Concordant mitochondrial and microsatellite DNA structuring between Polish lowland and Carpathian Mountain wolves

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Abstract Phylogeographic studies of highly mobile large carnivores suggest that intra-specific genetic differentiation of modern species might be the consequence of the most recent Pleistocene glaciation. However, the relative influence of biogeographical processes and subsequent human-induced population fragmentation requires a better understanding. Poland represents the western edge of relatively continuous distributions of many wide-ranging species, e.g. lynx (Lynx lynx), wolves (Canis lupus), moose (Alces alces) and, therefore, a key area for understanding historic and contemporary patterns of gene flow in central Europe. We examined wolf genetic structure in Poland and in a recently recolonized area in eastern Germany using microsatellite profiles (n = 457) and mitochondrial DNA sequencing (mtDNA, n = 333) from faecal samples. We found significant genetic structure and high levels of differentiation between wolves in the Carpathian Mountains and the Polish lowlands. Our findings are consistent with previously reported mtDNA subdivision between northern lowlands and southern mountains, and add new and concordant findings based on autosomal marker variation. Wolves in western Poland and eastern Germany showed limited differentiation from northeastern Poland. Although the presence of private alleles suggests immigration also from areas not sampled in this study, most individuals seem to be immigrants from northeastern Poland or their descendants. We observed moderate genetic differentiation between certain northeastern lowland regions separated by less than 50 km. Moreover, mtDNA results indicated a southeastern subpopulation near the border with Ukraine. The observed structure might reflect landscape fragmentation and/or ecological differences resulting in natal habitat-biased dispersal.
Keywords Canis lupus · Poland · Eastern Germany · Genetic structure · mtDNA lineages · Nuclear DNA · Non-invasive molecular techniques

Introduction

The Pleistocene glaciations have had profound implications for biogeographical processes such as distributions, population fragmentation and gene flow in wild species (e.g. Taberlet et al. 1998; Hewitt 2000). Data from highly mobile large carnivores indicate weak phylogeographic structuring prior to the last glacial maximum, suggesting that the current distributions of distinct genetic lineages arose in consequence of the most recent glaciation (Hofreiter et al. 2004). However, ensuing human activity has resulted in further and more recent fragmentation in populations of species such as European brown bears (Ursus arctos) (Zedrosser et al. 2001), Eurasian lynx (Lynx lynx) (Schmidt et al. 2011) and Canadian wolves (Canis lupus) (Stronen et al. 2012). The extent to which historical biogeographical processes and recent human-induced fragmentation have contributed to shape the extant population genetic subdivisions in vagile taxa requires further understanding.

An abrupt ecological transition between the Great European Plains and the Carpathian Mountains occurs in southern Poland. These areas represent potential contact zones between genetically divergent lineages of organisms that recolonized central Europe during the Holocene from different glacial refugia in the Balkans, the Carpathians, and in other parts of eastern Europe (Bhagwat and Willis 2008 and references therein). In several species, contact zones between different mitochondrial DNA (mtDNA) lineages have been detected in Poland. Examples include the Norway spruce Picea abies (Tollefsrud et al. 2008), the bank vole Myodes glareolus (Wójcik et al. 2010), and the red deer Cervus elaphus (Niedziałkowska et al. 2011). Poland represents the western edge of relatively continuous distributions of wide-ranging Palearctic species, including the lynx (Schmidt et al. 2009), moose (Alces alces) (Schmölcke and Zachos 2005), and wolf (Jędrzejewski et al. 2010). Some of these populations are now expanding westward, recolonizing parts of central Europe (e.g., the moose (Schmölcke and Zachos 2005, and references therein); the wolf (Nowak et al. 2011)). Other populations are geographically and perhaps genetically isolated. Lynx inhabiting northeastern Poland were reported to be genetically distinct from their conspecifics in Latvia and Estonia and are presumably isolated by unsuitable dispersal habitats (Schmidt et al. 2009). Moose appear to be expanding from their western range edge in Poland into the Czech Republic and Germany (reviewed in Schmölcke and Zachos 2005). Although the population structure of red deer in Poland and neighbouring countries has been mainly shaped by human translocation, some differentiation between populations in Carpathian Mountains and northern Poland is still detectable (Niedziałkowska et al. 2011, 2012 and references therein).

Wolves in eastern Europe show high levels of genetic diversity and distinct mtDNA haplogroups (Pilot et al. 2010). Two mtDNA haplogroups, which might have diverged about 200,000 years ago, converge in the Carpathian Mountains (Pilot et al. 2010). However, earlier results from microsatellite markers could not identify distinct populations in NE Poland and the Carpathian Mountain (Pilot et al. 2006), underlining the need to clarify patterns of gene flow and the origin of expanding populations.

Distinct genetic structuring of the Polish wolf population was suggested by an earlier large-scale study of central and eastern European wolves (Pilot et al. 2006). Although Pilot et al. (2006) sampled only a portion of the Polish wolf range, they detected 3–4 subpopulations delimited by mtDNA and 2 subpopulations based on microsatellite loci. Western Poland has abundant wolf habitat, and the country could support two to three times its current population of approximately 650 wolves (as estimated by Jędrzejewski et al. 2008, 2010). Wolves have recently recolonized parts of western Poland and eastern Germany, probably from northeastern Poland (Jędrzejewski et al. 2004, 2005a; Anstorge et al. 2006; Nowak et al. 2011). Consequently, Poland represents a key area for understanding historic and contemporary patterns of wolf gene flow in central Europe.

In this study we sampled the entire wolf distribution range in Poland and a portion of eastern Germany that now represents a natural extension of the population in western Poland. The wolf is a protected species, and we consequently based our investigations mainly on non-invasive molecular techniques using DNA extracted from faecal samples. The aim of our study was to answer the following questions: (1) Are wolf populations genetically structured across Poland? (2) Do mitochondrial and nuclear genetic markers show concordant results? Because the central European wolf range has expanded over the past decades, we also asked: (3) Were western Poland and eastern Germany recolonized by wolves from northeastern or southern Poland?

