



## A Multi-Forest Comparison of Dietary Preferences and Seed Dispersal by *Ateles* spp.

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Received December 10, 2003; revision February 19, 2004; accepted March 24, 2004

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*Investigations of coevolutionary relationships between plants and the animals that disperse their seeds suggest that disperser-plant interactions are likely shaped by diffuse, rather than species-to-species, coevolution. We studied the role of dietary plasticity in shaping the potential for diffuse coevolution by comparing dietary fruit preferences and seed dispersal by 3 species of spider monkeys (*Ateles* spp.) in 4 moist forests in Colombia, Ecuador, Panama, and Surinam. In all forests, spider monkeys were highly frugivorous and preyed upon seeds of few species. We estimated dietary use of fruiting taxa based on absolute consumption and preference, which accounts for resource availability. Of the 59 genera that comprised the 20 most frequently consumed genera summed in each forest, only 3—*Brosimum* (*Moraceae*), *Cecropia* (*Cecropiaceae*) and *Virola* (*Myristicaceae*)—ranked within the top 20 at every forest. Most genera were within the 20 most frequently consumed at only 1 or 2 forests. Based on preferences, only 4 genera ranked in the 20 most-preferred in all 4 forests: *Brosimum*, *Cecropia*, *Ficus* (*Moraceae*), and *Virola*. Patterns in fruit consumption and preference at the familial level were similar in that only 2 families—*Myristicaceae* and *Moraceae*—were in the 10 most-consumed or most-preferred in all 4 forests. Interforest variation in plant specific composition and abundances and supra-annual fruiting*

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*phenologies, combined with dietary flexibility of Ateles spp., may partly explain these patterns. Our results suggest that variation in plant community structure strongly influences dietary preferences, and hence, seed dispersal by spider monkeys. Thus, diffuse coevolution in spider monkey-plant relationships may be limited to few taxa at the generic and familial levels.*

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**KEY WORDS:** *Ateles*; plant-disperser interactions; coevolution; frugivory; seed dispersal.

## INTRODUCTION

Investigating coevolutionary relationships between plants and the animals that disperse their seeds has been an important focus of seed dispersal biology, especially in tropical systems (Snow, 1971; McKey, 1975). Tiffney (1984), and Tiffney and Mazer (1995) linked increases in seed size of tree species and the diversification of large-seeded angiosperm species to the evolution of large-bodied frugivores that could disperse such large seeds. Likewise, Sussman (1991) proposed that increases in the availability of food resources displayed on terminal branches of angiosperms (fruits) led to the morphological adaptations of some primates to an arboreal life and contributed to primate diversification.

However, coevolution in terms of reciprocal evolutionary change among pairs of species, or pairwise coevolution (Janzen, 1980; Thompson, 1994; Futuyma, 1998), among partners in seed dispersal systems, has rarely been demonstrated (Wheelwright and Orians, 1982; Howe, 1984; Herrera, 1985). Weak selective pressures between dispersers and plants, spatial and temporal unpredictability of favorable germination sites for dispersed seeds, and long generation times of woody plants relative to their dispersers, all reduce the likelihood for tight, pairwise coevolutionary relationships to develop (Wheelwright and Orians, 1982; Howe, 1984; Herrera, 1985). In particular, plastic foraging ecologies of dispersers may prevent pairwise coevolution if composition and abundances of fruit resources in forests change rapidly in space and time, as empirical data suggest is often the case (Delcourt and Delcourt, 1987; Campbell, 1994; Colinvaux *et al.*, 1996; Pitman *et al.*, 2001; Condit *et al.*, 2002).

Most evidence points toward disperser-plant relationships being shaped by diffuse, rather than pairwise, coevolution (Howe and Smallwood, 1982; Herrera, 1985). Diffuse coevolution occurs when either or both interacting populations are represented by an array of taxa that generate selective pressure as a group (Janzen, 1980; Thompson, 1994; Futuyma, 1998). In the case of seed dispersal, arrays of taxa involved in diffuse coevolution may be exemplified by groups of dispersal agents or fruiting plants at taxonomic levels above that of species. Evidence for diffuse coevolution comes from

studies describing dispersal syndromes in plants, i.e., suites of plant traits, such as fruit color, size, and hardness, that may have evolved in a coordinated fashion in response to frugivory by broad classes of dispersal agents (van der Pijl, 1982). For example, in the Neotropics, fruits adapted for primate dispersal often may be large, yellow-orange, brown, or green with a woody pericarp (Janson, 1983). However, African primates, tend to prefer red, in addition to yellow-orange, fruits and to avoid brown and green fruits (Gautier-Hion *et al.*, 1985). These differences between New and Old World primates may be related to their dichromatic versus trichromatic vision, respectively (SurrIDGE *et al.*, 2003). Nonetheless, our understanding of the functional mechanisms behind primate-plant seed dispersal relationships is incomplete. Furthermore, data have been relatively unavailable to analyze variation in plant-disperser relationships across large spatial scales, an effort that is fundamental to understanding the role of dietary plasticity in shaping the potential for coevolution.

Primates in general (Chapman and Onderdonk, 1998) and spider monkeys (*Ateles* spp.) in particular (Andresen, 1999) are important frugivores and seed dispersers in tropical forests and therefore may have developed strong interactions with particular plant taxa. We assembled data on the role of spider monkeys as seed dispersers in moist forests in Colombia, Ecuador, Panama, and Surinam. We evaluated inter forest and inter year variation in their frugivory and seed predation to assess the potential for diffuse coevolutionary relationships between spider monkeys and the plants for which they disperse seeds.

