



Three-dimensional mobility and muscle attachments in the pectoral limb of the Triassic cynodont *Massetognathus pascuali* (Romer, 1967)

Citation

Lai, Phil H., Andrew A. Biewener, and Stephanie E. Pierce. "Three-dimensional Mobility and Muscle Attachments in the Pectoral Limb of the Triassic Cynodont *Massetognathus Pascuali* (Romer, 1967)." *Journal of Anatomy* 232, no. 3 (2018): 383-406.

Permanent link

<http://nrs.harvard.edu/urn-3:HUL.InstRepos:41529913>

Terms of Use

This article was downloaded from Harvard University's DASH repository, and is made available under the terms and conditions applicable to Open Access Policy Articles, as set forth at <http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#OAP>

Share Your Story

The Harvard community has made this article openly available.
Please share how this access benefits you. [Submit a story](#).

[Accessibility](#)

1

1 Running heading: Cynodont pectoral limb musculoskeletal anatomy

2
3 **Title: Three-dimensional mobility and muscle attachments in the pectoral limb of**
4 **the Triassic cynodont *Massetognathus pascuali* (Romer, 1967)**

5
6 Phil H. Lai^{1,2*}, Andrew A. Biewener², Stephanie E. Pierce^{1*}

- 7 1. Museum of Comparative Zoology and Department of Organismic and Evolutionary Biology,
8 Harvard University, Cambridge, MA 02138, USA
9 2. Concord Field Station and Department of Organismic and Evolutionary Biology, Harvard
10 University, Bedford, MA 01730, USA

11
12 *Corresponding author: Phil H. Lai (phillai@g.harvard.edu) and Stephanie E. Pierce
13 (spierce@oeb.harvard.edu)

14
15 **ABSTRACT**

16 The musculoskeletal configuration of the mammalian pectoral limb has been heralded as a key
17 anatomical feature leading to the adaptive radiation of mammals, but limb function in the **non-**
18 **mammaliaform cynodont outgroup** remains unresolved. Conflicting reconstructions of abducted and
19 adducted posture are based on mutually-incompatible interpretations of ambiguous osteology. We
20 reconstruct the pectoral limb of the Triassic **non-mammaliaform** cynodont *Massetognathus pascuali* in
21 three dimensions, by combining skeletal morphology from micro-computed tomography with muscle
22 anatomy from an extended extant phylogenetic bracket. Conservative tests of maximum range of motion
23 suggest a degree of girdle mobility, as well as substantial freedom at the shoulder and the elbow joints.
24 The glenoid fossa supports a neutral pose in which the distal end of the humerus points 45°
25 posterolaterally from the body wall, intermediate between classically “sprawling” and “parasagittal” limb
26 postures. *M. pascuali* is reconstructed as having a near-mammalian complement of shoulder muscles,
27 including an incipient rotator cuff (m. subscapularis, m. infraspinatus, m. supraspinatus, and m. teres
28 minor). Based on close inspection of the morphology of the glenoid fossa, we hypothesize a posture-
29 driven scenario for the evolution of the therian ball-and-socket shoulder joint. The musculoskeletal
30 reconstruction presented here provides the anatomical scaffolding for more detailed examination of
31 locomotor evolution in the precursors to mammals.

32
33 **Keywords:** shoulder girdle; forelimb; synapsids; mammals; range of motion; musculoskeletal function;
34 postural evolution.

35
36 **INTRODUCTION**

37 Today's mammals **show** disparate **locomotor modes**, comprising cursorial, fossorial, aquatic, and even
38 volant forms (Hildebrand, 1989; Fischer et al. 2002; Vaughan et al. 2013). These varied lifestyles are
39 supported by modifications of the pectoral limb into anatomical structures as diverse as wings and
40 flippers. The evolution of the **therian-style shoulder girdle and forelimb**—mobile scapula, ball-and-socket
41 glenohumeral joint, “parasagittal” limb posture—has been suggested as a key innovation leading to the
42 adaptive radiation of the clade (Polly, 2007). Morphological diversification of this anatomical module
43 began early on in mammalian evolution (Ji et al. 2006; **Meng et al. 2017; Luo et al. 2017**), predating the
44 emergence of the crown group (Luo, 2007). Accordingly, interpreting morphological and functional
45 transformation of the pectoral limb in the sister group to mammals is key to understanding their
46 remarkable success.

47
48 The **non-mammaliaform** cynodonts (**henceforth referred to simply as "cynodonts"**) offer a glimpse **at an**
49 **ancestral condition from which mammalian locomotion evolved**. The osteology of the cynodont pectoral
50 girdle and forelimb is well known from the fossil record, and does not appear to have been particularly
51 disparate; in a series of papers, Jenkins (1970a, 1971a) synthesized a number of descriptions and
52 posited that most cynodonts shared a common appendicular morphology, and presumably similar
53 locomotor behaviors. In contrast to the hip articulation, where a socket-like acetabulum clearly
54 circumscribed range of motion (Jenkins, 1971a), the cynodont gleno-humeral joint possessed the
55 relatively unconstrained, hemisellar architecture on which late Permian archosaurs, lepidosaurs, and

56 synapsids converged (Jenkins, 1993)—the typical mammalian ball-and-socket articulation did not appear
57 until the Jurassic theriomorphs (Ji et al. 1999; Luo, 2015). Multiple reconstructions of the cynodont
58 pectoral limb have been advanced, drawing on skeletal morphology (Watson, 1917; Kühne, 1956;
59 Bonaparte, 1963; Jenkins, 1970b, 1971a; Kemp, 1980a, 1980b; Oliveira & Schultz, 2016) as well as
60 muscle anatomy as inferred from osteology and homology to extant taxa (Gregory & Camp, 1918; Romer,
61 1922).

62
63 Two competing hypotheses of cynodont posture and locomotion have emerged, with discrepancies
64 centered on divergent interpretations of shoulder mobility, and the position occupied along the classic
65 sprawling-to-upright continuum of tetrapod posture (Gatesy, 1991; Reilly & Elias, 1998). A cornerstone of
66 the upright, or adducted postural view, is Jenkins' work on *Massetognathus pascuali*, a traversodontid
67 cynodont from the Triassic Chañares Formation of Argentina (Romer, 1967; Jenkins, 1970b). As a
68 member of Cynognathia, the sister group to the anatomically-comparable probainognathians that gave
69 rise to mammals (Ruta et al. 2013), *M. pascuali* is a reasonable exemplar of early Mesozoic cynodont
70 organization (Liu & Olsen, 2010), and represents one of the last major transformational steps towards the
71 mammalian condition. Based on the postcranial skeleton of *M. pascuali*, Jenkins advanced a two-
72 dimensional reconstruction in a crouched, adducted pose with posteriorly-directed elbows, reminiscent of
73 a small, short-limbed therian (Jenkins, 1970b). Working from cynognathian (*Cynognathus*) and
74 probainognathian (*Trucidocynodon*) material, Watson (1917) and Oliveira and Schultz (2016) arrived at
75 similarly therian-like interpretations of posture and locomotion across eucynodonts.

76
77 On the other hand, Kemp's (1980a, 1980b) reconstructions of the basal Late Permian cynodont
78 *Procynosuchus* and the Middle Triassic traversodont *Luangwa* depicted the sprawling, abducted posture
79 thought to be plesiomorphic for amniotes. The humerus is held perpendicular to the animal's sagittal
80 plane, and the main stride component is furnished by protraction and retraction of the humerus around a
81 dorsoventral axis. Kemp posited that the basic structure and function of the forelimb remained unchanged
82 between the Permian and Triassic cynodonts, and that the limb and girdle transformations leading to
83 adducted posture were restricted to later, more crownward taxa. The close phylogenetic relationship
84 between *Massetognathus* and *Luangwa* (Liu & Abdala, 2014) means we currently have a reconstruction
85 with abducted posture in one traversodont, and adducted posture in another. The equivocal osteology of
86 the cynodont pectoral girdle has so far precluded consensus on forelimb function, hindering a deeper
87 understanding of locomotor evolution in this important clade.

88
89 Here we revisit cynodont forelimb morphology and function using modern computational methods to add
90 a third dimension (3D) to this classic problem. Using digital models of fossil material derived from micro-
91 computed tomography (μ CT), we interactively assess articular function at the shoulder and elbow joints
92 (e.g. Pierce et al. 2012; Nyakatura et al. 2015). Further, we reconstruct the origins and insertions of the
93 shoulder musculature using an updated, extended extant phylogenetic bracket, and map them onto the
94 3D pectoral limb skeleton of *Massetognathus pascuali*. The result is a robust, three-dimensional
95 reconstruction that will form the basis of future biomechanical analyses using musculoskeletal modeling
96 techniques (e.g. Hutchinson et al. 2005; Bates & Schachner, 2012; Hutchinson et al. 2015) to probe the
97 link between skeletal motion and muscle function.

98 MATERIALS AND METHODS

99 μ CT scanning and segmentation

100 A nodule containing the nearly-complete, articulated remains of *Massetognathus pascuali* (MCZVP 3691)
101 from the Museum of Comparative Zoology (Harvard University, USA) was scanned using a Nikon
102 Metrology (X-Tek) HMXST225 MicroCT unit located at Harvard University's Center for Nanoscale
103 Systems. Scanning parameters were 175kV 46 μ A, with a 0.01mm copper filter and a final voxel size of
104 127.22 μ m. The μ CT data were imported into Mimics v18 (Materialise NV, Leuven, Belgium) for
105 segmentation. Pectoral girdle (interclavicle, clavicles, scapulocoracoids) and forelimb (humeri, radii,
106 ulnae) skeletal elements were identified and assigned individual masks, from which high-resolution 3D
107 meshes were computed and exported for smoothing and repair (Fig. 1).

108 Bone repair

109
110

111 The pectoral limb exhibited no overall distortion, but certain bones had suffered taphonomic **fracture**,
112 necessitating repairs to their digital models. Fragments were manually aligned, using contralateral
113 elements as reference. In the case of the humerus, radius, and clavicle, damage to opposing ends of the
114 left- and right-side elements was remedied by taking the intact end of one element, mirroring it, and
115 grafting it onto the opposing end of its counterpart. Small gaps were filled using the “Wrap” and “Surface
116 Reconstruction” algorithms in 3-Matic v10 (Materialise NV, Leuven, Belgium), while larger breaks were
117 bridged and smoothed over using digital sculpting tools in Autodesk Mudbox (Autodesk, Inc., San Rafael,
118 CA, USA). As we were unable to locate all of the fragments of the interclavicle, we modeled missing
119 segments based on the preserved cranial portion, other **eucynodont** interclavicles in the MCZ collections,
120 and Jenkins’ (1970b, 1971a) description of the same element in other cynodonts.

121
122 The repaired bone meshes were smoothed and re-wrapped in MeshLab (ISTI-CNR, Pisa, Italy), to
123 eliminate artifacts introduced in scanning and segmentation, while preserving potentially informative
124 surface texture. To reduce noise, we performed a Poisson surface reconstruction (Hoppe, 2008), which
125 takes the vertex coordinates of the original mesh and outputs an optimized, re-triangulated mesh. We
126 then applied a single Laplacian smoothing step (Field, 1988) to correct any remaining polygonal
127 irregularities, and exported the finished meshes in Wavefront .OBJ format (Wavefront Technologies,
128 Santa Barbara, CA, USA) for examination and assembly.

129 130 **Re-articulation and rigging**

131 Using 3DS Max (Autodesk, Inc., San Rafael, CA, USA), centers of rotation for the acromio-clavicular,
132 humero-radial, and humero-ulnar joints were determined by fitting spherical primitives to their opposing
133 articular surfaces and then superimposing the spheres’ centroids in 3D space (Fig. 2). A cylinder was
134 used to model the planar clavo-interclavicular joint, and an ellipsoid of aspect ratio 24:13 was used to
135 model the gleno-humeral joint. Thus articulated, the pectoral girdle and forelimb skeleton were organized
136 as a kinematic hierarchy, wherein each bone was subordinated to the reference frame of its proximal
137 neighbor, and inherited all rotations and translations applied to the latter.

138
139 A local joint coordinate system (JCS) (Grood & Suntay, 1983) was defined for each articulation, with joint
140 axes positioned to reflect anatomically-informative rotations. Axes were oriented following the XYZ
141 rotation order convention, with Z capturing the axis of greatest expected mobility and X the least (Brainerd
142 et al. 2010).

143 144 **Range of motion testing**

145 Limb joint range of motion has been shown to be sensitive to assumptions of intra-joint spacing (Arnold et
146 al. 2014; Nyakatura et al. 2015; Pierce et al. 2012). To circumscribe this, we established an articular
147 cartilage thickness of 0.25 mm based on relationships for mammalian articular cartilage (Simon, 1970).
148 Assuming equal cartilage thickness on opposing articular surfaces, we modeled every joint with a total
149 joint space of 0.50 mm. This value is supported by the *in situ* spacing between limb elements in the fossil
150 specimen, and yielded good agreement between the curvature of opposing articular surfaces.

151
152 Osteological limits to joint motion were assessed by rotating the distal element of each joint until it
153 collided with another bony surface, and repeating in the opposite direction to give total osteological range
154 of motion around each axis. The clavo-interclavicular, acromio-clavicular, humero-radial, and humero-
155 ulnar joints were tested only in rotation, but the unusual morphology of the gleno-humeral joint has been
156 suggested to support coordinated translation and rotation, in the form of sliding (Jenkins, 1971a) or rolling
157 (Kemp, 1980b) kinematics. Accordingly, we opted to compare purely rotational range of motion at this
158 joint against combined translation-rotation. To do so, we imported the *Massetognathus pascuali* model
159 into SIMM (Software for Interactive Musculoskeletal Modeling: Delp & Loan, 1995) and defined kinematic
160 functions linking rotations around X, Y and Z (Fig. 2) with translations along those same axes. The
161 functions were tuned to maintain a constant 0.5mm offset between the humeral head and the glenoid
162 fossa. Due to the unconstrained morphology of the glenoid fossa, we found that extremes of gleno-
163 humeral rotation could result in disarticulation of the joint well before a collisional limit was reached. To
164 establish reasonable physiological limits for this joint, we defined a secondary constraint criterion of 50%
165 humeral head contact with the glenoid fossa. Results of range of motion testing are given in Table 1.

166

167 A neutral reference pose was defined, allowing joint rotations to be repeatedly measured and compared
168 (Brainerd et al. 2010; Gatesy et al. 2010). The gleno-humeral joint was rotated to the center of its
169 measured ranges of motion (Fig. 3). Doing so placed the approximate centroids of the glenoid fossa and
170 the humeral head in apposition, as has been hypothesized to best approximate *in vivo* utilization of joint
171 surfaces during locomotion (Fischer, 1994). The humero-radial and humero-ulnar joints were then rotated
172 to orient the antebrachium normal to the "substrate". Notably, MCZVP 3801 preserves the articulated left-
173 side pectoral limb of a different *M. pascuali* individual with the shoulder and elbow similarly flexed, lending
174 support to the viability of this pose.
175

176 Shoulder muscle reconstruction

177 Prior reconstructions of cynodont musculature have used monotremes (Gregory & Camp, 1918) as well
178 as saurians and therians (Romer, 1922; Jenkins, 1971b) as bookends to examine conservation and
179 transformation in muscle anatomy through mammal evolution. Following this practice, we defined an
180 extended Extant Phylogenetic Bracket (Witmer, 1995) encompassing a range of amniotes, and using
181 salamanders as an outgroup (see Table 2 for the full list of taxa). Due to the paucity of well-delineated
182 attachment sites on the radius and ulna, past studies have largely focused on muscles crossing the
183 shoulder joint. In the interest of parsimony and repeatability, we follow suit in limiting our set of muscles
184 under consideration to only those that span the gleno-humeral joint. This metric excludes the extrinsic
185 scapular muscles (such as m. serratus anterior, m. trapezius, and mm. rhomboidei) that originate on the
186 axial skeleton and insert on the shoulder girdle, as well as the various flexors, extensors, pronators, and
187 supinators that originate on the distal humerus and actuate the distal forelimb. M. latissimus dorsi and
188 mm. pectorales have insertions on the humerus, and were included in the analysis despite their origins on
189 the axial skeleton, due to their presumed first-order action on the brachium.
190

191 Muscle origins and insertions were taken from the primary literature (see Table 2 for list of references),
192 and reconstructed as likely present in *Massetognathus pascuali* (level I inference *sensu* Witmer, 1995) or
193 possibly present (level II inference). None of the muscles under consideration were determined to be
194 likely absent (level III inference). Bryant and Seymour's (1990) study of carnivorans was used as a
195 reference for muscle attachment type (direct/fleshy, aponeurotic, or tendinous). Osteological correlates
196 such as rugose ridges and fossae were identified on MCZVP 3691, and compared against other prepared
197 *M. pascuali* postcranial material (e.g. other individuals in MCZVP 3691, MCZVP 3801, 4000, 4001, 4018,
198 and 4037) to confirm observations; all were found to be consistently present across individuals. Absence
199 of clear osteological correlates was not considered grounds for elimination, as muscles inserting directly
200 into periosteum may not leave a mineralized scar (Bryant & Seymour, 1990). In the absence of bony
201 scars, we placed attachments on homologous regions of the bone (Holliday, 2009). Muscles were
202 homologized with reference to Abdala and Diogo (2010) and Diogo et al. (2009). Reconstructed muscle
203 attachment areas were then digitally painted onto the 3D bone meshes using Autodesk Mudbox, for
204 visualization and comparison.
205

206 RESULTS

207 An interactive 3D PDF of the left-side pectoral limb of *Massetognathus pascuali* is included for
208 visualization purposes as Supplementary Figure 1, and shows reconstructed muscle attachments in
209 detail.

