High frequency Barium profiles in shells of the Great Scallop Pecten maximus: a methodical long-term and multi-site survey in Western Europe
Aurélie Barats, David Amouroux, Laurent Chauvaud, Christophe Pécheyran, Anne Lorrain, Julien Thébault, T.M. Church, Olivier F.X. Donard

To cite this version:
Aurélie Barats, David Amouroux, Laurent Chauvaud, Christophe Pécheyran, Anne Lorrain, et al.. High frequency Barium profiles in shells of the Great Scallop Pecten maximus: a methodical long-term and multi-site survey in Western Europe. Biogeosciences, European Geosciences Union, 2009, 6 (2), pp.157-170. <hal-00484941>
High frequency Barium profiles in shells of the Great Scallop *Pecten maximus*: a methodical long-term and multi-site survey in Western Europe

A. Barats1,*, D. Amouroux1, L. Chauvaud2, C. Pécheyran1, A. Lorrain3, J. Thébault2,**, T. M. Church4, and O. F. X. Donard1

1Laboratoire de Chimie Analytique Bio-Inorganique et Environnement (LCABIE) – Institut Pluridisciplinaire de Recherche sur l’Environnement et les Matériaux (IPREM), UMR 5254 CNRS – Université de Pau et des Pays de l’Adour (UPPA), Hélioparc Pau-Pyrénées, 2 av. du Président Angot, 64053 Pau Cedex 9, France
2Laboratoire des sciences de l’Environnement MARin (LEMAR) – Institut Universitaire Européen de la Mer (IUEM), UMR 6539 CNRS – Université de Bretagne Occidentale (UBO), Technopole Brest-Iroise, Place Nicolas Copernic, 29280 Plouzané, France
3Institut de Recherche pour le Développement (IRD) – Laboratoire des sciences de l’Environnement MARin (LEMAR), Centre IRD de Brest, BP 70, 29280 Plouzané, France
4College of Marine and Earth Studies, University of Delaware, Newark, Delaware, 19716-3501, USA
*now at: Laboratoire de Radiochimie et des Sciences Analytiques et Environnement (LRSAE/E 1175) – Université de Nice Sophia Antipolis (USNA), 28 avenue Valrose, 06108 Nice Cedex 2, France
**now at: Johannes Gutenberg University, Institute of Geosciences, Department of Applied and Analytical Paleontology, Johann-Joachim-Becher-Weg 21, 55099 Mainz, Germany

Received: 23 July 2008 – Published in Biogeosciences Discuss.: 10 September 2008
Revised: 4 December 2008 – Accepted: 14 January 2009 – Published: 10 February 2009

Abstract. Skeletal barium/calcium ([Ba]/[Ca])shell ratios were measured every third daily striae in 39 flat valves of the Great Scallop *Pecten maximus* collected in temperate coastal environments of Western Europe. A methodical evaluation of the ([Ba]/[Ca])shell ratio was performed for the first time and demonstrates that ([Ba]/[Ca])shell profiles are reproducible for several scallop individuals from the same population (2-year old; 3 shells/year), over a 7-year period (1998–2004), and from different coastal environments in France (42°–49° N). As previously determined in the shells of other bivalve species, ([Ba]/[Ca])shell profiles generally exhibited a background ratio punctuated by two transient maxima occurring in early and late summer. Background partition coefficient (DBa=0.11±0.03, in 2000) was similar to that previously reported in *P. maximus* shells, suggesting a direct shell uptake of dissolved seawater Ba (Gillikin et al., 2008). The 7-year survey in the Bay of Brest of the high frequency ([Ba]/[Ca])shell profiles in the scallop’s shell was exploited to better constrain both the occurrence and the amplitude of the summer Ba relative enrichments as influenced by environmental processes. Seawater Ba contents in 2000 underlined significant particulate Ba inputs at the sediment water interface (SWI) during ([Ba]/[Ca])shell peak events. These Ba inputs are thus suggested to be subsequently induced by a pelagic biogenic process, which mainly occurs under summer post-bloom conditions in relationship to the cycling of particulate organic matter and associated Ba. The long term survey reveals that such pelagic Ba cycling processes are responsible for particulate Ba inputs to the sediment water interface (SWI). Subsequent indirect Ba uptake by the bivalve results in higher ([Ba]/[Ca])shell ratios, in that archived Ba within the shell cannot be used as a direct paleo productivity tracer. Our methodical approach, based on a multi-year and multi-site-survey of ([Ba]/[Ca])shell ratio in Scallop bivalves, allows us to establish the potential application of such high frequency archives for further biogeochemical and ecological investigations of bivalves in the coastal environment.