## STUDY SITES AND SPECIES

The 4 study sites (Table I) are moist forests, with varying degrees of seasonality ranging from the aseasonal forest at Yasuni, Ecuador, to the strongly seasonal forest on Barro Colorado Island (BCI), Panama. Two sites were relatively protected from human exploitation before and during the study periods. On BCI, spider monkeys were extirpated early in the 20th century and re-introduced in the 1960's (Milton, 1993). The spider monkey population at Yasuni has recently experienced moderate hunting pressure, losing *ca.* 10 females in a 2-year period (Suarez, 2003). Detailed descriptions of study sites were published by van Roosmalen (1985b) for Voltzberg, Stevenson (2002) for Tinigua, DiFiore and Rodman (2001) for Yasuni, and Leigh (1999) for Barro Colorado Island (BCI).

The different *Ateles* spp. have similar diets, social organization, and behavior, making them suitable for interspecific comparisons of diet in the context of understanding diffuse coevolution. Spider monkeys are large

**Table 1.** Study sites, species, and field methods used in comparison of diet and seed dispersal of *Ateles* spp.

	Voltzberg, Surinam	Tinigua, Colombia	Yasuni, Ecuador	BCI, Panama
Forest type	Seasonal, moist	Seasonal, moist	Asesonal, moist	Seasonal, moist
Species	<i>A. paniscus</i>	<i>A. belzebuth</i>	<i>A. belzebuth</i>	<i>A. Geoffroyi</i>
Population density (no. individuals/km <sup>2</sup> )	6.3	24.0	11.5 <sup>a</sup>	2.3
No. observation h	865	672	457 <sup>a</sup> and 1,268 <sup>b</sup>	1,200
Diet sampling method	Scan	Time	Scan <sup>a</sup> and time <sup>b</sup>	Scan
Data source: diet	van Roosmalen (1985b)	Stevenson (2000; 2002)	Dew (2001) Suarez (2003)	Campbell (2000)
Plant sampling method (total area)	1 plot (335 ha)	7 1-ha plots (7 ha)	5 1-ha transects (5 ha)	1 plot (50 ha)
Data source: plants	van Roosmalen (1985b)	Stevenson (unpublished data)	Di Fiore (unpublished data)	Condit <i>et al.</i> , (1996)

<sup>a</sup>Dew (2001).

<sup>b</sup>Suarez (2003).

(5–8 kg) arboreal primates that primarily occupy the canopy strata of neotropical forests (Hershkovitz, 1978). Groups usually consist of 25–40 individuals and are organized in a fission-fusion social structure (Hershkovitz, 1978; McFarland, 1986). Spider monkeys are highly frugivorous, but also eat leaves, flowers, and occasionally insects in quantities that vary seasonally (Klein and Klein, 1977; Chapman, 1987; Symington, 1987).

## METHODS

We gathered data on feeding ecology of spider monkeys and the abundances of fruiting tree species each site for 12 mo (Table I). Details of the field sampling methodologies are in the data source publications cited in Table I. We estimated absolute consumption of fruiting taxa at each site as either the percent of time (Tinigua and Yasuni) or scans (all other sites) spent by all focal individuals feeding on each species, summed at the generic and familial levels (hereafter, percent feeding). We estimated the abundances of fruit-producing species in spider monkey diets at each site based on tree census plots (Table I). Census plots varied in total area sampled and sampling method (Table I), making it inappropriate to make detailed quantitative comparisons among sites of the abundance of fruit resources consumed by spider monkeys. Therefore, we based estimates of similarity among sites in fruit resources on presence/absence and on ranks of abundances for taxa at multiple sites.

Data on fruit production was not available for all sites. Therefore, we estimated the availability at each site of species producing fleshy fruits consumed by spider monkeys as the percent of the total number of stems (>10 cm diameter at breast height) of the species in the site's census plot. Then we summed the percentages at the generic and familial levels (hereafter, percent availability).

We estimated dietary use of fruiting taxa based on both absolute consumption (percent feeding) and preference. Preference is based on absolute consumption, but takes into account the availability of the fruit resource. We calculated the preference index for each fruiting taxon in the diet as the difference between absolute consumption and availability (percent feeding minus percent availability). We standardized preference indices within sites so that they ranged from  $-1$  to  $1$ , where in  $-1$  indicates strongest avoidance,  $1$  indicates strongest preference, and zero indicates no preference (consumed according to abundance; Krebs, 1999).

We did not calculate preference indices in 2 cases: (1) for liana species, because only trees were censused in plots and (2) when there was no availability estimate for tree taxa. In both cases, we coded preference indices for the taxa as missing data and excluded them from analyses. Dew (2001), identified fruiting taxa generically only so we estimated availability for each genus by summing availability for all species in the genus present at the study site. For all other data sets, we summed only species in each genus that were included in the diet to estimate availability for the genus. In all data sets, we estimated availability at the familial level by summing all genera consumed at a site in each family. Dew (2001) and Suarez (2003), identified Lauraceae species only to family so we excluded their data on Lauraceae from our analyses.

We consider seeds to have been preyed upon if they were ingested, but were either not defecated intact or masticated. We also recorded seed species that the monkeys dropped under the parent tree.

## RESULTS

Spider monkeys were highly frugivorous in all 4 forests (Table II). They ingested fruits and seeds of a large number of species in each forest and preyed upon seeds of few species (Fig. 1, Table II). On average, 3.2% of feeding observations involved seed predation (range 1.1–6.5%).

