*Thalassemys bruntrutana* n. sp., a new coastal marine turtle from the Late Jurassic of Porrentruy (Switzerland), and the paleobiogeography of the Thalassemydidae

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**Background.** The Swiss Jura Mountains are a key region for Late Jurassic eucryptodiran turtles. Already in the mid 19th century, the Solothurn Turtle Limestone (Solothurn, NW Switzerland) yielded a great amount of Kimmeridgian turtles that are traditionally referred to Plesiochelyidae, Thalassemydidae, and Eurysternidae. In the past few years, fossils of these coastal marine turtles were also abundantly discovered in the Kimmeridgian of the Porrentruy region (NW Switzerland). These findings include numerous sub-complete shells, out of which we present two new specimens of *Thalassemys* (Thalassemydidae) in this study. **Methods.** We compare the new material from Porrentruy to the type species *Th. hugii*, which is based on a well preserved specimen from the Solothurn Turtle Limestone (Solothurn, Switzerland). In order to improve our understanding of the paleogeographic distribution of *Thalassemys*, anatomical comparisons are extended to *Thalassemys* remains from other European countries, notably Germany and England. **Results.** While one of the two *Thalassemys* specimens from Porrentruy can be attributed to *Th. hugii*, the other specimen represents a new species, *Th. bruntrutana* n. sp. It differs from *Th. hugii* by several features: more elongated nuchal that strongly thickens anterolaterally; wider vertebral scales; proportionally longer plastron; broader and less inclined xiphiplastron; wider angle between scapular process and acromion process. Our results show that *Th. hugii* and *Th. bruntrutana* also occur simultaneously in the Kimmeridgian of Solothurn as well as in the Kimmeridgian of England (Kimmeridge Clay). This study is an important step towards a better understanding of the paleobiogeographic distribution of Late Jurassic turtles in Europe.
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INTRODUCTION

*Thalassemys* Rütimeyer, 1873 is a coastal marine turtle from the Late Jurassic of Western Europe (Bräm, 1965; Lapparent de Broin, 2001). It is the only currently recognized representative of the family Thalassemydidae Zittel, 1889, a group that is potentially related to the Plesiochelyidae Baur, 1888 and the Eurysternidae Dollo, 1886 (Joyce, 2007). However, the exact relationships and systematics of these three groups are rather confused (e.g., Broin, 1994; Lapparent de Broin, Lange-Badré & Dutrieux, 1996; Joyce, 2003; Joyce, 2007; Anquetin & Joyce, 2014).

Rütimeyer (1859a; 1859b) first mentioned the name *Thalassemys* during a conference of the *Schweizerische Naturforschende Gesellschaft* at the University of Bern (Maack, 1869), but the name only became available in 1873 when he described and figured two species of *Thalassemys* from the Kimmeridgian of Solothurn, Canton of Solothurn, Switzerland: *Th. hugii* Rütimeyer, 1873 and *Th. gresslyi* Rütimeyer, 1873. The type species *Th. hugii* is based on a relatively flat shell with associated postcranial remains (NMS 8595–8609; formerly NMS 1), which is the largest turtle in Solothurn (preserved carapace length of 630 mm). Since Rütimeyer (1873), the fossil turtles of the so-called Solothurn Turtle Limestone have undergone two major revisions (Bräm, 1965; Anquetin, Püntener & Billon-Bruyat, 2014), both with important impacts on the taxonomy of *Thalassemys*. Bräm (1965) synonymized *Th. gresslyi* with *Th. hugii* and erected a new species, *Th. moseri* Bräm, 1965. Revealing the presence of lateral plastral fontanelles in *Thalassemys*, Anquetin, Püntener & Billon-Bruyat (2014) synonymized *Eurysternum ignorantum* Bräm, 1965 with *Th. hugii* and excluded ‘*Th.* moseri’ from *Thalassemys*.

There is some discussion regarding the validity and taxonomy of ‘*Th.* moseri’ in the literature (Lapparent de Broin, Lange-Badré & Dutrieux, 1996; Anquetin, Püntener & Billon-Bruyat,
but all authors agree that this taxon should be excluded from *Thalassemys*. This species was described by Bräm (1965) based on two shells from the Kimmeridgian of Solothurn, Switzerland. A specimen from the Tithonian of western France consisting of an association between a skull and a partial shell was later referred to this taxon (Rieppel, 1980). Without revising the material, Lapparent de Broin, Lange-Badré & Dutrieux (1996) stated that 'Th.' *moseri* was invalid since they referred the two specimens from Solothurn (type material) to juvenile forms of *Plesiochelys solodurensis* Rütimeyer, 1873, a plesiochelyid from the same locality. However, they considered the material described by Rieppel (1980) as a distinct and indeterminate species of *Plesiochelys* Rütimeyer, 1873. In 2014, Anquetin, Püntener & Billon-Bruyat (2014) revised the type material from Solothurn and concluded that 'Th.' *moseri* was indeed a valid taxon, distinct from *Plesiochelys solodurensis*. In the meantime, Pérez-García (2015) built upon the conclusions of Lapparent de Broin, Lange-Badré & Dutrieux (1996) and referred the material described by Rieppel (1980) to a new taxon, *Jurassichelon oleronensis*, without reassessing the validity of 'Th.' *moseri*. It should be noted that the latter author failed to see both the material from Solothurn and the specimen described by Rieppel (1980). Sorting out this situation would require a complete revision of the material, something that has not been done since 1980. It should notably be determined whether the material described by Rieppel (1980) and the material from Solothurn belong to the same species. This revision is ongoing, and the present paper on *Thalassemys* is certainly not the place to discuss further the systematics of this unrelated taxon.

Several other species have been attributed to *Thalassemys* in the late 18th and early 19th centuries, including *Thalassemys marina* Fraas, 1903 (Tithonian of Schnaitheim, Baden-Württemberg, Germany), *Thalassemys heusseri* Oertel, 1924 (Tithonian of Holzen im Hils,
Lower Saxony, Germany), and Thalassemys ruetimeyeri Lydekker, 1889 (Berriasian of Dorset, England). Because the type material is lost and the original description is insufficient, Th. heusseri must be considered a nomen dubium. Thalassemys ruetimeyeri has been recognized as a junior synonym of the pleurosternid Dorsetochelys typocardium (Seeley, 1869) (Milner, 2004; Pérez-García, 2014). Thalassemys marina has long been considered to represent a eurysternid (see Discussion), but Anquetin, Püntener & Billon-Bruyat (2014) recently challenged this conclusion and confirmed the validity of this taxon. As a result, Th. hugii and Th. marina are currently considered as the only two valid thalassemydids. Undetermined Thalassemys remains have been recently reported from the Kimmeridgian near Oker, Lower Saxony, Germany (Marinheiro & Mateus, 2011), and from the Kimmeridgian of the Isle of Purbeck, Dorset, southern England (Pérez-García, 2015).

