

## GENETIC STRUCTURE CONFIRMS FEMALE-BIASED NATAL DISPERSAL IN THE WHITE-TAILED EAGLE POPULATION OF THE CARPATHIAN BASIN

EDINA NEMESHÁZI<sup>1,2</sup>, KRISZTIÁN SZABÓ<sup>1</sup>, ZOLTÁN HORVÁTH<sup>3</sup> and SZILVIA KÖVÉR<sup>1</sup>

<sup>1</sup>*Department of Ecology, University of Veterinary Medicine, H-1077 Budapest, Rottenbiller u. 50, Hungary; E-mails: kr.szabo@gmail.com, kover.szilvia@univet.hu*

<sup>2</sup>*Lendület Evolutionary Ecology Research Group, Plant Protection Institute, Centre for Agricultural Research, Hungarian Academy of Sciences, H-1022 Budapest, Herman Ottó út 15, Hungary; E-mail: nemeshazi.edina@agrar.mta.hu; corresponding author*

<sup>3</sup>*Danube-Dráva National Park Directorate  
H-7625 Pécs, Tettye tér 9, Hungary; E-mail: retisas.hu@gmail.com*

Individuals can avoid inbreeding or competition with kin via long-distance natal dispersal. On the other hand, staying close to the well-known natal area may be a safer choice with respect to recruiting opportunities, reproductive success and the individual's survival probability as well. Natal dispersal strategy often differs between sexes, being generally female-biased in birds. We explored if the Carpathian Basin White-tailed Eagle population shows fine scale genetic structure and if it does, what is the extent of philopatry in the two sexes. We furthermore investigated sex bias in natal dispersal distance inferred from spatial distributions of genetically close relative breeding females and males. Spatial autocorrelation analyses failed to find fine-scale genetic structure, despite the species being known to be philopatric. Pairwise breeding distances of close relatives showed female bias according to Wilcoxon rank sum test. The median distance of two close relative females was 136 km, while it was only 38 km in males. Since White-tailed Eagles are known to be faithful to their breeding territory, we assumed that the breeding distance between an individual and its parents refers to the individual's natal dispersal distance. Due to the same reason, the breeding distance of two siblings should also be related to their individual dispersal distances from their shared natal area. Therefore, we argue that the difference we found between sexes in pairwise breeding distances of close relatives stands for a female-biased natal dispersal. This bias may be a consequence of the species' breeding strategy, and it decreases the inbreeding probability as well.

Keywords: *Haliaeetus albicilla*, raptor, natal dispersal, genetic relatedness, Carpathian Basin.

### INTRODUCTION

Individual dispersal strategy has substantial influence on the population or even species level traits such as population structure, genetic variability, level of inbreeding, metapopulation dynamics and speciation. Background of dispersal can be complex and is yet to be fully understood (MATTHYSEN 2005). The permanent movement that an individual makes from its natal area to the area where it first attempts to breed is referred to as 'natal dispersal', and its subsequent movements between its successive breeding areas are re-

ferred to as 'breeding dispersal' (GREENWOOD & HARVEY 1982). Estimation of natal dispersal distance (NDD; i.e. distance between the natal and first breeding place of an individual) can provide important information related to evolution, ecology and conservation of a species. Long-distance natal dispersal for example can be a successful strategy to avoid inbreeding or competition with kin or more experienced adults for resources and mates (GREENWOOD & HARVEY 1982, PERRIN & MAZALOV 2000). On the other hand, a strategy of short NDD (also referred to as philopatry) can be beneficial as well: moving within a restricted, familiar area, the individual can re-establish a territory just becoming vacant (SMITH 1978), knowing its available resources for feeding and reproduction; in addition, the costs derived from crossing less suitable or even dangerous areas during dispersal may decrease as well (YODER *et al.* 2004). On population level, long-distance dispersal can allow colonization of suitable distant habitats and facilitate gene flow between populations being far apart from each other. In general, long-distance dispersal increases the genetic diversity within (sub)populations and decreases the genetic differentiation between them. By contrast, populations of a strictly philopatric species exhibit strong population genetic structure and show heterozygote deficiency on a metapopulation level (e.g. see ORTEGO *et al.* (2008) for genetic structure of Lesser Kestrels *Falco naumanni* (Fleischer, 1818), and MONTI *et al.* (2018) for that of Osprey *Pandion haliaeetus* (Linnaeus, 1758)).

