MECHANISMS OF COMPETITION IN MAMMALS DURING THE LATE NEOGENE

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INTRODUCTION

Megaherbivore mammals are traditionally defined as species above 1000 kg (du Toit & Owen Smith, 1989; Fritz et al., 2011). In modern-day ecosystems, this ecological category includes few species such as rhinos, elephants, the hippopotamus, gaur, yak, and the giraffe, and it is geographically restricted to sub-Saharan Africa and Asia. However, before the Late Pleistocene megafauna extinction, megaherbivores were much more diverse. The so-called mammal megafauna was then represented by mastodons, mammoths, ground sloths, giant armadillos, a number of notoungulates, very large bovids, deer, and camels, plus giant kangaroos and other marsupials. Megaherbivores control the consumption of primary production excluding smaller species from accessing limited resources, and exposing them to an increased predation risk by clearing thickets of vegetation the small game usually use to hide from predators. Moreover, in virtue of their large body size, megaherbivores have virtually no predator (Fritz et al., 2011). Consequently, when abundant, megaherbivores also cause a negative effect on predators limiting their availability of prey. This ecological mechanism is known as “apparent competition” and describes the indirect competition for limiting resources. In this work, we verified the idea that apparent competition of megaherbivores on carnivores applies to Neogene large mammals. Carnivores were a factor in controlling the diversity of small prey.

MATERIALS AND METHODS

We downloaded from the Paleobiology Database, NOW Database and Pangaea Database, Neogene fossil occurrences of mammals belonging to Artiodactyla, Perissodactyla, Proboscidea, Carnivora and Creodonta. The geographical range of mammal occurrences covers the entire Eurasian region plus the Africa continent. We supplemented these data with occurrence records from Raia et al. (2009) and other published sources. For each species, age estimates and geographical paleocoordinates of individual occurrences at fossil sites were recorded. Overall, the occurrences dataset includes 655 artiodactyls, 114 carnivores, 4 creodonts, 163 perissodactyls, 67 proboscideans, for a total of 1003 species spanning from the earliest Miocene to the Holocene, distributed over 3021 fossil localities. For each species, we compiled body size estimates from different databases. Species were divided in four ecological categories based on both diet and estimated body mass: Megaherbivores (Mega), Herbivores...
(Herb), Carnivores (Carn) and Sabertooths (Sab).

Then, we divided all record in 2-million-years-long temporal intervals maintaining a reasonably dense record for each interval. For each species and within each time bin, we constructed minimum convex polygons (MCP, Carotenuto et al., 2010) starting from its fossil occurrences (Fig. 1). We overlaid a 500 × 500 km grid cell resolution on each projected continent, sampling regions in an equal area context.

The use of polygons overcomes problems generated by sampling inequality per species and geographic area, by adding cells to the species presence where no fossil occurrence is indeed present, but still within the minimal range of species geographic extent. Similarly, by using a geographic grid all the fossil sites are collapsed in a single faunal list. In this way, the effect of unequal sampling and taphonomic effects across fossil sites is avoided. We performed all analyses twice, both by using the raw fossil occurrence (henceforth “occurrence record”), and by using MCP polygons to attribute species to cells (henceforth “polygon record”). At this stage, for each cell and time bin, we excluded cells with < 5 species overall, or lacking any predator or prey, either.

After that, we computed the ratio between the number of “predators” (Carnivores plus Sabertooths) and “preys” (Meso- and Megaherbivores) for each geographic cell within a given time bin. We also computed the body size range of Predators and Prey for each cell within each time bin in order to retrieve the degree of overlap between Predators and Prey body mass distributions (PPR-overlap).

With these variables, we performed six different regressions using the number of species in each category and their estimated body sizes per cell, and separately per time bin. The regression (1) of predator-to-prey ratio (PPR) against the number of megaherbivores (Mega) was calculated to test for the effect of apparent competition of the latter on Predators. Predators (Pr) were regressed (2) against Herbivores (Herb) to verify for the relationship between the richness of predators and non-megaherbivore prey. The regression (3) between Mega and Herb was computed to test for competition between species belonging to these categories, to test the idea that megaherbivores did control the diversity of Herb. The richness of Sabertooths (Sab) was regressed (4) against Mega in order to test the idea that sabertooths preferentially preyed upon megaherbivores. Similarly, the diversity of large carnivores (Pr > 100) was regressed against Mega (5). Eventually, we regressed (6) PPR against Overlap in order to verify if an increment in degree of overlap is correlated to a higher chance of predation on megaherbivores, under the observation that larger predator might tackle down comparatively larger prey (Van Valkenburgh et al., 2016).

All of the six regressions were controlled for spatial autocorrelation by using GLS models. In details, we fitted the related empirical semivariograms with 4 models (Gaussian, Spherical, Rational Quadratic, Exponential) and then updated an OLS regression by these 4 spatial correlation structures. The outcomes of these five models (OLS and the 4 spatially structured) were then compared by means of ANOVA.

