1	Highly polymorphic microsatellite markers for the assessment of male reproductive skew										
2	and genetic variation in Critically Endangered crested macaques (Macaca nigra)										
3	Antje Engelhardt ^{1,2*} , Laura Muniz ^{3,4} , Dyah Perwitasari-Farajallah ^{5,6} , Anja Widdig ^{3,4,7*}										
4											
5											
6	¹ Junior Research Group of Primate Sexual Selection, German Primate Center, Germany										
7	² Courant Research Center Evolution of Social Behavior, Georg-August-University, Germany										
8	³ Junior Research Group of Primate Kin Selection, Department of Primatology, Max-Planck										
9	Institute for Evolutionary Anthropology, Germany										
10	⁴ Research Group of Behavioural Ecology, Institute of Biology, University of Leipzig, Germany										
11	⁵ Primate Research Centre, Bogor Agricultural University, Indonesia										
12	⁶ Department of Biology, Faculty of Mathematics and Natural Sciences, Bogor Agricultural										
13	University, Indonesia										
14	⁷ German Center for Integrative Biodiversity Research, Germany										
15											
16	*Corresponding authors:										
17	-Antje Engelhardt, School of Natural Sciences and Psychology, Liverpool John Moores										
18	University, Byrom Street, Liverpool L3 3AF, United Kingdom, email:										
19	A.Engelhardt@ljmu.ac.uk; phone: +44 151231 2434										
20	and										
21	-Anja Widdig, Junior Research Group of Primate Kin Selection, Department of Primatology,										

- 22 Max-Planck Institute for Evolutionary Anthropology, Deutscher Platz 6, D-04103 Leipzig,
- Germany, email: <u>anja.widdig@eva.mpg.de</u>; phone: +49 341 9736 707

24 Abstract

Genetic analyses based on non-invasively collected samples have become an important tool for 25 evolutionary biology and conservation. Crested macaques (Macaca nigra), endemic to 26 27 Sulawesi, Indonesia, are important for our understanding of primate evolution as Sulawesi macaques represent an exceptional example of primate adaptive radiation. Crested macaques 28 are also Critically Endangered. However, to date we know very little about their genetics. The 29 30 aim of our study was to find and validate microsatellite markers useful for evolutionary, conservation and other genetic studies on wild crested macaques. Using faecal samples of 176 31 32 wild macaques living in the Tangkoko Reserve, Sulawesi, we identified 12 polymorphic 33 microsatellite loci through cross-species PCR amplification with later modification of some of these primers. We tested their suitability by investigating and exploring patterns of paternity, 34 observed heterozygosity and evidence for inbreeding. We assigned paternity to 63 of 65 infants 35 with high confidence. Among cases with solved paternity, we found no evidence of extra-group 36 paternity and natal breeding. We found a relatively steep male reproductive skew B index of 37 38 0.330±0.267; mean±SD) and mean alpha paternity of 65% per year with large variation across groups and years (29-100%). Finally, we detected an excess in observed heterozygosity and no 39 evidence of inbreeding across our three study groups, with an observed heterozygosity of 40 0.766±0.059 and expected heterozygosity of 0.708±0.059, and an inbreeding coefficient of -41 0.082±0.035. Our results indicate that the selected markers are useful for genetic studies on 42 wild crested macaques, and possible also other Sulawesi and closely related macaques. They 43 further suggest that the Tangkoko population of crested macaques is still genetically variable 44 despite its small size, isolation and the species' reproductive patterns. This gives us hope that 45 46 other endangered primate species living in small, isolated populations may also retain a healthy gene pool, at least in the short term. 47

48

49 Keywords: microsatellite markers, *Macaca nigra*, Sulawesi, conservation, paternity,
50 reproductive skew, genetic variation, inbreeding, heterozygosity

51 Conflict of Interest: The authors declare that they have no conflict of interest

52

53 Introduction

54 The development of genetic analyses has revolutionized various fields in the medical and life sciences. More recently, genetic analyses based on naturally dropped animal waste such as fur, 55 feathers and faeces have created new opportunities for studies of wildlife under natural 56 conditions, particularly endangered and/or elusive species, and other species in which capturing 57 constitutes an ethical problem (e.g. Waits & Paetkau 2005). Potential applications of genetic 58 analyses for field studies include examining the occurrence, distribution and history of species 59 (e.g. Hewitt 2000; Leonard 2008; Ram et al. 2015), investigating taxonomic relationships and 60 speciation (e.g. Tosi et al. 2004), assessing hybridization (e.g. Roos et al. 2011; Charpentier et 61 62 al. 2012; Godinho et al. 2015), determining the level of heterozygosity, gene flow and the risk of inbreeding depression of isolated populations (Luikart et al. 1998; Nürnberg et al. 1998; 63 Widdig et al. 2004; Knief et al. 2015; Ram et al. 2015; Widdig et al. 2017), monitoring 64 65 population developments and movements (e.g. Nowak et al. 2014), identifying species (Harms et al. 2015), and studying reproductive patterns (Widdig et al. 2004; Engelhardt et al. 2006; 66 Syrůčková et al. 2015) and kin relationships in groups and populations (e.g. van Horn et al. 67 2008; Montague et al. 2014). Hence, studies of evolutionary biology, biogeography and 68 behavioural ecology greatly benefit from the availability of genetic analyses based on non-69 70 invasively collected samples, as does conservation management (Schwartz et al. 2007). The genetic markers used in such studies often need to be specified for the species in question, 71 although the same markers can be used for closely related species. 72

Genetic markers are not yet available but would be very important for the Sulawesi macaques. 73 74 The seven species of macaques on the island of Sulawesi (Macaca brunnescens, M. hecki, M. maurus, M. nigra, M. nigrescens, M. ochreata, M. tonkeana), the main island of the Wallacea 75 biodiversity hotspot, are an important group for our understanding of primate evolution. 76 Endemic to the island, they are a prominent example of primate adaptive radiation and 77 speciation in relation to the processes of geological change and colonization of new areas 78 79 (Groves et al. 1980). All seven species live in different habitats with only narrow overlapping contact zones, in which interbreeding occurs (Fooden 1982; Evans et al. 2003). Furthermore, 80 Sulawesi macaques are the only macaques classified as extremely socially tolerant with high 81 82 conciliatory tendencies and low degrees of power asymmetries (Thierry et al. 2000; Thierry 2004). Few studies have investigated Sulawesi macaques in the wild because their habitat is 83 very difficult to access. However, the rainforests of Sulawesi are now more accessible, and the 84 85 infrastructure on Sulawesi has improved, facilitating studies of Sulawesi wildlife. However, with these developments, the natural habitat of the macaques is shrinking and fragmented, and 86 heavily exploited by humans. As a result, all seven Sulawesi macaques are in danger of 87 extinction to various degrees (IUCN 2016). Given the precarious situation and geographic 88 isolation of Sulawesi macaques, genetic studies on these species are important not only for our 89 90 understanding of primate evolution (Evans et al. 1999, 2003), but also for their conservation management. 91

Crested macaques, *M. nigra*, are only found on the northern tip of Sulawesi. Habitat degradation and bushmeat hunting have brought this species to the edge of extinction, with the largest remaining population of less than 2000 animals seemingly occurring in Tangkoko Reserve (Palacios et al. 2012; Melfi 2010). There are at least two reasons why we need genetic studies of crested macaques. First, crested macaques are of particular interest for better understanding primate evolution since the species possesses features not found in any of the other Sulawesi macaques. For example, other Sulawesi macaques live in groups of up to 40 animals, while

crested macaques live in large groups sometimes containing over 100 individuals (Riley 2010; 99 100 Marty et al. 2015). Despite the large group size, crested macaques seem to be an extreme case in terms of male-male reproductive competition with males fighting fiercely for dominance 101 102 (Marty 2015) and dominant males able to monopolize matings with fertile females (Engelhardt et al. in revision). The male hierarchy, particularly the first three ranks, is so important that it is 103 104 clearly signalled in the occurrence and structure of loud calls (Neumann et al. 2010). Based on 105 these observations, we can expect male reproductive skew in favour of dominant males as 106 observed in other primates (reviewed in Widdig 2013), meaning that many infants sired during a male's tenure will share paternal genes. At the same time, the male hierarchy in crested 107 108 macaques is highly dynamic (Neumann et al. 2011), with high takeover rates resulting in a mean alpha tenure of only 12 months (Marty et al. 2015), so infants born in different years often have 109 110 different fathers. However, the genetic consequences of male reproductive strategies at the 111 population level remain unclear as no study has investigated male reproduction in crested macaques using genetic data. High reproductive skew may result in lower genetic variation as 112 only few, top-ranking males pass on their genes to the next generation; however, the high 113 takeover rate in alpha male position may counteract the effect of reproductive monopolisation 114 and contribute to the maintenance of genetic variation in the population. 115

