Not a jaguar after all? Phylogenetic affinities and morphology of the Pleistocene felid *Panthera gombaszoegensis*

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Abstract

*Panthera gombaszoegensis* is a fossil pantherine from the Pleistocene of Eurasia. It has been considered to be the closest ancestor the jaguar (*Panthera onca*) due to dental similarities, and has even sometimes been considered to be a subspecies of jaguar. However, our knowledge of this taxon is limited by the scarcity of cranial remains, which has made it difficult to properly assess the phylogenetic affinities and possible ecological role of this taxon. Here, we describe a new cranium of *P. gombaszoegensis* from Belgium, and present a morphometric analysis of the cranium and dentition of extinct and extant pantherines. Whereas the lower dentition of *P. gombaszoegensis* is similar to that of *P. onca*, similarities were not recovered in other parts of the skull. Some cranial traits of *P. gombaszoegensis* resemble those of other pantherines, especially larger species such as the tiger (*P. tigris*). Some similarities with taxa such as tigers (*P. tigris*), lions (*P. leo*), and leopards (*P. pardus*) in the skull of *P. gombaszoegensis*, suggesting a diet adapted to a wide prey spectrum. The first ever assessment of the phylogenetic placement of *P. gombaszoegensis* places this taxon closer to *P. tigris* than to *P. onca*, which considerably simplifies the biogeographic history of pantherines.

Key words: *Panthera*; morphometry; phylogeny; *gombaszoegensis*; jaguar.
Introduction

Pantherinae, also known as the ‘big cats’, is a group of apex predators that dominate food chains in every region they are found. The subfamily includes the living lion (Panthera leo), tiger (Panthera tigris), jaguar (Panthera onca), leopard (Panthera pardus) and snow leopard (Panthera uncia) (Turner & Antón 1997). Pantherines colonised a wide range of habitats since their first appearance, from the tropical rainforests of South America (jaguar, Panthera onca Seymour 1989) to the Himalayan peaks (snow leopard, Panthera uncia Janečka et al. 2008). The putatively oldest pantherine, ‘Panthera’ blythae, was found in the Zanda Basin of the north western Himalaya Range, and is approximately 6 million years (Myr) old (Tseng et al. 2013). However, it is not considered as a true pantherine by all authors (see Geraads & Peigné 2017), although pantherines likely originated in the central–northern Asia elsewhere in the Holarctic (Mazák 2010; Tseng et al. 2013). Pantherines became widespread during a series of Pliocene–Pleistocene migration events (Johnson et al. 2006), the first pantherinae arriving in Europe being the large-bodied cat known as Panthera gombaszoegensis during the Early Pleistocene (O’Regan & Turner 2004). Panthera gombaszoegensis had a wide geographical range (Fig. 1A), potentially due to the absence of competitors until the faunal turnover that took place during the Early/Middle Pleistocene transition when Panthera leo and Panthera pardus migrated into Europe (O’Regan & Turner 2004; Hemmer & Kahlke 2005). Often called the ‘European jaguar’ or the ‘Eurasian jaguar’, P. gombaszoegensis was a medium-to-large pantherine which went extinct approximately 350 thousand years (kyr) ago and is commonly considered as the ancestor of the extant jaguar, Panthera onca (Hemmer 1981; Hemmer et al. 2001; O’Regan & Turner 2004).

Initially regarded as a singular pantherine species (Kretzoi 1938a), P. gombaszoegensis was later reclassified as a subspecies of the modern jaguar: Panthera onca gombaszoegensis (Hemmer 1971). The status of the taxon is still debated, with a series of studies regarding P. gombaszoegensis as a fully valid species (Argant & Argant 2011; Reynolds 2013; Marciszak 2014; Stimpson et al. 2015; Jiangzuo & Liu 2020), whereas others note that the differences between P. gombaszoegensis and the living jaguar are not sufficient to warrant full specific distinction (Hemmer et al. 2001, 2010; Hankó 2007; Mol et al. 2011). The scarcity of P. gombaszoegensis material has made it difficult to unravel this taxonomic debate, especially as the species was erected solely based on isolated teeth described by Kretzoi (1938b) and almost all the material ever reported in the literature is composed of dental remains. It was recently demonstrated that dental traits of P. onca and P. gombaszoegensis are morphologically different (Jiangzuo & Liu 2020), although the cranial anatomy of P. gombaszoegensis remains insufficiently
described in the literature to draw definitive conclusions. As a result, there are very few truly multivariate morphometric analyses comparing *P. gombaszoegensis* to other pantherine species (Mazák, Christiansen, & Kitchener 2011), with most studies limited to bivariate analyses (Langlois 2002; O’Regan & Turner 2004; Argant & Argant 2011; Mol *et al.* 2011; Jiangzuo & Liu 2020; Marciszak & Lipecki 2021). Moreover, the very fragmentary state of fossils assigned to *P. gombaszoegensis* resulted in an absence of this taxon in phylogenetic analyses of pantherines. Indeed Christiansen (2008), Tseng *et al.* (2013), and King & Wallace (2014) have systematically excluded *P. gombaszoegensis* from cladistic datasets due to the amount of missing data.

In this study, we present well-preserved, as-yet undescribed material of *P. gombaszoegensis*, unearthed during the 1980’s from ‘La Belle-Roche’ in southern Belgium (Fig. 1B). We posit that this material can shed light on this obscure taxon, and may help resolve the phylogenetic relationships of this taxon within the Pantherinae. We hereby provide the first comparative description of the craniomandibular anatomy of *P. gombaszoegensis*, and assess its relationship with the modern jaguar *P. onca* and with other extant pantherines, both in terms of phylogeny, morphometry, and ecology.

### Material and methods

#### Institutional abbreviations

**AMNH** American Museum of Natural History (New York, USA); **IMNH** Idaho Museum of Natural History; **IVPP** Institute of Vertebrate Palaeontology and Palaeoanthropology; **MAV** Museo Anatómico de la Universidad de Valladolid; **MNCN** Museo Nacional de Ciencias Naturales (Madrid, Spain); **MNHN** Museum National d’Histoire Naturelle (Paris, France); **PMU** Paleontological Museum Uppsala universitet (Uppsala, Sweden); **RMCA** Royal Museum for Central Africa; **ULgPA** Université de Liège, palaeontology collections.

#### Fossil locality

‘La Belle-Roche’ is a fossil site located in the Belgian province of Liège (Municipality of Sprimont, Belgium), 20 km south of Liège, and a couple of kilometres away from the town of Sprimont (Fig. 1B). In ‘La Belle-Roche’ (translated as ‘The Beautiful Rock’), mudstone deposits filled a karstic cavity carved into Carboniferous limestone (Cordy *et al.* 1993). Uranium–thorium dating provides a minimum age of 350 kyr for the mudstone deposit and fossils it contains (Gascoyne & Schwarz 1985). The presence of *Ursus deningeri* and *P. gombaszoegensis* indicates the mudstone may have been deposited in an earlier part of the Middle Pleistocene (Roebroeks & Stapert 1986), with estimations reaching 500 kyr (Cordy & Ulrix-Closset 1991; Draily & Cordy 1997).
Figure 1: A, Occurrences of *P. gombaszoegensis* compared to the historic extent of the extant *P. onca* and *P. tigris*, estimated since 1900 based on Luo et al. (2004) and Seymour (1989). B, Location of the ‘La Belle-Roche’ fossil site, Belgium. Animal silhouettes were obtained from PhyloPic (phylopic.org). Image credits: Manabu Sakamoto (*P. onca*) and Sarah Werning (*P. tigris*).
Even though ‘La Belle-Roche’ is the only known Belgian fossil site were *P. gombaszoegensis* has been recorded, its presence can be inferred in Belgium before the deposits of ‘La Belle-Roche’, as suggested by the estimated distribution in Europe from 2 Myr in the surrounding countries (Fig. 1A and Fig. S1). Although most of the material from ‘La Belle-Roche’ is extremely fragmentary, this site has yielded a nearly complete cranium of *P. gombaszoegensis* (ULg PA BRII-81-146) and three fragmentary dentaries (ULg-PA-BR-III-M13-79; ULg-PA-20210823-01; ULg-PA-BRII-455). Well-preserved cranial material of *P. gombaszoegensis* is rare (see Plate 1 from Argant & Argant 2011 and Figure 3 A1–3 from Jiangzuo & Liu 2020), and the cranium from ‘La Belle-Roche’ is probably the most complete known to this date.

