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1	Description of the first definitive Corythosaurus (Dinosauria, Hadrosauridae)	
2	specimens from the Judith River Formation in Montana, USA and their	
3	paleobiogeographical significance	
4		
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21 ABSTRACT

22 The late Campanian Judith River Formation in northern Montana, USA has long been 23 recognized as a dinosaur-bearing rock unit. , eversince the first North American 24 dinosaur discovery from this unit. Despite the long study history of research in this 25 formation, most of the vertebrate fossils are represented by fragmentary remains, making precise taxonomic identifications difficult. Contrary to this, the partially 26 27 contemporaneous Dinosaur Park Formation, Alberta, Canada is known for its 28 tremendous fossil preservation, permitting rigorous studies of dinosaur diversity, 29 evolution, and biostratigraphy. Hadrosaurids comprise one of the most abundant 30 dinosaur clades in the Dinosaur Park Formation, but taxonomic affinities of hadrosaurid 31 specimens remain poorly understood in the Judith River Formation. Corythosaurus is 32 the most common hadrosaurid in the Dinosaur Park Formation and, to date,, and has 33 been restricted to this formation-to-date. This study reports the first definitive 34 Corythosaurus specimens from the Judith River Formation, which were discovered on 35 two private ranches in northern Montana. The attribution of the most complete skeleton to Corythosaurus is indicated by the combination of the following characters: wide 36 37 crest-snout angle, presence of premaxilla-nasal fontanelle, dorsoventrally expanded 38 nasal, laterally exposed ophthalmic canal of the laterosphenoid, and tall neural spines. A 39 second specimen preserves a largely ilium that can be positively identified as 40 Corythosaurus based on its associated skull, which is now in a private handhands due to 41 its association with a largely complete articulated skull. The specimens were recovered 42 from the Coal Ridge Member of the Judith River Formation, which is approximately 43 time equivalent to the Dinosaur Park Formation. Thus, the discovery of Corythosaurus 44 in the Judith River Formation extends the biogeographic range of this genus and

- 45 establishes a framework for future interformational biostratigraphic studies of Late
- 46 Cretaceous dinosaur faunas in North America. Further discoveries of the shared
- 47 dinosaur taxa between these two formations are expected to construct establish the an

48 interformational biostratigraphic framework.

- 50 Keywords: Late Cretaceous, Hadrosaur, Dinosaur, Judith River Formation
- 51

52 INTRODUCTION

53	Outcrops of the late Campanian Judith River Formation (JRF) exposed in
54	northern Montana, USA, have produced abundant vertebrate fossils. Since the discovery
55	of the first dinosaur in North America by Ferdinand Vandeveer Hayden in 1854 (Leidy,
56	1856), rigorous effort has been put into collecting vertebrate fossils from the JRF. As a
57	result of this, several relatively complete skeletal specimens have been collected and
58	described (Dodson, 1986; Prieto-Márquez, 2005; Murphy et al., 2006; Prieto-Marquez,
59	2007; Freedman, 2015; Mallon et al., 2016; Arbour and Evans, 2017) and have
60	contributed to understanding dinosaur evolution of the Late Cretaceous of North
61	America. Nonetheless, the majority of most of the dinosaur remains from the JRF are
62	represented by isolated or partial materials, hindering detailed taxonomic evaluations.
63	This pattern of fossil occurrences in the JRF is strikingly differents from those in the
64	partially contemporaneous Dinosaur Park Formation (DPF) of southern Alberta, where
65	a plethora of dinosaur remains have been discovered with various taphonomic modes,
66	including numerous complete skulls and skeletons (Eberth and Currie, 2005). Due to the
67	excellent preservation and rich fossil occurrences, the dinosaur faunas of the DPF have
68	been well-studied, and the the dinosaur biostratigraphy of dinosaurs has been
69	established at the species level (Ryan and Evans, 2005; Mallon et al., 2012; Cullen et
70	al., 2021), unlike the situation in the JRF.
71	Lambeosaurine hadrosaurids are one of the dinosaur groups that are-well-known
72	from the DPF, but <u>itstheir presence</u> remains ambiguous in the JRF. Currently, at least
73	five valid lambeosaurine taxa are known from the DPF: -(Corythosaurus casuarius,
74	C <u>.orythosaurus</u> intermedius, Lambeosaurus lambei, Lambeosaurus <u>L</u>. magnicristatus,

75 and Parasaurolophus walkeri). Corythosaurus is represented by over 25 relatively

76	complete skulls and skeletons (Currie and Russell, 2005) and is the most abundant
77	genus among the five lambeosaurine taxa (Ryan and Evans, 2005). Although abundant
78	in the DPF in the Dinosaur Park area, Alberta, Canada, there is no definitive record of
79	this genus from other time equivalent strata such as the JRF. Here, we report the first
80	well-preserved lambeosaurine skeleton identifiable to Corythosaurus, collected from the
81	JRF, northern Montana, USA. We also report a second, less complete specimen with a
82	complex provenance that can-be also be positively assigned to this genus based on an
83	articulated skull now lost to a private collection-also from the JRF of Montana. Both
84	specimens contribute significantly toward a more comprehensive understanding of the
85	dinosaur fauna of this formation and the paleobiogeographyical significance of
86	Campanian dinosaurs in North America.
87	
88	Institutional abbreviations
89	AMNH, American Museum of Natural History, New York; CMN, Canadian
90	Museum of Nature, Ottawa, Canada; MOR, Museum of the Rockies, Montana, USA;
91	NMMNH, New Mexico Museum of Natural History & Science, New Mexico, USA;
92	OUSM-FV, Okayama University of Science, Museum of Dinosaur Research, fossil
93	vertebrate collection, Japan; ROM, Royal Ontario Museum, Toronto, Canada; TMP,
94	Royal Tyrrell Museum of Palaeontology, Drumheller, Canada; UALVP, University of

95 Alberta Laboratory for Vertebrate Paleontology, Alberta, Canada; USNM, United States

5

96 National Museum, Smithsonian Institution, Washington, DC, USA.

97

98 MATERIAL AND METHODS

99	OUSM-FV-001, discovered and excavated from private land in Montana by the
100	commercial company Phoenix Fossils in 1990, was legally purchased by the
101	Hayashibara Museum of Natural Sciences in 1991. When the museum closed down-in
102	2015 subsequent toafter the bankruptcy of its parent company, OUSM-FV-001 was
103	transferred to the Okayama University of Science Museum. The pathological jugal and
104	the basisphenoid of OUSM-FV-001 were scanned using X-ray CT Latheta (LCT-200) at
105	the Okayama University of Science. The images were acquired at a voltage of 80kV and
106	a current of 0.5 mA. Both the pixel sizes and interslice thicknesses are 120 and 120 um
107	resulting in voxel sizes of 120 um (Supplemental Data S1). Photogrammetric models of
108	the partial nasals and the partial left premaxilla were made using Agisoft Metashape
109	Professional (Supplemental Data S2-S4). Selected measurements of OUSM-FV-001 are
110	provided as Supplemental Data S5.
111	ROM 77978 was also discovered and legally collected from a private ranch in
112	Hill County, Montana in 2015. The articulated skull and lower jaws were mounted and

sold into a private collection. Prior to the sale, the original skull was inspected firsthand
by DCE on January 29, 2016. The collector was not interested in obtaining the
associated postcranial elements, which were subsequently acquired by the Royal
Ontario Museum and accessioned as ROM 77978; and described here. Photographs, CT
scans, and a high-resolution surface scan together with a 3D print of the skull are also
reposited at the ROM. As part of the acquisition of ROM 77978, the landowner agreed

119 to allow scientific access to the field site in order for the ROM to gather detailed

120 contextual sedimentological and stratigraphic information and document the host

121 paleoenvironment. DCE and FF visited the quarry whe	here ROM 7/9/8 was collected on
---------------------------------------------------------	---------------------------------

- 122 August 10, 2017, along with the person who discovered and collected the specimen.
- 123
- 124 Systematic paleontology
- 125 Dinosauria Owen, 1842
- 126 Ornithischia Seeley, 1887
- 127 Lambeosaurinae Parks, 1923
- 128 Lambeosaurini Sullivan et al., 2011
- 129 Corythosaurus Brown, 1914
- 130 Corythosaurus sp.
- 131

132 Referred Specimen

133 OUSM-FV-001, a partial disarticulated skeleton including: partial left and right 134 premaxillae, a complete left and posterior two-thirds of the right maxillae, partial nasals 135 around the prefrontal-nasal fontanelle, a nearly complete left jugal and a partial right 136 jugal, posteromedial portions of both quadrates, partial right squamosal, partial left 137 postorbital, a left frontal, a right laterosphenoid, a left exoccipital other than missing the 138 paroccipital process, the main body of the basisphenoid, both dentaries, both 139 surangulars, both angulars, right splenial, partial axial elements, a nearly complete right 140 scapula, blade of the left scapula, both coracoids, nearly complete right sternum, a complete left humerus, nearly complete ulnae, nearly complete radiaeradii, both manus 141 142 mani missing a few phalanges, a nearly complete right ilium, both pubes with

143 incomplete anterior and posterior processes, nearly complete left ischium, nearly

