


Spatial mixing of mitochondrial lineages and greater genetic diversity in some invasive populations of the American mink (*Neovison vison*) compared to native populations

Karla García  · Yolanda Melero · Santiago Palazón · Joaquim Gosálbez · Jose Castresana

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Abstract The genetic characteristics of introduced populations have a relevant impact on their ability to establish and spread. The American mink (*Neovison vison*), native to North America, is an important invasive species in the Iberian Peninsula. Here, we used mitochondrial DNA sequences data to investigate the genetic diversity and phylogeographic structure of invasive versus native populations of this species. We also evaluated whether genetic diversity in invasive populations could be explained by the genetic characteristics of the native sources from which they derived. Phylogenetic analysis revealed two major

lineages in the native range, which indicated a clear separation between western and eastern populations. On the contrary, we found no evidence of genetic structure in the invasive range. This was probably the result of the diverse origins of the released specimens and the rapid expansion and encounters of the introduced populations. We detected spatial mixing of both North American lineages in several sampling localities of the north central area of the Iberian Peninsula, giving rise to high levels of genetic diversity in some areas compared to North American populations. This could potentially lead to higher fitness of these individuals and thus increase the population viability and invasiveness of this species. These results point to the need to better study the populations in which lineages mix and, if necessary, intensify control efforts in them.

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K. García (✉) · S. Palazón · J. Gosálbez
Department of Evolutionary Biology, Ecology and Environmental Sciences, University of Barcelona, 08028 Barcelona, Spain
e-mail: karla.garcia.cl@gmail.com

Y. Melero
CREAF, 08193 Cerdanyola del Vallès, Spain

S. Palazón
Fauna and Flora Service, Department of Territory and Sustainability, Government of Catalonia, 08017 Barcelona, Spain

J. Castresana
Institute of Evolutionary Biology (CSIC-Universitat Pompeu Fabra), 08003 Barcelona, Spain

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Introduction

Identifying the biological attributes of successful invaders is among the most pressing questions yet to be answered (Sol et al. 2012). Although there is evidence that species differ in their invasion potential, controversy exists about the characteristics that differentiate successful and unsuccessful invaders

(Blackburn et al. 2009). Other studies of introduced species show that genetics might play an integral role in the success of an invasion, and they suggest that this role is determined by the quality of the genetic variation introduced, rather than the total quantity of genetic variation (see Dlugosch et al. 2015). The genetic diversity of founding populations, along with potential increase in genetic diversity from subsequent introductions, may be important to the successful establishment and spread of introduced species (Suarez and Tsutsui 2008; Sol et al. 2012). Nevertheless, there is no consensus on whether invaders stemming from multiple native origins are more successful than those from single populations, or whether demographic bottlenecks may limit the invasion success of a species (Edelaar et al. 2015; Estoup et al. 2016).

Overall, introduced populations lose genetic variation relative to their source populations (Dlugosch and Parker 2008). However, numerous introduction events and a greater number of source populations may buffer against such genetic losses (Kolbe et al. 2004; Dlugosch and Parker 2008; Uller and Leimu 2011). In fact, in some cases successful invaders show large increases in genetic diversity in the introduced range, apparently caused by mixing of populations from different source regions (Genton et al. 2005; Novak and Mack 1993). Therefore, it is crucial to obtain information on the phylogeographic structure and genetic diversity throughout the native and introduced ranges to understand the factors affecting genetic diversity during invasion (e.g. multiple introduction events or admixture levels among populations) and the interaction between genetic diversity and invasive potential (Edelaar et al. 2015).

