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Trace element heterogeneity along isochronous growth layers in bivalve shell:

Consequences for environmental reconstruction

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ABSTRACT

To precisely reconstruct environmental changes from high-resolution bivalve shell geochemistry, the shell layers must represent environmental conditions at the time of deposition. Therefore, calcium carbonate along growth lines that formed at the same time should show the same geochemical signature. We test this assumption by evaluating the geochemical spatial homogeneity along growth layers (i.e., shell secreted at a given time) on shell cross-sections of modern shells of the bivalve *Protothaca thaca*, for ontogenically young and old specimens. Analyses were carried out using laser ablation ICP-MS and solution nebulization ICP-MS. Scanning electron microscope images showed signs of early diagenesis of old shell sections (close to the umbo) compared with recent (close to the ventral margin). The spatial distribution of Sr/Ca, Mg/Ca and Ba/Ca along growth layers differed consistently between old and recent shell sections. Elemental distribution at the ventral margin of young specimens was similar for the two specimens analyzed and similar to the element distribution in the older specimens at a similar ontogenic stage (closer to the umbo). If a balancing of the shell composition with the surrounding seawater between the old and recent part of the shell, supported by diagenesis, cannot be excluded, we assume that trace element incorporation process at a given time varied with the age of the specimen, in connection with changes of mantle characteristics. Whereas Ba/Ca and Sr/Ca were almost constant for shell secreted at a given time close to the current ventral margin, all individuals showed significant Mg/Ca heterogeneities, with reproducible patterns. We assumed that this would be due to a zonation of the outer epithelium of the mantle that would favor or inhibit the transport of Mg into the extrapallial fluid, and subsequently into the shell. Time-series obtained crossing successive growth layers using laser ablation ICP-MS and solution nebulization ICP-MS on a same-shell growth section were similar for Sr/Ca and Ba/Ca ratios and not for Mg/Ca, which reflected
both the different spatial resolution of the analytical techniques and the heterogeneous
distribution of Mg/Ca. In the *P. thaca* shells, Sr and Ba seemed to be most promising as
potential high-resolution environmental proxies. Our study showed that the location of high-
resolution analyses points in shell cross-section must be i/ precisely defined and ii/
consistently followed along the shell section to provide accurate environmental records; i.e.,
without bias related to shell layer heterogeneities.

Highlights:

1/ Evaluation of geochemical homogeneity of isochronous growth layer on modern bivalves.
2/ SEM observation of shell dissolution on shell portion close to the umbo.
3/ Differences in trace element distribution along isochronous layer for old and recent shell section.
4/ Trace element incorporation process may vary with the age of the specimen.
5/ Mantle zonation is assumed to explain Mg/Ca heterogeneity along isochronous layers of recent shell.
6/ High-spatial analytical technique procedure for molluscan shell study must be precisely defined.

Keywords: bivalve, shell, laser ablation, ICP-MS, trace elements
1. Introduction

Variations of trace element content through time (i.e., crossing successive growth layers) in bivalve shells have been used to provide records of environmental changes (e.g. Freitas et al., 2005; Hendry et al., 2001; Price and Pearce, 1997; Takesue and van Geen, 2004; Vander Putten et al., 2000). Nevertheless, various parameters influence the trace element incorporation into bivalve shells (Ford et al., 2010; Freitas et al., 2009; Gillikin et al., 2005; Takesue et al., 2008). To better understand what is recorded in the shell, studies on the trace element distribution within a given growth layer of the shell, i.e., shell material deposited at the same given time, have been undertaken (Carré et al., 2006; Foster et al., 2008; Freitas et al., 2009; Klein et al., 1996; Rosenberg et al., 1989). Indeed, shells will be reliable tracers of the environment only when their chemical composition at a given time point is homogeneous and reflects the environment.

Previous studies of intra-shell variability have focused essentially on different shell growth axes, each characterized by a different shell growth rate. Indeed, shell growth rate is lower for an axis going from the umbo to the posterior edge than for an axis going from the umbo to the ventral margin (see Fig. 2 from Klein et al., 1996, for example). Differences in Mg/Ca and Sr/Ca ratios between *Mytilus edulis* shell portions characterized by low or high growth rate were interpreted in terms of differences in mantle metabolism (Klein et al., 1996; Rosenberg and Hughes, 1991; Rosenberg et al., 1989). Recently, Carré et al. (2006) observed different Sr/Ca profiles in *Mesodesma donacium* shells between two different sections of the shell. These authors proposed a Sr incorporation model that took into account the shell "mineralization rate." Few works, however, concern the spatial distribution of the elements within the shell section made along the maximum growth axis, i.e., where the geochemical profiles for the environmental reconstitutions are carried out. In one bivalve shell section cut...
along the maximum growth axis, Carré et al. (2006) observed trace element heterogeneity within single growth layers, i.e., following an isochronous profile (Carré et al., 2006; electronic annex 2). Nevertheless, few analyses per growth layer (two in most cases) were carried out, yielding limited insight into the possible heterogeneity of trace element concentrations within the growth layer considered. Within one section of Arctica islandica, Foster et al. (2008; 2009) realized three parallel profiles at different distances from the periostracum and observed a Sr/Ca and Mg/Ca decrease while moving away from the periostracum. The authors proposed that these differences are related to crystal growth mechanisms or extrapallial fluid (EPF) heterogeneity. In Mytilus edulis and Pecten maximus shells, Freitas et al. (2009) evidenced Mg/Ca, Sr/Ca, and Mn/Ca heterogeneity along growth layers. To explain these heterogeneities, the authors evoked small differences in the mechanisms of element incorporation at the interface between the crystals and the EPF, or the presence of different amounts of organic matrix. These studies thus show that for a given growth layer of a given shell section, the distribution of trace elements can be heterogeneous. This means that, depending on the location where high-resolution trace element time-series analyses are carried out, even within the same section the results obtained can be different without this being due to environmental variability.

