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Perspectives on heterococcolith geochemical proxies based on high resolution X-ray fluorescence mapping

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ABSTRACT

Heterococcoliths are micron-scale calcite platelets produced by coccolithophores. They have been the most abundant and continuous fossil record over the last 215 million years (Myr), offering great potential for geochemical studies, although the heterococcolith fossil record remains underutilised in this domain. We have mapped heterococcoliths' composition using X-ray fluorescence (XRF) with a 100 nm resolution beam in order to decipher element distributions in heterococcoliths and to investigate the potential development of geochemical proxies for palaeoceanography. The study presents two Middle Jurassic Watznaueria britannica heterococcoliths from Cabo Mondego, Portugal. XRF analysis was performed with a 17 keV incident energy beam at the European synchrotron radiation facility ID22NI beamline in order to study elements from Sr down to S. Ca, Sr and Mn are distributed following the heterococcolith crystalline arrangement. Cl, Br and S display an homogeneous distribution, whereas K, Fe, Cu, Zn and Rb are concentrated at the edges and in the central area of the heterococcoliths. Distributions of K, Fe, Ti, Fe, Cu, Zn, Rb, and to a lesser extent V and Cr, are highly influenced by clay contamination and peripheral diagenetic overgrowth. Mn is related to diagenetic Mn-rich CaCO\textsubscript{3} overgrowth on top of or between heterococcoliths shields. Cl and Br are likely to be present in heterococcoliths inside interstitial nano-domains. We assume that the cytoplasm [Cl\textsuperscript{-}] and [Br\textsuperscript{-}] are mediated and constant during heterococcolithogenesis. Assuming a linear correlation between cytoplasm [Cl\textsuperscript{-}] and seawater [Cl\textsuperscript{-}], heterococcolith Cl may have potential as a salinity proxy. As S is incorporated into heterococcoliths by sulphated polysaccharides, and our study suggests a role for such polysaccharides in heterococcolithogenesis for at least 170 Myr. The low Sr/Ca in the W. britannica specimens studied here may either highlight an unusual cellular physiology of Mesozoic coccolithophores or result from low growth rates in oligotrophic environments.

INTRODUCTION

Coccoliths are micron-size platelets (< 15 μm) produced by the golden-brown algae coccolithophores. They are the most abundant calcareous nannofossils in marine sediments from coastal to open-ocean environments over the last 215 Myr (e.g. Winter & Siesser, 1994;
Bown, 1998; Thierstein & Young, 2004). Coccoliths are produced inside the cell and then extruded through the cell membrane to cover the cell (Taylor et al., 2007), forming an exoskeleton with multiple interlocking coccoliths called a coccosphere. Two types of coccoliths are distinguished: i) holococcoliths, which are produced during the haploid phase of the coccolithophore life-cycle and ii) heterococcoliths, which are produced during the diploid phase (e.g. Geisen et al., 2002). In the fossil record, heterococcoliths far outnumber holococcoliths, probably because they have greater preservation potential and/or different ecological preferences (Dunkley Jones et al., 2009; Cros & Estrada, 2013).

Heterococcoliths are generally made of very low Mg-calcite (Mg/Ca = 0.1-0.2 mmol/mol; Stoll et al., 2001). Low Mg-calcite is relatively resistant to dissolution in deep cold seawater because of its thermodynamic properties (Schlanger et al., 1973). As a consequence, heterococcoliths are well preserved in the geological record. However, because of their very small size (2 to 15 µm in diameter), and extremely low mass (5 to 650 × 10^{-12} g), analysing their chemical composition remains difficult, and fossil heterococcoliths are seldom used as a geochemical proxy substrate for palaeoceanography, compared to planktic foraminifera that are commonly used.

Studies of heterococcolith geochemistry first emerged in the late 1970's (e.g. Siesser, 1977; Dudley & Goodney, 1979; Dudley et al., 1986) and the number of such studies has grown over the past fifteen years (see review by Stoll & Ziveri, 2004), as it became possible to separate heterococcoliths from bulk sediment (Stoll & Ziveri, 2002; Minoletti et al., 2009; Stoll & Shimizu, 2009) and analyse single heterococcoliths (Stoll et al., 2007a; Prentice et al., 2014). Several geochemical proxies based on heterococcoliths are now calibrated, such as temperature proxies based on δ^{18}O (Candelier et al., 2013), Mg/Ca (Ra et al., 2010) or 'clumped isotopes' (Tripati et al., 2010); seawater composition proxies based on Mg concentration and Mg/Ca (Stanley et al., 2005; Müller et al., 2011), and productivity proxies based on Sr/Ca (Stoll & Schrag, 2000; Rickaby et al., 2002; Stoll & Bains, 2003). However, with the exception of the recent study by Prentice et al. (2014), heterococcolith geochemistry still lacks high spatial resolution studies that help to understand element incorporation mechanisms (e.g. for foraminifera (Sadekov et al. 2005)). In the present study, we analyse the chemical composition of single heterococcoliths at 100 nm spatial resolution using synchrotron X-ray fluorescence (XRF). The heterococcoliths analysed are two 170 Myr-old Watznaueria britannica heterococcoliths from Cabo Mondego, Portugal. W. britannica is an elliptical imbricated placolith made of several c-axis calcite single crystals and is abundant in middle Jurassic to Cretaceous deposits (Fig. 1A-B). The high-resolution 2D mapping allows us to discuss the distribution of elements, deduce mechanisms of elemental incorporation in heterococcoliths, and propose new perspectives on heterococcolith geochemistry.