Consecutive intervals within the same geographical place share a number of species. This means that the data could be temporally autocorrelated, thereby originating spurious associations between the variables. To address the issue of temporal autocorrelation, we used the autoregressive integrated moving average (ARIMA) model. ARIMA works by regressing a variable point value on previous (older) datapoints, at some (fitted) distance (i.e. lag). The best lag between variables was estimated via cross-correlation, and the existence (and removal thereof) of temporal autocorrelation was assessed by means of Breush-Godfrey test (McMurry & Politis, 2015). To produce ARIMA models, we used the polygons record to maximize the

![Fig. 1. The geographical distribution of fossil localities included in the analyses (left). Geographical coordinates were rotated to the present for plotting purposes. To the right, for each species it is plotted the minimum convex polygon including all species occurrences (right).](image-url)
number of datapoints. However, rather than using the 500 km wide cells we opted for 2000 × 2000 km cells and selected only cells possessing at least 10 species and at least one carnivore species per time bin, for at least 6 time bins. We performed the ARIMA regressions of PPR against Mega, Pr against Herb, Mega against Herb, and Pr > 100 against Mega.

RESULTS AND DISCUSSIONS
The results were qualitatively very similar using either the polygon and the occurrence record. The polygon record is much more dense and less affected by sampling issues, and was therefore used to perform ARIMAs.

As far as the six regression are concerned, the predator to prey ratio (PPR) is significantly and negatively correlated to the number of megaherbivores in one fourth of the intervals. No positive relationship applies. The number of predators (Pr) is negatively and significantly related to the number of mesoherbivores (Herb) in 6 out of 8 intervals (75%) (Tab. 1). Nearly two-thirds of the times (5 times in 8 intervals, 62.5%) the richness of megaherbivores (Mega) is inversely and significantly related to herbivore (Tab. 1). There is no positive relationship between the diversity of sabertooths (Sab) and Mega. The richness of large predators (species > 100 kg in body size) is not significant associated to the number of megaherbivores (Tab. 1). The regression between Mega and Pr > 100 is significant and negative. Finally, the degree of overlap between the body size distribution of predators and prey is positively and significantly related to PPR six times (66.7%) (Tab. 1). A significant and negative relationship occurs once (Tab. 1). As regards the first (most recent) interval, all the regression results are consistent with the apparent competition theory. The spatial distribution of PPR in the Old World during the last interval shows no significant spatial autoregression. The results of ARIMA regressions confirm the existence of a negative relationship between the number of predators (Pr) and mesoherbivores (Herb) in four out of five valid cells. The number of megaherbivores is negatively associated to the number of mesoherbivores (Mega-Herb) in three cells out of four. PPR is always negatively associated to Mega. Finally, the number of megacarnivores (i.e. predators above 100 kg in body mass) is negatively associated to the number of megaherbivores twice, and positively associated as many times. Cross-correlation applied to the residuals consistently shows a lag of 1 to 3 intervals, and mostly of 2 (i.e. 4 Myr) between herbivores and predators and megaherbivores respectively, meaning that Pr and Mega negatively affected the diversity of mesoherbivores in successive temporal intervals.

These results indicate that large carnivores paid a price to the ecological dominance of megaherbivores. As megaherbivores diversified, the biomass available to carnivores reduced, exactly because megaherbivores are hard to kill, and smaller herbivores were significantly outcompeted by the larger species. Indeed, rather than the apparent competition of megaherbivores on carnivores, the most robust generalizations we derive from this study is that the diversity of mesoherbivores is negatively affected by the diversity of both carnivores and megaherbivores. This implies that predation and direct competition increases extinction probability in small herbivores, as often suggested to occur for living prey species (Fritz et al., 2002; Malhi et al., 2016). Some sabertooths were probably able to dispatch juveniles of very large prey species such as mammoths or ground sloths. This was suggested to exercise a top-down control on megaherbivores (Van Valkenburgh et al., 2016). While killing juvenile megaherbivore is perfectly feasible for a top predator, and not contradicted by our data, we suggest these were probably not their most profitable prey, and more importantly, megaherbivores diversity did not sustain higher carnivore diversity.

CONCLUSIONS
We found evidence that, in the long run, such pervasive dominance of megaherbivores translates into higher extinction rate upon (primarily) small prey and (secondarily) predatory species, thereby altering predator to prey ratios (Raia et al., 2007; Meloro & Clauss, 2012). Such fact is further conceivable considering that small prey populations tend to be limited by predation, while larger species are mainly controlled by resources (Terborgh et al., 2010). It must be noted that very large predators and sabertooths were much less influenced by the diversity of megaherbivores than mesoherbivores. Together with the quite consistent, positive relationship between body size overlap and PPR, this indicates that very large carnivores did actually go for larger prey than other carnivores on average. However, this is also consistent with their larger average body size, and not just with any preference for megaherbivores. Indeed, large predators do not specialize on the larger prey, they just exploit a wider prey spectrum (Radloff & du Toit, 2004).

<table>
<thead>
<tr>
<th>Regression model (y-x)</th>
<th>Significant Positive</th>
<th>Non-significant</th>
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</thead>
<tbody>
<tr>
<td>PPR-Mega</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Pr-Herb</td>
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<td>6</td>
</tr>
<tr>
<td>Mega-Herb</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Sab-Mega</td>
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<tr>
<td>Pr &gt; 100-Mega</td>
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<td>1</td>
</tr>
<tr>
<td>PPR-Overlap</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

Tab. 1. Summary statistics for the regressions of the number of species within ecological categories per cell.
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REFERENCES


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