144 complete left femur, both tibiae missing the distal ends, nearly complete fibulae, a 145 complete right astragalus, and both pedes missing a few phalanges (Fig. 1C). 146 147 Locality and horizon 148 OUSM-FV-001 was discovered on a private ranch approximately 8 km north of 149 Winifred, Montana (Fig. 1). The locality data is are cataloged in the OUSM collection 150 database and is available upon request. The specific locality data was initially 151 unavailable since the private company that sold this specimen provided only a rough 152 quarry location (within a 4 x 4 km range) and has since ceased operation. Recently, the 153 authors noted that the location of a historical quarry on property now leased by Triebold 154 Paleontology, Inc. nicknamed "Japanese dig" by local ranchers fell exactly within this 155 area. According to recollections of those ranchers as well as from-dates found on 156 beverage cans found aint the quarry by one author, a 1990 date for that excavation 157 matches information provided with OUSM-FV-001. Together with the fact that a nearly 158 complete lambeosaurine skeleton is extremely rare from the Judith River Formation, we 159 tentatively consider that OUSM-FV-001 was collected from this quarry. The OUSM-160 FV-001 quarry is located near the reference section of the Judith River Formation 161 (Rogers et al., 2016). Since the quarry is positioned no lower than the top 50 m of the badlands, and the Coal Ridge Member occupies the upper ~ 90m of the formation, the 162 horizon yielding the quarry likely belongs to the Coal Ridge Member. The quarry is 163 164 located approximately 8.5 km NNE of the Spiclypeus shipporum holotype locality (Mallon et al., 2016), and is also several hundred meters away from the "Ava" 165 166 centrosaur quarry, which was collected by Triebold Paleontology, Inc. 167

168 Description of OUSM-FV-001

169	Premaxilla. The anterior right premaxilla and a small portion of the left
170	premaxilla are preserved (Fig. 2C-F). The oral margin of the right premaxilla is slightly
171	lost damaged but has a gently rounded outline. Posteriorly, the posterodorsal process is
172	laterally everted to form a ridge that dorsally defines the external naris. Although
173	incomplete, the dorsal process of the left premaxilla is gently eurvedconcave unlike in
174	Parasaurolophus which is straight or convex. The posterodorsal end of the dorsal
175	process is curved posterodorsally (Fig. 2D; Data S4), suggesting a wide crest-snout
176	angle as in Corythosaurus and Hypacrosaurus, rather than Lambeosaurus in which the
177	dorsal process is recurved anteriorly to form an acute crest-snout angle less than 120
178	degrees (e.g., ROM 794, TMP 66.04.01)and forms a crest-snout angle of approximately
179	150 degrees along its dorsal margin (Fig. 2D). The angle is markedly wider than that of
180	Lambeosaurus, in which the crest-snout angle is generally less than 120 degrees in large
181	specimens (e.g., ROM 794, TMP 66.04.01). Instead, the wide crest-snout angle of
182	OUSM-FV-001 resembles that of Corythosaurus and Hypacrosaurus. Internal to the
183	curvature, a septum intercepts the dorsal half of the nasal passage to form an S-loop, as
184	in other members of Lambeosaurini (Evans et al., 2009). The posterodorsal process of
185	the left premaxilla is medially flat to contact its counterpart and dorsally grooved,
186	possibly due to postmortem deformation.
187	Maxilla. A complete left maxilla and the posterior two-thirds of the right
188	maxilla are mounted for display (Fig. 2A, B, G). Although the anterior process of the
189	maxilla is dorsally covered by the reconstructed premaxilla, a part of the premaxillary
190	shelf exposed at the anterior end is angled at approximately 15 degrees relative to the

191 ventral margin. The lateral exposure of the dorsal process of the maxilla is located

192 approximately at the midline of the maxilla. It is dorsoventrally tall and tapers to a point 193 dorsally. Ventral to the dorsal process, the lateral surface of the left maxilla bears two 194 large and two small foramina, whereas the right maxilla bears one large and four small 195 foramina. A well-developed ectopterygoid ridge emerges posterodorsal to the most 196 posterior foramen, where the maxilla and the ventral margin of the anterior process of 197 the jugal contact. The ectopterygoid ridge occupies approximately 40% of the maxilla 198 length. It is oriented nearly parallel to the alveolar margin of the posterior maxilla, and 199 is approximately 1.5 times as wide as the height of the maxilla anterior to the pterygoid 200 process. The alveolar margin of the maxilla is shallowly concave. It should be noted 201 that the alveolar margin of the left maxilla is slightly swollen at around the midpoint of 202 the maxilla. The swollen area has a rugose surface that resembles the lesion identified 203 on a Parasaurolophus maxilla (Bertozzo et al., 2020). Medially, the gently arched row 204 of alveolar foramina are is located dorsal to the midline of the maxilla. At the posterodorsal end of the maxilla, a small, triangular pterygoid process projects 205 206 dorsomedially. 207 The left maxilla bears approximately 38 tooth positions within the 298 mm long dental battery. Although the labial-most tooth is not preserved in most of the tooth 208 209 positions, the remaining positions suggest that the majority of the tooth row had two 210 functional teeth in life. All of the preserved maxillary teeth have a prominent primary ridge and lack other ridges. The primary ridge is straight or slightly curved and 211 212 positioned at the midline of each tooth. The mesial and the distal margins of the coronal apical half of each tooth have mammilliform denticles. 213 214 Nasal. The posterior regions of both nasals are preserved (Fig. 2H-K). Both 215 nasals retain the posterior margin of the premaxilla-nasal fontanelle (PNF), unlike in

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216	Hypacrosaurus in which the PNF is closed early in ontogeny (Brink et al., 2014). The
217	nasal is thick around the PNF and becomes thinner distal to the PNF, forming a
218	medially concave space for the lateral diverticulum (Evans et al., 2009). Since the
219	broken posterior end is still mediolaterally thick, the nasal is likely to extend further
220	posteriorly in life. The elongation of the dorsoventrally high nasal posterior to the PNF
221	differs from the nasal of Corythosaurus intermedius, in which the cranial crest is
222	anteroposteriorly shorter than Corythosaurus C. casuarius and forms a rod-like
223	structure at the posteroventral end of the crest (e.g., ROM 776, 777, 845, UALVP 13).
224	Although the nasal is incomplete, the preserved region is much taller than the strip-like
225	nasal of Lambeosaurus (e.g., ROM 1218). The lateral surface of the nasal is generally
226	flat except for the concavity dorsal to the posteroventral margin of the crest. The lateral
227	and the ventral surfaces meet at an acute angle, forming a sharp ridge. Ventrally, the
228	nasal bears a shallow concavity for the prefrontal contact (Fig. 2I, Data S3) as in
229	Corythosaurus, intermedius (e.g., ROM 776). The ventral surface of the crest, posterior
230	to the prefrontal contact, is smooth and shallowly concave (Data S2, 3), as in
231	Corythosaurus casuarius (e.g., AMNH FARB 5240)
232	Jugal. A nearly complete left jugal and a partial right jugal are mounted (Fig
233	2A, B, Fig. 3A). The anterior process of the jugal is expanded dorsoventrally as in other
234	lambeosaurines (Evans, 2010). The medial palatine process is inclined posterodorsally,
235	resulting in the <u>a</u> posteriorly located dorsal apex of the anterior process relative to the
236	ventral apex. The ventral margin of the jugal forms a posteriorly skewed deep arch,
237	although this region may be slightly eroded. The ventral flange of the jugal posterior to
238	the ventral arch is trapezoidal unlike the pointed ventral flange of <i>Hypacrosaurus</i>

239	altispinus (Evans, 2010). The anterodorsal and posteroventral margins of the posterior
240	process are oriented nearly parallel to each other.
241	In addition to the mounted two-jugals, a posterior process of another hadrosaurid
242	jugal is present (Fig. 3B, C). The length along the ventral margin of the infratemporal
243	fenestra is nearly 1.5 times as long as the other two jugals, whereas the posterior
244	constriction of the third jugal is only half as wide as the other two. While the differences
245	may indicate the third jugal belongs to a different hadrosaurid taxon, these anomalies
246	are more likely pathological. The lateral and medial surfaces of the anterior region are
247	more porous than the posterior region, and the anterior fracture surface is rugose and
248	bears multiple small cavities which may suggest resorption (Supplemental Data S1).
249	Quadrate. The posteromedial portions of both quadrates are preserved (Fig 2A,
250	B, Fig. 3D, E). The quadrate is gently bowed posteriorly as in <i>Corythosaurus</i> <u>C</u> .
251	casuarius (e.g., AMNH FARB 5240) and Lambeosaurus-L. lambei (e.g., ROM 1218), at
252	approximately 144 degrees. Dorsally, a faint squamosal buttress is present on the
253	posterior margin of the quadrate. The posterolateral spur is located at about one-third
254	the height of the quadrate. The posterior surface of the pterygoid wing is slightly
255	excavated to contact the posteroventral ramus of the pterygoid. The ventral end of the
256	quadrate is much wider mediolaterally than the anteroposterior length of the lateral
257	condyle, which is typical in hadrosaurids (Prieto-Márquez, 2010). The lateral and the
258	medial condyles form a deep groove on the posterior surface of the quadrate as in
259	C.orythosaurus intermedius (e.g., ROM 777) and Hypacrosaurus stebingeri (e.g., MOR
260	549). The medial condyle appears to be elevated compared to the lateral condyle,
261	although the medial condule is slightly damaged