Materials and methods

Study area

We collected wolf scat and tissue samples throughout Poland (311,904 km², 49°00’–54°50’N, 14°08’–24°09’E). This territory extends throughout various geographical regions, with lowlands (<300 m a.s.l.) dominating in the northern and central part and uplands (301–500 m a.s.l.) and mountains (501–2,499 m a.s.l.) in the south. The climate of Poland is temperate with transitional character (oceanic in the north and west, continental in the east). The mean annual temperature...
decreases from south-west (8.5 °C) to north-east (6 °C). Annual precipitation is 500–650 mm in the lowlands and 1,200–1,500 mm in the mountains. The snow cover persists from an average of 40 days in south-western Poland to 100 days in the north-eastern part of the country and 200 days in the mountains (Concise Statistical Yearbook of Poland 2011). The mean human population density is 122 inhabitants/km² and ranges from 20 in north-western and north-eastern Poland to 500 inhabitants/km² in Upper Silesia in south-western Poland (Demographic Yearbook of Poland 2011).

Forests cover 29% of the country; the rest is primarily farmlands (60%) with predominance of arable land. Most forests are coniferous stands (51%) dominated by Scots pine (Pinus sylvestris, 60% of the forest area) and Norway spruce (Picea abies, 6%). Deciduous and mixed forest with oak (Quercus robur and Q. petraea), ash (Fraxinus excelsior), maple (Acer platanoides and A. pseudoplatanus), beech (Fagus sylvatica), hornbeam (Carpinus betulus), birch (Betula pubescens and B. pendula) and alder (Alnus glutinosa) constitute about one-fourth of all forest land. About 98% of woodlands are commercial stands and only 2% are national parks and reserves (Forestry 2011).

Almost all of Poland is inhabited by three native species of ungulates: red deer, wild boar (Sus scrofa), and roe deer (Capreolus capreolus). In north-eastern and central parts of the country, moose and three populations of European bison (Bison bonasus) occur. The Carpathian Mountains harbour isolated populations of chamois (Rupicapra rupicapra) and European bison. Populations of fallow deer (Dama dama), sika deer (Cervus nippon) and mouflon (Ovis musimon) have been introduced to certain forests (Wawrzyniak et al. 2010). Large predators—wolf and lynx—occur mainly in the north-eastern, eastern, and south-eastern parts of the country (Jędrzejewski et al. 2004, 2008; Niedziałkowska et al. 2006). However, wolves have recently been recolonizing the large woodlands of western Poland (Nowak et al. 2011) and the current range of wolves in Poland is shown in Fig. 1.

Sample collection

Wolf scat and tissue samples were collected during 2001–2009. Foresters, national park rangers, students, volunteers, and personnel of the Mammal Research Institute of the Polish Academy of Sciences (hereafter MRI PAS), the Association for Nature “Wolf”, and the Institute of Nature Conservation PAS surveyed areas within the presently known wolf range and gathered wolf scats throughout the year. Small fragments of fresh wolf faeces were either stored in plastic tubes (5–30 ml) filled with
Table 1 Wolf (*Canis lupus*) samples collected and analysed from Poland and Eastern Germany, 2001–2009

| No. | Region | Number of collected samples | Number of scat samples analysed | Number of wolf genotypes identified based on microsatellite markers | Number of individuals with mtDNA sequenced | Number of individuals genotyped at nine SNPs |
|-----|--------|-----------------------------|---------------------------------|---------------------------------------------------------------|------------------------------------------|--------------------------------------------|
| 1   | Bieszczady Mountains and Dynów Foothills | 115 (3) | 100 | 72 | 62 | 8 |
| 2   | Low and Śpidecki Beskid | 52 | 31 | 28 | 6 |
| 3   | Gorce and Tatry Mountains, Żywiec-Silesian Beskid | 107 | 100 | 44 | 12 |
| 4   | Roznowce | 161 (6) | 143 | 38 | 12 | |
| 5   | Parnawa, Sobirów and Włodawa Forests | 168 (8) | 184 | 56 | 10 |
| 6   | Knyszyn, Białowieża and Melnik Forests | 168 | 124 | 40 | 10 |
| 7   | Biebrza River Valley | 272 (6) | 220 | 84 | 4 |
| 8   | Augustów Forest | 272 (6) | 220 | 84 | 4 |
| 9   | Rumia and Borki Forests | 125 (4) | 87 | 26 | |
| 10  | Przele Forest | 125 (4) | 87 | 26 | |
| 11  | Biebrza River Valley | 125 (4) | 87 | 26 | |
| 12  | Western Poland (and Eastern Germany) | 231 (3) | 222 | 87 | 62 |

**Total** | 1715 (38) | 1407 | 576 | 333 | 66 |

Figure 1 shows the location of regions 1-12. Samples were scats and occasionally tissue (numbers of the latter in parentheses, quantities are included in the total numbers). A subsample from Eastern Germany included: 19 scats collected, 19 scats analysed, 9 wolf genotypes based on microsatellite markers, and 5 individuals with mtDNA sequenced.
96 % alcohol or ALS buffer (QIAGEN), or kept frozen at 
−20 °C. In total, 196 scat samples were collected in 
Poland (Table 1). In addition, 19 scat samples from eastern 
Germany were added to supplement data from recently 
colonized areas near the Polish-German border (Lower 
Silesia Forest). Information on dead wolves (mainly poa-
chloroformed or killed in vehicle collisions) was also recorded and 
tissue or skin samples (n = 38) were collected. We henceforth refer to geographic sampling areas as sampling 
regions (n = 12, Fig. 1) to distinguish these units from clustering results obtained from genetic analyses. Finally, 
twenty-five dog samples (blood and tissue), to be used as a 
reference group for detecting possible wolf-dog hybrids, 
were collected in Poland from private owners and from 
individuals killed by vehicles.