The American mink (*Neovison vison*) is a successful invader with a native range restricted to North America and with invasive populations distributed across Europe, Asia and South America (Bonesi and Palazón 2007; Dunstone 1993). In Europe, this species was introduced for commercial fur farming during the twentieth century. Populations established by escaped individuals are now naturalised in fresh water and coastal ecosystems (Bonesi and Palazón 2007), with detrimental consequences for the abundance of many native species of prey and competitors of conservation importance (Bonesi et al. 2004; Harrington et al. 2009; Macdonald and Harrington 2003; Melero et al. 2012a), including critically endangered species such as the

European mink (*Mustela lutreola*) (Maran et al. 2016) and the Pyrenean desman (*Galemys pyrenaicus*) (Fernandes et al. 2008). Fur farms were established in many countries and harboured mink stocks from different parts of their native range. Because of the large scale and long-term nature of this farming, the number of escapees, which started to breed in the wild and became the founder individuals of feral populations, was very high (Bonesi and Palazón 2007). In Spain, mink farming started at the end of the 1950s and the number of farms reached a peak in the 1980s with about 400 farms concentrated mainly in the North West (Galicia) (Ruiz-Olmo et al. 1997). Today, feral mink populations are widely distributed across almost all of the north and centre of the Iberian Peninsula. During the last few decades, these populations experienced a rapid expansion and increased the number of nuclei in almost every colonised area (Ruiz-Olmo et al. 1997; Santulli 2014). Although the number of mink farms in Spain was considerably reduced at the end of the 1980s, mink escapes from the last active farms are still ongoing (Bonesi and Palazón 2007).

In Europe, there is genetic information on feral American mink populations from several countries. Previous studies in Spain using microsatellite data determined that American mink populations from disjunct geographical areas appeared differentiated and not yet connected by gene flow (Lecis et al. 2008). In Scotland, Zalewski et al. (2009) and Fraser et al. (2013) found that landscape features restricted gene flow within this species. Other studies in Poland found that multiple introductions determine the genetic structure of American mink populations (Zalewski et al. 2010) and that these populations exhibit high mitochondrial genetic diversity (Zalewski et al. 2011). A study in France using microsatellite data suggested recent admixture between populations that had recently encountered each other (Bifolchi et al. 2010). Similarly, in Sweden it was found, also using microsatellite data, that genetic diversity decreased over six consecutive years of mink culling as part of a population control programme (Zalewski et al. 2016).

Despite the importance of understanding the genetic diversity within the native populations for investigating the evolutionary dynamics of invasions, no phylogeographic study has been performed so far in the North American native range of the American mink. Therefore, many genetic studies performed thus far are difficult to interpret. We used mitochondrial

DNA (mtDNA) sequence data from populations in the native range to elucidate whether invasive populations in the Iberian Peninsula have single or multiple origins from the North American sources. We aimed to determine the phylogeographic structure of native and invasive populations, and to evaluate whether mitochondrial genetic diversity observed in established invasive populations can be explained by genetic characteristics of native populations. We examined whether the genetic variability of the Iberian population represented only a small part of that in its native range, as expected for many invasive species or, due to the introduction history and particular origin of these populations, this genetic variability was larger. Overall, the study aims to provide recommendations for mink control in the Iberian Peninsula with respect to determining key control areas.

Materials and methods

Sample collection

We used hair and tissue samples from invasive populations provided by different Autonomous Communities of Spain. These samples were collected during population control operations conducted by the local administrations between 2007 and 2016 as part of eradication programs and were conserved in tubes containing absolute ethanol (ESM Appendix: Table S1). A total of 105 samples from Catalonia (16), La Rioja (15), Galicia (12), Basque Country (20), Navarra (11), Castellon (7) and Castilla-Leon (24) were included in the study.

Moreover, tissue samples of 21 specimens from North America (ESM Appendix: Table S2) were obtained from different biological collections: *University of Alaska Museum* (12), *Museum of Southwestern Biology* (5) and *Cornell University Museum of Vertebrates* (4).

DNA extraction, PCR and sequencing of mitochondrial DNA

We extracted DNA using the QIAGEN DNeasy Blood and Tissue Kit, following the manufacturer's instructions, in a final elution volume of 100 µl. These extractions were carried out in a separated UV-irradiated area with dedicated equipment. Samples

were incubated in a lysis buffer with proteinase K at 56 °C overnight to ensure maximum tissue lysis.

