

# So close and so different: comparative phylogeography of two small mammal species, the Yellow-necked fieldmouse (*Apodemus flavicollis*) and the Woodmouse (*Apodemus sylvaticus*) in the Western Palearctic region

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In Europe, concordant geographical distribution among genetic lineages within different species is rare, which suggests distinct reactions to Quaternary ice ages. This study aims to determine whether such a discrepancy also affects a pair of sympatric species, which are morphologically and taxonomically closely related but which have slight differences in their ecological habits. The phylogeographic structures of two European rodents, the Yellow-necked fieldmouse (*A. flavicollis*) and the woodmouse (*Apodemus sylvaticus*) were, therefore, compared on the basis of mitochondrial DNA cytochrome *b* (mtDNA *cyt b*) sequences (965 base pairs) from 196 specimens collected from 59 European localities spread throughout the species distributions. The results indicate that the two species survived in different ways through the

Quaternary glaciations. *A. sylvaticus* survived in the Iberian Peninsula from where it recolonized almost all Europe at the end of the last glaciation. Conversely, the refuge from which *A. flavicollis* recolonized Europe, including northern Spain, during the Holocene corresponds to the Italo-Balkan area, where *A. sylvaticus* suffered a serious genetic bottleneck. This study confirms that even closely related species may have highly different phylogeographic histories and shows the importance of ecological plasticity of the species for their survival through climate change. Finally, it suggests that phylogeographic distinctiveness may be a general feature of European species.

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## Introduction

Phylogeography has become a powerful field of research revealing the principles and processes determining the geographic distribution of genealogical lineages, within and among closely related species (Avice, 2000). An extension of the phylogeography of single species is to compare patterns in multiple codistributed species. The main goal of this 'comparative phylogeography' has been to search for concordant geographical distribution among lineages within different species, which would indicate the influence of common historical factors. Recent comparative phylogeographic studies have found unanticipated biogeographic patterns, which suggest that vicariance has played a more important role in the appearance of modern biotic assemblages than current taxonomy would indicate (Arbogast and Kenagy, 2001).

Examining the mitochondrial DNA (mtDNA) polymorphism in various animal species, including fishes,

turtles, birds and invertebrates, Avice (2000) reports the existence of similar phylogeographic patterns in South-Eastern USA, suggesting the influence of common historical factors in structuring the geographical distribution of the genetic variability of these taxa. Concordance between phylogeographic patterns was also observed in other regions, such as central America, in several rodent species (ie Bermingham and Martin, 1998; Riddle *et al*, 2000; Sullivan *et al*, 2000), southern America in various bat, rodent and marsupial species (ie da Silva and Patton, 1993, 1998; Ditchfield, 2000) as well as in Australian birds, reptiles (Joseph *et al*, 1995) and snails (Hugall *et al*, 2002). Furthermore, Lessa *et al* (2003) detected a strong concordance of demographic expansion in North American mammals after the last ice age, 10 000 years ago. Finally, in contrast to previous paleontological studies (Graham *et al*, 1996), recent fossil analyses strongly suggest that many North American species shifted their ranges in similar ways during the Quaternary climate changes (Lyons, 2003).

However, some other examples show that such a concordance should not be considered a general trend, incongruences appearing in the phylogeographic structure of different species, namely among different plants

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and animals from north western North America (Brunsfeld *et al*, 2001) or among various small mammals in Southern America (Costa, 2003). The most striking situation is found in Europe. Indeed, although some general trends can be identified concerning refuge regions, postglacial recolonization routes or location of contact zones (Bilton *et al*, 1998; Taberlet *et al*, 1998; Hewitt, 2001), important differences are detected between phylogeographic patterns throughout Europe suggesting that the taxa responded differently to the Quaternary ice ages (Taberlet *et al*, 1998; Hewitt, 1999, 2001).

This important lack of congruence between the phylogeographic histories of European species is probably linked to the particular topography of Europe. Indeed, strong biogeographic barriers such as the Mediterranean and the Black Seas or the Pyrenees, the Alps and the Carpathians Mountains run across Europe in an east–west orientation. They probably limited the southward movements of fauna and flora during the ice ages and the northward expansion from the Mediterranean peninsulas during interglacial periods. Depending on their ecological ability to cross these barriers as well as to stochastic effects (ie important environment changes or diseases over very short periods, which probably led to the disappearance of many populations, even in the potential refuges), the European species probably reacted to the Quaternary glaciations and survived them in many different ways.

This study aims to determine whether such incongruence between the phylogeographic patterns also occurs in closely related sympatric species with similar behavior and morphology but with some differences in their ecological habits: (1) Did they survive in the same regions during the Quaternary glaciations? (2) Did they expand rapidly in northern European regions from these refuges at the end of the last glaciation (like North American mammals)? (3) Did they recolonize these northern regions by the same routes?

In other terms, do these two species have a similar phylogeographic structure and is it possible to discover the influence of common historical factors or, on the contrary, do they have independent phylogeographic histories as a consequence of different ecological habits?

In order to answer these questions, we compared the well-known phylogeographic structure of the woodmouse (*Apodemus sylvaticus*) (Michaux *et al*, 1996, 1998, 2003) to that of the yellow-necked fieldmouse (*Apodemus flavicollis*). These species appear to be an interesting model, because they are phylogenetically and taxonomically closely related (Michaux *et al*, 2002), both have been present in Europe for at least 3 My (Michaux and Pasquier, 1974), their ecological preferences are slightly different but overlap, they are sympatric and even syntopic over a great part of their distribution, and their phenotypes are often so similar that species discrimination in the field is often problematic (Niethammer, 1978; Michaux *et al*, 2001).

## Materials and methods

A total of 98 *A. flavicollis* coming from 39 localities sampled throughout their European range have been analyzed (Table 1 and Figure 1). These were compared

with 102 *A. sylvaticus* from 40 localities and already described in Michaux *et al* (2003).

