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1 **New material of *Chirostenotes pergracilis* (Theropoda, Oviraptorosauria) from the**  
2 **Campanian Dinosaur Park Formation of Alberta, Canada**

3

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12 **New material of *Chirostenotes pergracilis* (Theropoda, Oviraptorosauria) from the**  
13 **Campanian Dinosaur Park Formation of Alberta, Canada**

14

15 **Abstract:**

16 The taxonomy of caenagnathids from the Dinosaur Park Formation of Alberta, Canada, has  
17 remained problematic because of incomplete, partial skeletons that do not overlap anatomically.  
18 This is particularly problematic for referring mandibular remains, which are the most abundant  
19 caenagnathid fossils recovered, but cannot be confidently tied to taxa known from postcranial  
20 remains. A new, partial skeleton of *Chirostenotes pergracilis* preserves the mandibles, cervical  
21 and caudal vertebrae, and parts of the hindlimb. Importantly, this is the first specimen with  
22 associated mandibles and postcrania of a caenagnathid from the Dinosaur Park Formation,  
23 allowing for unambiguous referral of mandibles to this taxon. The mandibles are remarkably  
24 similar to those previously suggested to pertain to *Chirostenotes pergracilis*, and support its  
25 distinction from *Caenagnathus collinsi*. An unfused distal tarsal IV distinguishes the skeleton  
26 from *Leptorhynchus elegans* and supports the referral of small, upturned mandibles to this taxon.  
27 Osteohistological analysis indicates that the individual was approaching maximum body size,  
28 and provides information on the growth patterns and size of *Chirostenotes pergracilis*.  
29 Accordingly, this supports the division of Dinosaur Park Formation caenagnathids into three taxa  
30 of varying body sizes.

31 **Keywords:**

32 Caenagnathidae; Oviraptorosauria; Late Cretaceous; Osteohistology; Dinosaur Park Formation.

33 **Introduction:**

34

35 Caenagnathidae is a clade of maniraptoran theropods from the Cretaceous of Asia and North  
36 America. The first known caenagnathid, *Chirostenotes pergracilis*, was named by Gilmore  
37 (1924) based on an articulated manus from the Upper Cretaceous (Campanian) Dinosaur Park  
38 Formation (DPF) of Alberta, Canada. Other caenagnathid material was described, mostly as new  
39 taxa, by C.M. Sternberg (1932; 1934), Parks (1933), and R.M. Sternberg (1940), although it was  
40 not clear until much later that this material pertained to the same group as *Chirostenotes*  
41 *pergracilis* (Osmólska 1976; Osmólska 1981; Currie 1989). A partial skeleton of *Chirostenotes*  
42 *pergracilis* including most of the hindlimb was described by Currie and Russell (1988). Its  
43 metatarsal structure suggested to Currie and Russell (1988) that “*Macrophalangia canadensis*”  
44 (Sternberg 1932) was the junior synonym of *Chirostenotes pergracilis*. Currie (1989) noticed  
45 similarities between the small, fused tarsometatarsus of ‘*Ornithomimus*’ *elegans* (Parks 1933)  
46 and those of *Elmisaurus rarus* (Osmólska 1981) from Mongolia. He suggested they were  
47 congeneric, and that *Elmisaurus elegans* and *Chirostenotes pergracilis* could be distinguished by  
48 fusion of the tarsometatarsus (Currie 1989). However, this was complicated by a partial skeleton  
49 (ROM 43250) from the Horseshoe Canyon Formation of Alberta, described by Sues (1997).  
50 Although the specimen cemented the synonymy of Caenagnathidae and Elmisauridae (Sues  
51 1997), it stimulated debate about which specimens were conspecific. Whereas Currie (1989)  
52 preferred to separate DPF caenagnathid material from into three genera: *Caenagnathus*,  
53 *Chirostenotes*, and *Elmisaurus*, Sues (1997) argued that all of this material should be united.  
54 Later, Sullivan et al. (2011) suggested, on the basis of stratigraphic separation, that the material  
55 described by Sues (1997) was distinct enough to merit a new name: *Epichirostenotes curriei*.



56 Taxonomic separation of ROM 43250 from *Chirostenotes pergracilis* called into question the  
57 assertion of Sues (1997) that *Caenagnathus collinsi*, *Chirostenotes pergracilis*, and *Elmisaurus*  
58 *elegans* were synonymous. Accordingly, most studies now follow Currie (1989) in the separation  
59 of DPF caenagnathids into three taxa (Longrich et al. 2013; Lamanna et al. 2014; Funston &  
60 Currie 2016).

61 In the last fifteen years, a number of new discoveries have ameliorated our understanding  
62 of the diversity and anatomy of caenagnathids, especially in Asia. Xu et al. (2007) described the  
63 appropriately named *Gigantoraptor erlianensis* on the basis of a giant skeleton from China with  
64 clear oviraptorosaur affinities. Subsequent cladistic analyses (Lamanna et al. 2014; Funston &  
65 Currie 2016) have placed *Gigantoraptor* as a basal caenagnathid, mostly on the basis of the  
66 mandible. Ma et al. (2017) further described the mandible of *Gigantoraptor* in more detail, and  
67 assessed the functional morphology of its intermediate shape. Sues and Averianov (2015)  
68 described additional material of the miniscule *Caenagnathasia martinsoni*, including vertebrae  
69 and a femur. Yao et al. (2015) described another partial mandible of *Caenagnathasia*, but from  
70 the Iren Dabasu Formation of China, expanding the stratigraphic and geographic range of  
71 *Caenagnathasia*. Tsuihiji et al. (2015) described a pair of giant mandibles from the Bayn Shiree  
72 Formation of Mongolia, noting similarities to *Gigantoraptor erlianensis*. The same authors then  
73 described much smaller, fused dentaries probably referable to *Elmisaurus rarus* from the Nemegt  
74 Formation exposed at Bugiin Tsav (Tsuihiji et al. 2016). Pu et al. (2017) published the long-  
75 awaited description of ‘Baby Louie’ from Henan Province in China, interpreting it as a new,  
76 giant caenagnathid closely related to *Gigantoraptor erlianensis*. Recently, Yu et al. (2018)  
77 named an intermediately-sized caenagnathid, *Anomalipes zhaoui*, from the Late Cretaceous of  
78 China, filling the former gap in body sizes of Chinese caenagnathids between *Caenagnathasia*

79 and *Gigantoraptor*. Wang et al. (2017; 2018) argued, based on tooth-loss patterns in a Jurassic  
80 ceratosaur and the Early Cretaceous bird *Sapeornis*, that caenagnathids underwent ontogenetic  
81 edentulism. On this basis, they argued that the complex structures on the occlusal surfaces of the  
82 dentaries were the vestiges of tooth-bearing structures. However, this finding was contested by  
83 Funston et al. (2019), who used osteohistology to show that caenagnathid dentaries lacked any  
84 evidence of tooth-bearing tissues. These studies have shown that caenagnathids were diverse and  
85 broadly distributed in the Late Cretaceous, but they have done little to clarify the taxonomy or  
86 relationships of the group.

87         However, better material from North America has provided some advances on this front.  
88 Lamanna et al. (2014) described *Anzu wyliei*, the largest caenagnathid known from North  
89 America, from three reasonably complete skeletons from the Upper Cretaceous (Maastrichtian)  
90 Hell Creek Formation. These specimens show that *Anzu* had a prominent cranial crest and a short  
91 tail with modified pygostyle-like distal vertebrae. Importantly, these specimens also provide  
92 concrete evidence that caenagnathid mandibles and postcrania pertain to the same animals.  
93 Funston and Currie (2016) described a relatively complete skeleton of a new taxon, *Apatoraptor*  
94 *pennatus*, from the Horseshoe Canyon Formation, which helped to resolve some aspects of  
95 caenagnathid phylogeny. Although less complete, numerous new caenagnathid specimens have  
96 also been described from the DPF (Longrich et al. 2013; Funston & Currie 2014; Bell et al.  
97 2015; Funston et al. 2015; Funston, Currie, & Burns 2016; Gregory F. Funston et al. 2019), and  
98 these help to improve our understanding of the anatomy and variation in caenagnathids.

99         Regardless of this considerable progress, the taxonomy of DPF caenagnathids has  
100 remained obscure and unresolved. This is due in large part to the fragmentary, non-overlapping  
101 nature of partial skeletons known thus far from the DPF—in spite of producing the greatest

102 abundance of caenagnathid specimens globally. Nonetheless, this issue remains central to  
103 unravelling the relationships and ecology of caenagnathids. Confident referral of isolated DPF  
104 specimens to their respective taxa would greatly improve skeletal representation and  
105 phylogenetic character scores.

106         In 2016, a partial caenagnathid skeleton (UALVP 59400) was found in the southeastern  
107 part of Dinosaur Provincial Park (Fig. 1). The specimen comprises a relatively complete  
108 mandible and elements of the axial and appendicular skeleton, providing the first associated  
109 mandibular and postcranial material from the DPF. The morphology of the mandible and a distal  
110 tarsal IV indicate that this specimen is attributable to *Chirosstenotes pergracilis*. In addition,  
111 reexamination suggests that some previously collected material can also be referred to  
112 *Chirosstenotes pergracilis* on the basis of overlap with known specimens. These specimens are  
113 described here with osteohistological analyses to assess relative maturity of the individual  
114 represented by UALVP 59400. The new material helps to resolve some of the ambiguities  
115 regarding taxonomy and diversity of DPF caenagnathids, and provides some insights into the  
116 growth patterns of *Chirosstenotes pergracilis*.

117 ***Institutional Abbreviations:***

118 **CMN**, Canadian Museum of Nature, Ottawa, ON, Canada; **TMP**, Royal Tyrrell Museum of  
119 Palaeontology, Drumheller, AB, Canada; **UALVP**, University of Alberta Laboratory for  
120 Vertebrate Palaeontology, Edmonton, AB, Canada.

121

122 **Materials and Methods:**

123

124 The material was excavated under provincial and parks collecting permits to PJC (UALVP  
125 59400) or the TMP. UALVP 59400 consisted of several badly crushed blocks that developed  
126 fracture planes through the bones. Because of the delicate nature of the specimen, the unprepared  
127 blocks were CT scanned using a Siemens Sensation 64 Medical CT scanner and visualized using  
128 Mimics 14.0 to guide preparation. The blocks were then mechanically prepared using  
129 conventional methods and photographed using a Nikon D7200 digital camera. Photographs were  
130 processed using Adobe Photoshop CC or Adobe Photoshop 2020. Any adjustments made to  
131 brightness, contrast, or colour balance were applied to the whole image. Osteohistological thin-  
132 sections of UALVP 59400 were made by vacuum-embedding a fragment of an indeterminate  
133 long bone in Castolite AC polyester resin, and cutting the billet using an Isomet 1000 Precision  
134 Sectioning Saw. Billets were adhered to plexiglass slides using 3M Cyanoacrylate glue. The  
135 mounted billet was resectioned to a thickness of 0.7mm, and then ground and polished by hand  
136 using 600-grit powder on a glass plate. The slides were polished on a buffing pad and mineral oil  
137 was applied to enhance optical clarity. Slides were imaged under plane polarized and cross-  
138 polarized light using NIS Elements on a Nikon Eclipse E600POL trinocular polarizing  
139 microscope with an attached Nikon DXM 1200F digital camera. For enhanced clarity and depth  
140 of field, some pictures (both microscopic and macroscopic) were generated using Z-stacked  
141 images. These were created manually, using either NIS Elements or Adobe Photoshop 2020.