For each sample, we amplified a 573 bp fragment of the 5' distal part of the mitochondrial DNA control region (D-loop). For this purpose, we designed novel primers using complete mitochondrial sequences of several mustelids: *Neovison_tRNA_pro* (5'-ATC AGC ACC CAA AGC TGA CAT TCT A-3') and *Neovison_Dloop_556r* (5'-TGT GTA TGT CCT GTG ACC ATT GAC T-3'). PCR reactions were set up in a dedicated PCR clean room that is physically separated from post-PCR working areas and regularly decontaminated by UV-irradiation. PCR reactions were performed in a final volume of 25 µl, containing 2–4 µl of genomic DNA, 1 µM of each primer, 0.2 mM dNTPs, 0.75 U of Promega GoTaq DNA polymerase and 17.5 µM of bovine serum albumin under the following conditions: an initial denaturation of 2 min at 95 °C, followed by 35 cycles of denaturation (30 s at 95 °C), annealing (30 s at 54 °C) and extension (30 s at 72 °C). PCR products were visualised by electrophoresis in a 1% agarose SYBR-Safe (Invitrogen) stained gel.

PCR products were purified using ExoSAP-It (Affymetrix), and both strands were sequenced using the original PCR primers at Macrogen Europe (Amsterdam, The Netherlands). Sequences were inspected, trimmed and assembled using Geneious Pro (Biomatters Ltd.). All sequences obtained in this study were deposited in European Nucleotide Archive/GenBank under accession numbers LT854061-LT854186.

Phylogenetic and genetic diversity analyses

Consensus sequences were aligned using MAFFT version 7.130 (Katoh et al. 2002) with default settings. Due to the presence of several gaps and ambiguous positions in a very specific region of the alignment, we manually removed positions 92–110, thus leaving 554 bp in the alignment.

A maximum-likelihood phylogenetic tree was calculated with RAxML version 8.0 (Stamatakis 2014) using a general time reversible substitution model and rate heterogeneity modelled with a gamma distribution plus a proportion of invariable sites, as recommended by the program. From this tree, a haplotype genealogy was generated using Haploviewer 1.0 (Salzburger et al. 2011).

The nucleotide diversity (π) was estimated using the R package PEGAS (Population and Evolutionary Genetics Analysis System, Paradis 2010). To analyse the variation in nucleotide diversity across the Iberian Peninsula, we used a visualization method that does not require the pre-definition of populations. The nucleotide diversity was estimated at each sampling location by using all samples present within 30 km of the location. This allowed the estimation of genetic diversity from a sufficient number of samples at each point, yet the resolution was sufficient to distinguish regional differences in genetic diversity. QGIS (<http://qgis.osgeo.org>) was used to plot the nucleotide diversity in the map. In the case of the North American native range, it was not possible to apply this analysis due to the limited number of samples. Therefore, we selected three areas with a sufficient number of samples: North West (British Columbia, six samples), South East (Texas, three samples) and East (New York, four samples), in order to estimate the nucleotide diversity in these specific areas.

Results

Mitochondrial phylogeographic analysis

A total of 16 different haplotypes of the mitochondrial control region were detected in 126 American mink, 11 of which were recorded in North America and 7 in the Iberian Peninsula (ESM Appendix: Fig. S1). The maximum-likelihood phylogenetic tree showed two large clades found throughout western and eastern North America, respectively (Fig. 1). The haplotype genealogy reconstructed from the maximum-likelihood tree revealed the same two large groups (Fig. 2a). As indicated in this genealogy, two mutations unambiguously differentiate both clades. The map of samples clearly shows the geographical separation of the two clades in North America (Fig. 2b). In the Iberian Peninsula, we found more samples belonging to the western than to the eastern clade of North America (68 vs. 37 samples, respectively). However, samples belonging to both clades were detected together in four regions of the north central area of the Iberian Peninsula: Basque Country, La Rioja, Navarra and Avila (South of Castilla-Leon) (Fig. 2c).

In addition, the phylogeographic structure was also different at a finer scale in the native and invasive ranges. While there was evidence of a geographical pattern for the individuals collected from the native range, with individual haplotypes concentrated in a single locality or close localities, many haplotypes were widely dispersed in the Iberian Peninsula (Fig. 1, Fig. S1), showing that haplotypes are not informative with regards to the geographical location of any sample in the invasive range.

