Genetic diversity within the Japanese badgers (*Meles anakuma*), as revealed by microsatellite analysis

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Abstract. To further understand the population structures of the Japanese badgers (*Meles anakuma*) on the Japanese islands, we analyzed their bi-parentally inherited microsatellites. Based on genotypes of nine microsatellite loci, the badgers were divided into five discrete clusters: three clusters from the Honshu Island, one from Kyushu and one from Shikoku. We propose that this genetic differentiation among badgers from the Honshu, Shikoku and Kyushu Islands is as a consequence of geographical isolation caused by the Seto Inland Sea. Furthermore, the cluster containing individuals from Shikoku was more differentiated from the other clusters, plausibly attributable to the earlier geological separation of the Shikoku Island from the Honshu and Kyushu Islands. The three clusters in Honshu, however, did not correspond precisely with geographical locations. As indicated in previous studies, based on mitochondrial DNA analysis, the genetic relationships within the Japanese badgers might reflect recent population expansion, occurring over a relatively short evolutionary time-scale. The findings preliminarily indicate that the Japanese badgers do not possess the high levels of philopatry seen in the European badger (*Meles meles*), a closely related species, although further analyses using balanced sample sizes from a wider range is required.

Key words: diversity, geographic barrier, Japanese badger, *Meles anakuma*, microsatellite.

The Japanese badger (*Meles anakuma*) is a medium-sized mustelid endemic to Japanese islands. *Meles anakuma* was thought to be a subspecies of the Eurasian badger (*M. meles*); this former species extending across the Palaearctic, from the Japanese islands to the United Kingdom and Ireland through the mid-latitude Eurasian Continent. Three subspecies groups were recognized: *M. m. meles*, *M. m. arenarius-leptorhyncus* and *M. m. amurensis-anakuma* (Heptner et al. 1967). Recently, however, these subspecies have been promoted to independent species status, the European badger (*M. meles*), the Asian badger (*M. leucurus*), and the Japanese badger (*M. anakuma*), based on morphological features such as mask coloration (Abramov 2003), baculum structure (Abramov 2002) and unique parasitic fleas (Abramov and Medvedev 2003). Moreover, phylogenetic analyses of mitochondrial DNA (mtDNA) (Kurose et al. 2001; Marmi et al. 2006) and nuclear DNA (Sato et al. 2003) have demonstrated the large differentiation between Japanese and continental badgers. Kurose et al. (2001) reported the low mtDNA cytochrome *b* genetic diversity within Japanese badgers, and found no relationships between genetic distances and the geographic distances between sampling locations.

Japanese badgers are distributed in the Honshu, Shikoku and Kyushu Islands, inhabiting various environments such as deciduous larch plantation, deciduous forests, cedar/cypress plantation, agriculture land, *inter alia* (see Kaneko 2009). The badger is currently treated as a game species in Japan although the numbers hunted

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per year have declined from 7,000 individuals in the
1970s to less than 2,000 in the late 1980s, Kaneko (2009)
proposing this decline to be as a result of loss of interest
in badgers as game animals, rather than intrinsic popula-
tion decline. Although this species is listed as ‘least
concern’ by the Japanese Ministry of the Environment, a
7% reduction in its distribution range occurred between
1978 and 2003 (Ministry of Environment, Japan 2004)
and 11 out of 46 prefectures list the Japanese badger in
the red data book (Kaneko 2009). Further survey work is
imperative to assess the species’ national status in Japan.

To further understand the genetic population struc-
tures of the Japanese badgers, here we investigated
variations between badgers from various regions in
biparentally inherited and allele frequencies. These
 genetic data are then used to construct a framework for
differentiation within the Japanese badgers.

Materials and methods

Samples and DNA extraction

Tissue samples, hairs and bloods were obtained from
85 Japanese badgers, which were roadkilled, hunted, and
captured at ecological studies between 1990 and 2009.
Muscle, liver and skin tissues were preserved in ethanol,
hairs were dried at room temperature, and bloods were
frozen at –80°C until use. Sampling locations and num-
bers are shown in Figure 1. DNA extraction was per-
formed by using QiAamp DNA Micro Kit (Qiagen) for

hair roots and DNeasy Blood & Tissue Kit (Qiagen) for
the other tissues. Extracted DNA samples were dis-
solved in 200 µl of 10 mM Tris/1 mM EDTA/pH 8.0
(TE) buffer, and preserved at 4°C.

