# UNIVERSITY<sup>OF</sup> BIRMINGHAM University of Birmingham Research at Birmingham

# An early chondrichthyan and the evolutionary assembly of a shark body plan

Coates, Michael; Finarelli, John; Sansom, Ivan; Andreev, Plamen; Criswell, Katherine; Tietjen, Kristen; Rivers, Mark; La Riviere, Patrick

DOI: 10.1098/rspb.2017.2418

License: Other (please specify with Rights Statement)

Document Version Peer reviewed version

### Citation for published version (Harvard):

Coates, M, Finarelli, J, Sansom, I, Andreev, P, Criswell, K, Tietjen, K, Rivers, M & La Riviere, P 2018, 'An early chondrichthyan and the evolutionary assembly of a shark body plan', *Royal Society of London. Proceedings B. Biological Sciences*, vol. 285, no. 1870, 20172418. https://doi.org/10.1098/rspb.2017.2418

Link to publication on Research at Birmingham portal

#### Publisher Rights Statement:

Final Version of Record published as: Coates, Michael I., et al. "An early chondrichthyan and the evolutionary assembly of a shark body plan." Proc. R. Soc. B. Vol. 285. No. 1870 - https://doi.org/10.1098/rspb.2017.2418

### **General rights**

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?) •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

#### Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.



# An early chondrichthyan and the evolutionary assembly of a shark body plan

Journal:	Proceedings B
Manuscript ID	RSPB-2017-2418.R1
Article Type:	Research
Date Submitted by the Author:	n/a
Complete List of Authors:	Coates, Michael; University of Chicago, Department of Organismal Biology and Anatomy Finarelli, John; University College Dublin, School of Biology and Environmental Science Sansom, Ivan; University of Birmingham, School of Geography, Earth and Environmental Sciences Andreev, Plamen; University of Birmingham, School of Geography, Earth and Environmental Sciences Criswell, Katharine; University of Cambridge, Department of Zoology Tietjen, Kristen; University of Chicago, Department of Organismal Biology and Anatomy Rivers, Mark; University of Chicago, Center for Advanced Radiation Sources La Riviere, Patrick; University of Chicago, Department of Radiology
Subject:	Palaeontology < BIOLOGY, Taxonomy and Systematics < BIOLOGY, Evolution < BIOLOGY
Keywords:	Chondrichthyes, gill skeleton, scales, Middle Devonian, gnathostome phylogeny, computed tomography
Proceedings B category:	Palaeobiology

SCHOLARONE<sup>™</sup> Manuscripts

1	An early chondrichthyan and the evolutionary assembly of a shark
2	body plan
3	
4	
5	Michael I. Coates <sup>1</sup> *, John A. Finarelli <sup>2</sup> , Ivan J. Sansom <sup>3</sup> , Plamen S. Andreev <sup>3</sup> , Katharine E. Criswell <sup>1,6</sup> ,
6	Kristen Tietjen <sup>1</sup> , Mark L. Rivers <sup>4</sup> , Patrick J. La Riviere <sup>5</sup>
7	
8	<sup>1</sup> Department of Organismal Biology and Anatomy, University of Chicago, Chicago, IL 60637-1508,
9	USA.
10	<sup>2</sup> UCD School of Biology and Environmental Science, UCD Science Education and Research Centre
11	(West), UCD Earth Institute, University College Dublin, Belfield, Dublin 4, Ireland.
12	<sup>3</sup> School of Geography, Earth and Environmental Sciences, University of Birmingham, Birmingham, B15
13	2TT, UK.
14	<sup>4</sup> Center for Advanced Radiation Sources, University of Chicago, Chicago, IL 60637-1508, USA.
15	<sup>5</sup> Department of Radiology, University of Chicago, Chicago, IL 60637-1508, USA.
16	<sup>6</sup> Department of Zoology, Cambridge University, XYZ 123, UK.
17	
18	Corresponding author: E-mail: mcoates@uchicago.edu
19	

### 21 Abstract

22 Although relationships among the major groups of living gnathostomes are well established, the 23 relatedness of early jawed vertebrates to modern clades is intensely debated. Here, we provide a new description of *Gladbachus*, a Middle Devonian (Givetian ~385-million-year-old) stem chondrichthyan 24 25 from Germany, and one of the very few early chondrichthyans in which substantial portions of the 26 endoskeleton are preserved. Tomographic and histological techniques reveal new details of the gill 27 skeleton, hyoid arch and jaws, neurocranium, cartilage, scales and teeth. Despite many features 28 resembling placoderm or osteichthyan conditions, phylogenetic analysis confirms *Gladbachus* as a stem 29 chondrichthyan and corroborates hypotheses that all acanthodians are stem chondrichthyans. The 30 unfamiliar character combination displayed by *Gladbachus*, alongside conditions observed in 31 acanthodians, implies that pre-Devonian stem-chondrichthyans are severely under-sampled and strongly 32 supports indications from isolated scales that the gnathostome crown group originated at the latest by the 33 early Silurian (~440 mya). Moreover, phylogenetic results highlight the likely convergent evolution of 34 conventional chondrichthyan conditions among earliest members of this primary gnathostome division, 35 while skeletal morphology points towards the likely suspension feeding habits of *Gladbachus*, suggesting 36 a functional origin of the gill slit condition characteristic of the vast majority of living and fossil 37 chondrichthyans. 38 39 40 Keywords:

41 Chondrichthyes, gnathostomes, gill skeleton, scales, Middle Devonian, jaws

42

## 44 **1. Introduction**

45 The early evolution of the Chondrichthyes (cartilaginous fishes) has long been obscured by an 46 impoverished fossil record [1-3]. This has only recently been improved through discoveries of partly 47 articulated bodies [4-7] and braincases from the Lower and Middle Devonian [8-10], coupled with 48 computed tomography (CT scanning) [11-16]. The Lower and Middle Devonian record of 49 chondrichthyans remains sparse, but the influx of high quality data from slightly younger material, 50 especially from the Upper Devonian and Carboniferous [14-17], combined with insights from earlier 51 studies [18-21] has transformed our understanding of the early evolution of the crown group (Holocephali 52 and Elasmobranchii). In a complementary manner, the origin of total group Chondrichthyes has been 53 amended by serial large-scale analyses of early gnathostome phylogeny [22-27], which consistently 54 recover acanthodians [28] as stem-chondrichthyans. However, because the acanthodian and 55 chondrichthyan taxa included in these data matrices are largely unaltered from Brazeau [29] and Davis et 56 al. [30], these results are not truly independent. 57 Here, we present a CT analysis and re-description of Gladbachus adentatus [31] from the Middle 58 Devonian (Givetian ~385-million-year-old) of Germany, one of the earliest chondrichthyans known from 59 articulated remains. Unlike *Doliodus* [1,4-7], the earliest and most completely described 'unambiguous' 60 stem chondrichthyan [32], *Gladbachus* has never been included among acanthodians, but, like *Doliodus*, 61 recent analyses and discussion [16,32] suggest that it might illuminate conditions bridging the 62 acanthodian-chondrichthyan transition. Thus, a primary aim of the present work is to test the assumed 63 chondrichthyan affinity of *Gladbachus* in light of the current acanthodians-as-stem-chondrichthyans 64 hypothesis. Here, we have constructed a new early gnathostome data base, with an augmented 65 chondrichthyan component with taxa and characters from analyses by Pradel et al. [14], Coates et al. [16] 66 and Coates & Tietjen [17]. This analysis presents the most detailed context, thus far, to reconstruct the 67 evolutionary assembly of the chondrichthyan morphotype: a body-plan that has persisted, more-or-less 68 conservatively, for at least 370 million years.

