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Kévin Le Verger, Laureano R González Ruiz, Guillaume Billet. Comparative anatomy and phylogenetic contribution of intracranial osseous canals and cavities in armadillos and glyptodonts (Xenarthra, Cingulata). *Journal of Anatomy*, 2021, pp.1-30. 10.1111/joa.13512 . hal-03291344

**HAL Id: hal-03291344**

<https://hal.sorbonne-universite.fr/hal-03291344v1>

Submitted on 19 Jul 2021

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## ORIGINAL PAPER

# Comparative anatomy and phylogenetic contribution of intracranial osseous canals and cavities in armadillos and glyptodonts (Xenarthra, Cingulata)

Kévin Le Verger<sup>1</sup>  | Laureano R. González Ruiz<sup>2</sup> | Guillaume Billet<sup>1</sup>

<sup>1</sup>Museum national d'Histoire naturelle, Centre de Recherche en Paléontologie – Paris, UMR 7207 CR2P MNHN/CNRS/UPMC, Sorbonne Universités, Paris, France

<sup>2</sup>Laboratorio de Investigaciones en Evolución y Biodiversidad (LIEB-FCNyCS sede Esquel, UNPSJB) y Centro de Investigaciones Esquel de Montaña y Estepa Patagónica (CIEMEP), CONICET, Universidad Nacional de La Patagonia San Juan Bosco (UNPSJB), Chubut, Argentina

**Correspondence**

Kévin Le Verger, Museum national d'Histoire naturelle, Centre de Recherche en Paléontologie – Paris, UMR 7207 CR2P MNHN/CNRS/UPMC, Sorbonne Universités, Paris, France.  
Email: kevin.le-verger@edu.mnhn.fr

**Abstract**

The evolutionary history of the Cingulata, as for many groups, remains a highly debated topic to this day, particularly for one of their most emblematic representatives: the glyptodonts. There is no consensus among morphological and molecular phylogenies regarding their position within Cingulata. As demonstrated by recent works, the study of the internal anatomy constitutes a promising path for enriching morphological matrices for the phylogenetic study of armadillos. However, internal cranial anatomy remains understudied in the Cingulata. Here we explored and compared the anatomy of intracranial osseous canals and cavities in a diverse sample of extant and extinct cingulates, including the earliest well-preserved glyptodont crania. The virtual 3D reconstruction (using X-ray microtomography) of selected canals, that is, the nasolacrimal canal, the palatine canal, the sphenopalatine canal, the canal for the frontal diploic vein, the transverse canal, the orbitotemporal canal, the canal for the capsuloparietal emissary vein and the posttemporal canal, and alveolar cavities related to cranial vascularization, innervation or tooth insertion allowed us to compare the locations, trajectories, and shape of these structures and to discuss their potential interest for cingulate systematics. We tentatively reconstructed evolutionary scenarios for eight selected traits related to these structures in which glyptodonts often showed a close resemblance to pampatheres, to the genus *Proeutatus*, and/or to chlamyphorines. This latter pattern was partly congruent with recent molecular hypotheses, but more research is needed on these resemblances and on the potential effects of development and allometry on the observed variations. Overall, these comparisons have enabled us to highlight new anatomical variation that may be of great interest to further explore the evolutionary history of cingulates and the origins of glyptodonts on a morphological basis.

**KEYWORDS**

alveolar cavities, cingulata, comparative anatomy, cranial canals, evolutionary scenarios, glyptodont, phylogeny

## 1 | INTRODUCTION

The peculiar morphology of xenarthrans, which include armadillos (Cingulata), sloths (Folivora), and anteaters (Vermilingua), has aroused the curiosity of anatomists since the end of the 18th century (Cuvier, 1798). Compared to other mammals, the sum of their unique characteristics has led them to be considered by some previous workers, the most unusual mammals (Vizcaíno & Loughry, 2008; for a summary see Superina & Loughry, 2015). They represent the only known mammals with a carapace formed by a mosaic of osteoderms (fixed and mobile) covering the head, body, and tail (Vizcaíno & Loughry, 2008). Cingulates include extant and extinct armadillos, among them the four extant subfamilies Dasypodinae, Euphractinae, Tolypeutinae, and Chlamyphorinae, as well as the extinct glyptodonts, a group of herbivorous and large-bodied mammals appearing in the late Eocene and disappearing at the beginning of the Holocene (Delsuc et al., 2016; see also Gaudin & Lyon, 2017 for an alternative classification). The phylogenetic position of glyptodonts within Cingulata is particularly difficult to resolve based on morphological data because of their highly specialized anatomy and the large morphological gap separating them from extant armadillos (Burmeister, 1874; Carlini & Zurita, 2010; Fariña et al., 2013). This difficulty was recently emphasized by ancient DNA studies showing that phylogenetic analyses of their mitogenomes support a position for glyptodonts at odds with that proposed in prior morphological studies (Billet et al., 2011; Delsuc et al., 2016; Engelmann, 1985; Gaudin & Wible, 2006; Herrera et al., 2017; Mitchell et al., 2016).

While it has long been accepted that glyptodonts were closely related to extant armadillos (Huxley, 1864), their exact relationships with the latter were never clearly resolved on the basis of morphological data. Flower (1882) initially proposed that glyptodonts were the ancestors of extant armadillos but Ameghino (1884, 1889) disputed this hypothesis, arguing that their great dental and postcranial complexity, as described by Owen (1839), was evidence of their derived nature. The nested position of glyptodonts among extant armadillos has been supported by many authors since then, including by recent phylogenetic analyses using morphological data (Billet et al., 2011; Castellanos, 1932, 1959; Fernicola, 2008; Gaudin & Wible, 2006; Herrera et al., 2017; Hoffstetter, 1958; Patterson & Pascual, 1972). However, the consensus stops here. Engelmann (1985) proposed a close relationship of glyptodonts with the extinct Eutatini (*Proeutatus* and *Eutatus*) and pampatheres, whereas the analysis of Gaudin and Wible (2006) retrieved glyptodonts in an apical position nested within a large clade gathering euphractines, chlamyphorines, the extinct pampatheres, and some extinct eutatines. A similar hypothesis was also supported by other studies, but with much variations in the composition of this large apical clade (Billet et al., 2011; Fernicola, 2008; Fernicola et al., 2017; Herrera et al., 2017; Porpino et al., 2010). Most of these studies, however, unvaryingly proposed a close relationship between the extinct giant pampatheres and the glyptodonts, in agreement with Patterson and Pascual (1972). Thanks to the progress made in ancient DNA studies and the recent extinction of glyptodonts—at the beginning of the

Holocene (Messineo & Politis, 2009)—the complete mitochondrial genome of the Pleistocene glyptodont *Doedicurus* (Burmeister, 1874) was successfully assembled, which gave rise to a completely new phylogenetic hypothesis (Delsuc et al., 2016; Mitchell et al., 2016). Based on analyses of their mitogenomes, glyptodonts were nested within extant armadillos and represented the sister group of a clade formed by chlamyphorines and tolpeutines (Delsuc et al., 2016; Mitchell et al., 2016). Although analyses of morphological matrices using a molecular backbone constraint can detect morphological characters congruent with the molecular pattern (Mitchell et al., 2016), the high level of disagreement between the two data partitions may call into question the phylogenetic signal provided by morphological matrices on cingulates.

External cranial and postcranial skeletal anatomy are already well known for many groups within Cingulata, but the internal cranial anatomy remains understudied. However, its study has already provided data of systematic interest for both the extant and extinct xenarthran diversity (Billet et al., 2015; Billet et al., 2017; Boscaini et al., 2020; Boscaini et al., 2018; Fernicola et al., 2012; Tambusso & Fariña, 2015a; Tambusso & Fariña, 2015b; Tambusso et al., 2021; Zurita et al., 2011). Intracranial osseous canals provide important pathways for innervation and vascularization of the head (Evans & de Lahunta, 2012). The diversity and phylogenetic signal of their intracranial trajectories are poorly known, as these hidden structures are rarely described in mammals, in general, including xenarthrans (e.g., Wible & Gaudin, 2004). In contrast, their external openings on the cranium are often described and scored in phylogenetic matrices (e.g., 37/131 of cranial characters—Gaudin & Wible, 2006).

Here we present a comparative investigation of intracranial osseous canals and cavities in a diverse sample of extant and extinct cingulates, including the earliest well-preserved glyptodont crania, using X-ray microtomography. The 3D virtual reconstruction of selected canals and cavities related to cranial vascularization, innervation, or upper tooth insertion enabled us to compare the locations, trajectories, and shape of each homologous structure and discuss their potential interest for cingulate systematics. We tentatively reconstructed the evolutionary history of these traits using a molecularly constrained phylogeny, and we discuss the potential effects of development and allometry on their variation. These comparisons allowed us to propose new potential characters that may be used to further explore the origins of glyptodonts within cingulates on a morphological basis.

## 2 | MATERIALS AND METHODS

### 2.1 | Sampling

We examined the crania of 33 extant and extinct xenarthran specimens. Two specimens belonging to the Pilosa (sloths and anteaters) were chosen as outgroups. The remaining 31 crania belonged to Cingulata (Table 1). This sample included specimens representing the nine extant genera as well as three small developmental series

TABLE 1 List of specimens. Symbol: †, extinct species; \*, perinatal stage; \*\*, juvenile; \*\*\*, subadult; \*\*\*\*, adult; (i), author of the genus

Family/Subfamily	Species	Institutional number	Locality	Period	References
Bradyrodidae	<i>Bradyrodus tridactylus</i>	MNH ZM-MO-1999-1065	French Guiana; Petit saut rive droite	Extant	Linnæus, 1758
Myrmecophagidae	<i>Tamandua tetradactyla</i>	NHMUK ZD-1903.77.135	Brazil; Mato Grosso; Chapada	Extant	Linnæus, 1758
Peltephilidae	<i>Peltephilus pumilus</i> †	YPM-VPPU 15391	Argentina; Santa Cruz; Santa Cruz Formation; Coy Inlet	Early Miocene	Ameghino, 1887
Dasypodinae	<i>Dasydus novemcinctus</i> *	AMNH 33150	Colombia; Quindí; Salento	Extant	Linnæus, 1758
Dasypodinae	<i>Dasydus novemcinctus</i> **	AMNH 133261	Brazil; Goiás; Anapolis	Extant	Linnæus, 1758
Dasypodinae	<i>Dasydus novemcinctus</i> ***	AMNH 133328	Brazil; Mato Grosso do Sul; Maracaju	Extant	Linnæus, 1758
Dasypodinae	<i>Dasydus novemcinctus</i> ****	AMNH 255866	Bolivia; Beni; Cercado	Extant	Linnæus, 1758
Dasypodinae	<i>Stegotherium tauberi</i> †	YPM-VPPU 15565 (type)	Argentina; Santa Cruz; Santa Cruz Formation; Killik Aike Norte	Early Miocene	González Ruiz & Scialto-Yané, 2008
Euphractinae	<i>Prozaedyus exilis</i> †	YPM-VPPU 15579	Argentina; Santa Cruz; Santa Cruz Formation; Killik Aike Norte	Early Miocene	Ameghino, 1887
Euphractinae	<i>Prozaedyus proximus</i> †	YPM-VPPU 15567	Argentina; Santa Cruz; Santa Cruz Formation; Coy Inlet	Early Miocene	Ameghino, 1887
Euphractinae	<i>Euphractus sexcinctus</i>	AMNH 133304	Brazil; Goiás; Anapolis	Extant	Linnæus, 1758
Euphractinae	<i>ChaetophRACTUS villosus</i>	AMNH 173546	South America; No more precision	Extant	Desmarest, 1804
Euphractinae	<i>Zaedyus pichiy</i> *	ZMB MAM-49039	Argentina; Oso Marino	Extant	Desmarest, 1804
Euphractinae	<i>Zaedyus pichiy</i> **	MHNG 1627.053	Argentina; Chubut; Punta Ninfas	Extant	Desmarest, 1804
Euphractinae	<i>Zaedyus pichiy</i> ***	MHNG 1276.076	Argentina; Chubut; Fof Cahuel	Extant	Desmarest, 1804
Euphractinae	<i>Zaedyus pichiy</i> ****	FMNH 23810	Argentina; Chubut; No more precision	Extant	Desmarest, 1804
Euphractinae	<i>Doellotatus</i> sp.†	UF 260533	Bolivia; unnamed formation; Potosí; Quebrada Honda	Middle Miocene	Bordas, 1932 (i)
Tolypeutinae	<i>Cabassous uncinatus</i> **	NBC ZMA-MAM-26326.A	Suriname; No more precision	Extant	Linnæus, 1758
Tolypeutinae	<i>Cabassous uncinatus</i> ***	AMNH 133386	Brazil; Mato Grosso do Sul; Maracaju	Extant	Linnæus, 1758
Tolypeutinae	<i>Cabassous uncinatus</i> ****	MNH CG-1999-1044	French Guiana; No more precision	Extant	Linnæus, 1758
Tolypeutinae	<i>Tolypeutes matacus</i>	FMNH 28345	Brazil; Mato Grosso; Descalvado	Extant	Desmarest, 1804
Tolypeutinae	<i>Priodontes maximus</i>	AMNH 208104	Zoo; No more precision	Extant	Kerr, 1792
Chlamyphorinae	<i>Chlamyphorus truncatus</i>	AMNH 5487	Argentina; Mendoza; No more precision	Extant	Harlan, 1825
Chlamyphorinae	<i>CalyptophRACTUS retusus</i>	NMNH 283134	Bolivia; Santa Cruz; No more precision	Extant	Burmeister, 1863
Eutatini	<i>Proeutatus lagenai</i> †	YPM-VPPU 15613	Argentina; Santa Cruz; Santa Cruz Formation; Coy Inlet	Early Miocene	Ameghino, 1887
Pampatheriinae	<i>Vassallia maxima</i> †	FMNH P14424	Argentina; Catamarca; Andalhuala Formation; Puerta del Corral Quemado	Late Miocene	Castellanos, 1946

(Continues)

TABLE 1 (Continued)

Family/Subfamily	Species	Institutional number	Locality	Period	References
Glyptodontinae	<i>Propalaeohoplophorus minus</i> †	AMNH-FM 9197	Argentina; Santa Cruz; Santa Cruz Formation; Cañadón de las Vacas	Early Miocene	Ameghino, 1891
Glyptodontinae	" <i>Cochlops</i> " <i>debilis</i> †	YPM-VPPU 15592	Argentina; Santa Cruz; Santa Cruz Formation; Cañadón Silva	Early Miocene	Ameghino, 1891
Glyptodontinae	<i>Eucinepeltus complicatus</i> †	AMNH-FM 9248 (type)	Argentina; Santa Cruz; Santa Cruz Formation; Killik Aike Norte	Early Miocene	Brown, 1903
Glyptodontinae	" <i>Metopotoxus</i> " <i>anceps</i> †	YPM-VPPU 15612	Argentina; Santa Cruz; Santa Cruz Formation; Lago Pueyrredón	Early Miocene	Scott, 1903
Glyptodontinae	<i>Glyptodon</i> sp.†	MNHN F-PAM-759	Argentina; Buenos Aires; No more precision	Pleistocene	Owen, 1839 (i)
Glyptodontinae	<i>Glyptodon</i> sp.†	MNHN F-PAM-760	Argentina; Buenos Aires; No more precision	Pleistocene	Owen, 1839 (i)

in phylogenetically distant species (*Dasybus novemcinctus*, *Zaedybus pichiy*, *Cabassous unicinctus*), with the aim of better understanding the ontogenetic variation of the selected anatomical structures. For the determination of relative ages within the developmental series of *Dasybus*, we used the protocol of Le Verger et al., (2020) to analyze the relationship among size, ossification, and dental eruption. Because members of the genus *Cabassous* and *Zaedybus* only have one dental generation (the possession of two dental generations seems to be restricted to *Dasybus*—Ciancio et al., 2012), dental eruption could not be used to determine ontogenetic stage. For these taxa, we used only the relationship between degree of ossification and size. The cingulate specimens also included 14 specimens belonging to fossil species, among them six glyptodonts (Table 1). This sample covered more than 18 million years of cingulate evolutionary history and includes all the major cingulate subfamilies. Our sample does not allow us to evaluate intraspecific variation in all taxa. However, we were able to estimate this variation in three species for which we possess developmental series. For each of them, we analyzed scans of four additional adult specimens. The adult intraspecific sampling is available in Table S1. Species identification was based on collection data, geographical origin, cranial anatomy, and the literature on cingulate taxonomy (e.g., Abba et al., 2015; Carlini et al., 2016; Gaudin & Lyon, 2017; Gaudin & Wible, 2006; Hayssen, 2014; McBee & Baker, 1982; Scott, 1903; Smith & Owen, 2017; Wetzel, 1985; Wetzel et al., 2007). Digital data for all specimens were acquired using X-ray micro-computed tomography ( $\mu$ CT). Specimens were scanned on X-ray tomography imagery platforms at the American Museum of Natural History (New York, USA); the Museum national d'Histoire naturelle (France) in Paris (AST-RX platform), the University of Montpellier (France—MRI platform), and the Museum für Naturkunde (ZMB) in Berlin (Phoenix nanotom [General Electric GmbH Wunstorf, Germany]). Three-dimensional reconstructions of the selected structures were performed using stacks of digital  $\mu$ CT images with MIMICS v. 21.0 software (3D Medical Image Processing Software, Materialize, Leuven, Belgium). The visualization of 3D models was also conducted with AVIZO v. 9.7.0 software (Visualization Sciences Group, Burlington, MA, USA). A large part of specimens scanned will be available on MorphoMuseum for 3D models and on MorphoSource for scans. Specimens scanned by LGR will be available upon request to LGR. The list of specimens is given in Table S2.