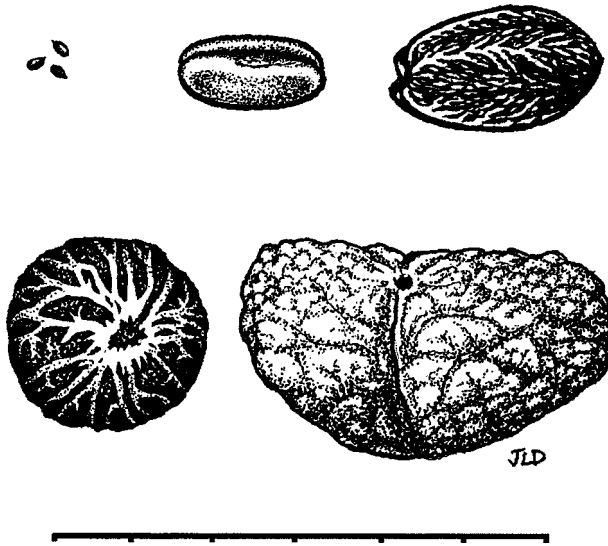
Substantial congruence existed among the 4 forests in terms of the presence of tree genera producing fleshy fruits that were important in spider monkey diets. Across all forests, 59 genera comprised the 20 fruit-producing tree genera most frequently consumed by spider monkeys. Among pairs of forests, 54–80% of the 59 genera occurred in both forests in each pair

**Table II.** Summary of the diets of 3 *Ateles* spp. from 4 neotropical forests

	Voltzberg, Surinam	Tinigua, Colombia	Yasuni, Ecuador <sup>a</sup>	Yasuni, Ecuador <sup>b</sup>	BCI, Panama
Species	<i>A. paniscus</i>	<i>A. belzebuth</i>	<i>A. belzebuth</i>	<i>A. belzebuth</i>	<i>A. geoffroyi</i>
Fruit (%)	82.9	74.0	87.0	78.8	82.2
Leaves (%)	6.4	12.0	9.0	7.7	17.2
Flowers (%)	7.9	5.0	1.0	1.3	1.0
Other (%)	2.7	9.0	0.7	12.2	0.6
Unidentified (%)	0	0	3.0	0	0.3
No. fruit spp. eaten	160	106	71	238	107
Seed predation (% species)	6.5	2.0	2.8	0.8	1.0

<sup>a</sup>Dew (2001).

<sup>b</sup>Suarez (2003).



**Fig. 1.** Seeds of trees dispersed by spider monkeys. Top row; left to right: *Cecropia*; *Inga*; *Spondias*, Bottom row; left to right: *Iriartea*; *Iryanthera*. Segments in scale bar correspond to 1 cm. Illustration by J. L. Dew.

(Table III). In terms of taxonomic composition, the least similar forests were Tinigua and Voltzberg, and the most similar forests were BCI and Yasuni (Table III). However, ranks of the abundance of fruit-producing genera are only significantly correlated between Voltzberg and Tinigua (Table IV). Hence, although they were dissimilar in terms of taxonomic composition (Table III), the abundances of the genera that occurred in both forests were congruent. Conversely, although BCI and Yasuni were most similar in terms of composition (Table III), the abundances of the genera present at both sites were not similar (Table IV).

Despite similarity among the 4 forests in the genera frequently consumed by spider monkeys, their diets were plastic. Few genera were

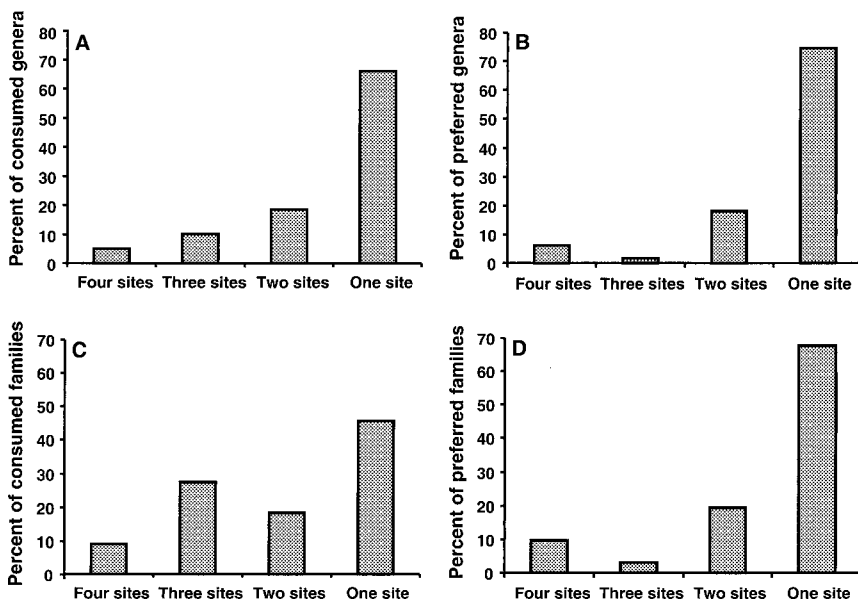
**Table III.** Percent of 59 genera producing fleshy fruits comprising the 20 most-consumed by spider monkeys that were present at both forests in each pair

	Voltzberg	Tinigua	Yasuni
Tinigua	54		
Yasuni	59	78	
Barro Colorado Island	56	71	80

**Table IV.** Spearman rank correlation coefficients (probability, sample size) for the association between abundances of fruit-producing genera consumed by spider monkeys in forest pairs. The correlation between Voltzberg and Tinigua remained significant after correcting for multiple comparisons using the Bonferroni method (Sokal and Rohlf, 1995).