262	Squamosal. The main body of the right squamosal is preserved (Fig. 3F). The
263	lateral surface of the postorbital process of the squamosal bears a contact surface for the
264	squamosal process of the postorbital. The contact surface extends dorsal to the apex of
265	the quadrate cotylus of the squamosal. The contact surface for the postorbital suggests
266	that the squamosal process of the postorbital is bifurcated, unlike in <i>Hypacrosaurus <u>H</u></i> .
267	altispinus (Evans, 2010). The preserved portion of the precotyloid process is shorter
268	than the anteroposterior width of the quadrate cotylus. In posterior view, the medial
269	process of the squamosal is dorsoventrally expanded as in other lambeosaurines (Evans,
270	2010).
271	Postorbital. The left postorbital is partially preserved (Fig. 3G). Although the
271 272	<i>Postorbital.</i> The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply
271 272 273	<i>Postorbital.</i> The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <i>Corythosaurus</i> - <i>C. casuarius</i> (Brown,
271 272 273 274	<i>Postorbital.</i> The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <i>Corythosaurus-C. casuarius</i> (Brown, 1914). Posterolateral to the depression, the dorsal surface of the base of the squamosal
271 272 273 274 275	<i>Postorbital.</i> The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <i>Corythosaurus C. casuarius</i> (Brown, 1914). Posterolateral to the depression, the dorsal surface of the base of the squamosal process forms a mediolaterally compressed protuberance as in some <i>Corythosaurus</i>
271 272 273 274 275 276	<i>Postorbital.</i> The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <i>Corythosaurus-C_casuarius</i> (Brown, 1914). Posterolateral to the depression, the dorsal surface of the base of the squamosal process forms a mediolaterally compressed protuberance as in some <i>Corythosaurus</i> (e.g., CMN 34825, ROM 776) and <i>Lambeosaurus</i> (ROM 1218) specimens. Although
271 272 273 274 275 276 277	Postorbital. The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <u>Corythosaurus C. casuarius</u> (Brown, 1914). Posterolateral to the depression, the dorsal surface of the base of the squamosal process forms a mediolaterally compressed protuberance as in some <i>Corythosaurus</i> (e.g., CMN 34825, ROM 776) and <i>Lambeosaurus</i> (ROM 1218) specimens. Although the squamosal process is incomplete, it makes a gentle curvature with the main body of
 271 272 273 274 275 276 277 278 	Postorbital. The left postorbital is partially preserved (Fig. 3G). Although the anteromedial process is partly missing, the base of the anteromedial process is deeply depressed, as in derived lambeosaurines such as <u>Corythosaurus C.</u> casuarius (Brown, 1914). Posterolateral to the depression, the dorsal surface of the base of the squamosal process forms a mediolaterally compressed protuberance as in some Corythosaurus (e.g., CMN 34825, ROM 776) and Lambeosaurus (ROM 1218) specimens. Although the squamosal process is incomplete, it makes a gentle curvature with the main body of the postorbital, unlike in Velafrons in which the squamosal process elevates abruptly

(Gates et al., 2007). Ventrally, the jugal process of the postorbital and the anteromedial

process form the semicircular posterodorsal margin of the orbit. The orbital margin has

a rugose texture. The anteromedial surface of the jugal process is shallowly concave.

(Fig. 3H-J). The anterior region of the frontal forms the anteroposteriorly striated

and is deeper laterally than medially. The contact surface for the nasal is angled at

contact surface for the nasal. The contact surface occupies the anterior half of the frontal

Frontal. The left frontal is nearly complete except for the posteromedial region

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2	286	approximately 20 degrees relative to the skull table. The anteromedial end of the nasal
2	287	contact surface does not reach the skull midline and forms a deep median cleft unlike in
2	288	Parasaurolophus (Evans et al., 2007). The posterolateral margin of the nasal sutural
2	289	surface is slightly everted to support the supracranial crest, unlike in large
2	290	parasaurolophines in which the nasal contact surface extends above the skull table
2	291	(Evans et al., 2007). Posteriorly, the skull table is slightly elevated to form a median
2	292	dome as in other lambeosaurines (Ostrom, 1961), although the region is partially
2	293	missing. Ventrally, the frontal bears three depressions: the olfactory depression, the
2	294	orbital depression, and the cerebral fossa. The cerebral fossa is the deepest among the
2	295	three depressions. Its posterodorsal surface shows vascular valleculae.
2	296	Laterosphenoid. A right laterosphenoid is missing its dorsolateral process (Fig.
2	297	<u>$3K_{4A}$</u>). The sulcus for the ophthalmic ramus (CN V ₁) is laterally fully open and forms a
2	298	horizontal groove as in Corythosaurus and Hypacrosaurus, but differs from
2	299	Lambeosaurus in which $\operatorname{CN} V_1$ is laterally covered by the laterosphenoid (Ostrom,
	300	1961; Evans and Reisz, 2007). <u>In lateral view</u> , <u>Relative to the ophthalmic sulcus</u> , tthe
	301	dorsal margin of the laterosphenoid where it contacts the parietal is nearly parallel or
	302	only slightly divergent from the line of the ophthalmic sulcus, resulting in a
	303	dorsoventrally short laterosphenoid where it contacts the parietal. This is consistent with
	304	the laterosphenoids of Corythosaurus C. casuarius (ROM 1933), Corythosaurus C.
	305	intermedius (ROM 776), and Hypacrosaurus-H. stebingeri (MOR 549). In contrast, it
	306	but differs from the laterosphenoid of <i>Parasaurolophus tubicen</i> (NMMNH P25100) _{15} in
	307	which the dorsolateral process is dorsally directed; therefore, the dorsal margin is
	308	strongly divergent from the ophthalmic sulcus. The In dorsal view, the dorsolateral
	309	process margin of the laterosphenoid strongly curves laterally as in <i>Prosaurolophus</i>

310	maximus (Horner, 1992), small Edmontosaurus annectens, and small Hypacrosaurus-H.
311	stebingeri (Takasaki et al., 2020).
312	Exoccipital. The left exoccipital is nearly complete other than its paroccipital
313	process (Fig. 3L, M4B, C). The occipital plate of the exoccipital is pierced by the
314	openings for the glossopharyngeal nerve (CN IX), the vagus nerve (CN X), and the
315	hypoglossal nerve (CN XII) (Ostrom, 1961; Evans et al., 2009). While CN IX and X are
316	associated close together laterally, they are well-separated medially. Dorsally, the
317	exoccipital has a horizontal shelf as the contact surface for the supraoccipital. The
318	horizontal shelf is slightly wider than its anteroposterior length. The laterodorsal margin
319	of the supraoccipital contact is medially overhung, indicating that the exoccipital
320	"locked" into the supraoccipital in articulation. Dorsolateral to the horizontal shelf, a
321	squamosal boss is present on the lateral half of the exoccipital unlike the indeterminate
322	lambeosaurine from the Prince Creek Formation, Alaska, in which the exoccipitals do
323	not participate in forming a squamosal boss (Takasaki et al., 2019).
324	Basisphenoid. The posterodorsal surface of the basisphenoid and the cultriform
325	process are missing (Fig. <u>3-4 ND-PF</u>). On the lateral surface of the long and massive
326	basipterygoid process, two ridges run proximodistally, emerging from the anteroventral
327	and posterodorsal margins of the foramen for the internal carotid artery at base of the
328	basipterygoid process. The alar process is bifurcated as in Amurosaurus riabinini
329	(Godefroit et al., 2004) and borders the anteroventral margin of the foramen for the
330	carotid artery. Posterodorsally, the basisphenoid is enlarged to form the sphenooccipital
331	tubera with the basioccipital. The basipterygoid processes meet at approximately 90
332	degrees in anterior or posterior views. A part of a small median protuberance is present
333	at the base of the basipterygoid processes as in derived lambeosaurines (Gates and

334 Sampson, 2007), although it is partly broken. Right lateral Tto-the median protuberance 335 is deeply incised on the right lateral side, and the surface surrounding the incision is 336 porous and slightly elevated, which may be pathological. 337 Mandible. Both dentaries are nearly complete, although most of the anterior 338 and the medial regions are reconstructed (Fig. 3Q4G-TJ). Anteriorly, the dentary is 339 curved ventrally, starting approximately from the level of the anteriormost reconstructed 340 tooth, at an angle of 159 degrees relative to the posterior half of the dentary. The 341 symphyseal process is markedly curved medially to contact the counterpart as in typical 342 lambeosaurines. The mediolateral width of the symphyseal process is slightly less than 343 twice the non-symphyseal anterior dentary mediolateral width. Posterior to the 344 symphyseal process, the edentulous diastema is shallowly sloped anteroventrally 345 relative to the main body of the dentary. The lateral surface of the main body of the 346 dentary is pierced by at least ten randomly -distributed foramina. Posteriorly, the 347 coronoid process is inclined anterodorsally at approximately 80 degrees relative to the 348 alveolar margin. The coronoid process is separated from the main body of the main 349 dentary body by a wide lateral shelf, typical for hadrosaurids. The dorsal apex of the 350 coronoid process is expanded anteroposteriorly, and its posterior surface is deeply 351 grooved to contact the anterodorsal process of the surangular. A fragmentary bone is cataloged as the left lateral process of the predentary, 352 but its detail is unavailable since it is embedded within the mounted skull (Fig. $\frac{3Q4G}{}$). 353 354 Both surangulars are preserved, although itstheir medial and anterior regions are also 355 embedded within the plaster<u>reconstruction</u>. The surangular is dorsoventrally flat at the 356 level of the glenoid. The posterior process of the surangular is strongly curved dorsally 357 and tapers into a point. Ventral to the surangular, the angular is visible in lateral view,

