### The comparative osteology of *Plesiochelys bigleri* n. sp., a new coastal marine turtle from the Late Jurassic of Porrentruy (Switzerland) (#16722)

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1

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### The comparative osteology of *Plesiochelys bigleri* n. sp., a new coastal marine turtle from the Late Jurassic of Porrentruy (Switzerland)

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**Background.** During the Late Jurassic several groups of eucryptodiran turtles inhabited the shallow epicontinental seas of Western Europe. Plesiochelyidae are an important part of this first radiation of crown group turtles into marine ecosystems. Fossils of Plesiochelyidae occur in many European localities, and are especially abundant in the Kimmeridgian layers of the Swiss Jura Mountains (Solothurn and Porrentruy). In the mid 19th century, the quarries of Solothurn (NW Switzerland) already provided a large amount of fossil turtles, most notably *Plesiochelys etalloni*, the best-known plesiochelyid species. Recent excavations in the Porrentruy area (NW Switzerland) revealed new fossils of *Plesiochelys*, including numerous well-preserved shells with associated cranial and postcranial material.

**Methods/Results.** Out of 80 shells referred to *Plesiochelys*, 41 are assigned to a new species, *Pl. bigleri* n. sp., including a skull-shell association. We furthermore refer 15 shells to *Pl. etalloni*, and 24 shells to *Plesiochelys* sp. Anatomical comparisons show that *Pl. bigleri* can clearly be differentiated from *Pl. etalloni* by cranial features. The shell anatomy and the appendicular skeleton of *Pl. bigleri* and *Pl. etalloni* are very similar. However, a statistical analysis demonstrates that the thickness of neural bones allows to separate the two species based on incomplete material. This study furthermore illustrates the extent of intraspecific variation in the shell anatomy of *Pl. bigleri* and *Pl. etalloni*. Our results represent an important point of comparison for future studies on Mesozoic turtle diversity.

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#### 15 ABSTRACT

Background. During the Late Jurassic several groups of eucryptodiran turtles inhabited the 16 shallow epicontinental seas of Western Europe. Plesiochelyidae are an important part of this first 17 radiation of crown group turtles into marine ecosystems. Fossils of Plesiochelyidae occur in 18 many European localities, and are especially abundant in the Kimmeridgian layers of the Swiss 19 20 Jura Mountains (Solothurn and Porrentruy). In the mid 19th century, the quarries of Solothurn (NW Switzerland) already provided a large amount of fossil turtles, most notably Plesiochelys 21 etalloni, the best-known plesiochelyid species. Recent excavations in the Porrentruy area (NW 22 Switzerland) revealed new fossils of Plesiochelys, including numerous well-preserved shells 23 with associated cranial and postcranial material. 24 Methods/Results. Out of 80 shells referred to *Plesiochelys*, 41 are assigned to a new species, *Pl*. 25 *bigleri* n. sp., including a skull-shell association. We furthermore refer 15 shells to *Pl. etalloni*, 26 and 24 shells to *Plesiochelys* sp. Anatomical comparisons show that *Pl. bigleri* can clearly be 27 28 differentiated from *Pl. etalloni* by cranial features. The shell anatomy and the appendicular skeleton of *Pl. bigleri* and *Pl. etalloni* are very similar. However, a statistical analysis 29 demonstrates that the thickness of neural bones allows to separate the two species based on 30 31 incomplete material. This study furthermore illustrates the extent of intraspecific variation in the shell anatomy of *Pl. bigleri* and *Pl. etalloni*. Our results represent an important point of 32

33 comparison for future studies on Mesozoic turtle diversity.

### 34 INTRODUCTION

The first radiation of crown-group turtles into marine environments occurred in Western Europe during the Late Jurassic. At that time, several groups of basal pan-cryptodiran turtles, traditionally referred to the families Plesiochelyidae Baur, 1888, Thalassemydidae Zittel, 1889, and Eurysternidae Dollo, 1886, colonized coastal ecosystems from restricted lagoons to more open seaways. These turtles eventually disappeared at the Jurassic-Cretaceous boundary following major sea-level changes that restricted their habitats (e.g., Bardet, 1994; Bardet, 1995; Bardet et al., 2014).

Plesiochelyids are relatively large coastal marine turtles known from the Late Jurassic of 42 Switzerland, France, Germany, England, Spain, and Portugal. They are notably characterized by 43 a fully ossified carapace and a series of derived basicranial features (Gaffney, 1975a; Gaffney, 44 1976; Anquetin, Püntener & Billon-Bruyat, 2015). Plesiochelys etalloni (Pictet & Humbert, 45 1857) is undoubtedly the best known plesiochelyid turtle, thanks notably to numerous specimens 46 47 found in Solothurn, Switzerland (Rütimeyer, 1873; Bräm, 1965; Gaffney, 1975a; Gaffney, 1976; Anquetin, Püntener & Billon-Bruyat, 2014; Anquetin, Püntener & Billon-Bruyat, 2015). 48 *Plesiochelys etalloni* is known by both skulls and shells from the Kimmeridgian of the Swiss and 49 50 French Jura Mountains, southern England, and northwestern Germany (Pictet & Humbert, 1857; Rütimeyer, 1873; Bräm, 1965; Karl et al., 2007; Anquetin, Deschamps & Claude, 2014, 51 52 Anquetin, Püntener & Billon-Bruyat, 2014; Anquetin, Püntener & Billon-Bruyat, 2015; Anquetin 53 & Chapman, 2016). Plesiochelys planiceps (Owen, 1842), the only other valid species in the genus, is known only from a single specimen (cranium, mandibule, and remains of the hyoids 54 55 and cervical vertebrae) from the Tithonian of the Isle of Portland, UK. The cranium of Pl.

*planiceps* differs in many aspects from that of *Pl. etalloni* (Anquetin, Püntener & Billon-Bruyat,
2015).

In the present study we describe new material of Plesiochelys from the Kimmeridgian of 58 Porrentruy, Canton Jura, Switzerland. The new specimens were found by the Paleontology A16 59 project, which rescued the paleontological material discovered during the construction of the 60 61 A16 Transjurane highway. These excavations yielded a great number of fossil vertebrates from Kimmeridgian layers, notably including dinosaur trackways (Marty & Hug, 2003; Marty et al., 62 2007; Marty, 2008; Marty & Billon-Bruyat, 2009) and numerous coastal marine turtles (Billon-63 Bruyat, 2005a). The rich and diverse turtle fauna from Porrentruy notably includes different 64 species of Plesiochelyidae and Thalassemydidae (see Geological setting) (Püntener et al., 2014; 65 Anguetin, Püntener & Billon-Bruyat, 2015; Püntener, Anguetin & Billon-Bruyat, 2015). 66 As in Solothurn, *Plesiochelys* is by far the most common turtle taxon in the Kimmeridgian 67 of Porrentruy. Slightly more than 100 relatively complete, but mostly disarticulated shells were 68 69 discovered during the excavations, out of which 80 can be referred to *Plesiochelys*. Among these shells, 41 are herein assigned to a new species, Plesiochelys bigleri n. sp. We furthermore refer 70 15 shells to *Plesiochelys etalloni*, and 24 shells to *Plesiochelys* sp. This material is described in 71 72 detail herein. The shell of *Pl. bigleri* shows only minor anatomical differences with that of *Pl. etalloni*, but cranial anatomy clearly distinguishes the two species. Two skulls, one associated 73 74 with a shell and the other found isolated, are known for *Pl. bigleri*. A statistical analysis confirms 75 that the thickness of neural bones allows to separate the two species and to tentatively identify otherwise indeterminate specimens. Based on abundant shell material, the intraspecific variations 76 77 in both species are discussed in details. Finally, we fully describe and illustrate elements of the

appendicular skeleton, which are otherwise rarely described in the literature, hoping that this willfacilitate future comparisons.

80

### 81 MATERIAL AND METHODS

82

#### 83 Material

The present study is based on a collection of 80 relatively complete, but mostly disarticulated shells (Table 1), most of which were found in a single stratigraphical layer (see Geological setting, below). Forty-one shells are referred to a new species, *Plesiochelys bigleri* n. sp. One of these specimens (MJSN TCH007-252) is a skull-shell association, which we designate as the holotype of the new species. An isolated cranium (MJSN TCH006-1451) is also referred to this new species and designated as its paratype.

90 Fifteen out of the 80 aforementioned shells are identified as *Plesiochelys etalloni*. The 91 identification of the remaining 24 shells is uncertain because they lack diagnostic features. These 92 specimens are therefore referred to *Plesiochelys* sp. However, a tentative identification of some 93 of these specimens is provided herein based on the statistical analysis of neural bone thickness 94 (see below).

95

#### 96 Geological setting

All of the specimens were collected between 2001 and 2011 near the small town of Courtedoux,
along the A16 Transjurane highway in the Ajoie Region, Canton of Jura, NW Switzerland (Fig.
1). The majority of the specimens come from the Lower *Virgula* Marls (Reuchenette Formation,
Chevenez Member; Comment et al., 2015) of the sites of Bois de Sylleux (BSY), Sur Combe

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Ronde (SCR), and Tchâfouè (TCH) (Fig. 2). These sites yielded a rich and diverse coastal
marine assemblage, including invertebrates (bivalves, gastropods, cephalopods, crustaceans, and
echinoderms), vertebrates (chondrichthyans, osteichthyans, turtles, crocodilians, and pterosaurs),
and wood remains (e.g., Billon-Bruyat, 2005a; Billon-Bruyat, 2005b; Marty & Billon-Bruyat,
2009; Philippe et al., 2010; Schaefer, 2012; Comment et al., 2015; Koppka, 2015; Leuzinger et
al., 2015).

*Plesiochelys* is the dominating turtle taxon in the Lower *Virgula* Marls. Other taxa occur 107 only in small numbers, including Tropidemys langii Rütimeyer, 1873, Portlandemys gracilis 108 Anquetin, Püntener & Billon-Bruyat, 2015, Thalassemys hugii Rütimeyer, 1873, and 109 Thalassemys bruntrutana Püntener, Anguetin & Billon-Bruyat, 2015 (Püntener et al., 2014; 110 Anguetin, Püntener & Billon-Bruyat, 2015; Püntener, Anguetin & Billon Bruyat, 2015). The 111 Lower *Virgula* Marls are dated from the Eudoxus ammonite zone (early late Kimmeridgian; 112 Comment et al., 2015) and are therefore slightly older than the Solothurn Turtle Limestone, 113 114 which forms the uppermost member of the Reuchenette Formation and is dated from the Autissiodorensis ammonite zone (Meyer, 1994; Comment, Ayer & Becker, 2011). 115 One specimen of *Plesiochelys bigleri* (MJSN CRT007-2) has been discovered within the 116 117 dinosaur track-bearing tidal laminites of the Crat site (CRT; Fig. 2) (Billon-Bruyat et al., 2012). These laminites represent the lowermost layers of the Corbis Limestones and are dated from the 118 119 Cymodoce ammonite zone (late early Kimmeridgian; Comment et al., 2015). Two other 120 specimens of Plesiochelys bigleri (MJSN VTT006-299 and MJSN VTT006-579) are stratigraphically slightly older and come from the Banné Marls of the Vâ Tche Tchâ site (VTT; 121 122 Cymodoce ammonite zone; Fig. 2), where *Tropidemys langii* is the dominating turtle taxon 123 (Püntener et al., 2014).

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Most turtle shells from Porrentruy have been discovered in a state of partial disarticulation, 124 which contrasts with the mostly articulated shells found in Solothurn. The latter were apparently 125 rapidly incorporated in a calcareous mud, while the turtle remains from Porrentruy remained for 126 a longer period on top of the sediment before being completely buried. This is confirmed by the 127 common presence of incrusting bivalves (oysters) on the bone remains from Porrentruy. 128 129 However, disturbance by predators or water movements was relatively limited since disarticulated elements were usually found relatively close together (Fig. 3). 130 131 **Anatomical comparisons** 132

133 As far as the cranium is concerned, *Plesiochelys bigleri* was compared to all plesiochelyids for

134 which that part of the skeleton is known: *Plesiochelys etalloni* (NMB 435, NMS 8738, NMS

135 8739, NMS 8740, NMS 9145, NMS 40870, NMS 40871, NHMUK R3370), Plesiochelys

136 planiceps (OUMNH J.1582), Portlandemys mcdowelli Gaffney, 1975a (NHMUK R2914,

137 NHMUK R3164), and Portlandemys gracilis (MJSN BSY009-708). When pertinent,

138 comparisons were also extended to PIMUZ A/III 514, a skull-shell association from the

139 Tithonian of the Isle of Oléron (Department of Charente-Maritime, France) initially referred to

140 *Thalassemys moseri* Bräm, 1965 by Rieppel (1980), but which was recently designated as the

141 holotype of a new taxon, Jurassichelon oleronensis Pérez-García, 2015. All of these specimens

142 have been studied first hand by the second author. The reader is referred to the primary literature

describing these specimens (Parsons & Williams, 1961; Gaffney, 1975a; Gaffney, 1976; Rieppel,

144 1980; Anquetin, Püntener & Billon-Bruyat, 2015; Anquetin & Chapman, 2016). Anatomical

descriptions in the present study follow the nomenclature established by Gaffney (1972, 1979) as

146 updated by Rabi et al. (2013).

Anatomical descriptions of shell material follow the nomenclature established by Zangerl (1969). Shell and non-shell postcranial material of *Pl. bigleri* was compared, when pertinent, to plesiochelyids (*Plesichelys etalloni, Tropidemys langii*, and *Craspedochelys jaccardi*) and thalassemydids (*Thalassemys bruntrutana* and *Thalassemys hugii*).

151

#### 152 Statistical analysis

153 The length and thickness of 119 neurals 2 to 5 pertaining to 43 selected specimens (25

154 *Plesiochelys bigleri*, 8 *Plesiochelys etalloni*, and 10 *Plesiochelys* sp.) were measured in order to

155 test the hypothesis that neural bones are significantly thinner in *Pl. bigleri* relative to *Pl. etalloni* 

156 (see Table S1). Length was measured as the maximal length on the dorsal surface of the neural

157 bone. Thickness was measured on the left and right sides and approximately at the mid-length of

each neural bone (see below). The mean of these two measurements was used as the thickness

value for each individual neural bone (Table S1). Measurements were taken using a digital

160 Vernier caliper by a single operator (CP).

Neural shape can be relatively variable within a single individual. For example, one neural can be disproportionately shorter, longer, thinner, or thicker in a given neural series. Specimens exhibiting extreme divergences from the common condition were not measured. For specimens included in this analysis, the mean neural length and mean neural thickness were computed for each individual, which had the effect of smoothing intra-individual discrepancies. The analyzed dataset therefore consists of the mean neural length, mean neural thickness, and corresponding length/thickness ratio measured for the 43 included specimens (see below).

