Eggs for breakfast? Analysis of a probable mosasaur biting trace on the Cretaceous echinoid Echinocorys ovata Leske, 1778

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Abstract. Fossil biting traces (praedichnia) represent indirect evidence of predation and shed light on fossil predator–prey interactions and fossil food webs. Especially from echinoderm skeletons, biting traces are well known. Here, we describe the oral surface of a large Cretaceous (Maastrichtian) holasteroid echinoid Echinocorys ovata Leske, 1778 from Hemmoor (northern Germany) which exhibits four circular punctures arranged in a semi-circular arc. Whereas three of the punctures penetrated the skeleton, one puncture only just hit the margin of the echinoid test at the ambitus, leaving a long incision furrow in the skeleton. The punctures were not lethal to the sea urchin as is indicated by progressed skeletal regeneration and closure of the fractures. The overall appearance of the punctures suggests that they were produced during a single mechanical event, most likely by the biting action of the teeth of a large vertebrate animal. We analysed the shape and arrangement of the biting trace and conclude that it was probably produced by a marine reptile possessing a prognath tooth position, most likely by a globidensine mosasauroid. Our finding not only sheds light on mosasaur feeding behaviour and prey selection but also increases the knowledge of the food webs in the chalk sea ecosystem during the uppermost Cretaceous.

1 Introduction

Predation plays an important role in modern and ancient ecosystems. Trophic relationships based on predation not only shape communities but also affect the evolution of both predators and their prey. Thus, fossil evidence of predator–prey interactions is important for the understanding of energy flow in ancient ecosystems, food webs, and the arms race between predator and prey. Direct evidence of predation, such as preserved gut contents, coprolites, or regurgitated masses containing identifiable prey remains are rare and exceptional findings. The study of the signs and traces preserved on the hard skeletal parts of prey organisms that reflect predatory behaviour is therefore of exceptional importance. These traces (praedichnia) include skeletal breakage, biting traces, borings, but also regeneration and repair patterns mirroring unsuccessful attacks.

Among marine invertebrates, the study of predation traces found in echinoids (Echinodermata) proves to be peculiarly rewarding. Echinoids are the diet of a wide variety of marine predators including gastropods, crustaceans, starfish, other echinoids, elasmobranch and teleost fish, turtles, birds and marine mammals (Kowalewski and Nebelsick, 2003). Previously, most authors focused on drilling predation by cassid gastropods (“helmet conchs”), since their predatory strategy leaves the echinoid test entirely intact except for a small but diagnostic borehole that can be easily recognized. In contrast, the study of shell-breaking predation is more challenging, assuming that a successful attack would result in the complete destruction of the echinoid test. However, the stereomic nature of the echinoid skeleton favours the preservation of biting traces (“tooth marks”) of unsuccessful attacks (Kähn, 1928; Gripp, 1929; Thies, 1985), allowing conclusions to be drawn as to the predator’s taxonomic identity. Biting trace analysis is based on the following two concepts or assumptions (Bowers, 2003): (i) the dental characteristics of teeth involved in biting are unique in species level and (ii) this asserted uniqueness is transferred and recorded in the biting trace. As a consequence, ideally preserved biting traces allow conclusions to be made about the predators’ dentition and consequently their identity (e.g., Neumann, 2000; Jacob-
The shape and pattern of biting traces is influenced by many factors, which not only include the biters tooth morphology and dental arrangement, which in addition may vary among individuals of one species due to ontogeny or individual wear, but also by biting angle, power of biting force, and nature of the substrate. Moreover, echinoids possess an enormous capability to survive and repair traumatic damage of the test, even when the test is penetrated (Bonasoro et al., 2004; Carnevali, 2006). Hence, skeletal regeneration process of non-lethal lesions produced by a biting attack may alter or even obscure the original shape of the biting trace.

Here, we describe an exceptional biting trace found on the basal–ambital region of a holasteroid echinoid Echinocorys ovata Leske, 1778 from the lower Maastrichtian chalk of Hemmoor. Repair features indicate that the echinoid survived this unsuccessful attack. The large size, circular outline, and arc-shaped arrangement of tooth traces suggests that the echinoid was bitten by a large predator possessing conical pointed teeth, most probably a marine reptile. We hypothesize that the echinoid was bitten by a mosasaur and experimentally tested this assumption by producing biting traces by applying a mosasaur jaw model and echinoid clay dummies. Of the trace from various angles has been undertaken with a digital DSLR after coating the specimen with ammonium chloride in order to bring out details of ornament.