**Material**

The cranium of *P. gombaszoegensis* ULg-PA-BR-II-81-146 described in this contribution (Fig. 2A–B and Fig. S2) belongs to the collections of the University of Liège, and is currently exhibited in Le Grand Curtius Museum (Liège, Belgium). Three mandibular fragments were also found at La Belle-Roche: two fragmentary right dentaries ULg-PA-BR-III-M13-79 and ULg-PA-20210823-01, and a fragmentary left dentary ULg-PA-BRII-455 (Fig. 2C–E and Fig. S3). The dentary fragments also belong to the collections of the University of Liège; however, ULg-PA-BRII-455 is currently on display at the Musée du Pays d'Oirthe-Amblève, with ULg-PA-BR-III-M13-79 housed in Le Grand Curtius Museum. For comparative analyses, different extant pantherines from the Museum National d'Histoire Naturelle (MNHN, Paris, France) and the Royal Museum for Central Africa (AfricaMuseum, Tervuren, Belgium) as well as *P. palaeosinensis* from the Paleontological Museum Uppsala universitet (PMU, Uppsala, Sweden), were studied (Table S2 ESM). A three-dimensional (3D) scanned model of a leopard *P. pardus* from the American Museum of Natural History (New York, United States) was downloaded from MorphoSource (media M7779; see Tseng *et al.* (2016) for the original publication of the 3D model), as well as a 3D model of the cranium of the cave lion *P. spelaea* from the Idaho Museum of Natural History, initially published in Melchionna *et al.* (2021). All specimens were adult based on their deciduous dentition; wild caught specimens were preferred for extant species, and we excluded animals with visible dental wear from the study.
Figure 2: Fossil material of P. gombaszogensis from la Belle-Roche. Cranium ULg PA BR-II-81-146 in A, dorsal and B, ventral view. C-E mandibular fragments. C ULg-PA-BR-III-M13-79, D ULg-PA-BRII-455, E ULg-PA-20210823-01. Scale bars represent 2cm
3D data acquisition

In order to compare the different pantherine specimens, material from fossil and extant species were scanned using a Creaform HandySCAN 300 laser surface scanner with a 0.2 mm resolution. 3D models associated with this study are available on Morphosource (Project ID: 000445179, https://www.morphosource.org/projects/000445179?locale=en); models include four specimens of *P. gombaszoegensis* (ULg-PA-BR-II-81-146; ULg-PA-BRII-455; ULg-PA-BR-III-M13-79; ULg-PA-20210823-01), two specimens of *P. spelaea* from the University of Liège (ULg-PA-SCHM-II-14-11; ULg-PA-BRIII-L14-5), and one specimen of *P. palaeosinensis* from the Paleontological Museum Uppsala (cranium PMU 21780/1 and mandible PMU 21780/2).

Phylogenetic analyses

Our phylogenetic analyses are based on the morphological character matrix published by Tseng *et al.* (2013) including fossil and extant pantherines with *Neofelis nebulosa* (the clouded leopard), *Leopardus pardalis* (the ocelot) and *Puma concolor* (The mountain lion) as outgroups. We added *P. gombaszoegensis* as a new OTU and added a new morphological character (Character 40: Contact between both occipital condyles on the ventral part of the cranium: 0 absent, 1 present). We also revised the score of *P. uncia* for the character 1 (position of the nasal-frontal suture), as we observed contradicting character states in one of our specimens. Character 1 for *P. uncia* is now scored as a polymorphism (state 0&1). Our first-hand observations of *P. palaeosinensis* (specimen PMU 21780/1-2) allowed us to score two additional characters states in the matrix of Tseng *et al.* (2013): character 27 (Jugal-Maxillary suture, medial side) as “suture extends posteriorly, then cuts ventrally at the base of the zygomatic arch” (state 0), and the character 58 (Upper canine labial ridges) as “present” (state 1). In addition, we discretised character 48 (P3 parastyle size compared to P3 length), placing the limit between ‘short’ and ‘long’ at 0.5 relative to P3 length as often seen in the literature. Finally, we modified character 42 (Mandibular symphysis angle relative to horizontal ramus) as the previous character states (0: Anteriorly inclined (> 130°) and 1: Weakly inclined (110-120°)) could not be scored for *P. gombaszoegensis* since all the angles we measured were comprised between 120 and 130° (120.400, 127.558, 128.867 124.764, 122.677, 127.335, 128.994, mean = 125.799°). We therefore slightly modified the states to 0: Anteriorly inclined (> 121°) and 1: Weakly inclined (110-120°). The matrix was generated using Mesquite V3 (Maddison & Maddison 2019).

We used TNT v 1.5 (Goloboff & Catalano 2016) to perform the phylogenetic analysis under implied weighting (see appendix S1 for the script). We employed a molecular scaffold to constrain the tree topology, according to the recent
molecular analyses which included fossil taxa (Barnett et al. 2016). We expanded the memory of TNT to a maximum of 100,000 trees. We set the search parameters at: New Technology Search, 200 ratchet iterations, 10 cycles of drifting, 5 hits and 5 replications for each hit. We then used the tree branch bisection and reconnection algorithm (TBR) to fully explore the tree islands identified by the ratchet. Nodal support was measured using symmetric resampling with 1000 replications, each replication involving a New Technology search with a change probability of 33%. We choose symmetric resampling over bootstrapping or jack-knifing, as this measure is not affected by character weighting and is thus more appropriate to deal with implied weights (Goloboff et al. 2003) (See ESM for the TNT scripts). To test for the influence of character weighting, we ran these analyses with increasing value of the concavity constant $K$ ($K=3, 6, 9, \text{ and } 12$). Increasing the value of $K$ reduces the penalty applied to homoplastic characters; however, all the described analyses retrieved the same tree topology. The best score was obtained when $K$ was set to 12 so we computed the time calibration on the strict consensus cladogram for $K=12$. To do so, we used the ‘equal’ method of the ‘timePaleoPhy’ function from the strap v1.4 package in R (Bell & Lloyd 2015) and a dataset describing the temporal biozones of each OTU (See table S9 for the FAD, LAD and each reference used to estimate the temporal distribution). We then generated the time tree using the ‘geoscalePhylo’ function from the paleotree v3.3.25 package (Bapst 2012). Ancestral states were reconstructed using Mesquite.

**Morphological data**

Anatomical descriptions are based on previous research (Barone 1986; Schaller et al. 2007; Evans & Lahunta 2013; Jennings & Reighard 2019) and we used the terminology recommended by the *Nomina Anatomica Veterinaria* (2012). Figure S4 shows the main terms used in the description, based on an extant specimen of *P. tigris* (collection ID: MNHN-ZM-AC1931-60).