	Voltzberg	Tinigua	Yasuni
Tinigua	0.53 ( $p < 0.01$ , $N = 26$ )		
Yasuni	-0.05 ( $p = 0.81$ , $N = 31$ )	0.29 ( $p = 0.12$ , $N = 33$ )	
Barro Colorado Island	0.32 ( $p = 0.21$ , $N = 17$ )	0.39 ( $p = 0.12$ , $N = 17$ )	-0.12 ( $p = 0.77$ , $N = 20$ )

important in their diets in multiple forests, either in terms of absolute consumption or in terms of preference, i.e., after accounting for the availability of the fruit resource (Fig. 2A, B). Based on absolute consumption, of the 59 genera that comprised the 20 most-consumed genera from each forest, only 3 ranked in the top 20 in all 4 forests: *Brosimum*, *Cecropia*,



**Fig. 2.** Percents of fruit-producing genera (A and B) or families (C and D) that were important in spider monkey diets in 4 neotropical forests either in terms of absolute consumption (A and C) or after the availability of the fruit resource was taken into account (B and D). (A) Percent of 20 most-consumed genera summed over all forests (59 genera) that ranked in the top 20 genera in 4, 3, 2, and 1 forest. (B) Percent of 20 most-preferred genera summed over all forests (56 genera) that ranked in the top 20 genera in 4, 3, 2, and 1 forest. (C) Percent of the 10 most-consumed families summed over all forests (20 families) that ranked in the top 10 families in 4, 3, 2, and 1 forest. (D) Percent of the 10 most-preferred families summed over all forests (27 families) that ranked in the top 10 families in 4, 3, 2, and 1 forest.



**Table V.** The 10 most-consumed genera and families, numbers of species, and percent feeding by spider monkeys in four neotropical forests

Genus	No. species eaten	Percent feeding	Family	No. species eaten	Percent feeding
<b>Voltzberg, Surinam</b>					
<i>Viola</i> (Myristicaceae)	2	12.5	Fabaceae	21	15.2
<i>Inga</i> (Fabaceae)	12	10.8	Myristicaceae	2	12.5
<i>Guarea</i> (Meliaceae)	2	4.9	Sapotaceae	7	9.7
<i>Tetragastris</i> (Bursaceae)	2	4.2	Moraceae	19	7.9
<i>Ecclinusa</i> (Sapotaceae)	1	3.8	Meliaceae	6	6.7
<i>Cecropia</i> (Cecropiaceae)	2	3.7	Bursaceae	4	6.3
<i>Dimorphandra</i> (Fabaceae)	1	3.6	Cecropiaceae	8	5.4
<i>Philodendron</i> (Araceae)	2	3.2	Annonaceae	5	4.0
<i>Bagassa</i> (Moraceae)	1	3.2	Araceae	4	3.4
<i>Achrouteria</i> (Sapotaceae)	1	3.0	Flacourtiaceae	1	2.5
<b>Tinigua, Colombia</b>					
<i>Ficus</i> (Moraceae)	8	15.4	Moraceae	21	25.5
<i>Oenocarpus</i> (Arecaceae)	2	9.6	Arecaceae	7	14.0
<i>Viola</i> (Myristicaceae)	3	7.3	Cecropiaceae	8	10.0
<i>Gustavia</i> (Lecythydaceae)	1	6.9	Bursaceae	7	9.1
<i>Protium</i> (Bursaceae)	4	5.8	Myristicaceae	5	7.5
<i>Sarcaulus</i> (Sapotaceae)	1	5.5	Lecythydaceae	1	6.9
<i>Brosimum</i> (Moraceae)	5	5.2	Sapotaceae	7	6.5
<i>Pourouma</i> (Cecropiaceae)	4	4.0	Fabaceae	8	3.0
<i>Pseudolmedia</i> (Moraceae)	3	3.8	Anacardiaceae	3	2.7
<i>Cecropia</i> (Cecropiaceae)	2	3.5	Meliaceae	2	2.2
<b>Yasuni, Ecuador<sup>a</sup></b>					
<i>Spondias</i> (Anacardiaceae)	—	21.5	Anacardiaceae	—	21.5
<i>Viola</i> (Myristicaceae)	—	14.6	Myristicaceae	—	18.7
<i>Iriarte</i> (Arecaceae)	—	9.0	Arecaceae	—	14.0
<i>Persea</i> (Lauraceae)	—	8.8	Lauraceae	—	8.9
<i>Matisia</i> (Bombacaceae)	—	7.7	Bombacaceae	—	8.0
<i>Guarea</i> (Meliaceae)	—	6.3	Meliaceae	—	6.9
<i>Oenocarpus</i> (Arecaceae)	—	4.9	Cecropiaceae	—	6.2
<i>Cecropia</i> (Cecropiaceae)	—	4.2	Moraceae	—	4.0
<i>Iryanthera</i> (Myristicaceae)	—	4.2	Fabaceae	—	2.4
<i>Naucleopsis</i> (Moraceae)	—	2.7	Sapindaceae	—	1.5
<b>Yasuni, Ecuador<sup>b</sup></b>					
<i>Viola</i> (Myristicaceae)	8	12.3	Moraceae	22	19.0
<i>Coccoloba</i> (Polygonaceae)	3	9.0	Myristicaceae	10	15.8
<i>Pseudolmedia</i> (Moraceae)	4	5.3	Polygonaceae	4	9.0
<i>Hyeronima</i> (Euphorbiaceae)	1	5.1	Euphorbiaceae	3	5.2
<i>Brosimum</i> (Moraceae)	3	4.4	Anacardiaceae	2	4.3
<i>Naucleopsis</i> (Moraceae)	2	4.4	Bombacaceae	3	4.2
<i>Matisia</i> (Bombacaceae)	3	4.2	Ulmaceae	2	3.7
<i>Ampelocera</i> (Ulmaceae)	1	3.7	Fabaceae	12	3.4
<i>Iryanthera</i> (Myristicaceae)	1	3.5	Arecaceae	5	3.3
<i>Tapira</i> (Anacardiaceae)	1	3.4	Malpighiaceae	1	3.3
<b>Barro Colorado Island, Panama</b>					
<i>Quararibea</i> (Bombacaceae)	1	12.4	Bombacaceae	1	12.4
<i>Spondias</i> (Anacardiaceae)	2	10.3	Moraceae	9	9.2
<i>Viola</i> (Myristicaceae)	1	8.5	Anacardiaceae	3	10.4
<i>Hyeronima</i> (Euphorbiaceae)	1	6.6	Myristicaceae	1	8.5
<i>Doliocarpus</i> (Dilleniaceae)	2	5.6	Euphorbiaceae	3	7.3

Table V. Continued

Genus	No. species eaten	Percent feeding	Family	No. species eaten	Percent feeding
<i>Ficus</i> (Moraceae)	6	4.8	Arecaceae	3	6.6
<i>Astrocaryum</i> (Arecaceae)	1	4.6	Rubiaceae	6	6.1
<i>Brosimum</i> (Moraceae)	1	4.3	Dilleniaceae	2	5.6
<i>Chrysophyllum</i> (Sapotaceae)	2	4.1	Burseraceae	3	5.3
<i>Tetragastris</i> (Burseraceae)	1	3.9	Sapotaceae	3	4.6

<sup>a</sup>Dew (2001).

