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Evidence for a composite organic-inorganic fabric of belemnite rostra

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Abstract: Carbonate skeletons of fossil marine organisms are widely used to reconstruct palaeoceanographic parameters. Specifically, the geochemistry of Jurassic and Cretaceous belemnite rostra is traditionally interpreted to represent near sea-surface seawater properties. More recently, an increasing number of workers, have reported significant scatter in geochemical data (e.g., $\delta 180$, $\delta 13C$, element/Ca ratio) when comparing rostra from the same stratigraphic level or within a single belemnite rostrum. This scatter is not explained by differential diagenetic overprint alone. Here we report petrographic evidence on the primary ultrastructure of rostra of Megateuthis (Middle Jurassic) and Belemnitella and Gonioteuthis (Late Cretaceous). The biogenic ultrastructure consists of a filigree framework of triaxial branches and tetrahedrons of variable size forming a honeycomb-like network. Data presented here suggest that these rostra yielded as much as 50 to 90% primary pore space. On the level of a working hypothesis - and in analogy with modern cephalopods - we propose that the pore space was formerly filled with body fluid and/or organic compounds during the life time of these organisms. Intra-rostral porosity was post mortem occluded by earliest diagenetic isopachous calcite cements of a non-biogenic origin. These may have been precipitated due to increased alkalinity related to the decay of organic matter. If this holds true, then the resulting fabric represents a composite biogenic/abiogenic structure. In order to optically separate the two calcite phases forming a single calcite fibre, we employed a wide range of state-of-the-art analytical tools to thin sections and ultra-thin sections of well-preserved specimens. Pending a verification of these well-supported ultrastructural data by means of high-resolution geochemical analyses from biogenic and abiogenic phases, we suggest that these findings have significance for those using belemnite rostra as archives of their palaeoenvironment.

Cover Letter

Dr. René Hoffmann Ruhr Universität Bochum Department of Earth Sciences Institute of Geology, Mineralogy, and Geophysics Branch Paleontology Universitätsstrasse 150, Building NA2/132 44801 Bochum Germany E-Mail: Rene.Hoffmann@rub.de

Bochum 2016-05-30

Dear Editor,

We greatly appreciate the input from two expert reviewers and have implemented the (minor) comments (see revision notes) in nearly all cases. Where we decided to not follow the reviewer's advice, we argue why we do so.

Thank you for your professional work!

Kind regards,

R. Hoffmann on behalf of the authors

Revision Notes

Dear Editor,

We greatly appreciate the input from two expert reviewers and have implemented the (minor) comments in nearly all cases. Where we decided to not follow the reviewer's advice, we argue why we do so.

General: We have included line numbering. Changed text passages are given in italics.

Reviewer SEDGEO5702_revDF - Reviewer 1

Comments made directly in the manuscript:

Line 94:

Belemnite rostra are traditionally considered to secrete their endoskeleton in oxygen isotope equilibrium with ambient seawater...

Belemnites are traditionally considered to secrete their endoskeleton in oxygen isotope equilibrium with ambient seawater...

Comment: We agree with the reviewer, not the belemnite rostra secrete their endoskeleton but the belemnite animal.

Line 127:

Thin sections (30 μ m) and ultra thin sections (< 10 μ m) of two well-preserved specimens of *Megateuthis gigantea...*

Thin sections (30 μ m) and ultra thin sections (< 10 μ m) of two well-preserved orthorostra of *Megateuthis gigantea...*

Comment: We agree with the reviewer that it is useful to distinguish between the orthorostrum and the epirostrum, both are present in *Megateuthis*.

Line 317-318:

Referring to the high porosity observed in both belemnite rostra and sepiid cuttlebone it is important to note that these structures are not homologous (Fuchs 2012).

Comment: We agree with the reviewer and added one sentence pointing out that the belemnite orthorostrum and the sepiid cuttlebone are not homologue structures. See below.

Line 334:

Apical region

Apical line region

Comment: We agree with the reviewer that it is better to refer to the apical line region connecting all apical regions during ontogeny instead of the apical region.

Line 359:

...outside of (i.e., the belemnite animal itself)...

... outside of the orthorostrum (i.e., the belemnite animal itself)...

Line 424:

shell

cuttlebone

Comment: To avoid further confusion between the belemnite rostrum and the tiny rostrum or apical spine at the posterior end of the sepiid cuttlebone, we used the general term cuttlebone for the latter which includes the highly porous phragmocone (see below).

Line 425:

...belemnite skeleton...

...belemnite orthorostrum...

Comment: As pointed out earlier by the reviewer it is more precise to refer to the orthorostrum instead of the whole belemnite skeleton which would include the phragmocone as well – which has a different mineralogy and ultrastructure and was not analysed in this study.

Line 428:

...living belemnites...

...living belemnite rostra

Comment: Changes accordingly to the comment by the reviewer.

Line 434:

...ommastrephid...

...onychoteuthid...

Comment: We changed the text accordingly to the comment by the reviewer.

Line 495:

Reviewer question: Does this concentration refers to the entire cuttlebone or only the sepiid spine ("rostrum")?

Comment: All cited references refer to the *Sepia* cuttlebone and do not specify a particular region, e.g., the spine.

Lines 518-521:

This tentative interpretation presented here with regard to belemnite rostra is arguably consistent with observations of 10-40% organic matrix in the *Sepia* cuttlebone (Birchall and Thomas, 1983; Florek et al., 2009). This is relevant as we suggest that the belemnite rostrum is structurally similar to the *Sepia* cuttlebone with regard to the primary intra-skeletal porosity.

This tentative interpretation presented here with regard to belemnite rostra is arguably consistent with observations of 10-40% organic matrix in the Sepia cuttlebone (Birchall and Thomas, 1983; Florek et al., 2009). This is relevant as we suggest that the belemnite rostrum is structurally similar but not homologous (Fuchs 2012) to the Sepia cuttlebone with regard to the primary intra-skeletal porosity.

Comment: The reviewer suggested to highlight here that the rostrum and sepiid cuttlebone are structurally similar, but not homologous. Accordingly, we added "but not homologous" to the text citing the work of Fuchs (2012). However, a discussion about the similarities and differences of the sepiid and belemnite rostrum is far beyond the scope of this paper.

Comments to the authors:

Reviewer #1: Dear authors, in some places, you compare the belemnite rostrum with sepiid cuttlebones. I recommend to provide a short explanation about the homologous shell parts of belemnites & sepiids. A clear differentiation between rostrum & phragmocone is essential to correctly interpret/comprehend observed porosities. In this context, it would be less confusing to compare the belemnite rostrum with the "rostrum" of sepiids and belosepiids (rather than their phragmocone).

Comment: We agree with the reviewer that the comparison of a belemnite rostrum with the Sepia cuttlebone might be misleading when it comes to the phylogenetic reconstruction of coleoid evolution. However, the reconstruction of coleoid phylogeny is far beyond the scope of our paper. Further, it should be noted that there is an open discussion about the homologous shell parts in fossil and modern coleoids and that authors cited in our manuscript did not distinguish between the different shell parts, e.g., Florek et al. 2009 reported on the amount of organic matter for the whole cuttlebone and did not distinguish between phragmocone, dorsal shield, and rostrum. In order to avoid additional confusion we decide to keep the comparison between the belemnite orthorostrum and the Sepia cuttlebone (which refers to the complete internal shell). We do so specifically because earlier authors, when describing porosity observed in belemnite rostra, compared it with the porous structure of the Sepia cuttlebone not distinguished between the phragmocone and other shell parts. For the "rostrum-problem" the reader is referred to Fuchs (2012). As far as we know no detailed description of the sepiid rostrum is available.

Apart from this, I would avoid the term "skeleton"; if then only in terms of the entire shell; not only shell parts, e.g. the rostrum.

We agree with the reviewer and delete whenever it seemed adequate the term skeleton.

I'm missing a paragraph discussing and verifying earlier observations/ ideas: who postulated a low porosity and what was wrong in the line argumentation?

We provide information about earlier observations of pore space or the varying amount of organic matter in belemnite orthorostra in the introduction part (lines 84-92) and in chapter 4.3 citing the most important articles by Müller-Stoll, Spaeth, and Saelen. Müller-Stoll (1936) argues for varying amounts of organic matter in the observed lighter/darker concentric rings, while Spaeth (1971, 1973, 1975) was the first who described a higher amount of porosity (up to 20%). In his extensive review Saelen (1989) comes to the conclusion that it is most likely that the belemnite rostrum was a dense structure already during life time of the belemnite animal. The latter statement (Saelen 1989) agrees with the majority of contemporaneous and all subsequent researchers (e.g., Veizer 1974, Podlaha et al. 1998) largely neglecting the porosity in belemnite rostrum (Neohibolites). To keep the fact that Spaeth could only present data for a single belemnite rostrum (Neohibolites). To keep the focus on our description of the new ultrastructure and to avoid duplication we do not provide an additional paragraph discussing and verifying earlier observations dealing with the belemnite porosity.

Reviewer SEDGEO5702 Review 27 April 2016 IJ comments – Reviewer 2

All minor corrections were implemented in the revised version of the manuscript.

Line 275:

Reviewer question: Why specifically choose these elements for mapping? Common substitutions in the calcite lattice such as Sr, Ba, Fe, Mn, Zn are not mentioned. Do these not show any variation?

Comment: Fe, Mn content was quantitatively assessed using the cathodoluminescence microscope. Both show slight variation between the biogenic precipitates and secondary cement phases. We specifically choose Mg, P, and S to collect arguments for the distribution of primary organic matter as we discussed in our manuscript.

Line 355:

... the dense fabrics observed in the case of most rostra collected in Mesozoic sedimentary successions...

... the dense fabrics observed in rostra collected in Mesozoic sedimentary successions...

Comment: We agree with the reviewer that it is surprising that no examples of preserved pore space exist as far as we know. However, fossil echinoderm remains, with a primary comparable porous endoskeleton, are also preserved as massive fossil structures with nearly no pore space left. Some belemnite rostra are preserved as hollow tubes thanks to strong diagenetic alteration processes. In some cases the outer margin of belemnite orthorostra show increasing porosity. This porosity is taken as sign of diagenetic alteration and does not resemble the described ultrastructure.

Line 384:

Reviewer statement: Yet echinoderm calcite commonly retains its intraparticle porosity in geological samples, despite being Mg-calcite!

Comment. We do not agree with the reviewers statement. Based on our own observations the majority of the primary pore space in echinoderm skeletons is occluded by secondary calcite (see also Dickson 2001, 2002, 2004). However, due to staining it is possible to reveal the original skeletal structure and distinguish it from the secondary cement phase.

Line 510:

Reviewer question: Are there any examples of cuttlebone being preserved in the way that you propose for belemnites?

Comment. As far as we know the ultrastructure of fossil sepiid cuttlebones is not described in comparable details. Further, no taphonomic framework for fossil cuttlebones is available.

Comments to the authors:

Reviewer #2: (1) The authors state (p14) that "The presence of a highly porous primary rostrum architecture during the life time of the belemnite organism as proposed here, is contrasted by the dense fabrics observed in the case of most rostra collected in Mesozoic sedimentary successions (Fig. 1A-B)."

They state "most", implying that not all belemnites have dense fabrics? I find it rather surprising that if their hypothesis is correct, why no examples of preserved porous calcite guards have ever been found. Is there any evidence of the existence of these, perhaps having been dismissed as 'altered' or 'leached' specimens? Such material would lend strong support to your hypothesis.

Comment. We agree with the reviewer that it is surprising that no examples of preserved pore space exist as far as we know. We have collected belemnite rostra from many different localities (Russia, Japan) while the primary intention was to collected hollow ammonites. Hollow ammonite preservation is very rare and requires special conditions. It can be assumed that belemnites collected from the same beds share a similar unique taphonomy. However, also those belemnite rostra collected together with hollow ammonites show a massive structure. Some belemnite rostra are preserved as hollow tubes thanks to strong diagenetic alteration processes. In some cases the outer margin of belemnite orthorostra show increasing porosity. This porosity is taken as a sign of diagenetic alteration and does not resemble the described ultrastructure. By the observation of darker and lighter concentric rings earlier researchers speculated about varying amounts of organic within these rings. Recently, Ullmann et al. (2015), based on geochemical analyses, postulated a 40% porosity for the apical line region. It was assumed by Ullmann et al. (2015) that the porosity decreases towards the rostral margin. Concluding, as far as we know no pore space had been described so far the fits with the herein described ultrastructure.

(2) The authors refer to the microstructure of echinoderm calcite (p16). They do not mention that echinoderm skeletons are also porous and subject to diagenetic infill. Yet echinoderm calcite may sometimes retain its intraparticle porosity even in Palaeozoic samples, despite originally being Mg-calcite! Why should belemnites be so uniform in their preservation style? A consequence of their unique crystallography?

Comment. We do not agree with the reviewers statement. Based on our own observations the majority of the primary pore space in echinoderm skeletons is occluded by secondary calcite (see also Dickson 2001, 2002, 2004). However, due to staining it is possible to reveal the original skeletal structure and distinguish it from the secondary cement phase.

(3) Cuttlebone is presented as the closest modern analogue to belemnite guards (e.g. p20). This seems to be perfectly logical. However, again, are there any examples recorded of cuttlebone being preserved in a similar manner to that proposed for belemnites? If the mechanism is valid, it seems rather surprising that no similar processes have been observed / described, even if only in a more superficial way. The different taphonomy a consequence of their aragonite mineralogy perhaps?

Comment. Sepiid cuttlebones are rather rare compared to the globally distributed and highly abundant belemnite rostra. We assume the different preservation originally lies in their different primary mineralogy (low Mg calcite for belemnites and aragonite for sepiids).

Thank you for your professional work!

Kind regards,

R. Hoffmann on behalf of the authors

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2	Implications for palaeoceanography and palaeoecology	Formatted: Numbering: Continuous
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4	Hoffmann, R. ¹ , Richter, D.K. ¹ , Neuser, R.D. ¹ , Jöns, N. ¹ , Linzmeier, B.J. ² , Lemanis, R.E. ¹ , Fusseis, F. ³ ,	
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27 Abstract

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29 Carbonate skeletons of fossil marine organisms are widely used to reconstruct palaeoceanographic 30 parameters. Specifically, the geochemistry of Jurassic and Cretaceous belemnite rostra is traditionally 31 interpreted to represent near sea-surface palaeoenvironmental parameters seawater properties. More recently, an increasing number of workers, however, report ahave reported significant scatter 32 in geochemical data (e.g., δ^{18} O, δ^{13} C, element/Ca ratio) when comparing rostra from the same 33 stratigraphic level or within a single belemnite rostrum. This scatter is not explained by differential 34 diagenetic overprint alone. Here we report petrographic evidence on the primary ultrastructure of 35 36 rostra of Megateuthis (Middle Jurassic) and Belemnitella and Gonioteuthis (Late Cretaceous). The 37 biogenic ultrastructure consists of a filigree framework of triaxial branches and tetrahedrons of 38 variable size forming a honeycomb-like network. Data presented here suggest that these rostra 39 yielded as much as 50 to 90% primary pore space. On the level of a working hypothesis - and in 40 analogy with Recent modern cephalopods - we propose that the pore space was formerly filled with 41 body fluid and/or organic compounds during the life time of these organisms. Intra-skeletonrostral porosity was post mortem occluded by earliest diagenetic isopachous calcite cements of a non-42 biogenic origin. These may have been precipitated due to increased alkalinity related to the decay of 43 organic matter. If this holds true, then the resulting fabric represents a composite biogenic/abiogenic 44 45 structure. In order to optically separate the two calcite phases forming a single calcite fibre, we 46 employed a wide range of state-of-the-art analytical tools to thin sections and ultra-thin sections of 47 well-preserved specimens. Pending a verification of these well-supported ultrastructural data by 48 means of high-resolution geochemical analyses from biogenic and abiogenic phases, we suggest that 49 these findings have significance for those using belemnite rostra as archives of their 50 palaeoenvironment.

51

52 Keywords: belemnite, ultrastructure, carbonate archive, diagenesis, Jurassic-Cretaceous

54 1. Introduction

55 The ultrastructure of recent biogenic carbonates is of great interest for those concerned with biomineralization research in general (Weiner and Addadi, 2011; Goetz et al., 2011), those studying 56 57 the primary biogenic skeletal structures of fossil skeletal hardparts (e.g., Coronado et al., 2013), and 58 for palaeoceanographers exploring these materials for their bearing on past climate dynamics (Saelen, 1989; Cochran et al., 2003; Parkinson et al., 2005; Jarvis et al., 2015; Immenhauser et al., 59 60 2016). Whereas the tests of planktonic and benthic foraminifera (and coccoliths -etc.) are important archives of open marine environments throughout the Cenozoic and beyond (e.g., Zachos et al., 61 62 2001), much of what is known about Cretaceous and Jurassic palaeoceanography has been deduced 63 from the geochemical archive of the calcareous rostra of extinct cephalopods, specifically belemnites 64 (Dutton et al., 2007). Reasons for the wide use of these archive materials include their abundance in 65 the fossil record, the diagenetically stable low-Mg calcite mineralogy of rostra (Veizer, 1974; Saelen, 66 1989), and the broad palaeobiogeographic distribution of belemnites in the marine Boreal and 67 Tethyan realms, (e.g., Urey-Urey et al., 1951; Voigt et al., 2003; Wierzbowski, 2004; McArthur et al., 2007; Dutton, 2007; Price and Page, 2008; Wierzbowski and Joachimski, 2009; Price et al., 2009 2011; 68 69 Li et al., 2012, 2013). As with all biogenic carbonates, however, these archives undergo post-mortem 70 diagenetic alteration, representing a major obstacle in carbonate research (Swart, 2015).