168 The statistical analysis was run using PAST 3.14 (Hammer, Harper & Ryan, 2001). Length 169 and thickness were plotted in a 2D space, whereas length, thickness, and length/thickness ratio

170 were tested for equal medians using a non-parametric Mann-Whitney test (with Monte Carlo

171 permutations). A discriminant analysis was also performed on the length and thickness

172 measurements and used to tentatively identify indeterminate specimens (*Plesiochelys* sp.; see

173 below).

174

### 175 **3D Models**

176 3D models of the holotype (MJSN TCH007-252) and paratype (MJSN TCH006-1451) crania

177 have been computed with the photogrammetry software Agisoft Photoscan 1.0.4 Standard

178 Edition using sets of high-quality photographs of the specimens. We followed the procedures

recently described by Mallison & Wings (2014). These models are provided herein as 3D PDFs

180 (reduced resolution; to be opened with Adobe Acrobat): MJSN TCH007-252 (Fig. S1), and

181 MJSN TCH006-1451 (Fig. S2). Scaled and textured high-resolution meshes in PLY format are

- also available freely on figshare (<u>http://figshare.com/authors/J\_r\_my\_Anquetin/651097</u>).
- 183 A 3D surface scan of the pelvis preserved with specimen MJSN BSY006-307 was
- 184 produced with an Artec Space Spider scanner (Artec Group, Luxembourg;
- 185 http://www.artec3d.com) and reconstructed with Artec Studio 10, the native scanner software.

186 The textured 3D mesh in PLY format is freely available on figshare

- 187 (http://figshare.com/authors/J\_r\_my\_Anquetin/651097).
- 188

### 189 Nomenclatural act

- 190 The electronic version of this article in Portable Document Format (PDF) will represent a
- 191 published work according to the International Commission on Zoological Nomenclature (ICZN),
- and hence the new name contained in the electronic version is effectively published under that

- 193 Code from the electronic edition alone. This published work and the nomenclatural act it
- 194 contains have been registered in ZooBank, the online registration system for the ICZN. The
- 195 ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed
- 196 through any standard web browser by appending the LSID to the prefix <u>http://zoobank.org/</u>. The
- 197 LSID for this publication is: urn:lsid:zoobank.org:pub:C5AE9DE8-9911-4CFD-AD09-
- 198 65850C35BDEC. The online version of this work is archived and available from the following
- 199 digital repositories: PeerJ, PubMed Central and CLOCKSS.
- 200

#### 201 SYSTEMATIC PALEONTOLOGY

- 202 TESTUDINES Batsch, 1788
- 203 PAN-CRYPTODIRA Joyce, Parham & Gauthier, 2004
- 204 EUCRYPTODIRA Gaffney, 1975c
- 205 PLESIOCHELYIDAE Baur, 1888

### 206 *Plesiochelys* Rütimeyer, 1873

- 207 Type species. Plesiochelys solodurensis Rütimeyer, 1873
- 208 Included valid species. Plesiochelys planiceps (Owen, 1842); Plesiochelys etalloni (Pictet &
- 209 Humbert, 1857); Plesiochelys bigleri n. sp.
- 210 Referred material and range. Kimmeridgian of Switzerland, France, Germany, and England
- 211 (Pictet & Humbert, 1857; Maack, 1869; Rütimeyer, 1873; Oertel, 1924; Bräm, 1965; Karl et al.,
- 212 2007; Anquetin, Deschamps & Claude, 2014; Anquetin, Püntener & Billon-Bruyat, 2014;
- 213 Anquetin, Püntener & Billon-Bruyat, 2015; Anquetin & Chapman, 2016), and Tithonian of
- England (Owen, 1842; Gaffney 1975a; Gaffney, 1976; Anquetin, Püntener & Billon-Bruyat,
- 215 2015). Indeterminate specimens are also signalled from the Kimmeridgian-Tithonian of Portugal

216 (Pérez-García et al., 2008) and late Tithonian of Spain (Pérez-García, Scheyer & Murelaga,

217 2013).

- 218 Diagnosis. See Anquetin, Püntener & Billon-Bruyat (2014, 2015).
- 219

#### 220 Plesiochelys etalloni (Pictet & Humbert, 1857)

- 221 Synonymy. Emys Etalloni Pictet & Humbert, 1857 (original description); Stylemys hannoverana
- 222 Maack, 1869 (subjective synonymy); Plesiochelys langii Rütimeyer, 1873 (subjective

223 synonymy); Plesiochelys sanctaeverenae Rütimeyer, 1873 (subjective synonymy); Plesiochelys

224 solodurensis Rütimeyer, 1873 (subjective synonymy); Plesiochelys solodurensis var.

- 225 langenbergensis Oertel, 1924 (subjective synonymy).
- 226 *Type material*. MAJ 2005-11-1, a shell missing a large part of the carapace medially.
- 227 Illustrations of type. Pictet & Humbert (1857: plates I-III); Anquetin, Deschamps & Claude
- 228 (2014: Figs. 1 and 2, S2); Anquetin, Püntener & Billon-Bruyat (2014: Figs. 2A–2D).
- 229 Type horizon and locality. "Forêt de Lect" (Lect is a small village) near Moirans-en-Montagne
- 230 (Department of Jura, France), Late Jurassic. See Anquetin, Deschamps & Claude (2014) for
- 231 details.
- 232 Referred material and range. Kimmeridgian of Oker and Hannover, Lower Saxony, Germany
- 233 (Maack, 1869; Oertel, 1924; Karl et al., 2007); Kimmeridgian of Solothurn, Canton of Solothurn,
- and Glovelier and Porrentruy, Canton of Jura, Switzerland (Rütimeyer, 1873; Bräm, 1965;
- 235 Gaffney, 1975a; Gaffney, 1976; Anquetin, Deschamps & Claude, 2014; Anquetin, Püntener &
- Billon-Bruyat, 2014; Anquetin, Püntener & Billon-Bruyat, 2015; Table 1); Kimmeridgian of
- 237 England, UK (Anquetin & Chapman, 2016).



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*Emended diagnosis. Plesiochelys etalloni* differs from other *Plesiochelys* spp. in a more 238 extensive flooring of the cavum acustico-jugulare by the pterygoid, the complete ossification of 239 the pila prootica, and a narrow, slit-like foramen nervi trigemini. In addition, Plesiochelys 240 etalloni differs from Plesiochelys planiceps in a smaller size, a lower lingual ridge on the 241 maxilla, a narrower distance between the lingual ridges of the maxilla at the level of the 242 243 pterygoid-vomer suture, a more rounded foramen palatinum posterius, a parietal-quadrate contact posterior to the foramen nervi trigemini, a less developed processus trochlearis oticum, a 244 superficial canalis caroticus internus often remaining partly open ventrally, an absent or reduced 245 contribution of the exoccipital to the condylus occipitalis, and the anterior portion of the lingual 246 ridge on the dentary curving medially, and from *Plesiochelvs bigleri* in a higher temporal skull 247 roof, a deeper pterygoid fossa, a more developed processus trochlearis oticum, an anterior 248 foramen nervi abducentis opening more posteriorly relative to the base of the processus 249 clinoideus, foramina anterius canalis carotici cerebralis opening almost vertically below the 250 dorsum sellae and usually more closely set, a processus paroccipitalis extending mainly 251 posteriorly, an increased neural and costal bone thickness, the presence of epiplastral bulbs, and 252 a more rounded or pointed anterior margin of the anterior plastral lobe. 253

254

255 *Plesiochelys bigleri* sp. nov.

#### 256 urn:lsid:zoobank.org:act:9A8EF46E-7DAA-4F5B-B727-58C559BA503C

257 Figs. 3–8, 9A–B, 10–15

*Etymology*. This species is dedicated to Pierre Bigler (Villars-sur-Fontenais, Canton of Jura,
Switzerland) who so skillfully prepared many of the fossil turtles from the Paleontology A16
collection, including the holotype specimen.

Holotype. MJSN TCH007-252, near complete disarticulated carapace, epiplastra, entoplastron, 261

- hypoplastra, and right xiphiplastron; basicranium; proximal part of right scapular process; 262
- 263 proximal parts of both humeri; one radius; one ulna; both ilia with acetabulum (Figs. 4, 7, 11,
- and 12). 264

272

- Type locality and horizon. Tchâfoué (TCH), Courtedoux, near Porrentruy, Canton of Jura, 265
- Switzerland. Lower Virgula Marls, Chevenez Member, Reuchenette Formation, late 266
- Kimmeridgian, Late Jurassic (Comment, Ayer & Becker, 2011; Comment et al., 2015). 267

Paratype. MJSN TCH006-1451, a plated partial cranium (Figs. 5 and 6). 268

*Referred material and range*. Late early and early late Kimmeridgian of Porrentruy, Canton of 269

Jura, Switzerland (See Table 1). 270

Diagnosis. Plesiochelys bigleri differs from other Plesiochelys spp. in a lower temporal skull 271

roof, a shallower pterygoid fossa, a reduced processus trochlearis oticum, a more rounded

foramen nervi trigemini, an anterior foramen nervi abducentis opening anteromedially to the 273

274 base of the processus clinoideus, and foramina anterius canalis carotici cerebralis opening more

anteriorly relative to the level of the dorsum sellae. In addition, Plesiochelys bigleri differs from 275

*Plesiochelys planiceps* in a smaller size, a lower lingual ridge on the maxilla, a parietal-quadrate 276

277 contact posterior to the foramen nervi trigemini, a less developed processus trochlearis oticum, a

superficial canalis caroticus internus that may have remained partly open ventrally, and an absent 278

279 or reduced contribution of the exoccipital to the condylus occipitalis, and from *Plesiochelys* 

280 *etalloni* in a less extensive flooring of the cavum acustico-jugulare by the pterygoid, the absence

of complete ossification of the pila prootica, a processus paroccipitalis extending 281

282 posterolaterally, a reduced neural and costal bone thickness, absent or poorly developed

283 epiplastral bulbs, and a more quadrangular anterior margin of the anterior plastral lobe.

284	Remarks. The better part of the 42 specimens referred to Plesiochelys bigleri will be amply
285	illustrated in the forthcoming "Catalogues du patrimoine paléontologique jurassien", which
286	document the numerous discoveries made by the Paleontology A16 team. Hence, the present
287	study concentrates on illustrating the most significant specimens only.
288	
289	Plesiochelys sp.
290	<i>Referred material</i> . Early late Kimmeridgian of Porrentruy, Canton of Jura, Switzerla (See
291	Table 1).
292	Remarks. Twenty-four shells among the 80 studied herein lack sufficient diagnostic characters to
293	be clearly identified as either Plesiochelys bigleri or Plesiochelys etalloni. This is no surprise
294	considering how close these two species are in terms of shell anatomy. These 24 indeterminate
295	specimens are therefore provisionally referred to Plesiochelys sp. Ten of these indeterminate
296	specimens are however tentatively identified herein based on a statistical analysis of neural bone
297	thickness (see below).
298	
299	DESCRIPTION OF PLESIOCHELYS BIGLERI
300	
301	Cranium
302	
303	General description. The cranium of <i>Plesiochelys bigleri</i> is known from two specimens. The
304	first one is the holotype specimen (MJSN TCH007-252) and consists of the posterior part of a
305	skull (Fig. 4) associated with a relatively complete shell and some limb and girdle elements. The
306	parts of the skull anterior to the pterygoids (including the nasal, palatal, and orbital regions) and

the lateral part of the left otic chamber are missing. For this reason, the following bones are 307 missing from that specimen: nasal, prefrontal, frontal, postorbital, premaxilla, maxilla, vomer, 308 and palatine. Post-mortem deformation is minor, but the basicranium is partly disarticulated 309 along the basisphenoid-pterygoid suture. The preservation of this specimen is not optimal. The 310 skull was initially heavily encrusted by ferruginous mineralizations. Although most of these 311 312 mineralizations were skillfully removed during preparation, sutures remain rather difficult to see in this specimen. The length of the skull as measured from the pterygoid-vomer suture to the 313 stem of the condylus occipitalis (the condyle itself is missing) is 38.5 mm, whereas the width at 314 the level of the condyli mandibularis is 60.6 mm (Table 2). 315 The second specimen (MJSN TCH006-1451) is an isolated, partial skull missing all of the 316 skull roof, the ethmoid region, the orbital area, the anterior part of the snout, and most of the 317 palate (Fig. 5). As a result, the following bones are missing from that specimen: nasal, prefrontal, 318 frontal, postorbital, and premaxilla. The skull has been severely flattened dorsoventrally during 319 fossilization. Crushing forces were not oriented exactly dorsoventrally and resulted in a slight 320 lean toward the right hand side, mostly apparent in posterior view (Figs. 5E–5F). As preserved, 321 the skull is 59.8 mm in length from the anteriormost part of the maxilla to the condylus 322 323 occipitalis (34.4 mm from pterygoid-vomer suture to condylus occipitalis; see Table 2). The width taken at the level of the condyli mandibularis is 60 mm. MJSN TCH006-1451 is therefore 324 325 slightly smaller than the holotype specimen (Table 2). Compared to other plesiochelyids, MJSN 326 TCH006-1451 is about the same size as NMS 8738 and NMS 9145, both referred to *Plesiochelys* etalloni, but it is much smaller than OUMNH J.1582, the holotype skull of Plesiochelys 327 328 planiceps (Anguetin, Püntener & Billon-Bruyat, 2015: table 1).

329

**Parietal.** The parietals are best preserved in the holotype specimen (MJSN TCH007-252), but 330 fragments of the ventral part of the processus inferior parietalis are preserved on the right hand 331 side of the paratype specimen (MJSN TCH006-1451). The parietals form the posterior part of the 332 skull roof and meet one another medially (Fig. 4). The anterior and lateral contacts of the 333 parietals are not preserved. Most of the posterior margin of each parietal is natural. The temporal 334 335 skull roof is relatively low compared to other *Plesiochelys* spp. The upper temporal emargination largely exposes the foramen stapedio-temporale in dorsal view, but does not extend as far 336 anteriorly as to expose the processus trochlearis oticum. This is similar to the condition in *Pl*. 337 etalloni (NMB 435, NMS 40870). The development of the upper temporal emargination is not 338 well known in other plesiochelyids. The preservation of the dorsal surface of the parietals in the 339 holotype (MJSN TCH007-252) does not allow to identify scute sulci with confidence. 340 As usual in plesiochelyids (Anguetin, Püntener & Billon-Bruyat, 2015), the anterior 341 braincase wall formed mostly by the processus inferior parietalis is shorter than in most turtles as 342 a result of the great development of the foramen interorbitale. As preserved, the ventral contacts 343 of the parietal in *Pl. bigleri* are as follows: epipterygoid and pterygoid anterior to the foramen 344 nervi trigemini; pterygoid, quadrate and prootic posterior to the foramen nervi trigemini; and 345 supraoccipital posteroventrally (Figs. 4I–4L). The parietal forms the dorsal half of the anterior 346 margin of the foramen nervi trigemini. There, the parietal and pterygoid have a broad contact 347 that excludes the epipterygoid from the margin of that foramen. The posterior margin of the 348 349 foramen nervi trigemini is formed entirely by a posteroventral process of the parietal that reaches the quadrate and prevents a contact between the pterygoid and prootic. This configuration of the 350 351 region of the foramen nervi trigemini is characteristic of plesiochelyids (except for the parietal-352 quadrate contact that is absent in *Pl. planiceps*) and *Jurassichelon oleronensis* (Anguetin,

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Püntener & Billon-Bruyat, 2015). In *Pl. bigleri*, the foramen nervi trigemini is somewhat more
rounded than in other plesiochelyids, in which the foramen is usually taller than wide and oval in
shape. In most specimens of *Pl. etalloni* (NMB 435, NMS 8738, NMS 40870, NMS 40871,
NHMUK R3370), the foramen nervi trigemini is very narrow and forms a slit-like opening.

