



Dietary Profile of *Rhinopithecus bieti* and Its Socioecological Implications

Cyril C. Grueter · Dayong Li · Baoping Ren ·
Fuwen Wei · Carel P. van Schaik

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Abstract To enhance our understanding of dietary adaptations and socioecological correlates in colobines, we conducted a 20-mo study of a wild group of *Rhinopithecus bieti* (Yunnan snub-nosed monkeys) in the montane Samage Forest. This forest supports a patchwork of evergreen broadleaved, evergreen coniferous, and mixed deciduous broadleaved/coniferous forest assemblages with a total of 80 tree species in 23 families. The most common plant families by basal area are the predominantly evergreen Pinaceae and Fagaceae, comprising 69% of the total tree biomass. Previous work has shown that lichens formed a consistent component in the monkeys' diet year-round (67%), seasonally complemented with fruits and young leaves. Our study showed that although the majority of the diet was provided by 6 plant genera (*Acanthopanax*, *Sorbus*, *Acer*, *Fargesia*, *Pterocarya*, and *Cornus*), the monkeys fed on 94 plant species and on 150 specific food items. The subjects expressed high selectivity for uncommon angiosperm tree species. The average number of plant species used per month was 16. Dietary diversity varied seasonally, being lowest during the winter and rising dramatically in the spring. The monkeys consumed bamboo shoots in the summer and bamboo leaves throughout the year. The monkeys also foraged on terrestrial herbs and mushrooms, dug up tubers, and

C. C. Grueter (✉) · C. P. van Schaik
Anthropological Institute and Museum, University of Zürich, 8057 Zürich, Switzerland
e-mail: cyril_grueter@eva.mpg.de

D. Li
Qinling Golden Snub-nosed Monkey Research Center, College of Life Science, Northwest University,
710069 Xi'an, China

B. Ren · F. Wei
Key Laboratory of Animal Ecology and Conservation Biology, Institute of Zoology,
Chinese Academy of Sciences, 100101 Beijing, China

Present Address:

C. C. Grueter
Department of Primatology, Max Planck Institute for Evolutionary Anthropology, 04103 Leipzig,
Germany

consumed the flesh of a mammal (flying squirrel). We also provide a preliminary evaluation of feeding competition in *Rhinopithecus bieti* and find that the high selectivity for uncommon seasonal plant food items distributed in clumped patches might create the potential for food competition. The finding is corroborated by observations that the subjects occasionally depleted leafy food patches and stayed at a greater distance from neighboring conspecifics while feeding than while resting. Key findings of this work are that Yunnan snub-nosed monkeys have a much more species-rich plant diet than was previously believed and are probably subject to moderate feeding competition.

Keywords diet · China · colobine · feeding ecology · food competition

Introduction

Researchers study primate food habits for a variety of reasons. Such studies may reveal a species' resource requirements (Litvaitis 2000), and diet-related factors such as nutritional quality, distribution, and spatiotemporal fluctuations in abundance of food may also have far-reaching influences on the emergence of primate social organizations and social structure (Chapman 1990; van Schaik 1989; Wrangham 1980). They may affect group size (Kirkpatrick 1996), tendency toward fission-fusion (Anderson *et al.* 2002; van Schaik and van Noordwijk 1988), and the intensity and the nature of feeding competition (scramble and contest; Janson and van Schaik 1988; Koenig *et al.* 1998; Saj and Sicotte 2007a; van Schaik and van Noordwijk 1988). Nonhuman primates exhibit clear preferences for particular food items (Milton 1981). Many factors—both internal and external—mold patterns of food choice in primates, e.g., energy needs, nutrient requirements, constraints of the digestive system, body size, chemical and structural properties of foods, and spatiotemporal availability of food resources (Barton and Whiten 1994; Garber 1987; Kay 1984; Lambert 2007; McKey *et al.* 1981; Milton 1984; Oates 1987; Oftedal 1991).

Colobines possess specialized capacious and partitioned stomachs (Caton 1998; Stevens 1988) where microbial fermentation of cellulose takes place (Bauchop and Martucci 1968; Hume 1989; Kay and Davies 1994; Kuhn 1964). This adaptation enables them to eat food containing high levels of structural polysaccharides, i.e., cellulose and related compounds. Overall, colobines can be classified as herbivores, ingesting flowers, fruits, leaves, and seeds to varying degrees (Bennett and Davies 1994; Fashing 2007; Kirkpatrick 2007; Oates 1994). Another common feature of colobine feeding ecology is their seasonally varying dietary spectrum as a response to temporal variation in food resource availability, of which switching to less nutritious plant foods (mature leaves) during periods of shortage of preferred food items is a key element (Dasilva 1994; Fashing 2001b; Koenig and Borries 2001; Struhsaker 1975). Although this is true for many tropical-dwelling colobines, some outliers within the Colobinae exemplify the subfamily's great plasticity pertaining to environmental conditions and demonstrate that dietary strategy is only partly determined by evolutionary history (*sensu* Struhsaker and Oates 1975). A case in point are the Yunnan or black-and-white snub-nosed monkeys (*Rhinopithecus bieti*)

of the temperate climate zone, which despite similar anatomy and presumably physiology, exhibit a somewhat different foraging strategy.

Previous work has revealed that Yunnan snub-nosed monkeys are highly dependent on lichens (Kirkpatrick 1996; Xiang *et al.* 2007), thus being the only anthropoid primate whose main food is not a plant. Their dietary regimen has been shown to vary geographically and to depend on overall habitat condition and productivity: in high latitude/high altitude habitats, lichens constitute the major fraction of the diet in virtually every month and are complemented with leaves from dicots and monocots (Kirkpatrick 1996; Xiang *et al.* 2007). The natural environment of *Rhinopithecus bieti* is characterized by striking seasonal variation in food resource availability. Winter is a period of plant food deprivation, and an almost exclusive dependence on carbohydrate-rich lichens has been regarded as a key ecological adaptation (Kirkpatrick 1996; Xiang *et al.* 2007; Grueter and Xiang 2008). Groups associated with more productive habitats at lower elevations and latitudes have a more species-rich diet and include a substantial proportion of nonlichen foods on a seasonal basis, such as immature leaves, fruits/seeds, buds, flowers, bamboo shoots, and bark of various plants (Ding and Zhao 2004; Yang and Zhao 2001).

We conducted the study in the Samage Forest, which is located in the central part of the geographical range of *Rhinopithecus bieti* and thus is intermediate in altitude and latitude compared to the populations in the north and south. In a companion paper, we documented seasonality in food use and fallback strategies of *Rhinopithecus bieti* at Samage (Grueter *et al.* 2009). We showed that lichens were chosen year-round and comprised *ca.* 67% of all the feeding records. Lichens were complemented with plant material, *viz.* 16% buds and young leaves, 11% fruits, 4% mature leaves, and 2% other items. Seasonal feeding patterns on plant items exactly matched the temporal variation in the availability of the main plant phenophases. The monkeys exploited immature leaves prolifically in spring and ingested heavy quantities of fruit in summer and fall. Our primary aim here is to document the overall dietary spectrum of *Rhinopithecus bieti* at Samage and to provide a thorough assessment of habitat composition. The findings are important for comprehending the species' resource requirements and the carrying capacity of the habitat and thus have implications for conservation management.