116 The second reasons why we need genetic studies of crested macaques is that they are the most threatened Sulawesi macaques, and are Critically Endangered (IUCN 2016). Genetic studies of 117 crested macaques are limited to mitochondrial and autosomal DNA phylogeny (Evans et al. 118 1999, 2003). The degree of gene flow and the risk of inbreeding depression remain unclear for 119 120 the remaining populations of crested macaques. Furthermore, many animals, rescued from 121 illegal captivity and currently held in sanctuaries, await release into the wild. We cannot determine the genetic value of these individuals for wild populations until genetic evaluations 122 are feasible. It is important to detect hybrids amongst these rescued individuals to avoid 123 124 releasing them into hybrid-free populations. Finally, we need to understand the genetic variation

in the largest population remaining in its natural distribution range, Tangkoko. This information
is highly relevant to conservation management. However, we still lack genetic markers useful
for such analyses in crested macaques.

128 The first aim of this study was to identify highly polymorphic microsatellite (short tandem repeats or STR) markers for reliable genotyping in crested macaques. Testing primers originally 129 designed for other, usually closely related species (cross-species amplification) is often the 130 131 cheapest and fastest way to define a set of useful markers. Our second aim was to test the suitability of the selected markers. To do this, we determined maker polymorphism and checked 132 for Hardy-Weinberg equilibrium and Mendelian inheritance between known mother-offspring 133 134 pairs. Our third aim was to assign paternity to the Tangkoko animals and determine the degree of male reproductive skew (using the B index, Nonacs 2000, 2003) which we predicted to be 135 high based on the observed mating skew (Engelhardt et al. in revision). We predicted a low 136 degree of extra-group paternities and natal breeding, given that a few males monopolize all 137 receptive females. As a final aim, we investigated whether this isolated population shows signs 138 139 of loss of heterozygosity by comparing observed and expected heterozygosity, as well as evaluated estimates of inbreeding in this fragmented population. 140

141

142 Methods

143 <u>Study population</u>

We studied crested macaques at Tangkoko Reserve (1N 32'39", 125E 12'42"), North
Sulawesi, Indonesia. A recent study in the reserve estimated the population size to be less than
2000 individuals (Palacios et al. 2012). Tangkoko Reserve borders another nature reserve,
Duasudara Reserve, but is disconnected from all other forested areas in North Sulawesi. The
number of crested macaques currently living in Duasudara Reserve is unknown, but preliminary

149 data suggest it to be very low (Palacios et al. 2012). However, there may be some genetic150 exchange between individuals in the two reserves.

As in other macaque species, female crested macaques stay in their natal groups for life, forming matrilines, while males emigrate from their natal group. Males are fully grown when they emigrate and frequently challenge alpha males in another group when immigrating (Marty et al. 2015). Although females give birth year-round, they are moderately seasonal (Marty et al. 2016) with an inter-birth interval of about 22 months (Marty et al. 2015).

The Macaca Nigra Project observes three groups (R1, R2, PB) almost daily (R1 and R2 since 156 2006 and PB since 2008 until present) collecting behavioural data including aggressive 157 interactions and their outcomes through focal animal and ad libitum sampling (Altmann 1974). 158 We also recorded births, deaths, and migration events. All adult individuals and sampled infants 159 were individually recognised. During our study period, the home range of group R1 overlapped 160 with that of R2 and PB. All three groups also overlapped with other, non-study groups. We 161 162 individually recognised all adult individuals of the three groups as well as infants used for 163 paternity analysis in this study. Group size ranged between 36 and >100 individuals across years. 164

We used the David's score (de Vries et al. 2006) to assess dominance rank on a matrix of proportions of wins calculated for each male-male dyad. We calculated David scores using the package "Steepness" (Leiva&de Vries 2011) in R (RTeam 2009). We used either hormonal data or data of sex skin swelling size to assess conception windows (for details see Higham et al. 2012). In addition, we combined demographic and hierarchy data to compute annual alpha tenure (A. Engelhardt, C. Neumann, P. Marty unpublished data).

171

172 <u>Sample collection</u>

We collected non-invasive faecal samples immediately after defecation from 176 individually 173 174 recognized animals from all three groups from 2006 onwards. We collected up to three samples for each individual. Following the two-step alcohol-silica storage protocol (Nsubuga et al. 175 2004), we placed 1-2 g from the surface of fresh faeces into a 50 ml plastic tube filled with 30 176 ml of 99% ethanol for at least 24 hrs. Subsequently, we placed the sample in another tube filled 177 with 30 ml of silica beads and stored it at room temperature until extraction. In a few cases, we 178 179 collected ejaculates from males, which we stored in 98% ethanol at room temperature until extraction. We considered any adult males present or immigrating into our study groups during 180 our study period as potential sires. We defined adult males as larger than fully grown females, 181 182 with fully erupted canines and completely descended testes. We obtained DNA samples for 54 of 56 potential sires (96%), including all adult males present in one of the three study groups 183 since 2006. For one male, however, we only obtained one sample and the DNA obtained was 184 185 of such low quality that it amplified successfully at only nine loci.

186 We also obtained faecal and blood samples during regular health checks of seven crested187 macaques (one of each per individual) from Dublin Zoo.

188

189 *DNA extraction*

We extracted DNA from 100-150 mg of faeces with the GEN-IAL® all-tissue DNA extraction
kit following the manufacturer's instructions with the exception that we eluted DNA in distilled
water.

193

194 *Identification of polymorphic markers*

195 *a: Testing potential markers via cross-species amplification*

196 We tested 39 microsatellite loci previously described to be polymorphic in rhesus (*M. mulatta*),

197 long-tailed (M. fascicularis) and Barbary (M. sylvanus) macaques (Nürnberg et al. 1998;

198 Engelhardt et al. 2006; Brauch et al. 2008; Widdig et al. 2017) for allele amplification and

polymorphism with a set of nine different PCR conditions to increase the chances of successful 199 200 cross-species amplification (cf. Moore et al. 1991) in crested macaques. For this, we combined three different magnesium salt concentrations (1.5 mM, 2.0 mM, 2.5 mM) with three different 201 202 annealing temperatures (56, 58 and 60 °C or 51, 53 and 55 °C, depending on primer pair). In this step, we used a high quality pooled DNA sample (from blood) from the seven Dublin Zoo 203 204 individuals. When we obtained a readable product for a primer pair, we selected the condition that yielded the highest concentration of the specific product and fewer stutters for individual 205 206 genotyping and polymorphism check. We included the matching faecal and blood samples from the seven Dublin crested macaques to confirm that genotypes obtained from faecal samples 207 matched those from blood samples. Finally, we tested Mendelian inheritance by individually 208 amplifying DNAs from known mother-offspring pairs. 209