359	the specimen. The right splenial is mediolaterally thin and becomes taller anteriorly,
360	although the anterior end is missing.
361	Vertebrae. Eleven post-axial cervical, 17 dorsal, and 34 caudal vertebrae are
362	cataloged as containing original bone (Fig. 45), although most retain only either the
363	centrum or neural spine. The preserved vertebrae exhibit general conditions of
364	Hadrosauridae. Centra of the cervical vertebrae are opisthocoelous and slightly concave
365	lateroventrally. The articular facets of the prezygapophyses are subcircular in the
366	anterior cervicals and become mediolaterally wide and elliptical posteriorly. The
367	postzygapophyses are acutely divergent from each other in the anterior cervicals, while
368	the angle becomes wider in the posterior cervicals. Dorsally, the neural spine is small
369	and becomes slightly more prominent in the posterior cervical vertebrae. All of the
370	preserved dorsal centra are amphiplatyan. The best-preserved dorsal centrum is a middle
371	dorsal centrum cataloged as "D-4". Although the neural spine corresponding to this
372	centrum is missing, a neural spine of a posterior dorsal vertebra (cataloged as "D-16") is
373	2.3 times longer than the height of the middle dorsal centrum ("D-4"). Since dorsal
374	centra of hadrosaurids become higher posteriorly, the ratio between neural spine length
375	and centrum of the OUS specimen is likely to be much less than 2.3. The short neural
376	spine differs from Hypacrosaurus, which has a long neural spine relative to the centrum
377	(Lull and Wright, 1942; Evans, 2010). The neural spine of the posterior dorsal vertebra
378	is long and gracile, being 5.5 times higher than its anteroposterior width. Centra of the
379	caudal vertebrae are amphiplatyan and are much higher and wider than its lengths in the
1	

although this configuration is likely to be an artifact of misplacement upon mounting

358

anterior caudal vertebrae, while they become elongated posteriorly. The transverse

381 processes lie nearly horizontally, unlike the acutely angled transverse processes of the 382 dorsal vertebrae. Pectoral girdle. The right scapula is nearly complete other than the glenoid and 383 384 the distal end of the scapular blade (Fig. 5A6A). The acromion of the scapula is oriented 385 laterally, forming a flat dorsal surface of the anterior end of the scapula. Posterior to the acromion, a poorly developed deltoid ridge is present along the lateral surface of the 386 387 scapular blade. The scapula curves ventrally at the neck, although the curvature is less 388 prominent than Brachylophosaurus canadensis (Prieto-Marquez, 2007) and 389 Gryposaurus latidens (AMNH FARB 5465). The posterior end of the scapular blade is 390 at least 1.5 times as high as the neck, although the further detail is unavailable due to the 391 poor preservation. Both coracoids are nearly complete, exhibiting a slightly shorter 392 contact surface for the scapula compared to the length of the glenoid (Fig. 5B6B). The 393 anterodorsal margin of the coracoid is concave and ventrally continues to a well-394 developed biceps tubercle. Distally, the ventral hook of the coracoid is 0.64 times as 395 high as its width. The medioventral margin of the coracoid is rugose for cartilage 396 contact. The right sternum is also nearly complete, except that the medial portion of the 397 proximal plate is missing -(Fig. 5C6C). The sternum is "hatchet" shaped, as isn typical 398 for hadrosaurids. The proximal plate of the sternum is thickest along the anterolateral 399 and the posterolateral margins, while the proximal plate is thin medially. The proximal 400 plate of the sternum is shorter than the distal process. The distal process is nearly 401 straight and slightly expanded in the distal enddistally. 402 403 Forelimb. The left humerus is complete (Fig. 5D6D). The deltopectoral crest is

half as wide as its length and reaches approximately 60% of the humerus length. The

405	anterior and the ventral margins of the deltopectoral crest meet at approximately 103
406	degrees. The ulna and radius are at least 1.2 times as long as the humerus, although the
407	ulna is missing a small portion of the distal midshaft, and the radius is missing its
408	proximal end (Fig. 5E6E, F). The ulna is slender, being more than 10 times longer than
409	its anteroposterior width as in other members of Lambeosaurini. The right manus is
410	missing only the phalanges II-2, III-1, IV-2, IV-3, and V-3 (Fig. 5G6G). The manus
411	shows general hadrosaurid features. Metacarpals II-IV are long and slender, with the
412	metacarpal III approximately eight times as long as its width. The manual phalanges are
413	stout other than II-1, which is mediolaterally flat and elongated. The ungual phalanges
414	II-4 and III-4 are hoof-shaped. It may be worth noting that the ungual phalanx II-4 is
415	notably smaller than the ungual phalanx III-4 unlike some hadrosaurids (Brown, 1912;
416	Parks, 1922; Suzuki et al., 2004).
417	Pelvic girdle. The right ilium is missing the distal half of the preacetabular
418	process, the distal end of the postacetabular process, the supracetabular process, the
419	acetabular margin, and the ischial peduncle (Fig. 6A7A). The preacetabular process is

420 curved downward at its base relative to the dorsal margin of the main plate. A small

421 ridge develops along the ventral margin of the base of the preacetabular process.

Posterior to the acetabulum, the ischial peduncle is composed of two protuberances. The postacetabular process is oriented posterodorsally. It is mediolaterally compressed in the dorsal half, whereas the ventral half is expanded to form the brevis shelf that is facing medioventrally. Medially, the sacral ridge is positioned along the dorsal rim of the ilium. The left public is missing most of its prepublic and postpublic processes (Fig. 6B7B). A faint ridge is present on the lateral surfaces of the iliac and the ischial

peduncles along the acetabulum. The left ischium is missing its pubic process and the

distal end (Fig. 667C). The proximal tip of the iliac peduncle is deflected posteriorly.
The short obturator process is located close to the base of the pubic peduncle. Although
the distal end of the ischium is missing, the ischial shaft is distally divergent, suggesting
the presence of the ischial boot.

433 Hindlimb. The left femur is nearly complete, although it is mediolaterally compressed (Fig. 6D7D). Proximally, the lesser trochanter is small and fused to the 434 435 greater trochanter. Although the distal end of the femur is posteriorly curved, the 436 curvature is likely due to postmortem distortion. The fourth trochanter is proximally 437 skewed and subtriangular in lateral view with a smooth apex. The fourth trochanter is 438 positioned slightly distal to the midpoint of the femur. The left tibia is missing its distal 439 end (Fig. 6E7E). The cnemial crest of the tibia is much more developed than the medial 440 condyle and distally extends over the midpoint of the tibia. The left fibula is complete, 441 and both the proximal and the distal ends are expanded anteriorly (Fig. 6F7F). The 442 anterodorsal process of the astragalus is medially skewed offset, and its anterior surface 443 is shallowly excavated (Fig. 6676). No calcaneum is preserved. Although both pedes 444 are missing several pedal phalanges, they show general hadrosaurid features (Fig. 445 6G7G). Metatarsal III is approximately six times as long as its width. The pedal 446 phalanges III-2, III-3, IV-2, and IV-3 are disc-like and more than three times as wide as their lengths. The ungual phalanges are hoof-shaped and have a flat ventral surface. 447 448 449 Corythosaurus sp.

- 450
- 451 Referred Specimen

ROM 77978 includes the angular, left ilium, several mid-caudal vertebrae, and
associated fragments of ossified tendons. An incomplete articulated skull and lower
jaws preserved from the orbits forward was also associated with this material, but is
currently in private hands.

456

457 Locality and Horizon

458 ROM 77978 was excavated from strata of the upper Judith River Formation 459 (Coal Ridge Member) approximately 9 kilometers to the East of the town of Havre, Hill 460 County, Montana; ... The quarry is located in badlands along the Little Boxelder Creek, 461 on the west arm of a north-south oriented coulee system and on the east side of a large 462 dry island, approximately 3 km upstream from its merger with the Milk River 463 (Supplemental Fig. S18). Detailed locality data is on file at the Royal Ontario Museum. The quarry itself is located in a shallow saddle low in the local section, approximately 464 465 10 m from the valley floor and about 10 m below prairie level of the dry island. The 466 partially disarticulated but closely associated specimen occurred at the base of a coarse-467 grained channel sandstone unit approximately 60 cm in thickness-specimen. The host 468 stratum is in the fourth finding fining upward sequence in the immediate section where 469 the quarry is located. The paleochannel is estimated to have been about 1.8 m in depth, and paleochannel flow direction was approximately to the north. 470 471 The general area is largely undocumented from a stratigraphic perspective, with 472 the closest published measured sections on the north side of the Milk River at Havre 473 (Eberth and Hamblin, 1993; Arbour and Evans, 2017). However, continuous bedrock

- 474 exposures along Little Boxelder Creek to the exposures at the Milk River allow
- 475 identification of key surfaces and marker beds useful to infer the stratigraphic

476	occurrence of ROM 77978; In ascending order (see Supplemental Fig. S18): 1. a sharp
477	lithological contact marking the onset of light-colored, Inclined Heterolithic Strata-
478	dominated, channelized sandstones on top of a thick, paleosol-dominated interval, and
479	2. a laterally extensive, coaly-lignite rich interval. Observations carried out north of
480	Havre (i.e. at the Zuul crurivastator quarry, Arbour and Evans, 2017; FF pers.obs.)
481	suggest the sand-dominated deposits as are part of the Coal Ridge Member of the Judith
482	River Formation (Rogers et al., 2016), and the coaly interval at the top of the Coal
483	Ridge Member as the palaeoenvironomental equivalent of the Lethbridge coal zone
484	(Eberth and Hamblin, 1993) occurring at the top of the Coal Ridge Member.
485	Furthermore, the overall thickness of strata in the the area ROM 77978 area is
486	approximately 45 meters, allowing comparison with the measured composite section of
487	Eberth and Hamblin (1993) to the north of Havre, where they inferred the top of the
488	sequence to represent the southernmost occurrence of the Dinosaur Park Formation
489	(Belly River Group), including the base of the Lethbridge Coal Zone.
490	Based on comparisons to with the more complete stratigraphic sequence
491	exposed on the east side the coulee system, the ROM 77978 host stratum is located
492	approximately 40 meters below the base of the coaly deposits, and by extrapolation
493	approximately 10-20 meters lower than the interval that hosted the skeleton of <u>Zuul-Z.</u>
494	crurivastator (Arbour and Evans, 2017; FF pers.obs.). Therefore ROM 77978 is from
495	approximately the middle of the Coal Ridge Member of the Judith River Formation
496	(Rogers et al., 2016). ROM 77978 is therefore chronostratigraphically correlative with
497	the uppermost beds of the Oldman Formation in southern Alberta and the lower
498	Dinosaur Park Formation in Dinosaur Provincial Park (Belly River Group; see Eberth
499	and Hamblin, 1993; Eberth, 2005; Freedman Fowler and Horner, 2015; Arbour and

Formattato: Tipo di carattere: Corsivo Formattato: Tipo di carattere: Corsivo Evans, 2017; Fowler, 2017), thus to be considered, from a stratigraphic perspective,
representative of the *Corythosaurus*-dominated zone of Ryan and Evans (2005) and
Mallon et al. (2012).