MATERIAL AND METHODS
Heterococcolith samples and picking
W. britannica heterococcoliths were picked from a Cabo Mondego (Portugal) marlstone sample (CM62 in Suchéras-Marx et al., 2012), and are dated to the Propinquans ammonite Zone in the Early Bajocian (Middle Jurassic, ca 170 Ma). This sample was selected because of the high abundance of well-preserved heterococcoliths, with overgrowth only observed under the SEM (Fig. 1B). The marlstone sample was powdered in an agate mortar using a
power hammer. A smear slide (Bown & Young, 1998) was then prepared with ethanol. Heterococcoliths were separated from the bulk sample following a picking method modified from Stoll & Shimizu (2009), using a 150 µm silicate tube thinned down to ~15 µm diameter by stretching in a flame. Individual heterococcoliths were picked by hand from the smear slide and deposited on sample holders using a Leica DM750P microscope at 400x magnification. Samples were mounted on 500 nm thick silicon nitride (Si₃N₄) TEM windows (Silson Ltd.). A droplet of ethanol deposited on the window helped to detach the heterococcolith from the silicate tube by surface tension. The W. britannica heterococcoliths presented in this study are called Wbrit1 and Wbrit2.

**Synchrotron radiation X-ray Fluorescence**

XRF measurements were performed at the European synchrotron radiation facility (ESRF), Grenoble, France on the beamline ID22NI. The XRF method is based on the excitation of elements by high energy X-rays. Due to the incident X-ray excitation, elements present in the sample emit secondary X-rays with constant energies per element and intensities depending on the concentration of each element. The incident X-ray beam energy was set at 17 keV to ensure the measure of Sr (Kα = 14.165 keV) and lighter elements. ID22NI uses an ESRF custom-made Kickpatrick-Baez double multilayer mirrors device focusing the X-rays into a beamspot of 100 nm × 100 nm. Analysis spots are adjacent, without overlapping or leaving space between two spots. The detectors were high-count rate twin SiI™ Vortex SDD (silicon drift diodes) detectors, capable of counting up to 200 kcps with no saturation and no peak shift or FWHM broadening, when operated below 10% dead time. Heterococcoliths were analysed with 2 s dwell time per pixel.

The Wbrit1 map was 81 x 71 pixels (~3 h 15) and the Wbrit2 map was 85 x 65 pixels (~3 h). The sample is orthogonal to the incident beam and the detector is located at a 15° angle relative to the sample surface. XRF is a penetrative method and one pixel sums the fluorescence signal throughout the whole heterococcolith thickness, the TEM windows and the air on both sides of the sample. The TEM windows were homogeneous in all the elements analysed and the air composition did not interfere with measured heterococcolith composition. The set up geometry influences the mapping results. The maps will give an image, which depends on the excitation angle *i.e.* orthogonal to the heterococcoliths shields in our set up. The intensity of photons per element depends on the concentration and the thickness of the analysed material. If the heterococcoliths have homogeneous composition, the thickness influence can be excluded because it would be the same for all elements. However, in case of layers with higher or lower concentrations in a given element inside or over (i.e. overgrowth crust) a heterococcolith, the thickness of the layer is a key parameter, which should be considered in the interpretations. Elemental mapping of the NIST 1577b Bovine Liver standard was performed under similar conditions in order to calibrate the exact analytical geometry. The fitted spectrum was constructed using the sum of photons for all heterococcolith pixels divided by the number of pixels, and the fitting calculations were made pixel by pixel. Fits were performed using the PyMCA 5.7.0 (Solé et al., 2007) software, which uses the Hypermet function coupled with the standard Levenberg-Marquardt non-linear fitting method. Peaks were routinely fitted by a least squares procedure, using Poisson weights based on their surfaces. The lines used in the fitting procedure are Kα and Kβ lines –
the main transition lines from orbitals with principal quantum number $n=2$ and 3 to $n=1$ – and also $L \alpha$ and $L \beta$ – main transition lines from orbital $n=3$ to $n=2$ lines. Absolute minimum detection limits (MDL) at ID22NI estimated in the standard 1000 s interval are 20-70 zg (2-7×10$^{-20}$ g) for intermediate Z elements. When mapping with a 1 s dwell time, the absolute MDL is approximately 1-2 fg, whereas relative limits are 1-2 ppm for Cr-Cu. PyMCA was also used to calculate the relative molar fraction of element $i$ to calcium Ca ($i$/Ca; mmol/mol).

Figure 1: A) Crystal organisation of *W. britannica* (derived from Young & Bown, 1991) in distal and proximal view and in cross-section. *W. britannica* is an elliptical imbricated placolith, the long axis of the ellipse is perpendicular to the central area bridge and called major axis; the axis perpendicular to the major axis is called the minor axis. *W. britannica* is made of several c-axis elongated calcite monocrystals organised in two shields – forming the rim – connected by a central inner tube partly filled by a bridge. The shield in contact with the cell in life position is called proximal shield, the other one is called distal shield. The shields are made in the outer part and most inner part of calcite crystals with a radial growth direction (from the centre to the edge) intersected by crystals with a vertical growth direction (from the proximal to the distal shield). In proximal view, the calcite crystals of the proximal and distal shields are imbricated anti-clockwise (laevogyre) and clockwise (dextrogyre), respectively. B) SEM picture of a *W. britannica* from the same rock sample of the heterococcoliths analysed in this study. The heterococcolith is presented in proximal view allowing the observation of the crystal organisation of the proximal shield.

