- The role of Japanese macaques (*Macaca fuscata*) as endozoochorous seed dispersers on Kinkazan
  Island, northern Japan
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12 Abstract

13 We studied the characteristics of seeds within faeces, an important aspect of endozoochorous 14seed dispersal, in Japanese macaques Macaca fuscata inhabiting Kinkazan Island, northern Japan. 15We intermittently collected faecal samples from 1999 to 2009 (N = 1294) and examined the rate of 16seed occurrence, species/life-form composition, number of seeds, and their intact rate . Seeds were 17found within faecal samples during every month, but their characteristics changed monthly: the rate 18of seed occurrence and the number of plant species within faecal samples were greater in summer 19and fall, and the intact rate and number of intact seeds observed within single faecal samples were 20also higher during these seasons than spring and winter. These results suggest that Japanese 21macaques on Kinkazan act as seed dispersers in summer and fall and that they disperse intact seeds 22into wider areas within the forest through defecation. During the study period, we observed seeds 23from a total of 35 plant species from 22 families in our samples. In addition to those of woody plants, 24we also observed seeds from as many as 13 herbaceous plants, for which sympatric sika deer 25(Cervus nippon) have historically been considered the sole seed dispersal agents. The intact rate of 26seeds was significantly negatively correlated with the seeds' mean cubic diameter, and this 27relationship was strengthened for non-fleshy fruits. We also conducted regional comparisons of the 28characteristics of defecated seeds in order to address whether regional variations in the diet of the 29macaques affect their efficacy as seed dispersers, both in terms of quantity and quality. Macaques 30 living in the natural habitats of Kinkazan and Yakushima dispersed more seeds of tall tree species 31than do macaques inhabiting the human-altered areas of in Shimokita and Kashima. The number of 32plant species represented within single faecal samples also varied geographically, being greater in 33Yakushima. This pattern likely resulted from Yakushima's warm-temperate climate, as the other 34 three study sites occur in the cool-temperate region. Our results suggest that the composition of 35seeds dispersed by Japanese macaques are not rigid, but are determined instead by the vegetation

36 found in a given habitat.

37 Keywords: Japanese macaque; intact rate; Kinkazan; seed dispersal; temperate region

38

39 Introduction

40Environmental and climatic conditions affect food availability, and can have an impact on 41 dietary composition/diversity of given animals (Hill and Dunbar 2002). Studies aimed at comparing 42the feeding habits of a single mammal species across different geographical regions have revealed 43that fruit diversity in the diets of various mammals living at lower latitudes is greater than at higher 44 latitudes, mainly because of differences in food availability and/or physical conditions (Virgòs et al. 451999; Zalewski 2004; Lozano et al. 2006; Zhou et al. in press). Investigating regional variation in the 46 food habits of a particular species is therefore important for understanding its ecological plasticity, 47especially when considering wide-ranging mammals living in environments where the type and 48 abundance of food resources are constantly changing (Rosalino and Santos-Reis 2009). With respect 49to seed dispersal, such regional variation in diet might affect that mammal's efficacy as a seed 50disperser, both in terms of quantity (the number of seeds dispersed) and quality (the number of plant 51species, proportion of intact seeds, and germination rate), leading to variation in forest ecosystem 52structuring. For example, frugivores inhabiting lower latitudes might disperse a larger number of 53intact seeds from a more diverse array of plant species than those inhabiting higher latitudes. 54Therefore, understanding the general role of a given mammal species as a seed disperser requires a 55comprehensive approach rather than surveying on a local scale. 56Primates constitute one of the groups of frugivorous mammals inhabiting temperate regions. The 57diets of almost all cercopithecinae monkeys inhabiting temperate regions include at least some fruits (Macaca mulatta: Goldstein and Richard 1989; M. cyclopis: Su and Lee 2003; M. fuscata: Tsuji et al. 58

59 2006; *M. thibetana*: Zhao et al. 1991), and these animals can thus deposit seeds in locations far from

60	the seeds' origins. Compared to their sympatric avian seed dispersers, primates consume similar
61	amounts of fruit (Howe 1980; Son et al. 2007; Nakashima 2008), have larger gape sizes (Noma and
62	Yumoto 1997), have larger home ranges (1.3—6.4 ha for birds: Fukui 1995, $<0.2$ to $>50$ km <sup>2</sup> for
63	primates: Clutton-Brock and Harvey 1977), and exhibit longer retention times in the gut (15-30
64	minutes for birds: Murphy et al. 1993; Fukui 1996; 1.6-190 hours for primates: Lambert 2002;
65	Tsuji et al. 2010), all of which contribute to a wider dispersal of seeds. Thus, primates in temperate
66	regions are likely to strongly affect the spatial and genetic structure of plant populations and thereby
67	contribute to plant community dynamics.
68	(Fig. 1 should appear here)
69	Japanese macaques (Macaca fuscata) have the northern-most distribution of extant non-human
70	primates (30° 2'-41° 1' N). Previous studies of the role of Japanese macaques as endozoochorous
71	seed dispersers have been conducted mainly in the warm-temperate regions of Yakushima Island
72	(hereafter Yakushima). These studies have compiled lists of intact seeds observed within faeces
73	(Otani and Shibata 2000), estimated dispersal distance (Yumoto et al. 1998; Terakawa et al. 2009),
74	and examined the effects of: dispersal on seed germination/seedling growth (Yumoto et al. 1998);
75	seed/fruit characteristics on seed destruction (Otani and Shibata 2000); and, foraging patterns on the
76	topography of defecation sites (Tsujino and Yumoto 2009).
77	The habitat of Japanese macaques is roughly divided into two major climate regions: the
78	warm-temperate region dominated by evergreen forests in the southern part of their range and the
79	cool-temperate region dominated by deciduous forests in northern areas (Fig. 1). The food habits and
80	habitat utilisation (e.g., home range size, daily moving distance, and degree of frequent use) of
81	macaques differ between the two climate regions (Takasaki 1981; Agetsuma and Nakagawa 1998;
82	Maruhashi et al. 1998; reviewed by Tsuji 2010), largely because of differences in forest type. For
83	example, tall trees with fleshy fruits occur in greater abundance (Otani 2005) and density