142

143 **Systematic Palaeontology:**

144

145 DINOSAURIA Owen, (1842)

146 SAURISCHIA Seeley, (1888)

147 THEROPODA Marsh, (1881)  
148 COELUROSAURIA von Huene, (1914)  
149 MANIRAPTORA Gauthier, (1986)  
150 OVIRAPTOROSAURIA Barsbold, (1976)  
151 CAENAGNATHIDAE R. M. Sternberg, (1940)  
152 *CHIROSTENOTES PERGRACILIS* Gilmore, (1924)  
153 ***Holotype:***  
154 CMN 2367, articulated manus (Gilmore, 1924)  
155 ***Referred Material:***  
156 CMN 8538, right pes (Sternberg 1932); TMP 1979.020.0001, partial skeleton (Currie and  
157 Russell 1988); TMP 1985.043.0070, partial dentaries (Gregory F. Funston et al. 2019); TMP  
158 1992.036.1237 (Gregory F. Funston et al. 2019); TMP 2001.012.0012, complete mandibles  
159 (Funston & Currie 2014).  
160 ***Newly Referred Material:***  
161 TMP 1990.056.0006, fused dentaries; TMP 1996.036.0181, partial tarsometatarsus (Funston,  
162 Currie, & Burns 2016); TMP 2002.012.0103, partial ilium; UALVP 59400, partial skeleton  
163 (GPS: UTM 12U 468540, 5621530).  
164 ***Locality and Horizon:***  
165 Dinosaur Park Formation (Campanian), Dinosaur Provincial Park, Alberta, Canada (Fig. 1).  
166 ***Revised Diagnosis (modified from Currie and Russell 1988 and Longrich et al. 2013):***  
167 Medium-sized (~65 kg) caenagnathid oviraptorosaur diagnosed by the following autapomorphies  
168 (\*) and combination of characters: occlusal tip of dentary upturned at approximately 45°\*;  
169 dentaries fused with well-developed symphyseal shelf; deep mandibular fossa; dentary excluded

170 from dorsal margin of external mandibular fenestra by surangular; articular ridge of mandible  
171 distinctly offset from dorsal margin of surangular; cervical vertebrae with low neural spines and  
172 small epiphyses; six sacral vertebrae with pleurocoels; distal caudal vertebrae with anteriorly-  
173 directed transverse processes; posterior chevrons anteroposteriorly elongate at proximal end, as  
174 long or longer anteroposteriorly than corresponding caudal vertebrae\*; digit III of manus longer  
175 than digit I, but with slender phalanges; tall, dolichoiliac ilium with reduced postacetabular  
176 blade\*; distal tarsals and proximal metatarsals not coossified at maturity; metatarsal III  
177 proximally pinched between metatarsals II and IV, but only the proximal tip is excluded from the  
178 anterior surface of the metatarsus; metatarsal V strongly procurving and not fused to distal tarsal  
179 IV\*.

180 **Description:**

181

182 ***UALVP 59400:***

183 UALVP 59400 (Table 1) consists of a partial articulated skeleton including the mandible, four  
184 cervical vertebrae, eleven caudal vertebrae and associated chevrons, a partial pubis and ischium,  
185 fragments of a femur and tibia, both astragali, and a right distal tarsal IV (Fig. 1C). The bones are  
186 transversely crushed but otherwise relatively well preserved.

187 ***Mandible:***

188 The mandible (Fig. 2) is relatively complete, consisting of badly crushed, fused dentaries and  
189 portions of both articular-surangular-coronoid (ASC) complexes. The angulars are preserved as a  
190 collection of fragments that likely represent a significant proportion of the bones, but cannot be  
191 reconstructed. The left dentary is more complete and can be rearticulated with the corresponding  
192 ASC complex (Fig. 2A–D), which is less complete than that of the right side (Fig. 2E–J). The

193 mandible is overall remarkably similar to TMP 2001.012.0012 in morphology, but it is slightly  
194 smaller in size. The dentary is transversely crushed but still preserves much of the morphology.  
195 The anterior occlusal grooves and ridges are shallow and lack nodules, in contrast to TMP  
196 1992.036.0390. There are four lateral occlusal ridges (Fig. 2C, D), which become successively  
197 smaller posteriorly. They border a deep lingual groove, which itself lies lateral to the well-  
198 developed lingual ridge. The tubercle of the lingual ridge is distorted by crushing, but it does not  
199 appear to have been as prominent as that of TMP 1992.036.1237, nor does it have the small  
200 nodules present in the latter. The symphyseal sulcus is mostly missing. The lateral surface of the  
201 dentary is pierced by numerous foramina (Fig. 2A, B), which, as in TMP 2001.012.0012, are  
202 arranged into three rows (Funston and Currie 2014). A mandibular fossa is present and  
203 apparently deep, but it is deformed by transverse and dorsoventral crushing. The ventral surface  
204 of the dentary is only preserved at its posterior end, near the attachment of *M. genioglossus*. This  
205 scar is roughly heart-shaped and foramina demarcate its posterior border. The Meckelian grooves  
206 converge towards the midline, and open posteroventrally on the posteromedial surface of the  
207 dentary. The rami of the dentaries bifurcate around a large external mandibular fenestra. The  
208 dorsal ramus is strap-like and forms an interfingering joint with the ASC complex (Currie et al.  
209 1993; Funston & Currie 2014). The ventral ramus tapers posteriorly and lies lateral to the  
210 angular, which has a deep groove for this contact. The ASC complex has a rugose, medially  
211 deflected coronoid process, which, as in TMP 2001.012.0012 (Funston & Currie 2014), has a  
212 distinct ventral suture (Fig. 2E, F). A small foramen pierces the medial surface of the surangular  
213 near its contact with the angular, but this does not form a fenestra in the way that it does in  
214 oviraptorids. The articular has a tall median ridge that presumably was flanked by tongue-like  
215 medial and lateral cotyles as in TMP 2001.012.0012, although these are missing in UALVP

216 59400. The slope of the anterior part of the articular ridge, where it meets the surangular, is less  
217 steep than in TMP 2001.012.0012, but it is more pronounced than in *Anzu wyliei* (Lamanna et al.  
218 2014) and *Caenagnathus collinsi* (Currie et al. 1993). It is most similar to that of *Aptoraptor*  
219 *pennatus* (Funston and Currie 2016).

220 *Cervical Vertebrae:*

221 Four cervical vertebrae are preserved, three of which are still in articulation (Fig. 3). Two mid-  
222 cervical vertebrae, probably representing postaxial cervical vertebrae 7–8 based on comparison  
223 to *Aptoraptor pennatus*, are better preserved. The morphologies of their centra are difficult to  
224 discern, but the neural arches are well preserved. The neurocentral suture is obliterated in all of  
225 the vertebrae. The centrum of the anterior vertebra appears to have a longitudinal ridge  
226 underlying the infrapostzygapophyseal fossa on the neural arch (Fig. 3). Ventral to this, there is a  
227 large lateral pleurocoel that opens posteriorly. The diapophysis is connected to the  
228 postzygapophysis by a broad lamina that forms the dorsal edge of the infrapostzygapophyseal  
229 fossa. The neural spine is dorsoventrally short and rounded in lateral view (Fig. 3). The  
230 postzygapophysis faces ventrally and the epipophysis is small or absent. The posterior cervical  
231 vertebra preserves the entire neural arch, but it is more damaged than the anterior one. The  
232 prezygapophysis is upturned, which may be taphonomically modified, although it also appears to  
233 be the case in postaxial cervical vertebra eight of *Aptoraptor pennatus* (Funston and Currie  
234 2016). The neural spine is low and rounded. The postzygapophyses face ventrally and also seem  
235 to lack epipophyses. Like the anterior vertebra, the neural arch has a broad lamina that connects  
236 the postzygapophysis to the diapophysis. The diapophysis is apparently fused to the cervical rib,  
237 although this region is damaged and the fusion cannot be determined without doubt. Dorsal to  
238 the two better preserved cervical vertebrae is a patch of matrix with filamentous stains (Fig. 3).



239 There is a distinct ‘tufted’ border between these stains and the matrix further away from the  
240 specimen, and the filaments appear to have a consistent orientation extending posterodorsally  
241 from the vertebrae. Most filaments preserved in this region are approximately 1 mm long and  
242 vary considerably in thickness. However, many of these filaments appear to form continuous,  
243 parallel lines that can be as long as 1 cm (Fig. 3C–E). The colouration of the filaments varies  
244 from black to dark brown to a rusty orange or red colour. Filaments of these various colours are  
245 interspersed in the same regions, creating a speckled pattern (Fig. 3C). The most posterior patch  
246 of impressions (Fig. 3E) has the highest proportion of black filaments, and these are generally  
247 arranged into parallel rows of smaller filamentous specks. Two predominant orientations of these  
248 rows can be discerned, one extending posterodorsally, and one extending anteroposteriorly.  
249 Some possible branching structures can be discerned close to the vertebra (Fig. 3E), but these  
250 may simply be patchy preservation of the filamentous stains.

251 *Caudal Vertebrae:*

252 A series of thirteen articulated caudal vertebrae (Figs 4, 5) were preserved with the skeleton, of  
253 which eleven remained in one block. The caudal vertebrae likely represent the penultimate part  
254 of the tail, and do not include the pygostyle. This is evident from the positions of the neural  
255 spines posterior to the articular faces of the corresponding vertebrae and the anteriorly directed  
256 transverse processes on the most posterior caudal vertebrae (Fig. 5). The more anterior caudal  
257 vertebrae (Fig. 4) have barrel-shaped centra with large lateral pleurocoels, which decrease in size  
258 posteriorly along the tail (Table 1). In the posterior caudal vertebrae, these pleurocoels are slit-  
259 like, underlying the infradiapophyseal fossa, and they are absent in the last three caudal  
260 vertebrae. The neural spines are low and triangular in lateral view. The transverse processes are  
261 elliptical in cross section, tongue-like in dorsal view, and extend posterolaterally in the anterior

262 caudal vertebrae (Fig. 4). Posteriorly, the transverse processes become more platelike, taper at  
263 their lateral ends, and become successively more anteriorly directed. In the anterior caudal  
264 vertebrae, there is a well developed infradiapophyseal fossa and a smaller infraprezygapophyseal  
265 fossa, but the presence of an infrapostzygapophyseal fossa cannot be determined because this  
266 area is overlain by the fingerlike prezygapophyses. The last two caudal vertebrae are slightly  
267 disarticulated and lie nearly perpendicular to the rest of the series (Figs. 4, 5). The centra of these  
268 vertebrae are transversely wider than they are dorsoventrally tall, and they have a groove along  
269 the ventral midline (Fig. 5). The transverse processes extend from the widest point of the  
270 centrum and curve anteriorly from their bases. In these respects, they are similar to the  
271 penultimate caudal vertebrae of *Nomingia gobiensis* (RZNCHEN Barsbold et al. 2000), and they  
272 likely represent the corresponding vertebrae. Specifically, the last caudal vertebra preserved with  
273 UALVP 59400 is nearly identical to caudal vertebra 17 of *Nomingia gobiensis* (Rinchen  
274 Barsbold et al. 2000). If this is the case, only four additional vertebrae would be missing from the  
275 anterior part of the caudal vertebral series, and seven from the posterior part of the tail.

276 *Chevrons:*

277 The chevrons (Fig. 6) are exceptionally large compared to other oviraptorosaurs, especially  
278 considering the distal positions of the corresponding caudal vertebrae. Anteriorly, the chevrons  
279 are elongate and taper towards their rounded distal ends. Posteriorly, they become dorsoventrally  
280 shorter and more platelike (Fig. 4), but do not decrease in anteroposterior length. The result is  
281 that the proximal ends of the posterior chevrons are nearly in contact, and the most posterior  
282 chevrons are longer anteroposteriorly than their corresponding vertebrae. A similar morphology  
283 is present in *Nomingia gobiensis* (RZNCHEN Barsbold et al. 2000), but not to the same extreme  
284 as in UALVP 59400.

285 *Pelvis:*

286 Small portions of a pubis and ischium (Fig. 7) are represented by fragments collected as float. A  
287 single fragment of the pubis is from the proximal end where it would have contacted the ilium  
288 and provides no information. The ischium is better represented, and most of a shaft can be  
289 reconstructed, although it is likely some fragments are from opposite sides. The reconstructed  
290 morphology of the ischium (Fig. 7) is similar overall to that of TMP 1979.020.0001 in that it is  
291 posteriorly concave and has a tab-like obturator process.

292 *Hindlimb:*

293 The hindlimb is known from fragments of the femur and/or tibia, both astragalocalcanea (Fig. 8),  
294 and a distal tarsal IV (Fig. 9). It is likely that some of the float fragments pertain to other  
295 hindlimb bones including metatarsals, but these cannot be identified with certainty. Based on the  
296 curvature of some fragments, they may represent the femur and/or tibia, but they do not provide  
297 any morphological information. The distal end of the right tibia is preserved in articulation with  
298 the badly crushed astragalocalcaneum (Fig. 8). All that can be said of these is that they were not  
299 fused, the postfibular flange of the tibia was small, and the astragalocalcaneum extended onto the  
300 posterior surface of the tibia as in CMN 8538. The left astragalocalcaneum is less crushed and  
301 shows that there was a transverse groove above the distal condyles (Fig. 8), as in other DPF  
302 caenagnathids (Currie & Russell 1988; Funston et al. 2015), rather than a median tubercle as in  
303 *Anzu wyliei* (Lamanna et al. 2014). The right distal tarsal IV (Fig. 9) is well preserved and  
304 relatively complete. It is roughly triangular in proximal view and tapers in proximodistal  
305 thickness towards its anterior edge. The proximodorsal process is broken off, but posterior to it  
306 there is a notch for metatarsal V (Fig. 9F), which contrasts the morphology of fused  
307 tarsometatarsi referable to *Leptorhynchos elegans* (Currie 1989; Funston, Currie, & Burns 2016).

