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1 **Forensic Science International: Genetics**

2 **SHORT COMMUNICATION**

3

4 **Y-chromosomal testing of brown bears (*Ursus arctos*): Validation of a multiplex PCR-**
5 **approach for nine STRs suitable for fecal and hair samples**

6

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22

23 **Abstract**

24 High-resolution Y-chromosomal markers have been applied to humans and other primates to
25 study population genetics, migration, social structures and reproduction. Y-linked markers
26 allow the direct assessment of the genetic structure and gene flow of uniquely male inherited
27 lineages and may also be useful for wildlife conservation and forensics, but have so far been
28 available only for few wild species. Thus, we have developed two multiplex PCR reactions
29 encompassing nine Y-STR markers identified from the brown bear (*Ursus arctos*) and tested
30 them on hair, fecal and tissue samples. The multiplex PCR approach was optimized and
31 analyzed for species specificity, sensitivity and stutter-peak ratios. The nine Y-STRs also
32 showed specific STR-fragments for male black bears and male polar bears, while none of the
33 nine markers produced any PCR products when using DNA from female bears or males from
34 12 other mammals. The multiplex PCR approach in two PCR reactions could be amplified with
35 as low as 0.2 ng template input. Precision was high in DNA templates from hairs, fecal scats
36 and tissues, with standard deviations less than 0.14 and median stutter ratios from 0.04 to 0.63.
37 Among the eight di- and one tetra-nucleotide repeat markers, we detected simple repeat
38 structures in seven of the nine markers with 9 to 25 repeat units. Allelic variation was found for
39 eight of the nine Y-STRs, with 2 to 9 alleles for each marker and a total of 36 alleles among
40 453 male brown bears sampled mainly from Northern Europe. We conclude that the multiplex
41 PCR approach with these nine Y-STRs would provide male bear Y-chromosomal specificity
42 and evidence suited for samples from conservation and wildlife forensics.

43

44 *Keywords:* Y-chromosome, Wildlife forensics, Microsatellite, population genetics,
45 conservation genetics, Non-invasive genetic sampling

46

47 **1. Introduction**

48 The brown bear (*Ursus arctos*) is an established model species in conservation genetics [1] and
49 has been extensively studied using both maternally inherited mitochondrial DNA (mtDNA) and
50 biparentally inherited autosomal STR-markers (e.g. [2-12]). Y-linked markers are important in
51 population genetics since they provide information on the male inherited lineages. Y-STR
52 testing may provide important information for a number of different applications including
53 paternity testing, forensic evidence examination, conservation genetics, **population and**
54 **geographic origin assignment** as well as studies of migration patterns. Bi-allelic loci like Y-
55 SNPs and multi allelic loci like Y-STRs are two broad classes of DNA markers that have been
56 used to examine Y-chromosome diversity. Results from combinations of the lower resolution
57 Y-SNPs are usually classified into haplogroups while combination of alleles from the multi-
58 allelic Y-STR loci are characterized as haplotypes [13]. In humans and primates Y-markers
59 have been used to study e.g. population genetics, migration, social structures, and reproduction
60 (e.g. [14, 15]) and Y-markers identified in domesticated animals have been applied in
61 phylogenetic studies, e.g. in sheep, horses and canids (e.g. [16-20]). **In human forensic science,**
62 **Y-markers have been a helpful tool to investigate the geographical/ethnic ancestry of the**
63 **DNA evidence [21, 22]. Despite the advantage of using a non-recombining marker in this type**
64 **of forensic application, in wildlife forensic science the use of autosomal STR and mtDNA**
65 **markers [6, 23-27] is still more common than the use of Y-markers [28] for determining the**
66 **geographic origin of an unknown sample.**

67 Recently, we identified multiple Y-STR loci from five Y-linked scaffolds in brown, polar and
68 American black bear genomes. **Nine** of these Y-STRs were applied in a phylogenetic and
69 phylogeographic study of brown, polar, and black bear [29]. Previously, we have developed a
70 DNA profiling system based on autosomal STRs that are commonly used for bear conservation
71 and in management as well as in forensic cases involving bears [6, 30]. We have also developed

72 a sensitive and specific multiplex PCR assay for sex identification in non-invasive samples
73 from bears [31]. Our aims for this study was to investigate the gene diversity in each of the nine
74 Y-STRs in a large population mainly representing northern Europe. In addition our goal to
75 establish a validated DNA profiling system for bear Y-chromosomal STRs suitable for typing
76 sample materials used in brown bear conservation genetics and wildlife forensics. Thus, we
77 have here performed tests for gender and species specificity, measurements of sensitivity and
78 precision for all the nine bear Y-STRs in agreement with recommendations from ISFG [32].
79 Also, we investigate tandem repeat structure and allele size variation by DNA sequencing
80 alleles from all loci. Finally, our validation include materials like hairs, and fecal samples (non-
81 invasive samples) to ensure that these challenging materials, which is also the most frequently
82 used sample materials in conservation genetics, may be properly analyzed using a validated
83 protocol.

