Spatio-temporal diversity of dietary preferences and stress sensibilities of early and middle Miocene Rhinocerotidae from Eurasia: impact of climate changes

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Abstract

Major climatic and ecological changes are documented in terrestrial ecosystems during the Miocene epoch. The Rhinocerotidae are a very interesting clade to investigate the impact of these changes on ecology, as they are abundant and diverse in the fossil record throughout the Miocene. Here, we explored the spatio-temporal evolution of rhinocerotids’ paleoecology during the early and middle Miocene in Europe and Pakistan. We studied the dental texture microwear (proxy for diet) and enamel hypoplasia (stress indicator) of 19 species belonging to four sub-tribes and an unnamed clade of Rhinocerotidae, and coming from nine Eurasian localities ranging from MN2 to MN7/8. Our results suggest a clear niche partitioning based on diet at Kumbi 4 (MN2, Pakistan), Sansan (MN6, France), and Villefranche d’Astarac (MN7/8, France), while dietary overlap and subtle variations are discussed for Béon 1 (MN4, France) and Gračanica (MN5/6, Bosnia-Herzegovina). All rhinocerotids studied were browsers or mixed-feeders, and none had a grazing nor frugivore diet. Regarding hypoplasia, the prevalence was moderate (~ 10%) to high (> 20 %) at all localities but Kumbi 4 (~ 6 %), and documented quite well the local conditions. Sansan and Devinska Nová Ves (MN6, Slovakia), both dated to the MN6 (i.e., by the middle Miocene Climatic Transition, ca. 13.9 Mya), had moderate hypoplasia prevalence. Besides locality, species and tooth locus were also important factors of variation for the prevalence of hypoplasia. The very large hippo-like Brachypotherium brachypus was
one of the most affected species at all concerned localities (but Sansan), while early-diverging
elasmotheriines were very little affected.

Keywords: paleoecology, Miocene Climatic Optimum (MCO), microwear (DMTA), enamel hypoplasia

Introduction

The Miocene is a key period in Earth and rhinocerotid evolutionary histories. Climatic conditions in
Eurasia during the Miocene epoch are globally tropical and the typical habitat is forested (Cerling et
al., 1997; Zachos et al., 2001; Bruch et al., 2007; Westerhold et al., 2020). It is the last warm episode
of the Cenozoic era, although marked by great climatic changes prefiguring the setup of modern cold
conditions (Westerhold et al., 2020). During early Miocene times, temperatures increased until
reaching the Miocene Climatic Optimum (MCO) between ~17 to 14 Mya (corresponding to the late
Burdigalian + Langhian standard ages; Westerhold et al., 2020). This optimum is followed by an
abrupt cooling (the middle Miocene climatic transition [mMCT]; Westerhold et al., 2020) associated
with faunal turnovers in Eurasia (Böhme, 2003; Maridet et al., 2007). The middle Miocene is marked
by a relative aridity, associated with a global cooling (Bruch et al., 2007; Böhme et al., 2008).

Concerning rhinocerotids, Miocene times witness peaks in their alpha diversity about 22–18 Mya and
11–10 Mya (Antoine et al., 2010; Antoine and Becker, 2013; Antoine, in press). During the early and
middle Miocene in Eurasia, four sub-tribes of Rhinocerotidae are encountered – Rhinocerotina,
Teleoceratina and Aceratheriina (Rhinocerotinae), and Elasmotheriina (Elasmotheriinae) – and
species of which are often found associated in fossil-yielding localities (Antoine et al., 1997, 2010;
Heissig, 2012; Becker and Tissier, 2020; Antoine, 2002, in press). This abundance and the potential
cohabitation of such large herbivores question habitat capacity and competition for food resources.
However, the ecology of the rhinocerotids has rarely been explored or only been assumed based on
morphological adaptations (Prothero et al., 1989; Prothero, 2005; Giaourtsakis et al., 2006). If the
Rhinocerotina appear to be ecologically varied, the literature suggests a similar ecology for most
elasmotheriines on one hand, as open environment dwellers adapted to though vegetation (Iñigo and
Cerdeño, 1997; Antoine and Welcomme, 2000), and for the teleoceratines on another hand, as hippo-
like rhinoceroses inhabiting lake side or swamps and probably browsing on low vegetation or even grazing (Prothero et al., 1989; Cerdeño, 1998).

In this article, we focused on the rhinocerotids from nine localities, covering wide temporal and geographical ranges (from MN2 to MN7/8 and from southwestern France to Pakistan). We assessed dietary preferences using dental microwear texture analysis, and stress sensibility via the study of enamel hypoplasia.

**Material and methods**

We studied the rhinocerotid dental remains from nine early and middle Miocene localities from France (Béon 1, Béon 2, Sansan, Simorre, and Villefranche d’Astarac), Germany (Steinheim am Albuch), Bosnia-Herzegovina (Gračanica), Slovakia (Devínska Nová Ves Spalte), and Pakistan (Kumbi 4, Balochistan), ranging from MN2 to MN7/8. The rhinocerotid assemblages are detailed in Table 1. The specimens are curated at the Naturhistorisches Museum Wien (NHMW), the Muséum de Toulouse (MHNT), and the Naturhistorisches Museum Basel (NHMB). For all details on the specimens included in this study see Supplementary S1. The localization of the studied localities is given in Figure 1. Further details on the localities are given in Supplementary S2.

**Dental Microwear Texture Analyses (DMTA)**

Dental Microwear Texture Analysis (DMTA) is a powerful approach to characterize dietary preferences at a short time scale (days to weeks prior the death of the individual; Hoffman et al., 2015; Winkler et al., 2020), widely used in paleontological and archeological studies (Grine, 1986; Rivals et al., 2012; Jones and DeSantis, 2017; Berlioz et al., 2018). We studied dental microwear texture on one well-preserved molar (germs and over-worn teeth excluded) per individual, preferentially the second molar (first or third otherwise), either upper or lower, left or right.

After cleaning the tooth with acetone or ethanol, two silicone (Regular Body President, ref. 6015 - ISO 4823, medium consistency, polyvinylsiloxane addition type; Coltene Whaledent) molds were made on a single enamel band, which shows two different facets acting as grinding and shearing (if present).
This shearing facet has a steep slope while the former is more horizontal and show several HSB band on the very enamel surface. To combine both type of facets with different functions indeed improves dietary reconstruction (Louail et al., 2021; Merceron et al., 2021). The enamel band on which we identified those two facets is localized labially near the protocone on upper molars and distally to the protoconid or hypoconid (if the protoconid is unavailable) on lower teeth (see supplementary S2).

