

Anguimorph lizards from the lower Eocene (MP 10–11) of the Cos locality, Phosphorites du Quercy, France, and the early evolution of Glyptosaurinae in Europe

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1 ARTICLE

2	Anguimorph lizards from the early Eocene (MP 10-11) of the Cos locality, Phosphorites du
3	Quercy, France and the early evolution of Glyptosaurinae in Europe
4	
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15	
16	ABSTRACT— Squamate faunas from the early Eocene of Europe are rare. A newly
17	discovered early Eocene (MP 10-11) Cos locality in southwestern France has yielded an
18	assemblage of anguimorph lizards that sheds light on the early evolution of this group.
19	Among them is a new glyptosaurine lizard Sullivania gallica gen. et sp. nov., based on frontal
20	material with a unique pattern and distribution of polygonal osteoderms that is distinct from
21	that of the middle and late Eocene Placosaurus from France. It slightly resembles the
22	stratigraphically older glyptosaurine Gaultia silvaticus from the earliest Eocene of North

America, but differs from it in several aspects. Although fossils of some members of 23 24 Glyptosaurinae (sensu this paper; Glyptosaurini in previous taxonomies – Glyptosaurinae is returned to the family level originally proposed by Marsh) have been documented rarely from 25 the middle and mainly from the late Eocene of Europe, they are virtually unknown from the 26 early Eocene. One isolated osteoderm is referred only to Anguioidea indet. This specimen 27 resembles osteoderms seen in Anguinae, provided that the absence of tuberclulate 28 ornamentation is not caused by abrasion. 29 Varanoids are represented by an isolated frontal referred to Palaeovaranidae indet. It 30 possesses a complex ornamentation composed of mounds on the dorsal surface. A maxilla is 31 32 identified as an indeterminate varanoid based on the plicidentine infolding along the bases of the preserved teeth. 33 Although incomplete, this Cos lizard assemblage is an important and rare discovery 34 that provides a glimpse into the ecosystems and paleobiodiversity of the early Eocene in 35 western Europe. 36 37 LSID urn:lsid:zoobank.org:pub:5C6376A3-CC26-4CF1-BE56-E7279D508D5D 38 39 40 **INTRODUCTION** The Cos fissure fill was the first "phosphatière" discovered in the Quercy by J. A. 41 Poumarède in 1865 (Pulou, 1980), but one of us (DV) discovered vertebrate fossils in this 42 locality only more recently. Cos constitutes the seventh pre-upper Eocene locality of 43 Phosphorites du Quercy [for others see Astruc et al. (2003): Pasturat (MP 8+9), Viélase and 44 45 Cazals (both MP 10-11), Cuzal, Parnac and Grélaou (all MP 13)]. Based on the studies of the primates (Godinot et al., 2021) and rodents (Vianey-Liaud et al., 2022), the vertebrate fauna of 46 the Cos site has been dated as MP 10–11, a zone that spans the late Ypresian and early Lutetian. 47

Thus, it corresponds to the Early Eocene Climatic Optimum (EECO; see Inglis et al., 2020), which is dated as 53.3 to 49.1 Ma. EECO was a time of major climatic and environmental change caused by perturbations to the global carbon cycle (Hyland et al., 2017). This makes the Cos locality particularly important because the early Eocene terrestrial fossil record in Europe is poorly known.

There have been only a few early Eocene sites where assemblages of amphibians and 53 reptiles have been described. These include: Dormaal (MP 7) in Belgium (Augé, 1990, 1992; 54 Augé & Smith, 1997, 2002; Augé et al., 2022; Čerňanský et al., 2022; Folie et al., 2013; 55 Hecht & Hoffstetter, 1962; Sullivan et al., 2012); Silveirinha (~MP 7) in Portugal (Rage & 56 57 Augé, 2003); the early Eocene of (different localities in) Catalonia, Spain (Bolet, 2017); and some localities in France, notably Prémontré (~MP 10; Augé, 2003, 2005; Augé et al., 1997), 58 Mutigny (MP 8–9; Čerňanský et al., 2020; Augé et al., 2022) and La Borie (MP 8–9) (Laurent 59 60 et al., 2010). There are also some other famous early Eocene localities, which yielded snakes, turtles, and crocodiles, such as Monte Bolca, Italy (Seghetti et al., 2022) and the English 61 62 localities in Isle of Sheppey, Bracklesham, and Kingston (Owen, 1841, 1850; Georgalis & Joyce, 2017; Smith & Georgalis, 2022). In contrast, such assemblages are relatively well-63 known for the middle and late Eocene in Europe, in part thanks to exceptional localities such 64 as Geiseltal and Messel in Germany (Čerňanský & Smith, 2018, 2019 Keller, 2009; Smith et 65 al., 2018; Sullivan et al., 1999; Villa et al., 2022) and in some of the localities of Phosphorites 66 du Quercy in France (Augé, 2005; Čerňanský et al., 2015; Georgalis et al., 2021). Here we 67 report on the fossil anguimorph lizards from the newly discovered Cos locality, on the 68 southern Quercy limestone plateau, southern France. 69 Modern Anguimorpha includes Anguidae, Diploglossidae, Xenosauridae, 70

71 Helodermatidae, Lanthanotidae, Varanidae and Shinisauridae. Anguidae represent a

72 diversified and widely distributed group of reptiles of which Anguinae is the most derived

clade (Augé, 2005; Uetz et al., 2023). Although there have been recent alterations to 73 phylogenetical conceptions that have been stable for decades (e.g., Burbrink et al., 2020; 74 Pyron et al., 2013) and the position of Diploglossidae is still debated (as a subclade of 75 Anguidae, Conrad 2008; Gauthier et al., 2012; Pyron et al., 2013; as a distinct clade outside of 76 Anguidae, Burbrink et al., 2020; Zheng & Wiens, 2016), aside from Anguinae three other 77 extant clades were traditionally included in Anguidae. These are Gerrhonotinae, Anniellinae, 78 and the extinct Glyptosaurinae (sensu McDowell & Bogert, 1954; see Gauthier et al., 2012; 79 Georgalis et al., 2021; Sullivan, 1979, 2019; note, however, that if Diploglossidae is treated at 80 the family level, then Glyptosaurinae must also be returned to the family level originally 81 82 proposed by Marsh, 1872, as members of this clade are much morphologically distinct 83 compared to Anguinae, see below). Glyptosaurinae (sensu McDowell & Bogert, 1954) inhabited North America, Europe, and Asia and are known from the Cretaceous to the late 84 85 Paleogene-Oligocene (Čerňanský and Augé, 2019; Conrad & Norell, 2008; Keller, 2009; Sullivan, 1979, 2019). This clade was traditionally divided into the tribes Glyptosaurini and 86 the paraphyletic "Melanosaurini" (Cicimurri et al., 2016; Estes, 1983; Sullivan, 1979, 2019). 87 Among the former, *Placosaurus* Gervais, 1848–1852 is the only known member of the tribe 88 in Europe (see Sullivan, 2019; Sullivan & Augé, 2006; Sullivan et al., 2012). It is represented 89 90 by two currently valid species from the late Eocene of France (only the two species are definitely members of the genus, whereas the others are only questionably referred to 91 *Placosaurus*), type of which are cranial remains. This is very low diversity compared to the 92 93 North American glyptosaurine genera such as *Glyptosaurus* Marsh, 1871, *Paraglyptosaurus* Sullivan, 1979, Helodermoides Douglass, 1903 and Gaultia Smith, 2009 (Smith, 2009, 2011; 94 Sullivan, 1979). Moreover, although members of the clade were well documented from the 95 early Eocene in North America, glyptosaurines are largely unknown for most of the early and 96 middle Eocene of Europe (Sullivan, 2019). In the early Eocene of Europe, the glyptosaurine 97