Jugal. In the holotype specimen (MJSN TCH007-252), the anterior part of the right jugal is 358 preserved as an isolated fragment. A small flake of bone (probably from the maxilla) is still 359 attached to the ventromedial margin of the isolated jugal. In the paratype specimen (MJSN 360 TCH006-1451), only the ventral part of the jugal is preserved, but the bone is more complete on 361 the right hand side (Fig. 5). The jugal forms the posteroventral corner of the orbit and contacts 362 the maxilla ventrally and the quadratojugal posteriorly. The other contacts of the jugal with 363 surrounding elements are unclear. As in *Pl. etalloni*, *Pl. planiceps*, and *J. oleronensis*, the jugal 364 lacks a medial process extending to meet the pterygoid and/or palatine. Similarly, the maxilla 365 lacks a corresponding posteromedial process and the foramen palatinum posterius remains open 366 posterolaterally. The ventral margin of the jugal and the posterior border of the maxilla show that 367 the lower temporal emargination was relatively well developed in *Pl. bigleri*, much like the 368 condition in Pl. etalloni, Pl. planiceps and J. oleronensis (note that the condition in 369 Portlandemys spp. is unknown). 370

371

Quadratojugal. The morphology of the quadratojugal is poorly known in *Pl. bigleri*. Only a
small part of the right quadratojugal is preserved anterior to the cavum tympani in the holotype
(MJSN TCH007-252). A larger portion of the right quadratojugal is preserved in the paratype
(MJSN TCH006-1451), but the bone is fragmented in several pieces and misses its dorsal and

anterior margins. Anteriorly, the quadratojugal articulates with the jugal. Its ventral margin 376 forms the posterior half of the lower temporal emargination. Posteriorly, the quadratojugal braces 377 the external margin of the cavum tympani along an extended, curved suture with the quadrate 378 (Figs. 4 and 5). It seems that the posteroventral process of the quadratojugal along the processus 379 articularis of the quadrate is proportionally shorter in *Pl. bigleri* than in *Pl. etalloni* and *Po.* 380 gracilis (unknown in other plesiochelyids). Posterodorsally, the quadratojugal has a broad 381 vertical contact with the squamosal just dorsal and slightly anterior to the level of the incisura 382 columellae auris. 383

384

Squamosal. The squamosal is best preserved in the holotype (MJSN TCH007-252), but the 385 paratype (MJSN TCH006-1451) provides additional information on the morphology of this 386 element. As usual in turtles, the squamosal forms the posterolateral corner of the otic chamber, 387 notably contributing to the formation of the antrum postoticum. The anterodorsal part of the bone 388 is unfortunately not preserved in any specimen. The contacts of the squamosal in the temporal 389 roof (notably with the postorbital and parietal) are therefore unknown in *Pl. bigleri*. The other 390 contacts of the squamosal are as follows: quadrate anteromedially and ventrally; quadratojugal 391 anterolaterally; and opisthotic posteromedially. Compared to other turtles, the antrum postoticum 392 of plesiochelyids is usually described as moderately developed (Gaffney, 1976; Rieppel, 1980; 393 Anquetin, Püntener & Billon-Bruyat, 2015). In Pl. bigleri, the antrum postoticum is also 394 395 moderately developed, but, compared to other plesiochelyids, the cavity is deeper both medially and posterodorsally. Remarkably, the anterior margin of the antrum postoticum is formed 396 397 entirely by the quadrate (MJSN TCH007-252; Fig. 4). This differs from the condition observed 398 in *Pl. etalloni* and *Po. gracilis* (as well as *J. oleronensis* and most basal pan-cryptodires) in

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which the squamosal contributes to the anterolateral margin of the antrum postoticum. The
posterodorsal parasagittal crest of the squamosal is well developed in *Pl. bigleri*, notably the
posterior part that forms an extended and somewhat pointed lamina (MJSN TCH007-252; Figs.
4I–4J). A wide concavity is present on the lateral surface of the squamosal just posterior to the
opening of the antrum postoticum. Posteroventrally, the squamosal contributes only to a limited
extent to the rugose area for the attachment of the M. depressor mandibulae (Werneburg, 2011).

Maxilla. The maxilla is not preserved in the holotype specimen. Only the posterior part of the 406 bone is preserved in the paratype (MJSN TCH006-1451), but about half of the original bone is 407 present on the right hand side in that specimen (Fig. 5). As preserved, the maxilla contacts the 408 jugal posterodorsally and the palatine posteromedially. As for the jugal, the maxilla lacks a 409 posteromedial process that would close the foramen palatinum posterius posterolaterally. The 410 maxilla seems to participate to the anterolateral margin of the foramen palatinum posterius both 411 in dorsal and ventral views. The labial ridge is slender and very high, notably posteriorly. 412 Anteriorly, the labial ridge is somewhat blunted in MJSN TCH006-1451, but this is possibly a 413 preservational artifact. The labial ridge is separated from the lingual ridge by a deep trough. As 414 415 in *Pl. etalloni*, *Pl. planiceps*, and *Po. mcdowelli* (unknown in *Po. gracilis*), the lingual ridge is broad and high. In these species, the lingual ridge is rugose, which contrasts with the condition in 416 417 J. oleronensis where the summit of the lingual ridge is acute and smooth. In *Pl. bigleri*, the 418 lingual ridge is closer to the condition in other plesiochelyids, but the bone surface is slightly eroded. In contrast to *Po. mcdowelli*, the lingual ridge is formed entirely by the maxilla. Due to 419 420 poor preservation and the presence of markings on the bone surface, we were unable to locate the 421 foramen supramaxillare in MJSN TCH006-1451.

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422

Vomer. The vomer is missing in the holotype (MJSN TCH007-252) and only the posteroventral part of the bone is preserved in MJSN TCH006-1451 (only visible in ventral view). Posteriorly, the vomer reaches the pterygoids and fully separates the palatines on the ventral surface of the palate, but not on the dorsal surface (Fig. 5). This configuration is similar to that of *Pl. etalloni* and *Po. mcdowelli*, but differs from that of *J. oleronensis*. The condition in *Pl. planiceps* is uncertain because the posteroventral part of the vomer is broken in the type and only known specimen (OUMNH J.1582).

430

**Palatine.** The palatine is only preserved in the paratype (MJSN TCH006-1451). The bone is not 431 complete and only the posterior and lateral parts can be observed (Fig. 5). As preserved, the 432 palatine contacts the pterygoid posteriorly, the vomer medially on the ventral surface of the 433 palate, the other palatine medially on the dorsal surface of the palate, and the maxilla 434 anterolaterally. Laterally, the palatine defines the anteromedial border of the foramen palatinum 435 posterius. The outline of the foramen palatinum posterius is impossible to assess in *Pl. bigleri* 436 since the lateral margin of the palatine and anterolateral part of the pterygoid are incomplete in 437 438 all of the specimens. The palatine was initially described as forming a small portion of the lingual ridge in *Plesiochelys* and *Portlandemys* (Gaffney, 1976). In fact, the palatine really 439 contributes to the lingual ridge only in Po. mcdowelli (condition unknown in Po. gracilis). In Pl. 440 441 etalloni, Pl. bigleri, Pl. planiceps and J. oleronensis, the palatine-maxilla suture extends along the medial base of the lingual ridge, i.e. on its dorsomedial edge, and the palatine therefore does 442 443 not take part into the formation of the ridge or triturating surface.

444

**Quadrate.** On the dorsal surface of the otic chamber, the quadrate contacts the prootic 445 anteromedially, the opisthotic posteromedially, and the squamosal posterolaterally. As usual, 446 guadrate and prootic contribute relatively equally to the formation of the foramen stapedio-447 temporale, but variability exists (see right side of the holotype MJSN TCH007-252). 448 Anterolaterally, the quadrate has a long curved suture with the quadratojugal. This suture is 449 450 mostly parallel to the anterior margin of the cavum tympani, but lies anterior to it (the quadratojugal does not enter the margin of the cavum tympani). The cavum tympani of Pl. 451 bigleri is deeper than in other plesiochelyids. This is notably apparent anteroventral and 452 posterodorsal to the incisura columellae auris. In contrast to other plesiochelyids, the lateral 453 margin of the cavum tympani faces more posterolaterally than laterally in *Pl. bigleri* (Figs. 4 and 454 5). The quadrate forms the entire anterior margin of the antrum postoticum, which is remarkable 455 (see Squamosal). The incisura columellae auris remains open posteroventrally. The processus 456 articularis of the quadrate is damaged in the holotype (MJSN TCH007-252), which may give the 457 misleading impression that this process is short. This structure is better preserved in the paratype 458 (MJSN TCH006-1451). A prominent ventrally-infolding ridge occurs on the posterior surface of 459 the processus articularis. Starting from the posterolateral corner of the condylus mandibularis, 460 461 this ridge extends dorsomedially toward the incisura columellae auris increasing in height. Ventral to the incisura, the ridge thickens and bends sharply medially. Pursuing its medial 462 course, the ridge thins progressively and finally merges with the posterolateral border of the 463 464 pterygoid. This ventrally-infolding ridge is found in all plesiochelyids, but also in J. oleronensis, Solnhofia parsonsi Gaffney, 1975b and Parachelys eichstaettensis Meyer, 1864, and is a strong 465 argument to support the monophyly of these Late Jurassic coastal marine turtles from Europe 466 467 (Anquetin, Püntener & Billon-Bruyat, 2015). The condylus mandibularis of Pl. bigleri is

remarkably narrow anteroposteriorly compared to other plesiochelyids. The condyle consists of 468 two slightly concave facets separated by a wide, but shallow parasagittal furrow. 469 The quadrate forms about half of the moderately developed processus trochlearis oticum, 470 which is less prominent than in *Plesiochelys etalloni*. On the anterior surface of the otic chamber, 471 the quadrate contacts the prootic dorsomedially, the parietal medially, and the pterygoid 472 473 ventromedially. This region is best preserved on the left hand side of the holotype MJSN TCH007-252. Inside the cavum acustico-jugulare, the quadrate forms the lateral part of the 474 aditus canalis stapedio-temporalis and canalis stapedio-temporalis, as well as the lateral half of 475 the posterior opening of the canalis cavernosus. Below the antrum postoticum, a rugose area 476 occurs on the ventral surface of the skull and probably served for muscular attachment. In Pl. 477 *bigleri*, this area is mostly formed by the quadrate, with minor contributions from the opisthotic 478 and squamosal. 479

480

481 **Epipterygoid.** The epipterygoid is best observable in the holotype (MJSN TCH007-252), whereas only the ventral part of the bone is preserved in the paratype (MJSN TCH006-1451). In 482 lateral view, the epipterygoid is a trapezoidal element located between the crista pterygoidea of 483 the pterygoid and the processus inferior parietalis of the parietal. The epipterygoid is exposed on 484 the medial surface of the anterior braincase wall, albeit to a more moderate extent. Posteriorly, a 485 broad contact between the parietal and pterygoid excludes the epipterygoid from the anterior 486 487 margin of the foramen nervi trigemini (Figs. 4I–4L). The same configuration seemingly occurs on the medial surface of the anterior braincase wall. Posteroventrally, a fossa cartilaginis 488 epipterygoidei is present and prevents a contact between the epipterygoid and the quadrate. The 489 490 anterolateral process of the epipterygoid that extends onto the dorsal surface of the pterygoid is

well developed. A shallow furrow prolongs this process on the dorsal surface of the pterygoid in *Pl. bigleri* (MJSN TCH006-1451). A similar process occurs in *Pl. etalloni* and *Po. mcdowelli*.
This process is more reduced in *Pl. planiceps* and *J. oleronensis*.

Pterygoid. Except for each processus pterygoideus externus, the pterygoids are complete in the paratype (MJSN TCH006-1451), but their ventral surface is somewhat abraded. Longitudinal striae extend from the pterygoid fossae posteriorly to the palatines and vomer anteriorly (Fig. 5C). In the holotype (MJSN TCH007-252), the anterior part of the pterygoids is more poorly preserved, although the left processus pterygoideus externus is complete. In contrast, the posterior part of the pterygoids (pterygoid fossa and quadrate process) is better preserved. The pterygoids are disarticulated from the basicranium in the holotype.

