#### **RESEARCH ARTICLE**

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# Non-invasive genetic study of the endangered Cantabrian brown bear (Ursus arctos)

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9 Abstract The Brown Bear (Ursus arctos) population 10 present in the Cantabrian Mountains has suffered a dra-11 matic decline in recent centuries and is now threatened 12 with extinction. This situation has led to the development 13 and implementation of a species recovery plan. To 14 accomplish this plan, we need to improve our knowledge 15 about the ecology, demography and genetics of this pop-16 ulation. This paper presents the genetic analysis of the 17 Cantabrian brown bear population using non-invasive 18 samples (faeces and hairs) collected between 2004 and 19 2006. It was necessary to optimize a set of 18 microsatellite 20 loci and a sex marker (several new multiplex reactions 21 were developed) to obtain a suitable probability of identity 22 among genotypes to work with this small, deeply structured 23 population. Genotyping of 48 individuals was carried out 24 using a two-step PCR protocol to increase the quality of the 25 multilocus genotypes. Validation of genotypes was per-26 formed using a multi-tube approach combined with 27 different software programmes to measure their error rate 28 and reliability. Diversity in the Cantabrian population was

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low ( $H_e = 0.51$ ) and the population was markedly subdi-29 vided into two subpopulations (western and eastern) 30 without current gene flow between them. The level of 31 divergence between the two subpopulations ( $F_{st} = 0.41$ ) 32 and the extremely low diversity in the eastern group 33  $(H_e = 0.25)$  indicate that this has had an extremely low 34 effective population size and had been isolated from the 35 main group during the last century. Connectivity between 36 the two subpopulations will be of prime importance for the 37 long-term survival of this species in the Cantabrian 38 Mountains. 39

Keywords Ursus arctos · Microsatellites ·	41
Non-invasive samples · Individual identification ·	42
Population structure	43
	44

## Introduction

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Brown bears were once found throughout Europe and even 46 inhabited the British Isles until the tenth century. However, 47 since the mid-1800's, populations in Europe have been 48 49 severely reduced due to habitat destruction and overexploitation by humans (Servheen 1990). Four very small, isolated 50 populations, all of which are endangered, can still be found 51 in southern and western Europe (the Pyrenees, Southern 52 53 Alps, Cantabrian Mountains and Apennine Mountains), 54 representing the remnants of a once widespread brown bear 55 population (Zedrosser et al. 2001). The Cantabrian population is found in two areas of the Spanish Cantabrian 56 57 Mountains separated by 30-50 km of mountainous terrain. Interchange between both subpopulations is unlikely (Naves 58 and Nores 1997), due to unsuitable habitat and a high speed 59 railway and motorway that bisect the area (Fig. 1). The 60 population estimate for the western subpopulation is around 61

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Fig. 1 Distribution map of the brown bear in Europe: (a) present distribution of the brown bear in Europe, and (b) distribution of the brown bear in the Cantabrian Mountains (based on Naves et al. 2003) and location of the 133 samples analysed in this study. White squares represent samples amplified for more than 14 markers, black squares represent samples with less than 14 loci amplified



50–60 individuals (Wiegand et al. 1998). The eastern Cantabrian Mountains subpopulation is estimated to contain
around 20 bears (Naves et al. 1999). Both Cantabrian bear
populations face similar conservation problems.

66 Mitochondrial DNA studies have shown that the Can-67 tabrian population belongs to the Iberian refugia clade of the western lineage of European brown bears. Only the 68 69 populations of the Cantabrian and Pyrenean Mountains and 70 small populations from the south of Sweden and Norway 71 belong to this clade (Taberlet and Bouvet 1994). It is 72 important to preserve this population if we wish to main-73 tain the most ancient lineage of the European brown bear.

74 In order to design an effective conservation plan, it is 75 necessary to understand the structure of threatened popu-76 lations, particularly those which, like this one, exist in 77 degraded or fragmented habitats (Lande and Barrowclough 78 1987; Simberloff 1988; Hanski and Gilpin 1997; Taylor 79 and Dizon 1999; Kraaijeveld-Smith et al. 2005). It is well 80 known that small populations are more vulnerable to 81 genetic factors, demographic and environmental stochas-82 ticity, genetic drift and inbreeding and have an increased 83 probability of extinction (Soulé 1987). When small popu-84 lations become fragmented and migration between 85 subpopulations decreases or is eliminated, consequent 86 increases in inbreeding and loss of genetic diversity can 87 have serious negative effects on the long-term viability of population fragments and, by extension, of the population 88 89 as a whole (Keller et al. 1994; Lacy 1997; Bjilsma et al. 90 2000; Sherwin and Moritz 2000; Coulon et al. 2004).

