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A Tournaisian (earliest Carboniferous) conglomeratepreserved non-marine faunal assemblage and its environmental and sedimentological context

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A conglomerate bed from the Tournaisian Ballagan Formation of Scotland preserves a rich array of vertebrate and other non-marine fossils providing an insight into the wider ecosystem and palaeoenvironment that existed during this pivotal stage of Earth history. It challenges hypotheses of a long-lasting post-extinction trough following the end-Devonian extinction event. The fauna recovered includes a wide size range of tetrapods, rhizodonts and dipnoans, from tiny juveniles or small-bodied taxa up to large adults, and more than one taxon of each group is likely. Some fauna, such as actinopterygians and chondrichthyans, are rare as macrofauna but are better represented in the microfossil assemblage. The fauna provides evidence of the largest Carboniferous lungfish ever found. The specimens are preserved in a localised, poorly-sorted conglomerate which was deposited in the deepest part of a river channel, the youngest of a group of channels. In addition to the fossils (micro- and macro-), the conglomerate includes locally-derived clasts of palaeosols and other distinctive elements of the surrounding floodplains. Charcoal fragments represent stem and possible trunk tissue from arborescent pteridosperms. Preservation of the fossils indicates some aerial exposure prior to transport, with abrasion from rolling. The findings presented here contrast with other published trends in vertebrate size that are used to interpret a reduction in maximum sizes during the Tournaisian. The richness of the fauna runs counter to the assumption of a depauperate non-marine fauna following the end-Devonian Hangenberg event, and charcoal content highlights the occurrence of fire, with the requisite levels of atmospheric oxygen during that stage.

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21	of vertebrate and other non-marine fossils providing an insight into the wider ecosystem and
22	palaeoenvironment that existed during this pivotal stage of Earth history. It challenges
23	hypotheses of a long-lasting post-extinction trough following the end-Devonian extinction event.
24	The fauna recovered includes a wide size range of tetrapods, rhizodonts and dipnoans,
25	from tiny juveniles or small-bodied taxa up to large adults, and more than one taxon of each
26	group is likely. Some fauna, such as actinopterygians and chondrichthyans, are rare as
27	macrofauna but are better represented in the microfossil assemblage. The fauna provides
28	evidence of the largest Carboniferous lungfish ever found.
29	The specimens are preserved in a localised, poorly-sorted conglomerate which was
30	deposited in the deepest part of a river channel, the youngest of a group of channels. In addition
31	to the fossils (micro- and macro-), the conglomerate includes locally-derived clasts of palaeosols
32	and other distinctive elements of the surrounding floodplains. Charcoal fragments represent stem

Abstract

35 The findings presented here contrast with other published trends in vertebrate size that 36

are used to interpret a reduction in maximum sizes during the Tournaisian. The richness of the fauna runs counter to the assumption of a depauperate non-marine fauna following the end-Devonian Hangenberg event, and charcoal content highlights the occurrence of fire, with the

and possible trunk tissue from arborescent pteridosperms. Preservation of the fossils indicates

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41 **Key Words** Tetrapods, rhizodonts, dipnoans, chondrichthyans, charcoal, micropalaeontology, 42 sedimentology, paleobotany 43 44 Introduction 45 The 12-15 million years following the end-Devonian mass extinction, also known as the 46 Hangenberg Event, was thought to show a depauperate post-extinction trough for non-marine 47 vertebrates, especially tetrapods. The hiatus in the tetrapod fossil record became known as 48 'Romer's Gap', and covered the entire Tournaisian stage and much of the Viséan stage. 49 Recently, studies of the Tournaisian stage Ballagan Formation in southern Scotland and northern 50 England have cast doubt on that assessment (Smithson et al. 2012, 2015, Clack et al. 2016). Here 51 we present further evidence of the potential richness of the Tournaisian stage for our 52 understanding of the evolution of life on land. 53 The coastal locality of Burnmouth, Scotland, about 5 miles north of Berwick on Tweed, 54 has long been known to geologists as part of the Berwick monocline (e.g. McAdam, Clarkson & 55 Stone, 1992; Roper, 1997). The section at Burnmouth (Fig. 1) consists of strata that dip almost 56 vertically along about a 500 m exposure through the mainly Tournaisian strata of the Ballagan 57 Formation, formerly part of the Cementstone series in older terminology. Cementstones are also 58 now known as dolostones. 59 The Devonian-Carboniferous boundary has been identified at the top of the Kinnesswood 60 Formation near the base of the exposed succession (Millward et al. 2018, Marshall et al. 61 unpublished data), and the transition to the Viséan Fell Sandstones at its top. Its fossils have been 62 little studied until recently. The geology was described for the British Geological Survey by 63 Grieg (1988 p.49) who noted that plant fragments were common but fish fragments had only



64 been found in two beds. McAdam, Clarkson and Stone (1992) also mentioned non-marine 65 invertebrates, fish spines and plant remains in the Carboniferous section of Burnmouth. In a field guide to the locality in part based on Grieg's Memoir, Scrutton and Turner (1995) included a 66 67 generalized geological map on which they numbered the large sandstone bodies 1-14 (1 being 68 the youngest) that lie within the succession. They also noted that "fish fragments" had been 69 found in the succession, but gave few details (Scutton and Turner, 1995 pp 39 and 40). 70 In 1993, TRS began a more focused effort to examine Tournaisian rocks of Scotland for fossils. This formed part of an exploration of the Early Carboniferous in the Scottish Borders that 71 72 had begun in 1988 at Coldstream (Smithson et al., 2012). Eventually, he discovered not only fish 73 remains, but those of tetrapods. Subsequently, from 2006, more regular collecting was 74 undertaken with the late Stan Wood, and in 2012, a NERC-funded consortium project focusing 75 on the Tournaisian Ballagan Formation of Scotland, known as the TW:eed Project (Tetrapod World: early evolution and diversification), began an intensive exploration of Burnmouth and 76 77 other Tournaisian localities. Several papers relating to the Burnmouth sediments and associated 78 Tournaisian and other Early Carboniferous sites have already resulted from this project 79 (Smithson, Richards and Clack, 2015; Clack et al., 2016; Bennett et al., 2016; Kearsey et al., 80 2016; Smithson et al., 2017; Chen et al., 2018; Clack, Porro & Bennett, 2018 (SI1); Clark et al., 81 2018; Coates & Tietjen, 2018; Millward et al., 2018; Ross et al. 2018; Smithson & Clack, 2018 82 (SI2)), (Challands et al.; Marshall et al.; Millward et al.; unpublished data). 83 At least 11 beds with vertebrate bones including 7 containing tetrapods (Clack et al., 2016) have been discovered in the Burnmouth succession so far. One particularly rich bed and its 84 85 context is described by B.K.A. Otoo and members of the consortium team (Otoo, 2015; Otoo et 86 al., in press 2018 (SI3).

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This account describes the faunal association of a single bed from macro- and microfossils, its floral association from plant remains, and sedimentological context from detailed studies of a conglomerate lens observed within the section at Burnmouth. Until recently, this was one of the best exposed and most accessible conglomerates present within the section at Burnmouth. It lies about 383 metres above the Devonian-Carboniferous boundary (Clack et al., 2016, Clack, Porro & Bennett, 2018; Marshall et al., unpublished data), as exposed at Burnmouth, henceforward referred to as Bed 383. Assuming, with caution, that the Tournaisian succession exposed at Burnmouth accumulated in proportion to the 12 million years of the stage, this would give an approximate date of 350 Ma for Bed 383, about 9 MY after the end of the Devonian. Other beds at Burnmouth mentioned here will be referred to by their heights above the D/C boundary as defined there. Unfortunately, exposure of and access to the Bed 383 was obliterated by a large rock fall between late March and early April 2018. The fauna of Bed 383 includes numerous isolated bones, including both very large and very small examples, of tetrapod, lungfishes, rhizodonts, gyracanth spines and girdle elements, actinopterygians, a few teeth of the stem chondrichthyan Ageleodus, and chondrichthyan spines and scales. Many of the vertebrate specimens are fragmentary and indeterminate, while others can be identified to major clades or in a few cases more precisely. One tetrapod specimen is the partial lower jaw of a Crassigyrinus-like tetrapod (Clack, Porro & Bennett, 2018). The bed also contains charcoalified plant stems, charcoal fragments and wood fragments. However, the more fragile ostracod and bivalve fossils that are a common element of sedimentary rocks are absent. The conglomerate described is from a succession that has produced one of the richest assemblages of vertebrates from Burnmouth and in particular of tetrapods (Clack et al., 2016). It epitomizes the richness of Tournaisian vertebrate fauna and highlights the occurrence of fire, and



the requisite levels of atmospheric oxygen during that stage. Among the aims of this paper is to draw attention to this richness, and its importance as a window into the terrestrial communities that existed in the earliest Carboniferous non-marine environments. The existence of these specimens is important for inclusion in future paleobiology databases, to give a more accurate picture of life in the Early Carboniferous.

Materials and methods

Collecting specimens presented difficulties as the beds at Burnmouth are almost vertically orientated and Bed 383 occurs between the overlying massive sandstone and an underlying dolostone, close to the cliff face (Fig 1C). There is a high risk of losing important fragments into the gap between these two more competent beds generated by erosion of the relatively soft conglomerate. The entire collection was retrieved from a volume of no more than about a cubic metre, but because of the geological situation and with collections made on several separate occasions, no more accurate an estimate is possible. Many are no more than small fragments, but about half of them are identifiable to a major clade. Some specimens were collected individually from the bed, others were collected as bulk matrix which was disassembled in the laboratory. Most of the latter specimens were prepared manually from the heterogenous matrix with a mounted needle, dental mallet or air-pen. Thus the list of specimens does not represent an exhaustive inventory of those elements present, nor should the collection of macrofossils be used in quantitative faunal analyses.

Sedimentological observations were made using a combination of detailed logging, and tracing of surfaces and packages in the field at low tide, then linked to an interpretation of outcrop photographs, satellite images and 'drone' footage. The 'drone' footage was used to aid



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identification of key erosion surfaces and analyse the internal structures exposed at low tide across the wave cut platform. This detailed imagery was important for comparison with the aerial imagery (1 pixel = 10 x 10 cm) supplied by the North East Coastal Observatory that allows the architectural relationships of the exposed sandstone bodies to be interpreted without any distortion. Drone footage was taken with a DJI Phantom 2 Vision+ carrying a camera with a sensor size of 1.2/inches, 14 megapixels, a resolution of 4384 x 3288, HD recording of 1080p30 / 1080i60 and a recording FoV of 110° / 85°. All specimens are housed in the University Museum of Zoology, Cambridge (prefixed UMZC). Most specimens have numbers with the formulae 2017.1.x and 2018.2.x (about half of those also with the earlier formula are from a lower bed (Bed 340.5) described by Otoo et al. (in press 2018). Some specimens were registered from 2011 onwards. There are about 600 registered specimens from this bed so far, but this is still in progress. Samples were collected by TRS from 1993 to 2007, and the rest have been collected by TW:eed team members from 2007 to 2017. Photographs of most macrofossil specimens were taken by JAC using a Panasonic Lumix DMC-LZ5 and processed using Adobe Photoshop CS suites 2017.1.1 release. Others were photographed by JAC or TRS with a Dino-lite Pro AM4000 digital microscope. Some specimen-bearing samples were micro-CT scanned in the UMZC X-Tek scanner and segmented using Materialise's Interactive Medical Image Control System (MIMICS R) Research v18 (biomedical materialise com/mimics, Materialise NV, Leuven, Belgium). Scan data: 2017.2.187, Resolution/Voxel size 59.8178um, Filter 1mm copper, kV 190, uA 160, Slices 1984; 2017.2.189G, Resolution/Voxel size 23.9724um, Filter 1mm copper, kV 190, uA 160, Slices 1983; 2017.2.51, Resolution/Voxel size 49.9248um, Filter None, kV 190, uA 175, Slices1671.

For the micropalaeontology, a sample weighing 55.74 g from the base of the bed was processed overnight in a 5% solution of H_2O_2 . The residue was then wet sieved at 1000, 425, 250, 125, 65 μ m fractions and oven dried at 40°C. All fossil specimens present were picked from the 1000, 425, 250 and 125 μ m fractions and total counts are recorded in SI4.

For the palaeobotanical analysis, sediment samples around 10 g in weight were dissolved firstly in dilute HCl followed by concentrated HF in line with standard techniques (Scott, 2010, Glasspool & Scott 2013). Charcoal fragments were picked under a binocular microscope and selected fragments were mounted on aluminium stubs and studied using a Hitachi S2400 SEM (Scott, 2010).

Geological context

The succession at Burnmouth covers most of the Tournaisian stage, somewhat condensed, but extending from slightly below the Devonian-Carboniferous boundary, located at the top of the red sandstone Kinnesswood Formation, through to the base of the Visean Fell Sandstone. The bed is dated as within the CM (claviger-macra) palynozone, although the palynozonation of the Tournaisian stage in Scotland is being reviewed as a result of work undertaken as part of the TW:eed project (Marshall et al., unpublished data).

The specimens are associated with a sandstone body ca.383 m above the base of the Ballagan Formation at Burnmouth (Kearsey et al., 2016; Clack, Porro & Bennett., 2018).

Analysis reveals 6 distinct sedimentary packages (Fig. 2A) that are exposed across the wave-cut platform at low tide and are broadly equivalent to sandstone 4 of Scrutton and Turner (1995).

Packages 1-3 are ca. 2 m in thickness, and comprise fine- to medium-grained sandstones with slighter finer-grained tops. The bodies have basal erosion surfaces and shallow channelised

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forms. Package 1 is characterised by low-angle cross-stratification dipping north that passes into small-scale ripple cross-laminated structures as the body thins northwards. Dolostone beds (including A on Fig. 2A and other overlying dolostones that are not illustrated separately due to imagery resolution) above packages 1 – 3 are truncated by the erosion surface at the base of package 4. This erosion surface has ca. 4 m of relief. Package 4 is sandstone-dominated with generally low-angle stratification, whereas package 5 is a grey, poorly-exposed siltstone that overlies an erosion surface that clearly truncates adjacent and underlying beds.

A red-brown, fine- to medium-grained sandstone (package 6) overlies the youngest erosion surface and is the thickest sandstone (minimum 6.5 m, Fig 2A). The basal erosion surface cuts down to, but not through, dolostone beds (including A). Chondrites burrows occur in the dolostone beds underlying the sandstone succession at 383 m (Bennett et al., 2017. Package 6 (including Bed 383) is characterised by large-scale (broadly southerly) dipping stratification (depicted on Fig. 2). Cross-stratification (0.5 m scale) is observed between these surfaces particularly near the base. A 0.2 m-thick conglomerate occurs at the base of package 6 above this erosion surface and is now present only at the cliff though formerly it extended along the erosion surface onto the wave-cut platform exposures. This very localised conglomerate is poorly sorted, generally matrix supported and variably cemented with dolomite (Fig. 2B). Centimetre-sized sub-rounded to angular clasts (typically up to 30 mm in length and elongate) comprise red and grey siltstone, organic material, and bioclasts, including the partial lower jaw (UMZC 2011.9.1, Clack, Porro & Bennett, 2018) and a range of microfossils. Clasts sit within a coarse-grained sandstone to granule grade sandstone matrix, but clasts of the underlying dolostone are not identified in it. The specimens described in this paper derive from this conglomerate.



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Plant material within the sediment comprises generally small fragments, both charred and uncharred and is scattered within the matrix rather than occurring in layers. The charcoal samples comprise predominantly fragments 1-3 mm in length but larger pieces around 5 mm in length also occur. Better-preserved charcoal from a slightly lower bed (Bed 362) was used for comparison. **Results Tetrapods: Large elements Figs 3-5** Braincase Two braincase elements have been identified. The anteriormost part of a parasphenoid cultrifom process (2017.2.187 Fig. 3E) of a large tetrapod shows a weakly denticulate ventral surface. The dorsal surface, seen from micro-CT scans (Fig. 3E), is strongly concave, and the body is pierced by canals and other foramina of uncertain function (Fig. 3F). The element is similar in size to that of Crassigyrinus, although the cultriform process in that animal has only a small oval area of denticulation further posteriorly (Panchen, 1985). A basal process (2011.9.12 Fig. 3A) from the right side of the braincase is preserved with part of the basisphenoid. Grooves for the carotid arteries are clearly shown, as are the two facets of the basipterygoid process and its lining of dermal bone from the paraspheniod. Dorsally, a cavity probably represents part of the dorsum sellae. The element is large, and would indicate an individual larger than known *Crassigyrinus* specimens.

