

## Digital preparation and osteology of the skull of *Lesothosaurus diagnosticus* (Ornithischia: Dinosauria)

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Several skulls of the ornithischian dinosaur *Lesothosaurus diagnosticus* (Lower Jurassic, southern Africa) are known, but all are either incomplete, deformed, or incompletely prepared. This has hampered attempts to provide a comprehensive description of skull osteology in this crucial early dinosaurian taxon. Using visualization software, computed tomographic scans of the *Lesothosaurus* syntypes were digitally segmented to remove matrix and identify and separate individual cranial and mandibular bones, revealing new anatomical details such as sutural morphology and the presence of several previously undescribed elements. Together with visual inspection of exposed skull bones, these CT data enable a complete description of skull anatomy in this taxon. Comparisons with our new data suggest that two specimens previously identified as *Lesothosaurus* sp. (MNHN LES 17 and MNHN LES 18) probably represent additional individuals of *Lesothosaurus diagnosticus*.

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2 Dinosauria)

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## 20 **Abstract**

21 Several skulls of the ornithischian dinosaur *Lesothosaurus diagnosticus* (Lower Jurassic,  
22 southern Africa) are known, but all are either incomplete, deformed, or incompletely prepared.  
23 This has hampered attempts to provide a comprehensive description of skull osteology in this  
24 crucial early dinosaurian taxon. Using visualization software, computed tomographic scans of  
25 the *Lesothosaurus* syntypes were digitally segmented to remove matrix and identify and separate  
26 individual cranial and mandibular bones, revealing new anatomical details such as sutural  
27 morphology and the presence of several previously undescribed elements. Together with visual  
28 inspection of exposed skull bones, these CT data enable a complete description of skull anatomy  
29 in this taxon. Comparisons with our new data suggest that two specimens previously identified as  
30 *Lesothosaurus* sp. (MNHN LES 17 and MNHN LES 18) probably represent additional  
31 individuals of *Lesothosaurus diagnosticus*.

32

## 33 **Introduction**

34 Ornithischian dinosaurs underwent major taxonomic and ecological radiations during the  
35 Jurassic (Serenó, 1997; Butler, Upchurch & Norman, 2008a) resulting in diverse craniodental  
36 morphologies and, presumably, disparate feeding strategies (e.g., Weishampel & Norman, 1989;  
37 Norman & Weishampel, 1991; Sereno, 1997; Barrett, 2014; Mallon & Anderson, 2014).  
38 Understanding the evolution of this trophic diversity requires detailed knowledge of skull  
39 anatomy (including the potential for cranial kinesis), cranial myology, jaw mechanism, and diet  
40 at the base of Ornithischia.

41 Triassic ornithischians are exceptionally rare, with only two recognized occurrences –  
42 *Pisanosaurus mertii* from Argentina and *Eocursor parvus* from South Africa, both of which

43 include isolated lower jaws but lack substantial cranial material (Casamiquela, 1967; Bonaparte,  
44 1976; Butler, Smith & Norman, 2007; Butler, 2010). A third potentially Triassic-aged taxon, an  
45 indeterminate heterodontosaurid from Argentina (Báez & Marsicano, 2001), may be Early  
46 Jurassic in age (Olsen, Kent & Whiteside, 2011). By contrast, the skull of the Early Jurassic  
47 heterodontosaurid *Heterodontosaurus tucki* from South Africa is well known, and has been  
48 described in detail on the basis of two almost complete skulls (Crompton & Charig, 1962; Santa  
49 Luca, Crompton & Charig, 1976; Norman et al., 2011; Sereno, 2012) and several incomplete  
50 skulls (Norman et al., 2011; Porro et al., 2011; Sereno, 2012), including that of a juvenile  
51 specimen (Butler, Porro & Norman, 2008b). Cranial and lower jaw material is preserved for at  
52 least four additional heterodontosaurid taxa from the Early Jurassic of South Africa (Haughton,  
53 1924; Thulborn, 1974; Hopson, 1975; Porro et al., 2011; Sereno, 2012) and one undescribed  
54 specimen from the Early Jurassic of western North America (Attridge, Crompton & Jenkins,  
55 1985; Sereno, 1986). Despite their early occurrence and basal position within ornithischian  
56 phylogeny (Butler et al., 2008a), most Early Jurassic heterodontosaurids exhibit cranial and  
57 dental specializations atypical of primitive ornithischians, including: a strongly heterodont  
58 dentition; closely-packed, chisel-shaped maxillary and dentary ‘cheek’ teeth (some species lack a  
59 distinct constriction between the crown and root or a cingulum); heavy tooth wear; a strongly  
60 developed coronoid process of the lower jaw; a strongly depressed jaw joint; and extensive  
61 fusion of cranial sutures (e.g. Norman et al., 2011; Sereno, 2012). Thus, the skull of  
62 *Heterodontosaurus* is probably not representative of skull morphology in the earliest  
63 ornithischians.

64 Skull material is also known for various Early Jurassic thyreophoran taxa, including  
65 *Emausaurus* (Haubold, 1990), *Scelidosaurus* (Owen, 1863; Barrett, 2001) and *Scutellosaurus*

66 (Colbert, 1981; Rosenbaum & Padian, 2000). However, cranial material of *Scutellosaurus* is  
67 disarticulated and fragmentary, and the skulls of *Emausaurus* and *Scelidosaurus* display a  
68 number of specialisations (such as the presence of curved tooth rows and unusual occlusal  
69 relationships) that are unlikely to have been present in more primitive ornithischians.  
70 Disarticulated cranial material of *Laquintasaura* from the Early Jurassic of Venezuela is also  
71 available (Barrett et al., 2014), but none of the currently available cranial material is amenable to  
72 functional analyses.

73 By contrast, the Early Jurassic taxon *Lesothosaurus diagnosticus* Galton, 1978, known on the  
74 basis of multiple specimens collected from the Upper Elliot and Clarens formations of South  
75 Africa and Lesotho, can serve as a useful model of the early ornithischian condition (Thulborn,  
76 1970; Sereno, 1991; Knoll, 2002a, 2002b). It has been incorporated into numerous studies of  
77 ornithischian phylogeny (e.g., Norman, 1984a; Sereno, 1984, 1986, 1999; Cooper, 1985;  
78 Maryanska & Osmolska, 1985; Butler et al., 2007, 2008a), locomotion (Maidment & Barrett,  
79 2011; Bates et al., 2012; Maidment et al., 2014) and feeding (e.g., Thulborn, 1971; Weishampel,  
80 1984; Galton 1986; Barrett, 1998; Norman, Witmer & Weishampel, 2004; Knoll, 2008).  
81 *Lesothosaurus* possesses a more generalised skull and tooth morphology than that exhibited by  
82 heterodontosaurids or thyreophorans, including: low, triangular ('leaf-shaped') teeth with a  
83 distinct neck and cingulum; coarse denticles on the mesial and distal tooth margins; sporadically  
84 developed high-angled marginal tooth wear and evidence of rapid tooth replacement; an  
85 inturned, 'spout-like' mandibular symphysis; straight tooth rows; and a jaw joint positioned only  
86 slightly below the occlusal plane of the tooth row (Thulborn, 1970; Crompton & Attridge, 1986;  
87 Norman & Weishampel, 1991; Sereno, 1991; Norman et al., 2004). Nevertheless, the lack of a  
88 complete, undistorted skull for *Lesothosaurus* has limited attempts to reconstruct the morphology

89 and arrangement of the jaw adductor musculature (Thulborn, 1971; Holliday, 2009) or carry out  
90 biomechanical analyses of the skull, which could serve as a baseline for comparisons with more  
91 derived ornithischian taxa (e.g., Bell, Snively & Shychoski, 2009).

92 Two skulls (NHMUK PV RU B17 and NHMUK PV RU B23) from the Lower Jurassic Elliot  
93 Formation of Lesotho were described by Thulborn (1970) and referred to *Fabrosaurus australis*.  
94 Galton (1978) later designated both specimens as syntypes of a new taxon, *Lesothosaurus*  
95 *diagnosticus*. These specimens, and other referred material (e.g., NHMUK PV R8501, NHMUK  
96 PV R11004, NHMUK R11956), served as the basis for several anatomical descriptions of the *L.*  
97 *diagnosticus* skull (Thulborn, 1970; Galton, 1978; Norman, 1984b; Weishampel, 1984;  
98 Weishampel & Witmer, 1990; Sereno, 1991; Norman et al., 2004). Two additional partial skulls  
99 from the Early Jurassic of Lesotho (MNHN LES 17, MNHN LES 18) described by Knoll  
100 (2002a, 2002b) share numerous characters with *L. diagnosticus*; however, various anatomical  
101 and proportional differences (as well as the larger size of MNHN LES 18) led to these specimens  
102 being excluded from *L. diagnosticus* and assigned to *Lesothosaurus* sp. Furthermore, the  
103 possibility has been raised that MNHN LES 18 may belong to a larger, sympatric neornithischian  
104 *Stormbergia dangershoeki* (Butler, 2005). The skulls of NHMUK PV RUB 23 (Fig. 1A, B),  
105 MNHN LES 17 and MNHN LES 18 are distorted and missing the anterior snout. NHMUK PV  
106 RUB 17 preserves the remains of at least two individuals in three separate blocks: the fully  
107 prepared 'snout' block contains the anterior ends of the premaxillae and dentaries, and the  
108 prementary (Fig. 1E); the 'palatal' block contains the bones of the palate, ventral facial region,  
109 posterior lower jaws, and ventral braincase (Fig. 1C, D). The 'snout' and 'palatal' blocks pertain  
110 to a single individual, while the partially prepared 'braincase' block contains the remains of a  
111 second individual that also includes several disarticulated skull and postcranial elements (Fig.

112 1F). Other isolated skull elements (e.g., a maxilla, jugal, squamosal) are also registered as part of  
113 NHMUK PV RU B17 and are identical in morphology to those elements preserved in the three  
114 blocks. Other specimens figured previously include NHMUK PV R8501, a nearly complete but  
115 badly crushed and disarticulated skull, including lower jaws, and NHMUK PV R11956, an  
116 articulated but crushed anterior skull with lower jaws, missing its posterior part (Serenó, 1991).  
117 One further undescribed specimen, a juvenile skull (NHMUK PV RU C109), provides only  
118 limited anatomical information. Unfortunately, no single specimen of *Lesothosaurus* preserves a  
119 skull that is complete, articulated and undistorted.

120 Computed tomography (CT) is increasingly applied to dinosaur skulls for various purposes:  
121 detailed anatomical description (Lautenschlager et al., 2014); digital preparation of fragile  
122 material (e.g., Butler et al., 2010, 2012; Porro et al., 2011); reconstruction of disarticulated skulls  
123 (Domínguez Alonso et al., 2004; Sampson & Witmer, 2007); imaging of internal cavities such as  
124 the endocranial cavity (e.g., Evans, Ridgely & Witmer, 2009; Zelenitsky et al., 2011; Walsh &  
125 Knoll, 2011; Knoll et al., 2012, 2013), semicircular canals (Serenó et al., 2007; Witmer et al.,  
126 2008; Walsh et al., 2009) and intracranial sinuses and nasal airways (Sampson & Witmer, 2007;  
127 Witmer & Ridgely, 2008; Miyashita et al., 2011; Bourke et al., 2014); and capturing skull  
128 morphology for biomechanical analyses (Rayfield et al., 2001; Bell et al., 2009; Lautenschlager  
129 et al., 2013; Cuff & Rayfield, 2013; Snively et al., 2013; Button, Rayfield & Barrett, 2014).  
130 Nearly all of these studies have been based on complete, minimally damaged specimens. This  
131 study uses CT scanning and visualization of the *Lesothosaurus* syntype skulls to: 1) to digitally  
132 prepare and provide an osteological description of the *Lesothosaurus* syntype skulls in  
133 combination with information from other referred specimens, supplementing and amending  
134 previous descriptions (Thulborn, 1971; Galton, 1978; Norman, 1984b; Weishampel, 1984;

135 Weishampel & Witmer, 1990; Sereno, 1991; Norman et al., 2004); and 2) to compare the  
136 syntype skulls with material referred to *Lesothosaurus* sp. (Knoll, 2002a, 2002b). The anatomical  
137 description is supplemented by 3D PDFs (Supporting Information, Figs. S1-S4); as previously  
138 noted by Lautenschlager et al. (2014), such documentation permits easier access and inspection  
139 of fossil material.

#### 140 **Institutional abbreviations**

141 MNHN, Muséum National d'Histoire Naturelle, Paris, France; NHMUK, The Natural History  
142 Museum, London, UK; SAM, Iziko South African Museum, Cape Town, South Africa.

143

#### 144 **Methods**

145 The two syntype skulls of *Lesothosaurus diagnosticus* (NHMUK PV RU B17 and NHMUK PV  
146 RU B23) were CT-scanned for this study. NHMUK PV RU B23 was CT-scanned at the Center  
147 for Quantitative X-ray Imaging at Pennsylvania State University (University Park, Pennsylvania,  
148 USA) using an X-Tek micro-focus subsystem of a Varian/BIR Omni-X HD-600 industrial high-  
149 resolution CT system at 180 kV and 600  $\mu$ A. The resulting reconstructions produced 1107 axial  
150 slices with a resolution of 0.15 mm/pixel and a slice thickness of 0.087 mm. Three blocks  
151 (described previously) of NHMUK PV RU B17 were scanned at the Ohio University MicroCT  
152 Facility (OU $\mu$ CT, Athens, Ohio, USA) using a General Electric (GE) eXplore Locus CT scanner  
153 at 80 kV and 500  $\mu$ A with 3600 views and frame-averaging of 8. Reconstruction of the 'snout'  
154 block produced 583 axial slices at an isotropic voxel size of 0.046 mm; reconstruction of the  
155 'palatal' block produced 828 axial slices at an isotropic voxel size of 0.092 mm; reconstruction  
156 of the 'braincase' block produced 634 axial slices at an isotropic voxel size of 0.092 mm.

157 Additionally, CT scans of MNHN LES 17 were examined to facilitate comparisons between this  
158 specimen and the syntype skulls; however, information from this specimen was not used in the  
159 anatomical description. This specimen was scanned at OU $\mu$ CT using the same GE scanner using  
160 the same parameters noted above for NHMUK PV RU B17; reconstruction produced 963  
161 transverse slices at a voxel size of 0.092 mm.

162 CT scans were processed using the 3D visualization software packages Amira 5.3.3 and  
163 Avizo 7.1.1 (FEI Visualization Sciences Group, Mérignac Cedex, France). Within the  
164 Amira/Avizo segmentation editor, density thresholding was used to separate higher density bone  
165 from lower density matrix. Scans were then processed slice-by-slice (interpolating across no  
166 more than five slices at a time) to separate bones from each other at sutures, which were  
167 identified as lower density areas between bones. (In some cases, minerals precipitated within  
168 sutures resulting in boundaries with higher density than surrounding bones.) Original specimens  
169 were used to confirm the location of sutures and to differentiate sutures from post-mortem  
170 damage. Individual bones were isolated and separately labeled within the segmentation editor.  
171 Three-dimensional surface models (.surf files) of each element were created that could be  
172 manipulated in isolation in 3D space; the following anatomical description is based on these  
173 surface models (Fig. 2).

174 Some portions of the CT scans could not be segmented, partly because the X-ray attenuation  
175 properties of the fossil bone and rock matrix were similar enough that contrast was relatively  
176 poor. Although the individual bones of the left lower jaw of NHMUK PV RU B23 were  
177 successfully isolated, scan resolution of the right lower jaw was too poor to separate individual  
178 elements. Scans did not penetrate the interior of NHMUK PV RU B23; it is likely that a  
179 complete braincase and palate are present but they cannot be identified. Teeth could not be

180 discerned in CT scans of NHMUK PV RU B23. Maxillary and dentary teeth were identified and  
181 segmented in the ‘palatal’ block of NHMUK PV RU B17 but, due to the presence of a high-  
182 density precipitate in their pulp cavities, resolution of tooth shape is poor.

183

## 184 **Results**

### 185 **Facial skeleton and skull roof**

186 General comments on overall skull morphology are based primarily on NHMUK PV RU B23  
187 (Fig. 2A, C), the most complete and least distorted skull in our sample, supplemented with  
188 information from the other available specimens. In lateral view, the cranium of NHMUK PV RU  
189 B23 is tallest just behind the orbit; the skull roof (frontals, parietal) is gently rounded in lateral  
190 profile and the snout tapers smoothly to the premaxillae; there is no break in slope along the  
191 snout anterior to the orbits as occurs in *Heterodontosaurus* (SAM-PK-K1332; Norman et al.,  
192 2011; Sereno, 2012). The orbits are circular in outline and are large relative to skull size,  
193 representing approximately 36% of basal skull length (i.e., as measured from the anterior margin  
194 of the premaxilla to the posterior margin of the basioccipital) . The antorbital fossa is sub-  
195 triangular in outline, with its apex pointing dorsally, and is relatively small, with a maximum  
196 (ventral) length that is approximately 13% of basal skull length. In dorsal view (Fig. 2C), the  
197 supratemporal fenestrae are anteroposteriorly longer than mediolaterally wide and have a sub-  
198 ovate to sub-triangular outline, whereas the infratemporal fenestrae are sub-rectangular in lateral  
199 view and extend for most of the height of the skull. The shape of the external narial opening can  
200 be estimated from the partially complete premaxillae of NHMUK PV RU B17 and NHMUK PV

201 R11956 which indicate that the bony narial openings were likely to have been small and sub-  
202 ovate in outline.

203 The craniomandibular joint is depressed relative to the maxillary alveolar margin. In dorsal  
204 view, the cranium is widest across the postorbitals, tapering anteriorly to the premaxillae;  
205 anterior to this, the shape of the nasals and medial curvature of the maxillae result in a short,  
206 strongly pointed muzzle (Fig. 2C). The postorbital portion of the skull has a box-like profile in  
207 dorsal view. In occipital view, the skull is widest across the midshafts of the quadrates.