210 Neutral reference pose

211 Digitally reassembling the pectoral limb into a neutral reference pose (Fig. 3) places the diaphysis of the
212 humerus at approximately 45° to the animal's body wall, with the proximal and distal articular surfaces in
213 roughly the same horizontal plane. The anteriorly-oriented scapulocoracoid, posterolateral placement of
214 the glenoid fossa, and caudally-pointing elbows of this pose are broadly consistent with Watson's (1917),
215 Jenkins' (1971a), and Kemp's (1980a) reconstructions of traversodont pectoral girdles/limbs.
216

217 Jenkins (1971a) contested Watson's (1917) reconstruction of the cynodont pectoral girdle, arguing that
218 angling the scapulocoracoids medially and cranially would force the humerus into a mechanically
219 untenable posture while compromising the weight-bearing suspensory function of the extrinsic scapular
220 musculature. Jenkins' reconstruction orients the scapulocoracoids more vertically and tilts them outward
221 from the midline; as a result, the glenoid fossae are directed more ventrally than in Watson's
222 reconstruction. Based on our 3D reconstruction, the geometry of the clavicles and interclavicle constrain

223 the scapulocoracoids such that the glenoid fossae must be oriented posterolaterally and slightly ventrally,
224 as in Jenkins' reconstruction.

225

226 **Joint range of motion**

227 As reconstructed here, the pectoral limb of *Massetognathus pascuali* has one possible degree of
228 rotational freedom (DOF) at the clavo-interclavicular joint, up to three at the acromio-clavicular joint, and
229 three each at the gleno-humeral, humero-radial, and humero-ulnar joints, for a total of 13 DOF. Table 1
230 presents measured ranges of motion at each joint.

231

232 The clavo-interclavicular articulation lacks the extensive, rigid transverse overlap seen in monotremes
233 and tritylodonts (Sun & Li, 1985; Sues & Jenkins, 2006; Luo, 2015), and there are no known instances of
234 tosis between these elements in *M. pascuali*. The planar geometry of the articular facets **constrains**
235 mobility at this joint to medial and lateral rotation of the clavicle around an axis normal to the interclavicle
236 (Figure 2) (Kemp, 1980b). Clavicular elevation-depression and long-axis rotation are improbable, as such
237 movements would require compressive deformation of a substantial thickness of soft tissue around the
238 joint.

239

240 Some degree of mobility at the acromio-clavicular articulation has been hypothesized (Kemp, 1980b), but
241 the extent of this is difficult to assess in our 3D model of *M. pascuali* as the acromion is small and **likely**
242 incomplete. Although the minimally-projecting acromion observed in MCZVP 3691 is consistent with
243 Jenkins' (1971a) and Kemp's (1980a) descriptions of eucynodont scapulocoracoids, MCZVP 4018
244 **contains a right-side *M. pascuali* scapulocoracoid with a larger acromion projecting 3mm cranially in the**
245 **same plane. Liu (2007) also documented several traversodont scapulocoracoids with more prominent**
246 **acromions, including a juvenile *M. pascuali*. We opted not to reconstruct a larger acromion onto MCZVP**
247 **3691; simply extending the existing acromial surface in a cranial direction is unlikely to significantly impact**
248 **collisional range of motion at this simple, convex-concave articulation, and reconstruction would require**
249 **extensive further CT scanning of *M. pascuali* material.** The distal tip of the clavicle likely presents a
250 shallowly, concave articular surface that is somewhat congruent with the medial surface of the
251 presumptive acromion. We cautiously propose that some sliding or translational motion may have been
252 possible between the clavicle and the acromion, allowing pitch, roll, and yaw rotations around the
253 acromio-clavicular joint, subject to soft-tissue constraints.

254

255 The glenoid fossa is dorsoventrally concave and anteroposteriorly convex, resembling one half of a sellar
256 joint. The humeral head has an approximately ellipsoidal morphology, with the major axis running
257 between the greater and lesser tubercles. Rotation appears to be somewhat restricted in the absence of
258 translation, totalling 40° in abduction-adduction, 30° in retraction-protraction, and 40° in pronation-
259 supination. By contrast, we found greatly increased mobility around all three rotational axes when
260 allowing for translation, suggesting—in line with Jenkins (1971a) and Kemp (1980b)—that the gleno-
261 humeral articulation is a six degree of freedom joint.

262

263 Moving distally to the elbow, we measured comparable amounts of total flexion-extension for the humero-
264 radial and humero-ulnar joints (135° for the former, 140° for the latter). Both the radius and the ulna are
265 capable of some amount of abduction-adduction (80° for the radius, 60° for the ulna), and long-axis
266 rotation of each of these bones is unrestricted with reference to the neutral pose. In life, interosseous
267 ligaments and pronator muscles running between the radius and ulna would likely have constrained
268 independent movement of these two bones, while permitting coordinated pronation and supination within
269 the maximum osteological ranges established here.

270

271 **Muscle reconstruction**

272 The full set of muscles and taxa considered is given in Table 2, along with primary literature references. A
273 total of 12 muscles were reconstructed for *Massetognathus pascuali*. Individual muscles and their
274 attachments are discussed in the text below.

275

276 **M. latissimus dorsi (Fig. 4)**

277 *M. latissimus dorsi* appears to be plesiomorphic for tetrapods (Romer, 1924), and is present across the
278 phylogenetic bracket (Table 2). This muscle originates aponeurotically from the dorsal and thoracodorsal

279 fascia, and sometimes takes multiple costal origins as well (Diogo et al., 2009). It inserts on the
280 proximodorsal surface of the humeral deltopectoral crest in all cases except for monotremes, where the
281 insertion follows the deltopectoral crest distally to terminate on the entepicondyle (Gambaryan et al.
282 2015). *M. latissimus dorsi* in cynodonts is a level I inference, given its presence on both sides of the
283 bracket. This muscle inserts adjacent to *m. teres major* on a linear area running parallel to the long axis of
284 the humerus (Fig. 4). Based on *Cynognathus*, Jenkins (1971a) situated the insertion of the cynodont *m.*
285 *latissimus dorsi* on a ridge running obliquely across the dorsal surface of the humerus. He did not identify
286 a corresponding ridge in *Thrinaxodon* or *Massetognathus*, and we are unable to locate this feature on
287 close examination of the latter. Instead, we follow Watson (1917), Romer (1922) and Kemp (1980a) in
288 reconstructing a linear insertion for *m. latissimus dorsi* running proximodistally along the dorsomedial
289 surface of the humerus, terminating on a tuberosity just proximal to the midpoint of the diaphysis.

291 **M. pectoralis (Figs. 4, 5)**

292 *M. pectoralis* is present in all extant tetrapods (Table 2). While non-mammals may have multiple *m.*
293 *pectoralis* heads (Jenkins & Goslow, 1983), the mammalian pectoralis complex comprises a cranial,
294 superficial pectoralis major and a caudal, deeper pectoralis minor (Jenkins & Weijjs, 1979). In most
295 mammals, these muscles insert together along the length of the humeral deltopectoral crest, but in some
296 taxa (including humans) the pectoralis minor inserts separately on the coracoid process of the scapula.
297 This is certainly a derived condition, and for the purposes of this study we will consider *m. pectoralis* as a
298 single functional unit, without major or minor divisions. Originating from the ventral midline of all tetrapods
299 on (where present) the interclavicle, sternal series, and sometimes the medial ends of the costal
300 cartilages, *m. pectoralis* inserts on the posteroventral surface or apex of the deltopectoral crest in all
301 cases. *Massetognathus pascuali* possesses a prominent deltopectoral crest running slightly more than
302 halfway along the humeral diaphysis. A “cruciate” interclavicle is plesiomorphic for synapsids (Jenkins,
303 1971a), and was considered by Romer (1940) to give origin to a pectoralis complex via the paired fossae
304 on the posterior ramus. Relative to its length, the interclavicle of *M. pascuali* is considerably broader
305 mediolaterally than that of pelycosaur, with a well-marked posterior ramus and ridge that may represent
306 an expanded attachment for *m. pectoralis* (Romer, 1940; Jenkins, 1971a). We reconstruct *m. pectoralis*
307 as a level I inference, originating all over the lateral surfaces of the posterior process, across the fossae
308 on the posterior ramus, and possibly also on the caudally-facing surfaces of the lateral ridge (Fig. 5). *M.*
309 *pectoralis* inserts as an aponeurosis on the posteroventral surface of the deltopectoral crest of the
310 humerus, spanning its proximodistal length (Fig. 4).

312 **M. deltoideus scapularis and m. deltoideus clavicularis (Figs. 4, 6, 7)**

313 *M. deltoideus* is present in all tetrapods as a scapular division (*m. deltoideus scapularis*) and a clavicular
314 division (*m. deltoideus clavicularis*), with mammals gaining an additional acromial division (*m. deltoideus*
315 *acromialis*) (Table 2, homology follows Diogo et al. 2009). The acromion appears to be variably-
316 developed in *Massetognathus pascuali* and other cynodonts, and offers no obvious site for muscle
317 attachment (Jenkins, 1971a; Kemp, 1978); accordingly, we have reconstructed *M. pascuali* with only the
318 scapular and clavicular heads common to all tetrapods.

319
320 Like monotremes, the cynodont scapula has a strongly reflected cranial border (rcb, Fig. 7), which is
321 probably homologous to the therian scapular spine (Romer, 1922; Jenkins, 1971a; Kemp, 1980a,
322 Gambaryan et al. 2015). The caudally-facing surface of this border is the likely site of origin for *m.*
323 *deltoideus scapularis*, in agreement with Gregory and Camp (1918), Romer (1922), and Jenkins (1971a),
324 but *contra* Kemp (1980a, 1980b), who attributed a broader origin to *m. deltoideus*, covering much of the
325 lateral surface of the scapula in addition to the reflected border. In living lepidosaurs and crocodylians, *m.*
326 *deltoideus scapularis* takes origin from the cranial or craniodorsal portions of the lateral scapular surface.
327 The presence of a pronounced reflected cranial border would functionally divide an *m. deltoideus*
328 *scapularis* spanning the entire lateral surface of the scapula into a posteriorly-facing portion and a
329 laterally-facing portion, with uncertain consequences for its resultant line of action. In the absence of any
330 instances across the phylogenetic bracket of such a functional division, we consider an *m. deltoideus*
331 *scapularis* origin restricted to the reflected cranial border more biomechanically plausible. As in all other
332 tetrapods (Table 2), the cynodont *m. deltoideus scapularis* inserts in conjunction with *m. deltoideus*
333 *clavicularis*, on the anterodorsal surface of the humeral deltopectoral crest (Fig. 4).

334

335 M. deltoideus clavicularis (*sensu* Diogo et al. 2009) originates on the ventral half of the cranial border of
336 the scapula surrounding the acromion in crocodylians (Meers, 2003), on the interclavicle extending onto
337 the clavicle in lepidosaurs (Romer, 1922; Jenkins & Goslow, 1983) and monotremes (Howell, 1937a;
338 Gambaryan et al. 2015), and solely along the length of the clavicle in therians (Parsons, 1896; Jenkins &
339 Weijs, 1979; Stein, 1981). The cranial edge of the clavicle in *Massetognathus pascuali* forms a distinct
340 ridge, which extends into a protruding, anteriorly-directed flange along the distal half of the bone (cf. Fig.
341 6). This is the likely origin of m. deltoideus clavicularis, though it may also extend ventrally beyond the
342 clavicle to the area of the scapula surrounding the acromion (not reconstructed). M. deltoideus
343 clavicularis inserts with m. deltoideus scapularis, on the anterodorsal surface of the humeral deltopectoral
344 crest (Fig. 4).

346 **M. supraspinatus and m. infraspinatus (Figs. 4, 7)**

347 There is some question as to whether m. supraspinatus and m. infraspinatus were present as separate,
348 differentiated muscles in cynodonts, though ontogeny shows that both are likely derivatives of the m.
349 supracoracoideus present in non-mammals (Cheng, 1955; Romer, 1956). Some workers regard the
350 majority of the lateral surface of the cynodont scapula as an infraspinous fossa for the origin of m.
351 infraspinatus, with m. supraspinatus occupying the area at the cranialateral base of the scapula and the
352 caudodorsal half of the procoracoid, where the ancestral supraspinatus attached (Gregory & Camp, 1918;
353 Romer, 1922; Jenkins, 1971a). On the other hand, Kemp (1980a, 1980b) considered the "infraspinous
354 fossa" an attachment site for m. deltoideus scapularis and m. teres minor. Under this hypothesis, the
355 ventral procoracoid area attributed to m. supraspinatus by others would instead be occupied by an
356 undifferentiated m. supracoracoideus. This latter interpretation more closely resembles the monotreme
357 condition, wherein m. supraspinatus and m. infraspinatus are located at the cranial base of the scapula
358 and on the procoracoid (Howell, 1937a; Gambaryan et al. 2015). However, despite their more stem-ward
359 position in the mammal phylogeny, the suitability of extant monotremes as cynodont analogues may be
360 compromised by modifications for a fossorial or aquatic lifestyle (Howell, 1937b; Jenkins, 1971a; Kemp,
361 1980b). The probable cranial migration of m. deltoideus scapularis in cynodonts (see above) likely
362 corresponded to a dorsal expansion of m. supracoracoideus along the large, laterally-facing surface of
363 the scapula, paralleling its origin from the lateral scapular base and procoracoid of lepidosaurs and
364 archosaurs (Table 2).

366 Kemp (1980a) further argued that m. supraspinatus preceded m. infraspinatus in differentiating from m.
367 supracoracoideus, via dorsal migration onto the anteriorly-facing surface of the reflected cranial scapular
368 border. The ventral border of the clavicle is closely juxtaposed with the dorsal border of the procoracoid in
369 the neutrally-posed pectoral girdle of *Massetognathus pascuali* (Fig. 3), leaving little space in between to
370 accommodate such a muscle or its tendon, which in any case would have had to wrap around the
371 acromion to reach Kemp's proposed insertion on the greater tubercle of the humerus. The anteriorly-
372 facing surface of the reflected cranial border was more likely occupied by various extrinsic muscles
373 inserting on the scapula, such as m. trapezius and m. levator scapulae, both of which are likely
374 plesiomorphic for amniotes (Jouffroy et al. 1971; Diogo et al. 2009).

376 We follow Romer (1922), Gregory and Camp (1918), and Jenkins (1971a) in reconstructing m.
377 infraspinatus on most of the lateral surface of the scapula, caudal to the origin of m. deltoideus scapularis
378 (Fig. 7). This muscle has a tendinous insertion on a rugosity on the distal portion of the greater tubercle,
379 between the insertion of m. supraspinatus and the proximalmost margin of the humeral deltopectoral
380 crest (Fig. 4). This position is intermediate between the insertion of m. supracoracoideus in *Varanus*
381 (Jenkins & Goslow, 1983) on the proximal border of the deltopectoral crest, and the insertion of mm.
382 spinati in mammals on the greater tubercle proper (Leach, 1977; Jenkins & Weijs, 1979; Warburton et al.
383 2014). An m. supracoracoideus/m. supraspinatus was likely present in *Massetognathus pascuali*,
384 originating on the rugose area around the cranial scapular base and the adjoining procoracoid (Fig. 7).
385 The presumptive m. supraspinatus inserts by a tendon on a rugosity on the proximal half of the humeral
386 greater tubercle, just proximal to the insertion of m. infraspinatus (Fig. 4).

