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# Insights into iguanodontian dental architecture from an Early Cretaceous Chinese basal hadrosauriform maxilla (Ornithischia: Iguanodontia)

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Basal hadrosauriform iguanodontian dinosaurs have been invaluable towards understanding the evolution of the complex and highly efficient advanced hadrosauriform tooth battery dental system. Here we report a new basal hadrosauriform maxilla specimen IVPP V22529 - from the Dashuiguo Formation of Maortu, Nei Mongol, China that preserves a corrugated middle ventrolateral margin that differs from the straight and undulating ventral margins found in most iguandontian and non-iguanodontian dinosaurs. The uniqueness of this ventrolateral margin relates to a new dental structure - cementum 'jackets' that wrap about the labial sides of the teeth. To our knowledge this is the first time that cementum has been described migrated onto the tooth crowns of iguandontians (and other dinosaurs), but this trait is common amongst mammals. This dental morphology - seen in a similar form in the basal hadrosauriform Equijubus – therefore broadens our knowledge of iguanodontian maxillary anatomy and shows that the basal hadrosauriform dental system was more morphologically complex than previously thought. IVPP V22529 resembles maxillae specimens of Probactrosaurus gobiensis, a contemporaneous taxon known from the same locality in North China, in sharing an inferred subtriangular shape, a relatively flat lateral surface bearing a low row of foramina as well as similar-looking teeth. However, the presence of a unique corrugated middle ventrolateral margin in IVPP V22529, a low row of foramina on its lateral surface that also open anteriorly and increase in size posteriorly as well as a prominent medial shelf suggests that this specimen does not belong to P. gobiensis. However, these differences could conceivably be related to ontogenetic and sexual variation, which have not been fully documented in *P. gobiensis*. More detailed comparisons of IVPP V22529 and *Probactrosaurus* are also hampered by the missing posterior portion of IVPP V22529 as well as the missing anterior ramii in *Probactrosaurus* maxillae specimens. It is clear though that IVPP V22529 is different from the more advanced Northern Chinese hadrosauriforms *Bactrosaurus* and *Gilmoreosaurus*. The latter lack well-developed maxillary grooves on their medial shelves, unlike IVPP V22529, but all three taxa possess less-developed ones on the medial surfaces of the anteromedial processes of the anterior ramii. Different to IVPP V22529, Gilmoreosaurus

also has foramina that are more highly-positioned on the lateral surface of its maxilla as well as a row of larger and more circular 'special' foramina on its medial surface. Thus, at this time, IVPP V22529 is identified as a basal hadrosauriform and not as a new genus or as a new species of *Probactrosaurus*.

# Insights into iguanodontian dental architecture from an Early Cretaceous Chinese basal hadrosauriform maxilla (Ornithischia: Iguanodontia)

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### Abstract

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Basal hadrosauriform iguanodontian dinosaurs have been invaluable towards understanding 30 the evolution of the complex and highly efficient advanced hadrosauriform tooth battery 31 dental system. Here we report a new basal hadrosauriform maxilla specimen - IVPP V22529 32 - from the Dashuiguo Formation of Maortu, Nei Mongol, China that preserves a corrugated 33 middle ventrolateral margin that differs from the straight and undulating ventral margins 34 found in most iguandontian and non-iguanodontian dinosaurs. The uniqueness of this 35 36 ventrolateral margin relates to a new dental structure - cementum 'jackets' that wrap about 37 the labial sides of the teeth. To our knowledge this is the first time that cementum has been described migrated onto the tooth crowns of iguandontians (and other dinosaurs), but this trait 38 39 is common amongst mammals. This dental morphology - seen in a similar form in the basal hadrosauriform *Equijubus* – therefore broadens our knowledge of iguanodontian maxillary 40 anatomy and shows that the basal hadrosauriform dental system was more morphologically 41 complex than previously thought. IVPP V22529 resembles maxillae specimens of 42 Probactrosaurus gobiensis, a contemporaneous taxon known from the same locality in North 43 China, in sharing an inferred subtriangular shape, a relatively flat lateral surface bearing a 44 45 low row of foramina as well as similar-looking teeth. However, the presence of a unique corrugated middle ventrolateral margin in IVPP V22529, a low row of foramina on its lateral 46 surface that also open anteriorly and increase in size posteriorly as well as a prominent 47 48 medial shelf suggests that this specimen does not belong to P. gobiensis. However, these differences could conceivably be related to ontogenetic and sexual variation, which have not 49 been fully documented in P. gobiensis. More detailed comparisons of IVPP V22529 and 50

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*Probactrosaurus* are also hampered by the missing posterior portion of IVPP V22529 as well 51 as the missing anterior ramii in Probactrosaurus maxillae specimens. It is clear though that 52 IVPP V22529 is different from the more advanced Northern Chinese hadrosauriforms 53 54 Bactrosaurus and Gilmoreosaurus. The latter lack well-developed maxillary grooves on their medial shelves, unlike IVPP V22529, but all three taxa possess less-developed ones on the 55 medial surfaces of the anteromedial processes of the anterior ramii. Different to IVPP 56 57 V22529, *Gilmoreosaurus* also has foramina that are more highly-positioned on the lateral surface of its maxilla as well as a row of larger and more circular 'special' foramina on its 58 medial surface. Thus, at this time, IVPP V22529 is identified as a basal hadrosauriform and 59 60 not as a new genus or as a new species of Probactrosaurus.

# 63 Introduction

64 Maortu (Chow & Rozhdestvensky, 1960: = Maorty; 毛尔图) is a fossil locality of Early

65 Cretaceous age (Dashuiguo Formation: Barremian to Albian stages (Rozhdestvensky, 1966;

66 Rozhdestvensky, 1974; van Itterbeeck *et al.*, 2001; van Itterbeeck *et al.*, 2004) located

approximately half-way along China's northern frontier in Nei Mongol Autonomous Region (

内蒙古自治区), ~500km west of the provincial capital Hohhot (呼和浩特市) (Fig. 1).