Philopatry was found in several species with high migration capability, such as many birds and mammals (GREENWOOD & HARVEY 1982, PUSEY 1987, ALCAIDE *et al.* 2009, WHITFIELD *et al.* 2009a). Average NDD can differ between sexes and if so, it is usually male-biased in mammals and female-biased in birds (GREENWOOD & HARVEY 1982, PUSEY 1987). Sex-biased natal dispersal strategies may predominantly be consequences of intrasexual or kin competition for territories, food resources or mates, given that these factors are unbalanced between sexes. Furthermore, sex-biased natal dispersal may serve inbreeding avoidance as well (GREENWOOD & HARVEY 1982, PERRIN & MAZALOV 2000). Territorial birds generally have a so called resource defence system, where males compete for breeding territories and pairing occurs upon female choice (GREENWOOD 1980). This may facilitate a sex bias in natal dispersal strategies as well: if the success of territory acquisition is more promising at the familiar natal area, philopatry may be a strategy especially useful for males, while females may reduce the probability of mating with kin via longer NDD. Female-biased NDD of course reduces the probability of inbreeding for the males as well.

The White-tailed Eagle (*Haliaeetus albicilla* (Linnaeus, 1758); WTE) is a large raptor species distributed across the Palaearctic and Greenland (BIRD-LIFE INTERNATIONAL 2015). While adult WTEs in most populations stay nearby

their breeding territory throughout the year, juveniles are vagrant and can move several hundred kilometres away from their natal area (HELANDER & STJERNBERG 2003, WHITFIELD *et al.* 2009b, NYGÅRD *et al.* 2010). Despite their juvenile vagrancy, similarly to other large raptor species, WTEs are philopatric (i.e. they start breeding relatively close to their natal area), as supported by both ringing data (HELANDER 2003, HELANDER & STJERNBERG 2003, STRUWE-JUHL & GRÜNKORN 2007, WHITFIELD *et al.* 2009a) and genetic structure of their wild populations (HAILER *et al.* 2007, HONNEN *et al.* 2010, NEMESHÁZI *et al.* 2016). Still, some level of long distance gene flow was recently suggested in the Carpathian Basin population (NEMESHÁZI *et al.* 2016). NDD values published from different WTE populations differ considerably, with mean values ranging from about 30 to 114 km (HELANDER 2003, STRUWE-JUHL & GRÜNKORN 2007, WHITFIELD *et al.* 2009a). WHITFIELD *et al.* (2009a) reported that females dispersed on average two times further than males in the reintroduced population of Western Scotland.

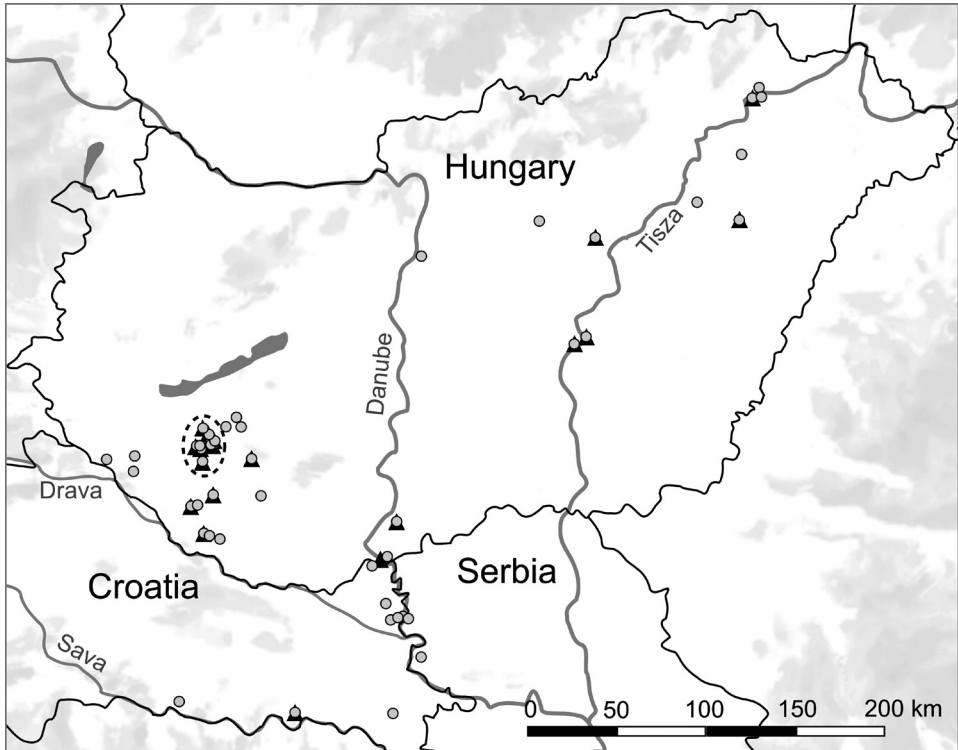
Investigation of large raptor species is challenging; capture of adults on the field is mostly not possible. Therefore, instead of conventional methods, genetic data recovered from non-invasive samples are increasingly used to investigate several topics of individual-based studies, such as annual turnover rate of breeding individuals, parentage, breeding dispersal, or territorial intrusions (RUDNICK *et al.* 2005, 2009, BOOMS *et al.* 2011, VILI *et al.* 2013b, NEMESHÁZI *et al.* 2018). Straightforward methods for investigation of natal dispersal require knowledge on both the hatching and first breeding place of the sampled individuals. Such investigations in large raptors require long-term investments, as several years pass between the hatching and recruitment of individuals (e.g. the first breeding attempt generally occurs around the 6th calendar year in WTEs: HELANDER & STJERNBERG 2003). However, spatial genetic structure of breeders can reflect the extent of philopatry as well, as shown in wild populations of several bird species (DOUBLE *et al.* 2005, TEMPLE *et al.* 2006, PONNIKAS *et al.* 2013). As observations suggest that WTEs are long-term faithful to their territories (HELANDER 2003, HELANDER & STJERNBERG 2003, KRONE *et al.* 2013), we assume that breeding dispersal is negligible in this species. Therefore, the scale of their natal dispersal can be inferred indirectly from the genetic relatedness of breeding individuals, even if their hatching places are unknown. This estimated level of natal dispersal can be biased by rare events of breeding dispersal, but such events would influence the estimations of the difference between sexes only if breeding dispersal itself was sex-biased.