Multiple measurements were taken capturing the overall shape of the upper dentition (10 measurements, Fig. 3A), lower dentition (9 measurements, Fig. 3B), and cranium (13 measurements, Fig. 3C–E) of the material from La Belle-Roche and other extant and fossil pantherines. To complete our dataset, measurements published by O'Regan (2002) and Jiangzuo & Liu (2020) on the upper dentition were also included in the morphometric analysis (see Table S2). All measurements and ratios on the cranium and on the upper and lower dentition are provided in the supplementary material (Tables S3–S8). All measurements were taken on 3D meshes derived from scanning using the software GOM Inspect suite 2020 (Gesellschaft für Optische Messtechnik, Germany 2020).
Figure 3: 3D model of the cranium and mandible of Panthera tigris MNHN-ZM-AC-1931-60 showing the different measurements taken on A, the upper dentition; B, the lower dentition and C-E, views of the cranium: C, lateral; D, ventral; E, dorsal. Abbreviations:Bal, Basioccipital + Basisphenoid length; BrW, Maximum braincase width; CBW, Condylobasal maximum width; CCL, Canine to choanae length; CH, Upper canine height; cL, lower canine anteroposterior length; CL, Upper canine anteroposterior length; COL, Canine to occiput length; CTL, Cheek teeth anteroposterior length; cW, lower canine width; CW, Upper canine width; diastema, diastema length; MaPW, Mastoid process maximum width; M1CL, length from M1 to C; m1L, lower first molar length; m1W, lower first molar width; NaW, Nasal width; OPL, Orbit to premaxilla length; POW, Post-orbital process width; p3L, lower third premolar length; P3L, Upper third premolar length; p3W, lower third premolar width; P3W, Upper third width; P4H, Upper fourth premolar height; p4L, lower fourth premolar length; P4L, Upper fourth premolar length; P4ML, Upper fourth premolar metacone length; P4PL, Upper fourth premolar paracone length; p4W, lower fourth premolar width; P4W, Upper fourth premolar width; TFL, Temporal fossa length; ZW, zygomatic width.

Morphometric analyses
All morphometric analyses were run using the R statistical environment (version 4.0.5; 2021-03-31) (R Core Team 2021). The R script used to run the analysis is provided in the electronic supplementary material. The morphological dataset was imported using the ‘read.csv’ function and following the protocol published in Fischer et al. (2017). We applied a 50% completeness threshold on every specimen to avoid distortion of morphospace due to missing data from incomplete specimens. After applying this threshold, the upper dentition dataset contained 72 specimens out of the 96 measured. All the specimens from the lower dentition (20 specimens) and the cranial measurements dataset (17 specimens) passed the completeness threshold.

The morphological variables of each dataset were then scaled (z-transform), and distance matrices (based on pairwise dissimilarities) were computed from these scaled datasets using the ‘dist’ function from the stats v4.3.0 package, which uses Euclidean distances as suggested by Legendre & Legendre (1998, pp 424-444). We generated morphospaces using two distinct ordination methods, both capable of handling missing values: a Principal Coordinates Analysis (PCoA), using the ‘pcoa’ function implemented in the ape v5.5 package (Paradis & Schliep 2019), and a non-metric multidimensional scaling (NMDS), using the ‘metaMDS’ function of the vegan v2.5-7 package (Dixon 2003). To estimate the influence of the allometric component in our dataset we used a loop to perform a linear regression between the first 10 PCo axes of each dataset and a variable representative of the size (Canine to occiput length, lower canine length and upper canine length). For the upper and lower dentition datasets, a permutational multivariate analysis of variance (PerMANOVA, formerly known as non-parametric / NP-MANOVA) (Anderson 2001) was performed using the ‘adonis2’ function from the vegan v2.5-7 package.
PerMANOVA was performed (1000 permutations using the ‘euclidean’ method) on the distance matrix of the ratios to test for significant differences between *P. gombaszoegensis* and the other pantherines and we performed post hoc corrections on the significant p-values using the False discovery rate (fdr) correction through the method argument from the ‘p.adjust’ function in the stats v3.6.2 package.

**Systematic palaeontology**

MAMMALIA Linnaeus, 1758  
CARNIVORA Bowdich, 1821  
FELIFORMIA Kretzoi, 1945  
FELIDAE Fischer, 1821  
FELINAE Fischer, 1821  
Genus *PANTHERA* Oken, 1816

*Panthera gombaszoegensis* Kretzoi, 1938b

*Holotype.* B991, a series of isolated teeth: C, c, P4, p3 and two m1 illustrated in Kretzoi (1938b).

*Emended diagnosis.* The P4 ectoparastyle is present with a straight anterior edge and a curved metacone. The P3 has variable anterior and posterior cusps, but usually large with a pronounced cingulum. Vertical groove on the upper canine may be present or indiscernible. Large sagittal crest and strong nuchal crest; vertically oriented occipital condyles; upper incisors positioned in a straight line, with I3 being the largest; large oval-shaped mystacial foramen; wide nasal aperture; nasal bones extend to or beyond the frontomaxillary suture. The m1 protoconid larger than the paraconid; cingulum usually present; talonid very rare. The p4 has a large protocone; anterior and posterior cusps and cingulum are distinct and usually large. The p3 is highly variable, although cusps usually ill-defined with a small posterior cusp. There are no determining features on the mandible, but all specimens have two mental foramina and usually have a straight (but not vertical) symphysis.

**Note.** A well-defined vertical groove on the upper canine was previously listed in the diagnosis of *P. gombaszoegensis*, but our observation suggests this feature is variable. For this reason, we have emended the diagnosis of *P. gombaszoegensis*.

**Type locality and horizon.** Gombasek, Slovakia. Gombasek Quarry, Cave deposit, Middle Pleistocene.

**Newly referred material.** ULg PA BR-II-81-146, ULg-PA-BRII-455, ULg-PA-BR-III-M13-79, ULg-PA-20210823-01.

**Comparative description**

As Jiangzuo and Liu (2020) recently provided a comprehensive description of the dentition of *P. gombaszoegensis*, we focused on the cranium for the description in this contribution. The mandibular fragments from ‘La Belle Roche’ are not well preserved enough to
be described here, but there is a recent description of mandibular remains of this taxon published by Jiangzuo et al. (2022). A more detailed and complete comparative work should be done on the mandibular anatomy could be done including the new material described by Jiangzuo et al. (2022) by also other well preserved material such as the complete left dentary figured in Langlois (2002).

**General state of preservation**

The cranium ULg PA BR-II-81-146 is relatively complete and is mostly three dimensionally preserved (Fig. 2A–B and S2). It lacks parts of both nasals, pieces of both premaxillae, both pterygoid flanges, and both auditory bullae. The skull is slightly distorted mainly due to a minimal rotation of the palatal region. Moreover, it is worth noting that the anterior-most section of both parietals is restored in plaster (Fig. 2). Due to some parts of bones being missing, some measurements for the morphometric analyses could not been taken on the cranium ULg PA BR-II-81-146; these were coded as missing values (NA). *Panthera gombaszoegensis* shows some typical pantherine morphological characters such as a robust cranium, a well-developed sagittal crest, a marked nuchal crest (Mazák et al. 2011), and a frontoparietal suture located close to the postorbital constriction (Christiansen 2008). The ULg PA BR-II-81-146 cranium is slightly larger than a living leopard (*P. pardus*), jaguar (*P. onca*) or a snow leopard (*P. uncia*) but relatively smaller than the two largest extant species: the tiger (*P. tigris*) and the lion (*P. leo*) in terms of cranial length. The mandibular material from ‘La Belle Roche’ (Fig. 2C–E) consisted of three fragments: the first fragment (ULg-PA-BR-III-M13-79), is as fragment of right dentary with an extremely worn canine, but with a well preserved p3 and m1 (Fig. 2C). The second mandibular (ULg-PA-BRII-455) is a left dentary with a worn canine, fairly well preserved p3 and p4, but extremely damaged m1 (Fig. 2D). The last fragment (ULg-PA-20210823-01) completely misses the lower canine, but has a complete p3, p4 and an almost complete m1 which is only lacking the protocone (Fig. 2E). All ULg-PA-20210823-01 cheek teeth have been worn to some extent.

