THE FUNCTIONAL AND PHYLOGENETIC IMPLICATIONS OF NASOALVEOLAR CLIVUS LENGTH IN ANTHROPOID PRIMATES.

by

Evan E. Bird

A thesis in the Department of Anthropology submitted to the faculty of the Graduate School of Arts and Sciences in partial fulfillment of the requirements for the degree of Master of Arts at New York University.

September 2015

ABSTRACT

The nasoalveolar clivus or subnasal clivus is the sagittally-located portion of the cranium that extends from the site of the anterior attachment of the nasal septum to the margin of the premaxillae at the osteometric point alveolar. In humans, the structure is the vertically-oriented area superior to the upper incisors and inferior to the nasal aperture, underlying the philtrum of the soft tissue. In the majority of primates the structure is formed by the premaxilla, which contain the roots of the upper incisors. Diversity in the relative length of the nasoalveolar clivus is evident among extant primates, with some taxa having a deep clivus and others a relatively short clivus. There is currently no comparative account of nasoalveolar clivus variation among primates. Reviewing the literature reveals that the feature is seldom referred to and little prior research has considered clivus depth as an informative variable in the phylogenetic analysis of the primate cranium. The aim of the present research is to investigate whether clivus depth is significantly different among anthropoid primates and to explore the possible structural and functional factors that influence clivus size. This research shows that the relationship between clivus size and body mass among anthropoid primates does not scale isometrically and that variations in clivus size relative to other traits appear to represent divergent specializations among anthropoids. Analyzing clivus depth measurements from 400 specimens representing 20 anthropoid taxa, the study examines clivus depth in relation to body mass and anterior tooth size, showing that functional constraints from masticatory stresses in the midface appear to influence clivus size. Further, the effects of allometric growth in clivus development and the influence of dietary adaptations are considered with regard to possible specializations in clivus size among Platyrrhini, Cercopithecidae, and Hominoidea. A better understanding of the relationship between these structural-functional variables will potentially have implications for interpreting

the evolutionary relationships and paleobiology of extant and fossil anthropoid primates, including early hominins.

INTRODUCTION

Much of the previous research related to the nasoalveolar clivus has focused more generally on the subnasal region, only considering the clivus among a suite of midfacial traits (Ward and Kimbel, 1983; Shea, 1985; McCollum et al., 1993; McCollum, 2007; Bastir and Rosas, 2004; Kaifu et al., 2011; Williams et al. 2012). Ward and Kimbel (1983) described the characteristics of the subnasal region (including the clivus) among extant hominoids. They noted differences in clivus shape while focusing on the phylogenetic significance of the African and Asian ape subnasal patterns. Ward and Kimbel also compared subnasal patterns of fossil hominins and discussed the potential utility of the morphology noted earlier by Robinson (1954a, b). Other researchers have studied the evolutionary significance of differences in midfacial forms between the African apes and *Pongo* (Ashley Montague, 1935; Gould, 1975; Jolly, 1970; Shea, 1985; Williams et al, 2012; Robinson, 1954a). Shea (1985) argued that, while differences in the nasal floor morphology in great apes may largely be due to differences in facial position, this explanation could be complicated by the possibility that enlarged incisors in orangutans directly influence the shape and orientation of the midfacial region, including the nasoalveolar clivus. Much of the past research on midfacial anatomy has linked variation in the region with that of the dentition, which is often considered in reference to dietary adaptations (Jolly, 1970; Hylander, 1975; Gould, 1975; Wolpoff et al., 1975; Smith, 1981; Gingerich, 1982; Eaglen, 1984; Wolpoff, 1985; Gingerich and Smith, 1985; Benefit and McCrossin, 1991; Anapol and Lee,

1994; Aiello and Wood, 1994; McCollum, 1997; Ungar, 1998; Godfrey et al., 2001; Weston et al., 2004; Strait et al., 2007; Strait et al., 2009; Deane, 2009; Godfrey et al., 2011; Kaifu et al., 2011; Deane 2012; Williams et al., 2012).

A decade after Ward and Kimbel described the shape of the subnasal region and the nasoalveolar clivus in hominoids, McCollum et al. (1993) compared the subnasal region of great apes and humans with those of Australopithecus and Paranthropus, but only presented clivus length as part of the qualitative description of the bony anatomy. This approach was similar to that of other researchers who compared clivus size and shape in fossil and extant taxa (Kaifu et al., 2011; Williams et al., 2012). McCollum and Ward (1997) also discussed clivus size when considering subnasal features, focusing on allometric and ontogenetic changes and sexual dimorphism. McCollum and Ward noted that the nasoalveolar clivus was significantly longer in males of *Pongo*, *Pan*, and *Gorilla* compared to females of the respective taxon. Additionally, they noted that body size, masticatory adaptations, and craniofacial orientation probably have varying influence on differing subnasal morphologies in hominoids. McCollum and Ward's focus on craniofacial form and orientation specifically refers to incisor size, following Shea's presumption that enlarged incisors in *Pongo* effect the shape and orientation of the midface, including the clivus. They comment only on the notably fast growth of the nasoalveolar clivus relative to the basic ranium among nonhuman primates and the positive allometric scaling of clivus length with the basicranium in humans. While previous research has aimed to separate primitive and derived facial forms, variation in the nasoalveolar clivus apart from the general midfacial region has been overlooked. Further evaluation of the nasoalveolar clivus is warranted, as quantification of the feature may be of relevance beyond the scope of this study.

MATERIALS & METHODS

This research used data from 400 individual anthropoid primate crania across 20 genera to consider the influence of body mass and upper incisor size on clivus depth. Measurements of the nasoalveolar clivus were recorded from specimens in the Mammalogy collection of the Vertebrate Zoology Department at the Smithsonian Institution's National Museum of Natural History in Washington, D.C., and the Mammalogy Department of the American Museum of Natural History in New York, N.Y. Additional measurements of Homo sapiens were obtained from crania in the Anthropology collection at the American Museum of Natural History. Extant primate genera in this study included representatives from Platyrrhini, Cercopithecidae, and Hominoidea. The taxa were selected to sample the diversity of nasoalveolar clivus depth among extant anthropoid primates. Each genus was represented by one species and one subspecies where possible [Table 1]. The target sample for each species was 10 adult male and 10 adult female specimens. Adult status was determined by the full eruption of the upper and lower M3 (or M2 in taxa that typically lack M3) of each specimen. Sex determination relied upon the noted assignment of the collector. Specimens without specified sex or likely inaccurate assignments were excluded. Additionally, specimens were included only if they were free of pathological conditions or traumatic damage that may have affected the anatomy of the subnasal and alveolar regions.

