Molar form, enamel growth, and durophagy in *Cercocebus* and *Lophocebus*

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**Abstract**

**Objectives:** To test the hypothesis that differences in crown structure, enamel growth, and crown geometry in *Cercocebus* and *Lophocebus* molars covary with differences in the feeding strategies (habitual vs. fallback durophagy, respectively) of these two genera. Relative to *Lophocebus* molars, *Cercocebus* molars are predicted to possess features associated with greater fracture resistance and to differ in enamel growth parameters related to these features.

**Materials and Methods:** Sample proveniences are as follows: *Cercocebus atys* molars are from the Taï Forest, Ivory Coast; *Lophocebus albigena* molars are from a site north of Makoua, Republic of Congo; and a *Lophocebus atterimus* molar is from the Lomako Forest, Democratic Republic of Congo. For μCT scans on which aspects of molar form were measured, sample sizes ranged from 5 to 35 for *Cercocebus* and 3 to 12 for *Lophocebus*. A subsample of upper molars was physically sectioned to measure enamel growth variables.

**Results:** Partly as a function of their larger size, *Cercocebus* molars had significantly greater absolute crown strength (ACS) than *Lophocebus* molars, supporting the hypothesis. Greater crown heights in *Cercocebus* are achieved through faster enamel extension rates. Also supporting the hypothesis, molar flare and proportional occlusal basin enamel thickness were significantly greater in *Cercocebus*. Relative enamel thickness (RET), however, was significantly greater in *Lophocebus*.

**Discussion:** If ACS is a better predictor of fracture resistance than RET, then *Cercocebus* molars may be more fracture resistant than those of *Lophocebus*. Greater molar flare and proportional occlusal basin thickness might also afford *Cercocebus* molars greater fracture resistance.

**KEYWORDS**

absolute crown strength, enamel thickness, fracture risk, hard-object feeding, teeth
1 | INTRODUCTION

The group of African monkeys referred to as mangabeys comprises species from two clades: Cercocebus and Lophocebus (Cronin & Sarich, 1976; Disotell, 1994; Groves, 1978; Harris, 2000). The two mangabey genera share several morphological and behavioral features, and it was this convergence that obscured the true phylogeny of African papionins for years (Fleagle & McGraw, 1999, 2002; Groves, 2000; Harris & Disotell, 1998; Jolly, 2001). Although similar in that they are both thickly enameled (Daegling et al., 2011; Lambert et al., 2004; McGraw et al., 2012; McGraw et al., 2014), Lophocebus and Cercocebus dentitions differ in the degree to which their premolars are molarized (Fleagle & McGraw, 1999). The premolars of Cercocebus are larger relative to their molars than are those of Lophocebus, a trait that proved informative for illuminating the diphyletic origins of these genera (Fleagle & McGraw, 1999). This difference in relative premolar size may also relate to different feeding regimes: the habitual consumption of hard foods in Cercocebus (Daegling et al., 2011; McGraw et al., 2012, 2014) versus the fallback consumption of hard foods in Lophocebus (Lambert et al., 2004).

Although the hardness of the foods these two genera consume has not been directly compared using the same methods, Cercocebus is believed to masticate harder foods than does Lophocebus (McGraw & Daegling, 2020). Moreover, these genera are known to eat hard foods in very different contexts and employ different oral processing behaviors: Lophocebus consumes hard foods as dietary fallbacks, when softer, preferred foods are unavailable (Lambert et al., 2004). Cercocebus is an habitual hard-object feeder that uses its posterior teeth to crush large, hard Saccoglottis seed casings not only when softer fruits are unavailable, but throughout the year (Daegling et al., 2011; McGraw et al., 2012, 2014).

Cercocebus’ more frequent consumption of hard foods would expose its molars to greater opportunities for fracture, and subsequent failure, over time. In addition, although enamel is more resistant than bone to crack propagation (Bajaj et al., 2008), like bone, enamel can suffer fatigue failure under cyclic loading (Gao et al., 2016). Data do not exist that would allow us to quantify differences between Cercocebus and Lophocebus in terms of the frequency and magnitude of the loads their molars experience, but more frequent exposure to high magnitude loading would put Cercocebus molars at greater risk of fatigue failure. We also note that Lophocebus, in contrast to Cercocebus, uses its incisors more often during ingestion and has deeper mandibular corpora, consistent with greater incisor use (McGraw & Daegling, 2020). For all these reasons, we suggest that Cercocebus molars are subject to greater risk of failure over their lifetimes.

Beyond a general similarity in having thick enamel, the molars of Lophocebus and Cercocebus have yet to be systematically compared in terms of structural features (e.g., enamel thickness, the distribution of enamel across the crown, and absolute crown strength [ACS; see below]), enamel growth correlates of these structural features, and aspects of crown geometry (e.g., degree of molar flare; see below). The goal of this study is to assess whether differences between the molars of these two genera in structure, growth and crown geometry are consistent with known divergences in their feeding strategies (habitual vs. fallback durophagy) and oral processing behavior.

2 | STRUCTURAL FEATURES AND THEIR ENAMEL GROWTH CORRELATES

Despite the centrality of these two mangabey lineages to arguments about the relationship of enamel thickness to fallback and habitual durophagy (Lambert et al., 2004; McGraw et al., 2012, 2014), their enamel thickness in known from only a handful of specimens for each genus (McGraw et al., 2012). Thus, it is not yet clear if molars of these two genera are endowed with similar degrees or distributions of enamel thickness. The present study uses 2D virtual sections from 3D reconstructions of μCT renderings in the largest sample of mangabey molars evaluated yet to compare a suite of molar geometric and structural features that have been purported to relate to durophagy.

Historically, functional inferences of molar form rely heavily on the single metric of relative enamel thickness (RET), first proposed by Martin (1985). By incorporating a dental proxy for differences in body size, RET is thought to reflect dietary adaptation more accurately than do measures of absolute enamel thickness (Martin, 1985). High values of RET are presumed to reflect the ability of a crown to resist fracture when consuming hard foods and/or forestall dentine exposure caused by consumption of abrasive foods (Molnar & Gant, 1977; Pampush et al., 2013; Rabenold & Pearson, 2011; Strait et al., 2013; Vogel et al., 2008). Thus, RET is related not only to food hardness but also to the abrasiveness of foods (Pampush et al., 2013; Rabenold & Pearson, 2011). Perhaps in part because RET relates to both food hardness and abrasiveness, Thiery et al. (2017) did not find a correlation between durophagy and RET in a sample of upper second molars drawn from 32 primate species.

Furthermore, it has recently been argued that RET is not as accurate a measure of how teeth withstand occlusal loads as a newly introduced metric, ACS, calculated as the product of tooth crown’s radius and its average enamel thickness (AET; Schwartz et al., 2020). Using an engineering approach, Schwartz et al. (2020) found that ACS more closely approximates a tooth’s fracture resistance than does RET, consistent with previous studies suggesting that absolute enamel thickness is directly related to crown strength (Lawn et al., 2009; Lawn & Lee, 2009; Lucas et al., 2008). Schwartz et al. (2020) suggest that RET, by contrast with ACS, is only “moderately informative” about fracture resistance and may even provide “misleading information.” For example, gorillas have relatively thin enamel, but their large tooth size endows them with high ACS (Schwartz et al., 2020). It is their high ACS values, despite their low RET values, that afford gorilla molar crowns a high degree of fracture resistance consistent with the discovery of seasonal hard-object feeding in gorillas from Loango National Park, Gabon (van Casteren et al., 2019).