**Dorsal view**

The morphology and layout of fronto-nasal and fronto-maxillary sutures are often cited as good criteria for distinguishing between pantherines (e.g. Boule 1906). Those sutures are slightly discernible in *P. gombaszoegensis* from ‘La Belle Roche’ and appear to extend to approximately the same level, comparable to *P. leo* or *P. onca* (Fig. 4, no. 1). In all the pantherines examined, the fronto-nasal suture extends posteriorly to the posterior fronto-maxillary suture, except in *P. uncia*. In dorsal view the length and width of the snout varies in our dataset (Fig. 4, nos. 2 and 3).
Figure 4: Crania of the different pantherines used for comparison in dorsal view. A P. tigris MNHN-ZM-AC-1931-60, B P. onca MNHN-ZM 2006-641, C P. leo RMCA-34836, D P. uncia MNHN-ZO-AC 1917-18, E P. pardin RMCA-29292, F P. gombaszoegensis ULg PA BR-II-81-146. Scale bars represent 2cm. 1, posterior extent of fronto-nasal and fronto-maxillary sutures; 2, snout length; 3, snout width; 4, mastoid process; 5, postorbital process of frontal; 6, curvature of braincase; 7, occipital shape.
It is least elongate and broadest in *P. tigris*, with the length increasing and width decreasing through modern pantherines in a generally size-associated pattern from *P. leo*, to *P. onca*, to *P. pardus*, and finally to *P. uncia*. The snout morphology of *P. gombaszoegensis* fits the pattern, falling between that of the lion (*P. leo*) and jaguar (*P. onca*). In addition, the snout of *P. gombaszoegensis* is wide and straight, it does not appear constricted as in *P. tigris*, as has been described by previous authors (Jiangzuo & Liu 2020). The mastoid processes protrude slightly in dorsal view (Fig. 4, no. 4), more so than in *P. uncia* or *P. pardus*, with similarities to the larger modern species *P. leo* and *P. tigris*. The post-orbital region is of a similar relative size in all pantherines described except in *P. uncia*, in which it is incredibly large. The post-orbital process of the frontal of *P. gombaszoegensis* (Fig. 4, no. 5) is only moderately developed, as in the extant pantherines except for *P. uncia*, where it is more prominent. The post-orbital process of the zygomatic of ULg PA BR-II-81-146 is badly damaged on the left side, but is fairly well preserved on the right side and appears more developed compared to modern pantherines. A considerable portion of the frontal region has been altered; however, as best as can be determined, it was originally slightly depressed as observed in *P. onca* and *P. tigris* rather than flat as in *P. leo*, *P. atrox* (Merriam 1909; Martin & Gilbert 1978) and *P. spelaea* (Martin & Gilbert 1978). The braincase appears less rounded than in most of the extant pantherines, except for *P. tigris* (Fig. 4, no. 6). The occipital is moderately developed (Fig. 4, no. 7) and rather rounded; this is most similar to the condition of *P. leo* within living felines. The zygomatic arch has been deformed, but it appears relatively wide. The partial cranium CHA.100-f.8-73 from Château Breccia, which still has the left zygomatic arch in its original state and position, seems to confirm this assertion (see Plate 1 Argant & Argant 2011). The *P. gombaszoegensis* cranium from Château Breccia also possess zygomatic arches which are more triangular in shape, thus appearing more similar to larger modern pantherines (*P. tigris* and *P. leo*).  

**Lateral view**

In lateral view, the braincase appears less rounded than is usually observed in mid-sized pantherine species (Fig. 5, no. 1) (e.g. *P. onca*, *P. uncia*, *P. pardus*) with a narrow intertemporal region, more comparable to the larger species *P. tigris* and *P. leo*. In *P. pardus* and *P. onca*, the dorsal profile tends to form less pronounced curve from the nasal aperture to the sagittal crest (Fig. 5, no. 2), whereas there is an abrupt angle between the nasal and the frontal region in both *P. tigris* and *P. gombaszoegensis*. This is more pronounced than in *P. leo*, although this feature remains most pronounced in *P. uncia*. Previous work also noted a similarly slight angle between those two planes in the American lion *P. atrox*, comparable to modern lions (*P. leo*) (Merriam 1909). The nuchal crest (Fig. 5, no. 3) is slightly prominent in lateral view, much less so than in *P. tigris* but still more pronounced than in *P. onca*.
Figure 5: Crania of the different pantherines used for comparison in lateral view. A P. tigris MNHN-ZM-AC-1931-60, B P. onca MNHN-ZM 2006-641, C P. leo RMCA-34836, D P. uncia MNHN-ZO-AC 1917-18, E P. pardus RMCA-29292, F P. gombaszoegensis ULg PA BR-II-81-146. Scale bars represent 2cm. 1, shape of the braincase; 2, dorsal profile shape; 3, nuchal crest profile; 4, postorbital process of the zygomatic; 5, post-glenoid process; 6, jugular process.

Also, the postorbital process of the zygomatic is wider than in P. onca (Fig. 5, no. 4), and more similar to those of larger pantherine species. The postorbital process of the frontal of ULg PA BR-II-81-146 is completely missing on the left side and broken on the right side. Nevertheless, from its base it is clearly thicker than in P. uncia or P. pardus, but thinner than in the two largest extant species P. leo or P. tigris, ultimately being most similar to P. onca. The post-glenoid
process (Fig. 5, no. 5) is relatively large and easily discernible in lateral view. The jugular process (Fig. 5, no. 6) is ventrally directed, contrary to that observed for *P. uncia* and *P. pardus* where it is more posteriorly projected. The posterior face of the zygomatic arch is inclined, as in all the other pantherines.