Measurement of nasoalveolar clivus length was taken as a chord distance from the osteometric point alveolar (infradentale superius) to the point of greatest distance from alveolare along the sagittal plane at the anterior attachment site of the nasal septum. The latter point is typically identifiable as the osteometric point nasospinale (White, 2000). In some genera, for example those lacking a nasal spine or those displaying more pronounced midfacial prognathism,

the superior demarcation of the clivus is less patent with greater retreat of the maxillae into the nasal cavity (McCollum et al., 1993). It is also necessary to note that definitions regarding the nasoalveolar clivus vary. Williams et al. (2012) reported using the chord length from infradentale superius to the anterior nasal spine, though they do not discuss the reduction of an anterior nasal spine among great apes. McCollum et al. (1993) refer to clivus length as the chord measurement from the anterior nasal attachment site to alveolare, clarifying that the anterior nasal attachment site is not developed as extensively in the great apes and most fossil hominins. In these latter cases, the presence of a small protuberance or tubercle on the clivus represents the most anterior attachment site of the nasal septum. McCollum and Ward (1997) define nasoalveolar clivus length simply as the chord measurement from nasospinale to prosthion (rather than alveolare). Kaifu et al. (2011) also referred to clivus length as the chord from nasospinale to prosthion. Prosthion is defined by White as "[T]he midline point at the most anterior point on the alveolar process of the maxillae." (White, 2000: p. 50). Thus, prosthion is an osteometric point defined by its maximum anterior position, not the more vertically-oriented, inferiormost position in relation to the nasal aperture that better captures the full length of the nasoalveolar clivus. In most cases, these points are synonymous, but in taxa with a verticallyoriented clivus, such as humans, the anteriormost and inferiormost points on the clivus are not necessarily the same. In this study, maximum clivus depth was measured to record greatest length. Therefore, clivus depth was recorded from the sagittal point at the anterior septal attachment site to alveolare, which White (2000) defines as "The midline point at the inferior tip of the bony septum between the upper central incisors." This replicates the method used by Williams et al. (2012) and McCollum et al. (1993).

Due to the possible structural-functional relationships between clivus depth, incisor size and body size noted by other researchers, data on body mass and incisor size for each taxon, presented in Table 2, were obtained from published (Swindler 2002) and unpublished sources (Bailey, personal communication). Incisor size was measured for each species following methods used by Eaglen (1984). Incisor size was recorded as the combined mesiodistal lengths of the upper central and lateral incisors from one side of the dental arcade. The right and left sides were measured and the mean was recorded when the sides differed [Table 3].

All data were analyzed and plotted using Microsoft Excel and Apple Numbers software. Data from males and females of each taxon were plotted separately. Measurements were transformed to base-10 logarithmic (log10) values for allometric analysis. Linear regression equations and R^2 values were calculated. The log10 values predicted by the plots for the dependent variables were then calculated for all data. The predicted log10 value of the dependent variable is found by multiplying the log10 value of the independent variable for each plot by the slope of the regression line then subtracting the y-intercept of the regression equation from the product [predicted $\log 10$ independent variable = (m x $\log 10$ independent variable) - b]. The predicted value of the independent variable was then used to calculate the statistical residual value, which measures the difference between the results obtained via observed measurement of the dependent variable and those predicted based on the overall sample body (Smith, 1981; Wolpoff, 1985; Aiello and Wood, 1994). Residual values are calculated by subtracting the predicted log10 value from the log10 value of the observed measurement, the difference from which is then divided by the same log10 value of the observed measurement and the quotient is then multiplied by 100. This formula [(observed measurement – predicted measurement) / predicted measurement x 100] provides a percentage deviation from the regression line,

indicating the extent to which the variable deviates positively or negatively from the regression line [data listed in appendix].

RESULTS

Mean clivus length was plotted against mean body mass to examine the scaling effects of clivus size. The bivariate plot of the two variables is fit by a line with a coefficient of determination $R^2 = 0.8625$ and a slope of m = 0.4447, displaying a positive allometric relationship [Fig. 2]. General trends in the main taxonomic groups are apparent. The majority of platyrrhines have a relatively large clivus on average for their body size with a mean residual value of +5.036 [range: -16.345 to +29.897]. Notably, *Callithrix*, the smallest primate included in this study, has the largest residual value for clivus length relative to body mass. Conversely, cercopithecoids generally have a short clivus relative to their body size compared to other anthropoids with a mean residual value of -10.850 [range: -36.924 to +12.399]. Only Macaca and Papio females had positive residual values. The trend for a shortened clivus in cercopithecoids is particularly evident in the colobines, with Nasalis and Presbytis having the most negative residual values. Even when colobines are excluded, the mean cercopithecine residual value remains negative at -3.534. Hominoids have a mean residual value of +1.987[range: -16.309 to +18.022]. It is noteworthy that hylobatids (which have a mean residual value of -10.116 and are more similar to the cercopithecids than to the great apes and humans) are the only hominoids with appreciably negative residual values. Symphalangus has the most negative residual, comparable to Ateles and Nasalis females. The mean residual value for the Hominidae is +8.039. Omitting *Homo*, which falls close to the regression line, increases the great ape

residual value to +10.975. This highlights the specialized condition among the Hominidae, particularly *Pongo* and *Pan*.

Comparisons of mean body size to mean incisor size among the 14 taxa for which incisor data were available exhibit a distribution pattern similar to the comparisons of body mass and clivus length, suggesting a relationship between incisor size and clivus depth [Fig. 3]. Many genera fall close to the regression line ($R^2 = 0.796$, m = 0.283). There is a negative allometric relationship for mean incisor size and mean body mass, which is the converse of the positive allometric relationship of mean clivus size and mean body mass. Most of the great apes cluster above the regression line, while the hylobatids are located below the line and are clustered with the colobines, which have small incisors relative to body mass. The cercopithecines fall above the regression line with larger upper incisors than expected for their body sizes. *Homo* lies below the line with small teeth relative to body size. Although only three platyrrhine genera have published incisor width data, they all fall on the positive sides of the regression line as they did for mean clivus length plotted against mean body mass. The presence of *Homo* below the regression line brings the hominoid mean residual value to -0.189 with a standard deviation of 7.224 but the value rises to +2.652 when the hylobatids are excluded. This rises further to +6.159 with the exclusion of *Homo*. The mean residual value for cercopithecoids is in the positive range at +0.937, driven primarily by the cercopithecines with large incisors relative to their body masses, though their standard deviation is also the highest at 9.683. The platyrrhines have an average residual value of -3.947, but their standard deviation is also high at 9.307. Similar to the comparison of mean clivus size to mean body mass, the large-bodied hominoids appear to have a strong influence on the upper end of the regression line due to having both large incisors and large clivuses relative to body mass.

