

Does trace element composition of bivalve shells record utra-high frequency environmental variations?

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Abstract

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Saint-Pierre and Miquelon (SPM) is a small archipelago where instrumental measures based on water column velocity and temperature profiles compiled comprehensive evidence for strong near-diurnal (25.8h) current and bottom temperature oscillations (up to 11.5°C) which is possibly the largest ever observed — at any frequency — on a stratified mid-latitude continental shelf. The main objective of our study was to identify if *Placopecten magellanicus* can record on its shell these high frequency environmental variations. To this end, we have tried to identify proxies for water temperature and food availability through development of a new ultra-high resolution LA-ICPMS analyses method capable of resolving shell surface elemental composition with a 10 µm resolution. This method was applied on two shell fragments, both representing the third year of growth and 2015 annual growth period, respectively coming from two environmentally contrasted sites, more (30 m depth) or less (10 m depth) affected by high frequency thermal oscillations. Our results strongly suggest a relationship between phytoplankton biomass and barium incorporation into P. magellanicus shells at both sites. Even if *P. magellanicus* might present a physiological control of magnesium incorporation, the shape of the two Mg/Ca profiles seems to illustrate that temperature also exerts a control on magnesium incorporation in P. magellanicus shells from SPM. While U/Ca and Mg/Ca profiles show a strong positive correlation for 30 m site shell, suggesting that uranium incorporation in *P. magellanicus* shell is at least partially temperature dependent. The absence of such correlation for 10 m site shell suggests differences in uranium environmental availability or in *P. magellanicus* biomineralization between these two sites. The resolution of this new analytical method raises questions about such data interpretation related to P. magellanicus growth dynamics and physiology or individual scale based environmental measurements.

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Keywords: Ultra-high resolution LA-ICPMS; *Placopecten magellanicus*; shell chemistry; trace elements; environmental change; bivalve; environmental proxies; North Atlantic; Saint-Pierre and Miguelon; Coastal Trapped Wave.

1. Introduction

Saint-Pierre and Miquelon (SPM) is a small archipelago at the confluence of major oceanic currents, marking the boundary between the North Atlantic Ocean subtropical and subpolar gyres. However, SPM archipelago hydrodynamics is poorly known and its physic observations (sensor deployments) only began very recently. In this context, instrumental measurements based on water column velocity and temperature profiles compiled comprehensive evidence for strong near-diurnal (25.8h) current and bottom temperature oscillations (up to 11.5°C) from July to October between 10 and 80m depth. This feature is possibly the largest ever observed, at any frequency, on a stratified mid-latitude continental shelf (Lazure *et al.*, 2018). The extremely unstable physical nature of this sub-tidal environment associated with the presence of poikilothermic organisms represents a true ecological paradox, making this site a relevant place to study benthic organism responses to chronic thermal variations.

Biogenic carbonate with recognizable periodic growth bands, such as bivalve molluscs can incorporate minor and trace elements into their shells, in amounts depending on their concentrations in the environment and on the physical and biological properties of the surrounding seawater. However, bivalve shell biomineralization is a complex process, subject to strong physiological and kinetic effects related to metabolism, growth rates, ontogenetic age, shell mineralogy, crystal fabrics and organic matrix (e.g. Carré et al., 2006; Freitas et al., 2008; Freitas et al., 2016; Klein et al., 1996; Lazareth et al., 2013; Lorens and Bender, 1977; Schöne et al., 2013; Shirai et al., 2014). Owing to their wide geographic distribution, economic importance, rapid growth rates, and the presence of annual growth lines on their shell, pectinid bivalves (aka. scallops) offer good opportunities to document past environmental conditions (Chauvaud et al., 1998). The occurrence of a clearly visible annual banding pattern on the upper valve of the Atlantic sea scallop, *Placopecten magellanicus*, and the presence of this species in SPM over a wide bathymetric gradient (5 to 80 m), make this species a good candidate to track high-frequency past environmental changes - reflected as variations in the shell geochemical properties – at extremely high temporal resolution.

Spatially-resolved geochemical analysis of biogenic carbonates deposited between two accurately dated growth lines can be performed with a wide set of methods, such as laser ablation inductively coupled mass spectrometry (LA-ICPMS), secondary ion mass spectrometry (nanoSIMS) or electron micro probe analyser (EMPA). Because of its potential

for rapid and accurate high-resolution *in situ* trace element analysis at relatively low cost and minimal sample preparation requirements, LA-ICPMS has become a routine analytical tool in a wide area of research applications (Warter and Müller, 2017).