84

85 **2. Materials and Methods**

86 *2.1 Materials*

87 Fecal and hair samples were obtained during monitoring programs, and tissue samples were from legally
88 shot bears during 2006-2012. Samples and sampling procedures have been previously described
89 elsewhere [6, 8, 10, 11, 30]. Samples were analyzed with a previously developed gender test [10] and
90 those identified as from male brown bears were included in our study. A total of 455 samples of male
91 brown bears were included from Norway (n=189), Sweden (n=96), Finland (n=95), northwest Russia
92 (n=65), Romania (n=5) and Canada (n=5). Sample from a male black bear were collected in Alberta in
93 Canada and two male polar bears were from Kolyma in Russia. A total of five samples from female
94 bears collected from the same materials described above were included to test gender specificity of the
95 Y-STRs. Duplicate samples known to be from same individuals of tissue and hair (n=10) or tissue and

96 scats (n=10) were used to demonstrate that all analyses from different materials provided identical Y-
97 haplotypes.

98

99 *2.2 DNA extraction, PCR and Y-STR analysis*

100 DNA was extracted from hair and tissue using Qiagen DNeasy Tissue kit (Qiagen) and from
101 feces using Invitek Stool kit (Stratec), following the manufactures` instructions. Hair samples
102 were stored in room temperature in paper envelopes, tissue in ethanol and feces in stool
103 collection tubes with DNA stabilizer (Stratec). The yield of DNA in tissue from male bears,
104 female bears and 12 other mammalian species (see chapter 2.3) was quantified using a
105 NanoDrop 2000 (Thermo Scientific). Single PCRs were performed in a 10 µl containing 1x
106 PCR Gold buffer (ABI), 200 µM dNTP (Eurogentec), 1.5 mM MgCl₂ (ABI), 0.5 µM of each
107 primer (Life technologies), 1 U Amplitaq Gold DNA polymerase (ABI), 1x BSA (NEB) and 1
108 µl template DNA. All samples have been typed with success using 1 µl template in autosomal
109 markers (lower threshold for successful typing 0.6 ng).

110 DNA amplification was on an ABI 2720 for 10 min at 95 °C, 35 cycles of 30 s at 94 °C, 30 s
111 at 58 °C, and 1 min at 72 °C, and ended with final extension for 45 min at 72 °C. Multiplex-
112 PCR development involved tests of different combinations of markers, primer concentrations
113 and DNA materials (details not shown). For the final analysis, the 9 Y-STR were split into one
114 pentaplex (A) and one tetraplex (B) (Table 1) in touchdown PCR-approach in 10 µl reaction
115 volumes using the following conditions: 5 µl 2x multiplex PCR master mix (Qiagen Multiplex
116 kit), 0.05 µg/µl BSA (NEB), adjusted primer set concentration (Table 1). PCR conditions on an
117 ABI2720 were 95 °C for 10 min followed by 10 cycles of 94 °C for 30 s, 69 °C (decreasing by
118 1 ° C per cycle) for 30 s and 72 °C for 60 s. This was followed by 20 cycles of 94 °C for 30 s,
119 59 °C for 30 s and 72 °C for 60 s. The final step was conducted for 45 min at 72 °C.

120 PCR products (1 μ l) were mixed with Genescan 500 LIZ (Applied Biosystems) size standard
121 (0.25 μ l) and Hi-Di formamide (9.75 μ l) following capillary electrophoresis on an ABI 3130xl
122 Genetic Analyzer (Applied Biosystems). The POP-7™ Polymer was used as separation matrix
123 and the sample injection time were set to 8 s/2kv. PCR fragments were analyzed in GeneMapper
124 4.1 (Applied Biosystems). Prior to PCR, all samples were verified to contain bear DNA as
125 described in [6]. We used 600 RFU as lower threshold for including results from any of the Y-
126 STRs in the Y-haplotypes produced from multiplex PCRs. To check for possible
127 contamination, negative controls were included for every 7th sample in all measurements in this
128 study.

129

130 *2.3 Testing of PCR specificity, sensitivity and precision*

131 To test for species specificity in the two multiplex PCR reactions (A and B), we used
132 approximately 1 ng template DNA from 12 other mammalian species; elk (*Alces alces*),
133 reindeer (*Rangifer tarandus*), wolverine (*Gulo gulo*), eurasian lynx (*Lynx lynx*), wolf (*Canis*
134 *lupus*), hare (*Lepus timidus*), red deer (*Cervus elaphus atlanticus*), domesticated cat (*Felis*
135 *catus*), badger (*Meles meles*) raccoon dog (*Nyctereutes procyonoides*), dog (*Canis familiaris*)
136 and human. We used 1 ng female template DNA from brown bears (*Ursus arctos*), Canadian
137 black bear (*U. americanus*) and polar bears (*U. maritimus*) to analyze gender specificity.