**Laboratory methods**

DNA from faecal samples was isolated using the QIAamp 
DNA Stool Mini Kit (QIAGEN) and DNA from tissue 
samples was extracted with the DNAeasy tissue kit 
(QIAGEN).

For a portion of the samples collected in southern 
Poland (n = 185), DNA was isolated using a phenol– 
chloroform method (Sambrook and Russel 2001) with two 
additional phenol extractions at the beginning of the pro-
cedure, at the Institute of Nature Conservation PAS in 
Cracow, Poland. In order to reduce contamination risk, scat 
DNA extraction was carried out in a room dedicated to non-
invasive samples. Negative controls were included in each 
DNA extraction set to monitor for contamination.

Amplification of a 230 bp mtDNA fragment of the HV1 
region was performed using primers described by Savo-
lainen et al. (1997). Amplifications were carried out in 
10 µl reaction volumes containing 1U Taq polymerase, 
200 µM dNTP, 0.2 µM of primers, 0.2 µl of BSA (Fer-
mentas) plus 0.2 µl PCR Anti-inhibitor (DNA Gdańsk, 
Poland) and 2 or 3.6 µl of DNA extract, respectively, for 
tissue or scat samples. The reaction conditions were as 
follows: 2 min at 94 °C of initial denaturation, 30 cycles 
(tissue) or 36 cycles (scats) of 20 s at 94 °C, 30 s at 69 °C, 
40 s at 72 °C, and the final extension step for 10 min at 
72 °C. We subsequently amplified mtDNA sequences for 
each region using a portion of the samples identified as 
having distinct microsatellite profiles. PCR products were 
purified using Clean Up (A&A Biotechnology, Gdansk, 
Poland). Sequencing reactions were carried out in 10 µl 
Volumes using the Big Dye ver. 3.1 sequencing kit 
(Applied Biosystems) with the forward primer. Products 
were purified with the Exterminator kit (A&A Biotech-
nology) and separated on an ABI PRISM 3100 Genetic 
Analyser (Applied Biosystems). Sequencing results were 
analysed with ABI PRISM DNA Sequencing Analysis 
software and aligned by eye in BioEdit ver. 7.0.9. (Hall 
1999). Haplotypes were compared to previously recorded 
sequences from Pilot et al. (2006) using Collapse ver. 1.1 
(D. Posada, http://darwin.uvigo.es/software/collapse.html).

Microsatellite genotyping was performed using 11 poly-
morphic loci: FH2001, FH2010, FH2017, FH2054, FH2079, 
FH 2088, FH2096, FH2137, and FH2140 (Francisco et al. 
1996), C213 (Ostrander et al. 1993) and VWF (Shibuya et al. 
1994). Amplifications were carried out in four multiplex 
reactions using Multiplex PCR Kit (QIAGEN) and PCR 
conditions described in the manufacturer’s instructions with 
modifications by adding 0.1 µl BSA (Fermentas) and 0.1 µl 
PCR Anti-inhibitor (DNA Gdansk). Cycling was performed 
on a DNA Engine Dyad Peltier Thermal Cycler (BIO RAD) 
with the following profile: 16 cycles of 94 °C for 30 s, 58 °C 
for 90 s, 72 °C for 60 s, followed by 10 cycles of 94 °C for 
30 s, 57 °C for 90 s, 72 °C for 60 s and 10 cycles of 94 °C for 
30 s, 56 °C for 90 s, 72 °C for 60 s with a final extension of 
60 °C for 30 min, after an initial denaturation step of 95 °C 
for 15 min. Fragments were separated using genetic analyzer 
ABI3100 and allele lengths were determined with GENEM-
APPER 3.5 (Applied Biosystems) and GENEMARKER 1.51 
(SoftGenetics LLC). Each scat sample was amplified at least 
three independent times through a multiple-tube approach 
(Taberlet et al. 1996). We accepted alleles confirmed by a 
minimum of two independent PCR amplifications. Only 
individuals for which six or more loci had been successfully 
amplified were included in subsequent analyses.

Additionally, 66 samples were analysed at nine single 
nucleotide polymorphism (SNP) loci: 372M9, BLA22, 
BLB52, 168J14, 1C06, 38K22, 182B11, 218J14, 309N24, 
using the TaqMan Assay protocol described in Fabbri et al. 
(2012) at the Laboratory of Genetics at Istituto Superiore 
per la Protezione e la Ricerca Ambientale (ISPRA), Italy, 
to verify individual identification, with focus on samples 
genotyped at ≤9 loci. If not stated otherwise, all analyses 
were done at MRI PAS, Poland.

Probability of identity and test for the presence of wolf-
dog hybrids

We used GIMLET 1.3.3 (Valière 2002) to identify false 
hozygotes (drop-out alleles) and false alleles by com-
paring repeated genotypes, construct a consensus genotype 
from the sets of PCR repetitions for each sample of faeces, 
and identify multiple samples from the same individual. 
Multilocus genotypes were analysed with 25 reference dog 
genotypes to detect any dogs or wolf-dog hybrids using the 
Bayesian clustering procedure implemented in STRUC-
TURE (Pritchard et al. 2000) as outlined in previous 
analyses of wolves from Italy (Verardi et al. 2006; Randi 
2008). We ran STRUCTURE using $10^5$ iterations,
following a burn-in period of $10^4$ iterations, to assign individuals to clusters. Five independent runs were completed for each $K$ value (1–10). We verified that likelihood values converged for each run of $K$. All samples with membership $q_i < 0.90$ in the wolf cluster (in total 17 dogs and possible wolf-dog hybrids, not concentrated in any particular region) were excluded, and we continued analyses only with samples considered to be wolves.