An additional aim relates to the question of how diet affects socioecology. For colobines, researchers have discussed the issue especially in relation to the importance of scramble competition. Scramble competition is the collective exhaustion of limited resources, leading to lower foraging efficiency for all group members (Janson and van Schaik 1988). Scramble competition increases as groups increase in size and is thought to limit group size for many primates (Janson and Goldsmith 1995). As a result of a more rapid depletion of food patches, larger groups are forced to travel farther to ensure procurement of a sufficient amount of the food (Chapman and Chapman 2000; Janson and van Schaik 1988; Majolo *et al.* 2008; van Schaik and van Noordwijk 1988). Scramble competition is usually inferred if there is a positive association between group size and home range size or day journey length. Based on weak relationships between these variables in combination with small group sizes and ubiquity of food resources, folivorous or frugivorous-folivorous primates such as colobines have traditionally been viewed as experiencing only a low intensity of intragroup scramble competition (Fashing

2001a; Isbell 1991; Janson and Goldsmith 1995; Sterck *et al.* 1997; Yeager and Kirkpatrick 1998; Yeager and Kool 2000). However, group size effects have recently been demonstrated among various folivores (Gillespie and Chapman 2001; Koenig *et al.* 2008; Majolo *et al.* 2008; Saj and Sicotte 2007b; Snaith and Chapman 2008; Teichroeb *et al.* 2003; *cf.* Steenbeek and van Schaik 2001).

This study of a colobine living in very large groups may help to shed some more light on the issue. *Rhinopithecus bieti* are also notable for having an unusual social organization: they live in large bands that are composed of distinct core one-male units (OMU). Given the fact that lichens occur in profusion in their natural habitat, at least currently, *Rhinopithecus bieti* are thought to be free to form large groups, and food competition would not be expected to be prevalent (*ibid.*). The evidence for scramble competition in *Rhinopithecus bieti* is scant: we have previously shown a positive correlation between group size and home range size, controlling for productivity, for different populations of *R. bieti*, indicating scramble competition effects (Grueter *et al.* 2008a). However, home range size is probably not as good a proxy for competition as day journey length, the latter correlating less strongly with group size (Grueter and van Schaik, *unpub. data*). A different picture may emerge when considering nonlichen foods, i.e. plant resources, which are the focus here. We assess the degree to which the lichenivorous-folivorous-frugivorous dietary regimen of *Rhinopithecus bieti* generates potential for scramble and also contest competition. The tests presented here are rudimentary given the challenges of observing wild *Rhinopithecus bieti*. If food competition occurs, then we predict that 1) preferred food species (species with high selection ratios) occur at low densities across the home range and are spatially clumped and thus can probably not accommodate all band members (interunit contest; *cf.* Koenig *et al.* 1998); 2) valued patches of food are being depleted (intra-band scramble; *cf.* Snaith and Chapman 2005), and 3) unit members avoid co-feeding (intra-unit scramble or contest; Saj and Sicotte 2007b) by having fewer nearest neighbors when feeding versus resting, assuming that dispersion reduces competition (*cf.* van Schaik and van Noordwijk 1988).

Methods

Study Site

We conducted the study in the predominantly temperate Samage Forest near the village of Gehuaqing (27°34'N, 99°17'E) in Yunnan's Baimaxueshan National Nature Reserve. Narrow valleys and steep hillsides characterize the topography at the site and land cover is a mosaic of mixed coniferous and deciduous-broadleaf forest (at 2900–3600 m), subalpine George's fir forest (3500–4000 m), montane sclerophyllous oak forests (3200–3500 m), subtropical evergreen broadleaf forest (2500–3000 m), Yunnan pine forest (2500–3100 m), as well as cattle pastures at various elevations. Umbrella bamboos (*Fargesia* spp.) and rhododendrons formed an important element of the underbush in all vegetation types. Parts of the Samage Forest have been selectively logged, and anthropogenic disturbance in the form of livestock grazing and collection of NTFP (nontimber forest products) is still widespread. The habitat of the monkeys at this locality ranged from 2500 m to 4000

m and included all major vegetation types, with mixed forest being the most used ecotype and clearcuts being unsuitable habitat for *Rhinopithecus bieti*. The semihabituated focal group comprised *ca.* 410 members.

Climate

Annual rainfall was 1004 mm, and mean annual temperature was 14.3°C at 2448 m (800 m below the altitude the focal group most frequently visited). Distribution of precipitation was highly irregular, but temperature varied strikingly with seasons: there was a steep increase in rainfall from spring onwards and a prolonged winter drought season with freezing nights (Grueter *et al.* 2009; Li *et al.* 2008). Complete snow cover rarely lasted for more than a few days within the frequently used zone of the group, as snowfall was followed by prolonged sunny days.

Data Collection

C. C. Grueter collected data on diet composition via scan sampling over 20 mo between September and July 2007. On 116 d, we obtained a total of 3872 feeding records: 1151 in fall (September–November), 772 in winter (December–February), 1314 in spring (March–May), and 635 in summer (June–August). The rugged terrain with steep-sided ravines and impenetrable undergrowth (bamboo, etc.) made tracking difficult, and thus distance observations from prominent topographical features (rocks, etc.) with the use of a spotting scope were the methods of choice. Occasionally we also observed the group at close range.

We took scans of all visible individuals at 15 min or 30-min intervals. If a large number of monkeys were in view, we chose 30-min scans; if only a small number was visible, we performed 15-min scans. Scans had to be completed ≥ 5 min before the beginning of the next scan. Every scan included information on date, time, and weather conditions. For every subject scanned, we recorded age, sex, activity, as well as distance and identity (age-sex class) of the nearest neighbor. Scan data were spread more or less evenly throughout the day. We divided age/sex classes into the following categories: adult male, adult female, juvenile (*ca.* 1–4 yr old), subadult male, and infant (<1 yr). We used the category SAMOF (subadult male or female) for cases wherein it was not possible to determine the sex of an individual whose body size was close to or larger than that of an adult female, but was not accompanied by an infant (*cf.* Bleisch *et al.* 1993).

Scan records of feeding behavior also included the food item, plant part, and its age as well as plant species whenever possible. We distinguished the following foods: lichens (fruticose vs. foliose), young leaves (including spring buds/shoots), mature leaves, buds (dormant leaf buds), flowers, flower buds, bark, pith, fruit or seeds (both ripe and unripe), invertebrates, snow, fungi, water, bamboo shoots, and tuber. It was usually difficult to see whether the small fruits were eaten wholly or whether the flesh was discarded. If we were unable to identify the tree taxon visually, we attempted to collect some samples from that feeding tree or a nearby tree of the same taxon for later identification.