138 Two positive controls (tissue) of male bear DNA (1 ng template DNA) were included for
139 species specificity and gender specificity. Sensitivity was evaluated by PCR amplification of
140 DNA from male brown bear muscle tissue in the range 20–0.1 ng. Measurements of within-run
141 precision and stutter ratios were performed in 10 independent amplifications and subsequent
142 runs of a single sample of feces, hair and tissue, respectively.

143

144 *2.4 DNA Sequencing*

145 The tandem repeat array and the immediate upstream and downstream sequences at each of the
146 nine loci were analyzed by DNA sequencing. PCR products amplified from DNA from male
147 brown bears were sequenced using the BigDye Terminator v3.1 Cycle Sequencing Kit (ABI)
148 as recommended by the manufacturer. Single PCR-primers from Table 1 were used as
149 sequencing primers in forward and reverse sequencing reactions, respectively. Forward and
150 reverse sequences from each sample were aligned in Sequencher 4.7. The allelic sequences
151 from each locus were aligned and the sequence and size variation at each locus was determined
152 by manual inspection.

153

154 **3. Results**

155 *3.1 Tests of species and gender specificity and multiplex PCR combinations.*

156

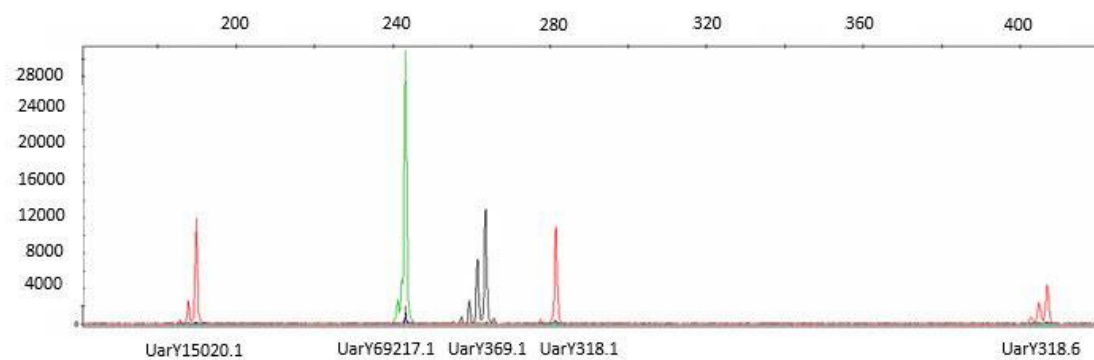
157 Template DNA from one black bear and two polar bears was tested in the different multiplex
158 combinations of the nine Y-STRs using the primers described in methods. **We found that all
159 nine markers were amplified for both species. The allele-sizes observed in one black bear and
160 two polar bears were within the allele-size range found in our brown bear populations,
161 indicating that alleles from these markers are not grouped into discrete allele sizes among
162 different bear species. This would also be consistent with findings in Bidon et al [29].**

163 The specificity of the primers to bear species was confirmed using template DNA from **males
164 of 12 other mammalian species** (see chapter 2.3). Our result showed that all species gave a
165 negative result in multiplex reactions tested.

166 Similarly, we could not detect any PCR products in multiplex reactions tested when we used
167 DNA from five female bears as template.

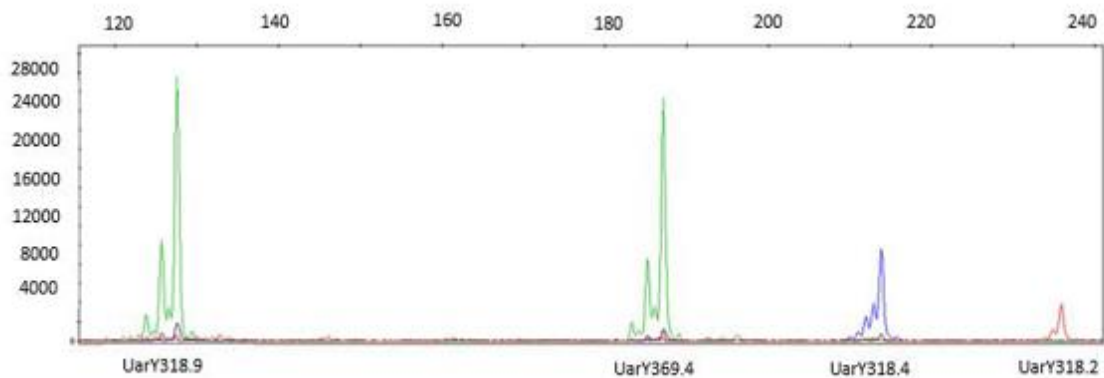
168 Different combinations of markers, primer concentration and DNA- material from male brown
169 bears was tested in multiplex PCR reactions (data not shown). The highest signal to noise ratio
170 was achieved when the nine Y-STR were split into one pentaplex (A) and one tetraplex (B)
171 PCR reaction. Figure 1 demonstrate results obtained using these two multiplex PCRs. The
172 combination of markers and primer concentrations used in multiplex reaction A and B are given
173 in Table 1.