In ventral view, the pterygoid contacts the vomer anteromedially, the palatine anteriorly, 502 the quadrate posterolaterally, and the basisphenoid posteromedially. Posteriorly, the pterygoid 503 504 probably also contacted the basioccipital, but the sutures in this region are poorly preserved in both specimens. As preserved, the pterygoid does not seem to contact the exoccipital posteriorly, 505 but this region is rarely well preserved in plesiochelyids and the presence/absence of this contact 506 507 is probably of poor systematic value (Anquetin, Püntener & Billon-Bruyat, 2015). In ventral view, the processus pterygoideus externus is similar in development and shape to that of *Pl*. 508 509 etalloni and Pl. planiceps. However, the parasagittal plate on the lateral margin of the processus 510 pterygoideus externus is more developed in Pl. etalloni than in the two other species. A distinct ridge extends posteromedially from the posterior edge of the processus pterygoideus externus to 511 512 the posterolateral part of the ventrally open canalis caroticus internus. This ridge forms the 513 medial margin of the pterygoid fossa. Compared to other plesiochelyids, J. oleronensis and

eurysternids (Anguetin, Püntener & Billon-Bruyat, 2015), the pterygoid fossa of *Pl. bigleri* 514 stands out as being remarkably shallow. The configuration of the canalis caroticus internus is 515 similar to the condition in *Pl. etalloni* and *J. oleronensis* (see Anguetin, Püntener & Billon-516 Bruyat, 2015 for a review). The canalis caroticus internus is superficial and open ventrally at 517 least along its anterior half (Figs. 4 and 5). The posterior half of the canalis caroticus internus 518 519 may have been floored by a thin ventromedial flap of the pterygoid, but the preservation of the paratype (MJSN TCH006-1451) prevents a definitive conclusion. Since no flooring is preserved 520 in any specimen, the position of the foramen posterius canalis carotici interni cannot be 521 determined with precision. The anterior part of the canalis caroticus internus follows the 522 basisphenoid-pterygoid suture. By comparison with *Pl. etalloni*, the split between the palatine 523 and cerebral branches of the internal carotid artery was probably not floored by bone, but the 524 preservation of the specimens prevents a clear observation of the foramen posterius canalis 525 carotici palatinum (possibly visible on the left hand side of MJSN TCH006-1451) and foramen 526 posterius canalis carotici cerebralis. The flooring of the cavum acustico-jugulare by the 527 posterolateral part of the pterygoid is not very extensive. The processus interfenestralis of the 528 opisthotic therefore remains largely visible in ventral view. For that matter, *Pl. bigleri* is closer to 529 the condition observed in *Pl. planiceps*. Although this contact is now disarticulated in the 530 holotype, there was a contact between the processus interfenestralis of the opisthotic and the 531 pterygoid. Based on the disarticulated surface, this contact was probably not sutural. Lateral to 532 533 this contact, the pterygoid forms the medial part of the floor of the posterior opening of the canalis cavernosus. 534

In the ethmoid region, the pterygoid forms the ventral margin of the foramen nervi trigemini, the rest of the foramen margin being formed by the parietal (Figs. 4I–4L). The

pterygoid-parietal contact anterior to the foramen nervi trigemini excludes the epipterygoid from 537 the margin of this foramen. Medial to the crista pterygoidea, the pterygoid forms the floor of the 538 sulcus cavernosus. This region is best preserved in the paratype specimen (MJSN TCH006-539 1451). The foramen anterius canalis carotici palatinum opens in the anterior part of the sulcus 540 cavernosus about halfway between the level of the foramen anterius canalis carotici interni and 541 542 the tip of the trabecula. After exiting the foramen anterius canalis carotici palatinum, the palatine branch of the internal carotid artery continues forward in a groove within the floor of the sulcus 543 cavernosus and is not obstructed anteriorly by a crest extending from the crista pterygoidea to the 544 midline shelf of the pterygoid, as seen in some plesiochelyids Anguetin, Püntener & Billon-545 Bruyat, 2015). Anterolaterally, a small foramen occurs on the dorsal surface of the pterygoid 546 medial to the anterolateral process of the epipterygoid. This foramen may correspond to the 547 foramen nervi vidiani, but this should be further investigated (see also Anquetin, Püntener & 548 Billon-Bruyat, 2015). Anterior to this small foramen, the dorsal surface of the pterygoid forms a 549 shallow, rounded depression, which likely served for the attachment of one of the eve muscles 550 (Gaffney, 1976). Finally, as noted above (see Epipterygoid), a shallow furrow prolongs the 551 anterolateral process of the epipterygoid on the dorsal surface of the pterygoid. 552

553

**Supraoccipital.** The supraoccipital contacts the parietal anteriorly, the prootic anterolaterally, the opisthotic posterolaterally, and the exoccipital posteriorly. A broad contact between the prootic and opisthotic separates the supraoccipital from the quadrate on the floor of the fossa temporalis superior (Figs. 4 and 5). This contrasts with the condition observed in *Po. mcdowelli* and most specimens referred to *Pl. etalloni*. Most of the crista supraoccipitalis is preserved in the holotype MJSN TCH007-252. This structure is relatively low, especially compared to *Pl.* 

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*planiceps* and *Po. mcdowelli*. The condition in *Pl. etalloni* is more difficult to appreciate based on the available material, but it seems to be intermediate in development between *Pl. bigleri* and the two aforementioned species. The posterior part of the crista supraoccipitalis is broken in MJSN TCH007-252, but it is unlikely that this structure projected far behind the level of the condylus occipitalis. As usual, the supraoccipital forms the dorsal part of the foramen magnum.

**Exoccipital.** The exoccipital contacts the supraoccipital dorsomedially, the opisthotic 566 laterodorsally, the basioccipital ventrally, and the processus interfenestralis of the opisthotic 567 anterolaterally. As preserved, there is no contact between the exoccipital and pterygoid, which 568 may be a difference with *Pl. etalloni*, but this contact is often difficult to observe in this taxon 569 (Anguetin, Püntener & Billon-Bruyat, 2015). An exoccipital-pterygoid contact is otherwise 570 present in *Portlandemys* ssp. and J. oleronensis. In contrast to *Pl. etalloni*, *Pl. planiceps*, *Po.* 571 mcdowelli (condition unknown in Po. gracilis), and J. oleronensis, the exoccipitals apparently do 572 not meet in the floor of the foramen magnum, but this feature can present some intraspecific 573 variability (Anguetin & Chapman, 2016). The contribution of the exoccipital to the condylus 574 occipitalis is uncertain, but probably reduced. There are two foramina nervi hypoglossi on each 575 576 side and they are formed exclusively by the exoccipital.

577

Basioccipital. The ventral aspect of the basioccipital is poorly preserved in all specimens.
However, it is apparent that the basioccipital contacts the basisphenoid anteriorly and the
pterygoid anterolaterally in this area. The tubercula basioccipitale are only moderately developed
and correspond to what is known in similarly-sized individuals of *Pl. etalloni*, but not in larger
specimens (Anquetin & Chapman, 2016). Posterodorsally, there is an extensive contact with the

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exoccipital, but the basioccipital nevertheless enters the ventral margin of the foramen magnum.
It appears that the basioccipital forms most of the condylus occipitalis, which is only preserved
(poorly) in MJSN TCH006-1451. In dorsal aspect, the basioccipital offers little remarkable
features. The basis tuberculi basalis is relatively low, and an oval depressed area occurs in front
of it. Finally, there is also a contact with the ventromedial margin of the processus
interfenestralis of the opisthotic in the floor of the recessus scalae tympani.

589

**Prootic.** On the dorsal surface of the otic chamber, the prootic contacts the parietal 590 anteromedially, the quadrate laterally and anteroventrally, the opisthotic posteriorly, and the 591 supraoccipital posteromedially. The prootic forms the medial half of the processus trochlearis 592 oticum, which is reduced compared to that of *Pl. etalloni*. Anteromedially, the prootic is 593 excluded from entering the posterior margin of the foramen nervi trigemini by an elongate 594 descending process of the parietal (Figs. 4I–4L). A broad contact between this descending 595 process of the parietal and the quadrate prevents a prootic-pterygoid contact in this area. This 596 configuration is found in all plesiochelyids but Pl. planiceps, as well as in J. oleronensis and 597 possibly also in eurysternids (Anguetin, Püntener & Billon-Bruvat, 2015). 598

Inside the cavum acustico-jugulare, the prootic forms the anterior half of the fenestra ovalis, the medial part of the aditus canalis stapedio-temporalis, and most of the roof of the posterior opening of the canalis cavernosus. As in other plesiochelyids, the aditus canalis stapedio-temporalis is located more posteriorly than the posterior opening of the canalis cavernosus rather than in the roof of the latter as in many turtles. In this area, the prootic contacts the opisthotic posterodorsally, the pterygoid ventrally, and the quadrate laterally. A C-shaped furrow occurs on the surface of the prootic dorsolateral to the fenestra ovalis. Inside the cavum

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cranii, the prootic contacts the supraoccipital posterodorsally, the parietal anterodorsally, the
basisphenoid ventromedially, and the pterygoid anteroventrally. In contrast to *Pl. etalloni*, the
pila prootica is not ossified. Medial to the foramen nervi trigemini, the prootic forms a welldefined recess that accommodated the ganglion of the trigeminal (V) nerve.

610

611 **Opisthotic.** On the dorsal surface of the otic chamber, the opisthotic contacts the prootic anteriorly, the quadrate anterolaterally, the squamosal posterolaterally, the supraoccipital 612 medially, and the exoccipital posteromedially. The anterior contact with the prootic is relatively 613 wide and clearly separates the supraoccipital and quadrate (Figs. 4 and 5). This condition is 614 found also in *Pl. planiceps*, *Po. gracilis*, and *J. oleronensis*. In contrast, the prootic-opisthotic 615 contact is reduced or absent in *Po. mcdowelli* and most specimens of *Pl. etalloni*. The posterior 616 aspect of the opisthotic is complete only on the right hand side of the paratype (MJSN TCH006-617 1451). The extremity of the processus paroccipitalis forms a distinct crest for muscular 618 attachment. In Pl. bigleri, as well as in Po. mcdowelli and J. oleronensis, the processus 619 paroccipitalis extends posterolaterally. This contrasts with *Pl. etalloni*, *Pl. planiceps*, and *Po.* 620 gracilis in which the processus paroccipitalis extends more posteriorly. This difference changes 621 622 the occipital outline as seen in dorsal view from a broad arch in *Pl. bigleri* to a more narrow arch in Pl. etalloni. 623

Ventrally, the opisthotic forms a large part of the roof of the cavum acustico-jugulare. In this area, the opisthotic contacts the prootic anteriorly, the quadrate anterolaterally, the squamosal posterolaterally, and the exoccipital posteromedially. The extremity of processus interfenestralis is triangular in shape and contacts the exoccipital, basioccipital, and pterygoid. The medial margin of the processus interfenestralis is pierced by a large fenestra perilymphatica,

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which is apparently entirely contained in bone thanks to a ventromedial contribution from the
basioccipital (MJSN TCH007-252). A well-defined foramen externum nervi glossopharyngei
opens at the base of the processus interfenestralis. It is noteworthy that this foramen opens more
laterally than in other plesiochelyids and *J. oleronensis*. Lateral to the base of the processus
interfenestralis and foramen externum nervi glossopharyngei, the opisthotic forms a strong
curving ridge oriented anteroposteriorly. As far as we know, this ridge is not present in other
plesiochelyids.

636

**Basisphenoid.** In ventral aspect, the basisphenoid contacts the pterygoid anteriorly and laterally, 637 and the basioccipital posteriorly (Figs. 4 and 5). As noted above, the basisphenoid-pterygoid 638 suture is disarticulated in the holotype, but this contact is better preserved in the paratype (MJSN 639 TCH006-1451). The posterior contact with the basioccipital is poorly preserved in both 640 specimens. The general outline of the basisphenoid in ventral view is triangular, as in most 641 plesiochelvids (Anguetin, Püntener & Billon-Bruvat, 2015). The ventral surface of the 642 basisphenoid is apparently slightly concave. The morphology of the canalis caroticus internus is 643 similar to the condition observed in *Pl. etalloni* and *J. oleronensis* (Anguetin, Püntener & Billon-644 Bruyat, 2015). The canalis caroticus internus is superficial and runs along the basisphenoid-645 pterygoid suture as a ventrally open canal. The posterior part of the canalis caroticus internus 646 may have been partly floored by a ventromedial flap formed by the pterygoid, but this region is 647 648 not well preserved in any specimen. Based on the available material, the position of the foramen posterius canalis carotici interni cannot be determined. As in *Pl. etalloni*, the split between the 649 cerebral and palatine branches of the internal carotid artery was probably not floored by bone. 650 651 However, the preservation prevents a clear observation of the foramen posterius canalis carotici

palatinum. In the holotype, a portion of each canalis caroticus cerebralis is preserved on the 652 anterior part of the basisphenoid. Anteriorly, these two canals penetrate deeply in the 653 basisphenoid and exit in the sella turcica dorsally. A small unnamed foramen opens 654 dorsomedially along the canalis caroticus cerebralis (only visible in MJSN TCH007-252; Fig. 4). 655 The dorsal aspect of the basisphenoid is better preserved in the paratype (MJSN TCH006-656 1451) and can be readily observed in that specimen thanks to the loss of the anterior part of the 657 roof of the cavum cranii (Fig. 6). The basioccipital contacts the basioccipital posteriorly, the 658 prootic posterolaterally, and the pterygoid anterolaterally and anteriorly. The part of the 659 basisphenoid that floors the cavum cranii is slightly concave. A moderately raised area located 660 sagittally on the posterior margin of the basisphenoid corresponds to the anterior part of the basis 661 tuberculi basalis. The posterior foramen nervi abducentis opens about midway along the part of 662 the basisphenoid that floors the cavum cranii. The anterior foramen nervi abducentis opens 663 ventral and slightly anteromedial to the base of the processus clinoideus, relatively close to the 664 basisphenoid-pterygoid suture. This is the condition usually found in plesiochelyids, with the 665 exception of Pl. etalloni (Anquetin, Püntener & Billon-Bruyat, 2015). The processus clinoideus 666 is formed just medial to the basisphenoid-prootic suture. In contrast to *Pl. etalloni*, the pila 667 prootica is not ossified. The dorsum sellae is high and does not overhang the sella turcica, as 668 usual in plesiochelyids. The surface below the dorsum sellae is devoid of ridge and slopes 669 relatively gently anteriorly. In that matter, Pl. bigleri is intermediate between Pl. etalloni, in 670 671 which the surface below the dorsum sellae is near vertical, and *Portlandemys* spp., in which that surface slopes very gently anteriorly resulting in a relatively long distance between the dorsum 672 673 sellae and the foramina anterius canalis carotici cerebralis (Anguetin, Püntener & Billon-Bruyat, 674 2015). The foramina anterius canalis carotici cerebralis are separated by a broad bar of bone,

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which is unusual in plesiochevids, and open slightly posterior to the level of the foramina 675 anterius canalis carotici palatinum. The trabeculae are relatively short and straight, and frame a 676 small, well-defined sella turcica. The rostrum basisphenoidale is short, and the anterior tip of the 677 trabeculae actually represents the anteriormost extension of the basisphenoid in dorsal view. 678 679 680 Carapace 681 General description. Elements of the carapace are preserved in 39 out of the 41 shells herein 682 referred to Plesiochelys bigleri (Figs. 7 and 8). The largest specimens reach a carapace length of 683 about 550 mm (MJSN BSY006-307, MJSN SCR011-30, MJSN TCH005-21), whereas the 684 smallest have a carapace length of about 220-250 mm (MJSN BSY008-848, MJSN SCR010-685 327, MJSN TCH007-516). Measuring carapace length with precision in disarticulated and 686 incomplete specimens is not possible, but we estimate that most specimens are about 450 mm in 687 carapace length, which corresponds approximately to what is known for *Pl. etalloni* in Solothurn. 688 Most shells of Pl. bigleri are disarticulated, but two shells (MJSN BSY006-307 and MJSN 689 TCH007-519) were found articulated. Bones were prepared out of the marly matrix individually 690 691 and we were able to reassemble them on a moldable sand bed and reconstruct the 3D shape of the shell. Post-mortem deformation occurred in most specimens and to variable extents. Our 692 reconstructions are therefore tentative. The resulting carapace outlines are either evenly oval 693 694 (e.g., MJSN TCH006-767), or roundish (e.g., MJSN TCH007-252; Figs. 7A-7B). In some specimens the posterior carapace part is pointed (e.g., MJSN BSY009-815, MJSN SCR011-30). 695 The shells of *Pl. bigleri* and *Pl. etalloni* are very similar. The following description 696 697 therefore primarily focuses on the few differences between the two species and the intraspecific

variations observed in *Pl. bigleri*. A general description of the shell morphology of *Pl. etalloni*and a discussion of its intraspecific variability can be found elsewhere (Anquetin, Deschamps &
Claude, 2014; Anquetin, Püntener & Billon-Bruyat, 2014).