taxon ?Placosaurus ragei Sullivan, Augé, Wille & Smith, 2012 and Placosauriops-like 98 "melanosaurine" are known from Dormaal (MP 7). The Dormaal glyptosaurine ?Placosaurus 99 ragei is known based on the holotype dentary and a referred parietal. However, the current 100 taxonomy of species of *Placosaurus* is based on the morphology of the frontal (Sullivan & 101 Augé, 2006), so its generic assignment is questionable (Sullivan et al., 2012). Indeterminate 102 material (frontal and parietal) of glyptosaurines is also described from Silveirinha (~MP 7; 103 104 Rage & Augé, 2003). In Asia, Stenoplacosaurus mongoliensis (Sullivan, 1979) was described from the middle Eocene of Mongolia (Sullivan & Dong, 2018; Helodermoides mongoliensis 105 in Sullivan 1979; Placosaurus mongoliensis in Sullivan and Augé, 2006). This taxon is also 106 107 based on the frontal morphology. Additionally, several isolated osteoderms of glyptosaurines, 108 plausibly belonging to this early Eocene lineage, were described from the early Oligocene of Mongolia (Čerňanský & Augé, 2019). Thus, Glyptosaurinae has a continuous history in Asia 109 110 at least until the Oligocene, although disappeared in Europe at the Eocene-Oligocene boundary (Rage, 2013). However, their early history in Europe remains largely unknown. All 111 the specimens in this study are from the Cos fissure fill, Phosphorites du Quercy, 2.5 km 112 southwest of Caylus, France. 113

114

Institutional Abbreviations— AMNH, American Museum of Natural History, New York;
MNHN, Museum National d'Histoire Naturelle, Paris; UCMP, University of California
Museum of Paleontology, Berkeley, USA; UM-COS, Université de Montpellier, France, Cos
collection.

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120

MATERIAL AND METHODS

121

122 Specimens examined and terminology

123 The studied material is housed at the University of Montpellier, cataloged under 124 individual UM-COS-numbers. The standard anatomical orientation system is used throughout 125 this paper, and terminology describing individual bone structures is based on Rage & Augé 126 (2010).

127

128 X-ray Microtomography, Three-Dimensional Visualization, Photography and 129 Reconstruction

The fossil specimens were imaged on micro-computed tomography (CT) at the micro-130 CT facility, Slovak Academy of Sciences in Banská Bystrica, Slovakia, using a Phoenix mikro-131 132 CTv/tome/x L240. The settings were as follows: VxSize = 0.019999999; Current = 150; Voltage = 130; Inttime = 20000; Average = 2; Steps = 1600; Steps 360 = 2200; Filter = 0.1 mm Cu. The 133 images were recorded over 360°. The CT data sets were analysed using VG Studio Max 3.1. 134 and Avizo 8.1. The photographs of the UM-COS-1011 maxilla were taken with a Leica M205 135 C binocular microscope with an axially mounted DFC 290 HD camera; software: LAS (Leica 136 Application Suite) 4.1.0 (build 1264). The image processing program ImageJ (Schneider et al., 137 2012) was used for measurements. For frontal reconstruction, the missing areas were restored 138 139 by mirror imaging of the holotype and paratype, and estimation of the posterior portion is based 140 on the preserved paratype. The holotype (UCMP 216000) and paratype (UCMP 150966) of Gaultia silvaticus were drawn from photographs published by Smith (2009: fig. 18). The 141 holotype of Stenoplacosaurus mongoliensis (AMNH 6669 left frontal) was drawn based on 142 photographs published by Sullivan & Dong (2018: fig. 1). 143

144

145 Geological Setting and Associated Fauna

Located in the southern part of the 'Causses du Quercy' plateau, the fissure fills of Coslie within the Middle Jurassic (late Bajocian/Bathonian) marine limestones of the Cajarc

Formation (44°13'11.20"N, 1°44'58.21"E; Fig. 1). The site is a northwest-southeast trending 148 large excavation (~80 m x 50 m), with the phosphate ore deposits nearly depleated. Positioned 149 at the eastern end of the excavation, the fossiliferous fissure-fill at Cos is unlike those found in 150 other localities of the Phosphorites du Quercy in that the strata that contain the fossiliferous unit 151 are composed of small beds of granular and porous stratified limestones, alternating with red 152 clays containing small pisolites. There are still uncertainties about geometric and stratigraphic 153 relationships between the limestones and the interbedded red clays. There are few scattered 154 vertebrate fossils through the entire section. The fauna recovered includes amphibians, 155 squamates, crocodilians, birds, and more than twenty mammal species among which only 156 157 primates and rodents have been studied (Godinot et al., 2021; Vianey-Liaud et al., 2022). All of the specimens described herein are from the Cos fissure fill of the Phosphorites du Quercy 158 region and are early Eocene (MP 10-11 interval) in age. 159

160

161

SYSTEMATIC PALEONTOLOGY

162

Taxonomic note—If most clades previously considered subfamilies of Anguidae sensu 163 Camp, 1923 - viz., Anguinae, Anniellinae and Diplglossinae - are now considered taxa at the 164 family level, and there is no evidence that glyptosaurs are part of any of those radiations, then 165 Glyptosaurinae sensu McDowell & Bogert, 1954 must also be returned to the family level 166 originally proposed by Marsh (1872). Accordingly, we consider Glyptosauridae Marsh, 1872 167 to contain the subfamilies Melanosaurinae Sullivan, 1979, which is paraphyletic, and 168 Glyptosaurinae Marsh, 1872, which correspond to the tribes Melanosaurini and Glyptosaurini 169 of Sullivan (1979). The name "Placosauridae" (e.g., Cope, 1877; Kuhn, 1940) is a junior 170 synonym of Glyptosauridae (Estes, 1983; McDowell & Bogert, 1954), so it does not affect 171 172 our new status.