For the execution of the biting experiment, a resin model skull with a movable jaw and a total length of 53 cm was produced using the skull of the type specimen of Prognathodon solvayi Dollo, 1889 as a reference (Fig. 2). We decided to build such a model, because the preservation of the original Prognathodon skull (isolated and fragile bones in a metal frame) appeared unsuitable for either producing casts or applying bite experiments. Our choice of Prognathodon solvayi is explained by its completeness, and because it matches the requirements of tooth morphology, tooth position, and stratigraphic and geographic appearance (Lingham-Soliar and Nolf, 1989). Differences in dental ornamentation occurring among different species of the genus (e.g., Lingham-Soliar and Nolf, 1989; Christiansen and Bonde, 2002; Grigoriev, 2013) are negligible in our experiment. In our experiment, biting traces from various angles were produced by pressing the lower and upper jaw of the model into artificial Echinocorys test “dummies” made of modeling clay. The resulting traces were measured, photographed, and compared with the fossil biting trace.

2 Material and methods

Our query for biting traces focussed on the echinoid genus Echinocorys, a common epifaunal deposit-feeding echinoid with a worldwide distribution known from the Turonian (Upper Cretaceous) to the Palaeogene. The applicability of Echinocorys as a model taxon for the ichnofossils-based study of biotic interactions (parasitism, predation) in deep-time has been emphasized in case studies by Neumann and Wissak (2006, 2009) and Wissak and Neumann (2006). We examined more than 7000 Echinocorys specimens obtained through our own fieldwork in the chalk of northern Germany as well as from collection specimens stored at the Natural History Museum, Berlin, the Geological Museum, Copenhagen, the Swedish Royal Museum of Natural History, Stockholm and the Federal Institute for Geosciences and Natural Resources (BGR) in Hanover, Germany. Our query resulted in the recognition of numerous predation traces. Among these, a peculiar and outstanding biting trace situated on the test of a large Echinocorys ovata from the Maastrichtian of Hemmoor (MB.E. 6565, stored in the Natural History Museum, Berlin) has been discovered which is described in the present study (Fig. 1a–c).

For descriptive purposes, each tooth imprint has been numbered (P1–P4, see Fig. 1a). Metric measurements of the test dimensions and the biting trace were taken with the aid of a precision slide caliper. Terms generally used to describe teeth (labial, lingual, mesial, distal) are used here to describe the tooth traces, a practice commonly used in forensic bite mark analysis (Bowers, 2003). Photographic documentation of the trace from various angles has been undertaken with a digital DSLR after coating the specimen with ammonium chloride in order to bring out details of ornament.

3 Geological context

The specimen of interest is of early Maastrichtian age (Belemnella sumensis Zone) and was collected in the inactive quarry at Hemmoor, which is situated about 60 km west of Hamburg (53°41′52″ N 9°08′07″ E). Previously, the quarry exposed a 141.5 m thick chalk succession of late early and early late Maastrichtian age (Belemnella sumensis Zone to Tylocidaris baltica/Oxytoma danica Zone, see Schmid et al., 2004). The chalk sediment is truly pelagic with minimal terrigenous influx and was deposited in a distal shelf environment far away from the coast (Schönfeld et al., 1995). Throughout this section, the epifaunal deposit-feeding holasteroid echinoid Echinocorys ovata Leske, 1778 is the most abundant macrofossil. Whereas Echinocorys tests from this locality often exhibit biting traces produced by sharks and teleosts (Gripp, 1929; Thies, 1985; Neumann, 2003), cases of probable mosasauroid predation have not been observed so far.

4 Results: Description of the biting trace found on Echinocorys ovata

The Echinocorys specimen exhibiting the biting traces measured 95 mm in length and 80 mm in width during life. The test is relatively stout and thick, measuring 4.3 mm at the base. Although somewhat fragmented with most of the aboral side missing, the oral (i.e., lower) surface of the echinoid is well preserved. In the posterior-central part of the oral surface, four conspicuous punctures (P1–P4) are visible which
Figure 1. The large deposit-feeding echinoid *Echinocorys ovata* (MB.E. 6565) from the lower Maastrichtian (Late Cretaceous) of Hemmoor, exposing biting traces: (a) oral surface (posterior to the right) with four tooth punctures (P1–P4); the anterior part of the test is not preserved. (b) Enlarged oblique view showing the broad linear score emanating from P4. (c) Enlarged view showing details of puncture shape and regeneration features of P1 (left) and P2 (right). Note slightly irregular outline of P2 due to chipping. Note that the echinoid’s periproct is of comparable size and shape and should not be confused with the punctures.