223 Skull bones



224 A robust but highly incomplete example of a tabular (2011.9.5 Fig. 3C) has worn but rugose 225 dermal ornament on the dorsal surface. The posterolateral margin carries a broken tabular horn, 226 grooved dorsally. The posterior margin carries a thin, narrow flange along its ventral edge, which 227 presumably contacted an extension from the postparietal. The posterior margin curves strongly 228 towards it medial edge, and with the postparietal would have formed a 'widow's peak' as 229 described in some embolomeres, but is more exaggerated in form. The bone is much larger than 230 the equivalent in the known specimens of *Crassigyrinus*. 231 Identified as a tetrapod postfrontal 2017.2.545 (Fig. 3B) has typically tetrapod pit and 232 ridge dermal ornament. It is rather strongly developed in the posterior part of the bone, but 233 lighter anteriorly. The bone is complete, with the orbital margin well defined, and sutural 234 surfaces on each of the other sides. It has a sutural surface anteriorly for the prefrontal, and there 235 is an excavation on the medial side for contact with the nasal. Medially is a contact for the 236 parietal, and posteriorly, two contacts, one presumably for the supratemporal and one more 237 laterally suggesting the presence of an intertemporal or a dorsal process of a postorbital. The 238 bone relationships more closely resemble those of Anthracosaurus (Panchen, 1977) than 239 embolomeres (Panchen, 1972) or *Eoherpeton* (Smithson, 1985). 240 241 Tooth 242 An isolated tooth (2017.2.384 Fig. 3D) is a blunt cone. It has labyrinthine infolding at the base, 243 and an acuminate tip. It does not resemble that of a rhizodont tooth and is interpreted as tetrapod. 244 245 Postcranial elements: axial skeleton



246 Centra, represented by two half-hoops (2017.2.374 Fig. 4D), show the typical morphology of 247 gastrocentrous intercentra, similar in size to those known from Crassigyrinus (Panchen 1985). 248 The ventral and lateral surfaces are lined with smooth periosteal bone. Internally the surfaces are 249 'unfinished'. 250 Neural arches are represented by one half of a neural arch (2017.2.189g Fig. 4 B,C), 251 probably an atlas arch somewhat similar to that of *Greererpeton* (Godfrey, 1989), and 252 2017.2.511, half of a neural arch (Fig 4.E). Specimen 2017.2.189g has little remaining evidence 253 of zygapophyses, and the neural spine is thin with the distal portion somewhat compressed, 254 leaving a transverse ridge across its base. Micro-CT scans show the mesial surface of unfinished 255 bone where it adjoined its antimere and the underlying centra. An atlas arch of this size probably 256 belonged to a moderately large individual, and it was found associated with the Crassigyrinus-257 like rib 2017.2.189a. 258 259 Postcranial elements: appendicular skeleton 260 Several large ribs or parts of ribs most closely resemble those of *Crassigyrinus*. A large example 261 (2017.2.189a Fig. 4A) has a striated surface. Its ends are somewhat crushed, but this seems to 262 result purely from compression that did not exaggerate the expansion of both ends. In the figure, 263 what is considered the proximal end is to the top. There is no apparent curvature, but that is usual 264 for cervical ribs in early tetrapods including Crassigyrinus, as is the expanded distal end 265 (Panchen, 1985). Rib 2017.2.49 (not figured) is of a similar size, but has lost both ends and is straight. The striation resembles that of 2017.2.189a, but identity as a lungfish rib is not ruled 266 267 out. 268 Several metapodials and phalanges including an ungual have been identified.



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269 A slightly asymmetrical autopodial bone (2017.2.388 Fig 4J, left pair) is smooth and convex in 270 dorsal view, and concave and grooved medially in ventral view. Others include 2017.2.380 (Fig. 4J, central image) exposed in ventral view only. It is more symmetrical but is also grooved 272 medially. A shorter element, 2017.2.61, is probably a phalanx (Fig. 4J right pair). A probable 273 ungual (2017.2.382) is exposed in dorsal view. It is flattened with an expanded proximal end and 274 a rounded distal end. It most closely resembles the unguals of *Greererpeton* (Godfrey, 1989). 275 Narrow body scales with longitudinal grooves and ridges very similar to those of 276 Crassigyrinus specimens (Panchen, 1985) have been found: 2017.2.107, 2017.2.175 and 277 2017.2.144 (Fig 4 F, G, H). 278 279 Small tetrapod elements 280 Alongside the large elements, are a few moderate sized and much smaller bones, showing the presence either of small-bodied individuals or juveniles. Delicate ribs, limb bones and small 282 phalanges have been found. 283 An almost complete jugal (2016.13 Clack et al., 2016) is exposed in internal view. The 284 postorbital region is relatively long, the orbit margin moderately deep, and the preorbital region 285 expands slightly anteriorly. The internal view shows no evidence of a lateral line. In terms of its 286 orbital component it falls somewhere between the shallow orbit of Ossirarus and the deep orbit with a narrow suborbital bar in *Diploradus*, and more closely resembles that of *Whatcheeria* in 287 288 proportion (Lombard and Bolt, 1995; Daeschler, Clack & Shubin, 2009) although this jugal is 289 from a much smaller individual. Whatcheeriid-like remains have been found elsewhere in the 290 Burnmouth sequence (Clack et al., 2016; Otoo, 2015; Otoo et al., in press 2018).

291 Two small partial jaws are interpreted as tetrapod on the basis of tooth shape and surface 292 ornament. Neither the smaller, 2017.2.117 (Fig. 5E) nor the larger, 2017.2.443 (Fig. 5H) is 293 identifiable to a higher taxon, but with other smaller elements, they show the presence of small 294 individuals. 295 A small tetrapod ulna 2017.2.51 (Fig 5C,E), is preserved in two counterpart pieces that 296 have been micro-CT scanned. It shows a moderate olecranon process and a thick cortex at mid-297 shaft. 298 Small, and some very small, ribs have been identified. A small almost parallel-sided rib 299 2017.2.476 (Fig. 5A) has an incipient uncinate process. Ribs 2017.2.149 (Fig. 5D) and 300 2017.2.170 are extremely slender and curved. 2017.2.170 (Fig. 5B) is more strongly so at its 301 thicker, and presumed proximal, end which is also grooved, and has a slight sigmoid bend at its 302 distal end. The rib head is not preserved. 303 304 **Lungfish Figs 6-10** 305 Skull bones 306 Conjoined J- and I- skull bones 2017.2.405 (Fig. 6A) have rugose ornament of pits and ridges 307 externally, but the strongly adhering matrix does not allow the lateral lines to be traced. The I-308 bone has a posterior process, which would have articulated with an anocleithrum. Internally, the 309 centres of growth can be seen from the direction of the striations. The sutures between the bones 310 are almost fused and difficult to trace in places. These bones would represent the skull of a very 311 large individual. 312 A large robust skull bone 2017.2.550 (Fig. 6H) is visible mainly in internal view. The 313 bone is complete except for wedge-shaped gaps at two places in the margin. Regions of sutural



overlap surface mark the edges indicating the contact with at least four different bones. The
possible course of a lateral line is indicated at one end by a groove and for the rest of its path, a
low smooth ridge. Its shape is comparable with the KL-bone of some Ctenodus specimens
(Sharp & Clack, 2013). The small area of exposed external surface shows low irregular
ornament.
A moderately large bone (2018.1.2 Fig. 6I) has an irregular hexagonal shape with sutural
overap areas on two sides, and an unsutured concave margin on another. It is identified as
lungfish on the lack of dermal ornamentation.
A probable lungfish skull bone 2017.2.72 (Fig 6B) has a smooth surface and lateral line
pits. It is a long oval with an overlap surface partly broken off. A rhizodont scale is adjacent. It is
not possible to identify the skull bone precisely within the skull.
Skull bone 2011.9.10 (Fig. 6E) is an irregular hexagon in internal view, with a more or
less smooth surface, whereas skull bone 2017.2.189i (Fig. 6C) is an oval bone with a plane
surface, no strong ornament, but marked with small pits.
Bones 2011.9.7 (Fig 6D), 2017.2.68, 2017.2.53 and 2017.2.547 (latter three not figured)
are all probably circumorbital bones, the first three from a large individual. 2011.9.7 shows
lateral line pores on the surface and has been micro-CT scanned (Fig 6G) to reveal the course of
the branches through the bone. All three bones are extremely robust. 2017.2.547 is more slender,
elongated and curved, showing two lateral line pores.
Numerous small skull bones are preserved, mainly hexagonal or somewhat elongated
with smooth surfaces and simple straight or curved margins, exemplified by 2018.2.193 (Fig.
6F).

337	Opercular series
338	Two very large opercula, 2017.2.188 (Fig. 7A) and 2017.2.667 (Fig. 7B) roughly similar in size,
339	both appear to represent the same side of an animal, so represent at least two large individuals.
340	The more complete 2017.2.188 is hexagonal in shape, with a small dorsally placed projection.
341	Specimen 2017.2.667 was collected as an in situ block positioned further seawards in the bed
342	than most of the other specimens, in a part of the bed that has now been completely removed by
343	erosion. Its matrix is more coherent with fewer inclusions than seen in the more cliff-wards
344	section, and may indicate one margin of the fossiliferous channel-fill. Potentially, the J+I- bones
345	and the circumorbitals are from an individual of a similar size to those relating to the opercula.
346	Smaller examples include 2018.1.5 (Fig. 7D) and 2011.9.6 (Fig.7C). The former is more
347	or less circular, whereas the latter is an elongate hexagon in shape. Alternatively, that may be a
348	suboperculum. A very small example 2017.2.720 (Fig. 7E) has a slightly striated surface.
349	A large suboperculum 2018.1.7 (Fig. 7F) is of a size commensurate with those of the
350	large operculars and 2018.1.5 (Fig. 7G) and 2017.2.543 (not figured) are parts of bones that may
351	be either subopercula or submandibulars.
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353	Shoulder girdle
354	Several elements of lungfish shoulder girdles are present, although only the anocleithrum is well
355	represented. The best preserved is 2017.2.57 (Fig. 8D), a large example. It is a large flattish bone
356	bearing two obvious processes, but otherwise fairly featureless. The broader of the two processes
357	is assumed to be that which contacted the I bone. The other has not been entirely exposed but is
358	narrower and more tapered. The presumed external surface onto which the cleithrum would have



fitted is not visible. It could have belonged to a similarly-sized individual as did the large operculum. Other smaller and incomplete examples are in the collection.

Partial clavicles and cleithra include 2017.2.70 (Fig. 8A), a partial clavicle and 2017.2.546 (not figured), the distal end of a clavicle. The clavicle is curved in cross section and probably represents the lower part of the bone, lacking the ventral and distal sections. Specimen 2017.2.546 (not figured) shows the rugose sutural surface for the cleithrum, with the external face showing low pustulate ornament. Alternatively, it may be the ventral end of the cleithrum. A small cleithrum is represented by 2017.2.494a (Fig. 8B).

A possible centrum element is 2017.2.398 (Fig. 8C). This bone consists of a shallow curved part that bears a grooved and ridged flange at right angles to it. A similar bone was indentified by the late Mahala Andrews (pers. comm. TRS 1979) from the Serpukhovian of the Dora Bone Bed near Cowdenbeath (see Smithson, 1985, for details of the excavation there). It does not resemble any element preserved on the articulated specimen of *Ctenodus interruptus* (Sharp & Clack, 2013) but is not otherwise identifiable.

Tooth plates

Five lungfish tooth plates are present. These include two incomplete specimens representing tooth plates approximately 24 mm long and three much smaller tooth plates up to 3.5 mm long. One of the largest specimens, 2014.1.2 (Fig. 9E) was referred to *Ballagadus rossi* (Smithson et al., 2015). It consists of a complete tooth ridge 1 and partial tooth ridges 2 and 3 from a right pterygoid tooth plate. Ridge 1 is 24 mm long and bears eight laterally compressed teeth. The six oldest teeth are worn to a narrow blade. Teeth seven and eight show little wear, are curved slightly anteriorly and separated from the older teeth by a distinct gap. The teeth on ridges 2 and

3 are longitudinally compressed. The second of the larger incomplete specimens 2017.2.54 (Fig. 9A) consists of three incomplete rows of teeth representing parts of ridges 1, 2 and 3. The teeth are similar in size and shape to those on 2014.1.2 with anteriorly curved, laterally compressed teeth on ridge 1 and longitudinally compressed teeth on ridges 2 and 3. The teeth are relatively large and probably formed part of the labial edge of the plate.

The three smaller tooth plates (Fig. 9B, C, D) are of similar size and morphology. The most complete is 2017.2.513 (Fig. 9D). It is approximately triangular-shaped, 3.5 mm long and 2.5 mm wide with a length to width ratio of 1.4:1. There are four tooth ridges radiating from a point immediately behind the posterior end of tooth ridge 1. The tooth ridge angle is approximately 80°. Ridge 1 has four laterally compressed teeth, ridges 2 and 3 have four longitudinally compressed teeth and ridge 4 has three. 2107.2.67 (Fig. 9C) is missing many of the labial teeth but has five ridges and the tooth ridge angle is approximately 110°. These smaller tooth plates are similar to those of *Ballagadus caustrimi* (Smithson, Richards & Clack, 2015) which have been found in two older beds at Burnmouth (Bed 333 and Bed 340.5).

Rhizodonts Figs 10-15

398 Skull roof

Postparietals include 2017.2.406 (Fig. 10B), which has part of its medial edge broken, but is otherwise similar in shape to that of *Strepsodus* (Andrews, 1985; Jeffery, 1999). It has the typical pustulate ornament externally as found in *Strepsodus* in this and other beds at Burnmouth. Others include 2017.2.186 (Fig. 10A), a postparietal with overall fine pustulate texture externally, but with irregular grooved 'vermiform' channels in which it is unlike that of 2017.2.406 or that of *Strepsodus* or *Screbinodus*. A probable partial postparietal, 2017.2.52 (Fig. 10E) has similar



405 surface texture to 2017.2.186, particularly on the longer, left side. The internal surface shows a 406 raised ridge. It is broader and shorter than that in 2017.2.406. A very small postparietal, 407 2017.2.643 (Fig. 10D), shows similar vermiform ornament invading a pustulate base, as 408 2017.2.186 and 2017.2.52. It is narrow with a straight edge, presumed to the medial edge 409 forming the midline. The lateral edge is somewhat broken, but may be a juvenile of that seen in 410 Fig 10D. 411 An extrascapula, 2017.2.94 (Fig. 10C), has tuberculate ornament which is eroded away 412 on the right hand side. 413 The stout hyomandibular bone, 2017.2.338 (Fig 10F), has a proximal articulation at an 414 angle of about 45° to the long axis of the bone. It appears relatively short compared to that of a 415 tristichopterid such as Eusthenopteron (Jarvik, 1980) and there is no sign of a foramen for the 416 chorda tympani. Although compressed, the bone shows a prominent longitudinal ridge down one side, and a groove, not quite opposite it, on the other. It is most similar to that of GLAHM 417 418 V8038 illustrated by Jeffery (1999) except that in the latter the proximal end appears incomplete, 419 and the distal end is wider. 420 421 Palate 422 A partial pterygoid, 2011.9.2 (Fig. 11), is one of the few known rhizodont pterygoids, which are 423 scarcely known in the literature. Young, Long and Ritchie (1992 figs 30 and 31)) described what 424 they considered to be the palatoquadrate and part of the lower jaw of the rhizodont Notorhizodon 425 but Johanson and Ahlberg (2001) concluded that these elements probably belonged to a 426 tristichopterid. However, 2011.9.2 bears ornament strikingly similar to that carefully illustrated 427 in Young, Long and Ritchie's figure 31. It consists of fine denticulations that morph gradually



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into vermiform grooves. There is an abrupt transition into a smooth overlap surface, exactly as depicted by Young and colleagues.

A problem arises in that tristichopterids are not known to occur in the Carboniferous, having become extinct at the end of the Devonian. Our specimen is thus unlikely to be a tristichopterid. An alternative explanation is that the awkward fit between parts of the jaw and palate described by Young and colleagues, and that was noted by Johanson and Ahlberg (2001), result from the parts belonging to different individuals rather than different taxa.

The bone described here is possibly much distorted. One end curves out of the plane of the denticulated surface and is smooth underneath. It appears to represent an articular surface, presumably part of the jaw joint. The surface then curves away from an edge rimmed by small teeth (unfortunately damaged during preparation and restored photographically from before and after photographs in Fig. 11D). Along the side opposite to the denticulated surface, the bone is folded downwards to a ledge, presumably also an overlap surface, and the row of small teeth runs along the edge of the fold. Comparison with the Young, Long and Ritchie's (1992) figure 31 shows that contrary to that bone, the articular end here is continuous with the denticulated ornament, whereas in the *Notorhizdon* bone, it is separated from the ornamented area by the smooth overlap surface. The relative positions of the row of small teeth and the denticulated region in 2011.9.2 is more similar to that of the palatoquadrate of *Notorhizodon*, although the entire extent of the more medial parts of the bone are missing in that specimen. There would have been some contact with the parasphenoid mesially, possibly though a quite extensive smooth overlap area, and the ledge beneath the row of small teeth was a contact for the marginal palatal bones.

