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Evolutionary patterns of cat-like carnivorans unveil drivers of the sabertooth morphology

Highlights

- There is a continuous spectrum of cat-like phenotypes with sporadic convergence
- Saber-toothed taxa show reduced integration and high early evolutionary rates
- Disparity is high among sabertooths, yet cat-like carnivorans decline after the Miocene
- The emergence of large sabertooths appears to act as a macroevolutionary ratchet

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In brief

Chatar et al. explore the tempo and mode of cat-like mammal evolution, highlighting a shape continuum from small cats to extreme sabertooths. They observe the highest disparities and evolutionary rates in clades with the most extreme sabertooth morphologies adaptations. These morphologies arose in clades with more plastic, less integrated skulls.

Article

Evolutionary patterns of cat-like carnivorans unveil drivers of the sabertooth morphology

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SUMMARY

The sabertooth morphology stands as a classic case of convergence, manifesting recurrently across various vertebrate groups, prominently within two carnivorans clades: felids and nimravids. Nonetheless, the evolutionary mechanisms driving these recurring phenotypes remain insufficiently understood, lacking a robust phylogenetic and spatiotemporal framework. We reconstruct the tempo and mode of craniomandibular evolution of Felidae and Nimravidae and evaluate the strength of the dichotomy between conical and saber-toothed species, as well as within saber-toothed morphotypes. To do so, we investigate morphological variation, convergence, phenotypic integration, and evolutionary rates, employing a comprehensive dataset of nearly 200 3D models encompassing mandibles and crania from both extinct and extant feline-like carnivorans, spanning their entire evolutionary timeline. Our results reject the hypothesis of a distinctive sabertooth morphology, revealing instead a continuous spectrum of feline-like phenotypes in both the cranium and mandible, with sporadic instances of unequivocal convergence. Disparity peaked at the end of the Miocene and is usually higher in clades containing taxa with extreme sabertoothed adaptations. We show that taxa with saberteeth exhibit a lower degree of craniomandibular integration, allowing to exhibit a greater range of phenotypes. Those same groups usually show a burst of morphological evolutionary rate at the beginning of their evolutionary history. Consequently, we propose that a reduced degree of integration coupled with rapid evolutionary rates emerge as key components in the development of a sabertooth morphology in multiple clades.

INTRODUCTION

Felidae, a highly diverse carnivoran family, emerged during the Oligocene in Eurasia and quickly spread worldwide during the Miocene.¹ Although Felidae represents the sole extant cat-like clade, the sabertoothed morphotype is a textbook example of iterative evolution (*sensu*^{2,3}), having evolved many times among placentals, marsupials, and gorgonopsians.^{4–8} Nimravidae is a well-diversified yet fully extinct carnivoran clade that extensively explored a range of cat-like morphologies,⁵ and although they have been commonly called “false cats” or “false sabertooth cats” in opposition to the “true cats,” they evolved cat-like morphologies, including both saber-toothed and non-saber-toothed forms, tens of millions of years prior to the appearance of felids. Saber-toothed taxa are thought to feed on larger prey than similar-sized conical-toothed cats,^{6,9–12} and a clear dichotomy between the craniomandibular shape of saber- and non-saber-toothed forms was expected based on this assumption. However, recent studies highlight that the functioning of the cat-like mandible^{13,14} or even the shape¹⁵ and functioning of the cat-

like cranium¹⁶ are more complex than previously expected. Even the assumption that saber-toothed taxa hunted significantly larger prey has been challenged, with some authors suggesting predation on species within a size range similar to those targeted by present-day carnivorans.¹⁷ One possible implication of this emerging continuum of shape and function across saber-toothed and conical-toothed cats is that niche overlap between those two morphotypes might be stronger than previously thought, as first noted in Werdelin et al.¹⁸

Although previous studies already used morphometry to quantify the cranial or mandibular shape variations within cat-like carnivorans^{15,19–23}, little attention has been given to the spatiotemporal component of such variations, and the phylogenetic context is often overlooked; previous studies usually focused either on the cranium or the mandible. This hinders a comprehensive understanding of the cat-like feeding apparatus, leaving key questions unanswered: is there a clear dichotomy between morphotypes/clades and how does craniomandibular shape change over time and space and across clades? Can we identify times, clades, or places characterized by a particularly high or low disparity?

To answer these questions, we reconstruct the tempo and mode of craniomandibular evolution within Felidae and Nimravidae within a contemporary phylogenetic framework. We gathered a large-scale three-dimensional (3D) morphometric dataset (189 3D models) spanning the entire evolutionary histories of both clades, covering a total 36 million years of evolution (Data S1). Our hypothesis posits that cat-like carnivorans have been “shoehorned” into various categories/morphotypes over the years, while their craniomandibular shape exhibits a nuanced disparity beyond prior conceptions.

To test this hypothesis, we adopt two distinct methodological approaches. Initially, we employ a qualitative approach grounded in multivariate morphospaces. Then, we started to explore the tempo by depicting the disparity across time and space in the different clades. Subsequently, we transition to a quantitative approach, employing a battery of widely used statistical tests to characterize the evolutionary dynamics governing morphological evolution. Specifically, we evaluate the tempo of evolution by quantifying evolutionary rates, while exploring the mode of evolution involves assessing various factors such as the impact of phylogeny (phylogenetic signal) on craniomandibular morphological variation, the presence of constraints shaping evolutionary trajectories (phenotypic integration), and the occurrence of convergent evolution among taxa with similar phenotypes across different clades (C metrics²⁴ and theta tests²⁵) as understanding the evolutionary context within which morphologies arise also requires knowledge of those constraints. Indeed, morphological modules (e.g., braincase, face, ascending ramus, mandibular corpus) evolving independently can lead to greater phenotypic disparity, whereas more integrated structures tend to constrain disparity.^{26–33} Nevertheless, highly integrated structures can also lead to a substantial disparity through amplification of extreme phenotypes when evolving hand in hand in a multidimensional space.³⁴ Therefore, we analyzed the strength of modularity both within and across the cranium and the mandible in felids and nimravids as well as their rate of phenotypic evolution per branch. The hypothesis we test here is that conical and sabertooth clades have distinct patterns of integration.

RESULTS

Morphospaces occupation, convergence, and global disparity

Global morphospace occupation of cat-like carnivorans (Figure 1; see Data S1A and S1D for the list of specimens used in those analyses; Figure S1 for the landmarks placed; Figure S2 for annotated morphospace with taxa names and back transform morphospace) do not show a clear subdivision between sabertooth and non-sabertooth taxa, although our cluster dendrograms retrieved a marked dichotomy separating both extreme morphotypes by a quite long branch length, even retrieving significant p values between the two groups ($p = 0.000999$ for both the skull and the mandible datasets) (Figure S3 for tanglegrams comparing phylogeny and cluster based on 3D data). Instead, for the mandibles (Figure 1A), morphospace occupation shows a clear continuum of shape from the small felines (*Felis* spp., *Prionailurus* spp., *Leopardus* spp., *Caracal* spp., etc.), with a curved mandibular body, short diastema, and a high coronoid process projected dorsally, to the most extreme saber-

toothed taxa (*Barbourofelis* spp.), with a short and straight coronoid process, long diastema, and the development of a mandibular flange. The peak showing the densest occupation in the morphospace is shared between middle- and large-sized felines (*Panthera* spp., *Lynx* spp., *Pseudaelurus* spp., etc.), metailurins (*Yoshi* spp., *Metailurus* spp.), and early machairodontines (*Paramachairodus* spp., *Promegantereon ogygia*, and *Machairodus aphanistus*), seemingly reflecting the most recurring shape in the evolution of cat-like carnivorans.