Figure 1: Geographical position of the studied Eurasian Miocene localities. Localization of all localities in Eurasia. Red square is a zoom on the southwestern French localities, modified from Antoine and Duranthon (1997). Color code by MN zones as detailed in A. Abbreviations from West to East: S- Steinheim am Albuch (MN7/8; Germany), D- Devínska Nová Ves Spalte (MN6; Slovakia), G- Gračanica (MN5-6; Bosnia-Herzegovina), K- Kumbi 4 (MN2; Pakistan).
Table 1: List of rhinocerotid species found at each locality studied

<table>
<thead>
<tr>
<th>Locality</th>
<th>Mesaceratherium welcommi</th>
<th>Pleuroceros blanfordi</th>
<th>Protaceratherium sp.</th>
<th>Protaceratherium minutum</th>
<th>Plesiaceratherium naricum</th>
<th>Plesiaceratherium mirallesi</th>
<th>Plesiaceratherium balkanicum</th>
<th>Plesiaceratherium sp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kumbi 4</td>
<td>x</td>
<td></td>
<td></td>
<td></td>
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<td>x</td>
<td>x</td>
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<td>Béon 2</td>
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<td></td>
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<td></td>
<td></td>
<td>x</td>
<td>x</td>
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<tr>
<td>Béon 1</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>x</td>
<td>x</td>
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<tr>
<td>Gračanica</td>
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<td></td>
<td></td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>Sansan</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>Devínska Nová Vés Spalte</td>
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<td></td>
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<td></td>
<td></td>
<td>x</td>
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<tr>
<td>Steinheim am Albuch</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>x</td>
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<tr>
<td>Simorre</td>
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<td></td>
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<tr>
<td>Villefranche d’Astarac</td>
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</tr>
</tbody>
</table>

Rhinocerotinae

Aceratheriina

Hoploaceratherium tetradactylum
Alicornops simorrense

Rhinocerotina

Gaindatherium cf. browni
Lartetotherium sansaniense
Dicerorhinus steinheimensis

Teleoceratina

Brachypotherium brachypus
Brachypotherium gajense
Diaceratherium fatehjangense
Prosantorhinus douvillei
Prosantorhinus shahbazi

Elasmotheriinae

Elasmotheriina

Bugtirhinus praecursor
Hispanotherium beonense
Hispanotherium cf. matritense

Total: 9 3 5 4 4 2 4 2 2

In this article we followed a protocol adapted from Scott et al. (2005, 2006) with sensitive-scale fractal analyses. Molds were scanned with a Leica DCM8 confocal profilometer (“TRIDENT” profilometer housed at the PALEVOPRIM, CNRS, University of Poitiers) using white light confocal technology with a 100× objective (Leica Microsystems; Numerical aperture: 0.90; working distance: 0.9 mm).
obtained scans (.plu files) were pre-treated with LeicaMap v.8.2. (Leica Microsystems) as follows: the surface was inverted (as scans were made on negative replicas), missing points (i.e., non-measured, less than 1%) were replaced by the mean of the neighboring points and aberrant peaks were removed (see details in the supplementary Information in Merceron et al., 2016b). The surface was then levelled, and we applied a polynomial of degree 8 removal of form to temper for Hunter-Schreger bands reliefs in the DMTA parameters. Eventually, we selected a 200×200-μm area (1551 × 1551 pixels) within the surface, which we saved as a digital elevation model (.sur) and used to extract DMTA parameters through Scale-Sensitive Fractal Analysis with SFrax (Surfract, www.surfract.com) and LeicaMap.

Here we focused on five classical DMTA parameters: anisotropy (exact proportion of length-scale anisotropy of relief; epLsar), complexity (area-scale fractal complexity; Asfc), heterogeneity of complexity (heterogeneity of area-scale fractal complexity here at 3×3 and 9×9; HAsfc9 and HAsfc81), and fine textural fill volume (here at 0.2 μm; FTfv). The description of these parameters is available in Scott et al. (2006).

To facilitate DMTA interpretation for fossil specimen, we used specimens of the five extant rhinocerotid species. This extant dataset was modified from that of Hullot et al. (2019), as precised below, and consists of 17 specimens of Ceratotherium simum (white rhinoceros), four of Dicerorhinus sumatrensis (Sumatran rhinoceros), 21 of Diceros bicornis (black rhinoceros), 15 of Rhinoceros sondaicus (Javan rhinoceros; one new specimen), and five of Rhinoceros unicornis (Indian rhinoceros; one new specimen).

**Enamel hypoplasia**

Hypoplasia is a common defect of the enamel resulting from a stress or a combination of stresses occurring during tooth development (Goodman and Rose, 1990). It is a permanent, sensitive, but non-specific indicator of stresses either environmental (e.g., drought or nutritional stress; Skinner and Pruetz, 2012; Upex and Dobney, 2012), physiological (e.g., disease or parasitism; Suckling et al., 1986; Rothschild et al., 2001; Niven et al., 2004), and/or psychological (e.g., depression in primates; Guatelli-Steinberg, 2001).
Enamel hypoplasia was studied with the naked eye and categorization of the defects followed the *Fédération Dentaire Internationale* (1982) as linear enamel hypoplasia (LEH), pitted hypoplasia, or aplasia. We studied all cheek teeth, both deciduous and permanent, but excluded 62 teeth to avoid false negative and uncalibrated defects, as enamel was obscured (e.g., tooth unerupted in bone, sediment occluding), broken or worn out, or as identification was impossible. This left 1401 teeth studied for the hypoplasia analysis – 294 milk molars and 1107 permanent premolars and molars – from the nine localities. In parallel, qualitative data (tooth locus affected, position of the defect on the crown, and severity) and caliper measurements (distance of the defect from enamel-dentine junction, width if applicable) were taken (details in Supplementary S3). Type of defects recorded, and caliper measurements are illustrated in Figure 2.

Figure 2: The three different types of hypoplasia considered in this study and the associated measurements

A- Lingual view of right M2 of the specimen MHNT.PAL.2004.0.58 (*Hispanotherium beonense*) displaying three types of hypoplasia

B- Interpretative drawing of the photo in A illustrating the hypoplastic defects: a- pitted hypoplasia, b- linear enamel hypoplasia, and c- aplasia

C- Interpretative drawing of the photo in A illustrating the measurements: 1- distance between the base of the defect and the enamel-dentine junction, 2- width of the defect (when applicable).

Figure from Hullot et al. (2021).
Statistics and GLMMs

Statistics were conducted in R (R Core Team, 2018: https://www.R-project.org/), equipped with the following packages: reshape2 (Wickham, 2007), dplyr (Wickham et al., 2019), lme4 (Bates et al., 2015), car (Fox et al., 2012), MASS (Venables and Ripley, 2002). According to the recent statement of the American Statistical Association (ASA) on p-values (Wasserstein and Lazar, 2016; Wasserstein et al., 2019), we avoided the use of the term “statistically significant” in this manuscript and the classical thresholds as much as possible. Figures were done using R package ggplot2 (Wickham, 2011) as well as Inkscape v.0.91.

General Linear Mixed Models (GLMM) on our data were constructed based on a R code modified from Arman et al. (2019) and adapted to each tested response variable. An example of this code applied to hypoplasia variable Hypo is given in Supplementary 4. DMTA response variables were the five DMTA parameters (epLsar, Asfc, FTfv, HAsfc9, and HAsfc81) and we selected Gaussian family for the GLMMs. Factors in the models were: specimen (number of the specimen; random factor), locality, province, age (MN zones), genus, tooth (e.g., second molar, fourth milk molar), position (upper or lower), side (left or right), cusp (protocone, protoconid, hypoconid), and facet (grinding or shearing).

