Evo-devo models of tooth development and the origin of hominoid molar diversity

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The detailed anatomical features that characterize fossil hominin molars figure prominently in the reconstruction of their taxonomy, phylogeny, and paleobiology. Despite the prominence of molar form in human origins research, the underlying developmental mechanisms generating the diversity of tooth crown features remain poorly understood. A model of tooth morphogenesis—the patterning cascade model (PCM)—provides a developmental framework to explore how and why the varying molar morphologies arose throughout human evolution. We generated virtual maps of the inner enamel epithelium—an indelibly preserved record of enamel knot arrangement—in 17 living and fossil hominoid species to investigate whether the PCM explains the expression of all major accessory cusps. We found that most of the variation and evolutionary changes in hominoid molar morphology followed the general developmental rule shared by all mammals, outlined by the PCM. Our results have implications for the accurate interpretation of molar crown configuration in hominoid systematics.

INTRODUCTION

Teeth are the most durable part of the skeletal system and form the dominant part of the hominin fossil record. Hence, analyses of hominin dental anatomy represent a cornerstone of paleoanthropological inquiry. Variation in one key aspect of tooth anatomy—molar crown configuration, or the expression in number, size, shape, and spatial disposition of molar cusps—has long been used for species diagnoses, phylogenetic and functional inferences, and population history reconstructions of Homo sapiens and our fossil hominin relatives (1–6). The developmental mechanisms that generate this variation, however, are poorly understood, obscuring interpretations of well-known evolutionary trends in molar cusp expression. Here, we test the applicability of a single morphogenetic rule to determine the number, size, and spatial distribution of molar cusps. In doing so, we contribute to a better understanding of the genotype-to-phenotype relationship driving variation in molar occlusal design over the past 15 million years (Ma) of hominoid evolution.

Over the past two decades, developmental studies of mammalian tooth germs suggest that growth of multicuspid teeth (premolars and molars) resembles a Turing-like mechanism governed by the iterative activation and silencing of embryonic signaling centers known as enamel knots (EKs) (7–10). EKs direct the growth and folding of the inner enamel epithelium, which acts as a “blueprint” for the final size and shape of the tooth crown (11, 12). EKs produce diffusible molecules that inhibit the formation of new knots nearby such that these new knots can only form outside the zones of inhibition of their previously initiated counterparts (10). The primary EK appears at the tip of the first cusp to initiate formation and controls the induction of secondary EKs arising along the inner enamel epithelium at the sites of the future cusps. The patterning cascade model (PCM) of tooth morphogenesis therefore postulates that molar cusp expression is determined by the interplay between the timing and spacing of EK initiation and the duration of crown growth before mineralization (7, 8, 13, 14). In a simplified example, a short duration of growth of the inner enamel epithelium, together with the late initiation of EK formation and increased inhibition, either reduces the likelihood of new cusp development or limits a cusp’s potential to attain a large size, assuming a constant rate of growth in time and space. The PCM has successfully reconstructed molar cusp variation of seal teeth and has been hypothesized to work across mammals (7–9, 13–15). Seal molars, however, are comparatively simple structures with a single row of mesiodistally oriented cusps and are unlike the quadrate molars of humans and most other primates. Thus, although predictions of the PCM in seals can be addressed using a single dimension (z axis), those in primates require mapping EK spatiotemporal relationships in three dimensions (3D), the approach taken in this paper. To date, supporting evidence of the PCM’s ability to explain hominoid molar morphological diversity has been equivocal and limited to one species, cusp, or molar type (16–20).

Here, we investigate the extent to which the PCM explains variation in molar cusp expression within Hominidae. Our sample comprises 763 molar teeth (M1 to M3) from six hominoid genera [Australopithecus (AUS), Paranthropus (PAR), Homo (HOM), Pan (PAN), Gorilla (GOR), and Pongo (PON)] and includes representatives of most hominin species currently recognized (see Materials and Methods). All major accessory cusps that are not primordially shared by all therian mammals were evaluated, including the hypocone, cusps 5 and 6, and Carabelli’s cusp (protoconal cingulum) of the upper molars, as well as cusps 5, 6 (“single” and “double”), and 7 of the lower molars (Fig. 1). Because the model was tested on fully formed teeth, we used the 3D form of the enamel-dentine junction (EDJ), which exactly reflects the endpoint of growth of the inner enamel epithelium, the embryonic anlage from which tooth crowns develop (11, 12, 18). Hence, dentine horns visible at the EDJ mark the precise location of EK initiation.

The 3D surface area of the EDJ and 3D intercusp distances were used as proxies for the size of the inner enamel epithelium and the spatiotemporal arrangement of EK formation, respectively. Following the logic of the PCM, the disposition of EKs across the epithelial landscape influences other aspects of molar crown configuration, such as cusp number and size (8, 13, 14). We classified all cusps according to their position on the crown: The hypocone and Carabelli’s cusp occur on the periphery, whereas all other accessory cusps were classified as central given that they develop between two previously initiated cusps. For peripheral cusps, we quantified the mean intercusp
distance (as a proportion of 3D surface area) between all earlier-initiating EKs. Consistent with the PCM, this approach assumes a close relationship between the size of the inhibitory fields and the proximity of EKs and predicts that a relatively small mean intercusp distance between previously initiated EKs makes an additional cusp more likely to initiate on the crown periphery (17, 19). Central cusps, on the other hand, face an additional spatial limitation such that they are more likely to be present and/or more strongly expressed when their two adjacent and earlier-initiated cusps are more widely spaced (Fig. 1) (18).