84	(Maruhashi et al. 1998) in warm- than cool-temperate regions. Considering the wide variation of
85	habitats throughout Japan, information from Yakushima represents only a small portion of the
86	variation in seed dispersal patterns by Japanese macaques. The characteristics of seeds within faeces,
87	an important aspect of endozoochorous dispersal, are likely to be affected by differences in the
88	habitat structure and feeding behaviour of these animals. Thus, in order to gain a more general
89	understanding of endozoochorous seed dispersal among Japanese macaques, regional comparisons
90	between the cool- and warm-temperate regions are necessary. However, at present, few studies
91	(Otani 2003) have addressed seed dispersal by macaques inhabiting cool-temperate regions.
92	The objectives of this study were to document the characteristics of seeds defecated by Japanese
93	macaques inhabiting the cool-temperate Island of Kinkazan and to compare the results to those from
94	other study sites. In terms of seed characteristics, we examined seed appearance, species/life-form
95	composition, intact rate, the numbers of intact seeds and species contained within single faecal
96	samples, and monthly changes in these parameters. Because macaques on Kinkazan feed on woody
97	fruits primarily in the summer and fall (Tsuji et al. 2006), they most likely act as seed dispersers
98	primarily during these two seasons, defecating large numbers of intact seeds from various plant
99	species. In addition, we examined the relationship between seed size and their rate of destruction. On
100	Yakushima, it is known that relatively larger seeds or seeds of non-fleshy fruits tend to be destroyed
101	more often than smaller ones (Otani and Shibata 2000). We therefore expected that differences in
102	forest type and consequent differences in plant species composition between the cool- and
103	warm-temperate zones would lead to differences in this relationship.
104	
105	Materials and methods
106	Study site and subject animals

107 Kinkazan (38°2'N, 141°4'E) is located 700 m from the Oshika Peninsula, northern Japan (Fig. 1).

108	The total area of the island is ca. $9.6 \text{ km}^2$ , and the highest peak is 450 m a.s.l. The monthly mean air
109	temperature on the island ranges from 2.5°C in February to 22.3°C in August. Based on the climate
110	conditions of the island, the year was divided into four seasons: spring (March-May), summer
111	(June-August), fall (September-November), and winter (December-February). On the island,
112	deciduous forests of Fagus crenata dominate the higher elevations (>150 m), whereas a mixture of
113	deciduous forests of Carpinus spp. (C. tschonoskii and C. laxiflora) and coniferous forests of Abies
114	firma cover the lower elevations (<150 m). Kinkazan also harbours a forest of planted Cryptomeria
115	japonica and patchy grassland communities of Zoysia japonica (Yoshii and Yoshioka 1949), the
116	latter of which has expanded due to heavy grazing and seed dispersal by sympatric sika deer (Cervus
117	nippon) (Takatsuki 2009). Approximately 200-250 Japanese macaques belonging to six troops (A, B <sub>1</sub> ,
118	B <sub>2</sub> , C <sub>1</sub> , C <sub>2</sub> , and D) inhabit the island (Izawa 2009).
119	
120	Observation of the macaques
121	Observations of the monkeys of Troop A were conducted from June 2004 to May 2005

122(88 days) (Tsuji et al. 2008). We recorded the foraging data of focal adult females (N = 14-17) with 1231-min instantaneous sampling methods. Observations focused on foraging and handling techniques 124for fruit and seeds (Otani and Shibata 2000). Fruit species eaten by focal animals were classified into 125three categories, depending on the macaques' handling techniques for seeds: swallowed, crunched or 126discarded. Species classified as swallowed were those for which whole fruits were eaten and seeds 127were ingested and then passed in the faeces. For species classified as discarded, although whole 128fruits were eaten, only the flesh was ingested while the seeds were spat out. Crunched seeds were 129those that were crushed during mastication, although the processing technique for the flesh of the 130 fruit varied by species. We could not identify a number of herbaceous plants (especially those of the 131family Gramineae) due to quick intake by the macaques.

132

133	Faecal sample collection and treatment
134	Fresh faecal samples were intermittently and opportunistically collected from the ground from
135	each troop between March 1999 and March 2009 (during 49 surveys in total), though we focused our
136	collection mainly within the home range of Troop A. Each faecal sample was placed into a plastic
137	bag and stored in a freezer at our field station. In the laboratory, each sample was thoroughly mixed
138	with water and rinsed through 0.5-mm sieves under fresh water. All contents were then placed into
139	plastic bottles with 70% alcohol and kept at room temperature until faecal analyses.
140	
141	Analyses of faecal samples and regional comparisons
142	We removed all seeds from the bottled faecal samples and identified them to the species level.
143	Seed identification was based on a reference collection established in previous years as well as on
144	Nakayama et al. (2000). We classified the identified species into five life forms: tall trees (>5 m in
145	height), shrubs (<5 m), vines, hemi-parasites, and herbaceous plants. In addition, seeds were
146	classified into two fruit types: fleshy- and non-fleshy fruits.
147	We measured the longest axis $(a_1)$ , second longest axis $(a_2)$ , and third longest axis $(a_3)$ of 10
148	randomly selected seeds for each plant species to the nearest 0.05 mm using vernier callipers. For
149	Torreya nucifera, we only measured one seed for each dimension due to a dearth of seeds in our
150	samples. From averaged values, we calculated the mean cubic diameter (MCD) of seeds (Otani and
151	Shibata 2000): $MCD = (a_1 \times a_2 \times a_3)^{1/3}$ . After taking these measurements, we counted the total
152	number of seeds and the number of "intact seeds", defined as those seeds that appeared to be
153	complete, within each bottled sample. For plant species whose seeds were relatively large ( $MCD \ge 1$
154	mm), we directly counted the numbers of both the total and intact seeds within the entire bottled
155	sample. Direct counts were difficult for smaller seeds ( $MCD < 1 \text{ mm}$ ), and we therefore sub-sampled