Outside of scan sessions, we recorded all partially consumed and discarded foods on the forest floor with tooth marks or other signs of having been handled by the

monkeys. We used evidence from such feeding signs as a complementary measure to estimate seasonal variance in diet composition. We used the diameter of a feeding litter to quantify remains roughly as small (<1 m; score 1), medium sized (1–3 m; score 2), or large (> 3 m; score 3). Observational sampling was usually biased toward arboreal feeding, so the importance of terrestrial foods such as bamboo shoots was likely underrepresented.

We investigated the composition of the forest via stratified random sampling, i.e., we subdivided the central part of the home range of the study group, which largely corresponds to the core area of the home range, into 5 distinct forest types or strata (Mueller-Dombois and Ellenberg 1974). We established a total of 67 plots of 20 m × 20 m each (area: 26,800 m²) in which we recorded species, total height, bole height, crown diameter, and circumference for all trees ($n=1851$) with girth >40 cm. The different strata and the exact vegetation sampling regime are described fully in Li *et al.* (2008).

On a monthly basis, we recorded presence/absence of fruits, flowers, and young leaves for 157 food trees and calculated the percentage of trees bearing each of the phenophases every month. For details on phenology monitoring, see Grueter *et al.* (2008a).

Data Analysis

One can obtain an indication of the degree to which primates are selective in their choice of food tree species by calculating a selection index (Krebs 1999). The index compares the proportion of feeding observations of a plant species with the relative abundance of the species concerned as estimated from the tree plots. We used basal area to express the relative species crown biomass and potential food abundance, and calculated the selection index, W_i , from the formula:

$$W_i = O_i/P_i$$

wherein O_i is the percentage of feeding observations for species i , and P_i is the percentage of total basal area accounted for by species i . $W_i > 1$ indicates preference and $W_i < 1$, avoidance.

Using tree data obtained from the plots, we measured the pattern of dispersion of important food tree species. We used the coefficient of dispersion (CD; *cf.* Koenig *et al.* 1998). The CD refers to the ratio of the variance to the mean of the number of species in a sample. If species were distributed randomly, their allocation across the plots in the sample would correspond to a Poisson distribution (mean equals variance; CD = 1). CD > 1 indicates a clumped/contagious distribution, while a CD < 1 shows a uniform distribution. We determined significant departures from randomness, i.e., departure from a variance-to-mean ratio of 1.0, using the χ^2 statistic (Brower *et al.* 1998; Perry and Mead 1979).

To examine if unit members avoided cofeeding, we tested whether OMU members had fewer nearest neighbors when feeding versus resting. To assess whether valued patches of food were being exhausted, we used opportunistic evidence of total patch depletion, i.e., leafy patches defoliated as a result of intensive foraging.

Results

Forest Composition

We recorded a total of 80 tree species of 23 families in the botanical plots (Fig. 1; Table I). An additional 9 species of rare woody plants —*Bretschneidera sinensis* (Bretschneideraceae), *Magnolia campbellii* (Magnoliaceae), *Corylus chinensis* (Betulaceae), *Populus yunnanensis* (Salicaceae), *Meliosma yunnanensis* (Sabiaceae), *Cerasus conadenia* (Rosaceae), *Clethra delavayi* (Clethraceae), *Ilex delavayi* (Aquifoliaceae), and *Rhododendron sinogrande* (Ericaceae)— are not represented

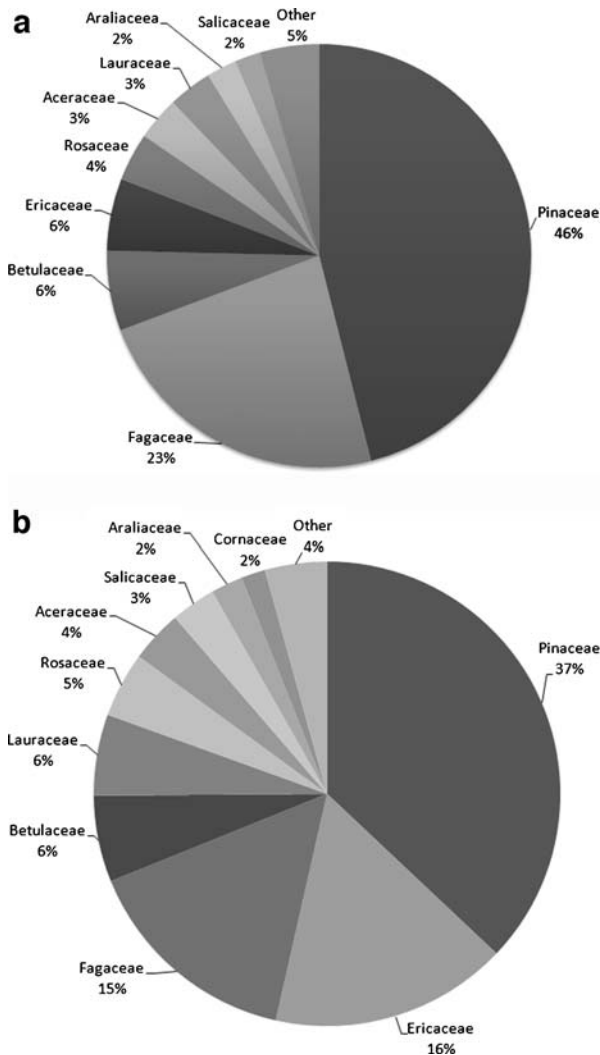


Fig. 1 The most common plant families at the Samage Forest, Baimaxueshan Nature Reserve. **(a)** Based on basal area. **(b)** Based on no. of stems ($n=1898$).

Table I The 20 most common tree species at the Samage Forest based on basal area

Rank	Species	Family	Basal area/ha (cm ² /ha)	% of total biomass
1	<i>Abies georgei</i>	Pinaceae	84,734	13.3
2	<i>Cyclobalanopsis cf. gambleana</i>	Fagaceae	83,359	13.0
3	<i>Picea likiangensis</i>	Pinaceae	80,014	12.5
4	<i>Pinus yunnanensis</i>	Pinaceae	57,869	9.0
5	<i>Tsuga dumosa</i>	Pinaceae	41,230	6.4
6	<i>Quercus pannose</i>	Fagaceae	37,128	5.8
7	<i>Abies ernestii</i>	Pinaceae	23,294	3.6
8	<i>Rhododendron rubiginosum</i>	Ericaceae	18,651	2.9
9	<i>Quercus rehderiana</i>	Fagaceae	18,185	2.8
10	<i>Alnus nepalensis</i>	Betulaceae	15,389	2.4
11	<i>Acanthopanax evodiaefolius</i>	Araliaceae	14,724	2.3
12	<i>Betula utilis</i>	Betulaceae	14,644	2.3
13	<i>Machilus microcarpa</i>	Lauraceae	14,333	2.2
14	<i>Acer laxiflorum</i>	Aceraceae	10,015	1.6
15	<i>Corylus cf. wangii</i>	Betulaceae	8802	1.4
16	<i>Sorbus cf. thibetica</i>	Rosaceae	8245	1.3
17	<i>Quercus aliena</i>	Fagaceae	8094	1.3
18	<i>Taxus yunnanensis</i>	Taxaceae	7315	1.1
19	<i>Tilia chinensis</i>	Tiliaceae	6288	1.0
20	<i>Cornus macrophylla</i>	Cornaceae	6166	1.0

in the plots. The Pinaceae contributed the greatest biomass at Samage based on both basal area and stem density. The 2 top families together, Pinaceae and Fagaceae, accounted for 69% of the total basal area. The 3 top families, these 2 plus Ericaceae, together accounted for 75% of the total basal area and 69% of the total stem density. The 3 dominant tree species by basal area at Samage were *Abies georgei*, *Cyclobalanopsis cf. gambleana*, and *Picea likiangensis*. Thirty-five percent of the tree species were evergreen, and 65% were deciduous. Of the conifers ($n=10$ species), 10% were deciduous (*Larix*), whereas 27% of the broadleaf trees were evergreen.