For hypoplasia, response variables were Hypo (1 or 0 for presence or absence of hypoplasia, respectively) for which we used Binomial family, Defect (e.g., LEH, Pits, Aplasia; converted to numbers), Localization (position of the defect on the crown; mostly labial or lingual), Multiple (number of defects), and Severity (0 to 4), modeled using Poisson family. The factors were: specimen (number of the specimen; random factor), locality, province, age (MN zones), genus, tooth (e.g., first molar, fourth premolar), position (upper or lower), side (left or right), and wear (low, average, high).

Additionally, for response variables Severity, Multiple, and Localization, defect was converted and used as a factor.

The models were built with a bottom-up approach, starting with the only random factor of our dataset alone (specimen) and adding factors incrementally for every set (e.g., 1|Specimen + Genus, 1|Specimen + Locality). New set was built has long as Akaike’s Information Criterion score (AIC) kept decreasing. Few interactions (e.g., Genus x Facet for microwear, Genus x Tooth for hypoplasia) were considered in the models, as most factors were considered independent and to avoid unnecessarily
complex and rarely selected models (Arman et al., 2019). We selected the best candidate model as the one with the lowest AIC and checked for over-dispersion (estimated through the ratio of deviance and degrees of freedom). If needed, we corrected it through quasi-Poisson or quasi-Binomial laws from the MASS package (Venables and Ripley, 2002) or by adjusting the coefficients table (multiply type error by square root of the dispersion factor and recalculate Z and p values accordingly). In total, 340 models were compared across the 10 response variables (see electronic supplementary material, S5, S6, and S7).

Results

Microwear

MANOVA (Species x Facet x Age x Locality) on all five main DMTA parameters (epLsar, Asfc, FTfv, HAfc9, HAfc81) revealed low p-values for Species (df = 14; p-value = 8.6 x 10^{-4}), Facet (df = 1; p-value = 6.5 x 10^{-4}), and Locality (df = 4; p-value = 0.014). The ANOVAs for each parameter, highlighted at least a marked influence of Species (all parameters; p-values between 7.3 x 10^{-4} and 0.027), Facet (Asfc, p-value = 0.028; FTfv, p-value = 6.22 x 10^{-6}), Age (Asfc, p-value = 7.57 x 10^{-3}), or Locality (epLsar, p-value = 0.01; Asfc, p-value = 1.7 x 10^{-4}). To precise the differences for Species and Locality (factors with more than two states) we ran post hocs, results of which are detailed in Table 2 and Table 3. The more conservative post hoc (Tukey's honestly significant difference; HSD) revealed very few noticeable differences in the microwear textures of the studied rhinocerotid specimens by Species or Locality (low p-values relatively to other pairs; Table 2). The DMT of Hoploaceratherium tetradactylum appears quite distinct from that of Plesiaceratherium spp. (epLsar, Asfc, HAfc9 and HAfc81). Concerning Locality, Gračanica specimens stood out with very low values of complexity compared to Sansan (p-value = 0.047), Simorre (p-value = 0.028), and Villefranche d’Astarac (p-value = 0.032).

The least conservative post hoc (Fischer’s least significant difference; LSD) highlighted more differences in the DMTA patterns of the specimens regarding Species and Locality (Table 3). Alicornops simorense and P. mirallesi cluster together with higher anisotropy than P. douvillei, B. brachypus, B. gajense, M. welcommi, D. steinheimensis, H. tetradactylum, and L. sansaniense (Table...
Concerning complexity, *B. gajense* and *H. tetradactylum* stood out for having higher complexities compared to all other species besides *M. welcommi* and *G. cf. browni*. Moreover, *G. cf. browni* was different from *B. brachypus*, both *Plesiaceratherium* species, and both *Hispanotherium* species regarding complexity and HAsfc81 (Table 3). For both DMTA parameters in which Locality had a noticeable effect (epLsar and Asfc), we found a cluster between Béon 1 and Gračanica opposed to one containing at least Simorre and Sansan (for Asfc: also Kumbi 4 and Villefranche d’Astarac; Table 3).

Table 2: Pairs (Species or Locality) with noticeable p-values after Tukey’s honestly significant difference (HSD) by DMTA parameters.

<table>
<thead>
<tr>
<th>DMTA parameter</th>
<th>Pair (Species or Locality) with differences</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anisotropy</strong></td>
<td><em>Plesiaceratherium mirallesi</em></td>
<td><em>Brachyptotherium brachypus</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Lartetotherium sansaniense</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Hoploaceratherium tetradactylum</em></td>
</tr>
<tr>
<td><strong>Complexity</strong></td>
<td><em>Hoploaceratherium tetradactylum</em></td>
<td><em>Hispanotherium beonense</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Prosantorhinus douvillei</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Plesiaceratherium mirallesi</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Plesiaceratherium balkanicum</em></td>
</tr>
<tr>
<td><strong>Gračanica</strong></td>
<td></td>
<td><em>Sansan</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Simorre</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Villefranche d’Astarac</td>
</tr>
<tr>
<td><strong>HAsfc9</strong></td>
<td><em>Hoploaceratherium tetradactylum</em></td>
<td><em>Plesiaceratherium balkanicum</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Plesiaceratherium mirallesi</em></td>
</tr>
<tr>
<td><strong>HAsfc81</strong></td>
<td><em>Hoploaceratherium tetradactylum</em></td>
<td><em>Plesiaceratherium balkanicum</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Plesiaceratherium mirallesi</em></td>
</tr>
</tbody>
</table>
Table 3: Fischer’s least significant difference (LSD) post hoc results by DMTA parameters

Groups (a, ab, abc, b, bc, and c) are indicated with a p-value threshold of 0.05, for the sake of clarity.

<table>
<thead>
<tr>
<th></th>
<th>a</th>
<th>ab</th>
<th>abc</th>
<th>b</th>
<th>bc</th>
<th>c</th>
</tr>
</thead>
<tbody>
<tr>
<td>epLsar Species</td>
<td>A. simorrense</td>
<td>P. mirallesi</td>
<td>G. cf. browni</td>
<td>H. beonense</td>
<td>H. cf. mtritensis</td>
<td>P. balkanicum</td>
</tr>
<tr>
<td>Locality</td>
<td>Gračanica</td>
<td>Villefranche d’Astarac</td>
<td>Devínska Nová Ves</td>
<td>Kumbi 4</td>
<td>-</td>
<td>Sansan Simorre</td>
</tr>
<tr>
<td>Asfc Species</td>
<td>B. gajense</td>
<td>H. tetradactylum</td>
<td>G. cf. browni</td>
<td>M. welcommi</td>
<td>-</td>
<td>B. brachypus</td>
</tr>
<tr>
<td>Locality</td>
<td>Simorre</td>
<td>Sansan</td>
<td>Kumbi 4</td>
<td>Villefranche d’Astarac</td>
<td>Devínska Nová Ves</td>
<td>-</td>
</tr>
</tbody>
</table>
Besides at Béon 1, the microwear sampling was very restricted (n < 5), either due to low numbers of exploitable teeth available, or to the lack of well-preserved microwear texture on molars. In order to facilitate the understanding, the results are presented by locality (chronologically) and by species. At Kumbi 4, four species were considered for DMTA: *Pleuroceros blanfordi*, *Mesaceratherium welcommi*, *Gaindatherium cf. browni* (grinding only), and *Brachypotherium gajense* (grinding only). Figure 3 shows that Kumbi rhinocerotids display a great variety of microwear patterns. Only one specimen, belonging to *G. cf. browni*, is above the high anisotropy threshold of 5 x 10^-3, while all specimens of *B. gajense* and *G. cf. browni* but none of *P. blanfordi* display values above the high complexity cutpoint of 2. *Gaindatherium cf. browni* and *P. blanfordi* have large variations in anisotropy, from low values (~ 1 x 10^-3) to high (about 5 x 10^-3), but consistent values of complexity (around 3 and 1.4 respectively). Such a pattern associated with moderate (*P. blanfordi*) to high (*G. cf. browsi*) values of HAsfc (Figure 4) point towards mixed-feeding diets, probably with the inclusion of harder objects for *G. cf. browsi*. The signature for *B. gajense* is suggestive of browsing with low mean anisotropy (1.82 x 10^-3), but high means of complexity (4.58), FTfv (7.89 x10^4), and HAsfc (HAsfc9 = 0.36; HAsfc81 = 1). Eventually, *M. welcommi* presents low to moderate anisotropy values (< 4 x 10^-3), a moderate complexity (~ 1.5) and HAsfc, but high FTfv (> 4 x 10^4) on both facets (Figure 4), which denotes browsing or mixed-feeding habits.

