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The rediscovery and redescription of the holotype of the Late Jurassic turtle *Plesiochelys*
etalloni

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ABSTRACT

Plesiochelyidae are a major component of Late Jurassic shallow marine environments throughout Europe. However, the taxonomy of the plesiochelyid turtles is rather confused. Over the years, many taxa have been synonymized with *Plesiochelys etalloni*, one of the first described species. However, the holotype of *P. etalloni* (and only specimen known from the type locality) was lost for more than 150 years. This specimen has been recently rediscovered in the collections of the Musée d'archéologie du Jura in Lons-le-Saunier, France. For the first time since its original description in 1857, the holotype of *P. etalloni* is redescribed and compared to relevant material. The taxonomical status of this taxon is revised accordingly. Based on the morphology of the newly rediscovered holotype, the species *P. solodurensis*, *P. sanctaeverenae* and *P. langii* are synonymized with *P. etalloni*. Known skull-shell associations for *P. etalloni* are re-evaluated in light of the new morphological information available since the rediscovery of this holotype specimen. Finally, we confirm that *Plesiochelys* is represented by a single species in Solothurn, Switzerland.

INTRODUCTION

Despite numerous historical discoveries dating from as early as the beginning of the nineteenth century (e.g., Cuvier, 1824; Pictet & Humbert, 1857; Meyer, 1860; Pictet, 1860; Wagner, 1861; Maack, 1869; Rüttimeyer, 1873), the diversity of Late Jurassic European turtles still eludes our understanding. Traditionally referred to the families Plesiochelyidae, Thalassemydidae and Eurysternidae, these forms are generally considered to be basal eucryptodires, but their exact relationships with one another and with other turtle groups remain largely unclear and usually vary among authors (e.g., Gaffney & Meylan, 1988; Hirayama, Brinkman & Danilov, 2000; Gaffney et al., 2007; Joyce, 2007; Sterli, 2010; Rabi et al., 2013). A number of reasons may be invoked to explain this situation, but at least two of these are the much needed revision of the rich historical material and the limited number of skull-shell associations. Cranial characters are important for turtle systematics, yet many Late Jurassic turtles from Europe are known only from postcranial material. There are few exceptions however, for which both the skull and the shell are known: notably *Solnhofia parsonsi* Gaffney, 1975b and *Plesiochelys etalloni* (Pictet & Humbert, 1857).

Emys etalloni Pictet & Humbert, 1857 was described based on a single shell found in the French Jura Mountains (see below). A few years later, Rüttimeyer (1873) correctly reassigned this species to his newly created genus *Plesiochelys*. The type species of *Plesiochelys* is *P. solodurensis* Rüttimeyer, 1873, a species typified based on material from the prolific quarries near Solothurn in the Swiss Jura Mountains. Rüttimeyer (1873) and Bräm (1965) afterwards both recognized the presence of *P. solodurensis* and *P. etalloni* in the Solothurn deposits. Although turtle skulls were known in Solothurn since as early as the 1820s (Cuvier, 1824; see Bräm, 1965 for an historical account), they were not fully prepared until the 1970s (Gaffney, 1975a). Based on this material, Gaffney (1975a) concluded that *Emys etalloni* Pictet & Humbert, 1857, *Emys jaccardi* Pictet, 1860, *Styemys lindenensis* Maack, 1869, *Plesiochelys*

65 *solodurensis* Rütimeyer, 1873, *Plesiochelys sanctaeverenae* Rütimeyer, 1873,
66 *Craspedochelys picteti* Rütimeyer, 1873, and *Craspedochelys crassa* Rütimeyer, 1873
67 represented a single species, which should be named *Plesiochelys etalloni* (Pictet & Humbert,
68 1857) in application of the Principle of Priority. The immediate effect was that *P. etalloni* was
69 henceforth included into phylogenetic analyses, which helped to improve our understanding
70 of the systematics and relationships of Late Jurassic and Early Cretaceous turtles from Europe
71 and Asia (e.g., Gaffney & Meylan, 1988; Hirayama, Brinkman & Danilov, 2000; Joyce,
72 2007). However, this relatively inclusive synonymy list was not generally accepted among
73 specialists (including ourselves), at least at the alpha level. For example, several subsequent
74 authors still considered *Craspedochelys* as a distinct form, and *P. etalloni* and *P. solodurensis*
75 as different species (e.g., Antunes, Becquart & de Broin, 1988; Lapparent de Broin, Lange-
76 Badré & Dutrieux, 1996).