308 In *Leptorhynchos elegans*, metatarsal V contacts and fuses with the proximodorsal process, and  
309 this coossification fills the notch for metatarsal V. The medial edge of the distal tarsal is mostly  
310 broken, but some intact surface indicates that this tarsal was not fused to distal tarsal III, which  
311 occurs in *Elmisaurus rarus* and *Leptorhynchos elegans* (Osmólska 1981; Currie 1989; Currie et  
312 al. 2016; Funston, Currie, & Burns 2016). Similarly, the intact ventral (distal) articular surface  
313 (Fig. 9D) indicates that distal tarsal IV had not fused to the proximal metatarsus, despite  
314 histological maturity of this individual (see subsequent discussion). This strongly suggests that  
315 this individual did not have a proximally fused metatarsus, which contrasts with all known  
316 specimens of *Leptorhynchos elegans* (Currie 1989; Funston, Currie, & Burns 2016), but is  
317 similar to the known specimens referred to *Chirostenotes pergracilis* (Sternberg 1932; Currie &  
318 Russell 1988; this study).

319

320 ***TMP 1993.036.0181:***

321 TMP 1993.036.0181 (Fig. 10) is a pathological, partial tarsometatarsus described by Funston et  
322 al. (2016), and includes metatarsals II and IV and distal tarsals III and IV. Although initially  
323 described as a representative of *Leptorhynchos elegans* (Funston, Currie, & Burns 2016), several  
324 features suggest it more likely represents *Chirostenotes pergracilis*. For example, it is the largest  
325 tarsometatarsus recovered from Alberta (~250 mm), comparable in size with the holotype of  
326 “*Macrophalangia*”, CMN 8538 (Sternberg 1932), but is not proximally coossified. This is most  
327 evident in the absence of the proximal end of metatarsal III, which shows it had not coossified,  
328 whereas metatarsal III is fused with metatarsals II and IV in all known individuals of  
329 *Leptorhynchos elegans* (Currie 1989; Funston, Currie, & Burns 2016). The clean contact  
330 surfaces for metatarsal III on the proximal ends of metatarsals II and IV further demonstrate the

331 lack of coossification in this individual. Upon reexamination, it is clear that the distal tarsals are  
332 not coossified to each other or with the metatarsals, and a clear suture is visible between them  
333 (Fig. 10). This contrasts with *Leptorhynchos elegans*, where the distal tarsals fuse to each other  
334 and the proximal metatarsus (Currie 1989; Funston, Currie, & Burns 2016). The absence of  
335 fusion between metatarsal V, which is missing, and the proximodorsal process of distal tarsal IV  
336 also contrasts with the condition in known specimens of *Leptorhynchos elegans* (Funston,  
337 Currie, & Burns 2016). In their initial description of the specimen, Funston et al. (2016) also note  
338 that the muscle scars for *M. tibialis cranialis* are less well developed in TMP 1993.036.0181 than  
339 in most specimens of *Leptorhynchos elegans*. In all of these features, TMP 1993.036.0181  
340 closely resembles *Chiostenotes pergracilis* (CMN 8538 and TMP 1979.020.0001).

341

342 ***TMP 2002.012.0103:***

343 TMP 2002.012.0103 is a partial right ilium (Fig. 10A–C) that includes the postacetabular blade  
344 and acetabulum. The pubic peduncle is incompletely preserved, but its ventral edge is  
345 distinctively inclined anterodorsally, similar to TMP 1979.020.0001 but not other DPF  
346 caenagnathid ilia. The acetabulum is slightly constricted transversely at its anteroposterior  
347 midpoint (Fig. 10C), but not to the same degree as other oviraptorosaurs. The ischiadic peduncle  
348 is small and triangular, extending to the same level ventrally as the pubic peduncle. Two large  
349 foramina pierce the posterior surface of the ischiadic peduncle. The postacetabular blade has a  
350 curved dorsal edge, similar to TMP 1979.020.0001, although it is broken along most of its  
351 length. There is a region of rugose bone near the ventral edge of the postacetabular blade that  
352 probably anchored musculature. The medial surface of the ilium (Fig. 10B) is excavated by three  
353 deep concavities, ventral to which a platform accommodates the sacral rib attachments. The

354 brevis fossa is shallow but the brevis shelf is broken, so its full extent cannot be determined.  
355 Overall the morphology of the ilium is remarkably similar to TMP 1979.020.0001, although  
356 TMP 2002.012.0103 is slightly larger and more robustly built.

357

### 358 **Osteohistology**

359 A fragment of an indeterminate long bone (probably the femur or the tibia) was thin-sectioned to  
360 assess the histological maturity of UALVP 59400 (Fig. 11). The element represented by the  
361 fragment cannot be determined with certainty, but considering its featureless external surface and  
362 the curvature of the fragment, it probably represents an area close to the midshaft of either the  
363 femur or the tibia. Other long bones, like the forelimb bones or metatarsals, have smaller  
364 diameters than the femur and tibia (Currie & Russell 1988), and thus would be expected to show  
365 tighter curvature than in this fragment. The exact position of the fragment relative to the midshaft  
366 cannot be established, which limits its usefulness for skeletochronology (Padian et al. 2013), but  
367 it clearly preserves some of the growth record and has little external evidence of muscular  
368 attachment. Because of the limited nature of the material available for sampling, only very  
369 limited insights can be made regarding the growth of this individual (Horner et al. 1999; Horner  
370 et al. 2001; Cullen et al. 2014; Woodward et al. 2014; Woodward et al. 2015). However, this  
371 information is nonetheless important for our understanding of caenagnathid diversity in the DPF.  
372 Future systematic study of caenagnathid osteohistology using better material may provide a more  
373 complete context within which UALVP 59400 can be placed.

374 Histological preservation of the specimen is excellent and several important features can  
375 be discerned (Fig. 11). The relative bone wall thickness is low, as expected of a theropod  
376 (Varricchio 1993; Horner & Padian 2004; Erickson et al. 2009; Funston, Currie, & Burns 2016),

377 but its exact value cannot be calculated without more complete material, as there is uncertainty in  
378 the diameter of the bone. The medullary cavity is large and lined by thick layers of endosteal  
379 lamellae (Fig. 11B), which suggests much of the earlier growth record has been resorbed (Horner  
380 & Padian 2004; Padian et al. 2013). There are no trabeculae in the medullary cavity, which is  
381 similar to other theropods (Varricchio 1993; Horner & Padian 2004; Cullen et al. 2014; Xu et al.  
382 2018). The cortex is composed of mostly of primary fibrolamellar bone (Francillon-Vieillot et al.  
383 1990) with abundant vascularization. Osteocyte lacunae are relatively dense ( $\sim 47,000/\text{mm}^3$ ),  
384 similar to other oviraptorosaurs and small dinosaurs (Stein & Werner 2013; Cullen et al. 2014;  
385 Funston & Currie 2018; Gregory F. Funston et al. 2019). Most osteocytes are flattened in shape,  
386 reflecting the well-developed primary osteons and the accompanying lamellar bone. The density  
387 and shape of the osteocytes is consistent throughout the entire cortex. Vascularity throughout the  
388 cortex is predominantly reticular in orientation, but in some areas of the inner cortex, it  
389 approaches a sub-plexiform orientation (Fig. 11E). Conversely, towards the periosteal surface  
390 there is a higher proportion of longitudinal canals (Fig. 11E). In these regions, the bone  
391 approaches a laminar organization (Francillon-Vieillot et al. 1990). Regardless, many radial  
392 anastomoses are still present in this region, and in most areas the vascularity is best described as  
393 reticular (Francillon-Vieillot et al. 1990; de Margerie 2004).

394         The cortex is marked by 11 lines of arrested growth (LAGs), two pairs of which are  
395 doublet LAGs (Fig. 11C)—common in non-avian theropods (Lee & O'Connor 2013; Cullen et  
396 al. 2014; Zanno et al. 2019). These doublet LAGs likely represent a single cyclical growth mark  
397 (CGM; Castanet et al. 1990), and taking this into consideration, nine CGMs can be counted  
398 throughout the cortex (Fig. 11E). Whereas the outer LAGs are each roughly parallel to the  
399 endosteal and periosteal surfaces, the innermost LAG is more tightly curved and intersects with

400 the medullary cavity at one end. Each LAG is accompanied endosteally by an annulus of bone  
401 with an increased proportion of parallel-fibered bone matrix and a reduction in vascularity (Fig.  
402 11D). An extra annulus can be discerned periosteal to the innermost LAG, but it does not  
403 encircle the entire cortex, nor is it associated with a rest line. The CGMs decrease in spacing  
404 towards the periosteal surface, but not in a regular fashion. Whereas the first, second and third  
405 CGMs are widely spaced, the remaining CGMs are much more closely spaced (Fig. 11E). These  
406 more periosteal CGMs are evenly spaced, each separated by a single circumferential row of  
407 predominantly longitudinal vascular canals (Fig. 11E). The close packing of LAGs and annuli  
408 towards the periosteal surface is associated with a higher proportion of parallel-fibered bone  
409 matrix (Fig. 11D), but the matrix is still predominantly fibrolamellar. These LAGs therefore do  
410 not represent a fully-developed external fundamental system (Horner et al. 1999; Woodward et  
411 al. 2014; Woodward et al. 2015). However, the bone right at the periosteal surface is parallel-  
412 fibered and has reduced vascularity (Fig. 11D), which suggests it may represent an incipient  
413 EFS. Alternatively, this may represent the formation of the annulus that precedes LAGs  
414 elsewhere in the cortex. Secondary remodelling is generally restricted, but there are some  
415 secondary osteons endosteal to the innermost LAG (Fig. 11B). These are concentrated in a small  
416 area and none are cross-cut by other secondary osteons, nor by the endosteal lamellae.

417

418 **Discussion:**

419

420 ***Taxonomy:***

421 A complete taxonomic review is beyond the scope of this study and is in preparation for  
422 publication elsewhere (GFF in prep). However, it is important to note that where DPF



423 caenagnathid elements are known from multiple specimens (i.e. mandibles, ilia, tarsometatarsi),  
424 in each case they support the distinction of only three morphotypes—likely corresponding to the  
425 three currently recognized genera. In addition to morphology, these groups appear to differ in  
426 adult body size (Bell et al. 2015; Funston et al. 2015; Gregory F. Funston et al. 2019), and  
427 currently known material from *Chirostenotes pergracilis* is intermediate in size between the  
428 smaller *Leptorhynchus elegans* and the larger *Caenagnathus collinsi*.

429 Evidence from the mandible is important for distinguishing DPF caenagnathids. Despite  
430 being the most completely known caenagnathid from the DPF, the lack of definitive mandibular  
431 material for *Chirostenotes pergracilis* has made it difficult to refer isolated specimens to this  
432 taxon, and has left its synonymy with *Caenagnathus collinsi* ambiguous (Sues 1997). The  
433 mandibles of CMN 8776 (*Caenagnathus collinsi*) are clearly distinct from all other mandibles  
434 collected from the DPF and their taxonomic distinction is unambiguous. Two morphotypes can  
435 be distinguished in the remaining mandibles from the DPF, but the taxa to which they pertain is  
436 unclear. All of these mandibles have dentaries that are upturned at their anterior ends, but some  
437 of the smaller specimens (e.g. TMP 1992.036.0390) have dentaries that are more strongly  
438 upturned, have a more square anteroventral margin in lateral view, and have more widely  
439 diverging rami in dorsal view (Currie et al. 1993; Longrich et al. 2013). Funston et al. (2019)  
440 showed that these morphotypes differed in adult body size, and therefore likely pertained to two  
441 different taxa: the larger *Chirostenotes pergracilis* and the smaller *Leptorhynchus elegans*.  
442 However, this conclusion was tentative because neither *Chirostenotes pergracilis* nor  
443 *Leptorhynchus elegans* are known from unambiguous mandibular material.

