

MAMMALIAN ENAMEL PRISM PATTERNS AND ENAMEL DEPOSITION RATES

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Abstract

Enamel prism patterns and enamel deposition rates were compared for specimens representing six mammalian orders. Enamel samples were characterized by either pattern 1 or pattern 3 prisms. Each prism pattern category contained prisms from at least two mammalian orders. Enamel deposition rate was estimated for each sample by measuring prism cross striation repeat intervals. Statistical analysis of cross striation repeat intervals illustrates significant differences in deposition rate between prism patterns 1 and 3. No statistically significant differences were found in deposition rate between the higher-level taxa represented within each prism pattern category. That enamel deposition rate is not taxon-specific reinforces the close association between deposition rate and prism morphology. In accord with previous studies, pattern 1 enamel is deposited more slowly than is pattern 3 enamel. Correlation analyses illustrated a lack of association between enamel deposition rate and body mass, tooth size, and estimated ameloblast size. Evidence that enamel deposition rate is associated with enamel prism morphology, coupled with evidence that deposition rate is not correlated with size parameters, points to developmental homology (i.e., homogeneous deposition rate) within each prism pattern.

Key Words: Enamel, prism patterns, deposition rates, cross striation repeat intervals, evolution.

Introduction

Mammalian dental enamel is mainly composed of submicroscopic hydroxyapatite crystallites. Discontinuities surrounding bundles of similarly oriented crystallites define enamel prisms, which extend from the dentine core of a tooth toward the outer surface of the enamel. Enamel prism patterns are divided into three basic categories based on their prism shape and packing patterns as seen in tangential sections of mature enamel (Fig. 1) (Korvenkontio, 1934-1935; Boyde, 1964, 1967, 1971). Prism pattern 1 consists of prisms with complete boundaries that are arranged in offset horizontal rows with respect to the apico-cervical axis of the tooth. In pattern 2, prisms are arc-shaped and arranged in offset vertical rows separated by a distinct inter-row sheet, while arc-shaped prisms arranged in offset horizontal rows constitute pattern 3.

A growing body of evidence suggests that enamel prism patterns are correlated with variations in enamel deposition rates (Moss, 1969; Osborn, 1970; Martin, 1983, 1985; Martin and Boyde, 1984; Fortelius, 1985). The consistency of this relationship has implications concerning the value of prism patterns in evolutionary studies. If prism patterns are rate dependent, changes in secretory rate could be the cause of apparent divergence, convergence, and parallelism in enamel prism pattern morphology. Alternatively, if prism patterns are not rate dependent, then variation in rate within prism pattern categories may serve to characterize specific mammalian lineages.

The goal of this study is to test two hypotheses concerning the relationship between enamel prism pattern morphology and enamel deposition rate. The first hypothesis states that enamel prism pattern morphology is associated with specific enamel deposition rates. To test this hypothesis, enamel deposition rates were compared between prism pattern categories that were each represented by several different mammalian orders. The combination of several orders in each category ensured that comparisons would reflect differences among prism patterns rather than differences among taxonomic groups.

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The second hypothesis states that, within a prism pattern category, different taxa are characterized by taxon-specific enamel deposition rates. This hypothesis was addressed through comparisons of enamel deposition rates between distantly related taxa within each of two prism pattern categories. Finally, associations between deposition rate and body mass, tooth size, and ameloblast size were assessed using correlation analyses.

Materials and Methods

Taxonomic sample

Taxa were selected based on a priori expectations of prism packing patterns that were drawn from the literature. The living scandentian *Tupaia glis* and the erinaceids *Atelerix albiventris* and *Erinaceus europaeus* were selected to represent pattern 1 taxa (Boyde, 1964; Silness and Gustavsen, 1969; Shellis and Poole, 1977; Shellis, 1984a). Recent human (*Homo sapiens*), dermopteran (*Cynocephalus variegatus*), and chiropteran (*Balantiopteryx plicata*, *Rhinopoma hardwickei*, and *Taphozous mauritianus*) enamel was selected to represent pattern 3 taxa, as was enamel from the fossil family Microsyopidae (*Microsyops* sp.) (Martin, 1983; Lester and Hand, 1987; Lester *et al.*, 1988; Martin *et al.*, 1988; Dumont, 1993). Single specimens of *Galagoides alleni*, *G. demidovii*, and the fossil primate *Notharctus* sp. were also anticipated to express prism pattern 3 (Dumont, 1993). The living cercopithecoid primates *Macaca fascicularis*, *Cercocebus torquatus* and *Cercocebus albigena* were chosen to represent prism pattern 2, which has been reported to occur within this primate family (Boyde and Martin, 1984a, 1984b, 1987; Martin *et al.*, 1988).

Microscopic techniques

Specimens were embedded in polymethylmethacrylate and sectioned longitudinally through the protoconid and metaconid tips. The sectioned specimens were then polished with successively finer diamond pastes (6 μm , 4 μm , 1 μm , and 0.25 μm). Between polishing treatments, specimens were cleaned with a detergent (Mr. Clean[®]) and rinsed with distilled water for approximately five minutes to insure that all grit and oils were removed. When polishing was complete, specimens were cleaned again, allowed to air dry overnight, and mounted with Duco Cement[®] on scanning electron microscope (SEM) stubs. Following an overnight curing period, specimens were etched with 0.5% H_3PO_4 for seven to 20 seconds; smaller recent specimens required less etching than did larger fossil specimens. The chemical reaction was stopped by immediately quenching the specimens, with agitation, in a distilled water bath for one minute. Specimens were then rinsed under running