Figure 2. The original skull of *Prognathodon solvayi*, IRSNB R33, holotype, from the lower Maastrichtian of Mesvin, Belgium, which has been used as a template for the reconstruction of the resin scale model.
Description and measurements

**P1:** positioned centrally on plastron area; outline ovoid. Measurements: 9.5 mm (labiolingually); 5.1 mm (mesiodistally).

**P2:** positioned centrally, mostly affecting plastron but also affecting periplastronal area; outline almost circular; slightly irregular probably because small skeletal fragments have been chipped off during the biting impact. Measurements: 9.4 mm (labiolingually); 8.7 mm (mesiodistally) (Fig. 1c).

**P3:** positioned on posterior part of plastron; outline almost circular, slightly irregular. Measurements: 8.7 mm (labiolingually); 9.7 mm (mesiodistally); associated with chipping.

**P4:** positioned on posterior close to periproct and margin; outline ovoid, emanating into broad linear score running along the ambitus towards the lateral side of the echinoid test (Fig. 1b). Measurements: 9.3 (labiolingually); 5.2 mm (mesiodistally). Length of score: > 13 mm (not completely preserved).

The interdental distance between P1 and P2 is 13.6 mm, between P2 and P3 6.7 mm, and between P3 and P4 14.6 mm. All four punctures are in an equally advanced stage of skeletal regeneration. Whereas the margins of the punctures still possess sharp outlines, the openings have been completely closed with skeletal tissue, and both newly formed primary and secondary tubercles are covering the surfaces of the regenerated skeletal parts.

5 Discussion

5.1 Predator bite vs. alternative explanations of trace formation

It is necessary to clarify if the observed set of punctures is actually a biting trace or if other biogenic processes can come into consideration. Circular holes or pits in echinoids are known to be produced post-mortem by the drilling activity of bioeroders (Rahman et al., 2015), whereas circular pits may be formed *syn vivo* by the attachment of parasitic or commensalistic symbionts (Neumann and Wisshak, 2006). A formation of the trace after the echinoids’ death can be excluded: the repair features (new formation of skeletal tissue including formation of tubercles) clearly prove that the trace has been formed in the lifetime of the echinoid and that the lesion was not lethal. The echinoid survived at least long enough to repair the injuries. Chipping associated with P2 and P3 and especially the linear score emanating from the well-defined P4 clearly indicate that a physical impact from below rather than a parasite infestation affected the sea urchin. The circumstance that all four punctures are in the same state of regeneration suggests that they were produced at the same time during a single event. Together with the serial, semi-circular arrangement of the punctures, it appears very likely that they were produced by the teeth of a large animal. To produce circular punctures, conical or cylindroconical teeth with pointed tips are required (Njau and Blumenshine, 2006).

However, it may appear puzzling that only P2 and P3 are circular in outline and larger while P1 and P4 are smaller and ovoid in outline. A standard mosasauroid jaw will produce the contrary pattern, as the first pair of teeth is generally smaller than the following pairs. Why are there no further punctures preserved? This pattern can be explained by the assumption that the trace was produced by an animal with a prognathous (forward pointing) tooth position. In this case, the frontal pair of teeth would penetrate much deeper and at a steeper angle than the following pair of teeth, thus producing large and circular punctures in contrast to ovoid and small punctures by the following teeth, which hit the echinoid in an oblique angle.

Thus, this distinctive biting trace pattern suggests that the bite must have been produced by a predator with large cone-shaped teeth arranged in a prognathous orientation.

The impact leading to the formation of P4 hit the echinoid marginally at the ambitus, causing not only a puncture, but also a long and broad score running along the ambitus towards the lateral side of the echinoid.