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As bivalve growth rates have often been related to environmental variables such as food availability or water temperature (Ballesta-Artero et al., 2017; Butler et al., 2010; Marali and Schöne, 2015; Witbaard et al., 1997) and because of the importance of these two parameters to track environmental and ecological changes, we then understand the interest to track and calibrate elemental proxy records of these two variables. For example, some authors proposed that magnesium to calcium ratios (Mg/Ca) can be used to record water temperature (Ullmann et al., 2013, Bougeois et al., 2014), while there are many reports of strong vital effects in bivalve shells for this element (Lorrain et al., 2005, Wanamaker et al., 2008, Surge and Lohmann, 2008). Uranium-to-calcium ratio has also been suggested as a proxy for temperature in shallow-water corals (Min et al., 1995; Shen and Dunbar, 1995) and in planktonic foraminiferal carbonates (Yu et al., 2008). Some authors also found a pH effect on U/Ca ratios in both inorganic aragonite and calcite (Kitano and Oomori, 1971; Chung and Swart, 1990). Indeed, U/Ca ratios in calcium carbonate are negatively correlated with pH and $[CO_3^{2-}]$, because in aqueous solutions the carbonate ion complexes with the uranyl ion (UO_2^{2+}) at higher pH (Langmuir, 1978), therefore less uranium is available to be incorporated in shell carbonate. Regarding U/Ca ratio on mollusc shells, Frieder et al. (2014) demonstrated that U/Ca varies as a function of pH in shells of living larvae in Mytilus californianus and Mytilus galloprovincialis. More recently, Zhao et al. (2018) demonstrated by measuring U/Ca ratio on Mya arenaria shells exposed to pCO₂-enriched environments the existence of efficient regulatory mechanisms to tightly control the pH at the site of calcification of this species. . Ba/Ca profiles in bivalve shells are typically characterized by a flat background signal interrupted by sharp peaks. Background level has been suggested to be linked with salinity (Gillikin et al., 2006, 2008). As for peaks, many authors suggested a synchronization with phytoplankton blooms (e.g. Elliot et al., 2009; Lazareth et al., 2003; Stecher et al., 1996; Thébault et al., 2009; Vander Putten et al., 2000). Building on the work of Stecher and Kogut (1999), Thébault et al. (2009) proposed two main hypotheses to explain the peaks: (1) ingestion of barite originating from assemblages of recently dead diatoms or (2) adsorption of barium onto iron oxyhydroxides associated with diatoms frustules.

The main objective of our study was to identify whether the calcitic shell of P. magellanicus can record the high frequency (25.8 h) environmental variations observed in SPM. To this end, we have developed a new ultra-high resolution LA-ICPMS analytical method in order to investigate skeletal trace element concentration with a 10 μ m resolution. Results of our investigation can contribute to a better understanding of environmental and physiological mechanisms associated to sub daily environmental variations on ions incorporation into fast growing marine bivalve shells.

2. Materials and Methods

2.1 Sample collection

Two live *P. magellanicus* were collected in September 2016 from Saint-Pierre Bay (Saint-Pierre and Miquelon – NW Atlantic) respectively at 10 m and 30 m depth (Figure 1). Both individuals were in their fourth year of growth. The deepest location consisted in a homogeneous substrate, made of compacted and stable fine sand. At the shallowest one, the substrate was more heterogeneous and consisted of a mixture of gravels, pebbles, and rocks with a seaweed cover. Soft tissues were removed immediately after collection. Both shells were carefully cleaned with freshwater to remove adherent sediment and biological tissues before sample preparation.



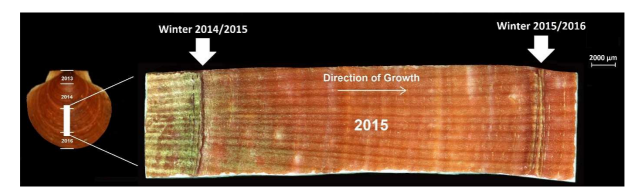
Figure 1: (A) Location of Saint-Pierre and Miquelon archipelago. (B) Satellite image of *P. magellanicus* sampling sites (red dots) in Saint-Pierre Bay.