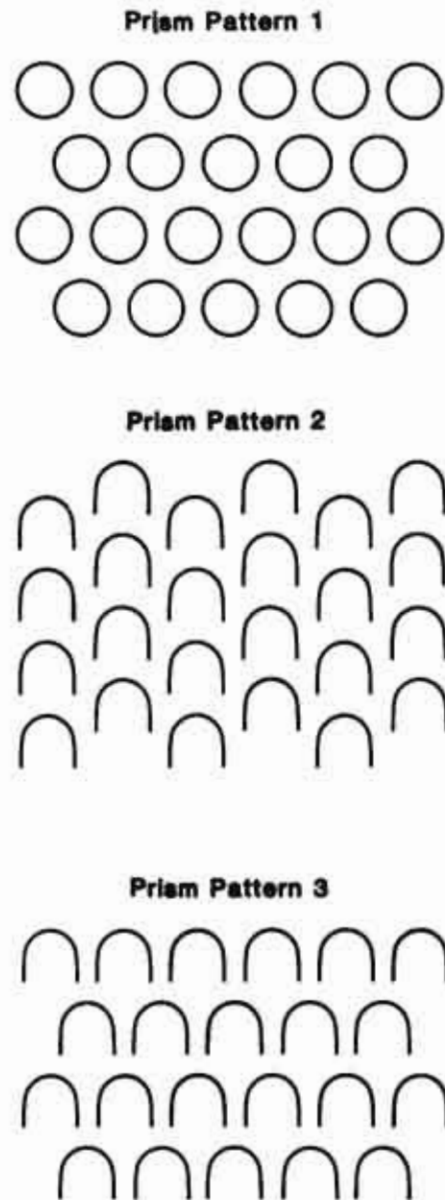


Figure 1. Diagram illustrating enamel prism patterns 1, 2 and 3. Each pattern exhibits a unique combination of enamel prism shape and spatial organization.

tap water for one minute and immediately rinsed again in fresh distilled water for 30 seconds. Following another overnight drying period, specimens were sputter-coated with silver for 60 to 70 seconds in ten second bursts in final preparation for SEM viewing.

All microscopy was accomplished using an AMRAY[®] model 1810D SEM equipped with a solid state backscattered electron detector. Accelerating voltages (kV) ranging from 25 to 30 were used in conjunction with a working distance of 9 to 12 mm, a condenser lens setting between 2.5 and 4.0, and either

Prism patterns and deposition rates

Table 1. Summary by specimen of methods used to couple prism pattern and cross striation repeat intervals. Taxon, specimen number (Number), number of micrographs (Nmicro), total number of measured cross striations (Nstriae), and sampled tooth position (Tooth) is listed for each specimen. Methods used to link prism pattern and cross striation listed in order of decreasing reliability are: direct evidence of prism pattern on the same micrograph (Direct), prism pattern determined by confocal microscopy of the same specimen (TSM), and prism pattern determined by confocal microscopy of other specimens of the same species (TSother, (N) = sample size).

Taxon	Number	Nmicro	Nstriae	Direct	TSM	TSother (N)	Tooth
PATTERN 1							
<i>Atelerix albiventris</i>	FSM 20551	3	30	x			right M ¹
	FSM 20552	3	50	x			right M ₁
	FSM 20553	4	55	x			right M ₁
<i>Erinaceus europaeus</i>	TT 49630	3	11	x			right M ₁
	TT 49631	1	14	x			right M ₁
<i>Tupaia glis</i>	SUSB (1)*	2	14			x (3)	right M ₁
	SUSB (2)*	3	40	x			right M ₁
	SUSB (3)*	1	7			x (3)	right M ₁
PATTERN 3							
<i>Balantiopteryx plicata</i>	TT 38121	4	56			x (4)	right M ₁
	TT 38122	3	60		x		right M ₁
	TT 38123	1	4		x		right M ₁
	TT 38128	2	27		x		right M ₁
<i>Cynocephalus variegatus</i>	DZUM 100	2	21			x (5)	right M ₁
	FMNH 56505	3	74			x (5)	right M ₁
	FMNH 56524	1	9			x (5)	right M ₁
	RVNH 15822	1	2			x (5)	right M ₁
<i>Homo sapiens</i>	SUSB (1)*	3	35	x			right M ₁
	SUSB (2)*	3	75	x			left M ₁
	SUSB (3)*	2	47			x	right M ₁
	SUSB (4)*	1	22			x	right M ₁
	SUSB (5)*	3	75	x			right M ₁
<i>Microsyops</i> sp.†	CM (1)*	1	4	x			right M ₁
	CM (3)*	1	2	x			M _x frag.
<i>Rhinopoma hardwickei</i>	TT 40638	2	22		x		right M ₁
	TT 40640	1	12	x			right M ₁
<i>Taphozous mauritanus</i>	CM 85237	3	72			x (11)	right M ₁
	CM 85241	3	65			x (11)	right M ₁
MIXED PATTERNS 2 AND 3							
<i>Cercocebus albigena</i>	SUSB 85-17	5	105			x (1)	right M ₃
	SUSB 85-7	1	5			x (1)	right M ₃
<i>Cercocebus torquatus</i>	ANSP 3072	1	18			x (1)	right M ₁
	ANSP 12645	4	97			x (1)	right M ₁
	ANSP 11840	3	75			x (1)	right M ₁
<i>Macaca fascicularis</i>	(1)*	1	10			x (1)	right M ₁
	(2)*	3	45			x (1)	right M ₁
SINGLE SPECIMENS ILLUSTRATING PATTERN 3							
<i>Galagoides demidovii</i>	SUSB PGa1	2	21			x (2)	right M ₁
<i>Notharctus</i> sp.†	CM*	1	6	x			M _x frag.
PRISM PATTERN NOT DETERMINED							
<i>Galagoides alleni</i>	CM 3898	3	66				right M ₁

* = uncataloged specimens;

† = fossil taxon.

