Mechanical Defenses in Leaves Eaten by Costa Rican Howling Monkeys (Alouatta palliata)

M.F. Teaford,1* P.W. Lucas,2 P.S. Ungar,3 and K.E. Glander4

1Center for Functional Anatomy and Evolution, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205
2Department of Anatomy, George Washington University, Washington, DC 20052
3Department of Anthropology, University of Arkansas, Fayetteville, AR 72701
4Department of Biological Anthropology and Anatomy, Duke University, Durham, NC 27708

KEY WORDS Alouatta; diet; fracture toughness; silica

ABSTRACT Primate species often eat foods of different physical properties. This may have implications for tooth structure and wear in those species. The purpose of this study was to examine the mechanical defenses of leaves eaten by Alouatta palliata from different social groups at Hacienda La Pacifica in Costa Rica. Leaves were sampled from the home-ranges of groups living in different microhabitats. Specimens were collected during the wet and dry seasons from the same tree, same plant part, and same degree of development as those eaten by the monkeys. The toughness of over 300 leaves was estimated using a scissors test on a Darvell mechanical tester. Toughness values were compared between social groups, seasons, and locations on the leaves using ANOVA. Representative samples of leaves were also sun-dried for subsequent scanning electron microscopy and energy dispersive x-ray (EDX) analyses in an attempt to locate silica on the leaves. Both forms of mechanical defense (toughness and silica) were found to be at work in the plants at La Pacífica. Fracture toughness varied significantly by location within single leaves, indicating that measures of fracture toughness must be standardized by location on food items. Monkeys made some food choices based on fracture toughness by avoiding the toughest parts of leaves and consuming the least tough portions. Intergroup and seasonal differences in the toughness of foods suggest that subtle differences in resource availability can have a significant impact on diet and feeding in Alouatta palliata. Intergroup differences in the incidence of silica on leaves raise the possibility of matching differences in the rates and patterns of tooth wear. Am J Phys Anthropol 129:99–104, 2006. ©2005 Wiley-Liss, Inc.

Until recently, very little information has been reported from the field about the physical properties of foods that primes eat, though this could be an important factor in food selection and in understanding the form and wear resistance of the dentition (Lucas, 1979; Lucas and Peters, 2000; Lucas and Teaford, 1994; Strait, 1997; Yamashita, 1996). In the last decade or so, the situation has changed considerably, and recent work has shown that primate species often eat foods with different physical properties (Kinsey and Norconk, 1990, 1993; Strait, 1997; Yamashita, 1996). As the resources available to primates may vary significantly from one habitat to another (Teaford and Glander, 1996), food choice, and thus dental function, may also vary within species.

While fruits, flowers, leaves, and seeds vary in their relative importance in primate diets, leaves are easier to study mechanically because of their two-dimensional form. There are two major mechanical defenses in leaves. The first is their toughness, or the energy required to propagate a crack through them (Vincent, 1990, 1992). In thin, sheet-like foods such as leaves, toughness is the major factor obstructing the rapid rate of food breakdown in the mouth. For any two given foods (all other features of their loading in the mouth being similar), a tougher food will require higher forces to fracture (Wright and Vincent, 1996). The second mechanical defense is provided by silica in the leaf. Silica phytoliths are now known to be rather common in angiosperm leaves (Piperno, 1989). In the presence of particles hard enough to plastically indent (and thus scratch or pit) dental enamel, higher forces must be associated with higher wear rates (Lucas and Teaford, 1995). The cause of these scratches and pits could be either opaline (amorphous) phytoliths or extraneous siliceous grit (either quartz or a variety of aluminosilicates), since any of these is hard enough to indent enamel (Lucas and Teaford, 1995; Ungar et al., 1995).

We previously reported on canopy-level dust particles, with a mean minimum diameter ranging from 1.6 µm in the wet season to 8.2 µm in the dry season, and a rate of accumulation ranging from 12.5 particles per mm² per day in the dry season to 117.0 particles per mm² per day in the wet season (Ungar et al., 1995). Here we supplement this information with a survey of phytoliths in leaves (collected in the wet season) and measurements of toughness of leaves eaten by the howlers in both seasons.