2.2 Environmental monitoring

Annual thermal profiles at 10 m and 30 m discussed were derived from Lazure *et al.* (2018) study. To refine our vision of thermal variations on the two collection sites (Figure 1), three multiparameter probes measuring temperature every 5 minutes were deployed at 8 m, 12 m and 30 m depth, between 28/08/2017 and 15/09/2017. The 2015 monthly satellite chlorophyll *a* measurements were downloaded from the GlobColour website (http://hermes.acri.fr) and are weighted monthly averages of single-sensor products (SeaWiFS/MERIS/MODIS/VIIRSN merged chlorophyll concentrations) over the area 46.6–47.3°N / 56.0–56.6°W (i.e., waters surrounding the SPM archipelago within ca. 30 km).

2.3 Sample preparation

All micro-chemical analyses were performed on *P. magellanicus* upper valves. Indeed, the lower valves might have been contaminated as a result of a prolonged contact with the sediment. For each individual, a fragment of shell of *ca.* 3.5 cm x 1 cm was cut with a diamond saw, including the axis of maximal growth (Figure 2).



<u>Figure 2:</u> Example of one *P. magellanicus* fragment used for LA-ICPMS analyses. White arrows indicate annual shell growth lines positions defining 2015 increment.

All ultra-high resolution LA-ICPMS analyses were performed one these two shell portions. These fragments represent, for each individual, the third year of growth corresponding to 2015 annual growth periods. The outer shell layer was ultrasonically cleaned with deionized water to remove organic matter and sediment particles. In addition, before LA-ICPMS analyses, the outer shell layer of each sample was chemically cleaned with a 15 seconds acetic acid (10 %) bath, soaked in deionized water during 10 seconds, and left to air dry in the LA-ICPMS clean room.

A UV high-repetition-rate femtosecond laser ablation (fs-LA) system (Nexeya SA, Canejan, France) was employed (Pulse duration: 360fs; wavelength: 257 nm). Each ICPMS measurement point represents an ablation transect with a 1-mm long arcuate trajectory, parallel to the ventral margin, made by fast round trips of a 10 μ m spot (Figure 3). All transects were adjacent in order to analyse the whole "2015 annual period of growth" for the two individuals. The area, covered by a 1 mm x 10 μ m transect is equivalent to the area covered by a 110 μ m diameter round spot.

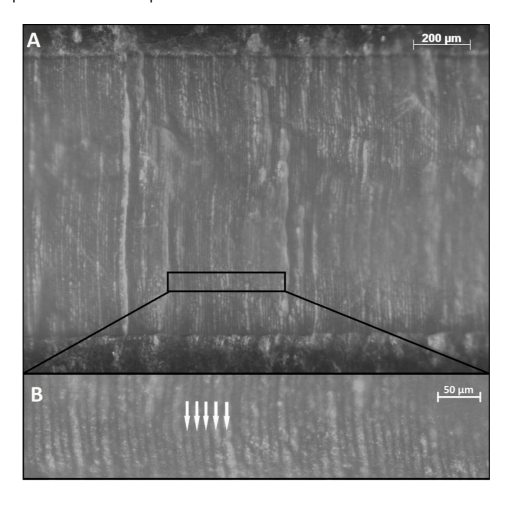


Figure 3: Post ablation picture of a 1.5 mm *P. magellanicus* section showing ca. 150 femtosecond laser ablation transects (A). Zoom on a small fraction of them, each white arrow points to a laser ablation transect (B). The visible lines represent ridges generated during the laser ablation process.

Outer shell layers were analysed for Mg/Ca, Ba/Ca and U/Ca ratios using a high-resolution inductively coupled plasma mass spectrometer fitted with a jet interface (Element XR, Thermo

Scientific, USA). A helium gas stream carried ablated material to the HR-ICP-MS (carrier gas flow rate 0.68 L.min⁻¹). Elemental ratios were quantified by monitoring 43 Ca, 24 Mg, 138 Ba, and 238 U. Calcium was used as an internal standard. Elements were standardized to calcium based on the stoichiometry of calcium carbonate (388 000 μ gCa.g⁻¹) outer shell layer), assuming 100 % CaCO₃: Mg/Ca (μ g.g⁻¹), Ba/Ca (μ g.g⁻¹) and U/Ca (μ g.g⁻¹). Quantification of trace elements in otoliths was achieved by external calibration using both carbonate pellets FEBS-1 (Barats et al., 2007) and 2 NIST glass standards (610, 612) to ensure the best accuracy. Each standard was analysed three times before and after each session with the laser to account for drifting during the day. The limits of detection (μ g.g-1 in shells) achieved in this study were 0.08, 0.01 and 0.002 for 24Mg, 138Ba, and 238U, respectively. They were based on a 3 σ criterion, where σ is the standard deviation of the mean blank count for each isotope. All the elemental concentrations in the outer shell layer were above the detection limits.