69

70	
71	2. Materials and methods
72	(a) Specimens
73	The subject of this study, <i>Gladbachus adentatus</i> [31] is known from a single, dorsoventrally compressed
74	individual, UMZC (University Museum of Zoology, Cambridge, UK) 2000.32 [31,33], collected from the
75	Lower Plattenkalk, Upper Givetian, Upper Middle Devonian, of Unterthal, Bergisch Gladbach
76	(Germany). The specimen (electronic supplementary material, figure S1) consists of three pieces
77	embedded within a rectangular slab of resin, with only the dorsal surface visible for direct inspection.
78	
79	(b) Computed tomography, anatomical reconstruction and histological thin sections.
80	Whole specimen scans: large scale scans were completed by the High Resolution X-ray Computed
81	Tomography facility at the University of Texas at Austin (UTCT, www.digimorph.org).
82	Scales, denticles, teeth, and calcified cartilage were examined using synchrotron $\mu CT$ : all data
83	were collected at beamline 13-BM-D at the Advanced Photon Source at Argonne National Laboratory.
84	Image reconstruction used GSECARS tomography processing software
85	(http://cars9.uchicago.edu/software/idl/tomography.html), which dark-current corrects and white-field
86	normalizes acquired data prior to performing gridding-based image reconstruction. Further details are
87	provided in the electronic supplementary material.
88	Histological thin sections: doubly polished thin sections were studied using a Zeiss Axioskop Pol
89	microscope equipped with Nomarski DIC optics.
90	Anatomical reconstruction: Mimics v. 17 (biomedical.materialise.com/mimics; Materialise, Leuven,
91	Belgium) was used for the three-dimensional modeling, including segmentation, three-dimensional object
92	rendering, STL polygon creation and kinematics. 3D Studio Max (Autodesk.com/products/3ds-max;
93	Autodesk, San Rafael, USA) was used for further editing of the STLs (color, texture, lighting), kinematics,
94	and mirroring for the final restoration.
95	

96	(c) Phylogenetic and phenetic analysis
97	The phylogenetic data matrix is developed from sources including iterations of the early gnathostome data
98	matrix by Brazeau [29], Davis et al. [30] and Zhu et al. [22]; most recently updated by Lu et al. [23],
99	Qiao et al. [26], Zhu et al. [27] and Burrow et al. [34]. Chondrichthyan content includes substantial
100	additions of new data drawn from Pradel et al. [14,15], Coates et al. [16], Coates & Tietjen [17], and
101	observation of original specimens (electronic supplementary material).
102	Phylogenetic Methods: The primary character matrix consists of 84 ingroup taxa and 2 outgroup
103	taxa (Galeaspida and Osteostraci) coded for 262 characters. Character and taxon sampling sources and
104	discussion are provided in electronic supplementary material. Phylogenetic analyses used maximum
105	parsimony implemented in PAUP*4.0.152 [35]. Nodal support was assessed via bootstrapping [36] and
106	Bremer Decay Indices [37], carried out using AutoDecay [38] and PAUP*. Details of phylogenetic
107	methods are provided in the electronic supplementary material. Character state transitions by node for the
108	strict consensus cladogram of the MPTs were reconstructed in PAUP* assuming hard polytomies with
109	DELTRAN [39] optimization (see Davis et al. [30], Coates et al. [16]).
110	A Principle Coordinate analysis (PCO) [40,41] was performed on the Hamming distance matrix
111	[42] of the character data. Computed dissimilarity was restricted to characters coded for both taxa, and
112	distances were normalized to the number of characters coded for both members in each taxon pair. For the
113	PCO, all characters were treated as equally weighted and unordered.
114	
115	
116	3. Results
117	(a) Specimen description, including (b) results of computed tomography, anatomical reconstruction
118	and thin section histology
119	Head length including the gill skeleton (figure $1a,b$ ) is ~21cm, and head plus trunk length as preserved

120 with the caudal region mostly absent is ~60cm (electronic supplementary material, figure S1), implying a

121 total body length of approximately 80cm.

122	Although considered one of the few 'unambiguous sharks' of the Lower and Middle Devonian
123	[32], details of Gladbachus anatomy do not conform in a straightforward manner with contemporary
124	models of early chondrichthyan anatomy. The internal skeleton consists of calcified cartilage with no
125	perichondral bone, yet the cartilage surface lacks the tightly connected tesserae that is a hallmark of
126	chondrichthyan skeletal anatomy [15,43]. Rather, most cartilage surfaces bear a mesh of continuously
127	calcified ridges (electronic supplementary material, figure S2), broadly resembling the 'wood-like' [44]
128	texture observed in some Mesozoic elasmobranchs. Discrete tesserae are visible only in the walls of the
129	semicircular canals, but these are irregularly sized and shaped, with broad intertesseral spaces. Thin
130	section histology and synchrotron microtomography show that poorly delineated tesserae are distributed
131	elsewhere in the skeleton, but concealed beneath the continuously mineralized cartilage surface.
132	The dermal skeleton includes no large plates. Head scales (figure $1d$ ) are mostly larger than trunk
133	scales (figure 1 <i>f</i> ) and the lateral line runs between scales. However, scale shape, composition and
134	histology are remarkable, as they resemble conditions observed in 'placoderms' [45,33], and differ
135	markedly from polyodontode scales like those of <i>Doliodus</i> [4,46] and mongolepids [47,48]. In
136	Gladbachus, scale and branchial denticle crowns consist of overlapping, mono-layered, cellular dentine
137	tubercles (electronic supplementary material, figure S3), lacking neck and basal canals. The standard,
138	total-group chondrichthyan scale growth pattern of areally apposed odontodes [32] is absent, as are
139	growing monodontode scales [49], and the non-growing placoid scales characteristic of modern
140	chondrichthyans. Instead, the reconstructed growth pattern of Gladbachus scales is linear and
141	bidirectional. Most unusually for a chondrichthyan, the cranial scales are asymmetric, with irregular and
142	inconsistent shapes. Fin spines, and spines associated with girdles and the flank region, are completely
143	absent.
144	The anterior section of the braincase is not preserved (figure $1a,b$ ), thus evidence of a precerebral
145	fontanelle is unknown, contra Heidtke and Kratschmer [31]. The right postorbital process includes traces of a
146	jugular canal, and a groove on the posterior surface, likely for articulation with the upper jaw (electronic
147	supplementary material, figure S4). The basicranium is compressed against the subjacent visceral arches, and

148 too poorly preserved to demonstrate presence or absence of a ventral cranial fissure, or canals for all or part of 149 the dorsal aorta network. Reconstructions of the vestibular, semicircular canals and ampullary spaces 150 (electronic supplementary material, figure S4c) demonstrate that the otic capsules were large and widely 151 separated across the midline. In extant gnathostomes, this degree of lateral separation is manifest only in 152 embryonic forms, and resemble adult conditions observed in 'placoderms' [50-53]. A pair of ring-shaped 153 structures flanking the dorsal ridge, next to the anterior lip of the persistent otico-occipital fissure probably 154 represents endolymphatic duct openings. This location for the endolymphatic ducts is consistent with the 155 absence of an endolymphatic fossa or single, median endolymphatic foramen, which characterizes all Recent 156 and fossil conventional chondrichthyans.