Microsatellites analysis

Nine polymorphic microsatellite loci were amplified
with the suite of primers previously reported by Carpenter
et al. (2003) (Mel101, Mel102, Mel104-Mel110). An
aliquot (10 µl) of polymerase chain reaction (PCR) solu-
tion was comprised by 1 µl of 10 × PCR buffer, 0.8 µl of
2.5 mM dNTP mixture, 0.1 µl of TaqDNA polymerase
(5 units/µl, Takara), 0.3 µl of above each primer and 1 µl
of extracted DNA, and 6.8 µl of distilled water. The
PCR amplifications were carried out in a PCR thermal
cycler (Takara TP600), one cycle of 94°C for 3 min and
35 cycles of 94°C for 15 sec; 54°C for 20 sec; 72°C for
30 sec; followed by one cycle of 72°C for 10 min. These
PCR products were then processed in an automated DNA
sequencer (Hitachi SQ5500), and resultant data were
analyzed using FRAGLYS v. 2.0 software (Hitachi).

Observed \(H_O\) and expected \(H_E\) heterozygosities,
mean numbers of alleles per locus and value of pairwise
\(F_{ST}\) were calculated using ARLEQUIN 3.1 software
(Excoffier et al. 2005). Departures from Hardy-Weinberg
equilibrium and linkage equilibrium were tested for each
of the nine loci using GENEPOP 3.4 software (Raymond
and Rousset 1995) applying the Markov chain method
according to the algorithm of Guo and Thompson (1992).
Based on values of pairwise \(F_{ST}\), a neighbor-joining tree
(Saitou and Nei 1987) was constructed using MEGA 4
software (Tamura et al. 2007). The most parsimonious
number of populations was estimated using STRUC-
TURE 2.2 software (Pritchard et al. 2000). This program
characterizes each population by allele frequencies at
each locus. Ten replicates were performed in order to
establish each value of \(K\), where \(K\) represents the optimal
number of genetic clusters (i.e. populations), testing a
range of \(K\) from 3 to 10 (with 100,000 burnin and
100,000 Markov Chain Monte Carlo replicates after bur-
nin). Individuals with greater than a 50% assignment
rate in favor of one population were assumed to belong
to that population. In addition, Nei’s standard genetic
distance \(D_S\) (Nei 1978) was calculated for each sub-
population using SPAGeDi 1.2 software (Hardy and
Vekemans 2002) and the phylogenetic tree based on the
neighbor-joining method (Saitou and Nei 1987) was
constructed based on \(D_S\) by using MEGA 4 software.
Results

Genetic diversity of Japanese badgers

Nine microsatellite loci in 85 Japanese badgers were genotyped. As only a single individual was genotyped in each of Iwate and Yamaguchi Prefectures, these data were excluded from heterozygosity calculation in Table 1. The Kanto population was comprised by individuals from Tokyo ($n = 29$), Chiba ($n = 1$) and Gunma ($n = 17$). Shikoku consists of individuals from Kochi ($n = 17$). Kyushu consists of individuals from Fukuoka ($n = 4$), Nagasaki ($n = 1$), Oita ($n = 9$) and Kumamoto ($n = 1$).

The number of alleles measured ranged from three (at Mel108) to 24 (at Mel105). The mean number of alleles per locus over all populations was 4.0 and the highest number was 7.9, from the Kanto population. Observed heterozygosity ($H_O$) ranged from 0.389 to 0.484, and the expected heterozygosity ($H_E$) ranged from 0.529 to 0.638 (Table 1). The value (1.4) of alleles per locus for both single individuals from Iwate and Yamaguchi Prefectures (Table 1) indicated multiple loci in each of the two badgers were homozygous.

The neighbor-joining tree based on $F_{ST}$ values showed that the Shikoku population is more distant from the other populations (Fig. 2). The highest value of pairwise $F_{ST}$ (0.235) occurred between the Shikoku and Kyushu populations. The lowest (0.104) occurred between Kanto and Gifu populations.

Population structures of Japanese badgers

The STRUCTURE analysis showed that the highest log-likelihood value was found at $K = 5$, consequently the most optimal number of populations of the Japanese badgers obtained in the present study was established to be five, referenced as clusters I–V. Of the 85 individuals genotyped, 81 were assigned to either of these five clusters with >50% assignment rates. Bayesian clusterings approximated with the populations which we defined on the basis of regions. Out of the five clusters, three clusters I, II and III were shared by the majority of individuals from the Kanto population. The Shikoku and Kyushu populations were assigned to cluster IV and V with >91% and >85% assignment rates, respectively (Table 2). Three individuals from Gifu, located in central Honshu, were assigned to cluster IV or V, and not to any of clusters I, II and III distributed in Kanto. The neighbor-joining tree based on $D_S$ values showed a broader differentiation of cluster IV from the other clusters (Fig. 3).

Discussion

Genetic relationships among the Japanese badgers

Regional populations defined by sampling locations corresponded well with genetic populations inferred by microsatellite genotyping (Table 2). Bayesian clusterings, established by STRUCTURE analysis, revealed that the Shikoku and Kyushu populations were separated from each other and from most individuals from the Honshu Island: that is, cluster IV consisted of all 17 individuals from Shikoku and one from Gifu, and cluster V consisted of all 15 individuals from Kyushu and two from Gifu (Table 2).