Grant sponsor: NSF; Grant numbers: DBC 9118876; SBR 9601766; Grant sponsor: COSEN-PEW.

*Correspondence to: Mark F. Teaford, Center for Functional Anatomy and Evolution, Johns Hopkins University School of Medicine, 1830 E. Monument St., Room 303, Baltimore, MD 21205. E-mail: mteaford@jhmi.edu

Received 19 April 2004; accepted 8 October 2004.

DOI 10.1002/ajpa.20225 Published online 31 August 2005 in Wiley InterScience (www.interscience.wiley.com).
More specifically, this study examines the physical properties of leaves eaten by *Alouatta palliata* from three different social groups at Hacienda La Pacifica in Costa Rica. The study area is a combination cattle ranch and rice farm of 1,990 ha, with pastures and fields broken by patches of dry forest serving as windbreaks (for further information on this site, see Teaford and Glander, 1996). The ranch is bounded by the Rios Corobicí and Tenorio, each being lined by rich gallery forest. Mantled howlers are the only nonhuman primates on the ranch and are found in virtually all forest patches. The nonriver habitats show lower tree diversity than do the riverine habitats (Glander and Nisbett, 1996). Furthermore, the most plentiful trees in each habitat differ significantly, showing that resource availability varies between microhabitats. In this study, we tested the hypothesis that primates living in less diverse habitats would be forced to consume leaves that were better defended mechanically. This would have important consequences for dental function in animals that occupy different habitats.

**MATERIALS AND METHODS**

Although 42 howler social groups have been identified at La Pacifica, the analyses presented here focus on one nonriver group (known as group 1) and one river group (known as group 19). In the wet season, another nonriver group (group 2) was also included in the study (for group descriptions and locations, see Teaford and Glander, 1996). For each group, focal animals were followed throughout the day, and each of the different food items consumed was noted, including plant part and color (if applicable). Food items were collected whenever possible during the day to ensure that the samples analyzed were from the same tree, same plant part, and same degree of development as those eaten by the monkeys (e.g., young leaves vs. mature leaves).

This study focuses on the commonest leaves eaten by the study groups during July 1997 (at the height of the wet season) and February–March 1998 (at the height of the dry season). Unfortunately, during data collection, it quickly became apparent that many trees were missing their tree tags, and that some had also been previously misidentified. Thus, we cannot present species-by-species summaries of physical properties at this time, and our selection of specimens was dictated solely by which ones the monkeys were eating. As a result, the plant samples probably include the same plant species from different home ranges. However, the number of specimens is a good indicator of the number of species tested within each social group.

Mechanical tests of leaves were made, within 24 hr of collection, using a portable fracture toughness tester described by Darvell et al. (1996). The tester can be linked to a notebook computer, and it offers true force-displacement integration built into its hardware, allowing quick estimates of the toughness of food items (measured in Joules/meters squared). Though capable of running a variety of tests, those run here utilized the scissors-cutting test. Cut 1 is an attempt to define the “least tough” path within the lamina (Fig. 1), for which there is a lot of comparable information (Choong et al., 1992; Hill and Lucas, 1996; Lucas and Pereira, 1990). The major nutrients in a leaf are found in the mesophyll tissues (the palisade and spongy layers), which are best sampled by this cut (Choong, 1996). Cut 2 crosses secondary veins, cut 3, the midrib, and cut 4, the petiole (Fig. 1). These transport tissues contain much more fiber (i.e., cell wall) per unit volume than photosynthetic tissues (Lucas et al., 1995). Veins are often sheathed by cells called fibers, which is the anatomically correct term for the typical cells of angiosperm woody tissue. These possess 10 times the toughness of other cells of the same fiber content (Lucas et al., 1997, 2000), where “fiber” here refers to the nutritional usage of the term. The last two cuts, 5 and 6, were made in an attempt to understand the apparent preference of some primates (including howlers) for leaf tips in terms of toughness selection. The procedure only involves two parallel cuts (Fig. 1). Thus, in order to establish whether some gradient might be found, a toughness map involving a series of cuts parallel to cuts 5 and 6 was made on one species (*Gliricidia sepium*) (Fig. 2).