At Béon 1, the DMT of the four rhinocerotids overlap contrary to that of Kumbi 4 rhinocerotids (Figure 3). The DMTA results are already detailed in Hullot et al. (2021). They suggest a mixed-feeding behavior for *H. beonense* with moderate anisotropy values (mostly < 4 x 10^-3), variable values of complexity (low-medium), moderate-high FTfv (around 4 x 10^4), and moderate HAsfc on both facets (Figure 4). *Plesiaceratherium mirallesi* is considered as a folivore due to low complexity (~ 1) and HAsfc values but relatively high anisotropy (above 5 x 10^-3), indicating an abrasive but not diversified diet. Concerning the teleoceratines, they display similar microwear textures (Figure 3; Figure 4), though *B. brachypus* has lower values of anisotropy (< 2 x 10^-3). This suggests that *B. brachypus* was probably a browser or a mixed-feeder, while *Pr. douvillei* was a browser favoring leaves.
Figure 3: Dental microwear results of early and middle Miocene rhinocerotids plotted as mean and standard deviation of anisotropy against that of complexity by facet, locality and species.
Localities organized chronologically. Steinheim am Albuch not shown as only one specimen was studied. Color code by species as indicated in the figure.

Figure 4: Dental microwear results of early and middle Miocene rhinocerotids plotted as boxplots of each DMTA parameter by facet and species. Time flows from left to right. DNV: Devínska Nová Ves, St: Steinheim am Albuch, Si: Simorre, V: Villefranche d’Astarac. Color code by species as indicated in the figure and consistent with Figure 3.
At Gračanica, we also observe a great overlapping in the DMT. The complexity is very low for all rhinocerotids studied (mostly below 1) suggesting soft food items. Anisotropy varies greatly but HAsfc81 is consistently low (< 0.5) for all species (Figure 4). This points towards soft browsing or folivory for all rhinocerotids at Gračanica.

At Sansan, the DMTA signatures of the rhinocerotids are more diversified and less overlapping, similarly to Kumbi 4 (Figure 3). Lartetotherium sansaniense and H. tetradactylum have low values of anisotropy (< 2.5 x 10^{-3}) and moderate (1-2; L. sansaniense) to high (> 2; H. tetradactylum) values of complexity, recalling browsers. The high values of HAsfc (Figure 4) for both species are compatible with a browsing diet. The other two species, B. brachypus and A. simorrense, are in the range of mixed-feeders (Figure 3), and have compatible moderate to high values of HAsfc.

At Devínska Nová Ves, our restricted sample suggest browsing habits for both species P. balkanicum and D. steinheimensis, with moderate values of both anisotropy (~ 2.5 x 10^{-3}) and complexity (mostly between 1 and 1.5). FTfv is high on both facets (> 4 x 10^{4}) and HAsfc moderate (Figure 3; Figure 4).

At Simorre, B. brachypus specimens display low values of anisotropy (< 2.5 x 10^{-3} except two specimens), and high values of complexity (> 2) and FTfv (> 4 x 10^{4}) on both facets. Values of HAsfc9 are high (> 0.3) on both facets, while that of HAsfc81 are moderate on the grinding facet (median = 0.45) and high on the shearing one (median = 0.7). These DMTA results suggest browsing preferences with the inclusion of hard objects, probably fruits.

At Villefranche d’Astarac, B. brachypus and A. simorrense present well-distinguished DMT (Figure 3). Brachypotherium brachypus has low anisotropy values (< 2.5 x 10^{-3}) and high complexity ones (> 2.5) corresponding to a browsing signal, while the opposite is true for A. simorrense. The moderate values of HAsfc for A. simorrense suggest that folivory is more likely than mixed-feeding for these specimens and the corresponding individuals.

Eventually the specimen of A. simorrense from Steinheim am Albuch has a moderate anisotropy (Grinding: 3.56 x 10^{-3}; Shearing: 2.27 x 10^{-3}), low (Shearing: 0.41) to moderate (Gringing: 1.6).
complexity, a high FTfv on the grinding facet ($8.35 \times 10^4$) but low on the shearing one ($0.63 \times 10^4$), and low HAsfc on the shearing facet but moderate-high on the grinding one (Figure 4). This pattern is consistent with browsing or mixed-feeding habits.

**GLMM**: For all response variables (epLsar, Asfc, FTfv, HAsfc9, and HAsfc81), model support increased (i.e., lower AIC) when intraspecific factors (e.g., Facet, Genus, Locality) were included. The final models contained three to seven factors, including Specimen, the random factor, by default in all models. Facet was in the final models of epLsar and FTfv, Locality and Age were found in the final models of Asfc and both HAsfc. Details and comparison of all models can be seen in electronic supplementary material S5 and S6. Differences by Locality were also observed. Béon 1 had a lower complexity than Kumbi 4, Sansan, Simorre, and Villefranche ($df = 119, \alpha = 0.05, |t-values| > 1.7$), while Tukey’s contrasts highlighted lower values of Asfc for Gračanica than for Kumbi ($p$-value < 0.004), Simorre ($p$-value = 0.027), and Villefranche ($p$-value < 0.001). Moreover, Béon 1 had lower HAsfc9 and HAsfc81 values than Kumbi 4 and Sansan ($df = 119, \alpha = 0.05, |t-values| > 1.7$). Tukey’s contrasts also showed that Sansan had higher HAsfc9 and HAsfc81 than Gračanica ($p$-value ≤ 0.001).

The sampling site (tooth locus, position, side) had sometimes a confounding effect. For instance, M2 had higher epLsar values than M3 ($df = 119, \alpha = 0.05, t$-value = -1.95).