DNA was extracted from ethanol-preserved tissue following Sambrook *et al* (1989). Tissues were taken from the *Apodemus* tissue collection of JR Michaux (JRM-numbers) and the mammal tissue collection housed at the Institut des Sciences de l'Evolution de Montpellier (Catzefflis, 1991; T-numbers).

A large portion of cytochrome *b* (965 bp) was amplified using the same primers and protocols previously used for *A. sylvaticus* (see Michaux *et al*, 2003).

## Sequence alignment

The published cytochrome *b* sequences for *A. flavicollis* ( $n=3$ : AB032853, AF159392 and AF127539), *A. sylvaticus* ( $n=2$ : AJ311148 and AJ311149), and *A. mystacinus* ( $n=2$ : AJ311146 and AJ311147) were downloaded from GenBank (see Table 1) and aligned to the new sequences using the ED editor (MUST package, Philippe (1993).

## Analyses

The aligned sequences were analyzed by distance (neighbor joining, NJ; Saitou and Nei, 1987) and maximum parsimony (MP) (Kluge and Farris, 1969) methods. We used PAUP 4.0b8 (Distance: ME criterion, Kimura's 2 parameter (K2P) and general time reversible (GTR) models; MP: heuristic search, TBR branch swapping option, Maxtrees = 1000 (Swofford, 1998)). The distance analysis was performed assuming a gamma distribution for substitution rates across sites, where the parameter alpha (Yang, 1996) and the proportion of invariant sites (I) was estimated by the maximum-likelihood method assuming the GTR phylogeny using the PAUP 4.0b8. The robustness of inferences was assessed by bootstrap resampling (BP) (Felsenstein, 1985) (1000 random repetitions). A minimum spanning network was constructed using the program TCS (Clement *et al*, 2000). Nucleotide diversity,  $\pi$ , was estimated using the DNASP program (Rozas and Rozas, 1997). A 'mismatch distribution' of substitutional differences between pairs of haplotypes was calculated within the main genetic lineages (clades A, B and C) and compared with a fit to the Poisson model using DNASP (Rozas and Rozas, 1997). This method can be used to assess the demographic history (patterns of population growth or stability) (Emerson *et al*, 2001) of the main clades. Moreover, a coalescent approach was also developed to validate the results revealed by the mismatch distributions (Emerson *et al*, 2001). This latter method is explained more extensively in Michaux *et al* (2003).

An approximate timing of divergence between the observed mtDNA lineages was calculated as explained in Michaux *et al* (2003). For this procedure, two calibration points derived from paleontological data were used for this analysis. First, the divergence time between the 'big' *A. mystacinus* and all the 'small' *Sylvaemus* (*A. flavicollis*, *A. sylvaticus*, *A. alpicola*, *A. uralensis*) was estimated from fossil data (cranial and dental characters) at approximately 7 My (Aguilar and Michaux, 1996; Michaux *et al*, 1997). Secondly, the same methods allowed an estimate of the divergence between *A. sylvaticus* and *A. flavicollis* at approximately 4 My (Michaux and Pasquier, 1974; Jacques Michaux, personal