## 2.2 | Selected regions of interest and anatomical nomenclature

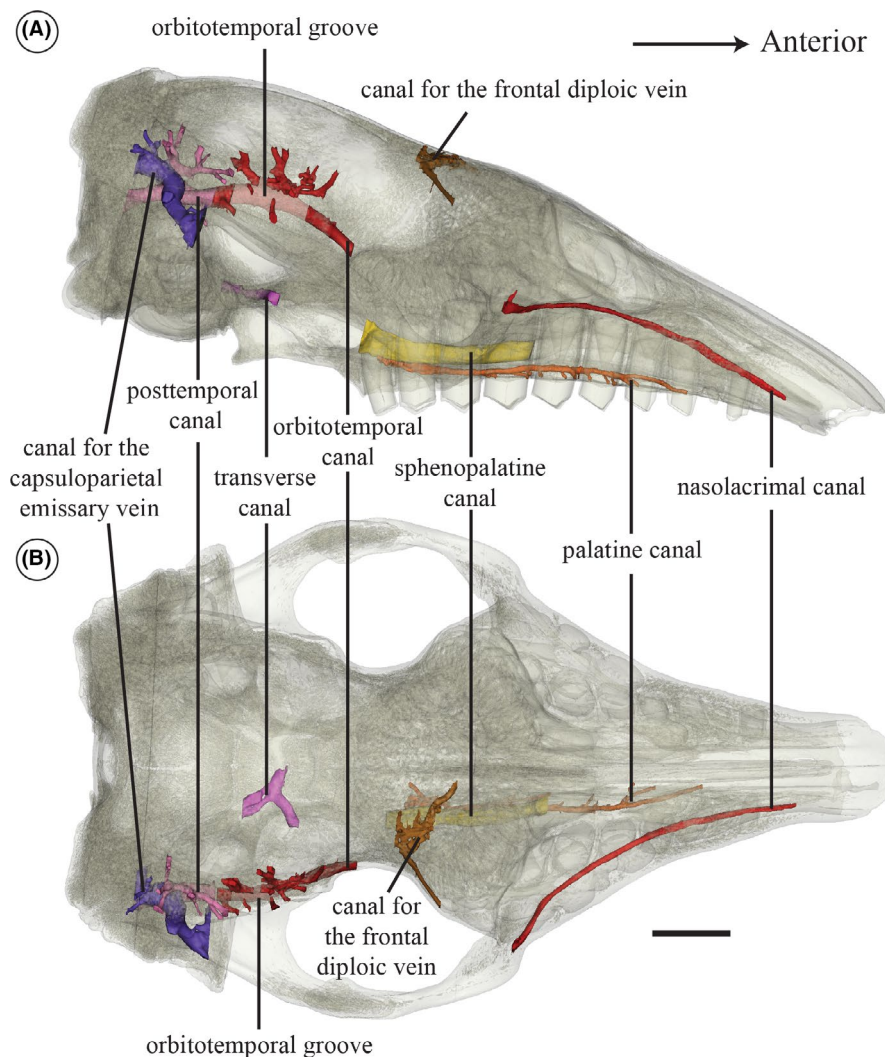
We selected several osseous anatomical complexes in the internal cranial anatomy of cingulates that are poorly known and of interest for this study, such as dental alveoli and several specific intraosseous canals. The latter mostly correspond to vascular pathways and/or the courses of cranial nerves involved in the innervation of various cranial areas: the nasolacrimal canal, the palatine canal, the sphenopalatine canal, the canal for the frontal diploic vein, the transverse

canal, the orbitotemporal canal, the canal for the capsuloparietal emissary vein, and the posttemporal canal.

Our study is focused only on the cranium and does not include the mandible. Therefore, we are only interested in upper dentition. Except for *Dasyus*, which represents a special case (see Ciancio et al., 2012), the homologies in the dental row among the different species of Cingulata are not known mainly because of the drastic reduction in dental complexity (Vizcaíno, 2009). For glyptodonts, previous studies conventionally designated the whole set of teeth as molariform but without further precision (e.g., González Ruiz et al., 2015). We, therefore, chose to treat all teeth as molariform (= Mf; see Herrera et al., 2017 for a different nomenclature and Table S3 for the dental formula of each specimen of our study with greater precision).

Only those intracranial canals whose sampled variation appeared to bear clear systematic information were selected in this work (Figure 1). Nonselected canals generally showed asymmetric and intraspecific variation according to our intraspecific sample (Table S1—e.g., hypoglossal canal; trajectory of the internal carotid artery—see

also Patterson et al., 1989; Gaudin, 1995) or provided no new data (e.g., infraorbital canal) with respect to what was already described or scored for the systematics of the group (Gaudin & Lyon, 2017; Gaudin & Wible, 2006; Wible & Gaudin, 2004). Other nonselected canals were rarely visible in all specimens (often due to taphonomy—not in place and/or obscured by a hard and dense matrix) and were therefore difficult to compare. Specimens that presented a canal that was too incomplete are not described in the relevant section. For the identification and nomenclature of the selected intracranial canals, our study used previous work describing intracranial anatomy in cingulates and eutherians in general (Evans & de Lahunta, 2012; Gaudin, 2004; Gaudin & Lyon, 2017; Muizon et al., 2015; Thewissen, 1989; Wible, 1993; Wible & Gaudin, 2004). We have indicated for each selected region of interest: the variation of these regions during ontogeny, a synthetic comparison among specimens, and the formalization of potential discrete or continuous characters to highlight potential evolutionary scenarios to be mapped onto the tree of cingulates. The formalization of new characters was performed based on observations that were stable among glyptodonts and were shared with



**FIGURE 1** Illustration of the selected internal canals on the transparent cranium of *Euphractus sexcinctus* AMNH 133304 in right lateral (A) and ventral (B) views. Scale = 1 cm

some nonglyptodont cingulates, so that they could provide pertinent information when investigating glyptodont origins.

### 2.3 | Virtual reconstruction of the selected regions

When present, the most dorsal part of the teeth did not always fill the whole alveolar cavity, leaving a void between the roof of the alveolar cavity and the dorsal edge of the tooth. Because we selected the internal orientation and curvature of the whole dental row for study, the reconstruction of alveoli was preferred over the modeling of teeth, which were sometimes absent for our specimens. In some cases, the distinction between the limit of the alveolar wall, the tooth, and the sediment was not identifiable. In this case, we modeled only the distinguishable structures to allow comparison (e.g., *Doellotatus* in Figure 2).

Within the cranium, a canal corresponds to a duct completely enclosed in one or more bones and which usually provides a passageway to vessels and/or nerves. In some cases, a canal may not be continuously enclosed by bone, and thus turns into a groove for part of its course (e.g., the orbitotemporal canal). In such a case, we have reconstructed the course of the groove in a manner similar to that of a canal, but we illustrated the areas corresponding to the groove with increased transparency. In some cases, two canals were confluent in such a way that it was no longer possible to distinguish which part of the duct corresponds to the passage of which vessel or nerve. These regions of confluence were specified in each of our reconstructions by a shaded area (e.g., the posttemporal canal and canal for the capsuloparietal emissary vein). Most selected canals were continuous with external or internal foramina. Figure S1 shows the location of each of these foramina in *Euphractus*.

### 2.4 | Measurements

A few specific measurements were taken for comparison where quantification seemed to offer a better account of the observed variation than a qualitative description. Cranium length, height, and width were measured to calculate the geometric mean of each specimen (= geometric mean, an estimator of the size of a specimen defined by the cubic root of the product of the three variables, Claude, 2008). The geometric mean was log-transformed to facilitate graphing of the data (Claude, 2008). Several other cranial measurements were selected, and their variation was compared to that of the geometric mean in our sample (Table S4 and Figure S2). These measurements were selected in order to quantify the following aspects that appeared interesting to us after our anatomical observations: (a) the relative height of the dental row (i.e., GDRH/GCH = the ratio

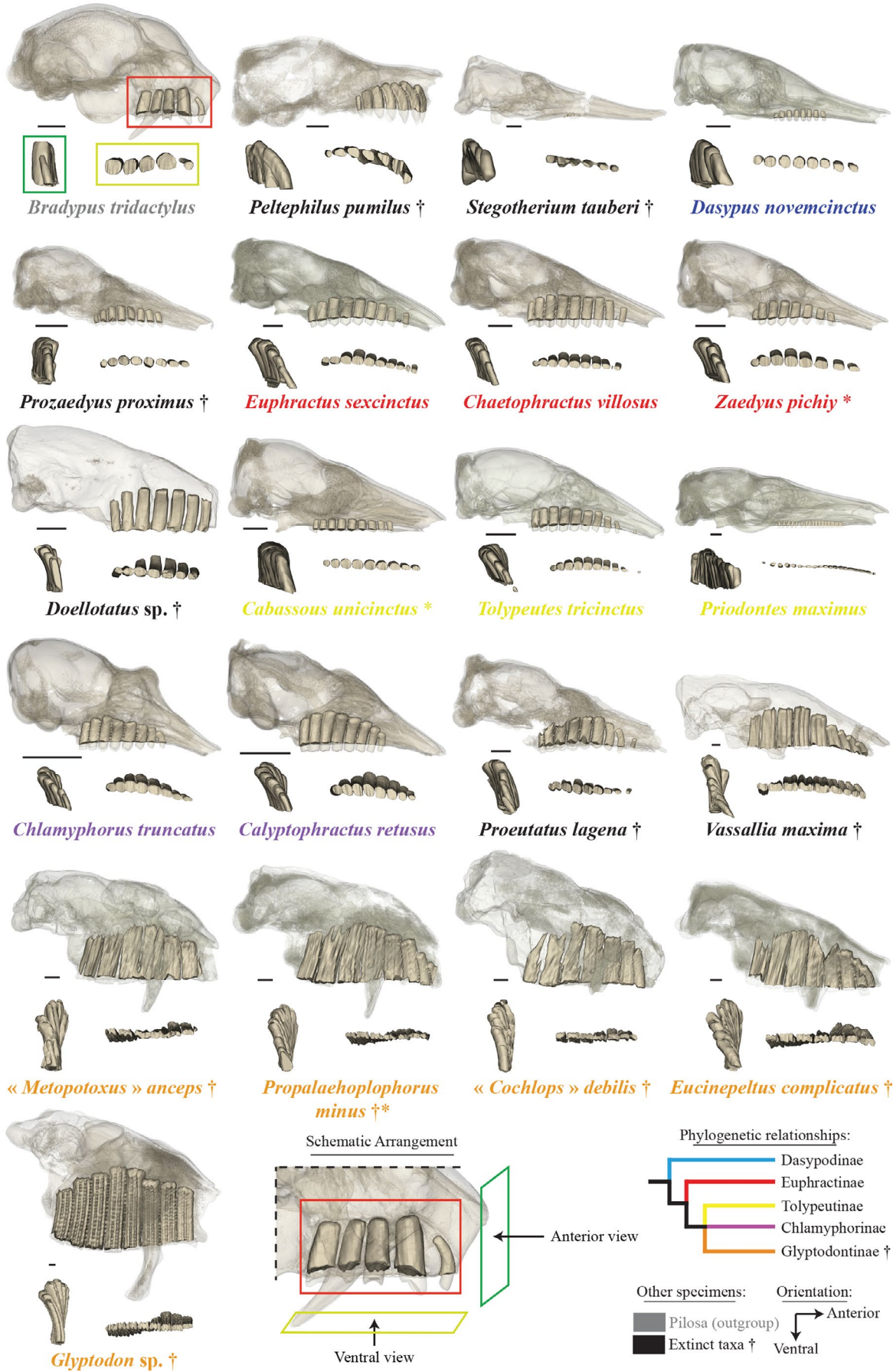
between the greatest dental row height and the greatest cranium height); (b) the angle between the straight line marked by the ventral most point of the tentorial process and the dorsal most point of the annular ridge (see Wible & Spaulding, 2013) and the horizontal anteroposterior axis, defined here by the straight line between the mesial edge of the first tooth and the distal edge of the last tooth in sagittal view. This allowed for a comparison of the internal vault inclination (IVI angle) among taxa. These measurements are further explained and summarized in Figure S2; they were taken using the linear distance and angle tools of MIMICS v. 21.0 software.

### 2.5 | Reconstructing evolutionary scenarios for intracranial traits

In order to visualize possible evolutionary scenarios for the discrete and continuous intracranial traits defined based on our observations, we reconstructed a cladogram that focused on the taxa in our sample by performing a parsimony analysis using PAUP v. 4.0a167 (Swofford, 2002). We used the morphological matrix of Billet et al., (2011) (largely based on that of Gaudin & Wible, 2006), which we limited to the genera in our sample. The initial matrix included all the genera of our sample except for one extant genus (*Calyptophractus*), along with four fossil genera of glyptodonts: "*Cochlops*," *Eucinepeltus*, "*Metopotoxus*," and *Glyptodon*. We have retained the quotation marks for the genera "*Cochlops*" and "*Metopotoxus*" in agreement with the notes of Scott (1903). They have otherwise not been revised since Scott's (1903) report. We scored these taxa (i.e., *Calyptophractus* and several glyptodonts) based on our CT-scanned specimens and added them to the analysis. After this stage, the matrix included 22 taxa scored for 125 characters (Data S1). We performed the same analysis as Billet et al., (2011), with a heuristic search involving 1000 random addition replicates, with 27 morphological characters treated as ordered and treating taxa with multiple states as polymorphic (Data S1). Following the recent molecular works of Delsuc et al., (2016) and Mitchell et al., (2016), we enforced a backbone constraint similar to that of Mitchell et al., (2016). Because this resulted in an unusual position for *Proeutatus* (i.e., as the sister group of euphractines), contrary to its usual placement close to the glyptodonts in morphological analyses (Billet et al., 2011; Engelmann, 1985; Gaudin & Wible, 2006; Herrera et al., 2017), we added a constraint on this taxon to assign it a priori to a position close to glyptodonts (Figure S3A). This choice, therefore, leads us to consider Eutatini as excluded from euphractines.

By using this approach, our aim was only to obtain a more consensual topology, based on recent morphological and molecular analyses, in order to discuss the relevance of our characters. This study is thus not intended to produce a new phylogenetic analysis. The strict consensus of the most parsimonious trees (with two polytomies—extant

**FIGURE 2** Diversity of the dental alveolar cavities in our cingulate sample in lateral (red square), ventral (yellow square) and anterior (green square) view. Crania are reconstructed with transparency to leave the cavities apparent. Colors of species names refer to their attribution at the subfamilial level according to molecular analyses (lower right panel); extinct species not allocated to one of these subfamilies are shown in black. † represents an extinct taxon. Scale = 1 operator # cm





euphractines and glyptodonts except for “*Metopotoxus*”) obtained was then used to calculate branch lengths in order to explore evolutionary scenarios for our qualitative and quantitative observations over the entire topology of the tree. The strict consensus was used as a baseline cladogram (Figure S3B—the strict consensus of the same analysis without constraint is illustrated in Figure S3C for comparison). As the optimization options of the analysis can affect the length of the branches, we have chosen to favor the hypothesis of convergence (= DELTRAN) rather than reversion because they have been regarded as more likely (Wake et al., 2011). For the reconstruction of evolutionary scenarios for intracranial discrete or continuous traits, one needs to complete missing data and resolve polytomies. For the latter, we used the *multi2di* function of the *ape* package of R (Paradis & Schliep, 2019). The function resolves polytomies by adding one or several additional node(s) and corresponding branch(es) of length 0. The duplicate nodes thus have the same values, allowing the removal of duplicates *a posteriori*. In order to facilitate the visualization of the evolutionary scenarios on the tree, we have rendered the tree ultrametric using *force.ultrametric* function of the *phytools* package of R (Revell, 2012). For the discrete traits, the missing data (i.e., 1 taxon for characters 1 and 2 [not applicable], 7 taxa for character 4; 1 taxon for character 5—see Results) were completed according to their ancestral node optimization. In case of optimization uncertainties, we also favored convergences for the same reason mentioned previously. The reconstruction and mapping of ancestral states were performed using stochastic mapping with a symmetric condition matrix (i.e., one for which all transformation rates are considered equivalent; see Bollback, 2006; Revell, 2012). This analysis was performed using the *make.simmap* function of the *phytools* package of R, with which we produced 100 simulations for each character. For continuous traits, missing data (i.e., 2 taxa for the character 8) were estimated using an approach combining the use of multiple imputations with procrustean superimposition of principal component analysis results performed on all measurements (Table S4; Clavel et al., 2014) with the *estim* function of the *mvMORPH* package of R (Clavel et al., 2015). Then, the reconstruction of the ancestral states was performed using maximum likelihood with the *contMap* function of the *phytools* package of R (Revell, 2012).

## 2.6 | Institutional Abbreviations

**AMNH**, American Museum of Natural History, New York, USA; **FMNH**, Field Museum of Natural History, Chicago, USA; **MHNG**, Muséum d'Histoire Naturelle de Genève, Genève, Switzerland; **MNHN**, Muséum National d'Histoire Naturelle, Zoologie et Anatomie comparée collections (**ZM**), Mammifères et Oiseaux collections (**MO**), fossil mammal collections, Pampean (**F.PAM**), Paris, France; **NBC**, Naturalis Biodiversity Center, Leiden, Holland; **NHMUK**, Natural History Museum, London; **NMNH**, National Museum of Natural History, Smithsonian Institution, Washington, DC, USA; **UF**, University of Florida, Gainesville, USA; **YPM-VPPU**, Princeton University collection housed at Peabody Museum, Yale University, USA; **ZMB**, Museum für Naturkunde, Berlin, Germany.

