Elsevier Editorial System(tm) for Journal of Human Evolution Manuscript Draft

Manuscript Number: T-307R1

Title: Enamel thickness trends in Plio-Pleistocene hominin mandibular molars

Article Type: Full Length Article

Keywords: enamel, relative enamel thickness, average enamel thickness, Australopithecus, Paranthropus, Homo, microCT

Corresponding Author: Dr. Matthew Skinner, Ph.D.

Corresponding Author's Institution: University of Kent

First Author: Matthew Skinner, Ph.D.

Order of Authors: Matthew Skinner, Ph.D.; Zeresenay Alemseged, Ph.D.; Charleen Gaunitz; Jean-Jacques Hublin

Abstract: Enamel thickness continues to be an important morphological character in hominin systematics and is frequently invoked in dietary reconstructions of Plio-Pleistocene hominin taxa. However, to date, the majority of published data on molar enamel thickness of Pliocene and early Pleistocene hominins derive from naturally fractured random surfaces of a small number of specimens. In this study we systematically analyze enamel thickness in a large sample of Plio-Pleistocene fossil hominins (n = 99), extant hominoids (n=57), and modern humans (n=30). Based on analysis of 2D mesial planes of section derived from microtomography, we examine both average and relative enamel thickness, and the distribution of enamel across buccal, occlusal, and lingual components of mandibular molars. Our results confirm the trend for increasing enamel thickness during the Pliocene that culminates in the thick enamel of the robust Australopithecus species, and then decreases from early Homo to recent modern humans. All hominin taxa, and Pongo, share a regional average enamel thickness pattern of thick occlusal enamel and greater buccal than lingual enamel thickness. Pan is unique in exhibiting thinnest average enamel thickness is a weak taxonomic discriminator. The data underlying these results are included as an appendix in the study.

We thank the editorial staff and the reviewers for their critical and helpful comments. Please find below a point by point explanation of how we addressed these comments with our revision.

Reviewers' comments:

AE Comments:

This is a very clean, easy to read manuscript that provides a thorough update to studies of enamel thickness in hominins. All three reviewers find the manuscript useful and praise its clarity, thoroughness, and brevity. However, the reviewers diverge in their opinions on how much revision is needed. Reviewer #3 finds the manuscript effectively acceptable as is, while the other two reviewers request some degree of revision. Reviewer #1 requests that the manuscript be revised with regard to "....inferences regarding the taxonomic valence and functional utility of enamel thickness..." Reviewer #2 focuses on the measurements of enamel thickness, and asks the authors especially to comment on why enamel is so thick in modern Homo, but not so in early Homo. The reviewer also offers some organizational suggestions.

The suggestions of the reviewers are certainly in the spirit of constructive advice, and should be relatively easy to manage. Given that the changes may involve extra analysis and an alteration in the primary thrust of the manuscript, I recommend that the authors revise the manuscript and reply to the reviewers comments. Pending the nature of the revisions, I suggest returning the ms to reviewer #1 if necessary.

Reviewer #1:

Overall, Skinner et al. provide a thorough and much needed compendium of enamel thickness in a range of early hominins, including early Homo. As such, it provides an extremely useful study that fills in the gap between good data available on hominoid enamel thickness and variation within species of Homo (i.e., Smith et al.). Examinations of this phenotype have a long history in both this field and this journal, and as such, topically it is appropriate for JHE. However, I do feel that in its current form, the ms tried to do too much, and in doing so, makes statements that may or may not be true; as there are not specific tests underlying such statements (see below), it leaves a reader not knowing how reliable these statements may be.

The one major drawback of this paper is the tension between being a wholly descriptive endeavor (and there's nothing wrong with that) and a series of inferential exercises (that would allow one to make specific statements about the taxonomic aspects of this feature) but are currently not present anywhere in the ms. I would strongly suggest that the ms be streamlined to focus on the descriptive part, as any inferences regarding the taxonomic valence and functional utility of enamel thickness are not tested, or hypotheses/predictions about these aspects stated explicitly. Engaging in the inferential part is essentially an exercise in making probabilistic statements, specifically about the likelihood that a species' phenotype (and variation therein) can confidently be distinguished from the range (and mean) expressed in other species. The authors do not do this - and again, that is okay - but instead conflate issues about degree of range overlap with probabilistic assessments of the taxonomic utility of that phenotype. So, statements such as "2D measurements of enamel thickness, which are commonly applied, are unreliable for definitive taxonomic distinction owing to the considerable overlap observed across taxa" (p. 15 1. 3-4) may be true. But as the authors know, two ranges can overlap quite a bit but still be statistically - and thus by extension, perhaps biologically - different.

We thank the reviewer for this comment. We believe strongly that including the statistical analysis of enamel thickness differences between species is an important contribution of this paper, as enamel thickness is so often implicated in taxonomic hypotheses. Furthermore, the samples for some of the hominin taxa are almost complete (e.g., Au. africanus and A. robustus) and are unlikely to increase soon and this lends credence to our conclusions about the likely lack of significant difference between taxa (on average). However, we accept that we may have overstated our results and therefore we have re-written the relevant parts of the manuscript to more accurately reflect the strength/weaknesses of our results. We also have tried to clarify/separate the descriptive and inferential aspects of our results.

Additionally, given the effort of others to produce 3D measures of enamel volume for comparative purposes, it might be useful to explain why these authors feel a 2D assessment is sufficient. Furthermore, is there a reason to suspect that different conclusions regarding species ranges, means, etc., would be reached by extending this into three dimensions?

It would certainly be ideal to include 3D data for these taxa as the reviewer is correct that the overall picture of taxonomic, metameric and crown specific enamel thickness results would likely differ when the whole enamel crown is scaled against the whole dentine crown. We restricted our analysis to 2D in order to maximize sample size as it is very difficult to reconstruct the 3D enamel tissue that is missing from much of the study sample. In a 2D section there is a limited range of possibilities for how the missing enamel can be reconstructed. But in 3D this becomes theoretically and practically (e.g., warping the vertices of a 3D surface model) very difficult. Acknowledging that a 3D study would be an excellent next step we have edited the manuscript to more clearly justify our reporting of 2D data only and added to the discussion the fact that our results might differ if based on 3D data (and that this should be the topic of future studies of this material). Having said this we believe that the two approaches can be complimentary to each other and possible discrepancies could open new avenue for further research.

I also have a series of other small editorial comments: Abstract: Not sure that "implicated" is the correct word. Perhaps "invoked"? Line 12: should read "early" not "Early" - that change should be made throughout the ms (e.g., p5, l. 7; p15, ls. 13, 15).

Both of these have been changed.

P4, 1. 4 - "mandibular" not "lower." This may be minor, and the editors may feel differently, but I always thought that "upper" and "lower" were too colloquial, and that "maxillary" and "mandibular" seemed more appropriate. Plus (as in p5, 1. 5, the authors use "mandibular"; they should pick one 'system' by which to indicate arcade and stick with that throughout the ms.

'lower' has now been changed to 'mandibular'

P4, 1. 7 - what does "thin section mounted" mean?

This sentence has been re-worded for clarity: Ward and colleagues (2001) report linear measurements of 1.0 - 2.1mm based on ground thin-sections of naturally fractured (and thin-sectioned mounted) specimens...

Is it possible to use a term other than "opposite-side cusps"? Perhaps stick with the same theme and say "non-functional cusps" - though, admittedly, I hate those terms as all cusps are "functional" to some degree.

Opposite-side has been changed to 'adjacent'.

P5, 1. 8 - remove comma after "as well as"

Done

P6, 1.8 - why is it relevant to point out that the authors are avoiding the debate over the monophyly of the robust australopiths? Whether or not they represent a monophyletic clade does not influence whether or not enamel thickness patterning is a useful taxonomic discriminator.

This sentence has been removed.

P8, 1. 7 - replace "we" with "was" - and I think the authors must provide more detailed information on how worn crowns were reconstructed. At the very least, a systematic methodology for doing so should be relayed. I appreciate the 'experiment' that they relay for Stw 308, but some readers would not characterize their measurement error of upwards of 5.2% as low, or acceptable. I am not suggesting that it isn't, I just think that there was likely some protocol that was followed that allowed the different researchers to reconstruct worn crowns, and that it is important for that protocol to be relayed in the Methods. And along those same lines, how much wear was tolerated; the authors use the phrase "partially worn" (p10, 1. 1), but this could mean anything from slight wear to a high degree of cuspal wear, as in KNM-ER 1802 (Fig. 2). Can more information be provided on their tolerance for wear as a limiting factor in their sample composition?

We agree with the reviewer that our method and criteria for reconstructing missing enamel was lacking. We have now added the following to the methods section:

As can be seen in the SI figures, in the majority of cases this involved very minor additions of missing enamel over one or more cusp tips. In a small number of cases between one half and one third of the enamel cusp was reconstructed. Reconstruction was guided by reference to the outer enamel surface (to determine based on the presence of wear facets, where enamel was missing), the curvature of the enamel cap cross-section. Also, specimens which did not preserve an intact central occlusal basin (in cross-section) were removed as having preserved enamel on each side of a worn cusp is necessary for a reasonable estimation of missing enamel. Only in the thinly enameled apes (i.e., Gorilla and Pan) whose enamel distribution is quite uniform and whose dentine horns are relatively sharp, was it deemed acceptable to reconstruct missing tips of dentine horns.

We have also edited the text on p.10 to point out that the degree of applied artificial wear used for our test is at the extreme end compared to most of the study sample and in some sense this is a worst case scenario (e.g., thick enamelled taxon and marked wear) for a specimen that would have fit our criteria for reconstruction.

P10, 1. 15 - should be singular, "hominin"; same for 1. 17.

Done

P12, 1. 21 - comma after "crown"

Done

P14, l. 21 - comma after "species." In fact, this first sentence of this section is a bit awkward and could do with it being broken into two sentences.

Split and reworded.

P15, l. 1 - "thick" not "think" Here the authors state the 2D measures are not reliable for taxonomic purposes. So, are 3D measures better in this regard? Or is enamel thickness per se just not a useful tool for taxonomic discrimination? Please see my general comments above about the descriptive vs. the inferential.

Agreed. We have changed this sentence as follows:

Overall, the results of our analysis suggest that within the hominin clade, 2D measurements of enamel thickness may be unreliable for definitive taxonomic distinction. Specifically, the results of our statistical analysis can be used as a guide (taking into account sample size and variation within taxa) as to which taxonomic comparisons within the hominin clade are likely to yield informative taxonomic discrimination.

We feel that our current data, which are 2D, do not allows us to comment on whether 3D measures are better or not. We have still added a paragraph to this section of the discussion to address 3D vs 2D data.

If the authors insist on a probabilistic assessment of 'lack of taxonomic utility' than I would urge them to address that specifically within their analytical design. Furthermore, I would caution the authors that they should not discount the value of their particular phenotype for making taxonomically meaningful inferences, and then go ahead and use it anyway (for the isolated specimens).

I am not suggesting that the observations and explanations about enamel thickness variation laid out on p15 are unimportant. On the contrary, they are quite useful. Just that this whole section is at odds with the beginning of the Discussion as it is set against a backdrop relaying the taxonomic uselessness of enamel thickness.

Agreed. This section has been re-worded and our consideration of the utility of 2D enamel thickness more accurately reflects the results of our analysis (and is less pessimistic).

P15, 1. 21 - "adaptation" should be plural

Done

P16, 1. 5 - the use of the bridge "and a further increase in" in this sentence makes it somewhat confusing.

Reworded

p.16, l. 20. "Our results indicate that the majority of fossil hominins do exhibit greater enamel thickness buccally than lingually..." Of course they found that, the authors only examined mandibular molars, and this is a basic structural feature of most mammalian molars given the nature of the chewing cycle.

The phrase 'as expected' has been added to this sentence.

P16 -- I don't find the last section particularly strong; there are just some very broad statements attempting to link this phenotype to some broad notion of 'diet' or 'function'. In the first paragraph, they attribute the trend towards increasing enamel thickness to an increase in the incorporation of C4 foods. Fine. Then in the following paragraph, they link the same trend (at least for some australopiths) to an increase in the complexity of enamel surfaces. How so the authors envisage these two things (increasing C4 signal and increasing complexity of occlusal surfaces) as being linked? Are C4 foods (grasses, sedges, succulents, etc. - if you include CAM pathways foods) things that would leave a greater complexity signal? Or is it perhaps that increasing enamel thickness (i.e., global enamel thickness, as that is what AET assesses) is linked to increasing dietary breadth, which could manifest as an increase in a C4/CAM signal? As it stands, it is just not clear how the authors are linked two very different measures (isotopes and microwear textures) to the same trend in enamel thickness.

This section has been reorganized and expanded. In particular, the discussion of relevant isotopic data and microwear complexity has been split from the discussion of regional distribution of enamel. We have also attempted to clarify how current isotopic/microwear data are associated with the trends in enamel thickness found in our study.

One note about the tables:

Do the blank cells in Table 4 indicate a non-significant difference between taxa, or that sample sizes were too small to perform any statistical test? (I imagine the latter given the sample sizes by molar position listed in Table 1, right?)

All pairwise comparisons were calculated (even in the few cases with sample sizes of 1-2) and blank cells indicate non-significant results. This has now been clarified in the caption of Table 4.

Reviewer #2: This paper presents comprehensive quantification of enamel thickness for a comprehensive taxonomic sample of pre-homo fossil hominins, a limited sample of archaic homo, and with an adequate hominoid comparative sample.

I am not enough of an expert to verify the accuracy of your claim about most enamel thickness measurements being based on opportunistic fractures.

In your review of published data (page 3 line 19 and all of page 4) a summary table might help. You should include means given for the previous studies in addition to ranges.

We acknowledge that a summary table would be ideal, however, the lack of data at particular tooth positions for most taxa, the range of methods employed, and the opportunistic nature of the available data (i.e., some naturally fractured surfaces, some from ground thin sections) limits, in our view, the utility of a summary table in this case.

The methodological description is easy to follow and seems sound. I think it is okay that you visually reconstruct broken segments of the enamel surface in some cases to increase the sample size, because your error study confirms that its repeatable and not inaccurate. The inclusion of individual measurement data, images showing section location and actual cross-section used for measurements of most specimens is a very positive feature of this work as it promotes incorporation of these data into future studies that wish to evaluate similar data collection methods on expanded samples. It also improves researchers' ability attempt to reproduce results if they desire.

One suggestion about enamel thickness evaluation... why not use a relative enamel thickness metric that is the square-root of enamel area divided by the length of the edj? This will give you a ratio that basically indicates whether the enamel cap looks more like a thin ribbon or a thick strap, which I think is what the human eye is gauging when it qualitatively makes an assessment about thickness. Your method for RET can be affected by additional un-investigated variables, like the shape of the EDJ. It is possible this different formulation of RET will even be somewhat independent of the presently used one as a result, and can be used in addition to it as a third way of evaluating enamel thickness.