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451 Other cranial elements

Mandibular bones include 2017.2.477 (Fig 12A), an elongate almost rectangular bone with pustular ornamentation best identified as an infradentary; 2017.2.404 (Fig. 11B) a tooth-bearing bone, probably a small, incomplete dentary with a symphysial tusk; and 2017.2.64 (Fig. 12E), an incomplete gular plate with pustular ornamentation.

Teeth are not usually well preserved in this bed, but a few isolated examples show characteristic features. Tooth 2017.2.99 (Fig. 12D) lacks its tip but shows the folded enamel at the base. The surface is damaged, but fine almost parallel striations are visible further away from the base. Tooth 2017.2.72b (Fig. 12C) has a complete tip, which shows no recurvature of the tip like that of *Strepsodus sauroides* (Jeffery, 2006). The striations more closely resemble those of *Archichthys*, but the tooth itself is more slender than those of the holotype series. Some of the external enamel is damaged, particularly along the convex (labial) surface, so that it is diffcult to say that it lacked striations as in that genus. The concave (lingual) surface shows fine parallel striations similar to those of 2017.2.99 and *Archichthys* (Jeffery, 2006).

Shoulder girdle elements

Cleithra (Fig 13) are among the elements more commonly recovered in the collection. A large example of a left cleithrum 2011.9.3 (Fig. 13C), is incomplete dorsally and ventrally. The external surface shows coarse meandering and anastomosing ridges on the lower blade and posterior edge, but is plain elsewhere. There is a deep groove along the posterior margin and up the middle of the mesial surface of the blade. The bone conforms with those of *Strepsodus* spp and *S. sauroides* (Andrews, 1985; Jeffery, 1999; Parker, Warren & Johanson, 2005).

473 An exceptionally large example, 2017.2.716 (Fig. 13A), in which the matrix has not been 474 completely removed from the external surface, where exposed shows similar ornament to 475 2011.9.3. A grooved and ridged cleithrum tip (not figured) collected at the same time and close 476 by does not fit onto the main specimen but was likely part of it. 477 The smallest cleithrum found in this bed is 2018.1.4 (Fig. 13B). It preserves only the 478 upper part of the blade. 479 Two incomplete cleithra, 2017.2.399 (Fig. 13D) and 2017.2.400 (see below), show 480 evidence of erosion by rolling during transport. They are similar to 2011.9.3 but are smaller. 481 482 Anocleithra Fig. 14 483 Two specimens are identified as rhizodont anocleithra, 2017.2.385 (Fig. 14B) and 484 2017.2.386 (Fig. 14A), by comparison with those illustrated by Parker, Warren and Johanson 485 (2005) for Strepsodus from Ducabrook, Australia, and an unidentified rhizodont from the Dora 486 Bone Bed, Cowdenbeath, Scotland. The two Burnmouth specimens differ somewhat in shape, 487 although the cleithral flange in the smaller example 2017.2.386 and the dorsal flange in the 488 larger example 2017.2.385 are both broken. Both have an anteroventral process, although that in 489 2017.2.386 is robust and rounded, whereas that in 2017.2.385 is pointed and more slender. Each 490 has a clear overlap region where the cleithrum would fit, but in 2017.2.386 it is defined by a 491 smooth rounded buttress, whereas in 2017.2.385 it is defined by a prominent ridge. Both are 492 probably from the left side of the animal, but their cleithral flanges have distinct processes in 493 differing positions. The lower margins of the cleithral flanges in the described examples are 494 highly variable. It is unlikely that these two belong to the same taxon. 495



496	Fin spines and scales
497	Fin spines both large and small are common in the bed. Fin spine 2017.2.494b, with about 75
498	mm preserved, shows a longtudinal groove, and tapers towards its distal end (Fig. 15F). Fin
499	spine 2017.2.568 (Fig. 15E) is circular in cross section along the approximately 80 mm of its
500	preserved length: it does not taper, but bears faint longitundinal striations. Cross sections show a
501	thick cortex with spongy medullary bone. It was originally identified as a plant stem. The best-
502	preserved examples of fin spines are probably attributable to rhizodonts, but may also pertain to
503	lungfish.
504	Rhizodont scales of many sizes are present, identified by the characteristic boss near the
505	centre of ossification (Fig. 15A-D) (Andrews, 1985).
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507	Gyracanths Fig. 16
508	Gyracanth spines are very common at this bed, although few are well preserved. Some eroded
509	examples were to be seen in the sandstone wall above this conglomerate bed prior to the rock fall
510	of 2018 (Fig. 16E,F). The most complete specimen collected is 2017.2.182, a pelvic spine
511	measuring 40 cm. The striations are narrow, close together, and bear closely spaced tubercles.
512	In other specimens such as 2017.2.185, an incomplete example, the ornamentation
513	appears coarser than in 2017.2.182. There are several other examples like this. Large
514	scapulocoracoid and procoracoid spines include 2011.9.11 and 2017.2.189b (Fig 16A,B).
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516	Other chondrichthyan elements Fig. 17



517	Agelodus teeth are occasionally found. The crown of Ageleodous tooth 2017.2.282 (Fig. 17A)
518	originally had 8 cusps, but one is missing. Cusps sit on a raised area, with the 'root' depressed
519	distally (see also the micropaleontology section for a different type of tooth).
520	Onychoselache-like spines are represented by 2017.2.397 (Fig 17B), a small spine
521	showing approximately longitudinal grooves and ridges, and 2017.2.51 (Fig. 17C) a smaller
522	example.
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524	Actinopterygian remains Fig. 18
525	Actinopterygian scales are not infrequent in this bed (see below), but are often fragmentary.
526	More complete examples include 2017.2.200 and 2018.1.3 (Fig 18, A, B). Isolated bones, though
527	rare, may include 2018.1.1 (Fig 18C).
528	
529	Coprolites Fig. 18
530	A few coprolites such as 2017.2.58 (Fig. 18D) have been found but cannot be associated with
531	any particular taxon.
532	
533	Elements of uncertain identity Fig. 19
534	A small tooth-shaped triangular element 2017.2.49 (Fig. 19A), probably flat on the unseen
535	surface, has a rounded profile towards the tapered end, but the wider end is characterized by a
536	raised horse-shoe-shaped rim with fluting, grooves and ridges. This bounds a depressed area. The
537	wider end is broken, and this bone may be a small part of a larger and more complex one.
538	Element 2017.2.327 (Fig 19B), appears to be from a rhizodont. It bears ornamentation
539	similar to skull bones, including some 'vermiform' grooves in a small area. One margin, the left



side in the figure, is strongly curved downwards. The other is broken. The ornamented area terminates in a distinct margin. Extending from this is an unornamented region with a smooth convex outline. It is marked on this surface by a deeply depressed area towards the side opposite the curved margin. On the underside, the whole surface is smooth, but there is a buttress beneath the curved margin that bounds a depressed and grooved area. There is no obvious distinction between where the ornamented segment of the external surface of the bone ends and the smooth area begins.

Suggestions for the identity of this bone include part of the shoulder girdle, a quadratojugal with attached quadrate, a surangular with attached articular, and a pterygoid with attached epipterygoid. We welcome other suggestions.

Abraded elements and plant macrofossils Fig. 20

Rolled bones such as 2017.2.214 of uncertain identity (Fig. 20B), appears smooth and featureless. It is convex in profile and slightly C-shaped. It may be part of a rib that has been rolled during transport. Other rolled and eroded bones include the two rhizodont cleithra listed above (Figs 13D, 20A).

Plant macrofossils are common with some large elements represented (Fig. 20C), suggesting that large elements of the flora existed during that part of the Tournaisian. However, they are usually poorly preserved such that identification is not usually possible. Some appear to be lycopod. They are often coated with iron oxide, as are some of the bones (see below).

Microfossil assemblage Figs 21, 22

In this study we compared the microfossil assemblage to the macrofossil assemblage in Bed 383, confining the identity of specimens to major groups. The micropalaeontological assemblage (SI4) is almost entirely composed of vertebrate material (5660 out of 5694 specimens), with some charcoal fragments (34 specimens). Microfossils are generally well preserved, although some abrasion, wear surfaces and cracks are observed. For vertebrates, 67.9% of specimens recovered have been identified. Rhizodont and actinopterygian material dominates the assemblage, with lesser occurrences of chondrichthyan and dipnoan microfossils (Fig. 21). Charcoal is identified by its black colour, brittle texture, fibrous external structure and hollow internal structure of preserved cellular tissue (see below and Scott, 2010).

Rhizodont scales have an exterior surface which is cream coloured, shiny and has a fibrous structure. When broken, the interior layers have a range of characteristic structural elements including sheets of either tubercules and pits or grooves and ridges that interlock together (Fig. 22A-B). These structures were identified by the examination of broken rhizodont scale macrofossils from this bed. 13.9% of the rhizodont scales have a red staining on the interior surfaces, likely due to the presence of iron oxide. Other rhizodont elements include rhizodont dermal bone (Fig. 22C) with a distinctive ornament of tubercules or ridges (cf. Jeffery, 2012) and three indeterminate rhizodont teeth (Fig. 22G), which have a shape similar to that of *Strepsodus*, but an ornament of well-defined striae similar to that of *Archichthys* (Jeffrey, 2006).

Actinopterygian scales have a rhombic shape with a smooth interior surface with keel, and a shiny, sculpted exterior surface (Fig. 22D) with transverse ridges or grooves (cf. Carpenter et al., 2014). Actinopterygian dermal bone and lepidotrichia fin bones (Fig. 22E) are also identified. Some of the dermal bone is similar to that of rhizodonts, with pustulate ornament on one side, but the fragments are small and they have a shiny, ganoine surface texture (Fig. 22F).



The lepidotrichia are most common in the 250 µm size fraction, so are more likely to be actinopterygian than rhizodont, although there are no physical characteristics to distinguish them other than size. They have a range of surface textures, from smooth to longitudinal striations or ridges. The interior medullary bone of the fine spines is infilled with calcite in some specimens. Actinopterygian teeth are present in all apart from the 1000 µm size fraction and are distinguished by their transparent apical cap (Fig. 22H) and shaft with cross-hatched surface ornamentation (Carpenter et al., 2011). Pharyngeal actinopterygian teeth are unornamented, curved, blunt at the tip, and occur in rows (Fig. 22I). In total 30 main teeth and 92 pharyngeal teeth are recorded.

Chondrichthyan teeth comprise three *Ageleodus* teeth with 5-6 tooth cusps (Fig. 22J) and one xenacanth (indeterminate) tooth (partly broken). Chondrichthyan material is dominated by denticles (140 specimens), which are common in the 1000 and 425 µm fractions and are mostly present as fragments in the 250 and 125 µm fractions. Three types of scales have been identified; elasmobranch (n = 43), hybodont (n = 28) and ?ctenacanth (n = 5). Indeterminate elasmobranch scales are identified by a flat or concave base and a top of curved spines (Fig. 22K), which in dorsal view form clusters of irregular height, or individual spines (Burrow, Long & Trinajstic, 2009; Carpenter et al., 2011). Hybodont scales have a concave base, spinose top and distinctive grouping of spines (Fig. 22L) which forms a single flat star shape, or multiple star shaped clusters in dorsal view (Garvey & Turner, 2006; Yazdi & Turner, 2000). Scales which can be putatively assigned to a ctenacanth origin have a flat base, spinose top and similar appearance to contemporaneous specimens described by Ivanov (1996) and Yazdi and Turner (2000). The spines are strongly curved, numerous and are of irregular height (Fig. 22M).

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Tetrapod microfossils have not been recorded from this bed and only one dipnoan element is identified (a toothplate fragment). This may result from problems in identification, given they are more common as macrofossils. Further direct comparative work of broken dipnoan and tetrapod macrofossils is necessary to determine their microfossil characteristics. Charcoal description and identification Figs 23-26 The charcoal fragments in Bed 383 range in size from a few millimetres to a centimetre in maximum dimensions (Fig. 23). All are woody comprising both secondary xylem and ray cell tissue. All specimens show evidence of pitting in the cell walls that often are circular pits that are arranged in single or in several rows (Fig. 23B-D). The cells also show evidence of homogenization (Fig. 23B) (Scott, 2010). It is often considered that secondary xylem (wood) that is dense or pycnoxylic is the most common feature of the arborescent pteridosperms (Galtier and Meyer-Berthaud, 2006). The tracheids are often less than 50µm wide and have short uniseriate rays but this is not easily visible. There is insufficient detail in the specimens found so far in Bed 383 to provide a definitive identifications. Charcoal fragments from Bed 362 (2018.4.13) are expected to preserved similar taxa to those of Bed 383, but they have a greater range of size, many specimens ranging from 10 to 20 mm in maximum length (Figs 24, 25). In addition, because of their larger size they show a wider

Charcoal fragments from Bed 362 (2018.4.13) are expected to preserved similar taxa to those of Bed 383, but they have a greater range of size, many specimens ranging from 10 to 20 mm in maximum length (Figs 24, 25). In addition, because of their larger size they show a wider range of characteristics including relatively small xylem cell sizes (Fig. 24 B-D; Fig. 25 C,D) but large numbers of wide rays (Fig. 24B). However, the specimens in Fig. 25 (A,B) and 26 (C) expose wide surfaces of longitudinal radial sections split at the level of the 'soft' ray tissue, demonstrating that rays were very high (i.e. tens of cells high). They may exhibit several types of wall pitting in the xylem as well as in the ray cells (Fig. 25 B,D). The pitting may comprise several rows of pits (Fig. 25D and occasionally bordered pits (Fig. 26C). In one specimen (Fig.

631 few rows of tracheids (Fig. 24 A,B). Cell wall homogenization is also seen (Fig. 26B). 632 Most of the charcoal appears to have come from stem and possible trunk tissue from a 633 range of arborescent pteridosperms (Galtier & Meyer-Berthaud, 2006). Such pteridosperms are 634 widely reported throughout the Mississippian of Scotland (see Scott et al., 1984; Scott & Galtier, 635 1988, 1996; Galtier & Meyer-Berthaud, 2006). Most common taxa that may be represented at 636 Burnmouth include the large tree *Pitus* that is also common in the Borders (Long 1979) and the 637 smaller Eristophyton (Scott & Galtier, 1996; Galtier & Scott, 1990). Other taxa reported from 638 Tournaisan rocks in the Midland Valley (Galtier & Meyer-Berthaud, 2006) include *Bilignia*, 639 Endoxylon, and Buteoxylon, although the size of the charcoalified specimens do not allow the 640 preservation of the combination of characters that will allow their certain identification. 641 Some of the specimens are of pycnoxylic wood that is compact with little parenchyma 642 between the rows of tracheids (Fig. 26A). Dadoxylon wood may also be a candidate as they have 643 uni- to biseriate rays. Calamopitys has been characterised as having manoxylic wood in the 644 stems (that is abundant parenchyma cells between rows of tracheids) and a thick cortex with a 645 range of cell sizes. The genus is more shrub-like with stems that are only 2-3 centimetres in 646 diameter but this genus has yet to be reported from the Tournaisian of Scotland. A related form 647 Stenomyelon has, however, been reported from Scotland (Scott & Galtier, 1988). All the specimens we have seen have a non-manoxylic wood and cannot be referred to calamopityans. In 648 649 Pitus (Pitvs) the tracheids of the secondary wood typically have pits arranged in multiseriate 650 rows (Gordon, 1935) and the trunks of this plant can be very large and may provide significant 651 quantities of wood that is available to produce charcoal.

24) there appears to be the presence of leaf traces (Fig. 24B) and also regular rays separating a



In many cases small fragments of wood charcoal are difficult to identify as the tracheid dimensions and pitting types occur in many different taxa. What is necessary are the ray size measurements and ray organization (number of cells wide and number of cells high) and this is only possible if, in addition to a good transverse section, radial and tangi-longitudinal sections are available. It is possible from the limited information available that the specimen in Fig. 24 has broad rays, but with no evidence that they are very high, that they may be comparable to *Eristophyton waltoni* ((cf. Galtier & Meyer-Berthaud, 2006, fig. 5E). Specimens in Figs 25 and 26 (possibly 26 but without ray data) have very high rays but no evidence they were broad (i.e. more than 3-4 cells wide) and therefore may be comparable to *Pitus antiqua* (Galtier & Meyer-Berthaud 2006, fig. 5H).