<sup>b</sup>Suarez (2003).

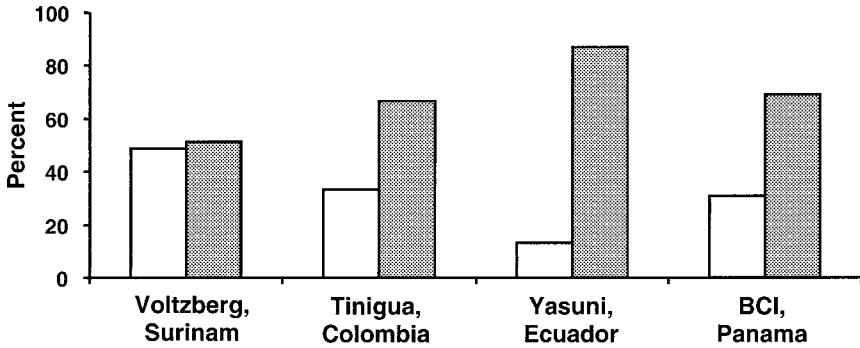
and *Viola* (Fig. 2A and Table V). After adjusting absolute consumption to account for fruit availability, 4 genera ranked in the 20 most-preferred genera in all 4 forests: *Brosimum*, *Cecropia*, *Ficus*, and *Viola* (Fig. 2B). The number of species in each genus varied somewhat among sites, ranging from only 1 species of *Brosimum* on BCI to 10 species of *Viola* and 10 species of *Ficus* at Yasuni and Voltzberg, respectively (Table VI).

As with genera, few families were important in spider monkey diets in multiple forests in terms of either absolute consumption or preference (Fig. 2C, D). Based on absolute consumption, of the 20 families comprising the 10 most-consumed families from each forest, only 2—Moraceae and Myristicaceae (Fig. 2C and Table V)—ranked in the top 10 in every forest. After adjusting absolute consumption to account for fruit availability, the same 2 families ranked in the 10 most-preferred families in every forest: Moraceae, and Myristicaceae (Fig. 2D and Table V). Few genera ranked in the 20 most-consumed or in the 20 most-preferred in 2 or 3 forests (Fig. 2A, B). However, at the familial level, a relatively greater proportion ranked within the 10 most-consumed families in 2 or 3 forests (Fig. 2C). Thus, which genera or families were important in spider monkey diets tended to be forest-specific (Fig. 2).

The importance of fruiting taxa in spider monkey diets may vary across forests due to differences in tree specific composition and abundance among them. Thirty-nine of the 59 genera comprising the 20

**Table VI.** Numbers of species in 4 genera that were important in spider monkey diets in 4 neotropical forests as ranked within the 20 most-preferred genera in all 4 forests

	Barro Colorado			
	Island	Voltzberg	Tinigua	Yasuni
<i>Brosimum</i>	1	2	6	4
<i>Cecropia</i>	2	2	2	2
<i>Ficus</i>	8	10	8	5
<i>Viola</i>	2	2	3	10



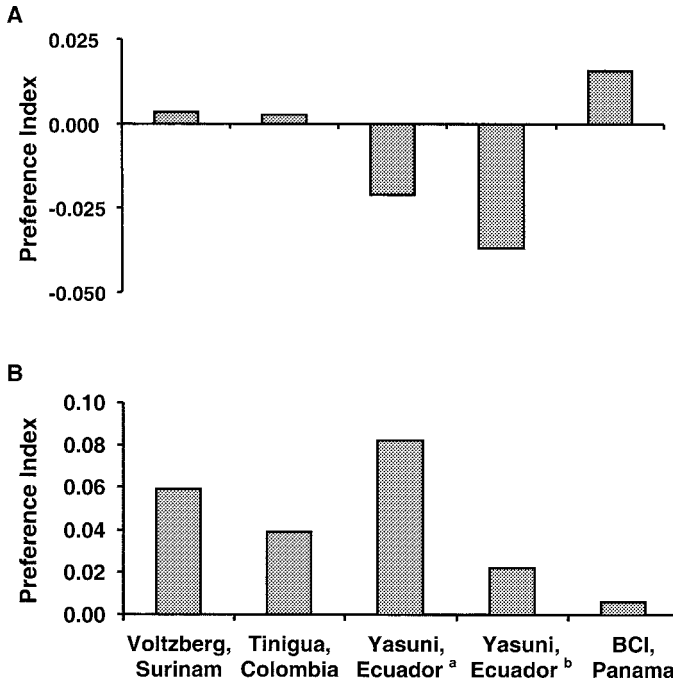
**Fig. 3.** Percent of fruit-producing genera that did not rank in the 20 most-consumed by spider monkeys in a forest and were either absent (open) or present (gray fill) at each forest. Details in text.

most-consumed genera did not rank in the top 20 in a particular forest. Of those genera, an average of 31% did not occur in that forest, and variation among forests in this quantity is related to variation in their floras (Fig. 3). Conversely, an average of 69% of the 39 genera were nonetheless present there (Fig. 3). Furthermore, preference indices for a genus are only weakly correlated between forests (Table VII). Only for BCI and Yasuni, the 2 most floristically similar forests in terms of tree genera producing fleshy fruits (Table III) is the preference index of each genus correlated (Table VII). Conversely, the 2 forests with greatest similarity in abundances of fruit-producing genera—Voltzberg and Tinigua, (Table IV)—did not have a significant correlation in preference indices (Table VII).