Tsuchiya et al. (2000) reported that the Shikoku and Kyushu populations of the large Japanese mole (Mogera wogura) are also separated from one another, as well as from Honshu populations, based on mtDNA cytochrome
The Seto Inland Sea (see Fig. 1), which presently separates the Honshu, Shikoku and Kyushu Islands, formed about 5,000 years ago (Ohshima 1990). Consequently it is highly plausible that the differentiation of the Shikoku and Kyushu populations from the Kanto and Gifu populations in the Honshu Island might have arisen as a result of the geographical isolation caused by the formation of the Seto Inland Sea. For the Japanese sika deer (*Cervus nippon*), however, the Shikoku Island is a sympatric region of the northern and southern mtDNA lineages (Yamada et al. 2006). In the present study, the individuals from Gifu at central Honshu were included in the clusters of Shikoku and Kyushu. Further investigation is required to examine the full ramifications of the ontogeny of the Seto Inland Sea on the extended population structures of the Japanese badgers, using more samples from central and western Honshu.

The present study demonstrates that the Shikoku population is the most differentiated (Fig. 2). The genetic separation of the Shikoku population could have been affected by formation of the Seto Inland Sea. During the formation process, the Shikoku and Honshu Islands first became separated about 7,000 years ago, with the Kyushu Island becoming separated from Honshu around about 5,000 years ago (Ohshima 1990). The earlier separation of Shikoku from the other main islands and consequent isolation, would support our observation of Shikoku having the most genetically differentiated population.

### Table 2. Assignment of the Japanese badgers to five clusters

<table>
<thead>
<tr>
<th>Clusters*</th>
<th>Iwate</th>
<th>Kanto</th>
<th>Gifu</th>
<th>Yamaguchi</th>
<th>Shikoku</th>
<th>Kyushu</th>
</tr>
</thead>
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<tr>
<td>I</td>
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<td>1</td>
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<td></td>
</tr>
<tr>
<td>II</td>
<td>9</td>
<td>3</td>
<td>1</td>
<td></td>
<td></td>
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<tr>
<td>VI</td>
<td></td>
<td></td>
<td>4</td>
<td>1</td>
<td>9</td>
<td>1</td>
</tr>
<tr>
<td>Non-assigned individuals</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

* Defined by STRUCTURE test using genotypes.

**Genetic structures within the Kanto and Gifu populations**

For the badgers from the Honshu Island, the Kanto populations consisted of 29 individuals from Tokyo, one from Chiba and 17 from Gunma Prefectures; the Gifu population consisted of four from Gifu Prefecture; and one badger from Iwate Prefecture and one badger from Yamaguchi Prefecture were examined. The STRUCTURE analysis showed that the badgers from the Honshu Island were divided into three genetic clusters. Most animals grouped within each of the three genetic clusters in Honshu were distributed in geographically proximate areas. For example, cluster III consisted of 12 individuals from Gunma Prefecture, but it also included three individuals from Tokyo. This genetic population structure could be explained by the recent population expansion. Based on the mtDNA analysis, Kurose et al. (2001) reported that the current distribution of the Japanese...
badgers had been formed through the repeated ‘increase and reduction’ of the total population size, with concomitant changes in distribution range due to the palaeoenvironmental changes, such as glacial-interglacial episodes.

In addition, Kurose et al. (2001) also suggested the possibility that the recent expansion of the Japanese badgers may have resulted in their unestablished geographical structures. In accord with the mtDNA data, the present study of microsatellites also did not identify any explicit regional genetic structure in the Japanese badgers of Honshu. By contrast, microsatellite analysis of the European badger (Meles meles) (Pope et al. 2006, 2007), a species closely related to the Japanese badger, established a clear population structure, which has been attributed to the high level of natal philopatry observed in study areas in the British Isles (Kruuk and Parish 1982; Cheeseman et al. 1987; Macdonald et al. 2008). If the Japanese badger were to observe strict natal philopatry, then resultant genetic population structures should be concordant with those described for the European badger (Pope et al. 2006, 2007): this, however, was not consistent with our findings among the Japanese badgers of Honshu, which did not show clear genetic structures. This indicates that the level of the natal philopatry of the Japanese badger may not be as high as that of the European badger. Moreover, in contrast to the European badger, the ecological traits of the Japanese badgers are not well known (Yamamoto 1986, 1991; Kaneko 2001, 2002; Tanaka et al. 2002). For another reason, the small number of sampling points and unbalanced sample sizes in the present study might have partly led to the resultant unestablished geographical structures of the Japanese badgers. To further understand the evolutionary ecology, breeding systems and life-history traits of the Japanese badgers, not only will genetic analyses using samples collected from a wider range be required, but also further ecological research into fundamental biology.

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