To avoid the extinction of this species in the Cantabrian 91 92 Mountains, the different regional governments from the areas where it is present have developed plans for the 93 conservation of brown bears since 1989. These plans have 94 95 been coordinated at a national level. In order to be effective, however, an action plan should be based on reliable 96 97 biological data, such as trustworthy estimates of population 98 size, population genetic status and connectivity with other 99 populations (Bellemain et al. 2007).

To assess the genetic status of the Cantabrian population 100 and provide guidelines for the conservation and manage-101 ment of this population, we used non-invasive genetic 102 techniques which have been successfully used in other 103 studies (Taberlet et al. 1997; Kohn et al. 1999; Woods 104 et al. 1999; Waits et al. 2000; Frantz et al. 2003; Smith 105 et al. 2006; Bellemain et al. 2007). However, this non-106 107 invasive approach has a major drawback, namely the problems associated with low DNA quantity and quality 108 (Taberlet and Luikart 1999). Under these circumstances, 109 the probability of critical genotyping errors (primarily 110 allelic dropout and false alleles) is high (Taberlet and 111 Luikart 1999; Taberlet et al. 1996, 1999). The solutions 112 proposed in these studies for addressing and reducing the 113 severity of genotyping errors and increasing the reliability 114 of genotypes include protocols for replicating amplification 115 and programmes for determining the magnitude of error in 116 117 a dataset and for calculating the reliability of genotype data (Taberlet et al. 1996; Miller et al. 2002; Bonin et al. 2004; 118 McKelvey and Schwartz 2004). 119 120 The goal of the present study was to ascertain the cur-121 rent genetic status of the Cantabrian brown bear population 122 using non-invasive genotyping techniques. The results of 123 this work will help us to provide guidelines for the con-124 servation and management of this population. To achieve 125 this goal, we optimized a set of suitable loci microsatellite 126 markers for carrying out population and individual identi-127 fication studies in this population.

## 128 Methods

#### 129 Sampling

130 This study presents the genetic data for the Cantabrian 131 brown bear population using non-invasive samples (faeces 132 and hairs) collected in the field between 2004 and 2006 using two sampling methods: opportunistic and systematic. 133 Most of the samples (n = 106) were collected in an 134 135 opportunistic manner from daily routine field work of 136 rangers and field biologists. Some samples (n = 27) were 137 collected through systematic surveys carried out seasonally 138 (summer: mid August-mid September; autumn: mid October-mid November) over a grid (2.5 km<sup>2</sup>) in the 139 western subpopulation in an area of roughly 750 km<sup>2</sup> 140141 covering around 630 km of on-foot surveys each season. 142 Although we were mainly looking for scats, some hair 143 samples were also collected. Since we wished to cover the 144 maximum area of distribution possible, we had to include 145 many opportunistic samples because the systematic surveys 146 only cover a partial area of the species distribution. 147 Figure 1b shows where the samples were collected.

148 Faeces samples were dry stored with silica after ethanol 149 soaking upon collection following Nsubuga et al. (2004) 150 and Roeder et al. (2004). Hairs were placed in individual 151 envelopes, no further manipulation being necessary until 152 extraction. A total of 133 samples were analysed: 88 from the western subpopulation and 45 from the eastern sub-153 154 population. Together with these non-invasive samples, 13 155 tissue/blood samples belonging to either dead or captured 156 individuals were used (all from the western subpopulation, 157 eight of which were found dead between 1989 and 2002). 158 The numbers from each type of sample used for the study 159 are listed in Table 1.

- 160 DNA extraction and typing
- 161 DNA extraction

162 DNA was extracted from faeces samples using the 'Qiamp
163 DNA stool kit' (Qiagen, Hilden, Germany) specially
164 developed for this type of material and following the
165 manufacturer's instructions. The only change introduced

 Table 1
 Number of samples analysed and percent of the total (between parentheses)

	Faeces	Hair	Skin/ blood	Total
No. DNA	25 (27.2%)	0 (0%)	1 (7.7%)	26 (17.8%)
Discarded genotypes <sup>a</sup>	27 (29.3%)	6 (14.6%)	3 (23.1%)	36 (24.7%)
Accepted genotypes <sup>b</sup>	40 (43.5%)	35 (85.4%)	9 (69.2%)	84 (57.5%)
Total number samples	92	41	13	146
a				

<sup>a</sup> Include all the incomplete genotypes (with less than 14 loci amplified)

<sup>b</sup> Genotypes with more than 14 loci amplified (all the hair, skin/blood samples and 24 of the faeces samples in this category were genotyped for the 18 loci)

was the incubation of the samples in ASL buffer ON at 166 25°C. For hair DNA extraction, 5–10 hair roots were used; 167 these were incubated ON with PCR buffer and proteinase 168 K (Allen et al. 1998; Vigilant 1999). Finally, tissue/blood 169 samples were extracted using the 'DNeasy Tissue kit' 170 (Qiagen, Hilden, Germany) following the manufacturer's 171 instructions. All hair and faeces extractions were carried 172 out in a separate room used exclusively for processing 173 samples of this type. 174