701

702 Nuchal. The shape of the nuchal in Pl. bigleri varies from almost rectangular (e.g., MJSN

SCR011-140; Figs. 8I–8J) to trapezoidal (e.g., MJSN BSY007-257). This element can be about

as wide as long (e.g., MJSN BSY009-815) or clearly wider than long (e.g., MJSN SCR011-160).

The nuchal notch is usually shallow, but can also be much reduced (e.g., MJSN SCR011-160) or

strongly pronounced (e.g., MJSN BSY007-257). In some specimens (e.g., MJSN TCH006-1420;

Figs. 8A–8B), the posteromedial part of the nuchal that articulates with the first neural projects

708 posteriorly.

709

Neurals, suprapygals and pygal. Like Pl. etalloni, Pl. bigleri usually has eight neurals, one 710 intermediate element, two suprapygals, and one pygal. However, this condition may vary. Neural 711 bones can be split in two (e.g., neural 2 in MJSN SCR010-342), fused together (e.g., neural 8 712 with the intermediate element in MJSN TCH005-21), or be reduced (e.g., neural 7 in MJSN 713 SCR011-140) or completely obliterated (e.g., neural 7 in MJSN SCR011-413 and MJSN 714 TCH006-1420; Figs. 8A-8B) by costals with midline contact. The first neural is rectangular to 715 oval in shape. It usually tapers posteriorly (not in MJSN BSY007-257 and MJSN TCH005-42; 716 717 Figs. 8M–8N). Neurals 2 to 6 are elongated hexagons with shorter sides facing anteriorly. However, these hexagons are often deformed (e.g., MJSN BSY007-257, MJSN SCR010-342, 718 719 MJSN SCR011-30, MJSN SCR011-276, MJSN SCR011-413, MJSN TCH005-42, MJSN 720 TCH006-767). Neurals 7 and 8 are shorter and more irregular in shape than the preceding
neurals. Their growth is often constrained by a midline contact of the costals. Neural 8 is usually
the smallest bone in the series. In some specimens, it is pentagonal with shorter sides facing
posteriorly (e.g., MJSN BSY009-815, MJSN TCH007-252; Figs. 7A–7B). The thickness of the
neural bones in *Pl. bigleri* is significantly reduced compared to *Pl. etalloni* (Fig. 9). This is
especially obvious for neurals 2 to 5. We statistically demonstrate this difference herein (see
below).

As relatively common in turtles (Zangerl, 1969), the posteromedial region of the carapace 727 is characterized by a great deal of intraspecific variation in *Pl. bigleri*. The element that follows 728 neural 8 corresponds to the intermediate element of Anguetin, Püntener & Billon-Bruyat (2014). 729 This intermediate element varies much in shape, but usually tapers anteriorly (e.g., MJSN 730 SCR011-140, MJSN SCR011-160, MJSN SCR011-276). In one specimen (MJSN TCH006-731 1420; Figs. 8A–8B) the intermediate element is fused to the first suprapygal. 732 Plesiochelys bigleri usually has two suprapygals that are clearly wider than long. Although 733 734 irregular in shape (usually from trapezoidal to sub-triangular), the first suprapygal usually tapers anteriorly and the second posteriorly (e.g., MJSN BSY009-815, MJSN SCR011-160, MJSN 735 TCH006-1420; Figs. 8A–8B). The two suprapygals can fuse into a single element (e.g., MJSN 736 737 TCH007-519, MJSN SCR011-148). In some specimens, the first suprapygal is divided in two (e.g., MJSN SCR011-30, MJSN TCH007-252; Figs. 7A–7B). The pygal is a trapezoidal element 738 739 that is wider than long. Its relative size varies substantially from one individual to another. 740 **Costals and peripherals.** *Plesiochelys bigleri* has eight costals and eleven peripherals. Their 741

shape and arrangement are relatively stable within the species. The length-width ratio of costal 4

corresponds to that of *Pl. etalloni* and differs notably from that of *Craspedochelys jaccardi* 

(Anquetin, Püntener & Billon-Bruyat, 2014). The costals in *Pl. bigleri* are not as thick as the 744 same elements in *Pl. etalloni*. In the latter the proximal part of the costals is very thick, matching 745 746 the thick neural bones, and the costals usually remain relatively thick on their distal margin. In Pl. bigleri, both the proximal and distal parts of the costals are thinner. A midline contact occurs 747 frequently between costals 7 (e.g., MJSN SCR011-140; Figs. 8I–8J), sometimes also between 748 749 costals 6 (e.g., MJSN SCR011-30, MJSN TCH006-1420; Figs. 8A-8B), and rarely between costals 8 (e.g., MJSN TCH005-42; Figs. 8M-8N). Such a reduction of posterior neurals and 750 midline contacts of costals can also occur in *Pl. etalloni* (e.g., MAJ 2005-11-1, MJSN TCH006-751 574), but more rarely than in *Pl. bigleri*. This condition appears to occur more commonly in 752 Craspedochelys jaccardi (Anquetin, Püntener & Billon-Bruyat, 2014). 753 754

Scutes of the carapace. There are three cervical scutes of about equal size in *Pl. bigleri* (best
preserved in MJSN CRT007-2 and MJSN TCH007-519). However, as common in plesiochelyids
(see Anquetin, Püntener & Billon-Bruyat, 2014), cervical scutes are difficult to discern in many
specimens.

There are usually five vertebrals, four preurals, and twelv parginals. Vertebrals 1 and 5 759 760 are always the shortest in the series. The second and third vertebrals are rectangular (e.g., MJSN BSY009-815) to hexagonal (e.g., MJSN SCR011-30) in shape and cover one third (e.g., MJSN 761 762 TCH007-519) to about half (e.g., MJSN TCH006-767) of the costal length laterally. Vertebral 4 763 is usually hexagonal in shape. In some specimens it almost extends as far as the peripherals laterally, significantly reducing the size of pleural 4 (e.g., MJSN BSY006-326 and MJSN 764 SCR011-160). The intervertebral scute sulci usually run on neurals 1, 3, and 5. The scute sulcus 765 766 between vertebrals 4 and 5 most often runs on the intermediate element (e.g., MJSN BSY009-

815), but can also cross neural 8 (e.g., MJSN SCR011-148) or the first suprapygal (e.g., MJSN 767 TCH007-252; Fig. 7B). The twelfth pair of marginals is either restricted to the pygal and 768 peripheral 11 (e.g., MJSN BSY006-326, MJSN TCH006-767), or extends on the second 769 suprapygal (e.g., MJSN TCH006-1420, MJSN SCR011-140; Figs. 8B and 8J). 770 One specimen (MJSN TCH005-42; Figs. 8M–8N) shows an anomalous scute pattern. It 771 772 has eight partially wedged vertebrals and five irregularly sized pleurals (maybe even six on the right side). This scute pattern shows no symmetry and we consider it as an anormal condition in 773 Pl. bigleri. 774 775 Plastron 776 777 General description. Elements of the plastron are preserved in 40 out of the 41 shells herein 778 referred to Pl. bigleri. With 481 mm, MJSN SCR10-1279 has the longest preserved plastron. In 779 contrast, the well preserved juvenile specimen MJSN SCR010-327 has a plastron length of only 780 194 mm. The anterior plastral lobe of *Pl. bigleri* is often somewhat quadrangular in outline (e.g., 781 MJSN SCR010-1196, MJSN SCR011-140; Figs. 8K–8L), but can also be rounded (e.g., MJSN 782 783 CRT007-2), or even pointed (e.g., MJSN TCH006-1420; Figs. 8C-8D) in some specimens. In Pl. etalloni, the anterior plastral lobe usually has a rather rounded, sometimes pointed, anterior 784 outline (Anquetin, Püntener & Billon-Bruyat, 2014). Like in Pl. etalloni, a central plastral 785

fontanelle (always longer than wide) is occasionally present (e.g., MJSN TCH006-1420; MJSN

787 BSY006-307; Figs. 8C–8D and 8G–8H).

788

**Epiplastra and entoplastron.** The shape of the epiplastra varies from rather rectangular (MJSN 789 SCR010-1196) to triangular (e.g., MJSN TCH006-1420; Figs. 8C-8D). The anterolateral border 790 of the epiplastra is often somewhat angular, mirroring the anterior outline of the anterior plastral 791 lobe. Epiplastral bulbs, as described in Pl. etalloni (Bräm, 1965; Anguetin, Püntener & Billon-792 Bruyat, 2014), are absent or only weakly expressed (e.g., MJSN TCH007-252; Fig. 7C). The 793 794 entoplastron is usually roundish (e.g., MJSN SCR011-140; Figs. 8K-8L) to roughly kite-shaped (e.g., MJSN TCH007-252; Figs. 7C–7D), but its relative size and length-width proportion varies 795 greatly from one individual to another. The entoplastron of the juvenile specimen MJSN 796 797 SCR010-327 is elongated and kite-shaped.

798

**Hyo-**, hypo-, and xiphiplastra. The hypolastra are always longer than wide, although only by a 799 small amount in a few individuals (e.g., MJSN BSY009-815). This corresponds to what is known 800 in *Pl. etalloni* and many other turtles, and contrasts with the condition in *C. jaccardi*, where the 801 802 hyoplastra are clearly wider than long (Anguetin, Püntener & Billon-Bruyat, 2014). The suture between the hyo- and hypoplastra is generally straight, but sometimes shows individual 803 symmetric anteroposterior projections (e.g., MJSN CRT007-2, MJSN BSY006-307; Figs. 8G-804 805 8H). There is a small supernumerary bone between the hypplastra of the juvenile specimen MJSN SCR010-327, but the sutures of this roundish bone are absent on the visceral side of the 806 plastron. The posterolateral borders of the hypoplastra often project so as to articulate with the 807 808 corresponding notches of the xiphiplastra (e.g., MJSN SCR011-413). The xiphiplastra are always longer than wide and often asymmetric. In most specimens, the xiphiplastra are relatively long 809 810 elements, possibly more elongated in proportion than in *Pl. etalloni*. However, in some

individuals the xiphiplastra appear to be significantly reduced in length, but still longer than wide
(e.g., MJSN BSY009-815).

813

Scutes of the plastron. The gular and extragular scute sulci are relatively shallow. In *Pl. etalloni*, the gular and extragular sulci are usually deeper, notably anteriorly, as a result of the
presence of the epiplastral bulbs. As in *Pl. etalloni*, the gular scute is either restricted to the
epiplastra (e.g., MJSN CRT007-2), or extends to the entoplastron (e.g., MJSN TCH007-252; Fig.
7D). In those specimens where the plastral midline sulcus is discernable, it is irregularly sinuous,
sometimes creating small supernumerary scutes (e.g., MJSN SCR011-30). A similar condition is
also known in *Pl. etalloni*.

Generally, the sulcus between the humeral and pectoral scutes is straight, while the sulcus between the pectoral and abdominal scutes is curving anteriorly. The sulcus between the femoral and anal scutes is usually restricted to the xiphiplastron, but extends to the hypoplastron in some specimens. In the latter case, the anal scutes may form a rectangular anterior projection on the hypoplastra (e.g., MJSN SCR011-30). A similar variability is present in *Pl. etalloni* (Anquetin, Püntener & Billon-Bruyat, 2014).

There are usually four pairs of inframarginal scutes. As in *Pl. etalloni*, the third one is generally the longest in the series. It covers the suture between hyo- and hypoplastron. The juvenile specimen MJSN SCR010-327 shows a variation of this pattern by having three inframarginal scutes on the left hypoplastron. It is however unclear whether this resulted in a total of five inframarginal scutes on the left side and whether this was also the case on the right side. The inframarginal scutes usually extend laterally on the peripherals. Occasionally some of

them are restricted to plastral elements (e.g., MJSN CRT007-2, MJSN TCH006-767). This is
also a condition known in *Pl. etalloni* (Anquetin, Püntener & Billon-Bruyat, 2014).

#### 836 **Pectoral girdle**

Eight partially preserved scapulae are associated with shells of *Pl. bigleri* (MJSN BSY008-567,

838 MJSN BSY009-815, MJSN SCR011-30, MJSN SCR011-148, MJSN TCH007-252, MJSN

839 TCH007-519, MJSN TCH005-42; Fig. 10). The two scapulae of MJSN BSY008-567 are

attached to the visceral side of the plastron by sediment, approximately in situ and in contact

841 with the humeri. The other six scapulae are disarticulated.

The dorsally projecting scapular process and anteromedially projecting acromion process 842 are elliptic cylinders. The distal parts of these processes are missing in most specimens. Only the 843 acromion process in the right scapula of MJSN BSY008-567 is probably complete. It measures 844 59 mm from its distal end to the notch between the scapular process and the coracoid. The two 845 processes form a scapular angle of 102° in MJSN TCH005-42 (Fig. 10), the only scapula of Pl. 846 bigleri where this angle can be measured with confidence. None of the scapulae associated with 847 *Pl. bigleri* provides clear information about the nature of the glenoid fossa or about the articular 848 849 surface for the coracoid. As in other plesiochelyids, as well as thalassemydids and eurysternids, the glenoid neck is well developed. 850

The posteromedially projecting coracoid itself is only partially preserved in MJSN TCH005-42 (proximal part) and MJSN TCH007-519 (distal part, partially covered by bones and sediment). It is unclear whether the disarticulated coracoid of MJSN TCH005-42 belongs to the left (preserved) or right (unpreserved) scapula. The short diaphysis is irregularly cylindric in cross section. It broadens proximally where it articulates with the scapular neck. The distal end

of the bone is missing, but the broadening diaphysis indicates yet the beginning of the coracoid 856 blade. The latter is partially observable in MJSN TCH007-519, where it forms a broad blade. 857 The scapula is also known in Pl. etalloni (NMS 8584, NMS 8731, NMS 9153 and NMB 858 435), as well as in *Thalassemvs bruntrutana* and *Thalassemvs hugii* (Püntener, Anguetin & 859 Billon-Bruyat, 2015). The scapular angle of *Pl. bigleri* (see above) corresponds to the angle 860 861 previously measured for *Pl. etalloni* in specimens from Solothurn and contrasts with the wider angles in Th. bruntrutana and Th. hugii (Püntener, Anquetin & Billon-Bruyat, 2015: table 1). 862 The scapula of *Pl. etalloni* does not show any significant anatomical differences to the scapula of 863 Pl. bigleri. 864

865

#### 866 Humerus

Four humeri are associated with shells of *Pl. bigleri* (the proximal parts of both humeri of MJSN
BSY008-567 and MJSN TCH007-252 respectively). While both humeri of MJSN TCH007-252
(Fig. 11) are disarticulated, the humeri of MJSN BSY008-567 are attached to the visceral side of
the plastron by sediment, approximately in situ and in contact with the scapulae.