388 **M. teres minor (Figs. 4, 7)**

389 According to Diogo et al. (2009), the origins of the mammalian m. teres minor are murky, with workers
390 proposing homology with either m. scapulothoracalis anterior based on development (Romer, 1944;

391 Cheng, 1955), or m. deltoideus scapularis based on the co-existence of m. teres minor with m.
392 scapulohumeralis anterior in extant monotremes (Howell, 1937a; Jouffroy et al. 1971). Presuming
393 homology with m. scapulohumeralis anterior, Romer (1922) reconstructed the origin of m. teres minor in
394 cynodonts at the caudolateral base of the scapula, just cranial to the origin of m. triceps brachii. Gregory
395 and Camp (1918) and Jenkins (1971a) also placed m. teres minor at the caudolateral base of the
396 scapula, but ventral to the origin of m. triceps brachii rather than adjacent to it. Following the deltoid origin
397 hypothesis for m. teres minor, Kemp (1980b) favored an origin high up near the vertebral border on the
398 lateral surface of the scapula, recalling this muscle's location in monotremes (Howell, 1937a; Gambaryan,
399 2015). *Massetognathus pascuali* presents no clear area of origin for m. teres minor on the dorsolateral
400 surface of the scapula, but does possess a scar at the base of the lateral scapular surface (Fig. 7). We
401 therefore agree with Gregory and Camp (1918) and Romer (1922) that this was the likely site of origin for
402 m. teres minor. It is worth noting that this location is compatible with both hypotheses of origin, via either
403 ventral differentiation of the deltoid complex or direct homology with m. scapulohumeralis anterior. In
404 mammals, m. teres minor inserts via a tendon on the greater tubercle of the humerus (Howell, 1937a;
405 Leach, 1977; Jenkins & Weijs 1979; Stein, 1981; Gambaryan et al. 2015). In extant lepidosaurs, m.
406 scapulohumeralis anterior inserts on the dorsal surface of the humerus near the insertions of m.
407 deltoideus and m. latissimus dorsi (Romer, 1922; Miner, 1925; Holmes, 1977; Jenkins & Goslow, 1983).
408 We follow Jenkins (1971a) and Kemp (1980b) in placing the insertion of m. teres minor on a short ridge
409 extending parallel to the long axis of the humerus from the junction of the deltopectoral crest and the
410 greater tubercle (Fig. 4).

411 412 **M. subcoracoscapularis/subscapularis (Figs. 4, 7)**

413 All tetrapods possess either m. subcoracoscapularis or m. subscapularis in the form of a muscle
414 originating over much of the medial surface of the scapulocoracoid or scapula (Table 2). In lepidosaurs
415 and monotremes, this muscle has an additional head originating on the medial surfaces of the coracoid
416 and the procoracoid (Jenkins, 1971a; Jenkins & Goslow, 1983; Gambaryan et al. 2015). Regardless of
417 origin, m. subcoracoscapularis/subscapularis always inserts via a tendon in the vicinity of the humeral
418 lesser tubercle (Table 2). M. subcoracoscapularis is found on the medial side of the scapulocoracoid in all
419 cases except for monotremes, where the subscapular fossa has migrated around the caudal border of the
420 scapula to face posterolaterally, exposing the subscapularis in lateral view. The cynodont scapula exhibits
421 no such torsion, and the presumptive fossa for m. subcoracoscapularis faces primarily medially, as is the
422 case for all other tetrapods. We follow Gregory and Camp (1918), Jenkins (1971a) and Kemp (1980a) in
423 reconstructing a two-headed subcoracoscapularis originating on the medial surfaces of the scapula and
424 coracoid (Fig. 7), and inserting via a tendon on a rugose area at the apex of the lesser tubercle on the
425 humerus (Fig. 4).

426 427 **M. teres major (Figs. 4, 7)**

428 M. teres major (Table 2) is present in crocodylians (Meers, 2003) and all mammals (Howell, 1937a;
429 Leach, 1977; George, 1977; Jenkins & Weijs, 1979; Stein, 1981, 1986; Abdala & Diogo, 2010;
430 Gambaryan et al. 2015), but is absent in lepidosaurs (Romer, 1944; Diogo et al. 2009; Abdala & Diogo,
431 2010). Abdala and Diogo (2010) considered m. teres major a derivative of m. subcoracoscapularis,
432 homologous across crocodylians and mammals, and secondarily lost in lepidosaurs and bird-line
433 archosaurs. In therians and crocodylians, the origin of m. teres major runs dorsoventrally along the
434 axillary border of the scapula from the caudal angle (Fig. 7), or on the lateral surface of the scapula
435 adjacent to the axillary border (Howell, 1937a; Leach, 1977; George, 1977; Taylor, 1978; Jenkins &
436 Weijs, 1979; Stein, 1981; Meers, 2003; Abdala & Diogo, 2010; Harvey & Warburton, 2010; Gambaryan et
437 al. 2015). In monotremes, the origin of m. teres major runs craniocaudally along the lateral surface of the
438 scapula, terminating at the caudal angle. Depending on whether the crocodylian m. teres major is
439 homologous to that of mammals, m. teres major is either a level I or a level II inference. In certain
440 cynodonts, such as *Cynognathus*, part of the axillary border of the scapula is reflected laterally into a
441 ridge dividing the caudalmost part of the lateral scapular surface from the infraspinous fossa (Romer,
442 1922; Jenkins, 1971a; Liu, 2007). Some workers have interpreted this clearly demarcated fossa as the
443 origin of m. teres major (Gregory & Camp, 1918; Jenkins, 1971a). In other cynodonts, including
444 cynognathians such as *Luangwa* (Kemp, 1980b) and *Massetognathus pascuali*, this caudal fossa is
445 absent, although the scapula does have a somewhat thickened area on the laterally-reflected axillary
446 border (Fig. 7). We follow Kemp (1980b) in reconstructing m. teres major as a straplike muscle originating

447 as a narrow strip along this thickened dorsal region of the scapula's axillary border (Fig. 7). *M. teres major*
 448 likely inserted along a ridge running proximodistally along the dorsal surface of the humeral diaphysis
 449 (Fig. 4), parallel to the insertion of *m. latissimus dorsi* but slightly proximal (Jenkins, 1971a).

450

451 **M. coracobrachialis (Figs. 4, 7)**

452 *M. coracobrachialis* (Table 2) is present in all extant tetrapods as a muscle running from the posterior part
 453 of the lateral coracoid surface—with a second head at the cranio-lateral base of the scapula in
 454 crocodylians (Meers, 2003)—to an insertion on the ventromedial surface of the humerus, extending onto
 455 the posteromedial surface of the humeral deltopectoral crest (Miner, 1925; Howell, 1937a; Holmes, 1977;
 456 Meers, 2003; Walthall & Ashley-Ross, 2006; Diogo et al. 2009; Abdala & Diogo, 2010; Gambaryan et al.
 457 2015). While the presence of *m. coracobrachialis* as a whole is conserved among tetrapods, its
 458 subdivisions and by extension its distal attachments are not. This muscle exists as longus, medius, and
 459 brevis divisions in most amphibians (but not all, see Walthall & Ashley-Ross, 2006 and Abdala & Diogo,
 460 2010). Lepidosauurs have lost the medius division (Jenkins & Goslow, 1983; Abdala & Diogo, 2010);
 461 crocodylians have lost all but the brevis division (Meers, 2003); monotremes seem to have lost either the
 462 brevis division (Diogo et al. 2009) or the medius (Gambaryan et al. 2015); while therians lose the longus
 463 and sometimes also the brevis (Leach, 1977; George, 1977; Diogo et al. 2009; Harvey & Warburton,
 464 2010). While the homology of the various *m. coracobrachialis* divisions among extant tetrapods is beyond
 465 the scope of this paper, it seems safe to say that cynodonts probably had some form of *m.*
 466 *coracobrachialis*. Here we have reconstructed two origins and two insertions, representing possible
 467 brevis/medius and longus divisions.

468

469 Romer (1922) and Jenkins (1971a) considered the cynodont *m. coracobrachialis* to originate just caudal
 470 to *m. biceps brachii* within a fossa on the lateral surface of the coracoid, while Gregory and Camp (1918)
 471 assigned that fossa to *m. biceps brachii* and placed *m. coracobrachialis* on the caudal tip of the coracoid
 472 instead. *Massetognathus pascuali* has a well-marked fossa on the coracoid immediately cranial and
 473 inferior to the glenoid, and a smaller, shallower scar on the procoracoid immediately cranial to the
 474 procoracoid-coracoid suture (Fig. 7). It seems likely that *m. coracobrachialis* originated on the former and
 475 *m. biceps brachii* on the latter, echoing the arrangement of these muscles in extant *Iguana* (Romer, 1922)
 476 and *Alligator* (Meers, 2003). A second *m. coracobrachialis* head may have originated on the lateral
 477 surface of the coracoid, caudal and inferior to the glenoid (Fig. 7). There is no rugosity associated with *m.*
 478 *coracobrachialis medius* on the humerus of *M. pascuali*, but insertion can reasonably be assumed to have
 479 occurred on the large fossa on the ventromedial surface, with a possible insertion for *m. coracobrachialis*
 480 longus occurring further distal on a ridge near the entepicondyle (Fig. 4), consistent with past
 481 reconstructions (Watson, 1917; Gregory & Camp, 1918; Romer, 1922; Miner, 1925).

482

483 **M. biceps brachii (Figs. 7, 8)**

484 *M. biceps brachii* is likely an amniote synapomorphy, derived from *m. coracobrachialis* (Abdala & Diogo,
 485 2010). The two heads of this muscle generally originate on adjacent areas of the lateral coracoid (Romer,
 486 1922; Miner, 1925; Howell, 1937a; Jenkins, 1971a; Holmes, 1977), although *m. biceps brachii brevis* is
 487 usually absent in *Alligator* (Meers, 2003), and the origin of *m. biceps brachii brevis* is shifted caudally to
 488 the tip of the coracoid in *Ornithorhynchus*, similar to the condition seen in *Tupaia* (George, 1977) and
 489 *Homo* (Netter et al. 1989). We follow Romer (1922) and Jenkins (1971a) in reconstructing an origin for *m.*
 490 *biceps brachii* in a depression on the lateral surface of the procoracoid, cranial to the procoracoid-
 491 coracoid suture and inferior to the procoracoid foramen (Fig. 7). A second head (*m. biceps brachii brevis*)
 492 may have originated on a scar on the lateral surface of the coracoid tip (Kemp, 1980b). *M. biceps brachii*
 493 inserts via a tendon on or near the radial tuberosity of the radius in all tetrapods (Table 2), and is
 494 reconstructed similarly in *Massetognathus pascuali* (Fig. 4).

495

496 **M. triceps brachii (Figs. 4, 7, 8)**

497 *M. triceps brachii* is present in all tetrapods (Table 2). Despite the name, triceps divisions vary in number
 498 from four in urodeles, lepidosaurs, and mammals (if *m. dorsoepitrochlearis* is an *m. triceps* derivative) to
 499 five in crocodylians (Diogo et al. 2009; Abdala & Diogo, 2010). Holmes (1977) and Abdala and Diogo
 500 (2010) considered a complement of four comprising a coracoid head, a scapular head, and two humeral
 501 heads to be plesiomorphic for amniotes. In extant *Iguana*, *m. triceps brachii coracoideus* originates from a
 502 scar on the medial side of the coracoid, close to its caudal tip, while *m. triceps brachii scapularis* takes

503 origin on a scar near the caudal base of the scapula (Romer, 1922). In extant monotremes, m. triceps
504 coracoideus is absent whereas m. triceps brachii scapularis originates along a ridge dividing the
505 infraspinous fossa from the subscapular fossa, a feature likely homologous with the axillary border of the
506 scapula in other tetrapods given the relocation of m. subscapularis from the medial surface of the scapula
507 to the posterolateral border (Gambaryan et al. 2015). *Massetognathus pascuali* has a scar on the
508 caudomedial surface of the scapula just superior to the supraglenoid buttress, and another on the medial
509 edge of the caudal tip of the coracoid. We follow Jenkins (1971a) in reconstructing an m. triceps brachii
510 scapularis on the former, and a possible m. triceps brachii coracoideus on the latter (Fig. 7). There is
511 reason to be skeptical about the presence of m. triceps brachii coracoideus in cynodonts: Romer (1922)
512 also favored the supraglenoid scar as the origin of the scapular head of m. triceps brachii, but considered
513 the coracoid head to have been lost, while Kemp (1980a) reasoned that a coracoid head for m. triceps
514 brachialis is incompatible with an extended, horizontal humerus, and placed m. biceps brachii at the tip of
515 the coracoid instead. There are no unambiguous sites of origin for the two humeral heads of m. triceps
516 brachii on the humerus of *M. pascuali*, but these are likely to have originated somewhere along the
517 medial and lateral surfaces of the humeral diaphysis as in the case of all tetrapods possessing them
518 (Table 2), distal to the insertions of m. teres major and m. teres minor (Romer, 1922). All divisions of m.
519 triceps brachii insert via a common tendon on the olecranon process of the ulna (Fig. 8).

520 521 DISCUSSION

522 The present study integrates evidence from an extant phylogenetic bracket with direct observation of
523 bony features, and corroborates earlier work (Gregory & Camp, 1918; Jenkins, 1971a) in recovering a
524 near-therian complement of shoulder-actuating muscles in cynodonts. All but five muscles were
525 reconstructed in *Massetognathus pascuali* as strong level I inferences, with the exceptions being m.
526 pectoralis minor, m. deltoideus acromialis, m. teres minor, m. supraspinatus, and m. teres major. The first
527 four are level II inferences, while the last is either level I or level II, depending on homology. Of these five
528 muscles, we opted to reconstruct those whose attachments indicate distinct actions on motions of the
529 forelimb at the shoulder (m. teres minor, m. supraspinatus, m. teres major), and omit those with similar
530 actions at the shoulder to muscles already reconstructed as definitely present (m. pectoralis minor~m.
531 pectoralis major; m. deltoideus acromialis~m. deltoideus scapularis). Our reconstructed attachment areas
532 encompass both the excluded potential muscles and their larger neighbors, so that these muscles may be
533 considered together as functional groups in future biomechanical analyses. Studies of extant amniotes
534 have shown that osteological correlates to muscle attachments differ between mammalian and non-
535 mammalian taxa (Holmes, 1977; McGowan, 1986), being more likely to manifest as rugosities in the
536 former, versus depressions and processes in the latter (Bryant & Seymour, 1990). *M. pascuali* exhibits a
537 combination of well-marked depressions (e.g. m. biceps brachii and m. coracobrachialis origins) and
538 rugosities (e.g. rotator cuff and m. latissimus dorsi insertions), while distinct processes seem to be rare
539 (coracoid origin of m. triceps brachii, if present). A mix of mammal-like and non-mammal-like muscle
540 scars is consistent with the intermediate phylogenetic position of cynodonts. Notably, several of the
541 muscles reconstructed (namely, m. pectoralis, m. deltoideus, m. latissimus dorsi, m. teres major, and m.
542 teres minor) have long, narrow insertions extending proximodistally along the humeral diaphysis, raising
543 the question of whether resistance to torsion in long, flat muscles might present constraints on humeral
544 movement.

545 546 Comparison with previous range of motion measurements

547 The mobility of individual joints has not been extensively documented for the cynodont forelimb, and most
548 workers have focused on the gleno-humeral joint over more proximal or distal articulations. Jenkins
549 (1971a) reported 30° of long-axis rotation and 40° of adduction at the gleno-humeral joint for cynodonts in
550 general, while Kemp (1980a, 1980b) reported 90° of long-axis rotation, nearly 90° of protraction-retraction
551 (“from almost transverse to fairly close to posteriorly directed”), and a “reasonable degree” of abduction-
552 adduction at the same joint for *Luangwa* and *Procynosuchus*. Both Jenkins’ and Kemp’s numbers fall
553 close to the limits reported here, with the exception of Kemp’s long-axis rotation measurement, which
554 exceeds ours by 20°. The discrepancy between Jenkins’ and Kemp’s measurements may be
555 phylogenetic, but may also reflect differing assumptions of joint kinematics. While both workers
556 hypothesized a translational component of humeral motion, Jenkins’ values resemble our rotation-only
557 measurements, while Kemp’s estimates are closer to our combined translation-rotation measurements
558 (Table 1). Oliveira and Schultz (2016) measured 70° of abduction-adduction, 15-20° of retraction-

559 protraction, and an unspecified amount of long-axis rotation at the gleno-humeral joint in *Trucidocynodon*.
560 Their measurements fall within our maximum ranges, although they do not report their joint space
561 assumptions and coordinate systems.