210

211 *b: Genotyping and determination of alleles*

To genotype the 176 subjects, we used a two-step multiplex polymerase chain reaction (PCR) 212 213 approach (modified from Arandjelovic et al. 2009). First, we amplified all loci in a multiplex approach using 4 µL of DNA extract (diluted 1:50 or 1:100), of 0.2 µL H20, 2 µL 10x Master 214 Taq Buffer with Mg2+ (5PRIME®, 500 mM KCl, 100 mM Tris-HCl pH 8.3, 15 mM 215 Mg(OAc)2), 2 µl 5x TaqMaster PCR Enhancer (5PRIME®), 0.8 µL dNTPs (10 mM), 1.2 µL 216 217 MgCl (25 mM), 0.4 µL (10 pmol) of 12 unlabelled forward and reverse primers, respectively, and 0.2 µL 5PRIME® Tag DNA Polymerase (5 U/µL, Enzyme storage Buffer: 20 mM Tris·HCl 218 219 pH 8.0, 100 mM KCl, 0.1 mM EDTA, 1 mM DTT, 50% glycerol, 0.5% Tween®20, 0.5% Igepal®CA-630) in an Eppendorf® Master Cycler Gradient. We started with 2 min of 220 denaturation at 94 °C, then ran 30 cycles of 20 sec denaturation at 94 °C, 30 sec of annealing at 221 54 °C, 30 sec of elongation at 70 °C and ended with 10 min of final elongation at 70 °C. 222 Following the multiplex approach, we ran single PCRs to amplify one locus at a time using 223 224 a similar protocol with specific annealing temperatures per primer pair (Table 1). Specifically,

we amplified 1 µL of multiplex PCR with 13.7 µL H20, 2 µL 10x Master Tag Buffer with 225 Mg2+ (5PRIME®, 500 mM KCl,100 mM 206 Tris-HCl pH 8.3, 15 mM Mg(OAc)2), 0.5 µl 5x 226 TaqMaster PCR Enhancer (5PRIME®), 0.8 µL dNTPs (10 mM), 0.8 µL MgCl (25 mM), 0.5 227 µL (10 pmol) of each primer labelled (HEX or FAM) forward and unlabelled reverse, and 0.2 228 µL 5PRIME® Taq DNA Polymerase (5U/μL, Enzyme storage Buffer: 20 mM Tris·HCl pH 8.0, 229 100 mM KCl, 0.1 mM EDTA, 1 mM DTT, 50% Glycerol, 0.5% Tween 20, 0.5% Igepal CA-230 630). We prepared singleplex PCR products for analysis by diluting PCR products between 231 1:25 and 1:500, and mixing 1.5 µL of diluted product into 14 µL of Hi-Di Formamide buffer 232 mixed with a size standard (HD400 from Applied Biosystems®). Finally, we ran amplicons on 233 234 an ABI 3130xL sequencer and determined allele sizes with PeakScanner (Applied 235 Biosystems[®]).

We analysed the samples in two laboratories (German Primate Center and Max-Planck Institute for Evolutionary Anthropology), with the identical protocols and equipment. We compared five individuals genotyped in both laboratories on the 12 markers and found genotype inconsistency in 2 of the 118 alleles, giving an error rate of 0.016.

240

241 *c: Modification of markers*

242 Many of the tested primer pairs produced unspecific products, typically detected as three or more differently sized amplicons resulting from the simultaneous amplification of two or more 243 loci (Smith et al. 2000). Since only 7 markers repeatedly produced up to two alleles per 244 individual, we modified specific primers for crested macaques for the other five identified 245 markers (Table 1). For this, we located sequences closer to the repetitive sequence than the 246 247 respective original primers. We then generated ligation of PCR products of the specific microsatellites into plasmid vector pCR®2.1-TOPO® with the TOPO TA Cloning®Kit 248 (INVITROGEN, Carlsbad, USA) followed by colony hybridisation as described in Takenaka 249 250 et al. (1993). We isolated plasmids containing the specific repeats from E. coli using the

QIAprep Spin Miniprep Kit (Qiagen). Next, we conducted fluorescent sequencing with the
Autocycle Sequence Kit Big Dye in the ABI Prism 3100 sequencer (Applied Biosystems, Foster
City, USA). Finally, we synthesised the selected primer sequences with Thermo Hybaid, Ulm,
Germany (Table 1). There may be further additional suitable markers among those we tested,
particularly if they are optimised for the species.

256

257 *d: Final marker selection*

We selected the 12 best markers using the following criteria: 1) we preferred markers with tetra-258 repeats over di-repeats, 2) amplification success at least 50%, 3) markers that were polymorphic 259 with at least 3 alleles) and 4) markers with reliable allele size scoring (no or few 260 stutters/multiple peaks). As faecal samples contain only a small amount of DNA and a high 261 level of allelic dropouts (Bayes et al. 2000), we genotyped three independent faecal samples for 262 263 each individual if available. Based on previous studies (Engelhardt et al. 2006; Brauch et al. 2008), we accepted a heterozygous genotype only if two different samples of the same 264 265 individual showed the same result in at least four amplifications; likewise, we accepted a 266 homozygous genotype if it was consistent in at least six amplifications (Taberlet et al. 1996). If we identified a third allele during analysis, we doubled the number of amplifications. 267

268

269 *Testing the suitability of selected markers*

270 a: Polymorphic information content and Hardy-Weinberg equilibrium

To investigate the suitability of our markers, we first calculated the polymorphic information content (PIC), an estimate of the discriminating power of markers (ranging from 0-1, from no allelic variation to only new alleles) (Botstein et al. 1980). We also tested markers for deviation from Hardy-Weinberg equilibrium (HWE). We considered that deviation from the HWE would indicate genotyping problems, such as segregating null alleles or incorrectly distinguished alleles. 277

278 b: Assessment of Mendelian inheritance

We investigated whether behavioural mothers (known from behavioural observations, i.e. association and nursing) were also the genetic mothers by testing Mendelian inheritance for 65 mother-offspring pairs through genotype matching using the 12 best markers (including the 5 specifically designed for crested macaques).

283

284 *Investigating paternity distribution*

a: *Paternity determination*

286 We used the 65 mother-offspring pairs in paternity analysis. Our paternity dataset included all offspring born into the three groups between 2006 and 2011 that we could sample. Following 287 a conservative approach, we assigned paternity only when exclusion and likelihood calculations 288 289 revealed the same father (cf. Widdig et al. 2017). In our exclusion method, we assigned paternity to the male who had no mismatches with a given mother-offspring pair across all loci 290 291 while all other potential sires mismatched the offspring at two or more loci (strict exclusion). 292 We also assigned paternity to the male with no mismatches with a given mother-offspring pair across all loci while one or more males mismatched the offspring at one locus only (relaxed 293 294 exclusion). We used the program FINDSIRE (https://www.uni-kiel.de/medinfo/ 295 mitarbeiter/krawczak/download/) to establish paternity exclusion. We used the same set of males (i.e., all potential sires) to calculate likelihood-odds (LOD) scores and confidence levels 296 and confirm sires using likelihood analyses in CERVUS 3.0. We used the following parameters 297 in CERVUS: simulated offspring: 100; number of candidate fathers: 56; proportion of candidate 298 fathers sampled: 0.96; proportion of loci typed: 0.99; proportion of loci mistyped: 0.01; 299 300 minimum number of typed loci: 10. To assess the proportion of extra-group paternities, we checked whether the assigned sire was a member of the infant's birth group at the time of 301 infant's conception using demographic and hormonal data (A. Engelhardt, unpublished data). 302

Given the delay in natal dispersal, we also investigated whether the assigned sire was natal to
the birth group of the infant to detect cases of natal breeding using demographic data (A.
Engelhardt, unpublished data).