When mean clivus size is compared to mean upper incisor size all hominoids except *Gorilla* plot above the line of regression ($R^2 = 0.896$, m = 1.687) [Fig. 4]. The hominoids have a mean residual value of +3.756 [range: -3.148 to +11.277] and the smallest standard deviation (4.191). This value rises slightly to +3.843 with the exclusion of the hylobatids. *Homo* has the greatest residual values among all the hominoids reaching the positive tens range. All other hominoids are in the range of positive single digits for residual values (*Pan* females being the only other taxon above +5). The hylobatids are slightly above the line of regression and they cluster near the cercopithecoids due to their similar clivus sizes. The cercopithecoids again have mostly negative residual values and are below the line of regression with only the male macaques plotting slightly above (with a residual value of +2.289). Among the cercopithecoids, Lophocebus show the greatest negative values, reaching in to the -30s for both sexes. Even with the exclusion of *Lophocebus*, however, the cercopithecoids retain a negative mean residual value (-5.874, range: -13.210 to +2.289, SD = 12.783). This drops to -11.479 when Lophocebus is included. While the platyrrhines are variable, with a mean residual value of +6.355 (ranging from -5.544 to +20.743 with a standard deviation of 10.179), Aotus plots below the regression line with the other platyrrhines above it. *Alouatta* has the largest residual values with males reaching +20.734. Cercopithecoids have the greatest standard deviation but with the exclusion of Lophocebus this drops, leaving platyrrhines with the greatest standard deviation (which remains even when Alouatta is excluded). The R^2 value is stronger than for the other comparisons, implying that a significant relationship exists between clivus depth and incisor size.

Plotting the residual values of clivus length with the residual values of incisor size (both calculated relative to mean body mass) removes the effects of allometry [Fig. 5]. The plot displays a relationship between clivus size and incisor size, showing that the majority of taxa fall

into two quadrants: those with both a large clivus and large incisors relative to body mass, and those with both a small clivus and small incisors relative to body mass. The large-bodied hominoids fall into quadrant 1 (large-large), where *Aotus* females and the macaques are also found. That *Macaca* males cluster closer to *Pan* is perhaps unsurprising as *Macaca* males have larger clivuses relative to body mass than other cercopithecoids, as well as larger clivuses relative to mean incisor size and larger incisors relative to mean body mass. *Nasalis* and Presbytis are both in quadrant 3 (small-small). Notably, Homo and the hylobatids are also in quadrant 3 but Homo plots near the neutral line of the x-axis, showing that their clivus is close to nearly proportional relative to body mass compared to the other taxa, but with small incisors. Quadrants 2 and 4 show those taxa that have clivus sizes that are counter to the association with incisor size seen in most anthropoid primates.. Cercopithecus and Lophocebus of both sexes are in quadrant 4 with incisors that are large relative to body mass but a small clivus and *Alouatta* males is the only sample that falls in quadrant 2. This again displays the character of cercopithecines to have a shortened clivus while the colobines are similar to several of the platyrrhines. Additionally, the hylobatids again show dissimilarity to the other hominoids, being in quadrant 3, but further away from the x-axis than *Homo*. The expanded clivus of the largebodied hominoids is also again displayed. This plot highlights the clustering of the taxa into specialized morphological groups, and that clivus size is strongly linked with upper incisor size.

DISCUSSION

The study documents the variation present in nasoalveolar clivus length in anthropoid primates. Relative to platyrrhines, cercopithecoids have reduced the length of the clivus with notable differences between the cercopithecines and the colobines, the latter having distinctly shortened clivuses relative to their body mass and incisor width. The presence of small incisors relative to body mass in platyrrhines has been discussed as a plausible primitive condition (Eaglen, 1984; Swindler, 2002). The hominoids exhibit clivus expansion, particularly among the hominids, although humans appear to have secondarily reduced the length of the clivus compared to the great apes in association with shortening of the lower face. The hylobatids are more similar to the cercopithecines, and the two groups arguably retain the ancestral catarrhine condition. The observed specializations in clivus length among anthropoid primates appear to be related, in part, to incisor size, though other structural-functional factors may contribute to variation in clivus length

The plot of mean clivus length to mean body mass in anthropoids exhibits a positive allometric relationship (m = 0.445). However, this relationship is largely determined by the very deep clivuses in the large-bodied great apes. When *Gorilla, Pan,* and *Pongo* are removed from the plot the slope is very close to isometry (m = 0.346). The plot of mean clivus size to mean incisor size also shows a positive allometric relationship (m = 1.687). When hominoids are excluded the slope value is m = 0.905. This value only rises to 0.930 when the hylobatids are included. The plot of mean incisor size to mean body mass is negatively allometric (m = 0.283). Additional analysis of the slope of mean clivus length to mean body mass for cercopithecoids and platyrrhines (excluding all hominoids) shows that many of the remaining taxa do not fall close to the regression line. These results confirm that clivus size variation in anthropoids is not simply proportionally scaled with body mass, but rather is influenced by adaptive specializations in certain clades.