In the present study we describe two new specimens of Thalassemys (MJSN SCR011-87 and MJSN BSY008-905) from the upper Kimmeridgian of Porrentruy, Canton of Jura, Switzerland. They were recently discovered within the scope of the Paleontology A16 project, which aims to rescue the paleontological material found during the construction of the A16 Transjurane highway. The excavations resulted in a rich and diverse vertebrate fossil collection from the Kimmeridgian, notably including extensive dinosaur trackways (Marty & Hug, 2003; Marty et al., 2007; Marty, 2008; Marty & Billon-Bruyat, 2009) and numerous coastal marine turtles (Billon-Bruyat, 2005; Püntener et al., 2014; Anquetin, Püntener & Billon-Bruyat, 2015). MJSN SCR011-87, an articulated, sub-complete shell with an associated scapula, is referred to a new species, Thalassemys bruntrutana n. sp. In contrast, MJSN BSY008-905, which consists of disarticulated shell elements and a partial scapula, is referred to the type species Th. hugii.
Furthermore, we discuss the implications of the new material from Porrentruy for the taxonomy and paleobiogeography of *Thalassemys*.

**GEOLOGICAL SETTING**

The new *Thalassemys* specimens were collected near the village of Courtedoux, along the A16 Transjurane highway in the Ajoie Region, Canton of Jura, NW Switzerland (Fig. 1). MJSN SCR011-87 was discovered in Sur Combe Ronde (SCR) in 2011, and MJSN BSY008-905 in Bois de Sylleux (BSY) in 2008. Both specimens come from the Lower *Virgula* Marls (Reuchenette Formation, Chevenez Member) that are dated from the Eudoxus ammonite zone (Comment et al., in press) (Fig. 2). The Lower *Virgula* Marls are slightly older than the Solothurn Turtle Limestone, which forms the uppermost member of the Reuchenette Formation and is dated from the Autissiodorensis ammonite zone (Meyer, 1994; Comment, Ayer & Becker, 2011).

MJSN SCR011-87 was embedded in a hardground within the Lower *Virgula* Marls (Fig. 2). The reddish and strongly encrusted (Ostreidae, Serpulidae) hardground is dated from the Orthocera ammonite horizon (Comment et al., in press). Its invertebrate fauna includes brachiopods (*Sellithyris*) and benthic bivalves (*Ceratomya, Pholadomya*), the latter being responsible for the hardgrounds undulated surface. Apart from this sub-complete shell, vertebrates are limited to isolated material (chondrichthyans, osteichthyans, turtles, crocodilians) and two well preserved crocodilian skeletons (*Metriorhynchus* sp. and *Steneosaurus* cf. *bouchardi*; Schaefer, 2012; Schaefer & Billon-Bruyat, 2014).

MJSN BSY008-905 comes from a marly interval of the Lower *Virgula* Marls that is slightly younger (Hibridus ammonite horizon) than the aforementioned hardground (Koppka, 2015;
Comment et al., in press) (Fig. 2). This about 1 m-thick brown marl 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 (Billon-Bruyat, 2005; Marty & Billon-Bruyat, 2009; Philippe et al., 2010; Koppka, 2015).

NOMENCLATURAL ACTS

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SYSTEMATIC PALEONTOLOGY

TESTUDINES Batsch, 1788

PANCRIPTODIRA Joyce, Parham & Gauthier, 2004

THALASSEMYDIDAE Zittel, 1889
Remark. Thalassemydidae is currently a monogeneric family. Since 2014, we are engaged in a
global revision of European Late Jurassic coastal marine turtles (this represents more than 60
named species in the literature), which are traditionally referred to three families:
Plesiochelyidae, Eurysternidae and Thalassemydidae. The characteristics and potential
relationships of these three families are still obscure. Early on, we decided to take a conservative
path and keep on using these three family names in their traditional definition until we have
gathered enough knowledge about their phylogenetic relationships (Anquetin, Püntener &
Billon-Bruyat, 2014). The purpose was to avoid havocking supra-generic classification at each
new paper, especially without phylogenetic justification.

*Thalassemys* Rütimeyer, 1873

_Type species._ *Thalassemys hugii* Rütimeyer, 1873

_Included valid species._ *Thalassemys hugii* Rütimeyer, 1873; *Thalassemys marina* Fraas, 1903;

*Thalassemys bruntrutana* n. sp.

_Occurrence._ Kimmeridgian of Switzerland (Rütimeyer, 1873; Bräm, 1965; this study), Germany
(Marinheiro & Mateus, 2011), and England (Pérez-García, 2015; Pérez-García, in press);
Tithonian of Germany (Fraas, 1903).

_Revised diagnosis._ Type and only genus of Thalassemydidae. Medium to large sized turtle
(estimated carapace length up to 700 mm); relatively flat carapace (as opposed to the more
domed plesiochelyid carapace); presence of clearly visible linear striations perpendicular to most
sutures on the carapace and plastron. Differing from *Plesiochelys*, *Craspedochoelys* Rütimeyer,
1873, and *Tropidemys* Rütimeyer, 1873 in: presence of small costo-peripheral fontanelles in the
adult; presence of lateral plastral fontanelles; non-sutural connection of epi- and entoplastron;
presence of a small fontanelle between the xhiphilastra; wider angle between scapular process...
and acromion process (only known in *Plesiochelys*). Differing from *Idiochelys* Meyer, 1839a, *Eurysternum* Meyer, 1839b, and *Solnhofia* Gaffney, 1975 in: larger size; narrower vertebral scales; osseous bridge; complete series of neurals (incomplete in *Idiochelys*); central plastral fontanelle present (absent in *Idiochelys*) and longer than wide (opposite in *Eurysternum*); small xiphiplastral fontanelle present (absent in *Idiochelys* and *Solnhofia*).

**Remarks.** *Thalassemys* differs from other Late Jurassic turtles from Europe by a combination of several features. A striking character is the presence of distinct linear striations perpendicular to sutures (Anquetin, Püntener & Billon-Bruyat, 2014). This somewhat recalls the condition known in the Early Cretaceous *Pleurosternon bullockii* Owen, 1842 (see Milner, 2004), but these striations are more pronounced in *Thalassemys*. In contrast, they are absent (or only very weakly expressed) in plesiochelyids and eurysternids. The shape of vertebral scales is also characteristic of *Thalassemys*. The anterolateral sides of vertebrae 2–4 are slightly concave, whereas the posterolateral sides are either slightly convex or sub-straight. The plastral anatomy of *Thalassemys* clearly differentiates it from plesiochelyids and eurysternids. In contrast to plesiochelyids, lateral plastral fontanelles occur in *Thalassemys* and there is no sutural connection of the epiplastra and entoplastron with the hyoplastra. In contrast to eurysternids, the bridge of *Thalassemys* is osseous. As previously proposed by Bräm (1965), the angle between the scapular and acromion processes of the scapula may also be a distinguishing feature of *Thalassemys*. This angle is more open in *Thalassemys* (113–130°) than in *Plesiochelys etalloni* (Pictet & Humbert, 1857) (103–105°), but a broader survey of Late Jurassic coastal marine turtles is needed to definitely conclude on this character variation (Table 1).