**Calculation of relative ratio $i$/Ca**
Relative ratio tables and maps for the two heterococcoliths are shown in the results section, and show the concentration of element $i$ with respect to the major element Ca. This procedure of normalisation allows correcting the element intensity for the sample thickness effect. The constant contribution of the TEM Si$_3$N$_4$ windows to the concentration values for each element was subtracted. The trace element composition of the TEM windows in element $i$ was calculated as the median of the concentration of element $i$ in all pixels void of heterococcolith contribution, *i.e.* around the heterococcolith. We use the median value rather than the arithmetic mean because it is less affected by potential outliers. Ratios are expressed in mmol/mol. The cut-off threshold contouring heterococcoliths was set for $[Ca] = 3 \times 10^5$ cps
and Ca ratios have been arbitrarily set to 0 below the latter threshold value. In Table 1, three regions of interest (ROI) are presented, the whole heterococcolith (Full), a region depleted in K, Fe, Cu, Zn and Rb (ROI 1) and a region more concentrated in K, Fe, Cu, Zn and Rb (ROI 2). Each ratio corresponds to the arithmetic mean of pixels in the ROI; the arithmetic mean is used because it is sensitive to the variance and allows more easily discussing the effect of outliers on heterococcoliths bulk composition (Full vs ROI 1 and ROI 2).

### Table 1: Wbrit1 and Wbrit2 elemental ratio (mmol/mol) of the whole heterococcolith (Full), in overgrowth-free region (ROI 1) and in overgrown region (ROI 2). Selected region of interest are presented in Fig. 6.

<table>
<thead>
<tr>
<th>Elemental ratio (mmol/mol)</th>
<th>Wbrit1</th>
<th>Wbrit2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Full</td>
<td>ROI 1</td>
</tr>
<tr>
<td>Si/Ca</td>
<td>4.843</td>
<td>3.776</td>
</tr>
<tr>
<td>K/Ca</td>
<td>12.325</td>
<td>0.877</td>
</tr>
<tr>
<td>Ti/Ca</td>
<td>1.091</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>V/Ca</td>
<td>0.111</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Cr/Ca</td>
<td>0.068</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Mn/Ca</td>
<td>0.961</td>
<td>0.820</td>
</tr>
<tr>
<td>Fe/Ca</td>
<td>9.750</td>
<td>2.126</td>
</tr>
<tr>
<td>Cu/Ca</td>
<td>0.361</td>
<td>0.044</td>
</tr>
<tr>
<td>Zn/Ca</td>
<td>0.495</td>
<td>0.12</td>
</tr>
<tr>
<td>Br/Ca</td>
<td>0.013</td>
<td>0.008</td>
</tr>
<tr>
<td>Rb/Ca</td>
<td>0.022</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sr/Ca</td>
<td>0.351</td>
<td>0.365</td>
</tr>
</tbody>
</table>

**RESULTS**

**Elemental spectrum**

The XRF spectrum displays the K-lines of 16 elements (Fig. 2), namely S, Cl, Ar, K, Ca, Ti, V, Cr, Mn, Fe, Cu, Zn, Br, Kr, Rb, Sr. The L-lines of Pb are also identified in the XRF spectrum. Due to the incident excitation beam energy of 17 keV, no element heavier than Sr has been identified aside from Pb. At low energy, the setup at ID22NI in air does not allow measuring elements lighter than S. Ar and Kr observed in the spectrum are from the air in the experimental hutch. Traces of Pb are usually considered as contamination, since high-energy experimental hutch at the ESRF are shielded with Pb. The least intense K-line identified, namely Cr K-line, is still about 30 times higher than the MDL. XRF is a penetrative method such that the incident beam goes across the sample with negligible absorption (≤ 0.004) while the fluorescent lines feature moderate self-absorption (corrected) as a function of their energy and exit angles in the sample. The Mn Kα-line features roughly 12% absorption in a 3 µm CaCO₃ sample while Ca displays features between 12% and 45% absorption. This progressive absorption as a function of the exit angle is a smooth dependence, which affects only the low Z element lines such as those of S, Cl, K and Ca. This dependence does not produce a visible effect on the element distribution but affects the whole map by underestimating the pixel values, which contain a partial grazing exit contribution. Hence, 14 trace elements were successfully identified and mapped in *W. britannica*. 
Figure 2: XRF spectrums with i) Data (sum) corresponding to the sum of all pixels within W. britannica Wbrit1, ii) Data (normalised) corresponding to the sum of all pixels normalised by the number of pixels within W. britannica Wbrit1, iii) Membrane corresponding to the sum of all pixels normalised by the number of pixels within the TEM window holding the heterococcolith, and iii) Fit corresponding to the fit calculated with PyMCA and used for calculating the concentration of element i relative to Ca (i/Ca) in Table 1 and Figs. 7 to 9.

Elemental maps
To clearly present the 14 elemental maps produced, they were separated in three groups based on a comparison between their 2D distribution and the morphology of W. britannica. The results in cps are presented in Figs. 3-5 with thresholds applied (see captions) in order to observe the distinctive details. The full datasets are presented on a log-scale in Supplementary Material 1 for Wbrit1 and Supplementary Material 2 for Wbrit2.