77 This extremely confusing situation is in part due to the fact that the holotype of
78 *Plesiochelys etalloni* was considered to be lost since the 1860s and was therefore unavailable
79 notably to Rütimeyer (1873), Bräm (1965), Gaffney (1975a), and Lapparent de Broin, Lange-
80 Badré & Dutrieux (1996). These authors based their conclusions on the original description
81 (Pictet & Humbert, 1857) and on plaster casts of the type specimen, which are available in
82 several European museums, notably in Paris and Geneva. We have been fortunate to locate
83 this historical specimen in the collections of the Musée d'archéologie du Jura in Lons-le-
84 Saunier, France. We have also been able to retrace the history of this specimen as it passed
85 from one owner to the other. This material is redescribed herein and the taxonomic status of
86 *Plesiochelys etalloni* is revised accordingly. Finally, this rediscovery allow us to re-evaluate
87 the known skull-shell associations for *P. etalloni*.

Institutional Abbreviations: **MAJ**, Musée d'archéologie du Jura, Lons-le-Saunier, France;
MH, Naturhistorisches Museum, Basel, Switzerland; **NMS**, Naturmuseum Solothurn,
Switzerland.

HISTORICAL BACKGROUND

Pictet and Humbert (1857) explained that the holotype of *Plesiochelys etalloni* was collected by a local priest in the forest close to the village of Lect, near Moirans-en-Montagne (Jura, France). When they studied the specimen, it was in possession of Joseph Célestin Girod, vicar general of the Saint-Claude diocese (France). Neither Rütimeyer (1873) nor Bräm (1965) gave indication relative to the repository of this specimen. Gaffney (1975a) indicated that H. Bräm told him the specimen was lost. Lapparent de Broin, Lange-Badré & Dutrieux (1996) explained that they actively looked for the type but were unable to locate it, but they figured the plaster cast housed in the Natural History Museum in Geneva, Switzerland. Finally, without further explanation, Lapparent de Broin (2001) stated that the holotype of *P. etalloni* had been located in the Natural History Museum of Besançon, France. After verification, it appears that this information is incorrect.

One of us (SD) rediscovered the original specimen a few years ago in the collections of the Musée d'archéologie du Jura in Lons-le-Saunier, France. Examination leaves no doubt whatsoever on the identity of this specimen (Figs 1 and 2). This specimen (MAJ 2005-11-1) was not always housed at the MAJ: it was donated to the museum by a private owner in 1994. The MAJ also houses a plaster copy of the fossil, which was offered by C-A Etallon, the renowned geologist, on March 30th, 1857. After a careful investigation, we were able to uncover most of the history of the fossil shell before it was finally donated to the MAJ.

The claim that the holotype of *P. etalloni* was housed in the Natural History Museum of Besançon (Lapparent de Broin, 2001) is not entirely incorrect. We have found evidences that

the specimen was actually part of the Besançon Museum collection, if only for a short time. This part of the story can be followed in the *Mémoires de la Société d'émulation du département du Doubs* dated from 1859 and 1860. Bishop Mabile, Vicar Girod's superior, offered the specimen to Mr Thiébaud, a member of the Société d'émulation du Doubs (a French scientific society), who gave it to the Besançon Museum. The exact date is unclear, but it was somewhere between 1857 and 1859. In 1859, Vicar Girod wrote to the French Minister of Public Education and Cults, who turned him down, then to the Rector explaining that he had never agreed for the fossil to be given for free to the Besançon Museum and claimed property on the specimen. The Rector abided and the holotype of *P. etalloni* was sent back to Saint-Claude (Jura, France). Joseph Célestin Girod died in 1863 and the track of the specimen was lost.

The last piece of the puzzle was revealed when Mr and Mrs Lacroix donated the specimen to the MAJ in 1994. After claiming the fossil as his own, the Vicar sold it to a private party, the ancestor of Mr and Mrs Lacroix, in order to finance the renovation of his church. The transaction must have occurred between 1859 and 1863. Until 1994, the holotype of *P. etalloni* remained in this family and was passed from one generation to another (Fig. 1).