Spider monkeys did not exhibit an overall tendency for strong preference for or avoidance of fruiting genera in their diets in each forest. The mean preference index in each forest is near zero, suggesting that, on average, genera are consumed according to their availability (Fig. 4A).

**Table VII.** Spearman rank correlation coefficients (probability, sample size) for the association between spider monkey feeding preferences (preference indices) for each genus in forest pairs. We calculated preferences for Yasuni per both Dew (2001) and Suarez (2003) by averaging the preference index for a genus when it was consumed in both studies, or, for genera only consumed in one study, by using the preference index from that study. The correlation between Yasuni and Barro Colorado Island remained significant after correcting for multiple comparisons using the Bonferroni method (Sokal and Rohlf, 1995).

	Voltzberg	Tinigua	Yasuni
Tinigua	0.28 ( $p = 0.17, N = 26$ )		
Yasuni	0.13 ( $p = 0.48, N = 31$ )	0.27 ( $p = 0.13, N = 33$ )	
Barro Colorado Island	0.34 ( $p = 0.18, N = 17$ )	0.25 ( $p = 0.33, N = 17$ )	0.63 ( $p < 0.01, N = 20$ )



**Fig. 4.** Mean (A) and variance (B) of preferences indices for all genera consumed by spider monkeys at each forest. The 2 studies at Yasuni, Ecuador, are in separate sets of bars; a refers to Dew (2001), and b refers to Suarez (2003).

Instead, variation in preference indices is high, indicating that a few taxa are strongly preferred and a few others strongly avoided (Fig. 4B). Genera that ranked within the 20 most-preferred in  $\geq 2$  forests are *Astrocaryum* (Arecaceae), *Brosimum* (Moraceae), *Cecropia* (Cecropiaceae), *Coccoloba* (Polygonaceae), *Ficus* (Moraceae), *Guatteria* (Annonaceae), *Hyeronima* (Euphorbiaceae), *Spondias* (Anacardiaceae), and *Virola* (Myristicaceae). Genera that the monkeys avoided include ones with high population densities in individual forests, such as *Guarea* (Meliaceae), *Iriartea* (Arecaceae), *Mouriri* (Melastomataceae), *Protium* (Burseraceae), *Pseudolmedia* (Moraceae), *Tetragastris* (Burseraceae), and *Trichilia* (Meliaceae). The mean preference index is highest and variance lowest on BCI.

Dietary patterns of spider monkeys indicate their roles as seed dispersers because seeds of few species were preyed upon (Table II) or dropped below the parent tree (Table VIII). Among forests, there are some similarities as to which tree species seeds were preyed upon (Table VIII).

**Table VIII.** Seeds experiencing predation by spider monkeys in 4 neotropical forests. Unless otherwise noted by abbreviations following species names, seeds were either digested or unripe fruits were eaten. Abbreviations: m, seed masticated; d, seed dropped under parent

Species	Family
Voltzberg, Surinam	
<i>Tanaecium nocturnum</i>	Bignoneaceae
<i>Anomoctenium granulosum</i>	Bignoneaceae
<i>Pachira insignis</i>	Bombacaceae
<i>Cheiloclinium cognatum</i>	Celastraceae
<i>Combretum rotundifolium</i>	Combretaceae
<i>Maripa glabra</i>	Convolvulaceae
<i>Maripa scandens</i>	Convolvulaceae
<i>Operculina hamiltoni</i>	Convolvulaceae
<i>Alchorneopsis floribunda</i>	Euphorbeaceae
<i>Couratari</i> spp.	Lecythidaceae
<i>Eschweilera</i> spp.	Lecythidaceae
<i>Eperua</i> spp.	Fabaceae
<i>Cedrelinga cateniformis</i>	Fabaceae
<i>Dioclea macrocarpa</i>	Fabaceae
<i>Ormosia paraensis</i>	Fabaceae
<i>Carapa procera</i>	Meliaceae
<i>Cedrela odorata</i>	Meliaceae
<i>Paullinia sphaerocarpa</i>	Sapindaceae
<i>Priourella</i> sp.	Sapotaceae
<i>Pouteria</i> sp.	Sapotaceae
Tinigua, Colombia	
<i>Astrocaryum chambira</i> (m)	Arecaceae
<i>Brosimum</i> spp.	Moraceae
<i>Oenocarpus bataua</i>	Arecaceae
<i>Pseudolmedia</i> spp.	Moraceae
Yasuni, Ecuador <sup>a,b</sup>	
<i>Astrocaryum chambira</i>	Arecaceae
<i>Iriartea deltoidea</i>	Arecaceae
Barro Colorado Island, Panama	
<i>Astrocaryum standleyanum</i> (d)	Arecaceae
<i>Beilschmedia pendula</i> (d)	Lauraceae
<i>Brosimum alicastrum</i>	Moraceae
<i>Dipteryx panamensis</i> (d)	Leguminosae

<sup>a</sup>Dew (2001).

<sup>b</sup>Suarez (2003).

Spider monkeys preyed upon seeds of *Brosimum* and palm species in 2 and 3 forests, respectively (BCI and Tinigua, and BCI, Yasuni, and Tinigua, respectively). Most seed predation involved consumption of unripe fruit. They dropped seeds of very large-seeded species below the parent, and therefore did not disperse them, e.g., *Dipteryx*, *Beilschmedia*.