Anquetin, Püntener & Billon-Bruyat (2014) suggested that a strong anterior widening of the first neural was diagnostic for *Thalassemys*. This feature is present in most specimens referred to
Th. hugii, as well as in the type specimens of Th. marina and Th. bruntrutana. However, Thalassemys specimens from England (NHMUK R8699 and OUMNH J.66966; see Discussion) and one specimen of Th. hugii from Solothurn (NMS 8555) lack a strong anterior widening of the first neural. This feature is therefore probably variable intraspecifically.

**Thalassemys hugii Rütimeyer, 1873**

*Synonymy.* Thalassemys Gresslyi Rütimeyer, 1873 and Eurysternum ignoratum Bräm, 1965 (Anquetin, Püntener & Billon-Bruyat, 2014).

*Type material.* NMS 8595–8609, almost complete and articulated carapace, disarticulated plastral fragments, postcranial remains. Lectotype designated by Bräm (1965: 143).

*Illustrations of type.* Rütimeyer (1873: plate I); Bräm (1965: plate 7); Anquetin, Püntener & Billon-Bruyat (2014: figs. 6A–6D); Figs. 3C–3D, 6C–6D and 7C–7D.

*Type horizon and locality.* Solothurn Turtle Limestone, uppermost member of the Reuchenette Formation (Autissiodorensis ammonite zone, upper Kimmeridgian, Late Jurassic), vicinity of Solothurn, Canton of Solothurn, Switzerland.

*Occurrence.* Kimmeridgian of Switzerland (Solothurn and Porrentruy) and England (Abingdon).

*Referred specimens.* All specimens referred by Bräm (1965) and Anquetin, Püntener & Billon-Bruyat (2014), except for NMS 9144 and NMS 37251; NMS 22286-22302 (material from the Solothurn St Niklaus quarry); MJSN BSY008-905 (Figs. 8A–8F); OUMNH J.66966 (Pérez-Garcia, in press; see Discussion).

*Diagnosis.* Differing from Th. bruntrutana in: proportionally wider nuchal with no anterolateral thickening on ventral surface; narrower vertebral scales; proportionally smaller plastron; narrower and more inclined xiphiplastron; smaller angle between scapular process and acromion
process. Differing from *Th. marina* in: narrower vertebrals scales; less pronounced lateral plastral fontanelles.

**Thalassemys bruntrutana** sp. nov.

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Figs. 3A–3B, 4A–4B, 5A–5C, 6A–6B, and 7A–7B

**Etymology.** The species name refers to the German form of the name Porrentruy (Pruntrut), possibly derived from Bruntrutum (meaning abundant springs) (Schweizerische Bundeskanzlei, 2015). This name is also a homage to Jules Thurmann (1804–1855), a renowned local geologist and botanist, for its fundamental paleontological work *Lethea bruntrutana* that was published after his death (Thurmann & Etallon, 1861–1864).

**Holotype.** MJSN SCR011-87, an almost complete and articulated carapace (peripherals and nuchal disarticulated), disarticulated plastron, and left scapula.

**Type horizon and locality.** Lower *Virgula* Marls, Chevenez Member, Reuchenette Formation (Eudoxus ammonite zone, upper Kimmeridgian, Late Jurassic), vicinity of Porrentruy (Courtedoux village), Canton of Jura, Switzerland (Comment et al., in press).

**Occurrence.** Kimmeridgian of Switzerland (Porrentruy and Solothurn) and England (Isle of Purbeck).

**Referred specimens.** NMS 9144 and NMS 37251; NHMUK R8699.

**Diagnosis.** Differing from *Th. hugii* in: more elongated nuchal (in anteroposterior direction) that strongly thickens anterolaterally on the ventral surface; wider vertebrals scales; proportionally longer plastron; broader and less inclined xiphiplastron; wider angle between scapular process
and acromion process. Differing from *Th. marina* in: wider vertebrals scales; less pronounced lateral plastral fontanelles.

**MJSN SCR011-87 (THALASSEMY BRUNTRUTANA)**

**General preservation**

MJSN SCR011-87 consists of an almost complete carapace and plastron associated with the left scapula. The neural series is almost complete and still in articulation with the costals. The nuchal and most of the peripherals are preserved as disarticulated elements. The remaining peripherals, the suprapygals and the pygal are missing. The different elements of the plastron are disarticulated. Epiplastra and entoplastron are missing. The uneven and fractured surface is yellowy-white in color with orange-to-reddish patches (iron mineralizations, mainly on the carapace) and black spots (manganese), the latter occasionally forming tiny dendrites along fractures. There are several traces of serpulids, mainly on the surface of the detached peripherals. Being broad but shallow, the scale sulci are not always easy to discern. Distinct linear striations extend perpendicular to sutures (mainly in anteroposterior direction), as observed in *Th. hugii* and *Th. marina* (Anquetin, Püntener & Billon-Bruyat, 2014). These striations are the most striking at the hyo-hypoplastron suture.

**Carapace**

As preserved, the (sub-complete) carapace measures 477 mm in length along the midline and 531 mm on the maximal width (at the level of costals 3–4). These dimensions are comparable to that of the lectotype of *Th. hugii* (NMS 8595–8609; Fig. 3). NMS 8595–8609 has a preserved length of about 630 mm (Bräm, 1965). However, without the nuchal and the two suprapygals
(elements that are disarticulated or missing in MJSN SCR011-87), it reaches a length of 470 mm.

With 530 mm (Bräm, 1965), NMS 8595–8609 has furthermore almost the same preserved width as MJSN SCR011-87 (both specimens missing the peripherals on either side of the carapace).

Individual bone measurements (neurals, costals) confirm that MJSN SCR011-87 and NMS 8595–8609 are approximately of the same size (Table S1). MJSN SCR011-87 is clearly larger than the holotype of *Th. marina* (SMNS 10817). However, the ontogenetic stage of SMNS 10817 is unknown.