This is also the case when looking at cranium shape (Figure 1B), with a continuum of shape in between middle- and large-sized felines, plus extreme saber-toothed taxa. However, in this case, small felines form a second peak of recurrent morphology separated from their larger counterparts by a valley of uncommon phenotypes. In this region of the morphospace, which is less densely populated, we still encounter the cheetah, *Acinonyx jubatus*, alongside certain fossil taxa, including *Machairodus aphanistus*, *Yoshi minor*, *Yoshi garevski*, and *Paramachairodus maximiliani*. Some small saber-toothed taxa (e.g., *Yoshi minor*) are quite close to that peak. Extreme saber-toothed taxa are more scattered in our morphospace as they do not form a peak of recurrent morphology. Although the most derived nimravids occupied regions in both morphospace that were never reached by felids (high PC1 on both Figures 1A and 1B), felids also have their own morphological innovation. Indeed, a small feline morphotype (taxa weighing only a few kilograms: *Felis* spp., *Prionailurus* spp., *Leopardus* spp., etc. with a really rounded skull) never occurred in nimravids or even in machairodontine felids. Allometry is strong enough in the dataset so that we retrieved a significant p value in our procD.lm test for the mandible (0.001) but is less marked in the cranium, resulting in a non-significant p value (0.086).

To test the strength of craniomandibular convergence, we used two different metrics designed to detect the occurrence of retained and/or evolved similarity. Stayton’s C1 is a distance-based measure quantifying how far two or more taxa have evolved to be more similar to one another than their ancestors were to each other, ranging from 0 (no convergence) to 1 (complete convergence leading to the same morphology).²⁴ The θ angle, on the other hand, represents the angle between phenotypic vectors and span from 0° (very similar phenotypes) to 180° (very dissimilar phenotype).²⁵ Although C metrics are found significant only in the case of convergence, Castiglione θ can be found significant both in the presence of convergence and conservatism; combining those two metrics in the same study therefore allows researchers to untangle those two evolutionary trends. More details about those convergence metrics are available in the STAR Methods section. Surprisingly, extremely few tested pairs were found to be significantly convergent according to the metrics used (Tables 1 and S3; Figures S2G and S2H). None of the tested pairs was found significant for both Stayton C1 and Castiglione θ questioning the potential convergence between those pairs. Also, for the three taxa pairs providing significant results, only one of the structures (either the cranium or the mandible) showed some degree of convergence. *Yoshi minor* and *Acinonyx jubatus*, although showing striking resemblances in the cranial shape (Figures 1B and S2H), were convergent only regarding their mandibular shape according to the Stayton C metrics. Stayton metrics

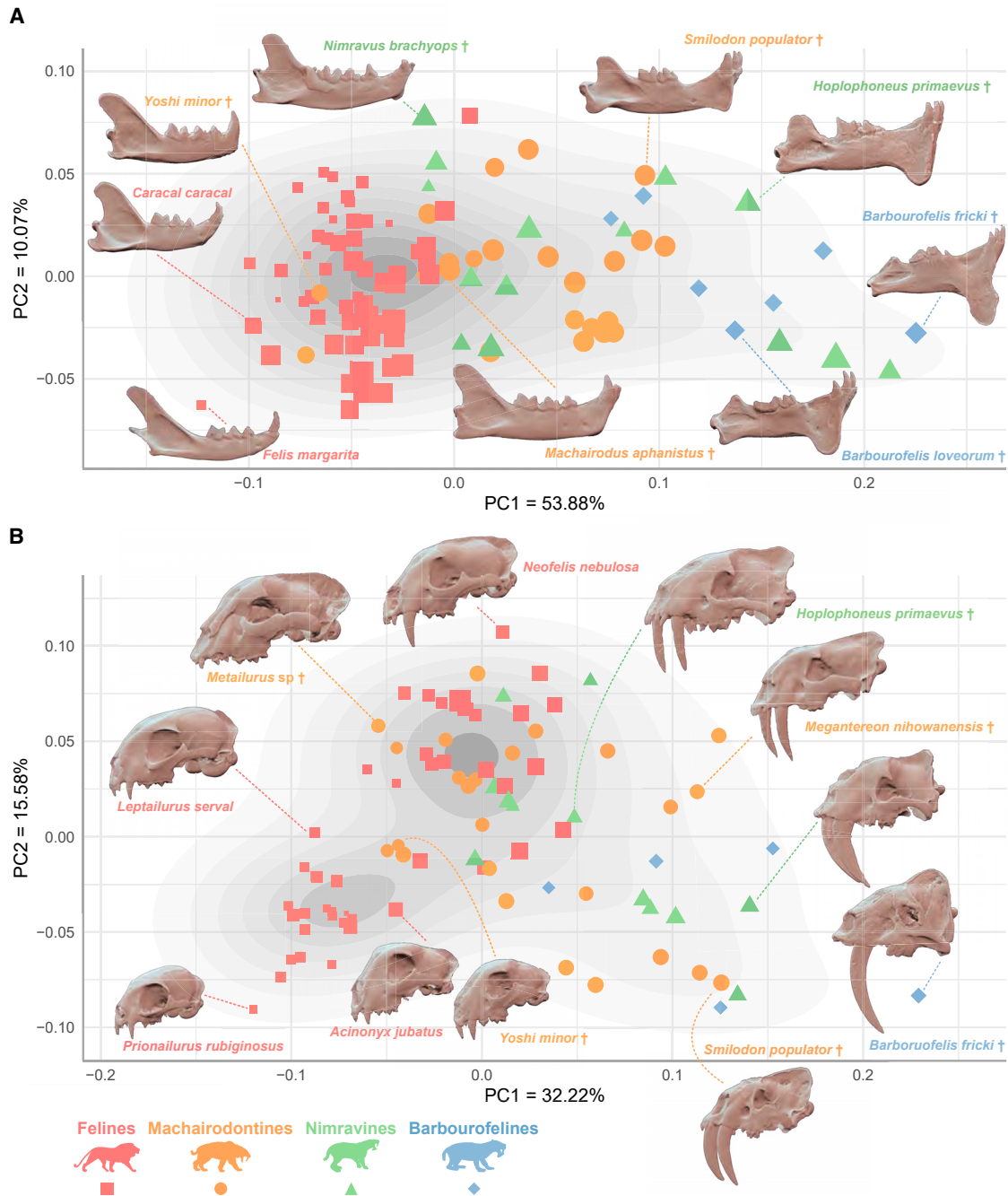


Figure 1. Morphospace occupation

Morphospace occupation of cat-like carnivorans for the mandible (A) and the skull (B). Point size was set relative to the log of the centroid size. See Figure S2 for backtransform shapes and annotated morphospaces with species names. Gray shading represents morphospace occupation density, and every point represents a specimen from Data S1A and S1B.

also retrieved the pair *Megantereon nihowanensis* and *Eusmilus adelos* to be significantly convergent, while Castiglione θ highlighted convergence or conservatism in the cranium of *Neofelis nebulosa* and *Nimravus brachyops* as well as the mandible of *Neofelis nebulosa* and *Metailurus major*. Although the clouded leopard was sometimes presented as an example of convergence^{35–37} with machairodontine felids, it was never compared

with nimravids. However, it is clear from Figures 1A and S9A that in this case the clouded leopard does not particularly show a saber-tooth-like craniomandibular shape, whereas *Nimravus brachyops* and *Metailurus major* exhibit an incredibly feline-like craniomandibular shape, falling close to extant felines in the morphospace, very close to the oldest felid, *Proailurus lemanensis*. Also, since only the Castiglione θ was found to be