Data from Yasuni collected in 2 study years allowed an evaluation of the interaction between temporal variation in fruit availability and spider

monkey diets. There is a significant correlation between the preference indices of genera consumed in both studies (Spearman  $r = 0.64$ ,  $p = 0.001$ ); however, there is considerable variation in the relationship. Of the 31 genera comprising the 20 most-consumed genera in the 2 studies at Yasuni, only 30% ranked in the top 20 in both. Hence, spider monkey diets, as measured by the most frequently consumed fruiting species, varied substantially on an annual basis in the same forest, even when comparing only 2 years. In at least 2 cases—*Brosimum lactescens* and *Spondias mombin*—variation in preference was due to low fruit production in one study year and high production in the other (S. A. Suarez and J. L. Dew, personal observations).

## DISCUSSION

Spider monkey diets varied across forests and years in terms of both absolute consumption and preference, which is consistent with results of other comparative studies of primates (Chapman and Chapman, 2002b). Our results suggest that variation in composition and abundance of fruiting taxa among forests and years strongly influences foraging choices, and hence seed dispersal by spider monkeys. Spider monkeys tended to consume most taxa according to their availability, as has been observed in at least one other study (Nunes, 1998). Nonetheless, they strongly preferred some genera, and avoided others. Strong interactions between spider monkeys and fruiting taxa that were consistent across all forests are limited to few genera—*Brosimum*, *Cecropia*, *Ficus*, and *Virola*—and families: Moraceae and Myristicaceae. From the plant's perspective, such dietary plasticity means that the selective pressures of spider monkeys on plant and fruit traits affecting dispersal are not consistent in space, even among forests with broad floristic similarity. Thus, diffuse coevolution in spider monkey-plant relationships may be limited to few taxa at the generic and familial levels.

Coevolution between a plant and a particular dispersal agent depends on spatial and temporal consistency in their interaction, as well as the interactions the plant has with other species that may also be important dispersers (Herrera, 2002). Hence, it is important to consider whether the 4 genera with which spider monkeys appear to have consistent interactions are substantially dispersed by other dispersal agents. *Cecropia* and *Ficus* spp. are well-known to be consumed (and likely dispersed) by many frugivores (Tello, 2003; Fleming and Williams, 1990). They fit the paradigmatic generalist dispersal syndrome in having sugar-rich, small-seeded fruits and being prodigious fruit-producers (McKey, 1975). *Brosimum* spp. also attract a diversity of frugivores (S. E. Russo, personal observation). However,

seeds of *Brosimum* are larger than those of *Cecropia* and *Ficus* (van Roosmalen, 1985a), which inevitably excludes smaller dispersal agents. In addition, spider monkeys preyed upon seeds of *Brosimum* in at least 2 forests. The relative proportions of ingested seeds that were dispersed versus preyed upon is unknown, but would likely be an important influence on the fruit traits of at *Brosimum* experiencing selection by spider monkeys, as well as the nature of that selection.

Conversely, *Virola* spp. appear to be dispersed by a relatively narrower assemblage of species (Howe, 1981; Howe and Vande Kerckhove, 1981; Russo, 2003), perhaps in part because of the generally larger seed sizes and the oily, rather than sugary, aril characteristic of the genus (van Roosmalen, 1985a). Hence, we predict that of the 4 genera, *Virola* would experience the strongest selection on tree and fruit traits as a result of seed dispersal by spider monkeys. However, a detailed analysis of this possibility in *Virola calophylla* in Perú showed that divergent selection from even a limited assemblage of other dispersal agents may counter any selection from spider monkeys (Russo, 2003). Thus, further research is necessary to understand responses to selection, but it appears that even consistent, strong interactions between dispersal agents and fruit-producing tree species may not be sufficient for dispersers to shape the traits of the plants they disperse.

Variation in spider monkey dietary consumption and preferences for fruiting taxa among forests may be explained by extrinsic and intrinsic factors. First, variation in both the composition and abundance of fruiting taxa is likely a strong influence on spider monkey foraging decisions. Three results support this conclusion. First, the 2 forests that shared the greatest number of genera most-frequently consumed by spider monkeys (BCI and Yasuni) showed the greatest correlation in preference indices, which suggests that taxonomic composition of fruiting taxa in a forest plays an important role in spider monkey foraging choices. Second, we also found that of the 39 genera comprising the 20 most-consumed genera that were not ranked in the top 20 in a particular forest, most were nonetheless still present in that forest, which suggests that a substantial amount of the variation among forests in spider monkey diets is due to foraging choices among fruiting taxa present and their abundances, rather than the absence of fruiting taxa. Furthermore, the fact that the preference indices for a genus were only weakly correlated between forests is consistent with this interpretation. Third, the 2 forests with the most correlated abundances of genera (Voltzberg and Tinigua) do not have significantly correlated preference indices. This result must be considered with caution because the tree abundance data were not designed for across-site comparisons. However, we can tentatively conclude that abundances of fruiting genera alone do not

determine spider monkey foraging decisions. Taken together, our analyses suggest that both taxonomic composition and abundances of fruiting taxa in forests strongly influence spider monkey foraging choices and, hence, seed dispersal.

Second, interspecific variation in feeding among species of *Ateles* may explain variation among forests in fruit consumption and preferences. If this were the case, then forests with the same species of *Ateles* should have the greatest correlation between preference indices, all else being equal. However, the correlation in preference indices between forests is not highest for Tinigua and Yasuni, the only forests that share the same species of *Ateles*. Given that current taxonomy reflects phylogeny, this result suggests that phylogeny may not be a dominant factor determining the patterns of variation in diet that we observed in *Ateles*.

