Diet and Activity Budget in *Colobus angolensis ruwenzorii* at Nabugabo, Uganda: Are They Energy Maximizers?

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Colobines · Diet composition · Food selection · Group cohesion · Fallback foods

**Abstract**

**Introduction:** Colobine monkeys are specialized folivores that use foregut fermentation to digest leaves. The slow process of fermentation forces them to spend a lot of time resting and to minimize their energy expenditure to subsist on a lower-quality diet. **Methods:** We recorded the diet and activity budget of *Colobus angolensis ruwenzorii*, which form a three-tiered multi-level society, at Lake Nabugabo, Uganda, over 12 months using scan sampling on adults and subadults, to determine whether they utilize the energy minimization strategy typical of colobines. **Results:** We found that the annual diet was primarily comprised of high-quality food resources (young leaves 65% and fruit 31%), and fruits were the only plant part the monkeys selected when available. Both the fruits and young leaves of some species were preferred food items in some months, and mature leaf consumption correlated negatively with preferred food availability. Mature leaves appear to be a fallback food for this population but are rarely relied upon (3%). The *C. a. ruwenzorii* at Nabugabo spent less time resting (40%) and more time moving (25%) than is typical for other species of black-and-white colobus. **Discussion/Conclusion:** The high-quality diet of this population appears to allow them to utilize an energy maximization strategy. Their reliance on food items that tend to be clumped in space and time likely explains the frequent fission-fusion behaviour that we observe between core units. Our findings demonstrate that the foraging strategies of colobines may be more flexible than was previously thought and illustrate how food availability and distribution can impact primate social organization.
how the need to meet nutritional requirements is balanced with other necessary activities and how foraging strategies impact social systems [e.g., Garber, 1987; Ganzhorn, 1988; Kappeler and van Schaik, 2002; Felton et al., 2009; Lambert and Rothman, 2015].

Primate foraging decisions are greatly dependent on the nutritional value (e.g., protein, carbohydrate/sugar and lipid/fat content) of the food items available [Kum-pan et al., 2019], as well as their physical and chemical defences. Physical defences make plants hard to process (e.g., thorns) or difficult to digest (e.g., fibre such as cellulose) [Waterman, 1984]. Chemical defences, also called secondary metabolites, can be distasteful (e.g., alkaloids), interfere with the efficiency of nutrient absorption and therefore inhibit digestibility (e.g., tannins), or be toxic (e.g., nicotine) [Griffiths, 1986; Waterman and Kool, 1994]. The concentration of secondary metabolites and the nutritional content both vary with the part of the plant that is being consumed, as well as its stage (e.g., ripeness or maturity) [Lambert and Rothman, 2015; reviewed by Garber, 1987]. For example, leaves tend to be a good source of protein [Ganzhorn et al., 2017] but also contain fibre and secondary metabolites, with the highest levels of these occurring in mature leaves [Waterman et al., 1980; Coley and Barone, 1996]. Fruits are high in sugar, particularly when they are ripe, seeds are a good source of protein and fat, and bark is primarily fibre [Lambert and Rothman, 2015]. Lichens are generally a poor source of protein, sugar and fat, but are used by some species as a seasonal fallback food [Liu et al., 2013; Grueter et al., 2009].

A number of foraging strategies have been proposed to explain what food items are selected and the quantity of each that are consumed [reviewed by Felton et al., 2009]. The foraging strategy a species utilizes has important consequences for the ways in which they allocate their time. Energy maximization strategies are often considered under optimal foraging theory and posit that individuals try to maximize their energy obtained per unit feeding time [Emlen, 1966; Schoener, 1971]. Primate species that are largely frugivorous are often thought to employ this strategy. Because fruits tend to occur in discrete clumps, frugivores typically spend more time moving and travel further between food patches, spend more time searching for and feeding on fruits, and spend comparatively little time resting [Rosenberger and Strier, 1989; Stier, 1992; Korstjens et al., 2010; Fashing, 2011]. Conversely, folivores are often thought to employ an energy minimization strategy [Dasilva, 1992; Wasserman and Chapman, 2003; Rangel-Negrín et al., 2018; Klass, 2020]. Many folivores have evolved specialized guts to optimize their digestion of leaves [Ankel-Simons, 2007]. In the case of colobines, this is a foregut fermentation system that allows them to neutralize some of the toxins present in their food and process energy from leaves high in fibre [Bau-chop and Martucci, 1968; Chivers, 1994; Kay and Davies, 1994]. However, the slow process of fermentation, and the associated problem of gut fill, results in high levels of forced resting time, necessitating an energy minimization strategy [Oates, 1977; Dasilva, 1992; Dunbar, 1992; Stier, 1992; Korstjens et al., 2010]. Consequently, highly folivo-rous species tend to spend more time resting than frugi-vores [Rosenberger and Stier, 1989; Stier, 1992; Korstjens et al., 2010; Fashing, 2011]. Furthermore, because leaves tend to be uniformly distributed, folivores are able to spend less time moving and searching for food [Isbell, 1991]. While these general patterns exist between folivores and frugivores, groups may adjust their time budgets and movement patterns, depending on the availability of different food items and their distribution [God-frey, 2003; Riley, 2007; Dunbar et al., 2009], with the same species showing an energy minimization strategy in one habitat and an energy maximization strategy in another [Alouatta cimitics, Jung et al., 2015; A. pigra, Rangel-Negrín et al., 2018; Klass, 2020; A. palliata mexicana, Asensio et al., 2007; Chlorocebus djamdjamensis, Mekon-nen et al., 2017].