For the genotyping, we used a total of 18 microsatellite 176 markers (G1A, G1D, G10B, G10C, G10J, G10L, G10O, 177 G10P, G10X (Paetkau and Strobeck 1994; Paetkau et al. 178 1995) and MU05, MU09, MU10, MU23, MU50, MU51, 179 MU59, MU61, MU64 (Taberlet et al. 1997)), as well as the 180 sex marker SRY (Bellemain and Taberlet 2004), all of 181 which were selected from markers previously used in 182 genetic tracking of Pyrenean brown bears (Taberlet et al. 183 1997). 184

DNA amplifications were performed in a two-step PCR 185 following Taberlet et al. (1997) for faeces samples and 186 using only one PCR for hair and tissue/blood samples. To 187 avoid the problem of running out of DNA template 188 (because of the low amount of DNA available) before 189 completing the genotyping for all the microsatellite 190 markers, the amplification was carried out following the 191 multiplex preamplification method (Piggot et al. 2004; 192 Bellemain and Taberlet 2004). This method was optimized 193 for six microsatellite loci and for the sex marker (MU10, 194 MU23, MU50, MU51, MU59, G10L and SRY) by Belle-195 main and Taberlet (2004). We designed three new 196 multiplex PCRs to amplify all the remaining loci 197 (MU64 + G1A + G10C + G10P)MU61 + G10J + -198 G100 + G10X, MU05 + MU09 + G1D + G10B). A 199

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200 total number of four-first-step PCRs, where we amplified 201 between 4 and 6 loci simultaneously, and nine-second-step 202 PCRs, where the number of loci amplified simultaneously 203 varied between 1 and 3, were sufficient to amplify the 18 204 loci microsatellites and the sex marker (Table 2).

205 For faeces samples, the first-step PCRs or preamplifi-206 cations were prepared in a 25 µl volume containing 5 µl 207 template DNA. 0.01 uM of each primer and 12.5 ul of 208 "Qiagen Multiplex PCR Kit" (Qiagen, Hilden, Germany). 209 The second-step, PCRs or amplifications were prepared in 210 a 13 µl volume containing 3 µl preamplified product, 0.1 mM of each dNTP, 0.5 µM of each primer, 2 mM MgCl<sub>2</sub>, 0.5 U Taq DNA Polymerase form Quiagen (Qia-212 213 gen, Hilden, Germany) and  $1 \times Taq$  Quiagen buffer 214 (Qiagen, Hilden, Germany). Amplifications were per-215 formed on a GeneAmp PCR 9600 (Applied Biosystems) 216 under the following conditions: for the first-step PCR 15 min at 95°C, 40 cycles composed of 30 s denaturing at 218 94°C, 90 s annealing at 60°C, 1 min extension at 72°C, and 219 as a final extension step, 30 min at 60°C. For the second-220 step PCR, 3 min at 94°C, 35 cycles composed of 30 s denaturing at 94°C, 30 s annealing at 60°C, 1 min exten-222 sion at 72°C, and as a final extension step, 7 min at 72°C.

For hair and tissue/blood samples, nine-one-step PCRs were carried out using the primers of the second-step PCRs in a 20 µl volume containing 5 µl (2 µl for tissue/blood samples) template DNA, 0.5 µM of each primer and 10 µl of "Qiagen Multiplex PCR Kit" (Qiagen, Hilden, Germany). Amplifications were performed on a GeneAmp 242

PCR 9600 (Applied Biosystems) under the following 229 230 conditions: 15 min at 95°C, 40 cycles composed of 30 s denaturing at 94°C, 90 s annealing at 60°C, 1 min exten-231 sion at 72°C, and as a final extension step, 30 min at 60°C. 232

PCR products were checked in a 2% agarose gel and the 233 234 product diluted between 0 and 100 times depending on the intensity of the signal. One microlitre of this product was 235 added to a 12 ul mix of formamide and ROX 400HD 236 (12:0.2) and then loaded on an automatic sequencer 237 238 ABI310 (Applied Biosystems). Microsatellite patterns were examined both visually and using GENESCAN ANALY-239 SIS 3.1 and GENOTYPER 2.5 software (Applied 240 Biosystems). 241

#### Reliability of genotyping results

To test the reliability of our genotyping results and to 243 reduce tracking error in our dataset, we followed different 244 recommendations already proposed in different studies 245 addressing non-invasive genotyping. The suggestions by 246 Bonin et al. (2004) for limiting potential errors in the 247 genotyping process were followed. All the genetic typing 248 was performed using a combination of the multi-tube 249 approach and software packages that assign a reliability 250 value to each multilocus genotype. Three positive PCRs 251 (for both types of non-invasive samples, hair and faeces) 252 were first analysed, a consensus genotype was assigned 253 254 using the GIMLET v.1.3.2 software (Valière 2002) and its reliability was tested using the RELIOTYPE software 255