Ventral view

The basicranial area of ULg PA BR-II-81-146 is quite well preserved and shows a clearly discernible opening of the oval foramen in ventral view (Fig. 6, no. 1), comparable to those of *P. onca*. The foramen is located at the level of the glenoid fossa, opening anterio-laterally as in other pantherines. This foramen is similar both in terms of size and shape to those of all pantherines, extant or fossil (see figures in Martin & Gilbert (1978); Sotnikova & Nikolskiy (2006); Christiansen & Harris (2009)), with the exception of the Miocene-Pliocene ‘*P.’ blythae’, where this foramen appears more developed than in more recent taxa (see Fig. 1 in Tseng et al. (2013)). In ventral view the choanae opens slightly posterior to the postorbital process of the zygomatic (see Fig. 2 in Sotnikova & Nikolskiy (2006)), with the opposite being true for ‘*P.’ blythae’ (anterior to the process; see Fig. 1 in Tseng et al. (2013)). The region between the auditory bullae (Fig. 6, no. 3) of *P. gombaszoegensis* is flat and wider than in modern large-size pantherines, which is also the case for the cranium from Château Breccia (Argant & Argant 2011). In *P. tigris* and *P. onca* the occipital condyles extend on the ventral part of the cranium and join but this is not the case in other extant pantherines (Fig. 6, no. 4). The junction between the condyles projects anteriorly in *P. tigris* and *P. onca*, but is even more anteriorly protruded in *P. gombaszoegensis*. The jugular process (Fig. 6, no. 5) is well marked in *P. gombaszoegensis*, falling between the morphology of *P. tigris* and that of *P. onca*, but is even more anteriorly protruded in *P. gombaszoegensis*. The jugular process (Fig. 6, no. 5) is well marked in *P. gombaszoegensis*, falling between the morphology of *P. tigris* and that of *P. onca*. The external occipital protuberance (Fig. 6, no. 6) is well pronounced, but the supraoccipital bone is almost invisible in ventral view. In extant pantherines, this condition is observed in *P. leo*, *P. pardus*, and *P. uncia*, although both *P. pardus* and *P. tigris* have a much more pronounced external occipital protuberance than *P. gombaszoegensis*. The palate is relatively wide in *P. gombaszoegensis*, and its shape resembles that of *P. uncia* and *P. onca*. However, the palate appears quite large; this represents one of the main differences between *P. gombaszoegensis* and *P. onca*, as noted by Argant and Argant (2011).
Figure 6: Crania of the different pantherines used for comparison in ventral view. A P. tigris MNHN-ZM-AC-1931-60, B P. onca MNHN-ZM 2006-641, C P. leo RMCA-34836, D P. uncia MNHN-ZO-AC 1917-18, E P. pardus RMCA-29292, F P. gombaszoegensis ULg PA BR-II-81-146. Scale bars represent 2cm. 1, opening of the oval foramen; 2, postorbital process of the zygomatic; 3, region between the auditory bullae; 4, occipital condyles extend on the ventral part of the cranium; 5, jugular process; 6, external occipital protuberance; 7, post-glenoid process; 8, shape of the mastoid process and the jugular process.
The post-glenoid process (Fig. 6, no. 7) is of moderate size, similar to *P. tigris* and *P. onca*, and more developed than in *P. leo*, *P. pardus* and *P. uncia*. The shape of the mastoid process and the jugular process (Fig. 6, no. 8) greatly resemble those of *P. tigris*, *P. onca* and *P. uncia* in ventral view; the jugular process is less developed than in *P. leo* but more so than *P. pardus*, whereas the mastoid process is much more developed than in *P. leo* and protrudes slightly more than in *P. pardus* or *P. uncia*.

**Posterior view**

The nuchal ridge of *P. gombaszoegensis* is relatively thick (Fig. 7, no. 1) and high relative to smaller pantherines (e.g. *P. pardus*; *P. onca*), comparable to *P. leo*. The occipital condyles (Fig. 7, no. 2) appear larger than in any extant pantherine (except for *P. uncia*), and also larger than in *P. atrox* (Figure 1 Merriam 1909) or in *P. spelaea* (Figure 2 and Figure 3 Sotnikova & Nikolskiy 2006). In posterior view, the zygomatic arches appear relatively wider than in *P. onca*, *P. uncia*, *P. leo* or *P. pardus* and more comparable to *P. tigris*. The intercondyloid notch is concave, similar to *P. tigris*, *P. pardus*, *P. uncia* or *P. onca*. The notch appears more deeply concave in *P. leo*. The gap between both occipital condyles (Fig. 7, no. 3) is moderately wide, comparable to that of *P. tigris*, *P. onca* and *P. uncia*, much wider than in *P. leo*, but narrower than in *P. pardus*. In posterior view the dorsal line of the *P. gombaszoegensis* appears slightly concave (Fig. 7, no. 4), more than in *P. onca* and *P. pardus*. It is similar to that observed in *P. tigris*. Yet this concavity is still much less pronounced than the condition in *P. leo* or *P. uncia*. The condyloid fossa (Fig. 7, no. 5) is marked and reassembles that of *P. onca*. It is somewhat less marked than in *P. tigris* and *P. uncia*, but more pronounced than those of the other pantherines studied. Both paracondylar process are missing. The largest extant pantherines (*P. tigris* and *P. leo*) exhibit a process above the supraoccipital bone, separated from the nuchal crest (Fig. 7, no. 6). This process seems to be present but is much smaller and closer to the nuchal crest in *P. onca*, whereas this process is completely absent in *P. gombaszoegensis*, reassembling the condition in the cranium of *P. pardus*.

**Anterior view**

In anterior view (Fig. S5) the zygomatic arches and the snout of *P. gombaszoegensis* appear wide. In particular, the snout of *P. gombaszoegensis* wide such that it obscures the infraorbital foramen, as in *P. tigris*. The infraorbital foramen is similar in terms of size and shape to the other pantherines. The lacrimal process is absent, or at least extremely reduced, as in *P. pardus*: this process is present and well developed in all other extant pantherines.
Figure 7: Crania of the different pantherines used for comparison in posterior view. A P. tigris MNHN-ZM-AC-1931-60, B P. onca MNHN-ZM 2006-641, C P. leo RMCA-34836, D P. uncia MNHN-ZO-AC 1917-18, E P. pardus RMCA-29292, F P. gombaszoegensis Ulg PA BR-II-81-146. Scale bars represent 2cm. 1, nuchal ridge; 2; occipital condyles 3, gap between both occipital condyles; 4, dorsal line of the cranium; 5, condyloid fossa; 6, process above the supraoccipital bone.
Upper dentition

Contrary to cranial traits, an extensive and well-documented comparative description of the dentition of *P. gombaszoegensis* has already been published (Jiangzuo & Liu 2020). Therefore, in this section we will focus on dental traits of the ULg-PA-BR-II-81-146 *P. gombaszoegensis* specimen which vary from previous descriptions, and highlight notable dental features of the specimen. Jiangzuo & Liu (2020) noted that upper dentition of *P. gombaszoegensis* differs mainly from that of *P. onca* by way of well-defined vertical grooves of the canine and less robust premolars (especially P3). In our sampling, the vertical groove is absent or indiscernible on the canine of ULg-PA-BR-II-81-146 from ‘La Belle-Roche’, and it is extremely reduced in the *P. onca* specimens we examined, whereas it is clearly present in the other pantherines. Thus, the presence of such a groove appears variable in *P. gombaszoegensis*, exactly as in *P. onca* (Seymour, 1989). Also, in our dataset, the P3 of *P. gombaszoegensis* appears more robust than in *P. onca* or *P. pardus*, but less so than in the other taxa, differing from previous descriptions (Jiangzuo & Liu 2020). Moreover, according to Jiangzuo & Liu (2020), the upper dentition of *P. gombaszoegensis* also differs from the lineages of *P. leo* and *P. spelaea* in several points, in particular a smaller and less marked anterior accessory cusps of P3. The third upper premolar of the specimen from ‘La Belle-Roche’ is damaged, but we can see that the anterior accessory cusp is undeniably extremely reduced (maybe the most reduced in our dataset with the exception of the living tiger, *P. tigris*). On the P3 of *P. gombaszoegensis*, the posterior cusp is large with a marked cingulum, but less clear than in extant pantherines.

*Panthera gombaszoegensis* ULg-PA-BR-II-81-146 also exhibits a well-developed P4 protocone, being less robust than in *P. leo*. The specimen from la Belle-Roche also seems to lack a well-defined ectoparastyle on the P4.