562
563 Kemp (1980b) reported over 90° of flexion-extension for the radius and ulna at their respective
564 articulations with the humerus. Oliveira and Schultz (2016) considered the radius and ulna as a functional
565 unit, measuring over 100° of flexion-extension at the elbow. Oliveira and Schultz additionally reported 25°
566 of mediolateral rotation at the clavo-interclavicular joint, and 15° of roll, 25° of yaw, and 30° of pitch at the
567 acromio-clavicular joint. Again, these values all fall well within our measured ranges (Table 1).

568
569 It should be stressed that the angular ranges reported in Table 1 represent maximum estimates of joint
570 mobility. It is well established that extrinsic soft tissues such as ligaments, joint capsules, labra, muscles,
571 and integument restrict range of motion in an intact animal to a subset of the mobility assessed from
572 manipulation of dry bones (Pierce et al. 2012; Hutson & Hutson, 2012, 2014; Arnold et al. 2014); although
573 the shoulder appears to be less constrained than the hip, and long-axis rotation seems to be the most
574 affected (Pierce et al. 2012). The aim of this analysis was to establish reasonable maximum ranges as a
575 basis for future validation and refinement, and we fully expect the maximum range of motion at all joints
576 modeled here to decrease substantially with the imposition of soft tissue constraints. Radiographic
577 studies of *in vivo* joint utilization (e.g. Fischer, 1994; Kambic et al. 2014) suggest that an even smaller
578 fraction of that mobility is actually employed during normal locomotion, with the remaining available joint
579 surface reserved for non-locomotor behaviors.

580

581 **Comparison with previous muscle reconstructions**

582 Using mammalian muscle anatomy as reference, Oliveira and Schultz (2016) reconstructed the pectoral
583 girdle and forelimb musculature of *Trucidocynodon riograndensis*, a Brazilian Triassic eucynodont. The
584 present reconstruction agrees with theirs in the location of the scapular and humeral heads of m. triceps
585 brachii, and differs in the relative arrangement of m. latissimus dorsi and m. teres major insertions (m.
586 latissimus dorsi inserts distal and medial to m. teres major in our reconstruction, whereas m. latissimus
587 dorsi is proximal and lateral in Oliveira and Schultz, 2016), and the presence of a humeral origin for m.
588 biceps brachii (absent in ours, present in theirs). We attribute these discrepancies to our use of extant
589 phylogenetic bracketing for determining muscle attachments, in contrast to their adherence to mammalian
590 anatomy. Unlike Oliveira and Schultz, we stopped short of recreating the morphology of the muscles
591 themselves. While Lautenschlager (2013) and others (Holliday, 2009; Cuff & Rayfield, 2015) have shown
592 the feasibility of using topography and spatial exclusion to establish the morphology of tightly juxtaposed
593 cranial muscles with direct attachments, limb muscles tend to be more widely spaced, and Bryant and
594 Seymour (1990) caution that architecture and non-uniform cross-sections can confound three-
595 dimensional reconstructions of muscles with tendinous attachments.

596

597 While Gregory and Camp (1918) recovered a similar muscle reconstruction to ours for the cynodont
598 *Cynognathus*, their skeletal reconstruction differed in one important respect. *Cynognathus* was
599 reconstructed with the scapulocoracoids much closer to the animal's sagittal plane, such that the
600 coracoids appear to contact the interclavicle along the ventral midline. This arrangement resembles that
601 seen in extant monotremes, wherein the coracoids articulate with the interclavicle and the procoracoids
602 are closely apposed, occasionally overlapping asymmetrically (Cave, 1970). This has been suggested to
603 be a derived condition allowing better resistance to compressive forces, and possibly related to fossorial
604 or swimming behaviors (Luo, 2015). In *Massetognathus pascuali*, the length and curvature of the clavicles
605 necessitate substantial separation between the scapulocoracoids, regardless of clavicular mobility. The
606 lateral separation between the scapulocoracoids and the interclavicle tends to get understated in two-
607 dimensional reconstructions, many of which depict a lateral view showing only the smaller vertical
608 component of the gap (e.g. Jenkins, 1971a; Kemp, 1980a, 1980b; Sun & Li, 1985).

609

610 Gregory and Camp (1918) noted that the suprascapular cartilages in their reconstruction are probably too
611 small, as the dorsalmost extent of these structures is still far ventral to the tops of the neural spines in the
612 vertebral column, possibly compromising the ability of m. rhomboideus and m. trapezius to suspend the
613 thorax. Greater separation between the scapulocoracoids would ameliorate this by placing the scapulae
614 higher up on the animal's body wall. Curiously, Gregory and Camp go on to hypothesize a thin

615 epicoracoid element in the Permian therapsid *Moschops*, spanning the gap between the clavicles,
616 procoracoids, and the interclavicle. It is unclear why they did not propose a similar structure in
617 *Cynognathus*, but the irregular ventromedial margins of the procoracoid and coracoid (Fig. 7) in
618 *Massetognathus pascuali* are consistent with having possibly articulated with unossified epicoracoid
619 cartilages in life, as seen in extant lepidosaurs (Fürbringer, 1900). The wide separation between the
620 scapulocoracoids and the midventral interclavicle in *M. pascuali* is suggestive of a functional
621 transformation away from the massive, heavily ossified, “U”-shaped girdles seen at the base of the
622 synapsid tree, possibly reflecting a shift in the loading regime experienced by the forelimb and pectoral
623 girdle from mediolateral compression towards more vertically-oriented reaction forces (Jenkins, 1971a).
624 Continued reduction of the endochondral primary girdle (scapulocoracoid) and dermal secondary girdle
625 (clavicle, interclavicle) throughout cynodont evolution may have resulted in progressively greater
626 separation between the scapulocoracoids, presaging the fully independent therian pectoral girdle,
627 dominated by a large scapula and a reduced, strutlike clavicle (Jenkins & Weijs, 1979; Luo, 2015).

629 **Mobility of the pectoral girdle**

630 Kemp (1980b) hypothesized that some degree of mobility in the cynodont pectoral girdle would have
631 been necessary in order to increase the functional length of the forelimb to match that of the hindlimb.
632 Hopson (2015) also considered the issue of stride length in his analysis of the pelycosaur *Dimetrodon*,
633 attributing it to mediolateral rotation of the pectoral girdle from increased bending of the trunk. Lateral
634 bending was suggested to have been lost or greatly reduced in therapsids (Jenkins, 1971a; Kemp, 2005),
635 and it seems unlikely that the cynodont pectoral girdle could have rotated as a single unit with respect to
636 the axial skeleton. It is worth noting that multiple derived amniote lineages seem to have converged on
637 some degree of pectoral girdle mobility—substantial translational coracosternal mobility has been
638 reported in several extant non-mammals, including lepidosaurs (Peterson, 1973; Jenkins & Goslow,
639 1983), ornithomirans (Baier et al. 2013), and pseudosuchians (Baier & Gatesy, 2013). Little is yet known
640 about the soft-tissue constraints acting on the joints of the **non-mammaliaform** synapsid pectoral girdle.
641 Based on osteology, there is no *a priori* reason to reject the possibility that rotation at the clavo-
642 interclavicular and acromio-clavicular joints may have furnished a degree of independent scapulocoracoid
643 mobility in *Massetognathus pascuali*. Experimenting with hypothetical, non-physiological poses reveals
644 that scapulocoracoid mobility allows a significant increase in maximum forelimb excursion, as well as
645 slightly more medial placement of the wrist (Fig. 9). Among derived probainognathian cynodonts,
646 tritylodonts (though not tritheledonts), basal mammaliaforms, and monotremes exhibit expanded lateral
647 processes on their interclavicles that overlap substantially with the proximal ends of their clavicles,
648 precluding clavo-interclavicular mobility (Sues & Jenkins, 2006; Luo, 2015). The ability to unilaterally
649 move one side of the pectoral girdle may have helped functionally separate limb pairs, possibly laying the
650 groundwork for the evolution of asymmetrical gaits. However, if clavo-interclavicular mobility was indeed
651 present in more stem-ward cynodonts, the robust, immobile pectoral girdle of monotremes would then
652 represent an interesting atavistic reversal (Ji et al. 1999).

654 **Epiphyses and range of motion**

655 Among amniotes, ossified epiphyses with well-defined articular surfaces arose convergently in mammals
656 and lepidosaurs (Haines, 1969); unossified epiphyses lacking growth plates are plesiomorphic for
657 cynodonts (Luo et al. 2007b), and the articular surfaces of long bones are thought to have been extended
658 and elaborated by cartilaginous structures in life (Jenkins, 1971a; Jenkins & Parrington, 1976).
659 Progressive removal of connective tissues in archosaurs has shown that cartilage may increase the range
660 of motion available at a joint (Hutson & Hutson, 2012), and that the surface morphology of cartilage caps
661 may differ sufficiently from that of the underlying bone to meaningfully alter joint action (Holliday et al.
662 2010). While it is unclear how cynodont cartilage caps compared to those of either extant crocodylians or
663 extant avians in thickness, both Jenkins and Kemp mentioned them in their reconstructions of cynodont
664 limb function. Jenkins (1971) noted the discrepancy in curvature between the “notch-shaped” hemisellar
665 glenoid and the convex, ovoid humeral head, and posited that a substantial thickness of cartilage must
666 have been present in life to increase congruence between the articular surfaces, much like in extant
667 archosaurs (Holliday et al. 2010). Kemp (1980b), on the other hand, argued that any cartilaginous
668 intermediary would have had to be “absurdly thick” to match the humeral head’s curvature to that of the
669 glenoid fossa, and interpreted the incongruity in articular morphology as representing a “rolling” rather
670 than sliding articulation at the gleno-humeral joint.

671
672 The 0.25 mm of epiphyseal cartilage modeled here (both proximal and distal) is based on mammalian
673 measurements from the literature. An articular cap measuring 0.25 mm on each epiphysis adds up to 0.5
674 mm per long bone, and represents an approximate 1% contribution to humeral length in *Massetognathus*
675 *pascuali*. This is comparable with measurements taken by Holliday et al. (2010) from extant avians
676 (0.07%–3.72%), but falls below the 7.99% measured in *Alligator*. However, it is unclear whether
677 archosaurs, and *Alligator* in particular, represent good models for reconstructing non-mammaliaform
678 synapsids. We were unable to locate cartilage measurements for lepidosaurs or amphibians in the
679 literature, but epiphyseal cartilage is likely to be thin in the former clade given the presence of extensive
680 secondary ossification centers (Haines, 1969), and extensive in the latter due to their aquatic tendencies.

681
682 Greater cartilage thickness—and consequently greater joint space—may indeed further increase range of
683 motion. An informal sensitivity test on this model revealed that a threefold increase in joint space (from
684 0.50 mm to 1.50 mm) increased rotation-only range of motion to ranges comparable to combined
685 translation-rotation levels measured with the original 0.50 mm joint space. However, thicker cartilage may
686 significantly alter functional articular morphology from that of the underlying bone (Holliday et al. 2010).
687 Since the superficial cartilage morphology is unpreserved, we are unable to satisfactorily bound this into a
688 testable problem, and instead present measurements based on a joint space of 0.5 mm (Table 1) a
689 conservative estimate.

690 691 **Evolution toward the therian glenoid fossa**

692 The separate scapular and coracoid facets that together comprise the glenoid fossa deserve closer
693 examination. The basal cynodont glenoid is commonly described as laterally or posterolaterally-facing
694 and either “hemisellar” (Jenkins, 1971a) or “notch”-shaped (Kemp, 2005), a morphology retained through
695 progressively more derived eucynodonts and mammaliaforms before finally evolving into the familiar,
696 ventrally-directed, mammalian ball-and-socket joint in the Jurassic theriomorphs (Luo, 2015). In
697 *Massetognathus pascuali*, the scapular and coracoid facets of the glenoid are dissimilar in morphology
698 (Fig. 10). Viewed together, the two create the impression of a unified notch, but close examination reveals
699 that the coracoid facet is mediolaterally convex, with a small, laterally projecting lip at its ventral apex
700 creating a small degree of dorsoventral concavity. The scapular facet is set at an obtuse angle (approx.
701 130°) to the coracoid facet, and is slightly concave at the center (*contra* Jenkins, 1970b, where it is
702 described as slightly convex in the same specimen). A procoracoid contribution to the glenoid seems
703 somewhat variable among eucynodonts. Within the cynognathian clade, it is present ancestrally in
704 *Cynognathus*, in which it makes up a small, cranial portion of the glenoid articulation (Jenkins, 1971a);
705 this condition is also retained in diademodonts (Hopson & Kitching, 2001). However, the procoracoid is
706 completely excluded from the glenoid in *Massetognathus pascuali* (as seen in our specimens) and other
707 traversodonts (Rowe, 1988). A procoracoid contribution to the glenoid is slightly present in basal
708 probainognathians (Hopson & Kitching, 2001), and completely absent from Mammaliaforma onwards
709 (Rowe, 1988).

710
711 Jenkins (1971a) interpreted the separate scapular and coracoid facets as representing a functional
712 division within the glenoid, with a possibly ventrally-facing scapular facet serving to transmit the vertical
713 component of ground reaction force during locomotion. The full-girdle reconstruction of *Massetognathus*
714 *pascuali* presented here confirms a ventral orientation for the scapular facet (Fig. 3). Taking Jenkins’ logic
715 a step further, one could hypothesize that the laterally-facing coracoid and procoracoid may have initially
716 functioned to accommodate a substantial lateral component of ground reaction force. Increasingly
717 parasagittal limb kinematics and increasingly vertically-oriented ground reaction forces in more crown-
718 ward taxa might have accompanied progressively reduced coracoid and procoracoid contributions to the
719 glenoid, until all that remained was a broadened, deepened scapular facet forming a ventrally-facing,
720 socket-shaped glenoid. Under this scenario, the characteristic therian ball-and-socket joint was achieved
721 not through reorientation of an ancestrally laterally-facing composite glenoid as a whole, but rather
722 progressive reduction, ventral reorientation, and possibly partial assimilation (Vickaryous & Hall, 2006) of
723 the plesiomorphic coracoid facet into a pre-existing, ventrally-facing, concave scapular facet. This
724 hypothesis of glenoid transformation is wholly compatible with macroevolutionary trends in coracoid
725 reduction and scapular expansion among derived cynodonts, as reported by Luo (2015) as well as
726 Jenkins and Weijs (1979).

727

728 Differences between monotreme and cynodont pectoral girdles

729 As the sister group to crown therians, extant monotremes have long been used as models for
730 understanding cynodont biology (Gregory & Camp, 1918; Kemp, 1980a; Luo, 2007). In addition to the
731 foregoing discrepancy in scapulocoracoid position, our reconstruction of *Massetognathus pascuali* differs
732 from monotremes in a number of important musculoskeletal respects, namely the length of the clavicles
733 and the extent of their overlap with the interclavicle; the significant mediolateral torsion present in the
734 monotreme scapula but absent in *M. pascuali*; the higher aspect ratio and weaker long-axis torsion of the
735 humerus in *M. pascuali*; the insertion of m. latissimus dorsi (much more distal in monotremes); the origin
736 of m. teres minor (far dorsal in monotremes); and the orientation of the fossa for m. subcoracoscapularis
737 (medially-facing in *M. pascuali* and all other amniotes, posterolaterally-facing in monotremes). In
738 reconstructing *M. pascuali*, we found that monotreme muscle attachments were frequently the exception
739 in the extant phylogenetic bracket, with the monotreme m. subcoracoscapularis in particular differing from
740 its location and attachments in therians and saurians alike. These discrepancies raise the question of
741 whether ecological specializations may have left a confounding signature on the musculoskeletal
742 organization of the monotreme pectoral girdle and forelimb. It seems reasonable to entertain the
743 possibility that niche adaptation and low ecological diversity in extant monotremes may render them
744 compromised locomotor analogues for cynodonts (Howell, 1936; Jenkins, 1971a; Jenkins, 1989). If this is
745 the case, it may be prudent to situate interpretations of postcranial function in cynodonts within a wide,
746 comparative context spanning mammalian and non-mammalian forms.