Given the close relationship between food material properties and tooth fracture mechanics, here we also compare Cercocebus and Lophocebus in terms of ACS. Specifically, we hypothesize that ACS, but not necessarily RET, will be greater in Cercocebus than in
Lophocebus, especially if Cercocebus molars have been selected to sustain high frequency, high magnitude loads without failing. Similarly, if Cercocebus molars simply afford them the ability to eat hard foods throughout the year (i.e., if they are exapted to their demanding feeding regimes), then they would also be expected to have greater ACS, but not necessarily RET, than do the molars of Lophocebus.

Although Thierry et al. (2017) did not find a relationship between RET and durophagy, these authors did find a relationship between the distribution of enamel over molar crowns and durophagy within Cercopithecoidea. Cercopithecoids that consumed hard objects had a more uneven distribution of enamel than did nondurophagous cercopithecoids (Thierry et al., 2017). Lucas et al. (2008) predicted that more unevenly distributed enamel would be found in durophagous primates, reflecting reinforcement of enamel in regions of the crown most likely to fail during hard-object feeding.

Reinforcement of enamel at the cusp tip, in particular, has been suggested to relate to the mastication of large, hard food objects like seeds and nuts (Lucas et al., 2008). Hard objects can induce subsurface cracks at cusp tips (i.e., radial cracks; Lawn & Lee, 2009) when teeth are brought together with high force during crushing (Lucas et al., 2008). Radial cracks initiate at the enamel-dentine junction beneath the cusp, where tensile stress is concentrated during hard food mastication (Lawn & Lee, 2009). Once initiated, radial cracks can develop into “ribbon fissures” that extend downward along cusp “shoulders” and through to the enamel surface, leading to catastrophic failure (Lawn & Lee, 2009). Such cracks are more likely to extend to the enamel surface when enamel is thin, causing it to fracture (Lucas et al., 2008). On this basis, Lucas et al. (2008) predicted that durophagous species should have relatively thicker enamel at their cusp tips. Unfortunately, we determined that measures of linear cusp tip thickness had low repeatability in our study (10%–12% error on average), and therefore do not include these measurements here. We do, however, include linear measures of occlusal basin thickness, which were more reliable. Given their intimate and reciprocal relationship—because cusps crush food in opposing occlusal basins—durophagy has been suggested to be associated with particularly thick occlusal basins, as appears to be the case in orangutans (Kono, 2004). O’Hara (2021) found that several measures of occlusal basin thickness scaled to AET (Martin, 1985) were associated with durophagy in extant carnitremes, including Cercocebus atys.

We also examined enamel thickness along the lateral wall of functional cusps, under the assumption that thicker lateral wall enamel offers greater resistance to margin fracture. Margin cracks can result from biting on either hard or soft foods and are initiated at the enamel-dentine junction of the crown base (cervical margin) as coronal dentine expands under vertical compression (Lawn & Lee, 2009). As the crown base bulges, it is subject to tensile “hoop” stresses that promote crack extension upward along the enamel walls. Like radial cracks, margin cracks can develop into ribbon fissures, leading to catastrophic failure. Again, we expected the habitually durophagous Cercocebus to have proportionally greater functional side lateral wall thickness than does Lophocebus.

We further hypothesized that potential differences between Cercocebus and Lophocebus in AET or crown size, both of which contribute to ACS, would be linked to differences in enamel growth. We therefore investigated potential differences in enamel growth variables such as daily secretion rates (DSRs), enamel extension rates (EERs), and overall enamel formation times. Thicker enamel can be produced by faster DSRs, longer crown formation times, or both (Grine & Martin, 1988). Larger crowns—specifically those with greater crown heights—can result from increases in EERs, crown formation time, or both (Guatelli-Steinberg et al., 2012).

### 3 | ASPECTS OF CROWN GEOMETRY

With respect to aspects of crown geometry, we evaluated the association of cuspal geometry with habitual hard-object consumption by quantifying cusp tip angles and degree of cusp flare. Blunt, low cusps (i.e., those with more obtuse cusp tip angles) are suggested to be less subject to breakage than tall sharp cusps when eating hard foods (Kay, 1978). We also note that blunt cusps are also expected to function more efficiently than tall, sharp cusps when cracking hard objects and reducing them through grinding (Constantino et al., 2009; Jolly, 1970; Kay, 1978, 1981; Luke & Lucas, 1983).

Cusp flare (Figure 1) refers to the angulation of the cusp from cervix to cusp tip (Shimizu, 2002; Singleton, 2003; Macho & Shimizu, 2009). In terms of function, Singleton (2003) found an association between molar flare and hard-object feeding in contemporary hominoids. More specifically, it has been suggested that molar flare might buttress crowns against laterally directed forces during chewing (Macho & Shimizu, 2009). However, Macho and Shimizu (2009) suggested that lateral flare is not necessarily expected in a dedicated hard-object feeder: omnivorous macaques, for instance, exhibit lateral flare. They also cited Woda et al. (2006) who found that in humans it is primarily jaw muscle activity, not jaw kinematics, that is related to food hardness. More recent experimental studies in humans, however, suggest that while muscular activity increases as a function of food hardness, so too does lateral jaw movement (Almotairy et al., 2021; Kitashima et al., 2015; Komino & Shiga, 2017). Thus, cusp flare may serve as a buttress against laterally-directed forces during the crushing or grinding of hard foods. Here we examined “functional” cusp flare, where functional cusps are those that are involved in Phase II of the chewing cycle (Kono, 2004; Schwartz, 2000). Cercocebus molar are expected to exhibit greater functional cusp flare, and thus stronger buttressing, than those of Lophocebus. Table 1 summarizes the predictions tested in this study.

### 4 | MATERIALS AND METHODS

#### 4.1 | Sample provenience and preparation

The C. atys sample used in this study derives from the Tai Forest of the Ivory Coast, collected by W.S.M. and members of the Tai Forest
The C. atys diets consist mostly of hard-foods, especially Sacoglottis gabonensis, which has seed casings with twice the hardness of cherry pits (Daegling et al., 2011). C. atys processes S. gabonensis seeds year-round, by placing them immediately posterior to their canines and crushing them on their premolars and molars with a powerful isometric bite (Daegling et al., 2011; McGraw et al., 2011, 2014).
nearby Dja Reserve in southcentral Cameroon (Poulsen et al., 2001; Poulsen et al., 2002), and Uganda’s Kibale Forest (Lambert et al., 2004; Olupot et al., 1997; Waser, 1984). At each of these sites, *L. albigena* prefers fruit but switches to seed-eating when fruit is scarce. For example, during the dry season, *L. albigena* at Lope rely on seeds of *Pentaclethera macrophylla* (Ham, 1994), which are protected by hard and tough pods (McGraw et al., 2016). Based on our reading of the literature, there is no indication that *L. albigena* is consuming hard-object foods year round, but rather is doing so seasonally. Because we are interested in generic differences (i.e., differences between Cercocebus and Lophocebus) and because our *L. albigena* sample is relatively small compared to our *C. atys* sample, we also include one *L. aterrimus* specimen from Democratic Republic of Congo’s Lomako Forest (McGraw, 1994).