3. Results

3.1 Environmental parameters:

Around 10 m depth, the temperature varied from 2 °C in May to a maximum of 16 °C in early September and then decreased to 8 °C in November. At this depth, seawater temperature presents a classic seasonal cycle with cold water intrusions (Figure 4). During the first two weeks of September at 8 m and 12 m in Saint-Pierre Bay, temperature showed high-frequency variations with cold water incursions leading to 4°C (8m) to 6°C (12m) amplitudes (Figure 4). Along these two weeks, temperatures were 70% of the time above 12 °C (Figure 4).

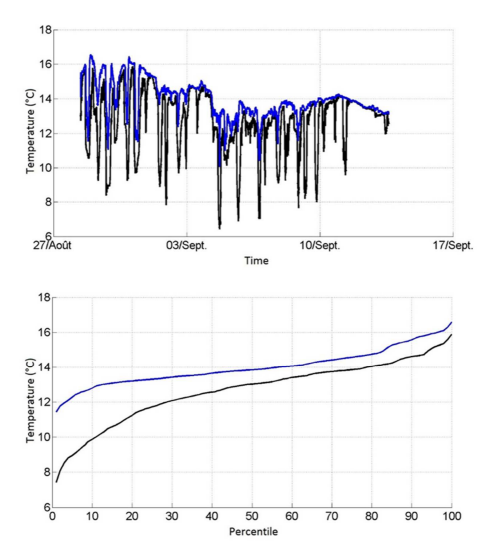
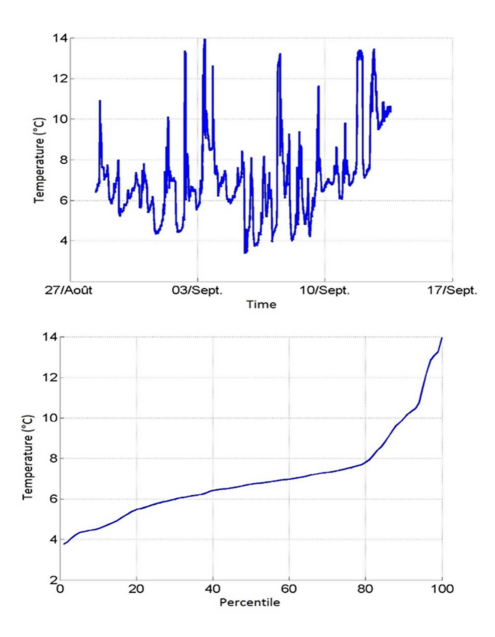


Figure 4: End of August and first two weeks of September 2017 time series of seawater temperatures at 8 m and 12m depth (blue and black) (top graphic). Percentile distribution of these temperatures at 8 m and 12m depth (blue and black).

At 30 m depth, the temperature annual profile was radically different (Figure 5). Seawater temperature baseline is mainly cold over the year, showing low seasonal amplitudes. However, during the stratified period, temperatures showed high-frequency variations whose amplitude increased with sea-surface temperature. During the first two weeks of September (Figure 5), oscillations were the largest in term of amplitude, reaching nearly 10°C. Along these two weeks, temperatures were 80% of the time below 8 °C (Figure 5).



<u>Figure 5:</u> End of August and first two weeks of September 2017 time series of seawater temperatures at 30m depth (top graphic). Percentile distribution of these temperatures during these two weeks.

Monthly mean satellite chlorophyll *a* concentrations ranged from 0.46 to 1.22 mg.m⁻³ (Figure 6). The annual time series-exhibited a background level around 0.7 mg.m⁻³, with one major peak in April - May 2015.

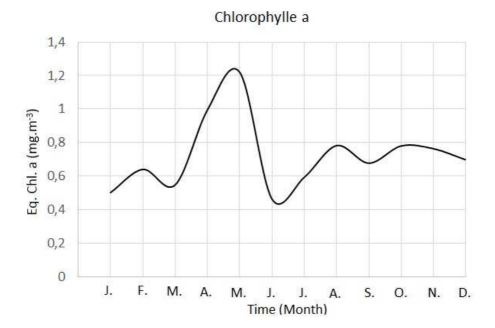


Figure 6: Monthly satellite equivalent chlorophyll a (mg.m⁻³) measurements over the year
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3.2 Ba/Ca ratio profiles in the shell carbonates

Outer shell layer Ba/Ca ratios ranged from 0.61 to 15.71 $\mu g.g^{-1}$ at 10 m and from 0.43 to 12.54 $\mu g.g^{-1}$ at 30 m (Figure 7). Both series have the same profile with one major peak occurring respectively 370 μ m and 830 μ m after the "winter 2014/2015" growth line. The main Ba/Ca peak covers 1370 and 2220 μ m of shell at 10 and 30 m, respectively. A secondary smaller Ba/Ca peak occurred immediately after the first one, covering respectively 1230 and 1160 μ m at 10 and 30 m. Ba/Ca baseline was the same for the two time-series (around 1.5 $\mu g.g^{-1}$).