Discussion

Macrofossil Assemblage

The partial *Crassigyrinus*-like lower jaw, discovered in 2010 (UMZC 2011.9.1), from Bed 383 (Clack, Porro & Bennett, in press 2018). is about 20 Myr older than *C. scoticus*, and although attribution to the same genus is not unreasonable, it is unlikely to represent the same species. Also found in 2010 were a partial parasphenoid and a basal process, not identified as *Crassigyrinus*-like at the time. These braincase elements indicate an animal at least 50% larger than *C. scoticus*.

Earlier-collected material includes central elements: one complete centrum, one half of a centrum and one fragment. Phalanges or metapodials of a large tetrapod are also preserved, though their attribution to *Crassigyrinus* is less secure. Subsequent discoveries include ribs and body scales. Found in close proximity, probably in less than one square cubic metre of the 15-20

cm thick matrix, it is possible that many of these elements belonged to the same individual. The fact that few other elements have been found more recently suggests that most of the animal's remains had been eroded away before collecting began.

Smaller elements among the macrofossils include a jugal (Clack et al., 2016 fig. 6), ribs with uncinate processes and an ulna. These much smaller and moderate sized bones show the presence either of small-bodied individuals or juveniles. Also included are delicate ribs, limb bones and small phalanges. Their shapes are quite different from those in *C. scoticus* and probably do not belong to a juvenile of the same taxon as the lower jaw.

As with the tetrapods at this bed, lungfish elements indicate a wide range of sizes of individuals, from tiny juveniles or small-bodied taxa up to large adults. At least two very large individuals are represented by opercula within this relatively small accumulation. To our knowledge, these represent the largest lungfishes found in the Carboniferous. A rough guide to the length of the fish can be obtained by a ratio of 15:1 from the diameter of the operculum (TRS pers obs.). By this measure, the large opercula belonged to fish up to 3 m in length. Those from the later Carboniferous found so far are no more than about half the size as judged by this ratio. This finding is in contrast to the trends in vertebrate size documented by Sallan and Gallimberti (2015), who suggested a reduction in maximum sizes during the Tournaisian, followed by very slow recovery of larger forms later in the Carboniferous.

By contrast, no large tooth plates have been found, but several very small ones are present, including probable representatives of two species of *Ballagadus* (Smithson, Richards & Clack, 2015). Skull bones with varying surface ornamentation or none suggest that more than one taxon is present in the assemblage. Large elements with ornamented external surfaces and deeply interdigitated sutural surfaces contrast with small often oval or hexagonal elements with

smooth or pitted surfaces and no obvious sutural overlap areas. Whether these differences could be accounted for by ontogeny is not clear.

Rhizodonts are represented mainly by bones from large or medium-sized individuals, although small scales, a small cleithrum and a small postparietal have been recovered. Although many specimens resemble bones of *Strepsodus* it is possible that two taxa of rhizodont were represented in Bed 383.

There is some question of the taxonomic status of the rhizodonts from Burnmouth. Otoo (2015, Otoo et al. in press 2018) suggest that two taxa might be present in a somewhat older assemblage from Burnmouth, possibly *Strepsodus* and *Archichthys*. This requires further investigation, but the anocleithra and the postparietals (e.g. Figs10 A, B, E and 14 A, B) do suggest the presence of two taxa in Bed 383. An isolated humerus from a lower bed (Bed 340.5) under study by JAC and co-authors is quite different from those figured for *Strepsodus*.

Gyracanth spines are among the most common elements in this bed, though they are difficult to collect and prepare and they are often broken. Some elements are exceptionally large. Similar spines, though not as long as 2017.2.182 were found in the base of the overlying sandstone (probably those noted in McAdam, Clarkson and Stone (1972), although they had been subject to erosion on the suface of the exposed wall. They were lost in the rock fall.

Rare chondrichthyan remains suggest their presence in the region at the time, and are better represented in the microfossil assemblage (see below).

One of the issues encountered in this investigation is the difficulty of identifying isolated elements. This is particularly the case with possible rhizodont elements that are known from very few other taxa. Where possible we therefore illustrate here both internal and external surfaces of individual bones. Internal surfaces of skull bones are often not accessible except by micro-CT



scanning, so that it is important to illustrate both where circumstances allow. Our rhizodont skull bones also show unusual surface features of the ornament such as the vermiform grooves illustrated above. Similar grooving is known on some lungfish skulls, such as *Ctenodus interruptus* (Watson & Gill, 1923; Sharp & Clack, 2013) but it is not combined with an otherwise pustular surface. The condition in the postparietals illustrated here was confirmed as rhizodont by J. Jeffery from his personal observations, but it is not illustrated in the literature.

Unknown morphologies have also been uncovered. One of our specimens is a particular puzzle and does not fit easily into known categories, and a second, a rhizodont pterygoid, has challenged information from the published literature.

Microfossil assemblage

Given their limited representation in the macrofossil assemblage, an unexpected actinopterygian abundance is revealed in the micropalaeontological sample. In total, actinopterygian microfossils comprise 55.3 specimens per gram, while rhizodont microfossils only 11.5 per gram (Supplemental information). Actinopterygian microfossils are more common than rhizodont microfossils in the 250µm and 125 µm size fractions (Fig. 21). This would imply that actinopterygians are more abundant than rhizodonts, especially given their difference in size - rhizodonts are approximately ten times larger than actinopterygians, so could be expected to produce a lot more microfossil fragments. The greater fragility and breakage of actinopterygian scales compared to rhizodont scales counters this interpretation, along with the taphonomic concentration or winnowing of a certain size of material during transport. However, the teeth of actinopterygians and rhizodonts recovered in this study are of a similar size and there are approximately ten times the number of actinopterygian teeth (excluding pharyngeal teeth) than



rhizodont teeth. This indicates that the relative abundance of actinopterygians was probably higher than that of rhizodonts.

Interpretation and Environment of Bed 383

The 6 packages are interpreted as channelised river deposits, the channel forms being clear from the aerial views. Packages 1 – 3 are laterally adjacent channels and each one is successively younger, potentially representing shallow river channels laterally migrating through time (Fig. 2A). The dolostones overlying these three packages indicate a time during which a standing body of water developed, probably a lake (Bennett et al., 2016). Package 4 represents the return of a river system to this area, with the associated erosional relief suggesting a more significant system. Sediment deposited in this channel was partially eroded, but package 5 may represent a phase of channel abandonment and the deposition of fine-grained sediments. The youngest package (6) has the greatest erosional relief. Only one side of the channel is exposed, therefore the large-scale stratification could represent bar progradation or lateral migration of the channel itself.

The localised conglomerate probably represents deposition in the deepest part of this channel (package 6) through bedload processes. The sedimentology of the conglomerate lag (with clasts of red and grey siltstone) indicate floodplain sediments were incorporated, including clasts of sandy siltstones (Bennett et al., 2016) and palaeosols (Kearsey et al., 2016). Erosion by these river systems of the surrounding vegetated floodplains is clear from the erosional relief observed at this location. Several of the specimens show evidence of abrasion from rolling, and most are chaotically distributed in the matrix. The iron oxide staining of many rhizodont scales indicates that some rhizodont carcases may have been subject to sub-aerial exposure. Staining

may have occurred during the desiccation of a floodplain lake and oxidation of fossil material, prior to the incorporation of these sediments into the river system.

The presence of ichnofauna, rare scolecodonts, orthocones and evaporites (gypsum and anhydrite) within the Ballagan Formation evidence that the floodplain was subject to occasional marine water input (Bennett et al., 2017; Millward et al., 2018). The *Chondrites* in dolostones below Bed 383 has been interpreted as short-lived colonisation of coastal lake sediments by opportunistic organisms that were transported from shallow marine environments during storm events (Bennett et al. 2017). These *Chondrites*-bioturbated dolostones suggest the development of a significant floodplain lake, prior to the fluvial systems (4-6) being re-established. This is consistent with the complex coastal to alluvial palaeoenvironment that characterises the Ballagan Formation. Dolostone and evaporite clasts have not been incorporated into the conglomerate. Xenacanths are often associated with freshwater deposits, and Carboniferous elasmobranchs, ctenacanths, and *Ageleodus* are also interpreted as having a euryhaline salinity tolerance and occur in a range of marine to continental environments (Carpenter et al., 2015). Elasmobranchs are more commonly associated with freshwater sedimentary deposits than contemporaneous holocephalan chondrichthyans (Friedman & Sallan, 2012).

The presence of plant and charcoal debris confirm that vegetated floodplains were eroded and were subject to fire. However other evidence for subaerial floodplain ecosystems, e.g. myriapods, (Ross et al., 2018) are absent. The bedload transport process is likely to have broken and winnowed away arthropod cuticle and the fragile calcite shells of ostracods and bivalves, which are more common within floodplain facies (Bennett et al., 2016). The scarcity of tetrapod and dipnoan microfossil material may again reflect taphonomic processes related to the transport, concentration and deposition of fossil material. The presence of charcoal may indicate forest



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fires that increased the potential for erosion of the floodplain.

Burnmouth is a locality well known for the occurrence of anatomically preserved plants that were described in several papers by Albert Long (see Scott, Galtier & Clayton, 1984). The majority of the plants described were from dolostone facies of presumed late Tournaisian age obtained from loose pebbles. Scott, Galtier and Clayton (1984) reported in situ plants from nearby localities such as the shore section at Partanhall, Burnmouth (NT959614) and Scott and Rex (1987) illustrated a permineralized Stauropteris berwickense in a silty-wackstone horizon that also contained *Lepidodendron calamopsoides* axes together with fish bones and ostracods. The flora described from loose blocks included a range of lycopsids, peridosperms and ferns. A similar flora was described in dolostone facies from the River Whiteadder, for example at Edrom (Scott, Galtier & Clayton 1984; Scott & Rex, 1987). In addition to permineralized plants, some preserved as charcoal were also recovered (Scott & Rex, 1987). Compression floras, together with permineralized plants and charcoal were also reported from Foulden, also of late Tournaisian age, and relatively near Burnmouth (Scott & Meyer-Berthaud, 1985; Scott & Rex, 1987; Scott & Galtier, 1996; Scott & Glasspool, 2006). Fire has been an important influence on the Earth system throughout history (Scott, 2018). For there to be fires, there needs to be not only fuel to burn, and an ignition source (generally lightning) but also an atmospheric oxygen level of over 17% (Scott, 2010; Scott et al., 2014). The occurrence of charcoal in the sediments of this conglomerate allows two immediate conclusions. The first is that the oxygen level in the atmosphere was at least above 17% corroborating the hypothesis that the Tournaisian stage was not one subject to lower oxygen levels. Secondly, there is evidence of wildfire that affected the floodplains. However, the occurrence of charcoal in the sediments offers a further possibility. If there are rainstorms



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following a wildfire then the area of the fire can experience post-fire erosion (Moody & Martin, 2001; Cannon, 2004). This is when the roots of the plants have been killed and the sudden influx of water leads to a sudden erosion and movement of both the sediment and plant material (as well as animal material on the soil surface). This slurry may move very quickly (Scott, 2010) and form almost instantaneous alluvial fans away from the place of the wildfire (Moody and Martin, 2001; Scott et al., 2014). Such sediment may be either be transported into a river or lake system directly (as following the frequent Yellowstone fires) (Meyer & Pierce, 1993; Pierce, Meyer & Jull, 2004;) or else the sediments are quickly eroded by fluvial processes such as occurred in the Buffolo Creek fires of 1996 (Moody & Martin, 2001, 2009). Post-fire erosion/depositional systems have only recently been recognized in the fossil record. Extensive sediment pulses with charcoal have been described from the Early Carboniferous (Mississippian) of Ireland where there has been a sudden influx of sediment in to a marine near-shore environment with consequences for the fauna living there (Nichols & Jones, 1992; Falcon-Lang, 2000). However, the occurrence of vertebrates that have been caught up in such post-fire flood events have been described in the Triassic (Havlik et al., 2013) and Cretaceous by several authors (Brown at al, 2013; Muir et al., 2014). The vertebrate remains alongside the charcoal, and the greater depth of erosion observed associated with package 6 suggest that post-fire erosion/deposition may have played a role in its formation. Charcoal often floats and takes time to settle in a water column, depending on size, type of organ and temperature of formation and hence can travel considerable distances by water transport (Nichols et al., 2000; Scott, 2010). Charcoal is often deposited by currents of a certain velocity where fragments are also well sorted (Nichols et al., 2000). The occurrence of charcoal and uncharred plants with siltstone and sand-sized grains alongside larger pebbles (palaeosols



and sandy siltstone) could suggest a more chaotic flow and rapid deposition for at least this part of the bed. However, the stratification in the channel fill above the conglomerate indicates large-scale bed-forms and sustained period of sandstone deposition. The widespread occurrence of charcoal suggests that fire appears to have been a regular occurrence through the Mississippian of Berwickshire and across the south of Scotland and to have had an impact on both the vegetation and the landscape (Scott, 1988; Brown, Scott & Jones, 1994; Scott & Glasspool, 2006).

Conclusions

This study has described and interpreted the fauna of an Early Carboniferous river system and aspects of its surroundings and environment. It has provided further evidence to demonstrate that the Tournaisian stage of the Early Carboniferous was by no means a depauperate time in the evolution of vertebrate life on land. Neither was it a time of low oxygen concentration. Instead, the richness of a single small channel-fill conglomerate demonstrates the diversity of vertebrate taxa and their potential abundance in previously understudied Tournaisian non-marine fossil localities. It also shows that large-bodied taxa, in some cases larger than those from later in the Carboniferous, lived in the Tournaisian, about 9 MY after the end of the Devonian. Thus large forms appear to have recovered more quickly following the Hangenberg extinction event than previous estimates of up to 36 MY (Sallan & Gallimberti, 2015) had suggested. The existence of the so-called 'Romer's Gap' thus largely resulted from lack of exploration of such localities, and efforts should be made to target equivalent localities in other parts of the world.

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872	Data archiving statement
873	Data currently available on
874	https://www.dropbox.com/s/ge4t18smipfehid/Dryad%20data%20Clack.zip?dl=0
875	On acceptance, data for this study will be available in the Dryad Digital Repository
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877	References
878	Andrews, S.M. 1985, Rhizodont crossopterygian fish from the Dinatian of Foulden,
879	Berwickshire, Scotland, with a re-evaluation of this group. Transactions of the Royal
880	Society of Edinburgh: Earth Sciences, 76, 67–95.



881	Bennett, C, Kearsey, T., Davies, S., Millward, D., Clack, J., Smithson, T., Marshall, J. 2016.
882	Early Carboniferous sandy siltstones deposited in seasonal flooding episodes preserve rare
883	non-marine fossils. Sedimentology 63, 1677-1700 https://doi.org/10.1111/sed.12280
884	Brown, S.A.E., Collinson, M.E., Scott, A.C. 2013. Did fire play a role in formation of dinosaur-
885	rich deposits? An example from the Late Cretaceous of Canada. Palaeobiodiversity and
886	Palaeoenvironments. 93, 317-326. Doi: 10.1007/s12549-013-123-y
887	Brown, R.E., Scott, A.C., Jones, T.P. 1994. Taphonomy of fossil plants from the Viséan of East
888	Kirkton, West Lothian, Scotland. Transactions of the Royal Society of Edinburgh: Earth
889	Sciences 84, 267-274.
890	Burrow, C.J., Long, J.A., Trinajstic, K., 2009. Disarticulated acanthodian and chondrichthyan
891	remains from the upper Middle Devonian Aztec Siltstone, southern Victoria Land,
892	Antarctica. Antarctic Science, 21, 71-88.
893	Cannon, S.H., 2001. Debris flow generation from recently burned watersheds. <i>Environmental and</i>
894	Engineering Geoscience 7, 321-341.
895	Carpenter, D.K., Falcon-Lang, H.J., Benton, M.J., Nelson, W.J. 2011. Fishes and Tetrapods in the
896	Upper Pennsylvanian (Kasimovian) Cohn Coal Member of the Mattoon Formation of
897	Illinois, United States: Systematics, Paleoecology, and Paleoenvironments. Palaios, 26,
898	639–657.
899	Carpenter, D.K., Falcon-Lang, H.J., Benton, M.J., Henderson, E. 2014. Carboniferous
900	(Tournaisian) fish assemblages from the Isle of Bute, Scotland: systematics and
901	palaeoecology. Palaeontology 57, 1215-1240.