Table 1. Results of the convergence tests using the Stayton C1 and Castiglione θ

Pairs	Bone	Stayton ²⁴		Castiglione et al. ²⁵			
		C1 Stayton	<i>p</i> value C1	Theta (θ)	<i>p</i> value θ	Time corr. θ	<i>p</i> value T corr. θ
<i>Yoshi minor</i> - <i>Acinonyx jubatus</i>	cranium	0.011	0.530	11.211	0.709	0.162	0.711
	mandible	0.398 ^a	0.029 ^a	4.881	0.176	0.060	0.088
<i>Hoplophoneus primaevus</i> - <i>Homotherium serum</i>	cranium	0	1	9.395	0.475	0.118	0.351
	mandible	0.037	0.469	6.19	0.112	0.149	0.412
<i>Machairodus aphanistus</i> - <i>Panthera pardus</i>	cranium	0	1	8.118	0.320	0.111	0.292
	mandible	0.033	0.461	6.198	0.382	0.149	0.382
<i>Megantereon nihowanensis</i> - <i>Eusmilus adelos</i>	cranium	0.342 ^a	0.03 ^a	8.007	0.320	0.098	0.177
<i>Megantereon cultridens</i> - <i>Eusmilus sicarius</i>	mandible	0.158	0.192	9.848	0.791	0.214	0.686
<i>Neofelis nebulosa</i> - <i>Nimravus brachyops</i>	cranium	0.001	0.611	7.329	0.205	0.087 ^a	0.050 ^a
	mandible	0.034	0.535	6.882	0.513	0.274	0.513
<i>Neofelis nebulosa</i> - <i>Metailurus major</i>	cranium	0.037	0.383	7.749	0.252	0.112	0.209
	mandible	0.247	0.102	4.987	0.295	0.0087 ^a	0.049 ^a
<i>Machairodus aphanistus</i> - <i>Nimravus brachyops</i>	cranium	0	1	10.004	0.581	0.131	0.482
	mandible	0	1	6.460	0.408	0.155	0.418

C2–C4 can be found in [Table S3](#); see [Figures S2G](#) and [S2H](#) for a more visual interpretation.

^aSignificant *p* values.

significant in this case while Stayton C1 was not, the resemblance between *Neofelis nebulosa* and *Nimravus brachyops* is likely the result of conservatism.

The highest disparity is observed in nimravines for the mandible and machairodontine felids for the cranium, being somewhat comparable to that of barbourofelines ([Figure S4](#)). The disparity is globally greater in nimravids than in felids. This is unsurprising considering the co-existence of extreme sabertooth taxa such as *Barbourofelis* spp. with more feline-like taxa such as *Eofelis* spp. in this group.

Spatiotemporal patterns

During the Paleogene, nimravids already exhibit a wide continuum of cranial shapes, spanning from the average cat-like shapes to some more extreme saber-toothed forms ([Figures 3](#) and [S5](#)), while their mandibular shape appears more far from the peak of recurring morphology ([Figure 3](#)). From the Miocene to the Pliocene, there is a wide distribution of specimens across our morphospace. However, in the Pleistocene, there was a pronounced divergence between machairodontine felids and felines sensu lato in terms of mandibular shape, with machairodontine felids consistently occupying more extreme regions of the morphospace, a pattern that is not retrieved for the cranial morphology. This differentiation is particularly noticeable in North America, where Pleistocene machairodontine (such as *Smilodon* spp., *Homotherium* spp., and *Xenosmilus hodsonae*) exhibit cranial shapes relatively similar to large cats in our dataset but display more extreme mandibular shapes ([Figures 3](#) and [S5](#)). Cat-like carnivorans reached the area of recurring mandibular morphology during the Miocene and more precisely at the end of this Epoch ([Figures 2](#), [S4C](#), and [S4D](#); [Figure 3](#) for the occupation of morphospace through time and space for the mandible; [Figure S5](#) for the cranium). From this Epoch, some taxa (i.e., extreme sabertooths and small felines) explored some unusual areas in the morphospace, but the area of recurring

shape was always occupied ([Figures 2](#), [3](#), [S4](#), and [S5](#)). Although similar regions of the morphospace were occupied by taxa from separate clades during the Miocene and Pliocene (i.e., felines and primitive machairodontines/metailurines), those morphologically similar taxa were usually quite well distributed worldwide, although the Miocene in Asia and North America saw highly convergent felines and machairodontines: *Yoshi minor*/*Felis* sp. Indet in Asia and *Machairodus catacopis*/*Pseudaelurus* spp. in North America. However, the size differences within those co-existing taxa pairs during the Miocene probably facilitated cohabitation just like in extant taxa, such as leopards and tigers in Asia.^{38,39} The peak of disparity in cat-like carnivorans is observed in the Late Miocene with the co-existence of barbourofelines and felids. The pattern of evolution of disparity through time is similar for both the cranium and mandible with a peak during the Miocene and a slow decline ever since. Although the disparity appears particularly low prior to the Miocene, this might be induced by a sampling bias despite bootstrapping, since a different pattern is visible in phylogenetically informed disparity ([Figures S4E–S4I](#)). Please note that our dataset currently exhibits bias toward European and North American specimens, and further sampling is necessary to comprehensively grasp the spatiotemporal trends in craniomandibular shape among cat-like carnivorans. Specifically, more specimens from Asia and Africa are needed to provide a deeper understanding, and we encourage future studies along these lines.

Patterns of phenotypic integration, rates of evolution, and phylogenetic signal

Overall, there is a significant phylogenetic signal in almost all groups tested, except for the cranium of nimravids, nimravines, or in the cranium and mandible of barbourofelines ([Table S4](#)). The high *K* values emerging from those tests might suggest that the low sample size in nimravines and barbourofelines potentially plays in reducing statistical power.



(legend on next page)

Also, our tanglegrams showing the results of our clustering analyses on the Procrustes coordinates against the phylogeny from the super tree (Figure S3) show no correspondences between phylogenetic relationships and craniomandibular shape in cat-like carnivorans unlike what is observed in the skull of extant carnivorans where some correspondences are still present in the cranium (see Law et al.⁴⁰), which suggests a high degree of homoplasy in the dataset. This is consistent with the continuum of shape described earlier. Our analyses retrieved different patterns of morphological integration between the different cat-like clades (Figure 4). Overall, the integration between the cranium and the mandible is present in all groups except machairodontines. Felinae shows the highest degree of morphological integration between the tested structures. Almost all clades show a significant phenotypic integration between the anterior and posterior portion of each structure except barbourufelines for which the mandible does not show any significant integration between its corpus and ramus. Although felines show a high degree of integration between the anterior portion of the cranium and the anterior portion of the mandible, as well as the posterior section of the cranium and posterior section of the mandible, those covariations are not found in any of the other groups. In felines, the anterior portion of the mandible even covaries with the posterior portion of the cranium, and this is the only group in our dataset showing this degree of integration. Still, due to the relatively low sample size for certain clades, particularly nimravines and barbourufelines, these results should be interpreted with caution. They should be viewed as an initial exploratory step toward understanding patterns of phenotypic integration in cat-like carnivorans.

Our branch-specific rate of morphological evolution (Figure 5 for the mandible; Figure S6 for the cranium) highlighted a rapid burst at the beginning of the nimravid evolutionary history. Some lineages of small felines (*Prionailurus* spp.) also show a higher rate of morphological evolution than their larger counterparts, and this pattern is more marked for the cranium (Figure S6) than for the mandible (Figure 5). There are also many acceleration phases during the evolutionary history of machairodontine felids, especially in branches reaching *Smilodon* spp. and *Homotheirus* spp. Except for some small taxa, as mentioned above, the rate is relatively stable and lower in felines. Branch-specific rates of morphological evolution remain relatively higher in nimravid compared with felids.