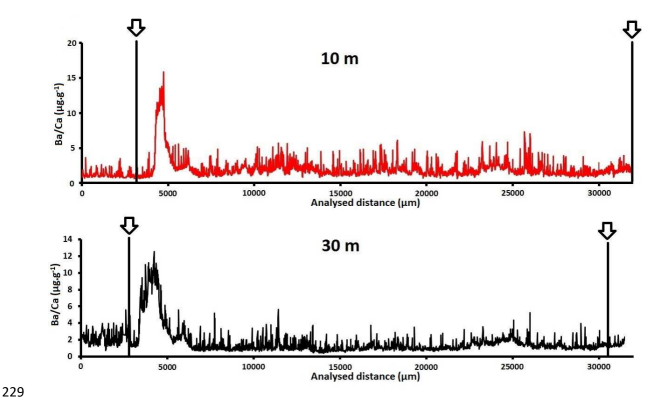
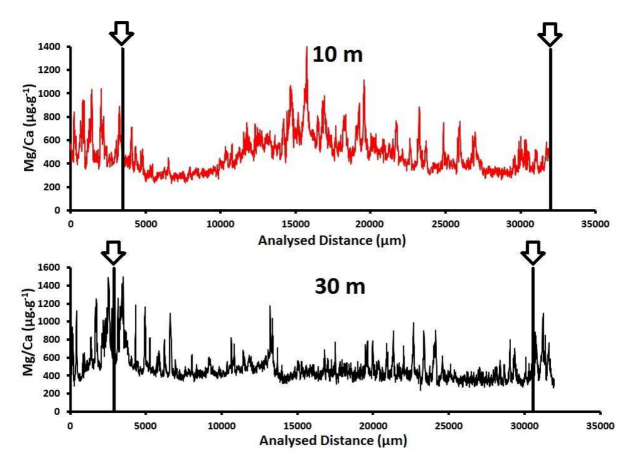


Figure 7: Ba/Ca (μg.g⁻¹) series at 10 m (red curve) and 30 m (black curve). Vertical lines placed under the arrows indicate the position of winter shell growth lines.

3.3 Mg/Ca ratio profiles

Outer shell layer Mg/Ca ratios ranged from 232 to 1408 $\mu g.g.$ at 10 m and from 233 to 1495 $\mu g.g.$ at 30 m (Figure 8). At 10 m, Mg/Ca profile followed a sinusoidal pattern with stronger high frequency variations between 15 000 and 25 000 μ m (top and decreasing phase, Figure 8). At 30 m depth, the Mg/Ca profile was radically different with a globally flat profile between the two growth lines and high frequency variations mainly between 15000 and 25000 μ m (Figure 8).



<u>Figure 8 :</u> Mg/Ca (μ g.g⁻¹) series at 10 m (red curve) and 30 m (black curve). Vertical lines placed under the arrows indicate the position of shell growth lines.

3.4 U/Ca ratio profiles

Outer shell layer U/Ca ratios ranged from 0.003 to 0.16 μ g.g-1 at 10 m and from 0.004 to 0.36 μ g.g-1 at 30 m (Figure 9). At 10 m depth, U/Ca time series had a relatively flat pattern with high-frequency variations all along the profile. At 30 m depth, U/Ca profile was close to Mg/Ca one. So we decided to compare those profiles in the next paragraph.