157 The jaws, hyoid arch, and gill skeleton are exceptionally complete (figures 1a,b and 2, electronic 158 supplementary material, figure S5), providing ready comparison with recently described in-group [15,54] 159 and out-group [55] examples. Mandibular arch morphology (electronic supplementary material, figure 160 S6) is more conventional than previously understood. *Contra* previous descriptions [31,56], there is no 161 palatal symphysis. The difference in length between the upper and lower jaws is considerable (figure 2e), 162 and comparable to conditions in *Acanthodes* [57,30], implying that a significant portion of the upper 163 dentition was born on the underside of the neurocranium; presumably on the internasal plate (cf. 164 *Ptomacanthus* [29]; *Doliodus* [5]). Notably, the scales and teeth (mixed) bordering the gape are preserved 165 as continuous, subparallel bands spanning the inter-orbital space (figures 1a,b). The palatoquadrate is 166 generally comparable to that of an early, conventional chondrichthyan (e.g. Orthacanthus [58]). The well-167 developed otic process bears a broad posterodorsal rim; the palatine process is broad and short, but there 168 is no evidence of a flange or process contributing to a palatobasal articulation (figure 2a). Preserved most 169 completely on the left side of the specimen, the anterior of the palate is thrust beneath the postorbital 170 process (figure 1a). The section of jaw visible in front of the preserved portion of the braincase is the 171 anterior extremity of Meckel's cartilage. The posterior portion of Meckel's cartilage is also exposed on the 172 dorsal surface of the specimen, but rotated through 90 degrees, such that the dorsal surface is compressed 173 against the mesial surface of the palatoquadrate.

174 The hyoid arch (figures 1a, b and 2c, d) is morphologically distinct from the gill arches. There is 175 no interhyal, and both the large and well-mineralized ceratohyal and slender first ceratobranchial 176 articulate with a broad basihyal. The five gill arches (electronic supplementary material, figure S7) are 177 positioned caudal to the braincase, as in non-holocephalan chondrichthyans. Epibranchials are present on 178 the first four arches, with anteriorly-directed simple pharyngobranchials (with no 179 suprapharyngobranchials) present in the first three gill arches (figures 1a and 2c) as in osteichthyans and 180 Ozarcus [15]. A pair of short, laterally directed cartilages medial to the base of the second gill provide the 181 only evidence of hypobranchials. The fifth arch ceratobranchials are unusually broad, nearly rectangular, 182 and keeled along the anterior margin. Remarkably, and uniquely, these resemble the posteriormost 183 ceratobranchials of *Paraplesiobatis*, a Lower Devonian 'placoderm' [55] (electronic supplementary 184 material, figure S8). The ceratobranchials of *Gladbachus* fourth and fifth gill arches articulate with a large 185 basibranchial copula, which is separated from the basihyal by a large gap. This revised description 186 provides the first accurate association of dorsal to ventral parts of each arch. 187 Although reported and named as toothless, Gladbachus possesses a dentition of small, mono-, bi-, 188 and tri-cuspid teeth lining the jaws, with branchial denticles lining gill arches I-IV (figure 1*a-c*, electronic 189 supplementary material, figure S3). The teeth are individually separate, and despite a suggestion of

lingual to labial alignment, there is no trace of whorl-like families as in conventional, non-holocephalan,chondrichthyans.

192

### 193 (c) Phylogenetic analysis and principal coordinates analysis.

194 Phylogenetic analysis of the data set returned 249,600 most parsimonious trees (TL = 691, CI = 0.396, RI

195 = 0.785, RCI = 0.311). The strict consensus cladogram of MPTs (figure 3*a*, electronic supplementary

196 material, figure S11) strongly corroborates recent phylogenetic hypotheses of early gnathostomes [23-27],

197 reconstructing all taxa usually referred to as acanthodians in a paraphyletic assemblage branching from

- 198 the chondrichthyan stem. Here, *Gladbachus* is also recovered as a stem-chondrichthyan, as the sister
- 199 taxon to a poorly resolved set of climatiid acanthodians and conventional chondrichthyans (including

crown clade Chondrichthyes). A monophyletic group uniting diplacanthid, ischnacanthid and acanthodid

201	acanthodians forms a clade that is the sister group of all other total-group chondrichthyans, including
202	Gladbachus.
203	The widely discussed Lower and Middle Devonian sharks Doliodus and Pucapampella branch
204	from close to the apex of the chondrichthyan stem. Pucapampella is recovered in an uncertain position
205	relative to several 'acanthodian' genera and the clade of conventional chondrichthyans. Doliodus is
206	recovered as a sister taxon to conventional chondrichthyans.
207	Contra Qiao et al. [26] and Zhu et al. [27], Ramirosuarezia is not recovered among 'acanthodian'
208	stem chondrichthyans, but rather, is nested among stem-gnathostomes, with other taxa, such as Qilinyu,
209	Entelognathus and Janusiscus branching crownward of the paraphyletic placoderms.
210	Within the chondrichthyan crown-group, xenacanths and ctenacanths (sensu lato) form a clade
211	branching from the elasmobranch stem. A further, poorly resolved cluster, including Homalodontus,
212	Tristychius, Acronemus and hybodontids branches from more crownward nodes, suggesting successive
213	sister groups to the elasmobranch crown and close relatives. Holocephalans include the symmoriids,
214	corroborating the arrangement found in Coates et al. [16].
215	A phylogenetic analysis on a reduced sample of chondrichthyans, focusing on the relationships
216	among stem members (figure 3b), recovered 24 MPTS, with a mostly resolved set of relationships among
217	acanthodians along the chondrichthyan stem. In this reduced analysis, Gladbachus again branches from
218	within the 'acanthodians', suggesting that the position of <i>Gladbachus</i> within this paraphyletic assemblage
219	is not the result of noise introduced into the data set by the large chondrichthyan sample.
220	Principal coordinates (PCO) analysis of the character data recovers all four of the traditional
221	gnathostome divisions ('placoderms', 'acanthodians', osteichthyans and chondrichthyans) as discrete
222	clusters in the space defined by the first three PCO axes (figure 3 <i>c</i> , <i>d</i> ). Notably, <i>Gladbachus</i> clusters with
223	chondrichthyans in the PCO, despite its phylogenetic position among 'acanthodians'. Gladbachus,
224	Doliodus and Pucapampella each occupy positions in PCO space between conventionally defined
225	chondrichthyans and 'acanthodians', however, these three genera are all significantly closer in PCO space

- to chondrichthyan taxa than to 'acanthodians' (t-tests of the intertaxon distances in PCO space for
- 227 *Gladbachus, Doliodus,* and *Pucapampella* yield p-values of 1.39\*10<sup>-6</sup>, 0.01, and 8.65\*10<sup>-8</sup>, respectively).
- 228
- 229

### 230 **4. Discussion**

231 (a) Tree shapes and implications for evolutionary timescale.