Group 1: morphological distribution (Ca, Sr and Mn)
Results for Ca, Sr and Mn are presented in Fig. 3. The Mn map in Wbrit1 shows 2 concentric rings, the wider outer ring is composed of radial features whereas the inner ring is very narrow, about 200 nm wide. In Wbrit2, only the external ring is clearly observed. The central parts of both heterococcoliths are almost entirely Mn-free. The Sr maps for both Wbrit1 and Wbrit2 show 2 concentric rings with radial features. The Sr outer ring is superimposed on the
Mn outer ring whereas the Sr inner ring has a slightly larger diameter than the Mn inner ring. On Ca maps, radial features are clearly visible in the outer ring and continue towards the central area. The central area of the heterococcoliths contains less Ca and Sr than the shield.

Figure 3: Elemental maps of Ca, Mn and Sr presented in photon counts per second (cps). Scales do not display the full range of values, thresholds are set to offer a better observation of the organisation.

Group 2: homogeneous distribution (S, Cl, Br, Ti, V, Cr)
Results for S, Cl, Br, Ti, V and Cr are presented in Fig. 4. S, Cl, Br and to a lesser extent Ti, V and Cr have an homogeneous distribution. Cl and Br have a very similar elemental distribution, whilst Br is less abundant. S does not show any organisation apart from a discontinuous ring with variable width that is slightly more concentrated. S is slightly less abundant in the central area and its distribution is more similar to Br than Cl. Cr and V have very similar elemental distributions, quite homogenous except for a zone of higher concentration close to the centre of the heterococcolith. Finally, Ti has an overall low abundance, but in ~15 spots ranging in size from 100 to 500 nm in which Ti concentration is very high. However, the Ti map drawn in log-scale displays a 2D distribution that is very similar to that of Cr and V except for the highly enriched spots (Supplementary material 1 and 2).
Figure 4: Elemental maps of S, Cl, Ti, V, Cr and Br presented in photon counts per second (cps). Scales do not display the full range of values, thresholds are set to offer a better observation of the organisation.
Figure 5: Elemental maps of K, Fe, Cu, Zn and Rb presented in photon counts per second (cps). Scales do not display the full range of values, thresholds are set to offer a better observation of the organisation.
**Group 3: peripheral and central distribution (K, Fe, Cu, Zn, Rb)**

Results for K, Fe, Cu, Zn and Rb are presented in Fig. 5. The distributions of these elements are very similar. They are highly concentrated on the external edge of the heterococcolith and in an area that covers the centre and extends along the heterococcolith's major axis. All these elements present comparable 2D distributions, however Rb and K are most similar to each other and Fe is similar to Cu and Zn.

**Element/Ca relative concentrations and maps**

**Relative concentration**

Relative concentrations of 13 elements normalised to Ca, hereafter designated as \( i/Ca \) ratios, are presented in Table 1, with the relative concentrations for the whole heterococcolith (Full) and for labelled regions of interest (ROI 1 and 2) (Fig. 6) shown. Maps of \( i/Ca \) ratios (in mmol/mol) are presented in Figs. 7-9.

In Table 1, the first column gives the mean \( i/Ca \) ratio for the whole heterococcolith; the second column labelled ROI 1 corresponds to the mean \( i/Ca \) ratio for a zone depleted in K, Fe, Cu, Zn and Rb. The last column labelled ROI 2 corresponds to the mean \( i/Ca \) ratio for a zone enriched in K, Fe, Cu, Zn and Rb. ROI 2 was selected because of the presence of metals identified in previous studies on calcite shells as contaminants (Pena et al., 2008; Prentice et al., 2014) while ROI 1 represents a more pristine section of the heterococcolith.

![Figure 6: Localisation of ROI 1 and ROI 2 in Wbrit1 and Wbrit2.](image)

The highest mean elemental ratios are observed for Cl/Ca, K/Ca and Fe/Ca. V/Ca, Cr/Ca, Br/Ca and Rb/Ca have very low elemental ratios, often below 0.1 mmol/mol, in the part per million range.

The comparison of \( i/Ca \) ratios between the whole heterococcolith and the selected ROI 1 and ROI 2 zones reveals two major features. i) Sr/Ca, Mn/Ca and Br/Ca have very comparable relative concentrations, ii) K/Ca, Ti/Ca, V/Ca, Cr/Ca, Fe/Ca, Cu/Ca, Zn/Ca, Rb/Ca are less abundant in ROI 1 than in the whole heterococcolith and ROI 2. In most case, \( i/Ca \) ratios in ROI 2 are similar to or very close to the mean values in the whole heterococcolith. Finally, in
Wbrit2, S/Ca and Cl/Ca are higher in ROI 1 than in ROI 2 and the whole heterococcolith whereas Br/Ca ratios are constant between ROI 1, ROI 2 and the whole heterococcolith.

Figure 7: Ratio maps of Mn/Ca and Sr/Ca presented in mmol/mol. Scales do not display the full range of values, thresholds are set to offer a better observation of the organisation.

**Group 1: morphological distribution (Sr/Ca and Mn/Ca)**

Results for Sr/Ca and Mn/Ca are presented in Fig. 7. Mn/Ca display exactly the same distributions as the Mn elemental maps, two concentric rings with pronounced radial structures in the outer ring in Wbrit1 and an outer ring with radial structures in Wbrit2. In Wbrit1 and Wbrit2, the radial structures of the external ring are marked by the alternation of high and low Mn/Ca. The Sr/Ca maps display a complex organisation in both heterococcoliths. In Wbrit1, Sr/Ca is organised in three concentric rings, with the external and the internal ones separated by a ring with lower Sr/Ca. In Wbrit2, four concentric rings are observed, with a supplementary ring located between the external ring and the intermediate depleted ring. This extra ring is very thin (i.e. ~200-300 nm) with high Sr/Ca. The Sr/Ca rings described on both heterococcoliths systematically display radial features.