We concentrated on two questions in the present study: 1) Is there a fine scale genetic structure within the White-tailed Eagle population of the Carpathian Basin as suggested by the philopatric behaviour of the species? 2) Is there a difference between sexes in pairwise breeding distance of close

relatives? We assumed that if there is a sex bias in pairwise breeding distance of close relatives, the same stands for a sex bias in natal dispersal distance as well. Both questions were investigated using genotypes of non-invasively sampled breeding individuals across the Carpathian Basin.

## MATERIAL AND METHODS

We used DNA samples collected between 2010 and 2016 by fellow workers of assigned national parks in the Carpathian Basin WTE population (Fig. 1). Moulded feathers were collected during the breeding season within approximately 100 m distance from nests where breeding attempts were observed. In most regions, moulded feathers were collected and stored as suggested by VILI *et al.* (2013a). Resident individuals of each territory were identified based on matching nestling genotypes or the largest number of matching moulded feathers (but at least three) shed by same-sex individuals at the same nest site. Feathers shed by intruders may also occur at occupied nest sites and misidentification of the intruders as breeders could significantly bias the results of investigations concentrat-



**Fig. 1.** Nesting locations of breeding White-tailed Eagles sampled across the Carpathian Basin. Black triangles indicate males and grey circles indicate females. In the area depicted by the dashed line overall 12 males and 12 females were sampled

ing on the breeding population. Nevertheless, our conservative criteria described above were assumed to be adequate for identification of resident WTEs based on non-invasive sampling (NEMESHÁZI *et al.* 2018). Individuals which did not meet the above criteria were excluded from the further analyses.

DNA was extracted from the superior umbilicus in large feathers (HORVÁTH *et al.* 2005), and the whole quill below the vane in small ones. DNA purification kits (Quiagen – DNEasy Blood and Tissue Kit or Thermo – GeneJet genomic purification kit) were used following the manufacturers' instructions, with 10  $\mu$ l of 1M dithiothreitol added during the digestion step.

Sex was identified from feather samples using the 2550F/2718R (FRIDOLFSSON & ELLEGREN 1999) or the GfUp/GErUp and GfLow/GErLow primer pairs (OGDEN *et al.* 2015). PCR profiles followed the original papers, but PCR profile of the latter was completed by an initial touch-down section where the annealing temperature decreased from 65 to 60°C during 7 cycles. PCR products were visualized under UV light (2% agarose gel stained with ECO Safe; Pacific Image Electronics Co., Ltd.).