SYSTEMATIC PALEONTOLOGY

TESTUDINES Batsch, 1788

EUCRYPTODIRA Gaffney, 1975c

PLESIOCHELYIDAE Rütimeyer, 1873

Plesiochelys Rütimeyer, 1873

Plesiochelys etalloni (Pictet & Humbert, 1857)

Emys Etalloni Pictet & Humbert, 1857 (original description)

Plesiochelys solodurensis Rütimeyer, 1873 (subjective synonymy)

138 *Plesiochelys sanctaeverenae* Rütimeyer, 1873 (subjective synonymy)

139 *Plesiochelys langii* Rütimeyer, 1873 (subjective synonymy)

140 Type material.—MAJ 2005-11-1, a shell missing a large part of the carapace medially.

141 Holotype by monotypy.

142 Type horizon and locality.—"Forêt de Lect" (Lect is a small village) near Moirans-en-

143 Montagne (Department of Jura, France), Late Jurassic. The exact horizon is uncertain, but

144 most outcrops in the vicinity of Lect are either Kimmeridgian or early Tithonian. According

145 to Etallon (1857), the specimen was found in the "calcaires portlandiens". *Gravesia gigas* was

146 also found in these limestones (Etallon, 1857), which led Lapparent de Broin, Lange-Badré &

147 Dutrieux (1996) to conclude that MAJ 2005-11-1 was from the early Tithonian.

148 Illustrations of type.—Pictet & Humbert (1857:plates I-III); Figs 1 and 2.

149 Referred specimens.—See Bräm (1965): specimens referred to *P. etalloni*, *P. solodurensis*,

150 *P. sanctaeverenae* and *P. langii* (except NMS 124). For cranial material, see Gaffney (1975a).

151 Revised diagnosis.—Based on shell only (see Gaffney, 1975a for a diagnosis based on

152 cranial characters): relatively large (up to 550 mm in carapace length) turtle with completely

153 ossified carapace; shell bones relatively thick; carapace oval in outline; wide and shallow

154 nuchal notch; nuchal wide and trapezoidal; additional trapezoidal element often present

155 between the neural series and first suprapygals; three cervical scales; wide vertebral scales,

156 usually extending approximately half the length of the costals; anterior marginal scales very

157 short and not extending onto costals; relatively long plastron (85-90% of carapace length)

158 sutured to the carapace along a long osseous bridge; entoplastron variable in size, usually

159 diamond-shaped with a more or less extended posterior part; hyoplastron and xiphioplastron

160 longer than wide; central plastral fontanelle retained in some adults; short gular and extragular

161 scales; long humeral scale; four inframarginal scales mostly covering the plastral elements.

Remarks.—The synonymy list is intentionally restricted to the *Plesiochelys* species described by Rütimeyer (1873) and later revised by Bräm (1965). The synonymy list proposed by Gaffney (1975a) is more inclusive, but testing it would require an extensive revision of historical material at the European scale, something that was done neither by Gaffney (1975a, 1976) nor any subsequent author (see Discussion).

DESCRIPTION

General Description

The holotype of *Plesiochelys etalloni* (MAJ 2005-11-1) is a large, oval shell with carapace and plastron still articulated (Fig. 2 and Video S1). The specimen may have been slightly flattened during fossilization, but there are no indications of severe deformation. The specimen is fairly complete, although part of the left bridge and central part of the carapace are missing. The part of the carapace that is missing reveals the steinkern, which probably explains why the locals regarded this specimen as the imprint of a human torso (Pictet & Humbert, 1857). The shell is filled with matrix. There are some indications in the right axillary and inguinal notches that some elements of the appendicular skeleton are preserved within the matrix, but as it stands these elements are undetermined.

Carapace

As preserved, the length of the carapace is 471 mm, but most of the pygal is missing (Figs 2A–C and S2). The carapace is evenly oval in outline, except anteriorly where there is a broad, shallow nuchal notch. A large part of the carapace is missing centrally. As a result, the neural series and the medial half of most costals are only visible as imprints on the steinkern.