**Table 1** Geographic distribution and references of *Apodemus* tissues used for the experiments

Geographic origin		Total no. of animals	Sample symbols (see Figures 1–3)	Tissue sample or GenBank accession numbers (for sequenced samples)	Haplotype number and name of clade (in bold) for each samples (see Figure 2)
<i>Apodemus flavicollis</i>					
Austria	Vorarlberg	3	A	JRM-203 to JRM-204	61 <b>C</b>
Belarus	Berezina Biosphere reserve	3	Bel	JRM-313 to JRM-315	8 <b>A</b> , 9 <b>A</b> , 57 <b>C</b>
Belgium	Ardennes	1	B	JRM-332	58 <b>C</b>
Czech Republic	Kašperské Nové Mlýny, Breclav (Bohemia)	4	CZ1	JRM-373, JRM-377, JRM-378, JRM-379	63 <b>C</b> , 64 <b>C</b> , 51 <b>C</b> , 40 <b>C</b>
		1	CZ2	JRM-255	33 <b>C</b>
Estonia	Tallin	2	Es	JRM-316, JRM348	7 <b>A</b> , 11 <b>A</b>
France	Allier	2	F1	JRM-343, JRM-344	27 <b>C</b> , 56 <b>C</b>
	Serandon (Corrèze)	2	F2	JRM-394, JRM-395	25 <b>C</b>
	Py Mantet (Pyrenées-Orientales)	5	F3	JRM-278 to 282	41 <b>C</b>
	Normandie	1	F4	JRM-1026	56 <b>C</b>
Germany	Bielefeld	1	G1	JRM-207	39 <b>C</b>
Greece	Macedonia region	2	Gr1	JRM-217, JRM-219	21 <b>C</b> , 29 <b>C</b>
	Peloponnesus	6	Gr2	JRM-216, JRM-481 to JRM-485	66 <b>C</b> , 72 <b>C</b> , 71 <b>C</b> , 77 <b>C</b> , 24 <b>C</b> , 3 <b>C</b>
	Epirus	2	Gr3	JRM-220, JRM-222	30 <b>C</b> , 35 <b>C</b>
	Mt Olympus	8	Gr4	JRM-460, JRM-701, JRM-704, JRM-707 to JRM-711	65 <b>C</b> , 76 <b>C</b> , 75 <b>C</b> , 79 <b>C</b> , 20 <b>C</b> , 34 <b>C</b> , 26 <b>C</b>
Hungary	Debrecen	1	Hu	JRM-307	41 <b>C</b>
Italy	Grosseto (Toscana)	1	I1	JRM-175	45 <b>C</b>
	Aspromonte (Calabria)	3	I2	JRM-192 to JRM-194	47 <b>C</b> , 50 <b>C</b> , 37 <b>C</b>
	Penne (Abruzzo)	2	I3	JRM-298, JRM-200	46 <b>C</b> , 49 <b>C</b>
Lituania	Unknown	1	Li	JRM-347	10 <b>C</b>
Macedonia	Mt Pellister	3	Ma1	JRM-213, JRM-214, JRM-492	31 <b>C</b>
	Bistra	4	Ma2	JRM-117 to JRM-120	32 <b>C</b> , 68 <b>C</b> , 55 <b>C</b>
Netherlands	Unknown	1	NL	ABO32853	52 <b>C</b>
Romania	Cheile Garlistei	2	R1	JRM-533, JRM-534	13 <b>B</b> , 17 <b>B</b>
	Piatra Craiului Mont (Natural Parc)	2	R2	JRM-572, JRM-573	74 <b>C</b> , 16 <b>B</b>
	Guarda	1	R3	JRM-532	36 <b>C</b>
	Saliste	1	R4	JRM-531	1 <b>A</b>
Russia	Samara	2	Ru1	JRM-405, JRM-408	5 <b>A</b> , 6 <b>A</b>
	Volgograd	2	Ru2	JRM-287, JRM-288	12 <b>B</b> , 15 <b>B</b>
	Voronezh	1	Ru3	JRM-309	14 <b>B</b>
Slovenia	Skofja	2	Sl1	JRM-418, JRM-420	22 <b>C</b> , 67 <b>C</b>
	Asan Cesma	1	Sl2	JRM-122	70 <b>C</b>
Spain	Eugi (Navarra)	10	E	JRM-124 to JRM-133	53 <b>C</b> , 54 <b>C</b> , 62 <b>C</b> , 60 <b>C</b> , 28 <b>C</b>
Sweden	Gotland	2	S1	JRM-445	38 <b>C</b> , 42 <b>C</b>
	Uppsala	2	S2	JRM-294, JRM-295	43 <b>C</b> , 4 <b>A</b> <b>C</b>
Switzerland	Geneva	1	Sw1	JRM-342	23 <b>C</b>
	Constanz	1	Sw2	AF159392	78 <b>C</b>
Turkey	Thrace	6	Tur	JRM-223, JRM-224, JRM-380 to JRM-382, JRM-235	48 <b>C</b> , 44 <b>C</b> , 59 <b>C</b> , 69 <b>C</b>
Ukraine	Southern Bug river	1	U	T-855	73 <b>C</b>
Yugoslavia	Susarra (Vojvodina)	3	Y	JRM-178 to JRM-180	18 <b>B</b> , 19 <b>B</b>
<i>Apodemus sylvaticus</i>					
102 specimens from 40 continental European localities (see Michaux <i>et al</i> , 2003)					
<i>Apodemus mystacinus</i>					
Syria		1		AJ311146	
Greece		1		AJ311147	

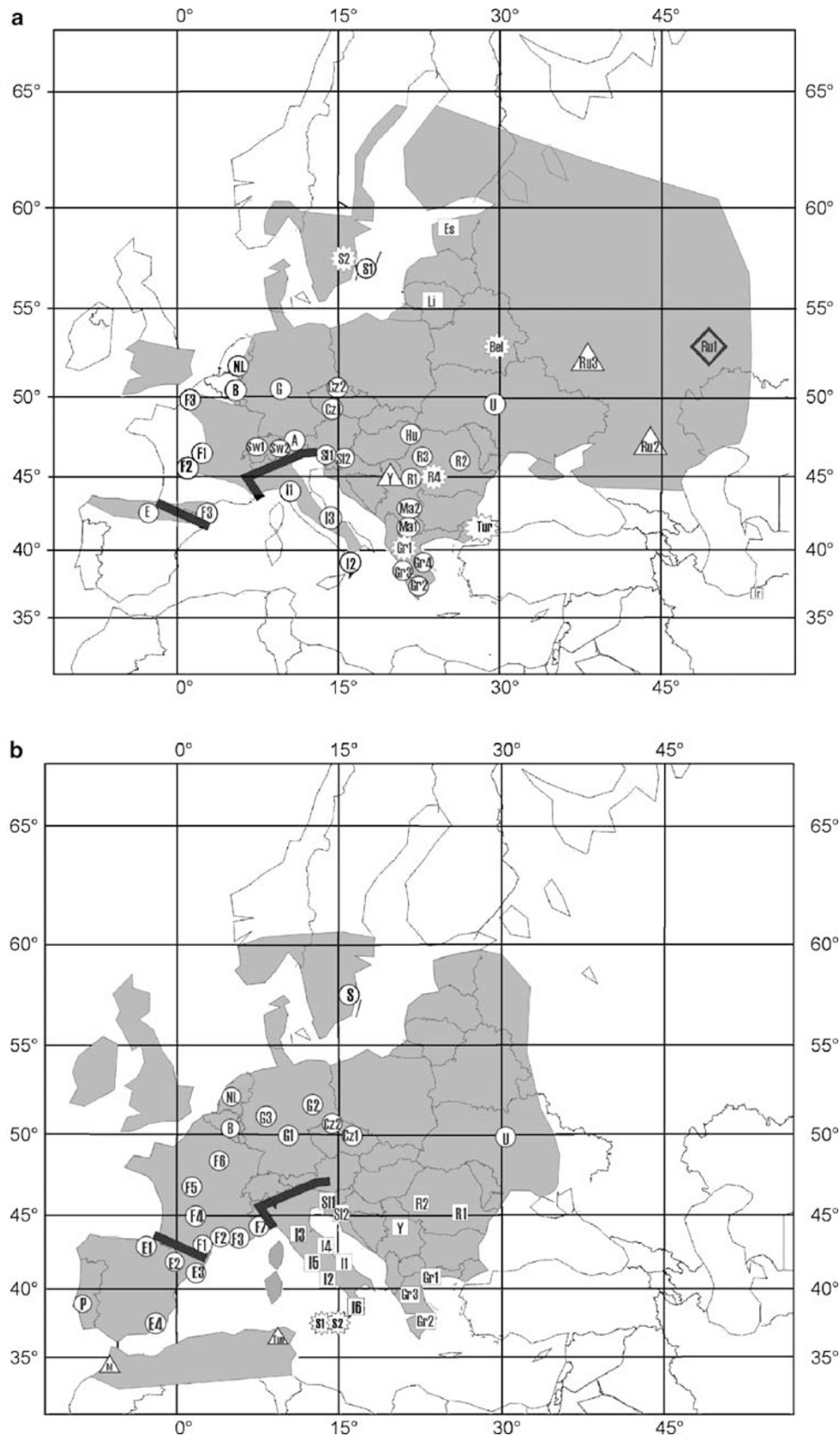
communication). Relative-rate tests were conducted according to Robinson *et al* (1998).