In this study, we analyzed the trace element concentration distribution along isochronous growth layers in several Protothaca thaca bivalve shells using laser ablation ICP-MS (LA-ICP-MS). For three shells, the trace element spatial distribution in two growth layers, close to and far from the ventral margin, was analyzed to determine whether the trace element distribution along growth layer remains the same during the lifetime of the animal. Then, to determine if the changes observed were related to modification of the shell (i.e., the biocarbonate as material) or to the animal ageing, shells of young and old specimens were studied. Lastly, short parallel time-series profiles over the same period of shell growth were
obtained on two shells, using LA-ICP-MS for one and LA-ICP-MS and SN-ICP-MS for the other, to examine these two techniques in connection with their application on molluscan shells in terms of paleoenvironmental reconstruction.

2. Material and methods

Table 1 presents a summary of the samples studied and of the geochemical analyses carried out on them.

2.1 Samples

Protothaca thaca (Molina, 1782) (Mollusca, Veneridae) is an upper-subtidal bivalve commonly living along the southeastern coasts of the Pacific Ocean (8°-45°S). Adult *P. thaca* shells were collected from two locations, Pocoma in the south of Peru (17° 25’ S, 71° 22’ W) and El Lenguado (23° 46’ S, 70° 28’ W), south of Antofagasta, Chile. The Pocoma specimens (PC77, PC134) were collected alive in July and in November 2003 respectively. These specimens belonged to a population used for a growth experiment in which a sclerochronology study was performed (Lazareth et al., 2006). Two specimens of around one year old, from the same Pocoma population, were also selected (PC173, PC180). PC173 and PC180 died between August and September 2003. Shells of recently dead specimens (C00-423-3, C00-423-5) were collected on a beach near Antofagasta, in 2000. The *P. thaca* shell is composed of two aragonite layers, an outer prismatic and an inner homogeneous layer. Hereafter, the shell part closer to the seawater will be termed the "external side of the shell" whereas the shell part in contact with, or closer to, the animal body will be called the "internal side of the shell." Shell sections, or shell growth layer, far from the ventral margin (therefore closer to the umbo, indicating that it was deposited when the animal was young) will be
termed "old." Contrarily, shell parts, or shell growth layer, close to the ventral margin (i.e., deposited shortly before death) will be termed "recent."

Shells were sectioned along their axis of maximum growth using a water-cooled diamond saw (Isomet 5000®, Buehler). The half-valve cross-sections were then polished, first using diamond grinding discs from 70 to 30 µm grain size, and then using diamond-suspensions with decreasing grain sizes (15 to 1 µm) on polishing cloths. Geochemical analyses were done in the prismatic layer. For the LA-ICP-MS analyses, 0.3 cm-thick slabs were cut from the half-valve polished cross-sections. LA-ICP-MS measurements on the C00-423-3 shell were compared to conventional digestion and solution nebulization ICP-MS (SN-ICP-MS) results obtained on an adjacent section of the same shell, for the same growth period. To collect samples for SN-ICP-MS, a shell section of C00-423-3 was embedded in epoxy resin and a slice ~ 5 mm thick was cut and polished before sampling using a micro-drilling device. A scanning electronic microscope (SEM) was used to examine the microstructure of the prismatic layer. Manually broken pieces of the PC77, PC134, and C00-423-3 specimens, one close to the ventral margin and one close to the umbo, were examined using a Cambridge Stereoscan SEM at 15 kV. In addition, for PC134 and C00-423-5, the microstructure of the prismatic layer was also examined at the location of the older isochronous profile (i.e., at around 1 cm from the ventral margin). All samples were sputter-coated with carbon before examination.