The nuchal is a wide and trapezoidal element. The nuchal notch is shallow, but it extends laterally on the medial part of the first peripheral. Only the anterior part of the first neural is

187 preserved. This element was apparently longer than wide and rectangular. Neural 2–6 are
 188 preserved as imprints on the steinkern. They are elongate, hexagonal elements with their
 189 shorter sides facing anteriorly. The sixth neural is shorter than the previous elements in the
 190 series. Behind the sixth neural, the imprint of the anterior part of the seventh neural is also
 191 preserved. Posteriorly, the steinkern is covered by the bony carapace, but the sutures in this
 192 area are hardly visible and it is uncertain whether or not there are additional elements to the
 193 neural series. Although it is impossible to be certain, the eighth costals may contact one
 194 another in the midline. Most specimens from Solothurn referred to *P. etalloni* (sensu this
 195 study) have an eighth neural and an additional trapezoidal element of uncertain identity
 196 between the seventh neural and the first suprapygal. This area is however relatively variable
 197 in plesiochelyids and neurals 7 and or 8 may be reduced or lost allowing a medial contact of
 198 costals 7 and/or 8 (Bräm, 1965; JA, unpublished data). There are eight costals. The first
 199 costals is relatively short compared to the following ones. Anteriorly, it contacts the nuchal
 200 and the three first peripherals. Costals 2–4 are wider and longer elements, with costal 3 being
 201 notably wide distally. Costals 5–8 decrease progressively in length and width. There were
 202 certainly 11 peripherals, even if they cannot be clearly all observed on the fossil. The sutures
 203 between peripherals 4, 5 and 6 are not preserved dorsally, but they are visible ventrally.
 204 Posteromedially, the suture between the tenth and eleventh peripherals is also not preserved,
 205 but it must have been there. Peripherals are longer than wide, rectangular elements. Most of
 206 peripheral 11 is missing on both sides. The posteromedial region of the carapace is rather
 207 poorly preserved. There are two large suprapygals. The first suprapygal is a broad element
 208 that contacts the costals 8 anteriorly along a long, anteriorly concave suture, the peripherals
 209 11 (probably) laterally, and the second suprapygal posteriorly along a more or less straight
 210 suture (poorly preserved). The exact outline of the second suprapygal is uncertain, because

most of its sutures with surrounding elements are effaced. Posteriorly, just in front of the broken margin of the carapace, the suture with the pygal is barely discernible.

Three cervical scales are clearly visible on the nuchal. Plesiochelyids have long been thought to be characterized by this character, but its distribution is actually wider. For example, several eurytetrasternids are known to have three cervical scales (Bräm, 1965; Joyce, 2003; Anquetin & Joyce, unpublished data). Scale sulci are clearly apparent on the carapace, but very little can be said about the vertebral scales because a large part of the carapace is missing. The first vertebral scale is a broad element, wider anteriorly than posteriorly. Its lateral margins extend on the first costal and first peripheral, but not on the nuchal. Laterally, the first vertebral scale reaches the lateral part of the first marginal. Nothing can be said about the second and third vertebral scales. The fourth vertebral scale is a broad element extending laterally about two-thirds of the length of the sixth and seventh costals. The outline of the fourth vertebral scale is somewhat unusual. Posterolaterally, its lateral margin extends abruptly onto the tenth peripheral. This unusual shape is symmetrical, but, based on our experience of the intraspecific variability in plesiochelyids, we grant it no systematic value. The fifth vertebral scale is a wide, pentagonal element extending onto costals 8, suprapygal 1 and 2, and peripherals 10 and 11. There are four pleural scales. The outlines of pleurals 1–3 are uncertain. The first pleural scale contacts marginals 1–4 and maybe also the fifth marginal scale. The first pleural scale is slightly shorter than pleurals 2 and 3. The second pleural scale reaches the seventh marginal scale posteriorly on the sixth peripheral. The fourth pleural scale is a reduced element covering only a small portion of the sixth and seventh costals and the medial part of the ninth and tenth peripherals. Marginals are only partly preserved. Marginals 1–6 are still partly visible on the right anterolateral part of the carapace. When preserved, the pleuro-marginal sulci are always on the peripherals and never extend onto the costals. It

should also be noted that the last marginal scales (probably the twelfth pair, although it is impossible to be sure) extend anteriorly onto the second suprapygal.