Each DNA sample was genotyped using 12 microsatellite loci: Hal01, Hal03, Hal04, Hal09, Hal13 (HAILER *et al.* 2005), Aa27, Aa35, Aa49 (MARTÍNEZ-CRUZ *et al.* 2002), IEAAAG04, IEAAAG05, IEAAAG12 and IEAAAG14 (BUSCH *et al.* 2005). PCR profiles followed the original papers for Aa and Hal loci, and HAILER *et al.* (2006) for IEAAAG loci, with some modifications (NEMESHÁZI *et al.* 2016). Forward primers were 5' labelled with fluorescent dyes (VIC<sup>TM</sup>, FAM6<sup>TM</sup>, PET<sup>TM</sup>, NED<sup>TM</sup>, or HEX) and fragment length analyses were performed on an ABI3130 Genetic Analyzer (Applied Biosystems, using Gene Scan<sup>TM</sup> -500LIZ<sup>TM</sup> Size Standard). Trace files were scored in Peak Scanner 1.0 (Applied Biosystems). We used Micro-Checker 2.2.3. (VAN OOSTERHOUT *et al.* 2004) to test for presence of null alleles and scoring errors due to large allele dropout or stutter bands. Genepop 4.2. (ROUSSET 2008) was used to estimate pairwise linkage disequilibrium across loci. Consensus individual genotypes were prepared manually, based on genotypes of feathers collected at the same nest site (as described in NEMESHÁZI *et al.* 2018). Probability of identity for combinations of increasing number of loci was calculated with GenAlEx 6.5 (PEAKALL & SMOUSE 2012).

Fine scale genetic structure within the study population was investigated by testing spatial genetic autocorrelation (SMOUSE & PEAKALL 1999) based on individual genotypes of resident WTEs. Spatial genetic autocorrelation was investigated by two methods: using pairwise genetic distance calculated for co-dominant loci in GenAlEx 6.5 (PEAKALL & SMOUSE 2012), and kinship coefficient described by LOISELLE *et al.* (1995) in SPAGeDi 1.5a (HARDY & VEKEMANS 2002). In these analyses, pairwise Euclidean distances between nest sites of sampled WTEs were used to test whether individual genotypes show random distribution in space; dyads of individuals were grouped into predefined spatial distance classes. GenAlEx calculated a genetic autocorrelation coefficient ( $r$ ) for each distance class; the statistical significance was tested based on random permutation of individuals across distance classes (no. permutations = 9999) and bootstrap estimates of  $r$  (no. bootstraps = 10000). In SPAGeDi, the association between genetic relationship and spatial distances was assessed by averaging the pairwise genetic relationship for each distance class, and statistical significance for spatial genetic structure was tested by 10000 permutations of locations and gene copies. Both in GenAlEx and SPAGeDi, we performed calculations on three datasets: (i) all breeding individuals, (ii) females and (iii) males. Distance classes were defined aiming for each class to contain a reasonably large number of pairwise data (minimum value was 380 for all breeding individuals, 117 for females and 37 for males). As spatial distribution of male-male dyads was aggregated (forming 4 groups: Fig. 2a), upper limit

of each distance class was adjusted according to these aggregations, but the first group was divided in two distance classes. We performed this latter subdivision because two of the three datasets (i, iii) contained significantly more dyads between the limits of the first aggregation than between the others'. Accordingly, 5 subsequent distance classes were analysed for spatial autocorrelation, with upper limits of 20, 80, 180, 280 and 400+ km, respectively. Individuals breeding between 0 and 20 km from each other were grouped in the 20 km class, those breeding for more than 20 km up to 80 km to the 80 km class, and so on. The last distance class (referred as 400+) differed among datasets, as the longest distance between two sampled males was 420 km, while it was 463 km between two females as well as between two sampled breeding individuals overall.

To test for a difference between sexes in pairwise breeding distance of close relatives, we calculated Maximum Likelihood pairwise relatedness in ML-Relate (KALINOWSKI *et al.* 2006) based on allele frequencies across all sampled individuals. We previously found this a reliable estimator inspecting real parent-offspring relationships in the Hungarian WTE population (NEMESHÁZI *et al.* 2017). The expected value of relatedness for close relatives (i.e. siblings or parent and offspring) is 0.5. We assumed individuals to be closely related if their estimated pairwise relatedness was above 0.4, since these estimated values scatter around the expected values of relatedness: using the same 12 loci dataset, across 67 real parent-offspring dyads from Hungary, 85% got a pairwise relatedness value of at least 0.5, and 91% got a value greater than 0.4. In contrast, from 2516 dyads of presumably not related WTEs, 96% got values below 0.4 (calculated from the results of NEMESHÁZI *et al.* 2017). Note, that the value 0.4 is closer to 0.5, than to the expected value of the next level of relatedness (0.25 for half-siblings, grandparent and grandchild, etc.). One-tailed Wilcoxon rank sum test was calculated in R 3.1.2. (R CORE TEAM 2015) to assess whether the distribution of geographic breeding distances of close relative females was shifted towards greater values than that of close relative males; using the function 'wilcox.exact' in the R package 'exactRankTests'. We used one-tailed test, because where NDD differs among sexes in birds, usually females disperse further (GREENWOOD & HARVEY 1982), and female bias was previously found in another WTE population as well (WHITFIELD *et al.* 2009a). To test for difference in variances across the two sampled groups, we performed Levene's test for homogeneity of variances in geographic distances across closely related males and closely related females; using the function 'leveneTest' in the R package 'car'.