Table 2 Observed number of         alleles (A) with the number of	Locus (i,j)	Eastern subpopulation $(n = 8)$		= 8)	Western sub	Western subpopulation $(n = 39)$		
single alleles of each		A (Ae)	H <sub>e</sub>	H <sub>o</sub>	A (Ae)	H <sub>e</sub>	H <sub>o</sub>	
parentheses (Ae), observed	MU10 (1,1)	2 (0)	0.50	0.50	4 (2)	0.42	0.28*	
heterozygosity $(H_o)$ , Nei's	G10L (1,1)	1 (0)	0.00	0.00	6 (5)	0.74	0.64*	
estimated heterozygosity $(H_e)$ and deviations from Hardy-	U50 (1,2)	1 (0)	0.00	0.00	5 (4)	0.67	0.69	
Weinberg equilibrium (HWE)	MU23 (1,2)	3 (1)	0.63	0.88	3 (1)	0.65	0.72	
by locus for each subpopulation	MU59 (1,3)	1 (0)	0.00	0.00	4 (3)	0.56	0.59	
with sample size in parentheses	MU51 (1,3)	2 (0)	0.49	0.63	4 (2)	0.37	0.38	
	G10C (2,1)	1 (0)	0.00	0.00	3 (2)	0.50	0.41	
	MU64 (2,1)	1 (1)	0.00	0.00	3 (3)	0.46	0.33*	
	G1A (2,2)	2 (1)	0.30	0.38	3 (2)	0.51	0.64	
	G10P (2,2)	2 (1)	0.22	0.25	2 (1)	0.45	0.49	
	G10J (3,1)	2 (0)	0.38	0.50	3 (1)	0.65	0.69	
	G10X (3,2)	2 (1)	0.49	0.13*	4 (3)	0.29	0.26	
	MU61 (3,2)	1 (0)	0.00	0.00	2 (1)	0.44	0.49	
	G100 (3,2)	1 (0)	0.00	0.00	1 (0)	0.00	0.00	
	MU05 (4,1)	3 (1)	0.63	0.75	4 (2)	0.58	0.56	
Next to the locus name, between	G1D (4,1)	2 (1)	0.30	0.38	1 (0)	0.00	0.00	
parentheses, the number of the	MU09 (4,2)	2 (1)	0.50	0.75	4 (3)	0.66	0.51*	
preamplification (i) and amplification (i) PCR	G10B (4,2)	1 (0)	0.00	0.00	3 (2)	0.17	0.13	
*Develop diverte D < 0.05	Mean	1.67	0.25	0.28	3.33	0.45	0.44	

\**P*-value significant P < 0.05



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256 (Miller et al. 2002). If we found a multilocus genotype 257 with reliability lower than 95%, more repetitions were 258 carried out until achieving said level of reliability. All the 259 samples that could not be reliably typed for at least 14 out 260 of the 18 loci after the entire process was completed were 261 discarded. To further identify any genotyping errors and 262 the relative magnitude of a problem within our multilocus 263 scores, we performed the tests proposed by McKelvev and 264 Schwartz (2004): Examining Bimodality (EB) and Differ-265 ence in Capture History (DCH). Finally, we determined 266 genotypic mismatches between all scores.

#### 267 Probability of identity

268 Using the software GIMLET version 1.3.2 (Valière 2002), 269 we computed the probability of identity  $(P_{\rm ID})$ , which is the 270 probability that two individuals drawn at random from a 271 given population share identical genotypes at all typed loci 272 (Paetkau and Strobeck 1994). We also computed the 273 probability of identity among siblings  $(P_{\text{ID-Sib}})$  (Waits et al. 274 2001). This value is the upper limit of the possible ranges 275 for the probability of identity in a population and thus 276 provides the most conservative number of loci required to 277 resolve all bears, including relatives. These calculations 278 were carried out for each subpopulation.

#### 279 Population genetic parameters and structure

280 We ran population genetic analyses using the software 281 programmes GENEPOP version 3.4 (Raymond and Rous-282 set 1995), GENETIX version 4.02 (Belkhir et al. 1996-283 2004) and STRUCTURE version 2.1 (Pritchard et al. 2000). Nuclear genetic diversity was measured as the 284 285 number of alleles per locus (A), the observed heterozy-286 gosity  $(H_0)$ , as well as Nei's unbiased expected 287 heterozygosity (H<sub>e</sub>) (Nei 1978). Deviations from Hardy-288 Weinberg equilibrium were tested using an exact test. 289 Global tests across loci for heterozygote deficiency and 290 heterozygote excess and pairwise tests for linkage disequilibrium were performed using Fisher's method (Sokal 291 292 and Rohlf 1994) with 10,000 batches and 10,000 iterations 293 per batch. Correspondence among individual genotypes 294 was studied by means of Factorial Correspondence Anal-295 ysis (FCA), performed with the GENETIX software. 296 Population substructure was detected with the programme 297 STRUCTURE, which uses a Markov chain Monte Carlo 298 (MCMC) algorithm to cluster individuals into populations 299 on the basis of multilocus genotype data (Pritchard et al. 300 2000). We used different values of K, from one to five. For 301 each K tested, we ran STRUCTURE 20 times for 100,000 302 steps, after a burn-in period of 50,000 steps. The correct 303 value of K was estimated following Evanno et al. (2005). 304 The programme also calculates the fractional membership of each individual in each cluster (Q). Quantification of<br/>variation among subpopulations was performed with<br/>GENEPOP using Wright's F-statistics (Weir 1996).305<br/>306