Correspondence to: D. Amouroux (david.amouroux@univ-pau.fr)
1 Introduction

Since 1965, trace elements in mollusc shells were investigated because bivalves form in their shells successive calcium carbonate layers, potential archives of conditions experienced by the organism during its life time (Dodd, 1965; Lorens and Bender, 1980; Klei et al., 1996a, 1996b; Stecher et al., 1996). The main aim of these studies was to use trace element records in shells as a bio indicator of environmental conditions. A recent increasing number of studies have investigated Ba content in biogenic carbonates, such as corals, foraminifera, and otoliths (Lea et al., 1989; Lea and Martin, 1996; McCulloch et al., 2003; Sinclair and McCulloch, 2004; de Vries et al., 2005; Sinclair, 2005). Coastal waters are enriched in Ba in the low to mid salinity ranges during estuarine mixing by either freshwater inputs of dissolved Ba or Ba release from river-born particulate phases (Coffey et al., 1997; Shaw et al., 1998; McCulloch et al., 2003). Alternatively Ba input can occur from the exchange of Ba-rich ground waters and pore waters within the tidal prism (Shaw et al., 1998). Coralline Ba content was reported as a proxy for discharge and/or sediment load in riverine systems (Sinclair and McCulloch, 2004; McCulloch et al., 2003). Some sharp Ba peaks in corals cannot however be assigned to any tracers of river discharge (Sinclair, 2005). Barium incorporation into carbonate parts of larval protoconchs or statoliths was also demonstrated to be promoted at lower temperature (Zacherl et al., 2003). In recent studies, there was much attention to the skeletal Ba content in mussels (Mytilus edulis; Vander Putten et al., 2000; Gillikin et al., 2006), Isognomon ephippium; (Lazareth et al., 2003)), clams (Mercenaria mercenaria, Spisula solidissima, Arctica islandica, Saxidomus giganteus, Asotic Corbula fluminea; Fritz et al., 1990; Stecher et al., 1996; Epplé, 2004; Gillikin et al., 2005, 2008), and scallops (Pecten maximus, Comptopallium radula, Argopecten purpuratus; Lorrain, 2002; Thébault, 2005; Gillikin et al., 2008). In all these studies, [Ba/Ca]_{shell} profiles were usually characterized by a relatively stable background ratio interrupted by sharp intense maxima. A simple interpretation achieved with several investigated bivalves (Mercenaria mercenaria, Mytilus edulis, Isognomon ephippium, Ensis siliqua) is also limited by an approximate dating of the shell record. In both lab and field experiments, the background [Ba/Ca]_{shell} in the common mussel shell (Mytilus edulis) was demonstrated to track the [Ba/Ca]_{water} aqueous concentration ratio (Gillikin et al., 2006). Considering the inverse relationship between [Ba/Ca]_{water} and salinity, background [Ba/Ca]_{shell} was considered as a specific indicator of salinity in estuarine environments (Gillikin et al., 2006). Barium was also established as a paleo productivity tracer in marine waters, suspended particles and sediments (Dymond and Collier, 1996; Paytan and Kastner, 1996). Stecher et al. (1996) postulated first that, in Mercenaria mercenaria and Spisula solidissima shells, these [Ba/Ca]_{shell} peaks might be induced by the ingestion of high levels of particulate Ba in estuarine waters. Bivalves are non-selective filter feeders and can assimilate these Ba-rich particles. Once inside the digestive tract of the bivalve, Ba may be metabolised, at least in part, by being shunted to the extra-pallial fluid (EPF) and sequestered into the shell (Stecher et al., 1996; Vander Putten et al., 2000; Lorrain, 2002; Gillikin et al., 2006). Time resolved [Ba/Ca]_{shell} peaks were suggested to be related to phytoplankton blooms (Stecher et al., 1996; Vander Putten et al., 2000; Lazareth et al., 2003). These Ba-rich particles were supposed to originate from diatom blooms as either biogenic matter, or barite precipitates (Stecher et al., 1996). Gillikin et al. (2006, 2008) indicated that these [Ba/Ca]_{shell} peaks could not be used as a direct proxy of [Ba/Ca]_{water} or phytoplankton production. These peaks were rather induced by ingestion of barite particles inputs to the sediment water interface (SWI) during the phytoplankton decay (Gillikin et al., 2006). Barium is recognized to be accumulated at high levels in phytoplankton, both in diatoms and in dinoflagellates (Fisher et al., 1991). A relatively large pool of labile Ba can be rapidly released during plankton decomposition and acts as the main source of Ba for barite formation in supersaturated microenvironments (Bishop, 1988; Ganeshram et al., 2003). Elevated levels of Ba, both as suspended particulate Ba and sedimentary barite occurs in high primary productivity oceanic regions (Dehairs et al., 1980; Bishop, 1988; Dehairs et al., 1991, 1997). Most probably, barite formation results from its passive precipitation through sulphate enrichment and/or release in barium containing biogenic material (Bishop, 1988; Stecher and Kogut, 1999; Jeandel et al., 2000; Ganeshram et al., 2003; Dehairs et al., 2000). The formation of specific aggregates or Ba-rich micro-environments is responsible for the Ba enrichment in biogenic particles (Bishop, 1988). These aggregates are related to the diatom extracellular polymeric substances (EPS) produced during summer blooms and their settling (Stecher and Kogut, 1999; Thornton, 2002). The subsequent vertical mixing and diatom decay provide a rapid flux and large abundance of biogenic particles transported down to the SWI (Sarthou et al., 2005). This was suggested to induce a sudden pulse of Ba to the epibenthic environment, subsequently ingested by molluscs to be incorporated into shells and recorded by ([Ba]/[Ca]_{shell} maxima (Gillikin et al., 2006). Phytoplankton Ba uptake and barite formation are however not sufficient to explain the vertical flux of Ba in oceanic waters (Sternberg et al., 2005). Barium is also scavenged in the presence of diatom biogenic particles or by adsorption onto mineral oxides (i.e. Fe, Mn) that contribute to significant Ba inputs to the SWI (Sternberg et al., 2005).

This study focuses on Ba content in the Great Scallop Pecten maximus (L.) to further establish how these bivalves can provide time resolved ([Ba]/[Ca]_{shell} profiles (daily scale) and valuable archives of environmental changes in temperate environments (Lorrain et al., 2000; Chauvaud et al., 1998, 2005; Lorrain et al., 2004; Barats 2000; Barats et al., 2007, 2008; Gillikin et al., 2008). High frequency
profiles of Ba content in the calcitic shell can be determined for each individual larva using laser ablation – inductively coupled plasma – mass spectrometry (LA-ICP-MS) (Barats et al., 2007). In this paper, the significance of $([\text{Ba}]/[\text{Ca}]_{\text{shell}}$ profiles in *P. maximus* is first evaluated for both background concentration and episodic sharp peaks in a single scallop population from the Bay of Brest (northwest France), among different coastal sites in western Europe, and over a 7-year period (1998–2004) in the Bay of Brest. Second, the demonstrated reproducibility, recurrence and ubiquity of $([\text{Ba}]/[\text{Ca}]_{\text{shell}}$ profiles is further utilized to clarify the biogeochemical processes influencing both the background and episodic sharp peaks of $([\text{Ba}]/[\text{Ca}]_{\text{shell}}$ content in scallop bi valves.

2 Materials and methods

2.1 Study sites and scallop sampling

Different sampling areas in Western Europe (Bay of Seine, Bay of Brest, Quiberon peninsula and Belle Ile, Ria de Vigo) were considered because they represent ecosystems with their own ecological characteristics (Barats, 2006). Shells originating from the Bay of Seine (France, English Channel, 49°30’N, 0°30’W), Belle Ile and Quiberon area (France, Bay of Biscay, respectively, 47°20’N, 3°10’W and 47°30’N, 3°00’W) were sampled in open coastal ecosystems influenced by large estuarine inputs (respectively, Seine and Loire rivers) (Fig. 1). Shells sampled in coastal environments such as the Ria de Vigo (Spain, Atlantic Ocean, 42°10’N, 8°50’W) or the Bay of Brest (France, Iroise Sea, Roscanvel bank, 48°20’N, 4°30’W) were subjected to low anthropogenic inputs. In this study, the main sampling site is the Bay of Brest (Fig. 1): a shallow embayment with more than a half of its surface (180-km$^2$) and with depth of less than 5 m. The hydrodynamics of this ecosystem is mainly influenced by tidal exchange with the Iroise Sea, but also by freshwater input mainly by two small riverine watersheds (80%): the Aulne (1842 km$^2$) with monthly flow rate ranging from 2 to 52 m$^3$/s and averaging 22 m$^3$/s, and the Elorn (402 km$^2$) with monthly flow rate ranging from 1 to 12 m$^3$/s and averaging 5.6 m$^3$/s. The sampling area (Roscanvel) is 30 m deep where the bottom consists of mixed sediments (mud, sand) and selected because scallop density is one of the highest in the Bay of Brest (Chauvaud et al., 2000, 2005).