156 ca. 1 g (dry weight) of the bottle contents and counted the numbers of both whole and intact seeds

157 within the sub-sample. From this sub-sample, the numbers of both whole and intact seeds within the

158 entire bottled sample were estimated.

159The evaluation of seed characteristics followed Kunz et al. (2008). We first calculated the 160number of dispersal events (DE), which was defined as the number of faecal samples containing 161 seeds, for each month and for each plant species. Because plant species have different fruiting 162periods and because we did not collect faecal samples in equal quantities across all months of the 163 study period (range: 10-374), we also calculated DE ratios ( $DE_r$ ), obtained by dividing DE by the 164number of faecal samples examined, for each month and for each plant species. To determine the 165primary month(s) of DE for each species, we compared the number of intact seeds in a given month 166to the seed's annual mean. Months during which the former was greater than the latter were defined 167as main seed dispersal months. From this calculation, we obtained the primary seed dispersal 168season(s). We also calculated an intact rate (IR) as a qualitative index of the efficacy of seed 169dispersal (Koike et al. 2008) for each month and for each plant species. The IR was obtained by 170dividing the total number of intact seeds in a given month by the total number of seeds in that 171corresponding month, and the efficacy index was obtained by dividing the total number of intact 172seeds of a given species during the entire study period by the total number of seeds of a given 173species. Finally, we compared the  $DE_r$ , life-form composition, percentage of fleshy fruits, MCD of 174seeds, and mean number of plant species within single faecal samples from Kinkazan to values from 175three other study sites: Yakushima (Otani 2005), Shimokita (Otani 2003), and Kashima (Otani 2003). 176 177Statistical analysis

To examine monthly changes in values of  $DE_r$  and IR, we employed binomial tests to compare the observed  $DE_r$  (or IR) in a given month to the expected  $DE_r$  (or IR) (obtained from the annual

180	mean). To determine monthly changes in the total number of intact seeds, the number of intact seeds
181	for each plant species, the number of species within single faecal samples, and to test the regional
182	variation in the MCD across study sites, we used non-parametric Kruskal-Wallis analyses of
183	variance (ANOVAs). Finally, to test the effects of the MCD, fruit type, and their interaction on IR,
184	we used a generalized linear model (GLM). We treated the sum of intact seeds collected from whole
185	faecal samples for each plant species as a dependent variable. The sum of all seeds of corresponding
186	plant species collected from whole faecal samples was used as an offset term. We assumed the
187	negative binomial distribution as an error structure. All data analyses were carried out using the
188	statistical software R. 2.9.1 (R Development Core Team 2009). For all analyses, significance levels
189	( $\alpha$ ) were set at 0.05. We did not conduct statistical analyses for the regional comparisons, except for
190	the MCD, because of small sample sizes at the other study sites.
191	(Table 1 should appear here)
192	Results
193	Observation of feeding behaviour
194	Fruit foraging was observed for a total of 659 h, during which the macaques foraged on the
195	fruits of 29 woody plant species (17 families) and at least 8 herbaceous plant species (6 families)
196	(Tsuji and Takatsuki in prep.). Consumed fruits were composed of 28 fleshy fruits and 9 non-fleshy
197	fruits.
198	
199	Monthly changes in seed occurrence and dispersal
200	A total of 1294 faecal samples were collected during 49 surveys (Table 1). The numbers of
201	faecal samples collected in April ( $N = 10$ ) and February ( $N = 19$ ) were lower than those for other
202	months, which may have affected the following results. The annual mean of the $DE_r$ was 0.58
203	(747/1294), and more than 66000 seeds were collected from faeces (Table 1). Compared to the

204	annual mean, values of $DE_r$ from June to December were significantly higher (binomial tests, $P <$
205	0.05), whereas the $DE_r$ s from February to May were significantly lower ( $P < 0.05$ ). In January, the
206	$DE_r$ did not significantly differ from the annual mean ( $P > 0.05$ ; Table 1). The annual mean of the
207	IR was 0.78 (66248/85042) (Table 1). Compared to the annual mean, the IR was significantly lower
208	in March (binomial tests, $P < 0.05$ ), but significantly greater in June and from November to January
209	(P < 0.05). In other months, the <i>IR</i> did not significantly differ from the annual mean $(P > 0.05)$ ;
210	Table 1).
211	

212 Monthly changes in seed species composition

213Table 2 shows the morphological characteristics and the handling techniques used for 36 214different plant species extracted from the faecal samples, from which 35 species from 22 families 215were identified (Table 2). These species included 11 tall trees, 5 shrubs, 5 vines, 1 hemi-parasite, and 21613 herbaceous plants. Seeds of shrubs were found during the entire year except for April. Seeds of 217herbaceous plants were found within faeces collected from May to December. Seeds of the 218hemi-parasite (Viscum album) were found in March and from November to February. Seeds of vines 219were found from June to January (Fig. 2). Identified plants included 20 fleshy-fruit and 15 220non-fleshy-fruit species (Table 2). The total number of accumulated plant species was highest in 221October and November (18 plant species in each month) and lowest in February (2 species each) and 222April (none) (Fig. 2). 223The seeds detected from the faecal samples contained many plant species whose seeds were 224swallowed by the macaques during our feeding observations, many of which being fleshy fruits (14/ 22520 species). We also detected the seeds of many plant species whose seeds were crunched by the 226macaques, about half of which were non-fleshy fruits (7/15 species). Furthermore, seeds of two

227 species (Cornus kousa and Diospyros lotus) that were discarded by the macaques during

228 observations were detected in faecal samples (Table 2).