#### Results

Reliability of non-invasive genotyping	309
for the identification of the Cantabrian bear	310

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A total number of 133 non-invasive samples were analysed 311 312 (92 faeces samples and 41 hair samples). We managed to obtain some amplification for all the hair samples and for 313 67 faeces samples (72.8%), but only those samples suc-314 cessfully amplified for more than 14 markers were included 315 in the analysis (Table 1). Another nine genotypes were 316 obtained from tissue/blood samples. For the genotyping, 317 we initially tested 24 loci previously used in genetic 318 tracking of Pyrenean bears (Taberlet et al. 1997; Bellemain 319 and Taberlet 2004). Six out these 24 loci did not give a 320 scorable product and so were discarded. Finally, we used 321 the remaining 18 loci and the sex marker; all the loci were 322 amplified using multiplex PCRs, three of which were 323 specifically developed for this study (Table 2). The prob-324 ability of identity values were 3.28E09 ( $P_{\rm ID}$ ) and 1.16E04 325  $(P_{\text{ID-Sib}})$  for the western subpopulation and 7.45E05 and 326 8.64E03 for the eastern subpopulation. The  $P_{\rm ID}$  and  $P_{\rm ID-Sib}$ 327 values for each marker and subpopulation are shown in 328 Table 3. 329

Once we had completed three repetitions per sample, we 330 331 found that out of the 67 faeces that gave a product, only 40 (59.7%) worked for at least 14 markers. For hair samples, 332 35 out of 41 (85.4%) gave a complete profile. The error 333 334 rates calculated using the GIMLET programme show that both the number of failed PCRs and percentage of total 335 error are higher for faeces samples than for hair samples 336 (see Fig. 2). These results indicate that the DNA recovered 337 from hair samples has better quality than that recovered 338 339 from faeces samples. Results from the RELIOTYPE programme showed that 76% of the samples reach 95% 340 reliability after three repetitions, 16% needed a fourth 341 repetition to reach this level of reliability and 8% needed 342 more than four repetitions to reach it (one of the samples 343 had to be repeated up to seven times to achieve 95% 344 345 reliability).

The results from the EB test, which examines the dis-346 tribution of the genetic differences between samples, and 347 the DCH test, which determines whether the number of 348 new individuals in the sample increased faster than would 349 350 be expected when additional loci are added (McKelvey and Schwartz 2004), indicated that our dataset had a low rate of 351 genotyping error (unimodal distribution for the EB test and 352 no locus-added new individuals for the DHC test). The 353

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Locus (i,j)	Eastern subpopulation	on $(n = 8)$	Western subpopulatio	n ( $n = 39$ )
	P <sub>ID</sub>	P <sub>ID-sib</sub>	P <sub>ID</sub>	P <sub>ID-sib</sub>
MU10 (1,1)	3.75E-01	5.94E-01	3.73E-01	6.34E-01
G10L (1,1)	1.00E+00	1.00E+00	1.10E-01	4.10E-01
MU50 (1,2)	1.00E+00	1.00E+00	1.60E-01	4.56E-01
MU23 (1,2)	2.12E-01	4.87E-01	1.93E-01	4.71E-01
MU59 (1,3)	1.00E+00	1.00E+00	2.71E-01	5.40E-01
MU51 (1,3)	3.79E-01	5.99E-01	4.32E-01	6.71E-01
G10C (2,1)	1.00E+00	1.00E+00	3.61E-01	5.89E-01
MU64 (2,1)	1.00E+00	1.00E+00	3.54E-01	6.10E-01
G1A (2,2)	5.30E-01	7.30E-01	3.58E-01	5.84E-01
G10P (2,2)	6.34E-01	7.99E-01	4.02E-01	6.24E-01
G10 J (3,1)	4.61E-01	6.78E-01	1.96E-01	4.73E-01
G10X (3,2)	3.79E-01	5.99E-01	5.18E-01	7.33E-01
MU61 (3,2)	1.00E+00	1.00E+00	4.09E-01	6.32E-01
G100 (3,2)	1.00E+00	1.00E+00	1.00E+00	1.00E+00
MU05 (4,1)	2.12E-01	4.87E-01	2.38E-01	5.20E-01
G1D (4,1)	5.30E-01	7.30E-01	1.00E+01	1.00E+01
MU09 (4,2)	3.75E-01	5.94E-01	1.82E-01	4.68E-01
G10B (4,2)	1.00E+00	1.00E+00	7.07E-01	8.44E-01
Accumulated	7.45E-05	8.64E-03	3.28E-09	1.16E-04