Figure 1. The basal hadrosauriform maxilla IVPP V22529 was found in the Early Cretaceous
Dashuiguo Formation of Maortu, Nei Mongol, China (map produced from a Google Maps
image). Maortu is the type locality of the non-euhadrosaurian hadrosauriform

74 Probactrosaurus gobiensis (Rozhdestvensky, 1966).

75

76 Maortu is the type locality of three dinosaurs: the non-euhadrosaurian hadrosauriform

77 Probactrosaurus gobiensis (Rozhdestvensky, 1966) [this study follows the ornithopod

- 78 classification of Norman (2015)], the advanced non-carcharodontosaurine
- carcharodontosaurid *Shaochilong maortuensis* (Brusatte *et al.*, 2009; Brusatte *et al.*, 2010)
- 80 and the basal non-ankylosaurine ankylosaurid *Gobisaurus domoculus* (Vickaryous *et al.*,
- 81 2001). It is also the type locality of the trionychine trionychid turtle *Dongania maortuensis*
- 82 (Hans-Volker, 1999; Yeh, 1965 [See Vitek & Danilov, 2010 for taxonomic discussion]). In
- the summer of 2014 a team including several of the authors (MP, JRA, JM and SDB) visited
- Maortu (and its surrounding areas) where they recovered a variety of fragmentary and mostly isolated dinosaur and mammal bones now housed at the Institute of Vertebrate Paleontology

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- and Paleoanthropology (IVPP), Beijing. Amongst the largest of these bones is an isolated,
- crushed and posteriorly broken right iguanodontian maxilla (~12cm tall and 22cm long; Fig.
  3).
- 89
- 90 Early Cretaceous Asian iguanodontian maxillae are known from nine Chinese taxa
- 91 (Bactrosaurus johnsoni [Prieto-Márquez, 2011], Bolong yixianensis [Wu et al., 2010; Zheng
- 92 et al., 2013], Equijubus normani [You et al., 2003c], Jinzhousaurus yangi [Wang & Xu,
- 93 2001], Lanzhousaurus magnidens [You et al., 2005], Probactrosaurus gobiensis [Norman,
- 94 2002; Rozhdestvensky, 1966], *P. mazongshanensis* [Lü, 1997; Norman, 2002],
- 95 Shuangmiaosaurus gilmorei [You et al., 2003a] and Xuwulong yueluni [You et al., 2011]),
- two Japanese taxa (*Fukuisaurus tetoriensis* [Kobayashi & Azuma, 2003; Shibata & Azuma,
  2015] and *Koshisaurus katsuyama* [Shibata & Azuma, 2015]), two Kazak taxa (*Altirhinus*)
- 2015] and *Koshisaurus katsuyama* [Shibata & Azuma, 2015]), two Kazak taxa (*Altirhinus kurzanovi* [Norman, 1998] and *Batyrosaurus rozhdestvenskyi* [Godefroit *et al.*, 2012]) and a
  Thai specimen identified to a higher taxonomic level (*Siamodon nimngami*; Buffetaut &
  Suteethorn, 2011; *nomen dubium*: Norman, 2015). Comparisons between IVPP V22529 and
  the aforementioned taxa (Table 1) identifies IVPP V22529 as a non-euhadrosaurian
  hadrosauriform based on the presence of at least two replacement maxillary crowns and the
  absence of a single median primary ridge on the teeth (see Description and Comparison and
  Discussion). The presence of marginal denticles comprising of parallel ledges with single
  - rows of ~6 relatively large mammillae suggests that IVPP V22529 is a basal hadrosauriform.

Taxon	Specimen number	Material	References
China			
IVPP V22529	IVPP V22529 (field number: JLT 20140622-1)	Partial right maxilla (posterior ramus missing)	This study (Pittman <i>et al.</i> 2015)
Bactrosaurus	AMNH 6553	Adult/subadult left	Prieto-Márquez,
johnsoni*^	(holotype)	maxilla	2011: Figs. 7, 8
	AMNH 6390-6393,	Juvenile left maxillae	Prieto-Márquez,
	6514	(AMNH 6393 is a partial specimen)	2011: Figs. 11, 12
	AMNH 6388, 6389,	Juvenile right	Prieto-Márquez,
	6583	maxillae	2011: Figs. 9, 10
	3 SBDE 1	?	Godefroit et al., 1998
Bolong yixianensis*	YHZ-001	Left maxilla	Wu <i>et al.</i> , 2010; Wu & Godefroit, 2012: Figs. 19.2, 19.3
	ZMNH-M8812	Juvenile left and right maxillae	Zheng <i>et al.</i> , 2013: Figs. 2, 4-6
Equijubus normani	IVPP V12534 (holotype)	Complete articulated maxilla exposed on both the right and left lateral sides.	You <i>et al.</i> , 2003c: Fig. 1
Jinzhousaurus yangi	IVPP V12691 (holotype)	Complete articulated maxilla exposed on its left lateral side; medial side embedded in matrix.	Wang & Xu, 2001: Figs. 1, 2; Barrett <i>et</i> <i>al.</i> , 2009: Fig. 1
Lanzhousaurus	GSLTZP01-001	Isolated maxillary	You et al., 2005: Fig.