306

307 *b: Degree of male reproductive skew*

We determined the degree of male reproductive skew using Nonacs' B Index (Nonacs 2000, 308 2003) with Skew Calculator 2003 (http://www.eeb.ucla.edu/Faculty/Nonacs/PI.htm). Positive 309 values of the B index suggest that the skew is higher than expected, while negative values 310 suggest that reproduction is more equally distributed than expected (Kutsukake & Nunn 2006). 311 312 Furthermore, an index close to 0 indicates a random distribution of paternities across potential sires, whereas values close to 1 suggest a high monopolization of reproduction by a single male. 313 314 The advantage of the B index is that it can incorporate the total number of days adult males 315 spent in a given group per year. We included information on group membership in the skew calculation based on demographic data. The program also computes 95% confidence intervals 316 317 (CI) with the width of the confidence interval revealing the precision of the estimates. If the CI 318 includes zero, then the distribution of paternity among group males is not significantly different from random. 319

320 As our sampling effort was not consistent across the study period, the skew analysis includes only years and groups in which we sampled at least 45% of offspring born 321 (mean±SD=66.8%+28.6%). Therefore, we restricted the skew analysis to offspring born 322 between 2007 and 2009 in R1 and R2 and born in 2009 in PB, giving 51 offspring with solved 323 324 paternity. Although crested macaques are only moderately seasonal, we calculated the annual 325 skew per group and year. Ideally, we should determine the degree of skew in successful conceptions during each alpha tenure, however, the number of offspring conceived per alpha 326 tenure was low due to the typically short tenure (mean 12 months; see Marty et al. 2015). 327

328

329 Assessing genetic variation and inbreeding

330 For each of the selected markers, we computed standard population genetic parameters of genetic variation within a population. First, we calculated the expected heterozygosity (He), 331 defined as the probability that an individual in a population is heterozygous at a given locus. 332 Second, we determined the observed heterozygosity (Ho) by counting the frequency of 333 heterozygous individuals per locus. If the observed heterozygosity is lower than expected, this 334 335 indicates inbreeding, while a higher than expected heterozygosity suggests a mixture of two previously isolated populations (Hartl & Clark 1997). Furthermore, we determined inbreeding 336 coefficients (FIS), where positive values indicate a deficit of heterozygosity (i.e., inbreeding) 337 while negative values indicate an excess of heterozygosity (Hedrick 2000). We conducted all 338 calculations (including PIC and HWE) in CERVUS 3.0 (Kalinowski et al. 2007) except the 339 Wright F statistics (FIS), which we computed in FSTAT (version 2.9.3.) (Goudet 2001). 340

341

342 Ethical note

343 Research complied with protocols approved by the Indonesian Institute for Science and Technology (RISTEK) and the Indonesian Ministry of Forestry (PHKA) and adhered to the 344 legal requirements of Indonesia and Germany. We received permits to collect samples and 345 346 export DNA extracts from the Indonesian Ministry of Forestry. Furthermore, we carried out our research in compliance with the animal care regulations and the principles of the American 347 Society of Primatologists and the German Primate Center for the ethical treatment of non-348 human primates. We collected faecal samples from wild and captive individuals non-invasively 349 after the animals left the site without disturbing, threatening or harming them in their natural 350 351 behaviour, and obtained blood samples as part of the regular health check.

352

353 **Results**

354 *<u>Identification of polymorphic markers</u>*

Overall, 31 % (12/39) of the markers we tested were suitable for investigating the crested macaque population at Tangkoko. These included 10 tetra-nucleotide and 2 di-nucleotide loci (Table 1) with 4-9 alleles per locus (Table 2). We typed 176 individuals at 12±0.3 (mean±SD) loci (Table 2).

- 359
- 360 *Testing the suitability of selected markers*
- 361 *a: Polymorphic information content and Hardy-Weinberg equilibrium*
- The PIC ranged 0.538 0.790 with a mean of 0.658±0.075 (mean±SD) (Table 2) suggesting our markers had high discriminating power. We detected no significant deviation from Hardy-Weinberg or evidence of null alleles (Table 1).
- 365
- 366 *b: Mendelian inheritance*
- We confirmed all 65 maternities (assigned by behavioural observations) through genotype
 matching (65 pairs * 10-12 loci) with one mismatch in one mother-offspring pair.
- 369

370 *Investigating paternity distribution*

a: *Paternity determination*

Our dataset included 65 offspring for which we could solve 63 paternities (97%). In 40 cases, we excluded all males on at least two loci, except for the assigned sire, who matched the offspring-mother pair at all loci (strict exclusion). In 14 cases, the assigned sire had no mismatch with the respective mother-offspring pair, but we excluded the next candidate sire at only one locus (relaxed exclusion). In 8 further cases, the assigned sire had one mismatch with the given infant, while the next likely sires had at least two mismatches (best match). In one case, two males matched the infant-mother pair at all loci (tie) and both males were also present in the group around the conception of the infant. In this case, we accepted the male assigned by CERVUS (Kalinowski et al. 2007) as the sire. In all cases, CERVUS supported the sires assigned based on exclusion rules (95% confidence level, see Supplement for an overview of genotypes and trios). In the remaining two cases, we did not assign paternity because the exclusion and likelihood approach did not reveal the same father. We found no evidence of extra-group paternity or natal breeding in the solved paternity cases.

385

386 *b: Degree of male reproductive skew*

Although 18 males sired the 63 infants investigated, the mean male reproductive skew per group 387 and year as assessed by the B index was relatively high (mean±SD: 0.330±0.267, range: 0.021 388 to 0.672). The B index was significantly different from a random distribution across groups and 389 years (e.g., very high for all years in group R2), except for two of three years in group R1 (Table 390 391 3). A posteriori analysis showed that the sex ratio (m/f) was negatively related to the B index; a female biased sex ratio significantly increased the B index (Spearman rho=-0.857, N=7, 392 393 p=0.014) (Table 3). Finally, the mean proportion of alpha paternity was 65% per year with high variation across groups (29-100%). 394

395

396 Assessing genetic variation and inbreeding

The observed heterozygosity (Ho) ranged from 0.665 to 0.856, and expected heterozygosity 397 398 (He) from 0.613 to 0.818 (Table 2). The observed heterozygosity mean (mean±SD=0.766±0.059) 399 was greater than the mean expected heterozygosity 400 (mean±SD=0.708±0.059) (Table 2) suggesting no risk of inbreeding at this point in time in our study groups (see Hartl & Clark, 1997, for comparison). In other words, while we expected 401 402 around 70% of individuals to be heterozygous at a given locus under random mating conditions, on average approximately 76% of individuals were heterozygous. Similarly, the mean FIS 403 across the three groups was -0.082±0.035 (mean±SD) with FIS consistently below zero for all 404

12 polymorphic loci, indicating an excess of observed heterozygosity (see Hedrick, 2000, for
comparison). In other words, individuals were less related than expected under random mating.
Finally, we found no major differences between groups in terms of number of alleles per locus
and degree of heterozygosity (Table 2), suggesting comparable estimates of genetic variability
despite different group size, degree of skew and duration of alpha tenure.

410

411 **Discussion**

Our results show that the 12 selected microsatellite markers provide reliable information on 412 individual genotypes in crested macaques and are useful for various applications in field studies 413 on this species. Specifically, they provided high confidence in paternity assignment, a relatively 414 high level of polymorphic information content and genetic variation (assessed by 415 heterozygosity and inbreeding coefficients) and a high accuracy of allele characterization (i.e., 416 low occurrence or absence of mutations). Furthermore, they mainly comprise tetra-nucleotide 417 418 repeats, which are usually easier to analyse and thus enhance the reliability of genotyping. 419 Altogether, the selected markers fulfil important genetic and technical criteria that are critical for the precision and efficacy of high-throughput genotyping (Butler et al. 2001). 420

We report highly polymorphic markers in Sulawesi macaques. Although we used primers 421 formerly applied to other macaque species, several markers did not generate satisfying PCR 422 products. We thus modified specific primers for crested macaques that produced much more 423 424 reliable amplification results. However, given that Sulawesi macaques split from their common ancestor with southern pig-tailed macaques from Borneo (M. nemestrina) only in the early to 425 middle Pleistocene (Fooden 1969; Evans et al. 1999), most, if not all, of the loci used in this 426 427 study are likely informative in the other Sulawesi macaque species too. With the validated markers and improved primers, we thus provide an important tool for conservation management 428 to assess gene flow, heterozygosity and inbreeding depression of small and/or isolated 429

populations across the whole island. Furthermore, with this set of markers, we will be able to
conduct more detailed studies of population genetics, sexual selection, behaviour and
sociobiology, including parentage data. We encourage the application of the selected markers
to other Sulawesi macaque species.

