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The oldest gnathostome teeth

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1 The oldest gnathostome teeth

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16 Summary paragraph

Mandibular teeth and dentitions are features of jawed vertebrates that were first 17 acquired by the Palaeozoic ancestors¹⁻³ of living chondrichthyans and osteichthyans. 18 The fossil record currently points to the latter part of the Silurian⁴⁻⁷ (circa 425 million 19 years ago) as a minimum date for the appearance of gnathostome teeth and to the 20 evolution of growth and replacement mechanisms of mandibular dentitions in the 21 subsequent Devonian Period^{2,8-10}. Here we provide the earliest direct evidence for 22 jawed vertebrates by describing Qianodus duplicis, a new genus and species of a 23 lower Silurian gnathostome based on isolated tooth whorls from Guizhou Province, 24 China. The whorls possess non-shedding teeth arranged in a pair of rows that 25 demonstrate a number of features found in modern gnathostome groups. These 26 include lingual addition of teeth in offset rows and maintenance of this patterning 27 throughout whorl development. Our data extend the record of toothed gnathostomes 28 by 14 million years from the upper Silurian into the lower Silurian (circa 439 million 29 years ago) and are significant in documenting the initial diversification of vertebrates. 30 They add to mounting fossil evidence supporting an earlier emergence of jawed 31 32 vertebrates as part of the Great Ordovician Biodiversification Event.

33

Jawed fishes have a patchy and controversial fossil record through the Ordovician 35 and the Sllurian^{11,12}. The unequivocal evidence for their first appearance is currently 36 constrained by the discovery of upper Silurian (c. 428-420 million years ago) 37 'placoderms'¹³⁻¹⁵ and stem sarcopterygians^{7,11} from South China and Vietnam and 38 the disarticulated remains of the stem osteichthyans Andreolepis and Lophosteus 39 from Europe and North America^{4,5}. The earliest remains of chondrichthyans 40 (elasmobranchs, holocephalans and their 'acanthodian' ancestors) include body 41 fossils from the upper Silurian¹⁶ and Lower Devonian¹⁷⁻²⁰, isolated teeth of upper 42 Silurian^{21,22} and basal Devonian²³ age and, more contentiously^{11,24}, isolated scales 43 and fin spines from the Ordovician¹² and Silurian^{25,26}. Convergence of scale-based 44 characters and the paucity of available data^{12,27} have further hampered the 45 assignment of these fragmentary specimens to jawed vertebrates. Their 46 incorporation into phylogenetic analyses has underscored this ambiguity by 47 producing tree topologies¹² that are incongruent with the composition of the 48 chondrichthyan stem and crown groups established from articulated fossils^{24,28}. 49 The present study describes tooth whorls from the Llandovery (Aeronian), 50 lower Silurian of China (Fig. 1) that provide much-needed data on the dental 51 conditions of early toothed vertebrates. With the investigation of the whorls we seek 52 to 1) characterise their tooth patterning and morphological features, 2) provide a 53 phylogenetic hypothesis for their affinities, and 3) highlight their implications for the 54 timing of the origin and the diversification of jawed vertebrates during the Palaeozoic. 55

56

57 Systematic palaeontology

58

Gnathostomata Gegenbaur, 1874 (ref.²⁹)

59	Chondrichthyes Huxley, 1880 (ref. ³⁰) sensu Coates et al. ²⁴
60	<i>Qianodus duplicis</i> gen. et sp. nov.
61	Etymology: The generic name is a composite of Qian, referring to the ancient
62	Chinese name of present-day Guizhou Province, and the Greek odus, meaning
63	tooth. The specific name <i>duplicis</i> (double) alludes to the paired tooth rows of the
64	whorls and derives from the genitive case of the Latin 'duplex'.
65	Holotype: Isolated tooth whorl IVPP V26641 (Figs. 1c–f and 2a, f and Extended
66	Data Fig. 2b, c, e).
67	Referred material: Twenty-three tooth whorls (including the holotype).
68	Locality and horizon: Section through the Rongxi Formation at Leijiatun village
69	(Shiqian County), Guizhou Province, China (Fig. 1a, b). Ozarkodina guizhouensis
70	conodont biozone (c. 439 Ma; late Aeronian, Llandovery, Silurian; see
71	supplementary information).
72	Diagnosis: A jawed vertebrate possessing tooth whorls with primary teeth organized
73	into two mutually offset rows. Lateral sides of the whorl base carry accessory tooth
74	rows positioned lower than the primary teeth.
75	Remarks: The recovered Qianodus specimens co-occur in the Rongxi Formation
76	together with disarticulated scales and fin spines of other yet-to-be-described
77	acanthodian-grade ²⁸ taxa. The available evidence does not allow us to categorically
78	confirm or rule out attribution of any of this material to Qianodus given the
79	preservation of the whorls as isolated elements and the development of tooth whorls
80	in a number of 'acanthodian' lineages ³¹ . Our decision to erect a new taxon for the
81	tooth whorls follows established practice ⁹ of formally describing isolated dental

elements of Palaeozoic chondrichthyans if they are identifiable by a unique set offeatures.

Tooth whorl morphology and tissue structure

Specimens referred to Qianodus duplicus gen. et sp. nov. are up to c. 2.5 mm long 85 (Extended Data Fig. 3) and possess a pair of primary tooth rows borne on a raised 86 medial crest of the whorl base (Figs. 1c, f, 2a, b, e and Extended Data Fig. 1g). 87 Primary teeth have a staggered arrangement (Fig. 2a-c, e, f) and their size 88 increases within the rows in a manner recorded in whorls with non-shedding teeth 89 where consistently larger elements are added lingually^{31,32}. Due to the offsetting, it 90 can be determined that the more labial (progenitor) row is initiated before the 91 adjacent (trailing) row (Fig. 2a, b), with the distinction between the two only possible 92 in specimens preserving the diminutive early tooth generations. The whorls exhibit 93 94 left and right configurations produced by 'mirror image' arrangements of the primary tooth rows (Fig. 2a, b). We interpret the progenitor row of either type as occupying a 95 mesial position based on comparison with modern elasmobranch dentitions^{33,34} that 96 97 possess alternate patterning of transverse tooth rows.

Most undamaged whorl bases have a deep, dome-like appearance, although 98 the two smallest specimens exhibit recurved, low profiles (Extended Data Fig. 1j-n). 99 The lateral faces of whorl bases carry rows of accessory teeth (Figs. 1, 2a-c, e, 3c 100 and Extended Data Fig. 1j, k) distributed on arched ridges that follow the curvature of 101 102 the whorl crest. The earliest formed generations of accessory teeth within each row are positioned at the inturned tip of the whorl spiral labially of the primary teeth (Fig. 103 2c, e and Extended Data Fig. 1a, d, j, k). Unlike the offset primary teeth, they are 104 organised in closely set symmetrical pairs and are distinguished on that basis. The 105

accessory tooth fields lack the alternate arrangement of primary teeth but
 demonstrate their characteristic labial increase in tooth size along individual rows. At
 corresponding positions accessory teeth are consistently smaller than the primary
 teeth of the whorl crest.

Due to dissolution and recrystallization, the microstructure of tooth whorl tissues is largely obscured by diagenetic artefacts. These preclude the identification of tissue types. Only large-calibre vascular spaces are apparent inside the whorls.

113 Whorl teeth demonstrate pulp-like spaces infilled by radiopaque crystalline 114 (Fig. 2d) and radiotransparent globular (Extended Data Fig. 2g, h) material. These 115 fabrics also account for the extensive recrystallisation of the tooth walls where they 116 exploit voids created by diagenetic dissolution of the mineralised tissue (Fig. 2d).

Tooth whorl bases are composites of a superficial spongiose tissue and an 117 inner compact tissue (Fig. 2d, f and Extended Data Fig. 2). The spongiosa is best 118 developed along the medial crest of the whorl and diminishes in thickness down its 119 sides and lingually (Fig. 2d, f and Extended Data Fig. 2a, b, d). Along the whorl crest 120 121 it forms socket-like depressions housing the ankylosed to the base attachment portion of primary teeth (Fig. 2d, f and Extended Data Fig. 2a–d). Volume renderings 122 of the spongiosa reveal an extensive system of canal-like spaces that follow the 123 curvature of the whorl (Fig. 2d, f and Extended Data Fig. 2c). The canals are 124 intercepted by the pulps of the whorl teeth and connect successive tooth generations 125 of the primary and accessory rows (Fig. 2f and Extended Data Fig. 2c, f). 126

127 The compact zone of the base is thickest labially and tapers off lingually. The 128 superimposed spongiosa extends over the compact zone on all sides to form the 129 base periphery (Fig. 2d, f and Extended Data Fig. 2a, b, d, e). No histological data

could be retrieved from the compact tissue, as its original structure is replaced byglobular diagenetic material (Fig. 2d and Extended Data Fig. 2a).

132

Growth of the tooth whorls

Two of the whorls (Extended Data Fig. 1j–n) are recognised as early developmental 134 stages, as these are of markedly smaller size, have fewer tooth rows/generations 135 and shallower base profiles than the rest of the recovered specimens (Extended 136 Data Fig. 3). This interpretation is further supported by the correspondence between 137 the spacing and size of their teeth and those carried by the labial end of the whorl 138 139 spiral of the larger ontogenetically older specimens. The early development whorls possess only the first, orally facing, pair of accessory rows, which contain one less 140 tooth per row in comparison to their primary tooth rows (2 versus 3). Mature whorls 141 (Fig. 2c) similarly have the largest number of tooth generations (maximum of 8) 142 present within the primary rows and exhibit progressively decreasing tooth counts in 143 the first, second and lower accessory rows (Fig. 2e). These observations suggest 144 that the first formed whorl teeth were those of the primary tooth rows, with the 145 inception of accessory rows occurring later in development in a sequence following 146 the expansion of the whorl base. 147

148

149 *Qianodus* in the context of vertebrate dentitions

Although the whorls of *Qianodus* show significant post-mortem abrasion and
 diagenetic alteration, the material allows direct comparison with the oropharyngeal
 teeth/denticles of gnathostomes. The pronounced curvature of the whorls and their

ordered tooth rows contrast with the aggregates of pharyngeal denticles of the type 153 developed by the lodont jawless stem-gnathostomes^{1,35}. The the lodont pharyngeal 154 denticles constitute `sheets` of fused scales¹ that, although exhibiting polarity, lack 155 the strictly unidirectional sequence of addition of the whorl teeth of Qianodus and 156 their arrangement into discrete rows. Compound buccopharyngeal denticles have 157 also been recorded in Palaeozoic symmoriid chondrichthyans^{36,37} but similarly lack 158 the arched profiles of tooth whorls. However, the spiral shape and longitudinal tooth 159 rows of Qianodus are consistent with an origin within the dental groove of the jaw as 160 elements that transverse the jaw rami^{32,33,38}. Similarly, identification of the whorls as 161 extraoral elements akin to the cheek scales found in ischnacanthid 'acanthodians'39 162 is considered implausible, as these do not exhibit the spiral-like morphology of the 163 whorls. 164

Possession of mandibular teeth, and by extension jaws, places *Qianodus* firmly within jawed vertebrates. Its phylogenetic position is further constrained by our parsimony analysis (Extended Data Figs. 4, 5) that recovers *Qianodus* as a derived stem chondrichthyan, in accord with a previously proposed origin of tooth whorls within crown gnathostomes^{24,31,40}.

Jawed stem gnathostomes, traditionally referred to as 'placoderms'¹¹, reveal that 170 primitive vertebrate dentitions lacked discrete dental elements such as tooth whorls, 171 but instead were composed of a few large toothed plates borne on the mandibular 172 arches^{3,8,15}. In one of their major divisions, the arthrodires, oral teeth develop radially 173 or bidirectionally along the length of gnathal bones^{10,41}. However, a recent 174 investigation³ of acanthothoracid 'placoderms' has identified lingual addition of 175 replacement teeth directed within rows in the gnathal plates of Kosoraspis. The 176 latter's labio-lingual tooth patterning is comparable to that of tooth whorls but, 177

similarly to the dentition of some arthrodires^{41,42}, *Kosoraspis* gnathal plates carry 178 multiple tooth rows whose number is not constantly maintained across all plates. 179 The disarticulated state of the material and its poor histological preservation 180 allows Qianodus to be coded only for a limited set of dental characters 181 (Supplementary Data 6). Improving the robustness of the phylogenetic hypothesis 182 (Extended Data Figs. 4, 5) will depend on new information on the detailed histology 183 of the whorls of *Qianodus*, as well as the currently missing data on the morphology 184 of the fish. These could potentially be collected by further sampling of the Rongxi 185 Formation and coeval assemblages¹⁴. 186

Data from whorl-forming chondrichthyans and osteichthyans (Fig. 3) places the 187 dentition of Qianodus within a broader phylogenetic context. Within osteichthyans, 188 tooth whorls are documented mostly among the stem sarcopterygians where they 189 are restricted to parasymphyseal positions^{38,43,44} of the inner dental arcade⁴⁴. 190 Sarcopterygian whorls carry a medial row of similar-sized teeth that are shed 191 repeatedly at the labial end of the whorl^{38,43}. Recurring tooth replacement through 192 hard-tissue resorption also occurs in the tooth cushions of the stem osteichthyans 193 Andreolepis and Lophosteus^{4,5}. These are, akin to tooth whorls, elements of the 194 inner dental arcade with a non-mineralised attachment to the jaw ramus⁵. Tooth 195 196 cushions support multiple tooth rows within which replacement teeth are deposited on top of resorbed older tooth generations^{4,5}. The resorptive tooth shedding of 197 osteichthyan parasymphyseal whorls and tooth cushions^{4,5,43}, nevertheless, 198 contrasts with the tooth retention seen in the whorls of Qianodus (Figs. 1, 2). Tooth 199 200 retention is seen in the tooth whorls of chondrichthyans³¹ and it is most parsimonious to look among these for a 'template' of the dentition of Qianodus. 201