444 The discovery of UALVP 59400 is thus particularly important because it preserves a  
445 relatively complete mandible in association with other postcranial bones. The morphology of the

446 mandible of UALVP 59400 shows that it is distinct from *Caenagnathus collinsi*, which lacks the  
447 upturned anterior tip of the beak present in UALVP 59400. The dentaries of UALVP 59400 are  
448 much larger than the small, histologically mature dentaries (UALVP 55639) described by  
449 Funston et al. (2019), and are more similar in size and morphology to the largest dentaries in that  
450 study (TMP 1992.036.1237). The correspondence in size, morphology, and histological signals  
451 in TMP 1992.036.1237 (Gregory F. Funston et al. 2019) and UALVP 59400 strongly support the  
452 referral of all this material to the same taxon. In this light, the isolated distal tarsal IV of UALVP  
453 59400 is important. The lack of fusion between distal tarsal IV and the metatarsus—despite  
454 histological maturity (see below)—shows that this specimen is distinct from *Leptorhynchos*  
455 *elegans*, where the distal tarsals fuse to each other and the metatarsus at or before maturity (G.  
456 Funston unpubl. diss.). Fusion of the hindlimb bones also appears to coincide with histological  
457 maturity in other oviraptorosaurs (Gregory F. Funston et al. 2019). There is variation in the  
458 ontogenetic sequence of some early theropods (C. T. Griffin & Nesbitt 2016; Christopher T.  
459 Griffin & Nesbitt 2016; Griffin 2018), particularly regarding the fusion of the tarsometatarsus  
460 (Griffin 2018). However, this variation is lost in more derived non-avian theropods, and is  
461 virtually absent in avian theropods (Christopher T. Griffin & Nesbitt 2016). The close  
462 phylogenetic position of caenagnathids to Aves suggests they probably had low variation in their  
463 ontogenetic sequences, and thus the onset of fusion of the tarsometatarsus is unlikely to have  
464 been highly variable through ontogeny. Indeed, in oviraptorosaurs with fused tarsometatarsi,  
465 there appears to be minimal size variation in fused specimens of each taxon (Funston, Currie, &  
466 Burns 2016; Funston, Currie, Eberth, et al. 2016; Gregory F. Funston et al. 2019), which  
467 tentatively suggests that fusion coincides with the achievement of maximum body size. The  
468 distinction of UALVP 59400 and, thus, the larger mandible morphotype from *Leptorhynchos*

469 *elegans* strongly supports the conclusion of Funston et al. (2019) that the larger mandible  
470 morphotype is referable to *Chirostenotes pergracilis*.

471 Therefore, considering that 1) UALVP 59400 is distinct from *Caenagnathus collinsi* on  
472 the basis of the mandible; 2) UALVP 59400 is separate from *Leptorhynchos elegans* on the basis  
473 of the unfused distal tarsal IV; and 3) there is no evidence for a second caenagnathid taxon  
474 intermediate in size between *Caenagnathus collinsi* and *Leptorhynchos elegans* in the Dinosaur  
475 Park Formation; it can be concluded that UALVP 59400 is referable to *Chirostenotes*  
476 *pergracilis*. A corollary of this is that, considering that 1) the smaller, upturned dentaries are  
477 distinct from both *Caenagnathus collinsi* (CMN 8776) and *Chirostenotes pergracilis* (UALVP  
478 59400); and 2) there is no evidence for a second, small caenagnathid in the Dinosaur Park  
479 Formation; the small, upturned dentary morphotype represents *Leptorhynchos elegans*. The  
480 further discovery of associated caenagnathid skeletons and the systematic osteohistological  
481 sampling of material may provide new lines of evidence to refute or support these hypotheses.

482

#### 483 ***Anatomy:***

484 The referral of new material to *Chirostenotes pergracilis* fills important gaps in the anatomy of  
485 this taxon. Referral of a relatively complete mandible (UALVP 59400) allows for more confident  
486 identification of isolated mandibular material, including TMP 2001.012.0012, an exceptionally  
487 complete mandible (Funston and Currie 2014). The ventral suture present on the surangular of  
488 both these specimens is noteworthy. Comparison to *Citipati osmolskae* (Clark et al. 2002), an  
489 undescribed *Conchoraptor gracilis*, and *Rinchenia mongoliensis* (Funston et al. 2018) shows that  
490 this suture corresponds in position to the suture between the coronoid and surangular of  
491 oviraptorids. This suggests that the coronoid is indeed present in caenagnathids, but is fused into

492 the ASC complex, as suspected by Currie et al. (1993). Combined with its presence in  
493 *Incisivosaurus gauthieri* (Balanoff et al. 2009), this suggests that the coronoid is present  
494 throughout Oviraptorosauria, but may appear absent because it fuses to the surangular.

495 UALVP 59400 includes cervical vertebrae, which have not yet been recovered from the  
496 DPF, as well as a series of articulated caudal vertebrae, which may be useful for future  
497 comparisons and referrals of isolated vertebrae. The cervical vertebrae are generally similar to  
498 those of *Anzu wyliei* (Lamanna et al. 2014), *Apatoraptor pennatus* (Funston & Currie 2016) and  
499 *Epichirostenotes curriei* (Sues 1997) in the reduced neural spines and anterodorsally directed  
500 prezygapophyses. However, they differ from those of *Apatoraptor pennatus* in the better  
501 development of the laminae connecting the postzygapophysis and transverse process, in which  
502 respect they are more similar to *Epichirostenotes curriei* (Funston & Currie 2016). Similarly, the  
503 cervical ribs appear unfused, despite the advanced skeletal maturity of UALVP 59400, which is  
504 more like *Epichirostenotes curriei* than *Anzu wyliei* or *Apatoraptor pennatus* (Sues 1997;  
505 Lamanna et al. 2014; Funston & Currie 2016). Compared to oviraptorids, the epipophyses of  
506 caenagnathids are reduced, whereas in oviraptorids they are generally large anteriorly (Funston  
507 et al. 2018). The caudal vertebrae of UALVP 59400 are similar to those of other oviraptorosaurs,  
508 but the chevrons are distinctive. In oviraptorids, the chevrons are dorsoventrally deep  
509 proximally, but become platelike, dorsoventrally shorter, and anteroposteriorly longer towards  
510 the distal end of the tail (Balanoff & Norell 2012; Persons IV et al. 2013). Those of *Anzu wyliei*,  
511 *Chirostenotes pergracilis*, and *Nomingia gobiensis*, in contrast, are much shorter in dorsoventral  
512 height, and become rectangular toward the distal end of the tail (RZNCHEEN Barsbold et al.  
513 2000; Lamanna et al. 2014). The anteroposteriorly elongate proximal end of each chevron in

514 *Chiostenotes pergracilis* is distinctive among caenagnathids, but whether it served a functional  
515 purpose is unclear.

516         The unusual stains and carbonaceous material preserved with the cervical vertebrae of  
517 UALVP 59400 (Fig. 3C–E) are similar to features described in the feathers of oviraptorosaurs  
518 and other theropods from China (Qiang et al. 1998; Xu & Zhang 2005; Xu et al. 2010; Xu et al.  
519 2012; Xu et al. 2017). In particular, the filamentous, ‘tufted’ appearance of the stains and the  
520 consistent orientation of the filaments parallel to the neck are similar to feather impressions on  
521 the neck and back of *Anchiornis huxleyi* (Zheng et al. 2018), *Caudipteryx zoui* (Qiang et al.  
522 1998), and *Sinosauropteryx prima* (Chen et al. 1998; Currie & Chen 2001; Smithwick, Nicholls,  
523 et al. 2017). Considering that evidence of feather preservation is known from other specimens in  
524 the Dinosaur Park and nearby Horseshoe Canyon Formations (Zelenitsky et al. 2012; van der  
525 Reest et al. 2016), this seems a plausible explanation for these structures. Although some authors  
526 have argued that these types of impressions represent collagen or the dermis (Lingham-Soliar et  
527 al. 2007; Lingham-Soliar 2016), this interpretation is doubtful (Smithwick, Mayr, et al. 2017). In  
528 any case, the filaments in UALVP 59400 are poorly preserved, probably as a result of  
529 taphonomic degradation, and they provide little information about the type or distribution of  
530 feathers in *Chiostenotes pergracilis*. Some possible branching structures can be detected (Fig.  
531 3E), but none of these is conclusive and therefore whether these feathers are filaments or  
532 pennaceous is ambiguous, although pennaceous feathers are known in other oviraptorosaurs  
533 (Qiang et al. 1998; Xu et al. 2010). More information about the feathers in UALVP 59400 might  
534 be gleaned by future chemical or microscopic analyses, which are outside of the scope of this  
535 study.

536           The other isolated specimens described here overlap with TMP 1979.020.0001 but reveal  
537 more of the morphology of those elements. Re-examination of a pathological tarsometatarsus  
538 (TMP 1993.036.0181) previously referred to *Leptorhynchos elegans* suggests that it more likely  
539 represents *Chirostenotes pergracilis*, because it lacks fusion of the distal tarsals to each other or  
540 the metatarsals. Although less complete than the metatarsals of TMP 1979.020.0001, the  
541 uncrushed nature of this specimen provides additional information on the pedal structure of  
542 *Chirostenotes pergracilis*. Importantly, this specimen elucidates the morphology of the  
543 proximodorsal process of distal tarsal IV, which is similar to—but smaller than—that of  
544 *Elmisaurus elegans* (Osmólska 1981; Currie et al. 2016) and *Leptorhynchos elegans* (Funston,  
545 Currie, & Burns 2016); it was probably a shared feature of most caenagnathids. The isolated  
546 ilium (TMP 2002.012.0103) provides a second example of the unusually tall ilium of  
547 *Chirostenotes pergracilis*, which is similar to that of *Rinchenia mongoliensis* (Funston et al.  
548 2018). This specimen also shows that the inclination of the ventral edge of the pubic peduncle  
549 may be useful for identifying more fragmentary ilia.

550           More of the anatomy and skeletal proportions of *Chirostenotes pergracilis* can be  
551 reconstructed based on TMP 1979.020.0001 and UALVP 59400 (Fig. 1B). This shows that the  
552 hindlimbs were exceptionally elongate, and the pelvis was relatively small. The tail was also  
553 relatively short, and the shorter chevrons would have reduced space available for musculature.  
554 Together with the elongate pedal digits adapted for grasping (Varricchio 2001; Longrich et al.  
555 2010; Longrich et al. 2013), these features support previous suggestions that the hindlimbs of  
556 *Chirostenotes pergracilis* were elongated to facilitate wading behaviour, rather than enhanced  
557 cursoriality.

558

559 ***Growth in *Chirostenotes pergracilis*:***

560 The fragmentary nature of the sampled material limits the scope of the conclusions that can be  
561 drawn. Regardless, the osteohistology of UALVP 59400 shows that this individual was  
562 approaching maximum body size, and that growth had slowed considerably in the final years of  
563 life. Assuming that doublet LAGs represent a single growth mark, UALVP 59400 was at least 9  
564 years of age at death. There is some evidence of shape change or cortical drift preserved in the  
565 cortex. For example, tighter curvature of the innermost LAG and its intersection with the  
566 medullary cavity indicates that this region of the bone changed in curvature throughout life.

567         Reticular to sub-plexiform vascularity in the inner cortex of the femur or tibia of UALVP  
568 59400 (Fig. 11) suggests that this individual had relatively rapid rates of growth at this stage of  
569 life. This is similar to the outer cortex of UALVP 57349, a small caenagnathid tibia from the  
570 HCF (Funston & Currie 2018), and to the bone in young avimimids (Gregory F. Funston et al.  
571 2019). The same is true as well of ornithomimids (Varricchio et al. 2008; Cullen et al. 2014;  
572 Skutschas et al. 2016), but appears not to have been the case in smaller non-theropods like  
573 alvarezsaurids and dromaeosaurids (Erickson et al. 2009; Xu et al. 2018). These animals appear to  
574 have had lower growth rates throughout life, reflected by their smaller body sizes (Erickson et al.  
575 2009). Despite similar body sizes, the reticular–plexiform vascular patterns in UALVP 59400  
576 and other caenagnathids (Funston & Currie 2018) suggest a more rapid growth rate than in  
577 oviraptorids (Erickson et al. 2007; Norell et al. 2018), which have femoral cortices with  
578 predominantly longitudinal vasculature. However, this may be the result of variation in element-  
579 specific growth rate (Horner et al. 1999; Cullen et al. 2014; Woodward et al. 2014), reflecting  
580 allometry rather than growth rates in the whole individual. Systematic sampling of caenagnathids

581 and oviraptorids of varying body sizes, controlling for sampling location, could clarify growth  
582 rate variation in Oviraptorosauria.

583         Towards the periosteal surface of the cortex of UALVP 59400, the spacing between  
584 CGMs is drastically reduced (Fig. 11E). This suggests that there was a transition to a slower  
585 growth rate, and that this individual had approached maximum body size. Although variation in  
586 LAG spacing between skeletal elements can pose problems for extrapolating maturity in isolated  
587 specimens (Cullen et al. 2014), the high number of closely-spaced LAGs in UALVP 59400  
588 strongly suggests that growth was diminished for several years, which is more likely to represent  
589 skeletal maturity than allometry. Indeed, the parallel-fibered bone at the periosteal surface may  
590 be indicative of an incipient EFS, signalling the cessation of growth. In extant amniotes, the  
591 onset of sexual maturity coincides with a stark decrease in linear growth rates (Castanet et al.  
592 2004; Lee & Werning 2008; Köhler et al. 2012; Botha-Brink et al. 2016). Although a broader  
593 sample of caenagnathids would be required to be certain, the transition preserved in UALVP  
594 59400 may indicate that this individual was sexually mature.