434 We assigned paternity to 97% of offspring sampled with 95% confidence, demonstrating the high analytical power of the marker set and its usefulness for studies of sexual selection and 435 436 reproductive success. Although we cannot draw conclusions for the two offspring with unsolved paternity, all cases of solved paternity show no indication of extra-group paternity and 437 438 natal breeding. This is interesting, given that male crested macaques do not disperse until they 439 fully developed, and their competitive ability is sufficient for challenging alpha males in nonnatal groups (Marty et al. 2015). Furthermore, groups are large enough for unrelated potential 440 mates to coexist in the natal group. It thus seems that male crested macaques need to migrate 441 and successfully take over the alpha position to reproduce (Marty et al. 2016). It is also 442 surprising that we found no extra-group paternity. Adjacent groups meet frequently and groups 443 444 are too large and the vegetation is too dense for males to oversee the whole group. This suggests that females ready to conceive are either well mate-guarded during inter-group encounters, or 445 refrain from mating with non-group males. More detailed behavioural observations during 446 447 intergroup encounters are needed to show which of these two explanations hold true for crested macaques. 448

As predicted from mating observations, we found a skew in male reproduction towards alpha males. The mean alpha paternity was 65% and ranged 29-100% across years and groups. Similarly, the degree of skew varied considerably across groups. Notably, our study on crested macaques found the highest B index reported so far for any primate (maximum: 0.672, mean: 0.330). In a study of free-ranging rhesus macaques, the skew in one large group varied 0.049-0.106 across six consecutive years (Widdig et al. 2004) and in one small group, the mean B index was 0.084 over two consecutive years (Dubuc et al. 2011). In wild Assamese macaques
(*M. assamensis*), the mean B index was only 0.087 over six years in one group, with the alpha
share of paternity limited to 29% (Sukmak et al. 2014).

458 Takeover rates had a negative effect on reproductive skew. The largest group, R1, generally had a lower skew and was subject to frequent alpha takeovers (i.e., the male hierarchy was 459 dynamic), while group R2 showed skew values as high as 0.672, but had fewer takeovers (i.e., 460 461 extended alpha tenure). These data are in line with results from species with extraordinary long alpha tenures, such as capuchin monkeys (Cebus capucinus), with an observed B index 462 calculated across eight alpha tenure periods varying from -0.125 to 0.473 (mean: 0.274) (Muniz 463 464 et al. 2010). Similarly, mountain gorillas (Gorilla beringei beringei) showed B indices between 0.337-0.432 in four groups containing multiple males of long tenure (Bradley et al. 2005). It is 465 surprising, however, that the skew in R2 study group was higher than in the gorilla study, where 466 a single male usually monopolizes all reproduction in his group. Skew calculations across these 467 three studies are comparable as they were calculated over the timeframe of alpha male tenure 468 469 typical for each species. In other words, for crested macaques with their extraordinary short alpha tenure we computed annual skew per group, while in the two other species with long 470 tenure, skew was computed over multiple years of alpha tenure per group. One potential reason 471 472 for the comparatively large skew in crested macaques is that male crested macaques need to maximize their reproductive effort in a short timeframe. Hence, alpha tenure length might affect 473 the inter-specific variation in reproductive skew. However, our study might also provide a 474 potential explanation for the intra-specific variation in skew. A more female biased sex ratio 475 476 significantly increased the B index which suggests that when more females are available, there 477 is more room for a few males to successfully monopolize receptive females, in contrast to when more male competitors are present. This supports the hypothesis that enhanced male 478 479 monopolization, among other factors, results in higher degree of reproductive skew (Ostner et 480 al. 2008; Gogarten & Koenig 2012).

The high degree of male reproductive skew observed in our study animals did not translate into lower genetic variation in the population than we would expected under random mating. This is interesting given that only a few dominant males pass their genes into the next generation. Most likely, the high rates of alpha male takeover reported for this population counterbalance this effect. We need more detailed data on genetic variation in relation to tenure length to understand this process better.

487 Our study animals reflect a geographically isolated population of a Critically Endangered species, but our analysis indicates no recent threat of considerable loss of heterozygosis and/or 488 of inbreeding depression in the study population. Compared to studies of other macaque 489 490 species, mainly using different markers (e.g. M. mulatta, Bercovitch and Nürnberg 1997; M. sinica, Keane et al. 1997; M. sylvanus, Kümmerli and Martin 2005; M. fuscata, Inoue and 491 Takenata 2008; M. assamensis, Sukmak et al. 2014), our markers were highly polymorphic. 492 Despite the small population size, it is possible that males migrate in and out of the Tangkoko 493 population, contributing to the genetic variability observed. 494

495 In contrast to our results, we found no polymorphism in a set of mtDNA markers in another study using a subset of the individuals included here (i.e., 12 females and 4 non-natal males 496 from two groups) (A. Engelhardt, unpublished data). This could indicate that the population of 497 498 Tangkoko may already be inbred or stems from one single matriline. To determine the degree of inbreeding in crested macaques at Tangkoko more precisely, we will need extended studies 499 over a broader range of groups. Furthermore, we need studies investigating the links between 500 501 reproductive patterns, genetic variation and population demography over time to expand our understanding of viability of threatened populations in the wild. 502

In conclusion, we provide genetic markers useful for studies on the conservation management
and evolutionary biology of crested macaques, and likely of Sulawesi macaques in general.
Parentage analysis of these species can contribute insights to the relationship between social

style, reproductive patterns and relatedness among macaque species (Schülke & Ostner 2008). 506 The fact that the Tangkoko population of crested macaques is still genetically variable despite 507 its small size, isolation and the species' reproductive patterns gives hope that other endangered 508 509 primate species living in small, isolated populations may also retain a healthy gene pool, at least in the short term. However, while the population in Tangkoko does not seem to be suffering 510 from genetic depletion, other isolated populations of crested macaques might. With the 511 512 described markers at hand, we will now be able to assess and manage genetic variation across all populations of crested macaques scattered over North Sulawesi. 513

514

515 Acknowledgment

We are grateful to all members of the Macaca Nigra Project that contributed to sample 516 517 collection, to Jan-Boje Pfeifer for continuous logistic support of the project and to Muhammad 518 Agil for support of sample export. Furthermore, we greatly thank the Dublin Zoo for providing samples. Kerstin Fuhrmann, Stefanie Bley and Maren Keller are kindly acknowledged for 519 supporting genetic data production and analysis. Furthermore, we thank Linda Vigilant for 520 providing laboratory access. The study was funded by the German Research Council within the 521 Emmy-Noether programme (grant No. EN 719/1, 2 to AE, WI 1801/3-1 to AW) and the 522 University of Leipzig (to AW), partly together with the German Federal Ministry for Economic 523 Cooperation and Development, and the German Academic Exchange Service (to AE). We are 524 525 grateful to Christof Neumann and Pascal Marty for providing male hierarchy and tenure data, 526 as well as to Joanna Setchell and two anonymous reviewers for fruitful comments on an earlier version of the manuscript. Finally, we thank the German Primate Center and the Max-Planck 527 Institute for Evolutionary Anthropology for logistic support. 528

529

530 **References**

531	Arandjelovic, M., Guschanski, K., Schubert, G., Harris, T.R., Thalmann, O., Siedel, H., et al. (2009).
532	Two-step multiplex polymerase chain reaction improves the speed and accuracy of genotyping
533	using DNA from noninvasive and museum samples. <i>Molecular Ecology Resources</i> , 9, 28–36.
534	Bayes, M.K., Smith, K.L., Alberts, S.C., Altmann, J., Bruford, M.W. (2000). Testing the reliability of
535	microsatellite typing from faecal DNA in the savannah baboon. Conservation Genetics, 1, 173–
536	176.
537	Bercovitch, F.B., Nürnberg, P. (1997). Genetic determination of paternity and variation in male
538	reproductive success in two populations of rhesus macaques. <i>Electrophoresis</i> , 18, 1701–1705.
539	Botstein, D., White, R.L., Skolnick, M., Davis, R.W. (1980). Construction of a genetic linkage map in
540	man using restriction fragment length polymorphisms. American Journal of Human Genetics,
541	32, 314–331.
542	Bradley, B., Robbins, M.M., Williamson, E.A., Steklis, H.D. Steklis, N.G., Eckhardt, N., Boesch, C.,
543	Vigilant, L. (2005). Mountain gorilla tug-of-war: Silverbacks have limited control over
544	reproduction in multimale groups. Proceedings of the National Academy of Sciences, 102,
545	9418–9423.
546	Brauch, K., Hodges, K., Engelhardt, A., Fuhrmann, K., Shaw, E., Heistermann, M. (2008). Sex-specific
547	reproductive behaviours and paternity in free-ranging Barbary macaques (Macaca sylvanus).
548	Behavioral Ecology and Sociobiology, 62, 1453–1466.
549	Butler, J.M., Ruitberg, C.M., Vallone, P.M. (2001). Capillary electrophoresis as a tool for optimization
550	of multiplex PCR reactions. Fresenius Journal of Analytical Chemistry, 369, 200–205.