Probability of identity (PI) for an increasing number of loci, i.e. the probability that different individuals share an identical multilocus genotype by chance, and the PI between sibs (PIsib) (Waits et al. 2001) were calculated in GenAlEx ver. 6.1 (Peakall and Smouse 2006). Additionally, the number of matches between wolf genotypes was calculated to determine the minimum number of loci needed for individual identification.

Statistical analyses

Genetic diversity statistics for mtDNA sequences including the number of haplotypes and polymorphic sites, haplotype and nucleotide diversity were estimated using DnaSP ver. 5 (Librado and Rozas 2009). To infer population genetic structure based on mtDNA haplotype frequency we used the spatial analysis of molecular variance implemented in the SAMOVA software (Dupanloup et al. 2002). This approach defines groups of populations that are geographically homogenous and maximally differentiated from each other. The method is based on a simulated annealing procedure that aims to maximize the proportion of total genetic variance due to differences between groups of populations. This approach requires the a priori definition of the number ($K$) of groups. We ran SAMOVA with $K$ ranging from 2 to 6. Each analysis was performed twice to check for consistency of results. In each run, 100 simulated annealing processes were performed. The recognition of the most probable number of groups was based on the pattern of changes in values of $\Phi$-statistic parameters with $K$.

Microsatellite variability statistics, including the number of alleles per locus and the number of private alleles, observed ($H_O$) and expected ($H_E$) heterozygosity, and tests for departures from Hardy–Weinberg equilibrium were calculated in GenAlEx. Allelic richness, which corrects the observed number of alleles for differences in sample size, and the inbreeding estimator Wright’s $F_{IS}$ were computed with FSTAT ver. 2.9.3 (Goudet 1995). To test for genetic differentiation among sampling regions, pairwise $F_{ST}$ values (Weir and Cockerham 1984) were calculated in Arlequin ver. 3.11 (Excoffier et al. 2005). Genetic differentiation between populations was characterized by exact tests (Raymond and Rousset 1995), using GENEPOP ver. 3.1. The levels of significance for multiple tests were adjusted by the sequential Bonferroni method (Rice 1989).

Two complementary Bayesian clustering algorithms, STRUCTURE ver. 2.3 (Pritchard et al. 2000) and GENELAND ver. 3.3 (Guillot et al. 2005), were applied to infer population structure and assign individuals to subpopulations (clusters) based on individual multilocus genotypes and—in GENELAND—their spatial location. The posterior probability of the data [LnP(D)] was estimated from four replicate runs for a number of groups $K = 1$ to 10, each with a burn-in of $10^4$ followed by $10^5$ iterations, using the admixture model and correlated allele frequencies (Falush et al. 2003). Inferences in GENELAND were done in a single step as recommended by the authors (Guillot 2008). This approach makes inferences faster and avoids the issue of ghost populations. Thus, we ran GENELAND 50 times allowing $K$ to vary from 1 to 15, with the following parameters: $20^5$ MCMC iterations, maximum rate of Poisson process fixed to 100, uncertainty attached to spatial coordinates fixed to 0.5°, and the maximum number of nuclei in the Poisson–Voronoi tessellation fixed to 300.

We computed the posterior probability of subpopulation membership for each pixel of the spatial domain and the modal subpopulation for each individual in all 50 runs. Finally, we examined all runs for consistency.

Subsequently, we evaluated population genetic structure performing principal component analyses (PCA) on individual genotypes from 4 subpopulations defined by STRUCTURE (with region 4 and 5 separated from regions 8–12 because of geographic distance and the GENELAND results) using the adegenet-package (Jombart 2008) in R 2.15.0 (R development Core Team 2012). The PCA does not assume Hardy–Weinberg equilibrium (HWE) and is highly efficient in revealing genetic structure in the form of clines, which is more difficult to detect than clusters (Jombart et al. 2009). We calculated pairwise genetic distances among wolves in the 12 regions based on mitochondrial and nuclear markers, and compared the results using the partial Mantel test in R with the vegan package (Oksanen et al. 2011).

Results

Genetic variability and structuring based on mtDNA

We found six mtDNA haplotypes (H1, H2, H3, H6, H8, H14—nomenclature consistent with Fig. 2 in Pilot et al. 2006) among 333 analysed samples. They are all known from previous studies (Vila et al. 1999; Randi et al. 2000; Jedrzejewski et al. 2005a; Pilot et al. 2006). H1, H2, H3 and H8 belong to haplogroup 1, which is widespread in north-eastern and central Europe and the Iberian Peninsula, whereas H6 and H14 belong to haplogroup 2 that dominates in south-eastern Europe and Italy (Pilot et al. 2010). The number of mtDNA haplotypes per
region ranged from one to four (Table 2). We observed no relationship between the number of samples and the number of haplotypes per region ($r^2 = 0.004$, $F_{1,10} = 0.038$, $P > 0.8$).

Notably, we found the highest number of haplotypes (4) in regions 4 ($n = 25$ individuals analysed) and 5 ($n = 10$ inds). The latter region also showed the highest number of polymorphic sites (Table 2). Conversely, samples from regions 2 ($n = 28$ inds) and 7 ($n = 27$ inds) revealed only one haplotype, which were H6 and H2, respectively.

Pairwise $\Phi_{ST}$ values between the 12 geographically predefined regions were generally very high (>0.25) and statistically significant, which indicated isolation among many regions according to the mtDNA (Table 3). Only seven of the 66 pairwise distance values were low (<0.05). These low values occurred among the regions located in the same part of the country within the lowlands and the Carpathian Mountains.