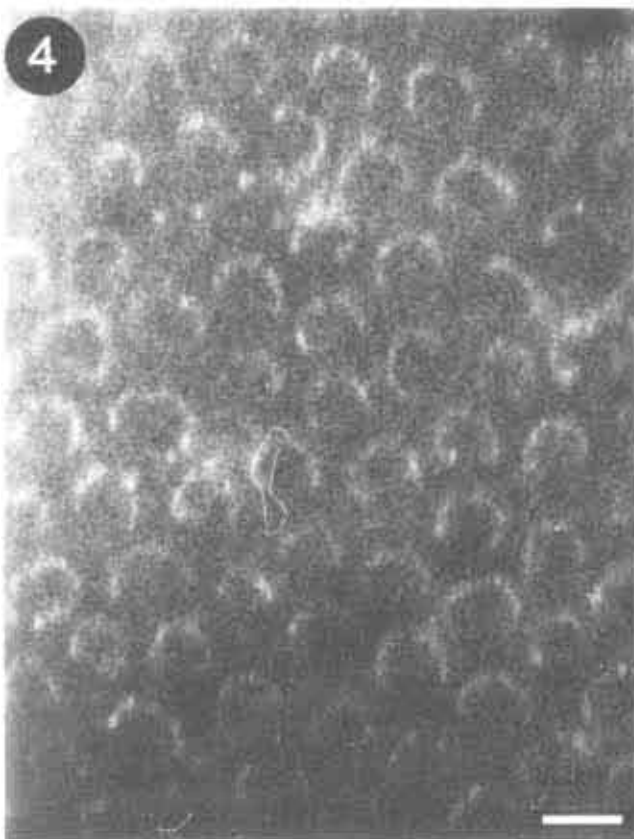
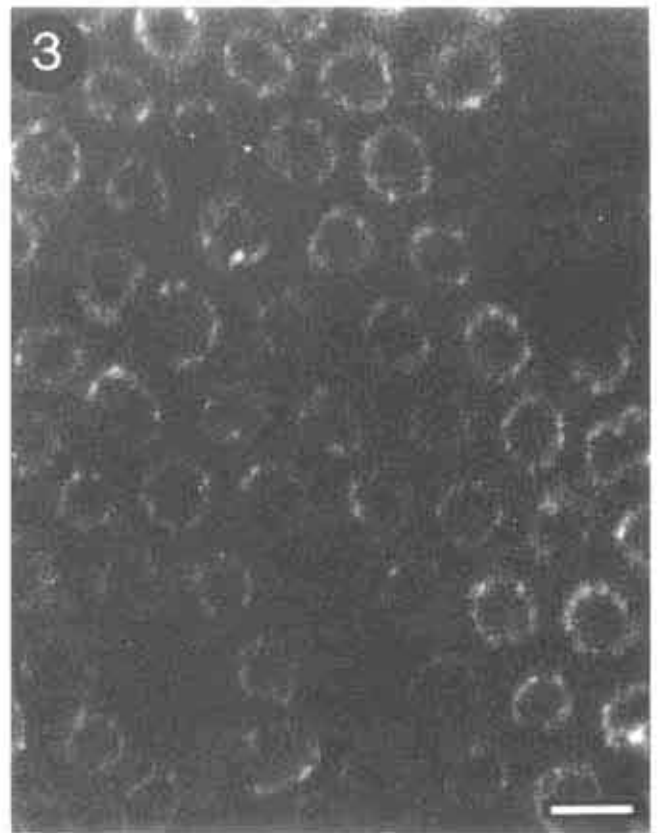
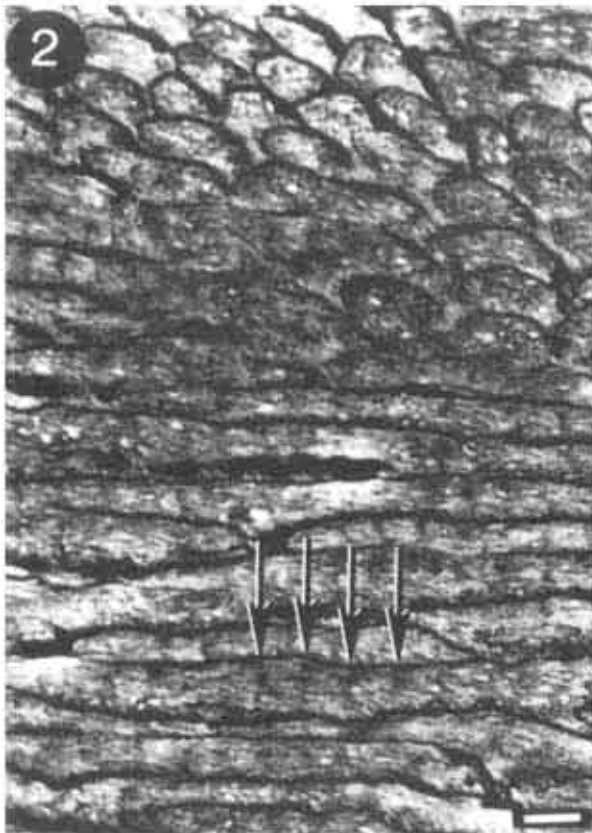


Figure 2. BSE image of longitudinally sectioned human enamel. Pattern 3 prisms are visible within the upper portion of the micrograph. Arrows point to a series of cross striations along a single prism. Bar = 10 μm .

Figure 3. Confocal image of *Tupaia glis* (SUSB) enamel taken at a depth of 25 μm below the buccal surface of the right M¹ hypoconid. Bar = 5 μm .

Figure 4. Confocal image of *Cynocephalus variegatus* enamel (RVNH 14516) taken at a depth of 25 μm below the buccal surface of the right M¹ protoconid. Bar = 5 μm .

a 200 μm or 300 μm aperture. This combination of settings was found through trial and error to produce the highest resolution and maximum contrast, while providing an acceptable signal to noise ratio.

Association between cross-striations and prism pattern

Enamel for analysis of enamel deposition rates was sampled from the central portion of the enamel thickness of 38 molar teeth, 77 areas from either the mid-thickness of the buccal aspect of the protoconid or the lingual aspect of the metaconid and 7 areas from the trigonid basin.