Results

Phylogenetic relationships of pantherines

A single most parsimonious tree was recovered in all analyses (Fig. 8A). These trees have a length of: 11.7 for K=3, 7.05952 for K=6, 5.06364 for K=9 and 3.94945 for K=12. As recently suggested by Goloboff, Torres, & Arias (2017), better results are obtained with a weaker concavity. Most of the taxa in our analysis were constrained based on the results of molecular analyses except for three fossil taxa: *P. gombaszoegensis*, *P. palaeosinensis* and ‘*P.’ blythae’. In the phylogenetic tree obtained through Bayesian inferences on a combined morphological and molecular dataset published by Tseng et al. (2013), *P. palaeosinensis* was recovered at the base of the Pantherinae, whereas in our topology it clusters with *N. nebulosa* which is not considered as a pantherine.
A

2 Frontoparietal suture
2:0 posterior to postorbital constriction
2:1 at the postorbital constriction

4 Parietal process at frontoparietal suture
20:1 present

20 Frontonasal depression
20:0 absent
20:1 present

39 Distance between bulla and glenoid ridge
39:0 wide
39:1 narrow
However, the node uniting *P. palaeosinensis* and *N. nebulosa* present a low symmetric resampling value (32), despite these two taxa being united by two synapomorphies: an intermediate nasal width at aperture (character 14:1), and a mandible depth anterior to p3 that is deeper than the depth at p4/m1 (character 44:1). In our analysis, and as in Tseng et al. (2013), ‘P.’ *blythae* forms a clade with *P. uncia* which is supported by a symmetric resampling value of 55.

The most interesting result of our phylogenetic analyses is that *P. gombaszoegensis* is systematically recovered as closer to *P. tigris* than to *P. onca*, no matter the importance of homoplastic characters. The *P. gombaszoegensis + P. tigris* clade is supported by four unambiguous but homoplastic synapomorphies: a frontoparietal suture posterior to the postorbital constriction (character 2:0), the presence of a parietal process at the dorsal frontoparietal suture (character 4:1), and the presence of a frontonasal depression (character 20:1) and finally the wide distance between the anterior bullae and the glenoid ridge (character 39:0) is also observed in *P. leo*. This very last character states might be influenced by allometry as states 0 was observed in the three largest species (*P. gombaszoegensis, P. tigris, P. leo*) but a larger dataset of large taxon and some correlations would be needed to test this hypothesis.

Multivariate morphometric analysis: upper dentition

A first two-dimensional morphospace was retrieved using a Principal Coordinates Analysis (PCoA) (Fig. 9A). This PCoA retrieved 19 axes (Fig. S6A), with the two first axes explaining 22.44% and 5.55% of variance respectively. The first axis is mainly influenced by the aspect ratio of the canine (CH/CL), the length of P3 (P3L/CL) and both measurements taken on the P4 (P4L/CL and P4W/CL), whereas the second axis is more influenced by the canine width (CW/CL) and the P3 width.
Figure 9: Results of the multivariate morphometric analyses. A, PCoA on the upper dentition dataset; B, NMDS on the upper dentition dataset; C, PCoA on the lower dentition dataset; D, NMDS on the lower dentition dataset; E, PCoA on the cranium dataset; F, NMDS on the cranium dataset; G, non-metric fit of the two-dimensional NMDS on the original dissimilarities for the upper dentition dataset (k = 2; stress = 0.08330135); H, non-metric fit of the two-dimensional NMDS on the original dissimilarities for the lower dentition dataset (k = 2; stress = 0.03082973); I, non-metric fit of the two-dimensional NMDS on the original dissimilarities for the cranium dataset (k = 2; stress = 0.1256406). Animal silhouettes were obtained from PhyloPic (phylopic.org). Image credits: Manabu Sakamoto (P. onca) and Sarah Werning (P. tigris).

While all pantherines specimens are well spread on the first axis, they strongly overlap on the second with only a few P. gombaszoegensis and one P. tigris specimens occupying the lowest portion of the axis and P. uncia showing the highest PC values. At first sight Axis 1 might seem to be well sorted by size; however, P. gombaszoegensis occupy much lower values than P. onca while being larger. Panthera onca overlap P. leo in the morphospace while P. pardus, which is supposed to be part of the ‘lion group’ in our phylogeny is closer to P. uncia.

The closest taxa to P. gombaszoegensis on the first axis are P. pardus and P. leo specimens. PERMANOVA retrieved a significant difference between P. onca and P. gombaszoegensis in terms of upper dentition proportions, with a p-value of 0.001 (FDR-corrected, p ≤ 0.003). However, according to our

PERMANOVA results P. gombaszoegensis differs from any of the extant pantherines in terms of dental proportions (P. tigris 0.002, FDR-corrected p ≤ 0.001; P. pardus 0.001, FDR-corrected p ≤ 0.001; P. leo 0.001, FDR-corrected p ≤ 0.001 and P. uncia 0.006, FDR-corrected p ≤ 0.001).

The only taxa to exhibit similar values to P. gombaszoegensis on Axis 2 is one specimen of P. tigris which could be explained by their phylogenetic affinities that we highlighted previously but may also be explained by intraspecific variation. The Pagel’s Lambda for the first axis was close to zero showing there is no phylogenetic signal on this axis (Table 1).

<table>
<thead>
<tr>
<th></th>
<th>Pagel’s Lambda</th>
<th>logL Lambda</th>
<th>p-value</th>
<th>Phylogenetic signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lower dentition</td>
<td>6.6107e-05</td>
<td>-11.7486</td>
<td>1</td>
<td>None</td>
</tr>
<tr>
<td>Upper dentition</td>
<td>6.6107e-05</td>
<td>-11.0724</td>
<td>1</td>
<td>None</td>
</tr>
<tr>
<td>Cranium</td>
<td>6.6107e-05</td>
<td>-6.31618</td>
<td>1</td>
<td>None</td>
</tr>
</tbody>
</table>
A second two-dimensional morphospace was performed using a non-metric multidimensional (NMDS) (Fig. 9B). In the NMDS plot, *P. gombaszoegensis* clearly plots apart from other pantherine species. The pantherine taxa occupy a wide range of values on the NMDS1 while they are confined to values between -1 and 2 on the second, the only exception being *P. gombaszoegensis* occupying the lower portion of the axis NMDS2. The non-metric fit of the two-dimensional NMDS on the original dissimilarities confirms that a two-dimensional plot represents the inter-specimen morphological distances very well (R² = 0.993 for k=2, stress value = 0.08330135) (Fig. 9G).

**Multivariate morphometric analysis: lower dentition**

Our PCoA computed on the 8 ratios from the lower dentition (Fig. 9C) retrieved 18 axes (Fig. S6B), with the first two PCo axes explaining 42.48% and 12.16% of variance respectively. The placement of specimens along the first axis is mainly influenced by measurements taken on the p3, p4 and m1 (p3l/cl, p3w/cl, p4l/cl, p4w/cl, m1w/cl and m1l/cl), whereas distribution across the second axis is mainly influenced by the check teeth length (ctl/cl) and the diastema length (diastema/cl). The three *P. gombaszoegensis* specimens included in this study are located in a region of morphospace with negative values on both axes, likely due to a high canine length compared to the measurements taken on the check teeth and the diastema. The closest species to *P. gombaszoegensis* in the morphospace are *P. onca* and *P. pardus* whereas the most dissimilar species are *P. tigris* and *P. uncia*. All the pantherines species are relatively occupies a relatively large part of the morphological space on the plot, except for *P. pardus* and *P. onca*. Interestingly, *P. spelaea* and *P. leo* are well separated from each other in the morphospace, with *P. spelaea* exhibiting negative values on the first axis while the modern lion shows positive values (yet values on the second axis are extremely similar). It should also be noted that the single specimen of *P. palaeosinensis* is comprised within the morphological space occupied by *P. pardus*. Resemblances between the lower dentition of *P. onca* and *P. gombaszoegensis* is corroborated using PERMANOVA, which did not retrieve any significant differences between those two taxa (p-value=0.4). PERMANOVA also did not retrieve any statistical difference between the lower dentition of *P. tigris* and *P. gombaszoegensis* (p-value=0.6), nor from *P. pardus* (p-value=0.1) but rather in *P. leo* (p-value=0.04) and *P. uncia* (p-value=0.01) although FDR corrections showed that those last two significant p-values were statistical artefActs (FDR-corrected p ≤ 0.1 for *P. leo* and p ≤ 0.3 for *P. uncia*). The Pagel’s Lambda for the first axis was close to zero showing there is no phylogenetic signal on this axis (Table 1).