### 208 **Premaxilla**

209 The premaxilla of *Lesothosaurus* is composed of a main body with narial, maxillary, posterior  
210 and palatal processes (premaxillary shelf). Both the left and right premaxillae are preserved in  
211 the ‘snout’ block of NHMUK PV RU B17 (Fig. 3), although both sides lack the maxillary  
212 process. Each premaxilla bears six alveoli in addition to several unerupted replacement teeth  
213 (two in the right, one in the left) revealed by CT-scanning (Fig. 3F). There is a short edentulous  
214 area anterior to the first premaxillary tooth, which is rugose and probably supported a keratinous  
215 rhamphotheca (Weishampel & Witmer, 1990; Sereno, 1991; Knoll, 2008; *contra* Thulborn,  
216 1970). The premaxilla forms the ventral margin of the external naris, and a weak excavation is  
217 present on the main body ventral to the narial opening, though this forms a smooth slope rather  
218 than a distinct fossa (NHMUK PV RU B17; NHMUK PV R8501; NHMUK PV R11956). The  
219 anterior premaxillary foramen (Fig. 3A, apmxf) lies at the anteroventral tip of the premaxilla,  
220 immediately dorsal to the first alveolus. A second opening, the premaxillary foramen, is  
221 positioned more posteriorly and dorsally, at a point just anteroventral to the external naris (Fig.  
222 3A, pmxf). The anterior premaxillary foramen and premaxillary foramen communicate via a

223 deep groove (Sereno, 1991:fig. 6D). CT scans of NHMUK PV RU B17 demonstrate that the  
224 anterior premaxillary foramen is connected to the anterior palatal foramen (Fig. 3D, apf), which  
225 opens on the palate, anterior and medial to the first premaxillary tooth; however, the  
226 premaxillary foramen does not communicate with the anterior palatal foramen on either side of  
227 NHMUK PV RU B17, instead leading to a short, blind-ending canal (Fig. 3F, pmxf) (*contra*  
228 Sereno, 1991). A second palatal foramen described by Sereno (1991) medial to the second and  
229 third premaxillary teeth can be seen on the surface of the segmented NHMUK PV RU B17  
230 ‘snout’ block but its route through the premaxilla cannot be traced. The narial processes (‘pre-  
231 narial process’ *sensu* Thulborn [1970]) of the premaxillae are anteroposteriorly broad at their  
232 bases and taper slightly as they extend dorsally and slightly posteriorly (Thulborn, 1970). Their  
233 dorsal portions are missing in all specimens. Nevertheless, they clearly separate the external  
234 nares anteriorly (Fig. 3E), though it is uncertain whether the internarial bar was complete and, if  
235 it was complete, how it might have contacted the anterior processes of the nasals. The maxillary  
236 processes (‘post-narial process’ *sensu* Thulborn [1970]) are preserved on both sides of NHMUK  
237 PV RU B23 (Fig. 2A, B, pmx), NHMUK PV R8501 and NHMUK PV R11956; CT scans  
238 demonstrate that in NHMUK PV RU B23 the maxillary process of the premaxilla extensively  
239 overlaps the dorsal margin of the maxilla and is dorsally overlapped by the ventrolateral edge of  
240 the nasals. The process is anteroposteriorly broad ventrally, but narrows slightly as it extends  
241 dorsally prior to angling sharply posterodorsally, giving it a kinked appearance in lateral view.  
242 The dorsal-most part of the process tapers to a sharp point that is wedged between the nasals and  
243 maxilla. A short posterior process of the premaxilla is preserved on the right side of NHMUK  
244 PV RU B17 and bears a dorsomedially-directed facet on its dorsal surface (Fig. 3B, E, a.mx);  
245 although not preserved in articulation in the scanned specimens, this facet probably fitted against

246 the anteroventral corner of the maxilla. The posterior process lacks alveoli, thereby forming a  
247 short diastema between the premaxillary and maxillary tooth rows of *Lesothosaurus*, a feature  
248 absent from most previous skull reconstructions (e.g., Thulborn, 1970; Weishampel & Witmer,  
249 1990; Sereno, 1991; Norman et al., 2004). CT scans of NHMUK PV RU B17 confirm that the  
250 anterior portions of the premaxillae meet along a dorsoventrally tall, vertical butt joint (i.e.,  
251 adjacent bones contact at straight, squared-off edges) (Weishampel & Witmer, 1990).  
252 Posteriorly, the premaxilla is an inverted ‘L’-shape in transverse section, due to the presence of a  
253 dorsally vaulted palate that forms an angle of  $\sim 120^\circ$  with the alveolar margin of the bone and  
254 meets its counterpart at a butt joint along the midline (Fig. 3E). The thickness of the premaxillary  
255 palate decreases posteriorly and its contact with the vomer (if any was present) is not preserved.

## 256 **Maxilla**

257 The maxilla is triangular in lateral view and encompasses all but the posterodorsal portion of the  
258 antorbital fossa (Figs. 2 and 4). Both maxillae are preserved in NHMUK PV RU B17 (‘palatal’  
259 block), NHMUK PV RU B23, NHMUK PV R8501 and NHMUK PV R11956 although all  
260 elements are slightly damaged. Most of the damage is found either at the anterior margin (e.g.,  
261 NHMUK PV RU B17) or dorsally (e.g., NHMUK PV R8501); however, the morphology of the  
262 entire maxilla is well-represented by these specimens when taken collectively. The number of  
263 maxillary teeth varies from a minimum of 12 (in the incomplete left maxilla of NHMUK PV  
264 R11956) to at least 15 (in the incomplete right maxilla of NHMUK PV R8501). CT scans of  
265 NHMUK PV RU B17 (which possesses 12 left and 14 right maxillary tooth positions) reveal the  
266 presence of four replacement teeth in the right maxilla, although it is likely that more teeth are  
267 present and cannot be resolved in the scans. In all specimens, the dentition extends for almost the  
268 full length of the alveolar ramus (Fig. 4F).

269 In lateral view, the rounded anteroventral corner of the maxilla is continuous with the  
270 ascending ramus of the maxilla dorsally (Fig. 4A, ascr), which forms the anterior margin of the  
271 antorbital fossa, and the alveolar margin posteriorly (Fig. 4A, alr). The dorsal margin of the  
272 ascending ramus is overlapped by the lateral margins of both the premaxilla and the nasal along  
273 short scarf joints (i.e., adjacent bones overlap along beveled edges) (Fig. 4A; Norman et al.,  
274 2004). The anterior margin of the ascending ramus forms an angle of approximately 45° with  
275 respect to the long axis of the alveolar ramus. The lateral surface of the alveolar margin is  
276 generally flat, weakly concave or bears only an indistinct longitudinal swelling; thus a maxillary  
277 buccal emargination is effectively absent in *Lesothosaurus diagnosticus* (Thulborn, 1970;  
278 Galton, 1973, 1978; Sereno, 1991; Knoll, 2008). This area is pierced by a line of up to six small,  
279 rounded neurovascular foramina of subequal diameter.

280 The maxilla forms most of the boundaries of the antorbital fossa (Fig. 4A, B, aof); CT scans  
281 of NHMUK PV RU B17 and NHMUK PV RU B23 reveal that the medial wall of the fossa is  
282 exceptionally thin in comparison to the alveolar ramus and ascending ramus. A distinct trough is  
283 present between the medial wall of the antorbital fossa and the supralveolar lamina (Fig. 4E, sal;  
284 Sereno, 1991; Witmer, 1997a; Norman et al., 2004). Within the floor of this trough is a foramen  
285 that communicates with the external neurovascular foramina, as is typically found in a diversity  
286 of dinosaurs (Witmer, 1997a). Both maxillae of NHMUK PV RU B17 and the left maxilla of  
287 NHMUK PV RU B23 feature a deeply depressed area in the anteroventral corner of the  
288 antorbital fossa (Fig. 4A, B, ad) perhaps indicative of an incipient pneumatic recess. Indeed, a  
289 “deep and hemispherical” depression occurs in this area in MNHN LES 17 (Knoll, 2002a:238).  
290 An opening is present in this region in *Heterodontosaurus* (anterior antorbital fenestra of  
291 Norman et al. [2011], accessory antorbital fenestra of Sereno [2012]), *Haya* (maxillary fenestra:

292 Makovicky et al., 2011), *Hypsilophodon* (Galton, 1974), and in theropods (promaxillary fenestra:  
293 e.g. Sampson & Witmer 2007). The posterodorsal edge of both maxillae in NHMUK PV RU  
294 B17 possess a rounded notch that forms the anterior margin of the small antorbital fenestra,  
295 which is bounded by the lacrimal posteriorly (Fig. 4B–D).

296 CT scans of NHMUK PV RU B23 demonstrate that the anterior process of the lacrimal fits  
297 into a slot in the tip of the ascending process of the maxilla (Sereno, 1991); posteriorly, the  
298 posterodorsal edge of the maxilla meets the lacrimal along a simple, rounded butt contact. The  
299 posteroventral maxilla features a gently everted surface above the tooth row and underlies the  
300 anterior ramus of the jugal, which it contacts via a short, dorsomedially-directed scarf joint (Fig.  
301 4A, B, a.j; Thulborn, 1970; Sereno, 1991). The maxilla also possesses a short anterior process  
302 (Fig. 4A, a.pmx) that is laterally overlapped by the maxillary process of the premaxilla and  
303 overlies the palatal process of the premaxilla (Weishampel & Witmer, 1990; Sereno, 1991;  
304 Norman et al. 2004).

305 The internal surface of the maxilla expands above the tooth row to form a longitudinal  
306 medial maxillary shelf that has extensive contact with the palatine (Fig. 4C–E, mxsh) as seen in  
307 NHMUK PV R8501 and in CT scans of NHMUK PV RU B17. Sereno (1991:fig.4) illustrated a  
308 line of ‘special foramina’ (*sensu* Edmund, 1957) in a referred specimen (SAM unnumbered) and  
309 in his skull reconstructions (Sereno, 1991:fig. 11). However, CT scans of NHMUK PV RU B17  
310 and NHMUK PV RU B23 do not resolve these features, nor is there any unambiguous indication  
311 of their presence in other specimens (e.g., NHMUK PV R8501).

312 **Nasal**

313 Both nasals are preserved in NHMUK PV RU B23 (Figs. 2C and 5A, B), NHMUK PV R8501  
314 and NHMUK PV R11956, although the majority are damaged: both nasals of NHMUK PV RU  
315 B23 and the right nasal of NHMUK PV R11956 are missing their anterior ends and the right  
316 nasal of NHMUK PV R8501 is broken posteriorly. However, almost complete left nasals are  
317 present in NHMUK PV R8501 and NHMUK PV R11956. A small fragment of the left nasal is  
318 present in the ‘palatal’ block of NHMUK PV RUB 17 (Fig. 2E, n). In dorsal view, the nasal is an  
319 elongate, subtriangular bone that expands laterally at its mid-section to form a short, laterally  
320 extending triangular process. The rest of the element tapers to form subrectangular processes  
321 anteriorly and posteriorly (Thulborn, 1970; Sereno, 1991). It is approximately 4.5 times as long  
322 as it is wide. In transverse section, each nasal is dorsally arched, so that the midline contact  
323 between them lies in a shallow depression that extends along the top of the snout. The nasals  
324 meet at a straight, vertical butt joint along the midline. The lateral margin of the nasal overlaps  
325 the dorsal edges of the premaxilla, maxilla and lacrimal via scarf joints (Fig. 5B, a.pmx, a.mx,  
326 a.l); the contact with the lacrimal is very short (Norman, 1984a; Sereno, 1991). The  
327 posterolateral margin of the nasal is deeply embayed to accommodate the prefrontal; CT scans  
328 demonstrate that the prefrontal extensively overlaps the nasal (Fig. 5A, a.pf). The posterior tip of  
329 the nasal overlaps the frontal (Fig. 5B, a.f), as described by Thulborn (1970) and Weishampel  
330 and Witmer (1990). In lateral view, the dorsal margin of the nasal is almost flat. Although all of  
331 the specimens are either crushed or incomplete, the anterior margin of the left nasal in NHMUK  
332 PV R8501 is smoothly concave and clearly formed the posterodorsal margin of the external  
333 naris.

### 334 **Lacrimal**

335 The lacrimal has an inverted 'L'-shape and it separates the antorbital fossa from the orbit. Both  
336 lacrimals are preserved in NHMUK PV RU B23 (Fig. 2A, B, I), NHMUK PV R8501 and  
337 NHMUK PV R11956; only the left element is present in NHMUK PV RUB 17 (Fig. 2E, G, I).  
338 The short anterior ramus of the lacrimal is rounded in lateral view and inserts into a slot at the tip  
339 of the ascending ramus of the maxilla (Fig. 5C, D, a.mx) (Serenó, 1991). The lacrimal shaft  
340 bears a dorsal facet that articulates with the ventral surface of the prefrontal via a broad  
341 ventrolaterally-inclined scarf joint (Fig. 5C–E, a.pf). The lateral surface of the shaft is gently  
342 convex anteroposteriorly while its posterior surface is dorsoventrally and mediolaterally concave  
343 (Fig. 5E). The medial surface bears a large triangular depression bounded anteriorly and  
344 posteriorly by distinct ridges (Fig. 5D). The opening for the nasolacrimal canal is visible on the  
345 posterior surface of the lacrimal shaft (Fig. 5E, lc) (Norman et al., 2004), but its path through the  
346 element cannot be fully traced, although the canal likely traversed the lacrimal's anterior ramus  
347 as in *Hypsilophodon*, *Plateosaurus*, and most other dinosaurs (Witmer, 1997a). A thin lamina  
348 arises from the anteromedial margin of the lacrimal shaft and the ventromedial margin of the  
349 anterior process. This sheet of bone forms the posterodorsal portion of the antorbital fossa medial  
350 lamina, and also defines the posterior margin of the antorbital fenestra. The ventral ramus tapers  
351 posteroventrally to a slender point, overlapping the dorsomedial aspect of the anterior process of  
352 the jugal in a long but narrow scarf joint (Fig. 5C, a.j) (Weishampel & Witmer, 1990; Sereno,  
353 1991).

#### 354 **Prefrontal**

355 Both prefrontals are preserved in NHMUK PV RU B23 (Fig. 2C, pf), NHMUK PV R8501 and  
356 NHMUK PV R11956. They form the anterodorsal margin of the orbit. Viewed dorsally, the  
357 prefrontal is a teardrop-shaped bone with a rounded anterior margin and tapering, slender

358 posterior process (Norman et al., 2004). In transverse section, the prefrontal is flat with a  
359 dorsoventrally thick main body and a thin lateral extension that forms the orbital margin; this  
360 extension bears a facet on its anterodorsal surface that articulates with the palpebral (Fig. 5F,  
361 a.pap; Thulborn, 1970). The main body of the prefrontal extensively overlaps the posterolateral  
362 margin of the nasal and the lateral margin of the frontal (Fig. 5G, a.n, a.f; Thulborn, 1970). A  
363 short, tapering ventral process arises from the posterolateral corner of the main body and forms  
364 an extensive contact with the dorsal surface of the lacrimal shaft along an oblique scarf joint  
365 (Fig. 5G, a.l).

### 366 **Palpebral**

367 Both palpebrals are preserved in NHMUK PV RU B23 (Figs. 2 and 6H–J) and NHMUK PV  
368 R8501; an isolated right palpebral is present in the ‘braincase’ block of NHMUK PV RU B17.  
369 They do not traverse the entire diameter of the orbit, although rugosities on the anterior margin  
370 of the postorbital suggest that in life the palpebral was connected to the postorbital by the  
371 supraorbital membrane. The palpebral shaft is ovoid in transverse section and is bowed laterally  
372 in dorsal view (Fig. 6I). The expanded base features short dorsal and anterior processes that  
373 articulate with the prefrontal in NHMUK PV RUB 23 (Fig. 6H); there is a point contact between  
374 the palpebral and lacrimal (Fig. 6J), but this is not extensive (Thulborn, 1970; Sereno, 1991;  
375 *contra* Weishampel & Witmer, 1990).

### 376 **Frontal**

377 Both frontals are preserved in articulation in NHMUK PV RU B23 (Figs. 2 and 5H–I) and  
378 NHMUK PV R8501. They are quadrilateral in dorsal view and the frontal is approximately 3.5  
379 times longer than it is wide (as measured at the midlength of the orbital margin). The frontals are

380 widest posteriorly, taper anteriorly, and bear shallow lateral embayments that form the dorsal  
381 margins of the orbits (Thulborn, 1970). In lateral view, the frontals are arched anteroposteriorly,  
382 giving the skull roof a rounded profile. In transverse section, the anterior half of the frontal is  
383 dorsoventrally thickest in its central part (corresponding to the position of the ventral ridge, see  
384 below) and tapers in thickness medially and laterally; posteriorly, the ventral ridge merges into  
385 the body of the bone. The frontals are dorsally arched in transverse section. The interfrontal  
386 suture is straight and the frontals contact each other via a vertical butt joint (Weishampel &  
387 Witmer, 1990). The pointed anterior tips of the frontals insert between and underlap the nasals  
388 and prefrontals (Fig. 5H, a.n, a.pf). The sharp lateral edge of the frontal forms the central third of  
389 the dorsal orbital margin. The orbital margins are smooth and lack the short striations seen in  
390 some other small ornithischians. Posterior to the orbit, the frontals expand posterolaterally to  
391 meet the postorbital in a complex, undulating suture (Fig. 5H, I, a.po). A shallow, well-defined  
392 supratemporal fossa excavates the posterolateral corner of the frontal (Thulborn, 1970) and the  
393 frontal makes a small contribution to the anteromedial margin of the supratemporal fenestra (Fig.  
394 5I, stf). The posterior margin of the frontal contacts the parietal along an undulating contact; fine  
395 interdigitations can be discerned along this contact in NHMUK PV RU B23, but these could not  
396 be segmented (Weishampel & Witmer, 1990). The ventral surface of frontal possesses a low  
397 rounded ridge (the crista cranii; Fig. I, vr) that extends parallel to the orbital margin and which  
398 continues onto the medial surface of the postorbital. This ridge defines the medial margin of the  
399 orbital cavity and the lateral margin of the shallowly concave endocranial cavity, in the region of  
400 the olfactory tract and cerebral hemispheres (Fig. 5I, ot, cer?). The trough for the olfactory tract  
401 extends along the medial part of the ventral surface of the anterior half of the frontals;  
402 posteriorly, the depression between the contralateral cristae cranii widens for the area occupied

403 by the cerebral hemispheres. The anterolateral (capitate) process of the laterosphenoid may have  
404 contacted the ventral surface of the skull roof near to the frontal-parietal-postorbital contact (Fig.  
405 5I, a.lat).