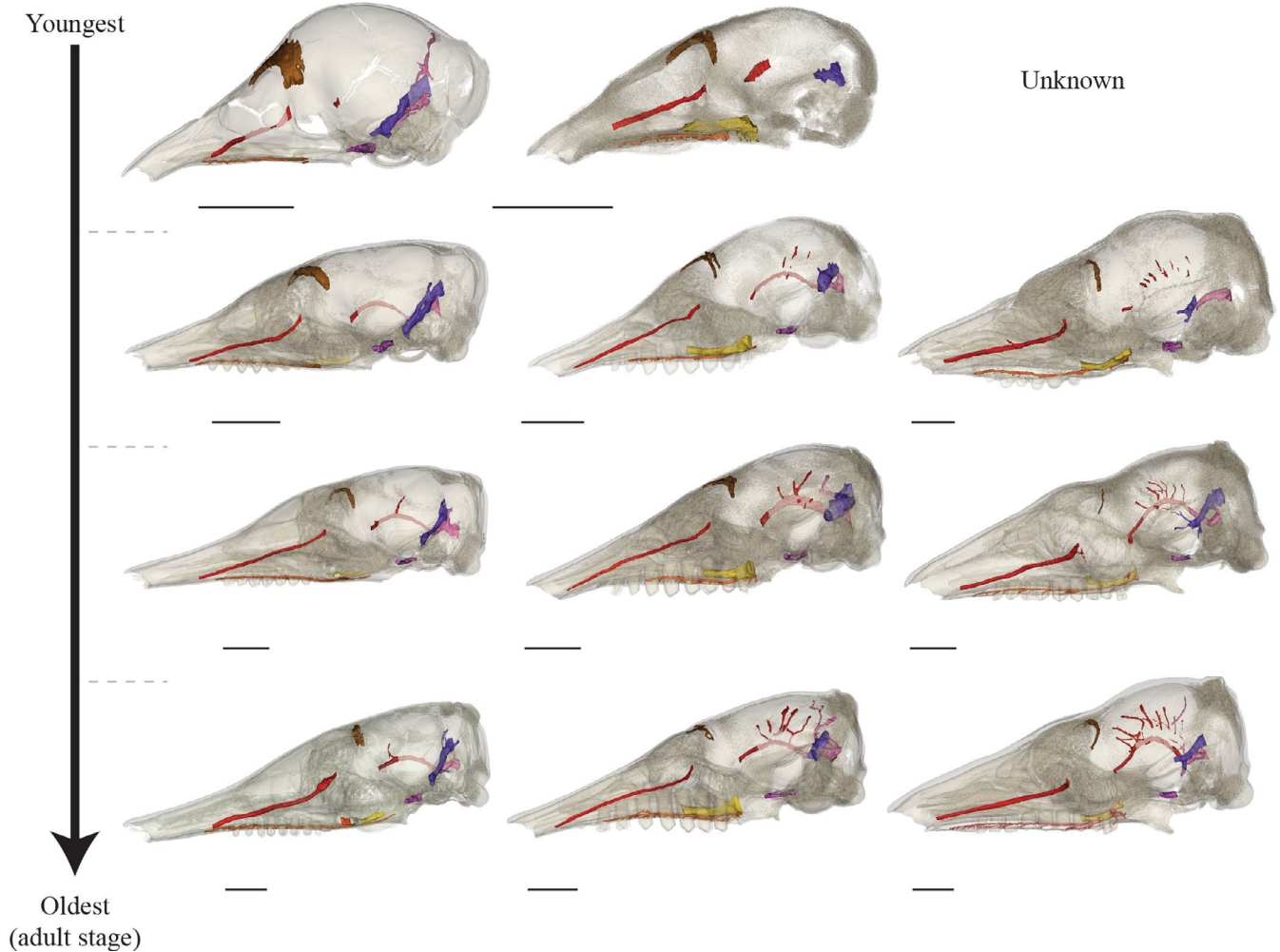
## 3 | RESULTS: ANATOMICAL DESCRIPTION AND COMPARISON

### 3.1 | Teeth and alveolar cavities: Orientation, curvature, and height

The three species sampled intraspecifically (*Dasyopus novemcinctus*, *Zaedyus pichiy*, and *Cabassous unicinctus*—Table 1) do not show any ontogenetic variation in the orientation and curvature of the teeth compared with the pattern observed in adults (see below). In *Dasyopus* and *Zaedyus*, the relative height of teeth shows an increase from the youngest to the oldest specimens, suggesting an increase in height during ontogeny (Figure 3). In *Cabassous*, which possess a reduced dentition as *Dasyopus*, no clear ontogenetic trend has been observed, but we lack a stage as young as those present in *Dasyopus* and *Zaedyus* (Figure 3).

**Height:** The relative height of the alveolar cavities distinguishes specimens with reduced teeth, such as *Stegotherium*, *Dasyopus*, *Cabassous*, and *Priodontes*, from other cingulate species, some of which may even show extremely enlarged teeth, as is the case in glyptodonts (Figure 2, Table S4).

In taxa with reduced teeth (unfunctional supernumerary teeth are not considered; see González Ruiz et al., 2014, 2015, 2017; González Ruiz & MacPhee, 2014), the height of the alveolar cavity in lateral view is practically the same for all teeth (Figure 2). In the other taxa, the dorsal profile of the dental row in lateral view is curved (dorsally convex): that is, teeth gradually increase in height backward until the middle of the dental series and then gradually decrease in height posteriorly. The most dorsal point is reached in the middle of the dental row (at the level of Mf5 in dental rows containing 7–10 Mf) in *Euphractus* (at Mf5/Mf9), *Chaetophractus* (at Mf5/Mf9), *Doellotatus* (at Mf5/Mf9–10) and *Tolypeutes* (at 5Mf/Mf9), and more anteriorly in *Peltephilus* (at Mf2/Mf7), and *Proeutatus* (at 4Mf/Mf9). In *Bradypus*, this dorsal point is attained more anteriorly (at Mf2/Mf5). It is reached more posteriorly in *Zaedyus* (at Mf5/Mf8), *Prozaedyus* (at Mf5/Mf8 and Mf6/Mf9), chlamyphorines (at Mf6/Mf8), and most glyptodonts (at Mf5–6/Mf8). In addition, the GDRH/GCH ratio, which expresses dental height relative to cranium height, seems to be correlated with body size ( $R^2 = 0.5363$ ;  $P$ -value =  $9.42E-06$ ), showing that large-sized taxa have relatively taller dentitions. This is reminiscent of the increase in height of the dental row during ontogeny in *Dasyopus* and *Zaedyus*. Most large-bodied taxa in the sample, that is, *Vassallia* and glyptodonts, but not *Priodontes*, do indeed show a strongly elevated dentition in relation to cranium depth (Figure 4). This is also the case in *Peltephilus* for which the height of the cranium is particularly low in relation to its length. Although they are small (i.e., GCL <43 mm), chlamyphorines show a tooth height comparable in proportion to the larger-sized *Euphractus*, *Chaetophractus*, and *Proeutatus* (Figure 4), meaning that they depart slightly from the detected allometric trend in having teeth taller than expected for their size. Relative tooth height is lower in *Prozaedyus*, *Zaedyus*, *Tolypeutes*, and *Cabassous* and extremely low in specimens with dental reduction—*Stegotherium*, *Dasyopus*, and *Priodontes* (Figure 4).

*Dasypus novemcinctus**Zaedyus pichiy**Cabassous unicinctus*

**FIGURE 3** Ontogenetic variation of internal canals selected in this study viewed in lateral view on transparent crania in each developmental series (*Dasypus*, *Zaedyus* and *Cabassous*). Note that early postnatal stages for *Cabassous* are not documented in our sample. See Le Verger et al., (2020) for the determination of stages. Phylogenetic relationships are indicated with colors following Figure S3. The color of canals follows that of Figure 1. Scale = 1 cm

**Curvature and Orientation:** As visible in anterior and ventral views (Figure 2), *Stegotherium*, *Priodontes*, and *Cabassous* exhibit a rather homogenous curvature and orientation of teeth along the dental row while most cingulates show gradual changes in these aspects anteroposteriorly. In most cingulates, anterior teeth are tilted lingually, that is, tilted with a medially offset apex, while most posterior teeth often tilt labially (character 5 in Gaudin, 2004). The tilt of posterior teeth is often much less pronounced than for anterior teeth. Similarly, taxa that show curved crowns generally show anterior teeth with an inward curvature (lingual convexity) which can be strongest in the middle of the dental row, while the two posterior-most teeth often show a lesser degree of inward curvature or even an outward curvature (labial convexity). A strong inward curvature of anterior teeth is observed in *Peltephilus*, *Doellotatus*, *Tolypeutes*, *Vassallia*, chlamyphorines, and glyptodonts in our sample, whereas other taxa show straighter crowns, and an outward curvature is

observed in *Dasypus*. In lateral view, the first tooth (Mf1; P1 for *Dasypus*) or teeth show a mesial curvature (mesial convexity) and a mesially offset apex in *Bradypus*, *Peltephilus*, *Proeutatus*, *Vassallia*, *Propalaeohoplophorus*, and *Eucinepeltus*. Mf1 is also tilted with a mesially offset apex in *Prozaedyus*. Finally, the two to three posteriormost teeth are distinctly tilted with a distally offset apex in *Bradypus* and glyptodonts. Most other loci and taxa exhibit a nearly vertical orientation of teeth in lateral view.

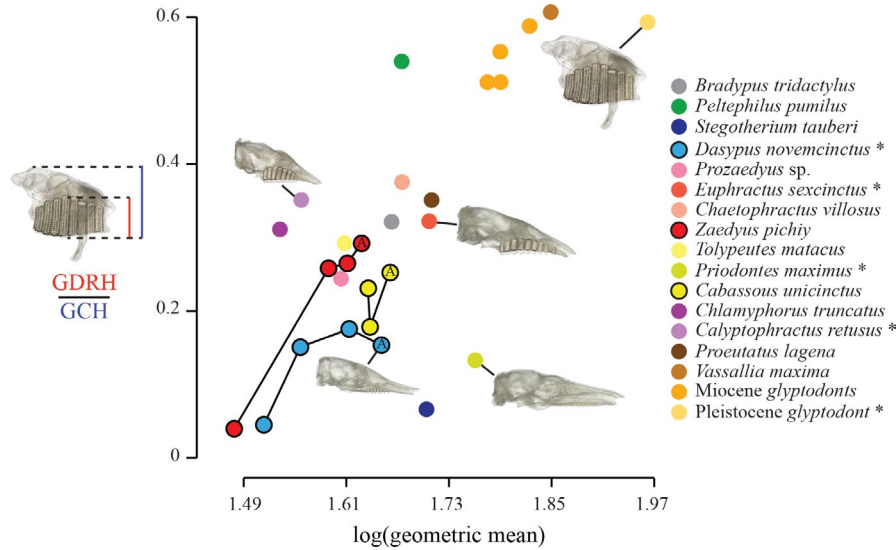
Based on these observations, we propose scrutinizing evolutionary scenarios for the following two characters of the dentition (see Table 2).

**Character 1** (discrete—unordered): Position of the most dorsal point of the dorsal convexity of the tooth row.

**State (0):** Toothrow not dorsally convex.

**State (1):** Anterior to middle of the tooth row.

**State (2):** At the middle of the tooth row.



**FIGURE 4** Distribution of the ratio of greatest dental row height (GDRH) to greatest cranium height (GCH) with respect to cranial size (= log(geometric mean); see Materials & Methods). For each of the three developmental series, A, symbolizes the adult specimen. Crania of specimens marked with an asterisk are illustrated in the graph

**TABLE 2** Data matrix with character scores for each genus. Characters 1 and 7 are discrete, and character 8 is continuous (see text). Symbols: -, not applicable; ?, missing data; †, extinct species.

Taxa	Character 1	Character 2	Character 3	Character 4	Character 5	Character 6	Character 7	Character 8
<i>Bradypus</i>	1	1	0	0	2	0	0	3.66
<i>Tamandua</i>	-	-	0	0	0	0	0	11.06
<i>Peltephilus</i> †	1	0	0	1	2	1	1	-4.85
<i>Dasybus</i>	0	2	1	1	0	1	1	17.73
<i>Stegotherium</i> †	0	1	1	1	2	1	1	20.83
<i>Prozaedyus</i> †	3	1	?	?	0	1	1	?
<i>Doellotatus</i> †	2	0	?	?	?	1	1	8.08
<i>Euphractus</i>	2	1	1	1	0	1	1	6.58
<i>Chaetophractus</i>	2	1	1	1	0	1	1	3.96
<i>Zaedyus</i>	3	1	1	1	0	1	1	1.67
<i>Chlamyphorus</i>	3	0	1	2	2	1	1	18.29
<i>Calyptophractus</i>	3	0	1	2	2	2	2	11.68
<i>Cabassous</i>	0	1	1	1	0	1	2	8.57
<i>Tolypeutes</i>	2	0	1	1	2	1	2	-1.04
<i>Priodontes</i>	0	1	1	1	0	0	2	-2.11
<i>Proeutatus</i> †	1	1	0	?	1	2	1	17.92
<i>Vassallia</i> †	3	0	0	?	1	1	2	0.08
<i>Propalaehoplophorus</i> †	3	0	0	?	1	2	2	18.56
" <i>Metopotoxus</i> " †	3	0	0	?	1	2	2	19.45
" <i>Cochlops</i> " †	3	0	0	2	1	2	2	21.2
<i>Eucinepeltus</i> †	3	0	0	?	1	2	2	?
<i>Glyptodon</i> †	3	0	0	2	2	2	2	35.84

**State (3):** Posterior to middle of the tooth row.

**Character 2** (discrete—unordered): Curvature of anterior teeth in anterior view.

**State (0):** Inward curvature.

**State (1):** Straight.

**State (2):** Outward curvature.

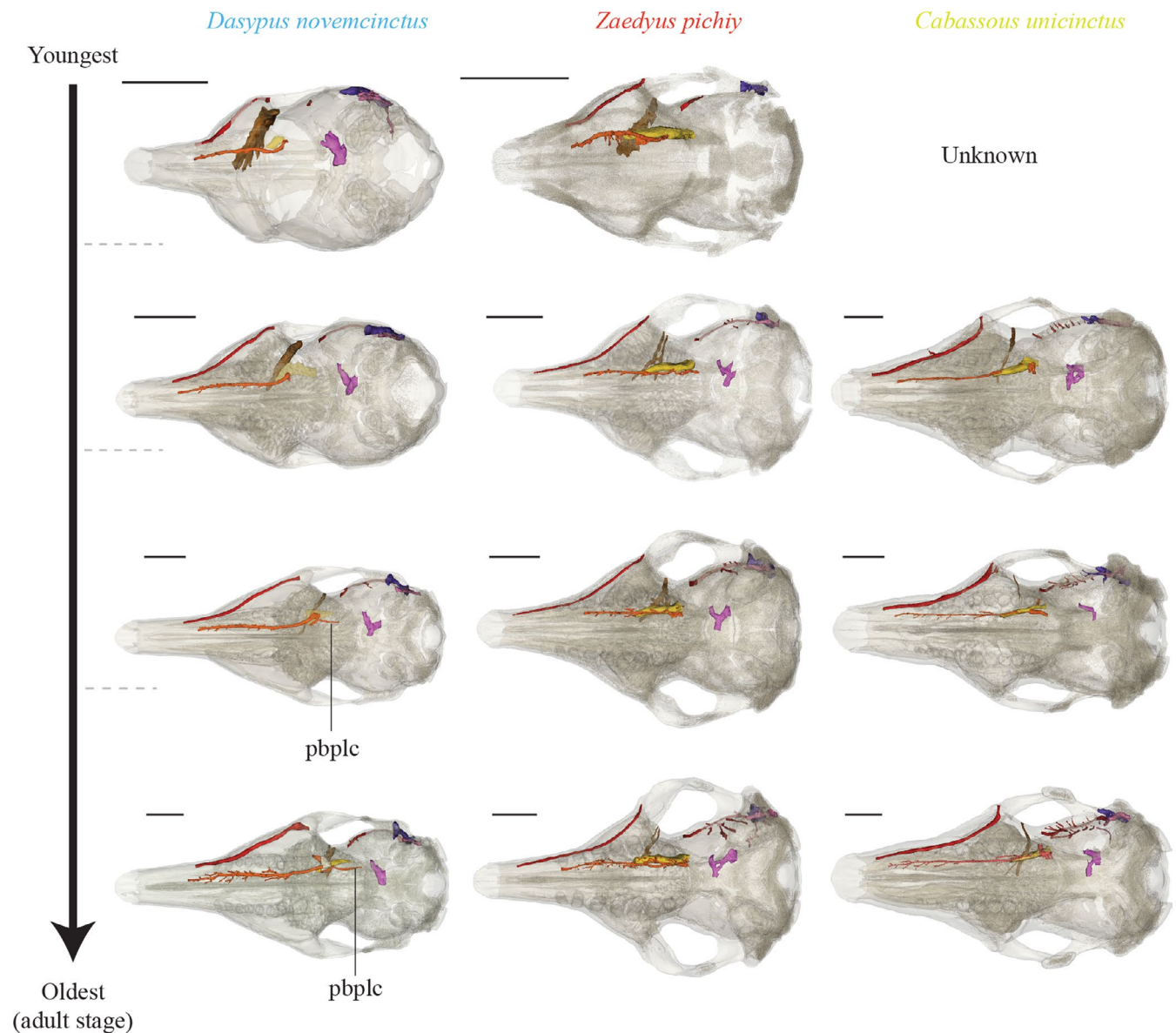
## 3.2 | Cranial Canals

### 3.2.1 | Frontal Diploic Vein canal

The course of this canal is entirely within the frontal bone. It opens externally at one or exceptionally two foramina located in the orbitotemporal fossa, slightly ventral to the most dorsal part of the orbital margin (Figure S1). Internally, the canal opens through one or more foramina located posterior to the annular ridge and near the midline without ever crossing it (Figure S4). This canal conveys the frontal diploic vein, an emissary of the dorsal cerebral vein/dorsal sagittal

sinus, or a vein issuing from the frontal diploë (Evans & de Lahunta, 2012; Muizon et al., 2015; Thewissen, 1989; Wible & Gaudin, 2004).