The results of SAMOVA indicated significant population genetic structure for each assumed number of groups. The highest increase in $\Phi_{CT}$ value occurred between $K = 3$ and $K = 4$ and all parameters of the $\Phi$-statistics stabilized from $K = 4$ (Appendix: Fig. 7). Thus, we assumed four subpopulations as the most parsimonious clustering configuration that maximized variation among groups. Wolves from the regions 1, 2, and 3 in the Carpathians formed subpopulation S1 (Fig. 2). Subpopulation S2 was comprised exclusively of individuals from region 4 in south-eastern Poland. Wolves from regions 5, 8, 10, 11, and 12 formed the geographically disjunct subpopulation S3. Finally, wolves from regions 6, 7, and 9 formed subpopulation S4. Subpopulation S1 in the Carpathian Mountains was strongly dominated by haplogroup 2 (97 % of the wolves), and four haplotypes from haplogroup 1 were present there at low frequencies. In contrast, subpopulations S2, S3 and S4 were primarily comprised of individuals carrying haplotypes from the previously identified haplogroup 1 (S2—88 %, S3—96 %, and S4—99 %) and included haplogroup 2 at very low frequencies (Fig. 2).

Pairwise $\Phi_{ST}$ values among four subpopulations (S1–S4) were all very high (>0.25), which indicates little or no mtDNA gene flow among them (Appendix: Table 5).

Genetic differentiation and structuring based on microsatellites

We identified a total of 457 wolf genotypes. The allelic drop-out rate varied from 0.079 to 0.360 among regions. The highest values were observed for the loci FH2010, FH2017 and FH2079 (Appendix: Table 6), which had

![Fig. 2 Four subpopulations (S1–S4) of wolves in Poland, based on mtDNA, delimited by SAMOVA (left panel) and haplotype frequencies for subpopulations (right). Sampling information is provided in Table 1](image)
Fig. 3 Results of STRUCTURE analysis for Polish wolves, based on 11 microsatellite loci, assuming $K = 2$ to 5 subpopulations of wolves. Black lines separate wolves from different sampling regions. Sampling information is provided in Table 1.

Table 2 Basic genetic variability parameters for wolves from 12 regions in Poland and eastern Germany genotyped at 11 microsatellite loci

| Parameter                        | Region | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|----------------------------------|--------|---|---|---|---|---|---|---|---|---|----|----|----|
| **Mitochondrial DNA**            |        |   |   |   |   |   |   |   |   |   |    |    |    |
| No. of haplotypes                |        | 3 | 1 | 3 | 4 | 4 | 3 | 1 | 2 | 2 | 2  | 2  | 2  |
| No. of polymorphic sites         |        | 8 | 0 | 10| 9 | 13| 10| 0 | 6 | 6 | 6  | 6  | 6  |
| Haplotype diversity (SD)         |        | 0.063| – | 0.123| 0.473| 0.733| 0.310| – | 0.313| 0.363| 0.363| 0.524| 0.133| 0.458 |
| Nucleotide diversity (SD)        |        | 0.002| – | 0.004| 0.011| 0.020| 0.009| – | 0.008| 0.010| 0.014| 0.004| 0.012 |
| **Microsatellites**              |        |   |   |   |   |   |   |   |   |   |    |    |    |
| Average no. of alleles/locus     |        | 6.000| 6.000| 5.909| 5.636| 4.818| 6.182| 4.909| 5.818| 4.273| 5.364| 5.636| 6.636|
| Allelic richness                 |        | 4.147| 4.561| 4.410| 4.478| 4.635| 4.163| 3.903| 4.294| 3.994| 4.508| 4.676| 4.769|
| Average number of private alleles|        | 0.091| 0.091| 0.091| 0.091| 0.091| 0.091| 0.091| 0.364| – | 0.182| 0.455|    |
| $H_O$                            |        | 0.524| 0.576| 0.557| 0.517| 0.538| 0.537| 0.513| 0.507| 0.452| 0.545| 0.475| 0.520|
| $H_E$                            |        | 0.650| 0.644| 0.626| 0.670| 0.671| 0.652| 0.629| 0.651| 0.628| 0.689| 0.661| 0.688|
| $F_{IS}$                         |        | 0.203| 0.122| 0.126| 0.242| 0.243| 0.186| 0.197| 0.235| 0.310| 0.228| 0.307| 0.260|

Figure 1 and Table 1 show sampling regions and sample sizes. $H_O$, observed heterozygosity; $H_E$, expected heterozygosity; $F_{IS}$, inbreeding coefficient.
fragment length >200 bp and thus higher probability of drop-out for low quality-samples (Broquet et al. 2007). Based on all 11 loci, the cumulative PI of all genotypes was $1.35 \times 10^{-11}$ and the probability of identity between sibs (PISib) equaled $5.0 \times 10^{-5}$ (detailed data available upon request). Data from seven microsatellites were sufficient to distinguish all 457 genotypes with PI = $3.13 \times 10^{-7}$. We used three samples genotyped at 6 loci. They were assigned to the region of origin, and so their inclusion is unlikely to have biased our results. SNP data for the 66 samples with ≤9 microsatellite loci amplified confirmed individual identifications with cumulative PI = 0.002 and PI-sibs = 0.042. Microsatellite data for these 66 samples were therefore included in subsequent analyses.