#### 406 **Parietal**

407 The parietals are preserved in articulation in NHMUK PV RU B23, NHMUK PV R8501 and  
408 NHMUK PV R11004, and form the roof of the braincase and the medial and posterior margins  
409 of the supratemporal fenestrae (Figs. 2C and 5J–L). They are strongly dorsally arched in  
410 transverse section (Thulborn, 1970), and there is no sagittal crest (Knoll, 2002a; *contra*  
411 Weishampel & Witmer, 1990; Sereno, 1991; Norman et al., 2004). The midline suture between  
412 the parietals is visible in CT scans of NHMUK PV RU B23, but is very faint compared to other  
413 sutures, suggesting that the elements may have partially fused. Nevertheless, the straight  
414 interparietal suture is visible externally in this specimen and also in NHMUK PV R11004. The  
415 gently undulating anterior margin of the parietal contacts the posterior edge of the frontal along  
416 an interdigitating suture. The short anterolateral process of the parietal contacts the frontal and  
417 the medial process of the postorbital via a rounded butt joint (Fig. 5J, K, a.f/a.po; Sereno, 1991).  
418 The parietals are weakly constricted between the supratemporal openings (Fig. 5K, stfe). The  
419 posterior margins of the parietals expand to form prominent posterolaterally extending processes  
420 that diverge from the midline of the skull at angles of approximately 45° in dorsal view: together  
421 these processes and the straight posterior margin of the main parietal body form a deep  
422 embayment that accommodates the dorsal and lateral margins of the supraoccipital (Fig. 5K, L,  
423 a.so). In dorsal view, the posterolateral process forms the posteromedial margin of the  
424 supratemporal fenestra, and in posterior view (Fig. 5L) it is dorsoventrally expanded and  
425 overlaps the dorsal and medial surfaces of the squamosal medial process. The ventral margin of

426 the posterolateral process contacts the dorsal margin of the paroccipital process (Fig. 5L, a.ot). It  
427 is likely that the ventrolateral margins of the parietal contacted the laterosphenoid, prootic and  
428 otoccipital; however, the nature of these contacts cannot be visualized in NHMUK PV RU B23,  
429 NHMUK PV R8501 or NHMUK PV R11004 due to disarticulation, crushing, the presence of  
430 matrix or difficulty with segmentation. There is some equivocal evidence for the presence of a  
431 small opening between the parietal and paroccipital process in NHMUK PV RU B23 (potentially  
432 indicative of the posttemporal foramen), but this cannot be confirmed due to deformation and  
433 bone loss in the region in this and all other specimens. There appears to be no evidence for the  
434 presence of a foramen between the parietal and supraoccipital (*contra* Sereno, 1991).

#### 435 **Jugal**

436 The jugal is incomplete in almost all specimens, although a nearly complete left jugal missing  
437 the ends of some processes is present in NHMUK PV R8501. Combining information from this  
438 specimen with partially preserved jugals from the left sides of NHMUK PV RU B17 and  
439 NHMUK PV RU B23 (Figs. 2 and 5M, N) allows most features to be reconstructed. The jugal  
440 consists of three processes that arise from a central main body: anterior (maxillary), dorsal  
441 (postorbital) and posterior (quadratojugal). In lateral view, the anterior process is slender, being  
442 longer than it is tall, and it forms the ventral margin of the orbit. The process tapers anteriorly  
443 and extends close to the antorbital fossa though it does not contribute to the margin of the  
444 fenestra. Dorsomedially, the anterior process is overlapped by the ventral ramus of the lacrimal  
445 (Fig. 5N, a.l). It contacts the posterior process of the maxilla ventrally along a ventrolaterally-  
446 directed scarf joint (Fig. 5M, N, a.mx). The lateral surface of the anterior process is rounded and  
447 continuous with the horizontal shelf of the maxilla. CT scans of NHMUK PV RU B17 reveal a  
448 hollow space within the main body of the jugal and a small foramen on the medial surface

449 connects with this space, though a similar foramen cannot be seen in NHMUK PV R8501. These  
450 structures likely represent evidence of blood vessels that passed through the floor of the orbit, as  
451 similar vascular features are found in a range of dinosaurs and extant diapsids (Sampson &  
452 Witmer, 2007). The anterior process is round in transverse section anteriorly, but becomes  
453 dorsoventrally expanded and transversely narrow posteriorly, with thickened dorsal and ventral  
454 margins. The dorsal process of the jugal forms the ventral part of the postorbital bar and the  
455 posteroventral margin of the orbit. It is robust, elongate, tapers dorsally, and bears a long,  
456 triangular facet on its anterolateral surface where it is overlapped by the ventral process of the  
457 postorbital (Fig. 5M, a.po; Weishampel & Witmer, 1990; Norman et al., 2004). A rounded ridge  
458 on its anteromedial surface is continuous with a ridge on the postorbital. The dorsal process of  
459 the jugal approaches but does not contact the squamosal (*contra* Knoll, 2002a, 2002b). A partial  
460 posterior process is present in NHMUK PV R8501 and NHMUK PV RU B17, which indicates  
461 that the depth of the posterior process was greater than that of the anterior process, and that the  
462 posterior process formed the ventral margin of the infratemporal fenestra. The medial surface of  
463 the jugal is gently concave with a thickened, ridge-like ventral border (Fig. 5N).

#### 464 **Postorbital**

465 Both postorbitals are preserved in NHMUK PV RU B23 (Figs. 2A–C and 5O–Q) and NHMUK  
466 PV R8501. It is a triradiate bone in lateral view with anterior, posterior and ventral processes.  
467 These processes radiate from a subtriangular main body that is laterally convex. All three  
468 processes taper distally. The short, stout anterior process joins the frontal via a ‘W’-shaped,  
469 undulating contact in dorsal view (Fig. 5P, a.f). The posterodorsal surface of the anterior process  
470 is excavated by the anterior corner of the supratemporal fossa (Fig. 5P, stfe). The posteroventral  
471 surface of the anterior process bears a small facet that receives the anterolateral process of the

472 parietal and laterosphenoid (Fig. 5Q, a.p/a.lat). The posterior process underlaps the squamosal in  
473 an extensive contact, forming the upper temporal bar; it bears a rounded ridge on its ventrolateral  
474 surface that dorsally bounds the infratemporal fossa. The ventral process overlaps the dorsal  
475 process of the jugal anteriorly and laterally via a long scarf joint, forming the dorsal part of the  
476 postorbital bar and the posterodorsal corner of the orbit. The medial surface of the ventral  
477 process possesses a ridge that is continuous with ridges on the medial surface of the jugal and on  
478 the ventral surface of the frontal (Fig. 5Q, vr). In NHMUK PV RU B23, a slightly rugose area is  
479 present along the orbital margin at the junction of the ventral and anterior processes (visible on  
480 both postorbitals), probably representing an attachment site for the supraorbital membrane that  
481 also would have attached to the palpebral bone anteriorly (Fig. 5O, rug; Maidment & Porro,  
482 2010).

### 483 **Quadratojugal**

484 Only a small fragment of the left quadratojugal is preserved in NHMUK PV RU B23 (Fig. 2A,  
485 qj), attached to the quadrate, but both elements are present in NHMUK PV R8501. The  
486 quadratojugal formed the posteroventral corner of the infratemporal fenestra and part of the  
487 lateral wall of the adductor chamber. In lateral view, it has an isosceles triangle-shaped outline,  
488 with the apex of this triangle pointing dorsally. The anterior and posterior margins are gently  
489 concave and the ventral margin is slightly convex, and the posterior margin is closely appressed  
490 to the quadrate. The lateral surface is smooth and there is no indication of a paraquadratic  
491 foramen. It cannot be determined if the quadratojugal made contact with the squamosal, but a  
492 point contact seems plausible as the quadratojugal extended dorsally for approximately half of  
493 the height of the quadrate. The contact between the quadratojugal and jugal is not preserved in  
494 any specimen.

**495 Squamosal**

496 Both squamosals are preserved in NHMUK PV RU B23 (Figs. 2A–C and 6A–C) and NHMUK  
497 PV R8501. It is a complexly-shaped, tetradial bone with anterior, medial, prequadratic and  
498 postquadratic processes. The anterior process is distally expanded (Serenio, 1991) and dorsally  
499 and medially overlaps the posterior process of the postorbital to form the upper temporal bar  
500 (Fig. 6A, a.po). The dorsal surface of the anterior process is drawn up into a rounded ridge that  
501 laterally bounds the supratemporal fossa (Fig. 6B, stfe). The medial process of the squamosal  
502 forms the posterior edge of the supratemporal fenestra and laterally and ventrally overlaps the  
503 posterolateral process of the parietal (Thulborn, 1970). A sharp lateral ridge between the body of  
504 the squamosal and the prequadratic process forms a well-defined sulcus for the origin of the *m.*  
505 *adductor mandibulae externus superficialis*, in the posterodorsal corner of the infratemporal  
506 fenestra. The long, tapering prequadratic process of the squamosal medially and dorsally  
507 overlaps the anterior surface of the quadrate (Fig 6A, a.q; Sereno, 1991). The much shorter  
508 postquadratic process overlies the anterior surface of the paraoccipital process (Fig. 6B, C, a.ot).  
509 The small cotylus receiving the head of the quadrate is deep, cup-shaped and ventrally-directed  
510 (Fig. 6A, cot), and is formed by the junction between the pre- and postquadratic processes  
511 (Norman et al., 2004). The main body of the squamosal faces dorsally and dorsolaterally, and is  
512 small, subtriangular in outline and has a gently convex external surface.

**513 Quadrate**

514 Only the left quadrate of NHMUK PV RUB 23 is preserved, but both are present in NHMUK PV  
515 R8501 (Figs. 2A, B and 6D–G). It is composed of a stout sub-vertically inclined shaft, whose  
516 anterior surface supports two thin sheets of bone: the medial pterygoid ramus and anterolateral

517 ramus (Norman et al., 2004). Consequently, the quadrate has a ‘V’-shaped horizontal section at  
518 mid-shaft . The ventral part of the anterior margin of the anterolateral ramus contacted the  
519 quadratojugal (Fig. 6D), whereas the dorsal portion formed a long overlapping contact with the  
520 prequadrate process of the squamosal, with these two elements excluding the quadrate from  
521 participation in the infratemporal fenestra. The transversely narrow head of the quadrate (Fig.  
522 6F) articulates with the ventral cotylus on the squamosal. The pterygoid ramus is dorsoventrally  
523 tall and has a rounded anterior margin (Fig. 6E, ptw); it laterally overlaps the quadrate wing of  
524 the pterygoid. The quadrate shaft of *Lesothosaurus* is strongly anteriorly arched (Weishampel &  
525 Witmer, 1990), resulting in the surface of the jaw joint being directed posteroventrally. In  
526 posterior view, the shaft is slightly medially-inclined, shallowly excavated, and mediolaterally  
527 narrowest above the jaw joint but transversely expanded to form the joint itself (Fig. 6G; Sereno,  
528 1991). There are distinct lateral and medial condyles, which are separated by a shallow groove  
529 (Norman et al., 2004; *contra* Weishampel & Witmer, 1990). The medial condyle is larger than  
530 (and ventrally displaced relative to) the lateral condyle.

### 531 **Palate**

532 The palate is completely preserved (except for the premaxillae) in the ‘palatal’ block of NHMUK  
533 PV RU B17 (Figs. 2G and 7). The right palatal complex is largely articulated; the left palatal  
534 complex is displaced dorsally relative to the ventral braincase and maxilla, and the ectopterygoid  
535 has separated from the pterygoid. The palate is almost certainly present in NHMUK PV RU B23  
536 but cannot be visualized in CT scans due to the presence of dense matrix that limits X-ray  
537 penetration. The pterygoids and a left ectopterygoid are present in NHMUK PV R8501, as well as  
538 some more anteriorly positioned elements that are badly crushed and difficult to interpret. All of  
539 the elements described in this section are based on NHMUK PV RU B17. The palate is dorsally

540 vaulted in transverse section, and dorsally arched in lateral view (Fig. 7B). The maxillary shelves  
541 laterally border the internal nares (choana) and appear to exclude the premaxillae. The palatines  
542 and anterior processes of the pterygoids form the posterior margin of the internal nares, while the  
543 vomers separate them at the midline. The palatal or suborbital fenestra ('postpalatine fenestra' of  
544 Thulborn [1970] and Sereno [1991]) is very small and bordered by the maxilla, ectopterygoid,  
545 pterygoid and palatine; the subtemporal opening is bordered by the posterior tip of the maxilla,  
546 jugal, ectopterygoid, pterygoid, quadrate and, presumably, the quadratojugal.

#### 547 **Vomer**

548 CT scans demonstrate that the vomers (Fig. 7,v) are elongate elements fused at the midline  
549 (Norman et al., 2004) to form the medial margins of the internal nares. They are transversely thin  
550 and dorsoventrally tall at their anterior ends (*contra* Sereno, 1991), being teardrop-shaped in  
551 lateral view. The vomers taper posteriorly to a point that lies between the anterior processes of  
552 the pterygoids. The vomers do not appear to contact the palatines. As neither the premaxillae nor  
553 the anterior portions of the maxillae are preserved in the 'palatal block' of NHMUK PV RU B17,  
554 it is unclear which of these elements the vomers contacted anteriorly.

#### 555 **Palatine**

556 Both palatines are complete in NHMUK PV RU B17 (Figs. 2G and 8A–D), with the right  
557 palatine in articulation with its respective maxilla, pterygoid and ectopterygoid. It roofs the  
558 posterior palate and forms the posterior margin of the internal nares, although the anterior margin  
559 of the palatine is not deeply embayed (*contra* Thulborn [1970]). The palatine of *Lesothosaurus*  
560 consists of an extensive horizontal lamina and a short vertical lamina (restricted to the posterior  
561 half of the element) that are joined laterally (Fig. 8A, hl, vl); thus, the posterior palatine is 'L'-  
562 shaped in transverse section. The vertical lamina possesses a concave facet on its lateral surface

563 that articulates with the medial shelf of the maxilla (Fig. 8C, a.mx); anteriorly, the thickened  
564 lateral edge of the horizontal lamina also contacts the maxilla, resulting in a long and extensive  
565 suture between these bones (Thulborn, 1970; Sereno, 1991; Norman et al., 2004). The lateral  
566 aspect of the vertical lamina also contacts the internal surface of the jugal on the disarticulated  
567 left side of NHMUK PV RU B17. Unfortunately, the jugal is missing from the articulated right  
568 side and, as a result, the position of the palatine-jugal contact cannot be firmly established (but  
569 see Sereno, 1991). A small portion of the vertical lamina may have contacted the internal surface  
570 of the ventral ramus of the lacrimal, but this is uncertain. A short, rounded ridge, formed by the  
571 vertical lamina, is prominent along the lateral edge of the dorsal surface of the horizontal lamina  
572 before bifurcating anteriorly (Fig. 8A). One branch continues along the lateral margin of the  
573 element; the other is anteromedially-directed and crosses the dorsal surface of the palatine. As a  
574 result, the dorsal surface of the palatine bears two prominent depressions (Fig. 8A, m.pt, pnr).  
575 The posterior depression is larger and has been identified as a muscular fossa for the *M.*  
576 *pterygoideus dorsalis* (Witmer, 1997a); the anterior depression is smaller and has been identified  
577 as potentially a palatine pneumatic recess (Witmer, 1997a). The medial margins (Fig. 8D) of the  
578 palatines closely approach each other and the midline elements (vomeres, parasphenoid), but  
579 contact appears unlikely. The posterior tip of the horizontal lamina makes a short contact with  
580 the anterior surface of the ectopterygoid (Fig. 8D, a.ect; Sereno, 1991; *contra* Weishampel &  
581 Witmer, 1990). The ventromedial aspect of the horizontal lamina extensively overlies the  
582 anterior process of the pterygoid, tapering posteromedially to a spike-like projection.

### 583 **Ectopterygoid**

584 The ectopterygoid (Fig. 8E-G) is a hooked, 'U'-shaped element that connects the pterygoid and  
585 palatine with the maxilla (and possibly jugal). The base of the ectopterygoid is broad and

586 extensively contacts the dorsal surfaces of the main body and flange of the pterygoid (not the  
587 quadrate ramus, *contra* Weishampel & Witmer [1990]), contributing to the base and posterior  
588 margin of the pterygoid flange. The anterior surface of the ectopterygoid makes a short contact  
589 with the posterior tip of the palatine on the right side of NHMUK PV RUB 17 (Fig. 8F, a.pl).  
590 The ectopterygoid tapers anterolaterally to a rounded articular facet but, due to disarticulation of  
591 this specimen, this surface is free on both sides of NHMUK PV RUB 17 (and also in NHMUK  
592 PV R8501). Most likely, it contacted the medial surface of the maxilla, although contact with the  
593 jugal cannot be ruled out. The dorsal edge of the ectopterygoid bears a sharp ridge (Serenó,  
594 1991). CT scans reveal a hollow cavity within both ectopterygoids of NHMUK PV RUB 17.  
595 Although this cavity might represent an ectopterygoid pneumatic recess, the cavity is fully within  
596 the bone and does not open externally, which is a requirement of pneumatic systems (Witmer,  
597 1997a), suggesting that it is simply an open cancellous structure, as seen in many extant diapsids,  
598 especially squamates (Fig. 8E–G; Witmer, 1997a).