174 **Multiplex A**



175

176 **Multiplex B**



177

178 **Figure 1. Chromatograms from capillary electrophoresis (ABI 3130XL) showing multiplex PCR reactions**
179 **A and B (see also Table 1) for the nine brown bear Y-chromosomal STRs. Template DNA (1 ng) were from**
180 **brown bear tissues. The names of the STR-markers are indicated below the peaks, and the peak height**
181 **(RFU) is indicated on the Y-axis.**

182

Table 1: PCR primers, alleles and multiplex PCR set-up for nine Y-STRs from brown bears

Locus	Primer sequences (5'-3') ^a	Repeat motif	Allele size range (bp) ^b	No. of alleles observed ^c	PCR multiplex	Primer conc., Dye
UarY318.4	F:TACCTGGCTGGCTTTCTTGG R:CACTGTTGGTTTTGGCTCCG	GA	213-215	2	B	1 μM, FAM
UarY318.2	F:CAGGCTGACACTGGGGATTT R:AAGAGGGAGTCATCTGGGGT	TA	233-235	2	B	3 μM, PET
UarY318.9	F:CACTCAGGCACCCCTCTATC R:TGGCCAGGATACAGAAACAAC	AC	127-131	3	B	1 μM, VIC
UarY369.4	F:AGGCATCCATTCTATCACCAC R:TGTGGATGTATCTGCCCAAC	AC	182-200	7	B	1 μM, VIC
UarY318.1	F:GGGATCAAGCCCCACATCAA R:ACTTGTAGATGCACATCTGTGGT	AAAT	281-289	3	A	2 μM, PET
UarY69217.1	F:CTCCACCTTGCTGCCACTC R:TTCCCTCCCTTTCTGTCCT	TG	243	1	A	1 μM, VIC
UarY318.6	F:GCTGGCTGTCTCTCTCTGA R:AAATTCCTTTGGAAACGTCCT	TG	400-410	6	A	3 μM, PET
UarY15020.1	F:TGCAATTTCTCTCAAACAACCTTCCT R:GCGATGAAGGTCAGAGCAGT	TG	185-189	3	A	1 μM, PET
UarY369.1	F:TCCCTGAATGAGCAGTAGCC R:GGGGTATTGCGTTGCATTGG	TG	249-273	9	A	2 μM, NED

183

^aF forward, R reverse, from Bidon et al. 2014

184

^b Allele Size range in base pairs observed in 455 male brown bears (see Table 4)

185

^c A total of 36 different alleles were observed among 455 male brown bears (see Table 4).

186

187 3.2 Sensitivity, precision and stutter ratios

188 We tested a concentration series of 20, 10, 1, 0.5, 0.2, 0.1, 0.05, 0.04, 0.03, 0.02 and 0.01 ng of
 189 template DNA in the two multiplex reactions. All markers in multiplex A were successfully
 190 typed with signals above the lower peak height threshold of 600 RFU with template DNA in
 191 the range 20–0.1 ng while multiplex B showed successful typing of all markers with template
 192 DNA in the range 20-0.2 ng. All samples representing duplicates from same individual, but
 193 template extracted from different materials, showed identical results for all parallel samples.

194 We also tested the within-run precision and stutter ratios using ten independent amplifications
 195 and subsequent runs of one sample of feces, hair and tissue (Table 2). These results show that
 196 the standard deviations (S.D.) from allele length measurements of all the nine loci tested were

197 between 0.05 bp and 0.14 bp. Within-run precision and stutter ratios showed very similar results
 198 for hair-, fecal- and tissue-samples (Table 2). Stutters at -1R was observed in all markers (Figure
 199 1). We found that these stutter ratios ranged from 0.04 to 0.63 from all materials (Table 2).
 200 Stutter in position -2R at loci UarY369.1, UarY318.4, UarY318, UarY369.4 and UarY318.9
 201 was observed, but at much lower proportions (<0.04) than stutter band at -1R. Stutter one repeat
 202 larger than the true allele (+1R) was also observed, but in very small proportion (<0.05).

