# The youngest ellimmichthyiform (clupeomorph) fish to date from the Oligocene of South China (#56381)

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# The youngest ellimmichthyiform (clupeomorph) fish to date from the Oligocene of South China

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A new ellimmichthyform, *Guiclupea superstes*, gen. et sp. nov., from the Oligocene Ningming Formation of Ningming Basin, Guangxi Zhuang Autonomous Region, South China is described herein. With relatively large body size, parietals meeting at the midline, anterior ceratohyal with a beryciform foramen in the center, a complete predorsal scutes series of very high number and about equally-size scutes with radiating ridges on dorsal surface, first preural centrum unfused with first uroneural but fused with the parhypural, and first ural centrum of roughly the same size as the preural centrum, Guiclupea superstes cannot be assigned to the order Clupeiformes. The phylogenetic analyses using parsimony and Bayesian inference methods with *Chanos/Elops* as outgroup respectively suggests that the new form is closer to ellimmichthyiform genus *Diplomystus* than to any other fishes, although there are some discrepancies between the two criteria and different outgroup used. It shares with Diplomystus the high supraoccipital crest, pelvic-fin insertion in advance of dorsal fin origin, and number of predorsal scutes more than 20. The new form is the youngest ellimmichthyiform fish record in the world so far. Its discovery indicates that the members of the Ellimmichthyiformes had a wider distribution range and a longer evolutional history than previously known.

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11	*Corresponding author: cgengjiao@aliyun.com
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28 Keywords Ellimmichthyiforms, Oligocene, southern China, phylogeny, paleobiology

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#### Introduction

The Ellimmichthyiformes is one of the two major clades of the Clupeomorpha (Nelson et al., 30 2016). The Recent Clupeomorpha is represented only by the order Clupeiformes, which is 31 32 amongst the most economically important fish species for food (Zhang, 2001), and containing both fossil and extant herrings, anchovies, and other relatives. The order Ellimmichthyiformes is 33 an extinct cosmopolitan clade (Nelson et al., 2016), established by Grande in 1982. It initially 34 included only a single family Ellimmichthyidae (=Paraclupeidae, see Chang & Grande 1997) 35 with only two genera- Diplomystus and Ellimmichthys, diagnosed by bearing a series of 36 predorsal scutes expanding laterally then taking a subrectangular-shape, and laking some derived 37 characters of the Clupeiformes, e.g., presence of recessus lateralis, parietal bones completely 38 separated by the supraoccipital, and loss of the 'beryciform' foramen in anterior ceratohyal 39 40 (Grande, 1982, 1985). Since the founding of Ellimmichthyiformes, especially in the last two decades, many new and previously known genera and species have been added or moved to this 41 42 order (Silva Santos, 1990, 1994; Bannikov & Bacchia, 2000; Chang & Maisey, 2003; Poyato-43 Ariza & García-Garmilla, 2000; Forey et al., 2003; Hay et al., 2007; Alvarado-Ortega & Ovalles-Damián 2008; Khalloufi et al., 2010; Newbrey et al 2010; Murray & Wilson, 2011, Malabarba et 44 45 al., 2004; Bannikov, 2015; Vernygora & Murray, 2015; Murrey et al., 2016; Alvarado-Ortega & 46 Melgarejo-Damián. 2017; Plock et al., 2020; etc), though some of them with the predorsal scutes

47 pattern only partially agree with or completely disagree with this order-level character, e.g., Ellimma branneri, whose anterior predorsal scutes is longer than broad; Scutatospinosus 48 itapagipensis and Codichthys carnavalii, without any subrectangular predorsal scutes at all, 49 completely disagree with the order-level character. Along with the increasing of the members of 50 this order, several families were erected, and the interest in the definition, classification, and 51 52 intra-relationship of the group has been increasing (Bannikov & Bacchia, 2000; Chang & Maisey, 2003; Zaragüeta-Bagils, 2004; Alvarado-Ortega et al., 2008; Murray & Wilson, 2013; Figueiredo 53 & Ribeiro 2016; Vernygora et al., 2016; Marramà & Carnevale 2017; Boukhalfa et al., 2019; 54 Vernygora & Murray, in press; etc), though no definitive consensus has been reached on these 55 issues. The main differences among the results of previous studies are the relationship of 56 Armigatus and Diplomystus [some analysis suggested that Armigatus is sister to Diplomystus 57 (Chang & Maisey, 2003, Murray & Wilson, 2013; others suggested that *Armigatus* is in a more 58 basal (Forey, 2004; Figueiredo & Ribeiro 2016) or derived position than Diplomystus 59 60 (Vernygora & Murray, 2016; Marramà & Carnevale 2017; Boukhalfa et al., 2019), or Armigatus is not an ellimmichthyiform member (Zaragüeta-Bagils, 2004)], and Ornategulum is an 61 ellimmichthyiforms or not (Murrey & Wilson, 2013; Marrama et al., 2018; Figuereido & Ribeiro 62 63 2017; Boukhalfa et al., 2019). To date, reported members of the group have reached 21 genera and 37 species at least, ranging from the Early Cretaceous to middle Eocene marine and non-64 65 marine strata of Eastern Asia, Middle East, North and South America, Africa, and Europe 66 (Figure 1). No ellimmichthyiform fish from the strata younger than the Eocene has ever been reported. Accordingly, it was believed that the ellimmichthyiforms have finally become extinct 67 68 after the middle Eocene. However, recently, an Oligocene ellimmichthyiform fish was 69 discovered from the non-marine deposits of Ningming Basin, Guangxi Zhuang Autonomous

Region, South China. Consequently, this new ellimmichthyiform fish is the youngest one to date. This discovery not only extends the spatial and temporal distribution of ellimmichthyiform, but also shed new light for our understanding of the evolutionary history and paleobiogeography of the order. Herein we provide descriptions of the new form, perform a phylogenetic analysis of the Ellimmichthyiformes, and discuss the taxonomic position and the paleobiographical implications of the new form.

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#### Abbreviations

Institutional Abbreviation—IVPP, Institute of Vertebrate Paleontology and
Paleoanthropology, Chinese Academy of Sciences, Beijing, China. NHMG, Natural History
Museum of Guangxi Zhuang Autonomous Region, China.