Three replicates of live juvenile *P. maximus* specimens were collected for each year and site (except in 2002 in the Bay of Brest, n=2) during the late autumn period by SCUBA diving in the Bay of Brest and by dredging for other sites. Only the juvenile stage of the shells (the second year of growth) was examined because it exhibits the longest annual growth period. An exception was made for a three-year old *P. maximus* specimen collected in the Bay of Brest in 2003 to examine its third year of growth. The shells were isolated and cleaned by submerging in glacial acetic acid (Merck, analytical grade) for 60 s to remove bio-fouling, rinsed with Milli-Q water ($R=18.2$ MΩ/cm) and dried. For convenience in visualizing the growth striae, only the flat valve was considered for LA-ICP-MS analysis. A 45×10 mm$^2$ cross section was cut from each shell using a diamond saw to fit into an ablation chamber.

2.2 Determination of Ba concentrations in shells by Laser Ablation – Inductively Coupled Plasma – Mass Spectrometry (LA-ICP-MS)

Quantitative analyses of 39 shells were performed by LA-ICP-MS method consisting of coupling a UV laser ablation unit (LSX 100 UV 266 nm, Cetac Tech.) to an ICP-MS (X7 serie, Thermo Fisher). The methodology and validation were described elsewhere and summarized here (Barats et al., 2007). Samples were analysed during 2 min at a scan speed rate of 5 µm/s. For shell analyses, a pre cleaning ablation of the calcite surface was conducted to avoid surface contamination (Stecher et al., 1996; Vander Putten et al., 1999; Lazareth et al., 2003; Wyndham et al., 2004). This pre cleaning step consisted in a quick (around 20 s) pre-ablation of shell surface at a rate of 50 µm/s. Twenty-eight isotopes including $^{135,137,138}$Ba, $^{55}$Mn and $^{43}$Ca were analysed. The intensity of each isotope signal was systematically normalized against the $^{43}$Ca signal to compensate for instrumental drift and instability. An external calibration was performed with lab prepared CaCO$_3$ standards containing Ba concentrations ranging from 0.005 to 19.6 µg/g (Barats et al., 2007). The Ba calibration curve displayed a good linearity with a regression coefficient $r^2$ above 0.998 (Barats et al., 2007). The detection limit was about 3.3 ng/g, and the relative standard deviations obtained for both analytical repeatability (5 integration zones during 1 analysis) and reproducibility (5 successive independent analyses) were below 5% (Barats et al., 2007).

Fig. 1. Maps of the investigated area (Atlantic Coast, France) showing the sampling zones: (a) the Bay of Seine (49°30’N, 0°30’W), the Bay of Brest (48°20’N, 4°30’W), Belle Ile (47°20’N, 3°10’W) and Quiberon (47°30’N, 3°00’W); (b) the Bay of Brest in detail with all the investigated sites (Roscanvel for the sampling, and the others for regular measurements of environmental parameters).
Analytical performances obtained for Mn were previously presented (Barats et al., 2007, 2008). The ([Ba]/[Ca])shell or ([Mn]/[Ca])shell ratios were calculated by dividing shell Ba or Mn concentrations by the Ca concentration in the shell (40%) and expressed in µmol/mol. Mollusc shells were recognized to contain less than 5% of organic matrix in calcite or aragonite shells (Carroll et al., 2006; Levi-Kalisman et al., 2001). The shell is thus composed by minimum 95% of CaCO₃, i.e. a minimum shell Ca content of 38%. In comparison with the Ca concentration of 40%, the error is only of 5%. This study thus used a shell Ca content of 40% because scallop shells have a calcite structure. While the exact percentage of organic matter remains unknown, yet likely less than 5%, the analytical error will be negligible. A recent study of Takesue et al. (2008) revealed higher organic content in clams (19%), but they demonstrated also that Ba assay in shell content was unchanged by the removal of organic matter. This conclusion underscores that while Ba occurs exclusively in shell aragonite of clams, the analytical result agrees with the use of Ca concentration about 40% to calculate our ([Ba]/[Ca])shell ratios.

Shell analyses were performed each third striae during the shell growth period (from April to November) to limit the analysis time to approximately 5 h per shell. A date of formation was assigned to each ablated sample, by backdating from the harvest date, based on the daily periodicity of the striae formation in P. maximus. An evaluation of the shell growth rate (dorso-ventral linear extension of the shell per unit time), expressed in µm/d, was also performed for each shell by measuring distances between successive striae (growth increment width) using an image analysis technique (Chauvaud, 1998). For each shell analysis, a profile of shell growth rate and a profile of ([Ba]/[Ca])shell ratio were obtained. For each year and site, a mean shell growth rate profile and a mean ([Ba]/[Ca])shell were then defined averaging the results of 3 shells from a same scallop population. Due to the uncertainties from the backdating of analysed striae and those from the intershell comparison, the uncertainty in the date for mean shell profiles was estimated to range from 1 to 7 days.

2.3 Environmental monitoring database

([Ba]/[Ca])shell time series were compared with environmental variables that may influence Ba biogeochemical cycle at the SWI. The Bay of Brest (Roscanvel station) was mainly investigated because of the regular environmental monitoring and the long time series obtained by shell analyses (1998 to 2004). Regular measurements (2–3 days resolution) of Ba and Mn concentrations in dissolved (<0.6 µm, Nuclepore) and particulate phases from bottom seawater (1 m above the SWI) were also performed at the sampling site (Roscanvel) in 2000 (from February to December). Filtrated samples (dissolved seawater phase) were acidified in 2% HNO₃ (69–70% Suprapur, Merck) and diluted 30 times to determine concentrations of dissolved elements. Two internal standards were added (Y and Bi) in diluted samples. Elemental concentrations (Ba, Mn, ...) were then determined by ICP-MS (X7 series, Thermo Fisher) by an external and internal calibration. Particulate samples (filters) were digested in closed vials (Savillex, PFA) by an acid mixture (1.5 ml HCl, 1 ml HNO₃, 0.5 ml HF; suprapur quality) at 95°C during one night. The digested samples were evaporated to dryness at 110°C under a laminar hood to eliminate the matrix. The residues were dissolved in 2.5 ml of nitric acid solution (HNO₃ 2%) and the volumes were adjusted to 11 ml. These samples were analyzed by ICP-AES (THERMO Optek Iris Advantage, Thermo Fisher) at the Royal Museum for Central Africa (Belgium). Samples were spiked with a mixture of internal standards (Au, Y) before its introduction into the spectrometer to compensate instrumental drift and instability. All these experiments were performed under clean conditions (clean vials, laminar flood, blank procedure...).