Howlers at La Pacifica also eat some unexpanded leaves and associated shoots. These leaves were too small to examine by the systematic method described above. Instead, cuts were taken at key locations along the shoot, such as with guanacaste shoots (*Enterolobium cyclocarpum*) (Fig. 3).

Of the major cuts, results from cuts 1, 5, and 6 are reported here. Toughness estimates for these cuts were
RESULTS

Intra-leaf differences in toughness

One difference consistently underlaid all the results of this study: measures of toughness varied between locations on the leaves. Thus, measurements taken at the basal end of the leaf (cut 6) were significantly higher than those taken elsewhere on the leaf (cuts 1 or 5) \( (P < 0.000, P < 0.003, \text{respectively}) \) (Tables 1–3, Fig. 2). This difference appeared irrespective of the other factors considered in our analyses, e.g., season of collection or social group. In addition, we also noted that the attachment site of leaves was generally much tougher than the rest of the food consumed. This is shown very clearly with *Gliricidia* leaves (Fig. 2) and guanacaste shoots (Fig. 3), where it appears that there is a very rapid fall-off in toughness away from the attachment site.

Seasonal variations in consumption of tough leaves

During the dry season, leaf toughness data were only collected for the foods eaten by two social groups (1 and 19). Thus, seasonal differences could only be tested for those two groups. As leaf toughness and season of collection varied independently of cut (Table 2), there was no need to do separate analyses for each cut. Group 1 simply ate tougher leaves in the wet season than in the dry season, and group 19, by contrast, ate tougher leaves in the dry season than in the wet season (Tables 1 and 4).

Intergroup differences in consumption of tough leaves

Results indicate significant differences in the toughness of leaves eaten by different social groups as well. Because of a significant interaction between group and season (Table 2), leaf toughness values for groups were compared separately for the wet and dry seasons. Again, as leaf toughness and social group varied independently of cut, there was no need for separate analyses for each cut. Wet-season results show a significant difference between groups 1 and 2 (Tables 1 and 5), and dry-season results show a significant difference between groups 1 and 19 \( (F = 16.849, \text{df} = 1, P < 0.000) \).

Silica in leaves in the wet season

About 40% of all leaves tested possessed silica, mostly associated with sharp-tipped trichomes (or hairs) projecting from the leaf surface (Fig. 4). Chi-square analyses indicated that leaves found in the home range of group 19 showed a significantly lower incidence of silica than did leaves from the home range of group 1 (Table 6).

DISCUSSION

Intra-leaf differences in toughness

The pattern of development of toughness of tropical leaves was studied in detail for *Castanopsis fissa* (Fagaceae) leaves by Choong (1996, 1997). During development, the leaf base toughens more quickly than the leaf tip, presumably to protect the transport system of the leaf from predators. Rupture that introduces even a single air bubble into a major vein is terminal for all parts of the leaf that depend on that vein. In a mature *C. fissa*
leaf, the midrib and secondary veins are tougher at their origin and at the margins of the leaf than elsewhere. Our data substantiate a major difference between the toughness of the leaf base and leaf tip in most leaves in our sample. In *Gliricidia sepium*, where we have a large sample (n = 94 cuts), the basal third is typically about 30% tougher than the apical third of the mature leaf. Since toughness is produced by the behavior of cell walls (Lucas et al., 1995, 1997), the frequent selection of the apical part of leaves by the howlers at La Pacifica is basically selection against toughness and fiber content (Fig. 5). A lower-volume fraction of fiber (cell wall) in a leaf means a higher-volume fraction of cytoplasm (the protoplasm) and thus greater nutritional value. Waterman and Kool (1994) showed that there can be a lower personal observations.