The shell is unusually flat. This can be partially explained by taphonomic compaction, which notably flattened the carapace medially along the neurals. However, there are no important openings between the costals, which suggests that the compaction was only moderate. The original carapace vaulting must have been similar to that of *Th. hugii* (NMS 8595–8609), which is less affected by taphonomic compaction and therefore slightly more domed. Compared to *Th. bruntrutana* and *Th. hugii*, the carapaces of plesiochelyids (e.g., *Plesiochelys etalloni* or *Tropidemys langii* Rütimeyer, 1873) are clearly more domed.

As preserved, the carapace is roundish in outline, but the missing rim elements (especially the suprapygals and pygal) prevent an accurate reconstruction of the original outline (Figs. 3A and 3B). Bräm (1965) reconstructed the carapacial outline of *Th. hugii* as slightly heart-shaped, but most of the specimens he had at hand were either too incomplete or missing their peripherals. In contrast, based on undescribed material from Solothurn, Lapparent de Broin, Lange-Badré & Dutrieux (1996) suggested that the carapace of *Th. hugii* had an oval outline. However, the exact carapacial outline of *Th. hugii* and *Th. marina* remains uncertain.

**Nuchal**
The exact outline of the disarticulated nuchal cannot be reconstructed (Fig. 4A). It is roughly rectangular and proportionally more elongated than the nuchal of *Th. hugii* (e.g., NMS 8595–8609; Fig. 3D). There is no nuchal notch. The uneven notch on the anterior border of the nuchal is clearly a fracture. A wide embayment on the posterior border of the nuchal once held the first neural. The nuchal bone is thickening strongly anterolaterally on the visceral side (Fig. 4B). This distinctive feature is absent in *Th. hugii* (see Discussion).

**Neurals**

There are eight neurals (Fig. 3A and 3B). The first one misses its anterior part, but the wide embayment on the posterior border of the nuchal (see above) indicates a strong widening of the first neural towards the anterior. The following neurals are roughly hexagonal in outline with the shorter sides facing anteriorly (except for neural 8). Neurals 2 and 3 are twice as long as wide. Neurals 4–8 are then successively shorter in proportion, neural 8 being clearly wider than long. The bone posterior to neural 8 is herein interpreted as the ‘intermediate’ element, as described in *Th. hugii* and several plesiochelyids (Anquetin, Püntener & Billon-Bruyat, 2014).

**Costals**

Costals 2–5 are completely preserved on each side of the carapace (Figs. 3A and 3B). In contrast, costals 1 have incomplete anterior borders, whereas costals 6–8 lack their lateral parts. The posterior border is convex posteriorly in costals 1 and 2, but concave posteriorly in costals 3–8. As an individual variation, the distal margin of the left costal 2 is clearly longer than that of its right counterpart. The lateral borders of costals 2–5 as well as the posterolateral border of the first costal do not show any sign of suture, indicating a cartilaginous contact with the peripherals.
(possibly with minor costo-peripheral fontanelles). Although damaged by abrasion, the preserved anterolateral border of the right costal 1 suggests a sutural connection with peripherals 1 and 2.

In *Th. hugii*, the first three peripherals are sutured to the first and second costals, but the condition for other peripherals remains uncertain (NMS 8595–8609; Figs. 3C and 3D).

**Peripherals**

Among the disarticulated bone elements, seventeen peripherals or parts of peripherals have been identified (Fig. 5). However, their precise position in the carapace cannot be evaluated, mainly due to the poor preservation and the absence of sutural contacts between most peripherals and costals. Some of these peripherals are still partly articulated with one another (Figs. 5A and 5B).

The largest peripheral of the bridge area reaches 100 mm in length and 80 mm in width.

**Scales of the carapace**

As often, scale sulci are difficult to discern on the nuchal bone. One cervical scale can be discerned on the anteromedial part of the nuchal (Fig. 4A). It is about 30 mm long and 50 mm wide. However, the poor surface preservation impedes a conclusion about the actual number of cervicals. Anquetin, Püntener & Billon-Bruyat (2014) suggested that three cervicals may have been present in *Th. hugii* (based on NMS 8595–8609).

Five vertebrals are present (Fig. 3B). The first vertebral is wider than the nuchal and would have reached the (missing) first peripherals anterolaterally. Vertebrals 2 and 3 are hexagonal in shape with posterolateral margins shorter than anterolateral ones. This can also be observed in *Th. marina* (SMNS 10817). In *Th. hugii*, the anterolateral and posterolateral sides tend to be of similar length (e.g., NMS 8595–8609, NMS 8555 and NMS 8997), but some variation may exist.
for this character (e.g., OUMNH J.66966). In *Th. bruntrutana*, vertebrals 2 and 3 are about twice as wide as long and cover about half of the costal width. They are proportionally slightly wider than in *Th. marina* and significantly wider than in *Th. hugii*, where they cover only about one quarter to one third of the costal width (e.g., NMS 8595–8609, NMS 8612–8627 and NMS 8733). Vertebral 4 is the longest in the series. Its anterolateral sides are mostly straight. The outline of vertebral 5 is not preserved. The intervertebral scale sulci are concave posteriorly and cross neurals 1, 3, 5, and the ‘intermediate’ element.

Pleurals 1–3 can partially be discerned (Fig. 3B). Although their lateral borders are missing, it is evident that they are narrower than vertebrals 2–4. The interpleural 1/2 and 2/3 sulci are situated on the posterior parts of costals 2 and 4, running parallel to the intercostal sutures.

Pleurals 4 are not preserved.

**Plastron**

The hyo-, hypo-, and xiphiplastra are preserved and mostly complete (Figs. 6A and 6B), but the epiplastra and entoplastron are missing. The lateral parts of the hyo- and hypoplastra now lie within the same plane as the rest of the plastron, due to taphonomic compaction. As preserved, the plastron is as wide as long (maximal preserved length = 510 mm; maximal preserved width = 510 mm). *Thalassemys bruntrutana* has a proportionally longer plastron than *Th. hugii*. Despite the almost identical carapace size (see above), the plastron of MJSN SCR011-87 is measurably longer than the one of NMS 8595–8609 (Figs. 6C and 6D), which has a maximal preserved length of 430 mm. The length of the bridge measured between the axillary and inguinal notches is also more important in *Th. bruntrutana* (225 mm in MJSN SCR011-87) than in *Th. hugii* (190 mm in NMS 8595–8609).
Hyo- and hypoplastra

The hyoplastra are wider than long (Figs. 6A and 6B). A several centimeters wide, obconical notch that once hold the missing entoplastron separates the hyoplastra anteromedially, whereas an elongated central plastral fontanelle separates them posteromedially. In between, on a length of about 50 mm, the hyoplastra meet along an undulating contact. Anteriorly, there is no evidence of sutural contact with the epiplastra and entoplastron.