Anatomical Abbreviations—aa, angulo-articular; ach, anterior ceratohyal; as, autosphenotic; 80 **br**, branchiostegal rays; **cl**, cleithrum; **cor**, coracoids; **cs**, caudal scute; **d**, dentary; **ect**, 81 ectopterygoid; en, epineural; enpt, entopterygoid; ep, epural; epl, epipleural; fr, frontal; h, 82 hypural; hy, hyomandibular; io, infraorbital; iop, interopercle; la, lacrimal; le, lateral ethmoid; 83 met, metapterygoid; ms, mesethmoid; msc, mandibular sensory canal; mx, maxilla; ns, neural 84 spine; op, opercle; pa, parietal; pch, posterior ceratohyal; pcl, postcleithra; pd, predorsal bones; 85 86 **ph**, parhypural; **pmx**, premaxilla; **pop**, preopercle; **pr**, pleural rib; **ps**, parapophysis; **pt**, posttemporal; pto, pterotic; pu, preural centrum; q, quadrate; ra, retroarticular; sc, scapula; scl, 87 supracleithrum; scr, sclerotic bone; so, supraorbital; sop, subopercle; smxa, anterior 88 89 supramaxilla; smxp, posterior supramaxilla; sp, sphenotic; spo, supraoccipital; sy, symplectic; u, ural centrum; **uh**, urohyal; **un**, uroneural; **vo**, vomer; **vs**, ventral scutes. 90

91

#### **Materials and Methods**

92 Fossil specimens except NHMG 038777, described herein, including articular skeletons and

93 detached bones, were collected from the outcrops about 2.5~3.5 km west of Ningming County, Guangxi, South China (Figure 1), about 40 km away from the boundary of China and Vietnam 94 and about 120 km northwest of the South China Sea, and now housed in the NHMG. The fossil-95 bearing strata positions in the middle-upper part of the Second Member of the Yongning Group 96 (Bureau of Geology and Mineral Resources of Guangxi Zhuang Autonomous Region, 1985), or 97 Ningming Formation (Li et al., 1995), which is a set of fossiliferous lacustrine sediments 98 dominated by light-gray, yellowish mudstones, occasionally containing fine sand grains. This 99 deposition also bears a variety of cyprinid and a few siluriform and perciform fishes, and a large 100 101 number of plant macrofossils (Chen et al., 2018). No volcanic material which can be used for isotopic dating was found at the locality. The geological age of the Ningming Formation, 102 according to palynologists (Wang et al., 2003), is Oligocene. Paleobotanists accepted this 103 opinion when studied macrofossil plant from the same strata (Li et al., 2003; Shi et al., 2010, 104 2012, 2014; Wang et al., 2014; Dong et al., 2017; Ma et al., 2017; etc). We applied this 105 geological age also when we studied *Huashancyprinus robustispinus* (Cyprinidae, Cypriniformes) 106 from the same locality and horizon (Chen & Chang, 2011) and adopt it herein. NHMG 038777 is 107 a disarticulated dentary collected from Yongning formation of Santang, Nanning basin, Guangxi. 108 109 The geological age of Yongning formation is Oligocene (Zhao, 1983, 1993; Quan et al., 2012) Fossil fish materials were prepared mechanically with steel needles of different sizes under a 110

binocular microscope. Line drawings were made based on the observations of the fossils under an Olympus SZ61 microscope and the photos.

The taxonomy terminology and the methods of counting and measurement used in here follow Grande (1982) and Forey et al. (2003). The descriptions of gill raker follow Bornbusch & Lee (1992). Specimens used for comparison include: 1) *Paraclupea chetungensis* Sun 1956,

including IVPP V816, V2986.2, V3002.1, 5-8, 10, 12, 15, 19, from the Lower Cretaceous
Chawan Formation, eastern China; 2) *Diplomystus shengliensis* Zhang et al. 1985, including
SOF 790001, SOF 790002, and SOF 790003, from top of series 4 to bottom of series 3 of the
Shahejie Formation, Middle Eocene, East China; 3) *Knightia bohaiensis* Zhang et al. 1985,
including SOF 790003, from top of series 4 to bottom of series 3 of the Shahejie Formation,
Middle Eocene, East China; and 4) dried skeleton and disarticular bones of *Ilisha elongate*,
NHMG, collected from Nanning Dancun Market.

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The Phylogenetic analyses are based on data matrix (see Appendix 1S-3S) consisted of 55 morphological characters and 40 taxa, including three Recent clupeiform species (*Denticep*, *clupeoides*, *Chirocentrus dorab*, and *Odaxothrissa vittata* (the first one is the only extant member of the Denticipitoidei; the latter two represent the Clupeoidei), a gonorynchiform or elopomorph (*Chanos chanos* or *Elops saurus*, being used as outgroup alternatively to polarize the characters and root the tree), the enigmatic fossil *Ornategulum sardinioides* Forey 1973, and

our new form (to test its position within the Clupeomorpha). Characters are mainly adopted from
Chang & Maisey (2003), and Murray & Wilson (2013).

The analyses use both parsimony and Bayesian inference methods, for both methods have 141 advantages and disadvantages for morphological data (Bai et al., 2020). The parsimony analyses 142 were performed with TNT 1.5 (Goloboff et al., 2008), using the Traditional Search method with 143 144 1000 replicates and tree bisection and reconnection (TBR) swapping algorithm. All characters are unordered and equally weighted. The most parsimonious trees (MPTs) generated by the 145 analysis were used to construct a strict consensus tree. Tree length, consistency index (CI), 146 147 retention index (RI), Bremer support and bootstrap values were then calculated for the strict consensus tree. 148

Bayesian analyses were conducted by MrBayes 3.1.2 (Ronquist & Huelsenbeck, 2003). For the substitution models, the Mkv model was used with an assumption of gamma rate variation across characters. Markov chain Monte Carlo analysis consists of four chains, which were run simultaneously with 2 000 000 trees, sampling 1/100 trees, with a burn-in value of 5000. The remaining trees were used to build a 50% majority rule consensus tree, and statistical support of each node was assessed by posterior probabilities.

155

156	Systematic Paleontology
157	Infraclass TELEOSTEI Müller, 1845
158	Cohort CLUPEOCEPHALA Patterson & Rosen, 1977
159	Superorder CLUPEOMORPHA Greenwood et al., 1966
160	Order ELLIMMICHTHYIFORMES Grande, 1982
161	Diplomystus clade

162	Genus GUICLUPEA gen. nov.
163	Diagnosis: A pretty large-sized, double-armoured ellimmichthyiform fish, differing from
164	other genera of the order with the following combination of characters: dorsal body margin
165	without marked angle at the dorsal fin insertion; posttemporal large; predorsal scutes series
166	complete, with scutes small, numerous (about 55), all about equal in size, and with ridges on
167	dorsal surface; number of predorsal bones ten or more; no diastema between second and third
168	hypural; proximal end of middle principal caudal fin ra
169	Etymology : 'gui', the abbreviation in Chinese of the Guangxi Zhuang Autonomous Region, a
170	province of China from where the fossil materials were collected; 'clupea', from the Latin, to
171	indicate clupeomorph affinities of the new taxon.
172	Type species : Guiclupea superstes gen. et. sp. nov.
173	
174	Guiclupea superstes gen. et. sp. nov.
175	(Figs. 2–7)
176	<b>Diagnosis</b> : See generic diagnosis. Pectoral fin rays 18, pelvic fin rays 5~6, dorsal fin rays 14,
177	anal fin rays 38, total number of vertebrae (include ural centra) about 46.
178	Etymology: 'superstes', Latin 'survivore'. The species name means that the species survived
179	in Oligocene when almost all menbers of the order Elijimmchthyiforms had become extinct.
180	Holotype: NHMG 005532, a nearly complete skeleton, part and counterpart (Figs. 2A-B).
181	Paratypes: NHMG 033659 (Fig. 3A), a relatively complete skeleton with the snout and the
182	caudal fin rays missing; NHMG 033658 (Fig. 6), a skeleton from the anterior margin of the
183	orbital to the caudal peduncle, part of the anterior portion and counterpart of the posterior portion;
184	NHMG 011648 (Fig. 7A), caudal peduncle to caudal fin, part and counterpart.