In the chondrichthyan total group tooth whorls are restricted to symphyseal 202 and/or parasymphyseal positions^{9,19} or can also be found along the lateral extent of 203 the jaw as the only elements of the dentition^{18,32,45,46} (Fig. 3). The asymmetry of tooth 204 offsetting and its mirroring in *Qianodus* whorls (Fig. 2a, b) points to their origin from 205 lateral positions on both jaw rami rather than formation on the jaw symphysis. An 206 alternative explanation for *Qianodus* whorls as symphyseal and/or parasymphyseal 207 208 elements is considered less credible as single tooth rows occupy each of these positions in crown gnathostomes with whorls confined to the jaw symphysis^{19,38}. 209 210 Moreover, the size differences between Qianodus whorls of comparable stage of development are consistent with the mesial enlargement of whorl dimensions along 211 the jaw ramus observed in the stem chondrichthyan *Doliodus*³² (Extended Data Fig. 212 3). Whorl-based dentitions occur in the climatiids (e.g. *Ptomacanthus* and 213 *Climatius*^{17,18}) and in *Doliodus*³² (Fig. 3) and represent the stem chondrichthyan 214 condition for developing tooth rows that traverse the jaw rami (Fig. 3). Tooth 215 retention in whorls is also documented in the chondrichthyan crown group within 216 holocephalans⁴⁷ (Extended Data Fig. 4), notably in lateral positions as inferred for 217 Qianodus. The latter's possession of compound whorls is indicative of the 218 competence of tooth rows to fuse into complex units and exhibit alternate patterning 219 during ontogeny at lateral positions of the dentition. It remains to be determined 220 221 whether similar offsetting occurs between adjacent single-tooth-row whorls in climatiids (e.g. *Ptomacanthus*¹⁷) and, potentially, in other more crown-ward²⁴ stem 222 chondrichthyans (Doliodus³²). 223

The presence of tooth whorls in the lower Silurian extends the minimum age for the origin of vertebrate jaws and dentitions back by approximately 13 million years (Extended Data Fig. 4) to close to the Ordovician–Silurian boundary. In addition,

Qianodus partially infills and explains a gap in the fossil record between putative 227 gnathostome scales and fin spines from the Upper Ordovician–lower Silurian^{11,12,26,48} 228 and the appearance of teeth in the upper Silurian^{6,7,21,22}. The paucity of teeth and 229 dental elements recorded through this interval may result from their extremely low 230 abundance (as evident from the Rongxi Formation collections - see Methods), 231 coupled with the challenges in differentiating between isolated oral and extraoral 232 elements of unconventional morphologies^{3,5,24,39}. *Qianodus* presents rare, and some 233 of the clearest to date, evidence supporting a proposed^{11,12} appearance of jawed 234 235 vertebrates during the Great Ordovician Biodiversification Event (c. 485 to 443 Ma⁴⁹). Our phylogenetic analysis places *Qianodus* in the chondrichthyan total group, 236 with the implication that a range of jawed fishes appeared in the Upper Ordovician 237 and lower Silurian and co-existed alongside jawless vertebrates shortly after the 238 inception of biomineralization in the gnathostome total group. 239



- Fig. 1 | *Qianodus duplicis* tooth whorls and their position in the Rongxi
- 244 Formation exposed at Leijiatun (Shiqian-Tunping section), Guizhou Province,
- **China. (a)** Chronostratigraphy and lithostratigraphy of the Shiqian-Tunping section
- and (**b**) a map showing its location. (**c** to **f**) Volume renderings of synchrotron X-ray
- tomography data depicting the holotype of *Qianodus* IVPP V26641 in (c) distal, (d)
- labial, (e) basal and (f) mesial lingual views. at, accessory teeth; la, labial; li, lingual;
- 249 pt, primary teeth. Scale bars, 0.5 mm.



Fig. 2 | Tooth patterning and tissue structure of *Qianodus duplicis* whorls. 251 Volume renderings of X-ray (a, f) synchrotron and (b) microcomputed tomography 252 data, (c) scanning electron microscopy and (d) Nomarski DIC optics microscopy. 253 Oral views of whorls with (a) left (holotype IVPP V26641) and (b) right (IVPP 254 V26647) progenitor tooth families. Black horizontal lines in (a) indicate the alignment 255 between the accessory tooth rows located closest to the primary teeth. (c) Mesial 256 view of a whorl specimen (IVPP V26649) preserving evidence for the position of all 257 primary and accessory teeth (black lines trace the course of accessory tooth rows 1 258 and 2). (d) Longitudinal thin section of a tooth whorl IVPP V26644. (e) Summary 259 diagram of tooth arrangement and growth pattern of Qianodus whorls. (f) Vascular 260 system of the holotype IVPP V26641 shown in labial distal view (purple, primary 261 teeth; yellow, accessory teeth). ct, compact tissue; di, distal; la, labial; li, lingual; m, 262 mesial; p, tooth pulp; st, spongiose tissue. Scale bars, 0.5 mm. 263



Fig. 3 | Examples of the earliest chondrichthyan and osteichthyan tooth

whorls. The diagram illustrates oral views of tooth whorl morphologies from the

267 Silurian–Middle Devonian and their position in lower jaw dentitions. Dashed-line

268 margins of dentition symbols indicate unknown elements. Question mark next to

269 *Qianodus* denotes indirect support for proposed dentition patterning. Presence of

- 270 parasymphyseal whorls in *Guiyu* is indirectly supported by possession of jaw
- articulations surfaces for parasymphyseal plates⁷. See Supplementary information

for details about the line art and data presented in the figure.



Extended Data Fig. 1 | Morphology of Qianodus tooth whorls. (a-c, i, j, m) Scanning 274 electron microscopy, volume renderings of (g, h) synchrotron and (d-f) microcomputed X-275 ray tomography datasets and (k, l, n) light microscopy. (a) Lateral (mesial or distal) view of a 276 heavily abraded tooth whorl (IVPP V26648). (b) Mesial view of a tooth whorl (IVPP V26649). 277 (c) Latero-posterior view of a tooth whorl (IVPP V26652). (d-f) Tooth whorl (IVPP V26647) 278 in (d) labial, (e) mesial and (f) basal views. (g, h) Incomplete tooth (IVPP V26645) whorl in 279 (g) mesial and (h) basal views. (i) Lateral view of a tooth whorl (IVPP V26654) with a flared-280 out base. (j–n) Two complete whorls with 6 recognizable primary teeth in (j, k) lateral (distal 281 and mesial) (IVPP V26650), (m) oral (IVPP V26651) and (I, n) basal (IVPP V26650, 51) 282 283 views. at, accessory teeth; la, labial, li, lingual; pt, primary teeth. Scale bars, 0.5 mm.



Extended Data Fig. 2 | Internal structure of Qianodus tooth whorls. (a, g) Nomarski DIC 285 286 optical microscopy and (**b**–**f**, **h**) volume renderings of synchrotron X-ray tomography datasets. (a) Longitudinal thin section through a whorl with partially preserved teeth IVPP 287 V26653. (b) Longitudinal virtual slice through the progenitor tooth row of the holotype IVPP 288 V26641. (c) Horizontal virtual slice through the holotype IVPP V26641 at the level of tooth. 289 (d) Transverse virtual slice through a partially preserved whorl IVPP V26645. (e) Basal view 290 291 of IVPP V26641 with highlighted compact tissue of the base. (f) Volume rendering of radiotransparent structures inside a tooth whorl fragment (IVPP V26646) shown in oral view. 292 (g) Longitudinal thin section and (h) longitudinal virtual section through IVPP V26646. at, 293 294 accessory teeth; ct, compact tissue; la, labial; li, lingual; p, tooth pulp; pt, primary teeth; st, spongiose tissue; stc, spongiose tissue canals; wbc, whorl base crest. Scale bars, 0.5 mm. 295



- 297 Extended Data Fig. 3 | Comparison of *Qianodus* tooth whorls with the whorl-based
- dentition of the stem chondrichthyan *Doliodus problematicus*. (a) *Qianodus* tooth
 whorls at a late stage of development (from top to bottom IVPP V26652, V26641, V26649,
- V26647, IV26655 and V26648). (b) Tooth whorls of the lower left jaw ramus of *Doliodus* at
- 301 positions 1 to 9 (P1–9) (adapted from Maisey et al.³²).



Extended Data Fig. 4 | **Phylogenetic position of** *Qianodus* **within early jawed vertebrates.** 50

percent majority-rule consensus tree from a parsimony analysis of 105 taxa and 294 characters. Tree
 time-adjusted using minimum branch length scaling. Taxon and tree root ages sourced from King et
 al.⁵⁰ and other studies (see Extended Data Table 1). Colour coding of cladogram branches: jawless

307 stem gnathostomes (purple), 'placoderms' (black), Osteichthyes (green), stem Chondrichthyes

308 (ochre), crown Chondrichthyes (blue). Pie charts represent Markov k-state 1 likelihood values for

tooth whorl/dentition characters at select internal nodes. Circles show character states at terminal
 nodes. Character numbers shown in parentheses.



Extended Data Fig. 5 | Results of the parsimony analysis described in the Methods section and
 in Extended Data Fig. 4. (a) 50% majority-rule consensus and (b) strict consensus tree topologies.
 Squares in (a) depict most-parsimonious character state reconstructions at select internal nodes
 (character numbers shown in parentheses). Numbers at internal branches represent bootstrap values

of 50 percent and above.

318 **Online content**

- Any methods, additional references, Nature Research reporting summaries, source
- 320 data, extended data, supplementary information, acknowledgements, peer review
- information; details of author contributions and competing interests; and statements
- of data and code availability are available at https://doi.org/XXXXX
- Donoghue, P. C. & Rücklin, M. The ins and outs of the evolutionary origin of teeth.
 Evol. Dev. 18, 19-30 (2016).
 Smith, M. M. Vertebrate dentitions at the origin of jaws: when and how pattern
 evolved. Evol. Dev. 5, 394-413 (2003).
- 327 3 Vaškaninová, V. *et al.* Marginal dentition and multiple dermal jawbones as the 328 ancestral condition of jawed vertebrates. *Science* **369**, 211-216 (2020).
- Chen, D., Blom, H., Sanchez, S., Tafforeau, P. & Ahlberg, P. E. The stem
 osteichthyan *Andreolepis* and the origin of tooth replacement. *Nature* 539, 237-241
 (2016).
- 5 Chen, D. *et al.* Development of cyclic shedding teeth from semi-shedding teeth: the
 inner dental arcade of the stem osteichthyan *Lophosteus*. *R. Soc. Open Sci.* 4,
 161084 (2017).
- Choo, B., Zhu, M., Zhao, W. & Jia, L. The largest Silurian vertebrate and its palaeoecological implications. *Sci. Rep.* **4**, 5242 (2014).
- Zhu, M. *et al.* The oldest articulated osteichthyan reveals mosaic gnathostome
 characters. *Nature* 458, 469-474 (2009).
- 339 8 Denison, R. H. *Placodermi*. Vol. 2 (Gustav Fischer Verlag, 1978).
- Ginter, M., Hampe, O., Duffin, C. J. & Schultze, H. Handbook of Paleoichthyology
 Volume 3D. Chondrichthyes. Paleozoic Elasmobranchii: Teeth (Verlag Dr Friedrich
 Pfeil, 2010).
- Rücklin, M. *et al.* Development of teeth and jaws in the earliest jawed vertebrates.
 Nature 491, 748-751 (2012).
- Brazeau, M. D. & Friedman, M. The origin and early phylogenetic history of jawed
 vertebrates. *Nature* 520, 490-497 (2015).
- Sansom, I. J. & Andreev, P. S. in *Evolution and Development of Fishes* (eds
 Johanson, Z., Underwood, C. & Richter, M.) 59-70 (Cambridge Univ. Press, 2019).
- Thanh, T.-D., Phuong, T. H., Boucot, A. J., Goujet, D. & Janvier, P. Silurian
 vertebrates from Central Vietnam. *C. R. Acad. Sci. Series IIA* 12, 1023-1030 (1997).
- 351 14
 352 Zhao, W.-J. *et al.* A review of Silurian fishes from north-western Hunan, China and
 352 related biostratigraphy. *Acta Geol. Pol.* 68 (2018).
- 353 15 Zhu, M. *et al.* A Silurian maxillate placoderm illuminates jaw evolution. *Science* 354, 334-336 (2016).
- Burrow, C. J. & Rudkin, D. Oldest near-complete acanthodian: the first vertebrate
 from the Silurian Bertie Formation Konservat-Lagerstätte, Ontario. *PLoS One* 9,
 e104171 (2014).
- Brazeau, M. D. A revision of the anatomy of the Early Devonian jawed vertebrate
 Ptomacanthus anglicus Miles. *Palaeontology* 55, 355-367 (2012).
- Burrow, C. J., Davidson, R. G., Den Blaauwen, J. L. & Newman, M. J. Revision of
 Climatius reticulatus Agassiz, 1844 (Acanthodii, Climatiidae), from the Lower
 Devonian of Scotland, based on new histological and morphological data. J. Vertebr.
 Paleont. 35, e913421 (2015).