Table 3 Probability of identity ( $P_{\rm ID}$ ) and Probability of identity for siblings ( $P_{\rm ID-sib}$ ) by locus for each subpopulation with sample size in parentheses and accumulated values

Next to the locus name, between parentheses, the number of the preamplification (i) and amplification (j) PCR



354 presence of a bimodal structure (Fig. 3) in the distribution 355 of the number of loci at which individuals differed (EB 356 test) for the eastern subpopulation is not necessarily a result of error; in this case, it indicates that one of the individuals 357 358 is highly different from the rest, showing that it could be a 359 migrant. The DHC test could not be carried out on the 360 eastern subpopulation, since all the polymorphic loci tested 361 are included in the genetic tag.

362 The average number of loci at which individuals differed was  $10.62 \pm 1.78$  for the western subpopulation and 363 364  $7.69 \pm 4.2$  for the eastern, although if we remove the 365 individual that appears to be different from the rest, the 366 average is lower (5.68  $\pm$  1.94).

A final number of 31 individuals (16 females, 15 males) 367 368 out of 45 samples were identified in the western subpopulation and 9 (5 females, 4 males) out of 30 samples in the 369 eastern subpopulation. 370

Microsatellite diversity and population substructure 371

The number of alleles for the total population was 67; of 372 373 these, 45 were uniquely sampled from one of the two subpopulations (8 for the eastern subpopulation and 37 for the 374 western one). The null hypothesis of uniform allelic and 375 genotypic frequencies in the two subpopulations was rejec-376 ted at 15 out of 17 polymorphic loci. The number of alleles 377 per locus for the western subpopulation ranged between 1 378 and 6, with an average of  $3.33 \pm 1.28$  and between 1 and 3 379 for the eastern subpopulation with an average of  $1.67 \pm 0.67$ 380 (Table 2). Two loci were monomorphic for the western 381

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Fig. 3 Number of loci at which one individual differed from all the other individuals from the same population based on 18 loci for both populations: (a) western and (b) eastern. Only individuals with completed genotypes were used in this calculation (25 for the western subpopulation and 9 for the eastern one). All the comparisons involving one of the individuals found in the eastern subpopulation (in a circle) differed in a higher number of loci from all the rest of the individuals from the same subpopulation

382 subpopulation (G10O and G1D), but up to eight loci were 383 monomorphic for the eastern subpopulation (G10L, Mu50, Mu59, G10C, Mu64, Mu61, G10O, and G10B). The locus 384 385 G10O was monomorphic for both subpopulations. The 386 average observed and expected heterozygosities for the total 387 population were 0.51 and 0.43, respectively, and were sig-388 nificantly different (P < 0.001). The partition of genetic 389 diversity among subpopulations was 42.96%; the values of 390 diversity found for each subpopulation were 0.45 for the 391 western subpopulation and 0.25 for the eastern one 392 (Table 2).

393 Global tests showed that both subpopulations are in 394 Hardy-Weinberg equilibrium, although four loci (Mu10, 395 G10L, Mu64, Mu09) for the western subpopulation and 396 one locus (G10X) for the eastern subpopulation had a 397 significant deficiency in heterozygotes at the P < 0.05398 level (Table 2). Statistical tests for linkage disequilibrium 399 were computed for all pairs of loci, though none of these 400 were significant.

401The canonical analysis based on factorial correspon-402dence analysis (FCA) divided all the samples in two clearly

differentiated groups (Fig. 4). Each group corresponded 403 404 with the two subpopulations present in the Cantabrian Mountains (western and eastern subpopulations). One 405 individual that was detected in samples collected in the 406 eastern area groups together with the individuals from the 407 western subpopulation. This is the same individual that 408 showed a strange pattern when we checked the distribution 409 of the number of loci at which two individuals differed for 410 the eastern subpopulation (Fig. 3). Furthermore, the pro-411 412 gramme STRUCTURE detected that Cantabrian brown 413 bears are structured in two subpopulations which corre-414 spond with the eastern-western groupings (Fig. 4). The mean Q value (cluster membership) is 0.998 for the wes-415 tern subpopulation and 0.888 for the eastern one. Once 416 more, one individual is detected in the eastern subpopula-417 tion that is more similar to the individuals from the western 418 group. The Q value of this individual is 0.010 for the 419 eastern subpopulation and 0.998 for the western one, 420 indicating that this individual (a male) is a migrant. If we 421 remove this individual, the mean Q for the eastern sub-422 population is 0.998. For all the aforementioned reasons, 423 this individual was removed from the basic statistical cal-424 culations regarding population differentiation. 425