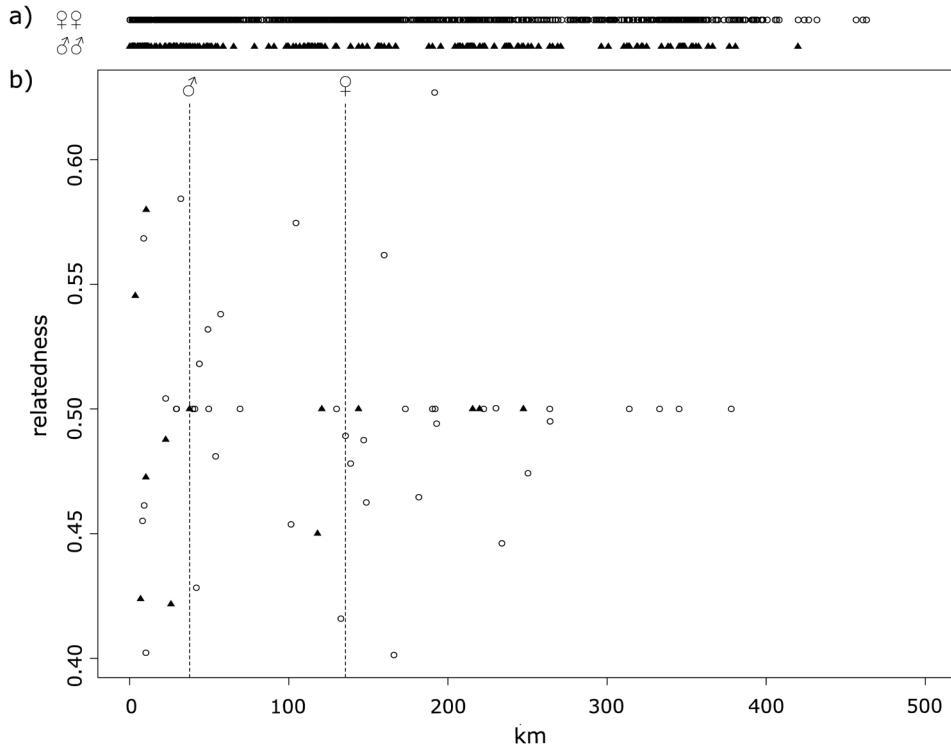
## RESULTS

We used DNA samples extracted from a total of 214 moulted feathers belonging to 73 resident WTEs across the Carpathian Basin (24 males and 49 females; Fig. 1). Each individual consensus genotype consisted of the full set of 12 loci.

A total of 59 alleles were found; the mean number of alleles across the 12 loci was 4.9, ranging from 2 on Hal03 to 10 on IEAAAG05. Analyses in Micro-Checker suggest that our dataset is not influenced by null alleles, large allele dropout or stutter bands. No evidence for pairwise linkage disequilibrium was found across the loci after Bonferroni correction. Probability of identity was estimated at  $3.3 \times 10^{-8}$  for 12 loci genotypes, and that for siblings was esti-

mated at  $5.2 \times 10^{-4}$  suggesting that our dataset is reliable for individual-based genetic analyses (i.e. being below or about the magnitude of the threshold of 0.0001, following *Waits et al.* (2001).