## Results

### Haplotype variation

We identified 79 haplotypes among the *A. flavicollis* cytochrome *b* sequences (GenBank Accession Numbers

AJ605600–AJ605665, AJ605672–AJ605675, AJ631968–AJ631970, AJ605686, AJ605687, AJ605691, AJ605692 + published sequences: ABO32853, AF159392). The complete data matrix comprised these 79 haplotypes as well as two *A. sylvaticus* and two *A. mystacinus* sequences used as outgroups. This matrix provided 965 bp of which 298 sites (31%) were variable and 189 (19%) were parsimony informative, respectively. The mean transition to transversion ratio was 3.49. The nucleotide frequencies



**Figure 1** Geographical distribution of the *Apodemus flavicollis* (a) and *A. sylvaticus* (b) samples. The shaded zones correspond to the distribution area of each species in Europe respectively (as described by Schilling *et al*, 1986; Mezhzherin, 1997; Zagorodnyuk *et al*, 1997). Thick lines correspond to the main biogeographic barriers: the Pyrenees and the Alps. In (a), squares, triangles and circles correspond to clades A, B and C, respectively. White stars and diamond shape correspond to localities where haplotypes from the clades A and C or from B and C are mixed, respectively. See Table 1 for locality symbols. In (b), squares, triangles, white stars and circles represent localities where the clades 1a and 1b, 2a and 2b were found, respectively (see Michaux *et al*, 2003).

were 0.272, 0.290, 0.314 and 0.123 for C, T, A and G, respectively.

### Phylogenetic analyses

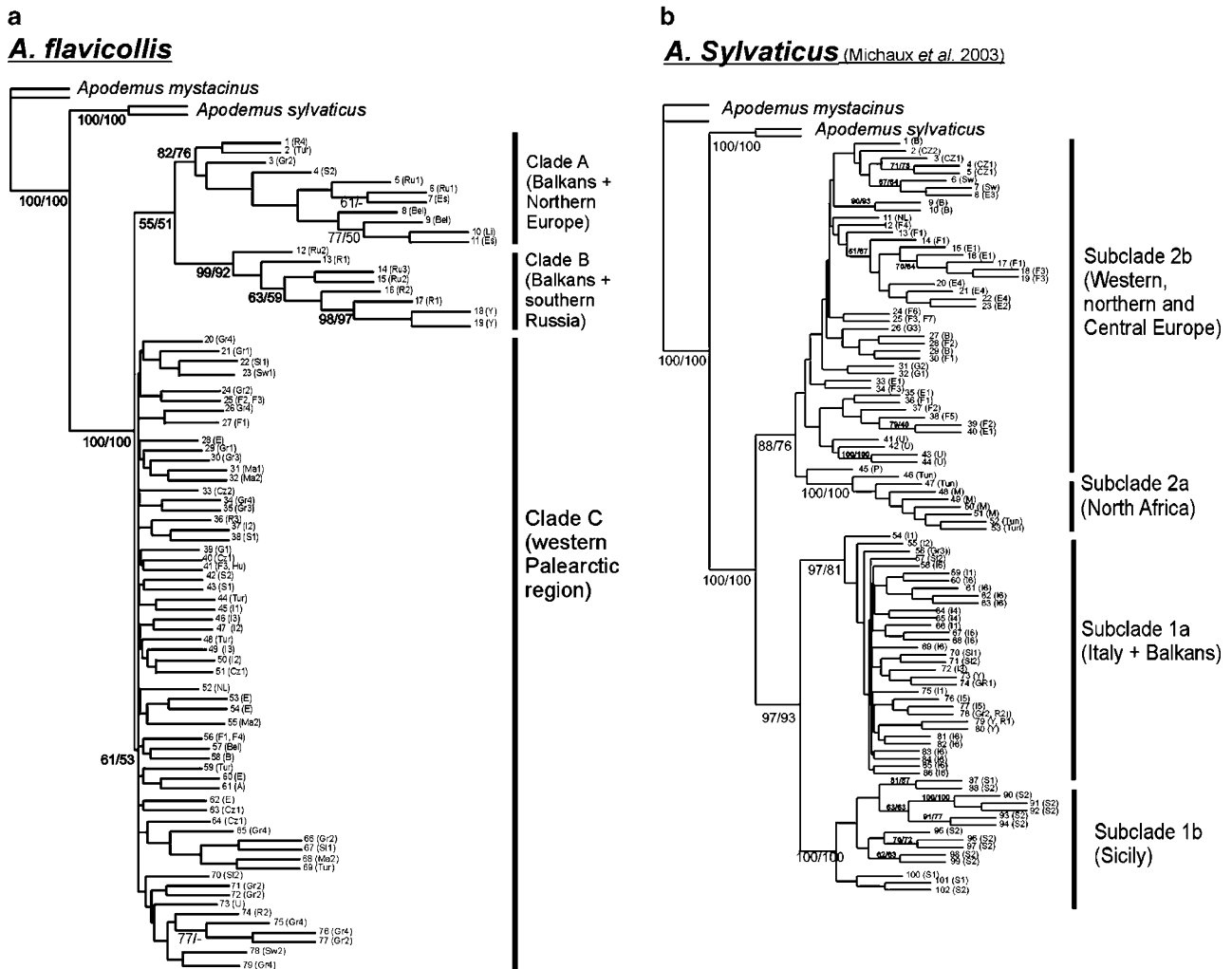
The *A. flavicollis* NJ tree (Figure 2a) is characterized by a weak genetic and geographic structure. However, three clades can be observed: the first one (clade A) is well supported (BP of 82%) and characterized by specimens from northern Europe (Russia, Estonia, Lithuania, Belarus, Sweden) and the Balkan region (Greece, Romania and European Turkey). The second one (B) (BP: 99%) corresponds to animals from southern Russia, Romania and Yugoslavia and appears more closely related to the first lineage than to the third one, which is less well supported (BP: 61%) and comprises individuals from the whole Western Palearctic region including Western (France, Belgium), Northern (Sweden) and Eastern Europe up to Ukraine as well as the three Mediterranean peninsulas (Iberian, Apennine and Balkan). The genetic divergence between these clades is low, varying between 1.3 and 1.9% K2P distance. The MP analyses yielded one most parsimonious tree for each