The proximal articulation is a hemispherical head that projects dorsally with approximately 871 135° from the horizontal plane of the humerus (only measurable in the humeri of MJSN 872 TCH007-252). The anteriorly expanding lateral process is the smaller one, as in most cryptodires 873 (Gaffney, 1990). A strong deltopectoral crest projects ventrally from its lateral border (best 874 875 visible in the left humerus of MJSN TCH007-252; Figs. 11F–11G). The larger, posteriorly expanding medial process is only slightly bulged ventrally, so that the intertubercular fossa is 876 877 mainly defined by the deltopectoral crest. More distally, the narrowing diaphysis forms a waist 878 almost circular in section.

The humerus is also partly known in *Pl. etalloni* (NMS 8584 and NMB 435) and 879 Tropidemys langii (MJSN VTT006-253 and MJSN VTT010-17). The observed features of the 880 humerus of *Pl. bigleri* correspond in all aspects to the humerus of *Pl. etalloni*. According to 881 Püntener et al. (2014), the humerus of Tr. langii (MJSN VTT006-253) has a more expanded 882 medial process and a broader and flatter diaphysis. However, a more recent discovery of a 883 884 humerus of Tr. langii (MJSN VTT010-17) contradicts this statement and suggests that the observed features of MJSN VTT006-253 are probably due to postmortem flattening. On the base 885 of the new material, the humerus of Tr. langii does not show significant anatomical differences 886 to the humerus of *Pl. bigleri* and *Pl. etalloni*. 887

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#### 889 Radius and Ulna

The radius and ulna are only known in the holotype specimen (MJSN TCH007-252; Figs. 12A– 12H). The ulna can be confidently identified as a right ulna, but the radius is more poorly preserved and is only tentatively identified as a right one. The radius is a slim bone with a cylindric diaphysis of only 5 mm diameter at its narrowest point (Figs. 12A–12D). The bone is twisted, so that the proximal and distal heads stand perpendicular to each other. Although not completely preserved, it is clear that the distal expansion was broader and flatter than the proximal one, which is a common condition in turtles.

The ulna is strongly bent along the long axis due to postmortem deformation (Figs. 12F and 12H). It is clearly broader than the radius. The proximal head forms a moderately concave sigmoid notch, and the olecranon is poorly developed. Below the proximal head, the medial surface of the bone bears a well developed bicipital tubercle (Figs. 12E–12G). The diaphysis is

901	flat and 7 mm broad at its narrowest point. Towards the distal end, the bone remains flat and
902	broadens up to 16 mm. The articulation for the carpals is not preserved.
903	Bräm (1965) briefly described one radius of Pl. etalloni (NMS 8731). Unfortunately, only
904	a fragment of this radius has been preserved into the present day, leaving only Bräm's
905	description as a reference. According to Bräm (1965), the distal expansion of the radius of <i>Pl</i> .
906	etalloni is larger and more stoutly built than the proximal expansion.
907	Both ulnae of the same specimen (NMS 8731) are preserved. The distal expansion of these
908	bones is quadrangular in shape. In contrast, this part is triangular shaped in <i>Pl. bigleri</i> . However,
909	it is not clear whether this is a real anatomical difference or whether it is due to an insufficient
910	preservation of MJSN TCH007-252.
911	

#### 912 Pelvic girdle

Ten partially preserved pelves are associated with shells of *Pl. bigleri* (MJSN BSY006-307, 913 MJSN BSY-147, MJSN BSY009-815, MJSN SCR010-30, MJSN SCR011-148, MJSN SCR011-914 413, MJSN TCH006-767, MJSN TCH006-1420, MJSN TCH007-252, MJSN TCH007-519). The 915 following description is mainly based on the sub-complete pelvis of MJSN BSY006-307 that is 916 917 still articulated in its original three dimensional shape, though it suffered minor postmortem deformation (Figs. 13). Right and left halves of the pelvis are still interconnected posteriorly by 918 the ischium plate. The anterior midline connection of the pubes is not preserved. Although both 919 920 acetabula are completely preserved on both sides of the pelvis, the ilium and pubis are much better preserved on the right side. 921

The acetabulum is a relatively deep and somewhat kidney-shaped cavity (Fig. 13). Its longer axis (about 35 mm long in MJSN BSY006-307) lies in the horizontal plane, its shorter

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axis (about 20 mm) in the vertical plane. The three bones of the pelvis (ilium, pubis, ischium) 924 form the acetabulum with their proximal parts. The sutures between these bones are best visible 925 on the medial surface of the right acetabular region in MJSN BSY006-307. 926 The ilium extends posterodorsally (Fig. 13). Proximally, where it contributes to the 927 acetabulum, its maximal width reaches 29 mm. More distally it narrows to form an irregularly 928 929 oval, slightly twisted shaft (about 15 mm wide in MJSN BSY006-307). At the distal end, the ilium is flatter and expands in the anteroposterior direction. However, the posterodorsal iliac 930 process and the anterodorsal articulation surface with the sacral rib are only very partially 931 preserved in *Pl. bigleri* (MJSN TCH007-519 and MJSN SCR011-30 respectively). 932 The proximal part of the pubis extends anteroventrally approximately in the same axis as 933 the ilium (Fig. 13). It narrows below the acetabulum, but the shaft remains much broader and 934 flatter than the ilium shaft. Distally, the pubis divides into two parts: a ventrally orientated lateral 935 pubic process and an anteriorly orientated part that normally forms the thyroid fenestra together 936 with the ischium. The lateral pubic process is about 24 mm long in MJSN BSY006-307. At the 937 distal end it has an oval articulation surface (only partially preserved in MJSN BSY006-307) that 938 rested on the dorsal surface of the xiphiplastra. The anteromedial part of the pubis is broken in 939 940 MJSN BSY006-307, and the shape of the thyroid fenestra remains unclear in *Pl. bigleri*. The ischium extends posteroventrally, first narrowing into a short, circular shaft, then 941 broadening again medially in order to form a broad plate that meets the other ischium medially. 942 943 This ischial plate is concave dorsally. The contact between ischium and pubis at the anterior margin of this plate is not preserved in any specimen. The lateral ischial process is strong and 944 extends posteriorly from the ischium. Due to postmortem deformation, the right process is 945 946 strongly bent dorsally in MJSN BSY006-307. The posterior margin of the ischium, between the

lateral ischial process and the midline contact with the other ischium, forms a shallow 947 depression. The ventral surface of the ischial plate is generally convex. It bears a shallow V-948 shaped rugose area that points anteriorly. 949 The exceptional preservation of the pelvis of MJSN BSY006-307 is unique among 950 plesiochelyids. Bräm (1965) described the pelvis of *Pl. etalloni* (mainly based on NMS 8731) 951 952 and C. jaccardi (based on NMS 8713–8718), but this material is strongly fragmented and deformed. The observable features correspond fairly well with the pelvis of *Pl. bigleri*, namely 953 the general shape of the ilium (well visible in NMS 8731). Bräm (1965) concluded that there are 954 no significant differences between the pelvis of *Pl. etalloni* and *C. jaccardi*. 955

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#### 957 Femur

Three femora are associated with shells of *Pl. bigleri* (both of MJSN SCR010-1279 and one of MJSN TCH005-21). The femur of MJSN TCH005-21 is strongly deformed and its proximal part is badly damaged. Only the proximal head of the left femur of MJSN SCR010-1279 is preserved. The right femur of MJSN SCR010-1279 is almost completely preserved, but most of the bone surface is covered by encrustation. The following description is therefore mainly based on the right femur of MJSN SCR010-1279 (Figs. 14A–14D).

The femur of *Pl. bigleri* is essentially a straight bone that is only slightly arched dorsally. The dorsally projecting proximal head and the ventrally projecting distal articulations give the left femur an elongated S-shape in posterior view (Fig. 14D). The right femur of MJSN SCR010-1279 is 134 mm long. The proximal head projects from the long axis of the femur at an angle of about 50° (Figs. 14B and 14D). It is hemispherical, but elongated along the long axis of the femur (about twice as long as wide), which is apparently more consistent with swimming than

walking (Zug. 1971). This contrasts with the more roundish femoral head of Tr. langii (Püntener 970 et al., 2014). In ventral view, the two trochanters form a deep, narrow, and V-shaped 971 intertrochanteric fossa (Fig. 14C). The posteriorly situated trochanter major is slightly shorter 972 than the trochanter minor (anteriorly), but expands more prominently on the horizontal plane. 973 Lateral ridges on both trochanters form a shallow V-shaped depression just distal to the 974 975 intertrochanteric fossa, giving the latter a terraced appearance. The diaphysis is oval to circular in cross section. The bone is gradually broadening towards 976 the distal end. The condyles are only slightly less expanded than the trochanters. The medial 977 condyle is strongly arched and tappers proximally. The lateral condyle is not completely 978 preserved, but it seems smaller and more roundish than the medial condyle. A deep fossa (about 979 6 mm deep) separates the two articulation surfaces from each other (Fig. 14C). 980 Within plesiochelyids, the femur is partly known in *Pl. etalloni* (NMS 8584 and MNS 981 8731), C. jaccardi (NMS 8713-8718), and Tr. langii (MJSN VTT010-13). Based on this 982 incomplete material, the femur of *Pl. bigleri*, *Pl. etalloni* and *C. jaccardi* cannot be 983 differentiated. On the other hand, these species differ from Tr. langii, in which the trochanters 984 expand more prominently along the horizontal plane and the intertrochanteric fossa is shallower, 985 986 wider, and more rounded at its base (Püntener et al., 2014). 987

#### 988 Fibula

789 Two left fibulae are associated with shells of *Pl. bigleri* (MJSN TCH005-21 and MJSN SCR010-

1279). The proximal third of the fibula of MJSN TCH005-21 is missing. The fibula of MJSN

991 SCR010-1279 is complete (Figs. 14E–14H). It is 84 mm long, which corresponds to 63% of the

992 femur length. Proximally, the fibula is only slightly expanded and has a small, hemispherical

articulation surface facing moderately ventrally. A swelling on the medial edge probably marks
the attachement site of the proximal tibiofibular ligament (Figs. 14E–14G). The shaft is a rather
flat, elliptic cylinder. At its narrowest point it has only one third of the width of the proximal
expansion. Distally, the fibula is almost twice as expanded as proximally. Here the bone is
slightly concave dorsally and has a somewhat triangular articulation surface facing moderately
ventrally (Figs. 14E and 14G). The attachement site for the distal tibiofibular ligament is again
marked by a swelling on the medial edge of the bone (Figs. 14E–14G).

The fibula is also partly known in *Pl. etalloni* (NMS 8584 and MNS 8731) and *C. jaccardi* (NMS 8713–8718). Based on this incomplete and deformed material, the fibula of *Pl. bigleri*, *Pl. etalloni* and *C. jaccardi* cannot be differentiated.

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#### 1004 Vertebral column

Two disarticulated cervical vertebrae are associated with shells of *Pl. bigleri*. The cervical 1005 vertebra of MJSN TCH005-21 is almost complete, but somewhat deformed (Fig. 15). The 1006 cervical vertebrae associated with MJSN SCR011-30 is more damaged and lacks three 1007 zygapophyses. The precise position of these two vertebrae in the cervical series is unclear. The 1008 centrum is moderately elongated and oval in cross section (slightly flattened dorsoventrally). 1009 There is a robust, but low ventral keel running all of the length of the centrum in MJSN 1010 TCH005-21 (Fig. 15D; not preserved in MJSN SCR011-30). The two known cervical vertebra 1011 1012 are amphicoelous, with oval and slightly concave central articulations (Figs. 15B and 15E). The neural arch is moderately high (especially posteriorly) and the neural spine is reduced to a low 1013 longitudinal ridge (MJSN SCR011-30). The prezygapophyses and postzygapophyses are widely 1014 1015 separated (Figs. 15A and 15D). The articular surface of the prezygapophyses is oriented dorsally

and slightly medially. The anterior margin of the neural arch forms a deep embayment between
the prezygapophyses (Fig. 15A). The articular surface of the postzygapophyses faces ventrally
and slightly laterally. A strong V-shaped ridge occurs on the dorsal surface of the
postzygapophyses and defines a deep triangular fossa between the postzygapophyses (Figs. 15A)
and 15E). The transverse processes is poorly developed and is situated anteriorly along the
centrum (MJSN SCR011-30).

Cervical vertebrae are known in Pl. etalloni (NMS 8584), C. jaccardi (NMS 8713-8718), 1022 and Th. hugii (NMS 8595-8609). The cervical vertebrae of Pl. etalloni and C. jaccardi are 1023 strongly deformed and broken. The few discernable features are consistent with what is known in 1024 *Pl. bigleri*. The cervical vertebrae of *Th. hugii* are much better preserved, even though they 1025 suffered strong lateral pressure. Taking postmortem deformation into account, these cervical 1026 vertebrae are also consistent with what is known in *Pl. bigleri*: moderately long amphicoelous 1027 centrum, robust but low ventral keel, moderately high neural arch (notably posteriorly), and 1028 widely separated zygapophyses. 1029

Thoracic vertebrae can best be observed in the articulated specimens MJSN BSY007-147 1030 and MJSN TCH007-519, as well as in the disarticulated specimen MJSN SCR011-30. The best 1031 1032 preserved centra in these specimens are biconcave, smoothly rounded ventrally and without keel: e.g., 6th thoracic vertebra in MJSN BSY007-147, 7th thoracic vertebra in MJSN TCH007-519, 1033 and the disarticulated thoracic vertebrae (probably 9th and 10th) of MJSN SCR011-30. In 1034 1035 contrast, the best preserved thoracic centra of *Pl. etalloni* are keeled: e.g., 1st thoracic vertebra of NMS 8723, 4th thoracic vertebra of NMS 8731, and 5th thoracic vertebra of MJSN TCH006-1036 1037 574. Bräm (1965) already pointed out the keeled anterior thoracic centra of *Pl. etalloni* and *C.* 

*iaccardi*. However, it remains unclear whether this keel is also present on posterior thoracic 1038 centra of these species, and whether Pl. bigleri also had keeled anterior thoracic centra. 1039 One sacral and two caudal vertebrae of MJSN SCR011-30 are preserved. The sacral 1040 vertebra is short, and has two narrow prezygapophyses and a small keel on the centrum. The 1041 centrum of one caudal vertebra bears a robust ventral keel, similar to the cervical vertebrae. Poor 1042 preservation prevents any conclusion on the type of central articulation in the caudal vertebrae. 1043 A still articulated series of caudal vertebrae is preserved in specimen NMS 8584 referred to 1044 *Pl. etalloni.* As far as observable, their shape is consistent with what is known in *Pl. bigleri*. 1045 Bräm (1965) described their centra as procoelous. However, the state of preservation of these 1046 centra does not allow to confirm Bräm's observation. 1047

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#### 1049 PLESIOCHELYS ETALLONI