2.2 Laser ablation analyses

The analyses were conducted on the shell cross-sections using two sampling strategies (Fig. 1): along single growth layers, i.e., sampling shell secreted at a given time (hereafter-called "isochronous profiles," Fig. 1b); and sampling successive shell growth layers (hereafter called "time-series profiles," Fig. 1c). The reproducibility of the time-series was examined on
the C00-423-5 specimen doing two parallel time-series profiles on the same portion of the shell. These LA-ICP-MS profiles are located at a mean distance of 350 µm (profile 1) and 600 µm (profile 2) from the external side of the shell.

Isochronous profiles were made on all specimens. To analyze shell material deposited at the same time on the shell cross-sections, we chose one darker growth line like a guide. The laser spots for the isochronous profiles were carefully placed at small distances along this noticeable growth line (i.e., the darker line was not analyzed). For PC77, PC134, and C00-423-3, we made two isochronous profiles, one close to the ventral margin (~0.5 cm), and slightly distant (~1 cm; PC134) or far (>2 cm; C00-423-3, PC77) from the ventral margin (see Fig. 1a and Table 1). One isochronous profile at around 1 cm from the ventral margin was done in C00-423-5. At last, two isochronous profiles were performed in each young Pocoma specimen, at 60 µm and 260 µm from the ventral margin, i.e., separated from each other by ca. one week of shell growth. To clarify, isochronous profile close to the current ventral margin corresponds to recent shell whereas the one far from the current ventral margin corresponds to old shell. The thickness of the outer shell layer increases towards the ventral margin of the shell and differs between specimens. Consequently, to compare the isochronous profiles of the different shells, the same number was allocated to the samples located at the same relative position with respect to the inflection point of the increment considered. The n°3 was arbitrarily allocated to the laser spot the closest to the inflection point (see example in Fig. 2c). Then, numbers decreased for spots closer to the external side of the shell and increased for spots closer to the internal side of the shell. Thereby, the same number on each isochronous profile should correspond to a similar location of the mantle, the organ of the animal secreting the shell. To compare the trace element variations along isochronous profiles of the four specimens, the geochemical data were centered (i.e., mean subtracted).
The laser was a frequency quadrupled Nd-YAG operating at 266 nm (LSX 200 Cetac®) coupled with an Elan 6000 (Perkin-Elmer—SCIEX) ICP-MS from the "Laboratoire des Mécanismes de Transfert en Géologie, LMTG" (now "Géosciences Environnement Toulouse, GTE", Toulouse, France). The carrier gas was helium, mixed with argon before the plasma torch. Each laser ablation analysis was done as a single spot, and lasted between 50 and 60 s. With the LA-ICP-MS operating conditions used (Table 2); the resulting laser holes were 62 µm in diameter. The washout time between spots was at least 30 s. Ion sensitivity, oxide rates, and doubly charged ions were optimized each day by adjusting the plasma power, gas flows, and lens settings during ablation of the NIST-610 standard. The signal intensities of $^{24}$Mg, $^{43}$Ca, $^{88}$Sr and $^{138}$Ba were recorded. To correct for daily instrumental drift, the NIST-610 was analyzed every 10 analyses. A pressed pellet of the certified otolith CRM-22 (National Institute for Environmental Studies, Japan) was analyzed every 10 to 20 samples to estimate the accuracy. The production of the CRM-22 pressed-pellet is described in Lazareth et al. (2007).

The LA-ICP-MS data processing was done using the GLITTER© software (van Achterbergh et al., 2001) available at the LMTG laboratory (now "Géosciences Environnement Toulouse, GTE"). The first ~ 5 seconds were integrated and subtracted from the analytical signal (background). For signal integration, the largest window, corresponding to the more stable signal, was chosen. Usually, this corresponded to the signal between 20 and 40 s. $^{43}$Ca was used as internal standard with a Ca shell content of 40 wt%. The NIST-610 was used as external standard, using NIST-610 element concentrations from Pearce et al. (1997) to calculate the quantitative element concentrations of the unknown samples, CRM-22, and shells.

In a first data processing step, all analyses carried out on NIST-610 during one day were considered to calibrate the unknown samples. The NIST-610 concentrations calculated
by the GLITTER© program were then checked for instrumental drift for each daily session. If a drift of more than 5% was observed, analyses for the considered day were split in several parts and reprocessed. After drift corrections, accuracy was checked using the CRM-22 data. Reproducibility and accuracy are in percentages, calculated on trace element concentrations in part per million. Then, element/Ca ratios were calculated assuming a constant shell Ca content of 40%.

The reproducibility of LA-ICP-MS was evaluated from the mean relative standard deviation (RSD, in %) of all certified reference material results. The shell Sr and Ba contents are close to those in CRM-22, whereas the shell Mg content is close to that of NIST-610. Consequently, the RSDs of Sr and Ba obtained on CRM-22 (n = 176) and the RSDs of Mg obtained on NIST-610 (n = 259) were taken as representative of shell trace element measurement reproducibility. The reproducibility was 3% for Mg, 3.6% for Sr, and 7.3% for Ba. The LA-ICP-MS accuracy was determined from CRM-22 results. Although shell reference material would have been desirable, none was available when this work started. For each analytical sequence, we calculated the difference between the measured and the reference values of CRM-22. The mean difference, expressed as a percentage, was then calculated. The mean accuracy on CRM-22 considering all sequences (n = 176) was 5.6% for Mg, 10.8% for Sr, and 8.7% for Ba.