203

204 **Table 2: Measurements of precision and stutter ratio for Y-chromosomal STRs from brown bears.**

Locus ^a		Allele/genotype ^a	Mean (bp) ^b	Stutter ratio ^c
UarY318.4	Feces	213	213.76 (0.06)	0.21 (0.23)
	Hair	213	213.71 (0.05)	0.23 (0.24)
	Tissue	213	213.72 (0.07)	0.24 (0.31)
UarY318.2	Feces	235	235.78 (0.14)	0.10 (0.11)
	Hair	235	235.66 (0.14)	0.10 (0.60)
	Tissue	235	235.76 (0.09)	0.10 (0.10)
UarY318.9	Feces	127	127.59 (0.14)	0.33 (0.35)
	Hair	125	125.66 (0.10)	0.24 (0.26)
	Tissue	127	127.51 (0.11)	0.33 (0.35)
UarY369.4	Feces	192	192.82 (0.13)	0.40 (0.43)
	Hair	184	185.12 (0.13)	0.20 (0.22)
	Tissue	186	187.04 (0.13)	0.29 (0.36)
UarY318.1	Feces	281	281.51 (0.10)	0
	Hair	285	285.38 (0.09)	0.04 (0.05)
	Tissue	285	285.40 (0.07)	0.04 (0.05)
UarY69217.1	Feces	243	243.13 (0.08)	0.08 (0.11)
	Hair	243	243.08 (0.05)	0.12 (0.15)
	Tissue	243	242.88 (0.07)	0.18 (0.23)
UarY318.6	Feces	406	406.87 (0.13)	0.50 (0.58)
	Hair	404	404.74 (0.08)	0.49 (0.52)
	Tissue	404	404.71 (0.07)	0.49 (0.50)
UarY15020.1	Feces	189	189.56 (0.11)	0.23 (0.28)
	Hair	187	187.52 (0.08)	0.20 (0.20)
	Tissue	187	187.61 (0.11)	0.21 (0.29)
UarY369.1	Feces	263	263.58 (0.08)	0.55 (0.61)
	Hair	267	267.67 (0.05)	0.63 (0.64)
	Tissue	267	267.69 (0.04)	0.62 (0.63)

205 Stutter ratio was calculated by dividing the peak height (RFU) of the stutter peak in position -1R (one repeat less than the true allele) by the
 206 peak height of the true allele

207 ^a Genotype nomenclature is based on PCR fragment sizes.

208 ^b Mean value allele sizes when measured with POP7 on ABI3730, with SD from in within-run measurement of 10 run per sample.

209 ^c Median stutter ratio with upper 95% percentile in parenthesis.

210

211 *3.3 Investigation of repeat structure and gene variation for the nine Y-STRs*

212
 213 DNA sequencing was performed on the largest and the smallest alleles observed for each
 214 marker. The sequencing revealed that all size variation observed between the two alleles
 215 selected from same loci could be explained by variation in repeat numbers in the tandem repeat
 216 arrays (Table 3). Eight loci were tandem arrays of dinucleotide repeats while one had a repeat
 217 array of tetranucleotide repeats (UarY318.1). Seven of the loci showed simple tandem repeat
 218 array structures while loci UarY318.2 and UarY69217.1 showed compound repeat structures
 219 (Table 3).

220

221 **Table 3. DNA sequencing of tandem repeat structures of nine Y-chromosomal STRs from brown bears**

Locus	Allele ^a	No. repeats	Repeat structure
UarY318.4	213	12R	(GA) ₁₂
	215	13R	(GA) ₁₃
UarY15020.1	187	11R	(GT) ¹¹
	189	12R	(GT) ₁₂
UarY318.1	281	9R	(AAAT) ⁹
	289	11R	(AAAT) ₁₁
UarY318.2	235	18R	(TA) ⁴ (TG)(TA) ² (TG)(TA)(T)(TA) ⁷ (A)(TA) ⁴
Uar318.9	127	14R	AC ¹⁴
	131	16R	AC ¹⁶
UarY369.1	259	18R	GT ¹⁸
	273	25R	GT ²⁵
UarY369.4	186	15R	(AC) ₁₅
	200	22R	(AC) ₂₂
UarY69217.1^b	243	11R	(TG) ⁶ (TA)(TG) ⁵
UarY318.6	400	15R	(TG) ²⁰
	410	22R	(TG) ²⁵

222 ^a Nomenclature of alleles is based on PCR fragment size.

223 ^b Monomorphic in this study.

224

225 Allele size measurements showed that there was allelic variation in eight of the nine Y-STRs
226 analyzed in our material. The number of alleles observed in each marker was from 2-9. We
227 found 36 different alleles among the 453 males analyzed. Allele frequencies for each locus are
228 given for each country in Table 4. Rare alleles represented by only one individual were alleles
229 249, 273 (UarY369.1) and 410 (UarY318.6). In addition, 10 other alleles were present at low
230 frequencies (>0.05). Locus UarY69217.1 did not reveal any size variation (monomorphic).
231 Locus UarY318.2 showed allele size variations in individuals from Canada and Romania (allele
232 233) while all bears from Northern Europe were identical (allele 235). A total of 45 different
233 Y-haplotypes were revealed in our material (Table 5). Some Y-haplotypes are fairly common
234 among brown bears included in our material (mostly representing northern Europe) with
235 frequencies ranging from 0.2 to 98.2 % in polymorphic markers. However, 19 different Y-
236 haplotypes were observed in single individuals only.