([Ba]/[Ca])shell time series were also compared with an environmental database: hydrological (S, T, suspended particulate matter), chemical (O₂, NO₃⁻, NO₂⁻, NH₄⁺, PO₄³⁻, Si(OH)₄, particulate organic carbon (POC) or nitrogen (PON)) and biological parameters. Biological parameters include total phytoplankton biomass, as reflected by chlorophyll-a and abundance of phytoplankton species. These measurements were performed in the Bay of Brest at the “Saine Anne” site near to Roscanvel sampling zone, as part of monitoring activities lead by the Institut Universitaire Européen de la Mer (http://www.univ-brest.fr/IUEM/observation/observation_riroise.htm; Fig. 1). All these measurements were performed at a weekly resolution. Nutrient dynamics in SOMLIT station were previously demonstrated to reflect environmental conditions in Roscanvel (Lorrain, 2002). Phytoplankton composition (a hundred species) were monitored from 1998 to 2002 at Lanvéoc (48°18'N, 4°27'W), located near to Roscanvel, as a part of the littoral environment monitoring program lead by the Institut Français de Recherche pour l’Exploitation de la MER (IFREMER database, http://www.ifremer.fr/envlif/), and from 2003 to 2004, at the seawater surface (0–1 m) in a reference SOMLIT station (Service d’Observation en Milieu Littoral: 48°22’N, 4°33’W) (Fig. 1). Phytoplankton identifications were performed in surface seawater (0–1 m) and every 15 days, except in 2004 (weekly resolution).

2.4 Statistical analyses

A statistical data treatment was performed to highlight environmental parameters influencing the occurrence and the amplitude of ([Ba]/[Ca])shell peaks. ([Ba]/[Ca])shell maximum events from their beginning to their end, usually last 20 days, whatever year studied. Averaged environmental parameters were thus examined over different periods: 4 weeks before–1 week after (−4 W+1 W), 3 weeks before–1 week after (−3 W+1 W), 2 weeks before–1 week after...
Table 1. Intershell comparison of ([Ba]/[Ca])shell profiles and interannual comparison (1998–2004) of both mean background and mean peak intensity (named by the time order of the peak) of ([Ba]/[Ca])shell ratios (n=3) in scallops from the Bay of Brest.

<table>
<thead>
<tr>
<th>Year</th>
<th>1 vs 2</th>
<th>1 vs 3</th>
<th>2 vs 3</th>
<th>p</th>
<th>Mean</th>
<th>SD</th>
<th>%RSD</th>
<th>Date (±3 d)</th>
<th>Mean</th>
<th>SD</th>
<th>%RSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1998</td>
<td>0.72</td>
<td>0.78</td>
<td>0.95</td>
<td>&lt;0.05</td>
<td>0.526</td>
<td>0.074</td>
<td>14</td>
<td>peak 1 12/06/1998</td>
<td>4.67</td>
<td>1.90</td>
<td>41</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.38</td>
<td>0.99</td>
<td>44</td>
<td>peak 2 04/09/1998</td>
<td>0.898</td>
<td>0.314</td>
<td>35</td>
</tr>
<tr>
<td>1999</td>
<td>0.92</td>
<td>0.95</td>
<td>0.96</td>
<td>&lt;0.05</td>
<td>0.377</td>
<td>0.031</td>
<td>8</td>
<td>peak 1 18/06/1999</td>
<td>1.27</td>
<td>0.48</td>
<td>38</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.12</td>
<td>0.78</td>
<td>0</td>
<td>peak 2 19/07/1999</td>
<td>2.12</td>
<td>0.84</td>
<td>40</td>
</tr>
<tr>
<td>2000</td>
<td>0.48</td>
<td>0.22</td>
<td>0.82</td>
<td>&lt;0.09</td>
<td>0.561</td>
<td>0.057</td>
<td>10</td>
<td>peak 1 23/07/2000</td>
<td>1.03</td>
<td>0.03</td>
<td>3</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.55</td>
<td>1.01</td>
<td>15</td>
<td>peak 1 26/08/2000</td>
<td>0.73</td>
<td>0.04</td>
<td>5</td>
</tr>
<tr>
<td>2001</td>
<td>0.18</td>
<td>0.24</td>
<td>0.46</td>
<td>&lt;0.2</td>
<td>0.657</td>
<td>0.101</td>
<td>19</td>
<td>peak 1 22/07/2001</td>
<td>0.971</td>
<td>0.431</td>
<td>44</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.42</td>
<td>0.134</td>
<td>32</td>
<td>peak 2 31/07/2001</td>
<td>1.39</td>
<td>0.91</td>
<td>65</td>
</tr>
<tr>
<td>2002</td>
<td>0.88</td>
<td>–</td>
<td>–</td>
<td>&lt;0.05</td>
<td>0.533</td>
<td>0.102</td>
<td>19</td>
<td>peak 1 14/06/2002</td>
<td>1.86</td>
<td>0.60</td>
<td>32</td>
</tr>
<tr>
<td>(n=2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.31</td>
<td>0.134</td>
<td>25</td>
<td>peak 2 01/07/2002</td>
<td>1.31</td>
<td>0.42</td>
<td>32</td>
</tr>
<tr>
<td>2003</td>
<td>0.93</td>
<td>0.96</td>
<td>0.87</td>
<td>&lt;0.05</td>
<td>0.535</td>
<td>0.134</td>
<td>25</td>
<td>peak 1 26/07/2002</td>
<td>1.55</td>
<td>0.18</td>
<td>12</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.18</td>
<td>0.55</td>
<td>19</td>
<td>peak 2 20/08/2002</td>
<td>2.83</td>
<td>0.55</td>
<td>19</td>
</tr>
<tr>
<td>2004</td>
<td>0.44</td>
<td>0.78</td>
<td>0.30</td>
<td>&lt;0.05</td>
<td>0.745</td>
<td>0.175</td>
<td>24</td>
<td>peak 1 20/07/2003</td>
<td>3.80</td>
<td>0.74</td>
<td>20</td>
</tr>
<tr>
<td>(n=3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.03</td>
<td>0.12</td>
<td>7</td>
<td>peak 2 26/08/2003</td>
<td>1.72</td>
<td>0.12</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.48</td>
<td>0.03</td>
<td>30</td>
<td>peak 1 18/06/2004</td>
<td>2.25</td>
<td>0.99</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.31</td>
<td>0.03</td>
<td>30</td>
<td>peak 2 05/09/2004</td>
<td>1.26</td>
<td>0.38</td>
<td>30</td>
</tr>
</tbody>
</table>

(−2 W+1 W), 1 week before–1 week after (−1 W+1 W) the maximum ([Ba]/[Ca])shell ratios. The choice of such integration periods around the maximum of ([Ba]/[Ca])shell was consistent with the lower resolution obtained for most of environmental parameters and the uncertainty due to the back-dating of shell striae. These integration periods took into account an eventual lag between changes in the water column, at the SWI, and its further transcription within the shell. This allowed a good overlap between ([Ba]/[Ca])shell maxima and environmental datasets. The influence of transient events was tested taking into account the maximum value during the considered period.

Bravais Pearson tests and multiple regression analyses were performed with Stabox Pro software for Windows Ver. 6 (Grimmer Software, France). Bravais Pearson tests were used to emphasize univariate correlations between the amplitude of ([Ba]/[Ca])shell maxima and other variables. Multiple regression analyses were considered to express the amplitude of ([Ba]/[Ca])shell maxima according to several independent factors. The relationships were significant if the probability was below p<0.05, and moderate-to-strong if r²>0.7. If all the ([Ba]/[Ca])shell maxima agreed with the relationship, the Cook’s distance was below $d_{\text{cook}}<1$.