**Group 2: homogeneous distribution (S/Ca, Cl/Ca, Br/Ca, Ti/Ca, V/Ca, Cr/Ca)**

Results for S/Ca, Cl/Ca, Br/Ca, Ti/Ca, V/Ca and Cr/Ca are presented in Fig. 8. All element ratios i/Ca in this group present artificially very high values in the central area due to the presence of a hole. In both heterococcoliths, the S/Ca ratio is very homogeneous and Cl/Ca and Br/Ca are very similar, displaying a homogenous pattern with some weak radial features. V/Ca and Cr/Ca display a general homogenous pattern with small zones that are highly enriched. Lastly, the Ti/Ca distribution is patchy with a few small, very Ti-rich zones (i.e. ~200-700 nm) and a general pattern that is similar to those of V/Ca and Cr/Ca ratios.
Group 3: peripheral and central distribution (K/Ca, Fe/Ca, Cu/Ca, Zn/Ca, Rb/Ca, Ti/Ca)
This third group composed of K/Ca, Fe/Ca, Cu/Ca, Zn/Ca and Rb/Ca elemental ratios is characterised by maps showing a relative enrichment in the centre and outer edge of heterococcoliths (Fig. 9). In the i/Ca maps of group 3, an elongated, enriched zone along the major axis (larger than the central area) is located in the centre of the heterococcoliths. Radial features are sometimes observed and are more pronounced in Wbrit1 than in Wbrit2. In Wbrit1 Cu/Ca, Zn/Ca and Fe/Ca, on the bottom part of the maps, those radial structures are closer to the heterococcolith external border than to the centre. But on the upper and left parts of the maps, the radial structures seem longer, radiating from the external border almost to the centre. In Wbrit2 Fe/Ca, Cu/Ca, Zn/Ca and Rb/Ca, there are fewer radial structures radiating from the heterococcolith external border to a depleted zone. Ti/Ca is also classified as group 3 because despite a generally homogenous pattern, it presents several spots with high Ti/Ca relative concentration as mentioned earlier.

DISCUSSION

Post-depositional overprint on the heterococcolith composition

Clay contamination
We consider as contamination any particle that was not originally a piece of the heterococcolith. We tried to minimise contamination from the matrix rock by using the careful picking method developed in the present study. In sedimentary carbonate deposits, contaminants could be either carbonate from the matrix (e.g. broken pieces of shells and tests), siliciclastic particles (e.g. clay, quartz) or organic matter (OM). In both heterococcoliths Wbrit1 and Wbrit2, the centre of the heterococcoliths is contaminated. This zone is depleted in Ca but highly enriched in K, Fe, Cu, Zn and Rb (Figs. 3 and 5). This contamination cannot be due to broken pieces of carbonate because Ca concentration is low. We also exclude the presence of micrometric grains of quartz because they were not observed in sample CM62 (see sample localisation in Suchéras-Marx et al., 2012). We also exclude bulk algal organic matter contamination because such organic matter is composed of Fe and Mn (Ho et al., 2003) and we only observe Fe and almost no Mn in the central area. We therefore confidently attribute this contamination to clay particles from the marlstones from which heterococcoliths were picked. Clays are generally composed of Si, Al, Fe, Mg, Ca, Na, and K as major elements, and contain traces of Ti, Cr, Cu, Zn, Rb and Sr (Köster, 1996); they are typically depleted in Ca and Sr in comparison to heterococcoliths (Stoll & Ziveri, 2004). High Fe/Ca was already proposed as a way to identify clay contamination in heterococcolith geochemistry studies (Halloran et al., 2009); with clay contamination being later confirmed by atomic force microscopy observations of sub-micron clay particles covering...
heterococcoliths (Skovbjerg et al., 2012) and secondary ion mass spectrometry (Prentice et al., 2014). In this study, Ti is particularly concentrated in small nodules with size ranging from 100 to 500 nm. In both heterococcoliths, Ti nodules are not correlated with Ca-rich spots but rather located at the junction between CaCO₃ crystals. Ti nodules could possibly be nanoscale Ti-rich clay particles, which size (down to 100 nm) is consistent with the clay particles observed by Skovbjerg et al. (2012). However, these Ti-rich clay nodules are different from the central area clay, highlighting at least two sources of contamination in our results.

Contamination from clays cannot be entirely avoided with the methodology we adopted to isolate heterococcoliths in the present study. We deliberately avoided chemical reagents in order to circumvent any potential cleaning contamination of the specimens. However, the advantage of the high-resolution mapping method adopted here is that it allows the unambiguous identification of contaminated zones, and therefore the selection of uncontaminated zones such as ROI 1 (Fig. 6) for further discussions on heterococcolith primary geochemical composition.

Diagenetic contribution

During diagenesis, heterococcoliths can be etched and/or overgrown. Overgrowth builds upon primary heterococcolith crystals (Adelseck et al., 1973), and tends to preserve the primary crystalline organisation. In Fig. 1B, the primary crystalline organisation of W. britannica is easily observable even if several monocrystals are thicker at the rim than at the inner tube due to overgrowth. During diagenesis, heterococcolith crystals larger than 1 µm and large heterococcoliths tend to overgrow whereas small crystals, small heterococcoliths and broken pieces of heterococcoliths tend to dissolve (Adelseck et al., 1973). Overgrowth of an heterococcolith from its etched parts was not observed and thus not further considered in this discussion. Dissolved CaCO₃ from heterococcoliths and other carbonate sources is mixed with the diagenetic fluids of the bulk rock, which is also composed of clay. Hence, the overgrown parts of heterococcoliths are likely to have different elemental compositions than the original one. In marlstones that contain clay, diagenetic fluids are typically enriched in metals. The absolute and relative i/Ca concentration maps of Wbrit1 and Wbrit2 show that heterococcoliths are enriched in K, Fe, Cu, Zn and Rb on the outermost rims that are interpreted as overgrown parts (Fig. 10). Those elements are incorporated in overgrown calcite on one or both sides of the heterococcoliths or between the proximal and distal shields (Fig. 10). In both heterococcoliths, radial overgrowth is also observed and is located between crystals, which form edges offering crystallization loci. These features define a specific pattern of overgrowth on the edges and between crystals, where K, Fe, Cu, Zn and Rb are concentrated.