2.3 Solution analysis

A part of the C00-423-3 shell, already analyzed using LA-ICP-MS, was analyzed using SN-ICP-MS. The solution analyses were carried out on an Ultramass Varian ® ICP-MS instrument at the IPSL/LOCEAN laboratory, Centre IRD France Nord (ex-UR055-Paléotropique). The operating conditions are listed in Table 3.
Powders were taken following growth increments (curved grooves), sampling almost all the external shell layer, using a Merchantek Micromill® device. The average width of a micro-drilled sample was 200 ± 6 µm and sampling was as continuous as possible along the shell section. The carbonate powders were dissolved in 2.5 to 5 ml of 2 wt% nitric acid solution (Suprapur 65 wt%, Merck) that contained 100 µg.l⁻¹ of scandium, yttrium, and indium, used as internal standards. For ICP-MS calibration, standard solutions with different increasing concentrations of the analytes were prepared by appropriate dilution of mono-elemental Mg, Ca, Sr, and Ba 1 000 mg.l⁻¹ certified stock solutions (Spex, Longjumeau, France) in 2 wt% nitric acid. One sample of the above standard solution was analyzed every 10 samples to check for instrumental drift, which was less than 5%. The CRM-22 was analyzed every 10 samples for accuracy assessment.

Sample matrix effect and long term fluctuations were corrected using the internal standard method (Le Cornec and Corrège, 1997). Scandium was used as an internal standard for Ca and Mg, yttrium for Sr, and indium for Ba. As the powders were not weighted, results are given as element/Ca ratios.

The Mg/Ca SN-ICP-MS ratios for micro-milled samples taken between 22 to 30 mm from the shell ventral margin were significantly higher than the Mg/Ca results obtained using LA-ICP-MS in the same shell section. Analyses on additional samples milled from the same shell section confirmed that the first samples had probably been contaminated (in the tubes?). Consequently, the Mg/Ca data between 22 to 30 mm were removed.

The SN-ICP-MS reproducibility was calculated from shell data. Fourteen C00-423-3 samples were analyzed two or three times. The reproducibility (n = 14) was 2.3% for Mg/Ca, 1.4% for Sr/Ca, and 2.3% for Ba/Ca. For Mg/Ca, the reproducibility obtained here is similar to that reported by Gillikin (2005). The reproducibility for Sr/Ca was two times better than the one reported by Gillikin et al. (2005) for an in-house standard analyzed by SN-HR-ICP-MS.
Lastly, our Ba/Ca reproducibility was similar to that obtained on *Saxidomus giganteus*, and more than twice that obtained on *Mytilus edulis* using a SN-HR-ICP-MS (Gillikin, 2005; Gillikin et al., 2006). Our results can thus be considered as highly reproducible. Accuracy was checked on CRM-22 (n=5) and was 2.9% on Mg/Ca, 0.9% on Sr/Ca, and 11% on Ba/Ca. For Ba, the SN-ICP-MS accuracy is poor (11%). This probably reflects that, compared to Sr and Mg, Ba concentration in CRM-22 is very low (less than 0.5 µg.l⁻¹ in solution).

### 2.4 Comparison of LA-ICP-MS and SN-ICP-MS profiles

To compare the results obtained in C00-423-3 using LA-ICP-MS and SN-ICP-MS, sample correspondence between each shell slice analyzed must be determined. The distance from the ventral margin of the shell for each sample was measured under an optical microscope. For these measurements, we took the center point of the micro-drill grooves or laser craters. Then, for each shell slice, dark growth bands were identified. Samples taken from these easily distinguishable features were used as references to correlate the LA-ICP-MS and SN-ICP-MS profiles.

Reduced major axis regressions, using the software from Bohonak and van der Linde (2004), were carried out to compare the analytical results obtained with LA-ICP-MS and SN-ICP-MS.