747

748 CONCLUSION

749 The musculoskeletal reconstruction of *Massetognathus pascuali* presented here recovers maximum
750 ranges of motion and muscle origins and insertions comparable to those found in previous studies, and
751 reveals features of the pectoral girdle not previously described for this genus, such as a wide separation
752 between the scapulocoracoids and the interclavicle, possible mobility at the clavo-interclavicular and
753 acromio-clavicular joints, and a ventrally-facing, concave scapular facet of the glenoid fossa. Extensive
754 functional interpretation of our 3D reconstruction awaits further comparison with extant mammalian and
755 non-mammalian taxa. Having already mapped well-defined muscle attachment areas onto a digital
756 skeleton, we have laid the groundwork for constructing interactive musculoskeletal models. Future
757 experiments that combine detailed musculoskeletal modeling with *ex vivo* and *in vivo* data from extant
758 taxa will provide the opportunity to investigate the relationship between skeletal posture and muscle
759 function, and shed further light on the cynodont pectoral limb and its significance to the rise of mammals.

760

761 ACKNOWLEDGEMENTS

762 We thank Katrina Jones for her help with μ CT scanning, and Jessica Cundiff for assistance with
763 specimens in the MCZ. The Pierce and Biewener labs provided valuable feedback. Financial support was
764 provided by Harvard University funds made available to S.E.P. μ CT data and bone models are repositied
765 in the Department of Vertebrate Paleontology, Museum of Comparative Zoology, Harvard University. The
766 authors declare no conflicts of interest.

767

768 AUTHOR CONTRIBUTIONS

769 All authors contributed to study conception and manuscript preparation. P.H.L. and S.E.P. contributed to
770 experimental design, data acquisition, and interpretation. All authors gave final approval for publication.

771

772 REFERENCES

773 Abdala, V., & Diogo, R. (2010). Comparative anatomy, homologies and evolution of the pectoral and
774 forelimb musculature of tetrapods with special attention to extant limbed amphibians and reptiles. *Journal*
775 *of Anatomy*, 217(5), 536–573.

776

777 Arnold, P., Fischer, M. S., & Nyakatura, J. A. (2014). Soft tissue influence on *ex vivo* mobility in the hip of
778 Iguana: comparison with *in vivo* movement and its bearing on joint motion of fossil sprawling tetrapods.
779 *Journal of Anatomy*, 225(1), 31–41.

780

781 Baier, D. B., & Gatesy, S. M. (2013). Three-dimensional skeletal kinematics of the shoulder girdle and
782 forelimb in walking Alligator. *Journal of Anatomy*, 223(5), 462–473.

- 783
784 Baier, D. B., Gatesy, S. M., & Dial, K. P. (2013). Three-dimensional, high-resolution skeletal kinematics of
785 the avian wing and shoulder during ascending flapping flight and uphill flap-running. *PloS One*, 8(5),
786 e63982.
- 787
788 Bates, K. T., & Schachner, E. R. (2012). Disparity and convergence in bipedal archosaur locomotion.
789 *Journal of the Royal Society, Interface / the Royal Society*, 9(71), 1339–1353.
- 790
791 Bonaparte, J. F. (1963). Descripción del esqueleto postcraneano de Exaeretodon (Cynodontia-
792 Traversodontidae). *Acta Geológica Lilloana*, 4, 5–52.
- 793
794 Brainerd, E. L., Baier, D. B., Gatesy, S. M., Hedrick, T. L., Metzger, K. A., Gilbert, S. L., & Crisco, J. J.
795 (2010). X-ray reconstruction of moving morphology (XROMM): precision, accuracy and applications in
796 comparative biomechanics research. *Journal of Experimental Zoology. Part A, Ecological Genetics and*
797 *Physiology*, 313(5), 262–279.
- 798
799 Bryant, H. N., & Seymour, K. L. (1990). Observations and comments on the reliability of muscle
800 reconstruction in fossil vertebrates. *Journal of Morphology*, 206(1), 109–117.
- 801
802 Cave, A. J. E. (1970). Observations on the monotreme interclavicle. *Journal of Zoology*, 160(3), 297–312.
- 803
804 Cheng, C.-C. (1955). The development of the shoulder region of the opossum, *Didelphys virginiana*, with
805 special reference to the musculature. *Journal of Morphology*, 97(3), 415–471.
- 806
807 Coues, E. (1871). *On the Myology of the Ornithorhynchus*.
- 808
809 Coues, E. & Wyman, J. (1872). *The osteology and myology of Didelphys virginiana* (p. 116). Boston :The
810 Society,.
- 811
812 Cuff, A. R., & Rayfield, E. J. (2015). Retrodeformation and muscular reconstruction of ornithomimosaurian
813 dinosaur crania. *PeerJ*, 3, e1093.
- 814
815 Davison, A. (1895). A contribution to the anatomy and phylogeny of *Amphiuma means* (Gardner). *Journal*
816 *of Morphology*, 11(2), 375–410.
- 817
818 Delp, S. L., & Loan, J. P. (1995). A graphics-based software system to develop and analyze models of
819 musculoskeletal structures. *Computers in Biology and Medicine*, 25(1), 21–34.
- 820
821 Diogo, R., Abdala, V., Aziz, M. A., Lonergan, N., & Wood, B. A. (2009). From fish to modern humans--
822 comparative anatomy, homologies and evolution of the pectoral and forelimb musculature. *Journal of*
823 *Anatomy*, 214(5), 694–716.
- 824
825 Field, D. A. (1988). Laplacian smoothing and Delaunay triangulations. *Communications in Applied*
826 *Numerical Methods*, 4(6), 709–712.
- 827
828 Fischer, M. S. (1994). Crouched posture and high fulcrum, a principle in the locomotion of small
829 mammals: The example of the rock hyrax (*Procavia capensis*) (Mammalia: Hyracoidea). *Journal of*
830 *Human Evolution*, 26(5–6), 501–524.
- 831
832 Fischer, M. S., Schilling, N., Schmidt, M., Haarhaus, D., & Witte, H. (2002). Basic limb kinematics of small
833 therian mammals. *The Journal of Experimental Biology*, 205(Pt 9), 1315–1338.
- 834
835 Fürbringer, M. (1900). Zur vergleichenden Anatomie des Brustschulterapparates und der
836 Schultermuskeln. IV Teil. *Jena Ische Zeitschr. Naturwiss.* , Jena, 34, 351.
- 837

- 838 Gambaryan, P. P., Kuznetsov, A. N., Panyutina, A. A., & Gerasimov, S. V. (2015). Shoulder girdle and
839 forelimb myology of extant Monotremata. *Russian Journal of Theriology*, 14(1), 1–56.
840
- 841 Gatesy, S. M. (1991). Hind limb movements of the American alligator (*Alligator mississippiensis*) and
842 postural grades. *Journal of Zoology*, 224(4), 577–588.
843
- 844 Gatesy, S. M., Baier, D. B., Jenkins, F. A., & Dial, K. P. (2010). Scientific rotoscoping: a morphology-
845 based method of 3-D motion analysis and visualization. *Journal of Experimental Zoology. Part A,*
846 *Ecological Genetics and Physiology*, 313(5), 244–261.
847
- 848 George, R. M. (1977). The limb musculature of the Tupaiidae. *Primates; Journal of Primatology*, 18(1), 1–
849 34.
850
- 851 Gregory, W. K., & Camp, C. L. (1918). *Studies in comparative myology and osteology*. American Museum
852 of Natural History.
853
- 854 Grood, E. S., & Suntay, W. J. (1983). A joint coordinate system for the clinical description of three-
855 dimensional motions: application to the knee. *Journal of Biomechanical Engineering*.
856
- 857 Haines, R. W. (1942). THE EVOLUTION OF EPIPHYSES AND OF ENDOCHONDRAL BONE. *Biological*
858 *Reviews of the Cambridge Philosophical Society*, 17(4), 267–292.
859
- 860 Harvey, K. J., & Warburton, N. (2010). Forelimb musculature of kangaroos with particular emphasis on
861 the tammar wallaby *Macropus eugenii* (Desmarest, 1817). *Australian Mammalogy*, 32(1), 1–9.
862
- 863 Hildebrand, M. (1989). The quadrupedal gaits of vertebrates. *Bioscience*, 39(11), 766–775.
864
- 865 Holliday, C. M. (2009). New insights into dinosaur jaw muscle anatomy. *Anatomical Record*, 292(9),
866 1246–1265.
867
- 868 Holliday, C. M., Ridgely, R. C., Sedlmayr, J. C., & Witmer, L. M. (2010). Cartilaginous epiphyses in extant
869 archosaurs and their implications for reconstructing limb function in dinosaurs. *PLoS One*, 5(9).
870 <https://doi.org/10.1371/journal.pone.0013120>
871
- 872 Holmes, R. (1977). The osteology and musculature of the pectoral limb of small captorhinids. *Journal of*
873 *Morphology*, 152(1), 101–140.
874
- 875 Hoppe, H. (2008). Poisson Surface Reconstruction and Its Applications. In *Proceedings of the 2008 ACM*
876 *Symposium on Solid and Physical Modeling* (pp. 10–10). New York, NY, USA: ACM.
877
- 878 Hopson, J. A. (2015). Fossils, Trackways, and Transitions in Locomotion: A Case Study of *Dimetrodon*. In
879 *Great Transformations in Vertebrate Evolution*.
880
- 881 Hopson, J. A., & Kitching, J. W. (2001). A probainognathian cynodont from South Africa and the
882 phylogeny of nonmammalian cynodonts. *Bulletin of the Museum of Comparative Zoology at Harvard*
883 *College*, 156(1), 5–35.
884
- 885 Howell, A. B. (1936). The phylogenetic arrangement of the muscular system. *The Anatomical Record*.
886
- 887 Howell, A. B. (1937a). Morphogenesis of the Shoulder Architecture. Part V. Monotremata. *The Quarterly*
888 *Review of Biology*, 12(2), 191–205.
889
- 890 Howell, A. B. (1937b). The Swimming Mechanism of the Platypus. *Journal of Mammalogy*, 18(2), 217–
891 222.
892

- 893 Hutchinson, J. R., Anderson, F. C., Blemker, S. S., & Delp, S. L. (2005). Analysis of hindlimb muscle
894 moment arms in *Tyrannosaurus rex* using a three-dimensional musculoskeletal computer model:
895 implications for stance, gait, and speed. *Paleobiology*, 31(4), 676–701.
896
- 897 Hutchinson, J. R., Rankin, J. W., Rubenson, J., Rosenbluth, K. H., Siston, R. A., & Delp, S. L. (2015).
898 Musculoskeletal modelling of an ostrich (*Struthio camelus*) pelvic limb: influence of limb orientation on
899 muscular capacity during locomotion. *PeerJ*, 3, e1001.
900
- 901 Hutson, J. D., & Hutson, K. N. (2012). A test of the validity of range of motion studies of fossil archosaur
902 elbow mobility using repeated-measures analysis and the extant phylogenetic bracket. *The Journal of*
903 *Experimental Biology*, 215(Pt 12), 2030–2038.
904
- 905 Hutson, J. D., & Hutson, K. N. (2014). A repeated-measures analysis of the effects of soft tissues on wrist
906 range of motion in the extant phylogenetic bracket of dinosaurs: Implications for the *The Anatomical*
907 *Record*. Retrieved from <http://onlinelibrary.wiley.com/doi/10.1002/ar.22903/full>
908
- 909 Jenkins, F. A. (1970a). Cynodont postcranial anatomy and the “prototherian” level of mammalian
910 organization. *Evolution; International Journal of Organic Evolution*, 230–252.
911
- 912 Jenkins, F. A. (1970b). The Chañares (Argentina) Triassic reptile fauna VII. The postcranial skeleton of
913 the traversodontid *Massetognathus pascuali* (Therapsida, Cynodontia). *Breviora*, 352, 1–28.
914
- 915 Jenkins, F. A. (1971a). *The postcranial skeleton of African cynodonts: problems in the early evolution of*
916 *the mammalian postcranial skeleton*. Harvard MCZ.
917
- 918 Jenkins, F. A. (1971b). Limb posture and locomotion in the Virginia opossum (*Didelphis marsupialis*) and
919 in other non-cursorial mammals. *Journal of Zoology*, 165(3), 303–315.
920
- 921 Jenkins, F. A. (1989). Monotremes and the biology of Mesozoic mammals. *Netherlands Journal of*
922 *Zoology*.
923
- 924 Jenkins, F. A. (1993). The evolution of the avian shoulder joint. *American Journal of Science*.
925
- 926 Jenkins, F. A., & Goslow, G. E. (1983). The functional anatomy of the shoulder of the savannah monitor
927 lizard (*Varanus exanthematicus*). *Journal of Morphology*, 175(2), 195–216.
928
- 929 Jenkins, F. A., & Parrington, F. R. (1976). The postcranial skeletons of the Triassic mammals
930 *Eozostrodon*, *Megazostrodon* and *Erythrotherium*. *Philosophical Transactions of the Royal Society of*
931 *London. Series B, Biological Sciences*, 273(926), 387–431.
932
- 933 Jenkins, F. A., & Weijs, W. A. (1979). The functional anatomy of the shoulder in the Virginia opossum
934 (*Didelphis virginiana*). *Journal of Zoology*, 188(3), 379–410.
935
- 936 Ji, Q., Luo, Z.-X., & Ji, S. A. (1999). A Chinese triconodont mammal and mosaic evolution of the
937 mammalian skeleton. *Nature*, 398(6725), 326–330.
938
- 939 Ji, Q., Luo, Z.-X., Yuan, C.-X., & Tabrum, A. R. (2006). A swimming mammaliaform from the Middle
940 Jurassic and ecomorphological diversification of early mammals. *Science*, 311(5764), 1123–1127.
941
- 942 Jouffroy, F. K. L., Saban, J., Souteyrand-Boulenger, R., Jouffroy, J., & Others. (1971). *Mammifères:*
943 *musculature des membres, musculature peaucière, musculature des monotrèmes*. *Arthrologie*. Retrieved
944 from [http://www.sidalc.net/cgi-](http://www.sidalc.net/cgi-bin/wxis.exe/?IsisScript=FCL.xis&method=post&formato=2&cantidad=1&expresion=mfn=001693)
945 [bin/wxis.exe/?IsisScript=FCL.xis&method=post&formato=2&cantidad=1&expresion=mfn=001693](http://www.sidalc.net/cgi-bin/wxis.exe/?IsisScript=FCL.xis&method=post&formato=2&cantidad=1&expresion=mfn=001693)
946
- 947 Kambic, R. E., Roberts, T. J., & Gatesy, S. M. (2014). Long-axis rotation: a missing degree of freedom in
948 avian bipedal locomotion. *The Journal of Experimental Biology*, 217(Pt 15), 2770–2782.