71 Characteristic geochemical patterns and fabrics in biogenic carbonate hardparts have been used 72 to test for example molluscks (e.g., Cochran et al., 2003, Sessa et al., 2015, Immenhauser et al., 73 2016), brachiopods (Parkinson et al., 2005), or foraminifera (e.g., Huber and Hodell, 1996; Kozdon et 74 al., 2011) for evidence of diagenetic alteration. In the case of ammonites, the degree of preservation 75 of nacre tablets provides evidence with regard to the preservation of these exoskeletons, to name 76 one example (e.g., Cochran et al., 2010). With regard to belemnite rostra, the observation of well-77 preservedan intact fibrous microfabrics in thin sections and polished rock surfaces in combination 78 with cathodoluminescence is commonly used to identify well_-preserved belemnite rostra (Rosales et

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al., 2001). In contrast, cloudy areas, exfoliation, fractures, stylolites, or boring traces are interpreted
as evidence for post_-mortem alteration (Saelen, 1989; Li, 2011; Benito and Reolid, 2012). A rigorous
discussion of screening techniques, including trace_-element concentrations, and isotopic ratios
applied to biogenic low-Mg calcite macrofossils, ishas been provided by Ullmann and Korte (2015).
Similarly, a "best practice" approach for the interpretation of mollusc and brachiopod carbonate
archives was presented in-by Immenhauser et al. (2016).

85 The ultrastructure of belemnite rostra was first studied by Müller-Stoll (1936). This author described organic-rich (laminae obscura) and carbonate-rich (laminae pellucidae) concentric growth 86 rings. Both of these were later shown to be calcitic but differ due to variable degrees of amounts of 87 88 occluded organic matter (Saelen, 1989). Growth rings are made up by fibrous calcite crystals 89 radiating from a central zone called apical line (Richter et al., 2011) forming what seems to be a low-90 porosity fabric (Saelen, 1989 and references therein; Fig. 1A-C). Single fibres (= radial structures of 91 Saelen, 1989) can be arranged in bundles traversing the concentric growth layers. Each fibre thickens 92 outwards and shows a sub-fibrous framework potentially first proposed - but not further explored -93 by Saelen (1989, Fig. 15a).

94 Belemnites rostra are traditionally considered to secrete their endoskeleton in oxygen isotope 95 equilibrium with ambient seawater (e.g., Anderson et al., 1994; Price and Sellwood, 1997; Voigt et al., 2003; Price et al., 2009; Wierzbowski and Joachimski, 2007, 2009). The main argument brought 96 97 forward is commonly the presence of what are considered cyclical oxygen isotope patterns 98 interpreted as seasonal seawater temperature variations (Urey et al., 1951). The PeeDee belemnite 99 used in Urey's study was considered as well-preserved based on the compact fabric and the optical 100 features of the calcite crystals (but see Li, 2011_for discussion). The assumption of equilibrium 101 precipitation was further supported by data sets from recent cephalopods including Nautilus, Sepia, and Spirula precipitating their skeletal hardparts in near-equilibrium with $\delta^{18}O_{seawater}$ (Lukeneder et 102 103 al., 2010). In contrast to this traditional view, there is an increasing amount of evidence suggesting 104 that belemnite rostra are problematic archives of their palaeoenvironment (see Immenhauser et al., 105 2016 for detailed discussion). For example, Price et al. (2015) reported an offset of about 5°C 106 between the aragonitic phragmocone and the calcitic rostrum of a single specimen of 107 *Cylindroteuthis*. This offset was regarded as vital effect but it remains unclear whether the 108 temperatures derived from the aragonite are too warm or from the calcite too cool. Similarly, high 109 intra-rostral variability of elemental (Ca, Mn, Mg, Fe, Sr) and isotopic (δ^{18} O, δ^{13} C) composition, the 110 latter with a scatter of up to 2‰, has been reported from belemnite rostra lacking evidence for 111 diagenetic alteration (Podlaha et al., 1998).

The significant discrepancy of these data sets forms a strong motivation for a reconsideration of 112 belemnite rostra as archives of their palagoeonvironment. Here, we report data from a wide set of 113 114 state-of-the-art analytical infrastructure applied to thin- and ultra-thin sections of exceptionally well-115 preserved Jurassic and Cretaceous belemnite rostra. The following aims guided this paper: First, to 116 present well-constrained petrographic evidence for the complex primary biogenic framework of 117 these rostra; second, to document evidence that points to athe highly porous nature of this biogenic 118 framework; third, to assess the relative proportion between primary skeleton and porosity; fourth, to 119 discuss the timing and nature of the pore-filling calcites phase. Evidence reported here has 120 significance for the interpretation of proxy data from ancient belemnite rostra and forms the 121 foundation of a detailed, high-resolution geochemical study that will be in the focus of forthcoming 122 work.

123

124 2. Materials and Methods

125 2.1. Belemnites

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Thin sections (30 μm) and ultra_-thin sections (< 10 μm) of two well-preserved specimens
 orthorostra of Megateuthis gigantea (Schlotheim, 1820) from the Middle Jurassic) and one specimen
 of Belemnitella mucronata (Schlotheim, 1813) and Gonioteuthis quadrata (Blainville, 1827) from the
 Upper Cretaceous) were studied. Specimens of Megateuthis were collected in Bajocian marly

131 limestone deposits from southern Germany. Belemnitella rostra are from the late early Campanian

(mucronata Zzone), and Gonioteuthis from the early Campanian lingua/quadrata – gracilis/mucronata Zzone from northwestern Germany. Gonioteuthis and Belemnitella rostra were embedded in calcareous (65-90%) epicontinental shelf deposits of the Misburg fromation (Niebuhr, 135 1995; Niebuhr et al., 2007). Specimens did not display evidence for exfoliation nor did they show boring traces of the surfaces of rostra. Specimens were sectioned along their long axis in a marginal position and perpendicular to the c-axis of their calcite fibres. One section cuts the rostrum of *Megateuthis* perpendicular to the long axis.

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140 2.2. Methods

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Surfaces of thin sections were chemo-mechanically etched using colloidal silica (OP-S) for 5-15 minutes to reduce surface irregularities on an atomic scale (Massonne and Neuser, 2005) and coated by a thin carbon layer. All coated thin sections have been studied under a high-resolution field emission scanning electron microscope (HR-FESEM) type LEO/ZEISS 1530 Gemini using a backscatter detector (BSD) at the Ruhr-Universität Bochum, Germany (Figs. 1-3).

147 Crystallographic orientation of belemnite calcite fibres was determined by electron backscattered diffraction (EBSD; Nordlys, OXFORD Instruments). The data acquisition and analysis 148 149 was performed using the software packages AZtec and Channel 5 by Oxford Instruments (Fig. 4). The 150 scanning electron microscope (SEM) was operated at beam energy of 20 kV, an aperture of 60 µm, a 151 working distance of 25 mm and a tilt angle of 70°. Thin sections were mapped at Bochum University 152 in the high-resolution mode using a grid matrix (1149x748 points) at a step rate-width of 2.017 µm 153 (Fig. 4A) and (543x266 points) at a step rate-width of 1.652 µm (Fig. 4D) and for the single calcite fibre 50 μ m to reduce artefacts and increase the reliability of the data. The orientations of the 154 crystals of in the individual maps were visualized using a rainbow colour coding ranging from blue 155 156 over green and yellow to orange and red, where identical colours indicate identical crystal axis Formatted: Font: Italic, No underline, Font color: Auto Formatted: Font: Italic, No underline, Font color: Auto Formatted: Font: Italic, No underline, Font color: Auto Formatted: Font: Italic, No underline, Font color: Auto

Formatted: Font: Italic, No underline, Font color: Auto orientations. For visualizing the weak angular deviations in the crystal lattice in a single calcite fibre,
we applied an angular resolution of 2 degrees for the complete rainbow colour range. In addition,
orientations of the measured crystallographic axes <u>awe</u>re plotted into the lower hemisphere of a
Schmidt net (Fig. 4E).

Rostra were further investigated under a cathodoluminescence microscope type HC1-LM by Lumic equipped with a hot cathode (Neuser et al., 1996) and a digital camera system (DP73 by Olympus) for recording digital images at Bochum. Beam energy of 14kV and a beam current density between 5 and 10 μ A/mm² were generally used for the CL-measurements. Integration times for CLspectra were commonly between 10 and 60 seconds (Fig. 5).

166 X-ray element distribution maps were acquired using a Cameca SX5FE field emission electron 167 microprobe at Bochum. The acceleration voltage was 15 keV with a probe current of about 80 nA and 168 a fully focused beam. The intensity of S Kα1 was recorded simultaneously on two wavelength 169 dispersive spectrometers equipped with LPET and PET analyzing crystals. The Mg Kα1 line was also 170 measured on two spectrometers (LTAP and TAP crystals), whereas P Kα1 was measured on a single 171 spectrometer equipped with a LPET crystal. The images were acquired in continuous stage scan 172 mode. They have a resolution of 2048 x 1536 pixel, and the dwell time was 17 ms per pixel (Fig. 6).

173 To test for the distribution of organic matter within the belemnite rostra, thin sections were 174 studied under thea fluorescence microscope (Leica DM4500P) equipped with a mercury short-arc 175 reflector lamp coupled with a Leica EL6000 compact light source. We used the blue light filter set 176 producing bright green fluorescence images (filter set I3 for blue light excitation: excitation 450-490 177 nm, emission 515 nm, voltage 100-240 VAC and 50-60 Hz frequency; Fig. 7). Fluorescence microscopy 178 in the manner applied here is an optical tool to qualitatively document the spatial distribution of 179 organic matter in shells. Fluorescence reflects organic matter and less commonly crystal lattice 180 defects and solid inclusions in crystals. Fuorther more details of this methods have been presented please refer toby Wanamaker et al. (2009) and Ritter et al. (subm.). 181

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182 Confocal laser fluorescence microscopy (CLFM) images on an uncoated *Megateuthis* mount 183 were made using a Bio-Rad MRC-1024 scanning confocal microscope at the W. M. Keck Laboratory 184 for Biological Imaging at UW-Madison (Fig. 8). The microscope was operated with a 40 mW laser at 185 wavelengths of 488 nm, 568 nm, and 647 nm. All three wavelengths were simultaneously rastered 186 across the sample. Naturally occurring compounds within the sample caused fluorescence at multiple 187 wavelengths. Images were collected through the following three emission filters: visible green light (λ 188 = 505 to 539 nm);_j visible red light (λ = 589 to 621 nm);_j and far-red light (λ = 664 to 696 nm).

189 In order to analyze the three dimensional filigree framework, synchrotron radiation based 190 micro-computed tomography was applied (Fig. 9). Data were collected at the bending magnet beam line 2-BM at the Advanced Photon Source, Argonne National Laboratory, USA. A double multilayer 191 192 monochromator of 1.5% band-width provided 27.2 KeV X-rays. Images were collected in transmission 193 mode by a CCD camera behind the sample in the hutch configuration. The sample-detector distance 194 was set to 300 mm to collect quantitative phase contrast data. 1440 projections were acquired while the sample was rotated over 180° in steps of 0.125°. A microtomographic data set with a size of 2048 195 196 x 2048 x 1948 voxels was reconstructed using a phase retrieval algorithm (Mokso et al., 2013).

197

198

199 **3. Results**

3.1 Optical-, cathodoluminescence-, fluorescence-, and confocal laser fluorescence microscopy (TL, CL,
 FL, CFLM)

202 Under transmitted light (TL), thin sections of rostra cut perpendicular to the c-axis of calcite 203 fibres reveal a banded distribution of calcite fibres containing brownish triangular elements of 204 variable size (Fig. 5A, D). Some of the larger fibres contain a triangular, organic-rich centre, with its 205 innermost domains occluded by translucent calcite. Bundling of neighbouring calcite fibres is 206 indicated by their subparallel orientation of their a-axis (Fig. 5D). A uniform extinction (orientation) pattern of adjacent fibres under crossed polarizers is observed (Fig. 5B, E) and is in agreement with EBSD data (Fig. 4A). Individual fibres display an undulatory extinction (converging or diverging c-axes) under crossed polarizers. Calcite occluding space between organic-rich elements and fibre reinforcement is translucent and contains little or no organic matter (Fig. 5D). Brownish, triangular elements have a relatively low optical relief, compared to the surrounding translucent calcite with a relatively higher optical relief (supplement Fig. 1A-B).

Three different luminescence patterns are observed under the cathodoluminescence microscope (CL): (i) A light blue luminescence of the brownish, triangular elements (Fig. 5C, F); (ii) dark blue, intrinsic luminescence of the translucent phase; (iii) locally, orange to red luminescent fractures and microstylolites are observed. In rare cases, fractures retrace the triangular outline of a fibre (Fig. 5C, F, supplement figure 1D).

Brownish triangular areas under transmitted light display light green fluorescence while the translucent areas show a dark green fluorescence under the fluorescence microscope (FL). Microfractures are darker under transmitted light compared to the triangular elements and show a light green fluorescence (Fig. 7A-B).

Different fluorescence patterns under the CLFM reveals calcite domains visible in BSE (and other) imaging techniques (Figs. 3, 5-7). Filled cracks fluoresce brightly in green and red wavelengths (Fig. 8B-C). Domains that are dark in BSE do not fluoresce in CLFM (Fig. 8A-D). Bright domains in BSE fluoresce brightly in CLFM at all wavelengths observed (Fig. 8A-D). Higher magnification reveals brighter fluorescence between adjacent domains that appear bright in BSE (Fig. 8D).

227

228 3.2 Scanning-electron microscopy equipped with a backscatter detector (SEM BSD)

Images collected with the scanning electron microscope equipped with a backscatter detectorpresent important evidence for the presence of two calcite phases (dark and bright) of different

231 chemical compositions building the rostra studied. The brownish triangular areas under transmitted 232 light are dark in BSE. Thin sections cut perpendicular to the c-axis of the radiaxal fibrous calcite (Fig. 233 1D I-IV) reveal a complex framework, dark in BSE, surrounded by relatively brighter calcite (Figs. 1-3). 234 Cross sections of the majority of fibres are polygonal or, less often, honeycomb shaped with a tri-235 radial (120°) symmetry representing an ultrastructure that has not yet-been described in previous 236 studies (Figs. 1C-G, Fig. 3A-E). Individual fibre diameters vary between 10-80 µm. Calcite fibres 237 terminate at individual concentric growth layers that also form the nucleation site for the overlying, 238 next fibre generation, displaying increasing thicknesses and occurring in increasing numbers towards 239 the outer portions of the rostrum (Fig. 1A-D). Brighter area in BSE are more likely to be composed of 240 near-stoichiometric CaCO₃ with higher average atomic masses (mainly Ca), whereas excess C, P, Mg, 241 or S will lower the average atomic mass. Accordingly, areas with decreasing amount of Ca and 242 increasing amounts of C, P, Mg or S are darker.

243 Four section planes are presented to describe the intricate bio-composite mineral present within a single fibre. The following description distinguishes between the darker framework (i), i.e., 244 245 micronmetre-thick branches terminating in wall-like reinforcements and (ii) tri-radial central 246 portions; and brighter fabric (iii) consisting of calcite crystals forming isopachous cement layers with individual crystallites coated by submicronmetre thick layers of matter darker in BSE (Fig. 1G). In all 247 248 sections, we observed isopachous calcite crystals oriented perpendicular to the inner dark walls (Fig. 249 3A-E). Section plane I consists of three simple, dark in BSE branches of variable lengths. Often, these 250 branches exhibit reinforcements of variable lengths and thicknesses (about 1 micrommetre) and 251 increasing widths towards neighbouring branches (Figs. 1D-I, 3A). Conversely, reinforcements being 252 connected to neighboring triangles are rarely observed. Section type II is characterized by an 253 enlarged, dark in BSE central element of variable diameter (2-30 μm, Figs. 1D-II, 3C). Some of the 254 larger central elements display an inner zone with additional tri-radial structures comprising of 255 smaller, isopachous, brighter in BSE area calcite crystals rotated by 60° relative to outer branches 256 (Figs. 1D-III, 3D; section type III). Section type IV represents the most complex fabric. Here, the inner 257 portions of larger central elements display alternating darker and brighter in BSE areas (Figs. 1D-IV,
258 3E). Generally, enlarged central elements correlate with a shortening of corresponding, darker
259 branches.

260 Thin sections cut parallel to the c-axis of the radiaxial fibrous calcite reveal an framework dark in 261 BSE of triangular shaped elements with their tips pointing towards the outer margin of the rostrum (Fig. 1C-D, supplement Fig. 2). Lighter and darker areas within a single calcite fibre show their 262 263 composite nature. Lighter and darker areas within concentric growth rings, as visible in transmitted 264 light, depend on the number and size of these elements. Higher magnification reveals a homogenous 265 central portion ("trunk" - white line) of pyramidal morphology dark in BSE surrounded by inclined 266 isopachous calcite crystals coated by material dark in BSE. This overall pattern results in a "*-Christmas tree like" structure (Fig. 3F). The dark framework forms the substratum upon which an 267 268 isopachous, translucent calcite phase nucleated (Fig. 1D, G 2-3). The boundary surface between the 269 framework dark in BSE and the surrounding carbonates brighter in BSE is, in places, corroded and 270 uneven (Fig. 2).