- Charpentier, M., Fontaine, M., Cherel, E., Renoult, J., Jenkins, T., Benoit, L., et al. (2012). Genetic
 structure in a dynamic baboon hybrid zone corroborates behavioural observations in a hybrid
 population. *Molecular Ecology*, 21, 715–731.
- 554 Duboscq, J., Agil, M., Engelhardt, A., Thierry, B. (2014). The function of postconflict interactions: new
- prospects from the study of a tolerant species of primate. *Animal Behaviour*, 87, 107–120.
- 556 Duboscq, J., Micheletta, J., Agil, M., Hodges, J.K., Thierry, B., Engelhardt, A. (2013). Social tolerance in
- 557 wild female crested macaques (*Macaca nigra*) in Tangkoko-Batuangus Nature Reserve,
- 558 Sulawesi, Indonesia. American Journal of Primatology, 75, 361–375. Dubuc, C., Muniz, L.,
- 559 Heistermann, M., Engelhardt, A, Widdig, A. (2011). Testing the Priority-of-Access model in a
- seasonally breeding primate species. *Behavioral Ecology and Sociobiology*, 65, 1615–1627.
- Engelhardt, A., Heistermann, M., Agil, M., Perwitasari-Farajallah, D., Higham, J.P. (in revision). A
 despotic mating system in a socially tolerant primate, the crested macaque.
- 563 Engelhardt, A., Heistermann, M., Hodges, J.K., Nuernberg, P., Niemitz, C. (2006). Determinants of
- 564 male reproductive success in wild long-tailed macaques (*Macaca fascicularis*)—male
- 565 monopolisation, female mate choice or post-copulatory mechanisms? *Behavioral Ecology and*566 *Sociobiology*, 59, 740–752.
- 567 Evans, B.J., Morales, J.C., Supriatna, J., Melnick, D.J. (1999). Origin of the Sulawesi macaques
- 568 (Cercopithecidae: *Macaca*) as suggested by mitochondrial DNA phylogeny. *Biological Journal of* 569 *the Linnean Society*, 66, 539–560.
- Evans, B.J., Supriatna, J., Andayani, N., Melnick, D.J. (2003). Diversification of Sulawesi macaque
 monkeys: decoupled evolution of mitochondrial and autosomal DNA. *Evolution*, 57, 1931–
 1946.
- Fooden, J. (1969). *Taxonomy and evolution of the monkeys of Celebes: (Primates: Cercopithecidae.)*.
 S. Karger.

- 575 Fooden, J. (1982). Ecogeographic segregation of macaque species. *Primates*, 23, 574–579.
- 576 Godinho R, López-Bao JV, Castro D, Llaneza L, Lopes S, Silva P, et al. (2015). Real-time assessment of
- 577 hybridization between wolves and dogs: combining noninvasive samples with ancestry
- 578 informative markers. *Molecular Ecology Resources*, 15, 317–328.
- 579 Gogarten, J.F. and Koenig, A. (2012) Reproductive seasonality is a poor predictor of receptive
- 580 synchrony and male reproductive skew among nonhuman primates, *Behavioral Ecology and*581 Sociobiology, 67, 123–134.
- 582 Goudet, J. 2001. FSTAT, a program to estimate and test gene diversities and fixation indices (version
 583 2.9.3)
- Groves, C.P. (1980). Speciation in *Macaca*: the view from Sulawesi. In D.G. Lindburg (Ed.), *The Macaques: Studies in Ecology, Behavior, and Evolution* (pp. 84-124). New York: Van Nostrand
 Reinhold Co.
- 587 Harms, V., Nowak, C., Carl, S., Muñoz-Fuentes, V. (2015). Experimental evaluation of genetic predator

identification from saliva traces on wildlife kills. *Journal of Mammalogy*, 96, 138–143.

- Hartl, D.L, Clark, A.G. (1997). Principles of population genetics, 3rd edition. Sinauer Associates,
 Sutherland.
- Hedrick, P.W. (2000). Genetics of populations. 2nd edition. Jones and Bartlett Publishers, Sudbury.
- Hewitt, G. (2000). The genetic legacy of the Quaternary ice ages. *Nature*, 405, 907–913.
- 593 Hill, W.C.O. (1974). Primates: Comparative Anatomy and Taxonomy: Cynopithecinae: Cercocebus,
- 594 *Macaca, Cynopithecus*. Edinburgh: Edinburgh University Press.
- 595 Inoue, E., Takenaka, O. (2008). The effect of male tenure and female mate choice on paternity in
- 596 free-ranging Japanese macaques. *American Journal of Primatology*, 70, 62–68.

- 597 IUCN (2016). *IUCN Red List*. <u>http://www.iucnredlist.org/initiatives/mammals/analysis/red-list-status</u>.
- 598 Accessed 15 July 2016.

Kalinowski, S.T., Taper, M.L., Marshall, T.C. (2007). Revising how the computer program CERVUS
accommodates genotyping error increases success in paternity assignment. *Molecular Ecology*,
16, 1099–1106.

- Keane, B., Dittus, W.P.J., Melnick, D.J. (1997). Paternity assessment in wild groups of toque macaques
 Macaca sinica at Polonnaruwa, Sri Lanka using molecular markers. *Molecular Ecology*, 6, 267–
 282.
- Knief, U., Hemmrich-Stanisak, G., Wittig, M., Franke, A., Griffith, S.C., Kempenaers, B., et al. (2015).
- 606 Quantifying realized inbreeding in wild and captive animal populations. *Heredity*, 114, 397–403.
- 607 Kümmerli, R., Martin, R.D. (2005). Male and female reproductive success in Macaca sylvanus in
- 608 Gibraltar: no evidence for rank dependence. International Journal of Primatology, 26, 1229–
- 609 1249. doi:10.1007/s10764-005-8851-0
- 610 Kutsukake, N., Nunn, C.L. (2006). Comparative tests of reproductive skew in male primates: The roles
- of demographic factors and incomplete control. *Behavioral Ecology and Sociobiology*, 60: 695–
 706.
- Leonard, J.A. (2008). Ancient DNA applications for wildlife conservation. *Molecular Ecology*, 17,
 4186–4196.
- Luikart, G., Sherwin, W.B., Steele, B.M., Allendorf, F.W. (1998). Usefulness of molecular markers for
 detecting population bottlenecks via monitoring genetic change. *Molecular Ecology*, 7, 963–
 974.
- 618 Marty, P.R. (2015). *Male migration and alpha male takeovers in crested macaques, Macaca nigra*.
- 619 PhD thesis, University of Göttingen.