Mitochondrial genetic diversity

The global genetic diversity of the American mink estimated with the control region was relatively low for all samples, with a value of nucleotide diversity of $\pi = 0.68\%$, whereas the nucleotide diversity for North America and the Iberian Peninsula was 0.78 and 0.64%, respectively (haplotype diversity was 0.81, 0.92 and 0.76, respectively). However, the nucleotide diversity varied considerably when different localities and populations were considered separately. Among the three areas of the native range in which π was calculated separately, the population of British Columbia showed the highest nucleotide diversity (0.53%) whereas New York (0.12%) and Texas (0%) registered lower values. In agreement with the spatial mixing of clades in the central areas of the Iberian Peninsula, the map derived from the genetic diversity values of the samples around each locality clearly showed maximum levels of genetic diversity in these regions (Fig. 3). When values of several sampling localities were averaged, the Basque population concentrated the highest nucleotide diversity (0.61%), followed by Avila (0.50%) and La Rioja (0.25%). From these areas, genetic diversity decreased towards the remainder of the mink distribution, reaching values of 0% in some areas.

Discussion

Our results show that the establishment of the invasive American mink population in the Iberian Peninsula is the product of the introduction of individuals from the two main genetic lineages found in western and eastern North America. In the invasive range, these lineages registered a complete lack of phylogeographic structure. This was probably the result of the