Calculating relative enamel thickness as the square-root of enamel divided by the length of the EDJ is an interesting suggestion. However, we believe it is most appropriate to report measurements based on a protocol that is well established in the anthropological community. In our opinion, a comparison of the suggested approach and current methods would be best served by a more technically oriented paper (e.g., Benazzi et al., 2014) and is beyond the scope of this project.

The error studies are well-constructed and imbue confidence that methods are repeatable.

Analysis - when group sizes are less than 3, they should probably not be included in ANOVA's (eg Au. afarensis is represented by n=2 for m1, but included in ANOVA comparisons of Table 4).

We acknowledge that for the Kruskal-Wallis test across the whole sample the inclusion of taxa with n=2 are problematic, however, we would like to include them as they do indicate statistically significant differences between a number of pairwise comparisons that will be of interest to the anthropological community (e.g., that Homo sp indet differs significantly from modern humans).

Page 6, line 7-9: must be a more professional way of saying this.

This sentence has been removed.

Page 7, line 6: please provide a proper citation for Aviso software and include it in the bibliography

Following the JHE article Smith et al., 2012, we have changed this to Avizo (v6.3, FEI Inc.)

Page 7, line 14: I think it is confusing to call area of a 2D cross-section "surface area" just say "area" or "exposed-section area" If you say "surface area" some readers will confused thinking you are looking at the whole enamel surface and dentine surface

'surface' has been removed.

Page 13, lines 11-22 and page 14, lines 1-18: I think this discussion may be more appropriate in the introduction to help explain the motivation behind your methodology. You might consider citing Boyer (2008) who justified using a more subjective method of defining the portion of the tooth crown to be used for relief index calculation, with regard to the point that using the more objective approach (of Ungar and others in the case of relief index) resulted in tooth orientations and measurements that were not broadly homologous for the sample at hand.

We thank the reviewer for this suggestion. As the main goal of the paper is to examine enamel thickness trends, rather than methodological issues, we would request to keep this topic in the discussion. We have now included Boyer (2008) in this section.

Page 15, lines 5-7 (as an example) - in this section you explain how certain variables they measured support or are consistent with a particular hypotheses, but say nothing about other, leaving the reader wondering. In the lines mentioned, you state that KNM WT-8556 falls in the range of Au africanus and Au afarensis in tooth size and AET. However, what about RET? Please at least mention that it is either outside the range - or inside but non-distinctive. Also this statement is vague - for the uninitiated, we don't know what your point is. Does this observation support an assignment to K. platyops or refute it? You don't actually tell us what your interpretation is. Apply the issue in this example to the other cases you mention in this section as well.

As also noted also by reviewer 1 we agree that this section was vague. We have added additional discussion of particular specimens and been specific about the implications of our results for each discussed specimen.

Page 16, lines 1-9 - would it be possible to do a correlation analysis on species mean enamel thickness and delta C13 values gleaned from the literature (like the Cerling papers?). It would be awesome if you could report a significant pearson correlation coefficient or something.

This is a great idea. Unfortunately, having consulted the supplementary information in Cerling et al., 2013 there is very little overlap in specimens (presumably because they were only given permission to destructively sample the less well preserved specimens). Only KNM-ER 1802B, 820 and 992, and KNM-WT 8556 were sampled. Therefore, we did not pursue this very interesting suggestion.

Coming to your conclusion, I think something is still missing from you analyses and discussion. You certainly demonstrate both relative and average enamel thickness increase through australopithecine evolution. But why is Homo sapiens so high and early homo so low in thickness?

We are slightly confused by this comment as our results state the opposite (i.e., recent Homo sapiens have thinner enamel than early Homo. We also refer to how are results for the Homo sp material from Omo is consistent with the findings of thick enamel in early Homo published by Smith et al., 2012.

Also it would still be nice to know how much of the AET variation is explained by tooth size versus RET. Couldn't you run a multiple regression with AET as the dependent variable and the other two as independents and comment on this more explicitly? If you showed that tooth size contributed substantially less to the variance, that would be interesting because you could argue that whatever dietary shifts happened put selective pressures on both absolute tooth size and on proportional enamel thickness. Winchester et al (2014) discuss increasing enamel thickness, tooth size, and hyspodonty as different strategies of achieving a similar goal - to put more enamel in the mouth and increase the "lifetime" of the tooth (and consequently its owner) in the face of an abrasive diet. So here you have an opportunity to comment on how much the evolutionary response for increasingly tough foods was expressed through tooth size increase versus proportional enamel thickness increase. This is an interesting idea, however, we are not convinced of the statistical meaning of including RET as one of the independent variables as it is derived from AET (being simply the quotient of AET/SQRT of dentine area. We tried using tooth size and dentine area, however, even the latter is not independent from AET which is derived using EDJ length. For the purpose of this manuscript, and given the theoretical difficulties in finding a ethick measure that is not correlated with tooth size, we have chosen not pursue this line of investigation for the moment.

I also really think that RET as a ratio of sqrt(enamel area)/(edj length) will be a better reflector than (enamel area/dentine area).

See above

I did not check your bibliography for errors.

Overall good work.

Reviewer #3: This is a nice paper on enamel thickness. I found it clearly written and informative. The review of the topic is certainly of interest to JHE readers. I have only the most minor comments: On page 4, perhaps the authors could explain linear measurements and radial linear measurements. I assume this is just taking a measurement of the exposed enamel, but I am not certain. Perhaps a figure could clarify?

Radial thickness measurements are now explained in the text.

On Page 5 I would refer to 'modern humans' as 'recent humans' just to be clear, since "modern" could include fossil H.s.

Done

On page 9, line 15 needs a comma after 'which' **Done**

Also on page 9 the last full sentence (lines 21 and 22) is a bit awkward to read.

This sentence has been reworded.

Page 10, there is a type-o in line 1 (ordedr)

Corrected

Page 10 lines 14-15 - this sentence is difficult to read.

Reworded

Page 14, line 22 needs a comma after 'which'

Reworded

Other than these minor suggests I see no issues with publishing the manuscript as is.

1	
2	Title: Enamel thickness trends in Plio-Pleistocene hominin mandibular molars
3	
4	
5	
6	
7	
0	
8 9	Matthew M. Skinner ^{1,2,3*} , Zeresenay Alemseged ⁴ , Charleen Gaunitz ² , Jean-Jacques Hublin ²
10	
11	
12 13	¹ School of Anthropology and Conservation, University of Kent, Canterbury, CT2 7NR, United Kingdom
14 15	² Department of Human Evolution, Max Planck Institute for Evolutionary Anthropology, Deutscher Platz 6, Leipzig 04103, Germany
16 17	³ Evolutionary Studies Institute and Centre for Excellence in PaleoSciences, University of the Witwatersrand, Private Bag 3, WITS 2050, South Africa.
18	⁴ Department of Anthropology, California Academy of Sciences, San Francisco, 94118, USA
19	
20	*corresponding author
21	
22	
23 24	Keywords: enamel, relative enamel thickness, average enamel thickness, <i>Australopithecus</i> , Paranthropus, <i>Homo</i> , microCT

1 Abstract

2 Enamel thickness continues to be an important morphological character in hominin systematics and is frequently invoked in dietary reconstructions of Plio-Pleistocene hominin taxa. 3 4 However, to date, the majority of published data on molar enamel thickness of Pliocene and early 5 Pleistocene hominins derive from naturally fractured random surfaces of a small number of specimens. In this study we systematically analyze enamel thickness in a large sample of Plio-6 7 Pleistocene fossil hominins (n = 99), extant hominoids (n=57), and modern humans (n=30). Based 8 on analysis of 2D mesial planes of section derived from microtomography, we examine both 9 average and relative enamel thickness, and the distribution of enamel across buccal, occlusal, and lingual components of mandibular molars. Our results confirm the trend for increasing enamel 10 11 thickness during the Pliocene that culminates in the thick enamel of the robust Australopithecus 12 species, and then decreases from early Homo to recent modern humans. All hominin taxa, and 13 Pongo, share a regional average enamel thickness pattern of thick occlusal enamel and greater buccal than lingual enamel thickness. *Pan* is unique in exhibiting thinnest average enamel 14 15 thickness in the occlusal basin. Statistical analysis indicates that among Pliocene hominins enamel thickness is a weak taxonomic discriminator. The data underlying these results are included as an 16 17 appendix in the study.

1 Introduction

The thickness and distribution of enamel tissue across tooth crowns remains an important 2 3 character in assessments of the taxonomy, phylogeny, and dietary reconstructions of fossil primates. Within the hominoid clade, over three decades of research has elucidated patterns of 4 5 enamel thickness variation in fossil hominins (e.g., Martin, 1985; Beynon and Wood, 1986; Grine 6 and Martin, 1988; Conroy, 1991; Macho and Thackeray, 1992; Schwartz et al., 1998; Brunet et al., 7 2002, 2005; Olejniczak and Grine, 2005; Smith et al., 2006b; White et al., 2006; Olejniczak et al., 8 2008a/b; Smith et al., 2009a/b, 2012a), fossil hominoids (e.g., Martin et al., 2003; Smith et al., 9 2003, Olejniczak et al., 2008c), and extant hominoids (Molnar and Gantt, 1977; Gantt, 1986; Grine, 10 1991; Schwartz, 2000; Kono, 2004; Tafforeau, 2004; Smith et al., 2005, 2006a; Kono and Suwa, 2008; Olejniczak et al., 2008d; Smith et al., 2012b). Many of these studies dating to the last decade 11 12 have utilized microtomography to systematically produce homologous mesial planes of section in 13 molars, which has led to more rigorous taxonomic comparisons (see review in Smith et al., 2012a). However, due to inherent practical and methodological difficulties in producing microtomographic 14 15 scans of their dentitions, systematic analysis has not been conducted on the majority of otherwise 16 extensively investigated Pliocene hominin taxa. In this contribution we fill in this gap for many 17 species of the genus Australopithecus and complement the extensive review recently published by 18 Smith and colleagues (2012a) for Pleistocene Homo. To date the majority of reported enamel thickness values for Pliocene hominins derive 19 20 from linear measurements taken on naturally cracked surfaces of molars. For example, White and

colleagues (1994) report linear measurements of Ardipithecus (Ar.) ramidus molars ranging from

22 1.1-1.2mm and for Australopithecus (Au.) afarensis of 1.4-2.0mm. Based on microtomography,

Suwa and colleagues (2009) reported Ar. ramidus as having enamel thickness greater than Pan but 1 2 thinner than later Australopithecus. Johanson and colleagues (1982) and White and colleagues 3 (2000) report linear dimensions for various Au. afarensis specimens but do not report any 4 measurements for mandibular molars. In the initial publication of the Au. anamensis specimens, 5 linear measurements of upper and mandibular molars ranged between 1.5 and 2.0mm (Leakey et 6 al., 1995). Ward and colleagues (2001) report linear measurements of 1.0 – 2.1mm based on 7 ground thin-sections of naturally fractured (and thin-sectioned mounted) specimens (upper molar 8 KNM-ER 30748 and mandibular molar KNM-ER 30749) in the occlusal basin, cusp tip and lingual 9 and buccal walls. The Au. anamensis finds from Asa Issie exhibit radial (i.e., measured not in a 10 mesial plane of section but rather along a trajectory running perpendicular from the dentine 11 surface to the enamel surface) linear measurements of 1.7 – 2.3mm for functional cusps (i.e., buccal cusps on mandibular molars and lingual cusps on upper molars) and 1.3 - 2.0 mm for 12 13 adjacent cusps (White et al., 2006). Haile-Selassie and colleagues (2010) assessed enamel thickness 14 in the Woranso-Mille material from naturally fractured molars and concluded that the range (1.5-2.1mm) falls within the range of reported measurements for Au. afarensis, Au. anamensis and Au. 15 africanus. In their analysis of crown formation times Lacruz and Ramirez Rozzi (2010) report linear 16 17 enamel thickness measurements of 1.95mm (AL 333-52), 2.13mm (AL 366-1), and 1.71mm (Omo L2-79). Examining Au. africanus specimens, Grine and Martin (1988) report average enamel 18 19 thickness values of 1.81mm (Stw 284; now referred to as Stw 280) and 1.78mm (Stw 402), and relative enamel thickness values of 21.27 (Stw 280) and 23.06 (Stw 402). Macho and Thackeray 20 21 (1992) used medical CT to examine the regional distribution of enamel thickness across the crowns 22 of Au. robustus, Au. africanus, and Homo sp. maxillary molars finding considerable overlap

between taxa in many regions of the crown. Finally, Olejniczak and colleagues (2008b) published
data on *Au. africanus* and *Au. robustus* from South Africa, expanding their analysis to 3D enamel
distribution across the crown. Collectively, however, the limited sample size, limited assessment of
enamel thickness (i.e., often linear measurements), and variation in location of measurement
result in a poor characterization of enamel thickness variation along the molar row in Pliocene
hominins.

7 Using microtomography and controlled mesial planes of section in mandibular molars, we 8 analyze enamel thickness to assess taxonomic differences in mandibular molar crowns of Au. 9 anamensis, Au. afarensis, Au. africanus, Au. boisei, Au. robustus, and specimens of early Homo. We compare these results to samples of Pan, Gorilla, and Pongo, as well as a sample of recent 10 11 humans. The goals of this study are to: 1) analyze enamel thickness variation among Plio-Pleistocene homining using a 2D mesial plane of section; 2) characterize the distribution of lingual, 12 13 occlusal and buccal enamel among hominin taxa; 3) assess the reliability of taxonomic 14 discrimination based on enamel thickness measured in a 2D section; 4) evaluate the affinity of 15 taxonomically ambiguous specimens based on their enamel thickness values; and 5) provide molar-specific enamel thickness measurements for extant apes and fossil hominins for use by 16 other researchers. 17

18

19 Materials

The study sample consists of mandibular molars (n = 186) belonging to both extant hominoids and fossil hominins and is detailed in full in Appendix 1. The number of first, second and third molars of each taxon is listed in Table 1. This sample is the largest compiled to date for a

systematic analysis of enamel thickness in Plio-Pleistocene hominins of Africa. Molars either derive
from mandibles or are isolated specimens. In the case of the latter, the justification for assigning a
molar to a particular position is also noted. Specimens were chosen for study that exhibited no
evidence of known pathology. Given that sex is unknown for the majority of fossil specimens it was
not incorporated into our analysis as a variable.

Hominoid taxa include *Pongo* sp., *Gorilla* sp., *Pan paniscus* and *Pan troglodytes* ssp. Due to
the small sample sizes for some molar positions no species delineation was made for *Pongo* and *Gorilla* and no subspecies delineation for *Pan troglodytes*. The Plio-Pleistocene hominin taxa
include *Au. anamensis*, *Au. afarensis*, *Au. africanus*, *Au. aethiopicus*, *Au. boisei*, *Au. robustus*, *Homo*sp. indet., *H. erectus*, and modern *H. sapiens*. A number of specimens of uncertain taxonomic
affinity were also analyzed and their taxonomic affinity assessed based on their measured enamel
thickness values.