The African colobines are largely considered to be folivorous, although the percentage of leaves in the diet varies both between and within species [Fashing, 2011]. This dietary flexibility is thought to allow for a large amount of lability in their social systems [Fashing, 2011]. In general, the red colobus (Piliocolobus spp.) consume more fruit than the other African colobines and live in larger groups (i.e., 15–75 individuals on average) [Korstjens and Dunbar, 2007]. Olive colobus (Procolobus verus) and most black-and-white colobus (Colobus spp.) are similar in that they tend to be more folivorous than red colobus and live in relatively small groups (i.e., 4–20 individuals on average) [Oates, 1988; Korstjens, 2001; Korstjens and Dunbar, 2007]. A striking exception to this pattern is Colobus angolensis, in which the Rwenzori sub-species can form groups that are extremely large. In the high-altitude forest of Nyungwe National Park (Rwanda) groups contain 300–500 individuals [Fimbel et al., 2001; Fashing et al., 2007; Miller et al., 2020c]. Although these groups are organized into multi-level societies [Miller et al., 2020c], groups are fairly cohesive [Fimbel et al., 2001; Fashing et al., 2007; Miller et al., 2020c]. Recent work from a mid-altitude forest at Lake Nabugabo (Uganda)
has found *C. a. ruwenzorii* aggregations of approximately 200 individuals also form a multi-level society, but groups are not cohesive, rather core units fission and fuse throughout the day [Stead and Teichroeb, 2019].

To date, *C. a. ruwenzorii* foraging strategies have only been studied at Nyungwe, where it is thought that they are able to live in a large and relatively cohesive society because they have access to an abundant, high-quality food resource [Fimbel et al., 2001; Fashing et al., 2007]. Here, their diet is primarily comprised of young and mature leaves (25 and 40%, respectively) [Fimbel et al., 2001], which is one of the highest documented levels of folivory among the Colobinae [Oates, 1977; Wasserman and Chapman, 2003; Saj and Sicotte, 2007a; Fashing, 2011]. However, the mature leaves at this high-altitude site are richer in protein and lower in fibre than has been observed at other black-and-white colobus field sites. When leaves are in low supply, the monkeys can fall back to feeding on lichens, which are also abundant [Miller et al., 2020a]. While other African colobines that are highly folivorous tend to dedicate much (i.e., >50%) of their time to resting [reviewed in Fashing et al., 2007; Fashing, 2011], *C. a. ruwenzorii* in Nyungwe spend only 32% of their time resting [Fashing et al., 2007]. Thus, this population does not seem to employ the energy minimization strategy typical of other highly folivorous black-and-white colobus [Fashing et al., 2007]. The low resting time seen in this population is suggested to be a manifestation of scramble competition, as so many individuals foraging together drives up search times and necessitates increased travel to find food [Fashing et al., 2007]. Indeed, scramble competition, inferred using patch depletion, appears to be an issue for this population for most food types with the exception of mature leaves [Miller et al., 2020b].

It is still unknown how generalizable an energy maximization foraging strategy is to other *C. a. ruwenzorii* populations and/or subspecies since *C. angolensis* is not well studied. It has been noted that *C. a. cottoni* in the Ituri Forest, DRC, are unusually active compared to other colobus monkeys [Bocian, 1997], which hints that this may be a species-wide pattern of behaviour. The goal of this study is to examine the diet and activity budget of *C. a. ruwenzorii* living in the mid-altitude forest at Nabugabo, Uganda, in order to determine the foraging strategies employed by this population. The habitat use of the Nabugabo population differs greatly from the Nyungwe population owing to differences in the availability of food at various forest strata and differences in the complement of predators [Adams and Teichroeb, 2020]. Although not as large as the group sizes observed in Nyungwe, bands of *C. a. ruwenzorii* at Lake Nabugabo are much larger than the group sizes observed in most black-and-white colobus. Therefore, we hypothesize that this population must also have access to abundant, high-quality food resources. Without nutritional analyses to confirm food quality, it is likely that high-quality resources at this lower altitude site will follow common patterns and consist of young leaves, seeds, or fruit, which are items that tend to be high in protein and carbohydrates but low in fibre [Lambert and Rothman, 2015; Ganzhorn et al., 2017]. To verify this supposition, we examined the extent to which these plant parts were consumed relative to their availability, and the extent to which *C. a. ruwenzorii* preferred these plant parts for each species present in their diet. We expect that the diet of the *C. a. ruwenzorii* at Nabugabo will largely consist of these high-quality, preferred foods. We also hypothesize that a high-quality diet will allow the *C. a. ruwenzorii* at Lake Nabugabo to employ an energy maximization foraging strategy. Therefore, we expect to see levels of resting time that are similar to those observed at Nyungwe [30–40%; Fashing et al., 2007] and lower than is typical of other black-and-white colobus species (>50%; Fashing et al., 2007; Fashing, 2011). Lastly, because there are known differences in group cohesion between the two sites, we do expect to see some differences in foraging strategies of the *C. a. ruwenzorii* at Nabugabo versus Nyungwe. The higher levels of group cohesion at Nyungwe are thought to be possible because both the preferred and fallback foods of the *C. a. ruwenzorii* at this site are abundant and uniformly distributed [Fimbel et al., 2001; Fashing et al., 2007], and therefore able to support large aggregations of animals [Janson and van Schaik, 1988; Johnson et al., 2002]. We hypothesize that the greater propensity for core units to fission and fuse at Nabugabo [Stead and Teichroeb, 2019] arises because they primarily consume foods that occur in a clumped distribution, and that these cannot always support large groups, necessitating group fission [Wrangham and White, 1988; de Sá and Strier, 1992; Asensio et al., 2007]. Therefore, we expect that both the preferred and fallback foods of the *C. a. ruwenzorii* living at Nabugabo will primarily consist of items such as fruits, seeds, or young leaves [Janson and van Schaik, 1988; Johnson et al., 2002].