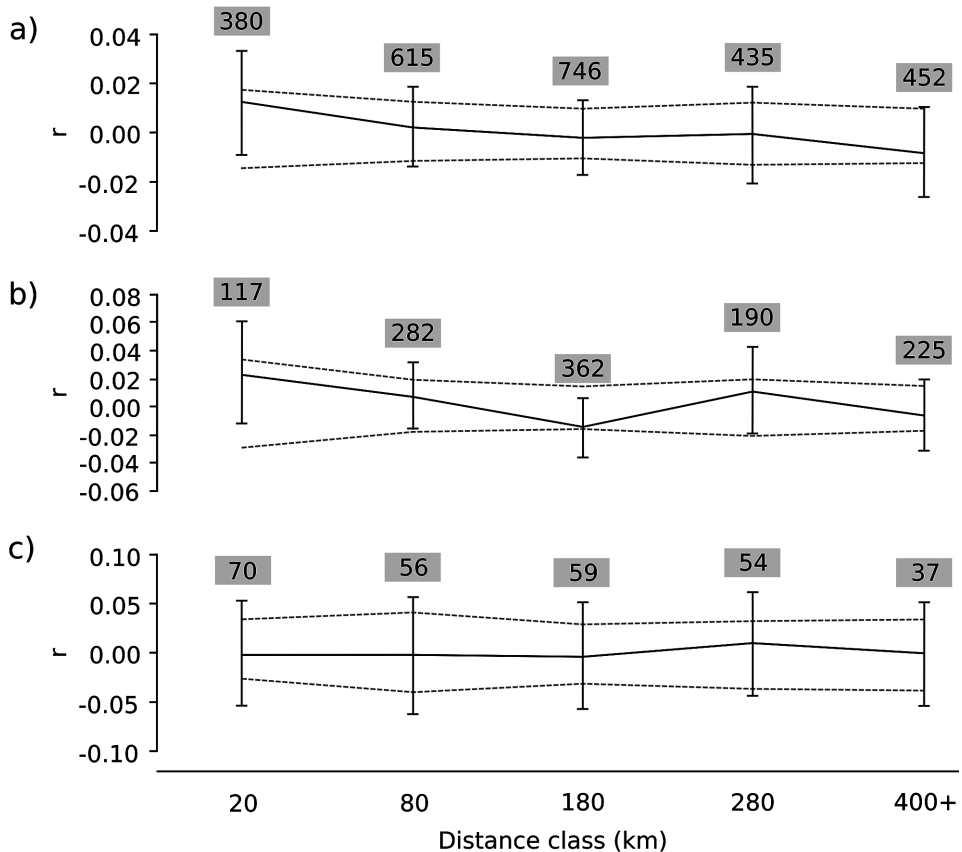
In SPAGeDi, spatial autocorrelation analysis of all breeding individuals ( $N = 73$ ) showed a moderate, but significant negative relationship between the average pairwise genetic similarity within each distance class and its logarithmic spatial distance (slope  $\ln(\text{dist}) = -0.0027$ ,  $p = 0.044$ ). However, no such relationship was found based on linear spatial distances, and the observed mean kinship values of each distance class did not significantly differ from that expected under random spatial distribution. GenAlEx also failed to find a relationship between the genetic autocorrelation coefficient  $r$  and spatial distance, and no significant deviation of  $r$  from zero was observed in any of the distance classes (Fig. 3a). When analysing females ( $N = 49$ , Fig. 3b) and males



**Fig. 2.** Pairwise breeding distance of White-tailed Eagles: a = spatial distributions of all investigated same-sex dyads (spatial scale is shown in b); b = pairwise spatial distances (km) of individuals with a pairwise Maximum Likelihood relatedness (relatedness) of minimum 0.4. Male-male (triangles) and female-female (circles) dyads are shown separately. Vertical dashed lines indicate the median pairwise breeding distance of related males and that of related females

( $N = 24$ , Fig. 3c) separately, both methods failed to find any spatial genetic structure and even the slope for logarithmic spatial distances was nonsignificant for each sex.

Overall 43 female-female and 13 male-male dyads were assumed to consist of closely related individuals, based on their pairwise Maximum Likelihood relatedness being at least 0.4 (Fig. 2b; note that these dyads were formed by overall 38 female and 18 male individuals). The maximum breeding distance was 378 km among related females and 247 km among related males; median values of breeding distance were 136 and 38 km, respectively. The



**Fig. 3.** Correlogram plots of the degree of genetic correlation coefficient ( $r$ ; calculated in GenAlEx) as a function of breeding distance: a = plot for dyads of all sampled breeders; b = plot for dyads of females; c = plot for dyads of males. Dashed lines indicate the permuted 95% confidence intervals. Bootstrapped 95% confidence error bars are also shown. Numbers with grey background indicate the numbers of pairwise comparisons (i.e. dyads) within each distance class. Note that the vertical axis scales differ among plots



Wilcoxon rank sum test revealed that distribution of pairwise geographic distances was significantly shifted towards greater values for close relative females than for close relative males ( $p = 0.038$ ,  $CI: 2.609$ - $Inf$ ,  $W = 371$ ). Levene's test found no significant difference in the variance of geographic distances across related males and related females ( $p > 0.6$ ).