**GLMM - Comparison to extant dataset**: When compared to the extant dataset (see S8 for all details), we noticed that all fossil species had lower anisotropy values than the extant grazer *Ceratotherium simum* (white rhinoceros) and the folivore *Dicerorhinus sumatrensis* (Sumatran rhinoceros), although the classic t-value threshold was not reached for a few species (*P. blanfordi*, *G. cf. browsi*, *B. gajense* [only regarding *C. simum*], *P. mirallesi*, and *A. simorrense*; $\alpha = 0.95, |t$-values| ≤ 1.7). On the contrary, *P. mirallesi* displayed higher values of anisotropy than the extant browsers *Diceros bicornis* (black rhinoceros; t-value = 1.93) and *Rhinoceros sondaicus* (Javan rhinoceros; t-value = 2.66). Regarding complexity, *C. simum* and *D. sumatrensis* had lower values than *B. gajense* and *H. tetradactylum*, while the extant browsers had higher values than *P. balkanicum*, *P. douvillei*, *P. mirallesi*, and *H. beonense* ($\alpha = 0.95, |t$-values| > 1.7). All other DMTA parameters showed less differences between the extant and fossil datasets: *C. simum* and *R. sondaicus* had higher FTfv, HAsfc9, and HAsfc81 than *B. brachypus*, *P. balkanicum*, and *P. mirallesi* ($\alpha = 0.95, |t$-values| > 1.7).
Hypoplasia

The overall prevalence of hypoplasia on rhinocerotid teeth from the early and middle Miocene localities studied is high, with 302 teeth affected out of 1401, corresponding to over 20 % (21.56 %).

There are, however, marked discrepancies between species, localities, and tooth loci (Figure 5). The most affected genera were Plesiaceratherium (104/357; 29.13 %), Prosantorhinus (97/370; 26.22 %), and Brachypotherium (46/178; 25.84 %), but this resulted mostly from the dominance of Béon 1 specimens in our sample. Brachypotherium brachypus was often one of the most affected species at all sites where the species was found, except Sansan (1/13; 7.69 %), contrary to A. simorrense often found associated with the latter species and relatively spared by hypoplasia (maximum 4/35 = 11.43 % of teeth affected at Simorre; Figure 6).

The prevalence was above 10 % for all localities except Kumbi 4, for which the overall prevalence is low (6/99; 6.06 %; Table 4). Hypoplasia defects are quite rare at Kumbi 4 for all species studied, and even null for the teleoceratine species (D. fayejiangense and B. gajense), Bugtirhinus praecursor, and Plesiaceratherium naricum (Figure 6). Only Pleuroceros blanfordi appears a little more affected (4/42; 9.52 %), totaling four of the six hypoplasias observed at the locality. Hypoplasia was also relatively limited at Sansan (14/132; 10.61 %) and Devinska Nová Ves (5/48; 10.42 %), with only B. brachypus and H. tetradactylum affected at Sansan, and D. steinheimensis from the latter (Table 4 ; Figure 6). On the contrary, the rhinocerotids from Béon 1, Béon 2, and Gračanica are very affected, with more than 25 % of the teeth presenting at least one hypoplasia at Béon 1 (216/832; 25.96 %) and Béon 2 (5/18; 27.78 %), and nearly 50 % at Gračanica (15/31; 48.39 %; Table 4). At these sites, the prevalence of hypoplasia is high for all species but the elasmotheriines (Figure 6). Indeed, the elasmotheriines of all sites were relatively spared (H. beonense at Béon 1: 13.04 %) or even not affected by hypoplasia (B. praecursor at Kumbi 4 and H. cf. matritense at Gračanica).
Figure 5: Number (A) and Frequency (B) of hypoplasia by Locality and Species

Numbers on barplot A indicate the number of hypoplastic teeth (dark colors) versus unaffected ones (light colors). Frequencies are calculated as the ratio of hypoplastic teeth on the total number of teeth (hypoplastic and normal). Sub-tribes colored in blue: Teleoceratina, in green: Rhinocerotina; in yellow: Aceratheriina, in pink: stem Rhinocerotinae, and in orange: Elasmotheriina.

Abbreviations: DNV: Devínska Nová Ves, Sim: Simorre, Vil: Villefranche d’Astarac


**Figure 6**: Prevalence of hypoplasia by locality, species and tooth locus plotted against phylogeny

Phylogenetic relationships follow formal parsimony analyses (Antoine, 2002; Antoine et al., 2010; Becker et al., 2013; Tissier et al., 2020).
Subtribes colored in blue: Teleoceratina, in green: Rhinocerotina; in yellow: Aceratheriina, in pink: stem Rhinocerotinae, and in orange: Elasmotheriina

Dark colors: hypoplastic teeth; Light colors: unaffected teeth

Table 4: Prevalence of hypoplasia by locality (number of specimens/percentages)

<table>
<thead>
<tr>
<th>Locality</th>
<th>Hypoplastic</th>
<th>Normal</th>
<th>Percentage of hypoplasia</th>
</tr>
</thead>
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<tr>
<td>Kumbi 4</td>
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<tr>
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<tr>
<td>Béon 1</td>
<td>616</td>
<td>216</td>
<td>25.96</td>
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<tr>
<td>Gračanica</td>
<td>16</td>
<td>15</td>
<td>48.39</td>
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<tr>
<td>Devínska Nová Ves</td>
<td>43</td>
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<td>10.42</td>
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<tr>
<td>Sansan</td>
<td>118</td>
<td>14</td>
<td>10.61</td>
</tr>
<tr>
<td>Simorre</td>
<td>60</td>
<td>11</td>
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<tr>
<td>Villefranche d’Astarac</td>
<td>111</td>
<td>23</td>
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<tr>
<td>Steinheim</td>
<td>29</td>
<td>7</td>
<td>19.44</td>
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</table>

Concerning the loci, milk teeth (47/294; 15.99 %) were overall less affected by hypoplasia than permanent ones (255/1107; 23.04 %; Table 5). Indeed, besides at Béon 1, very few milk molars are hypoplastic (one D1/d1 at Gračanica and Steinheim, two D1 at Simorre, two D2 and 1 d4 at Sansan, 4 D4/d4 at Villefranche d’Astarac). Upper and lower teeth were equally affected (Kruskal-Wallis, df = 1, p-value = 0.11), with respectively 19.86 % (144/725) and 23.37 % (158/676) of teeth bearing hypoplasia. The most affected locus was the fourth milk molar with 38.24 % (26/68), while the least affected were second and third milk molars with around 4 % affected (3/68 and 3/72 respectively; Table 5). Other loci particularly affected were fourth premolars (60/200; 30 %), third molars (50/188; 26.60 %), and second molars (49/202; 24.26 %; Table 5). Once again, these findings mostly result from the dominance of Béon 1 specimens in the sample, and great differences in the hypoplasia pattern are observed by locality (Figure 6). Indeed, if virtually all tooth loci are likely to be affected for Béon 1 rhinocerotids, the pattern is less varied at other localities although it seemingly diversifies with sample size (e.g., H. tetradactylum from Sansan and Villefranche d’Astarac). For instance, hypoplastic teeth are nearly exclusively molars at Kumbi 4 (with only one defect on a p3), and permanent teeth at Gračanica (only one defect on a D1; Figure 6).
Table 5: Prevalence of hypoplasia by tooth locus, regardless of provenance and taxon

Upper and lower teeth merged as they have a similar timing of development

<table>
<thead>
<tr>
<th>Tooth</th>
<th>Hypoplastic</th>
<th>Normal</th>
<th>Percentage of hypoplasia</th>
</tr>
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<tr>
<td>Total permanent teeth</td>
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GLMM: For all response variables (Hypo, Defect, Multiple, Localization, and Severity), model support increased (i.e., lower AIC) when intraspecific factors (e.g., Tooth Loci, Genus, Locality) were included. When Genus was not forced into the models, the final models contained three to six factors, including Specimen, the random factor, by default in all models. Defect (converted to a factor) was in the final models of all concerned variables (Multiple, Localization, and Severity). Genus was in the final models of all variables but Localization. Position was in the final models of Hypo, Defect, and Localization, while Tooth, and Wear were in that of Hypo and Defect. Details and comparison of all models can be seen in electronic supplementary material S5 and S7.