DISCUSSION

A continuum of shapes in cat-like carnivorans

Cat-like carnivorans have historically been classified using various dichotomies, such as true vs. false cats,^{41–43} scimitar vs. dirk-toothed cats,^{10,16,44,45} saber-toothed vs. non-saber-toothed cats,^{6,46} primitive vs. derived sabertooth,^{47,48} among others. However, our comprehensive morphospace analysis reveals a broad continuum of shapes (Figures 1, S2, and S3). This observation aligns with previous morphometric studies that have shown

that cat-like carnivorans evolved relatively conservative morphologies within distinct clades.^{15,19–21} Furthermore, recent research has highlighted the unexpected functional complexity within these taxa.^{13,14,16} Some taxa considered as “primitive” machairodontine felids (e.g., *Paramachairodus* spp., *Promeganteron ogygia*, and *Machairodus aphanistus*), metailurins (*Yoshi* spp. and *Metailurus* spp.), or nimravids (e.g., *Eofelis*), or even some “derived” scimitar-toothed, exhibit cranial shapes remarkably similar to those of pantherins and other large-bodied felines, although more differences can be found in the mandible (Figure 1). While it has been argued that *Paramachairodus* spp., *Promeganteron ogygia*, and *Machairodus aphanistus* exhibit clear machairodontinae characters^{48–50} and that those primitive taxa must have used the same killing bite as their more derived counterparts,^{11,51} their craniomandibular shapes (Figures 1, S2, S3, and S5) and the functioning of their mandible¹⁴ suggest that they likely employed a different killing bite compared with *Smilodon* or even *Homotheirus*. Our understanding of both the killing bite and hunting strategies employed by extinct cats remains limited. Throat and/or muzzle clamping in extant big cats would have been exceedingly challenging to predict using bone morphology alone. Furthermore, the concepts of the killing bite and hunting strategy represent distinct facets of prey acquisition, potentially evolving independently. Thus, understanding the precise methods by which extinct saber-toothed cats acquired their prey remains an ongoing challenge, likely without a definitive resolution.

Furthermore, nimravids, often labeled as false cats or sometimes “paleofelid” appeared long before felids^{52,53} and exhibited a greater range of morphological diversity (Figures 3, 4, and S5) spanning nearly every region of the cat-like morphospace occupation, except for the small feline cranial morphotype (Figure 1B). Notably, nimravids produced some of the most extreme craniomandibular phenotypes among saber-toothed carnivorans, exemplified by the genus *Barbourufelis* (Figure 1). The distinction between false cats and true cats can be traced back to the earliest descriptions of nimravid remains by Cope, who extensively noted the similarities between nimravids and felids and even remarked, “[...] As nothing but the characters of the canine teeth distinguished these from the typical feline [...]”⁴² However, subsequent authors argued that nimravids constituted a distinct clade⁵⁴ relegating this group to the status of false cats, despite being truly the first cats to have appeared on Earth.

In conclusion, our analyses reveal a continuous spectrum of shape variation, where primitive machairodontine felids, metailurins, primitive nimravids, and certain derived scimitar-toothed cats closely resemble pantherins and other large-bodied felines in terms of cranial shape. However, notable differences emerge in the mandible, underlining the intricate disparity within this group.

Convergence and competition in cat-like carnivorans

The results of our convergence tests (Tables 1 and S3; Figures S2G and S2H) did not provide statistical evidence for convergence, in contrast to some previous studies that found

Figure 2. Disparity through time

Disparity through time computed as a sum of ranges in (A) the cat-like mandible and (B) the cat-like cranium, with occupation of the morphospace at each Epoch. The Miocene was divided into two time bins for the mandible dataset due to the great number of Miocene specimens (see STAR Methods for details). See Figure S4 for other metrics, disparity in every clade, and phylogenetic inferred disparity and Data S1 for specimens used as well as the FAD and LAD (first and last appearance datum, respectively) used to calibrate the supertree.

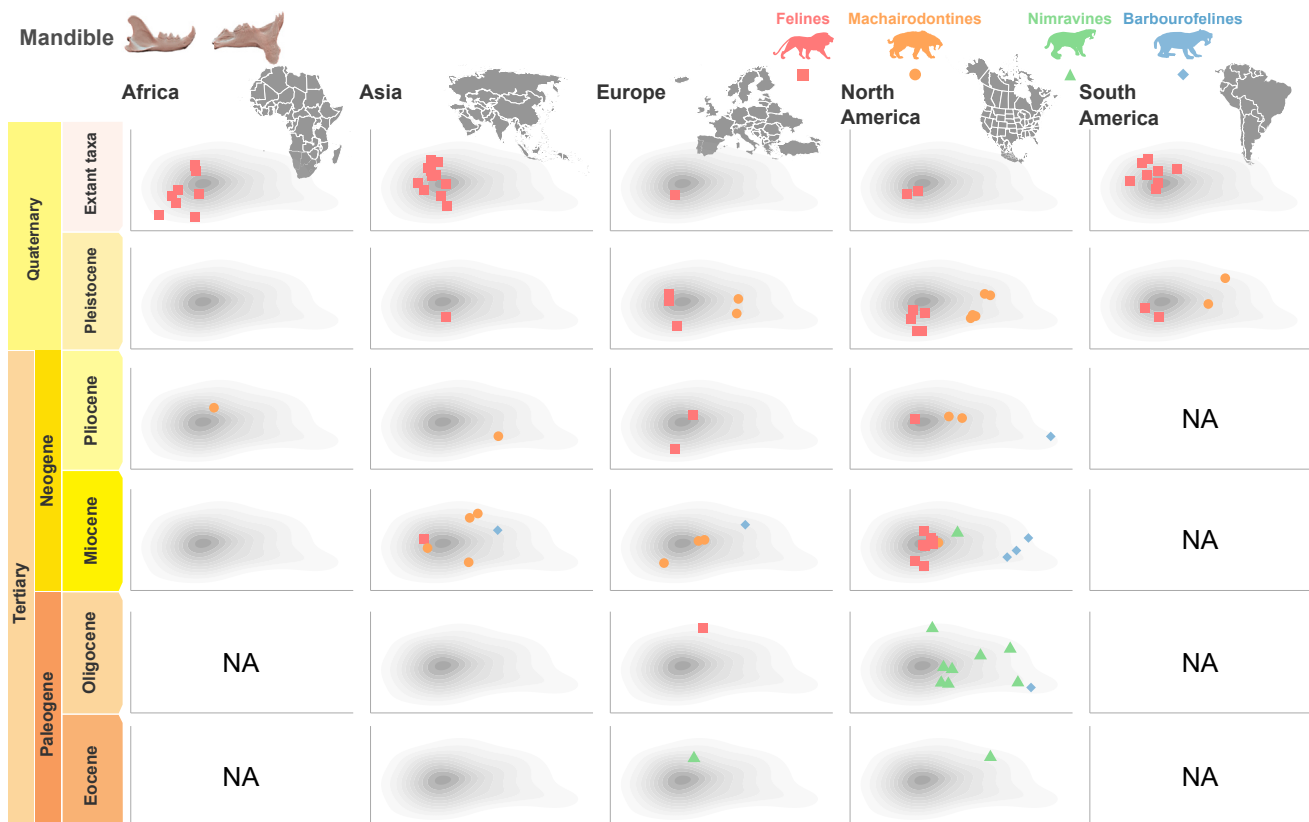


Figure 3. Occupation of the morphospace trough time and space for the mandible dataset
See [Figure S5](#) for the cranium. See also [Data S1A](#) and [S1B](#) for the age and locality of each specimen.

statistically significant results at the family level.⁵⁵ It is essential to consider the methodology used in these tests: Stayton's metrics rely on the maximum phenotypic distance between ancestors, whereas Castiglione's method also takes into account the time since cladogenetic divergence. Given the relatively short divergence time among the studied taxa, this could explain the lack of significance in the Castiglione tests. Furthermore, the overall conservative morphology within these groups may result in a low maximum phenotypic distance between lineages at various points in their evolutionary histories (Dmax), impacting the results concerning Stayton's metrics.