The fact that the bite pierced the echinoid without crushing it provides clues for the attacker’s prey handling behaviour and biting mechanics. A powerful snapping bite would pierce the echinoid, in contrast to a situation involving a careful, squeezing bite. In contrast to mollusc shells, where the plywood-like arrangement of aragonite or calcite crystals leads to a lateral deflection of the biting force and thus causes irregular breakage, the meshwork structure of the echinoderm skeleton prevents such lateral biting force deflection, leading to punctures mirroring the outline of the biter’s teeth (Thies, 1985; Neumann, 2000). Thus, biting traces observed on echinoid skeletons are generally excellently preserved as may be illustrated by a further example of another *Echinocorys ovata* specimen from the Maastrichtian of Hemmoor (Fig. 3a, b), which exhibits a set of scores produced by the teeth of a teleost or shark (Thies, 1985). Moreover, in this case a large fragment of the plastron has been snapped off during the attack, but the echinoid survived and was able to completely repair the fracture. Another example (Fig. 3c) shows a specimen of the extant spatangoid *Spatangus purpureus* Müller, 1776 from the Adriatic Sea near Hvar, Croatia, exhibiting a large traumatic fracture affecting a large part of the aboral interambulacrum 4. Amazingly, this large injury was not lethal to the echinoid but has been completely repaired. Echinoids, as with most other echinoderms possess a high potential to repair traumatic injuries of the test, a strik-
5.2 Probable biting trace agents

Various trace makers can be excluded on the basis of the overall morphology of their dentition. Examples of conceivable suspects of Late Cretaceous marine vertebrates which were unable to produce similar biting traces are listed in Table 1. Late Cretaceous large teleost fishes do not match the requirements needed to produce the kind of traces found in the described Echinocorys specimen. Potential teleost candidates have large, conical teeth that alternate with small teeth, a condition found in most predatory teleosts of the Late Cretaceous (see Friedman, 2009). Cretaceous sharks in question, e.g., the widely distributed Squallorax, do not possess the cone-shaped teeth able to induce rounded biting traces as seen in the Echinocorys in question (Reif, 1973; Schwimmer et al., 1997a; Motta, 2004; Becker and Chamberlain, 2012).

In dyrosaurid crocodylians, the premaxillae are relatively narrow and very slightly inflated laterally relative to the width of the maxillae, resulting in a narrower arc of front teeth. Each premaxilla bears four alveoli, the first tooth usually being the smallest, the third tooth being the largest (e.g., Jouve et al., 2006). Dyrosaurids are documented from Late Cretaceous to Eocene sediments along the Atlantic Ocean and Tethys Sea margins (Khosla et al., 2009). Certain remains of the Dyrosauridae of predominantly Maastrichtian age have been found on the east coast of North America (Denton et al., 1997), in Argentina (Gasparini and Spalletti, 1990), Mali (Brochu et al., 2002), Sudan (Salih et al., 2016), and India (Rana and Sati, 2000). No contemporary remains are recorded to date from European deposits. Only uncertain remains of Cenomanian age have been found in Portugal (Buffetaut and Lauerjet, 1978).

Pliosaurs disappeared more than 20 Ma before the Echinocorys record described here, with the Turonian Brachasuchus lucasi as the last survivor of the group (Hampe, 2005; Ketchum and Benson, 2010; Fischer et al., 2015). Other sauropterygians can also be excluded with high probability, although recent discoveries of elasmosaur stomach contents, mainly including benthic invertebrates, have been reported from the Lower Cretaceous in Australia (McHenry et al., 2005). The Australian research group suggests a bottom feeding habitus for these Late Mesozoic long-necked pliosaurians usually interpreted as fish- and squid-feeders.

Although some elasmosaurs possess prognathous anterior tooth pairs, tooth and skull sizes in most cases do not correspond to the traces found in the Echinocorys test. Moreover, these sauropterygians lived in a different time-slice (see Table 1). The fragility of the elasmosaurid skull and the needle-like associated teeth are probably more suitable for piscivorous feeding or soft prey diets (see above, and Massare, 1987; Everhart, 2005). For the time being we cannot prove the possibility of sauropterygians producing the biting trace because of the incompleteness of records in the latest Mesozoic. Several isolated vertebrae and insignificant teeth have been reported from the latest Cretaceous worldwide (Mulder et al., 2000; Vincent et al., 2011), but the discussion shows that the type of teeth and the usually significantly lower degree of prognath anterior dentition excludes the possibility that the echinoid was bitten by a pliosaur.