While non-human hominoids have a specialized clivus that is expanded compared with the inferred primitive catarrhine condition, *Homo* is notably different. Comparing mean clivus

depth to mean body mass, *Homo* is close to the regression line with residual values close to zero (+0.055 in males and -1.596 in females). The incisors of Homo have negative residual values relative to body mass, while mean clivus length in *Homo* has positive residual values relative to body mass. Thus, even though their teeth are relatively small, the clivus depth is neither notably large nor small relative to body size. Among hominids this is unusual, as *Pan* and *Pongo* both have large clivuses and large incisors relative to body mass. Residual values for clivus length relative to incisor size in Pan and Pongo are lower than Homo, though. Gorilla stands out among the great apes with a negative residual value for clivus size relative to incisor size. Gorilla males also have a residual value of only +1.235 for clivus size relative to body mass and a residual value of +0.584 for incisor size relative to body mass (Gorilla females have greater residual values than males for both these residual values, while female Gorilla have incisors and a clivus approximately the same size as males but a body mass that is less than half that of males). Like Homo, the hylobatids have a clivus that is large relative to incisors size, and incisors that are small relative to body mass. However, hylobatids have a clivus that is small relative to their body mass. This supports the inference that hominids evolved an enlarged clivus along with enlarged incisors relative to body size. The reduced incisor size in humans relative to body mass is documented by their positive residual values for clivus size relative to incisor size. Among hylobatids the clivus arguably may retain the ancestral condition, having incisors that are relatively small for their body mass. It is worth considering if the variations observed in the residual values among the great apes may be influenced by sexual dimorphism. While sexual dimorphism of subnasal form has only been noted in *Pongo* and *Gorilla*, overall dimorphism of tooth size relative to the face and body size are reported (McCollum and Ward, 1997; Weston et al., 2004). Males and females in Pan have relatively similar mean clivus lengths, while males are over 10 kg larger than females with a 2 mm difference in mean incisor size. *Pongo* males have a clivus which is on average 5 mm longer than females and mean incisor width 1 mm larger though they have nearly twice the body mass of females.

While the hominoids in general show expansion of the clivus, and many of the cercopithecoids show reduction, the platyrrhines are more diverse. All the platyrrhines included in this study have a clivus that is large relative to their body mass. If incisor size influences clivus size then it might be expected that the platyrrhines would have large incisors for their body mass as well. Incisor size data was only available for three of the platyrrhine taxa in this study. Among these, *Alouatta* and *Ateles* both have small incisors relative to their body mass as well as a generally small clivus size for their body mass (*Alouatta* males being an exception with a slightly larger clivus for their body mass). *Aotus* is the other platyrrhine with incisor width data available and they show incisors that are on average large relative to their body mass but a clivus that is small relative to their body mass as well as relative to incisor width. It has been inferred that platyrrhines may retain the ancestral anthropoid primate condition of generally small incisors relative to body size (Eaglen, 1984).

The relationship between clivus size and incisor size is more strongly correlation than the relationship between clivus size and body mass. This is particularly the case for cercopithecoids. For example, *Presbytis* is near the middle of the range for body mass among cercopithecoids but has the most negative residual values for clivus size relative to body mass. Their incisors are also small for their body mass, as is their mean incisor size relative to their mean clivus size, but each less markedly so than their mean clivus size relative to mean body mass. *Macaca*, the smallest cercopithecoid in this study, has the only positive residual values for clivus depth relative to incisor size among the males, whereas the similar-sized *Cercopithecus* has negative

residual values. *Macaca* also has upper incisors that are large relative to their body mass and a clivus that is also large relative to their body mass, while Cercopithecus have lower residual values for incisors relative to body mass. The platyrrhines show some diversity in clivus size in relation to body mass. Saguinus is only slightly larger than Callithrix, yet Saguinus has residual values slightly above +0 and +1 (respective of sex) while *Callithrix* has residual values at approximately +25 to +30 for clivus size relative to body mass. Additionally, three-quarters of the platyrrhines with negative residual values are among the larger genera of New World monkeys. However, this is not a general predictor of relatively smaller clivus length relative to body mass among the platyrrhines. Male Alouatta and both sexes of Cebus and Pithecia, three of the largest platyrrhines, all remain in the positive residual value range. Meanwhile, Ateles, the largest platyrrhine in the study, has the most negative residual value among the New World monkeys. Of the smaller-bodied (less than 1 kg) platyrrhines all except *Aotus* have positive residual values. Thus, many of the platyrrhines have larger clivuses relative to their body mass, but the largest taxa tend to have a relatively small clivus. Apparent here is the association between small incisors and a small clivus in relation to body size, respectively. Concurrently, incisors and clivus are large in relation to each other, which some researchers have explained in reference to dietary differences (Anapol and Lee, 1994).

The nasoalveolar clivus accommodates the roots of the upper incisors. The incisor roots extend superiorly into the subnasal clivus region. In some taxa, such as *Saguinus* and *Ateles*, the nasal aperture descends inferiorly into the region medial to the upper central incisors. Strait et al. (2007) noted that the strains experienced during mastication are elevated in regions of the face other than just the palate and immediate dentition, implying a suite of facial features would likely develop together for adaptive resistance to masticatory stresses. Given the important role of

incisors in food processing, it can be assumed that the anterior dentition will be influenced by selective pressures that arise from the functional mechanics of specific diets (Deane, 2009). Incisor root size, and the size of the incisors themselves, can be inferred to have a direct relationship on the development of the nasoalveolar clivus (Wolpoff, 1985). The effect which incisal preparation has on incisor size has received attention from numerous researchers for at least half a century (Robinson, 1954a; Jolly, 1970; Hylander, 1975; Eaglen, 1984; Gingerich, 1985; Anapol and Lee, 1994; Ungar, 1998; McCollum, 2007; Strait et al., 2009; Deane, 2009; Deane, 2012; Williams et al., 2012). Robinson (1954) and Jolly (1970) argued that small incisors relative to molars in fossil taxa could be indicative of a seed-eating diet. Deane (2009b) also contended that narrow incisors in Miocene catarrhines are indicative of greater folivory. Hylander (1975) noted a correlation between incisor size and incisal preparation, as increased preparation of food items would lead to greater tooth wear. Analyzing incisor size in relation to body mass and finding a near-isometric relationship, Hylander theorized that larger incisors are an adaptation to increased wear. Primates that eat food items, such as fruits, that require preparation prior to ingestion have larger incisors than those primates that eat leaves. Hylander also found a distinction between leaf-eating colobines and frugivorous cercopithecines, with smaller teeth relative to their body size in the former. Further, Hylander noted the division of hylobatids from other hominoids, and hypothesizes that the reduced dentition of Homo is a consequence of food preparation techniques. Eaglen (1984) found a negative allometric relationship between incisor size and body mass among the Platyrrhini. Noting that, regardless of diet, platyrrhines have smaller incisors relative to body mass compared to catarrhines. Eaglen argued that the smaller incisors of New World primates represent the primitive anthropoid condition. Further, Eaglen contends that increases in incisor size among other anthropoids are