503

504 Description of ROM 77978

505 Skull. Since the skull is in private hands, it will not be described in detail. 506 However, inspection of the original fossil before reconstruction by DCE, and the 507 reposited CT scans, surface scans, and 3D print in the ROM collection reveals key 508 morphological features that allow it to be identified as Corythosaurus (Supplemental 509 Fig S29). These include a cranial crest that extends supraorbitally and prescribes a 510 partial hemispherical semicircular outline in lateral view with its apex positioned above 511 the orbits. The preserved region of the crest is comprised of the premaxilla anteriorly 512 and nasal bones posteriorly with a prominent premaxilla-nasal fontanelle occurring 513 between these bones in the central region of the crest; below the PNF, the sinuous 514 dorsal margin of the posterolateral process of the premaxilla formed by a caudal lobe 515 that extends further dorsally than the rostral lobe, is diagnostic of Corythosaurus (Evans 516 et al., 2005). The wide crest-snout angle is approximately 142 degrees, which is within 517 the range of Corythosaurus (Evans, 2010). Since the caudal region of the crest is not preserved, and therefore the relative size and the shape of the caudal margin of the crest 518 519 are not known, ROM 77978 cannot be positively assigned to the specific level. The 520 morphology of the external narial opening, the shape of both the rostral process and free 521 ventral flange of the jugal, shape of the maxilla, and degree of deflection of the 522 edentulous rostral region of the dentaries are all within the range of variation of the

523 sample of Corythosaurus from the Dinosaur Park FormationDPF (e.g., Ostrom, 1961; 524 DCE pers.obs.; Evans, 2010) 525 Mandible. ROM 77978 includes an almost complete left angular (Fig. 7A10A-526 D), which is missing only the posteriormost region where it forms the base of the 527 retroarticular process. It is a thin, splint-like bone with contact surfaces for the dentary 528 anteriorly and laterally, the splenial medially, and the surangular posteriorly. The 529 exterior surface is smooth, and it is rugose internally at its posterior terminus, where it 530 contacts the surangular in an interdigitating joint. As preserved, the splenial has a 531 maximum length of 198 mm, with a maximum mediolateral width at the contact with 532 the surangular of 27 mm. 533 Vertebrae. Two relatively complete distal caudal vertebrae (Fig. 7E10E-J) are 534 typically hadrosaurid in morphology, with subhexagonal centrum faces, prominent 535 chevron facets ventrally, and highly raked neural spines. The vertebral bodies of both vertebrae lack facets or protrusions for caudal ribs, and the zygapophyses are present 536 537 but poorly developed. Based on comparisons with a mounted skeleton of Corythosaurus 538 C. intermedius (ROM 845), the largest caudal vertebra is approximately from 539 approximately caudal position 45, and the smaller vertebra is from position number 51. 540 The largest vertebra has a centrum length of 45mm, with an anterior width of 57 mm 541 and height of 47_mm. The incomplete, highly raked neural spine is 64 mm in length. 542 The left prezygapophysis is preserved and projects anteriorly from the neural arch, and 543 there are faint facets that represent postzygapophyses on the posterior base of the neural 544 spine. The smaller vertebra has a centrum length of 39.8 mm, anterior width of 47 mm 545 and height 41 mm. The incomplete highly raked neural spine is 63 mm as preserved.

546	Pelvic girdle. The almost complete right ilium is only missing the distal half of
547	the strongly downcurved preacetabular process (Fig. <u>10</u> -K, L). It has a total length of
548	855 mm in length as preserved, with a depth above the acetabulum of 198 mm. The
549	acetabular margin (175 mm in length) is a shallowly incised arch in the ventral margin
550	of the bone between the pubic and ischiac peduncle. The postacetabular process is
551	elongate, with a length of 270 mm and a dorsoventral height of 128 at its midpoint in
552	lateral view; the elongate postacetabular process (L/H ratio of 2.1) falls within the
553	expected range of Lambeosaurinae, and not Saurolphinae (Brett-Surman and Wagner,
554	2007). The dorsal margin of the ilium is strongly sigmoidal with <u>a</u> sharp dorsal
555	curvature over the supracetabular crest and a ventrally deflected preacetabular process,
556	as is characteristic of Lambeosaurinae (Horner et al., 2004; Brett-Surman and Wagner,
557	2007). The prominent supracetabular crest extends ventrally on the lateral surface of the
558	body to a point approximately halfway between the dorsal margin and the base of the
559	ischiac peduncle. The morphology the ilium in ROM 77978 is almost identical to the
560	type series of AMNH 5240 and AMNH 5338 and compares well to OUSM-FV-001,
561	with a strongly sigmoidal dorsal margin, robust suprailiac crest, and elongate
562	postacetabular process that falls into ilium type four of Brett-Surman and Wagner
563	(2007).
564	
565	DISCUSSION
566	Taxonomic assignments
567	Hadrosaurid remains have long been recognized from the Judith River
568	Formation, ever since the discovery of "Trachodon mirabilis" (Leidy, 1856). Since then,
569	three-two hadrosaurine taxa, Brachylophosaurus canadensis (Sternberg, 1953; Prieto-

570	Márquez, 2005; Murphy et al., 2006; Prieto-Marquez, 2007), (Sternberg, 1953; Horner,
571	1988; Prieto-Márquez, 2005; Murphy et al., 2006; Prieto-Marquez, 2007)
572	Brachylophosaurus goodwini (Horner, 1988), and Probrachylophosaurus bergi
573	(Freedman Fowler and Horner, 2015), have been described from the McClelland Ferry
574	Member of the Judith River Formation. On the other hand, lambeosaurines from the
575	formation are poorly understood, although their presence has been suggested by several
576	poorly preserved specimens. A maxilla and a squamosal (USNM 5457) collected from
577	Dog Creek (~10 km northwest of OUSM-FV-001 quarry) were originally reported as
578	"Hadrosaurus paucidens" (Marsh, 1889), and later renamed as "Lambeosaurus
579	paucidens" (Ostrom, 1964). While the bones clearly belong to Lambeosaurinae, further
580	their taxonomic status is now considered ambiguous due to their incompleteness
581	(Prieto-Márquez et al., 2006). Another such specimen, an isolated tooth (AMNH FARB
582	8527) from the Clambank Hollow Quarry (~25 km northwest of OUSM-FV-001 quarry)
583	originally identified as "Procheneosaurus altidens" (Sahni, 1972) obviously lacks
584	diagnostic features useful for identifying it to a specific genus. Another incomplete
585	specimen (pelvic and limb bones, AMNH FARB 3971) from near Cow Island, Montana
586	(~ 40 km northeast of the OUSM-FV-001 quarry) was reported as "Pteropelyx
587	grallipes" (Cope, 1889). Although this specimen was once treated as a senior synonym
588	of Corythosaurus (Brett-Surman, 1989), it was later considered a nomen dubium
589	(Horner et al., 2004). In addition to these dubious specimens, an ischium of an
590	indeterminate lambeosaurine is also known from a multitaxic bonebed from the
591	Careless Creek Quarry (Fiorillo, 1989; Prieto-Márquez and Gutarra, 2016).
592	Furthermore, lambeosaurine egg clutches and embryonic remains in ovo have been
593	described from the Judith River Formation (Horner, 1999), but their incompleteness and

594	embryonic nature obscure their taxonomic status. As such, while lambeosaurine	
595	material has continuously been reported from the Judith River Formation, their its	
596	taxonomic status remains ambiguous, largely due to the lack of taxonomically	
597	identifiable elements. The new lambeosaurine skeleton OUSM-FV-001 described in this	
598	study is therefore valuable in its reasonably well-preserved status, including part of the	
599	cranial crest, which makes genus-level taxonomic identification possible and	
600	contributes to a better understanding of the Judith River dinosaur fauna.	
601	OUSM-FV-001 is markedly different from the members of Parasaurolophini in	
602	the angled dorsal margin of the posterodorsal process of the premaxilla (Fig. 2D),	
603	although it is missing most of its cranial crest. The frontal also differs from	
604	Parasaurolophini in the absence of the posteriorly elongated nasal platform of the frontal	
605	and the long interfrontal suture relative to the frontal width (Evans et al., 2007). Instead,	
606	the presence of the PNF (Fig. 2H-K), the S-loop internal to the posterodorsal process of	
607	the premaxilla (Fig. 2E, F), and the dorsoventrally tall nasal suggest an affinity with	
608	Lambeosaurini. Among Lambeosaurini, OUSM-FV-001 differs from Hypacrosaurus in	
609	the absence of tall dorsal neural spines, which is a diagnostic character of	
610	Hypacrosaurus (Lull and Wright, 1942; Evans, 2010). The presence of a PNF also	
611	differs from Hypacrosaurus because the PNF of Hypacrosaurus closes at an early stage	
612	of ontogeny (Horner and Currie, 1994; Brink et al., 2014). Additionally, OUSM-FV-	
613	001 differs from Hypacrosaus-H. altispinus in the trapezoidal ventral margin of the	
614	ventral flange of the jugal and the bifurcated squamosal process of the postorbital	
615	(Evans, 2010). OUSM-FV-001 also differs from Lambeosaurus in the wide crest-snout	
616	angle and the laterally exposed ophthalmic canal of the laterosphenoid, both of which	
617	are diagnostic characters of Lambeosaurus (Evans and Reisz, 2007). OUSM-FV-001	