The 11 loci were polymorphic in all regions and the average number of microsatellite alleles per locus varied between 4.27 and 6.64 (Table 2). Observed heterozygosity ($H_O$) values (range 0.45–0.58) were lower than expected ($H_E$, range 0.63–0.69). In the total population of Polish wolves we found significant deviation from HWE at all 11 loci (Appendix: Table 6). We also tested for HWE in subpopulations defined by STRUCTURE (Fig. 3). Four to seven of 11 loci (depending on subpopulation) showed significant deviation from HWE (after correcting for multiple tests), except for subpopulation 5 (regions 8–12) where 10 loci showed significant deviation. Only locus FH2137 displayed consistent deviation from HWE across all subpopulations, the results for other loci varied among subpopulations. $F_{IS}$ values ranged from 0.122 to 0.310 and were not correlated to the number of wolf genotypes per group ($r^2 = 0.15$, $F_{1,10} = 1.779$, $P > 0.2$). We found no relationship between the number of mtDNA haplotypes and the number of microsatellite alleles per locus in the 12 regions (Pearson $r = 0.03$, $F_{1,10} = 0.002$, $P > 0.9$). Individuals from regions 5, 9, and 10 did not show private alleles (Table 2). The highest number of private alleles was found in region 12 (western Poland and eastern Germany), which comprised 57
Pairwise genetic distances (above diagonal $\Phi_{ST}$, based on mtDNA and below diagonal $F_{ST}$, based on microsatellite markers) wolves sampled from 12 regions in Poland and eastern Germany. All $\Phi_{ST}$ and $F_{ST}$ values (except those marked NS) were statistically significant at $P < 0.001$ after Bonferroni correction ($k = 66$).

| Region | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|--------|---|---|---|---|---|---|---|---|---|----|----|----|
| 1      | * | 0.002 NS | -0.007 NS | 0.743 | 0.565 | 0.814 | 0.935 | 0.831 | 0.846 | 0.777 | 0.897 | 0.768 |
| 2      | 0.052 | * | 0.011 NS | 0.743 | 0.584 | 0.818 | 1.000 | 0.844 | 0.875 | 0.769 | 0.954 | 0.756 |
| 3      | 0.055 | 0.044 | * | 0.681 | 0.438 NS | 0.763 | 0.931 | 0.776 | 0.790 | 0.698 | 0.869 | 0.698 |
| 4      | 0.087 | 0.109 | 0.127 | * | 0.317 | * | 0.614 | 0.711 | 0.575 | 0.564 | 0.481 | 0.639 | 0.509 |
| 5      | 0.102 | 0.085 | 0.095 | * | 0.049 NS | * | 0.465 | 0.747 | 0.248 NS | 0.363 NS | 0.169 NS | 0.349 NS | 0.180 NS |
| 6      | 0.212 | 0.166 | 0.202 | 0.139 | 0.096 | * | 0.109 NS | 0.560 | -0.042 NS | 0.229 NS | 0.687 | 0.388 |
| 7      | 0.160 | 0.171 | 0.205 | 0.067 | 0.064 NS | 0.156 | * | 0.803 | 0.246 NS | 0.538 | 0.949 | 0.633 |
| 8      | 0.136 | 0.111 | 0.144 | 0.075 | 0.069 | 0.108 | 0.111 | * | 0.498 NS | 0.132 NS | 0.009 NS | 0.01 NS |
| 9      | 0.121 | 0.111 | 0.176 | 0.065 | 0.099 | 0.119 | 0.095 | 0.047 | * | 0.129 NS | 0.672 NS | 0.298 NS |
| 10     | 0.163 | 0.153 | 0.176 | 0.082 | 0.052 NS | 0.066 | 0.091 | 0.055 | 0.071 | * | 0.288 NS | 0.003 NS |
| 11     | 0.149 | 0.129 | 0.172 | 0.053 | 0.063 | 0.063 | 0.081 | 0.076 | 0.043 NS | 0.025 NS | * | 0.130 NS |
| 12     | 0.159 | 0.132 | 0.184 | 0.078 | 0.076 | 0.052 | 0.114 | 0.080 | 0.054 NS | 0.040 | 0.01 NS | * |

Values of low level of differentiation (<0.05) are in italics, moderate (0.05–0.15) in regular font, high (>0.15), and very high (>0.25) are in bold.
(GENELAND, STRUCTURE with K = 5). We found concordance between pairwise genetic distances among wolves in 12 regions based on mitochondrial and nuclear markers (Fig. 6). A partial Mantel test (Smouse et al. 1986) where the geographic distance matrix was held constant, while the relationship of $F_{ST}$ and $\Phi_{ST}$ was determined, showed a significant positive correlation of the two measures of genetic distance ($r = 0.521$, $P = 0.001$, test with 999 permutations).

**Discussion**

Are wolf populations structured across Poland?

We found significant genetic structure between wolf populations in the Carpathians and the Polish lowlands. Our mtDNA results seem to correspond with those reported from earlier research encompassing a larger area of eastern Europe. A Carpathian subpopulation indicated in Pilot et al. (2006) corresponds with subpopulation S1 identified in our study. Similarly, S2 corresponds with a subpopulation spanning from southeastern Poland through southern Belarus and northern Ukraine into Russia. S3 and S4 appear to be part of a subpopulation extending from northeastern Poland and eastward into Russia (Pilot et al. 2006). A subsequent study examined the historical distribution of haplogroups in (primarily) eastern Europe (Pilot et al. 2010). These findings suggest that most haplotypes from S1 belong to haplogroup 2 from southern Europe, whereas S2, S3, and S4 are principally composed of wolves from haplogroup 1. The present contact zone between the two lineages in central Europe appears to be a result of one haplogroup (1) partially replacing a more ancient haplogroup (2) that had been predominant over the last several millennia and remains widely distributed in southern Europe (Leonard et al. 2007; Pilot et al. 2010). The presence of mainly one haplotype (H6) from haplogroup 2 in the Polish Carpathians indicates low genetic diversity at its current northern range. Consequently, Poland can be seen as a meeting zone between two lineages of wolves. Despite the geographical proximity well within wolf dispersal distance (e.g. Wabakken et al. 2007), our results suggest very limited gene flow between the two areas and, hence, a restricted contact zone between the north and the south. Pilot et al. (2006) observed a single population cluster encompassing northern and southern Polish wolves. Our microsatellite results showed strong genetic structure between the Carpathians and the Polish lowlands, and thus demonstrate additional substructuring within Poland when compared with previous findings. The observed deviation from HWE may be due several reasons: the presence of null alleles, high allelic dropout rate in non-invasive studies, the existence of local genetic structure (Wahlund’s effect), lack of random mating, or the presence of closely related individuals (members of the family groups) in a sample. Although genotyping error and null alleles may have contributed to the excess of homozygotes, these factors are unlikely to explain the observed mountain-lowland structure in Polish wolves. Other studies on wolf population structure (that used tissue samples) also showed significant heterozygote deficit and positive values of $F_{IS}$, which was explained by moderate inbreeding, the presence of closely related individuals, or the presence of additional undetected structure (Lucchini et al. 2004, Pilot et al. 2006, Jansson et al. 2012).