magnidens* Probactrosaurus gobiensis	(holotype) PIN 2232/9-2*	teeth Partial right maxilla (missing anterior ramus)	2A-H Norman, 2002: Fig. 5; Rozhdestvensky, 1966
	PIN 2232/10-2*	Partial right maxilla (missing anterior ramus as well as teeth from anterior and posterior aveolar slots)	Norman, 2002: Fig. 5; Rozhdestvensky, 1966
P. mazongshanensis	IVPP V1134.10-15 **	Isolated maxillary teeth	Lü, 1997; Norman, 2002: Fig. 4A
Shuangmiaosaurus gilmorei* <sup>#</sup>	LPM 0165	Left maxilla	You <i>et al</i> ., 2003a: Fig. 1
Xuwulong yueluni*	GSGM-F00001	Both maxillae	You <i>et al.</i> , 2011: Figs. 2, 3
Kazakhstan			
Altirhinus kurzanovi* Batyrosaurus rozhdestvenskyi*	PIN 3386/7 AEHM 4/1	Both maxillae ~30 maxillary teeth	Norman, 1998: Fig. 6 Godefroit <i>et al.</i> , 2012: Fig. 20.10C, D
Japan			
Fukuisaurus tetoriensis*	FPDM-V-40-1 (holotype)	Right maxilla	Kobayashi & Azuma, 2003: Fig. 6
	FPDM-V-40-5	Left maxilla	Kobayashi & Azuma, 2003: Fig. 2C-E
	FPDM-V-40-13	Isolated left maxillary tooth	Kobayashi & Azuma, 2003
Koshisaurus katsuyama*	FPDM-V9079	Right maxilla	Shibata & Azuma, 2015: Fig. 3
Thailand Siamodon nimngami* (nomen dubium: Norman, 2015	PRC-4	Left maxilla	Buffetaut & Suteethorn, 2011: Fig. 1; Norman, 2015 ( <i>nomen dubium</i> )

\*Taxa studied from the literature only; \*\*Specimens absent from host collection; ^Upper
Cretaceous taxa that were also found in Nei Mongol; <sup>#</sup>Suggested to be an Upper Cretaceous
taxon by You *et al.* (2003a) based on biostratigraphic evidence.

- 111 Table 1. Early Cretaceous Asian iguanodontian dinosaur maxillae studied.
- List of Early Cretaceous Asian iguanodontian maxillae used to describe IVPP V22529.
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# 115 Materials and Methods

IVPP V22529, an isolated partial right iguanodontian maxilla. This specimen was excavated, studied
 and described using standard palaeontological methods, in accordance with a fossil excavation permit
 (14-0620-JLT) obtained from the Department of Land and Resources, Nei Mongol, China.

#### Locality and Horizon 121

- Maortu (毛尔图), "Women's shoe" (妇女鞋) sublocality (40°12.109' N 105°42.957' E); 122
- ~60km north of Jilantai lake (吉兰泰盐湖), Alxa Left Banner, Alashan League, Nei Mongol, 123
- China (Figs. 1, 2); Dashuiguo Formation, Barremian to Albian, Early Cretaceous 124
- (Rozhdestvensky, 1966; Rozhdestvensky, 1974; van Itterbeeck et al., 2001; van Itterbeeck et 125 al., 2004).
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128 Figure 2. Stratigraphic log showing the position of IVPP V22529 in the local rock succession 129 at the "Women's shoe" (妇女鞋) sublocality (40°12.109' N 105°42.957' E). Grain size 130 abbreviations: c, clay; s, silt; fs, fine sand; ms, medium sand; cs, coarse sand; gt, grit; gv, 131 gravel. 132

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IVPP V22529 was recovered from the lower part of the exposure at the "Women's shoe" 134 sublocality (Bed 7 in Fig. 2) (40°12.109' N 105°42.957' E) located ~60km north of Jilantai 135 lake in the Alxa Left Banner of the Alashan League of Nei Mongol, China. The specimen 136 137 was found ~13 metres below where the Early Cretaceous succession is capped by recent gravel washout material. The sediments comprise vari-coloured beds (red-purple, pale green, 138 olive green and cream) that are typically 20–100 cm thick. The grain size is mostly clay or 139 silt, but coarse sands and grits are encountered. Some units show cross-bedding (Bed 3 in Fig. 140 2), but most are devoid of any internal layering, apart from a few that fine upwards. The 141 depositional setting is inferred to be a lake margin, the cross-bedded horizons probably 142 marking a time when a stream or small river, quite probably ephemeral, was discharging into 143 the system. 144

#### **Description and Comparison** 147

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IVPP V22529 is an isolated right iguanodontian maxilla that is missing its posterior ramus and 149

has a broken anterior one (Fig. 3). This section first describes the specimen's dentition and 150

compares it with other iguanodontians because this portion of the maxilla contains the most 151 diagnostic and unique information in this specimen. Then, the maxillary body will be

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153 described and compared with other iguanodontians.

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155 Figure 3. IVPP V22529 in lateral view showing the missing posterior ramus and broken 156 anterior one, the dorsal process and lateral surface, the labial view of the tooth row and the 157 corrugated middle ventrolateral surface formed by an unusual cementum 'jacket' tooth 158 159 morphology.

- 160 Maxillary Dentition 161 162
- 163 Tooth position count
- 164



Figure 4. Tooth row in ventral view showing the specimen's partial and heavily worn teeth, its unworn teeth as well as its missing ones.

IVPP V22529 preserves a partial tooth row (Figs. 2, 3) comprising of more than 18 vertical tooth positions, as indicated by 14 *in situ* teeth, an empty aveolar socket and a row of at least three empty parallel aveolar sockets at the anterior end of the maxilla (although the empty sockets may each have accommodated more than one tooth (Norman, 2002)). Without the posterior portion of the maxillary tooth row and no associated complete dentary row to estimate the number of vertical maxillary tooth positions, the latter is uncertain.

Nevertheless, by comparison, the non-euhadrosaurian hadrosauriform *Probactrosaurus* (PIN 2232/9-2, /10-2; Norman, 2002) preserves ~17 vertical maxillary tooth positions out of an estimated total of 22+ positions (22-23 positions estimated by Norman (2002); 23 or more total positions estimated by Rozhdestvensky (1966)). The 'iguanodontoid' styracosternan Jinzhousaurus is estimated to have fewer maxillary teeth than Probactrosaurus with only ~15-16 teeth [IVPP V12691, Barrett et al., 2009: Fig. 3C]), as in the hadrosauriform Koshisaurus which has 19 vertical tooth positions [FPDM-V907, Shibata & Azuma, 2015: Figs. 3, 8]. The basal non-euhadrosaurian hadrosauriform *Altirhinus* may have had a slightly 183 higher vertical tooth position count than Probactrosaurus as Norman (1998) estimated 26 184 positions on the basis of the 24 positions present in its dentary (PIN 3386/7, Norman, 1998: 185 Fig. 16). However, Altirhinus only preserves direct evidence of 21 vertical maxillary tooth 186 positions (right maxilla of PIN 3386/7; Norman, 1998: Fig. 6). Given the uncertain tooth 187 position count in IVPP V22529 it might even be possible that it has a high tooth position 188 count as in the basal non-euhadrosaurian hadrosauriform Eolambia (32 positions in the left 189 maxilla of CEUM 9758; Kirkland, 1998: Fig. 4A-C). Amongst iguanodontians the number of 190 maxillary teeth appears to increase during ontogeny (Horner et al., 2004; Hübner & Rauhut, 191 2010; Zheng et al., 2013), but the relatively large size of the maxilla fragment suggests that 192 IVPP V22529 probably has close to its maximum number of vertical tooth positions. 193