618	further differs from <i>Lambeosaurus</i> in the dorsoventrally tall nasal, which does not form	
619	the caudal process of the crest. OUSM-FV-001 differs from another member of	
620	Lambeosaurini, Magnapaulia (Prieto-Márquez et al., 2012) in the well-separated	
621	prezygapophyses of the proximal caudal vertebrae. OUSM-FV-001 also differs from the	
622	immature lambeosaurine Velafrons (Gates et al., 2007) in having the gently curved	
623	squamosal process of the postorbital (Fig. 3G). Although the incompleteness of OUSM-	
624	FV-001 hinders comparisons with the diagnostic characters of the Asian Lambeosaurini,	
625	the dentary of OUSM-FV-001 differs from those of Amurosaurus (Godefroit et al.,	
626	2004; Bolotsky et al., 2014) and Sahaliyania (Godefroit et al., 2008) in the absence of	
627	the strong ventral deflection of the anterior half of the dentary (Fig. 3Q4G-T]). OUSM-	
628	FV-001 also differs from another Asian Lambeosaurini lambeosaurine Olorotitan	
629	(Godefroit et al., 2012) in the relatively short diastema edentulous region of the dentary.	
630	Although the diagnostic character of Corythosaurus (i.e., large, bifurcated nasal; Evans	
631	et al., 2005) cannot be confirmed on OUSM-FV-001 due to incompleteness of the nasal,	
632	the unique set of characters listed above strongly suggests that OUSM-FV-001 is	
633	Corythosaurus. Whether OUSM-FV-001 is Corythosaurus-C. casuarius, Corythosaurus	
634	<u>C.</u> intermedius, or a distinct species cannot be determined at this point; thus, OUSM-	
635	FV-001 is referred to Corythosaurus sp	
636	ROM 77978, although less complete, also contributes important new	
637	information on the distribution of Corythosaurus. The identification of ROM 77978 is	
638	more straightforward but is scientifically hindered because the associated articulated	
639	skull is currently in private hands (Supplemental Fig. S29), although CT scans, surface	
640	scans, and photographs are reposited at the ROM. The skull clearly exhibits a number of	
641	diagnostic hallmarks of Corvthosaurus, including a rounded, helmet-like supracranial	

642	crest with its apex positioned above the orbits, a prominent premaxilla-nasal fontanelle
643	between the premaxilla and nasal bones in the central region of the crest, and most
644	importantly a sinuous dorsal margin of the posterolateral premaxillary process formed
645	by a caudal lobe that extends further dorsally than the rostral lobe. The shape of the
646	lateral process of ROM 77978 differs from that of <i>Hypacrosaurus <u>H</u>. stebingeri</i> , which
647	is dorsally arched (Brink et al., 2014). The wide crest-snout angle is approximately 142
648	degrees, which is within the range of Corythosaurus (Evans, 2010) and similar to H.
649	stebingeri (Brink et al., 2014), but outside that of most other lambeosaurines
650	(Lambeosaurus, Parasaurolophus and Hypacrosaurus-H. altispinus) known from
651	northern Laramidia during the Campanian-Maastrichtian (Evans, 2010). The caudal
652	region of the nasal is not preserved, and therefore the skull cannot be positively
653	assigned to either of the two known species of Corythosaurus, C. casuarius or C.
654	intermedius. Fortunately, the almost complete, well-preserved ilium of ROM 77978
655	(Fig. 7K <u>10K</u> , L) exhibits a distinctive suite of morphologies that allow its taxonomic
656	identification to the tribe Lambeosaurini, which is consistent with the identification of
657	the skull as Corythosaurus. The ilium of ROM 77978 is almost identical to the holotype
658	of Corythosaurus (AMNH 5240) and compares well to the plesiotype AMNH 5338 as
659	well as OUSM-FV-001;It has a strongly sigmoidal dorsal margin, large, pendant
660	suprailiac crest, and elongate postacetabular process and can clearly be categorized as
661	ilium type four of Brett-Surman and Wagner (2007). Recognizing that their five
662	hadrosaurid ilium morphotypes that in-form a morphocline and are not strictly
663	taxonomically 'useful', Corythosaurus is used to exemplify their fourth morphotype
664	(Brett-Surman and Wagner 2007). Although the ilia of the Lambeosaurini
665	Corythosaurus, Hypacrosaurus-H. stebingeri, and Lambeosaurus for example cannot be
I	

666	reliably distinguished from each other, the ilium in the ROM collections serves	
667	minimally as a detailed stratigraphic record of Lambeosaurini in Montana, which based	
668	on the reposited data for the skull can reasonably be identified as Corythosaurus sp.	
669	The taxonomic assignment of OUSM-FV-001 and ROM 77978 to	
670	Corythosaurus sp. in this study demonstrates that it is the first hadrosaurid specimen in	
671	the Coal Ridge Member identified below the subfamily level, and the first time a genus	
672	of lambeosaurine has been positively identified in the Judith River Formation as a	
673	whole.	
674		
675	Paleobiogeographical significance	
676	OUSM-FV-001 and ROM 77978 are notable for being the first Corythosaurus	
677	specimens from outside the DPF and for extending the biogeographic range of	
678	$Corythosaurus \sim 350$ km southward. The late Campanian vertebrate assemblage	
679	represented in the DPF is the most diverse terrestrial fauna of the Late Cretaceous,	
680	including over 50 dinosaurian taxa (Currie and Koppelhus, 2005). Hadrosaurids are the	
681	most abundant in the dinosaur fauna of the DPF (Dodson, 1983; Brinkman, 1990;	
682	Cullen and Evans, 2016), and Corythosaurus is the most common hadrosaurid genus in	
683	the formation (Ryan and Evans, 2005). Despite its abundance in the DPF, the	
684	geographical occurrence of Corythosaurus is-has been restricted to the Dinosaur	
685	Provincial Park area in Alberta, Canada, with one exception from Hilda, Alberta,	
686	approximately 80 km east of the park (Evans, 2002; Ryan and Evans, 2005). Although	
687	there are some ambiguities in the locality and stratigraphical data, the horizon yielding	
688	OUSM-FV-001 is supposed-likely to be within the Coal Ridge Member of the JRF,	
689	which is approximately time equivalent to the DPF (Rogers et al., 2016; Fowler, 2017).	

690	ROM 77978 can be definitively traced to the middle of the Coal Ridge Member of the	
691	JRF. Thus, OUSM-FV-001 and ROM 77978 are therefore at least	
692	penecontemporaneous to the Corythosaurus population from the DPF and likely	
693	represent an extension of its biogeographical range rather than a temporal habitat shift.	
694	The shared occurrence of Corythosaurus in the Coal Ridge Member of the JRF and the	
695	DPF is congruent with previous knowledge on of the paleoenvironment and the	
696	vertebrate fauna of these strata. The Coal Ridge Member of the JRF and the DPF are	
697	within a continuous clastic wedge, and both are interpreted to represent a wet and humid	
698	coastal plain environment along the Western Interior Seaway (Eberth and Hamblin,	
699	1993; Eberth, 2005; Rogers et al., 2016). Although there is only one shared dinosaur	
700	species (Mercuriceratops gemini, Ryan et al., 2014) between the two strata, plus some	
701	theropod teeth assigned to Dromaeosaurus albertensis, Troodon formosus, and Zapsalis	
702	abradens (Mallon et al., 2016), the terrestrial and freshwater fauna of the JRF and the	
703	DPF resemble each other as a whole compared to other Campanian formations in North	
704	America (Gates et al., 2010). Based on the limited biogeographical extent of	
705	Corythosaurus and other late Campanian megaherbivorous dinosaurs, it has been	
706	suggested that they have specific habitat preferences and/or limited environmental	
707	tolerances (Lehman, 2001). The extension of the Corythosaurus biogeographical	
708	distribution revealed in this study does not readily disagree with this idea because of the	
709	paleoenvironmental similarities of these two strata , <u>but Aa</u>dditional specimens and	
710	further taxonomic studies of megaherbivorous dinosaurs as well as the construction of a	
711	rigorous temporal framework of the time equivalent beds yet deposited in different	
712	environmental conditions, such as the more xeric Two Medicine Formation (Rogers,	
713	1990), are necessary to test this hypothesis.	