The pattern observed from the NMDS plot (Fig. 9D) is somewhat similar, although there are still some differences worth mentioning: *P. onca* and *P.
**gombaszoegensis** specimens almost completely overlap in the morphological space; *P. palaeosinensis* does not overlap with *P. pardus*; *P. spelaea* and *P. leo* are clearly separated from each other. The modern leopard (*P. pardus*) is clearly separated from *P. gombaszoegensis* in the morphospace, but the modern lion (*P. leo*) is relatively closer in this analysis. Again, the non-metric fit of the two-dimensional NMDS on the original dissimilarities shows a high $R^2$ of 0.996 for $= 2$, stress = 0.03082973 (Fig. 9H).

**Multivariate morphometric analysis: cranium**

The PCoA computed on the 12 cranial ratios (Fig. 9E) retrieved 15 axes (Fig. S6C), the first two explaining 26.71% and 17.38% of variance respectively. The first axis is mainly influenced by the post-orbital width (POW/COL), the braincase width (BrW/ COL), the palatal length (PaL/ COL), the basicranium length (BaL/COL) and the condylobasal width (CBW/COL) while the axis 2 is mostly influenced by the nasal width (NaW/COL), the mastoid process width (MaPW/COL), the orbital process length (OPL/COL) and the temporal fossa length (TFL/COL). Pantherines occupy a relatively large part of the morphological space on both PCo axes based on cranial ratios. *P. gombaszoegensis* is recovered close to *P. palaeosinensis* and ‘*P.’ blythae’ on the second axis of both the PCoA (Fig. 9E) and the NMDS2 (Fig. 9F). Cranial morphology of *P. gombaszoegensis* seems to be most removed from *P. uncia*. The Pagel’s Lambda for the first axis was close to zero showing there is no phylogenetic signal on this axis (Table 1). The pattern of morphospace occupation for the cranial dataset appears more similar between the PCoA and the NMDS than for the two previous datasets, although the non-metric fit of the two-dimensional NMDS on the original dissimilarities appears slightly lower than for the two previous datasets ($R^2 = 0.984$) (Fig. 9I).

**Allometry**

Our linear regressions between the cranial or canine length (upper and lower) and their respective scores on the first axis of the PCoA (Fig. S8-S10) show a clear allometric component in the features studied her, which is more marked ($R^2_{adj} = 0.82$ for the upper dentition; $R^2_{adj} = 0.84$ for the lower dentition) for dental measurements than for the cranium ($R^2_{adj} = 0.48$). However, this trend is not retrieved in the other PCoA axes. Still, it is clear than *P. gombaszoegensis* show lower values on PCo axis 1 for the upper dentation dataset than similar sized *P. onca* specimens in our dataset. The allometric influence is present in the first PCo axis of our analyses, but is not prevalent in subsequent axes, with adjusted $R^2$ values dropping lower than 0.1 for PCo axes 2–10 in each morphometric dataset (Fig. S8–S10).
Discussion

Several authors have pointed out the resemblances between *P. onca* and *P. gombaszoegensis* (e.g. Argant & Argant 2011; Marciszak 2014; Stimpson *et al.* 2015; Jiangzuo & Liu 2020), leading to the characterisation of the latter as the ‘European jaguar’ or ‘Eurasian jaguar’ (e.g. Argant and Argant 2011; Hemmer *et al.* 2001, 2010; Hankó 2007; Mol *et al.* 2011; Stimpson *et al.* 2015). Bonifay (1971) mentioned its resemblance to *P. leo*, and other felids, based on cranial fragments and upper teeth of young adults. Then, O'Regan (2002) noted that it was more similar to the modern tiger (*P. tigris*) and jaguar (*P. onca*) than to any extant or extinct lion (*P. leo*, *P. spelaea*, *P. atrox*) or leopard (*P. pardus*). Later, its resemblance with the modern tiger were forgotten as authors assigned it as a subspecies of *P. onca* (Hemmer *et al.* 2001, 2010; Hankó 2007; Mol *et al.* 2011; Stimpson *et al.* 2015). Our analysis (and that of Tseng *et al.* (2013)) retrieved *P. palaeosinensis* as a basal pantherine, with our topology placing this taxa outside of crown *Panthera* and as sister to *Neofelis* (Fig. 8A). Our analyses recognise the presence of a stem tiger (*P. gombaszoegensis*) in the Pleistocene of Europe which contradicts the previous hypotheses concerning the origin of *P. tigris*. *Panthera gombaszoegensis* was theorised to have migrated to North America from eastern Asia during the Early Pleistocene through Beringia, following different potential dispersal scenarios (Kurtén 1973; Argant & Argant 2011; Jiangzuo & Liu 2020). However, the affinities between *P. gombaszoegensis* and *P. onca* has always been unclear, especially given that *P. onca* is endemic to the American continent (Seymour 1989), with its first uncontested fossil remains being found in the Hamilton Cave (West Virginia) dating from 820-850 kya (Repenning *et al.* 1995). Yet, *P. gombaszoegensis* has never been identified on the American continent, nor even in eastern Asia. The first fossil that can be clearly attributed to *P. tigris* is approximately 850 kyr old (Hemmer 1971), and according to different sources, the evolutionary origin of the modern tiger is presumably located in the north of China (Hemmer 1981,
The Haro Quarry river fauna (including some *P. gombaszoegensis* specimens) is estimated to be between 1.77 and 1.07 Myr old (Jiangzuo & Liu 2020), indicating that *P. gombaszoegensis* likely reached Asia just before the first *P. tigris* appeared. This is in accordance with the results of our phylogenetic analysis, and provides a more coherent scenario in terms of geographic dispersion through time (see Fig. 1a). Also, Jiangzuo & Liu (2020) already recognized that the specific assignments of some fossils to *P. gombaszoegensis* in eastern Asia was complex, due to the presence of a similar-sized pantherine (early *P. tigris*), clearly pointing towards an imbricated fossil record of *P. gombaszoegensis* and *P. tigris*.

*P. gombaszoegensis* was thought to be significantly larger than the extant jaguar (Argant & Argant 2011), but also smaller than most extant tigers (Jiangzuo & Liu 2020). Some living tiger subspecies are thought to significantly overlap *P. gombaszoegensis* when considered only dental size (e.g. *P. tigris sumatrae*, *P. tigris sondaica* or *P. tigris balica* see Mazák & Groves (2006)). As evolutionary allometry can affect bone shape, and thus influence the results of morphometrics analyses (Klingenberg 1996), and different studies have already highlighted the importance of considering evolutionary allometry when dealing with felids cranial shape and proportions (e.g. Slater & Van Valkenburgh 2009; Tamagnini, Meloro, & Cardini 2017), it may not be surprising that *P. gombaszoegensis* presents similarities with both the extant tiger (*P. tigris*) and the extant jaguar (*P. onca*) with regards to size, falling in between those two taxa. Nevertheless, we describe in this contribution a medium-sized specimen of *P. gombaszoegensis* (large female or small male) which varies in numerous features from *P. onca*. The differences we observe cannot be solely explained by allometric variation. Our morphospaces do not reflect the phylogeny we obtained (see Pagel’s Lambda and p-values in Table 1) which could mean that our ratios are more driven by ecological factors than by the phylogenetic relationships between our taxa. Key differences were revealed between the upper and lower dentition proportions for *P. gombaszoegensis*: our morphospaces corroborate the similarities in lower dentition between *P. gombaszoegensis* and *P. onca*, but these are not recovered for the upper teeth or for the cranium (though with only a single cranial specimen of *P. gombaszoegensis*). This discovery reiterates that dental characters should be treated carefully when dealing with taxa identification or phylogenetic reconstructions (especially those from the mandible), since morphological adaptations for feeding may obscure phylogenetic affinities (as discussed in Jernvall 2000; Jernvall & Jung 2000; Naylor & Adams 2001; Dávalos *et al.* 2014; Sansom *et al.* 2017; Billet & Bardin 2019). Indeed, the differences between upper and lower dentition observed in *P. gombaszoegensis* were already
discussed as an example of mosaic evolution within this taxon (Hemmer 1981). Despite occlusal surfaces of mammalian teeth supposedly corresponding perfectly, modularity is present within mammalian dentition (Hemmer 1981), and other morphometric analysis already retrieved slightly different results for lower and upper dentition (e.g. Leroy et al. 2004; Bever 2005; Dumbá et al. 2022). Moreover, it seems that variations in dental proportions do not negatively affect the efficacy of occlusion (Ungar 2010).