13 Fossil hominin specimens derive from collections housed at the following institutions: 14 National Museum of Ethiopia, Addis Ababa, Ethiopia; National Museums of Kenya, Nairobi, Kenya; University of Witwatersrand, Johannesburg, South Africa; Ditsong National Museum of Natural 15 History, Pretoria, South Africa. The hominoid samples derive from the Museum for Natural History 16 17 (ZMB), Berlin, Germany; the Senckenberg Research Institute (SMF), Frankfurt, Germany; the Royal Museum for Central Africa (MRAC), Terverun, Belgium; and the Max Planck Institute for 18 19 Evolutionary Anthropology (MPI), Leipzig, Germany. The modern human sample derives from the 20 Leipzig University Anatomical Collection (ULAC), Leipzig, Germany; the "Francisc J. Rainer" 21 Anthropology Institute (R), Bucharest, Romania; and the Max Planck Institute for Evolutionary 22 Anthropology, Leipzig, Germany.

1

2 Methods

To obtain a 2D mesial plane of section each molar was non-destructively imaged using 3 computed tomography (using either a BIR Actis 300/225 FP or SkyScan 1172 microtomographic 4 scanner) with a resultant isometric voxel size of 15-65 μ m³. The CT data set of each specimen was 5 6 rotated manually in Avizo (v6.3, FEI Inc.) into anatomical position. Next, a plane was placed 7 perpendicular to the occlusal plane and passing through the tip of the protoconid dentine horn. 8 This plane was then rotated to pass through the tip of the metaconid dentine horn. This slice 9 image was then saved in TIFF format (Figure 1). Benazzi and colleagues (2014) have outlined a methodology to produce repeatable 2D planes of section. This methodology was not adopted for 10 11 this study because it is difficult to apply to many of the fragmentary hominin teeth used in this 12 study whose cervical line is not preserved (and see Discussion). 13 Four variables were measured on each mesial section using ImageJ (v1.47, NIH): area of the enamel cap (mm²), area of the coronal dentine crown (mm²) delimited by a line drawn between 14 15 the most cervical enamel extensions, length of the enamel-dentine junction, or EDJ (mm), and bicervical diameter (mm) also measured between the most cervical enamel extensions. These 16 17 measurements are listed for each specimen in Appendix 1. In order to assess regional differences 18 in enamel thickness buccolingually across the tooth crown the mesial crown section was divided 19 into lingual, occlusal and buccal components. This division was accomplished in ImageJ by connecting the tip of each dentine horn to the corresponding tip of the cusp at the outer enamel 20 21 surface. Figure 1 illustrates the measurement locations.

1 The Supplementary Information contains figures of the majority of hominin specimens 2 measured as well as a sample of extant hominoids and modern humans, illustrating the location of 3 the plane of section and the delineation of the enamel and dentine components of the section. 4 This is particularly important as it allows researchers to assess our placement of the plane of 5 section. In a number of cases, and particularly in the fossil hominin sample, missing enamel over 6 cusp tips was reconstructed in the mesial section. As can be seen in the SI figures, in the majority 7 of cases this involved very minor additions of missing enamel over one or more cusp tips. In a 8 small number of cases between one half and one third of the enamel cusp was reconstructed. 9 Reconstruction was guided by reference to the outer enamel surface (to determine based on the 10 presence of wear facets, where enamel was missing), the curvature of the enamel cap cross-11 section. Also, specimens which did not preserve an intact central occlusal basin (in cross-section) 12 were removed as having preserved enamel on each side of a worn cusp is necessary for a 13 reasonable estimation of missing enamel. Only in the thinly enameled apes (i.e., Gorilla and Pan) 14 whose enamel distribution is quite uniform and whose dentine horns are relatively sharp, was it 15 deemed acceptable to reconstruct missing tips of dentine horns. While this reconstruction of 16 missing enamel introduces a subjective component into our analysis, it is worthwhile for 17 characterization of enamel thickness trends within the hominin clade as the number of absolutely unworn hominin teeth is very small. In almost all cases a researcher can evaluate our 18 19 reconstruction of each specimen in the supplementary material and since we provide all of the raw 20 data, they are able to drop specimens from the sample and re-calculate sample statistics for their 21 own purposes.

22

1 Quantitative analyses

2 We calculated two standard measures of enamel thickness following well established 3 protocols (Martin, 1985; Olejniczak et al., 2008a). Average enamel thickness (AET) was calculated 4 as the area of the enamel cap divided by the length of the EDJ. This yields the average straight line 5 distance from the EDJ to the enamel surface in millimeters. Relative enamel thickness (RET) was 6 calculated as AET divided by the square root of dentine area and multiplied by 100. This yields a 7 scale-free value of enamel thickness that allows comparisons between taxa of differing tooth/body 8 size. For the assessment of regional variation in AET across the tooth crown we divided the surface 9 area of the enamel for each region by its corresponding EDJ length. Plots of the log of AET against the log of dentine area were used to illustrate the relationship between AET and tooth size and to 10 11 visualize the placement of specimens of uncertain taxonomic affinity. Significant differences in AET and RET between the study taxa were assessed in SPSS 20 using a Kruskal-Wallis Test with posthoc 12 13 pairwise comparisons.

14 Intraobserver error in AET and RET was assessed by MMS and CG each repeating the complete processing sequence (including rotation and mesial section derivation) for two 15 specimens 10 times each, over a period of three months. Interobserver error rates (calculated as 16 the difference in the measurement of MMS and CG divided by the average of their measurements) 17 were 2.4% (AET) and 3.56% (RET) for a modern human specimen and 0.6% (AET) and 1.66% (RET) 18 19 for an Au. robustus specimen. Intraobserver error (calculated as the average deviation from the mean of 10 measurements of a modern human specimen) was 1.3% (AET) and 1.4% (RET) for MMS 20 21 and 0.9% (AET) and 1.1% (RET) for CG. These values are considered acceptable and establish the 22 repeatability of the protocol. We also compared our measurements of particular specimens with

those from a previous study (Olejniczak et al., 2008a) and noted mean differences of between 3.1 1 2 7.1%. In most cases, variation is due to differences in locating the bi-cervical line, which can affect RET in particular due to the marked effect of changes in coronal dentine surface area. 3 4 We also tested the potential impact of the inclusion of partially worn teeth by artificially 5 wearing one of the unworn thick-enameled hominin teeth (STW 308) and blindly reconstructing the missing enamel (Supplementary Figure 1). This reconstruction was conducted five times on 6 7 different occasions. The range of measured enamel area was 40.1-43.0mm, resulting in a range of 8 calculated AET of 1.81-1.94mm. This results in a difference in AET of 1.0-5.2%. Given the high 9 degree of applied artificial wear (that is at the extreme end relative to the majority of specimens in the study sample) in this thick enameled specimen this test is essentially the worst case scenario 10 11 for a specimen that would have fit our criteria for reconstruction, and this level of error supports 12 the inclusion of partially worn specimens in the analysis in order to supplement sample size and 13 improve the characterization of enamel thickness in fossil hominin species. 14 Results 15 Figure 2 presents a selection of second molars from the majority of the study taxa. 16 Additionally, Supplementary Figures 2-19 illustrate 2D planes of section, measured enamel and 17 dentine area, and the position of the plane of section for all hominin molars and the majority of 18 19 the extant comparative sample. Table 2 lists the mean and standard deviation calculated for each of the four measured variables, AET, and RET for each taxon at each molar position. Table 3 lists 20 the values of measured variables for the specimens of unknown taxonomic affiliation. Table 4 lists 21 22 the results of the Kruskal-Wallis posthoc pairwise comparisons for AET and RET among the study

1	taxa at each molar position. Although across the study taxa all tests of AET and RET are highly
2	significant ($p = <0.001$), pairwise comparisons reveal that this result is driven primarily by
3	significant differences between the extant apes (Pan, Gorilla, Pongo, and recent humans) on the
4	one hand, and the thick enameled Australopithecus species on the other. Within fossil hominin
5	first molars, only Au. anamensis is significantly thinner in AET than Au. robustus. Within fossil
6	hominin second molars, Au. anamensis is significantly thinner than both Au. boisei and Au.
7	robustus in AET and RET. Additionally, second molars of Au. africanus are significantly thinner than
8	Au. robustus in RET. For third molars, Au. anamensis is significantly thinner than both Au. boisei
9	and Au. robustus in AET and thinner than Au. robustus in RET. Au. boisei third molars are
10	significantly thicker than <i>H. erectus</i> in AET.
11	Figure 2 here
12	Table 2 here
13	Table 3 here
14	Figure 3 presents box plots of AET across the study sample. There is a clear trend of
15	increasing AET from Au. anamensis to Au. africanus and a general increase in AET from first to
16	third molars in these taxa. Au. boisei and Au. robustus exhibit the highest AET values, with the
17	second molar being the thickest on average in Au. robustus. AET in Homo sp. is comparable with
18	Au. africanus and then there is a marked decrease in H. erectus and then modern humans. Of the
19	extant apes, <i>Pongo</i> presents the thickest AET at each molar position and <i>Pan</i> the thinnest (with
20	Gorilla intermediate). Figure 4 presents box plots of RET across the study sample and highlights a
21	broadly similar trend of increasing thickness in Australopithecus, followed by a decrease in Homo.
22	As RET is essentially scaled by tooth size there is greater overlap with large toothed taxa, such as

Au. africanus, being more similar to Au. anamensis in RET, than AET. Similarly, the position of 1 2 Gorilla and Pan shifts as the former's enamel thickness is relatively smaller than the latter's after scaling for tooth size. 3 Figure 3 here 4 5 Figure 4 here Table 4 here 6 7 Table 5 lists the mean and standard deviation of buccal, occlusal, and lingual 8 measurements of AET for each of the study taxa. Figure 5 illustrates the pattern of regional 9 variation in a combined molar sample for each taxon. The majority of study taxa present a consistent regional distribution with AET concentrated in the occlusal basin and a slight dominance 10 11 of the buccal side over the lingual side. Gorilla is unique in presenting thickest AET buccally (and 12 decreasing from occlusal to lingual), while *Pan* possesses thinnest enamel in the occlusal basin. 13 Figure 5 here Table 5 here 14 Figures 6-8 present bivariate plots of the log of AET against the log of dentine area for first 15 second and third molars, respectively. In essence, this is a visual representation of RET. Generally, 16 the distribution of taxa is consistent in first, second and third molars with Gorilla being 17 characterized by thin enamel covering a large dentine core. Although broadly overlapping in tooth 18 19 size, Au. boisei and Au. robustus exhibit thicker AET than Au. africanus. In second and third molars, Au. afarensis tends to exhibit relatively thicker AET than Au. anamensis. Pan is consistently 20 21 positioned and exhibits thin enamel over its relatively small molars. With regard to the first molar

specimens of uncertain taxonomic affinity (Table 3), KNM-WT 8556 falls within the range of Au.

1	africanus and close to Au. afarensis. Omo K7-19 has thick AET similar to Homo sp. and Au.
2	robustus, while Omo L26-1g sits between the convex hulls of Au. robustus and Au. africanus.
3	Second molars L28-31 and L795-1 have relatively thick AET for the size of their dentine crown and
4	fall in proximity to a cluster of Au. robustus, Au. boisei, and Au. aethiopicus. Finally, L28-30 exhibits
5	thick AET for its size, while Omo 75s-16 falls near modern humans and <i>H. erectus</i> .
6	Figure 6 here
7	Figure 7 here
8	Figure 8 here
9	
10	Discussion
11	Defining and quantifying enamel thickness
12	Since the first analyses of naturally fractured tooth crowns, the definition of enamel
13	thickness and the way it is measured have been in flux and varied substantially from one author to
14	another. As a consequence, though enamel thickness has frequently been used to interpret
15	taxonomy and diet in hominin fossils, methods used to quantify it have been less than satisfactory
16	in many cases. Researchers depended, short of other options, on naturally fractured surfaces
16 17	in many cases. Researchers depended, short of other options, on naturally fractured surfaces where the plane of breakage is random, resulting in non-comparable metric data. Also enamel
16 17 18	in many cases. Researchers depended, short of other options, on naturally fractured surfaces where the plane of breakage is random, resulting in non-comparable metric data. Also enamel thickness data were not necessarily derived from the same cusps, tooth type or sides of the teeth,
16 17 18 19	in many cases. Researchers depended, short of other options, on naturally fractured surfaces where the plane of breakage is random, resulting in non-comparable metric data. Also enamel thickness data were not necessarily derived from the same cusps, tooth type or sides of the teeth, questioning the biological homology of the measured data. None the less, recently developed
16 17 18 19 20	in many cases. Researchers depended, short of other options, on naturally fractured surfaces where the plane of breakage is random, resulting in non-comparable metric data. Also enamel thickness data were not necessarily derived from the same cusps, tooth type or sides of the teeth, questioning the biological homology of the measured data. None the less, recently developed imaging and visualization methods have resulted in more systematic assessments of enamel
16 17 18 19 20 21	in many cases. Researchers depended, short of other options, on naturally fractured surfaces where the plane of breakage is random, resulting in non-comparable metric data. Also enamel thickness data were not necessarily derived from the same cusps, tooth type or sides of the teeth, questioning the biological homology of the measured data. None the less, recently developed imaging and visualization methods have resulted in more systematic assessments of enamel thickness variation (e.g., Schwartz et al., 2000; Kono, 2004; Tafforeau, 2004; Kono and Suwa, 2008;

quantify enamel distribution across the tooth or in particular regions of the tooth crown, and allow
 for collecting and combing data on large fossil data sets.

3 Benazzi et al (2014) have published a revised CT-based methodology, the goal of which is to remove as much subjectivity in defining a mesial plane of section as possible. This method focuses 4 5 on the cervix as a means of defining a basal plane from which a perpendicular plane can be derived 6 and placed at the intersection of particular dentine horns. Their methodology is an important step 7 forward in developing published data that can be used by other researchers, however, it can be 8 challenging to apply the method to fragmentary fossil teeth. In particular, partial crowns may 9 preserve a mesial section but lack the distal portion of the cervix. In order to include such 10 specimens they would have to be oriented manually, albeit virtually, as is done in this study. Also, 11 when cervical enamel is missing (which is quite common in fossil specimens) these regions of the 12 cervix will have to be estimated.

13 There are also many examples of fossil/modern teeth whose pattern of enamel extension 14 around the circumference of the crown (and by consequence the cervix) is abnormal or results in 15 the creation of a plane of section that is not biologically homologous. Also, the approach of 16 Olejniczak (2006) that uses a plane fit to three dentine horns can result in the measurement of 17 non-homologous planes of section due to variation in relative dentine horn height (which should not be directly associated with enamel thickness). If the theoretical basis of the 2D plane of section 18 19 is to capture functionally and/or developmentally (sensu Butler, 1956) relevant enamel thickness 20 values associated with occlusion (and thus perpendicular to the occlusal plane), then the cervix 21 cannot always be relied upon to produce this plane. We would caution that in the absence of a 22 critical evaluation of the plane produced, systematic variation can be introduced in enamel

thickness measurements that exceeds normal levels of inter- and intraobserver error. Thus, we
stress the importance of biological homology as a defining principle in developing measurement
protocols (see also Boyer, 2008). Future studies should test the comparability of the method
outlined by Benazzi et al (2014) and a basal plane oriented manually (as was done in this study). If
it is found to result in acceptably small differences in measured AET and RET, then researchers can
be confident in combing manually oriented specimens when it is necessary to do so to produce an
homologous plane of section.