Formattato: Tipo di carattere: Corsivo

714	Even though our study demonstrates the paleobiogeographical extension of
715	Corythosaurus, the Corythosaurus remains of the genus in the JRF are still
716	overwhelmingly outnumbered by those in the DPF, which is potentially driven by the
717	limited temporal occurrence of Corythosaurus in the fossil record. Due to its rich fossil
718	record and intense research history, the stratigraphic distribution of dinosaur taxa in the
719	DPF is well understood (Ryan and Evans, 2005; Mallon et al., 2012; Cullen et al.,
720	2021), and Corythosaurus is only discovered inknown from the lower part of the
721	Dinosaur Park Formation (Centrosaurus-Corythosaurus faunal zone [Ryan and Evans,
722	2005], or the megaherbivore assemblage zone 1 [MAZ-1, Mallon et al., 2012]). If such
723	a tight stratigraphic distribution of megaherbivorous dinosaurs is also present in the
724	JRF, it is implied that the horizon directly comparable to the lower half of the DPF is
725	exposed in the outcrop around the OUSM-FV-001 and ROM 77978 localities, and so
726	would have high potential for new discoveries of dinosaurs known from these faunal
727	zones of the DPF. The faunal zones defined within the Dinosaur Park Formation at
728	Dinosaur Provincial Park have been shown to extend across a large regional area of
729	southern Alberta-(Evans et al., 2014)(Evans, 2002), and some ornithischian taxa such
730	as Centrosaurus apertus have been shown to occur in time equivalent portions of the
731	upper muddy unit of the Oldman Formation (Chiba et al., 2015), which is both
732	lithostratigraphically and chronostratigraphically correlative to the Coal Ridge Member
733	of the JRF. We therefore predict that many dinosaur taxa currently known only from the
734	Dinosaur Park Formation will be discovered in the upper Judith River Formation, which
735	will further elucidate paleobiogeographic and <u>32b</u> iostratigraphic comparisons across the
1 736	Western Interior Basin.

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l 757	

758 **REFERENCES**

Arbour VM, Evans DC. 2017. A new ankylosaurine dinosaur from the Judith River
Formation of Montana, USA, based on an exceptional skeleton with soft tissue
preservation. R Soc Open Sci 4:161086.

762	Bertozzo F, Manucci F, Dempsey M, Tanke DH, Evans DC, Ruffell A, Murphy E. 2020.
763	Description and etiology of paleopathological lesions in the type specimen of
764	Parasaurolophus walkeri (Dinosauria: Hadrosauridae), with proposed
765	reconstructions of the nuchal ligament. J Anat.
766	Bolotsky YL, Godefroit P, Bolotsky IY, Atuchin A. 2014. Hadrosaurs from the Far East:
767	Historical Perspective and New Amurosaurus Material from Blagoveschensk
768	(Amur Region, Russia). In: Hadrosaurs: Indiana University Press. p 315-331.
769	Brett-Surman MK. 1989. A revision of the Hadrosauridae (Reptilia: Ornithischia) and
770	their evolution during the Campanian and Maastrichtian. In: Graduate School of
771	Arts and Sciences. Washington, DC: George Washington University. p 1-272.
772	Brett-Surman MK, Wagner JR. 2007. Discussion of character analysis of the appendicular
773	anatomy in Campanian and Maastrichtian North American hadrosaurids-
774	variation and ontogeny. In: Carpenter K, editor. Horns and Beaks: Ceratopsian
775	and Ornithopod Dinosaurs. Bloomington: Indiana University Press. p 135-169.
776	Brink KS, Zelenitsky DK, Evans DC, Horner JR, Therrien F. 2014. Cranial morphology
777	and variation in Hypacrosaurus stebingeri (Ornithischia: Hadrosauridae). In:
778	Eberth DA, Evans DC, editors. Hadrosaurs: Indiana University Press. p 245-265.
779	Brinkman DB. 1990. Paleooecology of the Judith River Formation (Campanian) of
780	Dinosaur Provincial Park, Alberta, Canada: Evidence from vertebrate microfossil
781	localities. Palaeogeogr, Palaeoclimatol, Palaeoecol 78:37-54.
782	Brown B. 1912. The osteology of the manus in the family Trachodontidae. Bull Am Mus
783	Nat Hist 31:105-108.
784	Brown B. 1914. Corythosaurus casuarius, a new crested dinosaur from the Belly River
785	Cretaceous, with provisional classification of the family Trachodontidae. Bull Am

786 Mus Nat Hist 33:559-565.

787	Chiba K, Ryan MJ, Braman DR, Eberth DA, Scott EE, Brown CM, Kobayashi Y, Evans
788	DC. 2015. Taphonomy of a monodominant Centrosaurus apertus (Dinosauria:
789	Ceratopsia) bonebed from the Upper Oldman Formation of Southeastern Alberta.
790	Palaios 30:655-667.
791	Cope ED. 1889. Notes on the Dinosauria of the Laramie. The American Naturalist
792	23:904-906.
793	Cullen TM, Evans DC. 2016. Palaeoenvironmental drivers of vertebrate community
794	composition in the Belly River Group (Campanian) of Alberta, Canada, with
795	implications for dinosaur biogeography. BMC Ecol 16:52.
796	Cullen TM, Zanno L, Larson DW, Todd E, Currie PJ, Evans DC. 2021. Anatomical,
797	morphometric, and stratigraphic analyses of theropod biodiversity in the Upper
798	Cretaceous (Campanian) Dinosaur Park Formation. Can J Earth Sci 58:870-884.
799	Currie PJ, Koppelhus EB. 2005. Dinosaur Provincial Park: A Spectacular Ancient
800	Ecosystem Revealed. Bloomington: Indiana University Press.
801	Currie PJ, Russell DA. 2005. The geographic and stratigraphic distribution of articulated
802	and associated dinosaur remains. Dinosaur Provincial Park: A Spectacular

803 Ancient Ecosystem Revealed: Indiana University Press, Bloomington:537-569.

- Bodson P. 1983. A faunal review of the Judith River (Oldman) Formation, Dinosaur
 Provincial Park, Alberta. Mosasaur 1:89-118.
- Bodson P. 1986. Avaceratops lammersi: a new ceratopsid from the Judith River
 Formation of Montana. Proceedings of the Academy of Natural Sciences of
 Philadelphia 138:308-317.
- 809 Eberth DA. 2005. The geology. In: Currie PJ, Koppelhus EB, editors. Dinosaur Provincial

810	Park: A Spectacular Ancient Ecosystem Revealed. Bloomington and Indianapolis:
811	Indiana University Press. p 367-397.
812	Eberth DA, Currie PJ. 2005. Vertebrate taphonomy and taphonomic modes. In: Currie PJ,
813	Koppelhus EB, editors. Dinosaur Provincial Park, a Spectacular Ancient
814	Ecosystem Revealed. Bloomington: Indiana University Press. p 453-477.
815	Eberth DA, Hamblin AP. 1993. Tectonic, stratigraphic, and sedimentologic significance
816	of a regional discontinuity in the upper Judith River Group (Belly River wedge)
817	of southern Alberta, Saskatchewan, and northern Montana. Can J Earth Sci
818	30:174-200.
819	Evans DC. 2002. A juvenile Corythosaurus skull (Ornithischia: Hadrosauridae) from the
820	Dinosaur Park Formation, southeastern Alberta. Can Paleontol Conf Program
821	Abstracts 12:11-14.
822	Evans DC. 2010. Cranial anatomy and systematics of Hypacrosaurus altispinus, and a
823	comparative analysis of skull growth in lambeosaurine hadrosaurids (Dinosauria:
824	Ornithischia). Zoological Journal of the Linnean Society 159:398-434.
825	Evans DC, Forster C, Reisz R. 2005. The type specimen of Tetragonosaurus erectofrons
826	(Ornithischia: Hadrosauridae) and the identification of juvenile lambeosaurine.
827	In: Dinosaur Provincial Park: a spectacular ancient ecosystem revealed: Indiana
828	University Press Bloomington and Indianapolis. p 349-366.
829	Evans DC, McGarrity CT, Ryan MJ. 2014. A skull of Prosaurolophus maximus from
830	southeastern Alberta and the spatiotemporal distribution of faunal zones in the
831	Dinosaur Park Formation. In: Hadrosaurs: Indiana University Press. p 200-207.
832	Evans DC, Reisz RR. 2007. Anatomy and relationships of Lambeosaurus magnicristatus,
833	a crested hadrosaurid dinosaur (Ornithischia) from the Dinosaur Park Formation,
	36

834	Alberta. Journal of Vertebrate Paleontology 27:373-393.
835	Evans DC, Reisz RR, Dupuis K. 2007. A juvenile Parasaurolophus (Ornithischia:
836	Hadrosauridae) braincase from Dinosaur Provincial Park, Alberta, with comments
837	on crest ontogeny in the genus. Journal of Vertebrate Paleontology 27:642-650.
838	Evans DC, Ridgely R, Witmer LM. 2009. Endocranial anatomy of lambeosaurine
839	hadrosaurids (Dinosauria: Ornithischia): a sensorineural perspective on cranial
840	crest function. Anat Rec (Hoboken) 292:1315-1337.
841	Fiorillo A. 1989. The vertebrate fauna from the Judith River Formation (Late Cretaceous)
842	of Wheatland and Golden Valley Counties, Montana. The Mosasaur 4:127-142.
843	Fowler DW. 2017. Revised geochronology, correlation, and dinosaur stratigraphic ranges
844	of the Santonian-Maastrichtian (Late Cretaceous) formations of the Western
845	Interior of North America. PLoS ONE 12:e0188426.
846	Freedman EA. 2015. Evolution and ontoeny of hadrosaurs (Dinosauria: Ornithischia) in
847	the Judith River Formation (late Cretaceous: Campanian) of northcentral Montana
848	In: Earth Sciences: Montana State University. p 477.
849	Freedman Fowler EA, Horner JR. 2015. A new brachylophosaurin hadrosaur
850	(Dinosauria: Ornithischia) with an intermediate nasal crest from the Campanian
851	Judith River Formation of northcentral Montana. PLoS ONE 10:e0141304.
852	Gates TA, Sampson SD. 2007. A new species of Gryposaurus (Dinosauria:
853	Hadrosauridae) from the late Campanian Kaiparowits Formation, southern Utah,