Table 2 gives the specimen numbers and tooth types used in the current study. Not all specimens were used in every analysis; sample sizes for various measurements and statistical comparisons are provided in the Results section. Choice of right or left antimere was based on which crown appeared least worn. Sex was known for only a portion of the sample. To maintain adequate sample sizes, sex was not included as a covariate in our analyses, and as a result, remains an unknown source of potential variation. All teeth were manually extracted from maxillae and mandibles and sterilized with UV light.

### 4.2 μCT methods

Extracted teeth were scanned using a Bruker SkyScan 1172 High Resolution Ex Vivo 3D X-Ray Tomography Scanner located in the Do-Gyoon Kim Laboratory at the OSU College of Dentistry. The first ten teeth were scanned at a resolution of 13 μm, but since the boundary between enamel and dentine was clearly differentiable at 22 μm, the

| TABLE 2 Full dental sample |
|-----------------------------|
| **Species** | **Individual** | **Sex** | **Tooth types** |
| *C. atys* | TF 16–5 | M | UM1, UM2, LM1, LM2 |
| TF 16–9 | M | UM3 |
| TF 16–11 | F | UM3 |
| TF 22–26 | M | UM2, UM3, LM2, LM3 |
| TF 22–29 | M | UM2, UM3, LM3 |
| TF 22–46 | M | UM2, UM3 |
| TF 23–10 | M | UM2, UM3 |
| TF 24–3 | F | UM2, LM1 |
| TF 94–7 | F | UM2 |
| TF 94–9 | M | UM2, UM3, LM3 |
| TF 94–25 | M | UM1, UM2 |
| TF 2001 | F | UM1, UM2 |
| TF 2008 | F | UM3 |
| TF 2010–1 | F | UM1 |
| TF 2010–2 | F | UM2, UM3 |
| TF 2016 | F | UM2, UM3 |
| TF 2019 | M | UM1, UM2 |
| TF 2020 | M | UM1, LM1 |
| TF 2040 | M | UM1 |
| TF 2041 | M | UM1 |
| TF 2106 | M | UM3, LM3 |
| TF 2108 | M | UM2, UM3, LM3 |
| TF 2138 | M | UM3 |
| TF 22–46 | M | UM1, UM2 |
| TP-91 | M | UM2 |
| *L. albigena* | 85–1 | F | UM1, UM2, UM3, LM1, LM2, LM3 |
| 85–7 | M | UM1, UM2, UM3, LM1, LM2, LM3 |
| 85–17 | M | UM1, UM2, UM3, LM1, LM2 |
| 642 | M | UM1, UM2, UM3, LM1, LM2, LM3 |
| *L. aterrimus* | 81–7 | F | UM2 |
latter resolution was used for the remainder of the sample. RAW output files were processed with N.Recon v1.7.4.2 and then saved as TIFFs. Two-dimensional bucco-lingual planes of section (mesial and distal) were generated from 3D digital renderings of the teeth (following Skinner et al., 2015) using Dragonfly v.2021.1.0.977, passing through buccal and lingual dentine horns and perpendicular to the cervical margin. Each virtual section was then saved as a TIFF and imported into Adobe Photoshop where crown outlines were reconstructed (when necessary) prior to performing measurements.

Measurement reference lines and points are shown in Figure 1. AET was calculated as the enamel cap area divided by the EDJ length (Martin, 1985), while RET was calculated as AET divided by the square root of the dentine-pulp area bordered by the EDJ and bicervical diameter (BCD) and multiplied by 100 (Martin, 1985). ACS was calculated as the square root of the product of the coronal dentine radius (half of the BCD) and AET (Schwartz et al., 2020). Linear enamel thickness of the occlusal basin was measured as the distance between the lowest point of the occlusal basin at the EDJ and the lowest point of the occlusal basin at the outer enamel surface, or OES (Kono-Takeuchi et al., 1998; Macho & Berner, 1994; Macho & Thackeray, 1992, 1993; Olejniczak & Grine, 2006; Schwartz, 1997, 2000). Maximum lateral wall enamel thickness was measured at the widest point between the EDJ and OES, along a line perpendicular to the EDJ (Kono et al., 2002; Schwartz et al., 2020; Spoor et al., 1993; Suwa & Kono, 2005; Ulhaas et al., 1999). Proportional linear occlusal basin and lateral enamel wall thicknesses were obtained by dividing each linear measure by AET (O’Hara, 2021). Thus, “proportional thickness” scales the linear thickness of enamel in each of these regions (occlusal basin and lateral enamel wall) to the average thickness of enamel for a crown. Measures greater than 1 represent relatively thicker enamel in these regions.

To measure cusp tip angles, a line parallel to the BCD and tangent to the lowest point on the EDJ of the occlusal basin was drawn (Figure 1). Cusp tip angles were measured as the included angle between where that line intersected the OES and the angle’s vertex (Figure 1). To perform measurements of cusp flare, 2D slices were rotated in ImageJ so that their BCDs were horizontal; then, a line perpendicular to the BCD was drawn. Lastly, a connecting line from the CEJ to the apex of the cusp (or reconstructed cusp) on the OES was drawn (Shimizu, 2002). The angle between this line and the line perpendicular to the BCD was the angle measured for flare (Figure 1).

While there were some unworn crowns in the sample, most exhibited varying degrees of wear. Worn crowns were reconstructed following the recommendations of O’Hara and Guatelli-Steinberg (2021). These authors found that for AET and crown height measurements, crowns with limited wear (for which wear did not reach the dentine horns and/or the deepest point of the occlusal basin) accurate values were achievable with the Profile (Grine & Martin, 1988; Smith et al., 2011; Smith et al., 2012), Polynomial (Modesto-Mata et al., 2017), and Pen Tool methods (Guatelli-Steinberg et al., 2009; O’Hara et al., 2019; Saunders et al., 2007). O’Hara and Guatelli-Steinberg (2021) also found that, for crowns on which wear exposed the tips of dentine horns (what they termed “extensive wear”), it was still possible to obtain accurate AET values using the Profile or Polynomial methods. The O’Hara and Guatelli-Steinberg (2021) recommendations were followed here for AET, RET, and measurements of molar flare using the Profile method on teeth with extensive wear and incorporating the Pen Tool method if a reference tooth was not available, but wear was limited. Maximum functional lateral wall linear measurements were possible with slight wear on cusps since the dentine horn was used as a reference point. No measurements of cusp tip angles or linear occlusal basin thickness were performed on crowns with any occlusal basin wear and/or on any reconstructed crowns.