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- Replacement crowns 195

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Figure 5. IVPP V22529 appears to be a hadrosauriform iguandontian because it has at least two replacement teeth (Norman, 2015). This is indicated by a heavily worn tooth (right) that is supported by a replacement tooth, which is adjacent to a tooth that had just started to wear before the animal died.

201 One replacement crown is observed in five positions along the tooth row (Fig. 4), but the 202 preservation of the socket walls prevents the total number of replacement crowns from being 203 204 determined e.g. two replacement crowns are revealed by the broken posterior portion of the right medial wall of Altirhinus PIN 3386/7 (Norman, 1998: Fig. 6). However, there appears to 205 be indirect evidence of at least two replacement teeth in IVPP V22529 because one heavily 206 worn tooth is supported by a replacement tooth and anterior to it there is a taller erupted tooth 207 that had only just started to be worn prior to the animal's death (Fig. 5). The presence of at 208 least two replacement crowns implies that IVPP V22529 is a hadrosauriform iguandontian 209 (Norman, 2015: character 54, state 1) - non-hadrosauriform iguanodontians have one 210 211 functional crown supported by only one replacement crown (Norman, 2015: character 54, state 0)). 212 213

214 Denticles



Figure 6. Six heavily worn teeth in IVPP V22529 shown in ventromedial view.



Figure 7. The teeth of IVPP V22529 are dominated by a well-developed distally-offset primary ridge and lack any subsidiary (accessory) ridges. The unworn teeth of IVPP V22529 220 show marginal denticles comprising of parallel ledges with single rows of ~6 relatively large

- 221 mammillae. The latter suggests that IVPP V22529 is a basal hadrosauriform. 222
- 223

Six of the teeth preserved in IVPP V22529 have well-developed wear facets (Fig. 6) that are 224 up to 2cm shorter vertically than the tallest of the three teeth with slightly worn tips (the latter 225 are presumed to have erupted not long before the animal's death). There is a small fragment 226 of a seventh worn tooth located in the most posteriorly preserved position along the tooth row 227 (Fig. 6). One of the extensively worn teeth as well as all of the slightly worn or non-worn 228 teeth have marginal denticles comprising of parallel ledges with single rows of ~6 mammillae 229 230 (Fig. 7). Styracosternan iguanodontians have marginal denticles on both their maxillary and dentary teeth that form ledges with mammillations (Norman, 2015: character 58, state 2), but in IVPP V22529 the mammillae are comparatively large suggesting that it is a basal hadrosauriform as more advanced hadrosauriforms have smaller mammillae.

# Primary and subsidiary (accessory) ridges

In labial view, the enamelled surface of the crown is narrow and appears lozenge-like (elongated and asymmetrically diamond-shaped), as in *Probactrosaurus* (Norman, 2002). Altirhinus (Norman, 1998) and Iguanodon (Norman et al., 1987) [Fig. 7]. The crown's asymmetry is indicated by the distal offset of an enlarged primary ridge relative to the tooth's mid-line (a feature diagnostic of Iguanodontia (Norman, 2015: character 68, state 1) which includes Probactrosaurus, Altirhinus and Jinzhousaurus (Norman, 1998; Norman, 2002; Wang & Xu, 2001)) and the anterior position of the shoulder of the crown margin (as in Altirhinus (PIN 3386/7, Norman, 1998: Fig. 21B) and Jinzhousaurus (IVPP V12691, Wang & Xu, 2001) [Fig. 7]. The absence of a single median primary ridge indicates that IVPP V22529 is not a euhadrosaurian iguanodontian (Norman, 2015: character 68, state 3). As in *Probactrosaurus*, there appears to be little evidence of subsidiary ridges (Norman, 2002) [Fig. 7], unlike in Altirhinus (Norman, 1998: Fig. 21B) and Jinzhousaurus (IVPP V12691, Wang & Xu, 2001) where there is one anterior to the primary ridge and unlike in the styracosternan iguanodontian Lanzhousaurus (GSLTZP01-001; You et al., 2005: Fig. 2A) where the primary ridge is flanked by several subsidiary ones. There is no lingual ridge on the maxillary tooth crowns of IVPP V22529, so this ridge still appears to be an autapomorphy of Koshisaurus (FPDM-V907, Shibata & Azuma, 2015: Figs. 3C, 8B, 9B]). It is worth mentioning that the two isolated near-complete maxillary teeth (IVPP V1134.10) of 253 Probactrosaurus mazongshanensis Lü, 1997 - not a junior synonym of Probactrosaurus 254 gobiensis (Norman, 2002) - could not be located at the IVPP in May 2015 for direct 255 comparison with IVPP V22529. However, Lü (1997) noted that these teeth have a large 256 highly-developed primary ridge and his figure of one of them (his Figure 4) shows very 257 similar morphological traits to the maxillary teeth of IVPP V22529. 258

# 259260 Tooth root morphology

Relatively straight but poorly preserved tooth roots are exposed on the broken posteromedial surface of IVPP V22529. One portion of an exposed tooth root appears to be longitudinally

surface of IVPP V22529. One portion of an exposed tooth root appears to be longitud
 grooved (a synapomorphy of Iguanodontia (Norman, 2015: character 59, state 1 –