595

## 596 **Conclusions:**

597

598 A new partial skeleton referable to *Chirostenotes pergracilis* improves the skeletal representation  
599 of this taxon. The specimen preserves the first mandible associated with postcrania from the  
600 Dinosaur Park Formation, as well as the first cervical vertebrae and chevrons. Osteohistology  
601 shows that the individual represented by UALVP 59400 was skeletally mature and approaching  
602 maximum body size. Based on UALVP 59400, *Chirostenotes pergracilis* can be distinguished  
603 from *Caenagnathus collinsi* on the basis of the dentary, which is anteriorly upturned, and from



604 *Leptorhynchos elegans* by its larger adult body size and the absence of fusion of the distal tarsals  
605 and proximal metatarsals. This helps to clarify the taxonomy of Dinosaur Park Formation  
606 caenagnathids, and mandibular material can now be confidently referred to each of the three  
607 taxa.

608

609

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611

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622

623 **Declaration of interests:**

624 The authors declare there are no conflicts of interest, financial or otherwise.

625

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- 849
- 850



851 Table 1. Selected measurements of UALVP 59400 and other caenagnathids

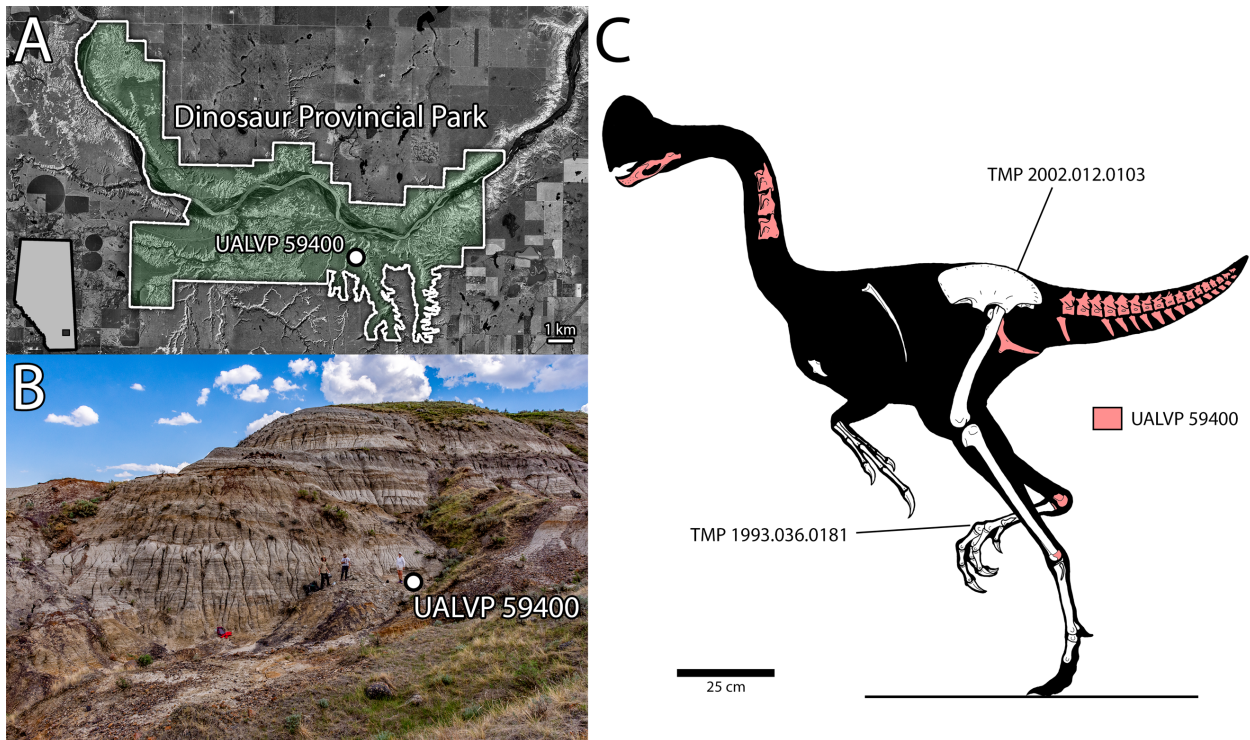
		<i>Chirostenotes pergracilis</i>		<i>Apatoraptor pennatus</i>	<i>Epichirostenotes curriei</i>	<i>Anzu wyliei</i>	<i>Nomingia gobiensis</i>
Measurement (mm)		UALVP 59400	TMP 2001.012.0012	TMP 1993.051.0001	ROM 43250	CM 78001	MPC-D 100/119
Mandible: length of articular ridge		23.5	24.7	21.1	-	35.7	-
Mandible: minimum height of dentary		20.7	20.9	13.6	-	25.8	-
Cervical Vertebrae (length of centrum)	Cv7	49e	-	54.5	81	86	-
	Cv8	57e	-	55	85	95	-
Caudal Vertebrae (length of centrum)	C8	27.1	-	-	-	34	22
	C9	25.2	-	-	-	32	21
	C10	22.3	-	-	-	31	20
	C11	21.4	-	-	-	31	18
	C12	19.5	-	-	-	-	18
	C13	19.4	-	-	-	-	16
	C14	16.1	-	-	-	-	16
	C15	20.4	-	-	-	-	16
	C16	20.9	-	-	-	-	17
	C17	16.9	-	-	-	-	19
Chevron 8, length		58.7	-	-	-	-	42

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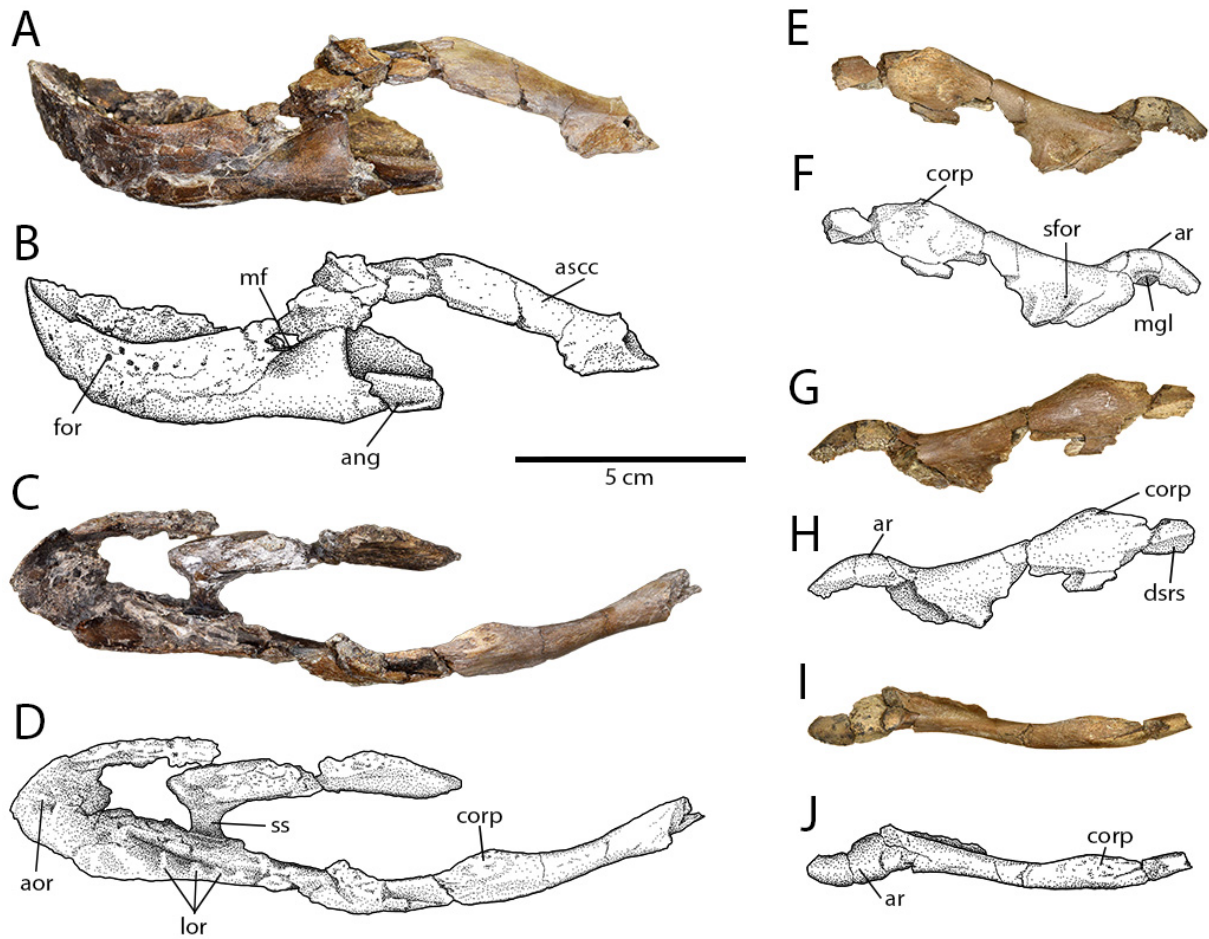
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857 FIGURE 1. **A**, Satellite image of Dinosaur Provincial Park (highlighted in green), showing the  
858 locality where UALVP 59400 was found. Inset shows location of satellite image within Alberta,  
859 Canada. **B**, Field photography of site, indicated by the white point. Note three people standing to  
860 the left of the point, for scale. **C**, Reconstruction of *Chirostenotes pergracilis*, showing known  
861 elements, with bones preserved in UALVP 59400 highlighted in red. Satellite imagery in (A)  
862 from Google Maps (map data: © Google Maps), used under fair use terms.

863



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865

866 FIGURE 2. Mandibles of *Chirostenotes pergracilis* (UALVP 59400). **A, B**, Photograph (A) and

867 illustration (B) of mandibles in left lateral view. **C, D**, Photograph (C) and illustration (D) of

868 mandibles in dorsal view. **E, F**, Photograph (E) and illustration (F) of right articular-surangular-

869 coronoid complex in medial view. **G, H**, Photograph (E) and illustration (F) of right articular-

870 surangular-coronoid complex in lateral view. **I, J**, Photograph (I) and illustration (J) of right

871 articular-surangular-coronoid complex in dorsal view. **Abbreviations:** **ang**, angular; **aor**,

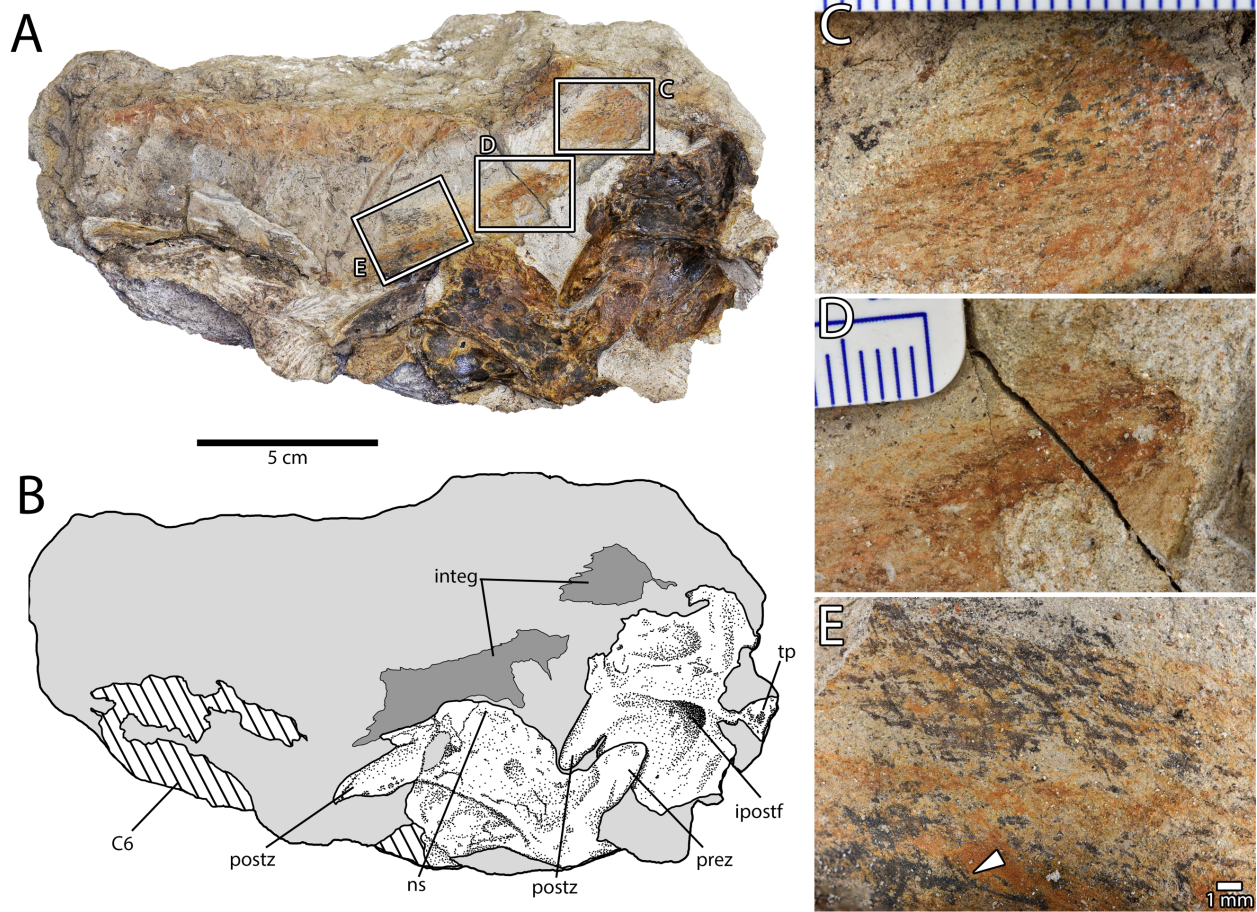
872 anterior occlusal ridge; **ar**, articular ridge; **ascc**, articular-surangular-coronoid complex; **corp**,

873 coronoid process; **dsrs**, dentary-surangular suture; **for**, foramen; **lor**, lateral occlusal ridges; **mf**,

874 mandibular fossa; **mgl**, medial glenoid; **sfor**, surangular foramen; **ss**, symphyseal sulcus.