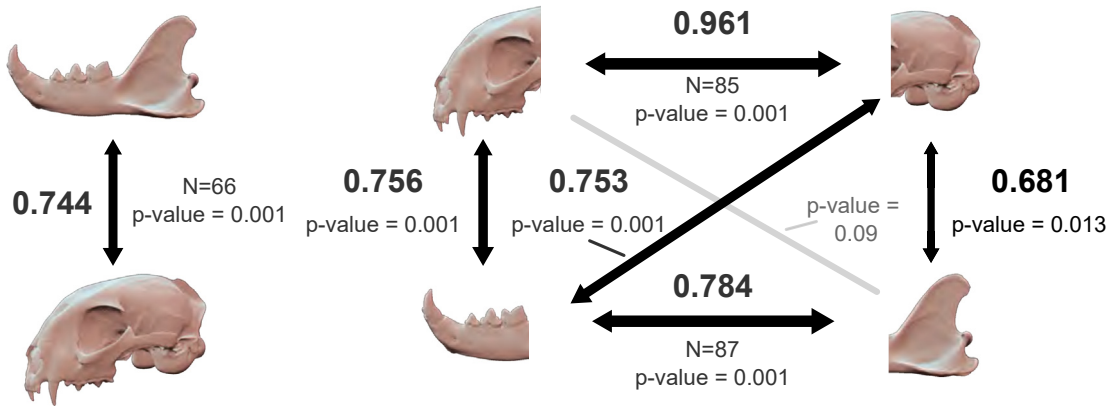
Nevertheless, our cluster dendrogram still emphasizes the substantial craniomandibular similarities between some taxa in both families. This suggests that cranial shape closely follows clade-based evolutionary shifts, with cluster dendrograms effectively reflecting the distinctions between different families. However, mandibular shape evolution appears to be more influenced by other factors and does not consistently align with these phylogenetic relationships.⁴⁰ In our cluster dendrograms, we identified two distinct groups for both crania and mandibles: one comprising taxa with the shortest upper canines and another containing taxa with the longest upper canines ([Figure S3](#)) with a mixture of felids and nimravids found within each group. Despite the apparent cranial similarities, there exists a well-structured spatial and temporal distribution of these morphologically analogous taxa. This arrangement minimizes potential competition

between morphologically similar species belonging to distinct clades, akin to the highly structured small felid guild observed in contemporary ecosystems, which acts to reduce competition.⁵⁶

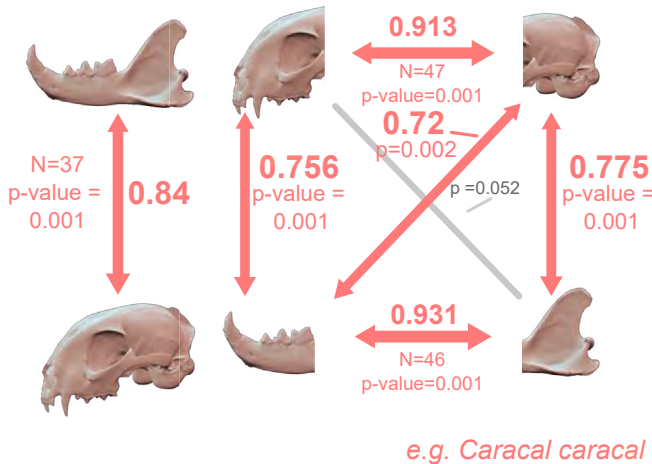
From an evolutionary perspective, small extant felines are as much derived as extreme sabertooth taxa

Small cat species, particularly in terms of cranial shape, form a second peak of recurring phenotypes. This demonstrates a deviation from the average/most common cat-like shape, comparable to the extremes seen in saber-toothed taxa like *Barbourofelis* spp. The small felines skull shape is characterized by a domed cranium, wide orbits with both postorbital processes nearly touching, a short and weak sagittal crest, and a low nuchal crest ([Figure 1B](#)). These morphological variations stand in stark contrast to the extreme saber-toothed carnivorans, such as *Barbourofelis fricki*, which display features like the rostrocaudal shortening of the sagittal crest and temporal fossa, the rotation of the occipital plane, and completely closed orbits, among others. Despite the differences, small felines find themselves equally distant from the main peak of recurring morphology, much like the extreme sabertooths. When considering both their mandibles and crania, it becomes evident that small felines are just as evolutionarily derived as the most extreme saber-toothed taxa and are equally distant from the earliest condition seen in the first felid, *Proailurus lemanensis*.^{57,58} Small cats are well separated from larger taxa by an area of uncommon phenotype

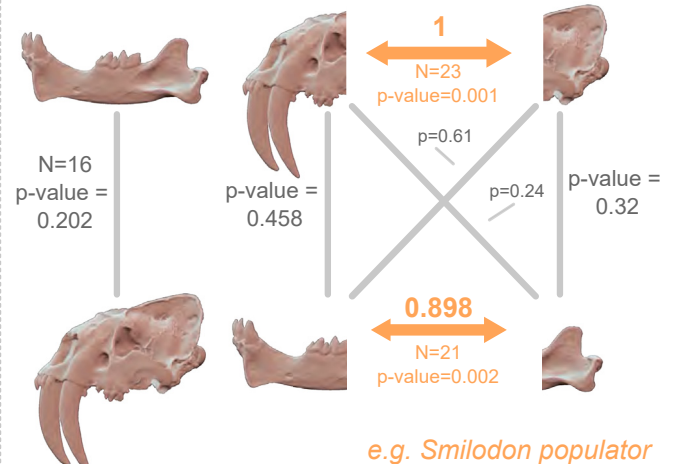
Whole dataset



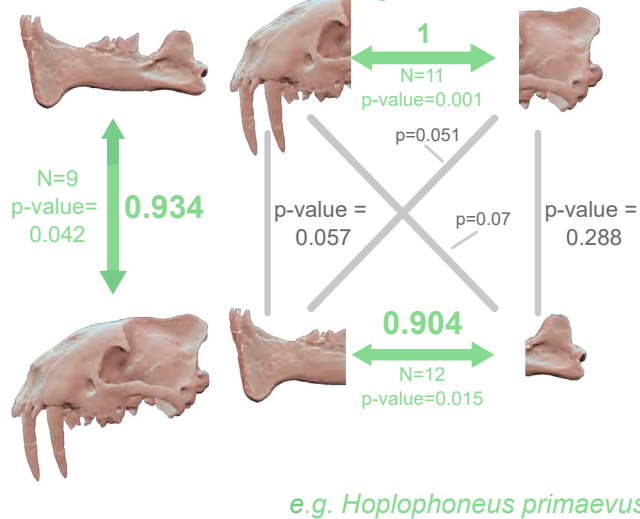
Integration per clade Felinae



Machairodontinae



Nimravinae



Barbourfelinae

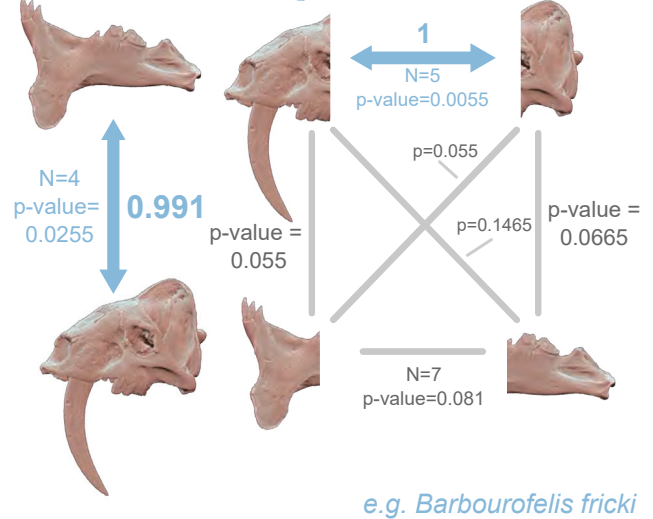


Figure 4. Phylogenetic integration tests

Line thickness depends on the r -partial least squares (PLS) values (indicated in bold next to each arrow as well as number of individuals n and p values). Gray lines without arrows indicate non-significant integration (p value > 0.05). See [Data S1C](#) and [S1F](#) for FAD and LAD used to calibrate the supertree.