**Methods**

**Study Site and Species**

Data for this study were collected on Rwenzori Angolan colobus monkeys (*Colobus angolensis ruwenzorii*) on the western edge of Lake Nabugabo, a small lake (8.2 × 5 km, elevation 1,136 m) ly-
ing just west of Lake Victoria in central Uganda (0°22’–12° S and 31°54’ E) (Fig. 1). The landscape is a mixed habitat of wetlands, grasslands, patches of primary and secondary forest, degraded forest, farmers’ fields, and a few buildings [Chapman et al., 2016]. There are two peaks in rainfall over the year from February to May and September to November, and the mean annual rainfall over a 5-year period was 1,117.21 mm (SD ±729.46) (2015–2019). The average temperature over the same 5-year span was 21.96 °C (SD ±1.21) (data from www.worldweatheronline.com for the nearby town (12.5 km away) of Masaka). The colobus monkeys were followed in a primary and secondary forest fragment, which is made up of the approximately 280-ha Manwa Forest Reserve and an unprotected forest fragment that borders the Juma River [Teichroeb et al., 2019]. Potential predators of the colobus consisted of venomous and constricting snakes (e.g., Dendroaspis polylepis, Dispholidus typus, Python sebae), crowned hawk eagles (Stephanoaetus coronatus), and local people’s dogs (Canis lupus familiaris) [Adams and Teichroeb, 2020].

We studied one large band of C. a. ruwenzorii (Band TR: approx. 120 individuals at the time of data collection) made up of 12 core units, of which 5 were uni-male/multi-female and 7 were multi-male/multi-female. We began following this band in 2015, and animals were well habituated. In the multi-level society that this species forms, there are at least three tiers of organization. Stable core units cluster into clans, and clans cluster together to share a home range in a band tier of organization. Core units fission and fuse with one another throughout the day, so although each core unit typically contains between 4 and 23 individuals, operational group size is often larger [Stead and Teichroeb, 2019]. No all-male unit was present during data collection for the current study but they do sometimes occur in the population [Adams and Teichroeb, 2020]. Preliminary findings suggest that while both sexes disperse, males tend to disperse within bands while females disperse between bands [Stead and Teichroeb, 2019]. Consequently, female-female bonds tend to be weaker than in other black-and-white colobus species [Arseneau-Robar et al., 2018].

**Behavioural Data Collection**

Diet and activity budget data were collected by a team of two field assistants (E. Mujjuzi and H. Kaketo) over 12 months, between May 2016 and April 2017 (mean 10.8 days per month, range: 5–14 days; mean 79.7 h/month). Field assistants were trained by J.A.T., and the same two observers collected data together throughout the data collection period and agreed on categories before recording them. Follows generally began at 07:30 and continued until 16:30. Sunrise at Nabugabo occurs just before 07:00, so the assistants usually began collecting data when the colobus were still resting in their sleeping trees. One caveat of our study is that assistants left the study band about 2 h prior to sunset. Thus, they likely missed at least one bout of each feeding, resting, and movement, which could have impacted the proportions in our calculated activity budget. Field assistants collected instantaneous scan samples [Altmann, 1974] via the “frequency method” [Struhsaker, 1975] every 30 min, on up to 5 individuals (n = 9,373 scans). Observers recorded each individual’s age-sex class, identity (if known), activity state (e.g., feeding, moving, resting, social, or other), and a GPS location was taken. When feeding, the food item (i.e., young leaf, mature leaf, fruit, flower, seed, seed pod, etc. – see below for definitions of

**Fig. 1.** The location of Uganda in Africa and the location of Lake Nabugabo in Uganda indicated by the red star. The satellite images show the forest fragment where the study band of *Colobus angolensis ruwenzorii* live with their home range outlined in red. The photo is of an adult female *C. a. ruwenzorii* (photo by Julie Teichroeb). Satellite image (Google Earth, 2018).
food items) and plant species were also noted (n = 2,277 feeding scans). Observers made an effort to move around the group between scans so that the same individuals were not scanned within the same 2-h period. In addition, observers swept through the visible individuals from left to right to avoid scanning the same individual twice in the same scan. Any fission and fusion events with other core units were recorded ad libitum. When the core units being followed underwent fission, observers stayed with either the largest subgroup or the one that contained core units with the most known individuals.

When recording food item type, the age of leaves was determined by their colour and texture. Compared to mature leaves, young leaves were either a lighter green, red, or pink colour, appeared softer in texture, and occurred at the tips of branches. If leaf buds or flower buds were eaten, this was recorded. Fruit feeding was recorded when fruit flesh and smaller seeds within the flesh were fed upon. Seed eating was recorded if larger pits within fruits were consumed or if small seeds were extracted out of pods. If the whole seed pod was fed upon, seed pod feeding was recorded.

Assessment of Food Availability

We used a line transect survey carried out in the year before behavioural data collection to assess the species composition of the forest patch where the colobus reside. We mapped the perimeter of the forest fragment by walking around it with a GPS and then cut 32 parallel, straight-line transects at 100-m intervals through the forest. Transects varied in length (range: 29–671 m) and were cut, either until the edge of the fragment was reached, or we moved into the swamp on the north-west side of the forest and standing water became more than 30 cm deep [Teichroeb et al., 2019]. To determine what tree species were available, we identified and measured all trees ≥ 10 cm diameter at breast height (DBH) within 5 m on each side of transects. Trees were identified to the species level with the assistance of a knowledgeable local botanist (M. Ponsio-no). For trees with multiple stems ≥ 10 cm DBH, we measured all stems and consolidated them by taking the square root of the sum of all squared DBHs [Nature Conservation, 2006]. On each transect, a scan was taken to assess average canopy height, and one tree that represented this height was measured using a clinometer. The total area sampled was 9,702 ha, which is 7% of the area covered by the forest fragment. Data from the ecological survey was used to determine the mean DBH (cm), stem density (stems/ha) and basal area (m²/ha) for each tree species. The basal area covered by each tree species was calculated by summing the area covered by each sampled tree (A = πr²) and dividing by the number of hectares sampled.