232 Due principally to its plesiomorphic scale conditions and absence of a dentition consisting of toothwhorls,

233 *Gladbachus* is reconstructed close to the base of chondrichthyan total-group (figure 3*a*,*b*), removed from

234 Doliodus and Pucapampella, which have traditionally been ascribed to the Chondrichthyes, but

interleaved among taxa normally referred to as 'acanthodians'. Accordingly, despite a phenetic similarity

236 to conventionally defined chondrichthyans, phylogenetically, *Gladbachus* is an acanthodian-grade stem-

chondrichthyan.

238 Support for the acanthodian branching pattern is weak (electronic supplementary material, figure 239 S11), but consistent with recent analyses [23-27]. Recent reconstructions have recovered traditional 240 acanthodian family-level sets: acanthodids, ischnacanthids, diplacanthids and climatiids (electronic 241 supplementary material, figure S9). In all of these trees, the climatilds group with conventional 242 chondrichthyans, echoing results of Brazeau [29] and Davis et al. [30]. Diplacanthids, ischnacanthids and 243 acanthodids fall into one of two arrangements, 1) as successive sister groups to more crownward taxa 244 [25,27], or 2) as in the present analysis, a monophyletic clade [23,24]. Here, we propose resurrecting the 245 term Acanthodii to define the diplacanthid-ischnacanthid-acanthodid clade. 246 Support for the chondrichthyan crown clade is strong (figure 3a), introducing new data for the 247 elasmobranch branch and corroborating the topology found in Coates et al. [16]. A time-calibrated

248 phylogeny using the strict consensus tree (figure 4) places the origin of the crown group at least as early

- as the end-Middle Devonian. The initial evolutionary radiation of crown chondrichthyans is primarily
- 250 post-Devonian, forming a significant component of the vertebrate recovery after the end-Devonian

251	Hangenberg extinction [59,60], which is evident from faunas recorded at Lower Carboniferous localities
252	such as Glencartholm [61], Bearsden [61], and Bear Gulch [62,63].
253	Conventional chondrichthyan conditions, exemplified by Doliodus [4], are present by the middle-
254	Lower Devonian (Pragian: ~410mya), and a minimum date for the origin of the chondrichthyan total-
255	group is currently tethered to the late Silurian (Ludlow: ~423 mya) by the earliest well-preserved
256	osteichthyan (Guiyu) [64]. However, the earliest 'acanthodian' stem-chondrichthyan body fossils
257	(Nerepisacanthus) are only slightly younger (Pridoli: ~419 mya) [65], and a wide variety of 'acanthodians'
258	are known from the Lower Devonian (e.g. Ptomacanthus, Brochoadmones, Cassidiceps and
259	Promesacanthus [29,66-69]). Such diversity, first apparent in the Lochkovian (figure 4) supported by a
260	taphonomically biased record of articulated specimens, implies either a sudden radiation in the early
261	Lower Devonian or a severely under-sampled history of Silurian stem-sharks. The latter hypothesis is
262	supported by isolated scales scattered through the Middle Ordovician to Silurian [48,49], including
263	strikingly characteristic, classically defined 'acanthodian' scales from the Rhuddanian (Llandovery
264	~440mya) of the Siberian Platform [70]. In the present phylogenetic context, we prefer to combine the
265	micro- and macro-/articulated fossil records, which strongly suggest that the chondrichthyan total group,
266	and, therefore, the gnathostome crown node, dates to at least the earliest Silurian, ~440 million-years-ago.
267	Thus, the early history of chondrichthyans consists of two phases (figure 4): a Silurian-Devonian
268	evolutionary radiation of micromeric, acanthodian-like taxa, and a subsequent Carboniferous radiation of
269	the crown clade, initially dominated by holocephalans [16,63].
270	

# 271 **(b)** Palaeobiological inferences

*Gladbachus* adds to an increasingly populated chondrichthyan stem lineage that also includes *Acanthodes* [24,30], *Ptomacanthus* [29,66], *Pucapampella* [8-10] and *Doliodus* [4-7]. The resultant data
on early chondrichthyan morphological diversity captures endoskeletal detail comparable to the content of
early osteichthyans, contributing to a more balanced interpretation of the initial gnathostome radiation.
However, there is no straightforward emerging sequence of character acquisition for the chondrichthyan

crown group (crown clade apomorphies are listed in electronic supplementary material, figure S10). The

277

278 current analysis highlights conflicting patterns of character-state distributions, implying repeated and 279 convergent evolution of chondrichthyan-like specializations among the earliest total-group members. For 280 *Gladbachus*, PCO analysis clearly identifies the chondrichthyan-like nature of its body plan (figure 3c), 281 but this stands in marked contrast to its reconstruction as representative of a previously unrecognized 282 'acanthodian' lineage (figures 3a, b and 4). Gladbachus approaches a quantifiably defined shark space, but 283 does so from a phylogenetically distinct origin (electronic supplementary material, figure S10). 284 The character combination observed in *Gladbachus*, alongside the array of contrasting conditions 285 observed in Early Devonian acanthodians (figure 4), defies conventional hypotheses of morphologically 286 segregated acanthodian and chondrichthyan morphotypes, reinforcing the hypothesis that pre-Devonian 287 stem-chondrichthyan diversity is fundamentally under-sampled. Reasons for the absence of substantial 288 Silurian remains of crown gnathostomes are unclear, although restricted environmental specificity has 289 been conjectured [71]. *Gladbachus* is a morphotypic outlier, in the sense that although phylogenetically 290 placed within the acanthodian grade, it lacks fin spines, its scales lack synapomorphies shared with any 291 acanthodian subgroup, and its estimated body length (electronic supplementary material, figure S1) is two 292 to three times greater than contemporary or earlier 'acanthodians', with the notable exception of 293 gyracanthids [60]. Furthermore, several features of the skeletal morphology suggest that *Gladbachus* was 294 a continuous ram suspension feeder [72], somewhat like modern basking sharks (*Cetorhinus*). The head, 295 including the gill skeleton, accounts for  $\sim 25\%$  of estimated total body length, the reconstructed oral 296 aperture is likely to have been in a near-perpendicular plane to the direction of forward movement, the 297 dentition is minimal, and the lower jaw is long and slender (figure  $2b_{c}$ ). To the best of our knowledge, 298 this is the earliest combination of such features known in any jawed vertebrate, adding to an emerging 299 picture of total-group chondrichthyans as early, nektonic specialists, in contrast to the reconstructed 300 demersal habits of their heavily skeletonized osteichthyan and 'placoderm' contemporaries [73]. Aspects 301 of this character-suite occur repeatedly among stem chondrichthyans, suggesting that the familiar gill slit

302	condition of sharks might originate from such early, and apparently multiple, natural experiments in
303	suspension feeding.
304	
305	
306	5. Conclusion
307	Gladbachus offers a glimpse of early chondrichthyan diversity yet to be discovered. Significantly,
308	Gladbachus scales, if discovered as isolated specimens, would be unrecognizable as chondrichthyan in
309	the new, total-group sense, unlike an increasing variety of Silurian and Ordovician [47-49,74] scale-based
310	taxa assigned with increasing confidence to the chondrichthyan total-group. Insights offered by
311	Gladbachus and other early chondrichthyans suggest that the morphological disparity in the early
312	members of the chondrichthyan total group was likely substantially greater than that which is observed in
313	the more-or-less stable shark-morphotype which has persisted from the Middle Devonian through to the

314 present. Accordingly, the importance of *Gladbachus* lies in its apparent morphological incongruence with

315 its phylogenetic position, hinting at multiple paths leading to the modern shark-like body plan.