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185 Additional material: NHMG 033660 (Fig. 5), disarticulated bones of skull and anterior part of the body; NHMG 004929, an incomplete skeleton with the posterior part of the body missing; 186 NHMG 033361, disarticulated bones of skull, the anterior ceratohyal and entopterygoid in this 187 specimen were showed in Fig. 4E; NHMG 033680, disarticulated bones of skull and anterior part 188 the posterior ceratohyals in this specimen were showed in Fig. 4F; NHMG 033685, 189 of the body. 190 premaxilla (Fig. 4C); NHMG 038778, an incomplete skeleton with the head and caudal skeleton and fin missing; NHMG 033681-033683 and NHMG 038777, dentary (NHMG 033681 showed 191 in Fig. 3D); NHMG 011647 and NHMG 011649, caudal skeleton and caudal fin; NHMG 192 193 011650-011651, caudal skeleton.

Localities and horizon: Gaoling Village (22°07N, 107°02′E), Ningming County, and Santang (22°52′N, 108°25′E), Nanning, Guangxi province, China; middle to upper portion of the second member of the Yongning Group or Ningming Formation and Yongning Formation; Oligocene.

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#### Description

(1) General Appearance. This new form is a pretty large-sized double-armored clupeomorph. 199 The total length of the holotype (Figs. 2A-B) is about 526 mm, and the known largest incomplete 200 201 specimen, NHMG 033659 (Fig. 3A), with the preserved portion reaches 638 mm in length, and the distance from the anterior margin of the lacrimal to the caudal fin base is about 585 mm. The 202 fish has an elongate fusiform body. The standard length in holotype is about 3.3 times of203 204 maximum body depth. The anterior dorsal margin of the fish body is rounded and convex, without a marked angle at the origin of the dorsal fin as the typical members of paraclupeids. The 205 206 ventral border in front of the insertion of the pelvic fin is also convex, but is straight and 207 obliquely rising upwards behind the insertion, making the posterior part of the body gradually

208 narrowing caudally. The origin of the dorsal fin is posterior to the level of the insertion of the 209 pelvic. The anal fin has a long base. The caudal fin is deeply forked. There are complete series of 210 predorsal and ventral scutes along the dorsal and ventral margin of the body respectively. The 211 meristic characters are listed in Table 1.

(2) **Skull Roof**. The head is slightly longer than deep (Figs. 2A; 4A). The skull roof above the 212 213 eye is narrow. The frontal is a long bone, with its posterior one-fourth expanding laterally. There is a longitudinal ridge of the supraorbital sensory canal running along the dorsal surface of this 214 bone (Figs. 4A-B; 5). The frontal sutures with the anterior edge of the parietal posteriorly. Two 215 parietal bones meet at the midline, at least in the anterior part (Fig. 4A-B), as in the case of the 216 primitive clupeomorphs (Grande, 1985). No fontanelle between the anterior portions of the 217 frontals is observed, which is often present in clupeoids and *Paraclupea chetungensis* (Chang 218 and Grande, 1997). Anterior to the frontal is the mesethmoid, which bears a lateral process on 219 each side (Fig. 5). Lateral ethmoid, contacting the frontal at its anteriolateral margin, is situated 220 anterior to the orbit, forming the lateral portion of the anterior wall of the orbit (Figs. 4A-B). The 221 outlines of pterotic and sphenotic are not clear. But the strong ventrally directed process of the 222 autosphenotic can be seen, lying in front of the head of the hyomandibula (Fig. 4A-B). The 223 224 posttemporal is a very large bone (Figs. 4A-B; NHMG 004929). Supraoccipital is situated posteriorly and sutures with the parietals anteriorly. Supraoccipital crest is well-developed, being 225 226 high and triangle-shaped, making the lateral profile of the skull roof with a distinct angle 227 between the anterior and the posterior parts (Fig. 4A-B). The external surface of all the skull roof bones lacks ornamentation, except for a longitudinal ridge containing the supraorbital sensory 228 229 canal (Figs. 2A-B, 4A, 5, 6A). No openings of the recessus lateralis are observed.

230 (3) **Opercular Series and Cheek Bones**. The opercle is trapezoidal in shape. Its ventral part is

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wider than the dorsal part, with the anteroventral corner protrudes downward and a little bit 231 forward. The depth of the opercle is about 1.5 times of its width. No ornamentation on the 232 surface of the opercle can be observed (Figs. 4A, 6). Two arms of the preopercle, with the dorsal 233 branch slightly longer than the ventral branch, form an obtuse angle. The preopercular sensory 234 canal runs along the mid-line of the bone sending out several branches backwards and 235 236 downwards. Interopercle and subopercle are long and thin, with smooth surface (Figs. 4A-B, 6A). About 8-9 branchiostegal rays can be detected in the holotype, although the outline of each ray is 237 not very clear (Fig. 2A). In NHMG 033659, five of the posterior branchiostegal rays of the right 238 239 side can be counted below the interopercular bone, while four displaced, slender anterior branchiostegal rays are discernible in the position anterior to the interopercle (Fig. 4A-B). 240

(4) Circumorbital Bones. There is an arched, long bone above the frontal bone in NHMG 033659 (Fig. 4A-B), we guess that it ought to be the supraorbital bone displaced from its original position. The sclerotic ring, consisting of two halves, can be observed in the posterior and anterior part of the orbit. Anterior to the orbit, there are two bones, which seem to bear sensory canals. The large, sub-trianglular, anterior thin bone is the lacrimal, whereas the posterior rectangle one may be the infraorbital 2 (Fig. 4A-B). Detached infraorbital bones are preserved in NHMG 033660 (Fig. 5).

(5) **Jaws and Palate**. The mouth is somewhat superterminal based on NHMG 004929. The oblique gap is relatively short, with the lower jaw articulation under the anterior border of the orbit (Fig. 4A; NHMG 004929). The upper jaw consists of a premaxilla, a maxilla and two supramaxillae. The premaxilla is a small, long, triangular bone, with a row of small conical teeth on its oral margin (Fig. 4C). The maxilla is a long bone, with its anterior part narrow and thick while its posterior part broadening into a thin blade and bearing a rounded ventral profile. The

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oral margin of the thin blade is finely serrated. The anterior end of the maxilla develops into a round ethmoid head and a round palatine head (Fig. 5, NHMG 004929, 033684, 033686, 033689). Along the dorsal edge of the maxilla two supramaxillae can be detected in NHMG 004929 and 033659 (Fig. 4A). In NHMG 033689, two disarticulated supramaxillae bones seem to be roughly equal in size and similar in shape, and their external surfaces are smooth, except for a low ridge extending along its midline.