620	Marty, P.R., Hodges, K., Agil, M., Engelhardt, A. (2015). Alpha male replacements and delayed
621	dispersal in crested macaques (Macaca nigra). American Journal of Primatology, 9999, 1–8.
622	Marty, P.R., Hodges, K., Agil, M., Engelhardt, A. (2016). Determinants of immigration strategies
623	in male crested macaques (<i>Macaca nigra</i>). Scientific reports, 6, 32028.
624	Melfi V. (2010). Selamatkan Yaki! Conservation of Sulawesi Crested Black Macaques Macaca nigra. In:
625	S. Gursky, J. Supriatna (Eds), Indonesian Primates (pp. 343–356). New York: Springer.
626	Montague, M.J., Disotell, T.R., Fiore, A. (2014). Population Genetics, Dispersal, and Kinship Among
627	Wild Squirrel Monkeys (Saimiri sciureus macrodon): Preferential Association Between Closely
628	Related Females and Its Implications for Insect Prey Capture Success. International Journal of
629	Primatology, 35, 169–187.
630	Moore, S.S., Sargeant, L.L., King, T.J., Mattick, J.S., Georges, M., Hetzel, D.J.S. (1991). The
631	conservation of dinucleotide microsatellites among mammalian genomes allows the use of
632	heterologous PCR primer pairs in closeley related species. <i>Genomics</i> , 10, 654–660.
633	Morin, P.A., Chambers, K.E., Boesch, C., Vigilant, L. (2001). Quantitative polymerase chain reaction
634	analysis of DNA from noninvasive samples for accurate microsatellite genotyping of wild
635	chimpanzees (Pan troglodytes verus). Molecular Ecology, 10, 1835–1844.
636	Muniz, L. Perry, S., Manson, J.H., Gilkenson, H., Gros-Louis, J., Vigilant, L. (2010). Male dominance and
637	reproductive success in wild white-faced capuchins (Cebus capucinus) at Lomas Barbudal, Costa
638	Rica. American Journal of Primatology, 72, 1118–1130.
639	Neumann, C., Assahad, G., Hammerschmidt, K., Perwitasari-Farajallah, D., Engelhardt, A. (2010). Loud
640	calls in male crested macaques (Macaca nigra) - a signal of dominance in a tolerant species.
641	Animal Behaviour, 79, 187–193.

- Neumann, C., Duboscq, J., Dubuc, C., Ginting, A., Irwan, A.M., Agil, M., Widdig, A., Engelhardt, A.
- 643 (2011). Assessing dominance hierarchies: validation and advantages of progressive evaluation
 644 with Elo-rating. *Animal Behaviour*, 82, 911–921.
- Nonacs, P. (2000). Measuring and using skew in the study of social behavior and evolution. *American Naturalist*, 156, 577–589.
- Nonacs, P. (2003). Measuring the reliability of skew indices: is there one best index? *Animal Behaviour*, 65, 615–627.
- Nowak, K., le Roux, A., Richards, S.A., Scheijen, C.P.J., Hill, R.A. (2014). Human observers impact
- habituated Samango monkeys' perceived landscape of fear. *Behavioral Ecology*, 25, 1199–1204.
- 651 Nsubuga, A.M., Robbins, M.M., Roeder, A.D., Morin, A., Boesch, C., Vigilant, L. (2004). Factors
- affecting the amount of genomic DNA extracted from ape faeces and the identification of an
- 653 improved sample storage method. *Molecular Ecology*, 13, 2089–2094.
- Nürnberg, P., Sauermann, U., Kayser, M., Lanfer, C., Manz, E., Widdig, A., et al. (1998). Paternity
- assessment in rhesus macaques (*Macaca mulatta*): Multilocus DNA fingerprinting and PCR
- 656 marker typing. *American Journal of Primatology*, 44, 1–18.
- Ostner, J., Nunn, C.L., Schülke, O. (2008). Female reproductive synchrony predicts skewed paternity
 across primates. *Behavioral Ecology*, 19, 1150–1158.
- Palacios, J.F.G., Engelhardt, A., Agil, M., Hodges, K., Bogia, R., Waltert, M. (2012). Status of, and
 Conservation Recommendations for, the Critically Endangered Crested Black Macaque, *Macaca*
- 661 *Nigra,* in Tangkoko, Indonesia. *Oryx*, 46, 290–297.
- Ram, M.S., Marne, M., Gaur, A., Kumara, H.N., Singh, M., Kumar, A., et al. (2015). Pre-Historic and
- 663 recent vicariance events shape genetic structure and diversity in endangered lion-tailed
- 664 macaques in the Western Ghats: implications for conservation. *PLoS ONE*, 10, e0142597.

- Riley, E.P. (2010). The endemic seven: four decades of research on the Sulawesi macaques.
- 666 *Evolutionary Anthropology*, 19, 22–36.
- 667 Roos, C., Zinner, D., Kubatko, L.S., Schwarz, C., Yang, M., Meyer, D., et al. (2011). Nuclear versus
- 668 mitochondrial DNA: evidence for hybridization in colobine monkeys. *BMC Evolutionary Biology*,
- 669 11, 77.
- Schülke, O., Ostner, J. (2008). Male reproductive skew, paternal relatedness and female social
 relationships. *American Journal of Primatology*, 70, 1–4.
- 672 Schwartz, M.K., Luikart, G., Waples, R.S. (2007). Genetic monitoring as a promising tool for
- 673 conservation and management. *Trends in Ecology and Evolution*, 22, 25–33.
- 674 Smith, K.L., Alberts, S.C., Bayes, M.K., Bruford, M.W., Altmann, J., Ober, C. (2000). Cross-species
- 675 amplification, non-invasive genotyping, and non-Mendelian inheritance of human STRPs in
 676 Savannah baboons. *American Journal of Primatology*, 51, 219–227.
- 677 Sukmak, M., Wajjwalku, W., Ostner, J., Schülke, O. (2014). Dominance rank, female reproductive
- 678 synchrony, and male reproductive skew in wild Assamese macaques. *Behavioral Ecology and*679 *Sociobiology*, 68, 1097–1108.
- 680 Syrůčková, A., Saveljev, A.P., Frosch, C., Durka, W., Savelyev, A.A., Munclinger, P. (2015). Genetic

relationships within colonies suggest genetic monogamy in the Eurasian beaver (*Castor fiber*). *Mammal Research*, 60, 139–147.

Taberlet, P., Griffin, S., Goossens, B., Questiau, S., Manceau, V., Escaravage, N., et al. (1996). Reliable
genotyping of samples with very low DNA quantities using PCR. *Nucleic Acids Research*, 24,
3189–3194.

- Takenaka, O., Takasaki, H., Kawamoto, S., Arakawa, M., Takenaka, A. (1993). Polymorphic
 microsatellite DNA amplification customized for chimpanzee paternity testing. *Primates*, 34,
 27–35.
- Thierry, B., Iwaniuk, A.N., Pellis, S.M. (2000). The influence of phylogeny on the social behaviour of
 macaques (Primates: Cercopithecidae, genus *Macaca*). *Ethology*, 106, 713–728.
- Thierry, B. (2004). Social epigenesis. In: B. Thierry, M. Singh, W. Kaumanns (Eds.). *Macaque societies: A model for the study of social organization* (pp. 267–289). Cambridge: Cambridge University
 Press.
- Tosi, A.J., Morales, J.C., Melnick, D.J. (2003). Paternal, maternal, and biparental molecular markers
 provide unique windows onto the evolutionary history of macaque monkeys. *Evolution*, 57,
 1419–1435.
- van Horn, R.C., Altmann, J., Alberts, S.C. (2008). Can't get there from here: inferring kinship from
 pairwise genetic relatedness. *Animal Behaviour*, 75, 1173–1180.
- 699 Waits, L.P., Paetkau, D. (2005). Non-invasive Genetic Sampling Tools for Wildlife Biologists: A Review
- of Applications and Recommendations for Accurate Data Collection. *Journal of Wildlife*
- 701 *Management*, 69, 1419–1433.
- Widdig, A. (2013). The impact of male reproductive skew on kin structure and sociality in multi-male
 groups. *Evolutionary Anthropology*, 22, 239–250.
- 704 Widdig, A., Bercovitch, F.B., Streich, W.J., Sauermann, U., Nürnberg, P., Krawczak, M. (2004). A
- 705 longitudinal analysis of reproductive skew in male rhesus macaques. *Proceedings of the Royal*
- 706 *Society of London Series B*, 271, 819–826.

- 707 Widdig, A., Muniz, L., Minkner, M., Barth, Y., Bley, S., Ruiz-Lambides, A., Junge O., Mundry, R. Kulik, L.
- 708 (2017). Low incidence of inbreeding in a long-lived primate population isolated for 75 years.
- 709 Behavioral Ecology and Sociobiology, 71, 1–15.

Table 1: Characterization of 12 primer pairs for amplifying polymorphic microsatellite loci in crested macaques	with PCR conditions, c	leviation from Hardy-
Weinberg equilibrium and estimated null allele frequency. F indicates forward primers and R indicates reverse p	orimers.	