## DISCUSSION

Juvenile White-tailed Eagles (WTE) are vagrant and can travel several hundred kilometres from their natal area (HELANDER & STJERNBERG 2003, WHITFIELD *et al.* 2009b, NYGÅRD *et al.* 2010), but they tend to start breeding relatively close to it, and the species is therefore assumed to be philopatric (HELANDER 2003, HELANDER & STJERNBERG 2003, WHITFIELD *et al.* 2009a). Mean natal dispersal distance (NDD) values reported in European WTE populations differ considerably, being below 60 km in the reintroduced population of Western Scotland (WHITFIELD *et al.* 2009a) and exceeding 100 km in Sweden (HELANDER 2003). Observations in some WTE populations suggest that NDD can be female biased (HELANDER 2003, WHITFIELD *et al.* 2009a). Lacking sufficient information published from conventional methods in the Carpathian Basin WTE population, we aimed to infer the spatial scale of natal philopatry based on microsatellite genotypes of breeding individuals.

First, we investigated fine scale genetic structure within the study population by testing spatial genetic autocorrelation. This method is known to be effective in highlighting genetic spatial patterns expected in a philopatric population, or under sex-biased dispersal, in the philopatric sex (DOUBLE *et al.* 2005, TEMPLE *et al.* 2006, BANKS & PEAKALL 2012, PONNIKAS *et al.* 2013). Namely, relatively high genetic similarity is expected among neighbouring individuals, or across the first few distance classes. Although analyses in SPAGeDi revealed that the slope of the relationship of genetic similarity and logarithmic spatial distance (slope  $\ln(dist)$ ) was significantly negative for dyads of WTEs breeding in the Carpathian Basin, both SPAGeDi and GenAlEx failed to find significant deviation from a random spatial distribution of pairwise genetic relatedness values. When we analysed females and males separately, similar lack of fine-scale genetic structure was found, with even the slope  $\ln(dist)$  being nonsignificant. Nevertheless, the species is philopatric, as showed both by mark-resight data (HELANDER 2003, HELANDER & STJERNBERG 2003, STRUWE-JUHL & GRÜNKORN 2007, WHITFIELD *et al.* 2009a) and the presence of genetic structure at wide geographical range inferred from microsatellite data (HAILER *et al.* 2007, HONNEN *et al.* 2010, NEMESHÁZI *et al.* 2016). Moreover, the Carpathian Basin population maintains a local, unique mitochondrial haplotype as well (HONNEN *et al.* 2010, NEMESHÁZI *et al.* 2016). Similar contradictions

of fine-scale spatial genetic and mark-resight data were reported in a lesser kestrel (*Falco naumanni*) population (ALCAIDE *et al.* 2009), where the authors concluded, that rare occurrence of long-distance dispersal events overrode the genetic structure expected in the otherwise philopatric population. Long-distance natal dispersal events have been observed in WTEs as well (STRUWE-JUHL & GRÜNKORN 2007), suggesting that our case with WTEs may be similar to that described by ALCAIDE *et al.* (2009) for the lesser kestrels.

In the Finnish WTE population, PONNIKAS *et al.* (2013) reported strong genetic spatial autocorrelation for nestlings, and the mean distance of the highest distance class with significantly positive kinship coefficient corresponded to both ringing data from the Finland population and the known mean natal dispersal distances in the neighbouring population of Sweden. Using the same method for calculating genetic spatial autocorrelation, we failed to find such pattern among breeding WTEs in the Carpathian Basin. Investigating several WTE populations across Europe, NEMESHÁZI *et al.* (2016) suggested that the impact of immigrants from distant populations may be higher in the Carpathian Basin than in populations to the North. Considering above data we hypothesize that the lack of fine-scale genetic structure in our study despite the known philopatric behaviour of the species is an outcome of some level of long-distance natal dispersal: if such events occur within the population, they increase the average genetic relatedness in higher distance classes, and immigrants may cause a background noise across all distance classes. Our limited sample size, especially in males where genetic similarity within a reduced spatial distance should be higher, may have further increased difficulties for finding fine-scale genetic structure (BANKS & PEAKALL 2012). Furthermore, the investigated spatial scale may be smaller than the general extent of natal dispersal: in this case, extension of sampling to wider geographical range would be needed to find genetic structure by spatial autocorrelation analyses. However, the fact that the Carpathian Basin maintains a unique genetic composition suggests that natal dispersal in this WTE population is generally restricted to the spatial scale of this region (NEMESHÁZI *et al.* 2016). Therefore, the extent of our sampling area should be sufficient for investigation of questions related to natal dispersal in the Carpathian Basin population. Similarly, ringing data of a limited number of individuals seem to confute this possibility: all seven individuals originating from Hungary with known natal dispersal distance remained within the country (unpublished data of the Hungarian Bird Ringing Centre).