Plastron

The plastron of MAJ 2005-11-1 is mostly complete (Figs 2D–F and S3). The anterior margin of the left epiplastron, the bridge area on the left hand side, and posterior tip of the xiphiplastra are broken. The matrix preserved the imprints of the broken parts of the bridge and xiphiplastra. The total length of the plastron is 431 mm, measuring from the epiplastra anteriorly to the imprint of the xiphiplastra posteriorly. As such, the plastron represents 91.5% of the length of the carapace (the true ratio would be slightly lower if the pygal had been preserved). The plastron is strongly sutured to the carapace. The bridge extends from the posterior part of the third peripheral to the anterior part of the eighth. The axillary and inguinal notches are deep. A small central fontanelle is present between the hyo- and hypoplastra. The anterior lobe is shorter than the posterior lobe, which is itself shorter than the bridge measured between the axillary and inguinal notches. The anterior lobe is trapezoidal in outline with a nearly straight anterior margin. The posterior lobe has a triangular outline with a slightly rounded posterior tip. The central part of the plastron is slightly concave. This concavity may have been natural.

None of the two epiplastra is complete. The left one is missing its anterior margin, whereas the lateral part of the right one is partly covered by matrix. As preserved, the epiplastra are relatively short, wider than long elements. They contact one another medially, the hyoplastra posteriorly, and the entoplastron medially. The epi-hyoplastron suture is straight and transverse. The entoplastron is a diamond-shaped, slightly longer than wide element with its posterior faces slightly more elongated than the anterior. The hyoplastron is a large, longer than wide element. Posteromedially, the hyoplastra form the anterior third of the central

260 plastral fontanelle. The hyo-hypoplastral suture is slightly concave anteriorly, more so
261 medially. The hypoplastra are shorter than the hyoplastra. They form the remaining two-thirds
262 of the central fontanelle. The suture between the hypoplastron and the xiphiplastron is mostly
263 straight and transverse medially. Laterally, it bends suddenly backwards, as it is so often
264 seen in turtles. The xiphiplastra are triangular, longer than wide elements with a slight
265 broadening where the femoro-anal sulcus meets their lateral margin, as correctly noted by
266 Pictet & Humbert (1857). The midline contacts between the different plastral elements are
267 partly disarticulated (Fig. S3), so that the exact position of the sutures is difficult to assess.
268 Probably as a result, Pictet & Humbert (1857) erroneously described and depicted a very
269 small fontanelle between the hypo- and xiphiplastra. Direct examination of the specimen and
270 observation of the 3D surface reconstruction (Video S1 and Fig. S3) both suggest that there is
271 no such fontanelle in MAJ 2005-11-1.

272 Gular and extragular scales are relatively small. The gular scales extend only a little onto
273 the anteromedial part of the entoplastron. The extragular scales are restricted to the epiplastra.
274 The long humeral scales cover the rest of the anterior plastral lobe. The pectoral scale is
275 nearly as long as the abdominal scale on the midline, but both are shorter than the humeral
276 scale. The abdominal-femoral sulcus is oblique and extends from the inguinal notch to the
277 posterior third of the central plastral fontanelle. The femoral is the longest scale of the
278 plastron. The femoral-anal sulcus is deeply concave posteriorly in its medial part. The anal
279 scales are restricted to the xiphiplastra. The medial sulcus between paired scales is unusually
280 irregular. The median sulcus diverges strongly from the midline between the humeral and
281 pectoral scales, being notably sinusoidal between the latter. The median sulcus is more poorly
282 preserved between the femoral and anal scales, but observation of the 3D surface
283 reconstruction (Video S1) suggests that it might also have been slightly sinusoidal, at least in
284 the posterior part of the femoral scales. The bridge area is covered by four inframarginal

scales increasing in length posteriorly. The two first and the last are restricted to the hyoplastron and hypoplastron, respectively. The third inframarginal scale covers the hyoplastron anteriorly, the hypoplastron posteriorly, and a small portion of the fifth peripheral laterally.

DISCUSSION

Skull-Shell Associations

Despite a profusion of material collected from the Late Jurassic of Europe, relatively few species are known from both skull and shell material. European lithographic limestone localities (especially Solnhofen, Kelheim, and Cerin) have produced a fair number of relatively complete, articulated specimens with shell, skull, and various elements of the skeleton (e.g., Meyer, 1860), but the cranial material is always badly crushed and very difficult to interpret. Hence, the skull is 'known' in species such as *Eurysternum wagleri*, *Idiochelys fitzingeri*, and *Palaeomedusa testa* (e.g., Jourdan, 1862; Joyce, 2003; Anquetin & Joyce, unpublished data), but only scarce information can be gathered from these examples.