316

317 Data accessibility. Data available from the Dryad Digital Repository:

- 318 https://doi.org/10.5061/dryad.84mh3
- 319

320 Authors' contributions. M.I.C. and J.A.F. conceived the idea and designed the research; K.E.C.

321 provided additional input. I.J.S. and P.S.A. provided all thin section histological analyses, related figure

322 preparations and comparative scale data. K.E.C. and M.I.C. completed initial CT renderings; M.K.T.

323 generated present CT renderings and produced the figures. K.E.C. provided comparative developmental

324 data. M.L.R. and P.J. La R. conducted the synchrotron CT scanning. J.A.F. and M.I.C. conducted

325 phylogenetic analyses; J.A.F. conducted PCO analysis. The manuscript was drafted by M.I.C. with

326 significant input from J.A.F., I.J.S. and K.E.C.

327

328 <b>Competing interests.</b> The authors declare no competing interest	ests.
---	-------

330	Fund	ling. This work was supported by grants DEB-0917922 and DEB 1541491 from the National		
331	Science Foundation (USA) (to MIC). GeoSoilEnviroCARS is supported by the National Science			
332	Foundation - Earth Sciences (EAR-1128799) and Department of Energy- GeoSciences (DE-FG02-			
333	94ER	14466). The Advanced Photon Source, a U.S. Department of Energy (DOE) Office of Science User		
334	Facili	Facility, is operated for the DOE Office of Science by Argonne National Laboratory under Contract No.		
335	DE-A	С02-06СН11357.		
336				
337	Ackn	owledgements. We thank J.A. Clack and M. Lowe for specimen loan, U. Heidtke for provision of		
338	materials recording early preparation of specimen, and K. A. Hardy for assistance with computed			
339	tomography renderings. We thank the GeoSoilEnviroCARS (Sector 13), Advanced Photon Source (APS),			
340	Argonne National Laboratory for access and support to complete significant portions of this work.			
341				
342				
343	References			
344	1.	Zangerl R. 1981 Chondrichthyes I. Handbook of Paleoichthyology 3A (ed. H-P Schultze) 115 pp.		
345		New York, USA: Gustav Fischer Verlag.		
346				
347	2.	Janvier P. 1996 Early Vertebrates. Oxford, UK: Oxford University Press.		
348				
349	3.	Stahl BJ. 1999 Chondrichthyes III: Holocephali. Handbook of Paleoichthyology 4 (ed. H-P Schultze)		
350		164 pp. München, Germany: Verlag Dr. Friedrich Pfeil.		
351				

352	4.	Miller RF, Cloutier R, Turner S. 2003 The oldest articulated chondrichthyan from the Early Devonian
353		period. Nature. 425, 501-504. (doi:10.1038/nature02001)
354		
355	5.	Maisey JG, Miller R, Turner S. 2009 The braincase of the chondrichthyan Doliodus from the
356		Lower Devonian Campbellton Formation of New Brunswick, Canada. Acta ZoolStockholm
357		Suppl. 90, 109-122. (doi:10.111/j.1463-6395.2008.00330.x)
358		
359	6.	Maisey JG, Turner S., Naylor GJP, Miller RF. 2013 Dental patterning in the earliest sharks:
360		implications for tooth evolution. J. Morphol. 2013, 1-11. (doi:10.1002/jmor.20242)
361		
362	7.	Maisey JG, Miller R, Pradel A, Denton JSS, Bronson A, Janvier P. 2017 Pectoral morphology in
363		Doliodus: bridging the 'acanthodian'-chondrichthyan divide. Am. Mus. Novit. 3875, 1-15.
364		(doi:10.1206/3875.1)
365		
366	8.	Maisey JG. 2001 A primitive chondrichthyan braincase from the middle Devonian of Bolivia. In
367		Major Events in Early Vertebrate Evolution (ed. PE Ahlberg), pp. 263-288 London, UK: Taylor
368		& Francis.
369		
370	9.	Maisey JG, Anderson ME. 2001 A primitive chondrichthyan braincase from the Early Devonian
371		of South Africa. J. Vertebr. Paleontol. 21, 702-713. (doi:10.1671/0272-
372		4634(2001)021[0702:APCBFT]2.0.CO;2)
373		
374	10.	Janvier P, Maisey JG. 2010 The Devonian vertebrates of South America and their biogeographical
375		relationships. In Morphology, Phylogeny and Paleobiogeography of Fossil Fishes (eds DK Elliot, JG
376		Maisey, X Yu, D Miao), pp. 431-459. München, Germany: Verlag, Dr. Freidrich Pfeil.
377		

378	11.	Maisey JG. 2005 Braincase of the Upper Devonian shark Cladodoides wildungensis
379		(Chondrichthyes, Elasmobranchii), with observations on the braincase in early chondrichthyans.
380		Bull. Am. Mus. Nat. Hist. 288, 1-103. (doi:10.1206/0003-
381		0090(2005)288<0001:BOTUDS>2.0.CO;2)
382		
383	12.	Maisey JG. 2007 The braincase in Paleozoic symmoriiform and cladoselachian sharks. Bull. Am.
384		Mus. Nat. Hist. 307,1-122.
385		
386	13.	Pradel A. 2010 Skull and brain anatomy of Late Carboniferous Sibyrhynchidae (Chondrichthyes,
387		Iniopterygia) from Kansas and Oklahoma (USA). Geodiversitas 32, 595-66.
388		(doi:10.5252/g2010n4a2)
389		
390	14.	Pradel A, Tafforeau P, Maisey JG, Janvier P. 2011 A new Paleozoic Symmoriiformes
391		(Chondrichthyes) from the Late Carboniferous of Kansas (USA) and Cladistic Analysis of Early
392		Chondrichthyans. PLoS ONE. 6, e24938. (doi:10.1371/journal.pone.0024938)
393		
394	15.	Pradel A, Maisey JG, Tafforeau P, Mapes RH, Mallatt JA. 2014 A Palaeozoic shark with
395		osteichthyan-like branchial arches. Nature. 509, 608-611. (doi:10.1038/nature13195)
396		
397	16.	Coates MI, Gess RW, Finarelli JA, Criswell KE, Tietjen K. 2017 A symmoriiform chondrichthyan
398		braincase and the origin of chimaeroid fishes. Nature. 541, 208-211. (doi:10.1038/nature20806)
399		
400	17.	Coates MI, Tietjen K. in press. The neurocranium of the Lower Carboniferous shark Tristychius
401		arcuatus (Agassiz, 1837). Earth Environ. Sci. Trans. R. Soc. Edinb.
402		