3 Results and discussion

3.1 Definition of a general pattern for ([Ba]/[Ca])shell profiles

3.1.1 Reproducibility of background and episodic maxima of ([Ba]/[Ca])shell ratios among a single scallop population

The variations of ([Ba]/[Ca])shell ratios were examined in 2003, for 2-year old shells and for 3-year old shells from the Bay of Brest (Fig. 2). Two-year old scallops, analysed for its juvenile stage, reveal a high reproducibility of ([Ba]/[Ca])shell profiles (r²>0.78, p<0.05, n>60) (Table 1, Fig. 2). Background ([Ba]/[Ca])shell ratio was obtained after the exclusion of peculiar data (i.e. Ba/Ca peaks) exhibiting higher ratio than the average values plus 3 times the standard deviation. A mean ([Ba]/[Ca])shell profiles exhibits a background ratio of 0.535±0.134 μmol/mol, and two significant enrichments from July to August (respectively, 3.80±0.74 μmol/mol the 20th of July, and 1.72±0.12 μmol/mol the 26th of August) (Fig. 2b, Table 1).

This ([Ba]/[Ca])shell profile demonstrates a high interindividual reproducibility among scallops of the same age for both background content and summer peak events with similar occurrence dates and amplitudes (Fig. 2a). The reproducibility of ([Ba]/[Ca])shell ratios was previously demonstrated for 2-year old scallops from the Bay of Brest (Barats et al., 2007). Analyses of a 3-year old scallop from the same population (shells collected the same year and site) were also performed for the third year of the shell growth.
3.1.2 Recurrence of \( [\text{Ba}]/[\text{Ca}] \) profiles over a 7-year period (1998–2004, Bay of Brest) and among different coastal sites (Western Europe)

The \( [\text{Ba}]/[\text{Ca}] \) profiles were then examined on a specific population (Bay of Brest) over a 7-year survey. Inter-shell comparison of \( [\text{Ba}]/[\text{Ca}] \) profiles obtained for 3 scallop replicates confirms first the significant reproducibility of these profiles among a single population \((p<0.05, n>60, \text{except in 2000 and 2001})\) \( (\text{Table 1}) \). Variations of mean \( [\text{Ba}]/[\text{Ca}] \) \( (n=3) \) over the 7-year period reveal a common pattern with a stable background ratio punctuated by 2 summer maxima with specific year-dependant intensities \( (\text{Fig. 3, Table 1}) \). The background \( [\text{Ba}]/[\text{Ca}] \) ratio is about \( 0.56\pm0.12 \mu \text{mol/mol} \) and rather homogenous along the year \((\%\text{RSD}<20)\). The first summer \( [\text{Ba}]/[\text{Ca}] \) peak occurs from mid June to July and the second one from end July to early September. The year 2000 exhibits the weakest \( [\text{Ba}]/[\text{Ca}] \) ratio \( (1.03–0.73 \mu \text{mol/mol}) \). The year 2002 remains unusual with four \( [\text{Ba}]/[\text{Ca}] \) peaks \( (1.86, 1.31, 1.55, 2.83 \mu \text{mol/mol}) \).

The background and maximum \( [\text{Ba}]/[\text{Ca}] \) ratios obtained in different ecosystems \( (\text{Bay of Seine, Bay of Brest, Belle Ile, Quiberon, Ria de Vigo}) \) were summarized in Table 2. This inter-site study shows a similar \( [\text{Ba}]/[\text{Ca}] \) profile with background content ranging from 0.44 to \( 0.68 \mu \text{mol/mol} \) and ubiquitous maxima in summer. The range of maximum \( [\text{Ba}]/[\text{Ca}] \) ratios is similar in all locations \( (0.73–5.3 \mu \text{mol/mol}) \) corresponding to a 1.3–9 fold enrichment. These results characterize a recurrent \( [\text{Ba}]/[\text{Ca}] \) profile typology at all sites and years, with a constant background content punctuated by a first intensive peak in early summer \( (\text{mid-June to July}) \) and a second weaker one in late summer \( (\text{August to early September}) \).

3.1.3 Comparability of archived Ba signals in other bivalve shells

These recurrent \( [\text{Ba}]/[\text{Ca}] \) profiles observed for \( P. \text{maximus} \) shells are compared to similar ones recorded in bivalves from other coastal environments \( (\text{Table 2}) \). The background \( [\text{Ba}]/[\text{Ca}] \) ratio and summer peaks \( (\text{max. 9 times the background in} \ P. \text{maximus} \text{shells and up to 34 times the background for other bivalves}) \) are slightly lower in \( P. \text{maximus} \) shells. Differences in \( [\text{Ba}]/[\text{Ca}] \) content can be related to the bivalve species, its physiology, or differences in analytical methods. For example, the maximum intensity of \( [\text{Ba}]/[\text{Ca}] \) ratios is particularly high in mussels and clams \( (\text{up to} 20 \mu \text{mol/mol}) \). This may be explained by the preferential precipitation of Ba under the aragonite structure of \( \text{CaCO}_3 \) \((\text{Rimstidt et al., 1998}) \). Higher \( [\text{Ba}]/[\text{Ca}] \) content can be influenced by the structure of this bivalve \( (\text{aragonite or calcite}) \). Gillikin et al. (2008) demonstrated however that shell mineralogy has no effect on \( [\text{Ba}]/[\text{Ca}] \) ratios \( (\text{Gillikin et al., 2008}) \).
Fig. 3. Variations of mean \( [\text{Ba}/\text{Ca}]_{\text{shell}} \) ratios (3 shells per year; except in 2002, \( n=2 \); black profile) according to the chlorophyll-\( a \) concentration (\( \mu g/l \), grey profile), over 7-year period (1998–2004) in scallops from Roscanvel station (Bay of Brest, France).

Differences in calibration steps can also induce differences in \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) ratios. Quantitative Ba analyses were performed with LA-ICP-MS using external calibrations with NIST glass standard reference materials at a minimum of 41 \( \mu g/g \) of Ba (Carré et al., 2006; Lazareth et al., 2003; Pearce and Mann, 2006; Stecher et al., 1996; Vander Putten et al., 1999, 2000). Such external calibration is probably not suitable to accurately measure shell Ba concentrations ranging from 1 to 10 \( \mu g/g \), especially considering the different matrix properties and response between glasses and CaCO\(_3\) shells when using some laser ablation units (Belotto and Mikeley, 2000; Barats et al., 2007). Based on these conclusions, external calibrations with CaCO\(_3\) standards was thus preferred in this study and others (Belotto and Mikeley, 2000; Thébault, 2005; Gillikin et al., 2006; Barats et al., 2007). Finally, taking into account the accurate dating of this study, and differences in analytical methods and in bivalve species, all these studies agree with a common profile of \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) ratio in bivalves; widespread over the world a background ratio punctuated by sharp summer episodic peaks. This observation thus supports specific and ubiquitous processes involved in the increase of \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) ratio, which must be directly related to changing conditions in the scallop environment.