8

9 Enamel thickness and its role in taxonomic determination

10 Enamel thickness has been widely used to diagnose hominin species. In particular, the 11 geochronologically earliest ones are expected to possess at least moderately thicker enamel compared to that of extant African great apes, while 'robust' australopiths are characterized as 12 13 having very thick enamel. Overall, the results of our analysis suggest that within the hominin clade, 14 2D measurements of enamel thickness may be unreliable for definitive taxonomic distinction. Specifically, the results of our statistical analysis can be used as a guide (taking into account 15 16 sample size and variation within taxa) as to which taxonomic comparisons within the hominin 17 clade are likely to yield informative taxonomic discrimination. And the database of individual measurements we provide can be used to statistically test particular taxonomic hypotheses or 18 19 determine whether enamel thickness can be used to inform the affinity of newly 20 discovered/measured specimens. Acknowledging these potential shortcomings, AET and RET 21 values do offer some insight into the affinity of taxonomically uncertain specimens. KNM-WT 8556 22 has been attributed to Kenyanthropus platyops (Leakey et al., 2001) and Au. africanus (Brown et

al., 2001). AET and tooth size (based on dentine surface area) of the mandibular first molar are 1 2 within the range of Au. africanus and in the vicinity of Au. afarensis in (Fig. 6). The RET value of 21.79 for this specimen is within the range of Au. africanus and within one standard deviation of 3 4 the mean of Au. afarensis. Unfortunately, until enamel thickness values in a sample of mandibular 5 first molars of K. platyops are published it is unlikely that enamel thickness can be used as a 6 primary criterion for the taxonomic affinity of this specimen. Omo K7-1969-19 is similar in size, 7 AET, and RET to KNM-ER 1802 and DNH 67 and would be consistent with a classification to Homo 8 sp. (but see Leakey et al., 2012 for a discussion of the taxonomic affinity of KNM-ER 1802). L26-1g. 9 was attributed to Au. africanus by Howell and colleagues (1987) and has AET and RET values that align it with Au. afarensis and/or Homo sp. L28-30 (M2) and L28-31 (M3) are attributed to Homo 10 11 sp. by Suwa (1996). Both present AET and RET values similar to Au. boisei but over a relatively 12 small dentine crown, which would be consistent with the recent finding of high AET values in early 13 Homo by Smith and colleagues (2012a). Omo 75s-16 has relatively low AET and moderately high RET, suggesting that while it may be early Homo (Suwa pers communication) it is different from 14 other potential early Homo specimens such as L28-30. An important possibility for further 15 clarifying the taxonomic status of these specimens from West Turkana, Kenya and Omo, Ethiopia 16 will be an analysis of enamel thickness in the sample of ~3Ma teeth from Lomekwi, Kenya (Brown 17 et al., 2001; Leakey et al., 2001). 18

An increasing number of studies are examining dental tissue proportions in 3D (e.g., Kono, 2004; Olejniczak et al., 2008a/b; Kono and Suwa, 2008; Suwa et al., 2009; Benazzi et al., 2014; 21 Zanolli et al., 2014) and it is clear that taxonomic differences in molar crown shape (e.g., being 22 mesiodistally wider or narrower) will result in differences in AET and RET between 2D mesial plane

1 of section and 3D whole crown calculations (Olejniczak et al., 2008a). As there are tooth crown 2 shape differences between Pliocene hominins it will be necessary to determine whether the 3 patterns of statistically significant differences (and lack thereof in some pairwise comparisons) 4 found in this study, hold for 3D analyses of these specimens. However, it should be noted that 5 reconstructing missing enamel in 3D (due to fragmentation of the enamel cap around the cervix, 6 missing portions of the enamel cap, and the difficulty of reconstructing the original outer enamel 7 surface) can be extremely difficult and sample sizes may drop precipitously for many hominin taxa. 8 Additional analyses of crown and/or EDJ morphology may further elucidate the taxonomic affinity 9 of these specimens (Skinner et al., 2008a/b, 2009).

10

11 Dietary adaptation and enamel thickness

12 Enamel thickness has also been the basis for interpreting hominoid/hominin dietary 13 adaptations (e.g., Kay, 1981; Dumont, 1995; Shimizu, 2002; Vogel et al., 2008; Smith et al., 2012b). 14 As such, thick enamel is commonly associated with the consumption of hard (Constantino et al., 15 2009; 2011) and/or abrasive (Rabenold and Pearson, 2011) grass-based food material. Recent 16 results from isotopic and microwear research (see review in Sponheimer et al., 2013), have 17 created favorable conditions for testing these longstanding hypotheses. Isotopic studies of dental enamel (e.g., Wynn et al., 2013; Cerling et al., 2013) demonstrate an increase in C4 consumption 18 19 from Au. anamensis to Au. afarensis, and an additional increase in C4 consumption from Au. aethiopicus and Au. boisei. Our results suggest that this increase in C4 consumption by hominins is 20 21 associated with a concomitant increase in enamel thickness; however, whether increased enamel 22 thickness is related to abrasion resistance (Rabenold and Pearson, 2011) or fracture resistance

1 (Constantino et al., 2009, 2011; Strait et al., 2013) continues to be debated. Ungar and Sponheimer 2 (2011) analyzed microwear texture of Plio-Pleistocene hominins and found a decrease in complexity from Au. anamensis, to Au. afarensis, to Au. boisei that could correlate to an increase 3 4 in the need to shear tough foods (such as C4 grasses). Thus, for these taxa in East Africa there is an 5 associated change from greater complexity, lower C4 isotopic signatures and thinner enamel in Au. 6 anamensis, to reduced complexity, higher C4 isotopic signatures and thicker enamel in Au. boisei. 7 However, a similar trend across Pliocene hominins is complicated by the findings of Ungar and 8 Sponheimer (2011) of relatively high microwear complexity and relatively high C3 plant 9 consumption found by in Au. robustus (which also overlaps somewhat with the patterns in H. 10 erectus). Since Au. robustus has similarly high AET/RET values as Au. boisei, our results support the 11 hypothesis that thick enamel in Au. robustus (and possibly early Homo) cannot be attributed to 12 similar selective pressures (Ungar and Sponheimer, 2011; Sponheimer et al., 2013). Delezene and 13 colleagues (2013) analyzed microwear texture of Au. afarensis and Au. africanus premolars and 14 molars and found an increase in complexity (a proxy for hard-object feeding) from premolars to 15 molars within each species. Future analyses should explore whether the microwear pattern along 16 the tooth row (premolars to molars) is also matched in enamel thickness distribution. 17 In his analysis of enamel thickness distribution in mesial sections of hominoid upper molars, Schwartz (2000) noted a strong taxonomic signal and a relationship between differential 18 19 enamel distribution and diet. A number of studies have highlighted the influence of differential distribution of enamel across the crown and tooth function (Macho and Thackeray, 1992; Shimizu, 20 21 2002; Kono, 2004; Kono and Suwa, 2008). Our results indicate that the majority of fossil hominins,

as expected, do exhibit greater enamel thickness buccally than lingually. However, unexpectedly,

1 the thickest enamel is usually found occlusally. Thus, any variation in diet associated with changes 2 in isotopic chemistry, microwear, tooth size, does not seem to be associated with variation in 3 differential distribution of enamel buccolingually across the crown. Future studies should consider 4 more detailed mapping of the 3D distribution of enamel across the crown (e.g., Kono, 2004; 5 Olejniczak et al., 2008) and correlations with primary facet orientation and position (Kullmer et al., 6 2009). Ultimately, these studies can be expanded to using FE models to test the interaction of 7 enamel thickness, dentine crown morphology, and tooth wear on tooth function (e.g., Benazzi et 8 al., 2011, 2013).

9

10 Conclusion

11 In this study we report on 2D mesial plane of section enamel thickness values in Pliocene and early Pleistocene fossil hominins and non-human large ape mandibular molars. Our findings 12 13 confirm a general trend for increasing enamel thickness throughout the Pliocene Australopithecus 14 species culminating in Au. boisei. The majority of hominin and non-human large ape species exhibit 15 thickest enamel in the occlusal basin, less thick buccally and thinnest lingually. Gorilla exhibits the thinnest enamel relative to its tooth crown size and has thickest enamel buccally. Pan species are 16 17 unique in exhibiting the thinnest enamel occlusally. While there is considerable overlap in average and relative enamel thickness values among hominins of similar geochronological age, enamel 18 19 thickness retains the potential to be a useful taxonomic indicator for particular genera and time periods. 20

21

22 Acknowledgments

1 For access to specimens we thank the following institutions: National Museum of Ethiopia 2 and the ARCCH, Addis Ababa, Ethiopia; Emma Mbua and the National Museums of Kenya, Nairobi, Kenya; Bernard Zipfel and the University of Witwatersrand, Johannesburg, South Africa; Stephany 3 4 Potze and the Ditsong National Museum of Natural History, Pretoria, South Africa.; Museum for 5 Natural History (ZMB), Berlin, Germany; Ottmar Kullmer and the Senckenberg Research Institute (SMF), Frankfurt, Germany; Emmanuel Gillisen and the Royal Museum for Central Africa (MRAC), 6 7 Terverun, Belgium; Christophe Boesch and the Max Planck Institute for Evolutionary Anthropology 8 (MPI), Leipzig, Germany; Leipzig University Anatomical Collection (ULAC), Leipzig, Germany; the 9 "Francisc J. Rainer" Anthropology Institute, Bucharest, Romania. Tomographic scans of some specimens were produced through a collaborative project between the Department of Human 10 11 Evolution, Max Planck Institute for Evolutionary Anthropology and the Evolutionary Studies 12 Institute and Centre for Excellence in PaleoSciences, Johannesburg, South Africa. We also thank Bill 13 Kimbel for access to material from Hadar, Ethiopia and Colin Menter for access to material from Drimolen, South Africa. For scanning and technical assistance we thank Heiko Temming and Patrick 14 15 Schoenefeld. This research was supported by the Max Planck Society.

1	References
2	Benazzi, S., Kullmer, O., Grosse, I.R., Weber, G.W., 2011. Using occlusal wear information and finite
3	element analysis to investigate stress distributions in human molars. J. Anat. 219, 259-72.
4	
5	Benazzi, S., Panetta, D., Fornai, C., Toussaint, M., Gruppioni, G., Hublin, J-J. 2014. Guidelines for the
6	digital computation of 2D and 3D enamel thickness. Am. J. Phys. Anthropol. 153, 305-13.
7	
8	Benazzi, S., Nguyen, H.N., Schulz, D., Grosse, I.R., Gruppioni, G., Hublin, J-J., Kullmer, O., 2013. The
9	Evolutionary Paradox of Tooth Wear: Simply Destruction or Inevitable Adaptation? PLoS One 8,
10	e62263.
11	
12	Beynon, A.D., Wood, B.A., 1986. Variations in enamel thickness and structure in east African
13	hominids. Am. J. Phys. Anthropol. 70, 177-193.
14	
15	Boyer, D., 2008. Relief index of second mandibular molars is a correlate of diet among prosimian
16	primates and other euarchontan mammals. J. Hum. Evol. 55, 1118-1137.
17	
18	Brain, C.K., 1981. The Hunters of the Hunted? An Introduction to African Cave Taphonomy.
19	University of Chicago Press, Chicago.
20	
21	Brown, B., Brown, F.H., Walker, A., 2001. New hominids from the Lake Turkana Basin, Kenya. J.
22	Hum. Evol. 41, 29-44.
23	
24	Brunet, M., Guy, F., Pilbeam, D., Lieberman, D.E., Likius, A., Mackaye, H.T., Ponce de León, M.S.,
25	Zollikofer, C.P.E., Vignaud, P., 2005. New material of the earliest hominid from the upper Miocene
26	of Chad. Nature 434, 752-755.
27	
28	Brunet, M., Guy, F., Pilbeam, D., Mackaye, H.T., Likius, A., Ahounta, D., Beauvilain, A., Blondel, C.,

29 Bocherens, H., Boisserie, J.R., De Bonis, L., Coppens, Y., Dejax, J., Denys, C., Duringer, P.,

1	Eisenmann, V.R., Fanone, G., Fronty, P., Geraads, D., Lehmann, T., Lihoreau, F., Louchart, A.,
2	Mahamat, A., Merceron, G., Mouchelin, G., Otero, O., Campomanes, P.P., de Leon, M.P., Rage, J.C.,
3	Sapanet, M., Schuster, M., Sudre, J., Tassy, P., Valentin, X., Vignaud, P., Viriot, L., Zazzo, A.,
4	Zollikofer, C., 2002. A new hominid from the upper Miocene of Chad, central Africa. Nature 418,
5	145-151.
6	
7	Butler, P.M., 1956. The ontogeny of molar pattern. Bio. Rev. 31, 30-70.
8	
9	Cerling, T.E., Manthi, F.K., Mbua, E.N., Leakey, L.N., Leakey, M.G., Leakey, R.E., Brown, F.H., Grine,
10	F.E., Hart, J.A., Kaleme, P., Roche, H., Uno, K.T., Wood, B.A., 2013. Stable isotope-based diet
11	reconstructions of Turkana Basin hominins. PNAS 110, 10501-10506.
12	
13	Coffing, K., Feibel, C., Leakey, M., Walker, A., 1994. Four-million-year-old hominids from East Lake Turkana,
14	Kenya. Am. J. Phys. Anthropol. 93, 55-65.
16	Conroy G 1991 Enamel thickness in South African australonithecines: noninvasive evaluation by
17	computed tomography Palaeontol Afr 28 53-59
18	
19	Constantino. P.J., Lucas. P.W., Lee, J.JW., Lawn, B.R., 2009. The influence of fallback foods on
20	great ape tooth enamel. Am. J. Phys. Anthropol. 140, 653-660.
21	
22	Constantino, P.J., Lee, J.JW., Morris, D., Lucas, P.W., Hartstone-Rose, A., Lee, WK., Dominy, N.J.,
23	Cunningham, A., Wagner, M., Lawn, B.R., 2011. Adaptation to hard-object feeding in sea otters and
24	hominins. J. Hum. Evol. 61, 89-96.
25	
26	Dart, R.A., 1925. Australopithecus africanus: the man-ape from South Africa. Nature 115, 195-199.
27	
28	Dart, R.A., 1948. The adolescent mandible of Australopithecus Prometheus. Am. J. Phys.
29	Anthropol., 6:391-411.
30	

1	Delezene, L.K., Zolnierz, M.S., Teaford, M.F., Kimbel, W.H., Grine, F.E., Ungar, P.S., 2013. Premolar
2	microwear and tooth use in Australopithecus afarensis. J. Hum. Evol., 65, 282-293.
3	
4	De Ruiter, D., 2001. A Methodological Analysis of the Relative Abundance of Hominids and other
5	Macromammals from the Site of Swartkrans, South Africa. PhD Dissertation, University of the
6	Witwatersrand, South Africa.
7	
8	Dumont, E.R., 1995. Correlations between enamel thickness and dietary adaptations among extant
9	primates and chiropterans. J. Mammal., 76, 1127-1136.
10	
11	Gantt, D.G., 1986. Enamel thickness and ultrastructure in hominoids: with reference to form,
12	function and phylogeny. In: Swindler, D.R., Erwin, J. (Eds.), Comparative Primate Biology. Alan R.
13	Liss, New York, pp. 453-475.
14	
15	Grine, F.E., 1981. A new composite juveline specimen of Australopithecus africanus (Mammalia,
16	Primates) from Member 4, Sterkfontein Formation, Transvaal. Ann. S. Afr. Mus. 84, 169-201.
17	
18	Grine, F.E., 1991. Computed tomography and the measurement of enamel thickness in extant
19	hominoid primates: implications for its paleontological application. Palaeont. Afr. 28, 61-69.
20	
21	Grine, F.E., 2004. Description and preliminary analysis of new hominid craniodental fossils from
22	the Swartkrans Formation. In: Brain, C.K. (Eds.), Swartkrans: A Cave's Chronicle of Early Man.
23	Transvaal Museum, Pretoria, pp. 75-116.
24	
25	Grine, F.E., Martin, L., 1988. Enamel thickness and development in Australopithecus and
26	Paranthropus. In: Grine, F.E. (Ed.), Evolutionary History of the 'Robust' Australopithecines. Aldine
27	de Gruyter, New York, pp. 3-42.