A closer look at the morphology of the jaws and dentition of the Late Cretaceous rulers of the epicontinental seas, the mosasauroids, reveals eliminating factors that leave only a few taxa as candidates for having caused the biting trace on the investigated Hemmoor echinoid. The Tylosaurinae, for example, could be withdrawn from the list because they do not possess terminal teeth. Tylosaurinae developed large, prolonged rostra anterior to the premaxillary teeth (e.g., Russell, 1967; Bullard and Caldwell, 2010). The Pliopleistaciridae, the second group of derived Russellosauri-
Table 1. Examples for excluded non-mosasaurid causers of the German *Echinocorys* bite traces. Positions which are not congruent with the finding are in bold.

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Representative source</th>
<th>Morphology</th>
<th>Age/Record</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elasmobranchii</td>
<td>Reif (1973); Motta (2004)</td>
<td>non-cone-shaped teeth</td>
<td>Cretaceous</td>
</tr>
<tr>
<td>Ichthyodectiformes</td>
<td>Schwimmer et al. (1997b)</td>
<td>not all species possess procumbent anterior teeth</td>
<td>unknown from European Maastrichtian</td>
</tr>
<tr>
<td>Euteleostei</td>
<td>Friedman (2009)</td>
<td>often numerous fanglike conical teeth alternated by small teeth</td>
<td>Late Cretaceous</td>
</tr>
<tr>
<td>Dyrosauridae</td>
<td>Khosla et al. (2009)</td>
<td>narrow arc of front teeth, different tooth sizes</td>
<td>no contemporary records in Europe</td>
</tr>
<tr>
<td>Pliosauridae</td>
<td>Ketchum and Benson (2010); Fischer et al. (2015)</td>
<td>procumbent anterior teeth known</td>
<td>disappeared more than 20 Ma before <em>Echinocorys</em> record</td>
</tr>
<tr>
<td><em>Aristonectes parvidens</em> (and the following = Elasmosauridae)</td>
<td>Cabrera (1941); Gasparini et al. (2003); O’Gorman (2016)</td>
<td>relatively large cranium (0.73 m length); jaw bones containing numerous tiny alveoles</td>
<td>Maastrichtian of the Weddellian Sea</td>
</tr>
<tr>
<td><em>Elasmosaurus platyurus</em></td>
<td>Sachs (2005)</td>
<td>prognathous dentition</td>
<td>early Campanian of Kansas</td>
</tr>
<tr>
<td><em>Hydrotherosaurus alexandriae</em></td>
<td>Welles (1943)</td>
<td>skull length only 0.33 m; remarkable irregular dentition, not procumbent</td>
<td>Campanian-Maastrichtian of California</td>
</tr>
<tr>
<td><em>Kaiwhekea katiki</em></td>
<td>Cruickshank and Fordyce (2002)</td>
<td>small and homodontous needle-shaped teeth; skull length 0.62 m</td>
<td>Maastrichtian of New Zealand</td>
</tr>
<tr>
<td><em>Libonectes morgani</em></td>
<td>Carpenter (1997, 1999)</td>
<td>premaxillae and dentary with prognathous anterior tooth pairs</td>
<td>early Turonian of Texas</td>
</tr>
<tr>
<td><em>Stryxosaurus snowi</em></td>
<td>Welles (1952); Carpenter (1999)</td>
<td>prognath, but skull too short (0.47 m) and slender teeth</td>
<td>Coniacian–Santonian of Texas</td>
</tr>
<tr>
<td><em>Terminonatator ponteixensis</em></td>
<td>Sato (2003)</td>
<td>skull length 0.26 m; anteriorly procumbent tooth</td>
<td>late Campanian of Saskatchewan</td>
</tr>
<tr>
<td><em>Tuarangisaurus keyesi</em></td>
<td>Wiffen and Moisley (1986); Carpenter (1999)</td>
<td>long and narrow teeth, skull length 0.37 m; premaxillary teeth interlocked with dentary teeth which are meagerly prognath</td>
<td>Maastrichtian of New Zealand</td>
</tr>
<tr>
<td><em>Dolichorhynchops herschelensis</em> (= Polycotylidae)</td>
<td>Sato (2005)</td>
<td>extremely narrow arrangement of teeth (in parallel)</td>
<td>late Campanian-early Maastrichtian of Saskatchewan</td>
</tr>
</tbody>
</table>

rina have short heads, but a predental rostrum is absent. As a typical representative, *Platecarpus*, which occurred from the Santonian to early Campanian, has recurved, not procumbent teeth (Russell, 1967; Konishi and Caldwell, 2007), and is also an older form. *Plioplatecarpus* has proportionally small anteriormost teeth but they can be somewhat procumbent (Holmes, 1996). *Halisaurus* has small, recurved premaxillary teeth with delicate crowns (Bardet et al., 2005a). *Clidastes* of the Mosasaurinae could also be released as the biting trace producer because of its small and not procumbent anterior teeth (Williston, 1925; Russell, 1967). Species of the eponymous genus *Mosasaurus* possess procumbent premaxillary teeth to a certain degree, but they are proportionally small (see Lingham-Soliar, 1995; Mulder, 1999; Konishi et al., 2014) and certainly not favoured to handle sea urchins with effortlessness.