The seasonal availability of plant parts was recorded during monthly phenology surveys of 124 trees from 44 species, which pilot behavioural data had indicated were part of the colobus diet. We attempted a random sample of at least 3 individual trees of each species but not all 44 species had 3 individuals in the sample due to low availability. In addition, because the phenology survey was created before behavioural data collection began, lianas were not sampled, since it was not yet known how important they are in the diet. To ensure that phenology surveys were separated by a similar amount of time they were typically carried out in the middle of each month (range: the 11th to 18th day of the month). Observers used binoculars and indexed the approximate abundance of plant parts (i.e., flower buds and flowers, unripe and ripe seeds, unripe and ripe fruit, young leaves, mature leaves) by noting the percent of the canopy comprised of each part using a scale from 0 to 4, where 0 = the plant part is not present, 1 = 1–25%, 2 = 26–50%, 3 = 51–75%, and 4 = 76–100% [Sun et al., 1996]. For each tree species, phenology scores were averaged among all of the trees sampled in a given month. Food availability indices (FAI) were calculated for each plant part, each month, using the following formula, and summing the scores for all tree species [Fashing, 2001a; Dasilva, 1994]:

\[ \text{FAI} = \text{average phenology score for species } i \times \text{basal area for species } i \]

Data Analyses

We calculated the proportion of the annual diet comprised of each tree species, as well as each plant part. Temporal patterns in diet were investigated by determining the proportion of the diet constituted by each plant part, each month. The average monthly rainfall was obtained from https://www.worldweatheronline.com/ for the nearby town of Masaka (12.5 km away) and visually compared to phenological patterns. Activity budgets were calculated as the proportion of scans that individuals spent in each of the five behavioural categories (i.e., feed, rest, move, social, other), both annually and monthly. We determined the extent to which plant parts were selected relative to their availability by using Spearman rank correlations to compare the proportion of each plant part in the monthly diet to the food availability index for that plant part in that same month. To assess the extent to which each tree species was selected, we calculated selection ratios by dividing the proportion of each tree species in the annual diet by the proportion of the total basal area contributed by that tree species [Fashing, 2001a]. We calculated food preference scores for each part of each species using Ivlev’s electivity index (EI) [Ganas et al., 2008; Watts, 1984]:

\[ EI = \frac{m \text{onthly rank of that item in diet} - m \text{onthly rank of availability of that item}}{m \text{onthly rank of that item in diet} + m \text{onthly rank of availability of that item}} \]

In the rare cases where a part of a species was consumed but it did not appear in our phenology survey, we gave it an availability rank of “1” to indicate it was present but had low availability. Fall-back foods are foods that are relied upon when preferred foods are scarce [Altman, 1998; Doran-Sheehy et al., 2009]. We investigated the use of fallback foods using Spearman rank correlations to test for negative correlations between the consumption of potential fallback foods in the diet and the availability of (1) plant part(s) that were highly selected for or (2) the parts of species that were preferred in at least 1 month of the study period. In investigating fallback food use, we only considered plant parts that were important in the diet (i.e., >1% of the annual diet). In all Spearman rank correlations, we present Monte-Carlo approximated p values as these are more robust in the face of ties [Hájek et al., 1999], which occurred when rarely eaten plant parts comprised 0% of the diet in 2 or more months of the study period. Where appropriate we controlled for multiple comparisons using a Bonferroni correction to decrease the likelihood of false positives [Field et al., 2012]. Statistics were conducted using the “coin” package [Hothorn et al., 2006] in R (version 3.3.2; R Core Team, 2017) with R Studio version 1.1.463 [R Studio Team, 2020].

Rwenzori Colobus Diet and Activity Budget

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39
Results

Young leaves (65%) and fruits (31%) were the items most frequently consumed by the C. a. ruwenzorii at this field site. Mature leaves comprised only 3% of the annual diet, and other food items like bark (1%), flowers (<1%), and seeds (<1%) were rarely eaten (Table 1). While young leaves were consistently available in large quantities, fruits were most abundant during the dry season (May to August) (Fig. 2a, b). There was a significant positive correlation between the availability of fruits and their consumption (Table 2), and as a result, fruits were the most common item in the diet of the C. a. ruwenzorii in the dry season, as well as in January when there was also little rain (Fig. 2c). Young leaf consumption was not related to availability (Table 2), but young leaves did appear to be the preferred food item when fruits were scarce, and so comprised the bulk of the diet in wetter months (Fig. 2c). Although mature leaves were abundant throughout the year, they tended to be eaten less than would be expected given their availability (Table 2) and were only consumed during the peak wet season (October and November) (Fig. 2c). There was little temporal pattern in the consumption of flowers and flowerbuds, seeds and seedpods, or bark (Fig. 2c).

The colobus monkeys were observed to feed from 20 tree species, 2 shrub species, and several liana species. Only a handful of tree species comprised the bulk of the diet of the C. a. ruwenzorii at this site (Table 3). The most frequently eaten tree species included Maesopsis eminii (24% of the annual diet), Antiaris toxicaria (19%), Newtonia buchananii (4%), and Pseudospondias microcarpa (4%). These species were common in the study area, and after accounting for availability, only A. toxicaria and M. eminii were highly selected (Table 3). The C. a. ruwenzorii ate the fruits, young and mature leaves, and the bark of these two species. Other species were highly selected (i.e., selection ratios >1), but because they were relatively rare within the study area, they comprised a smaller proportion of the diet: Prunus africana (2% of the annual diet), Rauvolfia vomitoria (<1%), Sapium ellipticum (3%), and Swietenia macrophylla (<1%) (Table 3). None of the consumed tree species were consistently important over time (i.e., consistently comprised more than 1% of the monthly diet), nor did any one species comprise the majority of their diet in any given month (Table 3). When we look more specifically at preferences for the parts of each species that were consumed, we find that only the fruits and young leaves of some species were preferred (Table 3). The fruits of A. toxicaria, M. eminii, P. microcarpa, S. ellipticum, Teclea nobilis, and the seeds of N. buchananii were preferred in at least 1 month of the study period. Similarly, C. a. ruwenzorii also showed a preference for the young leaves of A. toxicaria, P. africana, S. ellipticum, and S. macrophylla in at least 1 month of the study period. Lianas were an important food resource for the colobus making up 33.7% of the annual diet (Table 3); however, since they were not represented in the phylogeny survey, we could not assess the extent to which they were selected/preferred.