902 Chen, D., Alavi, Y., Brazeau, M., Blom, H., Millward, D., Ahlberg, P. E. 2018 Earth and 903 Environmental Transactions of the Royal Society of Edinburgh. https:// doi.org/10.1017/S1755691018000099 904 905 Clack J. A., Porro L.B., Bennett, C.E. In press 2018. A Crassigyrinus-like jaw from the 906 Tournaisian (Early Mississippian) of Scotland. Earth and Environmental Transactions of 907 the Royal Society of Edinburgh. 908 Clack, J. A., Bennett, C. E., Carpenter, D. K., Davies, S. J., Fraser, N. C., Kearsey, T. I, Marshall, 909 J. E. A., Millward, D., Otoo, B. K. A., Reeves, E. J., Ross, A. J., Ruta, M., Smithson, K. Z., 910 Smithson, T. R. Walsh, S. 2016 Phylogenetic and Environmental Context of a Tournaisian 911 Tetrapod Fauna., Nature Ecology and Evolution. doi:10.1038/s41559-016-0002. 912 Clark, N.D.L. Miller, R.F., Ross, A.J. The distribution of *Schamocaris* (Eumalacostraca, 913 Crustacea) along the northwestern coast of the Rheic Ocean during the Lower 914 Carboniferous. Earth and Environmental Transactions of the Royal Society of 915 Edinburgh. 107, 91-98. https://doi.org/10.1017/S1755691018000159 916 Coates, M.I., Tietjen, K. 2018 The neurocranium of the Lower Carboniferous shark Tristychius 917 arcuatus (Agassiz). Earth and Environmental Transactions of the Royal Society of 918 Edinburgh. https://doi.org/10.1017/S1755691018000130 919 Daeschler, E. B., Clack, J. A., Shubin, N. H. 2009. Late Devonian tetrapod remains from Red Hill, 920 Pennsylvania, USA: how much diversity? Acta Zoologica 90: 306-317. 921 Falcon-Lang, H. 1998. The impact of wildfire on an early Carboniferous coastal system. 922 Palaeogeography, Palaeoclimatology, Palaeoecology 139, 121-138. 923 Garvey, J.M., Turner, S., 2006. Vertebrate microremains from the presumed earliest 924 Carboniferous of the Mansfield Basin, Victoria. *Alcheringa*, 30, 43-62.

Press.

925 Galtier, J. and Mever-Berthaud, B. 2006. The diversification of early arborescent seed ferns. 926 Journal of the Torrey Botanical Society 133(1), 7-19 927 Galtier, J., Scott, A.C.1990. On *Eristophyton* and other gymnosperms from the Lower 928 Carboniferous of Castleton Bay, East Lothian, Scotland. Geobios 23, 5-19. 929 Glasspool, I.J., Scott, A.C. 2013. Identifying past fire events. in Belcher, CM ed. Fire Phenomena 930 in the Earth System – An Interdisciplinary Approach to Fire Science. Oxford: J. Wiley and 931 Sons 179-206 932 Godfrey, S. J. 1989. The postcranial skeletal anatomy of the Carboniferous tetrapod *Greererpeton* 933 burkemorani Romer 1969. Philosophical Transactions of the Royal Society of London (B) 934 323: 75-133. 935 Gordon, W.T. 1935. The genus Pitys, Witham, emend. Transactions of the Royal Society of 936 Edinburgh 58: 279-311. 937 Greig, D. C. 1988 Geology of the Eyemouth district. *Memoirs of the British Geological Survey*, 938 Keyworth, UK. 939 Havlik, P., Aiglstorfer, M., El Atfy, H., Uhl, D. 2013. A peculiar bonebed from the Norian 940 Stubensandstein (Löwenstein Formation, Late Triassic) of southern Germany and its 941 palaeoenvironmental interpretation. Neues Jahrbuch Für Geologie und Paläontologie – 942 Abhandlungen. 269/3, 321–33 943 Ivanov, A., 1996. The Early Carboniferous chondrichthyans of the South Urals, 944 Russia. Geological Society, London, Special Publications, 107: 417-425.

945 Jarvik 1980 Basic Structure and Evolution of Vertebrates. Volumes 1 and 2. New York, Academic



947 Jeffery J. E. 1999 The morphology and phylogeny of the European members of the Order 948 Rhizdontida (Pisces, Sarcopterygii). D. Phil. thesis, University of Cambridge. 949 Jeffery, J.E., 2006. The Carboniferous fish genera *Strepsodus* and *Archichthys* (Sarcopterygii: 950 Rhizodontida): clarifying 150 years of confusion. *Palaeontology*, 49, 113-132. 951 Jeffery, J.E., 2012. Cranial morphology of the Carboniferous rhizodontid Screbinodus ornatus 952 (Osteichthyes: Sarcopterygii). Journal of Systematic Palaeontology, 10, 475-519. 953 Johanson, Z., Ahlberg, P.E. 2001. Devonian rhizodonts and tristichopterids (Sarcopterygii: 954 Tetrapodomorpha) from East Gondwana. Transactions of the Royal Society of Edinburgh: 955 Earth Sciences 92, 43-74. 956 Kearsey, T.I., Bennett, C.E., Millward D., Davies, S.J., Gowing, C.J.B, Kemp, S.J, Leng, M.J. 957 Marshall, J.E.A, Browne, M.A.E. 2016 The terrestrial landscapes of tetrapod evolution in 958 earliest Carboniferous seasonal wetlands of SE Scotland. *Palaeogeography*, 959 Palaeoclimatology, Palaeoecology 457, 52-69 960 https://doi.org/10.1016/j.palaeo.2016.05.033 961 Lombard, R. E., Bolt, J. R. A new primitive tetrapod, Whatcheeria deltae, from the Lower 962 Carboniferous of Iowa. *Palaeontology* 38: 471-494. 963 McAdam, A. D., Clarkson, E. N. K., Stone, P. 1992 Scottish Borders Geology: an excursion 964 guide. Edinburgh: Scottish Academic Press, for Edinburgh Geological Society http://earthwise.bgs.ac.uk/index.php/Burnmouth - an excursion 965 966 Meyer, G.A., Pierce, J.L. 2003. Climatic controls on fire-induced sediment pulses in Yellowstone 967 National Park and central Idaho: a long term perspective. Forest Ecology and Management 178, 89–104. 968

- 969 Millward, D., Davies, S.J., Williamson, F., Curtis, R., Kearsey, T.I., Bennett, C.E., Marshall,
- J.E.A., Browne, M.A.E. 2018 Early Mississippian evaporites of coastal tropical wetlands.
- 971 *Sedimentology* 1-31 doi: 10.1111/sed.12465
- 972 Moody JA, Martin DA. 2001 Initial hydrologic and geomorphic response following a wildfire in
- 973 the Colorado Front Range. Earth Surace Processes and Landforms 26, 1049–1070.
- 974 Moody, J.A., Martin, D.A. 2009. Forest fire effects on geomorphic processes. In: Cerd_a, A.,
- Robichaud, P. (Eds.), Fire Effects on Soils and Restoration Strategies, pp.41–79. Enfield,
- 976 NH: Science Publishers, Inc.
- 977 Muir, R. A., Bordy, E. M., Prevec, R. 2015. Lower Cretaceous deposit reveals first evidence of a
- post-wildfire debris flow in the Kirkwood Formation, Algoa Basin, Eastern Cape, South
- 979 Africa. Cretaceous Research 56, 161-179.
- 980 Nichols, G., Cripps, J., Collinson, M.E., Scott, A.C. 2000. Experiments in waterlogging and
- sedimentology of charcoal:results and implications. *Palaeogeography, Palaeoclimatology*,
- 982 *Palaeoecology*, 164, 43-56.
- 983 Nichols, G., Jones, T. P. 1992. Fusain in Carboniferous shallow marine sediments, Donegal,
- 984 Ireland: The sedimentological effects of wildfire. *Sedimentology* **39**, 487-502.
- 985 Otoo, B.K.A. 2015. A taxonomic and Palaeoecological investigation of an earliest Carboniferous
- fauna from Burnmouth, Scotland, UK. M. Sci. thesis, University of Cambridge.
- 987 Otoo, B. K. A., Clack, J. A., Smithson, T. R., Bennett, C. E., Kearsey, T. I., Coates, M. I. In press
- 988 2018. An early post-Devonian vertebrate fauna preserved in Scottish Tournaisian flood
- 989 plain deposits. *Palaeontology*.



990 Panchen, A.L. 1972. The skull and skeleton of *Eogyrinus atthevi* Watson (Amphibia: 991 Labyrinthodontia). Philosophical Transactions of the Royal Society of London (B) 263: 992 279-326. 993 Panchen, A.L. 1977. On Anthracosaurus russelli Huxley (Amphibia: Labyrinthodontia) and the 994 family Anthracosauridae. Philosophical Transactions of the Royal Society of London (B) 995 279: 447-512. 996 Panchen, A.L. 1985. On the amphibian Crassigyrinus scoticus Watson from the Carboniferous of 997 Scotland. Philosophical Transactions of the Royal Society of London (B) 309: 461-568. 998 Parker, K., Warren, A., Johanson, Z. 2005 Strepsodus (Rhizodontida, Sarcopterygii) Pectoral 999 Elements From The Lower Carboniferous Ducabrook Formation, Queensland, Australia. 1000 *Journal of Vertebrate Paleontology* 25(1):46–62. 1001 Pierce, J.L., Meyer, G.A. & Jull, A.J.T. 2004. Fire induced erosion and millennial-scale climate 1002 change in northern Ponderosa pine forests. *Nature* 432, 87-90. 1003 Richards, K. R., Sherwin, J., Smithson, T. R., Bennion, R.F., Davies, S.J., Marshall, J.E.A. Clack, 1004 J. A. Diverse and durophaguous: Early Carboniferous chondrichthyans from the Scottish 1005 Borders. Earth and Environmental Transactions of the Royal Society of Edinburgh. https:// 1006 doi.org/10.1017/S1755691018000166 1007 Rimmer, S.M., Hawkins, S.J., Scott, A.C., Cressler III, W.L. 2015. The rise of fire: Fossil charcoal 1008 in Late Devonian Marine shales as an indicator of expanding terrestrial ecosystems, fire 1009 and atmospheric change. American Journal of Science. 315, 713–733, DOI 1010 10.2475/08.2015.01 1011 Roper, H. 1997 Origin of the 'Berwick Monocline': geometrical and geophysical considerations. 1012 Scottish Journal of Geology. 33, 133-148



Earth Sciences 75, 311-340.

1013 Ross, A. J., Edgecombe, G. D., Clark, N. L. D., Bennett, C. E., Carriò, V., Contreras-Izquierdo, 1014 R., Crichton B 2018. A new terrestrial millipede fauna of earliest Carboniferous 1015 (Tournaisian) age from southeastern Scotland helpf sill 'Romer's Gap'. Earth and 1016 Environmental Transactions of the Royal Society of Edinburgh. https:// 1017 doi.org/10.1017/S1755691018000142 1018 Sallan L. C., Gallimberti, A. K. 2015. Body-size reduction in vertebrates following the end-Devonian mass extinction. Science 350, 812-815. 1019 1020 Scott, A.C. 1988. Volcanoes, fires and the Lower Carboniferous vegetation of Scotland. *Journal* 1021 of the Open University Geological Society. 8, 27-31. 1022 Scott, A.C. 2000. The Pre-Quaternary History of Fire. *Palaeogeography*, *Palaeoclimatology*, 1023 Palaeoecology 164, 281-329. 1024 Scott, A.C. 2010. Charcoal recognition, taphonomy and uses in palaeoenvironmental analysis. 1025 Palaeogeography, Palaeoclimatology, Palaeoecology. 291, 11-39. 1026 Scott, A.C. 2018. Burning Planet. The story of fire through time. Oxford University Press, Oxford 224 pp. ISBN 978-0-19-873484-0 1027 1028 Scott, A.C., Bowman, D.J.M.S., Bond, W.J., Pyne, S.J., Alexander M. 2014. Fire on Earth: An 1029 *Introduction*. Oxford: J. Wiley and Sons. 1030 Scott, A.C., Galtier, J. 1996. A review of the problems in the stratigraphical, palaeoecological and 1031 palaeobiogeographical interpretation of Lower Carboniferous (Dinantian) floras from 1032 Western Europe. Review of Palaeobotany and Palynology 90, 141-153. 1033 Scott, A. C., Galtier, J., Clayton, G. 1984. Distribution of anatomically-preserved floras in the 1034 Lower Carboniferous in Western Europe. Transactions of the Royal Society of Edinburgh:

1036 Scott, A.C., Glasspool, I.J. 2006. The diversification of Paleozoic fire systems and fluctuations in 1037 atmospheric oxygen concentration. Proceedings of the National Academy of Sciences, 1038 U.S.A. 103, 10861-10865. 1039 Scott, A. C., Meyer-Berthaud, B. 1985. Plants from the Dinantian of Foulden, Berwickshire, 1040 Scotland. Transactions of the Royal Society of Edinburgh: Earth Sciences 76, 13-20 1041 Scott, A. C., Rex, G. M. 1987. The accumulation and preservation of Dinantian plants from 1042 Scotland and its borders. In: Adams A. E., Miller J. and Wright V. P. eds European 1043 Dinantian Environments. Geological Journal Special Issue 12, 329-344. 1044 Scrutton C, Turner B. 1995. The geology of Eyemouth and Burnmouth. In Northumbrian rocks and 1045 landscape: a field guide. Ed Scrutton C. Yorkshire Geological Society, York: UK 31-41. 1046 Sharp, E. L., Clack, J. A. 2013 A review of the Carboniferous lungfish genus *Ctenodus* Agassiz 1047 1838 from the United Kingdom, with new data from an articulated specimen of *Ctenodus* 1048 interruptus. Earth and Environmental Science Transactions of the Royal Society of 1049 Edinburgh, 104, 169-204 1050 Smithson, T. R., Richards, K. R., Clack, J. A. 2015. Lungfish diversity in Romer's Gap: reaction 1051 to the end-Devonian extinction. *Palaeontology* 59,29-44 doi: 10.1111/pala.12203 1052 Smithson, T.R., Wood, S.P., Marshall, J.E.A., Clack, J.A. 2012. Earliest Carboniferous tetrapod 1053 and arthropod faunas from Scotland populate Romer's Gap. Proceedings of the National 1054 *Academy of Sciences*. 109:4532-4537. 1055 Smithson, T. R., Browne, M. A., Davies, S. J., Marshall, J. E. A., Millward, D., Walsh, S. A., 1056 Clack, J. A. 2017. A new Mississippian tetrapod from Fife, Scotland, and its environmental 1057 context. Papers in Palaeontology. 2017, 1-11. doi: 10.1002/spp2.1086.

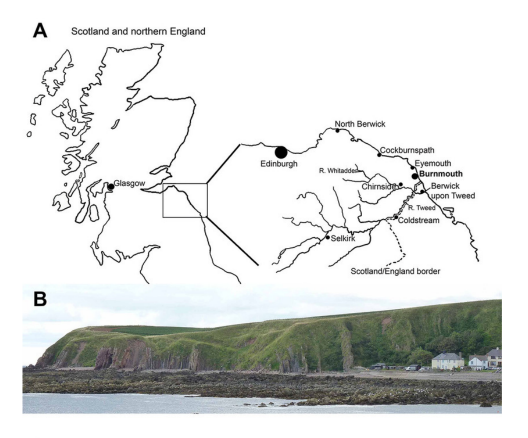


1058	Smithson, T. R., Clack, J. A. in press 2018. A new tetrapod from Romer's Gap reveals an early
1059	adaptation for walking. Earth and Environmental Science Transactions of the Royal
1060	Society of Edinburgh
1061	Watson, D. M. S., Gill, E. L. 1923. The structure of certain Palaeozoic Dipnoi. <i>Journal of the</i>
1062	Linnean Society of London, Zoology 35, 163–216.
1063	Young G. C., Long, J. A., Ritchie, A. 1992 Crossopterygian fishes from the Devonian of
1064	Antarctica: systematics, relationships and biogeographic significance. Records of the
1065	Western Australia Museum, Supplement 14, 1-77.
1066	Yazdi, M., Turner, S., 2000. Late Devonian and Carboniferous vertebrates from the Shishtu and
1067	Sardar formations of the Shotori Range, Iran. Records of the Western Australian Museum
1068	Supplement 58, 223-240.
1069	

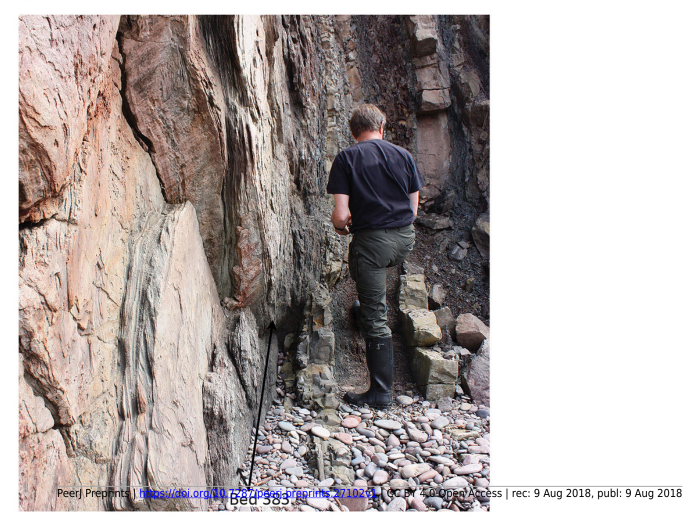


Geographical and geological context of Burnmouth and Bed 383.