403	18.	Schaeffer B. 1981 The xenacanth shark neurocranium, with comments on elasmobranch monophyly.
404		Bull. Am. Mus. Nat. Hist. 169, 1-66.
405		
406	19.	Coates MI, Sequeira SEK. 1998 The braincase of a primitive shark. Trans. R. Soc. Edinb. (Earth
407		Sci.) 89, 63-85. (doi:10.1017/S026359330000701X)
408		
409	20.	Coates MI, Sequeira SEK. 2001a A new stethacanthid chondrichthyan from the Lower
410		Carboniferous of Bearsden, Scotland. J. Vertebr. Paleontol. 21, 438-459 (2001a).
411		(doi:10.1671/0272-4634(2001)021[0438:ANSCFT]2.0.CO;2)
412		
413	21.	Coates MI, Sequeira SEK. 2001b Early sharks and primitive gnathostome interrelationships. In
414		Major Events in Early Vertebrate Evolution (ed.PE Ahlberg), pp. 241-262 London, UK: Taylor
415		& Francis.
416		
417	22.	Zhu M, Yu X, Ahlberg PE, Choo B, Lu J, Qiao QL, Zhao J, Blom H, Zhu Y. 2013 A Silurian
418		placoderm with osteichthyan-like marginal jaw bones. Nature. 502, 188-193.
419		(doi:10.1038/nature12617)
420		
421	23.	Lu J, Giles S, Friedman M, den Blaawen JL, Zhu M. 2016 The oldest actinopterygian highlights
422		the cryptic early history of the hyperdiverse ray-finned fishes. Curr. Biol. 26, 1-7.
423		(doi:10.1016/j.cub.2016.04.045)
424		
425	24.	Brazeau MD, de Winter V. 2015 The hyoid arch and braincase anatomy of Acanthodes support
426		chondrichthyan affinity of 'acanthodians'. Proc. R. Soc. B 282: e20152210 (doi:10.1098/rspb.
427		2015.2210)
428		

429	25.	King B, Qiao T, Lee MSY, Zhu M, Long JA. 2016 Bayesian morphological clock methods
430		resurrect placoderm monophyly and reveal rapid early evolution in jawed vertebrates. Syst. Biol.
431		<b>2016</b> , syw107 (doi:10.1093/sysbio/syw107)
432		
433	26.	Qiao T, King B, Long JA, Ahlberg PE, Zhu M. 2016 Early Gnathostome Phylogeny Revisited:
434		Multiple Method Consensus. PLoS ONE. 11, e0163157. (doi:10.1371/journal.pone.0163157).
435		
436	27.	Zhu M, Ahlberg P, Pan Z, Zhu Y, Qiao T, Zhao W, Jia L., Lu J. 2016 A Silurian maxillate
437		placoderm illuminates jaw evolution. Science. 354, 334-336. (doi:10.1126/science.aaah3764)
438		
439	28.	Denison R. 1979 Acanthodii. Handbook of Paleoichthyology 3D (ed. H-P Schultze) 62 pp. New
440		York, USA: Gustav Fischer Verlag.
441		
442	29.	Brazeau MD. 2009 The braincase and jaws of a Devonian 'acanthodian' and modern gnathostome
443		origins. Nature. 457, 305-308. (doi:10.1038/nature07436)
444		
445	30.	Davis SP, Finarelli JA, Coates MI. 2012 Acanthodes and shark-like conditions in the last common
446		ancestor of modern gnathostomes. Nature. 486, 247-250. (doi:10.1038/nature11080)
447		
448	31.	Heidtke UHJ, Krätschmer K. 2001 Gladbachus adentatus nov. gen. et sp., ein primitiver Hai aus dem
449		Oberen Givetium (Oberes Mitteldevon) der Bergisch Gladbach – Paffrath-Mulde (Rheinisches
450		Schiefergebirge). Mainzer geowiss. Mitt. 30, 105-122.
451		
452	32.	Brazeau MD, Friedman M. 2014 The characters of Palaeozoic jawed vertebrates. Zool. J. Linn. Soc.
453		<b>170</b> , 779-821. (doi:10.1111/zoj.12111)

454		
454		
455	33.	Burrow CJ, Turner S. 2013 Scale structure of the putative chondrichthyan Gladbachus adentatus
456		Heidtke & Krätschmer, 2001 from the Middle Devonian Rheinisches Schiefergebirge, Germany. Hist.
457		Biol. 25, 385-390. (doi:10.1080/08912963.2012.722761)
458		
459	34.	Burrow CJ, den Blaauwen, J, Newman M, Davidson R. 2016 The diplacanthid fishes
460		(Acanthodii, Diplacanthiformes, Diplacanthidae) from the Middle Devonian of Scotland.
461		Palaeontol. Electronica. 19, 1-83.
462		
463	35.	Swofford DL. 2003 PAUP*. Phylogenetic analysis using parsimony (*and other methods). Version
464		4.0a147 Sunderland, MA: Sinauer Associates.
465		
466	36.	Felsenstein J. 1985 Confidence-limits on phylogenies - an approach using the bootstrap. Evolution.
467		<b>39</b> , 783-791.
468		
469	37.	Bremer K. 1988 The limits of amino-acid sequence data in angiosperm phylogenetic reconstruction.
470		Evolution. <b>42</b> , 795-803.
471		
472	38.	Eriksson T. 2001 AutoDecay Version 5.0.
473		
474	39.	Swofford DL, Maddison WP. 1987 Reconstructing ancestral character states under Wagner
475		parsimony. Math. Biosci. 87, 199-229.
476		
477	40.	Wills MA. 1998 Crustacean disparity through the Phanerozoic: comparing morphological and
478		stratigraphic data. Biol. J. Linn. Soc. 65, 455-500.
479		

480	41.	Davis JC. 1986 Statistics and Data Analysis in Geology. New York, USA: John Wiley & Sons.
481		
482	42.	Creanza N, Schwarz JS, Cohen JE. 2010 Intraseasonal dynamics and dominant sequences in H3N2
483		influenza. PLoS ONE. 5, e8544 (doi:10.1371/journal.pone.0008544)
484		
485	43.	Seidel R, Lyons K, Blumer M, Zalansky P, Fratzl P, Weaver JC, Dean MN. 2016 Ultrastructural
486		and developmental features of the tessellated endoskeleton of elasmobranchs (sharks and rays). J.
487		Anat. 2016, 1-22 (doi:10.1111/joa.12508)
488		
489	44.	Maisey JG. 2013 The diversity of tessellated calcification in modern and extinct chondrichthyans.
490		<i>Rev. Paléobiol.</i> <b>32</b> , 355-371.
491		
492	45.	Burrow CJ, Turner S. (1999) A review of placoderm scales, and their significance in placoderm
493		phylogeny. J. Vertebr. Paleontol. 19, 204-219 (doi:10.1080/02724634.1999.10011135)
494		
495	46.	Ginter M, Hampe O, Duffin C. 2010 Paleozoic Elasmobranchii: Teeth. Handbook of
496		Paleoichthyology 3D (ed. H-P Schultze) 168 pp. München, Germany: Verlag Dr. Friedrich Pfeil.
497		
498	47.	Karatajūtė-Talimaa V. 1998 Determination methods for the exoskeletal remains of early vertebrates.
499		Fossil Record. 1, 21-51.
500		
501	48.	Andreev PS, Coates MI, Shelton RM, Cooper RM, Smith MP, Sansom IJ. 2015 Upper
502		Ordovician chondrichthyan-like scales from North America. Palaeontology 58, 691-704.
503		(doi:10.1111/pala.12167)
504		