237

238 **Table 4: Allele frequencies for nine Y-STRs for brown bears marker per country. Frequencies are first**
 239 **given per country of origin of samples and then for the total sample. The number of individuals analyzed**
 240 **for each country and in total, as well as the number of individuals carrying the respective allele are given in**
 241 **brackets.**

Marker	Allele	Allele frequency						
		Finland (95)	Norway (189)	Russia (65)	Sweden (96)	Romania (5)	Canada (5)	Total (455)
UarY318.4	213	0.842 (80)	0.899 (170)	0.846 (55)	0.846 (96)	1.000 (5)	1.000 (5)	0.903 (411)
	215	0.161 (15)	0.101 (19)	0.154 (10)	-	-	-	0.097 (44)
UarY369.1	249	-	-	-	-	0.200 (1)	-	0.002 (1)
	259	0.022 (2)	-	0.031 (2)	-	0.200 (1)	-	0.011 (5)
	261	0.043 (4)	0.333 (63)	0.015 (1)	0.427 (41)	-	-	0.241 (109)
	263	0.043 (4)	0.106 (20)	0.138 (9)	0.031 (3)	-	-	0.079 (36)
	265	0.527 (49)	0.312 (59)	0.385 (25)	0.313 (30)	0.200 (1)	-	0.362 (164)
	267	0.232 (22)	0.196 (37)	0.215 (14)	0.229 (22)	0.200 (1)	-	0.211 (96)
	269	0.054 (5)	0.005 (1)	0.092 (6)	-	0.200 (1)	1.000 (5)	0.040 (18)
	271	0.086 (8)	0.048 (9)	0.123 (8)	-	-	-	0.055 (25)
	273	0.011 (1)	-	-	-	-	-	0.002 (1)
UarY15020.1	185	-	-	-	-	-	1.000 (5)	0.011 (5)
	187	0.568 (54)	0.376 (71)	0.585 (38)	0.585 (39)	-	-	0.407 (185)
	189	0.441 (41)	0.624 (118)	0.415 (27)	0.771 (74)	1.000 (5)	-	0.585 (265)
UarY318.2	233	-	-	-	-	0.600 (3)	1.000 (5)	0.018 (8)
	235	1.000 (95)	1.000 (189)	1.000 (65)	1.000 (96)	0.400 (2)	-	0.982 (447)
UarY318.1	281	0.484 (45)	0.619 (117)	0.508 (33)	0.771 (74)	0.800 (4)	1.000 (5)	0.614 (278)
	285	0.505 (48)	0.381 (72)	0.492 (32)	0.229 (22)	0.200 (1)	-	0.385 (175)
	289	0.022 (2)	-	-	-	-	-	0.004 (2)
UarY369.4	186	0.337 (32)	0.265 (50)	0.308 (20)	0.229 (22)	-	-	0.273 (124)
	190	0.108 (10)	0.159 (30)	0.062 (4)	0.313 (30)	-	-	0.163 (74)
	192	0.269 (25)	0.545 (103)	0.338 (22)	0.458 (44)	0.200 (1)	1.000 (5)	0.442 (200)
	194	0.032 (3)	-	0.031 (2)	-	-	-	0.011 (5)
	196	0.204 (19)	0.032 (6)	0.154 (10)	-	0.200 (1)	-	0.079 (36)
	198	0.032 (3)	-	0.077 (5)	-	0.600 (3)	-	0.024 (11)
	200	0.032 (3)	-	0.031 (2)	-	-	-	0.011 (5)
UarY69217.1	243	1.000 (95)	1.000 (189)	1.000 (65)	1.000 (96)	1.000 (5)	1.000 (5)	1.000 (455)
UarY318.6	400	0.172 (16)	0.111 (21)	0.215 (14)	-	-	-	0.113 (51)
	402	0.258 (24)	0.032 (6)	0.246 (16)	-	-	-	0.102 (46)
	404	0.284 (27)	0.233 (44)	0.323 (21)	0.229 (22)	-	-	0.251 (114)
	406	0.280 (26)	0.608 (115)	0.169 (11)	0.771 (74)	1.000 (5)	1.000 (5)	0.521 (236)
	408	0.011 (1)	0.016 (3)	0.046 (3)	-	-	-	0.015 (7)
	410	0.011 (1)	-	-	-	-	-	0.002 (1)
UarY318.9	127	0.684 (65)	0.413 (78)	0.646 (42)	0.260 (25)	0.800 (4)	1.000 (5)	0.481 (219)
	129	0.062 (6)	0.005 (1)	0.138 (9)	-	0.200 (1)	-	0.038 (17)
	131	0.258 (24)	0.582 (110)	0.215 (14)	0.740 (71)	-	-	0.483 (219)

242 **Table 5: Frequencies of brown bear Y-chromosome haplotypes* per country. Frequencies are first given**
 243 **per country of origin of samples and then for the total sample. The number of individuals analyzed for each**
 244 **country and in total, as well as the number of individuals carrying the respective haplotype are given in**
 245 **brackets.**