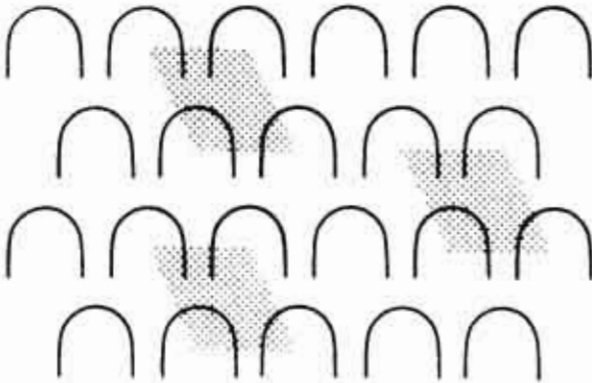


Figure 5. Using formulas developed by Fosse (1968a), the estimated area of a single ameloblast is calculated as the area of a parallelogram drawn between the centers of adjacent prisms. In this diagram, three parallelograms are represented as shaded areas.

Because enamel prism pattern morphology can vary within a tooth, it was important to associate measurements of enamel prism cross striations with cross-sectional prism morphology as directly as possible. Table 1 presents a summary of the sampled specimens and the methods used to couple prism pattern categories and photomicrographs of longitudinally sectioned prisms.

For 35% of the specimens, prism pattern assignments were confirmed by viewing cross-sectioned prisms exposed on longitudinal sections in regions adjacent to those in which prism cross striations were present (Fig. 2). For a few specimens, prism pattern was confirmed by examining tangential sections of subsurface enamel using confocal microscopy. Prism pattern could not be resolved for the single specimen of *Galagoides alleni*. Evidence for prism pattern assignments summarizing the remaining 50% of the specimens were based on the prism pattern found on tangential sections of homologous teeth from conspecific individuals; these sections were viewed using confocal microscopy (Figs. 3 and 4).

Measurement methods

Cross striations and micron scales on enlarged micrographs were traced and measured on acetate overlays. Distances between the centers of adjacent cross striation repeat intervals were measured parallel to prism long axes to the nearest 0.1 mm using dial calipers. Sampling of cross-striations on the periphery of micrographs was avoided whenever possible. The orientation of cross striations to larger incremental features was not investigated. Up to 25 cross striation repeat intervals were measured from each micrograph. Each cross striation was numbered for future reference. All measure-

ments were brought to the same scale prior to statistical comparisons.

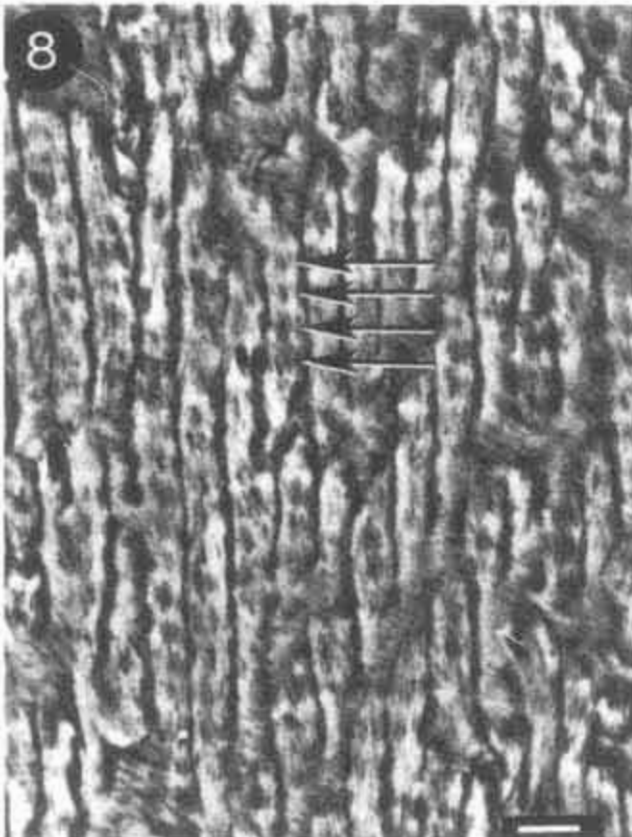
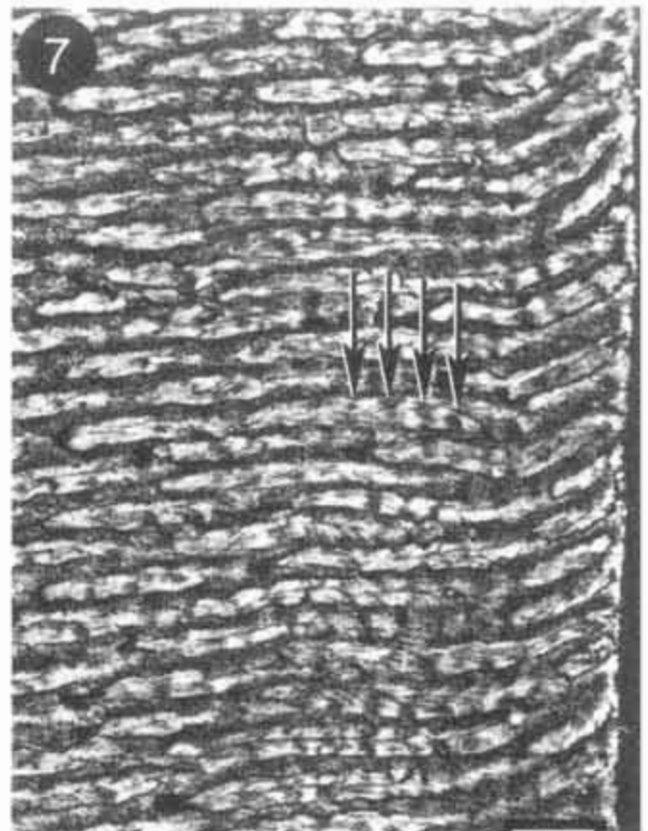
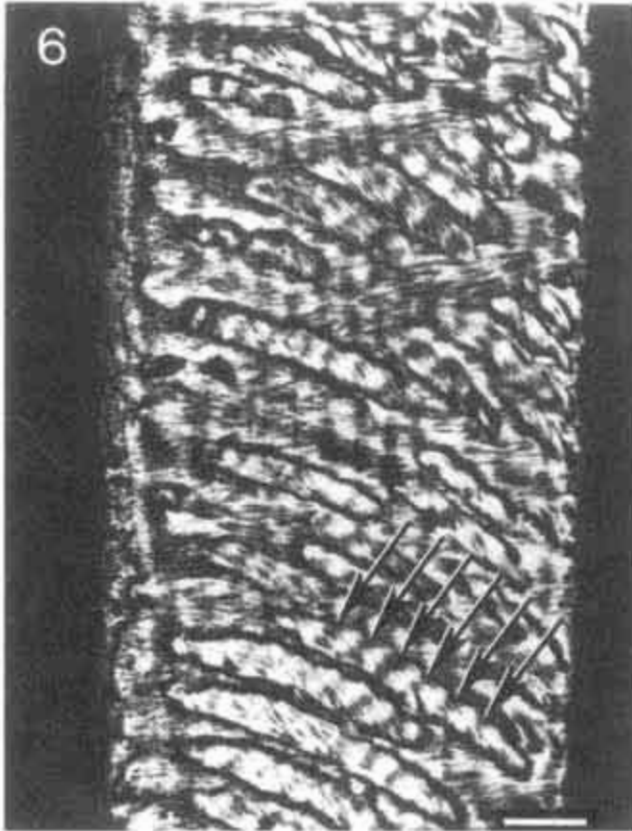
Statistical analysis