3.2 Biogeochemical processes influencing \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) profiles

3.2.1 \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) background content as an indicator of Ba aqueous concentration

The \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) background ratios in \( P. \text{maximus} \) shells were measured in the Bay of Brest over a 7-year period (Table 1). This reveals an average ratio of 0.56\( \pm 0.12 \mu \text{mol/mol} \). \( ([\text{Ba}]/[\text{Ca}]) \) ratio in the Bay of Brest was determined in both dissolved and particulate seawater collected at 1 m above the SWI from March to September 2000 at intervals of 2 or 3 times/week, except from 24 July to the 9 September (Lorrain, 2002). \( ([\text{Ba}]/[\text{Ca}])_{\text{shell}} \) profiles

www.biogeosciences.net/6/157/2009/  
Biogeosciences, 6, 157–170, 2009
were compared with variations of Ba concentrations in seawater (Fig. 4a–b). Variations of dissolved and particulate Ba exhibit concentrations ranging, respectively, from 45 to 100 nmol/l and 0.7 to 9 nmol/l, and averaging 50 and 2 nmol/l. The background ([Ba]/[Ca]) ratio in dissolved seawater in 2000 is about 5.2±0.5 µmol/mol (Lorrain, 2002) which is slightly different than the value reported in 2003 (3.8 µmol/mol) (Gillikin et al., 2008). The ratio calculated in 2003 corresponds to only 3 measurements, which is considered less than in 2000 (n=43) and may explain such difference. The major pool of Ba in the seawater at the SWI originates from the dissolved phase. Background ([Ba]/[Ca])\_shell ratios in Mytilus edulis mussel shells were previously demonstrated to be directly related to the ([Ba]/[Ca])\_sw ratios of the water in which they grew (Gillikin et al., 2006). For the same scallop species (Pecten maximus) and in the same sampling site (Bay of Brest), the partition coefficient \(D_{Ba}\) was reported to be 0.18 in 2003 and similar to those obtained for Mytilus edulis mussels or Saxidomus giganteus clams in other coastal temperate ecosystems (Gillikin et al., 2006, 2008). In our study, the shell Ba partition coefficient (\(D_{Ba}\), evaluated in 2000 using the ([Ba]/[Ca])\_sw ratio in the dissolved seawater phase and the background ([Ba]/[Ca])\_shell ratio measured in shells, is about \(D_{Ba}=0.11±0.03\) (Barats, 2006). Taking into account the difference of analytical methods, these results are all similar to those obtained previously as reported in the literature. Background ([Ba]/[Ca])\_shell ratios in P. maximus shells may thus be considered as a relevant proxy of Ba aqueous concentrations.

### 3.2.2 Transient processes related to ([Ba]/[Ca])\_shell maximum events

**The origin of ([Ba]/[Ca])\_shell scallop shell maxima**

Summer ([Ba]/[Ca])\_shell maxima underline an important additional source of Ba at the SWI, which in turn will influence both dissolved and particulate Ba concentrations in the scallop habitat. In 2000, dissolved and particulate Ba exhibited increasing seawater concentrations at the end of July concomitant to the first increase of ([Ba]/[Ca])\_shell ratio (Fig. 4a–b) (Lorrain, 2002). This maximum of ([Ba]/[Ca])\_shell ratio is 2.6 times higher than the background one, whereas dissolved seawater Ba concentration is 1.5 times higher than the background one. The increase of dissolved Ba concentration is probably not sufficient to explain the increase in skeletal ([Ba]/[Ca])\_shell content. At the end of July, particulate

---

**Table 2.** Barium archives in different bivalve shells according to different ecosystems. Both background and maximum ([Ba]/[Ca])\_shell ratios are examined. The period of ([Ba]/[Ca])\_shell maxima are also shown.

<table>
<thead>
<tr>
<th>Bivalve</th>
<th>Species</th>
<th>Structure of precipitation</th>
<th>Location</th>
<th>Ecosystem type</th>
<th>Time resolution</th>
<th>Investigated period</th>
<th>Background Ba/Ca ratios (µmol/mol)</th>
<th>Enrichment period</th>
<th>Maximum Ba/Ca ratios (µmol/mol)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scallop</td>
<td>Pecten maximus</td>
<td>calcite</td>
<td>Bay of Brest, France</td>
<td>Coastal</td>
<td>3 days + dating</td>
<td>1996–2004</td>
<td>0.56±0.12</td>
<td>late June to early September</td>
<td>0.73–1.7</td>
<td>1.3–3</td>
</tr>
<tr>
<td>Scallop</td>
<td>Comptopallium rubrum</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>4 days + dating</td>
<td>2002–2003</td>
<td>0.52</td>
<td>July–August</td>
<td>0.92–3.7</td>
<td>2–10</td>
</tr>
<tr>
<td>Scallop</td>
<td>Argopecten purpuratus</td>
<td>calcite</td>
<td>The Netherlands, Schelde estuary</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1995–2003</td>
<td>0.7</td>
<td>July–August</td>
<td>3–12</td>
<td>4–17</td>
</tr>
<tr>
<td>Mussel</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>Kenya, Tidur estuary</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late May–June</td>
<td>1.3–9</td>
<td>2–3</td>
</tr>
<tr>
<td>Clam</td>
<td>Mercenaria mercenaria</td>
<td>anaragite</td>
<td>Delaware Bay, USA</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1984–1992</td>
<td>2.5</td>
<td>Late May–August</td>
<td>5–20</td>
<td>3–10</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late June to 0.73–4.7</td>
<td>2–3</td>
<td>1.3–9</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>Bay of Brest, France</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1996–2004</td>
<td>0.56±0.12</td>
<td>late June to early September</td>
<td>0.73–1.7</td>
<td>1.3–3</td>
</tr>
<tr>
<td>Clam</td>
<td>Mercenaria mercenaria</td>
<td>anaragite</td>
<td>Delaware Bay, USA</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1984–1992</td>
<td>2.5</td>
<td>Late May–August</td>
<td>5–20</td>
<td>3–10</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late June to 0.73–4.7</td>
<td>2–3</td>
<td>1.3–9</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late June to 0.73–4.7</td>
<td>2–3</td>
<td>1.3–9</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late June to 0.73–4.7</td>
<td>2–3</td>
<td>1.3–9</td>
</tr>
<tr>
<td>Clam</td>
<td>Mytilus edulis</td>
<td>calcite</td>
<td>New Calamudia ion, Chile</td>
<td>Coastal</td>
<td>Monthly or better</td>
<td>1991–1998</td>
<td>1</td>
<td>Late June to 0.73–4.7</td>
<td>2–3</td>
<td>1.3–9</td>
</tr>
</tbody>
</table>

* represent analyses performed with CaCO\(_3\) standards.
Ba concentration increases up to 5 times the background content. This significant increase of particulate Ba may provide additional inputs of Ba at the SWI. As bivalves are non-specific filter feeders, these additional inputs of Ba-rich particles at the SWI, in the surrounding scallop environment, are supposedly ingested as food, in part digested, transferred to the extrapallial fluid, and finally archived in the shell (Stecher et al., 1996; Vander Putten et al., 2000; Lorrain, 2002; Gillikin et al., 2006).