due to dietary shifts that require increased incisal preparation of food items. This supports the results of the present study regarding colobines and their similarity to platyrrhines in measures of incisor size relative to body mass. Goldstein et al. (1978) used a similar argument for cercopithecoids, noting that frugivorous and omnivorous taxa have notably wider incisors than folivorous colobines. Regarding the negative allometric relationship of incisor size relative to body mass, Eaglen noted that the largest platyrrhines are the most folivorous and those with the largest incisors relative to body mass are frugivorous. In general, these studies and the present study show that relative incisor size tends to decrease proportionally as body size increases among platyrrhines. Additionally, for species where the comparison is possible, platyrrhines are shown to have relatively smaller incisors than catarrhines when diets overlap (Eaglen, 1984; Deane, 2009). Anapol and Lee (1994) also analyzed platyrrhine diet and morphological adaptations, noting that broader incisors, such as those of *Ateles*, are a result of feeding primarily on fruit mesocarps. Relatively wider incisors are found in platyrrhines that consume fruits in addition to animal protein (Cebus, Saimiri, and Saguinus), while narrower incisors are found in those where fruits are supplemented exclusively by non-animal foods (Alouatta and Pithecia). It has also been argued that small incisors may be the primitive condition for platyrrhines, while the opposite may be true for catarrhines (Eaglen, 1984). In this hypothesis, incisor reduction may be the evolutionary novelty with larger incisors being the primitive catarrhine condition. The reduced incisors of colobines is probably due to the demands of leaf-eating, associated with less extensive use of the incisors (Eaglen, 1984; Anapol and Lee, 1994; Ungar, 1998). The majority of small-bodied platyrrhines are frugivorous and insectivorous, while the larger platyrrhines combine folivory and frugivory. Among the hominoids, *Pan* and *Pongo* are both frugivorous while Gorilla and Hylobates have a mixed diet of fruits and leaves. Pan and Pongo

differ though, since orangutans feed on hard objects while chimpanzees consume soft objects. Some research has found greater incisor size among hard object feeding hominoids than soft object feeders (Deane, 2009a, b, 2012). The seasonal folivore-frugivores are intermediate in incisor size to dedicated leaf or fruit eaters. Pan, Pongo, and Gorilla are the only hominoids with large incisors relative to their body size. In Homo the incisors are reduced relative to body size, while the hylobatids are more similar to cercopithecoids. Additionally, *Gorilla* is the most folivorous hominoid with incisors that are closest to the size expected for their body mass among the hominoids. Platyrrhines are not dissimilar to hominoids in this regard, which may be expected when considering their tendency for have a relatively deep clivus. These observations are correspond to those made when comparing mean clivus length to mean body mass [Table 2]. Previous research on platyrrhine incisor size and diet also corresponds with the data in the present study (Deane, 2012). Aotus, an omnivore, is the only platyrrhine with a positive residual value for mean incisor size compared to body mass, while Ateles (a frugivore) is located well below the regression line, along with *Alouatta* (a folivore). Likewise, *Aotus* is the only platyrrhine with a large clivus relative to body size. Among cercopithecoids, the diets of cercopithecines include leaves and fruits supplemented with animal protein, while the colobines rely primarily on leaves. In the present research, the colobines are the only cercopithecoids with small incisors relative to their body size and have extremely negative residual values among the cercopithecoids for mean clivus length relative to body mass. Other cercopithecoids, other than *Macaca*, also have small clivuses relative to body mass, but with less marked residual values. It should be noted, though, that some researchers have expressed concerns about how informative the relationship is between diet and dental size and morphology. Eaglen (1984) warned that distinctions between platyrrhines and catarrhines may be clouded by the overall body size

differences between the two groups. Ungar (1998) also questioned whether diet and incisor size correlations are reliable across taxa, pointing to examples in which the frugivory/folivory distinction does not predict incisor size differences. For example, platyrrhines have relatively smaller incisors than cercopithecoids, but they are not necessarily more folivorous. This could complicate analyses of fossil taxa or make researchers wary of comparisons between taxa that are of unknown phyletic affinities. Godfrey et al. (2001) noted that consumption between different seasons or even forests may vary. They point out that among colobines, dedicated leaf consumption can vary from as little as 26% of total diet to as much as 94%, depending on season and geography. Additionally, seeds can make up anywhere from 0-50% of the same diets and frugivores at times may ingest large amounts of foliage (Godfrey et al., 2001). Using Hylander's (1975) study as a basis, McCollum (2007) sought to analyze African apes but found that incisor wear rates did not support the resistance hypothesis for development of broader incisor crowns. Ungar (1998) also found that incisors are used regularly by folivores during food preparation. However, Deane (2009a) points out that McCollum's analysis fails to distinguish the overlap in diets between the few taxa analyzed. More recent analyses have supported Hylander's original theory, that frugivores can be discriminated using incisor size as a guide (Deane, 2009a, 2012).

Although research on the nasoalvolar clivus in extant anthropoid primates has been sparse, some authors have considered the feature in the fossil record (McCollum et al., 1993; Deane, 2009b; Williams et al., 2012). Data on clivus length in fossil hominoids, while limited [Table 5], shows that the subnasal morphologies of extinct species are intermediate between those of extant great apes and humans (McCollum et al., 1993; Williams et a., 2012). The general subnasal morphology of *Homo erectus* is intermediate between African apes and modern humans. Measurements for *H. erectus* are close to *Gorilla* and *Pan* but smaller than *Paranthropus*. In comparison, the clivus length of *A. afarensis* shows that the species had a clivus slightly larger than humans but smaller than in African apes. Also, *Australopithecus africanus* has a similar clivus length to *A. afarensis*. Additionally, the measurements for *Homo habilis* show a clivus that is of comparable in length or slightly larger than humans. This supports the notion that the clivus size reduction trend observed in modern humans may have begun in early hominins.

CONCLUSION

The differences observed in mean nasoalveolar clivus length among anthropoid primates can be interpreted in a phylogenetic context. Previous research has lacked substantial consideration or quantification of the variability seen in the nasoalveolar clivus of anthropoids. The results present study shows that functional constraints of the anthropoid face appear to dictate the relationship between clivus size and other anatomical factors. Specifically, a relationship likely exists between upper incisor size and clivus depth. The relationship between incisor size and clivus development may be a consequence of dietary adaptations to resist masticatory stresses in the midface.