The hypoplastra are more than twice as wide as long (Figs. 6A and 6B). Separated anteromedially by the central plastral fontanelle, they meet on the posterior 65 mm of their length along an undulating contact. Only about one third of the central plastral fontanelle lies between the hypoplastra, where it is squarish to roundish in shape, in contrast to the longer, rather oval part between the hyoplastra. The lateral margins of the hyoplastra are severely damaged, but the hypoplastra show clear evidence for lateral fontanelles on their anterolateral borders (Figs. 6A and 6B). Based on NMS 8595–8609, Bräm (1965) reconstructed the plastron of Th. hugii without lateral fontanelles, but with an extensive central fontanelle that completely separates the hyo- and hypoplastra. In this reconstruction, only the posterior parts of the xiphiplastra are connected medially. However, the preservation of this specimen does not allow to conclude either to the absence of a lateral plastral fontanelle, or to the size and shape of the central plastral fontanelle (Figs. 6C and 6D). Isolated elements from Solothurn (e.g., NMS 22325) clearly indicate that Th. hugii indeed possesses a lateral plastral fontanelle and an interdigitating contact between the hyoplastra (Anquetin, Püntener & Billon-Bruyat, 2014), refuting Bräm's reconstruction. We have detected new plastral material of Th. hugii from the St Niklaus quarry in the Solothurn collections (NMS 22286, NMS 22287, and NMS 22296;
possibly from the same individual as NMS 22325) that confirm the observations of Anquetin, Püntener & Billon-Bruyat (2014), but it remains uncertain whether the hypoplastra met medially in Th. hugii, as in Th. bruntrutana and Th. marina. The lateral plastral fontanelles of Th. marina are clearly more pronounced than in Th. bruntrutana and Th. hugii.

Xiphiplastra

The xiphiplastra, which are still connected to the hypoplastra by suture, are long elements (about twice as long as wide) that are narrowing strongly posteriorly (Figs. 6A and 6B). They are clearly separated by a fontanelle anteromedially and by a small anal notch posteromedially. It is unclear whether the xiphiplastra actually met one another medially. The xiphiplastra are not as strongly inclined (in relation to the anteroposterior axis of the plastron) as in Th. hugii (e.g., NMS 8595–8609; Figs. 6C and 6D). Compared to Th. hugii, the xiphiplastra are also broader anteriorly at the contact to the hypoplastra.

Scales of the plastron

Scale sulci are only partially preserved (Fig. 6B). They show no important differences from Th. hugii (e.g., NMS 8595–8609; Fig. 6D). The limit between humeral and pectoral scales lies slightly anterior to the level of the deepest point of the axillary notches. The pectorals are slightly shorter than the preserved parts of the humerals. The hyo-hypoplastral suture divides the abdominal scale in about two equally sized parts. The femoral-anal sulcus is not preserved. Of the inframarginal scutes, only the medial borders are partly preserved.

Scapula
Only the left scapula is preserved (Figs. 7A and 7B). The glenoid fossa is only poorly preserved and partly filled with sediment. The dorsally projecting scapular process is complete. It measures 140 mm from the dorsal end to the notch between the acromion process and the coracoid. It is only slightly longer than the scapular process of NMS 8595–8609 (*Th. hugii*; 135 mm; Figs. 7C and 7D). The scapular process forms an angle of 130° with the acromion process. Due to minor post-mortem deformation, the scapular process and acromion process do not lie exactly in the same plane, which might have a slight influence on the measured angle. In *Th. hugii*, this angle is always smaller than in *Th. bruntrutana* (Table 1).

**MJSN BSY008-905 (*THALASSEMYS HUGII*)**

**Carapace**

The carapace of specimen MJSN BSY008-905 is represented by the nuchal, neurals 3–5, left costals 2–3, right costal 5, right peripherals 1–2, and nine other peripherals or parts of peripherals (Figs. 8A and 8B). All elements of the carapace are disarticulated. The bone surface is brownish-gray with distinct linear striations perpendicular to sutures. This specimen is smaller than the lectotype of *Th. hugii* (about 85% of the size of NMS 8595–8609 based on individual bone measurements).

**Nuchal**

The nuchal of MJSN BSY008-905 (Figs. 8A and 8B) is roughly trapezoidal in outline and about twice as wide as long. Anteriorly, there is a broad and very shallow nuchal notch. The ventral surface of the nuchal is flat and lacks anterolateral thickenings. These are characteristics of *Th.*
hugii and clearly exclude MJSN BSY008-905 from Th. bruntrutana. A wide embayment on the posterior border of the nuchal once held the first neural.

The anterolateral borders of the nuchal are emarginated in order to hold an anteromedial projection of the first peripherals. Such emarginations are usually absent in Th. hugii, but they occur in the specimen OUMNH J.66966 from the Kimmeridge Clay (Pérez-García, in press; see Discussion). Püntener et al. (2014) reported a comparable morphology in one specimen of Tropidemys langii from Porrentruy, although in this specimen small supernumerary bones articulate with the nuchal in this area. This morphology is therefore interpreted as an intraspecific variation.

Neurals

The three preserved neurals are hexagonal in outline with shorter sides facing anteriorly (Figs. 8A and 8B). The neural length decreases from neurals 3 to 5, but all neurals remain clearly longer than wide.

Costals

The lateral border of the second costal is rounded and smooth on the dorsal edge, but shows a sutural contact on the visceral edge, suggesting the transition from a cartilaginous to a sutural contact. Costals 3 and 5 do not show any sign of suture with the peripherals. Here, the contact to the peripherals was fully cartilaginous and minor costo-peripheral fontanelles may have been present. Costal 5 still possesses a laterally jutting rib (Figs. 8A and 8B). The closure of costo-peripheral fontanelles is slightly less advanced than in the only sub-complete adult specimen of Th. hugii (NMS 8595–8609; see Discussion).
Peripherals

Out of the eleven preserved peripherals, the precise position of only two could be identified with certainty: the right peripherals 1 and 2 (Figs. 8A and 8B). The posterior borders of peripherals 1–2 show a sutural contact to the anterior border of the missing first costal. The anteromedial corner of peripheral 1 is reaching out to fit in the anterolateral emargination of the nuchal. The second peripheral is squarish and contrasts with the elongated second peripheral in the lectotype of *Th. hugii* (NMS 8595–8609). However, the shape of peripherals is known to be variable in other closely related taxa, for example in *P. etalloni* (Anquetin, Püntener & Billon-Bruyat, 2014).