875

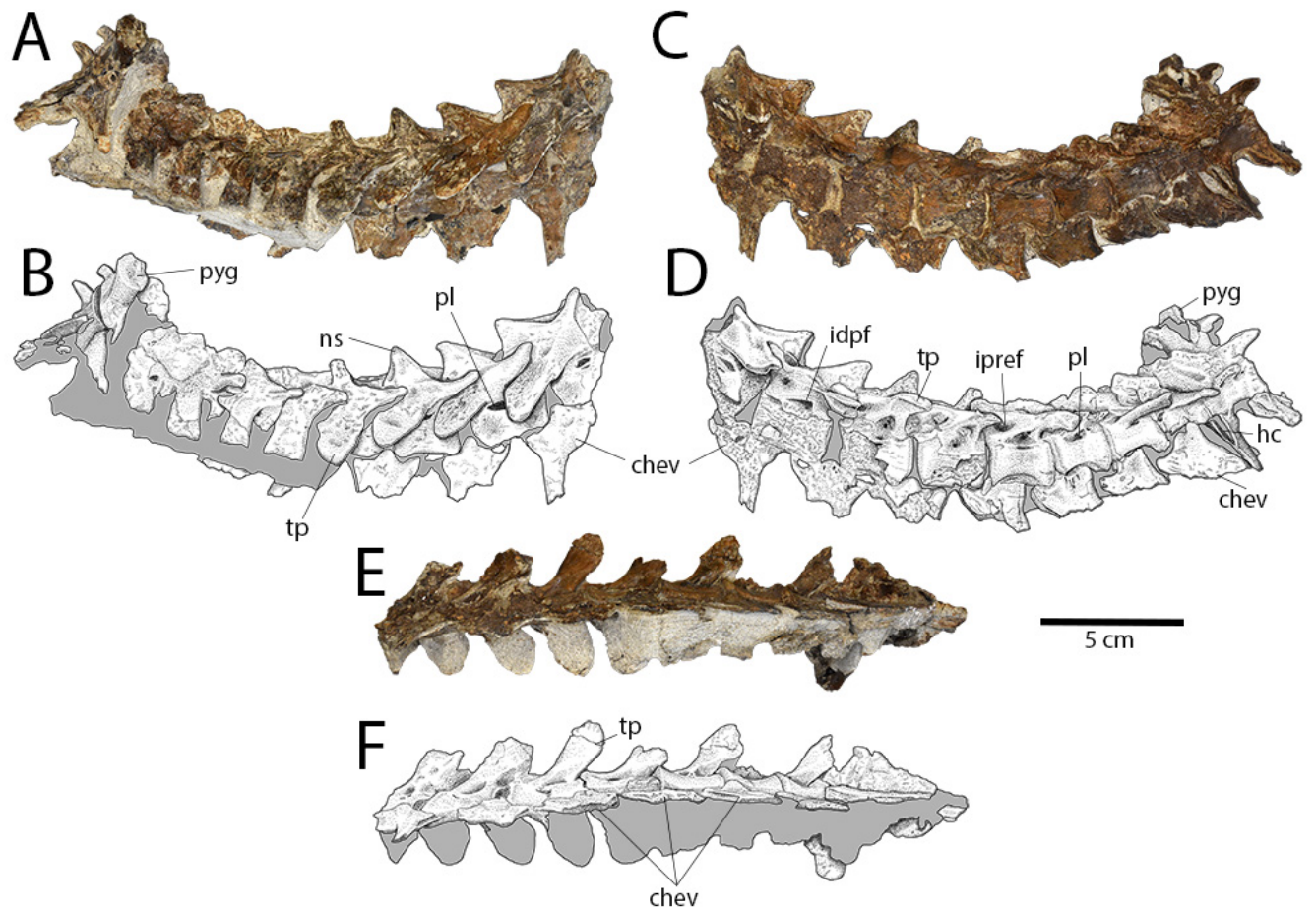
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877 FIGURE 3. Cervical vertebrae of *Chirostenotes pergracilis* (UALVP 59400). **A, B**, Photograph  
878 (A) and illustration (B) of postaxial cervical vertebrae four, five, and six in right lateral view.  
879 Boxes in (A) indicate the locations of close-up images (C–E). Light grey indicates matrix, dark  
880 grey indicates possible integument, and lines indicate broken bone. **C–E**, Close-up photographs  
881 of possible feather impressions dorsal to the cervical vertebrae. Note the filamentous impressions  
882 and the consistent orientation of the fibers. Arrow in (E) indicates possible branching structure.  
883 Scales in (C) and (D) are in millimetres. **Abbreviations:** **C6**, postaxial cervical vertebra six;  
884 **integ**, possible integumentary structures; **ipostf**, infrapostzygapophyseal fossa; **ns**, neural spine;  
885 **postz**, postzygapophysis; **prez**, prezygapophysis; **tp**, transverse process.



886



887

888 FIGURE 4. Caudal vertebrae of *Chirostenotes pergracilis* (UALVP 59400). **A, B**, Photograph

889 (A) and illustration (B) of articulated distal caudal vertebral series in right lateral view. **C, D**,

890 Photograph (C) and illustration (D) of articulated distal caudal vertebral series in left lateral

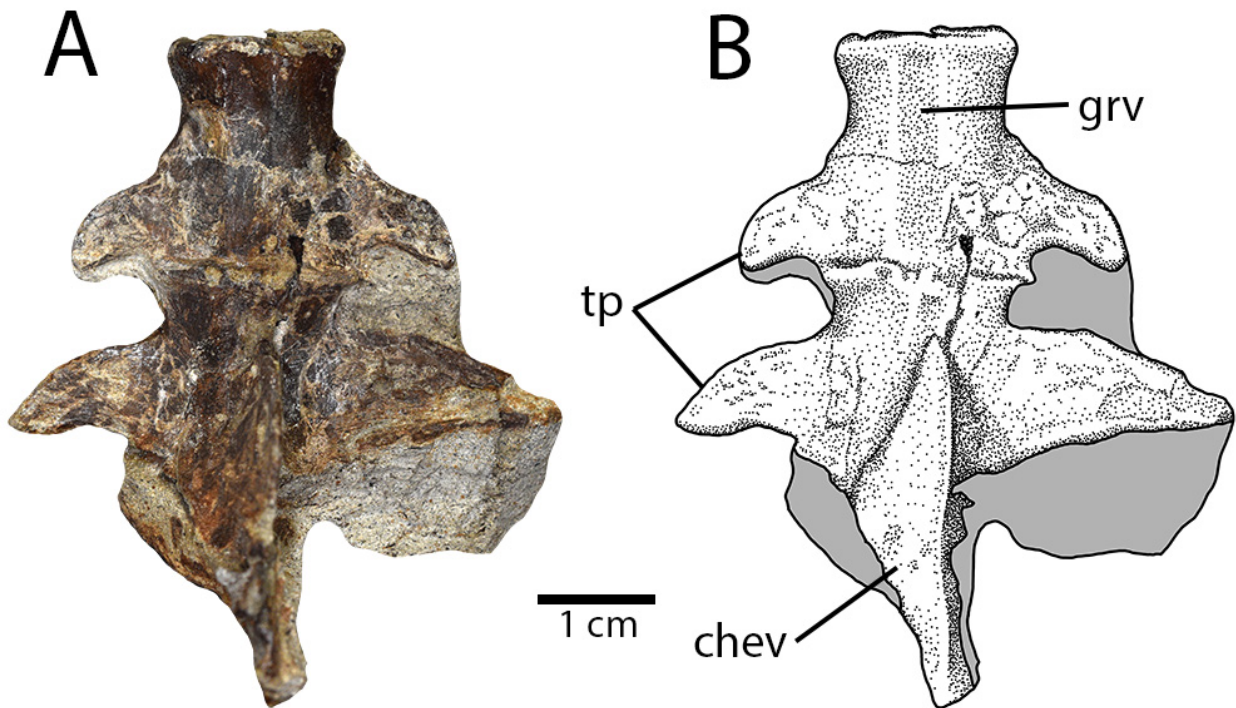
891 view. **E, F**, Photograph (E) and illustration (F) of articulated distal caudal vertebral series in

892 ventral view. **Abbreviations:** chev, chevron; hc, haemal canal; idpf, infradiapophyseal fossa;

893 ipref, infraprezygapophyseal fossa; ns, neural spine; pl, pleurocoel; pyg, pre-pygal vertebra; tp,

894 transverse process.

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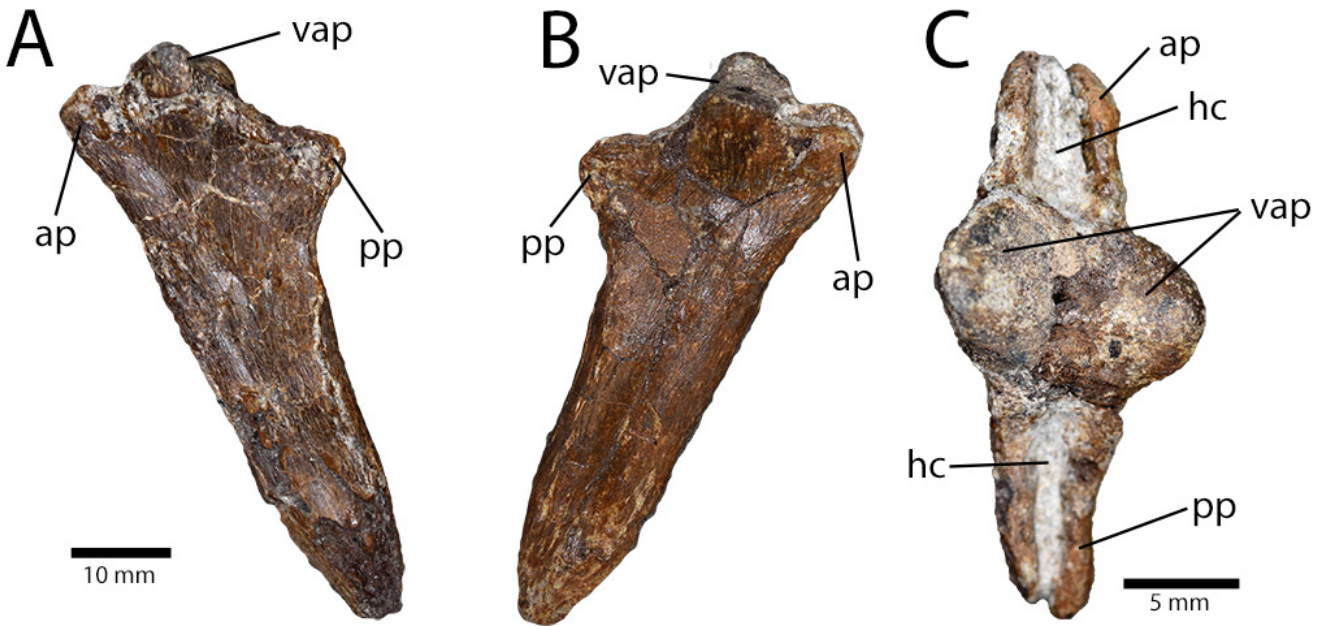
897 FIGURE 5. Distal caudal vertebrae of *Chirostenotes pergracilis* (UALVP 59400). **A, B,**

898 Photograph (A) and illustration (B) of pre-pygal caudal vertebrae of UALVP 59400 in ventral  
899 view, showing anteriorly-directed transverse processes, midline ventral groove, and large,

900 platelike chevrons. Anterior is downwards, distal (posterior) is upwards, and lateral is to either

901 side. **Abbreviations:** chev, chevron; grv, groove; tp, transverse process.