Among European Late Jurassic turtles, only *Solnhofia parsonsi* and *Plesiochelys etalloni* are sufficiently known from both skull and shell material. Additional skull and associated fragmentary shell remains were described by Rieppel (1980) and assigned to *Thalassemys moseri* Bräm, 1965, but the validity of both this taxon and this referral was questioned by subsequent authors (e.g., Gaffney & Meylan, 1988; Lapparent de Broin, Lange-Badré & Dutrieux, 1996). This material should therefore be revised. *Solnhofia parsonsi* was described by Gaffney (1975b) based on two isolated skulls, one from the Solnhofen region (Germany), one from Solothurn (Switzerland). Later, Joyce (2000) described a nearly complete skeleton that can be confidently referred to *S. parsonsi*.

309 Skulls of *P. etalloni* are known since the early nineteenth century (e.g., Cuvier, 1824;
 310 Rüttimeyer, 1873; Bräm, 1965), although they were not necessarily assigned to this species in
 311 those times. The Solothurn Turtle Limestone has produced four *Plesiochelys* skulls, which
 312 Gaffney (1975a) prepared and identified as belonging to a single species. Among these four
 313 skulls, only one (NMS 594) is associated with significant shell material (few disarticulated
 314 costals and peripherals and partial posterior half of a plastron). Bräm (1965) identified this
 315 specimen as *P. etalloni* based on the probable presence of a central plastral fontanelle.
 316 However, this material is too fragmentary to allow a definitive specific identification. Only
 317 one other skull-shell association exists for *P. etalloni*. It is a specimen (MH 435) that was
 318 found in the Kimmeridgian near Glovelier (Canton of Jura, Switzerland). Bräm (1965)
 319 referred this material to *P. etalloni* without further description and depicted the skull and a
 320 humerus (ibid.:plate 4, figs 1–4). The skull, one of the best for *P. etalloni*, was prepared by
 321 Gaffney (1975a), who followed the identification of Bräm (1965). Gaffney (1975a:7)
 322 examined the associated, incompletely prepared shell material and concluded that "the shell
 323 features as determinable at this time are consistent with [his] concept of *Plesiochelys*
 324 *etalloni*". Because Gaffney's (1975a) concept of *P. etalloni* is inclusive and not necessarily
 325 accepted among fossil turtle specialists, it was important to reassess the shell material of MH
 326 435 and compare it with the newly rediscovered holotype specimen of *P. etalloni*.

327 If the skull of MH 435 has been extensively studied (Gaffney, 1975a, 1976; Sterli et al.,
 328 2010; Carabajal et al., 2013), the associated shell material has never been described or
 329 illustrated. This material (Fig. 3) consists of the anterior half of a shell with carapace and
 330 plastron still in articulation. Everything posterior to the fifth costal on the carapace and
 331 inguinal notch on the plastron is missing. Both the carapace and the plastron are greatly
 332 fractured. Many fragments of the costals are missing. The carapace is oval in outline with a
 333 broad, shallow nuchal notch (Fig. 3A–B). The nuchal is a wide and trapezoidal element. The

first neural is rectangular, whereas neurals 2–4 are hexagonal with their shorter sides anteriorly. Cervical scale sulci are not preserved. The first vertebral scale is a broad, trapezoidal element that extends laterally onto the first peripheral and contacts the lateral border of the first marginal scale. The second and third vertebral scales are wide and hexagonal. Their sulci are moderately sinuous, as it is common in Solothurn specimens referred to *P. etalloni* (Bräm, 1965). The anterior margin of the anterior lobe of the plastron is rounded (Fig. 3C–D). The epiplastron is separated from the hyoplastron by a straight, transverse suture. The hyoplastron is longer than wide. There is an oval central fontanelle between the hyo- and hypoplastron. The hyo-hyoplastral suture is relatively straight and slightly oblique defining a small concavity toward the anterior. The bridge is long and osseous. It extends from the posterior half of the third peripheral to the anterior part of the eighth peripheral. The scale arrangement on the plastron is similar to that of MAJ 2005-11-1. The median sulcus between the humeral and pectoral scales diverges strongly from the midline, although it is not sinusoidal as in the holotype of *P. etalloni*. There are four inframarginal scales increasing in length posteriorly. Based on this description, MH 435 can be confidently referred to *P. etalloni* (sensu this study, not Gaffney, 1975a). This confirms the importance of this specimen, especially for phylogenetic reconstructions.