A proposed pelagic biogenic process as the initial cause of Ba-rich particles at the SWI and subsequent ([Ba]/[Ca])<sub>shell</sub> maxima

The occurrence and the amplitude of summer ([Ba]/[Ca])<sub>shell</sub> peaks were intensively examined in the Bay of Brest over 7 years (1998–2004) because of a regular and complete monitoring of physiological (growth) and environmental parameters (hydrological, biological, chemical).

([Ba]/[Ca])<sub>shell</sub> variation as a result of environmental changes: The influence of the shell growth rate on ([Ba]/[Ca])<sub>shell</sub> peaks is studied to take account a potential external environmental control. Decreases of shell growth rate were previously considered to be mainly induced by lower seawater temperature or the occurrence of specific phytoplankton blooms (Chauvaud et al., 1998; Lorrain et al., 2000). In 2002 (4th peak) or in 2004 (1st and 2nd peaks), such decreases of shell growth rate occur simultaneously to ([Ba]/[Ca])<sub>shell</sub> maxima. For other investigated years, ([Ba]/[Ca])<sub>shell</sub> maxima are not coincident to any shell growth anomaly. ([Ba]/[Ca])<sub>shell</sub> maxima cannot thus be directly related to minima of shell growth rate.

The specific survey in 2000 supports rather supplementary dissolved and particulate Ba inputs at the SWI in summer, subsequently taken up by the bivalve to explain ([Ba]/[Ca])<sub>shell</sub> maxima. There are different potential sources providing these Ba inputs at the SWI: either related to a benthic Ba recycling and remobilization processes, or pelagic Ba enrichments originating from anomalous ecological changes, or a coupling of both phenomena.

([Ba]/[Ca])<sub>shell</sub> variation as a result of a biogenic pelagic process: First, the examination of hydrological conditions (seawater temperature and salinity) reveals no significant similarity neither with the occurrence, nor with the amplitude of maximum ([Ba]/[Ca])<sub>shell</sub> ratios. In the Bay of Brest, the role of hydroclimatic events (flood and/or resuspension) was previously investigated in detail (Lorrain, 2002) and did not exhibit any influence on the shell growth and chemistry. Ba inputs due to such hydroclimatic events are thus improbably to explain these Ba particulate inputs at the SWI. Benthic release due to more reducing conditions at the SWI can also occur, but these should preferentially release dissolved components. More reductive condition at the SWI, as observed in a eutrophicated estuarine bay (Seine Bay), promote the benthic release of dissolved Mn which was demonstrated to induce increasing shell Mn content in summer (Barats et al., 2008). In 2000, skeletal Mn concentrations are relatively constant, concurrent with constant seawater particulate Mn content and a low increase of Mn concentration in the dissolved phase (Fig. 4c). The dissolved oxygen concentrations (annual average: 6.1±0.5 ml/l; and summer average from mid-May to mid-September: 6.0±0.3 ml/l) are constant in the Bay of Brest, suggesting well oxygenated seawater. For other years, even if seawater Mn measurements are not available, archived shell Mn supports stable content during ([Ba]/[Ca])<sub>shell</sub> maxima, contrary to benthic Ba remobilisation due to more reductive conditions. The origin of Ba inputs at the SWI is thus rather initiated by a pelagic biogenic process (Lorrain, 2002).

([Ba]/[Ca])<sub>shell</sub> variation as a result of phytoplankton biomass dynamic: The phytoplankton dynamics are
particularly examined because phytoplankton is an important vector for the vertical transport of Ba in oceans (Dehairs et al., 1980, 1991, 1997; Bishop, 1988). The Chl-α concentrations in the seawater that are supposed to reflect phytoplankton biomass, exhibited a general pattern with an intense maximum occurring in spring and later ones somewhat smaller in late summer (Fig. 3). Phytoplankton blooms generally occurred in the month preceding ([Ba]/[Ca])shell maxima. In 1998 and 2003, ([Ba]/[Ca])shell profiles revealed a surprising analogy with the variations of Chl-α concentrations: 2 maximum events with the first most intense one and during similar time periods. Time lag between the occurrence of maximum Chl-α concentration and maxima of ([Ba]/[Ca])shell ratio ranged from 1 week to 3 months (in 2003). In 1999 and 2002, ([Ba]/[Ca])shell maxima occurred coincident to periods of higher Chl-α concentration, but their profile shapes were totally different. In 2001, a particularly productive year, the maxima of ([Ba]/[Ca])shell peaks were surprisingly weak; whereas in 2004, the year with lowest productivity, the intensity of ([Ba]/[Ca])shell ratios is close to the background concentration (Fig. 5). These 2 specific blooms cannot thus influence the amplitude or the occurrence of maximum ([Ba]/[Ca])shell ratios. Neither specific species, nor these two genera ([Chaetoceros and Gymnödynium]), nor these two largest group of eukaryotic algae (diatoms or dinoflagellates), nor the total phytoplankton abundance are directly correlated with these ([Ba]/[Ca])shell maxima events.

([Ba]/[Ca])shell variation as a result of particulate biogenic material production: Among major chemical parameters examined (i.e. dissolved oxygen, nutrients, POC, PON), only POC and PON concentrations exhibit similar transient events such as maxima in ([Ba]/[Ca])shell ratio (Fig. 5). Considering maximum concentrations of POC and PON in the month preceding the maximum ([Ba]/[Ca])shell ratios, single correlations are obtained between the amplitude of maximum ([Ba]/[Ca])shell ratios and maxima of POC and PON concentrations (r²=0.58, p<0.05, n=16) (Table 3). Multiple regression analyses reveal a systematic relationship expressing the amplitude of maximum ([Ba]/[Ca])shell ratio as a function of nitrates, PON and nitrite contents.

Table 3. : Single correlations and multiple regression analyses outlining significant relationships between the amplitude of maximum [Ba/Ca]shell ratios (µmol/mol) and environmental variables (Stabox Pro software for Windows Ver. 6, Grimmer Software. France).