229	Of the 35 identified	plant species.	six (Vibrunum	dilatum, Ros	sa multiflora.	Vitis flexuosa	Malus
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- 230 tchonoskii, Berchemia racemosa, and Oxalis corniculata) exhibited significant monthly variation in
- their number of intact seeds (Kruskal–Wallis tests, P < 0.05), and two (*Pourthiaea villosa* and
- 232 *Cornus kousa*) showed marginally significant monthly changes (P < 0.10). We observed seasonal
- 233 variation in the DE among species, and the number of intact seeds exhibited peaks during two main
- seasons: summer (14 species) and fall (18 species) (Table 2).
- 235 (Table 2 and Fig. 2 should appear here)
- 236 Number of intact seeds and seed diversity within single faecal samples
- The mean ( $\pm$  SD) total number of intact seeds within single faecal samples was 58  $\pm$  176, and

this value exhibited significant monthly variation (Kruskal–Wallis test,  $\chi^2 = 759.4$ , df = 10, P < 10

239 0.001) (Table 3). The number of intact seeds was greatest in June and lowest in February, March, and

240 May. Of the 747 faecal samples that contained intact seeds, the mean seed diversity (i.e. number of

- plant species represented) was  $2.0 \pm 0.9$  (range:1-7) (Table 3). The mean seed diversity showed
- significant monthly variation as well (Kruskal–Wallis test,  $\chi^2 = 272.3$ , df = 10, P < 0.001), and was
- highest in June (4.1  $\pm$  1.7) and lowest in May (1.5  $\pm$  1.3), July (1.3  $\pm$  0.6), and February (1.3  $\pm$  0.5)
- 244 (Table 3).
- 245 (Table 3 should appear here)
- 246 Relationship between the intact rate and seed size/fruit type
- 247 The *MCD* of dispersed seeds was  $3.0 \pm 2.0$  mm (range: 0.8–10.0) (Table 2). *IR* was
- significantly affected by both the MCD (GLM: z = -4.32, P < 0.001) and by an interaction between
- fruit type and the MCD (z = 2.39, P = 0.017) in which the effect of the MCD on the IR was
- strengthened for non-fleshy fruits (Fig. 3).
- 251 (Fig. 3 should appear here)

252 Regional comparisons

253A regional comparison revealed that the  $DE_r$  for Kinkazan (0.58) was lower than that for Yakushima (0.99), Shimokita (0.93), and Kashima (0.79) (Table 4). The life-form composition of 254255plant species represented in faecal samples varied among the study sites such that, on Kinkazan, the 256percentage of tall trees (31%) was similar to that on Yakushima (38%), and greater than the other 257two sites (8% and 17%, respectively). The percentages of shrubs (17%) and vines (14%) represented 258in samples from Kinkazan were lower than those from other study sites, whereas the percentage of 259herbaceous plants (33%) was greater than that from other sites (Table 4). The percentage of fleshy 260fruits observed in our Kinkazan samples (56%) was lower than those from the other three study sites 261(Yakushima: 91%, Shimokita: 92%, Kashima: 89%). The MCD of seeds did not significantly vary 262among the four study sites (Kruskal-Wallis test,  $\chi 2 = 1.93$ , df = 3, P = 0.586) (Table 4). In contrast, 263although we were not able to test for the significance of this pattern, the mean seed diversity per 264faecal sample was greatest on Yakushima  $(2.8 \pm 1.4)$ , followed by Kashima  $(2.4 \pm 1.9)$  and Kinkazan 265 $(2.0 \pm 0.9)$ , and was lowest on Shimokita  $(1.9 \pm 0.8)$ . 266(Table 4 should appear here) 267Discussion 268Monthly changes in seed characteristics within faeces 269In summer and fall (strictly from June to December), values of  $DE_r$  and IR were greater than 270their annual means (Table 1), the seed diversity was high (Fig. 2), and the number of intact seeds 271peaked for many plant species (Table 2). Furthermore, the numbers of intact seeds and seed diversity 272were greater in summer and fall than spring and winter (Table 3). Thus, as we predicted, the role of 273Japanese macaques as seed dispersers on Kinkazan appeared to be greater in summer and fall, during

which macaques dispersed intact seeds of various plant species more frequently and into wider areas

within the forest than during winter and spring.

