized species have shared barcodes with closely-related species - many of them hybridize regularly
- in all groups studied so far, distinct barcode clusters with biologic co-variation suggest cryptic species

# Barcoding North American birds

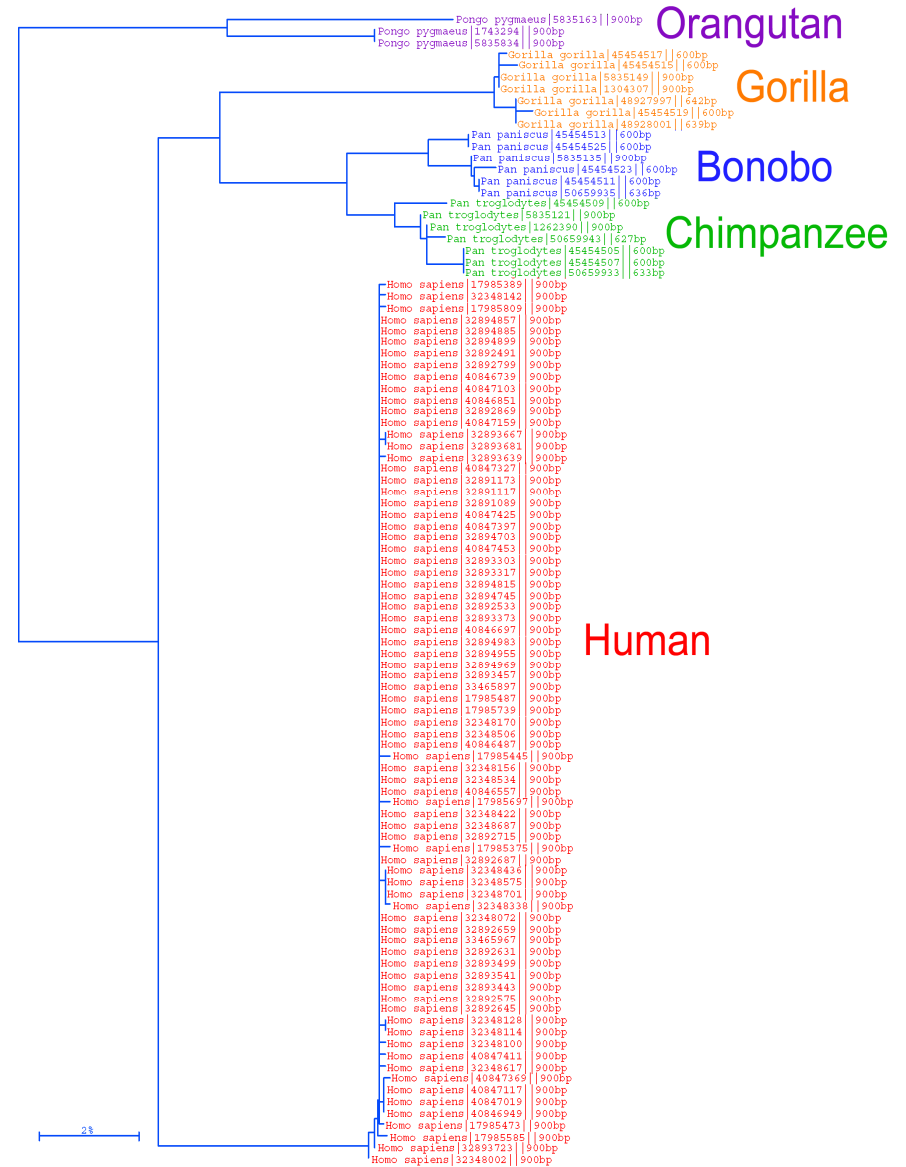
Interspecific vs. intraspecific COI barcode differences



Tak co třeba znamená toto?

Barcodes affirm the unity of the species *Homo sapiens*

Comparisons show we differ from one another by only 1 or 2 nucleotides out of 648, while we differ from chimpanzees at 60 locations and gorillas at 70 locations.





## A barcoder?



Mark Stoeckle The Rockefeller University  
Paul Waggoner Connecticut Agricultural Experiment Station  
Jesse Ausubel Alfred P. Sloan Foundation

# Next generation sequencing of amplicons

## 1. Metagenomika/metatranskriptomika

- celé společenstvo půdních, vodních mikroorganismů, střevní mikroflóra
- PCR genu 16S (18S) rRNA
- lze i kvantifikovat

## 2. Složení potravy (COI barcoding)

- COI barcoding (masožravci)
- cpDNA (býložravci)

## 3. Analýza kontaminovaných vzorků

OPEN ACCESS Freely available online

PLOS ONE

### Next-Generation Sequencing for Rodent Barcoding: Species Identification from Fresh, Degraded and Environmental Samples

Maxime Galan<sup>1\*</sup>, Marie Pagès<sup>1,2</sup>, Jean-François Cosson<sup>1</sup>

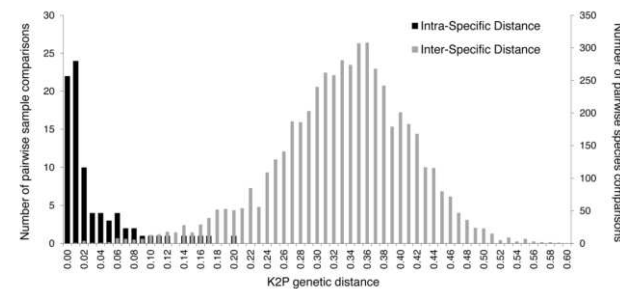


Figure 2. Distribution of pairwise K2P genetic distances within and between 103 rodent species (265 individuals) based on the 136 bp mini-barcode *cytb*.

|SE|S|AM|E| BARCODE

SEquences Seeker & AMPlicons Explorer for Barcoding

Genotyping and barcoding based on high-throughput multiplex amplicon sequencing

## Illumina sequencers

### Illumina MiSeq

4 millions reads/run  
150 bp/read



### Illumina GAIIx

300 millions reads/run  
150 bp/read



### Illumina HighSeq

1500 – 3000 millions reads/run  
100 bp/read



## A barcoder? ... COMING SOON



Mark Stoeckle The Rockefeller University  
Paul Waggoner Connecticut Agricultural Experiment Station  
Jesse Ausubel Alfred P. Sloan Foundation

## What *isn't* DNA Barcoding?



- it is not intended to, in any way, supplant or invalidate existing taxonomic practice
- it is not DNA taxonomy; it does not equate species identity, formally or informally, with a particular DNA sequence
- it is not intended to duplicate or compete with efforts to resolve deep phylogeny (e.g., Assembling the Tree of Life, ATOL)



# What are the main limits to barcoding encountered so far?

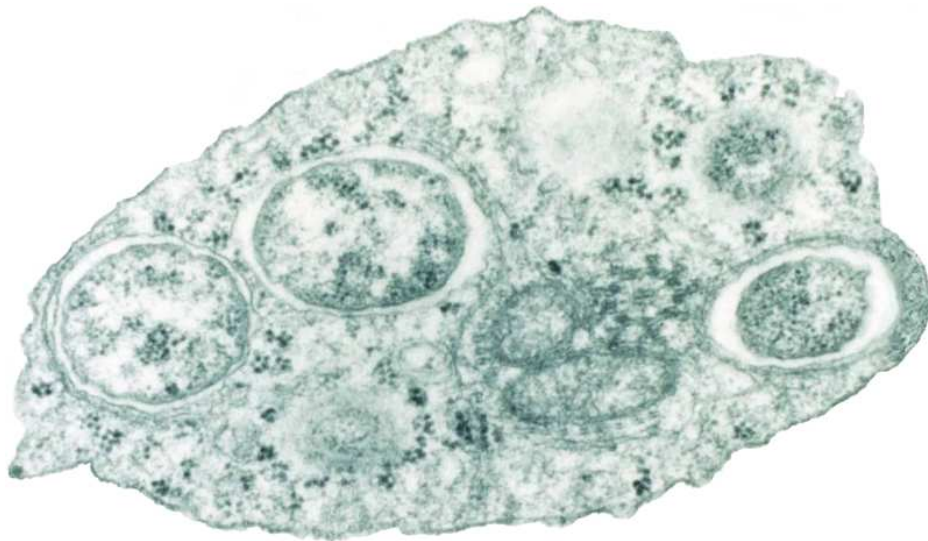
- horizontal gene transfer
- gene tree vs. species tree
- nuclear pseudogenes
- hybrids – mtDNA introgression

# 1. Horizontal gene transfer

## DNA barcoding cannot reliably identify species of the blowfly genus *Protophthora* (Diptera: Calliphoridae)

T.L Whitworth, R.D Dawson, H Magalon and E Baudry

*Proc. R. Soc. B* 2007 **274**, doi: 10.1098/rspb.2007.0062, published 22 July 2007



*Wolbachia* within an insect cell  
(25-70% druhů hmyzu)

# Výsledky jaderné a mitochondriální DNA si neodpovídají

AFLP

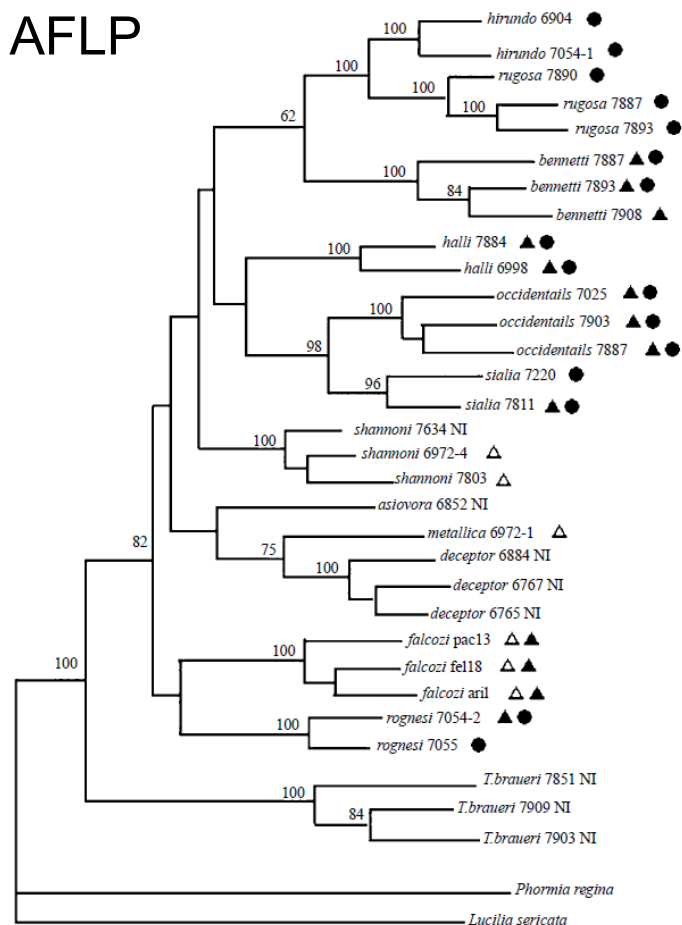
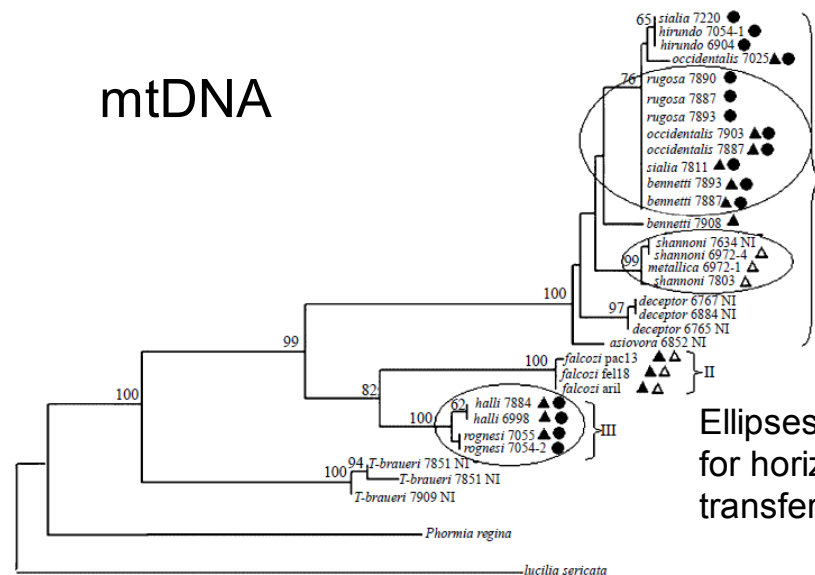


Figure 1. Phylogram of the *Protocalliphora* genus based upon AFLP data. The tree was generated by parsimony analysis using a heuristic search with tree bisection-reconnection. Bootstrap values are shown as percentage of 1000 replicates at each node only if they are 50% or greater. The *Wobachia* infection status of each individual is shown on the tree. Individuals infected with *wA1*, *wA2* or *wB* *Wobachia* strains are respectively represented by an open triangle, a solid triangle and a circle. Non-infected individuals are symbolized by NI.

mtDNA



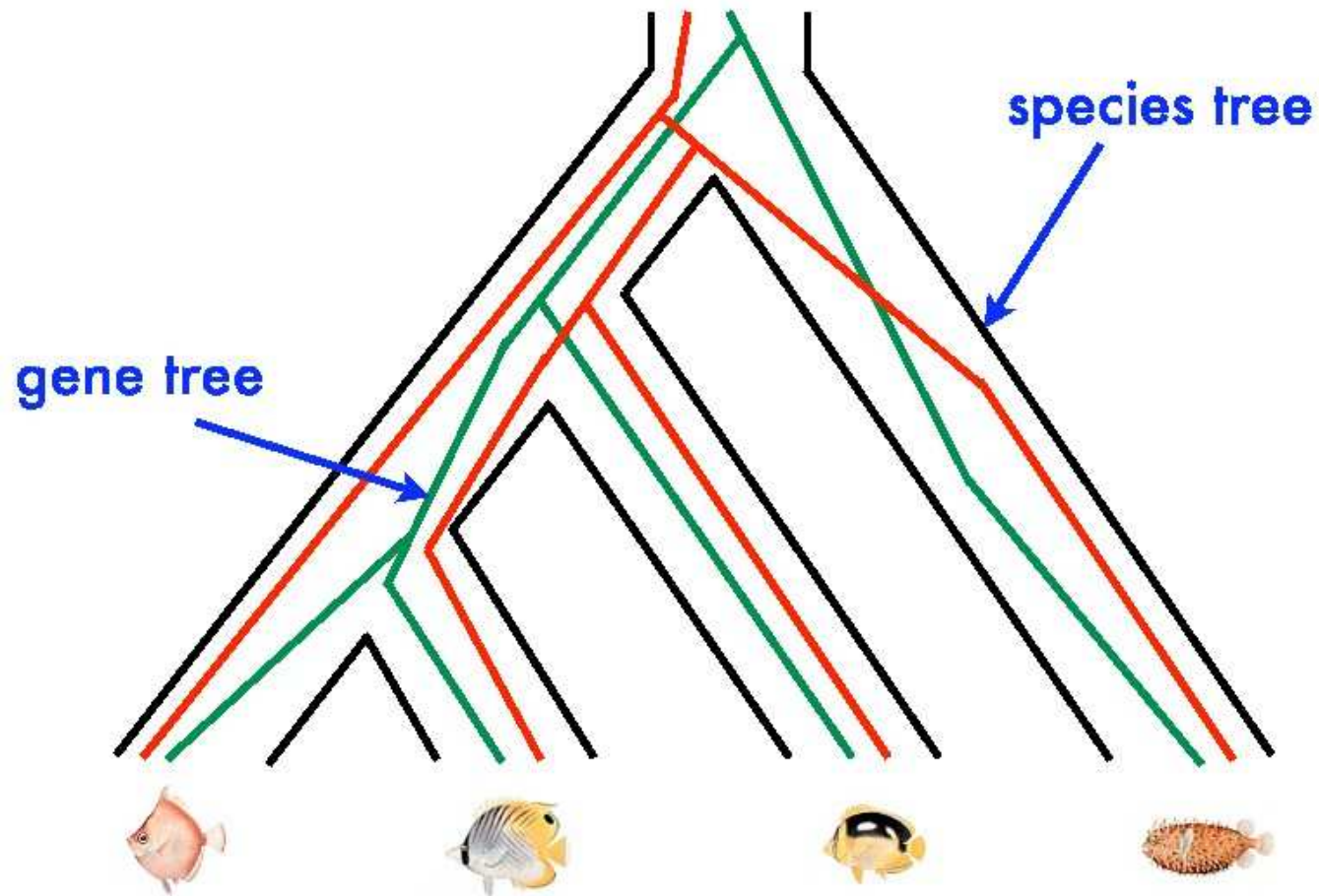
Ellipses = candidates for horizontal gene transfer

Figure 2. Phylogram of the *Protocalliphora* genus based upon COI and COII data (total of 953 bp). The tree was generated by maximum likelihood analysis using a heuristic search with tree bisection-reconnection. Bootstrap values are shown as percentage of 1000 replicates at each node only if they are 50% or greater. The *Wobachia* infection status of each individual is shown on the tree. Individuals infected with *wA1*, *wA2* or *wB* *Wobachia* strains are respectively represented by an open triangle, a solid triangle and a circle. Non-infected individuals are symbolized by NI. Three clusters defined using 3 or 1.8% divergence as threshold values (§3) are shown on the figure. The three ellipses indicate cases where horizontal transfer of *Wobachia* between species seems probable (§3).