Dietary Repertoire

Approximately 150 different vegetative food items from at least 94 species and 38 families contributed to the diet of the focal group. Foraging occurred both on the ground and in the canopy. Subjects obtained food items were obtained from 40 woody plant species (49% of all available tree species), 22 shrubs, 1 semiparasitic shrub, 7 vines, 2 root-parasitic herbs, and 14 species of terrestrial herbaceous vegetation (THV). Food lists are provided in Tables II and III. In terms of stem density, food trees, excluding species supporting lichen only, accounted for 30.4% ($n=40$ species) of the trees in the study area. In terms of relative dominance, the basal area of food trees comprised 35% of the total basal area in the study area. The

Table II Food repertoire of *Rhinopithecus bieti* at Samage: vascular plants

Predominant life form	Species	Part	Month
	Angiosperms		
	Fagaceae		
Tree	<i>Quercus aliena</i>	Shoot	Apr
Tree	<i>Quercus pannosa</i>	Seed	Sep
		Flower	Jul
Tree	<i>Cyclobalanopsis cf. gambleana</i>	Seed	Jan
		Pith	Apr
	Juglandaceae		
Tree	<i>Pterocarya delavayi</i>	Young leaf	Apr
	Aceraceae		
Tree	<i>Acer</i> sp.	Bud	Feb
Tree	<i>Acer</i> sp.	Seed	Sep
Tree	<i>Acer laxiflorum</i> var. <i>laxiflorum</i>	Young leaf	Apr, May
		Petiole	May
		Flower	May
Tree	<i>Acer mono</i> var. <i>macropterum</i>	Young leaf	Apr
Tree	<i>Acer caesium</i>	Young leaf	Apr
Tree	<i>Acer hookeri</i>	Young leaf	Apr
Tree	<i>Acer caudatum</i>	Flower bud	May
		Young leaf	May
	Bretschneideraceae		
Tree	<i>Bretschneidera sinensis</i>	Petiole	Oct
		Seed	Sep
	Araliaceae		
Tree	<i>Acanthopanax evodiaefolius</i>	Mature leaf	Jul, Aug, Sep, Oct
		Young leaf	Apr, May, Jun
		Flower bud	Feb, Apr
		Fruit	Aug, Sep, Oct, Nov, Jan, Feb
		Bark	Mar, May, Sep
		Bud	Jan, Feb, Mar, Dec
		Petiole	Jun, Jul
Shrub	<i>Acanthopanax leucorrhizus</i> var. <i>fulvescens</i>	Leaf	Oct
	Celastraceae		
Tree	<i>Euonymus theifolius</i>	Seed	Oct
	Hydrangeaceae		
Tree	<i>Hydrangea heteromalla</i>	Mature leaf	Jul, Sep, Oct, Nov
		Pith	Jul
Tree	<i>Hydrangea</i> sp.	Mature leaf	Aug
Tree	<i>Hydrangea</i> sp.	Bark	Jan

Table II (continued)

Predominant life form	Species	Part	Month
Tree	<i>Philadelphus delavayi</i> var. <i>delavayi</i>	Mature leaf	Oct, Sep, Aug
		Bark	Mar
Shrub	<i>Philadelphus</i> sp.	Young leaf	May
Shrub	<i>Deutzia glomeruliflora</i>	Mature leaf	Jul, Nov
		Young leaf	May
		Flower	May
Clethraceae			
Tree	<i>Clethra delavayi</i>	Petiole	May
		Young leaf	May
Cornaceae			
Shrub	<i>Helwingia japonica</i>	Mature leaf	Sep
Tree	<i>Cornus macrophylla</i>	Fruit	Aug, Sep
		Young leaf	Apr
Rosaceae			
Tree	<i>Sorbus rufopilosa</i>	Mature leaf	Jul, Oct
		Fruit	Aug, Sep
Tree	<i>Sorbus rehderiana</i> var. <i>cupreonitens</i>	Young leaf	May
Tree	<i>Sorbus</i> sp.	Young leaf	Jun
Tree	<i>Sorbus</i> sp.	Bud	Feb
Tree	<i>Sorbus oligodonta</i>	Young leaf	Apr
		Mature leaf	Oct, Nov
		Fruit	Aug, Oct, Nov
Tree	<i>Sorbus monbeigii</i>	Mature leaf	Jul, Sep
Tree	<i>Sorbus hupehensis</i>	Young leaf	Apr
Tree	<i>Sorbus macrantha</i>	Mature leaf	Oct
		Fruit	Jan, Oct
		Bud	Feb
Tree	<i>Sorbus thibetica</i>	Young leaf	Apr, May, Jun
		Fruit	Aug, Sep, Oct, Nov
		Flower, flower bud	Apr, May
Tree	<i>Sorbus epidendron</i>	Fruit	Feb
Shrub	<i>Stranvaesia davidiana</i>	Mature leaf	Mar, Apr, Dec
		Fruit	Oct
Tree	<i>Padus obtusata</i>	Fruit	Jun, Oct
		Young leaf	Apr, May, Jun
		Mature leaf	Jul, Oct
		Bud	Feb
		Flower bud	May
Tree	<i>Malus yunnanensis</i>	Young leaf	Apr, May
Tree	<i>Malus</i> cf. <i>prattii</i>	Fruit	Sep

Table II (continued)