### 3. Results

#### 3.1 Isochronous trace element profiles

The patterns of the isochronous trace element profiles close to the ventral margin of the adult specimen’s shells (3 profiles; corresponding to recent shell) differ according to the element considered (Fig. 2a). The Mg/Ca patterns are similar with first a decrease, then an increase,
and subsequently another decrease moving from the external towards the internal side of the shell. These variations are significant considering the analytical precision of 3% for Mg/Ca. The Sr/Ca decreases slightly along the analyzed growth lines. Along the isochronous profiles in the PC77 and PC134 shells, the Ba/Ca decreases slightly, whereas in C00-423-3 a sharp decrease is observed towards the internal side of the shell (Fig. 2a). Trace element distributions for the isochronous profiles far from the ventral margin of the shells (corresponding to older shell) differ from those observed for the isochronous profiles close to the ventral margin. Almost all trace element ratios decrease toward the internal side of the shell (Fig. 2b). These decreases range from 33% to 70% for Mg and Ba, and from 16% to 37% for Sr. Only the Sr/Ca and Ba/Ca patterns in C00-423-3 are different, with, from the external to the internal side of the shell, an increase for Sr/Ca and a decrease followed by an increase for Ba/Ca. In addition, the ranges of elemental ratios are higher for isochronous profiles far from the ventral margin (e.g., Mg/Ca from ~ -1.0 to 1.0 around the mean) compared with ranges for isochronous profiles close to the ventral margin (e.g., Mg/Ca from ~ -0.2 to +0.2 around the mean).

The trace element distribution along the four isochronous profiles done close to the ventral margin of the young Pocoma shells (PC173 and PC180) is similar for both shells and for all elements, with a decrease from the external to the internal side of the shells (Fig. 3a). The range of trace element/Ca ratio is higher for isochronous profile very close to the ventral margin of the shell (60 µm away), with higher ratios close to the external side of the shells, than that observed for the isochronous profiles at 260 µm from the ventral margin. Compared with isochronous profiles done close to the ventral margin in the older specimens, the range of trace element/Ca ratio is far higher in young shells (Fig. 3a). This is particularly striking for the Mg/Ca ratios which are comprised between -0.1 and 0.1 around the mean for the old specimens and between -0.5 and +0.6 around the mean for the young specimens. Contrarily,
the distribution and range of trace element ratios along the isochronous profiles done close to the ventral margin of the young Pocoma shells are similar to those done far from the ventral margin in the older specimens (Fig. 3b).

3.2 Time-series profiles

Trace element time-series profiles were obtained on two adjacent shell slices of C00-423-3 using LA-ICP-MS and SN-ICP-MS (Fig. 4). The Sr/Ca and Ba/Ca profiles obtained with the two methods correlate well ($R^2$ of 0.69 for Ba/Ca and 0.70 for Sr/Ca; $p<0.001$) whereas the Mg/Ca differences between LA-ICP-MS and SN-ICP-MS are more pronounced (Fig. 4). Even if the Mg/Ca values and variations are similar in the two profiles, the two time-series are not correlated ($R^2 = 0.15; p<0.001$).

Two time-series profiles for the same growth period, but at different distances from the external side of the shell, were carried out on C00-423-5 using LA-ICP-MS. The two profiles obtained are highly similar (Fig. 5a). For Sr/Ca and Ba/Ca, the determination coefficients are good ($R^2 = 0.61$ for Sr; $p<0.001$) or very good ($R^2 = 0.95$ for Ba; $p<0.001$) and points on the scatter diagrams are close to the 1:1 line (Fig. 5b). For Mg/Ca, the correlation coefficient between the two LA-ICP-MS profiles is 0.72 ($p<0.001$) and the regression line is slightly shifted from 1:1 (Fig. 5b).

3.4 "Recent" vs. "old" prismatic shell microstructure

The microstructure of the prismatic shell layer was examined close to the ventral margin (i.e., recent shell) and far from the ventral margin (i.e., close to the umbo = old shell). The surface of the prisms close to the ventral margin was relatively smooth, constituted by micrometer-sized scales (Fig. 6a & c). Older prisms showed an altered surface; the scale edges seemed to have been slightly dissolved and the surface of the prisms was granular (Fig. 6b & d). All
specimens examined presented the same patterns. The microstructure of the PC134 and C00-423-5 shells at around 1 cm from the ventral margin was well preserved.

4. Discussion

4.1 Intra-shell trace element distribution along isochronous growth layers

Few studies have examined the element distribution along individual growth layers in shell cross-sections (isochronous profiles; see Fig. 1a & c), and even fewer have examined this for several specimens of the same species. In our *Protothaca thaca* shells, with the exception of C00-423-3 for Sr/Ca and Ba/Ca, the isochronous trace element patterns are similar for individuals taken from the same site (PC77 and PC134, Pocoma) and for individuals taken from two different sites (~700 km apart) on various dates (C00-423-3 and 423-5, Antofagasta-2000 vs. PC77 and PC134, Pocoma-2003). Consequently, the way the elements are distributed in the growing shell at a given time is independent of the environment; the same kind of distribution is observed for two different sites and moments. The particular element distribution observed along isochronous profiles is thus more probably specific to the bivalve species, here *Protothaca thaca*. A species dependency on trace element incorporation into bivalve shells was also proposed by Freitas et al. (2009) following their observation of differences in small-scale heterogeneity between *Pecten maximus* and *Mytilus edulis*.