271

272 3.3 Electron microprobe analysis (EMPA)

273	The contrasting chemical composition of the two different calcite phases building these rostra is
274	also revealed by EMP data. Detailed WDS scans show that the strongest variability is displayed by
275	Mg, P, and S, whereas other elements are almost constant or containedpresent in amounts close to
276	the detection limit (e.g., SrO = 0.1-0.2 wt.%; FeO = <0.1 wt.%; MnO < 0.5 wt.%. X-ray element
277	distribution counts-maps show a generally low concentration of the elements Mg (MgO = $0.3-0.4$)
278	<u>wt.%</u>), P_($P_2O_5 = 0.1-0.2$ wt.%), and S_($SO_3 = 0.20-0.50$ wt.%). The Mg- and P-content is slightly
279	elevated in the darker, triangular areas in BSE images (MgO up to 0.6 wt.%; P_2O_5 up to 0.25 wt.%)
280	compared to the relatively brighter areas in BSE images in the centre of larger tetrahedrons and their
281	surroundingsvicinity, Lowest Mg values-concentrations were collected-found along microfractures

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that show adisplay bright luminescence in CL_colours (MgO < 0.2 wt.%; Fig. 6B). Sulfur has a higher concentration in the brighter in BSE areas in BSE images (SO₃ up to 0.65 wt.%) and lower concentrations in the darker areas in BSE areaimages ($\frac{1}{5O_3} = 0.20-0.50$ wt.%; Fig. 6D). For better comparison with BSE images from other samples and with the element distribution, we collect BSE data were obtained for the same field of view (Fig. 6A).

287

288 3.4 Electron Backscattered Diffraction imaging (EBSD)

289 Electron backscattered diffraction was used to document the biological controlled (sensu 290 Lowenstam and Weiner, 1989; Dupraz et al., 2009; Weiner and Addadi, 2011) formation of what is 291 here assumed to represent the primary biogenic skeletal carbonate. EBSD reveals a very low 292 variation of c-axes orientation of calcite fibres (Fig. 4A). The sub-parallel orientation of the c-axes 293 {001} is documented by their close fitting in the lower hemisphere of a Schmidt net (Fig. 4E, left plot). 294 A different characteristic is found for the crystallographic orientation of the a-axes {010} of calcite 295 fibres (Fig. 4E, right plot). All a-axes are aligned along a great circle whilst the colour code of the 296 according fibres is indicative of an arrangement in bundles. Adjacent fibres (10 to 100 fibres) share a 297 similar - if not identical - orientation (Fig. 4A-B). The majority of fibre bundles are characterized by 298 blue, green and red colour coding. Specifically, the spatial orientation of individual a-axes within a 299 single fibre bundle deviates by 20° or less from the bundle mean value. Variation in the orientation of 300 the c-axis of a single calcite fibre in the range of 1-3° was observed (Fig. 4D). As the angular 301 resolution from blue to red rainbow colour shown in EBSD maps (Fig. 4A-B) has a resolution of 50°, 302 minor angular deviations (<1°) are not visible in our image Fig. 4A-B.

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- 304

3.5 Synchrotron radiation based micro-computed tomography (SRµCT)

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305 Tetrahedral structures are visible in three dimensions in the tomographic dataset despite the 306 high signal to noise ratio and artefacts. There is a distinct difference in X-ray attenuation between the 307 inner tri-radial elements and the surrounding calcite.

308

309 4. Interpretation and Discussion

- 310 4.1 Reconstruction of the primary belemnite rostrum ultrastructure
- 311

312 Data presented here document a repeated pattern of triangular elements that seem to be 313 originally connected, building a concentric layered, highly complex, and porous framework. The 314 space between the former skeletal elements is interpreted as pore space probably filled by body 315 fluids or organic material during the life time of these organisms. The latter assumption is based on 316 analogous observations in the porous endoskeletons of recent Sepia (Sherrard, 2000; Guerra, 2006). 317 Referring to the high porosity observed in both, the belemnite rostraum and the sepiid cuttlebone, it 318 is important to note that these structures are not homologous (Fuchs 2012).

319 Accordingly, the belemnite rostra originally consisted of an organic-rich biogenic framework of 320 calcitic tri-radial tetrahedrons (triangular pyramids arranged perpendicular to the concentric layers 321 during lifetime). Tetrahedrons are elongated along their c-axis with the tip of the pyramid pointing 322 towards the precipitation site i.e., in growth direction towards the outer margin of the rostrum (Fig. 323 1C-D). Individual branches protruding from the edges of the tetrahedrons possess outer 324 reinforcements acting as stabilizers. The central portion of the tetrahedrons may yield a channel-like 325 cavity (Fig. 1D III-IV). Organic membranes at which nucleation of the tetrahedrons may have started and stopped are not preserved. However, it seems likely that such membranes were present because 326 327 the bases of the tetrahedrons follow a concentric layer. Structurally, the bundling of the tri-radial 328 elements forms a simple honeycomb-like framework in which the terminations of branches of

adjacent elements are connected. The mechanical stability of the honeycomb structure is enhanced
by reinforcement walls (Figs. 1D, G, 3C).

331 Based on the variable dimensions of skeletal elements seen in thin sections, intra-rostral pore 332 space was visually estimated to range between 50-90% of the total rostrum (Fig. 3A: 90% pore space 333 and 10% skeletal elements; Fig. 3B; 50% pore space and 50% skeletal elements). In this context, the 334 observation that pore space is not limited to the apical line region but is present across the bulk 335 rostrum is important. The secretion of a porous, but mechanically stable endoskeleton-orthorostrum 336 is probably best seen in the context of a considerable reduction of energy and building material required to form this structure compared to a massive structure endoskeleton (Sherrard, 2000). 337 338 Strict biological control, i.e. in the presence of organic templates (Chateigner et al., 2000; Richter et 339 al., 2011), over the precipitation of primary skeletal elements is demonstrated by a systematic 340 arrangement of adjacent elements into bundles of similar or identical orientation of their 341 crystallographic a-axes. This configuration results in a much higher mechanical load capacity and 342 torsion stiffness of the framework of the rostrum.

343 Alternating concentric Ca-rich (brighter; laminae obscurae) and C-rich layers (darker; laminae pellucidae) of Müller-Stoll (1936) are related with the banded distribution of larger and smaller 344 345 organic-rich triangular elements of the primary skeletonal structure of the belemnite orthorostrum 346 (Fig. 1D, supplement Fig. 2). These layers potentially indicate differences in calcite precipitation rates. 347 Based on petrographic and ultrastructural evidence, the Mg and Ca elemental concentrations of the 348 rostrum, the lack of secondary micro-dolomite, and the absence of blotchy luminescence, the widely 349 held assumption of a primary low-Mg calcite mineralogy for the studied belemnite rostra is 350 confirmed (see discussion in Richter et al., 2003).

351

352 4.2 Early and late diagenetic processes

353

354 The presence of a highly porous primary rostrum architecture during the life time of the belemnite organism as proposed here, is in contrasted byto the dense fabrics observed in the case of 355 356 most rostra collected in Mesozoic sedimentary successions (Fig. 1A-B). Hence, the diagenetic 357 pathway from porous to dense fabrics deserves attention. The marine diagenetic alteration of 358 biominerals is initiated directly after the death of a carbonate-secreting marine organism when 359 metabolic processes come to a halt. At this early stage, organic matter outside of the orthorostrum 360 (i.e., the belemnite animal itself) and in the pore space of rostra and between biominerals 361 decomposes (Saelen, 1989), triggering a series of complex bio-chemical processes. Essentially, decomposition of organic matter is mediated by microbial activity, and given the abundance of 362 363 marine microbial life, there is no reason to assume that this would have been different in the case 364 examples studied here.

365 Microbial metabolic products, the presence of microbial "mucus" (extracellular polymeric 366 substance) and charged surfaces represented by microbial bodies influence the micro-environment 367 in intra-rostrum pore space by altering the balance between more reduced and more oxidized forms 368 of carbon as previously summarized under the term "alkalinity engine" (Dupraz et al., 2009). 369 Specifically, Visscher and Stolz (2005) subdivided microbial species into 5-7 groups ("guilds") having a 370 similar metabolism. Some of these promote carbonate precipitation (e.g., cyanobacteria, sulphate 371 reducers), whilst others favour dissolution (e.g., aerobic heterotrophs, fermenters). Dupraz et al. 372 (2009) documented that the balance of microbial metabolic activities directly influences carbonate 373 precipitation or, vice versa, dissolution. Initial microbial decomposition of organic matter may result 374 in the production of organic acids lowering the pH-values in the pore space (Berner et al., 1978), 375 enhancing intra-skeleton-orthorostrum dissolution of biominerals (Fig. 2).

As soon as the bulk of organic matter is decomposed, alkalinity is increased due to microbial metabolic products enhancing the precipitation of carbonate minerals (intrinsic organomineralization) leading – in the view of the authors - to the formation of the translucent, isopachous calcite crystals that occlude primary pore space in rostra (Fig. 2). Obviously, any assumption regarding microbial processes in these ancient carbonates must remain speculative. Nevertheless, following the basic concepts laid out in Dupraz et al. (2009), we tentatively assume that a first phase of decomposition was dominated by aerobic heterotrophy, sulphide oxidation, and fermentation decreasing the saturation index and resulting in the corrosion of skeletal elements in the belemnite rostrum. Evidence for this comes from micro-corrosion features at the outer surfaces of the primary skeletal framework forming the substratum upon which the pore-filling, secondary calcite phase nucleates (Fig. 2).

387 The nucleation and precipitation of the secondary calcite phase (Fig. 1D, G, 2-3) was possibly 388 dominated by sulphate reduction increasing the saturation index and hence favoring precipitation of 389 CaCO3. During crystals growth, remnants of belemnite organic matter combined with microbial 390 mucus were likely trapped between single crystals or at the growth front of crystals and delineate 391 crystal boundaries. Specifically, the growth of fibrous calcitic crystals will proceed as long as growth 392 rate, fluid supersaturation, and temperature are low enough to discourage spontaneous nucleation 393 (Oti et al., 1989). Similar processes have been described from extant echinoderm endoskeletons 394 skeletal remainsthat share a comparable amount of primary porosity like the hereinwith the 395 described belemnite orthorostra (Richter et al., 2003).

396 During this early stage of diagenetic evolution, the rostra most likely preserved their original 397 morphology due to the biogenic calcite framework and abiogenic calcite progressively occluding 398 former pore space preventing, in combination, a collapse of the endoskeletonsrostra. The directly 399 comparable, dark blue intrinsic luminescence of the translucent secondary outer calcite layer and the 400 calcite infill of central pore space of many of the larger triangles (Figs. 3D-E, 5A, D, 6) suggest that 401 both phases precipitated from one fluid, or different fluids with near-identical geochemistry. This is, 402 of course, within the limitations of the geochemical resolution of the CL-cathodoluminescence method and with reference to elements that affect luminescence patterns (e.g., Mn²⁺, Fe²⁺ and REE; 403 see discussion in Barbin 1991, 1993; Ritter et al., 2015). Conversely, the organic matter-rich triangles 404 405 display a moderately brighter blue luminescence pattern (Fig. 5C, F). This feature is best explained by

calcite lattice deformation due to the incorporation of organic matter into the crystal lattice
(intracrystalline) and is not necessarily indicative of a different geochemical composition. This
<u>concept</u> is supported by x-ray diffractometry (Richter et al., 2011) documenting that the fibrous
fabric of the belemnite rostra, lacking late diagenetic Mn-rich cements, is composed of
stoichiometric or near-stoichiometric calcites (d (104) = 3.030 to 3.035 Å).
A late diagenetic (burial) stage of rostra is documented by dissolution and compaction features

as indicated by microfractures and microstylolites (Figs. 2, 5C, F; Rosales et al., $2004a_{f_{2}}^{\prime}$ b). The circulation of Mn²⁺-rich fluids caused the precipitation of a late calcite phase that occludes fractures and fissures. In some cases this late phase yields a bright luminescence and is zigzag or triangular shaped, tracing the morphology of the triangular biominerals (Fig. 5C).

416

417 4.3 Implications for the function of the rostrum

418

It is generally accepted that the belemnite rostrum acts as a counterweight to the soft body.
Based on the observation of a high primary porosity (see also Spaeth, 1971, 1973, 1975; Ullmann et
al., 2015), this interpretation requires renewed consideration. Rostra are commonly considered to
have the same density as inorganic calcite crystals, ranging from 2.5-2.7 g/cm³.

423 Sepia, the closest living relative of the extinct belemnites, incorporates a total of 10-40% organic 424 matter in its cuttlebone shell and comparably high amounts of intracrystalline organics were 425 observed for the biogenic belemnite skeletonorthorostrum. Accordingly, ignoring open pore space, a reduced density of the biogenic belemnite calcite of about 2.4 g/cm³ (10% organic) - 2.0 g/cm³ (40% 426 427 organic) results. Assuming that liquid or extracrystalline organic matter (1.03 g/cm³) - with density 428 comparable to that of seawater (1.026 g/cm³) - filled up the pore space of living belemnite rostra, an 429 overall density of the rostrum ranging between roughly 1.7-1.1 g/cm³ (mean 1.4 g/cm³) is tentatively 430 assumed on the level of a working hypothesis. If these assumptions hold true, then the belemnite rostrum had a cumulative density that is significantly lower than that of stoichiometric calcite.Therefore, questions regarding the locomotion of belemnites result.

433 A possible analogue may come from a structure reported from a modern onychoteutmmastrephid squid (Onykia) that has remarkable morphological similarity to the 434 435 Megateuthis rostrum. Onykia has a purely organic rostrum that due to its very low density does not 436 act as a counterweight for the soft body. It is assumed that it supports the posterior part of the 437 mantle and fins (= axial stability in Bizikov and Arkhipkin, 1997 and Arkhipkin et al., 2015). A function as a muscle attachment structure for belemnite rostra hwas been first put forward by Stevens 438 (1965). Direct evidence for the presence of fins in belemnites has recently reported by Klug et al. 439 (2015), favouring a squid-like high speed swimming inmode of life for Jurassic belemnites. 440 441 Noteworthy to report here, is the case of belemnites (Chitinoteuthis) with a non-calcified rostrum 442 (Müller-Stoll, 1936).

443

444 5 Open questions and suggestions for future research

445 5.1. Paragenesis of porosity-occluding calcite phase

446 Within individual belemnite rostra, data presented here differentiate: (i) athe biogenic, highly 447 porous skeletal framework secreted during the life time of the belemniteorganism; (ii) the an inorganic or organomineralic – arguably early diagenetic - calcite phase occluding the pore space of 448 449 the biogenic framework; and (iii) a late diagenetic, burial, Mn-rich carbonate phase filling fissures and 450 larger cracks within the rostra. Assuming that the above-discussed paragenetic succession is valid, a 451 series of open questions result.+ In the view of the authors, it is at least conceivable that portions of 452 the rostral pore space were occluded during the life time of the belemnites (remote 453 biomineralization sensu Hücker and Hemleben, 1976; Chinzei and Seilacher, 1993; Seilacher and 454 Chinzei, 1993). If this holds true, then the paragenetic sequence of primary framework and 455 secondary infill calcite is even more complicated than presented here and gradual in nature. 456 Moreover, individual growth increments within rostra are then not representative of specific 457 correlative time intervals. Specifically, each growth increment then represents a complex composite 458 structure of paragenetic phases representing temporally different stages in the belemnite 459 ontogenetic cycle. Obviously, this would render the interpretation of time series belemnite 460 geochemical data difficult. Evidence against a biogenic infill of the skeletal pore space by remote 461 biomineralization sensu Seilacher and Chinzei (1993), however, may or may not come from the 462 presence of a corroded outer surface of what is considered the primary skeletal ultrastructure of 463 these rostra (Fig. 2). It seems difficult to argue that intra-rostrum body fluids became corrosive at some stage during the life time of the belemnite animal. Clearly, these questions require further 464 465 detailed work.

466

467 5.2 Primary skeletal ultrastructure and preservation of organic matter

468

The authors acknowledge the fact that despite the very detailed information regarding the belemnite ultrastructure shown here, our study lacks direct evidence for a primary biogenic origin of the complex, highly porous framework and the subsequent cementation by an early diagenetic calcite phase. Open questions, however, remain. Specifically, the significance of preserved organic matter in the biogenic belemnite calcite deserves attention. In the following, we present several lines (petrographic, optical, and geochemical data) of circumstantial evidence suggesting the presence of preserved organic matter.

The sector-wise systematic arrangement of triangular elements - with their a-axis being oriented subparallel to parallel (Figs. 3A-C, 4A-B, 5A, D) - in bundles, indicate a biologically controlled origin (Lowenstam and Weiner, 1989) of these fabrics. The primary belemnite rostrum ultrastructure is brownish (Figs. 5A, 7A) in thin sections under transmitted light and has a relatively low optical relief. According to Ullmann et al. (2014), brownish areas in thin sections of biogenic carbonates are indicative of remnant organic matter (C_{org}). Under the cathodoluminescence- and fluorescence microscope, the primary filigree belemnite framework shows a light blue luminescence (CL) and light green fluorescence (FL). According to Wanamaker et al. (2009) and Pérez-Huerta et al. (2008), fluorescence in biominerals is triggered by organic macromolecules associated with chitin polysaccharides and proteins. Dark fluorescence patterns commonly refer to portions of the skeletal hardparts that are relatively depleted in organic matter. Bright green fluorescence patterns typify areas with increased amount of organic matter (Wanamaker et al., 2009).

488 Primary skeletal structures display darker colours in SEM-BSE images. Electron microprobe 489 analyses revealed that the biogenic skeletal calcites contain more P and Mg but less S compared to what is here considered an early diagenetic, pore-filling calcite phase. Higher concentrations of P (Fig. 490 6C) may be related to the presence of organic matter. Arguments for this have been presented by 491 492 Longinelli et al. (2002, 2003) and Gröcke et al. (2003) who found phosphate (PO₄³⁻) of presumed 493 biogenic origin being preferentially enriched along concentric growth rings. Generally, the phosphate 494 concentration of ancient belemnite rostra is variable but very low (less than 0.3%) comparable to 495 that in Recent-modern Sepia.