Horizontální přenos mtDNA přes *Wobachia* (většinou mezi blízce příbuznými druhy, na úrovni rodů je barcoding OK)

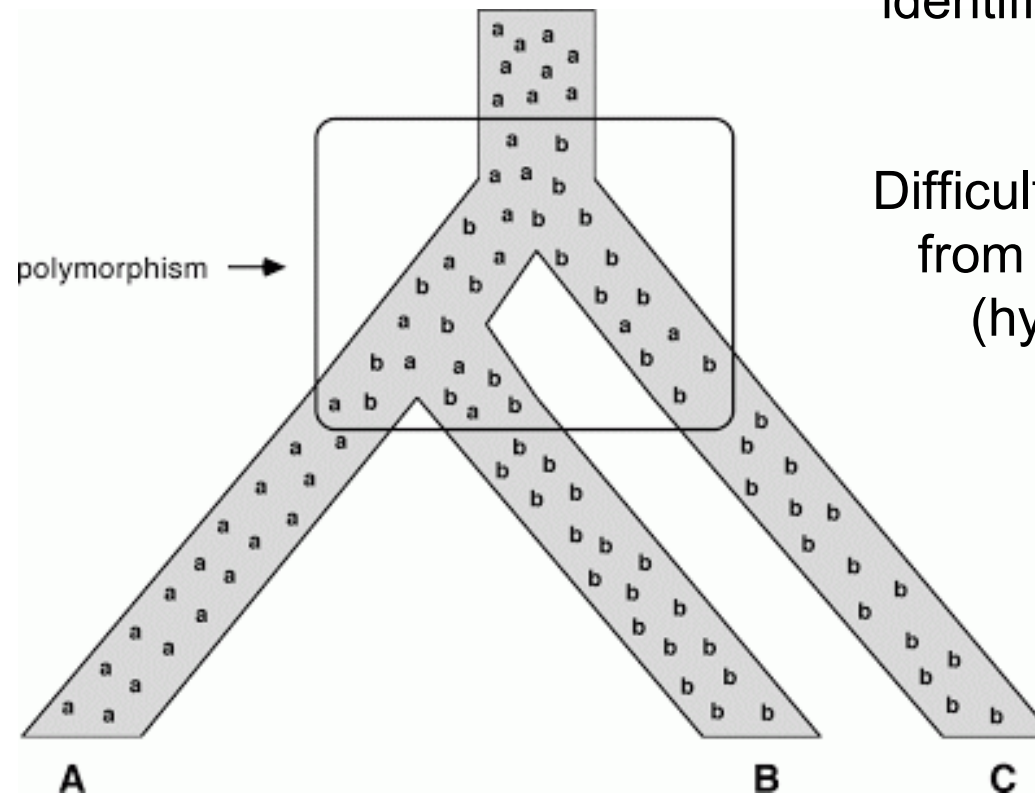
Symboły vpravo odpovídají typu *Wobachia* infekce

## 2. Gene tree vs. species tree



Jeden gen (navíc mitochondriální) není (někdy) dostatečný pro definici druhu

# Incomplete lineage sorting



Problems of species identification in young species

Difficult to distinguish from the gene flow (hybridization)

Time to fix the polymorphisms =  $2 N_e$  generations



# Statistical species delimitation

*Syst. Biol.* 56(3):400–411, 2007  
 Copyright © Society of Systematic Biologists  
 ISSN: 1063-5157 print / 1076-836X online  
 DOI: 10.1080/10635150701405560

## Estimating Species Phylogeny from Gene-Tree Probabilities Despite Incomplete Lineage Sorting: An Example from *Melanoplus* Grasshoppers

BRYAN C. CARSTENS AND L. LACEY KNOWLES

Department of Ecology and Evolutionary Biology, 1109 Geddes Avenue, Museum of Zoology, University of Michigan, Ann Arbor, MI 48109-1079, USA; E-mail: knowlesl@umich.edu (L.L.K.)

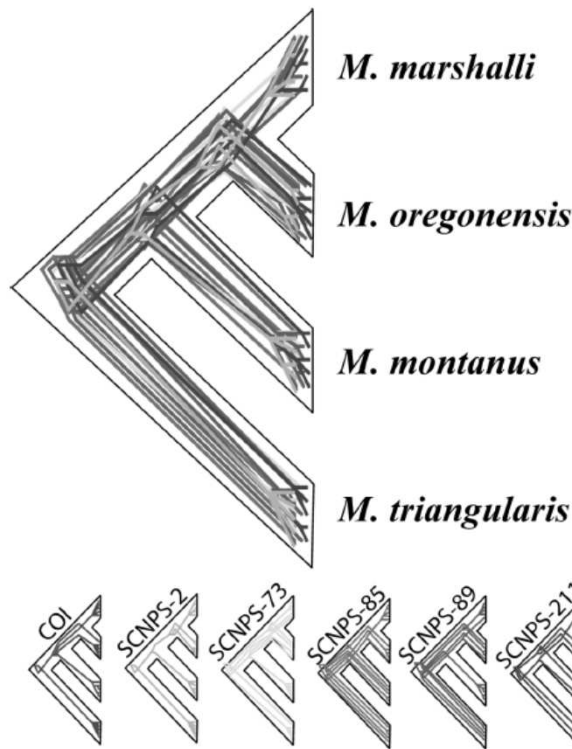


FIGURE 3. Estimate of the species phylogeny (ESP) with the highest likelihood score ( $-\ln L = -10,487.1938$ ) calculated with the coalescent-based approach from the probabilities of the gene trees (see Fig. 2), with each gene tree shown within the species tree.

\*BEAST  
 (species trees)

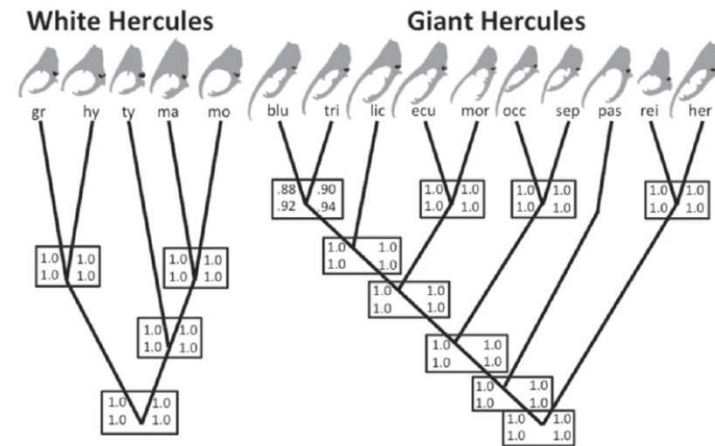


FIGURE 5. Results from iBPP analyses using combined genetic and morphological data sets; note that outlines of major male horn shape shown adjacent to the abbreviated taxonomic names are not drawn to scale (see Fig. 1 for relative body sizes). Support values reported for each node are based on the algorithm setting 0 for the rjMCMC, the algorithm setting 1 for the Brownian motion model, and for four different priors corresponding to large (vs small) ancestral population sizes with relatively deep (vs shallow) divergence times. Specifically, the support values in each box correspond to analyses with the following different priors: upper left,  $\theta = G(1, 10)$  and  $\tau = G(1, 10)$ ; upper right,  $\theta = G(2, 2000)$  and  $\tau = G(2, 2000)$ ; lower left,  $\theta = G(1, 10)$  and  $\tau = G(2, 2000)$ ; lower right,  $\theta = G(2, 2000)$  and  $\tau = G(2, 2000)$ . Taxonomic abbreviations and the corresponding names can be found in Figure 1.

BPP and iBPP

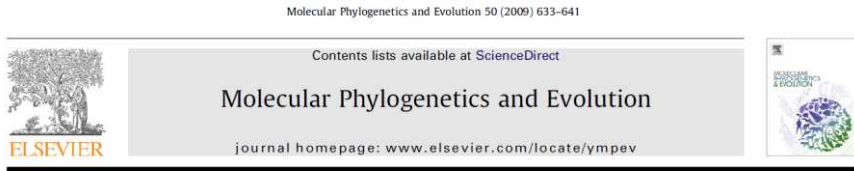
*Syst. Biol.* 0(0):1–15, 2016  
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 For Permissions, please email: journals.permissions@oup.com  
 DOI:10.1093/sysbio/syv119

## The Species versus Subspecies Conundrum: Quantitative Delimitation from Integrating Multiple Data Types within a Single Bayesian Approach in Hercules Beetles

JEN-PAN HUANG\* AND L. LACEY KNOWLES

Department of Ecology and Evolutionary Biology, 1109 Geddes Avenue, Museum of Zoology, University of Michigan, Ann Arbor, MI 48109-1079, USA  
 \*Correspondence to be sent to: Department of Ecology and Evolutionary Biology, 1109 Geddes Avenue, Museum of Zoology, University of Michigan, Ann Arbor, MI 48109-1079, USA; E-mail: huangjp@umich.edu.

# 3. Pseudogenes



False phylogenies on wood mice due to cryptic cytochrome-*b* pseudogene  
 Sylvain Dubey<sup>a\*</sup>, Johan Michaux<sup>b</sup>, Harald Br nner<sup>c</sup>, Rainer Hutterer<sup>d</sup>, Peter Vogel<sup>e</sup>

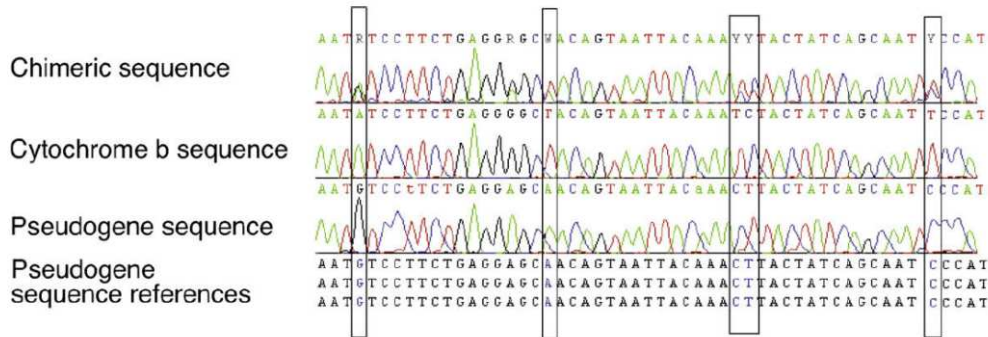
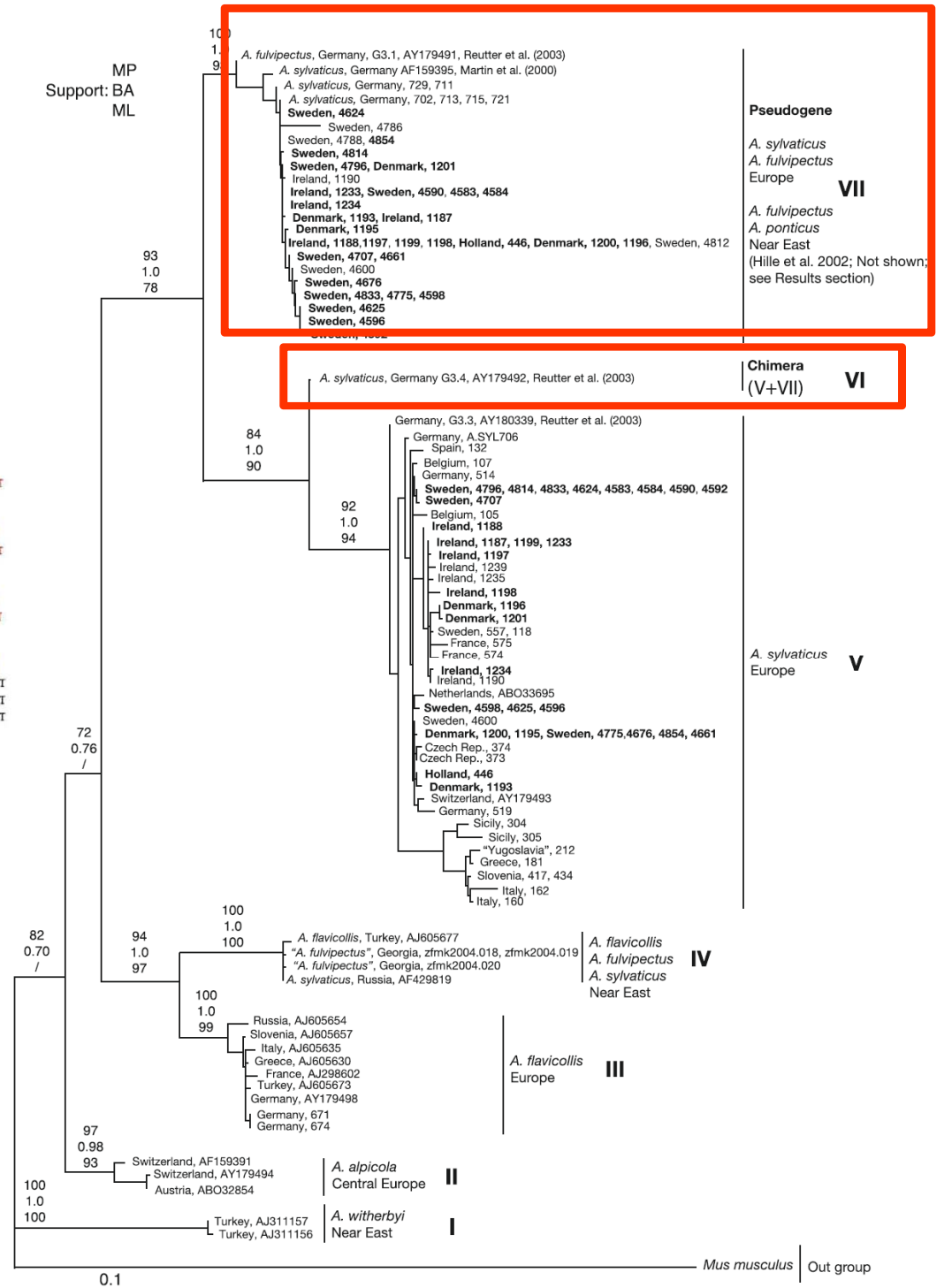


Fig. 2. Examples of electropherograms of a chimeric *cyt-b* sequence of lineage V/VII (VI) and of pure lineages V and VII.

Heterozygotes in mtDNA → be careful!