The two hypotheses concerning prism pattern and cross striation repeat intervals were addressed separately using single classification analysis of variance (ANOVA). This method was determined to be the most appropriate method of comparison, as the data are both normal and homoscedastic (Sokal and Rohlf, 1981). In analyzing whether cross striation repeat intervals are significantly different between pattern 1 and pattern 3 enamel, specimen means of cross striation repeat intervals from pattern 1 taxa (*Erinaceus*, *Atelerix*, and *Tupaia*) were compared to specimen means from pattern 3 taxa (*Taphozous*, *Balantiopteryx*, *Rhinopoma*, *Homo*, *Cynocephalus*, and *Microsyops*).

In order to test the second hypothesis, that significant variation in cross striation repeat intervals exists among taxa within prism pattern categories, scandentians were compared to erinaceids for the pattern 1 case, while chiropteran, *Cynocephalus*, *Homo*, and *Microsyops* specimens were compared simultaneously for the pattern 3 case. Again, specimen means were used to represent variation that occurs among individuals within each higher-level group.

Associations between cross striation repeat interval and body mass, tooth area (mesiodistal length x buccolingual breadth), and estimated ameloblast area were assessed using correlation analyses. Estimated ameloblast area was calculated using confocal images of tangentially sectioned enamel prisms from molar teeth of conspecific individuals. Following the method developed by Fosse (1968a) and used by many subsequent workers (Carlson and Krause, 1985; Fosse *et al.*, 1985; Grine *et al.*, 1986, 1987; Krause and Carlson, 1986), estimated ameloblast area was calculated as the area of a parallelogram drawn between the centers of four adjacent prisms (Fig. 5).

All variables were transformed to a linear scale by taking roots and logged to reduce the effects of magnitude on the correlation coefficient (Smith, 1984). Because the transformed data were normal and displayed homogeneous variances, the parametric Product-Moment correlation test was used to analyze each set of paired data (Sokal and Rohlf, 1981). These analyses do not rely on prism pattern assignments and data from the three sampled cercopithecoid primates were included. Samples from single specimens (i.e., *Galagoides demidovii*, *Galagoides alleni* and *Notharctus* sp.) were also incorporated into the analysis to provide a larger sample size. Due to missing data, one or more taxa were omitted from each analysis; *Homo* was excluded from analyses of tooth area; *Galagoides alleni* and the



Figures 6-9 are on the facing page.

Figure 6. BSE image of longitudinally sectioned prisms in *Taphozous mauritanus* (CM 85237) enamel. Arrows point to a series of cross striations along a single prism. Bar = 10 μm .

Figure 7. BSE image of longitudinally sectioned prisms in *Galagoides alleni* (CM 3898) enamel. Arrows point to a series of cross striations along a single prism. Bar = 10 μm .

Figure 8. BSE image of longitudinally sectioned prisms in *Homo sapiens* (SUSB, not cataloged) enamel. Arrows point to a series of cross striations along a single prism. Bar = 10 μm .

Figure 9. BSE image of longitudinally sectioned prisms in *Cercocebus torquatus* (ANSP 12645) enamel. Arrows point to a series of cross striations along a single prism. Bar = 10 μm .

cercopithecoid primates were excluded from analyses of ameloblast area; and fossil taxa were excluded from analyses involving body mass.

Results

When viewed using backscattering electron microscopy (BSE), cross striations appear as alternating light and dark bands (Figs. 6-9). With the exceptions of *Homo sapiens* and *Erinaceus europaeus*, prisms within these taxa exhibit relatively straight courses from the enamel-dentine junction to the outer enamel surface (i.e., there was no evidence of prism decussation zones).

Comparisons of cross striation repeat intervals between individuals of the same species using single classification ANOVA illustrated that in 8 of 9 cases, conspecific individuals differ significantly from one another. This indicates that cross striation repeat intervals from a single specimen are unlikely to represent the range of cross striation values for an entire species. Therefore, species represented by only one individual (e.g., *Notharctus* sp., *Galagoides demidovii*, and *Galagoides alleni*) were deleted from comparisons of cross striation repeat intervals between prism pattern categories. In addition, it was not possible to confirm the sole presence of pattern 2 prisms in any of the species that were expected to exhibit the prism pattern. Therefore, the prism pattern 2 category was deleted from further analyses.

Table 2 presents summary statistics for each sampled taxon, as well as results of statistical comparisons of prism cross striation repeat intervals within and between prism pattern 1 and 3 categories. Taxa charac-

terized by prism pattern 1 exhibit more narrowly spaced cross striations than do taxa that are characterized by prism pattern 3. Cross striation repeat intervals are significantly different between the two prism pattern categories ($p < .01$). Within each pattern category, values representing species were very similar. Comparisons between species within each prism pattern category illustrated that there were no significant differences between representatives of different mammalian orders.