- 264 DELTRAN)), but this observation is equivocal owing to the root's poor preservation.
- Hadrosauromorpha is characterised by highly angular-sided roots (hexagonally prismatic)
- that relate to close packing of the teeth in a functionally integrated multi-tooth magazine
- 267 (Norman, 2015: character 59, state 2). Given the poor state of tooth root preservation in IVPP
- V22529, the presence of the latter in the specimen cannot be excluded at present.
- 269



Figure 8. In labial view, the five middle maxillary crowns preserve a bone-like sheath over their base. This appears to be cementum owing to its rugose texture and the absence of the fibres expected in ossified periodontal ligaments. This cementum 'jacket' morphology - which is the first to be described amongst dinosaurs to our knowledge – originates within the tooth socket and extends below the crenulated ventral margin of the maxilla to form a corrugated ventrolateral surface created by the grooves that separate each 'jacket'.

278 Below the five middle maxillary crowns at the broken posterior end of the specimen, the 279 labial side of each tooth appears to be tightly enveloped by smooth to a slightly rugose material (Fig. 8). These bone-like sheaths do not form a continuous surface and appear 280 separate from the walls of the tooth socket. They extend beyond the crenulated ventral 281 margin of the lateral surface, but begin beneath the socket walls themselves as revealed 282 through a broken portion of the wall (Fig. 8). The separation of this structure from the tooth 283 284 socket walls and its non-uniform roughened texture suggests that it is cementum (Fig. 8). Cementum with similar textural characteristics has been identified in *Probactrosaurus* 285 (Norman, 2002) and other iguanodontians; this texture is presumably associated with 286 ligamentous scarring on the tooth root. However, the unknown structure could conceivably be 287 ossified periodontal ligaments as these bind the tooth root to its socket. However, this 288 hypothesis is poorly supported as no structures resembling ligamental fibres were observed in 289 IVPP V22529. If these fibres were observed they should also show differences in orientation 290 along the length of the ligament. The lack of pathologies on the 'jackets' suggests that tooth 291 292 eruption was probably relatively smooth and unhindered. 293



Figure 9. Laser-stimulated fluorescence imaging (LSF; Kaye *et al.*, 2015) of IVPP V22529 shows that the unknown structure has similar green fluorescence colours to dentine and maxillary bone. However, this does not help to constrain the identification of the unknown structure because cementum and ossified periodontal ligaments would probably fluoresce with similar colours since these materials are both made of fossilised hydroxylapatite. In the LSF image enamel reacts differently to the laser light even though it is also made of fossilised hydroxylapatite. Clearly, the mineralogy of the fossilised enamel is sufficiently different to the other parts of the fossil to give such a vividly different orange fluorescence colour.

Unfortunately, laser-stimulated fluorescence (LSF) analysis (using a 408nm violet laser; 304 Kaye et al., 2015) was unable to support either the cementum 'jacket' or ossified ligament 305 identifications. The LSF image (Fig. 9) shows similar greeny fluorescence colours for the 306 unknown structure, dentine and maxillary bone, but a vivid orange colour for the enamel. 307 This is interesting because all of these materials are varieties of fossilised hydroxylapatite 308 309 (the ligaments are expected to have been ossified prior to fossilisation), but clearly there is a marked mineralogical difference between the fossilised enamel and the other fossilised 310 materials that is probably related to mineral density. Thus, the unknown structures are 311 proposed as cementum 'jackets' given the current evidence available. 312

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- To our knowledge these cementum 'jackets' have not been described amongst
- iguanodontians and other dinosaurs, but this derived root attachment tissue is commonly
- found in mammals where it also migrates onto the crowns (Erickson *et al.*, 2012). Studies of
- existing iguanodontian specimens in person and from the literature revealed their presence in
- 318 *Equijubus normani* (IVPP 12534, You *et al.*, 2003c: Fig. 1E). Unlike IVPP V22529, these
- 319 'jackets' are found on alternating teeth rather than on each one.



Figure 10. A cementum 'jacket' tooth morphology is also present in the basal hadrosauriform *Equijubus normani* (IVPP 12534, You *et al.*, 2003c: Fig. 1E), but unlike IVPP V22529 this is observed in alternating teeth rather than on each tooth.

### Tooth orientation

The maxillary tooth row follows a laterally concave path (Fig. 4). The slightly worn and unworn teeth appear to be posteriorly inclined whilst the heavily worn teeth are anteriorly inclined, but the latter appears to be an artefact of the fragmentation of the bone sockets holding them in place (Fig. 3). Thus, the specimen's tooth orientation is considered to be consistent with that of *P. gobiensis* (Norman, 2002) – posteriorly inclined.

# Maxillary body



Figure 11. Maxilla in medial view showing the broken anterior ramus and missing posterior ramus as well as the maxillary grooves, medial shelf, dorsal process, lingual view of the tooth row and the broken posteromedial surface that exposes several fragmentary tooth roots.

The medial shelf (Fig. 11) has been artificially shifted ventrally partially obscuring the row of 339 'special' foramina such that their exact number and shapes are unclear; they are located 340 341 relatively low on the medial surface, as in other basal hadrosauriforms such as Altirhinus (PIN 3386/7; right maxilla [Norman, 2002]). In Altirhinus (PIN 3386/7; Norman, 1998) an 342 incomplete row of 14 regularly spaced foramina is preserved subparallel to the ventral margin 343 344 of the maxilla (the missing posterior portion of the maxilla truncates this row of foramina). In Bactrosaurus their appearance varies between individuals of similar and different ages 345 (adult/subadult: AMNH 6553, Fig. 7 Prieto-Márquez, 2011; juvenile: AMNH 6389, 6390, 346 347 Figs. 9, 11 Prieto-Márquez, 2011). In *Gilmoreosaurus* there are at least 17 large, evenlyspaced and circular 'special' foramina (AMNH FARB 30653; Prieto-Márquez & Norell, 348 2010). 349



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Figure 12. The dorsal process of IVPP V22529 is laterally-compressed, subtriangular and dorsally-rounded similar to the ones present in *Altirhinus* (PIN 3386/7; Norman, 2002: Fig. 6) and *Bactrosaurus* (AMNH 6389, 6390; Prieto-Márquez, 2011: Figs. 9, 11). Dorsal process in, A, lateral view; B, in dorsal view showing the lacrimal articular groove that is also present in *Koshisaurus* and *Fukuisaurus* [Shibata & Azuma, 2015: Fig. 8C].