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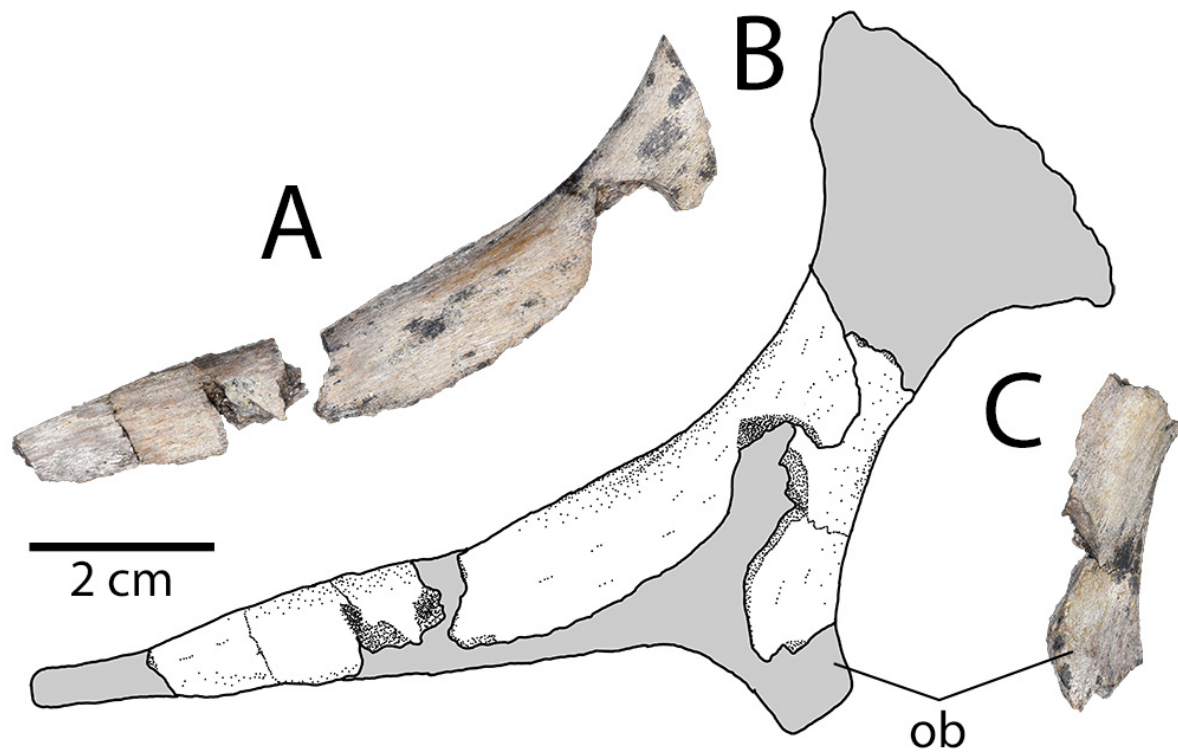
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905 FIGURE 6. Chevron of *Chirostenotes pergracilis* (UALVP 59400). **A–C**, mid-caudal chevron in

906 left lateral (A), right lateral (B), and proximal views. **Abbreviations:** **ap**, anterior process; **hc**,

907 haemal canal; **pp**, posterior process ;**vap**, vertebral articular processes.

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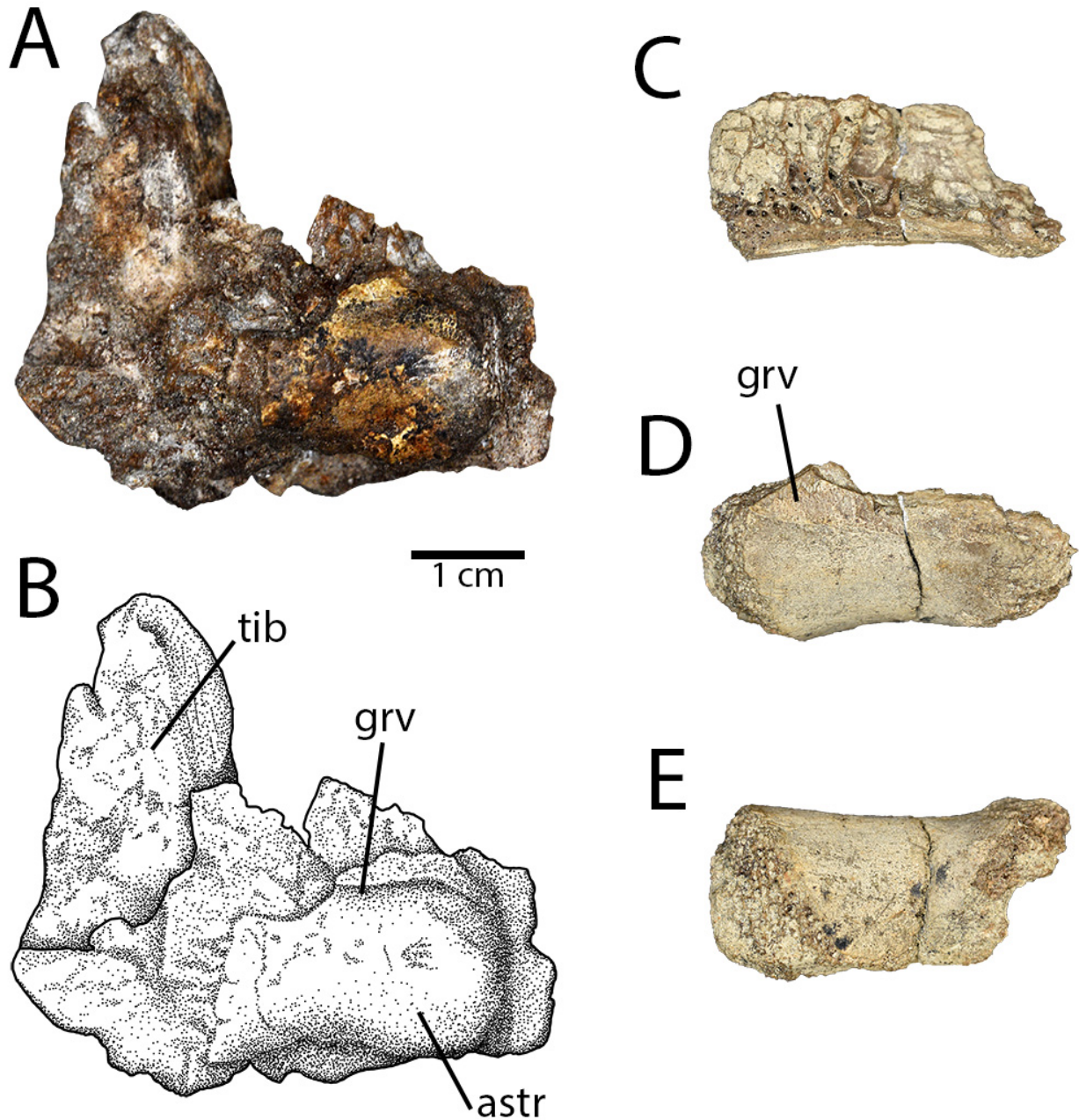
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910 FIGURE 7. Ischium of *Chirostenotes pergracilis* (UALVP 59400). **A**, Fragments of right  
 911 ischium in lateral view. **B**, Composite reconstruction of ischium based on fragments of the left  
 912 and right ischia, reconstructed using the complete ischium of TMP 1979.020.0001 (silhouette).  
 913 **C**, Fragment of right ischium in medial view. **Abbreviation: ob**, obturator process.

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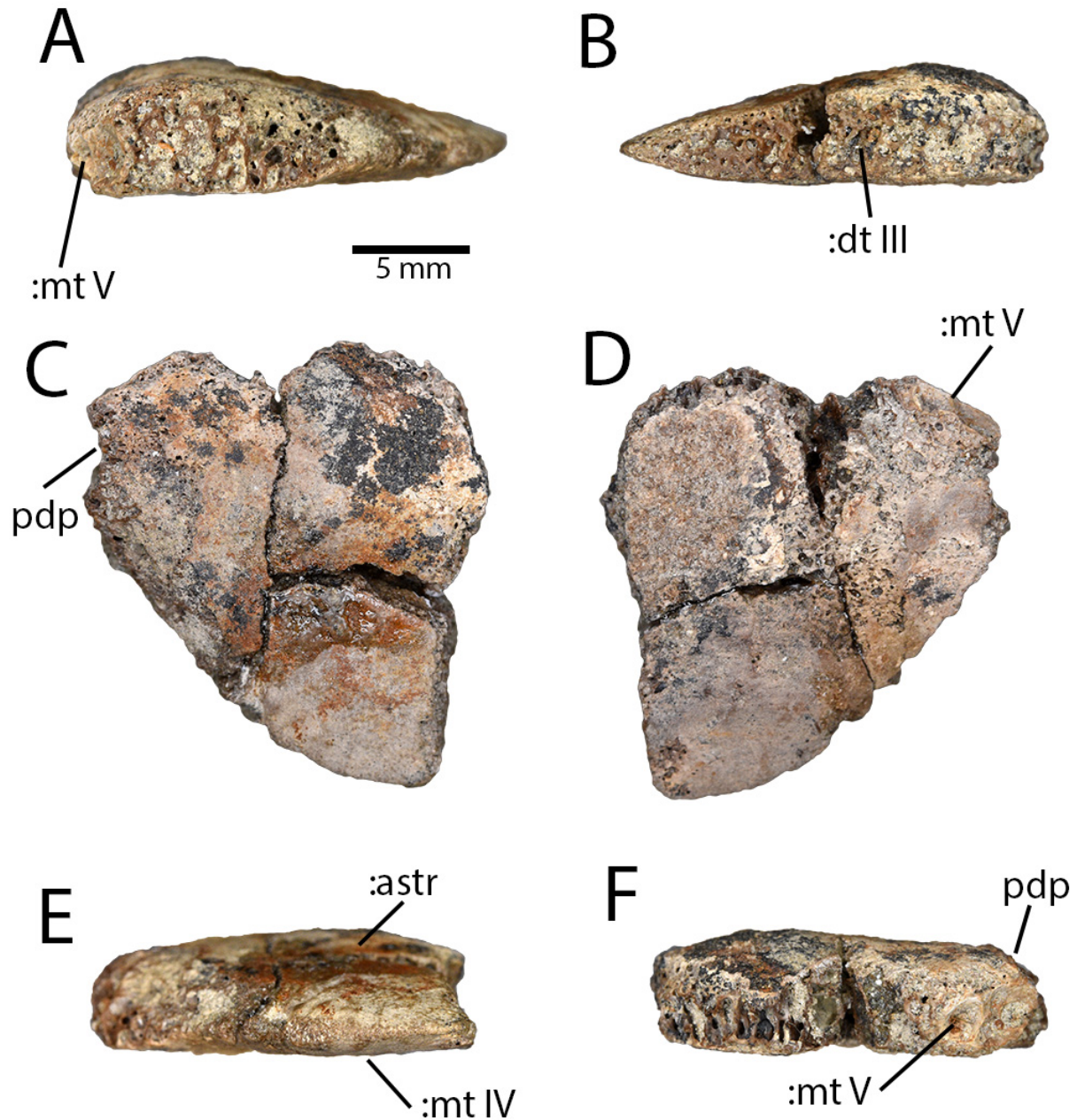
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918 FIGURE 8. Astragalocalcanei of *Chirostenotes pergracilis* (UALVP 59400). **A, B**, Photograph

919 (A) and illustration (B) of right distal tibia and astragalocalcaneum in anterior view. **C–E**, left

920 astragalocalcaneum in dorsal (C), anterior (D), and ventral (E) views. **Abbreviations: astr,**

921 astragalus; **grv**, groove; **tib**, tibia.