Predominant life form	Species	Part	Month
Shrub	<i>Rosa macrophylla</i>	Seed	Oct
Shrub	<i>Rubus alexeterius</i>	Fruit	Jul
Tree	<i>Cerasus conadenia</i>	Mature leaf	Oct
Tree	<i>Cerasus patentipila</i>	Flower bud	May
		Young leaf	May
Tree	<i>Cerasus clarofolia</i>	Young leaf	Apr
		Flower bud	Apr
	Caprifoliaceae		
Shrub	<i>Lonicera tangutica</i>	Mature leaf	Oct
Vine	<i>Lonicera acuminata</i>	Mature leaf, fruit	Mar
Shrub	<i>Viburnum betulifolium</i>	Fruit	Feb, Oct
Shrub	<i>Viburnum nervosum</i>	Young leaf	May
Tree	<i>Viburnum cylindricum</i>	Fruit	Oct
Herb	<i>Sambucus adnata</i>	Fruit	Sep
Shrub	<i>Leycesteria formosa</i>	Mature leaf	Aug
	Balanophoraceae		
Root-parasitic herb	<i>Balanophora involucrata</i>	Tuber	Oct
	Lililiaceae		
Shrub	<i>Polygonatum cirrhifolium</i>	Mature leaf	Aug
Herb	<i>Maianthemum atropurpureum</i>	Leaf	May
Herb	<i>Maianthemum forrestii</i>	All	Jun
	Berberidaceae		
Shrub	<i>Berberis</i> sp.	Young leaf	Apr
	Ranunculaceae		
Vine	<i>Clematis obtusidentata</i>	Mature leaf	Oct
Vine	<i>Clematis chrysocoma</i>	Mature leaf	May
Herb	<i>Thalictrum delavayi</i>	Leaf	Oct
	Orobanchaceae		
Root-parasitic herb	<i>Boschniakia himalaica</i>	Tuber	Nov
	Balsaminaceae		
Herb	<i>Impatiens arguta</i>	Leaf	Oct
Herb	<i>Impatiens xanthocephala</i>	Leaf	Oct
	Urticaceae		
Herb	<i>Urtica</i> sp.	Leaf	Oct
Herb	<i>Pilea angulata</i>	Leaf	May, Oct, Nov
Herb	<i>Elatostema obtusum</i>	Leaf	Nov
	Cruciferae		
Herb	<i>Cardamine macrophylla</i> var. <i>macrophylla</i>	Leaf	Nov

Table II (continued)

Predominant life form	Species	Part	Month
Herb	<i>Eutrema yunnanense</i> Schisandraceae	Leaf	Nov
Vine	<i>Schisandra rubriflora</i> Gramineae	Fruit Young leaf	Nov May
Shrub	<i>Fargesia cf. dura</i>	Leaf Shoot	Feb, Mar, Sep, Oct, Nov Jun, Aug
Shrub	<i>Fargesia cf. melanostachys</i>	Young leaf Mature leaf Shoot	Mar, Apr, May Jul, Nov, Dec May, Jun, Jul
Shrub	<i>Fargesia</i> sp.	Mature leaf	Jan, Aug
Shrub	<i>Yushania</i> sp. Tiliaceae	Mature leaf	Oct
Tree	<i>Tilia chinensis</i>	Mature leaf Young leaf	Oct Apr
Tree	<i>Salix rehderiana</i> Salicaceae	Young leaf Catkin	Apr Mar, Apr
Tree	<i>Populus davidiana</i>	Young leaf Flower bud Bud Bark	Apr Mar Mar Feb, Oct
Tree	<i>Betula utilis</i> Betulaceae	Young leaf	May
Tree	Unid. species	Young leaf	Apr
Tree	<i>Corylus ferox</i>	Young leaf	Apr
Tree	<i>Machilus yunnanensis</i> Lauraceae	Bud Mature leaf	Mar Apr, Sep
Tree	<i>Machilus microcarpa</i>	Seed	Sep
Tree	<i>Litsea chunii</i>	Young leaf Flower	Apr, May Apr
Shrub	<i>Ligustrum cf. delavayanum</i> Oleaceae	Fruit	Oct
Tree	<i>Syringa yunnanensis</i>	Mature leaf	Aug
Shrub	<i>Piptanthus nepalensis</i> Leguminosae	Fruit	Sep
Herb	<i>Senecio scandens</i> Compositae	Leaf	Oct
Herb	<i>Ligularia nelumbifolia</i> Ericaceae	Leaf	Aug

Table II (continued)

Predominant life form	Species	Part	Month
Tree	<i>Rhododendron rubiginosum</i>	Flower	May
Shrub	<i>Rhododendron yunnanense</i>	Young leaf	May, Jun
	<i>Rhododendron selense</i>	Petiole	May
Tree	<i>Enkianthus cf. deflexus</i>	Young leaf	Apr
Shrub	<i>Enkianthus chinensis</i>	Flower	Apr
Shrub	<i>Lyonia villosa</i>	Young leaf	Apr
	Loranthaceae		
Semi-parasitic shrub	<i>Arceuthobium pini</i>	All	Oct
	Laradizabalaceae		
Vine	<i>Holboellia angustifolia</i>	Mature leaf	Apr
Shrub	<i>Decaisnea fargesii</i>	Fruit	Aug
	Acrinidiaceae		
Vine	<i>Actinidia pilosula</i>	Young leaf	Apr
		Fruit	Oct
	Vitaceae		
Vine	<i>Cayratia cf. cardiospermoides</i>	Fruit	Oct
	Sabiaceae		
Tree	<i>Meliosma yunnanensis</i>	Fruit, mature leaf	Oct
	Saxifragaceae		
Herb	<i>Chrysplenium davidianum</i>	All	
	Aquifoliaceae		
Tree	<i>Ilex</i> sp.	Mature leaf	Jan, Feb, Apr, Nov, Dec
		Flower	Dec
		Young leaf	Feb
Tree	<i>Ilex delavayi</i>	Mature leaf	Jul, Aug
		Bark	Jan
	Gymnosperms		
	Cupressaceae		
Tree	<i>Sabina squamata</i>	Fruit	Mar
	Pinaceae		
Tree	<i>Larix speciosa</i>	Petiole	Apr, May, Jul, Sep

The list is based on systematic scan observations, opportunistic observations, and food remnants.

top 10 food tree species (Table IV) accounted for >90% of the total feeding time on plant foods. All top 10 food tree species had selection ratios >1, with a few species having extraordinarily high selection indices, i.e., *Pterocarya delavayi* (71.7; represented by a single specimen in the plots), *Padus obtusata* (41.8), and *Acanthopanax evodiaefolius* (20.4; Table IV). Many species listed in Table II were fed on infrequently. As estimated from feeding records, immature leaves of *Acanthopanax evodiaefolius* were the single most prominent food type (Table V).

Table III Food repertoire of *Rhinopithecus bieti* at Samage: cryptogams, i.e., nonvascular plants (lichens) and fungi

Species	Season
Lichens	
Parmeliaceae ^a	
<i>Usnea longissima</i>	Year-round
<i>Usnea</i> sp.	Year-round
<i>Bryoria confusa</i>	Year-round
<i>Bryoria</i> cf. <i>trichodes</i> cf. ssp. <i>americana</i>	Year-round
<i>Cetrelia</i> sp.	Apr, Dec
1 unk species	Nov
Fungi	
Russulaceae	
<i>Russula</i> sp. 1	Jul, Aug, Sep, Oct
<i>Russula</i> sp. 2	Sep
<i>Russula</i> sp. 3	Aug
<i>Russula senecis</i>	Sep
Amanitaceae	
<i>Amanita hemibapha</i> var. <i>ochracea</i>	Aug
<i>Amanita fritillaria</i>	Sep
<i>Amanita flavipes</i>	Sep
Gomphaceae	
<i>Gomphus floccosus</i>	Aug, Sep
Boletaceae	
<i>Boletus</i> sp.	Aug, Sep

The list is based on systematic scan observations, opportunistic observations, and food remnants.