The relationship between clivus size and body mass among anthropoid primates does not scale isometrically. There appears to be divergent specializations in relative clivus size and incisor size among anthropoid primates: those with large clivuses and large incisors relative to their body mass and those with small clivuses and small incisors. Great apes have a marked expansion of the nasoalveolar clivus relative to their body size, while a secondary reduction has occurred in *Homo*. Relative clivus size among fossil hominins shows that this secondary reduction may have begun among early hominins. Platyrrhines exhibit diversity in clivus length,

but the taxa included in this study tend towards small incisors and a small clivus relative to body mass. The cercopithecines, being similar to platyrrhines, retain what can be inferred to be the ancestral anthropoid condition. In contrast, the colobines show a specialized condition with their clivus being small relative to their body size. The hylobatids share a similar condition to most cercopithecoids, and may be inferred to retain the primitive catarrhine condition. As described above, some of the specializations seen in extant anthropoids may be attributable to upper incisor size and its relationship to dietary adaptation. The variability in the size of the nasoalveolar clivus in anthropoids provides insight into the evolutionary pressures that may have shaped aspects of the lower face. Further exploration of the relationship between clivus depth and other cranio-dental variables may prove useful for interpreting primate phylogenetic relationships and adaptations.

Table 1. Taxa Included in this Study				
Hominoidea	Cercopithecidae	Platyrrhini		
Gorilla gorilla	Cercopithecus mitis	Alouatta seniculus		
Homo sapiens	Lophocebus johnstoni	Aotus trivirgatus		
Hoolock hoolock	Macaca fascicularis	Ateles geoffroyi		
Pan troglodytes	Nasalis larvatus	Callithrix argentata		
Pongo pygmaeus	Papio anubis	Pithecia monachus		
Symphalangus syndactylus	Presbytis melalophos	Saguinus nigricollis		
		Saimiri boliviensis		
		Sapaju capucinus		

Table 2. Mean Clivus Length & Mean Body Mass (Smith and Jungers, 1997)							
	Taxon	Clivus (mm)	SD	Body Mass (g)	Residual		
	Gorilla gorilla ී	36.06	2.97	170400	1.014		
	Gorilla gorilla 🍳	30.79	2.50	71500	7.719		
	Homo sapiens ੈ	21.33	1.39	64240	-1.794		
H o m i n o i d e a	Homo sapiens ♀	21.14	1.23	56170	-0.136		
	Hoolock hoolock 💍	7.36	1.14	6870	-6.254		
	Hoolock hoolock $igodoldsymbol{arphi}$	7.66	0.67	6880	-4.201		
	Pan troglodytes ੈ	31.08	2.60	59700	12.575		
	Pan troglodytes ♀	34.27	2.45	45800	16.119		
	Pongo pygmaeus ്	37.76	1.83	78500	11.761		
	Pongo pygmaeus ♀	32.25	2.35	35800	17.879		
	Symphalangus syndactylus ೆ	7.96	0.88	11900	-14.017		
	Symphalangus syndactylus ្	7.32	0.86	10700	-16.433		
с	Cercopithecus mitis ै	6.49	0.94	5850	-14.277		
	Cercopithecus mitis \bigcirc	6.24	0.60	3930	-8.499		
	Lophocebus johnstoni 💍	6.87	1.25	8250	-9.579		
e r	Lophocebus johnstoni ${\mathbb Q}$	6.69	0.82	6020	-2.269		
c o	Macaca fascicularis 👌	9.91	0.88	5360	12.339		
p i	Macaca fascicularis ♀	7.29	1.61	3590	7.763		
t h	Nasalis larvatus <i>ै</i>	7.27	1.03	20400	-31.311		
c i	Nasalis larvatus ♀	7.07	0.72	9820	-16.561		
da	Papio anubis ୖ	13.88	2.65	25100	-2.534		
e	Papio anubis ${\mathbb Q}$	11.46	1.62	13300	0.992		
	Presbytis melalophos ੈ	5.05	0.90	6590	-30.467		
	Presbytis melalophos ${\mathbb Q}$	4.61	0.60	6470	-37.036		
	Alouatta seniculus ୖ	8.87	1.31	6690	3.371		
	Alouatta seniculus ${igsiremath{\mathbb Q}}$	6.14	1.24	5210	-10.088		
	Aotus trivirgatus ്	3.21	0.44	810	-0.335		
	Aotus trivirgatus ♀	3.41	0.46	740	7.885		
	Ateles geoffroyi ♂	6.48	1.21	7780	-16.455		
Р	Ateles geoffroyi ♀	6.47	0.90	7290	-15.002		
l a	Callithrix argentata ै	3.02	0.45	330	30.255		
t y	Callithrix argentata 🏻	2.97	0.24	360	25.360		
r r	Pithecia monachus 👌	5.56	0.48	2610	5.549		
i n	Pithecia monachus ${\mathbb Q}$	4.96	1.08	2110	10.109		
ï	Saguinus nigricollis 👌	2.94	0.36	470	1.462		
	Saguinus nigricollis 🏻	2.67	0.12	480	0.941		
	Saimiri boliviensis	3.99	0.50	910	13.935		
	Saimiri boliviensis 🍳	3.41	0.33	710	4.539		
	Sapaju capucinus 👌	7.04	0.56	3680	11.696		
	Sapaju capucinus 🎗	6.47	0.81	2540	9.385		

Table 3. Mean Clivus Length & Mean Incisor Width (Swindler, 2002; Bailey, 2013)						
	Taxon	Clivus Length (mm)	Incisor Width (mm)	Residual		
	Gorilla gorilla ି	36.06	25.5	-1.652		
	Gorilla gorilla ♀	30.79	23.9	-3.148		
H o m i	Homo sapiens ੇ	21.33	14.7	11.277		
	Homo sapiens ♀	21.14	15.0	9.900		
	Hoolock hoolock 3	7.36	9.2	3.597		
0	Hoolock hoolock \cap	7.66	9.3	4.592		
i d	Pan troglodytes 👌	31.08	22.7	2.201		
e a	Pan troglodytes ♀	34.27	20.7	6.834		
	Pongo pygmaeus 👌	37.76	23.5	3.432		
	Pongo pygmaeus ♀	32.25	22.2	1.899		
	Symphalangus syndactylus 👌	7.96	9.5	4.629		
	Symphalangus syndactylus	7.32 9.4		1.509		
	Cercopithecus mitis 🖑	6.49	9.6	-6.727		
C e r c o p it h e	Cercopithecus mitis $\stackrel{\frown}{\downarrow}$	6.24	9.5	-8.052		
	Lophocebus johnstoni 👌	6.87	14.1	-37.233		
	Lophocebus johnstoni ${\mathbb Q}$	6.69	12.2	-30.562		
	Macaca fascicularis 👌	9.91	11.1	2.289		
	Macaca fascicularis ♀	7.29	10.9	-11.269		
C	Nasalis larvatus 👌	7.27	10.0	-4.093		
d	Nasalis larvatus 🏳	7.07	9.5	-1.153		
e e	Presbytis melalophos ∂	5.05	8.0	-4.779		
	Presbytis melalophos ♀	4.61	8.2	-13.210		
Р	Alouatta seniculus 👌	8.87	8.2	20.734		
a I	Alouatta seniculus 🌻	6.14	7.5	12.964		
t y	Aotus trivirgatus 👌	3.21	6.1	-5.544		
rr h	Aotus trivirgatus ♀	3.41	6.3	-4.781		
i	Ateles geoffroyi 👌	6.48	8.2	7.417		
i	Ateles geoffroyi ♀	6.47	8.2	7.341		