496 The primary filigree belemnite framework does not fluoresce under the CLFM. The factors 497 that cause fluorescence in samples studied under the CLFM are poorly constrained (Fig. 8). Naturally-498 occurring organic compounds such as proteins or polysaccharides can cause fluorescence in other 499 biogenic carbonates, including brachiopods (Pérez-Huerta et al., 2008), gastropods (Guzman et al., 500 2007), or cephalopods (Linzmeier et al., 2016). In modern brachiopods and Nautilus, portions of the 501 exoskeletons with higher amounts of intracrystalline organic matter (Clark, 1999) appear dark under 502 CLFM (-Pérez-Huerta et al., 2008; Linzmeier et al., 2016; Pérez Huerta et al., 2008). This pattern lends 503 support to the argument that dark triangular areas in BSE images represent the primary biogenic 504 skeleton and contain remnants of organic matter. Marine sediments may contain abundant humic 505 substances resulting from the degradation of marine organic matter (Nissenbaum and Kaplan, 1972) 506 and evidence has been presented that the sulfphur content of the humic substances increases with 507 degradation (Francois 1987). Concluding, it is here proposed that humic substances caused elevated 508 S concentrations in the diagenetic calcite phase that occludes the skeletal pore space (Fig. 6D) and 509 causes the CLFM fluorescence in all three wavelengths (Blyth et al., 2008; Orland et al., 2009, 2012;
510 Fig. 8).

Similarly, microtomographic data indicate the former presence of organic matter in dark triangular areas of rostra (Fig. 9) as observed in BSE images. The brightness of a carbonate observed in CT image indicates the degree of attenuation of an X-ray passing through this material (Mobilio et al., 2015). As the inner tri-radial structures of rostra appear darker in colo<u>u</u>r relative to the calcite phase fringing these structures, we suggest that the fringing phase is made of a denser calcite phase compared to the inner structure. That observation is in line with the observation of a low optical relief of these features (supplement Fig. 1A-B).

518 Thise tentative interpretation presented here with regard to belemnite rostra is arguably consistent with observations of 10-40% organic matrix in the Sepia cuttlebone (Birchall and Thomas, 519 520 1983; Florek et al., 2009). This is relevant as we suggest that the belemnite rostrum is structurally 521 similar but not homologous (Fuchs 2012) to the Sepia cuttlebone with regard to the primary intra-522 skeletal porosity. Having said this, the presence of preserved organic matter in ancient biogenic 523 carbonates particularly, intra-crystalline organic matter is not uncommon (Clark, 1999, 2005). Excess 524 carbon observed for Megateuthis has been interpreted as evidence for a former organic matrix 525 within these low-Mg calcite biominerals (Dunca et al., 2006). Similarly, Florek (2004) argued for an 526 excess of carbon in the rostra of Belemnopsis and Hibolites. Summing up: Different lines of 527 circumstantial evidence point to the presence of remnant organic matter within biominerals. These 528 data require verification or rejection via the application of spatially highly resolved geochemical data. 529 This work is presently junder progress.

530

531	5. Conclusions	Formatted: No underline, Font color: Auto
532	Ultrastructural data documented here suggest that the calcitic rostra of Mesozoic belemnites	
533	yielded 50-90% primary porosity probably filled with body fluids and/or organic matter during the life	
534	time of the animal. Porosity was distributed throughout the rostrum as opposed to being limited to	

the central apical area. The primary biogenic rostrum framework consists of triaxial branches and
tetrahedrons of variable size forming a honeycomb-like network. This structure arguably combined
mechanical stability with an energy-efficient biomineralization strategy.

538 The recognition of belemnite rostra as a highly porous structure requires a re-interpretation of 539 the function of the rostrum as counterweight to the soft body and has implications for the swimming 540 mode of belemnites. On the level of a working hypothesis, we argue that the low-porosity fabric 541 found in fossil rostra collected in outcrops worldwide is the result of a syntaxial, early diagenetic 542 cement phase that nucleated upon the surface of the biogenic framework and subsequently occluded the pore space. The possibility of gradual occlusion of skeletal porosity by remote 543 544 biomineralization during later ontogenetic stages during the life of the animal is possible but seems 545 unlikely at present.

546 If the here-presented concepts hold true, then these new findings have significant implications 547 with regard to geochemical proxy data collected from fossil belemnite rostra. Specifically, the fact 548 that rostra may consist of biogenic and abiogenic calcite phases formed at different times may 549 explain the controversially low reconstructed seawater temperatures and the uncommonly high 550 scatter of proxy data even from well-preserved rostra collected in the same stratigraphic interval. 551 This is because seawater properties of surficial water masses, the habitat of nektobenthic 552 belemnites, are recorded in the biogenic portions of the rostrum whereas the early diagenetic phase 553 reflects cooler basinal bottom or marine pore water signatures. Findings presented here form a solid 554 and well-constrained petrographic data set but one that must be verified by high-resolution 555 geochemical data of all paragenetic calcite phases observed.

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558 6. Acknowledgement

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811 Figure captions

Fig. 1A-B) Structural and ultrastructural composition of belemnite rostra. Belemnitella mucronata, thin sections photographed under crossed polarizers. A) Cross section with pseudo-uniaxial cross indicating radially arranged calcite fibres, red box refers to C. B) Longitudinal section with central apical line and radiating fibres from the centre to the margin, stippled line refers to the position of cross section shown in A. C) Idealized bundle of calcite fibres, each fibre contains a stack of tetrahedral elements. D) SEM BSD image of the tetrahedral ultrastructure of Megateuthis gigantea, dashed lines (I-IV) indicate section planes and corresponding reconstructions. Primary skeletal framework is shown in blue, yellow and green whilst early diagenetic phase is shown in white and red for the crystal boundaries. The basis of tetrahedrons points toward the centre of the belemnite rostrum and its tip towards the rostrum margin i.e. the growth direction. E) Three dimensional reconstruction of a single tetrahedron of the belemnite endoskeleton. F) Reconstruction of the complex spatial arrangement of biogenic and early diagenetic phases. Colour code in lower right. G) SEM image of a single complex tetrahedron (black line) with indication of structural elements. Primary skeletal components: br = branch, trc = triradial centre, rf = reinforcement, ic = isopachous crystallites. (full page width; bw in print, colour in pdf)

837	Fig. 2A-B) SEM BSD images of Megateuthis gigantea. A-B) Section perpendicular to the c-axes of
838	calcite fibres (section plane II in Fig. 1D-II). White stippled line indicate dissolution features (early
839	diagenetic), black stippled line indicate microstylolites -(late diagenetic). (full page width; bw in print,
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842	Fig. 2A-B) SEM BSD images of Megateuthis gigantea. A-B) Section perpendicular to the c-axes of
843	calcite fibres (section plane II in Fig. 1D-II). Blue stippled line indicate dissolution features (early
844	diagenetic), red stippled line indicate microstylolites (late diagenetic). (full page width; bw in print,
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863	Fig. 3A-F) SEM BSD images of Megateuthis gigantea. A-E) Section perpendicular to the c-axes of
864	calcite fibres. A) Triangular structures with a relatively thick outer, light grey margin of abiogenic
865	early diagenetic cement and a small darker centre with branches giving rise to reinforcement
866	structures representing the primary biogenic skeletal framework. B) Larger dark grey, organic rich
867	triangular elements belonging to the biogenic skeletal framework, partly with light grey central
868	abiogenic calcite filling of variable sizes. Branches are often short and cut off at variable distances
869	from the centre. C) Some smaller and a few larger biogenic skeletal elements with cut off branches
870	(lower arrow) and reinforcement structures (upper arrow). Note the variable expression of early
871	diagenetic crystallites with sheaths of remnant organic matter. D) Close up of larger, biogenic skeletal
872	elements and abiogenic isopachous calcites coated by remnants of organic matter within the brighter
873	outer margin. Central portion of the biogenic skeletal elements shows abiogenic crystal; arrows point
874	to dissolved branches. E) Centre of biogenic skeletal element completely filled with abiogenic bright
875	calcite leaving only a thin dark inner margin. F) Same specimen, section subparallel to the c-axes
876	showing a homogenous central portion ("trunk" – white line) of pyramidal morphology rich in
877	intracrystalline organic matter surrounded by inclined isopachous calcite crystals coated by remnants
878	of organic matter. This overall pattern results in an "*Christmas-mas tree like" structure (compare
879	with Fig. 1D). (full page width; bw in print and pdf)
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889 of the fiber bundles. Same colours represent same crystallographic orientations. In A and B angular 890 deviation from blue to red is up to 40°, in D angular deviation is 2°. A) Overview map showing the 891 bundling of fibres with identical orientation of a-axes, black frame indicates area for close up in B; 892 blue frame refers to Fig. 5A-C. B) Close up map, within one bundle blue tinted fibres are mainly 893 neighboured by other blue fibres, red tinted fibres are surrounded by red fibres. C) Close up to 894 demonstrate slight-documenting minor angular deviation within one fibre (compare with D). D) Map 895 of a single fibre with an angular deviation of 2° from blue to red, showing a slight systematic shift of 896 axes orientation. E) Pole-plots of c-axes (001) and a-axes (012) from all fibres shown in A, all c-axes 897 show nearly the same orientation while the a-axes demonstrate the bundled structure of the rostral 898 fabric which may have improved the stability of the skeletal structure. (full page width; bw in print, 890 colour in pdf) 901 902 903 904 904 905 905 906 906 907 907 908 908 909 909	888	Fig. 4A-E) Megateuthis gigantea, EBSD map with colour code in sections perpendicular to the c-axes
 bundling of fibres with identical orientation of a-axes, black frame indicates area for close up in B; blue frame refers to Fig. 5A-C. B) Close up map, within one bundle blue tinted fibres are mainly neighboured by other blue fibres, red tinted fibres are surrounded by red fibres. C) Close up te demonstrate slight documenting minor angular deviation within one fibre (compare with D). D) Map of a single fibre with an angular deviation of 2° from blue to red, showing a slight systematic shift of axes orientation. E) Pole-plots of c-axes (001) and a-axes (010) from all fibres shown in A, all c-axes show nearly the same orientation while the a-axes demonstrate the bundled structure of the rostral fabric which may have improved the stability of the skeletal structure. (full page width; bw in print, colour in pdf) goo goo<!--</td--><td>889</td><td>of the fiber bundles. Same colours represent same crystallographic orientations. In A and B angular</td>	889	of the fiber bundles. Same colours represent same crystallographic orientations. In A and B angular
blue frame refers to Fig. 5A-C. B) Close up map, within one bundle blue tinted fibres are mainly neighboured by other blue fibres, red tinted fibres are surrounded by red fibres. C) Close up te demonstrate slight- <u>documenting minor</u> angular deviation within one fibre (compare with D). D) Map of a single fibre with an angular deviation of 2° from blue to red, showing a slight systematic shift of axes orientation. E) Pole-plots of c-axes (001) and a-axes (010) from all fibres shown in A, all c-axes show nearly the same orientation while the a-axes demonstrate the bundled structure of the rostral fabric which may have improved the stability of the skeletal structure. (full page width; bw in print, colour in pdf)	890	deviation from blue to red is up to 40°, in D angular deviation is 2°. A) Overview map showing the
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913	Fig. 5A-F) Transmitted light, polarized light and cathodoluminescence. Thin section of Megateuthis
914	gigantea, A-C refer to the blue frame in Fig. 4A, D-F are close ups (black frame in A) A and D) TL
915	image perpendicular to the c-axes of calcite fibres, filigree biogenic skeletal framework is indicated
916	by the dark tinted structures, primary porosity is represented by the abiogenic translucent calcites.
917	Note banded distribution of calcite fibre domains relating to larger and smaller organic-rich biogenic
918	skeletal elements, single fibres may contain a central portion of transparent calcite of varying size,
919	bundling of adjacent calcite fibres is indicated by the same orientation of the triangles. B and E)
920	Uniform extinction (orientation) of adjacent fibres under crossed polarizers. C and F) CL of abiogenic
921	calcite portions show a dark blue, intrinsic luminescence (pure stoichiometric calcite), CL of the
922	biogenic skeletal framework show light blue luminescence. (full page width; bw in print, colour in
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938	Fig. 6) Electron microprobe data for Megateuthis gigantea. A) Overview BSE map B) Shows higher
939	Mg concentrations within the triangular areas dark in BSE images and a lower Mg concentration in
940	the surrounding area bright in BSE images. C) Shows higher P concentrations within the triangular
941	areas dark in BSE images and a lower P concentration in the surrounding area bright in BSE images.
942	D) Shows lower S concentrations within the triangular areas dark in BSE images and higher S
943	concentrations in the surrounding region bright in BSE images. (full page width; bw in print, colour
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964	Fig. 7) Fluorescence microscope images for Megateuthis gigantea. A) Transmitted light shows
965	brownish triangular structures, rich in organic matter and dark in BSE images, partly with central
966	translucent areas (compare with Fig. 1D section plane IV, Fig. 3D, E, 5A, D). B) Shows brighter
967	fluorescent triangular area compared to the in transmitted light translucent calcite. (full page width;
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990	Fig. 8) Confocal laser fluorescence microscopy images of Megateuthis gigantea. A) CLFM images
991	showing fluorescence in far-red light (λ = 664 to 696 nm). B) CLFM images showing fluorescence in
992	visible green light (λ = 505 to 539 nm). C and D) CLFM images showing fluorescence in visible red light
993	(λ = 589 to 621 nm). Triangular structures visible in other imaging techniques (Fig. 3, 5-7) do not
994	fluoresce as brightly as cracks (B) or early diagenetic calcite (Fig. 3) separating the triangles (A, B, C,
995	D). D) Higher magnification shows some brighter fluorescing calcite between the triangular elements
996	dark in BSE images (Fig. 1, 3). Brightly fluorescent early diagenetic calcite separating triangles also
997	containsis enriched in-elevated S (Fig. 6),. which may support inclusion of S rich humic substances as
998	the cause of fluorescence. (full page width; bw in print, colour in pdf)
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1016	Fig. 9) Three-dimensional visualization of the filigree biogenic framework. Synchrotron radiation-
1017	based tomographic visualization of a sub-volume of the rostrum of Megateuthis gigantea. Specimen
1018	was scanned with an isotropic voxel size of 0.74 $\mu m.$ A) Multi-planar image of a sub-domain of the
1019	original dataset with dimensions of 447x592x663 voxels, triangular elements dark in BSE images
1020	appear here as dark elements due to reduced densities. B-D) Volumetric renderings of the same sub-
1021	domain with variable rendering settings. (full page width; bw in print, colour in pdf)
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1061	Fig. 1) Examples of the filigree framework from other belemnite species. A-B) Gonioteuthis
1062	quadrata, A) shows the Becke line outside of the triangular area with a relatively larger distance
1063	between the sample and objective. B) Shows-Image shows the Becke line within the triangular area
1064	while the distance between the sample and the objective was reduced, accordingly the triangular
1065	area (dark in BSE;{Fig. 3}) has a lower optical relief. C-D) Belemnitella mucronata, C) thin section
1066	under polarized light, D) same area under CL showing microfractures filled with Mn-rich calcite
1067	tracing the outline of triangular elements (encircled). (full page width; bw in print, colour in pdf)

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1081	Fig. 2) SEM BSD images of Megateuthis gigantea. A-F) Section plane parallel to the c-axes of calcite
1082	fibres. A-C) stepwise enlargement of a particular area. D-F) stepwise enlargement of a particular
1083	area. A and D give the impression of a concentric arrangement of distinct darker and brighter layers
1084	(black frames enlarged in B and E), arrow in D point to an organic rich layer (laminae obscura sensu
1085	Müller-Stoll, (1936)). B and E) Allow the recognition of single darker structures of tetrahedral
1086	morphology with their tips pointing towards the outer margin of the belemnite rostrum, i.e. the
1087	growth direction (black frames enlarged in C and F). C and F) show the intricate framework of
1088	biogenic (dark) and abiogenic (light) carbonate phase within the rostrum, larger dark grey, triangular
1089	elements belonging to the biogenic skeletal framework, partly with light grey central abiogenic
1090	calcite filling of variable size. (full page width; bw in print, colour in pdf)

1 Evidence for a composite organic-inorganic fabric of belemnite rostra:

2 Implications for palaeoceanography and palaeoecology

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- 27 Abstract
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29 Carbonate skeletons of fossil marine organisms are widely used to reconstruct palaeoceanographic parameters. Specifically, the geochemistry of Jurassic and Cretaceous belemnite rostra is traditionally 30 31 interpreted to represent near sea-surface seawater properties. More recently, an increasing number of workers, have reported significant scatter in geochemical data (e.g., δ^{18} O, δ^{13} C, element/Ca ratio) 32 33 when comparing rostra from the same stratigraphic level or within a single belemnite rostrum. This 34 scatter is not explained by differential diagenetic overprint alone. Here we report petrographic 35 evidence on the primary ultrastructure of rostra of Megateuthis (Middle Jurassic) and Belemnitella 36 and Gonioteuthis (Late Cretaceous). The biogenic ultrastructure consists of a filigree framework of 37 triaxial branches and tetrahedrons of variable size forming a honeycomb-like network. Data 38 presented here suggest that these rostra yielded as much as 50 to 90% primary pore space. On the 39 level of a working hypothesis - and in analogy with modern cephalopods - we propose that the pore 40 space was formerly filled with body fluid and/or organic compounds during the life time of these 41 organisms. Intra-rostral porosity was post mortem occluded by earliest diagenetic isopachous calcite cements of a non-biogenic origin. These may have been precipitated due to increased alkalinity 42 related to the decay of organic matter. If this holds true, then the resulting fabric represents a 43 44 composite biogenic/abiogenic structure. In order to optically separate the two calcite phases forming 45 a single calcite fibre, we employed a wide range of state-of-the-art analytical tools to thin sections 46 and ultra-thin sections of well-preserved specimens. Pending a verification of these well-supported ultrastructural data by means of high-resolution geochemical analyses from biogenic and abiogenic 47 48 phases, we suggest that these findings have significance for those using belemnite rostra as archives 49 of their palaeoenvironment.