Forty percent of the annual activity budget was dedicated to resting, 28% to feeding, 25% to moving, 7% to social interactions, and 1% to other activities (Table 1). There was very little temporal variation in the activity

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**Table 1.** Annual diet (% of the annual diet) and activity budget (% of the annual activity budget) for Colobus angolensis ruwenzorii living near Lake Nabugabo, Uganda

| Plant part       | Young leaves | Mature leaves | Ripe and unripe fruits | Seeds and seedpods | Flowers and flowerbuds | Bark |
|------------------|--------------|---------------|-----------------------|--------------------|------------------------|------|
| Behaviour        | Feeding      | Moving        | Resting               | Social             | Other                  |      |
|                  | 65%          | 3%            | 31%                   | <1%                | <1%                    | 1%   |

| Table 2. Spearman rank correlations between the proportion of each plant part in the monthly diet of Colobus angolensis ruwenzorii and the food availability index for that plant part, that month

| Plant part       | p              | Z              | MC-approximated p value |
|------------------|----------------|----------------|-------------------------|
| Flowers and flowerbuds | 0.044         | 0.145          | 1.000                   |
| Ripe and unripe fruits | 0.692         | 2.296          | 0.014                   |
| Seeds and seedpods   | 0.516          | –0.691         | 1.000                   |
| Young leaves       | –0.028         | –0.093         | 0.936                   |
| Mature leaves      | –0.574         | –1.904         | 0.052                   |

Because there were many ties in the data sets, we present Monte-Carlo (MC)-approximated p values. Significant relationships are in bold and trends are italicized.
budget of the *C. a. ruwenzorii* at this site over the year (Fig. 3). September is the only month with a notable change, with a decrease in moving and social behaviour and an increase in feeding and resting coinciding with the switch from more fruit feeding back to more young-leaf feeding.

In assessing the use of fallback foods, we found no significant correlation between the consumption of young leaves and the availability of ripe and unripe fruits, which were the plant parts that were highly selected when available (Spearman: $r_s = -0.580$, $Z = -1.925$, $p = 0.116$). However, we did find that mature leaves tended to be consumed when fruits were less available within the forest (Spearman: $r_s = -0.603$, $Z = -2.001$, $p = 0.084$). Similarly, mature leaf consumption was significantly negatively correlated with the availability of preferred food items, which included the fruits and/or young leaves of 8 tree species (Spearman: $r_s = -0.580$, $Z = -1.945$, $p = 0.047$). Importantly, this fallback food comprised only 3% of the annual diet (Table 1). Similarly, food items that had low preference scores accounted for only 15% of the annual diet (Table 3).

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**Fig. 2.** a Average daily rainfall (mm/day) for each month of the study period. b Food availability index scores for the different plant parts consumed. c The monthly diet of *Colobus angolensis ruwenzorii* near Lake Nabugabo, Uganda.
Table 3. The diet of the *Colobus angolensis ruwenzorii* living near Lake Nabugabo, Uganda, including the species eaten, their proportion in the annual and monthly diet (minimum and maximum proportions), and the selection ratio for each species