All measurements on µCT slices were made by either Kaita Gurian or Jess Rychel, whose inter-observer error was assessed. Forty-six mesial slices were measured by both Kaita Gurian and Jess Rychel for crown area, bicervical diameter, EDJ length, and maximum functional cusp linear thickness. For all these measures, average interobserver error ranged from 0.5% to 4.6%. Linear occlusal basin thickness measurements on 23 slices averaged 7.4% error, while errors for functional cusp tip angle on 24 slices and nonfunctional cusp tip angle on 23 slices averaged 2.8% and 2.9%, respectively. The higher error of the occlusal basin thickness measurements, we believe, reflects small variations in how the measurement line is angled from the lowest point of the occlusal basin to the EDJ. Flare was measured by Kaita Gurian only, whose average intra-observer error for 15 teeth was 0.8%. For calculated values (ACS, AET, and RET) average interobserver error was less than 5%.

4.3 | Histological methods

After µCT scans were made, a subset of Cercocebus and Lophocebus upper second and third molars were physically sectioned across mesial cusps following steps outlined in Reid et al. (1998), with some modifications. Using the µCT slices, we marked the plane of section on the tooth and then embedded it in epoxy resin (Buehler™ Epoxicure). We used a Buehler IsoMet low-speed saw equipped with a 5-in. diameter diamond-wafering blade. Teeth were mounted on the saw using an irregular sample chuck in order to line up the desired section plane with the blade. Despite efforts to directly cut along the desired plane, we found that we usually were able to obtain cuts directly through only one of the dentine horns of the two mesial cusps. The side of the cut block with the sharpest dentine horn was mounted on a slide with Gorilla™ epoxy and a second cut was made parallel to the first. The specimen was then ground using a target holder on a series of Buehler CarbiMet abrasive papers with successively finer grit until a thickness of approximately 100 μm was reached and microstructures (e.g., cross striations) were visible. The sample was then polished using Buehler alumina micropolish, after which the slide was dehydrated in an ethanol series and rinsed in the clearing agent Histoclear. The slide was then cover-slipped using immersion oil and imaged with a Motic BA 310 Microscope fitted with a Moticam camera. Montages at 4× were made for measurements of EDJ lengths used in calculating enamel extension rates. All enamel growth measurements were made in ImageJ by one of us (Debbie Guatelli-Steinberg).
To measure daily secretion rates (DSRs), total EDJ lengths along the crown walls were measured and divided into thirds. In each third of the crown, viewed at 40x, measurements along enamel prisms were taken across six cross-striations (5 days) within 200 μm of the EDJ. Three such measurements were taken in each third of the crown and averaged to obtain a DSR value for each region. Three measurements were taken and averaged in the occlusal basin as well, as an indicator of whether there were differences between genera in the rate of enamel secretion in this region of the crown. This set of measurements was performed because, as will be seen below, the two genera differed in the proportional thickness of their occlusal basins.

Enamel extension rates were measured in the following methods described by Dean (2009, 2012) and Guatelli-Steinberg et al. (2012). Just lateral to the tip of the dentine horn, a point along the EDJ was identified. A 200-μm line was drawn from this point outward, along an enamel prism. Where that line stopped, another line parallel to or coincident with a stria of Retzius was followed back to the EDJ (Figure 2). The point where that line intersected the EDJ was marked, and the distance along the EDJ was measured between this point and the first point (i.e., the point near the dentine horn tip). Because striae of Retzius mark the enamel-forming front at a point in time, this segment of the EDJ represents the distance over which ameloblasts differentiated during the same length of time it took for the 200 μm prism segment to form. That length of time varied depending on each third of the crown. For example, in the cuspal third of the crown, DSRs were faster than they were in the cervical third, such that the length of time it took to form the 200 μm prism length was shorter in the cuspal region than in the cervical region. The amount of time it took to form the first EDJ segment is 200 μm divided by the cuspal DSR. For example, if the cuspal DSR were 4 μm/day, then the time it took to form this first EDJ segment was 50 days. To obtain the rate of enamel extension corresponding to this segment, the EDJ segment length would then be divided by 50 days.

This “zigzag” procedure was followed down the full length of the EDJ along the crown walls (see Figure 2), with the appropriate DSR used in the enamel extension rate calculation based on crown location. In three of 26 cusps, there was a small portion (of approximately 200 μm or less of the EDJ missing at the dentine horn tip) owing to wear. In these three cases, the dentine horn was reconstructed (profile method) and a 200 μm line parallel with the first prism was drawn from the first point of the reconstructed EDJ toward the outer enamel surface. A stria of Retzius was followed from the EDJ back to the end of that line, establishing the boundaries of the first measurement.

To obtain total enamel formation time, the time it took to form each segment length was summed along the length of the entire EDJ. However, it is not possible to draw a 200 μm line along a prism near the cervix as enamel thins extensively in this region. To estimate the length of time it took to form the remainder of the crown (see Figure 2), the enamel extension rate of the preceding segment was applied to the remaining length, and then added to the summed enamel formation times of the preceding EDJ segments.

Measurement error was assessed on four molar crowns (two Lophocebus; two Cercocebus). For DSRs, there were 12 areas on these crowns where DSR had previously been measured three times and then averaged. For the 12 areas, the average DSRs from the first and second round of measurements differed by 2.1%. For EERs, 30 segments were remeasured and the new DSRs from the second round were applied to them. On average, EERS differed between the first and second round by 3.5%. The four total enamel formation times for these teeth differed between the first and second round by 1.3%.

4.4 | Statistical methods

Graphs were generated in SYSTAT version 13. Statistical analyses were carried out in SAS v. 9.4 (SAS Institute, 2015). Proc Mixed (the Mixed Procedure) was used to analyze differences between Cercocebus and Lophocebus in RET and ACS and was also used for the analysis of enamel growth variables. This procedure fits mixed linear models to
data, allowing a repeated measures analysis that was useful when using measurements from the same individual (e.g., more than one tooth type; more than one enamel extension rate measure). AET and tooth size were not analyzed statistically, as both are incorporated into ACS. The variables RET and ACS were modeled as a function of genus, tooth type, and the interaction of tooth type and genus. Each individual was treated as a subject with tooth type as the repeated measure. Models were run with four different variance–covariance structures and results from models with the lowest AICs are reported here. Since there were 4 ways to obtain significance for RET (for genus, and the genus by tooth type interaction, for upper and lower teeth), a Bonferroni-corrected alpha value of 0.0125 was used; for the same reason the alpha value used for ACS was 0.0125. Bootstrapped t-tests were used to compare the two genera in daily secretion rates, functional cusp flare, proportional linear basin occlusal enamel thickness, proportional maximum functional lateral enamel thickness, and crown shape. Bonferroni corrections were used given that there were multiple tests of the same variable. For example, there were four tests of DSR differences between Cercocebus and Lophocebus, so an alpha value of 0.05/4 (or 0.0125) was used in this comparison.