### 599 **Pterygoid**

600 The pterygoid is the largest bone of the palate and links the braincase and sidewalls of the skull  
601 (Figs. 2G, H and 8H–J). It consists of a main body, anterior process, a quadrate ramus, and the  
602 pterygoid flange. The anterior process is long and formed of a vertical septum of bone that is  
603 tallest anteriorly and tapers posteriorly. This process appears to have contacted the vomer  
604 medially (Thulborn, 1970), thus contributing to the medial margin of the internal naris (Fig. 8H–  
605 J). The anterior process of the pterygoid underlies the ventral surface of the horizontal lamina of  
606 the palatine (Fig. 8I, a.pl). Posteriorly, a thin, horizontal lamina of bone extends laterally from  
607 the anterior process and progressively widens and deepens to form the main body of the  
608 pterygoid. The left and right main bodies meet in a short, dorsoventrally deep midline butt joint

609 posterior to the pterygoid flange (Fig. 8H–J, a.pt), resulting in a long, narrow interpterygoid  
610 vacuity through which the parasphenoid is visible. As noted above for the ectopterygoid, CT  
611 scans reveal hollow cavities within the body of the pterygoid at the level of the pterygoid flange  
612 that remain fully within the confines of the bone, breached only by a minute vascular foramen or  
613 two. The relatively open cancellous structure of many of the skull bones in NHMUK PV RUB  
614 17 may reflect its subadult or even juvenile status. The quadrate ramus flares posterodorsally and  
615 laterally from the main body of the pterygoid and is transversely thin (as well as laterally arched)  
616 in transverse section (Fig. 8I, J, qw), with a thickened, inturned ventral edge and an undulating  
617 posterior margin. The quadrate ramus is overlapped laterally by the pterygoid ramus of the  
618 quadrate (Fig. 8H, a.q), forming the medial margin of the subtemporal fenestra. Medial to the  
619 base of the quadrate ramus is a deep, posterodorsally-facing concavity that articulates with the  
620 basiptyergoid processes (Fig. 8J, a.bs). This concavity is delimited ventromedially by a  
621 prominent bony projection. The pterygoid flange of *Lesothosaurus* is triangular in ventral and  
622 lateral views, with the apex directed anteriorly (Fig. 8H, I, ptf), and lacks strong excavations on  
623 its dorsal surface. A rounded concavity on the posterior margin of the pterygoid flange (ventral  
624 to the basal articulation) marks the origin of *M. pterygoideus ventralis* (Fig. 8J, m.pt).

## 625 **Braincase**

626 NHMUK PV RUB 23 almost certainly preserves a complete braincase but, with the exception of  
627 the supraoccipital and otoccipital (opisthotic and exoccipital), it could not be visualized in the  
628 CT scans due to high-density matrix within the braincase. NHMUK PV R8501 contains a  
629 potentially complete, but partially disarticulated and slightly distorted braincase, whereas the  
630 ‘palatal’ block of NHMUK PV RUB 17 preserves a disarticulated basisphenoid (with  
631 parasphenoid), basioccipital and a left laterosphenoid/prootic. Another basisphenoid is preserved

632 in the ‘braincase’ block of NHMUK PV RU B17 and an otoccipital is present in NHMUK PV  
633 R11004. Many braincase elements are unfused in all of the aforementioned specimens,  
634 suggesting juvenile status.

### 635 **Prootic and Laterosphenoid**

636 A left prootic with a fragment of laterosphenoid is preserved in the palatal block of NHMUK PV  
637 RU B17 (Fig. 9A, B). As noted by Sereno (1991) a left prootic is also present in NHMUK PV  
638 R8501, though the latter is largely obscured by overlying elements and provides very little  
639 anatomical information. In NHMUK PV RU B17, the left prootic is well preserved but is out of  
640 position such that its lateral surface is now facing dorsally as preserved. Otherwise, the  
641 morphology of the prootic is fairly typical. The exposed lateral surface (Fig. 9A) has a fossa for  
642 the adductor musculature dorsally, a long pointed process posterodorsally that would have  
643 attached to the anterior surface of the otoccipital’s paroccipital process (Fig. 9A, a.pp), and a  
644 ventral portion that would have articulated with the basisphenoid. The anterior margin of the  
645 prootic is strongly incised for the trigeminal foramen (Fig. 9A, CNV), which would have been  
646 completed anteriorly by the laterosphenoid. The contact for the laterosphenoid dorsal to the  
647 trigeminal foramen is well preserved as a stout facet (Fig. 9A, a.ls), as in most dinosaurs. A  
648 fragment of bone attached to this facet that wraps around to the medial side of the specimen is  
649 likely the remnants of the left laterosphenoid (Fig. 9B, ls). The posterior margin the prootic is  
650 marked by the long otosphenoidal crest(Fig. 9A, osp) that separated the adductor domain from  
651 the middle ear domain (Witmer, 1997b). The crest sweeps anteroventrally from the posterodorsal  
652 region (where it probably would have continued onto the otoccipital) down to the basisphenoid  
653 region, probably to the region of the basiptyergoid process, which is typical for diapsids  
654 (Witmer, 1997b). In fact, the dorsolateral wing of the basisphenoid that laterally covers the

655 basisphenoid recess (see below) is probably continuous with the otosphenoidal crest, which is  
656 again typical of other diapsids. The foramen for the facial nerve (cranial nerve VII) is just  
657 posterior to the otosphenoidal crest and thus within the middle ear cavity, which is by far the  
658 most common situation in archosaurs. The anterior margin of the fenestra ovalis (vestibuli) is  
659 also preserved posterior to the otosphenoidal crest, posterodorsal to the facial nerve foramen  
660 (Fig. 9A, fo). The CT scan data reveal the medial aspect of the bone (Fig. 9B), which is again  
661 very conservative. The internal acoustic meatus (Fig. 9B, iam) is an oval depression posterior to  
662 the trigeminal foramen that transmits the facial nerve canal, as well as the canals for the two  
663 major branches of the vestibulocochlear nerve. The prootic portion of the vestibular pyramid (the  
664 conical medial eminence formed by the prootic and otoccipital that houses the vestibule of the  
665 inner ear) is well preserved, opening into the substance of the prootic bone from its posterior  
666 surface and creating a large vestibular chamber. The CT scan data shows that the lateral  
667 (horizontal) semicircular canal opens into this space. Anterior to the vestibular pyramid and  
668 dorsal to the internal acoustic meatus is the prootic portion of the fossa for the floccular lobe of  
669 the cerebellum, which would have been completed by the otoccipital (Fig. 9B, fr). The preserved  
670 fragment of the laterosphenoid is largely uninformative.

### 671 **Otoccipital**

672 The exoccipitals and opisthotics are indistinguishably fused to form the otoccipitals in NHMUK  
673 PV RU B23, NHMUK PV R8501 and NHMUK PV R11004. Only the posterior portions could  
674 be resolved in scans of NHMUK PV RU B23 (Fig. 9C–E); anteriorly, the otoccipitals  
675 presumably articulated with the basisphenoid and prootic. In posterior view (Fig. 9D), the  
676 otoccipitals form the lateral margins of the foramen magnum (Fig. 9D, fm); they flare laterally to  
677 gently rounded, non-pendant paroccipital processes, the distal ends of which are slightly

678 dorsoventrally expanded. The anterior surface of the paroccipital process is convex and  
679 contacted the postquadratic process of the squamosal. The posterolateral processes of the parietal  
680 rested on the dorsal edge of the paraoccipital process (Fig. 9D, E, a.p) as did the ventrolateral  
681 margins of the supraoccipital (Fig. 9C,E, a,so). In NHMUK PV R8501, a small foramen pierces  
682 the anterior surface of the paroccipital process (Sereno, 1991:fig. 13B, labelled post-temporal  
683 foramen) and it is plausible that this continued posteriorly to open on the posterior surface of the  
684 process as there are indications of a foramen in this area in NHMUK PV R8501, though this  
685 canal is not detectable in CT scans of NHMUK PV RU B23. Ventrally, the margin of the process  
686 forms a distinct ridge, the otosphenoidal crest. The ventral process of the otoccipital tapers to  
687 form the medially concave margin of the foramen magnum prior to expanding lateromedially at  
688 its ventral end to form a footplate that articulated with the dorsal surface of the basioccipital. The  
689 otoccipital makes a very small contribution to the dorsolateral corner of the occipital condyle  
690 (NHMUK PV R8501). Due to the orientation of the otoccipital in NHMUK PV R8501, and  
691 damage in NHMUK PV RU B23, the morphology of the jugular foramen cannot be determined  
692 in these specimens. However, the anterior margin of the otoccipital in NHMUK PV R11004,  
693 although partially damaged and obscured by matrix, does appear to bear at least one small  
694 emargination that might represent the posterior margin of the jugular foramen: the dorsal margin  
695 of this embayment is formed by a distinct crest, which extends on to the ventral surface of the  
696 paroccipital process for a short distance and probably represents the posterodorsal part of the  
697 crista interfenestralis. The region where the foramen ovale might be situated is damaged and  
698 covered with matrix, so its preservation in this specimen is equivocal (*contra* Sereno, 1991).  
699 Three possible openings for cranial nerve exits are visible on the internal surface of the right  
700 otoccipital in NHMUK PV R8501 and it seems plausible that these represent the foramina for

701 cranial nerves X (one opening situated anterodorsally) and XII (two openings, situated ventrally)  
702 as proposed by Sereno (1991). However, the external openings of these foramina are obscured by  
703 overlying elements and matrix in NHMUK PV R8501. Nevertheless, the external openings of at  
704 least two, and possibly three foramina, are visible on the lateral surface of NHMUK PV R11004.  
705 Details of the semicircular canals are not determinable in the scans of NHMUK PV RU B23 and  
706 none of the openings in NHMUK PV R8501 can be confidently identified as semicircular canal  
707 openings (*contra* Sereno, 1991).

### 708 **Supraoccipital**

709 The supraoccipital (Fig. 9F) is a single median element forming the dorsal margin of the foramen  
710 magnum and is present in NHMUK PV RU B23 and NHMUK PV R8501. It is trapezoidal in  
711 occipital view, being narrowest dorsally and flaring ventrally. In lateral view, the CT scan of  
712 NHMUK PV RU B23 reveals an anteriorly tapering, thin sheet that underlies the parietal; it is  
713 possible this may represent an anterior process of the supraoccipital but is more likely a broken  
714 fragment of the parietal. The posterior surface bears a rounded median nuchal crest that is most  
715 prominent dorsally and merges gradually into the main body of the bone, disappearing at  
716 approximately midheight (Fig. 9F, nc). The areas to either side of the crest are gently concave.  
717 The ventrolateral margins of the supraoccipital bear small facets for articulation with the dorsal  
718 margins of the paraoccipital process (Fig. 9F, a.pp), while the dorsolateral margins of the  
719 element are bounded by the posterolateral processes of the parietal in posterior view (Sereno,  
720 1991; Norman et al., 2004). Its anterior contacts cannot be resolved in CT scans. The medial  
721 (anterior) surface is deeply concave.

### 722 **Basisphenoid**

723 The basisphenoid is fused to the parasphenoid anteriorly; it presumably joined the prootic,  
724 laterosphenoid and otoccipital dorsally, but these contacts are either not preserved, obscured in  
725 external view or cannot be visualized in CT scans. Most of this description is based on scan data  
726 from the ‘palatal’ and ‘braincase’ blocks of NHMUK PV RU B17 (Fig. 9G–I), but a  
727 parabasisphenoid is also present in NHMUK PV R8501. The basisphenoid is tallest and  
728 narrowest anteriorly, widening and shortening posteriorly. There is a deep excavation on the  
729 dorsal surface of the basisphenoid (Fig. 9G, pf), forming the anterior part of the floor of the  
730 endocranial cavity that communicates with the deep, anteroposteriorly narrow pituitary fossa,  
731 which opens posterodorsal to the base of the parasphenoid rostrum (cultriform process). On  
732 either side of the pituitary fossa, the bone flares posterodorsally and laterally. The sharp ventral  
733 edge of this lamina, almost certainly continuous with the otosphenoidal crest noted above with  
734 the prootic, forms the anterior and lateral margins of the deep basipterygoid recesses (Fig. 9H, I,  
735 bpr) of the middle ear cavity. The basipterygoid processes (Fig. 9G–H, bpp) are short,  
736 anteroposteriorly expanded at their distal ends, have a subtriangular cross-section and are  
737 rounded at their tips. They are deflected anteroventrally at an angle of  $\sim 55^\circ$  (relative to the  
738 parasphenoid) and laterally at angles of  $50^\circ$  (‘palatal’ block) to  $60^\circ$  (‘braincase’ block) from the  
739 midline. Rounded ridges, continuous with the basipterygoid processes, extend posteriorly along  
740 the lateroventral margins of the basisphenoid, helping to define a shallow midline depression that  
741 extends for the full length of the element. Posteriorly these ridges diverge laterally to form  
742 transversely expanded flanges that underlie the basioccipital and form the ventrolateral margins  
743 of the basal tubera; these flanges are separated along the midline by a deep, rounded embayment.  
744 The internal structure of the bone in NHMUK PV RU B17, as noted for other cranial bones in  
745 this specimen, contains relatively large cancellous spaces, presumably filled with marrow in life,

746 that do not open externally. These spaces make it challenging to trace the structures known to  
747 traverse the basisphenoid in other diapsids, such as the cerebral branch of the internal carotid  
748 artery and the abducens nerves (CN VI). The cerebral carotid artery canals should open into the  
749 pituitary fossa, and indeed paired canals open into the posterolateral aspect of the floor of the  
750 pituitary fossa from the basisphenoid recesses. This condition of having the cerebral carotids pass  
751 through the middle ear to enter the basisphenoid deep to the otosphenoidal crest within the  
752 basisphenoid recess is typical for diapsids (Oelrich, 1956; Witmer, 1997b; Sampson & Witmer,  
753 2007; Porter WR & Witmer LM, unpubl data). The canals for the abducens nerves are more  
754 difficult to trace, but there is a candidate pair of canals visible in the CT scan data, although their  
755 external apertures are difficult to see.

#### 756 **Parasphenoid**

757 The cultriform process of the parasphenoid in the ‘palatal’ block of NHMUK PV RU B 17 is  
758 complete and fused to the anterior aspect of the basisphenoid. It projects anteriorly along the  
759 midline between the orbits to the midpoint of the palatines. It is shaped like an inverted triangle  
760 in transverse section, tapers to a sharp, pointed tip and bears a deep dorsal groove (Norman et al.,  
761 2004) for the cartilaginous interorbital septum. CT scans reveal that this groove continues into  
762 the body of the basisphenoid, although it does not connect with the pituitary fossa. The long axis  
763 of the cultriform process is situated at the same level as the long axis of the basisphenoid, just  
764 dorsal to the bases of the basiptyergoid processes, and is not dorsally or ventrally offset.

#### 765 **Basioccipital**

766 A disarticulated basioccipital is preserved in the ‘palatal’ block of NHMUK PV RU B17 (Fig.  
767 9J, K); NHMUK PV R8501 also includes a basioccipital that is partially obscured by

768 surrounding cranial elements. In posterior view, the occipital condyle is shaped like a rounded,  
769 inverted triangle and formed the ventral margin of the foramen magnum. Anterior and lateral to  
770 the condyle are the deep concavities of the basioccipital recesses, which are separated by a low,  
771 sharp ridge at the midline. A pair of hook-shaped processes from the lateral aspect of the  
772 basioccipital form the dorsolateral margins of the tubera. The dorsal surface of the posterior  
773 basioccipital is slightly depressed; anteriorly, a low median ridge and paired lateral ridges form  
774 two distinct depressions. Facets on the anterolateral surfaces of the basioccipital mark its contact  
775 with the basisphenoid; as the basioccipital and basisphenoid are disarticulated in the ‘palatal’  
776 block and the former is lost in the ‘braincase’ block, it appears these elements were not strongly  
777 joined. The basioccipital contacted the otoccipitals dorsolaterally, via two large crescentic facets  
778 (NHMUK PV R8501).

#### 779 **Lower jaw**

780 The lower jaw of *L. diagnosticus* has a nearly straight ventral margin and is only slightly  
781 upturned at its anterior end (Fig. 2). The dentary forms over half of the length of the lower jaw in  
782 lateral view, and there is a well-developed coronoid eminence, though this is not extended  
783 dorsally into a tall, distinct coronoid process. The jaw joint is slightly depressed relative to the  
784 alveolar margin (Fig. 2A). There is an anteroposteriorly elongate external mandibular fenestra  
785 between the dentary, surangular and angular (Fig. 2A, E, F).

#### 786 **Predentary**

787 A predentary is preserved in the ‘snout’ block of NHMUK PV RU B17 (Figs. 2D and 10A–C)  
788 and in NHMUK PV R8501; both examples are preserved in articulation with the dentaries. As  
789 noted in previous descriptions, it is shaped like an arrowhead in ventral view, with a long median

790 ventral keel and slightly shorter lateral processes separated by deep embayments that  
791 accommodate the anterior ends of the dentaries (Fig. 10C, a.d). The oral margin of the  
792 prementary is smooth and straight in lateral view and the anterior tip is not curved dorsally (Fig.  
793 10B). In transverse section, the anterior prementary is shaped like an inverted triangle with a flat  
794 occlusal surface; posteriorly, it becomes 'V'-shaped. Two prominent foramina are visible in  
795 lateral view: the first at the junction between the lateral and ventral processes, and the second  
796 within the lateral process (Fig. 10B, fo). The presence of high-density precipitates at the  
797 prementary-dentary joint makes tracing these openings into the body of the prementary difficult;  
798 however, the abundance of these precipitates suggests that the prementary and its overlying  
799 rhamphotheca was richly supplied with blood vessels and nerves. The ventral keel is triangular in  
800 transverse section, with its dorsal apex fitting between the anterior ends of the dentaries (Norman  
801 et al., 2004). The lateral processes become mediolaterally thin and laterally overlap the dentaries.