The distinctiveness between the craniodental form of *P. gombaszoegensis* and *P. onca*, also has interesting palaeoecological implications. A number of authors have considered *P. gombaszoegensis* as a ‘generalist’ taxon (i.e. which would be able to hunt a wide spectrum of prey), certainly more so than *P. onca* (Jiangzuo & Liu 2020; Marciszak & Lipecki 2021) and comparable to the notably ubiquitous *P. pardus* (Marciszak 2014). The postcranial material available for *P. gombaszoegensis* has not recovered great similarities to *P. onca*. Indeed some similarities are clearly observable between these two taxa, but there is a large size difference and widespread resemblance with the postcrania of various pantherines (O’Regan 2002; Argant & Argant 2011). Argant & Argant (2011) insisted on the ability of *P. gombaszoegensis* to adapt to the harsh climates of glacial periods, as demonstrated by modern Siberian tigers (*Panthera tigris altaica*). However, *P. gombaszoegensis* has also often been described as an ecological equivalent of the modern jaguar, considered as a forest dweller with strong penchant for open water (Hemmer 1971; Hemmer et al. 2001). Marciszak and Lipecki (2021) suggested that, based on the observations published by Jiangzuo and Liu (2020), the moderately robust dentition of *P. gombaszoegensis* (compared to *P. onca*) would reflect its prey preferences. Marciszak and Lipecki (2021) also argued that the marked vertical grooves on the upper canines of *P. gombaszoegensis* and the poor development or even absence of this feature in *P. onca* are probably correlated with habitat and prey preference. However, we did not observe any significant variation in the robustness of the upper check teeth between *P. gombaszoegensis* and *P. onca*, and the vertical groove was not observed on the cranium of the *P. gombaszoegensis* ULg-PA-BR-II-81-146 specimen. This may be explained by intraspecific variation, and, if true, may further underline that those traits should not be used to infer any palaeoecological interpretations (and certainly not phylogenetic affinity). A wide prey spectrum could explain the presence of *P. gombaszoegensis* in various habitats (Marciszak and Lipecki 2021). Our observations show that *P. gombaszoegensis* shares similarities with not one but several different pantherines: both *P. pardus* and *P. leo* for the upper dentition; *P. onca* for the lower dentition; and generally larger pantherines for the cranium as a whole. The morphological evidence presented in
this study therefore advocates for an ‘ecological generalist’ niche for *P. gombaszoegensis*, which is highly consistent with the large geographical range of the taxon throughout the Pleistocene (cf. Fig. 1A and S1).

**Conclusions**

*Panthera gombaszoegensis* is a ubiquitous felid in the Pleistocene of Eurasia. However, many aspects of its morphology are mysterious, owing to the incompleteness of most published remains. Our examination of the near-complete cranium of the *P. gombaszoegensis* ULg-PA-BR-II-81-146 specimen revealed similarities with the extant jaguar *P. onca*, but also highlighted common morphological features with other extant members of Pantherinae. We also found a large number of evidences that support phenotypic differences between *P. onca* and *P. gombaszoegensis*, that comfort the status of *P. gombaszoegensis* as a valid pantherine species, rather than as a subspecies of jaguar. Our morphometric analyses provide a potential explanation for the historic interpretation of *P. gombaszoegensis* as a jaguar, with the lower dentition being extremely similar to that of *P. onca*. Nevertheless, the upper dentition and cranial proportions are far more variable and do not suggest strong affinity to *P. onca* as previously thought. The behaviour of *P. gombaszoegensis* has often been inferred based on its supposed relationship with *P. onca*; our description and analyses show a combination of different pantherine traits, advocating for a more ‘generalist’ taxa. Finally, our phylogenetic analyses place *P. gombaszoegensis* as the sister taxon to the extant tiger (*P. tigris*), further from the extant jaguar (*P. onca*). This phylogenetic hypothesis simplifies the biogeographic dispersal scenario of this taxon, which is only found in Eurasia, whereas *P. onca* is endemic to the American continent. Under this new scenario, *P. gombaszoegensis* reached Southern Asia approximately in between 1 and 1.8 Myr and did not reach North America. The close affinity between *P. tigris* and *P. gombaszoegensis* thus enables us to propose the presence of a stem tiger in the Pleistocene of Europe, casting doubt on the Asian origin of *P. tigris* and abolishing the misnomer of *P. gombaszoegensis* as a “European jaguar”

**Data archiving statement**

Data archiving statement. 3D models of the fossil material from the palaeontology collections of the University of Liège are available on MorphoSource ([https://www.morphosource.org/projects/000445179](https://www.morphosource.org/projects/000445179)); Appendix S5 lists all associated DOIs. Associated data (Excel file containing the measurements and ratios, nexus file containing the phylogenetic matrix) and the R script used are provided as Supporting Information, and are available on the University of Liège institutional repository ORBi ([https://hdl.handle.net/2268/294237](https://hdl.handle.net/2268/294237)) and MorphoBank
Authors’ contributions
NC designed the study. NC and MM collected scan data, photographs and drafted the figures. NC took the measurements and ran the morphometric analyses. VF wrote the TNT scripts for the phylogenetic analyses and NC performed the analyses. All authors contributed to writing the manuscript, and gave final approval for publication.

Supporting information
Additional Supporting Information can be found online (https://doi.org/10.1111/spp2.1464):

Appendix S1. Includes TNT script, Figures S1–S10 and Tables S1–S9.
Appendix S2. R script.
Appendix S3. Nexus file containing the phylogenetic matrix.
Appendix S4. README file to explain data.
Appendix S5. List of MorphoSource DOIs associated with this publication.
Data S1. Measurements used in this study.

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References


CORDY, J.-M., BASTIN, B., DEMARET-FAIRON, M., EK, C., GEERAERTS,


GOLOBOFF, P. A. and CATALANO, S. A. 2016. TNT version 1.5, including a full implementation of phylogenetic morphometrics. Cladistics, 32, 221–238.

———, TORRES, A. and ARIAS, J. S. 2017. Weighted parsimony
outperforms other methods of phylogenetic inference under models appropriate for morphology.


——— and LIPECKI, G. 2021. Panthera gombaszoegensis (Kretzoi, 1938) from Poland in the scope of the species evolution.


MOL, D., LOGCHEM, W. Van and VOS, J. De. 2011. New record of the


SOTNIKOVA, M. S. 1990. Pleistocene mammals from Lakhuti, Southern Tajikistan, USSR. *Quatärpaläontologie*, 237–244.