### RESULTS

#### 5.1 RET and ACS

Summary statistics by tooth type for RET, AET (mm.), BCD (mm.) and ACS for upper molar mesial and distal slices (mean ± 1SD)

| Taxon   | Slice | Tooth | n  | RET       | AET       | BCD       | ACS       |
|---------|-------|-------|----|-----------|-----------|-----------|-----------|
| C. atys | Mesial| UM1   | 7  | 12.4 ± 1.4| 0.576 ± 0.07| 7.6 ± 0.9| 1.48 ± 0.15|
|         |       | UM2   | 15 | 13.3 ± 1.6| 0.715 ± 0.08| 8.5 ± 0.7| 1.74 ± 0.14|
|         |       | UM3   | 12 | 14.8 ± 1.6| 0.747 ± 0.05| 8.1 ± 0.7| 1.74 ± 0.09|
|         |       | All uppers | 34 | 13.6 ± 1.8| 0.697 ± 0.09| 8.2 ± 0.8| 1.68 ± 0.16|
|         | Distal| UM1   | 9  | 14.1 ± 1.7| 0.565 ± 0.10| 6.6 ± 1.1| 1.37 ± 0.24|
|         |       | UM2   | 14 | 15.1 ± 2.3| 0.706 ± 0.09| 7.9 ± 0.8| 1.66 ± 0.16|
|         |       | UM3   | 12 | 16.2 ± 2.0| 0.712 ± 0.07| 7.0 ± 1.0| 1.60 ± 0.16|
|         |       | All uppers | 35 | 15.2 ± 2.1| 0.672 ± 0.11| 7.2 ± 1.1| 1.60 ± 0.21|
| L. albigena | Mesial| UM1   | 4  | 15.8 ± 1.8| 0.611 ± 0.08| 6.3 ± 0.5| 1.39 ± 0.10|
|         |       | UM2   | 4  | 14.3 ± 0.7| 0.627 ± 0.03| 7.4 ± 0.4| 1.52 ± 0.06|
|         |       | UM3   | 4  | 15.7 ± 0.3| 0.643 ± 0.04| 6.7 ± 0.4| 1.47 ± 0.07|
|         |       | All uppers | 12 | 15.3 ± 1.2| 0.627 ± 0.05| 6.8 ± 0.6| 1.46 ± 0.09|
|         | Distal| UM1   | 2  | 15.4 ± 1.3| 0.591 ± 0.03| 6.3 ± 0.4| 1.37 ± 0.00|
|         |       | UM2   | 3  | 15.9 ± 1.3| 0.632 ± 0.03| 6.8 ± 0.6| 1.46 ± 0.04|
|         |       | UM3   | 4  | 18.7 ± 2.6| 0.643 ± 0.07| 5.5 ± 0.8| 1.30 ± 0.10|
|         |       | All uppers | 9  | 17.0 ± 2.3| 0.628 ± 0.05| 6.1 ± 0.9| 1.37 ± 0.10|
| L. atterimus | Mesial| UM1 | –  | –          | –          | –         | –          |
|         |       | UM2   | 1  | 14.8      | 0.649      | 6.5       | 1.45       |
|         |       | UM3   | –  | –          | –          | –         | –          |
|         |       | All uppers | 1  | 14.8      | 0.649      | 6.5       | 1.45       |
|         | Distal| UM1   | –  | –          | –          | –         | –          |
|         |       | UM2   | 1  | 15.4      | 0.606      | 6.3       | 1.38       |
|         |       | UM3   | –  | –          | –          | –         | –          |
|         |       | All uppers | 1  | 15.4      | 0.606      | 6.3       | 1.38       |

Abbreviations: ACS, absolute crown strength; AET, average enamel thickness; BCD, bicervical diameter; RET, relative enamel thickness.
### TABLE 4
Summary statistics by tooth type for RET, AET (mm.), BCD (mm.) and ACS for mesial and distal slices of lower molars (mean ± 1SD)

| Taxon       | Slice | Tooth n | RET ± 0.8 | AET ± 0.04 | BCD ± 0.07 | ACS ± 0.09 |
|-------------|-------|---------|-----------|------------|------------|------------|
| C. atys     | Mesial| LM1 4   | 12.2      | 0.526      | 5.9        | 1.24       |
|             |       | LM2 2   | 12.0      | 0.680      | 7.8        | 1.63       |
|             |       | LM3 5   | 14.5      | 0.732      | 7.3        | 1.74       |
|             |       | All lowers | 11       | 13.2       | 6.9        | 1.49       |
|             | Distal| LM1 2   | 13.1      | 0.499      | 5.2        | 1.13       |
|             |       | LM2 1   | 14.1      | 0.704      | 7.5        | 1.62       |
|             |       | LM3 4   | 15.8      | 0.755      | 6.7        | 1.59       |
|             |       | All lowers | 7        | 14.8       | 6.4        | 1.46       |
| L. albigena | Mesial| LM1 4   | 16.3      | 0.584      | 5.2        | 1.23       |
|             |       | LM2 4   | 15.4      | 0.656      | 6.5        | 1.46       |
|             |       | LM3 3   | 14.4      | 0.602      | 6.3        | 1.38       |
|             |       | All lowers | 11       | 15.5       | 6.0        | 1.35       |
|             | Distal| LM1 –   | –         | –          | –          | –          |
|             |       | LM2 2   | 14.8      | 0.632      | 6.8        | 1.48       |
|             |       | LM3 2   | 18.7      | 0.660      | 4.7        | 1.46       |
|             |       | All lowers | 4        | 16.8       | 6.1        | 1.36       |

Abbreviations: ACS, absolute crown strength; AET, average enamel thickness; BCD, bicervical diameter; RET, relative enamel thickness.

FIGURE 3 Box plots comparing Cercocebus and Lophocebus for RET and ACS for mesial and distal cusp μCT slices for all upper molars combined and all lower molars combined.
Measurements on occlusal basins could only be performed on completely unworn molars and maximum functional cusp wall linear thickness required molars with intact dentine horns. For these reasons, the data are limited across tooth types (Table 6). Figure 4 compares the two genera for proportional linear occlusal basin thickness and proportional maximum linear functional cusp wall enamel thickness and Table 7 reports the results of bootstrapped t-tests for these variables. For these tests, to optimize sample sizes, second and third molars for each jaw were combined. First molars were omitted since they were less well balanced across the two genera.
Proportional occlusal basin thickness is significantly greater for Cercocebus than it is for Lophocebus, while the two genera do not differ significantly in proportional maximum linear functional cusp wall enamel thickness.

5.3 | Enamel growth variables

Summary statistics for upper molar (Upper M2 and Upper M3 mesial slices) enamel growth variables are given in Table 8. Bootstrapped t-tests were performed for the “nonfunctional” buccal cusps, for which we had the largest sample: 12 Cercocebus and 4 Lophocebus. For DSRs, four tests were performed, with a Bonferroni-corrected alpha of 0.0125. None of the DSR differences between Cercocebus and Lophocebus were statistically significant at either the 0.0125 or 0.05 levels. Nonfunctional cusp enamel formation time (in days) was also not statistically significantly different between the two genera (p < 0.2689), although the two genera differed significantly in nonfunctional cusp EDJ length (p < 0.0006), with Cercocebus having greater EDJ length than Lophocebus. Thus, although the two genera differ significantly in EDJ length for their functional cusps, they do not have a statistically significant difference in their enamel formation times (although as can be seen in Table 8, numerically, Lophocebus has slightly shorter enamel formation times). This finding suggests that the longer EDJ lengths of Cercocebus are forming at a faster rate. This possibility is explored in the growth curve analysis of enamel extension rates.