173	
174	SQUAMATA Oppel, 1811
175	ANGUIMORPHA Fürbringer, 1900
176	ANGUIOIDEA Gray, 1825
177	GLYPTOSAURIDAE Marsh, 1872 (emended status)
178	GLYPTOSAURINAE Marsh, 1872 (sensu this paper)
179	SULLIVANIA gen. nov.
180	
181	Type species— Sullivania gallica sp. nov.
182	Etymology—The genus is named in recognition of the American paleontologist
183	Robert M. Sullivan for his valuable contributions to glyptosaurine morphology,
184	paleobiogeography, and paleobiodiversity.
185	Diagnosis —As for <i>Sullivania gallica</i> sp. nov., the only known species.
186	
187	Sullivania gallica sp. nov.
188	(Figs. 2, 3, 4A)
189	
190	Etymology — The specific epithet <i>gallica</i> is based the Latin "gallicus" in reference to
191	the country of France, where the fossils were found.
192	
193	Holotype— UM-COS-1001, greater anterior portion of fused frontals.
194	Paratypes— UM-COS-1000, incomplete right frontal; and UM-COS-1003, posterior
195	region of the right frontal.
196	Differential Diagnosis— Glyptosaurine differing from all other Glyptosauridae in
197	having unique pattern of irregular cephalic osteoderms covering the frontal where the first two

anterior central osteoderms straddle the midline, and are coalesced or fused, narrow, 198 anteroposteriorly elongate, leaf-shaped, reaching only slightly further anteriorly than the 199 much larger, anteroposteriorly elongated lateral osteoderm with rounded lateral margin. This 200 is in a sharp contrast to several wide, chevron-shaped osteoderms in Gaultia (in Sullivania, 201 202 only the second lateral left and right osteoderms appear expanded medially, having contact in the mid-line) and high and regular division of dermal armor into discrete hexagonal 203 osteoderms in *Placosaurus*, *Stenoplacosaurus*, *Glyptosaurus*, *Paraglyptosaurus*, 204 Helodermoides, and Eoglyptosaurus. It is further distinguished from all other glyptosaurines 205 by the following combination of features of the frontal: (1) osteoderms are apically flat, as 206 207 Gaultia, not thick as in Placosaurus, and not inflated (or bulbous) as in Glyptosaurus, 208 Eoglyptosaurus, Helodermoides and Stenoplacosaurus; (2) generally rather fever osteoderms contra numerous in Placosaurus (especially Placosaurus estesi), Glyptosaurus, 209 210 Eoglyptosaurus and Helodermoides; (3) concave lateral margins unlike in the North American Helodermoides; (4) posterolateral process not distinctly laterally expanded and expansion 211 includes only the posterior 1/3 of the bone, unlike in Placosaurus estesi and 212 Stenoplacosaurus, in which the lateral expansion starts at the posterior 2/3 of the bone; and 213 214 (5) frontals fused only in the late ontogeny (fusion is present only in the larger, robust 215 specimen). In Stenoplacosaurus, the frontals are unfused. Type locality and horizon—Cos, fissure fill in the Quercy region (southwestern France); 216

early Eocene (MP 10–11 interval); the species is so far known exclusively from the type
locality.

219 **Description**

Frontal—The holotype (UM-COS 1001, Fig. 2) represents fused right and left
frontals (although traces of fusion are still recognizable). Its lateral margin is slightly concave
and gradually widens posteriorly. The dorsal area surrounding the mid-orbital margin slopes

slightly toward the orbit. The posterior portion of the bone is missing due to breakage. Thus, 223 only the anterior and mid-portions are preserved. The maximum width of the preserved 224 frontal is 11.7 mm. The element is more robust than UM-COS-1000 and is inferred to be from 225 a larger (older) individual. The external surface is largely covered by ornamented osteodermal 226 shields of various irregular, polygonal to roughly ovoid outline. They are completely fused to 227 the underlying bone and are separated one from another by sulci (note, however, that it is not 228 229 clear whether the osteodermal boundaries coincide with the epidermal scale boundaries). The ornamentation is formed by small rounded discrete tubercles that are regularly and densely 230 arranged. The exception is in its anterior section, where a broad semi-elliptical smooth surface 231 232 is located. It forms the articular facet for nasal (which is preserved on the right side). Here, the 233 bone expands into a prominent, pointed anteromedial (nasal) process. The process is triangular. The anterolateral process is only partly preserved on the right side, but this 234 preserved portion extends much less anteriorly than the anteromedial one. Posterior to this, 235 three osteodermal shields are present. The first one is an anterior central shield. Here, the left 236 and right frontal are fused along the mid-line. Thus, the first two anterior osteoderms straddle 237 the midline and are coalesced or fused into the one. The first central shield is narrow, 238 239 anteroposteriorly elongate and more or less leaf-shaped. It extends only slighty more 240 anteriorly than the lateral shields. The lateral osteodermal shields are large, much larger that the central one (completely preserved on the right side). The medial margin of the lateral 241 shields is V- shaped, whereas the lateral margin is rounded. The medial peak of V-shaped 242 243 medial margin indicates a mediolateral short sulcus, where the first anterior central shield ends. Posterior to the sulcus in the midline, there is a second central shield. The more 244 posterior region bears several cracks and thus, it is difficult to properly identified the original 245 borders of the osteodermal shields (Fig. 2A, B). However, it appears that the both second 246 lateral shields are medially expanded (best visible on the left side). The left and right shields 247

seem to have a short medial contact. If this is the case, these second lateral osteoderms are
mediolaterally wide rather than anteroposteriorly long (in contrast to the first lateral shield).
In this case, the second central osteodermal shield has truncated appearance of the posterior
portion due to this contact and thus, being trapezoidal in shape.

In ventral view, a large and robust frontal cranial crest (crista cranii frontalis) is 252 visible. Its anterior portion is expanded forming a well-defined and rounded prefrontal 253 (=subolfactory) process. It is, however, only partly preserved (its end is damaged; Fig. 2C). 254 The anteromedial margin of this crest is thin and sharp. Posteriorly, it widens, gradually 255 diminishing dorsally. The anterior portion of the frontal crest, anterior to the subolfactory 256 257 process, is less deep, forming a sharp, medially directed ridge. The right and left branches join 258 together in the anterior mid-line, forming a sharp angle. Note, however, that there is the matrix that is still adhering to the ventral side of the frontal between the anterior portions of 259 260 the left and right frontal cranial crests. It obscures much of the medial ventral surface of the frontal. In the anterior region, lateral to the frontal crest (including its lateral surface), a large 261 facet for the prefrontal is located (the right one is preserved; Fig. 2D). Prefrontal and 262 postfrontal are not in a contact, and did not exclude the frontal from the orbital border. 263

UM-COS-1000 and UM-COS-1003 (paratypes) ---UM-COS-1000 (Fig. 3A-F) is a 264 265 nearly complete right frontal, with only the anterior end broken. The maximum anteroposterior length of this incompletely preserved element is 15 mm. It is robust, although 266 less than the holotype. UM-COS-1003 (Fig. 3G, H) represents the posterior portion of the 267 right frontal and this specimen is much more robust than the UM-COS-1000. The 268 posterolateral section of UM-COS-1000 protrudes into a short, but robust, posterolateral 269 270 process (the expansion starts in the posterior 1/3 of the bone). Here, the bone is widest – the maximum mediolateral width of this part of the bone is 9.1 mm. Further anteriorly, the frontal 271 narrows and the lateral margins are slightly concave. Thus, the bone appears to be narrowest 272