Fifteen specimens from Porrentruy are referred to Plesiochelys etalloni (Table 1). They are all 1050 represented by elements of the carapace and most of them also by the plastron. The shells MJSN 1051 BSY007-205, MJSN TCH005-332, and MJSN TCH006-574 are still articulated, the latter being 1052 by far the best preserved specimen (Fig. 16). Non-shell post-cranial material is only poorly 1053 preserved. For example, small remains of the scapula and pelvis are associated with MJSN 1054 BSY003-347, but the poor preservation impedes any anatomical comparisons. 1055 As mentioned above, *Pl. bigleri* and *Pl. etalloni* mostly differ in their cranial anatomy. 1056 1057 However, some characteristics of the shell allow to tell the two species apart. The main characters we used for this study are the thickness of neural and costal bones, and the presence 1058 and development of the epiplastral bulbs. In Pl. bigleri, the epiplastral bulbs are reduced or 1059 1060 absent (Fig. 7C), whereas they are usually well developed in *Pl. etalloni* (Figs. 16D-16E). The

neural and costal bones of *Pl. etalloni* are usually remarkable for their great relative thickness 1061 (Fig. 9D). Specimens of adult size regularly have neurals reaching 15 to 20 mm in thickness. 1062 Similarly sized specimens of *Pl. bigleri* usually have a neural thickness ranging between 11 and 1063 14 mm. However, there is a great deal of variation in both species, and the difference between 1064 the two is less obvious in juvenile specimens. Neural (and costal) thickness alone is therefore not 1065 always sufficient to discriminante between the two species. In order to test whether differences 1066 in neural thickness are significant in the two species, we measured 43 specimens referred to Pl. 1067 *bigleri*, *Pl. etalloni*, or *Plesiochelys* sp. (see below). 1068

The newly discovered shells of *Pl. etalloni* show about the same range of variation as 1069 previously described for this species (Anguetin, Püntener & Billon-Bruyat, 2014). MJSN 1070 TCH006-574 is however remarkable in its extremely reduced fourth pleurals, which are 1071 restricted to the peripheral bones due to the great posterolateral development of the fourth 1072 vertebral scute (Fig. 16B). The fourth pleurals are similarly reduced in some specimens from 1073 Solothurn, but they always occupy at least a small part of the costals (e.g., NMS 8514 and NMS 1074 8517; Anquetin, Püntener & Billon-Bruyat, 2014: figs. 2, 8). In MJSN TCH006-574, the twelfth 1075 pair of marginal scutes is restricted to the pygal, which is a common variation in *Pl. bigleri* (see 1076 above), but is unknown in other specimens referred to Pl. etalloni (Anguetin, Püntener & Billon-1077 Bruyat, 2014). 1078

1079

#### 1080 NEURAL THICKNESS IN PLESIOCHELYS

1081 The neural length and thickness was measured on selected specimens (see Material and Methods;

1082 Fig. 17B; Table 3). The scatter plot of mean length and thickness measurements reveals a

1083 relatively clear separation between *Plesiochelys bigleri* and *Plesiochelys etalloni* (Fig. 17A).

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This separation is mostly due to the proportionally increased neural thickness observed in *Pl*. 1084 etalloni, as confirmed by the Mann-Whitney tests for mean thickness and length/thickness ratio 1085 (p < 0.0001, respectively). Mean neural length however is not significantly different in the two 1086 species (p = 0.1156; Fig. 17D). 1087

The discriminant analysis resulted in a relatively good classification of specimens (93.94%) 1088 1089 of individuals correctly identified; Fig. 17C). Only two specimens (MJSN BSY009-815 and MJSN TCH006-767) are incorrectly classified as *Pl. etalloni*. These two specimens happen to be 1090 the ones that lie the closest to the Pl. etalloni morphospace (Fig. 17A). The discriminant analysis

also provided tentative classifications for the 10 indeterminate specimens (Plesiochelys sp.),

which correspond fairly well with the conclusions we can draw based on the length/thickness 1093

scatter plot (Fig. 17A; Table 4). We consider the classification of only two of these indeterminate 1094

specimens to be dubious. MJSN BSY 009-310 is classified by the discriminant analysis as *Pl*. 1095

etalloni, but this specimen actually lies on the margin of the Pl. bigleri morphospace (Fig. 17A). 1096

SCR010-479 is classified by the discriminant analysis as *Pl. bigleri*, but lies so far from the 1097

known morphospaces of *Pl. bigleri* and *Pl. etalloni* that this classification must be regarded with 1098

caution for the moment. 1099

1100 Neural thickness is therefore a good character to help differentiating between *Pl. bigleri* 

and Pl. etalloni. In the analyzed sample, mean neural thickness ranges from 9.61 to 13.77 mm 1101

(mean = 11.95 mm) in *Pl. bigleri*, and from 12.82 to 17.76 mm (mean = 14.83 mm) in *Pl.* 1102

1103 etalloni (Fig. 17E). This difference between the two species is even more clearly expressed in the

length/thickness ratio, which ranges from 4 to 5.65 (mean = 4.79) in *Pl. bigleri*, and from 3.1 to 1104

1105 3.91 (mean = 3.67) in *Pl. etalloni* (Fig. 17F).

1106

#### DISCUSSION 1107 1108 1109 Alpha taxonomy There is a dual issue with the identification and distinction of Plesiochelys bigleri and 1110 *Plesiochelys etalloni*. First, the two species are so closely related that differences in their shell 1111 1112 and appendicular anatomy are minimal (see above). Second, each of the two species is known by tenths of shells from a single locality and horizon. These extensive collections reveal a great 1113 intraspecific variability in the two species (Anguetin, Püntener & Billon-Bruyat, 2014; this 1114 study). Therefore, differentiating the two species can be challenging. 1115 The holotype (MJSN TCH007-252) and paratype (MJSN TCH006-1451) specimens of Pl. 1116 *bigleri* are of paramount importance to establish the distinction between the two species. The 1117 isolated cranium MJSN TCH006-1451 (Figs. 5 and 6) exhibits a number of characteristics that 1118 clearly set it apart from *Pl. etalloni*: reduced processus trochlearis oticum, reduced posterior 1119 flooring of the cavum acustico-jugulare by the pterygoid, clear prootic-opisthotic contact on the 1120 floor of the fossa temporalis superior, pila prootica not ossified, processus paroccipitalis 1121 extending posterolaterally, anterior foramen nervi abducentis opening ventral and slightly 1122 1123 anteromedial to the base of the processus clinoideus, surface below the dorsum sellae sloping more gently anteriorly, and foramina anterius canalis carotici cerebralis widely separated. The 1124 1125 preservation of the cranium associated with the holotype specimen (Fig. 4) is not as good as that 1126 of the paratype, but it also exhibits most of the above differences with *Pl. etalloni*, in addition to the following: more rounded foramen nervi trigemini, and absence of midline contact of the 1127 1128 exoccipital in the floor of the foramen magnum. Most importantly, the cranium of the holotype is 1129 associated with a near-complete carapace and partial plastron (Fig. 7). The cranium was found

literally within the associated shell during preparation alongside elements of the appendicular 1130 skeleton. There is therefore no doubt regarding the natural state of this association. Interestingly, 1131 the holotype shell is remarkably similar to that of *Pl. etalloni*, except for the much thinner neural 1132 and costal bones and for the absence of epiplastral bulbs. 1133 Among the 80 relatively complete shells studied herein, 41 can be confidently referred to 1134 1135 the new species *Pl. bigleri* based on the reduced neural and costal thickness, the absence or great reduction of epiplastral bulbs, and a generally more quadrangular anterior plastral lobe. Among 1136 the remaining 39 shells, 15 exhibit features that are consistent with an identification as *Pl*. 1137 etalloni, notably the great thickness of neural and costal bones and/or the presence of well-1138 developed epiplastral bulbs. The remaining 24 shells are provisionally identified as *Plesiochelys* 1139 sp. because they lack sufficient diagnostic features. Among the 56 specimens referred either to 1140 *Pl. bigleri* or to *Pl. etalloni*, 33 with well-preserved neural bones were selected for a statistical 1141 analysis of neural thickness. This analysis confirmed that the mean thickness and 1142 length/thickness ratio were statistically different in the two species, with a mean length/thickness 1143 ratio of 4.79 for *Pl. bigleri* and 3.67 for *Pl. etalloni* (see above). The length/thickness ratio of 1144 neural bones (notably from neurals 2 to 5) is therefore an important additional feature to consider 1145 1146 in order to differentiate these two species.

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#### 1148 Is *Plesiochelys bigleri* also present in Solothurn?

1149 The Solothurn turtle assemblage is diversified and slightly younger than the one from Porrentruy.

1150 However, the two localities share a number of species in common: *Plesiochelys etalloni*,

1151 Tropidemys langii, Thalassemys hugii, and Thalassemys bruntrutana (Rütimeyer, 1873; Bräm,

1152 1965; Püntener et al., 2014; Püntener, Anquetin & Billon Bruyat, 2015). Given the similarity

observed between *Pl. bigleri* and *Pl. etalloni* in Porrentruy, the presence of *Pl. bigleri* among 1153 Solothurn specimens referred to Pl. etalloni would certainly not come as a surprise. 1154 Six skulls from Solothurn are referred to Pl. etalloni (NMS 8738, NMS 8739, NMS 8740, 1155 NMS 9145, NMS 40870, and NMS 40871; Gaffney, 1975a; Gaffney, 1976; Anguetin, Püntener 1156 & Billon-Bruyat, 2015 hey all appear to belong to that species, although there may be some 1157 1158 doubts regarding NMS 9145 which is associated with unprepared postcranial material including relatively thin costal bones. Remaining specimens referred to Pl. etalloni are otherwise often 1159 preserved as articulated shells, which complicates observation of neural and costal bones 1160 thickness in some cases and prevents a statistical analysis of neural bone thickness. Most of these 1161 specimens exhibit traits that are compatible with *Pl. etalloni*, notably: relatively thick neurals or 1162 costal bones (e.g., NMS 8461, NMS 8515, NMS 8517, NMS 8732), well-developed epiplastral 1163 bulbs (e.g., NMS 8533, NMS 8693, NMS 9150, NMS 9153, NMS 9173), or both features at the 1164 same time (e.g., NMS 8582). In some specimens, neither of the two main distinguishing features 1165 1166 can be observed due to preservation (e.g., NMS 8727). The attribution of other specimens might be questioned because one of the distinguishing feature is poorly expressed, whereas the other is 1167 impossible to check. For example, NMS 8731 has only moderately expressed epiplastral bulbs, 1168 1169 and the thickness of the neurals and costals is difficult to evaluate (the costals seem to be relatively thin). 1170

For the moment, none of the *Plesiochelys* specimens from Solothurn can be confidently referred to *Pl. bigleri*. However, this material should definitely be re-evaluated in the future in light of the new material from Porrentruy.

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#### 1175 CONCLUSIONS

*Plesiochelys bigleri* is a new plesiochelyid turtle known based on 41 relatively complete, but 1176 mostly disarticulated shells and two crania (one associated with a shell, and another one 1177 isolated). All of this material originates from a series of close by localities west of the small town 1178 of Courtedoux, near Porrentruy, Canton of Jura, Switzerland. Most of the specimens were 1179 collected from a single stratigraphically limited horizon, the Lower Virgula Marls, dated from 1180 1181 the early late Kimmeridgian. A few additional specimens were found in two underlying horizons, the Corbis Limestones and Banné Marls, dated from the late early Kimmeridgian. 1182 The shell morphology of *Pl. bigleri* is remarkably similar to that of *Pl. etalloni*, a species 1183 known based on tenths of shells and several crania from the Kimmeridgian of Switzerland, 1184 France, Germany, and England. These two closely related species however differ in the thickness 1185 of the neural and costal bones of the carapace (a difference that is statistically tested herein), and 1186 the presence and development of the epiplastral bulbs in the plastron. Differentiating the two 1187 species based only on shell morphology can be challenging in some incomplete or juvenile 1188 individuals. The two species co-occur in Porrentruy and 24 shells (30% of the shells referable to 1189 *Plesiochelys*) cannot be identified at the species level as a result of this great similarity. 1190 However, the two species are more easily separated based on cranial morphology. Actually, Pl. 1191 bigleri exhibits cranial features that clearly set it apart from *Pl. etalloni* and other plesiochelyids, 1192 such as: a rounded foramen nervi trigemini, a shallow pterygoid fossa, a reduced processus 1193 trochlearis oticum, the absence of ossification of the pila prootica, the surface below the dorsum 1194 1195 sellae sloping rather gently anteroventrally, and the widely separated foramina anterius canalis carotici cerebralis. 1196

For the moment, *Pl. bigleri* is known only in Porrentruy. However, Solothurn and
Porrentruy share several species of turtles in common (*Pl. etalloni, Tropidemys langii*,

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1199	Thalassemys	hugii, and Thalassemys bruntrutana), and these species are also known in the
1200	Kimmeridgia	n of southern England (Püntener et al., 2014; Püntener, Anquetin & Billon-Bruyat,
1201	2015; Anquet	in & Chapman, 2016). Finding Pl. bigleri in other localities would therefore not
1202	come as a sur	prise.
1203	The abu	indant material from Solothurn and Porrentruy referred to Pl. etalloni and Pl.
1204	<i>bigleri</i> illustra	ates the extent of intraspecific variability in these two species. Although these
1205	results may no	ot be blindly transposable to other groups of turtles, they represent an important
1206	point of comp	parison for other studies on Mesozoic turtle diversity.
1207		
1208	Institutional	Abbreviations
1209	MAJ	Musée d'archéologie du Jura, Lons-le-Saunier, France
1210	MJSN	JURASSICA Museum, Porrentruy, Switzerland
1211	NHMUK	Natural History Museum, London, UK
1212	NMB	Naturhistorisches Museum Basel, Switzerland
1213	NMS	Naturmuseum Solothurn, Switzerland
1214	OUMNH	Oxford University Museum of Natural History, Oxford, UK
1215	PIMUZ	Paläontologisches Institut und Museum, Universität Zürich, Switzerland
1216		
1217	Locality Abb	previations
1218	BSY	Bois de Sylleux, Courtedoux, near Porrentruy, Switzerland
1219	CRT	Crat, Chevenez, near Porrentruy, Switzerland
1220	SCR	Sur Combe Ronde, Courtedoux, near Porrentruy, Switzerland
1221	ТСН	Tchâfoué, Courtedoux, near Porrentruy, Switzerland

1222 VTT Vâ Tche Tchâ, Courtedoux, near Porrentruy, Switzerland

1223

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- 1230

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# Figure 1

Geographical map of the Ajoie Region, Canton Jura, Switzerland.

The five excavation sites Bois de Sylleux (BSY), Crat (CRT), Sur Combe Ronde (SCR), Tchâfouè (TCH), and Vâ Tche Tchâ (VTT) are situated along the Transjurane A16 highway (gray).



# Figure 2

Stratigraphic section of the Reuchenette Formation.

Most specimens were discovered within the Lower *Virgula* Marls (sites of BSY, SCR, and TCH). One specimen comes from dinosaur track-bearing tidal laminites (CRT), and two from the Banné Marls (VTT). Scheme modified after Comment et al. (2015).