923

924 FIGURE 9. Distal tarsal IV of *Chirostenotes pergracilis* (UALVP 59400). A–F, Right distal  
 925 tarsal IV in lateral (A), medial (B), proximal (C), distal (D), anterior (E) and posterior (F) views.  
 926 Note the intact distal surface, indicating that the distal tarsal was not fused to the proximal end of  
 927 metatarsal IV. **Abbreviations:** :astr, contact for astragalus; :dt III, contact for distal tarsal III;

928 **:mt IV**, contact for metatarsal IV; **:mt V**, contact for metatarsal V; **pdp**, base of proximodorsal

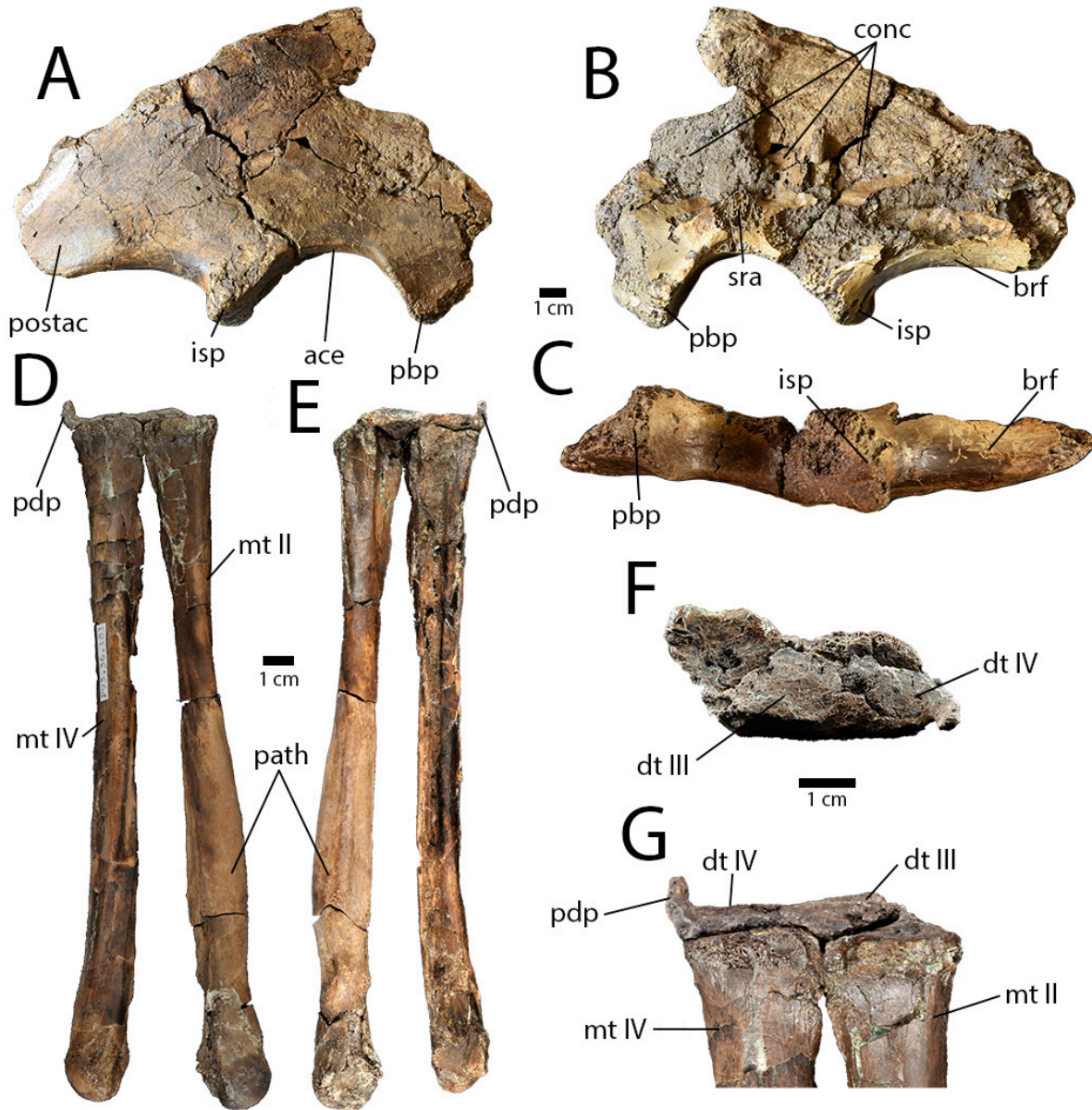
929 process.

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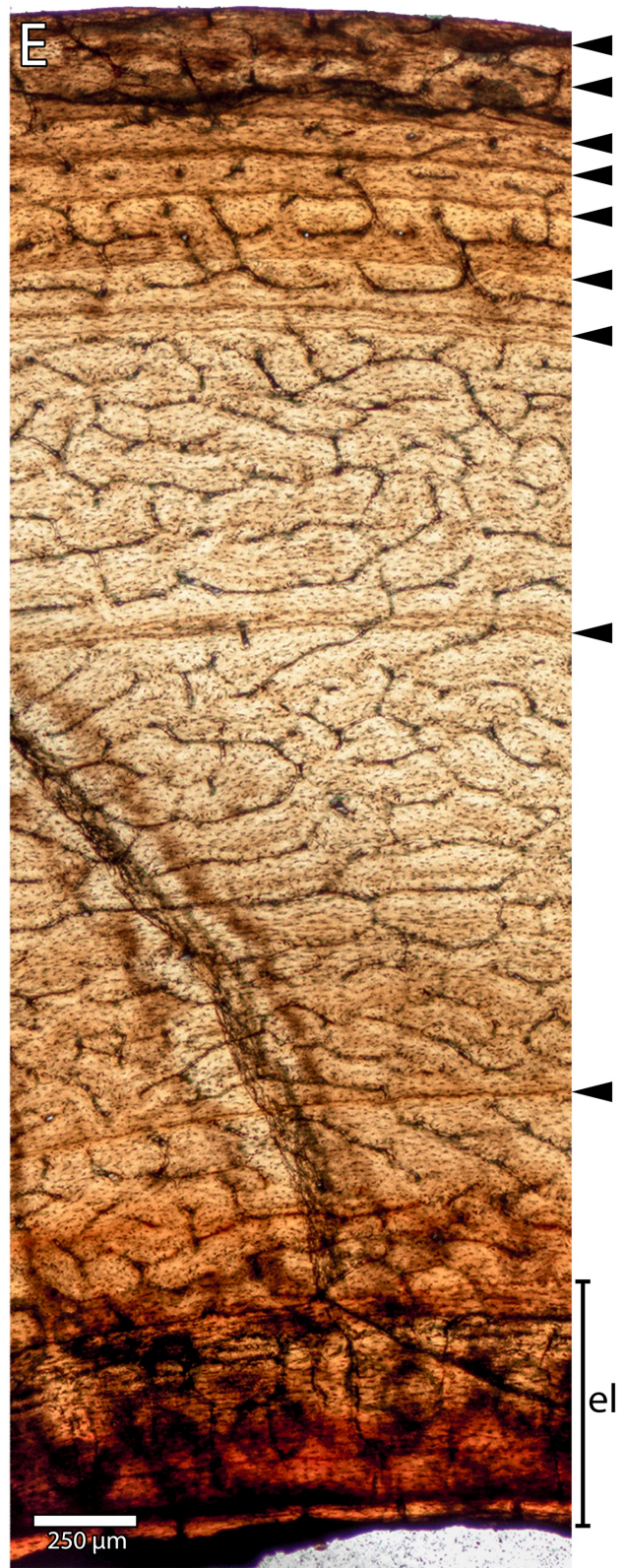
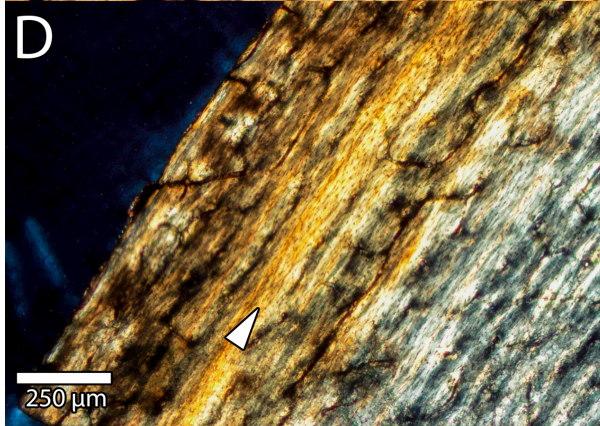
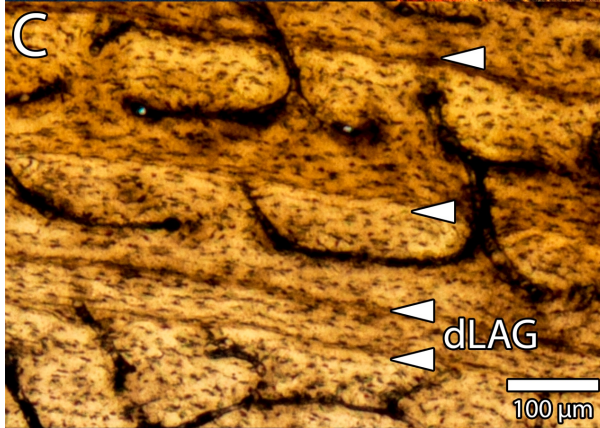
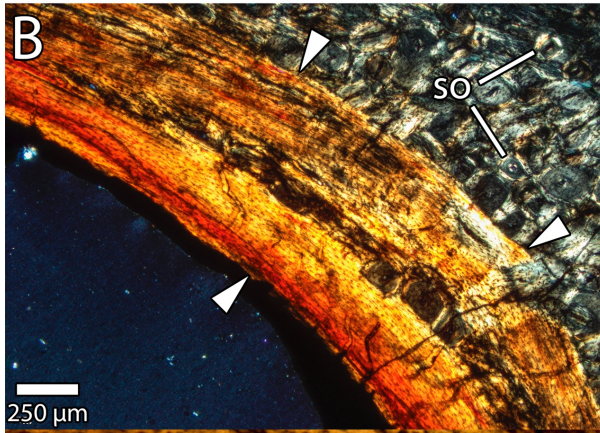
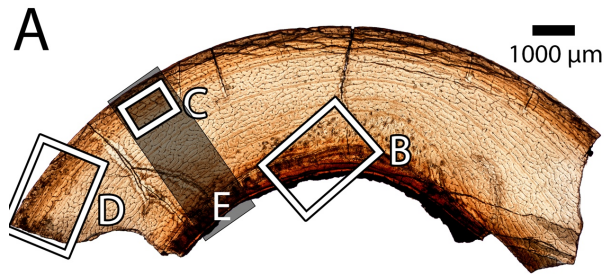
935 FIGURE 10. Other specimens referable to *Chirostenotes pergracilis* (TMP 2002.012.0103 and

936 TMP 1993.036.0181). A–C, Partial right ilium (TMP 2002.012.0103) in lateral (A), medial (B),

937 and ventral (C) views. D–F, Partial right tarsometatarsus (TMP 1993.036.0181) comprising

938 distal tarsals III and IV, and metatarsals II and IV in anterior (D), posterior (E), and proximal (F)  
939 views. **G**, closeup of proximal end of partial right tarsometatarsus (TMP 1993.036.0181),  
940 showing lack of fusion between metatarsals II and IV and between the distal tarsals and  
941 metatarsals. **Abbreviations:** : **ace**, acetabulum; **antr**, anterior ridge; **brf**, brevis fossa; **dt III**,  
942 distal tarsal III; **dt IV**, distal tarsal IV; ; **intf**, intermediate fossa; **isp**, ischiadic peduncle; **MT II**,  
943 metatarsal II; **MT IV**, metatarsal IV; **path**, pathological region; **pbp**, pubic peduncle; **pdp**,  
944 proximodorsal process; **postac**, postacetabular blade; **postf**, posterior fossa; **sra**, sacral rib  
945 attachment.  
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948

949 FIGURE 11. Osteohistology of *Chirostenotes pergracilis* (UALVP 59400). **A**, Transverse thin  
950 section of long-bone fragment from UALVP 59400 under normal light, showing the locations of  
951 other images. **B**, Detail of endosteal lamellae (arrows) and secondary remodeling along the  
952 endosteal margin and inner cortex under cross-polarized light. **C**, Detail of lines of arrested  
953 growth (arrows) in the outer cortex, showing doublet LAGs, under normal light. **D**, Detail of the  
954 periosteal region of the cortex, showing a high proportion of parallel-fibered bone and an  
955 annulus (arrow) associated with a line of arrested growth, under cross-polarized light. **E**, The  
956 cortex of UALVP 59400, showing resorption by the medullary cavity and endosteal lamellae  
957 (el), nine cyclical growth marks (arrows), and changes in the vascular pattern from reticular  
958 endosteally to longitudinal periosteally. **Abbreviations:** **dLAG**, doublet lines of arrested growth;  
959 **el**, endosteal lamellae; **so**, secondary osteon.

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962 **Word count: 8447**