In addition to the north–south structure, our results indicated clustering within the lowland area. $F_{ST}$ values between small groups may reflect social structure and can show relatively high values between local family groups. For example, Thiessen (2007) reported between-pack differentiation of $F_{ST} = 0.179$ based on $n = 36$ wolf packs in western Canada. Wolves typically live in social and territorial groups of 2–11 individuals (Fuller et al. 2003; Jędrezewski et al. 2010). Although wolf social structure may have contributed to the high $F_{ST}$ value (0.156) observed between region 6 ($n = 55$) and region 7 ($n = 45$), the samples from these regions represent members of multiple packs (Jędrezewski et al. 2004). Consequently, wolf pack structure alone is not expected to produce such high $F_{ST}$ values. The moderate $F_{ST}$ values (0.05–0.15) observed between a number of lowland regions require further investigation, as well as the high $F_{ST}$ value (0.156) seen between wolves in regions 6 and 7 that are separated by <50 km. Assessment of samples from contiguous regions, landscape features, and additional genetic markers could improve understanding of the extent to which these differences might be explained by landscape fragmentation or ecological differences resulting in natal habitat-biased dispersal (Geffen et al. 2004; Sacks et al. 2004; Pilot et al. 2006).

Factors that could maintain divisions between genetically distinct wolf populations in Poland

MtDNA and microsatellite results consistently showed differentiation between wolves in the Carpathian Mountains and the Polish lowlands. We examined only non-coding fragments of DNA, and future analyses of genes under selection (using e.g. SNP mapping within exons or regulatory sequences of functional genes) could help clarify whether adaptive genetic differences might play a role in the observed structure. Associations between environmental factors (habitat type, climate, prey abundance) and genetic variants might have resulted in the development of wolf ecotypes adapted to different habitats (e.g. Carmichael et al.
This is consistent with the findings that (1) ecological factors (habitat, prey, climate) appear to explain much of the spatial variation in wolf genetic diversity in east-central Europe (Pilot et al. 2006), and that (2) wolves from three genetically distinct populations in Poland show significant differences in prey composition and prey preferences (Jędrzejewski et al. 2012). In northeastern Poland, wolves prey on four ungulate species (moose, red and roe deer, wild boar), although only red deer was killed in a higher proportion than that expected from availability. In eastern Poland (regions 4 and 5) wolves preferred roe deer, whereas in the south-east (regions 1–3) they hunted mostly red deer and only a small proportion of roe deer and wild boar occurred in their diet (Jędrzejewski et al. 2012).

The genetic structure of wolves in Poland shows strong and abrupt spatial division, and the primary separation occurs between lowland and Carpathian Mountain wolves. Within the lowlands, both mtDNA and microsatellite results indicate a further division between region 4 (Roztocze) in the southeast and northern Poland. This appears to concur with previous mtDNA and microsatellite results that supported the presence of a subpopulation extending from southeastern Poland and eastward into Russia (Pilot et al. 2006). Interestingly, separation of Carpathian and lowland wolves by a ‘wolf-free belt’ has also been suggested to occur in Ukraine (Gursky 1985). Wolves are highly mobile (mean daily movement distance in Polish wolves: 23 km, max 64 km, Jędrzejewski et al. 2001). Moreover, long-distance dispersers can travel several hundred kilometers (Wabakken et al. 2001, 2007; Schede et al. 2010), so individuals from different populations should be able to meet and interbreed. The observed genetic isolation of Carpathian wolves is thus surprising given the species’ remarkable dispersal ability. Several factors could nevertheless contribute to the observed divisions. First, analyses of habitat structure in and around wolf ranges conducted in southern and northern Poland (Jędrzejewski et al. 2004, 2005a) showed very rapid deterioration of habitat connectivity immediately north of the Carpathians. In contrast, northeastern Poland provides better wolf habitat because the human density and network of transportation infrastructure is lower here than in southern Poland (Jędrzejewski et al. 2004, 2005a). In southern Poland, wolves occurred only in a narrow belt of <100 km, whereas in northern Poland stable wolf populations persist more than 200 km from the continuous wolf range (Jędrzejewski et al. 2004, 2005a).

Huck et al. (2010, 2011) analysed dispersal costs among patches of wolf habitat and modeled dispersal corridors in Poland, and found that dispersal from the Carpathians to any other patch would be much more costly than dispersal among other regions of the country. Densely populated and urbanized areas in southern Poland along the Carpathians may act as a serious barrier to wolf movement and limit wolf dispersal (Huck et al. 2011). Moreover, modeling of suitable wolf habitat showed that the eastern portion of Poland was already ‘filled’ by wolves (Jędrzejewski et al. 2008). Western Poland still has much suitable wolf habitat not occupied at present, which could support a large population of wolves (Jędrzejewski et al. 2008). Hence, dispersers would likely prefer to settle in western Poland than the more saturated east.