364 19 Burrow, C. J., Newman, M., Den Blaauwen, J., Jones, R. & Davidson, R. The Early Devonian ischnacanthiform acanthodian Ischnacanthus gracilis (Egerton, 1861) from 365 the Midland Valley of Scotland. Acta Geol. Pol. 68, 335-362 (2018). 366 367 20 Maisey, J. et al. in Evolution and development of fishes (eds Johanson, Z., Underwood, C. & Richter, M.) 87-109 (Cambridge Univ. Press, 2019). 368 21 Burrow, C. J. & Simpson, A. J. A new ischnacanthid acanthodian from the Late 369 Silurian (Ludlow, ploeckensis Zone) Jack Formation, north Queensland. Mem. 370 Queensl. Mus. 38, 383-396 (1995). 371 372 22 Gross, W. Mundzähne und hautzähne der acanthodier und arthrodiren. Palaeontogr. Abt. A, 1-40 (1957). 373 Martínez-Pérez, C. et al. Vascular structure of the earliest shark teeth. Acta Geol. 374 23 375 Pol. 68, 335-362 (2018). 24 Coates, M. I. et al. An early chondrichthyan and the evolutionary assembly of a shark 376 body plan. Proc. R. Soc B 285, 20172418 (2018). 377 Andreev, P. S. et al. The systematics of the Mongolepidida (Chondrichthyes) and the 378 25 Ordovician origins of the clade. PeerJ 4, e1850 (2016). 379 Andreev, P. S. et al. Early Silurian chondrichthyans from the Tarim Basin (Xinjiang, 380 26 China). PLoS One 15, e0228589 (2020). 381 Keating, J. N., Marquart, C. L. & Donoghue, P. C. Histology of the heterostracan 27 382 dermal skeleton: insight into the origin of the vertebrate mineralised skeleton. J. 383 384 Morph. 276, 657-680 (2015). 385 28 Dearden, R. P., Stockey, C. & Brazeau, M. D. The pharynx of the stem-386 chondrichthyan Ptomacanthus and the early evolution of the gnathostome gill skeleton. Nat. Commun. 10, 1-7 (2019). 387 388 29 Gegenbaur, C. Grundriss der vergleichenden Anatomie (Wilhelm Engelmann, 1874). Huxley, T. H. On the application of the laws of evolution to the arrangement of the 30 389 Vertebrata, and more particularly of the Mammalia. Proc. Sci. Meetings Zool. Soc. 390 391 Lond. 1880, 649-662 (1880). Rücklin, M. et al. Acanthodian dental development and the origin of gnathostome 392 31 dentitions. Nat. Ecol. Evol. 5, 919-926 (2021). 393 394 32 Maisey, J. G., Turner, S., Naylor, G. J. & Miller, R. F. Dental patterning in the earliest sharks: Implications for tooth evolution. J. Morph. 275, 586-596 (2014). 395 Underwood, C., Johanson, Z. & Smith, M. M. Cutting blade dentitions in squaliform 396 33 sharks form by modification of inherited alternate tooth ordering patterns. R. Soc. 397 Open Sci. 3, 160385 (2016). 398 Underwood, C. J. et al. Development and evolution of dentition pattern and tooth 399 34 400 order in the skates and rays (Batoidea; Chondrichthyes). PLoS One 10, e0122553 (2015). 401 Smith, M. M. & Coates, M. I. in Major Events in Early Vertebrate Evolution (ed 402 35 Ahlberg, P. E.) 223-240 (Taylor & Francis, 2001). 403 Coates, M. & Sequeira, S. A new stethacanthid chondrichthyan from the Lower 404 36 Carboniferous of Bearsden, Scotland. J. Vertebr. Paleontol. 21, 438-459 (2001). 405 Zangerl, R. & Case, G. Cobelodus aculeatus (Cope), an anacanthous shark from 406 37 Pennsylvanian black shales of North America. Palaeontogr. Abt. A 154, 107-157 407 408 (1976). 409 38 Andrews, M., Long, J., Ahlberg, P., Barwick, R. & Campbell, K. The structure of the sarcopterygian Onychodus jandemarrai n. sp. from Gogo, Western Australia: with a 410 functional interpretation of the skeleton. Earth. Env. Sci. Trans. R. Soc. Edinb. 96, 411 412 197-307 (2005). Blais, S. A., MacKenzie, L. A. & Wilson, M. V. Tooth-like scales in Early Devonian 39 413 eugnathostomes and the 'outside-in' hypothesis for the origins of teeth in vertebrates. 414 J. Vertebr. Paleont. 31, 1189-1199 (2011). 415 40 Frey, L. et al. The early elasmobranch Phoebodus: phylogenetic relationships, 416 ecomorphology and a new time-scale for shark evolution. Proc. R. Soc. B 286, 417 20191336 (2019). 418

419 41 Hu, Y., Lu, J. & Young, G. C. New findings in a 400 million-year-old Devonian placoderm shed light on jaw structure and function in basal gnathostomes. Sci. Rep. 420 421 7, 7813 (2017). Hu, Y.-Z., Young, G., Burrow, C., Zhu, Y.-a. & Lu, J. High resolution XCT scanning 422 42 reveals complex morphology of gnathal elements in an Early Devonian arthrodire. 423 Palaeoworld 28, 525-534 (2019). 424 Doeland, M., Couzens, A. M., Donoghue, P. C. & Rücklin, M. Tooth replacement in 425 43 early sarcopterygians. R. Soc. Open Sci. 6, 191173 (2019). 426 427 44 Zhu, M. & Yu, X. in: Recent Advances in the Origin and Early Radiation of Vertebrates (eds Arratia, G., Wilson, M. V. H. & Cloutier, R.) 271-286 (Verlag Dr. 428 Friedrich Pfeil, 2004). 429 430 45 Burrow, C. J., Newman, M. J., Davidson, R. G. & den Blaauwen, J. L. Redescription of Parexus recurvus, an Early Devonian acanthodian from the Midland Valley of 431 Scotland. Alcheringa 37, 392-414 (2013). 432 Dearden, R. P. & Giles, S. Diverse stem-chondrichthyan oral structures and evidence 433 46 for an independently acquired acanthodid dentition. R. Soc. Open Sci. 8, 210822 434 435 (2021). 47 Zangerl, R. & Case, G. R. Iniopterygia, a New Order of Chondrichthyan Fishes from 436 437 the Pennsylvanian of North America (Field Mus. Nat. Hist., 1973). 438 48 Andreev, P. S. et al. Upper Ordovician chondrichthyan-like scales from North America. Palaeontology 58, 691-704 (2015). 439 440 49 Servais, T. & Harper, D. A. The great Ordovician biodiversification event (GOBE): 441 definition, concept and duration, Lethaia 51, 151-164 (2018), King, B., Qiao, T., Lee, M. S., Zhu, M. & Long, J. A. Bayesian morphological clock 50 442 methods resurrect placoderm monophyly and reveal rapid early evolution in jawed 443 vertebrates. Syst. Biol. 66, 499-516 (2017). 444 445

446 Methods

- 447 A total of 23 *Qianodus* tooth whorls of varying degrees of completeness were
- recovered from residues of Rongxi sample 35SQTP following disaggregation of c.
- 300 kg of sediment with buffered 8% acetic acid. The *Qianodus* specimens are
- among thousands of vertebrate microremains isolated from 35SQTP that await
- 451 future investigation.

452

453 X-ray tomography

- 454 Synchrotron X-ray tomography analyses of three whorls (IVPP V26641
- 455 (Supplementary Data 3), V26645 (Supplementary Data 4), V26646)) were performed
- 456 at BL01A1 and BL01B1 beamlines of the Taiwan Light Source (TLS), National

457 Synchrotron Radiation Research Center (NSRRC), Taiwan. Acquisition at \geq 4 keV 458 with a parallel semi-white-light hard X-ray beam over a 180° rotation arc generated 459 datasets of 601 radiographs with pixel size of 2.76 µm. Post-acquisition, radiograph 460 alignment was enhanced in Matlab R2014b by using the fast projection matching 461 (Faproma) algorithm developed by Wang 2020⁵¹. Reconstructions of the radiograph 462 data in VGSTUDIO MAX 3.0 produced sets of 1200 tomographic slices (1600 x 1600 463 pixels) per specimen.

One tooth whorl (IVPP V26647) was imaged at the Institute of Vertebrate 464 Paleontology and Paleoanthropology, Chinese Academy of Sciences with an X-ray 465 micro-computed tomography scanner (225-3D-µCT) designed by the Institute of 466 High Energy Physics, Chinese Academy of Sciences ⁵². The analysis produced 720 467 radiographs over a 360-degree rotation cycle that were converted in VGSTUDIO 468 MAX 3.0 to a dataset of 1442 tomograms (each 1748 x 556 pixels at 5.33 µm per 469 pixel). Three-dimensional visualization of the tomogram data from IVPP V26641 and 470 IVPP V26645–V26647 was performed in Mimics 19.0 (Fig. 2a, f and Extended Data 471 Fig. 2e, f, h), VGSTUDIO MAX 3.0 (Extended Data Fig. 2b-d) and in Drishti 2.6.5 (Fig. 472 1c-f and Extended Data Fig. 1d-h) from Mimics segmentation masks exported as 473 DICOM files. 474

475

476 Scanning electron microscopy

477 Surface morphology of seven uncoated whorls (IVPP V26642, V26643, V26648–

478 V26652) was documented with a Phenom ProX Desktop SEM at 5 keV at the School

of Geography, Earth and Environmental Sciences, University of Birmingham.

480

481 Light microscopy

Doubly-polished sections of seven specimens (IVPP V26644, V26646, V26653–
V26657) were investigated under DIC polarized light with an Olympus BX51
Fluorescence Microscope and documented with an Olympus D12 digital camera at
Qujing Normal University, China.

Two tooth whorls (IVPP V26650, V26651) were imaged with a GXMXTL-3101
stereo microscope at the University of Birmingham, UK.

488

489 **Phylogenetic analysis**

TNT version 1.5⁵³ was used to perform a parsimony phylogenetic analysis based on 490 a matrix of 294 characters and 105 taxa (Supplementary Data 6). The matrix was 491 assembled from character data from Brazeau et al.⁵⁴, Coates et al.²⁴, Dearden⁵⁵, 492 Dearden et al.²⁸, Dearden and Giles⁴⁶, Giles et al.⁵⁶, King et al.⁵⁰, Qiao et al.⁵⁷, Zhu 493 et al.¹⁵, Zhu et al.⁵⁸ and this study (character scores colour coded in the matrix nexus 494 file, Supplementary Data 6). Characters were treated as unordered and of equal 495 weights with Galeaspida being designated as an outgroup. Tree reconstruction was 496 performed in TNT by a heuristic analysis using the mult search algorithm with tree-497 bisection-reconnection (TBR) branch swapping set to save 100 trees per replication 498 and limited to 100 random addition-sequences. The search was set to retain 100000 499 trees and returned 100000 most parsimonious trees (940 steps each) from which 500 50% majority-rule consensus (944 steps) and strict consensus trees (1022 steps) 501 were calculated (Supplementary Data 6). 502

503 Bootstrap support for internal tree nodes of the 50 percent majority-rule 504 consensus and strict consensus trees was calculated in TNT via a traditional search 505 method over 100 bootstrap replicates.

The 50 percent majority-rule consensus tree and its branch length values was imported into R (version 4.0.2) and time adjusted with the R package paleotree 3.3.25⁵⁹. This analysis used the timePaleoPhy function of paleotree with mbl-type time scaling and pre-assigned taxon and tree root ages taken from King et al.⁵⁰ and other studies (Table 1).

511 Maximum likelihood scores for characters at internal nodes of the 50 percent 512 majority-rule consensus tree were produced in Mesquite 3.51 (build 898) using a 513 Markov k-state¹¹ probability model that assumes equal rates of character change. 514

515 **Dentition data**

Figure 3 dentition data and line art from this study and adapted from Andrews³⁸ 516 (Onychodus jandemarrai), Botella et al.⁶⁰ (Obruchevacanthus ireneae), Burrow et 517 al.¹⁸ (Climatius reticulatus), Burrow et al.⁴⁵ (Parexus recurvus), Burrow et al.¹⁹ 518 (Ischnacanthus gracilis), Burrow and Simpson²¹ (Gomphonchus? turnerae), Gagnier 519 and Wilson⁶¹ (*Brochoadmones milesi*), Gross²² (*Gomphodus*), Jarvik⁶² (*Holoptychius*) 520 sp.), Jarvik⁶² (?Porolepis), Maisey et al.³² (Doliodus problematicus), Long 1988⁶³ 521 (Howqualepis rostridens), Miles⁶⁴ (Ptomacanthus anglicus), Mondéjar-Fernández 522 et al.⁶⁵ (*Durialepis edentatus*), Qu et al.⁶⁶ (unidentified acanthodian), Vergoossen⁶⁷ 523 (ischnacanthid), Wang⁶⁸ (*Gomphonchus liujingensis*) and Zhu et al.⁷ (*Guiyu oneiros*). 524 525