The degree of genetic differentiation between the two subpopulations was considerable. The proportion of variation in genetic frequencies,  $\theta$  ( $F_{st}$ ), accounted for by subdivision was 0.41, while Nei's standard distance between both subpopulations was 0.47. 428

#### Discussion

Quality of the genetic data

In order to maximize the success of the non-invasive 433 sample DNA extracts, we used storage and extraction 434 techniques that have previously shown their effectiveness. 435 We obtained 72.8% amplification success for faeces and 436 437 100% for hairs; these values are in the range of values 438 described in the literature (see Nsubuga et al. 2004; Chu et al. 2006). However, these success values dropped to 439 43.48% for faeces and 85.37% for hairs when we consid-440 ered only those samples that gave us a multilocus profile 441 for at least 14 out the 18 loci used. 442

Taberlet et al. (1999) suggest that non-invasive studies 443 444 should include a pilot study to assess the confidence level of the final result by quantifying the genotyping error rate. 445 The results of studies of this type are unique and cannot be 446 transferred to another species or even to another population 447 with different heterozygosity or sample quality. By com-448 bining several methodological and statistical methods for 449 tracking and reducing error previously used in different 450 non-invasive studies (Frantz et al. 2003; Bellemain and 451

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Fig. 4 (a) Distribution of individuals according to genotype based on factorial correspondence analysis (FCA), and (b) Graphical representation of the STRUCTURE programme. In both cases individuals belonging to the eastern and western populations are indicated in white and black, respectively



452 Taberlet 2004; Smith et al. 2006), we were able to thor-453 oughly assess the level of error in our data set (2.55% for 454 hair samples and 5.21% for faeces samples). These error 455 values are similar to the lowest found in the literature for 456 these types of samples (Bayes et al. 2000; Bellemain et al. 457 2004; Smith et al. 2006). The reliability of the multilocus 458 genotypes after three repeats was quite high (76% of the 459 samples had >95% reliability). A fourth repetition increa-460 ses this percentage up to 92%, which is in concordance with the result previously presented by Bellemain and 461 462 Taberlet (2004) in the brown bear.

The low diversity found in the Cantabrian population, mainly in the eastern subpopulation, together with the fact that the most informative loci are not the same in the two subpopulations, makes it necessary to use a high number of markers if we wish to unambiguously identify individuals for future demographic studies.

Genetic status of the bear population in the CantabrianMountains

471 The genetic diversity of the Cantabrian population is lower 472 than in other bear populations considered to have a good 473 conservation status (Paetkau et al. 1998; Waits et al. 1998) 474 and is comparable with those found in Yellowstone 475  $(H_e = 0.55;$  Paetkau et al. 1998) and Deosai National Park 476 (Pakistan) ( $H_e = 0.55;$  Bellemain et al. 2007). Yellowstone

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and Cantabrian populations have had a very similar history. 477 478 Both populations have gone from being embedded in a very 479 large continuous population to being an isolated remnant, separated from other brown bears for 300 years in the case of 480 the Cantabrian population (Naves et al. 1999) and 100 years 481 for the Yellowstone population (Paetkau et al. 1998). In both 482 483 cases, there is no prospect of renewed connections with other 484 populations. The low heterozygosity values can be explained either by a founder effect, which is not the case for the 485 Cantabrian brown bear, or for a sharp decline in population 486 size. The whole population probably began to lose genetic 487 diversity about 300 years ago, when it began to decline in 488 489 size from a larger population.

The brown bear habitat in the Cantabrian Mountains has 490 decreased considerably from  $\sim 9.000 \text{ km}^2$  at the turn of the 491 twentieth century to  $\sim 5,000 \text{ km}^2$  at present (Naves and 492 Nores 1997), while the population subdivided into two 493 apparently isolated subpopulations, the western and the 494 eastern. Genetic differentiation ( $F_{st} = 0.41$ , Nei's standard 495 D = 0.47) between the two subpopulations was found to 496 497 be very high. These values can be compared with the reported microsatellite-based estimates of differentiation 498 among other bear populations. For example, the degree of 499 genetic differentiation is considerably higher than the 500 501 values reported for pairwise comparisons among subpopulations in Scandinavia, where  $F_{st}$  ranged between 0.01 and 502 503 0.14 and Nei's standard distance ranged between 0.03 and

504 0.38. The genetic distance between the two Cantabrian 505 subpopulations, which are only 30 km apart, are compa-506 rable with the values reported for the most distant areas 507 within the continuous distribution of brown bears in North 508 America, which are several thousands of kilometres apart 509 (Paetkau et al. 1998). This result also shows that habitat 510 discontinuities such as roads and farmland play a larger 511 role in genetic substructuring of population than linear 512 distance, which has also been seen in populations of Cross 513 River gorilla (Bergl and Vigilant 2007).