- USA. Zoological Journal of the Linnean Society 151:351-376.
- 855 Gates TA, Sampson SD, De Jesús CRD, Zanno LE, Eberth D, Hernandez-Rivera R,
- 856 Martínez MCA, Kirkland JI. 2007. *Velafrons coahuilensis*, a new lambeosaurine
- 857 hadrosaurid (Dinosauria: Ornithopoda) from the late Campanian Cerro del Pueblo

858	Formation, Coahuila, Mexico. Journal of Vertebrate Paleontology 27:917-930.
859	Gates TA, Sampson SD, Zanno LE, Roberts EM, Eaton JG, Nydam RL, Hutchison JH,
860	Smith JA, Loewen MA, Getty MA. 2010. Biogeography of terrestrial and
861	freshwater vertebrates from the late Cretaceous (Campanian) Western Interior of
862	North America. Palaeogeogr, Palaeoclimatol, Palaeoecol 291:371-387.
863	Godefroit P, Bolotsky YL, Bolotsky IY. 2012. Osteology and relationships of Olorotitan
864	arharensis, a hollow-crested hadrosaurid dinosaur from the latest Cretaceous of
865	Far Eastern Russia. Acta Palaeontologica Polonica 57:527-560.
866	Godefroit P, Bolotsky YL, Van Itterbeeck J. 2004. The lambeosaurine dinosaur
867	Amurosaurus riabinini, from the Maastrichtian of Far Eastern Russia. Acta
868	Palaeontologica Polonica 49.
869	Godefroit P, Shulin H, Tingxiang Y, Lauters P. 2008. New hadrosaurid dinosaurs from
870	the uppermost Cretaceous of northeastern China. Acta Palaeontologica Polonica
871	53:47-74.
872	Horner JR. 1988. A new hadrosaur (Reptilia, Ornithischia) from the Upper Cretaceous
873	Judith River Formation of Montana. Journal of Vertebrate Paleontology 8:314-
874	321.
875	Horner JR. 1992. Cranial morphology of Prosaurolophus (Ornithischia: Hadrosauridae)
876	with descriptions of two new hadrosaurid species and an evaluation of hadrosaurid
877	phylogenetic relationships. Museum of the Rockies Occasional Paper 2:1-119.
878	Horner JR. 1999. Egg clutches and embryos of two hadrosaurian dinosaurs. Journal of
879	Vertebrate Paleontology 19:607-611.
880	Horner JR, Currie PJ. 1994. Embryonic and neonatal morphology and ontogeny of a new
881	species of Hypacrosaurus (Ornithischia, Lambeosauridae) from Montana and

882	Alberta. In: Carpenter K, Hirrsch KF, Horner JR, editors. Dinosaur Eggs and
883	Babies. New York: Cambridge University Press. p 312-336.

- 884 Horner JR, Weishampel DB, Forster CA. 2004. Hadrosauridae. In: Weishampel DB,
- Bodson P, Osmólska H, editors. The Dinosauria: Second Edition: University of
 California Press. p 438-463.
- Lehman TM. 2001. Late Cretaceous dinosaur provinciality. In: Tanke D, Carpenter K,
 editors. Mesozoic Vertebrate Live: Indiana University Press. p 310-328.
- 889 Leidy J. 1856. Notice of remains of extinct reptiles and fishes, discovered by Dr. FV
- Hayden in the Bad Lands of the Judith River, Nebraska Territory. Proceedings of
 the Academy of Natural Sciences of Philadelphia 8:72-73.
- Lull RS, Wright NE. 1942. Hadrosaurian dinosaurs of North America. Geological Society
 of America Special Papers 40:1-272.

894 Mallon JC, Evans DC, Ryan MJ, Anderson JS. 2012. Megaherbivorous dinosaur turnover

in the Dinosaur Park Formation (upper Campanian) of Alberta, Canada.
Palaeogeogr, Palaeoclimatol, Palaeoecol 350-352:124-138.

897 Mallon JC, Ott CJ, Larson PL, Iuliano EM, Evans DC. 2016. Spiclypeus shipporum gen.

898et sp. nov., a boldly audacious new chasmosaurine ceratopsid (Dinosauria:899Ornithischia) from the Judith River Formation (Upper Cretaceous: Campanian) of

900 Montana, USA. PLoS ONE 11:e0154218.

Marsh OC. 1889. Notice of new American Dinosauria. American Journal of Science s337:331-336.

- 903Murphy NL, Trexler DL, Thompson M. 2006. "Leonardo," a mummified904Brachylophosaurus (Ornithischia: Hadrosauridae) from the Judith River
- 905 Formation of Montana. In: Carpenter K, editor. Horns and Beaks Ceratopsians

and Orithopods. Bloomington an	d Indianapolis: Indiana University Press. p 117-
--------------------------------	--------------------------------------------------

- 907 133.
- 908 Ostrom JH. 1961. Cranial morphology of the hadrosaurian dinosaurs of North America.
 909 Bull Am Mus Nat Hist 122:33-186.
- 910 Ostrom JH. 1964. The Systematic Position of Hadrosaurus (Ceratops) paucidens Marsh.
 911 Journal of Paleontology 38:130-134.
- Owen R. 1842. Report on British fossil reptiles, Part 2. Report of the British Association
 for the Advancement of Science 11:60-204.
- Parks WA. 1922. *Parasaurolophus walkeri*: a new genus and species of crested
 trachodont dinosaur. University of Tronto Studies (Geological series) 13:1-32.
- Parks WA. 1923. *Corythosaurus intermedius*, a new species of trachodont dinosaur.
 University of Toronto Studies, Geological Series 15:1-57.
- 918 Prieto-Marquez A. 2007. Postcranial osteology of the hadrosaurid dinosaur
 919 Brachylophosaurus canadensis from the Late Cretaceous of Montana. In:
- 920 Carpenter K, editor. Horns and beaks. Ceratopsian and ornithopod dinosaurs.
- 921 Bloomington & Indianapolis, Indiana: Indiana University Press. p 91-115.
- Prieto-Márquez A. 2005. New information on the cranium of *Brachylophosaurus canadensis* (Dinosauria, Hadrosauridae), with a revision of its phylogenetic
 position. Journal of Vertebrate Paleontology 25:144-156.
- Prieto-Márquez A, Chiappe LM, Joshi SH. 2012. The lambeosaurine dinosaur
 Magnapaulia laticaudus from the Late Cretaceous of Baja California,
 northwestern Mexico. PLoS ONE 7:e38207.
- 928 Prieto-Márquez A, Gutarra S. 2016. The 'duck-billed' dinosaurs of Careless Creek
- 929 (Upper Cretaceous of Montana, USA), with comments on hadrosaurid ontogeny.

930 Journal of Paleontology 90:133-146.

931	Prieto-Márquez A, Weishampel DB, Horner JR. 2006. The dinosaur Hadrosaurus foulkii,
932	from the Campanian of the east coast of North America, with a reevaluation of
933	the genus. Acta Palaeontologica Polonica 51:77-98.
934	Rogers RR. 1990. Taphonomy of three dinosaur bone beds in the Upper Cretaceous Two
935	Medicine Formation of northwestern Montana: evidence for drought-related
936	mortality. Palaios 5:394.
937	Rogers RR, Kidwell SM, Deino AL, Mitchell JP, Nelson K, Thole JT. 2016. Age,
938	correlation, and lithostratigraphic revision of the Upper Cretaceous (Campanian)
939	Judith River Formation in its type area (North-Central Montana), with a
940	comparison of low- and high-accommodation alluvial records. The Journal of
941	Geology 124:99-135.
942	Ryan MJ, Evans DC. 2005. Ornithischian Dinosaurs. In: Dinosaur Provincial Park: A
943	Spectacular Ancient Ecosystem Revealed: Indiana University Press. p 312-348.
944	Ryan MJ, Evans DC, Currie PJ, Loewen MA. 2014. A new chasmosaurine from northern
945	Laramidia expands frill disparity in ceratopsid dinosaurs. Naturwissenschaften
946	101:505-512.
947	Sahni A. 1972. The vertebrate fauna of the Judith River Formation, Montana. Bull Am
948	Mus Nat Hist 147:321-412.

- Seeley HG. 1887. On the classification of the fossil animals commonly named Dinosauria.
 Proceedings of the Royal Society of London 43:165-171.
- 951 Sternberg CM. 1953. A new hadrosaur from the Oldman Formation of Alberta: discussion
 952 of nomenclature: Edmond Cloutier, Queen's Printer.
- 953 Sullivan R, Jasinkski S, Guenther M, Lucas S. 2011. The first lambeosaurin (Dinosauria,

954	Hadrosauridae, Lambeosaurinae) from the Upper Cretaceous Ojo Alamo
955	Formation (Naashoibito Member), San Juan Basin, New Mexico. New Mexico
956	Museum of Natural History and Science Bulletin 53:405-417.
957	Suzuki D, Weishampel DB, Minoura N. 2004. Nipponosaurus sachalinensis (Dinosauria;
958	Ornithopoda): anatomy and systematic position within Hadrosauridae. Journal of
959	Vertebrate Paleontology 24:145-164.
960	Takasaki R, Fiorillo AR, Kobayashi Y, Tykoski RS, McCarthy PJ. 2019. The first definite
961	lambeosaurine bone from the Liscomb Bonebed of the Upper Cretaceous Prince
962	Creek Formation, Alaska, United States. Sci Rep 9:5384.
963	Takasaki R, Fiorillo AR, Tykoski RS, Kobayashi Y. 2020. Re-examination of the cranial
964	osteology of the Arctic Alaskan hadrosaurine with implications for its taxonomic
965	status. PLoS ONE 15:e0232410.