Scales of the carapace

Cervical scales are only weakly expressed. There is probably a medial cervical scale that is about 15 mm long and 35 mm wide (Fig. 8B). To its left, a narrower lateral cervical scale may have been present. However, no trace of a lateral cervical scale is preserved on the right side. Hence, the actual number of cervicals is not conclusive in this specimen. The second vertebral covers about one quarter to one third of the width of costal 3 and the third vertebral covers slightly more than one third of the width of costal 5 (Fig. 8B). This is congruent with other specimens referred to *Th. hugii*, but clearly narrower than in *Th. bruntrutana* (where vertebral scales cover about half of the costal width) and *Th. marina*. There is no trace of the intervertebral 3/4 scale sulcus on costal 5, nor on neural 5, suggesting a shift of this sulcus onto costal 6 and neural 6 respectively. Such a shift is known to occur occasionally in other taxa, for example in *Tr. langii* (Püntener et al., 2014).
Plastron

Of the plastron, only parts of the left hypoplastron, right hypoplastron, and right xiphiplastron are preserved (Figs. 8C and 8D). The bridge length is estimated to be around 170 mm. As most borders are broken, the presence of neither lateral nor central fontanelles can be confirmed in this specimen. As in other specimens referred to Th. hugii, the lateral border of the xiphiplastron is strongly inclined relative to the anteroposterior axis of the plastron, which clearly contrasts with the condition in Th. bruntrutana.

Scapula

A small portion of the (left?) scapula is preserved (Figs. 8E and 8F). The distal part of the scapular and acromion processes is missing, as well as the glenoid. The angle between the scapular and acromion processes is about 116°. This falls within the range measured for Th. hugii and contrasts with the larger angle observed in Th. bruntrutana (Table 1).

DISCUSSION

Alpha taxonomy of Thalassemys

In the present study, three species of Thalassemys are considered valid: Th. hugii, Th. marina, and the new species Th. bruntrutana. Thalassemys hugii, the type species, is based on a relatively complete shell associated with some postcranial elements (NMS 8595–8609) from the late Kimmeridgian of Solothurn, Switzerland (Rütimeyer, 1873; Bräm, 1965). The validity of this species has never been questioned. Thalassemys marina is based on a partial carapace and plastron (SMNS 10817) from the Tithonian of Schnaitheim, Germany (Fraas, 1903). Based on
the presence of a lateral plastral fontanelle, Bräm (1965) and Maisch (2001) referred this species to the genus *Eurysternum*. However, it was later demonstrated that a lateral plastral fontanelle was also present in *Th. hugii* (Anquetin, Püntener & Billon-Bruyat, 2014). Joyce (2003) tentatively synonymized *Thalassemys marina* with Palaeomedusa testa Meyer, 1860 based on the purported presence of supernumerary pleural scales in SMNS 10817, but the specimen is reconstructed in this area and this assertion does not withstand direct observation. Since SMNS 10817 exhibits significant differences with specimens referred to *Th. hugii* and *Th. bruntrutana* (see below), *Th. marina* must be considered a valid species.

The nuchal of *Th. bruntrutana* is remarkable in many aspects. It is significantly more elongated than that of *Th. hugii* and it is characterized by the presence of a strong anterolateral thickening of the ventral surface on both sides. Such morphology has never been observed in other Late Jurassic coastal marine turtles from Europe. However, a similar condition has been described in the lindholmemydid freshwater turtle *Amuremys planicostata* (Riabinin, 1930) from the Late Cretaceous of Russia, for which it is considered a diagnostic feature (Danilov et al., 2002). The nuchal of *Th. marina* is not known.

The width of vertebral scales is another distinctive feature between species of the genus *Thalassemys*. The vertebral scales of *Th. hugii* are narrow (covering 1/4 to 1/3 of the costal bones), whereas the vertebral scales of *Th. bruntrutana* are distinctly wider (covering about 1/2 of the costals). The vertebral scales of *Th. marina* are somewhat intermediate in width between those of *Th. hugii* and *Th. bruntrutana*. The width of vertebral scales is known to change during ontogeny, usually decreasing from juveniles to adults (e.g., Joyce, 2007). However, it is noteworthy that juvenile specimens of *Th. hugii* (e.g., NMS 8612–8627, NMS 8733, and NMS 8997) have the same narrow vertebral scales as the adults (e.g., NMS 8595–8609 and NMS 8997).
It must also be considered that the type specimens of *Th. hugii* and *Th. bruntrutana* are of similar size. Therefore, ontogeny cannot explain the clearly different width of vertebral scales in the two taxa. The case of *Th. marina* is more complicated since this species is known only by a single individual that is about 65% the size of the type specimens of the two other species. However, two arguments allow the hypothesis that the holotype of *Th. marina* would be a juvenile of either *Th. hugii* or *Th. bruntrutana* to be rejected. First, as aforementioned, juveniles of *Th. hugii* have narrow vertebral scales. Second, given the general tendency toward the reduction of the vertebral scale width during ontogeny, juveniles of *Th. bruntrutana* would be expected to have vertebral scales that are wider than, or at least as wide as, those of the adults.

At comparable size, the suturing of costals 1 and 2 with adjoining peripherals is more advanced in *Th. hugii* than in *Th. bruntrutana*. For example, in the lectotype of *Th. hugii* (NMS 8595–8609) peripherals 1–3 are sutured to costals 1 and 2 at least up to the anterior half of peripheral 3. In the holotype of *Th. bruntrutana* (MJSN SCR011-87; a similarly-sized specimen), only peripherals 1 and 2 are sutured with costal 1, whereas the posterolateral border of costal 1 and costals 2–5 lack sutural contacts with the peripherals (see above). MJSN BSY008-905, a specimen we refer to *Th. hugii*, confirms this difference in the timing of costo-peripheral suturing between the two species. An incipient sutural contact between costal 2 and peripheral 3 is present in this specimen, although its size is only about 85% that of the holotype of *Th. bruntrutana*

This area is not preserved in *Th. marina*, which prevents comparison.

Lateral plastral fontanelles are present in all three species of *Thalassemys*. In *Th. hugii* and *Th. bruntrutana*, the lateral plastral fontanelle is relatively narrow, even in juvenile specimens. This probably explains why the presence of these fontanelles long went unnoticed in *Th. hugii*
(see Anquetin, Püntener & Billon-Bruyat, 2014). In contrast, the lateral plastral fontanelle is significantly broader in *Th. marina*.

*Thalassemys hugii* is remarkable in the strong inclination of the lateral border of its xiphiplastron relative to the anteroposterior axis of the plastron (Table 2). In *Th. bruntrutana* and most other turtles, the lateral border of the xiphiplastron is significantly less inclined in relation to the anteroposterior axis. The xiphiplastron is unknown in *Th. marina*. Two specimens from Solothurn previously referred to *Th. hugii* (NMS 9144 and NMS 37251; see Anquetin, Püntener & Billon-Bruyat, 2014) are herein tentatively referred to *Th. bruntrutana* based on the morphology of their xiphiplastra (Table 2).