514 The huge genetic differentiation shown between Canta-515 brian subpopulations may be related to total isolation 516 between them in conjunction with an extremely low popu-517 lation size in the eastern subpopulation. Diversity in this 518 subpopulation is among the lowest found in the literature and 519 can only be compared with the value reported in the isolated 520 population of the Kodiak islands in Alaska ( $H_e = 0.26$ ; 521 Paetkau et al. 1998). The eastern subpopulation, with a 522 diversity of 0.25, was detached from the main group a few 523 generations ago. Assuming that diversity at the time of the 524 split was comparable with the diversity of the western sub-525 population, the relative loss of diversity in the eastern 526 subpopulation (H<sub>subpopulation</sub>/H<sub>initial</sub>) is 0.56. This leads to a fixation index (1  $H_{subpopulation}/H_{initial}$ ) equal to 0.44, quite 527 close to the estimated value of  $F_{st}$ . If we assume a continent– 528 529 island model (given that the relative effect of drift in the 530 western subpopulation is low) with no migration, 531  $F_{st}=1$   $(1 \ 1/2N_e)^t$ , where  $N_e$  is the effective size and t the 532 time in generations. Considering that both subpopulations 533 have been isolated for 10 generations (50-75 years with a 534 generation time for the brown bear of 10-15 years, Allen-535 dorf and Servheen 1986; Craighead et al. 1995), the  $N_e$  for 536 the eastern subpopulation that would explain the obtained  $F_{st}$ 537 would be 4.99 individuals per generation. These figures 538 show that the two subpopulations have probably been totally 539 isolated without any effective migration during the last few 540 generations. Although the data point towards a total lack of 541 genetic flow between the two subpopulations, a migrant 542 male from the western subpopulation into the eastern 543 grouping was identified. This could either be interpreted as 544 the first signal that connectivity between both subpopula-545 tions is starting to occur after this long period of isolation, or 546 as the existence of a certain rate of migration, though not effective gene flow. It will be necessary to check in the future 547 548 whether more migrants are present in the population and 549 whether or not this migration is effective in terms of genetic 550 flux. The main genetic consequence of migration would be a 551 drop in genetic differentiation between the two subpopula-552 tions of the Cantabrian brown bear. If we once more assume 553 a continent-island model, the  $F_{ST}$  at equilibrium between 554 gene flow due to migration and genetic drift due to the small population size is  ${}^{F}ST \approx 1/(4N_{em} + 1)$ , where  $N_e$  is the 555 556 effective size and *m* is the migration rate per generation. This implies that with one effective migrant per generation  $(N_{\rm em} = 1)$ , the equilibrium  $F_{\rm ST}$  is 0.20, and 0.11 should there 558 be two effective migrants per generation. That would be sufficient to prevent the huge effects of genetic drift in the oriental subpopulation, despite its reduced population size. 561

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#### Conclusions and recommendations

The results of this study show that the population of 563 564 Cantabrian brown bears is effectively split into two sub-565 populations with a very high level of differentiation. Applying this unique criteria, it could be thought that they 566 should be treated as separate management units (MU) 567 (Moritz 1994) However, bearing in mind their contiguous 568 distribution range, the reduced population size of both 569 570 subpopulations and the real possibility of migration, these two subpopulations need to be managed as a unique unit. 571 The rate of inbreeding per generation in the eastern sub-572 population is around 10%, a value far exceeding the 573 maximum tolerable rate of 1% given for domestic animals 574 575 (Franklin 1980). Therefore, connectivity of both subpopu-576 lations should be the highest priority if we wish to maintain the diversity afforded by the eastern nucleus which is in 577 risk of immediate extinction. 578

The western subpopulation shows a moderate level of 579 diversity in the lowest range of values found in the species, 580 probably due to a sharp decline in population size that began 581 around 300 years ago, when it was isolated from a larger 582 population. A population size of 50-60 individuals was 583 estimated for this nucleus on the basis of the number of 584 585 females with cubs (Wiegand et al. 1998). It was suggested (Franklin 1980) that the minimum effective size for a pop-586 ulation to be viable in the short-term should be 50. Bearing in 587 mind that the ratio of  $N_e/N$  found in the brown bear from 588 Yellowstone was 0.27 (Miller and Waits 2003) and using this 589 value in our case, the minimum size of population to be 590 viable in the short-term is  $\sim$  186 individuals, and even the 591 western subpopulation is far from this number. 592

593 If the whole population is far from the minimum number 594 to be viable over the short-term, an even more difficult 595 situation can be depicted in the context of long-term protection of adaptative potential where the minimum 596 effective population size that has been put forward is 597 between 500 and 5,000 (Franklin 1980; Frankham and 598 599 Franklin 1998; Lynch and Lande 1998). It will be necessary to monitor the whole population to obtain a more 600 accurate estimate of population size and its trend. Habitat 601 loss and human-caused mortality should be avoided in 602 order to facilitate an increase in population size. 603

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