Table 3 presents cross striation repeat intervals, body mass, tooth area, and ameloblast area data for each species, as well as correlation coefficients summarizing associations between cross striation repeat interval and each of the size variables. The correlation coefficients for body mass and tooth area analyses are of similar magnitude at $r = 0.23$ and $r = 0.30$, respectively. The correlation between cross striation repeat intervals and ameloblast area is slightly lower at $r = 0.16$. None of these correlation coefficients differ significantly from zero.

Discussion

In his original description of prism patterns, Boyde (1964) described several subdivisions of the pattern 2 and 3 categories that were subsequently elaborated by Gantt (1982, 1983). Although many taxa within this study exhibit pattern 3 enamel, it proved difficult to assign pattern 3 prisms to any one subdivision because of minor variations in prism distribution within even limited areas of individual teeth. Rather than using ill-fitting subcategories, all taxa with unambiguous prism patterns were categorized as either prism pattern 1 or 3. Although Shellis (1984a) described dermopterian enamel as exhibiting pattern 2 prisms, no evidence of consistent vertical prism stacking or inter-row sheets was encountered in this analysis and dermopterians were retained in the prism pattern 3 category.

It proved impossible to verify the exclusive presence of pattern 2 prisms in the cercopithecoid primates. The specimens used in this study exhibited arc-shaped prisms that were arranged in both pattern 3 and pattern 2 spatial distributions. No clear indications of inter-row sheets were seen in any specimens. A similarly mixed distribution of arc-shaped prisms among cercopithecoid primates have been reported by several other workers (Shellis and Poole, 1977; Shellis, 1984a, 1984b; Grine *et al.*, 1985; Martin *et al.*, 1988), adding support to the current finding of mixed patterns 2 and 3 in these taxa.

Within any individual tooth, enamel prism pattern morphology exhibits variations attributable to changes in ameloblast (enamel matrix secreting cell) Tomes' process configuration during enamel deposition; a layer of aprismatic enamel resides at both the enamel-dentine

Table 2. Summary statistics for each sampled taxon and results of statistical comparisons of cross striation repeat intervals within and between prism patterns 1 and 3. Number of specimens (N), means and standard deviations ($X \pm SD$) are reported for each taxon. F-values and associated probability statements derived from ANOVA are given for comparisons within (WITHIN) and between (BETWEEN) prism pattern categories.

TAXON	N	$X \pm SD$ in μm	WITHIN	BETWEEN
PRISM PATTERN 1				
<i>Atelerix albiventris</i>	3	$2.51 \pm .098$	F = 3.648 n.s.	F = 11.400 p < .01
<i>Erinaceus europaeus</i>	2	$2.44 \pm .361$		
<i>Tupaia glis</i>	3	$1.88 \pm .730$		
PRISM PATTERN 3				
<i>Balantiopteryx plicata</i>	4	$2.86 \pm .895$	F = 1.181 n.s.	
<i>Rhinopoma hardwicki</i>	2	$2.88 \pm .828$		
<i>Taphozous mauritianus</i>	2	$3.40 \pm .286$		
<i>Cynocephalus</i> sp.	4	$3.06 \pm .667$		
<i>Homo sapiens</i>	5	$3.71 \pm .349$		
<i>Microsyops</i> sp.†	2	3.21 ± 1.29		
OTHER SAMPLED TAXA				
<i>Cercocebus albigena</i>	2	$4.83 \pm .968$		
<i>Cercocebus torquatus</i>	3	$5.21 \pm .525$		
<i>Macaca fascicularis</i>	3	$3.15 \pm .418$		
<i>Galagoides alleni</i>	1	4.98 ± 1.106		
<i>Galagoides demidovii</i>	1	$6.16 \pm .830$		
<i>Notharctus</i> sp.†	1	$2.39 \pm .362$		

† = fossil taxon;

n.s.: not significant.

junction and the outer enamel surface of most mammalian teeth and is typically underlain and overlain, respectively, by a layer of pattern 1 enamel. These layers were formed as ameloblasts began and ended their enamel-secreting cycles (Boyde, 1964; Ripa *et al.*, 1966; Gwinnett, 1967; Martin, 1983; Fortelius, 1985; Martin *et al.*, 1988). Variation in prism pattern beyond that already mentioned above is also common within single teeth (e.g., von Koenigswald, 1992; von Koenigswald and Clemens, 1992). However, several studies of primate enamel have demonstrated that prism patterns are relatively constant in mid-thickness enamel (e.g., Boyde and Martin, 1982, 1984a, 1984b; Martin *et al.*, 1988).

This level of consistency in mid-thickness prism patterns is also characteristic of the non-primate taxa included in this analysis (Dumont, 1993).

Enamel deposition rates may be estimated in mature enamel by measuring prism cross striation repeat intervals. This prism striation has been interpreted as representing circadian variation in enamel deposition rate (e.g., Gysi, 1931; Schour and Poncher, 1939; Boyde, 1964, 1979, 1989; Boyde and Martin, 1982, 1984a; Shellis, 1984b; Bromage and Dean, 1985; Risnes, 1986; Beynon and Reid, 1987; Dean, 1987a, 1987b). Although studies by several workers have suggested that cross striations are artifactual (e.g., Osborn, 1971;

Prism patterns and deposition rates

Table 3. Raw cross-striation repeat interval (C.S.R.I.), tooth area, mean ameloblast area and body mass data. Body mass data for sexually dimorphic species is reported as a mean of male and female values. Product-moment correlation coefficients (r) between (log) cross striation repeat interval and (log)($\sqrt{2}$ estimated ameloblast area), (log)($\sqrt{2}$ tooth area), and (log)($\sqrt{3}$ body mass) and the sample size (N) for each comparison are provided. None of the coefficients differ significantly from 0.