276	The observed seasonal variation in seed dispersal can be attributed to seasonal changes in the
277	feeding behaviour of macaques on Kinkazan: in summer and fall, these animals spend a great deal of
278	time feeding on various kinds of fruits (ca. 20-60%) (Tsuji et al. 2006). The macaques ingest large
279	amounts of fruit to accumulate body fat (Tsuji et al. 2008) and may incidentally swallow the entire
280	fruit without adequately chewing it. In contrast, when fruit is less available during spring and winter,
281	macaques opportunistically feed on fruits remaining on the ground and/or on tree branches
282	(Nakagawa 1989; Tsuji et al. 2006), and may thus take the time to chew the contents well before
283	swallowing.
284	
285	Plant species dispersed by macaques on Kinkazan
286	We found that macaques on Kinkazan defecate intact seeds of 35 species from 22 families,
287	corresponding to 50.7% of all species of fruits on which this population is known to feed (69 species,
288	Izawa 2009). We detected seeds of more than half of all woody plants observed to be ingested during
289	behavioural observations (22 out of 29 species). Interestingly, the macaques also dispersed the seeds
290	of 13 herbaceous plants (Table 2). Before the present study, baboons (Papio spp.) were the only
291	known primate species to disperse seeds of herbaceous plants (Slater and du Toit 2002; Kunz et al.
292	2008). On Kinkazan, sympatric sika deer (Cervus nippon) are typically considered the primary
293	dispersers of herbaceous seeds (Takatsuki 2009). Because the biomass of the macaques on Kinkazan
294	(ca. 2500 kg as estimated by multiplying mean body weight (ca. 10 kg) by population size (ca. 250);
295	Izawa 2009) is much lower than that of deer (ca. 30000 kg; Takatsuki 2009), the amount of seeds
296	dispersed by macaques in each instance would likely be less than that by deer. However, the home
297	range size of macaques (ca. 3 km <sup>2</sup> ; Tsuji and Takatsuki 2009) is much larger than that of deer (<20
298	ha; Ito and Takatsuki 2009), and therefore, macaques would be expected to disperse seeds into wider
299	areas than would deer.

We also found that the seeds of several species whose fruits were crunched (example.g. the nuts of Fagaceae) or discarded (*Cornus kousa* and *Diospyros lotus*) by the macaques were sometimes detected in faecal samples (Table 2).However, since both the  $DE_r$  and IR of these species were lower than those of species whose seeds were mainly swallowed, the contribution of macaques to the dispersal of these seeds is likely negligible.

305

306 Relationship between seed size and intact rate

307 The vast majority of ingested seeds in our faecal samples were intact (annual mean IR = 0.78) 308 (Table 1), having a proportion much greater than values observed in captive Japanese macaques 309 (0.04 in Otani 2004; 0.36 in Tsuji et al. 2010). The variation in the intact rate of seeds between wild 310 and captive conditions can be attributed to a difference in the macaques' adherence/handling to the 311plant seeds in each environment. The IR varied among plant species, and the IR and MCD were 312negatively correlated such that larger seeds tended to be destroyed more frequently (Fig. 3), similar 313to results obtained for forest guenons (Cercopithecus spp.; Gautier-Hion 1984). Because the 314probability of seed destruction appears to be directly proportional to seed size, our results are 315unsurprising. We also detected a significant interaction between MCD and fruit type such that the 316 effects of the seed size on the IR was greater for non-fleshy fruits (Fig. 3) The reason for this is 317unclear, but we speculate that that seeds of non-fleshy fruits are not covered by a fleshy pulp, and the 318risk of a given seed being destroyed might therefore increase in proportion to its size. In any case, 319since masticatory seed mortality may affect dispersal efficiency, it is necessary to consider the 320 possible effects of fleshy pulp on seed dispersal characteristics. 321

322 Regional comparisons

323 The annual  $DE_r$  on Kinkazan was lower than values observed at the other study sites (Table 4).

324	Our larger sample size may have contributed to this difference, but other possible causes remain
325	unclear. Our study on Kinkazan and the study on Yakushima by Otani (2003), both of which were
326	conducted in natural forest environments though in different climates (cool-temperate versus
327	warm-temperate, respectively), produced similar results in the percentage with which macaques
328	ingested seeds of tall tree species (31% and 38%, respectively). These values are higher than those
329	reported for Shimokita (8%) and Kashima (17%), both of which are located in cool temperate
330	regions has and have been affected by human activity (Otani 2003). Our results reveal that the
331	composition of seeds dispersed by Japanese macaques are not rigid, but are instead likely determined
332	by the vegetation within a given habitat, as noted by Corlett and Lucas (1990). Because the size of
333	intact seeds within faecal samples was similar across the four study sites, every plant whose seeds lie
334	within the 2–3-mm threshold has the potential to be dispersed intact by macaques, regardless of its
335	life form. Therefore, macaques differentially contribute to forest structure across different habitats.
336	The mean seed diversity per faecal sample varied from 1.9 to 2.9 across the four study sites, and
337	was greater at lower latitudes (i.e. the warm temperate regions of Yakushima) than at higher latitudes
338	(i.e. the cool temperate regions of Shimokita, Kinkazan, and Kashima). In general, macaques in
339	cool temperate regions utilise their home ranges less intensively thanthose in warm temperate
340	regions (Maruhashi et al. 1998). Thus, in warm temperate regions, the fruit contents of a single
341	feeding bout at one feeding site might be packaged in one single faecal sample, resulting in a greater
342	number of plant species within individual faeces samples.
343	On Yakushima, macaques disperse plant seeds not only through defecation, but also through
344	discarding behaviour (Yumoto et al. 1998; Otani and Shibata 2000). Discarded seeds exhibited
345	relatively large <i>MCD</i> values according to that study $(7.3 \pm 4.5 \text{ mm}, \text{ range: } 2.6 - 15.3; \text{ Otani and}$
346	Shibata 2000). On Kinkazan, in contrast, we rarely observe seed discarding behaviour, except with
347	seeds of Cornus kousa (MCD: 4.5 mm) and Diospyros lotus (MCD: 7.8 mm). Because the number of