species ( $L=472$  and 678 steps;  $CI=0.65$  and 0.58;  $RI=0.90$  and 0.82, respectively) identical to the NJ tree.

The network calculated using the TCS program (Clement *et al*, 2000) (Figure 3a) show similar results: almost all the haplotypes are connected together in a huge star-like topology (subclade C), whereas the clade A is separated from this group by three mutational steps. Between the clades A and B, there are nine further mutational steps.

### Nucleotide diversity and genetic structure

To assess whether genetic diversity was higher within the potential refuge regions as compared to northern populations, the samples were separated into different subgroups corresponding to these regions (Italy + Balkans, Iberian Peninsula + southern France, Western + Central Europe + Northern Europe, Ukraine + Russia) (Figure 1), then nucleotide and haplotype diversities were calculated for every group (see Table 2). The Italian and Balkan populations were combined as they were not genetically differentiated (mean K2P distance between the Balkan and Italian populations versus within the Balkan group = 1.0/1.2%). The results



**Figure 2** Neighbor-joining tree for the 79 *A. flavicollis* (a) and 102 *A. sylvaticus* (b) mtDNA haplotypes (labelled from 1 to 79 and 1 to 102). Numbers above the branches correspond to bootstrap support above 50% obtained in the NJ and MP analyses, respectively. Geographic origins (see Table 1 and Figure 1) are shown in parentheses.



**Table 2** Genetic variability observed within the potential refuge as well as in northern populations of *A. flavicollis*

	Sample size	Number of haplotypes	Genetic divergence within each clade (% K2P distance)	Nucleotide diversity ( $\pi$ )	SD	Haplotype diversity ( $h$ )	SD
<i>A. flavicollis</i>							
Italy+Balkans	49	43	1.20	0.0150	0.00110	0.988	0.007
Iberian peninsula+southern France	15	7	0.50	0.0054	0.00110	0.986	0.096
Western+Central+Northern Europe	27	25	0.78	0.0075	0.00111	0.992	0.026
Ukraine+Russia	7	6	1.11	0.0092	0.00140	0.952	0.096
<i>A. sylvaticus</i>							
Italy+Balkans (= subclade 1a, in Michaux <i>et al</i> , 2003)	26	25	1.22	0.0064	0.00080	0.968	0.022
Iberian peninsula+southern France (= Subclade 2b1)	27	23	1.41	0.0087	0.00070	0.981	0.023
Western+Central+Northern Europe (= Subclade 2b2)	26	20	1.09	0.0052	0.00100	0.965	0.028

Samples were pooled according to their geographic origin (Table 1). For comparison, the data obtained in *A. sylvaticus* (Michaux *et al*, 2003) are also given.

indicate that the Italo-Balkan region (Figure 1) has the highest nucleotide diversity ( $P < 0.05$ ,  $t$ -test) as compared to the populations from all the other regions. The animals from southwestern Europe were the least genetically diversified. Conversely, the Russian population is characterized by a higher level of nucleotide and haplotypes diversities, corresponding to the presence of two different clades (A and B) occurring syntopically in the region. For comparison, data obtained on *A. sylvaticus* (Michaux *et al*, 2003) are also included in Table 2.

A signature of population growth (ie a bell-shaped distribution) is clearly evident in the distribution of pairwise distribution within clade C (Figure 4). Moreover, the coalescence analysis rejects the null hypothesis of a stable population for this clade (Table 3). On the other hand, the clades A and B are bimodal suggesting constant population size for these groups. This is corroborated by the coalescence test that did not reject the constant population model. However, this result must be treated cautiously as the sample size for these groups is very low (seven individuals for each).

A population within a given area is the actual (resident) population. Thus, to test expansion of an actual population, it is obligatory to include its individuals, irrespective of genetic clades. Following this idea, we performed other mismatch distribution analyses considering different geographic populations of *A. flavicollis* (Italy + Balkans, Iberian Peninsula + southern France, Western + Central Europe, Northern Europe, Ukraine + Russia) and not genetic clades. The results (data not shown) evidence that the Italian-Balkan population is characterized by a bimodal curve suggesting constant population size, whereas the Western + Central European population is characterized by a bell-shaped curve (signature of population growth). More heterogeneous distribution appears in the other populations suggesting that they are more stable. However, this result was obtained on a very small sampling (maximum seven individuals for each population) and should be, again, considered carefully.

#### Divergence time

The relative rate test indicates no significant rate heterogeneity (both  $K_s$  and  $K_a$ ) for cytochrome *b* between

the different *A. flavicollis* clades and the *Apodemus* specimens.