Alpha Taxonomy

As mentioned above, comparisons for the present study are restricted to the *Plesiochelys* species described by Rütimeyer (1873) and later revised by Bräm (1965), i.e. forms first described from the Late Jurassic of the Swiss and French Jura Mountains. Many specimens from the Late Jurassic of France, Germany, England, Spain and Portugal have afterwards been either referred to *P. etalloni* and *P. solodurensis* or assigned to new species, but these need to be revised thoroughly. Kuhn (1964) listed 22 species of *Plesiochelys* typified based

on European material. It is far beyond the scope of the present study to revise the taxonomy of the genus *Plesiochelys*.

Another issue is the relatively inclusive synonymy list proposed by Gaffney (1975a), who synonymized the following species with *P. etalloni*: *Emys jaccardi*, *Stylemys lindenensis*, *P. solodurensis*, *P. sanctaeverenae*, *Craspedochelys picteti*, and *C. crassa*. *Stylemys lindenensis* is a form from the Late Jurassic of Hannover, Germany, and, along with many other specimens from the same region, it has never been properly revised since Oertel (1924). All other species but *E. jaccardi* were described based on material from Solothurn, Switzerland. *Emys jaccardi* was referred to *Plesiochelys* by Rüttimeyer (1873) and Bräm (1965). In contrast, Antunes, Becquart & de Broin (1988) and Lapparent de Broin, Lange-Badré & Dutrieux (1996) referred this species to the genus *Craspedochelys* Rüttimeyer, 1873, which they distinguished from *Plesiochelys* by a shell as wide as long and a shortened plastron. Gaffney (1975a) argued that variation in shell shape, especially relative width (as used to differentiate *E. jaccardi* and *C. picteti* from *P. etalloni*), was probably the result of postmortem deformation and should not be considered for systematic purposes. The objective of the present paper is not to settle this argument. The fact is that Bräm (1965) is the last author to have thoroughly reassessed the shell morphology of these forms. Gaffney (1975a) focussed essentially on skull description and did not describe shell morphology in detail. Lapparent de Broin, Lange-Badré & Dutrieux (1996) studied some of the Solothurn material, but they did not clearly formalized their views, instead proposing a general discussion as part of the description of new material from France. In contrast to these authors, we have thoroughly revised the Solothurn material. Our conclusions, which concern several additional species besides *P. etalloni*, will be presented elsewhere. For the purpose of the present paper, we restrict our comparisons to *P. solodurensis*, *P. sanctaeverenae* and *P. langii*.

According to Rütimeyer (1873) and Bräm (1965), both *P. etalloni* and *P. solodurensis* are present in Solothurn, the type locality of *P. solodurensis*. However, Bräm (1965) himself admitted that differentiating the two species was not easy. *Plesiochelys etalloni* was supposed to produce slightly larger individuals than *P. solodurensis* and to retain a small central plastral fontanelle in the adults (Bräm, 1965). The proposed difference in size is minor (about 10%) and is not interpreted as being significative. We have scrutinized all fairly complete specimens from Solothurn referred to both *P. etalloni* and *P. solodurensis*, representing about 30 individuals. We have extensively looked for additional characters that would confirm the presence of two species (one with a central plastral fontanelle and one without), but have found none. For example, a close comparison between MAJ 2005-11-1 (holotype of *P. etalloni*) and NMS 59 (lectotype of *P. solodurensis*) reveals only little differences: the shape of the posterolateral sulcus of the fourth vertebral (probably anomalous in MAJ 2005-11-1); the very minute extension of the fourth marginal onto costal 2 in NMS 59; the central plastral fontanelle in MAJ 2005-11-1; and the extension of the anal scale onto the hypoplastron in NMS 59. Anomalous scale shape is relatively common among Solothurn turtles, especially for vertebral scales. Similarly, both the extension of the fourth marginal onto costals and the extension of the anal scale onto the hypoplastron, characters that are otherwise diagnostic for Xinjiangchelyidae (e.g., Tong et al., 2012; Rabi et al., 2013; Pérez-García, Gasulla & Ortega, in press), are variable in *P. etalloni*. Hence, the retention of a central plastral fontanelle in adults is interpreted as an intraspecific variation of *P. etalloni*, and *P. solodurensis* is considered a subjective junior synonym of this species.