^a The species of lichens have previously been reported in Grueter *et al.* (2009).

As estimated from feeding remains, shoots of *Fargesia* spp. were the most important dietary item, followed by fruits of *Acanthopanax evodiaefolius*, fruits of *Sorbus* cf. *thibetica*, fruits of *Sorbus* spp., and mature leaves of *A. evodiaefolius*. The average number of plant species and specific plant food items used per month was 16 and 19, respectively. The richness of food species (controlled for observation time) peaked in April/May, August, and October (Fig. 2).

Mature leaves were chosen and ingested from both deciduous and evergreen trees, but only a few woody species were important sources of mature leaves, i.e., the deciduous *Philadelphus delavayi*, *Sorbus* spp., *Acanthopanax evodiaefolius*, and the evergreen *Ilex* sp. For some species, only petioles were eaten, e.g., *Bretschneidera sinensis*; for others, only the leaf blades, e.g., *Stranvaesia davidiana*; and for yet others both leaf blades and petioles, e.g., *Acanthopanax evodiaefolius*.

The snub-nosed monkeys fed on subterranean parts of *Boschniakia himalaica* and *Balanophora involucrata*. They spent a considerable amount of time unearthing unidentified tubers (hidden food items). Juveniles and females ate snow in winter, but only on rare occasions. The monkeys drank water from small ponds and streams.

Table IV Percentage of feeding records^a for the 10 top-ranked identified plant species (out of all records of plant feeding for which the plant species could be identified to at least genus level)

Rank	Species	Family	% of feeding records	Selection index
1	<i>Acanthopanax evodiaefolius</i>	Araliaceae	47.0	20.4
2	<i>Sorbus</i> spp. ^b	Rosaceae	15.3	9.5
3	<i>Acer</i> spp. ^c	Aceraceae	7.7	3.4
4	<i>Sorbus</i> cf. <i>thibetica</i>	Rosaceae	6.3	4.9
5	<i>Fargesia</i> spp. ^d	Gramineae	5.5	NA ^e
6	<i>Pterocarya delavayi</i>	Juglandaceae	4.5	71.7
7	<i>Cornus macrophylla</i>	Cornaceae	1.8	1.8
8	<i>Padus obtusata</i>	Rosaceae	1.5	41.8
9	<i>Tilia</i> cf. <i>chinensis</i>	Tiliaceae	1.2	1.2
10	<i>Litsea chunii</i>	Lauraceae	0.9	8.6

¹ In total, there were 2674 identified feeding records (1788 for lichens, 886 for plants).

^b Includes *Sorbus oligodonta*, *S. rufopilosa*, *S. rehderiana*, *S. monbeigii*, *S. hupehensis*, and *S. macrantha*; distinguishing among these species was difficult.

^c Includes *Acer laxiflorum*, *A. mono.* *A. hookeri*, *A. caesium*, and *A. caudatum*; distinguishing among these species was difficult.

^d Includes *Fargesia* cf. *melanostachys* and *F.* cf. *dura*. Identifying bamboos to species level was not possible.

^e Bamboo.

Table V Number of feeding records for the 10 top-ranked specific food items

Rank	Item	Family	No. of feeding records
1	<i>Acanthopanax evodiaefolius</i> (young leaves)	Araliaceae	207
2	<i>Acanthopanax evodiaefolius</i> (fruits)	Araliaceae	174
3	<i>Sorbus</i> spp. ^a (fruits)	Rosaceae	107
4	<i>Acer</i> spp. ^b (young leaves)	Aceraceae	63
5	<i>Fargesia</i> spp. ^c (mature leaves)	Gramineae	49
6	<i>Pterocarya delavayi</i> (young leaves)	Juglandaceae	42
7	<i>Acanthopanax evodiaefolius</i> (buds)	Araliaceae	40
8	<i>Sorbus</i> cf. <i>thibetica</i> (young leaves)	Rosaceae	35
9	<i>Sorbus</i> cf. <i>thibetica</i> (fruits)	Rosaceae	19
10	<i>Cornus macrophylla</i> (fruits)	Cornaceae	17

^a Includes *Sorbus oligodonta*, *S. rufopilosa*, *S. rehderiana*, *S. monbeigii*, *S. hupehensis*, and *S. macrantha*; distinguishing among these species was difficult.

^b Includes *Acer laxiflorum*, *A. mono.* *A. hookeri*, *A. caesium*, and *A. caudatum*; distinguishing among these species was difficult.

^c Includes *Fargesia* cf. *melanostachys* and *F.* cf. *dura*. Identifying bamboos to species level was not possible.

Contrary to Xiang *et al.* (2007), we never observed the monkeys of this population eating resin. We recorded a case of predation on bird eggs. One individual fed on the flesh of an unidentified flying squirrel (Sciuridae). We observed the monkeys biting into mushrooms in the fall. We also saw them removing the bark of dead fallen and standing trees, mostly *Abies georgei*, and disassembling rotten and brittle tree stumps. Although we never clearly saw an individual actually eating an insect, the latter observations may indicate foraging on invertebrates. We witnessed feeding on bamboo (*Fargesia* spp.) leaves in all seasons. The monkeys consumed large quantities of bamboo shoots (*Fargesia* spp.) in summer.

Food Competition

Although the overall density of trees was 708 stems/ha, the 3 main food trees comprised only 42.5 individuals/ha (6.0%; Table VI). The single most important woody food species, *Acanthopanax evodiaefolius*, occurred in only 16.4% of the vegetation plots. All top 3 food tree taxa —*Acanthopanax evodiaefolius*, *Sorbus thibetica*, and *Sorbus* spp.— were clumped in distribution. On several occasions, we encountered evergreen trees that were completely defoliated after the group of *Rhinopithecus bieti* had visited them, demonstrating full patch depletion (Table VII). The monkeys especially sought the leaves of *Ilex* and *Philadelphus*. OMU members had fewer nearest neighbors when feeding versus resting. When an OMU member was feeding ($n=5020$; excluding bachelors and infants), there was another member in proximity (0–2 arm's lengths) in only 4.6% of all records versus 41.5% when an individual was resting.

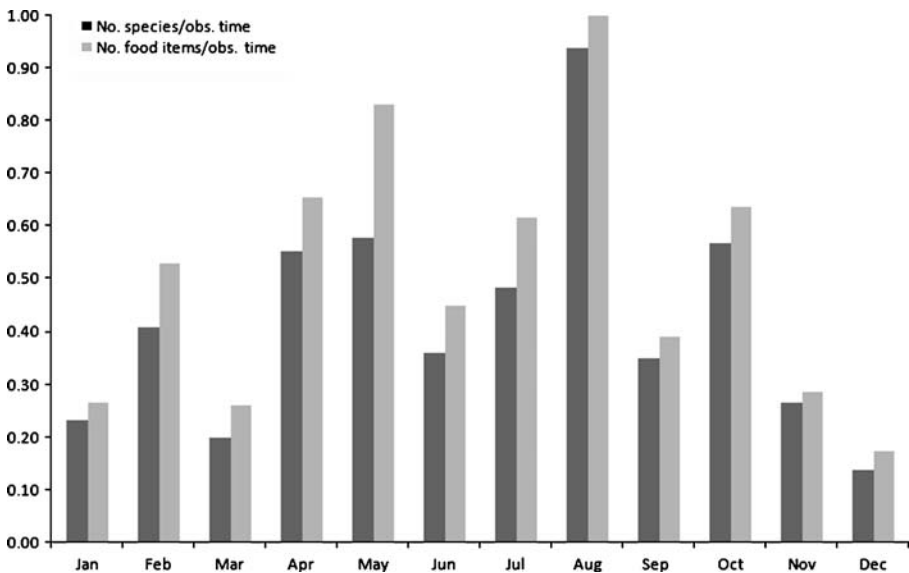


Fig. 2 Food species richness of *Rhinopithecus bieti* at Samage compared among months, calculated as the number of food species and food items per observation time (scan-based visual observation).