at about mid-orbit (the minimum width of the single right frontal in this area is 5.9 mm). The 273 external surfaces of both specimens are almost completely covered by tuberculated 274 osteoderms of various irregular, polygonal (rhomboidal, hexagonal) to roughly ovoid outlines. 275 The osteoderms are completely fused to the frontal and separated one from another by sulci. 276 The ornamentation is formed by small rounded discrete tubercles that are regularly and 277 densely arranged. The osteodermal shields on the posterior portion in UM-COS-1000, close to 278 279 the more or less straight contact with the parietal, are weakly ornamented. Only few tubercles are present in this section on an otherwise smooth surface; however, the dorsal surface in this 280 region seems to be slightly eroded. The posterolateral corner is smooth, although it appears 281 282 that some partial covering was originally here but now is lost. The evidence of this is that a 283 partly preserved medial margin of the shield still has several tubercles. Thus, the posterior portion of the right frontal possessed four osteodermal shields arranged in a mediolateral row. 284 285 Thus, the frontal is relatively broad. These osteroderms are anteroposteriorly long, roughly trapezoidal except for the third one (Fig. 3B), which has a triangular end due to hexagonal 286 shape of neighboring osteoderms in the second row. This second posterior row, located 287 anterior to the first one (at the level of the anterior beginning of the postfrontal facet), 288 possesses three osteoderms. They differ in shape from those forming the first posterior row, 289 290 being polygonal (roughly hexagonal). The same condition is present in UM-COS-1003, although the ornamentation is strongly developed here. In the mid-region of UM-COS-1000, 291 there is a shallow light groove close to the orbital margin. This groove extends almost parallel 292 293 to this margin (Fig. 3B).

In ventral view, a frontal cranial crest can be observed in both specimens. Posteriorly, it widens, gradually diminishing dorsally. On the lateral side of the crest, a foramen is visible. The crest fades out at the beginning of the posterolateral process of the frontal (in both paratypes) and does not continue to the posterior end of the bone. Here, the large rugose facet

for the postfrontal is clearly visible laterally. In UM-COS-1003, the wedge-shaped facet for parietal tab is visible dorsal to the frontoparietal suture. A facet for the prefrontal is preserved only in UM-COS-1000. The prefrontal and postfrontal are not in a contact, and thus did not exclude the frontal from the orbital border.

302 Virtual microanatomy and histology—The micro-CT scans of the frontals revealed
303 a bone structure with a very similar internal microanatomy in terms of a vascular network and
304 spongiosis (Fig. 2E, F; Fig. 3E, F). It revealed a large and complex meshwork of numerous
305 cavities. They are irregular, bubble-shaped, and some are interconnected. Thus, the bone
306 appears to be less compact in both axial and coronal sections. Note that the finer
307 histological details such as growth marks and cell lacunae of the bone are not visible.
308 Remarks— UM-COS-1001 is designated as the holotype because it is from a more

mature (adult) individual. The paratypes UM-COS-1000, the nearly complete right frontal, 309 310 and UM-COS-10003, the posterior part of the right frontal, are nearly identical, although UM-COS-1003 is slightly more robust. UM-COS-1000 and UM-COS-10003 have the identical 311 arrangement of osteoderms where they overlap the holotype UM-COS-1001. They can add to 312 the diagnosis a unique character state for Glyptosaurinae - the osteoderms on the 313 314 posteriormost portion of the frontal are anteroposteriorly elongate and roughly trapezoidal. 315 Four osteoderms of equal size are arranged in a mediolateral row. The second posterior row (located anterior to the first row) possesses three osteoderms, which are roughly hexagonal. 316 Although all frontals are incomplete, preservation of parts shared by all frontals permits for 317 318 reconstruction of the missing areas based on mirror imaging of the right and left elements (UM-COS-1000). The anterior portion is reconstructed based on the UM-COS-1001 (Fig. 319 2A). The reconstructed image clearly shows that it is very different from the earliest Eocene 320 North American glyptosaurine Gaultia, the late Eocene European Placosaurus and the middle 321 Eocene Asian Stenoplacosaurus (Fig. 4). 322

323	With regard to the virtual microanatomy, microanatomical studies on fossil anguioids
324	(i.g., anguids) have previously been conducted. These published works deal with parietals of
325	Ophisaurus holeci (Georgalis & Scheyer, 2021), Ophisaurus spinari (Syromyatnikova et al.,
326	2022), and Pseudopus pannonicus (Loréal et al., 2023).
327	
328	Glyptosauridae indet.
329	(Figs. 5, 6A-J)
330	
331	Referred Material—UM-COS-1002, medial portion of left maxilla; UM-COS-1011,
332	posterior portion of right maxilla; UM-COS-1004 and 1005, cephalic osteoderms and UM-
333	COS-1006 and 1007, dorsal trunk osteoderms.
334	
335	Description
336	Maxilla—Two incomplete maxillae were recovered (Fig. 5), representing much of the
337	posterior halves of the bone with the posterior ends broken on both specimens. The left
338	maxilla (UM-COS-1002, Fig. 5A-D) has seven tooth positions with four-and-a-half teeth
339	remaining in situ. The right maxilla (UM-COS-1011, Fig. 5E-H) has four posterior teeth and
340	two broken teeth. The nasal process of the maxilla is partly preserved in both specimens,
341	gradually decreasing in height posteriorly. Its external surface is ornamented with tubercles
342	and is best preserved in UM-COS-1011. The extent of this tuberculated surface is not well
343	delineated as the sulci are not recognizable (see Fig. 5A, E and remarks). The supralabial area,
344	below the ornamented field, is pierced by supralabial foramina (two are preserved in UM-
345	COS-1002 and one in 1011). In medial view, the supradental shelf is prominent. The shelf
346	slopes slightly posteroventrally from the level of the third preserved tooth position (counted
347	from posterior). The dorsal area of the shelf is concave, forming a longitudinal shallow

depression for articulation with the jugal. The shelf itself expands medially, and this
expansion (best preserved in UM-COS-1011) reaches its maximum at the level among third
and fifth tooth positions (counted from the posterior end). Here, the contact with the palatine
is present. UM-COS-1002 bears the posterior opening of the superior alveolar foramen (Fig.
5F, H). It is large, elliptical, and located at the level of the fifth tooth position (counted from
posterior).

Dentition—The tooth implantation is pleurodont. The teeth are heterodont with the lingual sides of the teeth expanding medially and gradually increase in robustness posteriorly. The apices of posterior three teeth are blunt, mesiodistally robust, and possess distinct, welldefined, mesiodistally straight cutting edges. The lingual and labial surfaces of the tooth apices are distinctly striated. The first preserved anterior tooth is less robust and labiolingually compressed.