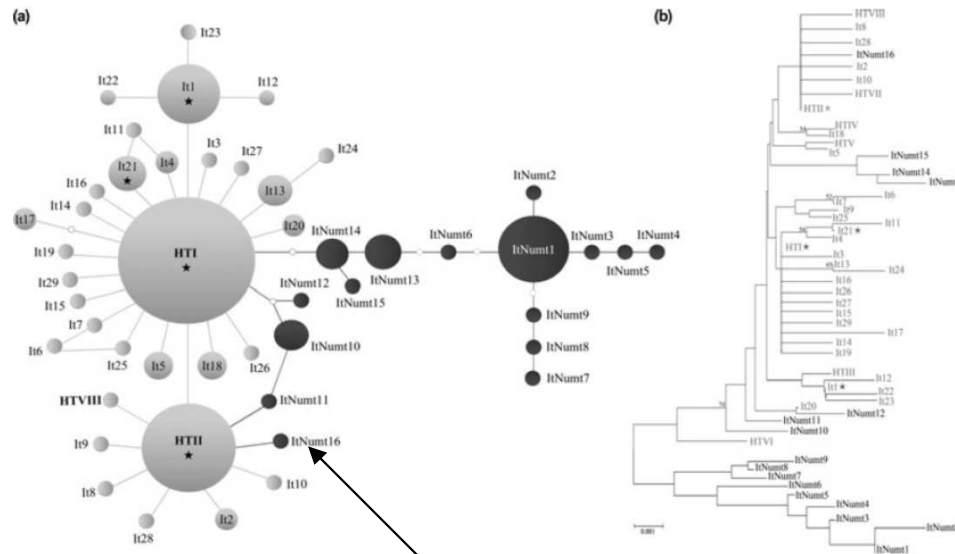
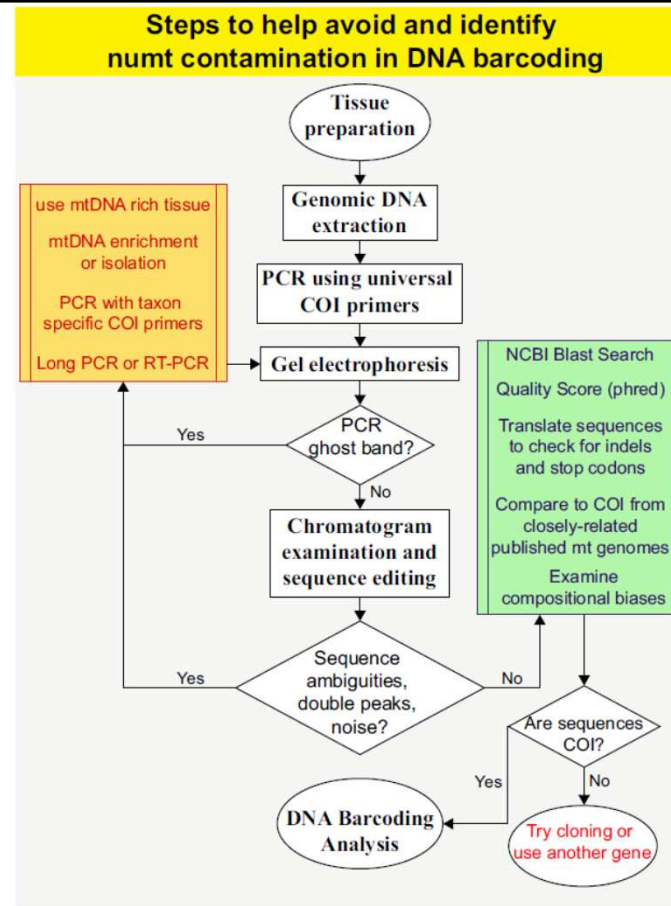
**NUMTS** = „nuclear copy of mtDNA sequences

Relatively often for cytochrome *b*



# How to recognize numt?

- ultracentrifugation (fresh samples required)
- the use of tissues with high proportion of mitochondria (e.g. muscles)
- *long-range PCR* (or sequence complete mtDNA)
- RT-PCR (pseudogenes are not transcribed)
- indels, stop codons
- cloning



cryptic numts



# Many species in one: DNA barcoding overestimates the number of species when nuclear mitochondrial pseudogenes are coamplified

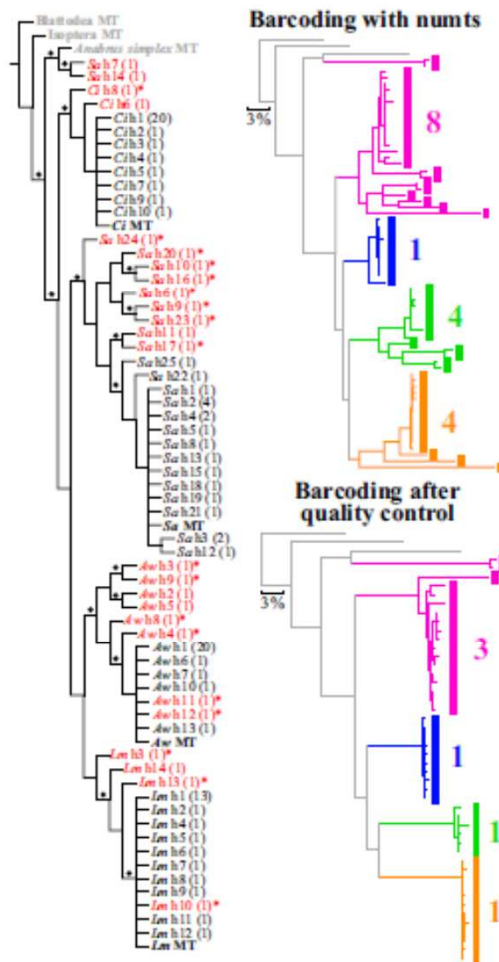
Hojun Song<sup>\*†</sup>, Jennifer E. Buhay<sup>\*‡</sup>, Michael F. Whiting<sup>\*</sup>, and Keith A. Crandall<sup>\*</sup>

<sup>\*</sup>Department of Biology, Brigham Young University, Provo, UT 84602; and <sup>†</sup>Belle W. Baruch Institute for Marine Sciences, University of South Carolina, Columbia, SC 29208

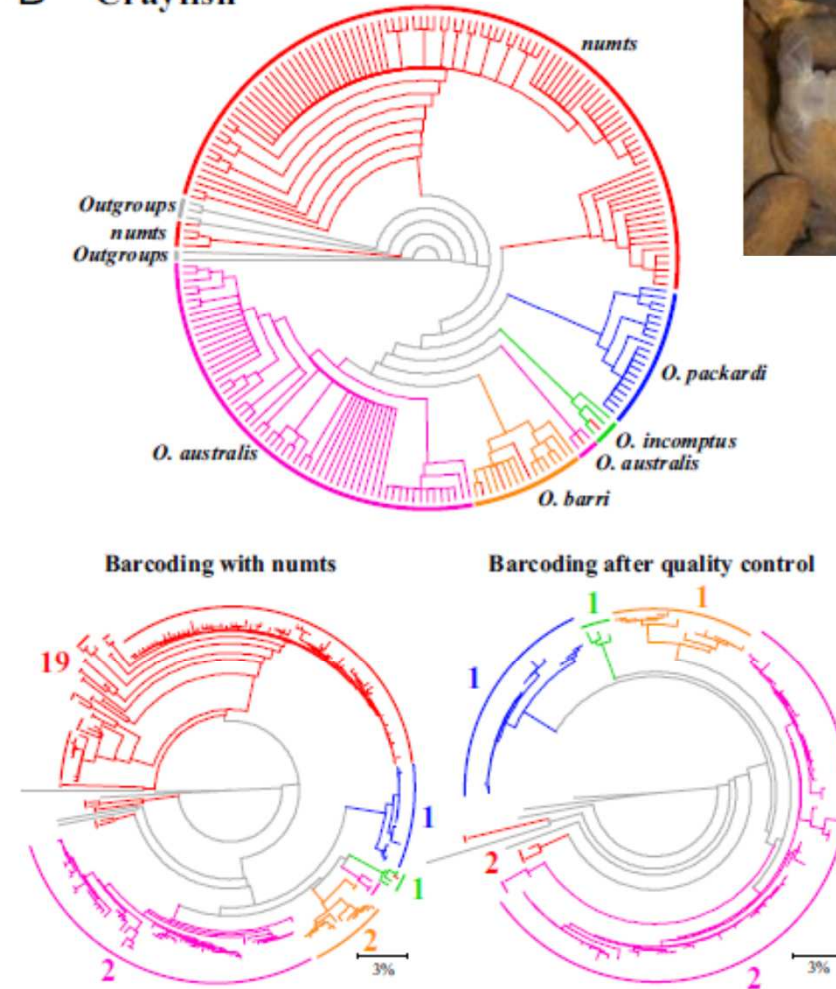
Edited by W. Ford Doolittle, Dalhousie University, Halifax, NS, Canada, and approved July 14, 2008 (received for review March 28, 2008)



## A Grasshoppers

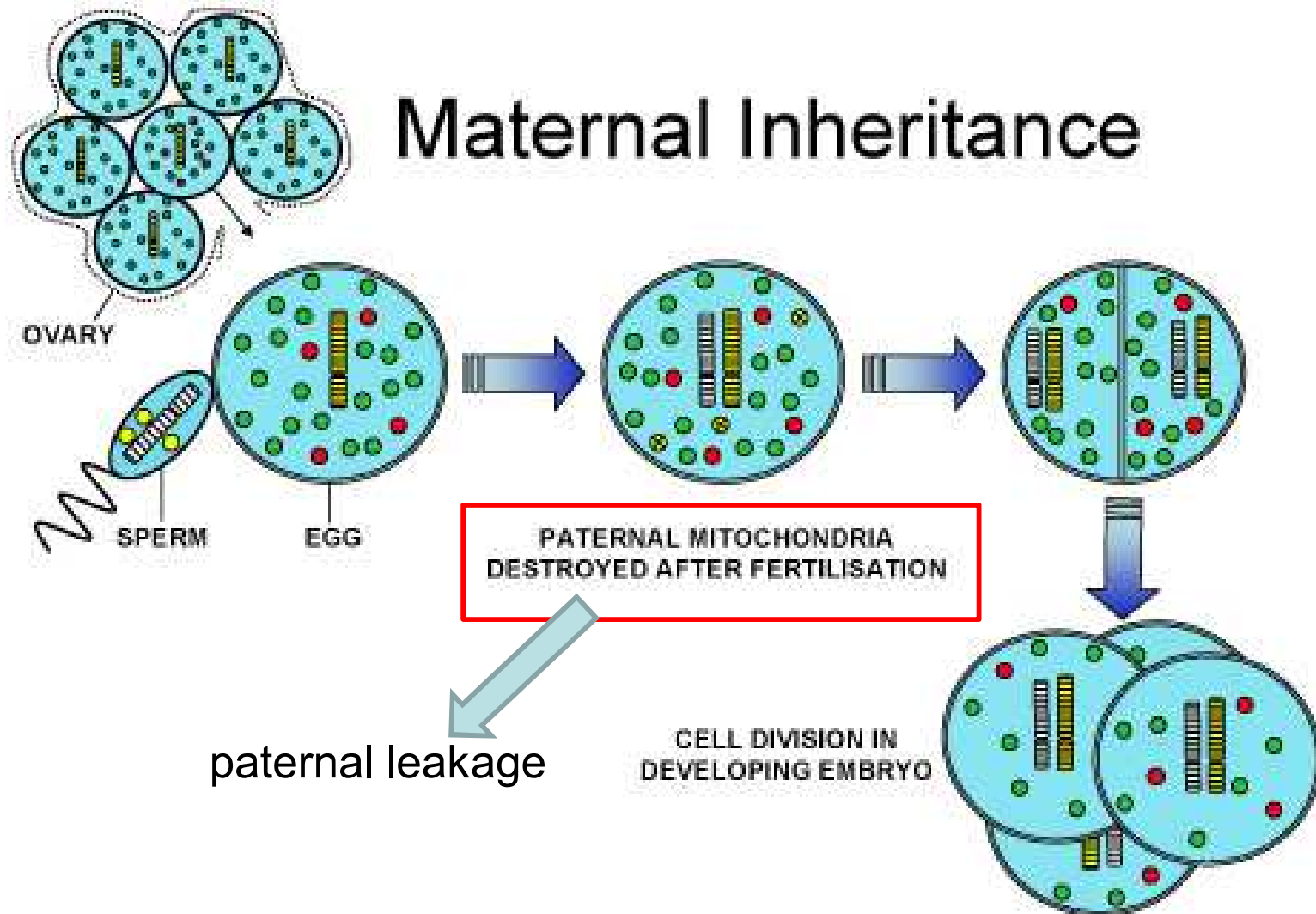


## B Crayfish



number of suggested barcoded taxa based on 3% divergence on COI with/without numts (identified by stop-codons and indels)

# Heteroplasmy



- well studied mitochondrial disorders in human
- low  $N_e$  of mtDNA → usually fast fixation of new mutations – mitochondrial bottleneck



# Paternal leakage

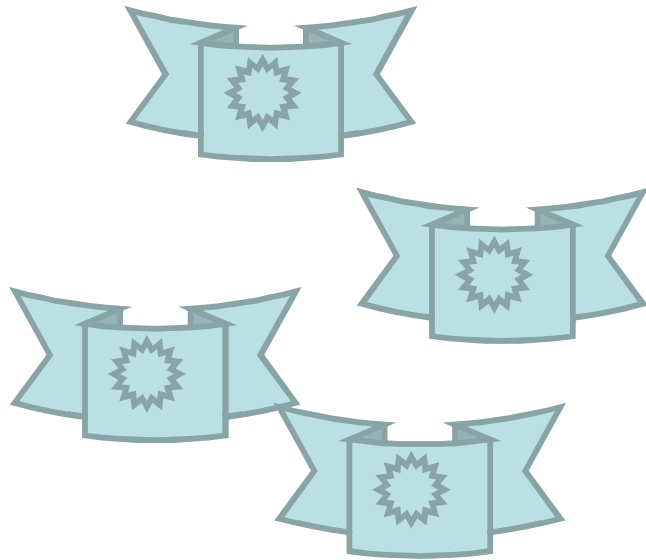
## Extensive paternal mtDNA leakage in natural populations of *Drosophila melanogaster*

MARIA D. S. NUNES,† MARLIES DOLEZAL and CHRISTIAN SCHLÖTTERER  
*Institut für Populationsgenetik, Vetmeduni Vienna, Veterinärplatz 1, A-1210 Vienna, Austria*

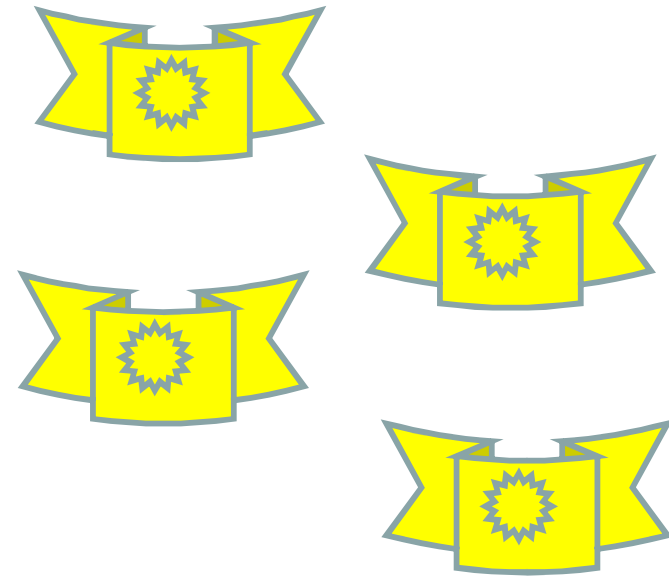
- allele-specific real-time quantitative PCR (RT-qPCR) → heteroplasmie je asi častý jev
- 14 % jedinců, ale velmi nízká frekvence druhého haplotypu
- paternal leakage 6 %