In early ontogeny, the canal of the frontal diploic vein is initially extremely thick in the youngest specimens of *Dasypus* (see also Billet et al., 2017) and *Zaedyus* and occupies a large part of the frontal bone (Figures 3 and 5). Its relative diameter is considerably reduced in older specimens of *Dasypus* and *Zaedyus*, with no change in its curved trajectory in *Zaedyus*, but becoming more strictly transverse in the adult stage in *Dasypus* (Figures 3 and 5). In *Cabassous*, it remains fairly similar from juvenile to adult stages in our sample (Figures 3 and 5).



**FIGURE 5** Ontogenetic variation of internal canals selected in this study viewed in ventral view on transparent crania in each developmental series (*Dasypus*, *Zaedyus* and *Cabassous*). Note that early postnatal stages for *Cabassous* are not documented in our sample. See Le Verger et al., (2020) for the determination of stages. Phylogenetic relationships are indicated with colors following Figure S3. The color of canals follows that of Figure 1. Abbreviations: pbplc, posterior branch of the palatine canal. Scale = 1 cm

In our adult sample, the canal is relatively straight, very thin, oriented posteromedially in dorsal view, and without ramifications in *Bradypus* (Figure 6). In *Tamandua*, *Peltephilus*, *Stegotherium*, *Dasypus*, *Euphractus*, *Priodontes*, and *Cabassous*, its overall course is transverse but with a posterior bend or convexity of varying degree (Figure 6). Ramifications and a posterior convexity are also found in *Prozaedyus*, *Chaetophractus*, *Zaedyus*, and *Proeutatus*, but the overall course of their canal is more anteromedially oriented, as is also observed in *Stegotherium*, *Priodontes*, and *Cabassous* (Figure 6). The same orientation is found in *Doellotatus* (incomplete), except that the course of the canal is completely straight and has no ramifications (Figure 6). *Tolypeutes*, on the other hand, show a very peculiar course with a forked structure in a dorsal view (Figure 6). The canal is merged with diploe in chlamyphorines, but its presence is observable in *Calyptophractus*, in which it has a straight and

anteromedially oriented trajectory (Figure 6). This canal was not observed in *Vassallia* or in any of the glyptodonts sampled. However, Gaudin (2004) notes the presence of this canal in glyptodonts and Gaudin and Lyon (2017) mention it in the pampathere *Holmesina* (Simpson, 1930) (but see below). We suspect a taphonomic bias for some specimens (e.g., some glyptodonts) and call for a deeper study of this canal in cingulates before its variation can be scored.

### 3.2.2 | Nasolacrimal Canal

The nasolacrimal canal originates posteriorly at the anterior orbital edge with the lacrimal foramen (Figure 1 and Figure S1). It runs anteriorly within the lacrimal and the maxillary bones in its posteriormost portion. More anteriorly, the canal is located between the inner wall

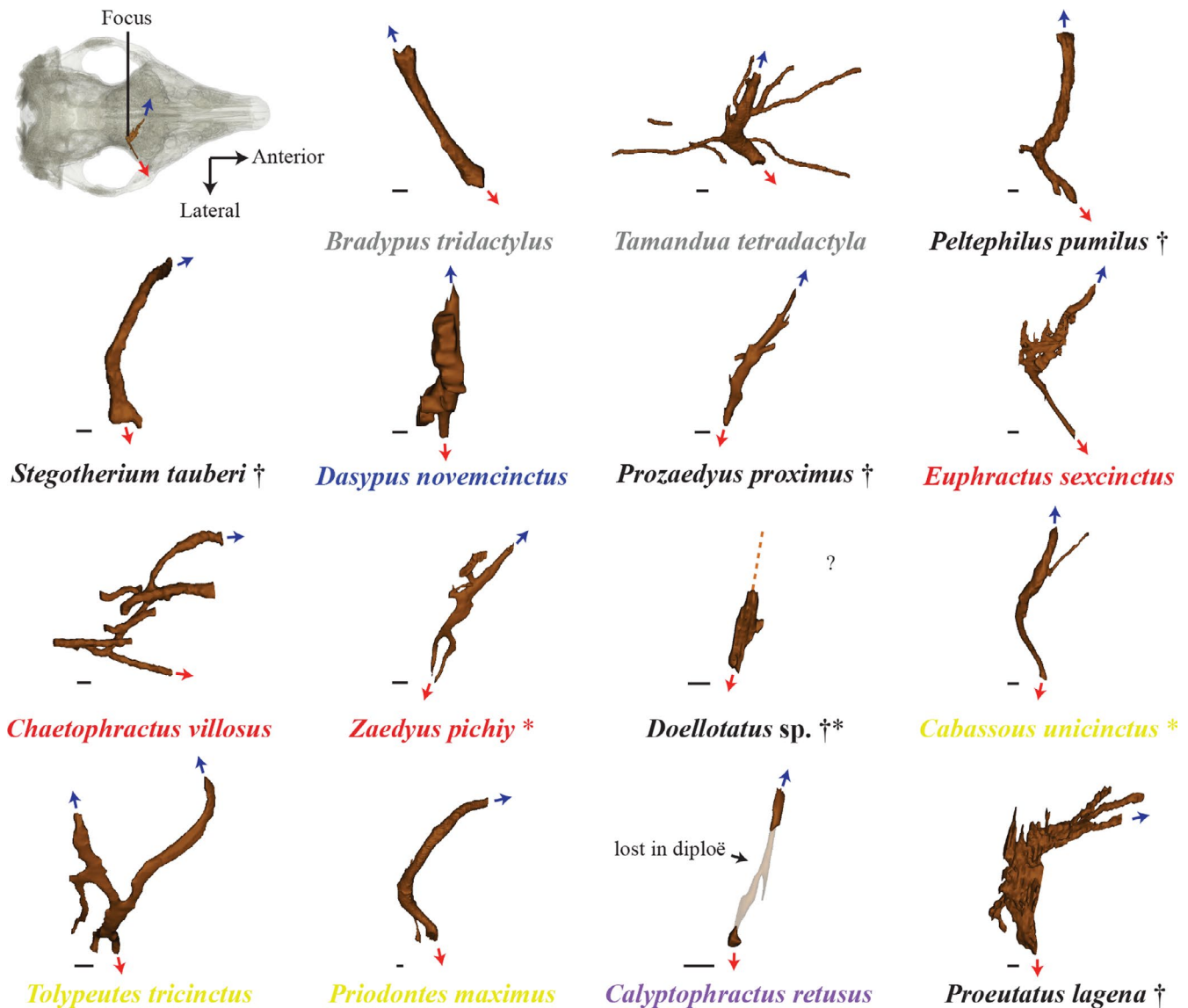


FIGURE 6 Interspecific variation of the canal for the frontal diploic vein in our sample, all illustrated in dorsal view. Red arrows mark the main external opening of the canal. Blue arrows mark the main internal path of the canal on the cranial midline. Colors of species names follow Figure 2. Scale = 1 mm

of the maxillary and turbinates. It opens anteriorly (in front of the tooth row, except in *Peltephilus*) and ventromedially into the nasal cavity (Figures S1E and S5). This canal allows the transmission of fluids from the lacrimal sac to the nasal cavity and is potentially accompanied by a vein in *Euphractus* (see Wible & Gaudin, 2004).

The course of the nasolacrimal canal barely changes during ontogeny in our sample (Figures 3 and 5). Its length changes and follows the lengthening of the snout that accompanies the growth of the cranium (Le Verger et al., 2020). In addition, *Cabassous* and, to a lesser degree, the adult stage of *Zaedyus* show a more medial orientation of the course of the nasolacrimal canal in its most posterior part, which differs slightly from *Dasybus* and other stages of *Zaedyus*, where it runs more parallel to the external surface of the cranium (Figures 3 and 5).

The position of the nasolacrimal canal and lacrimal foramen varies among taxa, although this feature was not scored in previous matrices (e.g., Gaudin & Wible, 2006). The lacrimal foramen is particularly high relative to the orbital edge (closer to the most dorsal point of the orbit than to the jugal bone in lateral view) in chlamyphorines, *Proeutatus*, and glyptodonts (Figure 7). The trajectory of the nasolacrimal canal is strongly sigmoid in *Tamandua* (Figure 7). In *Bradypus*, *Peltephilus*, *Proeutatus*, *Euphractus*, and *Chaetophractus*, the course of the nasolacrimal canal is slightly convex dorsally (Figure 7). The curvature is much pronounced in *Vassallia* in the posterior half of its course as it passes above the tooth row (Figure 7). In dasypodines, *Prozaedyus*, *Zaedyus*, tolpeutines, and chlamyphorines, the nasolacrimal canal is relatively straight or it bears a slight ventral convexity in its posterior part (Figure 7). In glyptodonts, the canal runs ventromedially and not anteriorly from the lacrimal foramen (Figures 7 and 8), and curves strongly inward anteriorly when approaching the sagittal plane of the cranium (Figures 7 and 8).

Whereas glyptodonts appear to show a unique condition in the course of their nasolacrimal canal, the anterior opening of this canal is located halfway up in their nasal cavity as in *Bradypus*, *Tamandua*, *Peltephilus*, *Doellotatus*, *Proeutatus*, and *Vassallia*. In comparison, this opening is closer to the palate in other specimens (Figure 7). This variation can be scored in the following character (see Table 2).

**Character 3** (discrete—unordered): Anterior opening of the nasolacrimal canal in nasal cavity.

**State (0):** Halfway up in the nasal cavity.

**State (1):** Ventral in the nasal cavity, close to the palate.

### 3.2.3 | Palatine and sphenopalatine canals

Two large canals are enclosed within the maxillary and palatine bones forming the hard palate in cingulates: the palatine canal and the sphenopalatine canal (Figure 1 and S1) (Wible & Gaudin, 2004). The palatine canal is generally thinner, longer than, and positioned medial to the sphenopalatine canal. The palatine canal transmits the major and minor palatine nerve, artery, and vein in *Euphractus* (Wible & Gaudin, 2004). This canal opens externally at the caudal palatine foramen posteriorly and at the multiple foramina scattered

throughout the hard palate (both in the palatine and maxillary) anteriorly (Figure S1). The sphenopalatine canal, which is often very wide in cingulates, transmits the caudal nasal nerve and the sphenopalatine artery and vein (Wible & Gaudin, 2004). The sphenopalatine canal opens anteriorly in the nasal cavity and posteriorly in the pterygopalatine fossa just posterior to the dental row and close to or confluent with the caudal palatine foramen (Wible & Gaudin, 2004; Figure S1). The palatine canal and sphenopalatine canal are confluent in part of their posterior course in many of the species in our sample.

The palatine canal and sphenopalatine canal vary little during ontogeny within the three species of our ontogenetic series (Figures 3 and 5). The branch of the palatine canal posterior to its region of confluence with the sphenopalatine canal in *Dasybus* and *Cabassous* appears in the subadult (or intermediate) stage and elongates further in the adult stage (Figures 3 and 5). In the youngest specimen of *Zaedyus*, the sphenopalatine canal is much wider relative to cranium size than in later stages, and there is no region of confluence with the palatine canal (Figures 3 and 5). In the youngest *Dasybus* specimen, the sphenopalatine canal is absent. It is present only as a wide groove in the youngest to subadult stages but is completely enclosed in the adult stage of *Dasybus* in our sample (Figures 3 and 5).

In our adult sample, the shape and trajectory of the palatine canal do not vary much among species (Figures 7 and 8). In glyptodonts, the palatine canal strongly ascends posteriorly (starting from Mf6) (Figure 7) and forms a groove running along the inner wall of the nasopharyngeal canal (Figure S6). In other cingulates, the palatine canal does not show such an ascending trajectory posteriorly. The sphenopalatine canal, which is thicker than the palatine canal, features a relatively straight trajectory in lateral view in *Peltephilus*, dasypodines, *Prozaedyus*, *Doellotatus*, *Euphractus*, *Zaedyus*, and "*Cochlops*," whereas it shows a strong ventral curvature in *Chaetophractus*, tolpeutines, chlamyphorines, and *Glyptodon* (Figure 7). It should be further noted that the dorsal position of the sphenopalatine canal relative to the nasopharyngeal canal is a feature shared by *Vassallia* and *Glyptodon* (Figure S7). In *Bradypus* and *Tamandua*, the palatine canal does not contact the sphenopalatine canal (Figures 7 and 8). The two canals are in contact in a small region of confluence in the dasypodines whereas the palatine canal becomes completely confluent with the sphenopalatine canal, and sometimes accompanied by ventrally oriented auxiliary branches of the palatine canal, in *Peltephilus*, tolpeutines, chlamyphorines, and *Glyptodon* (Figures 7 and 8). In extant euphractines, the confluence between the sphenopalatine canal and the palatine canal is not as clear-cut as in the above-cited cingulates (Figures 7 and 8). The palatine canal of euphractines runs ventrally along the sphenopalatine canal in the same figure eight-shaped canal while being completely conjoined only in their most posterior part and at their posterior opening (Figures 7 and 8). The extent of this confluence is highly variable in *Euphractus* according to Wible and Gaudin (2004) and in other cingulates as well (Gaudin & Wible, 2006—character 71). We observed the same variation as Gaudin and Wible (2006) concerning the extent of this confluence in the vicinity of the posterior opening

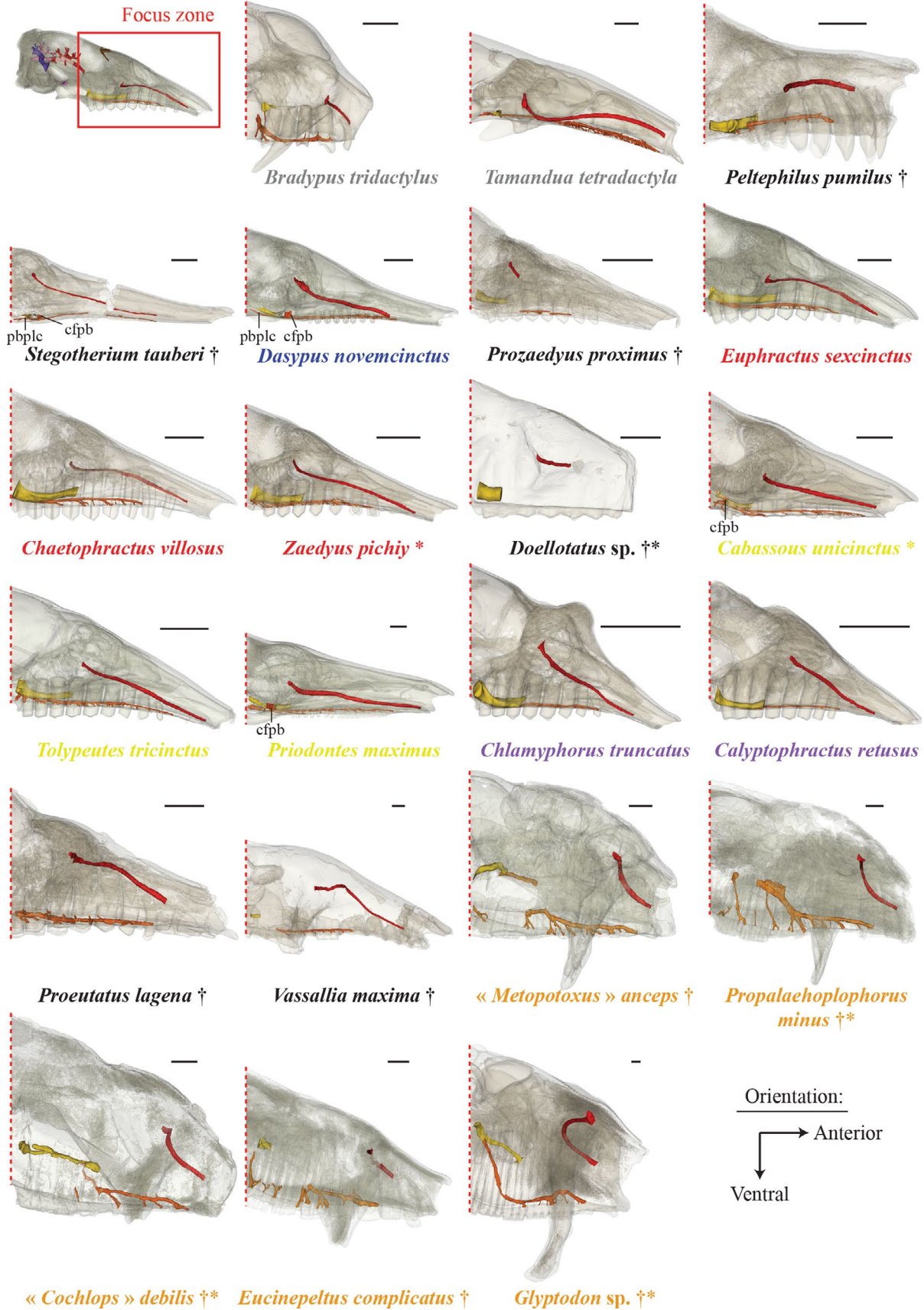


FIGURE 7 Interspecific variation in our sample of the reconstructed nasolacrimal (= red), sphenopalatine (= yellow) and palatine (= orange) canals shown in lateral view. Crania are reconstructed with transparency. Colors of species names follow Figure 2. Abbreviations: cfpb, caudal foramen for palatine canal branch; pbplc, posterior branch of palatine canal. Scale = 1 cm