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358 A relatively straight portion of the dorsal margin is preserved occupying around three-

quarters of the specimen's preserved anteroposterior length (Figs. 2, 11). This makes an  $\sim 30^{\circ}$ 

angle with the ventral margin of the maxilla and is grooved (as observed in *Koshisaurus* and

*Fukuisaurus* [Shibata & Azuma, 2015: Fig. 8C]), presumably for articulation with the

lacrimal (Fig. 12). The highest point of this section of preserved dorsal margin appears to

- preserve a laterally-compressed, subtriangular and dorsally-rounded dorsal (ascending)
- process similar to the ones present in *Altirhinus* (PIN 3386/7; Norman, 2002: Fig. 6) and

365 Bactrosaurus (AMNH 6389, 6390; Prieto-Márquez, 2011: Figs. 9, 11). Norman (2015) characterises this process shape as a 'laterally flattened subtriangular plate' (Character 17, 366 state 2) (Figs. 2, 11, 12). This process shape does not have a well-defined distribution 367 amongst iguanodontians unlike the 'narrow' and 'finger-like' dorsal process morphologies 368 that Norman (2015) recovered as synapomorphies of Iguanodontia and Ankylopollexia 369 respectively (state 0 of character 17 respectively under ACCTRAN [='narrow, figure-like 370 process']). The dorsal process of IVPP V22529 possesses a shallow subcircular depression on its lateral surface beneath its tip (Figs. 2, 11, 12) which is not observed in *Bactrosaurus* and Altirhinus. A comparison with the condition in Probactrosaurus is not possible as the dorsal region is not preserved e.g. in PIN 2232/9-2 and /10-2 (Norman, 2002) [coded as a '?' in Norman, 2015]. IVPP V22529 is not sufficiently well-preserved to confidently characterise the presence or absence of an antorbital fenestra, a feature that is used to diagnose clypeodont ornithischians and their subclades (Norman, 2015).

The anterior portion of the maxilla appears to be subtriangular (Figs. 2, 11, 13), as in most iguanodontians including Altirhinus (PIN 3386/7; Norman, 1998: Fig. 6B), Bactrosaurus (AMNH 6553; Prieto-Márquez, 2011) and Probactrosaurus (PIN 2232/9-2; Norman, 2002). The anterior maxillary ramus of IVPP V22529 is forked into a pointed anteromedial process at around the same level as the incompletely preserved anterolateral process (Figs. 2, 11, 13). Owing to the incomplete preservation of the latter process (Figs. 2, 11, 13), the relative length and size of these processes cannot be determined. Processes of different sizes are found in the 'iguanodontoid' styracosternan Iguandon (Weishamphel et al., 1993), Protohadros (Head, 1998: Fig. 3C, D), Bactrosaurus (AMNH 6553, 6389, 6390; Prieto-Márquez, 2011: Figs. 7-12) and Koshisaurus (Shibata & Azuma, 2015: Figs. 3A, C; 8A, B, E), but in the latter the anterior processes are actually of similar rather than different lengths (Shibata & Azuma, 2015: Fig. 3A). Bifurcated anterior processes are actually diagnostic of Iguanodontia (Norman, 2015: character 15, state 1; ACCTRAN), but in the left maxilla of Shuangmiaosaurus the anterolateral process appears to have become particularly enlarged with a dorsally placed nubbin at its base potentially being the remnants of the anteromedial process (LPM0165; You et al., 2003b: Fig. 1A). Teeth are present right up to base of the 394 anteromedial process, as in Protohadros (Head, 1998: Fig. 3C, D). The anterior half of the 395 anterodorsal margin of the anterolateral process of IVPP V22529 has a finger-shaped recess 396 (rostral foramen) (Fig. 13), as in non-hadrosaurid iguandontians. 397 398



Figure 13. Dorsal view of the anterodorsal process of IVPP V22529 showing the bifurcating anterior processes at roughly the same level. Only the pointed anteromedial process is complete. This is a finger-shaped recess (rostral foramen) in the anterior half of the anterodorsal margin of the anterolateral process.

404 Halfway up the medial side of the maxilla there is a well-developed medially projecting shelf 405 406 that originates from the dorsomedial portion of the anterior ramus (Fig. 11). Subhorizontal 407 ridges along the medial side of the anteromedial process and the medial shelf become increasingly well-developed dorsoposteriorly (although a portion of the intervening area is 408 409 broken). These ridges demark the boundaries of the maxillary grooves. Unlike IVPP V22529, 410 the maxillary grooves of *Koshisaurus* (Shibata & Azuma, 2015: Figs. 3C; 8B, E) are all poorly-developed where there are all well-developed in Protohadros (SMU 74582; Head, 411 1998: Fig. 3D) and Fukuisaurus (FPDM-V40-1; Shibata & Azuma, 2015; Fig. 8B, E; three 412 and five grooves respectively). Maxillary grooves presumably relate to the attachment of soft 413 tissues in the roof of the mouth, but these seems to be absent in the most derived 414 iguanodontians - the distribution of this feature is unclear across Iguanodontia. The evolution 415 of maxillary grooves is therefore of interest in further understanding iguanodontian feeding 416 and as a potential source of phylogenetic information. 417



Figure 14. The maxillary grooves in IVPP V22529 become better-developed along the medial surface of the anteromedial process and the medial shelf, unlike in *Protohadros* (SMU 74582; Head, 1998: Fig. 3D), *Fukuisaurus* (FPDM-V40-1; Shibata & Azuma, 2015: Fig. 8B, E) and *Koshisaurus* (Shibata & Azuma, 2015: Figs. 3C; 8B, E) where the grooves in each specimen are of similar sizes. However, the phylogenetic significance of maxillary groove morphologies is not understood.