Haplotype	Haplotype frequency						Total (455)
	Finland (95)	Norway (189)	Russia (65)	Sweden (96)	Romania (5)	Canada (5)	
1.01	0.137 (13)	0.101 (19)	0.092 (6)	-	-	-	0.084 (38)
1.13	0.011 (1)	-	-	-	-	-	0.002 (1)
1.19	-	-	0.015 (1)	-	-	-	0.002 (1)
1.23	0.011 (1)	-	0.046 (3)	-	-	-	0.009 (4)
1.25	-	-	0.031 (2)	-	-	-	0.004 (2)
1.37	0.011 (1)	-	-	-	-	-	0.002 (1)
2.02	0.021 (2)	0.101 (19)	0.077 (5)	0.031 (3)	-	-	0.064 (29)
2.05	0.063 (6)	0.148 (28)	0.031 (2)	0.313 (30)	-	-	0.147 (66)
2.06	0.021 (2)	0.005 (1)	0.046 (3)	-	-	-	0.013 (6)
2.07	0.189 (18)	0.026 (5)	0.123 (8)	-	-	-	0.069 (31)
2.08	0.021 (2)	0.333 (63)	-	0.427 (41)	-	-	0.236 (106)
2.10	-	-	0.031 (2)	-	-	-	0.004 (2)
2.11	0.021 (2)	-	0.015 (1)	-	-	-	0.007 (3)
2.12	0.021 (2)	-	0.031 (2)	-	-	-	0.009 (4)
2.14	0.032 (3)	-	0.031 (2)	-	-	-	0.011 (5)
2.16	-	-	0.015 (1)	-	-	-	0.002 (1)
2.17	-	-	0.031 (2)	-	-	-	0.004 (2)
2.18	-	-	0.015 (1)	-	-	-	0.002 (1)
2.22	0.021 (2)	-	-	-	-	-	0.004 (2)
2.24	-	0.005 (1)	-	-	-	-	0.002 (1)
2.27	0.011 (1)	-	-	-	-	-	0.002 (1)
2.28	-	0.005 (1)	-	-	-	-	0.002 (1)
2.29	-	-	0.015 (1)	-	-	-	0.002 (1)
2.30	-	-	0.015 (1)	-	-	-	0.002 (1)
2.31	0.032 (3)	-	-	-	-	-	0.007 (3)
2.32	0.011 (1)	-	-	-	-	-	0.002 (1)
3.03	0.074 (7)	0.016 (3)	0.015 (1)	-	-	-	0.024 (11)
3.04	0.021 (2)	0.048 (9)	0.077 (5)	-	-	-	0.036 (16)
3.09	0.179 (17)	0.185 (35)	0.123 (8)	0.229 (22)	-	-	0.182 (82)
3.15	-	-	0.031 (2)	-	-	-	0.004 (2)
3.20	0.011 (1)	0.011 (2)	0.031 (2)	-	-	-	0.011 (5)
3.21	0.011 (1)	0.016 (3)	0.031 (2)	-	-	-	0.013 (6)
3.26	-	-	0.015 (1)	-	-	-	0.002 (1)
3.33	-	-	0.015 (1)	-	-	-	0.002 (1)
3.34	0.011 (1)	-	-	-	-	-	0.002 (1)
3.35	0.021 (2)	-	-	-	-	-	0.004 (2)
3.36	0.032 (3)	-	-	-	-	-	0.007(3)
3.38	0.011 (1)	-	-	-	-	-	0.002 (1)
x.39	-	-	-	-	0.200 (1)	-	0.002 (1)
x.40	-	-	-	-	0.400 (2)	-	0.004 (2)

x.41	-	-	-	-	0.200 (1)	-	0.002 (1)
x.42	-	-	-	-	0.200 (1)	-	0.002 (1)
x.43	-	-	-	-	0.200 (1)	-	0.002 (1)
x.44	-	-	-	-	-	1.000 (5)	0.011 (5)

* Haplotypes are based on variation in 8 Y-STRs (see table 4). For haplogroups 1, 2 and 3, see [33],
x= haplogroup not determined.

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249 **4. Discussion**

250 In this study, we have validated nine Y-chromosomal STRs for use in conservation genetics
251 and wildlife forensic. We find that a dual multiplex PCR approach represents a fast and precise
252 assay for Y-chromosomal DNA profiling and haplotyping of hair- , fecal- and tissue samples
253 from brown bears. All nine Y-STRs used for this work had originally been identified and
254 aligned in genomic sequence data from polar and brown bears, and also tested successfully on
255 American black bears [29]. We recently showed that the patterns of PCR fragment sizes (allele
256 sizes) differ for the three bear species on a phylogeographic scale [29]. Additionally, we here
257 report a large and geographically extensive set of population data for the brown bear, while
258 detailed intra-population assessments of Y-chromosomal variation are still lacking for polar
259 bears and American black bears.