A. A map of Scotland with inset showing locality of Burnmouth and major towns and rivers. B. A view of the section exposed at Burnmouth. The horizon described lies close to the most prominent sandstone near the left hand side of the photograph. C. Photograph of the immediate region of Bed 383 prior to the 2018 rockfall, Rob Clack at the rock face (Photograph by SJD).



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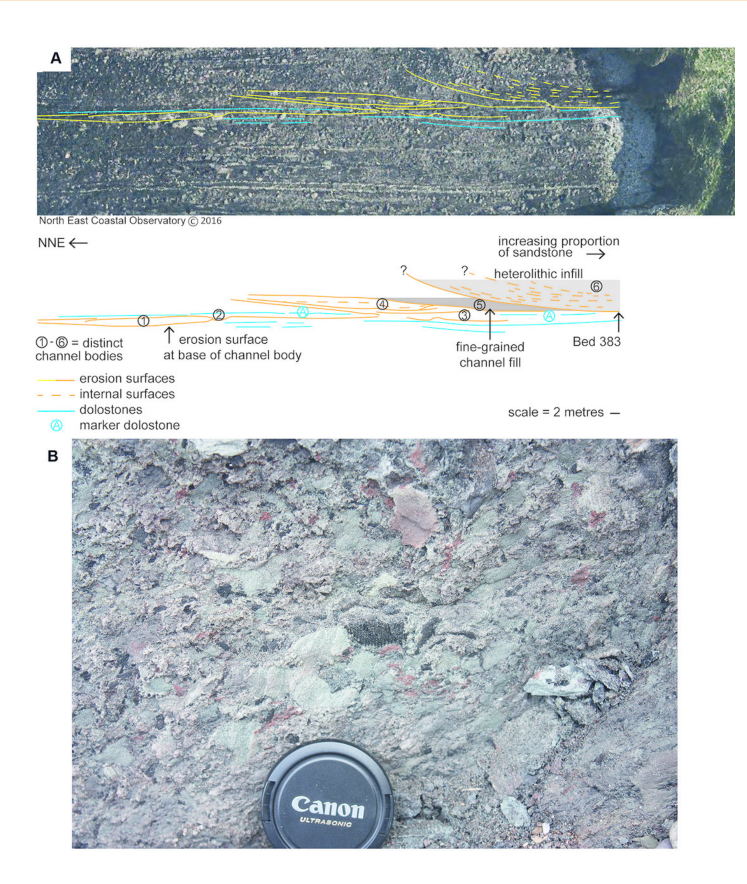




Geological context and channel interpretation.

A. Interpreted architecture of 6 distinct packages interpreted as river channel deposits. Bed 383 is located at the base of package 6 at the cliffward and barely extends onto the wave-cut platform. The erosion surface at the base of package 6 cuts down to a level below packages 4 and 5. The marker dolostone (A) can be traced into the cliff and is not removed by erosion associated with the base of package 6. B. Close-up photograph of a sample of Bed 383 matrix.



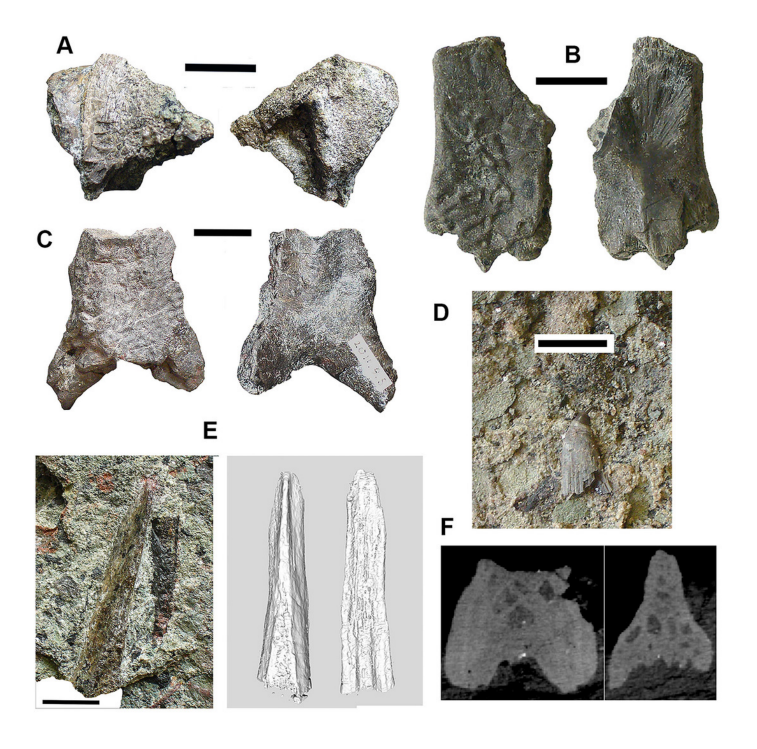




Large tetrapod skull bones.

A. Basipterygoid process 2017.2.12, dorsal view at left, ventral view at right. B. Postfrontal 2017.2.545 dorsal view at left, ventral view at right. C. Tabular 2011.9.5, dorsal view at left, ventral view at right. D. Tooth 2017.2.384. E. Parasphenoid 2017.2.187. Left image, photograph of the specimen, right image, micro-CT renderings, ventral view at left, dorsal view at right. F. Parasphenoid 2017.2.187 micro-CT scan sections near base (left) and tip (right). All scale bars 10 mm.



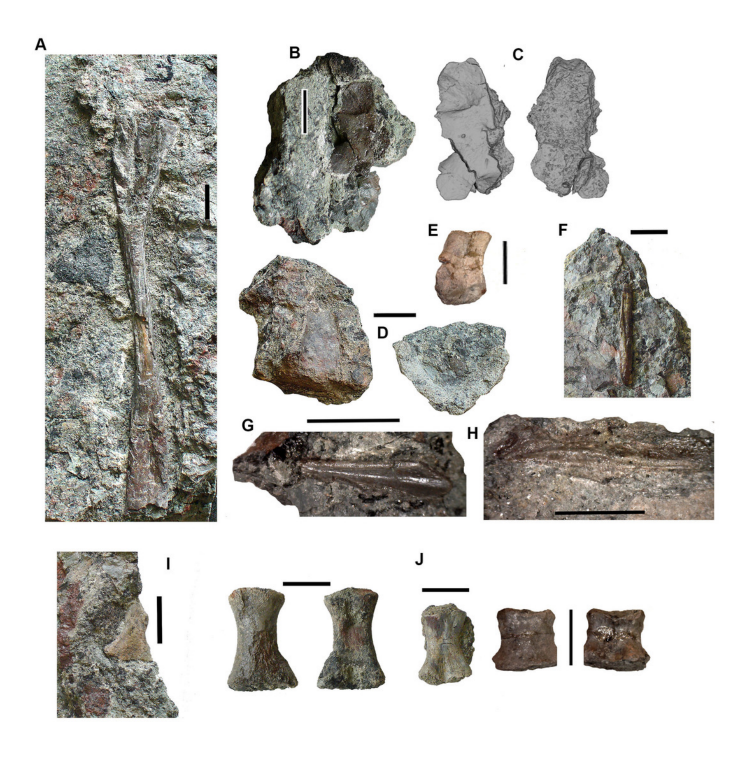




Large tetrapod postcranial bones.

A Rib 2017.2.189a. B. Neural arch 2017.2.189g. C. Micro-CT scans of 2017.2.189g, external view at left, internal view at right. D. Centra 2017.2.374, lateral view one centrum at left, anterior view of a second at right. E. Small neural arch 2017.2.511. F. Body scale 2017.2.107. G. Body scale 2017.2.175. H. Body scale 2017.2.144. I. Phalanx 2017.2.61. J. Metapodials or phalanges 2017.2.388, 2017.2.380 and 2017.2.61. All scale bars 10 mm.



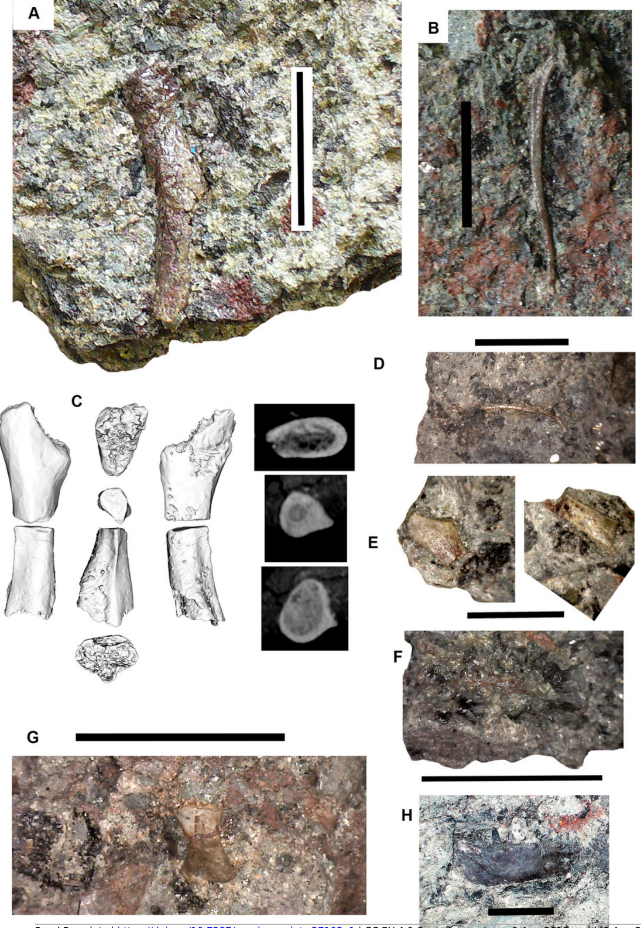




Small tetrapod elements.

A. Rib 2017.2.476. B. Rib 2017.2.170. C.Ulna 2017.2.51 from micro-CT scan. Views of the sections at right with the top image from near the proximal end, the middle image from midshaft and the lower image from near the distal end. D. Rib 2017.2.149. E Ulna 2017.2.51 in part and counterpart, proximal end at left. F. Jaw 2017.2.117. H. Jaw 2017.2.443. All scale bars 10 mm.



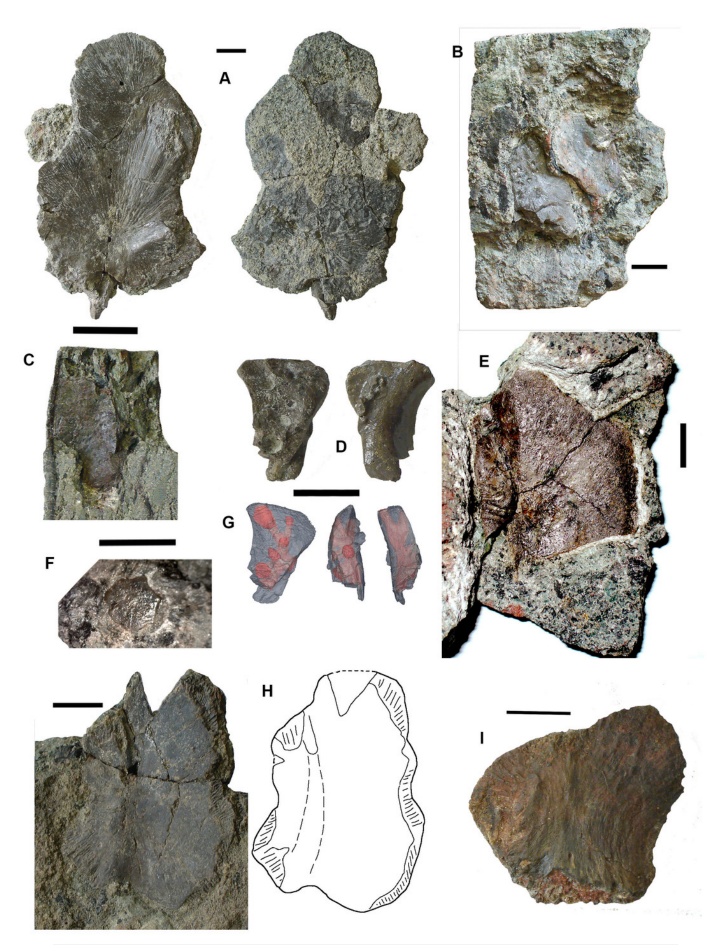


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Lungfish skull bones.

A. Conjoined J- and I-bones. Ventral view at left, dorsal view at right. B. Skull bone 2011.9.10, lying partially covered by a rhizodont scale. C. Skull bone 2017.2189i. D. Circumorbital bone 2011.9.7, dorsal view at left, ventral view at right. E. Skull bone 2011.9.10. F. Skull bone 2017.2.193. G. Micro-CT renderings showing lateral line canals. H. Skull bone 2017.2.443. I. Skull bone 2018.1.2. All scale bars 10 mm.



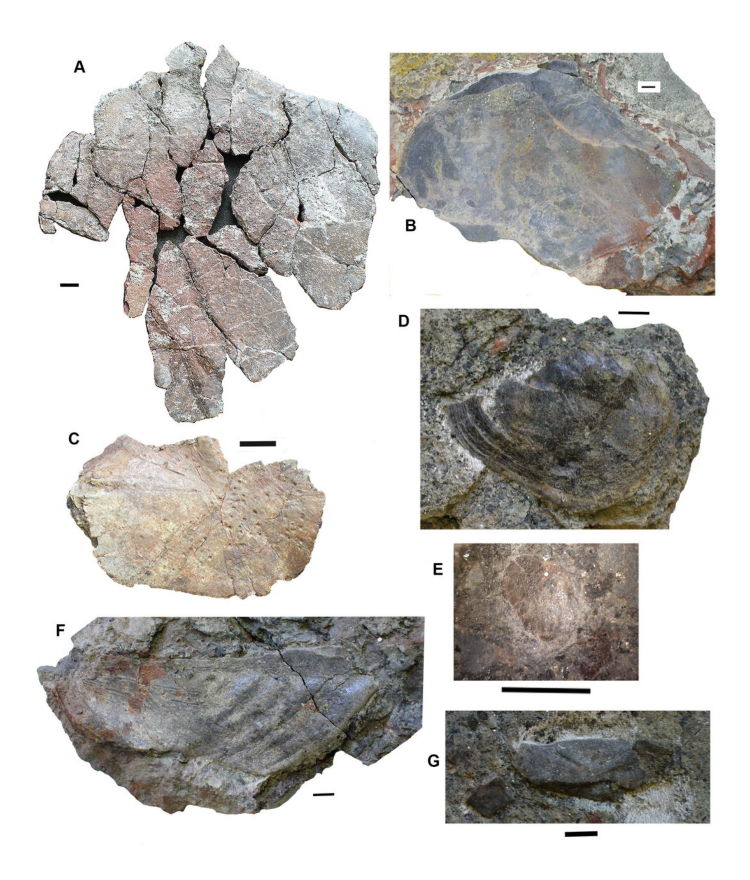
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Lungfish opercular series.

A. Operculum 2017.2.188. B. Operculum 2017.2.667. C. Operculum 2011.9.6. D. Operculum 2018.1.5. E. Operculum 2017.2.720. F. Suboperculum 2018.1.7. G. Possible suboperculum or submandibular 2018.1.5. All scale bars 10 mm.