The mediolateral width of the medial shelf is affected by dorsoventral diagenetic 427 428 compression, as evident from bone fragments that are thrusted upon each other (Fig. 11). However, the exact extent of this diagenetic artefact is unclear because a compression-429 corrected bone reconstruction is beyond the scope of this paper to produce. Bactrosaurus has 430 a comparatively less developed shelf in both adult (AMNH 6553; Prieto-Márquez, 2011: Fig. 431 7) and juvenile specimens (AMNH 6389, 6390; Prieto-Márquez, 2011: Figs. 9, 11) as well as 432 Altirhinus (PIN 3386/7; Norman, 1998: Fig. 6B) [but the degree of lesser development is 433 434 unknown for the aforementioned reason]. In these specimens and in IVPP V22529 the shelf is angled slightly dorsoposteriorly (Fig. 11). In contrast, Probactrosaurus was described by 435 Norman (2002) as having a vertical and planar medial wall. 436

437

The entire posterior portion of IVPP V22529 is missing so the morphology of the jugal-438 maxilla suture is unknown (Figs. 2, 11). In Altirhinus (PIN 3386/7; Norman, 1998; Fig. 6A) 439 the jugal sutural surface is a finger-like process that fits into a slot in the anterior ramus of the 440 jugal, a feature that unites styracosternan iguanodontians (Norman, 2015: character 20, state 441 1). The missing anterodorsal margin makes it impossible to infer the morphology of the 442 jugal's ventral margin which is sinusoidal in ankylopollexian iguanodontians (Norman, 2015: 443 character 21, state 1). The missing posterior ramus means it is unclear if this is dorsoventrally 444 tall with a rounded but slightly irregular tip, as in *Probactrosaurus* (PIN 2232/9-2,10-2; 445 Norman, 2002: Fig. 5), or if it is more 'finger-like', as in Altirhinus (PIN 3386/7; Norman, 446 447 2002, Fig. 6A, B).



Figure 15. Close-up of the relatively flat lateral surface of IVPP V25529 showing a row of anteriorly-opening foramina that increase in size posteriorly.

The lateral surface of the maxilla is relatively flat, a trait that is undoubtedly influenced by the fragmentation of the specimen (Figs. 2, 15). Ventral to the dorsal process is a broad shallow groove (Figs. 2, 12). There is a low scattered row of five anteriorly-opening neurovascular foramina on the lateral surface that increase in size posteriorly (Fig. 15). These 456 observations appear to be generally consistent with Probactrosaurus, notably the right 457 maxilla of PIN 2232-10-2 (Norman, 2002, Fig. 5A), except that the foramina in the latter do 458 not increase in size posteriorly. However, these arguments are weakened by the absence of 459 foramina in the larger left maxilla *Probactrosaurus* PIN 2232-9-2 (Norman, 2002, Fig. 5B) 460 suggesting that these foramina are not consistently expressed on the sides of the skull and/or 461 they may undergo changes with age or be different between sexes. 462 463

### 465 **Discussion**

466

464

467 Taxonomic status of IVPP V22529

- 468 469 *A basal hadrosauriform basal hadrosaurif*
- 469 *A basal hadrosauriform based on tooth anatomy*470 IVPP V22529 matches the maxillary teeth characteristics in the diagnosis of *Probactrosaurus*
- 470 (Norman, 2002): 'maxillary teeth narrow with prominent primary ridge and no subsidiary
- 472 ridges; tall and interlocking teeth that form a high, posteriorly inclined battery; marginal
- 473 denticles are mammillate.' However, these characteristics are not used to refer IVPP V22529
- 474 to *Probactrosaurus* (Norman, 2002) as the aforementioned characteristics are now
- 475 understood to have a wider distribution amongst iguanodontians than previously appreciated
- 476 (Norman, 2015). The tooth anatomy of IVPP V22529 identifies it as a non-euhadrosaurian

- hadrosauriform: two or more replacement crowns are a hadrosauriform iguandontian feature
  (Norman, 2015: character 54, state 1) whilst the absence of a single median primary ridge is a
  non-euhadrosaurian iguanodontian feature (Norman, 2015: character 68, state 3). The
  marginal denticles of IVPP V22529 comprise of parallel ledges with single rows of ~6
  relatively large mammillae, a feature that further constrains the specimen as a basal
  hadrosauriform.
- 482 ha 483

## 484 *IVPP V22529 compared to* Probactrosaurus

IVPP V22529 has a number of noteworthy differences with Probactrosaurus maxillae, 485 486 despite the former missing its posterior portion and the latter missing anterior ramii. Firstly, Norman (2002) noted that Probactrosaurus gobiensis has a vertical and planar medial wall 487 whereas in IVPP V22529 this in non-planar owing to its well-developed medial shelf (Fig. 488 489 11). However, a small proportion of the latter is ascribed to specimen deformation. Secondly, there is a low scattered row of five anteriorly-opening foramina on the lateral surface of IVPP 490 V22529 that increase in size posteriorly, but in Probactrosaurus PIN 2232-10-2 (Norman, 491 2002, Fig. 5A) these foramina do not increase in size posteriorly and are even absent in 492 493 specimen PIN 2232-9-2 (Norman, 2002, Fig. 5B). In Probactrosaurus (and iguanodontians more generally) there is incomplete knowledge of how maxillary foramina change with age 494 and how they can be different between sexes or individuals. Thus, further work is needed to 495 496 evaluate these types of variability so that the phylogenetic utility of maxillary foramina can 497 be established. Therefore, erring on the side of caution, the foraminal differences between IVPP V22529 and Probactrosaurus should be considered tentatively as differences with 498 499 taxonomic value, and certainly warrant lesser value than the aforementioned medial shelf difference. The presence and absence of the unique cementum 'jackets' in IVPP V22529 and 500 Probactrosaurus respectively could be strong evidence for differentiating them. However, 501 502 given the areas of uncertainty in the identification and formation mechanism of this structure as well as its presence in at least one iguanodontian, it would be inappropriate to place 503 phylogenetic value on this structure until it is more extensively investigated. The teeth of 504 505 IVPP V22529 and Probactrosaurus are very similar and their maxillae are both subtriangular. 506