Bräm (1965) found no significant difference between NMS 123 and NMS 126, two carapaces referred to *P. langii*, and NMS 59, and therefore synonymized *P. langii* with *P. solodurensis*. We agree and similarly find no significant difference between these specimens and MAJ 2005-11-1. Consequently, *P. langii* is synonymized with *P. etalloni*. *Plesiochelys*

sanctaeverenae was defined by Rütimeyer (1873) mainly based on NMS 118, a large, incomplete carapace. Bräm (1965) designated this specimen as the lectotype and considered this species as valid based on its larger size (carapace length = 565 mm) and elongate outline. However, observable characteristics do not allow to differentiate NMS 118 from others specimens we refer here to *P. etalloni*, especially neither from MAJ 2005-11-1 nor NMS 59. Concerning the outline of this specimen, Bräm (1965) was probably misled by the fact that the lateral parts of the carapace are largely missing. Consequently, *P. sanctaeverenae* is also considered a subjective synonym of *P. etalloni*.

From the above, we recognize only one species of *Plesiochelys* in Solothurn: *Plesiochelys etalloni*. Although this conclusion may appear superficially similar to that of Gaffney (1975a), we reached it through an extensive re-evaluation of the Solothurn material and a redescription of the type material of *P. etalloni*, which was unavailable for these past 150 years. Since Gaffney (1975a, 1976), we have an excellent knowledge of the cranial morphology of *P. etalloni*. Thanks to the present study, we now have a better understanding of the shell morphology and intraspecific variability of this species.

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FIGURE CAPTIONS

FIGURE 1—MAJ 2005-11-1, holotype of *Plesiochelys etalloni* (Pictet & Humbert, 1857). Specimen mounted on a wooden socle with the old label "*Emys Etalloni*, (Pictet et Humbert) - Portland - Moirans (Jura)". (A) carapace; (B) plastron. Note that the specimen is upside down.

FIGURE 2—MAJ 2005-11-1, holotype of *Plesiochelys etalloni* (Pictet & Humbert, 1857). (A) photograph of the carapace; (B) interpretative drawing of the carapace; (C) 3D surface reconstruction of the carapace; (D) photograph of the plastron; (E) interpretative drawing of the plastron; (F) 3D surface reconstruction of the plastron. Bones are white; striped lines indicate internal bone layers; green solid lines indicate scale sulci; matrix is gray.

Abbreviations: *ab*, abdominal scale; *an*, anal scale; *ce*, cervical scale; *co*, costal; *eg*, extragular scale; *epi*, epiplastron; *ento*, entoplastron; *fem*, femoral scale; *gu*, gular scale; *hyo*, hyoplastron; *hypo*, hypoplastron; *hum*, humeral scale; *m*, marginal scale; *n*, neural; *nu*,

532 nuchal; p, peripheral; *pect*, pectoral scale; *pl*; pleural scale; py, pygal; sp, suprapygal; v,
533 vertebral scale; xi, xiphiplastron.

534

535 FIGURE 3—MH 435, *Plesiochelys etalloni* (Pictet & Humbert, 1857). (A) photograph of the
536 carapace; (B) interpretative drawing of the carapace; (C) photograph of the plastron; (D)
537 interpretative drawing of the plastron. Bones are white; stripped lines indicate internal bone
538 layers; green solid lines indicate scale sulci; dotted areas indicate reconstructed parts; matrix
539 is gray. Abbreviations: *ab*, abdominal scale; *co*, costal; *eg*, extragular scale; *epi*, epiplastron;
540 *ento*, entoplastron; *fem*, femoral scale; *gu*, gular scale; *hyo*, hyoplastron; *hypo*, hypoplastron;
541 *hum*, humeral scale; *n*, neural; *nu*, nuchal; *p*, peripheral; *pect*, pectoral scale; *pl*; pleural scale;
542 *v*, vertebral scale.

543

544 FIGURE 1

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