The dentary has a well-developed coronoid process. In NHMG 033681-3 and NHMG 038777, 260 there is a single row of small conical teeth along the short tooth oral margin of the dentary (Fig. 261 4D-E). Teeth close to the symphysis of the two dentaries are slightly stouter than that in the rear<sub>1</sub> 262 But in specimen NHMG 033660, no teeth can be seen on the oral margin of the dentary, which 263 are probably missing during preparation or fossilization. Along the lower lateral margin of the 264 dentary, the mandibular sensory canal is well developed with  $6 \sim 7$  pores (Fig. 5). The angulo-265 articular is a triangular bone with mandibular sensory canal running along it lower margin of 266 267 lateral surface. Length of angulo-articular is about half of that of the dentary. Its posterior end forms the articular process for the quadrate (Figs. 4A-B, 5). The very small retroarticular bone is 268 located below the postarticular head of the angulo-articular bone (Fig. 4A-B). 269

The parasphenoid can be partly observed in NHMG 033658 and 004929. It is difficult to judge if a basipterygoid process and the "osteoglossid" tooth patch of the bone is present or not due to the preservation. The quadrate, as is generally for teleosts, consists of a fan-like plate at its dorsal side and a rod-like posterior process at its ventral side. The articular head of the quadrate fits into the socket at the postero-dorsal end of the angulo-articular (Fig. 4A-B).

(6) Hyoid Arch. The area for hyomandibular articulation with the braincase is broad. The
hyomandibular bears a thin, broad anterio-dorsal plate and a long, narrow ventral shaft, which

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ventrally connects the upper end of the symplectic (Figs. 4A-B, 5). The condyle for articulation 277 with the opercle is large. From that level, a prominent ridge runs ventrally along the posterior 278 margin of the inner surface of the shaft. The foramen for hyomandibular branch of facial never 279 (VII.hm) is clear. The symplectic inserts the notch between the plate of the quadrate and its 280 ventral process with its anterior end to complete the suspension of the lower jaw to the braincase 281 282 (Figs. 4A-B). Detached entopterygoid is shown in NHMG 033660 and 033661. It is a broad, triangular bone, with numerous fine conical teeth covering its buccal side (Figs. 4F, 5). The 283 metapterygoid is an expansive, trapezoid bone. Its anterior margin is posterior to the anterior 284 margin of the quadrate, and its posterior margin reaches a relatively more dorsal position, almost 285 in line with the hyomandibular condyle (Figs. 4A-B, 6A). 286

The anterior ceratohyal is a thick, sub-rectangular plate with its length about two times  $\mathbf{\Theta}$  its 287 depth. Its dorsal margin is slightly convex, whereas its ventral margin is slightly concave. Its 288 central part is pierced by a large elongated oval foramen (Figs. 4F, 5) as in primitive 289 clupeomorphs (Grande, 1985). The posterior ceratohyal is a triangle plate without foramen 290 within it. There is a small notch on its dorsal-posterior margin (Fig. 4G). The urohyal shows a 291 narrow ventral keel and a vertical crest. Height of the crest gradually increases posteriorly (Fig. 292 293 5). Gill arches are not well-preserved, but many dislocated, long and pointed gill rakers with a bifid base which embraced the gill arches are observed in several specimens (Figs. 3A, 4F-G, 5). 294 295 Length of the gill rakers varies from about one vertebral centrum to  $2 \sim 3$  time as long as a 296 vertebral centrum or even more. There are numerous fine conical denticles recurved posteriorly throughout almost the upper edge of the gill rakers, differing from the situation in *Diplomystus* 297 298 sp. from English chalk in which the rakers appear to be smooth throughout most of their length 299 (Forey, 2004)

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300 (7) Paired Fins and Girdles. Supracleithrum is small, lying below and posterior to the very developed posttemporal. The cleithrum is a long and S-shaped bone, with its upper end covered 301 by the supracleithrum. Below the cleithrum is the developed, laminate coracoid with a large 302 notch on its anterior margin (Figs. 4A, 5-6). The pectoral fin is located rather high in the flank. 303 The fin is long, extending over the insertion of the pelvic fin in NHMG 033658 (Fig. 6A). In 304 305 other specimens, the fin rays do not look so long, probably because of the distal ends of the fin rays were missing during the process of fossilization. Eighteen pectoral fin rays can be counted 306 307 (Fig. 6A).

The pelvic girdle cannot be observed due to the covering of the abdominal scutes. The pelvic fin is small, with about 5~6 fin rays, inserted at the level in front of the origin of the dorsal fin (Figs. 2A, 6A). The length of the longest pelvic fin ray is equivalent to the span of  $6\sim7$ postpelvic scutes.

(8) Dorsal and Anal Fins. The origin of the dorsal fin is situated slightly posterior to the midpoint of standard body length. There are about 14 dorsal fin rays (Figs. 2A, 6A). The first two are short and unbranched, while the third to the fifth rays are the longest ones. Twelve pterygiophores are preserved in the holotype. The first is comparatively long and broad, inserted between the tenth and eleventh neural spines, whereas those posterior to it are much narrower (Fig. 2A).

The origin of the anal fin lies behind the end of the dorsal fin base, closer to the pelvic fin insertion than to the caudal fin base. The anal fin base is comparatively long, containing about 38 rays, of which the anterior six are longer than the posterior ones. At least 36 pterygiophores are preserved in the holotype. In specimen NHMG 038778, 38 pterygiophores can be counted. Anterior pterygiophores are longer than the posterior ones. The first pterygiophore inserts

between the last <del>pleural</del> rib and the first hemal spine (Figs. 2A, 3A).

(9) Vertebral Column. Twenty-three caudal vertebrae, not including two ural centra, and 324 nineteen abdominal vertebrae are recognized in the holotype (Fig. 2A-B). We added two to our 325 counts for the vertebrae that normally lie under the superficial bones of the skull and pectoral 326 girdle (e.g., opercle, cleithrum), thus, the total number of the preural vertebrae is about 44 in the 327 328 holotype. The length and depth of the vertebra is about equal, except the last several ones which are shorter than the anterior ones (Figs. 2A, 3A). There are two longitudinal ridges along the 329 lateral side of each vertebra, forming two pits on their lateral side (Fig. 3A). Halves of the neural 330 arches fused medially. The hemal spines start from the 21<sup>st</sup> or 22<sup>nd</sup> centrum, and their length 331 decreases gradually until the fourth or fifth preural centrum where they increase greatly to 332 support the fin rays of the lower caudal lobe (Fig. 7). 333

Nineteen pairs of ribs are present in the holotype, but in NHMG 033659 only 18 pairs of ribs could be counted. All the pleural ribs insert deeply into the centra. Ventrally, these ribs touch the lateral wings of the abdominal scutes. There are numerous thin and long epineural and epipleural intermuscular bones. The epineural series extends from the occiput to the first preural centrum. The epipleural series starts from approximately under the last three abdominal vertebrae and extends to about the first preural centrum. Longest epineural reaches the length about five to six centra, and the epipleural is about the length of four to five centra (Figs. 2A-B, 3A, 6).