# 4. Introgression of mtDNA



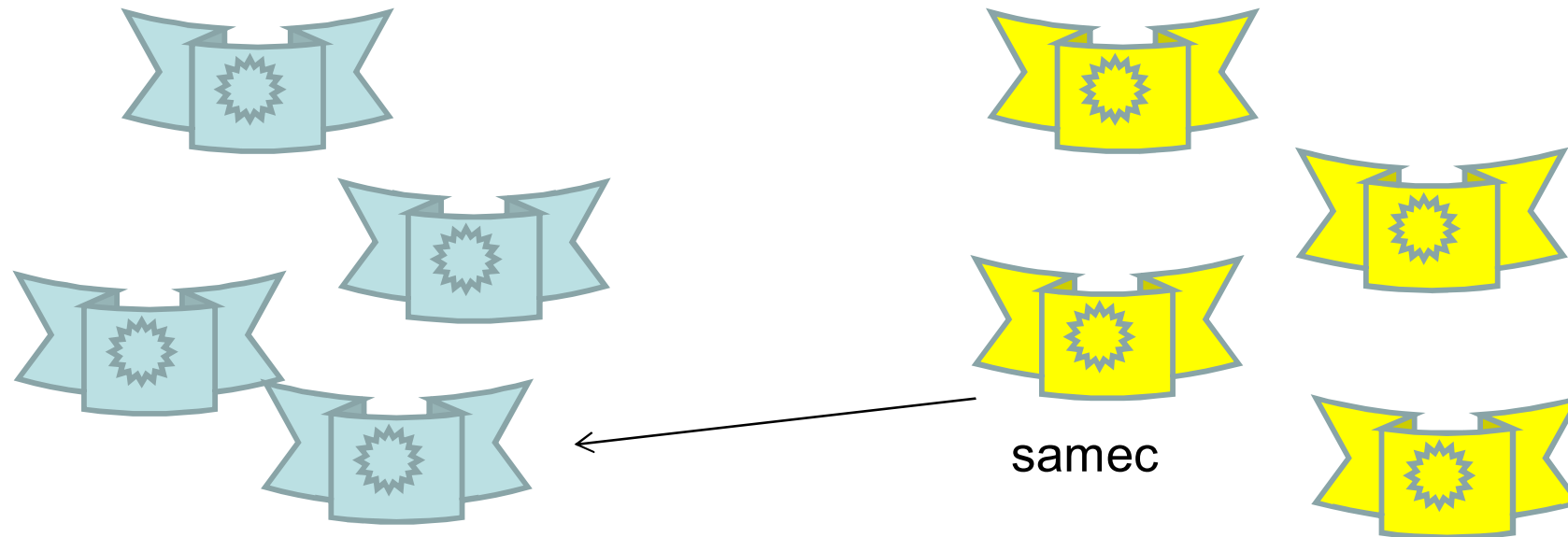
*Myotis myotis* - Evropa



*Myotis blythii* - Asie



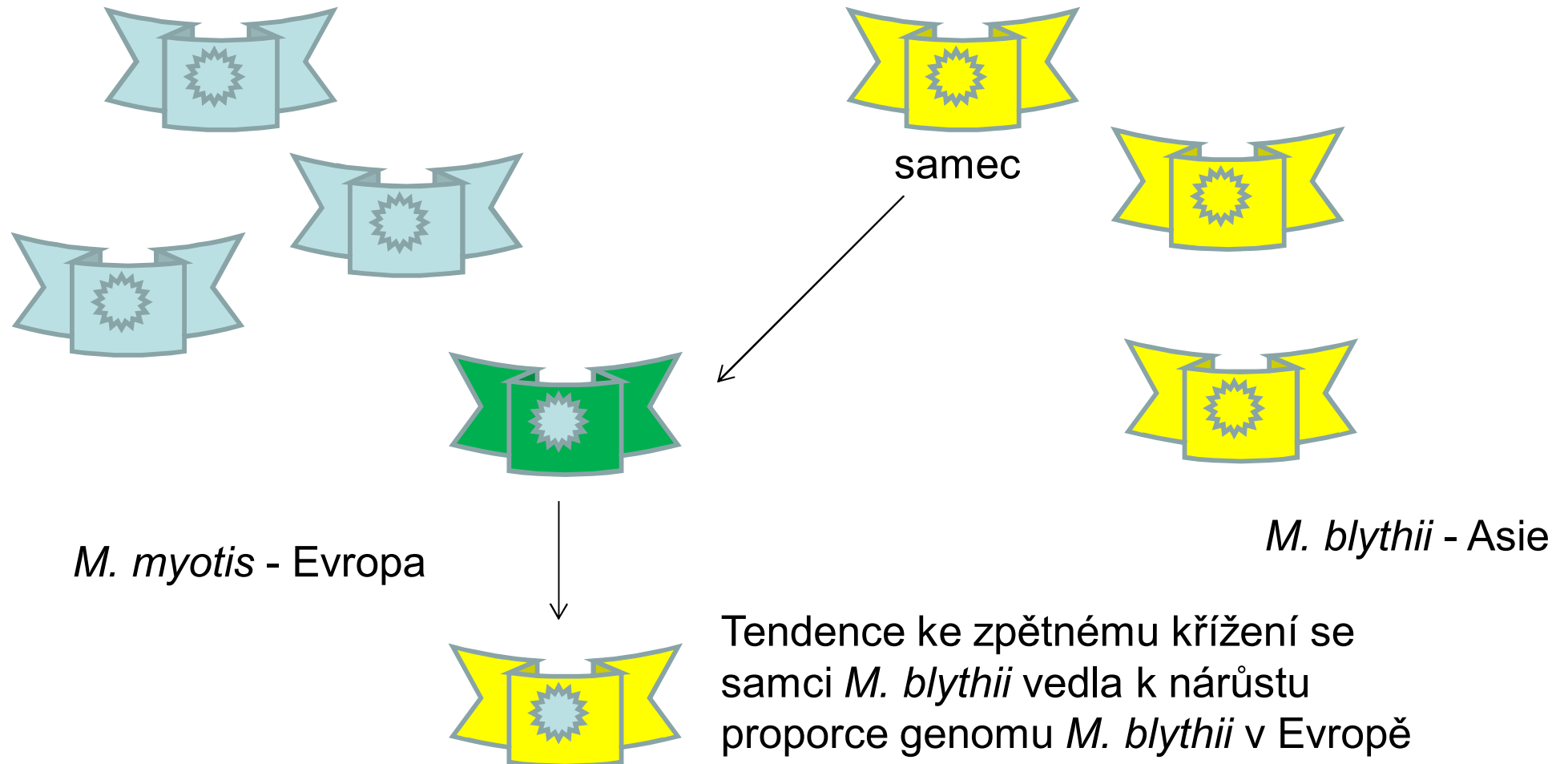
# Příklad: *Myotis blythii* vs. *Myotis myotis* - introgrese mtDNA



*M. myotis* - Evropa

*M. blythii* - Asie

# Příklad: *Myotis blythii* vs. *Myotis myotis* - introgrese mtDNA



Kolonizující (invazní) druh „ukradne“ mtDNA původnímu druhu (Currat et al. 2008)

# Plio-Pleistocene history of West African Sudanian savanna and the phylogeography of the *Praomys daltoni* complex (Rodentia): the environment/geography/genetic interplay

J. BRYJA,\* L. GRANJON,<sup>†1</sup> G. DOBIGNY,<sup>†2</sup> H. PATZENHAUEROVÁ,\* A. KONEČNÝ,\*<sup>†</sup>  
 J. M. DUPLANTIER,<sup>†</sup> P. GAUTHIER,<sup>†</sup> M. COLYN,<sup>‡</sup> L. DURNEZ,<sup>§</sup> A. LALIS<sup>¶</sup> and V. NICOLAS<sup>¶</sup>

How many species?

Čtyři podle mtDNA genotypu – cca 7% divergence (cyt b) -/97/-

Dva druhy podle fenotypu?

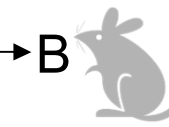
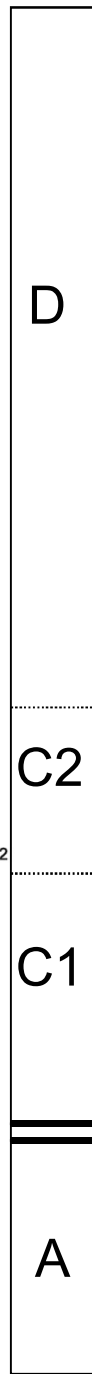
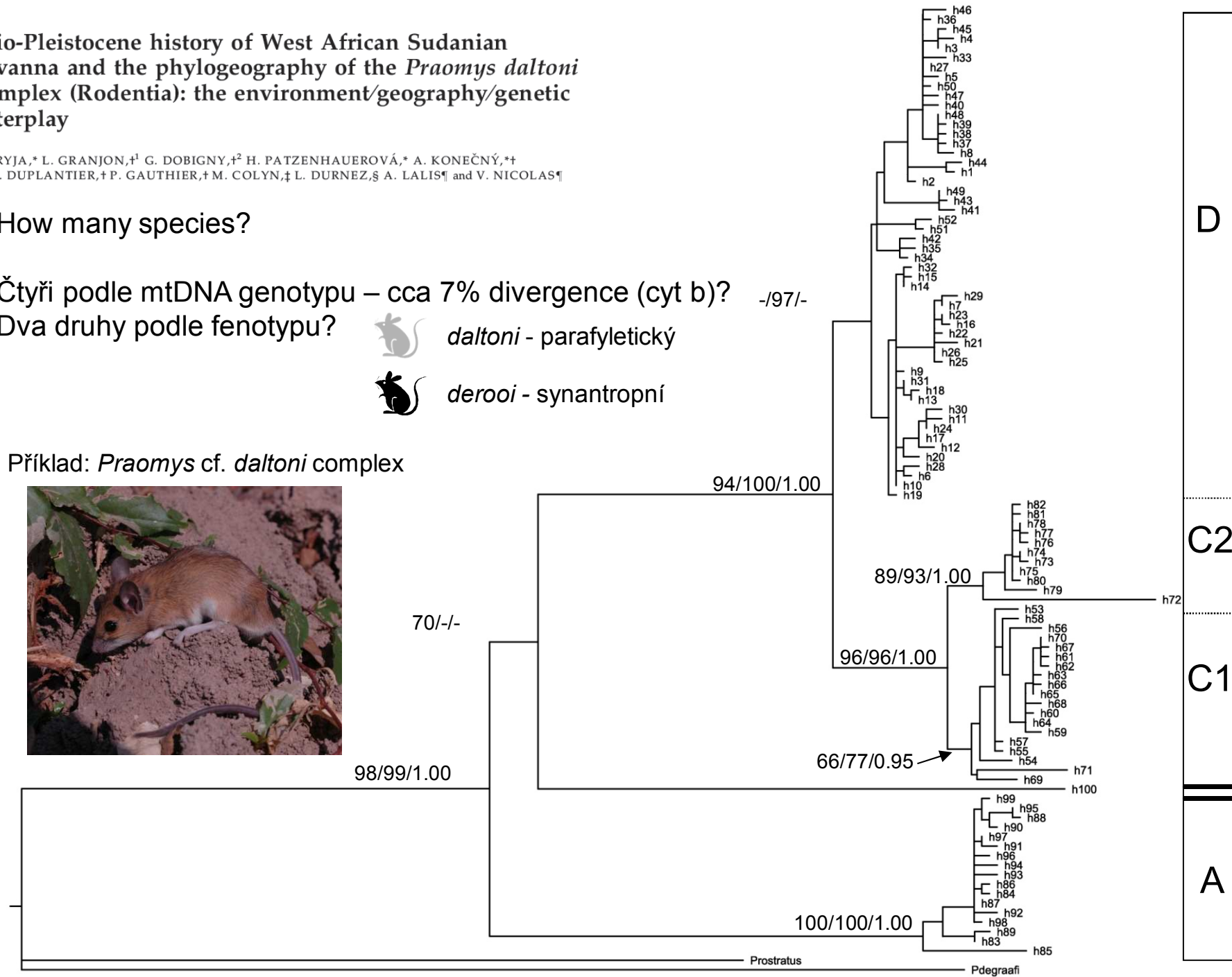