As noted by Bräm (1965), the angle between the scapular and acromion process of the scapula is more open in *Thalassemys* than in the plesiochelyid *Plesiochelys etalloni*. Within *Thalassemys*, the scapular angle ranges from 113° to 122° in specimens referred to *Th. hugii* (including MJSN BSY008-905), whereas this angle reaches 130° in the holotype of *Th. bruntrutana* (Table 1). Although the scapular angle of *Th. marina* is unknown, it appears that this measurement can be used to distinguish *Th. hugii* from *Th. bruntrutana*.

The potential effect of ontogeny and sexual dimorphism on our perception of the alpha taxonomy of *Thalassemys* must be considered. As discussed above, ontogeny can be easily dismissed to explain the differences between the three identified species. Sexual dimorphism was considered seriously notably for *Th. hugii* and *Th. bruntrutana*, which are contemporaneous species sometimes occurring in the same localities (see Paleobiogeographic considerations). In recent turtles, sexual dimorphism is primarily expressed in a difference in shell size between adult males and females (Berry & Shine, 1980; Pritchard, 2008). Furthermore, adult males often develop a concave plastron (in terrestrial species) and a shorter and wider anal notch (Pritchard,
In terms of fossil turtles, reports of sexual dimorphism are scarce and relate for instance to the shape of the anal notch, to the size of the central plastral fontanelle (Cadena, Jaramillo & Bloch, 2013), to plastral kinesis, and to tail length (Joyce et al., 2012). All these differences are directly linked to sexual selection (shell size), copulation (concave plastron, plastral fontanelle size, shape of anal notch, tail length) or oviposition (shape of anal notch, plastral kinesis) (Berry & Shine, 1980; Pritchard, 2008; Joyce et al., 2012; Cadena, Jaramillo & Bloch, 2013). In contrast, we are unable to link any observed anatomical difference between *Th. hugii* and *Th. bruntrutana* (e.g., the vertebral width or the nuchal shape) to reproductive behavior. Therefore, we consider these differences as specific and interpret *Th. hugii* and *Th. bruntrutana* as two closely related species.

**Thalassemys from the Kimmeridge Clay Formation**

Recently, Pérez-García (2015) discussed a relatively complete, but strongly flattened carapace of *Thalassemys* with associated postcranial remains from the Kimmeridge Clay Formation (late Kimmeridgian) of Egmont Bight, Isle of Purbeck, Dorset, England (NHMUK R8699). Based on the presence of linear striations perpendicular to sutures and the characteristic outline of the vertebral scales, we agree that this specimen belongs to *Thalassemys*. Pérez-García (2015) noted some differences between this specimen and the lectotype of *Th. hugii* (smaller size, wider vertebral scales, and more developed costo-peripheral fontanelles) and safely concluded that no specific determination was possible for this specimen at that time. NHMUK R8699 and the holotype of *Th. bruntrutana* (MJSN SCR011-87) have several features in common. Although NHMUK R8699 is only about 65% the size of MJSN SCR011-87, both specimens have vertebral scales of the same proportions (about twice as wide as long) and shape (clearly longer
anterolateral sides). As discussed above, the vertebral width likely represents a specific character within Thalassemys, in that case uniting NHMUK R8699 with Th. bruntrutana and distinguishing it from Th. hugii and Th. marina. The scapular angle measured on NHMUK R8699 is relatively small (about 103°; not 115° as incorrectly noted by Pérez-García, 2015), but it should be noted that this specimen has been severely flattened during fossilization: all bones and shell plates are flat and thin. This measured angle probably does not reflect the original scapular angle in this specimen. As discussed above, we can also observe a very slight deformation of the scapula of MJSN SCR011-87. However, NHMUK R8699 has clearly been more flattened during fossilization. There is almost no volume in the scapula, indicating a really strong deformation. Keeping in mind this uncertainty and the fact that several important parts of the shell (nuchal, plastron) are missing, we tentatively refer NHMUK R8699 to Th. bruntrutana and thereby report the presence of this species in the Kimmeridgian of southern England.

In 1992, Richard Wilkins, an amateur geologist, discovered the partial shell and some postcranial elements of a large turtle in the Kimmeridge Clay of Abington, Oxfordshire, England. He tentatively identified the specimen as a thalassemydid and donated it to the Oxford University Museum, where it still resides today (OUMNH J.66966). This specimen was later studied as part of a Master thesis and believed to be an indeterminate pleurodire (Harrison, 1999). However, this shell truly belongs to a thalassemydid and was recently referred to Th. hugii (Pérez-García, in press). Based on the morphology of the nuchal, width of the vertebral scales, and inclination of the xiphiplastron, we agree with this attribution.

Paleobiogeographic considerations
The paleobiogeographic distribution of Late Jurassic coastal marine turtles in Europe is largely unexplored. This is mainly the result of a poor understanding of the alpha taxonomy of these turtles. A global revision of these groups at the European scale is needed. Until recently, *Thalassemys hugii* was confidently identified only in Solothurn, Switzerland (Rütimeyer, 1873; Bräm, 1965), whereas *Th. marina* was known by a single specimen from Schnaitheim, Germany (Fraas, 1903). Recently, Pérez-García (in press) referred a specimen from the Kimmeridge Clay of England to *Th. hugii* and proposed that a second, unidentified species of *Thalassemys* was also present in the same formation.

In the present study, we identified a new thalassemydid from Porrentruy, Switzerland (*Thalassemys bruntrutana*), and tentatively proposed that this species was also present in Solothurn and the Isle of Purbeck (the unidentified specimen of Pérez-García, in press). We also described a new specimen from Porrentruy that can be confidently identified as *Th. hugii*. Our results therefore show that both *Th. hugii* and *Th. bruntrutana* are present in the Kimmeridgian of the Swiss Jura Mountains (Solothurn and Porrentruy) and of southern England (Isle of Purbeck and Abington; Fig. 9). This confirms the results of Pérez-García (in press). *Thalassemys hugii* and *Thalassemys bruntrutana* represent the first Late Jurassic coastal marine turtles that are demonstrated to have a Western European paleobiogeographic distribution. Such a result is not surprising since some thalattosuchian crocodylomorphs that inhabit the same environments also have a Western European distribution at that time, e.g., the large teleosaurid *Machimosaurus hugii* Meyer, 1837 (Young et al., 2014; Martin et al., in press). We expect that future studies will also extend the paleobiogeographic repartition of other Late Jurassic coastal marine turtles from Europe.
CONCLUSIONS