# 508 IVPP V22529 compared to Bactrosaurus and Gilmoreosaurus

The expression of neurovasucular foramina on the lateral surface of the maxilla is variably 509 expressed in *Bactrosaurus*, which has a row of different-sized ones in juvenile specimen 510 AMNH 6389 (Prieto-Márquez, 2011: Fig. 9) but more random arranged one in juvenile 511 512 specimen AMNH 6390 (Prieto-Márquez, 2011: Fig. 11). However, as in IVPP V22529 these foramina appear low on the lateral surface. In contrast, the maxillary foramina of 513 Gilmoreosaurus AMNH FARB 30653 (Prieto-Márquez & Norell, 2010) are more randomly 514 515 distributed but appear high as well as low on the lateral surface. On the medial surface of the latter specimen the 'special foramina' are larger and more circular than those in IVPP 516 V22529 (although these are partially obscured by the displaced medial shelf) and 517 Bactrosaurus (AMNH 6553, 6389, 6390; Prieto-Márquez, 2011; Figs. 8, 10, 12). 518 Gilmoreosaurus AMNH FARB 30653 (Prieto-Márquez & Norell, 2010) and Bactrosaurus 519 (AMNH 6553, 6389, 6390; Prieto-Márquez, 2011: Figs. 8, 10, 12) lack maxillary grooves 520 521 that become increasingly developed along the medial surface of the anteromedial process and the medial shelf, as in IVPP V22529 (Fig. 11). However, Gilmoreosaurus AMNH FARB 522 30653 (Prieto-Márquez & Norell, 2010) has less-developed maxillary grooves restricted to 523 524 the medial surface of the anteromedial process and so are less extensive than those of Koshisaurus (Shibata & Azuma, 2015: Figs. 3C; 8B, E). Bactrosaurus AMNH 6389 (Prieto-525

- 526 Márquez, 2011: Fig. 10) also appears to share this characteristic with *Gilmoreosaurus*, but 527 this needs to be confirmed by first-hand study of this specimen.
- 528
- 529 Cementum 'jackets'

Cementum 'jackets' are a dental structure presumably related to feeding style, but how it is 530 related to this is not yet obvious. The restriction of these 'jackets' to the labial side of the 531 532 tooth appears to be genuine and suggests this side was well-anchored to the tooth socket. The latter trait may have been important in allowing the 'jackets' to help resist lateral components 533 of bite forces, a force regime that is evident from the medially-directed slope of the wear 534 535 facets of the teeth. The 'jackets' probably reduced stress on the brittle tooth crests more generally as well by transmitting loads amongst the tissues of the teeth (Erickson et al., 536 2012). However, if these structures did have these roles then the alternating occurrence of the 537 'jackets' in *Equijubus* implies that not all taxa benefitted from this hypothesised function 538 equally. Histological analysis will no doubt be invaluable in testing the cementum 'jacket' 539 hypothesis at the microscopic level and should help to clarify both the composition and 540 morphology of these structures. Unfortunately, such work is beyond the scope of the current 541 study, but it should be a priority for future studies of IVPP V22529. Future opportunities to 542 compare feeding biomechanics in IVPP V22529 (and other basal hadrosauriformes) with 543 more derived hadrosauriforms (Erickson et al., 2012) will also be worthwhile to determine 544 545 how important these cementum 'jackets' were in iguandontian dental system evolution.

#### Conclusions

IVPP V22529 is an isolated Early Cretaceous partial right iguanodontian maxilla that 549 possesses tooth characteristics identifying it as a basal hadrosauriform. However, other parts 550 551 of this bone fail to convincingly support a referral to a new or existing taxon, including to a new or existing species of *Probactrosaurus*, a contemporaneous genus known from the same 552 locality in North China. Further work may better constrain the taxonomic status of this 553 specimen if characteristics differing from Probactrosaurus can be validated, namely: a 554 unique corrugated middle ventrolateral margin, a row of foramina on its lateral surface that 555 open anteriorly and increasing in size posteriorly as well as a prominent medial shelf. Despite 556 its coarse level of identification, IVPP V22529 has important implications for our 557 understanding of iguanodontian (and dinosaurian) dental architecture. In labial view, five 558 middle maxillary crowns each preserve a rugose cementum sheath over their basal portions 559 that are separate from the tooth socket but actually originate within them and extend ventrally 560 561 below the crenulated ventral margin of the maxilla. This arrangement forms a corrugated ventrolateral surface as grooves separate the sides of these sheaths. This structure - which we 562 propose to call a cementum 'jacket' structure - appears to be present in the basal 563 564 hadrosauriform *Equijubus* as well, but this differs from IVPP V22529 in being present in every other tooth rather than on each tooth. To our knowledge this structure has not been 565 described in other dinosaurs, but cementum commonly migrates onto the tooth crowns of 566 mammals (Erickson et al., 2012). The wider distribution of cementum 'jackets' amongst 567 iguanodontians (and dinosaurs more generally) warrants further attention as their morphology 568 could carry important phylogenetic information. The restriction of these 'jackets' to the labial 569 face of the teeth might indicate a structural role in resisting the lateral component of bite 570 forces and/or the stress on the brittle tooth crests, but these hypotheses and confirmation of 571 'jacket' composition and morphology would greatly benefit from future histological analysis 572 573 and biomechanical studies that were beyond the scope of this study. Despite, the further work required, IVPP V22529 provides important new insights into the dental architecture of basal 574

- 575 hadrosauriforms that deepens our understanding of the morphological diversity that preceded
- the revolutionary advanced hadrosauriform dental battery system (Erickson *et al.*, 2012).
- 577 578

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