*daltoni* - parafyletický



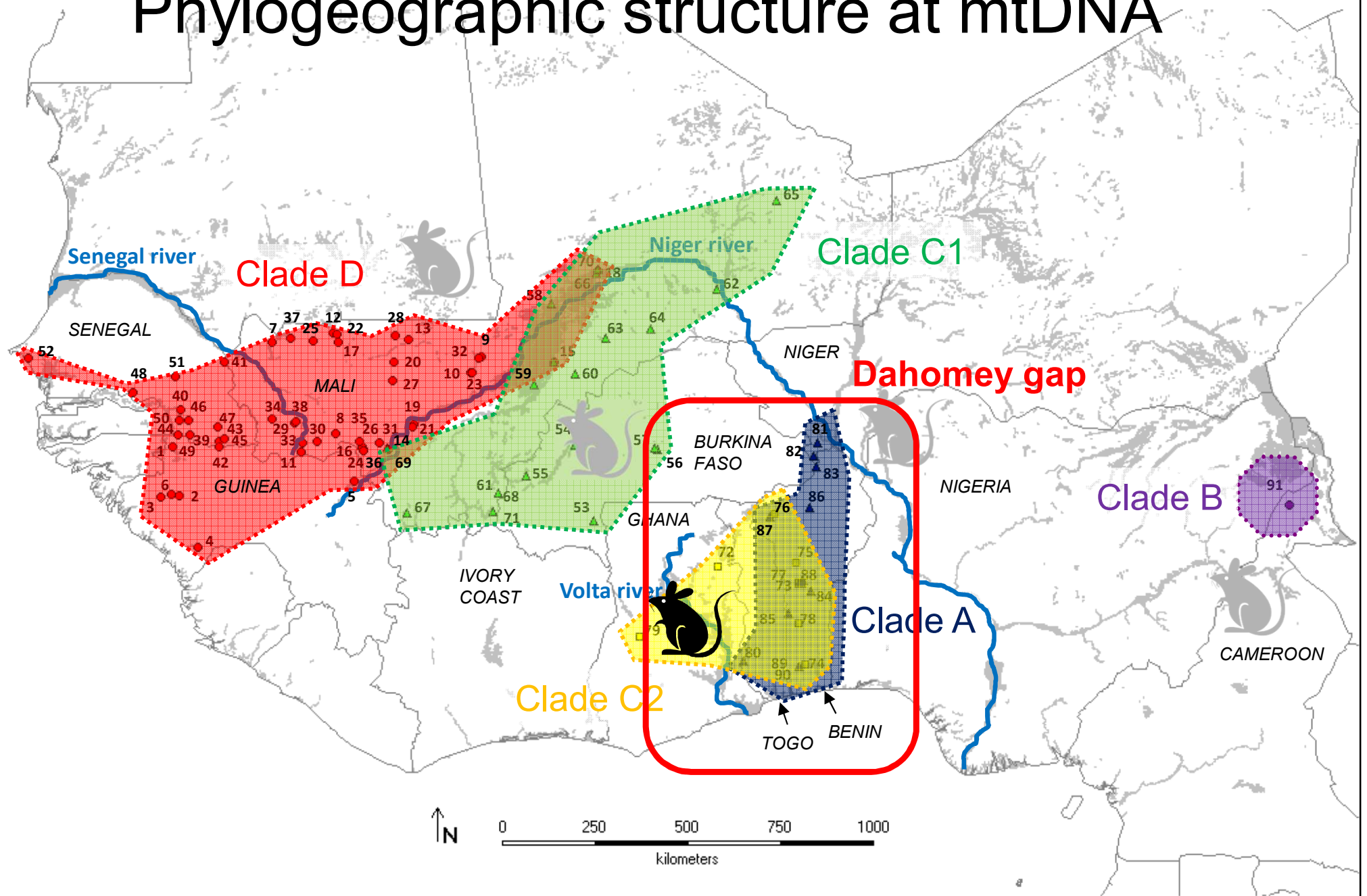
*derooi* - synantropní

Příklad: *Praomys* cf. *daltoni* complex





# Phylogeographic structure at mtDNA



# Partial mtDNA introgression in Dahomey gap

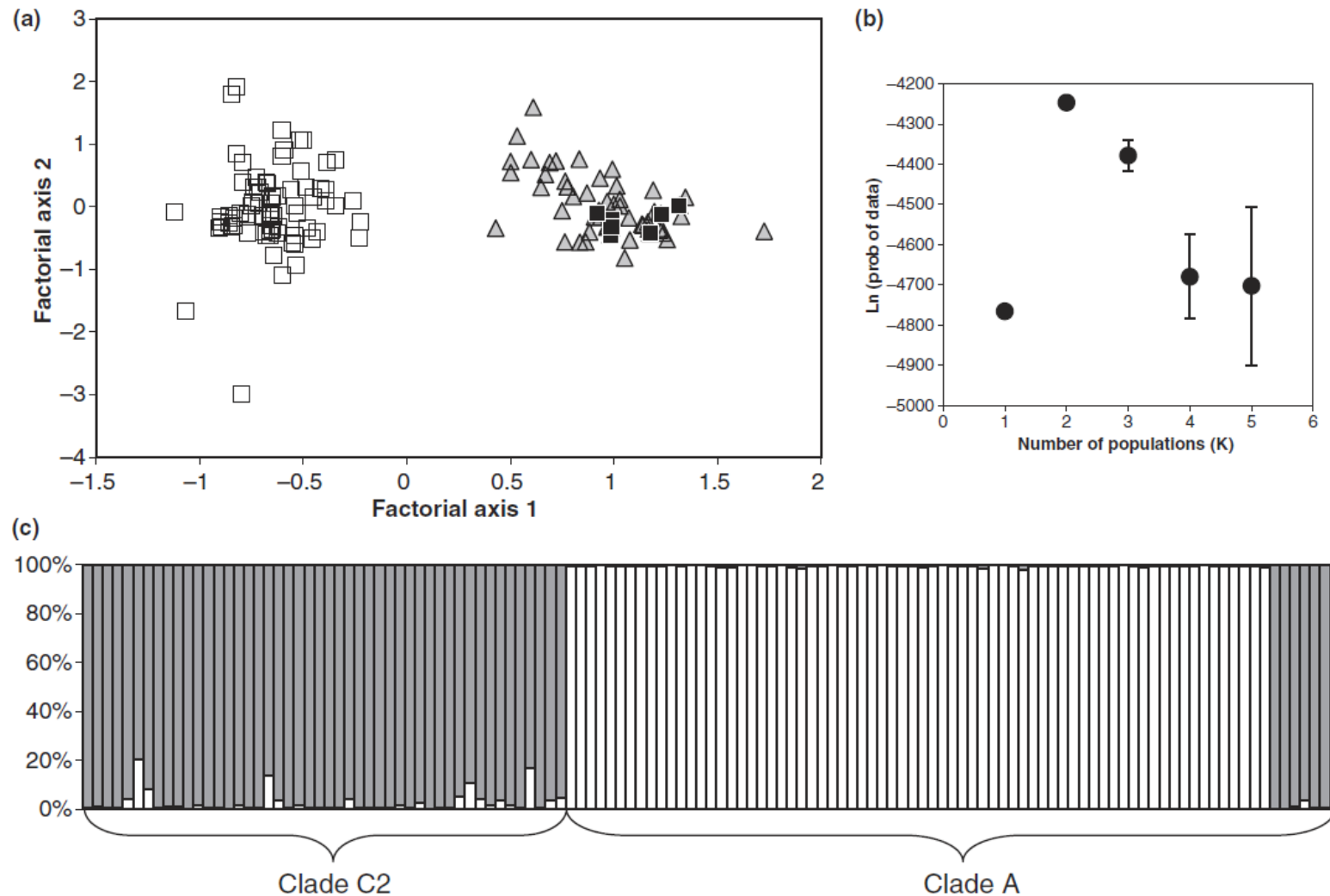
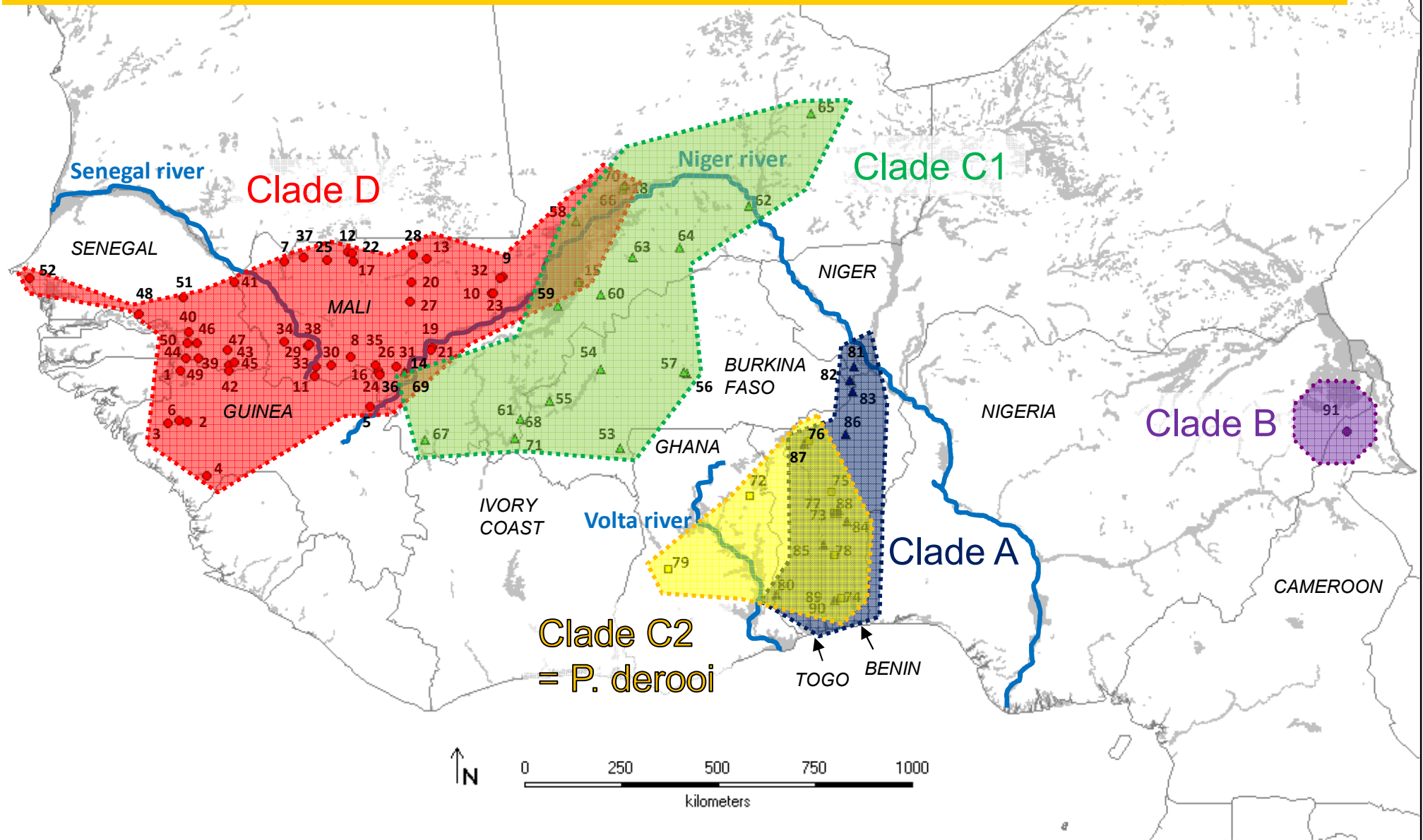
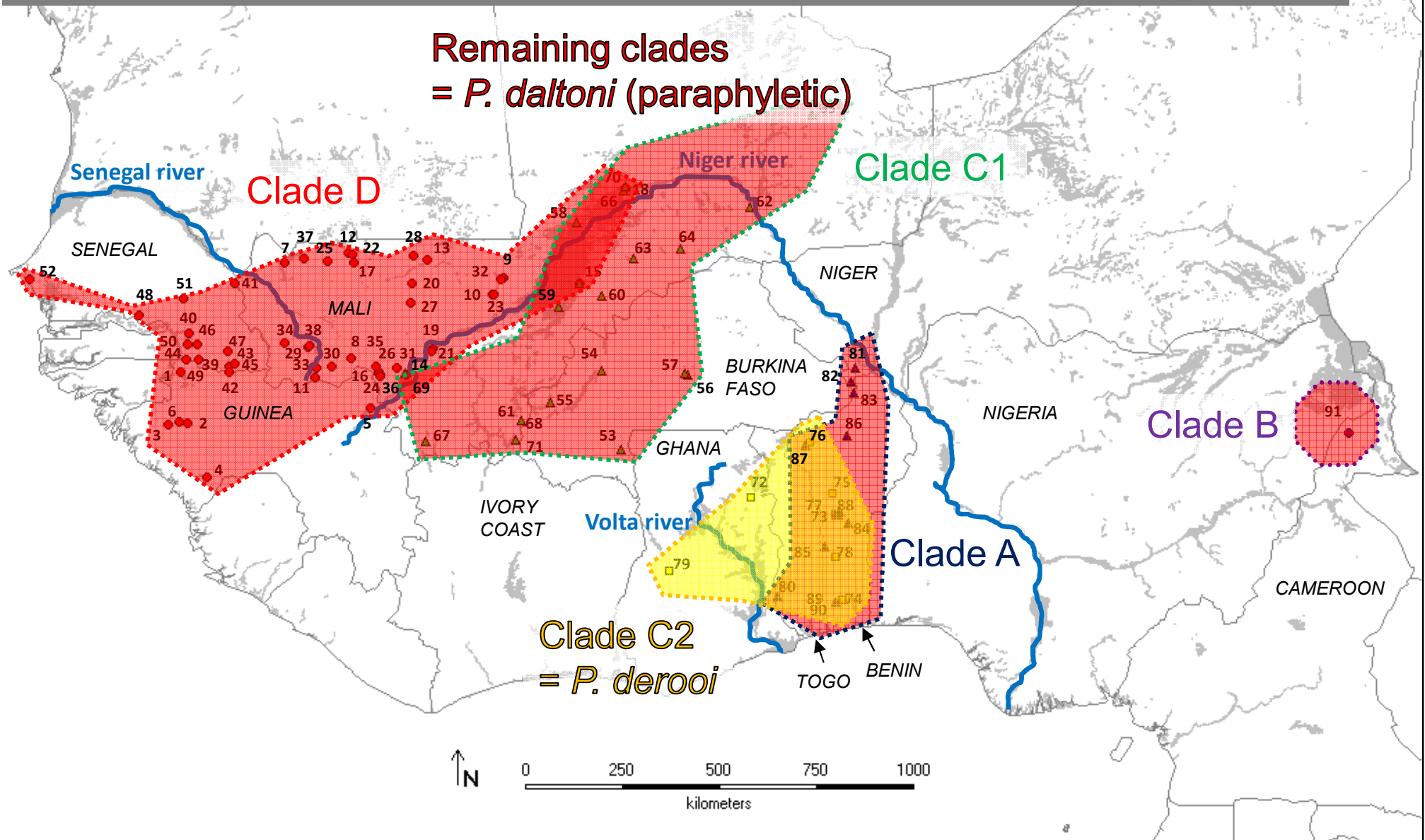


Fig. 6 Results of microsatellite analysis of individuals from Benin. (a) Factorial correspondence analysis in Genetix; *squares*—individuals with mtDNA from clade A, *triangles*—from clade C2. Note the seven individuals with mtDNA from clade A, but clustered within individuals with mtDNA from clade C2 (marked by *black squares*). (b) Likelihood of models in STRUCTURE for increasing number of populations ( $K$ ); the highest likelihood is observed for  $K = 2$ . (c) Assignment of individuals to particular populations using the best model (i.e.  $K = 2$ ; the run with the highest likelihood) in STRUCTURE.