RESULTS AND DISCUSSION

Our analyses of samples of Pleistocene to recent H. sapiens and 10 species of fossil hominins using rank correlation coefficients and odds logistic regression revealed that most of the diversity in molar cusp features can be explained by the PCM. A similar pattern was found for all great apes, suggesting that this developmental mechanism of cusp expression was present in the last common ancestor of extant hominoids. For peripheral cusps, we found that hypocone presence was significantly and negatively correlated with mean relative intercusp distance of previously formed cusps. Our results for Carabelli’s cusp, however, show a negative relationship between cusp expression and mean relative intercusp distance only in Australopithecus and Gorilla (Fig. 2, Table 1, figs. S1 and S2, and table S1). For central cusps of the upper molars, cusp 5 expression is positively correlated with relative metacne-hypocone spacing in Australopithecus, Paranthropus (P ≤ 0.05), and Gorilla (P ≤ 0.05), whereas the occasional presence of cusp 6 is also associated with the greatest metacne-hypocone distances.
Similarly, lower molar cusps 5 and 6 are more likely to develop and to be larger in teeth with expanded hypoconid-entoconid and entoconid-hypoconulid distances, respectively (\(P \leq 0.05\) in all six hominoid genera examined). Our results for double cusp 6—observed in our *Australopithecus*, *Paranthropus*, and *Homo* samples—are also suggestive of the validity of the PCM in that we found a positive relationship between cusp expression and hypoconulid–tuberculum sextum spacing, although results were significant only in *Paranthropus*. Results for cusp 7 were less clear, with a positive but nonsignificant relationship between cusp 7 and metaconid-entoconid distance in *Australopithecus* and *Homo* (Figs. 2 and 3, Table 1, and table S1).

Although direct experimental testing is impossible in fossils and unfeasible in most extant mammals, the application of a developmental-based model evaluating the complex form of the EDJ strongly suggests that the wide range of variation in hominoid cuspal patterns is primarily regulated by EKs such that the size of their inhibitory fields constrains the presence and size of subsequently developing cusps. The strongest support for the PCM comes from the hypocone and lower molar cusps 5 and 6 (single and double). Reported frequencies of cusp expression for hominoids provide an additional way to validate the PCM as tested in seals. When cusp variation is analyzed along the \(z\) axis and all other parameters are held constant, in high-cusped molars, EKs initiate farther apart than in bunodont molars. Hence, bunodont molars should exhibit both lower cusp height differentials and higher frequencies of accessory cusps (8, 9, 13, 14). Although the bunodont molars of *Paranthropus* and *Australopithecus* have higher frequencies of accessory cusps relative to later and higher-cusped hominin molars, high-cusped *Gorilla* molars do not have a distinctly high frequency of extra cusps (1, 21). Given the limited differential variability for selection to act upon, it is believed that going from high-cusped to a bunodont configuration is easier than the reverse (15). Thus, considering the highly variable and heritable nature of enamel thickness (22), it has been suggested that the evolution of hyperthick enameled teeth in *Paranthropus* was the “least resistant” adaptive response for fracturing tough food in otherwise suboptimal flat molars (23).

Although we highlight the power of the PCM to explain variation in cusp expression within the hominin lineage, our results also point to some deviations from this highly conserved developmental program. The PCM does not fully explain variation of Carabelli’s cusp and cusp 7, and, to a lesser extent, upper molar cusp 5. We hypothesize that one reason for these disparities in Carabelli’s cusp and cusp 7 is that mild expressions of these two features do not involve a dentine horn. In contrast, even the weakest degrees of expression of all other cusps analyzed are invariably associated with a dentine horn, and thus
with the unequivocal presence of an EK (24, 25). This suggests that a slightly different developmental pathway may account for Carabelli’s cusp and cusp 7 morphogenesis. A recent study has identified variants in the transcription factor FOXI3 that lead to incomplete development of lingually oriented cusps (26). The PCM derives from a wide range of studies on mice and seals, mammals with highly derived dentitions. Considering that humans and mice share an ancestry going back ~100 Ma and a shared ancestry with seals that goes even deeper in

**Table 1. Kendall’s rank correlation coefficient between accessory cusp expression and relative intercusp distance.** Sample size for each pairwise comparison in parentheses. Significant values (at $P < 0.05$) bolded. Blank cells indicate that the feature was invariably absent/present or that samples did not have enough grades of expression for statistical analysis. AUS, Australopithecus; PAR, Paranthropus; HOM, Homo; HSr, recent H. Sapiens; PAN, Pan; GOR, Gorilla; PON, Pongo.