<table>
<thead>
<tr>
<th>Significant correlations with maxima of shell [Ba/Ca] ratio (p&lt;0.05, n=16)</th>
<th>Multiple regressions expressing the amplitude of shell [Ba/Ca] maxima (µmol/mol) according to other variables (d&lt;sub&gt;cooker&lt;/sub&gt; &lt; 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Integration periods</td>
<td>POC</td>
</tr>
<tr>
<td>−4 w+1 w</td>
<td>r²</td>
</tr>
<tr>
<td>−3 w+1 w</td>
<td>r²</td>
</tr>
<tr>
<td>−2 w+1 w</td>
<td>r²</td>
</tr>
<tr>
<td>−1 w+1 w</td>
<td>r²</td>
</tr>
</tbody>
</table>

Ajusted 0.72 Intensity of shell 0.8 0.454
0.84 Intensity of shell 0.219 0.453
0.7 Intensity of shell
Fig. 5. Example of variations of \([\text{Ba}/\text{Ca}]_\text{shell}\) ratios for 3 years (1998, 2000, 2001) in shells from the Bay of Brest (France) according to: (a) two phytoplankton genera (Chaetoceros and Gymnodymium), (b) particulate organic carbon (POC) and particulate organic nitrogen (PON) concentrations (µg/l) in seawater.

Processes involved in Ba-enrichment at the SWI that evidence subsequent \([\text{Ba}/\text{Ca}]_\text{shell}\) maxima

This study underlines the influence of an initial pelagic biological bloom process on the occurrence and the amplitude of maximum \([\text{Ba}/\text{Ca}]_\text{shell}\). These processes are non-specific to either phytoplankton species or genera, and rather related to post-bloom summer conditions. The episodic \([\text{Ba}/\text{Ca}]_\text{shell}\) maxima in *P. maximus* shells are supposed to be induced by a trophic uptake of supplementary particulate Ba inputs at the SWI. These Ba-enriched particles may originate either from scavenging of phytoplanktton-derived particles or from a benthic post-bloom remobilization. A pathway leading to particulate Ba enrichment within the water column and scavenged to the SWI is most plausible (Ganeshram et al., 2003; Sternberg et al., 2005). Different pathways were proposed to explain Ba enrichment in particles at the SWI: such as Ba adsorption in phytoplankton cells, barite formation during phytoplankton decay, Ba enrichment in exopolymeric substances (EPS), or Ba adsorption onto mineral oxides formed within diatom biogenic particles.

in the seawater \((r^2>0.70, p<0.30, n=16)\) (Table 3). The amplitude of maximum \([\text{Ba}/\text{Ca}]_\text{shell}\) ratio is higher during increasing nitrate and PON contents in seawater and during decreasing concentrations of nitrates. These statistical relationships agree with an initial pelagic biogenic process involved in supplementary Ba inputs at the SWI to explain the maximum \([\text{Ba}/\text{Ca}]_\text{shell}\) events. In the month preceding \([\text{Ba}/\text{Ca}]_\text{shell}\) maxima, diatom blooms are usually dominant. These taxa grow rapidly when nitrate is available. They have a high uptake capacity for nitrate, even at low concentrations (Carter et al., 2005). Nutrient availability governs the amplitude of phytoplankton blooms and its composition (Chauvaud et al., 2000; Le Pape et al., 1996; Ragueneau et al., 2002). However, higher dissolved inorganic nitrogen in the seawater during the month preceding \([\text{Ba}/\text{Ca}]_\text{shell}\) maxima promotes the primary productivity (Carter et al., 2005). Higher phytoplankton biomass induces then the production of higher inputs of PON in seawater. Periods of higher summer productivity and post-bloom conditions are prone to amplify the maximum \([\text{Ba}/\text{Ca}]_\text{shell}\) ratio.
Chaetoceros spp. blooms are usually dominant during the month preceding maximum ((Ba)/(Ca))_{shell}. The agglomeration of EPS is specifically reported during bacterial decomposition of these Chaetoceros spp. blooms, as promoted in response to nutritional stress. These conditions are indeed collated in this study, which systematically display depleted concentrations of nitrates and nitrites in seawater before the most significant maximum [Ba]/[Ca]_{shell} (Passow and Alldredge, 1995; Stecher and Kogut, 1999; Alldredge et al., 1995; Thornton, 2002). Both colloidal organic material and EPS are particularly enriched in Ba by five orders of magnitude (Quigley et al., 2002). Ba-rich EPS may thus support inputs of Ba-enriched particles at the SWI.

Among the different pathways such as Ba adsorption in phytoplankton cells, barite formation during phytoplankton decay, Ba enrichment in EPS, or Ba adsorption onto mineral oxides formed within diatom biogenic particles, the exact processes inducing Ba enrichment in particles at the SWI cannot be clearly identified. This study however supports the concept that a pelagic biogenic process initiates the delivery and subsequent bivalve uptake of Ba-enriched particles leading to incorporation into the individual striae of scallop shells.

4 Conclusions

Barium shell profiles from Great Scallops obtained in this methodical survey in temperate waters agree with those of other bivalve species previously investigated and demonstrate their reproducibility, recurrence and ubiquity in such coastal environments. This study confirms that ([Ba]/[Ca])_{shell} profiles are characterized by a relatively constant background mainly governed by the ambient seawater dissolved Ba, and the occurrence of distinct summer ([Ba]/[Ca])_{shell} maxima. In 2000, Ba measurements in seawater allowed identifying that particulate Ba inputs can be the dominant pathway explaining ([Ba]/[Ca])_{shell} maxima. Pelagic biogenic processes are supposed to initiate seawater Ba enrichment at the SWI, subsequently taken up by scallops and translated by increased ([Ba]/[Ca])_{shell} ratios. Examination of the complete dataset (1998–2004) demonstrates that ([Ba]/[Ca])_{shell} maxima occurred under summer post-bloom conditions, such that their amplitude depends on nitrogen cycle: both particulate organic nitrogen and the turnover of dissolved nitrogen species. This whole dataset demonstrates that: (1) records of maximum ([Ba]/[Ca])_{shell} ratio cannot be used directly as a relevant paleo productive tracer, and (2) complex processes occur in the pelagic/benthic Ba cycle and are responsible for significant Ba inputs at the SWI. If these processes are better constrained, scallop ([Ba]/[Ca])_{shell} records could provide a proxy of Ba biogeochemistry with high temporal resolution in coastal environments.

Acknowledgements. This work was supported in the framework of two research programs: ACI PECTEN (French Ministère de la Recherche) and “Suivi Erika” (INERIS, French Ministère de l’Environnement). Thermo Electron Company is thanked for the loan of the ICP-MS. A. B. acknowledges the Aquitaine Region (ORQUE project) for her Doctoral fellowship. A. B. thanks G. Sinquin from the Groupe de Microscope Electrone (Université de Bretagne Occidentale, France) for his help concerning the SEM (scanning electron microscopy) photography, and D. Cardinal from Department of Geology (Royal Museum for Central Africa, Belgium) for his help in analyses of particulate Ba in seawater.

Edited by: S. Bouillon

References


Eptlé, V. M.: High-resolution climate reconstruction for the Holocene based on growth chronologies of the bivalve Arctica islandica from the North Sea, Department of Geosciences, University of Bremen, Bremen, Germany, 101 pp., 2004.


McCulloch, M., Fallon, S., Wyndham, T., Hendy, E., Lough, J., and