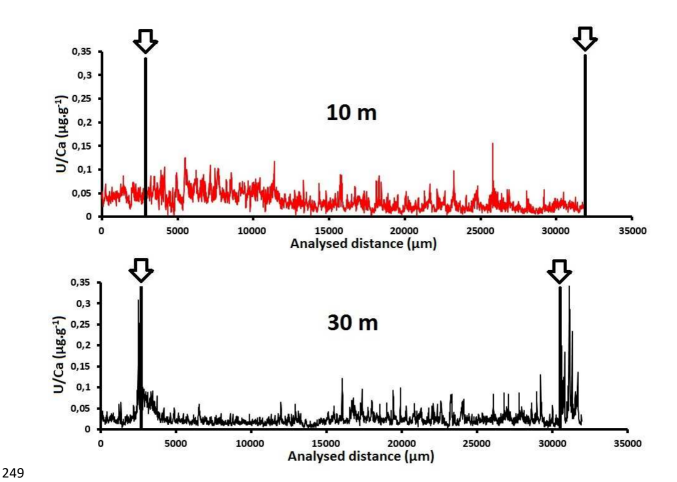


Figure 9: U/Ca ($\mu g.g^{-1}$) series at 10 m (red curve) and 30 m (black curve). Vertical lines placed under the arrows indicate the position of shell growth lines.

3.5 U/Ca and Mg/Ca comparison at 30m depth:

At 30 m depth, U/Ca and Mg/Ca profiles presented a strong positive correlation (N = 3135, r = 0.62, p < 0.001) (Figure 10). A closer examination of this relationship on a shorter time window (i.e. three high frequency cycles, between 22400 μ m and 24250 μ m) revealed an even stronger correlation (N = 186, r = 0.77, p < 0.001).

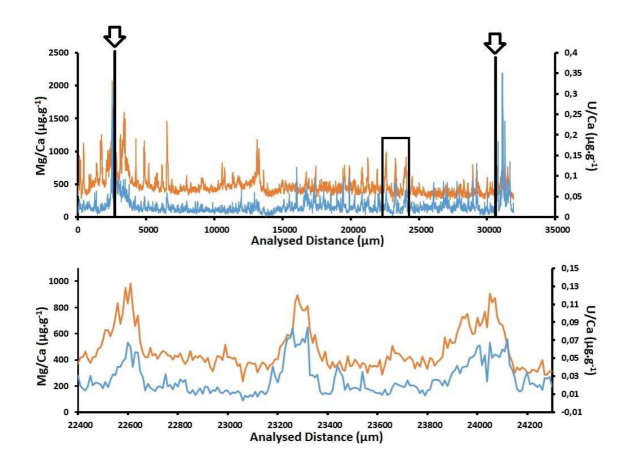


Figure 10: U/Ca ($\mu g.g^{-1}$) (blue curve) and Mg/Ca ($\mu g.g^{-1}$) (orange curve) series at 30 m. Vertical lines placed under the arrows indicate the position of shell growth lines. The graph bellow represents a close-up on three peaks (black box in the upper graph).

4. Discussion

This paper presents the first chemical analyses performed on P. magellanicus shells. Regarding the temporal resolution (25.82 h) of the environmental phenomena we wanted to track, it was necessary to develop a new analytical method. Our novel approach using ultra-high resolution fs-LA-ICPMS enables trace element analyses in bivalve shells with a 10- μ m resolution. This study gave us first insights about P. magellanicus ability to record high-frequency environmental variations within is shell.

4.1 Barium

The high degree of similarity between the two Ba/Ca profiles suggests that the occurrence of Ba/Ca peaks was controlled by one or multiple common environmental factors. The pattern

of these two Ba/Ca profiles is similar to those observed in cross sections of other bivalve species (e.g., Gillikin et al., 2008; Stecher et al. 1996; Vander Putten et al., 2000). This confirms the hypothesis which suggests that the choice of analyzing the shell surface or the outer shell layer in cross sections do not have significant influence on Ba/Ca records in shells. In this section, we will discuss several hypotheses to explain temporal variability of Ba/Ca in P. magellanicus shell. Background levels of Ba/Ca time-series in bivalve shells have been suggested to record salinity variations (e.g. Gillikin et al., 2006). There is generally a linear inverse relationship between seawater salinity and dissolved barium concentrations (Coffey et al., 1997; Gillikin et al., 2006). However, variability in seawater dissolved barium concentrations as a source for the Ba/Ca peaks in *P. magellanicus* from SPM is highly unlikely. Indeed, SPM islands, due to their offshore status, are not subjected to major riverine inputs and associated variations in salinity (Poitevin et al., 2018). Salinity usually ranges from 31.3 to 32.2 PSU (see Figure 4 in Poitevin et al., 2018) without a clear seasonal trend and, therefore, cannot explain Ba/Ca peaks measured in the shells. Many authors suggested a close relationship between these Ba/Ca peaks and phytoplankton blooms (e.g. Elliot et al., 2009; Lazareth et al., 2003; Stecher et al., 1996; Thébault et al., 2009; Vander Putten et al., 2000). In our study, the high similarity of chlorophyll α concentration (Figure 6) and Ba/Ca (Figure 7) profiles strongly suggest a relationship between phytoplankton biomass and barium incorporation into P. magellanicus shells. Indeed, the occurrence of this bloom, in May 2015, seems to be consistent with the starting of P. magellanicus annual growth from other Canadian regions (Chute et al. 2012; Kleinman et al., 1996). Elevated levels of suspended barite (BaSO₄) have been suggested to be linked with oceanic diatoms primary productivity (Dehairs et al., 1991). Most of the barium released by diatoms after a bloom is labile and only a minor fraction eventually forms barite crystals (Ganeshram et al., 2003). Therefore, if labile barium, either in phytoplankton or released into the dissolved phase, was the cause of the Ba/Ca peaks, these peaks should form near the end of the bloom or very shortly thereafter (Gillikin et al., 2008). Considering the absence of daily growth lines in P. magellanicus, we cannot conclude about chlorophyll a and Ba/Ca peaks timing. Finally, the two Ba/Ca profiles exhibited a double peak, with a first large amplitude one and a smaller second peak. This observation has also been made in P. maximus Ba/Ca profiles (Gillikin et al., 2008). One explanation for this double peak proposed in this study is based on Ganeshram et al. (2003). They found that barite formation can take several weeks to reach its maximum after the