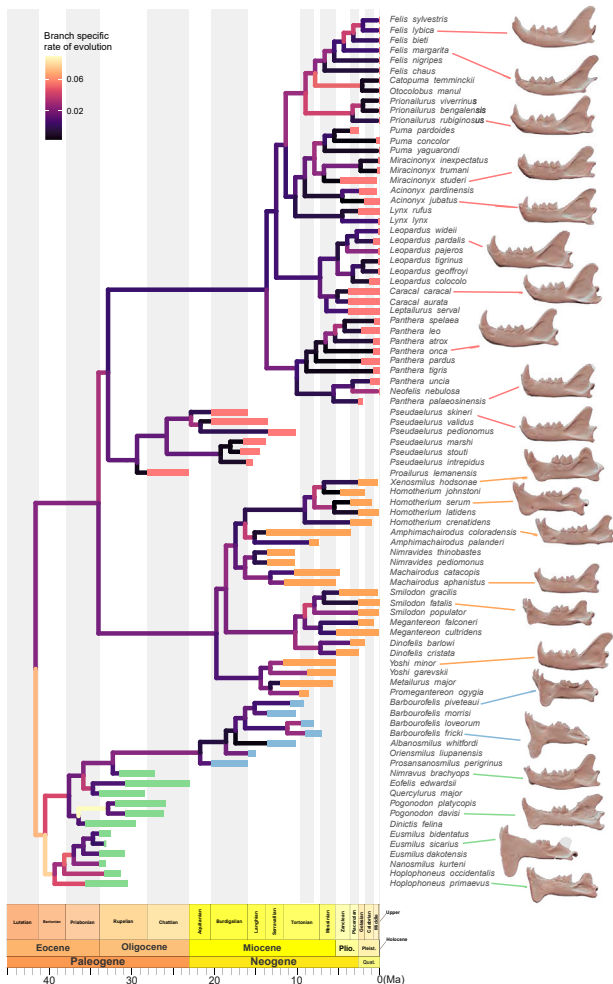


Figure 5. Branch-specific rate of morphological evolution for the mandible obtained from the RRPhylo function Plio, Pliocene; Pleist, Pleistocene; and Quat, Quaternary. See Figure S6 for the cranium. See Data S1C and S1F for FAD and LAD used to calibrate the supertree and STAR Methods for details.

which could be due to a functional constraint linked to the strong difference in prey size with big cats hunting prey much larger than them while smaller cats tend to hunt preys only a fraction of their body weight.

Interestingly, the small feline morphotype never occurred in nimravids and appeared quite recently in the fossil record (during the Pleistocene; see Figures S4 and S5). However, this observation should be approached cautiously, as it could be influenced by fossilization biases. In modern ecosystems, small cats primarily inhabit densely forested areas like jungles or even deserts,⁵⁹ which may reduce their chances of fossilization.⁶⁰ Additionally, their smaller size results in thinner and more fragile bones, making them less likely to be preserved over geological time scales.⁶¹

Cat-like carnivorans are on the decline since the Miocene

Despite showcasing a unique morphological innovation, the distinctive small morphotype, felids have, over time, become

confined to a considerably reduced area within the cat-like morphospace. This restriction has led to a significant loss in disparity, especially following the extinction of the machairodontine felids during the Late Pleistocene (Figures 2, 3, S4, and S5). Cat-like carnivorans reached their maximum disparity during the Miocene and, more precisely, at the end of the Miocene, characterized by the co-existence of extreme saber-toothed taxa and those with more feline-like characteristics. During this period, felids underwent substantial diversification.¹⁸ Although nimravids were already on the decline⁵⁴ the simultaneous presence of the first felids and the last nimravids resulted in a wide array of shapes among cat-like carnivorans.

The evolution of cranial and mandibular shape through time and space also reveals different patterns for each bone. Although the first nimravids quickly exhibited a cranial shape relatively close to the most recurring cat-like shape, their mandibles appeared more extreme. The Pleistocene witnessed a marked divergence between machairodontines and felines, with machairodontines exhibiting heavily derived saber-toothed forms that departed significantly from the average cat-like shape (Figures 2 and 3) in terms of mandibular shape. However, their cranial shape remained much more similar to large felines s.l. from the same Epoch. Machairodontinae were often cited as an example of over-specialization, making them unable to adapt to changing ecosystems and potentially contributing to their own extinction.⁶² Recently, Piras et al. found a higher extinction rate in saber-toothed taxa, attributing this pattern to a narrower ecological niche and less ecological flexibility.²¹ Although machairodontine felids displayed considerable diversity^{1,62,63} they ultimately succumbed to the various environmental shifts and faunal turnovers of the Late Pleistocene, vanishing alongside the rest of the megafauna.⁶⁴

However, our analyses reveal that despite the apparent diversity, machairodontine felids rapidly moved away from the most common cat-like morphology, with extreme saber-toothed taxa occupying more highly specialized niches by the Pliocene. These findings contrast with those of Romano,²² who observed a continuous increase in disparity among machairodontine felids, peaking during the Pleistocene. However, this discrepancy may be attributed to the limited number of taxa (17) and the use of a cladistic matrix from Christiansen,⁶³ which relies on character data rather than actual shape data. Although some authors support the use of cladistic data for characterizing disparity^{65–67} caution must be exercised when interpreting such results from matrices with limited characters and taxa. Romano²² interpreted this constant increase in morphospace as a “late saturation” (sensu⁶⁸). Nevertheless, with a more extensive dataset, it becomes evident that machairodontine felids, and cat-like carnivorans in general, had been experiencing a decline in disparity long before their eventual extinction. Despite the repeated evolution of large saber-toothed forms, this morphological specialization seems to act as a macro-evolutionary ratchet ultimately causing the decline of those lineages, as previously suggested in other studies.⁶⁹

Could a lower degree of phenotypic integration and a higher rate morphological evolution be potential drivers of the sabretooth morphology?

When examining the skull as a single unit, cranial shape appears to be more constrained by allometry in taxa with short upper

canines, while in taxa with elongated upper canines, it is more related to the length of these canines.¹⁵ Although not discussed as integration, Emerson and Radinsky⁶ already started to introduce the covariation between the coronoid process and other typical sabertooth traits. Later, Meloro and Slater⁷⁰ have highlighted distinct patterns of integration between the anterior (face) and posterior (braincase) portions of the cranium, these two modules being driven by body size in taxa with short upper canines while relative canine length appears to drive rostrum shape in saber-toothed taxa. More recently, Tamagnini et al.⁷¹ showed that extreme sabertooth adaptations of machairodontine break the craniofacial evolutionary allometry rule within felids. Similarly, Piras et al.²⁰ conducted integration tests on the cat-like mandible in their 2D geometric morphometric analyses, testing the hypothesis that sabertooth morphology entails covariation between the ascending ramus and the mandibular corpus.