Remarks— The maxillae from the Cos site may belong to Sullivania gallica, but are 360 here considered Glyptosauridae indet. owing to lack of definitive characters that would allow 361 362 positive identification. The maxilla is similar to that of the late Eocene species Paraplacosauriops quercyi in the lateral ornamentation of the surface of the nasal process of 363 maxilla and the presence of bulbous teeth (see Auge & Sullivan, 2006; Georgalis et al., 2021). 364 The ornamentation lacks discrete hexagonal/polygonal osteoderms that would clearly indicate 365 it was a glyptosaurine. Therefore, the maxilla may be from some unknown "melanosaurine" 366 similar to Paraplacosauriops. 367

Osteoderms—Four tuberculated osteoderms were recovered from the Cos site (Fig. 6A-J). Two are hexagonal/polygonal in shape (Fig. 6A-E) and two are rectangular/rhombic (Fig. 6F-M). The former are from the skull and the latter are from the body. The tubercles are prominent on the cephalic osteoderms and less so on the body osteoderms but this may be an artifact of preservation. The keel and the smooth overlap surface are absent on the first type of

osteoderms. These osteoderms are also slightly thicker than the second (rectangular) type. The 373 374 internal surfaces of the cephalic osteoderms are pierced by numerous foramina along with several grooves and ridges that extend anteroposterly. The rectangular body osteoderms (Fig. 375 6F-J) are distinguished by a prominent contact surface that is nearly one-third the 376 anteroposterior length. It is slightly higher than the posterior ornamented portion, from which 377 it is separated by a transverse groove. The latter is covered by discrete tubercles of various 378 sizes. A blunt, weak and slightly oblique keel is present midway along the posterior margin of 379 the osteoderm. The internal surface is pierced by three foramina located in the central region. 380 Remarks—Taxonomic assignment of isolated osteoderms below Glyptosauridae is 381 382 not possible (de Buffrénil et al., 2010; Estes, 1983; Gauthier et al., 2012; Rage, 1978; Sullivan, 1979, 2019). Although differences in general shape and absence/presence of the 383 overlap surface clearly exist among these osteoderms, their positions on the body plays more 384 385 likely a major role in this case. Although it seems to be unlikely that these osteoderms belong to a form for which no other elements have not been recorded in the locality, we do not 386 necessarily associate them with the frontals described above and identify them only as 387 Glyptosauridae indet. The hexagonal/polygonal osteoderms strongly suggest that they are 388 389 cranial. In the case of their cranial placement, they belong to Glyptosaurinae. However, the 390 fact that they may pertain to the cheek region of a "melanosaurine" cannot be ruled out. The rectangular body (trunk) osteoderms could also be equally be attributed to a "melanosaurine." 391 392 393 Anguioidea indet. (Fig. 6K-M) 394 395 Referred Material —One isolated osteoderm UM-COS-1008 396 397

398 **Description**

Osteoderm— UM-COS-1008, a single isolated rectangular osteoderm (Fig. 6K-M) is 399 thin rather than thick (although slightly thicker than the body osteoderms described above), 400 with a low medial ridge (which is slightly more pronounced in the posterior section) 401 extending along the entire central region, being restricted to the sculptured region. The 402 anterior overlap surface is well defined. It occupies about one quarter of the external surface. 403 The lateral bevel is narrow and the lateral imbrication is weak but present. The posterior 404 portion of the external surface is weakly ornamented. The ornamentation is formed by pits 405 and hardly distinguishable ridges diverging from the central region – pits are transformed into 406 407 short grooves at the periphery. The internal surface is pierced by several foramina (at least two of which are visible). Besides these, few short grooves are ridges are located on the 408 periphery in the mid-section of the osteoderm, but only on one side. 409

410 Remarks—The osteoderm UM-COS 1008 appears to differ from osteoderms of Glyptosauridae described above by its shape, thickness, length of overlap surface, presence of 411 412 lateral bevel, and the apparent absence of typical discrete tubercles forming the external ornamentation. This specimen resembles osteoderms present in members of Anguinae 413 (Čerňanský & Klembara, 2017; Estes, 1983; Georgalis et al., 2019; Hoffstetter, 1962; 414 415 Williams et al., 2022), provided that the absence of tuberclulate ornamentation is not caused by abrasion or corrosion. The osteoderm appears to be slightly water-worn. The fact that this 416 osteoderm is thicker might also support glyptosaurid affinity. 417

- 418
- 419 Varanoidea Gray, 1827 (sensu Estes et al., 1988).
 420 Palaeovaranidae Georgalis, 2017
 421 Palaeovaranidae indet.
 422 (Fig. 7)
 - 17

424

Referred Material — UM-COS-1009, nearly complete frontal.

425

426 **Description**

Frontal— The nearly complete frontal (Fig. 7) is an unpaired element, missing only 427 the anterior end and the left posterolateral corner. In the posterior section, the frontal bears 428 traces of the original midline suture on both ventral and dorsal surfaces (Fig. 7A, B). It is 429 narrow anteroposteriorly and flares outward posterolaterally on the right side. The frontal is 430 small, measuring only 4.6 mm along the mid-line, but was slightly longer in life as the 431 432 anterior portion is broken off. The lateral margin of the frontal is concave, with the narrowest part anterior to the mid-orbit region. Although the anterior portion is missing, the frontal 433 widens anteriorly, based on the lateral flaring of the crista cranii frontalis on the ventral side 434 435 (Fig. 7B).

The dorsal surface of the frontal is ornamented with irregular small, distinct mounds. Their structures are complex, having a shrunken (or rumpled) appearance with several longitudinal crests, whereas others are pointed, forming several small spike-like structures (they are of high relief). The posterior margin, which contacts the parietal, is anteriorly concave, giving it a slight irregular or wavy appearance in dorsal view (Fig. 7A, B).

On the ventral side, the frontal cranial crests are well-developed, forming a prominent orbital rim. The cranial crests are slightly deeper ventrally in the anterior region where the ventral margins are distinct. The crests extend to the posterolateral end of the frontal where they are slightly wider and blunt. They are pierced by three foramina on each side. The left lateral side preserves a wedge-shaped facet for the prefrontal at the anterior end (Fig. 7C). The right lateral side a bears a distinct facet at the posterior for the reception of the postfrontal (Fig. 7D). These articular surfaces are separated by a large portion of the orbital margin.

448
449 Varanoidea indet.
450 (Fig. 8)
451
452 Referred Material—UM-COS-1010, greater portion of left maxilla.
453

454 **Description**

Maxilla— The nearly complete left maxilla is preserved in matrix, with the lateral 455 side visible (Fig. 8A). The anterior portion is broken. The preserved portion has a maximum 456 anteroposterior length of 10.2 mm and a maximum height of 4.6 mm, measured from the 457 ventral edge of the maxilla to the uppermost extent of the nasal process. The nasal process of 458 the maxilla is rather thin and has a slightly concave smooth surface with a few fractures along 459 the anterior half of the process due to crushing. The posterior margin of the nasal process 460 slopes posteroventrally at an angle of 28° where it ends near a small slit at the level of the last 461 preserved tooth. The external surface of the process is pierced by a series of four large 462 supralabial foramina along the external labial margin of the tooth row. In addition, there are 463 several smaller foramina located anterodorsally to the anteriormost preserved tooth. The 464 maxilla bears five complete teeth and has positions for four others. Posterior to the last labial 465 foramen, four tooth positions are present. The supradental shelf is relative thin, but widens 466 anteriorly (Fig. 8B). It protrudes medially, reaching its maximum at the level between the 467 fourth and fifth tooth position (counted from posterior) because of its contact with palatine. 468 The posterior opening of the superior alveolar canal is large and located at the level of the 469 470 sixth tooth position (counted from posterior). The medial surface of the preserved portion of nasal process of maxilla is smooth and no nasal crest on the dorsomedial surface can be 471 observed; however, the dorsal portion of the maxilla is fractured. 472