On the other hand, patterns are different depending on the location of the isochronous profile within the same shell, in particular for Mg/Ca and Sr/Ca (Fig. 2; Table 1). We observed shell microstructure alteration (early diagenesis) with slight dissolution of the prisms for the old shell part (far from growing ventral margin), even on specimens collected alive (PC77 and PC134). This shell alteration might have favored some balancing of the shell composition with the surrounding seawater, the external part of the old shell becoming
relatively enriched in trace elements. Pyritization was observed at the surface and inside living shells collected in marshes (Clark and Lutz, 1980). Inside the shells, the pyrite was located in specific places where the shells were weak (e.g., holes, fractures, zones rich in organic matter). To our knowledge, however, our study is the first to mention shell dissolution on modern seawater bivalve shells, all the more collected alive, and to show its potential impact on trace element distributions. The occurrence of such early diagenesis, and of its consequences in terms of geochemical signature of the shell, should be evaluated on other molluscan species, in particular on long-lived ones.

Nevertheless, we also observed similar trace element distribution for isochronous profiles far from the ventral margin in old altered shell (PC77 and C00-423-3) and closer to the ventral margin in more recent shell with no clear microstructure alteration (PC134; C00-423-5). Therefore, early diagenesis alone is insufficient to explain the different trace element distribution along isochronous profiles done close to and far from the ventral margin. The isochronous trace element distributions and ranges at the ventral margin of the young specimens (PC173 and PC180) differ from those observed close to the ventral margin of the older specimens (Fig. 3a, Table 1), but are similar to the isochronous profiles done far from the ventral margin of the older specimens (Fig. 3). We hypothesize that the way the elements are incorporated along the growth layer (isochronous profiles) differs according to the age of the individual. Indeed, Jolly et al. (2004) showed that the outer epithelium of the mantle (responsible for shell formation) of *Haliotis tuberculata* has different characteristics according to the age of the animal (tubular zone more or less developed). Thus it could be that the differences observed between the isochronous profiles in the same shell (close to and far from the ventral margin) are related to an evolution of the mantle tissue during the life of the animal rather than to diagenesis. At last, one cannot exclude a redistribution of the trace elements along the isochronous layer shortly after its deposition (in the course of around one
week) as observed on the two isochronous profiles done close to each other in each young shell (PC173, PC180; Fig. 3a). We propose that this could be related to a maturation effect as observed in teeth enamel (Balasse, 2002 and references therein; Zazzo et al., 2005) but further studies are needed to test this hypothesis.

For isochronous profiles done close to the ventral margin (i.e., recent shell), Sr/Ca and Ba/Ca ratios are almost constant whereas Mg/Ca ratios vary significantly (Fig. 2a). The chemical components used to construct molluscan shell are transported into the EPF via the outer mantle epithelium (e.g., Simkiss and Wilbur, 1989; Wilbur, 1972; Wilbur and Saleuddin, 1983). The mantle and its outer epithelium are divided into zones (e.g., central and marginal or central, pallial, and marginal) with their own characteristics in terms of histology and matrix protein types (Auffret et al., 2003; Bubel, 1973; Crenshaw, 1980; Fang et al., 2008; Jolly et al., 2004; Neff, 1972; Wheeler, 1992; Wilbur, 1972). We hypothesize that some zonation in the outer epithelium may favor or limit the transport of Mg into the EPF, and subsequently into the forming shell (Fig. 7). Knowing that the distribution of Mg/Ca along the isochronous profiles is similar between our Protothaca thaca shells, the places where the magnesium content is higher could correspond to specific zones, dependant on the species and on the specimen age, of the outer epithelium of the mantle. To explain heterogeneities along growth layers in Mytilus edulis and Pecten maximus shells, Freitas et al. (2009) put forth the assumption of the existence of small differences in the mechanisms of element incorporation at the interface between crystals and EPF, or of various quantities of organic matrix. Our assumption of a zonation of the outer epithelium of the mantle would confirm the first hypotheses of Freitas et al. (2009), and even specify it via a physiological explanation.

Moreover, our assumption would be in favor of a very limited distance between the mantle outer epithelium cells and the forming shell, as proposed in other studies (Addadi et al., 2006; Freitas et al., 2009; Simkiss and Wilbur, 1989). Nevertheless, ultrastructural studies of the
outer epithelium of the *P. thaca* mantle, and more specifically of the zone responsible for the prismatic shell layer deposition, are needed to validate our zonation hypothesis.