**Table 7** Results of bootstrapped t-tests for proportional occlusal basin linear enamel thickness and proportional functional wall linear enamel thickness

| Variable                | Jaw    | Cercocebus (n) | Lophocebus (n) | Variances | Df   | T value | p-Value |
|-------------------------|--------|----------------|----------------|-----------|------|---------|---------|
| Proportional Occlusal Basin ET | Upper  | 16             | 4              | Equal     | 18   | 3.37    | 0.0030  |
|                         | Lower  | 5              | 3              | Equal     | 6    | 3.76    | 0.0094  |
| Proportional Functional Wall ET | Upper  | 26             | 9              | Equal     | 33   | 1.15    | 0.2604  |
|                         | Lower  | 7              | 7              | Equal     | 12   | −1.11   | 0.2908  |

Abbreviations: ET, enamel thickness. Note: Boldface denotes statistical significance.

**Table 8** Summary statistics for enamel growth variables for upper molars (mean ± 1SD)

| n       | Occlusal basin DSR (days per μm) | Cuspal DSR (days per μm) | Mid-crown DSR (days per μm) | Cervical DSR (days per μm) | Initial EER (μm/day) | Enamel formation time (days) |
|---------|---------------------------------|--------------------------|-----------------------------|---------------------------|----------------------|-----------------------------|
| Lingual cusp |                                 |                          |                             |                           |                      |                             |
| Cercocebus | 8                               | 3.9 ± 0.4                | 3.7 ± 0.4                   | 4.1 ± 0.1                 | 3.5 ± 0.3            | 40.2 ± 5.1                 | 738 ± 69                    |
| Lophocebus | 2                               | 4.2 ± 0.5                | 4.5 ± 0.4                   | 3.9 ± 0.4                 | 3.2 ± 0.3            | 29.5 ± 9.6                 | 662 ± 69                    |
| Buccal cusp |                                 |                          |                             |                           |                      |                             |                             |
| Cercocebus | 12                              | 3.9 ± 0.3                | 3.7 ± 0.3                   | 4.1 ± 0.3                 | 3.6 ± 0.4            | 32.4 ± 5.3                 | 626 ± 82                    |
| Lophocebus | 4                               | 4.3 ± 0.6                | 4.01 ± 0.2                  | 4.2 ± 0.4                 | 3.3 ± 0.3            | 22.7 ± 2.0                 | 571 ± 80                    |

Abbreviations: DSR, daily secretion rate; EER, enamel extension rate.
Growth curves were modeled in Proc Mixed for upper second and third molars combined for buccal (nonfunctional) cusps. Sample sizes were too small for statistical comparison for the lingual (functional) cusps; however, both nonfunctional and functional cusp data are plotted in Figure 5.

For the Proc Mixed growth comparison of nonfunctional cusps, there were two M2s and two M3s for *Lophocebus* and seven M2s and five M3s, for *Cercocebus*. The first nine enamel extension rate measures (interval 1 through interval 9) were treated as repeated measures for each individual, modeled as a quadratic, with the null model likelihood tests statistically significant (*p* < 0.0254). Individual variation was treated as a random effect. One individual had two molars in this test; the rest were represented by a single molar each. Tests for interval and genus are shown in Table 9. All predictors were statistically significant, including genus and the interval*genus predictors, indicating that the growth curves of these species differ. This difference can be seen in Figure 5, which plots mean enamel extension rates for upper and lower molars. Note that *Cercocebus* starts at a much higher rates than *Lophocebus* and maintains a slightly higher rate at most later time intervals. That is the basis for the significant effect of genus and the interaction of genus and interval in the mixed linear model results.

### Table 9: Analysis of nonfunctional cusp enamel extension rates (EERs)

| Effect               | Numerator df | Denominator df | F value | p Value |
|----------------------|--------------|----------------|---------|---------|
| Interval             | 1            | 14             | 281.49  | <0.0001 |
| Genus                | 1            | 107            | 9.54    | 0.0026  |
| Interval*Genus       | 1            | 107            | 5.4     | 0.0220  |
| Interval*Interval    | 1            | 107            | 178.3   | <0.0001 |

Note: Boldface denotes statistical significance.

Growth curves were modeled in Proc Mixed for upper second and third molars combined for buccal (nonfunctional) cusps. Sample sizes were too small for statistical comparison for the lingual (functional) cusps; however, both nonfunctional and functional cusp data are plotted in Figure 5.

For the Proc Mixed growth comparison of nonfunctional cusps, there were two M2s and two M3s for *Lophocebus* and seven M2s and five M3s, for *Cercocebus*. The first nine enamel extension rate measures (interval 1 through interval 9) were treated as repeated measures for each individual, modeled as a quadratic, with the null model likelihood tests statistically significant (*p* < 0.0254). Individual variation was treated as a random effect. One individual had two molars in this test; the rest were represented by a single molar each. Tests for interval and genus are shown in Table 9. All predictors were statistically significant, including genus and the interval*genus predictors, indicating that the growth curves of these species differ. This difference can be seen in Figure 5, which plots mean enamel extension rates for upper and lower molars. Note that *Cercocebus* starts at a much higher rates than *Lophocebus* and maintains a slightly higher rate at most later time intervals. That is the basis for the significant effect of genus and the interaction of genus and interval.

### 5.4 Cusp tip geometry and molar flare

Measurements on cusp tips could only be performed on completely unworn molars, while functional cusp flare required molars with intact dentine horns. Table 10 provides descriptive statistics for cusp tip angle and functional cusp flare in degrees.

As there were very few specimens with completely unworn functional cusp tips, no statistical comparisons were attempted for this variable, although Figure 6 suggests a tendency for *Cercocebus* to have sharper cusps than *Lophocebus*. For functional cusp flare, second and third molars for each jaw were combined to optimize sample sizes for bootstrapped t-tests comparing *Cercocebus* and *Lophocebus*. First molars were omitted since they were not well balanced across the two genera. There were two t-tests performed (upper and lower dentitions), so an alpha value of 0.025 was used. Table 11 summarizes results of the t-tests for molar flare. There was a statistically significant difference between these genera in functional cusp flare for lower molars, with *Cercocebus* exhibiting greater flare, but no significant difference was found for upper molars.

### 6 Discussion and conclusion

This study investigated the hypothesis that compared to the molars of *Lophocebus*, *Cercocebus* molars would possess features that afford them greater resistance to fracture, and that these features would have underlying enamel growth correlates. This hypothesis was partially supported by our analyses (see Table 12). The functionally relevant features of molar anatomy that supported our hypothesis were the following: *Cercocebus* exhibited greater values of ACS (upper molars), functional cusp flare (lower molars), and proportional occlusal basin enamel thickness (both upper and lower molars). At the same
time, *Cercocebus* possessed lower RET values than *Lophocebus* and displayed a tendency to exhibit sharper cusp tips (there were, however, insufficient sample sizes to perform statistical tests for this feature). Proportional functional lateral wall enamel thickness did not differ significantly between the two genera. Some of the differences between the molars of *Cercocebus* and *Lophocebus* can be seen in the μCT images shown in Figure 7.