473	Dentition —The dentition is subpleurodont (sensu Hoffstetter 1954, 1955), the maxilla
474	parapet is low, and the bases of the teeth are attached to a sloping, concave surface. The teeth
475	are pointed, tall, and recurved distally and slightly lingually. The mesial and distal cutting
476	edges are sharp. Serrations appear to be absent. Based on the CT images, the tooth bases are
477	mesiodistally broad and bear well preserved typical basal striae (i.e., plicidentine; Fig. 8F),
478	which is typical for both Varanidae and Palaeovaranidae (Kearney & Rieppel, 2006;
479	Georgalis & Scheyer, 2019).
480	
481	DISCUSSION
482	
483	The Cos locality is as a unique window into the early Eocene "greenhouse world" in
484	Europe, providing a glimpse into the paleobiodiversity of anguimorphs during this geological
485	time. The anguimorphs from this locality include a new glyptosaurine, an indeterminate
486	anguioid (potentially an anguine but its glyptosaurid affinity cannot be ruled out), and
487	varanoids. Although the Cos lizard fauna is important for our understanding of the Eocene
488	ecosystems, many aspects can be fully resolved only by future systematic research and studies
489	of new material from Europe.
490	
491	Glyptosauridae
492	The tuberculated cranial material and osteoderms from Cos present an autapomorphy of
493	glyptosaurids, i.g., tuberculate ornamentation (Camp, 1923; Gilmore, 1928; Estes, 1983;
494	Gauthier et al., 2012; Sullivan, 1979, 2019). Thus, this material clearly belongs to this clade. In
495	Glyptosaurinae (sensu this paper), subhexagonal (or polygonal) osteoderms are present on the
496	skull whereas they have rectangular osteoderms on the most of the dorsal and ventral portions

body (Sullivan, 1979). Members of the paraphyletic "Melanosaurinae" (sensu this paper) are

characterized by irregular plate-like osteoderms that cover the skull roof (Sullivan, 1979, 2019).
Thus, isolated glyptosaurine frontals (and parietals) can be identified based on the presence of
hexagonal/polygonal tuberculate osteoderms covering the skull roof. For this reason, the Cos
frontal specimens clearly represents a member of Glyptosaurinae.

Until now the only glyptosaurine known from Europe has been Placosaurus from the 502 Eocene (Depéret, 1917; Gervais, 1848-52; Sullivan, 1979; Sullivan & Augé, 2006). Today, it 503 comprises four species: Placosaurus rugosus Gervais, 1848–1852, Placosaurus estesi Sullivan 504 & Augé, 2006, "Placosaurus" europaeus (Filhol, 1876; based on a dentary with teeth, so its 505 generic assignment is equivocal; Sullivan & Augé, 2006), and ?Placosaurus ragei Sullivan et 506 507 al., 2012 (based on a dentary and a referred parietal). Most finds are from the late Eocene 508 (except for ?P. ragei), while some are also known from the late middle Eocene (Lavergne and Le Bretou, Quercy; Rage 1988 and Augé 2005, Sullivan & Augé, 2006). Placosaurus can be 509 510 distinguished from the North American glyptosaurine genera Glyptosaurus, Paraglyptosaurus, Proglyptosaurus, and Helodermoides by the following combination of features on frontal 511 (Sullivan & Augé, 2006): (1) frontals are fused and (2) slightly arched; and (3) cephalic 512 osteoderms usually form one or two complete rows of hexagonal osteoderms over each orbit. 513 514 However, the Cos frontals have a unique distribution of skull ostoderms and therefore cannot 515 be referred to this taxon or any of the other taxa (Fig. 4).

The early Eocene glyptosaurines compared to post-early Eocene glyptosaurines in Europe—All glyptosaurine frontals from Cos described here most likely belong to the same taxon as they have the same flat osteoderm morphology and display similar distribution pattern on the frontal where they correspond and are similar size. Moreover, they are comparable in size and come from the same locality. These frontals are markedly different from *Placosaurus* (Sullivan & Augé, 2006; see Diagnosis and descriptions here). For this reason, we proposed a new genus and species for the Cos taxon, *Sullivania gallica*.

Sullivania cannot be compared with the stratigraphically older ?*Placosaurus ragei*,
from the earliest Eocene (MP 7) from Dormaal, because it is based on an isolated dentary and
a parietal. For this reason, Sullivan et al. (2012) questionably assigned it to *Placosaurus*. New
glyptosaurine lizard material from Dormaal, including a frontal that is different from both *Placosaurus* and *Sullivania*, is currently under study of one of us (AČ).

The frontals described by Rage & Augé (2003:fig. 3A) from Silveirinha (~MP 7, 528 Portugal) as indeterminate "Glyptosaurini" (Glyptosaurinae sensu this paper) are also 529 different from Sullivania in the following features: (1) the dorsal surface is covered with 530 hexagonal osteoderms; (2) one osteoderm is located in the anterior central region, whereas 531 532 posterior large ones are arranged in two rows; and (3) the frontal is robust and fused, although it is much smaller than the specimens from Cos (the minimum width of the paratypic right 533 frontal of Sullivania is 5.9 mm, whereas the width of the fused paired frontals from 534 535 Silveirinha is 4.5 mm).

European Sullivania compared to North American Gaultia—The clade 536 Glyptosauridae first appears in the early Eocene of Europe (Rage, 2013; Sullivan, 2019). The 537 earliest European record of glyptosaurid lizards is from MP 7 at Silveirinha and Dormaal 538 539 (Rage and Augé, 2003; Sullivan, 2019; Sullivan et al., 2012). Thus, the presence of 540 glyptosaurids is another taxon between Europe and North America during the Eocene (e.g., the lizard Saniwa: Augé et al., 2022; the questionable iguanian Tinosaurus: Čerňanský et al., 541 2023; the turtle Axestemys: Georgalis & Joyce, 2017; many other non-congeneric but related 542 taxa on both continents, notably charinaine snakes; Smith & Scanferla, 2021). These imply 543 faunal exchange between North America and Europe during the late Paleocene to early 544 Eocene. Several episodes of faunal exchange occurred between Europe and North America 545 before and after the Palaeocene-Eocene Thermal Maximum PETM, based on mammals 546 (DeBast & Smith, 2017; Smith & Smith, 2013; Solé et al., 2016), but known coeval Asian 547

fossil lizard record is too poor to draw any definite conclusions regarding faunal exchanges.
Interestingly, the frontal of *Sullivania gallica* from France resembles *Gaultia silvaticus* from
the earliest Eocene (biozone Wa-0, Willwood Formation) of Wyoming (Smith, 2009).