505	49.	Andreev PS, Coates MI, Karatajūtė-Talimaa V, Shelton RM, Cooper PR, Sansom IJ. 2017
506		Elegestolepis and its kin, the earliest monodontode chondrichthyans. J. Vertebr. Paleontol. 37,
507		e1245664 (doi:10.1080/02724634.2017.1245664)
508		
509	50.	Stensiö EA. 1963 Anatomical studies on the arthrodiran head (Part I). K. Sv. Vet. Akad. Handl. Ser. 4.
510		9, 1-419.
511		
512	51.	Young, GC. 1980 A new early Devonian placoderm from New South Wales, Australia, with a
513		discussion of placoderm phylogeny. Palaeontographica. A167, 10-76.
514		
515	52.	Goujet D. 1984 Les Poissons Placodermes du Spitsberg. Paris, France: CNRS.
516		
517	53.	Hu Y, Lu J, Young GC. 2017 New findings in a 400 million-year-old Devonian placoderm shed
518		light on jaw structure and function in basal gnathostomes. Scientific Reports. 7: 7813.
519		(doi:10.1038/s41598-017-07674-y)
520		
521	54.	Heidtke UHJ, Schwind C, Krätschmer K. 2004 Über die Organisation des Skelettes und die
522		verwandschaftlichen Beziehungen der Gattung Triodus Jordan 1849 (Elasmobranchii:
523		Xenacanthida). Mainzer geowiss. Mitt. 32, 9-54.
524		
525	55.	Brazeau MD, Friedman M, Jerve A, Atwood RC. 2017 A three-dimensional placoderm (stem-group
526		gnathostome) pharyngeal skeleton and its implications for primitive gnathostome pharyngeal
527		structure. J. Morphol. 2017, 1-9 (doi:10.1002/jmor.20706)
528		
529	56.	Heidtke UHJ. 2009 Gladbachus adentatus, die Geschichte des weltweit ältesten Hais - untersucht
530		und beschriebenaus dem AK Geowissenschaften. Pollichia Kurrier. 25, 24-26.

531		
532	57.	Miles RS. 1973 Relationships of acanthodians. In Interrelationships of Fishes (eds PH
533		Greenwood, RS Miles, C Patterson), pp. 63–103 London, UK: Academic.
534		
535	58.	Hotton N. 1952 Jaws and teeth of American xenacanth sharks. J. Paleontol. 26, 489-500.
536		
537	59.	Sallan LC, Coates MI. 2010 End Devonian extinction and a bottleneck in the early evolution of
538		modern jawed vertebrates. Proc. Natl. Acad. Sci. USA. 107, 10131-10135.
539		(doi:10.1073/pnas.0914000107)
540		
541	60.	Sallan LC, Galimberti AK. 2015 Body-size reduction in vertebrates following the end-Devonian mass
542		extinction. Science. 35, 812-815. (doi:10.1126/science.aac7373)
543		
544	61.	Dineley DL, Metcalf SJ. 1999. Fossil Fishes of Great Britain. Geological Conservation Review
545		Series 16 (ed. D Palmer) 675 pp. Peterborough, UK: Joint Nature Conservation Committee.
546		
547	62.	Lund R, Poplin C. 1999 Fish diversity of the Bear Gulch Limestone, Namurian, Lower Carboniferous
548		of Montana, USA. Geobios. 32, 285-295.
549		
550	63.	Grogan ED, Lund R, Greenfest-Allen E. 2012 The origin and relationships of early
551		chondrichthyans. In Biology of Sharks and Their Relatives (eds JC Carrier, JA Musick, MR
552		Heithaus), pp. 3-29 Boca Raton, USA: CRC Press.
553		
554	64.	Zhu M, Zhao W, Jia L, Lu J, Qiao T, Qu Q. 2009 The oldest articulated osteichthyan reveals a
555		mosaic of gnathostome characters. Nature. 458, 469-474. (doi:10.1038/nature07855)
556		

557	65.	Burrow CJ, Rudkin D. 2014 Oldest near-complete acanthodian: the first vertebrate from the
558		Silurian Bertie Formation konservat-Lagerstätte, Ontario. PLoS ONE. 9, e104171.
559		(doi:10.1371/journal.pone.0104171)
560		
561	66.	Brazeau MD. 2012. A revision of the anatomy of the early Devonian jawed vertebrate
562		Ptomacanthus anglicus Miles. Palaeontology. 55, 355-367. (doi:10.1111/j.1475-
563		4983.2012.01130.x)
564		
565	67.	Hanke GF, Wilson MVH. 2006 Anatomy of the Early Devonian acanthodian Brochoadmones
566		milesi based on nearly complete body fossils, with comments on the evolution and development
567		of paired fins. J. Vertebr. Paleontol. 26, 526-537. (doi:10.1671/0272-
568		4634(2006)26[526:AOTEDA]2.0.CO;2)
569		
570	68.	Gagnier, P-Y, Wilson MVH. 1996 Early Devonian acanthodians from northern Canada.
571		Palaeontology. <b>39</b> , 241–258.
572		
573	69.	Hanke GF. 2008 Promesacanthus eppleri n. gen., n. sp., a mesacanthid (Acanthodii,
574		Acanthodiformes) from the Lower Devonian of northern Canada. Geodiversitas. 30, 287–302.
575		
576	70.	Karatajūtė-Talimaa V, Smith MM. 2003 Early acanthodians from the Lower Silurian of Asia. Trans.
577		<i>R. Soc. Edinb. (Earth Sci.)</i> <b>92</b> , 277–299.
578		
579	71.	Sansom IJ, Andreev P. in press The Ordovician enigma: fish, first appearances and phylogenetic
580		controversies. In Evolution and Development of Fishes (eds Z. Johanson, M. Richter, C. Underwood).
581		Cambridge, UK; Cambridge University Press.
582		

583	72.	Sanderson SL, Wassersug R. 1993 Convergent and alternative designs for vertebrate suspension
584		feeding. In The Skull Volume 3: Functional and Evolutionary Mechanisms (eds J Hanken, BK Hall).
585		pp. 37-112 Chicago, USA: University of Chicago Press.
586		
587	73.	Blieck A. 2011 From adaptive radiations to biotic crises in Palaeozoic vertebrates: a geobiological
588		approach. Geol. Belg. 14, 203-227. (doi:10.1007/s12549-016-0260-1)
589		
590	74.	Sansom IJ, Davies NS, Coates MI, Nicoll RS, Ritchie A. 2012 Chondrichthyan-like scales from
591		the Middle Ordovician of Australia. Palaeontology 55, 243-247. (doi:10.1111/j.1475-
592		4983.2012.01127.x)
593		
594	75.	Cohen KM, Finney SC, Gibbard PL, Fan J-X. 2017 The ICS International Chronostratigraphic
595		Chart (2013; updated) Episodes 36, 199-204.
596		
597		