In terms of enamel growth, no significant differences were found between genera in enamel daily secretion rates or crown formation times; however, extension rates in *Cercocebus* were significantly greater than they were in *Lophocebus*. *Cercocebus* had significantly longer EDJ lengths than *Lophocebus* and therefore appears to grow its longer EDJs primarily through faster rates of enamel extension rather than by significantly prolonging crown formation time. These functional and enamel growth details appear consistent with one another: larger (both taller and wider) crowns that are not significantly different in daily enamel secretion rates and crown formation time could be expected to have lower RET, all other factors being equal.

### Table 10

descriptive statistics for cusp tip angle and functional cusp flare in degrees (mean ± 1SD)

| Taxon       | Tooth | Functional cusp tip angle (°) | Functional cusp flare (°) |
|-------------|-------|-------------------------------|---------------------------|
| *C. atys*   | UM1   | 100.0 (1)                     | 28.5 ± 5.1 (6)            |
|             | UM2   | 88.1 ± 12.0 (8)               | 28.4 ± 3.6 (13)           |
|             | UM3   | 89.5 ± 8.1 (8)                | 29.3 ± 4.0 (12)           |
|             | All uppers | 89.5 ± 10.0 (17)         | 28.7 ± 4.0 (31)           |
|             | LM1   | 92.0 ± 6.4 (2)                | 31.9 ± 6.5 (4)            |
|             | LM2   | 68.5 (1)                      | 39.9 ± 1.6 (2)            |
|             | LM3   | 92.8 ± 7.7 (3)                | 36.1 ± 4.3 (5)            |
|             | All lowers | 88.5 ± 11.3 (6)         | 35.3 ± 5.4 (11)           |
| *L. albigena* | UM1   | –                             | 32.3 (1)                  |
|             | UM2   | 111.4 (1)                     | 28.5 ± 1.9 (4)            |
|             | UM3   | 98.7 (1)                      | 25.8 ± 2.5 (4)            |
|             | All uppers | 105.1 ± 9.0 (2)          | 27.7 ± 2.9 (9)            |
|             | LM1   | –                             | –                         |
|             | LM2   | –                             | 34.4 ± 1.94 (2)           |
|             | LM3   | 99.5 (1)                      | 27.4 ± 2.65 (3)           |
|             | All lower | 99.5 (1)                  | 30.2 ± 4.4 (5)            |

### Table 11

descriptive statistics for cusp tip angle and functional cusp flare in degrees (mean ± 1SD)

| Variable                | Jaw    | *Cercocebus* (n) | *Lophocebus* (n) | Variances | Df   | T value | p-value   |
|-------------------------|--------|------------------|------------------|-----------|------|---------|-----------|
| Functional cusp flare   | Upper  | 25               | 8                | Equal     | 31   | 1.16    | 0.2568    |
|                         | Lower  | 7                | 5                | Equal     | 10   | 2.88    | 0.0165    |

Note: Boldface denotes statistical significance.
Not all features of Cercocebus molars appear to provide greater fracture resistance than those of Lophocebus. This finding begs the question of whether it is the combination of features within each genus that might produce differences in how well their molars resist fracture. To address this question, molars with the features described here could be modeled using finite element analysis to assess their performance under a variety of loading regimes (e.g., Kupczik & Lev-Tov Chattah, 2014). Furthermore, this study only examined molars, yet premolars are obviously involved in mastication as well, and the P4s of Cercocebus are larger relative to their first molars than those of Lophocebus (Fleagle & McGraw, 1999). Few of the features examined in the present molar study have which been explored in mangabey premolars. Lastly, decussation, the crisscrossing of enamel prisms that resists crack propagation (Bajaj et al., 2008), was not examined in the present study. Comparison of the two mangabey genera in terms of degree of decussation would also be relevant to fully understanding how their molars resist fracture and might be especially relevant to the problem of fatigue stress (Gao et al., 2016).

Nevertheless, the fact remains that both genera can and do eat very hard objects. The present investigation aligns with what recent studies have suggested: there is more than one way to crack obdurate foods (Constantino et al., 2011; Thiery et al., 2017; van Casteren et al., 2019). Sea otters (Enhydra lutris) crack the shells of clams, mussels, and abalone with thin enameled but large, bunodont molars (Constantino et al., 2011). In Loango National Park, Gabon, western lowland gorillas (Gorilla gorilla), whose molars have thin enamel, consume hard seeds of Coula edulis throughout its annual fruiting season. In both of these mammals, large overall tooth size appears to compensate for relatively thin enamel. Indeed, despite their relatively thin enamel, gorillas have the highest ACS values among all living hominoids (Schwartz et al., 2020). It is therefore conceivable that for Cercocebus and Lophocebus, a similar dynamic is at work, such that the larger teeth of Cercocebus, which have greater ACS than Lophocebus, do not require relatively thicker enamel. In a reciprocal fashion, the relatively thicker enamel of Lophocebus molars may compensate for smaller size.

Proportionally thick occlusal enamel in Cercocebus relative to Lophocebus suggests that the former is better protected against fracture in this region of the tooth. An uneven distribution of enamel was predicted for primates consuming durophagous, as opposed to

### Table 12 Summary of results relative to predictions

| Feature                                      | Prediction          | Lower molar results                            | Upper molar results                      |
|----------------------------------------------|---------------------|------------------------------------------------|------------------------------------------|
| Relative enamel thickness (RET)              | Cercocebus Lophocebus | Cercocebus ≈ Lophocebus Supported               | Cercocebus < Lophocebus Not supported    |
| Absolute crown strength (ACS)                | Cercocebus > Lophocebus | Cercocebus Lophocebus Not supported             | Cercocebus > Lophocebus Supported        |
| Occlusal basin thickness (OBT)               | Cercocebus > Lophocebus | Cercocebus > Lophocebus Supported               | Cercocebus > Lophocebus Supported        |
| Lateral wall enamel thickness (LWET) of functional cusps | Cercocebus > Lophocebus | Cercocebus Lophocebus Not supported             | Cercocebus Lophocebus Not supported      |
| Enamel growth variables: DSR and crown formation time | Cercocebus ≠ Lophocebus | Not evaluated                                    | Cercocebus Lophocebus Not supported      |
| Enamel growth variables: EER                 | Cercocebus ≠ Lophocebus | Not evaluated                                    | Cercocebus > Lophocebus Supported        |
| Cusp angle                                   | Cercocebus > Lophocebus | Cercocebus might be less blunt Unclear           | Cercocebus might be less blunt Unclear   |
| Flare of functional cusps                    | Cercocebus > Lophocebus | Cercocebus > Lophocebus Supported               | Cercocebus Lophocebus Not supported      |

Note: Boldface denotes statistical significance.
abrasive diets (Lucas et al., 2008), and is associated with durophagy in cercopithecid primates (Thiery et al., 2017). Moreover, as previously noted, proportionally thicker occlusal basin enamel, specifically, has been found in orangutans (Kono, 2004) and other durophagous primates (O’Hara, 2021). We note that despite the differences between Cercocebus and Lophocebus in occlusal basin enamel thickness, there was no difference between the two genera in daily enamel secretion rates in this region. Thickened enamel in the occlusal basin of Cercocebus must be produced by some other mechanism, such as by an increase in ameloblast lifespan (e.g., Grine & Martin, 1988) or other complex mechanisms related to the diffusion-limited secretion of enamel matrix atop the underlying dentine core (e.g., Häkkinen et al., 2019).