_Thalassemys_ was hitherto mainly based on a relatively complete shell of _Th. hugii_ from the Kimmeridgian of Solothurn (Switzerland). Undetermined thalassemydid material was also reported from different Western European countries (e.g., Germany, England, and France), but poor preservation limited the value of these specimens. The new material from the Kimmeridgian of Porrentruy (Switzerland) offers new insights into the anatomy, taxonomy and paleobiogeographic distribution of _Thalassemys_. The new species _Th. bruntrutana_ shows important anatomical differences to _Th. hugii_ that cannot be explained by ontogenetic variation or sexual dimorphism. Both species are simultaneously present in the Kimmeridgian of Solothurn and Porrentruy (two localities from the Swiss Jura Mountains of slightly different ages) as well as in the Kimmeridgian of southern England. These results confirm that at least some Late Jurassic coastal marine turtles had a Western European paleobiogeographic distribution. _Thalassemys hugii_ and _Th. bruntrutana_ are currently not identified in the German fossil record, but the undetermined material of _Thalassemys_ from the Kimmeridgian of Oker (northern Germany; Marinheiro & Mateus, 2011) should be analyzed in the light of the Porrentruy material. So far, _Th. marina_ from the Tithonian of Schnaitheim (southern Germany) remains the only valid species of _Thalassemys_ in Germany.

Institutional abbreviations

- **MJSN** JURASSICA Museum (formerly Musée jurassien des sciences naturelles), Porrentruy, Switzerland
- **NHMUK** Natural History Museum, London, UK
- **NMS** Naturmuseum Solothurn, Switzerland
OUMNH  Oxford University Museum of Natural History, Oxford, UK
SMNS  Staatliches Museum für Naturkunde Stuttgart, Germany

Locality abbreviations

BSY  Courtedoux—Bois de Sylleux
SCR  Courtedoux—Sur Combe Ronde.

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

The excavation sites Sur Combe Ronde (SCR) and Bois de Sylleux (BSY) are situated along the Transjurane A16 highway (gray; dotted lines indicate tunnels).
Stratigraphic section of the Reuchenette Formation.

The two *Thalassemys* specimens were discovered within the Lower *Virgula* Marls (Eudoxus ammonite zone). Scheme modified after Comment et al. (in press).
Carapaces of *Thalassemys*.

(A, B) *Thalassemys bruntrutana*, specimen MJSN SCR011-87 (Kimmeridgian, Porrentruy, Switzerland); (C, D) *Thalassemys hugii*, specimen NMS 8595–8609 (Kimmeridgian, Solothurn, Switzerland). Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci; matrix is gray. Abbreviations: co, costal; n, neural; sp, suprapygal; v, vertebral scale, *, intermediate element (see text).
Nuchal of *Thalassemys bruntrutana*.

Specimen MJSN SCR011-87 (Kimmeridgian, Porrentruy, Switzerland). (A) dorsal view; (B) posterior view with the visceral side upward, showing the strong anterolateral thickening of the nuchal.
Peripherals of *Thalassemys bruntrutana*.

Specimen MJSN SCR011-87 (Kimmeridgian, Porrentruy, Switzerland). (A) group of peripherals in dorsal view; (B) two peripherals of the bridge area in ventral view; (C) peripheral of the bridge area in lateral view.
Plastra of *Thalassemys*.

(A, B) *Thalassemys bruntrutana*, specimen MJSN SCR011-87 (Kimmeridgian, Porrentruy, Switzerland); (C, D) *Thalassemys hugii*, specimen NMS 8595-8609 (Kimmeridgian, Solothurn, Switzerland). Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci. Abbreviations: cpf, central plastral fontanelle; lpf, lateral plastral fontanelle; hyo, hyoplastron; hypo, hypoplastron; xi, xiphiplastron.
Scapulae of *Thalassemys*.

Lateral (A) and medial (B) view of the left scapula of *Thalassemys bruntrutana*, specimen MJSN SCR011-87 (Kimmeridgian, Porrentruy, Switzerland); Lateral (C) and medial (D) view of the left scapula of *Thalassemys hugii*, specimen NMS 8595–8609 (Kimmeridgian, Solothurn, Switzerland).
Thalassemy brunntrutana

Thalassemy hugii

20 mm
MJSN BSY008-905, *Thalassemys hugii* (Kimmeridgian, Porrentruy, Switzerland).

(A, B) carapace; (C, D) plastron; lateral (E) and medial (F) view of the (left?) scapula. Line width indicates natural borders (thick lines), bone sutures (medium lines), and fractures (thin lines); double lines indicate scale sulci. Abbreviations: co, costal; n, neural; nu, nuchal; v, vertebral scale; hyo, hyoplastron; hypo, hypoplastron; xi, xiphiplastraon.
Paleobiogeographic distribution of *Thalassemys*.

Localities with *Thalassemys* species from the Kimmeridgian (black stars) and Tithonian (white star) on a Late Jurassic paleogeographic map of Western Europe. Map from Ron Blakey, Colorado Plateau Geosystems, Arizona, USA (http://cpgeosystems.com/paleomaps.html).
Table 1 (on next page)

The angle between scapular process and acromion process in specimens of *Thalassemys* and *Plesiochelys*.

When the two scapulae were preserved, both were measured. The high value for the right scapula of NMS 8631–8643 must therefore be treated with caution with regard to taphonomic compaction.
| Specimen       | Species               | Scapular angle |
|---------------|-----------------------|----------------|
| NMS 8595–8609 | *Th. hugii*           | 118°           |
| NMS 8612–8627 | *Th. hugii*           | 114° / 113°    |
| NMS 8631–8643 | *Th. hugii*           | 113° / 122°    |
| MJSN BSY008-905 | *Th. hugii*    | 116°           |
| MJSN SCR011-87  | *Th. bruntrutana*   | 130°           |
| NHMUK R8699   | *Th. bruntrutana* a | 103°           |
| NMS 8584      | *P. etalloni*        | 103°           |
| NMS 8731      | *P. etalloni*        | 103°           |
| NMS 9153      | *P. etalloni*        | 105°           |

1

2 a *Thalassemys* sp. in Pérez-Garcia (2015).

3
Table 2 (on next page)

The angle between the anteroposterior axis of the plastron and the lateral border of the xiphiplastron in specimens of *Thalassemys*. 
| Specimen           | Species            | Xiphplastral angle |
|--------------------|--------------------|--------------------|
| NMS 8595–8609      | Th. hugii          | 45°                |
| MJSN BSY008-905    | Th. hugii          | 49°                |
| OUMNH J.66966      | Th. hugii<sup>a</sup> | 48°              |
| NMS 9144           | Th. bruntrutana<sup>b</sup> | 30°              |
| NMS 37251          | Th. bruntrutana<sup>b</sup> | 29°              |
| MJSN SCR011-87     | Th. bruntrutana    | 30°                |

1. cf. Pérez-García (in press).

2. Thalassemys hugii in Anquetin, Püntener & Billon-Bruyat (2014).