| Species                        | Family              | Plant parts | Proportion of annual diet | Min. and max. proportion of monthly diet | Selection ratio Months consumed, n | Mean (SD) preference score | Preferred in at least 1 month |
|-------------------------------|---------------------|-------------|---------------------------|----------------------------------------|----------------------------------|---------------------------|-----------------------------|
| **Trees**                     |                     |             |                           |                                        |                                  |                           |                             |
| Albizia zygia                 | Fabaceae            | Total       | 0.000                     | 0.000 0.004 0.003 0.1                  |                                  |                           |                             |
|                               |                     | BK          | 0.000                     |                                          |                                  |                           |                             |
| Antiaris toxicaria            | Moraceae            | Total       | 0.191                     | 0.000 0.004 0.430 0.044 4.4             |                                  |                           |                             |
|                               |                     | BK          | 0.001                     | 0.000 0.005                             |                                  |                           |                             |
|                               |                     | FR          | 0.010                     | 0.000 0.163                            | 2 0.73 (0.04) *                   |                           |                             |
|                               |                     | YL          | 0.158                     | 0.000 0.440                            | 10 -0.63 (0.30) *                 |                           |                             |
|                               |                     | ML          | 0.018                     | 0.000 0.089                            | 2 -0.21 (0.03)                   |                           |                             |
| Apodytes dimidiata            | Metteniusiacae      | Total       | 0.003                     | 0.000 0.014 0.008 0.008 0.3           |                                  |                           |                             |
|                               |                     | BK          | 0.000                     |                                          |                                  |                           |                             |
|                               |                     | FR          | 0.001                     | 0.000 0.005                            | 1 -0.81                          |                           |                             |
|                               |                     | YL          | 0.001                     | 0.000 0.008                            | 2 -0.44 (0.48)                   |                           |                             |
|                               |                     | ML          | 0.001                     | 0.000 0.014                            | 2 -0.88 (0.06)                   |                           |                             |
| Canarium schweinfurthii       | Burseraceae         | Total       | 0.001                     | 0.000 0.006 0.017 0.1                  |                                  |                           |                             |
|                               |                     | BK          | 0.000                     | 0.000 0.006                            |                                  |                           |                             |
|                               |                     | YL          | 0.000                     | 0.000 0.004                            | 1 -0.85                          |                           |                             |
| Ficus congensis               | Moraceae            | Total       | 0.000                     | 0.000 0.005 0.005 0.1                  |                                  |                           |                             |
| Ficus nitidus                 | Moraceae            | Total       | 0.002                     | 0.000 0.015 0.009 0.2                  |                                  |                           |                             |
| Funtumia latifolia            | Apocynaceae         | Total       | 0.002                     | 0.000 0.008 0.097 0.0                  |                                  |                           |                             |
|                               |                     | BK          | 0.001                     | 0.000 0.007                            |                                  |                           |                             |
|                               |                     | YL          | 0.001                     | 0.000 0.009                            | 3 -0.60 (0.13)                   |                           |                             |
| Harungana madagascariensis    | Hypericaceae        | Total       | 0.000                     | 0.000 0.007 0.003 0.1                  |                                  |                           |                             |
|                               |                     | YL          | 0.000                     | 0.000 0.007                            | 1 -0.67                          |                           |                             |
| Lovoa trichilioides           | Meliaceae           | Total       | 0.000                     | 0.000 0.004 0.013 0.0                  |                                  |                           |                             |
|                               |                     | YL          | 0.000                     | 0.000 0.004                            | 1 -0.66                          |                           |                             |
| Macaranga schweinfurthii      | Euphorbiaceae       | Total       | 0.001                     | 0.000 0.005 0.071 0.0                  |                                  |                           |                             |
|                               |                     | YL          | 0.001                     | 0.000 0.005                            | 2 -0.84                          |                           |                             |
| Mecopsis emnii                | Rhamnaceae          | Total       | 0.242                     | 0.000 0.802 0.178 1.4                  |                                  |                           |                             |
|                               |                     | BK          | 0.001                     | 0.000 0.006                            |                                  |                           |                             |
|                               |                     | FR          | 0.196                     | 0.000 0.806                            | 8 0.09 (0.75) *                  |                           |                             |
|                               |                     | YL          | 0.040                     | 0.000 0.096                            | 10 -0.58 (0.19)                  |                           |                             |
|                               |                     | ML          | 0.002                     | 0.000 0.008                            | 3 -0.68 (0.19)                   |                           |                             |
| Newtonia buchananii           | Fabaceae            | Total       | 0.036                     | 0.003 0.066 0.070 0.5                  |                                  |                           |                             |
|                               |                     | BK          | 0.004                     | 0.000 0.017                            |                                  |                           |                             |
|                               |                     | SD          | 0.000                     | 0.000 0.005                            | 1 0.50                          |                           |                             |
|                               |                     | YL          | 0.031                     | 0.005 0.062                            | 12 -0.61 (0.19)                  |                           |                             |
|                               |                     | ML          | 0.000                     | 0.000 0.004                            | 1 -0.59                          |                           |                             |
| Prunus africana               | Rosaceae            | Total       | 0.018                     | 0.000 0.049 0.006 3.0                  |                                  |                           |                             |
|                               |                     | BK          | 0.000                     | 0.000 0.004                            |                                  |                           |                             |
|                               |                     | YL          | 0.017                     | 0.000 0.049                            | 7 -0.18 (0.48) *                 |                           |                             |
| Pseudopondias microcarpa      | Anacardiaceae       | Total       | 0.035                     | 0.000 0.131 0.196 0.2                  |                                  |                           |                             |
|                               |                     | BK          | 0.001                     | 0.000 0.009                            |                                  |                           |                             |
|                               |                     | FR          | 0.006                     | 0.000 0.043                            | 3 0.85 (0.06) *                  |                           |                             |
|                               |                     | YL          | 0.027                     | 0.000 0.136                            | 11 -0.65 (0.20)                  |                           |                             |
|                               |                     | ML          | 0.001                     | 0.000 0.005                            | 2 -0.70 (0.06)                   |                           |                             |
| Rauvolfia vomitoria           | Apocynaceae         | Total       | 0.005                     | 0.000 0.028 0.000 10.9                 |                                  |                           |                             |
|                               |                     | LV          | 0.005                     | 0.000 0.028                            | 5 0.52 (0.24) *                  |                           |                             |
| Sapium ellipticum             | Euphorbiaceae       | Total       | 0.025                     | 0.000 0.086 0.005 4.8                  |                                  |                           |                             |
|                               |                     | FR          | 0.005                     | 0.000 0.023                            | 3 0.48 (0.66) *                  |                           |                             |
|                               |                     | YL          | 0.019                     | 0.000 0.009                            | 8 -0.07 (0.32) *                 |                           |                             |
|                               |                     | ML          | 0.001                     | 0.000 0.009                            | 1 -0.17                          |                           |                             |
| Swietenia macrophylla         | Meliaceae           | Total       | 0.001                     | 0.000 0.005 0.001 2.6                  |                                  |                           |                             |
|                               |                     | BK          | 0.001                     | 0.000 0.004                            |                                  |                           |                             |
|                               |                     | YL          | 0.000                     | 0.000 0.005                            | 1 0.50                          |                           |                             |
| Teclea nobilis                | Rutaceae            | Total       | 0.002                     | 0.000 0.011 0.005 0.4                  |                                  |                           |                             |
|                               |                     | FR          | 0.001                     | 0.000 0.009                            | 2 -0.04 (0.33) *                 |                           |                             |
|                               |                     | YL          | 0.001                     | 0.000 0.011                            | 1 -0.14                          |                           |                             |
Discussion