526 Data availability

- 527 Supplementary files (Supplementary Data 1 to 6) are available at
- 528 https://www.dropbox.com/sh/eor2wvwwbui25sk/AACckwBWZ9vyX5ouSdJ1aq27a?dl
- 529 =0 and will be published online in a publicly accessible repository (Dryad) upon
- acceptance of the manuscript. Investigated *Qianodus* specimens were assigned
- accession numbers (IVPP V26641–V26663) and deposited at the Institute of
- 532 Vertebrate Paleontology and Paleoanthropology (IVPP), Chinese Academy of
- 533 Sciences, Beijing.
- 53451Wang, C.-C. Joint iterative fast projection Matching for fully Automatic Marker-free535Alignment of nano-tomography Reconstructions. Sci. Rep. 10, 7330 (2020).
- 53652Wang, Y. et al. Development and applications of paleontological computed537tomography. Vertebr. PalAsiat. 57, 84-92 (2019).
- 538 53 Goloboff, P. A. & Catalano, S. A. TNT version 1.5, including a full implementation of 539 phylogenetic morphometrics. *Cladistics* **32**, 221-238 (2016).
- 54054Brazeau, M. *et al.* Endochondral bone in an Early Devonian 'placoderm' from541Mongolia. *Nat. Ecol. Evol.*, **4**, 1477-1484 (2020).
- 542 55 Dearden, R. P. *The Anatomy and Evolution of "Acanthodian" Stem-chondrichthyans* 543 (Imperial College London, 2018).
- 54456Giles, S., Friedman, M. & Brazeau, M. D. Osteichthyan-like cranial conditions in an545Early Devonian stem gnathostome. *Nature* **520**, 82-85 (2015).
- 546 57 Qiao, T., King, B., Long, J. A., Ahlberg, P. E. & Zhu, M. Early gnathostome
 547 phylogeny revisited: multiple method consensus. *PLoS One* **11**, e0163157 (2016).
 548 58 Zhu, Y.-a., Lu, J. & Zhu, M. Reappraisal of the Silurian placoderm Silurolepis and
- 549 insights into the dermal neck joint evolution. *R. Soc. Open Sci.* **6**, 191181 (2019).
- 550 59 Bapst, D. W. paleotree: an R package for paleontological and phylogenetic analyses 551 of evolution. *Methods Ecol. Evol.* **3**, 803-807 (2012).
- Botella, H., Manzanares, E., Ferrón, H. & Martínez-Pérez, C. *Obruchevacanthus ireneae* gen. et sp. nov., a new ischnacanthiform (Acanthodii) from the Lower
 Devonian of Spain. *Paleontol. J.* 48, 1067-1076 (2014).
- 555 61 Gagnier, P.-Y. & Wilson, M. V. An unusual acanthodian from northern Canada: 556 revision of *Brochoadmones milesi. Mod. Geol.* **20**, 235-252 (1996).
- 557 62 Jarvik, E. Middle and Upper Devonian Porolepiformes from East Greenland with 558 special reference to *Glyptolepis groenlandica* n. sp. and a discussion on the structure 559 of the head in the Porolepiformes. *Medd. Grønl.* **187**, 1-307 (1972).
- Long, J. New palaeoniscoid fishes from the Late Devonian and Early Carboniferous
 of Victoria. *Mem. Assoc. Australas. Palaeontol.* 7, 1-64 (1988).
- Miles, R. S. Articulated acanthodian fishes from the Old Red Sandstone of England:
 with a review of the structure and evolution of the acanthodian shoulder-girdle. *Bull. Br. Mu. Nat. Hist. Geol.* 24, 111-213 (1973).
- 565 65 Mondéjar-Fernández, J., Friedman, M. & Giles, S. Redescription of the cranial 566 skeleton of the Early Devonian (Emsian) sarcopterygian *Durialepis edentatus* Otto (Dippomorpha, Porolepiformes), *Pap. Palaeontol.* (2020)
- (Dipnomorpha, Porolepiformes). Pap. Palaeontol. (2020).
 (Qu, Q., Sanchez, S., Blom, H., Tafforeau, P. & Ahlberg, P. E. Scales and tooth
 whorls of ancient fishes challenge distinction between external and oral 'teeth'. PLoS
 One 8, e71890 (2013).
- 571 67 Vergoossen, J. Late Silurian fish microfossils from Helvetesgraven, Skåne (southern 572 Sweden)(I). *Geol. Mij.* **78**, 267-280 (1999).

68 Wang, N.-Z. Microremains of agnathans and fishes from Lower Devonian of central Guangxi with correlation of Lower Devonian between central Guangxi and eastern Yunnan, South China. *Acta Palaeontol. Sin.* **31**, 280-303 (1992).

576

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596 Supplementary Information for:

597 The oldest gnathostome teeth

598

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612 Geological setting and biostratigraphy of the Rongxi

The Rongxi Formation forms part of the laterally extensive and loosely defined 'lower 613 marine red beds' that form a distinctive unit in South China. The Shigian-Tunping 614 (SQTP) section through the Rongxi is located N of Leijiatun village in Shiqian 615 County, Guizhou Province (Fig. 1b). Here the unit is 242 m thick and consists of 616 purple-red and grey mudstones interbedded with sporadic sandstone and carbonate 617 units. Sample 35SQTP was collected from a c. 90 cm sequence of massive 618 conglomerates alternating with mudstones within the upper half of the Rongxi 619 Formation (Fig. 1a). The 35SQTP limestone conglomerate consists of well-rounded 620 mudstone clasts (up to pebble size, c. 2 cm) and shell fragments/vertebrate remains 621 622 bound by calcite cement. Previous paleoenvironmental interpretations of the Rongxi have concluded that it represents a shallow water intertidal depositional tract, 623 referable to Benthic Assemblage Zone 1¹⁻³, and in this context the conglomerate 624 dominated lens is readily recognized as a tidal channel deposit. 625 Age diagnostic fossils from the Rongxi outcrop at Leijiatun have largely been lacking, 626 with invertebrate fossils being notably rare⁴ and the age of the formation at Leijiatun 627 and elsewhere has primarily been based upon the biostratigraphy of the underlying 628 and overlying units. In part, this has led to the Rongxi being assigned to the late 629 Aeronian⁵⁻⁷, straddling the Aeronian-Telychian boundary, or the lower Telychian^{3,8,9}. 630 Sample 35SQTP has yielded the first conodonts to be recovered from the Rongxi 631 and include platform elements of Ozarkodina guizhouensis¹⁰, an index taxon for the 632 Ozarkodina guizhouensis Biozone that covers the Aeronian-Telychian stage 633 boundary. However, Wang et al.⁷ and Wang⁵ considered the conodont and graptolite 634 evidence from the overlying Xiushan Formation to place the Rongxi in the Aeronian 635

regard it highly probable that the Lower Red Beds are diachronous throughout their 637 1000 kms of expression, with the outcrops in Guizhou, Hunan and Chongqing being 638 older than the basal Telychian Formations in Sichuan and Shaanxi^{3,8}. Thus, an 639 Aeronian age for the Rongxi Formation at Leijiatun seems the most likely. 640 641 Character list for the phylogenetic analysis 642 Abbreviations of character sources: [B] Brazeau et al.¹¹, [C] Coates et al.¹², [D] 643 Dearden et al.¹³, [G] Giles et al.¹⁴ and [K] King et al.¹⁵. 644 645 1. [B:1] Tessellate prismatic calcified cartilage. 646 0 absent 647 1 present 648 649 2. [B:2] Prismatic calcified cartilage. 650 0 single layered 651 1 multi-layered 652 653 654 3. [B:3] Perichondral bone. 0 present 655 1 absent 656 657 4. [B:4] Extensive endochondral ossification. 658 0 absent 659 1 present 660 661 [B:5] Enamel(oid) present on dermal bones and scales. 5. 662

within the Ozarkodina parahassi division of the O. guizhouensis Biozone. We also

663	0 absent				
664	1 present				
665					
666	6.	[B:6] Enamel.			
667	0 single-lay	rered			
668	1 multi-laye	ered			
669					
670	7.	[B:7] Enamel layers.			
671	0 applied d	irectly to one another (ganoine)			
672	1 separated	d by layers of dentine			
673					
674	8.	[B:8] Pore canal network.			
675	0 absent				
676	1 present				
677					
678	9.	[B:9] Dentinous tissue.			
679	0 absent				
680	1 present				
681					
682	10.	[B:10] Dentine kind (modified).			
683	0 mesoden	tine			
684	1 semidentine				
685	2 orthodent	ine			
686					
687	11.	[B:11] Bone cell lacunae in trunk scale bases.			
688	0 present				
689	1 absent				
690					
691	12.	[B:12] Main dentinous tissue forming fin spine.			
692	0 osteoden	tine			

693	1 orthodentine			
694				
695	13.	[B:13] Longitudinal scale alignment in fin webs.		
696	0 absent			
697	1 present			
698				
699	14.	[B:14] Differentiated lepidotrichia.		
700	0 absent			
701	1 present			
702				
703	15.	[B:15] Composition of trunk scale crowns (modified).		
704 705 706	Character 1 odontode so patterning.	5 of Brazeau et al. ¹¹ was reformulated to describe mono- and poly- cale crowns on the basis of odontode number without alluding to their		
707	0 comprisino	g single odontode unit/generation ("monodontode")		
708	1 comprisino	g a complex of multiple odontode generations/units ("polyodontode")		
709				
710	16.	[B:16] Concentric addition of trunk scale odontodes (modified).		
711 712 713	Modified cha by specifying generations	aracter 16 of Brazeau et al. ¹¹ distinguished from the original formulation g that the concentric growth of scales refers to their odontode		
714	0 absent			
715	1 present			
716				
717	17.	[B:17] Buried odontode generations (modified).		
718 719	We split character 17 of Brazeau et al. ¹¹ in order to code separately for overgrowth of odontode generations (character 17) and odontode resorption (character 289).			
720	0 present			
721	1 absent			
722				
723	18.	[B:18] Trunk scales with peg-and-socket articulation.		
724	0 absent			

1 present	
19.	[B:19] Scale peg.
0 broad	
1 narrow	
20.	[B:20] Anterodorsal process on scale.
0 absent	
1 present	
21.	[B:21] Trunk scale profile.
0 distinct cro	own and base demarcated by a constriction ("neck")
1 flattened	
22.	[B:22] Profile of scales with constriction between crown and base.
0 neck simil	ar in width to crown
1 neck grea	tly constricted, resulting in anvil-like shape
23.	[B:23] Trunk scales with bulging base.
0 absent	
1 present	
24.	[B:24] Trunk scales with flattened base.
0 present	
0 present 1 absent	
0 present 1 absent	
0 present 1 absent 25 .	[B:25] Basal pore in scales.
0 present 1 absent 25. 0 absent	[B:25] Basal pore in scales.
0 present 1 absent 25. 0 absent 1 present	[B:25] Basal pore in scales.
	 19. 0 broad 1 narrow 20. 0 absent 1 present 21. 0 distinct cross 1 flattened 22. 0 neck similar 1 neck great 23. 0 absent 1 present

26. 755 [B:26] Flank scale alignment. 0 vertical rows oblique rows or hexagonal 756 1 rhombic packing 757 2 disorganised 758 759 27. [B:27] Scute-like ridge scales (basal fulcra). 760 0 absent 761 1 present 762 763 28. [K:324] Sensory line canal. 764 0 passes between or beneath scales 765 1 passes over scales and/or is partially enclosed or surrounded by scales 766 2 perforates and passes through scales 767 768 29. [B:29] Dermal ornamentation. 769 0 smooth 770 1 parallel, vermiform ridges 771 2 concentric ridges 772 3 tuberculate 773 774 30. [B:30] Sensory line network. 775 0 preserved as open grooves (sulci) in dermal bones 776 1 sensory lines pass through canals in dermal bones (open as pores) 777 778 31. [B:31] Sensory canals/grooves. 779 0 contained within the thickness of dermal bones 780 1 contained in prominent ridges on visceral surface of bone 781 782 [B:32] Jugal portion of infraorbital canal joins supramaxillary 32. 783 canal. 784 0 present 785

786	1 absent		
787			
788	33.	[B:33] Dermal skull roof (modified).	
789	State 2 was added from character 25 of Coates el al. ¹²		
790	0 includes	large dermal plates	
791	1 consists	of undifferentiated plates, tesserae or scales	
792	2 naked or	largely scale free	
793			
794	34.	[B:34] Anterior pit line of dermal skull roof.	
795	0 absent		
796	1 present		
797			
798	35.	[B:35] Tessera morphology.	
799	0 large inte	rlocking polygonal plates	
800	1 microsqu	amose, not larger than trunk squamation	
801			
802	36.	[B:36] Cranial spines.	
803	0 absent		
804	1 present		
805			
806	37.	[B:37] Cranial spines.	
807	0 monocus	pid	
808	1 multicups	sid	
809			
810	38.	[B:38] Extent of dermatocranial cover.	
811	0 complete		
812	1 incomple	te (limited to skull roof)	
813			
814	39.	[B:39] Openings for endolymphatic ducts in dermal skull roof.	
815	0 present		
816	1 absent		
------------	---------------------------------------------------	---------------------------------------------------------------------------------	
817			
818 819	40. skull	[B:40] Endolymphatic ducts with oblique course through dermal bones.	
820	0 absent		
821	1 present		
822			
823 824	41. (i.e. r	[B:41] Endolymphatic duct relationship to median skull roof bone nuchal plate).	
825	0 within me	dian bone	
826	1 on bones	flanking the median bone (e.g. paranuchals)	
827			
828	42.	[B:42] Pineal opening perforation in dermal skull roof.	
829	0 present		
830	1 absent		
831			
832	43.	[B:43] Dermal plate associated with pineal eminence or foramen.	
833 834	0 contribute bones.)	es to orbital margin (plate(s) excluded from orbital margin by skull roofing	
835	1 plate bordered laterally by skull roofing bones		
836			
837	44.	[B:44] Broad supraorbital vaults.	
838	0 absent		
839	1 present		
840			
841	45.	[B:45] Median commissure between supraorbital sensory lines.	
842	0 absent		
843	1 present		
844			
845	46.	[B:46] Dermal cranial joint at level of sphenoid-otic junction.	
846	0 absent		