Having eliminated the above taxa, we are left with the globidensine mosasaurs as contenders, in which *Prognathodon* has large forward-pointed conical teeth and was widespread in late Campanian to early Maastrichtian strata of the Cretaceous North Sea basin (Dortangs et al., 2002; Machalski et al., 2003; Lindgren, 2004). Early ontogenetic stages of other Globidensini (*Carinodens, Globidens*) may also show prognath dentition (see below) and thus may also come into consideration as producers of the biting trace. At least, *Carinodens* also occurs in the Maastrichtian chalk of the North Sea
basin (Milàn et al., 2017). The likelihood that *Mosasaurus hoffmanni* (Mulder, 1999), a common inhabitant of the North Sea basin, is the agent is not very probable as outlined in the paragraph before.

5.3 Experimental testing of the globidensine mosasaur hypothesis

We tested our hypothesis using techniques adopted from forensic odontology (ABFO, 1986; Rai et al., 2006) applying biting experiments with an original-scale resin model of the skull based on the holotype of *Prognathodon solvayi* (Dollo, 1889) and *Echinocorys* clay dummies. Casts of original *Prognathodon* teeth from Maastrichtian phosphates of Morocco were applied imitating the dentition. We produced a series of biting traces applying varying biting angles and biting forces. The tooth traces on the fossil echinoid and those produced by the anteriormost teeth of the upper jaw (premaxillae) in our experiment show a striking resemblance (Figs. 1a, 3a) where P1 has been produced by the second left premaxillar tooth, P2 by the first left, P3 by the first right and P4 by the second right premaxillar tooth (Fig. 4a–c).

We could state that the producer was very probably a representative of the Globidensini (Fig. 5). In our experiment, the traces fit in size and arrangement with the fossil finding. Nevertheless, other globidensine taxa besides *Prognathodon* cannot generally be excluded. *Globidens* and *Carinodens* can also have anterior prognathid dentition (e.g., Bardet et al., 2005b; Schulp, 2005) and there is an ontogenetic change in tooth form documented for *Globidens* (Polcyn and Bell, 2005), with young animals possessing a very *Prognathodon*-like dentition. Regarding size and proportion, a producer of an early ontogenetic stage is very unlikely in our case.

5.4 Palaeoecological and palaeobiological implications

Mosasaur biting traces (including *Prognathodon*) on invertebrate skeletons have been documented so far from the shells of ammonitid and nautilitid cephalopods (Kauffman and Kesling, 1960; Kauffman, 2004), demonstrating that non-duphagous mosasaurs were able to feed on hard-shelled, nektonic prey. Although some authors questioned whether these traces were produced by biting mosasaurs (Seilacher, 1998), further work on this subject has accumulated much evidence that this interpretation was correct (Tsuji and Westermann, 2001). So far, no evidence of mosasaur predation on echinoids exists. Dollo (1913) mentioned a *Hemipneustes* echinoid as prey of *Prognathodon*. The echinoid was deposited above the mosasaur skeleton at a later time interval, but was superimposed on the mosasaur skeleton, due to low sedimentation rate (John W. M. Jagt, personal communication, 2008). Thus, this finding represents a preservation effect rather than predation evidence. Donovan et al. (2008) suggested mosasaur predation on *Hemipneustes*, but the evidence is poor and must be regarded as dubious since it is merely based on a single round pit on the echinoid test.