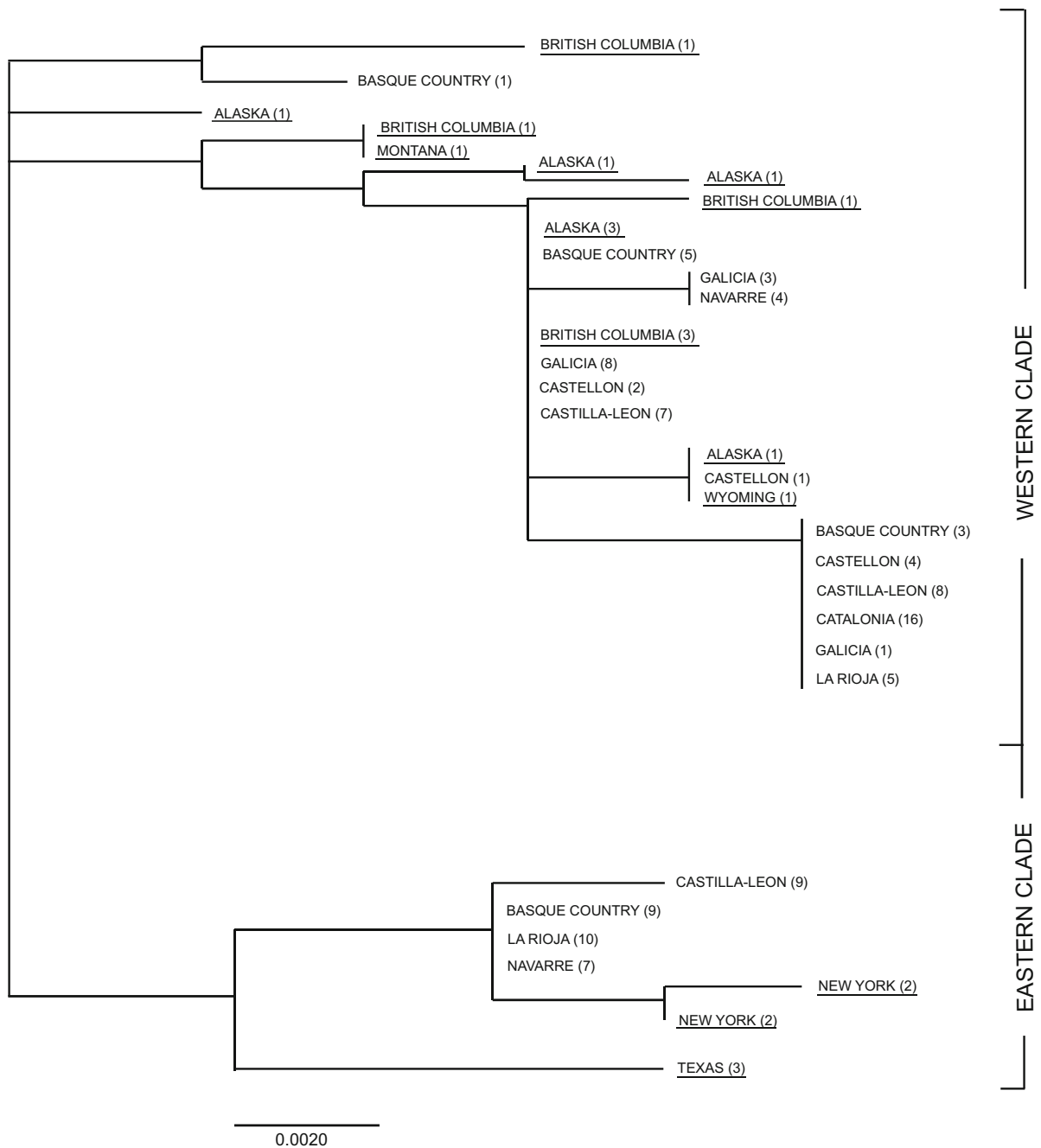


Fig. 1 Maximum-likelihood phylogenetic tree derived from the partial mitochondrial control region sequences for 126 American mink, with eastern and western clades indicated. The

number of individuals per locality is shown in *brackets*. Sequences from the native range are *underlined*. The *scale* indicates number of substitutions per position

additive effect of multiple continuous introductions, and the rapid expansion and encounters of the introduced populations. Overall, genetic diversity was lower in the introduced than in the native

populations, although in some introduced populations diversity was higher to any one native population, at least with respect to the reduced native area analysed so far. These results provide evidence for the potential

Fig. 2 Phylogeographic analysis of the mitochondrial control region sequences of *N. vison*. **a** Haplotype genealogy based on the maximum-likelihood tree. Circle sizes are proportional to the haplotype frequency and *small filled circles* are inferred haplotypes not present in the sample. **b**, **c** Map of native (**b**) and invasive (**c**) ranges showing the 126 samples of *N. vison* used in this study. *Red and blue colours* represent the eastern and western clades, respectively. *Grey areas* represent the original distribution of *N. vison* downloaded from the IUCN Red List of Threatened Species website (Reid et al. 2016) (**b**) and the occupancy area of mink in the Peninsula (MAGRAMA 2013) (**c**). *AL* Alaska, *BC* British Columbia, *MT* Montana, *WY* Wyoming, *TX* Texas, *NY* New York, *GA* Galicia, *CL* Castilla-Leon, *AV* Avila, *LR* La Rioja, *BC* Basque Country, *NAV* Navarra, *CAS* Castellon, *CAT* Catalonia