The current landscape structure and wolf distribution in Poland are important factors that are likely to limit gene flow between Carpathian and lowland wolves. However, wolves can move through highly heterogeneous and human dominated landscapes (Blanco and Cortes 2007) and cross a range of natural and anthropogenic barriers (Blanco et al. 2005; Wabakken et al. 2007; Ciucci et al. 2009). Other factors might therefore contribute to the observed structure. Natal-habitat biased dispersal seems a possible explanation in the case of Carpathian wolves, and is consistent with findings from Europe and North America (Geffen et al. 2004; Sacks et al. 2004; Pilot et al. 2006). Population genetic structure consistent with the presence of highland and lowland habitats has been reported in coyotes (Sacks et al. 2004, 2005) and merits further attention in wolf populations from the Carpathians Mountains and surrounding lowland areas. Differences in the legal status and protection of the species in the Carpathians might also influence genetic structure, as only wolves in the Polish and Czech parts of the north-western edge of the Carpathian Mountains are protected. In Slovakia and Ukraine, wolves are regularly hunted or persecuted. This probably causes a source-sink effect and thus dispersal southward from the Polish to the Slovakian and Ukrainian parts of the Carpathian Mountains, where dispersers repopulate vacant territories (Nowak et al. 2008).

Recolonization of western Poland and eastern Germany

Our results suggest that wolves colonizing western Poland and eastern Germany primarily originate from northeastern Poland. In particular, it appears that westward dispersal from regions 10 and 11 has been relatively frequent. The location of these sampling regions on the western border of the established wolf range in northeastern Poland, and the relatively contiguous forest habitat in this area (Huck et al. 2011) suggest that regions 10 and 11 represent a natural starting point for westward expansion (Jędrzejewski et al. 2008; Huck et al. 2010, 2011). Wolves in western Poland and eastern Germany appear to represent the expanding western edge of a vast, northeastern European wolf population that primarily inhabits boreal and temperate forests.
and extends through the Baltic States, northern Belarus, and northwestern Russia (Pilot et al. 2006, 2010).

Importantly, our study detected private alleles in region 12 (western Poland and eastern Germany). Although these alleles might be present in unsampled northeastern Polish wolves, the most likely explanation is immigration from areas not covered by our investigation. East-European countries (Belarus, Latvia) harbour the region’s largest wolf populations, although human harvest is high and potentially unsustainable in some areas (Jędrzejewski et al. 2010), which might affect source-sink dynamics (Jędrzejewski et al. 2005b). Recently documented movements of a radio-collared wolf between Germany and Belarus further support such dispersal (Schede et al. 2010). The apparent recolonization of western Poland and eastern Germany from various source populations should help ensure high levels of genetic variation and subsequent potential for adaptation to new and altered environments.

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Appendix

See Tables 4, 5, 6 and Fig. 7.

| K | L(K) | SD | ΔK |
|---|------|----|----|
| 1 | −15255 | 0.447 | 
| 2 | −14246 | 2.588 | 69.637 |
| 3 | −13808 | 6.457 | 27.913 |
| 4 | −13525 | 7.014 | 25.698 |
| 5 | −13278 | 3.435 | 52.473 |
| 6 | −13248 | 217.289 | 0.830 |
| 7 | −13029 | 90.833 | −0.014 |
| 8 | −12846 | 27.318 | 6.598 |
| 9 | −12729 | 32.935 | 5.473 |

In bold the most probable number of subpopulations and the highest values of ΔK

| Subpopulation | S1 | S2 | S3 |
|---|---|---|---|
| S2 | 0.81 | | |
| S3 | 0.73 | 0.50 | |
| S4 | 0.86 | 0.70 | 0.49 |

| Locus | H₀ | Hₑ | NA | Aᵣ | Fᵢₛ | ADO (SD) |
|---|---|---|---|---|---|---|
| FH2088 | 0.549 | 0.796 | 7 | 7.000 | 0.310 | 0.139 (0.082) |
| FH2017 | 0.382 | 0.654 | 7 | 6.990 | 0.417 | 0.222 (0.235) |
| FH2010 | 0.439 | 0.668 | 7 | 7.000 | 0.342 | 0.222 (0.091) |
| FH2054 | 0.642 | 0.797 | 13 | 12.928 | 0.195 | 0.148 (0.078) |
| FH2096 | 0.412 | 0.600 | 4 | 4.000 | 0.314 | 0.202 (0.081) |
| FH2079 | 0.428 | 0.653 | 12 | 12.000 | 0.345 | 0.245 (0.183) |
| VWF | 0.667 | 0.812 | 15 | 14.773 | 0.178 | 0.131 (0.081) |
| FH2001 | 0.589 | 0.727 | 6 | 6.000 | 0.190 | 0.182 (0.101) |
| C213 | 0.564 | 0.833 | 15 | 15.000 | 0.323 | 0.139 (0.116) |
| FH2137 | 0.537 | 0.782 | 15 | 14.890 | 0.313 | 0.160 (0.093) |
| FH2140 | 0.581 | 0.737 | 12 | 11.873 | 0.211 | 0.113 (0.056) |

All loci deviated significantly from HWE and all values of Fᵢₛ were statistically significant.

See Tables 4, 5, 6 and Fig. 7.

Table 4 Results of calculations for the number of genetic clusters indicated by STRUCTURE using the method of Evanno et al. (2005)

Table 5 Pairwise genetic distances for haplotypic data (Φₑₛ) between wolf subpopulations defined by SAMOVA

Table 6 Expected (Hₑ) and observed (H₀) heterozygosity, number of alleles (NA), allelic richness (Aᵣ), inbreeding coefficients (Fᵢₛ) and mean allelic dropout rate (ADO) in the studied Polish wolf population
Fig. 7 Changes in $\Phi$-statistics for $K = 2$ to 6 subpopulations of wolves in Poland, on the basis of mtDNA and inferred from SAMOVA. $\Phi_{SC}$—proportion of the variance among local populations within groups. $\Phi_{ST}$—proportion of the variance among local populations within the total population. $\Phi_{CT}$—proportion of the total variance explained by the grouping

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