- 949
950 Kemp, T. S. (1978). Stance and gait in the hindlimb of a theriocephalian mammal-like reptile. *Journal of*
951 *Zoology*, 186(2), 143–161.
952
953 Kemp, T. S. (1980a). Aspects of the structure and functional anatomy of the Middle Triassic cynodont
954 Luangwa. *J. Zool., Lond.*, 191, 193–239.
955
956 Kemp, T. S. (1980b). The Primitive Cynodont Procynosuchus: Structure, Function and Evolution of the
957 Postcranial Skeleton. *Philosophical Transactions of the Royal Society of London. Series B, Biological*
958 *Sciences*, 288(1027), 217–258.
959
960 Kemp, T. S. (2005). *The Origin and Evolution of Mammals*. OUP Oxford.
961
962 Kirsch, J. A. W. (1973). *Notes for the dissection of the opossum, Didelphis virginiana*. Madison, WI.
963
964 Kühne, W. G. (1956). *The Liassic therapsid Oligokyphus*. British Museum.
965
966 Lautenschlager, S. (2013). Cranial myology and bite force performance of *Erlisosaurus andrewsi*: a novel
967 approach for digital muscle reconstructions. *Journal of Anatomy*, 222(2), 260–272.
968
969 Leach, D. (1977). The forelimb musculature of marten (*Martes americana* Turton) and fishes (*Martes*
970 *pennanti* Erxleben). *Canadian Journal of Zoology*, 55(1), 31–41.
971
972 Liu, J. (2007). *New traversodontid materials from North Carolina, USA and the taxonomy, phylogeny of*
973 *Traversodontidae (Synapsida: Cynodontia)*. COLUMBIA UNIVERSITY.
974
975 Liu, J., & Abdala, F. (2014). Phylogeny and Taxonomy of the Traversodontidae. In C. F. Kammerer, K. D.
976 Angielczyk, & J. Fröbisch (Eds.), *Early Evolutionary History of the Synapsida* (pp. 255–279). Springer
977 Netherlands.
978
979 Liu, J., & Olsen, P. (2010). The Phylogenetic Relationships of Eucynodontia (Amniota: Synapsida).
980 *Journal of Mammalian Evolution*, 17(3), 151–176.
981
982 Luo, Z.-X. (2007). Transformation and diversification in early mammal evolution. *Nature*, 450(7172),
983 1011–1019.
984
985 Luo, Z.-X. (2015). Origin of the Mammalian Shoulder. In *Great Transformations in Vertebrate Evolution*.
986
987 Luo, Z.-X., Chen, P., Li, G., & Chen, M. (2007b). A new eutriconodont mammal and evolutionary
988 development in early mammals. *Nature*, 446(7133), 288–293.
989
990 Luo, Z.-X., Meng, Q.-J., Grossnickle, D. M., Liu, D., Neander, A. I., Zhang, Y.-G., & Ji, Q. (2017). New
991 evidence for mammaliaform ear evolution and feeding adaptation in a Jurassic ecosystem. *Nature*,
992 548(7667), 326–329.
993
994 McGowan, C. (1986). The wing musculature of the weka. *Gallirallus Australis*.
995
996 Meers, M. B. (2003). Crocodylian forelimb musculature and its relevance to Archosauria. *The Anatomical*
997 *Record. Part A, Discoveries in Molecular, Cellular, and Evolutionary Biology*, 274(2), 891–916.
998
999 Meng, Q.-J., Grossnickle, D. M., Liu, D., Zhang, Y.-G., Neander, A. I., Ji, Q., & Luo, Z.-X. (2017). New
1000 gliding mammaliaforms from the Jurassic. *Nature*, 548(7667), 291–296.
1001
1002 Miner, R. W. (1925). THE PECTORAL LIMB OF ERYOPS AND OTHER PRIMITIVE TETRAPODS.
1003 *Bulletin of the AMNH*, 51(7).
1004

- 1005 Mivart, S. G. (1869). Notes on the Myology of *Menopoma alleghaniense*. *Proceedings of the Zoological*
1006 *Society of London*, 37(1), 254–271.
- 1007
- 1008 Netter, F. H., Colacino, S., & Others. (1989). *Atlas of human anatomy* (Vol. 11). Ciba-Geigy Summit, NJ.
- 1009
- 1010 Nyakatura, J. A., Allen, V. R., Lauströer, J., Andikfar, A., Danczak, M., Ullrich, H.-J., ... Fischer, M. S.
1011 (2015). A Three-Dimensional Skeletal Reconstruction of the Stem Amniote *Orobates pabsti*
1012 (Diadectidae): Analyses of Body Mass, Centre of Mass Position, and Joint Mobility. *PloS One*, 10(9),
1013 e0137284.
- 1014
- 1015 Oliveira, T. V. D., & Schultz, C. L. (2016). Functional Morphology and Biomechanics of the Cynodont
1016 *Trucidocynodon riograndensis* from the Triassic of Southern Brazil: Pectoral Girdle and Forelimb. *Acta*
1017 *Palaeontologica Polonica*, 61(2), 377–386.
- 1018
- 1019 Parsons, F. G. (1896). 4. Myology of Rodents.—Part II. An Account of the Myology of the Myomorpha,
1020 together with a Comparison of the Muscles of the various Suborders of Rodents. In *Proceedings of the*
1021 *Zoological Society of London* (Vol. 64, pp. 159–192). Wiley Online Library.
- 1022
- 1023 Peterson, J. A. (1973). *Adaptation for arboreal locomotion in the shoulder region of lizards*. University of
1024 Chicago Press.
- 1025
- 1026 Pierce, S. E., Clack, J. A., & Hutchinson, J. R. (2012). Three-dimensional limb joint mobility in the early
1027 tetrapod *Ichthyostega*. *Nature*, 486(7404), 523–526.
- 1028
- 1029 Polly, P. D. (2007). Limbs in mammalian evolution. *Fins into Limbs: Evolution, Development and*
1030 *Transformation*, 245–268.
- 1031
- 1032 Reilly, S. M., & Elias, J. A. (1998). Locomotion in alligator mississippiensis: kinematic effects of speed and
1033 posture and their relevance to the sprawling-to-erect paradigm. *The Journal of Experimental Biology*, 201
1034 (Pt 18), 2559–2574.
- 1035
- 1036 Romer, A. S. (1922). The locomotor apparatus of certain primitive and mammal-like reptiles. *American*
1037 *Museum of Natural History*.
- 1038
- 1039 Romer, A. S. (1924). Pectoral limb musculature and shoulder girdle structure in fish and tetrapods. *The*
1040 *Anatomical Record*, 27(2), 119–143.
- 1041
- 1042 Romer, A. S. (1944). The development of tetrapod limb musculature—the shoulder region of *Lacerta*.
1043 *Journal of Morphology*.
- 1044
- 1045 Romer, A. S. (1956). *Osteology of the Reptiles*. 772 pp. University of Chicago Press, Chicago.
- 1046
- 1047 Romer, A. S. (1967). *The Chanares (Argentina) Triassic reptile fauna. III. Two New Gomphodonts,*
1048 *Massetognathus Pascuali and M. Teruggii* (Vol. 264). Museum of Comparative Zoology.
- 1049
- 1050 Romer, A. S., & Price, L. W. (1940). Review of the Pelycosauria. *Geological Society of America Special*
1051 *Papers*, 28, 1–534
- 1052
- 1053 **Rowe, T. (1988). Definition, diagnosis, and origin of Mammalia. *Journal of Vertebrate Paleontology*, 8(3),**
1054 **241–264.**
- 1055
- 1056 Ruta, M., Botha-Brink, J., Mitchell, S. A., & Benton, M. J. (2013). The radiation of cynodonts and the
1057 ground plan of mammalian morphological diversity. *Proceedings. Biological Sciences / The Royal*
1058 *Society*, 280(1769), 20131865.
- 1059
- 1060 Simon, W. H. (1970). Scale effects in animal joints. I. Articular cartilage thickness and compressive

- 1061 stress. *Arthritis and Rheumatism*, 13(3), 244–256.
- 1062
- 1063 Stein, B. R. (1981). Comparative Limb Myology of Two Opossums, *Didelphis* and *Chironectes*. *Journal of*
- 1064 *Morphology*, (169), 113–140.
- 1065
- 1066 Stein, B. R. (1986). Comparative limb myology of four arvicolid rodent genera (mammalia, rodentia).
- 1067 *Journal of Morphology*, 187(3), 321–342.
- 1068
- 1069 Sues, H.-D., & Jenkins, F. A., Jr. (2006). 5 The Postcranial Skeleton of *Kayentatherium wellsi* from the
- 1070 Lower Jurassic Kayenta Formation of Arizona and the Phylogenetic Significance of Postcranial Features.
- 1071 *Amniote Paleobiology: Perspectives on the Evolution of Mammals, Birds, and Reptiles*, 114.
- 1072
- 1073 Sun, A., & Li, Y. (1985). The postcranial skeleton of the late tritylodont *Bienotheroides*. *Vertebrata*
- 1074 *PalAsiatica*, 23(2).
- 1075
- 1076 Taylor, B. K. (1978). The anatomy of the forelimb in the anteater (*Tamandua*) and its functional
- 1077 implications. *Journal of Morphology*, 157(3), 347–367.
- 1078
- 1079 Vaughan, T. A., Ryan, J. M., & Czaplewski, N. J. (2013). *Mammalogy*. Jones & Bartlett Learning, LLC.
- 1080
- 1081 Vickaryous, M. K., & Hall, B. K. (2006). Homology of the reptilian coracoid and a reappraisal of the
- 1082 evolution and development of the amniote pectoral apparatus. *Journal of Anatomy*, 208(3), 263–285.
- 1083
- 1084 Walter, L. R. (1988). Appendicular Musculature in the Echidna, *Tachyglossus-Aculeatus* (Monotremata,
- 1085 *Tachyglossidae*). *Australian Journal of Zoology*, 36(1), 65–81.
- 1086
- 1087 Walthall, J. C., & Ashley-Ross, M. A. (2006). Postcranial myology of the California newt, *Taricha torosa*.
- 1088 *The Anatomical Record. Part A, Discoveries in Molecular, Cellular, and Evolutionary Biology*, 288(1), 46–
- 1089 57.
- 1090
- 1091 Warburton, N. M., Grégoire, L., Jacques, S., & Flandrin, C. (2014). Adaptations for digging in the forelimb
- 1092 muscle anatomy of the southern brown bandicoot (*Isodon obesulus*) and bilby (*Macrotis lagotis*).
- 1093 *Australian Journal of Zoology*, 61(5), 402–419.
- 1094
- 1095 Watson, D. M. (1917). The Evolution of the Tetrapod Shoulder Girdle and Fore-limb. *Journal of Anatomy*,
- 1096 52(Pt 1), 1–63.
- 1097
- 1098 Witmer, L. M. (1995). The Extant Phylogenetic Bracket and the Importance of Reconstructing Soft
- 1099 Tissues in Fossils. In J. Thomason (Ed.), *Functional Morphology in Vertebrate Paleontology*.
- 1100
- 1101 Zaaf, A., Herrel, A., Aerts, P., & De Vree, F. (1999). Morphology and morphometrics of the appendicular
- 1102 musculature in geckoes with different locomotor habits (Lepidosauria). *Zoomorphology*, 119(1), 9–22.
- 1103
- 1104
- 1105
- 1106
- 1107
- 1108
- 1109
- 1110
- 1111

1112 **Supplementary Figure 1.** Three-dimensional reconstruction of the left pectoral limb of *M. pascuali*.
1113 Reconstructed muscle origins/insertions are listed in legends of Figs. 4-8. This interactive PDF may be
1114 viewed in Adobe Acrobat (Adobe Systems Incorporated, San Jose, CA, USA).

For Peer Review Only

22

1115 **Table 1.** Pectoral limb joint range of motion in the cynodont *Massetognathus pascuali*. All measurements
 1116 are made using 0.5mm joint space (see text for details).
 1117

Clavo-interclavicular						
Z						
+/-medial rotation (°)	-/-lateral rotation (°)					
35	35					
Acromio-clavicular						
Z		Y		X		
+/-medial roll (°)	-/-lateral roll (°)	+/-lateral yaw (°)	-/-medial yaw (°)	+/-cranial pitch (°)	-/-caudal pitch (°)	
35	5	20	20	Unrestricted	5	
Gleno-humeral						
Z		Y		X		
+/-abduction (°)	-/-adduction (°)	+/-retraction (°)	-/-protraction (°)	+/-pronation (°)	-/-supination (°)	
w/o translation	20	20	15	15	15	25
w/ translation	75	45	45	45	35	35
Humero-radial						
Z		Y		X		
+/-extension (°)	-/-flexion (°)	+/-lateral rotation (°)	-/-medial rotation (°)	+/-adduction (°)	-/-abduction (°)	
90	45	Unrestricted	Unrestricted	40	40	
Humero-ulnar						
Z		Y		X		
+/-extension (°)	-/-flexion (°)	+/-lateral rotation (°)	-/-medial rotation (°)	+/-adduction (°)	-/-abduction (°)	
80	60	Unrestricted	Unrestricted	30	30	

1118
 1119
 1120

1121
1122
1123

Table 2. Extant Phylogenetic Bracket with muscle attachments and homologies. Abbreviations: sf., surface; dc, deltopectoral crest