## 802 **Dentary**

803 Partial or complete dentaries are known from many specimens (Fig. 2), including NHMUK PV  
804 R8501 (missing only the anterior end of the right dentary), NHMUK PV RU B17 (though  
805 divided between the 'snout' and 'palatal' blocks), NHMUK PV RU B23 (posterior two-thirds of  
806 both dentaries, although only the left dentary, and other individual bones of the lower jaw, could  
807 be resolved in CT scans), and NHMUK PV R11956 (partial posterior parts of both dentaries).  
808 The left dentary of NHMUK PV RU B17 (including portions in the 'snout' and 'palatal' blocks)  
809 preserves 12 tooth positions and the right dentary preserves 17 tooth positions; more teeth are  
810 likely to be present but cannot be resolved in scans. The dorsal and ventral margins of the  
811 dentary are parallel throughout its length (Fig. 10D, E; Sereno, 1991; Weishampel & Witmer,  
812 1990; Norman et al., 2004). The anterior end of the dentary tapers abruptly to a rounded point

813 (Fig. 10D) and twists about its long axis so it meets its opposite ventrally, forming a distinct  
814 ‘spout-shaped’ symphysis. The anterior dentary bears a convex, ventromedial facet that contacts  
815 the ventral prementary process (Fig. 10E, a.pd) and a smaller, flat dorsolateral facet for the lateral  
816 prementary process (Fig. 10D, a.pd). The contact between the lateral prementary processes and the  
817 dentaries are tight while the contact between the dentaries and the ventral prementary process are  
818 patent. CT scans reveal that the anterior dentaries meet each other at an anteroposteriorly short,  
819 flattened midline contact restricted to the lower third of the element (Fig. 10E, a.d). The lateral  
820 surface of the anterior dentary bears a prominent foramen (the ‘anterior dentary foramen’ of  
821 Sereno [1991]) between the lateral and ventral prementary processes; three additional large  
822 foramina are exposed on the ventrolateral surface of the anterior dentary (Fig. 10D, fo). All of  
823 these large foramina (as well as a number of smaller openings on the ventromedial aspect of the  
824 anterior dentary) can be traced to a precipitate-filled Meckelian canal that becomes taller  
825 posteriorly. CT scans confirm a short, edentulous area of the dentary between its contact with the  
826 lateral prementary process and the first dentary tooth.

827 Unlike the flat ventrolateral surface of the maxilla, the external surface of the dentary is  
828 convex and the dentary teeth are inset (Fig. 10D, be; Thulborn, 1970; Sereno, 1991). Posteriorly,  
829 the dentary bifurcates into long tapering dorsal and ventral processes that overlap the surangular  
830 and angular, respectively (Fig. 10E, dp, vp; Thulborn, 1970; Norman et al., 2004). The dorsal  
831 process forms the anterior border of the coronoid eminence. A deep embayment between these  
832 processes forms the anterior half of the external mandibular fenestra (Fig. 10D, emf). The middle  
833 and posterior parts of the dentary are ‘C’-shaped in transverse section, laterally arched and have  
834 thickened dorsal and ventral margins. The ventromedial edge of the dentary meets the  
835 ventrolateral margin of the splenial in a rounded butt joint along most of its length (Fig. 10E,

836 a.sp). A long anterior process of the angular contacts the ventromedial aspect of dentary. The  
837 posterior third of the dorsomedial margin of the dentary contacts the lateral surface of the  
838 coronoid (Fig. 10E,a.co). Replacement foramina on the medial surface of the dentary cannot be  
839 resolved in CT scans though they are clearly visible in NHMUK PV R8501. In medial view, the  
840 dentary forms the anterior boundaries of the internal mandibular fossa.

#### 841 **Splénial**

842 The splénial is a transversely flattened sheet of bone forming much of the medial aspect of the  
843 lower jaw and encloses the Meckelian canal medially; it is preserved on both sides of the  
844 ‘palatal’ block of NHMUK PV RU B17 and can be visualized on the left side of NHMUK PV  
845 RU B23 (Fig. 2). In CT scans the dorsal and ventral margins are slightly thickened and the  
846 ventral margin is inturned to contact the ventral edge of the dentary in a simple butt joint along  
847 much of its length (Fig. 11A, a.d). In medial view, the anterior margin of the splénial is gently  
848 rounded; it approaches but does not reach the symphysis (Fig. 11B). Posteriorly, the splénial  
849 bifurcates into a short, rounded dorsal process and a long, tapering ventral process. A rounded  
850 notch between the processes contributes to the anterior half of the small internal mandibular  
851 fenestra (Fig. 11B, imf). The dorsal process of the splénial laterally contacts the anterodorsal  
852 process of the prearticular; the dorsal margin of this process meets the ventral edge of the  
853 coronoid in a simple butt joint (Fig. 11A, a.co). The longer ventral process of the splénial  
854 extensively underlaps the anterior process of the angular; posteriorly, it also underlaps the main  
855 body of the prearticular (Fig. 11A, a.pa).

#### 856 **Coronoid**

857 The coronoid is preserved on both sides of the ‘palatal’ block of NHMUK PV RU B17 and the  
858 left side of NHMUK PV RU B23 (Fig. 2). Anteriorly, it is a mediolaterally flattened strip of  
859 bone applied to the dorsal margin of the dentary (Fig. 11C, a.d) with its ventral margin resting on  
860 the dorsal edge of the splenial (Fig. 11D, a.sp); it extends along the posterior third of the dentary.  
861 It increases in height and width posteriorly, forming the highest point of the coronoid eminence  
862 and developing a lateral process that overlies the dorsal ramus of the surangular (Fig. 11C, a.sa).  
863 A ventral tab of the coronoid dorsally and medially overlaps the anterodorsal process of the  
864 prearticular (Fig. 11C, a.pa). There is a rounded ridge on the dorsal aspect of the coronoid that  
865 marks an area of muscle attachment.

#### 866 **Surangular**

867 Both surangulars are preserved in NHMUK PV RU B23, the ‘palatal’ block of NHMUK PV RU  
868 B17 (Fig. 2) and NHMUK PV R8501; additionally, an isolated but well-preserved right  
869 surangular is present in the ‘braincase’ block of NHMUK PV RUB 17 (Supporting Information,  
870 Fig. S4). The tapering dorsal ramus of the surangular underlies the dorsal process of the dentary  
871 (Fig. 11E, G, a.d). Additionally, it features a dorsomedial facet that underlaps the lateral process  
872 of the coronoid (Fig. 11E-G, a.co) and, on the left side of the ‘palatal’ block (NHMUK PV RU  
873 B17), the medial margin of the dorsal surangular ramus has a short contact with the posterodorsal  
874 process of the prearticular (Fig. 11F, a.pa). Posteriorly, the surangular increases in dorsoventral  
875 height; in transverse section, the element is laterally arched, with the dorsal margin strongly  
876 inturned and thickened. In external view, this results in a flattened area (Fig. 11E, G, m.ae) on  
877 the dorsolateral aspect of the surangular that is bounded dorsally and medially by a strong ridge  
878 (Fig. 11E, F, dr); this ridge is continuous anteriorly with the dorsal ridge of the coronoid.  
879 Together, this flat area and ridge mark areas of attachment for portions of the *M. adductor*

880 *mandibulae externus* group. The posterior half of the lateral surface of the surangular bears a  
881 longitudinal ridge (Fig. 11E, G, lr); the surangular foramen opens immediately below the  
882 posterior end of this ridge (Fig. 11E, sf). Below the lateral ridge, the ventrolateral aspect of the  
883 surangular is overlapped by the angular (Fig. 11E, a.an); the isolated surangular in the  
884 ‘braincase’ block clearly exhibits a large facet for this contact. The anteroventral edge of the  
885 surangular forms the posterodorsal margin of the external mandibular fenestra. The dorsal  
886 margin of the surangular is convex in lateral view (Sereno, 1991) and the bone decreases in  
887 height posteriorly. Immediately anterior to the jaw joint, there is a robust, medial extension  
888 (‘medial flange’ of Thulborn [1970]) of the surangular with a strongly concave anterior surface  
889 (Fig. 11F, G, mf). This flange forms the posterior wall of the adductor fossa, contacts the  
890 dorsoventral expansion of the prearticular, contacts the anterior tip of the articular, and forms the  
891 anterior margin of the jaw joint (Fig. 11F). Posteriorly, the surangular is deflected medially and  
892 becomes a transversely thin sheet of bone that laterally overlaps the articular, forming the  
893 retroarticular process (Fig. 11E–G, rp). A faint ridge divides the lateral surface of the  
894 retroarticular process into dorso- and ventrolateral surfaces. In medial view, the dorsal margin of  
895 the surangular is slightly inturned to form the border of the internal mandibular fenestra.

## 896 **Angular**

897 Both angulars are preserved in NHMUK PV RU B23, the ‘palatal’ block of NHMUK PV RU  
898 B17 (Fig. 2) and NHMUK PV R8501; a fragment of a right angular (attached to the isolated  
899 surangular) and an articulated left angular, articular and prearticular are also preserved in the  
900 ‘braincase’ block (Supporting Information, Fig. S4). A tapering anterior process extends into the  
901 Meckelian canal and overlaps the internal surfaces of both the dentary and splenial (Fig. 11H, I,  
902 a.d, a.sp). Posteriorly, the angular increases in height and becomes ‘L’-shaped in transverse

903 section. The vertical lamina externally overlaps the surangular while the horizontal lamina  
904 underlies the prearticular (Fig. 11I, a.pa). The anterodorsal margin of the angular forms the  
905 posteroventral margin of the external mandibular fenestra (Fig. 11H, emf). In medial view, the  
906 angular forms the ventrolateral border of the internal mandibular fenestra.

### 907 **Prearticular**

908 Both prearticulars are preserved in NHMUK PV RU B23 (although only the left element could  
909 be visualized in CT scans) and in the ‘palatal’ block of NHMUK PV RU B17 (Fig. 2); an  
910 additional left prearticular is preserved in the ‘braincase’ block (Supporting Information, Fig.  
911 S4). Although no single element is completely preserved, collectively the entire prearticular is  
912 represented. Posteriorly, the prearticular is a thin sheet of bone that extensively overlaps the  
913 medial surfaces of the articular and medial expansion of the surangular (Fig. 11J, a.ar, a.sa). This  
914 sheet of bone is strongly depressed laterally and marks a possible muscle attachment site. The  
915 prearticular decreases in height anteriorly and lies in a trough formed by the angular and splenial  
916 to form the floor of the mandibular adductor fossa (Fig. 11K). Anteriorly, the prearticular  
917 becomes transversely thin and dorsoventrally tall to form a robust anterodorsal process that  
918 articulates with the ventral tab of the coronoid (Fig. 11K, a.co), medial margin of the surangular  
919 (Fig. 11J, a.sa), and dorsal process of the splenial (Fig. 11K, a.sp), forming the anterodorsal  
920 margin of the internal mandibular fenestra.

### 921 **Articular**

922 Both articulars are preserved in NHMUK PV RU B23 and in the ‘palatal’ block of NHMUK PV  
923 RU B17 (Fig. 2); an additional left prearticular is preserved in the ‘braincase’ block (Supporting  
924 Information, Fig. S4). The articular is nearly as wide as it is long; its dorsoventral height is

925 approximately half its mediolateral width. In dorsal view the articular is widest across its centre  
926 and tapers anteriorly and posteriorly (Fig. 11M). It is held in a cup formed by the prearticular,  
927 angular and surangular (Fig. 11L, N). The tapering anterior process extends medial and ventral to  
928 the medial extension of the surangular (Fig. 11L); the posterior processes of the articular and  
929 surangular form the retroarticular process. The dorsal surface forms the jaw joint (Fig. 11M,  
930 cmj); there is no median ridge separating the surfaces for the lateral and medial condyles of the  
931 quadrate. A strong ridge traverses the articular and forms the anterior margin of the joint surface.

932

### 933 **Discussion**

934 Digital preparation has clarified the cranial anatomy of *Lesothosaurus diagnosticus* as well as  
935 revealing new features, allowing fresh comparisons with two additional ornithischian specimens  
936 from the Upper Elliot Formation of Lesotho (MNHN LES 17 and MNHN LES 18). Knoll  
937 (2002a, 2002b) compared these skulls with those of *Lesothosaurus diagnosticus* and concluded  
938 that, despite sharing many features with the syntypes, MNHN LES 17 could be distinguished  
939 from them. Consequently, he assigned this specimen to *Lesothosaurus* sp. Differences between  
940 MNHN LES 17 and the syntypes identified by Knoll (2002a) include:

- 941 1. Possession of a deep, hemispherical depression in the anteroventral corner of the  
942 antorbital fossa in MNHN LES 17 that was thought to be absent from the syntypes of *L.*  
943 *diagnosticus*. However, CT scans demonstrate that this depression is present on both  
944 sides of NHMUK PV RU B17 ('palatal' block) and NHMUK PV RU B23.
- 945 2. The antorbital fenestra was described as reniform in outline and relatively larger in  
946 MNHN LES 17 than in *L. diagnosticus*. However, crushing has caused anterior

- 947 displacement of the lacrimal on both sides of NHMUK PV RU B23, effectively closing  
948 this fenestra. Its true size and shape can be better appreciated in segmented scans of the  
949 left side of NHMUK PV RU B17 ('palatal' block), which shows that this structure was  
950 similar in both MNHN LES 17 and *L. diagnosticus*.
- 951 3. The frontal-nasal suture is flush and the prefrontal-nasal sutures are offset in MNHN LES  
952 17, whereas in NHMUK PV RU B23 the prefrontal-nasal contacts are flush while the  
953 frontals are ventrally offset. Observations of extant crocodylian and squamate skulls  
954 suggest that these sutural contacts on the external skull roof were likely flush in life and  
955 that the offsets in both specimens are due to post-mortem deformation (Knoll, 2002a).
- 956 4. The ventral branch of the postorbital is straighter in MNHN LES 17 than in NHMUK PV  
957 RU B23. However, both skulls have suffered some degree of deformation and both  
958 postorbitals are incomplete in MNHN LES 17.
- 959 5. The quadratojugal is separated from the quadrate condyle in MNHN LES 17, whereas it  
960 closely approaches the level of the jaw joint in *L. diagnosticus*. Re-examination of this  
961 area indicates that this region is heavily damaged in both MNHN LES 17 and most *L.*  
962 *diagnosticus* specimens, suggesting that this interpretation of this feature may be  
963 ambiguous.
- 964 6. Presence of an interparietal suture in MNHN LES 17. CT scans reveal a partially patent  
965 interparietal suture in NHMUK PV RU B23 that is not apparent in external view. An  
966 open suture appears to be present in NHMUK PV R11004.
- 967 7. Embayment between the parietals in MNHN LES 17 forms a "sharper V" than in *L.*  
968 *diagnosticus* (Knoll, 2002a:239), as illustrated by Sereno (1991). Dorsal views of MNHN  
969 LES 17 and segmented CT data of NHMUK PV RU B23 do not support this claim.

- 970 8. The paroccipital processes of MNHN LES 17 are horizontally-directed, compared to  
971 those of NHMUK PV RU B23 as depicted by Sereno (1991). CT scans show that the  
972 paraoccipital processes of NHMUK PV RU B23 are more horizontally-directed than  
973 illustrated by Sereno (1991); this discrepancy was also noted by Knoll (2002b).  
974 Furthermore, the left paroccipital of NHMUK PV RU B23 is slightly *dorsally* inclined  
975 while the right is slightly *ventrally* inclined, suggesting deformation in this area.
- 976 9. The distal ends of the paraoccipital processes are more inflated in MNHN LES 17 than in  
977 NHMUK PV RU B23. CT scans reveal an expansion of the distal ends of the processes in  
978 the syntype.
- 979 10. Basispterygoid processes are less laterally-directed in MNHN LES 17 than figured by  
980 Sereno (1991). This does not appear to be the case when comparing segmented  
981 basisphenoids from the syntypes and MNHN LES 17 in posterior views.
- 982 11. The coronoid eminence is higher in MNHN LES 17 than in *L. diagnosticus*. Segmented  
983 CT scans of the left side of NHMUK PV RU B23 and both sides of NHMUK RU B17  
984 ('palatal' block) demonstrate that the coronoid eminence of *L. diagnosticus* is higher than  
985 previously depicted due to the tall coronoid.
- 986 12. Shorter retroarticular process of MNHN LES 17 than NHMUK PV RU B23. The  
987 quadrate has been anteriorly displaced in the latter; segmentation of CT data from both  
988 syntypes demonstrates the retroarticular processes were of similar relative length as in  
989 MNHN LES 17.
- 990 13. Relatively large (and more angular) orbit and larger supratemporal fenestrae in MNHN  
991 LES 17 than in NHMUK PV RU B23. Both skulls are deformed and the postorbital bars

992 and upper temporal bars in MNHN LES 17 are incomplete, rendering these observations  
993 qualitative.

994 Additional similarities between the *L. diagnosticus* syntypes and MNHN LES 17 include:  
995 a midline groove between the nasals; lack of a sagittal crest in MNHN LES 17, as observed by  
996 Knoll (2002a); concave ventral surface of the basisphenoid; and an anteriorly arched caudal  
997 margin of the quadrate. Remaining differences between MNHN LES 17 and the syntypes  
998 include: the presence of a midline suture between the supraoccipitals in the former (almost  
999 certainly due to damage as the supraoccipital is an unpaired element in diapsids); and a gap  
1000 between the dorsal margin of the supraoccipital and parietal (though such a gap occurs frequently  
1001 in dinosaurs and may be ontogenetic). Based on the overwhelming similarities between MNHN  
1002 LES 17 and the syntype skulls, we assign MNHN LES 17 to *Lesothosaurus diagnosticus*.