28

1	Haile-Selassie, Y., Saylor, B.Z., Deino, A., Alene, M., Latimer, B., 2010. New hominid fossils from
2	Woranso-Mille (Central Afar, Ethiopia) and taxonomy of early Australopithecus. Am. J. Phys.
3	Anthropol. 141, 406-417.
4	
5	Howell, F.C., Haesaerts, P., de Heinzelin, J., 1987. Depositional environments, archaeological
6	occurrences and hominids from Members E and F of the Shungura Formation (Omo basin,
7	Ethiopia). J. Hum. Evol. 16, 665-700.
8	
9	Johanson, D.C., White, T.D., Coppens, Y., 1982. Dental remains from the Hadar Formation,
10	Ethiopia: 1974-1977 collections. Am. J. Phys. Anthropol. 57, 545-603.
11	
12	Kay, R.F., 1981. The nut-crackers – a new theory of the adaptations of the Ramapithecinae. Am. J.
13	Phys. Anthropol. 55, 141-151.
14	
15	Kono, R., 2004. Molar enamel thickness and distribution patterns in extant great apes and humans,
16	new insights based on a 3-dimensional whole crown perspective. Anthropol. Sci. 112, 121-146.
17	
18	Kono, R.T., Suwa, G., 2008. Enamel distribution patterns of extant human and hominoid molars:
19	occlusal versus lateral enamel thickness. Bull. Natl. Mus. Nat. Sci. D 34, 1-9.
20	
21	Kullmer, O., Benazzi, S., Fiorenza, L., Schulz, D., Bacso, S., Winzen, O., 2009. Technical note:
22	occlusal fingerprint analysis: quantification of tooth wear pattern. Am. J. Phys. Anthropol. 139,
23	600-605.
24	
25	Lacruz, R.S., Ramirez Rozzi, F.V., 2010. Molar crown development in Australopithecus afarensis. J.
26	Hum. Evol. 58, 201-206.
27	

1	Leakey, M., Tobias, P.V., Martyn, J.E., Leakey, R.E.F., 1969. An Acheulian industry with prepared
2	core technique and the discovery of a contemporary hominid mandible at Lake Baringo, Kenya.
3	Proc. Prehist. Soc. 3, 48-76.
4	
5	Leakey, M.G., Feibel, C.S., McDougall, I., Walker, A., 1995. New four-million-year-old hominid
6	species from Kanapoi and Allia Bay, Kenya. Nature 376, 565-571.
7	
8	Leakey, M.G., Spoor, F., Brown, F.H., Gathogo, P.N., Kiarie, C., Leakey, L.N., McDougall, I., 2001.
9	New hominin genus from eastern Africa shows diverse middle Pliocene lineages. Nature 410:433-
10	440.
11	
12	Leakey, M.G., Spoor, F., Dean, M.C., Feibel, C.S., Anton, S.C., Kiarie, C., Leakey, L.N., 2012. New
13	fossils from Koobi Fora in northern Kenya confirm taxonomic diversity in early Homo. Nature 488,
14	201-204.
15	
16	Leakey, R.E.F., Walker, A., 1988. New Australopithecus boisei specimens from east and west Lake
17	Turkana, Kenya. Am. J. Phys. Anthropol. 76, 1-24.
18	
19	Macho, G.A., Thackeray, J.F., 1992. Computed tomography and enamel thickness of maxillary
20	molars of Plio-Pliestocene hominids from Sterkfontein, Swartkrans, and Kromdraai (South Africa):
21	an exploratory study. Am. J. Phys. Anthropol. 89:133-143.
22	
23	Martin, L.B., 1985. Significance of enamel thickness in hominoid evolution. Nature 314, 260-263.
24	
25	Martin, L.B., Olejniczak, A.J., Maas, M.C., 2003. Enamel thickness and microstructure in pitheciin
26	primates, with comments on dietary adaptations of the middle Miocene hominoid Kenyapithecus.
27	J. Hum. Evol. 45, 351-367.
28	

1	Moggi-Cecchi, J., Grine, F.E., Tobias, P.T., 2006. Early hominid dental remains from Members 4 and
2	5 of the Sterkfontein Formation (1966–1996 excavations): catalogue, individual associations,
3	morphological descriptions and initial metric analysis. J. Hum. Evol. 50, 239–328.
4	
5	Moggi-Cecchi, J., Menter, C., Boccone, S., Keyser, A., 2010. Early hominin dental remains from the
6	Plio-Pleistocene site of Drimolen, South Africa. J. Hum. Evol. 58, 374–405.
7	
8	Molnar, S., Gantt, D.G., 1977. Functional implications of primate enamel thickness. Am. J. Phys.
9	Anthropol. 46, 447-454.
10	
11	Olejniczak AJ. 2006. Micro-computed tomography of primate molars. PhD Dissertation, Stony
12	Brook University.
13	
14	Olejniczak, A.J., Grine, F.E., 2005. High-resolution measurement of Neandertal tooth enamel
15	thickness by micro-focal computed-tomography. S. Afr. J. Sci. 101, 219-220.
16	
17	Olejniczak, A.J., Smith, T.M., Feeney, R.N.M., Macchiarelli, R., Mazurier, A., Bondioli, L., Rosas, A.,
18	Fortea, J., de la Rasilla, M., García-Tabernero, A., Radovcic, J., Skinner, M.M., Toussaint, M., Hublin,
19	JJ., 2008a. Dental tissue proportions and enamel thickness in Neandertal and modern human
20	molars. J. Hum. Evol. 55, 12-23.
21	
22	Olejniczak, A.J., Smith, T.M., Skinner, M.M., Grine, F.E., Feeney, R.N.M., Thackeray, F.J., Hublin, J-J.,
23	2008b. Three-dimensional molar enamel distribution and thickness in Australopithecus and
24	Paranthropus. Biol. Lett. 4, 406-410.
25	
26	Olejniczak, A.J., Smith, T.M., Wei, W., Potts, R., Ciochon, R., Kullmer, O., Schrenk, F., Hublin, J-J.,
27	2008c. Molar enamel thickness and dentine horn height in Gigantopithecus blacki. Am. J. Phys.
28	Anthropol. 135, 85-91.

1	Olejniczak, A.J., Tafforeau, P., Feeney, R.N.M., Martin, L.B., 2008d. Three-dimensional primate
2	molar enamel thickness. J. Hum. Evol. 54, 187-195.
3	
4	Rabenold, D., Pearson, O.M., 2011. Abrasive, silica phytoliths and the evolution of thick molar
5	enamel in primates, with implications for the diet of Paranthropus boisei. Plos ONE 6: e28379,
6	doi:10.1371/journal.pone.0028379
7	
8	Schwartz, G.T., 2000. Taxonomic and functional aspects of the patterning of enamel thickness
9	distribution in extant large-bodied hominoids. Am. J. Phys. Anthropol. 111, 221-244.
10	
11	Schwartz, G.T., Thackeray, J.F., Reid, C., van Reenan, J.F., 1998. Enamel thickness and the
12	topography of the enamel-dentine junction in South African Plio-Pleistocene hominids with special
13	reference to the Carabelli trait. J. Hum. Evol. 35, 523-542.
14	
15	Shimizu, D., 2002. Functional implications of enamel thickness in lower molars of red colobus
16	(Procolobus badius) and Japanese macaque (Macaca fuscata). J. Hum. Evol. 43: 605-620.
17	
18	Skinner, M.M., Gunz, P., Wood, B.A., Hublin, J-J., 2008a. Enamel-dentine junction (EDJ)
19	morphology distinguishes the lower molars of Australopithecus africanus and Paranthropus
20	<i>robustus</i> . J. Hum. Evol. 55, 979-988.
21	
22	Skinner, M.M., Wood, B., Boesch, C., Olejniczak, A.J., Rosas, A., Smith, T.M., Hublin, J-J., 2008b.
23	Dental trait expression at the enamel-dentine junction of lower molars in extant and fossil
24	hominoids. J. Hum. Evol. 54, 173-186.
25	
26	Skinner, M.M., Gunz, P., Wood, A., Boesch, C., Hublin, J.J., 2009. Discrimination of extant Pan
27	species and subspecies using the enamel-dentine junction morphology of lower molars. Am. J.
28	Phys. Anthropol. 140, 234-243.
1	Smith, T.M., Martin, L.B., Leakey, M.G., 2003. Enamel thickness, microstructure and development
----	--
2	in Afropithecus turkanensis. J. Hum. Evol. 44, 283-306.
3	
4	Smith, T.M., Olejniczak, A.J., Martin, L.B., Reid, D.J., 2005. Variation in hominoid molar enamel
5	thickness. J. Hum. Evol. 48, 575-592.
6	
7	Smith, T.M., Olejniczak, A.J., Reid, D.J., Ferrell, R.J., Hublin, JJ., 2006a. Modern human molar
8	enamel thickness and enamel-dentine junction shape. Arch. Oral Biol. 51, 974-995.
9	
10	Smith, T.M., Olejniczak, A.J., Tafforeau, P., Reid, D.J., Grine, F.E., Hublin, JJ., 2006b. Molar crown
11	thickness, volume, and development in South African Middle Stone Age humans. S. Afr. J. Sci. 102,
12	513-517.
13	
14	Smith, T.M., Harvati, K., Olejniczak, A.J., Reid, D.J., Hublin, JJ., Panagopoulou, E., 2009a. Brief
15	communication: dental development and enamel thickness in the Lakonis Neanderthal molar. Am.
16	J. Phys. Anthropol. 138, 112-118.
17	
18	Smith, T.M., Olejniczak, A.J., Kupczik, K., Lazzari, V., Vos, J., Kullmer, O., Schrenk, F., Hublin, JJ.,
19	Jacob, T., Tafforeau, P., 2009b. Taxonomic assessment of the Trinil molars using non-destructive
20	3D structural and developmental analysis. Paleoanthropol, 117-129.
21	
22	Smith, T.M., Olejniczak, A.J., Zermeno, J.P., Tafforeau, P., Skinner, M.M., Hoffmann, A., Radovčić, J.,
23	Toussaint, M., Kruszynski, R., Menter, C., Moggi-Cecchi, J., Glasmacher, U.A., Kullmer, O., Schrenk,
24	F., Stringer, C., Hublin, J-J., 2012a. Variation in enamel thickness within the genus Homo. J. Hum.
25	Evol. 62:395-411.
26	
27	Smith, T.M., Kupczik, K., Machanda, Z., Skinner, M.M., Zermeno JP., 2012b. Enamel thickness in
28	Bornean and Sumatran orangutan dentitions. Am. J. Phys. Anthropol. 147:417-426.
29	

1	Sponheimer, M., Alemseged, Z., Cerling, T.E., Grine, F.E., Kimbel, W.H., Leakey, M.G., Lee-Thorp,
2	J.A., Manthi, F.K., Reed, K.E., Wood, B.A., Wynn, J.G., 2013. Isotopic evidence of early hominin
3	diets. PNAS 110, 10513-10518.
4	
5	Strait, D.S., Constantino, P., Lucas, P.W., Richmond, B.G., Spencer, M.A., Dechow, P.C., Ross, C.F.,
6	Grosse, I.R., Wright, B.W., Wood, B.A., Weber, G.W., Wang, Q., Byron, C., Slice, D.E., Chalk, J.,
7	Smith, A.L., Smith, L.C., Wood, S., Berthaume, M., Benazzi, S., Dzialo, C., Tamvada, K., Ledogar, J.A.,
8	2013. Diet and dietary adaptations in early hominins: the hard food perspective. Am. J. Phys.
9	Anthropol. 151:339-355.
10	
11	Suwa, G., 1996. Serial allocations of isolated mandibular molars of unknown taxonomic affinities
12	from the Shungura and Unso Formations, Ethiopia, a combined method approach. Hum. Evol. 11,
13	269-282.
14	
15	Suwa, G., Kono, R.T., Simpson, S.W., Asfaw, B., Lovejoy, C.O., White, T.D., 2009. Paleobiological
16	implications of the Ardipithecus ramidus dentition. Science 326, 70-99.
17	
18	Tafforeau, P., 2004. Phylogenetic and functional aspects of tooth enamel microstructure and
19	three-dimensional structure of modern and fossil primate molars. Ph.D. Dissertation, Montpellier
20	University II.
21	
22	Thackeray, J.F., de Ruiter, D.J., Berger, L.R., Van Der Merwe, N.J., 2001. Hominid fossils from
23	Kromdraai: a revised list of specimens discovered since 1938. Ann. Transv. Mus. 38, 43-56.
24	
25	Ungar, P.S., Sponheimer, M., 2011. The diets of early hominins. Science 334, 190-193.
26	
27	Vogel, E.R., van Woerden, J.T., Lucas, P.W., Atmoko, S.S.U., van Schaik, C.P., Dominy, N.J., 2008.
28	Functional ecology and evolution of hominoid molar enamel thickness: Pan troglodytes
29	schweinfurthii and Pongo pygmaeus wurmbii. J. Hum. Evol. 55, 60-74.