847	1 present	
848		
849	47.	[B:47] Otic canal extends through postparietals.
850	0 absent	
851	1 present	
852		
853	48.	[B:48] Number of bones of skull roof lateral to postparietals.
854	0 two	
855	1 one	
856	2 more than	i two
857		
858 859	49. placo	[B:49] Suture between paired skull roofing bones (centrals of derms postparietals of osteichthyans).
860	0 straight	
861	1 sinusoidal	
862		
863 864	50. corne	[B:50] Medial processes of paranuchal wrapping posterolateral ers of nuchal plate.
865	0 absent	
866	1 present	
867		
868	51.	[B:51] Paired pits on ventral surface of nuchal plate.
869	0 absent	
870	1 present	
871		
872	52.	[B:52] Sclerotic ring.
873	0 absent	
874	1 present	
875		
876	53.	[B:53] Consolidated cheek plates.
877	0 absent	

878	1 present	
879		
880	54.	[B:54] Cheek plate.
881	0 undivided	1
882	1 divided (i	.e., squamosal and preopercular)
883		
884	55.	[B:55] Subsquamosals in taxa with divided cheek.
885	0 absent	
886	1 present	
887		
888	56.	[B:56] Preopercular shape.
889	0 rhombic	
890	1 bar-shap	ed
891		
892 893	57. cana	[B:57] Vertical canal associated with preopercular/suborbital I.
894	0 absent	
895	1 present	
896		
897	58.	[B:58] Enlarged postorbital tessera separate from orbital series.
898	0 absent	
899	1 present	
900		
901	59.	[B:59] Extent of maxilla along cheek.
902	0 to posteri	or margin of cheek
903	1 cheek bo	nes exclude maxilla from posterior margin of cheek
904		
905	60.	[B:60] Dermal neck joint.
906	0 overlap	
907	1 ginglymo	id ('arthrodire'-type)
908	2 reverse g	inglymoid ('antiarch'-type)

909	3 longitudina	d d
910		
911	61.	[B:61] Sensory line scales/plates on head.
912	0 unspecializ	zed
913	1 apposed g	rowth
914	2 paralleling	canal
915	3 semicylind	rical C-shaped ring scales
916		
917	62.	[B:62] Bony hyoidean gill-cover series (branchiostegals).
918	0 absent	
919	1 present	
920		
921 922	63. jaw.	[B:63] Branchiostegal plate series along ventral margin of lower
923	0 absent	
924	1 present	
925		
926	64.	[B:64] Branchiostegal ossifications.
927	0 plate-like	
928	1 narrow and	l ribbon-like
929	2 filamentou	S
930		
931	65.	[B:65] Branchiostegal ossifications.
932	0 ornamente	d
933	1 unornamer	nted
934		
935	66.	[B:66] Imbricated branchiostegal ossifications.
936	0 absent	
937	1 present	
938		
939	67.	[B:67] Median gular.

940	0 absent	
941	1 present	
942		
943	68.	[B:68] Lateral gular.
944	0 absent	
945	1 present	
946		
947	69.	[B:69] Opercular (submarginal) ossification.
948	0 absent	
949	1 present	
950		
951	70.	[B:70] Shape of opercular (submarginal) ossification.
952	0 broad plat	te that tapers towards its proximal end
953	1 narrow, ro	od-shaped
954		
955	71.	[B:71] Size of lateral gular plates.
956	0 extending	most of length of the lower jaw
957 958	1 restricted to the anterior third of the jaw (no longer than the width of three or fou branchiostegals)	
959		
960	72.	[B:72] Gill arches.
961	0 largely res	stricted to region under braincase
962	1 extend far	posterior to braincase
963		
964	73.	[B:73] Basihyal.
965	0 absent	
966	1 present	
967		
968	74.	[B:74] Interhyal.
969	0 absent	
970	1 present	

971		
972	75.	[B:75] Hypohyal.
973	0 absent	
974	1 present	
975		
976	76.	[B:76] Endoskeletal urohyal.
977	0 absent	
978	1 present	
979		
980 981	77. of the	[B:77] Oral dermal tubercles borne on jaw cartilages or at margins e mouth.
982	0 absent	
983	1 present	
984		
985	78.	[B:79] Enamel(oid) on teeth.
986	0 absent	
987	1 present	
988		
989	79.	[B:80] Cap of enameloid restricted to upper part of teeth (acrodin).
990	0 absent	
991	1 present	
992		
993	80.	[B:81] Tooth whorls.
994	0 absent	
995	1 present	
996		
997	81.	[C:83] Tooth families/whorls
998	0 restricted	to symphysial region
999	1 distributed	d along jaw margin
1000		
1001	82.	[B:84] Distribution of tooth whorls.

1002	0 lower jaw	s only
1003	1 upper and lower jaws	
1004	2 upper jaw	/s only
1005		
1006	83.	[K:354] Parasymphyseal plate
1007	0 detachab	le tooth whorl
1008	1 long with	posterior corner, sutured to coronoid, denticulated or with tooth row
1009	2 absent	
1010		
1011	84.	[C:80] Bases of tooth families/whorls
1012	0 single, co	ntinuous plate
1013	1 some or a	all consist of separate tooth units
1014		
1015	85.	[B:85] Teeth ankylosed to dermal bones.
1016	0 absent	
1017	1 present	
1018		
1019	86.	[B:86] Plicidentine.
1020	0 absent	
1021	1 present	
1022		
1023	87.	[B:87] Dermal jaw plates on biting surface of jaw cartilages.
1024	0 absent	
1025	1 present	
1026		
1027	88.	[G:88] Maxillary and dentary marginal bones of mouth.
1028	0 absent	
1029	1 present	
1030		
1031	89.	[B:89] Premaxilla.

1032	0 extends under orbit		
1033	1 restricted anterior to orbit		
1034			
1035	90.	[B:90] Maxilla shape.	
1036	0 splint-sha	aped	
1037	1 cleaver-s	haped	
1038			
1039 1040	91. ethm	[B:91] Pair of tooth plates (anterior supragnathals or vomers) on noidal plate.	
1041	0 absent		
1042	1 present		
1043			
1044 1045	92. man	[K:353] Strong ascending flexion of symphysial region of dible.	
1046	0 absent		
1047	1 present		
1048			
1049	93.	[B:93] Extent of infradentaries.	
1050	0 along mu	ich of ventral margin of dentary	
1051	1 restricted	to posterior half of dentary	
1052			
1053	94.	[B:94] Coronoid fangs.	
1054	0 absent		
1055	1 present		
1056			
1057 1058	95. chee	[B:95] Position of upper mandibular arch cartilage (and associated k plate where present).	
1059	0 entirely s	uborbital	
1060	1 with a po	storbital extension	
1061			
1062	96.	[B:96] Position of mandibular arch articulations.	

1063	0 terminal		
1064	1 subtermin	al	
1065			
1066	97.	[B:97] Autopalatine and quadrate.	
1067	0 comineral	ized	
1068	1 separate r	mineralizations	
1069			
1070	98.	[B:98] Large otic process of the palatoquadrate.	
1071	0 absent		
1072	1 present		
1073			
1074	99.	[B:99] Insertion area for jaw adductor muscles on palatoquadrate.	
1075	0 ventral or	medial	
1076	1 lateral		
1077			
1078	100.	[B:100] Palatoquadrate fused with neurocranium.	
1079	0 absent		
1080	1 present		
1081			
1082 1083	101. palato	[B:101] Oblique ridge or groove along medial face of oquadrate.	
1084	0 absent		
1085	1 present		
1086			
1087 1088	102. articu	[B:102] Fenestration of palatoquadrate at basipterygoid lation.	
1089	0 absent		
1090	1 present		
1091			
1092 1093	103. portic	[B:103] Perforate or fenestrate anterodorsal (metapterygoid) on of palatoquadrate.	

1094	0 absent	
1095	1 present	
1096		
1097 1098	104. cartil	[B:104] Pronounced dorsal process on Meckelian bone or age.
1099	0 absent	
1100	1 present	
1101		
1102	105.	[B:105] Number of coronoids.
1103	0 four or mo	ore
1104	1 three or fe	ewer
1105		
1106	106.	[B:106] Preglenoid process.
1107	0 absent	
1108	1 present	
1109		
1110 1111	107. mano	[B:107] Jaw articulation located on rearmost extremity of dible.
1112	0 absent	
1113	1 present	
1114		
1115	108.	[B:108] Precerebral fontanelle.
1116	0 absent	
1117	1 present	
1118		
1119	109.	[B:109] Median dermal bone of palate (parasphenoid).
1120	0 absent	
1121	1 present	
1122		
1123	110.	[B:110] Parasphenoid.
1124	0 lozenge-s	haped

1125	1 splint-shaped		
1126	2 diamond-shaped		
1127			
1128	111.	[B:111] Multifid anterior margin of parasphenoid denticle plate.	
1129	0 absent		
1130	1 present		
1131			
1132	112.	[B:112] Enlarged ascending processes of parasphenoid.	
1133	0 absent		
1134	1 present		
1135			
1136	113.	[B:113] Buccohypophysial canal in parasphenoid.	
1137	0 single		
1138	1 paired		
1139			
1140	114.	[B:114] Nasal opening(s).	
1141	0 dorsal, pla	aced between orbits	
1142	1 ventral an	d anterior to orbit	
1143			
1144	115.	[B:115] External opening of posterior nostril and orbit.	
1145	0 separated	l by dermal bone(s)	
1146	1 confluent		
1147			
1148	116.	[B:116] Olfactory tracts.	
1149	0 short, with	n olfactory capsules situated close to telencephalon cavity	
1150	1 elongate a	and tubular (much longer than wide)	
1151			
1152 1153	117. neuro	[B:117] Prominent pre-orbital rostral expansion of the ocranium.	
1154	0 present, f	ormed of subethmoidal platform ('upper lip')	
1155	1 absent		

1156	2 present, formed of rhinocapsular block		
1157			
1158	118.	[B:118] Pronounced sub-ethmoidal keel.	
1159	0 absent		
1160	1 present		
1161			
1162	119.	[B:119] Internasal vacuities.	
1163	0 absent		
1164	1 present		
1165			
1166 1167	120. brain	[B:120] Discrete division of the ethmoid and more posterior case at the level of the optic tract canal.	
1168	0 absent		
1169	1 present		
1170			
1171	121.	[B:121] Position of myodome for superior oblique eye muscles.	
1172	0 posterior	and dorsal to foramen for nerve II	
1173	1 anterior a	nd dorsal to foramen	
1174			
1175	122.	[B:122] Endoskeletal intracranial joint.	
1176	0 absent		
1177	1 present		
1178			
1179	123.	[B:123] Spiracular groove on basicranial surface	
1180	0 absent		
1181	1 present		
1182			
1183	124.	[B:124] Transverse otic process.	
1184	0 present		
1185	1 absent		
1186			

1187	125.	[B:125] Jugular canal.
1188	0 long (inve	ested in otic region along length of skeletal labyrinth)
1189	1 short (res	tricted to short portion of region of skeletal labyrinth, or anterior to it)
1190	2 absent (ju	igular vein uninvested in otic region)
1191		
1192	126.	[B:126] Spiracular groove on lateral commissure.
1193	0 absent	
1194	1 present	
1195		
1196	127.	[B:127] Subpituitary fenestra.
1197	0 absent	
1198	1 present	
1199		
1200	128.	[B:128] Supraorbital shelf broad with convex lateral margin.
1201	0 absent	
1202	1 present	
1203		
1204	129.	[B:129] Orbit dorsal or facing dorsolaterally, surrounded laterally
1205	0 present	
1200	1 absent	
1207	raboont	
1200	130.	[B·130] Evestalk attachment area.
1210	0 absent	
1211	1 present	
1212	·	
1213	131.	[B:131] Postorbital process.
1214	0 absent	
1215	1 present	
1216		
1217	132.	[B:132] Canal for jugular in postorbital process.

1218	0 absent	
1219	1 present	
1220		
1221 1222	133. sens	[B:133] Series of perforations for innervation of supraorbital ory canal in supraorbital shelf.
1223	0 absent	
1224	1 present	
1225		
1226	134.	[B:134] Extended prehypophysial portion of sphenoid.
1227	0 absent	
1228	1 present	
1229		
1230 1231	135. along	[B:135] Narrow interorbital septum, with outer walls in contact g midline forming a single sheet.
1232	0 absent	
1233	1 present	
1234		
1235	136.	[B:136] The main trunk of facial nerve (N. VII).
1236	0 elongate	and passes anterolaterally through orbital floor
1237	1 stout, divides within otic capsule at the level of the transverse otic wall	
1238		
1239 1240	137. jugul	[B:137] Course of hyoid ramus of facial nerve (N. VII) relative to ar canal.
1241	0 traverses	jugular canal, with separate exit in otic region
1242	1 intersects	jugular canal, with exit through posterior jugular foramen
1243		
1244	138.	[B:138] Glossopharyngeal nerve (N. IX) exit.
1245	0 foramen s	situated posteroventral to otic capsule and anterior to metotic fissure
1246	1 through n	netotic fissure
1247		
1248	139.	[B:139] Relationship of cranial endocavity to basisphenoid.