Tab	Table 4. Mean Incisor Width & Mean Body Mass (Smith and Jungers, 1997; Swindler, 2002; Bailey, 2013)					
	Taxon	Mean Incisor Width (mm)	Mean Body Mass (g)	Residual		
	Gorilla gorilla ି	25.5	170400	0.584		
H o m	Gorilla gorilla ♀	23.9	71500	6.756		
	Homo sapiens 👌	14.7	64240	-8.977		
	Homo sapiens ♀	15.0	56170	-6.761		
	Hoolock hoolock 👌	9.2	6870	-3.483		
i n	Hoolock hoolock $\stackrel{\frown}{\downarrow}$	9.3	6880	-3.000		
o i	Pan troglodytes 👌	22.7	59700	6.853		
d	Pan troglodytes ♀	20.7	45800	6.493		
a	Pongo pygmaeus 👌	23.5	78500	5.421		
	Pongo pygmaeus ♀	22.2	35800	10.852		
	Symphalangus syndactylus 👌	9.5	11900	-8.914		
	Symphalangus syndactylus ${}^{\mathbb{Q}}_{+}$	9.4	10700	-8.086		
_	Cercopithecus mitis 👌	9.6	5850	0.475		
C e	Cercopithecus mitis \bigcirc	9.5	3930	5.013		
r C	Lophocebus Johnstoni 👌	14.1	8250	11.257		
o p	Lophocebus Johnstoni ${\mathbb Q}$	12.2	6020	11.388		
i t h	Macaca fascicularis 👌	11.1	5360	7.507		
	Macaca fascicularis ${\mathbb Q}$	10.9	3590	11.551		
e C	Nasalis lavartus 👌	10.0	20400	-13.112		
i d	Nasalis lavartus ♀	9.5	9820	-6.499		
a e	Presbytis melalophos 👌	8.0	6590	-9.872		
	Presbytis melalophos ${\mathbb Q}$	8.2	6470	-8.336		
P	Alouatta seniculus 👌	8.2	6690	-8.785		
a	Alouatta seniculus ${\mathbb Q}$	7.5	5210	-10.091		
t y	Aotus trivirgatus ♂	6.1	810	6.459		
r r	Aotus trivirgatus ♀	6.3	740	9.489		
h i	Ateles geoffroyi ♂	8.2	7780	-10.815		
n i	Ateles geoffroyi ♀	8.2	7290	-9.941		

Table 5. Mean Clivus Height Among Extant and Fossil Taxa Published Literature						
		Taxon	Mean Clivus Height (mm)	SD	EB Data	SD
	E	Gorilla gorilla	33.2	6.15	33.4	2.74
	t a	Homo sapiens	19.2	3.61	21.2	1.31
	n t	Pan troglodytes	31.6	4.93	32.7	2.53
		Australopithecus africanus	27.0	2.86	-	-
Williams et al. (2012)	_	Homo erectus	32.7	4.37	-	-
	Г 0	Homo habilis	23.7	8.36	-	-
	s s	Paranthropus aethiopicus	36.2	-	-	-
	i I	Paranthropus boisei	36.6	5.15	-	-
		Paranthropus robustus	29.0	0.94	-	-
		StW 53	26.4	-	-	-
	Е	Gorilla gorilla	31.1	5.8	33.4	2.74
	x t	Homo sapiens	19.3	2.5	21.2	1.31
	a n	Pan troglodytes	32.1	4.5	32.7	2.53
	t	Pongo pygmaeus	29.9	4.4	35.0	2.09
		Australopithecus africanus	25.2	4.2	-	-
		Australopithecus afarensis	24.3	1.5	-	-
		Paranthropus boisei	37.1	11.5	-	-
McCollum et		Paranthropus robustus	29.0	3.0	-	-
al. (1993)		StW 53 (Homo gautengensis)	25.5	-	-	-
	F O	KNM-WT 17000 (Paranthropus aethiopicus)	27.0			
	s s	KNM-ER 1470 (Homo rudolfensis)	33.5	-	-	-
	i I	KNM-ER 1805 (Homo habilis)	21.0	-	-	-
		KNM-ER 1813 (Homo habilis)	25.3	-	-	-
		OH 24 (<i>Homo habilis</i>)	25.7	-	-	-
		OH 62 (<i>Homo habilis</i>)	23.0	-	-	-
		KNM-ER 3733 (Homo ergaster)	31.00	-	-	-
		SK 847 (Early <i>Homo</i>)	32.00	-	-	-



Fig. 1: Measurement of maximum nasoalveolar clivus length in anthropoids. [Photo by author]



Figure 2. Log₁₀ Mean Clivus Height and Log₁₀ Mean Body Mass.



Figure 3. Log_{10} Mean Incisor Width and Log_{10} Mean Body Mass.



Figure 4. Log $_{10}$ Mean Clivus Length and Log $_{10}$ Mean Incisor Width



Figure 5. Clivus Residuals from Body Mass and Incisor Residuals from Body Mass.

References:

Aiello LC, Wood BA. 1994. Cranial Variables as Predictors of Hominine Body Mass. American Journal of Physical Anthropology 95: 409-426.

Anapol F, Lee S. 1994. Morphological Adaptation to Diet in Platyrrhine Primates. American Journal of Physical Anthropology 94: 239-261.

Ashley-Montagu MF. 1935. The Premaxilla in the Primates. The Quarterly Review of Biology 10 (1):. 32-59.

Bastir M, Rosas A. 2004. Facial Heights: Evolutionary Relevance of Postnasal Ontogeny for Facial Orientation and Skull Morphology in Humans and Chimpanzees. Journal of Human Evolution 47: 359-381.