The diet of the *C. a. ruwenzorii* living at Nabugabo, Uganda, largely consisted of ripe and unripe fruits and young leaves. Fruits were selected when they were available, and both fruits and young leaves of some species were preferred food items. Mature leaf consumption tended to correlate negatively with these selected/preferred food items, suggesting that mature leaves are a fallback food for this population. However, mature leaves comprised a very small proportion of the annual diet, signifying that the *C. a. ruwenzorii* at this site rely minimally on fallback foods. This is likely because young leaf availability was only slightly lower than mature leaf availability for much of the year, and fruits were seasonally abundant. Furthermore, preferred species parts were
available throughout the year. These findings suggest that the *C. a. ruwenzorii* living at Nabugabo had access to abundant, high-quality food resources year-round. This high-quality diet likely allowed them to employ an energy maximization foraging strategy, as the observed levels of resting time were comparable to those seen in more frugivorous African colobines and lower than typically observed in more folivorous species [Korstjens et al., 2010; Fashing, 2011]. Their reliance on food items that tend to have a clumped distribution likely explains why cohesion between core units is lower in the Nabugabo population than has been reported elsewhere [Fimbel et al., 2001; Fashing et al., 2007; Miller et al., 2020c].

The observed consumption of young leaves in this study is one of the highest values reported in the African colobines, being higher only in *Procolobus verus* at Tai National Park [Korstjens et al., 2007], and matched by *C. guereza* at Kibale National Park [Oates, 1977]. Mature leaves were avoided by the *C. a. ruwenzorii* at our study site and were only used as a fallback food that the monkeys relied upon to a minimal extent in the wettest months of the year. In this respect, *C. a. ruwenzorii* at Nabugabo are similar to most other black-and-white colobus populations that show avoidance of mature leaves except during seasonal bottlenecks [Colobus polykomos, Dasilva, 1994; *C. satanas*, Fleury and Gautier-Hion, 1999; *C. guereza*, Fashing, 2001a; Wasserman and Chapman, 2003; *C. vellerosus*, Saj and Sicotte, 2007a, b], with the notable exception of the *C. a. ruwenzorii* in Nyungwe [Fimbel et al., 2001].

Fruit was significantly positively selected for when it became available, and the fruits of *A. toxicaria*, *M. eminii*, *P. microcarpa*, *S. ellipticum*, and *T. nobilis* (as well as the seeds of *N. buchananii*) were preferred food items in this population. The fruits of *M. eminii* were particularly important as they comprised 20% of the annual diet. Colobines are not thought to be able to feed on large quantities of ripe fruit because the high acidity of the fruit flesh can kill the bacteria in the forestomach that aid in the breakdown of cellulose in leaves [Goltenboth, 1976]. To mitigate this, colobus monkeys are often described as feeding on unripe fruits [Fashing, 2011]. As Davies et al. [1999] put it “a colobus fruit can be described as: dull green or brown/black; unripe; weighing <50 g; generally lack flesh, but when it is present it is dry or fibrous or both” (p. 345). The fruits of *M. eminii* and *A. toxicaria*, as well as the shrub *Alchornea cordofolia*, fit this description well. Furthermore, for *M. eminii* fruit, the colobus at Nabugabo often eat off the outer layer of the fruit skin and leave the fleshier parts of the fruit intact. Importantly, *M. eminii* accounts for 18% of the total forest biomass, *A. toxicaria* for 4%, and *A. cordofolia* is also common. Thus, the high levels of frugivory observed in this study likely occur because the *C. a. ruwenzorii* at Nabugabo have access to abundant amounts of fruits that they are able to digest, despite being an anatomical folivore, and they are using behavioural strategies for exploiting the available fruit resources to the fullest.

While abundant food resources and behavioural strategies for exploiting them seem to allow the *C. a. ruwenzorii* to form large groups, scramble competition is very likely an issue in this population, since it has been shown to impact folivorous primates in groups that are comparatively much smaller [e.g., Steenbeek and van Schaik, 2001; Snaith and Chapman, 2008; Teichroeb and Sicotte, 2009]. Adjusting activity budgets is one of the main ways that animals can compensate for increased food competition [Caraco, 1979; Janson, 1988]. However, it has been suggested that folivoses may be limited in the degree that they can use this strategy [Borries et al., 2008; Korstjens et al., 2010]. Resting time may be dictated by a high-fibre diet paired with slow digestion that includes fermentation [Lambert, 1998], feeding time may be limited by gut capacity [Stephens and Krebs, 1986], and moving time may be limited by a diet with a low overall energy yield [Janson and Goldsmith, 1995]. If the diet is altered however, to include less fibrous food with more energy available from either carbohydrates or protein, colobus may be able to adjust activity budgets and behave more similarly to a typical energy-maximizing frugivore. Resting times for the *C. a. ruwenzorii* at Nabugabo (40%) were among the lowest observed in the African colobines [Fashing et al., 2007] and are most similar to those seen in the more frugivorous red colobus [reviewed in Fashing, 2011]. In support of our second hypothesis, this finding suggests that this population is employing an energy maximization foraging strategy. This strategy is likely possible because of the high-quality resources available. A diet concentrated on young leaves and fruit should result in a relatively low consumption of fibre [Waterman et al., 1980]. Consequently, digestion may not be inhibited as greatly as for those species that eat large quantities of mature leaves [Milton, 1979; Chivers, 1994; Lambert, 1998]. This inference needs to be investigated with more research on the nutritional composition of their foods; however, work in other species has shown that when more energy is available, animals can adjust their time budgets and movement patterns [Godfrey, 2003; Riley, 2007; Dunbar et al., 2009]. The same species can show energy-minimizing or energy-maximizing strategies depending on the quality...
of their current habitat [A. calmitans, Jung et al., 2015; A. pigra, Rangel-Negrin et al., 2018; Klass, 2020; A. palliata mexicana, Asensio et al., 2007; C. djamadjemensis, Meikonen et al., 2017].