1249	0 endocavity occupies full depth of sphenoid	
1250	1 enodcavity	/ dorsally restricted
1251		
1252	140.	[B:140] Subcranial ridges.
1253	0 absent	
1254	1 present	
1255		
1256 1257	141. caroti	[B:141] Ascending basisphenoid pillar pierced by common internal d.
1258	0 absent	
1259	1 present	
1260		
1261	142.	[B:142] Canal for lateral dorsal aorta within basicranial cartilage.
1262	0 absent	
1263	1 present	
1264		
1265	143.	[B:143] Entrance of internal carotids.
1266	0 through se	parate openings flanking the hypophyseal opening or recess
1267	1 through a	common opening at the central midline of the basicranium
1268		
1269 1270	144. basic	[B:144] Canal for efferent pseudobranchial artery within ranial cartilage.
1271	0 absent	
1272	1 present	
1273		
1274	145.	[B:145] Position of basal/basipterygoid articulation.
1275	0 same ante	roposterior level as hypophysial opening
1276	1 anterior to	hypophysial opening
1277	2 posterior to	o hypophysial opening
1278		

1279 1280	146. poste	[B:146] Articulation between neurocanium and palatoquadrate rodorsal to orbit (suprapterygoid articulation).
1281	0 absent	
1282	1 present	
1283		
1284	147.	[B:147] Labyrinth cavity.
1285 1286	0 separated wall	from the main neurocranial cavity by a cartilaginous or ossified capsular
1287	1 skeletal ca	apsular wall absent
1288		
1289 1290	148. orient	[B:148] Basipterygoid process (basal articulation) with vertically ted component.
1291	0 absent	
1292	1 present	
1293		
1294	149.	[B:149] Pituitary vein canal.
1295	0 dorsal to l	evel of basipterygoid process
1296	1 flanked posteriorly by basipterygoid process	
1297		
1298	150.	[B:150] External (horizontal) semicircular canal.
1299	0 absent	
1300	1 present	
1301		
1302	151.	[B:151] Sinus superior.
1303 1304	0 absent or saccular cha	indistinguishable from union of anterior and posterior canals with amber
1305	1 present	
1306		
1307	152.	[B:152] External (horizontal) semicircular canal.
1308	0 joins the v	estibular region dorsal to posterior ampulla
1309	1 joins level	with posterior ampulla
1310		

1311	153.	[B:153] Horizontal semicircular canal in dorsal view.
1312	0 medial to	path of jugular vein
1313	1 dorsal to j	jugular vein
1314		
1315	154.	[B:154] Lateral cranial canal.
1316	0 absent	
1317	1 present	
1318		
1319	155.	[B:155] Posterior dorsal fontanelle.
1320	0 absent	
1321	1 present	
1322		
1323	156.	[B:156] Shape of posterior dorsal fontanelle.
1324	0 approxima	ately as long as broad
1325	1 much long	ger than wide, slot-shaped
1326		
1327	157.	[B:157] Synotic tectum.
1328	0 absent	
1329	1 present	
1330		
1331	158.	[B:158] Dorsal ridge.
1332	0 absent	
1333	1 present	
1334		
1335 1336	159. fossa	[B:159] Shape of median dorsal ridge anterior to endolymphatic a.
1337	0 develope	d as a squared-off ridge or otherwise ungrooved
1338	1 bears a m	nidline groove
1339		
1340	160.	[B:160] Endolymphatic ducts in neurocranium.
1341	0 posteriod	orsally angled tubes

1342	1 tubes oriented vertically through median endolymphatic fossa	
1343		
1344	161.	[B:161] Position of hyomandibula articulation on neurocranium.
1345	0 below or	anterior to orbit, on ventrolateral angle of braincase
1346	1 on otic ca	psule, posterior to orbit
1347		
1348 1349	162. of sk	[B:162] Position of hyomandibula articulation relative to structure eletal labyrinth.
1350	0 anterior o	r lateral to skeletal labyrinth
1351	1 at level of	f posterior semicircular canal
1352		
1353	163.	[B:163] Hyoid arch articulation on braincase.
1354	0 single	
1355	1 double	
1356		
1357	164.	[B:164] Branchial ridges.
1358	0 present	
1359	1 reduced t	o vagal process
1360	2 absent (a	rticulation made with bare cranial wall)
1361		
1362	165.	[B:165] Craniospinal process.
1363	0 absent	
1364	1 present	
1365		
1366	166.	[B:166] Ventral cranial fissure.
1367	0 absent	
1368	1 present	
1369		
1370	167.	[B:167] Basicranial fenestra.
1371	0 absent	
1372	1 present	

1373		
1374	168.	[B:168] Metotic (otic-occipital) fissure.
1375	0 absent	
1376	1 present	
1377		
1378	169.	[B:169] Vestibular fontanelle.
1379	0 absent	
1380	1 present	
1381		
1382	170.	[B:170] Occipital arch wedged in between otic capsules.
1383	0 absent	
1384	1 present	
1385		
1386	171.	[B:171] Spino-occipital nerve foramina.
1387	0 two or mo	pre, aligned horizontally
1388	1 one or two	o, dorsoventrally offset
1389		
1390	172.	[B:172] Ventral notch between parachordals.
1391	0 present o	r entirely unfused
1392	1 absent	
1393		
1394	173.	[B:173] Parachordal shape.
1395	0 forming a	broad, flat surface as wide as the otic capsules
1396	1 mediolate	rally constricted relative to the otic capsules
1397		
1398	174.	[B:174] Stalk-shaped parachordal/occipital region.
1399	0 absent	
1400	1 present	
1401		
1402	175.	[B:175] Paired occipital facets.

1403	0 absent	
1404	1 present	
1405		
1406	176.	[B:176] Size of aperture to notochordal canal.
1407	0 much sma	ller than foramen magnum
1408	1 as large, c	r larger, than foramen magnum
1409		
1410	177.	[B:177] Canal for median dorsal aorta within basicranium.
1411	0 absent	
1412	1 present	
1413		
1414 1415	178. canal	[B:178] Hypotic lamina (and dorsally directed glossopharyngeal).
1416	0 absent	
1417	1 present	
1418		
1419	179.	[B:179] Macromeric dermal shoulder girdle.
1420	0 present	
1421	1 absent	
1422		
1423	180.	[B:180] Dermal shoulder girdle composition.
1424	0 ventral and	d dorsal (scapular) components
1425	1 ventral cor	mponents only
1426		
1427 1428	181. cleith	[B:181] Shape of dorsal blade of dermal shoulder girdle (either rum or anterolateral plate).
1429	0 spatulate	
1430	1 pointed	
1431		
1432 1433	182. trunk.	[B:182] Dermal shoulder girdle forming a complete ring around the

1434	0 present	
1435	1 absent	
1436		
1437 1438	183. armo	[B:183] Pectoral fenestra completely encircled by dermal shoulder ur.
1439	0 present	
1440	1 absent	
1441		
1442	184.	[B:184] Median dorsal plate.
1443	0 absent	
1444	1 present	
1445		
1446	185.	[B:185] Posterior dorsolateral (PDL) plate or equivalent.
1447	0 absent	
1448	1 present	
1449		
1450 1451	186. (i.e., d	[B:186] Pronounced internal median keel on dorsal shoulder girdle crista of median dorsal plate).
1452	0 absent	
1453	1 present	
1454		
1455	187.	[B:187] Crista internalis of dermal shoulder girdle.
1456	0 absent	
1457	1 present	
1458		
1459	188.	[B:188] Scapular infundibulum.
1460	0 absent	
1461	1 present	
1462		
1463	189.	[B:189] Scapular process of shoulder endoskeleton.
1464	0 absent	

1465	1 present	
1466		
1467	190.	[B:190] Ventral margin of separate scapular ossification.
1468	0 horizontal	
1469	1 deeply an	gled
1470		
1471	191.	[B:191] Cross sectional shape of scapular process.
1472	0 flattened o	or strongly ovate
1473	1 subcircula	ır
1474		
1475	192.	[B:192] Flange on trailing edge of scapulocoracoid.
1476	0 absent	
1477	1 present	
1478		
1479	193.	[B:193] Scapular process with posterodorsal angle.
1480	0 absent	
1481	1 present	
1482		
1483	194.	[B:194] Endoskeletal postbranchial lamina on scapular process.
1484	0 present	
1485	1 absent	
1486		
1487	195.	[B:195] Mineralisation of internal surface of scapular blade.
1488	0 mineralise	ed all around
1489	1 unmineral	ised on internal face forming a hemicylindrical cross-section
1490		
1491	196.	[B:196] Coracoid process.
1492	0 absent	
1493	1 present	
1494		

1495	197.	[B:197] Procoracoid mineralisation.
1496	0 absent	
1497	1 present	
1498		
1499	198.	[B:198] Fin base articulation on scapulocoracoid.
1500	0 deeper th	an wide (stenobasal)
1501	1 wider tha	n deep (eurybasal)
1502		
1503	199.	[B:199] Pectoral fin articulation.
1504	0 monobas	al
1505	1 polybasal	
1506		
1507	200.	[B:200] Number of basals in polybasal pectoral fins.
1508	0 three or n	nore
1509	1 two	
1510		
1511	201.	[B:201] Branching radials in paired fins.
1512	0 absent	
1513	1 present	
1514		
1515	202.	[B:202] Number of mesomeres in metapterygial axis.
1516	0 five or fev	ver
1517	1 seven or	more
1518		
1519	203.	[B:203] Biserial pectoral fin endoskeleton.
1520	0 absent	
1521	1 present	
1522		
1523	204.	[B:204] Perforate propterygium.
1524	0 absent	

1525	1 present	
1526		
1527	205.	[B:205] Filamentous extension of pectoral fin from axillary region.
1528	0 absent	
1529	1 present	
1530		
1531	206.	[B:206] Pelvic fins.
1532	0 absent	
1533	1 present	
1534		
1535	207.	[B:207] Pelvic claspers.
1536	0 absent	
1537	1 present	
1538		
1539	208.	[B:208] Dermal pelvic clasper ossifications.
1540	0 absent	
1541	1 present	
1542		
1543	209.	[B:209] Pectoral fins covered in macromeric dermal armour.
1544	0 absent	
1545	1 present	
1546		
1547	210.	[B:210] Pectoral fin base has large, hemispherical dermal
1548	0 absort	ionent.
1549	1 present	
1550	i present	
1221	244	[P:211] Doroal fin oningo
1552	211. O absort	[D.2 1] Dorsai ini spines.
1223		
1554	i present	
1555		

1556	212.	[B:212] Anal fin spine.	
1557	0 absent		
1558	1 present		
1559			
1560	213.	[B:213] Paired fin spines.	
1561	0 absent		
1562	1 present		
1563			
1564	214.	[B:214] Median fin spine insertion.	
1565	0 shallow, r	not greatly deeper than dermal bones/scales	
1566	1 deep		
1567			
1568	215.	[B:215] Prepelvic fin spines (modified).	
1569 1570 1571	We follow Burrow et al. ¹⁶ and Hanke & Wilson ¹⁷ in labelling the fin spine pairs developed between the pectoral and pelvic fin spines as 'prepelvic' instead of the 'intermediate' used by Brazeau et al. ¹¹		
1572	0 absent		
1573	1 present		
1574			
1575	216.	[B:216] Fin spine cross-section.	
1576	0 Round or	horseshoe shaped	
1577	1 Flat-sideo	l, with rectangular profile	
1578			
1579	217.	[B:217] Prepelvic spines when present (modified).	
1580	See comme	ents on character 215.	
1581	0 one pair		
1582	1 multiple p	airs	
1583			
1584	218.	[B:218] Paired prepectoral spines (modified).	
1585	Modified to	enable coding for lateral pairs of prepectoral spines.	
1586	0 absent		

1587	1 present	
1588		
1589	219.	[B:219] Fin spines with ridges.
1590	0 absent	
1591	1 present	
1592		
1593	220.	[B:220] Fin spines with nodes.
1594	0 absent	
1595	1 present	
1596		
1597	221.	[B:221] Fin spines with rows of large retrorse denticles.
1598	0 absent	
1599	1 present	
1600		
1601	222.	[B:222] Expanded spine rib on leading edge of spine.
1602	0 absent	
1603	1 present	
1604		
1605	223.	[B:223] Spine ridges
1606	0 converginę	g at the distal apex of the spine
1607	1 convergino	g on leading edge of spine
1608		
1609	224.	[B:224] Synarcual.
1610	0 absent	
1611	1 present	
1612		
1613	225.	[B:225] Series of thoracic supraneurals.
1614	0 absent	
1615	1 present	
1616		

1617	226.	[B:226] Number of dorsal fins, if present.
1618	0 one	
1619	1 two	
1620		
1621	227.	[B:227] Posterior dorsal fin shape.
1622	0 base appro	eximately as broad as tall, not broader than all of other median fins
1623 1624	1 base much other dorsal	longer than the height of the fin, substantially longer than any of the fins
1625		
1626	228.	[B:228] Basal plate in dorsal fin.
1627	0 absent	
1628	1 present	
1629		
1630 1631	229. plate.	[B:229] Branching radial structure articulating with dorsal fin basal
1632	0 absent	
1633	1 present	
1634		
1635	230.	[B:230] Anal fin.
1636	0 absent	
1637	1 present	
1638		
1639	231.	[B:231] Basal plate in anal fin.
1640	0 absent	
1641	1 present	
1642		
1643	232.	[B:232] Caudal radials.
1644	0 extend bey	rond level of body wall and deep into hypochordal lobe
1645	1 radials rest	ricted to axial lobe
1646		
1647	233.	[B:233] Supraneurals in axial lobe of caudal fin.