The greater functional cusp flare of Cercocebus lower molars might better buttress them against laterally-directed bite forces (Macho & Shimizu, 2009; Shimizu, 2002). Functional cusp flare in cercopithecoids appears to have evolved through the incorporation of the cingulum into the molar sidewall (Strasser & Delson, 1987). Molar flare is an ancient feature of Cercopithecoida, present in the 22 Ma nonbilophodont stem Cercopithecid Alopia, and becoming more pronounced in victoriapithecids in conjunction with increased bilophodonty, around 15.5 Ma (Rasmussen et al., 2019). Bilophodonty has been suggested to have originated in a seed-eating context, with the lophs acting as wedges (as per Lucas & Teaford, 1994) to initiate cracks in hard food items (Benefit, 1999). The apparent co-evolution of bilophodonty and well-developed molar flare in victoriapithecids supports the notion that both features originated within the context of durophagy.

Functional lateral cusp flare might also be linked to more pointed molar cusps, characteristic of Papionini relative to Cercopithecini (Kim, 2019). Perhaps greater molar flare is the reason why the cusps of Cercocebus tend to be more pointed than those of Lophocebus. Molar flare has also been suggested to relate to occlusal enamel thickness in that when cusp tips are narrowly spaced on a crown, the occlusal basin is constricted, arguably leading to the accumulation of thicker enamel during development in this region (Olejniczak et al., 2003). Figure 8 is a scatterplot of proportional linear occlusal basin thickness vs. functional cusp flare in mesial slices of Cercocebus and Lophocebus upper molars (all molar types combined). There does not appear to be a positive relationship between the two variables (if there is any relationship, it appears negative in this plot), but larger samples of each tooth type are required to address this question definitively. We note that the one Lophocebus molar with a high proportional linear occlusal basin thickness belongs to L. aterrimus, perhaps indicating that there may be differences in this feature among Lophocebus species.

Despite possessing lower RET, Cercocebus molars have greater ACS than Lophocebus molars, supporting the hypothesis that overall, Cercocebus molars are better protected against fracture. The greater ACS and lower RET of Cercocebus may both be related to its greater flare. With more pronounced flare, Cercocebus molars are wider, and therefore have both greater coronal dentine areas and longer BCDs (Tables 3 and 4; Figure 7). Given that AET is scaled to coronal dentine area in calculating RET, greater molar flare would tend to cause RET to decrease. On the other hand, by increasing the bicervical diameter of a molar, greater flare would tend to cause ACS to increase. Essentially, greater flare, as occurs in Cercocebus relative to Lophocebus molars, may entail a trade-off between ACS and RET.

The data presented here suggest that taller teeth of C. atys are formed primarily by increasing the rate of enamel extension, rather than the duration of enamel formation. Given that taller crowns can result from either faster rates of enamel extension, by lengthening the crown formation period, or both (Guatelli-Steinberg et al., 2012), it may be significant from a life-history perspective that C. atys molars extend at faster rates than those of Lophocebus. Two studies have found correlations between enamel-formation front angles, which are associated with rates of enamel extension, and life history variables, brain size, and/or body size across a range of primates (Guatelli-Steinberg et al., 2018; Hogg & Walker, 2011).

According to the Food Processing Hypothesis (Godfrey et al., 2001), food mechanical properties can directly select for rates of dental development. Specifically, this hypothesis predicts that folivores should have more advanced dental emergence schedules than frugivores because folivorous weanlings must be dentally endowed enough to process mechanically challenging leaves and/or seeds. Harvati (2000) found that colobines, which are folivorous, have relatively advanced M2 and M3 emergence compared to other catarrhines. There is evidence that seed-eating folivores have even more accelerated emergence schedules than those that do not include seeds in their diets (Bolter, 2004). Relative to nonseed eaters, seed-eating folivores (Presbytis rubicunda, Colobus angolensis, and Colobus satanas) emerge their the M2s and M3s earlier in their emergence sequences (Bolter, 2004). It would be interesting to know if other aspects of Cercocebus dental growth and development—besides their rates of
enamel extension—are accelerated relative to Lophocebus. Even newly weaned sooty mangabeys are capable of cracking hard Sacoglottis seed casings (McGraw et al., 2011), perhaps indicating that selection has targeted accelerated dental development in this taxon, although selection for accelerated dental development might also reflect more global influences (e.g., predation pressure) on the pace of Cercocebus growth and development. Additional research aimed at elucidating relationships among dental development, masticatory mechanics, and the ontogeny of ecological independence would provide an avenue for testing the application of the Food Processing Hypotheses to mangabeys.

7 | CONCLUSION

Cercocebus and Lophocebus are both known hard-object feeders, but the former eats hard foods more regularly than the latter. With a greater frequency of hard-food mastication, Cercocebus was hypothesized to have molars with features that would endow them with greater fracture resistance than those of Lophocebus, a fallback consumer of hard foods. This study compared RET, ACS, proportional linear occlusal basin thickness, functional lateral wall thickness, functional cusp tip angles and functional cusp lateral flare between molars of the two genera. Although some comparisons revealed no statistically significant differences between the two genera, Cercocebus molars were shown to have significantly greater ACS (upper molars only), proportional linear occlusal basin thickness (both upper and lower molars), and molar flare (lower molars only), consistent with greater fracture resistance. However, Cercocebus had significantly lower RET (upper molars only) than Lophocebus. We suggest that greater molar flare in C. atys contributes to its greater ACS and lower RET relative to Lophocebus by increasing its bi-cervical diameter and dentine core area. We further argue that because ACS is a better predictor than RET of the fracture resistance of a tooth (Schwartz et al., 2020), C. atys molars appear to be more fracture resistant than those of Lophocebus. Finally, we also examined aspects of enamel growth in these molars, finding that the larger (and taller) crowns of C. atys extended in height significantly more rapidly than those of Lophocebus but did not form in significantly shorter periods of time.

AUTHOR CONTRIBUTIONS

Debbie Guatelli-Steinberg: Conceptualization (lead); data curation (lead); formal analysis (lead); funding acquisition (lead); investigation (equal); methodology (equal); project administration (lead); resources (equal); software (equal); supervision (lead); validation (lead); visualization (equal); writing – original draft (lead); writing – review and editing (equal). Gary T. Schwartz: Conceptualization (supporting); funding acquisition (supporting); investigation (supporting); writing – review and editing (equal). Mackie C. O’Hara: Conceptualization (supporting); data curation (supporting); funding acquisition (supporting); investigation (supporting); methodology (supporting); software (lead); validation (supporting); writing – review and editing (equal). Kaita Guran: Data curation (supporting); investigation (supporting); methodology (supporting); supervision (supporting); validation (supporting). Jess Rychel: Data curation (supporting); investigation (supporting). W. Scott McGraw: Conceptualization (equal); data curation (supporting); formal analysis (supporting); funding acquisition (equal); investigation (equal); project administration (supporting); writing – review and editing (equal).

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CONFLICT OF INTEREST

The authors have no conflicts of interest.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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