The mean K2P distance between *A. mystacinus* and *A. flavicollis*/*A. sylvaticus* that diverged 7 My ago is 18.2%. The distance between *A. flavicollis* and *A. sylvaticus* that diverged 4 My ago is 11.4%. These values give a rate of 2.6–2.85% K2P distance per million years. Therefore, taking into account the correction for ancestral mtDNA polymorphism, the separation time between clades A–B and C and between clades A and B took respectively 0.6–0.5 and 0.4 My ago. For comparison, the separation time between clades 1 and 2 of *A. sylvaticus* dates back to 1.5–1.6 My (Michaux *et al*, 2003).

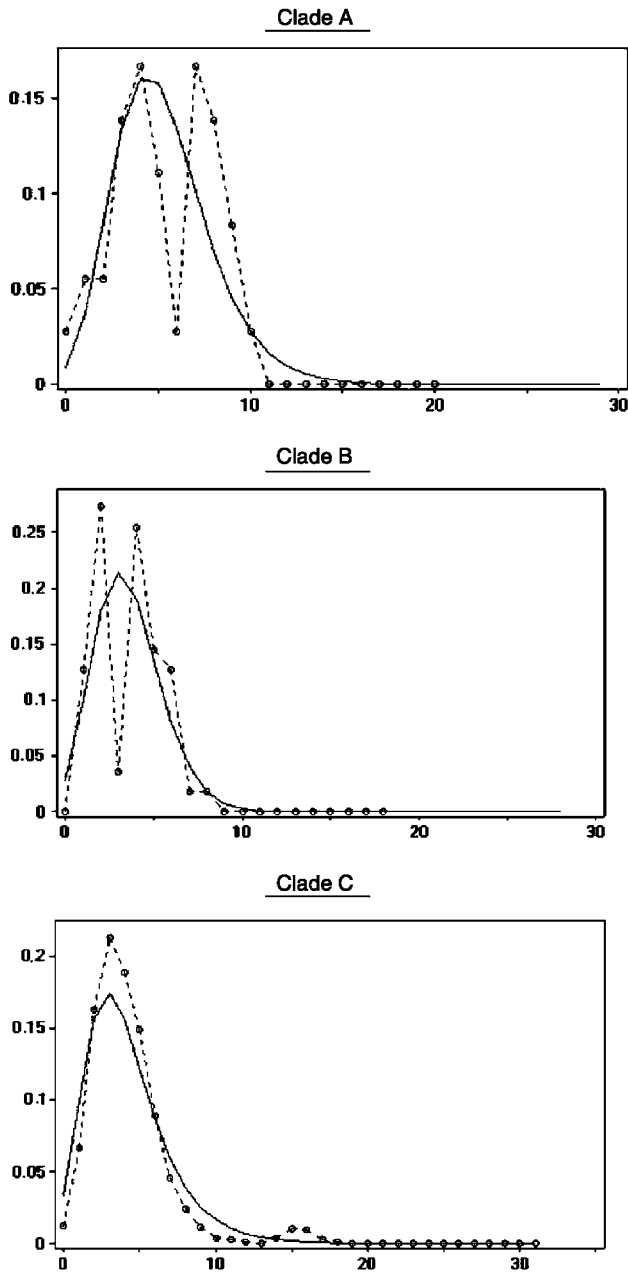
## Discussion

### Comparison of the genetic structure of *A. sylvaticus* and *A. flavicollis*

The distance, maximum parsimony and network analyses clearly show that the two *Apodemus* species have distinct genetic structure (Figure 2 and 3).

The *A. flavicollis* populations are split into three main genetic lineages separated by a low level of nucleotide divergence (between 1.3 and 1.9% K2P distance) and overlapping in a part of their geographical distribution in the Balkan region. The divergence time estimated on the basis of two paleontological calibration points (Aguilar and Michaux, 1996; Michaux *et al*, 1997) indicates a separation time of 0.6–0.5 My between the ancestral haplotypes that have led to these three lineages. Therefore, it appears that the intraspecific structure of *A. flavicollis* developed during this period. This result is consistent with the hypotheses of Avise (2000), Hewitt (2001) and Seddon *et al* (2001) who postulated that the majority of the intraspecific differentiation in vertebrates occurred during the Quaternary.

The European continental *A. sylvaticus* populations are split into two main genetic lineages nonoverlapping in their geographical distributions and separated by a high level of genetic divergence (5.4% of K2P genetic distance) (Figures 2b and 3b; Michaux *et al*, 2003): the first is widespread from southern Spain in the south to Sweden in the north and Ukraine in the East; the second one is limited to Italy and the Balkans (Figure 1b). As already



**Figure 4** Mismatch distribution for mtDNA types from the three major *A. flavicollis* genetic clades observed in Europe. The expected frequency is based on a population growth-decline model (for the clades A, B and C, respectively:  $\theta$  initial = 0, 2.4 and 0,  $\theta$  final = 1000,  $\tau$  = 4.4, 9.2 and 2.9), determined using the DNASP v3.5 program (Rozas and Rozas, 1997) and is represented by a continuous line. The observed frequency is represented by a dotted line.

proposed for several other European species (Taberlet *et al*, 1998; Hewitt, 2001; Seddon *et al*, 2001), this structure could be explained by the isolation of two woodmice groups in two different refugia (Iberian peninsula and Italy–Balkans) during one of the first Quaternary glacial periods. This hypothesis is corroborated by the molecular clock analysis that estimates the separation time between the ancestral haplotypes at 1.5–1.6 My ago. This isolation was probably reinforced by the presence of the Alps that played and probably continue to play the role

**Table 3** Results of the coalescence analysis on the three main genetic clades of *A. flavicollis*

Subclades	Log-likelihood			
	Stable population model	Exponential change model	$\chi^2$	P
Balkans+Northern Europe (A)	0.0141	1.0592	2.090	0.148
Balkans+southern Russia (B)	0.0026	0.1117	0.218	0.640
Western Palearctic region (C)	−0.2444	1.8731	4.235	0.040

of a biogeographic barrier for the woodmouse (Michaux *et al*, 1996). These two factors likely have prevented genetic exchanges between both groups for a long time and have led to the strong genetic differences observed within this species. In contrast, the frequent drops in the level of the Adriatic Sea during the Quaternary ice periods (Strahler and Strahler, 1989) apparently allowed the Italian and Balkan woodmice populations to remain in contact. This could also explain the great similarity observed between the *A. flavicollis* from Italy and the Balkans.