One of the most intriguing observation in the present study was the distinctive distribution of Mn. Mn²⁺ has strong affinities with carbonate and may substitute Ca²⁺ during calcification (Paquette & Reeder, 1995; Astilleros et al., 2002). It is often considered as a diagenetic contaminant resulting from secondary incorporation in carbonate fossils (Boyle et al., 1983; Pena et al., 2008). In the present XRF maps, the distribution of Mn in Wbrit1 and Wbrit2 is closer to Ca and Sr than to any other element (Fig. 3) questioning the often-assumed diagenetic origin of Mn in calcite. The Mn and Mn/Ca maps (Figs. 3 and 7) display with great
detail an outer Mn ring with clockwise orientation as well as anticlockwise radial features, which are more visible in the upper right corner of each heterococcoliths. Assuming a proximal view of the heterococcolith, the former ring could correspond to the proximal shield with clockwise (dextrogyre) orientation of the calcite crystals while the latter could correspond to the distal shield with anticlockwise (laevogyre) orientation of the calcite crystals (Fig. 1A). This distinctive organisation suggests that Mn is more concentrated in one or the other shield. This could be due to abiotic diagenetic overgrowth on the shields surfaces and/or between both shields (Fig. 11), rather than zonal biological incorporation. A differential enrichment in Mn in only part of a given monocrystal seems indeed difficult to explain during heterococcolithogenesis. Abiotic experiments have shown that Mn-rich carbonate has affinities with \{10\overline{1}4\} crystal surfaces (Astilleros et al., 2002), which actually correspond to the orientations of shield surfaces in *Watznaueria* (Saruwatari et al., 2008). This suggests that Mn is present in heterococcolith in the form of Mn-CaCO$_3$ rather than Mn-oxides.

![Figure 10](image)

*Figure 10: Diagenetic model of K, Fe, Cu, Zn and Rb overgrowth based on the Wbrit1 and Wbrit2 Fe/Ca maps (Fig. 9). The overgrowth is concentrated on the shields borders, in the central area and some nods covering the proximal and distal shields. Overgrowth at the connection between R- and V-units is not shown on the lateral view scheme but was also observed in the Wbrit1 and Wbrit2 Fe/Ca maps.*

As a concluding remark to this part of the discussion, we recommend that heterococcoliths are carefully selected and cleaned prior to any analysis focusing on K, Mn, Fe, Ti, V, Cr, Cu, Zn.
and Rb elemental composition and/or their isotopes. Oppositely Ca, Sr, S, Cl and Br appear to be affected by neither clay contamination nor secondary diagenetic processes in the heterococcoliths studied here, and are therefore of potential use in palaeobiology and palaeoceanography.

![Image](image1)

Figure 11: Diagenetic model of Mn-rich carbonate overgrowth based on the Wbrit1 and Wbrit2 Mn/Ca maps (Fig. 7). The Mn-rich overgrowth is concentrated between the shields and at the connection between R- and V-units in the inner part of the heterococcoliths.

**Trace elements identified as potential environmental proxies**

*Possible salinity proxy based on chlorine and bromine in heterococcoliths*

The model traditionally used in calcite elemental geochemistry is based on cations, with Ca$^{2+}$ being replaced by other cations in the calcite lattice like Sr$^{2+}$ or Mg$^{2+}$ (Finch & Allison, 2007). Anion geochemistry in biogenic calcite, however, is poorly studied and there is, to our best knowledge, no model available for anion incorporation and exchange in calcite. Cl and Br are homogenously present in our maps. Ca$^{2+}$ and CO$_3^{2-}$ form the crystalline lattice of CaCO$_3$. The present data cannot directly determine whether Cl$^-$ and Br$^-$ actually replace CO$_3^{2-}$ in the calcite lattice. If Cl and Br did replace CO$_3^{2-}$ in the calcite lattice, their distribution in the heterococcolith should be similar to that of Ca, which is not the case. Because of the differences in Cl and Br distribution and that of Ca, we hypothesize instead that Cl and Br are impurities in interstitial nano-domains.
In the coccolithophore species *Coccolithus pelagicus*, a Cl\(^-\) inward-rectifying current (also applicable to Br\(^-\)) has been observed in the cell plasma membrane, but this ion channel does not mediate large osmo-regulation and likely equilibrates HCO\(_3^-\) uptake and membrane voltage (Taylor & Brownlee, 2003; Mackinder et al., 2011). HCO\(_3^-\) is then used by the cell for photosynthesis and heterococcolithogenesis (Mackinder et al., 2011). Hence, once HCO\(_3^-\) is consumed, we can confidently hypothesize that there is an influx of anions through the cell wall that compensates for the loss of HCO\(_3^-\) and activates the Cl\(^-\) channel again. This anion influx that compensates the loss of HCO\(_3^-\) is most likely sustained by Cl\(^-\), the most abundant anion in seawater. This could be a passive diffusive influx through the cell wall by osmotic regulation or active pumping. Inward and outward Cl\(^-\) fluxes would then result in an active cyclic movement of Cl\(^-\) leading to a constant [Cl\(^-\)] in the cell at the scale of heterococcolith formation. This hypothesis is likely also applicable to Br\(^-\), which has the same behaviour as Cl\(^-\) in coccolithophore cells (Taylor & Brownlee, 2003).