Our hypothesis challenges the common assumption that the nature of the prey is tightly constrained to the prey type considered optimal for a given predator tooth morphology. Globidensine mosasaurs have a powerful jaw with large blunt to pointed conical teeth pre-adapted for large, fleshy but also osseous prey (see also Massare, 1987). Our example illustrates that Globidensini such as *Prognathodon* were also able to feed on benthic armoured invertebrates such as sea urchins. Indeed, the prognathous alignment of *Prognathodon*’s anteriormost teeth appears to be suitable for picking up benthic prey from the sea floor. Moreover, we argue that top predators such as *Prognathodon* probably possessed considerable variation in foraging and feeding techniques, including benthic foraging. We assume that in the pelagic and nutrient-poor chalk-sea (e.g., Voigt and Schön-
fied, 2010; Engelke et al., 2016; Linnert et al., 2016), advantage was taken of sea urchins, which were probably the most frequently encountered prey (Fig. 5). Predatory damage caused by vertebrates observed in sea urchin tests from the chalk (e.g., Gripp, 1929; Thies, 1985; Neumann, 2003) suggests that they probably represented a major food source for a variety of predators, and most likely played an important role in the benthic-pelagic coupling and the energy flow in the chalk-sea ecosystem. In modern seas, vertebrate predators such as the Atlantic wolfish (Anarhichas lupus) depend on sea urchins as a food source when their preferred prey is scarce or absent; even though the energy content of sea urchins is relatively low (Liao and Lucas, 2000). In the Late Cretaceous of NW Europe, mosasaur remains are more common in neritic sediments (Machalski et al., 2003; Lindgren and Jagt, 2005; Hornung and Reich, 2015; Sachs et al., 2015) than in the pelagic chalk, suggesting a preference for nearshore habitats. Although the European pelagic chalk-sea was probably not the optimal habitat for mosasaurs, their capability for opportunistic feeding allowed them to exploit available nutrient resources in this harsh environment, which otherwise might have represented a barrier for mosasaur migration and dispersal.

A modern analogy for this hypothesis is documented for other marine top predators, like tiger sharks (Galeocerdo cuvieri). They are able to change or modify their foraging tactics with changing habitats and changing food availability (Compagno, 1984). A large dietary range has also been reported, for example, from toothed whales (Orcinus orca: Tomilin, 1957; Jefferson et al., 1991; Visser, 2005). Furthermore, juvenile saltwater crocodiles (Crocodylus porosus) already show opportunistic food intake (Taylor, 1979). Among mosasaurs, Tylosaurus proriger has a well documented history of being an opportunistic feeder with stomach contents revealing teleost remains, other mosasaurs (Clidastes), plesiosaurs, and Hesperornis (Martin and Bjork, 1987; Everhart, 2004). Cephalopods are also reported from stomach contents of Tylosaurus and possibly from Prognathodon (Konishi et al., 2014). However, the dentition tells us that Globidens, Carinodens, and Prognathodon were likely better suited for crushing hard shelled prey than other mosasaurs (Lingham-Soliar, 1999; Bardet et al., 2005b; Schulp, 2005; Martin and Fox, 2007), but they could also process the ordinary mosasaur diet. Other mosasaurs mostly killed their prey with powerful bites and swallowed it whole (Everhart, 2017). Robbins et al. (2008) utilized carbon stable isotopes to demonstrate segregation of foraging habits in mosasaurs at both a taxonomic and ontogenetic level. According to their results, adult Prognathodon and Globidens both exhibit an enriched δ13C value, indicating a long submergence, suggesting bottom feeding for both genera (see for more detailed discussion to effects regarding body size and diving in Schulp et al., 2013).

6 Summary

A biting trace found on the oral surface of the holasteroid echinoid Echinocorys ovata from the Maastrichtian of Hemmoor is attributed to the attack from a globidensine mosasaur, most probably belonging to the genus Prognathodon. The plausibility of this hypothesis was successfully tested by executing biting experiments with a mosasaur jaw model and artificial echinoid dummies, resulting in similar trace patterns. Our finding is the first plausible record of mosasaur predation upon echinoids. The observations indicate that mosasaurs were opportunistic top predators taking advantage of all available prey, including benthic shelled invertebrates. The chalk sea ecosystem most likely represented a nutrient-starved environments where prey selection was quantity-selective, favouring predators with an opportunistic prey selection strategy.

Data availability. Fossils, the mosasaur resin model, and the clay dummies are stored in the fossil invertebrate collection of the Mu-

Figure 5. Artist’s view of a globidensine mosasaur (based on the Prognathodon model) at the bottom of the chalk-sea feeding on benthic Echinocorys sea urchins (artwork: Elke Siebert).
Due to the nature of the “special issue” for Competing interests.

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