348larger-seeded plants is lower in cool temperate than warm temperate regions (Otani 2005), situations 349 during which macaques might discard larger seeds are expected to be rare in the former. 350Furthermore, , many of the larger seeds on Kinkazan were nuts, which were [always OR typically 351OR generally OR usually OR most often OR etc] crunched rather than swallowed. Considering that 352the dispersal distance attained through discarding behaviour is shorter than that that via defecation 353 (Yumoto et al. 1998), regional differences in the handling of seeds would likely influence seeds 354shadows. In general, the home range sizes of macaques in cool temperate regions are greater than 355 those in warm temperate regions (Takasaki 1981). In combination with the fact that macaques in 356cool temperate regions utilise their home range less intensively (Maruhashi et al. 1998), this 357 suggests that macaques in cool temperate regions should disperse seeds more widely and sparsely 358than macaques in warm temperate regions. Such variation in the patterns of dispersal would 359ultimately lead to differences in the forest structures of the two climate regions. To confirm these 360 predictions, future studies should collect quantitative data regarding feeding behaviour and 361defecation, conduct experiments on seed germination/growth in both cool and warm temperate 362 regions, and make regional comparisons between the two climate types. 363 In this study we demonstrated regional variation in the role of Japanese macaques as seed 364dispersers. Our results can be generalized not only to other primate species but also to other 365 mammalian seed dispersers, especially concerning wide-ranging mammals (e.g. genets (Virgòs et al. 366 1999), martens (Zalewski 2004), baboons (Hill and Dunbar 2002)) living in environments in which 367the type and abundance of food resources are constantly changing (Rosalino and Santos-Reis 2009). 368 369 Acknowledgements

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487 Figure Legends

- 488 Figure 1. Distribution of the two main vegetation types in the natural habitats of Japanese
- 489 macaques (■: deciduous forest, □: evergreen forest) (from Tsuji 2010) and locations of
- 490 the four study sites (Shimokita, Kinkazan, Kashima, and Yakushima). Subalpine/alpine
- 491 zones are omitted from the map for convenience. The black star indicates the location of
- 492 Kinkazan, where the present study was conducted.
- 493 Figure 2. Monthly changes in the accumulated number of plant seeds found within faecal
- 494 samples in a given month. Life-form composition (categorized into tall trees, shrubs, vines,
- 495 hemi-parasites, and herbaceous plants) is also shown. Unidentified species are omitted. The
- 496 whole year was separated into four seasons: spring: March to May; summer: June to
- 497 August; fall: September to November; and winter: December to February. Numbers in
- 498 parentheses represent the number of collected faecal samples.
- 499 Figure 3. The relationship between mean cubic diameter (MCD) of seeds (mm) and their intact
- 500 rate (%).  $\bigcirc$ : fleshy fruit,  $\bigcirc$ : non-fleshy fruit.



Yakushima





Month	# fecal sample	DE	$DE_r^{a}$	$\Sigma$ of seeds	Σ of intact seeds	IR
Spring						
March	374	83	$0.22^{***}$	3209	1977	$0.62^{***}$
April	10	1	$0.10^{**}$	2	0	$0.00^{NS}$
May	158	21	0.13***	373	306	$0.82^{NS}$
Summer						
June	29	29	$1.00^{***}$	22327	2094	$0.94^{***}$
July	56	45	$0.80^{***}$	2806	2006	$0.72^{NS}$
August	93	90	$0.97^{***}$	9408	7743	$0.82^{NS}$
Fall						
September	76	56	$0.74^{**}$	3013	2365	$0.79^{NS}$
October	177	169	$0.96^{***}$	9439	6824	$0.72^{NS}$
November	134	130	$0.97^{***}$	22848	20198	$0.88^{***}$
Winter						
December	39	37	$0.95^{***}$	9535	8333	$0.87^{*}$
January	129	82	$0.64^{NS}$	2062	1122	$0.54^{***}$
Feburuary	19	4	$0.21^{*}$	20	20	$1.00^{NS}$
Annual mean	1294	747	0.58	85042	66248	0.78

Table 1. Monthly changes in dispersal event and total numbers of seeds defecated by the Japanese macaques on Kinkazan northern Japan

<sup>1</sup>Numbers in parentheses are percentage of total fecal samples. Results of binomial test are also shown. \*\*\*: P < 0.001, \*\*: P < 0.01, \*: P < 0.05, NS: not significant (P > 0.05).