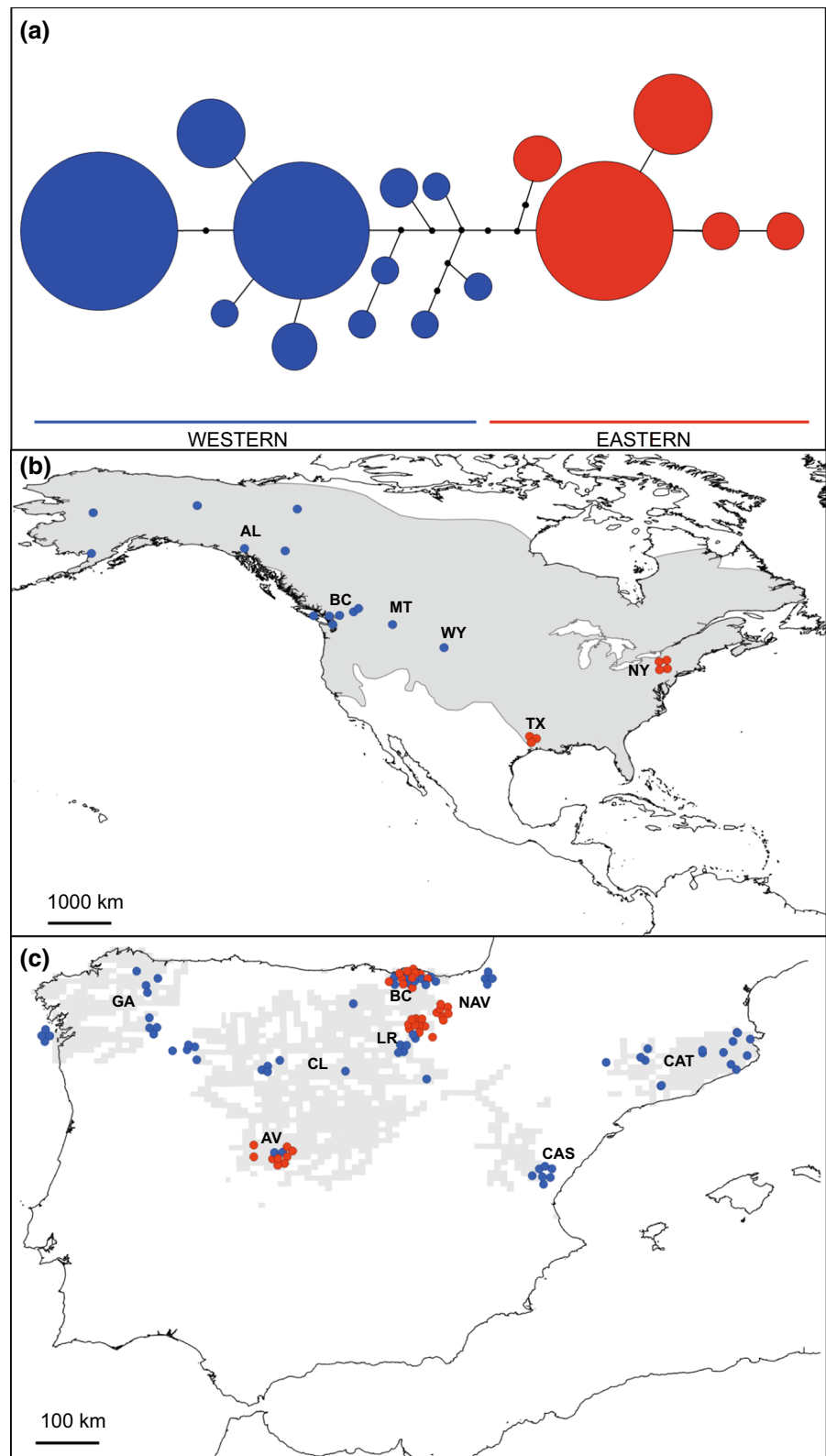
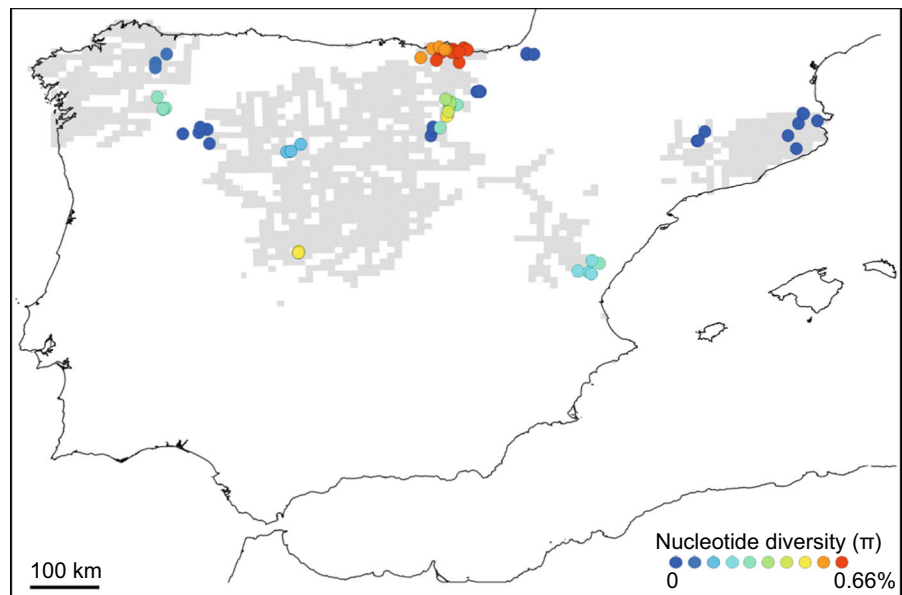