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51 Keywords: belemnite, ultrastructure, carbonate archive, diagenesis, Jurassic-Cretaceous

53 **1.** Introduction

The ultrastructure of recent biogenic carbonates is of great interest for those concerned with 54 biomineralization research in general (Weiner and Addadi, 2011; Goetz et al., 2011), those studying 55 56 the primary biogenic skeletal structures of fossil skeletal hardparts (e.g., Coronado et al., 2013), and 57 for palaeoceanographers exploring these materials for their bearing on past climate dynamics (Saelen, 1989; Cochran et al., 2003; Parkinson et al., 2005; Jarvis et al., 2015; Immenhauser et al., 58 59 2016). Whereas the tests of planktonic and benthic foraminifera and coccoliths are important 60 archives of open marine environments throughout the Cenozoic and beyond (e.g., Zachos et al., 2001), much of what is known about Cretaceous and Jurassic palaeoceanography has been deduced 61 62 from the geochemical archive of the calcareous rostra of extinct cephalopods, specifically belemnites 63 (Dutton et al., 2007). Reasons for the wide use of these archive materials include their abundance in the fossil record, the diagenetically stable low-Mg calcite mineralogy of rostra (Veizer, 1974; Saelen, 64 65 1989), and the broad palaeobiogeographic distribution of belemnites in the marine Boreal and Tethyan realms, (e.g., Urey et al., 1951; Voigt et al., 2003; Wierzbowski, 2004; McArthur et al., 2007; 66 67 Dutton, 2007; Price and Page, 2008; Wierzbowski and Joachimski, 2009; Price et al., 2009 2011; Li et 68 al., 2012, 2013). As with all biogenic carbonates, however, these archives undergo post mortem 69 diagenetic alteration, representing a major obstacle in carbonate research (Swart, 2015).

70 Characteristic geochemical patterns and fabrics in biogenic carbonate hardparts have been used 71 to test for example molluscs (e.g., Cochran et al., 2003, Sessa et al., 2015, Immenhauser et al., 2016), 72 brachiopods (Parkinson et al., 2005), or foraminifera (e.g., Huber and Hodell, 1996; Kozdon et al., 73 2011) for evidence of diagenetic alteration. In the case of ammonites, the degree of preservation of 74 nacre tablets provides evidence with regard to the preservation of these exoskeletons, to name one 75 example (e.g., Cochran et al., 2010). With regard to belemnite rostra, the observation of an intact 76 fibrous microfabrics in thin sections and polished rock surfaces in combination with 77 cathodoluminescence is commonly used to identify well-preserved belemnite rostra (Rosales et al., 78 2001). In contrast, cloudy areas, exfoliation, fractures, stylolites, or boring traces are interpreted as

evidence for post mortem alteration (Saelen, 1989; Li, 2011; Benito and Reolid, 2012). A rigorous
discussion of screening techniques, including trace-element concentrations, and isotopic ratios
applied to biogenic low-Mg calcite macrofossils, has been provided by Ullmann and Korte (2015).
Similarly, a "best practice" approach for the interpretation of mollusc and brachiopod carbonate
archives was presented by Immenhauser et al. (2016).

84 The ultrastructure of belemnite rostra was first studied by Müller-Stoll (1936). This author 85 described organic-rich (laminae obscura) and carbonate-rich (laminae pellucidae) concentric growth 86 rings. Both of these were later shown to be calcitic but differ due to variable amounts of occluded 87 organic matter (Saelen, 1989). Growth rings are made up by fibrous calcite crystals radiating from a central zone called apical line (Richter et al., 2011) forming what seems to be a low-porosity fabric 88 89 (Saelen, 1989 and references therein; Fig. 1A-C). Single fibres (= radial structures of Saelen, 1989) can 90 be arranged in bundles traversing the concentric growth layers. Each fibre thickens outwards and 91 shows a sub-fibrous framework potentially first proposed - but not further explored - by Saelen 92 (1989, Fig. 15a).

93 Belemnites are traditionally considered to secrete their endoskeleton in oxygen isotope equilibrium with ambient seawater (e.g., Anderson et al., 1994; Price and Sellwood, 1997; Voigt et 94 al., 2003; Price et al., 2009; Wierzbowski and Joachimski, 2007, 2009). The main argument brought 95 forward is commonly the presence of what are considered cyclical oxygen isotope patterns 96 97 interpreted as seasonal seawater temperature variations (Urey et al., 1951). The PeeDee belemnite 98 used in Urey's study was considered as well-preserved based on the compact fabric and the optical 99 features of the calcite crystals (but see Li, 2011 for discussion). The assumption of equilibrium 100 precipitation was further supported by data sets from recent cephalopods including *Nautilus*, *Sepia*, and Spirula precipitating their skeletal hardparts in near-equilibrium with $\delta^{18}O_{seawater}$ (Lukeneder et 101 102 al., 2010). In contrast to this traditional view, there is an increasing amount of evidence suggesting 103 that belemnite rostra are problematic archives of their palaeoenvironment (see Immenhauser et al., 104 2016 for detailed discussion). For example, Price et al. (2015) reported an offset of about 5°C

105 between the aragonitic phragmocone and the calcitic rostrum of a single specimen of 106 *Cylindroteuthis*. This offset was regarded as vital effect but it remains unclear whether the 107 temperatures derived from the aragonite are too warm or from the calcite too cool. Similarly, high 108 intra-rostral variability of elemental (Ca, Mn, Mg, Fe, Sr) and isotopic (δ^{18} O, δ^{13} C) composition, the 109 latter with a scatter of up to 2‰, has been reported from belemnite rostra lacking evidence for 110 diagenetic alteration (Podlaha et al., 1998).

111 The significant discrepancy of these data sets forms a strong motivation for a reconsideration of 112 belemnite rostra as archives of their palaeoenvironment. Here, we report data from a wide set of 113 state-of-the-art analytical infrastructure applied to thin- and ultra-thin sections of exceptionally wellpreserved Jurassic and Cretaceous belemnite rostra. The following aims guided this paper: First, to 114 115 present well-constrained petrographic evidence for the complex primary biogenic framework of 116 these rostra; second, to document evidence that points to the highly porous nature of this biogenic 117 framework; third, to assess the relative proportion between primary skeleton and porosity; fourth, to 118 discuss the timing and nature of the pore-filling calcites phase. Evidence reported here has 119 significance for the interpretation of proxy data from ancient belemnite rostra and forms the 120 foundation of a detailed, high-resolution geochemical study that will be in the focus of forthcoming 121 work.

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123 2. Materials and Methods

124 2.1. Belemnites

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Thin sections (30 μm) and ultra-thin sections (< 10 μm) of two well-preserved orthorostra of *Megateuthis gigantea* (Schlotheim, 1820) from the Middle Jurassic and one specimen of *Belemnitella mucronata* (Schlotheim, 1813) and *Gonioteuthis quadrata* (Blainville, 1827) from the Upper Cretaceous were studied. Specimens of *Megateuthis* were collected in Bajocian marly limestone deposits from southern Germany. *Belemnitella* rostra are from the late early Campanian (*mucronata*) Zone), and *Gonioteuthis* from the early Campanian *lingua/quadrata – gracilis/mucronata* Zone from northwestern Germany. *Gonioteuthis* and *Belemnitella* rostra were embedded in calcareous (65-90%) epicontinental shelf deposits of the Misburg Formation (Niebuhr, 1995; Niebuhr et al., 2007). Specimens did not display evidence for exfoliation nor did they show boring traces of the surfaces of rostra. Specimens were sectioned along their long axis in a marginal position and perpendicular to the c-axis of their calcite fibres. One section cuts the rostrum of *Megateuthis* perpendicular to the long axis.

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139 2.2. Methods

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Surfaces of thin sections were chemo-mechanically etched using colloidal silica (OP-S) for 5-15 minutes to reduce surface irregularities on an atomic scale (Massonne and Neuser, 2005) and coated by a thin carbon layer. All coated thin sections have been studied under a high-resolution field emission scanning electron microscope (HR-FESEM) type LEO/ZEISS 1530 Gemini using a backscatter detector (BSD) at the Ruhr-Universität Bochum, Germany (Figs. 1-3).

146 Crystallographic orientation of belemnite calcite fibres was determined by electron 147 backscattered diffraction (EBSD; Nordlys, OXFORD Instruments). The data acquisition and analysis 148 was performed using the software packages AZtec and Channel 5 by Oxford Instruments (Fig. 4). The 149 scanning electron microscope (SEM) was operated at beam energy of 20 kV, an aperture of 60 μ m, a 150 working distance of 25 mm and a tilt angle of 70°. Thin sections were mapped at Bochum University 151 in the high-resolution mode using a grid matrix (1149x748 points) at a step width of 2.017 µm (Fig. 152 4A) and (543x266 points) at a step width of 1.652 μ m (Fig. 4D) and for the single calcite fibre 50 μ m 153 to reduce artefacts and increase the reliability of the data. The orientations of the crystals in the 154 individual maps were visualized using a rainbow colour coding ranging from blue over green and yellow to orange and red, where identical colours indicate identical crystal axis orientations. For 155 156 visualizing the weak angular deviations in the crystal lattice in a single calcite fibre, we applied an

angular resolution of 2 degrees for the complete rainbow colour range. In addition, orientations of
the measured crystallographic axes were plotted into the lower hemisphere of a Schmidt net (Fig.
4E).

160 Rostra were further investigated under a cathodoluminescence microscope type HC1-LM by 161 Lumic equipped with a hot cathode (Neuser et al., 1996) and a digital camera system (DP73 by 162 Olympus) for recording digital images at Bochum. Beam energy of 14kV and a beam current density 163 between 5 and 10 μ A/mm² were generally used for the CL-measurements. Integration times for CL-164 spectra were commonly between 10 and 60 seconds (Fig. 5).

165X-ray element distribution maps were acquired using a Cameca SX5FE field emission electron166microprobe at Bochum. The acceleration voltage was 15 keV with a probe current of about 80 nA and167a fully focused beam. The intensity of S Kα1 was recorded simultaneously on two wavelength168dispersive spectrometers equipped with LPET and PET analyzing crystals. The Mg Kα1 line was also169measured on two spectrometers (LTAP and TAP crystals), whereas P Kα1 was measured on a single170spectrometer equipped with a LPET crystal. The images were acquired in continuous stage scan171mode. They have a resolution of 2048 x 1536 pixel, and the dwell time was 17 ms per pixel (Fig. 6).

172 To test for the distribution of organic matter within the belemnite rostra, thin sections were 173 studied under a fluorescence microscope (Leica DM4500P) equipped with a mercury short-arc 174 reflector lamp coupled with a Leica EL6000 compact light source. We used the blue light filter set 175 producing bright green fluorescence images (filter set I3 for blue light excitation: excitation 450-490 176 nm, emission 515 nm, voltage 100-240 VAC and 50-60 Hz frequency; Fig. 7). Fluorescence microscopy in the manner applied here is an optical tool to qualitatively document the spatial distribution of 177 178 organic matter in shells. Fluorescence reflects organic matter and less commonly crystal lattice 179 defects and solid inclusions in crystals. Further details of this method have been presented by Wanamaker et al. (2009) and Ritter et al. (subm.). 180

181 Confocal laser fluorescence microscopy (CLFM) images on an uncoated *Megateuthis* mount 182 were made using a Bio-Rad MRC-1024 scanning confocal microscope at the W. M. Keck Laboratory

for Biological Imaging at UW-Madison (Fig. 8). The microscope was operated with a 40 mW laser at wavelengths of 488 nm, 568 nm, and 647 nm. All three wavelengths were simultaneously rastered across the sample. Naturally occurring compounds within the sample caused fluorescence at multiple wavelengths. Images were collected through the following three emission filters: visible green light (λ = 505 to 539 nm); visible red light (λ = 589 to 621 nm); and far-red light (λ = 664 to 696 nm).

188 In order to analyze the three dimensional filigree framework, synchrotron radiation based 189 micro-computed tomography was applied (Fig. 9). Data were collected at the bending magnet beam 190 line 2-BM at the Advanced Photon Source, Argonne National Laboratory, USA. A double multilayer 191 monochromator of 1.5% band-width provided 27.2 KeV X-rays. Images were collected in transmission 192 mode by a CCD camera behind the sample in the hutch configuration. The sample-detector distance 193 was set to 300 mm to collect quantitative phase contrast data. 1440 projections were acquired while 194 the sample was rotated over 180° in steps of 0.125°. A microtomographic data set with a size of 2048 195 x 2048 x 1948 voxels was reconstructed using a phase retrieval algorithm (Mokso et al., 2013).

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198 3. Results

3.1 Optical-, cathodoluminescence-, fluorescence-, and confocal laser fluorescence microscopy (TL, CL,
FL, CFLM)

Under transmitted light (TL), thin sections of rostra cut perpendicular to the c-axis of calcite fibres reveal a banded distribution of calcite fibres containing brownish triangular elements of variable size (Fig. 5A, D). Some of the larger fibres contain a triangular, organic-rich centre, with its innermost domains occluded by translucent calcite. Bundling of neighbouring calcite fibres is indicated by their subparallel orientation of their a-axis (Fig. 5D). A uniform extinction (orientation) pattern of adjacent fibres under crossed polarizers is observed (Fig. 5B, E) and is in agreement with EBSD data (Fig. 4A). Individual fibres display an undulatory extinction (converging or diverging c-axes) 208 under crossed polarizers. Calcite occluding space between organic-rich elements and fibre 209 reinforcement is translucent and contains little or no organic matter (Fig. 5D). Brownish, triangular 210 elements have a relatively low optical relief, compared to the surrounding translucent calcite with a 211 relatively higher optical relief (supplement Fig. 1A-B).

Three different luminescence patterns are observed under the cathodoluminescence microscope (CL): (i) A light blue luminescence of the brownish, triangular elements (Fig. 5C, F); (ii) dark blue, intrinsic luminescence of the translucent phase; (iii) locally, orange to red luminescent fractures and microstylolites are observed. In rare cases, fractures retrace the triangular outline of a fibre (Fig. 5C, F, supplement figure 1D).

Brownish triangular areas under transmitted light display light green fluorescence while the translucent areas show a dark green fluorescence under the fluorescence microscope (FL). Microfractures are darker under transmitted light compared to the triangular elements and show a light green fluorescence (Fig. 7A-B).

Different fluorescence patterns under the CLFM reveals calcite domains visible in BSE (and other) imaging techniques (Figs. 3, 5-7). Filled cracks fluoresce brightly in green and red wavelengths (Fig. 8B-C). Domains that are dark in BSE do not fluoresce in CLFM (Fig. 8A-D). Bright domains in BSE fluoresce brightly in CLFM at all wavelengths observed (Fig. 8A-D). Higher magnification reveals brighter fluorescence between adjacent domains that appear bright in BSE (Fig. 8D).

226

227 3.2 Scanning-electron microscopy equipped with a backscatter detector (SEM BSD)

Images collected with the scanning electron microscope equipped with a backscatter detector present important evidence for the presence of two calcite phases (dark and bright) of different chemical compositions building the rostra studied. The brownish triangular areas under transmitted light are dark in BSE. Thin sections cut perpendicular to the c-axis of the radiaxal fibrous calcite (Fig. 232 1D I-IV) reveal a complex framework, dark in BSE, surrounded by relatively brighter calcite (Figs. 1-3). Cross sections of the majority of fibres are polygonal or, less often, honeycomb shaped with a tri-233 234 radial (120°) symmetry representing an ultrastructure that has not been described in previous studies 235 (Figs. 1C-G, 3A-E). Individual fibre diameters vary between 10-80 µm. Calcite fibres terminate at 236 individual concentric growth layers that also form the nucleation site for the overlying, next fibre 237 generation, displaying increasing thicknesses and occurring in increasing numbers towards the outer 238 portions of the rostrum (Fig. 1A-D). Brighter area in BSE are more likely to be composed of near-239 stoichiometric CaCO₃ with higher average atomic masses (mainly Ca), whereas excess C, P, Mg, or S 240 will lower the average atomic mass. Accordingly, areas with decreasing amount of Ca and increasing 241 amounts of C, P, Mg or S are darker.

242 Four section planes are presented to describe the intricate bio-composite mineral present 243 within a single fibre. The following description distinguishes between the darker framework (i), i.e., 244 micrometre-thick branches terminating in wall-like reinforcements and (ii) tri-radial central portions; 245 and brighter fabric (iii) consisting of calcite crystals forming isopachous cement layers with individual 246 crystallites coated by submicrometre thick layers of matter darker in BSE (Fig. 1G). In all sections, we 247 observed isopachous calcite crystals oriented perpendicular to the inner dark walls (Fig. 3A-E). 248 Section plane I consists of three simple, dark in BSE branches of variable lengths. Often, these 249 branches exhibit reinforcements of variable lengths and thicknesses (about 1 micrometre) and 250 increasing widths towards neighbouring branches (Figs. 1D-I, 3A). Conversely, reinforcements being 251 connected to neighboring triangles are rarely observed. Section type II is characterized by an 252 enlarged, dark in BSE central element of variable diameter (2-30 μm, Figs. 1D-II, 3C). Some of the 253 larger central elements display an inner zone with additional tri-radial structures comprising of smaller, isopachous, brighter in BSE area calcite crystals rotated by 60° relative to outer branches 254 (Figs. 1D-III, 3D; section type III). Section type IV represents the most complex fabric. Here, the inner 255 256 portions of larger central elements display alternating darker and brighter in BSE areas (Figs. 1D-IV,

3E). Generally, enlarged central elements correlate with a shortening of corresponding, darkerbranches.