In this contribution, we assessed integration degree between different subunits exploring new combinations, revealing that Felinae sensu lato is the clade with the highest degree of integration. This contrasts with the findings of Piras et al. who identified Barbourofelinae as the clade with the highest degree of integration, followed by Nimravinae, Machairodontinae, and Felinae, confirming their hypothesis. Saber-toothed taxa exhibit a wide range of morphologies within the same clade. For instance, within machairodontine felids, a similar reduction of the coronoid process can be associated with various degrees of development of a mandibular flange (see *Smilodon* spp., *Homotherium* spp., and *Megantereon* spp. or the development of the mandibular flange in *Barbourofelis* spp.; see Figure 1). Therefore, it is logical to assume that sabertooths have a lower degree of integration between the mandibular corpus and ramus to allow for such a wide range of possible combinations. By contrast, felines exhibit a more integrated morphology, with smaller taxa featuring slender and curved mandibular rami with a long and more posteriorly projected coronoid process, while larger taxa display a more robust corpus with a shorter and straighter coronoid process. It is the principle of morphological integration: if modules answer to different constraints, evolving independently, this eventually leads to a greater disparity, while more integrated structures tend to constraint phenotypic diversity.^{26–33} The greater disparity observed in clades with longer upper canines (Figures 1 and S4) further supports the idea of a lower degree of integration in these groups. This lower degree of integration might even represent the initial step that allowed some cat-like carnivorans to develop sabertooth morphologies. This reduced integration, coupled with bursts in the rates of morphological evolution during sabertooth diversification, appears to have played a pivotal role in shaping the diversity of cat-like carnivorans.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cub.2024.04.055>.

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The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
3D Surface scans	https://www.morphosource.org/projects/000548886?locale=en	See Data S1 for complete list with specimen numbers
Landmark coordinates	Online supplemental information + https://hdl.handle.net/2268/316060	Online supplemental information & Orbi.
Specimen list with metadata	Online supplemental information + https://hdl.handle.net/2268/316060	See Data S1 & Orbi.
Software and algorithm		
R Studio v 4.2.3	R Core Team ⁷²	https://www.r-project.org
VX Models	AMETEK, Inc. ⁷³	https://www.creaform3d.com/
Stratovan checkpoint v2018.08.07	Stratovan Corporation	https://www.stratovan.com/
Blender	Blender Foundation ⁷⁴	blender.org
Meshlab	ISTI – CNR ⁷⁵	http://www.meshlab.net/

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Narimane Chatar (narimane.chatar@uliege.be).

Materials availability

3D models used to perform the analyses are available on (Project ID: 000548886, <https://www.morphosource.org/projects/000548886>). For scans not available on MorphoSource (for instance, material that is not yet described) please contact Narimane Chatar (narimane.chatar@uliege.be) or the corresponding museum curator.

Data and code availability

All original code is available on Orbi, the ULiège Open Repository (<https://hdl.handle.net/2268/316060>) and Github (<https://github.com/cha-nar>) and is publicly available as of the date of publication as well as all the data: Landmark coordinates, Excel files containing the full material list with metadata, sliding sequence for landmarks, measurements, FAD and LAD.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

The dataset is composed of a total of 189 3D models, 99 mandibles (91 different taxa) and 90 crania (88 different taxa). Our dataset exclusively contains felids and nimravids: Respectively 32% and 37% of sampled mandibular and cranial specimens belong to living species (See [Data S1A](#) and [S1D](#) for detailed list of specimens including metadata), covering 33 out of the 42 (78%) recognized extant taxa. All specimens were adults and analyses were performed on the specimens themselves rather than species average.

Retrodeformation and missing data estimation

Some specimens were mirrored using VX models, as we chose to landmark only one side of each structure. Different repair tools were used to fix specimens that were slightly incomplete, this step was done in VX models and/or Blender. However, for some specimens lacking more consistent parts, we employed various reconstruction techniques. Figures showing specimens before and after retrodeformations are available on Orbi (<https://hdl.handle.net/2268/316060>).

Retrodeformation

Some fossil crania appeared relatively complete but underwent basic shearing induced by taphonomic processes. For those specimens, we retrodeformed them using the protocol described in Schlager et al.⁶⁹ To do so, we placed landmarks from our landmarking procedure that do not lie on the symmetry plane of the cranium (Landmarks 5, 6, 10, 11, 12, 13, 14, 15, 16, 17, 18, 20, 21 as well as curves 3, 4, 6, 7; see [Figure S1](#)). This procedure was performed using the Morpho package.⁷⁶ The specimens retrodeformed using this method were: *Panthera zdansky* (BC-294) *Dinofelis abeli* (AMNH-F-AM-50445) and *Panthera palaeosinensis* (PMU 21780-1) (See retrodeformations Figures on Orbi).

Other specimens underwent basic compression (along one axis). For those specimens, we used Blender to correct the deformations. Those specimens include *Eusmilus adelos* (USNM-12820), *Eusmilus sicarius* (YPM-VPPU-012953), *Metailurus* sp (AMNH-131854), *Metailurus parvulus* (NHMUK-PV-OR-28862) and *Megantereon cultridens* (MNHN.F.PET2001a) (See retrodeformations Figures on Orbi).

Missing landmarks estimation

Some specimens were still too incomplete to landmark fully, so we used missing landmark estimation. The proportion of specimens exhibiting missing data was 8% for the mandibles (8/99) and 21% for the crania (19/90) (See orange lines in [Data S1A](#) and [S1D](#)). However, we still excluded specimens that were too incomplete and used missing data estimation only when a fragment of the bone was missing, for instance a fragment of coronoid process missing on a mandible (curve 1 and potentially landmark 1) or the zygomatic arches missing on a cranium (Landmarks 18, 6 and curve 3). This estimation was performed using the `fixLMtps` function of the Morpho package⁷⁶ that estimates missing landmarks by mapping weighted averages from complete datasets onto the missing specimen using the three closest observations.

METHOD DETAILS

Digitization

Novel scans realized for this study were obtained using a Creafom HandySCAN 300 laser surface scanner with a resolution varying from 0.2 mm to 0.5 mm (see [supplemental information](#)). Scans were initially cleaned, merged, and exported in ply format using VXEelements v.6.0. Models were then treated using VXmodel. Additional models were obtained from (Robles et al.⁷⁷ (*Albanosmilus jourdani*); Tseng et al.⁷⁸ (*Panthera blytheae*); Spassov and Geraads⁷⁹ (*Yoshi garevskii*); Adams et al.⁸⁰ (*Dinofelis barlowi*, *Dinofelis piveteaui*) Geraads and Spassov⁸¹ (*Amphimachairodus giganteus*); Wang et al.⁸² (*Oriensmilus liupanensis*)) see [Data S1](#) for details.

Morphometric data collection

We placed eleven type-II landmarks⁸³ and three curves composed of 27 semi-landmarks on the mandible ([Figure S1](#); [Data S1A](#)) using Stratovan checkpoint v2018.08.07⁸⁴ and exporting the coordinates as '.pts' files. On the cranium, we placed a total of twenty-two type-II landmarks + type-III landmarks and seven curves composed of 50 semi-landmarks ([Figure S1](#); [Data S1D](#)). All analyses were then performed in the R v 4.2.3 statistical environment.⁷² See [Tables S1](#) and [S2](#) for the description of each landmark used. Landmark coordinates were imported into R using the custom function 'import.pts' (<https://github.com/cha-nar/importpts>). Stratovan checkpoint exports all the missing coordinates as '9999' missing values were replaced by 'NA' using imbricated loops to allow the `fixLMtps` function from the Morpho package⁷⁶ to estimate the missing coordinates using weighted averages from the three most morphologically similar, complete, configurations. which uses weighted averages.

Generalized Procrustes superimposition was performed using the `gpagen` function from the `geomorph` package v4.0.5⁸⁵ specifying the semi-landmarks to slide using the 'curve' argument ([Data S1B](#) and [S1E](#)). Log centroid size was extracted from cranial shape data during Procrustes superimposition and used as a proxy of overall size.