Fig. 3 Map plotting genetic diversity (π) in different sampling localities of *N. vison* in the Iberian Peninsula. Colours of the sample points indicate the nucleotide diversity according to the provided scale



of introduced populations to maintain or increase the genetic diversity.

Phylogeographic patterns

Phylogenetic analysis revealed two large native phylogeographic lineages, which indicated a clear separation between the western and eastern populations in North America. Therefore, the relationship between the mtDNA haplotypes and the location from which they were sampled provides evidence of phylogeographic structure among populations of American mink in their native range. Unfortunately, the lack of information about sequence data of mtDNA of the American mink in North America did not allow our results to be compared with those of similar studies. Other studies of plant and animal species from North America have illustrated a complex phylogeographic history of the biota of this region (Shafer et al. 2010; Soltis et al. 2006). The data currently available suggest that vicariance, dispersal and the existence of refugia have worked in concert to produce the species' distribution and genetic divergence patterns of the region (Brunsfeld et al. 2001). Future research focused on filling the gaps by sampling at locations between the two clades seen in this work is required to achieve a deeper understanding of the evolutionary history of the American mink in its native range.

The pattern of spatial structure recorded in North America allowed for a more precise identification of the origin of invasive populations in the Iberian Peninsula. Our analysis revealed that there were introductions from both western and eastern regions in North America. Additionally, the distribution of the invasive haplotypes in the Iberian Peninsula suggests that at least the north central area of the Iberian Peninsula (populations of the Basque Country, La Rioja, Navarra and Avila) experienced multiple introductions from both regions of North America, while the western and eastern Iberian populations (Galicia, Catalonia and Castellon) originated from introductions from western North America, contrary to what was previously believed (Palazón et al. 1997). This situation most likely occurred as a result of the multiple origins of mink from different farms and the fast range expansion of the occupied area since their introduction in the 1960s–1970s. It is also likely that farms received founder individuals from different source populations in the native range.

The occurrence of the two main mitochondrial lineages in the north central area of the Iberian Peninsula suggests that the intermixing of individuals of both lineages may potentially take place. However, it will be necessary to perform analysis with multiple genomic markers in the future to determine whether individuals belonging to these two lineages effectively interbreed and, if so, to estimate admixture levels in

these populations (Pritchard et al. 2000). It is also important to note that, due to the limited information derived from the mitochondrial DNA, which only reflects the maternal lineage, the existence of some admixture levels in other Iberian populations cannot be discarded until genomic analyses are performed. In fact, a previous study in the Iberian Peninsula based on microsatellites showed that the highest admixture levels between populations were found in the Basque Country population, which in principle is consistent with our results (Lecis et al. 2008). However, since no native specimens were available in that study, it was not possible to determine how the identified populations were related to the source populations. This confirms that it is crucial to obtain samples from the native range to shed light on the relationship between invasive and native populations and to identify potential source populations.

Genetic diversity in the native versus invasive range

Based on mitochondrial DNA control region sequence data, our results provide evidence that the introduced American mink populations in the Iberian Peninsula have overall less genetic diversity than native mink populations in North America. These results are in agreement a priori with the “genetic paradox” (Allendorf and Lundquist 2003; Estoup et al. 2016), which predicts that low genetic variability can be expected in invasive species because of the loss of rare alleles, as a result of the founder effect and population bottlenecks (Allendorf and Lundquist 2003; Grapputo et al. 2005). Nevertheless, when analysed at the population level within each range, nucleotide diversity in the population of the Basque Country and La Rioja in the Iberian Peninsula was found to be greater than that recorded for North American populations. The pattern registered here is consistent with other studies (e.g. Edelaar et al. 2015; Genton et al. 2005; Roman and Darling 2007), providing another example of an invasive species retaining genetic diversity. Similarly, previous studies have demonstrated that the genetic diversity of invasive populations in their new range may be increased through the admixture of lineages from multiple native populations (Keller and Taylor 2010; Kolbe et al. 2008). As already indicated, the pattern observed in the Iberian Peninsula could be the result of multiple introductions of mink deriving