# What is species???

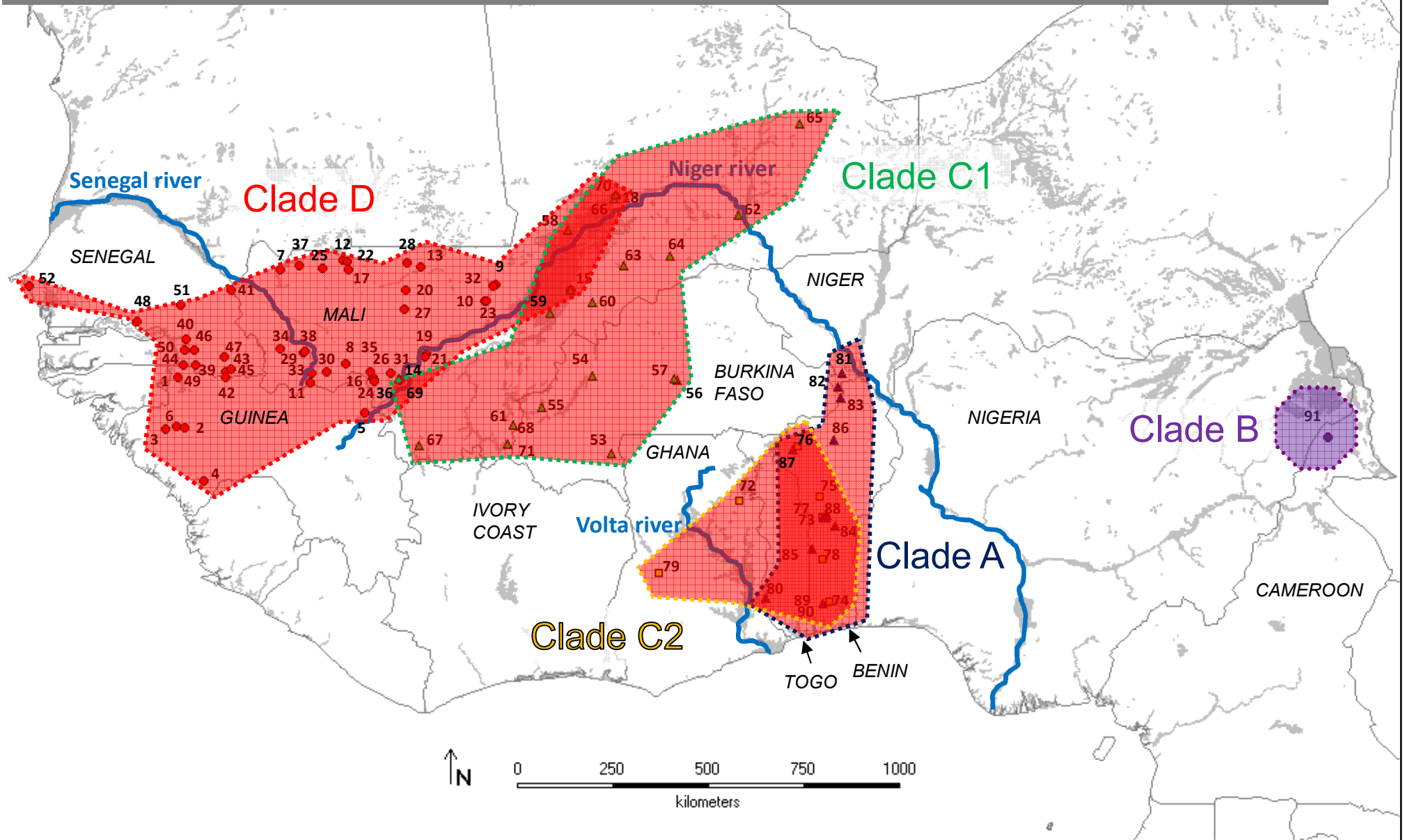


# Morphology and ecology



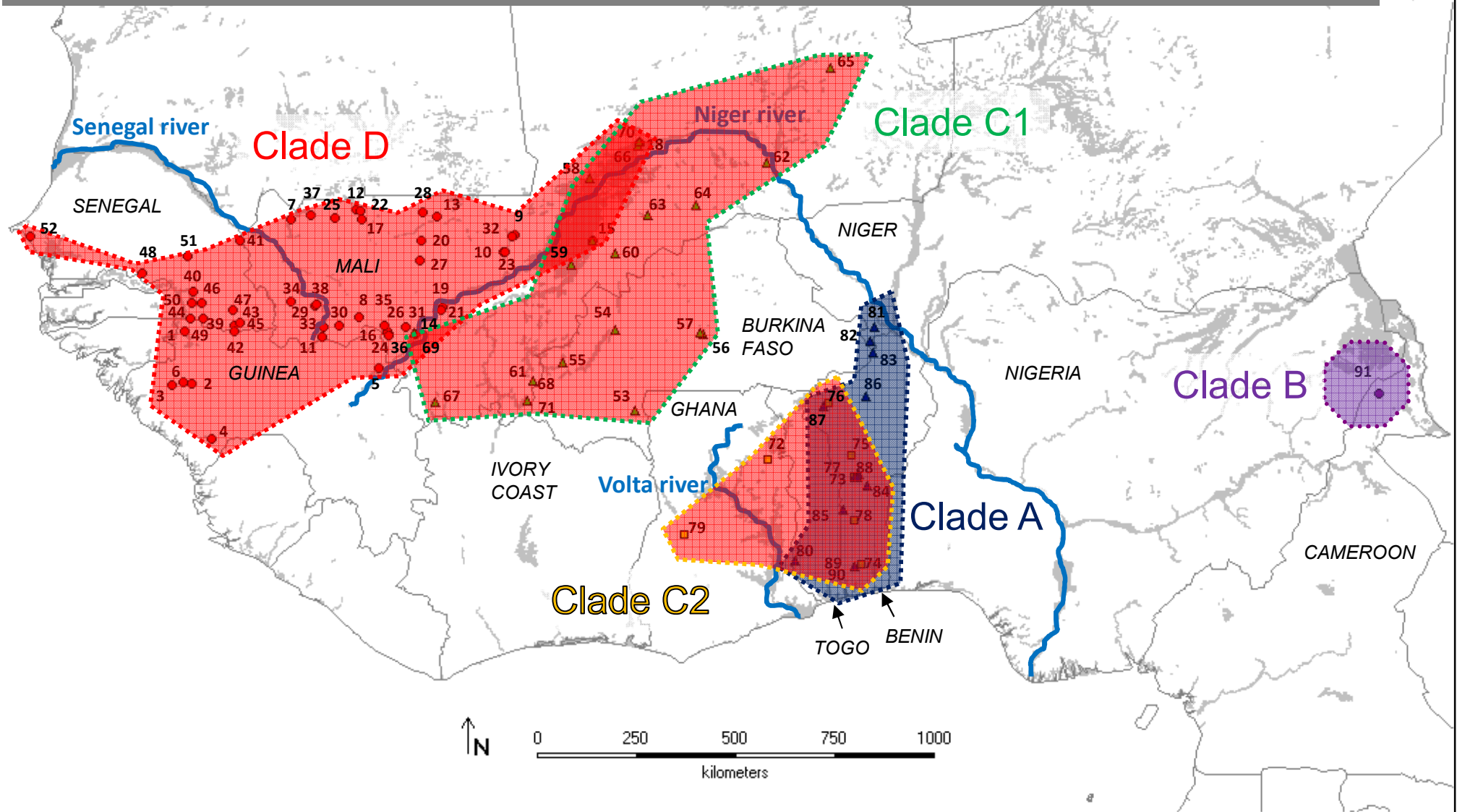


# Karyotypes

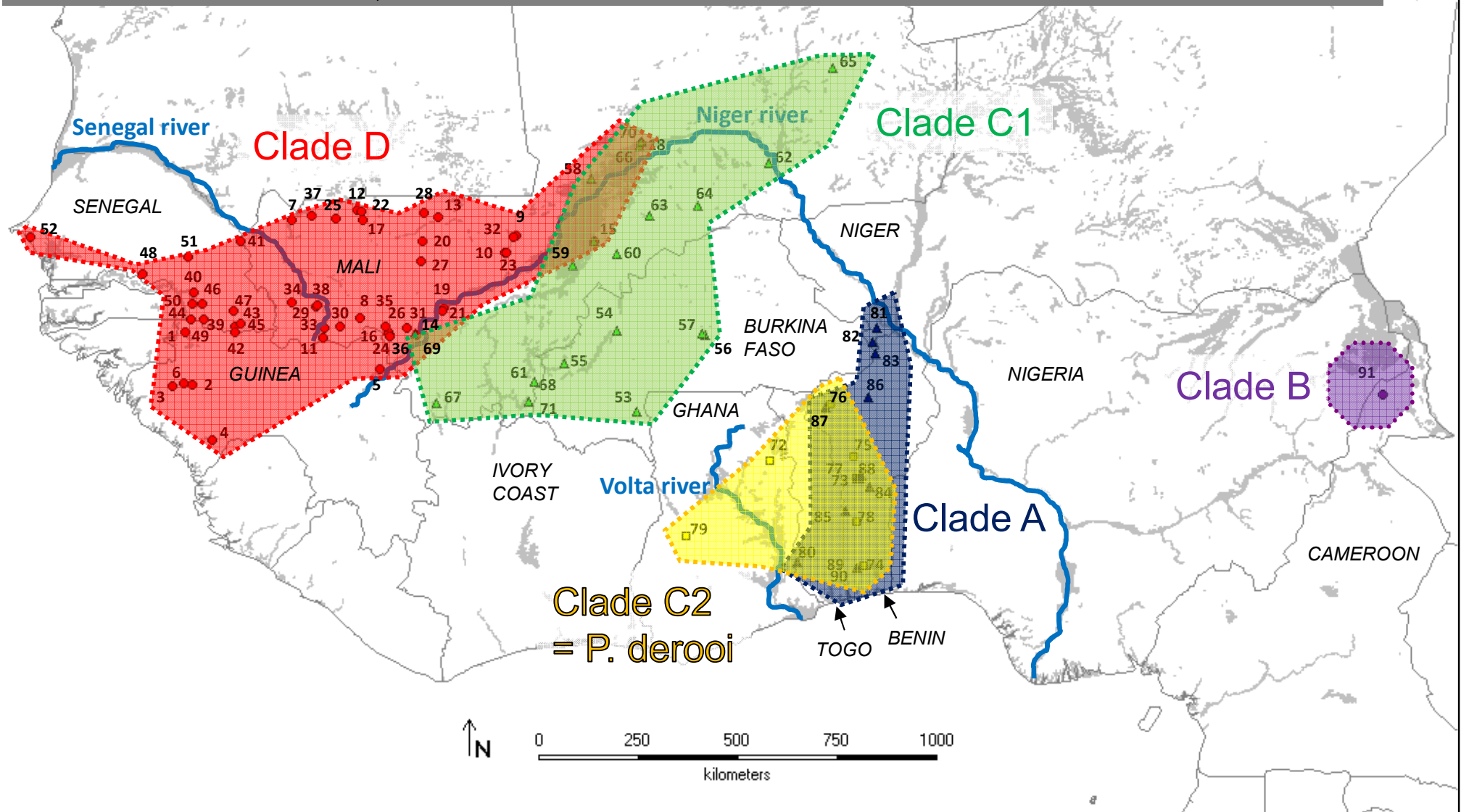




# Mitochondrial DNA + microsatellites in Benin + karyotypes



Splitting approach taking morphology and ecology into account – the reproductive barriers between clades A, B, C1, and D remains to be identified



# Identifikace jedinců

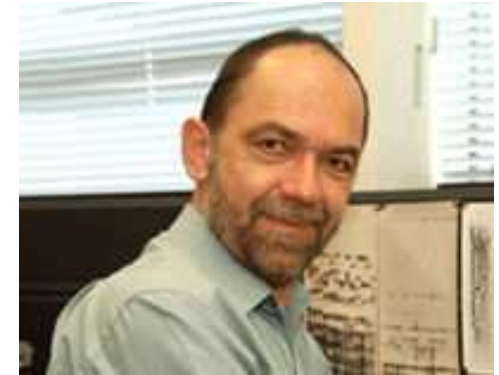
DNA fingerprinting  
(DNA profiling)

# Identifikace jedinců – kdy?

- většinou když jedince nevidíme (příp. u zpětných odchyťů, když chybí značka)
- neinvazivní genetika – skrytá zvířata, vzorky trusu, moči, chlupů atd. – dá se spojit např. s individuální variabilitou potravy
- forenzní genetika – identifikace DNA v živočišných produktech, pytláctví, atd.
- druhová ochrana – např. sokolnictví (potvrzení rodičovství) apod.

# Identifikace jedince – unikátní multilokusový genotyp

- minisatelitový fingerprinting
  - velké množství kvalitní DNA, technická náročnost
  - + univerzalita
- AFLP
  - kvalitní nedegradovaná DNA
  - + univerzalita



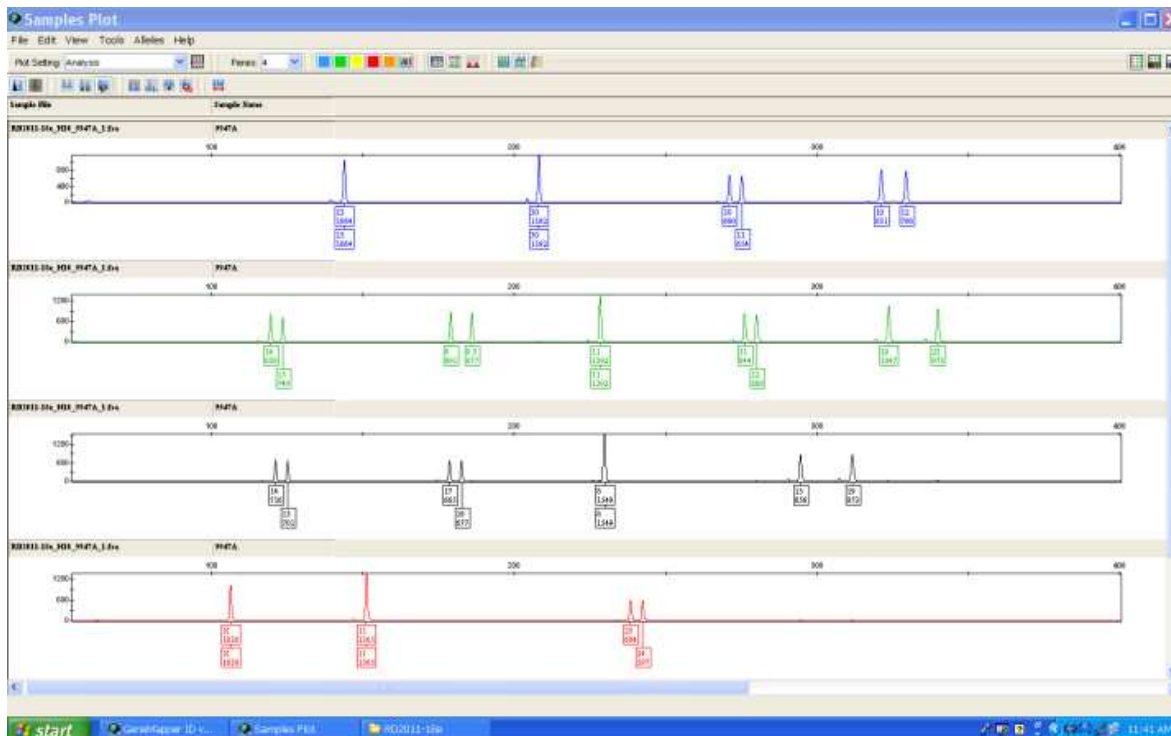
Alec Jeffreys, the pioneer of DNA profiling

- Mikrosatelity
  - + stačí malé množství nekvalitní DNA, **optimální pro neinvazivní přístupy**
  - je nutné znát konkrétní lokusy a sekvence specifických primerů
- SNPs
  - + velké množství znaků v genomu
  - malá variabilita, je potřeba hodně lokusů, nutno znát konkrétní lokusy

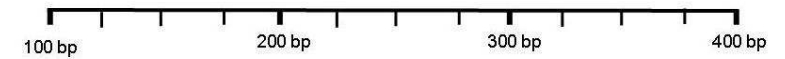




# Individual human identification



AmpFISTR® Identifiler™



D8S1179 D21S11 D7S820 CSF1PO

D3S1358 TH01 D13S317 D16S539 D2S1338

D19S433 vWA TPOX D18S51

A D5S818 FGA



GS500-internal lane standard

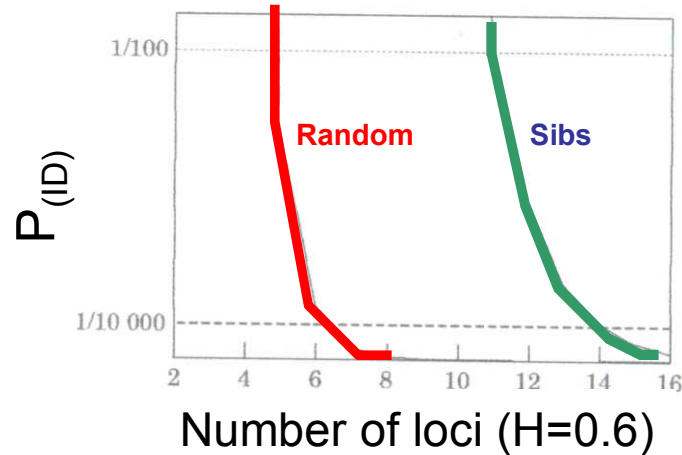
16 loci = reliable individual identification  
(Euro-American population)



# Identifikace jedinců závisí na stupni polymorfismu

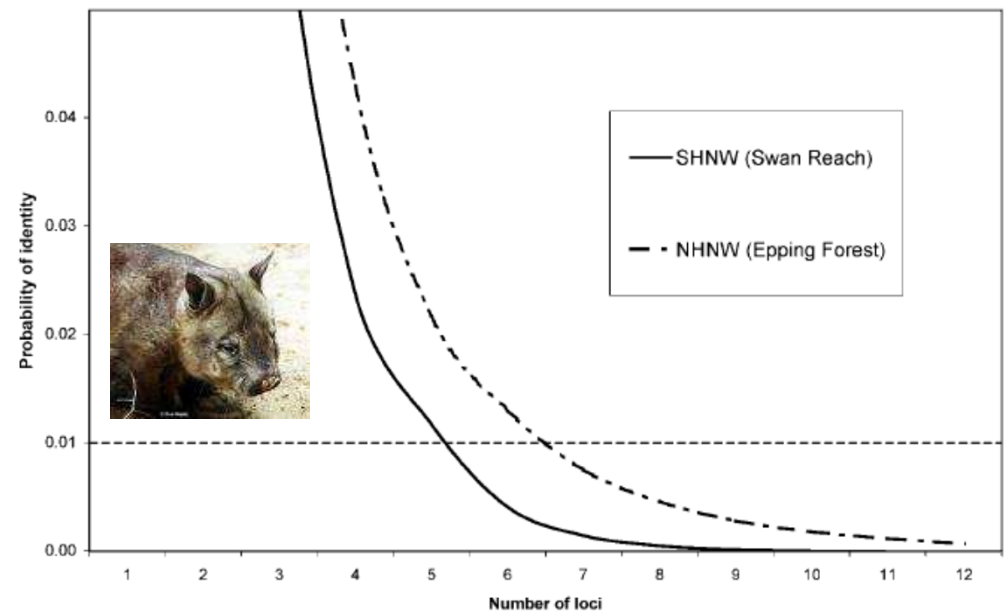
- multilocus microsatellite fingerprinting – power estimated as „probability of identity“ ( $P_{(ID)}$ ) (Waits et al. 2001) – např. program GenAlex

$$P_{(ID)} = \sum p_i^4 + \sum \sum (2 p_i p_j)^2$$



$$P_{(ID)sib} = 0.25 + (0.5 \sum p_i^2) + [0.5(\sum p_i^2)^2] - (0.25 \sum p_i^4)$$

- pilot studies with tissue samples are required to identify  $P_{(ID)}$  in a population studied by e.g. non-invasive methods





# Medvědi v Pyrenejích

*Taberlet et al. 1997*



- Trus a chlupy
- 24 mikrosatelitových lokusů
- 4 samci a jedna samice s unikátními multilokusovými genotypy (o jednoho víc než podle stop a fotografií)
- Multiple-tube approach, mnohonásobné opakování PCR reakcí

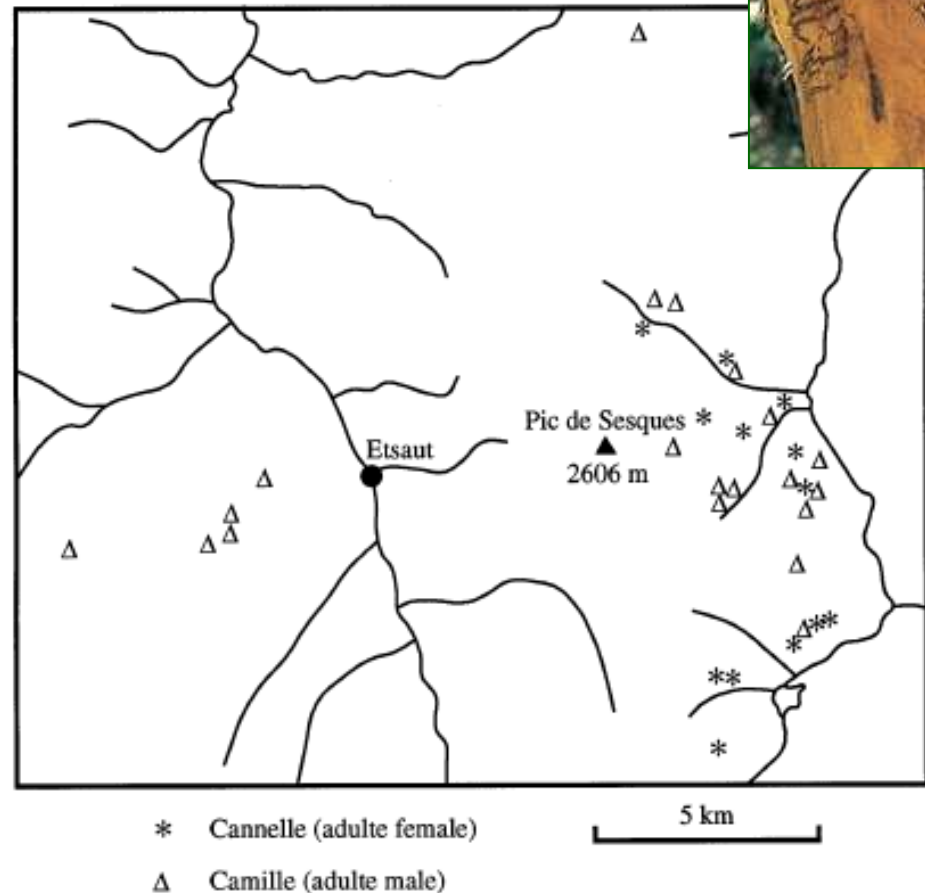


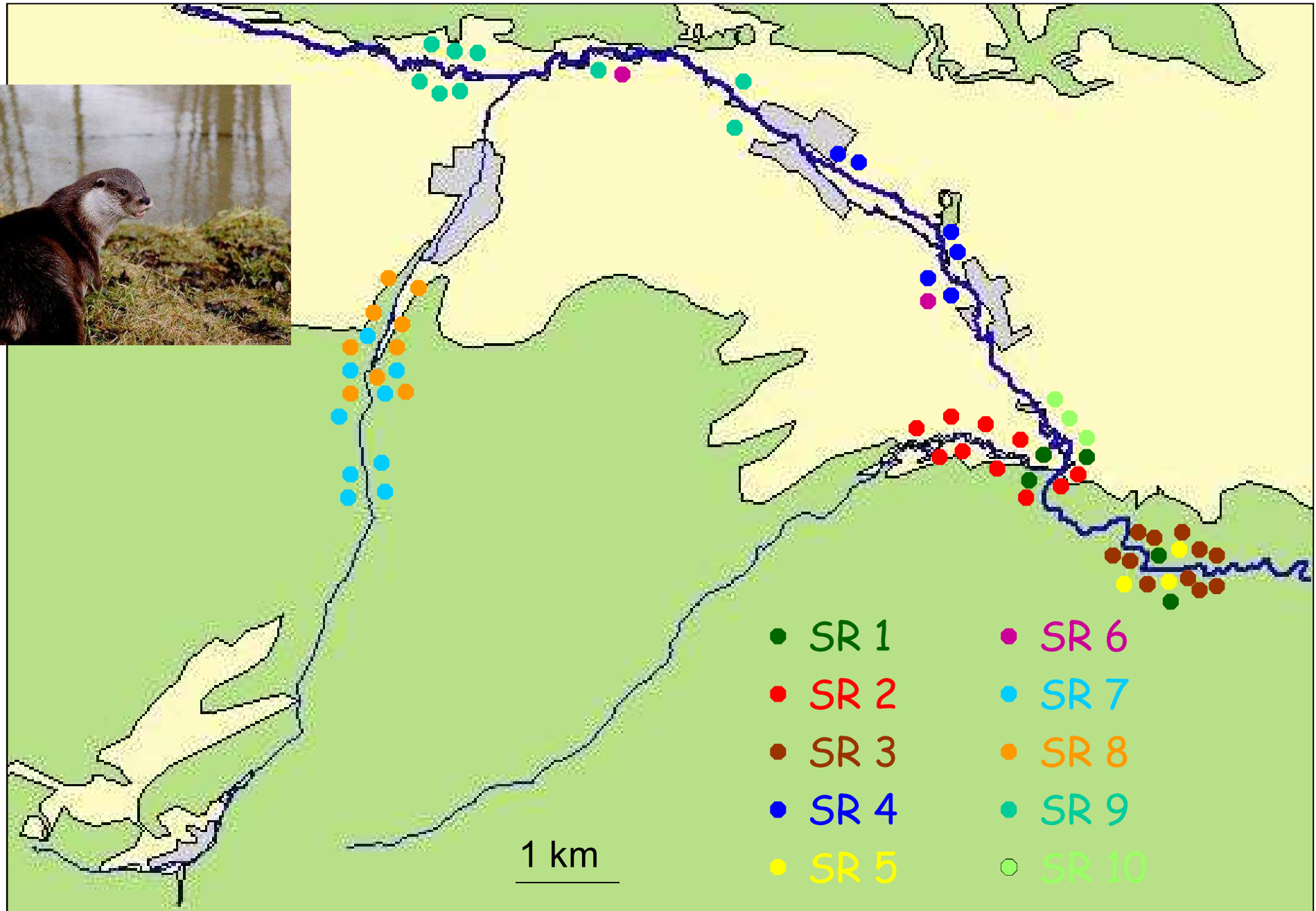
Fig. 3 Home range of two Pyrenean brown bears obtained by noninvasive genetic sampling and genotyping.