| Taxon | HYP | UMC5 | CC | LMC5 | LMC6 | LMC6D | LMC7 |
|-------|-----|------|----|------|------|-------|------|
| AUS   | —   | 0.21 (35) | — | — | 0.39* (48) | 0.29 (13) | 0.26 (19) |
| PAR   | —   | 0.22* (39) | — | — | 0.37* (46) | 0.38* (38) | —   |
| HOM   | —0.24* (94) | —0.03 (113) | —0.01 (79) | 0.55* (144) | 0.36* (127) | 0.17 (29) | 0.14 (39) |
| HSr   | —0.23* (48) | 0.07 (49) | 0.14 (35) | 0.56* (104) | 0.39* (67) | —       | —   |
| PAN   | —   | —0.02 (49) | 0.09 (35) | — | 0.28* (69) | —       | —0.04 (46) |
| GOR   | —0.51* (16) | —0.16 (11) | —       | — | —       | —       | —   |
| PON   | —   | —0.26 (16) | 0.13 (14) | — | 0.33* (29) | —       | —   |

*Adherence to PCM predictions (only significant results considered).
Our results are consistent with the emerging picture that a single basic molar crown configuration observed throughout human evolution. Developmental explanation for these and other long-standing patterns of cusp expression (30), and recently, it was found that malnutrition and systemic diseases in humans lead to developmental disruptions that increased variability in molar cusp expression, including the presence of extra cusps (30).

Broad themes in molar cuspal patterns exist and have shaped various phylogenetic or taxonomic schemes of human origins and diversity, including the differential expansion of the talonid in *P. robustus* and *Paranthropus boisei* (2), the higher frequency of lower molar cusps 6 and 7 in *Paranthropus* and *Australopithecus*, respectively (1, 31), the large and protruding hypocones of Neandertals (4), or the trend toward the reduction or loss of the hypocone and lower molar cusp 5 in recent Europeans (3). By testing the PCM, we have provided a developmental explanation for these and other long-standing patterns of molar crown configuration observed throughout human evolution. Our results are consistent with the emerging picture that a single basic developmental program (with the differential composition of its parameters) is responsible for the generation of morphological novelties and the myriad dental phenotypes found in mammalian teeth. These results also point to the highly homoplastic nature of accessory cusps and have implications for the accurate interpretation of dental phenotypic variation in hominoid systematics.

### MATERIALS AND METHODS

**Materials**

The study sample consisted of 276 and 487 upper and lower molars, respectively (Table 2). The extinct hominin sample included 12 *Australopithecus anamensis*, 13 *Australopithecus afarensis*, 80 *A. africanus*, 2 *Paranthropus aethiopicus*, 9 *P. boisei*, 83 *P. robustus*, 15 *Homo habilis sensu lato*, 9 *Homo erectus sensu lato*, 5 Middle Pleistocene hominins, 91 *Homo neanderthalensis*, and 37 Pleistocene *H. sapiens*. The complete list of the fossil specimens used is provided in tables S2 and S3. Reference samples of 177 recent *H. sapiens*, 115 *Pan troglodytes*, 27 *Pan paniscus*, 38 *Gorilla* spp., and 40 *Pongo* spp. molars were also examined. No antemers were included except for rare cases where cusp number differed between the left and right sides (tables S2 and S3).

**Data collection procedures and statistical analyses**

Each molar was subjected to microcomputed tomography to produce 3D surface models of the internal tooth structure at the EDJ. These models were subsequently manipulated on Avizo/Amira (FEI Visualization Sciences Group) for landmark digitizing and trait scoring. Accessory cusp expression was collected following standardized scoring protocols summarized in table S4. Intercusp distances (Euclidean distances) were derived from homologous landmarks placed on the dentine horns. Tooth size was calculated using the 3D surface area of the EDJ crown (32) following the parameters recommended by Pampush et al. (33). Following the study of Hunter et al. (17), both nonparametric Kendall’s rank correlation tests and ordered logistic regression models were used to examine the relationship between relative intercusp spacing (that is, intercusp distance divided by tooth size) and the presence and degree of expression of accessory cusps. Most analyses were performed at the genus level given the small samples of most fossil species available for study. A detailed explanation of the methods used can be found in the Supplementary Materials.

### SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/4/eaar2334/DC1

**Supplementary Materials and Methods**

fig. S1. Relationship between mean relative intercusp distance and Carabelli’s cusp per genus.

fig. S2. Relationship between mean relative intercusp distance and Carabelli’s cusp per molar type in *Homo*.

fig. S3. Right lower molar with example of homologous landmarks (yellow dots) placed at the cusp tips from which Euclidean distances were calculated.

fig. S4. Ordinary least squares regression of upper (top row) and lower (bottom row) molar size comparisons estimated from crown outline, centroid size, and 3D surface area. 

**table S1.** Ordinatory least squares regression of upper (top row) and lower (bottom row) molar size comparisons estimated from crown outline, centroid size, and 3D surface area.

**table S2.** Fossil hominin upper and lower molars used in this study including accession number, locality/ site, and source.

**table S3.** Fossil hominin lower molars used in this study including accession number, locality/ site, and source.

**table S4.** System used in this study for scoring accessory cusps.

**table S5.** Tooth size comparisons estimated from crown outline, centroid size, and 3D surface area.

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