4.2 Shell chemical sampling procedures and their consequences in terms of environmental reconstructions

To reconstruct environmental changes that occurred during the life of the animal, successive growth layers of the shell are analyzed using relatively low-spatial analysis techniques (e.g., micromill sampling followed by SN-ICP-MS) or high-spatial resolution techniques (e.g., microprobes, LA-ICP-MS). In this study, we obtained times-series profiles on the same shell, for the same period of growth, using LA-ICP-MS and SN-ICP-MS. The resolution of the laser and that of the solution method are different. Indeed, because of the sampling procedure, a larger shell area is analyzed using SN-ICP-MS than when using LA-ICP-MS. This applies in terms of the time interval covered, due to sampling size (microdrill 200 µm vs. crater 60 µm), and in terms of the part of the outer shell layer sampled (almost all the outer layer is sampled using the micromill). Consequently, SN-ICP-MS analysis is less sensitive to potential heterogeneity within growth layers (= isochronous layers), contrary to the LA-ICP-MS. For Sr/Ca and Ba/Ca, the results obtained on *Protothaca thaca* using SN-ICP-MS and LA-ICP-MS correlate well. Similar results were obtained by Gillikin (2005) on *Saxidomus giganteus*. As in the Gillikin (2005) study, the elevated Ba/Ca values that occur on small shell portions (corresponding to 3 to 4 laser points) are averaged when using SN-ICP-MS (due to the sampling procedure as mentioned above). Nevertheless, the good reproducibility observed between the two analytical methods for Sr/Ca and Ba/Ca in this study also reflects the good homogeneity of these element/Ca ratios along the isochronous profiles. Contrarily, for Mg/Ca, the correlation between the two analytical methods is poor. Gillikin (2005) made a similar observation and concluded that the certified material used for LA-ICP-MS calibration was
inadequate (glass instead of a shell). Even if a similar conclusion can be drawn here (the NIST-610 was used for trace element LA-ICP-MS data calibration), an additional explanation can be put forward. Indeed, the Mg/Ca fluctuations observed along the isochronous growth layers in *P. thaca*, captured by the LA-ICP-MS but not by the SN-ICP-MS method, certainly contribute to the poor correlation obtained between the two methods. Two conclusions arise from these results, i/ the LA-ICP-MS procedure provides accurate data and ii/ similar Sr/Ca and Ba/Ca time-series are obtained using relatively low-spatial resolution or high-spatial resolution analytical techniques, whereas Mg/Ca shell growth layer heterogeneities complicate high-spatial resolution sampling procedure of the shell.

Although SN-ICP-MS analyses provide accurate results, this procedure is time consuming. Moreover, when sampling shells using micromill, the spatial (and thus the temporal) resolution is reduced, which can be limiting, in particular to obtain paleo-environmental records with a high temporal resolution from slow-growing mollusks. We thus further evaluated the LA-ICP-MS sampling procedure on two time-series profiles realized on the same growth period but at different distances from the external side of the C00-423-5 shell. The Sr/Ca and Ba/Ca shell changes through time are reproducible. Consequently, environmental or paleo-environmental reconstructions based on high-spatial resolution Sr/Ca and Ba/Ca time-series profiles on *Protothaca thaca* shells will not suffer from bias related to the location of the analyzes with respect to the external side of the shell. These elements could thus be used as potential proxies for environmental changes (e.g., salinity, productivity), but only if an accurate relationship between environmental parameters can be established. Indeed, if the Ba/Ca potential as environmental tracer in bivalve shells has been shown in some species (Barats et al., 2009; Gillikin et al., 2006; Gillikin et al., 2008; Thébault et al., 2009), the use of Sr/Ca as tracer of environmental changes seems compromised (Foster et al., 2009; Gillikin et al., 2005; Takesue and van Geen, 2004).
For Mg/Ca, the deviation from 1:1 of the slope of the regression line between the two LA-ICP-MS time-series (made at a different distance from the external side of the shell) clearly reflects the Mg/Ca isochronous heterogeneity. The validity of environmental data extracted from high-spatial resolution Mg/Ca analyses, at least in *Protolithaca thaca*, is thus highly dependent on the careful location of the high-resolution analyses. One must locate the analysis points at the same relative position with respect to the thickness of the shell layer considered; e.g., always with 1/3 the thickness relative to the external part of the shell. Still, this procedure is appropriate only for shell parts where trace element distribution along a single growth layer remains the same. Concerning Mg/Ca, the complexity of its incorporation in the shell at a given moment shown in this study might provide additional clues to explain the difficulties encountered for the calibration of a SST proxy based on Mg/Ca in molluscan shells (e.g., Surge and Lohmann, 2008; Wanamaker Jr et al., 2008). Before interpreting shell geochemistry in terms of environmental changes, shell growth layer homogeneity, on the shell cross-section chosen, should be assessed for recent (close to the ventral margin) and old (closer to the umbo) shell section, and this more particularly concerning Mg/Ca records.

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growth experiment, and L. Ortlieb for providing the Antofagasta shells.
References


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Mytilus edulis (bivalvia) shell calcite precipitated at constant temperature.

Biogeosciences 6, 1209-1227.