Benefit BR, McCrossin ML. 1991. Ancestral Facial Morphology of Old World Higher Primates. Proceedings of the National Academy of Sciences 88 (12): 5267-71.

Deane A. 2009. First Contact: Understanding the Relationship Between Hominoid Incisor Curvature and Diet. Journal of Human Evolution 56: 263-274.

Deane AS. 2009. Early Miocene Catarrhine Dietary Behaviour: The Influence of the Red Queen Effect on Incisor Shape and Curvature. Journal of Human Evolution 56: 275-285.

Deane A. 2012. Platyrrhine Incisors and Diet. American Journal of Physical Anthropology 148: 249-261.

Eaglen RH. 1984. Incisor Size and Diet Revisited: The View From a Platyrrhine Perspective. American Journal of Physical Anthropology 64 (3): 63-275.

Fleagle JG. 1985. Size and Adaptation in Primates. In: Jungers WL, (Ed.), Size and Scaling in Primate Biology. Plenum Press, New York, NY. pp. 1-19.

Gingerich PD, Smith BH, Rosenberg K. 1982. Allometric Scaling in the Dentition of Primates and Prediction of Body Weight From Tooth Size in Fossils. American Journal of Physical Anthropology 58: 81-100.

Gingerich PD, Smith BH. 1985. Allometric Scaling in the Dentition of Primates and Insectivores. In: Jungers WL. (Ed.), Size and Scaling in Primate Biology. New York: Plenum Press. p. 257-272.

Godfrey LR, Samonds KE, Jungers WL, Sutherland MR. 2001. Teeth, Brains, and Primate Life Histories. American Journal of Physical Anthropology 114: 192-214.

Goldstein S, Post D, Melnick D. 1978. An Analysis of Cercopithecoid Odontometrics. American Journal of Physical Anthropology 49: 517-532.

Gould SJ. 1975. On the Scaling of Tooth Size in Mammals. American Zoology 15:. 351-362.

Hylander WL. 1975. Incisor Size and Diet in Anthropoids with Special Reference to Cercopithecidae. Science, New Series, 189 (4208): 1095-1098.

Jolly CJ. 1970. The Seed Eaters: A New Model of Hominid Differentiation Based on an Analogy. Man 5: 5–26.

Kaifu Y, Baba H, Sutikna T, Morwood MJ, Kubo D, Saptomo EW, Jatmiko, Awe RD, Djubiantono T. 2011. Craniofacial Morphology of *Homo floresiensis*: Description, taxonomic affinities, and Evolutionary Implications. Journal of Human Evolution 61: 644-682.

Kean MR, Houghton P. 1987. The Role of Function in the Development of Human Craniofacial Form-A Perspective. The Anatomical Record 218: 107-110.

McCollum MA, Grine FE, Ward SC, Kimbel WH. 1993. Subnasal Morphological Variation in Extant Hominoids and Fossil Hominids. Journal of Human Evolution 24: 87-111.

McCollum MA, Ward SC. 1997. Subnasoalveolar Anatomy and Hominoid Phylogeny: Evidence from Comparative Ontogeny. American Journal of Physical Anthropology 102: 377-405.

McCollum MA. 2007. Rethinking Incisor Size and Diet in Anthropoids: Diet, Incisor Wear and Incisor Breadth in the African Apes. American Journal of Physical Anthropology 133: 986-993.

Robinson JT. 1954a. Prehominid Dentition and Hominid Evolution. Evolution 8 (4): 324-334.

Robinson JT. 1954b. The Genera and Species of the Australopithecinae. American Journal of Physical Anthropology 12:. 181-200.

Shea BT. 1985. On Aspects of Skull Form in African Apes and Orangutans, With Implications for Hominoid Evolution. American Journal of Physical Anthropology 68: 329-342.

Smith RJ. 1981. On the Definition of Variables in Studies of Primate Dental Allometry. American Journal of Physical Anthropology 55: 323-329.

Smith RJ, Jungers WL. 1997. Body Mass in Comparative Primatology. Journal of Human Evolution 32: 523-559.

Strait DS, Richmond BG, Spencer MA, Ross CF, Dechow PC, Wood BA. 2007. Masticatory Biomechanics and its Relevance to Early Hominid Phylogeny: An Examination of Palatal Thickness Using Finite-Element Analysis. Journal of Human Evolution 52: 585-599.

Strait PW, Weber GW, Neubauer S, Chalk J, Richmond BG, Lucas PW, Spencer MA, Schrein C, Dechow PC, Ross CF, Grosse IR, Wright BW, Constantino P, Wood BA, Lawn B, Hylander WL, Wang Q, Byron C, Slive DE, Smith AL, Pilbeam D. 2009. The Feeding Biomechanics and

Dietary Ecology of *Australopithecus africanus*. Proceedings of the National Academy of Sciences of the United States of America 106 (7): 2124-2129.

Swindler DR. 2002. Primate Dentition: An Introduction to the Teeth of Non-Human Primates. Cambridge: Cambridge University Press.

Ungar P. 1998. Dental Allometry, Morphology, and Wear As Evidence for Diet in Fossil Primates. Evolutionary Anthropology: Issues, News, and Reviews. 6 (6): 205-217.

Ward SC, Kimbel WH. 1983. Subnasal Alveolar Morphology and the Systematic Position of Sivapithecus. American Journal of Physical Anthropology 61: 157-171.

Weston EM, Friday AE, Johnstone RA, Schrenk F. 2004. Wide Faces or Large Canines? The Attractive Versus the Aggressive Primate. Proceedings: Biological Sciences 271 (6): S416-S419.

White T. 2000. Human Osteology (Second Edition). San Diego: Academic Press.

Williams FL, Schroeder L, Ackermann RR. 2012. The Mid-Face of Lower Pleistocene Hominins and its Bearing on the Attribution of SK 847 and StW 53. HOMO - Journal of Comparative Human Biology 63: 245-257.

Wolpoff MH, Brace CL, Kay RF, Pilbeam D, Gould SJ. Allometry and Early Hominids. 1975. Science, New Series, 189 (4196): 61-64.

Wolpoff MH. 1985. Tooth Size-Body Size Scaling in a Human Population: Theory and Practice of an Allometric Analysis. In: Jungers WL, (Ed.), Size and Scaling in Primate Biology. New York: Plenum Press. p. 273-317.