Table VI Density and dispersion of major^a food of *Rhinopithecus bieti* at Samage

Species	<i>n</i>	Density (ha ⁻¹)	% plots ^c	Index of dispersion (variance-to- mean-ratio)	Distribution	Mean/plot (if <i>n</i> >0)
<i>Acanthopanax evodiaefolius</i>	41	15.3	16.4 (31.4)	5.2** (4.7**)	Clumped	3.7
<i>Sorbus thibetica</i>	16	6.0	17.9 (28.6)	1.4** (1.3)	Clumped ^d	1.3
<i>Sorbus</i> spp. ^b	57	21.3	26.9 (48.6)	3.8** (3.1**)	Clumped	3.1
<i>A. evodiaefolius</i> , <i>S. thibetica</i> & <i>Sorbus</i> spp.	114	42.5	40.3 (68.6)	5.7** (4.5**)	Clumped	4.2
All	1898	708.2				

^a “Major” here refers to the 2 most important food plant genera *Sorbus* and *Acanthopanax*, which together make up almost 69% of the feeding records.

^b Includes *Sorbus oligodonta*, *S. rufopilosa*, *S. rehderiana*, *S. monbeigii*, *S. hupehensis*, and *S. macrantha*; these species were lumped together for the analyses because distinguishing among them *in situ* was not straightforward.

^c Percentage of plots with species *i*.

^d Random in mixed forest.

** Designates a significant deviation from randomness ($p < 0.05$). Numbers in parentheses refer to the 35 plots in mixed forest only.

Discussion

We demonstrated that the dietary richness of this population of *Rhinopithecus bieti* encompassed 94 different plant species and a few unusual items: mammals, mushrooms, tubers, bamboo. Moreover, we showed that plant species were not consumed in accordance with their spatial abundance, but that the monkeys showed a strong preference for uncommon species with a clumped distribution and that, based on several lines of evidence, they may experience some scramble and contest competition.

Plant Food Selection and Diversity

Seventy-five percent of the woody stems at Samage were Pinaceae (pines, firs, hemlocks and spruces), Fagaceae (oaks), and Ericaceae (rhododendrons), none of

Table VII *Ad libitum* observations of broadleaf trees having been depleted through intense foraging by *Rhinopithecus bieti*

Species	No. of depleted trees and dates	BA (%)
<i>Philadelphus delavayi</i>	5 (Oct 05), 2 (Nov 05), 5 (Aug 06), 2 (Oct 06), 2 (Nov 06)	0.08
<i>Ilex</i> sp.	2 (Oct 05), 1 (Jan 07)	0.03
<i>Ilex delavayi</i>	3 (Oct 05), 3 (Nov 05), 1 (Feb 06), 1 (Mar 06), 2 (Aug 06), 2 (Oct 06), 4 (Nov 06)	0 ^a
<i>Hydrangea heteromalla</i>	1 (July 07)	0.03
<i>Malus yunnanensis</i>	1 (May 07)	0.09

^a Not recorded in plots.

which were of direct dietary importance to the snub-nosed monkeys (*cf.* Kirkpatrick 1996). However, conifers and oaks were important lichen-bearing trees and were almost exclusively exploited for this nonplant resource (Grueter *et al.* 2009). *Rhinopithecus bieti* at Samage derived its plant diet mostly from the deciduous angiosperms families Araliaceae, Rosaceae, Aceraceae, Juglandaceae, and Cornaceae. High selection ratios for most of these angiosperm plant species indicate strong selectivity for uncommon species, which is likely due to variability in nutritional quality.

Only a few fruit species provided the majority of the diet in summer and fall, *viz.* *Acanthopanax evodiaefolius*, *Sorbus* spp., *Sorbus cf. thibetica*, and *Cornus macrophylla*. Of *Sorbus* and *Acanthopanax*, the monkeys ate both fruits and mature leaves at the same time, often in an alternating fashion. Mature leaves contributed relatively marginally to the diet of *R. bieti* at Samage (4%; Grueter *et al.* 2009). Most of the ingested mature leaves (most notably *A. evodiaefolius*, *Sorbus* spp., *Padus obtusata*, *Hydrangea heteromalla*, *Philadelphus delavayi*, *Fargesia* spp., and *Ilex* spp.) were deciduous with the exception of *Ilex* spp. and *Fargesia* sp. Sayers and Norconk (2008) demonstrated a similar preference for broad-leaved deciduous mature leaves over evergreen mature leaves in Himalayan langurs. The digestibility of short-lived deciduous leaves is almost universally superior to that of the evergreen leaves (Coley 1988). *Rhinopithecus bieti* at Samage also displayed a preference for young over mature leaves (Grueter *et al.* 2009). Compared to mature foliage, young foliage generally has higher nutritional quality; it is higher in protein, lower in fiber and secondary compounds (Boonratana 1993; McKey *et al.* 1981; Milton 1979; Oates *et al.* 1980).

Intra- and Interspecific Differences

There has been some discrepancy with regard to the typical feeding strategy of *Rhinopithecus bieti*. Kirkpatrick (1996) stressed the species' relatively monotonous dependence on lichens (specialist) whereas Ding and Zhao (2004) accentuated its dietary diversity (generalist). This inconsistency is likely a consequence of different habitat characteristics. Populations of *Rhinopithecus bieti* occur in different ecological conditions, and findings from the Wuyapiya population, which inhabits one extreme of the habitat of *R. bieti* (Kirkpatrick 1996), are not representative of all other populations. In a gradient from south to north, precipitation and temperature decrease while average altitude of occupied habitat increases and the vegetation becomes progressively poorer. The limited diet of the Wuyapiya and also Xiaochangdu population (Xiang *et al.* 2007) in the north is a consequence of low species richness. The Samage forest is floristically richer and more productive than Xiaochangdu and Wuyapiya and foraging options are thus less constrained. Our study confirms that the dietary spectrum and key foods largely depend on the particular habitat, climate condition, botanical composition, and species richness. The number of species consumed clearly decreases with increasing altitude and latitude. At the northern end of the geographical range of *Rhinopithecus bieti* (Wuyapiya and Xiaochangdu), they consumed 20 and 25 plant species in 12 and 13 mo, respectively (Kirkpatrick 1996; Xiang *et al.* 2007). As for the central part of the species' range, the group at Tacheng-Xiagguqing fed upon 50 plant species over 14

mo of study while the group at Samage-Gehuaqing ate 75 plant species over the first 14 mo of study and 94 species over the total of 20 mo (Ding and Zhao 2004; this study). At the southern end (Longma), they incorporated an assumed 97 species into the diet (data derived from indirect evidence such as trail signs only; Huo 2005). Diet breadth (species richness) of the Samage and Longma populations is comparable to that of most tropical sites where researchers have studied Asian colobines (Curtin 1980; Davies 1991; Kool 1993).