#### Refuge regions and postglacial recolonizations

As predicted by the expansion/contraction model (Nichols and Hewitt, 1994; Santucci *et al*, 1998), the analysis of nucleotide diversity (Table 2) indicates that the Italo-Balkan region was a refuge for *A. flavicollis*. Indeed, these populations are characterized by a genetic diversity significantly higher ( $\pi = 0.015$ ,  $P < 0.05$ ) than in northern, eastern or western populations. This result is also observed for each of the three genetic lineages analyzed separately and is corroborated by the mismatch distribution analysis performed on different geographic populations of *A. flavicollis*. Indeed, this analysis suggests the existence of a stable population in the Italo-Balkan region. The three genetic lineages probably survive to the last Quaternary glaciations in three independent Balkan refuges where they diverged separately. They invaded later the entire Western Palearctic region during interglacial periods. These results rather correspond to the hypothesis of Hewitt (1999) and Taberlet *et al* (1998) which proposed a South to North expansion of the European species, in contrast to the hypothesis of Bilton *et al* (1998), suggesting a colonization of Europe from Eastern regions (Russia, Asia). Indeed, in this last case, the Eastern populations of *A. flavicollis* would be characterized by a higher level of genetic diversity as compared to the Balkan ones. However, despite their low number (seven individuals), the Russian–Ukrainian fieldmice are characterized by a relatively high level of nucleotide diversity and genetic divergence as well as by an important number of haplotypes (6). As high genetic diversity also arises when diverse clades occur syntopically, this result would be explained by the presence of two *A. flavicollis* clades (A and B) in this region. Nevertheless, without a larger sample for this region, the hypothesis of another Eastern refuge cannot be precluded.

Conversely, the population of the southwestern potential refuge (Iberian Peninsula and southern France) has the lowest level of nucleotide diversity and, in contrast to



*A. sylvaticus*, does not possess specific haplotypes absent from other regions. This can be explained by a recent colonization of this region from the Balkan refuge population. Indeed, as proposed by Hewitt (1999) and Avise (2000), a rapid expansion from refugial populations involves serial bottlenecks with progressive loss of allelic diversity resulting in less genetic diversity among populations living in the more recently colonized places. This hypothesis is corroborated by the intermediate level of nucleotide diversity characterizing the populations from Central and Western Europe as well as from Northern Europe.

Therefore, it seems that: (i) the Iberian peninsula did not constitute a refuge for this species during the last glaciation. This is corroborated by paleontological data, which attest the presence of *A. flavicollis* in this region only from the end of the Lower to the beginning of the Upper Pleistocene (Sese, 1994); (ii) *A. flavicollis* did not survive up to the last glaciation in Western, Eastern and Northern Europe. This hypothesis is also confirmed by paleontological data (Cordy, 1984; Aguilar and Michaux, unpublished data); (iii) Three genetic lineages of the Yellow-necked fieldmouse surviving in the Balkan region recolonized the main part of the Western Palearctic region after the last ice age, 10 000 years ago. This scenario of expansion is corroborated by a bell-shaped curve in the mismatch distribution analysis at least for the main genetic lineage (C) (Figure 4) (Luikart *et al*, 2001), the results from the coalescent analysis as well as from allozyme data (Filippucci *et al*, 2002). Moreover, the star-like topology characterizing the main genetic lineage (clade C) suggests that this expansion was probably very rapid and explosive for this group, which recolonized almost all Europe as compared to the other lineages, and took place after a bottleneck, which probably lowered its ancestral genetic diversity (Avise, 2000).

For *A. sylvaticus*, the results strongly suggest that southwestern Europe (the Iberian Peninsula and southern France) was a refuge region (Michaux *et al*, 2003). Indeed, these populations are characterized by a genetic diversity significantly higher ( $\pi = 0.0087$ ,  $P < 0.05$ ) than in the northern populations (Table 2). Therefore, it is probable that woodmice recolonized and expanded throughout the main part of the Western Palearctic region from southwestern Europe at the end of the last ice age. This scenario of expansion is supported by a bell-shaped aspect of the curve obtained in the mismatch distribution analysis for the main European lineage (Luikart *et al*, 2001; Michaux *et al*, 2003). The Pyrenees were therefore not an effective barrier to the northward expansion of the woodmouse (Michaux *et al*, 1998). In contrast to the southwestern group, the Italo-Balkan populations are characterized by a lower genetic diversity ( $\pi = 0.0052$ ,  $P < 0.05$ ). This observation is somewhat surprising as paleontological (Aguilar and Michaux, unpublished data) and paleoclimatological (Tzedakis *et al*, 1997) data attest that Italy and the Balkans were a refuge region for *A. sylvaticus* during the Quaternary glaciations. This low genetic variability could be explained by a genetic bottleneck that appeared during one of the last ice ages. This hypothesis is corroborated by three results: the very short branch lengths between haplotypes within this group in the distance analysis (Figure 2b); the star-like topology in the minimum spanning network (Figure 3b) suggesting a rapid expansion

from a relative small number of founder animals and a low value of haplotype ( $h$ ) and nucleotide ( $\pi$ ) diversity (Table 2) (Avise, 2000). The factors that could explain this genetic bottleneck are discussed in details in Michaux *et al* (2003). It could also explain why the Italo-Balkan populations did not contribute to the postglacial colonization of the northern regions, as expansion could only have occurred after population growth. During this time, the western group rapidly expanded towards Northern and Central Europe, thereby preventing a northern colonisation by the Italo-Balkan animals. Indeed, once established, resident rodents often aggressively exclude newcomers (Granjon and Cheylan, 1989). Moreover, the presence of topographic obstacles such as the Alps might slow down a postglacial expansion of the Italo-Balkan group.