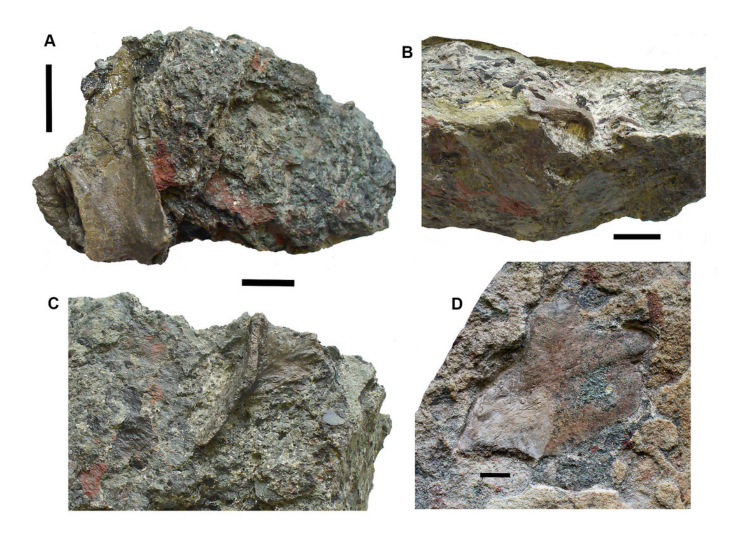






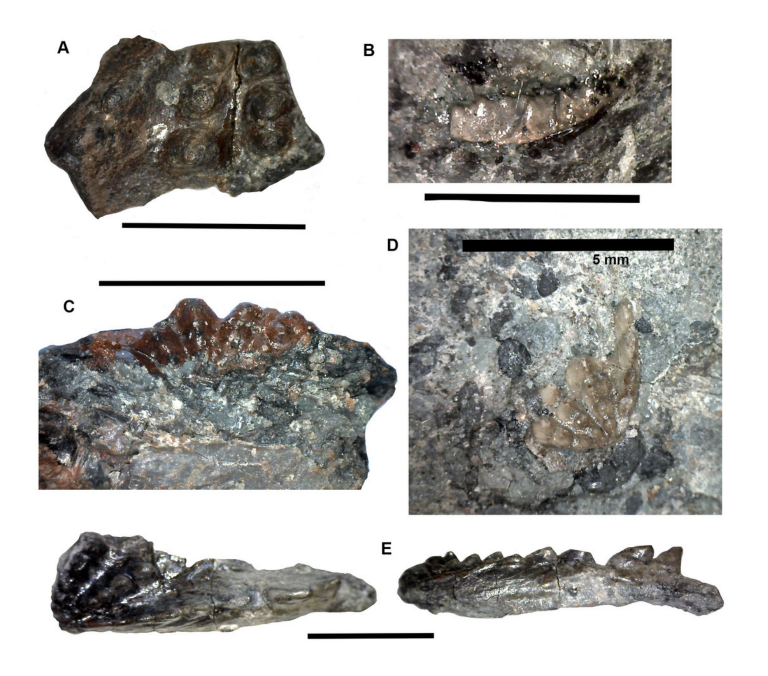
Lungfish shoulder girdle elements

A. Partial clavicle 2017.2.70. B.Partial cleithrum 2017.2.494a. C. Possible central element 2017.2.398. D. Anocleithrum 2017.2.57.



Lungfish tooth plates.

A. 2017.2.54. B. c.f *Ballagadus caustrimi* 2017.2.66. C. *B. caustrimi* 2017.2.67. D. *B. caustrimi* 2017.2.513. E. *Ballagadus rossi* 2014.1.2, buccal view at left, lingual view at right. All scale bars except D 10 mm, D, 5mm.

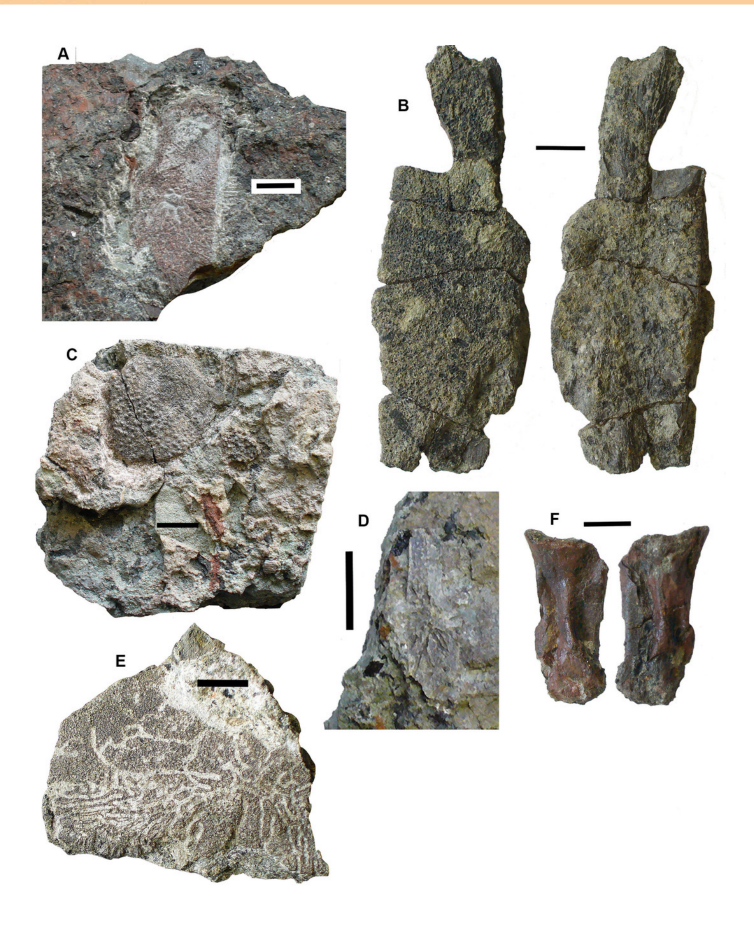




Rhizodont skull bones and hyomandibula.

A. Postparietal 2017.2.186. B. Postparietal 2017.2.406. External view at right, internal view at left. C. Extrascapula 2017.2..94. D. Postparietal 2017.2.643. E. Possible partial postparietal 2017.2.52. F.Hyomandibula 2017.2.338. All scale bars 10 mm.

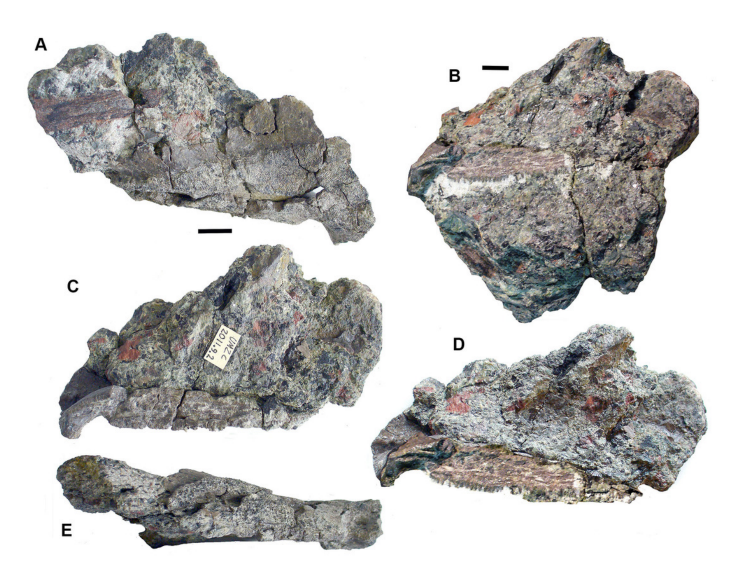






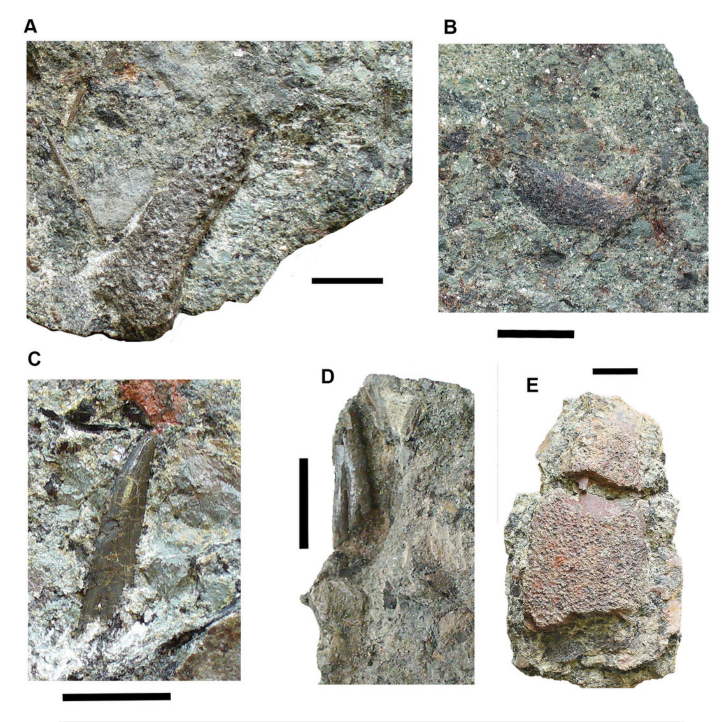
Rhizodont pterygoid 2011.9.2.

A. View of surface showing denticulated area, vermiform ornament and transition to plain surface, articular end at right. B. View of overlap surface with teeth along the edge before preparation. C. Same as B but after preparation, articular end at left. D. Restoration of B plus C. E. View of assumed ventral surface, articular end at left. All scale bars 10 mm.



Rhizodont jaw elements and teeth.

A. Infradentary 2017.2.477. B. Possible small dentary 2017.2.64. C. Tooth 2017.2.72b. D Tooth 2017.2.99. E. Gular plate 2017.2.64.

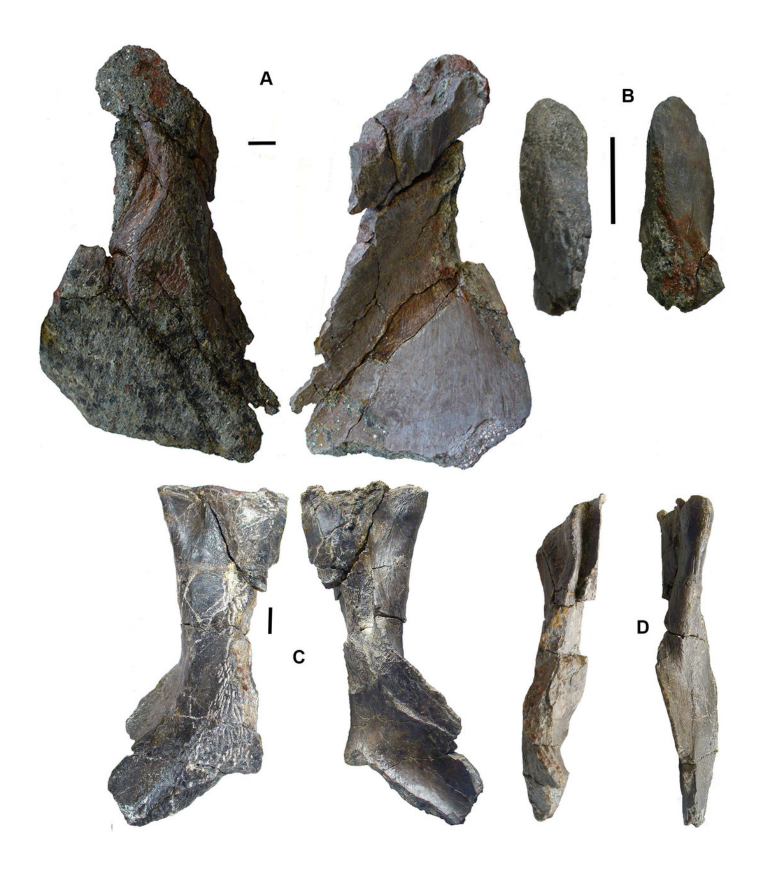




Rhizodont cleithra.

A. A very large example 2017.2.716, external view at left, internal view at right. B. The smallest example 2017.2.2018.1.4. C. 2011.9.3. External view at left, internal view at right. C. An eroded example 2017.2.399. All scale bars 10 mm.



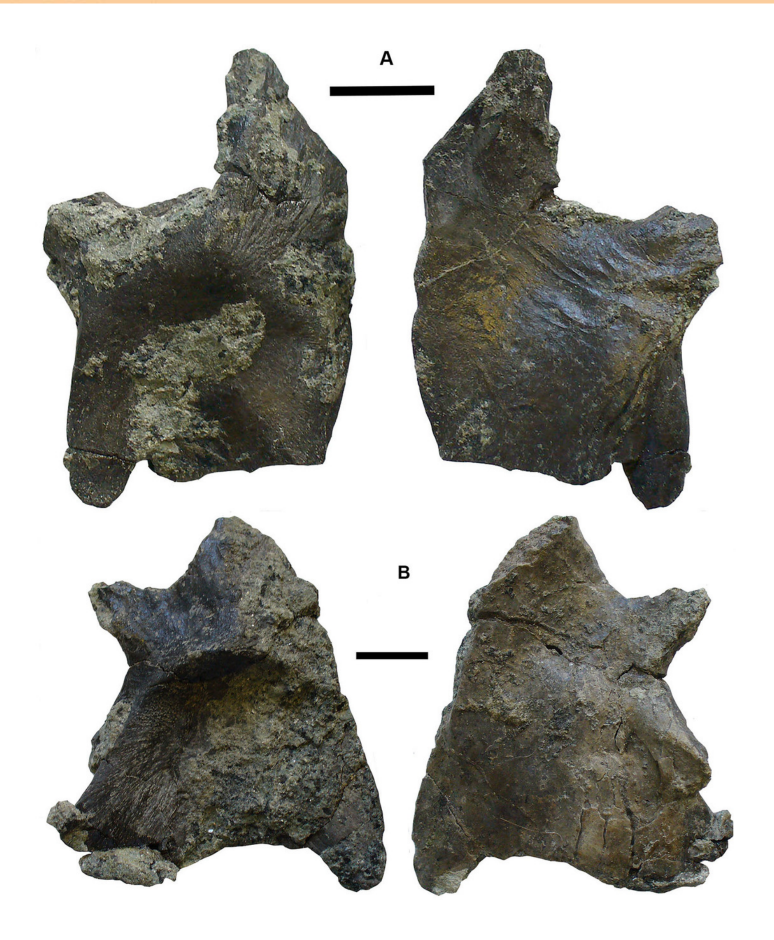




Rhizodont anocleithra.

A. 2017.2.386, probable internal view at left, external view at right B. 2017.2.385, probably internal view at left, external view at right. All scale bars 10 mm.



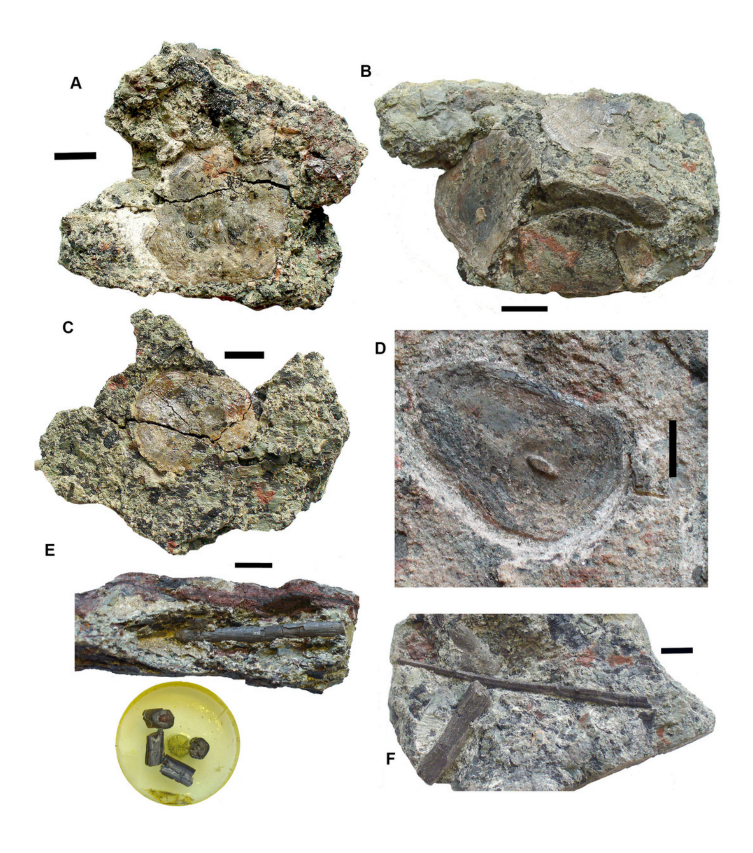




Rhizodont scales and fin spine.

A. Scale 2011.9.4. B. Scale 2017.2.95. C. Scale 2011.9.4. D. Scale 2011.2.11. E. Part of fin spine 2017.2.568, with mounted sections below. F. Fin spine 2017.2.494b, with a partial rib of uncertain identity adjacent. All scale bars 10 mm.