1003 A larger partial skull (MNHN LES 18) was also assigned to *Lesothosaurus* sp. by Knoll  
1004 (2002b) based on numerous similarities between it and the syntypes. Knoll (2002b)  
1005 acknowledged that many of the differences between MNHN LES 18 and the syntypes (e.g.,  
1006 relative size and shape of the orbits and fenestrae, shape of the skull roof) could be attributed to  
1007 dorsoventral crushing of Paris specimen and possibly ontogenetic changes. Other differences  
1008 identified by Knoll (2002b) include:

- 1009 1. The right postorbital of MNHN LES 18 is not flush with the frontals as in NHMUK  
1010 PV RU B23. As noted above, it is likely that all sutural contacts on the external  
1011 surface of the skull roof were originally flush and such offsets can be attributed to  
1012 deformation.
- 1013 2. Direct contact between the head of the quadrate and paroccipital process in MNHN  
1014 LES 18. Given that the squamosal is missing and that the specimen is strongly

1015 crushed, the absence of the postquadratic process of the squamosal between the  
1016 quadrate and paroccipital process cannot be confirmed and seems unlikely.

1017 3. Lack of a median ridge on the supraoccipital of MNHN LES 18. As noted by Knoll  
1018 (2002b) the supraoccipital of this specimen is broken at the midline; thus, it is  
1019 impossible to determine whether or not a nuchal crest was present.

1020 4. Ventrally-directed paraoccipital processes in MNHN LES 18. See comments above as  
1021 well as noting strong dorsoventral crushing of MNHN LES 18.

1022 The only remaining differences between MNHN LES 18 and the syntype skulls include  
1023 the presence in the former of a weak sagittal crest and the much larger size of MNHN LES 18  
1024 compared to the syntypes. Knoll (2002b) acknowledged that specimens assigned to *L.*  
1025 *diagnosticus* by Sereno (1991) already exhibit a large range of body sizes. Furthermore, recent  
1026 work has demonstrated that the largest specimen of the early ornithischian *Heterodontosaurus* is  
1027 nearly three times the body length of the smallest known individual (Porro et al., 2011). Thus, it  
1028 is likely that MNHN LES 18 represents a larger individual of *Lesothosaurus diagnosticus*.  
1029 Alternatively, MNHN LES 18 could represent the skull of the larger ornithischian *Stormbergia*  
1030 (Butler, 2005), also present in the Upper Elliot Formation and presently known from postcranial  
1031 material only; however, the validity of this taxon is in doubt and there is evidence that it may, in  
1032 fact, represent an adult *Lesothosaurus* (Knoll, Padian & de Ricqles, 2009).

1033

## 1034 **Conclusions**

1035 Together with visual inspection of specimens, CT-scanning and 3D visualization was used to  
1036 produce a detailed anatomical description of the skull of *Lesothosaurus diagnosticus*, revealing

1037 new anatomical details such as sutural morphology and internal structures. Elements obscured by  
1038 matrix or other bones were described for the first time. This new description was used to assign  
1039 two specimens previously identified as *Lesothosaurus* sp. MNHN LES 17 (and possibly MNHN  
1040 LES 18) to *Lesothosaurus diagnosticus*.

1041

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1056

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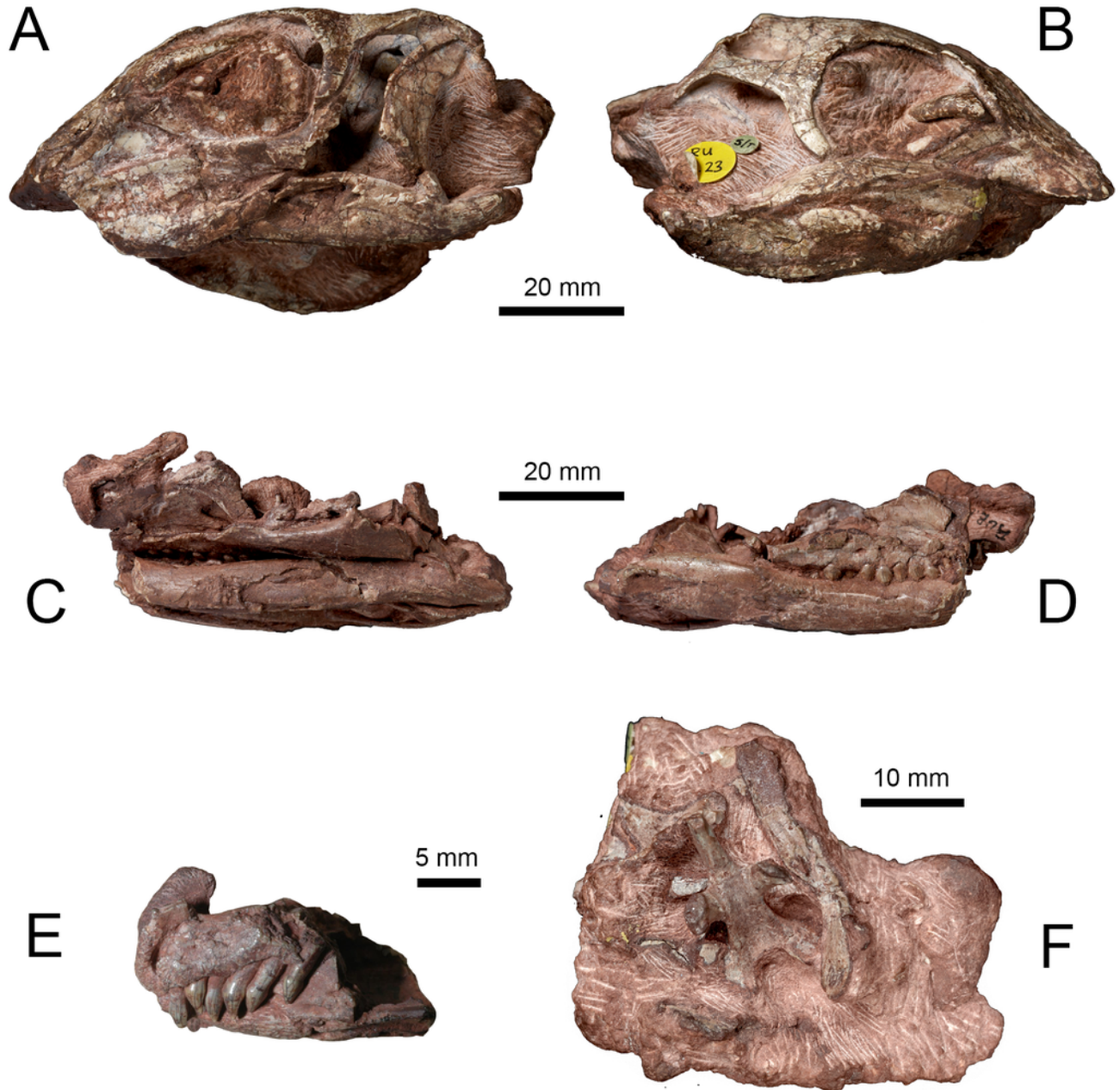
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# 1

Syntype specimens of *Lesothosaurus diagnosticus* examined CT-scanned in this study.

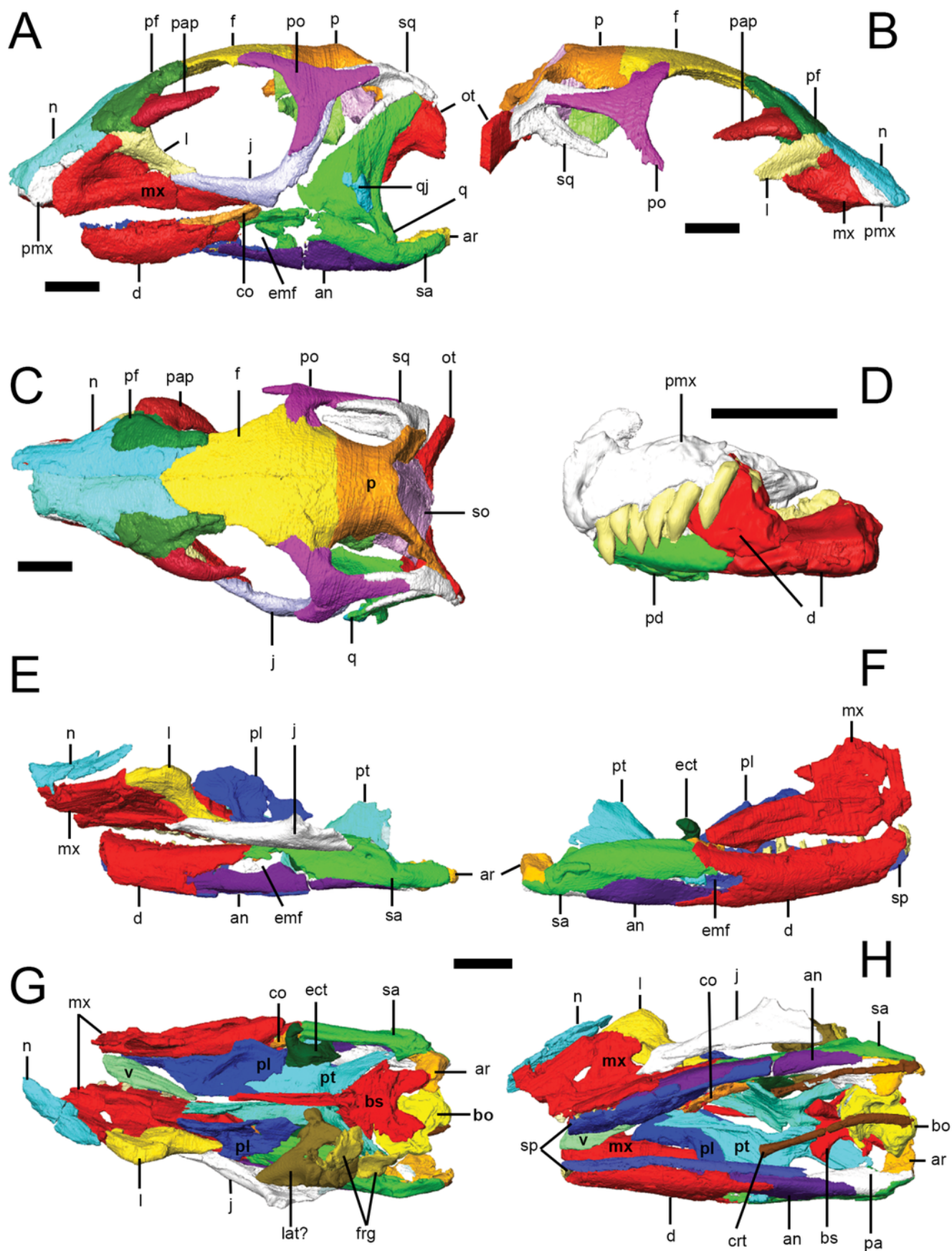
Left (A) and right (B) lateral views of NHMUK PV RU B23. Left (C) and right (D) lateral views of the 'palatal' block of NHMUK PV RU B17. Left lateral view of the 'snout' block of NHMUK PV RU B17 (E) and ventral view of the 'braincase' block of NHMUK PV RU B17 (F).



## 2

Surface models of *Lesothosaurus diagnosticus* specimens used in this study.

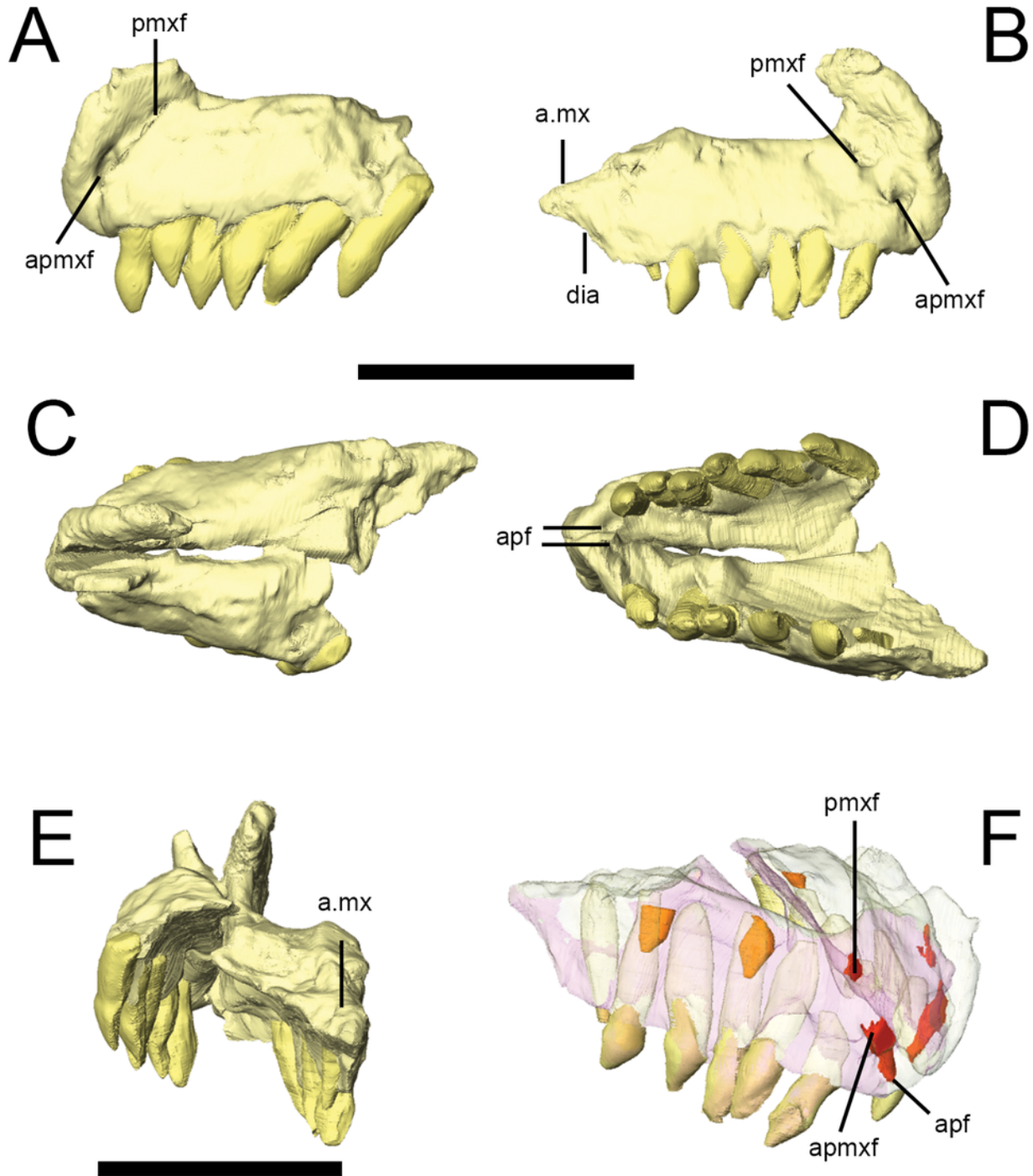
Left (A) and right (B) lateral views, and dorsal view (C) of segmented skull bones of NHMUK PV RU B23. Left lateral view of segmented bones in the 'snout' block of NHMUK PV RU B17 (D). Left (E) and right (F) lateral views, and dorsal (G) and ventral (H) views of the 'palatal' block of NHMUK PV RU B17. Individual bones are shown in various colours. Anatomical abbreviations: **an** , angular; **ar** , articular; **bo** , basioccipital; **bs** , basisphenoid; **co** , coronoid; **crt** , ceratohyals ; **d** , dentary; **ect** , ectopterygoid; **emf**, external mandibular fenestra; **f** , frontal; **frg**, braincase fragments; **j** , jugal; **l** , lacrimal; **lat**, laterosphenoid; **mx** , maxilla; **n** , nasal; **ot**, otoccipital; **p** , parietal; **pa** , prearticular; **pap**, palebral; **pd**, prementary; **pf** , prefrontal; **pl** , palatine; **pmx** , premaxilla; **po** , postorbital; **pt** , pterygoid; **q** , quadrate; **qj** , quadratojugal; **sa** , surangular; **so**, supraoccipital; **sp** , splenial; **sq** , squamosal; **v** , vomer. All scale bars equal 10 mm.



## 3

Premaxilla of *Lesothosaurus diagnosticus* (NHMUK PV RU B17).