Because Cl and Br are most likely impurities in heterococcolith calcite, they are not biologically concentrated in the heterococcolith and would therefore depend on the coccolithophore cytoplasm [Cl\(^-\)] and [Br\(^-\)]. Since there is a direct link between seawater [Cl\(^-\)] and [Br\(^-\)] and coccolithophore cytoplasm [Cl\(^-\)] and [Br\(^-\)] as described earlier, heterococcolith Cl and Br may prove useful as seawater [Cl\(^-\)] and [Br\(^-\)] proxies. Cl\(^-\) is the most abundant anion in seawater, and salinity is directly related to seawater [Cl\(^-\)] (Adkins et al., 2002). Thus, if heterococcolith Cl and Br prove to be good seawater [Cl\(^-\)] and [Br\(^-\)] proxies, heterococcolith Cl and Br may in turn be an interesting tool in the establishment of a salinity proxy.

**Record of heterococcolithogenesis based on sulfur**

The strong similarity between the distribution of Cl and S in heterococcolith could lead to the hypothesis that heterococcolith S/Ca has potential as a seawater [S] proxy, an important element in Earth geochemical dynamics (Berner, 2006). Unfortunately, S is present in many biological processes occurring inside the cell during heterococcolithogenesis, involving polysaccharides (PS) that transport Ca\(^{2+}\) to a base plate where the heterococcolith is built (Westbroek et al., 1983). In *Pleurochrysis carterae* and *Emiliania huxleyi* for instance, the polysaccharide known as PS3 is a sulfated galacturonomannan that is active in crystal growth and shaping (Marsh et al., 2002). Some polysaccharides have been observed trapped inside the heterococcolith calcite lattice (Takahashi et al., 2002). Consequently, we interpret the presence of S as PS residues inside heterococcoliths. Because of the disequilibrium between heterococcoliths and seawater caused by biological concentration during heterococcolith formation, S/Ca is not useful as a tracer for seawater composition. The presence of S in *W. britannica* nonetheless suggests that coccolithophores used sulfated PS for heterococcololithogenesis for at least 170 Myr. This is of interest because further isotopic analysis of S trapped within heterococcoliths may deliver palaeobiological information on heterococcolithogenesis.

**New insights of Sr/Ca in W. britannica**

In both Wbrit1 and Wbrit2, Sr/Ca maps display detailed features that follow *W. britannica's* crystalline organisation (Figs. 1, 7 and 12). Sr/Ca distribution perfectly reflects the arrangement of calcite crystals in the V-units of the mid-tube elements. Sr/Ca distribution in
calcite crystals in the R-units of the rim are less distinguishable, a result of overlapping of the R-unit crystals in the distal shield and/or the fact that the shields' crystals have different orientations, i.e. clockwise for the proximal shield and anti-clockwise for the distal shield in distal view (Fig. 1A). Conversely, limits between mid-tube elements are easily identifiable because those elements are made of V-unit monocrystals with vertical walls.

Although Sr/Ca maps clearly depict the organisation of calcite crystals in heterococcoliths, the Sr/Ca ratio varies only slightly between R- and V-units, with a mean value of 0.35±0.14 (1σ) mmol/mol and 0.36±0.15 (1σ) mmol/mol (Fig. 12). This important result confirms previous observations (Stoll and Ziveri, 2004) that the partitioning coefficient of strontium DSr between calcite and cytoplasm does not depend on the crystal growth direction. Despite the differences in crystal axes and shapes between V- and R-units (Saruwatari et al., 2008; Saruwatari et al., 2011), the factor controlling Sr incorporation is the elongation along the c-axis in both V- and R-units (Payne et al., 2006).

Figure 12: Identification of the W. britannica Wbrit1 and Wbrit2 crystal organisations in the Sr/Ca maps (in mmol/mol) and Sr/Ca transects showing the low variations in Sr/Ca along the minor axis.
Sr/Ca in *W. britannica* is twice to 12 times (Table 1) lower than in modern and Cenozoic species, which are characterized by a Sr/Ca ranging from 0.8 to 4 mmol/mol (Stoll & Bains, 2003; Stoll & Ziveri, 2004; Dedert et al., 2012; Dedert et al., 2014; Prentice et al., 2014). In one hand, Sr/Ca is homogenous in Wbrit1 and Wbrit2 (Fig. 12). On the other hand, calcite overgrowth in heterococcoliths is not homogeneous (Fig. 1B) and is documented to lower the original Sr/Ca (Dedert et al., 2014; Prentice et al., 2014, ). Hence, we consider that calcite overgrowth in Wbrit1 and Wbrit2 has a minor to negligible impact on Sr/Ca. In the absence of published *W. britannica* Sr/Ca to compare our results with, two hypotheses explaining the low Sr/Ca of our *W. britannica* are presented and could both occur together: i) the cellular physiology of Mesozoic coccolithophores and/or ii) a low cellular growth rate.