	Urodela		Crocodylia		Rhynchocephalia		Squamata		Cynodontia		Monotremata		Theria	
	Taricha: Walhall & Ashley-Ross (2006) Amphystoma: Diogo et al. (2009) Anphyma: Davison (1895) Menopoma: Mivart (1869)		Alligator, Crocodylus, Osteolemus, Gavialis, Meers (2005)		Sphenodon: Holmes (1977); Miner (1925)		Gekko: Zaaf et al. (1999) Timon: Abdala & Diogo (2010) Varanus: Jenkins & Goslow (1983) Iguana: Holmes (1977); Romer (1922)		Massetognathus pascuali: Present study		Ornithorhynchus: Coues (1871); Gambaryan et al. (2015); Howell (1937a) Tachyglossus: Walter (1988); Gambaryan et al. (2015) Zaglossus: Gambaryan et al. (2015)		Didelphis: Jenkins & Wejs (1979); Kirsch (1973); Stein (1981); Coues & Wyman (1872) Rattus: Parsons (1896) Tupaia: George (1977)	
	o	i	o	i	o	i	o	i	o	i	o	i	o	i
<i>m. latissimus dorsi</i>	+ ¹	o	o	o	+	o	+	o	+	(I)	+	o	+	o
		posterior sf. of humeral dc crest		thoracodorsal fascia		anterodorsal sf. of humerus near m. teres major		posterdorsal sf. of proximal humerus		thoracolumbar fascia		posterior sf. of distal humerus, near entepicondyle		postero-medial sf. of proximal humerus
<i>m. pectoralis major</i>	+ ²	o	+	+	+	o	+	o	+	(I)	+	o	+	o
		sternum and ventral fascia		sternum and sternal ribs		sternum, interclavicle, clavicle, sternocostate		lateral sf. of interclavicle, sternal ribs, and ventral midline fascia		all over ventral sf. of interclavicle		interclavicle and ventral sf. of sternal ribs		sternum
		humeral dc crest		apex of humeral dc crest, near m. deltoideus clavicularis		posterior sf. of humeral dc crest		humeral dc crest		entire length of humeral dc crest		entire length of humeral dc crest		entire length of humeral dc crest
<i>m. pectoralis minor</i>	-		-	-	-		-	-	-	(II)	-	+	+	o
												cranial 1/4 of abdominal midline		xiphoid process of sternum
												Humeral dc crest distal to M. pectoralis major insertion		proximal portion of humeral dc crest
<i>m. deltoideus scapularis</i>	+ ³	o	+	+	+	o	+	+	+	(I)	+	o	+	o
		dorsolateral sf. of scapula		lateral sf. of scapula		lateral sf. of scapula		lateral sf. of scapula		cranial border of scapula		cranial border of scapula		scapular spine
		anterior sf. of humeral dc crest		anterior sf. of humeral dc crest		along ridge on anterior sf. of humeral dc crest		anterodorsal sf. of proximal humerus		flange along distal 1/2 of humeral dc crest		dorsal sf. of humeral dc crest		humeral dc crest
				clavicle, cranio-lateral sf. of scapula around acromion process		clavicle		dorsal and ventral sf. of interclavicle, ventral 1/3 of proximal clavicle		surrounding acromion		ventral sf. of acromion, clavicle, and lateral process of interclavicle		most of clavicle distal to sternoclavicular joint
<i>m. deltoideus clavicularis</i>	+ ⁴	o	+	+	+	o	+	+	+	(I)	+	o	+	o
		dorsolateral sf. of scapula		anterior sf. of humeral dc crest, distal to m. deltoideus scapularis		along ridge on anterior sf. of humeral dc crest, adjacent to m. deltoideus scapularis		anterodorsal sf. of proximal humerus		humeral dc crest		humeral dc crest		humeral dc crest
		anterior sf. of humeral dc crest												
<i>m. deltoideus acromialis</i>	-		-	-	-		-	-	-	(II)	-	+	+	o
												posterior sf. of acromion		acromion
												anterodorsal sf. of humeral dc crest		humeral dc crest
<i>m. supraspinatus</i>	-		-	-	-		-	-	-	(II)	-	+	+	o
												roughly on cranial scapular base and adjoining procoracoid		supraspinous fossa of scapula
												cranio-medial base of scapula		anterolateral sf. of greater humeral tubercle
												proximal half of greater tubercle of humerus		greater tubercle of humerus
<i>m. infraspinatus</i>	+ ⁵	o	+	+	+	o	+	+	+	(I)	+	o	+	o
		ventrolateral sf. of coracoid		cranial tip of scapulo-coracoid, on both medial and lateral aspects of junction between scapula and coracoid		cranio-lateral sf. of coracoid		lateral sf. of coracoid		lateral sf. of scapula		cranio-lateral sf. of scapula		infraspinous fossa of scapula
		humeral dc crest adjacent to m. pectoralis		apex of humeral dc crest		distal half of greater tubercle of humerus		proximal margin of humeral dc crest and greater tubercle of humerus		distal half of greater tubercle of humerus		greater tubercle of humerus		lateral sf. of greater humeral tubercle
<i>m. teres minor</i>	-		-	-	-		-	-	-	(II)	-	+	+	o
												roughly on lateral scapular base		axillary border of scapula and adjacent infraspinous fossa
												ridge on dorsolateral sf. of proximal humerus		lateral sf. of greater humeral tubercle
<i>m. subcoracoscapularis</i>	+	o	+	+	+	o	+	+	+	(I)	+	o	+	o
		medial sf. of scapulo-coracoid		medial sf. of scapula		axillary border and medial sf. of scapula, medial sf. of coracoid		medial sf. of coracoid and scapula		medial sf. of scapula and coracoid		medial sf. of coracoid and procoracoid, caudolateral sf. of scapula		subscapular fossa on medial sf. of scapula
		humerus		lesser tubercle of humerus		lesser tubercle of humerus		lesser tubercle of humerus		lesser tubercle of humerus		lesser tuberosity of humerus		lesser tubercle of humerus
<i>m. teres major</i>	-		+	-	-		-	-	-	(III)	-	+	+	o
				caudolateral sf. of scapula		lateral sf. and caudolateral tip of coracoid		caudolateral and caudomedial sf. of coracoid		thickened axillary border of scapula		lateral sf. of caudal angle of scapula		caudal angle of scapula and dorsal half of scapular axillary border, sometimes extending onto adjacent infraspinous fossa
				anterodorsal sf. of humerus near m. latissimus dorsi				humeral dc crest, much of anteroventral humeral diaphysis as far as the entepicondyle		ridge on dorsomedial sf. of proximal humerus		near lesser tubercle of humerus		short crest on postero-medial sf. of humeral diaphysis
<i>m. coracobrachialis</i>	+	o	+	+	+	o	+	+	+	(I)	+	+	+	o
		caudolateral sf. of coracoid		cranio-lateral sf. of scapula and coracoid		lateral sf. of coracoid		caudolateral and caudomedial sf. of coracoid		two fossae on cranio-lateral and caudolateral sf. of coracoid		caudolateral border of coracoid		coracoid process
		medial sf. of distal humerus extending to the entepicondyle		ventral sf. of humerus proximal to dc crest		much of ventral sf. of humerus		ventromedial sf. of humeral diaphysis, along medial supracondylar ridge		ventromedial sf. of humeral diaphysis, along medial supracondylar ridge		entire ventral sf. of humerus, extending to base of entepicondyle		postero-medial sf. of humerus between m. subscapularis and m. latissimus dorsi insertions; humerus proximal to entepicondyle, along medial supracondylar ridge
<i>m. biceps brachii</i>	+ ⁶	o	+	+	+	o	+	+	+	(I)	+	+	+	o
		caudolateral sf. of coracoid		ventrolateral sf. of coracoid		lateral sf. of coracoid caudal to m. supracoracoideus origin		lateral sf. of coracoid		lateral sf. of procoracoid; lateral sf. of coracoid tip		ventral sf. of procoracoid and interclavicle; ventral sf. of caudal tip of coracoid		coracoid process; supraglenoid tubercle of scapula
		medial sf. of distal humerus extending to the entepicondyle		posterior sf. of proximal radius, near radial tuberosity		posterior sf. of proximal radius and ulna		radial tuberosity of radius		radial tuberosity of radius		posterior sf. of radius		radial tuberosity of radius
<i>m. triceps brachii</i>	+ ⁷	o	+	+	+	o	+	+	+	(I)	+	+	+	o
		caudolateral sf. of coracoid; ventrolateral sf. of scapula; proximal 2/3 of lateral humeral sf.; entire posterior sf. of humerus		caudolateral sf. of scapula superior to glenoid fossa; caudomedial sf. of scapula and coracoid around glenoid fossa; three additional heads all over humeral diaphysis except ventral area occupied by m. brachialis		caudal coracoid tendon; base of scapula; postero-dorsal sf. of humerus; antero-dorsal sf. of humerus		caudal coracoid tendon; base of scapula; postero-dorsal sf. of humerus; antero-dorsal sf. of humerus		caudomedial sf. of scapula; lateral sf. of humerus; dorso-medial sf. of humerus		lateral sf. of the scapula along crest running dorso-ventrally to the glenoid; postero-dorsal sf. of proximal humerus from lesser tubercle to mid-diaphysis; postero-dorsal sf. of distal humerus spanning from entepicondyle to entepicondyle		ventral portion of scapular axillary border; postero-lateral sf. of proximal humeral diaphysis; postero-medial sf. of distal humeral diaphysis
		olecranon process of ulna		olecranon process of ulna		olecranon process of ulna		olecranon process of ulna		olecranon process of ulna		olecranon process of ulna		olecranon process of ulna

1. As m. dorsohumeralis; 2. As m. pectoralis; 3. As m. dorsalis scapulae; 4. As m. procoracohumeralis; 5. As m. supracoracoideus; 6. As m. coracobrachialis; 7. As m. anconaeus; 8. As m. subscapularis

1124
1125
1126

1127 **FIGURE CAPTIONS**

1128

1129 **Figure 1.** Nodule containing the articulated remains of *Massetognathus pascuali* (MCZVP 3691) (A), with
 1130 lateral (B) and medial (C) views of pectoral limb 3D surface models, prior to mesh refinement and repair.
 1131 MCZVP, Museum of Comparative Zoology, Department of Vertebrate Paleontology, Harvard.

1132

1133 **Figure 2.** Cranial (A), lateral (B), and dorsal (C) views of the articulated left-side pectoral limb of
 1134 *Massetognathus pascuali*, showing rotational axes and primitives used to determine centers of rotation.
 1135 X, Y, and Z axes as labeled in Table 1. Axis colors: X-Red/Y-Green/Z-Blue.

1136

1137 **Figure 3.** Orthographic views of the pectoral limb of *Massetognathus pascuali* in an anatomically-neutral
 1138 reference pose (not in vivo posture), with all joints rotated to the centers of their measured ranges of
 1139 motion. The bones depicted comprise the bilateral scapulocoracoids, humeri, ulnae, and radii, as well as
 1140 the median interclavicle. Line drawing of *M. pascuali* adapted from Figure 9 in Jenkins (1970b).

1141

1142 **Figure 4.** Orthographic views of the left humerus of *Massetognathus pascuali*, with reconstructed muscle
 1143 origins/insertions. Abbreviations: cp, capitulum; dc, deltopectoral crest; ec, ectepicondyle; en,
 1144 entepicondyle; fen, entepicondylar foramen; gt, greater tubercle; h, humeral head; lt, lesser tubercle; th,
 1145 trochlea. Reconstructed muscles are listed in legend.

1146

1147 **Figure 5.** Repaired (A, B) and original (C, D) interclavicle of *Massetognathus pascuali* in ventral (A, C)
 1148 and lateral (B, D) views, with reconstructed muscle origins/insertions. Reference images (E) adapted from
 1149 Jenkins (1970b) (top, *M. pascuali*) and Jenkins (1971a) (bottom left *Thrinaxodon*, bottom right unidentified
 1150 cynodont.) Abbreviations follow Jenkins (1971a): ap, anterior ridge; cc, concavity for clavicle articulation;
 1151 lr, lateral ridge; pp, posterior ridge; pr, posterior ramus. Reconstructed muscles are listed in legend.

1152

1153 **Figure 6.** Orthographic views of the left clavicle of *Massetognathus pascuali*, with reconstructed muscle
 1154 origins/insertions. Abbreviations: ca, concavity for articulation with acromion; cf, clavicular flange; st,
 1155 rugose striations. Reconstructed muscles are listed in legend.

1156

1157 **Figure 7.** Orthographic views of the left scapulocoracoid of *Massetognathus pascuali*, with reconstructed
 1158 muscle origins/insertions. Abbreviations: acr, acromion; axb, axillary border of scapula; c, coracoid
 1159 (=metacoracoid *sensu* Vickaryous & Hall, 2006); cda, caudal angle of scapula; prc, procoracoid foramen;
 1160 g, glenoid fossa; prc, procoracoid; rcb, reflected cranial border of scapula; sb, scapular base; sc, scapula;
 1161 vb, vertebral border of scapula. Reconstructed muscles are listed in legend. Muscles are color-coded for
 1162 visual differentiation, not homology.

1163

1164 **Figure 8.** Orthographic views of the left radius (A) and ulna (B) of *Massetognathus pascuali*, with
 1165 reconstructed muscle origins/insertions. Note that the orientations for the radius are slightly rotated from
 1166 Jenkins (1971a). Abbreviations: pr ar f, proximal articular facet; rt, radial tuberosity; rd nt, radial notch.
 1167 Reconstructed muscles are listed in legend.

1168

1169 **Figure 9.** Potential pectoral limb excursion of *Massetognathus pascuali* with fixed (A) and mobile (B)
 1170 clavo-interclavicular and acromio-clavicular articulations. Neutral pose in white. Joint angles (protracted):
 1171 [clavicle rotated medially 15°; scapulocoracoid yawed laterally 15°, rolled laterally 5°, pitched caudally 5°];
 1172 humerus adducted 20°, supinated 15°, protracted 15°; radius and ulna flexed 45°. Joint angles (retracted):
 1173 [clavicle rotated laterally 15°; scapulocoracoid rolled medially 5°, pitched cranially 30°]; humerus pronated
 1174 10°, retracted 15°; radius and ulna extended 45°.

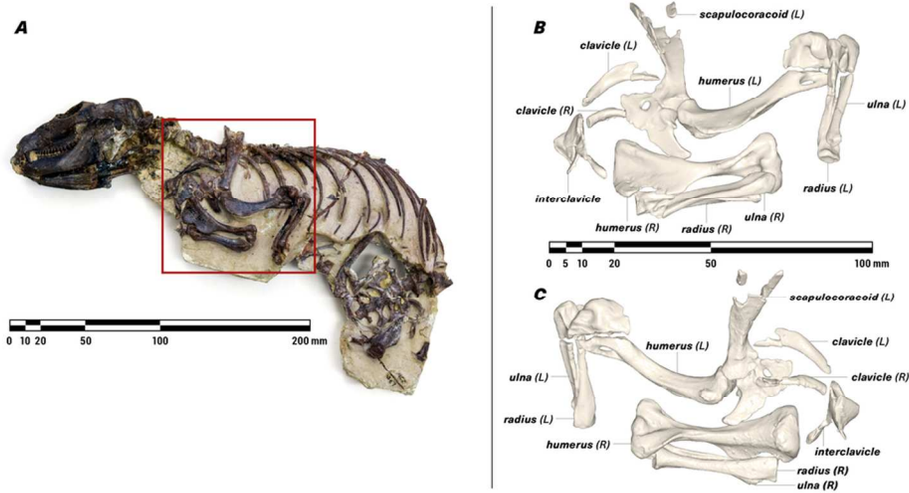
1175

1176 **Figure 10.** Close-up caudal (A), caudolateral (B) and ventrocaudolateral (C) views of the glenoid fossa of
 1177 *Massetognathus pascuali*, showing the convex coracoid facet and the concave scapular facet.

1178

1179

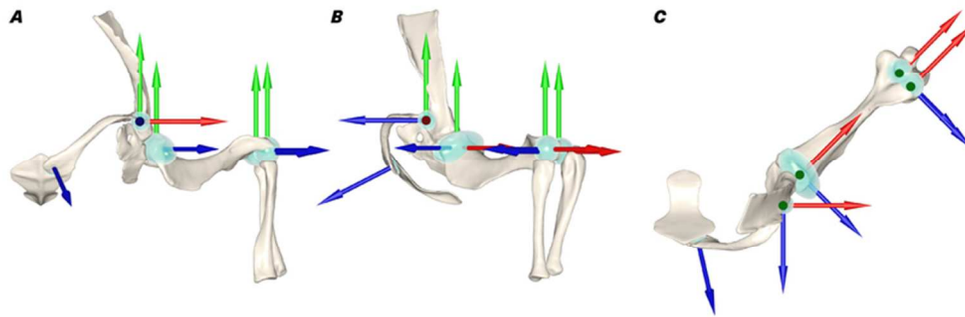
For Peer Review Only



Nodule containing the articulated remains of *Massetognathus pascuali* (MCZVP 3691) (A), with lateral (B) and medial (C) views of pectoral limb 3D surface models, prior to mesh refinement and repair. MCZVP, Museum of Comparative Zoology, Department of Vertebrate Paleontology, Harvard.

100x50mm (300 x 300 DPI)

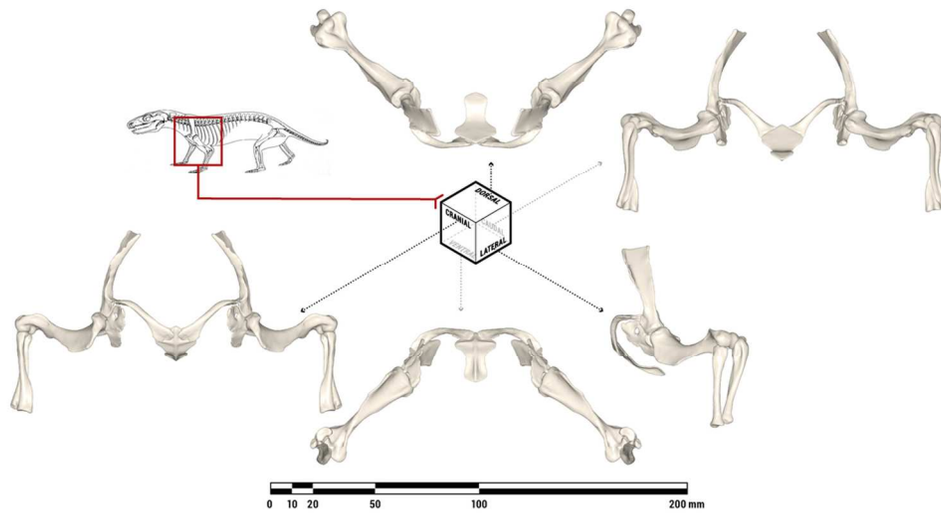
Review Only



Cranial (A), lateral (B), and dorsal (C) views of the articulated left-side pectoral limb of *Massetognathus pascuali*, showing rotational axes and primitives used to determine centers of rotation. X, Y, and Z axes as labeled in Table 1. Axis colors: X-Red/Y-Green/Z-Blue.

70x24mm (300 x 300 DPI)

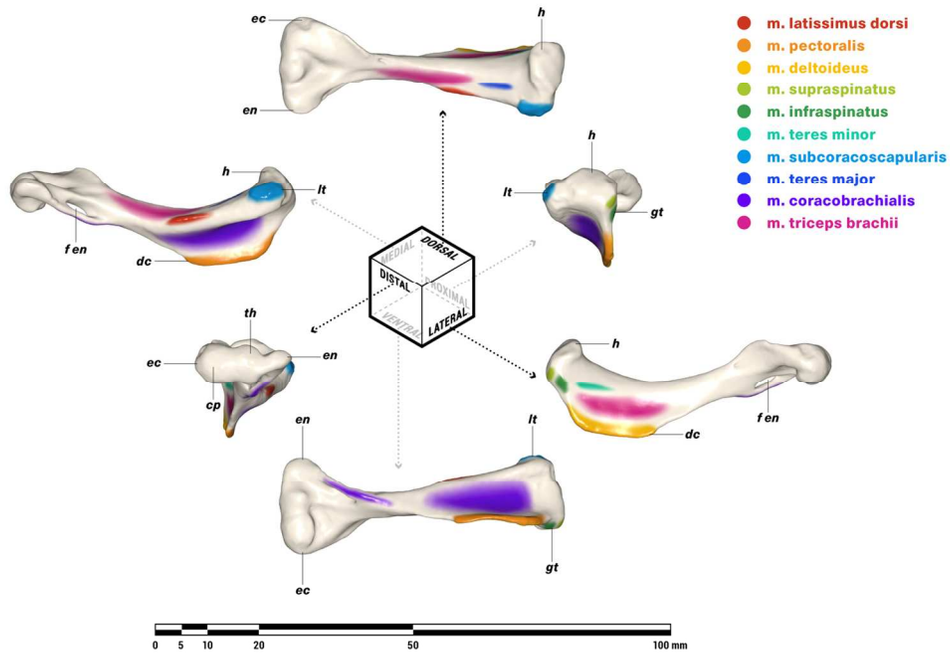
Peer Review Only



Orthographic views of the pectoral limb of *Massetognathus pascuali* in an anatomically-neutral reference pose (not in vivo posture), with all joints rotated to the centers of their measured ranges of motion. The bones depicted comprise the bilateral scapulocoracoids, humeri, ulnae, and radii, as well as the median interclavicle. Line drawing of *M. pascuali* adapted from Figure 9 in Jenkins (1970b).