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beginning of phytoplankton decay. Barite may be formed at the sediment surface and be ingested by *P. magellanicus* several weeks after the phytoplankton bloom ends.

These observations on Ba/Ca incorporation in *P. magellanicus* shell from SPM imply a real need for complementary information related to local *P. magellanicus* growth dynamics and physiology. It would also be crucial to get insights about the nature and the quantity of benthic and pelagic primary production over the year and along a bathymetric gradient.

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4.2 Magnesium

In bivalve shells, the relationship between seawater temperature and Mg/Ca ratio is still subject to controversy. Some authors proposed that Mg/Ca ratios can be used to record water temperature (e.g., Bougeois et al., 2014; Lazareth et al., 2003; Mouchi et al., 2013; Surge and Lohmann, 2008; Ullmann et al., 2013), while there are many reports of strong vital effects in bivalve shells for this element (e.g., Elliot et al., 2009; Lorrain et al., 2005, Wanamaker et al., 2008,). In this study, we can hardly discuss the importance of vital effects on trace elements incorporation into P. magellanicus shell. Indeed, our analyses were only carried out on one year of growth (ontogenetic and calendar) and one individual per site. The only thing we can say about physiological control of Mg incorporation in P. magellanicus shell is based on Mg/Ca level. In this study, the mean Mg/Ca ratio (~500 μg.g-1 corresponding to ~2mmol.mol-1) of the calcitic outer shell layer of P. magellanicus corresponds to a low value compared to other calcitic mollusc shells (Lazareth et al., 2007 and references therein). Given the absence of sclerochemical studies about trace element incorporation in P. magellanicus shells, we can only try to explain these low Mg concentrations relying on studies based on other calcitic bivalves with low Mg/Ca concentrations. For example, Lorens and Bender (1977) suggested that Mytilus edulis biologically regulates the amount of Mg entering the extrapallial fluid to produce low-Mg calcite. Perhaps a similar process occurs in P. magellanicus, suggesting a physiological control of Mg incorporation that could obscure Mg/Ca and seawater temperature relationship. This confirms the need for additional investigations on biomineralization, e.g. through experiments in controlled environments, in order to better understand trace elements incorporation in *P. magellanicus* shell. However, the shape of the two Mg/Ca profiles tend to highlight kind of a temperature control on magnesium

incorporation in our shells. At 10 m, the sinusoidal pattern of Mg/Ca ratio may reflect the seasonal seawater temperature annual cycle at 10 m depth. While at 30 m, the Mg/Ca profile presents a relatively flat baseline with high-frequency variations, which could mirror the seawater seasonal temperature trend, namely showing low seasonal amplitudes with highfrequency variations (Lazure et al., 2018). Other studies also point to the non-systematic relationship between Mg/Ca ratio and SST. From a one year study of *M. edulis* growth, Vander Putten et al. (2000) observed a positive correlation between Mg/Ca and SST only during spring. Small-scale variations in Mg concentrations in M. edulis calcite have also been shown to derive from Mg being concentrated along the margins of calcite prisms (Rosenberg et al., 2001). Indeed, the absence of interannual growth lines on the P. magellanicus shell is problematic to convert analysed distances in time. That is why enhance our knowledge on P. magellanicus growth dynamics along this bathymetric gradient in SPM would help us decipher physiological and environmental effects on trace element incorporation in P. magellanicus shell calcite. In addition, the lack of high frequency environmental data limits our ability to fully interpret our results and confirms the interest to set up a high frequency observatory along this bathymetric gradient.