# Analýza prostorové aktivity

- P. Hájková – PhD thesis

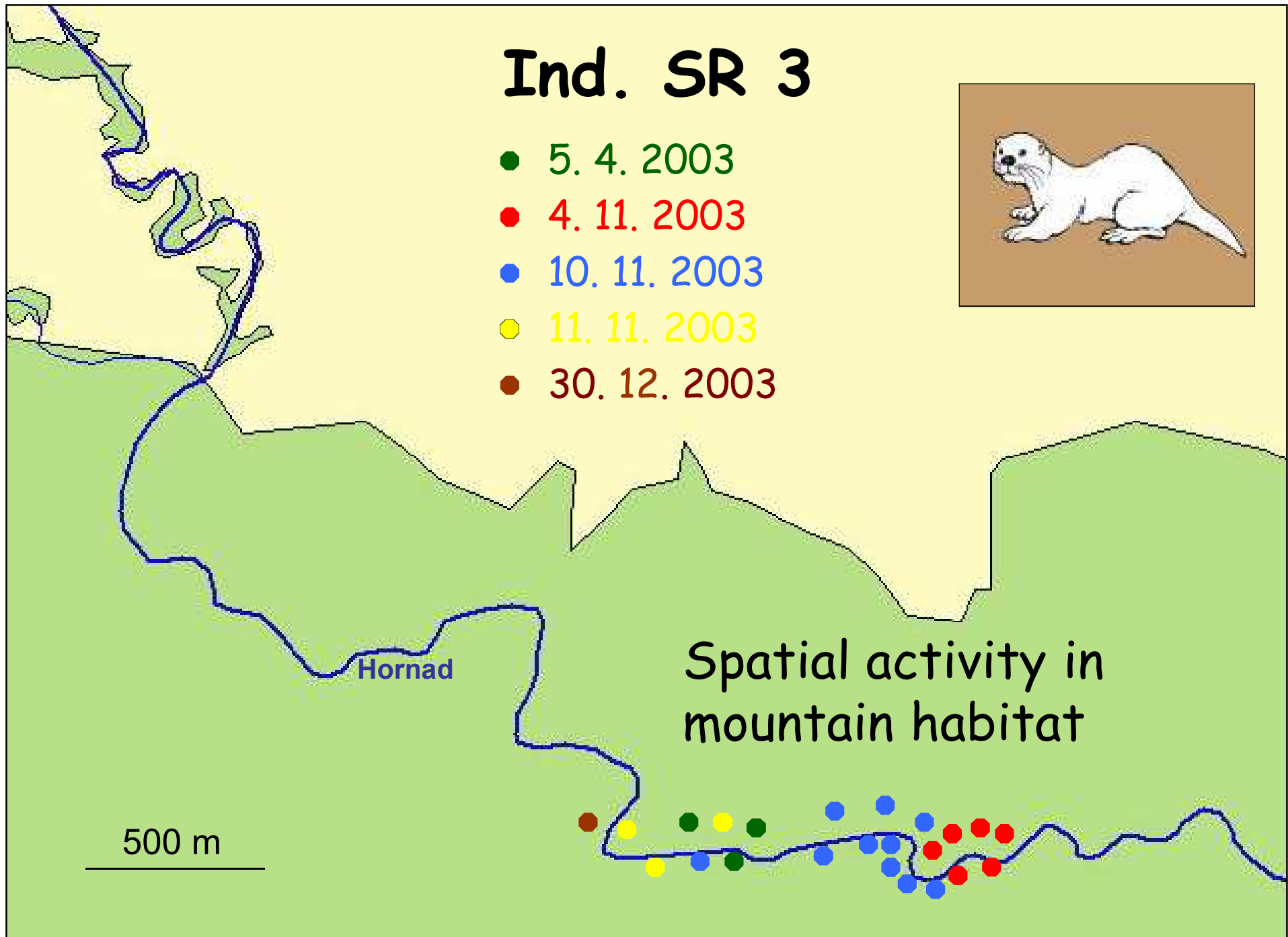
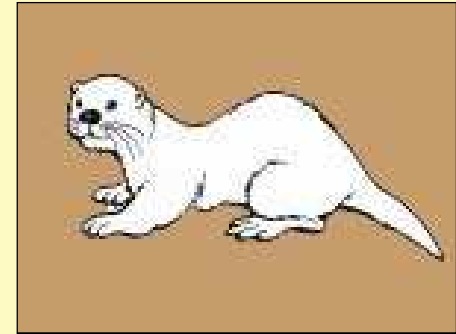


# Identified individuals - Hornád, NP Slovenský Raj



# Ind. SR 3

- 5. 4. 2003
- 4. 11. 2003
- 10. 11. 2003
- 11. 11. 2003
- 30. 12. 2003



Spatial activity in  
mountain habitat

500 m

Hornad



# Human forensic genetics

- **Pozůstatky vojáků z války** Vietnam a Korea

Identifikace na základě mtDNA příbuzných osob  
(lze jen někdy)

V současnosti: vzorek DNA (krve) při odvodu, jiné markery

Armed Forces Repository of Specimen Samples for the Identification of Remains



- **Soudní pře**  
Clinton-Lewinská  
Pozůstatky ruského cara Nikolaje II



- **Kriminalistika**

- **Oběti tragických událostí**



# Klony

## Bambus *Sasa senanensis*

- Suyama et al. 2000
- Plocha 10 hektarů
- AFLP
- 22 klonů
- Klon na ploše 300 m v průměru



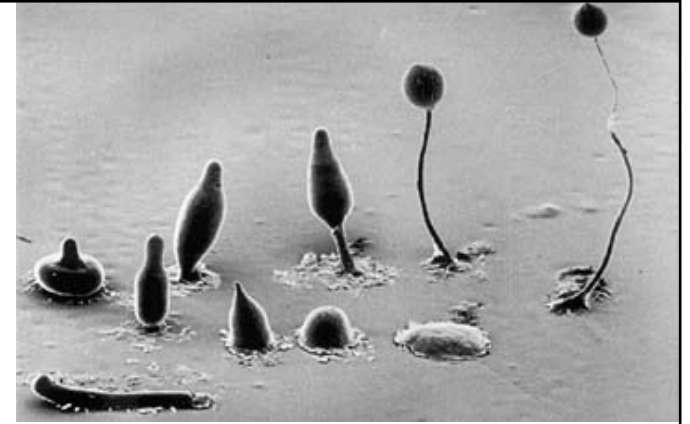
# Slavní klonální bezobratlí

- Rotifera – Bdelloidea
- Ostracoda  
(*Darwinula*)
- Partenogenetické  
klony vysokého stáří  
(milióny let)

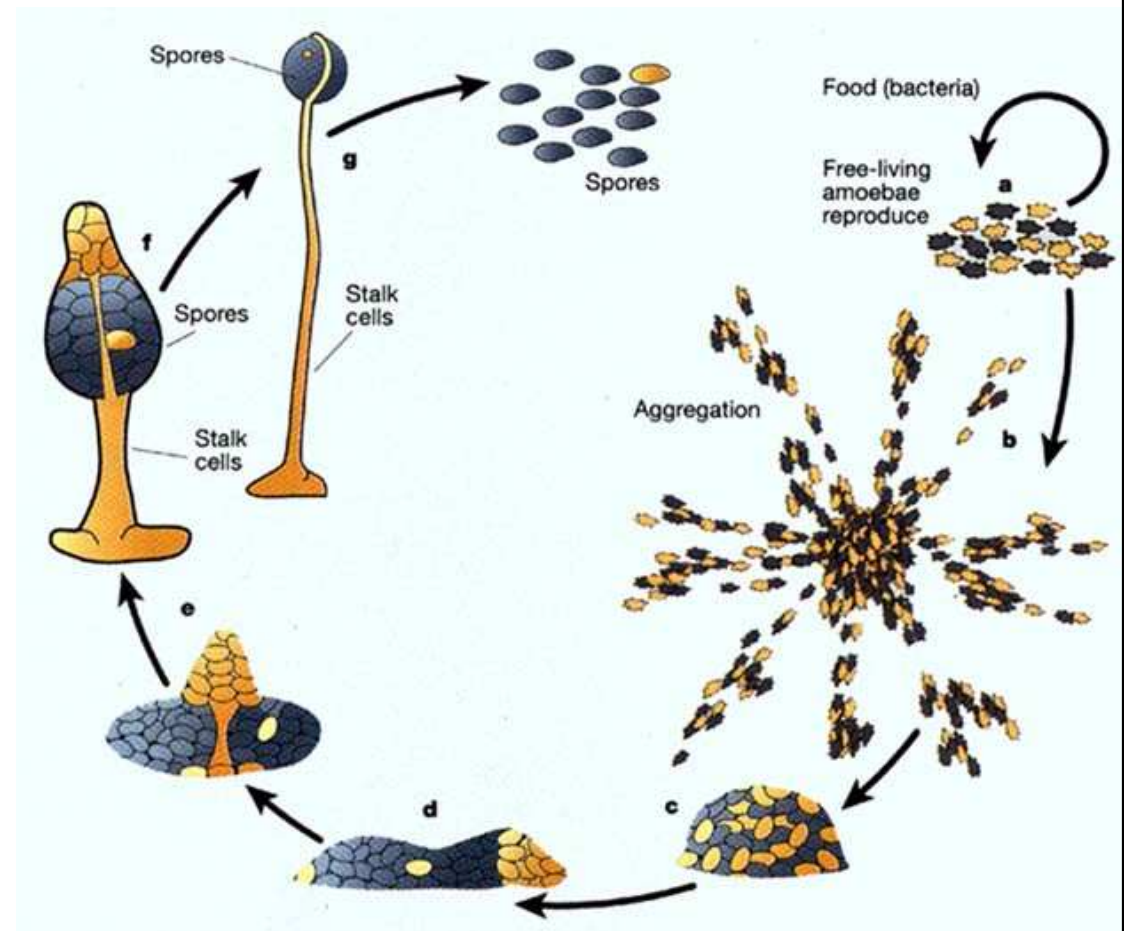


*Darwinula stevensoni*

# Genetické chiméry



- organismy složené z buněk s různými genotypy
- *Dictyostelium discoideum*  
chimérismus je pravidelná součást života





# Genetické chiméry

- *Ficus* srůst kořenů různých jedinců
- sumky *Botryllus schlosseri*  
chimérické kolonie  
příbuzní jedinci
- *Diplosoma listerianum*  
i nepříbuzní

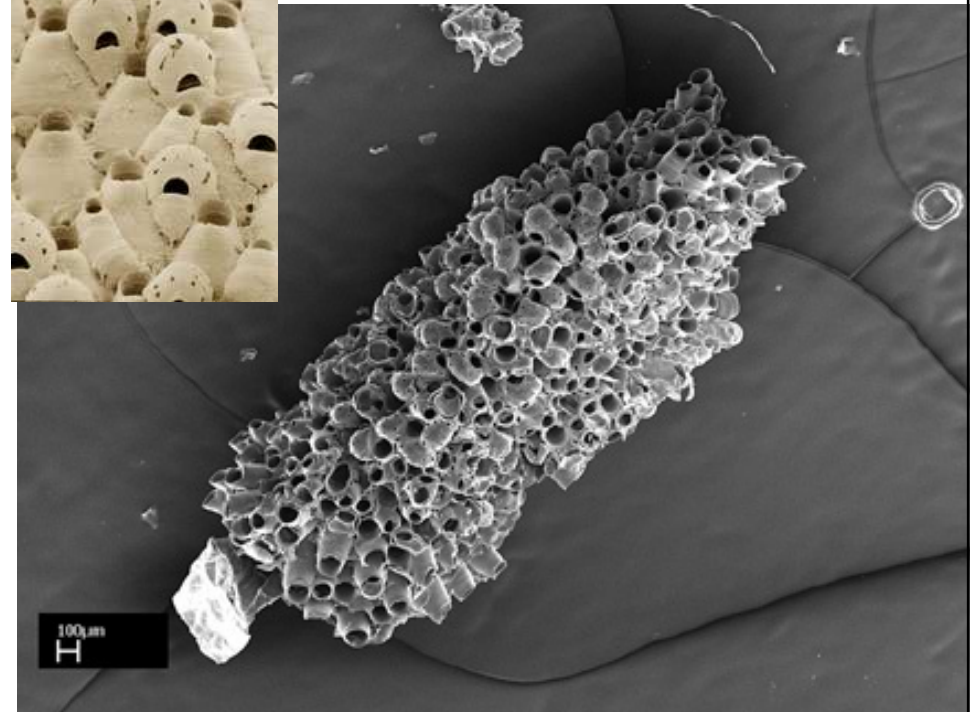
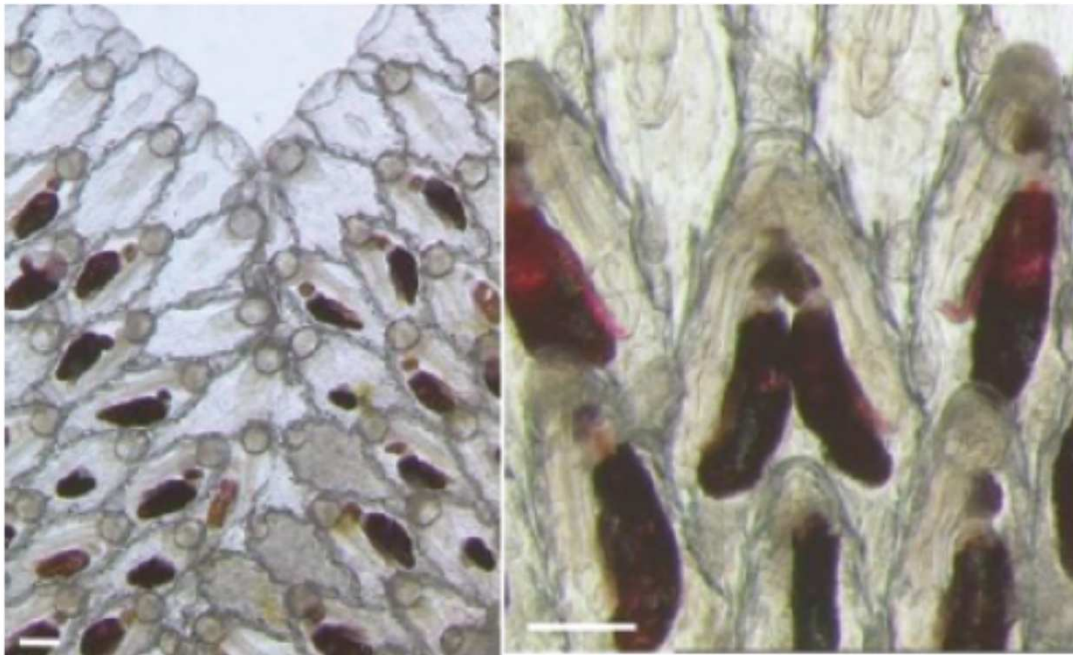