1648	0 absent	
1649	1 present	
1650		
1651	234.	[B:234] Epichordal lepidotrichia in caudal fin.
1652	0 absent	
1653	1 present	
1654		
1655	235.	[B:235] Enamel and pore canals.
1656	0 enamel at	osent from inner surface of pores
1657	1 enamel lir	es portions of pore canal
1658		
1659 1660	236. marg	[B:236] Canal-bearing bone of skull roof extends far past posterior in of parietals.
1661	0 no	
1662	1 yes	
1663		
1664	237.	[B:237] Pineal eminence (in taxa lacking pineal foramen).
1665	0 absent	
1666	1 present	
1667		
1668	238.	[B:238] Position of anterior pitline.
1669	0 on postpa	rietal
1670	1 on parieta	l
1671		
1672 1673	239. bone	[B:239] Opening in dermal skull roof for spiracular bounded by s carrying otic canal.
1674	0 absent	
1675	1 present	
1676		
1677	240.	[B:240] Median skull roof bone between postparietals.
1678	0 absent	

1679	1 present	
1680		
1681	241.	[B:241] Westoll lines.
1682	0 absent	
1683	1 present	
1684		
1685	242.	[B:242] Preoperculosubmandibular.
1686	0 absent	
1687	1 present	
1688		
1689	243.	[B:243] Hyomandibula.
1690	0 imperforat	e
1691	1 perforate	
1692		
1693	244.	[B:244] Urohyal shape.
1694	0 absent	
1695	1 vertical pla	ite
1696		
1697	245.	[B:246] Length of dentary.
1698	0 constitutes	a majority of jaw length
1699	1 half the ler	ngth of jaw or less
1700		
1701	246.	[B:247] Labial pit.
1702	0 absent	
1703	1 present	
1704		
1705	247.	[B:248] Prearticular symphysis.
1706	0 absent	
1707	1 present	
1708		
1707 1708	1 present	

1709	248.	[B:249] Mandibular sensory canal.
1710	0 extends th	nrough infradentaries
1711	1 extends th	nrough infradentaries and dentary
1712		
1713 1714	249. bone	[B:250] Extensive flange composed of prearticular and Meckelian that extends beyond ventral edge of outer dermal series.
1715	0 absent	
1716	1 present	
1717		
1718	250.	[B:251] Posterior coronoid.
1719	0 similar to	anterior coronoids
1720	1 forms exp	anded coronoid process
1721		
1722	251.	[B:252] Retroarticular process.
1723	0 absent	
1724	1 present	
1725		
1726	252.	[B:253] Inturned medial process of premaxilla.
1727	0 absent	
1728	1 present	
1729		
1730 1731	253. and s	[B:254] Anteriorly directed adductor fossae between neurocranium skull roof.
1732	0 absent	
1733	1 present	
1734		
1735	254.	[B:255] Vomerine fangs.
1736	0 absent	
1737	1 present	
1738		
1739	255.	[B:256] Number of dermopalatines.

1740	0 multiple	
1741	1 one	
1742		
1743	256.	[B:257] Entopterygoids.
1744	0 separated	i i i i i i i i i i i i i i i i i i i
1745	1 contact al	long midline
1746		
1747	257.	[B:258] Rostral tubuli.
1748	0 absent	
1749	1 present	
1750		
1751	258.	[B:259] Position of anterior nostril.
1752	0 facial	
1753	1 at oral ma	argin
1754		
1755	259.	[B:260] Posterior nostril.
1756	0 facial	
1757	1 at margin	of oral cavity
1758	2 palatal	
1759		
1760 1761	260. each	[B:261] Three large pores (in addition to nostrils) associated with side of ethmoid.
1762	0 absent	
1763	1 present	
1764		
1765 1766	261. ethm	[B:262] Ventral face of nasal capsule in taxa with mineralized oid.
1767	0 complete	
1768	1 fenestra v	ventrolateralis
1769	2 entire floo	or unmineralized
1770		

1771	262.	[B:263] Size of profundus canal in postnasal wall.
1772	0 small	
1773	1 large	
1774		
1775	263.	[B:264] Paired pineal and parapineal tracts.
1776	0 absent	
1777	1 present	
1778		
1779	264.	[B:265] Posterior of parasphenoid.
1780	0 restricted	to ethmosphenoid region
1781	1 extends to	o otic region
1782		
1783	265.	[B:266] Endoskeletal spiracular canal.
1784	0 open	
1785	1 spiracular	bar
1786	2 complete	enclosure in canal
1787		
1788	266.	[B:267] Barbed lepidotrichial segments.
1789	0 absent	
1790	1 present	
1791		
1792 1793	267. hyom	[B:268] Relative position of jugular groove/canal and and and and and bular articulation.
1794	0 hyomandi	bula dorsal
1795	1 hmd strad	dles
1796	2 hmd ventr	al
1797		
1798	268.	[B:269] Optic lobes.
1799	0 narrower f	han cerebellum
1800	1 same widt	h or wider than cerebellum
1801		

1802	269.	[B:270] Hypophyseal chamber.
1803	0 projects p	osteroventrally
1804	1 projects v	entrally or anteroventrally
1805		
1806 1807	270. canal	[B:271] Crus commune of anterior and posterior semicircular s.
1808	0 dorsal to b	praincase endocavity roof
1809	1 ventral to	braincase endocavity roof
1810		
1811	271.	[B:272] Horizontal semicircular canal.
1812	0 obliquely o	oriented
1813	1 horizontal	ly oriented
1814		
1815	272.	[B:273] Supraotic cavity.
1816	0 absent	
1817	1 present	
1818		
1819	273.	[B:274] Pelvic girdle with substantial dermal component.
1820	0 present	
1821	1 absent	
1822		
1823	274.	[B:275] Pelvic fin spines.
1824	0 absent	
1825	1 present	
1826		
1827	275.	[B:276] Pelvic fin.
1828	0 monobasa	al
1829	1 polybasal	
1830		
1831	276.	[B:277] Postparietals/centrals.
1832	0 absent	

1833	1 present		
1834			
1835	277.	[B:278] Condition of postparietals/centrals.	
1836	0 do not me	eet in midline	
1837	1 meet in midline		
1838	2 single mic	lline bone	
1839			
1840	278.	[B:279] Parietals.	
1841	0 absent		
1842	1 present		
1843			
1844	279.	[B:280] Condition of parietals.	
1845	0 do not me	eet in midline	
1846	1 meet in m	idline	
1847			
1848 1849	280. nostr	[B:281] Endoskeletal lamina (postnasal wall) separating posterior il and orbit.	
1850	0 absent		
1851	1 present		
1852			
1853	281.	[B:282] Pituitary vein canal.	
1854	0 discontinu	ious, enters the cranial cavity	
1855	1 discontinu	ious, enters hypophysial recess	
1856	2 continuou	s transverse vein	
1857			
1858	282.	[B:283] Sutures between dermal bones.	
1859	0 absent		
1860	1 present		
1861			
1862	283.	[B:284] Interolateral/clavicular margin.	
1863	0 angled an	terolaterally	

1864	1 mediolater	ally straight
1865		
1866 1867	284. odont	Scale odontodes added in a linear sequence within rows (linear ocomplexes) (new character).
1868	0 absent	
1869	1 present	
1870		
1871 1872	285. chara	Number of linear odontocomplexes in scale crowns (new cter).
1873	0 one	
1874	1 more than	one
1875		
1876 1877	286. assoc	[D:262] Anteriormost prepelvic fin spine (admedian fin spine) ated with the shoulder girdle (modified).
1878 1879 1880 1881 1882	We agree wi medially of the climatiids ^{18,1} Here these a other studies	th Dearden et al. ¹³ in recognising the shoulder girdle spines positioned he pectoral fin spines in a number of stem chondrichthyans (e.g. ⁹ , diplacanthids ²⁰ and gyracanthids ²¹) as the first prepelvic fin spine pair. are labelled admedian after Burrow et al. ^{18,20,22} , who contrary to this and s ^{13,23} consider them separate from the prepelvic series.
1883	0 absent	
1884	1 present	
1885		
1886	287.	[K:482] Median ventral prepectoral spine.
1887	0 absent	
1888	1 present	
1889		
1890	288.	[K:447] Median ventral trunk plates.
1891	0 absent	
1891 1892	0 absent 1 present	
1891 1892 1893	0 absent 1 present	
1891 1892 1893 1894 1895	0 absent 1 present 289. (modi	[B:17] Odontode resorption in the extra-oral dermal skeleton fied).

- 1897 0 absent
- 1898 1 present
- 1899

1900 **290.** Pinnal plates of the dermal shoulder girdle (new character).

Pinnal plate development differs from that of the dermal plates of jawed stem
gnathostomes ('placoderms')^{24,25} and osteichthyans²⁶⁻²⁸. The latter form as a single
unit through areal growth unlike the pinnals where independent dermal scales are
integrated into discrete elements fused together by a basal plate^{18,20,29} (see also this
study). On this basis, and in accordance with previous research³⁰, we code for
presence/absence of pinnal plates independently of the pectoral ventral plate pairs of
'placoderms' (ventrolateral plates) and osteichthyans (clavicles).

- 1908 0 absent
- 1909 1 present
- 1910

1911291. Dermal bone structure of scales and non-dental plates (new1912character). The stem chondrichthyans *Climatius*, *Diplacanthus* and1913*Ptomacanthus* possess extra-oral dermal elements (e.g. shoulder girdle plates,1914branchiostegal plates and scales) with two distinct bone architectures1915(compact and spongy)^{13,18,20}. They are therefore considered polymorphic for1916this character instead of possessing state (0) where the bone tissue has a1917tiered structure produced by the formation of compact and vascular bone.

- 1918 0 compact and vascular
- 1919 1 compact
- 1920 2 vascular
- 1921

1922 **292**. Size of primary teeth within whorls (new character)

- 1923 0 remains consistent
- 1924 1 increases gradually
- 1925
- 1926293. Accessory teeth in tooth whorls (new character). These have been1927documented^{31,32} to develop laterally and/or labially of the primary whorl teeth1928and are distinguished from the latter by their noticeably smaller size and1929different organisation/patterning.
- 1930 0 absent
- 1931 1 present
1932

- 1933294.Mandibular teeth (new character)
- 1934 0 monocuspid
- 1935 1 multicuspid

1936

1937

- **Table S1.** Tip (taxon) ages of the 50 percent majority-rule tree shown in Extended
- 1939 Data Fig. 4.

Taxon	Age in million years	Reference
Acanthodes	298	King et al. ¹⁵
Achoania	412	King et al. ¹⁵
Akmonistion	327	King et al. ¹⁵
Austroptyctodus	327	King et al. ¹⁵
Bothriolepis	383	King et al. ¹⁵
Brachyacanthus	415	King et al. ¹⁵
Brindabellaspis	401	King et al. ¹⁵
Brochoadmones	415	King et al. ¹⁵
Buchanosteus	408	King et al. ¹⁵
Campbellodus	383	King et al. ¹⁵
Cassidiceps	415	King et al. ¹⁵
Cheiracanthus	388	King et al. ¹⁵
Cheirolepis	388	King et al. ¹⁵
Chondrenchelys	338	King et al. ¹⁵
Cladodoides	375	King et al. ¹⁵
Cladoselache	360	King et al. ¹⁵
Climatius	415	King et al. ¹⁵
Cobelodus	325	King et al. ¹⁵
Coccosteus	388	King et al. ¹⁵
Compagopiscis	383	King et al. ¹⁵

Cowralepis	383	King et al. ¹⁵
Culmacanthus	385	King et al. ¹⁵
Debeerius	320	King et al. ¹⁵
Diabolepis	412	King et al. ¹⁵
Dialipina	401	King et al. ¹⁵
Dicksonosteus	411	King et al. ¹⁵
Diplacanthus	388	King et al. ¹⁵
Diplocercides	383	Long and Trinajstic ³³
Dipterus	388	Gross ³⁴
Doliodus	395	King et al. ¹⁵
Entelognathus	424	King et al. ¹⁵
Eurycaraspis	385	King et al. ¹⁵
Eusthenopteron	380	King et al. ¹⁵
Euthacanthus	415	King et al. ¹⁵
Galeaspida	436	King et al. ¹⁵
Gavinia	385	Long ³⁵
Gemuendina	408	King et al. ¹⁵
Gladbachus	388	Coates et al. ¹²
Gladiobranchus	415	King et al. ¹⁵
Glyptolepis	388	King et al. ¹⁵
Gogonasus	383	King et al. ¹⁵
Guiyu	424	King et al. ¹⁵
Gyracanthides	388	Warren et al. ²¹
Halimacanthodes	383	Burrow et al. ³⁶
Hamiltonichthys	302	King et al. ¹⁵
Helodus	311	King et al. ¹⁵
Homalacanthus	380	King et al. ¹⁵
Howqualepis	385	King et al. ¹⁵
Incisoscutum	383	King et al. ¹⁵
Iniopera	307	Pradel et al. ³⁷
Ischnacanthus	415	King et al. ¹⁵

Jagorina	375	King et al. ¹⁵
Janusiscus	415	King et al. ¹⁵
Kansasiella	303	King et al. ¹⁵
Kathemacanthus	415	King et al. ¹⁵
Kentuckia	347	King et al. ¹⁵
Kosoraspis	419	Vaškaninová et al.38
Kujdanowiaspis	411	King et al. ¹⁵
Latviacanthus	404	King et al. ¹⁵
Lawrenciella	303	Poplin 1984
'Ligulalepis'	401	King et al. ¹⁵
Lophosteus	423	Schultze and Märss ³⁹
Lunaspis	408	King et al. ¹⁵
Lupopsyrus	415	King et al. ¹⁵
Macropetalichthys	390	King et al. ¹⁵
Meemannia	412	King et al. ¹⁵
Mesacanthus	415	King et al. ¹⁵
Miguashaia	380	King et al. ¹⁵
Mimipiscis	383	King et al. ¹⁵
Minjinia	411	Brazeau et al. ¹¹
Moythomasia	383	King et al. ¹⁵
Nerepisacanthus	423	King et al. ¹⁵
Obtusacanthus	415	King et al. ¹⁵
Onychodus	383	King et al. ¹⁵
Onychoselache	336	King et al. ¹⁵
Orthacanthus	290	King et al. ¹⁵
Osteostraci	427	King et al. ¹⁵
Parayunnanolepis	412	King et al. ¹⁵
Parexus	415	King et al. ¹⁵
Poracanthodes	417	King et al. ¹⁵
Porolepis	411	King et al. ¹⁵
Powichthys	411	King et al. ¹⁵

Promesacanthus	415	King et al. ¹⁵
Psarolepis	416	King et al. ¹⁵
Pterichthyodes	389	King et al. ¹⁵
Ptomacanthus	415	King et al. ¹⁵
Ptyctolepis	411	Lu et al. ⁴⁰
Pucapampella	388	King et al. ¹⁵
Qianodus	439	this study
Qilinyu	424	Zhu et al. ⁴¹
Qingmenodus	411	Lu and Zhu ⁴²
Ramirosuarezia	392	King et al. ¹⁵
Raynerius	373	Giles et al. ¹⁴
Rhadinacanthus	388	Burrow et al. ²⁰
Rhamphodopsis	388	King et al. ¹⁵
Romundina	415	King et al. ¹⁵
Sparalepis	424	King et al. ¹⁵
Styloichthys	412	King et al. ¹⁵
Tamiobatis	360	King et al. ¹⁵
Tetanopsyrus	415	King et al. ¹⁵
Tristychius	336	King et al. ¹⁵
Uranolophus	411	Denison ⁴³
Vernicomacanthus	415	King et al. ¹⁵
Youngolepis	412	King et al. ¹⁵
Yunnanolepis	415	King et al. ¹⁵

1940

1941

1942 **Descriptions of Supplementary Files**

- 1943 Supplementary Data 1. Volume rendering of the holotype of *Qianodus* (IVPP
- 1944 V26641) based on synchrotron X-ray tomography data acquired at the Taiwan Light
- 1945 Source (TLS), National Synchrotron Radiation Research Center (NSRRC), Taiwan.