(10) Caudal Skeleton and Fin. Caudal skeleton and fin are preserved relatively well in specimens NHMG 011646-011651. The neural and haemal spines of the second to the fourth or fifth preural centrum are elongated and somewhat flattened in that of the second and third preural centrum, and support a few caudal fin rays and procurrent rays. The structure of the caudal skeleton, collectively, differs from that in clupeiforms but closely resembles that in

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ellimmichthyiforms, i.e., bearing at least two autogenous uroneurals, and the first one without 346 fusing with the first preural centrum as in elupeoidei, though it is long and thick, extending 347 anteriorly to reach the anterodorsolateral side of that centrum. The second uroneural is much 348 shorter than the first one, extending anteriorly only to the anterior end of the second ural centrum, 349 although its distal end reaches that of the first one. It cannot be confirmed if a third uroneural is 350 351 present or not. There are two free ural centra, the first one is about equal in size to the first preural centrun, but the second one is much shorter than the first one. Six hypurals are present. 352 The proximal end of first hypural is in contact with but not fused to the first ural centrum, 353 354 although the narrow second hypural is fused to this centrum with its proximal end. The hypural 3 is the largest. Anteriorly, its enlarged proximal end contacts with the distal end of the second ural 355 centrum entirely constraining the hypurals above it to go forwardly to the tact with this centrum; 356 posteriorly, it expands, filling the entire space between the second and the fourth hypurals, then 357 there is no diastema between them, as the case in Armigatus brevissimus and the Eocene 358 *Diplomystus* species. The fourth to sixth hypurals are narrower and shorter in turn. The proximal 359 end of parhypural fused with the first preural centrum, which has a long and broad neural arch 360 and spine. There are three epurals (NHMG 011650). The caudal fin is deeply forked with the 361 upper and lower lobes of about equal in length, containing 19 principal fin rays (I<sub>7</sub> 9-8, I), and 362 eleven and nine procurrent rays above and below the principal caudal fin rays, respectively (Figs. 363 2A, 7). The proximal ends of the middle principal fin rays are preserved as impressions in 364 NHMG 011648, but relatively well-preserved in NHMG 011649, from which we can see that 365 these ends enlarge obviously. Caudal scutes can be seen in NHMG 011647. 366

367 (11) Predorsal Bone and Scutes. There are ten or eleven predorsal bones with thin anterior
 368 and posterior bony expansions. The anterior bones are broader than the posterior ones, and the

369 first three stretches almost vertically (Figs. 2A-B, 3A) or somewhat postero-ventrally (Fig. 6),
370 whereas the rest antero-ventrally.

There is a series of scutes along the dorsal margin from the occiput to the origin of the dorsal 371 fin in the holotype. The entire series includes about 55 small, equally-sized scutes (Fig. 2). For 372 poor preservation, the details of the scutes cannot be observed in this specimen. In specimen 373 374 NHMG 038778, about six predorsal scutes from just behind the occiput and about nine ones just anterior to origin of the dorsal fin can be recognized. The detail of the scutes cannot be observed 375 for poorly preserved also. However, it can be detected in NHMG 033658 (Fig. 6), in which 376 numerous small dorsal scutes are preserved along the dorsal margin from the occiput to the 377 seventh predorsal bone. Most of the scutes are displaced, some of them are even upside-down 378 379 and showing their smooth ventral surface, but many scutes show their dorsal surface ornamented with several radial ridges. In NHMG 033659, a few predorsal scutes with weak ridges are 380 preserved anterior to the dorsal fin. In NHMG 033680, several displaced dorsal scutes bearing 381 radial ridges as NHMG 033658 can be detected. No dorsal scutes are seen behind the dorsal fin 382 383 base.

About 24 prepelvic scutes are counted from the posterior edge of the coracoid to the insertion 384 385 of pelvic fin in the holotype (Fig. 2A-B) and in NHMG 033658 (Fig. 6). Fourteen postpelvic scutes are present in the holotype. Only first 11 postpelvic scutes are preserved in NHMG 386 033658. Several much smaller ventral scutes can be detected below the coracoids in the holotype 387 388 or displaced in the lower part of the head in NHMG 033659. The scutes behind the coracoid bear strong ventral spine and much higher lateral wings. The lateral wings are wider at their ventral 389 390 edges, but get narrower gradually dorsally, extending from the ventral edge of the body up to 391 about one quarter of the way to the vertebral column (Figs. 2A, 3A, 6).

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392 (12) Squamation/Scales. In NHMG 038778, impression of some squamations can be seen in
 393 the body above the vertebral column. The squamation is small in shape. The detail of it is not
 394 clear.

395

#### DISCUSSION

#### **396 Phylogenetic Relationships of the New Form**

Although the predorsal scutes of the new form do not expand laterally as in the diagnosis given by Grande (1982) of the Ellimmichthyiformes when he established the order, the presence of two parietals meeting at the midline, a beryciform foramen within the anterior ceratohyal, ornamentations on the predorsal scutes, and the structure of the caudal skeleton suggest that it apparently differs from clupeiforms but resembles ellimmichthyiforms (Grande, 1982, 1985). To further assess the systematic position of the new form, phylogenetic analyses are conducted.

Two data matrices were constructed for the phylogenetic analyses. Data matrix 1 (D1) used *Chanos chanos* as outgroup taxon, whereas data matrix 2 (D2) used *Elopse saurus* taking the place of *Chanos chanos* as outgroup taxon. Each data matrix includes 55 morphological characters and 40 taxa (see Apeendix 2S-3S), being analyzed using parsimony and Bayesian inference methods respectively.

The analysis of the D1, using parsimony criteria, generated 4 equally most parsimonious trees (PMTs). A strict consensus tree of 189 steps was built, with a consistency index (CI) of 0.323 and retention index (RI) of 0.670 (Fig. 8, Appendix Figure 1S). The cladogram of the strict consensus tree (SCT1) shows two main clades of Clupeomorpha: Clupeiformes and Ellimmichthyiformes. The monophyly of the Ellimmichthyiformes, including *Ornategulum* as the most basal taxon, is supported by the following features: two parietals meeting at the midline (2:0), two supramaxillary bones (8:0), presence of the "basipterygoid" process (9:1), anterior

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ceratohyal with foramen (11:1), and three epurals (38:0). Gasteroclupea and Sorbinichthys lie in 415 the basal position of the Ellimmichthyiformes, but do not form sister groups as suggested by 416 Marramà et al. (2019) and Boukhalfa et al. (2019). Sorbinichthyidae sensu Murrey and Wilson 417 2013 including only two Sorbinichthys species, is strongly supported by a number of derived 418 characters: broad dorsal process of the posttemporal (15:2), posterior predorsal scutes laterally 419 420 expanded (41:1), the most posterior predorsal scutes enlarged (44:1), high number of abdominal scutes (51:2), but less number of postpelvic scutes (52:1). Monophyly of Armigatus is supported 421 by sharp proximal end of first hypural (27:1), predorsal scute series incomplete (39:0). 422 423 Monophyly of *Diplomystus* (exclude *D. solignaci* Gaudant 1971) is supported by the presence of sub-rectangular scutes in anterior and posterior predorsal series (40:1 and 41:1), and presence of 424 series of spines on posterior margin of lateral wings of predorsal scutes (42:1). Diplomystus 425 solignaci is a member of the paraclupeid clade as suggested by many previous studies (Alvarado-426 Oetega et al., 2008; Murrey & Wilson 2013; Figuereido & Ribeiro 2016; Marramà et al., 2019; 427 428 Boukhalfa et al., 2019). Armigatus is in a more basal position than that of Diplomystus as suggested by Alvarado-Oetega et al. (2008) and Figuereido & Ribeiro (2016). Our new form, 429 Guiclupea, forms sister group to Diplomystus. The synapomorphies of the two are high 430 431 supraoccipital crest (4:1), pelvic-fin insertion in advance of dorsal-fin origin (22:0), number of predorsal scutes more than twenty (46:1). The membership of Paraclupeidae, not include 432 Kwangoclupea and Codoichthys as some previous studies suggested (e.g., Murrey and Wilson, 433 434 2013; Francisco and Figueiredo, 2016; Marrama et al., 2018; Boukhalfa et al., 2019) but in some way consistent with the recent analysis conducted by Vernygona and Murray (in press), is 435 436 supported by the dorsal outline forming a marked angle at dorsal-fin insertion (1:1), first 437 uroneural extending forward to second preural centrum (32:0), and predorsal scutes with ridges

438 on the dorsal surface (44:1).