4.3 Uranium

In our study, U/Ca and Mg/Ca profiles show a strong positive correlation in shells collected at $30 \,\mathrm{m}$ (N=3135, r=0.62, p<0.001) (Fig. 10). However, this is not the case for the shallowest shells for which no significant correlation could be found. These results suggest that (i) environmental uranium availability for P. magellanicus are not the same between the two sites and/or (ii) that physiological differences between P. magellanicus from 10 m and 30 m sites could lead to differential incorporation of uranium in shells.

Since (i) we do not have information about *P. magellanicus* physiological differences between these two depths, and (ii) only few studies previously investigated U/Ca ratio as a potential paleo environmental proxy in bivalve shells (Frieder *et al.*, 2014; Gillikin and Dehairs, 2012; Zhao *et al.*, 2018), it seems difficult to draw conclusions about the kind of processes influencing uranium incorporation in *P. magellanicus* shells.

To our knowledge, U/Ca ratio as a paleo environmental proxy has been studied for the first time in mollusc shell by Gillikin and Dehairs (2012). In this study the authors tried to investigate U/Ca in Saxidomus gigantea shell as a potential acidification proxy and concluded that U/Ca may not reflect environmental variability and did not function as a paleo-pH proxy. More recently, Zhao et al. (2018) also found virtually unchanged U/Ca values in Mya arenaria shells with increasing seawater pCO_2 up to 2900 μ atm. However, in the same study the authors found a significant increase in U/Ca ratio in shells with the increase in seawater pCO_2 at 6600 µatm. These findings reveal the existence of certain compensatory mechanisms by which this species may partially mitigate the impact of high environmental pCO₂ on shell formation through modifying the calcifying fluid chemistry to maintain its pH homeostasis (Zhao et al., 2018). These conclusions lead us to consider P. magellanicus physiological responses induced by the repeated thermal variations occurring at the 31m site and their impact on the calcifying fluid pH of this species. Considering our study purpose and resolution, it seems difficult to conclude about U/Ca as a potential acidification proxy in P. magellanicus shell, especially as we do not have pH measurements on our study sites. Uranium-to-calcium ratios have also been suggested as a proxy for temperature in shallow water corals (e.g., Min et al., 1995; Shen and Dunbar, 1995) and in planktonic foraminiferal carbonates (e.g., Yu et al., 2008). In our study, the positive correlation between U/Ca and Mg/Ca profiles in the shell collected at 30 m would support this hypothesis. However, this correlation does not hold anymore at 10 m, suggesting that variations in uranium bioavailability differs between our two sites. Indeed, microorganisms have the ability to adsorb radionuclides/metals through extracellular binding involving physical adsorption, ion exchange, complexation and precipitation (Acharya et al., 2009). They also sequester the metal ions by passive/active transport to the interior of the cell, followed by its accumulation. Microbial cells have been shown to reduce, oxidize, adsorb, accumulate and precipitate uranium (Fredrickson et al., 1999; Macaskie et al., 2000). So differences in microbial communities between the two sites, related to the nature of the habitat or to depth, could lead to changes in environmental uranium availability and finally to shell U/Ca ratios.

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5. Conclusion

Our novel approach using ultra-high resolution fs-LA-ICPMS enables trace element analyses in bivalve shells with a $10-\mu m$ resolution. This study gave us first insights about *P. magellanicus* ability to record high-frequency environmental variations within is shell at a sub-hourly scale.

From an analytical point of view, it would be interesting to continue this study by applying this new analytical technique to more individuals. This would allow us to discuss about interindividual variability within those two sites. Moreover, combining this approach with nano-SIMS δ^{18} O measurements (temperature proxy) would help us to get insights about the temperature control of Mg and U incorporation in shells.

In term of data interpretation, these results also confirm a real need for complementary information. Some of them should be related to *P. magellanicus* intra-annual growth dynamics. Indeed, in this study the absence of visible intra-annual growth lines visible on *P. magellanicus* shells hindered the temporal alignment of our microchemical data. Others must concern *P. magellanicus* physiological responses and their impacts on the calcifying fluid chemistry of this species. All these additional studies will require multiple high frequency environmental data continuously recorded at an individual scale within those two sites.

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