### Seasonal and microhabitat variations in consumption of tough leaves

From one perspective, the differences documented in this study might be a bit surprising. In other words, if the nonriverine habitats show less plant diversity (Glander and Nisbett, 1996), we might expect animals in those habitats, such as those in group 1, to be forced to deal with a more limited series of options during the dry season and thus consume tougher leaves. However, the results show the opposite, with a riverine group (group 19) consuming tougher leaves in the dry season, and the nonriverine group (group 1) consuming tougher leaves in the wet season. These seasonal differences are mirrored by intergroup differences, with the riverine group (group 19) eating tougher leaves in the dry season than did the monkeys in the nonriverine group (group 1). This, however, for some plants, they are used to detach whole leaves, or groups of leaves, from the tree. If the point of detachment always has high toughness (as suggested here), then incisal bite forces will have to be high. In contrast, the molars will act on the full range of tissues, which will require a load that is more frequently lower than that on the incisors due to the aforementioned drop-off in toughness away from the basal end of leaves. This may have implications for the relative wear rates of incisors vs. molars: the former often show far more wear than do the latter on the howlers at La Pacifica.
ever, may ultimately be due to seasonal differences in leaf growth in the riverine vs. nonriverine habitats. During the dry season, all food trees in the nonriverine forests drop their leaves. Thus, in the dry season, group 1 would have no choice but to eat immature leaves. By contrast, many of the trees in the riverine habitats retain their leaves during the dry season, due to a better supply of moisture near the river. Thus, during the dry season, group 19 would have choices unavailable to group 1, and animals in group 19 take more mature leaves in the dry season as a result. Why that would be so, nutritionally, remains to be seen.

**Silica in leaves in the wet season**

We previously showed the presence of large quantities of very fine grit particles in the canopy at La Pacifica (Ungar et al., 1995). The high incidence of silica on leaves eaten during the wet season offers another possible cause of dental microwear in the howlers. Interestingly, the higher incidence of silica on leaves from the nonriverine habitat correlates with a higher incidence of microwear features previously documented from nonriverine groups at La Pacifica (Teaford and Glander, 1996). Clearly, a comparison of the incidence of phytoliths on leaves eaten during the wet and dry seasons in different microhabitats is now necessary.

**CONCLUSIONS**

1. Fracture toughness varies significantly by location within single leaves, and therefore, measures of fracture toughness must be standardized by location on food items. Thus, discussions about the toughness of a whole leaf may be misleading.

2. Monkeys may apparently make food choices based on fracture toughness, in some cases avoiding the toughest parts of leaves (such as the base of the leaf) and only consuming the least tough portions (such as the leaf tip).

3. Two forms of mechanical defense (toughness and silica content) are evidently at work in the plants at La Pacifica. The relative importance of phytoliths vs. grit in causing tooth wear will require more work, including in vitro studies of the effects of these abrasives on teeth.

4. Group 1, a nonriver group, ate tougher leaves in the wet season than in the dry season, and group 19, a riverine group, ate tougher leaves in the dry season than in the wet season, reflecting seasonal differences in food availability between these microhabitats.

**ACKNOWLEDGMENTS**

This research was supported by NSF grant DBC 9118876 to M.F.T. and K.E.G., and SBR 9601766 to M.F.T., K.E.G., and P.S.U. Support was also received from COSEN-PEW grants for undergraduate students working on the howler project. For access and continued accommodation, we express our thanks to Dr. Stephan Schmidheiny and the management of Hacienda La Pacifica. For continued help and support in Costa Rica, we thank Werner and Lily Hagnauer, Tony and Vreni Leigh, and Margaret Clarke. For their comments and sugges-

---

**TABLE 6. Chi-square analyses of incidence of silica on leaves from groups 1 and 19**

<table>
<thead>
<tr>
<th>Social group 1</th>
<th>With silica</th>
<th>Without silica</th>
<th>Row totals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Social group 19</td>
<td>15</td>
<td>10</td>
<td>25</td>
</tr>
<tr>
<td>Column totals</td>
<td>18</td>
<td>20</td>
<td>38</td>
</tr>
</tbody>
</table>

1 Chi-square = 4.68 (P < 0.05).
tions on the manuscript, we thank Clark Larsen and two anonymous reviewers.

LITERATURE CITED