259 Thin sections cut parallel to the c-axis of the radiaxial fibrous calcite reveal an framework dark in 260 BSE of triangular shaped elements with their tips pointing towards the outer margin of the rostrum 261 (Fig. 1C-D, supplement Fig. 2). Lighter and darker areas within a single calcite fibre show their 262 composite nature. Lighter and darker areas within concentric growth rings, as visible in transmitted 263 light, depend on the number and size of these elements. Higher magnification reveals a homogenous central portion ("trunk" - white line) of pyramidal morphology dark in BSE surrounded by inclined 264 265 isopachous calcite crystals coated by material dark in BSE. This overall pattern results in a "Christmas 266 tree like" structure (Fig. 3F). The dark framework forms the substratum upon which an isopachous, 267 translucent calcite phase nucleated (Fig. 1D, G 2-3). The boundary surface between the framework 268 dark in BSE and the surrounding carbonates brighter in BSE is, in places, corroded and uneven (Fig. 269 2).

270

271 3.3 Electron microprobe analysis (EMPA)

272 The contrasting chemical composition of the two different calcite phases building these rostra is 273 also revealed by EMP data. Detailed WDS scans show that the strongest variability is displayed by 274 Mg, P, and S, whereas other elements are almost constant or present in amounts close to the 275 detection limit (e.g., SrO = 0.1-0.2 wt.%; FeO = <0.1 wt.%; MnO < 0.5 wt.%. X-ray element distribution 276 maps show a generally low concentration of the elements Mg (MgO = 0.3-0.4 wt.%), P (P₂O₅ = 0.1-0.2277 wt.%), and S (SO₃ = 0.20-0.50 wt.%). The Mg- and P-content is slightly elevated in the darker, 278 triangular areas in BSE images (MgO up to 0.6 wt.%; P_2O_5 up to 0.25 wt.%) compared to the relatively 279 brighter areas in BSE images in the centre of larger tetrahedrons and their vicinity. Lowest Mg 280 concentrations were found along microfractures that display bright luminescence colours (MgO < 0.2 281 wt.%; Fig. 6B). Sulfur has a higher concentration in the brighter areas in BSE images (SO₃ up to 0.65 wt.%) and lower concentrations in the darker areas in BSE images (SO₃ = 0.20-0.50 wt.%; Fig. 6D). For better comparison with BSE images from other samples and with the element distribution, BSE data were obtained for the same field of view (Fig. 6A).

285

286 3.4 Electron Backscattered Diffraction imaging (EBSD)

287 Electron backscattered diffraction was used to document the biological controlled (sensu 288 Lowenstam and Weiner, 1989; Dupraz et al., 2009; Weiner and Addadi, 2011) formation of what is 289 here assumed to represent the primary biogenic skeletal carbonate. EBSD reveals a very low 290 variation of c-axes orientation of calcite fibres (Fig. 4A). The sub-parallel orientation of the c-axes 291 {001} is documented by their close fitting in the lower hemisphere of a Schmidt net (Fig. 4E, left plot). 292 A different characteristic is found for the crystallographic orientation of the a-axes {010} of calcite 293 fibres (Fig. 4E, right plot). All a-axes are aligned along a great circle whilst the colour code of the 294 according fibres is indicative of an arrangement in bundles. Adjacent fibres (10 to 100 fibres) share a 295 similar - if not identical - orientation (Fig. 4A-B). The majority of fibre bundles are characterized by 296 blue, green and red colour coding. Specifically, the spatial orientation of individual a-axes within a 297 single fibre bundle deviates by 20° or less from the bundle mean value. Variation in the orientation of the c-axis of a single calcite fibre in the range of 1-3° was observed (Fig. 4D). As the angular 298 299 resolution from blue to red rainbow colour shown in EBSD maps (Fig. 4A-B) has a resolution of 50°, 300 minor angular deviations (<1°) are not visible in Fig. 4A-B.

301

302

3.5 Synchrotron radiation based micro-computed tomography (SRµCT)

303Tetrahedral structures are visible in three dimensions in the tomographic dataset despite the304high signal to noise ratio and artefacts. There is a distinct difference in X-ray attenuation between the

305 inner tri-radial elements and the surrounding calcite.

306

- 307 4. Interpretation and Discussion
- 308 *4.1 Reconstruction of the primary belemnite rostrum ultrastructure*
- 309

Data presented here document a repeated pattern of triangular elements that seem to be originally connected, building a concentric layered, highly complex, and porous framework. The space between the former skeletal elements is interpreted as pore space probably filled by body fluids or organic material during the life time of these organisms. The latter assumption is based on analogous observations in the porous endoskeletons of recent *Sepia* (Sherrard, 2000; Guerra, 2006). Referring to the high porosity observed in both, the belemnite rostrum and the sepiid cuttlebone, it is important to note that these structures are not homologous (Fuchs 2012).

317 Accordingly, the belemnite rostra originally consisted of an organic-rich biogenic framework of 318 calcitic tri-radial tetrahedrons (triangular pyramids arranged perpendicular to the concentric layers 319 during lifetime). Tetrahedrons are elongated along their c-axis with the tip of the pyramid pointing 320 towards the precipitation site i.e., in growth direction towards the outer margin of the rostrum (Fig. 321 1C-D). Individual branches protruding from the edges of the tetrahedrons possess outer 322 reinforcements acting as stabilizers. The central portion of the tetrahedrons may yield a channel-like 323 cavity (Fig. 1D III-IV). Organic membranes at which nucleation of the tetrahedrons may have started 324 and stopped are not preserved. However, it seems likely that such membranes were present because 325 the bases of the tetrahedrons follow a concentric layer. Structurally, the bundling of the tri-radial 326 elements forms a simple honeycomb-like framework in which the terminations of branches of 327 adjacent elements are connected. The mechanical stability of the honeycomb structure is enhanced 328 by reinforcement walls (Figs. 1D, G, 3C).

Based on the variable dimensions of skeletal elements seen in thin sections, intra-rostral pore space was visually estimated to range between 50-90% of the total rostrum (Fig. 3A: 90% pore space and 10% skeletal elements; Fig. 3B; 50% pore space and 50% skeletal elements). In this context, the

332 observation that pore space is not limited to the apical line region but is present across the bulk 333 rostrum is important. The secretion of a porous, but mechanically stable orthorostrum is probably 334 best seen in the context of a considerable reduction of energy and building material required to form 335 this structure compared to a massive endoskeleton (Sherrard, 2000). Strict biological control, i.e. in 336 the presence of organic templates (Chateigner et al., 2000; Richter et al., 2011), over the 337 precipitation of primary skeletal elements is demonstrated by a systematic arrangement of adjacent 338 elements into bundles of similar or identical orientation of their crystallographic a-axes. This 339 configuration results in a much higher mechanical load capacity and torsion stiffness of the 340 framework of the rostrum.

Alternating concentric Ca-rich (brighter; laminae obscurae) and C-rich layers (darker; laminae 341 342 pellucidae) of Müller-Stoll (1936) are related with the banded distribution of larger and smaller 343 organic-rich triangular elements of the primary skeletal structure of the belemnite orthorostrum (Fig. 344 1D, supplement Fig. 2). These layers potentially indicate differences in calcite precipitation rates. 345 Based on petrographic and ultrastructural evidence, the Mg and Ca elemental concentrations of the 346 rostrum, the lack of secondary micro-dolomite, and the absence of blotchy luminescence, the widely held assumption of a primary low-Mg calcite mineralogy for the studied belemnite rostra is 347 348 confirmed (see discussion in Richter et al., 2003).

349

350 *4.2 Early and late diagenetic processes*

351

The presence of a highly porous primary rostrum architecture during the life time of the belemnite organism as proposed here, is in contrast to the dense fabrics observed in rostra collected in Mesozoic sedimentary successions (Fig. 1A-B). Hence, the diagenetic pathway from porous to dense fabrics deserves attention. The marine diagenetic alteration of biominerals is initiated directly after the death of a carbonate-secreting marine organism when metabolic processes come to a halt. At this early stage, organic matter outside of the orthorostrum (i.e., the belemnite animal itself) and

in the pore space of rostra and between biominerals decomposes (Saelen, 1989), triggering a series of complex bio-chemical processes. Essentially, decomposition of organic matter is mediated by microbial activity, and given the abundance of marine microbial life, there is no reason to assume that this would have been different in the case examples studied here.

362 Microbial metabolic products, the presence of microbial "mucus" (extracellular polymeric 363 substance) and charged surfaces represented by microbial bodies influence the micro-environment 364 in intra-rostrum pore space by altering the balance between more reduced and more oxidized forms 365 of carbon as previously summarized under the term "alkalinity engine" (Dupraz et al., 2009). 366 Specifically, Visscher and Stolz (2005) subdivided microbial species into 5-7 groups ("guilds") having a 367 similar metabolism. Some of these promote carbonate precipitation (e.g., cyanobacteria, sulphate 368 reducers), whilst others favour dissolution (e.g., aerobic heterotrophs, fermenters). Dupraz et al. 369 (2009) documented that the balance of microbial metabolic activities directly influences carbonate 370 precipitation or, vice versa, dissolution. Initial microbial decomposition of organic matter may result 371 in the production of organic acids lowering the pH-values in the pore space (Berner et al., 1978), 372 enhancing intra-orthorostrum dissolution of biominerals (Fig. 2).

373 As soon as the bulk of organic matter is decomposed, alkalinity is increased due to microbial 374 metabolic products enhancing the precipitation of carbonate minerals (intrinsic 375 organomineralization) leading - in the view of the authors - to the formation of the translucent, 376 isopachous calcite crystals that occlude primary pore space in rostra (Fig. 2). Obviously, any 377 assumption regarding microbial processes in these ancient carbonates must remain speculative. 378 Nevertheless, following the basic concepts laid out in Dupraz et al. (2009), we tentatively assume 379 that a first phase of decomposition was dominated by aerobic heterotrophy, sulphide oxidation, and 380 fermentation decreasing the saturation index and resulting in the corrosion of skeletal elements in 381 the belemnite rostrum. Evidence for this comes from micro-corrosion features at the outer surfaces 382 of the primary skeletal framework forming the substratum upon which the pore-filling, secondary 383 calcite phase nucleates (Fig. 2).

384 The nucleation and precipitation of the secondary calcite phase (Fig. 1D, G, 2-3) was possibly dominated by sulphate reduction increasing the saturation index and hence favoring precipitation of 385 386 CaCO₃. During crystals growth, remnants of belemnite organic matter combined with microbial 387 mucus were likely trapped between single crystals or at the growth front of crystals and delineate 388 crystal boundaries. Specifically, the growth of fibrous calcitic crystals will proceed as long as growth 389 rate, fluid supersaturation, and temperature are low enough to discourage spontaneous nucleation 390 (Oti et al., 1989). Similar processes have been described from extant echinoderm endoskeletons that 391 share a comparable amount of primary porosity with the described belemnite orthorostra (Richter et 392 al., 2003).

393 During this early stage of diagenetic evolution, the rostra most likely preserved their original 394 morphology due to the biogenic calcite framework and abiogenic calcite progressively occluding 395 former pore space preventing, in combination, a collapse of the rostra. The directly comparable, dark 396 blue intrinsic luminescence of the translucent secondary outer calcite layer and the calcite infill of 397 central pore space of many of the larger triangles (Figs. 3D-E, 5A, D, 6) suggest that both phases 398 precipitated from one fluid, or different fluids with near-identical geochemistry. This is, of course, 399 within the limitations of the geochemical resolution of the cathodoluminescence method and with reference to elements that affect luminescence patterns (e.g., Mn²⁺, Fe²⁺ and REE; see discussion in 400 401 Barbin 1991, 1993; Ritter et al., 2015). Conversely, the organic matter-rich triangles display a 402 moderately brighter blue luminescence pattern (Fig. 5C, F). This feature is best explained by calcite lattice deformation due to the incorporation of organic matter into the crystal lattice 403 404 (intracrystalline) and is not necessarily indicative of a different geochemical composition. This 405 concept is supported by x-ray diffractometry (Richter et al., 2011) documenting that the fibrous 406 fabric of the belemnite rostra, lacking late diagenetic Mn-rich cements, is composed of 407 stoichiometric or near-stoichiometric calcites (d (104) = 3.030 to 3.035 Å).

408 A late diagenetic (burial) stage of rostra is documented by dissolution and compaction features 409 as indicated by microfractures and microstylolites (Figs. 2, 5C, F; Rosales et al., 2004a, b). The

circulation of Mn²⁺-rich fluids caused the precipitation of a late calcite phase that occludes fractures
and fissures. In some cases this late phase yields a bright luminescence and is zigzag or triangular
shaped, tracing the morphology of the triangular biominerals (Fig. 5C).

- 413
- 414 4.3 In

4.3 Implications for the function of the rostrum

415

It is generally accepted that the belemnite rostrum acts as a counterweight to the soft body.
Based on the observation of a high primary porosity (see also Spaeth, 1971, 1973, 1975; Ullmann et al., 2015), this interpretation requires renewed consideration. Rostra are commonly considered to have the same density as inorganic calcite crystals, ranging from 2.5-2.7 g/cm³.

420 Sepia, the closest living relative of the extinct belemnites, incorporates a total of 10-40% organic 421 matter in its cuttlebone and comparably high amounts of intracrystalline organics were observed for 422 the biogenic belemnite orthorostrum. Accordingly, ignoring open pore space, a reduced density of the biogenic belemnite calcite of about 2.4 g/cm³ (10% organic) – 2.0 g/cm³ (40% organic) results. 423 Assuming that liquid or extracrystalline organic matter (1.03 g/cm³) - with density comparable to that 424 425 of seawater (1.026 g/cm³) - filled up the pore space of living belemnite rostra, an overall density of 426 the rostrum ranging between roughly 1.7-1.1 g/cm³ (mean 1.4 g/cm³) is tentatively assumed on the 427 level of a working hypothesis. If these assumptions hold true, then the belemnite rostrum had a 428 cumulative density that is significantly lower than that of stoichiometric calcite. Therefore, questions 429 regarding the locomotion of belemnites result.

A possible analogue may come from a structure reported from a modern onychoteuthid squid (*Onykia*) that has remarkable morphological similarity to the *Megateuthis* rostrum. *Onykia* has a purely organic rostrum that due to its very low density does not act as a counterweight for the soft body. It is assumed that it supports the posterior part of the mantle and fins (= axial stability in Bizikov and Arkhipkin, 1997 and Arkhipkin et al., 2015). A function as a muscle attachment structure for belemnite rostra was first put forward by Stevens (1965). Direct evidence for the presence of fins

in belemnites has recently reported by Klug et al. (2015), favouring a squid-like high speed swimming
mode of life for Jurassic belemnites. Noteworthy to report here, is the case of belemnites
(*Chitinoteuthis*) with a non-calcified rostrum (Müller-Stoll, 1936).

439

440 **5** Open questions and suggestions for future research

441 5.1. Paragenesis of porosity-occluding calcite phase

442 Within individual belemnite rostra, data presented here differentiate: (i) a biogenic, highly porous skeletal framework secreted during the life time of the organism; (ii) an inorganic or 443 444 organomineralic – arguably early diagenetic - calcite phase occluding the pore space of the biogenic 445 framework; and (iii) a late diagenetic, burial, Mn-rich carbonate phase filling fissures and larger 446 cracks within the rostra. Assuming that the above-discussed paragenetic succession is valid, a series 447 of open questions result. In the view of the authors, it is at least conceivable that portions of the 448 rostral pore space were occluded during the life time of the belemnites (remote biomineralization 449 sensu Hücker and Hemleben, 1976; Chinzei and Seilacher, 1993; Seilacher and Chinzei, 1993). If this 450 holds true, then the paragenetic sequence of primary framework and secondary infill calcite is even 451 more complicated than presented here and gradual in nature. Moreover, individual growth 452 increments within rostra are then not representative of specific correlative time intervals. 453 Specifically, each growth increment then represents a complex composite structure of paragenetic phases representing temporally different stages in the belemnite ontogenetic cycle. Obviously, this 454 455 would render the interpretation of time series belemnite geochemical data difficult. Evidence against 456 a biogenic infill of the skeletal pore space by remote biomineralization sensu Seilacher and Chinzei 457 (1993), however, may or may not come from the presence of a corroded outer surface of what is 458 considered the primary skeletal ultrastructure of these rostra (Fig. 2). It seems difficult to argue that 459 intra-rostrum body fluids became corrosive at some stage during the life time of the belemnite animal. Clearly, these questions require further detailed work. 460

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462 5.2 Primary skeletal ultrastructure and preservation of organic matter

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The authors acknowledge the fact that despite the very detailed information regarding the belemnite ultrastructure shown here, our study lacks direct evidence for a primary biogenic origin of the complex, highly porous framework and the subsequent cementation by an early diagenetic calcite phase. Open questions, however, remain. Specifically, the significance of preserved organic matter in the biogenic belemnite calcite deserves attention. In the following, we present several lines (petrographic, optical, and geochemical data) of circumstantial evidence suggesting the presence of preserved organic matter.