Colour coded features: orange, progenitor tooth row; green, trailing tooth row; yellow,
accessory teeth; grey, whorl base. Rendering generated in Mimics 19.0.

1948

Supplementary Data 2. Volume rendering of synchrotron X-ray tomography data
depicting radiotransparent spaces inside the holotype of *Qianodus* (IVPP V26641)
analysed at the Taiwan Light Source (TLS), National Synchrotron Radiation
Research Center (NSRRC), Taiwan. Colour coded features: purple, primary teeth;
yellow, accessory teeth; white, spongiose tissue of the whorl base; pink, compact
tissue of the whorl base. Rendering generated in Mimics 19.0.

1955

Supplementary Data 3. Tomographic slices from an X-ray tomography analysis of
the *Qianodus* holotype specimen (IVPP V26641) at the Taiwan Light Source (TLS),
National Synchrotron Radiation Research Center (NSRRC), Taiwan. Dataset
generated in Mimics 19.0.

1960

Supplementary Data 4. Tomographic slices from an X-ray tomography analysis of a *Qianodus* specimen IVPP V26645 at the Taiwan Light Source (TLS), National
Synchrotron Radiation Research Center (NSRRC), Taiwan. Dataset generated in
Mimics 19.0.

1965

Supplementary Data 5. Tomographic slices from an X-ray tomography analysis of a
 Qianodus specimen IVPP V26647 at the Institute of Vertebrate Paleontology and

76

Paleoanthropology, Chinese Academy of Sciences. Dataset generated in Mimics1969 19.0.

1971	Supplementary Data 6. Parsimony analysis files. Character-taxon matrix in TNT
1972	(.tnt) and nexus (.nex) file formats. Most parsimonious trees (.tre) produced by the
1973	parsimony analysis. 50 percent majority-rule consensus tree (.tre) and strict
1974	consensus tree (.tre) for the set of most parsimonious trees. Likelihood and
1975	parsimony reconstructions of character states at internal nodes of the 50 percent
1976	majority-rule consensus tree in excel (.xlxs) file format. TNT log file in rich text format
1977	(.rtf) of the parsimony analysis and the bootstrap resampling analysis. R script in rich
1978	text format (.rtf) used in the calculation of the time-scaled 50 percent majority-rule
1979	consensus tree.
1980	

References

1982 1983 1984	1	Aldridge, R. J. & Wang, CY. in <i>Telychian Rocks of the British Isles and China (Silurian, Llandovery Series: An Experiment to Test Precision in Stratigraphy National Museum of Wales Geological Series</i> (eds Holland, C. H. & Bassett, M. G.) 83-94 (Nat. Mus. Wales, 2002).
1985	2	Rong, J., Johnson, M. E. & Yang, X. Early Silurian (Llandovery) sealevel changes in the Upper
1986		Yangtze region of central and southwestern China. Acta Palaeont. Sin. 23, 672-694 (1984).
1987 1988	3	Rong, J., Wang, Y. & Zhang, X. Tracking shallow marine red beds through geological time as
1989		<i>China Earth Sci.</i> 55 , 699-713 (2012).
1990	4	Mu, E. Correlation of the Silurian Rocks of China. Vol. 202 (Geological Society of America,
1991		1986).
1992	5	Wang, CY. Restudy on the ages of Silurian red beds in South China. J. Stratigr. 35, 440-447
1993		(2011).
1994	6	Wang, CY. Silurian Conodonts in China (Univ. Sci. Tech. China Press, 2013).
1995	7	Wang, CY., Chen, L., Wang, Y. & Tang, P. Affirmation of Pterospathodus eopennatus Zone
1996		(Conodonta) and the age of the Silurian Shamao Formation in Zigui, Hubei as well as the
1997		correlation of the related strata. Acta Palaeont. Sin. 49, 10-28 (2010).
1998	8	Rong, J. et al. Silurian integrative stratigraphy and timescale of China. Sci. China Earth Sci. 62,
1999		89-111 (2018).
2000	9	Tang, P., Xu, H. & Wang, Y. Chitinozoan-based age of the Wengxiang Group in Kaili,
2001		southeastern Guizhou, Southwest China. J. Earth Sci. 21, 52 (2010).

- 200210Zhou, X., Zhai, Z.-Q. & Xian, S. On the Silurian conodont biostratigraphy, new genera and2003species in Guizhou Province. *Oil Gas Geol.* 2, 123-140 (1981).
- 200411Brazeau, M. *et al.* Endochondral bone in an Early Devonian 'placoderm' from Mongolia. *Nat.*2005*Ecol. Evol.* **4**, 1477–1484 (2020).
- 200612Coates, M. I. *et al.* An early chondrichthyan and the evolutionary assembly of a shark body2007plan. *Proc. R. Soc B* 285, 20172418 (2018).
- 200813Dearden, R. P., Stockey, C. & Brazeau, M. D. The pharynx of the stem-chondrichthyan2009Ptomacanthus and the early evolution of the gnathostome gill skeleton. Nat. Commun. 10,20102050 (2019).
- 201114Giles, S., Friedman, M. & Brazeau, M. D. Osteichthyan-like cranial conditions in an Early2012Devonian stem gnathostome. Nature **520**, 82-85 (2015).
- King, B., Qiao, T., Lee, M. S., Zhu, M. & Long, J. A. Bayesian morphological clock methods
 resurrect placoderm monophyly and reveal rapid early evolution in jawed vertebrates. *Syst. Biol.* 66, 499-516 (2017).
- 201616Brazeau, M. D. & Friedman, M. The origin and early phylogenetic history of jawed2017vertebrates. Nature 520, 490-497 (2015).
- 201817Hanke, G. & Wilson, M. in Morphology, Phylogeny and Paleobiogeography of Fossil Fishes.2019(eds Elliott, D. K., Maisey, J. G., Yu, X. & Miao, D.) 159-182 (Verlag Dr. Friedrich Pfeil, 2010).
- 202018Burrow, C. J., Davidson, R. G., Den Blaauwen, J. L. & Newman, M. J. Revision of Climatius2021reticulatus Agassiz, 1844 (Acanthodii, Climatiidae), from the Lower Devonian of Scotland,2022based on new histological and morphological data. J. Vertebr. Paleont. 35, e913421 (2015).
- Miles, R. S. Articulated acanthodian fishes from the Old Red Sandstone of England: with a
 review of the structure and evolution of the acanthodian shoulder-girdle. *Bull. Br. Mu. Nat. Hist. Geol.* 24, 111-213 (1973).
- 2026 20 Burrow, C., den Blaauwen, J., Newman, M. & Davidson, R. The diplacanthid fishes
 2027 (Acanthodii, Diplacanthiformes, Diplacanthidae) from the Middle Devonian of Scotland.
 2028 Palaeontol. Electron. 19, 1-83 (2016).
- 2029 21 Warren, A., Currie, B. P., Burrow, C. & Turner, S. A redescription and reinterpretation of
 2030 *Gyracanthides murrayi* Woodward 1906 (Acanthodii, Gyracanthidae) from the Lower
 2031 Carboniferous of the Mansfield Basin, Victoria, Australia. *J. Vertebr. Paleontol.* 20, 225-242
 2032 (2000).
- 203322Burrow, C. J., Newman, M. J., Davidson, R. G. & den Blaauwen, J. L. Redescription of *Parexus*2034recurvus, an Early Devonian acanthodian from the Midland Valley of Scotland. Alcheringa **37**,2035392-414 (2013).
- 203623Gagnier, P.-Y. & Wilson, M. V. Early Devonian acanthodians from northern Canada.2037Palaeontology **39**, 241-258 (1996).
- 203824Dupret, V., Sanchez, S., Goujet, D., Tafforeau, P. & Ahlberg, P. E. Bone vascularization and2039growth in placoderms (Vertebrata): The example of the premedian plate of *Romundina*2040stellina Ørvig, 1975. C. R. Palevol **9**, 369-375 (2010).
- 204125Giles, S., Rücklin, M. & Donoghue, P. C. Histology of "placoderm" dermal skeletons:2042Implications for the nature of the ancestral gnathostome. J. Morph. 274, 627-644 (2013).
- 26 Andrews, M., Long, J., Ahlberg, P., Barwick, R. & Campbell, K. The structure of the
 2044 sarcopterygian *Onychodus jandemarrai* n. sp. from Gogo, Western Australia: with a
 2045 functional interpretation of the skeleton. *Earth. Env. Sci. Trans. R. Soc. Edinb.* 96, 197-307
 2046 (2005).
- 2047 27 Mondéjar-Fernández, J., Friedman, M. & Giles, S. Redescription of the cranial skeleton of the
 2048 Early Devonian (Emsian) sarcopterygian *Durialepis edentatus* Otto (Dipnomorpha,
 2049 Porolepiformes). *Pap. Palaeontol.* (2020).
- 205028Zhu, M., Yu, X., Wang, W., Zhao, W. & Jia, L. A primitive fish provides key characters bearing2051on deep osteichthyan phylogeny. Nature 441, 77-80 (2006).

- 2052 29 Long, J. A new diplacanthoid acanthodian from the Late Devonian of Victoria. *Mem. Assoc.*2053 *Australas. Palaeontol.* 1, 51-65 (1983).
- 2054 30 Moy-Thomas, J. & Miles, R. S. Palaeozoic Fishes (Chapman and Hall, 1971).
- 205531Ahlberg, P. E. A new holoptychiid porolepiform fish from the Upper Frasnian of Elgin,2056Scotland. Palaeontology **35**, 813-828 (1992).
- 205732Qu, Q., Sanchez, S., Blom, H., Tafforeau, P. & Ahlberg, P. E. Scales and tooth whorls of2058ancient fishes challenge distinction between external and oral 'teeth'. *PLoS One* **8**, e718902059(2013).
- 2060 33 Long, J. A. & Trinajstic, K. The Late Devonian Gogo Formation lägerstatte of Western
 2061 Australia: exceptional early vertebrate preservation and diversity. *Annu. Rev. Earth Planet*2062 *Sci.* 38, 255-279 (2010).
- 206334Gross, W. Über die Randzähne des Mundes, die Ethmoidalregion des Schädels und die2064Unterkiefersymphyse von Dipterus oervigi n. sp. Paläontol. Z. **38**, 7-25 (1964).
- 206535Long, J. A. A new genus of fossil coelacanth (Osteichthyes: Coelacanthiformes) from the2066Middle Devonian of southeastern Australia. *Rec. West. Aust. Mus., Suppl* **57**, 37-53 (1999).
- 206736Burrow, C. J., Trinajstic, K. & Long, J. First acanthodian from the Upper Devonian (Frasnian)2068Gogo Formation, Western Australia. *Hist. Biol.* 24, 349-357 (2012).
- 2069 37 Pradel, A., Tafforeau, P. & Janvier, P. Study of the pectoral girdle and fins of the Late
 2070 Carboniferous sibyrhynchid iniopterygians (Vertebrata, Chondrichthyes, Iniopterygia) from
 2071 Kansas and Oklahoma (USA) by means of microtomography, with comments on
 2072 iniopterygian relationships. *C. R. Palevol* 9, 377-387 (2010).
- 207338Vaškaninová, V. *et al.* Marginal dentition and multiple dermal jawbones as the ancestral2074condition of jawed vertebrates. *Science* **369**, 211-216 (2020).
- 2075 39 Schultze, H.-P. & Märss, T. Revisiting *Lophosteus*, a primitive osteichthyan. *Acta Univ. Latv.*2076 679, 57-78 (2004).
- 207740Lu, J., Giles, S., Friedman, M. & Zhu, M. A new stem sarcopterygian illuminates patterns of2078character evolution in early bony fishes. Nat. Commun. 8, 1932 (2017).
- 207941Zhu, M. *et al.* A Silurian maxillate placoderm illuminates jaw evolution. *Science* **354**, 334-3362080(2016).
- 208142Lu, J. & Zhu, M. An onychodont fish (Osteichthyes, Sarcopterygii) from the Early Devonian of2082China, and the evolution of the Onychodontiformes. *Proc. Roy. Soc. B: Biol. Sci.* 277, 293-2992083(2010).
- 208443Denison, R. H. Early Devonian lungfishes from Wyoming, Utah, and Idaho. Field Mus. Nat.2085Hist. Bull. 17, 353–413 (1968).