260 To test for bear specificity we used test species that bears may predate upon, as well as hair and
261 fecal samples from other carnivores that may erroneously be collected as bear samples in the
262 field. We also included human DNA to assure that any contamination by handling samples
263 would not produce any false results. Even though DNA from more species may have been
264 included in the test, we may conclude that all these nine Y-markers show very strong male bear-
265 specific amplification. The negative results from female bear DNA (absence of amplification
266 signals) from all three species show that all nine Y-STRs are male-specific sequences that do
267 not have any close homologues on the X-chromosome or autosomes. This finding is also

268 supported by our DNA sequencing results on the alleles, i.e no indication of heterozygosity that
269 may be a result of a second copy interfering with the assay.

270 Sensitivity testing showed that the Y-specific multiplex PCR method worked well down to
271 amounts of 0.2 ng template DNA. This is more sensitive than for autosomal STRs on similar
272 samples that are successfully amplified down to 0.6 ng, but less sensitive than our previous
273 results on multiplex PCR assays for sex determination at DNA template levels as low as 0.02
274 ng [31]. The latter is based on very small amplicon sizes (100-160bp) from Y- and X-
275 chromosomes, while the PCR fragments for this study have a wider range (127-410 bp). Thus,
276 reducing the larger Y-STR amplicon sizes and then repeating the multiplex development may
277 further improve the sensitivity of our novel assay. However, this may also be unfavorable
278 because of Y-sequence structures and motifs that are not ideal for PCR priming (results not
279 shown).

280 Precision was in general high for both multiplex assays A and B, also when compared to
281 autosomal STRs (see [6]). Stutter ratios were as expected very low for the single tetranucleotide
282 repeat (UarY318.1), and more pronounced for the remaining eight dinucleotide repeats. Stutter
283 ratios were in general much lower than measured previously for brown bear autosomal STRs
284 [6]. We tested template DNA extracted from hairs, scats and tissues in the multiplex assays. All
285 replicated sample materials from the same individuals showed identical Y-haplotypes and very
286 low variation in precision and stutter ratios were observed. This justifies that larger monitoring
287 or conservation studies may combine Y-haplotypes generated from the three sample materials
288 tested in our validated multiplex assays.

289 Sequencing of alleles showed that the size variation observed could be explained as depending
290 on the number of repeats. A nomenclature of alleles based on the number of repeats instead of
291 PCR fragment size used in human forensics is also recommended for STR markers applied for

292 wildlife forensics [32]. Sequencing of all alleles observed could facilitate such a nomenclature
293 on alleles in the Y-STRs validated in our study. Independent of this, we offer sample DNA of
294 alleles sequenced in our study to other laboratories for use as inter-laboratory calibrators.

295 Allele frequencies for 36 different alleles showed only three different alleles that were present
296 in only one male, while 10 additional alleles were found to be rare in Northern Europe.
297 Including ten samples from other regions (Canada and Romania) only added one extra allele,
298 but showed that the corresponding marker (UarY318.2) also is polymorphic in brown bears.
299 Thus even if marker UarY69217.1 was monomorphic in Northern Europe, we still suggest that
300 this marker be kept in the multiplex, for application of our method in other bear populations
301 and species.

302 In wildlife forensics and illegal trade with endangered species, determination of population and
303 geographical origin of an unknown forensic sample can contribute to resolve cases [23, 34, 35],
304 but this requires the source population to be sufficiently genetically distinct from other
305 candidate populations, as well as large reference data [26]. Previous studies [29] and our present
306 results show region-specific Y-haplotype frequencies across Europe as well as some region-
307 specific haplotypes [33]. This indicates that the construction of a Y-STR-based profiling
308 system, especially in combination with one based on autosomal STRs [6, 30], may contribute
309 significantly to resolving geographical origin in brown bear forensic cases in the future.

310 Resolving wildlife forensic cases like poaching, illegal killing, collection and trade is most
311 commonly achieved using autosomal STR-profiling and mtDNA-sequencing [27]. The use of
312 Y-STR markers provides additional information and is already commonly used in human
313 forensics cases to resolve male-female and male-male mixtures, e.g., to reliably determine the
314 minimum number of male individuals presumable involved [22]. Our Y-STR marker developed
315 for brown bear could provide same useful data in wildlife forensic cases as well as in predation

316 events [28]. Another argument for the promise of Y-markers in bears is the strong male-biased
317 dispersal, which means that males are at larger risk of being victims of poaching, or being
318 involved in conflict with humans.

319

320 **5. Conclusion**

321 Our work is focused on validation of the Y-profile system for brown bears. In our opinion, such
322 a validation is a pre-requisite for its use in forensics, and such a standard is also desirable in
323 conservation genetics. To use this Y-profile system for geographical assignment there need to
324 be developed an even larger population database eventually containing bears from all over the
325 world. If the research communities use the Y-profile system presented here, such data may be
326 compared and thus our Y-system represents a means to achieve this rather than a proof that our
327 Y-system may be used in such a manner.

328

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