The analysis of the D1 using Bayesian inference method, generated a Bayesian Inference tree (BIT) (Fig. 9). As in SCT1, the monophyly of *Sorbinichthys*, *Armigutus*, *Diplomystus*, and Paraclupeidae were supported, and *Guiclupea* is sister to *Diplomystus*. Unfortunately, the monophyly of the Clupeiformes cannot be supported, and *Gasteroclupea*, *Sorbinichthys*, and *Kwangoclupea* lies in a more derived position than that in SCT1.

The analysis of the D2, using parsimony criteria, recovered 16 PMTs. The SCT of the 16 PMTs (see Appendix Figure 3S) is 188 in step, the CI is 0.324, and the RI is 0.671. The cladogram of the SCT2 shows that *Ornategulum* is in a basalmost position of the superorder, i.e., *Ornategulum* does not belong to the Ellimmichthyiformes as suggested by SCT1 and BIT1. The monophyly of the Clupeiformes, *Sorbinichthys*, and the remaining members of the Ellimmichthyiformes clade exclusive of *Gasteroclupea* was all supported, and the last one has the same topology as in SCT1.

The topology of the BIT2, resulted from the analysis of D2, is very similar with that of the 451 BIT1 except for the position of Ornategulum and Denticeps. In BIT2, Ornategulum lies in the 452 most basal position as in SCT2, and *Denticeps* is sister group to the two clupeoid genera. It 453 seems that the applying different outgroup taxa effects the position or the assignment of 454 Ornategulum, as demonstrated by previous analyses (Murrey and Wilson, 2013; Marramà et al., 455 2019; Figuereido and Ribeiro 2017; Boukhalfa et al., 2019). In addition, the positions of 456 457 Armigatus, Gasteroclupea, Sorbinichthys, Kwangoclupea, and Codoichthys are the same in the two SCTs and in the two BITs respectively, but different between the SCTs and the BITs. In 458 BITs, Armigatus lies in the basalmost position of the order but it is not so in SCTs. In BITs, 459 460 Gasteroclupea belongs to the Sorbinichthyidae as suggested by Marramà et al. (2019) and

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Boukhalfa et al. (2019); but in SCTs, Gasteroclupea does not form sister group to Sorbinichthys. 461 Kwangoclupea forms sister group to Codoichthys and lies in a relatively basal position in the two 462 SCTs, whereas it lies in a relatively derived position and belongs to the Paraclupeidae in the two 463 BITs. These differences between the general topologies of the SCTs and BITs probable due to 464 that the information contained in the dataset is insufficient to draw firm conclusions about their 465 466 relationships as pointed out in the recent analyses of the phylogeny of Ceratomorpha (Bai et al., 2020). To improve the understanding of relationships of the group, more phylogenetically 467 informative fossils and more complete data are needed. 468

Although there are discrepancies between the MPTs and BITs, and between the trees with 469 alternative outgroups, the general topologies of the four trees mentioned above are basically 470 similar, and all the four trees suggest that the new form is a member of the ellimmichthy forms 471 and forms a sister group to Diplomystus, The close relationship of them is supported by high 472 supraoccipital crest (4:1), pelvic-fin insertion in advance of dorsal fin origin (22:0), and number 473 of predorsal scutes  $\geq 20$  (46:1). Actually, in addition to these synapomorphies, the new form and 474 Diplomystus, especially the Eocene species, i.e., D.dentatus and D. shengliensis, share many 475 more similar characters, such as having an elongated fusiform body form, dorsal outline curved 476 477 gently, no ornamentation on the skull bones, with teeth on entopterygoid, high number of anal fin rays (23-25 in D. birdi, 27 in D. dubertreti, 38-41 in D. dentatus, about 39 in D. shengliensis and 478 55 in *Guiclupea superstes*), and no diastema between second and third hypural (but there is a gap 479 480 between second and third hypural in the Later Cretaceous species, i.e., D. birdi and D. dubertreti (Chang & Maisey 2003, p27)). The last character also occurs in pristigasteroids, and 481 482 osteoglossids, some elopomorphs, and a number of ostariophysans (Chang & Maisey, 2003). The 483 differentiation between *Diplomystus* and *Guiclupea* is in the shape and ornamentation of the

predorsal scutes (sub-rectangular vs. ovate, presence vs. absence of pectinate posterior border, 484 dorsal surface smooth vs. with radial ridges), and the number of predorsal bones (6-8 vs. 10-11). 485 Accordingly, the new form is a distinct genus and species, and can be easily distinguished from 486 Diplomystus. Comb-like teeth along the posterior edges of the dorsal scutes is a derived character 487 of Diplomystus. In addition to Diplomystus, sub-rectangular predorsal scutes also occur in most 488 489 members of the Paraclupeidae. As far as the radial ridges on the dorsal surface of predorsal scutes is concerned, they usually occur in paraclupeids, such as Paraclupea, Ellimmichthys, 490 Ellimma, Triplomystus, etc., these forms usually have a marked angle at the insertion of dorsal 491 fin, and sub-rectangular dorsal scutes at least in the posterior part of the scute series. The new 492 form is distinct from them in the shape of the body and predorsal scutes. Among species with 493 ornamentation on predorsal scute, the new form resembles *Scutatuspinosus itapagipensis* in the 494 shape of body and predorsal scutes (not laterally expanded), and posterior expansion of the third 495 hypural, leaving no gap or notch between the second and third hypurals. However, there are 496 obvious differences between the two forms in the number of predorsal scutes and anal fin rays, 497 ornamentation on the skull roof bones, and the size and shape of the abdominal scutes. On the 498 whole, the new form displays a mosaic combination of characters. It bears radial ridges on the 499 500 dorsal surface of predorsal scutes as the paraclupeids, but the scutes are all about the same size as in Armigatus and Diplomystus. Consequently, Guiclupea can easily be distinguished from all 501 502 known ellimmichthyiforms in number and morphology of the dorsal scutes. So that predorsal 503 scutes with ridges on the dorsal surface are not unique to Paraclupeidae.

#### 504 Body shape and size of the Ellimmichthyiformes

505 The ellimmichthyiforms are diverse both in general morphology and body size. Generally, 506 ellimmichthyiforms show two types of body form. One bears a deep body, with the maximum

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depth/standard length (MD/ST) larger than 50%, some of them even with the MD roughly equal to, or slightly larger than the ST in adult specimens, e.g., in *Tycheroichthys dunveganensis* and *Rhombichthys intoccabilis* (Hay et al., 2007; Khalloufi et al., 2010). Most of this kind of fishes are referred to paraclupeids. The other kind of fishes have an elongate fusiform shape, the MD/ST often lower than 50%. They occupy a relatively basal position in the Ellimmichthyiformes, such as *Armigatus, Diplomystus*, and *Guiclupea*.