471 The sector-wise systematic arrangement of triangular elements - with their a-axis being 472 oriented subparallel to parallel (Figs. 3A-C, 4A-B, 5A, D) - in bundles, indicate a biologically controlled 473 origin (Lowenstam and Weiner, 1989) of these fabrics. The primary belemnite rostrum ultrastructure 474 is brownish (Figs. 5A, 7A) in thin sections under transmitted light and has a relatively low optical 475 relief. According to Ullmann et al. (2014), brownish areas in thin sections of biogenic carbonates are 476 indicative of remnant organic matter (C_{org}). Under the cathodoluminescence- and fluorescence 477 microscope, the primary filigree belemnite framework shows a light blue luminescence (CL) and light 478 green fluorescence (FL). According to Wanamaker et al. (2009) and Pérez-Huerta et al. (2008), 479 fluorescence in biominerals is triggered by organic macromolecules associated with chitin 480 polysaccharides and proteins. Dark fluorescence patterns commonly refer to portions of the skeletal 481 hardparts that are relatively depleted in organic matter. Bright green fluorescence patterns typify 482 areas with increased amount of organic matter (Wanamaker et al., 2009).

Primary skeletal structures display darker colours in SEM-BSE images. Electron microprobe analyses revealed that the biogenic skeletal calcites contain more P and Mg but less S compared to what is here considered an early diagenetic, pore-filling calcite phase. Higher concentrations of P (Fig. 6C) may be related to the presence of organic matter. Arguments for this have been presented by

487 Longinelli et al. (2002, 2003) and Gröcke et al. (2003) who found phosphate (PO_4^{3-}) of presumed 488 biogenic origin being preferentially enriched along concentric growth rings. Generally, the phosphate 489 concentration of ancient belemnite rostra is variable but very low (less than 0.3%) comparable to 490 that in modern *Sepia*.

491 The primary filigree belemnite framework does not fluoresce under the CLFM. The factors 492 that cause fluorescence in samples studied under the CLFM are poorly constrained (Fig. 8). Naturally-493 occurring organic compounds such as proteins or polysaccharides can cause fluorescence in other 494 biogenic carbonates, including brachiopods (Pérez-Huerta et al., 2008), gastropods (Guzman et al., 495 2007), or cephalopods (Linzmeier et al., 2016). In modern brachiopods and Nautilus, portions of the 496 exoskeletons with higher amounts of intracrystalline organic matter (Clark, 1999) appear dark under 497 CLFM (Pérez-Huerta et al., 2008; Linzmeier et al., 2016). This pattern lends support to the argument 498 that dark triangular areas in BSE images represent the primary biogenic skeleton and contain 499 remnants of organic matter. Marine sediments may contain abundant humic substances resulting 500 from the degradation of marine organic matter (Nissenbaum and Kaplan, 1972) and evidence has 501 been presented that the sulphur content of the humic substances increases with degradation 502 (Francois 1987). Concluding, it is here proposed that humic substances caused elevated S 503 concentrations in the diagenetic calcite phase that occludes the skeletal pore space (Fig. 6D) and 504 causes the CLFM fluorescence in all three wavelengths (Blyth et al., 2008; Orland et al., 2009, 2012; 505 Fig. 8).

Similarly, microtomographic data indicate the former presence of organic matter in dark triangular areas of rostra (Fig. 9) as observed in BSE images. The brightness of a carbonate observed in CT image indicates the degree of attenuation of an X-ray passing through this material (Mobilio et al., 2015). As the inner tri-radial structures of rostra appear darker in colour relative to the calcite phase fringing these structures, we suggest that the fringing phase is made of a denser calcite phase compared to the inner structure. That observation is in line with the observation of a low optical relief of these features (supplement Fig. 1A-B).

513 The interpretation presented here with regard to belemnite rostra is arguably consistent with observations of 10-40% organic matrix in the Sepia cuttlebone (Birchall and Thomas, 1983; Florek et 514 al., 2009). This is relevant as we suggest that the belemnite rostrum is structurally similar but not 515 516 homologous (Fuchs 2012) to the Sepia cuttlebone with regard to the primary intra-skeletal porosity. 517 Having said this, the presence of preserved organic matter in ancient biogenic carbonates 518 particularly, intra-crystalline organic matter is not uncommon (Clark, 1999, 2005). Excess carbon 519 observed for Megateuthis has been interpreted as evidence for a former organic matrix within these 520 low-Mg calcite biominerals (Dunca et al., 2006). Similarly, Florek (2004) argued for an excess of carbon in the rostra of Belemnopsis and Hibolites. Summing up: Different lines of circumstantial 521 evidence point to the presence of remnant organic matter within biominerals. These data require 522 523 verification or rejection via the application of spatially highly resolved geochemical data. This work is 524 presently in progress.

525

526 **5.** Conclusions

Ultrastructural data documented here suggest that the calcitic rostra of Mesozoic belemnites yielded 50-90% primary porosity probably filled with body fluids and/or organic matter during the life time of the animal. Porosity was distributed throughout the rostrum as opposed to being limited to the central apical area. The primary biogenic rostrum framework consists of triaxial branches and tetrahedrons of variable size forming a honeycomb-like network. This structure arguably combined mechanical stability with an energy-efficient biomineralization strategy.

The recognition of belemnite rostra as a highly porous structure requires a re-interpretation of the function of the rostrum as counterweight to the soft body and has implications for the swimming mode of belemnites. On the level of a working hypothesis, we argue that the low-porosity fabric found in fossil rostra collected in outcrops worldwide is the result of a syntaxial, early diagenetic cement phase that nucleated upon the surface of the biogenic framework and subsequently occluded the pore space. The possibility of gradual occlusion of skeletal porosity by remote biomineralization during later ontogenetic stages during the life of the animal is possible but seemsunlikely at present.

If the here-presented concepts hold true, then these new findings have significant implications 541 542 with regard to geochemical proxy data collected from fossil belemnite rostra. Specifically, the fact 543 that rostra may consist of biogenic and abiogenic calcite phases formed at different times may 544 explain the controversially low reconstructed seawater temperatures and the uncommonly high 545 scatter of proxy data even from well-preserved rostra collected in the same stratigraphic interval. 546 This is because seawater properties of surficial water masses, the habitat of nektobenthic 547 belemnites, are recorded in the biogenic portions of the rostrum whereas the early diagenetic phase reflects cooler basinal bottom or marine pore water signatures. Findings presented here form a solid 548 549 and well-constrained petrographic data set but one that must be verified by high-resolution 550 geochemical data of all paragenetic calcite phases observed.

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805	Fig. 1A-B) Structural and ultrastructural composition of belemnite rostra. Belemnitella mucronata,
806	thin sections photographed under crossed polarizers. A) Cross section with pseudo-uniaxial cross
807	indicating radially arranged calcite fibres, red box refers to C. B) Longitudinal section with central
808	apical line and radiating fibres from the centre to the margin, stippled line refers to the position of
809	cross section shown in A. C) Idealized bundle of calcite fibres, each fibre contains a stack of
810	tetrahedral elements. D) SEM BSD image of the tetrahedral ultrastructure of Megateuthis gigantea,

811 dashed lines (I-IV) indicate section planes and corresponding reconstructions. Primary skeletal

812 framework is shown in blue, yellow and green whilst early diagenetic phase is shown in white and

813 red for the crystal boundaries. The basis of tetrahedrons points toward the centre of the belemnite

rostrum and its tip towards the rostrum margin i.e. the growth direction. E) Three dimensional

reconstruction of a single tetrahedron of the belemnite endoskeleton. F) Reconstruction of the complex spatial arrangement of biogenic and early diagenetic phases. Colour code in lower right. G) SEM image of a single complex tetrahedron (black line) with indication of structural elements. Primary skeletal components: br = branch, trc = triradial centre, rf = reinforcement, ic = isopachous crystallites. (full page width; bw in print, colour in pdf)

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Fig. 2A-B) SEM BSD images of *Megateuthis gigantea*. A-B) Section perpendicular to the c-axes of calcite fibres (section plane II in Fig. 1D-II). White stippled line indicate dissolution features (early diagenetic), black stippled line indicate microstylolites. (full page width; bw in print, colour in pdf)

Fig. 2A-B) SEM BSD images of *Megateuthis gigantea*. A-B) Section perpendicular to the c-axes of calcite fibres (section plane II in Fig. 1D-II). Blue stippled line indicate dissolution features (early diagenetic), red stippled line indicate microstylolites. (full page width; bw in print, colour in pdf)

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830 Fig. 3A-F) SEM BSD images of Megateuthis gigantea. A-E) Section perpendicular to the c-axes of 831 calcite fibres. A) Triangular structures with a relatively thick outer, light grey margin of abiogenic 832 early diagenetic cement and a small darker centre with branches giving rise to reinforcement 833 structures representing the primary biogenic skeletal framework. B) Larger dark grey, organic rich 834 triangular elements belonging to the biogenic skeletal framework, partly with light grey central 835 abiogenic calcite filling of variable sizes. Branches are often short and cut off at variable distances 836 from the centre. C) Some smaller and a few larger biogenic skeletal elements with cut off branches 837 (lower arrow) and reinforcement structures (upper arrow). Note the variable expression of early 838 diagenetic crystallites with sheaths of remnant organic matter. D) Close up of larger, biogenic skeletal 839 elements and abiogenic isopachous calcites coated by remnants of organic matter within the brighter 840 outer margin. Central portion of the biogenic skeletal elements shows abiogenic crystal; arrows point

to dissolved branches. E) Centre of biogenic skeletal element completely filled with abiogenic bright
calcite leaving only a thin dark inner margin. F) Same specimen, section subparallel to the c-axes
showing a homogenous central portion ("trunk" – white line) of pyramidal morphology rich in
intracrystalline organic matter surrounded by inclined isopachous calcite crystals coated by remnants
of organic matter. This overall pattern results in a "Christmas tree like" structure (compare with Fig.
1D). (full page width; bw in print and pdf)

Fig. 4A-E) Megateuthis gigantea, EBSD map with colour code in sections perpendicular to the c-axes of the fiber bundles. Same colours represent same crystallographic orientations. In A and B angular deviation from blue to red is up to 40°, in D angular deviation is 2°. A) Overview map showing the bundling of fibres with identical orientation of a-axes, black frame indicates area for close up in B; blue frame refers to Fig. 5A-C. B) Close up map, within one bundle blue tinted fibres are mainly neighboured by other blue fibres, red tinted fibres are surrounded by red fibres. C) Close up documenting minor angular deviation within one fibre (compare with D). D) Map of a single fibre with an angular deviation of 2° from blue to red, showing a slight systematic shift of axes orientation. E) Pole-plots of c-axes {001} and a-axes {010} from all fibres shown in A, all c-axes show nearly the same orientation while the a-axes demonstrate the bundled structure of the rostral fabric which may have improved the stability of the skeletal structure. (full page width; bw in print, colour in pdf)

Fig. 5A-F) Transmitted light, polarized light and cathodoluminescence. Thin section of Megateuthis gigantea, A-C refer to the blue frame in Fig. 4A, D-F are close ups (black frame in A) A and D) TL image perpendicular to the c-axes of calcite fibres, filigree biogenic skeletal framework is indicated by the dark tinted structures, primary porosity is represented by the abiogenic translucent calcites. Note banded distribution of calcite fibre domains relating to larger and smaller organic-rich biogenic skeletal elements, single fibres may contain a central portion of transparent calcite of varying size, bundling of adjacent calcite fibres is indicated by the same orientation of the triangles. B and E) Uniform extinction (orientation) of adjacent fibres under crossed polarizers. C and F) CL of abiogenic calcite portions show a dark blue, intrinsic luminescence (pure stoichiometric calcite), CL of the biogenic skeletal framework show light blue luminescence. (full page width; bw in print, colour in pdf)

Fig. 6) Electron microprobe data for *Megateuthis gigantea*. A) Overview BSE map B) Shows higher
Mg concentrations within the triangular areas dark in BSE images and a lower Mg concentration in
the surrounding area bright in BSE images. C) Shows higher P concentrations within the triangular
areas dark in BSE images and a lower P concentration in the surrounding area bright in BSE images.
D) Shows lower S concentrations within the triangular areas dark in BSE images and higher S
concentrations in the surrounding region bright in BSE images. (full page width; bw in print, colour
in pdf)

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Fig. 7) Fluorescence microscope images for *Megateuthis gigantea*. A) Transmitted light shows
brownish triangular structures, rich in organic matter and dark in BSE images, partly with central
translucent areas (compare with Fig. 1D section plane IV, Fig. 3D, E, 5A, D). B) Shows brighter
fluorescent triangular area compared to the in transmitted light translucent calcite. (full page width;
bw in print, colour in pdf)

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902 Fig. 8) Confocal laser fluorescence microscopy images of Megateuthis gigantea. A) CLFM images 903 showing fluorescence in far-red light (λ = 664 to 696 nm). B) CLFM images showing fluorescence in 904 visible green light (λ = 505 to 539 nm). C and D) CLFM images showing fluorescence in visible red light 905 $(\lambda = 589 \text{ to } 621 \text{ nm})$. Triangular structures visible in other imaging techniques (Fig. 3, 5-7) do not 906 fluoresce as brightly as cracks (B) or early diagenetic calcite (Fig. 3) separating the triangles (A, B, C, 907 D). D) Higher magnification shows some brighter fluorescing calcite between the triangular elements 908 dark in BSE images (Fig. 1, 3). Brightly fluorescent early diagenetic calcite separating triangles is 909 enriched in S (Fig. 6). (full page width; bw in print, colour in pdf)

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Fig. 9) Three-dimensional visualization of the filigree biogenic framework. Synchrotron radiation based tomographic visualization of a sub-volume of the rostrum of *Megateuthis gigantea*. Specimen was scanned with an isotropic voxel size of 0.74μm. A) Multi-planar image of a sub-domain of the original dataset with dimensions of 447x592x663 voxels, triangular elements dark in BSE images appear here as dark elements due to reduced densities. B-D) Volumetric renderings of the same subdomain with variable rendering settings. (full page width; bw in print, colour in pdf)

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920 Supplementary Figures

Fig. 1) Examples of the filigree framework from other belemnite species. A-B) *Gonioteuthis quadrata*, A) shows the Becke line outside of the triangular area with a relatively larger distance between the sample and objective. B) Image shows the Becke line within the triangular area while the distance between the sample and the objective was reduced, accordingly the triangular area (dark in BSE; Fig. 3) has a lower optical relief. C-D) *Belemnitella mucronata*, C) thin section under polarized light, D) same area under CL showing microfractures filled with Mn-rich calcite tracing the outline of triangular elements (encircled). (full page width; bw in print, colour in pdf)

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930 Fig. 2) SEM BSD images of Megateuthis gigantea. A-F) Section plane parallel to the c-axes of calcite 931 fibres. A-C) stepwise enlargement of a particular area. D-F) stepwise enlargement of a particular 932 area. A and D give the impression of a concentric arrangement of distinct darker and brighter layers 933 (black frames enlarged in B and E), arrow in D point to an organic rich layer (laminae obscura sensu 934 Müller-Stoll, 1936). B and E) Allow the recognition of single darker structures of tetrahedral 935 morphology with their tips pointing towards the outer margin of the belemnite rostrum, i.e. the 936 growth direction (black frames enlarged in C and F). C and F) show the intricate framework of 937 biogenic (dark) and abiogenic (light) carbonate phase within the rostrum, larger dark grey, triangular 938 elements belonging to the biogenic skeletal framework, partly with light grey central abiogenic 939 calcite filling of variable size. (full page width; bw in print, colour in pdf)

Highlights

We present well-constrained petrographic evidence for the complex primary biogenic framework of belemnite rostra.

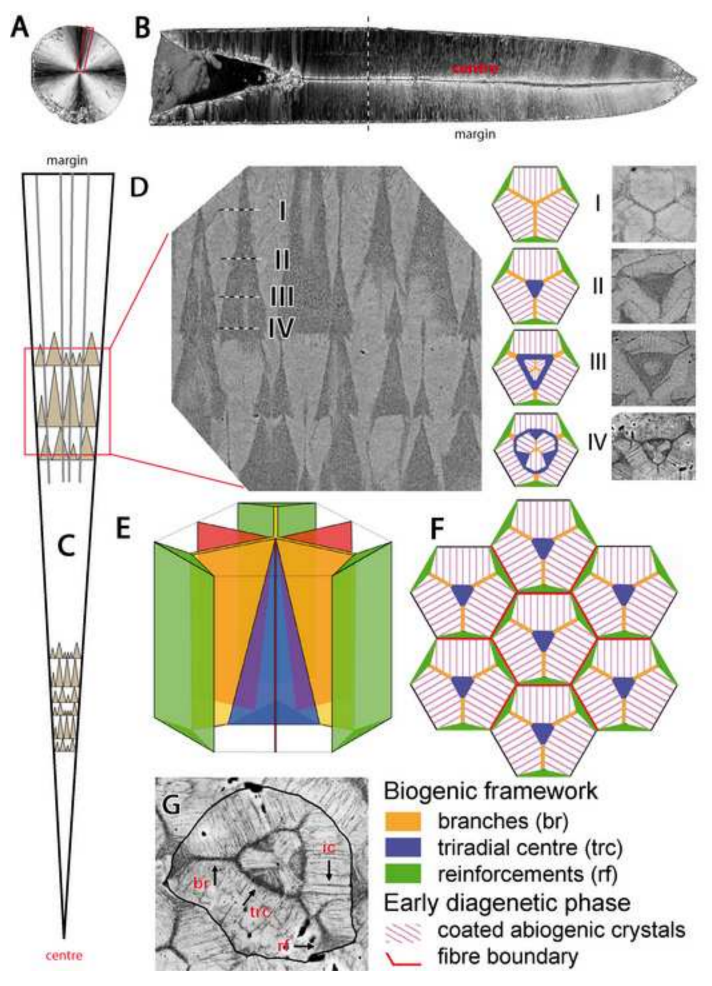
Petrographic evidence demonstrates a high (50-90%) primary porosity for the bulk of the belemnite rostrum.