Third, variation among forests in specific composition and interspecific interactions among sympatric frugivorous primates is also likely to influence spider monkey foraging choices. Studies of diets of frugivorous primates suggest that species partition fruit resources based on fruit characteristics (Kinzey and Norconk, 1990; Ungar, 1995; Dew, 2001). BCI, which has the lowest population density of spider monkeys and the fewest species of sympatric, frugivorous primates, had the highest mean preference index and lowest variance. In addition, the single group on BCI has an extremely large home range (Campbell, 2000). A potential explanation for these patterns is that spider monkeys on BCI may be less constrained by intra- and interspecific competition in their foraging choices, and therefore may be freer to use preferred fruit resources.

Fourth, the comparison between 2 study years at Yasuni indicated the importance of accounting for supra-annual fruiting phenology and the availability of fruit resources in understanding foraging choices of spider monkeys. Similarly, at least 1 genus on BCI (*Faramea*) had a very low preference index in the single year of data collection there, when the population of *Faramea* produced little fruit. However, in other years when *Faramea* fruited prodigiously on BCI, spider monkeys foraged extensively on this genus (C. J. Campbell, personal observation). More sampling years in all forests would probably reveal a greater number of genera and families ranking as important dietary components in all forests. However, the fact that *Brosimum*, *Cecropia*, and *Virola* are important in all forests, even in one year of sampling, suggests that the spider monkeys have strong relationships with them. In particular, in all forests, *Virola* is within the top 3 genera having the greatest consumption, followed by a large drop in consumption rates for lower-ranked genera (Table V). *Virola* is also important in the diets of spider monkeys in Manú National Park, Perú, where 92% of dispersed seeds of *Virola calophylla* were dispersed by spider monkeys (Russo, 2003).



Finally, primate foraging choices are influenced by many factors, including caloric, nutrient, and secondary chemical contents of resources, the abundance, spatial distribution, and temporal predictability of resources, and competition (Clutton-brock, 1977; Hladik, 1977; Terborgh, 1983; van Schaik *et al.*, 1993; Kaplin *et al.*, 1998; Chapman and Chapman, 2002a). Several factors may contribute to consistent preferences by spider monkeys for *Brosimum*, *Cecropia*, *Virola*, and *Ficus*. *Brosimum* may be preferred because of the generally large quantities of fruit produced and long periods of fruit availability on individual trees. *Cecropia* may be preferred because of its extended fruiting phenology, as has been suggested in a comparative study of the diets of avian dispersal agents (Carlo *et al.*, 2003). Similarly, *Virola* may be preferred for reliable annual fruit production. *Virola* had consistent fruit production every year for 4 years in Manú National Park, Perú (S. E. Russo, unpublished data) and for 10 years on BCI (S. J. Wright, personal communication). In contrast, 3 important genera in spider monkey diets—*Brosimum*, *Clarisia*, and *Pseudolmedia*—produced no fruit in at least 1 of the 4 years in Manú National Park (S. E. Russo, unpublished data). *Ficus* may be preferred because it often fruits in seasons of resource scarcity (Terborgh, 1986; Ahumada *et al.*, 1998).

Our findings have important conservation implications. For plants, spider monkeys play a particularly vital role in dispersing a large number of tropical genera. Spider monkeys are one of the first animals to be hunted out of forests (Redford, 1992; Peres, 2000), and their loss is likely to have important consequences for demography, community structure, and gene flow of trees and lianas in tropical forests (Chapman and Chapman, 1995; Chapman and Onderdonk, 1998; Pacheco and Simonetti, 2000). For spider monkeys, dietary plasticity may enable them to vary foraging in response to loss of an important food resource, e.g., as a result of selective logging or changes in resource abundance as a result of climate change or mild disturbance (Chapman and Chapman, 1990).

Evaluating potential coevolutionary relationships among plants and their animal seed dispersers requires using a comparative approach that accounts for both the ecological patterns across multiple closely related species in a geographic context and the evolutionary trends in those patterns (Thompson, 1999). Our research constitutes one of the few comparative studies of diet and seed dispersal in primates across large geographic areas. Nonetheless, its limitations must be acknowledged. We collected feeding data for only one year in all but one forest, yet many neotropical fruiting species demonstrate supra-annual variation in fruit production, and diets of neotropical primates are flexible on both seasonal and annual bases (Chapman, 1987; Chapman *et al.*, 2002). Methods of collection of feeding and tree abundance data varied among forests, which could

confound interpretation of patterns (Stevenson and Quiñones, in press). Ideally, availability of fruits would be measured directly, rather than by way of the correlate of population density. We hope that future comparative data sets can be assembled to examine further which factors influence frugivory and seed dispersal by primates.

### ACKNOWLEDGMENTS

Fieldwork on BCI by Campbell was supported by grants from the National Science Foundation (SBR-9711161), the L. S. B. Leakey Foundation, the Department of Anthropology, University of California, Berkeley, and the Smithsonian Tropical Research Institute. Campbell also thanks Antigone Thomas, Dr. S. Joseph Wright, and Osvaldo Calderon for help in collecting data. Dew thanks the National Science Foundation, the Douroucouli Foundation, Primate Conservation Incorporated, Tony Di Fiore, Kristin Phillips, and Peter Rodman. Stevenson thanks Jorge Ahumada and Marcela Quiñones, who helped in data collection, and Banco de la Republica, Centro de Investigaciones Ecologicas La Macarena, and Colciencias for their support. Suarez thanks the L. S. B. Leakey Foundation, the National Science Foundation, Dr. Peter Rodman for financial support, and Gorky Villa, Nigel Pitman, Jill Anderson, Jacob Nabe-Nielson, Robyn Burnham, and Robin Foster for botanical assistance. Russo was supported by a Dissertation Completion Fellowship from the University of Illinois while writing this article and thanks Carol K. Augspurger for constructive input. We also thank 2 anonymous reviewers for comments that improved an earlier version of this article.

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