<table>
<thead>
<tr>
<th>Elemental ratio</th>
<th>Palaeoceanographic proxy</th>
<th>Palaeobiologic proxy</th>
<th>Diagenetic or sedimentary contamination</th>
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</thead>
<tbody>
<tr>
<td>S/Ca</td>
<td>Sulphated polysaccharides</td>
<td></td>
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<tr>
<td>Cl/Ca</td>
<td>Salinity ?</td>
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<td></td>
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<tr>
<td>K/Ca</td>
<td>Diagenesis and clays</td>
<td></td>
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<tr>
<td>Ti/Ca</td>
<td>Diagenesis and clays</td>
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<td>V/Ca</td>
<td>Diagenesis and clays</td>
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<td>Cr/Ca</td>
<td>Diagenesis and clays</td>
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<td>Mn/Ca</td>
<td>Diagenesis</td>
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<td>Fe/Ca</td>
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<td>Cu/Ca</td>
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<td>Zn/Ca</td>
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<td>Br/Ca</td>
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<td>Rb/Ca</td>
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<tr>
<td>Sr/Ca</td>
<td>Productivity</td>
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*Table 2: Concise table resuming the interest of the different element ratios presented for palaeoceanographic proxy, palaeobiologic proxy and/or diagenetic or sedimentary contamination.*

*Watznaueria* are the most abundant heterococcoliths in the fossil record from the Middle Jurassic until the end of the Cretaceous and *W. britannica* is particularly abundant in Middle and Upper Jurassic sediments. Sedimentary evidence (microlaminae) of blooms of this species is documented (Lees et al., 2004) and an affinity for eutrophic conditions was proposed (Lees et al., 2006; Suchéras-Marx et al., 2015) without distinguishing morphotypes. Such high abundances in the fossil record are more likely related to a ubiquitous and eurytopic ecology. In comparison to modern environments, *W. britannica* in the Middle Jurassic is comparable to *E. huxleyi* in term of dominance in the nannofossil assemblage even if their morphology, size and size range are completely different. *W. britannica* has very low Sr/Ca with respect to *E. huxleyi* and other modern and Cenozoic species. These exceptionally low Sr/Ca values for the dominant Mesozoic placolith could be related to different cellular physiology between Mesozoic forms and Cenozoic to modern forms. The Cretaceous-Paleogene mass extinction, which has led to the most important turnover in coccolithophore evolutionary history (Bown, 2005), could have led to the extinction of the most Sr-depleted heterococcoliths replaced by coccolithophore producing Sr-enriched heterococcoliths. There is a need to establish the behaviour and geochemistry of Mesozoic nannofossils somewhat independently of what we assume from modern/Cenozoic taxa. In particular, new Sr/Ca
analyses on Mesozoic survivors still present in modern oceans, namely *Tergestiella adriatica* (Hagino *et al.*, 2015), could help to test this hypothesis.

Both *W. britannica* specimens studied here belong to a large-sized morphotype and different studies focusing on the palaeoecology of this species have shown that large-size morphotypes were adapted to stable and relatively oligotrophic environmental conditions (Olivier *et al.*, 2004; Giraud *et al.*, 2006; Giraud *et al.*, 2009). Today, extent heterococcolith species adapted to oligotrophic conditions (e.g. *Umbilicosphaera sibogae sibogae*) are also characterized by low cell growth rates and low D Sr (Stoll *et al.*, 2002), leading to low Sr/Ca. This also applies to heterococcoliths that are usually characterized by high growth rates during the stationary phase of the population growth (Rickaby *et al.*, 2002). Low Sr/Ca in *E. huxleyi* was recorded in nutrient-poor culture experiments (Stoll *et al.*, 2007b), stationary-phase population growth in culture experiments (Rickaby *et al.*, 2002), and in the Eastern Mediterranean Sea, which is considered as a “nutrient desert” (Auliaherliaty *et al.*, 2009). Hence, low Sr/Ca in our *W. britannica* suggests an adaptation to oligotrophic conditions of the large morphotype, as proposed by Giraud *et al.* (2006, 2009). A summary of potential uses of heterococcolith elemental geochemistry for palaeoceanography, palaeobiology and/or identification of diagenesis and clay contamination is given in Table 2.

**CONCLUSION**

We have confidently identified and mapped 14 elements in two Middle Jurassic *W. britannica* heterococcoliths using nanoscale XRF mapping. Amongst those 14 elements, K, Ti, V, Fe, Cu, Zn and Rb are linked to clay contamination and diagenetic overprint whereas Mn is present in calcite overgrowth, likely in the form of Mn-rich CaCO 3. Presence of S is interpreted as a residue of sulfated polysaccharides, which mediate crystal growth and shape in some modern coccolithophores. Cl and Br are also present in heterococcoliths, more likely in interstitial nano-domains than as substitutes for CO 3 2-. Heterococcolith Cl content may be interesting as a salinity proxy, but this would require demonstration of a linear correlation between [Cl] in heterococcoliths and [Cl] in seawater in culture experiments. Finally, Sr/Ca is unusually low (i.e. 0.35 mmol/mol) in the analysed *W. britannica* compared to Cenozoic/modern species. This low Sr/Ca could be related to different cellular physiology in Mesozoic coccolithophores, or it could also be the result of low growth rates in those individuals. The present study on fossilised heterococcoliths helps to identify the relative contributions of contamination, diagenesis and biology to the final elemental composition. This study, alongside Prentice *et al.* (2014), provides new insights in heterococcolith chemical composition and applicability to palaeoceanography but further effort is needed in terms of culture studies, cell biology and crystallographic studies.

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