Based on GLMMs results, we can assess the influence of Genus, Locality, and Tooth on the hypoplasia pattern. Alicornops was less affected than Brachypotherium (p-value = 0.015), while Plesiaceratherium was more prone to hypoplasia than Hispanotherium (p-value = 0.02). GLMMs revealed differences in the patterns of hypoplasia (i.e., type of defects and their frequencies) between Brachypotherium and the following taxa: Dicerorhinus (p-value = 0.0046), Alicornops (p-value < 0.001), and Protaceratherium (p-value = 0.034). Tukey’s contrasts also revealed the lowest p-values between Alicornops and the following taxa: Hispanotherium (p-value = 0.064), Plesiaceratherium (p-value < 0.01), Prosantorhinus (p-value < 0.01), Protaceratherium (p-value = 0.05). Eventually
Dicerorhinus had a different hypoplasia pattern than Plesiaceratherium (p-value = 0.034) and Prosantarhinus (p-value = 0.071).

Concerning tooth loci, all teeth but fourth premolars and third molars are less affected than fourth milk molars (p-values < 0.05). The results further suggested that the most commonly affected loci were third molars, fourth premolars and fourth milk molars, while the least affected were all milk molars but the fourth. Concerning localities, Gračanica teeth were significantly more touched than Béon 1 and Sansan specimens (p-values = 0.01). Middle Miocene rhinocerotids present a hypoplasia pattern distinct from that of early Miocene ones (p-value = 0.019). Similarly to GLMMs for DMTA, we observed confounding effects. Slightly-worn teeth had less hypoplasia than average worn (p-value = 0.005) and very worn teeth (p-value = 0.026).

Discussion

Dietary preferences and niche partitioning of the rhinocerotids studied

The comparison of the fossil specimens DMT to that of extant ones highlighted important differences. This suggests that the dietary spectrum of extinct rhinocerotids might have been very distinct from that observed in the living species (Hullot et al., 2019). However, the microwear textures of the fossils are critically distinct from that of the only extant strict grazer Ceratotherium simum, banning such dietary preferences for the studied fossil specimens. This finding is not surprising as grasses and associated grazing ungulates expended only during latest Miocene in Eurasia (Janis, 2008). The reconstructed dietary preferences based on DMTA are presented in Table 6 by locality and by species.

The DMTA results of fossil specimens on both facets (Figure 3; Figure 4) suggest a clear niche partitioning based on feeding preferences for the rhinocerotid specimens studied at Kumbi 4, Sansan, and Villefranche d’Astarac. Although DMT could only be explored in four out of the nine rhinocerotid species present at Kumbi 4, the patterns observed indicate clear differences in the feeding behaviors, even if leaf consumption seems to be a major component for all rhinocerotids studied but B. gajense.
Table 6: Dietary preferences inferred from textural microwear (DMTA) of the studied rhinocerotid specimens from different fossil localities of the lower and middle Miocene of Eurasia.

Color code: brown/B – browser, blue/M – mixed-feeder, light green/F – folivore, no color/x – not studied

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<th>Béon 2</th>
<th>Béon 1</th>
<th>Gračanica</th>
<th>Sansan</th>
<th>Devínska Nová</th>
<th>Vès Spalte</th>
<th>Simorre</th>
<th>Villefranche d’Astarac</th>
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</table>

This finding is in line with the inferred lush vegetation under warm and moist climatic conditions proposed for this locality, providing abundant and diverse feeding resources for the many large herbivores present (Antoine et al., 2010, 2013; Martin et al., 2011). The dietary preferences reconstructed for Sansan rhinocerotids suggest the co-occurrence of two browsers (L. sansaniense and H. tetractylum, the latter including harder items in its diet) and two mixed-feeders (A. simorrense and B. brachypus), coherent with the (sub-)tropical forested environment reconstructed for that locality (Costeur et al., 2012), but at odds with the recent-like Miocene coolhouse as depicted by Westerhold et al. (2020; ~415 parts per million CO₂). This niche partitioning is probably accentuated by different habitat preferences: H. tetractylum is mostly found in swamp or fluvial sediments.
indicating wet habitat preferences contrary to *A. simorrense*, while *B. brachypus* seems intermediate and *L. sansaniense* generalist (Heissig, 2012). Eventually, we observed obvious differences in the dietary preferences for *A. simorrense* (folivore or mixed-feeder favoring leaves) and *B. brachypus* (browser including hard objects) at Villefranche d’Astarac, where a humid forested environment is hypothesized (Bentaleb et al., 2006).

On the contrary, an overlap of microwear textures, especially for the grinding facet, is observed for the localities of Béon 1 and Gračanica. Besides diet, different habitats and feeding heights might result in niche partitioning (Hutchinson, 1959; Arsenault and Owen-Smith, 2008). Concerning Béon 1, a partial niche partitioning due to habitat differences has been hypothesized for the rhinocerotids – swamps for both teleoceratines *B. brachypus* and *P. douvillei*, open woodland for *P. mirallesii*, and savannah-like open environments for *H. beonense* (Bentaleb et al., 2006) – and subtle dietary differences are discussed in Hullot et al. (2021) in the light of the combination of molar mesowear and dental microwear texture analysis. At Gračanica, 2D microwear and mesowear score already revealed an overlap in the dietary preferences of *Pl. balkanicum* and *B. brachypus* as two browsing species, although the latter is labelled as “dirty browsing”, a dietary category (not defined by comparative datasets on intensively studied extant species with known diets) including species assumed to browse and incorporate soil particles. (Xafis et al., 2020). Although microwear sampling is restricted and includes premolars for the other two rhinocerotids (*L. sansaniense* and *H. cf. matritense*), it points towards different mixed-feeding behaviors, most likely with a dominance of grass in the diet of *H. cf. matritense* (Xafis et al., 2020). The very low values of complexity for all Gračanica rhinocerotids in our sample, combined to relatively high values of anisotropy (Figure 3; Figure 4), could suggest an important consumption of leaves for all species, as well as a very low amount of lignified tissues that would have required more grinding to get access to cell content. Interestingly, the reconstructed environment at this locality (based on mammal assemblage and flora) is a lowland swamp surrounded by a closed canopy-like environment (Butzmann et al., 2020; Xafis et al., 2020), meaning that leaves would have been an abundant resource. This recalls the feeding preferences and microwear textures of the extant *Dicerorhinus sumatrensis* (Sumatran rhino; Hullot et al., 2019). Eventually, the restricted DMTA samples from Devinska Nová Ves Spalte, Steinheim am Albuch, and Simorre suggest browsing...
or mixed-feeding habits for all specimens studied, but did not allow to conclude on potential competition for food resources.