As a second objective, we tested whether close relative females disperse at larger distances than close relative males, as suggested by the typical pattern of female-biased natal dispersal in birds. Similar method was used by HAAS *et al.* (2010) in a Brown-headed Nuthatch (*Sitta pusilla* Latham, 1790)

population. Note, that we could only investigate the pairwise breeding distance of individuals, which is not equivalent to the NDD, although related to that. For example, if two close relatives with a pairwise breeding distance of 200 km were parent and offspring, than the offspring probably indeed had an NDD value of 200 km; as it most probably breeds in its first territory (given that WTEs are long-term faithful to their breeding territories: HELANDER 2003, HELANDER & STJERNBERG 2003, KRONE *et al.* 2013). If they were siblings, then the fact that they bred 200 km apart means that one of them could have dispersed 200 km while the other one remained completely faithful to the natal area, or both could have dispersed 100 km, or even more, from an area located further from the bee line between the breeding sites of the two individuals. Therefore, although our findings on close relatives with considerable breeding distance up to 378 km do not explicitly correspond to the NDD, they indicate the occurrence of long-distance natal dispersal. The longest pairwise breeding distances of close relatives found in our study are comparable with the dimensions of the Carpathian Basin. This is in accordance with previous findings based on nestling microsatellite genotypes that the Carpathian Basin maintains one genetic WTE population (NEMESHÁZI *et al.* 2016). We presumed that the directions of natal dispersal are not sex-biased (WHITFIELD *et al.* 2009a), and therefore the relationship of NDD and breeding distance of close relatives should be similar for both sexes.

We found that the median distance of close relative breeding females is bigger than that of close relative males and this result suggests that long-distance natal dispersal events are female-biased in the Carpathian Basin WTE population. Similarly female-biased natal dispersal was found in the reintroduced population of Western Scotland based on mark-resight data, with females dispersing on average two times further than males (WHITFIELD *et al.* 2009a). HELANDER (2003) also reported somewhat higher mean NDD for females than for males in Sweden. Female-biased dispersal is common in bird species, and may serve inbreeding avoidance, or may be a consequence of different sex roles derived from their resource defence system (GREENWOOD & HARVEY 1982, PERRIN & MAZALOV 2000). Our findings suggest that sex-biased natal dispersal contribute to the generally low pairwise intersexual relatedness of WTEs breeding close to each other in Southwestern Hungary (NEMESHÁZI *et al.* 2018).

Despite the known philopatry of the species, our spatial autocorrelation analyses generally failed to show evidence of spatial genetic structure within the Carpathian Basin or gave only a weak evidence of it. We assume that this contradiction is due to occurrence of long-distance natal dispersal events together with our limited sample size (and maybe the limited spatial scale). On the other hand, the comparison of pairwise breeding distances of closely

related same-sex individuals revealed a significant female bias. We conclude that long-distance natal dispersal events in the Carpathian Basin are female-biased, in accordance with a limited number of previous reports on sex-biased NDD in other European WTE populations. This bias may be a consequence of the territorial behaviour of the males: if a recruiting young male has better chances to acquire a good territory near its natal place then male philopatry imposes a pressure on females to avoid mating with kin by dispersal. Furthermore dispersal can be advantageous if resources show spatio-temporal variation or because it reduces competition with kin. For WTEs these latter advantages of dispersal seem not to differ between the sexes thus we think these are less likely to account for the female-biased pattern we found.

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Acknowledgements – Moulded feathers in Eastern Hungary were provided by Gábor Tihanyi. Some of the feathers from Southwestern Hungary were collected by Attila Mórocz, Tamás Nyemcsok, Andrea Schulcz, Balázs Szélényi and Attila Bereczky. Genetic data of Croatian and Serbian individuals were available from our former study, in collaboration with Tibor Mikuska and István Hám. Nóra Vili helped with useful suggestions for the final touches of the manuscript. We also thank for the insightful suggestions of an anonymous reviewer. Laboratory work was funded by the Normative Research Funding Committee of the Szent István University.

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Received September 18, 2017, accepted June 8, 2018, published August 1, 2018