The holotype of Gaultia silvaticus is an incomplete fused frontal (although no 551 evidence of a suture is apparent on the ventral surface and most of the right half of the bone is 552 broken away, Smith, 2009: fig. 18D, Fig. 4B) diagnosed by: (1) apically flat osteodermal 553 shields covering the frontal and (2) different division of osteodermal cover relative to the 554 typical hexagonal one in other glyptosaurines. However, several differences are notable, 555 especially in the distribution of osteodermal shields and their shapes; in the anterior portion, 556 557 osteodermal shields are wide in *Gaultia*, mediolaterally elongated and chevron-shaped. They form several lines, with two prominent rows of osteoderms extending along the mediolateral 558 width of the frontal. In contrast, most frontal osteoderms in Sullivania are anteroposteriorly 559 560 elongate rather than wide and, moreover, there is the presence of leaf-shaped first central osteoderm. However, the second lateral left osteoderm crosses over the entire mediolateral 561 width of the frontal in Sullivania, having contact with the right one along the mid-line, similar 562 to that seen in Gaultia. The overall shape of these lateral osteoderms is different compared to 563 the shevron-shaped osteoderms in Gaultia; posteriorly, the osteodermal shields in Gaultia are 564 565 divided into convex polygons that are smaller than the chevrons. However, there is no indication of the presence of anteroposteriorly elongated, roughly trapezoidal osteoderms in 566 the posterior region of both paratype frontals of Sullivania. 567

A close relationship between *Gaultia* and *Sullivania* than to *Placosaurus* and *Glyptosaurus* cannot be ruled out. Assuming that they are closely related, *Sullivania* seems to retain some characteristics from its North American ancestor. It worth noting that half of the mammal taxa from the early Eocene of North America are closely related to the mammalian taxa known from Dormaal (Gingerich & Smith, 2006). The breakup of plate-like osteoderms

into smaller polygonal/hexagonal osteoderms on the skull roof is likely a derived feature 573 574 among glyptosaurines, in contrast to plesiomorphic type of the large dermal plates covering the skull roof (Sullivan, 1979). The Late Cretaceous Odaxosaurus piger retains the plate-like 575 osteoderm pattern seen in many anguids (Meszoely, 1970; Klembara et al., 2019) and "proto-576 glyptosaurines" (Sullivan, 2019). The fragmentation of cephalic osteoderms seen in the early 577 Eocene Sullivania and Gaultia may represent an intermediate stage. Gaultia, like Sullivania, 578 579 has its frontal roof armor divided into heterogenous plates as in Arpadosaurus (see Meszoely, 1970), not into discrete subequal hexagonal osteoderms as in *Glyptosaurus*, 580 Paraglyptosaurus, Placosaurus, and other taxa (Sullivan, 1979; Sullivan & Augé, 2006). 581 582 Sullivan (2019:754) considered Gaultia a sister taxon to Glyptosaurinae (sensu this paper, "Glyptosaurini" sensu Sullivan, 1979). Ultimately, it is a question of how Glyptosaurinae is 583 circumscribed. At present, no phylogenetic definition of Glyptosaurinae has been put forward. 584 585 There is no apparent disagreement about the intermediate morphology of Gaultia and its close relationship with Glyptosaurinae as circumscribed by Sullivan (1979). Sullivan (1979:15) 586 587 gave "hexagonal osteoderms that cover the entire skull" as a defining feature, and since such osteoderms are only present on the parietal, not the frontal, of Gaultia, it would support 588 589 Sullivan's (1979) position. However, Gaultia clearly does not fit the definition of "Melanosaurinae" (sensu this paper, "Melanosaurini" sensu Sullivan, 1979) either. For this 590 reason, given the close relationship between Gaultia and accepted members of 591 Glyptosaurinae, we continue to follow Smith's 2009 assignment ("Glyptosaurini" in Smith, 592 593 2009). We envision a time when a robust phylogenetic hypothesis of glyptosaurine relationships exists and clade names can be redefined on that basis. 594 Hypothetically, this might raise a question whether Sullivania is referable to 595 Glyptosaurinae or is a sister taxon to this clade. Again, however, the phylogenetic topology of 596 these taxa within Glyptosauridae can be resolved only based on phylogenetic analyses. For 597

now, we decided to tentatively assign Sullivania to Glyptosaurinae. The phylogenetic 598 599 relationships of Sullivania and Placosaurus are unknown. Taking account that the taxonomy is based on the frontal bone, Stenoplacosaurus from presumably the late-middle Eocene of 600 Mongolia is morphologically more similar to *Placosaurus* (previously, it was even placed to 601 Placosaurus [Sullivan & Augé, 2006] and later placed to its own genus [Sullivan & Dong, 602 2018]) than to Sullivania gen. nov. Thus, an Asian origin of Placosaurus cannot be excluded. 603 Owing to an incomplete glyptosaurine fossil record between the early and late Eocene in 604 Europe, their evolution and distribution patterns between these distant regions remain largely 605 unresolved. 606

607

608 Varanoidea

UM-COS 1009 represents a "miniature" palaeovaranid. It differs from currently 609 known genera by the following combination of features: (1) overall small size; the frontals of 610 Paranecrosaurus feisti (Stritzke, 1983), which is known only from the early-middle Eocene 611 612 of Messel in Germany, are much larger (18.2 mm in the holotype and even 30.8 in the paratype; see Smith & Habersetzer, 2021); (2) the non-paired (fused) narrow frontal, as in 613 Palaeovaranus Zittel, 1887–1890 (Georgalis et al., 2021) and Paranecrosaurus feisti 614 (Stritzke, 1983) (Smith & Habersetzer, 2021), unlike the paired frontals in Eosaniwa 615 Haubold, 1977 (Rieppel et al., 2007); (3) the type of an ornamentation. Its ornamentation, 616 which consists of small, densely arranged mounds, slightly resembles that of Palaeovaranus 617 lismonimenos Georgalis, Čerňanský & Klembara, 2021 (Georgalis et al., 2021). 618 619 Palaeovaranus lismonimenos was previously reported from the late Eocene (~ MP 17, see 620 Georgalis et al., 2021), whereas the Cos frontal is much older. However, the detailed surface texture is different - the ornamentation of the Cos specimen is more complex, having also 621 several spikes forming each mound (rather than possessing more-or-less simple mounds). The 622

ornamentation of UM-COS 1009 differs from the distinct sculpturing pattern observed on the 623 frontal of the holotype of Melanosauroides giganteus Kuhn, 1940 from Geiseltal. The dorsal 624 surface of the frontal of *M. giganteus* possesses broad, apically flat, rugose patches that 625 appear to be fused broad (not distinctly keeled) osteoderms (Georgalis, 2017; Smith & 626 Habersetzer, 2021). The dorsal surface of the frontal of Paranecrosaurus Smith & 627 Habersetzer, 2021 frontal is also different (Smith & Habersetzer, 2021); and (4) the long 628 posterolateral process relative to the the overall size of the frontal, as Paranecrosaurus (Smith 629 630 & Habersetzer, 2021), and unlike the short process in *Palaeovaranus* (Georgalis et al., 2021) and Eosaniwa (Rieppel et al., 2007). 631 632 UM-COS 1009 represents most likely a new taxon, but the taxonomy of palaeovaranids is now mainly based on the parietal morphology (Georgalis et al., 2021). For 633 this reason, we identified this frontal only as Palaeovaranidae indet. 634 It is unclear whether the maxilla (UM-COS 1010) and the frontal (UM-COS 1009) 635 represent a single taxon, although they are comparable in size and come from the same 636 locality. The overall morphology and plicidentine support allocation of the maxilla to a 637 varanoid (Kearney and Rieppel, 2006; Georgalis & Scheyer, 2019). It cannot be referred to 638 639 Shinisauridae, a group of anguimorph presence that is well-documente from the Eocene of 640 Europe (Smith, 2017; Smith et al., 2018). The maxilla has several teeth posterior to the last labial foramen where at least four tooth positions are preserved. This condition resembles 641 Saniwa orsmaelensis Dollo, 1923, which is known from Dormaal (Augé et al., 2022). Indeed, 642 643 the teeth and the concave lateral surface (the posterior portion of the maxilla is slightly bent laterally) of the Cos specimen resemble the condition in this taxon (Augé et al., 2022: fig. 2). 644 645 The geological age of the maxilla would also be consistent with the presence of Saniwa in Cos. However, several tooth positions posterior to the last labial foramen are also present in 646 Paranecrosaurus feisti (Smith & Habersetzer, 2021). The diagnostic parts of palaeovaranid 647