Interactions with co-occurring herbivores

Besides other rhinocerotid species, the individuals studied co-occurred with many other herbivore mammals. Although co-occurrence is not necessarily a good proxy for ecological interactions (Blanchet et al., 2020), it is possible that some of these herbivores were competing for or partitioning food resources with the rhinocerotids. Unfortunately, very little has been studied concerning the dietary preferences of the fauna at most of the studied localities with the notable exceptions of Gračanica and Sansan.

Indeed, recent studies on dental wear (micro- and meso-wear) or stable isotopy, suggested frugivory for some associated species such as tragulids (*Dorcatheirum* spp. at all localities but Simorre and Villefranche; Aiglstorfer et al., 2014; Xafis et al., 2020), the middle Miocene Moschidae (*Micromeryx* spp. found at Sansan, Simorre and Steinheim am Albuch; Aiglstorfer and Semprebon, 2019) or the chalicotherere *Metaschizotherium fraasi* (found at Steinheim am Albuch; Semprebon et al., 2011). Interestingly, no rhinocerotid specimens studied here seemingly favored fruits. Similarly, the lophodont suid *Listriodon splendens*, found at Gračanica, Devínska Nová Ves, and Simorre, might have favored grasses (Van der Made, 2003; Xafis et al., 2020), a resource not exploited by the rhinocerotids either.

Otherwise, the vast majority of herbivore species were probably browsers or mixed-feeders, in good agreement with the statement by Eronen and Rössner (2007) that these forms were dominant between MN4 and MN9. This is for instance the case of the associated perissodactyl species *Anisodon grande* (Chalicotheriidae), which 2D microwear signal at Devínska Nová Ves suggests folivory (Semprebon et al., 2011), and *Anchitherium* spp. (Equidae) ranging from generalists to “dirty browsers” (Kaiser, 2009; Xafis et al., 2020).

Within browsers and mixed-feeders, resources partitioning is still possible (e.g., consumption of different plant parts or species) but might be difficult to detect in fossil communities. Moreover, other strategies can lead to niche partitioning, such as different habitat, different body mass, or different feeding height (Hutchinson, 1959; Schoener, 1974; Arsenault and Owen-Smith, 2008). Regarding body mass, most rhinocerotids studied are megaherbivores *sensu* Owen-Smith (1988; terrestrial...
herbivores weighting more than 1000 kg), which implies specific feeding strategies and metabolic
requirements. Megaherbivores are often treated as a separate herbivore guild, mostly disturbing that
of mesoherbivores (4–450 kg; Fritz et al., 2002; Calandra et al., 2008; Landman et al., 2013). Within
megaherbivores, proboscideans frequently co-occurred with rhinocerotids at the studied localities and
were mostly browsers or mixed-feeders, placing them as direct competitors for rhinocerotids. Indeed,
the mesowear and 2D microwear suggest that Prodeinotherium bavaricum and Gomphotherium
angustidens were browsers at Gračanica (Xafis et al. 2020), while the mesowear angle categorizes P.
bavaricum from Sansan, D. giganteum from Villefranche d’Astarac and G. angustidens from Simorre
as browsers, but G. angustidens from Sansan and Villefranche d’Astarac as mixed-feeders (Loponen,
2020). Such overlapping in the diet of proboscideans and rhinocerotids is observed nowadays
between African elephants and black rhinoceroses (Landman et al., 2013). Interestingly, this
competition is detrimental to the rhinoceros, whose individuals shift towards the inclusion of more
grasses in presence of elephants (on a seasonal basis). Another possibility, as postulated by Xafis et
al. (2020) for Deinotherium spp. and Plesiaceratherium balkanicum at Gračanica, would be different
feeding heights between proboscideans and rhinocerotids, as the first ones were most likely feeding at
the top of trees due to their larger size (some of the biggest Neogene mammals; Larramendi, 2015).

Hypoplasia prevalence and environmental conditions

We found that the hypoplasia prevalence and pattern (i.e., tooth loci affected) were very different
depending on the locality and the species concerned. Except for Kumbi 4, the prevalence was
relatively high (> 10 %) at all sites of our early-middle Miocene sample. Even though nine species of
rhinocerotids are found at Kumbi 4, such a low prevalence is in agreement with previous results in the
region over the Cenozoic (Roohi et al., 2015), and coherent with the very favorable, low-stress context
hypothesized at this locality, that is a rich vegetation under a warm and humid climate (Antoine et al.,
2013).

The prevalence of hypoplasia is high at Béon 1 (>25 %) for all rhinocerotids except H. beonense, with
molars being particularly affected with respect to other dental loci. Second and third molars are the last
teeth to develop and erupt in rhinocerotids (Hitchins, 1978; Hillman-Smith et al., 1986; Böhmer et al.,
2016), and stresses on these late-developed teeth have been correlated with environmental, seasonal
stresses in sheep (Upex and Dobney, 2012). Although subtropical wet conditions are reconstructed at Béon 1 (just prior to the MCO), periodic droughts are also reported in the area at that time (Duranthon et al., 1999; Hullot and Antoine, 2020). Interestingly, the least affected species is the elasmotheriine *H. beonense*, an early representative of a clade adapted to relatively open and arid environments (Cerdeño and Nieto, 1995; Iñigo and Cerdeño, 1997), and which displays a mixed-feeding diet (Figure 4). On the contrary, both teleoceratine species, often considered swamp dwellers, display a high prevalence of hypoplasia (Figure 5).

We found a very high prevalence of hypoplasia at Gračanica, with nearly 50 % of the teeth bearing at least one hypoplastic defect. The proposed age for the locality ranges between 14.8 and 13.8 Ma (Göhlich and Mandic, 2020), which is an interval of great climatic changes. Indeed, though included in the MCO, the interval from 14.7 to 14.5 Ma present an increased seasonality in precipitations, with prolonged dry periods (Böhme, 2003). On the other hand, an abrupt cooling occurred between 14 and 13.5 Ma, correlating with the Mi-3 event (Zachos et al., 2001; Böhme, 2003; Holbourn et al., 2014; mMCT of Westerhold et al., 2020). Besides this challenging environmental context for the rhinocerotids, our DMTA results suggest a potential competition for food resources (Figure 3), that could have generated stressful conditions.

At Sansan, the prevalence of hypoplasia is overall moderate (~ 10 %) and defects are only found in two species out of four: *H. tetradactylum* and *B. brachypus* (only one M1). The pattern of hypoplasia for *H. tetradactylum*, with various loci affected, suggests different stresses and timing, from in utero (D2) to post-weaning (M3). It is quite remarkable, as the proximity of the MCO peak (Maridet and Sen, 2012) leading to seasonal warm and moist conditions (Costeur et al., 2012), would seemingly constitute relatively low stress conditions for the concerned rhinocerotids.