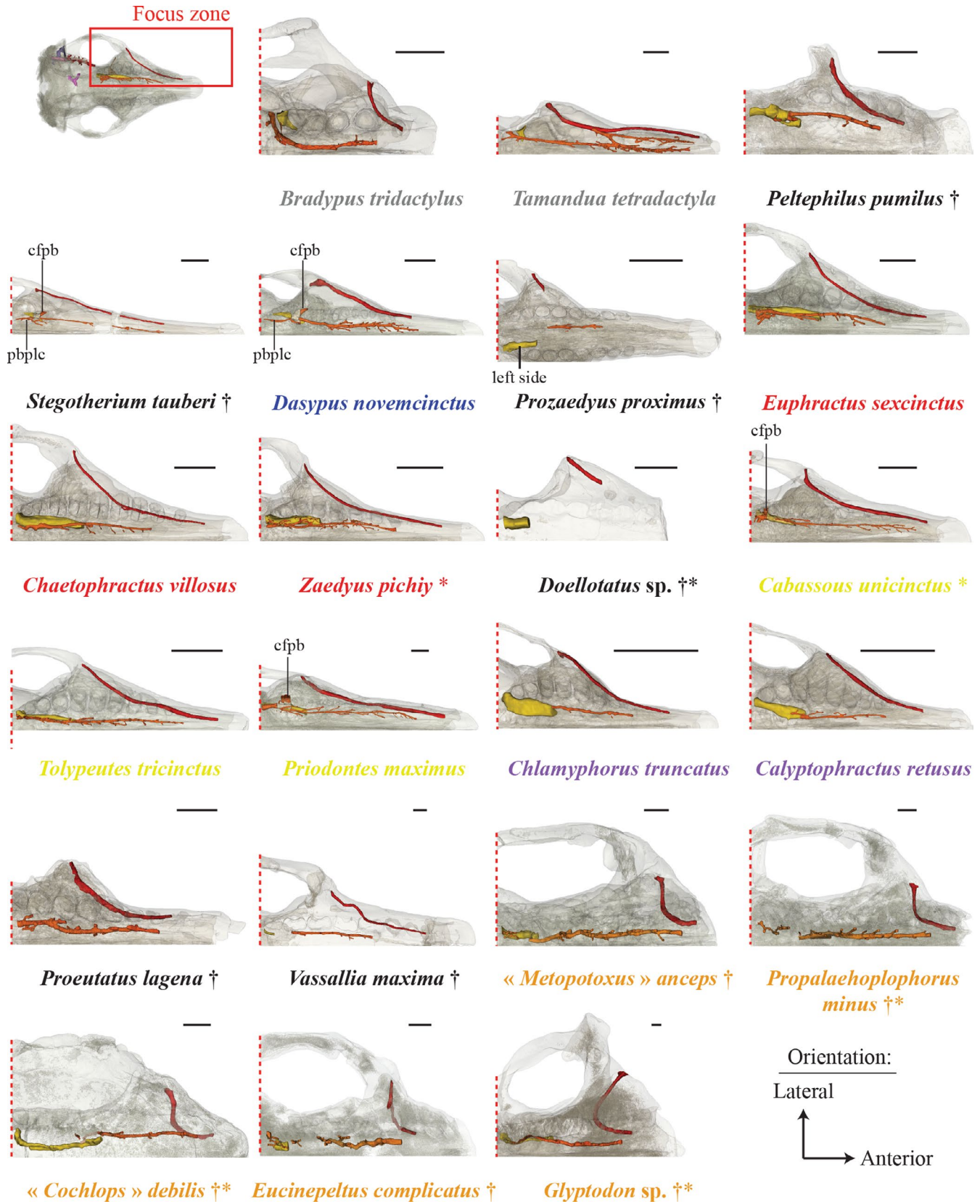


FIGURE 8 Interspecific variation in our sample of the reconstructed nasolacrimal (= red), sphenopalatine (= yellow) and palatine (= orange) canals shown in ventral view. Crania are reconstructed with transparency. Colors of species names follow Figure 2. Abbreviations: cfpb, caudal foramen for palatine canal branch; pbplc, posterior branch of palatine canal. Scale = 1 cm



of the sphenopalatine canal for all the taxa in our sample. Several branches may emerge from this region of confluence (Figures 7 and 8), some of which present an interesting pattern. One branch opens externally from the caudal foramen of the palatine canal in dasypodines, *Priodontes*, and *Cabassous* (Figures 7 and 8). A second branch extends posteriorly into the hard palate before disappearing in the bone diploe in dasypodines. In *Euphractus*, *Zaedyus*, *Priodontes*, and *Cabassous*, two or more branches extend posteriorly into the hard palate, but they open in the nasopharyngeal canal (Figures 7 and 8). Several other ramifications of the palatine canal are present in our sample, but it remains to be determined how variable these are at the intraspecific level with a data set larger than that of Table S1.

Based on the observation of these two canals, we proposed to scrutinize evolutionary scenarios for the following character (see Table 2).

**Character 4** (discrete—unordered): Sphenopalatine and palatine canal connection.

**State (0):** No contact.

**State (1):** Partial contact, with palatine canal running ventrally along the sphenopalatine canal and producing posterior branches.

**State (2):** Complete fusion.

### 3.2.4 | Transverse Canal

When present, the transverse canal is marked externally by a foramen positioned anteroventrally to or within the foramen ovale in the alisphenoid (Figure S1). This canal crosses the cranium transversally at the level of the basisphenoid and transmits a large vein issued from the cavernous sinus (Sánchez-Villagra & Wible, 2002; Wible & Gaudin, 2004). Often, one or more branches originate from this canal and extend posteriorly toward the lateral edge of the basisphenoid to join the inferior petrosal sinus (Wible & Gaudin, 2004).

The transverse canal appears early in *Dasybus* and varies little during ontogeny. Although, in *Zaedyus* and *Cabassous*, the connection between the transverse canal and the posterior branch extending toward the inferior petrosal sinus varies in both series, we have not observed a clear ontogenetic pattern of variation in our three species (Figures 3 and 5).

In *Bradypus*, *Peltephilus*, *Stegotherium*, *Doellotatus*, *Tolypeutes*, and *Glyptodon*, there is no transverse canal, or it is not detectable (scored as present in *Doellotatus* and *Stegotherium*, variable in *Tolypeutes* by Gaudin & Wible, 2006—character 111). Chlamyphorines have a transverse canal foramen opening directly within the braincase, and thus there is a very short canal that does not cross the midline of the cranium. *Tamandua* stands out in having a thin lateral part of the transverse canal which differs markedly from the condition in Cingulata (Figure 9). In *Tamandua*, *Dasybus*, *Prozaedyus*, extant euphractines, *Priodontes*, and *Cabassous*, the branch starting from the transverse canal foramen is oriented posteromedially in ventral view (Figure 9). In the Miocene glyptodonts, it is medially oriented (Figure 9). In *Vassallia*, it is anteromedially oriented (Figure 9). This lateral branch of the transverse canal splits into a branch that crosses

the midline in the basisphenoid bone and a branch that joins the inferior petrosal sinus (Figure 9). In *Proeutatus*, this bifurcation occurs much closer to the transverse canal foramen than in other taxa (Figure 9). It is relatively more medial in *Prozaedyus*, extant euphractines, *Priodontes*, and *Cabassous* (Figure 9). In *Tamandua*, *Dasybus*, *Vassallia*, and Miocene glyptodonts, this bifurcation is closer to the sagittal plane than to the foramen (Figure 9). The transverse medial branch is anteromedially oriented in ventral view in *Prozaedyus*, *Euphractus*, and *Proeutatus*, whereas it is medially oriented in the other taxa (Figure 9). Our specimens of *Tamandua*, extant euphractines (not complete in *Chaetophractus*), and *Priodontes* show an additional branch connecting the transverse medial branch to the posterior branch reaching the inferior petrosal sinus (Figure 9). In *Vassallia* and the Miocene glyptodonts, the posterior branch reaching the inferior petrosal sinus is much thinner than the rest of the canal, in contrast to the other taxa, in which the whole canal exhibits a relatively homogeneous width (Figure 9).

Based on these observations of the transverse canal, we propose to scrutinize evolutionary scenarios for the following character (see Table 2).

**Character 5** (discrete—unordered): Orientation of the branch starting from the transverse canal foramen.

**State (0):** Posteromedial.

**State (1):** Medial or anteromedial.

**State (2):** No canal.

### 3.2.5 | Orbitotemporal and posttemporal canals

Within the braincase, the orbitotemporal canal provides a passageway to the rostral extension of the *ramus superior* of the stapedia artery, or orbitotemporal artery (giving rise to the *ramus supraorbitalis* in the orbital region), and a few small veins (Wible & Gaudin, 2004), whereas the more posterior posttemporal canal transmits the *arteria diploëtica magna* and the large *vena diploëtica magna* (Wible, 1987; Wible & Gaudin, 2004). These two canals can give rise to the arterial and venous *rami temporales* along the lateral wall of the braincase, which exit via numerous foramina on the cranial roof to irrigate the *temporalis* muscle (Wible & Gaudin, 2004; Figure S1). The posttemporal canal is connected posteriorly to an external groove in the petro-occipital region marking the passage of the occipital artery (Wible & Gaudin, 2004; Figure S1). From there, the posttemporal canal extends forward as a canal enclosed between the petrosal and squamosal (or only in the squamosal) and is oriented mostly horizontally. More anteriorly, past the petrosal, the posttemporal canal is connected to the orbitotemporal canal, which runs further anteriorly within or on the inner surface of the lateral braincase wall (formed by the squamosal, frontal, and sometimes the parietal). The delimitation between the two canals generally occurs where the canal for the *ramus superior* joins them (Muizon et al., 2015). However, we were only able to observe a canal possibly transmitting the *ramus superior* in *Tolypeutes* (Figure S8) but not in the other specimens. Consequently, we followed the suggestion of Wible and Gaudin

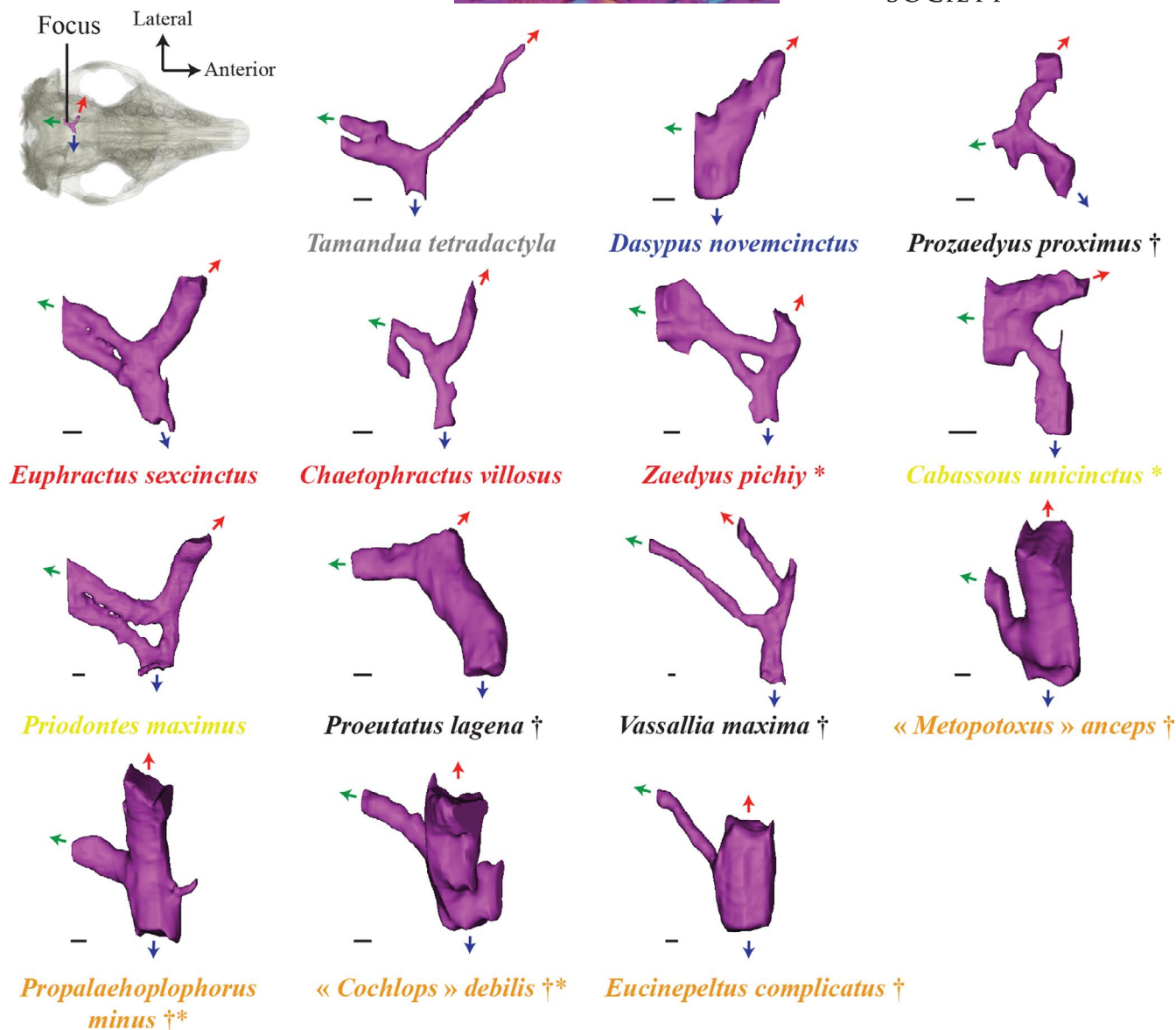


FIGURE 9 Interspecific variation of the transverse canal in ventral view in our sample. Red arrows mark the main external opening of the canal. Blue arrows mark the branch crossing the midline. Green arrows mark the main posterior branch which connects to the inferior petrosal sinus (see text). Colors of species names follow Figure 2. Scale = 1 mm

(2004), who separate the two canals at the level of the postglenoid foramen. In some taxa, the orbitotemporal canal is not completely enclosed for parts of its length and appears instead as a groove on the internal wall of the braincase. The anterior opening of the orbitotemporal canal is located in the orbitotemporal region, just posteroventral to the postorbital constriction (Figure S1; Wible & Gaudin, 2004; = cranioorbital foramen in Gaudin & Wible, 2006; see also Muizon et al., 2015 for multiple illustrations of these canals in another placental taxon).

The orbitotemporal canal only appears as a short canal near the orbitotemporal region in the youngest specimens of all three species, whereas the canal (or groove) is clearly visible for its entire length in later stages (Figures 3 and 5). The connections of the orbitotemporal canal with canals for the numerous *rami temporales* occur

in the two last stages in both *Zaedyus* and *Cabassous* (Figures 3 and 5). The posttemporal canal is already well formed in the youngest specimen of *Dasybus* and hardly changes during ontogeny (Figures 3 and 5). In *Zaedyus*, the posttemporal canal is not enclosed in the youngest stage. In *Cabassous*, the posttemporal canal is present in the youngest stage sampled, but it expands anteriorly during the two subsequent stages (Figures 3 and 5).

In our adult sample, *Bradypus*, *Tamandua*, extant euphractines, *Vassallia*, and glyptodonts possess a posttemporal canal emerging at the center of the occiput in lateral view and at the level of the jugular foramen (Figure 10). In *Peltephilus*, it originates more ventrally, even with the most dorsal margin of the occipital condyles in lateral view (Figure 10). In dasypodines, tolpeutines, and chlamyphorines, the posterior extremity of the canal is at an intermediate height

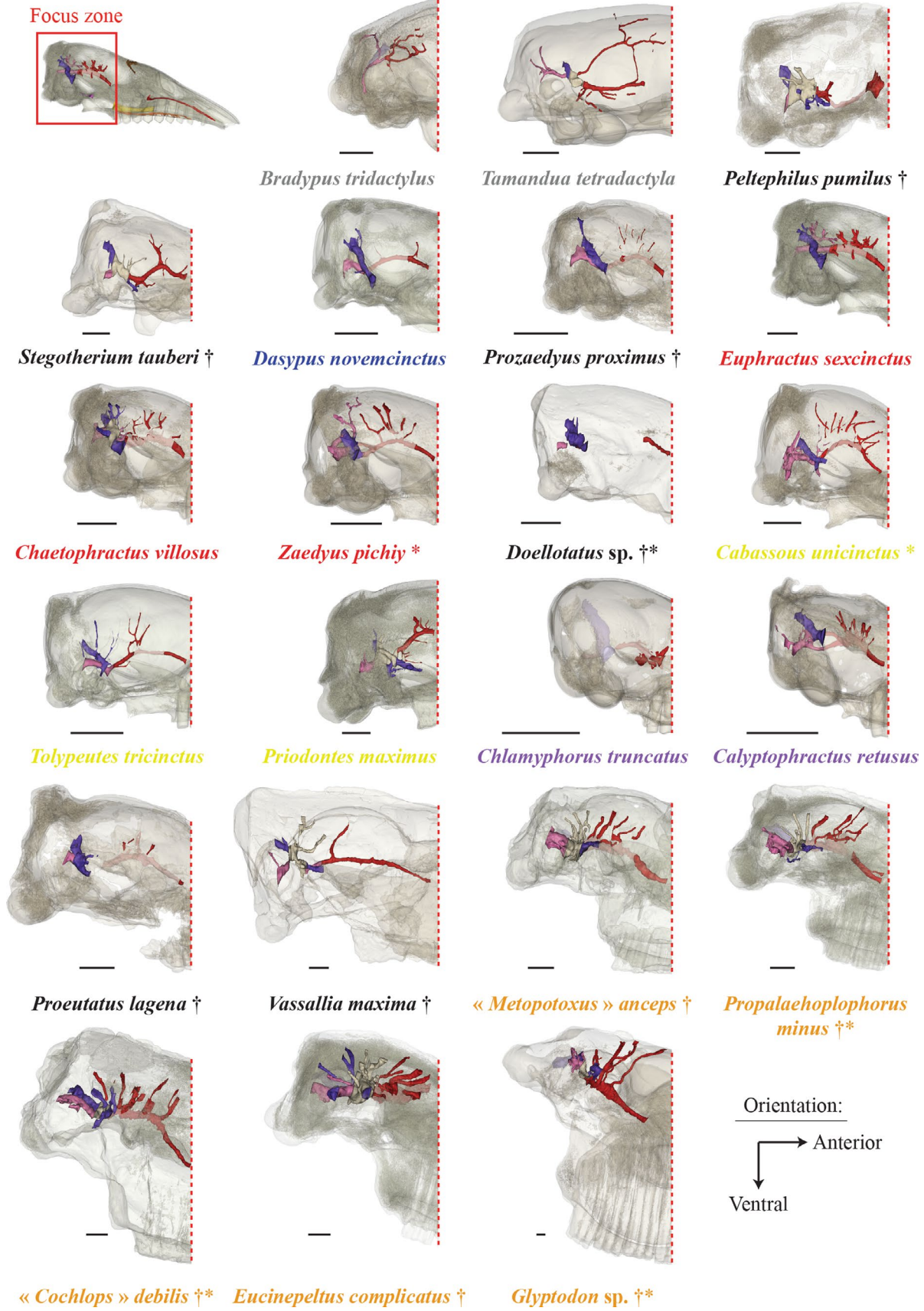


FIGURE 10 Interspecific variation in our sample of the orbitotemporal (= red), posttemporal (= pink) and capsuloparietal emissary vein (= blue) canals shown in lateral view. Caudal part of the cranium is transparent. Parts of the canals showing transparency symbolize a groove instead of a fully enclosed canal. Colors of species names follow Figure 2. Scale = 1 cm

(Figure 10). Apart from the position of the posterior opening of this canal (= posttemporal foramen of Wible & Gaudin, 2004), we were unable to determine any systematically significant variation regarding its direction, length, or width.