Figure captions

**Figure 1.** Illustration of the procedures followed for the high-spatial resolution intra-shell geochemistry study carried out in the *Protothaca thaca* specimens. (A) Three-dimensional diagram of a shell cross-section showing the isochronous (open circles) and the time-series (filled circle) laser-ablation ICP-MS profiles (not to scale). The numbers correspond to the isochronous profiles realized close to (1) and far from (2) the ventral margin, corresponding to recent and old shell respectively. Ext.: external side of the shell; Int.: internal side of the shell. (B) Example of a laser-ablation ICP-MS isochronous profile. Scale bar: 200 µm (optical microscope, diffuse light). (C) Example of a laser-ablation ICP-MS times-series profile (optical microscope, diffuse light).

**Figure 2.** Trace element distributions (LA-ICP-MS) along isochronous profiles done in the different *Protothaca thaca* shells (centered element/Ca ratios). "Ext." and "Int." : external and internal side of the outer shell layer. (A) Isochronous profiles close to the ventral margin (i.e., recent shell) (B) Isochronous profiles far from the ventral margin (i.e., old shell). See Table 1 for details. (C) Close up of C00-423-3 (left) and PC134 (right) isochronous profiles (thick slide, optical microscope, diffused light). Black numbers correspond to actual laser craters; white numbers are fictive laser craters that illustrate the choice of numbering to compare the different isochronous profiles (see text for details).

**Figure 3:** Comparison of the trace element distributions (LA-ICP-MS) along isochronous profiles between the young and the old *Protothaca thaca* specimens (centered element/Ca ratios). "Ext." and "Int." : external and internal side of the outer shell layer. (A) Isochronous
profiles done close to the ventral margin. (B) Isochronous profiles done far from the ventral margin (i.e., old specimens only; see Table 1 and Fig. 7).

**Figure 4.** Comparison of the LA-ICP-MS and SN-ICP-MS trace element variations from time-series profiles on the same growth period on two adjacent slices of the C00-423-3 shell. (A) Time-series trace elements content variations. (B) Correlations between the two analytical results using reduced major axis regression software from Bohonak and van der Linde (2004).

**Figure 5.** Comparison of LA-ICP-MS trace element variations of two time-series performed on the same portion of the C00-423-5 shell, at a different distance from the external side of the shell. (A) Time-series trace elements content variations. (B) Correlations between the two LA-ICP-MS profiles using reduced major axis regression, software from Bohonak and van der Linde (2004).

**Figure 6.** Scanning electron microscope images of recent and old outer (prismatic) layer of the *Protothaca thaca* shell (specimen PC77). (A) and (C) Prisms close to the ventral margin; i.e., recent shell. The surface of the prisms is regular and composed of micrometer scales. P: one prism. S: micrometric scale at the surface of the prisms. The white spots on the picture A are small remains remained following the preparation. (B) and (D) Prisms far from the ventral margin; i.e., old shell. The surface of the prisms is granular (see picture B) because of slight dissolution (arrows on D). H: micro boring holes.

**Figure 7.** Schematic representation of the mantle zonation hypothesis and of the resulting Mg distribution along isochronous growth layer in *Protothaca thaca* shells. Ext. external side of the shell; Int.: Internal side of the shell. (A) Picture of a young and of an older *Protothaca*
thaca shell section (common scale). (B) Sketch of the margin of a shell (grey) with its secreting mantle and extrapallial fluid (blue). Red line: mantle epithelium; OE: outer epithelium; IE: inner epithelium; M: mantle. AB: animal body. (C) and (D) Enlargement of the margin of an old specimen and Mg distribution observed in an old (C) and young (D) specimen. Sketch color and legend as in (B); detail of the Mg results in the text and in Figure 3. Arrows represent the amount of Mg going through the external epithelium to the EPF and shell. Black slashes on the outer epithelium represent the hypothetic mantle zonation; the more they are tightened, the more there is magnesium in the shell.
Table 1: Summary of the *Protothaca thaca* shells studied and of the geochemical analyses carried out. When unspecified, analyses were done using LA-ICP-MS. M, U: location of the isochronous profile relative to the ventral margin of the shells. M: close to the ventral margin; U: far from the ventral margin (i.e., closer to the umbo). The number in brackets is the distance from the ventral margin. For the C00-423-5 specimen, the measures specified for the time-series profiles correspond to the distance between the external side of the shell and the time-series profile considered. See text for details.

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<td>44.5</td>
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<td></td>
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<td>PC180</td>
<td>19</td>
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<td>C00-423-3</td>
<td>51</td>
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Table 2: Laser probe operating conditions.
**Laser**

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**ICP-MS (Elan 6000)**

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<tr>
<td>Carrier Ar</td>
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**Acquisition parameters**

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<td>Peak hopping, 1 point per peak</td>
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<tr>
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Readings / Replicate 350
Numbers of replicate 1
Blanc analysis 5 s
Analysis time 60 s

Table 3: ICP-MS plasma operating conditions and acquisition parameters.

ICP-MS for solution analyses

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Fig. 1
Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 6
Fig. 7