Body size of ellimmichthy forms ranges from several centimeter to about 65 cm in TL. The 513 known smallest fish is *Eoellimmichthys superstes* with the TL and ST about 17.5 mm and 13.7 514 mm, respectively (Marramà et al., 2019). Many species are with the TL shorter than 100 mm, 515 such as Armigatus alticorpus, A. dalmaticus, A. oligodentatus, Codoichthys carnavalii, 516 shengliensis, "Diplomystus" trebicianensis, 517 Diplomystus Ellimmichthys maceioensis, *Eoellimmichthvs* superstes, Gasteroclupea branisai. *Scutatuspinosus* 518 itapagipensis, Sorbinichthys africanus, Thorectichthys marocensis, T. rhadinus, and Tunisiaclupea speratus. 519 520 Species with TL larger than 200 mm are rare. The TL of *Rhombichthys intoccabilis* reaches about 230 mm in the holotype (Khalloufi et al., 2010, fig. 3). The predorsal length of 521 Horseshoeichthys armigserratus, from the Maastrichtian of Canada, is 172 mm, with the 522 estimated SL about 260-280 mm (Newbrey et al., 2010). Specimens with SL over 300 mm are 523 only seen in *Diplomystus dentatus* from the Eocene of the eastern coast of the Pacific and 524 525 Guiclupea superstes from the Oligocene of the western coast of the Pacific so far. The former 526 reaches a TL of about 650 mm (Grande, 1982) while the latter reaches a ST about 600 mm. They should be the largest ellimmichthy form fishes. It is noted that the order or the Diplomystus clade 527 528 seems to show a trend to enlarge their body size. In the Cretaceous, members of this order 529 usually have a small body length; it is until the end of the Cretaceous, some member (e.g.,

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530 Horseshoeichthys armigserratus) became relatively large body-sized; in the late Paleogene,
531 some members developed large body size. It is worth mentioning that the fishes with large body
532 size all are members of *Diplomystus* clade (*Horseshoeichthys* forms sister group to *Diplomystus*533 species (Veryngora & Murray, in press)) and occur along the coast of the Pacific.

A recent study based on a comparative analysis indicates that across the Clupeiformes 534 535 diadromous species are larger than non-diadromous species (Bloom et al., 2018). Another study, based on over 4500 migratory and non-migratory species of ray-finned fishes also shows that 536 migratory species are larger than non-migratory relatives in nearly all clades and across all 537 modes of migration (Burns & Bloom, 2020), No association of body size with trophic position 538 was found (Bloom et al., 2018). From these two studies, we have reason to speculate that the 539 wide-spread *Diplomystus* clade possibly involve some diadromous or migratory species which 540 have large-sized body and are able to be widely distributed along the coast of the Pacific, for 541 increased body size is an adaptation to mitigate the energetic expense of long-distance migration 542 (Bloom et al., 2018). Then the restriction of the species of the Sorbinichthyidae and Armigatidae 543 to the circum-Mediterranean region may be associated with their small body size. 544

545 Paleobiogeographic history of the Ellimmichthyiformes

The Ellimmichthyiformes is an extinct cosmopolitan group of fishes, with members known from marine and non-marine sediments of South and North America, Africa, coastal regions of Mediterranean, and East Asia (Fig. 1). The oldest known so far are from Hauterivian-Barremian. It includes *Scutatuspinosus itapagipensis* and *Eillmmichthys longicostatus* from northeastern Brazil (Figueiredo & Ribeiro 2017; Cope, 1886), *Ezkutuberezi carmenae* from northern Spain (Poyato-Ariza et al., 2000), *Tunisiaclupea speratus* from southern Tunisia (Boukhalfa et al.,

2019), and Paraclupea chedungensis from eastern China (Sun, 1956; Chang & Grande, 1997; Hu 552 et al., 2017). All these species belong to the Paraclupeidae. The paraclupeid fishes are abundant 553 during late Early Cretaceous (Aptian-Albian). There are *Ellimma branneri* and *Ellimmichthys* 554 maceioensis from Alagoas (Schaeffer, 1947; Chang and Maisey 2003; Malabarba et al., 2004), 555 and Ellimma longipectoralis from Santos Basin (Polck et al., 2019) of Brazil; Ellimmichthys 556 goodi from Equatorieal Guinea (Eastman, 1912); and Paraclupea seilacheri from Puebla, 557 Mexico (Alvarado-Ortega and Melgarejo-Damián 2017). In addition to paraclupeids, there are 558 Codoichthys carnavalii from the Aptian of Brazil (de Figueiredo and Ribeiro, 2016) and 559 Forevclupea loonensis from the Albian of Canada (Vernygora et al., 2016, the authors thought 560 this species should be closely related with *Scutatuspinosus itapagipensis*). The close relationship 561 between the Early Cretaceous fish faunas from northeastern South America and from western 562 Africa might have resulted from the contiguous margins of Brazil and West Africa during the 563 Early Cretaceous (Chang and Grande, 1997). But there is little geologic evidence to support an 564 Early Cretaceous non-marine paleogeographic connection between the eastern Asiatic margin 565 and western Gondwana. Consequently, the distribution pattern of paraclupeids in the Early 566 Cretaceous arouses a biogeographic conundrum. No favorable hypothesis well deciphers this 567 distribution pattern to date. Chang and Maisey (2003) suggested that either a substantial portion 568 of their non-marine fossil record is missing or their distribution involved marine dispersal. 569 During the early Late Cretaceous, the Ellimmichthy formes reaches its greatest diversity. 570 Not only the Paraclupeidae is highly diversified in the Mediterranean region and extended their 571

572 range to North America (*Tycheroichthys dunveganensis* from Canada, Hay et al., 2007), but all

573	other main ellimmichthyiform clades, i.e., Sorbinichthydae, Armigatidae or Armigatus, and
574	Diplomysus, occurred and flourished in the Cenomanian with their oldest record from the eastern
575	Tethys (Lebanon) (Woodward, 1895; Signeux, 1951; Grande, 1982; Zhang et al., 1985;
576	Bannikov & Bacchia, 2000; Murray & Wilson, 2011; 2013; Murray et al., 2016). With all main
577	clades first occurring there, undoubtedly, the circum-Mediterranean region is a hotspot $\mathbf{ef}$ the
578	evolution of ellimmichthyiforms. Species of <i>Diplomysus</i> are also found from the Cenomanian
579	English chalk (Forey, 2004). The diversity of ellimmichthyiforms during the early Late
580	Cretaceous was probably resulted from the global sea-level rising and the consequential land-sea
581	distribution which increased food input, dispersal routes, and habitat fragmentation for these
582	fishes (Boukhalfa et al., 2019). In the end of the Cretaceous, the diversity of the
583	Ellimmichthyiformes suddenly declined. Only Gasteroclupea branisai from South America and
584	Horseshoeichthys armigserratus from North America are known to date.
585	Rare ellimmichthyiforms survived after the Cretaceous-Tertiary boundary. Gasteroclupea
586	branisai, Eoellimmichthys superstes, and Diplomystus clade are the exception. Gasteroclupea
587	branisai firstly occurred in the Late Cretaceous of South America and survived to the Danian of
588	Argentina and Bolivia (Signeux, 1964; Marramà & Carnevale, 2017). Eoellimmichthys
589	superstes is a paraclupeid from the marine Eocene of Italy. Interestingly, Diplomystus clade not
590	only survived up to the Oligocene, but also had a relatively wide distribution range along both
591	sides of the Pacific eoast (see Fig. 1). Horseshoeichthys armigserratus occurred on west coast
592	of North America in the end of the Cretaceous (Newbrey et al., 2007). The Eocene Diplomystus
593	occurred on both sides of the Pacific Ocean (along the coast of the Bohai Gulf, east China and