In *Bradypus* and *Tamandua*, the orbitotemporal canal does not reach the orbitotemporal region (Figures 10 and 11). It reaches the orbitotemporal region in all the cingulates in our sample except for *Priodontes* (Figures 10 and 11). Gaudin and Wible (2006) mention the absence of the anterior opening of the orbitotemporal canal not only in *Priodontes* (character 78) but also in *Chlamyphorus* and *Vassallia* for which we observe a clear opening in the orbitotemporal region (Figure 10). In *Bradypus*, *Tamandua*, and *Peltephilus*, this canal bears a ventral curvature (strong in *Peltephilus*), whereas its trajectory exhibits a more or less strong dorsal curvature in all the other taxa (Figure 10). In *Peltephilus*, *Dasypus*, *Prozaedyus*, extant euphractines, *Tolypeutes*, *Cabassous*, chlamyphorines, *Proeutatus*, and Miocene glyptodonts, the orbitotemporal canal becomes a groove on the internal lateral wall of the braincase before being enclosed again as a canal close to its external anterior opening (Figure 10). In *Calyptopractus*, *Proeutatus*, and glyptodonts, the anterior half of the orbitotemporal canal displays a pronounced downward trajectory, and its anterior opening is located lower than its posterior connection with the posttemporal canal (Figure 10). *Calyptopractus* stands out, however, in having two branches that connect the orbitotemporal and posttemporal canals (Figure 10). A large region of confluence between the orbitotemporal and posttemporal canals and the canal for the capsuloparietal emissary vein is identified in *Tamandua*, *Peltephilus*, *Stegotherium*, *Dasypus*, *Chaetopractus*, *Priodontes*, *Vassallia*, and the glyptodonts (and potentially also in *Proeutatus* but the orbitotemporal canal is not distinct in this region; Figures 10 and 11). This confluent region is reminiscent of the petrosquamosal fossa in *Alcidedorbignya*, Muizon & Marshall, 1987 (Muizon et al., 2015). The orbitotemporal and posttemporal canals connect with the *rami temporales* through multiple small canals whose number remains quite variable at the intraspecific level. However, a high density of *rami temporales* for the orbitotemporal canal is particularly noticeable in extant euphractines, *Cabassous*, *Calyptopractus*, and glyptodonts (Figure 10). This density is coded by Gaudin and Wible (2006), who counted the number of foramina for *rami temporales* in the temporal fossa of parietal (0, equal to or less than five; 1, greater than five—character 97).

In their matrix, almost all cingulates are coded as having more than five foramina. The variations in the density of the canals for the *rami temporales* observed in our study call for further investigation of these structures, which could potentially carry an interesting phylogenetic signal among cingulates.

Based on these observations of the orbitotemporal canal, we propose to scrutinize evolutionary scenarios for the following character (see Table 2).

**Character 6** (discrete—unordered): Downward trajectory of the orbitotemporal canal.

**State (0):** Canal does not reach the orbitotemporal region.

**State (1):** The anterior part of the canal has a slight downward trajectory ending at the same or almost the same height as its origin from the posttemporal canal.

**State (2):** The canal has a strong downward anterior trajectory, and its anterior opening is much lower than its posterior connection to the posttemporal canal.

### 3.2.6 | Canal of the Capsuloparietal Emissary Vein

The canal for the capsuloparietal emissary vein opens anteroventrally at the postglenoid and suprimeatal foramina and connects to the groove for the transverse sinus posterodorsally (Muizon et al., 2015; Wible, 1993). From the cranial roof, the transverse sinus runs lateroventrally in a wide groove excavated in the inner surface of the parietal and extends anteroventrally within a canal formed by the petrosal medially and the squamosal laterally. This canal opens externally through the suprimeatal foramen, the postglenoid foramen and accessory foramina at the posterior base of the zygomatic arch (e.g., Wible & Gaudin, 2004; Figure S1). It transmits the capsuloparietal emissary vein (= retroarticular vein in *Canis*, Linnaeus 1758 (Evans & de Lahunta, 2012); postglenoid vein in *Cynocephalus*, Boddaert, 1768 (Wible, 1993), and *Dasypus*, Linnaeus 1758 (Wible, 2010)). In our sample, this canal is often partly confluent with the posttemporal and orbitotemporal canal in its posterodorsal portion (Figure S9).

The canal for the capsuloparietal emissary vein is already well formed in the youngest specimen of *Dasypus* and hardly changes during ontogeny, except that it becomes relatively thinner (Figures 3 and 5). In *Zaedyus*, it is only partly formed in the youngest specimen, and its trajectory is better marked in older stages (Figures 3 and 5). In the youngest *Cabassous*, this canal is noticeably short and only marked in its most ventral part (glenoid region). It is only in older stages that the canal elongates posterodorsally (Figures 3 and 5).

The canal is absent and the passage of the vein is only indicated by a groove in *Bradypus* and *Chlamyphorus*. The groove is very wide in the latter two (Figure 10). The canal of the capsuloparietal emissary vein opens through the postglenoid foramen and potentially in the suprimeatal foramen as well (see *rami temporales* opening in the squamosal whose number varies in our intraspecific sampling, Table S1) in *Peltephilus*, *Prozaedyus*, *Doellotatus*, extant euphractines, chlamyphorines, *Proeutatus*, *Vassallia*, and the glyptodonts (as notably scored for the glenoid region by Gaudin & Wible, 2006—character 119; Figures 10 and 11). In tolpeutines, the canal always opens by a suprimeatal foramen with a well-marked branch, whereas its opening via the postglenoid foramen may sometimes be absent, as we have occasionally observed the absence of a postglenoid foramen in our intraspecific sample of *Cabassous* (Table S1). For all our taxa except *Bradypus*, the canal extends anteroventrally from the cranial roof (Figure 10). However, its inclination varies considerably from one species to another. It is steeply inclined in dasypodines, slightly less so in *Prozaedyus*, *Doellotatus*, extant euphractines, and *Proeutatus*, and much less so in tolpeutines, *Calyptopractus*,

Focus zone

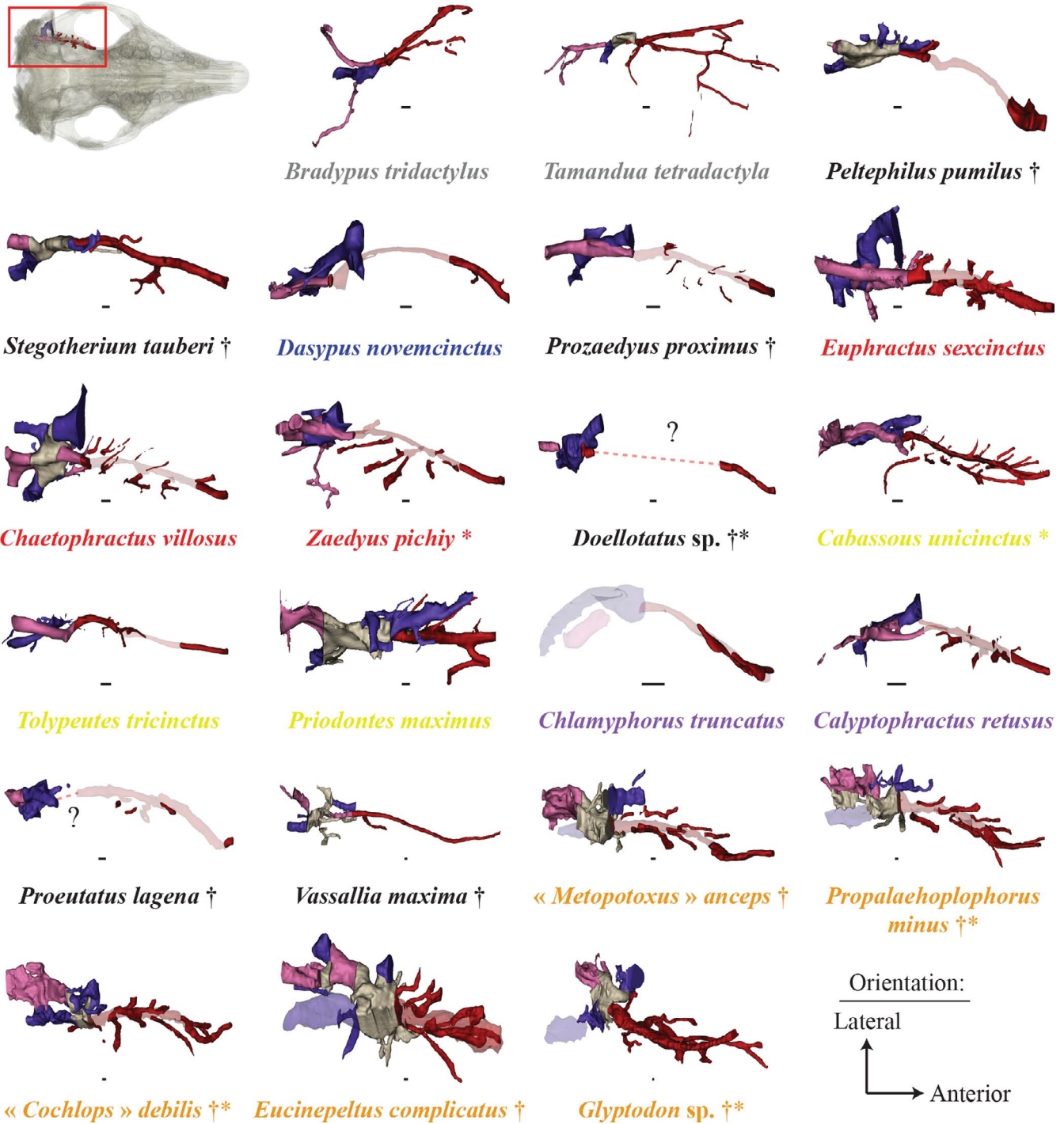


FIGURE 11 Interspecific variation in our sample of the orbitotemporal (= red), posttemporal (= pink) and capsuloparietal emissary vein (= blue) canals shown in ventral view. Transparent parts of the canals indicate a groove instead of a fully enclosed canal. Colors of species names follow Figure 2. Scale = 1 mm

*Vassallia* and glyptodonts (Figure 10). The canal is sinuous in *Euphractus* and *Chaetophractus*. In glyptodonts, the canal even has an almost anteroposterior orientation near its anterior opening (Figures 10 and 11). The canal (or groove) for the capsuloparietal emissary vein reaches a region of confluence (= petrosquamosal fossa, Muizon et al., 2015) with the other canals of the braincase

in *Tamandua*, *Peltephilus*, dasypodines (particularly thin in *Dasypus*), *Chaetophractus*, *Priodontes*, *Vassallia*, and glyptodonts (Figures 10 and 11). For the other taxa, the canal for the capsuloparietal emissary vein is only adjacent to the other canals (Figures 10 and 11—see Figure S9 for an illustration of the different cases in our sample). The variation in length and thickness of this canal does not provide

clear systematic information. A relatively short canal is present in *Tamandua*, *Doellotatus*, *Zaedyus*, and *Proeutatus* compared with dasyrodines (Figure 10). Chlamyphorines and *Zaedyus* show a relatively thicker canal compared with the very thin canal in *Vassallia*. The fact that small taxa have a relatively thicker canal is reminiscent of young specimens of *Dasybus* compared to older and larger specimens of the same species (Figures 3 and 5) and suggest a potential allometric pattern.

Based on these observations, we propose to scrutinize evolutionary scenarios for the following character related to the anterolateral extremity of the canal (see Table 2).

**Character 7** (discrete—unordered): Canal of the capsuloparietal emissary vein inclination in lateral view.

**State (0):** Anterodorsal orientation or canal very short.

**State (1):** Inclination anteroventral from the posterior opening to the anterior opening.

**State (2):** Less anteroventrally inclined, with an almost anteroposterior orientation close to the anterior opening.

### 3.3 | Inclination of the cranial roof

Digital study of the braincase of our sample enabled observation of notable variations in the internal vault inclination (= IVI) relative to the anteroposterior axis of the cranium. The values of the IVI angle (see Materials and Methods) show an interesting pattern in our sample that does not correlate ( $R^2 = 0.1768$ ;  $P$ -value = 0.3246) with size (Figure 12). Whereas *Glyptodon* is distinguished from the other cingulates by an exceedingly high inclination combined with a large size, the large-sized *Vassallia* has a very low cranial roof inclination (Figure 12). *Dasybus* (adult specimen), *Stegotherium*, *Chlamyphorus*, *Proeutatus*, and Miocene glyptodonts

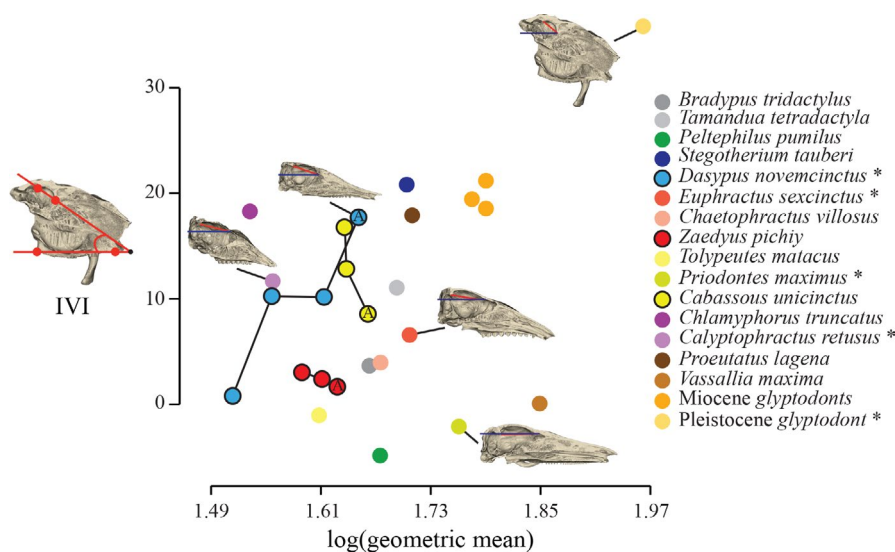
are also characterized by high inclinations (Figure 12). The juvenile *Cabassous* also displays a strong inclination, which decreases with age, whereas the inclination increases with age in *Dasybus* (Figure 12). *Calyptophractus* shows an intermediate cranial roof inclination (Figure 12). The outgroups, extant euphractines, and *Cabassous* (adult) exhibit low values, and the remaining tolypeutines and *Peltephilus* are even characterized by a negative tilting of IVI (Figure 12).

We propose to treat the IVI angle as a continuous character to scrutinize evolutionary scenarios (see Table 2).

**Character 8** (continuous): Internal Vault Inclination.

## 4 | DISCUSSION

Due to their very unusual anatomy, even within the Cingulata, the phylogenetic placement of glyptodonts has been long debated (Burmeister, 1874; Carlini & Zurita, 2010; Fariña et al., 2013; Scott, 1903). Recent molecular analyses have proposed a new hypothesis (i.e., as sister group of the clade tolypeutines + chlamyphorines—Delsuc et al., 2016; Mitchell et al., 2016) at odds with the multiple hypotheses based on morphological analyses (Billet et al., 2011; Engelmann, 1985; Gaudin & Wible, 2006; Herrera et al., 2017). The recent investigation of the internal anatomy in the *Dasybus novemcinctus* species complex (Billet et al., 2017) underlined the potential phylogenetic signal hidden in the internal cranial structures of cingulates. Comparative studies on the paranasal sinuses and on brain endocasts of cingulates have been performed but could not provide consistent information on the placement of glyptodonts within the Cingulata (Fericola et al., 2012; Tambusso & Fariña, 2015a; Zurita et al., 2011). Our comparative study complements these previous efforts by adding the analysis of the dental alveolar cavities (previously



**FIGURE 12** Distribution of the internal vault inclination (IVI) with respect to cranial size (= log (geometric mean)) in our sample. For the three developmental series, A symbolizes the adult specimen. Specimens marked with an asterisk are illustrated by cranial reconstructions in sagittal view in the graph. Scale = 1 cm

only explored in the glyptodont genus *Eucinepeltus* by González Ruiz et al., 2020) and of various canals involved in the vascularization and innervation of the cranium (see Wible & Gaudin, 2004). Based on an extensive sample of extant and extinct cingulates, our survey enabled us to describe many new aspects of the internal cranial anatomy in the group and to propose eight potential characters with helpful phylogenetic information to aid in determining the placement of glyptodonts within Cingulata.