Taxon	Mean C.S.R.I.	Tooth Area	Mean A.A.	Body Mass
Order Lipotyphla				
<i>Atelerix albiventris</i>	2.51	12.46	17.72	485 g ¹
<i>Erinaceus europaeus</i>	2.44	23.33	20.98	912.5 g ²
Order Scandentia				
<i>Tupaia glis</i>	1.88	7.51	30.80	200 g ²
Order Chiroptera				
<i>Balantiopteryx plicata</i>	2.86	1.10	21.44	7.5 g ³
<i>Rhinopoma hardwickei</i>	2.88	1.64	25.50	11 g ¹
<i>Taphozous mauritianus</i>	3.40	2.53	26.42	22.5 g ¹
Order Dermoptera				
<i>Cynocephalus variegatus</i>	3.06	14.75	29.38	1,250 g ³
Order Primates				
<i>Cercocebus albigena</i>	4.83	15.13	-	7,690 g ⁴
<i>Cercocebus torquatus</i>	5.21	62.88	-	10,625 g ⁴
<i>Galagoides alleni</i>	4.98	7.34	-	295 g ⁴
<i>Galagoides demidovii</i>	6.16	10.63	28.30	60 g ²
<i>Homo sapiens</i>	3.71	-	31.36 ⁵	60,000 g ²
<i>Macaca fascicularis</i>	3.15	28.84	-	4,030 g ⁴
<i>Notharctus</i> sp.†	2.39	12.01	34.11	-
Order incerta sedis				
<i>Microsyops</i> sp.†	3.21	11.90	39.94	-
(log) Cross Striation Repeat Interval Against		(log) (sqrt) Tooth Area	(log) (sqrt) A.A.	(log) (cbrt) Body Mass
		$r = .30$	$r = .16$	$r = .23$
		$N = 14$	$N = 11$	$N = .13$
		n.s.	n.s.	n.s.

† = fossil taxon;

n.s.: not significant.

¹Kingdon (1974); ²Eisenberg (1981) ³Walker (1975); ⁴Fleagle (1988); ⁵Fosse (1968b).

Weber and Glick, 1975; Warshawsky and Bai, 1983; Warshawsky *et al.*, 1984), Bromage (1991) has provided experimental evidence that cross striations indeed represent cyclical variation in enamel deposition. Similar studies have demonstrated that such rhythms also characterize dentine deposition and endochondral ossification (e.g., Yilmaz *et al.*, 1977; Simmons, 1974; Kawasaki *et al.*, 1980). This study assumes that prism cross striations are manifestations of a constant physiological rhythm. Provided that the periodicity is the same for all taxa, the length of the rhythm period is not relevant to this analysis.

Because cross striations are considered in part to be manifestations of variation in carbonate concentrations within fully mineralized enamel (Boyde, 1979; Boyde and Jones, 1983), BSE was selected as the most appropriate technique for observing prism cross striations morphology. Although contrast in BSE images of perfectly flat specimens that are oriented perpendicular to the electron beam indicated only variation in atomic number (Postek *et al.*, 1980), specimens used in this study were etched, and consequently exhibited some surface topography. Comparisons of secondary electron and BSE images of a subset of the specimens illustrated that contrast permitting visualization of the cross striations was primarily based on BSE signal.

Statistical comparisons of cross striation repeat intervals support the hypothesis that depositional rates differ significantly between prism pattern categories. Based on the data presented here, pattern 1 enamel is deposited significantly more slowly than is pattern 3 enamel. That is, within the same period of time, larger segments of pattern 3 enamel prisms are deposited by an ameloblast than along pattern 1 enamel prisms. These differences in depositional rates transcend ordinal boundaries and appear to point to a basic relationship between ameloblast activity and the structure of fully mineralized enamel.

In contrast to the first hypothesis, the data analyzed here failed to support the second hypothesis that higher-level taxa are characterized by taxon-specific cross striation repeat interval values. Among pattern 1 taxa, prism cross striation repeat intervals do not differ significantly between erinaceids and scandentians. Similarly, there are no significant differences in cross striation repeat intervals among the four sampled pattern 3 taxa (*chiropterans*, *Homo sapiens*, *Microsyops* sp., and *Cynocephalus variegatus*). That diverse taxa exhibiting the same prism patterns are homogeneous with respect to deposition rate again supports the conclusion that prism pattern and depositional rate are closely associated.

Despite the suggestions of previous workers that enamel deposition rate is correlated with either body or ameloblast size (Boyde, 1969; Martin, 1983), no evi-

dence for these assertions was found within the data presented here. Based on these data, deposition rates are not correlated with either tooth area, body mass, or estimated ameloblast area. By inference, these data also suggest that prism pattern is not correlated with any of these factors.

The association between prism pattern and cross striation repeat intervals obtained in this study are in general accord with those reported by other workers. Intervals for pattern 1 enamel have been reported in most instances to remain below 2.5 μm (Martin, 1983; Boyde and Martin, 1984; but see Shellis and Poole, 1977). This is supported by the species means for pattern 1 taxa within this study which range from 1.88 μm to 2.51 μm . Cross striation repeat interval values for pattern 3 enamel have been reported to range between 2 μm and 7 μm (Shellis and Poole, 1977; Martin, 1983; Martin and Boyde, 1984; Risnes, 1986). Although the mean cross striation repeat intervals reported here for pattern 3 enamel are on the low end of the reported range, they are nonetheless significantly higher than those found to characterize pattern 1 cross striation repeat intervals. In sum, the data presented here support the consensus opinion that pattern 1 cross striation repeat intervals are much smaller than those for pattern 3.

The length of cross striation repeat intervals for the cercopithecids are most similar to those of the pattern 3 species, though they are significantly larger ($p < .03$). While cercopithecoid prism patterns exhibit a large proportion of pattern 2 spatial distribution, they do not exhibit the clearly defined inter-row sheets that are characteristic of pattern 2 enamel. The degree of variability in the spatial distribution of cross-sectioned prisms also makes it difficult to assign these taxa to prism pattern 3. These data suggest that slightly modified definitions of prism patterns may more accurately reflect the variability in the spatial distribution of arc-shaped prisms that is encountered as more mammalian species are sampled. For example, the consistent presence of vertically oriented inter-row sheets could be the determining factor in categorizing an enamel as pattern 2 while prisms pattern 3 could encompass all other spatial distributions of arc-shaped prisms.