# *Celleporella hyalina* (Bryozoa)

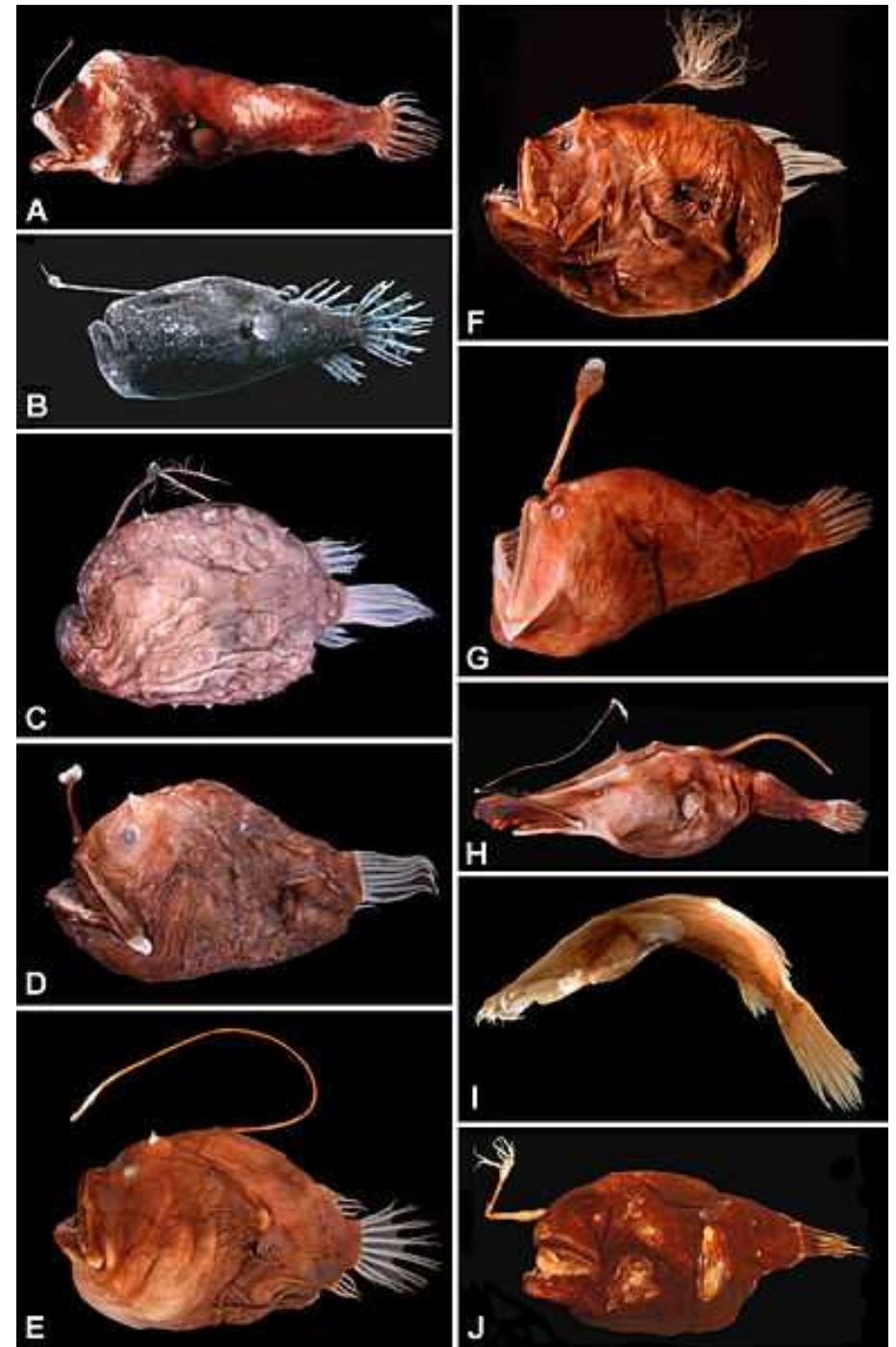
Hughes et al. 2004

- Pravděpodobnost fúze koreluje s příbuzností
- Histokompatibilita
- Lepší rozpoznávání v pokročilejších fázích  
→ dozrávání imunokompetence
- Speciální proteiny (spongikany...)



# Ceratioid anglerfish

- miniaturní samec po narození vyhledá samici, její kůže vyloučí hydrolytický enzym a samec přiroste
- vzniká hermafroditická chiméra



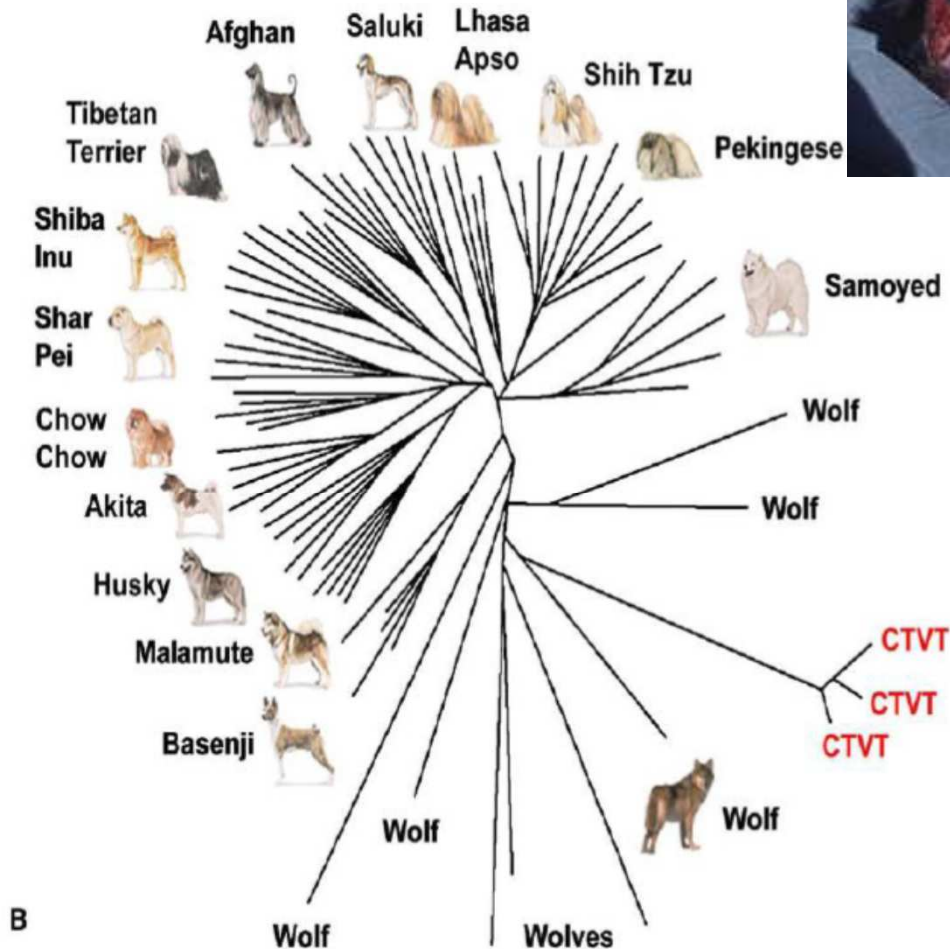
# Genetické chiméry – „microchimerism“



- kosman bělovousý *Callithrix jacchus* (asi i rod *Saguinus*)
- dizygotická dvojčata
- DNA fingerprinting krve - hematopoietické chiméry
- během embryonálního vývoje vzájemná výměna buněk kostní dřeně
- týká se to asi jen krve (neinvazivní metody – chlupy, trus → jeden genotyp)
- průnik embryonálních erytroblastů a volné DNA přes placentu i u člověka
- (pohlaví dítěte před narozením lze určit i pomocí PCR sekvencí typických pro Chr Y, jako templát je periferní krev matky)



# Canine transmissible venereal tumor (CTVT)



## Devil facial tumour disease

- parasitic cancer
- „single cell parasitic wolf“



# Známé „lidské chiméry“



## Foekje Dillema

46XX/46XY woman

- holandská atletka, mistryně na 100 a 200 m
- odmítla test na pohlaví
- mozaika zjištěna až posmrtně (v r. 2007)

### Lydia Kay Fairchild The Twin Inside Me - Chimera

#### The Twin DNA

Lydia Fairchild was twenty one when she had her first baby. Despite being separated from the baby's father, Jamie Townsend, she and Jamie had a second baby a year later. Another year on and she became pregnant for the third time after which she and Jamie split up again. With no steady work and unable to support herself and the children she applied for state benefit.

Her world was about to be turned upside down.

The State Prosecutor's Office required DNA tests from Jamie to prove that he was the father of the children and, as a matter of course, Lydia was also tested.

In December 2002 she received a phone call from the prosecutor's office asking her to come in for the results. This was unusual and it soon became apparent why. The results confirmed that Jamie was the father but they also revealed that Lydia was **not** the mother. A normal DNA test proving a mother-child link would show a 50% match between their DNA patterns. Yet Lydia's DNA showed no match at all.



Lydia Fairchild

- geneticky nepotvrzené mateřství
- chiméra matky



# Identifikace pohlaví

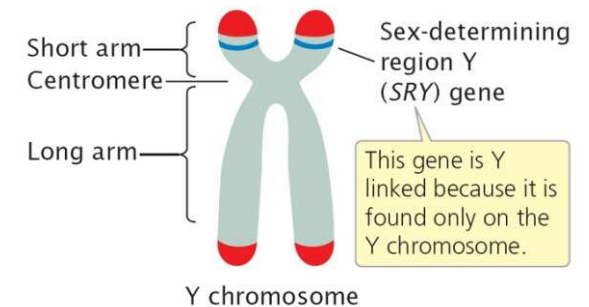
DNA sexing

# Genetická identifikace pohlaví - proč

- 1) druhy s nevýrazným pohlavním dimorfismem (ptáci, ale i řada savců)
- 2) zárodky v ranném stádiu ontogeneze (embrya, larvy)
- 3) neinvazivní metody (trus, skořápky, šupiny)

# Genetická identifikace pohlaví

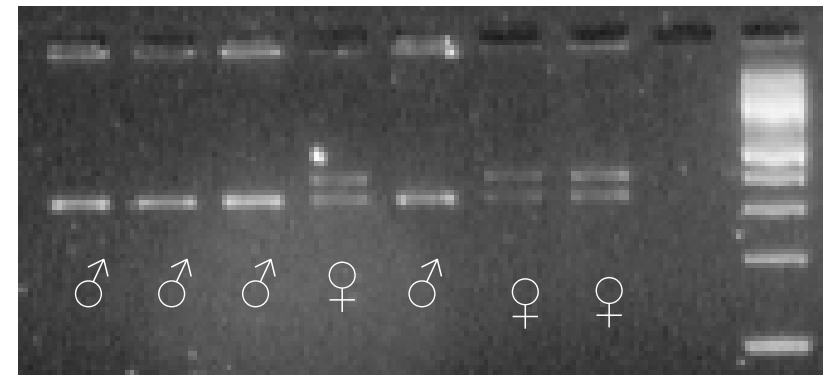
- druhy s genetickou determinací pohlaví (tj. nejčastěji pohlavní chromozómy)
- ptáci ( $\text{♂} = ZZ$ ,  $\text{♀} = ZW$ )
- savci ( $\text{♂} = XY$ ,  $\text{♀} = XX$ )
- amplifikace DNA oblasti specifické pro heterogametické pohlaví
- W, Y – malé chromozómy



# Určení pohlaví – ptáci

*Griffith et al. 1998*

- *CHD1W* a *CHD1Z*, geny na pohlavních chromosomech (chromobox-helicase-DNA-binding gene (CHD) – Griffiths & Tiwari 1995)
- Primery amplifikují introny obou genů
- Introny se mohou lišit délkou
- Existují už tři možnosti běžně používaných primerů
- Problematické druhy Struthioniformes



# *Manorina melanocephala*

(Meliphagidae) *Arnold et al. 2001*

- Synové fungují jako pomocníci
- U adultů  
2,31 samců na 1 samici
- Mláďata v hnízdě  
poměr pohlaví 1:1 (57:57)
- První se líhnou samci  
(v 17 hnízdech z 18)  
Při opouštění hnízda jsou větší a těžší

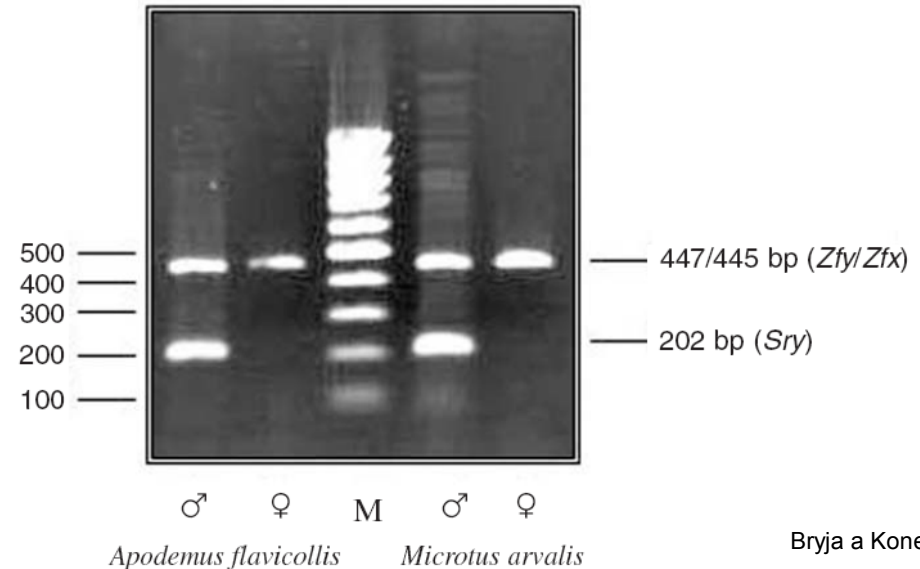


medosavka hlučná



# Určení pohlaví - savci

- Amplifikace genu na Chr Y (*Sry*)  
(nejlépe duplex PCR s genem na X nebo autosomech)



- *Microtus cabrerae*  
Sry na Chr X  
*Ellobius*, *Tokudaia*  
Sry zcela chybí
- *Nannomys*  
Velká variabilita

*M. cabrerae*



*Nannomys*

*Ellobius*



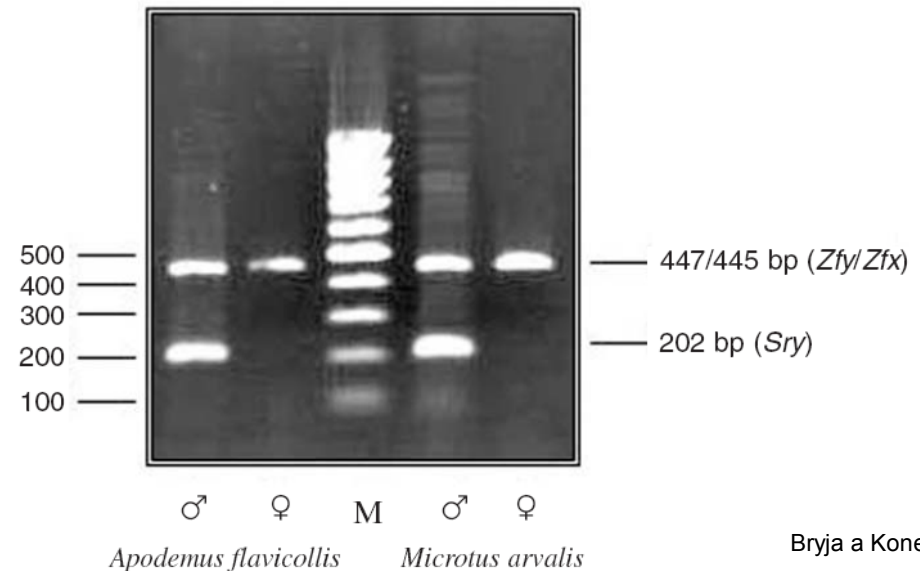
*Tokudaia osimensis*



© 奄美野生生物保護センター

# Určení pohlaví - savci

- Amplifikace genu na Chr Y (*Sry*)  
(nejlépe duplex PCR s genem na X nebo autosomech)

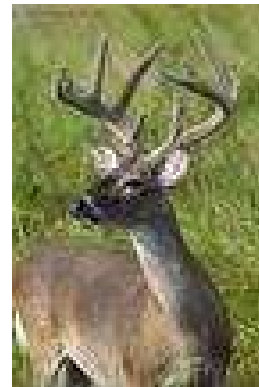


Bryja a Konečný 2003

- Analýzy z trusu: nutno používat druhově specifické markery  
(jinak cross-amplification s druhy tvořícími potravu)



**X**



Murphy et al. 2003

# Určení pohlaví – jiné skupiny

- Chr Y občas i u rostlin  
*Rumex*
- Plazi  
*Calotes versicolor*  
Sry má i 50% samic!
- Hledání markerů pomocí nespecifických metod (RAPD, AFLP)