There were important differences in the diet and activity budgets observed in this study, and those observed in previous work on the same subspecies living in Nyungwe National Park, Rwanda. While both populations had access to abundant high-quality resources, the types of resources, and therefore the diets in the two populations of C. a. ruwenzorii, differed greatly. Our study band at Nabugabo relied on young leaves (65%) and fruits (31%) while the Nyungwe population relied on fruit less (16%) and largely consumed lichens [21%, Miller et al., 2020a] and mature leaves that had high protein-to-fibre ratios [40%, Fimbel et al., 2001]. We found that the C. a. ruwenzorii at Nabugabo spent less time feeding (28%) and more time moving (25%) than their conspecifics at Nyungwe [Fashing et al., 2007]. These differences in activity budget are likely related to the distribution of each population’s important food resources. The large, relatively cohesive supergroups observed at Nyungwe are only possible if food resources are both abundant and uniformly distributed [Wrangham et al., 1993; Janson and Goldsmith, 1995; Janson and van Schaik, 1988]. The large valley bowls at Nyungwe are filled with lianas whose mature leaves are nutritious [Fimbel et al., 2001; Fashing et al., 2007]. Here, the colobus monkeys spend 1–3 h per day on the ground feeding on these uniformly distributed food resources [A. Miller, pers. commun.]. This “grazing-like” feeding pattern does not require a lot of between-patch movement, which could account for the lower amount of time spent moving and the higher amount of time spent feeding in Nyungwe. To minimize scramble competition, the large band maintains a very large home range (>20 km² compared to the much smaller <2 km² observed at Nabugabo) [Fashing et al., 2007; Adams and Teichroeb, 2020] and sometimes migrates large distances to access new food patches [Fashing et al., 2007]. Thus, after exploiting the resources in one area, the band moves on and allows a long regeneration time before returning to the same patch [Fashing et al., 2007]. Maintaining such large groups may be of paramount importance at Nyungwe, since chimpanzees (Pan troglodytes) are present as predators [Fashing et al., 2007].

Conversely, the C. a. ruwenzorii at Nabugabo rely on food resources that are not uniformly distributed. The seasonal flushing of young leaves and fruiting of trees typically leads to a resource base with a clumped distribution [Janson and van Schaik, 1988; Johnson et al., 2002], and even within food patches, young leaves and fruit are typically distributed in clusters near terminal branches [Snaith and Chapman, 2005]. As a result, the colobus here almost always forage in the canopy [Adams and Teichroeb, 2020] and frequently move both between discrete food patches and within food patches. This feeding pattern likely explains the lower time spent feeding and higher time spent moving in Nabugabo versus Nyungwe. It has been commonly observed that time spent moving increases in patchy habitats in both non-primate species [wild turkeys, Meleagris gallopavo silvestris, Marable et al., 2012; wood mice, Apodemus sylvaticus, Corp et al., 1997; racoons, Procyon lotor, Beasley and Rhodes, 2010] and in primates [A. clamitans, Jung et al., 2015; A. pigra, Ostro et al., 1999; Chlorocebus pygerythrus and Erythrocebus patas, Isbell et al., 1998; Lemur catta, Gabriel, 2013]. Importantly, the clumped distribution of resources at Nabugabo (e.g., fruits and young leaves) should result in high levels of scramble competition as group size increases, making it difficult for the large band of C. a. ruwenzorii at Nabugabo to remain cohesive. The fission-fusion social organization observed in this population [Stead and Teichroeb, 2019] appears to be an effective strategy for dealing with scramble competition as it would allow the C. a. ruwenzorii to adjust operational group size to the current abundance and distribution of resources. Supporting this supposition are recent findings that core units are most likely to fuse together into large aggregations when fruit resources are at their most abundant [Adams et al., in review].

Our findings, as well as the findings of others, highlight that C. a. ruwenzorii are able to form extremely large groups when their resource base allows it, but flexibly adjust grouping patterns to changing resource availability and/or distribution. We cannot be certain how generalizable these patterns are across other subspecies of C. angolensis, but “supertroops” or aggregations of groups have been observed in multiple studies, suggesting the potential to form large, hierarchically structured groups may be a species-wide trait [e.g., C. a. cottoni, Bocian, 1997; C. a. palliata, Moreno-Black and Bent, 1982]. This begs the question of why other species of black-and-white colobus only form groups averaging 4–20 individuals [Fashing, 2011], even when living in high-quality habitats [e.g., C. guereza, Fashing, 2001a; C. polykomos, Dasilva, 1994; Korstjens, 2001], instead of the hundreds of individuals observed in some C. angolensis studies. It is possible that predation rates vary for black-and-white colobus populations and that these tend to be higher for C. a. ruwenzorii populations providing an incentive for larger group sizes; however, we do not currently have data that suggest this.
We posit that even when ecology allows large groups to form, social behaviour and reproductive strategies may still keep groups small. The large group sizes observed in *C. a. ruwenzorii* can only occur when there is a high level of between-unit tolerance, particularly among adult males. In two other species of black-and-white colobus, adult males have been shown to defend important food-rich areas for the females in their group, and these males show little tolerance for nearby conspecific groups [C. vellerosus, Teichroeb et al., 2012; Teichroeb and Sicotte, 2018; *C. guereza*, Fashing, 2001b; Harris, 2006]. Conversely, it has been anecdotally noted that males in *C. angolensis* are more tolerant towards males from their own group, as well as extragroup males, than *C. guereza* [Bocian, 1997]. Therefore, an exciting avenue of future research would be to investigate the form of interunit interactions and male relationships in *C. angolensis*, including in the Ruwenzori subspecies.

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