DE: dispersal event,  $DE_r$ : rate of dispersal event, IR: intact rate

Food plant spacing	L ifa form	SUT	MCD	DE	DE	# intact seed	ds	Months in which	D 1 b	Main season(s) in	$\Sigma$ of goods	$\Sigma$ of intact	ID
Food plant species	Life form	501	(mm, Mean $\pm$ SD)	DE	$DL_r$	(Mean±SD	))	intact seeds appeared <sup>a</sup>	P-value	seed appearance <sup>c</sup>	$\geq$ of seeds	seeds	IK
Taxaceae													
Taxus cuspidata	high tree	С	$3.1 \pm 0.4$	1	0.00	1.0		Mar	—	Sp	1	1	1.00
Torreya nucifera*	high tree	С	10.8	3	0.00	$0.3 \pm$	0.6	Sep, Nov	0.157	F	3	1	0.33
Betulaceae													
Carpinus spp.*	high tree	С	$3.5 \pm 0.4$	117	0.09	$2.8$ $\pm$	6.2	Mar, Jun, Jul, Aug, Oct, Nov, Dec, Jan	0.270	Sp, W	756	328	0.43
Ulmaceae													
Zelkova serrata*	high tree	С	$3.4 \pm 0.3$	66	0.05	$1.9 \pm$	3.0	Mar, Aug, Oct, Nov, Jan	0.092	Su, F	354	125	0.35
Loranthaceae													
Viscum album	hemi-parasite	S	$4.0 \pm 0.4$	75	0.06	$13.5 \pm$	29.2	Mar, Nov, Dec, Jan, Feb	0.895	W	1113	1013	0.91
Schisandraceae													
Schisandra nigra	vine	S	$4.4 \pm 0.3$	49	0.04	$17.1 \pm$	25.0	Sep, Oct	0.297	F	838	838	1.00
Actinidiaceae													
Actinidia arguta	vine	S	$1.4 \pm 0.1$	12	0.01	$30.5 \pm$	33.0	<b>Jul</b> , Aug	0.381	Su	394	366	0.93
Rosaceae													
Rosa multiflora	shrub	S	$2.2 \pm 0.3$	221	0.17	83.9 ±	196.6	Mar, May, Oct, Nov, Dec, Jan, Feb	< 0.001	F, W	22475	18542	0.83
Pourthiaea villosa	shrub	S	$2.7 \pm 0.3$	85	0.07	$10.5 \pm$	18.1	Mar, Sep, Oct, Nov, Dec, Jan	0.088	F, W	1325	893	0.67
Prunus spp.	high tree	С	$5.2 \pm 0.2$	15	0.01	$1.7$ $\pm$	2.6	<b>Jun</b> , Jul	0.688	Su	80	26	0.33
Rubus microphyllus	shrub	S	$1.3 \pm 0.1$	13	0.01	$65.2 \pm$	147.8	<b>Jun</b> , Jul	1.000	Su	848	848	1.00
Sorbus japonica	high tree	S	$3.1 \pm 0.5$	13	0.01	$4.7$ $\pm$	5.6	Sep, Oct, Nov, Dec, Jan	0.353	F	91	61	0.67
Malus tschonoskii	high tree	S	$3.5 \pm 0.3$	74	0.06	$8.7$ $\pm$	9.7	Mar, Sep, Oct, Nov, Dec, Jan	< 0.001	F	665	644	0.97
Leguminosae													
Amphicarpaea bracteata*	herbaceous	С	$1.6 \pm 0.2$	33	0.03	$4.2$ $\pm$	8.0	Jul, Aug, Sep, Oct, Nov, Dec	0.310	Su, F	158	139	0.88
Desmodium podocarpum*	herbaceous	S	5.1	1	0.00	1.0		Nov		F	1	1	1.00
Rutaceae													
Zanthoxylum piperitum	shrub	С	$2.8 \pm 0.2$	49	0.04	$1.0 \pm$	1.0	Mar, Sep, Oct, Nov, Dec	0.563	F, W	81	49	0.61
Aquifoliaceae													
Îlex macropoda	high tree	S	$2.6 \pm 0.4$	55	0.04	11.3 ±	17.2	Mar, Aug, Nov, Dec, Jan	0.147	Sp	665	622	0.94
Rhamnaceae	C												
Berchemia racemosa	vine	S	$3.7 \pm 0.2$	126	0.10	$64.8 \pm$	62.5	Jun, Jul, <b>Aug</b> , Sp	0.001	Su	9897	8165	0.83
Vitaceae													
Vitis flexuosa	vine	S	$3.5 \pm 0.3$	79	0.06	$26.0 \pm$	29.9	Sep, Oct, Nov, Jan	0.023	F	2073	2054	0.99
Cornaceae													
Swida macrophylla	high tree	С	$3.4 \pm 0.1$	124	0.10	$1.4 \pm$	2.9	Mar, Jul, Sep, <b>Oct</b> , Nov, Dec, Jan	0.193	F	211	174	0.83
Cornus kousa	high tree	D	$4.5 \pm 0.4$	50	0.04	13.0 ±	25.2	Aug, Sep, Oct	0.071	F	1199	650	0.54
Araliaceae	C												
Acanthopanax trichodon	shrub	S	$2.5 \pm 0.2$	9	0.01	37.8 ±	53.2	Jul, Aug	0.437	Su	340	340	1.00
Ebenaceae													
Diospyros lotus	high tree	D	$7.8 \pm 0.4^{a}$	2	0.00	$1.0 \pm$	0.0	Nov	_	F	2	2	1.00
Solanaceae	0												
Tubocansicum anomalum	herbaceous	S	13 + 01	41	0.03	478 +	92.8	Mar Sep Oct Nov Dec	0 595	F	1994	1960	0.98
Caprifoliaceae	neroueeous	5	1.5 ± 0.1		0.02	17.0 1	2.0	11111, 5 <b>cp</b> , 5ct, 1101, 200	0.070	ľ	1771	1700	0.70
Viburnum dilatum	herbaceous	S	3.3 + 0.4	236	0.18	65.2 +	92.2	Mar. May. Oct. <b>Nov</b> . Dec. Jan. Feb	< 0.001	F	15945	15387	0.97
Gramineae	nereueeous	5		200	0.10		> 2.2	1111, 1111, 000, 1000, 200, 001, 100		•	10910	10007	0.97
Zovsia japonica*	herbaceous	С	1.3 + 0.2	17	0.01	14.5 +	19.4	May, Aug. Oct. Nov	0.300	Sp. Su	317	247	0.78
Oplismenus undulatifolius*	herbaceous	S	1.5	8	0.01	1.1 ±	0.4	Mar. Oct. Nov	0.435	~P, ~~ F	10	9	0.89
Digitaria ciliaris*	herbaceous	S	$12 + 01^{a}$	1	0.00	1.0	0.1	Ang		Su	1	1	1.00
Grominana 2*	herbaccous	S	$1.2 \pm 0.1$	20	0.00	454.0 +	440.5	Lun		Su	12102	12102	1.00
Gramineae 1*	herbaceous	S	$0.8 \pm 0.1$	29	0.02	$434.9 \pm$	449.5	Jun		Su	15192	15192	1.00
Gramineae 1*	herbaceous	S	$2.4 \pm 0.3$	10	0.01	$14.1 \pm 17.0$	13.8	Jun		Su	220	220	1.00
	nerbaceous	3	$1.4 \pm 0.4$	Z	0.00	$17.0 \pm$	1.4	Jun	_	Su	34	34	1.00
Clamatic aniifalia *	vina	C	10 00	2	0.00	1.0	0.0	I Oct		С., F	2	2	1.00
Ciemans apiijolla *	vine	C	$1.8 \pm 0.2$	2	0.00	$1.0 \pm$	0.0	Jun, Oct		Su, F	2	2	1.00
Oxandaceae	hauka	C	0.0 . 0.1	20	0.02	157 0	170 6	N# T	- 0.001	Q Q	(101	(121	1.00
Oxalis corniculata *	nerbaceous	3	$0.9 \pm 0.1$	39	0.03	137.2 ±	1/8.0	May, Jun	< 0.001	5p, 5u	0131	0131	1.00
v iolaceae	11	c	0.0	20	0.00		00 5	<b>T</b>		0	0001	0001	1.00
viola sp.*	nerbaceous	8	$0.8 \pm 0.1$	29	0.02	/0.0 ±	92.6	Jun		Su	2221	2221	1.00
	<b>1a1a</b>	C	17 . 0.2	A 77	0.02	4.0	6.0	Mar Oct Mar D	0.122	r.	200	100	0.07
rerula frutescens *	nerbaceous	C	$1.7 \pm 0.2$	45	0.03	$4.0 \pm$	6.3	Mar, Uct, Nov, Dec	0.133	F	209	180	0.86
unidentified	-		-	11	0.01			Mar, May, Jul, Aug, Oct, Nov					