1	
2	Ward, C.V., Leakey, M.G., Walker, A., 2001. Morphology of Australopithecus anamensis from
3	Kanapoi and Allia Bay, Kenya. J. Hum. Evol. 41, 255-368.
4	
5	White, T.D., Suwa, G., Asfaw, B., 1994. Australopithecus ramidus, a new species of early hominid
6	from Aramis, Ethiopia. Nature 371, 306-312.
7	
8	White, T.D., Suwa, G., Simpson, S., Asfaw, B., 2000. Jaws and teeth of Australopithecus afarensis
9	from Maka, Middle Awash, Ethiopia. Am. J. Phys. Anthropol. 111, 45-68.
10	
11	White, T.D., WoldeGabriel, G., Asfaw, B., Ambrose, S., Beyene, Y., Bernor, R.L., Boisserie, JR.,
12	Currie, B., Gilbert, H., Haile-Selassie, Y., Hart, W.K., Hlusko, L.J., Clark Howell, F., Kono, R.T.,
13	Lehmann, T., Louchart, A., Lovejoy, C.O., Renne, P.R., Saegusa, H., Vrba, E.S., Wesselman, H., Suwa,
14	G., 2006. Asa Issie, Aramis and the origin of Australopithecus. Nature 440, 883-889.
15	
16	Wood, B.A., 1991. Koobi Fora Research Project, Vol. 4: Hominid Cranial Remains. Clarendon Press,
17	Oxford.
18	
19	Wynn, J.G., Sponheimer, M., Kimbel, W.H., Alemseged, Z., Reed, K., Bedaso, Z.K., Wilson, J.N.,
20	2013. Diet of Australopithecus afarensis from the Pliocene Hadar Formation, Ethiopia. PNAS 110,
21	10495-10500.
22	
23	Zanolli, C., Bondioli, L., Coppa, A., Dean, M.C., Bayle, P., Candilio, F., Capuani, S., Dreossi, D., Fiore,
24	I., Frayer, D.W., Libsekal, Y., Mancini, L., Rook, L., Takle, T.M., Tuniz, C., Macchiarelli, R., 2014. The
25	late Early Pleistocene human dental remains from Uadi Aalad and Mulhuli-Amo (Buia), Eritrean
26	Danakil: macromorphology and microstructure. J. Hum. Evol. 74, 96-113.

2	Figure 1. Illustration of protocol used to collect enamel thickness data. Top left shows a surface
3	model of a mandibular first molar with a red line indicating the location of the 2D plane of section.
4	Top right shows the 2D plane of section for this specimen upon which measurements were
5	collected. Bottom right shows the surface area of the enamel cap (yellow) and the dentine (blue).
6	Bottom left shows the bi-cervical diameter measurement (red line), the enamel-dentine junction
7	length measurement (white line), and the black lines delimit the lingual, occlusal and buccal
8	regions used to measure the distribution of enamel across the crown.
9	
10	Figure 2. Selection of mandibular second molars of the study sample. The red line indicates slice
11	position and the segmented enamel and dentine image is overlaid on original slice to show areas
12	corrected for missing enamel. White scale bar = 5mm.
13	
14	Figure 3. Box plots of average enamel thickness values for each taxon.
15	
16	Figure 4. Box plots of relative enamel thickness values for each taxon.
17	
18	Figure 5. Patterns of regional (buccal, occlusal, lingual) average enamel thickness for the combined
19	molar sample of each taxon. Hominins and Pongo tend to exhibit thickest enamel in the occlusal
20	basin, while Gorilla and Pan exhibit thickest enamel on the lateral tooth crown.
21	

1	Figure 6. Plot of AET (log) against dentine surface area (log) for the first molar. Specimens of
2	uncertain taxonomic affinity are marked with stars.
3	
4	Figure 7. Plot of AET (log) against dentine surface area (log) for the second molar. Specimens of
5	uncertain taxonomic affinity are marked with stars.
6	
7	Figure 8. Plot of AET (log) against dentine surface area (log) for the third molar. Specimens of
8	uncertain taxonomic affinity are marked with stars.
9	
10	
11	Supplementary figure captions
12	
13	Supplementary Figure 1. Example of estimation of worn enamel in a 2D mesial section. The
14	unworn crown of STW 308 (top left) was artificially worn (top right) to remove a substantial
15	proportion of enamel (much greater than in most of the study specimens). The original
16	segmentation of enamel and dentine tissue (bottom left) can be compared to the blind estimation
17	of the original enamel (bottom right). Note that there is only a 2.2% difference in the calculated
18	average enamel thickness between the original and reconstructed specimen.
19	
20	Supplementary Figure 1. Au. anamensis M1 sample - mesial planes of section and their location for
21	each specimen. White scale bar = 5mm.
22	

1	Supplementary Figure 3. Au. anamensis M2 and M3 sample - mesial planes of section and their
2	location for each specimen. White scale bar = 5mm.
3	
4	Supplementary Figure 4. Au. afarensis M1 – M3 sample - mesial planes of section and their
5	location for each specimen. White scale bar = 5mm.
6	
7	Supplementary Figure 5. Au. africanus M1 sample - mesial planes of section and their location for
8	each specimen. White scale bar = 5mm.
9	
10	Supplementary Figure 6. Au. africanus M2 sample - mesial planes of section and their location for
11	each specimen. White scale bar = 5mm.
12	
13	Supplementary Figure 7. Au. africanus M3 sample - mesial planes of section and their location for
14	each specimen. White scale bar = 5mm.
15	
16	Supplementary Figure 8. Au. boisei M1 – M3 sample - mesial planes of section and their location
17	for each specimen. White scale bar = 5mm.
18	
19	Supplementary Figure 9. Au. robustus M1 sample - mesial planes of section and their location for
20	each specimen. White scale bar = 5mm.
21	
22	Supplementary Figure 10. Au. robustus M2 sample - mesial planes of section and their location for
23	each specimen. White scale bar = 5mm.
24	

1	Supplementary Figure 11. Au. robustus M3 sample - mesial planes of section and their location for
2	each specimen. White scale bar = 5mm.
3	
4	Supplementary Figure 12. <i>H. erectus</i> M1 – M3 sample - mesial planes of section and their location
5	for each specimen. White scale bar = 5mm.
6	
7	Supplementary Figure 13. <i>Homo sp.</i> M1 – M3 sample - mesial planes of section and their location
8	for each specimen. White scale bar = 5mm.
9	
10	Supplementary Figure 14. Modern <i>Homo sapiens</i> selection of the M1 – M3 sample - mesial planes
11	of section and their location for each specimen. White scale bar = 5mm.
12	
13	Supplementary Figure 15. Pan troglodytes (MPI sample) M1 – M3 - mesial planes of section and
14	their location for each specimen. White scale bar = 5mm.
15	
16	Supplementary Figure 16. Pan troglodytes (ZMB sample) M1 – M3 - mesial planes of section and
17	their location for each specimen. White scale bar = 5mm.
18	
19	Supplementary Figure 17. Pan paniscus M1 – M3 sample - mesial planes of section and their
20	location for each specimen. White scale bar = 5mm.
21	
22	Supplementary Figure 18. Gorilla M1 – M3 sample - mesial planes of section and their location for
23	each specimen. White scale bar = 5mm.
24	

- 1 Supplementary Figure 19. *Pongo* selection of the M1 M3 sample mesial planes of section and
- 2 their location for each specimen. White scale bar = 5mm.

1	Table 1. Composition of the study sample ¹
---	---

Taxon	M1	M2	M3	Total
Pongo	9	8	3	20
Gorilla	2	5	6	13
Pan paniscus	3	5	0	8
Pan troglodytes	6	7	3	16
Australopithecus anamensis	6	4	3	13
Australopithecus afarensis	2	4	2	8
Australopithecus africanus	9	13	12	34
Australopithecus aethiopicus	0	2	1	3
Australopithecus boisei	0	4	3	7
Australopithecus robustus	6	8	10	24
<i>Homo</i> sp. indet.	2	2	0	4
Homo erectus	1	3	2	6
Homo sapiens	8	15	7	30
Total	54	80	52	186

1. Not including specimens of uncertain taxonomic affinity listed in Table 4.

Taxon	Tooth	N	Enamel Area (mm ²)	SD	Dentine Area (mm²)	SD	EDJ Length (mm)	SD	BCD (mm)	SD	AET (mm)	SD	RET	SD
Pongo	M1	9	19.64	5.40	42.21	9.00	20.21	1.78	9.56	1.12	0.96	0.18	14.79	1.70
Gorilla	M1	2	23.94	1.12	76.27	4.01	27.96	1.34	12.77	0.16	0.86	0.08	9.84	1.19
Pan	M1	9	12.89	2.00	29.79	3.64	18.01	0.94	7.67	0.40	0.71	0.09	13.08	1.33
A. anamensis	M1	6	19.79	2.52	28.70	5.37	17.72	1.42	9.93	0.61	1.12	0.13	21.21	3.88
A. afarensis	M1	2	25.25	1.35	43.01	5.19	19.80	1.50	11.37	0.23	1.28	0.03	19.53	1.62
A. africanus	M1	9	29.10	3.52	43.41	9.21	20.81	2.01	11.04	0.92	1.40	0.16	21.67	3.97
A. aethiopicus	M1	0	-	-	-	-	-	-	-	-	-	-	-	-
A. boisei	M1	0	-	-	-	-	-	-	-	-	-	-	-	-
A. robustus	M1	6	39.53	4.59	47.46	10.15	21.56	2.03	11.64	0.86	1.84	0.18	27.10	4.12
H. sp. indet.	M1	2	33.35	0.95	34.33	2.83	19.24	1.21	11.66	1.87	1.73	0.07	29.61	2.35
H. erectus	M1	1	23.06	-	36.28	-	19.55	-	10.28	-	1.18	-	19.59	-
H. sapiens	M1	8	18.50	2.84	33.65	3.60	19.29	1.20	8.66	0.53	0.96	0.10	16.47	1.14
Pongo	M2	8	23.65	2.31	46.52	4.65	20.54	1.00	10.56	1.01	1.16	0.14	17.05	2.58
Gorilla	M2	5	30.10	4.55	82.84	13.54	29.91	1.74	14.00	0.91	1.00	0.12	11.10	1.33
Pan	M2	12	13.73	2.11	30.04	4.52	18.21	0.85	7.93	0.88	0.75	0.11	13.82	2.02
A. anamensis	M2	4	24.61	1.94	46.27	4.52	20.52	0.70	12.50	0.43	1.20	0.10	17.69	1.75
A. afarensis	M2	4	27.41	4.50	36.23	7.23	18.08	1.48	11.74	0.70	1.51	0.20	25.33	3.62
A. africanus	M2	13	36.66	5.12	51.84	9.92	22.13	1.82	13.08	1.29	1.66	0.21	23.27	3.30
A. aethiopicus	M2	2	50.52	3.15	66.77	6.30	23.73	1.46	13.86	1.31	2.13	0.00	26.09	1.21
A. boisei	M2	4	46.18	15.74	50.93	14.87	21.63	2.89	13.63	2.36	2.11	0.50	29.58	5.17
A. robustus	M2	8	44.47	5.46	49.14	9.57	20.71	1.79	12.76	1.21	2.15	0.21	31.09	4.54
H. sp. indet.	M2	3	31.66	6.12	37.95	6.47	20.09	1.49	11.76	1.34	1.57	0.24	25.54	2.74
H. erectus	M2	3	27.39	3.95	35.80	3.46	18.90	0.76	11.67	0.08	1.45	0.15	24.15	1.34
H. sapiens	M2	15	20.73	2.56	33.62	5.67	18.44	1.49	9.11	0.76	1.12	0.10	19.56	2.27
Pongo	M3	3	20.08	2.06	34.30	10.55	17.99	2.42	9.17	0.95	1.13	0.21	20.05	5.94
Gorilla	M3	6	26.55	3.43	65.66	6.76	26.49	1.24	13.35	0.98	1.01	0.14	12.43	1.73
Pan	M3	3	14.69	1.93	31.79	0.88	19.22	0.65	7.91	0.66	0.76	0.08	13.53	1.27
A. anamensis	M3	3	26.30	3.91	39.37	3.91	19.22	0.90	12.14	1.01	1.37	0.17	21.79	2.32
A. afarensis	M3	2	28.78	0.52	39.35	0.78	18.65	0.16	11.83	1.70	1.54	0.01	24.61	0.47
A. africanus	M3	11	41.08	3.99	55.71	9.53	22.93	1.87	13.63	1.32	1.79	0.16	24.33	3.32
A. aethiopicus	M3	1	57.47	-	73.35	-	25.79	-	15.49	-	2.23	-	26.02	-
A. boisei	M3	3	57.32	7.13	59.15	14.39	22.37	1.87	14.16	1.10	2.56	0.18	33.67	3.74
A. robustus	M3	10	43.43	3.34	52.57	5.25	21.63	0.93	13.21	1.15	2.01	0.17	27.86	3.10
H. sp. indet.	M3	0	-	-	-	-	-	-	-	-	-	-	-	-
H. erectus	M3	2	25.24	1.94	35.47	4.22	18.54	0.62	10.75	0.91	1.37	0.15	23.05	3.90
H. sapiens	M3	7	21.60	4.55	31.27	7.59	17.49	2.14	8.65	1.43	1.23	0.19	22.36	3.82

Table 2. Mean and standard deviation of selected measured variables for each taxon and tooth position.

Notes

Accession	Current taxon	Citation	Tooth	Basis ¹	Citation	Enamel Area (mm ²)	Dentine Area (mm²)	EDJ Length (mm)	BCD ² (mm)	AET ³	RET ⁴
KNM-WT 8556	A. afarensis	Α	LM1	1	Α	27.74	41.38	19.79	11.61	1.40	21.79
Omo K7-1969-19	<i>Homo</i> sp.	D	LM1	3	В	38.60	35.89	21.02	11.58	1.84	30.65
L26-1g	A. aff. africanus	E	RM1	3	В	34.29	50.82	21.78	11.77	1.57	22.09
L28-31	Homo sp.	В	RM2	3	В	44.45	36.41	19.29	9.77	2.30	38.18
Omo 75s-1969-16	Homo sp.	С	RM3	3	С	26.24	30.46	18.06	10.77	1.45	26.32
L28-30	Homo sp.	В	RM3	3	В	41.02	36.11	17.99	10.49	2.28	37.94
L795-1	Hominin	В	RM2	3	В	51.70	63.66	24.21	14.82	2.14	26.76

Table 3. Enamel thickness values for specimens with uncertain taxonomic affinity

Notes:

1. Basis – 1 = molar in jaw or from associated dentition, 2 = molar position based on morphology and possible association with other teeth, 3 = molar position is best estimation based on morphology; 2. BCD = bi-cervical diameter; 3. AET = average enamel thickness; 4. RET = relative enamel thickness

Citations: A – Brown et al., 2001; B – Suwa, 1996; C – Suwa pers. comm; D – Alemseged pers. comm; E – Howell et al. 1987

Table 4. Molar enamel thickness comparison (AET bottom/RET top). Kruskal-Wallis with posthoc pairwise comparisons. Light shading indicates comparisons between hominins and extant non-human apes. Blank cells indicate non-significant results.