This study also evidenced that Sicily is a refuge for *A. sylvaticus* and that this region is characterized by a genetically well-differentiated and stable population, which is separated from the Italian animals since 0.8/0.9 Myr (for more details, see Michaux *et al*, 2003). Moreover, this species invaded North Africa recently, probably via the Gibraltar Strait and anthropogenic introductions (Libois *et al*, 2001; Michaux *et al*, 2003). This wider distribution range of the woodmouse in Western Europe and North Africa is probably associated to its greater ecological plasticity as compared to *A. flavicollis*, enabling it to colonize regions covered by habitats other than forests (eg Mediterranean scrub and open fields with small groves).

## Conclusions

Although *A. sylvaticus* and *A. flavicollis* are closely related from a taxonomical point of view and that they have very similar morphology and distribution, they reacted and survived to the Quaternary glaciations in totally different ways. Indeed, from paleontological and genetic data, *A. flavicollis* did not survive, at least during the last glaciation (22 000–16 000 years ago), in the Iberian Peninsula, whereas *A. sylvaticus* persisted and recolonized almost all Europe from there at the end of the last glaciation. Conversely, the refuge region from which *A. flavicollis* recolonized Europe during the Holocene corresponds to the Balkan area where *A. sylvaticus* seems to have suffered a severe genetic bottleneck. This result corresponds to the Gleasonian model (Graham *et al*, 1996), which predicts that species respond in an individualistic manner to climate changes, and expand and contract their ranges independently from each other.

The differences between the two *Apodemus* species or could be explained by stochastic events or interspecific competition: As these species often live in the same habitats, high interspecific competition during the Quaternary ice ages may have played a role in the observed phylogeographic patterns. However, behavioral studies (Hoffmeyer, 1973; Montgomery, 1978, 1985; J Cassaing, unpublished data) showed that the overall level of interaction is weak and neither species can exclude the other. Moreover, in interspecific contests, *A. sylvaticus* is always dominated by *A. flavicollis*. Therefore, this hypothesis would better explain the decrease of the genetic variability of the Italo-Balkan *A. sylvaticus* than the disappearance of *A. flavicollis* in the Iberian region. Another explanation could be related to the evolutionary

history of these species. Indeed, according to the paleontological (Michaux and Pasquier, 1974) and molecular (Michaux *et al*, 2002) data, the differentiation between *A. flavicollis* and *A. sylvaticus* is probably the result of an allopatric speciation: the ancestors of *A. sylvaticus* were isolated in Spain and southern France, whereas those of *A. flavicollis* rather lived in Central Europe and in the Balkans. Each species could be more adapted to the environment of its ancestral region and, therefore, have had more chances of surviving there. However, as the differentiation between the two species is very old (4My), both had time to establish across the European regions during the Quaternary (Aguilar and Michaux, unpublished data) and they probably had time enough to adapt themselves to the different European environments. Moreover, the differences between Western and Eastern European environments are probably not sufficient (Blondel, 1996) to support this hypothesis, particularly for such species characterized by strong ecological adaptability (Renaud and Michaux, 2003).

Several ecological (eg Kaustuv *et al*, 2001; Holt, 2003) and paleontological (Barnosky *et al*, 2001) studies have suggested the importance of ecological factors in the changes of species ranges, particularly during climate variation. *A. sylvaticus* is characterized by a greater ecological plasticity than *A. flavicollis* which is more associated with forested habitat with a high canopy cover (Marsh and Harris, 2000). Thus, the fragmentation of forests during the last glaciation in the Iberian peninsula (Blondel, 1996) may have played a role in its extinction in this region. Moreover, every climate change involves important habitat reorganizations and new biological interactions (Graham *et al*, 1996). Therefore, a more adaptable species like *A. sylvaticus* was probably more resistant to these factors than *A. flavicollis*. Finally, *A. flavicollis* shows poorer winter survival than *A. sylvaticus* (Montgomery, 1980; Marsh and Harris, 2000). Therefore, the harsher climatic conditions that characterized the Iberian peninsula during the last glaciation as compared to the Balkan region (Tzedakis *et al*, 1997) could also explain the extinction of the Yellow-necked fieldmouse in Western Europe during this period.

The time of their respective intraspecific differentiation can be compared under the assumption that cytochrome b gene evolves at the same rate in the two species (Michaux *et al*, 2002) and that their ancestral effective population size was the average of the current population sizes (Edwards and Beerli, 2000). *A. sylvaticus* is then estimated to have diverged during Early and Middle Pleistocene (1.5–1.6My ago for the two continental lineages) (0.8–0.9My between the Sicilian and Italian lineages) whereas *A. flavicollis* diverged more recently (0.4–0.6My ago). This result could be explained by the disappearance of ancient *A. flavicollis* lineages in southwestern Europe during the Early Quaternary ice ages, which is corroborated by paleontological data (Sese, 1994). In contrast, the better ecological plasticity of the woodmouse probably enabled it to better survive the Quaternary glaciations, even if its populations were periodically fragmented leading to different and deep genetic lineages.

In conclusion, the present results confirm that different species, even if they are closely related, can have strongly different phylogeographic histories and suggest that the phylogeographic exception may be a general feature of

European species. This shows the importance of ecological plasticity of the species, for their survival to climate changes. Finally, the present results evidence that the search of the influence of common historical factors to explain the phylogeographic structure of the European species turns out to be very complex.

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