The prevalence of hypoplasia at Devínska Nová Ves Spalte is also moderate (5/48; 10.42 %) and restricted to *D. steinheimensis* (P3, M1, M2 only; Figure 6), although the locality dates from the mMCT. However, despite this transitional climatic system, pollen data from the Vienna Basin, to which the locality belongs, indicate that regional conditions remained tropical with few precipitation variations (Sabol and Kováč, 2006), coherent with the absence of hypoplasia on third molars, that can be
correlated with seasonal stresses. The paleogeographic context seems to have played a major role, as the taxonomic differences with Sansan are partly explained by different paleoenvironments:

Devínska Nová Ves Spalte was a forested area near the shoreline of the transgressive late Langhian sea (Sabol and Kováč, 2006).

The rhinocerotids from the localities of the MN7/8 (Simorre, Villefranche d’Astarac, and Steinheim am Albuch), a time of sea-level drop and comparatively dry climate (Legendre et al., 2005; Böhme et al., 2011; Heissig, 2012; Westerhold et al., 2020), present higher prevalences but contrasted patterns depending on the species and locality (Figure 5; Figure 6). However, contrary to what we could have been expected regarding the environmental conditions, the most affected loci (P2, P3, D1) document mostly early-life stresses (e.g., birth, juvenile disease), rather than environmental or seasonal stresses (Niven et al., 2004; Upex and Dobney, 2012). At Steinheim, only L. sansaniense has hypoplasia on other teeth than second and third premolars, suggesting mostly early life stresses. At Simorre, more loci are affected (D1, P2-P3, P4, and M1) and the pattern is relatively similar for both co-occurring species i.e., B. brachypus and A. simorrense. Hypoplasia on D1, that develops mostly in utero synchronously with D4, could indicate birth-related stresses (Hillman-Smith et al., 1986; Mead, 1999; Böhmer et al., 2016). Similarly, the M1 starts its development relatively early, attested by the presence of a neonatal line in some rhinocerotid teeth (Tafforeau et al., 2007), revealing particularly stressful conditions around birth. Eventually, the rhinocerotids from Villefranche d’Astarac document later-life stresses, with hypoplasia recorded from D4 to M2 (not P2-P3 for A. simorrense). The fourth premolars are particularly affected in B. brachypus, which could indicate harsh weaning or cow-calf separation conditions.

Paleoecologic implications and changes

Several species or genera are retrieved in various localities overtime, but B. brachypus clearly has the longest range (from Béon 1 [MN4] to Simorre + Villefranche d’Astarac [MN7/8], with Gračanica [MN5-6] and Sansan [MN6] in the meantime). We observe a clear shift in the DMT of B. brachypus over time from a mixed-feeding behavior at Béon 1, Gračanica, and Sansan to a clear browsing signal with the ingestion of harder items (fruits, seeds, or even soil) at Simorre and Villefranche d’Astarac (Figure 3; Figure 4). This result could be due to a change in the regional climatic conditions, from warm and
humid pre-MCO to cooler, more seasonal and arid post-MCO (Zachos et al., 2001; Böhme, 2003; Holbourn et al., 2014), perhaps leading to behavioral changes in this species (Cerdeño and Nieto, 1995), and/or changes in local conditions.

Contrastingly, the DMT of A. simorrense remains quite similar from Sansan (MN6) to Steinheim (MN7/8; Figure 3; Figure 4). Interestingly, there are clear differences in the hypoplasia prevalence of these two species, B. brachypus being one of the most affected species in our sample. Such differences in the hypoplasia prevalence could reveal the existence of a competition for food and/or water resources. The pattern of hypoplasia at Simorre (D1, P2-P3) and Villefranche d’Astarac (D4-M1, P4-M2) suggests early life stresses for both rhinocerotids (Figure 6), mostly before weaning (Mead, 1999).

Concerning other species found at more than one locality (L. sansaniense, D. steinheimhensis, and Plesiaceratherium spp.), the hypoplasia patterns seem to be different at each locality, denoting a greater effect of local conditions than species-related sensitivities. Only the Plesiaceratherium species from Béon 1 and Gračanica exhibit comparable patterns (Figure 6), although it could be related to the high prevalence of stresses for individuals belonging to the concerned taxa at these localities. Overall, the elasmotheriines (Bugtirhinus and Hispanotherium) were seemingly spared by hypoplasia. Indeed, no tooth was hypoplastic at Kumbi 4 (B. praecursor; 0/4) and Gračanica (H. cf. matritense; 0/6). At Béon 1, for which a greater sample is available, H. beonense is the least affected species with 13.04 % (12/92) of hypoplastic teeth, nearly exclusively permanent (only one hypoplasia on a D4). If this result was not surprising at Kumbi 4, where low-stress conditions were inferred and very little hypoplasia recorded for all studied species, the difference to other associated species was particularly striking at Béon 1 and Gračanica. The microwear study of elasmotheriines is restricted in the literature, but it suggests the inclusion of a non-negligible part of browse resources in the diet, at least seasonally (Rivals et al., 2020; Xafis et al., 2020). This finding is in line with our DMTA results for Hispanotherium species (Béon 1 and Gračanica) suggesting mixed-feeding preferences (Figure 3; Figure 4). The increasing crown height observed in this clade over time could allow for accommodating to a greater variety of food items (Semprebon and Rivals, 2007; Damuth and Janis, 2011; Tütken et al., 2013), thus limiting nutritional stress, as observed in hipparionine equids, with
respect to anchitherine equids (MacFadden, 1992; Janis, 2008; Mihlbachler et al., 2011). The classic view of elasmotherines as obligate open-environment rhinocerotids adapted to grazing – notably based on representatives from the arid Iberic Peninsula (Iñigo and Cerdeño, 1997) – is thus somehow challenged. This could mean that hypsocodony in this clade might counterbalance significant grit load induced by feeding low in open environments, thus reflecting more the habitat rather than the diet, an hypothesis that has already been proposed to explain hypsocodony evolution (Janis, 1988; Jardine et al., 2012; Semprebon et al., 2019).

Conclusions

The study of the paleoecology of rhinocerotids from the early and middle Miocene of Eurasia revealed clear differences over time and space between or within species. Though, DMTA results suggested only browsers and mixed-feeders (no grazers nor frugivores) in the studied rhinocerotid sample, they unraveled clear niche partitioning through food resources at several diachronous localities (Kumbi 4, Sansan, and Villefranche d’Astarac). At other localities (Béon 1, Gračanica), a significant overlap of microwear textures was observed, and more subtle differences in food preferences and other niche partitioning strategies (habitat, feeding height) may have existed. Regarding enamel hypoplasia, which is quite prevalent in the studied sample (except in the oldest and only South Asian locality, Kumbi 4), it revealed clear disparities between localities, species, and dental loci. While the effects of climate changes were not immediately obvious, we discussed more specific, local conditions that may explain the observed stresses. Regarding a potential phylogenetic effect, we were able to highlight very different sensitivities: while *B. brachypus* is highly affected by hypoplasia regardless of locality and conditions, elasmotheriines (*Bugtirhinus praecursor* at Kumbi 4, *Hispanotherium beonense* at Béon 1 and *H. cf. matritense* at Gračanica) are pretty spared in contrast. Over time and depending on the conditions, differences in DMT and/or prevalence of hypoplasia were observed for some species found in several localities. This is notably the case for the feeding preferences of *B. brachypus*, oscillating between browser and mixed feeder, or for the hypoplasia profiles of *L. sansaniense* and *D. steinheimensis* denoting different stress periods and local conditions.
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