Taxon	Pongo	Gorilla	Pan	Au. anam.	Au. afar.	Au. afric.	Au. boisei	Au. rob.	H. sp. indet.	H. erectus	H. sapiens	
First molars												
Pongo				0.013	0.003		-	< 0.001	0.004			
Gorilla				0.005	0.031	0.003	-	< 0.001				
Pan				0.001	0.031	< 0.001	-	< 0.001	< 0.001			
Au. anamensis			0.003				-					
Au. afarensis		0.012					-					
Au. africanus	0.006	0.035	< 0.001				-				0.047	
Au. boisei	-	-	-	-	-	-						
Au. robustus	< 0.001	0.005	< 0.001	0.025							0.005	
H. sp. indet.	0.013	0.022	< 0.001								0.021	
H. erectus												
H. sapiens			0.044		0.012			0.001	0.018			
Second molars												
Pongo					0.024	0.018	0.003	< 0.001	0.026			
Gorilla					< 0.001	< 0.001	< 0.001	<0.001	0.001	0.007	0.005	
Pan	0.015				0.001	< 0.001	< 0.001	<0.001	0.001	0.015	0.007	
Au. anamensis			0.032				0.014	0.002				
Au. afarensis		0.029	0.001									
Au. africanus	0.011	0.001	< 0.001					0.040				
Au. boisei	0.011	0.001	< 0.001	0.043							0.017	
Au. robustus	< 0.001	< 0.001	< 0.001	0.007							< 0.001	
H. sp. indet.		0.030	0.001									
H. erectus			0.013									
H. sapiens			0.016			< 0.001	0.002	< 0.001				
Third molars											-	
Pongo							0.013	0.036	-			
Gorilla					0.045	0.001	< 0.001	< 0.001	-		0.026	
Pan						0.020	< 0.001	0.001	-			
Au. anamensis							0.020		-			
Au. afarensis									-			
Au. africanus	0.029	0.001	0.001						-			
Au. boisei	0.002	< 0.001	< 0.001	0.015					-		0.011	
Au. robustus	0.003	< 0.001	< 0.001	0.030					-		0.025	
H. sp. indet.	-	-	-	-	-	-	-	-				
H. erectus							0.027					
H. sapiens						0.011	0.001	<0.001				

Note: *Australopithecus aethiopicus* specimens not included in statistical tests due to small sample size. A hyphen indicates no molars of that position for that taxon.

Taxon	Ν	Buccal	SD	Occlusal	SD	Lingual	SD
Pongo	20	1.02	0.17	1.15	0.27	1.02	0.13
Gorilla	13	1.05	0.12	0.99	0.20	0.88	0.10
Pan	24	0.79	0.11	0.67	0.09	0.78	0.13
A. anamensis	13	1.23	0.23	1.27	0.20	1.11	0.15
A. afarensis	8	1.43	0.25	1.59	0.26	1.36	0.13
A. africanus	34	1.62	0.21	1.77	0.31	1.50	0.21
A. boisei	7	2.15	0.48	2.57	0.52	2.14	0.38
A. robustus	24	1.93	0.26	2.27	0.27	1.80	0.21
H. sp. indet.	4	1.76	0.23	1.82	0.24	1.58	0.17
H. erectus	6	1.32	0.06	1.49	0.22	1.28	0.20
H. sapiens	30	1.09	0.17	1.18	0.22	1.05	0.12

Table 5. Regional average enamel thickness measurements for the combined molar sample.

Note: *Australopithecus aethiopicus* not included in regional AET analysis due to small sample size (n = 3).

	_		1		Enamel	Dentine	EDJ	BCD ²	3	4
Accession	Taxon	Tooth	Basis	Citation	Area	Area	Length	(mm)	AET	RET
					(mm ⁻)	(mm ⁻)	(mm)	· · /		
AL145-35	A. afarensis	LM1	1	A	26.17	46.68	20.85	11.53	1.26	18.37
AL333w-1a	A. afarensis	LM1	1	A	24.28	39.34	18.78	11.21	1.29	20.61
AL128-23	A. afarensis	RM2	1	A	20.66	29.68	16.37	11.18	1.26	23.17
AL145-35	A. afarensis	LM2	1	A	29.81	46.48	20.12	12.09	1.48	21.73
AL241-14	A. afarensis	LM2	3	A	29.35	33.35	17.87	12.56	1.64	28.44
AL333w-1a	A. afarensis	LM2	1	А	27.65	35.39	17.93	11.14	1.54	25.92
AL400-1a	A. afarensis	RM3	1	А	29.14	38.80	18.85	10.63	1.55	24.82
AL333w-32	A. afarensis	RM3	2	А	28.37	39.90	18.65	13.03	1.52	24.08
STW421B	A. africanus	LM1	2	В	30.70	58.83	23.72	12.34	1.29	16.87
STS9	A. africanus	RM1	3	Ν	34.82	40.34	19.56	11.66	1.78	28.03
Taung1	A. africanus	LM1	1	С	28.32	42.90	21.94	11.71	1.29	19.71
STW327	A. africanus	LM1	1	В	29.63	49.77	20.80	11.55	1.42	20.19
STW151	A. africanus	RM1	1	D	28.87	32.41	19.20	9.80	1.50	26.41
STW106	A. africanus	RM1	1	В	20.35	36.00	18.67	10.56	1.09	18.17
STW123a	A. africanus	RM1	1	В	31.75	37.45	18.96	10.86	1.67	27.36
STW309a	A. africanus	RM1	1	В	32.56	53.09	23.77	12.04	1.37	18.80
STW246	A. africanus	LM1	2	В	29.62	46.15	21.22	10.77	1.40	20.55
STS24	A. africanus	RM1	1	Е	29.11	34.07	18.99	9.81	1.53	26.26
STW3	A. africanus	LM2	2	В	40.98	46.55	21.43	12.89	1.91	28.03
STW412B	A. africanus	LM2	2	В	25.31	42.86	20.52	11.83	1.23	18.84
STW327	A. africanus	LM2	1	В	43.22	61.57	23.12	13.96	1.87	23.83
MLD2	A. africanus	RM2	1	Z	36.84	52.71	22.70	14.43	1.62	22.35
STW498c	A. africanus	LM2	1	В	37.64	76.41	26.40	14.69	1.43	16.31
STW404	A. africanus	RM2	1	В	33.00	44.89	19.60	11.78	1.68	25.13
STW61	A. africanus	RM2	2	В	35.12	43.53	21.23	13.31	1.65	25.08
STW555	A. africanus	LM2	2	В	31.33	50.19	23.53	11.18	1.33	18.80
STW109	A. africanus	RM2	1	В	38.61	51.70	22.41	14.89	1.72	23.96
STW537(269)	A. africanus	RM2	1	В	41.75	53.19	23.24	14.03	1.80	24.63
STW308	A. africanus	RM2	1	B	40.90	50.42	22.11	13.28	1.85	26.05
STW133	A. africanus	IM2	2	B	42.87	61.50	22.94	13.20	1.87	23.83
STW213	A. africanus	LM2	1	B	33.92	44.63	21.39	11.78	1.59	23.74
STW529(532)	A africanus	IM3	1	B	42 55	47 54	21.65	12 75	1 98	28 73
STW 560B	A africanus	IM3	1	B	38 78	59 64	23 10	14 14	1.50	21 74
STW 9002	A africanus	IM3	1	B	46 64	69.26	26 79	14 75	1 74	20.92
STW384	A africanus	RM3	1	B	42 95	72 46	24 98	15 41	1 72	20.20
STW14	A africanus	RM3	1	B	43 16	56 74	22 45	12.84	1 92	25 52
STW404	A africanus	RM3	1	B	39.40	51 70	21 27	11 97	1.85	25.52
STW109	Δ africanus	RM3	1	B	42 98	51.03	22.27	14.83	1.00	27.09
STW105	Δ africanus	RM3	2	B	31 33	48 53	22.21	12 65	1.34	20.22
STW526	Δ africanus	IM3	1	B	37 73	40.55	20.44	11 59	1.41	28.66
STW280(278)	Δ africanus	RM3	1	B	41 65	60.69	25.1	15 28	1.65	20.00
ST\N/527	A. africanus	RM2	1	B	41.05	62.05	23.1	11.17	1.00	21.50
VNIM ED 20422	A. anamonsis		2	<u>с</u>	21 24	21.12	10.27	10.44	1.00	22.74
KNM-ER 20422	A. unumensis		3	G	17 60	24.80	15.57	0 20	1.10	20.75
KNM-ER 35232	A. anamensis		3	G	2/ 21	24.00	18 3/	10 32	1 32	22.70
KNIM-KD 21772	A. Unumensis		2	G	18 26	20.00	15 70	10.3Z	1 16	24.92 25 71
KNM-KP 217121	Δ anamensis	RM1	1	G	17 90	23.40	17 56	9.20	1.10	17 72
KNM-KP 34725R	A anamensis	RM1	- 1	G	19.06	34 66	19 55	10.60	0.97	16 56
KNM-FR 35733	A anamensis	IM2	3	G	22 39	43 76	20.25	11 93	1 11	16 71
KNM-KP 20286	Δ anamensis	1 M2	1	G	24 13	A1 52	21 02	12 72	1 15	17.80
1. IVI IN 25200	. anamensis		-	9	24.13	41.02	21.00	12.72	1.10	17.00

Appendix A. Measured variables for study sample

KNM-KP 34725T	A. anamensis	LM2	1	G	26.99	48.16	20.30	12.43	1.33	19.16
KNM-KP 30500D	A. anamensis	RM2	1	G	24.53	51.64	18.77	12.91	1.31	18.19
KNM-ER 20428	A. anamensis	LM3	3	F	30.13	43.76	20.11	13.00	1.50	22.65
KNM-KP 29281	A. anamensis	LM3	1	G	22.30	38.09	19.15	11.03	1.16	18.87
KNM-KP 29286	A. anamensis	LM3	1	G	25.51	36.25	19.01	12.40	1.34	22.29
ZMB 31435	Gorilla sp.	LM1	1	Н	24.32	73.43	29.10	12.66	0.84	9.75
ZMB 83546	Gorilla sp.	LM1	1	Н	22.91	79.11	29.04	12.88	0.79	8.87
ZMB 30940	Gorilla sp.	RM2	1	Н	26.49	90.23	31.74	13.78	0.83	8.79
7MB 31435	Gorilla sp.	IM2	1	н	32.29	75.72	30.30	12.92	1.07	12.25
ZMB 83546	Gorilla sp.	IM2	1	н	36.65	103.03	29.60	15.41	1.24	12.20
ZMB 83581	Gorilla sp.	RM2	1	н	24.89	70.27	28.62	14.18	0.87	10.37
SMF 45713	Gorilla sp.	RM2	1	Н	29.28	74.94	28.98	13.72	1.01	11.67
ZMB 30940	Gorilla sp.	RM3	1	н	25.49	75.36	29.18	13.63	0.87	10.06
ZMB 30941	Gorilla sp.	RM3	1	Н	25.14	69.32	26.92	14.37	0.93	11.22
ZMB 31277	Gorilla sp.	LM3	1	Н	31.16	68.10	26.16	14.13	1.19	14.43
ZMB 31435	Gorilla sp.	LM3	1	Н	30.19	63.27	26.72	11.69	1.13	14.20
ZMB 31626	Gorilla sp.	RM3	1	н	24.95	62.03	25.31	12.82	0.99	12.52
ZMB 83581	Gorilla sp.	RM3	1	н	21.98	55.86	25.22	13.44	0.87	11.66
R123	Homo saniens	IM1	1	M	15 78	32.18	19.27	9.12	0.82	14 44
R1101 1498	Homo saniens	LM1	1	M	23.61	43.09	20.74	8 13	1 14	17 34
R1140 899	Homo saniens	RM1	1	M	16 14	32.02	18.01	8 73	0.90	15.83
R1989 1387	Homo saniens	IM1	1	M	25 50	/1 53	21.00	9.75	1 21	18.84
R2602 1673	Homo saniens		1	M	20.42	38 39	21.00	2.31 2.91	0.97	15.58
Relgian93a	Homo saniens		2	1 1	20.42	38 /1	21.15	8 5 8	1 1/	18.38
Beligian120a	Homo saniens		2	1	18 11	21 22	10 0/	7 57	0.05	16.00
Belgian A31	Homo saniens		2	1	10.11	36.52	10.71	8.64	0.95	16.39
102 1151	Homo saniens		2 1	I M	19.50	31 30	19.71	8.04	0.99	15.05
R123	Homo saniens		1	M	15.86	28.67	18.70	8 90	0.85	16.01
R125	Homo saniens		1	M	10.00	20.07	17 28	0.50	1 11	20.27
R258 144	Homo saniens	RM2	1	M	20.20	29.92	10.20	836	1.11	17 90
R600 1372	Homo saniens		1	N/	20.35	34.00	19.29	8.30 8.88	1.00	17.50
D012 750	Homo sapiens		1	N/	21.45	22.01	10.00	0.00	1.07	20.80
D1101 1/09	Homo sanians		1	IVI NA	21.49	22.01	10.10	0.22	0.97	20.89
R1101 1450 D1224	Homo sapiens		1	IVI NA	10.42	20.90	10.75	9.52	1.00	10.24
R1234 R124E 1006	Homo sapiens		1	IVI NA	19.77	52.74 52.15	10.11	0.94	1.09	19.08
P1620 1196	Homo sanians		1	IVI NA	10.25	23.12	19.05	7 70	1.11	19.02
D2/22 1156	Homo sanians		1	IVI NA	19.55	33.11 41 57	20.76	0.57	1.04	17.64
	Homo sapiens		1	IVI	23.01	41.37	10.00	3.37 10.27	1.14	10.60
	Homo sapiens		1	L .	22.55	22.52	19.09	10.57	1.10	19.00
Bolgian/1a	Homo sanians		1 2	L	20.31	26.80	16.22	0.10	1.15	24 70
Belgian41a Bolgian100f	Homo sanians		2	1	19 21	20.80	17.00	9.29	1.20	24.79
Delgianituui	Homo sapiens		2 1	1	10.21	29.84	10 51	9.39	1.04	19.11
	Homo sapiens		1	IVI NA	10.04	20.20	16.04	0.20	1.02	10.44
	Homo sapiens		1	IVI NA	17.19	23.07	10.40	0.41 9.60	1.05	21.40
R003 1183	Homo sapiens		1	IVI NA	10.00	35.00	16.39	0.05	1.01	22.09
R1580 2425	Homo sapiens		1		18.91	26.07	10.12	6.15	1.17	22.98
	Homo sapiens		1	IVI	21.19	20.03	15.10	0.95	1.40	27.37 10 EG
ULAC 799-27	Homo sapiens		т 2	L 1	20.50	40.50	10 20	0 50	1.55	19.50
			:	J	27.70	32.30	16.20	7.00	1.52	12.12
MRAC 29026	Pan paniscus	RIVII	1	ĸ	9.39	22.09	10.47	7.00	0.57	12.13
MRAC 84036W111	Pan paniscus		1	ĸ	12.39	28.36	18.55	7.54	0.67	12.54
MRAC 29030	Pan paniscus	RIVI1	1	ĸ	10.28	27.72	17.68	7.49	0.58	11.05
MRAC 22908	Pan paniscus	LIVI2	1	K	10.04	25.83	17.39	6.60	0.58	11.36
MIKAC 29030	Pan paniscus		1	ĸ	13.64	27.27	18.12	7.44	0.75	14.42
MRAC 29055	Pan paniscus	LM2	1	K	13.39	28.83	18.45	8.05	0.73	13.51
MRAC 84036M11	Pan paniscus	LIVI2	1	K	14.45	27.93	18.50	7.61	0.78	14./8
MIRAC 84036M03	Pan paniscus	LIVI2	1	К	13.05	21.14	16.35	1.//	0.80	17.36
ZIVIB 0A16207	Pan troglodytes	LM1	1	H	12.56	30.80	18.29	7.38	0.69	12.37
ZMB 20811	Pan troglodytes	RM1	1	н	13.41	33.11	18.05	7.92	0.74	12.92