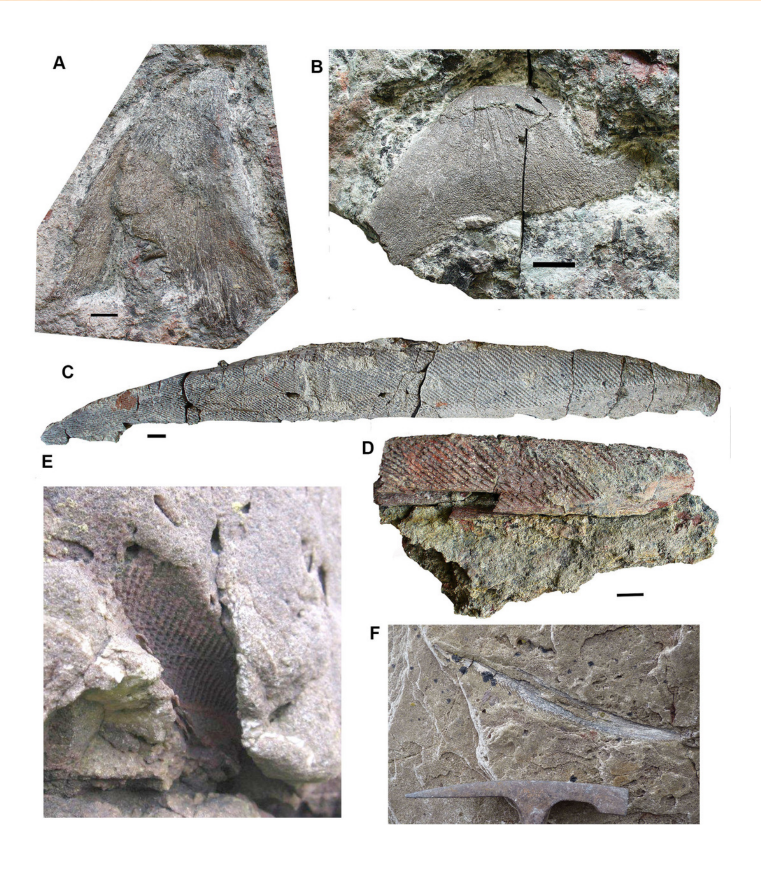




Gyracanth elements.

A. Scapulocoracoid spine 2011.9.11. B. Procoracoid spine 2017.2.189b. C. Pelvic spine 2017.2.182. D. Spine 2017.2.185. E and F. Spines preserved in situ, mostly in natural mould as seen in 2011. All scale bars 10 mm. Spine in E would be similar in size (when complete) to that in part C. Hammer head for scale in F.

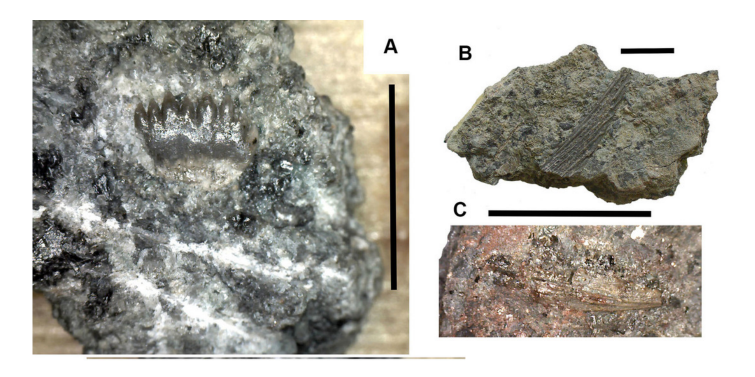






Chondrichthyan elements.

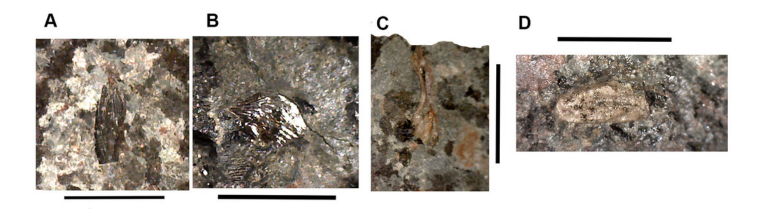
A. *Ageleodus* tooth 2017.2.282. B. *Onychoselache*-like spine 2017.2.282. C. Spine 2017.2.51. Scale bars for B and C, 10 mm, for A, 5 mm.





Actinopterygian elements and coprolite.

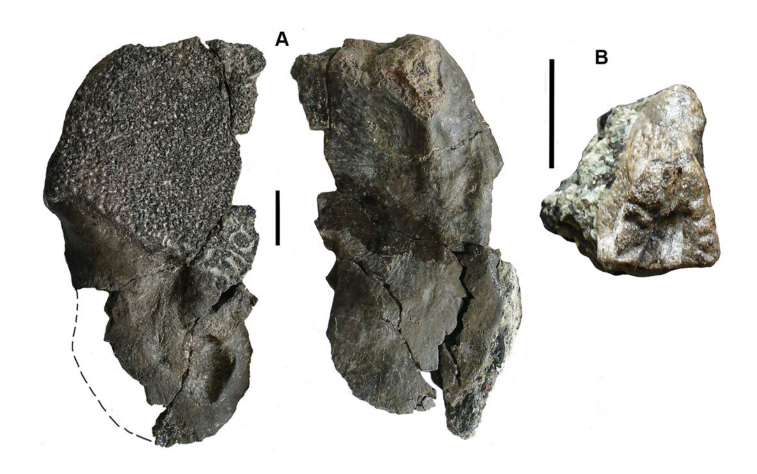
A. Scale 2017.2.200. B. Scale 2018.1.3. C. Possible skull or girdle bone 2018.1.1. D. Coprolite 2017.2.58. Scale bars in A and B, 5 mm, C and D, 10 mm.





Elements of uncertain identity.

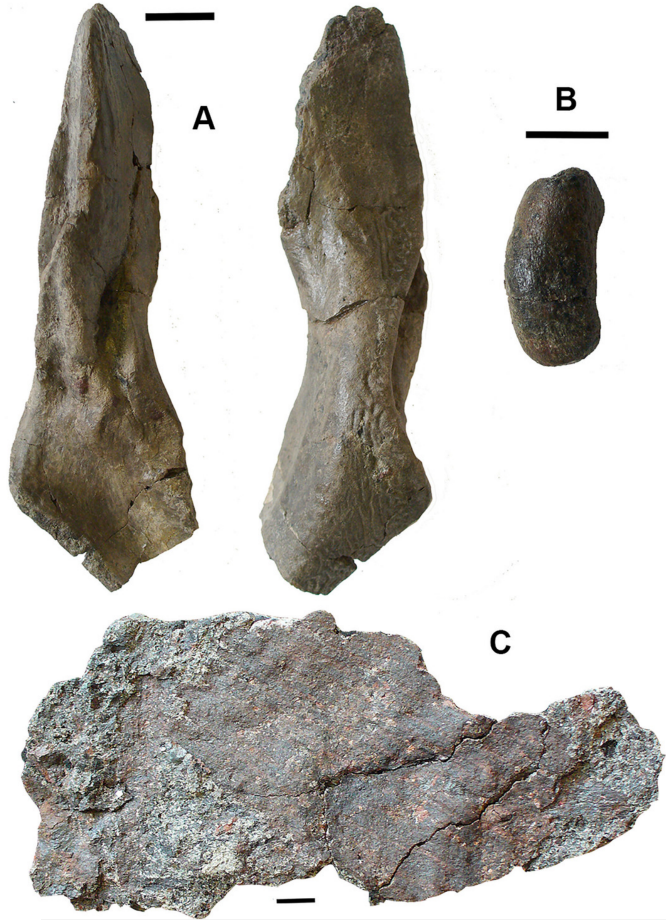
A. Large bone, possibly rhizodont 2017.2.327. B. Unknown element 2017.2.49. All scale bars 10 mm.





Abraded elements and plant remains.

A. Rhizodont cleithrum 2017.2.400. B. Rolled bone 2017.2.214. C. Plant stem, probably lycopod 2017.2.293. All scale bars 10 mm.



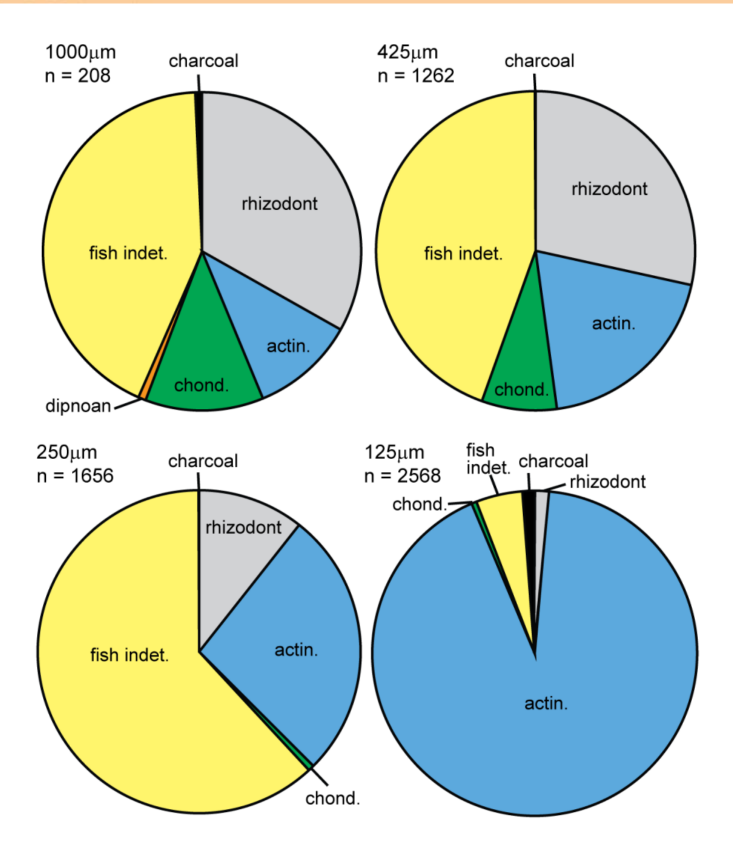
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Microfossil assemblage.

Percentage plots of microfossil counts for the four size fractions analysed. The full data table of counts for all size fractions is presented in Supplementary Information 4. Abbreviations: actin. - actinopterygian; chond. - chondrichthyan; indet. - indeterminate.







Selected microfossil specimens.

A. rhizodont scale with tuberculate internal structure and red staining. B. rhizodont scale with ridge and groove internal structure. C. rhizodont dermal bone, view of the exterior surface with irregular tuberculate ornamentation. D. actinopterygian scale fragment, exterior surface with transverse ridges and grooves. E. actinopterygian lepidotrichia fin bone fragment. F. actinopterygian dermal bone, exterior surface with irregular tuberculate ornament and gamine coating. G. rhizodont tooth with striations at the base, lateral view. H. actinopterygian tooth, with transparent cap, lateral view. I. pharyngeal actinopterygian tooth, double row, lateral view. J. *Ageleodus* tooth with broken tooth cusps, lateral view. K. Elasmobranch scale with a concave base and spinose top, lateral view. L. Hybodont scale with spines that are joined together into a star shape, lateral view. M. ?Ctenacanth scale with a top of curved spines. Scale bars 250 µm.



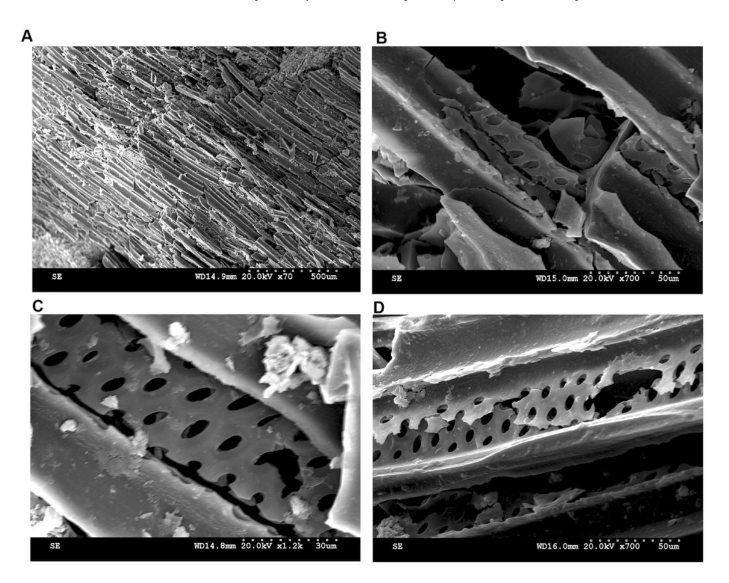




Scanning Electron Micrographs of charcoal fragment from Bed 383.

A. Longitudinal section showing pitting in the cell walls. B. Detail of a showing rows of circular pits. C. Detail of a showing three rows of circular pits. D. Detail of a showing three rows of circular pits.

*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.

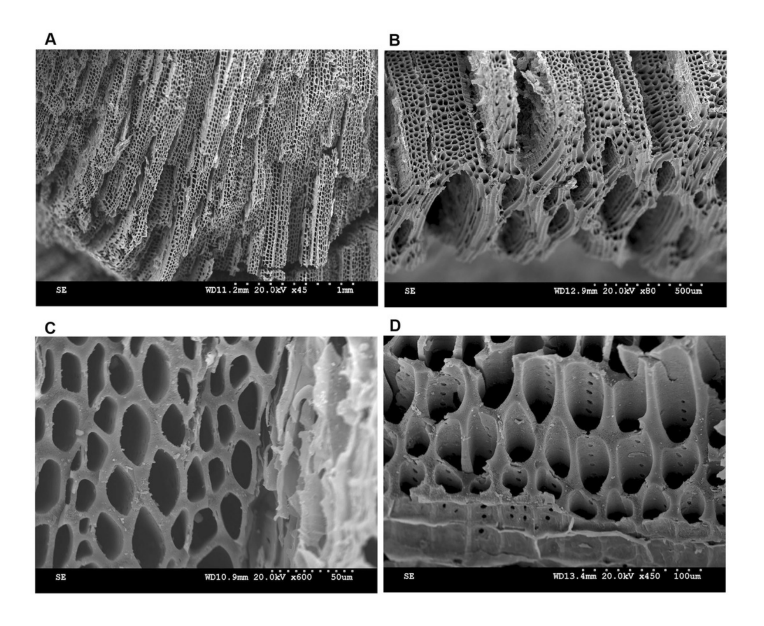




Scanning Electron Micrographs of large charcoal fragment from Bed 362.

A. Low power image showing rows of tracheids separated by ray tissue. B. Detail of a showing the position of short rays. C. Detail of tracheids showing cell wall homogenization. D. Detail of tracheids showing single rows of circular pits and ray cells with pits.

*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.

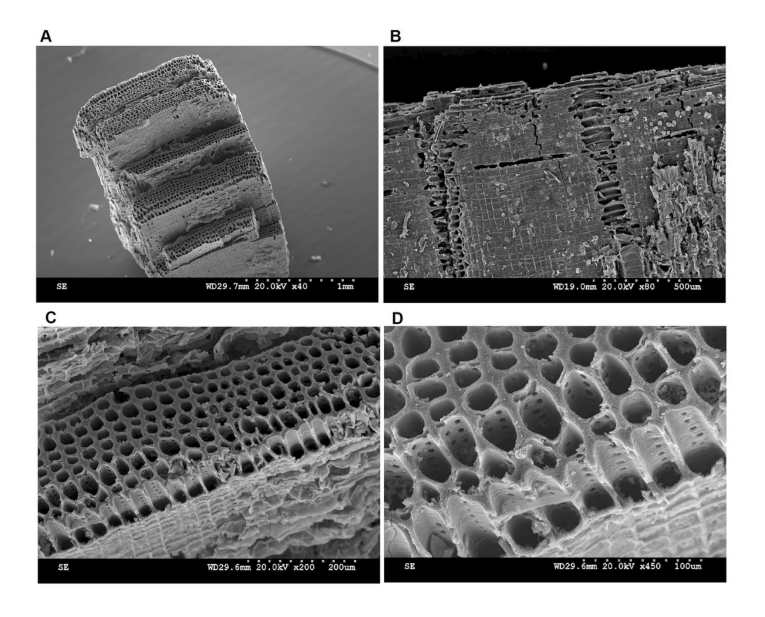




Scanning Electron Micrographs of charcoal fragment 1 from Bed 362

A. Wood fragment showing the position of ray tissue. B. Detail of a showing ray tissue. C. Detail of a showing pitting in tracheids. D. Detail of a showing rows of circular pits, homogenized cell walls and ray tissue.

*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.





Scanning Electron Micrographs of charcoal fragment 2 from Bed 362

A. Specimen showing rows of tracheids separated by ray cells. B. Detail of tracheids showing cell wall homogenization. C. Specimen showing both tracheids and ray tissue. D. Detail of A showing tracheids. E. Circular bordered pit in tracheid.