## Table 2. Seeds found within fecal samples of Japanese macaques on Kinkazan, northern Japan (1999-2009).

DE: dispersal event,  $DE_r$ : rate of dispersal event, IR: intact rate, MCD: mean cubic diameter of seeds, SHT: seed handring technique (C: crunching, D: discarded, S: swallowing). \* non fleshy-fruited species.

<sup>a</sup>: name of each months are abbribiated. Months in which number of intact seeds exceed annual mean are shown in bold.

<sup>b</sup>: *P* - values less than 0.05 are shown in bold.

<sup>c</sup>: F: fall, Sp: spring, Su: summer, W: winter.

Month	# fecal	DF	# Intact	per feces	# Spec	cies pe	r feces	
WOItti	sample	DL	(M	SD)	(Mean + SD)			
Spring								
March	374	83	7.4	±	46.7	1.5	±	1.3
April	10	1		-			-	
May	158	21	2.3	±	9.8	1.3	±	0.6
Summer								
June	29	29	765.2	$\pm$	583.1	4.1	±	1.7
July	56	45	33.1	$\pm$	45.2	1.2	±	0.4
August	93	90	75.5	$\pm$	67.3	1.3	±	0.5
Fall								
September	76	56	34.3	$\pm$	49.5	2.3	±	1.0
October	177	169	43.3	$\pm$	67.1	2.9	±	1.5
November	134	130	162.5	$\pm$	183.5	2.9	±	1.2
Winter								
December	39	37	196.2	$\pm$	354.4	2.6	±	1.0
January	129	82	11.2	$\pm$	26.9	2.6	±	1.3
Feburuary	19	4	1.1	±	2.8	1.3	±	0.5
Annual mean	1294	747	58.2	±	175.6	2.0	±	0.9

**Table 3.** Monthly changes in numbers of intact seeds and species within single feces defecated by the Japanese macaques on Kinkazan, northern Japan.

DE: dispersal event

Study site	Yakushima	Kashima	Kinkazan	Shimokita
Climate zone	W	С	С	С
Sampling period	1995-2004	2000-2001	2000-2009	2000-2001
# Fecal samples	132	107	1294	75
$DE(DE_r)$	131(0.99)	99 (0.93)	747 <b>(</b> 0.58 <b>)</b>	59 (0.79)
# identified plant species	33	18	35	12
Life-form composition (%)				
High trees	13 (38.2)	3 (16.7)	11 (30.6)	1 (8.3)
Shrubs	11 (32.4)	7 (38.9)	5 (13.9)	3 (25.0)
Vines	9 (26.5)	6 (33.3)	5 (13.9)	7 (58.3)
Hemi-parasites	0 (0.0)	0 (0.0)	1 (2.8)	0 (0.0)
Herbaceous plants	0 (0.0)	2 (11.1)	13 (36.1)	1 (8.3)
<pre># Freshy-fruit species (%)</pre>	31 (91.2)	16 (88.9)	20 (55.6)	11 (91.7)
$MCD$ (Mean $\pm$ SD) (mm)	2.5 <b>±</b> 1.7 (N=12)	2.7±1.2 (N=11)	3.0±2.0 (N=35)	2.3±1.1 (N=15)
Range	1.1-6.4	1.2-4.1	0.8-10.8	1.2-4.1
# species (Mean±SD)	2.8±1.4 (N=131)	2.4±1.5 (N=99)	2.0±0.9 (N=747)	1.9 <b>±</b> 0.8 (N=59)
References	Otani (2005)	Otani (2003)	This study	Otani (2003)

Table 4. Regional variation in characteristics of seeds detected from feces of Japanese macaques.

C: cool temperate zone, W: warm temperate zone, DE: dispersal event,  $DE_r$ : rate of dispersal event,

*MCD* : mean cubic diameter of seeds.