maxillae, however, are mainly found in medial view. The presence of a distinctly developed 648 nasal crest on the dorsomedial surface of the nasal process is a distinguishing feature of 649 Palaeovaranus (Georgalis et al., 2021) and Paranecrosaurus (nasolacrimal ridge sensu Smith 650 & Habersetzer, 2021). Unfortunately, this portion in the Cos maxilla is heavily damaged and 651 there is no indication of its presence. The teeth and the overall shape of the Cos specimen are 652 also similar to Melanosauroides Kuhn, 1940 (see Georgalis, 2017: fig. 4B). For all these 653 reasons, caution is needed and we decided to identify this maxilla only as Varanoidea indet. 654 (although we cannot be sure if palaeovaranids are indeed varanoids). More complete material 655 from Cos will shed light on the diversity of the varanoids here. 656

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- 658

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671

672 AVAILABILITY OF MATERIALS AND DATA

- All specimens are catalogued and accessible in the fossil reptile collection of the University of
- 674 Montpellier in France. Digital surface models of the figured fossil specimens are available on
- 675 Morphosource and Virtual Collections:
- 676 UM-COS-1001: https://www.morphosource.org/concern/parent/000514175/media/000514178
- 677 UM-COS-1000: https://www.morphosource.org/concern/media/000514185?locale=en
- 678 UM-COS-1003: <u>https://www.morphosource.org/concern/media/000514194?locale=en</u>
- 679 UM-COS-1002: https://www.morphosource.org/concern/media/000514200?locale=en
- 680 UM-COS-1004: <u>https://www.morphosource.org/concern/media/000514206?locale=en</u>
- 681 UM-COS-1005: <u>https://www.morphosource.org/concern/media/000514212?locale=en</u>
- 682 UM-COS-1006: <u>https://www.morphosource.org/concern/media/000514218?locale=en</u>
- 683 UM-COS-1007: <u>https://www.morphosource.org/concern/media/000514224?locale=en</u>
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- 685 UM-COS-1009: https://www.morphosource.org/concern/media/000514236?locale=en
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989 **Figure legend**

990

FIGURE 1. Location of the Cos locality in France. [planned for 2/3 of full-page width]992

993	FIGURE 2. <i>Sullivania gallica</i> gen. et sp. nov. from the early Eocene Cos locality of France.
994	UM-COS-1001 (holotype) anterior portion of fused frontals: in A, dorsal; B, outline drawing
995	showing the armour division pattern; C, ventral; and D, lateral views (all micro-CT
996	visualizations). E, axial section at the mid-level of the dorsoventral thickness; and F, coronal
997	section at the level of the frontal cranial crests. [planned for page width]
998	
999	FIGURE 3. Sullivania gallica gen. et sp. nov. from the early Eocene Cos locality of France.
1000	UM-COS-1000 (paratype), nearly complete right frontal: in A, dorsal; B, photograph with
1001	line drawing indicating the interpretation of osteoderm distribution; C, ventral; and D, lateral
1002	views (micro-CT visualizations). E, Axial section at the mid-level of the dorsoventral
1003	thickness; and F, coronal section at the level of the frontal cranial crests. UM-COS-1003
1004	(paratype), posterior portion of right frontal: in G, dorsal; and H, ventral views (all micro-CT
1005	visualizations). [planned for page width]

1006

1007 FIGURE 4. The reconstruction of complete frontal of *Sullivania gallica* gen. et sp. nov.

1008 comparing it to frontals of *Gaultia silvaticus*, from the earliest Eocene of Wyoming, USA;

- 1009 Placosaurus estesi and P. rugosus, from the middle and late Eocene of France; and
- 1010 Stenoplacosaurus mongoliensis from the late-middle Eocene of Mongolia (all are dorsal
- 1011 views, except G, which is left lateral view). A, Sullivania gallica, reconstruction based on

- 1012 UM-COS-1000 and 01; **B**, *Gaultia silvaticus* UCMP 216000, (holotype) middle portion of
- 1013 frontal and C, UCMP 150966 (paratype) partial left frontal (modified from Smith, 2009); D,
- 1014 Placosaurus estesi (MNHN QU-17735, holotype) nearly complete frontal; E, P. rugosus
- 1015 (MNHN 1906-25, holotype) nearly complete frontal and left prefrontal; and F, G,
- 1016 Stenoplacosaurus mongoliensis (AMNH 6669, holotype), left frontal (modified from Sullivan
- 1017 & Dong, 2018). The arrow in *Gaultia* (**B**) indicates the position of the mid-line between the
- 1018 left and right frontals (see Smith, 2009). The dotted line in *P. rugosus* (E) indicates the frontal
- 1019 prefrontal contact. [planned for page width]

- 1021 FIGURE 5. Glyptosauridae indeterminate from the early Eocene Cos locality of France. UM-
- 1022 COS-1002 left maxilla. A, left lateral view; B, medial (lingual) view; C, occlusal (ventral)
- 1023 view; and **D**, dorsal view. **A-D** are micro-CT visualizations; UM-COS-1011, right maxilla. **E**,
- 1024 right lateral view, F, medial (lingual) view, G, occlusal (ventral) view (with close-up of teeth
- in ventromedial view); and **H**, dorsal view. [planned for page width]

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- 1027 FIGURE 6. Glyptosauridae indeterminate (A-J) and Anguioidea indet. (K-M) from the early
- 1028 Eocene Cos locality of France. Isolated cephalic osteoderms. UM-COS-1004, hexagonal
- 1029 osteoderm: A, dorsal; B, ventral; and C, lateral views. UM-COS-1005, polygonal osteoderm:
- 1030 D, external and E, internal views. Isolated body osteoderms. UM-COS-1006: F, external; G,
- 1031 internal and H, lateral views. UM-COS-1007: I, external; and J, internal views. UM-COS-
- 1032 1008; K, external; L, internal; and M, oblique lateral views. All images are micro-CT
- 1033 visualizations. [planned for page width]

- 1035 FIGURE 7. Palaeovaranidae indeterminate from the early Eocene Cos locality of France.
- 1036 UM-COS- 1009, nearly complete frontal. A, dorsal; B, ventral; C, left lateral; D, right lateral;
- and E, anterodorsal views. All images are micro-CT visualizations. [planned for page width]

- 1039 FIGURE 8. Varanoidea indeterminate from the early Eocene Cos locality of France. UM-
- 1040 COS-1010, left maxilla. A, lateral; B, medial; C, dorsomedial; D, dorsal; and E, ventral views
- 1041 (A-E are micro-CT visualizations); and \mathbf{F} , μ CT slice of tooth bases showing the presence of
- 1042 plicidentine. [planned for page width]
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