Phylogeny

To assess the influence of phylogeny in our results and visualization purposes, we built a composite tree containing extant carnivorans, nimravids, and machairodontine felids in R by binding the trees published by Barrett,⁸⁶ Jiangzuo et al.,⁸⁷ and Slater and Friscia⁸⁸ using the 'bind.tree' function of the ape package.⁸⁹

We dropped the tips absent from our dataset using the 'drop.tip' function. In the machairodontine tree, we also removed *Panthera leo* and *Lynx rufus* as they were already present in the extant carnivoran tree and did the same for *Homotherium serum*, *Smilodon populator* in the extant carnivoran tree. Twenty nine taxa were absent from the sampled trees and had to be manually added in positions found in the literature: *Yoshi minor*, *Yoshi garevskii*, *Amphimachairodus palanderi*, *Nimravides pediomonus*, *Nimravides thinobastes*, *Machairodus catacopis*, *Smilodon gracilis*, *Pseudaelurus stouti*, *Pseudaelurus intrepidus*, *Pseudaelurus marshi*, *Pseudaelurus skineri*, *Pseudaelurus pedionomus*, *Pseudaelurus validus*, *Acinonyx pardinensis*, *Catopuma temminckii*, *Felis sylvestris*, *Leopardus pajeros*, *Leopardus wideii*, *Leptailurus serval*, *Miracinonyx studei*, *Miracinonyx inexpectatus*, *Panthera atrox*, *Panthera speleae*, *Panthera paleosinensis*, *Panthera gombaszoegensis*, *Puma yaguarondi*, *Panthera uncia*, and *Puma pardoides*.^{87,90–94}

We time-scaled the resulting supertree using the 'timePaleoPhy' function from the Paleotree package⁹⁵ through the minimum branch length (mbL) method. Divergence between Felidae and Nimravidae was set based on the results published by Barrett et al.⁹⁶ We generated the time tree using the 'geoscalePhylo' function from the paleotree package⁹⁵ with first and last occurrence dates obtained from the literature^{79,86,87,90,91,97,98} or from the PBDB (paleobiodb.org) (see [Data S1C](#) and [S1F](#)). The super tree is available in Newick format on Orbi (<https://hdl.handle.net/2268/316060>).

Spatiotemporal distribution

To understand the spatiotemporal distribution of cat-like carnivorans and assess the influence of environmental factors on the evolution of disparity and diversity through time, we collected time (epoch and age) and localization data (continent and

country or state) for all sampled species (Data S1A and S1D). We duplicated entries in the dataset only when a species is recorded in multiple time bins or multiple geographical bins. Due to its length compared to other Epoch, the Miocene was divided into two sub-Epoch for the disparity analyses as it might have affected our results. Early Miocene corresponding to Aquitanian, Burdigalian and Langhian (23.03 to 13.82 Ma) and Late Miocene corresponding to Serravallian, Tortonian and Messinian (13.82 to 5.333 Ma).

QUANTIFICATION AND STATISTICAL ANALYSIS

Craniomandibular shape variation

A principal components analysis (PCA) was performed using the `gm.prcomp` function of the `geomorph` package v4.0.5⁸⁵ to assess the primary aspects of shape variation and morphospace occupation. To visualise the areas of recurring morphologies (in the sense of frequently colonised) in our morphospace, we computed the kernel density of occupation by modifying the script from Fischer et al.⁹⁹ To better visualize shape changes in the morphospace we used the `plot_3D_bt_shape` custom function https://github.com/cha-nar/plot_3D_bt_morphospace (see Figure S2) to create a 3D backtransform morphospace.

To ensure that the larger sample size of living cats does not disproportionately influence the variance of the eigenvectors, we implemented a loop to randomly exclude 20 out of the 33 extant felid species from our set of Procrustes coordinates for the mandibular dataset. This reduction brought the number of extant species to 13, a sample size comparable to that of our nimravines (N=12) and barbourufelines (N=9). Subsequently, we conducted a Principal Component Analysis (PCA) on this artificially reduced dataset (those morphospaces are available on Orbi <https://hdl.handle.net/2268/316060>). Notably, the predominant shape variation observed was attributed to fossil taxa, even when only a dozen extant species were included. We used the `procD.lm` function from the `geomorph` package v4.0.5 to test the influence of the size on shape (allometry) on our shape data as previous studies shown that allometry is pervasive in felids.^{100–102} We used the `physignal` function from the `geomorph` package v4.0.5⁸⁵ to quantify the phylogenetic signal from Procrustes shape variables in both crania and mandibles within the whole dataset and per clades (See Table S4).

Disparity

We used the `dispRity` package v1.7.0¹⁰³ to compare clade disparities (Felinae, Machairodontinae, Nimravinae, and Barbourufelinae). Morphological disparity was estimated as a bootstrapped sum of variances (1000 bootstraps). Then we computed the global disparity through time in cat-like carnivorans as both a bootstrapped sum of ranges and a bootstrapped sum of variances (1000 bootstraps), for each time bin (Eocene, Oligocene, Miocene, Pliocene, Pleistocene, and Extant). Sum of variances captures the dispersal of taxa within morphospace and can be influenced with the intensity of sampling, while the sum of ranges indicates the amount of overall morphospace occupation and can be biased by extreme yet rare morphologies. Those two metrics are therefore commonly combined to yield an honest picture of disparity (e.g; Wills et al.,⁶⁶ Brusatte et al.,¹⁰⁴ Butler et al.,¹⁰⁵ and Prentice et al.¹⁰⁶). For comparison, we also computed the phylogenetically informed disparity through time using the `dtl` function from the `Geiger` package¹⁰⁷ using 1000 simulations and with a 95% CI. See Figure S4.

Convergence

We assessed the significance of the evolutionary convergence pairs of taxa using the `CalconvCt` function from the `convevol` package v2.0.0 to extract C-metrics (Stayton²⁴; Grossnickle et al., 2023¹⁰⁸) as well as the method implemented by Castiglione et al.²⁵ in the `RRPhylo` package.¹⁰⁹ Convergence and conservatism are two evolutionary processes resulting in distantly related phylogenetic tips with similar morphology. While Stayton's traditional C-measures are significant only in presence of convergence, the theta by Castiglione et al.²⁵ can be significant with both convergence or conservatism. Using a combination of those metrics can therefore help us distinguish between the two evolutionary patterns: significant C1 and theta indicates convergence, non-significant C1 and significant theta stands for conservatism, and non-significant C1 and theta indicate the absence of both convergence and conservatism. Traditional C-measures became widely used as they can distinguish between convergence and conservatism, which both those process resulting in distantly related phylogenetic tips with similar morphology. However, those metrics are extremely slow to compute, and Castiglione's method considers the time spent since their cladogenetic divergence where the temporal is thus important. We applied both those on the whole set of Procrustes coordinates as the occurrence of spurious patterns of convergence can increase in reduced shape spaces. See Figures S2G and S2H for a graphical representation of the convergence pairs tested, only C1 results are shown in the main document (see Table 1), C2–C4 results are available in Table S3. See Figure S3 for tanglegrams showing the correspondences between the phylogenetic supertree and the hierarchical cluster build from the distance matrix based on the Procrustes coordinates.

Patterns of phenotypic integration

To assess the potential role of phenotypic integration in sabertooth disparity we investigated patterns of integration between different craniomandibular modules (seven combinations in total). To do so, we performed phylogenetic two-block partial least squares analyses (2B-PLS) using the `phylo.integration` function from the `geomorph` package⁸⁵ on the residuals from shape on size as size-related shape differences could inflate integration results. We then use the pls coefficient (*r*-PLS) as an estimation of

the degree of phenotypic integration between the two structures. The significance of the r-PLS coefficient is evaluated using 1000 permutations against the hypothesis of a complete absence of covariation between blocks.

Rate of morphological evolution

We investigated branch-specific evolutionary rates of evolution through time in the craniomandibular complex using the RRphylo function from the RRPhylo package.¹⁰⁹