The single specimens representing *Galagoides demidovii* and *Notharctus* sp. exhibit pattern 3 prisms. Although prism pattern could not be resolved for *G. alleni*, it is likely that it too exhibits the pattern 3 enamel that characterizes its congener. The cross striation repeat intervals of the two galago species accord most closely with those of other pattern 3 taxa. In contrast, the mean cross striation repeat interval of *Notharctus* falls into the range of prism pattern one taxa. Nevertheless, it also lies within one standard deviation of other pattern 3 cross striation repeat interval means. Because the cross

striation repeat interval mean for *Notharctus* is based on a sample of only six cross-striations, it is likely that further study of *Notharctus* enamel will lead to a more reliable assessment of the range of cross striation repeat interval values for the species.

Summary and Conclusions

Prism patterns are significantly associated with enamel deposition rates. Therefore, fluctuations in rate among lower-level taxa may be associated with the heterogeneous prism patterns often seen at higher taxonomic levels. The immediate mechanism producing changes in deposition rate is unknown, although such changes could arise through the pleiotropic effects of alterations in any number of physiological systems.

Variation in prism patterns is common within orders and, to a lesser extent, within families (Boyde, 1964; Carlson and Krause, 1985; Skobe *et al.*, 1985; Martin *et al.*, 1988; von Koenigswald and Clemens, 1992; Dumont, 1993). Despite this variation, many family-level taxa can be characterized by a typical prism pattern. This conservation of prism pattern suggests that although prism patterns are associated with depositional rates, these rates and the factors controlling them do not vary extensively within families. It seems likely, therefore, that prism patterns are potentially reliable phylogenetic characters when they are applied below the ordinal level. However, recognizing the presence of variation in prism pattern within and between individuals, one must sample enamel prism patterns at homologous tooth positions and depths for several individuals from each taxon in order to accurately assess the predominant prism pattern of any taxonomic group.

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Discussion with Reviewers

S. Risnes: In the paper, it is suggested that the prism pattern is dependent on the rate of enamel production. Is it not possible that the opposite is the case, i.e., that the rate of enamel production is dependent on the prism pattern, i.e., on the spatial arrangement of the ameloblasts?

Author: I have attempted to refrain from confusing correlation with causation when discussing the relationship between the rate of enamel deposition and prism pattern. The data presented here do not address the causal relationship between these two variables. It is entirely possible that either the rate of enamel deposition or the spatial arrangements of ameloblasts is the factor that drives the relationship between the two. It is also possible that these factors are not causally related but that prism pattern and deposition rate are both mediated by other, yet undocumented, variables.

S. Risnes: The term "enamel deposition rate" is ambiguous until the direction of the incremental growth is defined. Generally, this term means the growth in thickness of the enamel layer along a direction perpendicular to the incremental lines (Retzius lines). Since enamel prisms often deviate from such a direction, the rate of enamel deposition along the direction of the prisms will have to be higher than the rate of enamel deposition along a direction perpendicular to the incremental lines. To what extent will a distinction between these two rates affect your interpretations and discussion?

Author: In this study, "enamel deposition rate" was considered to be the rate of enamel accretion along individual prism long axes. The relationship of this rate of enamel deposition was not studied relative to larger incremental features (i.e., Retzius lines). Clearly, the data presented here do not directly address the issue of incremental growth as defined by an increase in enamel thickness. For the reasons that you cite, the distinction between prism accretion rate and the rate of increase in enamel thickness is critical for structurally complex enamels. However, most of the taxa included in this study do not exhibit prism decussation and, because of the relatively simple course taken by the prisms, it is possible that the rates of prism deposition reported here are, at least in part, reflective of the rate of enamel

increase. Certainly, a detailed assessment of the relationship between the two rates is required before definitive statements can be made.

W.A. Clemens: In your analysis of cross striation repeat intervals, you make use of mean values for specimens. Did you detect any repeated patterns in variation in the length of interval related to the area of the tooth (cusp slope or trigonid basin) sampled?

Author: For the one instance in which data collected from the buccal aspect of the protoconid slope and the trigonid basin were combined to generate a species mean (*Galagoides alleni* CM3898), cross striae from within the trigonid basin were significantly smaller (3.92 ± 0.162 versus 5.92 ± 0.1 ; $p < .001$). Because this study was designed to focus on enamel sampled from the external aspect cusp slopes, this is the only individual for which this comparison can be made. This result is intriguing, however, since it suggests that separate functional surfaces may develop in different ways (at least in respect to enamel deposition rates).

W.A. Clemens: Did you detect any correlation between thickness of mature enamel and prism type? Were thicker enamels usually characterized by the presence of pattern 3 prisms?

Author: Enamel thickness data for several species of known enamel prism type are available (Dumont, in press). Species that exhibit pattern 3 prisms exhibit on average slightly higher relative enamel thickness values (6.44 ± 1.76 , $N = 7$) than do species that exhibit pattern 1 prisms (5.0 ± 0.57 , $N = 2$). However, these values are not significantly different ($p = 0.31$). Clearly, these sample sizes are quite small and additional data is required to accurately assess the relationship between prism pattern and enamel thickness.

M.A. Saunder: If prism pattern and deposition rate are correlated, could prism patterns be a by-product of different prism deposition rates necessary to form certain enamel types? For example, maybe Hunter-Schreger Bands require faster deposition than radial enamel. In an enamel organ depositing both enamel types at the same time (as is commonly the case), two different prism patterns in the same enamel would be the necessary result. This could be an explanation of the limited phylogenetic information content of prism patterns.

Author: It is entirely possible that variation in prism patterns is associated with variation in deposition rates needed to form different types of enamel. This is a very interesting hypothesis in that it proposes a functional requirement as the driving force for variation enamel deposition rates. I would venture that further investigation along these lines would provide some interesting results.

Additional Reference

Dumont ER. Enamel thickness and dietary adaptation among extant primates and chiropterans. *J. Mammal* (in press).