The anatomical variation of the dental alveolar cavities in cingulates comprises many aspects, including the height, curvature, and orientation of teeth, along with their number and distribution on the rostrum (Vizcaíno, 2009). Among our observations, we have proposed two characters with potential bearing on the affinities of glyptodonts. The first corresponds to the position of the most dorsal point of these cavities in lateral view (character 1). The dorsal margins of the alveoli form a dorsally convex line for the whole dental row in most cingulates, except in taxa with reduced teeth. This most dorsal point is situated posteriorly (i.e., at Mf5-6) in *Prozaedyus*, *Zaedyus*, chlamyphorines, pampatheres, and glyptodonts in our sample. The distribution of this character would be congruent with the hypothesis of a close relationship between pampatheres and glyptodonts, as suggested by many previous authors (Billet et al., 2011; Gaudin & Wible, 2006; Herrera et al., 2017; Patterson & Pascual, 1972), but also supports the clade formed by the latter two with chlamyphorines, as proposed by the morphological analysis of Mitchell et al., (2016) constrained by the molecular backbone (Figure 13). On the other hand, *Proeutatus*, generally considered to be a close relative of the clade pampatheres + glyptodonts (Billet et al., 2011; Engelmann, 1985; Gaudin & Lyon, 2017; Gaudin & Wible, 2006), has its most dorsal point situated much further anteriorly (i.e., ca. Mf1-3).

The second character derived from dental alveolar cavities corresponds to the curvature of the anterior teeth in anterior view (character 2). As for character 1, chlamyphorines and pampatheres resemble glyptodonts in this aspect, exhibiting an inward curvature. *Peltephilus*, *Doellotatus*, and also *Tolypeutes* show a similar condition (Figure 13). As shown by the reconstructed evolutionary scenario (Figure 13), this character could still support the morphological hypothesis of Mitchell et al., (2016) of a close relationship between chlamyphorines and glyptodonts. The distribution of this character on the baseline cladogram shows some homoplasy, however, with the condition in *Proeutatus* contrasting with that of glyptodonts, while *Peltephilus* resembles the latter (Figure 13). Another point of interest concerning the variation of dental alveolar cavities is the potential allometric pattern identified by the correlation between the relative height of the dental row and cranium size (Figure 4). This needs to be analyzed further and compared with other allometric patterns seen in the cranium. Mammals generally show an allometric elongation of the rostrum as body size increases, known as craniofacial allometry (Cardini, 2019; Cardini & Polly, 2013). In *Dasybus*, this allometric pattern mostly affects the proportions of the rostrum horizontally but less so vertically (Le Verger et al., 2020). The possible allometric increase in the height of the dentition in cingulates

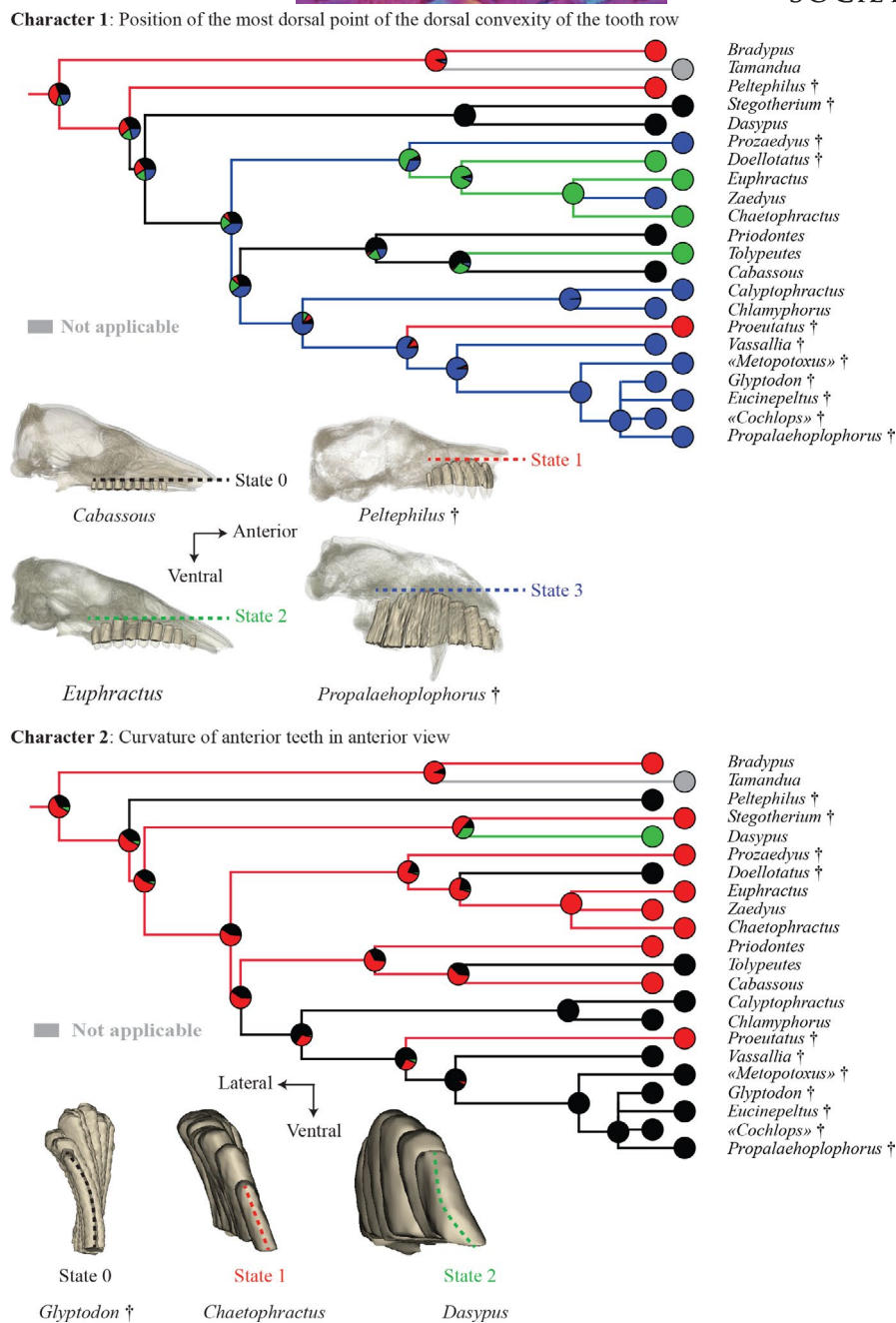
thus seems unrelated to craniofacial allometry, but this certainly requires further testing. In addition, we do not know whether allometry could affect the variation of the two aforementioned alveolar cavity characters, but the resemblance of small taxa (i.e., chlamyphorines) and large taxa (i.e., pampatheres and glyptodonts) seems to argue against this. In any case, our study brings to light interesting new variation regarding the upper tooth roots/alveolar cavities of cingulates, which have in the past been compared to one another on the basis of their number, the presence/absence of teeth in the premaxillary, tooth wear, histology, and the orientation of the long axis of the teeth relative to the long axis of the dental row (characters 1, 3–8—Gaudin & Wible, 2006; see discussion for glyptodonts regarding these characters—Ferigolo, 1985; González Ruiz et al., 2015; Scott, 1903).

Our initial observations on the overall spatial organization of the alveolar cavities point toward a resemblance among chlamyphorines, pampatheres, and glyptodonts, which would be partly compatible with molecular results (Delsuc et al., 2016; Mitchell et al., 2016). However, we believe further information could be gained by more detailed quantified analyses of shape to more fully reflect the complex spatial variation of the dental alveolar cavities.

Our analysis of selected intracranial canals described various aspects of their variation, which were unknown in cingulates until now. Further, it enabled us to propose five characters that provide potentially significant phylogenetic information. This analysis also demonstrated the need to further examine the variation of several structures (e.g., canal of the frontal diploic vein).

Both the trajectory and position of the nasolacrimal canal vary within the sample. A trajectory that is first directed medially at its posterior opening (i.e., the lacrimal foramen) represents a unique characteristic of glyptodonts within the Cingulata (Figures 7 and 8). The most informative feature on this canal regarding the relationships of glyptodonts with other cingulates is the relative height of its anterior opening within the nasal cavity (character 3—Figure 14). In most cingulates, this opening is positioned ventrally in the nasal cavity, close to the hard palate, whereas in glyptodonts, it is near the vertical midpoint of the cavity. The distribution of this character on the baseline cladogram supports the node linking *Proeutatus*, pampatheres, and glyptodonts (Figure 14), as in several previous morphological analyses (Billet et al., 2011; Engelmann, 1985; Gaudin & Lyon, 2017; Gaudin & Wible, 2006).

The course and connection between the sphenopalatine canal and the palatine canal is characterized by a complex pattern which is very peculiar in glyptodonts (Figures S6 and S7). The degree of fusion between the two canals was variable within our sample. Because of taphonomic issues, only "*Cochlops*" and *Glyptodon* allowed us to observe a fusion of the two canals in glyptodonts. This fusion was also present in the chlamyphorines. The uncertain condition in *Proeutatus* and pampatheres does not allow us to draw clear conclusions (character 4—Figure 14), but the distribution of this character on the baseline cladogram could be congruent with the morphological hypothesis of Mitchell et al., (2016) (Figure 14). However, caution is warranted, because the confluence between



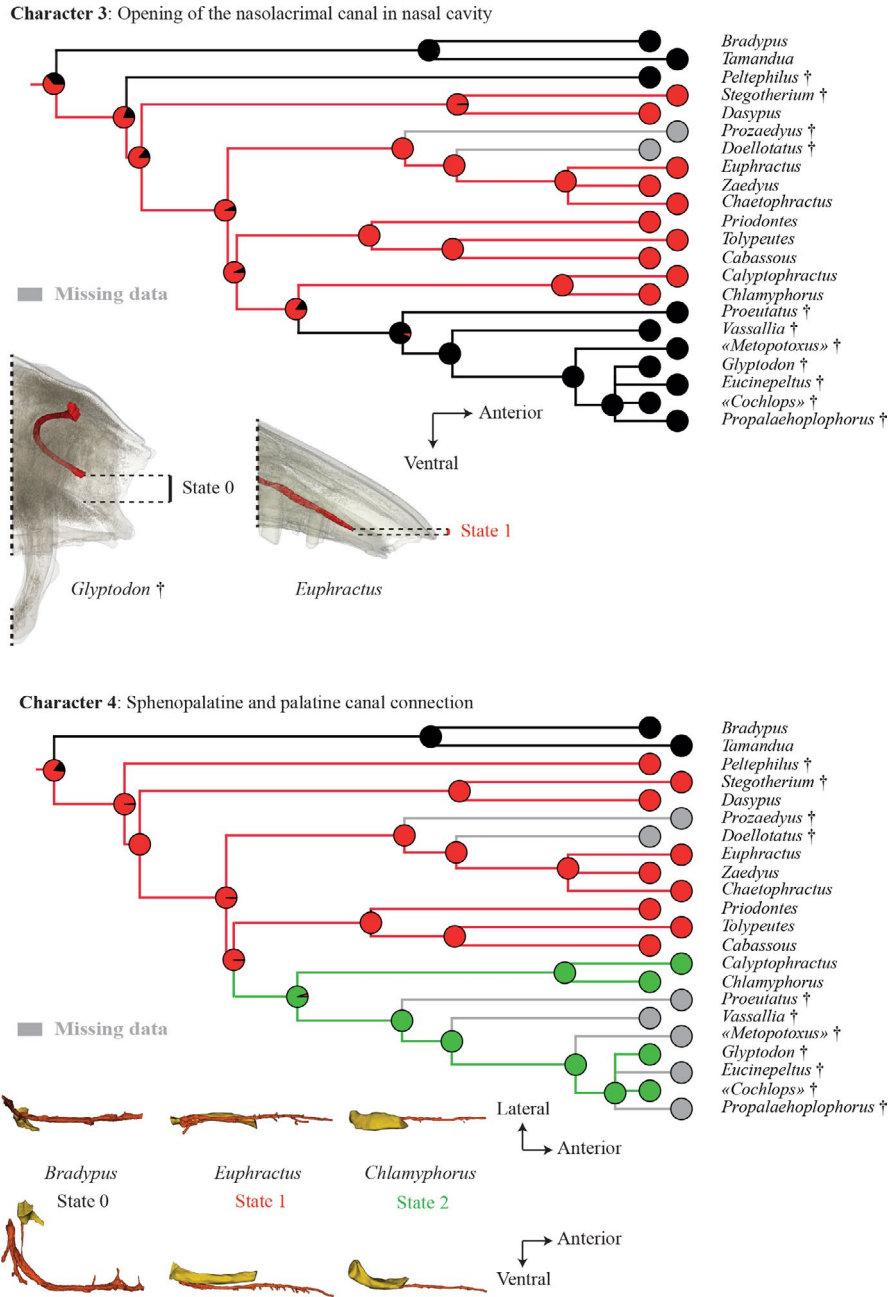
**FIGURE 13** Reconstructed evolutionary scenarios for the endocranial characters 1 and 2 plotted on the baseline cladogram with ancestral state estimation for internal nodes (see Materials & Methods, Data S1 and Figure S3). Each character is illustrated, and the relationship between color and coding is indicated within the figure itself. † represents extinct genera

the sphenopalatine foramen and the caudal palatine foramen is known to exhibit substantial intraspecific variation in several taxa (see character 71, Gaudin & Wible, 2006) including *Euphractus* (see Wible & Gaudin, 2004). In addition, the condition in pampatheres needs further investigation because a caudal palatine foramen has been identified in the floor of the sphenopalatine canal in *Holmesina* (see Gaudin & Lyon, 2017). Our study also reveals a strong resemblance between pampatheres and glyptodonts, which share a dorsal position of the sphenopalatine canal in relation to

the nasopharyngeal canal (Figure S7). This feature could have been coded as a character, but an investigation of potential allometry for this trait is required before coding.

The course, orientation, and connections of the transverse canal also vary in our sample. We were unable to identify this canal in *Tolypeutes*, chlamyphorines, and *Glyptodon* as well as *Bradypus*, *Peltephilus*, and *Stegotherium*. Its presence in *Proeutatus*, pampatheres, Miocene glyptodonts and many other cingulates suggests that it has been lost in *Glyptodon*. Apart from this potential loss,

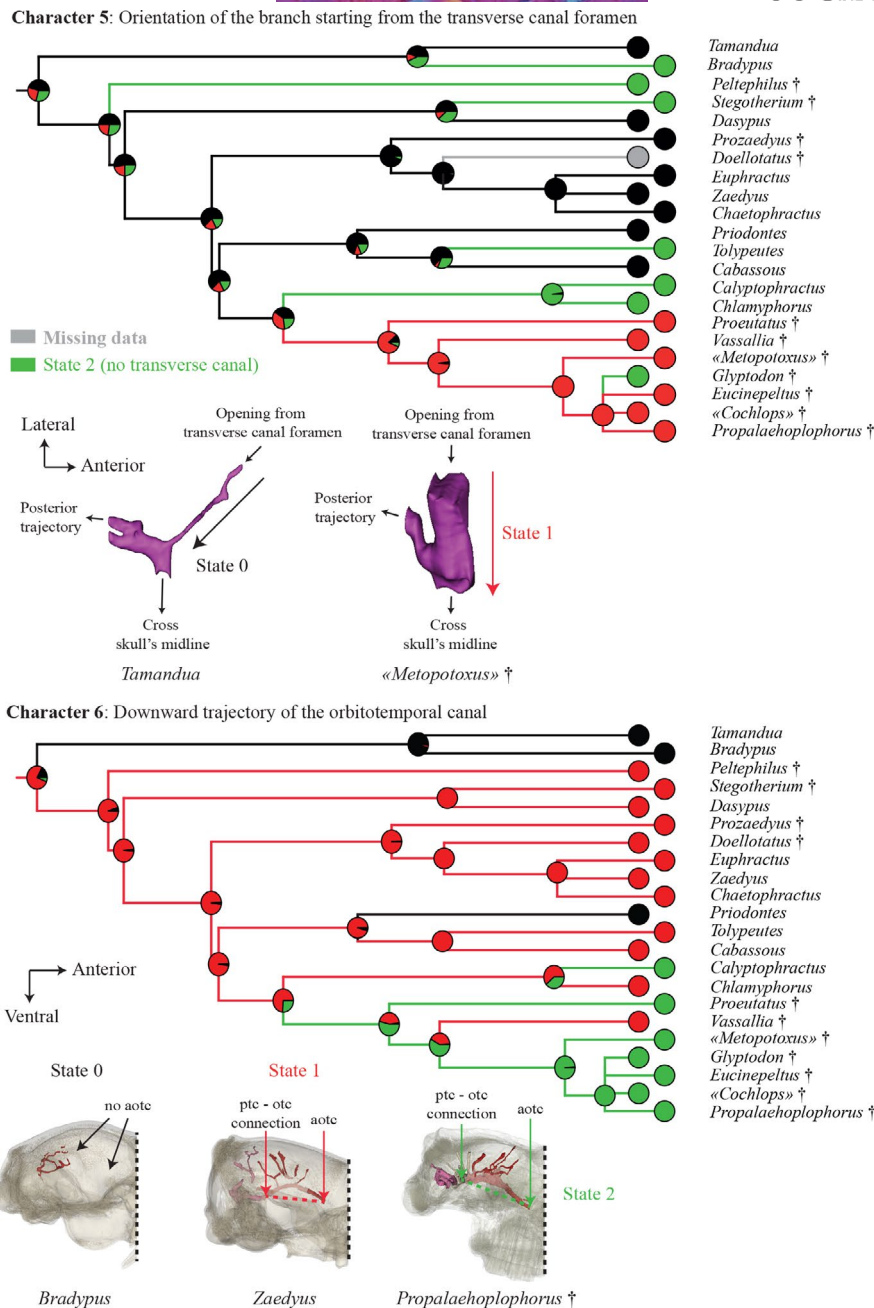




**FIGURE 14** Reconstructed evolutionary scenarios for the endocranial characters 3 and 4 plotted on the baseline cladogram with ancestral state estimation for internal nodes (see Materials & Methods, Data S1 and Figure S3). Each character is illustrated, and the relationship between color and coding is indicated within the figure itself. † represents extinct genera

the distribution on the baseline cladogram of the character corresponding to the orientation of the branch directly connected to the transverse canal foramen may support the common morphological hypothesis of a close relationship among *Proeutatus*, pampatheres, and glyptodonts (character 5—Figure 15; see also Billet et al., 2011; Engelmann, 1985; Gaudin & Wible, 2006; Gaudin & Lyon, 2017). Moreover, our work is congruent with the matrix of Gaudin and Wible (2006) regarding the presence/absence of the transverse canal foramen for most taxa (character 111) with the exceptions of *Tamandua* (coded as absent in Gaudin & Wible, 2006), *Stegotherium*