598	FIGURES
599	
600	
601	Figure 1. Gladbachus adentatus Heidtke and Krätschmer [31]. (a) Rendering of cranial and pectoral girdle
602	remains in dorsal view and (b) ventral view. (c) mandibular tooth; (d) cranial roof scale; (e) branchial
603	denticle; (f) trunk scale. All denticles and scales rendered semitransparent from micro-computed tomography
604	scans. Abbreviations: bhy, basihyal; chy, ceratohyal; hb, hypobranchial; mc, Meckel's cartilage; na, neural
605	arches; nc, neurocranium; or, orbital ring; pop, postorbital process; pq, palatoquadrate; sco, scapulocoracoid.
606	
607	
608	Figure 2. <i>Gladbachus adentatus</i> Heidtke and Krätschmer [31]. (a) Palatoquadrate; (b) Meckel's cartilage; (c)
609	jaws, hyoid arch and gill arches restored, dorsal view; (d) gills reconstructed ventral view; (e) reconstruction
610	of articulated mandibular arch, left lateral view. Abbreviations: adf, adductor fossa; bhy, basihyal; cbr,
611	ceratobranchial; chy, ceratohyal; cop, copula; ebr, epibranchial; end, endolymphatic duct; fm/oc, foramen
612	magnum/occipital cotylus; gl, glenoid; hb, hypobranchial; hy, hyomandibula; mp, mesial process; opr, otic
613	process; pbr, pharyngobranchial; pop, postorbital process; ppr, palatine process; pq, palatoquadrate; q,
614	quadrate condyle, unmineralized site of; ssc, semicircular canal network; ocf, otico-occipital fissure.
615	
616	
617	Figure 3. Phylogenetic placement of <i>Gladbachus adentatus</i> Heidtke and Krätschmer [31]. (a) parts i and
618	ii join at arrow heads. Strict consensus, complete tree. (b) Strict consensus of chondrichthyan total group
619	obtained from reduced taxon set; arrow head joins arrow head of (a) part <i>i</i> . Branch colours: black, stem
620	group gnathostomes; green, Osteichthyes; magenta, acanthodid stem Chondrichthyes; red, non-acanthodid
621	stem Chondrichthyes; purple, Holocephali (crown Chondrichthyes); blue, Elasmobranchii (crown
622	Chondrichthyes). Circles mark nodes with bootstrap support greater than 50% and/or decay values greater
623	than 1; filled circles mark nodes with bootstrap support greater than 75% and/or decay values greater than

624	3. (c) Phenetic results: PCO 1 (18.1% explained variance) is plotted on the vertical axis and PCO 2
625	(9.6%) is plotted on the horizontal axis. (d) PCO 1(vertical) vs. PCO 3 on the horizontal axis (8.1%).
626	Details of the PCO analysis are presented in the Supplementary Notes. The four traditionally named
627	groups (placoderms in purple, acanthodians in green, chondrichthyans in blue, osteichthyans in red)
628	cluster in distinct non-overlapping regions on the first three PCO dimensions. Relevant stem
629	chondrichthyan taxa are indicated in each plot.
630	
631	
632	Figure 4. Early chondrichthyan phylogeny: simplified strict consensus of phylogenetic analysis results
	<b>Figure 4.</b> Early chondrichthyan phylogeny: simplified strict consensus of phylogenetic analysis results calibrated against Ordovician-Carboniferous chronostratigraphic chart. Consensus computed from matrix with
632	
632 633	calibrated against Ordovician-Carboniferous chronostratigraphic chart. Consensus computed from matrix with
632 633 634	calibrated against Ordovician-Carboniferous chronostratigraphic chart. Consensus computed from matrix with 86 taxa and 262 characters. Taxon bar colour: black, non-chondrichthyan; magenta, acanthodid stem
<ul><li>632</li><li>633</li><li>634</li><li>635</li></ul>	calibrated against Ordovician-Carboniferous chronostratigraphic chart. Consensus computed from matrix with 86 taxa and 262 characters. Taxon bar colour: black, non-chondrichthyan; magenta, acanthodid stem chondrichthyan; red, non-acanthodid stem chondrichthyan; purple, holocephalan crown chondrichthyan; blue,

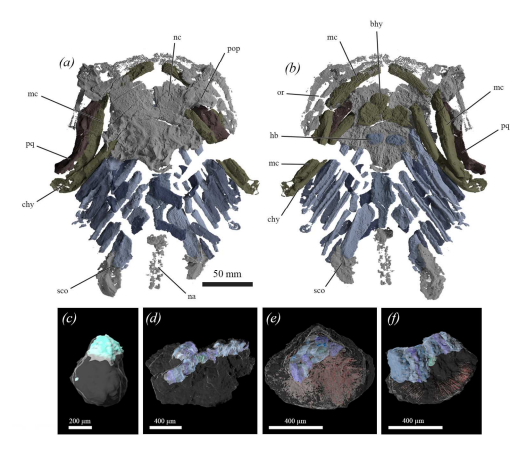
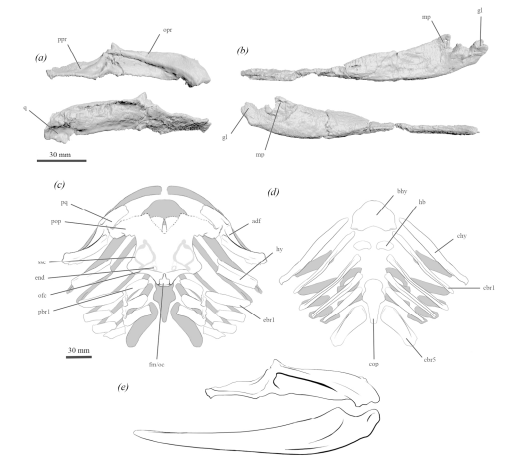


Figure 1 146x128mm (300 x 300 DPI)





184x174mm (300 x 300 DPI)

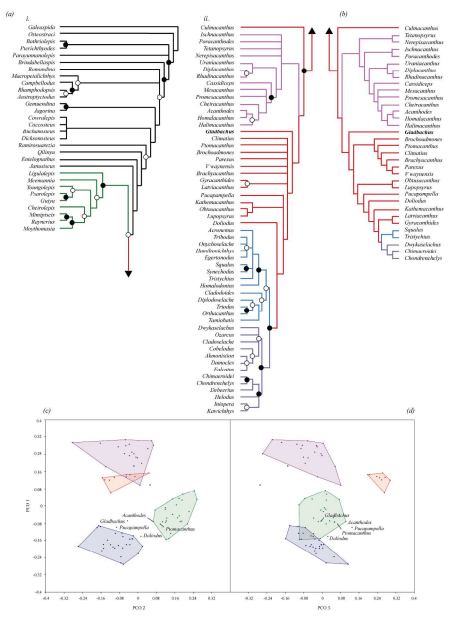


Figure 3

303x416mm (300 x 300 DPI)