The top ranking dietary genus of *Rhinopithecus bieti* at Samage was *Acanthopanax* (alternatively named *Eleutherococcus*), of which the monkeys consumed all major phytophases, viz. young leaves in spring; mature leaves in summer and fall; fruits in summer, fall, and winter; buds in winter; and bark all year round. *Acanthopanax* was not part of the diet of *Rhinopithecus bieti* at Wuyapiya and Xiaochangdu, probably due to the genus' low density there (Kirkpatrick 1996), but it is harvested by populations of *R. bieti* living to the south of Samage (Liu *et al.* 2004). The closely related allopatric and ecologically comparable *Rhinopithecus roxellana* has a similar diet, both in terms of food plant genera and species (Guo *et al.* 2007; Li 2006). Among the non-*Rhinopithecus* taxa, Himalayan langurs of Nepal are of great comparative value owing to their association with biogeographically, botanically, and topographically similar habitats. While *Acanthopanax* is a key food plant species for *Rhinopithecus bieti*, Himalayan langurs did not include *Acanthopanax* at all into the diet despite the genus being relatively common there (Sayers and Norconk 2008). Conversely, the Himalayan langurs ate the shrub *Gaultheria* in considerable quantities, and despite being superabundant at Samage, evidence for inclusion in the diet of Himalayan snub-nosed monkeys is lacking.

What Do These Data Tell Us About the Possibility of Food Competition?

We demonstrated for this species and population that preferred food trees were spatially clumped, occurred at low densities across the home range, and were possibly not large enough to accommodate a fairly cohesive 400-member band at the same time. The distributional features are a prerequisite for contest competition (Koenig *et al.* 1998; van Schaik 1989; cf. Isbell *et al.* 1998). Researchers have so far not addressed for this population food quality, which also impacts the competitive potential (van Schaik 1989).

Some comparatively rare and highly preferred, and supposedly high-quality, plant resource patches, e.g., leaves of *Ilex* and *Philadelphus*, became depleted after the group of *Rhinopithecus bieti* had encountered them. Along the same lines, Kirkpatrick (1996) noted that ≥ 3 *Sorbus* trees were defoliated in the path of the band of *Rhinopithecus bieti*. *Ilex* spp. (holly) and *Philadelphus delavayi* (mock orange) occurred at very low stem densities, and also in terms of basal area, they all accounted for only 0.15% of the total basal area of all trees. These opportunistic data indicate that *Rhinopithecus bieti* undeniably depleted some plant resources, although full depletion was mostly restricted to a few scarce species.

Some authors have proposed that an efficient way to alleviate the costs of food competition is to avoid cofeeding (Saj and Sicotte 2007b; Siex and Struhsaker 1999; van Noordwijk and van Schaik 1987) by spreading out while feeding and consequently having fewer neighbors while feeding versus resting. Our results

confirm that feeding individuals had fewer conspecifics in proximity than resting ones. However, we never observed a female behaving competitively toward other female unit members while feeding in a patch or supplanting another female from a food source, behaviors that would indicate the presence of contest competition.

While competition within units was perhaps more of the scramble type, the low density and clumped distribution of preferred resources, e.g., fruiting trees, waterholes, are suggestive of interunit contest competition. We previously showed that male aggressive behavior correlates positively with temporal availability of fruit (Grueter 2009), which, however, could have been confounded by some other factor. Further, the only recorded band fission event occurred in late winter when valued fruit resources became rare and extremely patchy (Grueter *et al.* 2008a, b). *Ad libitum* observations substantiate the possibility of contest competition among units: in May 2006, large, dominant OMUs appeared to defend leafing trees (rare *Pterocarya* trees) from other nearby units. The lower ranking units appeared to wait in nearby conifer trees eating lichens until the more dominant units left the leafing trees. In January 2007, 1 unit chased away another unit from an *Acanthopanax* tree that still bore fruits. These observations recall Isbell's (1991) proposition that species that feed on both dispersed and clumped resources might reduce competition by shifting from clumped to dispersed foods. Researchers have also reported competition among social units for access to food trees/feeding areas for *Rhinopithecus roxellana* (Zhang *et al.* 2003), *Colobus guereza* (Harris 2006), and *C. polykomos* (Korstjens 2001).

Although still preliminary, our results demonstrate the pervasiveness of moderate food competition in *Rhinopithecus bieti*, at least with regard to nonlichen foods (plants). We would expect this to have a constraining effect on the species' socioecology, i.e., limit group size. However, groups are unusually large, which implies that either feeding competition is not severe enough to constrain group size, or that there is an advantage of forming large groups that counteracts the disadvantage associated with feeding competition, such as threats stemming from nearby roaming all-male units (Grueter 2009; Grueter *et al.* 2008b).

Conclusion and Areas for Future Research

Owing to earlier observations suggesting that *Rhinopithecus bieti* feed almost exclusively on lichens, researchers originally characterized the species as having one of the most specialized diets of all primates. Subsequent studies including this one have confirmed that lichens are indeed a key dietary component, but also underpinned the dietary elasticity of *Rhinopithecus bieti* in response to variation in availability, abundance, and diversity of plant food supply. *Rhinopithecus bieti* at Samage exhibit a rather broad usage of the resources in their habitat (>90 plant species) and thus can be viewed as generalists. However, the dominant evergreen plant families did not offer many palatable foodstuffs to the monkeys, which instead relied heavily on a few rather uncommon deciduous hardwood species. Some highly sought food trees occurred at very low densities and were irregularly distributed in space and time, which are preconditions for the emergence of contest competition. In line with recent studies, our study provides preliminary evidence consistent with the recent contention that feeding competition may be more widespread among

colobines than previously thought. Future studies should aim to obtain a better understanding of food competition in the species, for instance by gathering data on patch residence time, patch size, and unit size to evaluate patch depletion, a measurable behavioral indicator of the presence or absence of intragroup scramble competition (Snaith and Chapman 2005).

A salient finding of our study is that some mature leaves were totally avoided by *Rhinopithecus bieti* whereas others were highly sought after. It is widely known that protein content and fiber have a strong influence on leaf choice in colobines (Davies *et al.* 1988; Fashing *et al.* 2007; Oates *et al.* 1980; Waterman and Choo 1981). Future research should investigate the phytochemical components associated with preferred and avoided food items, thereby contributing to a fuller comprehension of the feeding ecology of *Rhinopithecus bieti*.

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