from different local lineages of North America, their rapid expansion, and the encounter of lineages in the north central area of the Iberian Peninsula that may potentially hybridise. However, we cannot discard that cross-breeding of the different North American subspecies or populations took place within farms (Bellevue et al. 1999; Dunstone 1993).

High values of genetic diversity have also been registered in some mink population of Poland (Zalewski et al. 2011). In this case, mean π for all feral mink analysed was 0.94%, with values for different populations ranging between 0 and 1.41%. Although the mitochondrial control region fragment used in that work was different to ours and a direct comparison cannot be performed, the values found in some populations in Poland were much higher than in others. It is therefore likely that the same processes suggested for the Iberian Peninsula (i.e. multiple introductions, rapid expansion and encounters of individuals of different lineages) can be applied to other areas with invasive populations of American mink.

Implications for management

This study provides an understanding of the genetic structure and diversity of the populations of the American mink in the Iberian Peninsula, which can be used to identify key control areas. Thus, one way to improve the effectiveness of management programmes of an invasive species such as the American mink is by incorporating molecular-based information into these programs (Hampton et al. 2004). In this regard, the current spatial mixing of individuals from two different native lineages in the north central area of the Iberian Peninsula (Basque Country, La Rioja, Navarra and Avila populations) and the potential interbreeding between such lineages that may take place could translate, among other outcomes, into a higher fitness in these individuals (e.g. Vandewoestijne et al. 2008). In fact, previous studies have found a positive correlation between genetic diversity and colonization success as measured by the reproduction rate (e.g. Crawford and Whitney 2010). Interestingly, Melero et al. (2012b) found that mink of the Basque Country were bigger in size than mink of the rest of the Iberian Peninsula. According to Melero et al. (2012b), this phenotypic variation could be due to a mixing of individuals from different

origins, among other possibilities. If the link between spatial mixing of mitochondrial lineages, genome admixture and higher fitness is demonstrated for some populations, then higher efforts to control and eradicate the American mink should be undertaken in such populations in order to impede or minimise their expansion.

Current controls of the American mink in the Iberian Peninsula focus on reducing their effects on ecosystems and biodiversity (i.e., areas with local populations of threatened species, wetlands where they can cause high mortalities in waterfowl and protected natural areas, etc.). In this regard, it is important to consider that the areas of the Iberian Peninsula with presence of both American mink lineages coincide with critical areas for the conservation of two of the most threatened mammals of the Iberian Peninsula, the European mink and the Pyrenean desman. The American mink compete successfully for habitat and prey with the European mink (Sidorovich et al. 1999; Sidorovich and MacDonald 2001), whose population of the North of Spain and South West of France is highly isolated from the rest of the distribution (Michaux et al. 2005). On the other hand, the populations of Pyrenean desman of the Central System have experienced a strong reduction in the last few years and are threatened with extinction. Although the reasons for this decline are not well known, predation by the American mink may have contributed to it (Fernandes et al. 2008). Thus, the presence of the two native lineages of the American mink in the areas with these threatened species should be taken into account in future conservation plans due the putative higher expansive potential of the American mink in them.

The results of this work are in line with previous studies suggesting that mink control should also be directed to minimise the contact between genetically differentiated populations (Bifolchi et al. 2010; Fraser et al. 2013; Lecis et al. 2008; Robertson and Gemmill 2004), and so counteract a possible increase in mink fecundity and dispersal (Melero et al. 2015; Oliver et al. 2016). The present study has shown how molecular-based information obtained from the invasive range, together with that from the native range, was critical to characterise the sources from which the Iberian populations of the American mink originated. Further studies are now necessary to obtain more detailed information, particularly in areas where

admixture of lineages may potentially occur, to help prioritise efforts to control and manage this invasive species.

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