	ZMB 32356	Pan troglodytes	RM1	1	н	14.37	28.93	18.36	7.41	0.78	14.55
	ZMB 35526	Pan troalodytes	RM1	1	н	13.73	33.34	19.56	8.15	0.70	12.15
	ZMB 83623	Pan troalodytes	LM1	1	н	15.17	30.10	17.86	8.08	0.85	15.48
	7MB 30847	Pan troalodytes	IM2	1	н	14 76	32.08	19.00	7 54	0.78	13 72
	7MB 31279	Pan troalodytes	RM2	1	н	11 93	29.60	17 58	7.63	0.68	12.48
	ZMB 728//	Pan troalodytes	RM2	1	н	13.60	32.16	18 76	9.03	0.00	12.40
	ZIVID 72044	Pan troglodytes		1	н Ц	10.74	32.10	10.70	5.05 6.6E	0.72	10.57
		Pull troglodytes		1	п	10.74	29.91	10.00	0.05	0.56	10.57
	ZIVIB 83655	Pan trogloaytes	LIVI3	1	н	16.93	32.81	19.82	7.23	0.85	14.91
	MPI 13437	Pan troglodytes	RM1	1	J	14.83	33.63	19.60	8.08	0.76	13.04
	MPI 13433	Pan troglodytes	RM2	1	J	16.58	31.31	17.69	9.05	0.94	16.75
	MPI 13437	Pan troglodytes	RM2	1	J	17.17	37.89	19.36	9.13	0.89	14.41
	MPI 11800	Pan troglodytes	RM2	1	J	15.12	36.55	19.74	8.63	0.77	12.67
	MPI 11800	Pan troglodytes	LM3	1	J	13.76	31.30	18.54	7.96	0.74	13.26
	MPI 11779	Pan troglodytes	RM3	1	J	13.58	31.26	21.16	8.48	0.64	11.48
	ZMB 6954	Pongo sp.	LM1	1	Н	22.58	39.90	20.03	8.19	1.13	17.85
	ZMB 6957	Pongo sp.	RM1	1	Н	15.27	34.63	19.35	8.46	0.79	13.41
	ZMB 6987	Pongo sp.	RM1	1	н	17.90	40.77	19.81	9.16	0.90	14.15
	ZMB 30946	Pongo sp.	RM1	1	Н	32.10	62.09	24.56	11.55	1.31	16.59
	ZMB 67173	Pongo sp.	RM1	1	н	18.79	39.93	19.34	9.04	0.97	15.37
	SMF 1113	Ponao sp.	LM1	1	н	13.35	30.46	17.62	8.78	0.76	13.73
	SMF 1577	Pongo sp.	RM1	1	Н	19.46	47.58	22.65	10.74	0.86	12.46
	SMF 2654	Pongo sp.	IM1	1	н	16 76	39.42	19.69	10.01	0.85	13 56
	SME 28206	Pongo sp.		1	н	10.70	45 10	20.76	10.01	0.05	14.12
	7MR 605/	Pongo sp.		1	н	25.82	43.10	10 52	9.65	1 22	14.12
		Pongo sp.		1		23.82	27.00	10.41	9.05	1.52	19.09
		Ponyo sp.		1	п	25.92	57.09	19.41	9.29	1.25	20.24
	SIVIF 1117	Pongo sp.	RIVIZ	1	н	22.99	45.03	20.34	11.21	1.13	10.85
	SIME 2639	Pongo sp.	LIVI2	1	н	19.09	50.45	21.42	10.13	0.89	12.55
	SMF 15837	Pongo sp.	RM2	1	Н	22.17	48.89	22.32	9.90	0.99	14.20
	SMF 38296	<i>Pongo</i> sp.	LM2	1	н	25.32	32.34	20.94	12.19	1.21	21.26
	SMF 59140	<i>Pongo</i> sp.	RM2	1	Н	23.65	49.09	21.52	10.52	1.10	15.69
	SMF 59142	<i>Pongo</i> sp.	LM2	1	Н	25.34	51.66	21.11	11.59	1.20	16.70
	ZMB 6957	<i>Pongo</i> sp.	RM3	1	Н	22.42	28.76	17.36	8.22	1.29	24.09
	ZMB 83515	<i>Pongo</i> sp.	LM3	1	Н	18.83	46.46	21.19	10.11	0.89	13.03
	ZMB 12209	<i>Pongo</i> sp.	RM3	1	Н	18.86	27.67	16.19	9.19	1.16	22.15
	KNM-ER 820	Homo erectus	RM1	1	Р	23.06	36.28	19.55	10.28	1.18	19.59
	KNM-BK 67	Homo erectus	RM2	1	Q	25.75	30.51	18.55	10.25	1.39	25.13
	KNM-ER 992A	Homo erectus	RM2	1	Р	24.56	33.35	18.76	11.61	1.31	22.67
	KNM-ER 1507	Homo erectus	LM2	1	Р	29.98	38.25	19.40	11.72	1.55	24.99
	KNM-BK 67	Homo erectus	RM3	1	Р	26.41	32.48	17.89	10.10	1.48	25.89
	KNM-ER 992A	Homo erectus	RM3	1	Р	23.89	38.45	19.10	11.39	1.25	20.17
-	DNH 67	Homo sp. indet.	RM1	2	0	33.43	33.04	18.08	9.87	1.85	32.18
	KNM-FR 1802	Homo sp. indet	RM1	1	P	33.65	36 51	20.20	11 32	1 67	27 57
	KNM-FR 1802	Homo sp. indet	RM2	1	P	34.85	37.05	20.07	12 52	1 74	28 53
	KNM-ER 15064	Homo sp. indet.	RM2	1	P	30.88	/1 12	21 31	12.32	1 / 5	22.60
-	162-17	A gethionicus	RM2	3	N	48.06	62.32	21.51	12.24	2.12	22.00
		A. aethiopicus		2	IN N	40.00	71 22	22.02	14 70	2.12	20.92
	LI37-33	A. aethiopicus		с С	IN N	52.55	71.25	24.02	14.70	2.15	25.20
-		A. Uethopicus	RIVI3	2	<u>N</u>	57.04	73.35	25.81	15.49	2.21	25.80
	KMIN-ER 1820	A. boisei	LIMI	1	P	51.24	53.53	24.37	12.93	2.10	28.74
	L427-7	A. boisei	LM1	1	N	51.38	46.59	20.54	12.81	2.50	36.65
	Umo47-1973-1500	A. boiseí	RM1	3	N	36.05	46.45	21.31	13.72	1.69	24.82
	KNM-ER 3230	A. boisei	LM1	1	Р	65.00	72.45	25.89	16.80	2.51	29.50
	KNM-ER 15930	A. boisei	LM1	1	R	31.23	38.22	18.93	11.19	1.65	26.69
	L628-3	A. boisei	LM1	3	Ν	63.04	75.26	23.98	14.30	2.63	30.30
	KNM-ER 3230	A. boisei	RM1	1	Р	59.65	54.59	21.93	15.19	2.72	36.81
	KNM-ER 15930	A. boisei	LM1	1	R	49.37	47.59	20.72	13.00	2.38	34.53
•	DNH60B	A. robustus	RM1	1	0	31.62	31.62	18.61	10.80	1.70	30.22
	SK3974	A. robustus	RM1	2	S	39.92	41.17	18.99	10.39	2.10	32.76
	SK6	A. robustus	RM1	1	S	31.15	53.22	21.73	12.46	1.43	19.65

SK61	A. robustus	RM1	1	S	44.23	57.72	23.94	12.80	1.85	24.32
SK62	A. robustus	LM1	1	S	43.62	43.95	20.76	10.89	2.10	31.69
SK 63	A. robustus	RM1	1	S	38.67	41.58	20.15	10.99	1.92	29.77
SK(826b)828	A. robustus	LM1	2	S	40.79	55.35	24.27	12.01	1.68	22.59
DNH60C	A. robustus	RM1	1	0	38.93	32.92	17.98	11.77	2.17	37.74
SK6	A. robustus	LM1	1	S	49.42	54.13	21.655	14.17	2.28	31.02
SKW5	A. robustus	LM2	1	Т	43.46	49.06	20.56	12.40	2.11	30.18
SKX4446	A. robustus	RM2	1	Т	45.40	61.11	23.34	14.31	1.95	24.88
SK1587a	A. robustus	LM2	1	S	36.03	37.86	18.34	10.81	1.96	31.93
SK25	A. robustus	RM2	1	S	53.23	49.06	21.11	12.11	2.52	36.00
SK843.846a	A. robustus	LM2	1	S	42.51	51.00	21.09	13.03	2.02	28.22
SK1	A. robustus	LM2	2	S	45.20	58.01	23.13	13.44	1.95	25.66
SK6	A. robustus	LM3	1	S	48.99	48.18	21.94	13.91	2.23	32.18
SK23	A. robustus	LM3	1	S	42.52	51.97	22.40	12.71	1.90	26.33
SKW5	A. robustus	RM3	1	Т	45.34	49.44	20.46	12.49	2.22	31.51
SK843.846a	A. robustus	LM3	1	S	45.70	53.76	22.22	11.40	2.06	28.06
SK75	A. robustus	RM3	2	S	44.86	54.85	22.47	13.13	2.00	26.96
SK81	A. robustus	LM3	1	S	42.28	54.77	21.93	12.64	1.93	26.04
SKX10643	A. robustus	RM3	2	U	39.96	43.56	20.48	12.47	1.95	29.56
SKX5014	A. robustus	RM3	2	Т	38.94	57.27	22.64	14.06	1.72	22.73
TM1600	A. robustus	LM3	1	V	38.70	49.56	21.41	13.75	1.81	25.67
SK851	A. robustus	RM3	3	S	45.97	62.33	23.37	15.55	1.97	24.92
KNM-WT 8556	Uncertain	RM1	3	W	27.74	41.38	19.79	1.40	21.79	11.61
OmoK7-1969-19	Uncertain	LM1	3	х	38.60	35.89	21.02	1.84	30.65	11.58
OmoL26-1g	Uncertain	RM1	3	Ν	34.29	50.82	21.78	1.57	22.09	11.77
L795-1	Uncertain	RM2	3	Ν	51.70	63.66	24.21	2.14	26.76	14.82
L28-31	Uncertain	RM2	3	Y	44.45	36.41	19.29	2.30	38.18	9.77
Omo75s-1969-16	Uncertain	RM3	3	Ν	26.24	30.46	18.06	1.45	26.32	10.77
L28-30	Uncertain	RM3	3	Ν	41.02	36.11	17.99	2.28	37.94	10.49

Notes:

1. Basis -1 = molar in jaw or from associated dentition, 2 = molar position based on morphology and possible association with other teeth, 3 = molar position is best estimation based on morphology

2. BCD = bi-cervical diameter

3. AET = average enamel thickness

4. RET = relative enamel thickness

Citations: A – Johanson et al., 1982; B – Moggi-Cecchi et al., 2006; C – Dart, 1925; D – Mogg-Cecchi et al., 1998; E – Grine, 1981; F – Coffing et al., 1994; G – Ward et al., 2001; H – ZMB records; I – Michel Toussaint pers. Comm ; J – MPI-EVA records ; K – MRAC records ; L – ULAC records; M – FJR records; N – Suwa, 1996; O – Moggi-Cecchi et al., 2010; P – Wood, 1991; Q – Leakey et al., 1969; R – Leakey and Walker, 1988; S – Brain, 1981; T – Grine, 2004; U – De Ruiter, 2001; V – Thackeray et al., 2001; W – Brown et al., 2001; X – Alemseged personal communication; Y – Howell et al., 1987; Z – Dart, 1948.





Au. anamensis - KNM-ER 34725T LM2



Au. afarensis - AL128-23 RM2



Au. africanus - STW537 RM2



Au. boisei - L427-7 LM2





Homo sp indet - KNM-ER 1802 RM2



Homo erectus - KNM-ER 1507 LM2



Homo sapiens - R913-759 RM2



Pan troglodytes - ZMB 31279 RM2



Gorilla sp. - ZMB 31435 LM2





Pongo sp. - SMF 15837 RM2













Supplementary Material 1 Click here to download Supplementary Material: Supplementary Figure 1_Reconstructed_enamel.pdf Supplementary Material 2 Click here to download Supplementary Material: Supplementary Figure 2_Anamensis_M1.pdf Supplementary Material 3 Click here to download Supplementary Material: Supplementary Figure 3_Anamensis_M2_M3.pdf Supplementary Material 4 Click here to download Supplementary Material: Supplementary Figure 4_A_afarensis_all_teeth.pdf Supplementary Material 5 Click here to download Supplementary Material: Supplementary Figure 5_A_africanus_M1.pdf Supplementary Material 6 Click here to download Supplementary Material: Supplementary Figure 6_A_africanus_M2.pdf Supplementary Material 7 Click here to download Supplementary Material: Supplementary Figure 7_A_africanus_M3.pdf Supplementary Material 8 Click here to download Supplementary Material: Supplementary Figure 8_A_boisei_M1_M2_M3.pdf Supplementary Material 9 Click here to download Supplementary Material: Supplementary Figure 9_A_robustus_M1.pdf Supplementary Material 10 Click here to download Supplementary Material: Supplementary Figure 10_A_robustus_M2.pdf Supplementary Material 11 Click here to download Supplementary Material: Supplementary Figure 11_A_robustus_M3.pdf
Supplementary Material 12 Click here to download Supplementary Material: Supplementary Figure 12_Homo_erectus_all.pdf Supplementary Material 13 Click here to download Supplementary Material: Supplementary Figure 13_Homo_sp_all.pdf Supplementary Material 14 Click here to download Supplementary Material: Supplementary Figure 14_Homo_sap_modern_M1_M2_M3.pdf Supplementary Material 15 Click here to download Supplementary Material: Supplementary Figure 15_Pan troglodytes MPI_M1_M2_M3.pdf Supplementary Material 16 Click here to download Supplementary Material: Supplementary Figure 16_Pan troglodytes ZMB_M1_M2_M3.pdf Supplementary Material 17 Click here to download Supplementary Material: Supplementary Figure 17_Pan paniscus_M1_M2.pdf Supplementary Material 18 Click here to download Supplementary Material: Supplementary Figure 18_Gorilla_M1_M2_M3.pdf Supplementary Material 19 Click here to download Supplementary Material: Supplementary Figure 19_Pongo_M1_M2_M3.pdf