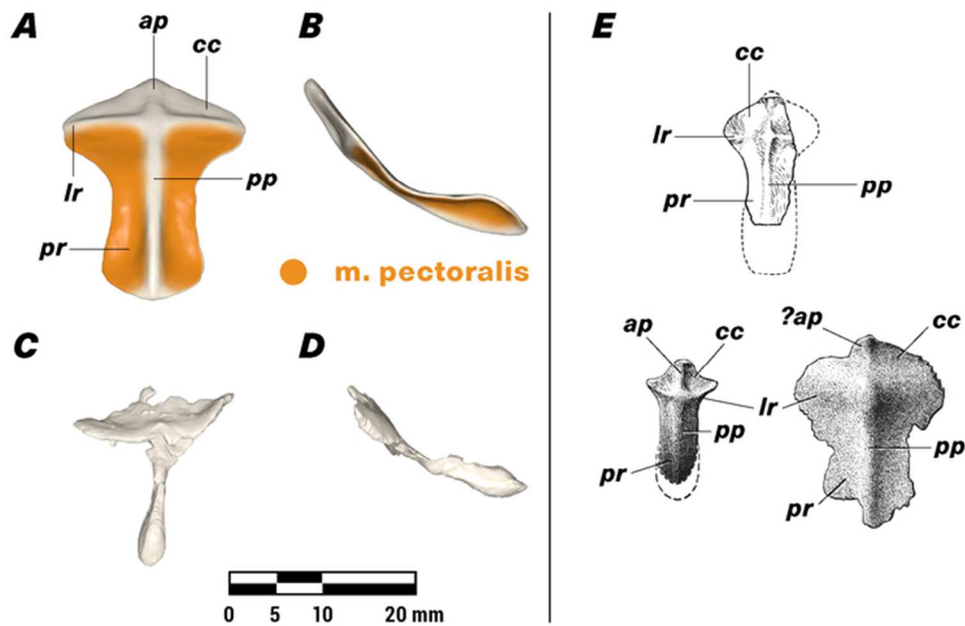
110x60mm (300 x 300 DPI)

Review Only



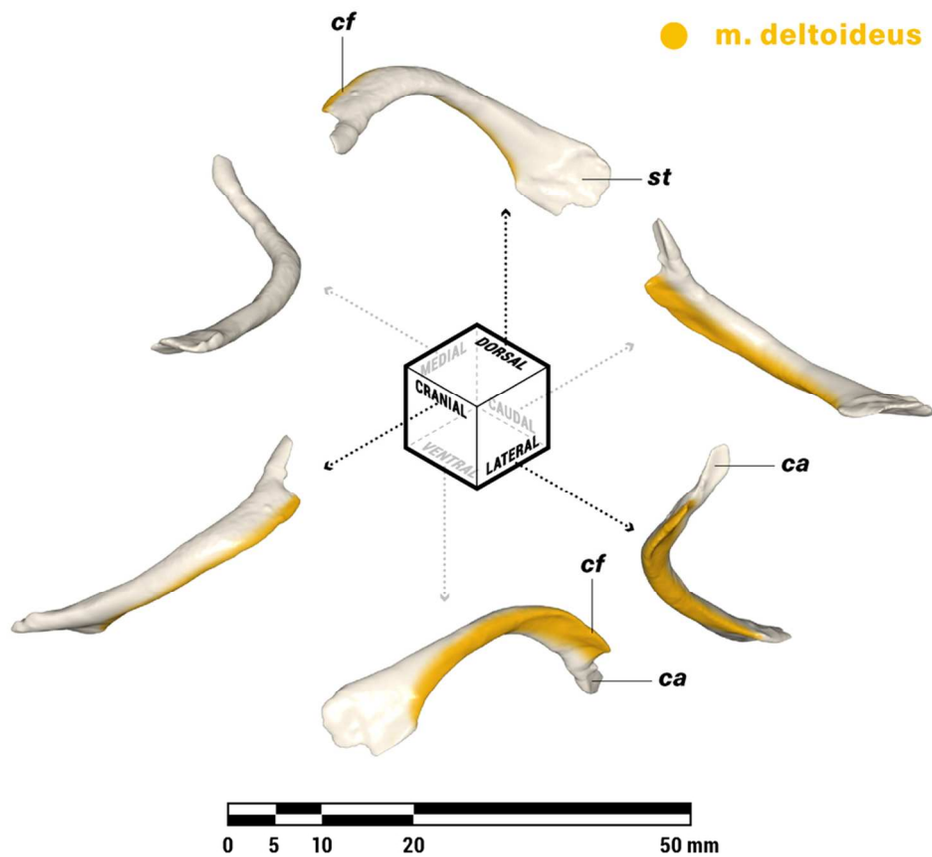
Orthographic views of the left humerus of *Massetognathus pascuali*, with reconstructed muscle origins/insertions. Abbreviations: cp, capitulum; dc, deltopectoral crest; ec, ectepicondyle; en, entepicondyle; fen, entepicondylar foramen; gt, greater tubercle; h, humeral head; lt, lesser tubercle; th, trochlea. Reconstructed muscles are listed in legend.

140x98mm (300 x 300 DPI)



Repaired (A, B) and original (C, D) interclavicle of *Massetognathus pascuali* in ventral (A, C) and lateral (B, D) views, with reconstructed muscle origins/insertions. Reference images (E) adapted from Jenkins (1970b) (top, *M. pascuali*) and Jenkins (1971a) (bottom left *Thrinaxodon*, bottom right unidentified cynodont.) Abbreviations follow Jenkins (1971a): ap, anterior ridge; cc, concavity for clavicle articulation; lr, lateral ridge; pp, posterior ridge; pr, posterior ramus. Reconstructed muscles are listed in legend.

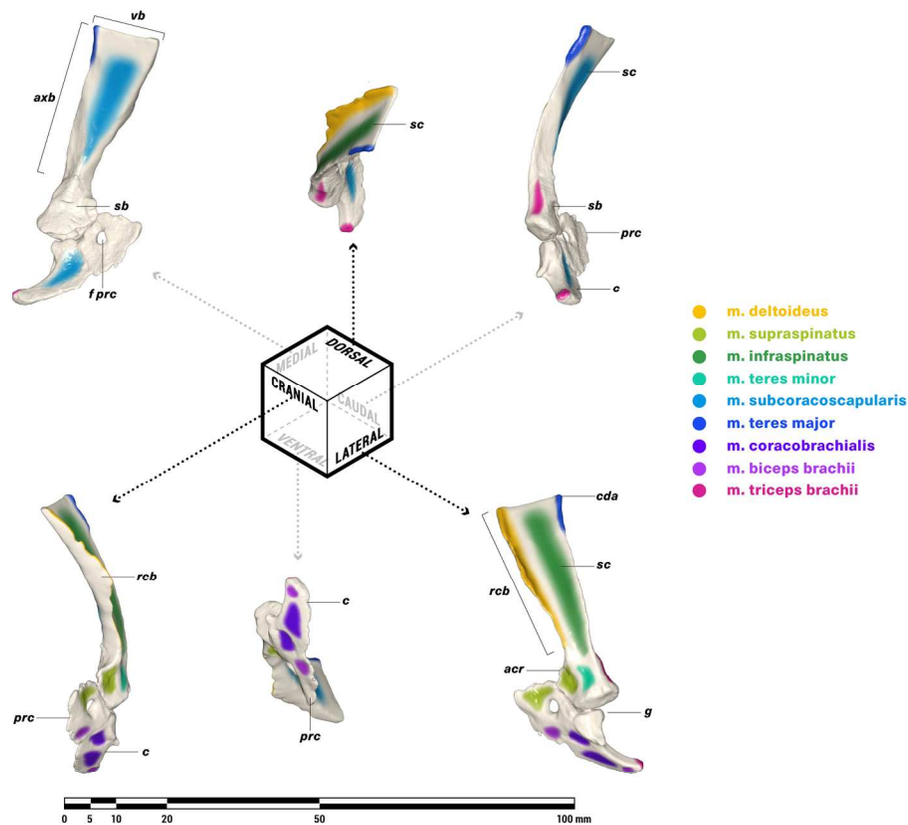
65x42mm (300 x 300 DPI)



Orthographic views of the left clavicle of *Massetognathus pascuali*, with reconstructed muscle origins/insertions. Abbreviations: ca, concavity for articulation with acromion; cf, clavicular flange; st, rugose striations. Reconstructed muscles are listed in legend.

90x80mm (300 x 300 DPI)

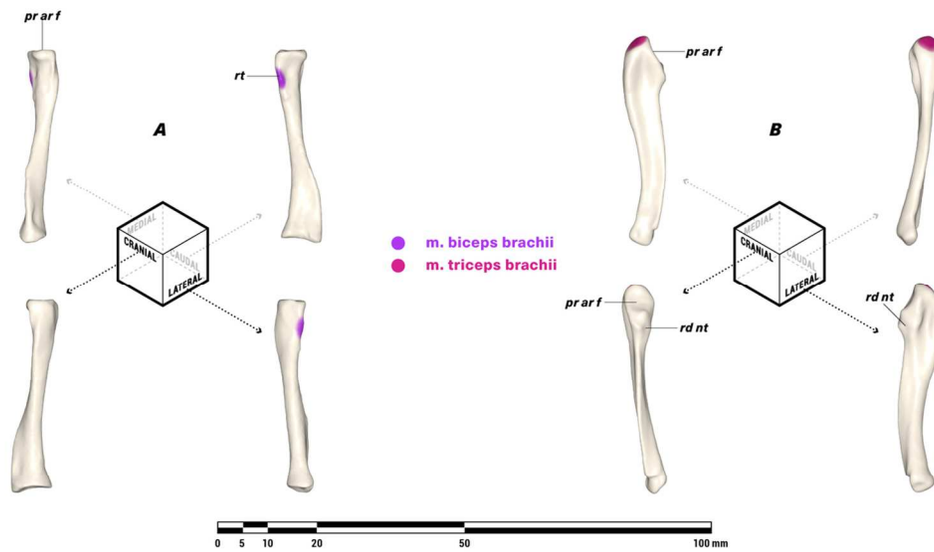
Only



Orthographic views of the left scapulocoracoid of *Massetognathus pascuali*, with reconstructed muscle origins/insertions. Abbreviations: acr, acromion; axb, axillary border of scapula; c, coracoid (=metacoracoid sensu Vickaryous & Hall, 2006); cda, caudal angle of scapula; prc, procoracoid foramen; g, glenoid fossa; prc, procoracoid; rcb, reflected cranial border of scapula; sb, scapular base; sc, scapula; vb, vertebral border of scapula. Reconstructed muscles are listed in legend. Muscles are color-coded for visual differentiation, not homology.

175x153mm (300 x 300 DPI)

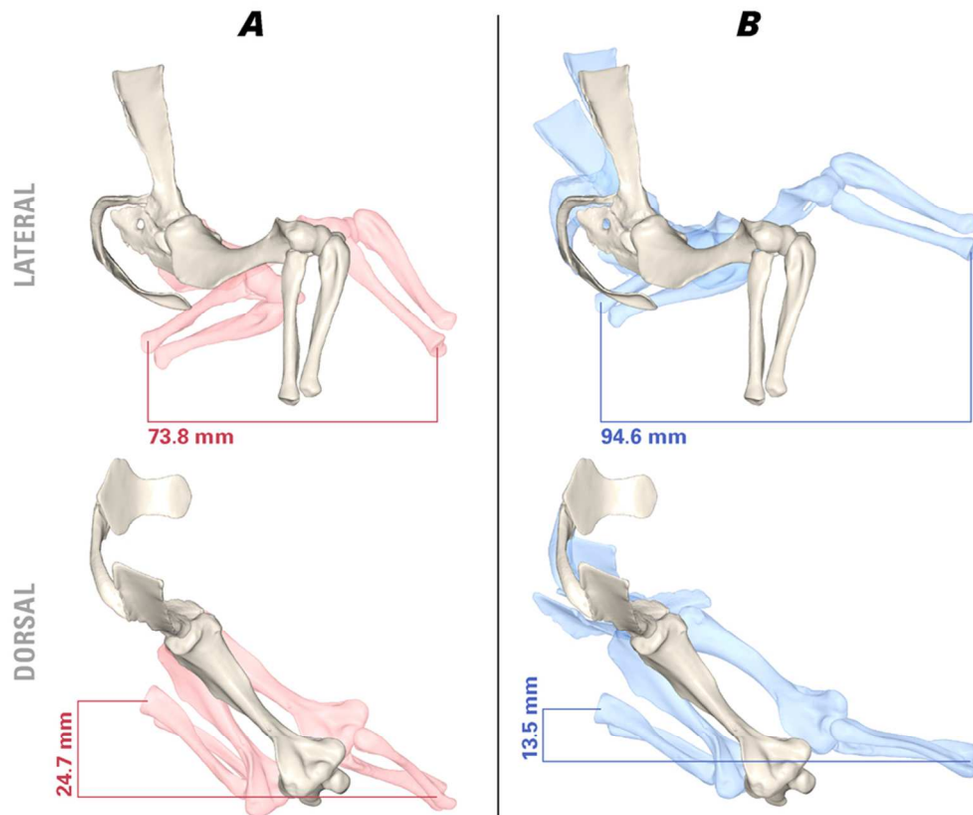




Orthographic views of the left radius (A) and ulna (B) of *Massetognathus pascuali*, with reconstructed muscle origins/insertions. Note that the orientations for the radius are slightly rotated from Jenkins (1971a). Abbreviations: prarf, proximal articular facet; rt, radial tuberosity; rdnt, radial notch. Reconstructed muscles are listed in legend.

115x66mm (300 x 300 DPI)

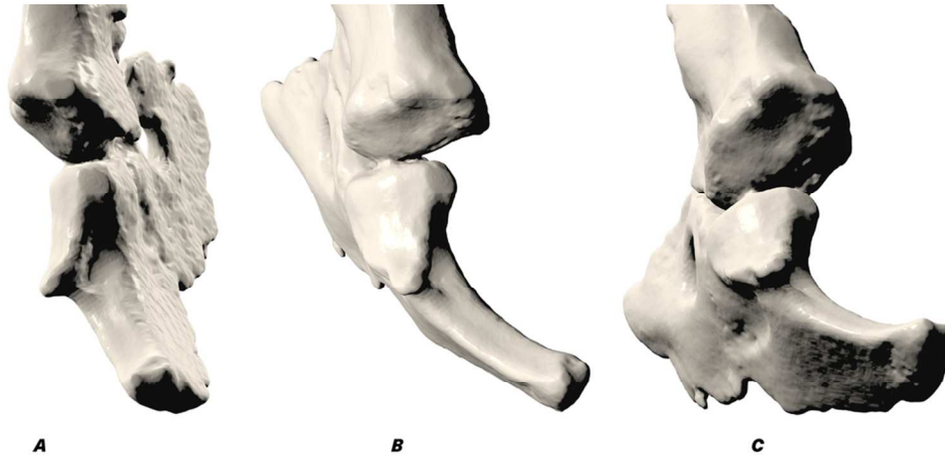
Review Only



Potential pectoral limb excursion of *Massetognathus pascuali* with fixed (A) and mobile (B) clavo-interclavicular and acromio-clavicular articulations. Neutral pose in white. Joint angles (protracted): [clavicle rotated medially 15°; scapulocoracoid yawed laterally 15°, rolled laterally 5°, pitched caudally 5°]; humerus adducted 20°, supinated 15°, protracted 15°; radius and ulna flexed 45°. Joint angles (retracted): [clavicle rotated laterally 15°; scapulocoracoid rolled medially 5°, pitched cranially 30°]; humerus pronated 10°, retracted 15°; radius and ulna extended 45°.

85x72mm (300 x 300 DPI)

AMN



Close-up caudal (A), caudolateral (B) and ventrocaudolateral (C) views of the glenoid fossa of *Massetognathus pascuali*, showing the convex coracoid facet and the concave scapular facet.

95x45mm (300 x 300 DPI)

Review Only