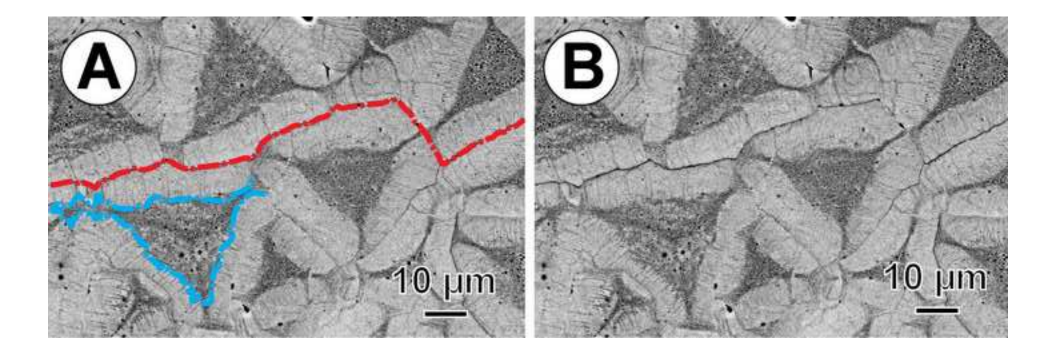
The relative proportion of biogenic skeletal framework versus abiogenic, earliest diagenetic calcite occluding the former pore space in rostra is assessed.

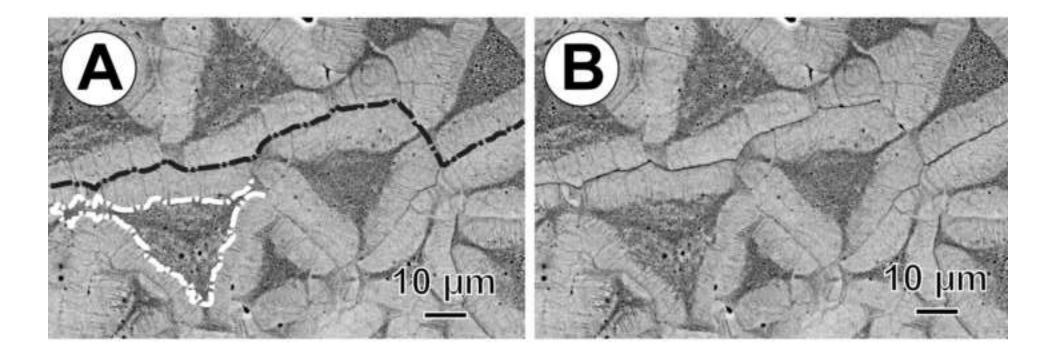
We discuss the timing and formation modes of pore-filling calcites.

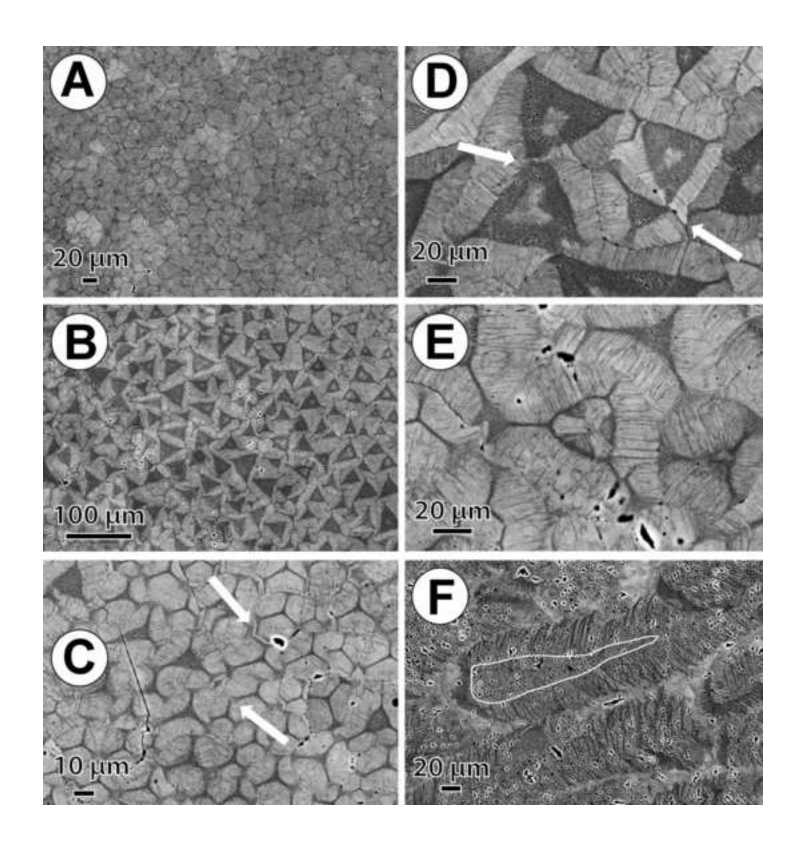
The significance of these findings for reconstructions of marine palaeo-environments and the function of the rostrum is discussed.

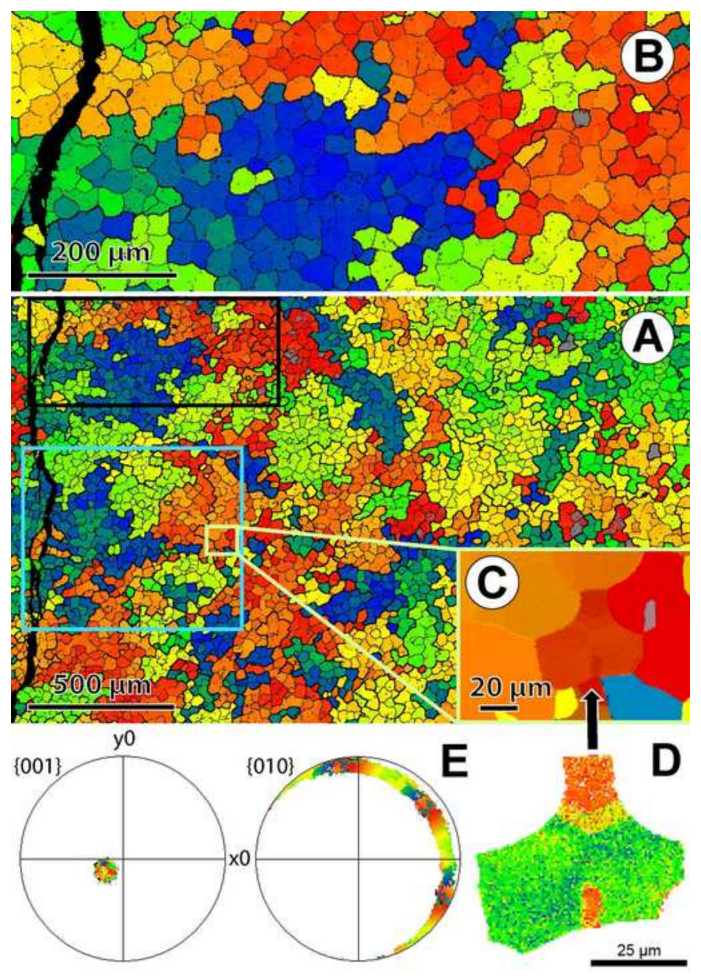
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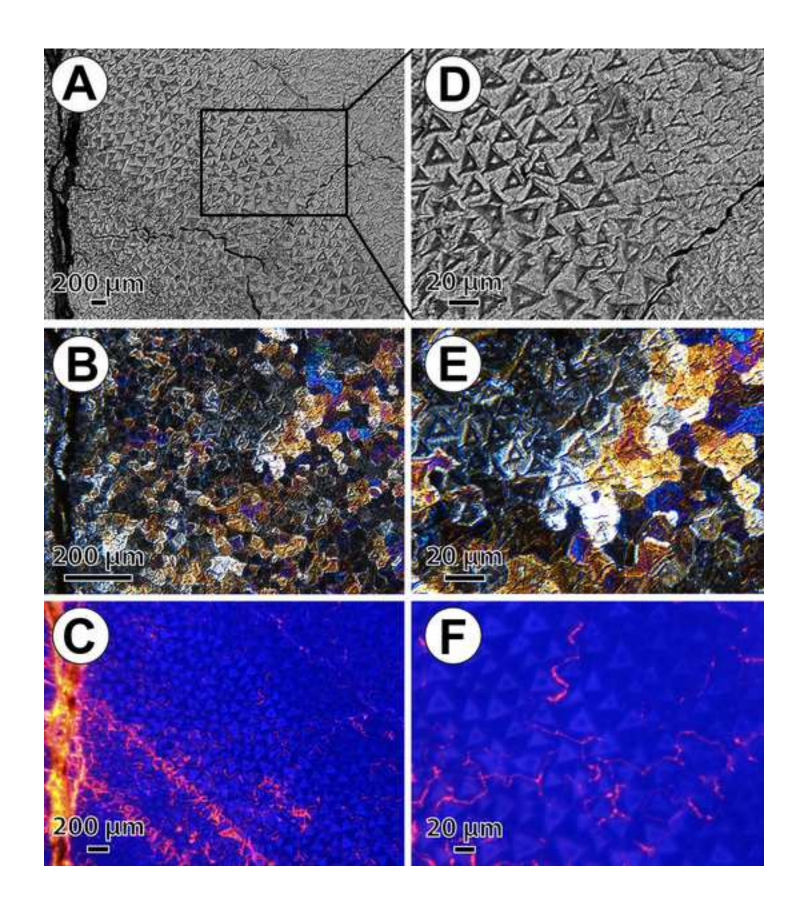


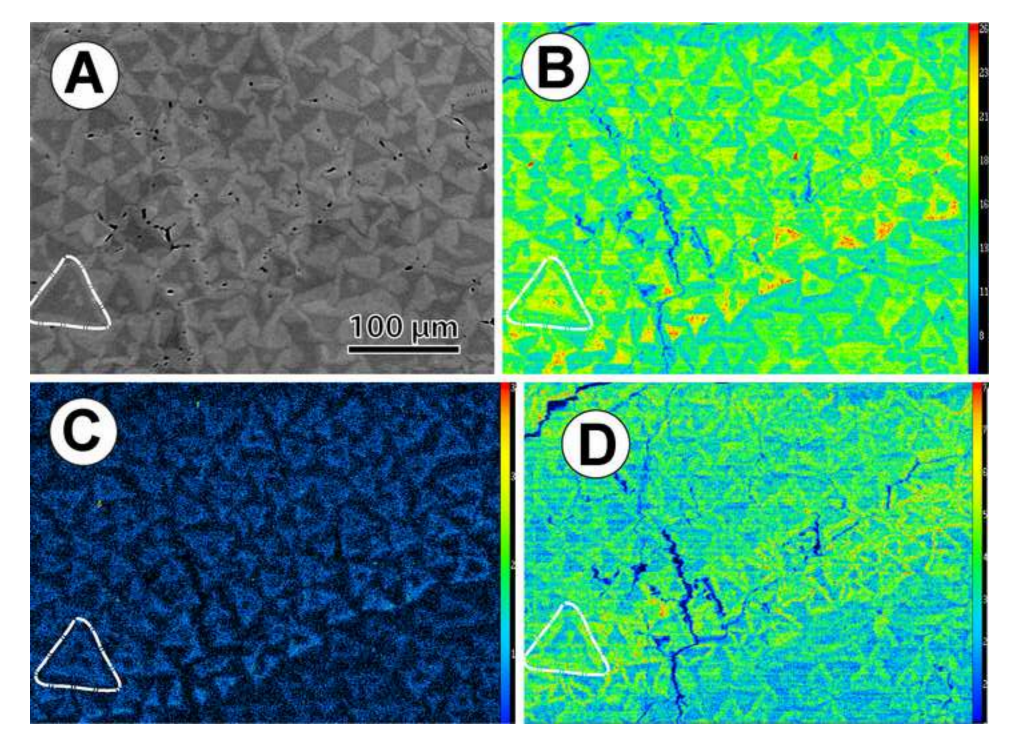


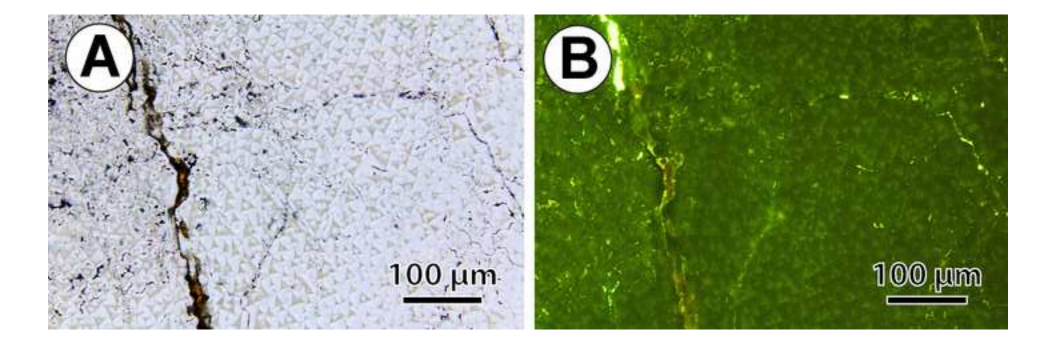












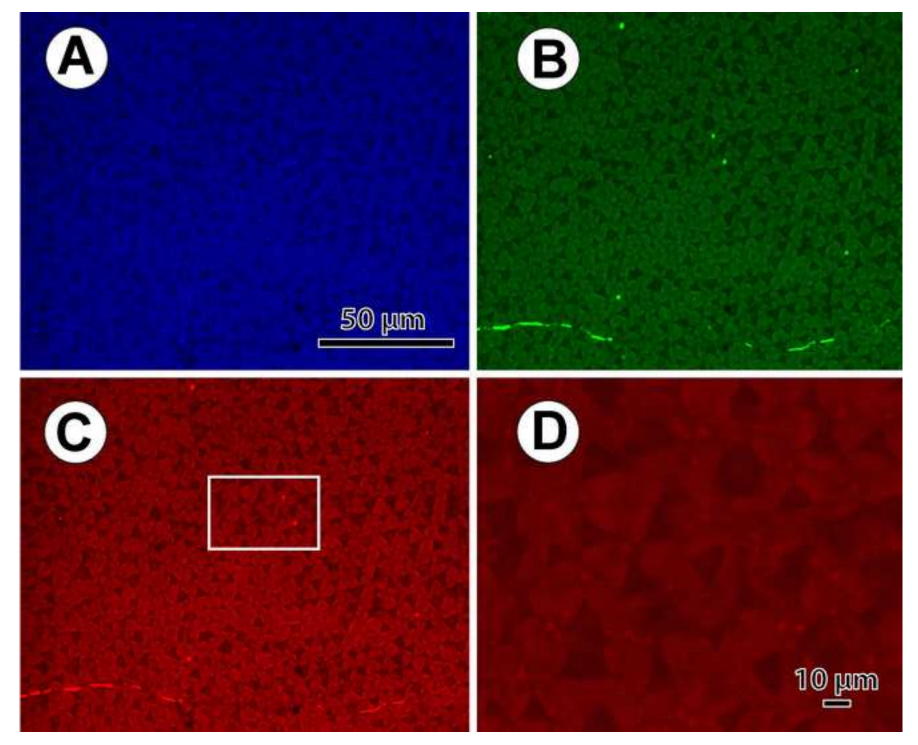
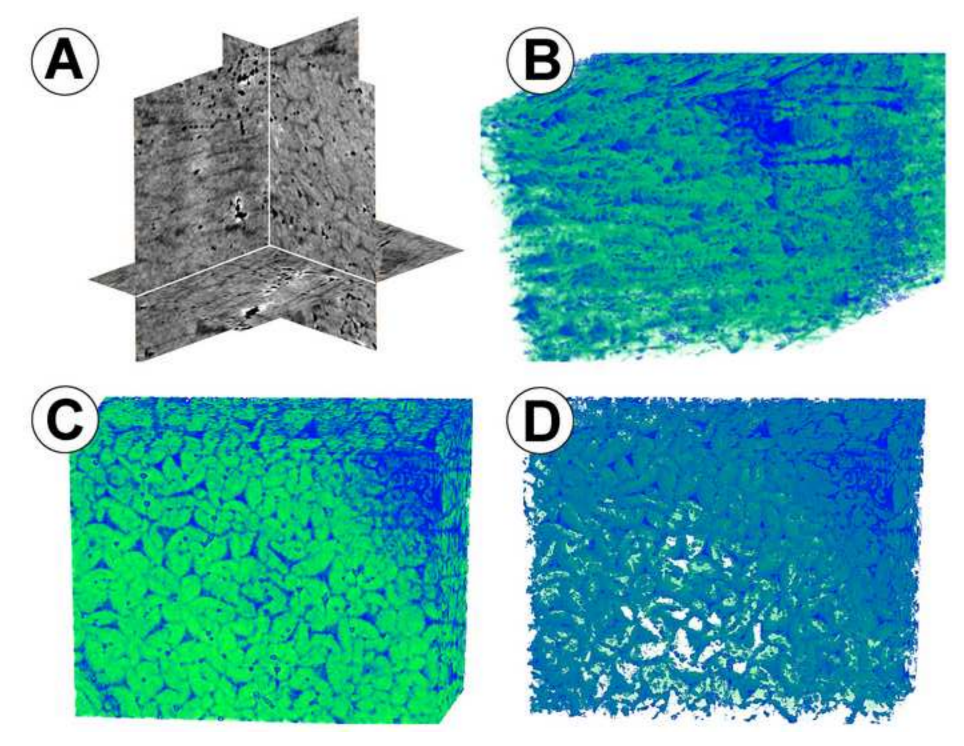


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Supplementary material for on-line publication only Click here to download Supplementary material for on-line publication only: Supplement Fig. 2 - full width.jpg