594	the west coast of North America) bearing striking similarity in morphology (Zhang et al., 1985;
595	Chang & Maisey, 2003). Their sister group, Guiclupea superstes, survived to the Oligocene as
596	the youngest ellimmichthyiform fish. The Eocene "transpacific" distribution pattern of
597	Diplomystus and other fishes and terrestrial vertebrates has long been noted by paleontologists
598	(Chang & Chow, 1978; Grande, 1982, 1985; Zhang et al., 1985). A broad connection between
599	Asia and North America in the Bering Strait area and temporary desalination of the Arctic
600	Ocean could have facilitated the dispersal of these fishes (Chang & Maisey, 2003). It is
601	interesting to find that all ellimmichthyiform fossil localities (Fig. 1) are near to the recent coast.
602	Many contemporaneous fish fauna have been found from inland areas of China, Mongolia, and
603	East Kazakhstan (Tang, 1959; Liu et al., 1962; Wang et al., 1981; Sytchevskaya, 1986), but no
604	ellimmichthyiforms have been reported from there to date. In addition, most Recent
605	Clupeomorpha inhabit in the sea. It seems reasonable to suggest that the origin and dispersal of
606	these fishes must have something to be related with the sea.
607	CONCLUTION
608	Guiclupea superstes from the Oligocene of southern China is the youngest
609	ellimmichthyiform known to date. Its occurrence indicates that the Ellimmichthyiformes had a
610	wider distribution range and a longer evolutionary history than previously knew. Guiclupea
611	superstes is closely related to Diplomystus, which suggests that the dorsal scutes with ridges on
612	dorsal surface is not a character unique to paraclupeids.
613	Paraclupeids is very diverse and widely distributed during the Early Cretaceous, and

614 survived to the Eocene in circum-Mediterranean area. Sorbinichthys and Armigatus are restricted

to the Mediterranean region in the early Late Cretaceous. *Diplomystus* clade might originated not
later than early Late Cretaceous, and disappeared from Europe and Middle East after early Late
Cretaceous, but still prosperous in the Pacific coast in the end of the Cretaceous to Eocene and
survived there until the Oligocene. This made the Ellimmichthyiformes to obtain a particularly
long distribution range along the West Pacific.

There is still no consensus among ichthyologists on the phylogenetic relationships of the Ellimmichthyiformes, especially the relationship of *Armigatus* and *Diplomystus*, the position of *Gasteroclupea* and *Codoichthys*. Besides, there are discordances between the most parsimonious tree and the Bayesian Inference tree. To solve those problems, more informative specimens and characters are needed to add to the dataset.

625

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# Figure 1

Map of the main localities of ellimmichthyiform fossils

Map of the main localities of ellimmichthyiform fossils



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#### Guiclupea superstes, gen. et sp. nov

*Guiclupea superstes*, gen. et sp. nov. **A**, and **B**, photograph of the holotype (NHMG 005532); **C**, close up of the complete predorsal scutes series maeked by the white box in A.



Guiclupea superstes, gen. et sp. nov

**A**, photograph of the NHMG 033659; **B**, tentative restoration mainly based on the holotype and paratypes.



Guiclupea superstes, gen. et sp. nov

**A**, photograph, and **B**, line drawing of the head in NHMG 033659. Anterior facing right. **C**, photograph of a left premaxilla, NHMG 033685; **D**, photograph of an incomplete dentary, NHMG 033682, showing the oral teeth; **E**, photograph of a dentary, NHMG 033683; **F**, photograph, showing anterior ceratohyal and entopterygoid in NHMG 033661; **G**, photograph, showing posterior ceratohyals in NHMG 033680.

#### Manuscript to be reviewed



Guiclupea superstes, gen. et sp. nov.

**A**, photograph and **B**, line drawing of the disarticular skull and anterior trunk bones of NHMG 033660.



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Guiclupea superstes, gen. et sp. nov

**A**, photograph of NHMG 033658, **B**, and **C**, close up of the predorsal scutes in the black boxes of B and C in A, respectively.



*Guiclupea superstes*, gen. et sp. nov.

**A**, photograph and **B**, line drawing of NHMG 011648. Anterior facing left.

### Manuscript to be reviewed



Strict consensus of most parsimonious trees

Strict consensus of most parsimonious trees retrieved in TNT 1.5 based on 55 morphological characters and 40 taxa, with *Chanos chanos* being used as outgroup taxa. Characters supporting each node are listed along corresponding branches. Characters above black spot have consistency index of 1. Numbers at each node represent Bremer support/bootstrap values. Only bootstrap values higher than 50% are shown on the tree.

#### Manuscript to be reviewed



#### Cladogram

Cladogram resulting from Bayesian phylogenetic analyses based on 55 morphological characters and 40 taxa, with *Chanos chanos* was used as outgroup taxon. The numbers at the internal nodes are the posterior probabilities of the corresponding clades.

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#### Table 1(on next page)

TABLE

Measurements and counts for the specimens of the † *Guiclupea superstes*, gen. et sp. nov. Measurements are in millimeters.

- 1 TABLE 1. Measurements and counts for the specimens of the *†Guiclupea superstes*, gen. et sp.
- 2 nov. Measurements are in millimeters.
- 3

	004929	005532	033658	033659
total length		526.0		630+
standard length (SL)	502+*	453.0	410+	585+
head length	175	132.0		140+
head length/SL		29.1%		
head depth	169.0	116.0		165+
body depth	213	137.0	150	210
body depth/SL		30.2%		
predorsal length	333.5	252.8		310+
predorsal length/SL		55.8%		
prepelvic length	293.7	242.0		260+
prepelvic length/SL		53.4%		
preanal length	387.0	309.3		358+
Preanal length/SL		68.3%		
dorsal fin rays		Ii, 12	at least 13	
anal fin rays		~35		
pterygophores of anal fin		~36	29+	22+
pectoral fin rays		at least 12	18	12+
pelvic fin rays		5~6	5~6	
abdominal vertebrae	20	22	20	20
caudal vertebrae (exclude u1, u2)	12+	23		24
total vertebrae (exclude u1, u2)	32+	45		44
predorsal bones	10	10 or 11	10	10
abdominal scutes	at least 36	~38	35+	27+
pre-pelvic scutes	~24	~24	~24	16+
post-pelvic scutes	at least 12	14	11+	11+
pre-dorsal scutes		55	27+	
pairs of ribs	18	20	18	19

4 \*: "+" stand for the actual digital larger than this digital because of the specimen is incomplete or not well-

5 preserved.