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# Figure 3

MJSN TCH006-1420, Plesiochelys bigleri (Kimmeridgian, Porrentruy, Switzerland).

Field photograph of the specimen embedded in the Lower Virgula Marls.



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# Figure 4

MJSN TCH007-252, holotype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

Cranium in dorsal (A, B), ventral (C, D), anterior (E, F), posterior (G, H), right lateral (I, J), and left lateral (K, L) views. Gray areas indicate disarticulated sutural surfaces. Hatchings represent damaged areas. Abbreviations: ap, antrum postoticum; bo, basioccipital; bs, basisphenoid; ccc, canalis caroticus cerebralis; cci, canalis caroticus internus; cs, crista supraoccipitalis; ct, cavum tympani; ds, dorsum sellae; epi, epipterygoid; ex, exoccipital; fce, fossa cartilaginis epipterygoidei; feng, foramen externum nervi glossopharyngei; fna, foramen nervi abducentis; fnh, foramen nervi hypoglossi; fnt, foramen nervi trigemini; fo, fenestra ovalis; fp, fenestra perilymphatica; fst, foramen stapedio-temporale; ica, incisura columellae auris; op, opisthotic; pa, parietal; pi, processus interfenestralis; pcl, processus clinoideus; ppe, processus pterygoideus externus; pr, prootic; pt, pterygoid; ptf, pterygoid fossa; pto, processus trochlearis oticum; qj, quadratojugal; qr, quadrate ridge; qu, quadrate; so, supraoccipital; sq, squamosal; tra, trabecula; un. f., unnamed foramen.

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# Figure 5

MJSN TCH006-1451, paratype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

Cranium in dorsal (A, B), ventral (C, D), posterior (E, F), and right lateral (G, H) views. Hatchings represent damaged areas. Abbreviations: ap, antrum postoticum; bo, basioccipital; bs, basisphenoid; cci, canalis caroticus internus; cm, condylus mandibularis; cs, crista supraoccipitalis; ct, cavum tympani; ds, dorsum sellae; epi, epipterygoid; ex, exoccipital; faccc, foramen anterius canalis carotici cerebralis; faccp, foramen anterius canalis carotici palatinum; feng, foramen externum nervi glossopharyngei; fm, foramen magnum; fna, foramen nervi abducentis; fnh, foramen nervi hypoglossi; fnv, foramen nervi vidiani; fo, fenestra ovalis; fpp, foramen palatinum posterius; fst, foramen stapedio-temporale; ica, incisura columellae auris; ju, jugal; lar, labial ridge; lir, lingual ridge; mx, maxilla; op, opisthotic; pal, palatine; pi, processus interfenestralis; pcl, processus clinoideus; pr, prootic; pt, pterygoid; pto, processus trochlearis oticum; qj, quadratojugal; qr, quadrate ridge; qu, quadrate; so, supraoccipital; sq, squamosal; vo, vomer.
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MJSN TCH006-1451, paratype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

Dorsal view of the dorsum sellae and sella turcica region. Hatchings represent damaged areas. The lattice pattern represents matrix infilling. Abbreviations: bs, basisphenoid; ds, dorsum sellae; epi, epipterygoid; faccc, foramen anterius canalis carotici cerebralis; faccp, foramen anterius canalis carotici palatinum; fna, foramen nervi abducentis; fnv, foramen nervi vidiani; pcl, processus clinoideus; pr, prootic; pt, pterygoid; sc, sulcus cavernosus; st, sella turcica; tra, trabecula.



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MJSN TCH007-252, holotype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

(A, B) carapace; (C, D) plastron. Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci.Abbreviations: co, costal; ent, entoplastron; epi, epiplastron; hypo, hypoplastron; n, neural; sp, supragygal; v, vertebral scale; xi, xiphiplastron; \*, intermediate element (see text).

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## Figure 8

Shells of Plesiochelys bigleri.

Carapace (A, B) and plastron (C, D) of specimen MJSN TCH006-1420 (Kimmeridgian, Porrentruy, Switzerland); carapace (E, F) and plastron (G, H) of specimen MJSN BSY006-307 (Kimmeridgian, Porrentruy, Switzerland); carapace (I, J) and plastron (K, L) of specimen MJSN SCR011-140 (Kimmeridgian, Porrentruy, Switzerland); carapace (M, N) of specimen MJSN TCH005-42 (Kimmeridgian, Porrentruy, Switzerland). Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci. Abbreviations: co, costal; ent, entoplastron; epi, epiplastron; hyo, hyoplastron; hypo, hypoplastron; nu, nuchal; pl, pleural scale; sp, supragygal; v, vertebral scale; xi, xiphiplastron; \*, intermediate element (see text).

С

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vc

hypo

hyo

hypo

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к



М

100 mm



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Neural bones of *Plesiochelys bigleri* and *Plesiochelys etalloni* (Kimmeridgian, Porrentruy, Switzerland).

Neural 4 of specimen MJSN TCH006-1420 (*Plesiochelys bigleri*) in dorsal (A) and lateral left view (B); neural 4 of specimen MJSN BSY006-347 (*Plesiochelys etalloni*) in dorsal (C) and lateral left view (D).



MJSN TCH005-42, Plesiochelys bigleri (Kimmeridgian, Porrentruy, Switzerland).

Left scapula in lateral (A) and medial view (B).



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MJSN TCH007-252, holotype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

Right humerus in dorsal (A), anterior (B), ventral (C), and posterior view (D); left humerus in dorsal (E), anterior (F), ventral (G), and posterior view (H).





MJSN TCH007-252, holotype of *Plesiochelys bigleri* (Kimmeridgian, Porrentruy, Switzerland).

Right radius in dorsal (A), medial (B), ventral (C), and lateral view (D); right ulna in dorsal (E), medial (F), ventral (G), and lateral view (H).

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MJSN BSY006-307, Plesiochelys bigleri (Kimmeridgian, Porrentruy, Switzerland).

Pelvis in right lateral view  $\bigcirc$ 



MJSN SCR010-1279, Plesiochelys bigleri (Kimmeridgian, Porrentruy, Switzerland).

Right femur in dorsal (A), anterior (B), ventral (C), and posterior view (D); left fibula in dorsal (E), medial (F), ventral (G), and lateral view (H).

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MJSN TCH005-21, Plesiochelys bigleri (Kimmeridgian, Porrentruy, Switzerland).

Cervical vertebra in dorsal (A), anterior (B), left lateral (C), ventral (D), posterior (E), and right lateral view (F).



MJSN TCH006-574, Plesiochelys etalloni (Kimmeridgian, Porrentruy, Switzerland).

Carapace in dorsal (A, B) and right lateral view (C); plastron in ventral view (D, E). Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci. Abbreviations: epi, epiplastron; pl, pleural scale; py, pygal.

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# Figure 17

Neural bone thickness in Plesiochelys etalloni and Plesiochelys bigleri.

Statistical analysis of mean neural length, mean neural thickness, and corresponding length/thickness ratio (see text) for 25 specimens of *Plesiochelys bigleri* (green), 8 specimens of *Plesiochelys etalloni* (blue), and 10 indeterminate specimens (*Plesiochelys* sp.; gray), (A) Mean length/thickness scatter-plot (specimen numbers are indicated for indeterminate specimens). (B) Length and thickness measurements on the fifth neural bone of specimen BSY006-307 (scale bar = 20mm). (C) Discriminant histogram. (D–F) Box-and-whisker plots for mean length, mean thickness, and length/thickness ratio, respectively.



#### Table 1(on next page)

The new *Plesiochelys* material from Porrentruy.

42 specimens are attributed to the new species *Plesiochelys bigleri*, 15 specimens to *Plesiochelys etalloni*, and 24 specimens to *Plesiochelys* sp.

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#### Plesiochelys bigleri

MJSN BSY006-307	MJSN SCR011-37	MJSN BSY006-347
MJSN BSY006-326	MJSN SCR011-140	MJSN BSY006-376
MJSN BSY007-147	MJSN SCR011-148	MJSN BSY007-205
MJSN BSY007-257	MJSN SCR011-160	MJSN BSY009-694
MJSN BSY008-206	MJSN SCR011-276	MJSN SCR003-1011
MJSN BSY008-242	MJSN SCR011-413	MJSN SCR008-33
MJSN BSY008-512	MJSN TCH005-16	MJSN SCR010-382
MJSN BSY008-567	MJSN TCH005-21	MJSN SCR011-415
MJSN BSY008-848	MJSN TCH005-42	MJSN TCH005-216
MJSN BSY009-639	MJSN TCH005-464	MJSN TCH005-332
MJSN BSY009-743	MJSN TCH005-819	MJSN TCH005-457
MJSN BSY009-815	MJSN TCH006-145	MJSN TCH006-574
MJSN BSY009-892	MJSN TCH006-767	MJSN TCH007-265
MJSN CRT007-2	MJSN TCH006-1420	MJSN TCH007-505
MJSN SCR010-327	MJSN TCH006-1451	MJSN TCH007-771
MJSN SCR010-342	MJSN TCH007-252	
MJSN SCR010-1009	MJSN TCH007-371	
MJSN SCR010-1047	MJSN TCH007-516	
MJSN SCR010-1196	MJSN TCH007-519	
MJSN SCR010-1279	MJSN VTT006-299	
MJSN SCR011-30	MJSN VTT006-579	

#### Plesiochelys sp.

Plesiochelys etalloni

MJSN BSY003-1
MJSN BSY008-240
MJSN BSY008-484
MJSN BSY008-674
MJSN BSY009-171
MJSN BSY009-310
MJSN BSY009-619
MJSN SCR010-413
MJSN SCR010-450
MJSN SCR010-479
MJSN SCR010-559
MJSN SCR010-560
MJSN SCR010-561
MJSN SCR010-562
MJSN SCR011-111
MJSN SCR011-525
MJSN TCH005-286
MJSN TCH005-817
MJSN TCH006-776
MJSN TCH006-787
MJSN TCH007-62
MJSN TCH007-272
MJSN TCH007-541
MJSN TCH007-580

1

#### Table 2(on next page)

Length and width measurements of the skull in *Plesiochelys bigleri*.

These measurements should be compared with those of other plesiochelyid skulls (Anquetin, Püntener & Billon-Bruyat, 2015: table 1).

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		Total length from	Length from pt-vo/pal	Width at the level of	
	condylus occipitalis to		suture to condylus	the condyli	
	Specimen	tip of the snout (mm)	occipitalis (mm)	mandibularis (mm)	
	TCH007-252	-	38.5ª	60.6	
	TCH006-1451	59.8 <sup>a</sup>	34.4	60.0	
1					

2 <sup>a</sup>Specimen incomplete

#### Table 3(on next page)

Measurements used for the analysis of neural thickness in *Plesiochelys* spp.

Mean neural length, mean neural thickness, and mean length/mean thickness ratio measured for selected specimens referred to *Plesiochelys bigleri*, *Plesiochelys etalloni*, and *Plesiochelys* sp. Measurements are expressed in millimeters. See Table S1 for original measurements. All specimens are housed at the MJSN.

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Specimen	Identification	Mean length	Mean thickness	Ratio
BSY006-307	Pl. bigleri	60.62	12.75	4.75
TCH007-252	Pl. bigleri	56.99	10.15	5.62
SCR011-140	Pl. bigleri	57.48	10.76	5.34
BSY009-815	Pl. bigleri	57.12	13.77	4.15
BSY007-257	Pl. bigleri	45.27	9.61	4.71
SCR011-148	Pl. bigleri	52.89	12.53	4.22
SCR011-413	Pl. bigleri	54.98	11.99	4.59
TCH006-1420	Pl. bigleri	56.09	11.65	4.81
SCR011-276	Pl. bigleri	60.57	13.22	4.58
VTT006-299	Pl. bigleri	61.39	12.17	5.04
TCH005-16	Pl. bigleri	61.07	11.50	5.31
TCH005-464	Pl. bigleri	57.14	11.70	4.89
TCH005-21	Pl. bigleri	63.02	13.16	4.79
SCR011-37	Pl. bigleri	52.90	11.46	4.61
BSY008-206	Pl. bigleri	59.31	12.88	4.60
SCR010-1279	Pl. bigleri	65.20	11.59	5.62
VTT006-579	Pl. bigleri	54.98	11.87	4.63
TCH006-145	Pl. bigleri	52.32	11.34	4.61
BSY009-892	Pl. bigleri	60.16	12.85	4.68
TCH005-819	Pl. bigleri	60.40	12.16	4.97
BSY006-326	Pl. bigleri	54.37	10.79	5.04
SCR010-1009	Pl. bigleri	56.53	11.34	4.99
SCR011-160	Pl. bigleri	58.74	12.34	4.76
SCR010-1196	Pl. bigleri	55.12	12.52	4.40
TCH006-767	Pl. bigleri	50.46	12.62	4.00
BSY009-694	Pl. etalloni	57.77	14.83	3.90
SCR011-415	Pl. etalloni	48.07	13.61	3.53
BSY006-347	Pl. etalloni	56.45	14.70	3.84
BSY007-205	Pl. etalloni	60.21	15.41	3.91
SCR008-33	Pl. etalloni	55.01	17.76	3.10
SCR010-382	Pl. etalloni	49.05	12.82	3.83

TCH007-265	Pl. etalloni	52.57	14.89	3.53
TCH007-505	Pl. etalloni	54.88	14.64	3.75
BSY009-310	Plesiochelys sp.	53.24	13.12	4.06
TCH007-272	Plesiochelys sp.	48.59	11.64	4.18
TCH005-817	Plesiochelys sp.	49.50	14.81	3.34
SCR010-450	Plesiochelys sp.	51.98	11.80	4.40
SCR010-479	Plesiochelys sp.	44.50	11.34	3.92
SCR011-525	Plesiochelys sp.	47.86	12.81	3.74
BSY009-619	Plesiochelys sp.	53.38	15.22	3.51
BSY008-240	Plesiochelys sp.	50.52	10.22	4.95
SCR011-111	Plesiochelys sp.	48.35	13.70	3.53
BSY008-484	Plesiochelys sp.	57.80	14.77	3.91

1

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#### Table 4(on next page)

Classification of indeterminate specimens.

Comparison of the tentative classifications of indeterminate specimens (*Plesiochelys* sp.) based on the length/thickness scatter plot (Fig. 17A) and the discriminant analysis (Fig. 17C). All specimens are housed at the MJSN.

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Specimen	Scatter plot	Discriminant analysis
BSY009-310	Pl. bigleri	Pl. etalloni
ТСН007-272	Pl. bigleri	Pl. bigleri
TCH005-817	Pl. etalloni	Pl. etalloni
SCR010-450	Pl. bigleri	Pl. bigleri
SCR010-479	?	Pl. bigleri
SCR011-525	Pl. etalloni	Pl. etalloni
BSY009-619	Pl. etalloni	Pl. etalloni
BSY008-240	Pl. bigleri	Pl. bigleri
SCR011-111	Pl. etalloni	Pl. etalloni
BSY008-484	Pl. etalloni	Pl. etalloni

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