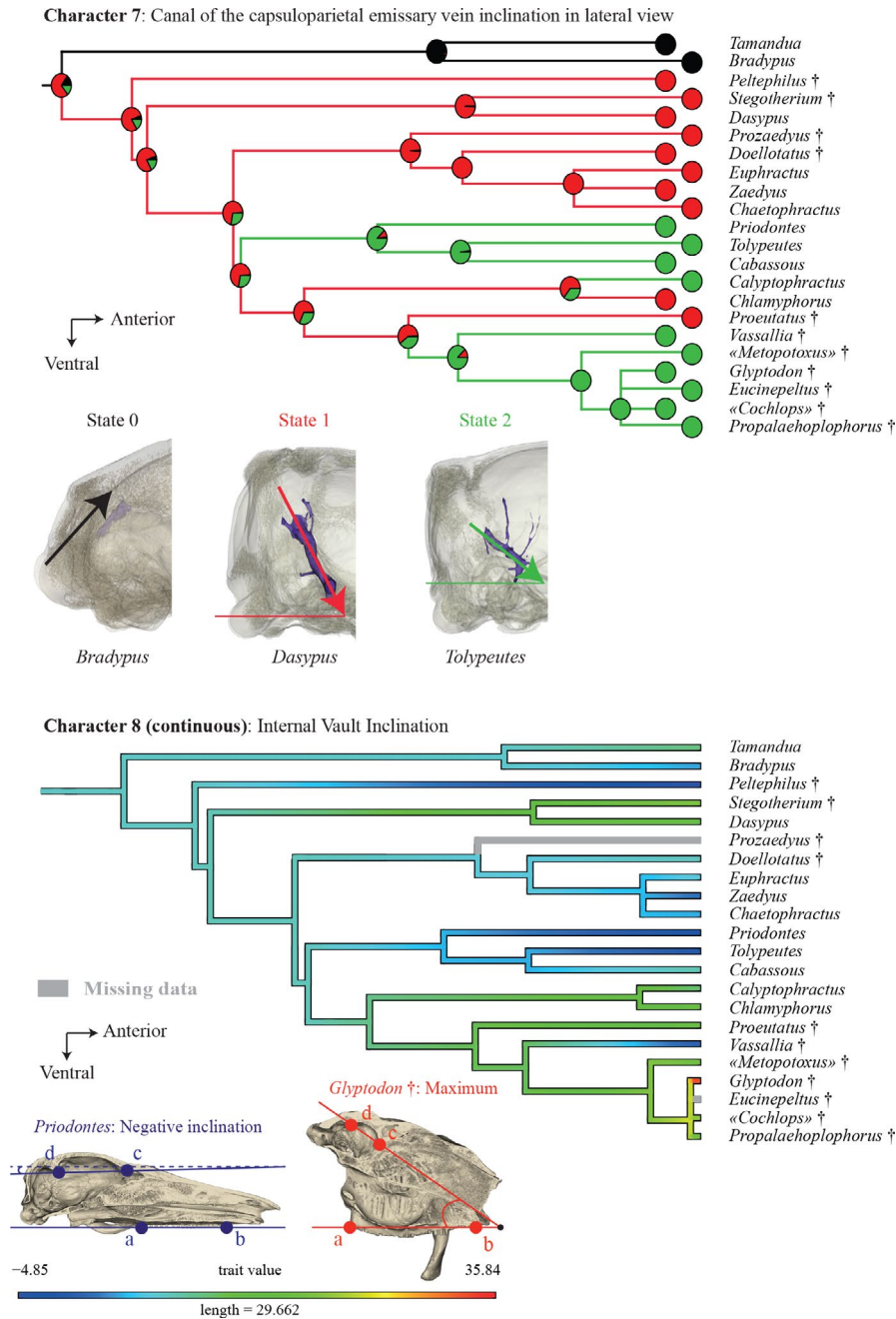
(coded as present in Gaudin & Wible, 2006), and *Doellotatus*. In the case of the latter, the specimen available for our study was poorly preserved for this character (coded as present in Gaudin & Wible, 2006). Our study also allows us to confirm the hypothesis of Wible and Gaudin (2004), which suggested that the canal crosses the midline of the cranium in *Euphractus sexinctus*. In their fetuses, they did not observe this transverse pattern and doubted whether it existed in adult specimens (Wible & Gaudin, 2004). We can confirm this pattern in our specimen (Figure 9). This circulation pattern is well known in several marsupials, but its putative presence is much less clear in



**FIGURE 15** Reconstructed evolutionary scenarios for the endocranial characters 5 and 6 plotted on the baseline cladogram with ancestral state estimation for internal nodes (see Materials & Methods, Data S1 and Figure S3). Each character is illustrated, and the relationship between color and coding is indicated within the figure itself. † represents extinct genera

placentals, for which the occurrence of a foramen for the transverse canal (often without knowing whether it crosses the cranium midline) is most often interpreted as a case of convergence (Sánchez-Villagra & Wible, 2002). In addition, our analysis of the youngest specimen of *Zaedyus* also suggests that the transverse canal is not formed in young euphractine individuals, in accordance with the observations of Wible and Gaudin (2004). However, in young dasypodines, we clearly observe this feature (Figures 3 and 5). For a large majority of taxa in our sample, we can confirm that a transverse canal crossing the midline is present in cingulates.

The distribution on the baseline cladogram of the character corresponding to the anterior course of the orbitotemporal canal also provided interesting information (character 6—Figure 15). *Calyptophractus*, *Proeutatus*, and glyptodonts share a strong downward trajectory of this canal, with an anterior opening located much lower in lateral view than its posterior connection of the orbitotemporal canal to the posttemporal canal. The evolutionary scenario for this character is, however, unclear for glyptodonts and their close allies, and homoplasy seems to be present (Figure 15). Our study also highlights the presence of a foramen for the anterior opening of



**FIGURE 16** Reconstructed evolutionary scenarios for the endocranial characters 7 and 8 plotted on the baseline cladogram with ancestral state estimation for internal nodes (see Materials & Methods, Data S1 and Figure S3). Each character is illustrated and the relationship between color and coding is explained within the figure itself. † represents extinct genera. For character 8 (= IVI), the angle between the anteroposterior axis of the cranium (defined by the line connecting the most posterior (a) and anterior (b) edges of the tooth row) and the straight line connecting the most dorsal point of the annular ridge (c) and the most ventral point of the tentorial process (d). Scale =1 operator # cm

the orbitotemporal canal in all cingulates except *Priodontes*, whereas this foramen was scored as absent in *Vassallia* and *Chlamyphorus* in Gaudin and Wible (2006, character 78) and is often unknown for other cingulate taxa. Based on our observation, there is no doubt that the canal opens in the orbitotemporal region in *Vassallia*, whereas this opening is described as absent in *Holmesina floridanus* Robertson, 1976 by Gaudin and Lyon (2017). The opposite discrepancy exists regarding the foramen for the frontal diploic vein, absent

in *Vassallia*, and described as present in *Holmesina floridanus* by Gaudin and Lyon (2017). As these two foramina are usually located relatively close to each other in the orbitotemporal region (a region that itself bears many foramina), we suggest that misidentifications may have occurred between the opening of these two canals, if they are present, or with ethmoidal foramina (Figure S4). The only way to confirm this hypothesis would be to scan Gaudin and Lyon's (2017) specimens of *Holmesina floridanus*.

Another character we investigated concerned the inclination of the canal for the capsuloparietal emissary vein in lateral view (character 7—Figure 16). All tolpeutines, *Calyptophractus*, *Vassallia*, and glyptodonts share a less steep inclination than that observed in other cingulates, with a part close to the anterior opening oriented almost directly to the anterior direction. The distribution of this character could provide support for the molecular hypothesis of a closer relationship of the glyptodonts with chlamyphorines and tolpeutines (Delsuc et al., 2016; Mitchell et al., 2016; Figure 16). However, the condition is reversed in *Chlamyphorus* and *Proeutatus*, creating homoplasy that casts doubt on the evolution of this trait. The trajectory of this canal has never been scored in prior phylogenetic analyses of the Cingulata. In the literature, only two characters have been scored that indirectly concern this canal, characters 119 and 120 of Gaudin and Wible (2006), related to the presence/absence of the postglenoid foramen and suprêmeatal foramen, respectively. These characters deliver contrasting groupings among cingulates (Gaudin & Wible, 2006).

Finally, the internal inclination of the cranial vault relative to the anteroposterior axis of the cranium also helps identify some cingulate taxa that may resemble glyptodonts in this regard (character 8; Figure 16). The distribution of this quantitative trait on the baseline cladogram indicates that the dasypodines, chlamyphorines, *Proeutatus*, and glyptodonts exhibit a strong IVI angle, whereas *Vassallia* possesses a very weak angle. This strong difference between glyptodonts and pampatheres was already highlighted by Tambusso and Fariña (2015b). The variation of this trait was not explained by allometry in our sample. A highly inclined vault may have been evolved independently in the dasypodines and in glyptodonts and their allies. Its distribution could provide support for a relationship among chlamyphorines, *Proeutatus*, and glyptodonts to the exclusion of pampatheres or the latter may have undergone a reversal (Figure 16). The vault inclination in these taxa seems to be partly congruent with the distribution of a character proposed by Billet et al. (2011) and discussed by Delsuc et al. (2016), that is, the dorsal position of the ventral surface of the auditory region relative to the palate (character 78—Billet et al., 2011). This is a characteristic of chlamyphorines, eutatines, pampatheres, and glyptodonts in Billet et al. (2011)'s analysis. The potential relationship of these two aspects of the braincase should be tested in future studies.

In conclusion, our investigation of the cranial canals and alveolar cavities highlighted several endocranial characters that support a resemblance of glyptodonts with chlamyphorines, *Proeutatus*, and pampatheres. Except for character 7, our investigation did not bring to light new, and strong resemblances between tolpeutines and glyptodonts that would support their close relationship as suggested by molecular studies (Delsuc et al., 2016; Mitchell et al., 2016). In short, our study of internal anatomy lends further credence to recent reports of a greater morphological resemblance between glyptodonts and chlamyphorines than tolpeutines (Delsuc et al., 2016; Mitchell et al., 2016). In this regard, it is worth noting that the chlamyphorines + tolpeutines relationship received distinctly lower statistical support than most other clades in mitogenomic analyses of cingulate phylogeny (Delsuc et al., 2016; Mitchell et al., 2016).

The congruence of these new internal characters with past morphological phylogenetic analyses of the group is difficult to evaluate since the results of the latter have been rather unstable (e.g., Abrantes & Bergqvist, 2006; Billet et al., 2011; Engelmann, 1985; Gaudin & Wible, 2006). However, a close resemblance of glyptodonts to the eutate *Proeutatus* and the pampatheres is supported by several characters in our study, which would be congruent with several recent phylogenetic hypotheses based on morphological data (Billet et al., 2011; Engelmann, 1985; Gaudin & Lyon, 2017; Gaudin & Wible, 2006). More generally, internal cranial anatomy seems to show similarities among chlamyphorines, *Proeutatus*, pampatheres, and glyptodonts (see also Tambusso et al., 2021), but new analyses are needed that incorporate this new data before firmer phylogenetic conclusions can be reached. Our study provides a non-exhaustive account of internal canals of the cranium. Internal structures still need to be studied to select new characters, in particular for endocranial traits that we briefly describe but do not explore in detail. This is notably the case with the canal for the frontal diploic vein or the position of the sphenopalatine canal in relation to the nasopharyngeal canal, whose patterns of variation remain unclear at the moment. Another promising way forward was also revealed by our developmental series, in which *Cabassous* varies in a manner different from that of *Dasypus* and *Zaedyus* as far as the inclination of the cranial roof is concerned (Figure 12). A better exploration of the developmental series with more specimens per species and more species sampled could provide a better understanding of their ontogenetic trajectories, which could then be coded for phylogenetic analyses (e.g., Bardin et al., 2017). Further studies are also needed to better understand the patterns of variation of the internal characters highlighted here, which in some respects may be affected by allometry, as we have highlighted for alveolar cavities and for the canal for the capsuloparietal emissary vein.

#### ACKNOWLEDGMENTS

We are grateful to Christiane Denys, Violaine Nicolas, and Géraldine Véron, (Muséum National d'Histoire Naturelle, Paris, France); Roberto Portela Miguez, Louise Tomsett, and Laura Balcells (British Museum of Natural History, London, UK); Neil Duncan, Eileen Westwig, Eleanor Hoeger, Ross MacPhee, Marisa Surovy, and Morgan Hill Chase (American Museum of Natural History, New York, USA); Nicole Edmison and Chris Helgen (National Museum of Natural History, Washington, DC, USA); Jake Esselstyn (Louisiana State University, Museum of Natural Sciences, Baton Rouge, USA); Manuel Ruedi (Muséum d'Histoire Naturelle, Geneva, Switzerland); Pepijn Kamminga, Arjen Speksnijder, and Rob Langelaan (Naturalis Biodiversity Center, Leiden, Holland); April Isch and Zhe-Xi Luo (University of Chicago, Chicago, USA); Adrienne Stroup and Bill Simpson (Field Museum of Natural History, Chicago, USA); Steffen Bock, Christiane Funk, Frieder Mayer, Anna Rosemann, Lisa Jansen, Kristin Mahlow, Johannes Müller, and Eli Amson (Museum für Naturkunde, Berlin, Germany); and Camille Grohé (University of Poitiers, Poitiers, France) for access to comparative material and/or to CT scans. We would like

to warmly thank Daniel Brinkmann (Yale Peabody Museum, New Haven, USA) for his precious help in organizing KLV's visit to the collections of the Yale Peabody Museum. Many thanks also to Sergio Ferreira-Cardoso (Institut des Sciences de l'Evolution) for his help with data acquisition during the trips of KLV. We thank Benoit de Thoisy (Institut Pasteur de la Guyane) and Clara Belfiore for their help with data acquisition, Olivia Plateau (University of Fribourg, Switzerland) for her help on the R script, and Cyril Le Verger for his help with segmentation. We thank Renaud Lebrun (Institut des Sciences de l'Evolution), Farah Ahmed (British Museum of Natural History), Miguel García-Sanz, Marta Bellato, Nathalie Poulet, and Florent Goussard (Platform AST-RX—Muséum National d'Histoire Naturelle) who generously provided help with CT scanning. We would like to warmly thank Christian de Muizon (Centre de Recherche en Paléontologie—Paris), Timothy Gaudin (University of Tennessee), Robert J. Asher (University of Cambridge), Sophie Montuire (University of Bourgogne), Isabelle Rouget (Centre de Recherche en Paléontologie—Paris), Allowen Evin and Lionel Hautier (Institut des Sciences de l'Evolution) for reviewing a previous version of this work in the PhD dissertation of KLV. Finally, we thank the two anonymous reviewers for their corrections and comments that helped us improve our manuscript. Some of the analyses were performed using the  $\mu$ -CT facilities of the Montpellier Rio Imaging (MRI) platform of the LabEx CeMEB. KLV acknowledges the financial support provided by the Sorbonne Universities/ED227 transhumance international grant program. In addition, the authors declare that they have no conflicts of interest.

#### AUTHOR CONTRIBUTIONS

KLV contributed to data acquisition, data analyses/interpretations, drafting of the manuscript, and study design. LGR contributed to data acquisition and critical revision of the manuscript. GB contributed to data acquisition, study design, and critical revision of the manuscript.

#### DATA AVAILABILITY STATEMENT

A large part of specimens scanned will be available on MorphoMuseum for 3D models and on MorphoSource for scans. Specimens scanned by LGR will be available upon request to LGR. The list of specimens is given in Table S2.

#### ORCID

Kévin Le Verger  <https://orcid.org/0000-0001-8498-7372>

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## SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

**How to cite this article:** Le Verger, K., González Ruiz, L.R.&Billet, G. (2021) Comparative anatomy and phylogenetic contribution of intracranial osseous canals and cavities in armadillos and glyptodonts (Xenarthra, Cingulata). *Journal of Anatomy*, 00, 1–30. <https://doi.org/10.1111/joa.13512>