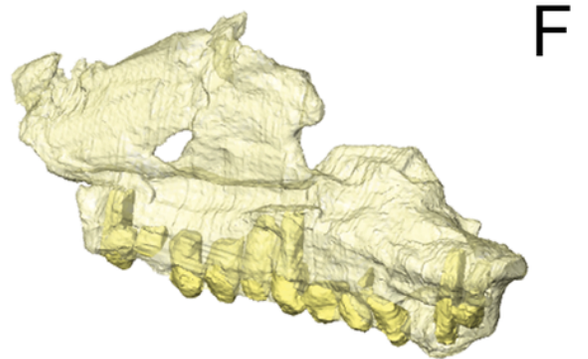
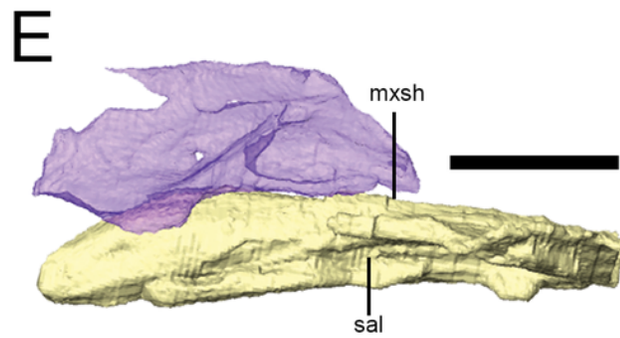
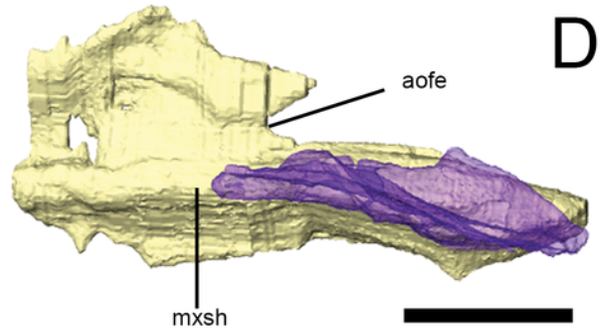
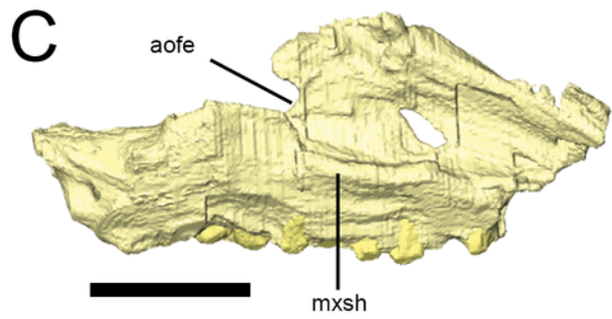
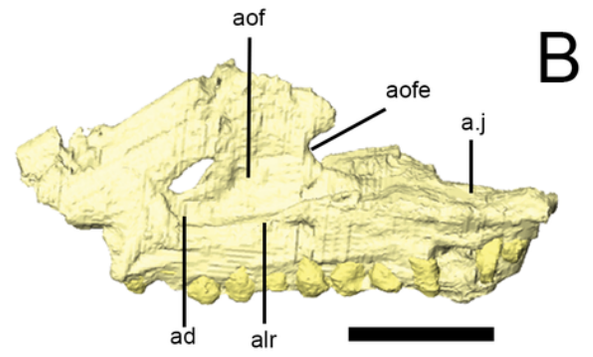
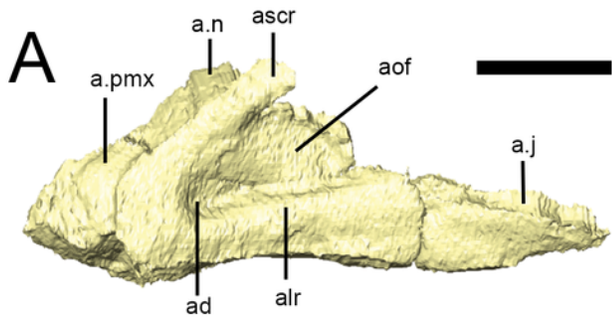
Left lateral (A) view of left premaxilla and right lateral (B) view of right premaxilla. Dorsal (C), ventral (D) and posterior (E) views of articulated left and right side elements. Right dorsal oblique view (F) of articulated elements with the bones transparent to visualize the tooth roots, replacement teeth (shown in orange) and canals linking various premaxillary foramina (red). Anatomical abbreviations: **a.mx**, articulation surface for maxilla; **apf**, anterior palatal foramen; **apmxf**, anterior premaxillary foramen; **dia**, diastema; **pmxf**, premaxillary foramen. All scale bars equal 10 mm. Scale bar cannot be provided for oblique view (F).



## 4

The maxilla of *Lesothosaurus diagnosticus*.

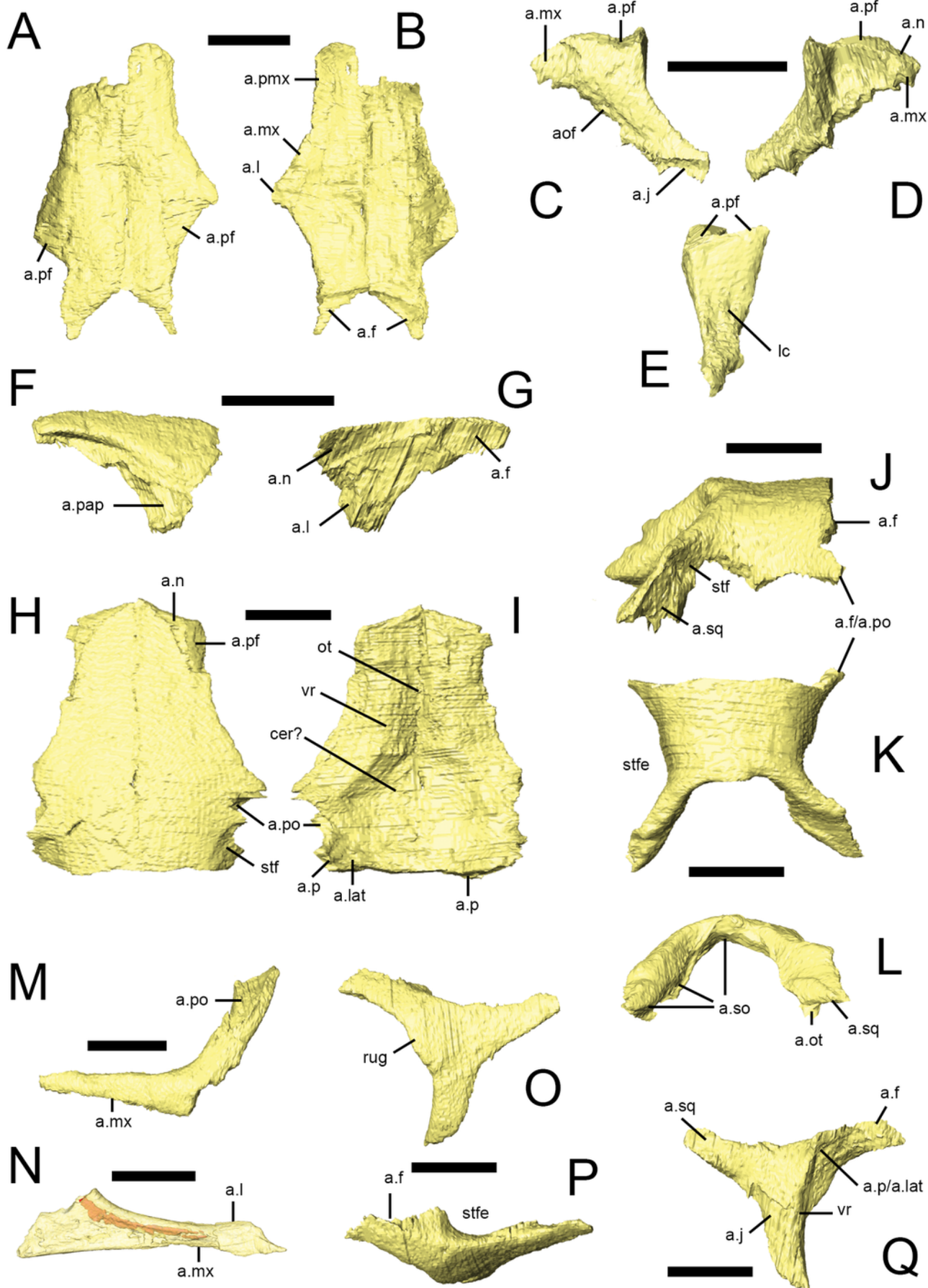
Surface renderings from NHMUK PV RU B23 (A) and NHMUK PV RU B17 (B-F). The left maxilla in left lateral view (A B); the left maxilla in medial view (C); the right maxilla in medial view (D); the right maxilla in dorsal view (E); and the left maxilla in posterolateral oblique view (F). The right palatine (blue semi-transparent) is shown in articulation with the right maxilla in D and E. Bones are rendered transparent to visualize tooth roots in F. Anatomical abbreviations: **ad**, deep depression in anteroventral corner of antorbital fossa; **a.j**, articulation surface for jugal; **alr**, alveolar ramus; **a.n**, articulation surface for nasal; **aof**, antorbital fossa; **aofe**, antorbital fenestra; **a.pmx**, articulation surface for premaxilla; **ascr**, ascending ramus; **mxsh**, medial maxillary shelf; **sal**, supralveolar lamina. All scale bars equal 10 mm. Scale bar cannot be provided for oblique view (F).



## 5

Facial bones and skull roof of *Lesothosaurus diagnosticus*.

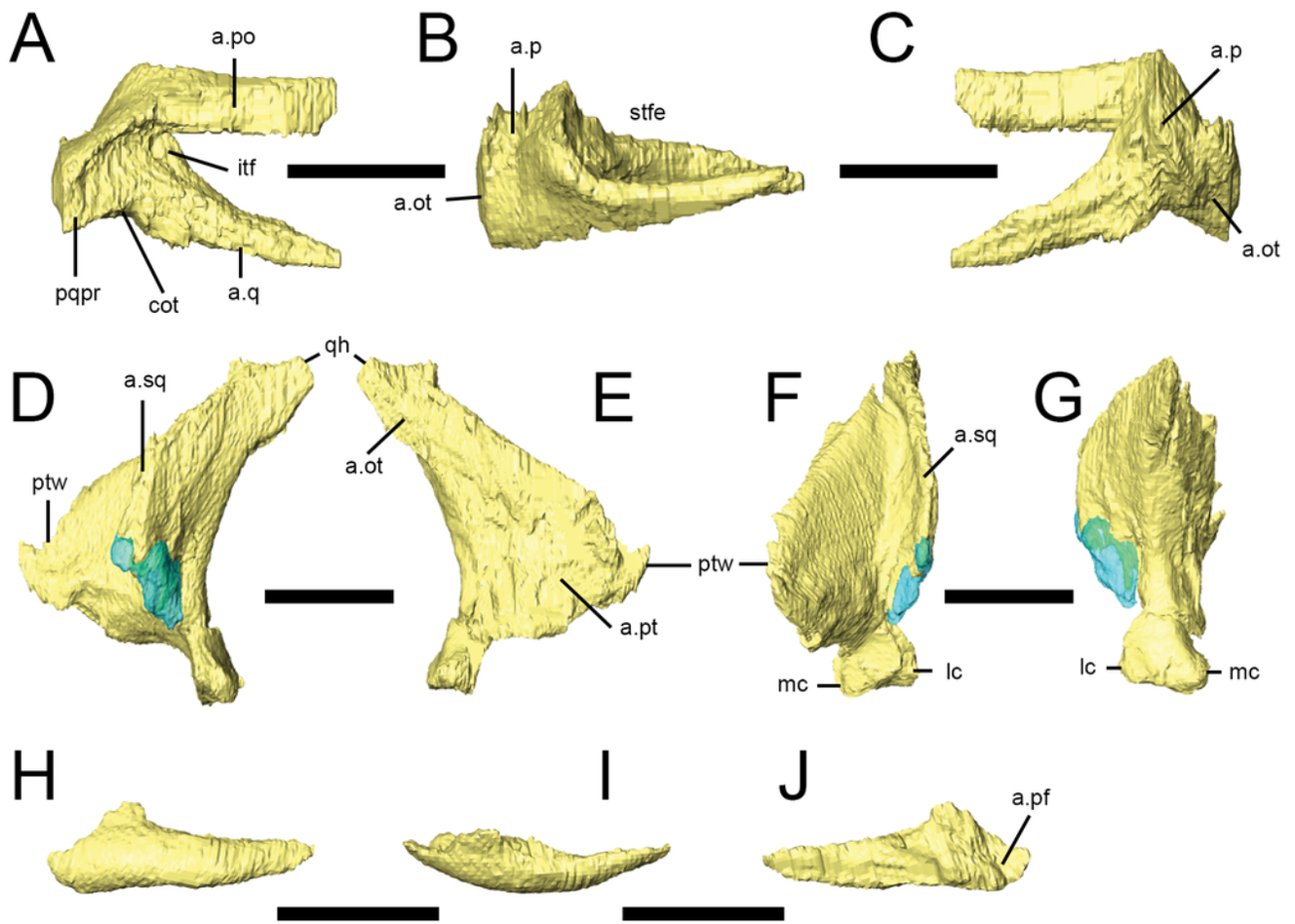
All surface renderings are from NHMUK PV RU B23 with the exception of N, taken from NHMUK PV RU B17. Paired nasals in dorsal (A) and ventral (B) views. The left lacrimal in lateral (C), medial (D) and posterior (E) views. The right prefrontal in lateral (F) and medial (G) views. Paired frontals in dorsal (H) and ventral (I) view. Parietal in right lateral (J), dorsal (K) and posterior (L) views. Left jugals in lateral (M) and medial (N) views. Left postorbital in lateral (O), dorsal (P) and medial (Q) views. Jugal (N) is shown transparent to visualize the internal cavity (indicated in red). Elements are shown at different scales. Anatomical abbreviations: **a.f**, articulation surface for frontal; **a.j**, articulation surface for jugal; **a.l**, articulation surface for lacrimal; **a.lat**, articulation surface for laterosphenoid; **a.mx**, articulation surface for maxilla; **a.n**, articulation surface for nasal; **a.ot**, articulation surface for paroccipital process, part of the otooccipital; **aof**, antorbital fossa; **a.p**, articulation surface for parietal; **a.pap**, articulation surface for palpebral; **a.pf**, articulation surface for prefrontal; **a.pmx**, articulation surface for premaxilla; **a.po**, articulation surface for postorbital; **a.so**, articulation surface for supraoccipital; **a.sq**, articulation surface for squamosal; **cer?**, possible impression for cerebral hemispheres; **lc**, lacrimal canal; **ot**, impression for olfactory tract; **rug**, rugosity on postorbital; **stf**, supratemporal fossa; **stfe**, supratemporal fenestra; **vr**, ventral ridge. All scale bars equal 10 mm.



## 6

Squamosal, quadrate and palpebral of *Lesothosaurus diagnosticus*.

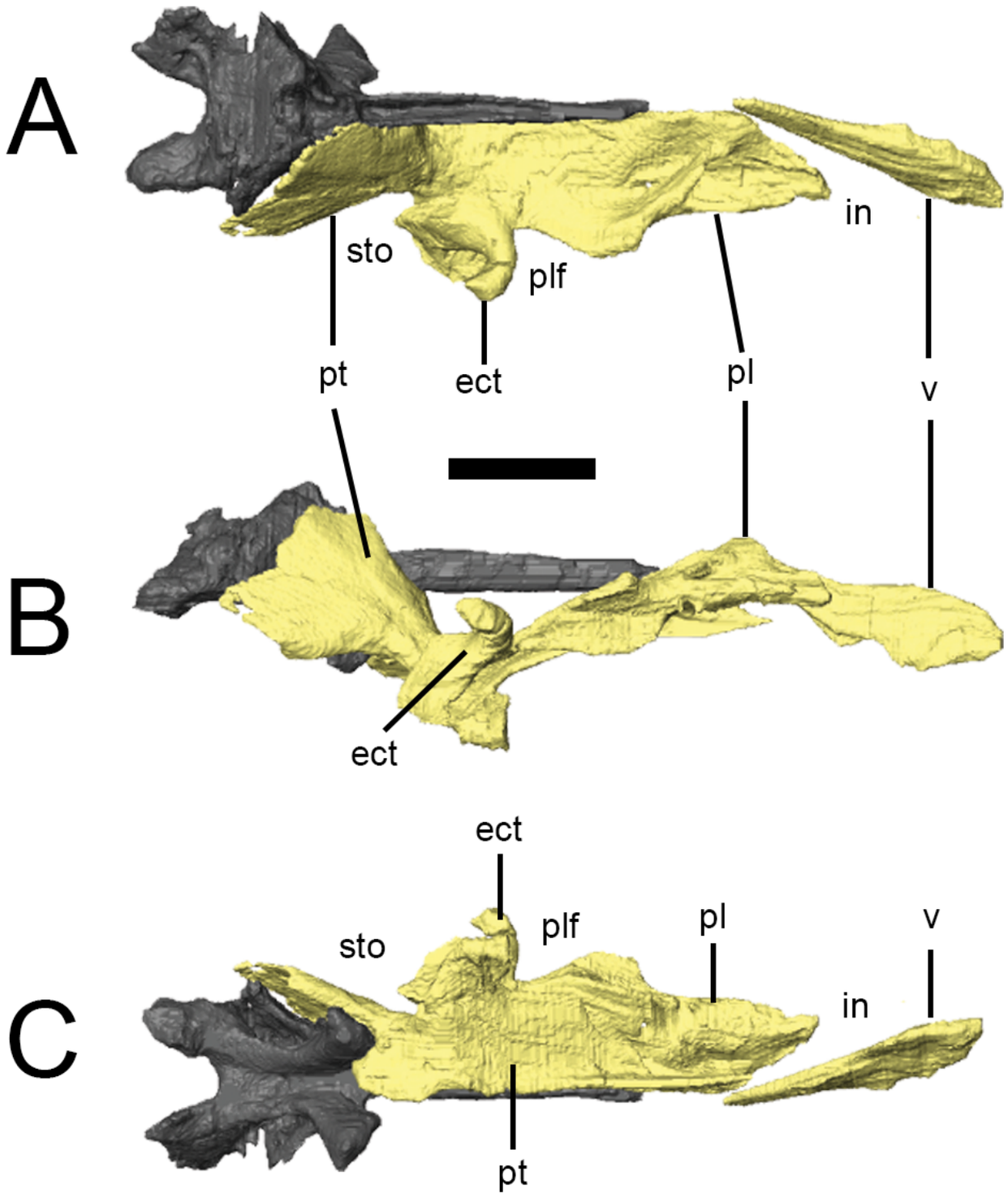
All surface renderings are from NHMUK PV RU B23. Right squamosal in lateral (A), dorsal (B) and medial (C) views. Left quadrate in lateral (D), medial (E), anterior (F) and posterior (G) views. The left palpebral in lateral (H), dorsal (I) and medial (J) views. Elements are shown at different scales. Anatomical abbreviations: **a.ot**, articulation surface for paraoccipital process, part of the otoccipital; **a.p**, articulation surface for parietal; **a.pf**, articulation surface for prefrontal; **a.po**, articulation surface for postorbital; **a.pt**, articulation surface for pterygoid; **a.q**, articulation surface for quadrate; **a.sq**, articulation surface for the squamosal; **cot**, cotylus for head of quadrate; **itf**, infratemporal fossa; **lc**, lateral condyle; **mc**, medial condyle; **pqpr**, postquadratic process of the squamosal; **ptw**, pterygoid wing of quadrate; **qh**, head of quadrate; **stfe**, supratemporal fenestra. All scale bars equal 10 mm.