Locus	Repeat pattern	Length of PCR product Zoo [bp]	Length of PCR product Tangkoko [bp]	Annealing temperature [°C]	Hardy- Weinberg deviation ^a	Estimated null allele frequency	Primer sequence (5'-3') (including modified primers)	Reference	
D1S548	Tetra	181-201	185-209	58	n.s.	-0.0394	F: GAACTCATTGGCAAAAGGAA R: GCCTCTTTGTTGCAGTGATT	Lathuilliere and Menard 2001	
D3S1768	Tetra	129-137	129-157	58	n.s.	-0.046	F: GGTTGCTGCCAAAGATTAGA R: AACTACATGATTCTAGCACA	Lathuilliere and Menard 2001	
D5S1457	Tetra	123, 127, 131	123-139	60	n.s.	-0.0609	F: TAGGTTCTGGGCATGTCTG R: TTGCTTGGCACACTTCAGG	Bayes et al. 2000	
D6S493*	Tetra	261-269'**	139-159***	58	n.s.	-0.0374	F: GCAACAGTTTATGCTAAAGC R: TTCCATGGCAGAAATTGTTT	Nürnberg et al., 1998	
D6S501*	Tetra	163-179**	129-145***	58	n.s.	-0.0345	F: GCTGGAAACTGATAAGGGCT R: CTTTATCTTTAATATAGGATTATTGG	Lathuilliere and Menard 2001	
D7S2204	Tetra	171-247	220-268	58	n.s.	-0.0579	F: TCATGACAAAACAGAAATTAAGTG R: AGTAAATGGAATTGCTTGTTACC	Lathuilliere and Menard 2001	
D10S1432	Tetra	137-145	132-148	58	n.s.	-0.0773	F: CAGTGGACACTAAACACAATCC R: TAGATTATCTAAATGGTGGATTTCC	Lathuilliere and Menard 2001	
D11S925	Di	205-221	179-237	60	n.s.	-0.0379	F: GAACCAAGGTCGTAAGTCC R: TAGACCATTATGGGGGCAAA	Lathuilliere and Menard 2001	
D12S67*	Tetra	135,177-193**	159-185***	58	n.s.	-0.0262	F: GCAACAGTTTATGCTAAAGC R: TGTTGTTCAAGGGTCAAATG	Nürnberg et al., 1998	
D13S765*	Tetra	220,224,232**	137-165***	58	n.s.	-0.0512	F: TGTAACTTACTTCAAATGGCTCA R: ATTTACCTAACATTTCACCCATC	Zhang et al. 2001	
D14S255*	Di	173-185**	91-113***	60	n.s.	-0.0142	F: AGCTTCCAATACCTCACCAA R: CTCTTAGTGGTCATTCTCAC	Nürnberg et al., 1998	
D18S536	Tetra	144-152	144-164	58	n.s.	-0.0491	F: ATTATCACTGGTGTTAGTCCT R: CACAGTTGTGTGAGCCAGT	Kümmerli and Martin 2005	

^an.s.=no significant deviation

*primers of this marker were modified to be specific to crested macaques

**before primer modification

***after primer modification

Table 2: Number of alleles, observed and expected heterozygosity, polymorphic information content and inbreeding coefficient for twelve selected markers overall (all) and per group (R1, R2, PB), with the mean and standard deviation (SD) across all markers. The analysis is based on 176 crested macaques from three groups in the Tangkoko population in North Sulawesi, Indonesia

	Nur	Number of alleles Observed heterozygosity					Ехр	Expected heterozygosity Polymorphic information conten			content	Inbreeding coefficient								
Locus	all	R1	R2	РВ	all	R1	R2	РВ	all	R1	R2	РВ	all	R1	R2	РВ	all	R1	R2	РВ
D1s548	6	5	6	5	0.784	0.726	0.833	0.881	0.736	0.726	0.765	0.736	0.697	0.681	0.725	0.690	-0.065	0.000	-0.090	-0.199
D3s1768	7	7	6	6	0.851	0.855	0.881	0.833	0.781	0.757	0.776	0.768	0.744	0.713	0.734	0.721	-0.089	-0.131	-0.137	-0.086
D5s1457	6	5	5	5	0.727	0.714	0.717	0.714	0.649	0.674	0.645	0.609	0.589	0.613	0.581	0.541	-0.121	-0.060	-0.112	-0.175
D6s493	5	4	5	3	0.688	0.683	0.627	0.780	0.643	0.648	0.614	0.658	0.579	0.580	0.553	0.577	-0.070	-0.054	-0.021	-0.190
D6s501	5	4	5	4	0.727	0.679	0.783	0.714	0.682	0.675	0.692	0.669	0.614	0.602	0.621	0.598	-0.067	-0.006	-0.133	-0.068
D7s2204	6	6	6	6	0.805	0.831	0.817	0.756	0.724	0.727	0.69	0.721	0.674	0.673	0.633	0.668	-0.112	-0.144	-0.185	-0.049
D10s1432	4	4	4	4	0.710	0.690	0.833	0.548	0.613	0.615	0.628	0.567	0.538	0.542	0.545	0.476	-0.159	-0.124	-0.332	0.035
D11s925	9	9	8	9	0.792	0.805	0.746	0.810	0.748	0.754	0.731	0.758	0.725	0.731	0.701	0.714	-0.059	-0.068	-0.020	-0.069
D12s67	9	9	8	7	0.856	0.869	0.879	0.762	0.818	0.825	0.779	0.806	0.790	0.796	0.735	0.768	-0.047	-0.054	-0.130	0.055
D13s765	7	7	7	6	0.795	0.762	0.800	0.810	0.727	0.691	0.703	0.762	0.693	0.655	0.656	0.713	-0.095	-0.104	-0.140	-0.063
D14s255	3	3	3	3	0.665	0.774	0.550	0.619	0.651	0.669	0.601	0.626	0.575	0.591	0.529	0.537	-0.021	-0.158	0.085	0.011
D18s536	6	6	5	5	0.787	0.771	0.767	0.805	0.723	0.711	0.705	0.702	0.672	0.655	0.651	0.635	-0.089	-0.085	-0.089	-0.149
Mean	6.1	5.8	5.7	5.3	0.766	0.763	0.769	0.753	0.708	0.706	0.694	0.699	0.658	0.653	0.639	0.637	-0.082	-0.082	-0.109	-0.079
SD	1.7	1.9	1.4	1.6	0.059	0.064	0.095	0.089	0.059	0.054	0.059	0.070	0.075	0.069	0.072	0.087	0.035	0.049	0.097	0.082

Table 3: Degree of male reproductive skew in three groups of crested macaques at Tangkoko Reserve, Indonesia, 2007-2009. We provide the number of potential group sires, number of group sires, number of adult females, number of determined paternities, proportion of alpha-male paternity, proportion of alpha-male tenure across the year, the observed B value, the lower and upper confidence interval (each 0.95%) together with the P value that the observed B value is due to chance (significant values in bold). The B index incorporates male residency in days per group and year. This analysis includes a total of 51 offspring.

Group and year	Number of potential group sires	Number of group sires	Number of adult females	Number of determined paternities	Proportion of alpha- male paternity [%]	Proportion of alpha- male tenure across the year [%]	Observed B index	P level	Lower confidence interval	Upper confidence interval
R1 2007	15	4	20	9	55.56	73.15	0.179	0.001	0.033	0.455
R1 2008	20	2	21	3	33.33	73.42	0.139	0.165	-0.303	0.562
R1 2009	21	5	25	7	28.57	97.26	0.021	0.250	-0.133	0.289
R2 2007	14	3	18	9	77.78	18.38	0.527	0.000	0.192	0.865
R2 2008	7	1	19	7	100.00	100.00	0.672	0.000	0.214	0.672
R2 2009	10	1	20	9	100.00	100.00	0.621	0.000	0.251	0.621
PB 2009	16	3	17	7	57.14	31.51	0.153	0.016	0.016	0.506
Mean	14.7	2.7	20.0	7.3	64.63	70.53	0.330			
SD	5.0	1.5	2.6	2.1	29.14	33.44	0.267			