## 7

Surface renderings of the right palate of *Lesothosaurus diagnosticus* from NHMUK PV RU B17 in dorsal (A), lateral (B) and ventral (C) views.

The anterior tip of the vomer has been displaced to the right. Anatomical abbreviations: **ect**, ectopterygoid; **in**, internal naris; **pl**, palatine; **plf**, palatal fenestra; **pt**, pterygoid; **sto**, subtemporal opening; **v**, vomer. All scale bars equal 10 mm. .

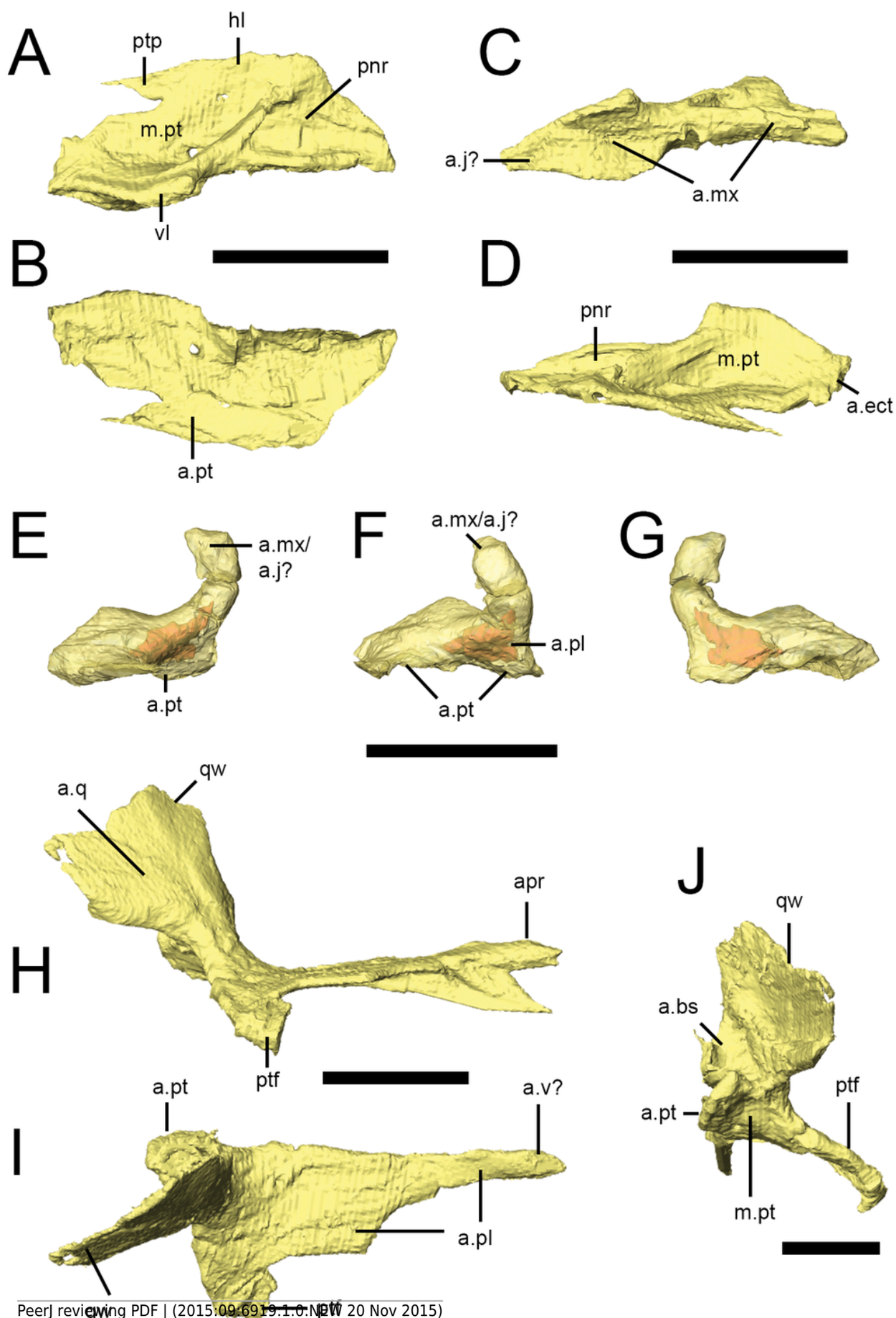


## 8

Palatal bones of *Lesothosaurus diagnosticus*.

All surface renderings are from NHMUK PV RU B17. Right palatine in dorsal (A), ventral (B), lateral (C) and medial (D) views. Right ectopterygoid in lateral (E), anterior (F) and medial (G) views (bone shown transparent to reveal hollow cavity within, shown in red). Right pterygoid in lateral (H), dorsal (I) and posterior (J) views. Elements are shown at different scales.

Anatomical abbreviations: **a.bs**, articulation surface for the basiptyergoid process; **a.ect**, articulation surface for ectopterygoid; **a.j?**, possible articulation surface for jugal; **a.mx**, articulation surface for maxilla; **a.pl**, articulation surface for palatine; **apr**, anterior process of the pterygoid; **a.pt**, articulation surface for pterygoid; **a.q**, articulation surface of the quadrate; **a.v?**, possible articulation surface for the vomer; **hl**, horizontal lamina of the palatine; **m.pt**, attachment site of *m. pterygoideus ventralis*; **pnr**, palatine pneumatic recess; **ptf**, pterygoid flange; **ptp**, pterygoid process of the palatine; **qw**, quadrate wing of the pterygoid; **vl**, vertical lamina of the palatine. All scale bars equal 10 mm.

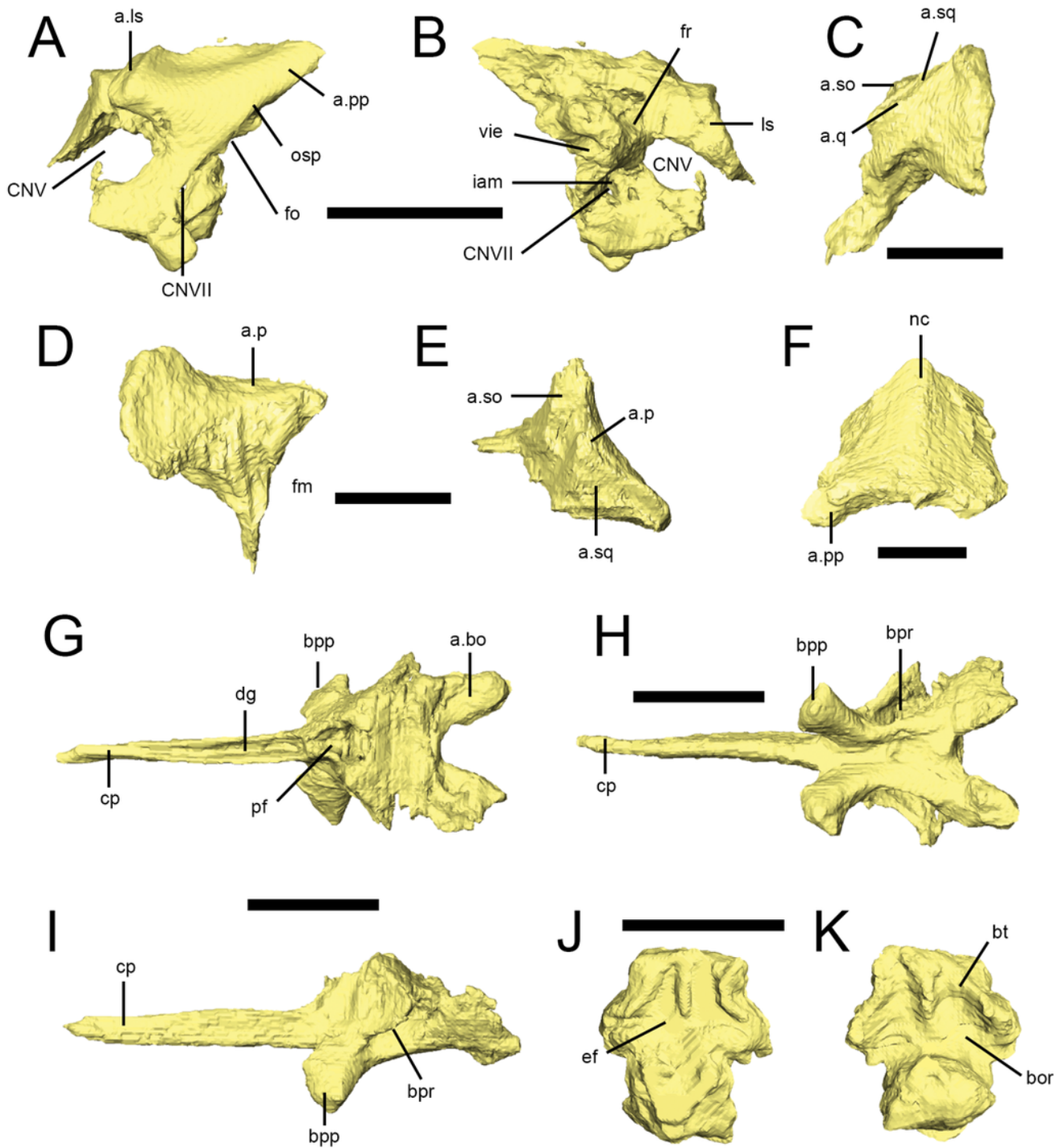


## 9

Braincase elements of *Lesothosaurus diagnosticus*.

Surface renderings from NHMUK PV RU B17 (A, B, G-K) and NHMUK PV RU B23 (C-F). Left prootic in lateral (A) and medial (B) views. Left paroccipital process (part of the otoccipital) in lateral (C), posterior (D) and dorsal (E) views. Supraoccipital in posterior (F) view.

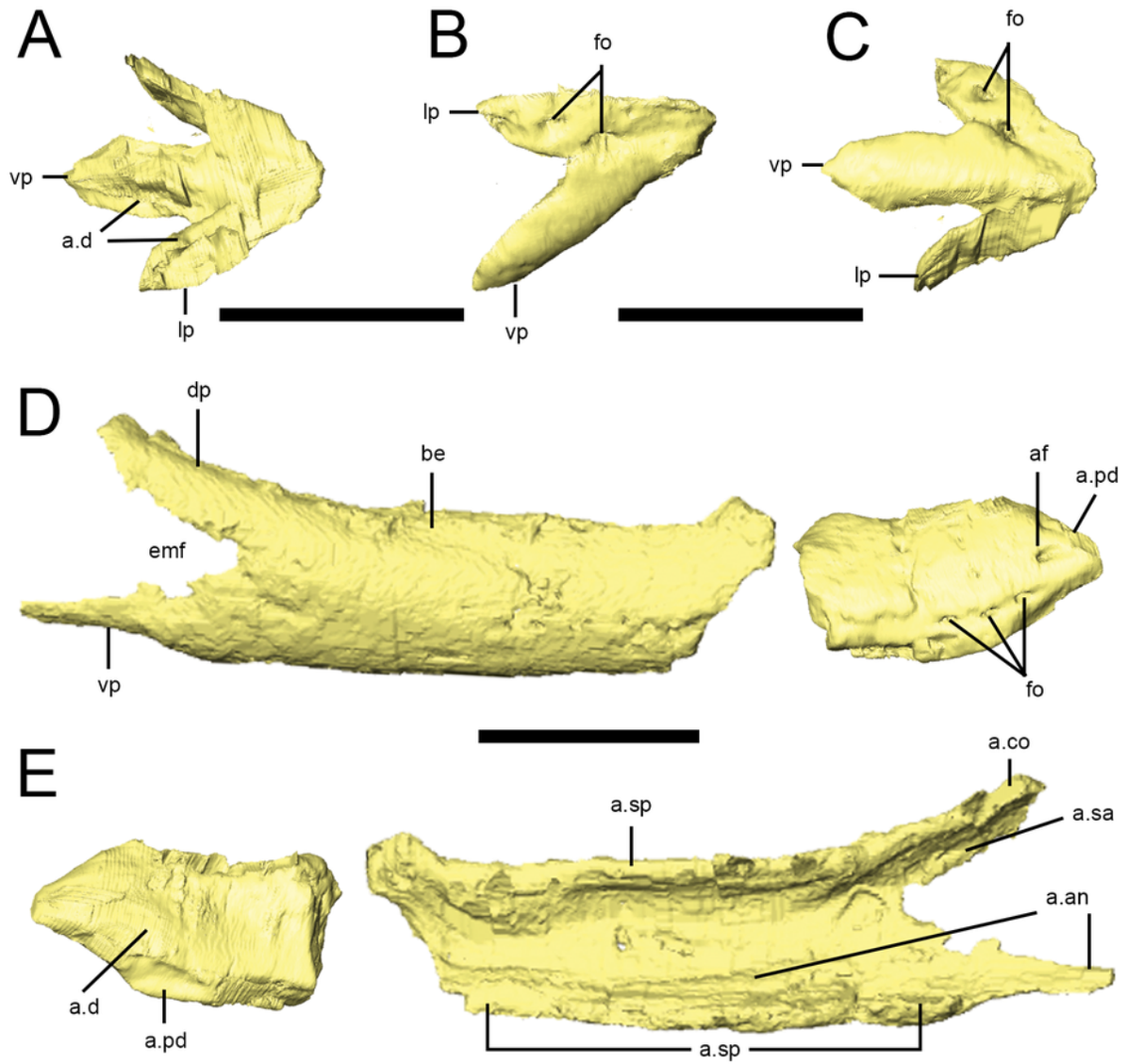
Basisphenoid in dorsal (G), ventral (H) and left lateral (I) views. Basioccipital in dorsal (J) and ventral (K) views. Elements are shown at different scales. Anatomical abbreviations: **a.bo**, articulation surface for the basioccipital; **a.ls**, articulation surface for the laterosphenoid; **a.p**, articulation surface for the lateral process of the parietal; **a.pp**, articulation surface for the paroccipital process (otoccipital); **a.so**, articulation surface for supraoccipital; **a.q**, articulation surface for quadrate; **a.sq**, articulation surface for the squamosal; **bor**, basioccipital recesses; **bpp**, basipterygoid process; **bpr**, basipterygoid recess; **bt**, basal tubera; **cp**, cultriform process of the parasphenoid; **CNV**, opening of the trigeminal nerve; **CNVII**, opening for facial nerve; **dg**, dorsal groove; **ef**, endocranial floor; **fm**, foramen magnum; **fo**, fenestra ovalis; **fr**, floccular recess; **iam**, internal acoustic meatus; **ls**, displaced fragment of laterosphenoid; **nc**, nuchal crest; **osp**, otosphenoidal crest; **pf**, pituitary process; **vie**, vestibule of inner ear. All scale bars equal 10 mm.



## 10

Predentary and dentary of *Lesothosaurus diagnosticus*.

Surface renderings are from NHMUK PV RU B17: the predentary and anterior right dentary is found in the 'snout' block while the posterior right dentary is preserved in the 'palatal' block. Predentary in dorsal (A), lateral (B) and ventral (C) views. Anterior and posterior portions of the right dentary in lateral (D) and medial (E) views. Elements are shown at different scales. Anatomical abbreviations: **a.an**, articulation surface for angular; **a.co**, articulation surface for coronoid; **a.d**, articulation surfaces for dentary; **af**, anterior foramen; **a.pd**, articulation surface for predentary; **a.sa**, articulation surface for surangular; **a.sp**, articulation surface for splenial; **be**, buccal emargination; **dp**, dorsal process of the dentary; **emf**, external mandibular fenestra; **fo**, foramina; **lp**, lateral process of the predentary; **vp**, ventral process. All scale bars equal 10 mm.



## 11

Lower jaw bones of *Lesothosaurus diagnosticus*.

Surface renderings are from the 'palatal' block of NHMUK PV RU B17 (A, B, E-K) and NHMUK PV RU B23 (C, D). Right splenial in lateral (A) and medial (B) views. Left coronoid in lateral (C) and medial (D) views; anteroventral margin of this element is damaged. Left surangular in lateral (E), medial (F) and dorsal (G) views; ventral margin of the anterior process is damaged. Left angular in lateral (H) and medial (I) views. Left prearticular in lateral (J) and medial (K) views; the anteroventral margin and posterior tip of this element are missing. Left articular in lateral (L), dorsal (M) and medial (N) views. Elements are shown at different scales. Anatomical abbreviations: **a.an**, articulation surface for angular; **a.ar**, articulation surface for articular; **a.co**, articulation surface for coronoid; **a.d**, articulation surface for dentary; **a.pa**, articulation surface for prearticular; **a.sa**, articulation surface for surangular; **a.sp**, articulation surface for splenial; **cmj**, craniomandibular joint; **dp**, dorsal process; **dr**, dorsal ridge; **emf**, external mandibular fenestra; **imf**, internal mandibular fenestra; **lr**, lateral ridge; **m.ae**, attachment site of *m. adductor externus superficialis* group; **maf**, mandibular adductor fossa; **mf**, medial flange; **rp**, retroarticular process; **sf**, surangular foramen; **vp**, ventral process. All scale bars equal 10 mm.

